# Water Quality Criteria Data Book Volume 3

**Effects of Chemicals on Aquatic Life** 

ENVIRONMENTAL PROTECTION AGENCY RESEARCH AND MONITORING

#### WATER POLLUTION CONTROL RESEARCH SERIES

The Water Pollution Control Research Series describes the results and progress in the control and abatement of pollution in our Nation's waters. They provide a central source of information on the research, development, and demonstration activities in the Environmental Protection Agency, through inhouse research and grants and contracts with Federal, State, and local agencies, research institutions, and industrial organizations.

Inquiries pertaining to Water Pollution Control Research Reports should be directed to the Chief, Publications Branch, Research Information Division, R&M, Environmental Protection Agency, Washington, D.C. 20460.

# Water Quality Criteria Data Book - Vol. 3

EFFECTS OF CHEMICALS ON AQUATIC LIFE Selected Data From the Literature Through 1968

by

Battelle's Columbus Laboratories

for the

### ENVIRONMENTAL PROTECTION AGENCY

Project No. 18050 GWV Contract No. 68-01-0007

May 1971

For sale by the Superintendent of Documents, U.S. Government Printing Office, Washington, D.C. 20402 - Price \$3.75

### **EPA Review Notice**

This report has been reviewed by the Environmental Protection Agency and approved for publication. The data are listed as reported in the literature without collaboration or evaluation of their validity. Therefore, these data must and cannot be used indiscriminately for the establishment of water quality criteria for the aquatic environment. These data should be used only as a guideline for the base of action. Approval does not signify that the contents necessarily reflect the views and policies of EPA, nor does mention of trade names or commercial products constitute endorsement or recommendation for use.

### ABSTRACT

Original data from more than 500 technical publications concerning the specific effects of chemicals on individual species of aquatic biota were collected and summarized in uniform format. Alphabetical assembly of the data by chemical allows rapid access to considerable detailed information. A Species Index facilitates search for information on the toxicity of chemicals to individual aquatic species.

The details of major procedures in laboratory bioassay and field assessment of chemical toxicity in water are discussed. Freshwater and marine procedures are included. A total of approximately 1000 references were utilized in preparing this report.

Recommendations include:

- (1) Establishment of an information-analysis center on chemical water pollution based to some extent on the report prepared.
- (2) Preparation of a listing of chemical constituents of effluents and continued up-dating of this list.
- (3) Development of a pattern of bioassays for evaluating the effects of a chemical on aquatic life. Data from these evaluations would be used in developing mathematical models for predicting chemical toxicity in a wide range of environmental circumstances.
- (4) Development of *in situ* bioassay procedures for more realistic assessment of chemical toxicity to aquatic life.

# TABLE OF CONTENTS

Section		Page	
Ι	Introduction	1	
II	Objectives	4	
III	Literature Search and Bibliographies	5	
IV	Fish Bioassay	6	
V	Bioassay of Aquatic Organisms Other Than Fish	18	
VI	Biochemical Oxygen Demand (BOD) and Related Microbiological Procedures	19	
VII	Marine Bioassay	25	
VIII	Field Assessment	26	
IX	Factors Affecting Chemical Toxicity in Water	43	
Х	Industrial Wastes		
XI	Extracted Data – The Effect of Chemicals on Aquatic Biota		
XII	Summary and Conclusions	66	
XIII	Recommendations	69	
XIV	Bibliography	70	
XV	Appendices		
	A. Chemicals and Mixtures of Chemicals	A-1	
	B. Commercial Chemical Products	B-1	
	C. Species Index	C-1	
	D. Identification of Commercial Chemicals	D-1	

## LIST OF FIGURES

Figure		Page
1	Food Web in Western Lake Erie Leading to the Sheepshead Fish	27

### LIST OF TABLES

Table		Page
1	Fish Used in Bioassays, Frequency of Use, and Type of Water in Which They Occur	12
2	Laboratory Methods for Studying the Effect of Chemicals on Fish Other Than Bioassay Lethality	15
3	A Partial Listing of References Using Freshwater Aquatic Organisms Other Than Fish for Bioassay	18
4	Toxicity of Various Compounds as Determined by BOD	23
5	Collecting Equipment in Common Usage in Limnological Studies and the General Purpose for Which Each is Used	34
6	Partial Listing of Organisms Commonly Associated With Pollution	35
7	Thermal Death Points of Fish Acclimized at the Indicated Tempera- tures (Freshwater = F, Marine $-$ Atlantic = A, Pacific = P)	45
8	Minimum Oxygen Values at Various Temperatures at Which Fish Can Exist Under Laboratory Conditions	51
9	Usual Fisheries Hazards of 30 Common Types of Municipal and Industrial Effluents	56
10	General Comments on Selected Industrial Effluents	57

### SECTION I

### INTRODUCTION

The internal and external chemical environment of an organism determines whether that organism will survive, grow, and perpetuate itself. Internal chemical balance is mediated by the genetic makeup of the organism, the external chemical milieu in which it lives, and all other environmental factors. The effect of chemicals on living organisms is an especially important factor in aquatic environs where organisms are in intimate contact with chemicals in solution and suspension. Water passes into and through the body of an organism primarily via the integument, membranes, gills, or mouth. Toxic chemicals in the water may cause immediate lethality although in many instances sublethal quantities of deleterious chemicals may be accumulated within the body. In time, the chemical residues in an organism may cause drastic effects of varying types, also including mortality. Complicating this situation is the effect of chemicals on lower animal forms which provide part or all of the food chain leading to higher aquatic organisms. Thus, sport fish may leave polluted areas not to avoid chemical pollutants or to escape death but rather to seek food, for example, when bottom fauna upon which they feed are obliterated. Low dissolved-oxygen concentrations in water caused by release of oxygenconsuming chemicals can also have equally drastic impact on aquatic organisms.

This then is the basic problem today in water pollution and is the primary subject of this report. A closely related problem, considering aquatic biota as indicators of chemical toxic effect, is the consideration of whether or not such water is safe for use by humans. At the moment fish bioassay appears to be the best method available for determining the toxic effect of chemicals on aquatic life.

In a report section entitled "Recommendations for the Use of Bioassays and Application Factors to Denote Safe Concentrations of Wastes in Receiving Streams", the National Technical Advisory Committee (Interim Report, 1967), has made the following recommendations in part for the use of bioassays:

"1. For the determination of acute toxicities, flow-through bioassays are the first choice. Methods for carrying out these flow-through tests have been described by Surber and Thatcher, 1962; Lemke and Mount, 1963; Henderson and Pickering, 1963; Jackson and Brungs, 1966; Mount and Warner, 1963; Mount and Brungs, 1965; and Brungs and Mount, 1967. Flow-through bioassays should be used for unstable volatile or highly toxic wastes and those having an oxygen demand. They also must be used when several variables such as pH, DO, CO<sub>2</sub> and other factors must be controlled.

2. When flow-through tests are not feasible, tests of a different type or duration must be used. The kinds of local conditions affecting the procedure might be single application of pesticides or lack of materials and equipment.

3. Acute static bioassays with fish for the determination of  $TL_m$  values should be carried out in accordance with Standard Methods for the Examination of Water and Waste Water. Such tests should be used for the determination of  $TL_m$  values only for persistent, nonvolatile, highly soluble materials of low toxicity which do not have an oxygen demand as it is necessary to use the amount added as the concentration to which the test organisms are exposed.

4. When application factors are used with  $TL_m$  values to determine safe concentrations of a waste in a receiving water, the bioassay studies to determine  $TL_m$  values should

be made with the most sensitive local species and life stages of economic or ecological importance and with dilution water taken from the receiving stream above the waste outfall. In the absence of knowledge concerning the most sensitive of the important local species or life stages or due to difficulty in providing them in sufficient numbers, other species whose relative sensitivity is known can be used or tests may be carried out using one species of diatom, one species of an invertebrate and two species of fish, one of which should be a pan or game fish. Further, these bioassays must be performed with environmental conditions at levels at which the waste is most toxic. Tests should be repeated with one species at least monthly and when there are changes in the character or volume of the waste.

5. Concentration of materials with noncumulative toxic effects should not exceed 1/10 of the 96-hour TL<sub>m</sub> value at any time or place. The 24-hour average of the concentration should not exceed 1/20 of the TL<sub>m</sub> value. For toxicants with cumulative effects, the concentrations should not exceed 1/10 and 1/100 for the above respective values."

The need for water of better quality by improved pollution control has been chronicled broadly with considerable justification in news media, scientific journals, and government reports. The result of this attention has been the establishment of water quality criteria and federal requirements for states, localities, and consequently industries to set minimum water standards within certain time limits, and to enforce these standards. The basic Federal Water Pollution Control Act (1956) was provided and later amended in 1961, by the Water Quality Act of 1965, and by the Clean Water Restoration Act of 1966. In the years given, these amendments were approved as public laws. Water quality requirements are becoming more stringent each year. Carpenter (1968) has outlined federal policy and organization in regard to this problem. In Water Quality Criteria (1968), the various problems of water pollution control are discussed in detail and recommendations are made for measures to improve pollution management. Earlier, these and related problems were discussed in publications by the National Research Council (1966), the Department of Health, Education, and Welfare (Public Health Service Publication No. 999-WP-25, 1965), ORSANCO (Ohio River Valley Water Sanitation Commission, 1967), and the Environmental Pollution Panel (1965). Establishment of water quality criteria in the U.S. has been recently considered by the Aquatic Life Advisory Committee (1955, 1956, 1960), the American Society for Testing Materials (Katz and Woelke, 1967; Woelke, 1967), Bartsh and Ingram (1959, 1966), Carter (1968), Ettinger and Mount (1967), Okum (1968), Smith (1961), Tarzwell (1957, 1959, 1962), Weston (1964), and Wilhm and Dorris (1968). The Manufacturing Chemists Association (1967) listed the sources of information on water quality criteria. The number of meetings increases each year as announced in such periodicals as Water and Sewage Works. The problems of industrial water utilization and effluent management of chemical wastes are generally discussed by Bower (1965), Cairns (1965, 1967), in Public Works (Anonymous, 1968), and in various texts, as well as briefly in the section of this report entitled "Industrial Wastes". Engdahl and Croxton (1962) have discussed the economics of pollution, a matter further treated in such journals as Chemical Week and Chemical and Engineering News.

Eutrophication of lakes is a special pollution problem that is not discussed in this report. Excellent documents pertaining to eutrophication are by Fruh, et al (1966) and bibliographies by the U.S. Public Health Service (Mackenthun, 1962, 1965). Similarly, thermal effluents were not considered as a topic for this report, due primarily to the magnitude of research in this field. Useful, extensive bibliographies have been recently published, including ones by the American Society for Civil Engineering (1967), Kennedy and Mihurksy (1967), Raney and Menzel (1967), and Wurtz and Renn (1965).

Another special problem is pesticide contamination of the environment. This is discussed to a considerable extent throughout this report, but especially in the section "Field Assessments". Reviews or general references concerning the effect of pesticides in the environment or other agricultural problems of this nature include an article in <u>Environmental Sciences and Technology</u> (Anonymous, 1968); papers by Cottam (1961), Langer (1964), Moore (1967), and Robinson (1967); and periodicals such as <u>Residue Reviews</u> (Springer-Verlag New York Inc., Vol 1<sup>+</sup>, 1962<sup>+</sup>) and Pesticides Documentation Bulletin (U.S. Department of Agriculture, Vol 1<sup>+</sup>, 1965<sup>+</sup>).

Other useful reference sources on trends in water pollution control are the chemical industry trade journals, <u>Chemical Week and Chemical and Engineering News</u>, and such publications as the <u>Conservation Foundation Letter</u>, and the <u>Environmental Health Letter</u> (Vol 1<sup>+</sup>, 1961<sup>+</sup>).

This is something of the background in which this report was prepared in late 1968 and early 1969.

### SECTION II

### **OBJECTIVES**

The objectives of this program were to:

- (1) Collect and summarize in standardized format the available information from the scientific literature concerning:
  - (a) The specific effects of chemicals on individual species of aquatic biota. (This study was limited to studies of single chemicals or simple mixtures of chemicals and does not include industrial effluents that contain highly complex chemical mixtures.)
  - (b) Details of the procedures and environmental factors important in the observation or the measurement of these effects.
- (2) Review the existing information on aquatic life as it is applicable or related to the study of water pollution.
- (3) Review the methodology used in studying the effects of chemicals on aquatic life.

### **SECTION III**

### LITERATURE SEARCH AND BIBLIOGRAPHIES

Some 3500 papers, mostly from the period 1950 through 1968, were screened and about 2000 obtained for direct examination. Foreign language publications were not included. About 500 contained original data, from which extracts were prepared (Appendices A and B). An attempt was made to be comprehensive for the years 1958 through 1968 with only selected references included preceding this period. Of these selected references, the majority were published after 1950, with only a few being from the older literature.

The primary source for identifying the references used in this study were the literature reviews published annually by the Water Pollution Control Federation Committee in the Journal of the Water Pollution Control Federation (1958-1968), which proved to be excellent. The reference list was checked against Chemical Abstracts, Biological Abstracts, Water Pollution Abstracts, and numerous recent special subject bibliographies. Very few additional references were added to the list from these other sources. Personal visits were made to selected governmental and industrial organizations to secure pertinent data. Information was also requested from the Science Information Exchange (Smithsonian Institution) and National Referral Center (Library of Congress). Letter requests for publications not commonly available were sent to a number of scientists in this field.

### SECTION IV

### FISH BIOASSAY

Fish bioassay of industrial wastes and other potentially toxic materials has evolved in the past 50 or so years from nonstandardized procedures by individual scientists to the present where standardized assay procedures now are available to researchers in this field. Early work on fish bioassays was done in Europe and Asia nearly 60 years ago. Pioneering work in the U.S. on developing procedures and methods for bioassay of fish was conducted by Shelford, Bilding, Carpenter, and Ellis. In 1945, Hart, Doudoroff, and Greenbank in a book now out of print described a standardized fish bioassay procedure, which Doudoroff, et al (1951) recommended as a standard method for use by industry, government agencies, and others. This method with comparatively few modifications, e.g., continuous flow exposure of fish in addition to static exposure, has been widely used and today is used more or less in its original form. The fish bioassay procedure outlined in the 12th edition of Standard Methods (American Public Health Association, 1967) is basically that described by Doudoroff, et al. Procedures developed by W. E. Martin of the Pesticides Regulation Division and by Burdick (1960) at the N.Y. Conservation Department are quite similar. A prepublication copy of fish bioassay procedures that is to appear in the forthcoming 13th edition of Standard Methods (1971) was kindly provided by Professor M. C. Rand. The following discussions are based primarily on this document.

### Static Bioassay

Briefly, the static bioassay procedure can be described as follows:

- (1) After determination of an approximate toxic range of a chemical or effluent, appropriate concentrations are prepared on a logarithmic or geometric scale within the toxic range.
- (2) Small (5.0-7.5 cm) fish, which have been quarantined 10-30 days (min-max) to assure no disease problems and acclimatized to the chosen assay water, are placed in the chemical or effluent solutions prepared with dissolved oxygen in concentrations not less than 4 mg/l (warm water fish) and 5 mg/l (cold water fish) at a constant temperature. Temperatures of 25 ± 2 C and 15 ± 2 C are recommended for warm water and cold water species, respectively.
- (3) Observation and recording are made of dead fish which should be removed at 8, 24, 48, and 96 hours after the assay is initiated. Notation of other effects, such as intoxication, distress, loss of equilibrium, and other abnormal behavior, should also be made.
- (4) Calculation or estimation of a TL<sub>50</sub> or  $TL_m$  for various time periods is made by interpolation of the data plotted on semilogarithmic coordinate paper.

The  $TL_m$  of a compound is not considered as representing the concentration of a chemical or effluent that is safe in fish habitats. It is merely a relative measure of the acute, lethal toxicity of the material to a certain fish under controlled environmental conditions and must be used with a mathematical application factor to determine safe concentrations of effluents to be released. This has been discussed by Doudoroff (1951), Warren and Doudoroff (1958), and the National Technical Advisory Committee (1967).

A further distinction between LD50, LC50, and EC50 is made in the prepublication copy of <u>Standard Methods</u> as follows:

"The expressions 'lethal dose' (LD) and 'lethal concentration' (LC) have also been frequently used, the term 'lethal dose' often incorrectly. The expression 'lethal dose' is not appropriate when designating a certain concentration in an external medium, inasmuch as a dose, strictly speaking, is a measured quantity administered. Unlike 'lethal dose' and 'lethal concentration', the term 'tolerance limit' is universally applicable in designating a level of any measurable lethal agent, including high and low temperatures, pH, and the like. The expression 'effective concentration' (EC) applies to concentrations only and is generally used in connection with effects other than death."

The APHA procedure describes in excellent detail the selection and preparation of fish and diluent water, effluent samplings or preparation-dilution of test substances, use of aeration, controls, etc.

Static, acute fish bioassay has been shown to be inadequate for estimating the effect of chemicals on fish. Lack of reproducibility between laboratories is the rule rather than the exception. Reasons for this include chemical and microbiological degradation of toxic compounds, volatility of some compounds, utilization of oxygen by microorganisms as well as by fish, water quality variability, accumulation of fish metabolic by-products in assay containers, and uptake of toxicants by the test animals.

Periodic (daily or more often) renewal of test solutions is a variation of the static, acute fish bioassay that can be utilized to overcome some of the objections of this type of evaluation. Continuous test solution renewal must be used in long-term, chronic exposures of fish to chemical solutions where sublethal effects are to be studied. This variation is recommended in the <u>Standard Method</u> especially "when there is evidence or expectation of a rapid change of toxicity of the test solution".

Also recommended in the procedure is the determination of temperature, DO, and pH of the samples under evaluation at various times during the experiment as well as of the chemical properties or dissolved mineral content of the diluent water. To quote, "A rather complete mineral analysis of the water is advisable". Furthermore, chemical analysis for the toxicant under study is suggested throughout the exposure period. Seldom is this type of information reported in the literature as is shown and discussed in subsequent sections of this report.

The U.S. Fish and Wildlife Service, Circular 185 (1964) describes static bioassay procedures in relation to piscicide studies being carried out by the U.S. Bureau of Sport Fisheries and Wildlife. Freeman (1953) discussed use of standardized diluent water in static bioassay of fish and aquatic invertebrates. Other authors have also discussed or used synthetic or defined water for bioassays (Cairns and Scheier, 1955, 1958, 1963, 1968; Doudoroff, 1956; Dowden and Bennett, 1965; Fitzgerald et al, 1952; Trama, 1955; and Whitley, 1968). Handling and maintenance of bioassay fish was described by Hunn, et al (1968). A number of authors have discussed mathematical treatment of fish toxicity data including Burdick (1957) and Henderson and Tarzwell (1957). Excellent general discussions of static fish bioassays have been published by Burdick (1960, 1967), Cairns (1957, 1966), McCall (1961), Tarzwell (1959), Wuhrmann and Woker (1959), and Wuhrman (1955). Cope (1961) suggested standards for reporting fish toxicity tests which apparently have not been accepted widely. Essentially his appeal dealt with correct identification, size, and condition of the test fish; complete description of the procedure involved and of chemical, physical, and biological factors; volume of water and number of fish for that volume; etc. Many of these data are lacking in most of the papers reviewed in the present report.

### **Continuous Flow Bioassay**

The majority of the factors discussed under static bioassay apply to the continuous flow procedure with the added requirement of automatic intermittent or continual metering of the test substance dissolved or suspended in diluent water into the test chambers and continuous flow-through of water. Problems associated with dissolved oxygen and test chemical content in static exposures can be obviated in the continuous flow technique since the water added contains these materials in constant concentrations.

Briefly, a continuous flow system is composed of:

- (1) Diluent water reservoir from which water flows into the
- (2) Constant head diluent supply where the water is cooled or heated to the desired temperature and then metered along with
- (3) The effluent or toxicant (added with a chemical pump, Mariotte bottle, etc.) into
- (4) The test container in which fish are exposed, and which
- (5) Overflows into an appropriate drain.

An acclimatizing tank for test fish can also receive water from the reservoir and constant head diluent supply. Water flow is by gravity and the recommended flow rate is equal to a complete volume change of test containers in 6 hours.

Data are taken usually over a 5-day period and plotted as for the static bioassay. Five-day supplies of water and toxicant are required.

The procedure as it is outlined allows ample latitude for assembling the apparatus according to individual requirements. As guides, the work of Jackson and Brungs (1966), Surber and Thatcher (1963), Lemke and Mount (1963), Mount and Warner (1965), and Mount and Brungs (1967), and others are referred to. These reports deal in part with information concerning valve control systems, chemical metering pumps, serial dilution apparatus, and the proportional diluter as utilized in various types of studies.

The earliest paper found on continuous flow bioassay was by Merkens (1957), a British scientist, who devised an automatically controlled apparatus for monitoring and adjusting temperature, pH, dissolved oxygen, and toxicant concentration in the test water added. This system was ingenious for its time.

Alabaster and Abram (1965) have more recently described British continuous flow techniques. Flow rate is adjusted to maintain an adequate level of dissolved oxygen. The apparatus and treatment of data are described in considerable detail. Other recent procedures or innovations on the continuous flow technique have been reported by Betts, et al (1967), Burke and Ferguson (1968), Grenier (1960), Hendersen and Pickering (1963), and Solon, et al (1968).

The use of the continuous flow procedure in chronic exposures (Mount, 1962, 1968; Mount and Stephan, 1967), piscicide development (Parker and Wurth, 1965), residue accumulation (Holden, 1966), tracer studies (Holden, 1962), spawning (Mount and Stephan, 1967), and avoidance (Foster, 1967; and Warner et al, 1966) is discussed in other sections of this report.

Burdick (1960, 1967) and Jackson and Brungs (1966) have thoroughly discussed the continuous flow technique and its applicability to current water pollution problems. There can be no doubt that continuous flow fish bioassay simulates the field situation more closely than does static bioassay.

### **Fish Selection**

The selection of fish for bioassay depends in part on the species of appropriate size available for study that can be maintained in the laboratory and also on the native fish present in the receiving water under study. Lennon (1967) has recommended development of inbred strains of test fish for standard reference in much the same manner as inbred mouse strains are used in mammalian toxicology. Cope (1966) has also made similar recommendations.

Small, preferably juvenile, fish are generally used so that sufficient numbers may be accommodated in the laboratory. Mount (1968) has briefly listed fish species that might be used as appropriate test organisms. This listing was prepared at the National Water Quality Laboratory, Duluth, Minnesota. The fish were selected on the basis of the following criteria:

- (1) Sport, commercial or forage value
- (2) Potential for exposure to pollution
- (3) Geographical distribution and abundance
- (4) Suitability for laboratory studies
- (5) Existing knowledge in regard to toxicity.

The fish selected were:

Primary list – all pollutants Threadfin shad (Dorosoma petenense) Brook trout (Salvelinus fontinalis) Rainbow trout (Salmo gairdneri) Northern pike (Esox lucius) Emerald shiner (Notropis atherinoides) Fathead minnow (Pimephales promelas) White sucker (Catostomus commersoni) Channel catfish (Ictalurus punctatus) White bass (Roccus chrysops) Bluegill (Lepomis macrochirus) Largemouth bass (Micropterus salmoides) Yellow perch (Perca flavescens)

Special list - for selected pollutants

Coho salmon (Oncorhynchus kisutch)
Lake trout (Salvelinus namaycush)
Mountain whitefish (Prosopium williamsoni)
American smelt (Osmerus mordax)
Smallmouth bass (Micropterus dolomieui)
Walleye (Stizostedion vitreum)

The goldfish (Carassius auratus) was the selected equivalent of the "white rat".

Hunn, et al (1968) list the bioassay species used by the Bureau of Sport Fisheries and Wildlife as follows:

Rainbow trout (Salmo gairdneri) Brown trout (Salmo trutta) Brook trout (Salvelinus fontinalis) Lake trout (Salvelinus namaycush) Northern pike (Esox lucius) Goldfish (Carassius auratus) Carp (Cyprinus carpio) Fathead minnow (Pimephales promelas) White sucker (Catostomus commersoni) Black bullhead (Ictalurus melas) Channel catfish (Ictalurus punctatus) Green sunfish (Lepomis cyanellus) Bluegill (Lepomis macrochirus) Smallmouth bass (Micropterus dolomieui) Largemouth bass (Micropterus salmoides) Yellow perch (Perca flavescens) Walleye (Stizostedion vitreum)

Henderson and Pickering (1963) state that many species are suitable for bioassays, including:

Guppy (Lebistes reticulatus) Mosquito fish (Gambusia affinis) Goldfish (Carassius auratus) Fathead minnow (Pimephales promelas) Bluegill (Lepomis macrochirus) On the basis of research usage as determined by the papers reviewed in the present study, an even wider variety of fish has been used experimentally. These, along with their frequency of use and type of water in which they may be found, are summarized in Table 1. Only those found in more than one paper are listed.

### Chronic Bioassay

Evaluation of sublethal concentrations of various chemicals in long-term fish exposures is probably the most reliable bioassay method for determining safe levels at which chemicals may be released into receiving water. The exposure may be either static in which periodic solution renewal is required or continuous flow in which the concentration of the chemical is maintained at a constant level. The latter is by far the method of choice. Both procedures have been discussed in previous sections.

### **Chronic Static Exposure**

A few recent papers serve to illustrate the variations that may be employed in conducting this type of exposure. The long-term effect of a 2-hour exposure to Dieldrin on the reproduction of guppies (*Lebistes reticulatus*) was studied by Hubble and Reiff (1967) over a 12-month period. The fish were placed in a standardized water following the exposure. No harmful effect on reproduction was observed.

Weiss and Gakstatter (1964) studied the long-term effect of various pesticides on acetylcholinesterase activity of bluegill, golden shiner, and goldfish by daily replenishing the test solutions over periods up to 30 days. The pesticides studied could be detected at concentration levels down to 0.1 x  $10^{-3}$  mg/l.

Test water containing subacute concentrations of copper or zinc was used by Grande (1967) to expose trout eggs, fry, and fingerlings. The test solutions were renewed during 28-day periods every second day in experiments with eggs and daily for young trout.

The effect of sublethal concentrations of Dieldrin on laboratory populations of guppies (*Poecilia reticulata*) in aquaria was studied by Cairns, et al (1967). Weekly renewal of test solutions over a 14-month period was employed.

Dugan (1967) studied the combined effects of sublethal concentrations of detergents and pesticides on goldfish. The test water was cleaned by filtering periodically and the chemical concentrations adjusted to desired levels. Four-month exposure periods to the surfactants and up to 51-day exposure periods to Dieldrin were studied. Synthetic water and 100-gal epoxy-coated, galvanized water tanks were used.

In a study of the effect of Diquat on bluegill and bluegill food organisms, Gilderhus (1967) exposed the animals to the chemical during a 24-week period with varied frequencies of sublethal concentrations.

None of these authors used the static, acute fish bioassay procedure outlined in <u>Standard</u> Methods.

# TABLE 1. FISH USED IN BIOASSAYS, FREQUENCY OF USE, AND TYPE OF WATERIN WHICH THEY OCCUR

### (Freshwater = F; Marine - Atlantic = A, Pacific = P)

\_

Scientific Name	Common Name	Occurrence	
Abramis brama	Bream	F	
Ambloplites rupestris	Rock bass	F	
Ameiurus nebulosus	Brown bullhead	F	
Brachydanio rerio	Zebrafish	F	
Campostoma anomalum	Stoneroller	F	
Carassius auratus*	Goldfish	F	
C. carassius	European carp	F	
Catastomus commersoni*	White sucker	F	
Cyprinodon variegatus	Longnose killifish	Α	
Cyprinus carpio*	Carp	F	
Ericymba buccata	Silverjaw minnow	F	
Esox lucius	Northern pike	F	
Eucalia inconstans	Brook stickleback	F	
Fundulus similis	Striped mullet	A	
Gambusia affinis*	Mosquitofish	A-F	
Gasterosteus aculeatus*	Threespine stickleback	A-F-P	
Gobio gobio	Gobie	F	
Hyborhynchus notatus	Bluntnose minnow	F	
Ictalurus melas*	Black bullhead	F	
I. natalis*	Yellow bullhead	F	
I. nebulosus*	Brown bullhead	F	
I. punctatus*	Channel catfish	F	
Lagodon rhomboides	Pinfish	А	
Lebistes reticulatus*	Guppy	F	
Leiostomus xanthurus	Spot	Α	
Lepomis auritus	Redbreast sunfish	F	
L. cyanellus <b>*</b>	Green sunfish	F	
L. gibbosus*	Pumpkinseed	F	
L. macrochirus**	Bluegill	F	
L. megalotis	Longear sunfish	F	
L. microlophus*	Redear sunfish	F	
Micropterus dolomieui*	Smallmouth bass	F	
M. salmoides*	Largemouth bass	F	
Mugil cephalus	Striped mullet	А	
Notemigonus crysoleucas*	Golden shiner	F	
Notropis atherinoides	Emerald shiner	F	
N. cornutus	Common shiner	F	
N. hudsonius	Spottail shiner	F	
N. lutrensis	Red shiner	F	
N. stramineus	Sand shiner	F	
N. umbratilis	Redfin shiner	F	
Oncorhyncus kisutch*	Coho salmon	P-F	
O. tshawytscha*	Chinook salmon	P-F	
Perca flavescens*	Yellow perch	F	
Petromyzon marinus*	Sea lamprey	A-F	
Phoxinus phoxinus*	Red-sided shiner	F	
Pimephales notatus*	Bluntnose minnow	F	
P. promelas**	Fathead minnow	F	
Rhinichthys atratulus	Blacknose dace	F	
Rutilus rutilus Selves existenciet	Roach	F	
Salmo gairdneri**	Rainbow trout	A-F-P	
S. salar*	Atlantic salmon	A-F	
S. trutta#	Brown trout	A-F	
Salvelinus fontinalis*	Brook trout	A-F	
S. namaycush	Lake trout	F	
Semotilus atromaculatus •	Creek chub	F	
Stizostedion vitreum*	Walleye	F	

All species listed were found in two or more papers.

-----

\*Found in more than 5 papers. \*\*The most commonly used species.

### **Chronic Continuous Flow Exposure**

Brown, et al (1968), Butler (1965, 1967), Cairns and Scheier (1963), Cope (1965), Jensen and Gaufin (1966), Mount (1962, 1968), Mount and Stephan (1967), Olsen and Foster (1958), Raymont and Shields (1964), Surber and Thatcher (1963), and Weiss (1965) have utilized continuous flow techniques of their own creation for the study of a variety of aquatic organisms in long-term, continuous flow exposure to a variety of chemicals. Exposure periods up to 11 months were employed in these studies. The reports cited above represent less than 5 percent of the total number of papers from which data were extracted for Appendices A and B.

Generally, chemicals are toxic at lower concentrations in continuous flow exposures, especially long-term ones, than in static exposures. Furthermore, nonlethal effects occur more readily in continuous flow bioassays. For example, Mount (1968) reported for this type of bioassay that the "safe concentration" was 3-7 percent of the 96-hour TL<sub>m</sub> (static exposure) in studying the chronic toxicity of copper to fathead minnows. Furthermore, Mount and Stephan (1967) have stated that the biologically safe concentrations for Malathion and butoxyethanol ester of 2,4-D as determined in a continuous flow, chronic study are 1/45 and 1/9, respectively, of the 96-hour TL<sub>m</sub> for each of these compounds as determined in static bioassay. However, Cairns and Scheier (1963) found in a study of the acute and chronic effects of sodium alkyl benzene sulfonate on sunfish that results from the two types of exposure at equivalent concentrations of ABS were quite close although not identical.

As further requirements to improve water quality are imposed, the need for chronic continuous flow data concerning the effects of sublethal concentrations of potential pollutants on aquatic biota will increase.

### In situ Bioassay

The need for standardizing fish bioassay laboratory procedures has led to environmental laboratory conditions unlike those found in streams and lakes. Factors such as fluctuating sunlight, temperature, DO, pH, pollutant and nutrient concentration, etc., cannot be taken into account or compensated for in the laboratory. *In situ* evaluation of a chemical solution in the stream or body of water in which it is to be released is a method of determining with an improved degree of accuracy the concentration effects of a discharge released into that particular body of water. Exposures to the chemical in question of native species of fish can be conducted by means of portable live cars, cages, plastic pools, or raceways. Thus, the fish species of concern for a given stream can be studied in conditions approaching their particular complex ecological situation.

There is no standard procedure for this type of bioassay, but it has been employed to some extent as briefly discussed later in the section, "Field Studies". Burdick (1967) has recommended this approach and pointed out that automated water quality monitoring equipment now available can provide continuous recording of physical and chemical changes in water conditions which may allow correlation of bioassay data with ecological conditions. Raceways with disposable vinyl liners are used in advanced evaluation of piscicides as well as 9-10-ft-diameter vinyl wading pools with bottom soils of various types, pond or ground waters, aquatic plants and invertebrates, fish, and amphibians, as required. Hawskley (1967) speculated on the advent of "continuous bioassay" in which effluent and receiving water in varied ratios will be circulated into and out of test containers and noted that this almost of necessity will have to be performed at the plant site. Standard method fish bioassays are conducted in this laboratory in conjunction

with routine chemical analyses and analyses with an atomic absorption spectrophotometer. Hawkins stated that a mobile unit for conducting fish bioassays and chemical analyses at the plant site was in the design stage in 1964. A mobile bioassay unit was used in developing selective larvicides for control of sea lamprey (Howell and Marquette, 1963). Automatic water quality monitors can provide continuous and depth-profile data acquisition for water temperature, dissolved oxygen, pH, conductance, dissolved chlorides, oxidation-reduction potential, and turbidity. These parameters are indirect but excellent physical-chemical indicators of water pollution. In conjunction with fish bioassays, they can provide data suitable for mathematical modeling and simulation. More than 200 monitors of this type are now in operation in the United States. The monitor can be housed in a trailer for portability. Weather data recording for air temperature, solar radiation, wind speed and direction, and total precipitation can be integrated into the continuous recorder.

### Fish Responses Other Than Bioassay Lethality

Methods for laboratory study of fish response to chemicals in freshwater environments vary nearly as much as the number of investigators in this field of research. These range from simple observations (as suggested in <u>Standard Methods</u> and other sources); to sophisticated determinations of chemical residues, ACHE blood content, etc.; to the highly sophisticated Conditioned Avoidance Response Apparatus (CARA). These methods are identified in Table 2. One of these procedures may become a "standard method" for aquatic laboratory studies, but this does not appear likely to occur in the near future. Standard static and continuous flow fish bioassay methods will probably remain the principal laboratory tools for developing toxicity data with chronic exposures becoming more widely used. Some of the methods, notably, the avoidance, life stage, fish tissue culture, and CARA techniques, may be very useful in determining more precisely the "safe concentration" levels for chemical effluent release. Texts, such as those by Brown (1957) describe physiological methods for studying fish. Some of these methods would be highly applicable to the study of the effect of chemicals on aquatic life and could form the basis for the development of new procedures.

### TABLE 2. LABORATORY METHODS FOR STUDYING THE EFFECT OF CHEMICALS ON FISH OTHER THAN BIOASSAY LETHALITY

Туре	Comments	References
Observations of abnormal behavior	<ul> <li>Observations may be made on the following: Quiescence, excitability, or irritability Surfacing or sounding Tetanic or flaccid movement</li> <li>Swimming – erratic, convulsive, gyrating, inverted on side, etc.</li> <li>Changes in pigmentation</li> <li>External mucosa – exudate, shedding, etc.</li> <li>Integument hemorrhagia</li> <li>Rate of respiration – slow, irregular, gulping, etc.</li> <li>Gill hemorrhaging or mucous discharge</li> <li>Defecating or regurgitating mucous or other material</li> <li>Sensitivity to stimuli such as light, sound, touch, electric probe, etc.</li> <li>Moribundity – distended operculum, opaque eyes, etc.</li> <li>Recovery – complete, or not.</li> </ul>	Brown, et al (1968), Cairns, et al (1967), Cope (1966), Fromm and Schiffman (1958), Grindley (1946), Mount (1962), and Olsen and Foster (1958)
Autopsy and histology	Tissue and organ pathology are studied by appropriate methods. Decrease of glycogen and RNA, tissue dissociation, necrosis, lesions, and secretions may also be noted.	Blumenkratz (1956), Cairns (1966), Cairns and Scheier (1963), Cope (1965), Eng. Science, Inc. (1964), Gilderh (1967), Herbert and Shurben (1964), Mount (1964), Moun and Stephen (1967), Van Val et al (1968), and Warner, et a (1966)
Avoidance	Raceways or similar laboratory structures are generally used so that a chemical solution can be metered into the bioassay water to estab- lish a concentration gradient. Fish have been trained to discriminate between very low con- centrations of selected chemicals.	Cairns (1957), Costa (1965), Hasler and Wisby (1949), and Ishio (1965)
Growth retardation	Chronic exposure was the most effective tech- nique utilized.	Crandall and Goodnight (1962), Olsen and Foster (1958), and Royer (1966)
Residue analysis	Following exposure, organs of the fish are removed and analyzed for specific chemical content. This technique is used most often in studies of pesticide accumulation, and is also quite useful in field studies to show previous exposure. Whole fish homogenates have also been analyzed as well as animal feeds and processed sea foods prepared from various types of marine fish species.	Butler (1965, 1967), Cope (1965), Eisler (1967), Gilderhus (1966, 1967), Godsil and Johnson (1968), Holden (1966), Mahdi (1956), Moubry, et al (1968) Mount (1962), Mount and Stephan (1967), Pagan and Hageman (1950), Ullman, et al (1961), Weiss (1965), and Welch and Spindler (1964)

Туре	Comments	References	
Acetylcholinesterase (ACHE) activity of brain	This method is used primarily in the study of organophosphorus pesticides in both labora- tory and field studies of freshwater and marine types. The utility of this method is somewhat limited because of its near specificity for organophosphates.	Butler (1965), Cope (1965), Fromm and Schiffman (1958), Weiss (1959, 1961, 1964, 1965), and Weiss and Gakstatter (1964)	
Radiotracers	This technique is used primarily in the study of pesticides and metal ions where labelling can be successfully accomplished. Tissue and organ analyses of radiotracer accumulation have been conducted. Among the radio-isotopes used in fish studies are Ca <sup>45</sup> , Cl <sup>4</sup> , P <sup>32</sup> , and Zn <sup>35</sup> . Acetates, chlorides, Bayer 22408, DDT, Dieldrin, Dimethoate, Lindane and Parathion are some of the compounds studied in this manner. Wet combustion of tissues and measurement of Cl <sup>4</sup> O <sub>2</sub> release has also been employed.	Butler (1965), Douglas and Irwin (1963), Fujiya (1965), Gakstatter and Weiss (1967), Holden (1962), Joyner (1961), Marchetti (1965), Miller, et al (1966), and Schmidt and Weidhaas (1961)	
Effects on various life stages of fish	Effects of chemicals on sperm, eggs, yearling, and adult fish as well as fry are often studied to determine the relative resistance of these life stages to chemicals. Embryos from fertilized eggs have also been studied with the finding that fertilized egg membranes provide some resistance to the effects of chemicals.	Cairns and Scheier (1959), Cope (1966), Crandall and Goodnight (1962), Goodman (1951), Grande (1967), Hiltibran (1967), Marchetti (1965), Mount (1968), Piavis (1962), and Skidmore (1966)	
Spawning (reproductive behavior)	This may be studied in the laboratory by pro- viding suitable objects, such as pieces of cement-asbestos tile; and proper environ- mental conditions, including a controlled photoperiod, for this activity. Spawning in several studies was shown to be affected by concentrations of chemical much lower than those for the $TL_m$ (96 hr). A "Laboratory Fish Production Index" (LFPI) has been proposed and is gaining acceptance.	Cairns, et al (1967), Cohen, et al (1961), Gilderhus (1967), Holden (1966), Hubble and Reiff (1967), Mount (1962, 1968), and Mount and Stephan (1967)	
Swimming or cruising speed and oxygen consumption while swimming	Specifically designed raceways, cages, or "current trays" are required to determine rate of speed. Oxygen utilization can be determined by means of an oxygen- consumption chamber or respirometer. This is a useful technique for studying fish larger than fry. Current velocity can be controlled and is an important factor in studying large fish which require sufficient speed for oxygen transfer in their gills.	Cairns and Scheier (1963), Doudoroff and Warren (1962), Herbert and Shurben (1963), Mount (1962), and Ogilvie and Anderson (1965)	
Chemical resistance of fish	After sublethal exposure, fish acquire specific resistance to certain chemicals. This has been demonstrated in the laboratory and the field most frequently for pesticides and metals.	Boyd and Ferguson (1964), Darsie and Corriden (1959), Fairchild (1955), Ferguson, et al (1954, 1955), and Mount (1968)	

TABLE 2. (Continued)

Туре	Comments	References	
Blood studies	Changes in erythrocyte count, hemoglobin, sodium and calcium levels, microhematocrit, and hematocrit have been used in a variety of studies. The latter has been suggested as a measure of the state of health of bioassay fish prior to testing.	Cairns and Scheier (1963), Cope (1965, 1966), Gilderhus (1967), Hatch (1957), and Hunn, et al (1968)	
Glucose transport	This is an <i>in vitro</i> type of study using dissected fish gut.	Stokes and Fromm (1965)	
Fish tissue culture	Epithelial cells of fathead minnow cultured on modified Eagle's MEM medium, were found to have a reduced mitotic index at the calculated "safe concentration" of zinc. It was concluded that one-tenth of the 96-hr $TL_m$ is probably closer to the safe concentration.	Rachlin and Perlmutter (1968)	
Environmental stress	Reduced DO or increased temperature caused increased toxicity of various chemicals.	Cairns (1957), Lloyd (1961), and Pickering (1968)	
Thermal acclimatization	In studies of the effect of DDT on salmon, it was found that DDT interferes with the normal thermal acclimation mechanism. Fish exposed to 10 ppm DDT and acclimated to warm water were extremely sensitive to cold water. Acclimatization also affected chemical toxicity.	Cope (1963, Keenleyside (1958) and Greer and Paim (1968)	
Fish taste	The taste of sport fish can be drastically changed by chemical pollutants.	Hynes (1966) and Rachlin and Perlmutter (1968)	
Conditioned avoidance response apparatus (CARA)	Toxicant-induced behavior of fish exposed to sublethal concentrations of chemicals was studied in raceways by means of photo- graphing the fish at various intervals and calculating response in terms of relative position. A large mirror facilitated photog- raphy. At concentration levels $1/2000$ of the 96-hr TL <sub>m</sub> value for tetraethyl pyro- phosphate (TEPP), aberrant behavior of goldfish was noted. A ratio of $1/25$ was obtained for Toxaphene.	Eng. Science, Inc. (1964) and Warner, et al (1966)	

### SECTION V

# BIOASSAY OF AQUATIC ORGANISMS OTHER THAN FISH

Surprisingly few aquatic organisms other than fish have been used as test organisms in bioassays. The organisms most commonly used are numerous species of algae and the crustaceans, *Daphnia magna* and *D. pulex*. Other freshwater invertebrates used in bioassays include protozoa (*Paramecium* and *Tetrahymena*), planaria (*Planaria* and *Dugesia*), crustacea (*Gammarus*), gastropods (*Lymnaea* and *Physa*), stonefly and mayfly naiads, and caddisfly and midge larvae. Oysters and shrimp are the principal test animals other than fish in marine bioassays. The oyster (*Ostrea*) are quite sensitive to low concentrations of some chemicals as determined by retarded shell growth. The brown, pink, and white shrimp (*Penaeus*) are the most commonly used crustacea in seawater bioassays. Barnacles (*Balanus*) are also used. These are discussed in the section, Marine Bioassay. Table 3 is a listing of references using various organisms other than fish for freshwater bioassay studies.

Procedures developed by C. M. Palmer and T. E. Maloney (1955) at the Taft Engineering Center in Cincinnati, Ohio, and by G. P. Fitzgerald, et al (1952, 1958, 1963) are widely used for laboratory study of freshwater algae.

There are no generally accepted or standard procedures for bioassays using these other types of organisms, although the procedures developed by Bertil Anderson (1944, 1945, 1948, 1960) in his studies of *D. magna* are commonly used.

In evaluating papers from which data were extracted (Appendices A and B), it was evident that a much broader spectrum of species are studied in the field than under laboratory conditions.

Туре	References
Algae: (Chlorella pyrenoidosa, Microcystis aeruginosa, and numerous other species)	Abram (1967), Alabaster and Swain (1963), Beak (1958), Elson and Kerswill (1967), Ganelin, et al (1964), Holden (1964), Hopkins, et al (1966), Kallman, et al (1962), Kemp, et al (1966), Khan (1964), Merkens (1958), Nejedly (1967), Palmer and Maloney (1955), and Sprague, et al (1965)
Invertebrates: (Daphnia magna, D. pulex, Gammarus pulex, Culex spp, etc.)	<ul> <li>Abram (1967), Anderson (1946), Burdick (1965), Cairns, et al (1965) Chadwick (1960), Clarke (1947), Fromm (1965), Gaufin (1961), Gaufin, et al (1961), Henderson, et al (1961), Ingols (1959), Kable (1957), Naylor (1965), Shaw and Grushkin (1967), Sprague (1965) Tarzwell (1957), Tarzwell and Henderson (1960), Turnbull, et al (1954), Weiss and Botts (1957), Wilber (1965), Williams (1964), and Wood (1957)</li> </ul>
Vertebrates: (Rana pipiens, R. catesbieana, Bufo valliceps – sperm, eggs, tadpoles, and adults)	Cairns, et al (1965), Lackey (1957), Shaw and Grushkin (1967), and Stroud (1967)

TABLE 3.	A PARTIAL LISTING OF	REFERENCES	USING	FRESHWATER	AQUATIC	ORGANISMS
	OTHER THAN FISH FOR	BIOASSAY				

### SECTION VI

### BIOCHEMICAL OXYGEN DEMAND (BOD) AND RELATED MICROBIOLOGICAL PROCEDURES

### **Biochemical Oxygen Demand**

The biochemical oxygen demand (BOD) test is a test which is designed to determine the relative oxygen requirement of a municipal and/or industrial effluent. The determination of BOD of an effluent for the purpose of regulating the rate of discharge into a stream or sewerage system with minimal adverse effects on the oxygen resources of the receiving water will be at best an analytical starting point. BOD has several very limiting criteria which must be adequately understood for this technique of possible waste dilution to be useful. The procedure for BOD determinations as described in the 13th Edition of the Standard Methods for the Examination of Water and Waste Water (American Public Health Association, 1967) provides the basis for this discussion. This procedure has been essentially the same for more than 10 years with comparatively minor changes.

Although basically a simple bioassay to execute, the exceptions and precautions given in the BOD procedure make it somewhat formidable to the uninitiated. Briefly without specific details, the procedure consists of:

- (1) Microbial seeding (if needed) of appropriate water dilutions of the chemical or effluent and initial determination of the dissolved oxygen (DO) of the sample by the iodometric method, azide modification. Sample dilutions are prepared with distilled water saturated with dissolved oxygen and buffered at pH 7.2 with a phosphate buffer solution.
- (2) Incubation of the seeded samples at 20 C for 5 days and in darkness in standard BOD bottles which are water-sealed to exclude oxygen.
- (3) DO determination of the diluted samples after the 5-day incubation period. The most reliable results are said to be for that dilution which shows a residual DO of at least 1 mg/l and a depletion of at least 2 mg/l. For toxic chemicals or effluents, toxic effect is indicated by lack of oxygen utilization by the microorganisms. When the lag period for microbial growth is prolonged, incubation periods of up to 20 days or longer may be employed.
- (4) When substances are evaluated that are oxidizable by molecular oxygen, then an immediate dissolved oxygen demand (IDOD) should be determined and taken into consideration when calculating the BOD. The IDOD is a short-term assay in which DO is determined 15 minutes after the sample is added to the dilution water.

Carbon compounds utilizable by aerobic microorganisms, oxidizable nitrogen compounds utilizable by nitrogen bacteria, and certain chemical reducing compounds (ferrous iron, sulfites, sulfides, and aldehydes) are the three main types of chemicals that influence oxygen demand. The latter can be taken into consideration by the IDOD determination. Solubility and volatility of chemicals must also be considered. Some organic wastes are not oxidizable and thus are not amenable to the BOD bioassay. When such wastes are suspected, chemical oxygen demand (COD) and total carbon (TC) analyses would be conducted for comparison with BOD results.

According to the procedure: "In many cases, particularly in food processing wastes, a satisfactory seed may be obtained by using the supernatant liquor from domestic sewage which

has been stored at 20 C for 24-36 hr", but it goes on to state that "acclimated" seed and receiving water below a point (2-5 miles) of effluent discharge may be used since "many industrial wastes contain compounds which are not amenable to oxidation by domestic-sewage seed". If the concern is with dissolved oxygen depletion, then an "acclimated" seed would seem most appropriate whether it is acclimated in the laboratory or collected downstream from a discharge. If the concern is with the toxic level of an effluent, then both acclimated and domestic-sewage seed evaluations might be made to establish a type of index for safe discharge. In the event of evaluation of a new type of discharge, seed acclimated in the laboratory to that particular discharge undoubtedly would be most desirable.

In regard to the amount of seed to be added, it is stated that, "Only past experience can determine the actual amount of seed to be added per liter." It would be more precise to add exact amounts of seed, e.g., Zintgraff, et al (1968) added 0.5-2.0 mg/l of seed in their studies.

The BOD bioassay suffers as do most laboratory procedures from lack of correlation between laboratory results and those obtained in the field. The need for a standardized procedure is recognized, but many factors enter into the behavior of a chemical in the aquatic environment that cannot be taken into account in the laboratory. Some of these objectionable features are alluded to and briefly discussed in <u>Standard Methods</u>, but others are overlooked and should be considered in attempting to apply the results of BOD determinations. The principal uncontrolled variable in the BOD procedure is the nonstandardized microbial inoculum or acclimated microbial seed as the case may be. Briefly, other factors include:

- (1) Temperature and pH seldom is the aquatic environment at precisely one temperature or pH.
- (2) Fluctuating solids and dissolved solids content in receiving water these can greatly influence the effect of a chemical on aquatic biota.
- (3) Algae although BOD determinations are conducted in a dark incubator, algae can grow heterotrophically and utilize oxygen, as do bacteria and other microorganisms. Dead algal cells can also affect BOD. Wisniewski (1958) has discussed the effect of algae on BOD determinations and DO in streams.
- (4) Protozoa these are known to be present in domestic sewage seed, and according to Bhatla, et al (1965) protozoa are responsible for approximately 30 percent of the BOD exerted under normal seeding conditions in 5-day BOD tests.
- (5) Total aquatic biomass all plants and animals other than the ones discussed above significantly influence the effect of chemicals on the aquatic environment.
- (6) Mixed nutrient substrates these are the rule rather than the exception in receiving water.
- (7) Mixed toxicants in sublethal concentrations already present in receiving water this problem has received comparatively little attention as judged by reports in the literature. Exceptions in non-BOD studies are the pesticides where the effect or accumulation of mixtures of these compounds and their decomposition products on and in aquatic biota have been documented. Additive, antagonistic, or synergistic effects probably do occur.
- (8) Photochemical oxidation by ultraviolet from sunlight.

- (9) Mixing due to currents the BOD laboratory assay is static and therefore no mixing occurs.
- (10) Other factors briefly mentioned in various papers as important in oxygen depletion are reduction of nitrates, anaerobic microbial alteration of organic compounds, secondary oxygen uptake, and decomposition of chemical intermediates.

All of these factors should be recognized by the analyst who should take them into account when applying data from BOD determinations.

BOD can be utilized to advantage by an experienced researcher in determining the oxygen depletion potential or the effect on microorganisms of an effluent containing toxic chemicals. Both are important considerations in effluent management for minimal effect on receiving waters.

On studying the various papers concerned with reporting BOD data, it was found that a wide variety of methods for reporting the data are utilized. As examples, Ingols (1954, 1955, 1956) plotted BOD values to show oxygen depletion in percent of control BOD with increasing concentrations of mercuric chloride, copper, zinc, etc., in ppm. Oberton and Stack (1957) using acclimated seed in studying the BOD of acrolein, diethanolamine, and methyl vinyl ketone reported their results as observed BOD in percentage of theoretical oxygen demand plotted with days of incubation. Randall (1966) reports the effect of acclimated seed on the pesticides, Malathion and Parathion, in terms of net oxygen utilization and time in hours. In an article entitled "The BOD of Textile Chemicals, Updated List -1966", the data presented on nearly 400 chemicals and commercial chemical products are given as percent of 5-day BOD (Anon., 1966). In another paper (Anon., 1958), data for mercuric chloride, sulfuric acid, formaldehyde, and phenol are presented as the median toxic concentration in mg/l, i.e., the concentration at which 50 percent inhibition of oxygen utilization occurred; Zintgraff, et al (1968) reported BOD data using acclimated and nonacclimated seed for potassium cyanide in molar concentrations plotted against oxygen uptake in ppm or with time in hours. Rudolfs, et al (1950) reviewed the literature in 1950 on toxic materials affecting sewage treatment processes, streams, and BOD determinations and made general statements concerning this subject but with scant tabular material.

Since such a variety of methods for presenting data are found in the BOD literature, no attempt has been made to summarize BOD results in this report. The reader is referred to the various articles cited for information pertinent to his own interests, and to the summaries of chemical data shown in Appendixes A and B.

Herman (1959) proposed a toxicity index based on BOD data. Depending on the BOD curves obtained (percent available oxygen utilized plotted against concentration in mg/l), a series of "toxigrams" (Types 1 through 5) were devised, which were:

Toxigram Type 1 -simple poisons (the curve drops at toxic concentrations)

Toxigram Type 2 - no effect (the "curve" is flat)

Toxigram Type 3 – immediate dissolved oxygen demand (IDOD) by reducing substances (the curve rises to 100 percent oxygen utilization at higher concentrations)

Toxigram Type 4 – oxygen demand at low concentrations, inhibition of oxygen utilization at relatively high concentrations (the curve rises at low concentrations and drops at toxic levels)

Toxigram Type 5 — same as Type 4 except that at still higher concentrations oxygen utilization rose to 100 percent again. The author noted that the rise in oxygen utilization was due probably to simple chemical oxygen demand.

By designating the median toxic concentration (TC50) and indicating the appropriate toxigram type, a convenient index for characterizing that particular chemical was obtained.

Despite its disadvantages, i.e., slowness, lack of correlation between the lab and the receiving stream, empirical application, and lack of reproducibility between laboratories, the BOD bioassay or some variation of it can be a useful tool in pollution control. An effort should be made by those who depend on BOD determinations to arrive at a common method for reporting results and possibly to develop a toxigram index similar to that proposed by Herman (1959).

Data for 33 chemicals from Herman's study are summarized in Table 4. This index approach has not been widely adopted, but probably should be in view of the confusing data presentations revealed in the present critique. Herman pointed out that toxic concentrations other than the median, e.g.,  $TC_{10}$ ,  $TC_{25}$ ,  $TC_{75}$ , etc., can be chosen to suit individual industrial needs for release of chemicals.

Correlations of BOD with other data have also been attempted with varying success as follows:

Chemical data on phenols, heavy metals, etc. (Lloyd and Jordan, 1964) Respirometric methods (Vernimmen, et al, 1967; Montgomery, 1967) Aquatic biota (Burlington, 1962) Coliforms (Burlington, 1962)

Hynes (1959) has diagramatically depicted the effect of an organic effluent on a river by plotting the BOD rate from an effluent outfall downstream and its relationship to dissolved oxygen, salt, suspended solids, concentration of nitrogen (NH4 and NO3) and phosphate (PO4), and populations of algae, bacteria, sewage fungi, *Cladophora*, Protozoa, Tubificidae, *Chironomus*, *Asellus*, and clean water fauna. These diagrams are quite general and Hynes pointed out that the detailed relationship of the various parameters plotted varies with the type of effluent.

### Short-Term Oxygen Demand

The short-term oxygen demand (STOD) bioassay is a variation of BOD which requires time in the order of minutes or a few hours to conduct rather than 5 days or longer. The STOD requires a relatively sophisticated respiration cell with an oxygen electrode, continuous recorder, and ancillary equipment compared to that required for BOD determinations. However, endogenous growth rate, effect of substrate addition, and oxygen demand to the point of substrate exhaustion can be determined within 40 minutes for some types of compounds. When oxygen is fully utilized, the system may be aerated and further oxygen utilization followed. Vernimmen, et al (1967) reviewed previous research on this subject and described the equipment, procedure, and some results on such chemicals as sodium acetate, formaldehyde, methanol, isopropanol, isobutanol and phenol. In this study various types of acclimated and domestic sewage seed were used. Vernimmen and co-workers suggest establishing a suitable correlation factor between STOD and BOD for a given waste and predicting BOD by means of a STOD/BOD ratio in the same manner as COD is used in predicting BOD. Although appealing because of immediate results, the STOD bioassay has not received wide acceptance.

Ammonium thiosyante         NH4SCN         5000         2           Boric acid         H3B04         1000         2           Cadmium sulfate         Cd <sup>++</sup> 142         1           Chromic sulfate         Ct <sup>+3</sup> 117         1           Cobatt chloride         CoC2         64         1           Copper sulfate         CuS04         21         1           Mercuric chloride         HgC2         0.61         1           Potassium cyanide         KCN         15         1           Suffuric acid         H2S04         58         1           Inorganic Reducing Agents         under Certain Conditions         -         3           Ferrous sulfate         FeS04         -         3         3           Sodium metarsenite         NaA902         -         3         3           Sodium oxalate         Na2C204         -         3         3           Inorganic Oxidizing Agents         under Acid Conditions         -         3           Under Acid Conditions         -         3         3         3           Sodium narsenate         Na3A904         100         2         3           Organic Acids and Derivatives	Substance Tested	Reported As	TC50, mg/l*	Toxigram Type
Boric acid         HgBO4         1000         2           Cadmium sulfate         Cd <sup>++</sup> 142         1           Chromic sulfate         Cd <sup>++</sup> 142         1           Chromic sulfate         CoSQ         64         1           Coper sulfate         CuSQ         21         1           Mercuric chloride         HgC12         0.61         1           Potassium cyanide         KCN         15         1           Sulfurie acid         H2SQ         -         3           Inorganic Reducing Agents         -         3         3           under Certain Conditions         -         -         3           Sodium metaarsenite         NaA02         -         -         3           Sodium oxalate         Na2C2O4         -         3         3           Sodium asenate         Na2C2O4         -         -         3           Inorganic Axidizing Agents         -         -         3         3           under Acid Conditions         -         -         3         -           Potassium dichromate         Cr <sup>46</sup> 17         1         Sodium asenate         Na3AsO4         100         2      <	Simple Inorganic Poisons			
Boric acid         H3B04         1000         2           Cadmium sulfate         Cd <sup>++</sup> 142         1           Chromic sulfate         Cx <sup>+3</sup> 117         1           Cobatt chloride         CoCl2         64         1           Coper sulfate         CuS04         21         1           Mercuric chloride         HgCl2         0.61         1           Potassium cyanide         KCN         15         1           Sulfuric acid         H2S04         58         1           Inorganic Reducing Agents	Ammonium thiocyanate	NH4SCN	5000	2
Cadmium sulfate       Cd <sup>++</sup> 142       1         Chromic sulfate       Cit <sup>+3</sup> 117       1         Cobat chloride       CoCl2       64       1         Copper sulfate       CuSO4       21       1         Mercuric chloride       HgCl2       0.61       1         Potassium cyanide       KCN       15       1         Inorganic Reducing Agents       under Certain Conditions       1       1         Inorganic Reducing Agents       under Certain Conditions       1       3         Ferrous sulfate       FeSO4       -       3       3         Sodium metaarsenite       NaAsO2       -       3       3         Sodium oxalate       Na2C2O4       -       3       3         Under Acid Conditions        2       2       2       3         Inorganic Oxidizing Agents       under Acids and Derivatives       -       3       3         Vitrobenzene       CdH5NH-COCH3       -       3       3         Sodium obenzoel sulfinide       CdH5NH-COCH3       -       3       3         Formic acid       HCO2H       550       4       4         Sodium obenzoel sulfinide       CdH5ND2 <td>-</td> <td></td> <td>1000</td> <td>2</td>	-		1000	2
Cobalt chloride         CoCl2         64         1           Copper sulfate         CuSO4         21         1           Mercuric chloride         HgC2         0.61         1           Potassium cyanide         KCN         15         1           Sulfuric acid         H2S04         58         1           Inorganic Reducing Agents         under Certain Conditions         -         3           Vinder Certain Conditions         Ferrous sulfate         FeSO4         -         3           Sodium metaarsenite         NaAS02         -         3         3           Sodium metaarsenite         NaN02         -         3         3           Inorganic Oxidizing Agents         under Acid Conditions         -         3           Potassium dichromate         Cr <sup>46</sup> 17         1         1           Sodium assenate         Na3sO4         100         2         0           Organic Acids and Derivatives         -         3         3         4           Acetanilide         C6H5NP-COCH3         -         3         4           Sodium bezoate         C6H5NO2         630         4         4           Sodium bezoate         C6H5CO2Na*H2O	Cadmium sulfate		142	1
Cobait chlorideCoCl2641Copper sulfateCuSO4211Mercuric chlorideHgC20.611Potassium cyanideKCN151Sulfuric acidH2SO4581Inorganic Reducing Agents11under Certain ConditionsFerous sulfateFeSO4-Ferrous sulfateFeSO4-3Oxalic acidH2C2O4431Sodium metaarseniteNaNO2-3Sodium moxalateNaNO2-3Inorganic Oxidizing Agents-3under Acid ConditionsCr <sup>+6</sup> 171Potassium dichromateCr <sup>+6</sup> 171Sodium senateNa3ASO41002Organic Acids and Derivatives-3AcetanilideC6H5NH-COCH3-3Formic acidH'CO2H5504NitrobenzeneC6H5OQ2H -33Sodium benzoateC6H5CO2Na*H2O-3Sodium benzoateC6H5CO2Na*H2O-3Sodium benzoateCH3-CH2-3AcetaldehydeCH3-CH3-3Acetonin10002(soluble saccharin)-3and Derivatives-3AcetoneCH3-CH3-3AcetoneCH3-CH3-3AcetoneCH3-CH3-3AcetoneCH3-CH3-3AcetoneCH3-CH4 </td <td>Chromic sulfate</td> <td>Cr+3</td> <td>117</td> <td>1</td>	Chromic sulfate	Cr+3	117	1
Copper sulfate         CuSO4         21         1           Mercuric chloride         HgCl2         0.61         1           Potassium cyanide         KCN         15         1           Sulfuric acid         H2SO4         58         1           Inorganic Reducing Agents         -         3         3           Under Certain Conditions         -         3         3           Ferrous sulfate         FeSO4         -         3           Sodium netaarsenite         NaNO2         -         3           Sodium oxalate         NaNO2         -         3           Inder Acid Conditions         -         3         3           Under Acid Conditions         -         3         3           Potassium dichromate         Cr <sup>+6</sup> 17         1           Sodium arsenate         Na3AsO4         100         2           Organic Acids and Derivatives         -         -         3           Acetanilide         C6H5NH-COCH3         -         3           Acetanilide         C6H5NO2         630         4           Nitrobenzene         C6H5NO2         630         4           Sodium bezoate         C6H2CN2H4DO		CoCl <sub>2</sub>	64	1
Mercuric chloride         HgCl         0.61         1           Potassium cyanide         KCN         15         1           Sulfuric acid         HyS04         58         1           Inorganic Reducing Agents			21	1
Potassium cyanide $KCN$ 151Sulfuric acidH2SO4581Inorganic Reducing Agents under Certain Conditions-3Ferrous sulfateFeSO4-3Sodium metarseniteNaASO2-3Sodium metarseniteNaASO2-3Sodium netarseniteNaASO2-3Sodium oxalateNa2C2O4-3Inorganic Oxidizing Agents under Acid Conditions-3Potassium dichromateCr <sup>+6</sup> 171Sodium arsenateNa3ASO41002Organic Acids and Derivatives-3AcetanilideC6H5NH-COCH3-3Formic acidH-CO2H5504NitrobenzeneC6H5NO26304Sodium benzoateC9HC6H4OH1104Sodium benzoateC6H5NO2-3Sodium benzoateC6H5NO2-3Sodium benzoateC6H5CO2Na <sup>+</sup> H2O-3Sodium benzoateC6H5CO2Na <sup>+</sup> H2O-3Sodium benzoateC6H3-CHO2305AcetaldehydeCH3-CHO2305AcetaldehydeCH3-CHO7404Hexamethylenetetramine(CH2)6N4-3MethanolCH3-GH4-OH9404HexamethylenetetramineCH3-GH4-OH9404ocresolCH3-GH4-OH9404mathydroxybenzeneC6H4(OH)2-3Ocresol <td></td> <td>•</td> <td>0.61</td> <td>1</td>		•	0.61	1
Sulfuric acid $H_2SO_4$ 58         1           Inorganic Reducing Agents under Certain Conditions         Inorganic Reducing Agents under Certain Conditions         Inorganic Acid         12000           Ferrous sulfate         FeSO_4         -         3         3           Sodium netaarsenite         NaAO2         -         3         3           Sodium netaarsenite         NaAO2         -         3         3           Sodium oxalate         Na2C2O4         -         3         3           Inorganic Oxidizing Agents         under Acid Conditions         -         3           Potassium dichromate         Cr <sup>+6</sup> 17         1         5           Sodium arsenate         Na3ASO4         100         2         2           Organic Acids and Derivatives         -         3         3         3           Acetanilide         C6H5NP-COCH3         -         3         3         5           Sodium obenzoyl sulfimide         C7H4O3NSNa-H2O         1000         2         2         6         3           Sodium obenzoyl sulfimide         C7H4O3NSNa-H2O         0         3         3         3           Sodium obenzoyl sulfimide         CH3-CHO         230			15	1
under Certain Conditions           Ferrous sulfate         FeSO4         -         3           Oxalic acid         H2C2O4         43         1           Sodium metaarsenite         NaASO2         -         3           Sodium nitrite         NaNO2         -         3           Sodium oxalate         Na2C2O4         -         3           Inorganic Oxidizing Agents         -         3           Under Acid Conditions         -         3           Potassium dichromate         Cr <sup>+6</sup> 17         1           Sodium arsenate         Na3ASO4         100         2           Organic Acids and Derivatives         -         -         3           Acetanlide         C6H5NH-COCH3         -         -         3           Formic acid         H-CO2H         550         4           Nitrobenzene         C6H5NO2         630         4           Solium obenzoate         C6H5NO2         -         3           Sodium obenzoate         C6H5C02Na <sup>+</sup> H2O         -         3           Sodium benzoate         CH4C02H         1000         2           (soluble saccharin)         Tannic acid         (HO)3C6H2·CO         -         3 </td <td></td> <td></td> <td>58</td> <td>1</td>			58	1
Ferrous sulfate         FesO4         -         3           Oxalic acid         H2C2O4         43         1           Sodium metaarsenite         NaAsO2         -         3           Sodium nitrite         NaNO2         -         3           Sodium oxalate         NayC2O4         -         3           Inorganic Oxidizing Agents         -         3           Under Acid Conditions         -         3           Potassium dichromate         Cr <sup>+6</sup> 17         1           Sodium arsenate         Na3AsO4         100         2           Organic Acids and Derivatives         -         -         3           Acetanlide         C6H5NH-COCH3         -         -         3           Formic acid         HCO2H         550         4           Nitrobenzene         C6H5NO2         630         4           Sodium benzoate         C6H5-CO2Na <sup>+</sup> H2O         -         3           Sodium benzoate         C6H5-CO2Na <sup>+</sup> H2O         -         3           Sodium benzoate         C6H3-CH2         0000         2           (soluble saccharin)         Tannic acid         (HO)3C6H2-CO         -         3           Acetone	Inorganic Reducing Agents under Certain Conditions			
Oxalic acid $H_2C_2O_4$ 43       1         Sodium metarsenite       NaAsO2       -       3         Sodium nitrite       NaNO2       -       3         Sodium oxalate       Na2C2O4       -       3         Inorganic Oxidizing Agents       -       3         under Acid Conditions       -       3         Potassium dichromate       Cr <sup>46</sup> 17       1         Sodium arsenate       Na3AsO4       100       2         Organic Acids and Derivatives       -       3         Acetanlilde       C6H5NH-COCH3       -       3         Formic acid       HCO2H       550       4         Nitrobenzene       C6H5NO2       630       4         Sodium benzoate       C6H5CO2Na <sup>+</sup> H2O       -       3         Sodium benzoate       C6H5CO2Na <sup>+</sup> H2O       -       3         Sodium benzoate       CH3 <sup>+</sup> CO2Na <sup>+</sup> H2O       -       3         Sodium benzoate       CH3 <sup>+</sup> CHO       230       2         (soluble saccharin)       -       3       3         Tannic acid       (HO)3C6H2 <sup>-</sup> CO       -       3         Acetaldehyde       CH3 <sup>+</sup> CHO       230       5		FasO	_	3
Sodium metaarsenite         NaASO2         -         3           Sodium nitrite         NaNO2         -         3           Sodium oxalate         NayC2O4         -         3           Inorganic Oxidizing Agents         -         3           Under Acid Conditions         -         3           Potassium dichromate         Cr <sup>46</sup> 17         1           Sodium arsenate         NajAsO4         100         2           Organic Acids and Derivatives         -         3           Acetanilide         C6H5NH-COCH3         -         3           Formic acid         H-CO2H         550         4           Nitrobenzene         C6H5NO2         630         4           Sodium obenzoate         C6H5NO2         -         3           Sodium obenzoate         C6H5C02Na:H2O         -         3           Sodium obenzoate         C6H3-CO2Na:H2O         -         3           Acetanili         Tannic acid         (HO)3C6H2-CO         -         3           Acetonein)         Tannic acid         CH3-CO-CH3         -         3           Acetone         CH3-CO-CH3         -         3         3           Acetone <t< td=""><td></td><td>•</td><td>43</td><td></td></t<>		•	43	
Sodium nitriteNaNO2-3Sodium oxalateNa2C2O4-3Inorganic Oxidizing Agents under Acid Conditions-3Potassium dichromateCr <sup>46</sup> 171Sodium arsenateNa3AsO41002Organic Acids and Derivatives-3AcetanilideC6H5NH-COCH3-3Formic acidHCO2H5504NitrobenzeneC6H5NO26304Sodium benzoateC6H5CO2Na H2O-3Sodium benzoateC6H5-CO2Na H2O-3Sodium o-benzoyl sulfimideC7H4O3NSNa H2O10002(soluble saccharin)-3Tannic acid(HO)3C6H2-CO-3Alcetoles, Aldehydes, Ketones, and Derivatives-3AcetadehydeCH3-CH3-3FormaldehydeCH3-CH3-3Phenols and Cresols-3OresolCH3-C6H4-OH-3PhenolC6H4(OH)2-3Acetalde HydrocarboneC6H4(OH)2-3OresolCH3-C6H4-OH9404MethanolCH3-C6H4-OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons-3				
Sodium oxalateNa $2C_{2}O_{4}$ -3Inorganic Oxidizing Agents under Acid ConditionsInorganic Acid Conditions171Potassium dichromate $Cr^{+6}$ 171Sodium arsenateNa $3AsO_4$ 1002Organic Acids and Derivatives-3Acetanilide $C_{6}H_5NH+COCH_3$ -3Formic acidHCO2H5504Nitrobenzene $C_{6}H_5NO_2$ 6304Sodium obenzoate $C_{6}H_5CO_2Na+H_2O$ -3Sodium obenzoate $C_{6}H_5CO_2Na+H_2O$ -3Sodium obenzoate $C_{7}H4O_3NSNa+H_2O$ 10002(soluble saccharin)-33Tannic acid(HO) $_3C_6H_2\cdotCO$ -3Alcohols, Aldehydes, Ketones, and Derivatives-3AcetaldehydeCH_3·CHO2305AcetaldehydeH-CH:O7404Hexamethylenetetramine(CH_2) $6N_4$ -3Phenols and Cresols-33O-cresolCH_3·ChH9404mdhydroxybenzeneC6H_4(OH)-32,4-dinitrophenol(NO2) $_2C_6H_3OH$ 1001PhenolC6H_5OH16004Chlorinated Hydrocarbons-3				3
under Acid ConditionsPotassium dichromate Sodium arsenateCr <sup>+6</sup> 171Potassium dichromate Sodium arsenateNa3AsO41002Organic Acids and Derivatives2AcetanilideC <sub>6</sub> H <sub>5</sub> NH-COCH <sub>3</sub> -3AcetanilideC <sub>6</sub> H <sub>5</sub> NO26304NitrobenzeneC <sub>6</sub> H <sub>5</sub> NO26304Salicylic acidCO2H-C <sub>6</sub> H <sub>4</sub> -OH1104Sodium benzoateC <sub>6</sub> H <sub>5</sub> -CO2Na·H <sub>2</sub> O-3Sodium o-benzoyl sulfimideC7H4O <sub>3</sub> NSNa·H <sub>2</sub> O10002(soluble saccharin)Tannic acid(HO) <sub>3</sub> C <sub>6</sub> H <sub>2</sub> -CO-3Alcohols, Aldehydes, Ketones, and Derivatives-33AcetaldehydeCH <sub>3</sub> ·CHO2305AcetaldehydeH-CH:O7404Hexamethylenetetramine(CH <sub>2</sub> ) <sub>6</sub> N <sub>4</sub> -3Phenols and Cresols-33o-cresolCH <sub>3</sub> ·C <sub>6</sub> H <sub>4</sub> ·OH9404n-dihydroxybenzeneC <sub>6</sub> H <sub>4</sub> (OH) <sub>2</sub> -32,4-dinitrophenol(NO <sub>2</sub> ) <sub>2</sub> C <sub>6</sub> H <sub>3</sub> OH1001PhenolC <sub>6</sub> H <sub>5</sub> OH16004	_		_	3
Sodium arsenateNa3AsO41002Organic Acids and DerivativesAcetanilide $C_{6}H_5NH$ -COCH3-3Formic acidH·CO2H5504Nitrobenzene $C_6H_5NO_2$ 6304Sodium benzoate $C_6H_5\cdotO_2Na\cdotH_2O$ -3Sodium o-benzoyl sulfimide $C_7H_4O_3NSNa\cdotH_2O$ 10002(soluble saccharin)773Tannic acid(HO)3C_6H_2·CO-3Alcohols, Aldehydes, Ketones, and Derivatives-3AcetaldehydeCH3·CH02305AcetoneCH3·CO·CH3-3FormaldehydeH·CH·O7404Hexamethylenetetramine(CH2)/6N4-3Phenols and CresolsCH3·C6H4·OH9404o-cresolCH3·C6H4·OH9404m-dihydroxybenzeneC_6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons	Inorganic Oxidizing Agents under Acid Conditions			
Sodium arsenateNa3AsO41002Organic Acids and DerivativesAcetanilide $C_6H_5NH$ -COCH3-3Formic acidH·CO2H5504Nitrobenzene $C_6H_5NO_2$ 6304Sodium benzoate $C_6H_5CO_2NaH_2O$ -3Sodium o-benzoyl sulfimide $C7H_4O_3NSNa+H_2O$ 10002(soluble saccharin)-3Tannic acid(HO)3C6H_2·CO-3Alcohols, Aldehydes, Ketones, and Derivatives-3AcetaldehydeCH3·CHO2305AcetaldehydeHCH:O7404Hexamethylenetetramine(CH2)6N4-3Phenols and Cresols-33Phenols and CresolsCH3·C6H4·OH9404o-cresolCH3·C6H4·OH9404m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons	Potassium dichromate	Cr+6	17	1
AcetanilideC6H5NH-COCH3-3Formic acidH-CO2H5504NitrobenzeneC6H5NO26304Salicylic acidCO2H-C6H4-OH1104Sodium benzoateC6H5·CO2Na·H2O-3Sodium benzoateC6H5·CO2Na·H2O-3Sodium benzoateC6H5·CO2Na·H2O-3Sodium benzoateC6H5·CO2Na·H2O-3Sodium benzoateC6H5·CO2Na·H2O-3Moluble saccharin)Tannic acid(HO)3C6H2·CO-3Tannic acid(HO)3C6H2·CO-3Alcohols, Aldehydes, Ketones, and Derivatives-3AcetaldehydeCH3·CHO2305AcetaldehydeH·CH:O7404Hexamethylenetetramine(CH2)6N4-3Phenols and Cresols-33o-cresolCH3·C6H4·OH9404m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons-3		Na3AsO4		
Formic acidHCO2H5504Nitrobenzene $C_6H_5NO_2$ 6304Salicylic acid $CO_2H \cdot C_6H_4 \cdot OH$ 1104Sodium benzoate $C_6H_5 \cdot CO_2Na \cdot H_2O$ –3Sodium o-benzoyl sulfimide $C_7H_4O_3NSNa \cdot H_2O$ 10002(soluble saccharin)Tannic acid(HO)_3C_6H_2 \cdot CO–3Tannic acid(HO)_3C_6H_2 \cdot CO–3Alcohols, Aldehydes, Ketones, and Derivatives–3AcetaldehydeCH_3 \cdot CHO2305AcetoneCH_3 \cdot CO \cdot CH_3–3FormaldehydeH \cdot CH: O7404Hexamethylenetetramine(CH2)_6N4–3Phenols and CresolsCH3 \cdot C_6H4 \cdot OH9404o-cresolCH3 \cdot C_6H4 \cdot OH9404m-dihydroxybenzeneC_6H4(OH)_2–32,4-dinitrophenol(NO2)_2C_6H3OH1001PhenolC_6H5 OH16004Chlorinated Hydrocarbons4	Organic Acids and Derivatives			
Formic acidHCO2H5504Nitrobenzene $C_6H_5NO_2$ 6304Salicylic acid $CO_2H \cdot C_6H_4 \cdot OH$ 1104Sodium benzoate $C_6H_5 \cdot CO_2Na \cdot H_2O$ –3Sodium o-benzoyl sulfimide $C_7H_4O_3NSNa \cdot H_2O$ 10002(soluble saccharin)Tannic acid(HO)_3C_6H_2 \cdot CO–3Tannic acid(HO)_3C_6H_2 \cdot CO–3Alcohols, Aldehydes, Ketones, and Derivatives–3AcetaldehydeCH_3 \cdot CHO2305AcetoneCH_3 \cdot CO \cdot CH_3–3FormaldehydeH \cdot CH: O7404Hexamethylenetetramine(CH2)_6N4–3Phenols and CresolsCH3 \cdot C_6H4 \cdot OH9404o-cresolCH3 \cdot C_6H4 \cdot OH9404m-dihydroxybenzeneC_6H4(OH)_2–32,4-dinitrophenol(NO2)_2C_6H3OH1001PhenolC_6H5 OH16004Chlorinated Hydrocarbons4	Acetanilide	C6H5NH·COCH3	-	3
Nitrobenzene $C_6H_5NO_2$ $630$ 4Salicylic acid $CO_2H \cdot C_6H_4 \cdot OH$ 1104Sodium benzoate $C_6H_5 \cdot CO_2Na \cdot H_2O$ -3Sodium o-benzoyl sulfimide $C7H4O_3NSNa \cdot H_2O$ 10002(soluble saccharin)Tannic acid(HO)_3C_6H_2 \cdot CO-3Tannic acid(HO)_3C_6H_2 \cdot CO-3Alcohols, Aldehydes, Ketones, and Derivatives-3AcetaldehydeCH_3 \cdot CHO2305AcetoneCH_3 \cdot CO \cdot CH_3-3FormaldehydeH \cdot CH: O7404Hexamethylenetetramine(CH_2)_6N4-3Phenols and Cresols-33o-cresolCH_3 \cdot C_6H_4 \cdot OH9404m-dihydroxybenzeneC_6H4(OH)_2-32,4-dinitrophenol(NO_2)_2C_6H_3OH1001PhenolC_6H5OH16004Chlorinated Hydrocarbons	Formic acid		550	4
Salicylic acid $CO_2H-C_6H_4-OH$ 1104Sodium benzoate $C_{6H5} \cdot CO_2Na \cdot H_2O$ -3Sodium o-benzoyl sulfimide $C7H4O_3NSNa \cdot H_2O$ 10002(soluble saccharin)Tannic acid(HO)_3C_6H_2 \cdot CO-3Alcohols, Aldehydes, Ketones, and Derivatives-33AcetaldehydeCH_3 \cdot CHO2305AcetaldehydeHoho CH_3 \cdot CO \cdot CH_3-3FormaldehydeH·CH:O7404Hexamethylenetetramine(CH_2)_6N4-3MethanolCH_3 OH-3Phenols and CresolsCH_3 \cdot C_6H_4 \cdot OH9404o-cresolCH_3 \cdot C_6H_4 · OH9404m-dihydroxybenzeneC_6H_4(OH)_2-32,4-dinitrophenol(NO_2)_2 \cdot C_6H_3 OH1001PhenolC_6H5 OH16004Chlorinated Hydrocarbons		-	630	
Sodium benzoate $C_6H_5 \cdot CO_2Na \cdot H_2O$ -3Sodium o-benzoyl sulfimide $C_7H_4O_3NSNa \cdot H_2O$ 10002(soluble saccharin)Tannic acid(HO)_3C_6H_2 \cdot CO-3Alcohols, Aldehydes, Ketones, and Derivatives-33AcetaldehydeCH_3 \cdot CHO2305AcetaldehydeCH_3 \cdot CO \cdot CH_3-3FormaldehydeH \cdot CH: O7404Hexamethylenetetramine(CH2)6N4-3MethanolCH3 \cdot Cfl + OH9404o-cresolCH3 \cdot C_6H4 \cdot OH9404m-dihydroxybenzeneC_6H4(OH)_2-32,4-dinitrophenol(NO2)2C_6H3OH1001PhenolC_6H5OH16004Chlorinated Hydrocarbons4-				
Sodium o-benzoyl sulfimide (soluble saccharin) Tannic acid $C7H4O_3NSNa \cdot H_2O$ 10002Match 103 C6H2 \cdot CO-3Alcohols, Aldehydes, Ketones, and Derivatives-3AcetaldehydeCH3 · CHO2305AcetoneCH3 · CO· CH3-3FormaldehydeH· CH: O7404Hexamethylenetetramine(CH2)6N4-3MethanolCH3 · C6H4 · OH9404o-cresolCH3 · C6H4 · OH9404m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons			_	
(soluble saccharin) Tannic acid (HO) <sub>3</sub> C <sub>6</sub> H <sub>2</sub> ·CO – 3 Alcohols, Aldehydes, Ketones, and Derivatives Acetaldehyde CH <sub>3</sub> ·CHO 230 5 Acetone CH <sub>3</sub> ·CO·CH <sub>3</sub> – 3 Formaldehyde H·CH:O 740 4 Hexamethylenetetramine (CH <sub>2</sub> ) <sub>6</sub> N <sub>4</sub> – 3 Methanol CH <sub>3</sub> OH – 3 Phenols and Cresols o-cresol CH <sub>3</sub> ·C <sub>6</sub> H <sub>4</sub> ·OH 940 4 m-dihydroxybenzene C <sub>6</sub> H <sub>4</sub> (OH) <sub>2</sub> – 3 2,4-dinitrophenol (NO <sub>2</sub> ) <sub>2</sub> C <sub>6</sub> H <sub>3</sub> OH 100 1 Phenol C <sub>6</sub> H <sub>5</sub> OH 1600 4 Chlorinated Hydrocarbons			1000	
Tannic acid $(HO)_3C_6H_2 \cdot CO$ -3Alcohols, Aldehydes, Ketones, and Derivatives $and Derivatives$ 5AcetaldehydeCH3 \cdot CHO2305AcetaldehydeCH3 · CO·CH3-3FormaldehydeH·CH:O7404Hexamethylenetetramine(CH2)_6N4-3MethanolCH3 · C6H4 · OH-3Phenols and Cresolso-cresolCH3 · C6H4 · OH9404m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004		07-4-5		-
and DerivativesAcetaldehydeCH3·CHO2305AcetoneCH3·CO·CH3-3FormaldehydeH·CH:O7404Hexamethylenetetramine(CH2)6N4-3MethanolCH3OH-3Phenols and Cresolso-cresolCH3·C6H4·OH9404m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004		(HO) <sub>3</sub> C <sub>6</sub> H <sub>2</sub> ·CO	-	3
AcetoneCH3·CO·CH3-3FormaldehydeH·CH:O7404Hexamethylenetetramine(CH2)6N4-3MethanolCH3OH-3Phenols and Cresolso-cresolCH3·C6H4·OH940o-cresolCH3·C6H4·OH940m-dihydroxybenzeneC6H4(OH)2-2,4-dinitrophenol(NO2)2C6H3OH100PhenolC6H5OH1600	Alcohols, Aldehydes, Ketones, and Derivatives			
AcetoneCH3·CO·CH3-3FormaldehydeH·CH:O7404Hexamethylenetetramine(CH2)6N4-3MethanolCH3OH-3Phenols and Cresolso-cresolCH3·C6H4·OH940o-cresolCH3·C6H4·OH940m-dihydroxybenzeneC6H4(OH)2-2,4-dinitrophenol(NO2)2C6H3OH100PhenolC6H5OH1600	Acetaldehyde	СНасСНО	230	5
FormaldehydeH·CH:O7404Hexamethylenetetramine $(CH_2)_6N_4$ -3MethanolCH_3OH-3Phenols and CresolsCH_3·C_6H_4·OH9404o-cresolCH_3·C_6H_4·OH9404m-dihydroxybenzeneC_6H_4(OH)_2-32,4-dinitrophenol(NO2)_2C_6H_3OH1001PhenolC_6H_5OH16004	-		_	
Hexamethylenetetramine Methanol $(CH_2)_6N_4$ $CH_3OH$ -3Phenols and Cresols-3o-cresol m-dihydroxybenzeneCH_3·C_6H_4·OH C_6H_4(OH)_294042,4-dinitrophenol Phenol(NO2)_2C_6H_3OH C_6H_5OH1001Chlorinated HydrocarbonsChlorinated Hydrocarbons4			740	4
MethanolCH3OH-3Phenols and CresolsCH3·C6H4·OH9404o-cresolCH3·C6H4·OH9404m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated HydrocarbonsChlorinated HydrocarbonsCH3·C6H4/OH				
o-cresol $CH_3 \cdot C_6H_4 \cdot OH$ 9404m-dihydroxybenzene $C_6H_4(OH)_2$ -32,4-dinitrophenol $(NO_2)_2C_6H_3OH$ 1001Phenol $C_6H_5OH$ 16004			—	
m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons33	Phenols and Cresols			
m-dihydroxybenzeneC6H4(OH)2-32,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons33	o-cresol	СНассенаюн	940	4
2,4-dinitrophenol(NO2)2C6H3OH1001PhenolC6H5OH16004Chlorinated Hydrocarbons			_	
Phenol     C <sub>6</sub> H <sub>5</sub> OH     1600     4       Chlorinated Hydrocarbons			100	
Chlorinated Hydrocarbons				
	Chloroform	HCCl <sub>3</sub>		3

 $TC_{50}$  = Concentration at which oxygen utilization is reduced 50 percent.

### **Related Microbiological Methods**

Montgomery (1967) and Ludzack and Ettinger (1963) thoroughly reviewed respirometric methods for the determination of biochemical oxygen demand, including the STOD procedure, Warburg respirometry, Barcroft differential manometry, Wilson six-unit recording respirometry, electrolytic respirometry, the Sierp apparatus, the Nordell odeometer, the oxyutilometer, and Sapromat A6 respirometry. Malaney, et al (1959) presented data on the toxic effects of metallic ions on sewage microorganisms using the Warburg procedure.

Biodegradability of organic chemicals in the aquatic environment is another important factor related to biochemical oxygen demand. This is of increasing concern because of the accumulation of chemicals, especially pesticides and detergents, in the beds of rivers, lakes, and estuaries. The behavior of organic chemicals in the aquatic environment was reported in a recent study by Buzzell, et al (1968). At sublethal concentrations, the BOD, COD, total organic carbon (TOC), and toxicity as determined by microbial and fish bioassay were all determined for a selected group of 20 compounds representing a variety of types of chemicals. Bacterial enumeration was used to indicate bacterial growth in biodegradation units. Theoretical oxygen demand (TOD) for each compound was compared with 5-day and 20-day BOD results. The comparison showed that seldom was TOD reached in the BOD determinations. Graphs showing all of the data obtained were plotted. Each compound had its own characteristic set of curves for BOD, COD, TOD, etc. A sound basis resulted from this study to further evaluate BOD and other measures of chemical effect on aquatic organisms. This approach might well be used in the study of chemical toxicity in the aquatic environment.

Earlier, Ludjack and Ettinger (1963) reviewed methods of estimating the biodegradability and treatability of organic water pollutants and how various types of data from BOD, respirometry, etc., procedures can be applied in practice to various contact treatment units.

Several excellent papers (Beak, 1957; Dobbins, 1964; Gannon, 1966; Nejedly, 1967; and Smith, et al, 1962) discuss laboratory BOD determination in relation to receiving stream BOD and the multiple factors that are involved in calculating or estimating downstream dissolved oxygen drop. In particular, papers by Dobbins (1964), Gannon (1966), Goodman and Dobbins (1966), and Smith, et al (1962) would be particularly useful in developing mathematical modeling or simulation of stream problems associated with dissolved oxygen depletion.

Other microbiological techniques for study of various types of water pollution are described in standard texts too numerous to mention here. Bacteria and other microorganisms are usually studied as indicators of fecal pollution. Papers by Kabler (1957, 1961), Khan (1964), Bonde (1966), Morrison and Fair (1966), O'Connell and Thomas (1965), Cooke and Bartsch (1959), Burman (1966), and Bick (1963) describe studies in which enumerations were made of Escherichia coli, coliforms, fecal streptococci, salmonellae, Aeromonas, Pseudomonas, Clostridia, microfungi, actinomycetes, and algae. Bick (1963) extended this list of organisms to include protozoa and other aquatic invertebrates in reviewing Central European ecological approaches in studying water pollution. According to this approach, organisms characteristically occur in various "saprobic zones" which are used to describe the degree of pollution. The procedures involved in the papers cited above are concerned primarily with sewage pollution or taste and odor problems. Burman (1966) reviewed the various procedures, media, equipment, etc., in bacteriological examination of water and describes a technique in which C14-labelled compounds are incubated, the C14O2 evolved is absorbed on barium hydroxide, and counts of radioactivity are used to quantitate respiration. Since only 4 hours are required for completion, this technique might be a useful, more rapid variation of the standard BOD assay. A similar technique, using C14O2 in the study of photosynthetic activity of algae in the field, is used to determine trophic levels in various types of water (Butler, 1965).

### SECTION VII

### MARINE BIOASSAY

Any report or critique to be made of the methods used in bioassaying the effects of chemical pollutants on marine and estuarine forms can be presented concisely and to the point. That is, those bioassay techniques in which the flowing-seawater method is not used fall short of obtaining accurate tolerance limits, etc., for marine and estuarine species in regard to chemical pollutants. The flowing-seawater technique for both acute and chronic toxicity studies developed at the Bureau of Commercial Fisheries at Gulf Breeze, Florida, as described by Lowe (1964) comes closely to providing the necessary data regarding chemical toxicants to marine and estuarine forms.

In this technique, the chemical solution is contained in a stock solution bottle and is metered by means of a stopcock into a slanted mixing trough which contains running fresh seawater. The fresh seawater is kept in a holding tank at a constant level and is siphoned at a constant rate into the trough. From the trough, the toxicant-containing water flows by gravity over baffles into the chamber containing the test animals. A drain is situated at one end of the chamber to allow overflow and maintenance of a constant level of toxicant-containing water. The author states that this constant-flow system eliminated the need for aeration and that no attempt was made to control temperature and salinity. A record of the latter two values was kept however.

Data on marine studies are included in Appendixes A and B and may be identified by the names of the marine species listed in the second (Organism) column. Further identification is afforded by the Species Index (Appendix C).

Marine species most frequently used in bioassay include: :

Ostrea spp – oyster

Algae	Fish
Dunaliella euchlora Platymonas sp Crustacea Artemia salina – brine shrimp Callinectes sapidus – blue crab Carcinus spp – decapod crab Peneaus aztecus – brown shrimp P. duorarium – pink shrimp P. setiferus – white shrimp	Cyprinodon variegatus — sheepshead minnow Fundulus similis — longnose killifish Lagodon rhomboides — pinfish Leiostomus xanthurus — spot Mugil curema — white mullet M. cephalus — striped mullet Oncorynchus kisutch — coho salmon Petromyzon marinus — sea lamprey Salmo gairdneri — rainbow trout S. salar — Atlantic salmon
Molluscs Balanus spp – barnacle Crassostrea virginica – oyster Mercenia mercenia – hard clam Mya spp – soft shell clam	S. trutta – brown trout

References to marine studies are made throughout the various sections of this report. It is of some interest to note that somewhat less than 10 percent of all papers reviewed were concerned with studies on the effect of chemicals on the marine organisms.

### SECTION VIII

### FIELD ASSESSMENT

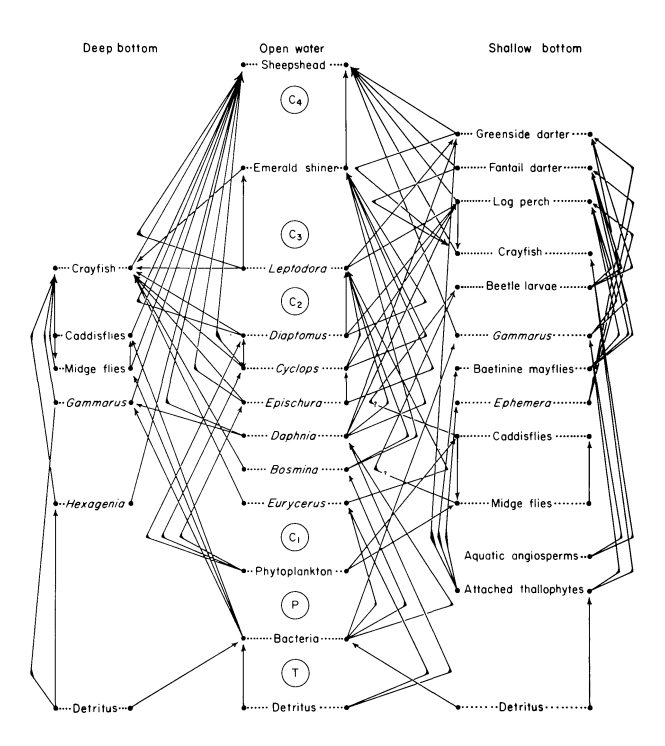
Many ecological parameters must be taken into consideration when field studies are conducted. Even minor variations in most environmental factors such as temperature, rainfall, pH, dissolved oxygen, and sunlight can significantly affect the toxicity of many chemical compounds. Full discussion of these factors is presented in texts by Hutchinson (1957, 1967), Welch (1952), Ruttner (1953), and Odum (1959). One consideration of major importance is the food web. The introduction of toxic substances at any point in the web may interfere with the reproduction and well-being of higher animal forms.

### Study of Residues in Aquatic Animals

The transfer of food energy from plants (the producers) through various animal organisms (the consumers) with repeated eating and being eaten is referred to as a food chain. The links in the chain seldom number more than five and usually many chains are interconnected with one another with the resulting pattern being called a food web. Figure 1 is a simplified diagram of a food web in western Lake Erie leading to the sheepshead. This diagram, modified from Daiber (1952) by Kendeigh (1961) shows the producers and consumers organized into nutritional or trophic levels. The lowest level (P) is composed of the producers that are able to use solar energy for the manufacture of food. At the second level (C1) are the primary consumers or grazing herbivores; at the third level (C2) the secondary consumers or small-size carnivores; and the fourth level (C3) the larger carnivores. It is possible that additional consumers may be present (C4). The consumer levels are not sharply defined because feeding behavior of some species may involve them in more than one level. Generally, the farther removed from the producers an organism is, the greater the likelihood it will feed on more than one level. Bacteria and fungi act as transformers (T) or decomposers and break down dead organic matter into nutrients that may be utilized by the producers (Ingols, 1959; Odum, 1959; Phillipson, 1966; and Welch, 1952).

Food webs are studied in a variety of ways including direct observation which is probably the least reliable. Stomach analysis of higher animal forms has been widely used for a great many years and has provided some useful information. When using this method, a major problem arises when plant juices and soft tissues must be considered because these are rapidly digested and practically impossible to identify. Precipitin tests have recently been used. An extract is made from a prey organism and this is injected into a rabbit which produces antibodies against this foreign protein. An extract is then made from a predator species and mixed with the rabbit antibodies. If this predator organism has been feeding on the prey organism, a white precipitate of antigen and antibody will be formed. In recent years, radioactive isotopes have also proven to be a most valuable tool in the study of the transfer of energy through trophic levels (Fujiya, 1965; Gakstatter and Weiss, 1967; and Miller, et al, 1966).

Meeks (1968) studied food chain organisms and how chemical contaminants can accumulate in the various trophic levels. A marsh adjacent to Lake Erie was treated with 3.9 millicuries of chlorine-36, ring-labeled DDT at a rate of 0.2 lb of technical DDT per acre. Radiolabeled DDT residues were traced until 15 months after the application. In his discussion of the work, Meeks stated that plankton and larger organisms rapidly removed the DDT from the water. Producer organisms contained their maximum residues between 1-3 days and most invertebrates contained their maximum residues several days later. These residues could have come directly from the



### FIGURE 1. FOOD WEB IN WESTERN LAKE ERIE LEADING TO THE SHEEPSHEAD FISH

Species are separated into their different trophic levels (as modified from Daiber, 1952, by Kendeigh, 1961).

water or could have been picked up through the food web. There are several factors that indicate that the food web is the most important contributor. Herbivorous snails, at the second trophic level, contained their maximum levels at the same time as most primary producers. Odonata naiads and backswimmers, both carnivorous invertebrates occupying the third trophic level, reached their peak accumulation at 1 week. The red leech was probably the invertebrate closest to a secondary carnivore, fourth trophic level, and it had the latest and highest DDT levels of any invertebrate. Most vertebrates attained their maximum DDT residues after the invertebrates had their highest levels.

The DDT applied in this project would equal 0.07 ppm in water if all of the DDT had been available at the same time. Meeks used this figure as a base level for determining magnitudes of accumulation and recorded a sample of *Cladophora* collected at 3 days which exceeded this level by a factor of 3125. For a tadpole at 4 hours and a northern water snake at 13 months accumulation was over 500 times this base level. Concentrations ranging from 200 to 500 times occurred in some duckweed and bladderwort samples during the first week as it did in samples of carp and tadpole tissues. Most plant and invertebrate species exceeded the 0.07 ppm by a factor of 50 during the first week and throughout the project, vertebrate tissue often concentrated DDT more than 50 times the base level.

Miller, et al (1966) noted that molluscs characteristically accumulate pesticidal compounds at levels far above those present in the surrounding water. In laboratory experiments, Butler (1966) showed that oysters exposed to one ppb of DDT in flowing seawater may store 25 ppm in its tissues within 10 days. Terriere, et al (1966) reported concentration factors from water to plant of 500, water to aquatic animals other than fish of 1,000 to 2,000, and for rainbow trout, 10,000 to 20,000. Odum, et al (1969) found that suspended particulate organic matter may be a reservoir of DDT and some particles may contain residues thousands of times greater than the concentration occurring in the water. Fiddler crabs and other organisms that utilize plant detritus for food concentrate the pesticide in their tissues.

Nicholson (1967) stated that any DDT which is not excreted or metabolized can accumulate in tissues to some degree. It may then be passed on to the next higher trophic level by way of the food chain. Pesticides have been detected in aquatic animal tissues far removed from where the chemicals were actually used. Sladen, et al (1966) cited examples of Adelie penguins and a crabeater seal whose tissues contained DDT residues. These species reportedly do not leave the Antarctic ice pack. The pathway to these animals is probably the marine crustaceans upon which they feed.

Cade, et al (1968) reported finding high levels of pesticides in the eggs and tissues of fish-eating peregrine falcons of the Yukon area of Alaska, and Enderson and Berge (1968) reported similar findings in peregrines in northern Canada.

Hunt and Bischoff (1960) believed that DDD residues in fish caused the deaths of grebes in Clear Lake, California. Investigations showed the following DDD concentrations in samples taken 13 months after application of the DDD: in plankton, 10 mg/kg; in fat from plankton-eating fish, 902 mg/kg; in fat from carnivorous fish, 2690 mg/kg; and in fat from fish-eating birds, 2134 mg/kg (Nicholson, 1967). It is believed that grebes are unable to tolerate as high a level of DDD as some species of fish.

Fay and Youatt (1967) concluded that various pesticide residues found in tissues of aquatic birds in Lake Michigan did not appear to be an important factor in bird die-offs in this lake. Studies by Keith (1966), however, suggest that unusual mortality of aquatic birds in California was due to pesticide poisoning. Pesticides have also been linked with the declining population of fish-eating ospreys in Connecticut (Ames, 1966).

Within a given species there may be strains or populations in existence which are resistant to, or have a greater tolerance for, a particular chemical and, therefore, will survive under conditions that would normally prove fatal for this species. Populations of yellow bullhead, golden shiner, green sunfish, and bluegill have been found that were resistant to Endrin (Ferguson and Bingham, 1966), while some mosquito fish (Ferguson and Bingham, 1966; Ferguson, et al, 1966; and Toohey, et al, 1965), and black bullhead (Ferguson, 1967) have been found resistant to DDT. The resistance of fish to these chemicals appears to be genetic, i.e., passed on from one generation to the next. This resistance, however, may be lost unless the fish are kept in continual contact with the chemical. While these populations are now geographically limited, the possibility exists that eventually they could become widespread. Ferguson (1967) concluded that although selection of a resistant fishery may permit fish exposed to toxic chemicals to survive, it may ultimately produce a biological product dangerous to consumers of all sort, including man himself.

In recent years, numerous investigations have been carried out on the accumulation of chemicals in both vertebrates and invertebrates. Emphasis has been placed primarily on pesticides (see Appendix B).

#### Field Methodology

Field assessment studies may be divided into two general types although a clear-cut distinction is not always possible. The first type consists of field observations made on the effects of chemicals on aquatic life with little prior manipulation or study of the environment by the investigator. In many cases, the exact concentration of the chemical is unknown and may not be fully identified but may be simply referred to as a pesticide, an eradicant, an industrial pollutant, an organic pollutant, etc. These studies are usually made when a body of water becomes polluted from a pesticide-spraying operation, effluents from an industrial site, or from the application of chemicals directly into the body of water.

The effects of these chemicals are often expressed as a reduction in numbers of a particular species or the total absence of a species or population. Dead organisms are sometimes identified and counted, as in fish kills, or estimations made of percent mortality of a given population. Effects may sometimes be expressed by noting the presence of particular organisms, usually considered to be undesirable, such as *Sphaerotilus, Chironomus,* and tubificids. Sometimes pre-pollution studies have been made or comparisons made between similar bodies of water. This type of approach has been widely used in assessing the effect of thermal pollution on aquatic life.

The second type of field assessment consists of actual toxicity studies of the effects of known chemical concentrations on particular organisms. The studies are sometimes made in conjunction with laboratory toxicity tests and implies some prior manipulation of the environment. Results are usually expressed in lethal concentrations of the chemical studied. Field assessments of this type are conducted in various sizes and types of water bodies. The smallest are simple pools or channels, such as man-made troughs or tanks. Ponds, man-made or natural, are widely used for this type of assessment. Lakes and reservoirs are also used but allow the minimum control in a lentic environment due to size. Streams are used, but less than lentic bodies of water. The following discussion deals with the methods used in these toxicity studies.

Chemicals are applied to bodies of water for the purpose of assessing their effects on aquatic organisms in several different ways. A uniform distribution is of primary concern and, therefore, the size and depth of the body of water will be a major factor in determining which method to use. Cloth bags containing chemicals may be submerged at various depths and the chemicals allowed to diffuse out into the water or the bags may be towed from a boat. A common method is to pour or drip chemicals from the stern of a power boat into the wake caused by the motor. Power sprayers are used from boats in smaller bodies of water or from the shore. In the largest bodies of water, airplanes or helicopters are used.

Gjullin, et al (1949) studied the effects of DDT on trout, blackfly, and caddisfly larvae from Alaskan streams using 6-ft-long galvanized metal troughs set up adjacent to a stream. Water from the stream was pumped into the troughs and DDT was administered by a 1-gallon aspirator bottle calibrated with a stopcock to deliver the desired concentration per minute. Darsie and Corriden (1959) used bushel-sized galvanized tubs placed at various points along a stream filled with stream water at that point. Fish from the stream were placed in the tubs and the entire area was sprayed with Malathion by plane. Control tubs were covered during spraying and mortality of fish in all tubs was recorded after 4 hours. A similar method using aquaria was used by Schouwenberg and Jackson (1966). Snow (1963) treated pails of water from a stream with Simazine and then bass fry were placed in the pails and mortality recorded over a 96-hour period. Field studies were conducted on the toxicity of Lindane using 60 large fish tanks (1.5 m x 1.5 m x 30 cm) made from corrugated metal sheets. Each contained 50 fish and a different concentration of Lindane was used in each tank (Kok and Pathak, 1966). Gannon, et al (1966) used an experimental outdoor channel 640 feet long for water pollution studies. The channel consisted of 4-feet-long aluminum units that supported a waterproof plastic liner.

Attempts to approach more natural conditions in man-made devices have been made by other investigators. Applegate, et al (1961) and Howell, et al (1964) used running water raceways with an artificial stream bed constructed of materials from local streams, to test sea lamprey larvicides. These raceways were 6 feet wide and over 60 feet long. Productivity studies using artificial streams, supplied with water from an underground spring, were reported by Haydu (1968). The streams were 4 feet wide and ranged up to 700 feet long. Yeo (1967) used plastic pools (4 feet square by 2 feet deep) with a 2-inch layer of clay on the bottom. The pools held 180 gallons of water and aquatic plants, clams, and fish were added. A liter of natural pond water was added to introduce naturally-occurring microorganisms. These pools were used to study the influence of water hardness on dissipation and toxicity of Diquat. Parka and Worth (1965) also used plastic pools (6 feet in diameter and 15 inches deep) to study the effects of Trifluralin on fish. These pools were placed in form-fitting holes at the lowest point of a sloping field to form a catch basin. The pools were stocked with fish and the field was sprayed with a known quantity of Trifluralin. Over the next three days a sprinkler system soaked the field with ten inches of water which resulted in Trifluralin being carried into the basin in runoff water.

A more direct inethod, and one commonly used is to take qualitative and quantitative data on biota, apply the chemical to the body of water, and resample the populations. A control body of water may or may not be used. Numerous researchers have used this general approach with varied modifications (Eipper, 1959; Hoffman and Drooz, 1953; Hilsenhoff, 1966; and Surber, 1943).

Some investigators desire more control over the organisms being used in field assessments, and various methods are used to contain them. Live boxes or screened cages are commonly used. Patterson and Von Windeguth (1964) confined fish in live boxes and placed these in three shallow ponds that were sprayed with Baytex. Additional live boxes were placed in three control ponds and mortality was recorded after 24 hours. Mulla, et al (1963) and Wollitz (1963) did similar work in ponds using fish and frogs. The same technique has also been used in lakes (Jackson, 1960; Johnson, 1966; and Kallman, et al, 1962) and streams (Davis, 1954; Elson and Kerswill, 1967; Graham and Scott, 1958; Kerswill, 1967; Kerswill and Edwards, 1967; Schoenthal, 1963; and Schouwenberg and Jackson, 1966).

Another method used to restrict the movement of organisms is to enclose sections of the body of water. Harp and Campbell (1964) studied benthos in a farm pond by using plastic enclosures that divided the pond into sections measuring 12 by 18 feet. Different concentrations of Silvex were used in each section. Walker (1964) studied the effects of Dichlobenil on fish and aquatic plants in enclosures and open plots in selected farm ponds. Copeland and Woods (1969) also studied herbicidal effects on aquatic plants and used plots staked out in shallow areas of a lake. The plots were screened in with chicken wire to prevent plants from drifting away. Bonn and Holbert (1961) blocked off entire coves in a Texas lake with one-inch mesh nylon net to prevent movement of fish into and out of the cove. The coves were then treated with rotenone products.

A unique method to assess industrial pollution in a stream was used by Tatum (1966). A sampler, similar to the one designed by Hester and Dendy (1962) consisting of masonite plates, was placed in a fertilized pond for about one month to accumulate a dense growth of chironomid larvae (Diptera). These samplers were then placed in a river at stations above and below the outfall of an industrial site. Counts of larvae were made on each sampler after 1 week and comparisons were made between the average number of organisms on the samplers at stations above the outfall and on the samplers below the outfall. Williams and Mount (1965) measured the effect of zinc on periphytic communities by using a glass slide method. Periphyton populations were monitored by allowing periphyton to accumulate on glass slides submerged in running water canals for 2-week periods. One canal was used as a control and three other canals were treated with different concentrations of zinc.

The effects of chemicals sprayed into streams have been studied by monitoring the rate of downstream drifting of aquatic insects (Binns, 1967; Burdick, et al, 1960; Coutant, 1964; and Reed, 1966). Insects were continuously collected by Surber square-foot bottom samplers both before and after spraying and also in control streams. In another assessment, the effects of DDT sprayed in a stream were studied by determining the abundance of aquatic insects (Reed, 1966). An index was developed for those benthic insects found attached on rocks measuring approximately 15.2 centimeters in diameter. Butler (1965) studied the toxicity of pesticides by measuring primary productivity. By mixing known amounts of  $C^{14}$  with two suspensions of phytoplankton, one of which contains a known concentration of pesticide, it is possible to measure the interference of the pesticide with growth in a given period of time. Decreased carbon fixation provides an index of productivity, from which the relative toxicities of various pesticides may be compared. Other field methods used to detect the effects of chemicals on aquatic life include the use of other more specific radioactive tracers, the measurement of the effects of chemicals on the biochemical oxygen demand (BOD), and the fish brain cholinesterase inactivation technique. All of these methods have been discussed previously.

#### Sampling Equipment

Quantitative population samples taken to determine the effects of external factors are difficult to obtain. The effects of the external factors must be great enough to override the natural changing of the population brought about by migration, temperature, availability of dissolved oxygen, food supply, etc. Studies that require collecting organisms for evaluation also face the problem of valid sampling techniques because by definition a sample must be representative. Dimond (1967) stated that sampling procedures for stream insects are crude, and so much variation in the data results from their use that only major shifts in population size and structure can be detected. Lauer, et al (1966) said it was difficult to collect water samples that are truly representative of the concentration of the toxic agent to which the organism has been exposed.

Ricker (1968) in reference to collecting fish for productivity studies said that four truisms emerge: (1) most collecting methods are selective, with respect to species and size of individuals; (2) soundness of collecting procedures has too often been assumed and has too seldom been evaluated experimentally; (3) vast opportunities remain for discovering and developing new methods; (4) there is no substitute for operation experience on the part of the collector.

Several books provide valuable information on equipment and collecting procedures. <u>Standard Methods for the Examination of Water and Wastewater</u> (American Public Health Association, 1967), <u>Limnological Methods</u> (Welch, 1948), and <u>Ecological Methods</u> (Southwood, 1966) provide detailed information on the physical and chemical examination of water, information on equipment and methods for collecting biological material, and information on population sampling in freshwater habitats. Books by Ricker (1968) and Bennett (1962) give techniques for collecting and examining fish. The brief discussion that follows concerns only the most common methods used in the studies previously considered.

Though a wide variety of devices exist for sampling stream and lake bottoms, the three most widely used are the Ekman and Peterson dredges for lake bottoms, and the Surber square-foot sampler for shallow streams. Dredges take relatively shallow samples which are usually disturbed before they reach the surface and, therefore, the devices are not suitable for use in stratification studies. After the material is brought to the surface it is washed through a No. 30 mesh screen and the organisms sorted out. The screen collects only macroscopic bottom fauna. The Ekman dredge relies on its own weight to sink, has a rather weak spring to close the jaws and is, therefore, limited to use on bottoms which are soft and consist of finely divided mud. Large bivalves, sticks, or small rocks interfere with the closing of the jaws. The Peterson dredge is heavier, has additional attached weights, and can be used in sand and gravel. This dredge is sufficiently heavy, however, that it must be raised by a hoist. The Surber square-foot sampler is by far the most widely used stream sampler and is especially suitable for sampling on rocky bottoms which are shallow and possess current enough to hold the net in an open position. It has limited use in water deeper than three feet and again only macroscopic organisms are collected (Libby, 1964; Mackenthun, 1966; Mackenthun and Ingram, 1967; Southwood, 1966; and Welch, 1948).

Benthic and periphytic organisms are also collected by emplacement of a removable substrate. According to Southwood (1966), this is one of the most accurate collection methods. Collecting devices of this type are in various forms including building bricks (Elvins, 1962), asbestos-cement plates (Southwood, 1966), Plexiglas substrata (King and Ball, 1967), glass slides (Welch, 1948), and wire boxes containing rocks and sticks (Bull, 1968; Mason, et al, 1967; and Scott, 1958). N. W. Britt (1955) used concrete blocks on a rubble and gravel bottom to collect mayfly naiads. Unattended concrete block and Hester-Dendy multiple plate samplers are sometimes disturbed by anglers. This can be a problem when collecting devices must be left unattended in areas where large numbers of people use the water for recreational purposes. An additional problem encountered using this type of sampler especially in deep water, is that organisms not firmly attached may be lost when the sampler is raised.

The Kemmerer water sampler is probably the most widely used water collecting device and is also suitable for quantitative plankton samples. An advantage that the Kemmerer sampler has over the Juday plankton trap is that nannoplankton as well as net plankton is collected. A possible disadvantage of the Kemmerer is that motile zooplankters may tend to avoid it. The Juday plankton trap is a commonly used quantitative sampler which collects and removes the plankton in one operation. When the trap is brought to the surface, the water drains out and concentrates the plankters in a small net container. This collects only net plankton as the nannoplankton are so small they pass through the bolting cloth filter. The Juday trap is bulky, awkward to handle, and usually must be raised with a hoist. Qualitative plankton samples may also be collected with a bolting cloth tow net or with a plankton pump (Southwood, 1966; and Welch, 1948).

Ricker (1968) states that the use of electricity for capturing fish is one of the least selective of all active fishing methods. Too strong an electrical current, prolonged exposure, or contact with the electrodes, however, can kill fish, or cause damage that later proves fatal, and is of potential danger to the operators. Electrofishing can be done in both lakes and streams but water resistivity, variations in fish size, shape, or species, temperature, and fish mortality factors all have a bearing on the effectiveness of the shocker (Patten and Gillespie, 1966). Seining is the most common way to collect fish but is limited to shallow waters and bottoms that have few large boulders and few aquatic plants. Hoop and fyke nets are commonly used and according to Ricker (1968) can be both strongly selective and differently efficient in collecting fish species. For example, a net set parallel to the shoreline can be either more or less efficient than one perpendicular to it, depending on the species. Gill and trammel nets tend to be more efficient in capturing fishes adorned with external roughnesses, teeth, etc. Since these nets are stationary and depend on the fish moving to them, the fishing success may depend on abrupt changes in barometric pressure, wind-driven currents, water-level fluctuations, turbidity, and transmitted light. In very large bodies of water, purse seining and trawling are the most practical collection methods.

Table 5 shows the most commonly used items of collecting equipment, exclusive of dip nets and simple seines, with the general purpose for each item indicated. Of course, the quantitative samples may also be used to collect qualitative samples. The various traps and nets used for collecting fish result in acquiring qualitative information only. For fish population studies, some form of the capture-mark-recapture method must be used. There are many kinds of collecting devices in use though no single one is suitable for all types of habitats; a fact which complicates attempts to make comparative determinations (Anderson, 1962).

#### **Indicator Organisms**

Thieneman (Patrick, 1965) was the first to emphasize the fact that certain groups or associations of species were characteristic of a given type of environment. This does not mean however, that individual species are necessarily reliable indicators of environmental conditions in a particular area. Various researchers (Beak, 1965; Beck, 1957; Brinkhurst, 1966; Gaufin and Tarzwell, 1956; Lackey, 1957; Lackey, 1961; Mackay, 1969, Olson, 1957; Palmer, 1959; Palmer, 1963; Patrick, 1957; and Patrick, et al, 1967) have concluded that few individual species as indicators of pollution exist, but when a number of kinds of organisms are used in conjunction with chemical, physical, and bacteriological methods, the combination may be a reliable index. Table 6 is a list of organisms that have been associated with pollution of various types. When considering this table, it must be borne in mind that a number of ecological factors may influence the presence or absence of an organism and, therefore, changes in distribution and abundance of a species may not be related to pollution (Paine and Gaufin, 1956; Patrick, 1965; Lackey, 1957). Lackey (1957) pointed out that a cause and effect relationship does not necessarily exist simply because of abundance of an organism and occurrence of a defined pollutant.

Beak (1965) proposed a biotic index of water pollution based on presence and density of certain macrobenthic organisms. There were six stages in the index from normal fauna to total absence of fauna corresponding to increasing degrees of pollution. In most cases organisms were

# TABLE 5. COLLECTING EQUIPMENT IN COMMON USAGE IN LIMNOLOGICAL STUDIESAND THE GENERAL PURPOSE FOR WHICH EACH IS USED

## (Bennett, 1962; Ricker, 1968; Southwood, 1966; and Welch, 1948)

Equipment	General Purpose		
	Benthos		
Ooze sucker	Microfauna (qualitative) in uppermost layers		
	Macrofauna (qualitative) on soft bottoms		
Ekman dredge	Macrofauna (quantitative) on hard bottoms		
Peterson dredge	Macrofauna (quantitative) on smooth bottoms		
Triangle bottom dredge	Macrofauna (quantitative) on soft or hard bottoms		
Wilding square-foot sampler	Macrofauna (quantitative) on solt of hald bottoms Macrofauna (quantitative) shallow moving streams		
Dendy inverting sampler	Macrofauna (quantitative) shallow moving streams Macrofauna (quantitative) shallow moving streams		
Surber square-foot sampler	Macrofauna (quantitative) shallow moving streams		
Hess circular sampler	Macrorauna (quantitativo) sitairon moving stroums		
	Periphyton		
Hollow square-foot-sampler	Macrofauna (qualitative) from hard objects having large areas		
Wisconsin trap	Macrofauna (qualitative) from plants in shallow water		
	Plankton		
Kemmerer water sampler	Net and nannoplankton (quantitative)		
Birge cone net	Net plankton (quantitative)		
Wisconsin plankton net	Net plankton (quantitative)		
Closing net	Net plankton (quantitative) from deep water verticle tows		
Juday plankton trap	Net plankton (quantitative)		
Clarke-Bumpus sampler	Net plankton primarily deep water		
	Fish		
Hoop and Fyke traps	Quiet shallow waters		
Gill and tangle nets	Pelagic fish, various depths		
Sunken trap nets	Lower depths in relatively shallow waters		
Electric shocker	Shallow streams and lakes		
Purse seine	Open water surface seining in large bodies of water		
Trawl	Bottom, surface, or midwater depths in large bodies of water		

Organism	Type of Pollution	References
Insects		
Chironomus riparius	Organic	Gaufin, 1957; Learner and Edwards, 1966; Paine and Gaufin, 1956
C. plumosus	**	Ingram, 1957
Culex pipiens	"	Gaufin, 1957; Ingram, 1957; Paine and Gaufin, 1956; and Gaufin, 1958
C. tentans	**	Gaufin and Tarzwell, 1952
Eristalis bastardi	"	Gaufin, 1957; Gaufin and Tarzwell, 1952; and Paine and Gaufin, 1956; Gaufin, 1958
E. tenax	"	Ingram, 1957
Glyptotendipes spp	"	Paine and Gaufin, 1956
Oligochaetes		
Limnodrilus spp	Organic	Brinkhurst, 1966; Gaufin, 1957;
Tubifex spp	"	1958; and Shrivastava, 1962 Brinkhurst, 1966; Gaufin, 1957, 1958; and Gaufin and Tarzwell, 1952
Fungi		
Fusarium aquaeductum Geotrichum candidum	Organic "	Cooke, 1957
Leptomities lacteus	**	"
Penicillium lilacinum	"	»» »
P. ochrochloron	Copper	
Bacteria		
Aerobacter aerogenes	Fecal pollution	Kabler, 1957, 1961
A. cloacae	» »	Kabler, 1961
Escherichia coli		Kabler, 1957, 1961 Curtia, 1960: Harbert and Bisharda
Sphaerotilus natans	Organic	Curtis, 1969; Herbert and Richards, 1963; and Patrick, 1968
Streptococcus durans	Fecal pollution	Kabler, 1961
S. faecalis	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	"
S. liquefaciens	"	"
S. zymogenes	**	.,
Bryozoa		
Ctenostomata sp	Organic	Lackey, 1961
Protozoa		
Bodo caudatus	Organic	Lackey, 1957
Caenomorpha medusula	<b>3</b> 7	Lackey, 1961
Chaenea spp	"	"
Colpoda spp	"	Lackey, 1957
Colpidium spp	27 27	77 77
Dimastigamoeba gruberi	37 35	22
Diplophrys archeri		

## TABLE 6. PARTIAL LISTING OF ORGANISMS COMMONLY ASSOCIATED WITH POLLUTION

TABLE 6. (Continued)

Organism	Type of Pollution	References		
Protozoa (Continued)				
Enchelyomorpha vermicularis	Organic	Lackey, 1957		
Glaucoma pyriformis	,,	Lackey, 1961		
G. schintillans	"	Lackey, 1957		
Hexamitus spp	"	Lackey, 1961		
H. crassus	"	Lackey, 1957		
H. inflatus	"	"		
Loxodes vorax	**	Lackey, 1961		
Mastigamoeba spp	"	27		
Mastigella spp	"	**		
Metopus spp	>>	>>		
M. sigmoides	>>	Lackey, 1957		
Opercularia spp	"	**		
Paramecium putrinum	>>	Lackey, 1961		
Pelomyxa palustris	,,	Lackey, 1957		
Polytoma uvella	"	Lackey, 1961		
Poteriodendron petiolatum	,,	Lackey, 1957		
Saprodinium putrinum	"	**		
Spirostomum spp	"	Lackey, 1961		
Strombidium spp	**	22		
Tetramilus spp	"	Lackey, 1957		
T. pyriformis	"	Lackey, 1957, 1961		
Tillina magna	••	Lackey, 1961		
Trachelocerca coluber	••	"		
Trepomonas spp	"	Lackey, 1957, 1961		
Trigonomonas compressa	>>	"		
Trimyema compressa	"	>>		
Uahlkampfia guttalu	"	Lackey, 1957		
U. limax	"	"		
Urocentrum turbo	"	Lackey, 1957, 1961		
Uroleptus spp	"	Lackey, 1961		
Urophagus rostratus	"	Lackey, 1957		
Urotricha spp	"	Lackey, 1961		
Urozona butschlii	"	"		
Algae				
Achanthes affinis	Hydrogen sulfide	Palmer, 1959		
A. minutissima	Calcium carbonate	Patrick, 1965		
Achnanthidium brevipes	Salt brine	Palmer, 1959		
var intermedia	(principally NaCl)	, "		
Actinastrum hantzschii	"	"		
Actinella spp	High acidity	Palmer, 1959, and Patrick, 1957		
Agmenellum quadriduplicatum	Organic	Palmer, 1959, and Patrick, 1957		
Amphora coffeiformis	Salt brine (principally NaCl)	"		
Amphora cojjeljornus A. ovalis	Paper mill wastes, salt brine, oil	"		
Anabaena constricta	Organic	**		
	Salt brine (principally NaCl)	"		
Anacystis spp A. montana	· · · · ·	"		
A. montana Anomoeoneis serians var.	Organic Iron	"		
	1011			
brachipira	Organic	>>		
Arthrospira jinneri	Organic			

Туре	of Pol	lution	n	

Organism

References

Algae (Continued)		
Astasia spp	Organic	Lackey, 1957
Asterionella formosa	Copper	Palmer, 1959
Caloneis amphisbaena	Paper mill wastes, hydrogen sulfide	,,
Calothrix spp	Salt brine (principally NaCl)	"
C. braunii	Copper	,,
Camphlodiscus spp	Hydrogen sulfide	27
Carteria multifilis	Organic	••
Ceratoneis arcus	Phenolic wastes	,,
Chaetomorpha spp	Salt brine (principally NaCl)	**
Chlamydobotrys spp	Distillery wastes	,,
Chlamydomonas spp	High acidity	**
C. ehrenbergii	Salt brine	**
C. reinhardi	Organic	33
Chlorella pyrenoidosa	"	"
C. vulgaris	"	**
C. variegata	Iron	"
Chlorobrachis spp	Organic	Lackey, 1957
C. gracillina	Distillery wastes	Palmer, 1959
Chlorococcum botryoides	Copper	, , , , , , , , , , , , , , , , , , ,
C. humicola	Organic	"
C. numicola Chlorogonium euchlorum	Distillery wastes, organic	,,
Chromulina spp	Iron	>>
C. ovalis	High acidity	>>
C. ovaiis Closterium acerosum	Chromium	,,
Coccachloris elabens	Salt brine (principally NaCl)	,,
(Aphanothece halophytica)	Sait office (principally Nacr)	
Cocconeis diminuta	Paper mill wastes	"
C. pediculus	"	"
C. placentula	Phenolic wastes	"
Cryptoglena pigra	Organic	"
Cryptomonas erosa	High acidity	,,
Cyclotella kiitzingiana	Phenolic wastes	"
C. meneghiniana	Hydrogen sulfide, salt brine	>>
Cymatopleura solea	Phenolic wastes, paper mill wastes	"
Cymbella lacustris	Salt brine (principally NaCl)	,,
-	Copper, phenolic wastes	,,
C. naviculiformis C. ventricosa	Salt brine, paper mill wastes, copper,	,,
C. veniricosa	hydrogen sulfide	
Diatoma elongatum	Salt brine (principally NaCl)	"
D. vulgare	Phenolic wastes, paper mill	"
Diploneis elliptica	Wastes, oil	"
Dunaliella salina	Salt brine (principally NaCl)	27
Enteromorpha intestinalis	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	,,
E. prolifeia	,,	**
Entophysalis deusta	"	**
(Aphanocapsa littoralis)		
Euglena spp	>>	"
E. acus	Chromium	,,
E. adhaerens	High acidity	>>
E. agilis	Organic	>>

Organism	Type of Pollution	References
Algae (Continued)		
E. deses	Organic	Palmer, 1959
E. gracilis	31	22
E. hiemalis	High acidity	"
E. mutabilis	"	Lackey, 1957; Palmer, 1959; and Sundaresan, et al, 1965
E. o <b>xguris</b>	Organic, chromium	Palmer, 1959
E. polymorpha	Organic	Lackey, 1959, and Palmer, 1959
E. sociabilis	Chromium	Palmer, 1959
E. stellata	Chromium, high acidity	Lackey, 1959, and Palmer, 1959
E. tatrica	High acidity	**
E. viridis	Chromium, high acidity, organic	**
Eunotia spp	Iron, high acidity	Palmer, 1959, and Patrick, 1957
E. exigua	High acidity	Lackey, 1957, and Palmer, 1959
E. lunaris	······································	"
E. trinacria	"	"
Fragilaria virescens	Phenolic wastes	Palmer, 1959
Frustulia rhomboides var	Salt brine (principally NaCl)	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
saxonica	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	"
<i>Fomphonema</i> spp	,,	>>
F. acuminatum	Iron	"
F. herculaneum	Paper mill wastes, oil	**
F. olivacuum	Calcium	Patrick, 1965
F. parvulum	Phenolic wastes, organic	Palmer, 1959
Fyrosigma attenuatum	Salt brine (principally NaCl)	, , , , , , , , , , , , , , , , , , ,
		"
lantzschia amphioxys Lalauaata	Hydrogen sulfide, organic	>>
I. elongata	Salt brine (principally NaCl)	
epocinclis ovum	High acidity, organic	Lackey, 1957, and Palmer, 1959
. texta	Organic	Palmer, 1959
yngbya astuarii	Salt brine (principally NaCl)	22
. digueti	Organic	>>
Ielosira arenaria	Salt brine (principally NaCl)	>>
1. varians	Oil, organic	22
Ieridion circulare	Salt brine (principally NaCl)	
Aicrocoleus chthonoplastic	,,	"
lavicula anglica		**
l. atomus	Chromium	37
l. cincta var heufleri	Salt brine (principally NaCl)	**
l. c <b>r</b> yptocephala	Salt brine, organic, phenolic wastes, paper mill wastes	"
V. gregaria	Salt brine (principally NaCl)	>>
V. linearis	Chromium	"
I. longirostris	Salt brine (principally NaCl)	"
V. minima	Hydrogen sulfide	"
V. minuscula	Salt brine (principally NaCl)	>>
V. palea	Chromium, organic	"
V. pygmaea	Salt brine (principally NaCl)	Palmer, 1959, and Patrick, 1957
N. radiosa	Paper mill wastes, oil	Palmer, 1959
N. salinarum	Salt brine (principally NaCl)	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
subtilissima	High acidity, salt brine	Lackey, 1957, and Palmer, 1959
V. viridis	High acidity, copper	, , , , , , , , , , , , , , , , , , ,

Algae (Continued) Veidium bisulcatum Vitzschia acicularis V. apiculata V. epithemoides V. frustulum	Copper Organic	Palmer, 1959		
Nitzschia acicularis V. apiculata V. epithemoides	Organic	Palmer, 1959		
Nitzschia acicularis V. apiculata V. epithemoides	Organic	- <b></b> ,		
N. apiculata N. epithemoides	-	**		
V. epithemoides	Salt brine (principally NaCl)	33		
-	"	33		
	**	>>		
V. ignorata	Hydrogen sulfide	**		
V. palea	Phenolic wastes, hydrogen sulfide, salt brine	"		
N. trybliowella var debilis	Hydrogen sulfide	**		
Ochromonas spp	High acidity	Lackey, 1957, and Palmer, 1959		
Oscillatoria spp	Paper mill wastes, salt brine	Palmer, 1959		
D. chalybea	Organic	"		
D, chlorina	»	>>		
D. formosa	"	"		
0. lauterbornii	"	"		
0. limosa	>>	"		
0. princeps	>>	"		
	"	>>		
O. putrida	>>	**		
O. tenuis Brudoring ann	Demon mill streeter	**		
Pandorina spp	Paper mill wastes	Laskey 1057 and Dalman 1050		
P. morum	Organic	Lackey, 1957, and Palmer, 1959		
Pediastrum spp	Paper mill wastes	Palmer, 1959		
P. simples	Salt brine (principally NaCl)	27		
Penium cucurbitinum	High acidity			
Phacus parvulus	Organic	Lackey, 1957		
P. pyrum	22	Lackey, 1957, and Palmer, 1959		
Phormidium autumnale		Palmer, 1959		
P. tenue	Salt brine (principally NaCl)	22 22		
P. uncinatum	Organic	>>		
Pinnularia spp	High acidity, iron, salt brine	22		
P. borealis	Phenolic wastes			
P. subcapitata var helseana	Iron	**		
Platymonas spp	Organic	Lackey, 1957		
Polytoma citri	32	**		
P. uvella	**	"		
Pyrobotrys gracilis	**	Lackey, 1957, and Palmer, 1959		
P. stellata	"	"		
Scenedesmus spp	Paper mill wastes	Palmer, 1959		
S. bijugatus	Salt brine (principally NaCl)	"		
S. obliquus	Copper	"		
S. quadricauda	Organic	"		
Spirogyra communis	»	"		
Spirulina subsalsa	Salt brine (principally NaCl)	22		
Spondylomorum spp	Paper mill wastes	"		
Stauroneis anceps	High acidity	Lackey, 1957, and Palmer, 1959		
S. phoenicentern	Iron	Palmer, 1959		
Stenopterobia intermedia	27	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
Stephanaptera gracilis	Salt brine (principally NaCl)	"		
Stichococcus bacillaris	Organic	"		

Organism	Type of Pollution	References
Algae (Continued)		
Stigeoclonium tenue	Organic	Curtis, 1969, and Palmer, 1959
Surinella delicatissima	Iron	Palmer, 1959
S. linearis	"	**
S. ovata	Paper mill wastes, phenolic wastes,	"
	organic	
S. ovata var salina	Paper mill wastes, phenolic wastes,	"
S. Ovata val satina	hydrogen sulfide, organic	
Symploca erecta	Copper	"
Synedra acus	Oil, salt brine	"
-	Salt brine (principally NaCl)	Palmer, 1959, and Patrick, 1957
S. affinis S. mulahalla	Paper mill wastes, salt brine	, <u>, , , , , , , , , , , , , , , , , , </u>
S. pulchella S. ulna	Paper mill wastes, phenolic wastes, oil	Palmer, 1959
	<b>.</b>	, , , , , , , , , , , , , , , , , , ,
Tabellaria flocculasa	High acidity	"
Tetraedron muticum	Organic	,,
Tetraspora spp	Chromium	"
Trachelomonas spp	Salt brine (principally NaCl)	57
T. hispida	Iron	22
Trichodesmium spp	Salt brine (principally NaCl)	22
Ulothrix spp	Salt brine, paper mill wastes	
U. zonata	High acidity	Lackey, 1957, and Palmer, 1959
Vanheurckia rhomboides var	"	33
crassenervia	"	
Xanthidium antilopaeum	**	Palmer, 1959

identified only to Family and were grouped according to feeding type and sensitivity to pollution.

In using groups of aquatic organisms as indicators of pollution, the absence or reduction in numbers of "clean-water" species may be as important, if not more so, than the presence of known pollutional forms (Anderson, 1962; Fremling, 1964; Gaufin, 1958, 1965; Gaufin and Tarzwell, 1952; and Leonard, 1965). Aquatic organisms usually considered to be "clean-water" organisms include mayflies, stoneflies, caddisflies, molluscs of the family Unionidae, and beetles of the family Elmidae. The absence of these organisms and the presence of physid snails, tubificids, *Eristalis tenax*, and *Chironomus pipiens* would indicate water highly degraded by organic wastes (Hinshaw, 1967; Ingram, 1957; Paine and Gaufin, 1956; and Young, 1961). Palmer (1959) lists over 40 species of algae that he considers "clean-water" forms. He also said that blue-green algae and flagellates are the algal groups most frequently encountered in the portion of a stream containing organic pollution. Palmer (1963) has compiled a listing of more than 600 species that are said to be tolerant of pollution.

The presence of large number of tubificids usually indicates a high concentration of organic matter. These worms can live in water low enough in oxygen that most other fauna will not survive (Brinkhurst, 1966, and Curry, 1965). King and Ball (1964) used wet weight ratios of tubificids to aquatic insects to indicate changes in water quality. Their results indicated that this technique may be useful in measuring organic pollution. Among the mayflies, there seems to be an order of sensitivity to organic waste and as pollution increases sensitivity declines in the following order: *Rhithrogena, Heptagenia, Ecdyonurus, Ephemerella,* and *Baetis.* An amphipod, *Gammarus pulex,* lives quite well even in badly polluted water as long as the oxygen content is not greatly lowered (Hynes, 1959). Ingram (1957) in discussing clams and snails, said that not enough is known about molluscan ecology to name any species a pollution indicator and though species such as *Psidium idanoensis, Physa integra, P. heterosteopha,* and *Musculium transversum* are found associated with organic waste.

Coliform bacteria are constantly present in alimentary discharges, are comparatively easy to enumerate, have long been considered indicative of fecal pollution (Gilderhus, 1966; and Kabler, 1957, 1961). Owing to special nutritional requirements a few species of fungi have been associated with certain types of pollution (Servizi, et al, 1966). Generally, however, there has been little correlation found between pollution and populations of aquatic fungi (Cooke and Bartsch, 1959).

Brinkhurst (1966) said that fish are not particularly easy to use as indicators because they are relatively difficult to sample, and their mobility makes it possible for them to avoid those parts of the environment which become intolerable for short periods of time. Katz and Gaufin (1953) studied the effects of organic pollution on fish distribution in a small Ohio stream. No species of fish were regarded as indicators of pollution although several were relatively tolerant of unfavorable conditions. They concluded that the number of species present and their relative abundance are the most important considerations when pollutional conditions are being evaluated.

Williams (1964) concluded that the search for biota or communities of biota which might be useful as indicators of water quality has been hampered by the lack of information on the environmental requirements of the various species and their resistance to specific chemical substances.

#### Concluding Remarks (Field Assessment)

The value of field studies lies in the fact that more natural conditions are approached in the field than in the laboratory. This is important because the reaction of an organism to a chemical in the laboratory is not necessarily the same as it would be in nature. A price is paid for these natural conditions, however, because it is impossible to control or even to ascertain all of the variables in a field study. To complicate this further, in most field work there is a conspicuous lack of detailed water-quality data taken in support of the field observations. In this report, for example, approximately 220 papers dealing with field projects were carefully studied and evaluated. Of these, only about 50 contained definitive water quality information. It has long been recognized that the toxicity of a compound may depend on a number of interrelated factors, including temperature, pH, water hardness, dissolved oxygen content, and exposure time. For example, Cairns (1957) showed that considerable increases in toxicity may result during periods of low dissolved oxygen content, and that this may occur even when the oxygen supply is not low enough to be directly harmful to the organism. Burdick (1967) states that toxicants react with detritus, and organic or inorganic materials in the water or bottom sediments and that bacterial decomposition may alter chemicals to substances of greater or less toxicity. He concluded that even light penetration may have an effect. Only rarely are all or even a majority of these factors taken into consideration in conducting field studies of water pollution.

#### SECTION IX

#### FACTORS AFFECTING CHEMICAL TOXICITY IN WATER

Depending on the nature of a chemical, environmental factors influencing water quality may also affect the inherent toxicity of that compound to aquatic biota. Similarly, water quality itself can affect chemical toxicity. For these reasons, chemical-physical characterization of water is important whether it is used in a bioassay or studied in the field. Experimentation may have little significance without minimal characterization, that is, measurement of water temperature, pH, dissolved oxygen (DO), conductivity, oxidation-reduction potential, dissolved chlorides, and turbidity. Furthermore, when potentially toxic ions, e.g., heavy metals or halogens, are known or suspected to be present, analysis for these should be made. Without such data for an aquatic experiment, the toxicity of a chemical to an aquatic organism means only that for the conditions of that experiment is the chemical toxic at the concentration level reported, i.e., the toxicity data cannot be extended to any other type of water.

As pointed out previously in other sections of this report, this type of water characterization data was seldom given in the publications reviewed. Use of an unspecified, "standard water" throughout a bioassay study helps very little when an attempt is made to extrapolate from the study and predict how a chemical may behave in an entirely different water. If there is to be a serious attempt to employ multivariate analysis or mathematical modeling in predictive studies of chemical pollution problems, then the suggested type of water data must be taken, or completely standardized experimental conditions including chemically defined water must be employed. The following discussions concern the more important water-quality factors that may affect the toxicity of a chemical in aquatic environments.

#### Temperature

The biological significance of temperature in the aquatic environment has been recognized for many years. It was once said that a limnologist could obtain more information about a body of water with a thermometer than any other single instrument. Reid (1962) believes "from the broad and basically ecological point of view, the thermal properties of water and the attending relationships are doubtless the most important factors in maintaining the fitness of water as an environment." In several limnology texts (Reid, 1961, Ruttner, 1953, and Welch, 1952), accounts are given of thermal stratification, thermoclines, heat budgets, general thermal dynamics of water bodies, and the effects these factors have on aquatic life. Hutchinson (1957) gives an in-depth account of the thermal properties of lakes. In recent years as the use of streams and lakes by industry has increased, more investigators have been concerned with the effects of increased temperatures on aquatic organisms. There are several very recent, extensive bibliographies (over 1500 references) available on heated effluents and their effects on aquatic life (American Society of Civil Engineers, 1967; Kennedy and Mihursky, 1967; and Raney and Menzel, 1967). A reference manual on thermal effects on aquatic organisms was prepared by Wurtz and Renn (1965).

A great deal of attention has been placed on thermal effects on fish. Fish, like most aquatic organisms, are poikilotherms and therefore lack the means of maintaining an independent body temperature. Needless to say, water temperature is a critical factor in the life of a fish and in fish production. Each species has a thermal zone in which it can function in a normal manner with a higher and lower zone in which it can survive for certain lengths of time. The degree of

success the fish will have in these less than optimal zones will depend on a multitude of factors including the health of the fish, stage of development, sex, diet, season of the year, and various water quality parameters (Alabaster, 1967; Alabaster and Welcomme, 1962; Brett, 1956; Hoar, 1956; Huet, 1965; Mihursky and Kennedy, 1967; Tarzwell, 1957; and Tyler, 1966).

A major factor affecting the ability of an organism to adapt to a new temperature is the previous temperature to which it has been exposed. Prosser and Brown (1961) define acclimation as the compensation by animals to persistent change in temperature, usually in the laboratory. Though not all authors make the distinction between acclimation and acclimatization, Prosser and Brown refer to acclimatization as compensations under field conditions which come about more slowly. Upper lethal temperatures tend to be closer to the acclimation temperature than lower lethal temperatures (Colton, 1959). Upper or lower lethal temperatures obviously have more meaning when the acclimation temperature is indicated. Table 7 lists the thermal death points of a number of species of freshwater and marine fish in relation to the acclimation temperatures. The table is a summary of work conducted by Brett (1956) and Jones (1964).

Laboratory studies conducted on thermal death points of various organisms may be of two basic types. These are acute or shock tests in which large temperature increases are usually completed in a few hours, and the chronic tests in which temperature increase is only a degree or two a day and the overall test lasts several months. Shock tests are of value in studying fish movements or when thermal loading is confined to a limited area. In these situations fish are likely to move rapidly from one temperature zone to another. Chronic tests are designed to approximate a condition of gradual exposure over considerable periods of time (Cairns, 1955, 1956).

Generally, fish of temperate regions are able to tolerate temperatures from 0 C to 30 C but resistance to the highest and lowest temperature varies with different species. Salmonids and other cold water fishes do not tolerate higher temperatures while warm water forms, such as the cyprinids, tolerate higher temperatures quite well. Marine species may be more sensitive to temperature change than freshwater species and immatures of both types are more sensitive than adults. In general, all abrupt changes in temperature can be harmful even if the changes are short lived.

Temperature may affect the fish directly or it may have an indirect effect. A change may be within the toleration limits of a fish but may alter the environment to the point where it is more suitable for another species (Tarzwell, 1957). This may come about in a number of ways including a reduction or an increase in food supply, interference with the spawning process, or alteration of the dissolved oxygen content of the water. Though other factors are also involved, fish only spawn when the water reaches a suitable temperature and this varies with different species. Water temperature may affect growth. For example, carp growth is very good between 20 C and 28 C, average between 13 C and 20 C, poor between 15 C and 13 C, and non-existent below 5 C (Alabaster, 1967; Colton, 1959; Fry, 1960; Huet, 1965; and Swift, 1965).

Though the physiological effects of heat on an organism are discussed in some detail by Brown (1957) and Prosser and Brown (1961), the actual cause of death by either heat or cold is not well understood. Various theories have been put forth concerning the mechanism of heat death including coagulation of protoplasm, inactivation of enzyme systems, lack of oxygen due to inactivation of the respiratory center, and the release of toxic materials from heat affected cells (Brett, 1956; Brown, 1961; Cairns, 1955; and Jones, 1964). Though the exact causes of death at high temperatures may not be clear, most investigators agree that multiple factors are involved.

## TABLE 7. THERMAL DEATH POINTS OF FISH ACCLIMIZED AT THE INDICATED TEMPERATURES(FRESHWATER = F, MARINE – ATLANTIC = A, PACIFIC = P)

Fish	Acclimation Temperature, C	Thermal Death- Point, C	Occurrence	
Atlantic salmon	_	29.5-30.5	A-F	
Atlantic salmon (grilse)	_	32.5-33.8	F	
Atlantic salmon (parr)	_	29.8	F	
Blacknose dace	10	28.8	F	
Blacknose dace	20	29.3	F	
Bluegill	15	30.7	F	
Bluegill	20	31.5	F	
Bluegill	30	33.8	F	
Bluntnose minnow	25	33.3	F	
Brook stickleback	25-26	30.6	F	
Brook trout	5	23.7	A-F	
Brook trout	10	24.4	A-F	
Brook trout	15	25	A-F	
Brook trout	20	25.3	A-F	
Brook trout	25	25.3	A-F	
Brown bullhead	15	31.8	F	
Brown bullhead	20	33.4	F	
Brown bullhead	30	36.5	F	
Brown trout	26	26	A-F	
Brown trout (fry)	5-6	22.5	F	
Brown trout (fry)	20	23	F	
Brown trout (yearling)	_	25.9	A-F	
Brown trout (parr)		29	A-F	
Carp	20	31-34	F	
Chinook salmon (fry)	15	25	F	
Chinook salmon (fry)	20	25.1	F	
Chum salmon (fry)	15	23.1	F	
Chum salmon (fry)	20	23.7	F	
Coho salmon (fry)	15	24.3	F	
Coho salmon (fry)	20	25	F	
Common shiner	15	30.3	F	
Common shiner	30	31.0	F	
Creek chub	10	27.3	F	
Creek chub	15	29.3	F	
Creek chub	25	30.3	F	
Emerald shiner	10	26.7	F	
Emerald shiner	15	28.9	F	
Emerald shiner	25	30.7	F	
Fathead minnow	10	28.2	F	
Fathead minnow	20	31.7	F	
Fathead minnow	30	33.2	F	
Gizzard shad	25	34.3	A-F	
Gizzard shad	30	35.9	A-F A-F	

#### (Brett, 1956; and Jones, 1964)

TABLE 7. (Continued)

Fish	Acclimation Temperature, C	Thermal Death- Point, C	Occurrence
	15	30.5	F
Golden shiner	15 25	33.2	F
Golden shiner	30	34.7	F
Golden shiner			F
Goldfish	10	30.8	F
Goldfish	20	34.8	F
Goldfish	30	38.6	
Guppy	30	34	F
Largemouth bass	20	32.5	F
Largemouth bass	25	34.5	F
Largemouth bass	30	36.4	F
-	15	35.4	A-F
Mosquito fish Mosquito fish	20	37.3	A-F
Mosquito fish	30	37.3	A-F
•		31.4	Р
Opaleye	20 30	31.4	P
Opaleye	30		
Perch	_	23-25	F F
Perch	10	25.0	r F
Perch	15	27.7	г F
Perch	25	29.7	
Pink salmon (fry)	5	21.3	F
Pink salmon (fry)	10	22.5	F
Pink salmon (fry)	20	23.9	F
Pumpkinseed	25-26	34.5	F
Rainbow trout	_	28	A-F-P
Rainbow trout (Kamloops var)	11	24	P-F
Roach	20	29.5	F
Roach	25	30.5	F
Roach	30	31.5	F
Sockeye salmon (fry)	5	22.9	F
Sockeye salmon (fry)	10	23.4	F F
Sockeye salmon (fry)	20	24.8	
Tench	—	29-30	F
White sucker	25	29.3	F
Yellow Perch	15	27.7	F

When the temperature goes beyond the thermal zone optimal for the organism, evidence indicates the general resistance to other adverse conditions is reduced. Hynes (1959) stated that several workers have shown that a rise of 10 C may halve the survival time of test animals. It has been reported that an increase in temperature caused an increase in toxicity in fluorides (Angelovic, et al, 1961), cyanide (Cairns and Scheier, 1963), sodium pentachlorophenate (Crandall and Goodnight, 1959), phenol (Brown, et al, 1967), various pesticides (Mahdi, 1966, and Macek, et al, 1969), as well as a possible reduction in resistance to disease (Cairns, 1955, and Turnbull, et al, 1954). It has also been reported that anesthesia with alcohol was induced more rapidly in fish when the temperature was increased. Though it may not appreciably affect the toxic threshold, an increase in temperature may affect the length of time required for a given concentration to kill an organism. Hester (1959) found that if 40 F tests were continued beyond 3 days, the kill of fish by the end of the twenty-first day was approximately the same as 70 F tests conducted for 3 days. When all tests were run at 3 days, however, more rotenone was required to kill fish at 40 F than at 70 F. Similar findings were reported by Lloyd (1965) and Cairns and Scheier (1957). The rate of uptake of chemicals by aquatic organisms increases with an increase in temperature (Das and Needham, 1961). This occurs probably because of the increase in metabolic rate which accompanies the increase in temperature.

An interesting example of the effects of temperature on fish behavior was reported by Loeb, et al (1966). Brown bullheads *(Ictalurus nebulosus)* were killed when exposed to 50 ppb of 4-iodo-3-salicylanilide at temperatures of 5 C or 21 C. When bottom sediments were added, the bullheads would bury themselves in the sediment at 5 C and thus escape the toxic chemical. At 21 C, however, the fish would not bury themselves and were killed by the chemical.

Results of field studies conducted to determine the effects of increased temperatures on aquatic life are usually recorded as a reduction in numbers of individual organisms, reduction in species (with or without reduction in numbers of individuals), or the presence of indicator organisms (Geen and Andres, 1961; Mann, 1965; Trembley, 1960; and Wurtz and Dolan, 1961). Various types of organisms are useful in these studies. Trembley (1965) conducted a five year study of heated discharges in a Pennsylvania river and outlined the types of useful organisms and made some brief remarks about each group. The numbers of species of periphyton tended to be reduced in high temperatures but individual species were often present in great numbers. Most aquatic invertebrates tended to increase during winter months and undergo reduction in the summer. Insect larvae of the family Tendipedidae were the most tolerant invertebrates in the heated water areas. A rooted aquatic plant, Potomogeton, was found growing well in temperatures ranging from 35 C to 37 C. Certain species of blue-green algae, primarily Oscillatoria, were found to be the most heat-tolerant and were observed growing well in temperatures up to 45 C. During the summer, fish left the heated-water zone and were apparently attracted to the heated water areas during the winter months. Plankters drifted with the current and because of this were not considered suitable organisms to work with in lotic environments.

The Aquatic Life Advisory Committee (1956) in discussing water quality requirements for freshwater fish concluded that "any change in the temperature of the aquatic habitat will affect the animals and plants living in it, even though the change remains within their ranges of thermal tolerance. Because there is a relationship between temperature and the solubility, dissociation and stability of the substances dissolved or suspended in water, a change in temperature will have an indirect effect upon aquatic organisms, entirely apart from any direct effect, through alteration of the physical and chemical characteristics of their environment. Since body temperature of a fish or lower aquatic organism is very close to that of the water, a change in temperature will have direct effect by action upon the metabolic rate, growth, reproduction and other vital processes. It should be pointed out further that, as a consequence of the temperature effect upon one species, a change in temperature might alter the biotic environment of another species, thereby affecting the latter indirectly through an increase or decrease in food or shelter. The complexity of the problem is increased by the fact that the nature and magnitude of the effects upon aquatic organisms are related, not only to the temperature itself, but also to the rate at which it is changed and to the duration of the altered level".

#### pН

The most frequently used index of hydrogen ion activity is pH. The pH of natural waters may range from extremes of 1.7, found in an African lake, to 12.0 recorded from some Japanese lakes. Normally however, surface water pH is between 6.0 and 9.0. Factors influencing pH in unpolluted bodies of water are currents, which serve to keep the waters mixed; biological processes such as photosynthesis and respiration; and the composition of the rocks and sediments of the substrate (Jordan and Lloyd, 1964; National Technical Advisory Committee, 1968; and Reid, 1961). Hutchinson (1957) states that in practically every case where the water is neither very acid nor very alkaline, it may be assumed that the pH is regulated by the carbon dioxide-bicarbonate-carbonate system.

Determination of pH is not a measure of total acidity or alkalinity in water. Many compounds may be in water in unionized portions of weakly ionizing acids such as phosphoric, carbonic, fatty acids, protein compounds, or as hydrolyzing salts such as ferrous or aluminum sulfate. The latter are referred to as acid buffers. When acidity is measured by titration using a dye like methyl orange with an end-point at pH 4.5, the value is termed "free acidity". If the titration is carried by alkali addition to the end point of phenolphthalein at a pH of 8.3, the value is called "total acidity" and will include the weak acids, acid salts, and with sufficient time for reaction between alkali additions, some acidity due to slowly hydrolyzable compounds.

Alkalinity is usually imparted by the bicarbonate, carbonate, and hydroxide components of a natural or treated water supply. These ions are the so-called alkali buffers. In determining alkalinity, if the solution is titrated to the phenolphthalein end point of 8.3, the alkali fraction measured is that contributed by the hydroxide and half of the carbonate. Indicators responding in the pH range of 4-5 are used to measure the "total alkalinity" contributed by the hydroxide, carbonate, and bicarbonate.

Alkaline buffering capacity of water in some limestone areas, for example, may partially neutralize acidic components of an effluent. Where carbon dioxide content is high, alkali components of a waste effluent may be partially neutralized. Total acidity and alkalinity are features of water quality that are often overlooked in considering effluent release, and also in conducting bioassay or field studies of chemical toxicity.

When pH is the only factor considered, the toleration limit of most organisms falls in the range of 5.0 to 9.0 (Jones, 1964; Doudoroff and Katz, 1950; and Hynes, 1966). Fry (1960) concluded that the general range for good fish production was 6.7 to 8.6. McKee and Wolf (1963) state that of waters which support a good fish fauna, only 5 percent have a pH of less than 6.7 and only 5 percent have a pH over 8.3. The permissible range for fish depends on several factors including temperature, age, dissolved oxygen, prior acclimatization, and the content of various anions and cations.

The exact cause of death of fish in low or high pH waters is unclear though Tarzwell (1957) has stated that an unsuitable pH may interfere with oxygen uptake. It has been reported (Jones, 1964, and Aquatic Life Com., 1955) that fish are killed in acid waters by precipitation and coagulation of the mucous on the gills and by coagulation of the gill membranes themselves.

The pH of water may have considerable influence on the toxicity of certain chemicals. The pH value will determine the degree of dissociation of weak acids and bases, some of which may be more toxic in molecular than ionic form (McKee and Wolf, 1963; Hynes, 1966; and Cairns and Scheier, 1963). Highly dissociated inorganic acids do not appear to be toxic at pH values above 5.0 and highly dissociated inorganic alkalies do not appear to be toxic below 9.0 (Aquatic Life Com., 1955).

The effect of pH on the toxicity of specific compounds has been reported. An increase in toxicity brought about by a decrease in pH was reported for pentachlorophenol and sodium pentachlorophenate (Goodnight, 1942, and Crandall and Goodnight, 1959), nickel cyanide (McKee and Wolfe, 1963), and sodium sulfide (McKee and Wolfe, 1963, and Tarzwell, 1957). Within certain ranges, pH may have little or no effect on toxicity. Henderson, et al (1958, 1959) reported no differences in toxicity for several chlorinated hydrocarbon insecticides when the pH was varied from 7.4 to 8.2. Loeb, et al (1965) conducted studies on ergot derivatives on surfacing behavior of fish, and found no change in response when pH was changed from 6.3 to 7.2. Marking and Hogan (1967) found little difference in toxicity of Bayer 73 to fish in a pH range between 6.4 to 8.0. At a higher pH (10.0) and a lower pH (5.0), the toxicity of this compound was reduced. Mount (1966) in a flow-through study showed that zinc was always more toxic at a high pH than at a low pH, and further that water hardness was also an important factor.

#### **Dissolved Oxygen**

The amount of dissolved oxygen (DO) present is one of the most significant chemical parameters in the study of surface waters. The amount of oxygen that can be dissolved in water at any one time is dependent upon (1) water temperature, (2) partial pressure of the oxygen in the atmosphere in contact with the water, and (3) salinity.

Photosynthesis in algae and higher aquatic plants is one source of DO in natural waters. The rate of photosynthesis depends on many factors but the major one is light. The depth that light penetrates the water (euphotic zone) is determined by turbidity, color, and the absorptive effect of the water itself. Another important source of oxygen is the atmosphere. Factors which will influence the rate at which oxygen will dissolve into the water from the atmosphere include (1) wave action, or other surface disturbances, (2) the difference in partial pressure between the atmosphere and the water, and (3) the moisture content of the atmosphere.

There may be considerable diurnal and seasonal fluctuations in DO in a stream or lake primarily due to changes in water temperature and photosynthetic rates. Water temperatures vary from one season to another and deep lake water may vary considerably from the surface to the bottom, e.g., during thermocline formation. Though photosynthesis does not occur at night, aquatic plant respiration continues and oxygen is utilized. The amount of oxygen that is used in aerobic biochemical action in the decomposition of organic matter (BOD) also causes extreme fluctuations in DO available for aquatic organisms.

Oxygen requirements of fish and other aquatic organisms vary with the species and are affected by age, degree of activity, size, prior acclimatization, and health of the organism. Environmental factors influencing DO requirements or interfering with oxygen uptake are temperature, pH, carbon dioxide, and dissolved solids. Temperature appears to be the major factor because as the temperature increases, the metabolic rate of cold-blooded animals increases along with oxygen uptake. At the same time, the solubility of oxygen in water decreases as temperature increases. This is discussed in excellent detail with a tabulation of the water solubility of oxygen in Standard Methods (American Public Health Association, 1967).

Jones (1964) summarized the work of various investigators (Table 8) who conducted laboratory studies on DO requirements of fish at various temperatures. Jones pointed out that these figures were somewhat low compared with observations made in the field at similar temperatures. It follows, however, that while fish may survive short periods of stress under laboratory conditions, this does not mean they will be able to survive indefinitely, feed, reproduce, grow, and compete with other organisms.

Doudoroff and Warren (1962) found that sublethal adverse effects of low DO on fish included reduction in swimming speed and loss of weight. The gross efficiency of food conversion was not greatly reduced in fish maintained on an unrestricted diet until the DO level dropped below 4 ppm. The reduction in growth rate was attributed to loss of appetite. It was also found that sac fry hatched from eggs in waters with a low DO content were small and weak.

A low level of DO may in itself be a lethal factor for various aquatic organisms and may also cause an increased toxicity in a variety of chemicals. Several investigators have reported an increase in the toxicity of chemicals due to decreased DO including various petroleum products (Tagatz, 1961), unionized ammonia (Downing and Merkens, 1955), potassium dichromate (Cairns, 1965), potassium cyanide (Downing, 1954; and Cairns, 1965) zinc, lead and copper salts (Reiff, 1964), and various other inorganic salts (McKee and Wolf, 1963).

#### Suspended Solids and Turbidity

Turbidity may be defined as the degree of opaqueness produced in water by suspended particulate matter. In much of the literature, turbidity and suspended solids (or suspensoids) are used as synonyms. The particle size, shape, and refractive index have more influence on turbidity than weight composition (American Public Health Association, 1967). The interplay of light on the suspended material along with the reflection from the sky or bottom are also responsible for the apparent color of the water. This is distinguished from true color which is derived from substances in solution or in the colloidal state.

Turbidity is measured in Jackson turbidity units (JTU) which is the distance through a column of water at which the image of a standard flame from a candle is no longer visible. The standard unit is that condition produced by 1 ppm Fullers earth in distilled water. Turbidity has a profound effect on natural light penetration which can be determined by the use of a photronic cell or a Secchi disk. The measure of natural light penetration, however, is not a good measure of turbidity because other factors affect light penetration including intensity, cloud cover, water disturbance, and direction of the sunlight.

Suspended solids that occur naturally in water bodies include plankton, organic and inorganic detritus, and silt. These suspended solids are augmented by a multitude of materials in discharges from population centers, agricultural, and industrial sites. McKee and Wolfe (1963) note that differentiation between suspended and settleable solids are often not clear because the terms are sometimes confused in the literature. Until settled to the bottom, all settleable solids are suspended solids and the rate of settling is dependent on quiescence, temperature, density, flocculation, and other factors.

Fish	Oxygen, ppm	Temperature, C	
Bleak	0.68-1.44	16	
Blunt-nosed minnow	2.25	20-26	
Brook trout	2.0 10		
Brook trout	2.2	15	
Brook trout	2.5	20	
Brook trout	1.52	3.5	
Brook trout	2.4	23	
Brook trout	2.5	19-20	
Brook trout	1.35-2.35	15.6	
Brown bullhead	0.3	30	
Brown trout	1.13	6.4	
Brown trout	1.16	9.5-10	
Brown trout	2.13	18	
Brown trout	2.8	24	
Brown trout	1.28-1.6	9.4	
Brown trout	1.64-2.48	17.2	
Brown trout	2.9	_	
Carp	1.1	30	
Carp (mirror)	0.59-2.5	16	
Coho salmon	1.3	16	
Coho salmon	1.4	20	
Coho salmon	2.0	24	
Dace	0.57-1.1	16	
Eel	1.0	17	
Goldfish	0.5	10	
Goldfish	0.6	20	
Goldfish	0.7	30	
Perch	1.1-1.3	16	
Rainbow trout	2.4-3.7	16	
Rainbow trout	2.5	19-20	
Rainbow trout	0.83-1.42	11.1	
Rainbow trout	1.05-2.06	18.5	
Roach	0.67-0.69	16	
Salmon parr	2.0-2.2	8	
Smallmouth bass	0.63-0.98	15-16	
Steel-colored shiner	2.25	20-26	
		20-20	
3-spined stickleback Fench	0.25-0.50	-	
	0.35-0.52	16	
Cellow perch	2.25	20-26	
ellow perch	0.37-0.88	15.5	

## TABLE 8. MINIMUM OXYGEN VALUES AT VARIOUS TEMPERATURES AT<br/>WHICH FISH CAN EXIST UNDER LABORATORY CONDITIONS

(Jones, 1964)

\_

Cairns (1967) described the adverse effects of suspended solids on aquatic biota and acknowledged that the effects would vary with the species and stage of development. A brief summary of this discussion follows:

- (1) Reduction of light penetration This may restrict the growth of photosynthetic forms and, as they are the base of the food web, this could have widespread effects on all other organisms.
- (2) Mechanical or abrasive action This is of particular importance to gill-breathing organisms, such as fish and mussels, because gill impairment not only effects respiration and excretion but may have other widespread metabolic effects.
- (3) Blanketing action or sedimentation This has a deleterious effect on fish spawning sites and in fact may make large areas useless for spawning. Benthic organisms which are a valuable food source for fish may be eradicated.
- (4) Availability as a surface for growth of fungi and bacteria The presence of particulate matter may enable the environment to support substantially increased populations of microorganisms.
- (5) Adsorption and/or absorption of various chemicals This may lead to a buildup of toxic substances in a limited area with a possibility of sudden release.
- (6) Reduction of temperature fluctuations Probably of little importance since particulate concentration would have to be extremely high.

Reduced light penetration will greatly influence productivity. Little plant or benthic productivity can be expected when the turbidity exceeds 200 JTU (National Technical Advisory Committee, 1968). Buck (Tarzwell, 1957) reported the average volume of net plankton in clear ponds was eight times greater than from turbid ponds. Buck also stated (Fry, 1960) that virtually no light is transmitted beyond three inches when suspended solids reach 150 ppm. Most predacious fish feed by sight and in turbid waters have difficulty competing with such bottom-feeder fish as carp, buffalo, and carpsuckers.

Heavier particles of suspended material will settle out and may in this way reduce benthic production. Generally, benthic productivity increases with a change from fine to coarse substrates. Only small amounts of sand and silt shifting in and around the gravel will eliminate much of an area suitable for aquatic insects and other benthic organisms (Aquatic Life Advisory Committee, 1956). Spawning sites for fish are greatly altered by silting, and fish eggs may not receive enough oxygen when covered with fine sediments. A covering of silt may also prevent metabolites from being washed away (Trama and Benoit, 1960).

Reviewing data from other investigations, Tarzwell (1957) stated that in order for suspended solids to be directly harmful to fish the material must be present in very large amounts. Herbert and Merkens (1961) exposed trout to suspensions of kaolin and diatomaceous earth at concentrations of 270 ppm, and substantial numbers of the fish died. Concentrations of 90-100 ppm were less harmful and concentrations of 30 ppm had no observable effect. Wallen (Aquatic Life Advisory Committee, 1956) reported that fish lived for at least short periods (approximately a week) in silt concentrations of 100,000 ppm. The fish died in a few hours when exposed to concentrations of 175,000 to 225,000 ppm.

MacLeod and Smith (1966) found that the rate of metabolism and swimming endurance were reduced in minnows exposed to sublethal concentrations (100-800 ppm) of suspended wood fibers. Herbert and Richards (1963) reported reduced growth in trout kept in pulp suspensions of 50 and 100 ppm for 40 weeks, but concluded that streams containing concentrations of these suspended solids as high as 200 ppm and sometimes higher may support a "reasonable" fish population. They also stated that a fishery is likely to be seriously harmed if the average concentration is greater than 600 ppm.

Herbert, et al (1961) reported a reduction in numbers of trout in a stream polluted with suspended solids (1000 ppm) which was the only polluting material in the stream. He attributed trout reduction to effects on spawning sites, reduction in available food organisms, and some harmful effects directly to the fish.

Smith, et al (1963, 1965, 1966) and Kramer and Smith (1966) have conducted a series of studies on the effects of suspended material from industrial sites. They stated that fish in streams receiving woodfiber wastes may suffer deleterious effects from exposure to sublethal concentrations of suspended fibers. They further concluded that the effects of suspended fibers on fish mortality would depend on the species of fish, type of wood fiber, processing method, DO, concentration, and to a lesser degree, temperature.

When high concentrations of suspended solids are present, death of fish may be due to clogging of the gills (Brown, 1957; Thompson, 1963; and McKee and Wolfe, 1935). Large populations of planktonic organisms such as diatoms and protozoans may produce irritation of fish gills, a condition referred to as sestonosis (Fry, 1960).

There is little information on the effect of turbidity on the toxicity of chemicals. Though the effects of the turbidity are not known, many investigators acknowledge its importance and it is often measured in both laboratory and field studies (see Appendices A and B). Wallen, et al (1957) conducted toxicity studies on a variety of chemicals and carefully measured the turbidity both before and after the tests. They concluded their paper by stating that it would be important to determine if variations in turbidity would significantly affect the toxicity of chemicals, especially those that react to reduce turbidity. Schoenthal (1963) found that mortality in trout exposed to DDT was reduced when turbidity and alkalinity were increased. This may have been due to adsorption of the DDT by the sediment. Brungs and Bailey (1966) have shown that Endrin toxicity to fish is not greatly reduced unless a highly absorptive material such as activated carbon is present.

#### **Other Factors**

Among other water quality factors affecting chemical toxicity in the aquatic environment, water hardness and CO<sub>2</sub> content are probably the most important.

Hardness of water is chiefly attributed to calcium and magnesium ions. Water containing more than 40 ppm total hardness is generally considered hard water while less than this amount indicates soft water. Hardness in natural water can also be correlated with dissolved solids, and sometimes with alkalinity. Increased toxicity of the following chemicals has been reported for hard water: antimony potassium tartrate (Tarzwell and Henderson, 1960), Dipterex (Henderson and Pickering, 1968), and Fermate (Pickering and Henderson, 1966). Soft water increased the toxicity of the following chemicals: Sarin (Pickering and Henderson, 1959), copper and zinc (Sprague and Ramsay, 1965), fifteen metal compounds (Tarzwell and Henderson, 1960). hexavalent chromium (Trama and Benoit, 1960), methyl methacrylate, styrene and vinyl acetate (Pickering and Henderson, 1966), zinc (Mount, 1966, and Cairns and Scheier, 1958), Cumate (Pickering and Henderson, 1966), and copper sulfate (McKee and Wolfe, 1963). Water hardness had little or no effect on the toxicity of the following chemicals: antimony trioxide (Tarzwell and Henderson, 1960), ten organic phosphorus compounds (Henderson and Pickering, 1958, 1959), twelve petrochemicals (Pickering and Henderson, 1966), eight organic cyanides (Henderson, et al, 1961), cyanide (Cairns and Scheier, 1963), and ten phosphorus and chlorinated hydrocarbon pesticides (Pickering and Henderson, 1966).

Dissolved carbon dioxide is important in the aquatic environment, especially to plants. Although a product of respiration, the amount of CO<sub>2</sub> in the body of many animals determines respiration rate. Its primary role in photosynthesis has long been known along with its importance in the carbon-dioxide-bicarbonate system that determines the pH of many natural bodies of water. Carbon dioxide can also affect the toxicity of chemicals in water. At concentrations below 30 ppm, carbon dioxide is generally not toxic to fish. Above this level, it may be limiting in various ways, or lethal at high concentrations depending on the fish species involved. The effect of carbon dioxide on aquatic organisms is closely associated with DO and is mediated largely by ambient water temperature. The significance of carbon dioxide in aquatic environs is discussed fully by Brown, 1957; Doudoroff and Warren, 1962; Fry, 1960; Tarzwell, 1957; and in Water Quality Criteria, 1968. No information was found on carbon dioxide enhancement of the toxicity of chemicals, but when carbon dioxide is present in amounts sufficient to alter pH, this is a distinct possibility.

Natural environmental factors that may affect chemical toxicity directly or indirectly by contributing to water quality changes are:

- (1) Air temperature contributes to water temperature
- (2) Solar irradiation and cloud cover affects surface evaporation rate and water temperature as well as varying incident ultraviolet which may photooxidize chemicals in water
- (3) Precipitation diluting factor
- (4) Wind speed and direction affects atmospheric O<sub>2</sub> uptake of water by surface roiling and also causes varied rates of mixing
- (5) Solids and rock substrata provide dissolved chemicals that primarily constitute the chemical make-up of water
- (6) Plant and animal detritus present in a body of water and from drainage areas provide suspended and dissolved solids and nutrients.

Another important part of the environment that may affect chemical toxicity but not one created by nature, is the extremely wide diversity of water pollutants added to natural waters by man. Synergistic or antagonistic effects can and do occur in dilute chemical concentrations. Mixed pollutants are discussed briefly in the section <u>Industrial Wastes</u>.

#### SECTION X

#### INDUSTRIAL WASTES

The problem of maintaining desirable water quality increases with advancing technological development. One of the most serious water quality problems facing industry with respect to effluent discharges is the effect of toxic wastes on aquatic life. The many substances carried in solution and suspension determine whether water will be suitable for supporting aquatic organisms. Chemical contents of some wastes may be freely soluble or miscible in water, such as acids, alkalies, organic solvents, etc.; or nonsoluble, such as slurries from mining operations, soil washings, or wood pulp fibers. Adverse effects may be direct and immediate or they may be chronic and deleteriously affect the environment only gradually over a long period of time. Mixed, the wastes may be synergistic or they may reduce the damaging effects each would have individually (Garrett, 1957; Keup, et al, 1967; and Neel, 1963).

Complex wastes such as pulp mill effluents, wastes from oil refineries, and chemical plants are neither constant in content nor in concentration and this further complicates tests to determine their toxicities. Not only will a waste vary in toxicological and chemical characteristics from day to day, but also within any given day variations will occur due to process changes, raw materials, and end products. These wastes contain many known but often many unknown toxic substances (Clemens and Clough, 1965; Keup, et al, 1967; and National Technical Advisory Committee, 1968). Ellis in 1937 summarized the hazards of 30 common types of municipal and industrial effluents. This list was republished 30 years later by Keup, et al (1967) as shown in Table 9. No updating of this data summary or anything similar to it was found. For these reasons, less emphasis was placed in the present study on acquiring mixed effluent data. However, during the course of literature acquisition, considerable information on this subject area was obtained. These are briefly abstracted in Table 10. Although merely a token selection of papers on this subject, the abstracts serve to show the wide diversity of problems associated with industrial waste effluents.

For research to be effective, the scientist must know the materials he works with. McKee and Wolfe (1963) in their summaries of potential chemical pollutants discuss 39 chemicals as originating from textile wastes, while another (Anon., 1966) listed 386 compounds. This type of situation probably exists for most other industries. In all likelihood, even the latter listing is not complete since some process changes have undoubtedly been made since 1966. One of the first orders of business should be the establishment of listing of effluent components from industrial plants. These listings should be continually updated.

١

			Changes in Water Aff	ecting Fish				
			Concentration	Increase in				
Turner of Montes	Decrease in	Increase in	Increase in	Specific	Increase in	Increase in	Bottom Pollution	Specific Toxic
Types of Wastes	Dissolved Oxygen	Acidity	Alkalinity	Conductance	Tutbidity	Ammonia	Blanket	Action on Fishes
			Mineral Wastes,	Little Bacterial	Action			
Frosion silt	None	None	None	None	Critical	None	Critical	None
Limestone sawmills			Possible	Moderate				
Ashestos works								Possible
Mine flotation	Possible	Possible						Possible to critica
Coal- and iron-mine drains		Critical	None		None		Possible	Possible
Crude oil		None		None			Possible to critical	Possible to critica
Salt water from oil wells	None		Possible	Critical			None	
			Organic,	Bacterial Action	<u>n</u>			
Municipal sewage	Critical	Possible	Possible	Possíble	Possible	Critical	Possible to critical	Possible to critica
Dairy industries		Critical	None	Moderate	Moderate	Moderate		Possible
Packing plants		Moderate				Critical	Critical	
Canning factories		Critical	Possible				Possible to critical	
Breweries and distilleries		None to moderate	None to moderate		Possible	Possible		
Beet sugar, pulp wastes		Critical	None					Possible to critica
Paper pulp	Possible to critical	Possible	Possible	Possible			Critical	Possible
Sawdust								
			Chemi	cal Processes				
Cool-gas wastes	Possible	Possible	Possible	Moderate	None	Critical	Critical	Critical
Spent lubricants			None	Possible		None		Possible to critica
Metal refineries	None		Possible		Possible	Possible	Possible to critical	Critical
Laundries and wool washings	Moderate	None	Moderate to critical	Moderate	Moderate	Moderate to critical	Possible	Possible
Steffens house waste			Critical	Critical		None		Critical
Sulphite <b>pul</b> p	Moderate to critical	Possible	Moderate to critical	Moderate		Possible	Possible to critical	
Strawbound waste		None	Critical		1			
Themical works (1)	None				Possible		None	
hemical works (2)	Possible	Critical	None		None	None		
Fanneries	Moderate	Possible to critical	Possible to critical		Possible	Possible to critical	Critical	
Dye works	Possible	None to moderate	None to moderate		None	None	Possible	
Bittem liquors	None	Critical	None	Critical			None	Possible to critic
in-plate and wire mills	None to possible			Moderate	None to possible		Possible to critical	
tarch factories	Possible to critical				Possible	Possible		
		Possible to critical		Possible	Moderate			Possible

(a) Increases in both acidity and alkalinity are noted in some cases, due to the fact that two or more kinds of effluents are mixed, with one predominating at times, and to changes which take place in the stream after the effluent is added.

#### TABLE 10. GENERAL COMMENTS ON SELECTED INDUSTRIAL EFFLUENTS

Type of Waste	Remarks		Reference
eneral			<b>TT</b> . 1.1.1.1
Industrial wastes	A discussion of methods for studying to:	kicities of industrial wastes.	Heukelekia (1948)
Organic wastes	Bottom communities found in streams show characteristics reac-		Hirsch
	tions to pollution, i.e., grossly polluted streams contain tubificid and chironomids, etc. Various streams in New Zealand were surveyed.		(1958)
Unspecified chemical waste	A complex chemical waste containing such toxicants as fluorides, arsenic, copper, zinc, tin, lead, and SO <sub>2</sub> was shown to lower pH and cause fish kill at a loading of about 0.5% of the waste in sea- water at pH 5.5 and lower. Maximum toxicity occurred when superphosphate was being produced.		Chanin and Dempster (1958)
Industrial wastes	Fifty percent reduction in photosynthesis in kelp resulted from exposures to the following chemicals in four days: Inorganic		Clendenning and North (1960)
	Mercury	0.05 ppm	(1700)
	Copper	0.1 ppm	
	Nickel	2.0 ppm	
	Chromium	5.0 ppm	
	Chlorine	5.10 ppm	
	Zinc	10.0 ppm	
	Organic		
	Sodium pentachlorophenate	0.3 ppm	
	Zephiran chloride	1.0 ppm	
	Sodium dodecyl sulfate	5-10 ppm	
	Cresols	5-10 ppm	
	Phenol	10.0 ppm	
	Emulsified fuel oils	10-100 ppm	
Organic wastes	Evaluation was made of the various approaches to the problems of organic pollution in tidal estuaries.		Pyatt (1964)
Industrial wastes	A summary of the ways in which industrial wastes may affect aquatic life.		Neel (1963)
Organic wastes from industrial sites	Stream had DO depletion for about a 45-mile stretch with heavy loss of fish and plankton organisms.		George, et a (1966)
Industrial wastes	Methods of studying industrial wastes are described.		Jackson and Brungs (1966)
Industrial wastes	An attempt is made to estimate future industrial discharges into the Eems Estuary, The Netherlands.		Eggink (1967)
Various polluting agents in rivers	A summary of problems arising from suspended solids, toxic materials and nutrients from sewage pollution.		Patrick (1968)

Type of Waste	Remarks	Reference
etroleum		Turnbull, et al
Refinery wastes from:	Effects on bluegill, 24-hr TL <sub>m</sub> , % vol were:	(1954)
Fractionation area	Nontoxic:	(1)5()
Cracking area	31.0	
Lube oil treating area	Nontoxic:	
Paraffin treating area	37.0	
Acid plant area	3.1	
Naphtha treating area	75.0	
Fluid catalyst unit	3.1	
Sulfuric acid alkylation	0.4	
unit	29.0	
Combination unit Distillate tank drawoff	12.0	
Oil field brine water	Average number of aquatic species found in a stream with varying chloride concentrations was: 4 - 13,000-20,000 ppm 6 - 10,000-13,000 ppm 7 - 8,000-10,000 ppm 8 - 4,000- 8,000 ppm 10 - 1,000- 4,000 ppm 13 - 1,000 ppm	Clemens and Finnell (1957)
Oil field brine water	The 24-hr $TL_m$ of fish at various concentrations of chlorides showed a marked reduction in deaths as the concentration neared 7,000 ppm. One test at 7,000 ppm for 192 hr showed 90% survival.	Wood (1957)
Oil field brine water	<i>Fundulus</i> and <i>Lagodon</i> may survive salinities up to 2.7%. <i>Leistomus</i> did well above 2.0%.	Cole, et al (1958)
Petroleum products: Gasoline Diesel fuel oil Bunker oil	Effects on American shad, 48-hr TL <sub>m</sub> (mg/l), were: 91 167 2417 Lethality increase was accompanied by low DO.	Tagatz (1961)
Refinery effluent	Based on 24-hr TL <sub>m</sub> , <i>Lebistes reticulatus</i> was most resistant fish of several tested.	Bunting and Irwing (1965)
Refinery effluent (hydrogen sulfide and phenolics)	No correlation between sulfide concentration and lethal dosage to fish was found. For phenolics, the LD50 for goldfish was 33.1%, LD50 for red shiners was 18.8%, and LD50 for <i>Daphnia</i> was 19.0% lower than that for red shiners.	Clemens and Clough (1965)
Petroleum oil	Pollution resulted from an underground storage tank leak. At the beginning, the concentration in the water was 221.3 ppm and after one year, 1.4 ppm. Toxic effect was pronounced on micro-fauna in sediments.	McCauley (1966)

Note: Further references on this general subject area includes papers by Copeland and Dorris (1964), Douglas, et al (1960, 1962, 1963), Gould and Irwin (1965), Johnson (1968), Smith (1968), Tubb and Dorris (1965), Ward and Irwin (1961), and Zobell (1964).

Type of Waste	Remarks	Reference
Pulp and Paper		
Sulfite waste liquor	Decrease in feeding rate in oysters was observed.	Galtsoff, et al (1947)
Sulfite waste	Marked avoidance by juvenile chinook salmon was observed with little or no avoidance by juvenile coho salmon.	Jones, et al (1956)
Kraft mill effluent	A 100% survival of young salmon was recorded in seawater with effluent concentration under 4.8% with adequate oxygen.	Alderdice and Brett (1957)
Sulphate waste liquor	Reduced DO in river water to 1.0 mg per liter was recorded. Prawns and Apocryptes lanceolatus died in 3 minutes or less when exposed to the waste liquor.	Chowdhury (1957)
Kraft mill effluent	Live car bioassays showed wastes were lethal to game fish during periods of high water temperature. In mid-July, pollution-sensitive bottom fauna decreased from 54 to 17%.	Spindler and Whitney (1960)
<b>P</b> aper mill effluent (chlorine)	A 13-hr TL <sub>m</sub> of 32% concentration of the effluent was obtained for <i>Salmo salar</i> .	Betts and Wilson (1966)
Paper mill effluent	Silver salmon did not avoid sulfite liquor or kraft wastes in low enough concentrations to be "safe". Toxicity data are too numerous to summarize here.	Holland, et al (1960)
Sulphite waste liquor	In fluviarum experiments, avoidance reactions were exhibited by <i>Phoxinus phoxinus, Leuciscus rutilus, L. idbarus, Perca</i> <i>fluviatilis, Coregonus nasus, Salmo salar,</i> and <i>Gasterosteus</i> <i>aculeatus.</i>	Hoeglund (1961)
Kraft mill effluent	A significant decrease in <i>Sphaerotilus natans</i> growth was accom- plished by the intermittent discharge of the waste using a five- or six-day holding period with a one- or two-day release.	McKeown (1962)
Kraft mill effluent	Induced spawning in mussels Mytilus edulis and M. californianus was observed.	Breese, et al (1963)
Paper mill effluent	<ul> <li>Maximum survival of walleye eggs above mill: On bottom 1.2%; off bottom 49.1%</li> <li>Maximum survival of walleye eggs below mill: On bottom 1.2%; off bottom 3.5%.</li> <li>The principal cause of mortality below the mill was Sphaerotilus natans.</li> </ul>	
Sulfite waste	Regeneration studies of bisected planaria indicated: At 550 ppm – no regeneration occurred At 50 ppm – regeneration was 75% of control.	Eng. Science, Inc. (1964)
Pulp mill waste	Histological examination of three species of fish showed decrease in RNA, glycogen in liver, necrosis in kidney, and accelerated secretion of mucus in gills. In bivalve livers, decrease in RNA and glycogen occurred, and nuclei disappeared in kidney cells.	Fujiya (1965)

Type of Waste	Remarks	Reference
Pulp and Paper (Continued) Sulfite wastes	It was not clearly demonstrated that sulfite waste in the area studied was the only cause of deaths of oysters, but it was con- cluded that the amounts were sufficient to cause stresses which may have long-term adverse effects.	Woelke (1965)
Neutralized kraft process effluents: Brown stock screen- ing and deckering Recausticizing	Effects on guppies were: 96-hr TL <sub>m</sub> , % vol of effluent – 51.3 92.5	Howard and Walden (1965)
Bleach plant acid sewer Bleach plant caustic sewer Neutralized whole effluent Unneutralized whole	29.5 41.1 52.5 9.2	
effluent Neutralized kraft pulp bleach waste	Reduced growth in sockeye and pink salmon alevins was found in concentration of $1/10$ to $1/20$ the average 96-hr TL <sub>m</sub> .	Servizi, et al (1966)
Kraft effluent	A 75% concentration was required to kill 100% of Salmo salar in less than 10 hr.	Betts, et al (1967)
<b>Sewage</b> Sewage	This is a summary of the problems of toxic materials and nutrients from sewage pollution.	Lackey (1958)
Sewage	A 10% concentration caused reduction on photosynthetic capacity of kelp. A concentration of 1% gave no such indication.	Clendenning and North (1960)
Sewage	Flagellates, protozoa, diatoms, and filamentous green algae showed highest sensitivity to pollution while rotifers, <i>Sarcodina</i> , and Volvocales were most tolerant.	Farmer (1960)
Sewage	A resume of sewage pollution of streams and beaches on Oahu.	Lam (1964)
Sewage	Low surface productivity at point of discharge was observed. Increase in productivity downstream in about 6 hr was recorded with maximum values in about 10 hr. This was followed by a decrease toward normal levels.	Calif. State Water Quality Control Board (1965)
Sewage	In samples of surface water from marine stations, the numbers of <i>Escherichia coli</i> depended primarily on the amount of sewage and direction of flow. Results varied enormously.	Bonde (1967)

Type of Waste	Remarks	Reference
Suspended Solids Suspended mineral solids	Concentrations of 90 to 810 ppm made trout more susceptible to other adverse factors in the environment.	Herbert and Merkens (1961)
China-clay suspended waste	Concentrations of 1000 ppm reduced abundance of brown trout in an otherwise unpolluted stream. Suspensions of 60 ppm had no observable adverse effects.	Herbert, et al (1961)
Suspended solids	Laboratory experiments did not indicate that suspensions of 30 ppm kaolin and diatomaceous earth and suspensions of 50 ppm wood fiber and coal-washery wastes make well-grown trout more susceptible to disease.	Herbert and Richards (1963)
Pulpwood fibers	Significant changes occurred in blood of fathead minnows exposed to wood fibers. Increased hematocrit was highest for conifer groundwood, followed by aspen groundwood, kraft conifer, and sulfite conifer.	Smith, et al (1965)
Suspended conifer groundwood	Survival of walleye fingerlings decreased when DO was reduced.	Smith and Kramer (1965)
Suspended groundwood	Rainbow and brown trout eggs survived in suspensions of 60, 125, and 200 ppm conifer groundwood. Trout alevins survival rate decreased to a minimum of 0 in 250 ppm. The growth rate of survivors was reduced.	Smith and Kramer (1965)
	Fathead minnows which were held for 96 hr in 0 to 2000 ppm of aspen groundwood showed no effects to this exposure. A similar series run in conifer groundwood showed increased mortality at 738 and 2000 ppm.	
Conifer ground- wood fiber	Reduced growth was recorded for walleye fingerlings held in con- centrations of 50 to 150 ppm.	Smith, et al (1966)
Suspended wood fibers	Walleye eggs survived at concentrations of 250 ppm.	Kramer and Smith (1966)
Paper fiber sludge	Low DO, high CO <sub>2</sub> , and presence of dissolved sulfides in streams were recorded.	Colby, et al (1967)
Miscellaneous Unspecified chem- ical waste	A complex chemical waste containing such toxicants as fluorides, arsenic, copper, zinc, tin, lead, and SO <sub>2</sub> was shown to lower pH and cause fish kill at a loading of about 0.5% of the waste in seawater at pH 5.5 and lower. Maximum toxicity occurred when superphosphate was being produced.	Chanin and Dempster (1958)
Electroplating wastes	A midgefly, <i>Cricotopus bicinctus</i> , survived and matured in con- centrations of chromium as great as 25 ppm, in copper at 2.2 ppm, and in cyanides at 3.2 ppm.	Surber (1959)

Type of Waste Remarks		Reference
<b>liscellaneous (Continued)</b> Spent still liquors from coal distillation		
Smelter wastes	Near the smelter, the aquatic flora and productivity was greatly reduced. <i>Leptodictyum riparium</i> and <i>Eleocharis acicularis</i> v. <i>submersa</i> appeared to be the most tolerant organisms.	Gorham and Gordon (1963)
Acid mine drainage	Twenty states have streams affected by acid mine drainage. Pennsylvania has 2,906 miles of streams polluted with acid mine drainage, Virginia has 1,150, and Kentucky has 590. The remain- ing states have less than 300 each.	Kinney (1964)
Alkaline water	The pH of water passing through asbestos-cement pipeline was increased to 9.5 with no immediate lethal effect on salmonids.	Sprague (1964)
Lurgi process wastes (bituminous coal)	Treatment of effluent reduced permanganate value to less than 50 ppm and BOD to less than 25 ppm. The residual organic matter had little direct toxic effect on fish.	Cooke and Graham (1965)
Uranium mill wastes	wastes The radioactive element in this study was radium; the nonradio- active materials included sulfates, nitrates, chlorides, manganese, iron, lead, arsenic, and various organics. These wastes were im- portant in limiting aquatic biota below uranium mills. Changes in composition of the wastes and water flow make it difficult to calculate the radioactive and nonradioactive components of the mill wastes.	
Coal washer wastes	As long as the coal washer wastes were intermittent, there was little effect on biological productivity.	Charles (1966)
Uranium mine	The effluent did not appear to have any adverse effect on plankton, periphyton, benthos, and fish species other than trout (reduced numbers).	Mitchum and Moore (1967)
Landfill pollution	Groundwater was polluted with CO <sub>2</sub> from decomposing refuse in a landfill.	Bishop, et al (1967)
Sulfuric acid water	furic acid water Considerable reduction in survival percentage was found in herring eggs and embryos at dilutions of 1:32,000.	
Photographic wastes	Common chemicals found in these wastes are potassium ferri- cyanide, sodium ferricyanide, boron, chromium, and sodium thiosulfate. Release of this type of waste into streams and the Los Angeles sewage system is discussed.	Hennessey ar Rosenberg (1968)

#### SECTION XI

#### EXTRACTED DATA – THE EFFECT OF CHEMICALS ON AQUATIC BIOTA

Extracted information from originally published data are divided in two sections, both alphabetically arranged by chemical name. One section (Appendix A) concerns listing by chemical name, and the other a similar listing by commercial designation (Appendix B). In all cases, the chemical names and names (common or scientific) of organisms designated by the authors were used in this compilation. None of the nomenclature was changed or corrected in any manner, e.g., when authors used the common name of a fish, this and this alone was used. The abbreviations and other designations are discussed later in this report section and described in footnotes to the Appendices. In using the data compilations, care should be exercised in searching varied alternative names for a given compound.

Since many papers contained large amounts of data, the most significant toxicity level was chosen for inclusion in this compilation. In most cases, data presented at 96-hr TL<sub>m</sub> (designated T4:  $T = TL_m$  or TL<sub>50</sub>, and 4 = four days or 96 hours) were selected when available. With few exceptions, the T value at 4 days was lower than the values for 1 or 2 days. The T4 value is generally accepted as a realistic indication of toxic effect and the best one to use (lacking data from chronic studies) in estimating safe levels for effluent release. T1 or T2 data were usually not included unless these were the only data given. A and C following these designations indicate acute or chronic bioassays, respectively. Since the data are presented as brief summaries, the reader is referred to the original report for additional information. When EC<sub>50</sub>, LC<sub>50</sub>, and LD<sub>50</sub>\*, were known or described as being concerned with lethal effects, these abbreviations were judged to be essentially the same as TL<sub>m</sub> or TL<sub>50</sub> and designated as such (T) in the data extracts for consistency. We acknowledge that this is not standard practice, and that there are important differences in these designations.

The conditions noted by the researchers are designated by lower case letters. When the conditions were controlled, these letters were underlined. In some cases, the authors briefly referred to previous papers as a simple means for describing experimental conditions. No underlines were made in these instances, although in all likelihood some conditions were controlled.

Comments, in general, are brief, with the expectation that interested readers would consult the original article for further information.

Since the chemical nature of most industrial effluents is very complex and seldom analyzed or reported, there is little information on the effect of mixed effluents or mixtures of chemicals in the data presented. For this reason, this document must be described merely as pertaining to the effect of single chemicals or simple mixtures of chemicals on aquatic life.

There was no attempt to extract data from various reviews available, since these rarely contained descriptive information concerning experimental conditions. Among others, the reader is referred to:

<sup>\*</sup> $EC_{50}$  = median effective concentration,  $LC_{50}$  = median lethal concentration, and  $LD_{50}$  = median lethal dosage.

American Public Health Assoc. (1960) Anon. (1968) Aquatic Life Advisory Committee (1955, 1960, 1967) Averett and Brinck (1960) Beak (1958) Bick (1963) Breidenback, et al (1967) Breidenback and Lichtenberg (1963) Brown (1961) Burdick (1965) Butcher (1959) Butler (1966) Buzzell, et al (1968) Byrd (1960) Carter (1962) Cope (1963, 1965) Cope and Springer (1958) Cottam (1961) Delaporte (1958) Dewey (1958) Doudoroff (1951) Doudoroff and Katz (1950, 1953) Faust and Aly (1964) Ferguson (1967) Ferguson, et al (1966) Fromm (1965) Fruh, et al (1966) Ganelin, et al (1964) George (1959) Graham (1960) Hawkes (1963) Henderson and Tarzwell (1957) Hirsch (1958) Hoffman (1960) Holden (1964, 1965) Hughes and Davis (1967) Hunt (1965) Hynes (1966)

Ingram and Towne (1960) Jackson (1966) Johnson (1968) Johnson, et al (1967) Jones (1964) Kerswill, et al (1960) Keup, et al (1966, 1967) King (1968) Langer (1964) Lawrence (1962) Llovd (1964, 1965) MacMullen (1968) Mackenthum and Ingram (1962, 1964) Malina (1964) McKee and Wolfe (1963) McFarland (1959) Moore (1967) National Technical Advisory Committee (1968)Neel (1963) Newsom (1967) Nicholson (1959, 1967) Nicholson, et al (1964) Patrick (1968) Powers (1918) Reymonds (1962) Rudolphs, et al (1950) Ryckman, et al (1966) Schoettger (1967) Skidmore (1964) Snow (1958) Spiller (1961) Sproul and Ryckman (1963) Surber and Taft (1965) Tarzwell (1959, 1962) Water Pollution Control Federation Research Committee (1958-1968) Weaver, et al (1965) Webb (1961) Wilson (1968)

Doudoroff (1951) states that certain references with literature summaries are particularly helpful in providing pertinent information published before 1954 on water pollutants toxic to fish. These references are:

Redeke, H. C., "Report on the Pollution of Rivers and Its Relation to Fisheries", Rapp. Conseil Permanent Intern. Exploration Mer, 43, 1 (1927).

Steinmann, P., "Toxikologie der Fische", Handbuch Binnenfischerei Mitteleuropas (Germany), 6, 289 (1928).

- Helfer, H., "Giftwirkungen auf Fishe; ihre Ermittelung der Versuche und die Bewertung der Ergebnisse", Kleine Mitt. Mitglied. Ver Wasser-Boden-u. Lufthyg., 12, 32 (1936).
- Cole, A. E., "The Effects of Pollutional Wastes on Fish Life", in a Symposium on Hydrobiology, University of Wisconsin Press, Madison, Wisconsin, 241 (1941).
- Southgate, B. A., "Treatment and Disposal of Industrial Waste Water", Department of Scientific and Industrial Research, London, England, 23 (1948).
- Harnisch, O., "Hydrophysiologie der Tiere", in "Die Binnengewasser", Vol. 19, Ed. A. Thienemann, Schweizerbart'sche, Erwin Nagele, Stuttgart, Germany (1951).
- "Water Quality Criteria", California Water Pollution Control Board, Pub. No. 3, Sacramento, California (1952). (Also, Addendum No. 1, 1954, and Pub. No. 3, 1963).

Not to demean past contributions from ecological investigators, but rather to suggest how the data they develop in the future can be made more valuable for engineering application, it may be stated that problems of interpretation encountered in this review would be minimized or eliminated by the following:

- Positive identity of chemicals under test
- Precise description of test organisms
- Use of standard test methods, where applicable, or full details of procedure if standard methods are not used
- Closer definition and control of test conditions.

Apparent differences in results among investigators of the same chemical on the same fish species may have resulted from different methods of handling specimens prior to and during tests, different stages in the life cycle of specimens, variations in physical and chemical properties of the water, excursions in time-temperature pattern of exposures to the chemical, and different methods of evaluating effects.

We believe the manner in which this report is compiled will serve the industrial community and others as well. Since each reader will undoubtedly have a specific applied situation for using the data, there was no attempt to summarize in narrative form the data for each compound. The compilation gives pertinent data for each chemical for which information was found, tempered by the comments on bioassay or field conditions, as well as providing a bibliography of the more recent information available in the literature through 1968. Additionally, a Species Index is presented in Appendix C and the chemical nature of commercial chemicals is given when available in Appendix D.

In handling large numbers of references, an occasional document may be overlooked and not included. The authors would sincerely appreciate being informed by the readers of such omissions for the principal time period covered (1958-1968). An updating effort of this report is now under consideration and will likely be completed by early 1972.

# SECTION XII

# SUMMARY AND CONCLUSIONS

Fish, representing one of the highest trophic levels in the aquatic environment, are the animals of choice in studying the toxicity of chemical effluents in natural waters. Their importance is further emphasized since man may be the next highest trophic level where edible fish are concerned. Furthermore, considering fish as indicator organisms, their presence probably indicates that the water in which they survive is suitable for consumption or other uses by man, except in some situations, for example, where a cumulatively toxic material is present in small amounts and the fish develop resistance to that material.

With the magnitude of pollution problems today, standard fish bioassay procedures (particularly, flow-through) are adequate for the task at hand. This is especially true for evaluation of chemicals that are acutely or immediately toxic although these procedures can also be used in studying the chronic toxicity of chemicals at sublethal levels. These standard procedures must be employed in conjunction with other evaluations, especially specific residue analyses, when a chemical or ion causes a drastic problem such as a large-scale fish kill. The chronic continuous flow exposure of fish is preferable for determining more precisely acceptable concentrations for chemical release.  $TL_m$  data should be a baseline for comparison of data from either type of evaluation. Adequate reporting of data and experimental conditions, especially water quality data, would greatly enhance the value of published information.

For field investigation of chemical toxicity in the aquatic environment, the *in situ* bioassay is desirable. Exposure of native fish or highly sensitive fish from other sources would give a better representation of the toxicity of a given chemical in a given situation. This should be supplemented with chemical analysis of the effluent in question as well as a recording of receiving water quality data. *In situ* evaluation of water from above and immediately below an effluent addition could provide an elegant proof of lack of complicity in a fish kill by a manufacturer.

With the present situation of gross pollution in many localities, study of fish responses other than lethality are of little direct utility except in cases where a chemical has long-term, sublethal effects, such as DDT and other chlorinated hydrocarbons. All such procedures would be best employed in conjunction with standard bioassays so that appropriate comparisons can be made. These procedures include:

- (1) Observations of abnormal behavior
- (2) Autopsy and histology
- (3) Avoidance
- (4) Growth retardation
- (5) Radiotracers
- (6) Effects on various life stages
- (7) Spawning
- (8) Swimming or cruising speed and oxygen consumption
- (9) Blood studies
- (10) Glucose transport
- (11) Environment stress
- (12) Thermal acclimitization
- (13) Fish taste
- (14) Conditioned avoidance response.

For a careful limnological approach in bioassay studies, several researchers have suggested toxicity evaluations of aquatic organisms representing at least three trophic levels of the food web. Fish would, of course, be one level. Another could be bioassay using *D. magna* and the techniques described by Anderson (1944-1946, 1948, 1960). The third type of bioassay could be with algae, using the technique of Palmer and Maloney (1955) or of Fitzgerald and Faust (1963). BOD determination by the standard method (American Public Health Association, 1967) could be another bioassay procedure. More rapid, alternative methods (e.g., STOD) are also available for estimating BOD. BOD data alone can provide a useful index of toxicity or of oxygen depletion in receiving water.

Marine bioassay utilizing various organisms primarily including fish, oyster, clams, and shrimp in a flow-through type of system lags considerably behind reports of freshwater bioassays in the amounts of data reported. The procedure is practical but could be improved upon by maintenance of water temperature, DO, and other water factors. The sensitivity of shell regrowth in bioassay and field studies of oyster (*Crassostrea virginia*), clam (*Mercenaria mercenaria*), and related marine mollusks to low concentrations of pesticides suggests that a bioassay using a freshwater mollusc should be developed.

Reports on field studies of pollution problems include some of the classic examples of disruption of the aquatic environment by polluting effluents and pesticide applications. Although the results of such research are irrefutable in most instances, improvement is needed in recording and reporting correlative data, e.g., water quality, weather, and other environmental factors. Collecting devices are generally adequate for their designed purposes if used by experienced field scientists, but some mechanical changes could improve collection and ease of manipulation in the field.

Evaluation in the field in a given pollution situation can yield more realistic results than evaluation by laboratory bioassay. Consider, for example, change in chemical toxicity due to seasonal temperature change. This is the reason *in situ* bioassay (using live cars or wire cages and plastic pools or raceways with suitable bioassay species in conjunction with automatic water quality monitoring) appears to be the method of choice for an individual industry to evaluate the effect of its particular effluent(s) on a given waterway.

The complex, highly interrelated factors in the aquatic environment may have profound effect on the toxicity of a chemical. Of these, the most important are temperature, dissolved oxygen, pH, turbidity (suspended solids), and water hardness. Their importance in aquatic studies and their effect on chemical toxicity were discussed in some detail.

In addition to conclusions and comments made throughout this report, the following remarks are made in direct response to the objectives outlined earlier in this report:

(1) Collect and summarize in standardized format the available information from the scientific literature. The extracted data presented in Appendices A and B show that there is a considerable lack of adequate reporting of experimental conditions concerning the effect of chemicals on aquatic life. The complexity of factors in both laboratory and field studies in aquatic biology is such that control or description of them is most difficult. The specific effects of chemicals on individual species of aquatic biota are voluminously shown in Appendices A and B in a standardized format. A Species Index (Appendix C) facilitates assembling all data for any given species. Procedural details and environmental factors important in the observation or measurement of these effects are discussed in appropriate sections of this report. Except for standard fish bioassays (static,

continuous flow, and chronic exposures) and BOD, the wide variety of procedures utilized for these studies were not discussed in detail. References are cited to allow the individual reader to obtain these procedures when needed.

- (2) Review the existing information on aquatic life as it is applicable or related to the study of water pollution. The existing, more recent information on aquatic life as it is applicable or related to the study of water pollution was reviewed. Discussion of test species, lack of species variety identification, short-comings of procedural details in reporting bioassay and field results, etc., is presented in various report sections.
- (3) Review the methodology used in studying the effects of chemicals on aquatic life. Similarly, a review of the more important aspects of aquatic life methodology is presented. Briefly, except for the standardized bioassays, experimental procedures vary almost directly and specifically with the number of researchers reporting data in the literature.

We believe the requirements described in the objectives for this study were fulfilled.

### SECTION XIII

# RECOMMENDATIONS

We recommend:

(1) Establishment of a chemical pollution effect information-analysis center as a means of continuously updating the information summarized here. This report has shown the large volume of information available on the effects of chemicals on aquatic life. The amount of information is unwieldy and difficult to work with. A computerized information-analysis center would be capable of quickly identifying all pertinent data and would allow rapid preparation of reports summarizing data on any chemical or group of chemicals in given situations for various aquatic biota. Establishment of a prototype information center on analytical methodology related to the aquatic environment is now in progress at Battelle's Columbus Laboratories. Bioassay data not now published but held by individual manufacturers could be anonymously submitted for inclusion into the information pool. Only data obtained by a standard procedure or a well-described one would be greatly expanded in this manner. The information content of this prototype center is to be continually updated so that it would always be current as well as immediately responsive as required.

As data are accumulated, the chances for predicting potential problems by mathematical modeling and simulation of the effect of chemicals on aquatic life will be improved. This report should provide a sound base for pursuing this approach.

(2) Preparation of listings of chemical constituents present in effluents by cooperative input from the chemical industry. Data inputs could be submitted anonymously. The listings should be continuously updated and made easily available to anyone who requests updated copies.

(3) Development of a standard pattern of laboratory evaluations, not limited to but primarily based on fish bioassay, for estimating more accurately the effect of chemicals on aquatic life. Data from such evaluations could then be used in mathematical modeling studies which would be used for predicting chemical toxicity under widely varied environmental conditions.

(4) Development of *in situ* field bioassay procedures for more realistic results than those obtained from laboratory bioassays.

We suggest that researchers publishing in this field be encouraged to positively identify the chemicals evaluated; to precisely describe test organisms; to use standard methods, if possible, or to fully describe experimental procedures; and to more closely define and control experimental conditions. This improved reporting would greatly enhance the utility of the data, and allow more precise development of multivariate analyses and mathematical modeling for predictive assessments of chemical pollution problems.

#### SECTION XIV

### BIBLIOGRAPHY

Abedi, Z. H. and D. E. Turton. (1968). Note on the response of zebra fish larvae to Folpet and Difolatan. J. Assoc. Ohio Anal. Chem. 51(5): 1108-1109.

Abram, F.S.H. (1964). An application of harmonics to fish toxicology. Intern. J. Air Water Pollut. 8: 325-338.

Abram, F.S.H. (1967). The definition and measurement of fish toxicity thresholds. In: Advances in Water Pollution Research, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 1. Washington, D.C., Water Pollut. Contr. Fed., pp. 75-95.

Alabaster, J. S. (1956). The toxicity of certain weed killers to trout. Proc. 3rd Brit. Weed Contr. Conf. 2: 807-808.

Alabaster, J. S. (1967). The survival of salmon (Salmo salar L.) and sea trout (S. trutta L.) in fresh and saline water at high temperatures. Water Res. 1(10): 717-730.

Alabaster, J. S. and F.S.H. Abram. (1965). Development and use of a direct method of evaluating toxicity to fish. In: Advances in Water Pollution Research, Proc. 2d International Conference held in Tokyo, August 1964. Vol. 1. N.Y., Pergamon Press, pp. 41-60.

Alabaster, J. S. and D. M. Herbert. (1954). Influence of carbon dioxide on the toxicity of ammonia. Nature 174: 404.

Alabaster, J. S. and A. Swain. (1963). Heater water and fish. Ann. Rept. Challenger Soc. 3(15): 20.

Alabaster, J. S. and R. L. Welcomme, (1962). Effect of concentration of dissolved oxygen on survival of trout and loach in lethal temperatures. Nature 194: 107.

Alderdice, D. F. (1963). Some effects of simultaneous variation in salinity, temperature and dissolved oxygen on the resistance of young coho salmon to a toxic substance. J. Fish. Res. Bd. Can. 20: 525-550.

Alderdice, D. F and J. R. Brett. (1957). Some effects of Kraft mill effluent on young Pacific salmon. J. Fish. Res. Bd. Can. 14: 783-795.

Alderdice, D. F and M. E. Worthington. (1959). Toxicity of a DDT forest spray to young salmon. Can. Fish Cult. 24: 41-48.

Allison, D., B. J. Kallman, O. B. Cope, and C. V. Valin. (1964). Some chronic effects of DDT on cutthroat trout. U.S. Fish and Wildl. Serv. Res. Rept. No. 64, 30 p.

American Public Health Association, American Water Works Association and Water Pollution Control Federation. (1971). Bioassays to evaluate toxicity to fish. To be published in: Standard methods for the examination of water and wastewater. 13th ed., 23 p. American Public Health Association, American Water Works Association, and Water Pollution Control Federation. (1967). Standard methods for the examination of water and wastewater; including bottom sediments and sludges. 12th ed. N.Y., American Public Health Association, Inc., 769 p.

American Public Health Association. Water Supply Committee. (1960). What we do and do not know about chemical pollutants in water. Public works 91(8): 194, 196, 198-200.

American Society of Civil Engineers. Committee on Thermal Pollution, Sanitary Engineering Division. (1967). Bibliography on thermal pollution. J. San. Eng. Div., Proc. Amer. Soc. Civil Eng. 93(SA3): 85-113.

Ames, P. L. (1966). DDT residues in the eggs of the osprey in the north-eastern United States and their relation to nesting success. J. Appl. Ecol. 3(Suppl.): 87-97.

Anderson, B. G. (1944). The toxicity thresholds of various substances found in industrial wastes as determined by the use of *Daphnia magna*. Sewage Works J. 16: 1156-1165.

Anderson, B. G. (1945). The toxicity of DDT to daphnia. Science 102: 539.

Anderson, B. G. (1946). The toxicity thresholds of various sodium salts determined by the use of *Daphnia magna*. Sewage Works J. 18: 82-87.

Anderson, B. G. (1948). The apparent thresholds of toxicity to *Daphnia magna* for chlorides of various metals when added to Lake Erie water. Trans. Amer. Fish. Soc. 78: 96-113.

Anderson, B. G. (1960). The toxicity of organic insecticides to *Daphnia*. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center. Tech. Rept. W603, pp. 94-95.

Anderson, J. B. (1962). Evaluation of streams by biological studies. Proc. 16th Ind. Waste Conf. Purdue Univ. 46(2): 1-7.

Anderson, J. M. and M. R. Peterson. (1969). DDT: Sublethal effects on brook trout nervous system. Science 4(3878): 440-441.

Angelovic, J. W., W. F. Sigler, and J. M. Neuhold. (1961). Temperature and fluorosis in rainbow trout. J. Water Pollut. Contr. Fed. 33: 371-381.

Anon (1958). Toxicity index proposed. Chem. Eng. News 36(52): 26.

Anon (1964). Toxicity test on disinfectants. Water Wast Treat. 10: 210.

Anon (1966). The BOD textile chemicals, updated list – 1966. Amer. Dyestuff Rept. 55(18): 39-41.

Anon (1967). Aquatic life water quality criteria. Environ. Sci. Tech. 1(11): 888-897.

Anon (1967). New York WPCF conference. Water Sewage Works 114(12): 483-485.

Anon (1968). Waste problems of agriculture and forestry. Environ. Sci. Tech. 2(7): 498-503.

Anon (1968). Exploiting and polluting oceans. Nature 219: 840-842.

Anon (1968). Pollution show sparks a debate. Chem. Week 102: 60-61.

Anon (1968). The cost of clean water. Environ. Sci. Tech. 2(4): 257-266.

Applegate, V. C., J. H. Howell, J. W. Moffett, B.G.H. Johnson, and M. A. Smith. (1961). Use of 3-trifluormethyl-4-nitophenol as a selective sea lamprey larvicide. Ann Arbor, Mich., Great Lakes Fish. Comm., Tech. Rept. No. 1, 35 p.

Applegate, V. C., J. H. Howell, and M. A. Smith. (1958). Use of mononitrophenols containing halogens as selective sea lamprey larvicides. Science 127: 336-337.

Applegate, V. C. and E. L. King. (1962). Comparative toxicity of 3-trifluormethyl-4-nitrophenol (TFM) to larval lampreys and eleven species of fishes. Trans. Amer. Fish. Soc. 91: 342-345.

Aquatic Life Advisory Committee of the Ohio River Valley Water Sanitation Commission. (1955). Aquatic life water quality criteria; First progress report. Sewage Ind. Wastes 27: 321-331.

Aquatic Life Advisory Committee of the Ohio River Valley Water Sanitation Commission. (1956). Aquatic life water quality criteria; Second progress report. Sewage Ind. Wastes 28: 678-690.

Aquatic Life Advisory Committee of the Ohio River Valley Water Sanitation Commission. (1960). Aquatic life water quality criteria; Third progress report. J. Water Pollut. Contr. Fed. 32(1): 65-82.

Arnold, G. E. (1962). Thermal pollution of surface supplies. J. Amer. Water Works Assoc. 54: 1332-1346.

Averett, R. C. and C. W. Brinck. (1960). Macroinvertebrates of the Clark Fork River, Montana; a pollution survey. Helena, Montana State Bd. Health and Montana State Fish and Game Dept., Water Pollut. Contr. No. 61-1, 27 p.

Bagenal, T. B. (1963). Propylene phenoxetol as a fish anaesthetic, Nature 197: 1222-1223.

Ball, I. R. (1966). Toxicity of dimethyl sulphoxide to the goldfish, *Carassius auratus*. Nature 210(5036): 639-640.

Ball, I. R. (1967). The relative susceptibilities of some species of freshwater fish to poisons. I. Ammonia. Water Res. 1: 767-775.

Ball, I. R. (1967). The relative susceptibilities of some species of freshwater fish to poisons. II. Zinc. Water Res. 1: 777-783.

Ball, I. R. (1967). The toxicity of cadmium to rainbow trout (Salmo gairdnerii Richardson). Water Res. 1: 805-806.

Barnhart, R. A. (1958). Chemical factors affecting the survival of game fish in a Western Colorado reservoir. (Abstract of thesis.) Colorado Cooperative Fish. Unit, Quart. Rept. 4: 25-27.

Bartsch, A. F. and W. M. Ingram. (1959). Stream life and the pollution environment. Public Works 90(7): 104-110.

Bartsch, A. F. and W. M. Ingram. (1966). Biological analysis of water pollution in North America. Verh. Internat. Verein. Limnol. 16: 786-800.

Batte, E. G. L. E. Swanson, and J. B. Murphy. (1951). New molluscicides for the control of freshwater snails. Amer. J. Vet. Res. 12(43): 158-160.

Beak, T. W. (1957). Industrial aqueous pollution problems. Chem. Can. 9: 38-40.

Beak, T. W. (1958). Toleration of fish to toxic pollution. J. Fish. Res. Bd. Can. 15: 559-572.

Beak, T. W. (1965). A biotic index of polluted streams and its relationship to fisheries. In: Advances in Water Pollution Research, Proc. 2d International Conference held in Tokyo, August 1964. Vol. 1. N.Y., Pergamon Press, pp. 191-219.

Beauchamp, R.S.A. (1967). The effect of biological factors on the design and operation of power stations. In: M. Brook (ed.), Biology and the Manufacturing Industries, Proceedings of a Symposium held at the Royal Geographical Society, London, 29-30 September 1966. London, Academic Press, pp. 43-52.

Beck, W. M. (1957). The use and abuse of indicator organisms. In: C. M. Tarzwell (ed. & comp.), Biological problems in water pollution, Trans. of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, pp. 175-177.

Bedford, J. W., E. W. Roelofs, and M. J. Zabik. (1968). The freshwater mussel as a biological monitor of pesticide concentrations in a lotic environment. Limnol. Oceanogr. 13(1): 118-126.

Bennett, G. W. (1962). Management of artificial lakes and ponds. N.Y., Reinhold Publ. Corp. 283 p.

Benville, P. E., C. E. Smith, and W. E. Shanks. (1968). Some toxic effects of dimethyl sulfoxide in salmon and trout. Toxicol. Appl. Pharm. 12: 156-178. Betts, J. L., T. W. Beak, and G. G. Wilson. (1967). A procedure for small-scale laboratory bioassays. J. Water Pollut. Contr. Fed. 39(1): 89-96.

Betts, J. L. and G. G. Wilson. (1966). New methods for reducing the toxicity of Kraft mill bleachery wastes to young salmon. J. Fish. Res. Bd. Can. 23: 813.

Beyerle, G. B. and J. E. Williams. (1967). Attempted control of bluegill reproduction in lakes by the application of copper sulfate crystals to spawning nests. Prog. Fish-Cult. 29(3): 150-155.

Bhatla, M. N. and A. F. Gaudy. (1965). Role of protozoa in the diphasic exertion of BOD. J. San. Eng. Div., Proc. Amer. Soc. Civil Eng. 91(SA3): 63-87.

Bick, H. (1963). A review of Central European methods for the biological estimation of water pollution levels. Bull. World Health Organ. 29: 401-413.

Binns, N. A. (1967). Effects of rotenone treatment on the fauna of the Green River, Wyoming. Wyoming Game Fish Comm., Fish. Res. Bull. No. 1, 114 p.

Bishop, E. L. (1947). Effects of DDT mosquito larviciding on wildlife. Pt. III. The effects on the plankton population of routine larviciding with DDT. Public Health Repts. 62: 1263-1268.

Bishop, W. D., R. C. Carter and H. F. Ludwig. (1967). Water pollution hazards from refuse produced carbon dioxide. In: Advances in Water Pollution Research, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 1. Washington, D.C., pp. 207-225.

Blackburn, R. D. and L. W. Weldon. (1965). The sensitivity of duckweeds (Lemnaceae) and *Azolla* to Diquat and Paraquat. Weeds 13: 147-149.

Blumenkrantz, B. L. (1956). The effects of Chloreton on developing fish embryos. Proc. Oklahoma Acad. Sci. 35: 62-65.

Bohmont, B. L. (1967). Toxicity of herbicides to livestock, fish, honey bees, and wildlife. Proc. West. Weed Contr. Conf. 21: 25-27.

Bond, C. E., J. D. Fortune, and F. Young. (1965). Results of preliminary bioassays with Kurosal SL and Dicamba. Progr. Fish-Cult. 27: 49-51.

Bond, C. E., R. H. Lewis and J. L. Fryer. (1960). Toxicity of various herbicidal materials to fishes. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center. Tech. Rept. W60-3, pp. 96-101.

Bonde, G. J. (1966). Bacteriological methods for estimation of water pollution. Health Lab. Sci. 3(2): 124-128.

Bonde, G. J. (1967). Heterotrophic bacteria in a polluted marine environment. In: Advance in Water Pollution, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 3. Washington, D.C., Water Pollut. Contr. Fed., pp. 87-103.

Boni, P. (1965). Acute toxicity and elimination of phenol injected into fish (*Carassius auratus* L.). Experientia 21: 222-223.

Bonn, E. W. and B. J. Follis. (1967). Effects of hydrogen sulfide on channel catfish (*Ictalurus punctatus*): Proc. 20th Ann. Conf., Southeast Assoc. Game Fish Comm., pp. 424-432.

Bonn, E. W. and L. R. Holbert. (1961). Some effects of rotenone products on municipal water supplies. Trans. Amer. Fish. Soc. 90: 287-297.

Boschetti, M. M. and T. F. McLoughlin. (1957). Toxicity of sodium arsenite to minnows. Sanitalk 4: 14-18.

Boudreaux, J., K. Strawn and G. Callas. (1959). Fire ants, Heptachlor, & fish kill. Southwest. Natur. 3: 7-12.

Bower, B. T. (1965). Industrial water utilization. Ind. Water Eng. 2(9): 10-15.

Boyd, C. E. and D. E. Ferguson. (1964). Spectrum of crossresistance to insecticides in the mosquito fish, *Gambusia affinis*. Mosquito News 24: 19-21.

Boyd, C. E. and D. E. Ferguson. (1964). Susceptibility and resistance of mosquito fish to several insecticides. Econ. Entomol. 57: 430-431.

Boyd, C. E., S. B. Vinson, and E. E. Ferguson. (1963). Possible DDT resistance in two species of frogs. Copiea, No. 2: 426-429.

Breese, W. P., R. E. Millimann and R. E. Dimic. (1963). Stimulation of spawning in the mussels, *Mytilus edulis* Linnaeus and *Mytilus californianus* Conrad, by Kraft mill effluent. Biol. Bull. 125: 197-205.

Breidenbach, A. W., C. G. Gunnerson, F. K.Kawahara, J. J. Lichtenberg, and R. S. Green. (1967). Chlorinated hydrocarbon pesticides in major river basins, 1957-65. Public Health Repts. 82(2): 139-156.

Breidenbach, A. W. and J. J. Lichtenberg. (1963). DDT and Dieldrin in rivers: A report of the National Water Quality Network. Science 141: 899-901.

Brett, J. R. (1956). Some principles in the thermal requirement of fishes. Quart. Rev. Biol. 31(2): 75-87.

Bridges, W. R. (1958). Sodium cyanide as a fish poison. U.S. Fish Wild. Serv., Spec. Sci. Rept., Fish. No. 253, 11 p.

Bridges, W. R. (1961). Disappearance of Endrin from fish and other materials of a pond environment. Trans. Amer. Fish. Soc. 90: 332-334.

Bridges, W. R. and O. B. Cope. (1965). The relative toxicities of similar formulations of pyrethrum and rotenone to fish and immature stoneflies. Pyrethrum Post 8: 3-5.

Bridges, W. R., B. J. Kallman, and A. K. Andrews. (1963). Persistance of DDT and its metabolites in a farm pond. Trans. Amer. Fish. Soc. 92: 421-427. Brinkhurst, R. O. (1966). Detection and assessment of water pollution using oligochaete worms. Part 1. Water Sewage Works 113: 398-401.

Brinkhurst, R. O. (1966). Detection and assessment of water pollution using oligochaete worms. Part 2. Water Sewage Works 113: 438-441.

Britt, N. W. (1955). New methods of collecting bottom fauna from shoals or rubble bottoms of lakes and streams. Ecology 36(3): 524-525.

Brown, A. C. (1964). Effect of hydrogen sulfide on *Bullia* (Gastropoda). Nature 203: 205-206.

Brown, A.W.A. (1961). The effect of chemical control of insects on wildlife conservation: evaluation of the present in the light of past experience. Proc. Entomol. Soc. Ont. 91: 40-46.

Brown, M. E. (ed.) (1957). The physiology of fishes. Vol. I. Metabolism. N.Y., Academic Press, 447 p.

Brown, M. E. (ed.) (1957). The physiology of fishes. Vol. II. Behavior, Academic Press, 500 p.

Brown, V. M. (1968). The calculation of the acute toxicity of mixtures of poisons to rainbow trout. Water Research 2: 723-733.

Brown, V. M., V. V. Mitrovic, and G.T.C. Stark. (1968). Effects of chronic exposure to zinc on toxicity of a mixture of detergent and zinc. Water Res. 2: 255-263.

Brown, V. M., D.H.M. Jordan, and B. A. Tiller. (1967). The effect of temperature on the acute toxicity of phenol to rainbow trout in hard water. Water Research 1: 587-594.

Brown, V. M., D. G. Shurben, and J. K. Fawell. (1967). The acute toxicity of phenol to rainbow trout in saline waters. Water Research 1: 683-685.

Brungs, W. A. and G. W. Bailey. (1966). Influence of suspended solids on the acute toxicity of Endrin to fathead minnows. Proc. 21st Ind. Waste Conf. Purdue Univ. 50(2): 4-12.

Brungs, W. A. and D. I. Mount. (1967). Lethal Endrin concentration in the blood of gizzard shad. J. Fish. Res. Bd. Can. 24: 429-432.

Bull, C. J. (1968). A bottom fauna samples for use in stony streams. Prog. Fish-Cult. 30(2): 119-120.

Bunting, D. W. and W. H. Irwin. (1965). The relative resistances of seventeen species of fish to petroleum refinery effluents and a comparison of some possible methods of ranking resistances. Proc. 17th Annu. Conf., Southeast. Assoc. Game Fish Comm., pp. 293-307.

Burden, E.H.W.J. (1956). A case of DDT poisoning in fish. Nature 178: 546-547.

Burdick, G. E. (1957). A graphical method for deriving threshold values of toxicity and the equation of the toxicity curve. N.Y. Fish Game J. 4(1): 102-108.

Burdick, G. E. (1960). The use of bioassays by the Water Pollution Control Agency. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, Tech. Rept. W60-3, pp. 145-148.

Burdick, G. E. (1965). Some problems in the determination of the cause of fish kills. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 289-292.

Burdick, G. E. (1967). Use of bioassays in determining levels of toxic wastes harmful to aquatic organisms. In: A symposium on water quality criteria to protect aquatic life. Trans. Amer. Fish. Soc., Spec. Publ. No. 4, pp. 7-12.

Burdick, G. E., H. J. Dean, and E. J. Harris. (1956). Toxicity of emulsifiable rotenone to yellow perch. N.Y. Fish Game J. 3: 75-80.

Burdick, G. E., H. J. Dean, and E. J. Harris. (1958). Toxicity of cyanide to brown trout and smallmouth bass. N.Y. Fish Game J. 5: 133-163.

Burdick, G. E., H. J. Dean, and E. J. Harris. (1960). Effect of Sevin upon the aquatic environment. N.Y. Fish Game J. 7: 14-25.

Burdick, G. E., H. J. Dean, and E. J. Harris. (1964). Toxicity of Aqualin to fingerling brown trout and bluegills. N.Y. Fish Game J. 11: 106-114.

Burdick, G. E., H. J. Dean, E. J. Harris, J. Skea, and D. Colby. (1965). Toxicity of Sevin (Carbaryl) to fingerling brown trout. N.Y. Fish and Game J. 12: 127-146.

Burdick, G. E., E. J. Harris, H. J. Dean, T. M. Walker, J. Skea, and D. Colby. (1964). The accumulation of DDT in lake trout and the effect on reproduction. Trans. Amer. Fish. Soc. 93: 127-136.

Burke, W. D. and D. E. Ferguson. (1968). A simplified flow-through apparatus for maintaining fixed concentrations of toxicants in water. Trans. Amer. Fish Soc. 97(4): 498-501.

Burlington, R. F. (1962). Quantitative biological assessment of pollution. J. Water Pollut. Contr. Fed. 34(2): 179-183.

Burman, N. P. (1966). Recent advances in the bacteriological examination of water. In: C. H. Collins, (ed.), Progress in Microbiological Techniques. N.Y., Plenum Press, pp. 185-212.

Butcher, R. W. (1959). Biological assessment of river pollution. Proc. Linn. Soc. London 170: 159-165.

Butler, P. A. (1965). Effects of herbicides on estuarine fauna. Proc. Southern Weed Conf. 18: 576-580.

Butler, P. A. (1965). Commercial fishery investigations. In: Effects of pesticides on fish and wildlife, 1964 research findings of the Fish and Wildlife Service. U.S. Fish Wildl. Serv., Circ. 226, pp. 65-77.

Butler, P. A. (1966). Pesticides in the marine environment. J. Appl. Ecol. 3(Suppl.): 253-259.

Butler, P. A. (1966). The problem of pesticides in estuaries. In: A symposium on Estuarine Fisheries. Amer. Fish. Soc., Spec. Publ. No. 3, pp. 110-115.

Butler, P. A. (1966). Fixation of DDT in estuaries. Trans. 31st N. Amer. Wildl. Conf., pp. 184-189.

Butler, P. A. (1967). Pesticide residues in estuarine mollusks. In: P. L. McCarty & R. Kennedy. Proceedings of the National Symposium on Estuarine Pollution. Stanford, California, Stanford University, Department of Civil Engineering, pp. 107-121.

Butler, P. and R. F. Johnson. (1967). Report of the Bureau of Commercial Fisheries Biological Laboratory, Gulf Breeze, Florida; Fiscal Year 1966. Washington, D.C., U.S. Fish Wildl. Serv., Bur. Comm. Fish. Circ. 160. iii, 15 pp.

Buzzell, J. C., R.H.F. Young, and D. W. Ryckman. (1968). Behavior of organic chemicals in the aquatic environment. Part II. Behavior in dilute systems. Research report from the Environmental and Sanitary Engineering Laboratories of Washington University, St. Louis, Mo., for the Manufacturing Chemists Association. Washington, D.C., Manufacturing Chemists Association, 81 pp.

Byrd, I. B. (1960). What are the side effects of the imported fire ant control programme? In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center. Tech. Rept. W60-3, pp. 46-50.

Cade, T. J., C. M. White, and J. R. Haugh. (1968). Peregrines and pesticides in Alaska. Condor 70(2): 170-178.

Cairns, J. (1955). The effects of increased temperatures upon aquatic organisms. Proc. 10th Ind. Wastes Conf. Purdue Univ. 40(1): 346-354.

Cairns, J. (1956). Effects of heat on fish. Ind. Wastes 1: 180-183.

Cairns, J. (1957). Environment and time in fish toxicity. Ind. Wastes 2: 1-4.

Cairns, J. (1965). Biological concepts and industrial waste disposal problems. Proc. 20th Ind. Waste Conf. Purdue Univ. 49(4): 49-59.

Cairns, J. (1965). Pollution's eternal triangle. ASB Bull. 12(2): 35-37.

Cairns, J. (1966). Don't be half-safe. The current revolution in bioassay techniques. Proc. 21st Ind. Waste Conf. Purdue Univ. 50(2): 559-567.

Cairns, J. (1967). The use of quality control techniques in the management of aquatic ecosystems. Water Resour. Bull. 3(4): 47-53.

Cairns, J. (1967). Suspended solids standards for the protection of aquatic organisms. 22nd Purdue Industtrial Waste Conference, May 2-4, 1967, Purdue University, 21 pp.

Cairns, J. (1968). The effects of Dieldrin on diatoms. Mosquito News 28(2): 177-179.

Cairns, J. and J. J. Loos. (1966). Changes in guppy populations resulting from exposure to Dieldrin. Progr. Fish-Cult. 28: 220-226. Cairns, J., N. R. Foster, and J. J. Loos. (1967). Effects of sublethal concentrations of Dieldrin on laboratory populations of guppies, *Poecilia reticulata* Peters. Proc. Acad. Natur. Sci. Phila. 119: 75-91.

Cairns, J. and A. Scheier. (Feb. 1955). Bioassay studies for the Manufacturing Chemists' Association; the relationship of body size of the bluegill sunfish to the acute toxicity of some common chemicals. Philadelphia, Penn., Acad. Natur. Sci. Phila., Dept. of Limnol., 49 pp. (Unpubl. rept.).

Cairns, J. and A. Scheier. (1957). The effects of temperature and hardness of water upon the toxicity of zinc to the common bluegill (*Lepomis macrochirus* Raf.) Notulae Natur., No. 299, 12 pp.

Cairns, J. and A. Scheier. (1958). The effects of temperature and hardness of water upon the toxicity of zinc to the pond snail, *Physa heterostropha* (Say). Notulae Natur., No. 308, 11 pp.

Cairns J. and A. Scheier. (1958). The effects of periodic low oxygen upon the toxicity of various chemicals to aquatic organisms. Proc. 12th Ind. Waste Conf. Purdue Univ. 42(3): 165-176.

Cairns, J. and A. Scheier. (1959). The effects of temperature and hardness of water upon the toxicity of potassium dichromate to the common bluegill sunfish. In: Trans. Northeast Wildl. Conf., 10th Ann. Meeting, Jan. 4, 5, 6, and 7, 1958. Montreal, Quebec, Canada, pp. 86-98.

Cairns, J. and A. Scheier. (1959). The relationship of bluegill sunfish body size to tolerance for some common chemicals. Proc. 13th Ind. Waste Conf. Purdue Univ. 43(3): 243-252.

Cairns, J. and A. Scheier. (1962). The effects of temperature and water hardness upon the toxicity of naphthenic acids to the common bluegill sunfish, *Lepomis macrochirus* Raf., and the pond snail, *Physa heterostropha* Say. Notulae Natur., No. 353, 12 pp.

Cairns, J. and A. Scheier. (1963). The acute and chronic effects of standard sodium alkyl benzene sulfonate upon the pumpkinseed sunfish, *Lepomis gibbosus* (Linn.) and the bluegill sunfish *L. macrocjoris* Raf. Proc. 17th Ind. Waste Conf. Purdue Univ. 47(2): 14-28.

Cairns, J. and A. Scheier. (1963). Environmental effects upon cyanide toxicity to fish. Notulae Natur., No. 361, 11 pp.

Cairns, J. and A. Scheier. (1964). The effects of sublethal levels in zinc and of high temperature upon the toxicity of a detergent to the sunfish *Lepomis gibbosus* (Linn.). Notulae Natur., No. 367, 3 pp.

Cairns, J. and A. Scheier. (1964). The effect upon the pumpkinseed sunfish *Lepomis gibbosus* (Linn.) of chronic exposure to lethal and sublethal concentrations of Dieldrin. Notulae Natur., No. 370, 10 pp.

Cairns, J. and A. Scheier. (1966). The influence of exposure to ABS detergent upon the blood chloride regulation of the pumpkinseed. Progr. Fish-Cult. 28: 128-132.

Cairns, J. and A. Scheier. (1968). A comparison of the toxicity of some common industrial waste components tested individually and combined. Progr. Fish-Cult. 30(1): 3-8.

Cairns, J., A. Scheier, and N. E. Hess. (1964). The effects of alkyl benzene sulfonate on aquatic organisms. Ind. Water Wastes 9(1): 22-28.

Cairns, J., A. Scheier, and J. J. Loos. (1965). A comparison of the sensitivity to certain chemicals of adult zebra danios, *Brachydanio rerio* (Hamilton-Buchanan) and zebra eggs with that of adult bluegill sunfish, *Lepomis macrochirus* Raf. Notulae Natur., No. 381, 9 pp.

California State Water Quality Control Board. (1964). An investigation of the effects of discharged wastes on kelp. Sacramento, Calif., State Water Quality Control Board, Publ. No. 26, 125 pp.

California State Water Quality Control Board. (1965). An investigation on the fate of organic and inorganic wastes discharged into the marine environment and their effects on biological productivity. Sacramento, Calif., State Water Quality Control Board, Publ. No. 29, 118 pp.

Carlson, C. A. (1966). Effects of three organophosphorus insecticides on immature *Hexagenia* and *Hydropsyche* of the Upper Mississippi River. Trans. Amer. Fish. Soc. 95: 1-5.

Carpenter, R. A. (1968). Federal policy and environmental chemistry. Environ. Sci. Tech. 2(7): 518-523.

Carter, L. (1962). Bioassay of trade wastes. Nature 196: 1304.

Carter, L. J. (1968). Water Pollution: officials goaded into raising quality standards. Science 160(3283): 49-51.

Casper, V. L. (1967). Galveston Bay pesticide study – water and oyster samples analyzed for pesticide residues following mosquito control program. Pestic. Monit. J. 1(3): 13-15.

Chadwick, G. G. and U. Kiigemagi. (1968). Toxicity evaluation of a technique for introducing Dieldrin into water. J. Water Pollut. Contr. Fed. 40(1): 76-82.

Chadwick, H. K. (1960). Toxicity of tricon oil spill eradicator to striped bass (*Roccus saxatilis*). Calif. Fish Game 46(3): 371-372.

Chang, P. S. (1965). Effects of pollution on oysters and fish in Taiwan. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 368-369.

Chanin, G. and R. P. Dempster. (1958). A complex chemical waste and its toxicity to fish. Ind. Wastes 3: 155-158.

Charles, J. R. (1966). Effects of coal-washer wastes on biological productivity in Martin's Fork of the upper Cumberland River. Kentucky Dept. Fish Wildl. Resour., Fish. Bull. No. 27-B, 48 pp. (Kentucky Proj. F-18-R). Chen, C. W. and R. E. Selleck. (1968). A kinetic model of fish toxicity threshold. A paper presented at the 41st Annual Conference of the Water Pollution Control Federation in Chicago, Sept. 22-27, 1968, 29 pp.

Chin, E. and D. M. Allen. (1957). Toxicity of an insecticide to two species of shrimps, *Penaeus aztecus* and *Penaeus setiferus*. Texas J. Sci. 9: 270-278.

Chowdhury, S. H. (1957). An investigation of the Karnaphuli paper mill effluent and its detrimental effects of fish and other aquatic life of the river. Agr. Pakistan 8: 138-153.

Claffey, F. J. and J. E. Ruck. (1967). The effect of rotenone on certain fish food organisms. Proc. 20th Ann. Conf., Southeast. Ass. Game Fish Comm., pp. 278-283.

Clarke, G. L. (1947). Poisoning and recovery in barnacles and mussels. Biol. Bull. 92: 73-91.

Clean Water Restoration Act of 1966. (PL89-753), approved 11/3/66.

Clemens, H. P., S. Bhinyoying, and N. Youngstead. (1966). An evaluation of Dylox medication on growth and reproduction of the goldfish and guppy. Progr. Fish-Cult. 28: 159-161.

Clemens, H. P. and C. Clough. (1965). Check effluent with bioassay. Petrol. Refiner 35(6): 197-201.

Clemens, H. P. and J. C. Finnell. (1957). Biological conditions in a brine-polluted stream in Oklahoma. Trans. Amer. Fish. Soc. 85: 18-27.

Clemens, H. P. and K. E. Sneed. (1958). The chemical control of some diseases and parasites of channel catfish. Progr. Fish-Cult. 20: 8-15.

Clemens, H. P. and K. E. Sneed. (1958). Effect of temperature and physiological condition on tolerance of channel catfish to pyridylmercuric acetate. Progr. Fish-Cult. 20: 147-152.

Clemens, H. P. and K. E. Sneed. (1959). Lethal doses of several commercial chemicals for fingerling channel catfish. U.S. Fish and Wildl. Serv., Spec. Sci. Rept.: Fish. No. 316, 10 pp.

Clendenning, K. A. and W. J. North. (1960). Effects of wastes on the giant kelp, *Macrocystis pyrifera*. In: E. A. Pearson (ed.), Proc. of the 1st International Conference on Waste Disposal in the Marine Environment. Univ. of Calif., Berkeley, July 22-25, 1959. N.Y., Pergamon Press, pp. 82-91.

Cohen, J. M., L. J. Kamphake, A. E. Lemke, C. Henderson, and R. L. Woodward. (1961). Effect of fish poisons on water supplies. Pt. 1. Removal of toxic materials. J. Amer. Water Works Assoc. 52(12): 1551-1566.

Colby, P. J. and L. L. Smith. (1967). Survival of walleye eggs and fry on paper fiber sludge deposits in Rainy River, Minnesota. Trans. Amer. Fish. Soc. 96: 278-296.

Cole, B. T., H. J. Bennett, and J. D. Miller. (1958). Tolerance of euryhaline fish forms to dilution of oil field bleed water. Proc. La. Acad. Sci. 20: 13-23.

Cole, H., D. Barry, D.E.H. Frear, and A. Bradford. (1967). DDT levels in fish, streams, stream sediments, and soil before and after DDT aerial spray application for fall cankerworm in Northern Pennsylvania. Bull. Environ. Contam. Toxicol. 2: 127-146.

Colton, J. B. (1959). A field observation of mortality of marine fish larvae due to warming. Limnol. Oceanogr. 4: 219-222.

Colvin, H. J. and A. T. Phillips. (1968). Inhibition of electron transport enzymes and cholinesterases by Endrin. Bull. Environ. Contam. Toxicol. 3(2): 106-115.

Cook, S. F. and J. D. Conners. (1963). The short-term side effects of the insecticidal treatment of Clear Lake, Lake County, California in 1962. Ann. Entomol. Soc. Amer. 56(6): 819-824.

Cooke, W. B. (1957). Use and value of fungi as biological indicators of pollution. In: C. M. Tarzwell (ed. and comp.), Biological problems in water pollution. Trans. of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, pp. 84-93.

Cooke, W. B. and A. F. Bartsch. (1959). Aquatic fungi in water with high waste loads. Sewage Ind. Wastes 31(11): 1316-1322.

Cooke, R. and P. W. Graham. (1965). The biological purification of the effluent form a lurgi plant gasifying bituminous coals. Int. J. Air Water Pollut. 9(3): 97-112.

Cooke, W. B., H. J. Phaff, M. W. Miller, M. Shifrine, and E. P. Knapp. (1960). Yeasts in polluted water and sewage. Mycologia 52: 210-230.

Cope, O. B. (1960). The retention of DDT by trout and whitefish. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center. Tech. Rept. W60-3, pp. 72-75.

Cope, O. B. (1961). Effects of DDT spraying for spruce budworm on fish in the Yellowstone River system. Trans. Amer. Fish. Soc. 90: 239-251.

Cope, O. B. (1961). Standards for reporting fish toxicity tests. Progr. Fish-Cult. 23: 187-189.

Cope, O. B. (1963). Sport fishery investigations. In: Pesticide-wildlife studies, a review of Fish and Wildlife Service investigations during 1961 and 1962. U.S. Fish Wildl. Serv., Circ. 167, pp. 26-42.

Cope, O. B. (1965). Some responses of freshwater fish to herbicides. Proc. Southern Weed Conf. 18: 439-445.

Cope, O. B. (1965). Sport fishery investigation. In: Effects of pesticides on fish and wildlife, 1964 research findings of the Fish and Wildlife Service. U.S. Fish Wildl. Serv., Circ. 226, pp. 51-63.

Cope, O. B. (1966). Contamination of the freshwater ecosystem by pesticides. J. Appl. Ecol. 3(Suppl.): 33-44.

Cope, O. B. and P. F. Springer. (1958). Mass control of insects: The effects of fish and wildlife. Bull. Entomol. Soc. Amer. 4(2): 52-56.

Copeland, B. J. and T. C. Dorris. (1964). Community metabolism in ecosystems receiving oil refinery effluents. Limnol. Oceanogr. 9: 431-447.

Copeland, J. B. and J. W. Woods. (1959). Preliminary results of several herbicides on aquatic vegetation in Florida. Proc. 12th Ann. Conf. Southeast. Assoc. Game Fish Comm., pp. 199-205.

Corbet, P. S. (1958). Effects of *Simulium* control on insectivorous fishes. Nature 181(4608): 570-571.

Corner, E.D.S. (1959). The poisoning of *Maria squinado* (Herbst) by certain compounds of mercury. Biochem. Pharmacol. 2: 121-132.

Corner, E.D.S. and B. W. Sparrow. (1956). The modes of action of toxic agents. I. Observations on the poisoning of certain crustaceans by copper and mercury. J. Mar. Biol. Assoc. U. K. 35: 531-548.

Costa, H. H. (1965). Responses of freshwater animals to sodium cyanide solutions. I. Fish. Ceylon J. Sci. (Biol. Sci.) 5(2): 41-87.

Costa, H. H. (1965). Responses of freshwater animals to sodium cyanide solutions. II. *Gammarus pulex*. Ceylon J. Sci. (Biol. Sci.) 5(2): 88-96.

Costa, H. H. (1965). Responses of freshwater animals to sodium cyanide solutions. III. Tadpoles of *Rana* temporaria. Ceylon J. Sci. (Biol. Sci.) 5(2): 97-104.

Cottam, C. (1961). Pesticides, chemicals and water pollution. Public Works 92: 206-209.

Cottam, C. and E. Higgins. (1946). DDT and its effect on fish and wildlife. J. Econ. Entomol. 39: 44-52.

Coutant, C. C. (1964). Insecticide Sevin: Effect of aerial spraying on drift of stream insects. Science 146: 420-421.

Cowell, B. C. (1965). The effects of sodium arsenite and Silvex on the plankton populations in farm ponds. Trans. Amer. Fish. Soc. 94: 371-377.

Crance, J. H. (1963). The effects of copper sulfate on *Microcystis* on zooplankton in ponds. Progr. Fish-Cult. 25: 198-202.

Crandall, C. A. and C. J. Goodnight. (1959). The effect of various factors on the toxicity of sodium pentachlorophenate to fish. Limnol. Oceanogr. 4: 53-56.

Crandall, C. A. and C. J. Goodnight. (1962). Effects of sublethal concentrations of several toxicants on growth of the common guppy, *Lebistes reticulatus*. Limnol. Oceanogr. 7: 233-239.

Crosby, D. G. and R. K. Tucker. (1966). Toxicity of aquatic herbicides to Daphnia magna. Science 54: 289-291.

Curry, L. L. (1965). A survey of environmental requirements for the midge (Diptera: Tendipedidae). In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 127-141.

Curtis, E.J.C. (1969). Sewage fungus: Its nature and effects. Water Res. 3(5): 289-311.

Cushing, C. E. and J. R. Olive. (1957). Effects of Toxaphene and rotenone upon the macroscopic bottom fauna of two Northern Colorado reservoirs. Trans. Amer. Fish. Soc. 86: 294-301.

Daiber, F. C. (1952). The food and feeding relationships of the freshwater drum, *Aplodinotus grunniens* Rafinesque in western Lake Erie. Ohio J. Sci. 52: 35-46.

Darsie, R. F. and F. E. Corriden. (1959). The toxicity of Malathion to kill fish (Cyprinodontidae) in Delaware. J. Econ. Entomol. 52(4): 696-700.

Das, M. and P. H. Needham. (1961). Effect of time and temperature on toxicity of insecticides to insects. I. Tests on DDT on larvae of *Aedes aegypti* L. Ann. Appl. Biol. 49(1): 32-38.

Daugherty, F. M. and J. T. Garrett. (1951). Toxicity levels of hydrocyanic acid and some industrial by-products. Texas J. Sci. 3: 391-396.

Davis, A. N., J. B. Gahan, J. A. Fluno, and D. W. Anthony. (1957). Larvicide tests against blackflies in slow-moving streams. Mosquito News 17: 261-265.

Davis, J. T. and W. S. Hardcastle. (1959). Biological assay of herbicides for fish toxicity. Weeds 7: 397-404.

Davis, J. T. and J. S. Hughes. (1963). Further observations on the toxicity of commercial herbicides to bluegill sunfish. Proc. Southern Weed Conf. 16: 337-340.

Davis, R. C. (1954). *Gambusia*-industrial effluent monitors. Water Sewage Works 111: 259-261.

Dawood, I. K. and B. C. Dazo. (1966). Field tests on two new molluscicides (Molucid and WL 8008) in the Egypt-49 project area. Bull. World Health Org. 35: 913-920.

de Calventi, I. B. (1965). Copper poisoning in the snail *Helix pomatia* and its effect on mucous secretion. Ann. N.Y. Acad. Sci. 118(24): 1015-1020.

Degani, J. G. (1943). Studies of the toxicity of ammunition plant wastes to fishes. Trans. Amer. Fish. Soc. 73: 45-51.

Delaporte, A. V. (1958). Pollution of waters by industrial wastes. Can. J. Public Health 49: 154-156.

Dewey, J. E. (1958). Utility of bioassay in the determination of pesticide residues. J. Agr. Food Chem. 6: 274-281.

Dimond, J. B. (1968). Persistence of DDT in crayfish in a natural environment. Ecology 49(4): 759-762.

Dimond, J. B. (1967). Pesticides and stream insects. Maine Forest Serv. [and The Conservation Foundation (Washington, D.C.)] Bull. No. 23, 21 pp.

Dobbins, W. E. (1964). BOD and oxygen relationships in streams. J. San. Eng. Div., Proc. Amer. Soc. Civil Eng. 90(SA3): 53-78.

Domogalla, B. (1935). Eleven years of chemical treatment for Madison Lakes:—Its effect on fish and fish foods. Trans. Amer. Fish. Soc. 65: 115-121.

Doudoroff, P. (1951). Biological observations and toxicity bioassays in the control of industrial waste disposal. Proc. 6th Ind. Waste Conf. Purdue Univ. 35(6): 88-104.

Doudoroff, P. (1956). Some experiments on the toxicity of complex cyanides to fish. Sewage Ind. Wastes 28(8): 1020-1040.

Doudoroff, P., B. G. Anderson, G. E. Burdick, P. S. Galtsoff, W. B. Hart, R. Patrick, E. R. Strong, E. W. Surber, and W. M. Van Horn. (1951). Bioassay methods for the evaluation of acute toxicity of industrial wastes to fish. Sewage Ind. Wastes 23: 1380-1397.

Doudoroff, P. and M. Katz. (1950). Critical review of literature on the toxicity of industrial wastes and their components to fish. I. Alkalies, acids and inorganic gases. Sewage Ind. Wastes 22(11): 1432-1458.

Doudoroff, P. and M. Katz. (1953). Critical review of literature on the toxicity of industrial wastes and their components to fish. II. The metals, as salts. Sewage Ind. Wastes 25(7): 802-839.

Doudoroff, P., G. Leduc, and C. R. Schneider. (1966). Acute toxicity to fish of solutions containing complex metal cyanides, in relation to concentrations of molecular hydrocyanic acid. Trans. Amer. Fish. Soc. 95: 6-22.

Doudoroff, P. and C. E. Warren. (1962). Dissolved oxygen requirements of fishes. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 145-156.

Douglas, N. H. (1960). A study of the comparative use of different species of fish in the toxicity bioassay of petroleum refinery effluent. Proc. 14th Ann. Conf. Southeast. Assoc. Game Fish Comm., pp. 215-222.

Douglas, N. H. and W. H. Irwin. (1962). Relative resistance of fish species to petroleum refinery. Part I. Ind. Water Wastes 7(6): 171-175.

Douglas, N. H. and W. H. Irwin. (1962) Evaluation and relative resistance of sixteen species of fish as test animals in toxicity bioassays of petroleum refinery effluents. Proc. 17th Ind. Waste Conf. Purdue Univ. 47(2): 57-76.

Douglas, N. H. and W. H. Irwin. (1963). Relative resistance of fish to petroleum refinery wastes. Part II. Ind. Water Wastes 8: 23-27.

Douglas, N. H. and W. H. Irwin. (1963). Relative resistance of fish to petroleum refinery wastes. Part III. Ind. Water Wastes 8: 22-25.

Dowden, B. F. (1962). Toxicity of commercial waste-oil emulsifiers to *Daphnia magna*. J. Water Pollut. Contr. Fed. 34(10): 1010-1014.

Dowden, B. F. and H. J. Bennett. (1965). Toxicity of selected chemicals to certain animals. J. Water Pollut. Contr. Fed. 37(9): 1308-1316.

Downing, K. M. (1954). The influence of dissolved oxygen concentration on the toxicity of potassium cyanide to rainbow trout. J. Exp. Biol. 31: 161-164.

Downing, K. M. and J. C. Merkens. (1955). The influence of dissolved-oxygen concentration on the toxicity of unionized ammonia to rainbow trout (*Salmo gairdnerii* Richardson). Ann. Appl. Biol. 43(2): 243-246.

Duffy, J. R. and D. O'Connell. (1968). DDT residues and metabolites in Canadian Atlantic Coast fish. J. Fish. Res. Bd. Can. 25(1): 189-195.

Dugal, L. C. (1968). Pesticide residue in freshwater fish oils and meals. J. Fish. Res. Bd. Can. 25(1): 169-172.

Dugan, P. R. (1967). Influence of chronic exposure to anionic detergents on toxicity of pesticides to goldfish. J. Water Pollut. Contr. Fed. 39(1): 63-71.

Dupree, H. K. (1960). The arsenic content of water, plankton, soil and fish from ponds treated with sodium arsenite for weed control. Proc. 14th Ann. Conf., South-east. Assoc. Game Fish Comm., pp. 132-136.

Edwards, R. W., H. Egan, M. A. Learner, and P. J. Maris. (1964). The control of chironomid larvae in ponds, using the TDE (DDD). J. Appl. Ecol. 1: 97-117.

Eggink, H. J. (1967). Predicted effects of future discharges of industrial wastes into the Eems estuary. In: Advances in Water Pollution Research, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 3. Washington, D.C., Water Pollut. Contr. Fed. pp. 1-27.

Eggler, W. A. (1953). The use of 2,4-D in the control of water hyacinth and alligator weed in the Mississippi Delta, with certain ecological implications. Ecology 34: 409-414.

Eide, P. M., C. C. Deonier, and R. W. Burrell. (1945). Toxicity of DDT to certain forms of aquatic life. J. Econ. Entomol. 38: 492-493.

Eipper, A. W. (1959). Effects of five herbicides on farm pond plants and fish. N.Y. Fish Game J. 6(1): 46-56.

Eisler, R. (1967). Acute toxicity of zinc to the killifish, Fundulus heteroclitus. Chesapeake Sci. 8: 262-264.

Eisler, R. and M. P. Weinstein. (1967). Changes in metal composition of the Quahaug clam, *Mercenaria mercenaria*, after exposure to insecticides. Chesapeake Sci. 8(4): 253-258.

Elson, P. F. (1967). Effects on wild young salmon of spraying DDT over New Brunswick forests. J. Fish. Res. Bd. Can. 24: 731-767.

Elson, P. F. and C. J. Kerswill. (1967). Impact on salmon of spraying insecticide over forests. In: Advances in Water Pollution Research, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 1. Washington, D.C., Water Pollut. Contr. Fed. pp. 55-74.

Elvins, B. J. (1962). Investigation of the animal population in polluted streams. J. Proc. Inst. Sewage Purif. London. 6: 569.

Enderson, J. H. and D. D. Berge. (1968). Chlorinated hydrocarbon residue in peregrines and their prey species from Northern Canada. Condor 70(2): 149-153.

Engdahl, R. B. and F. C. Croxton. (1962). Pollution-a problem in economics. Pulp Pap. Mag. Can. 63(12): 75-78.

Engineering-Science, Inc. (1964). Toxicant-induced behavioural and histological pathology. A quantitative study of sublethal toxication in the aquatic environment. Final Report, Project Year 1962-63. Arcadia, Calif., Engineering-Science, Inc. 128 pp.

Ettinger, M. B. and D. I. Mount. (1967). A wild fish should be safe to eat. Environ. Sci. Tech. 1(3): 203-205.

Everhart, W. H. and W. W. Hassler. (1945). Aquarium studies on the toxicity of DDT to brown trout, Salmo trutta. Trans. Amer. Fish. Soc. 75: 59-64.

Fairchild, E. J. (1955). Low dissolved oxygen: Effect upon the toxicity of certain inorganic salts to the aquatic invertebrate *Daphnia magna*. In: Proc. 4th Ann. Water Symp., Mar. 22-23, 1955. Baton Rouge, La. Eng. Exp. Station, Bull. No. 51, pp. 95-102.

Farmer, J. A. (1960). The effects of industrial and domestic pollution of the Black Warrior River on the net-plankton population. J. Ala. Acad. Sci. 31(6): 447-458.

Faust, S. D. and O. M. Aly. (1964). Water pollution by organic pesticides. J. Amer. Water. Works Assoc. 56(3): 267-279.

Fay, L. D. and W. G. Youatt. (1967). Residues of chlorinated hydrocarbon insecticides in loons, grebes, a gull, and a sample of alewives from Lake Michigan. Mich. Dept. Conserv., Res. Develop. Rept. No. 109, 7 pp.

Federal Water Pollution Control Act (PL84-660), approved 7/9/56.

Federal Water Pollution Control Act Amendments of 1961 (PL87-88), approved 7/20/1961.

Ferguson, D. E. (1963). Notes concerning the effects of Heptachlor on certain poikilotherms. Copeia, No. 2: 441-443.

Ferguson, D. E. (1967). The ecological consequences of pesticide resistance in fishes. Trans. N. Amer. Wildl. Conf. 32: 103-107.

Ferguson, D. E. and C. R. Bingham. (1966). The effects of combinations of insecticides on susceptible and resistant mosquito fish. Bull. Environ. Contam. Toxicol. 1: 97-103.

Ferguson, D. E. and C. R. Bingham. (1966). Endrin resistance in the yellow bullhead, *Ictalurus natalis*. Trans. Amer. Fish. Soc. 95: 325-326.

Ferguson, D. E., W. D. Cotton, D. T. Gardner, and D. D. Culley. (1965). Tolerances to five chlorinated hydrocarbon insecticides in two species of fish from a transect of the lower Mississippi River. J. Miss. Acad. Sci. 11: 239-245.

Ferguson, D. E., D. D. Culley, and W. D. Cotton. (1965). Tolerances of two populations of fresh water shrimp to five chlorinated hydrocarbon insecticides. J. Miss. Acad. Sci. 11: 235-237.

Ferguson, D. E., D. D. Culley, W. D. Cotton, and R. P. Dodds. (1964). Resistance of chlorinated hydrocarbon insecticides in three species of fresh-water fish. BioScience 14(11): 43-44.

Ferguson, D. E., D. T. Gardner, and A. L. Lindley. (1966). Toxicity of Dursban to three species of fish. Mosquito News 26(1): 80-82.

Ferguson, D. E. and C. P. Goodyear. (1967). The pathway of Endrin entry in black bullheads, *Ictalurus melas*. Copiea, No. 2: 467-468.

Ferguson, G. E., L. W. Grayson, C. M. Ogborn, and P. Weir. (1966). Oil pollution water supplies. J. Amer. Water Works Assoc. 58: 813-821.

Ferguson, D. E., J. L. Ludke, and G. G. Murphy. (1966). Dynamics of Endrin uptake and release by resistant and susceptible strains of mosquito fish. Trans. Amer. Fish. Soc. 95: 335-344.

Ferguson, D. E., J. L. Ludke, J. P. Wood, and J. W. Prather. (1965). The effects of mud on the bioactivity of pesticides on fishes. J. Miss. Acad. Sci. 11: 219-228.

Fitzgerald, G. P. (1958). Control of growth of algae with CMU. Trans. Wis. Acad. Sci. Arts Lett. 46: 281-294.

Fitzgerald, G. P. and S. L. Faust. (1963). Bioassay for algicidal vs. algistatic chemicals. Water Sewage Works 110: 296-298.

Fitzgerald, G. P. and S. L. Faust. (1963). Factors affecting the algicidal and algistatic properties of copper. Appl. Microbiol. 11: 345-351.

Fitzgerald, G. P., G. C. Gerloff, and F. Skoog. (1952). Studies on chemicals with selective toxicity to blue-green algae. Sewage Ind. Wastes 24(7): 888-896.

Foster, N. R., A. Scheier, and J. Cairns. (1966). Effects of ABS on feedings behavior of flagfish, *Jordanella floridae*. Trans. Amer. Fish. Soc. 95: 109-110.

Foster, R. F. (1967). Problems of water pollution associated with thermal energy industries. Richland, Wash. Battelle Memorial Institute, Pacific Northwest Laboratory, BNWL-SA-1546, 8 pp.

Foster, R. F. and P. A. Olson. (1951). An incident of high mortality among large rainbow trout after treatment with pyridylmercuric acetate. Progr. Fish-Cult. 13: 129-130.

Frank, P. A., N. E. Otto, and T. R. Bartley. (1961). Techniques for evaluating aquatic weed herbicides. Weeds 9(4): 515-521.

Frear, D.E.H. (1965). Pesticide index, 3rd ed. State College, CPa., College Sci. Publ. 295 p.

Frear, D.E.H. (1968). Pesticide handbook-Entoma. 20th ed. State College, Pa., College Sci. Publ. 323 p.

Freeman, L. (1953). A standardized method for determining toxicity of pure compounds to fish. Sewage Ind. Wastes 25: 845-848.

Freeman, L. (1953). Toxicity thresholds of certain sodium sulfonates for *Daphnia magna* Straus. Sewage Ind. Wastes 25: 1331-1335.

Freeman, L. and I. Fowler. (1953). Toxicity of combinations of certain inorganic compounds to *Daphnia magna* Straus. Sewage Ind. Wastes 25: 1191-1195.

Fremling, C. R. (1964). Mayfly distribution indicates water quality on the Upper Mississippi River. Science 146: 1164-1166.

Fribourgh, J. H. (1965). The effects of potassium tellurite on goldfish spermatozoan activity. Trans. Amer. Fish. Soc. 94: 399-402.

Fromm, P. O. (1965). Physiological considerations in studies of the action of pollutants on aquatic animals. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 316-319.

Fromm, P. O. and J. R. Gillette. (1968). Effect of ambient ammonia on blood ammonia and nitrogen excretion of rainbow trout (*Salmo gairdneri*). Comp. Biochem. Physiol. 26: 887-896.

Fromm, P. O. and R. H. Schiffman. (1958). Toxic action of hexavalent chromium on largemouth bass. J. Wildl. Manage. 22(1): 40-44.

Fromm, P. O. and R. M. Stokes. (1962). Assimilation and metabolism of chromium by trout. J. Water Pollut. Contr. Fed. 34(11): 1151-1155.

Fruh, E. G., K. M. Stewart, G. F. Lee, and G. A. Rohlich. (1966). Measurements of eutrophication and trends. J. Water Pollut. Contr. Fed. 38(8): 1237-1258.

Fry, F.E.J. (1960). The oxygen requirements of fish. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, R. A. Taft, San. Eng. Center, Tech. Rept. W60-3, pp. 106-109.

Fry, F.E.J. (1960). Requirements for the aquatic habitat. Pulp Pap Mag. Can. 61: T.61-T.66.

Fujiya, M. (1965). Physiological estimation on the effects of pollutants upon aquatic organisms. In: Advances in Water Pollution Research, Proc. 2nd International Conference held in Tokyo, August 1964. Vol. 3. N.Y., Pergamon Press. pp. 315-331. Fukano, K. G. and F. F. Hooper. (1958). Toxaphene (chlorinated camphene) as a selective fish poison. Progr. Fish-Cult. 20(4): 189-190.

Gakstatter, J. H. (1968). Rates of accumulation of 14C-dieldrin residues in tissues of goldfish exposed to a single sublethal dose of 14C-aldrin. J. Fish. Res. Bd. Can. 25(9): 1797-1801.

Gakstatter, J. H. and C. M. Weiss. (1967). The elimination of DDT- $C^{14}$ , dieldrin- $C^{14}$ , and lindane- $C^{14}$ , from fish following a single sub-lethal exposure in aquaria. Trans. Amer. Fish. Soc. 96: 301-307.

Galtsoff, P. S., W. A. Chipman, J. B. Engle, and H. N. Calderwood. (1947). Ecological and physiological studies of the effect of sulfate pulp mill wastes on oysters in the York River, Virginia. U.S. Fish and Wildl. Serv., Fish. Bull. 51: 59-186.

Ganelin, R. S., C. Cueto, and G. A. Mail. (1964). Farm use of pesticides not to blame for massive river fish kills. Chem. Eng. News 42(23): 23-24.

Gannon, J. J. (1966). River and laboratory BOD rate considerations. J. San. Eng. Div., Proc. Amer. Soc. Civil Eng. 92(SA1): 135-161.

Gannon, J. J., H. A. Dirasian, and J. D. Phaup. (1966). A versatile outdoor channel for water pollution investigations. Proc. 21st Ind. Waste Conf., Purdue Univ. 50(2): 234-247.

Garrett, J. T. (1957). Toxicity considerations in pollution control. Ind. Wastes 2(1): 17-19.

Garrett, J. T. (1957). Toxicity investigations on aquatic and marine life. Public Works 88(12): 95-96.

Garrison, R. L. (1968). The toxicity of pro-nox fish to salmonid eggs and fry. Prog. Fish-Cult. 30: 35-38.

Gaufin, A. R. (1957). The use and value of aquatic insects as indicators of organic enrichment. In: C. M. Tarzwell (ed. and comp.), Biological problems in water pollution. Trans. of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, R. A. Taft San. Eng. Center, pp. 136-143.

Gaufin, A. R. (1958). The effects of pollution on a midwestern stream. Ohio J. Sci. 58(4): 197-208.

Gaufin, A. R. (1961). Bioassays to determine the toxicity of pesticides to aquatic invertebrates. Proc. 15th Ind. Waste Conf. Purdue Univ. 45(2): 94-98.

Gaufin, A. R. (1965). Environmental requirements of plecoptera. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 105-110.

Gaufin, A. R., L. D. Jensen, A. V. Nebeker, T. Nelson, and R. W. Teel. (1965). The toxicity of ten organic insecticides to various aquatic invertebrates. Water Sewage Works 112: 276-279. Gaufin, A. R., L. Jensen, and T. Nelson. (1961). Bioassays determine pesticide toxicity to aquatic invertebrates. Water Sewage Works 108: 355-359.

Gaufin, A. R. and C. M. Tarzwell. (1952). Aquatic invertebrates as indicators of stream pollution. Public Health Repts. 67(1): 57-64.

Gaufin, A. R. and C. M. Tarzwell. (1956). Aquatic macroinvertebrate communities as indicators of organic pollution in Lytle Creek. Sewage Ind. Wastes 28(7): 906-924.

Gaylord, W. E. and B. R. Smith. (1966). Treatment of East Bay, Alger County, Michigan, with Toxaphene for control of sea lampreys. In: Investigations in fish control. No. 7. U.S. Fish and Wildl. Serv., Bur. Sport Fish. and Wildl., Resour. Publ. 11, 7 p.

Geen, G. H. and F. J. Andres. (1961). Limnological changes in Seton Lake resulting from hydroelectric diversions. New Westminster, B.C., Int. Pac. Salmon Fish. Comm., Progr. Rept. No. 8, 76 p.

George, J. L. (1959). Effects on fish and wildlife of chemical treatments of large areas. J. Forest. 57: 250-254.

George, J. L., R. F. Darsie, and P. F. Springer. (1957). Effects on wildlife of aerial applications of Strobane, DDT, and BHC to tidal marshes in Delaware. J. Wildl. Manage. 21: 42-53.

George, J. L. and D.E.H. Frear. (1966). Pesticides in the Antarctic. J. Appl. Ecol. 3(Suppl.): 155-167.

George, M. G., S. Z. Qasim, and A. Q. Siddipi. (1966). A limnological survey of the River Kali with reference to fish mortality. Environ. Health 8: 262-269.

Gersdorff, W. A. (1943). Effect of introducing the carboxyl group into the phenol molecule on toxicity to goldfish. Amer. J. Pharm. 115: 159-167.

Gersdorff, W. A. and L. E. Smith. (1940). Effect of halogenation of phenol on its toxicity to goldfish. I. Monchlorophenols. Amer. J. Pharm. 112: 197-204.

Gersdorff, W. A. and L. E. Smith. (1940). Effect of introduction of the halogens into the phenol molecule on toxicity to goldfish. II. Monobromophenols. Amer. J. Pharm. 112: 316-322.

Gersdorff, W. A. and L. E. Smith. (1940). Effect of introduction of the halogens into the phenol molecule on toxicity to goldfish. III. Monoiodophenols. Amer. J. Pharm. 112: 389-394.

Gilderhus, P. A. (1966). Some effects of sublethal concentrations of sodium arsenite on bluegills and the aquatic environment. Trans. Amer. Fish. Soc. 95: 289-296.

Gilderhus, P. A. (1967). Effects of Diquat on bluegills and their food organisms. Progr. Fish-Cult. 29: 67-74.

Gillespie, D. M. (1964). Some toxic effects of Malathion on *Daphnia*. Proc. Montana Acad. Sci. 24: 11-17.

Gillette, L. A., D. L. Miller, and H. E. Redman. (1952). Appraisal of a chemical waste problem by fish toxicity test. Sewage Ind. Wastes 24: 1397-1401.

Ginsburg, J. M. (1945). Toxicity of DDT to fish. J. Econ. Entomol. 38: 274-275.

Ginsburg, J. M. (1947). Results from feeding mosquito larvae, killed by DDT, to goldfish. J. Econ. Entomol. 40: 276-276.

Gjulian, C. M., O. B. Cope, B. F. Quisenberry, and F. R. DuChanois. (1949). The effects of some insecticides on black fly larvae in Alaskan streams. J. Econ. Entomol. 42: 100-105.

Godsil, P. J. and W. C. Johnson. (1968). Pesticide monitoring of the aquatic biota at the Tule Lake National Wildlife Refuge. Pestic. Monit. J. 1(4): 21-26.

Gohar, H.A.F. and H. El-Gindy. (1961). Tolerance of vector snails of bilharziasis and fascioliasis to some chemicals. Proc. Egypt. Acad. Sci. 16: 37-48.

Goodman, A. S. and W. E. Dobbins. (1966). Mathematical model for pollution control programs. J. San. Eng. Div., Proc. Amer. Soc. Civil Eng. 92(6): 1-19.

Goodman, J. R. (1951). Toxicity of zinc for rainbow trout (Salmo gairdnerii). Calif. Fish Game 37(2): 191-194.

Goodnight, C. J. (1942). Toxicity of sodium pentachlorophenate and pentachlorophenol to fish. Ind. Eng. Chem. 34: 868-872.

Gorham, E. and A. G. Gordon. (1963). Some effects of smelter pollution upon aquatic vegetation near Sudbury, Ontario. Can. J. Bot. 41: 371-378.

Gould, W. R. and W. H. Irwin. (1965). The suitabilities and relative resistances of twelve species of fish as bioassay animals for oil-refinery effluents. Proc. 16th Ann. Conf., Southeast. Assoc. Game Fish Comm., Charleston, South Carolina, pp. 333-348.

Graham, J. E. and R. D. Anderson. (1958). The effects of mosquito larviciding on other organisms in Slat Lake County. Utah Acad. Sci. Arts Lett., Proc. 35: 43-48.

Graham, R. J. (1960). Effects of forest insect spraying on trout and aquatic insects in some Montana streams. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, R. A. Taft San. Eng. Center, Tech. Rept. W60-3. pp. 62-65.

Graham, R. J. and D. O. Scott. (1958). Effects of forest insect spraying on trout and aquatic insects in some Montana streams. Final report. Helena, Mont., Montana Fish and Game Dept., 40 p.

Graham, R. J. and D. O. Scott. (1959). Effects of an aerial application of DDT on fish and aquatic insects in Montana. Final report. Helena, Mont., Montana State Fish and Game Dept., 35 p.

Grande, M. (1967). Effect of copper and zinc on salmonid fishes. In: Advances in Water Pollution Research, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 1. Washington, D.C., Water Pollut. Contr. Fed. pp. 97-111.

Gray, L. (1950). The toxicity of Aldrin and Dieldrin to goldfish, *Carasius auratus*, and bluegill sunfish, *Lepomis macrochirus*. Little Rock, Ark., Arkansas Game Fish Commission. 4 p. (unpubl.)

Greenbank, J. (1940). Selective poisoning of fish. Trans. Amer. Fish. Soc. 70: 80-86.

Greer, G. L. and U. Paim. (1968). Degradation of DDT in Atlantic salmon (*Salmo salar*). J. Fish. Res. Bd. Can. 25(11): 2321-2326.

Grenier, F. (1960). A constant flow apparatus for toxicity experiments on fish. J. Water Pollut. Contr. Fed. 32(10): 1117-1119.

Grindley, J. (1946). Toxicity to rainbow trout and minnows of some substances known to be present in waste waters discharged to rivers. Ann. Appl. Biol. 33: 103-112.

Grzenda, A. and H. P. Nicholson. (1965). Distribution and magnitude of insecticide residues among various components of a stream system. Proc. S. Water Resour. Pollut. Contr. Conf. 14: 165-174.

Harp, G. L. and R. S. Campbell. (1964). Effects of the herbicide Silvex on benthos of a farm pond. J. Wildl. Manage. 28: 308-317.

Harrington, R. W. and W. L. Bidlingmayer. (1958). Effects of Dieldrin on fishes and invertebrates of a salt marsh. J. Wildl. Manage. 22: 76-82.

Harrison, J.W.E. and E. W. Rees. (1946). 2,4-D toxicity. I. Toxicity towards certain species of fish. Amer. J. Pharm. 118: 422-425.

Harukawa, C. (1922-23). Preliminary report on the toxicity of colloidal sulphur to fish. Trans. Amer. Fish. Soc. 52: 219-224.

Hasler, A. D. and W. J. Wisby. (1949). Use of fish for the olfactory assay of pollutants (phenols) in water. Trans. Amer. Fish. Soc. 79: 64-70.

Hassall, K. A. (1962). A specific effect of copper on the respiration of *Chlorella vulgaris*. Nature 193: 90.

Hastings, E., W. H. Kittams, J. H. Pepper. (1961). Repopulation by aquatic insects in streams sprayed with DDT. Ann. Entomol. Soc. Amer. 54: 436-437.

Hatch, R. W. (1957). Relative sensitivity of salmonids to DDT. Progr. Fish-Cult. 19: 89-91.

Hawkes, H. A. (1963). Effects of domestic and industrial discharges on the ecology of riffles in midland streams. Int. J. Air Water Pollut. 7: 565-586.

Hawksley, R. A. (1967). Advanced water pollution analysis by a water laboratory. Analyzer 8(1): 13-15.

Haydu, E. P. (1968). Biological concepts in pollution control. Ind. Water Eng. 5(7): 18-21.

Haydu, E. P., H. R. Amberg, and R. E. Dimick. (1952). The effect of Kraft mill waste components on certain salmonoid fishes of the Pacific Northwest. Tappi 35: 545-549.

Hayes, M. L. (1955). Evaluation of Ryania and Malathion as fish toxins. Colorado Cooperative Fish. Unit, Quart. Rept. 1(3): 40-42.

Haynes, H. L., H. H. Moorefield, A. J. Borash, and J. W. Keays. (1958). Toxicity of Sevin to goldfish. J. Econ. Entomol. 51: 540.

Hayton, W. L. and N. A. Hall. (1968). Apparent pH dependence of ethanol absorption rate in the common guppy. J. Pharm. Sci. 57(1): 158-161.

Hazeltine, W. E. (1963). The development of a new concept for control of the Clear Lake gnat. J. Econ. Entomol. 56: 621-626.

Helms, D. R. (1967). Use of formalin for selective control of tadpoles in the presence of fishes. Progr. Fish-Cult. 29: 43-47.

Hemens, J. (1966). The toxicity of ammonia solutions to the mosquito fish (*Gambusia affinis* Baird and Girard). J. Proc. Inst. Sewage Purif. 3: 265-271.

Hemphill, J. E. (1954). Toxaphene as a fish toxin. Progr. Fish-Cult. 16: 41-42.

Henderson, C. and Q. H. Pickering. (1958). Toxicity of organic phosphorus insecticides to fish. Trans. Amer. Fish. Soc. 87: 39-51.

Henderson, C. and Q. H. Pickering. (1963). Use of fish in the detection of contaminants in water supplies. J. Amer. Water Works Assoc. 55(6): 715-720.

Henderson, C., Q. H. Pickering, and J. M. Cohen. (1959). The toxicity of synthetic detergents and soaps to fish. Sewage Ind. Wastes 31(3): 295-306.

Henderson, C., Q. H. Pickering, and A. E. Lemke. (1961). The effect of some organic cyanides (nitriles) on fish. Proc. 15th Ind. Waste Conf. Purdue Univ. 45(2): 120-130.

Henderson, C., Q. H. Pickering, and C. M. Tarzwell. (1959). Relative toxicity of ten chlorinated hydrocarbon insecticides to four species of fish. Trans. Amer. Fish. Soc. 88(1): 23-32.

Henderson, C., Q. H. Pickering, and C. M. Tarzwell. (1960). The toxicity of organic phosphorus and chlorinated hydrocarbon insecticides to fish. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, Tech. Rept. W60-3, pp. 76-88. Henderson, C. and C. M. Tarzwell. (1957). Bio-assays for control of industrial effluents. Sewage Ind. Wastes 29(9): 1002-1017.

Hendrick, R. D., F. L. Bonner, T. R. Everett, and J. E. Fahey. (1966). Residue studies on Aldrin and Dieldrin in soils, water and crawfish from rice fields having insecticide contamination. J. Econ. Entomol. 59: 1388-1391.

Hendrick, R. D. and T. R. Everett. (1965). Toxicity to the Louisiana red crawfish of some pesticides used in rice culture. J. Econ. Entomol. 58: 958-961.

Hendrick, R. D., T. R. Everett, and H. R. Caffey. (1966). Effects of some insecticides on the survival, reproduction, and growth of the Louisiana red crawfish. J. Econ. Entomol. 59: 188-192.

Henegar, D. L. (1966). Minimum lethal levels of toxaphene as a piscicide in North Dakota lakes. In: Investigations in fish control. No. 3. Washington, U.S. Fish and Wildl. Serv., Bur. Sport Fish. and Wildl., Resour. Publ. 7, 16 p.

Hennessey, P. V. and D. G. Rosenberg. (1968). Photographic wastes laboratory treatment. Water and Sewage Works, 115: 131.

Hepworth, W. G. (n.d.). Toxicity of the non-ionic detergent Kyro-eo to brook trout, rainbow trout and nymphal mayflies, stoneflies, damselflies and larval midges. Wyoming Game and Fish Comm., Federal Aid in Fish and Wildl. Restoration, Project No. FW-3-R-8, Analysis No. 21, pp. 40-44.

Herbert, D.W.M. (1955). Measuring the toxicity of effluents to fish. Analyst 80: 896-898.

Herbert, D.W.M. (1961). Freshwater fisheries and pollution control. Proc. Soc. Water Treat. Exam. 10: 135-161.

Herbert, D.W.M. (1962). The toxicity to rainbow trout of spent still liquors from the distillation of coal. Ann. Appl. Biol. 50: 755-777.

Herbert, D.W.M., J. S. Alabaster, M. C. Dart, and R. Lloyd. (1961). The effect of china-clay wastes on trout streams. Int. J. Air Water Pollut. 5: 56-74.

Herbert, D.W.M. and K. M. Downing. (1955). A further study of the toxicity of potassium cyanide to rainbow trout (*Salmo gairdnerii* Richardson). Ann. Appl. Biol. 43(2): 237-242.

Herbert, D.W.M., D.H.M. Jordan, and R. Lloyd. (1965). A study of some fishless rivers in the industrial Midlands. J. Proc. Inst. Sewage Purif. London 6: 569-582.

Herbert, D.W.M. and J. C. Merkens. (1952). The toxicity of potassium cyanide to trout. J. Exp. Biol. 29: 632-649.

Herbert, D.W.M. and J. C. Merkens. (1961). The effect of suspended mineral solids on the survival of trout. Int. J. Air Water Pollut. 5: 46-55.

Herbert, D.W.M. and J. M. Richards. (1963). The growth and survival of fish in some suspensions of solids of industrial origin. Int. J. Air Water Pollut. 7: 297-302.

Herbert, D.W.M. and D. S. Shurben. (1963). A preliminary study of the effect of physical activity on the resistance of rainbow trout (*Salmo gairdnerii* Richardson) to two poisons. Ann. Appl. Biol. 52: 321-326.

Herbert, D.W.M. and D. S. Shurben. (1964). The toxicity of fluoride to rainbow trout. Water Waste Treat. J. 10: 141-142.

Herbert, D.W.M. and D. S. Shurben. (1964). The toxicity to fish of mixtures of poisons. I. Salts of ammonia and zinc. Ann. Appl. Biol. 53: 33-41.

Herbert, D.W.M. and A. C. Wakeford. (1962). The effect of calcium sulfate on the survival of rainbow trout. Water Waste Treat. J. 8: 608-609.

Hermann, E. R. (1959). Toxicity index for industrial wastes. Ind. Eng. Chem. 51: 84A-87A.

Hervey, R. J. (1949). Effect of chromium on the growth of unicellular chlorophyceae and diatoms. Bot. Baz. 111(1): 1-11.

Hess, A. D. and G. G. Keener. (1947). Effect of airplane distributed DDT thermal aerosols on fish and fish food organisms. J. Wildl. Manage. 11(1): 1-10.

Hester, F. E. (1959). The tolerance of eight species of warm-water fishes to certain rotenone formulations. Proc. 13th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 121-133.

Hester, F. E. (1959). The toxicity of Noxfish and Pronoxfish to eggs of common carp and fathead minnows. Proc. 13th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 325-331.

Hester, F. E. and J. S. Dendy. (1962). A multiple-plate sampler for aquatic macro-invertebrates. Trans. Amer. Fish. Soc. 91: 420-421.

Heukelekian, H. (1948). Method for studying the toxicity of industrial wastes. Water Sewage Works 95: 285-287.

Hickey, J. H., J. A. Keith, and F. B. Coon (1966). An exploration of pesticides in a Lake Michigan ecosystem. J. Appl. Ecol. 3(Suppl): 141-154.

Hicks, C. E. and J. M. Neuhold. (1966). Alkyl benzene sulfonate effects on stream algae communities. Bull. Environ. Contam. Toxicol. 1: 225-236.

Hildebrand, E. M. (1946). Herbicidal action of 2,4-dichlorophenoxyacetic acid on the water hyacinth, *Eichornia crassipes*. Science 103: 477-479.

Hilsenhoff, W. (1966). Effect of diquat on aquatic insects and related animals. J. Econ. Entomol. 59: 1520-1521.

Hilsenhoff, W. L. (1962). Toxicity of granular Malathion to walleyed pike fingerlings. Mosquito News 22: 14-15.

Hilsenhoff, W. L. (1965). The effect of Toxaphene on the benthos in a thermally-stratified lake. Trans. Amer. Fish. Soc. 94: 210-213.

Hiltibran, R. C. (1965). The effects of Diquat on aquatic plants in Central Illinois. Weeds 13: 71-72.

Hiltibran, R. C. (1967). Effects of some herbicides on fertilized fish eggs and fry. Trans. Amer. Fish. Soc. 96: 414-416.

Hinshaw, R. N. (1967). The pollutional degradation of the Jordan River as shown by aquatic invertebrates. Utah State Dept. Fish Game, Publ. No. 66-11, 121 p.

Hirsch, A. (1958). Biological evaluation of organic pollution of New Zealand streams. N. Z. J. Sci. 1(4): 500-553.

Hisaoka, K. K. (1958). The effects of 2-acetylaminofluorene on the embryonic development of the zebrafish. I. Morphological studies. Cancer Res. 18: 527-535.

Hisaoka, K. K. (1958). The effects of 2-acetylaminofluorene on the embryonic development of the zebrafish. II. Histochemical studies. Cancer Res. 18: 664-667.

Hoar, W. S. (1956). Photoperiodism and thermal resistance of goldfish. Nature 178: 364-365.

Hoeglund, L. B. (1961). The reactions of fish in concentration gradients. A comparative study based on fluvarium experiments with special reference to oxygen, acidity, carbon dioxide, and sulphite waste liquor (SWL). Inst. Freshwater Res., Drottningholm, Rept. No. 43, 147 p.

Hoff, J. G. and J. R. Westman. (1965). Experiments with Dibrom-Malathion formulation as a selective piscicide. N.Y. Fish Game J. 12: 99-107.

Hoffman, C. H. (1960). Are the insecticides required for insect control hazardous to aquatic life? In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, Tech. Rept. W60-3, pp. 51-61.

Hoffman, C. H. and A. T. Drooz. (1953). Effects of a C-47 airplane application of DDT on fish-food organisms in two Pennsylvania watersheds. Amer. Midland Natur. 50: 172-188.

Hoffman, D. A. and J. R. Olive. (1961). The effects of rotenone and Toxaphene upon plankton of two Colorado Reservoirs. Limnol. Oceanogr. 6: 219-222.

Hoffman, C. H. and E. W. Surber. (1945). Effects of an aerial application of wettable DDT on fish and fish-food organisms in Back Creek, West Virginia. Trans. Amer. Fish. Soc. 75: 48-58.

Hoffman, C. H. and E. W. Surber. (1949). Effects of feeding DDT-sprayed insects to fresh-water fish. U.S. Fish Wildl. Serv., Spec. Sci. Rept.-Fish., No. 3, 9 p.

Hoffman, R. A. (1957). Toxicity of three phosphorus insecticides to cold water game fish. Mosquito News 17: 213.

Holden, A. V. (1962). A study of the absorption of 14C-labelled DDT from water by fish. Ann. Appl. Biol. 50: 467-477.

Holden, A. V. (1964). The possible effects on fish of chemicals used in agriculture. J. Proc. Inst. Sewage Purif. London 4: 361-368.

Holden, A. V. (1965). Contamination of fresh water by persistent insecticides and their effects on fish. Ann. Appl. Biol. 55: 332-335.

Holden, A. V. (1966). Organochlorine insecticide residues in salmonid fish. J. Appl. Ecol. 3(Suppl.): 45-53.

Holland, H. T., D. L. Coppage, and P. A. Butler. (1966). Increased sensitivity to pesticides in sheepshead minnows. Trans. Amer. Fish. Soc. 95(1): 110-112.

Holland, H. T., D. L. Coppage, and P. A. Butler. (1967). Use of fish brain acetylocholinesterase to monitor pollution by organophosphorus pesticides. Bull. Environ. Contam. Toxicol. 2: 156-162.

Holland, G. A., J. E. Lasater, E. D. Neumann, and W. E. Eldridge. (1960). Toxic effects of organic and inorganic pollutants on young salmon and trout. Wash. Dept. of Fish., Res. Bull. 5, 264 p.

Hooper, F. F. and A. R. Grzenda. (1955). The use of Toxaphene as a fish poison. Trans. Amer. Fish. Soc. 85: 180-190.

Hopkins, C. L., H. V. Brewerton, and H.J.W. McGrath. (1966). The effect on a stream fauna of an aerial application of DDT prills to pasture land. N. Z. J. Sci. 9: 236-248.

Houser, A. (1962). Loss in weight of sunfish following aquatic vegetation control using the herbicide Silvex. Proc. Okla. Acad. Sci. 43: 232-237.

Howard, T. E. and C. C. Walden. (1965). Pollution and toxicity characteristics of kraft pulp mill effluents. Tappi 48: 136-141.

Howell, J. H., E. L. King, A. J. Smith, and L. H. Hanson. (1964). Synergism of 5,2'-dichloro-4'-nitro-salicylanilide and 3-trifluormethyl-4-nitrophenol in a selective lamprey larvicide. Ann Arbor, Mich., Great Lakes Fish. Comm., Tech. Rept. No. 8, 21 p.

Howell, J. H. and W. M. Marquette. (1963). Use of mobile bioassay equipment in the chemical control of sea lamprey. U.S.Fish Wildl. Serv., Spec. Sci. Rept.-Fish., No. 418, 9 p.

Hubble, D. R. and B. Reiff. (1967). Reproduction of guppies (*Lebistes reticulatus*) after a single exposure to Dieldrin-a 12 months' study. Bull. Environ. Contam. Toxicol. 2: 57-63.

Hubschman, J. H. (1967). Effects of copper on the crayfish *Orconestes rusticus* (Girad). I. Acute toxicity. Crustaceana 12: 33-42.

Huet, M. (1965). Water quality criteria for fish life. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 160-167.

Hughes, J. S. (1966). Use of the red crawfish, *Procambarus clarki* (Girad), for herbicidal assays. Proc. 20th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 437-439.

Hughes, J. S. and J. T. Davis. (1963). Variations in toxicity to bluegill sunfish of phenoxy herbicides. Weeds. 11(1): 50-53.

Hughes, J. S. and J. T. Davis. (1965). Comparative toxicity to bluegill sunfish of granular and liquid herbicides. Proc. 16th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 319-323.

Hughes, J. S. and J. T. Davis. (1967). Effects of selected herbicides on bluegill sunfish. Proc. 18th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 480-482.

Huish, M. T. (1961). Toxaphene as a fish eradicant in Florida. Proc. 15th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 200-205.

Hunn, J. B., R. A. Schoettger, and E. W. Whealdon. (1968). Observations on the handling and maintenance of bioassay fish. Progr. Fish-Cult. 30(3): 164-167.

Hunn, J. B., R. A. Schoettger, and W. A. Willford. (1968). Turnover and urinary excretion of free and acetylated M. S. 22 by rainbow trout, *Salmo gairdneri*. J. Fish. Res. Bd. Can. 25(1): 25-31.

Hunt, E. G. and A. I. Bischoff. (1960). Inimical effects on wildlife of periodic DDD applications to Clear Lake. Calif. Fish Game 46(1): 91-106.

Hunt, G. S. (1965). The direct effects on some plants and animals of pollution in the Great Lakes. BioScience 15: 181-186.

Hutchinson, G. E. (1957). A treatise on limnology. Vol. 1. Geography, physics, chemistry. N.Y., Wiley and Sons, 1015 p.

Hutchinson, G. E. (1967). A treatise on limnology. Vol. 2. Introduction to lake biology and the limnoplankton. N.Y., Wiley and Sons, 1115 p.

Hynes, H.B.N. (1959). The biological effects of water pollution. In: W.B. Yapp (ed.), The effects of pollution on living material. London, Institute of Biology, pp. 11-24.

Hynes, H.B.N. (1959). The use of invertebrates as indicators of river pollution. Proc. Linn. Soc. London 170: 165-169.

Hynes, H.B.N. (1961). The effect of sheep-dip containing the insecticide BHC on the fauna of a small stream, including *Simulium* and its predators. Ann. Trop. Med. Parasitol. 55(2): 192-196.

Hynes, H.B.N. (1966). The biology of polluted waters. Liverpool University Press, 202 p.

Hynes, H.B.N. and F. W. Roberts. (1962). The biological effects of synthetic detergents in the River Lee, Hertfordshire. Ann. Appl. Biol. 50: 779-790.

Hynes, H.B.N. and T. R. Williams. (1962). The effect of DDT on the fauna of a Central African stream. Ann. Trop. Med. Parasitol. 56: 78-91.

Ide, F. P. (1957). Effect of forest spraying with DDT on aquatic of salmon streams. Trans. Amer. Fish. Soc. 86: 208-219.

Ide, F. P. (1967). Effects of forest spraying with DDT on aquatic insects of salmon streams in New Brunswick. J. Fish. Res. Bd. Can. 24: 769-802.

Ingols, R. S. (1954). Toxicity of mercuric chloride, chromic sulfate, and sodium chromate in the dilution B.O.D. test. Sewage Ind. Wastes 26: 536-538.

Ingols, R. S. (1955). Evaluation of toxicity. Sewage Ind. Wastes 27: 26-33.

Ingols, R. S. (1956). Toxicity of copper and zinc ions in the dilution B.O.D. tests. Sewage Ind. Wastes 28: 1168-1169.

Ingols, R. S. (1959). Effect of impoundment on downstream water quality, Catawba River, S. C. J. Amer. Water Works Assoc. 51(1): 42-46.

Ingols, R. S. and R. H. Fetner. (1961). Toxicity of chromium compounds under aerobic conditions. J. Water Pollut. Contr. Fed. 33(4): 366-370.

Ingols, R. S. and P. E. Gaffney. (1965). Biological studies of halophenols. Proc. S. Water Resour. Pollut. Contr. Conf. 14: 175-181.

Ingram, W. M. (1957). Use and value of biological indicators of pollution: Fresh water clams and snails. In: C. M. Tarzwell, (ed., comp.), Biological problems in water pollution. Trans. of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, pp. 94-135.

Ingram, W. M. and W. W. Towne. (1960). Effects of industrial wastes on stream life. Proc. 14th Ind. Waste Conf. Purdue Univ. 44(5): 678-710.

Irukayama, K. (1966). The pollution of Minamata Bay and Minamata disease. Adv. Water Poll. Res. 3: 153-180.

Irwin, W. H. (1965). Fifty-seven species of fish in oilrefinery waste bioassay. Trans. N. Amer. Wildl. Conf. 30: 89-99.

Ishio, S. (1965). Behavior of fish exposed to toxic substances. In: Advances in Water Pollution Research, Proc. 2d International Conference held in Tokyo, August 1964. Vol. 1. N.Y., Pergamon Press, pp. 19-40.

Isom, B. G. (1960). Toxicity of elemental phosphorus. J. Water Pollut. Contr. Fed. 32(12): 1312-1316.

Iyatomi, K., T. Tamura, Y. Itazawa, I. Hanyu, and S. Sugiura. (1958). Toxicity of Endrin to fish. Progr. Fish-Cult. 20(4): 155-162.

Jackson, C. F. (1956). Control of the common sunfish or pumpkinseed, *Lepomis gibbosus*, in New Hampshire. N. H. Fish Game Dept., Tech. Circ. 12: 16 p.

Jackson, K. J. (1960). A field experiment to determine the effect upon coho salmon fry (*Onchorhynchus kisutch*) from spraying sawlogs with an emulsified mixture of benzene hexachloride. Can. Fish Cult. 27: 33-42.

Jackson, K. J. (1966). The role of the Department of Fisheries of Canada in dealing with problems presented by the use of pesticides in British Columbia. Can. Fish Cult. 37: 45-50.

Jackson, H. W. and W. A. Brungs. (1966). Biomonitoring industrial effluents. Ind. Water Eng. 3(7): 14-18, 45.

Jamnback, H. and H. S. Eabry. (1962). Effects of DDT, as used in black fly larval control, on stream arthropods. J. Econ. Entomol. 55(5): 636-639.

Jamnback, H. and J. Frempong-Boadu. (1966). Testing blackfly larvicides in the laboratory and in streams. Bull. World Health Organ. 34(3): 405-421.

Jensen, L. D. and A. R. Gaufin. (1964). Effects of ten organic insecticides on two species of stonefly naiads. Trans. Amer. Fish Soc. 93: 27-34.

Jensen, L. D. and A. R. Gaufin. (1966). Acute and longterm effects of organic insecticides on two species of stonefly naiads. J. Water Pollut. Contr. Fed. 38(8): 1273-1286.

Johnels, A. G., T. Westermark, W. Berg, P. I. Persson, and B. Sjoestrand. (1967). Pike (*Esox lucius* L.) and some other aquatic organisms in Sweden as indicators of mercury contamination in the environment. Oikos 18(2): 323-333.

Johnson, D. W. (1968). Pesticide and fishes-a review of selected literature. Trans. Amer. Fisheries Soc. 97(4): 398-424.

Johnson, M. W. (1959). The effectiveness of Toxaphene at low temperatures. Minn. Dept. Conserv., Div. Game Fish, Invest. Rept. No. 197, 3 p.

Johnson, W. C. (1966). Toxaphene treatment of Big Bear Lake, California. Calif. Fish Game 52: 173-179.

Johnson, W. D. (1968). Identification of residual oil pollutants in surface waters of southern end of Lake Michigan. Chicago, Ill., U.S. Federal Water Pollution Control Administration, Great Lakes Region, Chicago Program Office, 32 p.

Johnson, W. D., F. D. Fuller, and L. E. Scarce. (1967). Pesticides in the Green Bay area. Conf. Great Lakes Res., Proc. 6: 262-374. Jones, B. F., C. E. Warren, C. E. Bond and P. Doudoroff. (1956). Avoidance reactions of salmonid fishes to pulp mill effluents. Sewage Ind. Wastes 28(11): 1403-1413.

Jones, B. R. and J. B. Moyle. (1963). Populations of plankton animals and residual chlorinated hydrocarbons in soils of six Minnesota ponds treated for control of mosquito larvae. Trans. Amer. Fish. Soc. 92(3): 211-215.

Jones, J.R.E. (1938). The relative toxicity of salts and lead, zinc and copper to the stickleback (*Gasterosteus aculeatus* L.) and the effect of calcium on the toxicity of lead and zinc salts. J. Exp. Biol. 15: 394-407.

Jones, J.R.E. (1939). The relation between the electrolytic solution pressures of the metals and their toxicity to the stickleback (*Gasterosteus aculeatus* L.). J. Exp. Biol. 16: 425-437.

Jones, J.R.E. (1941). A study in the relative toxicity of anions, with *Polycelis nigra* as test animal. J. Exp. Biol. 18: 170-181.

Jones, J.R.E. (1947). The reactions of *Pygosteus pungitius* L. to toxic solutions. J. Exp. Biol. 24: 110-122.

Jones, J.R.E. (1948). A further study of the reactions of fish to toxic solutions. J. Exp. Biol. 25: 22-34.

Jones, J.R.E. (1951). The reactions of the minnow, *Phoxinus phoxinus* (L.), to solutions of phenol, orthocresol and para-cresol. J. Exp. Biol. 28: 261-270.

Jones, J.R.E. (1958). A further study of the zinc-polluted River Ystwyth. J. Anim. Ecol. 27: 1-14.

Jones, J.R.E. (1964). Fish and river pollution. London, Butterworths, 203 pp.

Jones, R. O. (1965). Tolerance of the fry of common warm-water fishes to some chemicals employed in fish culture. Proc. 16th Ann. Conf. Southeast. Assoc. Game Fish. Comm. 1962, pp. 436-445.

Jordan, H. D. (1955). Control of crabs with crude BHC. Nature 175(4460): 734-735.

Jordan, D.H.M. and R. Lloyd. (1964). The resistance of rainbow trout (*Salmo gairdnerii* Richardson) and loach (*Rutilus rutilus* (L.)) to alkaline solutions. Int. J. Air Water Pollut. 8: 405-409.

Joyner, T. (1961). Exchange of zinc with environmental solutions by the brown bullhead. Trans. Amer. Fish. Soc. 90: 444-448.

Kabler, P. W. (1957). Use and value of bacterial indicators of pollution. In: C. M. Tarzwell (ed. & comp.), Biological problems in water pollution. Transactions of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, pp. 43-49. Kabler, P. W. (1961). Bacterial parameters of water quality. In: Water Quality Measurement and Instrumentation. Proc. of the 1960 Seminar at Cincinnati, Ohio, August 29-31, 1960. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, SEC-TR-W61-2, pp. 45-50.

Kallman, B. J. and O. B. Cope, and R. J. Navarre. (1962). Distribution and detoxication of Toxaphene in Clayton Lake, New Mexico. Trans. Amer. Fish. Soc. 91: 14-22.

Kaplan, H. M. and J. G. Overpeck. (1964). Toxicity of halogenated hydrocarbon insecticides for the frog, *Rana pipiens*. Herpetologica 20: 163-169.

Kaplan, H. M. and L. Yoh. (1961). Toxicity of copper for frogs. Herpetologica 17: 131-135.

Katz, M. (1961). Acute toxicity of some organic insecticides to three species of salmonids and to the threespine stickleback. Trans. Amer. Fish. Soc. 90: 264-268.

Katz, M. and G. G. Chadwick. (1961). Toxicity of Endrin to some Pacific Northwest fishes. Trans. Amer. Fish. Soc. 90: 394-397.

Katz, M. and A. R. Gaufin. (1953). The effects of sewage pollution on the fish population of a midwestern stream. Trans. Amer. Fish. Soc. 84: 157-165.

Katz, M. and C. E. Woelke. (1967). Water quality requirements of estuarine organisms. In: Water quality criteria. Philadelphia, Pa., Amer. Soc. Testing and Materials, Spec. Tech. Publ. No. 416, pp. 90-99.

Keenleyside, M.H.A. (1958). Comparative effects of the insecticides DDT and Malathion on young Atlantic salmon. In: Progress Reports of the Atlantic Coast Stations, Fish. Res. Bd. Can., No. 69, pp. 3-6.

Keenleyside, M.H.A. (1967). Effects of forest spraying with DDT in New Brunswick on food of young Atlantic salmon. J. Fish. Res. Bd. Can. 24: 807-822.

Keith, J. O. (1966). Insecticide contaminations in wetland habitats and their effects on fish-eating birds. J. Appl. Ecol. 3(Suppl.): 71-85.

Kemp, H. T., R. G. Fuller, and R. S. Davidson. (1966). Potassium permanganate as an algicide. J. Amer. Water Works Assoc. 58(2): 255-263.

Kendeigh, S. C. (1961). Animal Ecology. Englewood Cliffs, N.J., Prentice-Hall, Inc., 468 p.

Kennedy, V. S. and J. A. Mihursky. (1967). Bibliography on the effects of temperature in the aquatic environment. University of Maryland, Natural Resources Institute, Contrib. No. 326, 89 p.

Kerswill, C. J. (1967). Studies on effects of forest sprayings with insecticides, 1952-63, on fish and aquatic invertebrates in New Brunswick streams: Introduction and summary. J. Fish. Res. Bd. Can. 24: 701-708. Kerswill, C. J. and H. E. Edwards. (1967). Fish losses after forest sprayings with insecticides in New Brunswick, 1952-62, as shown by caged specimens and other observations. J. Fish. Res. Bd. Can. 24: 709-729.

Kerswill, J. C., P. J. Elson, M.H.A. Keenleyside, and J. B. Sprague. (1960). Effects of young salmon of forest spraying with DDT. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1969 Seminar. Cincinnati, Ohio. Robt. A. Taft San. Eng. Center, Tech. Rept. W60-3, 71 p.

Ketchum, B. H. (1960). Marine pollution problems in the North Atlanta area. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Transactions of the 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, Tech. Rept. W60-3, pp. 212-217.

Keup, L. E., W. M. Ingram, and K. M. Mackenthun. (1966). The role of bottom-dwelling macrofauna in water pollution investigations. U.S. Public Health Serv. Publ. No. 999-WP-38, 23 pp.

Keup, L. E., W. M. Ingram and K. M. Mackenthun. (1967). Biology of water pollution a collection of selected papers on stream pollution, waste water, and water treatment. Cincinnati, Ohio, Federal Water Pollut. Control Administration, CWA-3, 290 p.

Khan, K. R. (1964). Potentialities of algae in the bio-assay of microchemical pollutants in water systems. Environ. Health 6:274-477.

King, D. L. and R. C. Ball. (1964). A quantitative biological measure of stream pollution. J. Water Pollut. Contr. Fed. 36(5): 650-653.

King, D. L. and R. C. Ball. (1967). Comparative energetics of a polluted stream. Limnol. Oceanogr. 12(10): 27-33.

King, S. F. (1962). Some effects of DDT on the guppy and the brown trout. U.S. Fish Wildl. Serv., Spec. Sci. Report.—Fish., No. 399, 22 p.

King, P. H. (1968). Distribution of pesticides in California. J. San. Eng. Div., Amer. Soc. Civil Eng. 92(SA2): 444-446.

Kinne, O. and H. Rosenthal. (1967). Effects of sulfuric water pollutants on fertilization, enbryonic development and larvae of the herring, *Clupea harengus*. Marine Biol. (Berlin) 1(1): 65-83.

Kinney, E. C. (1964). Extent of acid mine pollution in the United States affecting fish and wildlife. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Fish and Wildl. Cir. 191. iii, 27 p.

Kiser, R. W., J. R. Donaldson, and P. R. Olson. (1963). The effect of rotenone on zooplankton populations in freshwater lakes. Trans. Amer. Fish. Soc. 92(1): 17-24.

Klassen, C. W., W. A. Hasfurther, and M. K. Young. (1948). The toxicity of hexavalent chromium to sunfish and bluegills. Proc. 4th Ind. Waste Conf. Purdue Univ. 33(4): 229-237.

Klock, J. W. (1956). A field technique for quantitative estimation of the molluscicide sodium pentachlorophenate based on fish mortality rates. Amer. J. Trop. Med. 5: 286-289. Knoll, J. and P. O. Fromm. (1960). Accumulation and elimination of hexavalent chromium in rainbow trout. Physiol. Zool. 33(1): 1-8.

Knowles, F. L., W. V. Parker, and H. A. Johnson. (1941). Observations on the use of "phenol" larvicides for mosquito control. Public Health Rep. 56: 1627-1641.

Koeman, J. H. and H. van Genderen. (1966). Some preliminary notes on residues of chlorinated hydrocarbon insecticides in birds and mammals in the Netherlands. J. Appl. Ecol. 3(Suppl.): 99-106.

Kok, L. T. and M. D. Pathak. (1966). Toxicity of Lindane used for Asiatic rice borer control to three species of fish. J. Econ. Entomol. 59: 659-663.

Konar, S. K. (1968). Experimental use of Chlordane in fishery management. Progr. Fish-Cult. 30(2): 96-99.

Kott, Y., G. Hershkovitz, A. Shemtob, and J. B. Sless. (1966). Algicidal effect of bromine and chlorine on *Chorella pyrenoidosa*. Appl. Microbiol. 14(1): 8-11.

Kramer, R. H. and L. L. Smith. (1966). Survival of walleye eggs in suspended wood fibers. Progr. Fish-Cult. 28: 79-82.

Krombach, H. and J. Barthel. (1963). Investigation of a small watercourse accidentally polluted by phenol compounds. Int. J. Air Water Pollut. 7: 39-46.

Krumholz, L. A. and W. L. Minckley. (1964). Changes in the fish population in the Upper Ohio River following temporary pollution abatement. Trans. Amer. Fish. Soc. 93: 1-5.

Lackey, J. B. (1957). Protozoa as indicators of the ecological condition of a body of water. In: C. M. Tarzwell (ed. and comp.), Biological Problems in Water Pollution. Trans. of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, pp. 50-59.

Lackey, J. B. (1958). Effects of fertilization on receiving waters. Sewage Ind. Wastes 30(11): 1411-1416.

Lackey, J. B. (1961). Aquatic organisms. In: Water Quality Measurement and Instrumentation. Proc. of the 1960 Seminar at Cincinnati, Ohio, August 29-31, 1960. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, SEC-TR-W61-2, pp. 37-44.

Lam, R. (1964). A resume of pollution of streams, ponds, and ocean water on Oahu. J. Water Pollut. Contr. Fed. 36(9): 1152-1158.

Lammering, M. W. and N. C. Burbank. (1961). The toxicity of phenol, o-chlorophenol, and o-nitrophenol to bluegill sunfish. Proc. 15th Ind. Waste Conf. Purdue Univ. 45(2): 541-555.

Langer, E. (1964). Pesticides: Minute quantities linked with massive fish kills; federal policy still uncertain. Science 144: 35-37.

Lauer, G. J., H. P. Nicholson, W. S. Cox, and J. T. Teasley. (1966). Pesticide contamination of surface waters by sugar cane farming in Louisiana. Trans. Amer. Fish. Soc. 95: 310-316.

Lawrence, H. M. (1950). Toxicity of some new insecticides to several species of pondfish. Progr. Fish-Cult. 12: 141-146.

Lawrence, J. M. (1962). Aquatic herbicide data. U.S. Dep. Agr., Agr. Handb. No. 231, 133 p.

Lawrence, J. M. (1958). Further investigations on the use of Delrad as an algaecide in fishponds. Progr. Fish-Cult. 20: 89-91.

Lawrence, J. M. (1958). Recent investigations on the use of sodium arsenite as an algicide and its effects on fish production in ponds. Proc. 11th. Conf., Southeast. Assoc. Game Fish Comm., pp. 281-287.

Lawrence, J. M., H. H. Funderburk, R. D. Blackburn, and P. G. Beasley. (1965). The status of Diquat and Paraquat as aquatic herbicides. Proc. 16th Ann. Conf., Southeast. Assoc. Game Fish Comm., Charleston, South Carolina, pp. 247-257.

Learner, M. A. and R. W. Edwards, (1963). The toxicity of some substances of *Nais* (Oligochaeta). Proc. Soc. Water Treat. Exam. 12(3):161-168.

Learner, M. A. and R. W. Edwards. (1966). The distribution of the midge *Chironomus riparius* in a polluted river system and its environs. Int. J. Air Water Pollut. 10: 757-768.

Lemke, A. E. and D. I. Mount. (1963). Some effects of alkyl benzene sulfonate on the bluegill, *Lepomis macrochinus*. Trans. Amer. Fish. Soc. 92(4): 372-378.

Lennon, R. E. (1967). Selected strains of fish as bioassay animals. Progr. Fish-Cult. 29:129-132.

Lennon, R. E. and C. R. Walker. (1964). Laboratories and methods for screening fish-control chemicals. In: Investigations in Fish Control. No. 1. Washington, D.C. (GPO), U.S. Fish Wildl. Serv., Bur. Sport Fish. and Wildl., Cir. 185.

Leonard, J. W. (1938). Notes on the use of derris as a fish poison. Trans. Amer. Fish. Soc. 68: 269-280.

Leonard, J. W. (1965). Environmental requirements of Ephemeroptera. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 110-117.

Lewallen, L. L. (1959). Toxicity of several organophosphorus insecticides to *Gambusia affinis* (Baird and Girard) in laboratory tests. Mosquito News 19(1): 1-2.

Lewallen, L. L. and W. H. Wilder. (1962). Toxicity of certain organophosphorus and carbamate insecticides to rainbow trout. Mosquito News 22: 369-372.

Lewallen, L. L. and W. H. Wilder. (1963). Laboratory tests of insecticides on mosquito larvae in polluted and tap water. J. Econ. Entomol. 56: 834-835.

Lewis, W. M. (1968). Isobornyl thiocyanoacetate as a fish drugging agent and selective toxin. Progr. Fish-Cult. 30: 29-31.

Libby, R. W. (1964). Methods and facilities for conducting water quality studies in the Great Lakes Conf. Great Lakes Res., Proc. 7: 100-109.

Lichtenstein, E. P., K. R. Schulz, R. F. Skrentny, and Y. Tsukano. (1966). Toxicity and fate of insecticide residues in water. Arch. Environ. Health 12: 199-212.

Lindgren, P. E. (1960). About the effect of rotenone upon benthonic animals in lakes. Inst. Freshwater Res., Drottningholm, Rept. No. 41, pp. 172-184.

Linduska, J. P. and E. W. Surber. (1948). Effects of DDT and other insecticides on fish and wildlife. Summary of investigations during 1947. U.S. Fish Wildl. Serv., Circ. 15, 19 p.

Lipschuetz, M. and A. L. Cooper. (1955). Comparative toxicities of potassium cyanide and potassium cuprocyanide to the western black-nosed dace. (*Rhinichthys atratulus meleagris*). N.Y. Fish Game J. 2: 194-204.

Lloyd, R. (1960). The toxicity of zinc sulphate to rainbow trout. Ann. Appl. Biol. 48(1): 84-94.

Lloyd, R. (1961). Effect of dissolved oxygen concentrations on the toxicity of several poisons to rainbow trout (Salmo gairdnerii Richardson). J. Exp. Biol. 38: 447-455.

Lloyd, R. (1961). The toxicity of mixtures of zinc and copper sulphates to rainbow trout (*Salmo gairdnerii* Richardson). Ann. Appl. Biol. 49: 535-538.

Lloyd, R. (1964). Sewage effluents and their effects on fisheries. Ann. Appl. Biol. 53: 508-509.

Lloyd, R. (1965). Factors that affect the tolerance of fish to heavy metal poisoning. In: Biological Problems in Water Pollution. Third. Seminar, 1962. Washington, D.C. (GPO), Public Health Serv. Publ. No. 999-WP-25, pp. 181-187.

Lloyd, R. and D.W.M. Herbert. (1960). The influence of carbon dioxide on the toxicity of un-ionized ammonia to rainbow trout (*Salmo gairdnerii* Richardson). Ann. Appl. Biol. 48(2): 399-404.

Lloyd, R. and D.H.M. Jordan. (1964). Some factors affecting the resistance of rainbow trout (*Salmo gairdnerii* Richardson) to acid waters. Int. J. Air Water Pollut. 8: 292-403.

Lloyd, R. and D.H.M. Jordan. (1964). Predicted and observed toxicities of several sewage effluents to rainbow trout: A further study. J. Proc. Inst. Sewage Purif. London 2: 183-186.

Loeb, H. A., H. H. Gettner, and H. A. Abramson. (1965). Effect of 48 derivatives of d-lysergic acid on the surfacing behavior of laboratory fish. N.Y. Fish Game J. 12: 79-98.

Loeb, H. A. and P. J. Starkey. (1966). Survival of buried bullheads subjected to 4'-iodo-3-nitrosalicylanilide. N.Y. Fish Game J. 13(2): 196-205. Louder, D. E. and E. G. McCoy. (1965). Preliminary investigations of the use of Aqualin for collecting fishes. Proc. 16th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 240-242.

Lowe, J. I. (1964). Chronic exposure of spot, *Leiostomus xanthurus*, to sublethal concentrations of Toxaphene in sea water. Trans. Amer. Fish. Soc. 93(4): 396-399.

Lowe, J. I. (1966). Some effects of Endrin on estuarine fishes. Proc. 19th Ann. Conf., Southeast. Assoc. Game Fish. Comm., pp. 271-276.

Lowman, F. G. (1965). A control for crayfish. Progr. Fish-Cult. 27: 184.

Ludke, J. L., D. E. Ferguson, and W. D. Burke. (1968). Some Endrin relationships in resistant and susceptible populations of golden shiners, *Notemigonus crysoleucas*. Trans. Amer. Fish Soc. 97(3): 260-263.

Ludzack, F. J. and M. B. Ettinger. (1963). Estimating biodegradability and treatability of organic water pollutants. Biotechnol. Bioeng. 5: 309-330.

Macek, K. J. (1968). Reproduction in brook trout (Salvelinus fontinalis) fed sublethal concentrations of DDT. J. Fish. Res. Bd. Can. 25(9): 1787-1796.

Macek, K. J., C. Hutchinson, and O. B. Cope. (1969). The effects of temperature on the susceptibility of bluegills and rainbow trout to selected pesticides. Bull. Environ. Contam. Toxicol. 4(3): 174-184.

Mack, G. L., S. M. Corcoran, S. D. Gibbs, W. H. Gutenmann, J. S. Reckahn, and D. J. Lisk. (1964). The DDT content of some fishes and surface waters of New York State. N.Y. Fish Game J. 11: 148-153.

Mackay, R. J. (1969). Aquatic insect communities of a small stream on Mont. St. Hilaire, Quebec. J. Fish. Res. Bd. Can. 26(5): 1157-1183.

Mackenthun, K. M. (1962). A review of algae, lake weeds, and nutrients. J. Water Pollut. Contr. Fed. 34(10): 1077-1085.

Mackenthun, K. M. (1965). Nitrogen and phosphorus in water, an annotated selected bibliography of their biological effects. U.S. Public Health Serv. Publ. No. 1305, 111 p.

Mackenthun, K. M. (1966). Biological evaluation of polluted streams. J. Water Pollut. Contr. Fed. 38(2): 241-247.

Mackenthun, K. M. and H. L. Cooley. (1952). The biological effect of copper sulfate treatment on lake ecology. Trans. Wis. Acad. Sci. Arts Lett. 41: 177-187.

Mackenthun, K. M. and W. M. Ingram. (1964). Limnological aspects of recreational lakes. U.S. Public Health Serv. Publ. No. 1167, 176 p.

Mackenthun, K. M. and W. M. Ingram. (1964). Pollution and the life in water. U.S. Public Health Serv. Publ. No. 999-WP-20, 16 p. Mackenthun, K. M. and W. M. Ingram. (1967). Biological associated problems in freshwater environments, their identification, investigation and control. Washington, D.C. (GPO), U.S. Dept. Int., Fed. Water Pollut. Contr. Admin., 287 pp.

Mackereth, F. J. and W.J.P. Smyly. (1951). Toxicity of copper in solution to the stone-loach. Nature 168: 1130.

MacLeod, J. C. and L. L. Smith. (1966). Effect of pulpwood fiber on oxygen consumption and swimming endurance of the fathead minnow, *Pimephales promelas*. Trans. Amer. Fish. Soc. 95: 71-84.

MacMullan, R. A. (1968). The case against hard pesticides. Mich. Conserv. Dept., 6 p. [Repr. from Mich. Conserv., Jan-Feb.].

MacPhee, C. and R. Pruelle. (1968). Fish culture by squawfish population eradication. U.S. Patent 3,389,685, 5 p.

Mahdi, M. A. (1966). Mortality of some species of fish to Toxaphene at three temperatures. In: Investigations in fish control. No. 6. U.S. Fish Wildl. Serv., Bur. Sport Fish. and Wildl. Resour. Publ. 10, 10 p.

Malaney, G. W., W. D. Sheets, and R. Quillin. (1959). Toxic effects of metallic ions on sewage microorganisms. Sewage Ind. Wastes 31(11): 1309-1315.

Malina, J. F. (1964). Toxicity of petrochemical in the aquatic environment. Water Sewage Works 111(10): 456-560.

Maloney, T. E. (1966). Detergent phosphorus effect on algae. J. Water Pollut. Contr. Fed. 38(1): 38-45.

Maloney, T. E. and C. M. Palmer. (1956). Toxicity of six chemical compounds to thirty cultures of algae. Water Sewage Works 103: 509-513.

Mann, K. H. (1965). Heated effluents and their effects on the invertebrate fauna of rivers. Proc. Soc. Water Treat. Exam. 14: 45-53.

Manufacturing Chemists Association. Water Resources Committee. 1967. Source materials on water quality criteria. (Rev.) Washington, D.C., Manufacturing Chemists Association, Water Resources Committee, 35 p.

Marchetti, R. (1965). The toxicity of nonyl phenol ethoxylate to the developmental stages of the rainbow trout, *Salmo gairdnerii*, Richardson, Ann. Appl. Biol. 55: 425-430.

Marking, L. L. (1966). Evaluation of p,p'-DDT as a reference toxicant in bioassays. In: Investigations in fish control. No. 10. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 14, 10 p.

Marking, L. L. (1967). Toxicity of MS-222 to selected fishes. In: Investigations in Fish Control. No. 12. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 18, 10 p.

Marking, L. L. and J. W. Hogan. (1967). Toxicity of Bayer 73 to fish. In: Investigations in Fish Control. No. 19. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl. Resour. Publ. 36, 13 p.

Martin, W. E. (n.d.). Fish-pesticide acute toxicity test method. U.S. Dep. Agr. Res. Serv., Pestic. Regul. Div., Animal Biol. Sect. (Fish), 11 p. (unpubl.)

Mason, W. T., J. B. Anderson, and G. E. Morrison. (1967). A limestone-filled, artificial substrate sampler-float unit for collecting macroinvertebrates in large streams. Progr. Fish-Cult. 29: 74.

Mathur, D. S. (1963). Observations of the toxicity of Dieldrin and Lindane to certain fishes. Zool. Pol. 13: 99-108.

Mathur, D. S. (1964). Determination of the accumulation of DDT and BHC in the liver and intestine of certain fishes. Proc. Nat. Acad. Sci. India Sect. B, Biol. Sci. 34(4): 337-338.

Mawdesley-Thomas, L. E. and J. S. Leahy. (1967). Organochlorine pesticide residues in pike. Progr. Fish-Cult. 29: 64.

McCall, G. W. (1961). Bioassay of an industrial waste. Sanitalk 9: 18-20.

McCauley, R. N. (1966). The biological effects of oil pollution in a river. Limnol. Oceanogr. 11: 475-486.

McDonald, S. (1962). Rapid detection of chlorinated hydrocarbon insecticides in aqueous suspension with *Gammarus lacustris lacustris* (Sars.). Can. J. Zool. 40: 719-723.

McFarland, W. N. (1959). A study of the effects of anaesthetics on the behavior and physiology of fishes. Publ. Inst. Mar. Sci. 6: 23-55.

McKee, J. E. and H. W. Wolf (eds.). (1963). Water quality criteria. 2d ed. Sacramento, Calif. State Water Quality Control Board, Resources Agency of California, Publ. No. 3-A, 548 p.

McKeown, J. J. (1962). The control of *Sphaerotilus natans* by a southern Kraft mill. Proc. 17th Ind. Waste Conf. Purdue Univ. 47(2): 440-453.

McLean, L. A. (1968). DDT residues and Bermuda petrels. Science 161: 397.

Meehan, W. R. and W. L. Sheridan. (1966). Effects of Toxaphene on fishes and bottom fauna of Big Kitoi Creek, Afognak Island, Alaska. In: Investigations in fish control. No. 8. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 12, 9 p.

Meeks, R. L. (1968). The accumulation of <sup>36</sup>Cl ring-labeled DDT in a freshwater marsh. J. Wildl. Manage. 32(2): 376-398.

Merkens, J. C. (1957). Controlled aqueous environments for bioassay. Lab. Pract. 6(8): 456-459, 471.

Merkens, J. C. (1958). Studies on the toxicity of chlorine and chloramines to the rainbow trout. Water Waste Treat. J. 7: 150-151.

Merkens, J. C. and K. M. Downing. (1957). The effect of tension of dissolved oxygen on the toxicity of un-ionized ammonia to several species of fish. Ann. Appl. Biol. 45(3): 521-527.

Meyer, F. P. (1965). The experimental use of Guthion as a selective fish eradicator. Trans. Amer. Fish. Soc. 94: 203-209.

Meyer, F. P. (1965). The effect of formulation differences on the toxicity of benzene hexachloride to golden shiners. Proc. 17th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 186-190.

Mihursky, J. A. (1962). Patuxent River study with special reference to the effects of heated steam electric station condenser water upon estuarine ecology. Prince Frederick, Md., Natur. Res. Inst., 29 p.

Mihursky, J. A. and V. S. Kennedy. (1967). Water temperature criteria to protect aquatic life. Amer. Fish. Soc., Spec. Publ. 4, pp. 20-32.

Miller, C. W., B. M. Zuckerman, and A. J. Charig. (1966). Water translocation of Diazinon- $C^{14}$  and Parathion- $S^{35}$  of a model cranberry bog and subsequent occurrence in fish and mussels. Trans. Amer. Fish. Soc. 95: 345-349.

Mitchum, D. L. and T. D. Moore. (1967). Study of water pollution problems which affect fish and other aquatic forms. Wyoming Game and Fish Comm., Federal Aid in Fish and Wildl. Restoration, Project No. FW-3-R-13, 32 p.

Mitrovic, V. V., V. M. Brown, D. G. Shurben, and M. H. Berryman. (1968). Some pathological effects of sub-acute and acute poisoning of rainbow trout by phenol in hard water. Water Res. 2: 249-254.

Montgomery, H.A.C. (1967). The determination of biochemical oxygen demand by respirometric methods. Water Res. 1: 631-662.

Moore, N. W. (1967). A synopsis of the pesticide problem. Advan. Ecol. Res. 4: 75-129.

Moore, N. W. and J. O. Tatton. (1965). Organochlorine insecticide residues in the eggs of sea birds. Nature 207: 42-43.

Moore, S. L. and S. R. Kin. (1969). Train wreck causes cyanide pollution. Water Sewage Works 116: 35-40.

Morrill, J. B. (1963). Morphological effects of cobaltous chloride on the development of *Limnaea stagnalis* and *Limnaea palustris*. Biol. Bull. 125(3): 508-522.

Morrison, S. M. and J. F. Fair. (1966). Influence of environment on stream microbial dynamics. Ft. Collins, Colorado State Univ., Hydrology Pap. No. 13, 21 p. Moubry, R. J., J. M. Helm, and G. R. Myrdal. (1968). Chlorinated pesticide residues in an aquatic environment located adjacent to a commercial orchard. Pesticides Monit. J. 1(4): 27-29.

Mount, D. I. (1962). Chronic effects of Endrin on bluntnose minnows and guppies. U.S. Fish and Wildl. Serv., Bur. Sport Fish. and Wildl. Res. Rept. 58, 38 p.

Mount, D. I. (1964). An autopsy technique for zinc-caused fish mortality. Trans. Amer. Fish. Soc. 93(2): 174-182.

Mount, D. I. (1966). The effect of total hardness and pH on acute toxicity of zinc to fish. Int. J. Air Water Pollut. 10: 49-56.

Mount, D. I. (1967). Considerations for acceptable concentrations of pesticides for fish production. Amer. Fish. Soc., Spec. Publ. 4, pp. 3-6.

Mount, D. I. (1968). Test animals for water quality. Amer. Fish. Soc. Newslett. 12(54): 3, 6.

Mount, D. I. (1968). Chronic toxicity of copper to fathead minnows (*Pimephales promelas*, Rafinesque). Water Res. 2: 215-223.

Mount, D. I. and W. A. Brungs. (1967). A simplified dosing apparatus for fish toxicology studies. Water Res. 1: 21-29.

Mount, D. I. and C. E. Stephan. (1967). A method for detecting cadmium poisoning in fish. J. Wildl. Manage. 31(1): 168-172.

Mount, D. I. and C. E. Stephan. (1967). A method for establishing acceptable toxicant limits for fish – Malathion and the butoxyethanol ester of 2,4-D. Trans. Amer. Fish. Soc. 96(2): 185-193.

Mount, D. I., L. W. Vigor, and M. L. Schafer. (1966). Endrin: Use of concentration in blood to diagnose acute toxicity to fish. Science 152: 1388-1390.

Mount, D. I. and R. E. Warner. (1965). A serial-dilution apparatus for continuous delivery of various concentrations of materials in water. U.S. Public Health Serv. Publ. No. 999-WP-23, 16 p.

Moye, W. C. and W. H. Luckmann. (1964). Fluctuations in populations of certain aquatic insects following application of Aldrin granules to Sugar Creek, Iroquois County, Illinois. J. Econ. Entomol. 57(3): 318-322.

Mulla, M. S. (1963). Toxicity of organochlorine insecticides to the mosquito fish *Gambusia affinis* and the bullfrog *Rana catesbeiana*. Mosquito News 23: 299-303.

Mulla, M. S. (1966). Toxicity of new organic insecticides to mosquito fish and some other aquatic organisms. Mosquito News 26: 87-91.

Mulla, M. S. and L. W. Isaak. (1961). Field studies on the toxicity of insecticides to the mosquito fish, *Gambusia affinis*. J. Econ. Entomol. 54: 1273-1242.

Mulla, M. S., L. W. Isaak, and H. Axelrod. (1963). Field studies on the effects of insecticides on some aquatic wild-life species. J. Econ. Entomol. 56: 184-188.

Mulla, M. S., J. St. Amant, and L. D. Anderson. (1967). Evaluation of organic pesticides for possible use as fish toxicants. Progr. Fish-Cult. 29: 36-42.

Muncy, R. J. and A. D. Oliver. (1963). Toxicity of ten insecticides to the red crawfish, *Procambarus clarki* (Girard). Trans. Amer. Fish. Soc. 92(4): 428-431.

Murphy, S. D. (1966). Liver metabolism and toxicity of thiophosphate insecticides in mammalian, avian, and piscine species. Proc. Soc. Exp. Biol. Med. 123: 392-398.

Murphy, S. D., R. R. Lauwerys, and K. L. Cheever. (1968). Comparative anticholinesterase action of organophosphorus insecticides in vertebrates. Toxicol. Appl. Pharm. 12: 22-35.

National Academy of Sciences-National Research Council. Committee on Pollution. (1966). Waste management and control. A report to the Federal Council for Science and Technology. Nat. Acad. Sci.-Nat. Resour. Counc. 257 p.

National Technical Advisory Committee. (1967). Interim Report on Water Quality Criteria, Federal Water Pollution Control Administration, 766 p.

National Technical Advisory Committee. (1968). Water quality criteria. Report to the Secretary of the Interior. Washington, D.C., Federal Water Pollution Control Administration, 234 p.

Naylor, E. (1965). Biological effects of a heated effluent in docks at Swansea, S. Wales. Proc. Zool. Soc. London 144: 253-268.

Nebeker, A. V. and A. R. Gaufin. (1964). Bioassays to determine pesticide toxicity to the amphipod crustacean, *Gammarus lacustris*. Proc. Utah Acad. Sci. 41(1): 64-67.

Needham, R. G. (1966). Effects of Toxaphene on plankton and aquatic invertebrates in North Dakota lakes. In: Investigations in fish control. No. 4. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 8, 16 p.

Neel, J. K. (1963). Industrial wastes and pesticides – avenues through which they affect water quality and aquatic life. Trans. Kans. Acad. Sci. 66: 42-48.

Nejedly, A. (1967). An explanation of the difference between the rate of the BOD progression under laboratory and stream conditions. In: Advances in Water Pollution Research, Proc. 3rd International Conference held in Munich, Germany, Sept. 1966. Vol. 1. Washington, D.C., Water Pollut. Contr. Fed., pp. 23-53.

Neuhold, J. M. and W. F. Sigler. (1962). Chlorides affect the toxicity of fluorides to rainbow trout. Science 135: 732-733.

Newsom, L. D. (1967). Consequences of insecticide use on nontarget organisms. Ann. Rev. Entomol. 12: 257-286.

Nicholson, H. P. (1959). Insecticide pollution of water resources. J. Amer. Water Works Assoc. 51(8): 981-986.

Nicholson, H. P. (1967). Pesticide pollution control. Science 158: 871-876.

Nicholson, H. P., A. R. Crzenda, G. J. Lauer, W. S. Cox, and J. I. Teasley. (1964). Water pollution by insecticides in an agricultural river basin. I. Occurrence of insecticides in river and treated municipal water. Limnol. Oceanogr. 9: 310-317.

Nicholson, H. P. and J. R. Thoman. (1964). The problem of pesticides as pollutants of water. Mosquito News 24: 169-172.

Nightingale, H. W. and V. L. Loosanoff. (1928). The effects of waste sulphite liquor on the early stages of chinook salmon, and means of prevention by disposal methods. Trans. Amer. Fish. Soc. 58: 232-244.

Norris, L. A. (1967). Chemical brush control and herbicide residues in the forest environment. In: Symposium Proceedings: Herbicides and Vegetation Management in Forests, Ranges, and Noncrop Lands. Oregon State University, pp. 103-123.

North, W. J., K. A. Clendenning, and H. L. Scotten. (1960). The effects of waste discharges on kelp. Quarterly Progress Report 1 July 1959 - 30 September 1959. San Diego, Univ. of Calif., Inst. Mar. Resour., IMR Ref. 60-4, 54 p.

Oberton, A.C.E. and V. T. Stack. (1957). Biochemical oxygen demand of organic chemicals. Sewage Ind. Wastes 29(11): 1267-1272.

O'Connell, R. L. and N. A. Thomas. (1965). Effect of benthic algae on stream dissolved oxygen. J. San. Eng. Div., Proc. Amer. Soc. Civil Eng. 91(SA3): 1-16.

Odum, E. P. (1959). Fundamentals of ecology. 2nd ed. Philadelphia, W. B. Saunders Co., 546 p.

Odum, E. P. and W. T. Sumerford. (1946). Comparative toxicity of DDT and four analogues to goldfish, *Gambusia*, and *Culex* larvae. Science 104(2708): 480-482.

Odum, W. E., G. M. Woodwell, and G. F Winster. (1969). DDT residues absorbed from organic detritus by fiddler crabs. Science 164: 576-577.

Ogilvie, D. M. and J. M. Anderson. (1965). Effect of DDT on temperature selection by young Atlantic salmon, *Salmo salar*. J. Fish. Res. Bd. Can. 22: 503-512.

Oglesby, L. C. (1964). Mortality of a freshwater polychaete, *Nereis limnicola* Johnson, attributed to rotenone. Calif. Fish Game 50: 268-270.

Ohio River Valley Water Sanitation Commission. (1956). Aquatic life water quality criteria. 2d Progress Report. Sewage Ind. Waste 28(5): 678-690.

Okum, D. A. (1968). The hierarchy of water quality. Environ. Sci. Tech. 2(9): 672-675. Olsen, P. A. and R. F. Foster. (1958). Effect of Separan on rainbow trout. Richland, Wash., General Electric Co., HW-55292, 5 p.

Olson. T. A. (1957). Biological indicators in stream ecology. Proc. 11th Ind. Waste Conf. Purdue Univ. 41(2): 601-610.

Olson, T. A. and F. J. Burgess, (eds.). (1967). Pollution and marine ecology. N.Y., Interscience Publ. 364 p.

Pagan, C. and R. H. Hageman. (1950). Determination of DDT by bioassay. Science 112: 222-223.

Paine, G. H. and A. R. Gaufin. (1956). Aquatic diptera as indicators of pollution in a midwestern stream. Ohio J. Sci. 56: 291-304.

Palmer, C. M. (1959). Algae in water supplies. U.S. Public Health Serv. Publ. No. 657, 88 p.

Palmer, C. M. (1963). The effect of pollution on river algae. Ann. N.Y. Acad. Sci. 108: 389-395.

Palmer, C. M. and T. E. Maloney. (1955). Preliminary screening for potential algicides. Ohio J. Sci. 55(1): 1-8.

Parka, S. J. and H. M. Worth. (1965). Effects of Trifluralin on fish. Proc. S. Weed Conf. 18: 469-474.

Parkhurst, Z. E. and H. E. Johnson. (1955). Toxicity of Malathion 500 to fall chinook salmon fingerlings. Progr. Fish-Cult. 17: 113-116.

Patrick, R. (1957). Diatoms as indicators of changes in environmental conditions. In: Biol. Probl. Water Pollut., Trans. of the 1956 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, pp. 71-83.

Patrick, R. (1965). Algae as indicators of pollution. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 225-231.

Patrick, R. (1968). Effect of suspended solids, organic matter and toxic materials on aquatic life in rivers. Water Sewage Works 115: 89-92.

Patrick, R., J. Cairns, and S. S. Roback. (1967). An ecosystematic study of the fauna and flora of the Savannah River. Proc. Acad. Natur. Sci. Philadelphia 118(5): 109-407.

Patrick, R., J. Cairns, and A. Scheier. (1968). The relative sensitivity of diatoms, snails, and fish to twenty common constituents of industrial wastes. Progr. Fish-Cult. 30(3): 137-140.

Patten, B. G. and C. C. Gillaspie. (1966). The Bureau of Commercial Fisheries type IV electro fishing shocker-its characteristics and operations. U.S. Fish Wildl. Serv., Spec. Sci. Report-Fish., No. 529, 15 p.

Patterson, R. S. and D. L. Von Windeguth. (1964). The effects of Baytex on some aquatic organisms. Mosquito News 24: 46-49.

Phillippy, C. L. (1961). Preliminary results of herbicides tested on certain aquatic plants in Florida, Proc. 15th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 288-295.

Phillips, A. M. (1944). The physiological effect of sodium chloride on brook trout. Trans. Amer. Fish. Soc. 74: 297-309.

Phillipson, J. (1966). Ecological Energetics. N.Y., St. Martins Press, 57 p.

Piavis, G. W. (1962). Exposure of several developmental stages of the sea lamprey, *Petromyzon marinus*, to selective larvicides. Copeia, No. 3: 652-653.

Pickering, Q. H. (1966). Acute toxicity of alkyl benzene sulfonate and linear alkylate sulfonate the eggs of the fathead minnow, *Pimephales promelas*. Int. J. Air Water Pollut. 10: 385-391.

Pickering. Q. H. (1968). Some effects of dissolved oxygen concentrations upon the toxicity of zinc to the bluegill, *Lepomis macrochirus*, Raf. Water Res. 2: 187-194.

Pickering, Q. H. and C. Henderson. (1959). Monitoring for BW and CW Agents in Water With Fish. In: R. L. Woodward and G. G. Robeck, Removal of Radiological, Biological, and Chemical Contaminants From Water, Cincinnati, Ohio, Robert A. Taft San. Eng. Ctr., Tech. Rpt. W59-2, pp. 119-133.

Pickering, Q. H. and C. Henderson. (1965). The acute toxicity of some heavy metals to different species of warm water fishes. In: Proc. 19th Ind. Waste Conf. Purdue Univ. 49(2): 578-591.

Pickering, Q. H. and C. Henderson. (1966). The acute toxicity of some pesticides to fish. Ohio. J. Sci. 66(5): 508-513.

Pickering, Q. H. and C. Henderson. (1966). Acute toxicity of some important petrochemicals to fish. J. Water Pollut. Contr. Fed. 38(9): 1419-1429.

Pickering, Q. H., C. Henderson, and A. E. Lemke. (1962). The toxicity of organic phosphorus insecticides to different species of warm-water fishes. Trans. Amer. Fish. Soc. 91: 175-184.

Pickering, Q. H. and W. N. Vigor. (1965). The acute toxicity of zinc to eggs and fry of the fathead minnow. Progr. Fish-Cult. 27(3): 153-157.

Pomelle, C. S. (1953). Toxicity of beryllium. Sewage Ind. Wastes 25: 1424-1428.

Portmann, J. E. and P. M. Connor. (1968). The toxicity of several oil-spill removers to some species of fish and shell-fish. Marine Biol. (Berlin) 1(4): 322-329.

Post, G. (1959). A preliminary report on the use of Nitrofuran compounds for furunculosis of trout, with special emphasis on Furoxone. Progr. Fish-Cult. 21: 30-33.

Post, G. and R. E. Keiss. (1962). Further laboratory studies on the use of Furazolidone for the control of furunculosis of trout. Progr. Fish-Cult. 24: 16-21.

Powers, E. B. (1918). The goldfish (*Carassius carassius*) as a test animal in the study of toxicity. Ill. Biol. Monogr. 4(2): 123-193.

President's Science Advisory Committee. Environmental Pollution Panel. (1965). Restoring the quality of our environment. Report of the Environmental Pollution Panel, President's Science Advisory Committee, Washington, D.C., The White House, 317 p.

Proffitt, M. A. (1966). Some factors affecting the toxicity of Aldrin to fishes. Proc. Indiana Acad. Sci. 75: 325-329.

Prosser, C. L. and F. A. Brown. (1961). Comparative animal physiology. 2nd ed. Philadelphia, W. B. Saunders Co., 688 p.

Pyatt, E. E. (1964). On determining pollutant distribution in tidal estuaries. Hydrology of tidal streams. U.S. Geol. Surv., Water-Supply Pap. 1586-F, 56 p.

Rabinowitz, J. L. and R. M. Myerson. (1966). Exposure of aquarium fish to dimethyl sulfoxide (DMSO) with special reference to toxicity and effects on uptake of radioactive dyes. Proc. Soc. Exp. Biol. Med. 121: 1065-1067.

Rachlin, J. W. and A. Perlmutter. (1968). Fish cells in cultures for the study of aquatic toxicants. Water Res. 2: 409-414.

Randall, C. W. (1966). Toxicity of organophosphate insecticides to freshwater microorganisms. In: K. L. Bowden (ed.), Proceedings of the 2nd Annual American Water Resources Conference at Univ. of Chicago, Center for Continuing Education, Chicago, Illinois, November 21-22, 1966. Urbana, Illinois. American Water Resources Association, Proc. Ser. No. 2, pp. 352-364.

Raney, E. C. and B. W. Menzel. (1967). A bibliography: Heated effluents and effects on aquatic life with emphasis on fishes. Philadelphia Electric Company and Ichthyological Associates, Bull. No. 1, 90 p.

Rao, T. S., S. Dutt, and K. Mangaiah. (1967).  $TL_m$  values of some modern pesticides to the fresh-water fish – *Puntius puckelli*. Environ. Health 9(2): 103-109.

Raymont, J.E.G. and J. Shields. (1964). Toxicity of copper and chromium in the marine environment. In: Advances in Water Pollution Research, Proc. International Conference held in London, Sept. 1962. Vol. 3. N.Y., MacMillan, pp. 275-290.

Reed, R. J. (1966). Some effects of DDT on the ecology of salmon streams in Southeastern Alaska. U.S. Fish Wildl. Serv., Spec. Sci. Rept.-Fish. 542, 15 p.

Reich, K. (1955). The effect of cyanide and azide on the respiration of the amoeba, *Mayorella palestinensis*. Physiol. Zool. 28: 145-151.

Reid, G. K. (1961). Ecology of inland waters and estuaries. N.Y., Reinhold Publ. Co. 375 pp.

Reiff, B. (1964). Factors influencing the testing of chemicals and effluents for toxicity to fish. J. Sci. Technol. 10(4): 167-171.

Renn, C. E. (1955). Biological properties and behaviors of cyanogenic wastes. Sewage Ind. Wastes 27: 297-310.

Reymonds, T. D. (1962). Pollutional effects of agricultural insecticides and synthetic detergents. Water Sewage Works 109: 352-355.

Ricker, W. E. (ed.). (1968). Methods for assessment of fish production in fresh waters. Oxford, Blackwell Sci. Publ. 313 p. (IBP Handbook No. 3.)

Roback, S. S. (1965). Environmental requirements of Trichoptera. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 118-126.

Roberts, H. (1963). Cadmium toxic to rainbow trout. Progr. Fish-Cult. 25: 216.

Robinson, J. (1967). Dynamics of organochlorine insecticides in vertebrates and ecosystems. Nature 215: 33-35.

Robinson, J., A. Richardson, A. N. Crabtree, J. C. Coulson, and G. R. Potts. (1967). Organochlorine residues in marine organisms. Nature 214: 1307-1311.

Rodgers, E. O., B. H. Hazen, S. B. Friddle, and S. F. Sneiszko. (1951). The toxicity of pyridylmercuric acetate technical (PMA) to rainbow trout (*Salmo gairdnerri*). Progr. Fish-Cult. 13(2): 71-73.

Rosato, P. and D. E. Ferguson. (1968). The toxicity of Endrin-resistant mosquitofish to eleven species of vertebrates. BioScience 18(8): 783-784.

Rose, E. T. (1958). Further notes on toxaphene in fish population control. Iowa State Conserv. Comm., Fish Game Div., Quart. Biol. Rept. 10(2): 5-7.

Royer, L. M. (1966). Bioassay method for the determination of Toxaphene in lake water. J. Fish. Res. Bd. Can. 23: 723-727.

Rudolfs, W., G. E. Barnes, G. P. Edwards, H. Heukelekian, E. Hurwitz, C. E. Renn, S. Steinberg and W. F. Vaughan. (1950). Review of literature on toxic materials affecting sewage treatment processes, streams, and B.O.D. determinations. Sewage Ind. Wastes 22:1157-1191.

Ruttner, F. (1953). Fundamentals of limnology. Univ. Toronto Press. 242 p.

Ryckman, D. W., A.V.S. Prabhakara-Rao, and J. C. Buzzell. (1966). Behavior of organic chemicals in the aquatic environment; a literature critique. Research report from the Environmental and Sanitary Engineering Laboratories of Washington University, St. Louis, for the Manufacturing Chemists Association. Washington, D.C. Manufacturing Chemists Association, 164 p.

Sanborn, N. H. (1945). The lethal effect of certain chemicals on freshwater fish. Canner 101(5): 13.

Sanders, H. O. and O. B. Cope. (1966). Toxicities of several pesticides to two species of cladocerans. Trans. Amer. Fish. Soc. 95: 165-169.

Sanders, H. O. and O. B. Cope. (1968). The relative toxicities of several pesticides to naiads of three species of stoneflies. Limnol. Oceanogr. 13(1): 112-117.

Sanders, R. L. and J. B. Sprague. (1967). Effects of copperzinc mining pollution on a spawning migration of Atlantic salmon. Water Res. 1: 419-432.

Schaumburg, F. D., T. E. Howard, and C. C. Walden. (1967). A method to evaluate the effects of water pollutants on fish respiration. Water Res. 1(10): 731-737.

Scheier, A. and J. Cairns. (1967). Persistence of gill damage in *Lepomis gibbosus* following a brief exposure to alkyl benzene sulfonate. Notulae Natur., No. 391, 7 p.

Scheier, A. and J. Cairns. (1968). An apparatus for estimating the effects of toxicants on the critical flicker frequency response of the bluegill sunfish. Proc. 23rd Ind. Waste Conf. Purdue Univ. 10 p.

Schiffman, R. H. and P. O. Fromm. (1959). Chromiuminduced changes in the blood of rainbow trout, *Salmo* gairdnerii. Sewage Ind. Wastes 31(2): 205-211.

Schmid, O. J. and H. Mann. (1961). Action of a detergent (dodecylbenzene-sulfonate) on the gills of the trout. Nature 192: 675.

Schmidt, C. H. and D. E. Weidhaas. (1961). The toxicological action of three organophosphorus insecticides with three species of mosquito larvae. J. Econ. Entomol. 54: 583-586.

Schoenthal, N. D. (1963). Some effects of DDT on cold water fish and fish-food organisms. Proc. Montana Acad. Sci. 23: 63-95.

Schoettger, R. A. (1967). Annotated Bibliography on MS-222. In: Investigations in Fish Control. No. 16, Washington, D.C., U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 22, 15 p.

Schoettger, R. A. and A. M. Julin. (1967). Efficacy of MS-222 as an anesthetic on four salmonids. In: Investigations in Fish Control. No. 13. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 19, 15 p.

Schoettger, R. A. and J. R. Olive. (1961). Accumulation of Toxaphene by fish-food organisms. Limnol. Oceanogr. 6: 216-219.

Schoettger, R. A., C. R. Walker, L. L. Marking, and A. M. Julin. (1967). MS-222 as an anesthetic for channel catfish: Its toxicity, efficacy, and muscle residues. In: Investigations in Fish Control. No. 17. Washington, U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 33, 14 p.

Schouwenberg, W. J. and K. J. Jackson. (1966). A field assessment of the effects of spraying a small coastal coho salmon stream with Phosphamidon. Can. Fish Cult. 37: 35-43.

Scott, D. C. (1958). Biological balance in streams. Sewage Ind. Wastes 30(9): 1169-1173.

Seiffer, E. A. and H. F. Schoof. (1967). Tests of 15 experimental molluscicides against *Australorbis glabratus*. Public Health Report 82(9): 833-839.

Servizi, J. A., E. T. Stone, and R. W. Gordon. (1966). Toxicity and treatment of Kraft pulp bleach plant waste. New Westminster, B.C., Int. Pac. Salmon Fish. Comm., Progr. Rept. No. 13, 34 p.

Shane, M. S. (1948). Effect of DDT spray on reservoir biological balance. J. Amer. Water Works Assoc. 40: 333-336.

Shaw, W.H.R. and B. Grushkin. (1967). The toxicity of metal ions to aquatic organisms. Arch. Biochem. Biophys. 67(2): 447-452.

Sheets, W. D. (1957). Toxicity studies of metal-finishing wastes. Sewage Ind. Wastes 29: 1380-1384.

Shrivastava, H. M. (1962). Oligochaetes as indicators of pollution. Water Sewage Works 109: 387-890.

Sigler, W. F., W. T. Helm, J. W. Angelovic, D. W. Linn, and S. S. Martin. (1966). The effects of uranium mill wastes on stream biota. Utah Agr. Exp. Sta., Bull. 462, 76 p.

Skidmore, J. F. (1964). Toxicity of zinc compounds to aquatic animals with special reference to fish. Quart. Rev. Biol. 39(3): 227-248.

Skidmore, J. F. (1965). Resistance to zinc sulphate of the zebrafish (*Brachydanio rerio* Hamilton-Buchanan) at different phases of its life history. Ann. Appl. Biol. 56: 47-53.

Skidmore, J. F. (1966). Resistance to zinc sulphate of zebrafish (*Brachy danio rerio*) embryos after removal or rupture of the outer egg membrane. J. Fish. Res. Bd. Can. 23: 1037-1041.

Skidmore, J. F. (1967). Oxygen uptake by zebrafish (*Brachydanio rerio*) of different ages in relation to zinc sulphate resistance. J. Fish. Res. Bd. Can. 24(6): 1253-1267.

Sladen, W.J.L., C. M. Menzie, and W. L. Reichel. (1966). DDT residues in adelie penguins and a crabeater seal from Antarctica. Nature 210(5037): 670-673.

Smith, A. J. (1967). The effect of the lamprey larvicide, 3-trifluoromethyl-4-nitrophenol, on selected aquatic invertebrates. Trans. Amer. Fish. Soc. 96(4): 410-413.

Smith, A. L., J. Layell, and J. C. Grey. (1962). How to predict downstream DO drop. Wastes Eng. 33: 456-458.

Smith, A. O. and B. R. Woodson. (1965). The effects of fluoride on the growth of *Chlorella pyrenoidosa*. Va. J. Sci. 16(1): 1-8.

Smith, G. E. and B. G. Isom. (1967). Investigation of effects of large-scale applications of 2,4-D on aquatic fauna and water quality. Pestic. Monit. J. 1(3): 16-21.

Smith, J. E. (ed.). (1968). 'Torrey Canyon' pollution and marine life. A report by the Plymouth Laboratory of the Marine Biological Association of the United Kingdom. Cambridge, University Press, 196 p.

Smith, J. W. and S. G. Grigoropoulos. (1968). Toxic effects of odorous trace organics. J. Amer. Water Works Assoc. 60: 969-979.

Smith, L. L. (1961). Water quality data and sustained fish harvests. In: Water Quality Measurement and Instrumentation. Proc. of the 1960 Seminar at Cincinnati, Ohio, August 29-31, 1960. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, SEC-TR-W61-2, pp. 97-102.

Smith, L. L. and R. H. Kramer. (1963). Survival of walleye eggs in relation to wood fibers and *Sphaerotilus natans* in the Rainy River, Minnesota. Trans. Amer. Fish. Soc. 92: 220-234.

Smith, L. L. and R. H. Kramer. (1965). Survival of walleye fingerlings in conifer ground wood fiber. Trans. Amer. Fish. Soc. 94: 402-404.

Smith, L. L. and R. H. Kramer. (1965). Some effects of paper fibers in fish eggs and small fish. Proc. 19th Ind. Waste Conf. Purdue Univ. 49(2): 369-378.

Smith, L. L., R. H. Kramer, and J. C. MacLeod. (1965). Effects of pulpwood fibers on fathead minnows and walleye fingerlings. J. Water Pollut. Contr. Fed. 37(1): 130-140.

Smith, L. L., R. H. Kramer, and D. M. Oseid. (1966). Longterm effects of conifer-groundwood paper fiber on walleyes. Trans. Amer. Fish Soc. 95: 60-70.

Smith, M. W. (1939). Copper sulfate and rotenone as fish poisons. Trans. Amer. Fish. Soc. 69: 141-157.

Snow, J. R. (1958). A preliminary report on the comparative testing of some of the newer herbicides. Proc. 11th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 125-132.

Snow, J. R. (1963). Simazine as an algicide for bass ponds. Progr. Fish-Cult. 25: 34-36. Sollman, T. (1949). Correlation of the aquarium goldfish toxicities of some phenols, quinones, and other benzene derivatives with their inhibition of autooxidative reactions. J. Gen. Physiol. 32: 671-679.

Solon, J. M., J. L. Lincer, and J. H. Nair. (1968). A continuous flow, automatic device for short-term toxicity experiments. Trans. Amer. Fish Soc. 97(4): 501-502.

Southwood, T.R.E. (1966). Ecological methods. London, Methuen and and Co., p. 391.

Sparr, B. I., W. G. Appleby, D. M. DeVries, J. V. Osmun, J. M. McBride, and G. L. Foster. (1966). Insecticide residues in waterways from agricultural use. In: Organic Pesticides in the Environment. Advan. Chem. Ser. No. 60: 146-162.

Spiller, D. (1961). A digest of available information on the insecticide Malathion. Advan. Pest Contr. 4: 249-335.

Spindler, J. C. and A. M. Whitney. (1960). Changes in bottom fauna composition and a fish kill resulting from pulp mill wastes. Proc. Montana Acad. Sci. 19: 107-111.

Sprague, J. B. (1964). Lethal concentrations of copper and zinc for young Atlantic salmon. J. Fish. Res. Bd. Can. 21(1): 17-26.

Sprague, J. B. (1964). Highly alkaline water caused by asbestos-cement pipeline. Progr. Fish-Cult. 26: 111-114.

Sprague, J. B. (1964). Avoidance of copper-zinc solutions by young salmon in the laboratory. J. Water Pollut. Contr. Fed. 36: 990-1004.

Sprague, J. B. (1965). Effects of sublethal concentrations of zinc and copper on migration of Atlantic salmon. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 332-333.

Sprague, J. B., P. F. Elson, and R. L. Saunders. (1965). Sublethal copper-zinc pollution in a salmon river – a field and laboratory study. In: Advances in Water Pollution Research, Proc. 2d International Conference held in Tokyo, August 1964. Vol. 1. N. Y., Pergamon Press, pp. 61-82.

Sprague, J. B. and B. A. Ramsay. (1965). Lethal levels of mixed copper-zinc solutions for juvenile salmon. J. Fish. Res. Bd. Can. 22(2): 425-432.

Sproul, O. J. and D. W. Ryckman. (1963). Significant physiological characteristics of organic pollutants. J. Water Pollut. Contr. Fed. 35(9): 1136-1145.

Sreenivasan, A. and G. K. Swaminathan. (1967). Toxicity of six organophosphorus insecticides to fish. Curr. Sci. 36(15): 397-398.

Srivastava, U. S. and S. K. Konar. (1966). DDVP as a selective toxicant for the control of fishes and insects. Progr. Fish-Cult. 28: 235-238.

St. Amant, J. A., W. C. Johnson, and M. J. Whalls. (1964). Aqualin as a fish toxicant. Progr. Fish-Cult. 26: 84-88. Starkey, R. J. and J. H. Howell. (1966). Substituted nitrosalicylanilides: A new class of selectively toxic sea lamprey larvicides. Ann Arbor, Mich. Great Lakes Fishery Commission, Tech. Rept. No. 11, pp. 21-29.

Stewart, J. W. and J. W. Cornick. (1964). Lobster (Homanus americanus) tolerance for tris buffer, sodium fluoride, and sea water extracts of various woods. J. Fish. Res. Bd. Can. 21: 1549-1551.

Stickel, L. F. (1968). Organochlorine pesticides in the environment. U.S. Fish Wildl. Serv., Spec. Sci. Rept., Wildl. No. 119, 32 p.

Stokes, R. M. and P. O. Fromm. (1965). Effects of chromate on glucose transport by the gut of rainbow trout (Pisces). Physiol. Zool. 38: 202-205.

Stringer, G. E. and R. G. McMynn. (1958). Experiments with Toxaphene as fish poison. Can. Fish-Cult. 23: 39-47.

Stroud, R. H. (1967). Water quality criteria to protect aquatic life: A summary. Amer. Fish. Soc., Spec. Publ. 4, pp. 33-37.

Strufe, R. (1968). "Problems and results of residue studies after application of molluscicides." Farbenfabriken Bayer AG, Pharma Wiss. Abteilung, Leverkusen, Germany, 168 p.

Summerfelt, R. C. and W. M. Lewis (1967). Repulsation of green sunfish by certain chemicals. J. Water Pollut. Contr. Fed. 39(12): 2030-2038.

Sundaresan, B. B., E. C. Bovee, D. E. Wilson, and J. B. Lackay. (1965). Effects of wood reduction waste pollution on the microbial ecology of a small creek. J. Water Pollut. Contr. Fed. 37(11): 1536-1544.

Surber, E. W. (1943). Weed control in hard-water ponds with copper sulfate and sodium arsenate. Trans. N. Amer. Wildl. Conf. 8: 132-141.

Surber, E. W. (1959). Cricotopus bicinctus, a midgefly resistant to electroplating wastes. Trans. Amer. Fish. Soc. 88(2): 111-116.

Surber, E. W. and M. H. Everhart. (1950). Biological effects of Nigrosine used for control of weeds in hatchery ponds. Progr. Fish-Cult. 12: 135-140.

Surber, E. W. and D. D. Friddle. (1946). Relative toxicity of suspension and oil formulations of DDT to native fishes in Back Creek, West Virginia. Trans. Amer. Fish. Soc. 76: 315-321.

Surber, E. W. and C. H. Hoffman. (1949). Effects of various concentrations of DDT on several species of fish of different sizes. U.S. Fish Wildl. Serv., Spec. Sci. Rept.-Fish., No. 4(1), 19 p.

Surber, E. W. and O. L. Meehan. (1931). Lethal concentrations of arsenic for certain aquatic organisms. Trans. Amer. Fish. Soc. 61: 225-239.

Surber, E. W. and Q. H. Pickering. (1962). Acute toxicity of Endothal, Diquat, Hyamine, Dalapon, and Silvex to fish. Progr. Fish-Cult. 24: 164-171.

Surber, E. W. and R. A. Taft. (1965). Water quality criteria for freshwater fishes. Proc. 16th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 435-436.

Surber, E. W. and T. O. Thatcher. (1963). Laboratory studies of the effects of alkyl benzene sulfonate (ABS) on aquatic invertebrates. Trans. Amer. Fish. Soc. 92(2): 152-160.

Swift, D. R. (1965). Effect of temperature on mortality and rate of development of the eggs of the Windermere char (*Salvelinus alpinus*). J. Fish. Res. Bd. Can. 22: 913-917.

Swisher, R. D., J. T. O'Rourke, and H. D. Tomlinson. (1964). Fish bioassays of linear alkylate sulfonates (LAS) and intermediate biodegradation products. J. Amer. Oil Chem. Soc. 41: 746-752.

Tagatz, M. E. (1961). Reduced oxygen tolerance and toxicity of petroleum products to juvenile American shad. Chesapeake Sci. 2: 65-71.

Tang, Y. (1961). The use of saponin to control predaceous fishes in shrimp ponds. Progr. Fish-Cult. 23: 43-45.

Tarpley, W. A. (1958). Studies on the use of the brine shrimp *Artemia salina* (Leach) as a test organism for bioassay. J. Econ. Entomol. 51(6): 780-783.

Tarzwell, C. M. (1957). Water quality criteria for aquatic life. In: C. M. Tarzwell (ed. and comp.), Biological Problems in water pollution. Trans. of a Seminar on Biological Problems in Water Pollution, April 23-27, 1956. Cincinnati, Ohio, R. A. Taft San. Eng. Center, pp. 246-272.

Tarzwell, C. M. (1959). Pollutional effects of organic insecticides. Trans. N. Amer. Wildl. Conf. 24: 132-142.

Tarzwell, C. M. (1959). Disposal of toxic wastes. Ind. Wastes 4(5): 136-139.

Tarzwell, C. M. (1959). The toxicity of some organic insecticides to fishes. Proc. 12th Ann. Conf., Southeast Assoc. Game Fish Comm., p. 233-239.

Tarzwell, C. M. (1962). The need and value of water quality criteria with special reference to aquatic life. Can. Fish-Cult. 31: 35-41.

Tarzwell, C. M. (1962). Development of water quality criteria for aquatic life. J. Water Pollut. Contr. Fed. 34(11): 1178-1185.

Tarzwell, C. M. and C. Henderson. (1957). Toxicity of Dieldrin to fish. Trans. Amer. Fish. Soc. 86: 245-257.

Tarzwell, C. M. and C. Henderson. (1960). Toxicity of less common metals to fishes. Ind. Wastes 5: 12.

Tatum, W. M. (1966). Bioassay of industrial pollution by use of masonite plate samplers populated with chironomids. Proc. 19th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 253-258. Tatum, W. M. and R. D. Blackburn. (1965). Preliminary study of the effects of Diquat on the natural bottom fauna and plankton in two subtropical ponds. In: Proc. 16th Ann. Conf. Southeast. Assoc. Game Fish. Comm. 1962, pp. 301-307.

Teeter, J. W. (1965). Effects of sodium chloride on the sago pondweed. J. Wildl. Manage. 29: 838-845.

Terriere, L. C., U. Kiigemai, A. R. Gerlach, and R. L. Borovicka. (1966). The persistence of Toxaphene in lake water and its uptake by aquatic plants and animals. J. Agr. Food Chem. 14: 66-69.

Thatcher, T. O. (1966). The comparative lethal toxicity of a mixture of hard ABS detergent products to eleven species of fishes. Int. J. Air Water Pollut. 10: 585-590.

Thatcher, T. O. and J. F. Santner. (1967). Acute toxicity of LAS to various fish species. Proc. 21st Ind. Waste Conf. Purdue Univ. 50(2): 996-1002.

Thomaston, W. W., P. C. Pierce, and H. N. Wyatt. (1959). Experimental use of Silvex and other aquatic herbicides in Georgia farm ponds. Proc. 13th Ann. Conf. Southeast. Assoc. Game Fish Comm. pp. 101-107.

Thompson, J. M. 1963. Mortality thresholds of fish in fly ash suspensions. Aust. J. Sci. 25: 414-415.

Toohey, J. I., C. D. Nelson, and G. Krotkov. (1965). Toxicity of phenazine carboxylic acids to some bacteria, algae, higher plants, and animals. Can. J. Bot. 43: 1151-1155.

Trama, F. B. (1954). The acute toxicity of some common salts of sodium, potassium and calcium to the common bluegill (*Lepomis macrochirus* Rafinesqu). Proc. Acad. Nat. Sci. Philadelphia 106: 185-205.

Trama, F. B. (1954). The acute toxicity of copper to the common bluegill (*Lepomis macrochirus* Rafinesque). Notulae Natur. No. 257: 1-13.

Trama, F. B. (1955). The acute toxicity of phenol of the common bluegill (*Lepomis macrochirus* Rafinesque). Notulae Natur., No. 269: 1-10.

Trama, F. B. and R. J. Benoit. (1960). Toxicity of hexavalent chromium to bluegills. J. Water Pollut. Contr. Fed. 32(8): 868-877.

Trembley, F. J. (1960). Research project on effects of condenser discharge water on aquatic life. Progress Rept. 1956-1959. Bethlehem, Penn., Lehigh Univ., Inst. of Res. 154 p.

Trembley, F. J. (1965). Effects of cooling water from steam-electric power plants on stream biota. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 334-345.

Tubb, R. A. and T. C. Dorris. (1965). Herbivorous insect populations in oil refinery effluent holding pond series. Limnol. Oceanogr. 10: 121-134.

Turnbull, H., J. G. DeMann, and R. F. Weston. (1954). Toxicity of various refinery materials to freshwater fish. Ind. Eng. Chem. 46: 324-333.

Turnbull-Kemp, P.St.J. (1958). Trout in Southern Rhodesia. V. On the toxicity of copper sulfate to trout. Rhodesia Agr. J. 55(6): 637-640.

Turner, N. (1965). DDT in Connecticut wildlife. Conn. Agr. Exp. Sta. Bull. 672, 11 p.

Tyler, A. V. (1966). Some lethal temperature relations of two minnows of the genus *Chrosomus*. Can. J. Zool. 44: 349-364.

Ukeles, R. (1962). Growth of pure cultures of marine phytoplankton in the presence of toxicants. Appl. Microbiol. 10(6): 532-537.

Ukeles, R. (1965). Inhibition of unicellular algae by synthetic surface-active agents. J. Phycol. 1: 102-110.

Ullman, W. W., R. W. Schaefer, and W. W. Sanderson. (1961). Arsenic accumulation by fish in lakes treated with sodium arsenite. J. Water Pollut. Contr. Fed. 33(4): 416-418.

U.S. Fish and Wildlife Circular 185 (1964). See Lennon and Walker (1964).

Van Overbeek, J., W. J. Hughes, and R. Blondeau. (1959). Acrolein for the control of water weeds and diseasecarrying water snails. Science 129: 335-336.

Van Valin, C. C., A. K. Andrews, and L. L. Eller. (1968). Some effects of Marex on two warm-water fishes. Trans. Amer. Fish. Soc. 97(2): 185-196.

Velsen, F.J.P. and D. F. Alderdice. (1967). Toxicities of two insecticides to young coho salmon. J. Fish. Res. Bd. Can. 24: 1173-1175.

Vernimmen, A. P., E. R. Henken, and J. C. Lamb. (1967). A short-term biochemical oxygen demand test. J. Water Pollut. Contr. Fed. 39(6): 1006-1020.

Vinson, S. B., C. E. Boyd, and D. E. Ferguson. (1963). Resistance to DDT in the mosquito fish, *Gambusia affinis*. Science 139: 217-218.

Vivier, P. and M. Nisbet. (1965). Toxicity of some herbicides, insecticides and industrial wastes. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 167-169.

Von Windeguth, D. L. and R. S. Patterson. (1966). The effects of two organic phosphate insecticides on segments of the aquatic biota. Mosquito News 26: 377-380.

Waldichuk, M. (1960). Effects of pulp and paper mill wastes on the marine environment. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Transactions of the 1959 Seminar. Cincinnati, Ohio, Robt. A. Taft San. Eng. Center, Tech. Rept. W60-3, pp. 160-176.

Walker, C. R. (1963). Endothal derivatives as aquatic herbicides in fishery habitats. Weeds 11: 226-332.

Walker, C. R. (1964). Dichlobenil as a herbicide in fish habitats. Weeds 12: 267-269.

Walker, C. R. (1965). Diuron, Fenuron, Monuron, Neburon, and TCA mixtures as aquatic herbicides in fish habitats. Weeds 13: 297-301.

Walker, C. R., R. E. Lennon, and B. L. Berger. (1964), Preliminary observations on the toxicity of Antimycin A to fish and other aquatic animals. In: Investigations in Fish Control. No. 2. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Circ. 186, 18 p.

Walker, C. R. and R. A. Schoettger. (1967). Method for determining MS-222 residues in fish. In: Investigations in Fish Control. No. 14. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 20, 10 p.

Walker, C. R. and R. A. Schoettger. (1967). Residues of MS-222 in four salmonids following anesthesia. In: Investigations in Fish Control. No. 15. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 21, 11 p.

Walker, C. R., P. J. Starkey, and L. L. Marking. (1966). Relation of chemical structure to fish toxicity in nitrosalicylanilides and related compounds. In: Investigations in fish control. No. 9. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl. Resour. Publ. 13, 12 p.

Wallen, I. E., W. C. Greer, and R. Lasater. (1957). Toxicity to *Gambusia affinis* of certain pure chemicals in turbid waters. Sewage Ind. Wastes 29(6): 695-711.

Ward, C. M. and W. M. Irwin. (1961). The relative resistance of thirteen species of fishes to petroleum refinery effluent. Proc. 15th Ann. Conf., Southeast. Assoc. Game Fish Comm., pp. 255-275.

Ware, G. W., M. K. Dee, and W. P. Cahill. (1968). Water flora as indicators of irrigation water contamination by DDT. Bull. Environm. Contam. Toxicol. 3(6): 333-338.

Warner, K. and O. C. Fenderson. (1962). Effects of DDT spraying for forest insects on Maine trout streams. J. Wildl. Manage. 26: 86-93.

Warner, R. E., K. K. Peterson, and L. Borgman. (1966). Behavioural pathology in fish: A quantitative study of sublethal pesticide toxication. In: N. W. Moore (ed.), Pesticides in the Environment and Their Effects on Wildlife. Oxford, Blackwell Scientific Publ. and J. Appl. Ecol. 3(Suppl.): 223-247.

Warnick, D. C. (1966). Growth rates of yellow perch in two North Dakota lakes after population reduction with Toxaphene. In: Investigations in fish control. No. 5. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 9,9 p.

Warren, C. E. and P. Doudoroff. (1958). The development of methods for using bioassays in the control of pulp mill waste disposal. Tappi 41(8): 211A-216A.

Warren, J. W. (1963). Toxicity tests of erythromycin thiocyanate in rainbow trout. Progr. Fish-Cult. 25: 88-92.

Water Pollution Control Federation Research Committee. (1958). A review of the literature of 1957 on sewage, waste treatment, and water pollution. J. Water Pollut. Contr. Fed. 30(7): 839-873.

Water Pollution Control Federation Research Committee. (1959). A review of the literature of 1958 on sewage, waste treatment, and water pollution. J. Water Pollut. Contr. Fed. 31(7): 763-803.

Water Pollution Control Federation Research Committee. (1960). A review of the literature of 1959 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 32(7): 681-720.

Water Pollution Control Federation Research Committee. (1961). A review of the literature of 1960 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 33(7): 681-710.

Water Pollution Control Federation Research Committee. (1962). A review of the literature of 1961 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 34(7): 629-703.

Water Pollution Control Federation Research Committee. (1963). A review of the literature of 1962 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 35(7): 819-876.

Water Pollution Control Federation Research Committee. (1964). A review of the literature of 1963 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 36(7): 791-863.

Water Pollution Control Federation Research Committee. (1965). A review of the literature of 1964 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 37(7): 887-979.

Water Pollution Control Federation Research Committee. (1966). A review of the literature of 1965 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 38(7): 1049-1137.

Water Pollution Control Federation Research Committee. (1967). A review of the literature of 1966 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 39(7): 1049-1154.

Water Pollution Control Federation Research Committee. (1968). A review of the literature of 1967 on wastewater and water pollution control. J. Water Pollut. Contr. Fed. 40(6): 897-1219.

Water Quality Act of 1965 (PL89-234), approved 10/2/1965.

Weatherholtz, W. M., G. W. Cornwell, R. W. Young, and R. E. Webb. (1967). Distribution of Heptachlor residues in pond ecosystems in southwestern Virginia. J. Agr. Food Chem. 15: 667-670.

Weaver, L. C., G. Gunnerson, A. W. Breidenbach, and J. J. Lichtenberg. (1965). Chlorinated hydrocarbon pesticides in major U.S. River basins. Public Health Rept. 80(6): 481-493.

Webb, F. E. (1960). Aerial forest spraying in Canada in relation to effects on aquatic life. In: C. M. Tarzwell (comp.), Biological Problems in Water Pollution, Trans. 1959 Seminar, Cincinnati, Ohio, R. A. Taft San. Eng. Center, Tech. Rept. W60-3, pp. 66-70.

Webb, W. E. (1961). Toxicity of certain pesticides to fish. Idaho Fish Game Dept. D-J Prog. Rept. F-34-R-2 (2), 36 p.

Webbe, G. and G. T. Shute. (1959). A further note on the action of fish of chlorinated hydrocarbons when used as larvicides. Ann. Trop. Med. Parasitol. 53(1): 47-50.

Wedemeyer, G. (1968). Uptake and distribution of  $Zn^{65}$  in the coho salmon egg (*Oncorhynchus kisutch*). Comp. Biochem. Physiol. 26: 271-279.

Weiss, C. M. (1959). Response of fish to sublethal exposures of organic phosphorus insecticides. Sewage Ind. Wastes 31(5): 580-593.

Weiss, C. M. (1961). Physiological effect of organic phosphorus insecticides on several species of fish. Trans. Amer. Fish. Soc. 90(2): 143-152.

Weiss, C. M. (1964). Organic pesticides and water pollution. Public Works 95(12): 84-87.

Weiss, C. M. (1965). Use of fish to detect organic insecticides in water. J. Water Pollut. Contr. Fed. 37(5): 647-658.

Weiss, C. M. and J. L. Botts. (1957). Factors affecting the response of fish to toxic materials. Sewage Ind. Wastes 29(7): 810-818.

Weiss, C. M. and J. L. Botts. (1957). The response of some freshwater fish to isopropyl methylphosphorofluoridate (Sarin) in water. Limnol. Oceanogr. 2: 363-370.

Weiss, C. M. and J. H. Gakstatter. (1964). Detection of pesticides in water by biochemical assay. J. Water Pollut. Contr. Fed. 36(2): 240-253.

Welch, E. B. and J. C. Spindler. (1964). DDT persistence and its effect on aquatic insects and fish after an aerial application, J. Water Pollut. Contr. Fed. 36(10): 1285-1292.

Welch, P. S. (1948). Limnological methods. N.Y., McGraw-Hill Book Co., Inc., 381 p.

Welch, P. S. (1952). Limnology, 2nd ed. N.Y., McGraw-Hill Book Co., Inc., 538 p. Weston, R. F. (1964). The value and use of water quality criteria to protect aquatic life. Ind. Water Wastes 9(1): 14-17.

Whitley, L. S. (1968). The resistance of tubificid worms to three common pollutants. Hydrobiologia 32(1-2): 193-205.

Whitten, B. K. and C. J. Goodnight. (1966). Toxicity of some common insecticides to tubificids. J. Water Pollut. Contr. Fed. 38(2): 227-235.

Wilber, C. G. (1965). A mathematical description of the toxicity of Sernyl to goldfish. Ohio J. Sci. 65: 43-46.

Wilber, C. G. (1965). The biology of water toxicants in sublethal concentrations. In: Biological Problems in Water Pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 326-331.

Wilhm, J. L. and T. C. Dorris. (1968). Biological parameters for water quality criteria. BioScience 18(6): 477-481.

Willford, W. A. (1966). Toxicity of 22 therapeutic compounds to six fishes. In: Investigations in Fish Control. No. 18. U.S. Fish Wildl. Serv., Bur. Sport Fish Wildl., Resour. Publ. 35, 10 p.

Willford, W. A. (1967). Toxicity of dimethyl sulfoxide (DMSO) to fish. In: Investigations in fish control. No. 20. U.S. Fish Wildl. Serv., Bur. Sport Fish. Wildl., Resour. Publ. 37, 8 p.

Williams, L. G. (1964). Possible relationships between plankton diatom species numbers and water-quality estimates. Ecology 45: 809-823.

Williams, L. G. and D. I. Mount. (1965). Influence of zinc on periphytic communities. Amer. J. Bot. 52(1): 26-34.

Wilson, B. R. (ed.). (1968). Environmental problems; Pesticides, thermal pollution, and environmental synergisms. Philadelphia, J. B. Lippincott Co., 183 p.

Wisniewski, T. F. (1958). Algae and their effects on dissolved oxygen and biochemical oxygen demand. In: Oxygen relationships in streams. Cincinnati, Ohio, U.S. Public Health Serv. Robt. A. Taft Eng. Center, Tech. Rept. W58-2, pp. 157-180.

Wisniewski, T. F. (1958). Algae and the effect on D.O. and B.O.D. Part II. Water Sewage Works 105: 300-305.

Woelke, C. E. (1965). Bioassays of pulp mill wastes with oysters. In: Biological problems in water pollution. Third Seminar, 1962. U.S. Public Health Serv. Publ. No. 999-WP-25, pp. 67-77.

Woelke, C. E. (1967). Measurement of water quality with the Pacific oyster embryo bioassay. In: Water quality criteria. Philadelphia, Pa., Amer. Soc. Testing and Materials. ASTM Spec. Tech. Publ. No. 416, pp. 112-120.

Wollitz, R. E. (1963). Effects of certain commercial fish toxicants on the limnology of three cold-water ponds, Montana. Proc. Montana Acad. Sci. 22: 54-81.

Wood, M. L. (1957). Biological aspects of stream pollution control in Arkansas. Proc. 10th Ann. Conf., Southeast Assoc. Game Fish Comm., pp. 136-138.

Woodwell, G. M., C. F. Wurster, and P. A. Isaacson. (1967). DDT residues in an east coast estuary: A case of biological concentration of a persistent insecticide. Science 156(3776): 821-824.

Workman, G. W. and J. M. Neuhold. (1963). Lethal concentrations of Toxaphene for goldfish, mosquito fish, and rainbow trout, with notes on detoxification. Progr. Fish-Cult. 25: 23-30.

Wuhrmann, K. (1959). Concerning some principles of the toxicology of fish. Transl. from: Bulletin du Centre Belge d'Etude et de Documentation des Eaux 15: 49, 1952. Nanaimo, B. C., Fish. Res. Bd. Can., Biol. Station Transl. Ser. No. 243, 19 p.

Wuhrmann, K. and H. Woker. (1955). Influence of temperature and oxygen tension on the toxicity of poisons to fish. Int. Ver. Theor. Angew. Limnol. Verh. 12: 795-801.

Wurster, C. F. (1968). DDT reduces photosynthesis by marine phytoplankton. Science 159: 1474-1475.

Wurtz, C. B. (1962). Zinc effects on fresh-water mollusks. Nautilus 76(2): 53-61.

Wurtz, C. B. and C. H. Bridges. (1961). Preliminary results from macroinvertebrate bioassays. Proc. Pa. Acad. Sci. 35: 51-56.

Wurtz, C. B. and T. Dolan. (1961). A biological method used in the evaluation of effects of thermal discharge in the Schuylkill River. Proc. 15th Ind. Waste Conf. Purdue Univ. 45(2): 461-472.

Wurtz, C. B. and C. E. Renn. (June 1965). Water temperature and aquatic life. (The Johns Hopkins University, Cooling water studies for Edison Electric Institute, Research Project RP-49.) N.Y., Edison Electric Inst. EEI Publ. 65-901, 99 p.

Yeo, R. R. (1967). Dissipation of Diquat and Paraquat, and effects on aquatic weeds and fish. Weeds 15: 42-46.

Young, F. N. (1961). Effects of pollution on natural associations of water beetles. Proc. 15th Ind. Waste Conf. Purdue Univ. 45(2): 373-380.

Zintgraff, G. D., C. H. Ward and A. W. Busch. (1968). Cyanide inhibition of mixed microbial populations. Presented in part at the Annual Meeting of the Society for Industrial Microbiology held at the Ohio State University, Sept. 3-6, 1968. Columbus, Ohio, 34 p.

ZoBell, C. E. (1964). The occurrence, effects, and fate of oil polluting the sea. In: Advances in Water Pollution Research, Proc. International Conference held in London, Spet. 1962. Vol. 3. N.Y., MacMillan, pp. 81-118.

## SECTION XV

## APPENDICES

- A. EXTRACTED DATA FROM ORIGINAL PAPERS CHEMICALS AND MIXTURES OF CHEMICALS
- B. EXTRACTED DATA FROM ORIGINAL PAPERS COMMERCIAL CHEMICAL PRODUCTS
- C. SPECIES INDEX FOR APPENDICES A AND B
- D. IDENTIFICATION OF COMMERCIAL CHEMICALS

Note: Both scientific and common names should be checked for complete retrieval of information for a given organism.

## EXTRACTED DATA FROM ORIGINAL PAPERS – CHEMICALS AND MIXTURES OF CHEMICALS

- Note: Names of chemicals and organisms are as given by the various authors. Readers should search for alternate, common, and/or scientific names of both chemical and aquatic species; and refer to report section on Extracted Data for further discussion of this appendix.
- Footnotes for Appendices A and B:
- (1) Letters represent:
  - B = bioassay, used in combination with S = static, CF = continuous flow, A = acute, and CH = chronic.
  - L = laboratory bioassay.
  - BOD = biochemical oxygen demand.
    - F = field study, used in combination with R = river, stream, creek, etc., L = lake or pond, M = marine, E = estuarine, and O = other (port facility, flooded area, etc.).
- (2) Field location is indicated by abbreviation of the state or country.
- (3) The number indicates ppm (mg/l), unless otherwise indicated by appropriate designations or (O). The letters within parentheses following indicate  $T = TL_m$ , K = kill, SB = sublethal effects, NTE = no toxic effect, or O = other. The number following these indicates the time in days at which observations were made. EC50, LC50, and similar designations for 50 percent lethality were all considered as  $TL_m$  and designated as such. The numbers within parentheses following these designations indicate the time in days when the effect was observed.
- (4) The following indicate (when underlined the variable was controlled):
  - a = water temperature
  - b = ambient air temperature
  - c = pH

```
d = alkalinity (total, phenolphthalein or caustic)
```

```
e = dissolved oxygen
```

```
f = hardness (total, carbonate, Mg, or CaO)
```

- g = turbidity
- h = oxidation-reduction potential
- i = chloride as Cl

```
j = BOD, 5 day; (J) = BOD, short-term
```

```
k = COD
```

```
l = nitrogen (as NO<sub>2</sub> or NO<sub>3</sub>)
```

- m = ammonia nitrogen as NH3
- n = phosphate (total, ortho-, or poly)
- o = solids (total, fixed, volatile, or suspended)

```
p = CO_2
```

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Acetaldehyde	Lagodon rhomboides	BSA	<u> </u>	70.0 (T1A)	а	Aerated sea water was used.	Daugherty and Garrett (1951)
Acetaldehyde	Lagodon rhomboides	BSA	_	70.0 (T1A)	-	Experiments were conducted in aerated salt water.	Garrett (1957)
Acetaldehyde	Sewage organisms	BOD	-	230 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxy- gen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Acetaldehyde	Lepomis macrochirus	BSA	-	53.0 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
Acetaidehyde	Nitzschia linearis Lepomis macrochirus	BSA	-	236.6- 249.1 (T5A) 53.0 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Acetaldehyde (a)- acetone (b)- copper (c)- acetic acid (d) mixture	Lepomis macrochirus	BSA	-	(a) 5.2 (T4A) (b) 5.2 (T4A) (c) 1.04 (T4A) (d) 26.0 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
Acetamide	Gambusia affinis	BSA	_	26,300 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Acetanilide	Sewage organisms	BOD	_	(NTE)	_	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Acetic acid	Daphnia magna	BSA	_	150 (O)	<u>a</u> e	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)

	Acetic acid	Semotilus atromaculatus	BSA	-	100 to 200 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
	Acetic acid	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T=toxic, NT=nontoxic, PT= partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl - NT Ma - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
	Acetic acid	Gambusia affinis	BSA	-	251 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Acetic acid	lctalurus punctatus	BSA	-	388 (T2A) 629 (K2)	<u>a</u> cfi	The experiment was conducted at 77 C.	Clemens and Sneed (1958)
•	Acetic acid	Channel catfish (fingerlings)	BSA	-	446 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Acetic acid	Culex sp (larvae) Daphnia magna Lepomis macrochirus	BSA	-	1500 (T1A) 47 (T1A) 100 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
CHEMICALS	Acetic acid	Lepomis macrochirus	BSA	-	75 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
ALS AND MIXTU	Acetic acid	Nitzschia linearis Lepomis macrochirus	BSA	-	74 (T5A) 75 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
TURES OF CHE	Acetic acid (a)- acetaldehyde (b)- acetone (c)- copper (d)- mixture	Lepomis macrochirus	BSA	-	(a) 26.0 (T4A) (b) 5.2 (T4A) (c) 5.2 (T4A) (d) 1.04 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES	Acetone	Daphnia magna	BSA	_	9280 (O)	<u>a</u> e	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
OF CHE	Acetone	Gambusia affinis	BSA	-	13,000 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
CHEMICALS	Acetone	Sewage organisms	BOD	_	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depict- ing the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	Acetone	Daphnia magna	BSA	-	10 (T2A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
А	Acetone	Lepomis macrochirus	BSA	-	8300 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
A-4	Acetone	Nitzschia Iinearis Lepomis macrochirus	BSA	-	11,493 to 11,727 (T5A) 8,300 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
	Acetone (a)- copper (b)- acetic acid (c)- acetaldehyde (d)- mixture	Lepomis macrochirus	BSA	-	(a) 5.2 (T4A) (b) 1.04 (T4A) (c) 26.0 (T4A) (d) 5.2 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
	Acetonitrile	Pimephales promelas Lepomis macrochirus	BSA	-	(H+S) 1000 (T4A) (S) 1850 (T4A)	c d e f	<ul><li>(H) Value in hard water</li><li>(S) Value in soft water</li></ul>	Henderson, et al (1960)
		Lebistes reticulatus			(S) 1650 (T4A)		The chemical caused no change in flavor of the cooked bluegill.	
	2-acetylamino- fluorene (AAF)	Zebrafish	BSA	-	(0)	_	The results of this investigation show that definite changes in the concentration of RNA and glycogen accompany the cell- ular disorganization in abnormal embryos induced by AAF. In embryos treated with AAF, there was a consistent decrease of RNA content of the liver, nervous tissue, sense organs, and the mucosal lining of the digestive tract. In general, this only occurred when concentrations of the chemical exceeded 0.03 percent.	Hiszoka (1958)

	Acetyl phenyl- hydrazine	Microcystis aeruginosa	L	-	100 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75-ml total volume. Chu Nø. 10 medium was used.	Fitzgerald, et al (1952)
	Acrolein	Sewage microorganisms	BOD	_	1.5 (0)	-	The chemical was studied as to how low levels (ppm) may affect BOD in domestic sewage. The chemical was toxic to sewage microorganisms at the level stated. To acclimated organisms the toxicity was 18 ppm.	Oberton and Stack (1957)
	Acrolein	Oyster	BCF	_	0.055 (O)	а	The value reported is a 96-hr $\mathrm{EC}_{50}$ (decreased shell growth).	Butler (1965)
	Acrolein	Fundulus similis (juvenile)	BSA	-	0.24 (O)	а	Water temperature was 21 C. The figure reported is a 48-hr $EC_{50}$ .	Butler (1965)
	Acrolein	Penaeus aztecus	L	-	0.19 (O)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality.	Butler (1965)
	Acrolein	Crassostrea virginica Penaeus aztecus Fundulus similis Phytoplankton	BCFA & BSA	-	0.05 (O) 0.1 (O) 0.24 (T2CFA)	-	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
	Acrolein	Salmon	BSA		0.08 (T2A)	_	Data are given as LC50.	Bohmont (1967)
CHEMICA	Acrylaldehyde (acrolein)	Potamogeton	BSA	-	100 (O)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. Injury rating of 8.3.	Frank, et al (1961)
		modosus Potamogeton pectinatus Elodea canadensis			100 (O) 100 (K4wk)		Injury rating of 9.6.	
	Acrylonitrile	Lagodon rhomboides	BSA		24.5(T1A)	а	Aerated seawater was used.	Daugherty and Garrett (1951)
	Acrylonitrile	Lepomis macrochirus Pomoxis annularis	BSA & CH	~	0.05-0.1 (100% KS) 0.1-1.0 (100% KCH) 6.0-10.0 (100% KCH)	<u>a</u>	Additional data are presented for less than 24 hr.	Renn (1955)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Acrylonitrile	Lagodon rhomboides	BSA	-	24.5 (T1A)	_	Experiments were conducted in aerated salt water.	Garrett (1957)
Acrylonitrile	Pimephales promelas Lepomis macrochirus Lebistes	BSA	-	(S) 18.1 (T4A) (H) 14.3 (T4A) (S) 11.8 (T4A) (S) 33.5 (T4A)	c d e f	(H) Value in hard water (S) Value in soft water The chemical did not change the flavor of the cooked bluegill.	Henderson, et al (1960)
Adipic acid	reticulatus Lepomis macrochirus	BSA	_	330 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Adiponitrile	Pimephales promelas	BSA	_	(S) 1250 (T4A) (H) 820 (T4A)	c d e f	(H) Value in hardwater (S) Value in softwater	Henderson, et al (196.0)
	Lepomis macrochirus			(S) 720 (T4A)			
	Lebistes reticulatus			(S) 775 (T4A)		The chemical produced no change in the flavor of the cooked bluegill.	
Alkyl aryl bromide	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u> _	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following {T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl $-$ NT Ma $-$ T So $-$ NT Cv $-$ NT Gp $-$ NT Np $-$ NT	Palmer and Maloney (1955)
Alkyl-dimethyl- ammonium chlorides	Cylindrospermum licheniforme (Cl) Gleocapsa sp (G) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzchia palea (Np)	L	_	2.0 (O)	<u>a</u> _	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl $=$ PT (7) G $=$ NT So $=$ PT (7) Cv $=$ PT (3) Gp $=$ NT Np $=$ PT (3)	Palmer and Maloney (1955)
Alkyl sulfate	Pimephales promelas (juveniles)	BSA	_	(S) 5.1-5.9 (T1-4A) (H) 5.9-6.1 (T1-4A)	<u>acdf</u>	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard water (H). The surfactant rather than the builder contained the toxicant.	Henderson, et al (1959)

A-6

Alkyl benzene sulfate - See ABS in Appendix B.

Aluminum ammonium sulfate	Daphnia magna	BSA	_	190 (O)	<u>a</u> e	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Aluminum chloride	Gambusia affinis	BSA	-	135 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Aluminum chloride	Daphnia magna	BSA	_	<6.7 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
Aluminum nitrate	Gasterosteus aculeatus	BSA	-	0.07 (K10)	_	Solutions were made up in tap water. 3.0 to 5.0 cm stickleback fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
Aluminum potassium sulfate	Daphnia magna	BSA		206 (O)	<u>a</u> e	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Aluminum sulfate	Daphnia magna	BSA	-	136 (O)	<u>a</u> e	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Aluminum sulfate	Micropterus salmoides Lepomis machrochirus Goldfish	BSA	-	100 (O) 100 (O) 100 (O)	<u>a</u> cfpi	The disposal of cannery wastes frequently involves the use of chemicals for treatment purposes. Ferrous sulphate, alum, and lime are used in chemical coagulation; sodium carbonate for acidity control in biological filters; and sodium nitrate in lagoons for odor control. Lye (sodium hydroxide) peeling of certain fruits and vegetables is not uncommon. These chemicals, in whole or part, are discharged in most cases to a stream. The concentrations listed permitted all fish to survive indefinitely.	Sanborn (1945)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Aluminum sulfate	Sewage organisms	BOD	_	18.0 (O)	_	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treatment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
Aluminum sulfate	Gambusia affinis	BSA	-	240 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
p-aminodi- ethylaniline HCl	Microcystis aeruginosa	L	-	100 (K)	<u>a</u> , etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/mł, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
p-aminodi- methylaniline	Microcystis aeruginosa	L	_	100 (K)	<u>a</u> , etc	Comment same as above.	Fitzgerald, et al (1952)
p-aminodi- methylaniline HCl	Microcystis aeruginosa	L	-	100 (K)	<u>a</u> , etc	Comment same as above.	Fitzgerald, et al (1952)
η-(3-amino- propyl) rosinamine D diacetate (28 percent active)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u> _	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T (14) Ma - T So - PT Cv - T (14) Gp - T Np - T	Palmer and Maloney (1955)
p-aminophenol	Daphnia magna	BSA	-	2 (K2A)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
4-amino-m toluene- sulfonic acid	Gambusia affinis	BSA	-	410 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

Ammor	nia	Trout	BSA	_	(O)	a e	No quantitative data are reported. 30 ppm of nitrogen was added as ammonium chloride. Carbon dioxide in concentrations up to 30 ppm reduced the toxicity of the ammonia by lowering the pH of the water. Concentrations of 60 ppm of $CO_2$ were toxic but not lethal when the concentration of dissolved oxygen was low. A concentration of 240 ppm of $CO_2$ was lethal to trout in little more than one hour.	Herbert (1955)
Ammo	nia	Pimephales promelas	BSA	-	(H) 8.2 (T4A) (S) 5.9 (T4A)	cdef	<ul><li>(H) Value in hardwater</li><li>(S) Value in softwater</li></ul>	Henderson, et al (1960)
Ammo (unior		Salmo gairdnerii	BSA	-	0.4 (T1A)	a b c d e	Toxicity of ammonia or of ammonium salts was increased by a rise in pH value from 7.0 to 8.2. Toxicity of such solutions to fish apparently depended upon the concentration of the un- ionized ammonia molecule present. Variation was attributed to the increase in the concen- tration of free carbon dioxide at the gill surfaces.	Lloyd and Herbert (1960)
Ammo	nia	Salmo gairdnerii	BSA	_	100-200 (O)	<u>a</u> сер	The major factor determining the toxicity of ammonia is the pH of the water. Temperature, dissolved oxygen, and bicarbonate alkalinity are also important. Only unionized ammonia was toxic to fish. At a pH of 7.0 the threshold value for ammonia ranges between 100 and 200 ppm (as N), depending on the bicarbonate hardness.	Lloyd (1961)
Ammor	nia	Gambusia affinis	BSA	-	(0)	a c d i	The pH value and temperature had a marked effect upon the toxicity of ammonia solutions. As the pH was raised, the toxicity increased markedly. The concentration of unionized ammonia present in each test was calculated using the mean temper- ature and the pH value. The absence of toxic action by tests at a total ammonia concentration equivalent to 120 mg/IN.	Hemens (1966)
CHEMICALS A	nia	Green sunfish	BSA	-	(0)	-	Ammonia or ammonium hydroxide was found to repel fish at 8.5, 10, and 20 mg/l. At 1.7 mg/l no repellency was noted. In concentrations of 10 and 22 mg/l, ammonia killed the fish in repellent studies before they had the opportunity to move out of the area containing the substance.	Summerfelt and Lewis (1967)
	nia	Abramis brama Perca fluviatillis Rutilis rutilis scardinius erythrophthalmus Salmo gairdnerii	BCF	-	0.41 (T7CF) 0.29 (T7CF) 0.35 (T5CF) 0.36 (T6CF) 0.41 (T2CF)	a c d e f	The T at LC50 values are asymptotic values of undissociated ammonia (mgN/l). Additional data are presented.	Ball (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Ammonia	Salmo gairdneri	BSA	_	(0)	acim	After 24-hr exposure the mean blood levels for total ammonia showed a direct linear correlation with ambient ammonia and ranged from 38 to 70 g/ml. Fish exposed to 0-1 g/ml nonionic ammonia had mean blood levels which ranged from 0.6 to 1.3 g/ml. Ammonia in concentrations up to 10 g NH3/ml was found to have no significant effect on the ability of hemoglobin to combine with oxygen in vitro.	Fromm and Gillette (1968)
Ammonia	Salmo gairdneri	BSA	-	34-47 (T2A)	a c d e f o	The concentration killing a half batch of fish in 2 days pro- vides a reasonable estimate of the threshold concentra- tion. The lethality of this chemical depends upon all the experimental variables listed and the concentration of undissociated ammonia which is present.	Brown (1968)
Ammonia (unionized)	Salmo gairdnerii	FR	Stevenage Herts.	(O)	<u>a c e l m</u>	Survival of rainbow trout in concentrations of unionized ammonia in the range of 0.86-1.96 ppm of nitrogen in- creased as the concentration of dissolved oxygen was raised from 1.5 to 8.5 ppm. The effect of dissolved oxy- gen in increasing survival time was greater in the lower concentrations of unionized ammonia.	Downing and Merkens (1955)
Ammonia (unionized)	Salmo gairdnerii Perca fluviatilis Rutilus rutilus Gobio gobio	BSA	-	(O)	асеор	The resistance to rapidly lethal concentrations of un- ionized ammonia ranging from about 2.0 to 8.8 ppm nitrogen was determined in tensions of dissolved oxygen 53.4 and 96.7% of air saturation value at 15.2 C. Period of survival decreased with rise in concentration of unionized ammonia. The effect of oxygen tension on period of survival was greatest in the lowest concentra- tions of unionized ammonia.	Markens and Downing (1957)
Ammonia plus carbon dioxide	Rainbow trout	BSA	-	(0)	<u>a e m n</u>	The reduction of toxicity of ammonia solutions by the addition of carbon dioxide, was due to lowering the pH of the solution. $60-240$ ppm CO <sub>2</sub> in solution was toxic within 12 hr. 30 ppm ammonia nitrogen was toxic, but up to 30 ppm CO <sub>2</sub> increased fish survival time.	Alabaster and Herbert (1954)
Ammonium acetate	Gambusia affinis	BSA	-	238 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Ammonium borofluoride	Sewage organisms	BOD	-	87.0 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treatment plant, BOD was used as the parameter to mea- sure the effect of the chemical. The chemical concentra- tion cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
Ammonium carbonate	Gambusia affinis	BSA	-	238 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

	Ammonium chloride	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chem- icals, possible mechanism of action of some, the effect of temperature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.224N solution, fish survived 99 minutes.	Powers (1918)
	Ammonium chloride	Daphnia magna	BSA	-	<134 (0)	<u>a</u> e	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Ammonium chloride	Salmo gairdnerii	BSA	-	(O) Tap water 1000 ppm – 27.3 min 1000 ppm – 52.5 min 50 ppm – ≥1000 min	acef	Tap or distilled water used as diluent. Toxicity defined as the average time when the fish lost equilibrium when exposed to the test chemical (ppm ammonia).	Grindley (1946)
					Distilled water 3000 ppm – 292 min 1000 ppm – 725 min 100 ppm > 4320 min			
	Ammonium chloride	Daphnia magna	BSA	-	91 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
CHEMIC	Ammonium chloride	Lepomis macrochirus	BCFA	-	6.0 (T4A)	a c e f	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. Toxicity was dependent upon the concentration of un- dissociated NH4OH which is dependent upon pH. The initial pH was 9.0 and after four days it was 7.5.	Cairns and Scheier (1955)
ALS AND MINTUR	Ammonium chloride	Daphnia magna	BSA	-	246,6 (O)	a C	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of in- organic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
	Ammonium chloride	Gambusia affinis	BSA	-	510 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Ammonium chloride	Lepomis macrochirus	BSA	_	7.7 (T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Ammonium chloride	Rainbow trout	BSA		(O)	acd	The 48-hour LD <sub>50</sub> of ammonium chloride (as N) as interpo- lated from three graphs may be 30, 24, or 12 ppm. The effect of dissolved oxygen is also discussed.	Herbert (1961)
(as N) Ammonium chloride (as N)	Salmo gairdnerii	BSA	-	24.6 (T2A)	a <b>c d</b> f	A mathematical equation was derived to explain the com- bined toxicities of this salt and zinc sulfate.	Herbert and Shurben (1964)
Ammonium chloride	Carassius carassius Daphnia magna Lepomis macrochirus Lymnaea sp (eggs)	BSA	-	202 (T1A) 161 (T2A) 50 (T4A) 139 (T4A) 725 (T1-4A) 241 (T1A) 173 (T2A) 70 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Ammonium chromate	Gambusia affinis	BSA	-	270 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Ammonium dichromate	Gambusia affinis	BSA	—	212 (T2A)	acdeg	Comment same as above.	Wallen, et al (1957)
Ammonium hydroxide	Daphnia magna	BSA	-	<8.75 (0)	<u>a</u> e	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Ammonium hydroxide	Gasterosteus aculeatus	BSA	_	(0)	c e	Tap water was used to make up the solutions. The fish avoided concentrations of 0.04 and 0.01N, but seemed attracted to concentrations of 0.001 and 0.0001N.	Jones (1948)
Ammonium hydroxide (as ammonia)	Semotilus atromaculatus	BSA	-	5 to 15 (CR)	<u>a</u> e 	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Ammonium hydroxide	Gambusia affinis	BSA	_	37 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Ammonium hydroxide	Fish	BSA	-	4.3 x 10 <sup>-5</sup> M (K)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)

	Ammonium hydroxide	Daphnia magna	BSA	-	60 (T1A) 32 (T2A) 20 (T4A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
	Ammonium nitrate	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chem- icals, possible mechanism of action of some, the effect of temperature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.213N solution, fish survived 78 minutes.	Powers (1918)
	Ammonium salt	Nitzschia linearis Physa heterostropha Lepomis macrochirus	BSA	-	420 (T5A) 90.0 (T4A) 3.4 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
	Ammonium salts	Salmo gairdnerii	BSA	_	(0)	<u>a e</u>	This is a study of the effect of varying dissolved oxygen concentrations on the toxicity of selected chemicals. The toxicity of heavy metals, ammonia, and monohydric phenols increased as the dissolved oxygen in water was reduced. The most obvious reaction of fish to increase the volume of water passed over the gills, and this may increase the amount of poison reaching the surface of the gill epithelium. The concentration of the chemical in the water was not specified.	Lloyd (1961)
	Ammonium sulfate	Daphna magna	BSA	-	<106 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentra- tion was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
CHEMICALS AND	Ammonium sulfate	Daphnia magna	BSA	-	288.5 (O)	a c	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inor- ganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
	Ammonium sulfate	Salmo gairdnerii	BSA	-	(O) Tap water 1000 ppm – 29.8 min Distilled water 3000 ppm – 318 min 1000 ppm – 847 min 100 ppm >5760 min	<u>a</u> c <u>e</u> f	Tap or distilled water used as diluent. Toxicity defined as the avg. time when the fish lost equilibrium when ex- posed to the test chemical (ppm ammonia).	Grindley (1946)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Ammonium U sulfate	Gambusia affinis	BSA	-	1,400 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
M XT Ammonium sulfate m S	Daphnia magna	BSA	-	423 (T1A) 433 (T2A) 292 (T4A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Ammonium Sulphate Ammonium	Biomorpholaria a. alexandrina Bulinus truncatus	BSA	-	800 (K1A) 300 (K1A)	а	The degree of tolerance for vector snails of biharziasis chem- icals is somewhat dependent upon temperature. The tem- perature at which (K1A) occurred was 28 C.	Gohar and El-Gindy (1961)
≤ Ammonium > sulfide	Gambusia affinis	BSA	-	248 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented,	Wallen, et al (1957)
Ammonium sulfite	Gambusia affinis	BSA	-	240 (T2A)	<u>a</u> cdeg	Comment same as above.	Wallen, et al (1957)
Ammonium sulfite	Daphnia magna	BSA	-	299 (T1A) 273 (T2A) 203 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Ammonium thiocyanate	Gambusia affinis	BSA	_	420 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented,	Wallen, et al (1957)
Amyl acetate	Semotilus atromaculatus	BSA	-	50 to 120 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
N-amyl-acetate	Gambusia affinis	BSA	-	65 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented,	Wallen, et al (1957)
n-amyi alcohol	Semotilus atromaculatus	BSA	_	350 to 500 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
t-amyl alcohol	Semotilus atromaculatus	BSA	-	1,300 to 2,000 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Aniline	Daphnia magna	BSA	-	279 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoreti- cally infinite) exposure	Anderson {1944}

cally infinite) exposure.

APPENDIX A

	Aniline	Microcystis aeruginosa	L	-	50 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
	Aniline hydrochloride	Daphnia magna	BSA	_	5.5 (K2)	a	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
	Antimony potassium tartrate	Pimephales promelas	BSA	-	12 (T4A) H 20 (T4A) S	acdf	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
	Antimony trichloride	Daphnia magna	BSA	-	37 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	Antimony trichloride	Pimephales promelas	BSA	~	17 (T4A) H 9 (T4A) S	a c d f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
	Antimony trìoxide	Pimephales promelas	BSA	-	≫80 (T4A) H ≫80 (T4A) S	acdf	Comment same as above.	Tarzwell and Henderson (1960)
	Arsenite	Lepomis macrochirus (eggs) L. cyanellus (eggs) Micropterus dolomieui (eggs)	L	-	15/7 (O), 8 (NTE) 15 (NTE), 8 (NTE) 15/6 (O), 8 (NTE)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (O).	Hiltibran (1967)
_	Barium carbonate	Gambusia affinis	BSA	_	10,000 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
CHEMICAI C AN	Barium chloride	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.172N solution, fish survived 169 minutes.	Powers (1918)
	Barium chloride	Daphnía magna	BSA	-	<83 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Barium chloride	Daphnia magna	BSA	-	29 (O)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen ts	Reference (Year)
Barium chloride	Gambusia affinis	BSA		3,200 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Barium chloride	Rana sp (eggs)	BSA	_	24,430 K	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Barium nitrate	Gasterosteus aculeatus	BSA	-	400 (K10)	~	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
Benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their rela- tive position(s) in the molecule. At 10 ppm, there was no toxicity to goldfish or trout.	Walker, et al (1966)
Benzene	Gambusia affinis	BSA	-	395 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Benzene	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	31 (T4A) 22 (T4A) 32 (T4A)	<u>a</u> c d e f	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
Benzidine	Microcystis aeruginosa	L	-	50 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Benzoic acid	Carassius auratus	BSA	-	0.165 (K)	<u>a</u>	Goldfish weighed between 2 and 4 g. Temperature was maintained at 27.0 $\pm$ 0.2 C.	Gersdorff (1943)
Benzoic acid	Daphnia magna	BSA	_	146 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)

Benzoic acid	Gambusia affinis	BSA	-	225 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)	
Benzonitrile	Pimephales promelas	BSA		(S) 135.0 (T4A) (H) 78.0 (T4A)	c d e f	<ul><li>(H) Value in softwater</li><li>(S) Value in softwater</li></ul>	Henderson, et al (1960)	
	Lepomis macrochirus			(S) 78.0 (T4A)				
	Lebistes reticulatus			(S) 400.0 (T4A)		The chemical did not change the flavor of the cooked bluegill.		
2-benzoyl-1,3- dichloropropane	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): Cl - T (7), PT (21) Ma - T So - PT (7) Cv - T Gp - T Np - T	(1955)	≥
3-benzyl-5,5- dimethyl-2- imidazolinethione	Microcystis aeruginosa	L	_	10.0 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)	APPENDIX
bis-benzyl ethylene diamine diacetate	Semotilus atromaculatus	BSA	-	5 to 20 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	(1332)	NX A
Beryllium chloride	Pimephales promelas	BSA	-	(H) 15 (T4A) (S) 0.15 (T4A)	a c d f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)	
Beryllium nitrate	Pimephales promelas	BSA	-	(H) 20 (T4A) (S) 0.15 (T4A)	acdf	Comment same as above.	Tarzwell and Henderson (1960)	
Beryllium sulfate	Pimephales promelas Lepomis macrochirus	BSA	_	(H) 11 (T4A) (S) 0.2 (T4A) (H) 12 (T4A) (S) 1.3 (T4A)	a c d f	Comment same as above.	Tarzwell and Henderson (1960)	
Beryllium sulfate plus sodium tartrate	Goldfish Minnow Snails Water plants	BSA	-	(O)	c <u>e</u>	After 10 days of incremental additions of the chemicals to the aquarium, the final concentrations were: beryllium – 28.5 ppm; sulfate – 302 ppm; sodium tartrate – 664 ppm. No toxic effect to the animals or plants was observed after 10 days of exposure.	Pomelee (1953)	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Boric acid	Sewage organisms	BOD		480 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treatment plant. BOD was used as the parameter to mea- sure the effect of the chemical. The chemical concentra- tion cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
D Boric acid	Gambusia affinis	BSA		10,500 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
OHEM Boric acid CO ALS	Sewage organisms	BOD	_	>1000 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxy- gen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Bromine	Chlorella pyrenoidosa	BSA	_	0.18 (O) 0.42 (O)		<ul> <li>At 0.18 ppm, 2,100 cells/mm<sup>3</sup> remained at the end of 4 days as compared with a count of 2,383 cells/mm<sup>3</sup> in control.</li> <li>At 0.42 ppm, 270 cells/mm<sup>3</sup> remained at the end of 4 days as compared with 2,383 cells/mm<sup>3</sup> in controls.</li> <li>Bromine showed no inhibitory effect in the first 48 hr.</li> <li>Experiments were carried out in seven-liter containers of tap water.</li> <li>By maintaining a constant level of 0.2 ppm of bromine, it would be possible to kill algae in water.</li> </ul>	Kott, et al (1966)
3'-bromo-3, 5-dinitro- benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biolog- ical activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 1.0 ppm, this chemical was toxic to 4 out of 10 trout; but at the concentrations (.1, 1.0, 10.0) there was no toxicity to goldfish.	Walker, et al (1966)
4'-bromo-3, 5-dinitrobenz- anilide	Salmo gairdnerii	BSA	-	(O)	<u>a</u>	Comment same as above except that at 10 ppm the chem- ical was not toxic to trout or goldfish.	Walker, et al (1966)
	Carassius auratus			(0)			
4'-bromo-2- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10 (K2) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm, this chem- ical was toxic to 2 out of 10 goldfish in 48 hours.	Walker, et al (1966)

	2'-bromo-3- nitrosalicyl-	Sea lamprey (larva)	BSA	-	1.0 (K)	See Applegate,	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout	Starkey and Howell
	anilide	Salmo gairdneri (fingerling)			1.0 (K)	et al (1957-1958)	as a function of substituent loci.	(1966)
	3'-bromo-3- nitrosalicyl-	Sea lamprey (larva)	BSA	-	0.3 (K)	Ditto	Comment same as above.	Starkey and Howell
	anilide	Salmo gairdneri (fingerling)			(0)		1.0 ppm killed 25%.	(1966)
	4'-bromo-3- nitrosalicyl-	Sea lamprey (larva)	BSA	-	0.3 (K)	"	Comment same as above.	Starkey and Howell
	aniline	<i>Salmo gairdneri</i> (fingerling)			(0)		1.0 ppm killed 25%.	(1966)
	4'-bromo-5- nitrosalicyl-	Sea lamprey (larva)	BSA		0.5 (K)	"	Comment same as above.	Starkey and Howell
	anilide	<i>Salmo gairdneri</i> (fingerling)			(O)		1.5 ppm killed 25%.	(1966)
	3-bromo-4- nitrophenol	Petromyzon marinus	BSA	-	5 (K 100%)	<u>a</u>	Mortality occurred in approximately 24 hr. This was a study on controlling sea lamprey larvae.	Ball (1966)
	(free phenol)	Lepomis macrochirus	BSA		15 (K 10%)	<u>a</u>		
		Salmo gairdnerii	BSA	-	11 (K 10%)	<u>a</u>		
	2-bromo-4- nitrophenol	Petromyzon marinus	BSA	-	5 (K 100%)	<u>a</u>	Comment same as above.	Ball (1966)
	(free phenol)	Salmo gairdnerii	BSA	-	13 (K 10%)	<u>a</u>		
2		S. trutta	BSA	-	11 (K 10%)	<u>a</u>		
	2-bromo-4- nitrophenol	Petromyzon marinus	BSA	-	7 (K 100%)	<u>a</u>	Comment same as above.	Ball (1966)
	(Na salt)	Salmo gairdnerii	BSA	_	15 (K 10%)	<u>a</u>		
	2-bromo-4- nitrophenol	Petromyzon marinus (Iarvae)	BSA	-	10 (K14)	<u>a</u>	Additional data are presented.	Piavis (1962)
Ì	3-bromo-4-	Petromyzon	BSA	-		<u>a</u>	Comment same as above.	Piavis
	nitrophenol	<i>marinus</i> (embryos and prolarvae)			10 (K5-18)			(1962)
2		(larvae)			10 (K2-4 hr)			

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES OF CHEMICALS		Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	ä	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the sali- cylanilides and benzanilides increased toxicity to fish. Sim- ilar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
	3'-bromo-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)
	4'-bromo-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)
A	2-butanone	Gambusia affinis	BSA	_	5,600 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
A-20	n-butyl alcohol	Semotilus atromaculatus	BSA	-	1,000 to 1,400 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
	t-butyi alcohol	Semotilus atromaculatus	BSAq	_	3,000 to 6,000 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
	Butyric acid	Daphnia magna Lepomis macrochirus	BSA	-	61 (T2A) 200 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
	Cadmium	Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	BSA	-	1.0 (K) 1.0 (K) 0.01 (K)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide — a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)

Cadmium	Salmo gairdnerii	BCFA	-	(O)	_	A small, cone-shaped, cadmium-plated metal screen was used to cover a 2-inch pipe outlet. Recirculating 2,500 gallons of water through the screen at the rate of 50 gallons per min- ute killed 16-per-pound rainbow trout in 24 hours. Rainbow trout placed in a 15-gallon tub of water, with recirculation through the cadmium screen were dead within 10 hours.	Roberts (1963)
Cadmium	Lepomis macrochirus Ictalurus nebulosus	BSCFCH	_	0.1-100.0	• acdef	Fish were exposed to 8, 16, and 20 ppm of cadmium for varying periods of time (up to 90 days). In living fish the accumulation of cadmium never exceeded 130 $\mu$ g/g of gill tissue, based on dry weight. In fish that died of poisoning, the accumulation of cadmium was a maximum of 634 $\mu$ g/g of gill tissue. The authors state that high cadmium content (3-400 $\mu$ g/g) in the liver of a fish would indicate a past history of exposure.	Mount and Stephan (1967)
Cadmium	Salmo gairdnerii	BCFA	_	0.008- 0.01 (T7A) 30 mg (T1A)	abf	The data show that even at high concentrations, the toxic effect to the fish was very slow. Experiments were con- ducted in hard water.	Bali (1967)
Cadmium	Salmo gairdnerii	BCFA	-	30 (T1A)	abf	A 7-day TL <sub>m</sub> may be between 0.008 and 0.01 ppm. Despite this high toxicity, the response of the fish to the poison was initially very slow, even at high concentrations.	Velsen and Alderdice (1967)
Cadmium chloride	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemi- cals, possible mechanism of action of some, the effect of temperature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a 0.157N solution, fish survived 70 minutes; in a solu- tion of 0.000000037N, they survived 442 minutes.	Powers⊷ (1918)
Cadmium chloride	Daphnia magna	BSA	_	<0.0026 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
Cadmium chloride <u>O</u>	Pimephales promelas	BSA	-	5 (T4A) H 0.9 (T4A) S \	a c d f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
Cadmium Cadmium Chloride	Limnaea palustris (eggs)	BSA	-	6 x 10 <sup>-6</sup> m (K1)	<u>a</u> c	Toxicity is given in molar concentrations for maximum direct mortality (kill) in 4 hours.	Morrill (1963)
Cadmium A chloride D S S S S S S S S	Pimephales promelas Lepomis macrochirus Lebistes reticulatus Green sunfish	BSA	-	(S) 1.05 (T4A) (H) 72.6 (T4A) (S) 1.94 (T4A) (S) 1.27 (T4A) (S) 2.84 (T4A) (H) 66.0 (T4A)	c d e f	(S) Soft water (H) Hard water Values are expressed as mg/I of metal.	Pickering and Henderson (1965)
Cadmium cyanide complex	Lepomis macrochirus (juveniles)	BSA	-	0.64 (O)	<u>a</u> cdf <u>p</u>	For the concentration given, the median resistance time was 134 minutes.	Doudoroff, et al (1966)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES	Cadmium cya- nide complex, sodium cya- nide (439 ppm CN), and cad- mium sulfate (528 ppm Cd)	Pimephales promelas	BSA	-	0.17 (T4A)	a c	Synthetic soft water was used. Toxicity data given as number of test fish surviving after exposure at 24, 48, and 96 hr. TL <sub>m</sub> values were estimated by straight-line graphical inter- polation and given in ppm CN <sup>-</sup> .	Doudoroff, et al (1956)
OF CHEMICALS	Cadmium nitrate	Gasterosteus aculeatus	BSA	-	0.2 (K10)	_	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
ALS	Cadmium sulfate	Sewage organisms	BOD	-	142 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxigrams de- picting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
A-22	Caffeine	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 0.285 g/liter, fish survived 94 minutes.	Powers (1918)
	Calcium c irbonate	Gambusia affinis	BSA	_	56,000 (T2A)	a c d e g	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Calcium chloride	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.249N solution, fish survived 174 minutes.	Powers (1918)
	Calcium chloride	Daphnia magna	BSA	_	1332 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Calcium chloride	Daphnia magna	BSA	-	920 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)

	Calcium chloride	Lepomis macrochirus	BSA	-	10,650 (T4A)	a d e f	This paper reports the $LD_{50}$ in 96 hours for 8 common in- organic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured. If this salt is toxic to fish, this experiment did not demonstrate it.	Trama (1954)
	Calcium chloride	Lepomis macrochirus	BCFA	_	9,500 (T4A) small 11,300 (T4f) large	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period.	Cairns and Scheier (1955)
	Calcium chloride	Daphnia magna	BSA	-	3,972 (O)	a c	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inor- ganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
	Calcium chloride	Gambusia affinis	BSA	-	13,400 (T2A)	acdeg	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Calcium chloride	Lepomis macrochirus	BSA	-	11,300 (T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
	Calcium chloride	Daphnia magna Lepomis macrochirus Lymnaea sp (eggs)	BSA	-	3,526 (T1A) 3,005 (T2A) 8,350 (T1A) 4,485 (T1A) 3,094 (T2A) 2,373 (T3A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
CHEMICALS	Calcium chloride	Nitzschia linearis Lepomis macrochirus	BSA	-	3,130 (T5A) 10,650 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Calcium hydroxide	Micropterus salmoides Lepomis machrochirus Goldfish	BSA	_	100 (O) 100 (O) 100 (O)	<u>a</u> cfpi	The disposal of cannery wastes frequently involves the use of chemicals for treatment purposes. Ferrous sulphate, alum, and lime are used in chemical coagulation; sodium carbonate for acidity control in biological filters; and sodium nitrate in lagoons for odor control. Lye (sodium hydroxide) peeling of certain fruits and vegetables is not uncommon. These chemicals, in whole or part, are dis- charged in most cases to a stream. The concentration listed permitted large mouth bass to sur- vive 3 to 5 hours, bluegills to survive 2 to 4.5 hours, and goldfish to survive 3 to 3.5 hours.	Sanborn (1945)
Calcium hydroxide	Gambusia affinis	BSA	-	220 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Calcium hydroxide	Biomorpholaria alexandrina Bulinus truncatus Lymnaea caillaudi	BSA	-	300 (K1) 300 (K1) 300 (K1)	а	The degree of tolerance for vector snails of bilharziasis to various chemicals is somewhat dependent upon tempera- ture. The temperature at which (K1) occurred was 28 C.	Gohar and El-Gindy (1961)
Calcium hypochlorite	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u> _	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI $-T$ (3) Ma $-T$ (3) So $-T$ (3), PT (7) Cv $-T$ (3) Gp $-T$ (3) Np $-T$ (3)	Palmer and Maloney (1955)
Calcium hypochlorite	Blue-green algae Cylindrospermum Anabaena Anacystis Calothrix Nostoc Oscillatoria Plectonema Green algae Ankistrodesmus Chlorella Closterium Oocystis Scenedesmus Stigeoclonium Zygneme	L	_	2.0 (O)	_	Ca(OCI) <sub>2</sub> was toxic or partially toxic to all of the algae species at the indicated concentration for 28 days.	Kemp, et al (1966)

		Green flagellate and yellow algae Chlamydomonas Pandorina Tribonema Gomphonema Navicula Nitzchia						
	Calcium nitrate	Carassius carassius	BSA	_	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.192N solution, fish survived 186 minutes.	Powers (1918)
	Calcium nitrate	Gasterosteus aculeatus	BSA	_	800 (K10)	_	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
>	Calcium nitrate	Gasterosteus aculeatus Phoxinus phoxinus	BSA	-	(O)	C e	Tap water was used to make up the solutions. The fish were indifferent to dilute solutions – 0.001 N, but they were attracted to a solution of 0.04 N. At 0.0004 N they swam out of the solution to a water zone. At 0.00002 N they still showed an avoidance reaction.	Jones (1948)
20	Calcium nitrate	Lepomis macrochirus	BSA	-	10,000 (T4A)	a d e f	This paper reports the LD <sub>50</sub> in 96 hours for 8 common in- organic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured. If this salt is toxic to fish, this experiment did not demonstrate it.	Trama (1954)
	Calcium sulfate	Lepomis macrochirus	BSA	_	2,980 (NTE)	a d e f	Comment same as above.	Trama (1954)
CHEM	Calcium sulfate	Gambusia affinis	BSA	-	56,000 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
ICAL	Calcium sulphate	Biomorpholaria a. alexandrina	BSA	-	(O)	а	The degree of tolerance for vector snails of biharziasis chemicals is somewhat dependent upon temperature.	Gohar and El-Gindy
ა გ	sulphate	Bulinus			(O)		The tolerance for Bulinus, Lymnaea, and Biomphalaria	(1961)
ND MI		truncatus Lymnaea caillaudi			(O)		was up to saturation.	
CHEMICALS AND MIXTURES OF (	Calcium sulfate	Rainbow trout	BSCHA	-	(O)	<u>a c e</u> 	This report concludes that fish in contact with a system con- taining 10,000 ppm of gypsum could survive for a day, or could survive in a system containing 6.820 ppm for three weeks. Also, that concentrations of 3.163 ppm or less should be safe for four weeks or much longer periods.	Herbert and Wakeford (1962)

CHEMICA Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Calcium Calcium Sulphate	Nitzschia linearis Lepomis macrochirus	BSA	_	3,200 (T5A) 2,980 (T4A)	9 C e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Capric acid Caproic Caproic acid	Lepomis macrochirus	BSA	-	(O)	_	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Chemical is only slightly soluble in water. No toxicity data were obtained.	Dowden and Bennett (1965)
Caproic acid	Lepomis macrochirus	BSA	-	150 - 200 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Caprylic acid	Lepomis macrochirus	BSA	_	(O)	_	Comment same as above except that compound was very insoluble in water. No toxicity data were obtained.	Dowden and Bennett (1965)
Carbon chloroform extract (CCE)	Trout Golden Shiner Sunfish	BSA	_	36 (T1A) 32 (T2A) 28 (T4A) 24 (T5A) 59 (T1A) 52 (T2A) 39 (T4A) 33 (T5A) 56 (T1A) 49 (T2A) 45 (T4A) 39 (T5A)	a c d e f i m	The objects of this investigation were the recovery of or- ganic micropollutants from subsurface and surface Missouri waters, characterization and identification of these substances, and evaluation of their toxic effects, both acute and long-term, in order to develop methods for their destruction or removal.	Smith and Grigoropoulos (1968)
Carbon chloroform extract (CCE)/ carbon alcohol extract (CAE) 1/1.48		BSA	_	130 (T1A) 125 (T2A) 95 (T4A) 82 (T5A) No effect up to 305 (T5A) 166 (T1A) 144 (T2A) 115 (T4A) 103 (T5A)	acdefim	Comment same as above.	Smith and Grigoropoulos (1968)
Carbon chloroform extract (CCE)/ carbon alcohol extract (CAE) 1/1.56		BSA	-	138 (T1A) 130 (T2A) 96 (T4A) 92 (T5A) No effect up to 240 (T5A)	acdefim	Comment same as above,	Smith and Grigoropoulos (1968)

	Carbon dioxide	Trout	BSA	-	(0)	a c	No quantitative data are reported. 30 ppm of nitrogen was added as ammonium chloride. Carbon dioxide in concen- trations up to 30 ppm reduced the toxicity of the ammonia by lowering the pH of the water. Concentrations of 60 ppm of CO <sub>2</sub> were toxic but not lethal when the concen- tration of dissolved oxygen was low. A concentration of 240 ppm of CO <sub>2</sub> was lethal to trout in little more than one hour.	Herbert (1955)
	Carbon dioxide plus ammonia	Rainbow trout	BSA	-	(0)	<u>a e m n</u>	The reduction of toxicity of ammonia solutions by the addi- tion of carbon dioxide was due to lowering the pH of the solution. $60-240$ ppm CO <sub>2</sub> in solution was toxic within 12 hr. 30 ppm ammonia nitrogen was toxic, but up to 30 ppm CO <sub>2</sub> increased fish survival time.	Alabaster and Herbert (1954)
	Carbon disulfide	Gambusia affinis	BSA	_	135 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Carbonic acid	Fish	BSA	-	6.5 x 10 <sup>-4</sup> М (К)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)
	Cetyldimethyl ammonium bromide plus alkylate ether alcohol	Cylindrospermum licheniforme (CI) Gleocapsa sp (G) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No num- ber indicates observation is for entire test period of 21 days): CI - NT G - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
CHEMICALS	Cetylpyridinum- bromide	Microcystis aeruginosa	L	-	2.0 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
ALS AND	Cetyltrimethyl- ammonium bromide	Microcystis aeruginosa	L	-	2.0 (K)	a, etc	Comment same as above.	Fitzgerald, et al (1952)
D MIXTURES	Chlorauric acid	Gasterosteus aculeatus	BSA	-	0.4 (K10)	-	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
OF CHEMI	Chloride plus fluoride	Rainbow trout	BSA	_	(0)	ai	When trout were exposed to 30 ppm CI <sup>-</sup> for 48 hours and then challenged with fluoride, the LC50 of the fluoride was 6 ppm. No exposure to CI <sup>-</sup> resulted in an LC50 of 22 ppm FI <sup>-</sup> .	Neuhold and Sigler (1962)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chlorinated benzene	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Cittorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	a	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No num- ber indicates observation is for entire test period of 21 days): CI = T Ma = T So = T (3), PT (21) Cv = T Gp = T Np = T	Palmer and Maloney (1955)
Chlorinated camphene (60 percent)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Comment same as above except that: CI – PT Ma – T (14), PT (21) So – PT (14), NT Cv – PT Gp – T (3) Np – PT (7)	Palmer and Maloney (1955)
Chlorine (from mono- and di- chloramines)	Salmo gairdnerii	BCFA	-	0.08 (T7A)	<u>ace</u>	The purpose of this paper was to investigate the toxicity of chlorine to the rainbow trout in solutions containing ammonia. The toxicity of residual chlorine was dependent upon the relative proportions of free chlorine and chloramines.	Merkens (1958)
Chlorine	Nais spp	BSA	_	1.0 (K)	a f	All tests were conducted in hard water. At 1.0 ppm of chlo- rine, 95% of the worms were killed after 35 minutes. There was considerable variation in chlorine tolerance below 2 ppm and contact times from 1-3 hours may be necessary for a complete kill.	Learner and Edwards (1963)
Chlorine	Chlorella pyrenoidosa	BSA	_	0.18 (O) 0.42 (O)	a c i	<ul> <li>At 0.18 ppm, 1,900 cells/mm<sup>3</sup> remained at the end of 4 days as compared with a count of 2,383 cells/mm<sup>3</sup> in controls.</li> <li>At 0.42 ppm, 500 cells/mm<sup>3</sup> remained at the end of 4 days as compared with a count of 2,383 cells/mm<sup>3</sup> in controls.</li> <li>Chlorine showed an inhibitory effect in 48 hr.</li> <li>Experiments were carried out in seven-liter containers of tap water.</li> <li>By using 0.2 ppm of free chlorine, one might expect not to reduce the numbers of algae appreciably but to keep the population constant.</li> </ul>	Kott, et al (1966)

3'-chloro-5- acetamidosali- cylanilide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (K 3 hr) 10.0 (K2)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
p-chlorobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	Comment same as above except that at 10 ppm this chemi- cal was not toxic to trout or goldfish.	Walker, et al (1966)
Chlorobenzene	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	29 (T4A) 20 (T4A) 45 (T4A) 44 (T4A)	<u>a</u> c d	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
Chlorobenzilate	Daphnia magna	BSA	-	1.4 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
Chlorobenzilate	Simocephalus serrulatus Daphnia pulex	BSA	-	0.550 (SB) 0.870 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. "Water Chemistry" (Unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
4'-chloro-5- bromo-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	-	0.1 (K2) 1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biologi- cal activity against fish. Meta nitro substitution on the salicylanilides are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
4'-chloro-2,5- dihydroxy diphenyl sulphone	Daphnia magna	BSA	_	28.9 (K2A)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
4 chlorohexyl- 2,6-dinitro- phenol, tech.	Lymnaeid snails	BSA	-	(O)		Each test container, 500-ml beaker, was filled with ditch water. 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	2'-chloro-5'- methyl-3-nitro- salicylanilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.7 (LD <sub>100</sub> ) 1.0 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howeli (1966)
S OF	2'-chloro-3- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	3.0 (K) (O)	Ditto	Comment same as above. 70 ppm killed 25%.	Starkey and Howell (1966)
n N	2'-chloro-5- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.9 (K) (O)	,	Comment same as above. 3.0 ppm killed 25%.	Starkey and Howell (1966)
	3'-chloro-3- nitrosalicyl- anilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	0.3 (K) (O)	,	Comment same as above. 0.9 ppm killed 25%.	Starkey and Howell (1966)
	3'-chloro-5- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	15.0 (K) (O)	"	Comment same as above. 15.0 ppm killed 25%.	Starkey and Howell (1966)
	4'-chloro-3- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.3 (K) (O)	"	Comment same as above. 0.7 ppm killed 25%.	Starkey and Howell (1966)
	1'-chloro-5- nítrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.5 (K) (O)	"	Comment same as above. 1.0 ppm killed 25%.	Starkey and Howell (1966)
'n	n-chlorophenol	Carassius auratus	BSA	_	70.5 to 219 (K 8 hr) 61.7 (O) 20.6 (O)	<u>a</u>	Temperature in test containers was maintained at 27 $\pm$ 0.2 C. Goldfish tested weighed between 2 and 4 g. m-chlorophenol, 61.7 mg per liter, killed 93% of the fish in 8 hr; 20.6 mg per liter killed 62% in 8 hr.	Gersdorff and Smith (1940)
0-	-chlorophenol	Carassius auratus	BSA	~	142 to 311 (K 8 hr) 104 (O) 82.8 (O) 10.0 (O)	<u>a</u>	Comment same as above except that o-chlorophenol, 104 mg per liter, killed 83% of the fish in 8 hr; 82.8 mg per liter killed 64% in 8 hr; and 10.0 mg per liter killed 20% in 8 hr.	Gersdorff and Smith (1940)

p-chlorophenol	Carassius						
	auratus	BSA	-	54.3 to 190 (K 8 hr) 47.5 (O) 12.7 (O) 6.3 (O)	<u>a</u>	Comment same as above except that p-chlorophenol, 47.5 mg per liter, killed 85% of the fish in 8 hr; 12.7 mg per liter killed 75% in 8 hr; and 6.3 mg per liter killed 54% in 8 hr.	Gersdorff and Smith (1940)
4'-chloro-2', 5'-dimethoxy- 3-nitrosali- cylanilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of sali- cylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
5'-chloro-3, 5-dinitro-2- benzanilide	Salmo gairdnerii Carassius auratus	BSA	-	(O) (O)	<u>a</u>	Comment same as above except that at 10 ppm the chemical was not toxic to trout. At 1.0 ppm, 1 out of 10 goldfish died. This may not be valid since at 10 ppm, no fish were killed.	Walker, et al (1966)
2'-chloro-3, 5-dinitro- benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	Comment same as above except that at 10 ppm this chemical was not toxic to trout or goldfish.	Walker, et al (1966)
3'-chloro-3, 5-dinitro- benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm the chem- ical was toxic to 7 out of 10 trout in 48 hours. No goldfish were killed at this and lower concentrations.	Walker, et al (1966)
3'-chloro-3,5- dinitro-o- benzotoluidide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K 3 hr) (O)	<u>a</u>	Comment same as above except that at 10 ppm the chemical was not toxic to goldfish. Precipitation occurred at 10 ppm.	Walker, et al (1966)
X 3'-chloro-3, 5-dinitro-p- benzotoluidide	Salmo gairdnerii Carassius auratus	BSA	-	(O) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm the chem- ical was toxic to 2 out of 10 trout in 48 hours. The chem- ical was not toxic to goldfish at 10.0 ppm.	Walker, et al (1966)
5'-chloro-3, 5-dinitro-3- benzotoluidide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K 3 hr) (K 3 min.) (O)	<u>a</u>	Comments same as above except that at 10 ppm the chem- ical was not toxic to goldfish.	Walker, et al (1966)
X 2'-chloro-3', 4'-dinitro- 5 salicylanilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K 3 hr) 1.0 K (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
Chloroform	Pygosteus pungitius	BCF	-	(O)	<u>a</u>	A 1/2000 solution anaesthetized or killed very rapidly. 1/5000 and 1/10000 induced an avoidance reaction in the fish.	Jones (1947)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
S AND MIXTURES	Chloroform	Sewage organisms	BOD	_	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams de- picting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
ES OF CHEMICALS	3'-chloro- 3-hydroxy- benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K2) (O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 10.0 ppm, the chemical was toxic to 7 out of 10 goldfish at 48 hours.	Walker, et al (1966)
	4'-chloro-3- hydroxybenz- anilide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (К2) (О)	<u>a</u>	Comment same as above except that at 10.0 ppm the chem- ical was toxic to 2 out of 10 goldfish in 48 hours.	Walker, et al (1966)
	2'-chloro-2- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	-	(O) (O)	<u>a</u>	Comment same as above except that this chemical was not toxic to trout or goldfish at 10 ppm.	Walker, et al (1966)
	3'-chloro-2- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (K2) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm the chem- ical was toxic to 6 out of 10 goldfish at 48 hours.	Walker, et al
	2'-chloro-3- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (K2) (O)	<u>a</u>	Comment same as above except that at 10 ppm the chem- ical was toxic to 1 out of 10 fish in 48 hours.	Walker, et al (1966)
	2'-chloro-4- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	Comment same as above except that at 10 ppm this chem- ical was not toxic to trout or goldfish.	Walker, et al (1966)
	3'-chloro-3- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K 3 hr) 10.0 (K2)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
	3'-chloro-4- nitrobenz- anilide	Salmo gairdneri Carassius auratus	BSA	-	(O) (O)	<u>a</u>	Comment same as above except that no fish were killed at 10 ppm.	Walker, et al (1966)

4'-chloro-2- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (К2) 10.0 (К2)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
5'-chloro-4- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K2) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm the chem- ical was toxic to 6 out of 10 goldfish in 48 hours.	Walker, et al (1966)
3'-chloro-3- nitro-p-benzo- toluidide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	Comment same as above except that chemical precipitated at 10 ppm, and the chemical was not toxic to trout. At 0.1 ppm the chemical was toxic to 1 out of 10 goldfish.	Walker, et al (1966)
5'-chloro-2- nitrophenol (free phenol)	Petromyzon marinus Salmo	BSA BSA	<b>-</b> 	3 (K 100%) 5 (K 10%)	<u>a</u> a	Mortality occurred in approximately 24 hr. This was a study on controlling sea lamprey larvae.	Ball (1966)
	gairdnerii S. trutta	BSA	_	5 (K 10%)	 <u>a</u>		
Chloronitro- propane	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	_	80 (K) 80 (K) 80 (K) 80 (K)	_ a	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants.	Ukeles (1962)
5'-chloro-3- nitro-o-sali- sylanilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K3A) 10.0 (K3A)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogen and their relative position(s) in the molecule.	Walker, et al (1966)
2'-chloro-5- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K 3 hr) 10.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)
3'-chloro-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K2) 10.0 (K 3 hrs) 10.0 (K 3 hrs) 1.0 (K2)	<u>a</u>	Comment same as above.	Walker, et al (1966)
4'-chloro-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K 3 hr) 0.1 (K2) 1.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)

CHEMICALS	Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
S AND MIXTURES OF CHEMIC	4'-chloro-5- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K2) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
AICALS	3'-chloro-2- nitro-o-benz- otoluidide	Salmo gairdnerii Carassius auratus	BSA	-	(O) (O)	<u>a</u>	Comment same as above except that this chemical was not toxic to trout or goldfish at 10 ppm.	Walker, et al (1966)
	3'-chloro-3- nitro-o- salicylotolu- idide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K2) 10.0 (K 3 hr) 10.0 (K2)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
	6'-chloro-3- nitro-o-salicy- lotoluidide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (K2) (O)	<u>a</u>	Comment same as above except that this chemical was not toxic to goldfish at 10 ppm.	Walker, et al (1966)
	4'-chloro-3- nitro-o-salicyl- otoluidide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K 3 hr) 10.0 (K 3 hr)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
	2'-chloro-3- nitro-p-sa- licylotoluidide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K 3 hr) 1.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)
	Chlorophenol (meta)	Minnows	BSA	-	18.0 (T1A)	e	In the halophenols, the ortho was less toxic than the meta or para. All of the monohalophenols were less toxic than the 2,4,6-trihalophenols. Some data on biodegradability of halophenols were presented.	Ingols and Gaffney (1965)
	o-chloro- phenol	Lepomis macrochirus	BSA	-	8.1 (T2A)	acdefgio 	Assays are completely described and autopsy data are reported.	Lammering and Burbank (1961)
	o-chloro- phenol	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	12 (T4A) 10 (T4A) 14 (T4A) 23 (T4A)	<u>a</u> cd	Most fish survived at test concentrations of about one half or slightly more of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)

Chlorophenol							
(ortho)	Minnows	BSA		58 (T1A)	e	In the halophenols, the ortho was less toxic than the meta or para. All of the monohalophenols were less toxic than the 2,4,6-trihalophenols. Some data on biodegradability of halophenols were presented.	Ingols and Gaffney (1965)
p-chlorophenol	Hyborhynchus notatus	BSA	-	(O)	_	Fish in aquaria were trained to detect and distinguish between phenol and p-chlorophenol at levels as low as 0.0005 ppm. The fish could also distinguish o-chlorophenol from the two other compounds. The training method is described.	Hasler and Wisby (1949)
Chlorophenol (para)	Minnows	BSA	-	14 (T1A)	e	In the halophenols, the ortho was less toxic than the meta or para. All of the monohalophenols were less toxic than the 2,4,6-trihalophenols. Some data on biodegradability of halophenols were presented.	Ingols and Gaffney (1965)
3-(p-chloro- phenol)-1,1- dimethyl- urea	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): Ci - PT (7), T (21) Ma - T So - T (7), PT (21) Cv - T (3), PT (14) Gp - T Np - T	Palmer and Maloney (1955)
Bis (p-chloro- phenoxy)	Bluegill	BSA	_	(O)	-	No mortality occurred at 0.05 ppm and very low mortality at 0.10 ppm. All fish died when the concentration was	Linduska and Surber
methane						0.2 ppm.	(1948)
methane P-chloro- phenyl-p- chloroben- zenesulfamate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	a	0.2 ppm. Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T $\approx$ toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): Cl - PT (3) Ma - PT (14) So - PT (7) Cv - NT Gp - PT (7) Np - T (3)	
P-chloro- phenyl-p- chloroben-	licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np) Pimephales promelas	L BSA	_	24 (T4A)	a a c d	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = PT (3) Ma = PT (14) So = PT (7) Cv = NT Gp = PT (7) Np = T (3) Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made	(1948) Palmer and Maloney (1955) Pickering and Henderson
P-chloro- phenyl-p- chloroben- zenesulfamate	licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np) Pimephales promelas Lepomis macrochirus		_	24 (T4A) 42 (T4A)		Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = PT (3) Ma = PT (14) So = PT (7) Cv = NT Gp = PT (7) Np = T (3) Most fish survived at test concentrations of about one half,	(1948) Palmer and Maloney (1955) Pickering and
P-chloro- phenyl-p- chloroben- zenesulfamate 3-chloro-	licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np) Pimephales promelas Lepomis		_	24 (T4A)		Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = PT (3) Ma = PT (14) So = PT (7) Cv = NT Gp = PT (7) Np = T (3) Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made	(1948) Palmer and Maloney (1955) Pickering and Henderson

	Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
4,	, chloro-o- toloxy- acetic acid	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = T (3) Ma = NT So = NT Cv = NT Gp = T (3) Np = T (3)	Palmer and Maloney (1955)
	hromic acid	Daphnia magna	BSA	-	<0.6 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to im- mobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	hromic chloride	Daphnia ma <b>g</b> na	BSA	-	<<3.6 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	hromic sulfate	BOD	L	-	1.0 (O)	j	"Toxicity" is expressed as 10 percent reduction in oxygen utilization.	Ingols (1955)
	hromic sulfate	Sewage organisms	BOD	-	(O)	_	Chromate ion is less toxic than chromic. 1.0 ppm produced a 10% oxygen depletion as compared to a control, and 10 ppm produced a 30% depletion.	Ingols (1954)
	hromic sulfate	Sewage organisms	BOD	-	117 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	hromic sulfate	Daphnia magna	BSA	-	0.1 (T1A) 0.03 (T2A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
s s	hromic sulfate plus sodium di- chromate	Lymnaea sp (eggs)	BSA	-	0.2 (T1A)	a c	Comment same as above.	Dowden and Bennett (1965)
	hromium, nexavalent	Bluegill, pumpkinseed sunfish, and orangespots	F	Wood- stock, III.	(0)	c	At chromium concentrations above 50 ppm, the range of survival was such that no general curve could be applied to the data plotted on the chart.	Klassen, et al (1948)

	Chromium	Chlorococcum variegatus C. humicola Scenedesmus obliquus Lepocinclis steinii	L	-	6.4-16.0 (O) 3.2-6.4 (O) 3.2-6.4 (O) 0.32-1.6 (O)	а	Chromium as dichromate was evaluated in two different tests. The concentrations reported are a range which completely inhibited growth for 56 days. Concentra- tions as low as 0.0001 to 0.032 ppm stimulated growth up to 33 days of <i>C. humicola</i> , <i>S. obliquus</i> , and <i>L. steinii</i> . Data for a flagellate and two diatoms are also presented.	Hervey (1949)
	Chromium (hexavalent)	Salmo gairdnerii	L	_	2.5 (O)	<u>a</u>	For accumulation studies, fish were exposed for periods up to 24 days. For elimination studies, fish were exposed for 12 days, then placed in fresh water from 5 to 25. Chro- mium in the blood never exceeded the concentration of the surrounding water. All other tissues except muscle accumulated concentrations in excess of that in the water. Chromium was eliminated rapidly from blood, liver, stomach, pyloric caeca, and posterior gut. The spleen lost little of its chromium even after being in fresh water for 25 days. The kidney lost about 50% of its chromium in 25 days of fresh water exposure.	Knoll and Fromm (1960)
	Chromium (hexavalent)	Lepomis macrochirus	BSA	-	110 (T4A)	<u>a</u> cdfq	Soft water was used. Alkalinity and hardness significantly reduced the toxicity of hexavalent chromium.	Trama and Benoit (1960)
A-37	Chromium (as chromate)	Salmo gairdnerii	BSCH	-	5 (K15)* 10 (K15)** 12.5 (K15)** * 40% kill **80% kill	<u>a</u>	This study is concerned with the measurement of chromium in trout before and after exposure. Chromium uptake is passive, and the amount accumulated is dependent on the concentration in water and duration of exposure.	Fromm and Stokes (1962)
	Chromium	Salmo gairdnerii	L	-	2.5 (O)	-	Trout were exposed to 2.5 ppm of chromium as chromate in tap water for one week. The <i>in vitro</i> glucose transport by gut segments from these animals was compared to that of segments from untreated fish. The values from the treated animals was 40 percent lower than the controls.	Stokes and Fromm (1965)
CHE	Chromium	Rainbow trout	FR	Scotland	20 (NTE)	acefim	This work represents an extension of laboratory studies of the toxicity of complex effluents to investigations of rivers.	Herbert, et al (1965)
CHEMICALS A	Chromium	Gasterosteus aculeatus	BSA	-	1.0 (O)	<u>a c e</u>	This is a discussion of a bioassay method using stickleback fish and spectrophotometric determinations of the chem- icals evaluated. The number listed is said to be the "toxic limit" for the fish.	Hawksley (1967)
AND MIXTURES	Mixture: Chromium (a)- naphthenic acids (b)-cyanide (c) Mixture	Lepomis macrochirus	BSA	-	(a) 0.019 (T4A) (b) 4.74 (T4A) (c) 0.26 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilu- tion water.	Cairns and Scheier (1968)
IES OF CHEMICALS	Chromium chloride	Sewage organisms	BOD	-	0.18 (O)	-	Various metal salts were studied in relation to how they af- fected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treatment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chem- ical was tested in an unbuffered system.	Sheets (1957)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND	Chromium chromate	Lepomis macrochirus	BSA	_	170 ( <b>T4</b> A)	<u>a</u> cdfq	Soft water was used. Alkalinity and hardness significantly reduced the toxicity of this form of chromium.	Trama and Benoit (1960)
MIXTURES	Chromium dichromate	Lepomis macrochirus	BSA	_	113 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
S OF CHEMICALS	Chromium oxide	Sewage organisms	BOD	_	4.0 (O)	-	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Sheets (1957)
ST S	Chromium potassium sulfate	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	(S) 5.07 (T4A) (H) 67.4 (T4A) (S) 7.46 (T4A) (H) 71.9 (T4A) (S) 4.10 (T4A) (S) 3.33 (T4A)	c d e f	(S) Soft water (H) Hard water Values are expressed as mg/l of chromium.	Pickering and Henderson (1965)
	Chromium sulfate	Gasterosteus aculeatus	BSA	-	1.2 (K10)	-	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
	Citric acid	Daphnia magna	BSA	_	153 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson {1944}
	Citric acid	Biomorpholaria a. alexandrina Bulinus truncatus Lymnaea	BSA	-	1200 (K1A) 1000 (K1A) 800 (K1A)	а	The degree of tolerance for vector snails of biharziasis chem- icals is somewhat dependent upon temperature. The tem- perature at which (K1A) occurred was 27 C for <i>Bulinus</i> and <i>Biomphalaria</i> and 28 C for <i>Lymnaea</i> .	Gohar and El-Gindy (1961)
		caillaudi			000 ((()))			
	Cobalt	Lebistes reticulatus Bufo valliceps	L	-	100.0 (K) 100.0 (K)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are	Shaw and Grushkin (1967)
		(tadpoles) Daphnia magna			50.0 (K)		the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide – a treatment that does not lend itself to tabulation. The cation toxicity cited is only an ap- proximate concentration interpolated from a graph. Time of death was not specified.	

Cobalt	- · ·						
chloride	Daphnia magna	BSA	-	<3.1 (S)	<u>a</u>	Lake Eria water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
Cobalt chloride	Sewage organisms	BOD	-	64.0 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depict- ing the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Cobalt chloride	Limnaea palustris (eggs)	BSA	-	4 x 10 <sup>-5</sup> M (K1)	<u>a</u> c	Toxicity is given in molar concentrations for maximum direct mortality (kill) in 4 hours.	Morrill (1963)
Cobaltous chloride	Daphnia magna	BSA	-	<26 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to im- mobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Cobalt nitrate	Gasterosteus aculeatus	BSA	-	10 (K10)	~	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
Copper	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of temperature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In water distilled from a copper still with block-tin leads, the fish survived 352 to 597 minutes – perhaps the effect of copper.	Powers (1918)
Cu	Nemacheilus barbatulus	ВСН ВСН	England England	0.28 (K) 0.20-0.30 (K)	-	Fresh water input was through Cu pipes into an aquarium. All fish died within 24 hours at concentrations of 0.20 ppm and above.	Mackereth and Smyly (1951)
Copper ion (copper chloride and copper sulfate	Lepomis macrochirus	BSA	-	0.74 (T4A) 0.94 (T2A)	acde	Modified Chu 14 diluent made of distilled water was used with aeration toxicity of copper ion was found to be de- pendent upon pH. Below pH 5.3, all copper is in solution, above this the copper precipitates and is less toxic.	Trama (1954)
Copper	Sewage organisms	BOD	_	(O)	_	Copper was more toxic than zinc in all concentrations from 0.1 to 10.0 ppm. The presence of the element could result in errors in BOD tests. At 1.0 ppm the oxygen demand in percent of the control was 65%.	Ingols (1956)
	chloride Cobalt chloride Cobalt chloride Cobaltous chloride Cobalt nitrate Copper Cu Cu Copper ion (copper chloride and copper sulfate	chloridemagnaCobalt chlorideSewage organismsCobalt chlorideLimnaea palustris (eggs)Cobaltous chlorideDaphnia magnaCobaltous chlorideDaphnia magnaCobalt nitrateGasterosteus aculeatusCopper CopperCarassius carassius carassiusCuNemacheilus barbatulusCopper ion (copper chloride and copper sulfateLepomis macrochirusCopper sulfateSewage	chlorideDaphmaDOXCobalt chlorideSewage organismsBODCobalt chlorideLimnaea palustris (eggs)BSACobaltous chlorideDaphnia magnaBSACobalt nitrateGasterosteus aculeatusBSACobalt nitrateGasterosteus aculeatusBSACopperCarassius carassiusBSACuNemacheilus barbatulusBCH BCHCopper ion chloride and copper sulfateLepomis macrochirusBSACopperSewageBOD	chloride magna bur - chloride Sewage BOD - Cobalt Sewage Organisms BOD - Cobalt Limnaea BSA - chloride palustris (eggs) Cobaltous Daphnia BSA - chloride magna BSA - Cobalt Gasterosteus BSA - chloride Gasterosteus BSA - Copper Carassius Carassius BSA - Copper Sewage BOD -	chloride     magna     DON     Co. 1 (S)       Cobalt chloride     Sewage organisms     BOD     -     64.0 (TC <sub>50</sub> )       Cobalt chloride     Limneea pelustris (eggs)     BSA     -     4 x 10 <sup>-5</sup> M (K1)       Cobalt chloride     Daphnia magna     BSA     -     4 x 10 <sup>-5</sup> M (K1)       Cobaltous     Daphnia magna     BSA     -     26 (O)       Cobaltous     Daphnia magna     BSA     -     26 (O)       Cobalt chloride     Gasterosteus aculeatus     BSA     -     10 (K10)       Copper     Carassius carassius     BSA     -     (O)       Cu     Nemacheilus barbatulus     BCH     England England     0.28 (K) 0.20-0.30 (K)       Copper ion chloride and copper sulfate     Lepomis macrochirus     BSA     -     0.74 (T4A) 0.94 (T2A)       Copper     Sewage     BOD     -     (O)	chloride       magna       DON       Col. (G)       a         Cobalt chloride       Sewage organisms       BOD       -       64.0 (TC <sub>50</sub> )       a         Cobalt chloride       Limnaea palustris (eggs)       BSA       -       4 x 10 <sup>-5</sup> M (K1)       a c         Cobalt chloride       Daphnia magna       BSA       -       4 x 10 <sup>-5</sup> M (K1)       a c         Cobalt chloride       Daphnia magna       BSA       -       <26 (O)	ehloride         magan         EXA         Curves         Example         Example <thexample< th=""> <thexample< th=""> <thexample< td=""></thexample<></thexample<></thexample<>

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Copper	Chlorella vulgaris	L	-	(O)	a c e	This was a respiration study using a shake culture technique. 10 <sup>-1</sup> M copper sulfate was not inhibitory for 7-20 hours. Concentrations of 10 <sup>-3</sup> M copper sulfate were toxic to un- shaken cultures.	Hassall (1962)
Copper	Nereis sp Carcinus maenas Leander squilla	BSA	-	1.5 (K2-3) 0.5 (K4) (O) (O)	а	The threshold of copper for <i>Nereis</i> worms was about 0.1 ppm. The copper toxicity threshold for the shore crab was 1-2 ppm. The copper toxicity threshold for prawns was below 0.5 ppm.	Raymont and Shields (1964)
Copper	Salmo salar	BCFA	-	0.034 (T1A)	<u>a</u> cf	The laboratory water in which the experiment was performed contained 3 $\mu$ g/liter of zinc, as judged by analyses over several years, and 2 $\mu$ g/liter of copper. Lethal concentrations of mixtures activities or three times as fast as the metals singly, a somewhat greater potentiation than was found in the previous tests with salmon.	Sprague (1965)
Copper	Rainbow trout	FR	Scotland	0.8 (T2)	aceflm	This work represents an extension of laboratory studies of the toxicity of complex effluents to investigations of rivers.	Herbert, et al (1965)
Copper	Gasterosteus aculeatus	BSA	-	0.02 (O)	<u>a c e</u>	This is a discussion of a bioassay method using stickleback fish and spectrophotometric determinations of the chemi- cals evaluated. The number listed is said to be the "toxic limit" for the fish.	Hawksley (1967)
Copper	Orconectes rusticus	BCFA	_	3.0 (T4A) 1.0 (T1A) 1.0 (K <sub>6</sub> )(T6A) 1.0 (T <sub>6</sub> )(T6A)	<u>a</u> c <u>e</u> f	<ul> <li>All experiments were conducted at 20 C.</li> <li>Crayfish in the intermolt adult stage.</li> <li>Adult crayfish.</li> <li>Juvenile crayfish.</li> <li>Recently hatched young which remained clinging to pleopods of the female during the first molt.</li> <li>An acute toxicity threshold existed between 0.6 and 0.125 mg/l for newly hatched young. At a concentration of 1 mg/l, 50% mortality among newly hatched young was reached with an exposure time of 1/50th required for adults.</li> </ul>	Hubschman (1967)
Copper	Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	BSA	_	1.0 (К) 0.1 (К) 0.1 (К)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product constant of the metal sulfide – a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)

	Copper	Pimephales promelas	BCFCH	_	0.43 (T4A)	acdef	The paper discusses growth rate, number of spawnings, num- ber of eggs produced and hatchability of eggs in water con- taining 4.4 to 95 ppm copper. Results indicated that the sublethal concentrations of copper affecting growth and reproduction lies between 3 and 7 percent of the 96-hr median tolerance limit.	Mount (1968)
	Copper	Salmo gairdnerii	BSA	-	0.4 to 0.5 (T2A)	acdef	The concentration killing a half batch of fish in 2 days pro- vides a reasonable estimate of the threshold concentration. The lethality of this chemical depends upon the total hardness and dissolved oxygen concentration.	Brown (1968)
	Copper (Cu <sup>++</sup> )	Lepomis macrochirus	BSA	-	1.25 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
	Copper (a)- acetic acid (b)- acetaldehyde (c)-acetone (d) mixture	Lepomis macrochirus	BSA	-	(a) 1.04 (T4A) (b) 26.0 (T4A) (c) 5.2 (T4A) (d) 5.2 (T4A)	a c d e	Comment same as above.	Cairns and Scheier (1968)
	Copper para- amino benzoate	Balanus eberneus	BSA	-	0.9 (O)	_	The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
	Copper carbonate (basic)	Balanus balanoides Balanus eberneus	BSA	-	0.41 (O) 0.28 (O)	-	The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
	Copper citrate	Balanus balanoides Balanus eberneus	BSA	_	0.60 (O) 0.55 (O)	-	Comment same as above.	Clarke (1947)
_	Copper cyanide complex	Lepomis macrochirus (juveniles)	BSA	-	4.0 (O)	<u>a</u> cdf <u>p</u>	For the concentration given, the median resistance time was 226 minutes.	Doudoroff, et al (1966)
CHEMICALS AND MIX	Copper cyanide complex Sodium cyanide (533 ppm CN <sup>-</sup> ) and Cupric sulfate (427 ppm Cu)	Pimephales promelas	BSA	-	1.5 (T4) CN <sup>-</sup> 1.2 (T4) Cu	<u>ac</u>	Synthetic soft water was used. Toxicity data given as number of test fish surviving after exposure at 24, 48, and 96 hr. TL <sub>m</sub> values were estimated by straight-line graphical inter- polation and given in ppm CN <sup>-</sup> .	Doudoroff, et al (1956)
TURES	Copper disodium versenate	Channel catfish (fingerlings)	BSA	-	1881 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES OF CHEMICALS	Copper naphthenate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th. and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - T (3) So - PT (3) Cv - PT (3) Gp - T (7), PT (14) Np - NT	Palmer and Maloney (1955)
ST A:	Copper nitrate	Gasterosteus aculeatus	BSA	-	1.0 (T6.5A)	a c	Death of the fish resulted from an interaction between the metallic ion and the mucus secreted by the gills. Coagulated mucus formed on the gill membranes and impaired respira- tion to such a degree that the fish asphyxiated.	Jones (1938)
	Copper salicylate	Balanus eberneus	BSA	-	0.90 (O)		The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
	Copper salts	Salmo gairdnerii	BSA	_	(O)	<u>a e</u>	This is a study of the effect of varying dissolved oxygen con- centration on the toxicity of selected chemicals. The toxicity of heavy metals, ammonia, and monohydric phenols increased as the dissolved oxygen in water was reduced. The most obvious reaction of fish to lowered oxy- gen content is to increase the volume of water passed over the gills, and this may increase the amount of poison reach- ing the surface of the gill epithelium. The concentration of the chemical in the water was not specified.	Lloyd (1961)
	Copper salt plus citrate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T (3) Ma - T (3) So - PT (7) Cv - T (3) $Gp^- T (3)$ Np - T (3)	Palmer and Maloney (1955)
	Copper sodium citrate	Artemia salina Acartia clausi	BSA	-	0.005 (O) 0.01 (O)	ас	All tests were conducted in seawater, Toxicity values reported are relative to that of mercuric chloride expressed as unity. Mechanism of action is discussed, as well as synergistic action	Corner and Sparrow (1956)
		Elminus modestus			0.002 (0)		of two poisons administered simultaneously.	

Copper tartrate	Balanus balanoides	BSA	-	0.58 (O)	_	The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
Copper and zinc	Atlantic salmon	FR	Canada	(O)	f	"Toxicity index" for copper and zinc combined was de- scribed in connection with disturbed salmon migration. Toxicity index $\geq$ 1.0 indicates lethality to "young salmon after long exposure". A toxicity index of 0.15 or 15% of lethal concentration of copper and zinc seemed to be the maximum safe level for salmon migration.	Sprague (1964)
Copper and zinc	Salmo salar	BSA	_	0.048 Cu (O) 0.600 Zn	а	The values given are for an ILL (incipient lethal level) and in this instance only in water of 20 mg/liter of hardness. Concentrations above this are lethal in about one day. These values were determined by bioassay. Salmon parr in the laboratory avoided less than one tenth of incipient lethal levels. Avoidance thresholds were 0.09 ILL of zinc, 0.05 ILL of copper and 0.02 ILL of equitoxic mixtures. In equitoxic mixtures of these compounds, the ILL was additive.	Sigler, et al (1966)
Copper chloride	Carassius carassius	BSA	_	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of temper- ature, effect of dissolved oxygen, the efficiency of the gold- fish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 0.66N, fish survived 78 minutes; at a concentration of 0.0000011N, fish survived 300 minutes — truly a very wide variation.	Powers (1918)
Copper chloride (tech)	Bluegill	BSA?	-	0.980 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
Copper chloride	Nitzschia linearis Lepomis macrochirus	BSA	_	0.795-0.815 (T5A) 1.25 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Copper sulfate	Algae zooplankton		Lakes in Wisc.	(O)	aegin	Copper sulfate was applied when deemed necessary to control algae (0.50 pounds of copper sulfate per million gallons of water). Applications of copper sulfate were made as re- quired over an eleven-year period. Zooplankton was not effected by these applications. The spray applied for control of algae also kept fish fungal diseases under control.	Domogalla (1935)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
Copper sulfate	Morone americana Perca flavescens All fish Mesocyclops obsoletus Macrobdella decora Nymphaea Juncus Pontederia Scirpus Eriocaulon Potamogeton Algae	FL	4 lakes, Nova Scotia	1 (K) 1 (K) 3 (K) 3 (SB) 3 (SB) 3 (NTE) 3 (NTE) 3 (NTE) 3 (NTE) 3 (NTE) 3 (NTE) 3 (NTE) 3 (NTE) 3 (NTE)	acdf	The work was undertaken to test the feasibility of utilizing poisons as a direct means of studying the production of fish in streams and lakes. Caution must be used to prevent irreparable damage by indiscriminate poisoning.	Smith (1939)
Copper sulfate	Morone americana Perca flavescens All fish Mesocyclops obsoletus Macrobdella decora Nymphaea Juncus Pontederia Scirpus Eriocaulon Potamogeton Algae	FL	4 lakes, Nova Scotia	1 (K) 1 (K) 3 (K) 3 (SB) 3 (SB) 3 (SB) 3 (NTE) 3 (	a c d f	Comment same as above.	Smith (1939)
Copper sulfate	Smallmouth black bass <i>Chara</i> sp	FL	Leetown, Va.	2.0 (0)	d	Treatment of a series of ponds resulted in control of <i>Chara</i> spp but no or slight fish kill due to copper sulfate. Some kill occurred because of suffocation caused by decaying vegetation.	Surber and Everhart (1950)
Copper sulfate	Pygosteus pungitius	BCF	-	(O)	a c	Fish were exposed to 0.1, 0.04, and 0.01N copper sulfate. pH of the solutions was 5.0, 5.4, and 5.8. Survival times were 55, 62, and 75 minutes, respectively.	Jones (1947)
Copper sulfate (anhydrous)	Lymnaeid snails	BSA	-	1.0 (K1A)	-	Each test container (500-ml beaker) was filled with ditch water.	Batte, et al (1951)

Copper sulfate	Tendipes plumosus Pisidium idahoense and other bottom- dwelling organisms	FL & BSA	Wisc.	(O)	_	The bottom muds of Lake Morona contained up to 480 milli- grams of copper per kilogram of mud on a dry-weight basis. Lakes Nagawicka and Pewaukee contain up to 22 and 55, respectively. All contained thriving populations of aquatic organisms despite years of CuSO <sub>4</sub> application for algal con- trol. From laboratory bioassays of muds containing CuSO <sub>4</sub> , it was concluded that 9,000 parts per million copper on a dry-weight basis precipitated and accumulated in bottom muds was toxic to bottom organisms. From the results of these studies, it is indicated that differences occurring in the population density of bottom organisms in the four lakes studied are due to ecological variables within these separate bodies of water.	Mackenthun and Cooley (1952)
Copper sulfate	BOD	L	-	1.0 (O)	j	"Toxicity" is expressed as 39 percent reduction in oxygen utilization.	Ingols (1955)
Copper sulfate	Microcystis aeruginosa	L	-	100 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Copper sulfate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = PT (7), T (14) Ma = T (3) So = PT (7) Cv = T (3) Gp = T (3) Np = T (3)	Palmer and Maloney (1955)
Copper si (with st lizing ag CHEMICALS AND	abi- licheniforme (Cl)	L	_	2.0 (0)	<u>a</u>	Comment same as above except that CI - T (3) Ma - T (3) So - PT (3) Cv - T (3) Gp - T (3) Np - T (3)	Palmer and Maloney (1955)
Copper sulfate	Pimephales promelas	BSA	-	0.18 (T4A)	acd e f	Toxicity to 30 species of algae is also presented. CuSO <sub>4</sub> was algicidal in the range 0.5 to 2.0 ppm.	Palmer and Maloney (1956)
Copper sulfate	Sewage organisms	BOD	-	0.4 (0)	-	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.0. Solutions were renewed every 12 hours.	Sheets (1957)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location <sup>(2)</sup>	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Copper sulfate	Gambusia affinis	BSA	_	84 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Copper sulphate	Salmo gairdneri (fry)	BSA	-	3.8 (T1A) 10 (O)	acefip	Five hatchery troughs were employed with 6 Imperial gallons (27.276 liters) of hatchery water. The water used in the experiments was reportedly typical of Inyanga Rhodesia trout streams and dams. Concentra- tions of 10 ppm of copper sulphate caused 90-100% mortality.	Turnbull-Kemp (1958)
Copper sulfate	Salvelinus fontinalis x Salmo trutta Notemigonus crysoleucas Micropterus salmoides Lepomis macrochirus	FPA	N.Y.	1.0 (S23) 1.0 (К) 1.0 (S23) 1.0 (S23)	a c d	Conventional farm ponds were used having an average surface area of 0.3 acre and a maximum depth of 7-9 ft. Toxicity (in ppm) to fish as maximum safe concentration (S) for 23 days was determined. Concentration of 0.5 ppm was required to control algae.	Eipper (1959)
Copper sulfate	Sewage organisms	BOD	-	21 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Copper sulfate	Pimephales promelas Lepomis macrochirus	BSA	_	(H) 1.4 (T4A) (S) 0.05 (T4A) (H) 10 (T4A) (S) 0.2 (T4A)	acd f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
Copper sulfate	Limnodrilus hoffmeisteri Gyraulus circumstriatus Physa heterostropha Tendipes decorus	BSA	_	0.40 (T4A) 0.425 (T4A) 0.27 (T4A) 1.0 (K 60%) 0.032 (K 40%)	<u>a</u> cdi	Hard water only was used in this study for all but <i>T. decorus</i> which was also studied in soft water.	Wurtz and Bridges (1961)
Copper sulfate	Rana pipiens	BSCH	-	16 (K)	<u>a c</u>	CuSO <sub>4</sub> was toxic to this frog at various temperatures in concentrations $\geq$ 0.0015 percent.	Kaplan and Yoh (1961)
Copper sulfate	Physa heterostropha	BSA	-	0.56 (T1A)	<u>a</u> cf	These tests were conducted in hard and soft water. Data indicated small if any differences in toxicity of copper sulfate due to water hardness.	Wurtz (1962)

\_\_\_\_

APPENDIX A

	Copper sulfate	Microcystis sp Zooplankters Copepods Cladocerans Rotifers Chaoboridae Ostracods etc.	FL	Auburn, Aia.	0.5-0.8 (O)	d	In a series of ponds, CuSO <sub>4</sub> at the indicated concentration range reduced the growth of <i>Microcystis</i> spp by as much as 95 percent in 5-20 days. This reduction lasted for as long as 30 days in some cases. According to the authors, generally there was an inverse relation between the abundance of <i>Micrycystis</i> and the number of zooplankters.	Crance (1963)
	Copper sulfate	Nais spp	BSA	_	1.0 (K)	a f	Around pH 7.0, copper was more toxic in soft than in hard water. At 1.00 ppm the average median survival time for the worms was reduced from 70 to 35 minutes. It is inter- esting that copper is less toxic at a pH of 4.0 than at 7.0.	Learner and Edwards (1963)
	Copper sulfate	Chlorella pyrenoidosa	L	-	20 (AS1)	-	Describes a bioassay method to differentiate between an algi- cide (AC) and an algistat (AS). The treated culture was sub- cultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)
>	Copper sulfate (Algeeclear) (Cuprose)	Microcystis aeroginosa Chlorella pyrenoidosa Anabaena circinalis Gloeotrichia echinulata Phormodinium inundatum	L	-	(0)	c e	Different sources of copper appeared to be equally effective as toxic agents for algae. The medium in which toxicity tests are carried out had a great influence on the toxicity of cop- per. It was pointed out that in copper compounds, the range in toxic action can vary from algicidal activity at concentra- tions of 0.05 to 0.4 ppm of CuSO <sub>4</sub> , or algistatic activity at 2 to 24 ppm of CuSO <sub>4</sub> with certain algae, to situations in which the growth of algae is only slightly inhibited by a con- centration of copper sulfate as high as 30 ppm.	Fitzgerald and Faust (1963)
L V	Copper sulfate	Gammarus Iacustris	BSA	_	1.5 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
	Copper (copper sulfate)	Salmo salar	BCF	-	0.048 (O)	<u>a</u> cdef	The experiments were carried out in soft water. Values are reported as micrograms of metal and toxicity as $LT_{50}$ . In solutions containing copper and zinc, fish died twice as fast as would occur if the two metals were simply additive in their lethal action.	Sprague (1964)
CHEMIC	Copper sulfate	Salmo salar	BSA	-	(0)	<u>acdef</u>	The EC <sub>50</sub> or the effective concentration that elicited as avoidance reaction in the fish was 0.052 x the ILL (incipient lethal level), or 0.052 x 44 $\mu$ g/L, or 2.28 $\mu$ g/L.	Sprague (1965)
CHEMICALS AND MIXTURES OF CHEMIC	Copper sulfate	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	(S) 0.025 (T4A) (H) 1.76 (T4A) (S) 0.66 (T4A) (S) 0.036 (T4A) (S) 0.036 (T4A)	c d e f	(S) Soft water (H) Hard water Values are expressed as mg/l of metal.	Pickering and Henderson (1965)
ES OF CHEMIC	Copper sulfate	Carp Tench Ephemeropterae Iarvae Trichopterae Iarvae	FR	France	0.1 (75% K6) 0.2 (75% K6) 0.2 (100% K)	-	Field studies conducted. Two streams were studied; one was used for testing, the other for control. Trichopterae were not affected, i.e., they were active even at concentra- tions of 0.30 ppm.	Vivier and Nisbet (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Copper sulfate	Helix pomatia	BSA	-	0.01-0.1 (O)	C	This paper was concerned with the effect of the chemical on mucous secretion in the snail. Snails exposed to the indicated copper sulfate solutions showed severe signs of toxicity. There was an increase in mucous secretion and the animals did not respond to tactile stimuli.	de Calventi (1965)
Copper sulfate (tech)	Bluegill	BSA	-	2.8 (T4A)	а	This is an estimated LC $_{\rm 50}$ value at temperatures from 55 to 75 F	Cope (1965)
Copper sulfate	Blue-green algae Cylindrospermum Anabaena Anacystis Calothrix Nostoc Oscillatoria Plectonema Green algae Ankistrodesmus Chlorella Closterium Oocystis Green algae Scenedesmus Stigeoclonium Zygnema Green flagellate and yellow algae Chlamydomonas Pandorina Tribonema Gomphonema Navicula Nitzchia	L	_	2.0-4.0 (O)	_	CuSO4 was generally toxic or partially toxic to blue- green algae for 28 days at the indicated concentrations. At 2.0 ppm, it was similarly toxic to the green algae, green flagellates, and yellow algae.	Kemp, et al (1966)
Copper sulfate	Salmo gairdneri Lepomis macrochirus	BSA	-	0.150 (T2A) 2.800 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Соре (1966)
Copper sulfate	Lepomis macrochirus	FL	Various lakes, Michigan	13-140 (K)	a d	For controlling bluegill reproduction, copper sulfate crystals were directed toward nests where eggs and fry were the primary target. The estimated copper sulfate concentrations were estimated to be 13-140 ppm. All eggs and fry were dead in some 200 samplings. Fish other than bluegill fry apparently were not killed by this copper sulfate treatment. Treatment throughout the 3-month spawning period was required for significant reduction of the bluegill population.	Beyerle and Williams (1967)

	Copper sulfate (as Cu)	Salmo salar S. trutta S. Salmo gairdnerii	BSCH	-	0.06 (K)	c f	The reported figure is a reported lethal concentrate as found in polluted lakes and streams in Norway. Organic matter apparently has a masking effect that reduces toxicity. 50% of rainbow trout eggs survived to hatch in 0.05 ppm of Cu. Rainbow trout and Atlantic salmon acted similarly to the chemical. Brown trout were slightly more resistant.	Grande (1967)
	Copper sulfate plus zinc sulfate (various ratios)	Salmo gairdnerii	BSA	-	(0)	<u>a e p</u>	Both hard and soft water were used. Median period of sur- vival in hard water was 3 days – 3.5 ppm Zn, and 1.1 ppm Cu; in soft water – 7 days, 0.56 ppm Zn and 0.044 ppm Cu.	Lloyd (1961)
	Cresol	Lepomis macrochirus	BCFA	-	13.6 (T4A) small 10.9 (T4A) med. 10 (T4A) large	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period.	Cairns and Scheier (1955)
	Cresol	Gambusia affinis	BSA	-	24 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Cresol	Lepomis macrochirus	BSA	-	10.0 (T4A)	acdei 	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
	Ortho- cresol	Phoxinus phoxinus	BCFA	_	0.04% (K 13 min)	<u>a</u> c	Tap water used as a diluent. The apparatus used was a 34 mm diameter tube fitted to permit sharp vertical separation of water and test solution. With this system, avoidance data could be obtained. Toxicity is given as average survival time of replicates. Fish avoided concentrations of 0.03 to 0.04%.	Jones (1951)
<u>0</u>	O-cresol	Sewage organisms	BOD	_	940 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
CHEMICA	O-cresol	Channel catfish (fingerlings)	BSA	_	66.8 (K 69 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
A S	O-cresol	Pimephales promelas	BSA	_	13 (T4A)	acdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made	Pickering and Henderson
		Lepomis macrochirus			24 (T4A)		to estimate 100 percent survival.	(1966)
AND MIXTURE		Carassius auratus			23 (T4A)			
URES		Lebistes reticulatus			29 (T4A)			
	p-cresol	Fish	BSA	-	5.1 x 10 <sup>-5</sup> M (K)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Cryolite Crystal violet	Simocephalus serrulatus Daphnia pulex	BSA	_	10.0 (SB) 5.0 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. "Water Chemistry" (Unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
Crystal violet	Microcystis aeruginosa	L	-	100 (K)	<u>a</u> , etc	The chemical was tested on a 5-day algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Cumene hydroperoxide	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T=toxic, NT=nontoxic, PT= partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - T (7) So - NT Cv - PT (7) Gp - PT (7) Np - T (7)	Palmer and Maloney (1955)
Cupric ammonium chloride	Daphnia magna	BSA	-	0.039 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
Cupric chloride	Daphnia magna	BSA	-	0.08 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoreti- cally infinite) exposure.	Anderson (1944)
Cupric chloride	Daphnia magna	BSA	-	0.027 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
Cupric citrate	Mytilus edulis	BSA	_	0.55 (O)		When the mussels were placed in the test solution for one day, and then in fresh sea water, they died in 2, 3, and 4 days.	Clarke (1947)
Cupric oxide	Gambusia affinis	BSA	-	56,000 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Cupric sulfate	Daphnia magna	BSA	-	0.1 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)

	Cyanide	Mayorella palestinensis (soil amoeba)	BSA	-	(O)	<u>a c</u>	<ul> <li>The experiments were carried out in Warburg manometers at 27 C for 4 hr at a pH of 8.0.</li> <li>Cyanide in concentrations up to 5 x 10<sup>-3</sup> M were shown to have lethal effects on the organism.</li> <li>Results were compared with controls and expressed in percent of respiration.</li> <li>Compared with normal respiration, nonlethal concentrations of cyanide increased the respiration of the organism in glucose-containing solutions.</li> <li>It was concluded that the respiration of the organism depends on at least three enzyme systems, which may be distinguished by their behavior toward cyanide.</li> </ul>	Reich (1955)
А	Cyanide	Lepomis auritus L. macrochirus Micropterus salmoides Pomoxis annularis	BSA & CF	-	0.06 (T1SA) 0.01-0.06 (T<1SA) 0.05-0.06 (T<1CFA) 0.06 (T<11SA) 0.05-0.07 (T<1SA) 0.02-0.04 (T<1CFA)	<u>a</u>	Additional data for less than 24 hr are given and also for the disappearance and breakdown of cyanide in anaerobic soil systems.	Renn (1955)
г-51 CH	Cyanide	Brown trout Small mouth bass	BSA BCF BCF	_	0.31-0.96 (O) 0.32-1.06 (O) 0.175-1.98 (O)	<u>a</u> c d <u>e</u>	The pH of the water varied from 7.5-8.28 in the test solu- tions. Dissolved oxygen was controlled by aeration. In the report, time of death is plotted against cyanide concentra- tion. In a continuous flow apparatus, a range of concentrations from 0.32 to 1.06 ppm killed in 17-48 minutes and 4.2 to 15.2 minutes, respectively. In a static test, 0.31 to 0.96 ppm killed in 33-230 and 6.0-18.7 minutes, respectively. These data are for brown trout. For small mouth bass, in a continuous flow apparatus, concentrations of 1.98 ppm down to 0.175 ppr killed in 6-10 and 213-477 minutes respectively. The effect of dissolved oxygen is discussed.	,
IEMICA	Cyanide	Lepomis macrochirus	BSA	_	0.18 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
LS AND MIXTU	Cyanide	Physa heterostropha Lepomis macrochirus	BSA	-	0.432 (T4A) 0.18 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
CHEMICALS AND MIXTURES OF CHEMICALS	Cyanide	Lebistes reticulatus	BSA	-	(O)	acfno	A series of equations was devised to describe the toxicity of a system containing two toxicants — zinc - zinc and cyanide. Concentrations of cyanide, 0.42 ppm, 0.28 ppm, and 0.26 ppm, killed 50 percent of the animals in 20, 30, and 43 hours, respectively. Toxicity of the two-component system was then determined using varying ratios of the two components.	Chen and Selleck (1968)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURE	Cyanide	Fish (unidentified)	FR	Dunreith, Indiana	0.05-0.1 (K)	_	Tests for cyanide pollution were made following a train- car collision. Five tank cars carrying acetone cyanohydrin, vinyl chloride, ethylene oxide, and methyl methacrylate were involved.	Moore and Kin (1969)
TURES OF	Cγanide (a)- chromium (b)- naphthenic acids ( mixture	Lepomis macrochirus c)	BSA	_	(a) 0.026 (T4A) (b) 0.019 (T4A) (c) 4.74 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
CHEMICALS	Cyanide (a)- zinc (b)- mixture	Lepomis macrochirus	BSA	-	(a) 0.26 (T4A) (b) 3.90 (T4A)	a c d e	Comment same as above.	Cairns and Scheier (1968)
CALS	Cychohexane	Gambusia affinis	BSA	-	15,500 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Cyclohexane	Pimephales promelas Lepomis macrochirus Carassius	BSA	-	30 (T4A) 31 (T4A) 33 (T4A)	<u>a</u> c d e f	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
A-S		auratus Lebistes reticulatus			48 (T4A)			
2	1, cyano-1,3- butadiene	Lagodon rhomboides	BSA	-	71.5 (T1A)	а	Aerated seawater was used.	Daugherty and Garrett (1951)
	1, cyano-1,3- butadiene	Lagodon rhomboides	BSA	-	71.5 (T1A)	-	Experiments were conducted in aerated salt water.	Garrett (1957)
	Cymeme thiocyanate	Green sunfish	BSA	-	(0)	-	Fish were moderately repelled at concentrations of 20 mg/l but the response to 10 mg/l was indifferent. The chemical has apparent high toxicity.	Summerfelt and Lewis (1967)
	2,4-diamino- phenol dihydro- chloride	Microcystis aeruginosa	L	-	100 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
	2,4-diamino- phenol hydro- chloride	Daphnia magna	BSA	_	80 (K2)	a	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical sub- stances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
	Diamylamine	Semotilus atromaculatus	BSA	_	5 to 20 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional date are presented.	Gillette, et al (1952)

2',5'-dibromo- 3-nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K2) 10.0 (K 3 hr) 10.0 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the sali- cylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
3,5-dinitro- 2',3'-benz- oxylidide	Salmo gairdnerii Carassius auratus	BSA	-	(O) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm the chemical was toxic to 1 out of 10 trout in 48 hr. At 10 ppm the chemical was not toxic to goldfish.	Walker, et al (1966)
4',5-dibromo- 3-nitrosalicy!- anilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
Di-sec- butylamine	Semotilus atromaculatus	BSA		15 to 40 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Di-n- butylamine	Semotilus atromaculatus	BSA	-	20 to 60 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
1,3-dibutyl- thiourea	Semotilus atromaculatus	BSA	-	30 to 100 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Orthodichloro- benzene	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	-	13 (NG) 13 (NG) 13 (NG) 13 (NG) 13 (NG)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were expressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG=no growth, but the organisms were viable.	Ukeles (1962)
2,6-dichloro- benzine acid (tech)	Rainbow trout Bluegill	BSA	-	140 (T4A) 120 (T4A)	а	This is an estimated LC $_{\rm 50}$ value at temperatures from 55 to 75 F.	Соре (1965)
2,4-dichloro- benzyl- nicotinium chloride	Microcystis aeruginosa	L	_	5.0 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
n 1,2-dichloro- O ethane	Lagodon rhomboides	BSA	_	150-175 (O)	-	Experiments were conducted in aerated salt water. Toxicity range given as the concentrations which produced $\leq$ 1/2 deaths and $\geq$ 1/2 deaths.	Garrett (1957)
3,6-dichloro- 2,5-dimethoxy- benzoquinone	Microcystis aeruginosa	L	_	75 (K)	<u>a,</u> etc	The chemical was tested on a 5-day-old algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
1,1-dichloro- ethane	Lagodon rhomboides	BSA	-	250-275 (O)	_	Experiments were conducted in aerated salt water. Toxicity range given as the concentrations which produced $<1/2$ deaths and $>1/2$ deaths.	Garrett (1957)
1,4-dichloro- 2-nitro- benzene	Green sunfish	BSA	-	6.5 (T1A) 4.5 (T2A)	a e p	The main purpose of this experiment was to determine the repellent characteristics of certain chemicals. Experiments were conducted in a wooden trough. The toxic action of this chemical appeared to involve suffocation.	Summerfelt and Lewis (1967)
4,4-dichloro- alpha- methyl- benzhydrol	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl - PT (3) Ma - NT So - NT Cv - NT Gp - PT (14) Np - NT	Palmer and Maloney (1955)
2,3-dichloro- naphtho- quinone	Fish: Pomoxis nigromaculatus Notropis antherinoides Hyborhynchus notatus Ambloplites rupestris Huro salmoides Water Plants: Ceratophyllum Myrophyllum Elodea Invertebrates: Snails Daphnia Rotifers	BSA		(O)	<u>e</u>	Aerated spring water was used as the test medium. No effect was observed on fish after 2 days of exposure, even with excess solid dispersed in water. No effect was observed on higher aquatic plants and green algae. At concentrations in excess of saturation level (100 mg/l), no toxic effect was observed. At algicidal concentrations, no toxic effect was noted on any of the species studied.	Fitzgerald, et al (1952)

	2,3-dichloro- napthoqui- none	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - T So - NT Cv - PT (7) Gp - T (7), PT (14) Np - T	Palmer and Maloney (1955)
	2,3-dichloro- naphtho- quinone	Pimephales promelas	BSA	-	0.15 (T4A)	a c d e f	Toxicity to 30 species of algae also presented. 2,3 DNQ was algicidal in the range 0.5 to 2.5 ppm.	Maloney and Palmer (1956)
	2,5-dichloro- 4-nitrophenol	Petromyzon marinus (larvae)	BSA	-	10 (K<1)	<u>a</u>	Additional data are presented.	Piavis (1962)
	2,5-dichloro-	Petromyzon	BSA	-	5 (K 100%)	a	Mortality occurred in approximately 24 hr. This was a	Ball (1966)
	4-nitrophenol (Na salt)	marinus Salmo trutta	BSA	-	17 (K 10%)	a	study on controlling sea lamprey larvae.	(1900)
	2,5-dichloro- 4-nitrophenol	Petromyzon marinus	BSA	_	3 (K 100%)	a	Comment same as above.	Ball (1966)
i	(free phenol)	Salmo gairdnerii	BSA	_	13 (K 10%)	a		
		S. trutta	BSA		7 (K 10%)	a		
CHEMICALS	3',4'-dichloro- 3-nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative posi- tion(s) in the molecule.	Walker, et al (1966)
S AND MIXT	Dichloro- phenoxy- butyric acid	Pteronarcys sp (nymphs)	BSA	_	15.0 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES OF CHEMICALS	Di (p-chloro- phenyl) methyl carbinol	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - NT So - T (3) Cv - T (3) Gp - T (3) Np - T (3)	Palmer and Maloney (1955)
SALS	Diethanol- amìne	Gambusia affinis	BSA		1,550 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Diethanol- amine	Sewage microorganisms	BOD	-	(O)	-	The chemical was studied as to how low levels (ppm) may affect the BOD in domestic sewage. This compound was not toxic to sewage organisms, but responded readily to acclimated seed and contributed to the biochemical oxy- gen demand.	Oberton and Stack (1957)
A-56	Diethylamine	Semotilus atromaculatus	BSA	-	70 to 100 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentra- tion in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952).
	Diethylamino- hydrochloride	Semotilus atromaculatus	BSA	-	4,000 to 6,000 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
	2',5'-diethyl- 3,5-dinitro- benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative posi- tion(s) in the molecule. At 10 ppm the chemical was toxic to 1 out of 10 goldfish in 48 hours.	Walker, et al (1966)
	Diethylene glycol	Gambusia affinis	BSA	-	32,000 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

Diethyl- ethanol- amine	Semotilus atromaculatus	BSA	-	80 to 120 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Diethyl nitrosoamine	Semotilus atromaculatus	BSA	-	900-1,100 (CR)	ae	Comment same as above.	Gillette, et al (1952)
1,3-diethyl- thiourea	Semotilus atromaculatus	BSA	-	100 to 300 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Diglycolic acid	Lepomís macrochirus	BSA	-	105 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerable limit for a selected group of herbicides.	Hughes and Davis (1967)
m-dihydroxy- benzene	Sewage organisms	BOD	-	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Di-isobutyl- amine	Semotilus atromaculatus	BSA	-	20 to 40 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Di-isopropyl- amine	Semotilus atromaculatus	BSA	-	40 to 60 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Dimethyl- amine	Semotilus atromaculatus	BSA	_	30 to 50 (CR)	a e	Comment same as above.	Gillette, et al (1952)
Dimethylamino- benzaldehyde	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI = NT Ma = NT So = NT Cv = NT Gp = NT Np = NT	Palmer and Maloney (1955)
0,0-dimethyl dithiophos- phate (47.7 per- cent)	Lymnaeid snails	BSA	_	(0)	_	Each test container, 500-ml beaker, was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
4,5-dimentyl- 2-mercapto- thiazole	Daphnia magna	BSA	-	56 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
2',3'-dimethyl- 3-nitrosalicyl- anılide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	3.0 (LD <sub>100</sub> ) 5.0 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)
2',4'-dimethyl- 3-nitrosalicyl- anilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	3.0 (LD <sub>100</sub> ) 7.0 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	Comment same as above.	Starkey and Howell (1966)
2',5'-dimethyl- 3-nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	_	1.0 (LD <sub>100</sub> ) 0.7 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	Comment same as above.	Starkey and Howell (1966)
2',6'-dimethyl- 3-nitrosalicyl- anilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	_	>10.0 (LD <sub>100</sub> ) >10.0 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	Comment same as above.	Starkey and Howell (1966)
Dimethyl sulphoxide	Carassius auratus	BSA	_	(0)	a <u>f</u>	At 32 ppt DMSO, five goldfish survived for 10 days without exhibiting signs of respiratory stress or symptoms of toxic reaction. In a similar concentration of acetone the median period of survival was about 90 minutes.	Ball (1966)
Dimethγl sulfoxide	Hemigrammus erythrozonus Paracheinodon innesi Xiphophorus maculatus Pescilia Iatipinna Poecilia reticulata Brachydanio rerio Corydoras paleatus	BSA	_	(O)	<u>a</u> ce	According to the authors, the LD50 concentration in 0-5 days was found to be 1.9% for <i>P. innesi</i> , <i>H. erythrozonus</i> , <i>P. reticulata</i> , <i>P. latipinna</i> , and <i>X. maculatus</i> . <i>B. rerio</i> and <i>C. poleatus</i> tolerated higher concentrations of DMSO for longer periods of time.	Rabinowitz and Myerson (1966)

	Dîmethyl sulfoxide	Salmo gairdneri Salvelinus fontinales S. namaycush Cyprinus carpio Ictalurus melas I. punctatus Lepomis cyanellus L. macrochirus Perca flavescens	BSA	_	53,000 (T1A) 32,300 (T3A) 54,500 (T1A) 36,500 (T3A) 47,800 (T1A) 37,300 (T3A) 44,000 (T1A) 41,700 (T3A) 42,500 (T1A) 36,500 (T3A) 39,000 (T1A) 32,500 (T3A) 65,000 (T1A) 33,500 (T2A) 65,000 (T1A) 33,500 (T2A) 65,000 (T1A) 37,000 (T2A)	<u>a</u> i	Water quality had little effect on toxicity of DMSO but increased temperature increased the toxicity to rainbow trout.	Willford (1967)
	Dimethyl sulfoxide	Oncorhynchus tshawytscha O. nerka O. kisutch Salmo gairdneri	BSA	-	12 (L)	а	LD <sub>50</sub> values were reported in g DMSO/kg body wt. Fish usually died within 24 hr after intraperitoneal injection.	Benville, et al (1968)
	l ,3-dimethyl- urea	Semotilus atromaculatus	BSA	_	7,000 to 15,000 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
-	8,5-dinitro- benzanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	This paper deals with the relations between chemical structures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 10 ppm, this chemical was not toxic to trout or goldfish.	Walker, et al (1966)
2	n-dinitro- benzene (tech)	<i>Lymnaeid</i> snails	BSA	-	(O)		Each test container (50-ml beaker) was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
<b>^</b>	n-dinitro- benzene	Microcystis aeruginosa	L	-	50 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)

CHEMICALS AND MIXTURES OF CHEMICALS

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
3,5-dinitro-2',3'- benzoxylidide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the sali- cylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 10.0 ppm, the chemical was toxic to 1 out of 10 trout in 48 hours. At 10 ppm the chemical was not toxic to goldfish.	Walker, et al (1966)
3,5-dinitro-o- benzotoluidide	Salmo gairdnerii Carassius auratus	BSA	-	10.1 (К2) (О)	<u>a</u>	Comment same as above except at 10.0 ppm, the chemical was toxic to 8 out of 10 goldfish at 48 hours.	Walker, et al (1966)
Dinitro-o-sec- butylphenol (tech)	Lymnaeid snails	BSA	-	(O)	-	Comment same as above except 100% mortality occurred at 1:200,000 and greater.	Batte, et al (1951)
Dinitro-o-sec- butylphenol	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cy) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - NT Ma - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
2,6-dinitro-4- chlorophenol (tech)	Lymnaeid snails	BSA	-	(0)	_	Each test container (500-ml beaker) was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
Dinitrocresol (tech)	Pteronarcys californica (naiads)	BSA	-	0.00032 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
3,5-dinitro-o- cresol (tech)	Lymnaeid snails	BSA	-	(0)	_	Each test container (500-ml beaker) was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
4,6-dinitro-o- cresol acetate (tech)	L <i>ymnaeid</i> snails	BSA	-	(0)	_	Comment same as above.	Batte, et al (1951)

4,6-dinitro-o- cresol methyl ether (tech)	Lymnaeid snails	BSA	-	(O)	-	Comment same as above.	Batte, et al (1951)
Dinitro-o-cyclo- hexylphenol (38 percent)	Lymnaeid snails	BSA	-	(0)	-	Comment same as above except 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)
Dinitro-o-cyclo- hexylphenol, d cyclohexylamir salt (tech)		BSA		(0)	-	Comment same as above except 100% mortality occurred in concentrations of 1:200,000 and greater.	Batte, et al (1951)
Dinitro-o-cyclo- hexylphenol	Lymnaeid snails	BSA	-	1.0 (K1)	-	Each test container (500-ml beaker) was filled with ditch water.	Batte, et al (1951)
Dinitro-o-cyclo- hexylphenol, dicyclohexyl- amine salt (20 percent)	Lymnaeid snails	BSA	-	(O)	_	Comment same as above except 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)
2,4-dinitro- phenol	Sewage organisms	BOD	_	100 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
2,4-dinitro- phenol (tech)	Lymnaeid snails	BSA		(0)	-	Each test container (500-ml beaker) was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
2,4-dinitro- phenolhydrazir (tech)	Lymnaeid ne snails	BSA	-	(0)	-	Comment same as above.	Batte, et al (1951)
2,4-dinitro- phenol, sodium salt { (tech)	<i>Lymnaeid</i> snails	BSA	-	(O)	-	Comment same as above.	Batte, et al (1951)
2,4-dinitro- phenyl- hydrazine	Microcystis aeruginosa	L	_	100 (K)	<u>a</u> , etc.	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
2,4-dinitro- phenyl- hydrazine	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI = NT Ma = NT So = NT PT (7) Cv = NT Gp = NT Np = NT	Palmer and Maloney (1955)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES OF CHEMICA	2',3-dinitro-m- salıcylanılıde	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K2) 10.0 (K 3 hr) (O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 10.0 ppm, the chemical was not toxic to goldfish.	Walker, et al (1966)
2	2',3-dinitro-p- salicylotoluidide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K2) 10.0 (K 3 hr) 10.0 (K 2)	<u>a</u>	Comment same as above except data cited.	Walker, et al (1966)
	3,5-dinitro-o- salicylotoluidide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (K 3 hr) (O)	<u>a</u>	Comment same as above except that at 10.0 ppm, the chem- ical was toxic to 9 out of 10 goldfish at 48 hr.	Walker, et al (1966)
	2,4-dinitro- thymol (tech)	Ly <i>mna</i> eid snails	BSA	-	(O)	_	Each test container (500-ml beaker) was filled with ditch water. 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)
	2,4-dinitro- toluene (tech)	L <i>ymnaeid</i> snails	BSA	_	(O)	_	Comment same as above except less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
	Di-n-propylamine	Semotilus atromaculatus	BSA	_	20 to 60 (CR)	<u>a</u> e 	Test water was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
	Disodium copper salt of ethylene diamine-tetra acetic acid	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl $-$ NT Ma $-$ PT (14) So $-$ NT CV $-$ NT Gp $-$ NT NP $-$ NT	Palmer and Maloney (1955)

Disodium ethylene bisdithio- carbamate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Comment same as above except that: CI - NT Ma - PT (14) So - NT Cv - T (3) Gp - T (3) Np - T (3)	Palmer and Maloney (1955)
Disodium octoborate tetrahydrate	Salmo gairdnerii	BSA	-	4200 (T1A) 2750 (T2A)	<u>a</u> e 	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
Dodecylaceta- mido-dimethyl benzyl ammonium chloride	Cyclindrospermum licheniforme (Cl) Gleocapsa sp (GP) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI $-$ PT (7) G $-$ T (3), PT (14) So $-$ T Cv $-$ T Gp $-$ T Np $-$ T	Palmer and Maloney (1955)
Ethanol	Lesbistes reticulatus Carassius auratus	BSA	_	(O)	a c	The uptake of ethanol from buffered solution by guppies has been studied. There was an apparent increase in the rate of absorption with increasing pH. Experiments with goldfish failed to show an increase in absorption rate as the pH was increased.	Hayton and Hall (1968)
Ethyl alcohol	Carassius carassius	BSA	_	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tempera- ture, effect of dissolved oxygen, the efficiency of the gold- fish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 16 cc per liter, fish survived 98 minutes.	Powers (1918)
Ethyl alcohol	Daphnia magna	BSA	-	18,400 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna.</i> Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure	
Ethyl alcohol	Pygosteus pungitius	BCF	_	(O)	<u>a</u>	A concentration of 4 percent ethyl alcohol immediately intox- icated the fish, which recovered when placed in fresh water. A 1 percent solution caused the fish to exhibit an avoidance reaction.	Jones (1949)

-

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Ethyl alcohol	Semotilus atromaculatus	BSA	-	7,000 to 9,000 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hrs. and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Ethyl benzene	Pimephales promelas	BSA	-	40 (T4A)	<u>a</u> cdef	Most fish survived at test concentrations of about one half, or slightly more, of the $TL_m$ value. No attempt was made	Pickering and Henderson
	Lepomís macrochirus			29 (T4A)		to estimate 100 percent survival.	(1966)
	Carassius auratus			73 (T4A)			
	Lebistes reticulatus			78 (T4A)			
Ethyldietha- nolamine	Semotilus atromaculatus	BSA	_	160 to 200 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Ethylene diamine	Semotilus atromaculatus	BSA	_	30 to 60 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Ethylene thiourea	Semotilus atromaculatus	BSA	_	6,000 to 8,000 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
2,ethyl-1,3- hexanediol	Channel catfish (fingerlings)	BSA	-	624 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
1-(2-ethyl- hexyl)-2- undecyl- 1,4,5,6- tetrahydro- pyrimidine	Microcystis aeruginosa	L	_	2.0 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Ethylmercuric chloríde	Artemia	BSA	-	24.0 (0)	ac	All tests were conducted in seawater.	Corner and Sparrow
Chionae	salina Acartia clausi	Acartia clausi		2.0 (O)		Toxicity values reported are relative to that of mercuric chloride expressed as unity.	(1956)
	Elminius modestus			4.4 (O)		Mechanism of action is discussed, as well as synergistic action of two poisons administered simultaneously.	

	2'-ethyl-3-nitro- salicylanilide	Salmo gairdnerii Carassius auratus	BSA	_	(O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. No affect occurred for rainbow trout or goldfish at 0.1 and 1.0 ppm.	Walker, et al (1966)
	O-ethyl-s- pentachloro- phenyl thiocarbamate	Petromyzoń marinus (larvae)	BSA	-	10 (K<1)	<u>a</u>	Additional data are presented.	Piavis (1962)
	Ferric chloride	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compared this work with earlier work, and lists an extensive bibliography. In a concentration of 0.284N, fish survived 29 minutes; in a concentration of 0.0000166N, they survived 1200 minutes.	Powers (1918)
	Ferric chloride	Daphnia magna	BSA	-	130 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Ferric chloride	Daphnia magna	BSA	-	<18 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1944)
CHEN	Ferric chloride	Gambusia affinis	BSA	-	74 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
CHEMICALS AND MIXTURES	Ferric chloride	Biomorpholaria alexandrina Bulinus truncatus	BSA	_	200 (K1) 200 (K1)	а	The degree of tolerance for vector snails of bilharziasis to various chemicals is somewhat dependent upon tempera- ture. The temperature at which (K1) occurred was 26 C.	Gohar and El-Gindy (1961)
ID MIXT	Ferric chloride	Daphnia magna	BSA	-	36 (T1A) 21 (T2A) 15 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
<b>URES O</b>	Ferric sulfate	Gambusia affinis	BSA	-	133 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Ferrocyanide complex Sodium cyanide (482 ppm CN <sup>-</sup> ) and	Pimephales promelas	BSA	_	10 (K ≪48 hr)	a c	Synthetic soft water was used. Toxicity data given as number of test fish surviving after exposure at 24, 48, and 96 hr.	Doudoroff, et al (1956)
Ferrous sulfate (193 ppm Fe <sup>++</sup> )							
Ferrous chloride	Daphnia magna	BSA	-	<38 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
Ferrous disodium versenate	Channel catfish (fingerlings)	BSA	_	>500 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Ferrous oxide	Gambusia affinis	BSA	-	10,000 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Ferrous sulfate	Daphnia magna	BSA	-	< 152 (0)	a c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Ferrous sulfate	Micropterus salmoides Lepomis machrochirus Goldfish	BSA	-	100 (O) 100 (O) 100 (O)	<u>a</u> cfpi	The disposal of cannery wastes frequently involves the use of chemicals for treatment purposes. Ferrous sulphate, alum, and lime are used in chemical coagulation; sodium carbonate for acidity control in biological filters; and sodium nitrate in lagoons for odor control. Lye (sodium hydroxide) peeling of certain fruits and vegetables is not uncommon. These chemicals, in whole or part, are dis- charged in most cases to a stream. The concentrations listed permitted large mouth bass to survive 2.5 to 3.5 days, and goldfish to survive indefinitely.	Sanborn (1945)
Ferrous sulfate	Sewage organisms	BOD	-	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Ferrous sulphate	Biomorpholaria alexandrina Bulinus truncatus	BSA	-	900 (K1) 900 (K1)	а	The degree of tolerance for vector snails of bilharziasis to various chemicals is somewhat dependent upon tempera- ture. The temperature at which (K1) occurred was 27 C.	Gohar and El-Gindy (1961)

Ferrous sulfide	Gambusia affinis	BSA	-	10,000 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Ferrous sulfite	Gambusia affinis	BSA	_	350 (T2A)	<u>a</u> cdeg	Comment same as above.	Wallen, et al (1957)
Fluoride	Salmo gairdnerii	BSA	-	(H) 250 (K21) (H) 150 (90% K21) (H) 100 (NTE 21) (S) 253 (K21) (S) 113 (K21) (S) 75 (NTE 21)	a d	Aerated lake and well water were used as diluents. Toxicity data are given as percentage killed at various concentrations of fluoride in both hard (320 ppm) and soft water (45 ppm). Threshold for 50% mortality was 8.5 ppm F in 504 hr (21 days).	Herbert and Shurben (1964)
Fluoride	Chlorella pyrenoidosa	L	-	(O)	_	Fluoride caused growth inhibition in cultures of <i>Chlorella pyrenoidosa</i> . This antimetabolite had its greatest effect at concentrations greater than 10 <sup>-3</sup> M. No proportionality could be established between the concentrations of fluoride and the percentages of inhibition occurring at these concentrations.	Smith and Woodson (1965)
2'-fluoro-3',5'- dinitrobenz- anìlide	Salmo gairdnerii Carassius auratus	BSA	_	10 (K2) 10 (K2)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the sali- cylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
3'-fluoro-5- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K2) 10.0 (K 3 hr) 10.0 (K2)	<u>a</u>	Comment same as above.	Walker, et al (1966)
3'-fluoro-3- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.5 (K) (O)	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci. 0.9 ppm killed 25%.	Starkey and Howell (1966)
2'-fluoro-3- nitrosalicyl- anilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	1.0 K	See Applegate, et al (1957-1958)	Comment same as above. 3.0 ppm killed 25%.	Starkey and Howell (1966)
4'-fluoro-3- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	_	1.0 (K)	See Applegate, et al (1957-1958)	Comment same as above. 3.0 ppm killed 25%.	Starkey and Howell (1966)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	4-fluoro-5- nitrosalicyl- anilide	Sea Iamprey (Iarva)	BSA	_	3.0 (K)	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)
AND MIXTURES OF CHEMICALS	Fluosilicic acid	Sewage organisms	BOD	-	2.6 (O)	_	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treatment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chem- ical was tested in an unbuffered system.	Sheets (1957)
	Formaldehyde (40% soln)	Pygosteus pungitius	BCF	_	(O)	<u>a</u>	Concentrations of 0.1 to 0.4 percent (v/v) caused the fish to show a negative reaction and appear to be irritated.	Jones (1947)
0,	Formaldehyde	Sewage organisms	BOD	-	740 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depict- ing the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
•	Formaldehyde	Daphnia magna	BSA	_	100 1000 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
5	Formaldehyde	Salmo gairdneri Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	_	168 (T2A) 185 (T2A) 157 (T2A) 167 (T2A) 96 (T2A) 140 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.1.) were also determined.	Willford (1966)
	Formalin	lctalurus punctatus	BSA	-	126 (K2A) 87 (T2A)	<u>a</u> cfi	The experiment was conducted at 77 C.	Clemens and Sneed (1958)
	Formalin (by volume)	Channel catfish (fingerlings)	BSA	-	87 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Formalin	Tadpoles Various fish	FL	111.	25-30 (K)	a c	After preliminary tests in aquaria, nine pond treatments were made in six different ponds ranging in size from 0.03 to 0.5 acre. Formalin treatments caused oxygen depletion, which, in turn, resulted in a fish kill. The ponds were treated with formalin at 25 to 30 ppm. The authors recommend that	Helms (1967)

with formalin at 25 to 30 ppm. The authors recommend that when fish are present, not more than 30 ppm should be used to kill tadpoles in ponds.

A-68

Formalin	Rana catesbeiana R. pipiens Bufo sp Notemigonus crysoleucas Cyprinus carpio Ictalurus melas Largemouth bass Lepomis macrochirus L. cyanellus Tilapia sp	BSA	_	80 (K), 53 (L1) 30 (K), 22 (L1) 50 (K), 45 (L3) 87 (L1), 67 (L2), 62 (L3) 70 (L3) 70+ (L1), 49 (L2), 45 (L3) 100 (L3) 100+ (L2), 80 (L3) 90 (L3)	a c	Data are reported as LD50, although TL <sub>m</sub> or LC50 might have been more appropriate. The (K) represents minimum con- centration for 100 percent kill.	Helms (1967)
Formic acid	Sewage organisms	BOD	-	550 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depict- ing the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Formic acid	Lepomis macrochirus	BSA	-	175 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Furfural	Gambusia affinis	BSA	_	24 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Glutaric acid	Lepomis macrochirus	BSA	-	330 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Heptane	Gambusia affinis	BSA	-	4,924 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Hexamethylene- tetramine	Sewage organisms	BOD	_	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Hydrochloric acid	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.0000313N solution, fish survived 1200 minutes.	Powers (1918)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Hydrochloric acid	Daphnia magna	BSA	_	62 (O)	a c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wates as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Hydrochloric acıd	Semotilus atromaculatus	BSA	_	60 to 80 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentra- tion in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Hydrochloric acid	Lepomis macrochirus	BCFA	_	(O)	a c e f	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. Toxicity was dependent upon pH. At pH 3.90 to 4.05, 10 percent of the fish died after 2 days. At pH 3.65, 50 percent survived after 3 days.	Cairns and Scheier (1955)
Hydrochloric acid	Gambusia affinis	BSA	_	282 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Hydrochloric acid	Lepomis macrochirus	BSA	-	3.5 (pH, T4A)	<u>acdei</u>	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data re- ported are for larger fish, app 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Hydrocyanic acid	Lagodon rhomboides	BSA	-	0.069 (T1A)	а	Aerated sea water was used.	Daugherty and Garrett (1951)
Hydrogen cyanide	Lagodon rhomboides	BSA	-	0.069 (T1A)	-	Experiments were conducted in aerated salt water.	Garrett (1957)
Hydrogen cyanide	Fish	BSA	_	7.7 x 10 <sup>-6</sup> M (K)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given, Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)
HCN	Lepomis macrochirus (juveniles)	BSA	_	0.16 (T3A)	<u>a</u> cdf <u>p</u>	The solutions were prepared with NaCN, but the data given are calculated as free HCN.	Doudoroff, et al (1966)
Hydrogen cyanide	Salmo gairdnerii	BSA	-	0.07 (T2A)	a c d e f o	The concentration killing a half batch of fish in 2 days pro- vides a reasonable estimate of the threshold concentration. The toxicity of cyanide is related to the concentration of molecular hydrogen cyanide, and not of the cyanide ion (CN <sup>-</sup> ). The lower the pH value the greater the proportion of molecular HCN.	Brown (1968)

	H ion	Fish	BSA	_	1.0 × 10 <sup>-5</sup> M (K)	<u>a</u> .c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)
	Hydrogen sulphide	Oncorhyncus tshawytscha Oncorhyncus kisutch Salmo clarkii clarkii	BSA	-	1.0 (К5) 1.2 (К5) 1.0 (К5)	a d e	This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et al (1952)
	Hydrogen sulfide	Bullia (Gastropoda)	BSA	-	(O)	_	No quantitative data are reported. H <sub>2</sub> S was bubbled through sea water. When animals of this species were exposed to the H <sub>2</sub> S solution more than half an hour, they were killed. Animals removed after 15 minutes, then placed in fresh aerated sea water, recovered.	Brown (1964)
	Hydrogen sulfide (undissociated)	Fish	BSA	-	1.9 x 10 <sup>-5</sup> М (К)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)
	Hydrogen sulfide	lctalurus punctatus	FL	Texas	_	a c g	One hundred cat fish were placed in a pen in one lake and in less than 48 hours, all the test fish fry were dead. Tests showed that total hydrogen sulfide to be 0.96 ppm and a pH of less than 6.0. This gave an unionized H <sub>2</sub> S concentra- tion of at least 0.797 ppm, which was lethal to the catfish. Based on the results of extensive tests, it was evident that the production of unionized H <sub>2</sub> S was seasonal, and often very erratic.	Bonn and Follis (1967)
CHEMICALS 4	Hydrogen sulfide	lctalurus punctatus Lepomis macrochirus	BSA	_	(O)	a c	The quantity of total sulfides necessary to produce a $TL_m$ of the test catfish varied from 1.82 to approximately 7.0 ppm, depending upon the pH of the water. Most of the catfish fry died in approximately 10 minutes at the concentration range given above. At a pH of 7.0 the $TL_m$ of unionized hydrogen sulfide was found to be 1.0 ppm for fingerling channel catfish, 1.3 for advanced fingerlings and 1.4 for adult catfish. The finger- lings died in approximately 20 minutes while the $TL_m$ for advanced fingerlings and adults was attained after about 45 minutes. No $TL_m$ was reached for bluegill in the fingerling tests.	Bonn and Follis (1967)
	Hydroquinone	Microcystis aeruginosa	L	_	100 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
ATURES OF CH	Hydroquinone	Daphnia magna	BSA	_	0.287 (K2)	<u>a</u>	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
EMICAL	Hydroquinone diacetate	Microcystis aeruginosa	L	-	100 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 $\times$ 10 <sup>6</sup> to 2 $\times$ 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Hydroquinone monobenzył ether	Daphnia magna	BSA	-	2.5 (K2)	a	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
Hydroquinone monomethyl ether	Daphnia magna	BSA	-	200 (K2)	а	Comment same as above.	Sollman (1949)
Hydroxyl ion	Fish	BSA	_	1.0 × 10 <sup>-5</sup> М (К)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	lshio (1965)
Hydroxyl ion	Moroco steindachnerii Pungtungia	L	_	11.23 to 9.74 (O) 10.62 to 9.16 (O)		The values given are the $ ho H$ range avoided by the fish.	lshio (1965)
	herzi Acheilognathous			10.12 to 9.03 (O)			
	limbata Cyprinus			10.13 to 8.62 (O)			
	carpio Zaccho			10.12 to 8.62 (0)			
	platypus Sarcocheilichthys			9.63 to 8.71 (O)			
	variegratus Lebistes			9.38 to 8.44 (O)			
	reticulatus Carassius			10.38 to 8.24 (O)			
	auratus (wild) Carassius			10.25 to 7.38 (O)			
	auratus Gnathepogon			10.38 to 7.40 (O)			
	gracilis Pimephalus			9.56 to 9.05 (O)			
	promelas Lepomis macrochirus			9.62 to 8.76 (O)			
Hydroxyl- amine- HCl	Microcystis aeruginosa	L		50 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et a 1 (1952)
Hydroxyl- ammonium benzoate	Microcystis aeruginosa	L	-	100 (K)	<u>a, etc</u>	Comment same as above.	Fitzgerald, et a (1952)
Hydroxyl- ammonium chloride	Microcystis aeruginosa	L	_	100 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et a (1952)

Hydroxyl- ammonium phosphate	Microcystis aeruginosa	L	-	100 (K)	a, etc	Comment same as above.	Fitzgerald, et al (1952)
Hydroxyl- ammonium sulfate	Microcystis aeruginosa	L		100 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et al (1952)
2'-hydroxy- phenazine-1- carboxylic acid	Microcystis aeruginosa Anabaena flos-aquae Notemogonous crysoleucas	L	-	0.1 (O) 1.0 (O)	_	Concentrations noted are for complete inhibition of <i>M. aeruginosa</i> and <i>A. flos-aquae</i> . No harmful effects to <i>N. crysoleucas</i> were noted at the concentrations evaluated.	Toohey, et al (1965)
o-hydroxybenzoic acid	Carassius auratus	BSA	-	0.254 (K)	<u>a</u>	Goldfish weighed between 2 and 4 g. Temperature was maintained at 27.0 $\pm$ 0.2 C.	Gersdorff (1943)
p-hydroxybenzoic acid	Carassius auratus	BSA	-	0.0230 (K)	<u>a</u>	Comment same as above.	Gersdorff (1943)
m-hydroxybenzoic acid	Carassius auratus	BSA	_	0.0363 (K)	<u>a</u>	Comment same as above.	Gersdorff (1943)
p-hydroxyphenyl- glycine	Daphnia magna	BSA	_	20 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949) Fitzgerald, et al
8-hydroxy- quinoline	Microcystis aeruginosa	L	-	100 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Imidazoline	Microcystis aeruginosa	L	-	2.0 (K)	a, etc	Comment same as above.	Fitzgerald, et al (1952)
lodoacetic acid	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - T (3) So - T (3) Cv - NT Gp - PT (14) Np - NT	Palmer and Maloney (1955)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
4'-ıodo-3,5- dinıtrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	(O) (O)	a	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative posi- tion(s) in the molecule. Precipitation occurred at 10 ppm. At 10 ppm the chemical was not toxic to trout or goldfish.	Walker, et al (1966)
2'-iodo-3- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	1.0 (К) (О)	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)
2'-iodo-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10.0 (K 3 hr) 1.0 (K2 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative posi- tion(s) in the molecule.	Walker, et al (1966)
3'-iodo-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	_	1.0 (K 3 hr) 1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)
3'-iodo-3- nitrosalicyl- anilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.3 (K) (O)	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)
4'-iodi-nitro- salicylanilide	lctalurus nebulosus	BSA	_	0.005 (K) 0.0025 (SB) at 47 and 71 F	<u>a</u> c g	The chemical was dissolved in dimethyl sulfoxide for test- ting. Non-aerated, turbid and non-turbid test waters at 47 and 71 F were used. Lodging of the fish in sediment increased survival.	Loeb and Starkey (1966)

¥	4'-iodo-3- nitrosalicyl- anilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	_	0.3 (K) (O)	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci. 0.7 ppm killed 25%.	Starkey and Howell (1966)
	4'-iodo-3- nitrosalicyl- anilide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K 3 hr) 1.0 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative posi- tion(s) in the molecule.	Walker, et al (1966)
	4'-iodo-5- nìtrosalicyl- anilide	Salmo gairdnerii	BSA	-	1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)
A-75	4'-iodo-5- nitrosalicyl- anilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	0.5 (K) (O)	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci. 1.0 ppm killed 25%.	Starkey and Howell (1966)
	m-iodophenol	Carassius auratus	BSA	_	51.7 to 155.0 (K 8 hr) 38.8 (O) 10.3 (O)	<u>a</u>	Temperature in test containers was maintained at 27 ± 0.2 C. Goldfish tested weighed between 2 and 4 g. m-iodophenol, 38.8 ppm, killed 75% of the fish in 8 hr; 10.3 ppm killed 33% in 8 hr.	Gersdorff and Smith (1940)
우	o-iodophenol	Carassius auratus	BSA	-	45.8 to 91.6 (K 8 hr) 36.6 (O) 26.2 (O)	<u>a</u>	Comment same as above except that o-iodophenol, 36.6 ppm, killed 83% of the fish in 8 hr; 26.2 ppm killed 8% in 8 hr.	Gersdorff and Smith (1940)
EMICALS AND	p-iodophenol	Carassius auratus	BSA	-	12.5 to 100 (K 8 hr) 11.8 (O) 10.0 (O) 7.5 (O)	<u>a</u>	Comment same as above except that p-iodophenol, 11.8 ppm, killed 92% of the fish in 8 hr; 10.0 ppm killed 77% in 8 hr; and 7.5 ppm killed 46% in 8 hr.	Gersdorff and Smith (1940)
CHEMICALS AND MIXTURES OF CHEMICALS	Iron	Daphnia magna	L	-	100 (K)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide — a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
lso-amyl alcohol	Daphnia magna	BSA	_	881 (O)	a c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
lsoamyl alcohols, mixed primary	Semotilus atromaculatus	BSA	_	400 to 600 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concen- tration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
IsobornyI thiocyano- acetate	Green sunfish Largemouth bass Black bullhead Golden shiner Mosquito fish Tadpoles Crayfish Bluegill Channel catfish Redear sunfish White crappie	FL	111.	(O)	а	<ul> <li>Ponds were treated with concentrations of 0.7, 0.8, and 1.5 ppm of the chemical. The ponds were drained or poisoned after the removal of isobornyl thiocyanoacetate-affected fish were removed. This was done to determine the numbers of each species that had survived.</li> <li>Water temperature in the ponds ranged from 50 to 87 F. Pond sizes ranged from 0.1 to 455 acres.</li> <li>Results were quite similar to the results obtained in bioassay studies.</li> <li>Centrarchids were selectively killed in the presence of ictalurids and cyprinids.</li> </ul>	Lewis (1968)
lsobornyl thiocyano- acetate		BSA	_		<u>a</u>	Twenty liter-glass aquaria were employed for the experi- ments. Temperature was maintained at 20 to 23 C. Results are recorded as 24-hr lethal minimum dose of the chemical.	Lewis (1968)
	Green			0.6 (O)		24-hr lethal minimum dose at 20 to 23 C.	
	sunfish Rainbow			<0.7 (0)		24-hr lethal minimum dose at 11 C.	
	trout Golden		1.5 (O)		24-hr lethal minimum dose at 20 to 23 C.		
	shiner Channel			1.5 (O)		24-hr lethal minimum dose at 20 to 23 C.	
	catfish Black			>1.5 (0)		24-hr lethal minimum dose at 20 to 23 C.	
	bullhead Bluegill			0.4 (O)		24-hr lethal minimum dose at 20 to 23 C.	

lsobutyl alcohol	Carassius Carassius	BSA	_	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 5.85 cc per liter, fish survived 61 minutes.	Powers (1918)
Isoprene	Pimephales promelas Lepomis macrochirus	BSA	_	75 (T4A) 39 (T4A)	<u>a</u> cdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
	Carassius auratus			180 (T4A)			
	Lebistes reticulatus			140 (T4A)			
p-isopropoxy diphenyl	Daphnia magna	BSA	-	5.7 (K2)	a	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
p-isopropoxy diphenylamine	Daphnia magna	BSA	-	5.7 (K2)	а	Comment same as above.	Sollman (1949)
Isopropyl ałcohol	Semotilus atromaculatus	BSA	_	900 to 1,100 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
1-isopropyl-2- (8,11-hepta- decadienyl)- 4,4-dimethyl- 2-imidazoline	Microcystis aeruginosa	L	-	2.0 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
1-isopropyl- 2-(S-hepta- decenyl)- 4,4-dimethyl- 2-imidazoline	Microcystis aeruginosa	L	-	1.0 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et al (1952)
1-isopropyl- 2-nonyl-4, 4-dimethyl- 2-imidazoline	Microcystis aeruginosa	L	-	2.0 (K)	<u>a</u>	Comment same as above.	Fitzgerald, et al (1952)
1-isopropyl- 2-undecyl- 4,4-dimethyl- 2-imidazoline	Microcystis aeruginosa	L	-	2.0 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et al (1952)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Lactic acid	Daphnia magna	BSA	-	243 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Lactonitrile	Lagadon rhomboides	BSA	-	0.215 (T1A)	а	Aerated seawater was used.	Daugherty and Garrett (1951)
Lactonitrile	Lagadon rhomeboides	BSA	-	0.215 (T1A)	-	Experiments were conducted in aerated salt water.	Garrett (1957)
Lactonitrile	Lepomis auritus Lepomis macrochirus Pomoxis annularis	BSA & CF	-	0.06-0.1 (100% KCF) 0.03-0.1 (100% KS) 0.055-0.07 (100% KF) 0.075 (100% KS) 0.065-0.07 (100% KS)	<u>a</u>	Additional data are presented for less than 24 hr.	Renn (1955)
Lactonitrile	Pimephales promelas Lepomis macrochirus Lebistes reticulatus	BSA	_	(100% K3) (S) 0.90 (T4A) (S) 0.90 (T4A) (S) 1.37 (T4A)	c d e f	(H) Value for hard water. (S) Value for soft water.	Henderson, et a (1960)
	Pimephalus promelas			(H) 0.90 (T4A)		The chemical did not change the flavor of the cooked bluegill.	
Laurylisoquino- linium bromide	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T (3) PT (7) Ma - PT (14) So - T (3) Cv - PT (7) Gp - PT (7) Np - PT (7)	Palmer and Maloney (1955)

	Lead	Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	BSA	_	1.0 (К) 100.0 (К) 10.0 (К)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide – a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)
	Lead	Gasterosteus aculeatus	BSA	_	0.1 (O)	<u>ace</u>	This is a discussion of a bioassay method using stickleback fish and spectrophotometric determinations of the chem- icals evaluated. The number listed is said to be the "toxic limit" for the fish.	Hawksley (1967)
	Lead acetate	Pimephales promelas Lepomis macrochirus	BSA	_	(S) 7.48 (T4A)	c d e f	(S) Soft water. Values are expressed as mg/l of lead.	Pickering and Henderson (1965)
	Lead chloride	Daphnia magna	BSA	-	1.25 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
j •	Lead chloride	Pimephales promelas	BSA	-	(H) >75 (T4A) (S) 2.4 (T4A)	acdf	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
,	Lead chloride	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	(S) 5.58 (T4A) (H) 482.0 (T4A) (S) 23.8 (T4A) (H) 442.0 (T4A) (S) 31.5 (T4A) (S) 20.6 (T4A)	c d e f	(S) Soft water. (H) Hard water. Values are expressed as mg/l of metal.	Pickering and Henderson (1965)
CHEMICALS	Lead nitrate	Gasterosteus aculeatus	BSA	_	0.3 (TL4-3/4A)	ac	Death of the fish resulted from an interaction between the metallic ion and the mucus secreted by the gills. Coagu- lated mucus formed on the gill membranes and impaired respiration to such a degree that the fish asphyxiated. The addition of 50 mg/l of calcium chloride to the tank protected against the toxic effect of this metal salt.	Jones (1938)
AND MIXTURES	Lead nitrate	Gasterosteus aculeatus Phoxinus phoxinus	BSA	-	(O)	C e	Tap water was used to make up the solutions. The animals were attracted to a solution 0.04N - a positive reaction, they tended to swim into it. They tended to show avoidance reactions at concentrations of 0.004N down to 0.00002N. The minnow detected and avoided a 0.000004N solution. <i>P. phoxinus</i> minnows were much more sensitive to this chemical than <i>G. aculeatus</i> .	Jones (1948)
OF CHE	Lead nitrate	Gambusia affinis	BSA	-	240 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
CHEMICALS	Lead nitrate	Lebistes reticulatus	BSCH	-	2.0 (27% K90)	<u>a</u> cde	Sublethal effects found were retarded growth, increased mortality, and delayed sexual maturity.	Crandall and Goodnight (1962)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND	Lead nitrate	Tubificid worms	BSA	-	49.0 (T1A) 27.5 (T1A)	a c 	Knop's solution was used. TL <sub>m</sub> levels for various pHs were determined for the tubificids and were found to be 5.8 to 9.7. Lead nitrate was more toxic at pH extremes of 6.5 and 8.5 than at 7.5.	Whitley (1968)
MIXTURES (	Lead oxide	Gambusia affinis	BSA	_	56,000 (T2A)	a c d e g	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
OF CHEMICALS	Lead salts	Salmo gairdnerii	BSA	_	(O)	<u>a e</u> 	<ul> <li>This is a study of the effect of varying dissolved oxygen concentrations on the toxicity of selected chemicals.</li> <li>The toxicity of heavy metals, ammonia, and monohydric phenols increased as the dissolved oxygen in water was reduced. The most obvious reaction of fish to lowered oxygen content is to increase the volume of water passed over the gills, and this may increase the amount of poison reaching the surface of the gill epithelium.</li> <li>The concentration of the chemical in the water was not specified.</li> </ul>	Lloyd (1961)
	Lithium chloride	Carassius carassius	BSA	_	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.166N solution, fish survived 234 minutes.	Powers (1918)
)	Lithium chloride	Daphnia magna	BSA	-	<7.2 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	D-lysergic acid	Notemigonis crysoleucas Cyprinus carpio Carassius auratus Rhinichthys atratulus Semotilus atromaculatus Notropis cornutus Lepomis gibbosus Lebistes reticulatus Perca flavescens Catostomus commersoni Ameiurus nebulosus	BSA	_	(O)	а	Lysergic acid and 45 of its derivatives were tested on a wide variety of aquatic animals. Various concentrations of the chemicals were used, from 0.5 to as high as 12.0 ppm. In nearly all cases, the chemical caused involuntary surfacing of the fish with no mortality at the above concentrations.	Loeb, et al (1965)

		Salmo trutta Cottus cognatus Boleosoma nigrum Rana pipiens						
	Magnesium chloride	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.313N solution, fish survived 88 minutes.	Powers (1918)
	Magnesium chloride	Daphnia magna	BSA	-	740 (O)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	Magnesium chłoride	Gambusia affinis	BSA	_	17,750 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
A-81	Magnesium chloride	Daphnia magna	BSA	_	3,391 (T1A) 3,489 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalu- ations were made in various types of water.	Dowden and Bennett (1965)
81	Magnesium nitrate	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemi- cals, possible mechanism of action of some, the effect of temperature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.229N solution, fish survived 107 minutes.	Powers (1918)
CHE	Magnesium nitrate	Gasterosteus aculeatus	BSA	-	300 (K10)	-	Solutions were made up in tap water 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
CHEMICALS A	Magnesium nitrate	Biomorpholaria 'a. alexandrina	BSA	-	(0)	а	The degree of tolerance for vector snails of biharziasis chemicals is somewhat dependent upon temperature. <i>B. a. alexandrina</i> tolerated a 24-hour exposure to 6200 ppm at 20 C.	Gohar and El-Gindy (1961)
AND MIX	Magnesium sulfate	Gambusia affinis	BSA	-	15,500 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
MIXTURES OF	Magnesium sulfate	Biomorpholaria a. alexandrina Bulinus truncatus	BSA	-	(O) 4000 (K1A)	а	The degree of tolerance for vector snails of biharziasis chemicals is somewhat dependent upon temperature. The temperature at which (K1A) occurred was 26 C for <i>Bulinus</i> . The tolerance for <i>Biomorpholaria</i> was 6200 ppm.	Gohar and El-Gindy (1961)
- CHEMICALS	Magnesium sulfate	Daphnia magna Lepomis macrochirus	BSA	-	3,803 (T4A) 19,000 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalu- ations were made in various types of water.	Dowden and Bennett (1965)
ຂັ		L <i>ymnaea</i> sp (eggs)			10,530 (T1A)			

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Malachite green	lctalurus punctatus	BSA	_	0.19 (K2) 0.14 (T2A)	<u>a</u> cfi	The experiment was conducted at 77 C.	Clemens and Sneed (1958)
Malachite green	Microcystis aeruginosa	L	_	100 (K)	a, etc	The chemical was tested on a 5-day algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Malachite green (oxalate salt)	Channel catfish (fingerlings)	BSA	_	0.14 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Malachite green	Micropterus salmoides (fry) Lepomis macrochirus (fry)	BSA	-	0.025 (SB3) 0.001 (SB3)	a c d e f p	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
Malachite green	Salmo gairdnerii	BSA	-	0.39 (T2A)	f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
Salmo trutta			0.34 (T2A)			,	
	Salvelinus fontinalis			0.26 (T2A)			
	Salvelinus namaycush			0.40 (T2A)			
	lctalurus punctatus			0.20 (T2A)			
	Lepomis macrochirus			0.11 (T2A)			
Malachite green	Salmo gairdnerii Rasbora heteromorpha	BCFA	_	0.04 (threshold)	<u>a</u> d <u>e</u>	<ul> <li>Aerated hard water was used. Threshold concentrations were examined by 4 methods.</li> <li>1. Long term - survival related to concentration.</li> <li>2. Short term - percentage kill in narrow range of concentrations.</li> <li>3. Comparison of survival times.</li> <li>4. Extrapolation of short-term results by plotting velocity of death against log of concentration.</li> </ul>	Abram (1967)
Malachite	Salmo gairdnerii	BSA	_	(0)	f	This report derives a mathematical equation for determining a threshold concentration for a toxicant. A concentration	
green	gaironern Rasbora heteromorpha			(0)		of 0.048 ppm of the compound will kill 50% of trout in about 18 days. 0.122 ppm was lethal to 50% in two and a half days.	
Maleic anhydride	Gambusia affinis	BSA	-	240 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Maleic hydrazide	Salmo gairdnerii	BSA	-	85 (T1A) 56 (T2A)	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)

	Malonic acid	Lepomis macrochirus	BSA	-	150 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalu- ations were made in various types of water.	Dowden and Bennett (1965)
	Manganese	Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	L	-	10,000 (К) 10,000 (К) 1,000 (К)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide — a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)
	Manganese chloride	Daphnia magna	BSA	-	50 (O)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	Manganese chloride	Limnaea palustris (eggs)	BSA	-	5 x 10 <sup>-5</sup> M (K1)	ac	Toxicity is given in molar concentrations for maximum direct mortality (kill) in 4 hours.	Morrill (1963)
	Manganese disodium versenate	Channel catfish (fingerlings)	BSA	-	>500 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Manganese nitrate	Gasterosteus aculeatus	BSA	_	40 (K10)	_	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
CHEMICALS AN	Mercuric acetate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T (3) Ma - T (3) So - T (3) Cv - T (3) Gp - T (3) Np - T (3)	Palmer and Maloney (1955)
	Mercuric chloride	Gasterosteus aculeatus	BSA	-	0.008 (K10)	_	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
S D F	Mercuric chloride	Balanus balanoides	BSA	-	1.0 (0)	-	The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
CHEMIC	Mercuric chloride	Pygosteus pungitius	BCF	-	(O)	ac	The fish were immersed in solutions of 0.003, 0.002, 0.0003, and 0.00004N mercuric chloride. Survival times in these solutions were respectively, 14, 22, 31, and 100 minutes.	Jones (1947)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Roferonco (Year)
AND	Mercuric chloride	Daphnia magna	BSA	_	<0.006 (0)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
MIXTURES	Mercuric chloride	BOD	L		1.0 (O)	j	"Toxicity is expressed as 80 percent reduction in oxygen utilization.	Ingols (1955)
о Г	chloride	Sewage organisms	BOD	-	(O)	-	There was a slow increase in toxicity of mercury from 0.02 to 0.2 ppm. Beyond this there was a sharp rise in the toxicity until at approximately 2.0 ppm there was com- plete bacteriostasis or an absence of BOD at this concentration.	Ingols (1954)
CHEMICALS	Mercuric chloride	Sewage organisms	BOD	-	0.61 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxy- gen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
A	Mercuric iodide	Artemia salina Acartia clausi Elminius	BSA	-	31.0 (O) 1.7 (O) 2.6 (O)	a c	All tests were conducted in seawater, Toxicity values reported are relative to that of mercuric chloride expressed as unity. Mechanism of action is discussed, as well as synergistic action of two poisons administered simultaneously.	Corner and Sparrow (1956)
A-84	Mercury	modestus Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	BSA	-	0.01 (K) 0.1 (K) 0.1 (K)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide – a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)
	Mercury	Maia squinado	BSA	-	10 (SB 28)	_	Results showed that the highest mercury concentrations oc- curred in the gills and internal organs. Concentrations were minute in the blood and there was none in the urine.	Corner (1959)
	Mercury compounds	Esox leucius	FL	Denmark	(O)	_	Mercury may become a water contaminant from seed dress- ings in agriculture, fungicides in pulp and paper mills, and from the chlorine alkali industry. Pike was chosen as an indicator organism, and many analyses were given for mer- cury content of pike. In water with a mercury content of 0.07 ppb, pike were found with a concentration of 3000 times that concentration. Analyses were reported of pike containing from 60 to 2500 ppb. One value as high as 8000 ppb was reported. There are many organisms capable of accumulating mercury from water.	Johnels, et al (1967)

	Methanol	Sewage organisms	BOD	-	(NTE)	-	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition $(TC_{50})$ of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	2'-methoxy-5'- chloro-3-nitro- salicylanilide	Sea lamprey (larva) <i>Salmo gairdneri</i> (fingerling)	BSA	-	0.7 (LD <sub>100</sub> ) 1.0 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)
	Methyl alcohol	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 25 cc per liter, fish survived 206 minutes.	Powers (1918)
	Methyl alcohol	Daphnia magna	BSA	_	32,000 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was de- fined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Methyl alcohol	Semotilus atromaculatus	BSA	_	8,000 to 17,000 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
CHEN	Methylamine HCl	Microcystis aeruginosa	L	-	100 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 × 10 <sup>6</sup> to 2 × 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
AICALS AND M	p-methylamino- phenol	Daphnia magna	BSA	-	0.5 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
IXTURES OF C	2'-methyl-3'- chloro-3-nitro- salicylanilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	0.7 (LD <sub>100</sub> ) 1.0 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
2'-methyl-4'- chloro-3-nitro- salicylanilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	0.5 (LD <sub>100</sub> ) 0.7 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	This paper deals with the comparative toxicity of halonitro- salicylanilides to sea lamprey and fingerling rainbow trout as a function of substituent loci.	Starkey and Howell (1966)
2'-methyl-5'- chloro-3-nitro- salıcylanilide	Sea lamprey (larva) Salmo gairdneri (fingerling)	BSA	-	0.5 (LD <sub>100</sub> ) 0.9 (LD <sub>25</sub> )	See Applegate, et al (1957-1958)	Comment same as above.	Starkey and Howell (1966)
Methyldodecyl- benzyl trimethyl ammonium chloride	Cylindrospermum licheniforme (Cl) Gleocapsa sp (G) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T (3), PT (7) G - PT (3) So - T (14) Cv - PT (7) Gp - T (14) Np - T (14)	Palmer and Maloney (1955)
Methył dodecyl benzyl trimethyl ammonium chloride plus tridecyl methyl hydroxy ethyl imidazolinium chloride	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Comment same as above except that: CI - NT Ma - NT So - PT (14) Cv - PT (14) Gp - NT Np - NT	Palmer and Maloney (1955)
1,1'-methylenedi- 2-naphthol (bis(2-hydroxy- naphthyl) methane)	Ptychocheilus oregonensis	FR	Idaho	(O)	<u>a</u>	<ul> <li>The creek was treated with 0.75 lb of chemical. Surface temperature remained at 61 F during the 3-hr treatment. The inlet of the stream was treated with 0.05 ppm for 2 hr after the lagoon was treated.</li> <li>Four and one-half hours after the start of the treatment, four northern squawfish were found dead. The next morning numerous dead squawfish were observed on the bottom of the lagoon.</li> <li>No live squawfish were seen and no dead fish of any other species were observed.</li> </ul>	MacPhee and Ruelle (1968)

	1,1'-methylenedi- 2-naphthol [bis(2-hydroxy- naphthyl) methane] Methylene blue — s	Ptychocheilus oregonensis Onchorhynchus tshawytscha Onchorhynchus kisutch Salmo gairdneri see Appendix B	BSA	-	0.006 (K4A) 0.008 (K4A) 0.010 (K4A) 0.015 (K4A)	<u>a</u> e	<ul> <li>Experiments were conducted in vessels containing 10 liters of water.</li> <li>Temperature was held at 65 F.</li> <li>Temperature was held at 60 F.</li> <li>Temperature was held at 55 F.</li> <li>Temperature was held at 50 F.</li> <li>This chemical had no toxic effect upon Chinook salmon, Coho salmon or steelhead trout at the temperature and</li> </ul>	McPhee and Ruelle (1968)
	Methyl mercaptan	Onchorlynchus tshawytscha Oncorhyncus kisutch Salmo clarkii clarkii	BSA	_	0.9 (K5) 1.75 (K5) 1.2 (K5)	<u>a</u> de	concentration indicated for squawfish. This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et al (1952)
	Methyl mercury chloride	Venus japonica Hurmomya mutabilis	F	Japan	(O)	-	Human beings, cats, and waterfowl eating shellfish from Minamata Bay all succumbed to a strange poisoning. At autopsy, clinicopathological changes similar to those induced in mercury poisoning, were found in the cerebellum, and the cerebral cortices. The shellfish were examined chemically and were found to contain as much as 85 mg/kg. The mercury compound was identified and found in the effluent waste from a chemical plant making acetyldehyde. A treatment was found to eliminate the pollutant.	lrukayama (1966)
	Methyl mercury dicyandiamide	Procambarus clarkii (juvenile)	BSA	-	0.083 (T5A)	acdo	The pesticides studied in this report are widely used in rice culture in Louisiana and are toxic to crawfish.	Hendrick and Everett (1965)
CHEMICALS AND	Methyl methacrylate	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	150 (T4A) 250 (T4A) 240 (T4A) 420 (T4A)	_acdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
D MIXTURES OF CHEMICAL	2-methyl- naphtho- quinone	Pomoxis nigromaculatus Notropis atherinoides atherinoides Hyborhynchus notatus Ambloplites rupestris rupestris Huro salmoides	BSA	_	0.3 to 0.6 (K1-2)	<u>e</u>	Aerated spring water was used as the test medium. Effective algicidal concentrations were also toxic to fish.	Fitzgerald, et al (1952)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
5'-methyl-o- salıcylanisidide	Salmo gairdnerii Carassius auratus	BSA		10 (K2) 10 (K2)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)
Methyl vinyl ketone	Sewage microorganisms	BOD	_	1.5 (O)	_	The chemical was studied as to how low levels (ppm) may affect BOD in domestic sewage. The chemical was toxic at the level stated.	Oberton and Stack (1957)
Molybdic anhydride	Pimephales promelas	BSA	-	(H) 370 (T4A) (S) 70 (T4A)	acd f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
Monoamyl- amine	Semotilus atromaculatus	BSA	-	30 to 50 (CR)	<u>a</u> e 	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Mono-n- butylamine	Semotilus atromaculatus	BSA	-	30 to 70 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Monoethyl- ethanolamine	Semotilus atromaculatus	BSA	-	40 to 70 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Mono- isobutylamine	Semotilus atromaculatus	BSA	_	20 to 60 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Mono-iso- propylamine	Semotilus atromaculatus	BSA	_	40 to 80 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Mono- methylamine	Semotilus atromaculatus	BSA	_	10 to 30 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Mono-n- propylamine	Semotilus atromaculatus	BSA	-	40 to 60 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Mono-sec- butylamine	Semotilus atromaculatus	BSA	-	20 to 60 (CR)	<u>a</u> e	Comment same as above.	Gillette, et al (1952)
Naphthenic acid	Lepomis məcrochirus	BSA	_	5.6 (T4A)	ace	Increase in temperature seemed to increase toxicity of this chemical. Low dissolved oxygen reduced toxicity of some chemicals in this study. Toxicity values may be 20% higher in hard versus soft water.	Cairns (1957)

	Naphthenic	Lepomis	BSA	_	(N) 5.6 (T4A)			Coirpo and
	acid	macrochirus Physa heterostropha		-	(N) 5.6 (14A) (L) 2.0 (T4A) (N) 6.6-7.5 (T4A) N (L) 2.0 (T4A) L	<u>a e</u>	Modified Chu No. 14 test medium was used. Toxicity is given both for "normal" O <sub>2</sub> (5-9 ppm), (N), and with "low" O <sub>2</sub> (2 ppm DO), (L). High and low threshold concentration and concentration percent of survival are also presented.	Cairns and Scheier (1958)
	Naphthenic acid	Lepomis macrochirus Physa heterostropha	BSA	-	5.6 (T4A) 2.0 (T4A) 6.6-7.5 (T4A) 2.0 (T4A)	a e	Normal oxygen content in water. Low oxygen content in water. Normal oxygen content in water. Low oxygen content in water.	Cairns (1965)
	Naphthenic acid	Nitzschia linearis Physa heterostropha Lepomis macrochirus	BSA	_	43.1 (T5A) 6.6-7.5 (T4A) 5.6 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
	Naphthenic acids	Lepomis macrochirus Physa heterostropha	BSA	_	5.79 (T4A) 6.60 (T1A)	<u>acdf</u>	This chemical is a mixture of compounds with a general formula of CnH <sub>2</sub> N-O <sub>2</sub> , CnH <sub>2</sub> N-4O <sub>2</sub> , or CnH <sub>2</sub> N-6O <sub>2</sub> , which are widely used in insecticidal formulations. The experiments were conducted in a synthetic dilution water of controlled chemical composition. In hard water, the chemical was somewhat less toxic.	Cairns and Scheier (1962)
	Naphthenic acids	Brachydanio rerio (adults) (eggs) Lepomis macrochirus	BSA		16.3 (T2A) 3.5 (T2A) 5.6 (T2A)	acdef	The test dilutions were made up from distilled water and ACS grade chemicals. Temperature was held at 24 C and the solution was aerated to maintain a dissolved oxygen content of 5-9 ppm.	Cairns, et al (1965)
	Naphthenic acids	Lepomis macrochirus	BSA	-	5.6 (T4A)	acde	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
CHEMI	Naphthenic acid (a) - cyanide (b) - chromium (c) mixture	Lepomis macrochirus	BSA	-	(a) 4.74 (T4A) (b) .026 (T4A) (c) 0.019 (T4A)	a c d e	Comment same as above.	Cairns and Scheier (1968)
CALS A	Naphthalene	Gambusia affinis	BSA	-	165 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	a-naphthol	Microcystis aeruginosa	L	-	100 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture. 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. CHU No. 10 medium was used.	Fitzgerald, et al (1952)
יוד	b-naphthol	Microcystis aeruginosa	L	_	100 (K)	<u>a</u>	Comment same as above.	Fitzgerald, et al (1952)
	1,4-naphtho- quinone	Microcystis aeruginosa	L	_	100 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et al (1952)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
1,4-naphtho- quinone a-naphthylamine	Pomoxis nigromaculatus Notropis atherinoides Hyborhynchus notatus Ambloplites rupestris Huro salmoides	BSA	-	0.3 to 0.6 (K1-2)	e	Aerated spring water was used as the test medium. Effective algicidal concentrations were also toxic to fish.	Fitzgerald, et al (1952)
a-naphthylamine	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI $-$ PT (7) Ma $-$ T So $-$ T (7) Cv $-$ T (7), PT (21) Gp $-$ T (3), PT (7)	Palmer and Maloney (1955)
b-naphtha- quinoline	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Comment same as above except that CI = PT Ma = NT So = PT Cv = PT (7) Gp = T (7), PT (21) Np = T (3), PT (7)	Palmer and Maloney (1955)
Nickel	Rainbow trout	FR	Scotland	25 (T2)	acefim	This work represents an extension of laboratory studies of the toxicity of complex effluents to investigations of rivers.	Herbert, et al (1965)
Nickel	Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	L	-	10 (K) 100 (K) 10 (K)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product constant of the metal sulfide — a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)

	Nickel ammonium sulfate	Sewage organisms	BOD	-	134 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
	Nickel chloride	Daphnia magna	BSA	-	<0.7 (0)	a	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	Nickel chloride	Sewage organisms	BOD	-	38 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
	Nickelous chloride	Pimephales promelas	BSA		(H) 24 (T4A) (S) 4 (T4A)	a c d f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
	Nickel chlorìde	Limnaea palustris (eggs)	BSA	-	8 x 10 <sup>-6</sup> M (K1)	ac	Toxicity is given in molar concentrations for maximum direct mortality (kill) in 4 hours.	Morrill (1963)
	Nickel chloride	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	<ul> <li>(S) 5.18 (T4A)</li> <li>(H) 42.4 (T4A)</li> <li>(S) 5.18 (T4A)</li> <li>(H) 39.6 (T4A)</li> <li>(S) 9.82 (T4A)</li> <li>(S) 4.45 (T4A)</li> </ul>	c d e f	(S) Soft water (H) Hard water Values are expressed as mg/l of metal.	Pickering and Henderson (1965)
CHEM	Nickel- cyanide complex	Lepomis macrochirus (juvenile)	BSA	-	(0)	acdfp_	In solution with a calculated CN content of 100 to 500 ppm, the median resistance time was 143 to 540 min. There was no apparent correlation between median resistance time and concentration.	Doudoroff, et al (1966)
ICALS AND MIXTU	Nickel cyanide complex [sodium cyanide (600 ppm CN <sup>-</sup> ) plus nickelous sulfate (355 ppm Ni)]	Pimephales promelas	BSA	-	0.95 (T4A)	<u>a</u> c <u>d</u>	Synthetic soft water was used. Toxicity data given as number of test fish surviving after exposure at 24, 48, and 96 hr. $TL_m$ values were estimated by straight-line graphical interpolation and given in ppm CN <sup>-</sup> . Additional toxicity data in which total alkalinity was varied, 730 (T-4) with 192 ppm CaCO <sub>3</sub> alkalinity.	Doudoroff, et al (1956)
RES OF CHEN	Nickel- ferrocyanide complex	Pimephales promelas	BSA	-	1.0 ppm CN 0.8 ppm Cu 0.4 ppm Fe (non-toxic after 4 days)	<u>a</u> c	Synthetic soft water was used. Toxicity data given as number of test fish surviving.	Doudoroff, et al (1956)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Nickel nitrate	Gasterosteus aculeatus	BSA	-	0.8 (K10)	_	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
AND MIXTURES OF CHEMICALS	Nickel nitrate	Sewage organisms	BOD	_	64 (0)	_	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
S S	Nickel sulfate	Sewage organisms	BOD	_	16 (O)	_	Comment same as above.	Sheets (1957)
	Nickel sulfate	Salmo gairdneri	BSA	-	160 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
		Salmo trutta			270 (T2A)			
		Salvelinus fontinalis			242 (T2A)			
>		Salvelinus namaycush			75 (T2A)			
2		lctalurus punctatus Lepomis macrochirus			165 (T2A) 495 (T2A)			
	Nitric acid	Daphnia magna	BSA	-	107 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposures.	Anderson (1944)
	Nitric acid	Gambusia affinis	BSA	-	75 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	3-nitro-4 acetoxybenzoic acid	Cylindorspermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema pervulum (Gp) Nitzschie pelee (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - NT Ma - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)

	3-nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	-	10 (K2) 10 (K2)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their rela- tive position(s) in the molecule.	Walker, et al (1966)
	Nitrobenzene	Sewage organisms	BOD	-	630 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	3-nitro-4- methoxy- benzoic acid	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl = NT Ma = PT (3) So = PT (7) Cv = PT (3) Gp = T (3) Np = NT	Palmer and Maloney (1955)
CHEMICALS A	4'-nitro-o- salicylanisidide	Salmo gairdnerii Carassius auratus	BSA	-	10 (K 3 hr) 10 (K2)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their rela- tive position(s) in the molecule.	Walker, et al (1966)
	o-nitro- phenol	Lepomis macrochirus	BSA	-	46.3 - 51.6 (T2A)	<u>acdefgio</u>	Assays are completely described and autopsy data are reported.	Lammering and Burbank (1961)
TINES	p-nitrophenyl- hydrazine hydrochloride	Microcystis aeruginosa	L	-	50 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
	p-nitrophenyl- hydrazine	Microcystis aeruginosa	L	-	100 (K)	a, etc	Comment same as above.	Fitzgerald, et al (1952)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)	
AL CHEMICAL PRODUCTS	2' nitro-p- salicy lanilide	Salmo gairdnerii Carassius auratus	BSA	_	10 (K 3 hr) 10 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical structures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	Walker, et al (1966)	
	3-nitro-2',6'- salicyloxylidide	Salmo gairdnerii Carassius auratus	BSA	-	10 (K2) 10 (K2)	<u>a</u>	Comment same as above,	Walker, et al (1966)	
A-94	3-nitrosali- cylanilide	Salmo gairdnerii Carassius auratus	BSA	-	10 (K2A) 10 (K2A)	a	Comment same as above.	Walker, et al (1966)	APPENDIX A
	3-nitro-2',3- salicyloxylidide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K2A) 10.0 (K2A)	a	Comment same as above.	Walker, et al (1966)	IX A
	3-nitro-2',5- salicyloxyl- idide	Salmo gairdnerii Carassius auratus	BSA	-	10.0 (K 3 hr) 10.0 (K2)	<u>a</u>	Comment same as above.	Walker, et al (1966)	
	3-nitro-2',4'- salicyloxyl- idide	Salmo gairdnerii Carassius auratus	BSA	-	1.0 (K2) 10.0 (K 3 hr)	<u>a</u>	Comment same as above.	Walker, et al (1966)	

Nonyl phenol ethoxylate	Salmo gairdnerii (12 days after hatching) (25 days after hatching, fry) (210 days after hatching, fingerling)	BCFA	_	13.5 (K) 3 hr 5.2 (K) 6 hr 4.4 (K) 3 hr 2.3 (K) 6 hr 8.0 (K) 3 hr 5.2 (K) 6 hr	a c d e i	Successive developmental stages of the organism showed marked differences in resistance to the chemical. Changes in resistance could not be correlated with changes in respiratory activity of the fish but rather with their water metabolism.	Marchetti (1965)
p-octyl diphenylamine	Daphnia magna	BSA	_	≫40 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical sub- stances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
Oxydipro- pionitrile	Pimephales promelas Lepomis macrochirus Lebistes reticulatus	BSA	-	(H) 3600 (T4A) (S) 3900 (T4A) (S) 4200 (T4A) (S) 4450 (T4A)	c d e f	<ul> <li>(H) Value in hardwater</li> <li>(S) Value in softwater</li> <li>The chemical produced no change in flavor of the cooked bluegill.</li> </ul>	Henderson, et al (1960)
Oxalic acid	Daphnia magna	BSA	-	95 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentra- tion was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Oxalic acid	Sewage organisms	BOD	_	43 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Pentachloro- phenol	Green sunfish	BSA	-	(O)	-	Pentachlorophenol was repellent to the green sunfish at 20 mg/l but the fish were indifferent in response to 5.0 mg/l.	Summerfelt and Lewis (1967)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Gasterosteus aculeatus	BSA	-	(0)	c e	Tap water was used to make up the solutions. The fish avoided water more acid than a pH of 5.6 or one more alkaline than 11.4.	Jones (1948)
рН рН	Salmo gairdnerii	BSA	_	(0)	abcdefp	The pH value at which acid solutions proved lethal to rainbow trout within 1 day was unaffected by the pH value to which the fish had been acclimatized (pH 6.5-8.4). Fifty percent of a population of yearling rainbow trout were killed in about 1 day at a pH value of 3.6 when little free CO <sub>2</sub> was present; where in the presence of 50 ppm free CO <sub>2</sub> , a pH value of 5.6 killed 50 percent of a population of fingerling trout in 15 days. In water of low free CO <sub>2</sub> content, the relation between pH value and log median period of survival was linear for survival times between about 3 hr and 15 days. Exposure to pH values below 5.0 for about 3 months might be harmful to rainbow trout when little free CO <sub>2</sub> is present in the water.	Lloyd and Jordan (1964)
Phenanthra- quinone	Pomoxis nigromaculatus Notropis atherinoides atherinoides Hyborhynchus notatus Ambloplites rupestris rupestris Huro salmoides	BSA	_	(0)	<u>e</u>	Aerated spring water was used as the test medium. No effect was observed on fish after 2 days of exposure, even with excess solid dispersed in water. At algicidal concentrations, this compound was not toxic to the fish studied.	Fitzgerald, et al (1952)
o-phenanthro- line	Microcystis aeruginosa	L	-	100 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Phenazine-1- carboxylic acid	Anabaena flos-aquae Notemigonous crysoleucas	L	-	100 (O) 0.1 to 10.0 (O)	-	Value given is concentration for complete inhibition of <i>A. flos-aquae.</i> No harmful effect to <i>N. crysoleucas</i> was noted at the concentrations evaluated.	Toohey, et al (1965)
Phenol	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tempera ture, effect of dissolved oxygen, the efficiency of the gold- fish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 0.259 g/liter, fish survived 104 minutes.	
Phenol	Carassius auratus	BSA	-	125 to 372 (K 8 hr) 83.2 (O) 41.6 (O)	<u>a</u>	Temperature in test containers was maintained at $27 \pm .2$ C. Goldfish tested weighed between 2 and 4 g. Phenol, 83.2 ppm (mg per liter), killed 86% of the fish in 8 hr; 41.6 (mg per liter) killed 67% in 8 hr.	Gersdorff and Smith (1940)

	Phenol	<i>Anopheles quadrimaculatus</i> Goldfish Shiner minnows	BSA		(0)	-	Under the conditions of this experiment, this chemical (diluted 1 to 30) applied at rates of 10 to 95 gallons per acre was less effective than kerosene in controlling mosquitos. In the laboratory, at the rate of 50 gallon per acre, 100 percent of fish were killed but only 16 per- cent of the larvae. Phenol did not appear to be a desirable larvacide for general mosquito control.	Knowles, et al (1941)
	Phenoi	Carassius auratus	BSA		0.103 (K)	<u>a</u>	Goldfish weighed between 2 and 4 g. Temperature was maintained at 27.0 $\pm$ 0.2 C.	Gersdorff (1943)
	Phenol	Daphnia magna	BSA	-	94 (O)	<u>a</u> C	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Phenol	Hyborhynchus notatus	BSA	-	-	-	Fish in aquaria were trained to detect and distinguish between phenol and p-chlorophenol at levels as low as 0.0005 ppm. The fish could also distinguish o-chlorophenol from the two other compounds. The training method is described.	Hasler and Wisby (1949)
A-97	Phenol	Daphnia magna	BSA	-	28.9 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
	Phenol	Phoxinus phoxinus	BCFA	-	0.04% (K 4 min) 0.01% (K 8 min) 0.004% (K 24 min) 0.0004% (K 40-50 hr)	<u>a</u> c	Tap water was used as diluent. The apparatus used was a 34 mm diameter tube fitted to permit sharp vertical separation of water and test solutions. With this system, avoidance data could be obtained. Toxicity is given as average survival time of replicates. Fish did not avoid phenol in the $\leq$ 0.04% range.	Jones (1951)
CHEMICALS	Phenol	Semotilus atromaculatus	BSA	_	10 to 20 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
AND MI	Phenol	Lepomis macrochirus	BSA	-	20.5 (T4A) 19.3 (T2A)	<u>a</u> cde	Chu No. 14 modified medium was used as dilution water. The fish were transferred each 24 hours into new test solutions because of phenol loss due to aeration.	Trama (1955)
AND MIXTURES OF	Phenol	Lepomis macrochirus	BCFA	_	11.5 (T4A)	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. The phenol concentration was kept constant during the test period.	Cairns and Scheier (1955)
CHEMICALS	Phenol	Gambusia affinis	BSA	-	56 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Phenol	Sewage organisms	BOD	-	1600 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Phenol	Channel catfish (fingerlings)	BSA	_	16.7 (K 48 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Phenol	Lepomis macrochirus	BSA	-	11.5 (T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Phenol	Lepomis macrochirus	BSA	_	22.2 (T2A)	acdefgio	Assays are completely described, and autopsy data are reported.	Lammering and Burbank (1961)
Phenols (monohydric)	Salmo gairdnerii	BSA	_	(0)	<u>a e</u>	This is a study of the effect of varying dissolved oxygen concentrations on the toxicity of selected chemicals. The toxicity of heavy metals, ammonia, and monohydric phenols increased as the dissolved oxygen in water was reduced. The most obvious reaction of fish to lowered oxygen content is to increase the volume of water passed over the gills, and this may increase the amount of poison reaching the surface of the gill epithelium. The concentration of the chemical in the water was not specified.	Lloyd (1961)
Phenol	Hydropsyche Stenonema	BSA	-	30.0 (T2A) 14.5 (T2A)	а	Soft water used as diluent water.	Roback (1965)
Phenol	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	-	500 (K) 500 (K) 500 (K) 100 (NG) 100 (NG)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were expressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
Phenol	"Aquatic flora and fauna"	FR	Luxembourg	5.0-10.0 (O)	C e	Destruction of all flora and fauna of the river occurred in highly polluted zone (10 ppm), slight affects occurred at 3.0-10 ppm, and practically no damage occurred at con- centrations below 3.0 ppm.	Krombach an Barthel (1963)

\_

	Phenol	Rasbora heteromorpha	BSA	-	6.0 (O)	-	For many toxins the rate of mortality is found to be a linear function of the logarithm of the concentration of the poison; whereas the comparable relation between the logarithms of the survival time and the concentration is nonlinear. The linear function can be exploited to provide comparatively simple methods of estimating long-term survival concentra- tions. An application of this is suggested for defining realistic standards of toxicity. At the concentration listed, there was a 30 percent mortality in about 2 weeks.	Abram (1964)
	Phenol	Fish	BSA	-	1.4 x 10 <sup>-4</sup> M (K)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentrations are presented and discussed.	lshio (1965)
A-99	Phenol	Fish	FR	Ohio	.016 (O)	_	Following shut-down of steel mills due to a strike, phenols were 3.0 ppb in the Ohio River during the shut-down as compared to 16.0 ppb after the mills resumed operation. Threshold odor intensity and dissolved-iron content were 2 to 8X greater after start-up of the mills than during the shut-down period. Appearance or increased abundance of such "clean-water fish" as big-eye chub, common sucker, stoneroller, creek chub, sand shiner, mimic shiner, common shiner, and bluntnose minnow occurred while mills were shut down. Additionally, small minnows increased 20X during this period. The authors note that these facts are indicative of a marked betterment of the environment. Further, they suggest that the faunal monotony of the upper Ohio River is more closely related to industrial than to domestic discharges.	Krumholz and Minckley (1964)
G	Phenol	Carassius auratus	BCSA	_	(O)	<u>a</u>	A 5% solution of phenol in water was injected in the muscular masses of the fish tails at various levels. The MLD (minimal lethal dose) of phenol was found to be 230 mg/kg. Goldfish are unable to conjugate phenol, while showing a high efficiency in excreting the drug unchanged.	Boni (1965)
CHEMICALS	Phenols	Rainbow trout	FR	Scotland	4.4 (T2)	acefim	This work represents an extension of laboratory studies of the toxicity of complex effluents to investigations of rivers.	Herbert, et al (1965)
ALS AND MIXTURES OF	Phenol	Daphnia magna (young) Daphnia magna (adult) Lepomis macrochirus Mollienesia latopinna	BSA	-	17 (T1A) 7 (T2A) 61 (T1A) 21 (T2A) 63 (T1A) 22 (T2A)	<u>a</u> C	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
S OF CHEMICALS	Phenol	Pímephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	29 (T4A) 26 (T4A) 46 (T4A) 44 (T4A)	<u>a</u> cdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>M</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Phenol	Salmo gairdnerii	BSA	_	1.5 (T2A)	acdef	Test solution used in this study was sea water collected from the North Sea, then diluted with distilled water. Sensitivity of fish to poisoning by phenol increased as salinity increased.	Brown, et al (1967)
Phenol	Salmo gairdnerii Salmo salar	BSA	_	5.2 (T2)	a c d e f	Fish were acclimatized to 14 days in salt water.	Brown, et al (1967)
Phenol	Salmo gairdnerii	BSA	-	(O)	acdefp	Fish were acclimatized to the temperature of the test water over a period of 24-36 hr and then held at the test temper- ature without being fed for 24 hr before testing. Results showed that the resistance to poisoning by phenol increases with increase in temperature up to at least 18 C, at which the L2 is almost twice that at 6 C. A similar relationship exists with gas-liquor phenols. The response of test popula- tions showed the least viability at 12 C.	Brown, et al (1967)
Phenol	Nitzschia linearis Physa heterostropha Lepomis macrochirus	BSA	_	258 (T5A) 94.0 (T4A) 13.5 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Phenol	Salmo gairdnerii	BCFA	_	7.5 (T2A)	a c d e f	Phenol rapidly damaged the gills of trout. Experiments were conducted at levels above and below the LC <sub>50</sub> and for varying periods of time. Even at the level which killed only 20% of the fish in 48 hours, sufficient damage was done within one week to impair survival of the individual and affect reproduction. (This concentration was not specified, but was probably 6.5 ppm.)	Mitrovic, et al (1968)
Phenol	Salmo gairdnerii	BSA	_	4.58 to 5.8 (T2A)	acdefo	The concentration killing a half batch of fish in 2 days provides a reasonable estimate of the threshold concer- tration. The lethality of this chemical depends upon the temperature and concentration of dissolved oxygen.	Brown (1968)
Phenylhydra- zine hydro- chloride	Microcystis aeruginosa	L	~	100 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et a (1952)
4'-phenylazo- 3-nitrosali- cylanilide	Salmo gairdnerii Carassíus auratus	BSA	_	0.1 (K2A) 1.0 (K 3 hr) 1.0 (K2A) 10.0 (K2A)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicylanilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their rela- tive position(s) in the molecule.	Walker, et al (1966)

	p-phenylene- diamine	Daphnia magna	BSA	_	5.74 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
	PhenyImercuric acetate (10% soln.)	lctalurus punctatus	BSA	-	2.30 (K2) 1.46 (T2A)	acfi	The experiment was conducted at 68 C.	Clemens and Sneed (1958)
	Phenylmercuric acetate	Channel catfish (fingerlings)	BSA	-	4.1 (K1A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	PhenyImercuric hydroxide	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): C1 $-$ T (3) Ma $-$ T (3) So $-$ T (3) Gp $-$ T (3) Np $-$ T (3)	Palmer and Maloney (1955)
CHEMI	Pheny Imercuric nitrate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Comment same as above, including data cited.	Palmer and Maloney (1955)
CALS AND MI	n-phenyl-naphthyl- amine	Daphnia magna	BSA	-	4.4 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
XTURES	Phenylthiourea	Microcystis aeruginosa	L	~	50 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x $10^6$ to 2 x $10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
OF CHEMICAL	Phloroglucinol	Dəphnia magna	BSA	~	630 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Phosphoric acid	Gambusia affinis	BSA	-	138 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Phosphorus	Lepomis macrochirus	BSA	-	0.105 (T2A) 0.053 (T3A) 0.025 (T7A)	acdef ghijk no	Colloidal phosphorus compounds were removed by filtra- tion, so that the effect of elemental phosphate toxicity was studied.	lsom (1960)
o-ph thalic anhydride	Pimephales promelas	BSA		>56 (T4A)	<u>a</u> cdef	o-phthalic anhydride is very slightly soluble in water.	Pickering and Henderson (1966)
Picrıc acıd	Cylindrospermum licheniforme (CI) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI NT Ma NT So NT Cv NT Gp NT Np NT	Palmer and Maloney (1955)
Polyethylene glycol	Sewage microorganisms	BOD	_	(O)	_	The chemical was studied as to how low levels (ppm) may affect BOD in domestic sewage. This compound was not toxic to sewage microorganisms. No concentration of the chemical was given. Apparently this glycol is bio- chemically inert because it did not respond even to acclimated seed.	Oberton and Stack (1957)
Polyoxy- ethylene ester	Pimephales promelas (juveniles)	BSA	-	(S) 37-42 (T1-4A) (H) 38-56 (T1-4A)	<u>acdf</u>	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard water (H).	Henderson, et a (1959)
Potassium azide	Procambarus clarki Lepomis macrochirus	BSA	-	1 (K1)* 2 (K1)** <1.5 (T1A)* <1.8 (T1A)** *Technical formulation **Granular	a	In general, when mud was added to the tank the toxicity of the chemical decreased.	Hughes (1966)
Potassium azide	Pteronarcys californica (naiads)	BSA	-	0.008 (T4A)	<u>acdef</u>	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Potassium chloride	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography.	Powers (1918)

I	Potassium chloride	Daphnia magna	BSA	-	373 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
I	Potassium chloride	Daphnia magna	BSA	-	432 (O)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hr.	Anderson (1948)
	Potassium chloride	Lepomis macrochirus	BSA	-	2,010 (T4A)	a d e f	This paper reports the LD <sub>50</sub> in 96 hours for 8 common inorganic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured.	Trama (1954)
	Potassium chloride	Gambusia affinis	BSA	-	4,200 (T2A)	acdef	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Potassium chloride	Biomorpholaria a. alexandrina Bulinus truncatus	BSA	-	1800 (K1A) 1200 (K1A)	а	The degree of tolerance for vector snails of biharziasis chemicals is somewhat dependent upon temperature. The temperature at which (K1A) occurred was 26 C.	Gohar and El-Gindy (1961)
	Potassium chloride	Daphnia magna Lepomis macrochirus Lymnaea sp	BSA		679 (T1A) 5,500 (T1A) 1,941 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
5	Potassium chloride	Nitzschia linearis Lepomis macrochirus Physa heterostropha	BSA	_	1,337 (T5A) 940 (T4A) 2,010 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
HEMICALS AND	Potassium chromate	Salmo gairdnerii	BSA	-	(O) 2000 ppm (42.0 min) 1000 ppm (79 min) 20 ppm (3580 min)	<u>a</u> c <u>e</u> f	Tap or distilled water used as diluent. Toxicity defined as the avg. time when the fish lost equilibrium when exposed to the test chemical (ppm Cr).	Grindley (1946)
MIXTURES OF	Potassium chromate	Lepomis macrochirus	BCFA	-	450 (T4A) smali 630 (B4A) medium 5.50 (T4A) large	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. Beginning pH was 7.9 to 8.6, pH after four days was 7.0 to 7.94.	Cairns and Scheier (1955)
CHEMIC	Potassium chromate	Gambusia affinis	BSA	-	480 (T2A)	acdeg	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

nd

APPENDIX A

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen <b>ts</b>	Reference (Year)
Potassium chromate	Sewage organisms	BOD	-	10.5 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
Potassium chromate	Micropterus salmoides	BSA	-	195 (T2A)	<u>a c d e</u>	The mechanism for poisoning is discussed. Exposure to chromium caused severe pathological change in the intestine immediately posterior to the pyloric caeca that in all probability completely destroyed its digestive function.	Fromm and Schiffman (1958)
Potassium chromate	Lepomis macrochirus	BSA	_	550 (T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app 14-24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Potassium chromate	Salmo gairdnerii	BSA	_	100 (T1)	acdg	Trout exposed to 20 ppm chromium had a mean hematocrit of 43.8, as compared to unexposed trout of 31.8. Addi- tional data are presented.	Schiffman and Fromm (1959)
Potassium chromate	Pimephales promelas	BSA	-	(S) 45.6 (T4A)	c d e f	(S) Soft water Values are expressed as mg/l of chromium.	Pickering and Henderson (1965)
Potassium chromate	Nitzschia linearis Physa heterostropha Lepomis macrochirus	BSA	_	7.8 (T5A) 16.8 (T4A) 168.8 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Potassium cuprocyanide	Rhinichthys atratulus	BCFA	-	0.38, 0.47 and 0.71 (T1A)	a c e	The three values given are for cyanide to copper ratios of 4.0, 3.7, and 3.0, respectively.	Lipschuetz and Cooper (1955)
Potassium cyanide (as CN)	Rainbow trout (yearling)	BCFA	-	0.14 (K-160 min)	<u>a</u> c <u>e</u>	Toxicity was determined in terms of survival time. Acclimatization of fish to test conditions and fish size was studied.	Herbert and Merkens (1952)
Potassium cyanide	Microcystis aeruginosa	L	-	90 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Potassium cyanide	Rainbow trout (yearling)	BSA	-	0.105-0.155 (O)	a c <u>e</u>	Tap water was used as diluent. Study related oxygen con- centration effect to cyanide toxicity. As an example, control fish in 1.11 ppm $O_2$ were affected in 18 min; at 0.105 ppm CN <sup>-</sup> , fish survived only 3.3 min at 10% $O_2$ concentration.	Downing (1954)

Potassium cyanide	Salmo gairdnerii	BCFA		(0)	<u>a</u>	Time-survival curves are plotted for seven concentrations of cyanide, from 0.14 to 10 ppm. At 10 ppm, all fish died in less than 3 minutes. At 0.14 ppm all fish died in 165 minutes.	Herbert and Downing (1955)
Potassium cyanide	Rhinichthys atratulus meleagris	BCFA	_	0.22 (T1A)	асе	This report contains a comparison of the toxicities of KCN and potassium cuprocyanide of three different composi- tions. Four-hour median tolerance limits are also given.	Lipschuetz and Cooper (1955)
Potassium cyanide	Lepomis macrochirus	BCFA	_	0.55 (T46) smail 0.45 (T46) medium 0.57 (T46) large	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. The cyanide ion concentration was controlled.	Cairns and Scheier (1955)
Potassium cyanide	Gambusia affinis	BSA	-	1.6 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Potassium cyanide	Lepomis macrochirus	BSA	-	0.45 (T4A)	<u>a c e</u>	Increase in temperature seemed to increase toxicity of this chemical. Low dissolved oxygen reduced toxicity of some chemicals in this study. Toxicity values may be 20% higher in hard versus soft water.	Cairns (1957)
Potassium cyanide	Lepomis macrochirus Physa heterostropha	BSA	-	(N) 0.45 (T4A) (L) 0.12 (T4A) (N) 1.08 (T4A) (L) 0.48 (T4A)	<u>a e</u>	Modified Chu No. 14 test medium was used. Toxicity is given both for "normal" O <sub>2</sub> (5-9 ppm), (N), and with "low" O <sub>2</sub> (2 ppm DO), (L). High and low threshold concentration and concentration percent of survival are also presented.	Cairns and Scheier (1958)
Potassium cyanide	Sewage organisms	BOD	-	15 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Potassium cyanide (as CN <sup>-</sup> )	Brachydanio rerio (adults) (eggs) Lepomis macrochirus	BSA	-	0.49 (T2A) 117 (T2A) 0.16 (T2A)	<u>a c d e f</u>	The test dilutions were made up from distilled water and ACS grade chemicals. Temperature was held at 24 C and the solution was aerated to maintain a dissolved oxygen content of 5-9 ppm.	Cairns, et al (1965)
D Potassium cyanide	Lepomis macrochirus Physa heterostropha	BSA	-	0.45 (T4A) 0.12 (T4A) 1.08 (T4A) 0.48 (T4A)	a e	Normal oxygen content in water. Low oxygen content in water. Normal oxygen content in water Low oxygen content in water.	Cairns (1965)
Potassium Cryanide	Lepomis macrochirus	BSA	-	0.57 (T4A)	<u>acdei</u>	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app 14-24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Potassium cyanide	Lepomis macrochirus	BSA	-	0.43 (T4A)	acdef	The experiments were conducted in a water of controlled chemical composition. The TL <sub>m</sub> concentration of KCN was slightly affected bv increased temperature (more toxic at 30 C than at 18 C), but not by water hardness.	Cairns and Scheier (1963)
Potassium cyanide	Rasbora heteromorpha	BSA	-	0.072 (O)	-	For many toxins the rate of mortality is found to be a linear function of the logarithm of the concentration of the poison; whereas the comparable relation between the logarithms of the survival time and the concentration is nonlinear. The linear function can be exploited to provide comparatively simple methods of estimating long-term survival concentra- tions. An application of this is suggested for defining realistic standards of toxicity. At the concentration re- ported, there was a 20 percent mortality in 7 days.	Abram (1964)
Potassium cyanide	Daphnia magna Lymnaea sp (eggs)	BSA	-	2 (T1A) 0.7 (T3A) 0.4 (T4A) 796 (T1A) 147 (T3A) 130 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Potassium cyanide as (CN <sup>°</sup> )	Hydropsyche Stenonema	BSA	-	2.0 (T2A) 0.5 (T2A)	а	Soft water used as diluent water.	Roback (1965)
Potassium dichromate	Daphnia magna	BSA	-	<0.6 (0)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentra- tion was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Potassium dichromate	Salmo gairdnerii	BSA	-	2000 ppm – 23.8 min 1000 ppm – 54.6 min 200 ppm – 188 min 20 ppm – 4342 min	<u>a</u> c <u>e</u> f	Tap or distilled water used as diluent. Toxicity defined as the avg time when the fish lost equilibrium when exposed to the test chemical (ppm Cr).	Grindley (1946)
Potassium dichromate	Lepomis macrochirus	BCFA	-	320 (T4A)	a c e f	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. The pH of the test water was about 6.2, which was determined by the concentration of the test chemical.	Cairns and Scheier (1958)
Potassium dichromate	Gambusia affinis	BSA	-	320 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

	Lepomis macrochirus	BSA	_	320 (T4A)	ace	Increase in temperature seemed to increase toxicity of this chemical. Low dissolved oxygen reduced toxicity of some chemicals in this study. Toxicity values may be 20% higher in hard versus soft water.	Cairns (1957)
	Lepomis macrochirus	BSA		(N) 320 (T4A) (L) 320 (T4A)	<u>a e</u>	Modified Chu No. 14 test medium was used. Toxicity is given both for "normal" $O_2$ (5-9 ppm), (N), and with "low" $O_2$ (2 ppm DO), (L). High and low threshold concentration and concentration percent of survival are also presented.	Cairns and Scheier (1958)
Potassium dichromate	Lepomis macrochirus	BSA	_	320-384 (T4A)	acdef	The concentration of $K_2Cr_2O_7$ which resulted in 50 percent kill in 96 hours was 320 ppm in soft water at both 18 and 30 C, 382 ppm in hard water at 18 C, and 369 ppm in hard water at 30 C.	Cairns and Scheier (1959)
Potassium dichromate	Lepomis macrochirus	BSA	_	320 (T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app 14-24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Potassium dichromate	Sewage organisms	BOD	_	17.0 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Potassium dichromate	Hydropsyche Stenonema	BSA	-	28.0 (T2A) 3.5 (T2A)	а	Soft water used as diluent water.	Roback (1965)
Potassium dichromate	Lepomis macrochirus	BSA	-	320 (T4A) 320 (T4A)	a e	Normal oxygen content of water. Low oxygen content of water.	Cairns (1965)
Potassium dichromate	Carassius carassius Daphnia magna Lepomis macrochirus	BSA	-	705 (T1A) 0.4 (T4A) 739 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Potassium dichromate	Brachydanio rerio (adults) (eggs) Lepomis macrochirus	BSA	-	180 (T2A) 1500 (T2A) 440 (T2A)	acdef 	The test dilutions were made up from distilled water and ACS grade chemicals. Temperature was held at 24 C and the solution was aerated to maintain a dissolved oxygen content of 5-9 ppm.	Cairns, et al (1965)
Potassium dichromate	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	(S) 17.6 (T4A) (H) 27.3 (T4A) (S) 118.0 (T4A) (H) 133.0 (T4A) (S) 37.5 (T4A) (S) 30.0 (T4A)	c d e f	(S) Soft water (H) Hard water Values are expressed as mg/l of chromium.	Pickering and Henderson (1965)
	Potassium dichromate Potassium dichromate Potassium dichromate Potassium dichromate Potassium dichromate	dichromate macrochirus Potassium Lepomis dichromate macrochirus Potassium Lepomis dichromate macrochirus Potassium Lepomis dichromate Sewage organisms Potassium Lepomis dichromate Stenonema Potassium Lepomis dichromate carassius Daphnia magna Lepomis macrochirus Potassium Brachydanio dichromate rerio (adults) (eggs) Lepomis macrochirus Potassium dichromate Pimephales macrochirus Carassius auratus Lebistes	dichromate     macrochirus       Potassium dichromate     Lepomis macrochirus     BSA       Potassium dichromate     Sewage organisms     BOD       Potassium dichromate     Kenonema organisms     BSA       Potassium dichromate     Lepomis macrochirus     BSA       Potassium dichromate     Lepomis macrochirus     BSA       Potassium dichromate     Rearassius macrochirus     BSA       Potassium dichromate     BSA     BSA       Potassium dichromate     Praney Silve macrochirus     BSA       Potassium dichromate     Brachydanio (eggs) Lepomis macrochirus     BSA       Potassium dichromate     Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes     BSA	dichromate macrochirus BSA – Potassium Lepomis BSA – dichromate macrochirus BSA – Potassium Lepomis BSA – dichromate macrochirus BSA – Potassium Lepomis BSA – dichromate macrochirus BSA – Potassium Sewage BOD – of tassium Lepomis BSA – Potassium Lepomis BSA – dichromate Stenonerma BSA – Potassium Lepomis BSA – dichromate macrochirus BSA – Potassium Lepomis BSA – dichromate BSA – Potassium Lepomis BSA – Potassium Carassius BSA – Potassium Brachydanio BSA – dichromate Promis macrochirus BSA – Potassium Brachydanio BSA – dichromate Promis BSA – dichromate Brachydanio	dichromate       macrochirus       Dot       Dot         botassium       Lepomis       BSA       -       (N) 320         dichromate       macrochirus       BSA       -       (T4A)         botassium       Lepomis       BSA       -       320-384         dichromate       macrochirus       BSA       -       320 (T4A)         Potassium       Lepomis       BSA       -       320 (T4A)         dichromate       macrochirus       BSA       -       320 (T4A)         Potassium       Lepomis       BSA       -       320 (T4A)         dichromate       organisms       BOD       -       17.0 (TC50)         Potassium       Lepomis       BSA       -       320 (T4A)         dichromate       Stenonema       3.5 (T2A)       320 (T4A)         otassium       Lepomis       BSA       -       320 (T4A)         otassium       Lepomis       BSA       -       320 (T4A)         otassium       Lepomis       BSA       -       320 (T4A)         otassium       Lepomis       739 (T1A)       magna         Lepomis       739 (T1A)       180 (T2A)       1600 (T2A)         dichromate	dichromate macrochirus BSA – (N) 320 sec (T4A) sec (T4A) (L) 320 (T4A) sec (T4A) sec	dichromate       mecrochirus       BSA       -       (N) 320       a chemical. Low disolved oxygen reduced toxicity of some chemicals in this stury. Toxicity values may be 20% higher in hard versus 507 water.         votassium       Leponis       BSA       -       (N) 320       (2 pm D) (L) (L) sign and low threshold concentration in full water.         votassium       Leponis       BSA       -       320.384       a c d e f       The concentration of KycryD which resulted in 50 percent distribution percent of survival are stop presented.         votassium       Leponis       BSA       -       320 (T4A)       a c d e f       The concentration of KycryD, which resulted in 50 percent distribution percent of survival are stop presented.         votassium       Leponis       BSA       -       320 (T4A)       a c d e i       A "control" was prepared by adding required chemicals to distribution water at 10 C. and 30 C. 320 pm in hard water at 30 C.         votassium       Leponis       BSA       -       17.0 (TC5p)       a       A "control" was prepared by adding required chemicals to distribution are also in the report.         votassium       Sewage       BOD       -       17.0 (TC5p)       a       The purpose of this paper was to devise a toxicity index concentration at system content in histing to the concentration at system content in histing to the paper distribution at the system content in histing to the concentric tration producing 50 percent inhibition tore in length. Dat

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Potassium dichromate	Nitzschia linearis Physa heterostropha Lepomis macrochirus	BSA	_	0.208 (T4A) 17.3 (T4A) 113.0 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Potassium ferricyanide	Daphnia magna	BSA	-	905 (T1A) 549 (T2A) 0.6 (T3A) 0.1 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Potassium hydroxide	Gambusia affinis	BSA	-	80 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Potassium hydroxide	Biomorpholaria a. alexandrina Bulinus truncatus Lymnaea caillaudi	BSA	-	500 (K1A) 300 (K1A) 150 (K1A)	а	The degree of tolerance for vector snails of biharziasis to chemicals is somewhat dependent upon temperature. The temperature at which (K1A) occurred was 27 C.	Gohar and El-Gindy (1961)
Potassium nitrate	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of temper- ature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.00002N solution, fish survived 2135 minutes.	Powers (1918)
Potassium nitrate	Gasterosteus aculeatus	BSA	_	50 (K10)	-	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
Potassium nitrate	Lepomis macrochirus	BSA	_	3,000 (T4A)	a d e f	This paper reports the LD <sub>50</sub> in 96 hours for 8 common inorganic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured.	Trama (1954)
Potassium nitrate	Gambusia affinis	BSA	-	224 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Potassium nitrate	Biomorpholaria a. alexandrina Bulinus truncatus	BSA	-	2600 (K1A) 1800 (K1A)	а	The degree of tolerance for vector snails of biharziasis to chemicals is somewhat dependent upon temperature. The temperature at which (K1A) occurred was 28 C for <i>Bulinus</i> and 25 C for <i>Biomorpholaria</i> .	Gohar and El-Gindy (1961)

	Potassium nítrate	Daphnia magna	BSA		900 (T4A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations	Dowden and Bennett
		Lepomis macrochirus			5,500 (T1A)		were made in various types of water.	(1965)
		Lymnaea sp (eggs)			1,941 (T1A)			
	Potassium permangante	Daphnia magna	BSA	-	0.63 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Potassium permanganate	Gambusia affinis	BSA	-	12 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Potassium permanganate	Channel catfish (fingerlings)	BSA	-	<3.2 (K1A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Potassium permanganate	Lepomis macrochirus Semotilus atromaculatus	BSA	_	4.2 (T1,2,4A) 3.7 (T4A)	-	The values given are for a laboratory study. However, when concentrations as high as 32 ppm were applied in a pond, no fish deaths occurred.	Kemp, et al (1966)
CHEMICALS AND MIXTURES OF CH	Potassium permanganate	Blue-green algae Cylindrospermum Anabaena Anacystis Calothrix Nostoc Oscillatoria Plectonema Green algae Ankistrodesmus Chlorella Closterium Oocystis Green algae Scenedesmus Stigeoclonium Zygnema Green flagellate and yellow algae Chalmydomonas Pandorina Tribonema Gomphonema Navicula Nitzchia	L		4.0-8.0 (O)	-	KMnO4 was toxic or partially toxic at the indicated concentra- tions to blue-green and green algae. A concentration of 8.0 ppm was usually required to control green, flagellate, and yellow algae.	Kemp, et al (1966)
FMICA	Potassium phosphate	Gambusia affinis	BSA		750 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Potassium sulfate Potassium	Lepomis macrochirus	BSA	_	3,550 (T4A)	a d e f	This paper reports the LD <sub>50</sub> in 96 hours for 8 common inorganic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured.	Trama (1954)
	Carassius auratus	BSA	-	(O)	ac	A 0.5% solution in water prolonged the mortality of sperm for at least 5 minutes in all samples tested. A 0.5% solution in frog Ringer's produced similar mortility patterns but average activity was lower after 10 minutes than in water solution.	Fribourgh (1965)
Propion- hydroxamic acid	Microcystis aeruginosa	L	-	100 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
, Propionic acid	Culex sp (larvae) Daphnia magna Lepomis macrochirus	BSA	_	1000 (T2A) 50 (T2A) 188 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
n-propyl alcohol	Semotilus atromaculatus	BSA	-	200 to 500 (CR)	a e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concen- tration in ppm below which the 4 test fish lived for 24 hrs. and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Propylene phenoxetol	Pleuronectes platessa	BSA	-	(O)	<u>a</u>	<ul> <li>Fish were tested at 6.5 C in aquariums of 3-liter capacity. At 0.05% solution, the fish were able to survive if removed to fresh water within 1 hour after exposure.</li> <li>At 15 C and 0.005% solution, the fish took 2 hours to become completely anesthetized and were unable to recover after 3 hours of exposure.</li> <li>At 15 C and 0.025% solution, the fish were not able to survive if not removed within 1 hour. The chemical can be used as an anesthetic for periods of up to 1 hour when a solution of 0.01-0.025% is used.</li> </ul>	Bagenal (1963)
n-propyl-N,N- di-n-propyl thiol-carbamate	Elodea canadensis Potamogeton nodosus Potamogeton pectinatus	BSA	_	5 (O) 100 (O) 5 (O) 100 (O) 5 (O) 100 (O)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. No toxic effect. Injury rating of 9.4. No toxic effect. Injury rating of 7.4 No toxic effect. Injury rating of 8.3	Frank, et al (1961)

	Pyridine	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of temper- ature, effect of dissolved oxygen, the efficiency of the gold- fish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In a concentration of 3.187 cc per liter, fish survived 180 minutes.	Powers (1918)
	Pyridine	Gambusia affinis	BSA	-	1,350 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Pyridine	Daphnia magna	BSA	-	2,114 (T1A) 944 (T2A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
	Pyridyl- mercuric acetate	Rainbow trout	FL	Wash.	2.0 (0)	-	After the first treatment with the chemical the ponds were partially emptied, flushed, and refilled. After a second treatment, one pond showed a "catastrophic mortality". The authors were unable to explain this unusual phenomenon.	Foster and Olson (1951)
A_111	Pyridyl- mercuric acetate (tech.)	Salmo gairdnerii	BSA	-	10 (K 17% 1 hr) 47 F 10 (K 50% 1 hr) 56 F 5 (K 1-1/2% 1 hr) 47 F 5 (K 18% 1 hr) 56 F 2.5 (K 0% 1 hr) 47 F 2.5 (K 1% 1 hr) 56 F	<u>a</u>	Temp concentration data presented on groups of 200 fingerlings. Brook and Brown trout not affected by the test conc. of 10, 5, and 2.5 ppm at either 47 F or 56 F for 1 hr.	Rodgers, et al (1951)
유	Pyridyl- mercuric acetate (80% active)	lctalurus punctatus	BSA	-	5.0 (K2) 3.8 (T2A)	acfi	The experiment was conducted at 75 C.	Clemens and Sneed (1958)
CHEMICALS AND MIXTURES OF CHEMICALS	Pyridyl- mercuric acetate	Channel catfish (fingerlings)	BSA	-	4.12 (T2A) 2.81 0.49 2.81 (T3A) 1.81 <.37 2.43 (T4A) <.37 <.37	<u>a</u>	The toxicity of this compound increased as the temperature was increased. In the data shown, the values for each T level is for temperatures of 10, 16.5 and 24 centigrade. These values were selected from a table presenting con- centrations for T levels from one to 153 hours. Fish of different ages were also studied.	Clemens and Sneed (1959)
JRES OF	Pyridyl- mercuric acetate	Channel catfish (fingerlings)	BSA	-	3.8 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
- CHEMICALS	Pyrocatechol	Daphnia magna	BSA	-	14 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)

14

A-111

CHEMICALS	Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES	Pyrogallol	Daphnia magna	BSA	_	18 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
ES O	Quinacrine hydro-	Salmo gairdneri	BSA	_	17.2 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
т С	chloride	Salmo trutta			230 (T2A)			(1000)
ĒM		Salvelinus			230 (T2A)			
OF CHEMICALS		fontinalis Salvelinus			21.0 (T2A)			
S		namaycush Ictalurus			70.0 (T2A)			
		punctatus Lepomis macrochirus			79.0 (T2A)			
	Quinine sulphate	Channel catfish (fingerlings)	BSA	-	42 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
•	Quinhydrone	Microcystis aeruginosa	L	_	100 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
,	Quinone	Microcystis aeruginosa	L	_	100 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et al (1952)
	Resorcinol	Daphnia magna	BSA	_	56.4 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
	Salicylaldehyde	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlcrella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI $-$ PT (3) Ma $-$ PT (3) So $-$ PT (3) Cv $-$ PT (3) Gp $-$ T (3), PT (21) Np $-$ T (3), PT (21)	Palmer and Maloney (1955)

	Salicylic acid	Sewage organisms	BOD	_	110 (ТС <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	Selenium	Black bullhead Bluegill Channel catfish Largemouth bass Rainbow trout White crappie Yellow walleye	FL	Sweitzer Lake, Colo.	_	a c	It was tentatively concluded on the basis of the available data that fish kill probably resulted from the toxic effects of selenium, possibly acting in synergism with other ions such as uranium or zinc. Arsenic was also found in the lake. Samples of flora and fauna of the lake were analyzed and found to contain greater than 300 ppm selenium. It was believed that selenium is passed up the food chain to the fish which accumulated the element in lethal concentrations.	Barnhart (1958)
	Silver, colloidal	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI = PT (3) Ma = PT (14) So = NT Cv = NT Gp = NT Np = NT	Patmer and Maloney (1955)
CHEMICALS AND MIX	Silver, colloidal, (33 percent silver nitrate)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Comment same as above except that: CI - T (3) Ma - T (3) So - T (3) Cv - T (3) Gp - T (3) Np - T (3)	Palmer and Maloney (1955)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
Silver	Lebistes reticulatus Bufo valliceps (tadpoles)	BSA	_	0.01 (K) 0.1 (K)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the	Shaw and Grushkin (1967)
	Daphnia magna			0.1 (K)		metal ion is plotted against the log of the solubility product constant of the metal sulfide – a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	
Silver- cynaide complex	<i>Lepomis macrochirus</i> (juveniles)	BSA	-	(K <1.0)	<u>a</u> cdf <u>p</u>	With 10 ppm as cyanide content, the median resistance time varied from 391 to 789 minutes.	Doudoroff, et a (1966)
Silver nitrate	Gasterosteus aculeatus	BSA	-	0.003 (K10)	-	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
Silver nitrate	Daphnia magna	BSA	-	0.0051 (O)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hours.	Anderson (1948)
Silver nitrate	Sewage organisms	BOD	-	0.3 (O)	-	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Sheets (1957)
Silver sulfate	Balanus balanoides	BSA	_	0.4 (O)	_	The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
Sodium acetate	Polycelis nigra	BSA	_	0.15 M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.2. Solutions were renewed every 12 hours.	Jones (1941)
Sodium acetate	Daphnia magna	BSA	-	<5800 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This salt may be toxic only when the concentration is great enough to exert an unfavorable osmotic effect.	Anderson (1946)
Sodium acetate	Lepomis macrochirus Culex sp. (Iarvae)	BSA	-	5,000 (T1A) 7,500 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Sodium aluminate	Gambusia affinis	BSA	_	126 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

Sodium anthra- quinone alpha- sulfonate	<i>Daphnia magna Lymnaea</i> sp (eggs)	BSA	-	12 (T1A) 186 (T1-4A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Sodium anthra- quinone- a-sulfonate	Daphnia magna	BSA	-	(0)	ac	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 7.1. The 100-hr threshold was 12%, with 0 percent toxicity at 10% and 100 percent toxicity at 30%.	Freeman (1953)
Sodium arsenate	Polycelis nigra	BSA	-	0.0048 M (L2)	C	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.2. Solutions were renewed every 12 hours.	Jones (1941)
Sodium arsenate (as As <sub>2</sub> O <sub>3</sub> )	Smallmouth black bass Largemouth black bass Bluegill sunfish White crappie Potomogeton crispus P. foliosus Najas flexilis Anarchis canadensis Nymphea sp Scirpus validus Chara sp Hydrodictyon sp Oedogonium sp Cladophora sp	FL	Leetown, Va.	5.0 (O)	d	Treatment of a series of ponds resulted in control of <i>P. crispus</i> , <i>P. foliosus</i> , <i>N. flexilis</i> , and <i>A. canadensis</i> . <i>Nymphea</i> sp, <i>S. validus</i> , and <i>Chara</i> sp were not controlled. Scum algae ( <i>Hydrodictyon</i> sp, <i>Oedogonium</i> sp, and <i>Cladophera</i> sp) in solid mats were effectively destroyed by the arsenate. Decomposing vegetation stimulated growth of more desirable algae. No fish mortality occurred due to toxic effect of chemical, but some fish suffocated due to decay- ing vegetation.	Surber and Everhart (1950)
Sodium arsenate	Daphnia magna	BSA	-	31 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Sodium arsenate	Phoxinus phoxinus	BSA	-	2970 ppm (205 min) 820 ppm (467 min) 234 ppm (951 min)	<u>ace</u> f	Tap or distilled water used as diluent. Toxicity defined as the avg time when the fish lost equilibrium when exposed to the test chemical (ppm As).	Grindley (1946)
Sodium arsenate	Daphnia magna	BSA	-	<20 (0)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This salt may be toxic only when the concentration is great enough to exert an un- favorable osmotic effect.	Anderson (1946)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium arsenate	Sewage organisms	BOD	-	>100 (TC <sub>50</sub> )	a	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
Sodium arsenite or arsenious oxide	Caenis sp Callibaetis sp Libellula sp Ischnura verticalis Chironomidae Asellus communis Hydracarina sp Hyalella knickerbockeri Colpidium sp Paramecium sp Stylonichia sp Spirogyra sp	BSA	_	3.0 (K) 4.0 (K) 14.0 (56% survival) 11.2 (85% survival) 2.96 (83% survival) 21 (81% survival) 10.5 (94% survival) 5.88 (30% survival) 3.5 (100% survival) 1.75 (plasmolysis but no kill)	а	River water was used as test media with room temperature and natural sunlight as environmental conditions. Considerable additional data are presented.	Surber and Meeham (1931)
Sodium arsenite	Phoxinus phoxinus	BSA	_	953 ppm (54.6 min) 290 ppm (186 min) 17.8 ppm (2174 min)	<u>a c e</u> f	Tap or distilled water used as diluent. Toxicity defined as the avg time when the fish lost equilibrium when exposed to the test chemical (ppm As).	Grindley (1946)
Sodium arsenite	Daphnia magna	BSA	-	9.1 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This salt may be toxic only when the concentration is great enough to exert an un- favorable osmotic effect.	Anderson (1946)
Sodium arsenite	Notropsis hudsonius	BSA	-	45 (T1A) 29 (T2A) 27 (T3A)	a c d e	Some of the fish were not killed in 72 hours by the higher doses of arsenic (30-35 ppm), had extensive damage to the fins, while others had scale damage, severe diarrhea, heavy breathing and hemorrhaging of the body areas around the caudal, dorsal, and ventral fins.	Boschetti and McLoughlin (1957)

	Sodium arsenite	Pithophora sp Hydrodictyon sp Bottom organisms Lepomis macrochirus Microcrustacea Rotifers	FL	Ala. ponds	4.0 (O) 4.0 (O) 4.0 (O)	-	The purpose of this experiment was to determine the effec- tiveness of sodium arsenite as a control agent for <i>Pithophora</i> and to determine the effects of repeated applications of 4 and 8 ppm arsenious oxide as sodium arsenite on bottom organisms and fish production in treated ponds. <i>Pithophora</i> was controlled by one or more applications of sodium arsenite at a concentration of 4.0 ppm arsenious oxide. Best results were obtained when sodium arsenite was applied while the alga was in an active growing stage. The alga <i>Hydrodictyo</i> was also controlled at 4.0 ppm. The applications of 4 ppm applied 1 month apart reduced the number of bottom organisms an average of 34 percent and reduced bluegill pro- duction an average of 42 percent as compared with those of the controlled ponds.	Lawrence (1958) n
	Sodium arsenite	Notemigonus crysoleucas Pimephales promelas Lepomis macrochirus	FPCH	N.Y.	4.0 (S23) 4.0 (S23) 4.0 (S23)	a c d	Conventional farm ponds were used having an average surface area of 0.3 acre and a maximum depth of 7-9 ft. Toxicity (in ppm) to fish as maximum safe concentration (S) for 23 days was determined. Concentration of 0.5 ppm was required to control algae.	Eipper (1959)
	Sodium arsenite	Channel catfish (fingerlings)	BSA	-	47.9 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Sodium arsenite	<i>Lepomis</i> sp	FL	Ponds in Ala.	(0)	-	Fish from ponds treated with sodium arsenite were analyzed for arsenic when the concentration in the water had declined to less than 1.0 ppm arsenious oxide. Bluegill sunfish analyzed for arsenic were recovered by seining when the arsenious oxide concentration in the pond water had declined to less than 1.0 ppm. Arsenic in the digestive tract of bluegills from the ponds ranged from 2.1 to 6.6 ppm arsenious oxide (wet weight). However, no detectible arsenic or only a trace amount was found in the tissue of the digestive tract, liver, or muscle.	Dupree (1960)
CHEMICALS	Sodium arsenite	Calico fish	FL	N.Y.	(0)	-	Fish were analyzed for arsenic, before and after the lakes were treated with this herbicide. No differences in residues were noted.	Ullmann (1961)
	50-51 (sodium arsenite)	Water Hyssop Parrot's Feather Bladderwort	FL	Lakes in Fla.	(O) (O) (O)	-	A concentration of 10.0 ppm controlled the indicated species.	Phillippy (1961)
AND MIX	50-52 (sodium arsenite)	Water Hyssop Parrot's Feather Bladderwort	FL	Lakes in Fla.	(O) (O) (O)	-	Comment same as above.	Phillippy (1961)
MIXTURES C	Sodium arsenite (tech.)	Rainbow trout Bluegill	BSA	-	26 (T4A) 30 (T4A)	а	This is an estimated LC $_{\!$	Cope (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium	Filamentous algae Cladophora Spirogyra Zygnema Submerged plants	FL	N.Y.	4 (K) 4 (K) 4 (K)	a c	Complete decomposition in about 2 weeks. Complete decomposition in about 2 weeks. Complete decomposition in about 2 weeks.	Cowell (1965)
) )	Chara Potamogeton			(O) (O)		Sodium arsenite, 4 ppm, did not cause any kill. Sodium arsenite, 4 ppm, caused 95% kill. Decomposition occurred in about 1 month.	
	Emergent plants <i>Alisma</i> <i>Sagittaria</i> Zooplankton			(0) (0) (0)		Sodium arsenite, 4 ppm, caused 15% kill. Sodium arsenite, 4 ppm, did not cause any kill. Applications of 4 ppm sodium arsenite produced significant reduction.	
Sodium arsenite	<i>Pteronarcys</i> sp (nymphs)	BSA	_	45 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
Sodium arsenite	Salmo gairdnerii	FL	La Cross, Wis.	25 (T4A)	acfim	The herbicide used was a commercial formulation containing 40 percent sodium arsenite by weight. Substantial residues	Gilderhus (1966)
	Carassius auratus Lepomis macrochirus			34 (T4A) 35 (T4A)		of arsenic were found in the water, bottom soil, and throughout the organs and flesh of the bluegills at the termination of the experiment. Treatments totaling 4.0 ppm or more resulted in reduced numbers of bottom fauna, and a concentration of 1.2 ppm of the chemical controlled rotifers.	
Sodium arsenite	<i>Daphnia magna</i> Rainbow trout Bluegill	BSA	_	6.5 (5.7-7.3) (O) 60 (O) 60 (O) 44 (O)	acdiq	Toxicity, in terms of median immobilization concentration $(IC_{50})$ , is presented for <i>Daphnia;</i> median lethal concentration ( $LC_{50}$ ) values for rainbow trout and bluegill are reported.	Crosby and Tucker (1966)
Sodium arsenite	Salmo gairdneri Lepomis macrochirus	BSA	-	36.5 (T2A) 44.0 (T2A)	а	This paper reports acute toxicity of a number of com- pounds, and discusses subacute mortality as well. Effects on reproduction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Соре (1966)
	Pteronarcys californicus Daphnia			80.0 (T2A) 1.8 (T2A)			
	pulex Simocephalus serrulatus			1.4 (T2A)			
Sodium arsenite	Simocephalus serrulatus	BSA	-	1.4 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr.	Sanders and Cope
0.301110	Daphnia pulex			1.8 (SB)		Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	(1966)

Sodium arsenit		Blue-green algae Cylindrospermum Anabaena Anacystis Calothrix Nostoc Oscillatoria Plectonema Green algae Ankistrodesmus Chlorella Closterium Oocystis Green algae Scenedesmus Stigeoclonium Zygnema Green flagellate and yellow algae Chalmydomonas Pandorina Tribonema Gomphonema Navicula Nitzchia	L	_	2.0 (0)	_	NaAsO2 was generally nontoxic or only partially toxic briefly for all algae species. Growth of <i>Cylindrospermum</i> and <i>Nitzchia</i> was apparently stimulated. This compound was the least effective of four evaluated as algicides.	Kemp, et al (1966)
Sodium arsenit		Lepomis macrochirus	BSA	-	0.7 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerable limit for a selected group of herbicides.	Hughes and Davis (1967)
Sodium arsenit (tech.)	te )	Pteronarcys californica (naiads)	BSA	_	0.038 (T4A)	acdef	Data reported as LC $_{\!$	Sanders and Cope (1968)
Sodium azide CHEMICALS	1	Procambarus clarki Lepomis macrochirus	BSA	-	1.0 (K1)* 1.0 (K1)** 1.5 (T1A)* 1.8 (T1A)** *Technical formulation **Granular	а	In general, when mud was added to the tank the toxicity of the chemical decreased.	Hughes (1966)
Sodium azide	1	Pteronarcys californica (naiads)	BSA	-	0.0092 (T4A)	<u>acdef</u>	Data reported as LC $_{\!$	Sanders and Cope (1968)
RES	า nesulfonate	Daphnia magna	BSA	-	(O)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 7.1. The 100-hr threshold was 2840%, with 0 percent toxicity at 1895% and 100 percent toxicity at 8000%.	Freeman (1953)
OF Sodium CHEMICA		Daphnia magna	BSA	_	<650 (0)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This salt may be toxic only when the concentration is great enough to exert an unfavorable osmotic effect.	Anderson (1946)

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURE	Sodium benzoate	Sewage organisms	BOD	-	(NTE)	-	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC50) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
S OF	Sodium o-benzoyl sulfimide (soluble saccharin)	Sewage organisms	BOD	-	>1000 (тс <sub>50</sub> )	<u>a</u>	Comment same as above.	Hermann (1959)
CHEMICALS	Sodium bicarbonate	Polycelis nigra	BSA	_	0.085 M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.4. Solutions were renewed every 12 hours.	Jones (1941)
A-120	Sodium bicarbonate	Daphnia magna	BSA	-	4200 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
20	Sodium bicarbonate	Daphnia magna	BSA	_	2350 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This report toxic value may be due to an unfavorable osmotic effect.	Anderson (1946)
	Sodium bicarbonate	Lepomis macrochirus	BCFA	_	8,250 (T4A) small 8,600 (T4A) medium 9,000 (T4A) large	a c e f	Test water was composed of distilled water through CP grade chemicals and was aerated throughout the 96-hour exposure period. At pH 7, the ratio of bicarbonate to carbonate was 2270:1.	Cairns and Scheier (1955)
	Sodium bicarbonate	Gambusia affinis	BSA	-	7,550 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Sodium bicarbonate	Lepomis macrochirus	BSA	-	9000 (T4A)	<u>acdei</u>	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app. 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
	Sodium bicarbonate	Culex sp (larvae)	BSA	_	2,000 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)

	Sodium bicarbonate	Nitzschia linearis Lepomis macrochirus	BSA	-	650 (T5A) 8,600 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
	Sodium bisulfate	Daphnia magna	BSA	_	190 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Toxic effect may be a result of lowering the pH below 6.0.	Anderson (1946)
	Sodium bisulfate	Daphnia magna	BSA	-	153.4 (O)	a C	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inorganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
	Sodium bisulfate	<i>Culex</i> sp (larvae)	BSA		300 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
A-121	Sodium bisulfite	Daphnia magna	BSA	-	<145 (0)		This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
21	Sodium bisulfite	Daphnia magna	BSA	-	102 (O)	ac	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
	Sodium bisulfite- Sodium sulfate	Daphnia magna	BSA	_	82 (O) 3642 (O)	a c	Comment same as above.	Freeman and Fowler (1953)
CHEMICALS	Sodium bisulfite- Sodium carbonate	Daphnia magna	BSA	_	850 (O) 436 (O)	a c	Comment same as above.	Freeman and Fowler (1953)
CALS AND MIXTURES	Sodium bisulfite- Sodium carbonate- Sodium chromate	Daphnia magna	BSA	_	87 (O) 440 (O) 0.35 (O)	a c	Comment same as above,	Freeman and Fowler (1953)
ę	Sodium bisulfite- Sodium carbonate- Sodium silicate	Daphnia magna	BSA	-	38 (O) 194 (O) 92 (O)	a c	Comment same as above.	Freeman and Fowler (1953)
CHEMICALS	Sodium bisulfite- Sodium silicate	Daphnia magna	BSA	-	177 (O) 427 (O)	a c	Comment same as above.	Freeman and Fowler (1953)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium bisulfite- Sodium chromate	Daphnia magna	BSA	_	70 (O) 0.286 (O)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium bisulfite- Sodium silicate- Sodium sulfate	Daphnia magna	BSA	-	52 (O) 126 (O) 2308 (O)	a c	Comment same as above.	Freeman (1953)
Sodium bisulfite- Sodium chromate- Sodium silicate	Daphnia magna	BSA	-	144 (O) 0.861 (O) 506 (O)	a c	Comment same as above.	Freeman (1953)
Sodium bisulfite- Sodium carbonate- Sodium sulfate	Daphnia magna	BSA	-	58 (O) 295 (O) 2562 (O)	a c	Comment same as above.	Freeman (1953)
Sodium bisulfite- Sodium chromate- Sodium sulfate	Daphnia magna	BSA	-	75 (O) 0.306 (O) 3312 (O)	a c	Comment same as above.	Freeman (1953)
Sodium bisulfite	Daphnia magna	BSA	-	61.4 (O)	a c	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inorganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
Sodium bisulfite	Gambusia affinis	BSA	-	240 (T2A)	a c d e g	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium bisulfite	Daphnia magna (young) Daphnia magna	BSA	-	116 (T2A) 102 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
	(adult) <i>Dugesia</i> sp			179 (T4A)			

		Lymnaea sp (eggs) Mollienesia			179 (T1A) 241 (T1A)			
	Sodium bisulfite plus sodium silicate	latopinna Daphnia magna	BSA	_	950-14,210 (T1A) 785-11,723 (T2A) 15-22 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. The two TL <sub>m</sub> values are the respective concentration of each of the chemicals listed.	Dowden and Bennett (1965)
	Sodium bisulfite plus sodium carbonate	Daphnia magna	BSA	-	436 (T4A) 85 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
	Sodium bisulfite plus sodium chromate	Daphnia magna	BSA	-	68 (T4A) 0.278 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
	Sodium bisulfite plus sodium sulfate	Daphnia magna	BSA	_	82 (T4A) 3,654 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
A-123	Sodium bisulfite plus sodium carbonate and sodium chromate	Daphnia magna	BSA	-	86 (T4A) 441 (T4A) 0.354 (T4A)	ac	Comment same as above.	Dowden and Bennett (1965)
ç	Sodium bisulfite plus sodium chromate and sodium sulfate	Daphnia magna	BSA	-	78 (T4A) 0.32 (T4A) 3,443 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
CHEMICALS AND	Sodium bisulfite plus sodium carbonate and sodium silicate	Daphnia magna	BSA	-	39 (T4A) 198 (T4A) 93 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
MIXTURES OF	Sodium bisulfite plus sodium chromate and sodium silicate	Daphnia magna	BSA	_	224 (T4A) 0.086 (T4A) 506 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
F CHEMICALS	Sodium bisulfite plus sodium carbonate and sodium sulfate	Daphnia magna	BSA	-	57 (T4A) 296 (T4A) 2,869 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium bisulfite plus sodium silicate and sodium sulfate	Daphnia magna	BSA	-	52 (T4A) 126 (T4A) 2,326 (T4A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each of the three $TL_m$ values represents the concentration of each of the chemicals, respectively.	Dowden and Bennett (1965)
Sodium borate Sodium borate	Polycelis nigra	BSA	-	0.026 M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.8. Solutions were renewed every 12 hours.	Jones (1941)
Sodium borate	Daphnia magna	BSA	_	<240 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Threshold value may be only half of that reported.	Anderson (1946)
Sodium borate (ore)	Salmo gairdnerii	BSA	-	2800 (T1A) 1800 (T2A)	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
Sodium borate	Gambusia affinis	BSA		8,200 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium bromate	Polycelis nigra	BSA	-	0.020 M (L2)	c	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Jones (1941)
Sodium bromate	Daphnia magna	BSA	_	210 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium bromide	Polycelis nigra	BSA	_	0.14 M (L2)	C	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Jones (1941)
Sodium bromide	Daphnia magna	BSA	-	8200 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This salt may show toxicity when the concentration is high enough to exert unfavorable osmotic effect.	Anderson (1946)
Sodium p-bromo- benzene- sulfonate	Daphnia magna	BSA	-	843 (K)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 6.9.	Freeman (1953)

Sodiu p-br	ım omo-	Daphnia magna	BSA		523 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations	Dowden and Bennett
benz		Lepomis macrochirus			1,560 (T1A)		were made in various types of water.	(1965)
		Lymnaea sp (eggs)			2,590 (T1-4A)			
Sodiu n-bu sulfo		Daphnia magna	BSA	_	7,827 (K)	a c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 7.1.	Freeman (1953)
Sodiu buty sulfe		Daphnia magna	BSA	-	8,000 (T1A) 5,400 (T3A) 2,700 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Sodiu buty		Lepomis macrochirus	BSA	-	5,000 (T1A)	ac	Comment same as above.	Dowden and Bennett (1965)
Sodiu carb	ım onate	Daphnia magna	BSA	-	424 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Sodiu carb	ım onate	Micropterus salmoides Lepomis macrochirus Goldfish	BSA	-	500 (O) 500 (O) 500 (O)	<u>a</u> cfpi	The disposal of cannery wastes frequently involves the use of chemicals for treatment purposes. Ferrous sulphate, alum, and lime are used in chemical coagulation; sodium carbonate for acidity control in biological filters; and sodium nitrate in lagoons for odor control. Lye (sodium hydroxide) peeling of certain fruits and vegetables is not uncommon. These chemicals, in whole or part, are dis- charged in most cases to a stream. The concentrations listed permitted largemouth bass to survive 7 to 9 hours, bluegills to survive 4.5 to 11 hours, and goldfish to survive indefinitely.	Sanborn (1945)
Sodiu carb	m onate	Daphnia magna	BSA	-	<424 (0)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Toxic effect may be due in part to the rise in pH to 9.2.	Anderson (1946)
Sodiu Carb	im onate	Oncorhyncus tshawytscha Oncorhyncus kisutch Salmo	BSA		68 (K5) 70 (K5) 80 (K5)	<u>a</u> de	This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et al (1952)
Sodiu Corb	im onate	clarkii Daphnia magna	BSA	~	524 (O)	ас	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodiu carb	m onate	Lepomis macrochirus	BCFA	-	300 (T4A)	a c e f	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hr exposure period. Toxicity was essentially determined by pH. At pH 10 the carbonate to bicarbonate ratio was 1:2.27.	Cairns and Scheier (1955)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium carbonate	Daphnia magna	BSA	_	552.4 (O)	a c	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inorganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmo- spheric dissolved oxygen.	Fairchild (1955)
Sodium carbonate	Gambusia affinis	BSA	-	840 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with ''high'' turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium carbonate	Lepomis macrochirus	BSA	-	300 (T4A)	a c e d i	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, app. 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Sodium carbonate	Amphipoda Culex sp (larvae) Daphnia magna Dugesia sp Lepomis macrochirus Lymnaea sp. (eggs) Mollienesia latopinna	BSA	-	360 (T1A) 1,820 (T1A) 347 (T1A) 607 (T1A) 384 (T1A) 385 (T1A) 403 (T1A) 405 (T2A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
Sodium carbonate	Nitzschia linearis Lepomis macrochirus	BSA	_	242 (T5A) 320 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Sodium carbonate- Sodium chromate	Daphnia magna	BSA	-	408 (O) 0.33 (O)	ac	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium carbonate plus sodium chromate	Daphnia magna	BSA	-	420 (T4A) 0.34 (T4A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each value represents the concentration of each respective chemical.	Dowden and Bennett (1965)
Sodium carbonate- Sodium silicate	Daphnia magna	BSA	_	180 (O) 85 (O)	ас	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)

Sodium	Daphnia	BSA		005 (74 4)			
carbonate plus sodium silicate	magna	BSA	-	265 (T1A) 130 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each TL <sub>m</sub> value is equal to the concentration of each respective chemical.	Dowden and Bennett (1965)
Sodium carbonate- Sodium sulfate	Daphnia magna	BSA	_	221 (O) 1,918 (O)	ac	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium carbonate plus sodium sulfate	Daphnia magna	BSA		198 (T1A) 666 (T1A) 172 (T2A) 577 (T2A) 66 (T3A) 222 (T3A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each $TL_m$ value is equal to the concentration of each respective chemical.	Dowden and Bennett (1965)
Sodium carbonate- Sodium chromate- Sodium silicate	Daphnia magna	BSA	_	182 (O) 0.146 (O) 86 (O)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium carbonate plus sodium chromate and sodium silicate	Daphnia magna	BSA	-	187 (T4A) 0.15 (T4A) 88 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each $TL_m$ value represents the concentration of each respective chemical.	Dowden and Bennett (1965)
Sodium carbonate- Sodium chromate- Sodium sulfate	Daphnia magna	BSA	-	240 (O) 0.192 (O) 2079 (O)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period,	Freeman and Fowler (1953)
Sodium carbonate plus sodium chromate and sodium sulfate	Daphnia magna	BSA	-	240 (T4A) 0.19 (T4A) 2,078 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each TL <sub>m</sub> value represents the concentration of each respective chemical.	Dowden and Bennett (1965)
Sodium carbonate- Sodium silicate- Sodium sulfate	Daphnia magna	BSA	_	155 (O) 73 (O) 1343 (O)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium carbonate plus sodium silicate and sodium sulfate	Daphnia magna	BSA	-	161 (T4A) 76 (T4A) 1,396 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Each $TL_m$ value is equal to the concentration of each respective chemical.	Dowden and Bennett (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen <b>ts</b>	Reference (Year)
Sodium carboxyethyl rosin amine	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	ä	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl $-$ PT (14) Ma $-$ PT (14) So $-$ NT Cv $-$ NT Gp $-$ T (3) Np $-$ NT	Palmer and Maloney (1955)
Sodium chlorate	Polycelis nigra	BSA	-	0.15 M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.4. Solutions were renewed every 12 hours.	Jones (1941)
Sodium chlorate	Daphnia magna	BSA	~	4240 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium chlorate	Salmo gairdnerii	BSA	_	4200 (T1A) 2750 (T2A)	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
Sodium chloride	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of temper- ature, effect of dissolved oxygen, the efficiency of the gold- fish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.27N solution, the fish survived 178 minutes.	Powers (1918)
Sodium chloride	Polycelis nigra	BSA	-	0.19 M (L2)	c	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.0. Solutions were renewed every 12 hours.	Jones (1941)
Sodium chloride	Daphnia magna	BSA	_	6143 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various substances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)

	Sodium chloride	Brook trout	BSA	_	(O)	_	Fish were fed NaCl in gelatin capsules in amounts of 5.0 to 25.0 mg. Fish averaged 5.6 grams in weight. Physical effects of the salt were exhibited rather than true toxicity. Fish were also immersed in NaCl solution. Immersion in a 2.5% solution produced no increase in blood salt concentration. A 30-minute bath in 3.0% salt or a 10-minute bath in 5.0% salt caused a rise in blood salinity that quickly returned to normal when the fish were placed in fresh water. A 60-minute bath in 3.0% salt resulted in a very high blood salt level that required 48 hours to return to normal. A 15-minute bath in a 5.0% solution resulted in the loss of the majority of the fish.	Phillips (1944)
	Sodium chloride	Daphnia magna	BSA	-	<4200 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
	Sodium chloride	Daphnia magna	BSA	-	3,680 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hours.	Anderson (1948)
A-129	Sodium chloride	Lepomis macrochirus	BSA	-	12,946 (T4A)	a d e f	This paper reports the LD <sub>50</sub> in 96 hours for 8 common inorganic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured. If this salt is toxic to fish, this experiment did not demonstrate it. A saturated solution of 2,980 ppm produced no significant mortalities.	Trama (1954)
	Sodium chloride	Daphnia magna	BSA	-	5,093 (O)	a c	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inorganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
CHEMICALS AND MIXTURES OF	Sodium chloride	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - NT Ma - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
S OF CHEMICALS	Sodium chloride	Biomorpholaria a. alexandrina Bulinus truncatus Lymnaea caillaudi	BSA	-	4100 (K1A) 2600 (K1A) 2600 (K1A)	а	The degree of tolerance for vector snails of biharziasis chem- icals is somewhat dependent upon temperature. The tem- perature at which (K1A) occurred was 26 C.	Gohar and El-Gindy (1961)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium chlori <b>d</b> e	Gambusia affinis	BSA	-	18,100 (T2A)	a c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium chloride	Limnodrilus hoffmeisteri Erpobdella punctata Helisoma campanulata Gyraulus circumstriatus Physa heterostropha Sphaerium cf. tenue Asellus communis Argia sp	BSA	_	6200 (T4A) 7500 (T4A) 6150 (T4A) 3200 (T4A) 3500 (T4A) 5100 (T4A) 6200 (T4A) 1100 (T4A) 1150 (T4A) 8250 (T4A) 24,000 (T4A)	<u>a</u> cdi	Most of the data developed was with hard water, but experi- ments with soft water were also conducted. Additional TL <sub>m</sub> data are presented.	Wurtz and Bridges (1961)
Sodium chloride	Hydropsyche Stenonema	BSA	-	9,000 (T2A) 2,500 (T2A)	а	Soft water used as diluent water,	Roback (1965)
Sodium chloride	Cyprinidae Asellus sp Hydropsyche sp Dressenia sp Calliriche sp Helosciadium sp Nodiflorum sp Oenanthe fluviatilis Lemna	BSA	_	10,000 (L10A) 10,000 (L7 and K4FA) 10,000 (L6 and K17A) 10,000 (L5A) 10,000 (K13A)	<u>a</u>	<i>L. trisulca</i> was not affected at 10,000 ppm.	Vivier and Nisbet (1965)
Sodium chloride	trisulca Nais spp	BSA	-	1.0% (T 36 min)	a f	All tests were conducted in hard water. Time given is median survival time of the worms.	Learner and Edwards (1963)
Sodium chloride	Potamogeton pectinatus	BSA	-	(0)	-	Increasing NaCl solutions produced a proportional adverse effect on vegetative growth and seed production, but a concentration of 3000 ppm stimulated the production and growth of tubers. 9000 ppm completely inhibited the growth of one-week-old plants. 15,000 ppm reduced growth completely and was fatal to many plants.	Teeter (1965)

	Sodium chloride	Carassius carassius	BSA		13,750 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua-	Dowden and Bennett
		Culex sp (larvae)			10,500 (T1A)		tions were made in various types of water.	(1965)
		Daphnia magna			6,447 (T1A)			
		Lepomis macrochirus			14,125 (T1A)			
		<i>Lymnaea</i> sp			3,412 (T1A)			
		(eggs) Mollienesia latopinna			18,735 (T1A)			
	Sodium chloride	Nitzschia linearis	BSA	-	2,430 (T5A)	ace	The purpose of this experiment was to determine whether there was a constant relationship between the responses of	Patrick, et al (1968)
		Lepomis macrochirus			12,940 (T4A)		these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	
	Sodium p- chlorobenzene sulfonate	Daphnia magna	BSA		3,007 (K)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 7.1.	Freeman (1953)
	Sodium p- chlorobenzene	Daphnia magna	BSA	-	2,394 (T4A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua-	Dowden and Bennett
	sulfonate	Lepomis macrochirus			3,219 (T1A)		tions were made in various types of water.	(1956)
		Lymnaea sp (eggs)			8,600 (T1A)			
	Sodium 2- chlorotoluene- 4-sulfonate	Lepomis macrochirus	BSA	_	1,374 (T1A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
	Sodium 2- chlorotoluene- 5-sulfonate	Daphnia magna (young)	BSA	-	0.8 (T1A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
CHEM	5-surronate	Daphnia magna (adult)			3.3 (T1A)			(1905)
		Lymnaea sp			30. (T1A)			
CHEMICALS AND		(eggs) Mollienesia latopinna			115.2 (T1A)			
MIXTURES	Sodium chromate	Polycelis nigra	BSA	-	0.0028M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.2. Solutions were renewed every 12 hours.	Jones (1941)
	Sodium chromate	Sewage organisms	BOD	-	1.0 (O)	j	"Toxicity" is expressed as 10 percent reduction in oxygen utilization.	Ingols (1955)
HEMICA	Sodium chromate	Daphnia magna	BSA	-	<0.32 (0)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
_								

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium chromate	Daphnia magna	BSA	_	0.42 (0)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium chromate	Sewage organisms	BOD	-	(O)	_	A concentration of 1.0 ppm produced an oxygen depletion in percent of the control of 90%. It required 10.0 ppm to pro- duce 38% oxygen depletion. There is an apparent relation- ship between toxicity of chromium and the organic matter concentration in that higher amounts of organic matter com- plex with the chromium thus reducing its apparent toxicity.	Ingols (1954)
Sodium chromate	Daphnia magna	BSA	-	0.51 (O)	a c	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inorganic salts known to be present in various industrial effluents. Anal- ysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
Sodium chromate	Gambusía affinis	BSA	-	500 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium chromate	Escherichia coli Saccharomyces ellipsoides	L	-	(0)	_	This study suggests that the chromates have an effect on microbial genetic expression. Toxicity appeared to be in the range of 100 to 500 mg/I.	Ingols and Fetner (1961)
Sodium chromate	Nereis sp Carcinus maenas Leander squilla	BSA	-	0.5 (SB 21) 1.0 (SB 21) 60.0 (T12A) 50.0 (SB 12) 5.0 (SB 35)	а	The threshold toxicity for shore crabs was in the range of 40 to 60 ppm for a 12-day period of exposure. The threshold toxicity for prawns was a little less than 10 ppm in adults and 5 ppm in young.	Raymont an Shields (1964)
Sodium chromate- Sodium silicate-	Daphnia magna	BSA	_	0.201 (O) 119 (O)	ac	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium sulfate				2180 (O)			
Sodium chromate-	Daphnia magna	BSA	-	0.276 (O)	a c	Comment same as above.	Freeman an Fowler
Sodium sulfate				2984 (O)			(1953)
Sodium chromate- Sodium silicate	Daphnia magna	BSA	-	0.159 (O) 93 (O)	ас	Comment same as above.	Freeman an Fowler (1953)

Sodium chromate plus sodium silicate	Daphnia magna	BSA	-	0.21 (T4A) 130 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water. Each TL <sub>m</sub> value is equal to the concentration of each respective chemical.	Dowden and Bennett (1965)
Sodium chromate plus sodium sulfate	Daphnia magna	BSA	-	0.28 (T4A) 3,044 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
Sodium chromate plus sodium silicate and sodium sulfate	Daphnia magna	BSA	_	0.28 (T4A) 122 (T4A) 2,255 (T4A)	<u>a</u> c	Comment same as above.	Dowden and Bennett (1965)
Sodium citrate	Polycelis nigra	BSA	-	0.015M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Jones (1941)
Sodium citrate	Daphnia magna	BSA	-	825 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium cyanide	Polycelis nigra	BSA	_	0.0006M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 4.8. Solutions were renewed every 12 hours.	Jones (1941)
Sodium cyanide	Daphnia magna	BSA	-	<3.4 (0)		This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium Cyanide	Pimephales promelas	BSA	-	0.23 (T4A)	<u>a</u> c	Synthetic soft water was used. Toxicity data given as number of test fish surviving after exposure at 24, 48, and 96 hr. TL <sub>m</sub> values were estimated by straight-line graphical in- terpolation and given in ppm CN <sup>-</sup> .	Doudoroff, et al (1956)
Sodium cyanide	Sewage organisms	BOD	-	3.6 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
Sodium cyanide	Lepomis cyanellus	FL	Carbon- dale, III.	1.0 (K1)	а	Green sunfish placed in cages in ponds 1 and 2 days after application of the chemical suffered 100 percent mortality at 1.0 ppm. Toxicity seemed to be less in waters exhibiting high pH or low temperature.	Bridges (1958)

;

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium cyanide	Lepisosteus osseus Carassius auratus Cyprinus carpio Ictalurus natalis Micropterus salmoides Lepomis cyanellus	BSA	_	1.0 (K <1)	a c e	After application of 1 ppm of the chemical to small farm ponds, fish began to surface within 5 to 30 minutes. At concentrations of 1 ppm and at a variety of temperature and pH conditions, effective kills of a number of different species of warm-water fishes were produced. Concentrations of 1 ppm produced complete kill of all species of fish within 8 hr.	Bridges (1958)
Sodium cyanide	Pimephales promelas Lepomis macrochirus	BSA	-	(H) 0.35 (T4A) (S) 0.23 (T4A) (H) 0.15 (T4A)	c d e f	(H) Value in hardwater. (S) Value in softwater.	Henderson, et al (1959)
Sodium cyanide	Gasterosteus aculeatus Anguilla anguilla Phoxinus phoxinus Salmo trutta Carassius auratus	BSA	-	0.49 (K 8 hr) 0.49 (K 12 hr) 0.49 (K 6 hr) 0.49 (K 2 hr) 4.9 (K 12 hr)	<u>a</u> ce	This rather long paper deals more with behavior (avoidance reaction time, etc.) than other aspects of toxicity. However, interpolation from several curves resulted in the concentra- tions quoted. Avoidance occurred at concentrations as low as 10 <sup>-6</sup> N.	Costa (1965)
Sodium cyanide	Gammarus pulex	BCFA	-	(O)	<u>a</u> ce	Temperature and pH were important factors determining the behavior and reaction time of <i>Gammarus</i> during exposure to solutions of this chemical. Most of the data were de- scribing behavioral responses. However, in a solution of 0.00005N, the fish survived 1-1/2 hours. <i>Gammarus</i> were somewhat more resistant to sodium cyanide than fish.	Costa (1965)
Sodium cyanide	Rana temporaria	BCFA	-	(O)	<u>a</u> e	This report deals more with behavioral aspects than strict toxicity. The response limit for frog tadpoles is about 0.49 ppm. Increased temperature, a higher pH, and the amount of dissolved oxygen were critical. The response limit for tadpoles was 0.00001N. The tadpoles were less sensitive than fish but more sensitive than <i>Gammarus</i> .	Costa (1965)
Sodium cyanide	Green sunfish	BSA and FL	Okla.	(O)	_	Sodium cyanide was found to be moderately effective as a repellent at 5 mg/l and to produce an avoidance response at 1.0 mg/l. No response was noted at or below 0.5 mg/l.	Summerfelt and Lewis (1967)
Sodium 2,5- dichloro- benzene- sulfonate	Daphnia magna	BSA	-	3,890 (K)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 7.1.	Freeman (1953)

Sodium 2,5- dichloro-	Daphnia magna	BSA	-	1,468 (T4A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua-	Dowden and Bennett
benzene sulfonate	Lepomis macrochirus			3,750 (T4A)		tions were made in various types of water.	(1965)
	Lymnaea sp (eggs)			4,513 (T4A)			
Sodium dichromate	Gambusia affinis	BSA	-	420 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium dichromate	Daphnia magna	BSA	-	22 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium dinitrophenate	Phoxinus phoxinus	BSA	-	250 ppm (17.7 min) 100 ppm (61.0 min) 50 ppm (209.0 min)	<u>a</u> c <u>e</u> f	Tap or distilled water used as diluent. Toxicity defined as the avg time when the fish lost equilibrium when exposed to the test chemical (ppm dinitrophenate).	Grindley (1946)
Sodium ferrocyanide	Polycelis nigra	BSA	-	0.0008M (L2)	c	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.4. Solutions were renewed every 12 hours.	Jones (1941)
Sodium ferrocyanide	Daphnia magna	BSA	-	<600 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium fluoride	Polycelis nigra	BSA	~	0.0011M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.2. Solutions were renewed every 12 hours.	Jones (1941)
Sodium Afluoride	Daphnia magna	BSA	~	504 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium A fluoride	Gambusia affinis	BSA	-	925 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium fluoride MXX TCC RE	Rainbow trout	BSA	-	5.9-7.5 (T2A)* 2.6-6.0 (T2A)** *45 F *55 F	<u>a</u>	This study postulates that temperature affects the toxicity of fluoride concentration because of its effect on the metabolic rate of the fish. $TL_m$ values are given as $LC_{50}$ .	Anonymous (1966)
Sodium G fluoride G	Homarus americanus	BSA	-	0.9-4.5 (SB10)	a c e	Fluoride was not toxic even at levels five times those gen- erally used in municipal water supplies. The lobsters employed weighed 500 grams.	Stewart and Cormick (1964)
Sodium formate	Daphnia magna	BSA	-	<5200 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Toxic effect may be a result of unfavorable osmotic effect.	Anderson (1946)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Røference (Year)
Sodium formate	Lepomis macrochirus	BSA	-	5,000 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium hydrosulfide	Gambusia affinis	BSA	-	206 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium hydrosulfide	Semotilus atromaculatus	BSA	-	4 to 10 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentra- tion in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Copeland and Woods (1959)
Sodium hydroxide	Polycelis nigra	BSA	-	0.000004M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.8. Solutions were renewed every 12 hours.	Jones (1941)
Sodium hydroxide	Daphnia magna	BSA	_	240 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Sodium hydroxide	Micropterus salmoides (large mouth bass) Lepomis macrochirus Goldfish	BSA	_	50 (O) 50 (O) 50 (O)	<u>a</u> cfpi	The disposal of cannery wastes frequently involves the use of chemicals for treatment purposes. Ferrous sulphate, alum, and lime are used in chemical coagulation; sodium carbonate for acidity control in biological filters; and sodium nitrate in lagoons for odor control. Lye (sodium hydroxide) peeling of certain fruits and vegetables is not uncommon. These chemicals, in whole or part, are discharged in most cases to a stream. The concentrations listed permitted fish to survive indefinitely.	Sanborn (1945)
Sodium hydroxide	Daphnia magna	BSA	_	156 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Toxic effect may be due to the rise in pH to 9.1-9.5.	Anderson (1946)
Sodium hydroxide	Oncorhyncus tshawytscha Oncorhyncus kisutch Salmo clarkii clarkii	BSA	-	48 (K5) 20 (K5) 35 (K5)	<u>a</u> d e	This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et a (1952)
Sodium hydroxide	Semotilus Atromaculatus	BSA	_	20 to 40 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional date are presented.	Gillette, et a (1952)

	Sodium hydroxide	Lepomis macrochirus	BCFA	-	(O)	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period. At pH 9.8, all fish survived. At pH 9.9 to 10.1 after 4 days, only one-half survived. At pH 10.41 to 10.50, only 10 percent survived after 3 days.	Cairns and Scheier (1955)
	Sodium hydroxide	Lepomis gibbosus	BSA	-	5 (K 3-5 min)	C	The author suggests placing pellets of sodium hydroxide in the nests of the sunfish when eggs or fry are present. This method for controlling sunfish was developed first in the laboratory in petri dishes and later conducted in the field.	Jackson (1956)
	Sodium hydroxide	Lepomis gibbosus	FL	Durham, N. H.	5 (K 3-5 min)	а	The chemical must be applied after spawning begins and before the fry leave the nest. The author suggests placing pellets of sodium hydroxide in the nest of the sunfish when eggs or fry are present.	Jackson (1956)
	Sodium hydroxide	Gambusia affinis	BSA	-	125 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Sodium hydroxide	Lepomis macrochirus	BSA	-	9.9 (pH, T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, approximately 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
A-137	Sodium hydroxide	Biomorpholaria a. alexandrina Bulinus truncatus Lymnaea caillaudi	BSA	-	450 (K1A) 150 (K1A) 150 (K1A)	а	The degree of tolerance for vector snails of biharziasis to chemicals is somewhat dependent upon temperature. The temperature at which (K1A) occurred was 27 C.	Gohar and El-Gindy (1961)
Q	Sodium iodate	Polycelis nigra	BSA	-	0.0013M (L2)	C	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 8.0. Solutions were renewed every 12 hours.	Jones (1941)
HEMI	Sodium iodide	Polycelis nigra	BSA	-	0.044M (L2)	с	Comment same as above.	Jones (1941)
CHEMICALS AN	Sodium iodide	Daphnia magna	BSA	_	3.3 (0)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
D M	Sodium iodate	Daphnia magna	BSA	-	<158 (0)	-	Comment same as above except value may be only half of that reported.	Anderson (1946)
AND MIXTURES OF C	Sodium metaarsenite	Sewage organisms	BOD	-	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
CHEMICALS	Sodium mono- hydrogen phosphate	Daphnia magna	BSA	-	1,154 (T1A) 1,089 (T2A) 426 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)

CHEMICALS	Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTURES	Sodium mono- hydrogen phosphate plus sodium pyrophosphate	Daphnia magna Lymnaea sp (eggs)	BSA	_	3,580 (T1A) 433 (T1A) 2,685 (T1A) 63 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water. Each TL <sub>m</sub> value is equal to the concentration of each re- spective chemical.	Dowden and Bennett (1965)
9 F	Sodium napthalene B-sulfonate	Daphnia magna	BSA	-	308 (K)	<u>a</u> c	Assay water was not characterizied chemically or otherwise described. The pH at 100 percent toxicity was 7.1.	Freeman (1953)
CHEMICALS	Sodium nitrate	Carassius carassius	BSA	-	(O)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.220N solution, fish survived 171 minutes.	Powers (1918)
	Sodium nitrate	Gasterosteus aculeatus	BSA	-	500 (K10)	-	Solutions were made up in tap water. 3.0 to 5.0 cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (1939)
	Sodium nitrate	<b>P</b> olycelis nigra	BSA	-	0.043M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.2. Solutions were renewed every 12 hours.	Jones (1941)
	Sodium nitrate	Daphnia magna	BSA	-	8,500 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Sodium nitrate	Micropterus salmoides	BSA	-	4,000 (O)	acfpi	The disposal of cannery wastes frequently involves the use of chamicals for tractment ourposes. Farrow subbate alum	Sanborn
	muale	Lepomis			2,000 (O)		chemicals for treatment purposes. Ferrous sulphate, alum, and lime are used in chemical coagulation; sodium carbonate	(1945)
		<i>macrochirus</i> Goldfish			2,000 (O)		for acidity control in biological filters; and sodium nitrate in lagoons for odor control. Lye (sodium hydroxide) peeling of certain fruits and vegetables is not uncommon. These chemicals, in whole or part, are discharged in most cases to a stream. The concentrations listed permitted large mouth bass to survive indefinitely, bluegills to survive 3 days to indefinitely, and goldfish to survive 4 days.	
	Sodium nitrate	Daphnia magna	BSA	-	5,000 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Toxic effect may be caused when the chemical concentration is high enough to exert unfavorable osmotic effect.	Anderson (1946)

APPENDIX A

	Sodium nitrate	Lepomis macrochirus	BSA	-	12,000 (T4A)	a d e f	This paper reports the $LD_{50}$ in 96 hours for 8 common inorganic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured. If this salt is toxic to fish, this experiment did not demonstrate it.	Trama (1954)
	Sodium nitrate	Lepomis macrochirus	BCFA	-	9,500 (T4A)	acef	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour exposure period.	Cairns and Scheier (1955)
	Sodium nitrate	Gambusia affinis	BSA	-	10,000 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity on the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Sodium nitrate	Lepomis macrochirus	BSA	_	9,000 (T4A)	acdei	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data re- ported are for larger fish, approximately 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
	Sodium nitrite	Sewage organisms	BOD	_	(NTE)	_	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic con- centration producing 50 percent inhibition (TC <sub>50</sub> ) of oxygen utilization as compared to controls. Five toxi- grams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	Sodium nitrate	Biomorpholaria a. alexandrina Bulinus truncatus	BSA	-	6,000 (K1A) 3,100 (K1A)	а	The degree of tolerance for vector snails of biharziasis to chemicals is somewhat dependent upon temperature. The temperature at which (K1A) occurred was 28 C for <i>Bulinus</i> and 26 C for <i>Biomophalaria</i> .	Gohar and El-Gindy (1961)
CHEM	Sodium nitrate	Carassius carassius Daphnia magna Lepomis macrochirus Lymnaea sp (eggs)	BSA	-	12,150 (T1A) 4,206 (T4A) 12,800 (T1A) 6,375 (T1A) 5,950 (T2A) 3,251 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
CHEMICALS AND	Sodium nitrite	Polycelis nigra	BSA	-	0.0006M (L2)	C	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.0. Solutions were renewed every 12 hours.	Jones (1941)
MIXTUR	Sodium nitrite	Daphnia magna	BSA	-	<20 (0)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in entrifuged Lake Erie water.	Anderson (1946)

1

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen <b>ts</b>	Reference (Year)
Sodium nitrite	Semotilus atromaculatus	BSA	_	400 to 2000 (CR)	<u>a</u> e 	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Sodium nitrite	Gambusia affinis	BSA	-	7.5 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium m- nitrobenzene sulfonate	Daphnia magna Lepomis macrochirus	BSA	-	2,235 (T4A) 1,350 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium m- nitrobenzene sulfonate	Daphnia magna	BSA	_	5,618 (K)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 8.6.	Freeman (1953)
Sodium 4- nitrochloro- benzene-2- sulfonate	Daphnia magna Lepomis macrochirus Lymnaea sp (eggs)	BSA	-	1,474 (T4A) 6,375 (T4A) 3,532 (T1A) 3,208 (T2A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium 4- nitrochloro- benzene-2- sulfonate	Daphnia magna	BSA	_	3,187 (K)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 6.9.	Freeman (1953)
Sodium nitroprusside	Polycelis nigra	BSA	-	0.0008M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.0. Solutions were renewed every 12 hours.	Jones (1941)
Sodium nitroprusside	Daphnia magna	BSA	-	<210 (0)	-	This assay is based on concentration of the chemical re- quired to immobilize the test animal. Assays were con- ducted in centrifuged Lake Erie water. Value may be half of that reported.	Anderson (1946)
Sodium 4- nitrotoluene- 2-sulfonate	Lepomis macrochirus	BSA	-	1,440 (T1A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium oxalate	Polycelis nigra	BSA	-	0.011m (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.2. Solutions were renewed every 12 hours.	Jones (1941)

	Sodium oxalate	Daphnia magna	BSA	_	214 (O)	_	This assay is based on concentration of the chemical re- quired to immobilize the test animal. Assays were con- ducted in centrifuged Lake Erie water.	Anderson (1946)
	Sodium oxalate	Gambusia affinis	BSA		1,350 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Sodium oxalate	Sewage organisms	BOD	_	(NTE)	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concen- tration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depict- ing the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	Sodium oxalate	Lepomis macrochirus	BSA	-	4,000 (T1A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
	Sodium pentachloro- phenate	Erisymba buccata (EB) Notropis	BSA	-	(O)	acei	Survival time in minutes for each species at 5.0 ppm was: EB – 23 minutes	Goodnight (1942)
	<b>P</b>	umbratilis (NU) Pimephales					NU – 16 minutes	
		notatus (PN) Campostoma					PN – 42 minutes	
		anomalum Notropis					CA – 13 minutes	
		whipplii (NW) Semotilus					NW- 15 minutes	
		atromaculatus (SA) Fundulus					SA 30 minutes	
		notatus (FN) Lepomis					FN – 90 minutes	
CHEN		<i>humilis (LH)</i> Tadpole					LH — 25 minutes Tadpole — 75 minutes Crayfish, amphipods, cladocera, dragon fly nymphs, damsel fly nymphs and isopods all survived 5.0 ppm, but this concentration killed blocdworms.	
MICALS AND N	Sodium pentachloro- phenate (88 percent)	Lymnaeid snails	BSA	_	(0)	_	Each test container (500-ml beaker) was filled with ditch water. 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)
=								

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium pentachloro- phenate Sodium	Cylindrospermum licheniforme (CI) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl $-T$ (3) Ma $-T$ (3) So $-PT$ (7) Cv $-NT$ Gp $-PT$ (7) Np $-T$ (3)	Palmer and Maloney (1955)
Sodium pentachloro- phenate	Lebistes reticulatus	BSA	-	2 (K 94%- 1440 min) 4 (K 100%- 300 min) 8 (K 100%- 90 min) 15 (K 100%- 40 min) 25 (K 100%- 25 min)	_	Standard curves are developed for use in determining concen- trations for molluscicidal use in field conditions.	Klock (1956)
Sodium pentachloro- phenate	Pimephales promelas	BSA	-	0.32-0.35 (T1A)	acdf	Temperature and pH were studied as variables. The lower the pH, the more toxic the chemical was to the fish. As temperature was increased the toxicity rose proportionately.	Crandall and Goodnight (1959)
Sodium pentachloro- phenolate	Channel catfish (fingerlings)	BSA	-	0.46 (K1A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Sodium pentachloro- phenate	Lebistes reticulatus	BSCH		0.5 (44.6% К 90)	<u>a</u> cde	Sublethal effects found were retarded growth.	Crandall and Goodnight (1962)
Sodium pentachloro- phenate	Oncorhynchus kisutch	BSA	-	3.0 (O)	<u>a e</u> 	The value reported is obtained by a complex mathematical treatment and is for "median resistance times" of juvenile salmon with varying levels of salinity, temperature, and dissolved oxygen. At 3.0 mg/l pentachlorophenate, the maximum response (toxicity) was calculated to be 17.68% salt concentration, 4.86 c, and 7.66 mg/l of dissolved oxygen.	Alderdice (1963)
Sodium pentachloro- phenate	Tubificid worms	BSA	-	0.31 (T1A)	<u>a c</u>	Knop's solution was used. TL <sub>m</sub> levels for various pH's were determined. This compound was more toxic at the lower pH levels studied.	Whitley (1968)

Sodium pentachloro- phenate plus sodium salts of other phenols	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI = PT Ma = PT (14), NT (21) So = PT (14), NT (21) Cv = NT Gp = PT (7) Np = T (3)	Palmer and Maloney (1955)
Sodium perborate	Daphnia magna	BSA	-	<5.2 (0)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium p- phenol sulfonate	Daphnia magna	BSA	_	5,623 (K)	<u>a</u> c	Assay water was not characterized chemically or otherwise described. The pH at 100 percent toxicity was 6.7.	Freeman (1953)
Sodium p- phenol sulfonate	Daphnia magna Lepomis macrochirus Lymnaea sp (eggs)	BSA	-	1,471 (T4A) 19,616 (T4A) 8,828 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium phosphate	Polycelis nigra	BSA	-	0.026M (L2)	c	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Jones (1941)
Sodium phosphate	Gambusia affinis	BSA	-	720 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium phosphate	Daphnia magna	BSA	-	237 (T1A) 177 (T2A) 126 (T3A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium picrate	Phoxinus phoxinus	BSA	_	2000 ppm (192 min) 1000 ppm (369 min) 200 ppm (1563 min)	<u>ace</u> f	Tap or distilled water used as diluent. Toxicity defined as the average time when the fish lost equilibrium when exposed to the test chemical (ppm picrate).	Grindley (1946)
Sodium propionate	Culex sp (larvae) Lepomis macrochirus	BSA	-	2,320 (T2A) 5,000 (T1A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium pyrophosphate	Gambusia affinis	BSA	-	1,380 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

A-143

CHEMICALS AND MIXTURES OF CHEMICALS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium pyrophosphate	Daphnia magna	BSA	_	433 (T1A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium salicylate	Daphnia magna	BSA	_	1, <b>450</b> (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium silicate	Daphnia magna	BSA	_	2.47 (O)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium silicate	Gambusia affinis	BSA	_	2,400 (T2A)	<u>a</u> c d e g	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium silicate	Amphipoda Daphnia magna Lymnaea sp	BSA	-	895 (T1A) 263 (T2A) 160 (T4A) 247 (T4A) 630 (T1-4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
	(eggs)			• • •			
Sodium silicate- Sodium sulfate	Daphnia magna	BSA	-	158 (O) 2,899 (O)	a c	Standard reference water used. Toxicity threshold is defined as that concentration which immobilizes 50 percent in a 100-hr exposure period.	Freeman and Fowler (1953)
Sodium stearate	Pimephales promelas (juveniles)	BSA	-	(S) 200 (T1-4A) (H) 1,800 (T1-4A)	acdf	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard water (H). Pure compound was less toxic than packaged soap products.	Henderson, et a (1959)
Sodium sulfate	Polycelis nigra	BSA	-	0.048M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Jones (1941)
Sodium sulfate	Daphnia magna	BSA	-	7,105 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
Sodium sulfate	Daphnia magna	BSA	-	5,960 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. This salt may be innocuous until the concentration exerts an unfavorable osmotic effect.	Anderson (1946)

Sodium sulphate	Oncorhyncus kisutch Salmo clarkii	BSA	-	16,500 (K5A) 6,700 (K5A)	<u>a</u> de	This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et al (1952)
Sodium sulfate	clarkii Lepomis macrochirus	BSA	-	13,500 (T4A)	a d e f	This paper reports the LD <sub>50</sub> in 96 hours for 8 common inorganic salts. A synthetic dilution water of controlled hardness was prepared for use in the experiments. Among other variables, specific conductivity, as mhos at 20 C, was measured.	Trama (1954)
Sodium sulfate	Lepomis macrochirus	BCFA	-	12,500 (T4A)	acef	Test water was composed of distilled water with CP grade chemicals and was aerated.	Cairns and Scheier (1955)
Sodium sulfate	Daphnia magna	BSA	-	5,514 (O)	ac	The primary aim of this study was to determine the effects of lowered dissolved oxygen concentration upon an aquatic invertebrate when exposed to solutions of inorganic salts known to be present in various industrial effluents. Analysis of data conclusively shows the <i>D. magna</i> tested under lowered oxygen tension exhibited lower threshold values for the chemicals studied than when tested at atmospheric dissolved oxygen.	Fairchild (1955)
Sodium sulfate	Gambusia affinis	BSA	-	17,500 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium sulfate	Lepomis macrochirus	BSA	-	12,500 (T4A)	<u>acdei</u>	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, approximately 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Sodium sulfate	Pimephales promelas (juveniles)	BSA	-	(S) 9,000- 13,000 (T1-4A) (H) 13,500- 14,000 (T1-4A)	<u>acdf</u>	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard water (H). The surfactant rather than the builder contained the toxicant.	Henderson, et al (1960)
Sodium sulphate	Biomorpholaria a. alexandrina Bulinus truncatus Lymnaea caillaudi	BSA	-	4,800 (K1A) 900 (K1A) 1,000 (K1A)	а	The degree of tolerance for vector snails of biharziasis to chemicals is somewhat dependent upon temperature. The temperatures at which (K1A) occurred was 27 C for <i>Bulinus</i> and <i>Lymnaea</i> and 26 C for <i>Biomorpholaria</i> .	Gohar and El-Gindy (1961)
Sodium sulfate	Hydropsychidae Stenonema ares S. heterotarsale	BCFA	-	320 (K 15%-4 da) 320 (K 50%-4 da) 320 (K 30%-4 da)	a c d e	Soft water used as diluent. Additional data are presented.	Surber and Thatcher (1963)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium sulfate	Amphipoda Culex sp (larvae) Daphnia magna (adult) Daphnia magna (young) Lepomis macrochirus Lymnaea sp (eggs) Mollienesia latopinna	BSA		2,380 (T1A) 1,110 (T2A) 880 (T4A) 11,430 (T1A) 4,547 (T4A) 6,800 (T1A) 17,500 (T1A) 5,401 (T1A) 3,553 (T4A) 20,000 (T1A) 15,996 (T2A)	a c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium sulfate	Nitzschia linearis Lepomis macrochirus	BSA	_	1,900 (T5A) 13,500 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)
Sodium sulfhydrate	Oncorhyncus tshawytscha Oncorhyncus kisutch Salmo clarkii clarkii	BSA	-	3.3 (K5) 3.5 (K5) 1.8 (K5)	<u>a</u> d e	This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et al (1952)
Sodium sulfide	Daphnia magna	BSA	_	9.4 (O)	_	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
Sodium sulfide	Gasterosteus aculeatus	BSA	-	(O)	c e	Tap water was used to make up the solutions, which made up a pH of 6.8 with sulfuric acid. At a concentration of 0.0007N, the fish displayed much distress. At 0.00008N, the animal showed very little reaction. The test animal survived 72 hours in a solution of 0.0003N.	Jones (1948)
Sodium sulphide	Oncorhyncus tshawytscha Oncorhyncus kisutch Salmo clarkii clarkii	BSA	-	3.5 (K5) 3.1 (K5) 3.0 (K5)	a d e	This chemical is one of a number that may be found in Kraft mill waste effluents. Data are expressed as minimum lethal concentration for 5 days.	Haydu, et al (1952)
Sodium sulfide	Gambusia affinis	BSA	-	750 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)

3	Sodium sulfide	Rasbora heteromorpha	BSA	-	3.0 (O)	_	For many toxins the rate of mortality is found to be a linear function of the logarithm of the concentration of the poison; whereas the comparable relation between the logarithms of the survival time and the concentration is nonlinear. The linear function can be exploited to provide comparatively simple methods of estimating long-term survival concentrations. An application of this is suggested for defining realistic standards of toxicity. At the concen- tration listed for the chemical, the mean survival time was was 173 minutes.	Abram (1964)
	Sodium sulfide	Daphnia magna	BSA	-	16 (T1A) 13 (T2A) 9 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
	Sodium sulfite	Polycelis nigra	BSA	-	0.048M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.8. Solutions were renewed every 12 hours.	Jones (1941)
A-147	Sodium sulfite	Daphnia magna	BSA	-	3,784 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
7	Sodìum sulfite	Daphnia magna	BSA	-	440 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
	Sodium sulfite	Gambusia affinis	BSA	-	2,600 (T2A)	acdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
CHE	Sodium sulfite	Daphnia magna	BSA	-	299 (T1A) 273 (T2A) 203 (T4A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
CHEMICALS AND MIXTURES	Sodium tartrate	Polycelis nigra	BSA	-	0.065M (L2)	C	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 7.4 Solutions were renewed every 12 hours.	Jones (1941)
	Sodium tartrate	Daphnia magna	BSA	~	<3,500 (O)	~	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)
ទួ	Sodium thiocyanate	Polycelis nigra	BSA	-	0.012M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 9.6. Solutions were renewed every 12 hours.	Jones (1941)
CHEMICALS	Sodium thiocyanate	Daphnia magna	BSA	-	<11.3 (0)		This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water.	Anderson (1946)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sodium thiosulfate	Polycelis nigra	BSA	-	0.053M (L2)	с	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.4. Solutions were renewed every 12 hours.	Jones (1941)
D n	Daphnia magna	BSA	_	<520 (O)		This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Value may be only half of that reported.	Anderson (1946)
Sodium thiosulfate Sodium	Gambusia affinis	BSA	-	26,000 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
n Sodium thiosulfate	Daphnia magna	BSA	-	2,245 (T1A) 1,223 (T2A) 805 (T4A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Sodium dibasic phosphate	Daphnia magna	BSA	_	<59 (O)	-	This assay is based on concentration of the chemical required to immobilize the test animal. Assays were conducted in centrifuged Lake Erie water. Value may be only half of that reported.	Anderson (1946)
Sodium tribasic phosphate	Daphnia magna	BSA	-	<52 (0)	_	Comment same as above. Toxic effect may be the result of the precipitate formed which may obstruct the straining mechanism of the <i>Daphnia</i> .	Anderson (1946)
Sodium triphosphate	Gambusia affinis	BSA		467 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Sodium tripoly- phosphate	Pimephales promelas (juveniles)	BSA	-	(S) 400 (T1-4A) (H) 1,300- 1,350 (T1-4A)	<u>acdf</u>	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard ware (H). The surfactant rather than the builder contained the toxicant. Additional data are given.	Henderson, et al (1959)
Sodium valerate	Lepomís macrochirus	BSA	-	5,000 (T1A)	ac	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
Stannic chloride	Daphnia magna	BSA	-	146 (O)	a	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hours.	Anderson (1948)
Stannous chloride	Daphnia magna	BSA	-	<25 (0)	<u>a</u>	Comment same as above.	Anderson (1948)
Strontium chloride	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography.	Powers (1918)

work, and lists an extensive bibliography. In 0.237N solution, fish survived 168 minutes. APPENDIX A

۰.

-

Strontium chloride	Daphnia magna	BSA	-	114 (O)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hours.	Anderson (1948)
Strontium nitrate	Carassius carassius	BSA	-	(0)	<u>a</u>	This old, lengthy paper discusses toxicity of many chemicals, possible mechanism of action of some, the effect of tem- perature, effect of dissolved oxygen, the efficiency of the goldfish as a test animal, compares this work with earlier work, and lists an extensive bibliography. In 0.165N solution, fish survived 300 minutes.	Powers (1918)
Strontium nitrate	Gasterosteus aculeatus	BSA	-	0.2 (K10)	-	Solutions were made up in tap water, 3.0 to 5.0-cm stickle- back fish were used as experimental animals. This paper points out that there is a marked relationship between the toxicity of the metals and their solution pressures. Those with low solution pressures were the most toxic.	Jones (193 <del>9</del> )
Styrene	Pimephales promelas	BSA	-	51 (T4A)	acdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made	Pickering and Henderson
	Lepomis macrochirus			22 (T4A)		to estimate 100 percent survival.	(1966)
	Carassius auratus			68 (T4A)			
	Lebistes reticulatus			68 (T4A)			
4-styryl- pyridine	Petromyzon marinus (larvae)	BSA	-	10 (NTE)	<u>a</u>	<ul> <li>Additional data are presented.</li> </ul>	Piavis (1962)
Sulfide	Polycelis nigra	BSA	-	0.00045M (L2)	С	This is part of a report listing 27 anions and their toxicities on a planarian. Mode of action of the anions is discussed. Water distilled in glass was used to prepare the solutions. The pH of this solution was 6.6. Solutions were renewed every 12 hours.	Jones (1941)
Sulfoxide	Pimephales promelas	BSA	-	0.74 (T4A)	acdfg	Test water was spring water diluted with distilled water. Removal of toxic chemicals by carbon adsorption, chlorine and chlorine dioxide treatment, and alum coagulation was studied. The most effective method to remove fish poisons was by use of activated charcoal adsorption.	Cohen, et al (1961)
Sulfur	Gambusia affinis	BSA	-	10,000 (T2A)	acdeg _	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al
Sulfur (colloidal)	Carassius carassius	BSA	_	(O)	<u>a</u>	Sulfur concentrations were toxic from 0.016 to 0.210 per- cent. Survival time is reported in minutes, from 45 to 315.	Harukawa (1922-23)
Sulfur,	Simocephalus serrulatus	BSA	-	11.0 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hours.	Sanders and Cope
XT Sulfur, Lime Da So Op	Daphnia pulex			10.0 (SB)		Data cited are for 60 F, but assays were performed at varied temperatures. "Water Chemistry" (unspecified) was "controlled" during the assay period.	(1966)

-	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
	Sulfuric acıd	Daphnia magna	BSA	-	88 (O)	a c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Sulfuric acid	Gambusia affinis	BSA	-	42 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
-	Sulfuric acid	Sewage organisms	BOD	-	58 (TC <sub>50</sub> )	<u>a</u>	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	Fannic acid	Daphnia magna	BSA	_	<26 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)
	Fannic acid	Gambusia affinis	BSA	-	41 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
	Fannic acid	Sewage organisms	BOD	-	(NTE)	_	The purpose of this paper was to devise a toxicity index for industrial wastes. Results are recorded as the toxic concentration producing 50 percent inhibition ( $TC_{50}$ ) of oxygen utilization as compared to controls. Five toxigrams depicting the effect of the chemicals on BOD were devised and each chemical classified.	Hermann (1959)
	artaric acid	Daphnia magna	BSA	-	135 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to immobilize the animals under prolonged (theoretically infinite) exposure.	Anderson (1944)

; (	erpine alcohol (85 percent bine oil)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with numbers of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI - T (3) Ma - PT (14) So - NT Cv - T (3) Gp - T (3) Np - NT	Palmer and Maloney (1955)
	-tertiary-butyl- 4,6 dinitro- phenol	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Comment same as above except CI – NT Ma – NT So – NT Cv – PT (7) Gp – NT Np – PT	Palmer and Maloney (1955)
	2,3,4-tetra- chlorobenzene	Australorbis glabratus	BSA and FL	Puerto Rico	Failed	c	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
	etrachloro- nydroquinone	Microcystis aeruginosa	L		100 (K)	a, etc	The chemical was tested on a 5-day-old algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume, Chu No. 10 medium was used.	Fitzgerald, et al (1952)
<b>5</b> c	etramethyl-p- ohenylene- diamine nydrochloride	Microcystis aeruginosa	L	-	100 (K)	a, etc	Comment same as above.	Fitzgerald, et al (1952)
<b>`</b>	,3',4'-5-tetra- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10 (K2) (O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 10 ppm the chemical was not toxic to goldfish.	Walker, et al (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Thiocarbamide Titanium sulfate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI - NT Ma - PT (14) So - PT (14) Cv - PT (14) Gp - PT (7) Np - PT (7)	Palmer and Maloney (1955)
Titanium sulfate	Pimephales promelas	BSA	-	(H) 120 (T4A) (S) 8.2 (T4A)	acdf	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
Toluene	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	44 (T4A) 24 (T4A) 62 (T4A) 66 (T4A)	<u>a</u> cdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
Toluene	Gambusia affinis	BSA		1,260 (T2A)	<u>a</u> cdeg	The effect of turbidity on the toxicity of the chemicals was studied. Test water was from a farm pond with "high" turbidity. Additional data are presented.	Wallen, et al (1957)
Tribromo- phenol	Bacteria (sewage)	BSA	-	97 (O)	e	In the halophenols, the ortho was less toxic than the meta or para. All of the monohalophenols were less toxic than the 2,4,6-trihalophenols. Some data on biodegradability of halophenols were presented. The figure reported is for a TL <sub>m</sub> value for cumulative gas production for 7 days.	Ingols and Gaffney (1956)
Tri-n- butylamine	Semotilus atromaculatus	BSA	-	20 to 40 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hours and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
Tri-n- butyltin acetate	Australorbis glabratus	BSA and FL	Puerto Rico	Failed	С	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
bis-(tri-n- butyltin) oxide	Australorbis glabratus	BSA and FL	Puerto Rico	0.41-0.84 (L)	с	Comment same as above.	Seiffer and Schoof (1967)

1,2,4-trichloro- benzene	Elodea canadensis Potamogeton nodosus Potamogeton pectinatus	BSA	_	5 (O) 100 (O) 5 (O) 100 (O) 5 (O) 100 (O)	a	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. No toxic effect. Injury rating of 9.5. No toxic effect. Injury rating of 9.8. No toxic effect. Injury rating of 9.8.	Frank, et al (1961)
1,1,1-trichloro- ethane	Lagodon rhomboides	BSA	-	75-100 (O)	-	Experiments were conducted in aerated salt water. Toxicity range given as the concentrations which produced $<1/2$ deaths and $>1/2$ deaths.	Garrett (1957)
1,1,2-trichloro- ethane	Lagodon rhomboides	BSA	-	150-175 (O)	-	Comment same as above.	Garrett (1957)
Trichloro- phenol	Bacteria (sewage)	BSA	-	60 (O)	e	In the halophenols, the ortho was less toxic than the meta or para. All of the monohalophenols were less toxic than the 2,4,6-trihalophenols. Some data on biodegradability of halophenols were presented. The figure reported is for a TLM value for cumulative gas production for 7 days.	Ingols and Gaffney (1965)
Trichloro- toluene	Elodea canadensis Potamogeton nodosus Potamogeton pectinatus	BSA	-	5 (K 4 wk) 100 (K 4 wk) 5 (O) 100 (O) 5 (O) 100 (O)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. Injury rating of 8.4. Injury rating of 9.1. Injury rating of 8.5. Injury rating of 9.5.	Frank, et al (1961)
3,4,6-trichloro- 2-nitrophenol (free phenol)	Petromyzon marinus Salmo gairdnerii S. trutta	BSA	_	5 (K 100%) 17 (K 10%) 15 (K 10%)	<u>a</u>	Mortality occurred in approximately 24 hours. This was a study on controlling sea lamprey larvae.	Ball (1966)
3,4,6-trichloro- 2-nitrophenol	Petromyzon marinus (prolarvae) (larvae)	BSA	_	10 (K15) 10 (K1)	<u>a</u>	Additional data are presented.	Piavis (1962)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
3,4,6-trichloro- 2-nitrophenol (Na salt)	Petromyzon marinus (larvae)	BSA FS	_ Mich.	13 (K 100%) 12 (K 100%)	<u>a</u>	Mortality occurred in approximately 24 hours. This was a study on controlling sea lamprey larvae. The "field" study involved use of simulated lake water in large raceways.	Applegate (1958)
	Salmo gairdnerii (fingerlings)	BSA FS		23 (K 10%) 40 (NTE)		involved use of simulated have watch in harge faceways.	
	S. trutta	FS		40 (NTE)			
	Salvelinus fontinalis	FS		40 (NTE)			
	Ambloplites rupestris	FS		40 (NTE)			
	Lepomis gibbosus	FS		40 (NTE)			
	Coesius plumbeus	FS		40 (NTE)			
	Semotilus atromaculatus	FS		40 (NTE)			
	Percina caprodes	FS		40 (NTE)			
	<i>Cambarus</i> spp Aquatic Iarvae	FS FS		40 (NTE) 40 (NTE)			
	Catostomus commersoni	FS		32 (NTE)			
	lctalurus melas	FS		00 (14)			
	l. nebulosus Perca flavescens	FS FS		20 (K) 32 (NTE)			
2',4',6'-tri- chloro-3- nitrosali-	Salmo gairdnerii Carassius	BSA	_	10 (K 3 hr)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of	Walker, et al (1966)
cylanilide	auratus			10 (K2)		salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule.	)
Triethylamine	Semotil <b>us</b> atromaculatus	BSA	-	50 to 80 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hours and above which all test fish died. Additional data are presented.	Gillette, et al (1952)

	3-trifluoro- methyl-4-	Petromyzon marinus	BSA	-	2 (K 100%)	<u>a</u>	Mortality occurred in approximately 24 hours. This was a study on controlling sea lamprey larvae.	Applegate (1958)
	nitrophenol	Salmo gairdnerii			7 (K 10%)			
	3-trifluoro- methyl-4- nitrophenol	Petromyzon marinus (larvae)	BSA	-	10 (K 1-2 hr)	<u>a</u>	Additional data are presented.	Piavis (1962)
	a,a,a-trifluoro- 4-nitro-m-	Salmo gairdnerii	BSA	-	9 (K 10%)	<u>a</u>	Mortality occurred in approximately 24 hours.	Applegate (1958)
	cresol	S, trutta	BSA		7 (K 19%)	<u>a</u>		
	Triiodophenol	Bacteria (sewage)	BSA	-	83 (O)	e	In the halophenols, the ortho was less toxic than the meta or para. All of the monohalophenols were less toxic than the 2,4,6-trihalophenols. Some data on biodegradability of halophenols were presented. The figure reported is for a TL <sub>M</sub> value for cumulative gas production for 7 days.	Ingols and Gaffney (1965)
	2,2,4-trimethyl (β-phenyliso- propyl)-1,2- dihydro- quinoline	Daphnia magna	BSA	-	1.8 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on systemic administration.	Sollman (1949)
-	Trimethyl and trimethyl- octadecadienyl ammonium chlorides	Cylindrospermum licheniforme (CI) Gleocapsa sp (G) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = NT G = NT So = PT Cv = NT Gp = NT Np = NT	Palmer and Maloney (1955)
LEMICALS AND MICTUDES	3,3',5-tri- nitrobenz- anilide	Salmo gairdnerii Carassius auratus	BSA	_	10 (K 3 hr) (O)	<u>a</u>	This paper deals with the relations between chemical struc- tures of salicylanilides and benzanilides and their toxicity to rainbow trout and goldfish. The chemical structure of salicylanilides and benzanilides was related to toxicity and selectivity to rainbow trout and goldfish. Salicylanilides were more toxic than benzanilides to the fishes. The ortho hydroxy substitution of salicylanilide accelerated biological activity against fish. Meta nitro substitution on the salicyl- anilides and benzanilides increased toxicity to fish. Similar findings are reported for halogens and their relative position(s) in the molecule. At 10 ppm the chemical was not toxic to goldfish.	Walker, et al (1966)
	Trinitro- toluene	Microcystis aeruginosa	L		100 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75-ml total volume. Chu No. 10 medium was used.	Fitzgerald, et a (1952)
	2,3,5-tri- phenyltetra- zolium chloride	Microcystis aeruginosa	L	-	2.5 (K)	<u>a,</u> etc	Comment same as above.	Fitzgerald, et a (1952)

CHEMICALS AND MIXTURES OF CHEMICALS

A-155

APPENDIX A

al

al

CHEMICALS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AND MIXTI	Triphenyltin acetate	Australorbis glabratus	BSA	Puerto Rico	(O)	с	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
MIXTURES (	Triphenyltin chloride	Australorbis glabratus	BSA	Puerto Rico	(O)	c	Comment same as above.	Seiffer and Schoof (1967)
OF CHEMICALS	Tri-n-pro- pylamine	Semotilus atromaculatus	BSA	_	30 to 70 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentra- tion in ppm below which the 4 test fish lived for 24 hours and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
0	Tri-n-propyltin oxide	Australorbis glabratus	BSA	Puerto Rico	(0)	С	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
	Trypaflavine (acriflavine neutral)	lctalurus punctatus	BSA	-	17.9 (K2) 11.5 (T2A)	acfi	The experiment was conducted at 66 C.	Clemens and Sneed (1958)
<b>A-15</b> 6	Uranyl acetate	Pimephales promelas	BSA	-	(S) 3.7 (T4A)	acdf	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
	Uranyl nitrate	Pimephales promelas	BSA	-	(S) 3.1 (T4A)	acdf	Comment same as above.	Tarzwell and Henderson (1960)
	Uranyl sulfate	Pimephales promelas	BSA	-	(H) 135 (T4A) (S) 2.8 (T4A)	a c d f	Comment same as above.	Tarzwell and Henderson (1960)
	Urea	Semotilus atromaculatus	BSA	-	16,000 to 30,000	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentration in ppm below which the 4 test fish lived for 24 hours and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
	Valeric acid	Daphnia magna	BSA	-	45 (T2A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evalua- tions were made in various types of water.	Dowden and Bennett (1965)
	Vanadium pentoxide	Pimephales promelas	BSA	-	(H) 55 (T4A) (S) 13 (T4A)	acd f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
	Vanadyi sulfate	Pimephales promelas Lepomis macrochirus	BSA	-	(H) 30 (T4A) (S) 4.8 (T4A) (H) 55 (T4A) (S) 6 (T4A)	acdf	Comment same as above.	Tarzwell and Henderson (1960)

	Vanillin	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): Ci - NT Ma - NT So - PT (3) Cv - PT (3) Gp - T (3), PT (21) Np - PT (7)	Palmer and Maloney (1955)
	Vinyl acetate	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	22 (T4A) 18 (T4A) 42 (T4A) 26 (T4A)	<u>a</u> cdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
A-157	Xanthic acid, ethyl sodium salt	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI - NT Ma - PT (7) So - NT Cv - PT (7) Gp - NT Np - NT	Palmer and Maloney (1955)
오	Xylene	Daphnia magna	BSA	_	100 1000 (T1A)	<u>a</u> c	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water.	Dowden and Bennett (1965)
CHEMICALS AND MIXTURES	Xylene	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	21 (T4A) 22 (T4A) 24 (T4A) 39 (T4A)	<u>a</u> cdef	Most fish survived at test concentrations of about one half, or slightly more, of the TL <sub>m</sub> value. No attempt was made to estimate 100 percent survival.	Pickering and Henderson (1966)
TURES OF	Zinc	Salmo gairdnerii	BSA	_	6 (K2)	a e	The concentration given was fatal to fingerlings. Young fish 2 and 4 weeks old could not tolerate concentrations of 4 ppm, but with increasing age showed a tendency to develop a tolerance to solutions of this concentration.	Goodman (1951)
- CHEMICALS	Zinc	Sewage organisms	BOD	-	(0)	-	Zinc was toxic to sewage organisms in concentrations as low as 0.001 ppm. This could result in errors in BOD tests. At 1.0 ppm, the oxygen demand in percent of the control was 83%.	Ingols (1956)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Zınc	Lepomis macrochirus (adult)	BSA	_	18 C (H) 10.1- 12.5 (T4A) (S) 2.9- 3.8 (T4A) 30 C (H) 10.2- 12.2 (T4A) (S) 1.9- 3.6 (T4A)	<u>a</u> cdefing	The results of these experiments indicated that in dilution water of the same quality there was little difference in toxicity at 18 C and 30 C. A considerable difference in toxicity was apparent between hard (H) and soft (S) water. A greater amount of zinc in solution was required in hard water than in soft water. Hardness of the dilution water had a greater effect upon the toxicity of zinc than did temperature.	Cairns and Scheier (1958)
Zinc	Salmo salar	BCFA	-	0.042 (T1A)	a c f	The laboratory water in which the experiment was per- formed contained 3 $\mu$ l/liter of zinc, as judged by analysis over several years, and 2 $\mu$ g/liter of copper. Lethal concen- trations of mixtures acted two or three times as fast as the metals singly, a somewhat greater potentiation than was found in the previous tests with salmon.	Schoenthal (1963)
Zinc	Lepomis macrochirus Lepisosteus Oorosoma petenense Dorosoma cepedianum Alosa chrysochloris Cyprinus carpio Carassius auratus	BCFA	-	0.0-5.0 (O)	a c f	An autopsy method for acute zinc toxicity in fish was devel- oped. Thirty to 90-day exposures to sublethal concentra- tions indicated that the opercular bone accumulates zinc at the same rate as gill tissue. By using the ratio of zinc in the gill to zinc in the bone a reasonably constant value was obtained by nonlethal exposures. This value increased up to a hundredfold in acute exposures.	Mount (1964)
Zinc	Lepomis macrochirus	BSA	-	2.86-3.78 (O) 0.90-2.10 (O) 6.60-9.47 (O) 6.18-9.50 (O)	a f	At the given concentration 50% survival occurred at 18 C in soft water. At the given concentration 50% survival occurred at 30 C in soft water. At the given concentration 50% survival occurred at 18 C in hard water. At the given concentration 50% survival occurred at 30 C in hard water.	Cairns (1965)
Zinc	Rainbow trout	FR	Scotland	3.9 (T2)	acefim	This work represents an extension of laboratory studies of the toxicity of complex effluents to investigations of rivers.	Herbert, et al (1965)

	Zinc	Pimephales promelas	BSA	_	(O)	a c d	Zinc sulfate was added to tap water for the experiments. $TL_m$ values for minnow eggs: 1 day - 3.95 ppm 2 day - 2.55 ppm 4 day - 1.83 ppm 7 day - 1.71 ppm 12 day - 1.63 ppm $TL_m$ values for minnow fry: 1 day - 0.95 ppm 2 day - 0.95 ppm 4 day - 0.87 ppm 7 day - 0.87 ppm From the experimental data, it appeared that animals ex- posed to a dilute zinc solution developed a tolerance to this metal. The duration of the tolerance was not investigated.	Pickering and Vigor (1965)
	Zinc	Fathead minnow	BCFA	_	4.9 to 32.3 (T4A)	a c d e	Zinc was most toxic at a pH of 8.0 and a water hardness of 50 ppm and least toxic at pH 6.0 and a hardness of 200 ppm. At any given hardness, zinc was always more toxic at a high pH than at a low pH. The results are in disagreement with most published work possibly because a flow-through system would keep any precipitated zinc in suspension. The first value reported is for a pH of 8.0 and a hardness of 50, and the second for a pH of 6.0 and a hardness of 200.	Mount (1966)
1	Zinc	Fundulus heteroclitus	BSA	-	157-180 (K)	асеі	Fish subjected to the concentration reported died in 24 to 48 hours. The dead fish contained 7 and 8 times more zinc in the whole fish and in the gill arch than untreated control fish.	Eisler (1967)
CHEMICALS AND	Zinc	Lebistes reticulatus Bufo valliceps (tadpoles) Daphnia magna	L	-	10.0 (К) 10.0 (К) 1.0 (К)	a c e	It is assumed in this experiment that the cations considered are toxic because they combine with an essential sulfhydryl group attached to a key enzyme. This treatment indicates that the metals which form the most insoluble sulfides are the most toxic. The log of the concentration of the metal ion is plotted against the log of the solubility product con- stant of the metal sulfide — a treatment that does not lend itself to tabulation. The cation toxicity cited is only an approximate concentration interpolated from a graph. Time of death was not specified.	Shaw and Grushkin (1967)
LS AND M	Zinc	Gasterosteus aculeatus	BSA	_	0.1 (O)	ace	This is a discussion of a bioassay method using stickleback fish and spectrophotometric determinations of the chem- icals evaluated. The number listed is said to be the "toxic limit" for the fish.	Hawksley (1967)
MIXTURES OF CHEI	Zinc	<i>Lebistes reticulatus</i> (guppy)	BSA	_	(O)	acfno	A series of equations was devised to describe the toxicity of a system containing two toxicants — zinc and cyanide. A concentration of 1 ppm of Zn killed 50% of the fish in 32 hours. 0.75 ppm killed 50% in 63 hours, and 0.56 ppm killed 50% in 96 hours. Toxicity of the two-component system was then determined using varying ratios of the two components.	Chen and Selleck (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Zinc	Lepomis macrochirus	BCFA	_	7.2-12.0 (T20CF)	<u>a c</u> d e f	The toxicity of zinc was largely dependent upon the dis- solved oxygen in the water. Bluegills showed an increased mortality to zinc as a result of an environmental stress of low dissolved oxygen concentration. The lowest toxic zinc concentration was for a system containing 1.8 mg/l of dissolved oxygen, and the highest for a system containing 5.6 mg/l.	Pickering (1968)
Zinc	Salmo gairdnerii	BSA	_	2.8-3.5 (T4A)	acdefo	The concentration killing a half batch of fish in 2 days pro- vides a reasonable estimate of the threshold concentration. The lethality of this chemical depends upon the total hard- ness of the water and the dissolved oxygen concentration.	Brown (1968)
Zinc	Oncorhynchus kisutch	BSA	-	(O)	ac	Zinc uptake and distribution in the developing coho salmon egg was measured using radioisotope tracer techniques. About 70% of the total accumulated zinc was bound rather firmly to the chorion, 26% was found in the perivitelline fluid, 2% in the yolk, and 1% in the embryo.	Wedemeyer (1968)
Zinc (Zn <sup>++</sup> )	Lepomis macrochirus	BSA	-	4.2 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
Zinc and copper	Atlantic salmon	FR	Canada	(O)	f	"Toxicity index" for zinc and copper combined was de- scribed in connection with disturbed salmon migration. Toxicity index $\geq$ 1.0 indicates lethality to "young salmon after long exposure" A toxicity index of 0.15 or 15% of the lethal concentration of zinc and copper seems to be the maximum safe level for migration.	Sprague (1964)
Zinc and copper	Salmo salar	BSA	-	0.048 Cu (O) 0.600 Zn	а	The values given are for an ILL (incipient lethal level) and in this instance only in water of 20 mg/liter of hardness. Concentrations above this are lethal in about one day. These values were determined by bioassay. Salmon parr in the laboratory avoided less than one tenth of incipient lethal levels. Avoidance thresholds were 0.09 ILL of zinc, 0.05 ILL of copper and 0.02 ILL of equitoxic mixtures. In equitoxic mixtures of these compounds, the ILL was additive.	Sigler, et al (1966)
Zinc acetate	Pimephales promelas	BSA	_	(S) 0.88 (T4A)	c d e f	(S) Soft water. Values are expressed as mg/l of metal.	Pickering an Henderson (1965)
Zinc boro- fluoride	Sewage organisms	BOD	-	55 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)

Zinc chloride	Daphnia magna	BSA	-	≪0.15 (S)	<u>a</u>	Lake Erie water was used as diluent. Toxicity given as threshold concentration producing immobilization for exposure periods of 64 hours.	Anderson (1948)
Zinc chloride	Lepomis macrochirus	BCFA	-	6.91 (T4A)	a c e f	Test water was composed of distilled water with CP grade chemicals and was aerated throughout the 96-hour ex- posure period.	Cairns and Scheier (1955)
Zinc chloride	Lepomis macrochirus	BSA	-	20 (T4A)	<u>a c e</u>	Increase in temperature seemed to increase toxicity of this chemical. Low dissolved oxygen reduced toxicity of some chemicals in this study. Toxicity values may be 20% higher in hard versus soft water.	Cairns (1957)
Zinc chloride (as Zn <sup>++</sup> )	Lepomis macrochirus	BSA	-	(N) 8.02 (T4A) N (L) 4.9 (T4A) L	<u>a e</u>	Modified Chu No. 14 test medium was used. Toxicity is given both for "normal" $O_2$ (5-9 ppm), (N), and with "low" $O_2$ (2 ppm DO), (L). High and low threshold concentration percent of survival are also presented.	Cairns and Scheier (1958)
Zinc chloride	Lepomis macrochirus	BSA	-	6.91 (T4A)	<u>acdei</u>	A "control" was prepared by adding required chemicals to distilled water, and this was constantly aerated. Data reported are for larger fish, approximately 14.24 cm in length. Data for smaller fish are also in the report.	Cairns and Scheier (1959)
Zinc chloride (tagged with zinc 35)	lctalurus nebulosus	BS	_	(O)	cdfil	Bullheads showed an initial rapid uptake of zinc for the first several hours followed by a short period of decline. Z <sup>35</sup> was used to measure zinc uptake. The fish exposed to 6.0 ppm of zinc for 96 hours, when placed in flowing, fresh water, lost 43 percent of their total accumulated zinc after 1 day. Fish exposed to 12 ppm of zinc for 14 days all survived.	Joyner (1961)
Zinc chloride (as Zn <sup>++</sup> )	Brachydanio rerio (adults) (eggs) Lepomis macrochirus	BSA	-	28 (T2A) 105 (T2A) 5.2 (T2A)	<u>acdef</u>	The test dilutions were made up from distilled water and ACS grade chemicals. Temperature was held at 24 C and the solution was aerated to maintain a dissolved oxygen content of 5-9 ppm.	Cairns, et al (1964)
Zinc chloride	Lepomis macrochirus	BSA	-	(S) 5.37 (T4A)	c d e f	(S) Soft water. Values are expressed as mg/I of metal.	Pickering and Henderson (1965)
Zinc chloride	Nitzschia linearis Physa heterostropha Lepomis macrochirus	BSA	-	4.3 (T5A) 0.79-1.27 (T4A) 2.86-3.78 (T4A)	a c e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors advise that bioassays on at least 3 components of the food web be made in any situation.	Patrick, et al (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Zinc-copper- cyanide complex	Pimephales promelas	BSA	_	1.0 ppm Zn 0.025 ppm Cu 0.05 ppm CN (non-toxic 4 days) 1.0 ppm Zn 0.25 ppm Cu 0.33 ppm CN (non-toxic 4 days) 1.0 ppm Zn 0.025 ppm Cu (K ≤ 14 hr)	<u>a</u> c	Synthetic soft water used.	Doudoroff, et al (1956)
Zinc cyanide	Sewage organisms	BOD	-	0.75 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treat- ment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chemical was tested in an unbuffered system.	Sheets (1957)
Zinc cyanide complex	Lepomis macrochirus (juveniles)	BSA	-	0.4 (O)	<u>a</u> cdf <u>p</u>	For the concentration given, the median resistance time in minutes was 256.	Doudoroff, et a (1966)
Zinc (a)- cyanide (b) mixture	Lepomis macrochirus	BSA	_	(a) 3.90 (T4A) (b) 0.26 (T4A)	a c d e	All fish were acclimatized for 2 weeks in a synthetic dilution water.	Cairns and Scheier (1968)
Zinc cyanide complex {sodium cyanide (564 ppm CN <sup>-</sup> ) and zinc sulfate (394 ppm Zn)]	Pimephales promelas	BSA	-	0.18 (T4A)	<u>a</u> c	Synthetic soft water was used. Toxicity data given as number of test fish surviving after exposure at 24, 48, and 96 hr. TL <sub>m</sub> values were estimated by straight-line graphical inter- polation and given in ppm CN <sup>-</sup>	Doudoroff, et a (1956)
Zinc dimethyl- dithio- carbamate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partial toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI = PT (7) Ma = T (7), PT (14) So = NT Cv = PT (14) Sp = T (14) Np = T (3)	Palmer and ly Maloney (1955)

Zinc dimethyl- dithio- carbamate (100 percent)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Comment same as above except that: CI – T (3) Ma – T (3) So – T (3) Cv – T (3) Gp – T (3) Np – T (3)	Palmer and Maloney (1955)
Zinc ion	Physa heterostropha	BSA	-	20 C (S) 0.79- 1.27 (T4A) (H) 2.66- 5.57 (T4A) 30 C (S) 0.62- 0.78 (T4A) (H) 2.36- 6.36 (T4A)	<u>a</u> cd <u>e</u> q	The objective of these experiments was to determine the effects of water temperature and hardness on the toxicity of zinc ion to pond snails. (H) = hard water, (S) = soft water.	Cairns and Scheier (1958)
Zinc ion	Fish	BSA	_	1.5 x 10 <sup>-4</sup> M (K)	<u>a</u> c	Avoidance behavior of test fish to toxic chemicals is given. Toxicity is given as the lowest lethal concentration (molar). Ratios of avoidance and lowest lethal concentration are presented and discussed.	łshio (1965)
Zinc-nickel cyanide complex	Pimephales promelas	BSA	-	1.0 ppm CN <sup>-</sup> 0.6 ppm Zn (K <16 hr) 0.13 ppm Ni	a_c	Synthetic soft water was used. Toxicity data given as number of test fish surviving.	Doudoroff, et al (1956)
Zinc nitrate	Balanus balanoides	BSA	_	32.0 (O)	-	The concentration listed was lethal to 90% of adult barnacles in 2 days.	Clarke (1947)
Zinc salts	Diatoms Platyhelminths Many insects	FR	Ystwyth, Wales	0.2-0.7 (O)	-	Zinc salts were from mine drainage. The flora above the mines was rich, but below the sources of pollution was poor in quantity and variety of lithophilous insects.	Jones (1958)
Zinc salts	Salmo gairdnerii	BSA	-	(O)	<u>a e</u>	<ul> <li>This is a study of the effect of varying dissolved oxygen concentrations on the toxicity of selected chemicals.</li> <li>The toxicity of heavy metals, ammonia, and monohydric phenols increased as the dissolved oxygen in water was reduced. The most obvious reaction of fish to lowered oxygen content is to increase the volume of water passed over the gills, and this may increase the amount of poison reaching the surface of the gill epithelium.</li> <li>The concentration of the chemical in the water was not specified.</li> </ul>	Lloyd (1961)
Zinc stearate	Lepomis macrochirus	BSA	-	(O)	-	"Standard reference water" was described and used as well as lake water. Varied results were obtained when evaluations were made in various types of water. Compound is very slightly soluble in water. No toxicity data given.	Dowden and Bennett (1965)

CHEMICALS AND MIXTURES OF CHEMICALS

A-163

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Gasterosteus aculeatus	BSA	-	0.7 (TL4½A)	a c	Death of the fish resulted from an interaction between the metallic ion and the mucus secreted by the gills. Coagulated mucus formed on the gill membranes and impaired respira- tion to such a degree that the fish asphyxiated. The addition of 50 mg/l of calcium chloride to the tank pro- tected against the toxic effect of this metal salt.	Jones (1939)
Zinc sulfate MXTURES Zinc Sulfate CHEMICALS Zinc	Daphnia magna	BSA	_	<48 (O)	<u>a</u> c	This paper deals with the toxicity thresholds of various sub- stances found in industrial wastes as determined by the use of <i>D. magna</i> . Centrifuged Lake Erie water was used as a diluent in the bioassay. Threshold concentration was defined as the highest concentration which would just fail to im- mobilize the animals under prolonged (theoretically infinite) exposure.	Anderson {1944}
<sup>to</sup> Zinc sulfate	Salmo gairdnerii	BSA	-	25 ppm (0, 133 min)	acef	Tap or distilled water used as diluent. Toxicity defined as the average time when the fish lost equilibrium when exposed to the test chemical (ppm Zn).	Grindley (1946)
Zinc sulfate	Pygosteus pungitius	BCF	-	(0)	<u>a</u> c	Fish were exposed to 0.04, 0.003, 0.0003, and 0.0001N zinc sulfate. Survival times at these concentrations were, re- spectively: 85 minutes, 190 minutes, 7 hr, and 15 hr.	Jones (1947)
Zinc sulfate	Sewage organisms	BOD	-	920 (O)	-	Various metal salts were studied in relation to how they affected the BOD of both raw and treated sewage as well as how they affected the processing of sewage in the treatment plant. BOD was used as the parameter to measure the effect of the chemical. The chemical concentration cited is the ppm required to reduce the BOD values by 50%. This chem- ical was tested in an unbuffered system.	Sheets (1957)
Zinc sulfate	Salmo gairdnerii	BSA	_	(O)	a c	Zinc sulfate was less toxic in hard water than in soft water; more toxic in hard water with increased temperature; and more toxic when DO decreased. Survival curves are presented.	Lloyd (1960)
Zinc sulfate (as Zn)	Rainbow trout	BSA	-	(0)	a c d	The 48-hour LD <sub>50</sub> as interpolated from a graph was 4 ppm. A method for prediction of toxicity of spent liquor from a coke oven before and after biological treatment is briefly discussed.	Herbert (1961)
Zinc	Tendipes	BSA	_	56 (K 40%)	<u>a</u> cdi	Kill data for <i>T. decorus</i> is presented on other concentrations	Wurtz and
sulfate	decorus Limnodrilus			10 (T4A)		in either hard or soft water.	Bridges (1961)
	hoffmeisteri Physa			14 (T4A)			
	heterostropha Asellus			38.5 (T4A)			
	<i>communis</i> Argia sp			56 (T4A)			
Zinc sulfate	Lebistes reticulatus	BSCH	-	5.0 (41% K 90)	acde	Sublethal effects found were retarded growth, increased mortality, and delayed sexual maturity.	Crandall and Goodnight (1962)

Zinc sulfate	Physa heterostropha Helisoma companulata	BSA	_	4.2 (T1A)* 1.9 (T2A) 1.9 (T3A) 1.9 (T4A) 49.0 (T1A) 49.0 (T2A) 13.4 (T3A) 13.4 (T4A)		These tests were conducted in hard and soft water at varied temperatures. Generally, this chemical was more toxic in soft water. At temperatures up to 90 F, zinc sulfate was less toxic than at 51 F for <i>P. heterostropha</i> .	Wurtz (1962)
Zinc sulfate	Salmo salar	BSA	-	(O)	<u>acdef</u>	The EC <sub>50</sub> or the effective concentration that produced an avoidance response in 50% of the fish was 0.092 x the ILL (incipient lethal level), or 0.092 x 580 µg/I, or 53.3 µg/I.	Sprague (1964)
Zinc (zinc sulfate)	Salmo salar	BCF	-	0.6 (O)	<u>a</u> cdef	The experiments were carried out in soft water. Values are reported as micrograms of metal and toxicity as $LT_{50}$ . In solutions containing copper and zinc, fish died twice as fast as would occur if the 2 metals were simply additive in their lethal action.	Sprague (1964)
Zinc sulfate (as Zn)	Salmo gairdnerii	BSA	-	3.86 (T2A)	acdf	A mathematical equation was derived to explain the com- bined toxicities of this salt and ammonium chloride.	Herbert and Shurben (1964)
Zinc sulfate	Periphyton	FL	Newtown, Ohio	1.1-6.5 (O)	acdf	Fungi and slime-forming bacteria grew abundantly in the high Zn concentrations, apparently due to nutrient release from decaying periphyton.	Williams and Mount (1965)
Zinc sulphate	Brachy danio rerio	BSA	_	(O)	<u>a</u> cef	Survival time for adult fish (aged 40 days) in 168 hours was 10 ppm. The chemical was more toxic to newly hatched fish.	Skidmore (1965)
Zinc sulfate	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	(S) 0.96 (T4A) (H) 33.4 (T4A) (S) 5.46 (T4A) (H) 40.9 (T4A) (S) 6.44 (T4A) (H) 1.27 (T4A)	c d e f	(S) Soft water. (H) Hard water. Values are expressed as mg/l of metal.	Pickering and Henderson (1965)
Zinc sulfate	Brachydanio rerio (embryos)	BSCH	-	20 (K 15 hr)	<u>a</u> ce	Embryos with the outer membranes removed survived longer than natural embryos – the action of zinc sulfate on mem- branes is unknown. Additional data are presented.	Skidmore (1966)
Zinc sulfate	Brachy danio rerio	BSA	-	20 (K1)	<u>a</u> cde	Data are given for several concentrations of zinc. The authors also measured oxygen uptake of the fish plotting this value against the dry weight of the fish. Toxicity of zinc to fish of different ages was also measured. An equation was de- rived to express toxicity of zinc to these fish.	Skidmore (1967)
Zinc sulfate (as Zn)	Salmo salar S. trutta S. gairdnerii S. trutta S. gairdnerii	BSCH	-	0.1 (K) (O) (O) (O) (O)	c f	The reported figure is a reported lethal concentrate as found in polluted lakes and streams in Norway. Apparently organic matter has a masking effect that reduces toxicity. Rainbow trout and Atlantic salmon reacted similarly to the chemical. Brown trout was only slightly more tolerant. The value given is for a 21-day median survival period. 50% of brown trout eggs survived to hatch in 0.3 ppm Zn. Eggs of rainbow trout behaved similarly.	Grande (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Zinc sulfate	Salmo gairdnerii Perca	BSA	_	4.6 (T4A) 16.0 (T5A)	<u>c e</u>	Data given as $LC_{50}$ which was taken as equivalent to $TL_m$ . Additional data for other exposure periods are presented.	Ball (1967)
	fluviatilis Rutilus rutilus			17.3 (T5A)			
	Gobio gobio			8.4 (T7A)			
	Abramis brama			14.3 (T5A)			
inc sulphate	Salmo gairdnerii	BCFA	-	3.8-5.5 (K5)			
hydrated)	Perca fluviatilis Rutilus	fluviatilis Rutilus rutilus Gobio		14.8-17.3 (K5)	асе	Data confirmed that experiments of short duration are not necessarily reliable for ranking the ultimate sensitivities of several species of fish to a given poison.	Ball (1967)
rutilı Gobio gobic Abran				15.4-19.4 (K5) 9-15 (K5)			
	gobio Abramis brama			12.5-16.3 (K5)			
inc sulfate	Tubificid worms	BSA	_	46.0 (T1A)	<u>a</u> c	Knop's solution was used. TL <sub>m</sub> levels for various pHs were determined for the tubificids and were found to be 5.8 to 9.7. Zinc sulfate was more toxic at pH extremes of 6.5 and 8.5 than at 7.5.	Whitley (1968)
Zinc sulfate plus copper sulfate (vari- ous ratios)	Salmo gairdnerii	BSA	-	_	<u>a e p</u>	Both hard and soft water were used. Median period of sur- vival in hard water was 3 days – 3.5 ppm Zn, and 1.1 ppm Cu; in soft water 7 days, 0.56 ppm Zn and 0.044 ppm Cu.	Lloyd (1961)
nc ulfate blus alkyl- benzene ulfonate	Salmo gairdnerii	BCFCH & A	-	0.3* (T4A) *ABS + 0.8 ppm Zn	a b c d e f	For a concentration of 0.45 ppm of alkyl benzene sulfonate alone, the median tolerance limit was recorded in 4 days. The zinc concentration was 0.08 ppm in the combined zinc-detergent solution. The ABS appeared to block devel- opment of resistance to Zn in the trout in chronic studies.	Brown, et al (1968)
rconium xychloride	Pimephales promelas Lepomis macrochirus	BSA	-	(H) 240 (T4A) (S) 18 (T4A) (H) 270 (T4A) (S) 15 (T4A)	a c d f	Both hard (H) and soft (S) water were used.	Tarzwell and Henderson (1960)
rconium ulfate	Pimephales promelas	BSA	-	(H) 145 (T4A) (S) 14 (T4A)	a c d f	Comment same as above.	Tarzwell an Henderson (1960)

CHEMICALS AND MIXTURES OF CHEMICALS

## APPENDIX B

## EXTRACTED DATA FROM ORIGINAL PAPERS – COMMERCIAL CHEMICAL PRODUCTS

Note: Names of chemicals and organisms are as given by the various authors. Readers should search for alternate, common, and/or scientific names of both chemical and aquatic species; and refer to report section on Extracted Data for further discussion of this appendix.

Footnotes for Appendices A and B:

- (1) Letters represent:
  - B = bioassay, used in combination with S = static, CF = continuous flow, A = acute, and CH = chronic.
  - L = laboratory bioassay.
  - BOD = biochemical oxygen demand.
    - F = field study, used in combination with R = river, stream, creek, etc., L = lake or pond, M = marine, E = estuarine, and O = other (port facility, flooded area, etc.).
- (2) Field location is indicated by abbreviation of the state or country.
- (3) The number indicates ppm (mg/l), unless otherwise indicated by appropriate designations or (O). The letters within parentheses following indicate  $T = TL_m$ , K = kill, SB = sublethal effects, NTE = no toxic effect, or O = other. The number following these indicates the time in days at which observations were made. EC50, LC50, and similar designations for 50 percent lethality were all considered as  $TL_m$  and designated as such. The numbers within parentheses following these designations indicate the time in days when the effect was observed.
- (4) The following indicate (when underlined the variable was controlled):

```
a = water temperature
```

```
b = ambient air temperature
```

c = pH

```
d = alkalinity (total, phenolphthalein or caustic)
```

```
e = dissolved oxygen
```

```
f = hardness (total, carbonate, Mg, or CaO)
```

g = turbidity

```
h = oxidation-reduction potential
```

```
i = chloride as Cl
```

```
j = BOD, 5 day; (J) = BOD, short-term
```

```
k = COD
```

```
l = nitrogen (as NO_2 or NO_3)
```

- m = ammonia nitrogen as NH3
- n = phosphate (total, ortho-, or poly)
- o = solids (total, fixed, volatile, or suspended)

 $p = CO_2$ 

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
2389 (10%)	Chlorella pyrenoidosa	L	_	100 (AC 1/2 hr)	_	Describes a bioassay method to differentiate between an algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)
Abate	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	_	4.0 (O)	а	Stones heavily populated with larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 56 percent.	Jamnback and Frempong- Boadu (1966)
Abate	Micropterus salmoides Lepomis	BSA	_	200+ (L1A)	<u>a</u>	Abate was toxic to fish at a dosage rate necessary to control the larvae of the chironomid midge.	Von Windeguth and Patterson
(Am.Cy.52, 160)				200+ (L1A)			(1966)
	macrochirus Gambusia			200+ (L1A)			
	affinis Lebistes			200+ (L1A)			
	reticulatus Paleomonetes			1.0 (L1A)			
	paludosus Hyalella			0.65 (L1A)			
	azteca Plankton (Euglena,			50.0 (K2)			
	<i>Coleps)</i> Rotifers			50.0 (K2)			
Abate	Callinectes sapidus	BCFCH	-	0.01 (K)	а	Little or no information was given about test procedures and further results.	Butler and Johnson (1967)
Abate	Microp terus salmoides	BSA	_	5.0 (T 1 hr) 5.0 (K 2 hr)	a e	Experiments were carried out in plastic tubs lined with saran plastic. Fish weights averaged 217 grams.	Mulla, et al (1967)
Abate	Pteronarcys californica (naiads)	BSA	-	0.01 (T4A)	<u>a c d e f</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Alkyl benzene	Cladophora	BSA	_	10 (K21A)		Within the range of reduction of ABS detergent concentration	Hynes and
sulphonate (ABS)	glomerata Eurhynchium			10 (K21A)	•	which has been achieved by the Luton experiment there was very little biological effect on the river.	Roberts (1962)
	rusciforme Ranunculus pseudofluitans Potamogeton			2.5 (K14)			,
				2-3 (SB14)			
	pectinatus P. densus			2.5 (SB14)			
Sodium alkylary) sulfonate	Rainbow trout (fry)	BSA	-	3.0-5.0 (T1A)	<u>a c d e</u>	The 24-hr TL <sub>m</sub> was very near the highest concentration that was nonlethal in 6 hr. Additional data are discussed.	Vivier and Nisbet (1965)

Sodium alkyl benzene sulfonate	Lepomis macrochirus L. gibbosus	BSA and CFCH	_	17.4 (T4A) 17.4 (T4CF) 21.9 (T4A)	<u>a</u> c d e	Both hard and soft water were used. Data from both were similar. TL <sub>m</sub> for 24 and 48 hr are given. Gill damage occurred at 5-6 ppm after 3 months of exposure. Data on cruising speed and active oxygen consumption are also presented in addition to erythrocyte count and histological examination of gills. Similar gill damage for <i>L. macrochirus</i> occurred in acute and chronic studies.	Cairns and Scheier (1963)
ABS	Lepomis macrochirus	BCF	_	19.7 (T1A) 18.1 (T4A) 17.3 (T30A)	acdefq	Toxicities are recorded as an average for 3 tests. Test fish exhibited some degree of acclimation to the chemical after exposure to sublethal concentrations.	Lemke and Mount (1963)
Alkyl benzene sulfonate (25 percent)	Hydropsychidae	BCFACR	-	32 (60% K)	acde	Concentration, time and percent survival are given. Con- siderable additional data are also presented.	Surber and Thatcher (1963)
	Stenonema sp S. ares S. heterotarsale Isonychia bicolor Orconectes rusticus			16 (K) 16 (K) 16 (K) 4.0 (K) 32 (K)			
ABS	Goniobasis sp Lepomis gibbosus	BSCHA	-	32 (K) 12 (O)	<u>a</u>	Fish were exposed to the ABS solution for two weeks, and subsequently to a sublethal concentration of ZnCl <sub>2</sub> (2.4 ppm). Limited tests indicated that exposure to ABS in excess of 5.6 ppm caused marked gill damage but produced no gross changes in zinc tolerance. Other fish were exposed to ABS as above, then in dilution water alone, and the temperature was raised to 35° for 96 hours. Exposure to ABS apparently caused no changes in the tolerance of the fish to the higher temperature.	Cairns and Scheier (1964)
Alkyl benzene sulfonate (54.8%)	Nitzchia linearis Navicula seminulum	BSA		(S) 10 (T5) (S) 5.6 (T5) (H) 39.4 (T5)	ac <u>e</u>	Effects in hard (H) and soft (S) waters were compared. Two compositions of ABS were used. TL <sub>m</sub> is given in ppm of ABS composition.	Cairns, et al (1964)
2	Physa heterostropha			(S) 34.2 (T5) (H) 35.8 (T5)			
	Lepomis macrochirus and Lepomis gibbosus			5.6 - 18.0 (survived, but extensive gill damage occurred)			
Sodium alkyl benzene sulfonate	lctalurus natalis	BSCH	_	0.5 (SB1CH)	<u>a</u> cdf	At 0.5 ppm, bullhead chemoreceptor damage occurred. Detergent concentration was monitored by the methylene-blue technique.	Bennett (1962)

APPENDIX B

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
ABS	Brachydanio rerio (adults) (eggs) Lepomis macrochirus	BSA	-	42.0 (T2A) 75.0 (T2A) 17.4 (T2A)	<u>acdef</u>	The test dilutions were made up from distilled water and ACS grade chemicals. Temperature was held at 24 C and the solution was aerated to maintain a dissolved oxygen content of 5-9 ppm.	Cairns, et al (1965)
ABS (54.8% active)	Lepomis gibbosus	BSCH	-	18 (O)	<u>acdei</u>	Chloride content of the water was adjusted to 60 ppm – and the fish were exposed to the test solution for 21 days. At this time, the chloride content was raised to 6500 ppm, and the test was continued another 21 days. ABS generally damaged the gill structure. Since salt exchange as well as oxygen exchange takes place here, it would not be surprising that gill damage would correlate with chloride content of the blood. However, there was little difference in the blood chloride in control and experimental animals.	Cairns and Scheier (1966)
Alkyl benzene sulfonate	Vaucheria Cladophora	BSA		(O)	<u>a</u> filn	<ul> <li>Experiments were conducted in five 1-gal. containers. Algal communities were subsampled and the samples were placed in 60-ml bottles at 4 time periods: 12 hr, 24 hr, 48 hr, and 96 hr.</li> <li>Results showed that ABS has a negative effect on C<sup>14</sup> uptake for both algae communities, the communities appear to partly recover their ability to assimilate C<sup>14</sup> at extended exposures to high concentrations, and a slight stimulation of C<sup>14</sup> uptake appears to occur at abbreviated exposures to low concentrations.</li> </ul>	Hicks and Neuhold (1966)
ABS	Chlorella pyrenoidosa	L	-	0-20 mg/l increased growth rate	acep	Growth rates of the <i>Chlorella</i> were followed when supplied synthetic detergents as the phosphorus source. Sodium triphosphate was responsible for increased growth.	Maloney (1966)
ABS	Pimephales promelas (eggs)	BCF	-	6.4 (T9)	acdef	Mortality range is given for exposure (days 1-9) with various concentrations and controls. Additional data are presented.	Pickering (1966)
ABS (54.8%)	Jordanella floridae	BSCH	_	10 to 65 (NTE)	-	Aquaria were prepared containing 0, 10, 28, 42, 56, 65 ppm of ABS. The major effect found was on the feeding habits of the fish. Apparently the chemical made worms in the aquaria unpalatable. Time required for the consumption of the worms varied with the concentration of the chemical.	Foster, et al (1966)
ABS	Lepomis gibbosus	BSA	-	(O)	<u>a e</u>	Gill damage in pumpkinseed sunfish resulting from 24-hr exposure to 18 ppm of this chemical was not reversible, even after the test fish were removed to fresh dilution water for an eight-week period.	Scheier and Cairns (1967)
ABS	Notropis antherinoides	BCFA	-	7.4 (T4A)	acdef	Differences in sensitivity to ABS between closely related species was studied. Since bluntnose and fathead minnows	Thatcher
	Pimephales	BCFA	-	7.7 (T4A)		are closely related phylogenetically and ecologically, one	(1966)
	notatus Lepomis macrochirus	BCFA	_	8.2 (T4A)		might expect them to be very similar in their response to a given toxicant. However, from the data in this report, this is not necessarily true since the two species were significantly different in ABS sensitivity. The differences between several species of <i>Notropis</i> also illustrate this.	

		Campostoma anomalum	BCFA	_	8.9 (T4A)			
		Notropis	BCFA	_	9.0 (T4A)			
		stramineus Ericymba buccata	BCFA	-	9.2 (T4A)			
		Notropis ardens	BCFA	-	9.5 (T4A)			
		Pimephales promelas	BCFA	-	11.3 (T4A)			
		Notropis cornutus	BCFA	-	17.0 (T4A)			
		Cyprinus carpio	BCFA	-	18.0 (T4A)			
		lctalurus melas	BCFA	-	22.0 (T4A)			
	Alkyl benzene sulfonate	Lepomis macrochirus	BSA	-	8.2 (T4A)	acde	In all of these tests, the LAS stock powder contained 60.8% LAS. The values reported were calculated on a basis of	Thatcher and Santner
		Pimephalus promelas			11.3 (T4A)		pure LAS.	(1967)
		lctalurus melas			22.0 (T4A)			
		Notropis atherinoides			7.4 (T4A)			
1		Notropis cornutus			17.0 (T4A)			
	ABS (54.8%)	Nitzschia linearis	BSA	-	10.0 (T5A)	a e	The purpose of this experiment was to determine whether there was a constant relationship between the responses of	Patrick, et al (1968)
		Physa heterostropha			34.2 (T4A)		these organisms. From the data presented, there was no apparent relationship of this type. Therefore the authors	
		Lepomis macrochirus			17.44 (T4A)		advise that bioassays on at least 3 components of the food web be made in any situation.	
	Alkyl benzene sulfonate	Salmo gairdneríi	BCFCH & A	-	0.3* (T4A)	abcdef	For a concentration of 0.45 ppm of alkyl benzene sulfonate alone, the median tolerance limit was recorded in 4 days.	Brown, et al (1968)
	plus zinc sulfate				*ABS + 0.8 ppm Zn		The zinc concentration was 0.08 ppm in the combined zinc-detergent solution. The ABS appeared to block	
8							development of resistance to Zn in the trout in chronic studies.	
MMERCIA	AC-5727 (15 percent EC)	Gambusia affinis	FL	Ponds — Bakers- field, Calif,	(O)	a c	At 0.2 lb/acre, 2 percent mortality occurred in 24 hours. At 0.8 lb/acre, 20 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
L CHEMICAL PRO	AC-5727	<i>Salmo gairdnerii</i> (one wk old sac fry)	BSA	-	0.5 (K 0%) 5.0 (K 0%)	<u>a e</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
ÄLF		(one mo old feeding fry)			0.5 (K 0%) 5.0 (K 0%)			
Ř								

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
	Pimephales	BSA	_	0.32 (T4A)	acdlef	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson
Cyanamid 12009	promelas Lepomis			0.075 (T4A)		quality of the water (pri, hardness, arkannity).	(1966)
(tech)	macrochirus Lebistes reticulatus			0.010 (T4A)			
American Cyanamid	Leiostomus xanthurus (juvenile)	BSA	_	(O)	а	Water temperature was 13 C. 20% mortality at 1.0 ppm occurred.	Butler (1965)
American Cyanamid 12009 (tech) American Cyanamid 43,913	Oyster	BCF		0.20 (O)		The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	
AC-43913	Australorbis glabratus	BSA and FL	Puerto Rico	(O)	с	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm, They were not used in field tests. Field tests showed WL 8008 to be highly effective, All others failed.	Seiffer and Schoof (1967)
AC-47031 (EC4)	Gambusia affinis	FL	Cal.	0.5 (K1)	-	Toxicity value is in Ib/acre.	Mulla (1966)
AC-47921 (EC4)	Gambusia affinis Rana catesbeiana	FL	Cal.	0.4 (O) (O)	_	At a concentration of 0.4 lb/acre, 96% mortality of the fish occurred in 24 hours. No mortality in tadpoles of <i>R. catesbeiana</i> occurred during an exposure period of one week. Toxicity value is in lb/acre.	Mulla (1966)
AC-47921 (EC4)	Gambusia affinis	FL	Cal.	0.1 (K1)	-	Toxicity value is in Ib/acre.	Mulla (1966)
Amer, Cyan- amid 52,160	Oyster	BCF	-	0.042 (0)	а	The value reported is a 96-hr $\text{EC}_{50}$ (decreased shell growth).	Butler (1965)
	Oyster	BCF		(O)		Exposure to a concentration of 1 ppm caused a 35.0% decrease in shell growth.	
	Leiostomus xanthurus (juvenile)	BSA		(O)		Water temperature was 13 C. Fish showed irritation at 1.0 ppm.	
Amer. Cyan. 52160	Rainbow trout	-	-	1.0 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
ACP-M-569	Onchorynchus tshawy tscha	BSA	_	185 (T1A) 155 (T2A)	a c d e	Concentrations were based on percent active ingredient.	Bond, et al (1960)
ACP (a-chloroaceto⊷ phenone)	Green sunfish	BSA and FL	Okla.	1.1 (T1A) 1.05 (T2A)	a e p	The main purpose of this experiment was to determine the repellent characteristics of certain chemicals. Tests were conducted at 22 C to 23 C. BSA experiments were made in a wooden trough.	Summerfelt and Lewis (1967)
Acriflavin	Microcystis aeruginosa	L	-	1.0 (K)	a, etc	The chemical was tested on a 5-day algae culture, 1 x 106 to 2 x 106 cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et (1952)

	Acriflavine	lctalurus punctatus (fry)	BSA	-	5.0 (SB3)	acdef P	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
		Lepomis macrochirus (fry)			5.0 (SB3)			
	Acriflavine	Salmo gairdnerii	BSA	-	19.9 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
		Salmo trutta Salvelinus			27.0 (T2A) 14.8 (T2A)			
		fontinalis Salvelinus			28.0 (T2A)			
		namaycush lctalurus			33.2 (T2A)			
		punctatus Lepomis macrochirus			13.5 (T2A)			
	Acrylaldehyde (acrolein)		BSA	-		а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks.	Frank, et al (1961)
1	Acti-dione	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indi- cates observation is for entire test period of 21 days): CI - PT (7) Ma - NT So - T Cv - PT (7) Gp - T Np - T	Palmer and Maloney (1955)
COMMERCIAL CHEMICAL	Aerosporin- Polymyxin B (sulfate)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Comment same as above except that: CI - T Ma - T So - T (14) Cv - T Gp - T Np - T	Palmer and Maloney (1955)
PRODUCTS	Aldrin (hexa- chloro- hexahydro- dimeth- anonaphthalene, 48 percent)	Lymnaeid snails	BSA	_	(O)	-	Each test container (500-ml beaker) was filled with ditch water. 100% mortality occurred at 1:100,000 and greater.	Batte, et al (1951)

APPENDIX B

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL PRODUCTS	Aldrin	Fathead minnow Bluegill Goldfish Guppy	BSA	-	0.033 (T4A) 0.013 (T4A) 0.028 (T4A) 0.033 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds.	Tarzwell (1959)
JCTS	Aldrin	Fathead minnow	BSA	-	0.028 (T4A)	<u>a</u>	Comments same as above except that the experiment was performed in hard water.	
	Aldrın	Pimephales promelas Lepomis macrochirus Carassius auratus	BSA	-	0.03 (T4A) 0.01 (T4A) 0.03 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient,	Henderson, et al (1959)
	Lebistes reticulatus Aldrin Pimephales (88.4%) promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus		BSA	_	0.03 (T4A) 0.032 (T4A)	a b e c d f	Dilution water was usually soft although some studies were	Henderson, et al
5				0.015 (T4A) 0.032 (T4A) 0.037 (T4A)		conducted with hard water.	Henderson, et al (1959)	
	Aldrin	Pimephales promelas Lepomis macrochirus	BSA	_	0.033 (T4A) 0.013 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)
	Aldrin (dust)	Tilapia <del>me</del> lanopleura	FLCH	Tangan- yika	1.0 lb (3.3% K) 5.0 lb (3.3% K - 3 wks) 10.0 lb (60.0% K - 3 wks)	-	Trial periods were for 20 weeks. Sublethal effects such as impaired breeding, retarded growth, or altered taste were not detected. Dosages are given as Ib/acre of surface water.	Webbe and Shute (1959)
	Aldrin	Daphnia magna	BSA	-	0.0292 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
	Aldrin	Oncorhynchus kisutch Oncorhynchus tshawytscha Salmo gairdnerii Gasterostaus aculeatus	BSA	_	45.9 (T4A) 7.5 (T4A) 17.7 (T4A) 39.8 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)

Aldrin	Gammarus Iacustris Iacustris	BSA	_	(O)	a e p	The mortality might have been partially due to the susceptibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Results were expressed as $LT_{50}$ ; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 200 (±35) min.	McDonald (1962)
Aldrin (EC 2)	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a c	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and resulted in a 90 percent mortality for the fish, and a 80 percent mortality for the tadpoles in 24 hours.	Mulla, et al (1963)
Aldrin	Lepomis macrochirus Salmo gairdneri	BSA	-	10 (T1A) 6 (T2A)	а	The experiment was conducted at 65 F.	Cope (1963)
	garoneri Aquatic insects: Ephemeroptera Trichoptera Chironomidae Fish: Moxostoma erythrurum Hypentelium nigricans Catostomus commersoni Pimephales notatus Notropis chrysocephalus Semotilus atromaculatus Campostoma anomalum Ericymba buccata Etheostoma	FR	111.	(O)	_	Dosage application rate was 2 lb aldrin/acre. After initial application a great number of fish and insects of indicated species were killed. A collection, made 7 months later, showed this stream contained a diversity of insect species and sizes of fish.	Moye and Luckmann (1964)
	zonale Hybopsis biguttata Percina						
COMMERCIAL CHEMICAL PRODUCT	maculata Notropis spilopterus N. stramineus N. volucellus Etheostoma caeruleum Notropis umbratilis						

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
drin	Bluegill	BSA	_	0.013 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
<b>j</b> rın	Notemigonus crysoleucas Lepomis macrochirus L. cyanellus	BSA	_	<ul> <li>(B) 0.080 (T 1.5)</li> <li>(A) 4.750 (T 1.5)</li> <li>(B) 0.038 (T 1.5)</li> <li>(A) 3.0 (T 1.5)</li> <li>(B) 0.062 (T 1.5)</li> <li>(A) 3.26 (T 1.5)</li> </ul>	acf	Chemical was dissolved in acetone. Final concentration of acetone was $\leq 2 \text{ mI/I}$ . Data shows $TL_m$ ppb for insecticide-resistant (A) and insecticide non-resistant (B) strains of the test fish.	Ferguson, et al (1964)
Irin	Gammarus Iacustris	BSA	-	38.5 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
lrın	Gambusia affinis affinis	BSA	_	0.05 to 2.1 (O)	а	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are exposed later.	Boyd and Ferguson (1964)
lrın	Paleomonetes kadiakensis	BSA	_	(N) 85 (T1½A) (TB) 185 (T1½A)	acf	Test organisms were collected from 2 locations, Twin Bayou (TB), Sunflower Co., Miss. (Agricultural area) and Noxubee National Wildlife Refuge (N), Noxubee Co., Miss. (non- agricultural area) and evaluated in laboratory bioassays. The Twin Bayou shrimp were more resistant.	Ferguson, et al (1965)
İrin	Acroneuria pacifica	BSA	_	0.143 (T4A)	ac	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
	Ephemerella grandis			0.009 (T4A)			
	Gammarus Iacustris			38.5 (T4A)			
	Pteronarcys californica			0.18 (T4A)			
Irin	Procambarus clarkii (juvenile)	BSA	-	0.038 (T5A)	acdo	The pesticides studied in this report are widely used in rice culture in Louisiana and are toxic to crawfish.	Hendrick and Everett (1965)
Irin	Gambusia	BSA	-	0.02-0.06 (T3A)	acde	Test fish were collected from 8 different locations of the Mississioni River. The 3 day TL - values were made to	Ferguson, et a (1965)
	affinis Ictalurus melas			0.013-0.185 (T3A)	)	determine if a resistance gradient existed. The data indi- cated that there was none.	(1903)
Irin ech)	Rainbow trout	BSA	-	0.031 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
lrin Irin	californica Procambarus clarkii (juvenile) (adult) Gambusia affinis Ictalurus melas Rainbow	BSA	-	0.038 (T5A) 0.60 (T5A) 0.02-0.06 (T3A) 0.013-0.185 (T3A)	a c d e	rice culture in Louisiana and are toxic to crawfish. Test fish were collected from 8 different locations of Mississippi River. The 3-day TL <sub>m</sub> values were made determine if a resistance gradient existed. The data cated that there was none.	f the e to

	Aldrin	Pteronarcys californica	BSA	-	0.18 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were determined by standard methods. General comments were	Gaufin, et al (1965)
		Acroneuria pacifica			0.1 (T4A)		made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional data are	
		Ephemerella grandis			0.009 (T4A)		presented.	
		Daphnia magna			0.03 (T 50 hr A)			
		Gammarus Iacustris			38.5 (T4A)			
	Aldrin (tech, 93 percent	Pteronarcys californica (naiad)	BSA	-	0.180 (T4A)	c d e f	A. pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than P. californica.	Jensen and Gaufin (1964)
	active in acetone)	Acroneuria pacifica (naiad)			0.143 (T4A)			(1504)
	Aldrin	Bluegill	BSA	-	9.7 (T4A) 7.7 (T4A) 6.2 (T4A)	<u>a</u>	These experiments were performed to demonstrate that at increased temperatures the toxic effect of most chemicals is increased.	Cope (1965)
Ð					5.6 (T4A)		For the toxicant concentrations listed, the temperatures were respectively, 45, 55, 65, 75, and 85 F. Data on the effect of time as well as temperature was also reported. The experimental animals all were approximately one gram in weight.	
-	Aldrin	Acroneuria pacifica	BSA & CFCH	_	0.143 (T4A) 0.022 (T30CH)	<u>acde</u>	Additional data are presented,	Jensen and Gaufin
		Pteronarcys californica			0.180 (T4A) 0.0025 (T30CH)			(1966)
0	Aldrin	Procambarus clarkii	FO	Crowley, La.	(O)	c d e p	Experiments were conducted in a flooded rice field. Area was divided into 4 blocks with a fence, restricting crawfish to desired areas. The rearing of crawfish in rice fields is of considerable commercial importance in Louisiana. No untoward effect on the crawfish occurred. Aldrin was used on the rice seed at the rate of 0.25 lb/100 lb seed. Even with the addition of carbonyl solution 0.8 lb/acre showed no more effect.	Hendrick, et al (1966)
COMMERCIAL CHEMICAL PRI	Aldrin	Daphnia magna	BSA	-	0.030 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Aldrin	Notropis umbratilis N. umbratilis (2 in.)	BSA	-	0.02-0.08 (T4A) 0.4 (T4A)	acd <u>e</u>	Aerated pond water was used as diluent. Both aquarium and a "boat" were used as test vessels. Other experiments with oxygen concentration variations are reported.	Proffitt (1966)
ALPR		N. cornutus			0.02-0.08 (T4A)			

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	N. blennius			0.6 (T6A)			
	(2 in.)						
	Spotfins						
	(2 in.)			0.6 (T6A)			
	(3 in.)			0.6 (T8A)			
	Lepomis						
	macrochirus						
	(<1-1/2 in.)			0.2 (T2A)			
	(1-1/2 in.)			0.4 (T4A)			
	(2 in.)			0.6 (T6A)			
	L. cyanellus						
	(1-1/2 in.)			0.4 (T4A)			
	(3 in.)			0.6 (T6A)			
	Microptera						
	salmoides						
	(2-1/2 in.)			0.4 (T4A)			
	Fundalus						
	notatus						
	(1-1/2 in.)			0.6 (T8A)			
	Etheostoma						
	flabellare						
	(2 in.)			0.6 (T8A)			
	Noturus			0.6 (T8A)			
	miurus						
	Etheostoma						
	nigrum						
	(2 in.)			0.8 (T10A)			
	E. caeruleum						
	(2 in.)			0.8 (T10A)			
	E. blennioides						
	(2-1/2 in.)			0.8 (T10A)			
	Campostoma						
	anomalum						
	(5 in.)			0.8 (NTE)			
	Hypentelium						
	nigricans						
	(5-1/2 in.)			0.8 (NTE)			
	Ericymba			0.21 (K2A)			
	buccata						
	Hypognathus			0.25 (K2A)			
	nuchalis						
Idrin	Símocephalus	BSA	-	0.023 (SB)	-	Concentration reported is for immobilization.	Sanders and
	serrulatus					Time for immobilization was 48 hr.	Cope
	Daphnia pulex			0.028 (SB)		Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	(1966)
Aldrin	Catfish Buffalo fish Perch Blu <del>eg</del> ill	L	~	(O)	-	The chemical was found from 0.02 to 0.21 ppm as residues in catfish, and 0.01 to 0.04 in buffalo fish — after a soil treatment nearby of 5 lb/acre.	Sparr, et al (1966)

	Aldrin	Oyster	BCF	-	0.001 (SB4) 1.0 (SB4)	a	Seawater was employed in this experiment.	Butler (1966)
	Aldrin	Daphnia carinata	BSA	-	0.0040 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Aldrin	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	-	(O)	-	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentra- tion of 350-4500X resulted.	Butler (1967)
	Aldrin	Oyster	FE	Galveston Bay, Texas	(0)	-	Oysters from the area were found to contain from none to 0.03 ppm.	Casper (1967)
D 13	Aldrin	Lampsilis siliquoidea L. ventricosa Anodonta grandis	F	Red Cedar River, Mich.	(0)	-	The mussels listed were analyzed for the toxicant and its metabolites. Mussels may be used as detectors for this toxicant, because they tend to concentrate the chemical in much higher concentrations than it is ever found in the water. The amount of chemical applied as a spray was not specified.	Bedford (1968)
	14 <sub>C</sub> -Aldrin	Carassius auratus (Linnaeus)	BSA	_	0.05 (SB)	а	Immediately after 8-hr exposure <sup>14</sup> C-Dieldrin was detected in various tissues; percentages increased with time until at 32 days they were 93.9% or more except for visceral fat; 50 and 100% of the residues were Dieldrin within 2.5-5.4 and 31.5-92.4 days, respectively; in visceral fat the corresponding times were 46.9 and 14,733 days.	Gakstatter (1968)
	Aldrin	Pteronarcys californica (naiads)	BSA	-	0.0013 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
COMMERCIAL	Algeeclear	Chlorella pyrenoidosa	L		20 (AS 1)	-	Describes a bioassay method to differentiate between an algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)
CIAL CH	Algibiol	Phoxinus phoxinus	BSA	-	25 (K2A) 20 (T1A)	<u>a</u> cde	The assays were conducted in a dual aquarium with aeration. The highest dilution that was nonletal was 7.5 ppm.	Vivier and Nisbet (1965)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL	Algımaster	Chlorella py renoidosa	L.	_	3.0 (AC<1/2)	_	Describes a bioassay method to differentiate between an algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)
70	Algimycin (MT-4)	Chlorella pyrenoidosa	L	_	3.0 (AC < 1/10)	-	Comment same as above.	Fitzgerald and Faust (1963)
RODUCTS	Algimycin 200	Chlorella pyrenoidosa	L	_	3.0 (AC ≤ 1/2)	_	Comment same as above.	Fitzgerald and Faust (1963)
•	Allethrin	<i>Pteronarcys</i> sp (nymphs)	BSA	_	0.0021 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
	Allethrin (tech)	Rainbow trout	BSA	-	0.019 (T4A)	-	The values reported are given as $LC_{50}$ .	Соре (1965)
	Allethrin	Simocephalus serrulatus Daphnia pulex	BSA	_	0.056 (SB) 0.021 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Allethrin	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 67 percent.	Jamnback and Frempong- Boadu (1966)
	Allethrin	Pteronarcys californica (naiads)	BSA	-	0.0021 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
	Altacide 2,4-D	Spatterdock	FL	Fla.	(O)	-	At 10.0 lb/acre, 2 percent control of spatterdock was obtained.	Copeland and Woods (1959)
	p-aminophenol	Daphnia magna	BSA	-	2 (K2A)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical sub- stances. Results indicated a considerable correlation between the aquarium fish toxicity and antiautocatalytic potency of the chemicals in marked contrast to their toxicity on sustain a completeration	Soliman (1949)

toxicity on systemic administration.

APPENDIX B

	Amiton oxalate	Carassius auratus Lepomis macrochirus	BSCH	-	10 (O)* 10 (O)* *in response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
	Amitrol-T	Lepomis macrochirus	BSA	-	(0)	а	No mortality in 2-in. fish was noted with concentrations of 10,000 mg/I over 100 hr at 65 F.	Cope (1963)
	Ametryne	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	1.0 (NTE) 1,0 (0, 10%) 1.0 (NTE) —	_	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Butler (1965)
B-15	Ametryne	Penaeus aztecus	L	-	(0)	а	Toxicant chemicals were evaluated in sea water at temper- atures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
01		Oyster	BCF	_	(O)	а	Exposure to a concentration of 1.0 ppm caused a 14.0% decrease in shell growth.	
		Leiostomus xanthurus (juvenile)	BSA	-	(0)	а	Water temperature was 28 C. No effect was noticed on exposure to 1.0 ppm.	
	Aminotriazol	<b>P</b> anicum hemitomum	FL	Fla.	(O)	-	At 10 ib/acre, 5-7 percent control of <b>P</b> . hemitomum was obtained.	Copeland and Woods (1959)
8	Aminotriazole	Oncorhynchus kisutch Micropterus salmoides	BSA and CF	-	325 (T1A) 325 (T2A) (O)	a c d e	Concentrations were based on percent active ingredient. In the constant-flow (CF) apparatus, 1000-ppm amino- triazole killed all test fish in 6 days.	Bond, et al (1960)
COMMERCIAL	Aminotriazole	Salmon	BSA	-	325 (T2A)	-	Data are given as $LC_{50}$ .	Bohmont (1967)
	Amitrole	Daphnia magna	BSA	-	23 (15.3- 44.4) (O)	acdiq	Toxicity, in terms of median immobilization concentration ( $IC_{50}$ ), is presented.	Crosby and Tucker (1966)
CHEMICAL PRODUCTS	Amitrole	Lepomis macrochirus (eggs) L. cyanellus (eggs) Micropterus dolomieui (eggs)	L	_	50 (NTE) 50 (NTE) —	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Erimyzon sucetta (eggs) L. macrochirus (fry)			50 (NTE) 25 (S)			
Amitrole T	Daphnia magna	BSA	_	40 (14.3- 112.0) (O)	acdiq	Toxicity, in terms of median immobilization concentration $(IC_{50})$ , is presented.	Crosby and Tucker (1966)
Ammate	Althernanthera philoxeroides Typha latifolia	FL	Fla.	(O)		At 76 and 120 lb/acre, respectively, 1 percent control of alligator weed was obtained while 80 percent control of cattail was obtained with the higher application rate.	Copeland and Woods (1959)
Ammate	Channel catfish (fingerlings)	BSA	_	259 (K1A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Amopyroquin	Salmo gairdneri	BSA	_	35.3 (T2A)		Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
	salmo trutta			36 (T2A)		were also determined.	(1986)
	Salvelinus fontinalis			40 (T2A)			
	Salvelinus			14 (T2A)			
	namaycush Ictalurus			12.5 (T2A)			
	punctatus Lepomis macrochirus			18.5 (T2A)			
Antimycin A	Salmo gairdneri	BSA	-	0.25 (T18 hr)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)
Antimycin A	Dorosoma cepedianum	BSA	-	800 (K1A) 100 (K4A)	adefil mp	Results were reported at 12 C. All fish were killed in 24 hr by 40 ppm at 22 C.	Walker, et al (1964)
	Salmo gairdneri			600 (K1A) 80 (K4A)		Results were reported at 12 C.	
	S. trutta			400 (K1A) 80 (K4A)		Results were reported at 12 C.	
	Esox Ieucius			800 (K1A)		Results were reported at 12 C. All fish were killed in 24 hr by 200 ppm at 17 C; by 100 ppm at 22 C.	
	Compostoma anomalum			1,000 (K1A)			
	Carassius auratus			100,000 (K1A) 2,000 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 4,000 ppm at 22 C.	
	Notemigon us crysoleucas			2,000 (K1A) 600 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 500 ppm at 22 C.	

	Pimephales promelas			2,000 (K1A) 400 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 800 ppm at 22 C.	
	Catostomus commersoni			220 (K4A)		Results were reported at 12 C.	
	lctiobus cyprinellus			400 (K4A)		Results were reported at 12 C.	
	lctalurus melas			120,000 (K1A) 80,000 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 40,000 ppm at 22 C.	
	l. natalis			80,000 (K1A)		Results were reported at 12 C.	
	l. punctatus			20,000 (K1A)		Results were reported at 12 C. All fish were killed in 124 h by 6,000 ppm at 22 C.	r
	Eucalia inconstans			5,000 (K1A)		Results were reported at 12 C.	
	Lepomis cyanellus			2,000 (K1A) 800 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 800 ppm at 22 C.	
	L. gibbosus			2,000 (K1A) 200 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 200 ppm at 22 C.	
	L. macrochirus			1,000 (K1A) 400 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 200 ppm at 22 C.	
	L. megalotis			2,000 (K1A) 400 (K4A)		Results were reported at 12 C. All fish were killed in 24 hr by 4,000 ppm at 22 C.	
	Micropterus salmoides			6,000 (K1A) 800 (K4A)		Results were reported at 12 C.	
	Poxomis annularis			2,000 (K1A)		Results were reported at 12 C.	
	Etheostoma exile			660 (K1A)		Results were reported at 12 C.	
	Perca flavescens			660 (K1A)		Results were reported at 12 C. All fish were killed in 24 hr by 660 ppm at 22 C.	
	Stizostedion vitreum			660 (K1A)		Results were reported at 12 C.	
	Cyprinus carpio			2,000 (K1A) 600 (K4)		Results were reported at 12 C.	
Antimycin A	Salmo gairdneri	FL	Wisc.	600 (K1A) 80 (K4A)	acdg	Results were recorded at 12 C. All fish were killed in 24 hr by 80 ppm at 17 C.	Walker, et al (1964)
2	S. trutta			400 (K1A) 80 (K4A)		Results were recorded at 12 C. All fish were killed in 24 hr by 60 ppm at 17 C.	
Aquaherb (2,4-D ester)	Althernanthera philoxeroides	FL	Fla.	(O)	-	At 14.2 pounds per acre, only 1-2 percent control of alligate weed was obtained.	or Copeland and Woods (1959)
Aqualin (acrolein)	Salmo trutta	BCFA	-	0.046 (T1CFA)	<u>a</u> c <u>e</u>	Spring water was used as dilution water. The chemical was found to be toxic to the test fish at concentrations below	Burdick, et al (1964)
	Lepomis machrochirus			0.079 (T1CFA)		that recommended to control aquatic vegetation.	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Aqualin Aqualin Aqualin	Carassius auratus	BSA	-	1.0-2.0 (K 3 hr)	а	Fiber glass tanks were used as test containers. Goldfish were acclimated to the tank habitat for 2 weeks before testing. Detoxification of the tank occurred within 43 hours when a concentration of 3.0 ppm was applied.	St. Amant, et al (1964)
Aqualin	Carassius auratus	FL	California	3.0 (K1) 2.0 (K 18 hr) 1.0 (K1)	a c e	The chemical was applied to Big Bear Lake at 3 ppm. Within 24 hours all fish in the area died. Fish were placed in Mentone pond in 3 live cars. An area between 2 dams separate from Big Bear Lake was tested. This area was made up of 26 acre-feet of water at a surface temperature of 72 F between the two dams. At 1.0 ppm distress of fish was evident in 1 hour and most visible fish died in 2 hours. In 1 day, all fish in live cars were dead.	St. Amant, et al (1964)
	lctalurus nebulosus			2.0 (K 18 hr)		Fish were placed in Mentone pond in 3 live cars.	
Aqua San	Pestia stratiotes	FL	Fla.	(0)	-	At 32.0 lb/acre, 2 percent control of water lettuce was obtained.	Copeland and Woods (1959)
Aquasan (colloidal Ag)	Microcystis aeruginosa	L	-	100 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Aqua San 2,4-D	<i>Pestia stratiotes</i> Spatterdock	FL	Fla.	(O)	-	At 20.0 and 25.0 lb/acre, respectively, 8 percent control of water lettuce was obtained while spatterdock was not controlled at the higher rate.	Copeland and Woods (1959)
Aquathol	Gammarus Iacustris	BSA	-	>320 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Aquatic	Richardsonius balteatus hydroflox	BSA	-	83 (T1A) 75 (T2A) 75 (T4A)	acdef	Results given were in soft water. Results in hard water were as follows: 57 (T1A), 83 (T2A), and 78 (T4A).	Webb (1961)
Aramite -chloro- ethyl, -(P-tertiary- butylphenoxy) methty-ethyl sulfite, 15 percent		BSA	-	(O)	-	Each test container, 500-ml beaker, was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
Aramite (15%)	Channel catfish (fingerlings)	BSA	-	>100 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Aramite (tech)	Rainbow trout	BSA?		0.320 (T4A)		The values reported are given as $LC_{50}$ .	Соре (1965)

	Aramite	Simocephalus serrulatus Daphnia pulex	BSA	-	0.180 (SB) 0.160 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)	
	Armazide	Chlorella pyrenoidosa	L	-	3.0 (AC 1)	-	Describes a bioassay method to differentiate between an algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)	
	ATA (3 amine- 1,2,3, trizole as the active ingredient)	Richardsonius balteatus hydroflox	BSA	-	1330 (T1A) 1163.3 (T2A) 983.3 (T4A)	acdef	Results given were in soft water. Results in hard water were as follows: >3600 (T1A), >3600 (T2A), and 1370 (T4A).	Webb (1961)	
	Atabrine	Microcystis aeruginosa	L	-	100 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)	
	Atabrine	Channel catfish (fingerlings)	BSA	-	0.93 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	
	Atlas "A"	Najas quadalupensis	FL	Fla.	(0)		At 50.3 lb/acre, <i>N. quadalupensis</i> was not affected.	Copeland and Woods (1959)	APF
	Atlas 1901	Pandalus montagni Crangon crangon Carcinus maenas Cardium edule	BSA	-	87.2 (T2A) 120.0 (T2A) 150.0 (T2A) 48.5 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C. It was shown that the toxicity of this solvent emulsifier decreased with time, due to evaporation of the solvent. Atlas 1901 at a concentration of 33.3 ppm killed 95% of <i>Crangon crangon</i> larvae in 3 hr.	Portmann and Connor (1968)	APPENDIX B
	Atlox 2082 A (spray emulsifier for DDT)	Oncorhyncus kisutch	BSA	-	20.7 (T2A)	а	The figure cited is calculated from the data. The compound is an alkyl sulfonate.	Alderdice and Worthington (1959)	
COMMERCIAL CHEMI	Atrazine	Micropterus salmoides (fry) Ictalurus punctatus (fry) Lepomis macrochirus (fry)	BSA	-	5.0 (SB3) 10 (SB3) 10 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)	
≦									

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Atrazine	Phoxinus phoxinus	BSA		5.0 (K2A) 1.25 (K2A)	<u>a</u> cde	The assays were conducted in a dual aquarium with aera The chemical was still toxic to minnows at 2.5 and 5.0 in the presence of plants. Kill occurred between 8-15	ppm Nisbet
Atrazine (gesaprime)	Phoxinus phoxinus	BSA	~	10 (K2A)	<u>a</u> d c e	The maximum nonlethal dose in 48 hours was 2.5 pp	n. Vivier and Nisbet (1965)
Atrazine	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	(0)	а	Water temperature was 28 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Oyster	BCF		(0)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
Atrazine	Crassostrea virginica Penaeus aztecus Leiostomus	BCFA & BSA	-	1.0 (NTE) 1.0 (NTE)	-	Seawater was pumped continuously into test aquaria. Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:	Butler (1965)
	xanthurus Phytoplankton					Oyster -96-hr EC50 - Conc. which decreas shell growth.Shrimp -48-hr EC50 - Conc. which killed or paralyzed 50% of test animals.Fish -48-hr EC50 - Conc. which killed 50%.Phytoplankton -Percent decrease of CO2 fixation t 4-hr exposure at 1.0 ppm chemical concentration.	or O a
Atrazine (WP)	Lepomis macrochirus (eggs) Micropterus dolomieui (eggs) Erimyzon sucetta (eggs) L. macrochirus (fry)	L	-	(O) 10/3 (O) 10 (NTE) 5.0 (S)		Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number o days survival. Maximum length of test was 8 days. No was added. Small bluegill were tested to find the high concentration of chemical which did not cause death 12 days (S).	o food est
Atrazine (granular)	Lepomis macrochirus (eggs)	L	_	10 (NTE)	-	Comment same as above.	Hiltibran (1967)
	L. Cyanellus (eggs)			10 (NTE)			
	L. macrochirus (fry)			10 (S)			

	Azida		564		(0)			Balah
	Azide	Mayorella palestinensis soil amoeba	BSA	_	(O)	<u>a c</u>	The experiments were carried out in Warburg manometers at 27 C for 4 hr as a pH of 8.0 Azide in concentrations up to 2 x 10 <sup>-3</sup> M were shown to have lethal effects on the organism. Results were compared with controls and expressed in percent of respiration. Compared with normal respiration, nonlethal concentrations of azide increased the respiration of the organism in glucose- containing solutions. It was concluded that the respiration of the organism depends on at least three enzyme systems, which may be distinguished by their behavior toward azide.	Reich (1955)
	Bayer 29493 (Baytex)	Procambarus simulans simulans	FL	Texas	0.25 (K2) 0.37 (K2) 0.50 (K2)	acdp	Bluegills held in wire boxes were not affected at the indi- cated concentrations. Water temperature was 58 F at the time of treatment, 49 F at drainage. Largemouth bass showed distress and some crappies died in waters treated with 0.33 ppm at 85 F. No deaths were noted in waters at 80 F.	Lowman (1965)
	Banvel D	Lepomis macrochirus	BSA	-	(L) 410 (T2A) (G) 20 (T2A)* (G) 67.5 (T2A)** *vermiculite **attapulgite	<u>a</u> cdeg	Toxicity data for 24 and 48 hours are presented for liquid (L) and granular (G) formulations. Various commercial formulations were tested. The liquid formulations were almost invariably more toxic than the granular ones.	Hughes and Davis (1965)
B-21	Baron	Onchorynchus tshawytscha Micropterus salmoides	BSA	-	2.62 (T1A) 2.3 (T2A) 4.6 (T1A)	a c d e	Concentrations were based on percent active ingredient.	Bond, et al (1960)
	Baron	Channel catfish (fingerlings)	BSA	_	7.2 (K1A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Bay 73 (WP 71)	Micropterus salmoides	BSA	_	0.05 (O) 0.10 (K1)	a e	At 0.05 ppm, 12 percent mortality occurred in 1 day. Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.	Mulla, et al (1967)
COMMER	Bay 73 (WP 71)	Micropterus salmoides Cyprinus carpio	FL	Chino Fishery bass pond, Cal.	0.10 (O) 0.25 (K2) 0.10 (O) 0.25 (O)	а	For bass: At 0.10 ppm, no mortality occurred in 1 day. At 0.25 ppm, 50 percent mortality occurred in 1 day, and 100 percent occurred in 2 days. For carp: At 0.10 and 0.25 ppm, 10 percent mortality occurred.	Mulla, et al (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Bayer 73	Salmo	BSA	_	0.052 (T2A)*	<u>a c</u>	Various temperatures (12 and 17 C) and water qualities in	Marking and
	gairdnerii Salvelinus fontinalis			0.016 (T2A)*		static bioassays did not influence the toxicity greatly, but pH variations in chemically buffered solutions did.	Hogan (1967)
	Carassius auratus			0.279 (T2A)			
1	Cyprinus			0.139 (T2A)*			
1	carpio			0.148 (T2A)			
				0.103 (T2A)			
	Pimephales						
	promelas Catastomus			0.081 (T2A)*			
	conmersoni			0.061 (12A)			
	Ctiobus			0.064 (T2A)			
	cyprinellus						
	lctalurus			0.096 (T2A)*			
	melas			0.084 (T2A)			
	l. punctatus						
	Pylodictis			0.043 (T2A)			
	olivaris Lepomis			0.115 (T2A)			
	Cyanellus			0.115 (12A)			2
	L. macrochirus			0.098 (T2A)*			1
	2			0.082 (T2A)			1
	L. microlophus			0.153 (T2A)			
	Micropterus			0.089 (T2A)			
	dolomieui			0.007 (To 4)			
	M. salmoides			0.097 (T2A)			
	Perca flavescens			0.081 (T2A)*			
	Talapia			0.150 (T2A)			
	mossambica			0.100 (12) ()			
	I. nebuilosus			0.071 (T2A)			
				*12 C, other			
				data at 17 C			
Bayer 73 (tech)	Rainbow trout	BSA	-	0.320 (T4A)	-	The values reported are given as $LC_{50}$ .	Соре (1965)
Bayer 73 (WP 71%)	Gambusia affinis	FL	Cal.	1.0 (K1)	-	Toxicity value is in Ib/acre.	Mulla (1966)
Bayer 73 (tech)	Pteronarcys californica (naiads)	BSA	~	0.0002 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
Bayer 4731	Leiostomus xanthurus (juvenile)	BSA	-	0.032 (O)	а	Water temperature was 13 C. The figure reported is a 48-hr $\text{EC}_{\overline{50}}.$	Butler (1965)
Bayer 9018 (tech)	Rainbow trout	BSA	-	0.320 (T4A)	-	The values reported are given as LC <sub>50</sub> .	Соре (1965)

	Bayer 22408	Anopheles quadrimaculatus	BSA	-	0.04 (K1)	-	4th instar larvae of mosquitos were used in this bioassay. Adsorption was determined by use of P <sup>32</sup> labeled Bayer 22488.	Schmidt and Weidhaas (1961)
	Bayer 22408 (EC2)	Gambusia affinis	FL	Ponds in III.	(0)	-	When applied at 2.0 pounds per acre active ingredients, 12 percent fish mortality occurred in 1 day.	Mulla, et al (1963)
	(202)	Rana catesbeiana					No bullfrog mortality occurred at 2.0 pounds per acre in 1 day.	
	Bayer 25198 (50 percent EC)	Gambusia affinis	FL	Ponds - Bakers- field, Cal.	(O)	a c	At 0.1 lb/acre, 4 percent mortality occurred in 24 hours. At 0.4 lb/acre, 8 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
	Bayer 25141 (50 percent EC)	Gambusia affinis	FL	Ponds - Bakers- field, Cal.	(0)	a C	At 0.2 lb/acre, 34 percent mortality occurred in 24 hours, and at 0.8 lb/acre, 100 percent kill occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
	Bayer 25141 (tech)	Lepomis macrochirus	BSA	-	0.056 (T4A)	acdef	The toxicity of this substance was not influenced by the the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson (1966)
	Bayer 29492 (EC2)	Gambusia affinis	FL	Ponds in III.	(0)	~	When applied at 0.2 pound per acre active ingredient, 100 percent mortality occurred in 1 day.	Mulla, et al (1963)
	Bayer 29493 (Baytex)	Carassius auratus Lepomis macrochirus Notemigonus crysoleugus	BSCH	-	20 (0)* 20 (0)** * no response, 15 days ** response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
	Bayer 29493	Gammarus Iacustris	BSA	-	0.0138 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
COMMERCIAL	Bayer 29493 (tech, 93 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica (naiad)	BSA	-	0.0265 (T4A) 0.0051 (T4A)	c d e f	<i>A. pacifica</i> was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than <i>P. californica</i> .	Jensen and Gaufin (1964)
CHEMICAL	Bayer 29493 (25 percent EC)	Gambusia affinis	FL	Ponds Bakers- field, Cal.	(0)	a c	At 0.1 lb/acre, 6 percent mortality occurred in 24 hours. At 0.4 lb/acre, 16 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulia and Isaak (1961)
_								

B-23

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Bayer 29493	Chaoborus astictopus Lepomis macrochirus	BSA	_	(0)	а	Tests were conducted on bluegill sunfish, <i>C. astictopus</i> first instar larvae, and fourth instar larvae, results on larvae were as follows: Fourth instar 0.007 (T1A) First instar 0.0043 (T1A)	Hazeltine (1963)
Bayer 29493 (tech)	Rainbow trout	BSA	_	0.760 (T4A)		The values reported are given as $LC_{50}$ .	Cope (1965)
Bayer 29493 (Baytex)	Procambarus simulans simulans	BSA	-	0.18 (K1A)	a c d	Bioassays showed that concentrations to 5.6 ppm in 96 hr did not kill fingerling channel fish, largemouth bass, and redear sunfish.	Lowman (1965)
Bayer 29493	Pteronarcys californica Acroneuría	BSA	_	0.03 (T4A) 0.005 (T4A)	a	Unspecified chemical characteristics of assay water were determined by standard methods. General comments were made concerning "standardized" conditions, use of "soft"	Gaufin, et al (1965)
	pacifica Ephemerella grandis Gammarus lacustris			0.02 (T4A) 0.01 (T4A)		water, and use of emulsifying agents. Additional data are presented.	
Bayer 29493	Acroneuria pacifica Ephemerella grandis Gammarus lacustris Pteronarchys californica	BSA	-	0.005 (T4A) 0.025 (T4A) 0.014 (T4A) 0.026 (T4A)	<u>a</u> c	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
Bayer 29493	Acroneuria pacifica Pteronarcys californica	BSA & CFCH		0.0051 (T4A) 0.00064 (T30A) 0.00265 (T4A) 0.00360 (T30A)	<u>a c d e</u>	Additional data are presented.	Jensen and Gaufin (1966)
Bayer 29493 (Baytex)	Pimephales promelas Lepomis macrochirus Lebistes reticulatus	BSA	-	3.3 (T4A) 3.1 (T4A) 3.1 (T4A)	acdef	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, and alkalinity).	Pickering and Henderson (1966)
Bayer 29952 (EC2)	Gambusia affinis Rana catesbeiana	FL	Ponds in III.	(0)	_	When applied at 0.025 pound/acre active ingredient, 100 percent fish mortality occurred in 1 day. When applied at 0.4 pound/acre, 5 percent bullfrog mortality occurred in 1 day.	Mulla, et al (1963)
Bayer 30749 (EC4)	Gambusia affinis	FL	Ponds in III.	(0)		When applied at 0.8 pound/acre active ingredient, 100 percent fish mortality occurred in 1 day.	Mulla, et al (1963)

	Bayer 34042 (EC4)	Gambusia affinis Rana Catesbeiana	FL	Ponds in III.	(0)	_	When applied at 0.025 pound/acre active ingredient, 100 percent fish mortality occurred in 1 day. No bullfrog mortality occurred at 0.4 pound/acre in 1 day.	Mulla, et al (1963)
	Bayer 37289 (EC4)	Gambusia affinis Bufo boreas Scophiopus hammondi	FL	Ponds in III.	(0)	_	When applied at 0.8 pound/acre active ingredient, 52 percent fish mortality occurred in 1 day. No toad mortality occurred at 0.4 pound/acre in 1 day.	Mulla, et al (1963)
	Bayer 37289	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.0001 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
	Bayer 37289	<i>Leiostomus xanthurus</i> (juvenile)	BSA	_	0.32 (O)	а	Water temperature was 13 C. The figure reported is a 48-hr ${\sf EC}_{50}.$	Butler (1965)
	Bayer 37289 (tech)	Rainbow trout	BSA	-	0.240 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
	Bayer 37289	Pteronarcys californica (naiads)	BSA	-	0.0001 (T4A)	<u>acdef</u>	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
;	Bayer 37342 (EC4)	Gambusia affinis	FL	Ponds in III.	(0)	-	When applied at 0.4 pound/acre active ingredient, 24 percent mortality occurred in 1 day.	Mulla, et al (1963)
	Bayer 37343 (EC2)	Gambusia affinis	FL	Ponds in III.	(0)		When applied at 0.025 pound/acre active ingredient, 0 percent mortality occurred in 1 day.	Mulla, et al (1963)
	Bayer 37344 (tech)	Rainbow trout	BSA	-	0.640 (T4A)	-	The values reported are given as $LC_{50}$ .	Соре (1965)
	Bayer 37344	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.0054 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
g	Bayer 37344	Pteronarcys californica (naiads)	BSA	_	0.0054 (T4A)	<u>acdef</u>	Data reported as LC $_{\!$	Sanders and Cope (1968)
COMMERCI	Bayer 38156 (50 per- cent EC)	Gambusia affínis	FL	Ponds — Bakers- field, Cal.	0.1 (K1) 0.4 (K1)	a c	Toxicity values indicate application rates in Ib/acre. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
AL CHEMICAL	Bayer 38156	Leiostromus xanthurus Cyprinodon variegatus Mugil cephalus	BCFCH	-	0.001 (O) 0.001 (O) 0.001 (O)	а	At a concentration of 0.001 ppm, the following percent acetylcholinesterase activity as compared to controls was found: <i>L. xanthurus</i> – 76 <i>C. variegatus</i> – 82 <i>M. cephalus</i> – 58	Butler and Johnson (1967)
Ð								

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Bayer 38819 (tech)	Rainbow trout	BSA		0.450 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
Bayer 38920 (EC4)	Gambusia affinis Rana catesbeiana	FL	Ponds in 111.	(O)	-	When applied at 0.5 pound per acre active ingredient, 100 percent mortality of both species occurred in 1 day.	Mulla, et al (1963)
Bayer 41831 (EC4)	Gambusia affinis Rana catesbeiana	FL	Ponds in III.	(O)	-	When applied at 1.6 pounds per acre active ingredients, 44 percent fish mortality occurred in 1 day. No bullfrog mortality occurred at 0.8 pound per acre in 1 day.	Mulla (1963)
Bayer 41831	Pteronarcys sp (nymphs)	BSA	-	0.0038 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
Bayer 41831 (tech)	Rainbow trout	BSA	_	0.700 (T4A)	_	The values reported are given as $LC_{50}$ .	Cope (1965)
Bayer 41831	Cyprinodon variegatus (juvenile)	BSA	_	(0)	а	Water temperature was 9 C. Fish showed irritation at 1.0 ppm.	Butler (1965)
Bayer 41831	Pteronarcys californica (naiads)	BSA	-	0.004 (T4A)	<u>acdef</u>	Data reported as LC $_{\!$	Sanders and Cope (1968)
Bayer 46676 (EC2)	Gambusia affinis	FL	Cal.	0.2 (K1)	_	Toxicity value is in Ib/acre.	Mulla (1966)
Baygon	Prosimulum spp Cnephia spp Simulium spp (Iarvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 83 percent.	Jamnback and Frempong- Boadu (1966)
Baγgon	Pteronarcys californica (naiads)	BSA	-	0.013 (T4A)	<u>acdef</u>	Data reported as LC $_{\!$	Sanders and Cope (1968)
Bay tex	<i>Salmo gairdnerii</i> (one wk old sac fry)	BSA	-	0.2 (K 0%) 2.0 (K 0%)	<u>a e</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
	(one mo old feeding fry)			0.2 (K 0%) 2.0 (K 0%)			

	Baytex	Culex pipiens quadrimaculatus	BSA	_	(0)	С	Tests were conducted in tap water and artificially polluted tap water. The values reported are the concentration range for an LC90, 0.0015 to 0.0080 ppm in polluted and 0.0060 to 0.0160 in tap water.	Lewallen and Wilder (1963)
	Baytex	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.0044 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
	Baytex	Gambusia affinis Fundulus chrysotus Lepomis macrochirus Lepomis microlophus Chaenobryttus gulosus	BSA & FL	-	Not affected 5 (K2) 5 (K2) 5 (K2) 5 (K2)	a C	None of the fish showed overt symptoms of Baytex poisoning at a concentration of 0.025 ppm which is the equivalent of an application rate of 0.2 pound per acre. Some mortality occurred at 2.5 ppm concentration after 48 hours. There was little danger of acute poisoning to these species of fish when it was applied at 0.2 pound per acre. Long range effects of the chemical on other aquatic organisms were studied in plastic-lined ponds of 300 gallon capacity. Baytex was applied at 0.2 pound (tech) per acre. All of the <i>Cladocera</i> and chironomid population at 0.2 pound per acre were almost completely eliminated in the treated pond within a week. Copepods, ostracods, hydra, and annelid worms exhibited no noticeable population change.	
	Baytex	Oyster	BCF	-	1.0 (SB4)	а	Seawater was employed in this experiment.	Butler (1966)
1	Baytex	Simocephalus serrulatus Daphnia pulex	BSA	-	0.00092 (SB) 0.00080 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Ben Venue 35 (tech)	Rainbow trout	BSA	-	3.0 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
	Ben Venue 3835 (tech)	Rainbow trout	BSA	-	0.380 (T4A)	-	Comment same as above.	Cope (1965)
	Ben Venue 54 (tech)	Rainbow trout	BSA	-	0.480 (T4A)	-	Comment same as above.	Cope (1965)
8	Benzene hexachloride	Bluegill	BSA	-	0.45 (SB)	(0)	Bluegills tolerated concentrations of 0.45 ppm. A field study is also described.	Linduska and Surber (1948)
COMMERC	a-benzene hexachloride	Microcystis aeruginosa	L	_	50 (K)	<u>a</u>	The chemical was tested on a 5-day algae culture, 1 x $10^6$ to 2 x $10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
CIAL CHEMICAL	Benzene hexa- chloride (various isomers, tech)	Lymnaeid snails	BSA	-	(0)	-	Each test container (500-ml beaker) was filled with ditch water. Less than 100% mortality occurred in concentrations of 1:100,000.	Batte, et al (1951)
ICAL PROD	Benzene hexa- chloride (gamma-isomer, 5 percent)	Lymnaeid snails	BSA	-	(0)	-	Each test container (500 ml-beaker) was filled with ditch water. 100% mortality occurred in concentrations of 1:600,000 and greater.	Batte, et al (1951)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)	
AL CHEMIC	Benzene hexa- chloride (99.8 percent isomer)	Lymnaeid snails	BSA	_	(0)	_	Each test container (500 ml-beaker) was filled with ditch water. 100% mortality occurred in concentrations of 1:600,000 and greater.	Batte, et al (1951)	
CAL PRODUCTS	Benzene hexa- chloride	Oncorhynchus kisutch (fry)	FL	Sproat Lake, Canada	(O)	_	Tests were in fresh water and seawater. No difference in toxicity was observed due to water type. The chemical was sprayed as an emulsion from a plane, at the rate of 12.3 lb per 34,848 sq ft. The fish were exposed in boxes submerged in the water. At a concentration of 1.38 ppr., 5 of 15 fish survived 15 minutes; at 0.36 ppm 3 of 15 sur- vived 32 minutes; at 0.031 ppm, 3 of 15 survived 105 minutes; at 0.034 ppm all fish were dead in 10 hours. The calculated initial concentration of 6.0 ppm had decreased to 1.38 in 15 minutes, and to 0.081 in 105 minutes, and 0.34 ppm in 10 hours. It is interesting that fish held more than a foot below the surface were unharmed.	Jackson (1960)	
B-28	Benzene hexa- chloride (alpha isomer)	Cylindrospermum licheniforme (CI) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl - NT Ma - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)	APPENDIX B
	Benzene hexa- chloride (beta isomer)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Comment same as above except that: CI - PT (7) Ma - PT (7) So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)	

	Benzene hexa- chioride (delta isomer)	Cylindrospermum licheniforme (CI) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Comment same as above except that: CI - NT Ma - PT (7) So - PT (7) Cv - NT Gp - T Np - PT (14)	Palmer and Maloney (1955)
	Benzene hexa- chloride (gamma isomer)	Black fly (larvae) Rainbow trout Caddisfly	FR	Alaskan streams	0.5 (O) 10 (O) 10 (O)	_	The chemical was applied for control of black flies, and because the acetone solution was most effective, only that data is reported here. The figures reported are for minimum effective dosages for black fly larvae and maximum nonlethal dosages for rainbow trout and caddisfly larvae. The value given for black flies was the highest dosage tested and was ineffective.	Gjulian, et al (1949)
	Benzene hexa- chloride (gamma isomer)	Lepomis macrochirus Micropterus salmoides	B F	Auburn, Ala.	0.1 to 2.0 (K) 0.1 to 0.2 (K)	-	Aquarium test. In an earthen pond, 0.18 ppm failed to kill bluegill, bass, golden shiner and several species of minnows.	Lawrence (1950)
	Benzene hexa- chloride (gamma isomer)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella Variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - NT Ma - NT So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
COMMERCIAL CHEMIC	Benzene hexa- chloride (gamma isomer, tech)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Comment same as above except that: CI – NT Ma – PT So – PT (14) Cv – NT Gp – PT Np – NT	Paimer and Maloney (1955)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL PRODUCTS	внс	Black fly (larvae) Rainbow trout Caddisfly	F	Alaskan streams	0.5 (O) 5.0 (O) 0.5 (O)	_	The chemical was applied for control of black flies, and because the acetone solution was most effective, only that data is reported here. The figures reported are for minimum effective dosages, for black fly larvae and maximum non- lethal dosages for rainbow trout and caddisfly larvae. The value given for black flies was the highest dosage tested, and was ineffective.	Gjulian, et al (1949)
RODUCTS	ВНС	Blue crab Marsh fiddler crab Red-jointed fiddler crab Cyprinodon variegatus Leiostomus xanthurus Mugil curema	FE	Bombay Hook Island, Del.	(O) (O) (O) (O) (O)	-	<ul> <li>The location under study was a salt marsh bounded by Delaware Bay.</li> <li>Organisms were confined in cages within the test area.</li> <li>BHC was applied at 0.1 pound per acre. C. variegatus, L. xanthurus, and M. curema showed 35 percent mortality in 7 days.</li> <li>Blue crabs showed 10 percent mortality when exposed for 7 days in streams and 10 percent mortality in ponds.</li> <li>Marsh fiddler crabs and red-jointed fiddler crabs showed mortalities of 80 and 35 percent, respectively, in 7 days.</li> </ul>	George, et al (1957)
R-30	внс	Fathead minnow Bluegill Goldfish Guppy	BSA	-	2.3 (T4A) 0.79 (T4A) 2.3 (T4A) 21.7 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish" It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in soft water.	Tarzwell (1959)
	внс	Fathead minnow	BSA	-	2.0 (T4A)	a	Comment same as above.	Tarzwell (1959)
	внс	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	2.30 (T4A) 0.79 (T4A) 2.3 (T4A) 2.17 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
	внс	Pimephales promelas Lepomis macrochirus	BSA	-	2.3 (T4A) 0.79 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)

	внс	Algae Salmo trutta Invertebrates	FR	Isle of Man, Eng.	_	-	Report notes that the fish were not killed, but may have moved from polluted areas when their normal insect food was no longer available. Tables give percentage composition of fauna at 8 collecting stations, given in yards above and below point of origin.	Hynes (1961)
	внс	_	FR	Flint Creek, Ala.	0.456 (K)	_	Conventional treatment in a water purification plant did not reduce the amount of chemical found in the stream. Data are given for 4 years 1959-1962, with a range of concentra- tions. Only the highest value is reported here. Some fish kill is reported, but species are not identified here. Data for different seasons are reported. The one listed here is for summer 1961.	Nicholson, et al (1964)
	внс	Heteropneustes fossilis	BSA	_	(0)	-	Experiments were conducted in a small battery jar containing 5 liters of water sprayed with 25 cc of BHC (20%). The fish died in 1 hour and 30 min.	Mathur (1964)
	внс	Błuegill	BSA		0.79 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
B-31	внс	Golden shiner <i>Cyclops</i> sp	BSA	-	0.062-1.5 (O) 0.062-0.5 (O)	acdep	The 1.5 ppm value cited is for a 2-day period with the active ingredient added as a wettable powder to water. Threshold values, (LD/O) for the BHC dissolved in a number of solvents were somewhat lower. Some of the solvents caused a 25-fold increase in toxicity of BHC to golden shiner. A $TL_m$ 48 hr of 0.125-0.25 ppm BHC was obtained for <i>Cyclops.</i> Formulations containing oil were more toxic than dust formulations of BHC.	Meyer (1965)
	внс	Tubifex spp Limnodrilus spp	BSA		3.0-15 (L4A)	a c e	Toxicity is reported as the mean lethal dose (LD <sub>50</sub> ) for 24, 48, and 96 hours.	Whitten and Goodnight C (1966)
	внс	Puntius puckelli	BSA	_	3.8 (T4A)	<u>a</u> cdelm	Tap water was used as diluent. Toxicity data are given as TL <sub>m</sub> 's in ppm for 24, 48, 96 hr. The pH of the water averaged 8.3. The study was conducted in India.	Rao, et al (1967)
	внс	Oncorhynchus kisutch	BSA	_	0.2 (T2A)	а	The rate of decay of the gamma isomer of BHC is suspected to be appreciable. The half-life in fresh water would be somewhere in the vicinity of 7-8 days.	Velsen and Alderdice (1967)
COMME	BHC (crude)	Sesarma africanum	BSA	_	65 (K≤1) 6.5 (K1) 0.65 (SB) 0.065 (NTE)		BHC caused complete lack of coordination within 24 hours.	Jordan (1955)
COMMERCIAL CHEMICAL PRODUCTS			FR	Sierra Leone	325	-	In rice fields, sprays with as low as 325 ppm BHC gave adequate protection from crabs to young rice seedlings. Initial results given were derived from contact with aqueous suspensions of varying concentrations in bioassay evaluations.	
EMICAL	BHC (emulsion)	Micropterus salmoides (fry)	BSA	_	0.05 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
. PROD		lctalurus punctatus (fry)			0.2 (SB3)			
UCTS		Lepomis macrochirus (fry)			0.1 (SB3)			

Chemical BHC (45%	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Oyster	BCF	_	1.0 (O)	a	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
gamma isomer) BHC (45% gamma isomer) BHC (tech, 15.5%)	Oyster	BCF	-	0.36 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
BHC (tech,	Pimephales promelas	BSA	-	15 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et al (1959)
15.5%)	Lepomis macrochirus			5.i (T4A)			
	Carassius auratus			15 (T4A)			
	Lebistes reticulatus			14 (T4A			
BHC (WP)	Micropterus salmoides (fry)	BSA	_	0.2 (SB3)	a c d e f	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
	lctalurus punctatus			0.4 (SB3)			
	(fry) Lepomis macrochirus (fry)			0.5 (SB3)			
Bidrin (tech)	Procambarus clarki	BSA	-	3.0 (T3A)	a c d o	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
Bidrin	<i>Pteronarcys californica</i> (naiads)	BSA	-	0.43 (T4A)	<u>a c d e f</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
Bomyl (EC <sub>4</sub> , GC-3707)	Gambusia affinis	FL	Cal.	(O)	-	At an application rate of 2.0 lb/acre, 66% mortality of the fish occurred in 24 hours.	Mulla (1966)
Borate	Salmo gairdnerii	BSA	-	2300 (T1A) 2050 (T2A)	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
BP 1002	Pandalus montagui	BSA	-	5.8 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C.	Portmann and Connor
	Crangon crangon			5.8 (T2A)		It was shown that the toxicity of this solvent emulsifier decreased with time due to evaporation of the solvent.	(1968)
	Carcinus maenas			15.0 (T2A)		BP 1002 at a concentration of 3.3 ppm killed 100% of Crangon crangon larvae in 3 hr; at 10 ppm it killed 95%	
	Cardium edule			81.0 (T2A)		of Carcinus maenas in 3 hr.	
Buramine	Semotilus atrom <del>a</del> culatus	BSA	-	1,000 to 1,500 (CR)	<u>a</u> e	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that concentra- tion in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)

	C-56	Lepomis macrochirus Micropterus salmoides	BSA	-	30 (T2A) 35 (T2A)	<u>a</u> co	The response of bluegill and bass fingerlings to nine agricul- tural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
	C-2059	Lepomis macrochirus	BSA	_	55 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
	C-2059	Lepomis macrochirus	BSA	-	90 (T1A)	<u>a</u> b e	Comment same as above.	Hughes and Davis (1967)
	C-8514	Oncorhynchus kisutch	BSA	_	21.5 (T2A)	а	Physical instability of this formulation would suggest that toxicity in the aquatic environment could be a problem if the preparation is to be used in or near water courses under field conditions.	Velsen and Alderdice (1967)
	Camphene (chlorinated)	Silverling minnows Spotfin shiner Creek chub Fall fish Blacknosed dace	BSA	-	0.04 (K)	-	The "Threshold" for bluegills was 0.01 ppm. A field study in W. Va. is also described.	Linduska and Surber (1948)
2	Captan (N-trichloro- methylthio-4- cyclohexene-1,2- dicarboxyimide)	Brachydanio rerio (larvae)	BSA	_	30 (T1A) 1.0 (O) (T 70 min)	<u>a</u>	TL <sub>m</sub> was 70 min for larvae. Ninety-eight percent of the larvae died in 90 min.	Dawood and Dazo (1966)
	Carbaryl	Procambarus clarkii	FO	Crowley La.	(O)	c d e p	Experiment was conducted in a flooded rice field Area was divided into 4 blocks with a fence restricting crawfish to the desired area. The rearing of crawfish in rice fields is of considerable commercial importance in Louisiana. It is fortunate that the chemicals discussed in this report had no untoward effect at the levels used. The chemical was applied at the rate of 0.8 lbs/acre.	Hendrick, et al (1966)
COMMERCIAL	Carbaryl	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 47 percent.	Jamnback and Frempong- Boadu (1966)
CHEMICAL	Carbaryl	Australorbis glabratus	BSA and FL	Puerto Rico	(0)	с	Seven of the tested compounds failed to meet acceptability criteria-that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective. All others failed.	Seiffer and Schoof (1967)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Carbaryl	Pteronarcys californica (naiads)	BSA	_	0.0048 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
	Pteronarcella badia (naiads)			0.0017 (T4A)			
	Claasenia sabulosa (naiads)			0.0056 (T4A)			
Carbo- phenothion (Trithion)	Culex pipiens quadrimaculatus	BSA	-	(0)	с	Tests were conducted in tap water and artifically polluted tap water. The values reported are the concentration range for an LC <sub>90</sub> , 0.085 to 0.280 ppm for polluted, and 0.017 to 0.034 ppm for tap water.	Lewallen and Wilder (1963)
Carbo- phenothion (EC4)	Gambusia affinis	FL	Ponds in 111.	(O)	-	When applied at 0.2 and 1.6 pounds per acre active ingredient, 98 and 100 percent fish mortality occurred respectively in 1 day.	Mulla, et al (1963)
	Rana catesbeiana					When applied at 0.4 pound per acre, 100 percent bullfrog mortality occurred in 1 day.	
Carbo- phenothion (EC4)	Micropterus salmoides	BSA	-	1.0 (O)	a e	At 1.0 ppm, 10 percent mortality occurred in 1 day. Experi- ments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.	Mulla, et al (1967)
Casoron	Redear sunfish	BSA	-	(O)	а	No mortality was noted in fish weighing 3 g with concentra- tions of 20,000 mg/1 at 48 hr.	Соре (1963)
Casoron	<i>Pteronarcys</i> sp (nymphs)	BSA	-	6.6 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
Casoron	Simocephalus serrulatus	BSA	_	5.8 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr.	Sanders and Cope
	Daphnia pulex			3.7 (SB)		Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	(1966)
Casoron	Rainbow trout	BSA	_	22 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
Casoron	Bluegill Rainbow	BSA	_	20 (T2A) 18 (T4A)	a	This is an estimated LC <sub>50</sub> value at temperatures from 55 to	Соре
(WP)	trout Bluegill			10 (T4A)	_	75 F.	(1965)
Catechol	Microcystis aeruginosa	L	-	100 (K)	<u>a</u> , etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et a (1952)
Ceresan M	Channel catfish (fingerlings)	BSA	-	1.8 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)

	Chem Ban	Channel catfish (fingerlings)	BSA	-	26 (K1A)	<u>a</u>	Comment same as above.	Clemens and Sneed (1959)
	Chem Fish Special	Channel catfish (fingerlings)	BSA	_	0.56 (K1A)	<u>a</u>	Comment same as above.	Clemens and Sneed (1959)
	Chem Mite	Channel catfish (fingerlings)	BSA	-	1.29 (K1A)	<u>a</u>	Comment same as above.	Clemens and Sneed (1959)
	Chem Sen 56	Channel catfish (fingerlings)	BSA	-	97.7 (K 30 hr A)	<u>a</u>	Comment same as above.	Clemens and Sneed (1959)
	Chemagro	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	0.032 (O)	а	Water temperature was 21 C. The figure reported is a 48-hr EC <sub>50</sub> .	Butler (1965)
	Chemagro 4497	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	0.032 (O)	а	Comment same as above.	Butler (1965)
	Chłorax	Channel catfish (fingerlings)	BSA	_	3157 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
B-35	Chlordane	Bluegill	F	_	(O)	-	At 1 lb per acre, 87 percent of the bluegill sunfish were killed. At 0.5 lb per acre most of the bluegills as well as other species survived.	Linduska and Surber (1948)
	Chlordane	Black fly (larvae) Rainbow trout Caddisfly	FR	Alaskan streams	0.5 (O) 20 (O) 10 (O)	-	The chemical was applied for control of black flies, and because the acetone solution was most effective, only that data is reported here. The figures reported are for mini- mum effective dosages of black fly larvae and maximum nonlethal dosages for rainbow trout and caddisfly larvae. The value given for black flies was the highest dosage tested. The value given for trout was also the highest tested.	Gjulian, et al (1949)
	Chlordane	Lepomis macrochirus	BSA	Auburn, Ala.	0.1 (NTE)	-	At this concentration, there was no apparent effect.	Lawrence (1950)
COMMERCIAL		Carassius auratus Micropterus salmoides			0.2 (K)		At this concentration, bluegill and bass were killed, but goldfish survived.	
RCIAL CHEMICAL PRODUCTS	Chiordane	Fathead minnow Bluegill Goldfish Guppy	BSA	_	0.052 (T4A) 0.022 (T4A) 0.082 (T4A) 0.19 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compound. This experiment was performed in soft water.	Tarzwell (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chlordane	Fathead minnow	BSA		0.069 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish" It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. The experiment was performed in hard water.	Tarzwell (1959)
Chlordane	Channel catfish (fingerlings)	BSA	-	0.74 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Chlordane (100%)	Pimephales promelas Lepomis	BSA	-	0.069 (T4A) 0.022 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et a (1959)
	macrochirus Carassius auratus			0.082 (T4A)			
	Lebistes reticulatus			0.19 (T4A)			
Chlordane	Pimephales promelas Lepomis	BSA	-	0.05 (T4A) 0.01 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et a (1959)
	macrochirus Carassius			0.08 (T4A)			
	auratus Lebistes reticulatus			0.19 (T4A)			
Chlordane (75%)	Pimephales promelas	BSA	-	0.18 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et a (1959)
Chlordane	Pimephales promelas	BSA	-	0.052 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as	Tarzwell (1959)
	Lepomis macrochirus			0.022 (T4A)		toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	
Chlordane	Oncorhynchus kisutch	BSA	-	56 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
	Oncorhynchus tshawytscha			57 (T4A)			(1961)
	Salmo gairdneríi Gasterosteus			44 (T4A) 90 (T4A)			
	Gasterosteus acu/eatus			JU (14A)			

	Chlordane	Gammarus Iacustris Iacustris	BSA	_	(O)	a e p	The mortality might have been partially due to the susceptibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Results were expressed as LT <sub>50</sub> ; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 235 ( $\pm$ 35) min.	McDonald (1962)
	Chlordane (EC 7.5)	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a C	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and resulted in a 70 percent mortality for the fish, and a 0 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
	Chlordane	Bluegill	BSA	-	0.022 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
	Chlordane	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.015 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
	Chlordane (tech)	Rainbow trout Bluegill	BSA	-	0.0078 (T4A) 0.040 (T4A)	_	The values reported are given as $LC_{50}$ .	Cope (1965)
	Chlordane	Simocephalus serrulatus Daphnia pulex	BSA	_	0.020 (SB) 0.029 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Chlordane (75% emulsion)	Green sunfish	BSA	_	1.0 (NTE)	-	Fish were repelled by concentrations of 5, 10, and 20 ppm. No lethal effects were noted at concentrations less than 5 ppm.	Summerfelt and Lewis (1967)
COM	Chlordane	Vascular plants Algae Chubs Largemouth bass Clams	FL	Tule Lake, Ore.	(O)	-	The amount of chemical applied as a spray was not specified. Plants contained 1.5 to 6.0 ppb. Algae contained 1.7 to 50.0 ppb. Chubs were analyzed to show a content of 8.0 to 24.0 ppb. Bass contained 7.5 to 43.0 ppb. Clams contained 2.0 to 25.0 ppb. The water contained 0.01 to 0.51 ppb.	Godsil and Johnson (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chlordane	Labeo rohita Chamna punctatus Mastocembelus pancalus Trichogaster fasciatus Mystus vittatus Nandus Nandus Puntius sophore Heteropneustes fossilis Amphipnous cuchia Phytoplankton: Volvox Pando	BSA	-	0.0000709 (T7A) 0.0001 (K 40 hr) 0.001 (K 115 hr) 0.0001 (T7) 0.0008 (T7) 0.0008 (T7) 0.0016 (K 51 hr) 0.0002 (K 130 hr) 0.0002 (K 10 hr) 0.0025 (K 25 hr) 0.0008 (T7) 0.005 (K 25 hr) 0.0008 (T7) 0.005 (K 51 hr) 0.001 (T7) 0.005 (K 45 hr) 0.01 (T7) 0.001 (T7) 0.001 (T7) 0.001 to 0.50 (K7)	a e	Chlordane was found to be highly toxic but not selective to fish. Its prolonged toxicity makes it unsuitable for fishery management and it is recommended that chlordane not be used in fields adjacent to fishery reservoirs.	Konar (1968)
	Zooplankton: Cyclops Nauplius Daphnia Cypris	Ceriodaphnia Diaptomus Gastrotrica Brachionus		0.10 to 0.50 (K7)		<i>Gastrotrica</i> were not affected and <i>Brachionus</i> was not killed at 0.10 ppm.	
Chlordane	Pteronarcys californica (naiads)	BSA	-	0.015 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Chlorea (granular)	Althernanthera philoxeroides Najas quadalupensis Spatterdock	FL	Fla.	(O)	-	The degree of control was as follows:A. philoxeroides(393 lb/acre) - 30 percentN. quadalupensis(393 lb/acre) - nonespatterdock(454 lb/acre) - 2 percent	Copeland and Woods (1959)

Chloretone	<i>Bræchydenio</i> <i>rerio</i> (fertilized eggs, 2 hr) (fertilized eggs, 24 hr)	BSA	_	100* (K 37 min) 50* (K 3-1/4 hr) 10* (K 5 hr) 5* (SB 1-1/2 hr) 1* (NTE) 100* (K 37 min) 50* (K 7-1/2 hr) 10* (K 119 hr) 5* (SB 1-1/2 hr) 1* (NTE) *% of saturated	<u>a</u>	Saturated solutions and dilutions of saturated solutions were used. 4.5 g of the chemical saturated 500 ml of water at RT. Percent dilutions used were 100, 50, 10, 5.0, 4.0, 3.0, 2.0, and 1.0. Immobilization times are given. Histological observations were also made.	Blumenkrantz (1956)
Chlorophenyl	Salmo gairdnerii	BSA	-	975 (T1A) 925 (T2A)	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
Chlorothion	Pimephales promelas	BSA	-	3.3 (T4A)	acdef	Tests were performed in both hard and soft water. Additional tolerance limit values are given.	Henderson and Pickering (1958)
Chlorothion	Fathead minnow	BSA	-	3.2 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959) )
Chlorothion	Pimephales promelas	BSA	_	3.2 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)
Chlorothion	Pimephales promelas	BSA	-	3.2 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
Chlorothion	Lepomis macrochirus Micropterus salmoides Notemigonus crysoleucas Carassius auratus	BSA	-	0.1 (O) 0.1 (O) 0.1 (O) 0.1 (O)	<u>acdf</u>	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The inhibition of the enzyme was a function of the concentra- tion of the insecticide, extent of exposure, and specific chemical nature of the compound.	Weiss (1959)
	Chlorophenyl Chlorothion Chlorothion Chlorothion	rerio (fertilized eggs, 2 hr)Chloropheny!Salmo gairdneriiChloropheny!Salmo gairdneriiChlorothionPimephales promelasChlorothionFathead minnowChlorothionPimephales promelasChlorothionPimephales promelasChlorothionPimephales promelas	rerio (fertilized eggs, 2 hr)DOA(fertilized eggs, 24 hr)(fertilized eggs, 24 hr)ChlorophenylSalmo gairdneriiBSAChlorothionPimephales promelasBSAChlorothionFathead minnowBSAChlorothionPimephales promelasBSAChlorothionPimephales promelasBSAChlorothionPimephales promelasBSAChlorothionPimephales promelasBSAChlorothionPimephales promelasBSA	rerio       (fertilized eggs, 2 hr)         (fertilized eggs, 24 hr)         (fertilized eggs, 24 hr)         Chlorophenyl       Salmo gairdnerii         Chlorophenyl       Pimephales promelas         BSA       -         Chlorophenyl       Pimephales promelas         BSA       -	TerioDefChTerio(fertilized50°eggs, 2 hr)(K 37 min)(fertilized(K 5 hr)(fertilized10°eggs, 24 hr)(K 37 min)50°(K 37 min)50°(K 37 min)60°(K 37 min)50°(K 37 min)51°(K 37 min)52°(K 37 min)53°(K 37 min)54°(K 37 min)55°(K 37 min)60°(K 37 min)56°(K 37 min)57°(K 37 min)58(K 37 min)59(K 37 min)50°(K 37 min)	rerio       Exh       I       IX 37 min)       IX 37 mi	improvement       improvement

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chlorothion	Pimephales promelas	BSA	_	0.5 (O)	_	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. The $LT_{50}$ for the fathead was 72 hr.	Weiss (1961)
Chlorothion (tech,	Pimephales promelas	BSA	-	2.8 (T4A)	<u>a</u> c d <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent	Pickering, et al (1962)
98 percent)	Lepomis macrochirus			0.71 (T4A)		or carrier in most cases.	(1552)
	Carassius			2.3 (T4A)			
	auratus Lebistes reticulatus			1.2 (T4A)			
Chlorothion	Carassius auratus Lepomis macrochirus Notemigonus crysoleucus	BSCH	_	1.0 (O)* * no response, 15 days	<u>a</u> c d <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
Chlorothion	Bluegill	BSA		0.7 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Chloroxuron (granular)	Lepomis macrochirus	BSA	-	60 (T1A)	<u>a</u> b e	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
Chloroxuron (WP)	Lepomis macrochirus	BSA	_	25 (T1A)	<u>a</u> b e	Comment same as above.	Hughes and Davis (1967)
CIPC	Lepomis macrochirus	BSA	-	12 (T2A)	<u>a</u> co	This response of bluegill and bass fingerlings to nine agricultural chemicals as determined by bioassay using	Davis and
	Micropterus salmoides			10 (T2A)		river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Hardcastle (1959)
Cleanosol	Pandalus montagni	BSA	-	32 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of of seawater at 25 C.	Herbert, et al (1965)
	Crangon			44 (T2A)		It was shown that the toxicity of this solvent emulsifier	(1905)
	crangon Carcinus			102 (T2A)		decreased with time, due to evaporation of the solvent. Cleansol at a concentration of 33.3 ppm killed 100% of	
	maenas Cardium edule			19.2 (T2A)		<i>Crangon crangon</i> larvae in 3 hr.	

	Clostridium botulinum (Type A toxin)	Pimephales promelas	BSA	-	(O)	<u>a</u> cde	Fish survived high does rates of 102,000 mouse LD $_{50}$ /ml for 24 hr and 17,000 mouse LD $_{50}$ /ml for 96 hr. Fish cannot be used to detect this chemical at levels critical for man.	Pickering and Henderson (1959)
	СМU	Algae (Mixed culture) Lepomis macrochirus Lepomis cyanellus Pomoxis nigromaculatus Hyborhyncus notatus Lebistes spp	LBSA	-	0.5-1.0 (K) (O)	<u>a e</u> _	Tests with 22 species of algae indicated that 0.5 to 1.0 ppm CMU prevented growth. No adverse effects on the fish were found in 23 days with concentrations of 10, 20, and 40 ppm.	Fitzgerald (1958)
	CMU	Channel catfish (fingerlings)	BSA	_	75.9 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Conco LCP-12	Daphnia magna	BSA		694 (T1A) 290 (T2A) 204 (T3A)	е	When emulsifier was mixed with crude oil, the TL <sub>m</sub> value was one-half the values cited.	Dowden (1962)
j	Co-ral	Pimephales promelas Lepomis macrochirus	BSA	-	≥18 (T4A) 0.18 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
	Co-ral	Pimephales promelas Lepomis macrochirus	BSA	-	≥18 (T4A) 0.18 (T4A)	<u>a</u>	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)
COMME	Co-Ral	Oncorhynchus kisutch Salmo gairdnerii Gasterosteus aculeatus	BSA	_	15,000 (T4A) 1,500 (T4A) 1,862 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
RCIAL CHEMICAL P	Co-Ral	Micropterus salmoides Pimephales promelas	BSA	-	0.5 (O) 0.5 (O)	-	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the $LT_{50}$ was 36 hr and for the fathead 72 hr.	Weiss (1961)

CO Chemical Co-Ral	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Carassius auratus Lebistes reticulatus	BSA	_	18 (T4A) 0.56 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TLm's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al {1962)
Co-Ral	Chaoborus astictopus	BSA	-	0.39 (T1A)	<u>a</u>	Toxicity value given is for the fourth instar larvae.	Hazeltine (1963)
CP (tech, 97.5 MC percent) Co-Ral PP Co-Ral Co-Ral	Carassius auratus Lepomis macrochirus Notemigonus crysoleucus	BSCH	_	1.0 (0)* 1.0? (0)** 1.0 (0)* * no response, 15 days **response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
CO-RAL	Cyp <i>rinodon</i> variegatus (juvenile)	BSA	-	0.28 (O)	а	Water temperature was 12 C. The figure reported is a 48 <del>-</del> hr EC <sub>50</sub> .	Butler (1965)
Co Ral	Salmo gairdenerii Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	_	0.55 (T2A) 0.73 (T2A) 0.8 (T2A) 4.0 (T2A) 6.8 (T1A) 8.0 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.1.) were also determined.	Willford (1966)
Co-ral (97,5% active in acetone)	<i>Hexagenia</i> sp <i>Hydropsyche</i> sp (Iarva) Bluegill	BSA	_	0.43 (T1A) 0.005 (T1A) 1.4 (T1A)	a e	Dissolved oxygen was measured before and after assay. Assays were conducted in Mississippi River water.	Carlson (1966)

Čube powder	Cyprinus carpio	BSA	_	0.115 (L3)	acdei	Such variables as temperature, species, and size of fish were studied. Toxicity is expressed as $LD_{50}$ for 72 hr.	Hester (1959)
(7.3% rotenone)	Micropterus salmoides			0.164 (L3)		Smaller concentrations of rotenone were required when used in conjunction with sulfoxide. The data shown are	
	Pimephales promelas			0.200 (L3)		for 70 F. The chemical was considerably more toxic at this temperature, than at 40 F for all fish species.	
	Carassius auratus			0.218 (L3)			
	Lepomis macrochirus			0.268 (L3)			
	L. cyanellus			0.246 (L3)			
	Notemigonus			0.620 (L3)			
	crysoleucas						
	Ictalurus			0.346 (L3)			
	nebulosus						
	marmoratus						
Cube root	Pimephales promelas	BSA	-	0.066 (T4A)	acdfg	Test water was spring water diluted with distilled water. Removal of toxic chemicals by carbon adsorption, chlorine and chlorine dioxide treatment, and alum coagulation was studied. The most effective method to remove fish poisons was by use of activated charcoal adsorption.	Cohen, et al (1961)
Cumate	Pimephales promelas	BSA	-	0.071 (T4A)	acdef	The toxicity of this substance was influenced by the quality of the water (pH, hardness, alkalinity). The	Pickering and Henderson
	Lepomis macrochirus			0.32 (T4A)		chemical was more toxic in soft water.	(1966)
Cuprose	Chlorella	L	_	20 (AS 1)	_	Describes a bioassay method to differentiate between an	Fitzgerald and
	pyrenoidosa	_				algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Faust (1963)
2, 4-D	Killifish (minnow)	BSA	-	2000 (O)	<u>a e</u>	Temperature was held at 20-25 C, and the water was aerated by circulating water pumps. Data reported as	Harrison and Rees
	Eupomotis gibbosus			1000 (O)		deaths in 7 days. Upper safe limit concentrations were established.	(1946)
	Ameiurus nebulosus			2000 (O)			
2, 4-D	Eichornia crassipes	FR	Fla.	(O)	_	Control of water hyacinth was affected with 1:1140 dilution applied to 100-150 sq ft plots. Author notes that no adverse effects to the water fauna (fingerling fish, etc.) were observed up to the time of disappearance of the water hyacinth. The roots were not killed, but did not readily produce shoots. Addition of carbowax as a wetting atent did not improve herbicidal action.	Hildebrand (1946)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
2.4-D	Eichhornia crassipes Alternanthera philoxeroides	FRLO	Mississippi R., Delta, La.	(O)	_	2,4-D was applied over weed-infested areas of the following types: borrow pits, drainage ditches, a shallow lake, a small tributary of the Mississippi River, and land areas. Water hyacinth was killed and sank under any condition in which it grows in South Louisiana by application of 8 Ib/acre (free acid equivalent) of the amine salt of 2,4-D. A single application of 2,4-D at 8 Ib/acre did not give complete elimination of alligator weed from borrow pits and deep ditches but it did reduce the popula- tion considerably.	Eggler (1953)
2,4-D	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl = PT (3) Ma = NT So = NT Cv = NT Gp = NT Np = T (3)	Palmer and Maloney (1955)
2,4-D	Salmo gairdnerii	BSA	-	2300 (T1A) 2050 (T2A)	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
2,4-D	Salmo gairdnerii	BSA	-	4.4 (T1A) 3.3 (T2A)	a e	Comment same as above.	Alabaster (1956)
2,4-D	Salmo gairdnerii	BSA	_	3.0 (T1A) 2.2 (T2A)	<u>a</u> e	Comment same as above.	Alabaster (1956)
2,4-D	Lepomis macrochirus Micropterus salmoides	BSA	-	375 (T2A) 350 (T2A)	<u>a</u> c o	The response of bluegill and bass fingerlings to nine agri- cultural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
2,4-D (granular)	Nympheae sp Myriophyllum brasiliense Myriophyllum heterphyllum Brasenia schreberi Utricularia sp	FL	Farm ponds in Ga.	(O)	-	Granular 2,4-D controlled Nympheae sp., Myriophyllum heterphyllum, Brasenia schreberi, and Utricularia sp. at the rate of 100 lb/acre (20 lb acid).	Thomaston, et a (1959)
2,4-D/ 2,4-5T + TCA	Althernanthera philoxeroides	FL	Fla.	(0)	-	At 19.2 pounds per acre, only 1-2 percent control of alligator weed was obtained.	Copeland and Woods (1959)

	2,4-D/ 2,4-5T	Althernanthera philoxeroides Typha I Iatifolia Spatterdock	FL	Fla.	(O)	_	The degree of control was as follows:A. philoxeroides(5.0 lb/acre) - 1-2 percentT. latifolia(10.0 lb/acre) - 80 percentSpatterdock(5.0 lb/acre) - 5 percent	Copeland and Woods (1959)
	2,4-D (pellets)	<i>Najas quadalupensis</i> Spatterdock	FL	Fla.	(0)	-	At 80.0 and 43.6 lb/acre, only 2 percent control of both species was obtained.	Copeland and Woods (1959)
	2,4-D + TCA	Panicum hemitomum	FL	Fla.	(0)	-	At 90.0 lb/acre, 75 percent control of <i>P. hemitomum</i> was obtained.	Copeland and Woods (1959)
	2,4-D (ester)	Althernanthera philoxeroides Pontederia cordata Spatterdock	FL	Fla.	(O)	_	At 4.2 lb/acre, the degree of control was as follows: A. philoxeroides - 2 percent P. cordata - 85 percent Spatterdock - 3 percent	Copeland and Woods (1959)
	2,4-D (pellets)	Bushy pondweed	FL	Lakes in Fla.	(O)	_	Concentrations of 1.5 to 2.5 ppm controlled the bushy pondweed.	Phillippy (1961)
	2,4-D (butoxyethanol ester)	Lepomis macrochirus	BSA	-	2.1 (T2A)	a e	The various salts of the chemicals showed wide variations in toxicity.	Hughes and Davis (1963)
,	2,4-D (acid, with emulsifiers)	Lepomis macrochirus	BSA	-	8.0 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
	2,4-D (jsopropyl ester)	Lepomis macrochirus	BSA	_	0.8 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
	2,4-D (propylene glycol butylether ester)	Lepomis macrochirus	BSA	_	2.1 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
0	2,4-D (ethylester)	Lepomis macrochirus	BSA	_	1.4 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
COMMERCIAL	2,4-D (di-n,n-dimethyl- cocoamine ester)	Lepomis macrochirus	BSA	-	1.5 (T2A)	a e	The various salts of the chemicals showed wide variations in toxicity.	Hughes and Davis (1963)
	2,4-D (isoocty) ester)	Lepomis macrochirus	BSA	-	36 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
CHEMICAL PR	2,4-D {mixed butyl and isopropyl esters)	Lepomis macrochirus	BSA	-	1.5 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen <b>ts</b>	Reference (Year)
AL CHEMICAL	2,4-D (dimethylamine ester)	Lepomis macrochirus	BSA	_	416 (T2A)	a e	The various salts of the chemicals showed wide variations in toxicity.	Hughes and Davis (1963)
AICAL PRODUCTS	2,4-D (alkanolamine, ethanol and isopropanol series)	Lepomis macrochirus	BSA	-	580 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
UCTS	2,4-D (oleic -1,3-propylene diamine)	Lepomis macrochirus	BSA	-	4.0 (T1A)	_	The bioassay methods employed in this experiment were not given in the paper but it was stated that the same procedures were employed as in previous work.	Davis and Hughes (1963)
	2,4-D (butyl ester, oil soluble)	Lepomis macrochirus	BSA	-	4.9 (T1A)	-	Comment same as above.	Davis and Hughes (1963)
	2,4-D (butyl ester)	Lepomis macrochirus	BSA	-	10 (T1A)	_	Comment same as above.	Davis and Hughes (1963)
	2,4-D (butyl ester)	Lepomis macrochirus	BSA	_	1.3 (T2A)	a e	The various salts of the chemicals showed wide variations in toxicity.	Davis and Hughes (1963)
<b>B-4</b> 6	2,4-D {acid}	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	2.0 (NTE) 2.0 (K10%) 50 (NTE) –	-	<ul> <li>Sea water was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
	2,4-D (propylene glycol butyl ether ester)	Crassostrea virginica Penaeus duorarum Leiostomus xanthurus Fundulus similis Mugil cephalus Cyprinodon variegatus Phytoplankton	BCFA & BSA	_	1.0 (0,39%) 1.0 (NTE) 4.5 (T4A) 44% (O)	-	Comment same as above.	Butler (1965)

	2,4-D (2-ethyl hexyl ester)	Crassostrea virginica	BCFA & BSA	-	5.0 (0,38%)	-	Comment same as above.	Butler (1965)
	,	Penaeus aztecus			2.0 (0,10%)			
		<i>Mugil cephalus cephalus</i> Phytoplankton			10 (NTE) 49% (O)			
	2,4-D (iso-octył ester)	Lepomis macrochirus	BSA	_	49% (0) (L) 8.8-59.7 (T2A) (G) 116-1000 (T2A)	acdeg	Toxicity data for 24 and 48 hours are presented for liquid (L) and granular (G) formulations. Various commercial formula- tions were tested. The liquid formulations were almost invariably more toxic than the granular ones.	Hughes and Davis (1965)
	2,4-D (propy- lene glycol butyl ether ester)	Lepomis macrochirus	BSA	_	(L) 2.1 (T2A) (G) 9.3 (T2A)	<u>a</u> cdeg	Comment same as above.	Hughes and Davis (1965)
	2,4-D (butoxy ethanol ester)	Lepomis macrochirus	BSA	-	(L) 2.1 (T2A) (G) 34.5 (T2A)	<u>a</u> cdeg	Comment same as above.	Hughes and Davis (1965)
,	2,4-D (butoxy ethanol ester)	Crassostrea virginica Penaeus duorarum Fundulus similis	BCFA & BSA	-	3.75 (O) 1.0 (NTE) 5.0 (T2A)		Sea water was pumped continuously into test aquaria. Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following: Oyster – 96-hr EC50 – Conc. which decreased shell growth.	Butler (1965)
i		Phytoplankton			16% (O)		Shrimp – 48-hr EC <sub>50</sub> – Conc. which killed or paralyzed 50% of test animals. Fish – 48-hr EC <sub>50</sub> – Conc. which killed 50%. Phytoplankton – Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	
	2,4-D (dimethyl- amino salt)	Crassostrea virginica Penaeus aztecus Fundulus	BCFA & BSA	-	2.0 (NTE) 2.0 (0,10%) 15 (T2A)	-	Comment same as above.	Butler (1965)
Š		<i>similis</i> Phytoplankton						
-	2,4-D (butoxy ethanol ester)	Crassostrea virginica	BCFA & BSA	-	2.0 (O)	-	Comment same as above.	Butler (1965)
MERCIAL		Pleurobena cordatum			5.0 (NTE)			
	2,4-D	Daphnia magna	BSA		>100 (0)	acdiq	Toxicity, in terms of median immobilization concentration (IC <sub>50</sub> ), is presented.	Crosby and Tucker (1966)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Referenc (Year)
2,4-D	Salmo gairdneri	BSA	-	1.1 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on repro-	Cope (1966)
	Lepomis macrochirus			0.9 (T2A)		duction and behavior are also discussed. Data presented as EC50.	
	Pteronarcy s californica			1.8 (T2A)			
	Daphnia pulex Simocephalus serrulatus			3.2 (T2A) 4.9 (T2A)			
2,4-D	Lepomis macrochirus Elliptis crassidens	FO	Tenn. and Ala.	(0)	_	There was little uptake of 2,4-D (treatment was with a 20% granular material at the rate of 100 lb. of 2,4-D acid equivalent per acre) by fish but some by mussels.	Smith and Isom (1967)
2,4-D (dimethylamine)	Lepomis macrochirus	BSA	-	188 (T1A)	a b e	This report is a simple and straightforward determination of a median tolerable limit for a selected group of herbicides.	Hughes and Davis (1967)
2,4-D (isooctylester)	Lepomis macrochirus	BSA	_	453 (T1A)	a b e	Comment same as above.	Hughes an Davis (1 <del>9</del> 67)
2,4-D (butoxy- ethanol ester)	Pimephales promelas	BCFCH	-	0.2-1.5 (O)	a c d e q	Carbon-filtered tap water was used as diluent. Growth and reproduction were not affected by 2,4-D at range of concentrations indicated. No mortalities occurred.	Mount and Stephan (1967)
2,4-D (esters)	Lepomis macrochirus (eggs) L. cyanellus (eggs) Micropterus	L	-	50 (S), 10 (NTE), 5.0 (NTE) 10/4 (O) 10/5 (O)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)
dolomieui (e Erimyzon sucetta L. macrochiru	Erimyzon			5.0 (NTE) 50 (S)			
2,4-D (dimethyl- amine salt)	Lepomis macrochirus (eggs)	L	-	25 (NTE)	-	Comment same as above.	Hiltibran (1967)
amine sait)	L. cyanellus			25 (NTE)			
	(eggs) Micropterus dolomieui (eggs)			25 (NTE)			
	Erimyzon sucetta (eggs)			25 (NTE)			
	L. macrochirus (fry)			40 (S)			

• ••••

2,4-D (buto ethar ester)	oxy- Iol	Pimephales promelas	BSA & CH	-	5.6 (T4A)	acdef	The fish could tolerate 1/19 this amount of 2,4-D for a ten-month test.	Mount and Stephan (1967)
2.4-D (Na s	alt)	Lepomis macrochirus (fry)	L	-	100 (S)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)
2,4-D (ester	rs)	Lepomis macrochirus (eggs) L. cyanellus (eggs Micropterus dolomieui (eggs) L. macrochirus (fry)	L	-	4/2 (O) 4.0 (NTE) 4.0 (NTE) 3.0 (S)	-	Comment same as above.	Hiltibran (1967)
2,4-D		Lepomis macrochirus (eggs)	L	-	5/1 (O)	-	Comment same as above.	Hiltibran (1967)
2,4-D		Rainbow trout Bluegill	BSA	-	1.1 (T2A) 3.7 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
2,4-D (PGE ester		Lepomis macrochirus (eggs) L. cyanellus (eggs) Micropterus dolomieui (eggs) Erimyzon sucetta (eggs) L. macrochirus (fry)	L	_	1/2 (O) 1/5 (O) 1/5 (O) 1/5 (O) 2 (S)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)
COMMERCIAL CHEMICAL PRODUC	oxy- nol	Anopheles quadri- maculatus (4th instar) Lepomis macrochirus Elliptis crassidens Hexagenia Tendipedidae Heleidae Chaoborus Oligochaeta Corbicula and others	BSA FS	Watts Bar Reservoir T.V.A. Gunters- ville Reservoir T.V.A.	(O)	_	The Watts Bar test site was treated with a 20% granular material at the rate of 100 lb of 2,4-D acid equivalent per acre. The Guntersville area was treated at the rate of 40 lb per acre. The applications were made for control of Eurasian watermilfoil, <i>Myriophyllum spicatum</i> . The toxic effect of 2,4-D was evaluated by sampling the benthic invertebrate communities of both reservoirs before treatment and at least twice after treatment. Residue analysis of water, fish, plants, mussels, and sediment were used to study diffu- sion, accumulation, translocation, and/or degradation of 2,4-D. In both areas at both concentrations, a monitoring device showed some movement of lake fish out of the treated area, but no mortality of fish occurred. A total of 50 assorted frozen samples of plants, animal tissue, and mud were analyzed for 2,4-D. Application of 2,4-D at the given concentrations caused no measurable toxic effect on benthic fauna.	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
2,4-D	Pteronarcys californica (naiads)	BSA	_	0.015 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
2,4-D (butoxy ethanol ester)	Pteronarcys californica (naiads)	BSA	-	0.0016 (T4A)	<u>acdef</u>	Comment same as above.	Sanders and Cope (1968)
2,4-DA (pellets)	Water lettuce	FL	Lakes in Fla.	(0)	_	An application rate of 10 lb/acre controlled water lettuce.	Phillippy (1961)
DAC, dodecylacetamido dimethyl benzyl ammonium chloride	Pimephales promelas	BSA	_	0.65 (T4A)	acdef	Toxicity to 30 species of algae also presented. DAC was algicidal in the range 0,25 to 2.0 ppm.	Maloney and Palmer (1956)
Dacthal	Lepomis macrochirus	BSA	_	1000 (T1A)	<u>a</u> b e	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
Dacthal	Crassostrea virginica Penaeus aztecus Cyprinodon variegatus Phytoplankton	BCFA & BSA	_	0.25 (O) 1.0 (NTE) 1.0 (NTE) 37% (O)	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC50 – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC50 – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC50 – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Dalapon (sodium salt)	Fundulus similis (juvenile)	BSA	-	(0)	а	Water temperature was 20 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Oyster	BCF				No effect on exposure to the chemical at 1.0 ppm.	
	Penaeus aztecus	L				Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality. A concentration of 1.0 ppm caused 30 percent mortality.	

	Dalapon (sodium salt)	Crassostrea virginica Penaeus aztecus Fundulus similis Phytoplankton	BCFA & BSA	-	1.0 (NTE) 1.0 (0,40%) 1.0 (NTE) (O)	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
	Dalapon (sodium salt)	<i>Pteronarcys</i> sp (nymphs)	BSA	-	>1000 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
	Dalapon (Na salt)	Pteronarcys californica (naiads)	BSA	_	100 (NTE)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
_	Dalapon	Cattails and other aquatic plants	FL	Farm ponds in Ga.	(O)	-	Dalapon was used primarily to control marginal grasses and cattails. Cattails can be eradicated at the rate of 1 lb to 5 gal of water or 20 lb/acre. It was indicated that 1 lb to 7.5 gal (15 lb per acre) proved satisfactory for control of cattails.	Thomaston, et al (1959)
B-51	Dalapon (Radapon)	Lepomis macrochirus Pimephales promelas	BSA	-	(S) 440 (T4A) (S) 390 (T4A) (H) 290 (T4A)	ace	Bioassay method in Standard Methods for Examination of Water was used. Both hard (H) and soft (S) water were used. TL <sub>m</sub> values for 24 and 48 hr are also presented.	Surber and Pickering (1962)
	Dalapon (tech)	Bluegill	BSA	-	105 (T4A)	а	This is an estimated LC $_{\!$	Cope (1965)
	Dalapon	Simocephalus serrulatus Daphnia pulex	BSA	-	16 (SB) 11 (SB)	_	Concentration reported is for immobilization. Time for immobilization was <b>64</b> hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
COMMERCIAL CHEMICAL PRODUCTS	Dalapon	Lepomis macrochirus (eggs) L. cyanellus (eggs) Micropterus dolomieui (eggs) Erimyzon sucetta (eggs) L. macrochirus (fry)	L	-	50 (S), 50 (NTE) 50 (NTE) 50 (NTE) 50 (NTE) 50 (S)		Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days.	Hiltibran (1967)
	Dalapon	Salmon Bluegill	BSA	-	340 (T2A) 115 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
4-(2,4-DB)	Lepomis macrochirus Micropterus salmoides	BSA	_	8.0 (T2A) 10 (T2A)	<u>a</u> co	The response of bluegill and bass fingerlings to nine agri- cultural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
4 (2,4) DB, (tech)	Rainbow trout	BSA	-	5.4 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F	Cope (1965)
DBrDT (DDT analogue)	Goldfish Gambusia affinis Culex apicalis (Iarvae)	BSA	-	0.06 (K) 0.01 (K) 0.0015 (K)	-	Experiments were run a maximum of 3 days. No other time data were reported.	Odum and Sumerford (1946)
DBS	Trout	ВСНА	_	5.0-20 (SB)	_	Fish exposed to 5.0 ppm of the chemical suffered a reduc- tion of the epithelium and a loss of mucous cells on top of the gill laminae. Exposure to 20 ppm for one hour caused great destruction of the epithelium — followed by death from suffocation.	Schmid and Mann (1961)
DDD	Channel catfish (fingerlings)	BSA	-	<2.6 (K 25 hr A)	<u>a</u>	Tap water was used. Conserable additional data are presented.	Clemens and Sneed (1959)
DDD	White catfish Largemouth bass Brown bullhead Black crappie Bluegill Hitch Sacramento blackfish Carp	FLCH	Cal.	0.014 (SB, application rate for 1949, -51, -57)	a g	Gnat control program with follow-up on accumulation in various species of wildlife. Ppm of DDD in edible flesh white catfish in 1958 was 30.4-129.0, Concentrations in remaining fish were 5.4-115 ppm.	Hunt and Bischoff (1960)
DDD (TDE, tech)	Salmo gairdneri	BSA	_	30 (T1A)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)

	TDE (DDD)	Chironomus riparius Asellus aquaticus Salmo gairdnerii	FL	Essendon Hertfordshire	(O)	a c m	Initial laboratory tests suggested that settleable powder formulations of TDE and DDT at application rates greater than 0.5 lb/acre would be effective in controlling <i>Chironomus</i> larvae, and that while trout would not be killed either directly (0.5-1 lb/acre) or by feeding on TDE- treated larvae, other invertebrates, e.g. <i>Asellus</i> , would be affected. Tubificids were not killed at application rates up to 2 lb/acre. TDE was considered a more useful insecticide than DDT because of its lower toxicity to fish. Carp were kept in cages for 11 months following the insecticide treat- ment; 35% died but the survivors grew well. By November 1962, fish tissues contained about 15 ppm TDE. In lab studies a 50 percent kill occurred in 7 days, at 0.1 lb per acre for chironimid larvae, at 0.5 lb per acre for <i>A.</i> <i>aquaticus</i> , and 2 lb per acre killed 10% of the tubificid worms. No deaths of rainbow trout occurred in 7 days at 2 lb per acre.	Edwards, et al (1964)
	DDD	Gambusia affinis affinis	BSA	-	0.46 to (L 1-1/2)	а	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are exposed later.	Boyd and Ferguson (1964)
ቅ	DDD (tech)	Bluegill	BSA		0.042 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
53	DDD	Pygosciles adeloriae Lobodon carcinophagus	FM	Ross Island, Antarctic	(O)	а	Adult penguins assayed showed no residue. The pre-molts examined had residues ranging from 0 to 16 ppb in the liver and 0 to 2 ppb in the fat. The crabeater seal examined showed residues of 2 ppb in the liver and 7 ppb in the fat.	George and Frear (1966)
	TDE (DDD)	Simocephalus serrulatus Daphnia pulex	BSA	-	0.0045 (SB) 0.0032 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
COMMERCIAL CHEMICAL PRODUCTS	pp'DDD	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto alba Strix aluco	FO	Netherlands	(O)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecticides. In most cases the liver had the highest con- centration of toxicant, ranging from 0.08 to 8.6 ppm. Most chlorinated hydrocarbons tend to accumulate in the fat depots of the body. In instances where mesenterial fat was analyzed the concentration of toxicant was found to be as high as 5.1 ppm.	Koeman and van Genderen (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
TDE (DDD)	Daphnia magna	BSA	-	0.0046 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
DDD	Atlantic salmon Brook trout	FR	New Brunswick	(O)	-	Spraying with this chemical at 0.25 to 0.5 lb/acre was no more harmful than with DDT at 0.25 lb/acre.	Kerswill and Edwards (1967)
DDD	Limnephilus rhombicus Sialis sp Gammarus sp	_	Knights Creek, Wisc.	(0)	_	Pesticide usage in an orchard did not significantly contaminate the aquatic environment of this creek adjacent to the treat- ment as determined by residue analysis.	Moubry, et al (1968)
TDE (DDD)	Pteronarcys californica (naiads)	BSA	_	0.38 (T4A)	acdef	Data reported at LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968
DDE	Penaeus aztecus	L	-	0.0068 (O)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
DDE	Oyster	BCF		0.014 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
DDE	Leptonychotes weddelli Pygosalis adeloriae Catharacta skua maccormicki	FM	Antarctic	(0)	а	<ul> <li>All residues are expressed as ppm wet weight. It was established that residues in the water were less than 0.0005 ppm. No detectible residues were found in tissues of <i>L. weddelli</i>.</li> <li>No detectable residues were found in tissues of <i>P. adelias</i>.</li> <li>Residues ranging from 0.01 to 0.73 ppm were found in tissues of <i>C. skua maccormicki</i>.</li> </ul>	George and Frear (1966)
DDE	Pygosciles adeloriae Lobodon carcinophagus	FM	Ross Island, Antarctic	(O)	а	Adult penguins assayed had residues ranging from 20 to 28 ppb in the liver. The fat residues in the pre-molts penguins ranged from 19 to 45 ppb. Crabeater seal examined showed residues of 7 ppb in the liver and 17 ppb in the fat.	George and Frear (1966)
pp'DDE	Platalea leucorodia Haematopus ostralegus Sterna sandvicensis Sterna hirundo Larus ridibundus Somateria mollissima tadorna tadorna	FO	Netherlands	(0)	-	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecticides. In most cases the liver had the highest con- centration of toxicant, ranging from 0.1 to 6.0 ppm. Birds feeding predominantly on crustacea, molluses, and fish contained significant amounts.	Koeman and van Genderer (1966)

• • • • • • • • •

	pp′DDE	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto alba Strix aluco Osio otus Falco pereginus	FO	Netherlands	(O)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecticides. In most cases the liver had the highest con- centration of toxicant, ranging from 1.2 to 75.2 ppm. Most chlorinated hydrocarbons tend to accumulate in the fat depots of the body. In instances where mesenterial fat was analyzed the concentration of toxicant was found to be as high as 68.3 ppm.	Koeman and van Genderen (1966)
	pp'DDE	Esox Iucius	FR	River Nene, Eng.	(O)	-	Higher concentrations were found in larger fish, indicating that they had been exposed to the pesticides for a longer time. Tissue extracts from the pike were analyzed for organochlorine pesticide residues by gas liquid chroma- tography. The average of six determinations was: 0.72 ppm muscle 96.0 ppm fat	Mawdesley- Thomas and Leahy (1967)
B-55	opʻDDE	Esox Iucius	FR	River Nene, Eng.	(O)	-	Comment same as above except that values for large fish only were: 0.042 ppm muscle 6.6 ppm fat	Mawdesley- Thomas and Leahy (1967)
	DDE	Limnephilus rhombicus Sialis sp Gammarus sp	_	Knights Creek, Wisc.	(O)	-	Pesticide usage in an orchard did not significantly con- taminate the aquatic environment of this creek adjacent to the treatment as determined by residue analysis.	Moubry, et al (1968)
	DDE	Oyster	FE	Galveston Bay, Texas	(0)	-	The chemical was found in the water at a concentration of $<$ 0.001 ppm. Oysters from the area were found to contain $<$ 0.01 to 0.05 ppm.	Casper (1967)
COMMERCIAL	DDE	Vascular plants Algae Chubs Largemouth bass Clams	FL	Tule Lake, Ore.	(0)	-	The amount of chemical sprayed in this area was not specified. The residue found was in the range of 0.6 to 1.0 ppb. Residue in chubs was from 2.5 to 45.0 ppb. The bass contained 11.0 to 38.0 ppb. Clams contained 0.75 to 6.3 ppb. The water contained 0.003 to 0.027 ppb.	Godsil and Johnson (1968)
AL CHEMICAL PRODUCTS	DDE	Alosa pseudoharengus Aplodinotus grunniens Coregonus artedii Lota lota	BSA	_	(O)	_	The study showed that the levels of chlorinated hydrocarbon pesticide residues in the fish meals and oils were, with the exception of the oil sample taken from the Lake Michigan alewife, below the regulatory tolerances established by the Food and Drug Directorate of Canada (1965) for certain foods intended for human consumption. Pesticide levels were interpreted as being representative for each species.	Dugal (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDE	Potamogeton pectinatus Cladophora Oscillatoria Cynodon dactylon Arundo donax	FR	Arizona	(O)	_	Irrigation canals were examined for plants which might serve as DDT collectors or indicators of DDT usage by con- centrating this material and its metabolites. Highest residues were found in <i>Cladophora</i> (19 ppm), followed by <i>Potamogeton</i> (9 ppm), and finally <i>Oscillatoria</i> (5 ppm).	Ware, et al (1968)
DDT	Carassius auratus	BSA	_	(O)	_	<ul> <li>DDT was used in 2 forms, as a dust containing 5 percent of DDT and in acetone solutions as a water suspension. The dusts were applied to the surface of the water by sifting through a fine-mesh screen.</li> <li>DDT produced characteristic symptoms in the goldfish in 24 hours when applied as either of the above formulations at concentrations of 0.2-2.0 ppm, but not at lower concentrations. The nervous system was affected causing a loss of equilibrium.</li> <li>When physa sp were tested, only the DDT from acetone solution was employed.</li> </ul>	Eide, et al (1945)
DDT	Huro salmoides Lepomis macrochirus Notemigonus crysoleucas Carassius auratus	FR	Back Creek, Glengary, W. Va.	(O)	d	<ul> <li>Aerial application of 1 pound per acre was made by plane. Only 0.39 pounds of DDT per acre reached the stream surface.</li> <li>Open live-boxes of fish were placed above, below, and within the sprayed section at five points. Of the 452 fish in these boxes, only 5 died from toxicity of the DDT. Predators removed a considerable number of fish from one live-box, and the handling of the fish was responsible for other losses. However, the survival, even with these losses, amount to 89.8%.</li> </ul>	Hoffman and Surber (1945)
	Invertebrates Orders: Annelida Megaloptera Ephemeroptera Odonata Plecoptera Coleoptera Trichoptera Diptera Mollusca					<ul> <li>Application of the chemical showed a rapid paralyzing effect on invertebrates.</li> <li>Application upon the bottom fauna revealed good survival (67%) at the first station and poorer survival (26% and 33%) at locations down stream.</li> <li>Wettable DDT applied at 1 pound per acre is not so toxic to fish and fish-food organisms as the same amount of DDT applied in an oil spray.</li> </ul>	
DDT	Daphnia magna	BSA	-	0.001 (SB1A)	-	Sublethal effect observed was immobilization of the <i>Daphnia.</i> Lake Erie water was used.	Anderson (1945)
DDT	Carassius auratus	BSA	-	2.0 (K1) 1.0 (O)	e	Ethyl alcohol was used as a solvent for 2 percent DDT. At 1 ppm, 90 percent of the fish were killed in 1 day.	Ginsburg (1945)

DDT (mosquito larvicide 50-D)	Carassius auratus	BSA	-	0.4 (O)	e	Mosquito Larvacide 50-D is water-white kerosene con- taining 3 percent DDT in solution and emulsified with sodium lauryl sulfate. A concentration of 0.4 ppm caused a mortality of 30 percent in 3 days at the spraying rate of 0.42 pounds per acre.	Ginsbury (1945)
DDT (Dust)	Carassius auratus	BSA	-	0.1 (O)	e	A concentration of 0.1 ppm killed 17 percent of the fish in 3 days at a spraying rate of 0.1 pounds per acre. 1.0 ppm killed 17 percent of the fish in 3 days at a spraying rate of 1.0 pounds per acre.	Ginsburg (1945)
DDT	Salmo trutta	BSA	Ithaca, N.Y.	0.25 (T1A)	<u>a</u>	LD <sub>50</sub> was determined in hours of survival time. The chemical was added as a wettable powder, as a solution in xylene with an emulsifying agent, and as a kerosene solution. It was most toxic as the emulsion.	Everhart and Hassler (1945)
DDT	Fall fish Common shiner Bluegill sunfish Eastern madtom Silverling minnow	FRK	Patuxent River, Md.	(0)	_	Fish kill occurred after an area was sprayed with an oil solu- tion of DDT at the rate of 2.0 lb/acre. Several other field studies were discussed in this report, but without much quantitative data. It is interesting that this paper presented as the first recommendation "Don't use DDT unless you must."	Cottam and Higgins (1946)
DDT		FR	W. Va.	(O)	-	An oil preparation of DDT was applied from a plane. The average deposit was 0.27 pounds per acre even though it was applied at 1 pound per acre. Thirteen stations were set up for sampling, Nrs. 1 and 2 were above point of application and the remainder were at and below the application point up to 2.0 miles. In general the closer the station to the point of application the more toxic the chemical.	Surber and Friddle (1946)
	Hyborhynchus notatus			1.0 (O)		Bluntnose minnows were not affected by the spray.	
	Micropterus salmoides			1.0 (O)		Greatest toxicity was noted at Station 10 (0.8 miles from point of application) which showed 46% survival. 100% survival occurred at Station 13 (2 miles from point of application).	
	Lepomis macrochirus			1.0 (O)		Greatest toxicity was noted at Station 10 (0.8 miles from point of application) which showed 74% survival. All other stations except 13, which had 100% survival, showed 92% survival of bluegill.	
	Pimephales promelas					Blackhead minnows showed 68-74% survival at most stations; 100% at Station 13.	
						DDT proved to be more toxic when applied as an oil spray rather than a suspension or a powder.	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDT	Goldfish Goldfish plus <i>Elodea</i> spp and <i>Cabomba</i> spp	BSA	_	0.07 (K) 0.16 (K)	-	Data are also given for analogues of DDT. Experiments were run a maximum of 3 days. No other time data were reported.	Odum and Sumerford (1946)
	Gambusia affinis			0.1 (K<1)			
	<i>Culex apicalis</i> (larvae)			<0.0001 (K)			
	Culex apicalis (pupae)			0.025 (K)			
DDT	Carassius auratus	BSA	_	0.2 (K5)	-	DDT fed to mosquito larvae at 1 part per 20 million killed 100 percent of the mosquitos in 2 days. DDT fed to gold-	Ginsburg (1947)
	Aedes aegypti			0.05 (K2)		fish at 1 part per 5 million killed 100 percent of the gold- fish in 5 days. Mosquito larvae killed by DDT at the above concentration when fed to goldfish did not have a toxic effect.	
DDT (Velsicol NR-70)	Numerous fish species and other aquatic organisms	FL	Tenn.	(O)	-	DDT was applied by thermal aerosol at the rate of 0.1 lb/acre. Rate at center of swath was 0.012 lb/acre. Anopheline and culicine mosquitoes were almost eliminated. After 16 appli- cations, the conclusion was that fish populations were unchanged when compared to controls.	Hess and Keener (1947)
DDT	Mastigophora Infusoria Hydrocarina Diatomaceae Synura Dinoflagellata Phacus Rototoria Copepoda Chroococcaceae Scenedesmaceae Chlamydomonas Euglena Trachelomonas Sarcodina	FL	Savannah, Ga.	(O)	_	No drastic killing of any specific group of organisms occurred from DDT treatment. At 0.5 lb/acre DDT spray, the growth of <i>Mastigophera, Synura</i> , Dinoflagellata, and Copepoda was inhibited; while <i>Phacus</i> , Rototaria, Chroococcaceae, and <i>Euglena</i> , appeared to be stimulated. All others were apparently unaffected. Similar results were obtained with a 0.1-0.2 lb/acre dust application of DDT. The author uses line-graphs to indicate trends of populations before and after treatment.	Bishop (1947)
DDT- copper sulfate	<i>Synedr</i> a spp <i>Daphnia</i> spp <i>Cyclops</i> spp	FL	Del.	(O)	-	Describes conditions, after DDT aerial spraying of a city water reservoir. Zooplankton disappeared with an over abundance of <i>Synedra</i> spp. Control of <i>Synedra</i> spp with 0.25 ppm CuSO4 was not effective. A possible antagonism of DDT to copper sulfate is noted.	Shane (1948)

	DDT	Bluegill Largemouth black bass Smallmouth black bass Golden-shiner Black crappies	FL & CF	Kearneysville, W. Va.	(0)	а	<ul> <li>Small bluegills, largemouth black bass, and smallmouth black bass one inch in length were killed by DDT in oil formulations in applications ranging from 0.25 to 1.0 pound per acre.</li> <li>Golden shiner fry were killed by oil sprays in excess of 0.25 pound per acre in dirt-bottomed ponds.</li> <li>Young black crappies 1.2 inches in length were killed by 0.5 pound per acre of DDT in both suspension and oil formulations.</li> <li>Fingerlings 2 inches or more in length were better able to withstand the higher rates of application.</li> <li>Fingerling bluegills, smallmouth black bass, and black crappies were found to be more sensitive to DDT than largemouth black bass, golden shiners, and trout.</li> <li>In continuous flow raceways, brook and rainbow trout, smallmouth bass, and golden shiners were relatively unaffected by a 1 pound per acre application of DDT.</li> </ul>	Surber and Hoffman (1949)
	DDT	Black fly (larvae) Rainbow trout Caddis fly	FR	Alaskan streams	0.3 (O)	-	The chemical was applied for control of black flies, and because the acetone solution was most effective, only that data is reported here. The figures reported are for minimum effective dosages for black fly larvae and maximum nonlethal dosages for rainbow trout and caddis fly larvae.	Gjulian, et al (1949)
	DDT	Lebistes reticulatus	BSA	-	0.025 (K1- 15%)	-	This is a bioassay method for determining DDT residue extracted from vegetables.	Pagan and Hageman (1950)
	DDT	Lepomis macrochirus Micropterus salmoides Notemigonus crysoleucas	FL	Auburn, Ala.	0.1 to 0.5 (K) –	-	Adult fish were not killed at 0.2 ppm. All were killed at 0.5 ppm in earthen ponds, in concrete pools 0.04 ppm was lethal. Adult fish were killed at concentrations greater than 0.1 ppm. This species withstood a concentration of 0.18 ppm.	Lawrence (1950)
		auratus Pomoxis			_	-	In concrete pools. 0.18 ppm was lethal to this species.	
~		nigromaculatus Megastomatobus			_	-	Withstood 1.0 ppm in earthen ponds.	
ŇMŇ		cybrinella Pimephalas promelas			_	_	The last two species withstood 0.4 ppm, but were killed at 2.0 ppm.	
COMMERCIAL		Carassius auratus			-	-	The toxicity of DDT to all species seems to be partly dependent on the form in which the chemical is added — as wettable powder or emulsion.	
L CHEMICAL PRODUCT	TDD	Bottom organisms: Ephemeroptera Odonata Plecoptera Megaloptera Coleoptera Trichoptera Diptera	FR	Wilkes Barre, Pa.	(O)	-	The DDT application of 1 lb/acre was made for control of <i>Porthetria dispar</i> , gypsy moth. Aquatic insects of the orders Megaloptera and Odonata appeared to be resistant to DDT poisoning at the dosage applied. Trichoptera were affected severely. Insect mortality increased as DDT moved down stream.	Hoffman and Drooz (1953)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDT	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): Cl $-$ NT Ma $-$ T (7) So $-$ NT Cv $-$ NT Gp $-$ PT (7) Np $-$ NT	Palmer and Maloney (1955)
DDT	Labea sp Synodontis schall	FR	Khartoum	(O)	_	A section of the Blue Nile was sprayed by air with an emulsified oil containing 30% DDT and 0.5% "Lissapol". The material was applied about 10 miles above Khartoum. It was hoped that by the time the treated water reached Khartoum, the DDT concentration would have been diluted 4 times. The concentration at the time of arrival was 0.17 ppm, and was maintained at a level of 0.003 ppm for 6 hr. It was reported that hundreds of fish were found dying 2 miles above Khartoum. <i>Labea</i> sp died in 8 hr and <i>Synodontis schall</i> died in 31 hr. The fish were analyzed for DDT residue the next day. The results are given below:	Burden (1956)
						Flesh Fatty	
						<u>Gills</u> <u>Viscera</u> <u>Deposit</u> Labea sp 0.9 ppm 2.5 ppm Nil Synodontis 2.7 ppm 7.9 ppm 64 ppm schall	
ססד	Labea sp Synodontis schall	F	Khartoum	0.09 (K)	_	Various levels of DDT were found in dead fish from 0.017 to 0.003 ppm downstream from the application. Undeter- mined degree of kill occurred.	Burdick, et al (1965)
тот	Young salmon	F	Canada	(0)	-	No toxicity data on fish were reported. The report deals primarily with reduction of insects available as fish food.	lde (1957)
DT	Salmo salar Salvelinus fontinalis Salmo gairdnerii	BSA	_	0.08 (L3) 0.16 (L<1) 0.08 (L3) 0.16 (L<2) 0.08 (L3) 0.16 (L<2)	a e	100 percent mortality occurred at 0.16 ppm in 18 hours for landlocked salmon, in 54 hours for rainbow trout, and 26 hours for brook trout.	Hatch (1957)
DT	Simulium sp (Iarvae)	FR	Streams, S. C. and Fla.	0.1-3.4 (0)	-	In slow-moving streams in Florida and South Carolina, DDT at the indicated concentrations controlled blackfly larvae for up to 0.28 mile. Control lasted for approximately 2 weeks. Data are presented as percent larval detachment in 1, 2, and 3 days time. Emulsion (0.1 ppm), oil (0.1 ppm), and granule (0.5-1.0 lb/acre) formulations had about the same degree of effectiveness.	Davis, et al (1957)
	A	-	"haimkey"				

APPENDIX B

.8

	DDT	Blue crab Marsh fiddler crab Red-jointed fiddler crab Cyprinodon variegatus Leiostomus xanthurus Mugil curema	FE	Bombay Hook Island, Del.	(O)		<ul> <li>The location under study was a salt marsh bounded by Delaware Bay.</li> <li>DDT was applied at 0.2 pound per acre.</li> <li>Organisms were confined in cages within the test area.</li> <li><i>C. variegatus, M. curema,</i> and <i>L. xanthurus</i> showed no mortality when exposed for 7 days.</li> <li>Blue crabs showed 17 percent mortality when exposed for 7 days in streams and 10 percent mortality in ponds.</li> <li>Marsh fiddler crabs and red-jointed fiddler crabs showed 75 and 36 percent mortality, respectively, in 7 days.</li> </ul>	George, et al (1957)
	DDT	Insectivorous fish Ephemeroptera Trichoptera	FR	Uganda	(O)	-	Rapid recolonization of aquatic insect populations decreased the possibility of accumulation of DDT by fish. Applica- tion rate was not given.	Corbet (1958)
<b>J</b>	DDT	Various aquatic and terrestrial organisms	FL	Salt Lake Co., Utah	(O)	-	The chemical was applied at 0.3 lb/acre. This concentration was sufficient for mosquito larvae control. At the above concentration no ill effects were observed in mammals, birds, reptiles, and amphibians. Invertebrates were not affected uniformly. Crustaceans were not harmed, nor were larvae of the insect family Ephydridae. Spiders and aquatic insects other than Ephydridae were adversely affected in varying degrees. Aquatic beetles seemed to be affected more seriously than other insects except mosquito larvae.	Graham and Anderson (1958)
-	DDT	Atlantic səlmon	BSA	_	0.049 (T1A) 0.047 (T2A)	a e	Results are recorded in ppm of insecticide by weight in water. Changes in temperature had an effect on the toxicity of the chemical.	Keenleyside (1958)
0	DDT	Brook, rainbow and cutthroat trout and whitefish Aquatic insects	FR	Montana	_	a c d	This study involves 13 rivers and streams following aerial spraying of DDT at rate of 1 lb/acre for control of spruce bud worm. The DDT recovery rate varied from 0.19-0.32 lb/acre. Significant amounts of DDT in fish tissue were found 16 months after spraying. The concentrations varied from 0.01 $\mu$ g/mg to 4.0 $\mu$ g/mg. Aquatic bottom invertebrates and adult insects were materially reduced in number but recovered in 1 year.	Graham and Scott (1958)
OMME	DDT	Artemia salina	BSA	-	0.142 (K <1)	ai	Rock salt was used in rearing all cultures employed in bio- assay work. The optimum salt concentration was 3.5%.	Tarpley (1958)
COMMERCIAL CHEMICAL PROD	DDT	Fathead minnow Bluegill Goldfish Guppy	BSA	_	0.032 (T4A) 0.016 (T4A) 0.027 (T4A) 0.043 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish" It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds.	Tarzwell (1959)
PRODU	DDT	Fathead minnow	BSA	-	0.034 (T4A)	<u>a</u>	Comment same as above except that this experiment was per- formed in hard water.	Tarzwell (1959)

COMMERCIAL CHEMICAL PRODUCTS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEI	DDT (50% dust)	Channel catfish (fingerlings)	BSA	-	>2.0 (K 25 hr A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
MICA	DDT	Pimephales promelas	BSA	-	0.03 (T4A)	a d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
PR		Lepomis macrochirus			0.02 (T4A)			
0 C		Carassius auratus			0.03 (T4A)			
CTS		Lebistes reticulatus			0.04 (T4A)			
	DDT	Pimephales promelas	BSA	-	0.042 (T4A)	abecdf	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et al (1959)
		Lepomis macrochirus			0.021 (T4A)			(1000)
		Carassius auratus			0.036 (T4A)			
		Lebistes reticulatus			0.056 (T4A)			
	DDT (screened)	Pimephales promelas	BSA	-	0.026 (T4A)	<u>a</u> becdf	Comment same as above.	Henderson, et al (1959)
	DDT	Oncorhynchus kisutch	BSA	-	(O)	а	This study provides information relating to an extensive field survey conducted to assess the effect of DDT spray deposition on aquatic fauna within the sprayed area. Times to 50% mortality (ET <sub>50</sub> ) were 850 minutes for 0.31 DDT, and 1750 minutes for 0.08 ppm of DDT. Levels of 0.05 ppm may be "safe" for coho salmon inas- much as this level did not produce death in one week.	Alderdice and Worthington (1959)
	DDT (dust)	Tilapia melanopleura	FLCH	Tanganyika	1 lb (6.6% К)	_	Trial periods were for 20 weeks. Sublethal effects such as impaired breeding, retarded growth, or altered taste were not detected. Dosages are given as lb/acre of surface water.	Webbe and Shute (1959)
	DDT	Pimephales promelas	BSA	_	0.032 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as	Tarzwell (1959)
		Lepomis macrochirus			0.016 (T4A)		toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	
	DDT	Sucker Trout	FR	Mont.	(O)	_	Aerial application rate was 1 lb/acre. Recovery of DDT from streams was 0.01 to 1.2 lb/acre. Surface water 15 minutes after spraying contained 1.35 ppmg. Sub-surface 0.08 ppm, and zero DDT before spraying. Initially, dead fish were mostly suckers, but no trout. Dead trout began appearing 5-6 months after spraying. Trout body tissue contained DDT. Considerable variation was found and no conclusions could be made.	Graham and Scott (1959)

	DDT	Gambusia affinis Huro salmoides	FL	Fla.	(O)	_	Surface applications of DDT as a dust and in oils were not harmful to fish in dosages used for mosquito control (0.1 pound per acre) other cold-blooded aquatic life, such as frogs, snakes, crayfish, spiders, and insects, were obviously affected by suspensions of DDT. No harmful effects on warm-blooded animals in the area were noted. When a small pond was treated with 2.0 ppm of a suspension containing 10 percent each of DDT and Nopco 1216 (sulfonated sperm oil) in cellosolve, at the end of 1 week all fish were killed. A third pond treated with the same formulation at 0.2 ppm killed all the fish in 4 days.	Dupree (1960)
	DDT	Salvelinus fontinalis Salmo clarki Prosopium williamsoni Salmo gairdnerii Salmo trutta Rhinichthys cataractae	FR	Mont. & Wyo.	(O)	_	This paper deals with the accumulation of DDT in trout and whitefish after exposure to DDT sprayed over large areas in Montana and Wyoming. The chemical was applied at 1 lb/acre with an average of approximately one-quarter pound per acre reaching the ground and the water. The greatest concentration of DDT was found in fat, followed by kidney, pyloric caecum and brain, in that order. At the given rate of application, this chemical was toxic to all the fish listed.	Cope (1961)
3	TOD	Daphnia magna	BSA		0.0014 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
Ĵ	DDT	Salmon	FR	Mitamichi R., N.B., Can.	(O)	-	Spraying with DDT in unspecified amounts markedly re- duced the salmon population in this river. In 1954, salmon fry were virtually eliminated and most of the parr were killed.	Kerswill, et al (1960)
	DDT (technical,	Acroneuria pacifica	BSA	-	0.18 (T4A)	acefin	Assays were conducted in hard water.	Gaufin (1961)
	25% active in xylene)	Pteronarcys californica			0.33 (T4A)			(1301)
		Claassenia sabulosa			0.01 (T4A)			
0		Arctopsyche grandis			0.1 (T4A)			

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDT	Brook trout White suckers Crayfish	F	Potter and Tioga Counties, Penn.	(0)	-	<ul> <li>DDT was applied as an aerial spray at 0.5 lb/acre. Thirty-two days after spraying, 10.6 ppm was found in brook trout, but 122 days after treatment the amount was at pretreatment level (0.7 ppm).</li> <li>In white suckers, 32 days after treatment, 6.9 ppm was found; and 122 days post-treatment, the concentration had dropped to a pretreatment level (0.24 ppm).</li> <li>Analyses of crayfish were anomalous – in some instances, the pretreatment specimens contained 1.9 ppm or more than 32 days later when the value was 1.1 ppm.</li> <li>This same paper gave some data on the DDE, TDE, and dieldrin content of these same animals.</li> </ul>	Cohen, et al (1961)
DDT	Oncorhynchus kisutch Oncorhynchus tshawytscha Salmo gairdnerii Gasterosteus aculeatus	BSA	-	44 (T4A) 11.5 (T4A) 42 (T4A) 18 (T4A)	<u>a</u> c d <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
DDT	Salmo gairdneri	BSA	-	0.410 (T1A) 0.410 (T2A) 0.395 (T4A)	acdfg	Hatchery artesian well water was employed for this experiment.	Webb (1961)
TDD	Aedes aegypti (larvae)	BSA	-	-	<u>a</u>	Increase in temperature during exposure to DDT (0.02 ppm — 1 hr) increased the toxic action. Additional data are presented.	Das and Needham (1961)
DDT (25 percent EC)	Gambusia affinis	FL	Ponds — Bakersfield, Cal.	(0)	ac	At 0.5 lb/acre, 20 percent mortality occurred in 24 hours. At 2.0 lb/acre, 40 percent mortality occurred in 24 hours, The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
DDT (tech grade emulsified in xylene)	Hydropsyche californica Arctopsyche grandis Acroneuria pacifica Pteronarcys californica	BSA	-	0.048 (T4A) 0.175 (T4A) 0.41 (T4A) 0.56 (T4A)	<u>a</u> cdeln	Test water was obtained from a mountain stream.	Gaufin (1961)
DDT	Ephemeroptera Trichoptera Plecoptera	FR	Mont.	-	_	A large area in Montana was sprayed with 1 pound of techni- cal DDT in 1 gallon of No. 2 diesel fuel per area. The streams draining this area were assayed. It was found that a drastic reduction in all biota took place in Hellroaring Creek and Tower Creek. Pebble Creek, situated away from the test area, served as a control and no reduction occurred. In both Hellroaring and Tower Creeks, repopulation of plecopteran populations occurred leveled by the following year while this did not occur for the other organisms until 3 years had elapsed	

	DDT	Lebistes reticulatus (Adult) (Young) Salmo trutta (2 wk) (10 wk) (11 wk)	BSA	_	0.018 (T14 A) 0.0024 (T14A) <0.18 (T14A) 0.00056 (T14A) 0.014 (T14A) fingerlings	<u>a</u>	<ul> <li>Deep well water was used as diluent. Histological observations were found to be similar for guppies and trout despite age differences, DDT concentration, and exposure periods.</li> <li>Liver: degeneration</li> <li>Kidney: no change in guppy - 1-2 days tubules occluded in trout.</li> <li>Other data and observations presented.</li> </ul>	King (1962)
	DDT	Salmo trutta	BSA	-	0.5 and 0.1 (O)	-	C <sup>14</sup> labeled DDT was placed in the water at the concentra- tions listed. Various organs and tissues of the trout were analyzed for DDT. The analytical method is outlined. At 0.5 ppm, one fish died in 18 hours, and another at 160 hours. At 0.1 ppm, one fish died in 30 hours and another at 230 hours. Only two fish were used at each concentration.	Holden (1962)
	TOD	Ephemeroptera	FR	Adirondack Mountains	(O)	-	DDT was applied as low as 0.1 lb/acre for effective control of blackfly. There was a small number of Ephemeroptera and Diptera in regularly treated streams, but the reduction in overall numbers did not reach a significant level. The treated and untreated streams were sampled in 1950-52 and in 1961.	Jamnback and Eabry (1962)
	DDT	Various insects	FS	Africa	(O)	-	This study showed that the effect of a single application of DDT on an African stream eliminated the majority of aquatic insect species for varying distances. It seems fairly certain, however, that almost all the species survived as eggs, from which the population was replenished. The three major predators were, however, among the most severely affected, and this led to an increase in the principal prey organisms, Baetidae and <i>Simulium</i> . This was an effect which has been observed in streams in Europe and North America, and indicates that DDT treatments could lead to severe outbreaks of <i>Simulium</i> species. The amount of the DDT sprayed on an area was not specified.	Hynes and Williams (1962)
COMMERCIAL CI	DDT	Protococcus sp Chlorella sp Dunaliella euchlora Phaedactylum tricornutum Monochrysis lutheri		-	0.6 (O)* 0.6 (O)* 0.6 (O)* 0.6 (O)* 0.6 (O)* *obvious, but inhibited growth.	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were expressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants.	Ukeles (1962)
COMMERCIAL CHEMICAL PRODU	DDT	Cutthroat trout	FL	Wyo.	(O)	а	Five lots of fish were given DDT once a week in their diets at different rates for each lot. Fish were fed in mg/kg of body weight. Analysis showed that residues of DDT, DDE, and DDD were present in fish fed 3 and 1 mg/kg per body weight. Greatest number of deaths occurred in fish lots fed 3 mg/kg of body weight.	Соре (1963)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chemical	Rainbow trout Bullhead Crayfish	FL	Colo.	0.02 (O)	_	<ul> <li>The pond was treated with DDT at a rate of 0.02 ppm.</li> <li>The concentration of DDT in the pond was at its highest point 30 min after treatment. None could be detected after 21 days.</li> <li>Bullheads and trout contained the greatest amounts of chlorinated hydrocarbon 30-40 days after treatment, with concentrations over 4 ppm. Levels slowly declined after that, averaging 3.5 ppm in samples taken 9 and 10 months after treatment in both species, and 3 ppm in rainbow trout taken 14 months after treatment.</li> <li>Crayfish developed lower DDT residues than did trout, and contained 0.33 ppm after 14 weeks.</li> </ul>	Соре (1963)
DDT	Acris crepitans A. gryllus	Lab	_	(0)	<u>a</u>	Possible resistance to DDT was demonstrated in natural populations of frogs from several Mississippi localities by exposure to different DDT concentrations for 36 hours. Frogs living near cottonfields heavily treated with DDT for several years tended to be less susceptible to the chemical than individuals having little or no prior contact with DDT. The animals were placed on filter paper impregnated with varying amounts of DDT. The data are difficult to average because of the range. It would appear that 30.0 g per liter was the TL <sub>m</sub> for 36 hours for <i>A. crepitans</i> , and 9.0 g per liter for <i>A. gryllus</i> (sic).	Boyd, et al (1963)
DDT	Cladocerans Copepods Ostrocods Rotifers <i>Volvox</i>	FL	Minn.	(O)	-	At an application rate of 1 lb/acre, depression of micro- crustacean populations occurred a few days after application. but this was followed by an apparent rapid recovery.	Jones and Moyle (1963)
DDT	Culex pipiens quadrimaculatus	BSA	-	(0)	с	Tests were conducted in tap water and artificially polluted tap water. The values reported are the concentration range for an LC <sub>90</sub> , 0.12 to 0.55 ppm for polluted and 0.33 to 128.0 ppm for tap water.	Lewallen and Wilder (1963)
DDT EC 2	Rana catesbeiana (tadpoles)	FL	Cal .	1.0 (O)	a c	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and resulted in a 30 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
DDT (tech)	Procambarus clarki	BSA		0.6 (T3A)	acdo	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
DDT (anti- resistant 50 percent WP)	Salmo gairdneri	BSA	-	24 (T1A) 21 (T2A) 16 (T4A)	а	The experiments were conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)

DDT (anti- resistant 25 percent EC)	Salmo gairdneri	BSA	-	10 (T1A)	а	Comment same as above.	Cope (1963)
DDT, p-p'	Səlmo gairdneri	BSA	-	18 (T 18 hr) 11 (T 32 hr) 10 (T 56 hr)	а	Comment same as above.	Cope (1963)
				10 (T1A) 10 (T1A)		Comment same as above. The experiment was conducted at 65 F. Fish weighed 0.6 g.	
				6.0 (T1A)		The experiment was conducted at 65 F. Fish weighed 0.4 g.	
				5-6 (T1A)		The experiment was conducted at 75 F. Fish weighed 1.5 g.	
	Lepomis macrochirus			6.0 (T1A)		The experiment was conducted at 75 F. Fish weighed 0.4 g.	
	Redear			19 (T1A) 15 (T1A)		The experiment was conducted at 75 F. Fish weighed 3 g.	
DDT	Rainbow trout Long-nose sucker Cutthroat trout Brown trout Brook trout Mountain whitefish	BSA	_	(O)	a d g	<ul> <li>Hatchery trout under 3.1 inches in length had a mortality rate of 100% at all concentrations for 0.5 to 10.0 ppm. The mortality rate decreased as size increased (66% for those over 5 inches). Wild rainbow trout under three inches had a mortality of 24%. Hatchery trout showed a 50 to 75% higher mortality than the wild trout.</li> <li>A comparison of six species of cold-water fish over four inches in length tested in 1 ppm DDT showed that the long-nosed sucker had a mortality rate of 94%, while the rainbow trout, cutthroat trout, brown trout, brook trout, and mountain whitefish had a mortality rate of less than 10%.</li> </ul>	Schoenthal (1963)
DDT	Gambusia affinis	BSA	-	0.05-0.10 (O)	-	<ul> <li>Mosquito fish from waters near cotton fields that have a long history of treatment with chlorinated hydrocarbon pesticides exhibited a marked resistance to DDT compared with fish from areas which had had no past exposure to insecticides. As an example, for a DDT concentration of 0.05 ppm:</li> <li>72 hours 90% mortality occurred for fish from untreated areas</li> <li>72 hours 25-28% mortality occurred for fish from DDT treated ponds.</li> </ul>	Vinson, et al (1963)
DDVP (tech)	Salmo gairdneri	BSA	-	500 (T1A)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDT	Salmo Iewisi clarki	BSA	_	(0)	a c e f	This experiment concerns the periodic exposure of fish to different levels of DDT in bath and in food over a 20-month period involving one spawning cycle. It was decided that a threshold level exists around 0.1 ppm monthly in contact form and around 0.3 mg DDT per Kg of fish weekly in the diet for the toxic effects of DDT. Fish lots given 0.1 ppm DDT monthly in bath form ex- hibited significantly higher mortality, similar size, and similar reproductive success when compared with the control group. Fish lots treated weekly with DDT in the diet at the rate of 0.3 milligrams per kilogram of body weight did not differ from the control except, in a highly variable manner, residue buildup, and a nonsignificant increase in mortality during the last few months of the experiment.	Allison, et al (1964)
DDT	Ophicephalus punctatus Barbus stigma	BSA	-	1.0 (K) (O)	_	<ul> <li>Experiments were conducted in a trough containing 3500 cc of water sprayed with 1 cc of a 25% DDT emulsion. Liver and intestines were examined for residues and both were found to contain 0.08083 g of DDT.</li> <li>When 5 cc of a 25% DDT emulsion was sprayed, the fish died in 2 hr. The fish were found to contain 0.1344 g of DDT residue in the intestine and 0.1292 g in the liver.</li> <li>When 5 cc of a 25% DDT emulsion was sprayed in a small jar containing 2 liter of water, death occurred in 2 hr and 30 min. Residues found were 0.06523 g of DDT in liver and 0.07799 g in intestines.</li> </ul>	Mathur (1964)
DDT	Notemigonus crysoleucas Lepomis macrochirus L. cyanellus	BSA	_	(B) 0.032 (T 1.5) (A) 0.028 (T 1.5) (B) 0.028 (T 1.5) (A) 0.033 (T 1.5) (B) 25 (T 1.5) (A) 22 (T 1.5)	acf	Chemical was dissolved in acetone. Final concentration of acetone was $\leq 2 \text{ ml/l}$ . Data shows TL <sub>m</sub> ppb for insecticide-resistant (A) and insecticide non-resistant (B) strains of the test fish.	Ferguson, et al (1964)
DDT (tech, 98 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica (naiad)	BSA	-	1.8 (T4A) 0.32 (T4A)	c d e f	<i>A. pacifica</i> was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than <i>P. californica.</i>	Jensen and Gaufin (1964)
DDT	Gammarus Iacustris	BSA	_	0.009 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
DDT	Bluegill	BSA	-	0.016 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)

	DDT	Trout: brown and rainbow <i>Cottus</i> sp Trichoptera Plecoptera Coleoptera Diptera Ephemeroptera	FR	Mont.	0.02 mg/l	_	The application rate was 0.5 lb/acre. Before and after spraying, determination of DDT concentrations was made on fish homogenates. Dead fish found after spraying were found to contain up to 0.6 mg/kg DDT. However, live fish caught were found to contain up to 3.4 mg/kg DDT. Fish kills occurred up to 48 hours after spraying. Exten- sive mortality of aquatic insects occurred, but not consistently throughout sprayed area of river.	Welch and Spindler (1964)
	DDT	Salmo gairdnerii lctalurus punctatus Lepomis macrochirus Pteronarcys californica	BSA	-	1.5 (T4A) 3.3 (T4A) 4.7 (T4A) 7.0 (T4A)	<u>a</u> cd	Toxicity values reported as median lethal conc. (LC <sub>50</sub> ) for 24, 48, 96 hr.	Bridges and Cope (1965) ;
	DDT + Toxa- phene	Oyster	BCF	_	0.030 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth),	Butler (1965)
7	DDT	Oyster	ВСН	_	(O)	~	Oysters exposed to DDT at a concentration of 0.0001 ppm contained 70,000 times that concentration after 40 days. Oysters exposed to the chemical at 0.001 ppm for 12 days contained 12 to 20 ppm. Because the oyster can concen- trate DDT so readily, it is an organism of choice to use in monitoring for pesticide pollution.	Butler (1965)
	DDT	Cyprinodon variegatus (juvenile) Fundulus similis (juvenile)	BSA	-	0.005 (O) 0.018 (O)	а	Water temperature was 9 C. The figure reported is a 48-hr EC <sub>50</sub> . Water temperature was 21 C.	Butler (1965)
	DDT + Strobane	Oyster	BCF	-	0.022 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
	DDT	Blue crab	всн	-	(O)	~	DDT at a concentration of 0.001 ppm kills crabs in 8 days. Crabs can live and grow in a concentration of 0.00025 ppm.	Butler (1965)
	DDT	Rainbow trout	BSA	_	4.1 (T4A) 5.0 (T4A) 6.0 (T4A)	<u>a</u>	These experiments were performed to show the effect of temperature on the toxicity. For the toxicant concentrations listed, the temperatures were respectively 45, 55, and 65 F. The fish all were approximately one gram in weight. Toxicant concentrations for one and 2-day times were also listed.	Cope (1965)
	DDT	Palaemonetes kadiakensis	BSA	_	(N) 4.5 (T1-1/2A) (TB) 10 (T1-1/2A)	acf	Test organisms were collected from 2 locations, Twin Bayou (TB), Sunflower Co., Miss. (agricultural area) and Noxubee National Wildlife Refuge (N), Noxubee Co., Miss. (non- agricultural area) and evaluated in laboratory bioassays. The Twin Bayou shrimp were more resistant.	Ferguson, et al (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDT	Gambusia affinis Ictalurus melas	BSA	_	0.008-0.023 (T3A) 0.009-0.275 (T3A)	<u>a</u> c d e	Test fish were collected from 8 different locations of the Mississippi River. The 3-day TL <sub>m</sub> values were made to determine if a resistance gradient existed. The data indicated that there was none.	Ferguson, et al (1965)
DDT	Lepomis cyanellus Gambusia affinis Ictalurus melas	F	Miss.	0.52 (0)	-	Muds reduced the toxicity of chlorinated hydrocarbon insecticides to fish. Lethal quantities of pesticides enter national waters and muds may contain sorbed pesticides in excess of lethal quantities. Although the chemicals can be leached with organic solvents, they were either not released in lethal amounts or slowly released in standing water.	Ferguson, et al (1965)
DDT	Arctopsyche grandis Pteronarcys californica Acroneuria pacifica Ephemeralla grandis Hydropsyche californica	BSA	-	0.18 (T4A) 1.8 (T4A) 0.3 (T4A) 0.03 (T4A) 0.05 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were determined by standard methods. General comments were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Addi- tional data are presented.	Gaufin, et al (1965)
	Daphnia magna Gammarus Iacustris Bluegill Fathead minnows			0.001 (T 50 hr A) 0.009 (T4A) 0.03 (T4A) 0.03 (T4A)			
DDT	Acroneuria pacifica Ephemerella grandis Gammarus Iacustris Pteronarcys californica	BSA	-	0.32 (T4A) 0.025 (T4A) 0.009 (T4A) 1.8 (T4A)	<u>a</u> C	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
DDT	Salmo salar	BCFCH	_	0.005-0.05 (O)	<u>a</u> e	The fish were exposed to the reported sub-lethal doses for 24 hours. Low dosages of DDT produced a downward shift in the temperature response whereas higher doses produced an upward shift.	Ogilvie and Anderson (1965)
DDT	Fish Shell fish Birds Mice	FLR	Conn.	(O)	-	The results showed that DDT was present in animals in areas where no spraying had been conducted. Analyses showed the following: Whole fresh fish - 0.1 to 0.9 ppm DDT Shell fish - 0.031 to 0.07 ppm Birds - 0.1 to 0.8 ppm Mouse kidney fat - 1.01 to 8.19 ppm	Turner (1965)

	DDT	Brown trout	BSCH	_	0.10 (O)	-	Data given on DDT concentrations in various tissues on Weiss weight basis are as follows: (1965 Brain 0.6 ppm Muscle 0.7 ppm	5)
	DDT	Oysters Adult Larvae	BCF	-	— (O) 1.0 (K6)	а	Sea water was employed in this experiment. As theButlerconcentration of DDT increased from levels of 1.0 ppb to(1966)1.0 ppm, there was a logarithmic decrease in the rate ofoyster shell growth from about 20 to 90 percent.	
	DDT	Crassostre <del>a</del> virginica	BCF	-	(0)	a	Tests were conducted in flowing seawater. DDT in levels as low as 0.001 ppm caused marked reduction in oyster (1966 growth.	
1	DDT	Salmo gairdnerii Lepomis macrochirus Ictalurus punctatus Pteronarcys californica Baetus sp Daphnia pulex Simocephalus serrulatus	BSA	-	0.005 (T2A) 0.005 (T2A) 0.012 (T2A) 0.016 (T2A) 0.012 (T2A) 0.0004 (T2A) 0.0002 (T2A)	a	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	;)
I	DDT	Cyprinodon variegatus	BSA	-	0.020 (O) 0.030 (O) 0.040 (O)	aei	A concentration of 0.020 ppm caused 80% mortality, Holland 0.030 caused 87% mortality, and 0.040 caused 97% (1966 mortality in 24 hr.	
	DDT	Gambusia affinis	BSA	-	(O)	<u>a</u>	The effect of combinations of pesticides was studied. In general, the results reflected the extreme levels of Endrin and Toxaphene resistance in the resistant population. The results failed to indicate additive effects wherein the com- bination mortality exceeded the sum of the mortalities produced by the individual insecticides.Fergus Bingh (1966)	am
COMM	DDT	Pygosciles adeloriae Lobodon carcinophagus	FM	Ross Island, Antarctic	(O)	а	Adult penguins assayed had residues ranging from 0 to 8 ppb.GeorgeThe pre-molts examined had residues ranging from 1 to 16Frearppb in the liver, and 0 to 69 ppb in the fat.(1966)eater seal examined showed residues of 4 ppb in the liverand 15 ppb in the fat.	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)	
p,p' -DDT	Salmo gairdnerii S. trutta Salvelinus fontinalis S. namaycush Esox lucius Carassius auratus Chrosomus eos Cyprinus carpio Pimephales promelas Ictalurus punctatus Eucalia inconstans Lepomis cyanellus Lepomis gibbosus L. macrochirus L. megalotis Micropterus salmoides Perca flavescens Aplodinotus grunniens I, melas	BSA		0.0107 (T4A) 0.0109 (T4A) 0.0115 (T4A) 0.093 (T4A) 0.0077 (T4A) 0.0587 (T4A) 0.0680 (T4A) 0.0082 (T4A) 0.0082 (T4A) 0.0075 (T4A) 0.0045 (T4A) 0.0045 (T4A) 0.0045 (T4A) 0.0045 (T4A) 0.0087 (T4A) 0.0009 (T4A) 0.0009 (T4A) 0.0100 (T4A)	<u>a</u>	Data are given for various lots within each species of fish obtained from different sources. However, a mean was given which is reported here. Data for fathead minnows were not consistent. The toxicity did not increase uniformly with increased concentrations up to 1.0 ppm.	Marking (1966)	
DDT	Simocephalus serrulatus Daphnia pulex	BSA	-	0.0025 (SB) 0.00036 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)	
DDT	Daphnia magna	BSA	-	0.0044 (SB)	_	Comment same as above.	Sanders and Cope (1966)	
DDT	Daphnia carinata	BSA	-	0.0022 (SB)	-	Comment same as above.	Sanders and Cope (1966)	

DDT	Oncorhynchus kisutch	BSA	-	0.024 (T2A) 0.013 (T4A)	<u>a</u> cejk	DDT at the given concentrations seemed to cause a coughing reaction in the fish. It was theorized that this coughing reaction was a reversal of the water flow over the gills as a gill cleansing reaction.	Schaumberg (1967)
DDT	<i>Tubifex</i> spp <i>Limnodrilus</i> spp	BSA	-	100 (L4A)	a c e	Toxicity is reported as the mean lethal dose (LD <sub>50</sub> ) for 24, 48, and 96 hours.	Whitten and Goodnight (1966)
DDT	Salmo gairdnerii Rasbora heteromorpha	BSA	-	(O) (O)	f	This report derives as mathematical equation for deter- mining a threshold concentration of a toxicant. For many toxins, the rate of mortality is a linear function of the concentration. The value of 0.02 ppm of DDT was obtained by interpolation from three different curves for the trout. The tests were conducted in hard water. A value of 0.04 ppm for harlequin fish was also obtained by interpolation from a graph.	Abram (1967)
DDT (mixed isomers)	Salmo gairdnerii Rasbora heteromorpha	BCFA	_	0.0015 (threshold)	<u>a</u> d <u>e</u>	<ul> <li>Aerated hard water was used. Threshold concentrations were examined by 4 methods.</li> <li>1. Long term - survival related to concentration.</li> <li>2. Short term - percentage kill in narrow range of concentrations.</li> <li>3. Comparison of survival times.</li> <li>4. Extrapolation of short-term results by plotting velocity of death against log of concentration.</li> </ul>	Abram (1967)
DDT	Mya arenaria Crassostrea virginica	BSCH	_	(0)	-	Oysters were exposed to 2.0 to 4.0 ppm DDT and then fed to shrimp and fish. At the end of 2 to 4 weeks, at least 50 percent of the experimental animals died. <i>M. arenaria</i> (soft clam) proved to be the most sensitive or efficient in storing organochloride residues.	Butler (1967)
DDT	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	-	(0)	-	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentration of 1200-9000X resulted.	Butler (1967)
DDT	Aquatic insects: Ephemeroptera Diptera, Simulidae Plecoptera Trichoptera	FR	Ontario, Can.	(0)	_	This is a review paper on the effect of DDT in the reduction of insects, and the time it takes a population to reestablish itself. The area was sprayed with DDT at a rate of 0.5 lb/acre Most organisms recovered to normal populations within 2-3 years, but caddisflies required 4 or more years.	lde (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDT	Leptonychotes weddelli Pygoscelis adeloriae Catharacta skua maccormicki Aptenodytes forsteri Rhigophila dearborni Trematomus bernacchii T. hansoni Invertebrate samples Arthropoda Echinodermata Nermertinea Mollusca	FM	Antarctic	<ul> <li>(O)</li> </ul>	a	<ul> <li>All residues are expressed as ppm wet weight.</li> <li>It was established that residues in the water were less than 0.0005 ppm. L. weddelli contained residues ranging from 0.042 to 0.12 ppm in fat. No residues were found in other tissues. Adult P. adeliae contained residues of 0.015 to 0.018 ppm of DDT in fat.</li> <li>C. skua maccormicki contained residues ranging from 0.01 to 0.68 in 9 tissues examined. A. forsteri adults were examined for residues and found to contain none.</li> <li>Ten R. dearborni were examined and found to contain an average of 0.44 ppm DDT residue. T. bernacchii and T. hansoni were examined and contained no residues.</li> <li>It was established that there were no residues at levels as high as 0.005 ppm in invertebrates.</li> </ul>	George and Frear (1966)
DDT (prills)	Amphipoda Elmidae Grypopterygidae Turbellaria Oligochaeta Gastropoda Decapoda Odonata Plecoptera Ephemeroptera Hemiptera Trichoptera Megaloptera Coleoptera Diptera Anguillidae Galaxiidae	FR	New Zealand (4 streams)	(O)	_	DDT prills were applied from the air to 200 acres of sheep pasture. The mean weight of active DDT reaching the ground was 61.3 mg/square meter. Changes in the fauna of streams draining the treated pasture were studied for 12 months after the application. A high mortality was found in most of the aquatic insect fauna with the excep- tion of Elmidae (Coleoptera) and Grypopterygidae (Plecoptera). Amphipoda were virtually wiped out. At the end of the investigation large number of insects were again present but with a completely altered species com- position. Mortality to freshwater crayfish was doubtful, and unproven for fish. The levels of DDT found in whole fish are discussed.	Hopkins, et al (1966)

	DDT and "organo- chlorine	Pontoporeia affinis Alosa	FLCH	Wisc.	-	-	Pesticide residues were determined in mud sediments, insects, fish, and birds. Conclusions were that the pesticides do not stay on land	Hickey, et al (1966)
	residues"	pseudoharengus Coregonus clupeaformis Leucichthys sp					but are accumulated and concentrated in moderately large lakes. Residues found in mud bottoms were: DDT = 0.14 ppm DDE = 0.24 ppm TDE = 0.03 ppm Residues found in: Gull food Insect - DDT = 0.12 ppm; DDE = 0.49; TDE = 0.06 Alewives - DDT = 1.13 ppm; DDE = 1.77; TDE = 0.43 Fish - Chub - DDT = 1.6; DDE = 2.3; TDE = 0.29 Whitefish - DDT = 1.7; DDE = 2.7; (muscle) TDE = 0.75. Additional residue data are presented.	
1	DDT	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.04 (O)	a	Stones heavily populated with larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number de- tached amounted to 52 percent.	Jamnback and Frempong- Boadu (1966)
	DDT	Fish	F	Cal.	(0)	-	This study was primarily concerned with insecticides found in fish-eating birds. Limited fish studies were also conducted. DDT was found in trace to 1.6 ppm concen- trations in whole fish (wet weight).	Keith (1966)
COMMERCIAL CHEMICAL	pp DDT	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto alba Strix aluco Osio	FO	Netherlands	(O)	-	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecticides. In most cases the liver had the highest concentration of toxicant, ranging from 1.6 ppm.	Koeman and van Genderen (1966)
- CHEMICAL		osio otus Falco pereginus						

**B-**75

COMMERCIAL CHEMICAL PRODUCTS

COMMERCIA	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Røference (Year)
COMMERCIAL CHEMICAL PRODUCTS	DDT	Crassostrea virginica Pseudomonas piscicida	BCFCH	_	(O) 10 (SB)	а	At 0.00001 ppm concentration, 0.11 ppm residue was found in 20 days. At 0.000001 ppm concentration, 0.085 ppm residue was found in 20 days. After extended periods of time, small amounts of DDT actually stimulated growth of <i>P. piscicida</i> .	Butler and Johnson (1967)
L PROD	DDT	Oyster	FE			_	The chemical was found in the water at a concentration of $\leq$ 0.001 ppm. Oysters from the area were found to contain $\leq$ 0.01 to 0.05 ppm.	Casper (1967)
UCTS	DDT	Stream insects: Ephemeroptera Odonata Plecoptera Trichoptera Neuroptera Coleoptera Diptera	FR	Maine	(O)	-	At an application rate of 1/2 to 1.0 pounds per acre, DDT produced marked reductions in the quality and quantity of the invertebrate fauna. Normally present fauna usually repopulate in 2-4 years. Thirty-nine genera of aquatic insects were studied.	Dimond (1967)
	p-p' DDT	Goldfish	BSA & CH	_	50 m g/ml (K)	-	Test fish were conditioned to alkyl benzene sulfonate $(4 \ \mu g/ml)$ or sodium lauryl sulfate $(4 \ \mu g/ml)$ , for various periods of time, then exposed to pesticides. Chronic exposure to the detergent increased the toxicity of the pesticide.	Dugan (1967)
D 76	DDT	Atlantic salmon	F	St. Andrews, New Brunswick	(O)	-	After spraying DDT at the rate of 1/2 lb/acre, all young salmon were reduced in number. Underyearlings were only 2-10% as abundant, small parr 30% as abundant, and long parr 50% as abundant. Spraying at the rate of 1/4 lb/acre reduced the numbers of underyearlings by 50%, small parr by 20%, but hardly affected large parr. Spraying with DDT at 1/4 lb per acre, applied twice, was followed by low numbers of underyearlings, similar to the effect of spraying at 1/2 lb/acre. No equivalent data for parr was available.	Elson (1967)
	DDT-C <sup>14</sup>	Lepomis macrochirus Carassius auratus	BCFCH	_	(0)	а	Fish were treated with carbon-labeled insecticides (0.03 ppm) from 5 to 19 hr and uptake rates were determined. They were placed in recovery tanks for up to 32 days. Whole body samples were then made. It was found that in both fish species >50 percent of the DDT absorbed was present after 32 days.	Gakstatter and Weiss (1967)
	DDT	Aquatic insects: Diptera Trichoptera Ephemeroptera Chironomidae	FR	Mitamichi River, New Brunswick Canada	_ ,	-	Young salmon, both fry and parr, were seined and the stomachs removed. Insects were removed from the stomachs, identi- fied and counted. Forage ratios of the insects were deter- mined and reported for 1953-1961. Reduction of all insects by DDT was soon followed by resurgence of Chironomidae and other Diptera. Five years after the spray- ing, the pre-spraying complexity of food for younger salmon was approached. Trichoptera were the slowest to reappear.	Keenleyside (1967)

	DDT	Salmo salar	FR	St. Andrews, New Brunswick	(0)	-	DDT at 0.25 lb/acre killed many insects and some fish, but was only about half as damaging to aquatic fauna as a heavier dosage (0.5 lb). After spraying at 0.5 lb/acre, young salmon of all size groups were found in reduced numbers.	Elson and Kerswill (1967)
	DDT	Atlantic salmon Brook trout Sucker Cyprinids	FR	New Bruns- wick	(O)		When DDT in an oil emulsion was sprayed at 0.5 lb/acre, heavy losses in underyearling salmon and parr were observed. Wild young salmon were found in streams when autumn water temperatures approached freezing after June sprayings with DDT. Spraying with DDT at 0.25 lb/acre had no effect on caged, or native fish during a period of 2 or 3 weeks after spraying. In one area suckers and cyprinids were extensively killed after 0.5 lb/acre spraying of DDT.	Kerswill and Edwards (1967)
	pp'DDT	Esox lucius	FR	River Nene, Eng.	(0)	-	Higher concentrations were found in larger fish, indicating that they had been exposed to the pesticides for a longer time. Tissue extracts from the pike were analyzed for organochlorine pesticide residues by gas liquid chroma- tography. The values for large pike were: 0.068 ppm muscle 6.7 ppm fat	Mawdesley- Thomas and Leahy (1967)
;	opʻDDT	Esox lucius	FR	River Nene, Eng.	(O)	-	Comment same as above except that: 0.38 ppm muscle 52.0 ppm fat	Mawdesley- Thomas and Leahy (1967)
1	DDT	Puntius puckelli	BSA	_	0.048 (T4A)	<u>a</u> cdelm	Tap water was used as diluent. Toxicity data are given as $TL_m$ 's in ppm for 24, 48, 96 hr. The pH of the water averaged at 8.3. The study was conducted in India.	Rao, et al (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)	_
DDT	Plankton Cladophora gracilis Shrimp Opsanus tau (immature) Menidia menidia Crickets Nassarius obsoletus Gasterosteus aculeatus Anguilla rostrata Flying insects, mostly Diptera Spartina patens Mercenaria mercenaria Cyprinodon	FECH	Long Island, N'. Y.	0.040 (O) 0.083 (O) 0.16 (O) 0.17 (O) 0.23 (O) 0.23 (O) 0.23 (O) 0.26 (O) 0.28 (O) 0.30 (O) 0.33 (O) 0.42 (O) 0.94 (O)	_	DDT residues in soil of the defined estuary area averaged more than 13 pounds per acre with a maximum of 32 pounds per acre. These values are based on the wet weight of the whole organism.	Woodwell, et al (1967)	APP
	variegatus Fundulus heteroclitus			1.24 (0)				APPENDIX B
	Paralich thys dentatus Esox			1.28 (O) 1.33 (O)				5
	niger Strongylura marina Spartina			2.07 (O) 2.80 (O)				
	patens			2.80 (0)				
DDT	Lampsilis siliquoidea L. ventricosa Anodonta grandis	FR	Red Cedar River, Mich.	(O)	-	The amount of the chemical sprayed in the area was not specified. Residue in plants ranged from 0.7 to 10.0 ppb. Algae contained 0.4 to 3.0 ppb. Chubs contained 2.5 to 17 ppb. Bass – 6.0 to 50.0 ppb. Clams 1.0 to 4.0 ppb. The water contained 0.002 to 0.027 ppb.	Godsil and Johnson (1968)	
DDT	Vascular plants Algae Chubs Largemouth bass Clams	FL	Tule Lake, Ore.	(0)	-	The mussels listed were analyzed for the toxicant and its metabolites. Mussels may be used as detectors for this toxicant, because they tend to concentrate the chemical in much higher concentrations than it is ever found in the water. The amount of chemical applied as a spray was not specified.	Bedford, et al (1968)	

DDT (and analogues)	Limnephilus rhombicus Sialis sp Gammarus sp Salvelinus fontinalis Semotilus atromaculatus Cottus bairdi Rhinichthys atratulus		Knights Greek, Wisc.	(O)	_	Pesticide usage in an orchard did not significantly contaminate the aquatic environment of this creek adjacent to the treatment as determined by residue analysis.	Moubry , et al (1968)
DDT	Alosa pseudoharengus Aplodinotus grunniens Coregonus artedii Lota lota	BSA	-	(O)	-	The study showed that the levels of chlorinated hydrocarbon pesticide residues in fish meals and oils were, with the exception of the oil sample taken from the Lake Michigan alewife, below the regulatory tolerances established by the Food and Drug Director- ate of Canada (1965) for certain foods intended for human consumption. Pesticide levels were interpreted as being representative for each species.	Dugal (1968)
DDT	Salmo salar L.	BCFA	-	(O)	а	Fish were conditioned for at least 60 days at 12-16 C in flowing water in the laboratory. The largest group, 20 parr with an average weight of 2.5 g, was killed in 9 hr by an aqueous suspension of 2.0 ppm p,p-DDT; 9 parr of the same size-group died in about 75 hr while in a 0.1 ppm suspension; 7 smaller parr (1.4 g) kept in a 0.01 ppm suspension, died in about 23 hours	Greer and Paim (1968)
DDT	Salvelinus fontinalis	BSA	-	(SB)	acdep	When sexually maturing yearling brook trout were fed for 156 days with DDT at sublethal rates, fish fed at the lower dosages produced more mature ova than untreated fish. Those fed at highest dosages produced fewer mature ova than untreated fish. The size of the male fish at the end of the feeding period tended to increase according to the dosage of DDT.	Macek (1968)
o,p-DDT	Potamogeton pectinatus Cladophora Oscillatoria Cynodon dactylon Arundo donax	FR	Ariz.	(O)	-	Irrigation canals were examined for plants which might serve as DDT collectors or indicators of DDT usage by concentrating this material and its metabolites. Highest residues were found in <i>Cladophora</i> (19 ppm), followed by <i>Potamogeton</i> (9 ppm), and finally <i>Oscillatoria</i> (5 ppm).	Ware, et al (1968)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
p,p'-DDT	Potamogeton pectinatus Cladophora Oscillatoria Cynodon dactylon Arundo donax	FR	Ariz.	(O)		Irrigation canals were examined for plants which might serve as DDT collectors or indicators of DDT usage by concen- trating this material and its metabolites. Highest residues were found in <i>Cladophora</i> (19 ppm), followed by <i>Potamogeton</i> (9 ppm), and finally <i>Oscillatoria</i> (5 ppm).	Ware, et al (1968)
p,p'-DDT	Alosa pseudoharengus Aplodinotus grunniens Coregonus artedii Lota lota	BSA	_	(O)	_	The study showed that the levels of chlorinated hydrocarbon pesticide residues in fish meals and oils were, with the excep- tion of the oil sample taken from the Lake Michigan alewife, below the regulatory tolerances established by the Food and Drug Directorate of Canada (1965) for certain foods intended for human consumption. Pesticide levels were interpreted as being representative for each species.	Dugal (1968)
DDT	Pteronarcys californica (naiads) Pteronarcella badia (naiads) Claasenia sabulosa (naiads)	BSA	-	0.0070 (T4A) 0.0019 (T4A) 0.0035 (T4A)	<u>a c d e f</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
DDT	Skeletonema costatum Coccolithus hyxleyi Pyramimonas sp Peridinium trochoideum	L & CH	-	(O)	<u>a</u>	Algal photosynthesis was reduced, as measured by 14CO <sub>2</sub> uptake. It was decreased at concentrations of a few ppb of DDT.	Wurster (1968)
DDT	Salvelinus fontinalis	всн	-	0 to 0.60 (SB)	<u>a e</u>	All experiments were conducted in 6 liters of water. When brook trout are exposed for 24 hr to sublethal doses of DDT, the cold-blocking temperature for a simple reflex, which shows lability related to thermal history, is altered in a way suggesting that DDT is affecting the thermal acclimation mechanism. Sublethal dosage of DDT also prevents the establishment of a visual conditioned	Anderson an Peterson (1969)

avoidance response.

Micropterus salmoides Pimephales promelas Salmo gairdnerii {one wk old sac fry } (one mo old feeding fry)	BSA BSA	_	0.5 (O) 0.5 (O) 1.0 (K 0%)	-	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the $LT_{50}$ was 48 hr and for the fathead 72 hr.	Weiss (1961)
<i>gairdnerii</i> (one wk old sac fry) (one mo old feeding fry)	BSA	_	1.0 (K 0%)			
feeding fry)			10 (K 100%)	<u>a e</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder {1962}
	BSA	-	1.0 (K 0%) 10 (K 100%)			
Bluegill	BSA	-	0.480 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
Pteronarcys sp (nymphs)	BSA	_	0.0001 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Соре (1965)
Simocephalus serrulatus Daphnia pulex	BSA	_	0.00026 (SB) 0.000066 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
Trichogaster fasciatus Chama punctatus Mastocembelus pancalus Macrognathus aculeatum Nandus nandus Rita rita Amphipnous cuchia Mystus vittatus Puntius sophore Esomus danrica Labeo rohita Sphaerodema annulatum Nepa sp Ranatra filiformis Dytiscus sp Hydrophilus sp	BSA		3 (K7) 3 (K7) 5 (K7) 5 (K7) 5 (K7) 5 (K7) 5 (K7) 10 (K7) 10 (K7) 30 (K7) 30 (K7) 0.1 (K7) 0.5 (K7) 0.2 (K7) 0.5 (K7)	а	All the organisms listed are detrimental to culture of carp. At the concentrations listed, there was 100 percent kill of the organisms in 7 days.	Srivastava and Konar (1966)
	Mastocembelus pancalus Macrognathus aculeatum Nandus nandus Rita rita Amphipnous cuchia Mystus vittatus Puntius sophore Esomus danrica Labeo rohita Sphaerodema annulatum Nepa sp Ranatra filiformis	Mastocembelus pancalus Macrognathus aculeatum Nandus nandus Rita rita Amphipnous cuchia Mystus vittatus Puntius sophore Esomus danrica Labeo rohita Sphaerodema annulatum Nepa sp Ranatra filiformis Dytiscus sp	Mastocembelus pancalus Macrognathus aculeatum Nandus nandus Rita rita Amphipnous cuchia Mystus vittatus Puntius sophore Esomus danrica Labeo rohita Sphaerodema annulatum Nepa sp Ranatra filiformis Dytiscus sp	Mastocembelus5 (K7)pancalus5 (K7)Macrognathus5 (K7)aculeatum5 (K7)Nandus5 (K7)nandus5 (K7)Rita rita5 (K7)Amphipnous5 (K7)cuchia10 (K7)Wystus10 (K7)vittatus10 (K7)Puntius10 (K7)sophore30 (K7)danrica30 (K7)Labeo30 (K7)rohita0.1 (K7)Sphaerodema0.5 (K7)Ranatra0.2 (K7)filiformis0.1 (K7)Dytiscus sp0.1 (K7)	Mastocembelus5 (K7)pancalus5 (K7)Macrognathus5 (K7)aculeatum5 (K7)Nandus5 (K7)nandus5 (K7)Rita rita5 (K7)Amphipnous5 (K7)cuchia0 (K7)Wystus10 (K7)vittatus10 (K7)Puntius10 (K7)sophore30 (K7)danrica30 (K7)Labeo30 (K7)rohita0.1 (K7)Sphaerodema0.5 (K7)Ranatra0.2 (K7)filiformis0.1 (K7)Dytiscus sp0.1 (K7)	Mascoembelus         5 (K7)           pancalus         5 (K7)           aculeatum         5 (K7)           nandus         5 (K7)           nandus         5 (K7)           nandus         5 (K7)           Amphipnous         10 (K7)           vitatus         10 (K7)           vitatus         10 (K7)           sophore         10           Esomus         30 (K7)           danrica         10 (K7)           sphaerodema         00 (K7)           rohita         10 (K7)           sphaerodema         30 (K7)           nanulatum         10 (K7)           Nepa sp         0.5 (K7)           Ranatra         0.2 (K7)           Ranatra         0.2 (K7)           Ranatra         0.2 (K7)           filiformis         10 (K7)

**B-**81

F F F

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
DDVP	Cyprinus carpio C	BSA	-	15.0 (T2A) 5.5 (T2A)	acdefp	The test animals were conditioned for 48 hours prior to use.	Sreenivasan and Swaminathan (1967)
	carpio Tilapia mossambica			3.0 (T2A)			
	Cirrhina mrigala			25.0-30.0 (T2A)			
	Labeo fimbriatus			18.0 (T2A)			
	Frog tadpoles			10.0 (T2A)			
Dead X (95 percent naphtha)	<i>Pteronarcys</i> sp (nymphs)	BSA	_	2.0 (T4A)	a	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Соре (1965)
Dead-X (EC)	Rainbow trout Bluegill	BSA	_	8.8 (T4A) 9.2 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F	Cope (1965)
Dead-X	Sim ocephalus serrulatus	BSA	_	7.60 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr.	Sanders and Cope
	Daphnia pulex			3.70 (SB)		Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	(1968)
Dead-X (95 percent naphtha)	Pteronarcys californica (naiads)	BSA	-	0.0023 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Deet	Prosimulum spp Chephia spp Simulium spp (larvæ)	LCFA	-	4.0 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time, the number detached amounted to 0.0 percent.	Jamnback and Frempong- Boadu (1966)
DEF	<i>Leiostomus</i> <i>xanthurus</i> (juvenile)	BSA	_	0.24 (O)	а	Water temperature 27 C. The figure reported is a 48-hr $\rm EC_{50}.$	Butler (1965)
DEF	Penaeus aztecus	L	-	0.028 (O)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)

	DEF	Crassostrea virginica Penaeus aztecus Penaeus duorarum Penaeus setiferus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	0.1 (O) 0.03 (O) 0.24 (T2CFA) 75% (O)	_	Seawater was pumped continuously into test aquaria.Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following: Oyster — 96-hr EC50 — Conc. which decreased shell growth.Shrimp — 48-hr EC50 — Conc. which killed or paralyzed 50% of test animals.Fish — 48-hr EC50 — Conc. which killed 50%.Phytoplankton — Percent decrease of CO2 fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	Butler (1966)
	DEF	Oyster	BCF	-	0.1 (SB4)	а	Seawater was employed in this experiment.	Butler (1965)
	DEF (tech)	Pteronarcys californica	BSA	-	0.0021 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
	Delnav	Gambusia affinis	BSA	_	0.05 (K 3%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)
1	Dełnav	Pimephales promelas	BSA	-	0.5 (O)	-	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentration but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as LT <sub>50</sub> which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. The LT <sub>50</sub> for the fathead was 72 hours.	Weiss (1961)
CON	Delnav (emulsible concentrate, 47 percent)	Pimephales promelas Lepomis macrochirus Lebistes reticulatus Largemouth bass Green sunfish	BSA	-	12.0 (T4A) 0.063 (T4A) 0.57 (T4A) 0.076 (T4A) 0.13 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)
COMMERCIAL CHEMICAL	Delnav (tech, 100 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	10.0 (T4A) 0.034 (T4A) 32.0 (T4A) 0.21 (T4A)	a c d e	Comment same as above.	Pickering, et al (1962)
	Delnav	Chaoborus astictopus	BSA	_	0.052 (T1A)	а	Toxicity value given is for the fourth instar larvae.	Hazeltine (1963)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Delnav	Bluegill	BSA	_	0.034 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Delnav	Carassius auratus Lepomis macrochirus	BSCH	_	1.0 (O)* 1.0 (O)** * no response, 15 days **response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or no response in 15 or 30 days.	Weiss and Gakstatter (1964)
Deirad	Pithophora spp Lepomis macrochirus Micropterus salmoides	FL	Ponds, Ala.	(0)	-	0.25 to 0.3 ppm of the chemical killed large amounts of <i>Pithophora</i> in ponds, but the effects were of short duration (1 to 3 weeks). As many as 4 applications of Delrad at a concentration of 0.3 ppm did not affect reproduction or production of bluegill in experimental ponds. As many as 3 applications of Delrad at the concentration of 0.3 ppm in bass brood ponds did not affect spawning of bass, hatching of eggs, or survival of fry and small fingerlings. The author states that "the minimal lethal dose" of this chemical for bluegill and large-mouth bass fingerlings is approximately 0.65 ppm. Microcrustaceans suffered approximately a 50-percent mortality when the concentra- tion of the chemical reached 0.5 ppm. The addition of equal parts of Roccal (10%) to Delrad 50-S produced no better kill of <i>Pithophora</i> in ponds, and the effects were usually of longer duration (2 to 4 weeks).	Lawrence (1958)
Delrad 70	Channel catfish (fingerlings)	BSA	-	0.74 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Deirad 70	Salvelinus fontinalis Salmo trutta Notemigonus crysoleucas	FPCH	N.Y.	0.50 (S23)	a c d	Conventional farm ponds were used having an average sur- face area of 0.3 acre and a maximum depth of 7-9 ft. Toxicity (in ppm) to fish as maximum safe concentra- tion (S) for 23 days was determined. Concentration of 0.5 ppm was required to control algae.	Eipper (1959)

Delrad 50S	Salvelinus fontinalis	FPCH	N.Y.	1.0 (S23)	acd	Comment same as above.	Eipper (1959)
	Salmo gairdneri			1.0 (S23)			
	Catostomus commersoni			0.5 (S23)			
	Notemigonus			0.25-1.0 (S23)			
	crysoleucas Ictalurus punctatus			1.0 (S23)			
	punctatus Micropterus salmoides			1.0 (S23)			
	Lepomis macrochirus			0.5 (S23)			
Demeton	Lepomis macrochirus	BSA	-	0.1 (O)	<u>acdf</u>	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The	Weiss (1959)
	Micropterus salmoides			0.1 (O)		inhibition of the enzyme was a function of the concentra- tion of the insecticide, extent of exposure, and specific	(1000)
	Notemigonus crysoleucas			0.1 (O)		chemical nature of the compound,	
	Carassius auratus			0.1 (O)			
Demeton	Carassius auratus	BSCH		1.0 (O)*	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetycholinesterase activity in the brains of fish. Concentrations are given in	Weiss and Gakstatter
	Lepomis macrochirus			1.0 (O)*		ppb as either response or not response in 15 or 30 days.	(1964)
	Notemigonus crysoleucas			1.0 (O)*			
	crysoleucas			* no response, 15 days			
Dermol	Pandalus montagni	BSA	-	148 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C.	Portmann and Connor
	Crangon crangon			156 (T2A)		It was shown that the toxicity of this solvent emulsifier decreased with time, due to evaporation of the solvent.	(1968)
	Carcinus maenas			435 (T2A)			
Į	Cardium edule edule			148 (T2A)			
Derris	Lepomis macrochirus Lepomis gibbosus Catostomus commersonii Notemigonus crysoleucas No tropis cornutus frontalis Eucalia inconstans	BSA	-	1.0 (K) 0.5 (K) 1.0 (K) 0.5 (K) 0.5 (K) 0.5 (K) 0.5 (K)	<u>a</u> d	The action of Derris root appeared to be somewhat faster in acid than alkaline waters. The derris employed in this experiment contained 5 percent rotenone.	Leonard (1938)
2	Umbra limi Carassius			0.5 (K) 0.5 (K)			
	auratus			0.0 (10)			

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Derris root (5% rotenone content)	Ambloplites rupestris Huro salmoides Perca flavescens Salmo gairdneri Salvelinus fontinalis Salmo trutta Catostomus c. commersonii Lepomis gibbosus Semotilus astromaculatus Cristivomer namaycush Umbra limi Hyborhynchus notatus Eucalia inconstans Poecilichthys exilis Fundulus diaphanus menona Notemigonus crysoleucas auratus Entosphenus Iamottenii	FL	Mich.	0.35-0.56 (O)	a	The results of this experiment indicate that in certain trout waters which are overrun with warm water fish, trout and other cold water fish can withstand an application of derris root at the concentration given, while other fish (warm water fish) can be eliminated. Derris is much less effective in colder water. It is likely that a thermocline prevents the penetration of the poison to deeper water. No trout were killed in these experiments. All other fish listed showed some mortality.	Greenbank (1940)
Derris	Tendipedidae	FL	Colo.	1.0 (S)	a c d e	This is a study of lake bottom fauna. Oligochaeta were not affected. Collection data of bottom fauna are given.	Cushing and Olive (1957)
Detergents	Pimephales promelas (juveniles)	BSA	-	(S) 61-63 (T1-4A) (H) 39-44 (T1-4A)	<u>a c d f</u>	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard water (H).	Henderson, et al (1959)
Dexon	Oyster	BCF	-	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
Dexon	Pteronarcys californica (naiads)	BSA	-	0.024 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1966)

Dexon	Penaeus aztecus	L	_	(0)	а	Toxicant chemicals were evaluated in sea water at temperatures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or	Butler (1965)
Dexon	<i>Cyprinodon</i> <i>variegatus</i> (juvenile)	BSA	_	(O)	а	mortality. No effect occurred at 1.0 ppm. Water temperature was 21 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
DFDT (DDT analogue)	Goldfish Gambusia affinis Culex apicalis (larvae)	BSA	-	0.9 (K) 0.175 (K) 0.0015 (K)	_	Experiments were run a maximum of 3 days. No other time data were reported.	Odum and Summerford (1946)
Diazinon	Daphnia magna	BSA	-	0.0043 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
Diazinon	Lepomis macrochirus Micropterus salmoides Notemigonus crysoleucas Carrasius auratus	BSA	-	0.1 (O) 0.1 (O) 0.1 (O) 0.1 (O)	<u>acdf</u>	This paper reports the effects of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The inhibition of the enzyme was a function of the concentration of the insecticide, extent of exposure, and specific chemical nature of the compound.	Weiss (1959)
Diazinon	Micropterus salmoides Pimephales promelas	BSA	-	0.5 (O) 0.5 (O)	_	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the $LT_{50}$ was 1 hr and for the fathead 80 min.	Weiss (1961)
Diazinon (EC2)	Gambusia affinis	۴L	Ponds in 111.	(O)	-	When applied at 0.3 pound per acre active ingredient, 100 percent mortality occurred in 1 day.	Mulla, et al (1963)
Diazinon	Carassius auratus Lepomis macrochirus Notemigonus crysoleucus	BSCH	-	1.0 (O)* 1.0 (O)* 1.0 (O)* *no response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss (1964)
Diazinon	<i>Pteronarcys</i> sp (nymphs)	BSA		0.025 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
Diazinon, Tech.	Rainbow trout Bluegill	BSA	-	0.090 (T4A) 0.022 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Diazinon	Salmo gairdnerii	BSA	_	0.170 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well.	Cope (1966)
	Lepomis macrochirus Pteronarcys californicus Daphnia pulex Simocephalus			0.030 (T2A)		Effects on reproduction and behavior are also dis- cussed. Data presented as EC50.	
				0.074 (T2A)			
				0.0009 (T2A) 0.002 (T2A)			
	serrulatus			0.002 (12) ()			
Diazinon	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 13 percent.	Jamnback and Frempong- Boadu (1966)
Diazinon	Simocephalus serrulatus	BSA	-	0.0018 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr.	Sanders and Cope
	Daphnia pulex			0.00090 (SB)		Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	(1966)
	Daphnia carinata		_	0.0008 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed a varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	
Diazinon	Leiostromus xanthurus	BCFH	-	0.001 (0)	а	At a concentration of .001 ppm, the following percent acetylcholinesterase activity as compared to controls	Butler and Johnson
	Cyprinodon variegatus			0.001 (O)		was found: L. xanthurus — 100 C. variegatus — 74	(1967)
Diazinon	Pteronarcys californica (naiads)	BSA	-	0.025 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Díbrom	Gambusia affinis	BSA	-	0.03 (K 3%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)
Dibrom	<i>Salmo gairdnerii</i> (one wk. old	BSA	-	1.0 (K 23%) 10.0 (K 100%)	<u>a</u> e	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
	(one mo. old feeding fry)			1.0 (K 0%) 10.0 (K 100%)			

					•			
	Dibrom	Salmo gairdneri	BSA	-	80 (T 18 hr)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)
	Dibrom (tech)	Salmo gairdneri	BSA	_	70 (T1A)	а	Comment same as above.	Cope (1963)
	Dibrom (tech)	Procambarus clarki	BSA	_	4.0 (T3A)	acdo	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
	Dibrom (tech)	Bluegill	BSA	_	0.180 (T4A)	-	The values reported are given as $LC_{50}$ .	Соре (1965)
	Dibrom	Oyster	BCF		0.1 (SB4) 1.0 (SB4)	а	Sea water was employed in this experiment.	Butler (1966)
	Dibrom	Simocephalus serrulatus Daphnia pulex	BSA	_	0.0011 (SB) 0.00035 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Dibrom	Leiostromus xanthurus Cyprinodon variegatus Mugil cephalus	BCFCH	-	0.05 (O) 0.05 (O) 0.001 (O)	а	At a concentration of 0.05 or 0.001 ppm, the following percent acetylcholinesterase activity as compared to controls was found: <i>L. xanthurus</i> – 10 <i>C. variegatus</i> – 79 <i>M. cephalus</i> – 76	Butler and Johnson (1967)
	Dicamba (Banvel D)	Oncorhynchus kisutch Salmo gairdneri	BSA	_	151 (T1A) 120 (T2A) 320 (O)	a c d	The active ingredient of Dicamba is 2-methoxy-3, 6- dichlorobenzoic acid (dimethylamine salt). Tests were conducted in glass jars holding 15 liters of water. Concentrations of 320 ppm produced no mortalities in rainbow trout.	Bond, et al (1965)
	Dicamba (Banvel D)	Rainbow trout Bluegill	BSA	_	35.0 (T2A) 130.0 (T2A)	_	Data are given as LC <sub>50</sub> .	Bohmont (1967)
	Dicapthon	Chaoborus astictopus	BSA	-	0.0057 (T1A)	a	Toxicity value given is for the first instar larvae.	Hazeltine (1963)
COMMEDO	Dichlobenil	Lepomis macrochirus	BSA	_	17.0 (T2A) L 30.0 (T2A) G	<u>a</u> cdef	Toxicity data for 24 and 48 hours are presented for liquid (L) and granular (G) formulations. Various commercial formulations were tested. The liquid formulations were almost invariably more toxic than	Hughes and Davis (1965)

the granular ones,

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)
Dichlobenil	Salmo	BSA	-	22.0 (T2A)	а	This paper reports acute toxicity of a number of com- pounds, and discusses sub-acute mortality as well.	Cope (1966)
	gairdnerii Lepomis macrochirus			20.0 (T2A)		Effects on reproduction and behavior are also discussed.	(1900)
	Pteronarcys californicus			8.4 (T2A)		Data presented as $EC_{50}$ .	
	Daphnia pulex			3.7 (T2A)			
	Simocephalus serrulatus			5.8 (T2A)			
Dichlobenil (Casoran)	Daphnia magna	BSA	-	9.8 (8.8-10.7) (O)	acdiq	Toxicity, in terms of median immobilization concentration $(IC_{5\Omega})$ , is presented for <i>Daphnia</i> ; median lethal concentra-	Crosby and Tucker
(0030/0//	Rainbow trout			22 (O)		tion (LC50) values for rainbow trout and bluegill are reported.	(1966)
	Bluegill			20 (O)			
Dichlobenil	Lecomis macrochirus (eggs)	L	-	20 (S), 10 (NTE)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days sur-	Hiltibran (1967)
	L. cyanellus (eggs)			25 (NTE)		vival. Maximum length of test was 8 days. No food was was added. Small bluegill were tested to find the highest	
	(eggs) Micropterus dolomieui (eggs)			25 (NTE)		concentration of chemical which did not cause death in 12 days (S).	
	Erimyzon sucetta (eggs)			25 (NTE)			
	L. macrochirus (fry)			20 (S)			
Dichlobenil	Pteronarcys californica	BSA	-	0.007 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Dichlone (Phygon)	Daphnia magna	BSA	-	0.014 (O)	acdiq	Toxicity, in terms of median immobilization concentration (IC50), is presented for <i>Daphnia;</i> medium lethal concen-	Crosby and Tucker
(, ,,,ge,,,,	Bluegill			0.04 (O)		tration (LC50) values for bluegill are reported.	(1966)
Dichlorvos	Prosimulum spp Cnephia spp Simulium spp (Iarvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 41 percent.	Jamnback and Frempong- Boadu (1966)
Dichlorvos	Pteronarcys californica (naiads)	BSA	-	0.001 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1966)

Diethanol rosinamine D acetate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI $\sim$ T Ma $\sim$ T So $-$ T (3), PT (21) Cv $-$ T (7), PT Gp $-$ T Np $-$ T	Palmer and Maloney (1955)
DIDT (DDT analogue)	Goldfish Gambusia affinis Culex apicalis (larvae)	BSA	-	0.175 (K) 0.025 (K) 0.003 (K)	-	Experiments were run a maximum of 3 days. No other time data were reported.	Odum and Sumerford (1946)
Dieldrin	Carassius auratus Lepomis macrochirus	BSA	-	4.0 (K) 2 hr	_	The toxicity threshold for the 2 species was 0.031 ppm. Water taken from rice field that had been treated with 0.1 pound of Dieldrin per acre, 10 to 20 days previously, killed fish unless diluted 1/2 with pond water.	Gray (1950)
Dieldrin (hexa- chloroepoxy- octahydrodi- methanonaptha- lene, 25 percent)	Lymnaeid snails	BSA	-	(0)	а	Each test container, 500-ml beaker, was filled with ditch water. 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)
Dieldrin (granules)	Simulium sp (Iarvae)	FR	Streams, S. C.	0.04 (0)	_	Dieldrin at a 0.04 lb/acre rate of application controlled blackfly larvae for up to 4 weeks. Data are presented as percent larval detachment for this period of time.	Davis, et al (1957)
Dieldrin	Artemia salina	BSA	-	1.172 (L 1)	ai	Rock salt was used in rearing all cultures employed in bioassay work. The optimum salt concentration was 3.5%.	Tarpley (1958)
Dieldrin	Fish Crustacea fiddler crabs, etc. Mollusks snails oysters, etc.	FO (salt marsh)	Fla.	(O)	-	Two thousand acres of marsh were treated for sandfly control at rate of 1 lb/acre. Fish reacted to the chemical within a few minutes after treatment (list of fish names given). An estimated 20-30 tons of fish of about 30 species died. Crustaceans, mollusks seemed to be un- harmed. Fish repopulation began after the 4th week and was climactic at the 10th week.	Harrington and Bidlingmayer (1958)
Dieldrin	Fathead minnow Bluegill Goldfish Guppy	BSA	-	0.016 (T4A) 0.0079 (T4A) 0.037 (T4A) 0.022 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in soft water.	Tarzwell (1959)

COMMERCI	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
COMMERCIAL CHEMICAL PRODUCTS	Dieldrın	Fathead minnow	BSA	_	0.016 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish" It is interesting that the dif- ferent tables from the book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in soft water.	Tarzwell (1959)
CTS	Dieldrin (dust)	Tilapia melanopleura	FLCH	Tangan- yika	1 lb (3.3% K) 5 lb (6.6% K - 3 wks) 10 lb (66.6% K - 3 wks)	-	Trial periods were for 20 weeks. Sublethal effects such as impaired breeding, retarded growth, or altered taste were not detected. Dosages are given as Ib/acre of surface water.	Webb and Shute (1959)
	Dieldrin (granules)	Tilapia melanopleura			1 Ib (0% K)	-	Comment same as above.	Webb and Shute (1959)
	Dieldrin 50	Channel catfish (fingerlings)	BSA	-	2.5 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Dieldrin	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	0.01 (T4A) 0.01 (T4A) 0.04 (T4A) 0.02 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
	Dieldrin	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	0.018 (T4A) 0.0088 (T4A) 0.041 (T4A) 0.025 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et al (1959)
	Dieldrin	Pimephales promelas Lepomis macrochirus	BSA	-	0.016 (T4A) 0.0079 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)

Dieldrin	Pimephales promelas Lepomis macrochirus	BSA	Cincin- natí, O.	0.1 to 5.0 (T4A) 0.0056-0.042 (T4A)	acdefp	Toxicity of run-off water from areas treated with Dieldrin was evaluated. Three different Dieldrin formulations were used: powder, emulsion, and acetone solution. The acetone formulation was generally the most toxic.	Tarzwell and Henderson (1960)
Dieldrin	Daphnia magna	BSA	-	0.33 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
Dieldrin	Oncorhynchus kisutch Oncorhynchus tshawytscha Salmo gairdnerii Gasterosteus aculeatus	BSA	-	10.8 (T4A) 6.1 (T4A) 9.9 IT4A) 15.3 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
Dieldrin	Salmo gairdneri	BSA	_	0.0355 (T1A) 0.0233 (T2A) 0.0233 (T4A)	<u>a</u> cdfg	Hatchery artesian well water was employed for this experiment.	Webb (1961)
Dieldrin	Gammarus Iacustris Iacustris	BSA	_	(O)	a e p	The mortality might have been partially due to the susceptibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Results were expressed as $LT_{50}$ ; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 74 ( $\pm$ 7) min.	McDonald (1962)
Dieldrin	Ophicephalus punctatus Heteropneustes fossilis Barbus stigma Trichogaster fasciatus	BSA	_	4000-8000 (K ≪4 hr) 2000-8000 (K ≪9 hr) 4000 (K ≪3 hr) 2000-4000 (K ≪4 hr)	а	The dosage to produce toxic symptoms varied with each species. At the very low dosage, these insecticides did not produce observable changes, but at the higher dosage changes were pronounced.	Mathur (1963)
Dieldrin (EC 1.5)	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a c	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and re- sulted in a 100 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
Dieldrin	Gambusia affinis affinis	BSA	-	0.016 to .50	а	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are later placed in clean water.	Boyd and Ferguson (1964)
Dieldrin	Lepomis gibbosus	BSA & CH	-	0.0067 (T4A)	<u>a c d</u> e	Other medium tolerance limits were: 0.0155 ppm — 24 hours 0.012 ppm — 48 hours 0.0075 ppm — 72 hours. Chronic exposure to 0.00168 ppm for the period of 12 weeks affected the oxygen consumption and the cruising speed ability.	Cairns and Scheier (1964)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Dieldrin	Notemigonus crysoleucas Lepomis macrochirus L. cyanellus	BSA	-	(B) 0.025 (T 1.5) (A) 0.90 (T 1.5) (B) 0.025 (T 1.5) (A) 0.900 (T 1.5) (B) 0.033 (T 1.5) (A) 1.25 (T 1.5)	<u>a</u> cf	Chemical was dissolved in acetone. Final concentration of acetone was $\leq 2 \text{ ml/l}$ . Data shows $TL_m$ in ppb for insecticide-resistant and insecticide non-resistant strains of the test fish.	Ferguson, et al (1964)
Dieldrin (tech, 100 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica	BSA	-	0.03900 (T4A) 0.02400 (T4A)	c d e f	A. pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides that P. californica.	Jensen and Gaufin (1964)
Dieldrin	Gammarus Iacustris	BSA	-	0.70 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Dieldrin	Bluegill	BSA	-	0.0079 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Dieldrin (tech)	Rainbow trout Bluegill	BSA	-	0.013 (T4A) 0.0028 (T4A)	~	The values reported are given as $LC_{50}$ .	Cope (1965)
Dieldrin	Bluegill	BSA	-	16 (T4A) 18 (T4A) 14.5 (T4A) 9.3 (T4A) 7.1 (T4A)	<u>a</u>	These experiments were performed to demonstrate that at increased temperatures the toxic effect of most chemicals is increased. For the toxicant concentrations listed, the temperatures were respectively, 45, 55, 65, 75, and 85 F Data on the effect of time as well as temperature was also reported. The experimental animals all were approximately one grain in weight.	Cope (1965)
Dieldrin	Palaemo netas kadiakensis	BSA	-	(N) 50.0 (T1-1/2A) (TB) 135.0 (T1-1/2A)	acf	Test organisms were collected from 2 locations, Twin Bayou (TB), Sunflower Co., Miss. (agricultural area) and Noxubee National Wildlife Refuge (N), Noxubee Co., Miss. (non- agricultural area) and evaluated in laboratory bioassays. The Twin Bayou shrimp were more resistant.	Ferguson, et a (1965)
Dieldrin	Gambusia affinis Ictalurus melas	BSA	_	0.001-0.025 (T3A) 0.003-0.028 (T3A)	<u>a</u> cde	Test fish were collected from 8 different locations of the Mississippi River. The 3-day TL <sub>M</sub> values were made to determine if a resistance gradient existed. The data indicated that there was none.	Ferguson, et a (1965)

	Dieldrin	Pteronarcys calífornica	BSA	-	0.04 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were determined by standard methods. General comments	Gaufin, et al (1965)
		Acroneuria pacifica			0.02 (T4A)		were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional	
		Ephemerella			0.008 (T4A)		data are presented.	
		grandis Daphnia			0.3 (T 50 hr A)			
		magna Gammarus Iacustris			0.7 (T4A)			
	Dieldrin	Acroneuria pacifica	BSA	_	0.024 (T4A)	a c	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
		Ephemerella grandis			0.008 (T4A)			
		Gammarus lacustris			0.7 (T4A)			
		Pteronarchys californica			0.039 (T4A)			
	Dieldrin	Carassius carassius	BSCH	-	0.1 (SB-2 hrs) 0.075 (SB-4 hrs)	_	Data given on chemical residue found in tissue computed from C <sup>14</sup> activity was: 37.2 mg/g in blood 10.5 mg/g in muscle.	Weiss (1965)
)	Dieldrin	Poecilia reticulata	BSA		0.021 (T4A)	ai	Light was controlled in this experiment. All tests were conducted in soft, synthetic dilution water.	Cairns and Loos (1966)
•	Dieldrin	Salmo gairdneri	BSA	-	0.005 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute ntoxicity as well. Effects on repro-	Cope (1966)
		Lepomis macrochirus			0.006 (T2A)		duction and behavior are also discussed. Data presented as $EC_{50}$ .	
		lctalurus punctatus			0.025 (T2A)		50-	
		Pteronarcys californicus			0.001 (T2A)			
		Baetis sp			0.064 (T2A)			
		Daphnia pulex			0.250 (T2A)			
8		Simocephalus serrulatus			0.250 (T2A)			
COMMERCIAI	Dieldrin	Salmo gairdnerii	BCFA	-	0.016 (K1)	_	The gills, muscles, and livers of the fish were examined for concentrations of the chemical. The trout tended to accumulate this compound in all tissues studied.	Holden (1966)
IAL CHEMIC	Dieldrin	Acroneuria pacifica Pteronarcys californica	BSA & CFCH	_	0.024 (T4A) 0.0002 (T30A) 0.039 (T4A) 0.002 (T30CH)	acde	Additional data are presented.	Jensen and Gaufin (1966)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL PRODUCTS	Dieldrin	Platalea leucorodia Haematopus ostralegus Sterna sandvicensis Sterna Hirundo Larus ridibundus Somateria mollissima Tadorna tadorna	FO	Netherlands	(0)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecticides. In most cases the liver had the highest concentration of toxicant, ranging from a trace to 9.5 ppm of endrin. Birds feeding on predominantly crustacea, molluscs, and fish contained significant amounts.	Koeman and van Genderen (1966)
R OK	Dieldrın	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto alba Strix aluco Osio otus Falco pereginus	FO	Netherlands	(O)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecticides. In most cases the liver had the highest concentration of toxicant, ranging from 0.4 to 44.0 ppm. Most chlorinated hydrocarbons tended to accumulate in the fat depots of the body. In instances where mesenterial fat was analyzed the concentration of toxicant was found to be as high as 17.0 ppm.	Koeman and van Genderen (1966)
	Dieldrin	Daphnia magna Daphnia carinata Simocephalus serrulatus	BSA	-	0.740 (SB) 0.250 (SB) 0.240 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
		Daphnia pulex			0.250 (SB)			NATE 1.
	Dieldrin (20% active)	Tubifex spp Limnodrilus spp	BSA	_	6-71 (T4A)	a c e	Toxicity is reported as the mean lethal dose (LD <sub>50</sub> ) for 24, 48, and 96 hours.	Whitten and Goodnight (1966)

Dieldrin	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	-	(0)	-	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentra- tion of 700-1800X resulted.	Butler (1967)
Dieldrin	Poecilia reticulata	всн	_	0.0018 (O) 0.0056 (O) 0.01 (O)	<u>a</u>	The three levels of toxicant reported are near "the estimated biologically safe concentration" for acute exposure of gup- pies to dieldrin. The period of exposure was fourteen months, during which time, conditions of "food, water, temperature, and photo period" were controlled. At the higher concentration, there was suggestive evidence that long-term exposure to Dieldrin had a deleterious effect on the reproductive process.	Cairns, et al (1967)
Dieldrin	Oyster	FE	Galveston Bay, Texas	(0)	~	Oysters from the area were found to contain $<0.01$ to 0.01 ppm.	Casper (1967)
Dieldrin	Goldfish	BSA & CH	-	50 mµg/mi (K)	~	Test fish were conditioned to alkyl benzene sulfonate $(4 \ \mu g/m)$ , or sodium lauryl sulfate $(4 \ \mu g/m)$ , for various periods of time, then exposed to pesticides. Chronic exposure to the detergent increased the toxicity of the pesticide.	Dugan (1967)
Dieldrin-C <sup>14</sup>	Lepomis macrochirus Carassius auratus	BCFCH	-	(0)	а	Fish were treated with carbon-labeled insecticides (0.03 ppm) from 5 to 19 hr and uptake rates were determined. They were placed in recovery tanks for up to 32 days. Whole body samples were then made. It was found that of the dieldrin absorbed, >90% was eliminated from the fish after 2 weeks.	Gakstatter and Weiss (1967)
Dieldrin	Lebistes reticulatus	BSA	-	(O)	<u>a</u> cei	The median lethal concentration for a 2 hour exposure to Dieldrin for guppies was approximately 0.05 ppm. The median concentration for Harlequin fish ( <i>Rasbora hetero- morpha</i> ) and trout ( <i>Salmo</i> sp) was approximately 0.01 ppm. This short exposure to dieldrin for guppies up to the median lethal concentration had no harmful effects on the reproduction of surviving guppies.	Hubble and Reiff (1967)
Dieldrin	Salmo spp Lebistes reticulatus Rasbora heteromorpha	BSA & CH	-	(0)	<u>a</u> cdefi	The median lethal concentration for a 2 hour exposure to Dieldrin for trout was approximately 0.01 ppm. Reproduc- tive capacity of surviving trout could not be determined. The median lethal concentration for a 2 hour exposure to Dieldrin for guppies was approximately 0.05 ppm. Reproduc- tive capacity of surviving guppies was apparently unaffected. The median lethal concentration for a 2 hour exposure to Dieldrin for Harlequin fish was approximately 0.01 ppm. Reproductive capacity of the surviving fish could not be determined.	Hubble and Reiff (1967)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Dieldrin	Esox Iucius	FR	River Nene, Eng.	(0)	_	Higher concentrations were found in larger fish, indicating that they had been exposed to the pesticides for a longer time. Tissue extracts from the pike were analyzed for organochlorine pesticide residues by gas liquid chroma- tography. The values for large pike were: 0.24 ppm muscle 28.0 ppm fat	Mawdesley- Thomas and Leahy (1967)
Dieldrin	Navicula seminulum var. Hustedtii	BSA	_	12.8 (T5A)	а	This diatom species survived concentrations of Dieldrin considerably greater than those reported for fish and aquatic invertebrates. Fish feeding on these algae could receive lethal amounts of Dieldrin.	Cairns (1968)
Dieldrin	Limnephilus rhombicus Sialis sp Gammarus sp Salvelinus fontinalis Semotilus atromarulatus Cottus bairdi		Knights Creek, Wisc.	(O)	_	Pesticide usage in an orchard did not significantly contami- nate the aquatic environment of this creek adjacent to the treatment as determined by residue analysis.	Moubry, et al (1968)
Dieldrın	Alosa pseudoharengus Aplodinotus grunniens Coregonus artedii Lota lota	BSA	-	(O)	_	The study showed that the levels of chlorinated hydrocarbon pesticide residues in fish meals and oils were, with the exception of the oil sample taken from the Lake Michigan alewife, below the regulatory tolerances established by the Food and Drug Directorate of Canada (1965) for certain foods intended for human consumption. Pesticide levels were interpreted as being representative for each species.	Dugal (1968)
Dieldrın	Poecilia reticulata	BSA	_	(O)	а	The paper describes a method for continuously producing a supply of an aqueous solution of Dieldrin by passing water at a constant rate through a column of sand coated with the insecticide. The concentration of HEOD, the active ingredient of Dieldrin, was nearly constant over a period of several months, but the toxicity of the water declined steeply during the first few weeks until a relatively stable level was attained.	Chadwick and Kiigemagi (1968)
Dieldrin	Pteronarcys californica (naiads)	BSA	-	0.0005 (T4A)	<u>acdef</u>	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
	Pteronarcella badia (naiads)			0.0005 (T4A)			(1900)
	<i>Claasenia</i> <i>sabulosa</i> (naiads)			0.00058 (T4A)			

Difo	latan	Oyster	BCF	_	0.034 (O)	а	The value reported is a 96-hour EC <sub>50</sub> (decreased shell growth).	Butler (1965)
		<i>Fundulus similus</i> (juvenìle)	BSA		0.032 (O)	а	Water temperature was 20 C. The figure reported is a 48-hr EC <sub>50</sub> .	
{N- tetu thio he>	latan -(1, 1, 2, 2- rachlorethyl- o)-4-cyclo- kane-1,2-di- boximide]	Brachydanio rerio	BSA	-	1 (0)	<u>a</u>	At 1 ppm all larvae were killed within 48 min. The TL <sub>50</sub> was 34 min. LC <sub>50</sub> was 0.21 ppm.	Abedi and Turton (1968)
Difo	latan	Pteronarcys californica (naiads)	BSA	_	0.0004 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Dila	n	Channel catfish (fingerlings)	BSA		0.5 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Dila	n	Gambusia affinis	BSA	-	0.5 (L1)* 0.4 (L1)** * Resistant fish **Nonresistant fish	а	This paper deals with the resistance of mosquito fish to chlorinated hydrocarbon compounds. Resistant fish were not always less sensitive to these chemicals.	Boyd and Ferguson (1964)
Dim	ecron	Cyprinus carpio	BSA	_	51.5 (T2A)	acdefp	The test animals were conditioned for 48 hours prior to use.	Sreenivasan and Swaminathan (1967)
Dim	ethoate	Anopheles quadrimaculatus Aedes aegypti	BSA	-	3.5 (O) 4.0 (O)	-	<ul> <li>4th instar larvae of mosquitos were used in this bioassay.</li> <li>At the indicated concentrations, the following mortalities occurred:</li> <li>Anopheles quadrimaculatus 79%</li> <li>Aedes aegypti 29%</li> <li>Adsorption was determined by use of P<sup>32</sup> labeled dimethoate.</li> </ul>	Schmidt and Weidhaas (1961)
Dim	ethoate	Pteronarcys sp (nymphs)	BSA	-	0.043 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC50.	Cope (1965)
Dim (te	ethoate ch)	Bluegill	BSA		6.0 (T4A)	_	The values reported are given as $LC_{50}$ .	Cope (1965)
	ethoate	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	_	4.0 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxication. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 4 percent.	Jamnback and Frempong- Boadu (1966)
Dim Dim	ethoate	Pteronarcys californica (naiads)	BSA		0.043 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Dimethrin	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	_	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 89 percent.	Jamnback and Frempong- Boadu (1966)
Dimethyl urea	Salmo gairdnerii	BSA	-	975 (T1A) 925 (T2A) 180 (T1A)* 100 (T2A)* *with adjuvant	<u>a</u> e	Most of the weed-killer formulations in this study consisted of more than one substance, i.e., oils, emulsifiers, stabilizers, and other adjuvants.	Alabaster (1956)
Dimetilan	Prosimulum spp Cnephia spp Simulium spp (Iarvae)	LCFA	_	4.0 (O)	а	Stones heavily populated with with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 2 percent.	Jamnback and Frempong- Boadu (1966)
Dimeton	Pimephales promelas	BSA	-	0.5 (O)	_	The degree of reaction to the cholinesterase-inhibiting insecti- cides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. The $LT_{50}$ for the fathead was 72 hr.	Weiss (1961)
Dipterex	Rainbow trout Eastern brook trout	BCFA	-	1-10 (K 0%)	<u>a</u>	Spring water (46 F) was used. The flow rate was 10 GPM. The chemical was added by continuous drip dispenser. 0.02 ppm for 180 hr showed toxic effects, but no kill.	Hoffman (1957)
Dipterex	Fathead minnow	BSA	_	51 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish" It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959)
Dipterex	Pimephales promelas	BSA	-	180 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et (1959)

	Dipterex	Pimephales promelas	BSA	_	180 (T4A)	<u>a</u>	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)
	Dipterex	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	-	1000 (К) 500 (К) 500 (К) 500 (К) 100 (К)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants.	Ukeles (1962)
	Dipterex	Chaoborus astictopus	BSA	_	0.60 (T1A)	<u>a</u>	Toxicity value given is for the fourth instar larvae.	Hazeltine (1963)
	Dipterex	Bluegill	BSA	_	3.8 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
	Dipterex	Daphnia magna	BSA	_	0.00012 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Dipterex (99%)	Pimephales promelas	BSA		180 (T4A)	<u>a c d e f</u>	Tests were performed in both hard and soft water. Additional tolerance limit values are given.	Henderson and Pickering (1958)
	Dipterex	Daphnia carinata Simocephalus serrulatus Daphnia pulex	BSA	_	0.00025 (SB) 0.00070 (SB) 0.00018 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
COMME	Diquat	Onchorynchus tshawytscha	BSA	-	29.5 (T1A) 28.5 (T2A)	acde	Concentrations were based on percent active ingredient.	Bond, et al (1960)
MERC	Diquat	Salmo gairdneri	BSA	-	(0)	а	No mortality was noted with concentrations of 10,000 mg/1 at 55 F for 100 hr. Fish were 2-3 in. long.	Cope (1963)

CHEMICAL PRODUCTS B-102	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
CHEMICAL	Diquət (1 1'-ethylene- 2:2'-depyridylium dibromide)	Elodea canadensis Potamogeton	BSA	-	5 (O) 100 (O) 5 (O)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. Injury rating of 8.5. Injury rating of 9.0. (njury rating of 7.5.	Frank, et al (1961)
DUCTS		nodosus Potamogeton pectinatus			100 (O) 5 (O) 100 (O)		Injury rating of 9.0. Injury rating of 8.8. Injury rating of 9.4.	
	Diquat	Lepomis macrochirus Pimephales promelas Micropterus salmoides	BSA	_	140 (T4A) H 72 (T4) S 130 (T4A) H 14 (T4) S 78 (T4A) S	<u>a</u> c <u>e</u>	Bioassay methods in Standard Methods for examination of water was used. Both hard (H) and soft (S) water were used. TL <sub>m</sub> values for 24 and 48 hr are also presented.	Surber and Pickering (1962)
	Diquat	Micropterus salmoides (fry)	BSA	-	1.0 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
		Ictalurus punctatus (fry) Lepomis			10.0 (SB3) 4.0 (SB3)			
15		macrochirus (fry)						
	Diquat	Lemna minor Spirodela polyrhyza Wolffia columbiana	FL	Fla.	(O)	_	Common duckweed and watermeal in small ponds can be controlled with diquat at rates as low as 0.25 ppmw, but rates greater than 0.5 ppmw are required for control in ponds infested with watermeal.	Blackburn and Weldon (1965)
	Diquat	Spirodela polyrhyza	BSA	-	(O)	<u>a</u>	0.01 ppm caused 80% chlorosis in 7 days.	Blackburn and Weldon
		Lemna minor					0.01 ppm caused 90% chlorosis in 7 days.	(1965)
		Wolffiella floridana					0.01 ppm caused 72% chlorosis in 7 days.	
		Azolla					0.01 ppm caused 50% chlorosis in 7 days.	
		caroliniana Wolffia					0.01 ppm caused 3% chlorosis in 7 days.	
		columbiana					Light intensity was kept at 500 foot-candles for 14 hours per day. Light has been shown to increase the rate of kill with diquat. Test containers were plastic petri dishes.	

Diquat (1,1'-ethylene- 2,2'-dipyridy- lium dibromide)	Plankton Oligochaeta Chironomids	FL	Fla.	0.5 (O)	_	Water samples from ponds taken at 3 and 11 days after application showed concentrations of diquat at 0.25 ppm and 0.001 ppm, respectively. No diquat was present after 16 days. Plankton appeared to be adversely affected by 0.5 ppm of diquat, but recovered rapidly. <i>Oligochaeta</i> showed a subtle chronic sensitivity to diquat.	Tatum and Blackburn (1965)
Diquat	Lepomis macrochirus Micropterus salmoides Pimephales promelas Ictalurus punctatus Salmo gairdneri	BSA	-	9-10 (L10) 10 (L10) 10 (L10) 10 (L10) 5 (L10)	- - - -	Toxicity to fish was determined as the threshold concentra- tion (LD10) in 96 hr at 75 F (65 F for trout). Herbicidal evaluations are also presented.	Lawrence, et al (1965)
Diquat	Oyster <i>Fundulus similus</i> (juvenile)	BCF BSA	-	(O) (O)	a ð	No effect on exposure to the chemical at 1.0 ppm. Water temperature was 19 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Diquat	Penaeus setiferus	L	_	(0)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
Diquat	Crassostrea virginica Penaeus setiferus Fundulus similis Phytoplankton	BCFA & BSA	-	1.0 (NTE) 1.0 (NTE) 1.0 (NTE) 45% (O)	-	Seawater was pumped continuously into test aquaria.Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:Oyster -96-hr EC50 - Conc. which decreased shell growth.Shrimp -48-hr EC50 - Concl which killed or paralyzed 50% of test animals.Fish -48-hr EC50 - Conc. which killed or paralyzed 50% of test animals.Fish -48-hr EC50 - Conc. which killed 50%.Phytoplankton -Percent decrease of CO2 fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	Butler (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Diquat	Ephemeroptera Caenidae Odonata Coenagrionidae Hemiptera Hebridae Mesoveliidae Gerridae Veliidae Pleidae Belostomaladae Corixidae Coleoptera Halipidae Dytiscidae Diptera Chironomidae Amphipoda Talitridae Basommatophora Planorbidae Physidae Copepoda Ostracoda <i>Eucalia</i> <i>inconstans</i> Salvelinus fontinalis	FL	Price Co., Wisc.	(O)	-	An application of 1.0 ppm of Diquat was made to control a nuisance weed, <i>Elodea canadensis</i> , in a pond. The effect of this treatment on aquatic insects and related animals was monitored. At 1.0 ppm, Diquat appeared to be harmless to the aquatic fauna, but caused sharp changes in the numbers of most arthropods and mollusks by destroying their habitats. Organisms affected to some degree are recorded in the organisms column.	Hilsenhoff (1966)
Diquat	Salmo gairdneri Lepomis macrochirus	BSA	_	20.000 (T2A) 19.000 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute toxicity as well. Effects on repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Cope (1966)
Diquat	Daphnia magna	BSA	-	7.1 (6.3-8.0) (O)	acdiq	Toxicity, in terms of median immobilization concentration (IC <sub>50</sub> ), is presented.	Crosby and Tucker (1966)

Diquat	Potamogeton crispus P. foliosus P. pectinatus P. pusillus Myriophyllum exalbescens Ranunculus trichophyllus Elodea canadensis Ceratophyllium demersum Najas flexilis Cabomba caroliniana Typha latifolia T. angustifolia Justicia americana J. repens var. glabrescens Sagittaria latifolia Scirpus acutus	FL	Ponds, Central Illinois	0.5 (K) 0.5 (K) 0.5 (K) 0.5 (K) 0.5 (K) 1.0 (K) 1.0 (K) 1.0 (K) 4.0 (NTE) 25.0 (O) 25.0 (O) 25.0 (O) 25.0 (O) (O)		Liquid formulations containing two pounds of the Diquat cation per gallon as the dibromide salt were used in all experiments. Most applications were based on the Diquat cation content in ppm, but some rates were expressed as the volume of herbicide per gallon of water, e.g., the 25.0 rate of application resulted in a kill by a foliage application of 25 ml of chemical diluted to 1 gallon with water. The paper states that recently <i>Scirpus acutus</i> was killed by a foliage application containing 10 ml of Diquat and 3 ml of X-77, a non-ionic wetting agent, diluted to 1 gallon with water.	Hiltibran (1967)
Diquat Diquat	Salmon Lepomis macrochirus Carassius auratus Esox lucius Salmo gairdnerii Stizostedion vitreum vitreum Daphnia pulex Cladocera Lepomis macrochirus	BSA BSA BSCH	_	28.5 (T2A) 35 (T4A) 35 (T4A) 16 (T4A) 11.2 (T4A) 2.1 (T4A) 3 (K8) (O) (O)	— acdfilmnp acdfimp	<ul> <li>Data are given as LC<sub>50</sub>.</li> <li>The herbicide showed an acute toxicity to <i>Cladocera</i>. At 3 ppm the <i>Cladocera</i> population was reduced to a level of 102 as opposed to a maximum in the control of 150.</li> <li><i>Elodea canadensis</i> was used as an indicator of herbicidal activity and to provide a nearly natural habitat in artificial ponds. Fingerling and adult bluegills were used in this experiment.</li> <li>The pools were treated with 1 and 3 ppm of Diquat at various frequencies. No fish kill occurred. After 24 weeks no residues of the chemical were found in the bluegills.</li> </ul>	Bohmont (1967) Gilderhus (1967) Gilderhus (1967)

.

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)	
AL CHEMICAL PRODUCTS	Dıquat	Lepomis macrochirus (eggs) Micropterus dolomieui (eggs) Erimyzon sucetta (eggs) L. macrochirus (fry)	L	_	2.5/3 (O) 2.5/1 (O), 1.3/4 (O) 2.5/2 (O), 1.3/2 (O) 10 (S)	_	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)	
ſS	Di-Syston	Pimephales promelas Lepomis macrochirus	BSA	_	3.7 (T4A) 0.064 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1959)	
	Di-Syston	Pimephales promelas Lepomis macrochirus	BSA	-	3.7 (T4A) 0.064 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly in- fluenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)	APF
B-106	Di-Syston	Micropterus salmoides Pimephales promelas	BSA	-	0.5 (O) 0.5 (O)	_	The degree of reaction to the cholinesterase-inhibiting insecti- cides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the $LT_{50}$ was 24 hr and for the fathead 72 hr.	Weiss {1961}	APPENDIX B
	Di-syston (tech, 90 percent)	Carassius auratus Lebistes reticulatus	BSA	-	7.2 (T4A) 0.28 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. $TL_m$ 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)	
	Di-syston, (emulsible concentrate, 20 percent)	Lepomis macrochirus	BSA	_	0.082 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)	
	Di-Syston (tech, 89 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica (naiad)	BSA	-	0.0285 (T4A) 0.0082 (T4A)	c d e f	A pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than P. californica.	Jensen and Gaufin (1964)	

Di-Syston	Gammarus Iacustris	BSA	-	0.24 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Di-Syston	Bluegill	BSA	_	0.063 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Di-Syston	Bluegill	BSCH	_	1.0 (O)* *response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
Di-Syston	Pteronarcys californica Acroneuria pacifica	BSA		0.03 (T4A) 0.008 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were determined by standard methods. General comments were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional	Gaufin, et al (1965)
	Ephemerella grandis Gammarus			0.08 (T4A) 0.2 (T4A)		data are presented.	
	<i>lacustris</i> Bluegill sunfish			0.07 (T4A)			
	Fathead minnow			4.1 (T4A)			
Disulfoton	Pteronarcys californica (naiads)	BSA	_	0.005 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
Dowacide A	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis	BSA	_	100 (NG) 100 (NG) 100 (NG) 100 (K) 50 (K)	<u>a</u>	This paper concerns the growth of pure cultures on marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
Diuron	lutheri Oncorhynchus kisutch Micropterus salmoides	BSA	-	33 (T1A) 16 (T2A) (O)	a c d e	Concentrations were based on percent active ingredient. Low Low toxicity occurred with <i>M. salmoides</i> in the solubility range of this compound.	Bond, et al (1960)
Diuron	Protococcus sp Chlorella sp Dunaliella euchlora	L	_	0.004 (K) 0.04 (NG) 0.004 (NG)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no odded toxicants to optical density in the basal	Ukeles (1962)
	Phaeodactylum tricoruntum Monochrysis lutheri			0.004 (K) 0.00002 (K)		with no added toxicants. NG = no growth, but the organisms were viable.	
Diuron	Penaeus aztecus	L	_	(0)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Diuron	Crassostrea virginica Penaeus aztecus Mugil cephalus Phytoplankton	BCFA & BS/	A –	1.8 (O) 1.0 (NTE) 6.3 (T2A) 87% (O)	-	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster - 96-hr EC<sub>50</sub> - Conc. which decreased shell growth.</li> <li>Shrimp - 48-hr EC<sub>50</sub> - Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish - 48-hr EC<sub>50</sub> - Conc. which killed 50%.</li> <li>Phytoplankton - Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Diuron (tech)	Bluegill	BSA	-	4.0 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Соре (1965)
Diuron-TCA (3 Ib/gal)	Lepomis macrochirus	BSA	-	5.7 (T4A)	_	Laboratory bioassays indicated that toxicity of the different formulations evaluated in this varied greatly with the fish used. Mortality data are expressed as EC10, EC50, etc.	Walker (1965)
Diuron (80 percent, WP)	lctalurus nebulosis Lepomis macrochirus	BSA	-	11.0 (T4A) 25.0 (T4A)	_	Comment same as above.	Walker (1965)
Diuron	Salmo gairdneri Lepomis macrochirus Daphnia pulex Simocephalus serrulatus	BSA	-	4.300 (T2A) 7.400 (T2A) 1.400 (T2A) 2.000 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Соре (1966)
Diuron	<i>Daphnia magna</i> Bluegill	BSA	-	47 (41.6- 53.1) (O) 7.4 (O)	acdiq	Toxicity, in terms of median immobilization concentration (IC <sub>50</sub> ), is presented for <i>Daphnia</i> ; median lethal concentra- tion for (LC <sub>50</sub> ) values for bluegill are reported.	Crosby and Tucker (1966)
Diuron	Simocephalus serrulatus Daphnia pulex	BSA	-	2.00 (SB) 1.40 (SB)	~	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during during the assay period.	Sanders and Cope (1966)
Diuron (Karmex)	Salmon Bluegill	BSA	-	16.0 (T2A) 74.0 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
Diuron (tech)	Pteronarcys callfornica (naiads)	BSA	-	0.0012 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)

Dowco 139 (25 percent EC)	Gambusia affinis	FL	Ponds— Bakers- field, Cal.	(0)	a C	At 0.1 lb/acre, 2 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulia and Isaak (1961)
Dowicide 31 (chloro-2- phenyl phenol, tech)	Lymnaeid snails	BSA	_	(0)	-	Each test container (500-ml beaker) was filled with ditch water. 100% mortality occurred at 1:200,000 and greater.	Batte, et al (1951)
Dowicide F (sodium 2,3,4,6- tetrachlorophe- nate, 80 percent)	Lymnaeid snails	BSA	-	(0)	-	Each test container (500-ml beaker) was filled with ditch water. 100% mortality occurred in concentrations of 1:600,000 and greater.	Batte, et al (1951)
Dowpon (2,2-dichloro- propionic acid)	Richardsonius balteatus hydroflox	BSA	-	444 (†1A) 412 (T2A) 395 (T4A)	ac de f	Results given were in soft water.	Webb (1961)
Dawpon	Typha latifolia Panicum hemitomum	FL	Fla.	(0)	-	At 51.0 and 17.0 lb/acre, 95 percent control of cattail was obtained and 3-5 percent control of <i>P. hemitomum.</i>	Copeland and Woods (1959)
Dowpon	Oncorhynchus kisutch Micropterus salmoides	BSA & CF	_	340 (T1) 340 (T2)	a c d e	Concentrations were based on percent active ingredient. In constant flow experiments, no bass survived 48 hours' exposure at 1000 ppm.	Bond, et al (1960)
2-(2,4DP) (dimethylamine)	Lepomis macrochirus	BSA	-	165 (T2A)	a e	The various salts of the chemicals showed wide variations in toxicity.	Hughes and Davis (1963)
2-(2,4-DP) (butoxyethanol ester)	Lepomis macrochirus	BSA	_	1.1 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
2-(2,4DP) (isooctyl ester)	Lepomís macrochirus	BSA	_	16 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
2,4-DP {ester}	Lepomis macrochirus Erimyzon sucetta L. macrochirus (fry)	L	-	10 (NTE) 1.5 (NTE) 20 (S)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)
DPT (DDT analogue)	Goldfish Gambusia affinis Culex apicalis (larvae)	BSA	-	10.0 (К) 2.0 (К) 0.1 (К)	-	Experiments were run a maximum of 3 days. No other time data were reported.	Odum and Sumerford (1946)
Drummer (pine)	Guppy	BSA	_	100 (K1)	<u>a</u>	Those fish that survived at lower concentrations were still very active several days after they had been taken out and placed in fresh water.	Anonymous (1964)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Dursban	Notemigonus crysoleucas Gambusia affinis Lepomis cyanellus	BSA	_	1.0 (T <1A) 1.0 (T <1A) 1.0 (T <1A)	e f	Fish used in the experiment were obtained from two sources, Mississippi Delta and State College pond. The fish obtained from the State College pond had not been exposed to insecti- cides. The fish obtained from the Mississippi Delta were known to be contaminated by cotton crop insecticides. Higher tolerance of Delta populations was evident in longer survival to 1.0 ppm Dursban.	Ferguson, et al (1966)
Dursban	<i>Prosimulum</i> spp <i>Cnephia</i> spp <i>Simulium</i> spp (larvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 32 percent.	Jamnback and Frempong- Boadu (1966)
Dursban	Leiostromus xanthurus Callinectes sapidus	BCFCH	_	0.001 (O)	а	At a concentration of .001 ppm, the following percent acetylcholinesterase activity as compared to controls was found: <i>L. xanthurus</i> – 38	Butler and Johnson (1967)
Dursban	Callinectes sapidus	BCFCH	-	0.010 (K)	а	Little or no information was given about test procedures and further tests.	Butler and Johnson (1967)
Dursban	Pteronarcys californica (naiads)	BSA	_	0.01 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Dursban	Pteronarcys californica (naiads)	BSA	-	0.010 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
	Pteronarcella badia (naiads) Claasenia sabulosa (naiads)			0.0038 (T4A) 0.0057 (T4A)			
DVP- iodine	Photococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum	BSA	_	100 (O)* 100 (O)* 50 (NG) 50 (NG)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but	Ukeles (1962)
	tricornutum Monochrysis lutheri			100 (K) *obvious, but inhibited growth.		the organisms were viable.	

Dylox	Gambusia affinis	BSA	_	0.5 (K 0%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)
Dylox	Salmo gairdnerii (one wk old sac fry) (one mo old feeding fry)	BSA	-	1.0 (K 0%) 10.0 (K 0%) 1.0 (K 0%) 10.0 (K 0%)	<u>a e</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
Dylox (Dipterex), (tech, 99 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	110 (T4A) 3.8 (T4A) 100 (T4A) 7.2 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)
Dylox (tech, 89 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica (naiad)	BSA	-	0.0690 (T4A) 0.0165 (T4A)	c d e f	A. pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than P. californica.	Jensen and Gaufin (1964)
Dylox	Gammarus Iacustris	BSA	_	0.050 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Dylox	<i>Cyprinodon variegatus</i> (juvenile)	BSA	-	(0)	а	Water temperature was 13 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Dylox, (tech)	Bluegill	BSA	-	0.260 (T4A)	а	This is an estimated LC $_{50}$ value at temperatures from 55 to 75 F.	Соре (1965)
Dylox, (tech)	Rainbow trout Bluegill	BSA	_	1.40 (T4A) 0.260 (T4A)	-	The values reported are given as LC <sub>50</sub> .	Cope (1965)
Dγlox	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.035 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
Dylox	Pteronarcys californica Acroneuria pacifica Ephemerella grandis Gammarus Iacustris Fathead	BSA	_	0.07 (T4A) 0.02 (T4A) 0.14 (T4A) 0.050 (T4A) 180.0 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were determined by standard methods. General comments were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional data are presented.	Gaufin, et al (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Dylox	Acroneuria pacifica	BSA	-	0.017 (T4A)	ac	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
	Ephemerella grandis			0.14 (T4A)			(,
	Gammarus Iacustris			0.05 (T4A)			
	Pteronarcys californica			0.07 (T4A)			
Dylox (99% active in water)	Hexagenia Hydropsyche (larva)	BSA	-	0.91 (T1A) 0.017 (T1A)	a e	Dissolved oxygen was measured before and after assay. Assays were conducted in Mississippi River water.	Carlson (1966)
,	Bluegill			12.0 (T1A)			
Dylox	Carassíus auratus Lebistes reticulatus	BCFA		0.4% (O)	<u>a</u>	The fish were fed on a diet to which the indicated amount of the chemical was added. During a 90-day period female goldfish showed no difference from the controls, while male fish showed a marked decrease in growth rate but this does not necessarily demonstrate that the chemical inhibited growth in the males. Reproduction was appar- ently not affected. Guppies under Dylox seemed to grow and reproduce normally.	Clemens, et (1966)
Dymid (WP)	Bluegill Rainbow trout	BSA?	_	75.0 (T4A) 97.0 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
EDB	Lepomis macrochirus Micropterus salmoides	BSA	_	18 (T2A) 15 (T2A)	aco	The response of bluegill and bass fingerlings to nine agri- cultural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
EN 18133 (50 percent EC)	Gambusia affinis	FL	Ponds — Bakers- field, Cal.	0.1 (K1) 0.4 (K1)	a c	Toxicity values indicate application rates in Ib/acre. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
Endosulfan EC2	Micropterus salmoides Cyprinus carpio	BSA	-	0.01 (O) 0.025 (K1) 0.005 (K 5 hr) 0.01 (K 3 hr) 0.02 (O)	a e	At 0.01 ppm, 50 percent mortality occurred in 1 day. For bass: Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds. At 0.02 ppm, 94 percent mortality occurred in 3 hr. For carp: Fish weights averaged 217 grams.	Mulla, et al (1967)
Endosulfan EC2	Micropterus salmoides Cyprinus	FL	Chino Fishery bass pond, Cal	0.05 (K1) 0.10 (K1) 0.05 (K1) 0.10 (K1)	а	The activity of Endosulfan when applied at 0.10 ppm lasted about 3 or 4 days. On the fifth day of treatment, only low mortality of the fish occurred in the treated pond.	
Endosulfan	carpio Pteronarcys californica (naiads)	BSA	Cal. —	0.00023 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)

	Endothal	Semotilus atromaculatus	BSA	-	1,600 to 3,200 (CR)	<u>a</u> e 	Test water used was freshly aerated Detroit River water. A typical water analysis is given. Toxicity is expressed as the "critical range" (CR), which was defined as that con- centration in ppm below which the 4 test fish lived for 24 hr and above which all test fish died. Additional data are presented.	Gillette, et al (1952)
	Endothal	Onchorynchus tshawytscha Micropterus salmoides	BSA, CF, and FL	-	155 (T1A) 136 (T2A) 200 (T1A) 200 (T2A)	a c d e	Concentrations were based on percent active ingredient. In 96 hours of exposure in the constant-flow apparatus, no largemouth bass mortalities were observed at 135 ppm, which was the highest concentration tested. In experimental field studies, Endothal controlled <i>Potamogeton pusillus</i> at about 0.3 ppm with no loss of largemouth bass or bluegills which were present in the pond.	Bond, et al (1960)
	Endothal (pellets)	Bushy pondweed Pondweed Coontail	FL	Lakes in Fla.	(O) (O) (O)	_	Concentrations of 0.5 to 2.0 ppm showed the best results in a variety of lakes, and in one lake 16.0 ppm was required to control bushy pondweed. 1.0 ppm controlled pondweed and coontail.	Phillippy (1961)
D 112	Endothal [3,6-endoxohexa- hydrophthalic acid (endotha), di-N,N-dimethyl- cocoamine salt]	Elodea canadensis Potamogeton nodosus Potamogeton pectinatus	BSA	-	5 (O) 100 (O) 5 (O) 100 (K 4 wk) 5 (O) 100 (K 4 wk)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. Injury rating of 8.5. Injury rating of 9.0. Injury rating of 9.1. Injury rating of 9.8.	Frank, et al (1961)
	Endothal and Silvex	Lepomis macrochirus	BSA	-	400 (T2A) L 600 (T2A) G	<u>a</u> cdeg	Toxicity data for 24 and 48 hours are presented for liquid (L) and granular (G) formulations. Various commercial formulations were tested. The liquid formulations were almost invariably more toxic than the granular ones.	Hughes and Davis (1965)
	Endothal	Lepomis macrochirus	BSA	-	280 (T2A) L 280 (T2A) G	<u>a</u> cdeg	Comment same as above.	Hughes and Davis (1965)
COMMERCIAL CHE	Endothal (liquid)	Micropterus salmoides (fry) Ictalurus punctatus (fry) Lepomis macrochirus (fry)	BSA	-	10 (SB3) 100 (SB3) 50 (SB3)	a c d e f p	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Endothal (granular)	Micropterus salmoides (fry) Ictalurus punctatus (fry)	BSA	-	2.0 (SB3) 50.0 (SB3)	a c d e f p	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
	Lepomis macrochirus (fry)			2.0 (SB3)			
Endothal	Lepomis macrochirus Pimephales promelas Micropterus salmoides	BSA	_	(H) 160 (T4A) (H) 610 (T4A) (S) 320 (T4A) (H) 200 (T4A)	<u>a</u> c <u>e</u>	Bioassay method in Standard Methods for examination of water was used. Both hard (H) and soft (S) water were used, TL <sub>m</sub> values for 24 and 48 hr are also presented.	Surber and Pickering (1962)
Endothal (dipotassium)	Lepomis macrochirus	BSA	_	428 (T1A)	-	The bioassay methods employed in this experiment were not given in the paper but it was stated that the same procedures were employed as in previous work.	Davis and Hughes (1963)
Endothall	Daphnia magna	BSA	-	46 (36-57) (O)	acdiq	Toxicity, in terms of median immobilization concentration ( $IC_{50}$ ), is presented.	Crosby and Tucker (1966)
Endothal	Salmon	BSA	-	136 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
Endothal	Lepomis macrochirus (eggs)	L	-	10 (NTE)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of	Hiltibran (1967)
	L. cyanellus (eggs)			10 (NTE)		days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the	
	Micropterus dolomieui (eggs)			10 (NTE)		highest concentration of chemical which did not cause death in 12 days (S).	
	Erimyzon sucetta (eggs)			10 (NTE)			
	L. macrochirus (fry)			50 (S)			

	Endrin	Carassius auratus Cyprinus carpio	FL	Japan	(O)	-	Four days after spraying at an application rate of 1 lb/acre, all fish placed in pond were dead after 8 hours of exposure. Endrin toxicity may persist in paddy fields as long as 1 month.	lyatomi, et al (1958)
	Endrin	Carassius auratus Channa argus (eggs & larvae) Cyprinus carpio (adult) (adult) (eggs & larvae) Moina	BSCH	_	0.003 (T1CH) 0.002 (T1CH) 0.0065-100 (T1CH) 0.003-0.42 (T1CH) 0.002-0.14 (T2CH) 0.046-19.9 (T1CH) 3.2 (T1CH)	<u>a</u>	Endrin became less toxic as temperature was lowered. Eggs and larvae of fishes were more resistant than adults, and the granular form of Endrin persisted longer than dust form.	lyatomi, et al (1958)
D 110	Endrin	macrocopa Fathead minnow Bluegill Goldfish Guppy	BSA	_	0.056 (T2CH) 0.001 (T4A) 0.00060 (T4A) 0.0019 (T4A) 0.0015 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds.	Tarzwell (1959)
	Endrin	Fathead minnow	BSA		0,0013 (T4A)	a	This experiment was performed in soft water. Comment same as above except that this experiment was performed in hard water.	Tarzwell (1959)
	Endrin	Daphnia magna	BSA	-	0.352 (O)	a	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
COMMERCIA	Endrin	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	0.001 (T4A) 0.0006 (T4A) 0.002 (T4A) 0.002 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
RCIA	Endrin (75%)	Pimephales promelas	BSA	~	0.0032 (T4A)	abecdf	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et al (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Røferøncø (Year)
Endrin (91%)	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	_	0.0014 (T4A) 0.00066 (T4A) 0.0021 (T4A) 0.0016 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et a (1959)
Endrin (19.5%)	Pimephales promelas Lepomis macrochirus	BSA	-	0.0038 (T4A) 0.0037 (T4A)	<u>a</u> becdf	Comment same as above.	Henderson, et al (1959)
Endrin	Pimephales promelas	BSA	_	0.0010 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)
Endrin	Live cars: Pimephales promelas Ictalurus melas Lepomis cyanellus Fish kill: Perca flavescens Lepomis gibbosus L. macrochirus Pomixis nigromaculatus Cyprinus carpio	FL	Pond	(O)	a c d	A beet field located adjacent to the study pond was treated with Endrin at the rate of 6 ounces of active ingredient per acre. Many fish were found dead after application. Live cars of the fish listed were placed in the pond to assess the residual toxic effects of Endrin. No mortality occurred in the live cars for 4 days, but the fish did accumulate Endrin in concentrations up to 1.0 ppm. The acute toxicity of Endrin appears to be less under these field conditions than in the laboratory.	Bridges (1961)
Endrin	Oncorhynchus kisutch Oncorhynchus	BSA	-	0.51 (T4A) 1.2 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
	tshawytscha Salmo gairdnerii Gasterosteus			0.58 (T4A) 0.44 (T4A)			

Endrin	Salmo	BCA		0.00 (744)		River water was diluont. The competization is given in onb	Katz
	gairdnerii	BJA	_		<u>a</u> cue		(1961)
	tshawytscha						
	kisutch						
	•			0.60 (T4A)			
	Gambusia			0.75 (T4A) 20 C			
	attinis			8.26 (14A) 3 C 0.33 (T4A) 25 C			
	Lebistes			0.90 (T4A)			
	Gasterosteus aculeatus			1.65 (T4A) Salin- ity 1.65 pp thousand.			
Endrin	Gammarus Iacustris Iacustris	BSA	_	(0)	аер	The mortality might have been partially due to the suscept- ibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Results were expressed as $LT_{50}$ ; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 27 (±1) min.	McDonald (1962)
Endrin	Pimephales notatus Lebistes reticulatus Adult đ Adult Q Adult Q Adult Q	BCFA	_	0.00047 (T4A) 0.0009 (T8A) 0.0009 (T10.6A) 0.00075 (T15A) 0.00075 (T15.5A)	<u>a c</u> d f	Chronic toxicity was also studied, as well as the effect of the toxicant on swimming and oxygen consumption. In the chronic study, bluntnose minnows survived for extended periods in 0.0001 ppm. Experiment on guppies was dis- continued because they succumbed to a kidney disorder.	Mount (1962)
Endrin	Lepomis	BSA	_	0.4 (T1A)	а	The experiment was conducted at 75 F. Fish weight was	Cope
(tech)	macrochirus			0.4 (T1A)		0.4 g. The experiment was conducted at 75 F. Fish weight was 0.6 g.	(1963)
Endrin EC 1.6	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a c	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals ap- plied as dilute sprays to ponds 1/16 acre in size. The indi- cated toxicant concentration is in Ib/acre, and resulted in a 100 percent mortality for the fish, and a 100 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
Endrin (tech)	Procambarus clarki	BSA	-	0.3 (T3A)	acdo	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
Endrin	Gambusia affinis affinis	BSA	_	0.001 to 0.12	а	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are later placed in clean water.	Boyd and Ferguson (1964)
	Endrin (tech) Endrin EC 1.6 Endrin (tech)	gairdneriiOncorhynchus tshawytschaOncorhynchus kisutch Lepomis macrochirus Gambusia affinisLebistes reticulatus Gasterosteus aculeatusEndrinGammarus lacustris lacustris lacustris lacustrisEndrinPimephales notatus Lebistes reticulatus Gambusia aculeatusEndrinPimephales notatus Lebistes reticulatus Adult of Adult Q Adult QEndrinLepomis macrochirusEndrinCammarus lacustris lacustrisEndrinPimephales notatus Lebistes reticulatus Adult of Adult QEndrinLepomis macrochirusEndrinCambusia affinis	gairdnerii       Oncorhynchus         tshawytscha       Oncorhynchus         tshawytscha       Oncorhynchus         kisutch       Lepomis         macrochirus       Gambusia         affinis       Lebistes         reticulatus       Gasterosteus         aculeatus       BSA         Endrin       Gammarus         Jacustris       BSA         Iacustris       BCFA         notatus       Lebistes         reticulatus       Adult of         Adult of       Adult of         Adult Q       Adult Q         Endrin       Lepomis       BSA	gairdnerii       Oncorhynchus         tshawytscha       Oncorhynchus         kisutch       Lepomis         macrochirus       Gambusia         affinis       Lebistes         Lebistes       reticulatus         Gasterosteus       aculeatus         Endrin       Gammarus       BSA         Indextris       lacustris         Lebistes       reticulatus         Gasterosteus       aculeatus         Endrin       Gammarus       BSA         Lebistes       reticulatus         Adult Q       Adult Q         Adult Q       Adult Q         Adult Q       BSA         Endrin       Lepomis         BSA       -	geirdherii       D. S. (T4A)         Oncorhynchus tshawytscha       0.92 (T4A)         Oncorhynchus kisutch       0.27 (T4A)         Oncorhynchus kisutch       0.60 (T4A)         Description       0.60 (T4A)         macrochirus       0.33 (T4A) 20 C         Gambusia       0.75 (T4A) 3 C         affinis       0.33 (T4A) 25 C         0.90 (T4A)       0.90 (T4A)         reticulatus       0.90 (T4A)         Gasterosteus       1.65 (T4A) Salin-         ity 1.65 pp       thousand.         Endrin       Gammarus         Lebistes       Primephales         reticulatus       BSA       -         Adult of       0.0009 (T8A)         Adult of       0.0009 (T10.6A)         Adult of       0.00075 (T15A)         Adult of       0.00075 (T15A)         Adult of       0.00075 (T15A)         Adult of       0.44 (T1A)         Endrin       Gembusia       FL       Cal.       0.5 (O)	geirdnerii     Oncorhynchus     0.92 (T4A)       tshewytscha     0.92 (T4A)       Oncorhynchus     0.27 (T4A)       kisutch     0.60 (T4A)       Lepomis     0.60 (T4A)       macrochirus     0.33 (T4A) 25 C       Jacustris     0.33 (T4A) 25 C       Jubistes     0.33 (T4A) 25 C       Jubistes     0.90 (T4A)       reticulatus     1.65 (T4A) 3alin-       aculeatus     1.65 (T4A) Salin-       ity 1.65 pp     thousand.       Endrin     Gammarus       BSA     -       Icoustris     0.000047 (T4A)       aculeatus     1.65 (T4A) Salin-       ity 1.65 pp     thousand.       Endrin     Gammarus       BSA     -       Icoustris     0.00009 (T8A)       Adult 3     0.0009 (T10.6A)       Adult 4     0.000075 (T15.A)       Adult 9     0.00075 (T15.A)       Adult 9     0.4 (T1A)       Endrin     Lepomis       BSA     -       0.4 (T1A)     a	sircherit     0.92 (TAA)       Oncorthynchus     0.92 (TAA)       Oncorthynchus     0.27 (TAA)       Kautch     0.80 (TAA)       Oncorthynchus     0.27 (TAA)       Kautch     0.80 (TAA)       macrochirus     0.75 (TAA) 20 C       Safinis     0.33 (TAA) 25 C       Lebistes     0.90 (TAA)       Gambusia     0.75 (TAA) 26 C       Safinis     0.33 (TAA) 25 C       Lebistes     0.90 (TAA)       Gasteroateus     1.65 (TAA) 56 C       seculaetus     1.65 (TAA) 56 C       Lebistes     0.90 (TAA)       freiculatus     Gasteroateus       Gasteroateus     1.65 (TAA) 56 C       seculaetus     1.65 (TAA) 56 C       Jacutris     BSA       Jacutris     BSA       Jacutris     BSA       Jacutris     BSA       Jacutris     BCFA       Jacutris     BCFA       Jacutris     BCFA       Jacutris     Chronic toxicity was also studied, as well as the effect of the chronic taxicy was also studied, as well as the effect of the chronic taxicy was also studied, as well as the effect of the chronic taxicy was also studied, as well as the effect of the chronic taxicy was also studied, in 0.0000 (TISA)       Adult Q     0.000075 (TISA)       Adult Q     0.000075 (TISA)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Endrin	Notemigonus crysoleucas Lepomis macrochirus L. cyanellus	BSA	-	(B) 0.003 (T1.5) (A) 0.310 (T1.5) (B) 0.0015 (T1.5) (A) 0.300 (T1.5) (B) 0.0034 (T1.5) (A) 0.160 (T1.5)	acf	Chemical was dissolved in acetone. Final concentration of acetone was $\leq 2 \text{ ml/l}$ . Data shows $\text{TL}_{m}$ in ppb for insecticide-resistant and insecticide non-resistant strains of the test fish.	Ferguson, et al (1964)
Endrin (technical, 100 percent active in acetone)	Pteronarcys californica (naìad) Acroneuria pacifica (naìad)	BSA	-	0.00240 (T4A) 0.00039 (T4A)	c d e f	A. pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than P. californica.	Jensen and Gaufin (1964)
Endrin	Gammarus Iacustris	BSA	_	0.0115 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Endrin	Bluegill	BSA	-	0.0006 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Endrin	Bluegilł	BSA	-	0.7 (T4A) 0.7 (T4A) 0.4 (T4A) 0.4 (T4A) 0.2 (T4A)	<u>a</u>	These experiments were performed to demonstrate that at increased temperatures the toxic effect of most chemicals is increased. For the toxicant concentrations listed, the temperatures were respectively, 45, 55, 65, 75, and 85 F. Data on the effect of time as well as temperature was also reported. The experimental animals all were approximately one grain in weight.	Соре (1965)
Endrin (tech)	Rainbow trout Bluegill	BSA	-	0.00086 (T4A) 0.00025 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
Endrin	Rainbow trout	BSA	-	2.4 (T4A) 1.4 (T4A) 1.1 (T4A) 0.75 (T4A)	<u>a</u>	These experiments were performed to show the effect of temperature on the toxicity. For the toxicant concentrations listed, the temperatures were respectively, 45, 55, and 65 F. The fish all were approximately one grain in weight. Toxicant concentrations for one and 2-day times were also listed.	Cope (1965)
Endrin	<i>Leiostomus xanthurus</i> (juvenile)	ВСН	_	(O)	-	Endrin at 0.0006 ppm killed half the shrimp exposed in 24 hours. The fish survived a concentration of 0.00005 ppm for 8 months, but a concentration of 0.0001 was usually lethal in 5 days.	Butler (1965)
Endrin	Fundulus similis (juvenile)	BSA	_	0.000079 (O)	а	Water temperature was 21 C. The figure reported is a 24-hr $EC_{50}$ .	Butler (1965)

	Endrin	Dorosoma cepedianum	BSA	_	(0)	<u>a</u> de	The critical level of Endrin in the blood (0.10 $\mu$ g/g) of gizzard shad as a result of laboratory exposures was also applicable for shad exposed to Endrin in a natural system.	Brungs and Mount (1967)
	Endrin	Palaemonetes kadiakensis	BSA	-	(N) 6.5 (T1½A) (TB) 9.5 (T1½A)	a c f	Test organisms were collected from 2 locations, Twin Bayou (TB), Sunflower Co., Miss. (agricultural area) and Noxubee National Wildlife Refuge (N), Noxubee Co., Miss. (non- agricultural area) and evaluated in laboratory bioassays. The Twin Bayou shrimp were more resistant.	Ferguson, et al (1965)
	Endrin	Gambusia affinis Ictalurus melas	BSA	_	0.0005- 0.002 (T3A) 0.0004- 0.002 (T3A)	<u>a</u> cde	Test fish were collected from 8 different locations of the Mississippi River. The 3-day $TL_m$ values were made to determine if a resistance gradient existed. The data indicated that there was none.	Ferguson, et al (1966)
	Endrin	Lepomis cyanellus Gambusia affinis Ictalurus melas	BSA	-	0.001-0.048 (O)	-	Muds reduced the toxicity of chlorinated hydrocarbon insecticides to fish. Lethal quantities of pesticides enter national waters and muds may contain sorbed pesticides in excess of lethal quantities. Although the chemicals can be leached with organic solvents, they were either not re- leased in lethal amounts or slowly released in standing water.	Ferguson, et al (1965)
<b>D</b> 110	Endrin	Acroneuria pacífica Ephemerella grandis Gammarus lacustris Pteronarcys californica	BSA	-	0.0003 (T4A) 0.005 (T4A) 0.0115 (T4A) 0.0024 (T4A)	<u>a</u> c	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
	Endrin	Pteronarcys californica Acroneuria pacifica Ephemerella grandis Daphnia magna Gammarus lacustris	BSA	-	0.002 (T4A) 0.0004 (T4A) 0.005 (T4A) 0.4 (T 50 hr A) 0.01 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were de- termined by standard methods. General comments were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional data are presented.	Gaufin, et al (1965)
COMMERCIAL	Endrin	Gambusia affinis	BSA	_	(O)	a	The effect of combinations of pesticides was studied. In general, the results reflected the extreme levels of Endrin and Toxaphene resistance in the resistant population. The results failed to indicate additive effects wherein the combination mortality exceeded the sum of the mortalities produced by the individual insecticides.	Ferguson and Bingham (1966)
COMMERCIAL CHEMICAL PRODUCTS	Endrin	Leiostomus xanthurus Mugil cephalus Brevoortia patronus Fundulus similis Cyprinodon variegatus	BCFA	_	0.00045 (T1A) 0.0026 (T1A) 0.00080 (T1A) 0.00023 (T1A) 0.00032 (T1A)	ai	The duration of exposure was important when determining the sublethal concentrations of Endrin to fish. Data are given as LC50.	Lowe (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted(4)	Comments	Reference (Year)	_
Chemical Endrin Endrin Endrin	Fathead minnow	BCFA	_	0.47 ppb (T4A) 0.50 ppb (T4A)* 0.66 ppb (T4A)** * clay **charcoal	<u>a</u> cde	The effect of suspended particles on Endrin toxicity was studied. Presence of clay particles had no effect, while activated charcoal reduced toxicity.	Brungs and Bailey (1966)	
Endrin	Oyster	BCF	-	0.01-1.0 (SB4)	а	Seawater was employed in this experiment.	Butler (1966)	
Endrin	Salmo gairdneri Lepomis macrochirus Ictalurus punctatus Pteronarcys californicus Baetis sp Daphnia pulex Simocephalus serrulatus	BSA	-	0.0005 (T2A) 0.0003 (T2A) 0.001 (T2A) 0.001 (T2A) 0.005 (T2A) 0.020 (T2A) 0.026 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects of repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Cope (1966)	F
Endrın	lctalurus natilis I. melas Gambusia affinis Lepomis cyanellus Notemigonus chrysoleneas	FR	Miss.	(O)	а	This paper deals with Endrin resistance in fish. The $TL_m$ 36 hours values for yellow bullheads from a contaminated area was 75 ppb, while the value for fish from an unsprayed area was only 1.25 ppb. Mosquito fish from a contaminated area tolerated 1500 ppb of Endrin, while golden shiners and green sunfish tolerated 1000 to 250 ppb, respectively. Bullheads were apparently more susceptable to the poison, but there seemed to be evidence that fish can develop a tolerance to the toxicant.	Ferguson and Bingham (1966)	APPENDIX B
Endrin	Acroneuria pacifica Pteronarcys californica	BSA & CFCH	-	0.00032 (T4A) 0.000035 (T30CH) 0.0024 (T4A) 0.0012 (T30CH)	<u>a c d e</u>	Additional data are presented.	Jensen and Gaufin (1966)	
Endrin	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto alba Strix aluco	FO	Netherlands	(O)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecti- cides. In most cases the liver had the highest concentration of toxicant, ranging from 0.3 ppm. Most chlorinated hydro- carbons tended to accumulate in the fat depots of the body. In instances where mesenterial fat was analyzed the con- centration of toxicant was found to be as high as 15.7 ppm.	Koeman and van Genderen (1966)	

	Osio otus Falco pereginus Platelea leucorodia Haematopus ostralegus Steran sandvicensis Sterna hirundo Larus ridibundus Somateria mollissima Tadorna tadorna		Netherlands	(0)	-	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecti- cides. In most cases the liver had the highest concentration of toxicant, ranging from a trace to 3.0 ppm. Birds feeding predominantly on crustacea, molluses, and fish contained significant amounts.	Koeman and van Genderen (1966)
Endrin	lctalurus punctatus	BCFCHA	-	(O)	<u>a</u>	Catfish blood content acutely toxic in 10 days or less- 0.25-0.3 ppb, nonlethal (exposure during 44 days)- 0.1-0.2 ppb. Endrin was not stored in blood and 0.3 ppb appeared to be the critical concentration level in the blood.	Mount, et al (1966)
Endrin	Daphnia carinata	BSA	_	0.050 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Simocephalus serrulatus Daphnia pulex	BSA	-	0.026 (SB) 0.020 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Daphnia magna	BSA	_	0.900 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Simocephalus serrulatus Daphnia pulex	BSA	-	4.90 (SB) 3.20 (SB)		Comment same as above.	Sanders and Cope (1966)
Endrin	Catfish Buffalo fish Perch Bluegill Carp	L	-	(O)	_	Chemical analysis showed residues of $<0.01$ to 0.04 ppm in catfish, $<0.01$ in buffalo fish, and 0.05 ppm in carp. Perch were reported to contain 0.02 and bluegills 0.01 ppm, but this may be doubtful inasmuch as the method of anaylsis uses was thought to have an interference from toxaphene which had been used previously. The treated area had been sprayed with Endrin at 0.3 lb/acre.	Sparr, et al (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Endrin	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	_	(O)	_	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentration of 500-1250X resulted.	Butler (1967)
Endrin	Fundulus similis Leiostomus xanthurus	BCFCH	-	0.0001 (SB 6 mo) 0.1 (SB 3 mo)	-	When these fish were examined for residues at the end of 6 months, 92.0 ppb Endrin residue was found.	Butler and Johnson (1967)
Endrin	Oyster	FE	Galveston Bay, Texas	(0)	_	Oysters from the area were found to contain $<$ 0.01 to 0.02 ppm.	Casper (1967)
Endrin	Puntius puckelli	BSA	_	0.00125 (T4A)	<u>a</u> c d e l m	Tap water was used as diluent. Toxicity data are given as TLm's in ppm for 24, 48, 96 hr. The pH of the water averaged 8.3. The study was conducted in India.	Rao, et al (1967)
Endrin	lctalurus melas	BSA	_	50 ppb (T1/2A)	а	The principal mode of Endrin entry in the body of the fish is by way of the gill surfaces. The toxicity figure cited is for 10 hr.	Velsen and Alderdice (1967)
Endrin	Vascular plants Algae Chub Largemouth bass Clam	FL	Tule Lake, Ore.	(0)	_	Endrin was applied at the rate of 1.6 lb/acre/year. Plants contained 1.6 to 12.5 ppb. Algae contained 2.0 to 22.3 ppb. Chubs contained 4.0 to 198.0 ppb. Bass contained 2.0 to 107.0 ppb. Clams contained 1.7 to 90.0 ppb. Concentrations of 0.007 to 0.01 ppb occurred in the water.	Godsil and Johnson (1968)
Endrin	Noternigonus crysoleucas	BSA	-	(O)	_	This paper deals with the resistance and susceptibility of populations of golden shiners to Endrin. Two populations of golden shiners from agricultural areas possessed different levels of resistance to Endrin. At 1.0 ppm, Endrin killed 50 susceptible golden shiners in 75 min, but only 40 of 50 resistant shiners in 40 hours. Endrin residues in whole bodies of resistant shiners killed by Endrin were as much as 82 times those of the susceptible shiners. It was concluded that the use of a critical concentration in the blood for diagnosis of Endrin-caused mortality must be based on the tolerance of local populations.	Ludke, et al (1968)
Endrin	Limnephilus rhombicus Sialis sp Gammarus sp		Knights Creek, Wisc.	(0)	-	Pesticide usage in an orchard did not significantly contaminate the aquatic environment of this creek adjacent to the treat- ment as determined by residue analysis.	Mowbry,et (1968)

Endrin	Esox americanus Micropterus salmoides Lepomis macrochirus Rana catesbeiana Pseudemys scripta elegans Natrix erythrogaster flavigaster Natrix rhombifera Ancistrodon piscivorus	BSA	_	(K) 7.1 hr (O) (K) 12.6 hr (K) 9.4 hr (K) 15.6 hr (K) 65.4 hr (K) 54.0 hr (O)	Not given	Mosquitofish ( <i>Gambusia affinis</i> ) were exposed to 2 ppm Endrin solutions for 7 days. The fish were somewhat resistant to Endrin. These fish were then force-fed to the 8 species of vertebrates listed. Survival time is listed. Mortality was 100 percent in this time period except for <i>P. scripta elegans</i> and <i>A. piscivorus</i> . For the <i>P. scripta elegans</i> mortality was 72 percent in 112.8 hr and for the <i>A. piscivorus</i> the mortality was 91% in 27.1 hr.	Rosato and Ferguson (1968)
Endrin	Pteronarcys californica (naiads) Pteronarcella badia (naiads) Claasenia sabulosa (naiads)	BSA	-	0.00025 (T4A) 0.00054 (T4A) 0.00076 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
EPN miticide (ethyl-p-nitro- phenyl thio- benzenephos- phonate, 31.5 percent)	Lymnaeid snails	BSA		(O)	-	Each test container (500-ml beaker) was filled with ditch water. Less than 100% mortality occurred in concentra- tions of 1:100,000.	Batte, et al (1951)
EPN-300 (25%)	Pimephales promelas	BSA	-	0.80 (T4A)	<u>acdef</u>	Tests were performed in both hard and soft water. Addi- tional tolerance limit values are given.	Henderson and Pickering (1958)
EPN	Fathead minnow	BSA	-	0.25 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959)
EPN	Daphnia magna	BSA	-	0.0001 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
EPN	Pimephales promelas	BSA	_	0.2 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
EPN	Pimephales promelas	BSA	-	0.20 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)
EPN	Lepomis macrochirus	BSA	-	0.1 (O)	<u>acdf</u>	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The	Weiss (1959)
	Microp terus salmoides			0.1 (O)		inhibition of the enzyme was a function of the concentra- tion of the insecticide, extent of exposure, and specific	
	Notemigonus			0.1 (O)		chemical nature of the compound.	
	crysoleucas Carassius auratus			0.1 (O)			
PN	Micropterus salmoides	BSA	-	0.5 (O)	-	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentra-	Weiss (1961)
	Pimephales promelas			0.5 (O)		tion, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the $LT_{50}$ was 9 hr 30 min and for fatheads 72 hr.	(1001)
EPN-300, wettable	Pimephales promelas	BSA	-	1.1 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent	Pickering, et al (1962)
powder (25 percent)	Lepomis macrochirus			0.44 (T4A)		or carrier in most cases.	
(25 percent)	Carassius auratus			2.3 (T4A)			
EPN (tech,	Pimephales promelas	BSA	-	0.25 (T4A)	<u>a</u> cd <u>e</u>	Comment same as above.	Pickering, et al (1962)
100 percent)	Lepomis			0.10 (T4A)			(1902)
	macrochirus Carassius			0.45 (T4A)			
	auratus Lebistes reticulatus			0.032 (T4A)			
EPN	Chaoborus astictopus	BSA	-	0.0036 (T1A)	<u>a</u>	Toxicity value given is for the fourth instar larvae.	Hazeltine (1963)
EPN	Bluegill	BSA	-	0.1 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)

EPN	Carassius auratus Lepomis macrochirus Notemigonus crysoleucas	BSCH	-	10.0 (0)* 1.0 (0)** 0.05 (0)*** 1.0 (0)* 0.05 (0)*** 1.0 (0)* 1.0 (0)* * response, 15 days ** no re- sponse, 15 days ***no re- sponse, 30 days	<u>a</u> c d <u>e</u>	Toxicity was determined by measuring acetylcholin- esterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
Eptam	Crassostrea virginica Penaeus setiferus Mugil cephalus Phytoplankton	BCFA & BSA		5.0 (0.43%) 0.63 (O) 20.0 (10% T2A)	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Erythromycin thiocyanate	Salmo gairdnerii	BSA	_	(O)	aceip	A dosage of 500 milligrams of erythromycin thiocyanate per kilogram per day (five times the usual therapeutic level) was required to produce overt symptoms of toxicity in rainbow trout.	Warren (1963)
Erythromycin thiocyanate	Salmo gairdnerii Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	_	100 (NTE)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
Essolvene	Pandalus montagni Crangon crangon Carcinus maenas Cardium edule	BSA	-	8.6 (T2A) 9.6 (T2A) 17.5 (T2A) 63.0 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C. It was shown that the toxicity of this solvent emulsifier decreased with time due to evaporation of the solvent. Essolvene at a concentration of 10 ppm killed 100% of <i>Crangon crangon</i> larvae in 3 hr; at 33.3 ppm it killed 100% of <i>Carcinus mænas</i> in 3 hr.	Portmann and Connor (1968)

ę

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Estron (2,4-5T)	Pontederia cordata	FL	Fla.	(0)	_	At 6.4 lb/acre, 80 percent control of pickerel weed was obtained.	Copeland and Woods (1959)
Esteron 99	Lepomis macrochirus	BSA	-	1,200 (T 18 hr)	а	The experiment was conducted at 65 F Fish were 2 in. long.	Cope (1963)
Esteron 99	<i>Lepomis</i> sunfish	FL	Okla.	(O)	-	Three ponds were partitioned with polyvinyl chloride sheeting to provide 6 test spaces for fish. Mortality of the fish was 19% in the 10-ppm pond in the first week.	Cope (1963)
Esteron 99 (EC)	Lepomis macrochirus	BSA	_	700 (T1A)	а	The experiment was conducted at 75 F. Fish weighed 0.6 g.	Cope (1963)
Esteron 99 (2, 4-D)	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	0.055 (O) 0.55 (O) 1.5 (T2SA)	_	Seawater was pumped continuously into test aquaria. Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following: Oyster - 96-hr EC <sub>50</sub> - Conc. which decreased shell growth. Shrimp - 48-hr EC <sub>50</sub> - Conc. which killed or paralyzed 50% of test animals. Fish - 48-hr EC <sub>50</sub> - Conc. which killed 50%. Phytoplankton - Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical	Butler (1965)
Esteron 99	Pteronarcys sp	BSA	-	1.6 (T4A)	a	Experiments were all conducted at 60 F in 1964. The values were listed as LC50.	Соре (1965)
Ethion	(ny mphs) <i>Cy prinodon variegatus</i> (juvenile)	BSA	-	0.064 (O)	а	Water temperature was 12 C. The figure reported is a 48-hr $EC_{50}$ .	Butler (1965)
Ethion (tech)	Pimephales promelas Lepomis macrochirus Lebistes reticulatus	BSA	-	2.4 (T4A) 0.13 (T4A) 0.13 (T4A)	a c d e f	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson (1966)
Ethion	Pteronarcys californica (naiads)	BSA	-	0.0028 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Ethyl carbo- phenothion	Chaoborus astictopus	BSA	-	0 <b>.044 (</b> T1A)	<u>a</u>	Toxicity value given is for the fourth instar larvae.	Hazeltine (1963)
Ethyl guthion (EC2)	Gambusia affinis Rana catesbeiana	FL	Ponds in III.	(0)	-	When applied at 0.01 and 0.025 pound per acre active ingredient, 88 and 100 percent fish mortality occurred respectively in 1 day. No bullfrog mortality occurred at 0.8 pound per acre in 1 day.	Mulia, et al (1963)

	Ethyl parathion	Chaoborus astictopus Lepomis macrochirus	BSA	-	(O) 0.021 (T9A)	<u>a</u>	Tests were run on bluegill sunfish, <i>C. astictopus</i> first instar larvae, and fourth instar larvae, results for larvae were as follows: Fourth instar 0.017 (T1A) First instar 0.0018 (T1A)	Hazeltine (1963)
	Ethyl parathion	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.0051 (T4A)	a	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> ,	Cope (1965)
	Ethyl parathion	Simocephalus serrulatus Daphnia pulex Daphnia carinata Daphnia magna	BSA	-	0,00037 (SB) 0.00060 (SB) 0.0005 (SB) 0.0008 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Ethyl guthion	<i>Pteronarcys</i> sp (nymphs)	BSA		0.002 (T4A)	a	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
	Ethyl guthion, (tech)	Rainbow trout	BSA	-	0.019 (T4A)	-	The values reported are given as LC <sub>50</sub> .	Cope (1965)
	Ethyl guthion	Salmo gairdneri Lepomis macrochirus Pteronarcys californica Daphnia pulex Simocephalus serrulatus	BSA	-	0.023 (T2A) 0.002 (T2A) 0.008 (T2A) 0.003 (T2A) 0.004 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Cope (1966)
	Ethyl guthion	Simocephalus serrulatus Daphnia pulex	BSA	_	0.0042 (SB) 0.0032 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
COMMERCIAL CHEMI	Ethyl guthion EC4	Micropterus salmoides Cyprinus carpio	BSA	-	0.05 (O) 0.10 (O) 0.50 (K1) 0.01 (K1) 0.05 (K 5 hr) 0.10 (K 5 hr)	a e	<ul> <li>At 0.05 ppm, 25 percent mortality occurred in 1 day.</li> <li>At 0.10 ppm, 90 percent mortality occurred in 1 day.</li> <li>For bass:</li> <li>Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.</li> <li>For carp:</li> <li>Experiments were carried out in plastic tubs lined with saran plastic. Fish weights averaged 217 grams.</li> </ul>	Mulla, et al (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Exalgae	Chlorella pyrenoidsa	L	-	5 (AC 1/2 hr)	-	Describes a bioassay method to differentiate between an algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)
F-98	Onchory nchus tshawy tscha	BSA	-	0.08 (T1A)	a c d e	Concentrations were based on percent active ingredient.	Bond, et al (1960)
Fairfield 279	Salmo gairdneri	BSA	_	360 (T1A)	а	The experiment was conducted at 55 F Fish weighed 0.5 g.	Cope (1963)
Fairfield OT 60-6	Salmo gairdneri	BSA	_	100 (T1A)	а	The experiments were conducted at 55 F. Fish weighed 0.8 g.	Cope (1963)
Fenac (sodium salt)	Redear sunfish Salmo gairdneri	BSA	-	(O) 10,000 (T1A) 7,500 (T2A)	а	The experiment was conducted at 75 F. Fish weighed 3 g. No mortality was noted with concentrations of 12,000 mg/1 at 48 hr. The experiments were conducted at 65 F Fish weighed 0.6 g.	Соре (1963)
Fenac	Lepomis macrochirus	BSA	_	22.5 (T2A) L 15.0 (T2A) G	<u>a</u> cdeg	Toxicity data for 24 and 48 hours are presented for liquid (L) and granular (G) formulations. Various commercial formulations were tested. The liquid formulations were almost invariably more toxic than the granular ones.	Hughes and Davis (1965)
Fenac (sodium salt)	<i>Pteronarcys</i> sp (ny mphs)	BSA	_	47 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
Fenac acid (tech)	Bluegill	BSA	_	41 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Соре (1965)
Fenac (sodium salt, WP)	Bluegill	BSA	-	14 (T4A)	а	Comment same as above.	Cope (1965)
Fenac (acid)	<i>Pteronarcys</i> sp (nymphs)	BSA	-	56 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
Fenac, Na	Salmo gairdneri	BSA	-	7.500 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses subacute mortality as well. Effects on repro-	Cope (1966)
	Lepomis macrochirus			19.000 (T2A)		duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	
	Pteronarcys californica			80.000 (T2A)			
	Daphnia pulex Simocephalus			4.500 (T2A) 6,600 (T2A)			
Fenac	serrulatus Daphnia	BSA		100 (O)	acdiq	Toxicity, in terms of median immobilization concentration	Crosby and
(Na salt)	<i>magna</i> Rainbow			7.5 (O)		(IC <sub>50</sub> ), is presented for <i>Daphnia</i> ; median lethal concentra- tion (LC <sub>50</sub> ) for rainbow trout and bluegill are reported.	Tucker (1966)
	trout Bluegill			19 (O)			

Fenac (sodium	Simocephalus serrulatus	BSA	_	6.6 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr.	Sanders and Cope
salt)	Daphnia pulex		-	4.5 (SB)		Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was ''controlled'' during the assay period.	(1966)
Fenac (sodium salt, WP)	Rainbow trout	BSA	-	7.5 (T2A)	_	Data are given as LC <sub>50</sub> .	Bohmont (1967)
	Bluegill			19.0 (T2A)			
Fenac	Lepomis macrochirus (eggs)	L	_	20/5 (O)	_	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of	Hiltibran (1967)
	Erimyzon sucetta (eggs)			20 (NTE)		days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause	
	<i>L. macrochirus</i> (fry)			50 (S)		death in 12 days (S).	
Fenac	Pteronarcys californica (naiads)	BSA	-	0.06 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
Fenac (Na salt)	Pteronarcys californica (naiads)	BSA	-	0.055 (T4A)	<u>acdef</u>	Comment same as above.	Sanders and Cope (1968)
Fenthion	Prosimulum spp Cnephia spp Simulium spp (Iarvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 89 percent.	Jamnback and Frempong- Boadu (1966)
Fenthion (Baytex)	Micropterus salmoides	BSA	—	1.75 (L1A)	a	Abate was toxic to fish at a dosage rate necessary to con- trol the larvae of the chironomid midge.	Von Windeguth and Patterson
(	Lepomis macrochirus			1.75 (L1A)		-	(1966)
	Gambusia affinis			2.0 (L1A)			
	Lebistes reticulatus			1.75 (L1A)			
	Palomonetes paludosus			0.011 (L1A)			
	Hyalella azteca			0.016 (L1A)			
	Plankton (Euglena, Coleops)			1.0 (K2)			
	Rotifers			1.0 (K2)			
Fenthion	Micropterus salmoides	BSA		5.0 (K 3 hr)	a e	Experiments were carried out in plastic tubs lined with saran plastic. Fish weights averaged 217 grams.	Mulia, et al (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Fenthion	Pteronarcys californica (naiads)	BSA	_	0.0045 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
~enuron	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodectylum tricornutum Monochrysis lutheri	BSA	-	29.0 (K) 2.9 (NG) 2.9 (NG) 2.9 (NG) 2.9 (K)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organ- isms were viable.	Ukeles (1962)
enuron	<i>Leiostomus</i> <i>xanthurus</i> (juvenile)	BSA		(O)	a	Water temperature was 25 C. Fish showed irritation at 1.0 ppm.	Butler (1967)
enuron	Penaeus aztecus	L	-	(0)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. A concentration of 1.0 ppm caused 10 percent mortality.	Butler (1965)
-enuron	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	2.0 (NTE) 2.0 (0, 10%) 1.0 (NTE) 41% (0)	_	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Butler (1965)
enuron (25 percent pellet)	Lepomis macrochırus	BSA	-	53.0 (T4A)	_	Laboratory bioassays indicated that toxicity of the different formulations evaluated in this varied greatly with the fish used. Mortality data are expressed as EC <sub>10</sub> , EC <sub>50</sub> , etc.	Walker (1965)
enuron-TCA (tech)	Lepomis macrochirus	BSA	-	5.3 (T4A)	_	Comment same as above,	Walker (1965)
Fenuron-TCA (3 Ib/gal)	Lepomis macrochirus Micropterus salmoides	BSA	-	4.8-6.5 (T4A) 7.4 (T4A)	-	Comment same as above.	Walker (1965)

Fenuron TCA	Lepomis macrochirus L. cyanellus Micropterus dolomieui Erimyzon sucetta L. macrochirus (fry)	L	-	10 (NTE) 10 (NTE) 10 (NTE) 10 (NTE) 20 (S)	_	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)
Fermate	Channel catfish (fingerlings)	BSA	-	12.6 (K 27 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Fermate	Salvelinus fontinalis x Salmo trutta Catostomus commersoni Micropterus salmoides	FPCH	N.Y.	0.5 (S23) 0.5 (S23) 0.5 (S23)	a c d	Conventional farm ponds were used having an average surface area of 0.3 acre and a maximum depth of 7-9 ft. Toxicity (in ppm) to fish as maximum safe concentration (S) for 23 days was determined. Con- centration of 0.5 ppm was required to control algae.	Eipper (1959)
Fermate	Pimephales promelas Lepomis	BSA	-	3.1 (T4A)	a c d e f	The toxicity of this substance was influenced by the quality of the water (pH, hardness, alkalinity). The $TL_m$ was lower in hard water.	Pickering and Henderson (1966)
Folidol	Tilapia massambica Gambusia affinis	BSA	_	0.6 (T2A) 0.1 (T2A)	acdefp	The test animals were conditioned for 48 hours prior to use.	Sreenivasan and Swaminathan (1967)
Folpet [N-(trichloro- methylthio)- phythalimide]	Brachydanio rerio	BSA	-	1 (O)	а	At 1 ppm all larvae were killed within 48 min. The $\rm TL_{50}$ was 34 min and $\rm LD_{50}$ was 0.71 ppm.	Abedi and Turton (1968)
Folithion or Sumithion (=fenitrothion)	Prosimulum spp Cnephia spp Simulium spp (Iarvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 65 percent.	Jamnback and Frempong- Boadu (1966)
Furron	Water lettuce	FL	Lakes in Fla.	(O)	_	Application rates of 4 to 5 lb/acre controlled water lettuce.	Phillippy (1961)
Furazolidone	Salmo trutta Salmo gairdnerii Salvelinus fontinalis	ВСН	-	(O)	a c	The chemical was nontoxic to brown, rainbow, and brook trout at levels up to 500 mg per kg of body weight per day when force-fed for 14 consecutive days. Therapeutic levels for control of furnuculosis appear to be as low as or lower than 10 milligrams of chemical per kilogram of body weight per day for 14 days. To have complete con- trol of the disease, a dosage of at least 75 milligrams of furazolidone per kilogram of body weight was given.	Post and Keiss (1962)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chemical Furoxone G-27365 (EC2)	Salvelinus fontinalis	BSA	_	(0)	<u>a e</u>	Aeromonas salmonicida is the bacterium causing furunculosis in fish. The toxicity level of this chemical indicates that this drug could be used as a therapeutic measure for A. salmonicide. 500 milligrams per kilogram per day used for 15 successive days showed no pathological effect on brook trout.	Post (1959)
G-27365 (EC2)	Gambusia affinis Rana	۴L	Ponds in III.	(0)	-	When applied at 0.2 and 0.4 pound per acre active ingredient, 8 and 100 percent fish mortality occurred respectively in 1 day. No bullfrog mortality occurred at 0.8 pound per acre in	Mulla, et al (1963) Mulla, et al
	catesbeiana					1 day.	(1963)
G- <b>28029</b> (EC2)	Gambusia affinis Rana castesbeiana	FL	Ponds in III.	(0)	_	When applied at 1.6 pounds per acre active ingredient, 6 per- cent fish mortality occurred in 1 day. No bullfrog mortality occurred at 1.6 pounds per acre in 1 day.	Mulla, et al (1963)
G-30493 (EC2)	Gambusia affinis Bufo boreas	FL	Ponds in III.	(O) (O)	-	When applied at 0.8 pound per acre active ingredient, 2 per- cent fish mortality occurred in 2 days. No toad mortality occurred at 0.8 pound per acre in 1 day.	Mulla, et al (1963)
	Scophiopus hammondi			(O)			
G-30494 (EC2)	Gambusia affinis Bufo	FL	Ponds in III.	(O) (O)	-	When applied at 0.8 pound per acre active ingredient, 100 percent fish mortality occurred in 1 day. When applied at 0.4 pound per acre, 5 percent toad mortal-	Mulla, et al (1963)
	boreas Scophiopus hammondi			(0)		ity occurred in 1 day.	
GC-405 (zinc nicotinyl fluosilicate)	Australorbis glabratus	BSA and FL	Puerto Rico	Failed	С	Seven of the tested compounds failed to meet acceptability criteria — that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1968)
GC-2131 [1-chloro-2,4- phenylene-bis- (0,0-diethyl phos- phorothiolate]	Australorbis glabratus	BSA and FL	Puerto Rico	Failed	С	Comment same as above.	Seiffer and Schoof (1968)
GC-3582 (EC4)	Gambusia affinis Rana catesbeiana	FL	Ponds in III.	(0)	_	When applied at 0.025 and 0.05 pound per acre active in- gredient, 4 and 80 percent fish mortality occurred in 1 day. When applied at 1.6 pounds per acre, 100 percent bullfrog mortality occurred in 1 day.	Mulla, et al (1963)

	GC-3582 (EC4)	Micropterus salmoides Cyprinus carpio	BSA	-	0.05 (O) 0.10 (K1) 0.005 (K1) 0.01 (K 5 hr) 0.05 (K 3 hr)	a e	At 0.05 ppm, 60 percent mortality occurred in 1 day. For bass: Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds. For carp: Experiments were carried out in plastic tubs lined with saran plastic. Fish weights averaged 217 grams.	Mulla, et al (1967)
	GC-3583 (EC4)	Micropterus salmoides	BSA	-	0.10 (O) 0.50 (K1)	a e	No mortality occurred at 0.10 ppm in 4 days. Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.	Mulla, et al (1967)
	GC-3707 (tech)	Lepomis macrochirus	BSA	-	600 (T1A)	а	The experiment was conducted at 75 F.	Соре (1963)
	GC-3707 (EC)	Salmo gairdneri	BSA	-	170 (T1A)	а	The experiment was conducted at 65 F.	Соре (1963)
	GC-3707 (WP)	Salmo gairdneri	BSA	-	95 (T1A)	а	The experiment was conducted at 65 F.	Cope (1963)
	GC-4072 (50 percent EC)	Gambusia affinis	FL	Ponds Bakers- field, Cal.	0.2 (K1) 0.8 (K1)	a c	Toxicity values indicate application rates in Ib/acre. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
J	GC-4072 (tech)	Lepomis macrochirus	BSA	_	3 (T1A)	а	The experiments were conducted at 75 F.	Cope (1963)
	GC-4072 (EC4)	Micropterus salmoides	BSA	_	0.50 (O) 1.00 (O)	a e	At 0.50 ppm, 6 percent mortality occurred in 1 day. At 1.00 ppm, 66 percent mortality occurred in 1 day. At 1.50 ppm, 100 percent mortality occurred in 2 days. Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.	Mulla, et al (1967)
	GC-9160 (EC2)	<i>Gambusia affinis</i> Tadpole shrimp	FL	Cal.	(O) (O)	-	At an application rate of 2.0 lb/acre, 62% mortality of the fish occurred in 24 hours. Tadpole shrimp survived this treatment.	Mulla (1966)
	GS-12968 (EC4)	Gambusia affinis	FL	Cal.	(O)	-	At a concentration of 0.4 lb/acre, 72% mortality of the fish occurred in 24 hours.	Mulla (1966)
COMN	GS-13005 (EC4)	Gambusia affinis Rana catesbeiana	FL	Cal.	0.4 (K1) (O)	-	Toxicity value in Ib/acre. No mortality in tadpoles of <i>R. catesbeiana</i> occurred during an exposure period of one week.	Mulla (1966)
ERCIAL CHEM	GS-13005	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours.	Jamnback and Frempong- Boadu (1966)
COMMERCIAL CHEMICAL PRODUCTS	Gamlen CW	Pandalus montagní Cardium edule	BSA	-	14.6 (O) 69.5 (O)	<u>a</u> e	<ul> <li>Experiments were conducted in tanks holding 10 liters of sea water at 15 C.</li> <li>It was shown that the toxicity of this solvent emulsifier decreased with time due to evaporation of the solvent.</li> <li>Gamlen CW at a concentration of 33.3 ppm killed 95% of Crangon crangon larvae in 3 hr.</li> </ul>	Portmann and Connor (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Gamlen D	Pandalus montagni Crangon crangon Cardium edule	BSA	-	11.5 (T2A) 9.6 (T2A) 38.8 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of sea water at 15 C. It was shown that the toxicity of this solvent emulsifier de- creased with time due to evaporation of the solvent. Gamlen D at a concentration of 33.3 ppm killed 100% of <i>Crangon crangon</i> larvae in 3 hr.	Portmann and Connor (1968)
Gamlen OSR	Pandalus montagni Crangon crangon Carcinus maenas Cardium edule	BSA	-	12.5 (T2A) 8.8 (T2A) 20.4 (T2A) 15.8 (T2A)	<u>a</u> e	Comments same as above except that Gamien OSR at a concentration of 10 ppm killed 95% of <i>Crangon crangon</i> larvae in 3 hr.	Portmann and Connor (1968)
Gammexane powder (larvicide)	Tilapia melanopleura	FLCH	Tanganyika	1 lb (0% K)	-	Trial periods were for 20 weeks. Sublethal effects such as impaired breeding, retarded growth, or altered taste were not detected. Dosages are given as Ib/acre of surface water,	Webbe and Shute (1959)
Gamosol solvent "D"	Daphnia magna	BSA	-	13.7 (T1A) 2.9 (T2A) 1.5 (T3A)	е	Crude oil plus emulsifier had the following values. 24.4 (T1A) 10.7 (T2A) 9.1 (T3A)	Dowden (1962)
Garlon	Myriophyllum heterophyllum Utricularia sp	FL	Farm ponds in Ga.	(0)	-	Garlon was developed as an overall herbicide containing 4 Ib/gal dalapon and 1/2 Ib/gal Silvex acid. It has given in- dications of control of several species of weeds, such as <i>Myriophyllum heterophyllum</i> and <i>Utricularia</i> sp. However, present results are inconclusive and this herbicide warrants further investigation and experimentation.	Thomaston, et al (1959)
Guthion (25% WP)	Green sunfish Orange spotted sunfish White crappie Bluegill Largemouth bass Gizzard shad Freshwater drum Gar Carp Longnose gar Golden shiners	FL	Ponds in Ark.	0,25-1.8 (O)	d g	Catfish were more tolerant to Guthion than the other species of fish tested. All other species of fish were quickly affected by applications of 1.0 ppm. Field studies were conducted in ponds ranging from 0.25-1.8 acre-feet in volume. Survival values were determined by draining the ponds whenever pos- sible. Residue studies indicated that the chemical disappeared from the water in less than 2 weeks and that the chemical is no longer detectable in catfish flesh after 6 weeks. Cladocera and rotifers were not eliminated from the treated ponds and, in many field collections, were more numerous than in un- treated ponds.	Surber (1943)

	Fathead minnows Bigmouth buffalo Black bullheads Channel catfish Warmouth							
Guthion	Pimephales promelas	BSA	-	0.09 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)	
	Lepomis macrochirus			0.005 (T4A)				
Guthion	Gambusia affinis	BSA	_	0.05 (K 53%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)	
Guthion	Pimephales promelas	BSA	-	0.093 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as	Tarzwell (1959)	
9	Lepomis macrochirus			0.0052 (T4A)		toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	(1959)	APPENDIX
Guthion	Lepomis macrochirus	BSA		0.1 (O)	acdf	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The	Weiss (1959)	Ï
	Micropterus salmoides			0.1 (O)		inhibition of the enzyme was a function of the concentra- tion of the insecticide, extent of exposure, and specific	(1999)	
	Notemigonus crysoleucas			0.1 (O)		chemical nature of the compound.		Β
	Carassius auratus			0.1 (O)				
Guthion	Oncorhynchus kisutch	BSA	_	4.2 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone, TL <sub>m</sub> expressed in ppb.	Katz (1961)	
	Oncorhynchus tshawytscha			4.3 (T4A)				
8	Salmo gairdnerii			3.2 (T4A)				
OMME	Gasterosteus aculeatus			12.1 (T4A)				
RC Guthion	Micropterus salmoides	BSA	-	0.5 (O)	-	The degree of reaction to the cholinesterase-inhibiting insecti- cides is not only a function of time and concentration, but	Weiss (1961)	
COMMERCIAL CHEMICAL PRODUC	Pimephales promelas			0.5 (O)		also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is ex- pressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chem- ical to kill 50 percent of the fish. For bass the $LT_{50}$ was 40 min and for the fathead 40 min.	(1991)	
PRODUC	Cyprinodon variegatus	BCFCH	_	0.01 (SB1)	а	Little or no information was given about test procedures and further results.	Das and Needham (1961)	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Guthion (tech, 90 percent)	Carassius auratus Lebistes reticulatus	BSA	~	1.4 (T4A) 0.12 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)
Guthion (EC1.5)	Gambusia affinis Bufo boreas Scophiopus hammondi	FL	Ponds in III.	(O) (O) (O)	_	When applied at 0.1 pound per acre active ingredient, 100 percent fish mortality occurred in 1 day. No toad mortality occurred at 0.4 pounds per acre in 1 day.	Mulla, et al (1963)
Guthion	Carassius auratus Lepomis macrochirus Notemigonus crysoleucas	BSCH	~	1.0 (O)* 0.1 (O)** 10.0 (O)* 1.0 (O)* 0.1 ? (O)* *response, 15 days **no response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
uthion	Gammarus Iacustris	BSA	-	0.000126 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Guthion	Bluegill	BSA	_	0.0052 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Guthion	Acroneuria pacifica Ephemerella grandis Gammarus lacustris Pteronarcys californica	BSA	_	0.0085 (T4A) 0.014 (T4A) 0.00013 (T4A) 0.022 (T4A)	<u>a</u> c	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
Suthion	Pteronarcys californica Acroneuria pacifica Ephemerella grandis Gammarus lacustris	BSA	~	0.02 (T4A) 0.009 (T4A) 0.01 (T4A) 0.0001 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were de- termined by standard methods. General comments were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional data are presented.	Gaufin, et al (1965)

	Guthion (25% WP)	Ictiobus cyprinellus Ictalurus puntatus Lepomis cyanellus Rana catesbeiana Notemigonus crysoleucas Lepomis macrochirus Micropterus salmoides	BSA	-	(O) 9.0 (T2A) 9.0 (T2A) 0.025 (T2A) (O) 0.10 (T2A) 0.025 (K2) 0.025 (T2A)	a c d e	At 1.0 ppm concentration, bullfrog tadpoles and bigmouth buffalo were not affected. The compound performed effectively under various water conditions which included water from a bayou, lake, and in ponds filled with well water. 1.0 ppm of Guthion effectively controlled green sunfish without apparent effect on channel catfish.	Meyer (1965)
	Guthion (tech, 95 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica (naiad)	BSA	-	0.0220 (T4A) 0.0085 (T4A)	c d e f	<i>A. pacifica</i> was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than <i>P. californica.</i>	Jensen and Gaufin (1966)
P 13	Guthion	Acroneuria pacifica Pteronarcys californica	BSA & CFCH	-	0.0085 (T4A) 0.00024 (T30CH) 0.022 (T4A) 0.0013 (T30CH)	a c d e	Additional data are presented.	Jensen and Gaufin (1966)
ř	Guthion	Lepomis gibbosus	BSA	-	1/4 (O)	_	The figures given are for mortality in 2 hours when the amount of chemical was 16 mg/kg, given by injection.	Murphy (1966)
	Guthion	Lepomis gibbosus lctalurus melas Pseudopleuronectes americanus Myoxocephalus scorpius		-	8.74 ± 1.72 (O) 3.64 ± 0.67 (O) 11.24 ± 1.60 (O) 0.03 ± 0.01 (O)	-	This paper is a study of the amounts of organic thiophosphate and their oxygen analogues which accumulate in liver slices in an <i>in vitro</i> study of insecticides. The numbers given are for m $\mu$ m of chemical (in the case of Parathion, Malathion, and Guthion — the oxygen analogue) accumulated in 100 mg (dry weight) of liver in 30 minutes.	Murphy (1966)
COMM	Guthion	Leiostromus xanthurus Cyprinodon variegatus	BCFCH	_	0.01 (O) 0.01 (O)	а	At a concentration of 0.01 ppm, the following percent acetyl- cholinesterase activity as compared to controls was found: <i>L. xanthurus</i> – 79 <i>C. variegatus</i> – <10.	Butler and Johnson (1967)
ERCIAL	Guthion (EC2)	Micropterus salmoides	BSA	-	1.0 (O) 1.50 (K1)	a e	At 10 ppm no mortality occurred in 1 day. Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.	Mulla, et al (1967)
COMMERCIAL CHEMICA	Guthion	Pteronarcys californica (naiads)	BSA	_	0.0015 (T4A)	acdef	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Guthion	Lepomis gibbosus Ictaluras melas Micropterus dolomieui Myxocephalus scorpius Pseudopleuronectes americanus	BSA	_	(O)	a p	The chemicals were poor inhibitors of brain cholinesterases <i>in vitro</i> ; their oxygen analogs were potent inhibitors.	Murphy, et al (1968)
Gutoxon	Lepomis gibbosus Ictalurus melas Pseudopleuronectes americanus Myxocephalus scorpius		-	0.205 ± 0.010 (O) 0.101 ± 0.044 (O) 0.039 ± 0.030 (O) 0.109 ± 0.020 (O)	_	This paper is a study of the amounts of organic thiophos- phate and their oxygen analogues which accumulate in liver slices in an <i>in vitro</i> study of the insecticides. The numbers given are for mµm of the chemical accumulated in 50 mg of liver (wet weight) in 10 minutes.	Murphy (1966)
Gutoxon	Lepomis gibbosus	BSA	-	4/4 (O)	-	The figures given are for mortality in 2 hours when the amount of chemical was 1.0 mg/kg, given by injection.	Murphy (1966)
HCA	Panicum hemitomum Pontederia cordata Spatterdock	FL	Fla.	(0)	-	The degree of control was as follows:P. hemitomum(80 lb/acre) - 85 percentP. cordata(80 lb/acre) - 85 percentspatterdock(160 lb/acre) - none.	Copeland and Woods (1959)
Hept	Channel catfish (fingerlings)	BSA	-	12.4 (K 25 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Heptachlor	Various	FL	Salt Lake Co., Utah	(O)	-	<ul> <li>The chemical was applied at 0.1 lb/acre.</li> <li>At the above concentration no ill effects were observed in mammals, birds, reptiles, and amphibians.</li> <li>Invertebrates were not affected uniformly. Crustaceans were not harmed, nor were larvae of the insect family Ephydridae.</li> <li>Spiders and aquatic insects other than Ephydridae were adversely affected in varying degrees. Aquatic beetles seemed to be affected more seriously than other insects excenting mosquito larvae</li> </ul>	Graham and Anderson (1958)

excepting mosquito larvae.

He	eptachior	Dorosoma cepedianum Esox americanus Erimyzon bucetta Notemigonus crysoleucas Opsopoeodus emiliae Ictalurus melas Fundulus chrysotus Gambusia affinis Aphredoderus sayanus Microp terus salmoides Chaenobryttus coronarius Lepomis symmetricus L. megalotis L. megrochirus Pomoxis	FR	Texas	(0)	_	<ul> <li>Experiments were conducted in fish streams and canals which bisect a farm treated with 10% Heptachlor at the rate of 20 pounds per acre to control fire ants.</li> <li>Fish were showing symptoms of distress 3 days after application of the Heptachlor. In a depression filled with water in a rice field 100% kill was noted for <i>Gambusia</i> and bantam sunfish (the most abundant species). There was no effect on tadpoles.</li> </ul>	Boudreaux, et al (1959)	APPENDIX
		nigromaculatus Etheostoma gracile							EX B
не С0	eptachlor	gracine Fathead minnow Bluegill Goldfish Guppy	BSA	-	0.094 (T4A) 0.019 (T4A) 0.230 (T4A) 0.170 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in soft water.	Tarzwell (1959)	
2	eptachlor	Fathead minnow	BSA	-	0.056 (T4A)	<u>a</u>	Comment same as above except that this experiment was performed in hard water.	Tarzwell (1959)	
₽ (	eptachlor (25%)	Channel catfish (fingerlings)	BSA	_	1.8 (K 24 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	
I CHEMICAL PRODUCT	eptachlor	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	0.09 (T4A) 0.02 (T4A) 0.23 (T4A) 0.11 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)	

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Heptachlor	Pimephales	BSA	_	0.18 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were	Henderson, et al
(72%)	promelas Lepomis			0.026 (T4A)		conducted with hard water.	(1959)
	macrochirus Carassius			0.320 (T4A)			
	auratus Lebistes reticulatus			0.148 (T4A)			
Heptachlor	Pimephales promelas	BSA	-	0.094 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as	Tarzwell (1959)
	Lepomis macrochirus			0.019 (T4A)		toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly in- fluenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	(1959)
Heptachlor	Daphnia magna	BSA	-	0.05777 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
Heptachlor	Oncorhynchus kisutch	BSA	-	59.0 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in PPB.	Katz (1961)
	Oncorhynchus tshawytscha			17.3 (T4A)			(,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
	Salmo gardnerii			19.4 (T4A)			
	Gasterosteus aculeatus			111.9 (T4A)			
Heptachlor (heptachloro- 4,7-methano- tetrahydro- indene)	Richardsonius balteatus hydroflox	BSA	-	>0.13 (T1A) 0.11 (T2A) 0.096 (T4A)	acdef	Results given were in soft water. Results in hard water were as follows: 0.15 (T1A), 0.12 (T2A), and 0.11 (T4A).	Webb (1961)
Heptachlor	Lepomis microlophus	BSA	-	0.02-0.09 (T1A)	<u>a</u>	This is a time-temperature study with considerable additional data presented.	Brown (1961)
Heptachlor	Gammarus Iacustris Iacustris	BSA	-	(O)	a e p	The mortality might have been partially due to the suscept- ibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Re- sults were expressed as $LT_{50}$ ; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 240 (±30) min.	McDonald (1962)
Heptachlor	Salmo gairdneri	BSA	-	150 (T1A) 90 (T2A) 70 (T4A)	а	The experiments were conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)
	Redear sunfish			70 (14A) 0.092 (T1A) 0.064 (T1A) 0.047 (T1A) 0.034 (T1A) 0.032 (T1A)		The experiment was conducted at 45 F. The experiment was conducted at 55 F. The experiment was conducted at 65 F. The experiment was conducted at 75 F. The experiment was conducted at 85 F. Higher temperatures caused a moderate increase in toxic effects.	

	Heptachlor	Lepomis cyanellus Lepomis macrochirus Rana catesbeiana	FL	Miss.	(0)	_	Limited mortality of fish and amphibians occurred as a result of Heptachlor applications used to control fire ants. At a concentration of 2.0 (Ib/acre) only one bullfrog was killed during the entire study. At a concentration of 0.25 plus 0.25 lb/acre (2 applications approximately 4 months apart) there were 8 dead green sunfish.	Ferguson (1963)
	Heptachlor (EC2)	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	ac	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and resulted in a 28 percent mortality for the fish, and a 50 percent mortality for the tadpoles in 24 hours.	Mulla (1963)
	Heptachlor	Gambusia affinis affinis	BSA	-	0.07 to 1.3 (O)	а	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are later placed in clean water.	Boyd and Ferguson (1964)
	Heptachlor	Bluegill	BSA		0.019 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
	Heptachlor (tech)	Rainbow trout	BSA	-	0.008 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
B-141	Heptachlor	Salmo gairdneri Lepomis macrochirus Pteronarcys californica Baetis sp Daphnia pulex Simocephalus serrulatus	BSA	-	0.009 (T2A) 0.026 (T2A) 0.006 (T2A) 0.032 (T2A) 0.042 (T2A) 0.047 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on re- production and behavior are also discussed. Data pre- sented as EC <sub>50</sub> .	Cope (1966)
COMME	Heptachlor	Simocephalus serrulatus Daphnia pulex	BSA	-	0.047 (SB) 0.042 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
COMMERCIAL CHEN	Heptachlor	Daphnia carinata	BSA	-	0.02 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr, Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during	Sanders and Cope (1966)

the assay period.

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Heptachlor	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	_	(O)	_	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentration of 250-2500X resulted.	Butler (1967)
Heptachlor	Oyster	FE	Galveston Bay, Texas	(O)	_	The chemical was found in the water at a concentration of $<$ 0.001 ppm. Oysters from the area were found to contain $<$ 0.01 ppm.	Casper (1967)
Heptachlor	Micropterus salmoides salmoides Lepomis macrochirus Chelydra serpentine	FL	Va.	(O)	_	The amount of chemical applied was not specified in this report. None of the chemical was found in the tissues of either bass or bluegill taken from polluted ponds. However, a snapping turtle taken from a pond that had no residue of chemical in the water or bottom mud was found to contain 5100 ppb in the body fat, egg yolk, and liver tissues.	Weatherholtz, et al (1967)
Heptachlor	Pteronarcys californica (naiads) Pteronarcella badia (naiads) Claasenia sabulosa (naiads)	BSA	-	0.0011 (T4A) 0.0009 (T4A) 0.0028 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
Heptachlor	Alosa pseudoharengus Aplodinotus grunniens Coregonus artedii Lota lota	BSA	-	(0)	_	The study showed that the levels of chlorinated hydrocarbon pesticide residues in fish meals and oils were, with the ex- ception of the oil sample taken from the Lake Michigan alewife, below the regulatory tolerances established by the Food and Drug Directorate of Canada (1965) for certain foods intended for human consumption. Pesticide levels were interpreted as being representative for each species.	Dugal (1968)
Heptachlor epoxide	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto Albe	FO	Netherlands	(0)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon insecti- cides. In most cases the liver had the highest concentration of toxicant, ranging from 0.07 to 4.7 ppm. Most chlorinated hydrocarbons tend to accumulate in the fat depots of the body. In instances where mesenterial fat was found the con- centration of toxicant was found to be as high as 3.0 ppm.	Koeman and van Genderen (1966)

		strix aluco Osio otus Falco pereginus						
	Heptachlor epoxide	Alosa pseudoharengus Aplodinotus grunniens Coregonus artedii Lota lota	BSA	-	(O)	-	The study showed that the levels of chlorinated hydrocarbon pesticide residues in fish meals and oils were, with the ex- ception of the oil sample taken from the Lake Michigan alewife, below the regulatory tolerances established by the Food and Drug Directorate of Canada (1965) for certain foods intended for human consumption. Pesticide levels were interpreted as being representative for each species.	Dugal (1968)
	Hercules 528	Lepomis macrochirus	BSA	-	0.1 (O)	acdf	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The	Weiss (1959)
		Micropterus salmoides			0.1 (0)		inhibition of the enzyme was a function of the concentra- tion of the insecticide, extent of exposure, and specific	
		Notemigonus crysoleucas			0.1 (0)		chemical nature of the compound.	
		Carassius auratus			0.1 (O)			
B-143	Hercules 3895 G	Gambusia affinis	BSA	-	0.05 (K 0%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hours.	Lewallen (1959)
ώ	Hercules 7175 (tech)	Lepomis macrochirus	BSA	_	40,000 (T1A)	а	The experiment was conducted at 75 F. Fish weighed 0.4 g.	Cope (1963)
	Hercules 7531 (tech)	Lepomis macrochirus	BSA	-	25,000 (T1A)	а	Comment same as above.	Cope (1963)
	HRS-1622 (octachloro- propane)	Australorbis glabratus	BSA and FL	Puerto Rico	Failed	с	Seven of the tested compounds failed to meet acceptability criteria — that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
8	Hyamine	Oncorhynchus kisutch	BSA		57 (T1A) 53 (T2A)	a c d e	Concentrations were based on percent active ingredient.	Bond, et al (1960)
COMMERCIAL CHEMICAL PRODUCTS	Hyamine 1622	Lepomis macrochirus Pimephales promelas	BSA	-	(S) 1.6 (T4A) (H) 3.8 (T4A) (S) 1.6 (T4A) (H) 3.8 (T4A)	ace	Bioassay method in Standard Methods for examination of water was used. Both hard (H) and soft (S) water were used. TL <sub>m</sub> values for 24 and 48 hr are also presented.	Surber and Pickering (1962)
. CHEMICA	Hyamine 2389	Lepomis macrochirus Pimephales promelas	BSA	-	(S) 1.2 (T4A) (H) 4.8 (T4A) (S) 2.4 (T4A) (H) 4.2 (T4A)	<u>a</u> c <u>e</u>	Comment same as above.	Surber and Pickering (1962)
L PR	Hydram	Oyster	BCF	-	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
ODUCTS	Hydram	Penaeus aztecus	L	-	(O)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. A con- centration of 1.0 ppm caused 10 percent mortality.	Butler (1965)

Strix

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Hyd <i>r</i> am	Leiostomus xanthurus (juvenile)	BSA	-	(0)	a	Water temperature was 25 C. 20% mortality at 1.0 ppm.	Butler (1965)
Hydram	Crassostrea virginica Penaeus az tecus Leiostomus xanthurus Phy toplankton	BCFA & BSA	-	1.0 (NTE) 1.0 (0, 30%) 1.0 (20% T2CFA) 9% (O)	_	<ul> <li>Sea water was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Hydram	<i>Pteronarcys</i> sp (nymphs)	BSA	_	0.370 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Соре (1965)
Hydram (tech)	Rainbow trout Bluegill	BSA	-	0.200 (T4A) 0.355 (T4A)	а	This is an estimated LC $_{\rm 50}$ value at temperatures from 55 to 75 F.	Cope (1965)
Hydram	Salmo gairdneri Lepomis macrochirus Pteronarcys californica	BSA	-	0.290 (T2A) 0.475 (T2A) 0.700 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on reproduction and behavior are also discussed. Data presented as $EC_{50}$ .	Cope (1966)
Hydram (tech)	Rainbow trout Bluegill	BSA	-	0.29 (T2A) 0.475 (T2A)	_	Data are given as LC <sub>50</sub> .	Bohmont (1967)
Hydrothal 191	Rainbow trout	BSA	-	1.5 (T2A)	-	Comment same as above.	Bohmont (1967)
Hydrothal plus	Lepomis macrochirus	BSA	-	3.5 (T1A)	a b e	This report is a simple and straightforward determination of a median tolerable limit for a selected group of herbicides.	Hughes and Davis (1967)
Ibcol	Guppy	BSA	-	100 (K1)	<u>a</u>	Those fish that survived at lower concentrations were still very active several days after they had been taken out and placed in fresh water.	Anonymous (1964)
lmidan	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	4.0 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxica- tion. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 85 percent.	Jamnback and Frempong- Boadu (1966)

	Inverton (2,4,5-T)	<i>Nympheae</i> sp Parrot feathers	FL	Farm ponds in Georgia	(0)	-	Nympheae sp and parrot feathers were killed at the recom- mended application rate of one gallon Inverton mixed with 15 gallons of fuel, and 84 gallons of water per acre killed completely in less than a week.	Thomaston, et ( (1959)	al
	lodophor	Guppy	BSA	_	6250 (K1)	<u>a</u>	Those fish that survived at lower concentrations were still very active several days after they had been taken out and placed in fresh water.	Anonymous (1964)	
	IPC 50%	Channel catfish (fingerlings)	BSA	-	>100 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	
	IPC (tech)	Bluegill	BSA?	_	29.0 (T4A)	а	This is an estimated LC $_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)	
	IPC	Simocephalus serrulatus	BSA	-	10.0 (SB)	-	Concentration reported is for immobilization.	Sanders and	
		pulex			10.0 (SB)		Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Cope (1966)	
	IPC (tech)	Bluegill	BSA	_	32.0 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)	
	lsodrin (EC 1.6)	Gambusia affinis	FL	Cal.	0.5 (O)	a c	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in lb/acre, and resulted in a 100 percent mortality for these fish.	Mulla (1963)	APPENDIX
	lsolan (EC2)	Gambusia affinis	FL	Cal.	2.0 (K1)	-	Toxicity value is in Ib/acre.	Mulla (1966)	χ Φ
	lsotex 25	Channel catfish (fingerlings)	BSA	_	0.54 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	
	K-6882 (25 percent EC)	Gambusia affinis	FL	Ponds — Bakersfield, Cal.	(O)	ac	No fish mortality occurred at 0.2 to 0.8 lb/acre rates of application. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)	
COMMERCI	Karmex W <sub>C</sub> [3-(p-chloro- phenyl)-1,1- dimethylurea) 80% active ingredient]	Richardsonius balteatus hydroflox	BSA	_	42.5 (T1A) 41.5 (T2A) 41.5 (T4A)	a c d e f	Results given were in soft water. Results in hard water were as follows: 60.3 <sup>+</sup> (T1A), 41.2 (T2A), and 40.1 (T4A).	Webb (1961)	
	Kelthane	Salmo gairdneri	BSA	_	110 (T1A)	а	The experiment was conducted at 55 F. Fish weighed 0.7 g.	Cope (1963)	
CHEMICAL	Kelthane	Gambusia affinis	BSA	-	2.1 (L1)* 1.9 (L1)** *Resistant fish **Nonresistant fish	а	This paper deals with the resistance of mosquito fish to chlorinated hydrocarbon compounds. Resistant fish were not always less sensitive to these chemicals.	Boyd and Ferguson (1964)	
PRODU	Kepone	Lepomis microlophus	BSA	_	0.1-0.6 (T1A)	<u>a</u>	This is a time-temperature study with considerable addi- tional data presented.	Brown (1961)	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Kepone	Lepomis macrochirus Redear sunfish	BSA	_	380 (T 18 hr) 240 (T 32 hr) 110 (T 56 hr) 0.62 (T1A) 0.54 (T1A) 0.34 (T1A) 0.24 (T1A) 0.12 (T1A)	а	The experiment was conducted at 65 F Fish were 2 in. long. The experiment was conducted at 45 F. The experiment was conducted at 55 F. The experiment was conducted at 65 F. The experiment was conducted at 75 F. The experiment was conducted at 85 F. Higher temperatures caused a moderate increase in toxic effects.	Cope (1963)
Керопе (EC 2)	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (0)	a C	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and re- sulted in a 18 percent mortality for the fish, and a 0 percent mortality for the tadpole in 24 hr.	Mulla (1963)
Kepone (tech)	Rainb <sup>o</sup> w trout	BSA	_	0.020 (T4A)	-	The values reported are given as $LC_{\overline{50}}$ .	Соре (1965)
Knoxweed 42	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	(0)	а	Water temperature was 25 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Knoxweed 42	Oyster	BCF	_	(0)	а	Exposure to a concentration of 1 ppm caused a 44.0% decrease in shell growth.	Butler (1965)
Knoxweed 42	Penaeus aztecus	L	-	(O)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. A concentration of 1.0 ppm caused 40 percent mortality.	Butler (1965)
Knoxweed 42	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	A _	1.0 (O) 0.48 (O) 1.0 (NTE) 1.0 (NTE)	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Korlan	Gambusia affinis	BSA	-	0.1 (K 3%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given average percent fish killed in 24 hr.	Lewallen (1959)

	Kuron	<i>Najas quadalupensis</i> Spatterdock	FL	Fla.	(0)	-	At 20 lb/acre, <i>N. quadalupensis</i> was not controlled while 5-10 percent control of spatterdock was obtained.	Copeland and Woods (1959)
	Kuron .	Onchorynchus tshawytscha Micropterus salmoides	BSA	_	1.35 (T1A) 1.23 (T2A) 3.5 (T1A)	acde	Concentrations were based on percent active ingredient.	Bond, et al (1960)
	Kuron (silvex acid equivalent)	Micropterus salmoides (fry) Ictalurus punctatus (fry) Lepomis macrochirus (fry)	BSA	_	1.0 (SB3) 0.5 (SB3) 0.3 (SB3)	ac de f p	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
	Kuron	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.320 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
	Kuron	Chinook salmon Bluegill	BSA	_	1.35 (T1A) 1.23 (T2A) 2.9 (T1A) 2.4 (T2A)	ac d	Tests were conducted in glass jars holding 15 liters of water. Toxicity of Kuron varies with the supplier.	Bond, et al (1965)
	Kuron	Simocephalus serrulatus Daphnia pulex	BSA	_	2.4 (SB) 2.00 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Kurosal	Lepomis macrochirus	BSA	-	120,000 (T1A)	а	The experiment was conducted at 75 F. Fish weighed 0.6 g.	Соре (1963)
	Kurosal G	Bluegill	BSA	-	21 (T1A) 15 (T2A)	acd	Tests were conducted in glass jars holding 15 liters of water.	Bond, et al (1965)
_	Kurosal G (silvex acid equivalent)	<i>Lepomis macrochirus</i> (fry)	BSA	_	150 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
OMMEDO	Kurosal SL	Oncorhynchus kisutch	BSA	-	290 (T1A) 240 (T2A) 83 (T1A) 83 (T2A)	a c d	Tests were conducted in glass jars holding 15 liters of water. Active ingredient of Kurosal SL is silvex 2-{2,4,5-trichloro- phenoxy) propionic acid, potassium salt.	Bond, et al (1965)
	Kurosal SL (silvex acid equivalent)	Lepomis macrochirus (fry)	BSA	-	100 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Kurosal SL (60% silvex)	Crassostrea virginica Fundulus similis Phy toplankton	BCFA & B	SA —	1.0 (NTE) 25.0 (NTE) 1.0 (NTE)	_	Seawater was pumped continuously into test aquaria.Salinity, temperature, and plankton fluctuated with tide,and ambient weather conditions. Some bioassays withfish were static. Toxicity was reported for the following:Oyster –96-hr EC50 – Conc. which decreasedshell growth.Shrimp –48-hr EC50 – Conc. which killed orparalyzed 50% of test animals.Fish –48-hr EC50 – Conc. which killed50%.Phytoplankton – Percent decrease of CO2 fixation toa 4-hr exposure at 1.0 ppm chemicalconcentration.	Butler (1965)
Kyro-eo (nonionıc)	Brook trout Rainbow trout Mayflies (Ephemeroptera naiads) Stoneflies (Plecoptera naiads) Damsel flies (Odonata naiads) Diptera larva (chiefly Tendipedidae)	BSA	_	5.2 (L1) 4.8 (L2) 4.6 (L3) 5.5 (L1) 5.3 (L2) 5.1 (L3) 5.6 (L2) 5.4 (L3) 5.2 (L4) 5.1 (L2) 4.8 (L3) 4.7 (L4) 5.2 (L3) 4.9 (L4) (O)	_	Kyro-eo is a synthetic, non-sulfonated detergent. Experi- mental water was not supplemented with oxygen because bubbling caused suds. Control tanks were also static. Control organisms apparently did not suffer from lack of oxygen in the static aquarium for 120 hr. Dipteran larvae withstood 10 ppm of Kyro-eo with no mortality.	Hepworth (no date)
LAS (degradation product sulfophenyl- undecanoic acid, disodium salt)	Bluegill (fingerlings)	BSA	-	75.0 (T4A)	c d <u>e</u> f o	The fish killed all showed severe hematomas of the respiratory folds of the gills. This was followed by the stripping of the mucous layers. Following this, soft tissue beneath was completely destroyed in most cases.	Swisher, et al (1964)
LAS C 12 (alkylbenzene sodium sulfonate)	Bluegill (fingerlings)	BSA	-	3.0 (T4A)	c d <u>e</u> f o	Comment same as above.	Swisher, et al (1964)
LAS C 14 (alkylbenzene sodium sulfonate)	Bluegill (fingerlings)	BSA	-	0.64 (T4A)	c d <u>e</u> f o	Comment same as above.	Swisher, et al (1 <del>964</del> )

	LAS	Pimephales promelas (eggs)	BCF	-	2.3 (T4A)	<u>a</u> cdef	Mortality range is given for exposure (days 1-9) with various concentrations and controls. Additional data are presented.	Pickering (1966)	
1	Linear alkyl sulfonate	Lepomis macrochirus	BSA	-	4.0 (T4A)	acde	In all these tests the LAS stock powder contained 60.8% LAS. The values reported were calculated on a basis of	Thatcher and Santner	
		Pimephales promelas			4.2 (T4A)		pure LAS.	(1967)	
		Ictalurus melas			6.4 (T4A)				
		Notropis atherinoides			3.3 (T4A)				
		Notropis cornutus			4.9 (T4A)				
	Lethane 384	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	4.0 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 2 percent.	Jamnback and Frempong- Boadu (1966)	
	Lexone	Channel catfish (fingerlings)	BSA	-	5.2 (K 30 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	VDD
	Lignasan	Channel catfish (fingerlings)	BSA	_	2.0 (K 28 hr A)	<u>a</u>	Comment same as above.	Clemens and Sneed (1959)	APPENDIX
	Lignasan	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	-	0.006 (K) 0.006 (K) 0.06 (K) 0.06 (K)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants.		(B
COMMERCIAL CHEM	Lindane	Fathead minnow Bluegill Goldfish Guppy	BSA	-	0.062 (T4A) 0.077 (T4A) 0.152 (T4A) 0.138 (T4A)	<u>a</u> _	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in soft water.	Tarzwell (1959)	
HEMIC	Lindane	Fathead minnow	BSA	-	0.056 (T4A)	<u>a</u>	Comment same as above, except experiment was conducted in hard water.	Tarzwell (1959)	

COMMERCIA	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
r	Lindane 3% (Methoxychlor 50%)	Channel catfish (fingerlings)	BSA	_	2.0 (K1A)	a	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
AICAL PI	Lindane	Pimephales promelas Lepomis	BSA	-	0.06 (T4A) 0.09 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
CHEMICAL PRODUCTS		macrochirus Carassius auratus Lebistes			0.15 (T4A) 0.14 (T4A)			
0,	Lindane (100%)	reticulatus Pimephales promelas	BSA	-	0.062 (T4A)	<u>a</u> becd <u>f</u>	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et al (1959)
		Lepomis macrochirus Carassius auratus			0.077 (T4A) 0.152 (T4A)			
		Lebistes reticulatus	204		0.138 (T4A)			<b>T</b>
<b>B-1</b> 50	Lindane	Pimephales promelas Lepomis mæcrochirus	BSA	-	0.062 (T4A) 0.077 (T4A)	a	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not signifi- cantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)
	Lindane	Oncorhynchus kisutch	BSA	-	50.0 (T4A) 40.0 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
		Oncorhynchus tshawytscha Salmo			38.0 (T4A)			
		gairdnerii Gasterosteus aculeatus			44.0 (T4A)			
	Lindane	Gammarus Iacustris Iacustris	BSA	_	(O)	a e p	The mortality might have been partially due to the susceptibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Results were expressed as LT <sub>50</sub> ; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 11 ( $\pm$ 2) min.	McDonald (1962)

	Lindere	8						
	Lindane	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	-	9.0 (O)* 9.0 (O)* 9.0 (O) 9.0 (NG) 7.5 (NG) *obvious, but inhib- ited growth	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medi- um with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
	Lindane	Ophicephalus punctatus Heteropneustes fossilis Barbus stigma Trichogaster fasciatus	BSA	_	4000-5000 (K < 1 hr) 2000-5000 (K < 7 hr) 1000 (K < 2 hr) 2000-3000 (K < 1 hr)	а	The dosage to produce toxic symptoms varied with each species. At the very low dosage, these insecticides did not produce observable changes, but at the higher dosage observable changes were pronounced. Lindane at low concentrations had no noticeable effect but at higher concentrations the rate or mortality was very high.	Mathur (1963)
	Lindane	Golden shiner	BSA	_	≫0.062- 0.125 (O)	ac de p	A number of values for a threshold limit (LD/O) of the toxicant in various solvents are given. Values from 0.062 ppm to 0.125 ppm for Lindane solvents in addition to water were obtained. Acetone and mixed solvents caused the greatest Lindane lethality.	Meyer (1965)
B-151	Lindane EC 1.65	Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a c	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and re- sulted in a 10 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
	Lindane	Gambusia affinis affinis	BSA	-	0.15 to 1.7 (O)	а	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are later placed in clean water.	Boyd and Ferguson (1964)
	Lindane (tech)	Rainbow trout	BSA	-	0.022 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
8	Lindane	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.001 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
COMMERCIAL CHEMICAL PRO	Lindane	Bluegill	BSA	-	65 (T4A) 53 (T4A) 56 (T4A) 38 (T4A) 25 (T4A)	<u>a</u>	These experiments were performed to demonstrate that at increased temperatures the toxic effect of most chemi- cals is increased. For the toxicant concentrations listed, the temperatures were respectively, 45, 55, 65, 75, and 85 F. Data on the effect of time as well as temperature was also reported. The experimental animals all were approximately one gram in weight.	Соре (1965)
CAL PROL	Lindane	Pteronarcys (stone fly nymphs)	BSA	-	0.001 (T4)	<u>a</u>	These experiments were all conducted at 60 F. The values were listed as $\text{LC}_{50}.$	Snow (1958)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Lindane	Salmo gairdnerii Lepomis macrochirus Pteronarcys californica Daphnia pulex Simocephalus serrulatus	BSA	-	0.022 (T2A) 0.053 (T2A) 0.002 (T2A) 0.460 (T2A) 0.520 (T2A)	а	This paper reports acute toxicity of a number of com- pounds, and discusses subacute mortality as well. Effects on reproduction and behavior are also dis- cussed. Data presented as EC <sub>50</sub> .	Соре (1966)
Lindane	Buteo buteo Accipiter gentilis Accipiter nisus Falco tinnunculus Tyto alba Strix aluco Asio otus Falco pereginus	FO	Netherlands	(0)	_	The results of this study show that birds of prey and fish- eating birds found dead in the Netherlands accumulated large amounts of different chlorinated hydrocarbon in- secticides. Most chlorinated hydrocarbons tended to accumulate in the fat depots of the body. In instances where mesenterial fat was analyzed the concentration of toxicant was found to be as high as 89 ppm.	K <i>o</i> eman and van Genderen (1966)
_indane	Puntius javanicus Tilapia mossambica Cyprinus carpio	FL	Japan	1.0% (O) 4.0% (O) 1.0% (O) 4.0% (O) 1.0% (O) 4.0 % (O)	-	No fish deaths occurred at the 1.0 percent concentration. The following mortality occurred at the 4.0 percent level: P. javanicus - 56.5 percent (2 days) T. mossambica - 86.0 percent (2 days) C. carpio - 7.5 percent (2 days)	Kok and Pathak (1966)
_indane	Puntius javanicus Tilapia mossambica Cyprinus carpio	BSA	_	2.0 (K2) 2.0 (K2) 2.0 (0)	e	The purpose of this experiment was to determine the effect of Lindane on three species of fish. The Lindane was Dol granule, a granular formulation containing 6 percent Lindane and 94 percent carrier. With <i>C. carpio</i> , the 2.0 ppm killed 77.5 percent of the test fish in 2 days. The data given are concerned with exposure in water solutions. When soil was added to the water, the mortality was reduced.	Kok and Pathak (1966)
_indane	Simocephalus serrulatus Daphnia pulex	BSA	_	0.520 (SB) 0.460 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)

	Lindane	Daphnia magna	BSA	-	1.1 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Lindane	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	-	(0)	_	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentra- tion of 10-250X resulted.	Butler (1967)
	Lindane	Oyster	FE	Galveston Bay, Texas	(0)	-	The chemical was found in the water at a concentration of $<0.001$ ppm. Oysters from the area were found to contain $<0.01$ to 0.01 ppm.	Casper (1967)
φ	Lindane-C <sup>14</sup>	Lepomis macrochirus Carassius auratus	BCFCH	-	(0)	a	Fish were treated with carbon-labeled insecticides (0.03 ppm) from 5 to 19 hr and uptake rates were determined. They were placed in recovery tanks for up to 32 days. Whole body samples were then made. Almost all of Lindane absorbed was eliminated in 2 days.	Gakstatter and Weiss (1967)
-153	Lindane	Esox Iucius	FR	River Nene, Eng.	(0)	-	Higher concentrations were found in larger fish, indicating that they had been exposed to the pesticides for a longer time. Tissue extracts from the pike were analyzed for organochlorine pesticide residues by gas liquid chroma- tography. The values for large pike were: 0.042 ppm muscle 7.5 ppm fat	Mawdesley- Thomas and Leahy (1967)
	M-502	Althernanthera philoxeroides Pestía stratiotes Spatterdock	FL	Fla.	(0)	-	At 1.0 lb/acre, the degree of control was: <i>A. philoxeroides</i> – 85-90 percent <i>P. stratiotes</i> – 80 percent spatterdock – 3 percent	Copeland and Woods (1959)
COMMERCIAL	M-1499 (granular Silvex)	Bushy pondweed Water Hyssop Parrot's Feather Bladderwort	FL	Lakes in Fla.	(0)	-	Concentrations of 2.3 to 2.5 ppm controlled bushy pond- weed while 1.0 to 4.0 ppm controlled the other species indicated.	Phillippy (1961)
	M-1500	Bushy pondweed	FL	Lakes in Fla.	(O)	-	A concentration of 1.5 ppm controlled bushy pondweed.	Phillippy (1961)
CHEMIC	M-1845 (liquid Silvex)	Spatterdock Bushy pondweed	FL	Lakes in Fla.	(O)	_	A concentration of 0.5 ppm controlled the spatterdock while 1.0 ppm per acre controlled the bushy pondweed.	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
MCP (Amine)	Crassostrea virginica Penaeus aztecus Fundulus similis Phytoplankton	BCFA & BSA	-	1.0 (NTE)	_	$\begin{array}{rcl} \mbox{Seawater was pumped continuously into test aquaria.}\\ \mbox{Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions.}\\ \mbox{Some bioassays with fish were static.}\\ \mbox{Toxicity was reported for the following:}\\ \mbox{Oyster - } & 96\text{-hr EC}_{50} - \text{Conc. which decreased shell growth.}\\ \mbox{Shrimp - } & 48\text{-hr EC}_{50} - \text{Conc. which killed or paralyzed 50\% of test animals.}\\ \mbox{Fish - } & 48\text{-hr EC}_{50} - \text{Conc. which killed 50\%.}\\ \mbox{Phytoplankton - Percent decrease of CO}_2 fixation to a 4-hr exposure at 1.0 ppm chemical concentration.}\\ \end{array}$	Butler (1965)
MCPA (alkyl amine)	Lepomis macrochirus	BSA	-	163.5 (T1A)	_	The bioassay methods employed in this experiment were not given in the paper but it was stated that the same procedures were employed as in previous work.	Davis and Hughes (1963)
МСРА	Lepomis macrochirus	BSA	-	1.5 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
МСРА	Daphnia magna	BSA	_	100 (O)	acdiq	Toxicity, in terms of median immobilization concentration $(1C_{50})$ , is presented.	Crosby and Tucker (1966)
4-(MCPB)	Lepomis macrochirus Micropterus salmoides	BSA	_	15 (T2A) 10 (T2A)	<u>a</u> c o	The response of bluegill and bass fingerlings to nine agricultural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
Malamar-50	Cyprinus carpio C. carpio Tilapia mossambica Cirrhina mrigala Labeo fimbriatus Danio sp Labeo rohita	BSA	-	10.0 (T2A) 8.5 (T2A) 8.3 (T2A) 7.0 (T2A) 8.5 (T2A) 13.5 (T2A) 8.0 (T2A)	acdefp	The test animals were conditioned for 48 hours prior to use.	Sreenivasan and Swaminathan (1967)
Malaoxon	Lepomis gibbosus	BSA	-	2/4 (O)	-	The figures given are for mortality in 2 hours when the amount of chemical was 0.25 mg/kg, given by injection.	Murphy (1966)
Malaoxon	Lepomis gibbosus Ictalurus melas Pseudopleuronectes americanus Myxocephalus scorpius	L	-	1.59 ± 0.17 (O) 0.97 ± 0.28 (O) 0.81 ± 0.09 (O) 1.27 ± 0.14 (O)	~	This paper is a study of the amounts of organic thiophosphate and their oxygen analogues which accumulate in liver slices in an <i>in vitro</i> study of the insecticides. The numbers given are for m $\mu$ m of the chemical accumulated in 50 mg of liver (wet weight) in 10 minutes.	Murphy (1966)
Malathion (25 percent wettable powder)	Cyprinus carpio	BSA	-	(O)	a c e	100 percent mortality occurred in 6 days at 5 ppm, 80 percent mortality occurred in 4 days at 7 ppm.	Hayes (1955)
Malathion (emulsifiable)	Cyprinus carpio	BSA	-	(0)	асе	60 percent mortality occurred in 143 hr at 3 ppm, 0 per- cent mortality occurred in 4 days at 0.01 ppm, 0 percent mortality occurred in 140 hr at 1 ppm, 100 percent	Hayes (1955)

,	Malathion	Fall chinook salmon (fingerlings)	BSA	-	0.17 (T1A) 0.15 (T2A) 0.12 (T4A)	â	At 0.32 ppm there were no survivals after 48 hours.	Parkhurst and Johnson (1955)
N	falathion	Various	FL	Salt Lake Co., Utah	(0)	_	<ul> <li>The chemical was applied at 0.5 lb/acre.</li> <li>At the above concentration no ill effects were observed in mammals, birds, reptiles, and amphibians.</li> <li>Invertebrates were not affected uniformly. Crustaceans were not harmed, nor were larvae of the insect family Ephydridae.</li> <li>Spiders and aquatic insects other than Ephydridae were ad- versely affected in varying degrees. Aquatic beetles seemed to be affected more seriously than other insects excepting mosquito larvae.</li> </ul>	Graham and Anderson (1958)
ſ	Malathion	Pimephales promelas	BSA	-	22.0 (T4A)	<u>acdef</u>	Tests were performed in both hard and soft water. Additional tolerance limit values are given.	Henderson and Pickering (1958)
I	Malathion	Atlantic salmon	BSA	-	0.033 (T1A) 0.033 (T2A)	a e	Results are recorded in ppm of insecticide by weight in water. Changes in temperature had an effect on the toxicity of the chemical.	Keenleyside (1958)
1	Malathion	Fathead minnow	BSA	-	12.5 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. The experiment was performed in hard water.	Tarzwell (1959)
I	Malathion	Daphnia magna	BSA	-	0.0009 (O)	<u>a</u>	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
ł	Malathion 25%	Channel catfish (fingerlings)	BSA	-	>100 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
COMMERC	Malathion	Fundulus ocellaris	FL(E)	Odessa, Del.	(O)	_	The extent of mortality at an application rate of 0.5 Ib/acre in tidal marshes while another 33% of the fish were not affected at all. The fate of those individuals that were poisoned, but remained alive, is in question. The tests indicated that 56% would recover if transferred to fresh water following the exposure period.	Darsie and Corriden (1959)
CIAL	Malathion	Pimephales promelas	BSA	-	12.5 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
CHEMIC	Malathion	Gambusia affinis	BSA	-	0.05 (K 40%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)
AL PRODUCTS	Malathíon	Pimephales promelas Lepomis macrochirus	BSA	_	17 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be ex- pected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Malathion	Lepomis macrochirus Micropterus salmoides Notemigonus crysoleucas Carassius auratus	BSA	_	0.1 (O) 0.1 (O) 0.1 (O) 0.1 (O)	<u>acdf</u>	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The inhibition of the enzyme was a function of the concentra- tion of the insecticide, extent of exposure, and specific chemical nature of the compound.	Weiss (1959)
lalathion (tech, 57% active in xylene)	Acroneuria pacifica Pteronarcys californica Claassenia sabulosa Arctopsyche grandis	BSA	-	0.0056 (T4A) 0.1 (T4A) 0.056 (T4A) 0.032 (T4A)	<u>a</u> c <u>e</u> fln	Assays were conducted in hard water.	Gaufin (1961)
falathion (57% concen- trate emulsified in xylene)	Acroneuria pacifica Hydropsyche californica Arctopsyche grandis Claassenia sabulosa Pteronarcys californica	BSA	_	0.0072 (T4A) 0.0225 (T4A) 0.032 (T4A) 0.056 (T4A) 0.1 (T4A)	<u>a</u> cde!m	Test water was obtained from a mountain stream.	Gaufin, et al (1961)
lalathion	Oncorhynchus tshawytscha Gasterosteus aculeatus	BSA	-	23 (T4A) 94 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961)
lalathion (81 percent EC)	Gambusia affinis	FL	Ponds— Bakers- field, Cal.	(0)	a c	At 0.5 lb/acre, 48 percent mortality occurred in 24 hours. At 2.0 lb/acre, 54 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
falathion (0,0-dimethyl dithiophosphate of diethyl mercapto- succinate)	Richardsonius balteatus hydroflox	BSA	_	13.6 (T1A) 11.4 (T2A) 8.9 (T4A)	ac de f	Results given were in soft water. Results in hard water were as follows: 11.7 (T1A), 9.6 (T2A), and 9.6 (T4A).	Webb (1961)

COMMERCIAL CHEMICAL PRODUCTS

**B-15**6

	Malathion	Killifish Cyprinodon Fundulus Gambusia	F	-	0.2-0.75 Ib/acre (O)	-	Extensive mortality.	Spiller (1961)
		Mollienesia Salmon (fingerlings) Carp Bluegill (fingerlings) Goldfish Rainbow trout Sunfish Yellow perch Fathead minnow	BSA	-	0.1 (K 100%) 3.0 (K 60%) 5.0 (K 100%) 25 (O)		Decidedly toxic. Killed 60% of test fish. Lethal dose. LD <sub>50</sub> in 24 hours.	
B-157	Malathion	Micropterus salmoides Pimephales promelas	BSA	_	0.5 (O) 0.5 (O)	-	The degree of reaction to the cholinesterase-inhibiting insecticides is not only a function of time and concentration, but also of chemical and biological species. This paper reports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as LT <sub>50</sub> which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the LT <sub>50</sub> was 2 hr and 40 min and for fatheads 72 hr.	Weiss (1961)
7	Malathion (granular)	Stizostedion vitreum	BSA		1.84 (O)	a e	Five percent of the fish survived 24 hours at the indi- cated concentration. Emulsions were more toxic than granular formulations of the chemical.	Hilsenhoff (1962)
	Malathion	Salmo gairdnerii (one wk old sac fry) (one mo old feeding fry)	BSA BSA	_	1.0 (K 26%) 10.0 (K 100%) 1.0 (K 100%) 10.0 (K 100%)	<u>a e</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
COMMER	Malathion, emulsible concentrate (20 percent)	Lepomis macrochirus Green sunfish Largemouth bass	BSA	_	Large 1.2 (T4A) Smail 0.55 (T4A) 0.60 (T4A) 0.25 (T4A)	<u>a</u> c d <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)
COMMERCIAL CHEMICAL PRODUCTS	Malathion, (tech, 100 percent)	Pimephales promelas Lepomis macrochirus Lebistes reticulatus	BSA	_	23 (T4A) 0.090 (T4A) 0.84 (T4A)	<u>a</u> cd <u>e</u>	Comment same as above.	Pickering, et al (1962)
L PRODUCTS	Malathion, emulsible concentrate (57 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus	BSA		25 (T4A) 0.088 (T4A) 0.79 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Malathron	Lepomis macrochirus	BSA		0.28 (T1A) 0.22 (T1A) 0.135 (T1A) 0.124 (T1A) 0.07 (T1A)	а	The experiment was conducted at 45 F The experiment was conducted at 55 F The experiment was conducted at 65 F. The experiment was conducted at 75 F. The experiment was conducted at 85 F.	Cope (1963)
Malathion	Culex pipiens quadrimaculatus	BSA	-	(0)	с	Tests were conducted in tap water and artificially polluted tap water. The values reported are the concentration range for an LCg0, 0.045 to 0.120 ppm in polluted water and 0.100 to 0.240 in tap water.	Lewallen and Wilder (1963)
Malathion (tech)	Salmo gairdneri Redear Lepomis macrochirus	BSA	-	100 (T1A) 170 (T1A) 100 (T2A) 60 (T4A) 45 (T1A) 35 (T2A) 120 (T1A)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long. The experiment was conducted at 75 F Fish weighed 3 g. The experiment was conducted at 75 F. Fish weighed 0.4 g. The experiment was conducted at 75 F Fish weighed 0.6 g.	Соре (1963)
Malathion	Daphnia magna	BSA	-	0.002 (T2A) 0.010 (K)	<u>a</u> ce	Acetone was used as a solvent for the Malathion. Each test solution contained 0.1% acetone.	Gillespie (1964)
Malathion	Gammarus Iacustris	BSA	_	0.00162 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Malathion (tech, 95 percent active In acetone)	Pteronarcys californica (naiad) Acroneuria p'acifica (naiad)	BSA	-	0.0500 (T4A) 0.0070 (T4A)	c d e f	A. pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than P. californica.	Jénsen and Gaufin (1964)
<b>Aalathion</b>	Bluegill	BSA	-	0.090 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Aalathion	Carassius auratus Lepomis macrochirus No ternigonus crysoleucus	BSCH	-	1.0 (O)* 1.0 (O)* 1.0 (O)* *response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
Malathion	Rainbow trout	BSA	-	77 (T4A) 68 (T4A) 110 (T4A)	<u>a</u>	These experiments were performed to show the effect of temperature on the toxicity. For the toxicant concentrations listed, the temperatures were respectively 45, 55, and 65 F. The fish all were approximately one g in weight. Toxicant concentrations for one and 2-day times were also listed.	Cope (1965)

and 2-day times were also listed.

	Malathion	Acroneuria	BSA	-	0.007 (T4A)	<u>a</u> c	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
		pacifica Ephemerella grandis			0.10 (T4A)			(1905)
		Gammarus			0.0016 (T4A)			
		lacustris Pteronarcys californica			0.05 (T4A)			
	Malathion	Pteronarcys californica	BSA	-	0.05 (T4A)	a	Unspecified chemical characteristics of assay water were determined by standard methods. General comments	Gaufin, et al (1965)
		Acroneuria pacifica			0.007 (T4A)		were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Addi-	
		Ephemerella grandis			0.100 (T4A)		tional data are presented.	
		Arctopsyche grandis			0.02 (T4A)			
		Hydropsyche californica			0.007 (T4A)			
		Daphnia magna			0.009 (T 50 hr A)			
		Gammarus Jacustris			0.002 (T4A)			
		Fathead minnow			12.5 (T4A)			
1	Malathion, 3,2,dibrom	Lepomis gibbosus	BSA	-	0.075 (T2A)	ac	This paper contained both bioassay and field studies. The tests revealed that a mixture of 3 parts of actual Dibrom	Hoff and Westman
	3,2,000000	Lepomis macrochirus			0.075 (T2A)		and 2 parts of actual Malathion (by weight) applied at 0.10 ppm was more toxic to bluegills and pumpkinseeds	(1965)
		Enneacan thus gloriosus			0.075 (T2A)		than to largemouth bass.	
		Esox niger			0.075 (T2A)			
		Enneacanthus chaetodon			0.075 (T2A)			
		Lepomis microlophus			0.08 (T2A)			
_		Salmo trutta			0.08 (T2A)			
ŝ		Salmo gairdnerii			0.08 (T2A)			
MER		Lepomis auritus			0.085 (T2A)			
		Perca flavescens			0.09 (T2A)			
CHE		Micropterus salmoides			0.10 (T2A)			
M N N		Fundulus diaphanus			0.10 (T2A)			
ALP		Catostomus commersoní			1.5 (T2A)			
ROD		Notemigonus crysoleucas			3.2 (T2A)			
COMMERCIAL CHEMICAL PRODUCTS		Erimyzon oblongus			4.2 (T2A)			

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL PRODUCTS	Malathion, 3,2,dibrom	Yellow perch Pumpkinseed Bluegill Golden shiner Brown bullhead Chain pickerel Largemouth bass Black crappie	FL	Silver Lake, Conn.	(O)	a C e	At 0.10 ppm sunfish kill was extremely heavy, the fish were still dying after 7 days. Many young of the year fish of all species were killed.	Hoff and Westman (1965)
DUCTS		Pumpkinseed Yellow perch White perch Largemouth bass Brown bullhead		Pooley Lake, Conn.	(0)		Kill of pumpkinseeds was slight. A few young of the year bass were killed at a concentration of 0.10 ppm.	
		Rainbow trout Pumpkinseed Bluegill Largemouth bass		A lake in N.J.	(O)		At a concentration of 0.10 the kill of bluegills was very heavy by the second day after treatment. Seining before treatment indicated that about 70 percent of the bluegills were killed, largemouth bass were still common.	
B-160		White perch Bluegill Largemouth bass Yellow perch Black crappie Channel catfish Gizzard shad Carp		Green- wich Lake, N.J.	(0)		The first application at 0.10 percent concentration appeared to kill about 80 percent of the white perch population and about 50 percent of bluegills.	
		Largemouth bass Black crappie Chain pickerel Bluegill Yellow perch Golden shiner Pumpkinseed Banded sunfish Brown bullhead		2 lakes in Mass.	(0)		The results were quite similar to those noted above at a concentration of 0.10. Largemouth bass were abundant and apparently minimally affected. Bioassay results are also presented.	
	Malathion, (95% active in acetone)	Hexagenia Hydropsyche (larva)	BSA	_	0.63 (T1A) 0.102 (T1A) 0.14 (T1A)	ae	Dissolved oxygen was measured before and after assay. Assays were conducted in Mississippi River water.	Carlson (1966)
		Bluegill	50.4				This paper reports on the truinity of a number of second state	0
	Malathion	Salmo gairdnerii Lepomis	BSA	_	0.079 (T2A) 0.086 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses subacute mortality as well. Effects on repro- duction and behavior are also discussed. Data presented as	Cope (1966)
		macrochirus Ictalurus			8.900 (T2A)		EC <sub>50</sub> .	
		punctatus Pteronarcys			0.020 (T2A)			
		californica Baetis sp			0.006 (T2A)			

Baetis sp

		Daphnia pulex Simocephalus			0.002 (T2A) 0.003 (T2A)			
Malathi		serrulatus Lepomis					This serves is a study of the amounts of organic this.	Murphy
Maiathi	ion	gibbosus Ictalurus		-	161.0 ± 19.5 (O) 11.6 ± 2.0 (O)	-	This paper is a study of the amounts of organic thio- phosphate and their oxygen analogues which accumulate in liver slices in an <i>in vitro</i> study of insecticides. The	(1966)
		melas Pseudopleuronectes			16.9 ± 3.8 (O)		numbers given are for m $\mu$ m of chemical (in the case of Parathion, Malathion, and Guthion – the oxygen analogue)	
		americanus Myxocephalus scorpius			6.1 ± 0.8 (O)		accumulated in 100 mg (dry weight) of liver in 30 minutes.	
Malath	ion	Lepomis gibbosus	BSA	_	2/4 (O)	_	The figures given are for mortality in 2 hours when the amount of chemical was 100.0 mg/kg, given by injection.	Murphy (1966)
Malath	ion	Sewage organisms	BOD	_	(O)	_	Shock loadings of the chemical as high as 100.0 mg/l were assimilated by microbial systems of 2000.0 mg/l with no observable toxic effect. Organophosphate insecti- cides in low concentration stimulated microbial respira- tion; however, greater concentrations inhibited the system and eventually destroyed the organisms. The amount of organophosphate insecticide required to inhibit the respiration of freshwater microorganisms was shown to be function of the amount of organisms present and not the volume of water in which the organisms are dispersed.	Randall (1966)
Malath	ion	Daphnia magna	BSA	-	0.0009 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
Malathi	ion	Daphnia carinata	BSA	_	0.0002 (SB)	_	Comment same as above.	Sanders and Cope (1966)
Malathi	ion	Simocephalus serrulatus Daphnia pulex	BSA	-	0.0035 (SB)	-	Comment same as above, except that time for immobili- zation was 48 hr and data were cited for 60 F.	Sanders and Cope (1966)
Malathi	ion	Tubifex spp Limnodrilus spp	BSA	-	16-7 (L4A)	ace	Toxicity is reported as the mean lethal dose (LD <sub>50</sub> ) for 24, 48, and 96 hours.	Whitten and Goodnight (1966)
Malathi	ion	Leiostomus xanthurus	BCFCH	_	0.01 (SB 182)	а	A concentration of 0.050 ppm killed juvenile spat in 14 days.	Butler and Johnson (1967)
Malathi	ion	Leiostomus xanthurus Cyprinodon variegatus	BCFCH	-	0.1 (O) 0.1 (O)	а	At a concentration of 0.1 ppm, the following percent acetylcholinesterase activity as compared to controls was found: <i>L. xanthurus</i> –76 <i>C. variegatus</i> –39	Butler and Johnson (1967)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Malathion	Stream insects: Ephemeroptera Odonata Plecoptera Trichoptera Diptera Coleoptera Neuroptera	FR	Maine	(O)		At an application rate of 1.2 pounds per acre, percent reduc- tion of the number of major insects is given with 0 to 66% reduction estimated. Population modifications were not severe and were considered transitory. Malathion is a safer chemical than DDT because it is not known to leave per- sistent residues. Thirty genera of aquatic insects were studied.	Dimond (1967)
Malathion	Mercenaria mercenaria American eel Mummichog Striped mullet Northern puffer Atlantic silverside Grass shrimp Sand shrimp Hermit crab	BSA	-	(O)	a c e	A 37,000 ppb exposure for 4 days caused no mortality of quahog clams. The LC <sub>50</sub> values (96 hour) for this chemi- cal extended from 8 to 3,250 ppb for fish, and from 33 to 83 ppb for crustaceans.	Eisler and Weinstein (1967)
Malathion	Atlantic salmon Brook trout	FR	New Bruns- wick	(0)	-	Spraying with this chemical at 1/8 lb per acre was no more harmful than the application of DDT at 1/4 lb per acre.	Kerswill and Edwards (1967)
Malathion	Pimephales promelas	BSA & CH	-	9.0 (T4A)	a c d e f	The fish could tolerate 1/45 this amount of malathion for a 10-month test.	Mount and Stephan (1967)
Malathion	Pimephales promelas	BCFCH	-	0.2-0.58 (O)	a c d e q	Carbon-filtered tap water was used as diluent. Malathion at indicated range of concentrations did not affect growth and reproduction, although 20% of fish died at the 0.58 ppm concentration during 7 weeks of exposure.	Mount and Stephan (1967)
Malathion	Puntius puckelli	BSA	-	3.7 (T4A)	<u>a</u> cdelm	Tap water was used as diluent. Toxicity data are given as TL <sub>m</sub> 's in ppm for 24, 48, 96 hr. The pH of the water averaged 8.3. The study was conducted in India.	Rao, et al (1967)
Malathion	Pteronarcys californica (naiads)	BSA	_	0.01 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Malathion	Pteronarcys californica (naiads) Pteronarcella badia (naiads)	BSA	-	0.010 (T4A) 0.0011 (T4A)	<u>acdef</u>	Comment same as above.	Sanders and Cope (1968)
	(nalads) Claasenia sabulosa (nalads)			0.0028 (T4A)			

Malathion	Trout Red shiner	BSA	-	0.0050 (T1A) 0.0046 (T2A) 0.0028 (T4A) 0.0023 (T5A) 0.040 (T1A) 0.036 (T2A) 0.025 (T4A) 0.023 (T5A)	a c d e f i	The objects of this investigation were the recovery of organic micropollutants from subsurface and surface Missouri waters, characterization and identification of these substances, and evaluation of their toxic effects, both acute and long-term, in order to develop methods for their destruction or removal.	Smith and Grigoropoulos (1968)	
Malathion	Lepomis gibbosus Ictalurus melas Micropterus dolomieui Myxocephalus scorpius Pseudopleuronectes americanus	BSA	-	(0)	ар	The chemicals were poor inhibitors of brain cholinesterases <i>in vitro</i> ; their oxygen analogs were potent inhibitors.	Murphy, et al (1968)	
Manzate	Channel catfish (fingerlings)	BSA	_	2.7 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	
2-mercaptobenzo- thiazole	Daphnia magna	BSA	-	2 (K2)	а	An attempt was made to correlate the biological action with the chemical reactivity of selected chemical substances. Results indicated a considerable correla- tion between the aquarium fish toxicity and anti- autocatalytic potency of the chemicals in marked con- trast to their toxicity on systemic administration.	Sollman (1949)	APPENDIX
Merthiolate	Salmo	BSA	-	21.2 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford	ω
	gairdnerii Salmo			54.0 (T2A)		were also determined.	(1966)	
	trutta Salvelinus			74.5 (T2A)				
	fontinalis Salvelinus			2.13 (T2A)				
	namaycush Ictalurus			5.65 (T2A)				
	punctatus Lepomis macrochirus			64.5 (T2A)				
Metacide (dialkyl nitroaryl thio- phosphate, 33.4 percent)	Lymnaeid snails	BSA	_	(0)	-	Each test container, 500-ml beaker, was filled with ditch water. 100% mortality occurred at 1:300,000 and greater.	Batte, et al (1951)	
Meta-Systox R	Oyster	BCF	-	(0)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)	
Metasystox	Cyprinus carpio C. carpio Tilapia mossambica Cirrhina mrigala Labeo fimbriatus	BSA	-	9.0 (T2A) 20.0-25.0 (T2A) 12.0-12.5 (T2A) 17.0 (T2A) 16.0 (T2A)	ac de f p	The test animals were conditioned for 48 hours prior to use.	Sreenivasan and Swaminathan (1967)	

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Methoxychlor	Fathead minnow	BSA	_	0.035 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish" It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959)
Methoxychlor	Fathead minnow Bluegill Goldfish Guppy	BSA	_	0.064 (T4A) 0.062 (T4A) 0.056 (T4A) 0.120 (T4A)	a	Comment same as above except that experiment was con- ducted in soft water.	Tarzwell (1959)
Methoxychlor	Daphnia magna	BSA	-	0.0036 (0)	a	The indicated concentration immobilized Daphnia in 50 hours.	Anderson (1960)
Methoxychlor	Pimephales promelas	BSA	-	0.06 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
	Lepomis macrochirus Carassius auratus Lebistes			0.06 (T4A) 0.06 (T4A) 0.12 (T4A)			
Methoxychlor	reticulatus Pimephales	BSA	_	0.064 (T4A)	a b e c d f	Dilution water was usually soft although some studies were	Henderson, et al
(100%)	promelas Lepomis	2011		0.062 (T4A)	<u> </u>	conducted with hard water.	(1959)
	macrochirus Carassius			0.056 (T4A)			
	auratus Lebistes reticulatus			0.120 (T4A)			
Methoxychlor	Pimephales	BSA	-	0.064 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as	Tarzwell
	promelas Lepomis macrochirus			0.062 (T4A)		toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	(1959)
Methoxychlor	Oncorhynchus kisutch	BSA	-	66.2 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz
	Oncorhynchus tshawytscha			27.9 (T4A) (196	(1961)		
	Salmo gairdnerii			62.6 (T4A)			
	Gasterosteus aculeatus			86.4 (T4A)			

	Methoxychlor (EC 2)	Gambusia affinis	FL	Cal.	2.0 (O)	ac	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in lb/acre, and re- sulted in a 6 percent mortality for these fish.	Mulla (1963)
	Methoxychlor (tech)	Salmo gairdnerii	BSA		20 (T1A)	а	The experiment was conducted at 55 F. Fish weighed 0.7 g.	Cope (1963)
	Methoxychlor	Gambusia affinis	BSA	_	0.6 (L1)* 0.9 (L1)** * Resistant fish **Nonresistant fish	а	This paper deals with the resistance of mosquito fish to chlorinated hydrocarbon compounds. Resistant fish were not always less sensitive to these chemicals.	Boyd and Ferguson (1964)
	Methoxychlor	Bluegill	BSA	-	0.062 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
	Methoxychlor	Pteronarcys sp (nymphs)	BSA	-	0.0014 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
5	Methoxychlor	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.04 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 33 percent.	Jamnback and Frempong- Boadu (1966)
100	Methoxychlor	Daphnia magna	BSA	_	0.0037 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Methoxychlor	Simocephalus serrulatus Daphnia pulex	BSA	-	0.005 (SB) 0.00078 (SB)	_	Comment same as above except that time for immobiliza- tion was 48 hr and data were cited for 60 F.	Sanders and Cope (1966)
COMMERCIAL CHE	Methoxychlor	Mya arenaria Crassostrea virginica Corbicula manillensis Mercenaria mercenaria Rangia cuneata	BCFCH	-	(O)	-	Results are recorded as a range of uptake of the chemical by 5 species of aquatic mollusks. An uptake or concentra- tion of 300-1500 ppb resulted.	Butler (1967)

5

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Methoxychlor	Mercenaria mercenaria American eel Mummichog Striped mullet Northern puffer Atlantic silverside Grass shrimp Sand shrimp Hermit crab	BSA	-	(0)	a c e	At 1,100 ppb exposure for 4 days caused no mortality of the quahog clam. Although this organism was quite resis- tant to this chemical, other organisms were susceptible. A 4 day TL <sub>m</sub> of 12 to 150 ppb was found for such fish as the American eel, mummichog, striped mullet, northern puffer, Atlantic silverside; and between 4 and 12 ppb for crustaceans (including grass shrimp, sand shrimp, and hermit crab).	Eisler and Weinstein (1967)
Methoxychior	Lampsilis siliguoidea L. vertricosa Anodonta grandis	FR	Red Cedar River, Mich.	(O)	-	The mussels listed were analyzed for the toxicant and its metabolites. Mussels may be used as detectors for this toxicant because they tend to concentrate the chemical in much higher concentrations than it is ever found in the water. The amount of chemical applied as a spray was not specified.	Bedford, et al (1968)
Methoxychlor	Pteronarcys californica (naiads)	BSA	-	0.0014 (T4A)	<u>a c d e f</u>	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Methylene blue	Microcystis aeruginosa	L	_	100 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Methylene blue	Micropetrus salmoides (fry) Ictalurus punctatus (fry) Lepomis mæcrochirus (fry)	BSA	_	5.0 (SB3) 5.0 (SB3) 5.0 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
Methylene blue	Salmo gairdneri Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	-	10.0 (T2A) 32.8 (T2A) 22.9 (T2A) 34.0 (T2A) 104 (T2A) 33.0 (T2A)	<u>ə</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)

Methyl carbo- phenothion	Chaoborus astic topus	BSA	_	0.0064 (T1A)	<u>a</u>	Toxicity value given is for the fourth instar.	Hazeltine (1963)	
Methyl green	Microcystis aeruginosa	L	-	100 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, 1 x 10 <sup>6</sup> to 2 x 10 <sup>6</sup> cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)	
Methyl parathion	Pimephales promelas	BSA	-	10.4 (T4A)	<u>acdef</u>	Tests were performed in both hard and soft water. Addi- tional tolerance limit values are given.	Henderson and Pickering (1958)	
Methyl parathion	Fathead minnow	BSA	_	7.5 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959)	
Methyl parathion	Pimephales promelas	BSA	-	8.3 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1960)	
Methyl parathion	Pimephales promelas	BSA	-	8.3 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)	APPENDIX B
Methyl parathion	Pimephales promelas	BSA	-	9.5 (T4A) 2.4 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)	~
(tech, 80 percent)	Lepomis macrochirus	ochirus						
	Carassius auratus			12.0 (T4A)				
	Lebistes reticulatus			9.8 (T4A)				
Methyl parathion (tech)	Lepomis macrochirus	BSA		8,500 (T1A)	а	The experiment was conducted at 75 F. Fish weighed 0.6 g.	Cope (1963)	
Methyl parathion	Chao borus astic topus	FL	Clear Lake, othe ponds & lakes, Cal.	(O) rr	-	Methyl parathion applied to 2.3 ppb and 3.3 ppb at intervals within 2 months was sufficient to control gnats in clear lake.	Hazeltine (1963)	
Methyl parathion	Chaoborus astictopus	BSA	-	(0)	_	Tests were conducted on bluegill, sunfish, <i>C. astictopus</i> first instar larvae, and fourth instar larvae, results on	Hazeltine (1963)	
	Lepomis macrochirus			0.115 (T10A)	а	larvae were as follows: Fourth instar 0.0058 (T1A) First instar 0.0012 (T1A)		

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen ts	Reference (Year)
Methyl parathion (EC7.5)	Gambusia affinis Bufo boreas Scophiopus hammondi	FL	Ponds in III.	(O) (O) (O)	_	When applied at 0.8 pound per acre active ingredient, 10 percent fish mortality occurred in 1 day. No toad mortality occurred at 0.4 pound per acre in 1 day.	Mulla, et al (1963)
Methyl parathion (tech grade)	Procambarus clarki	BSA	-	0.04 (T3A)	acdo	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
Methyl parathion	Bluegill	BSA	-	1.9 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Methyl parathion	Carassius auratus Lepomis macrochirus Notemigonus crysoleucus	BSCH	_	10 (0)* 1.0 (0)* 10.0 (0)** 1.0? (0)** 1.0 (R-30 da) 1.0 (0)** * no response, 15 days **response, 15 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or not response in 15 or 30 days.	Weiss and Gakstatter (1964)
Methyl parathion	Phy toplank ton Zooplank ton Chironomids Oligochaetes Fish	FL	Clear Lake, Cal.	-	а	The purpose of this field study was to determine the effect of methyl parathion at 3 ppb or .003 ppm (used to control gnats <i>Chaoborus astictopus</i> ) on other organisms in the treated area. It appears that the treatments of Clear Lake had minimal influence upon the biota of the lake with the exception of the Clear Lake gnat larvae and to a lesser degree, perhaps, species of the zooplankton. This is a very general paper and there are no numerical data given.	Cook and Conners (1963)
Methyl parathion	<i>Procambarus clarkii</i> (juvenile)	BSA	-	<1.0 (T5A)	acdo	The pesticides studied in this report are widely used in rice culture in Louisiana and are toxic to crawfish.	Hendrick and Everett (1965)
Methyl parathion	G <b>amb</b> usia affinis	BSA	_	(0)	<u>a</u>	The effect of combinations of pesticides was studied. In general, the results reflected the extreme levels of Endrin and Toxaphene resistance in the resistant population. The results failed to indicate additive effects wherein the combi- nation mortality exceeded the sum of the mortalities pro- duced by the individual insecticides.	Ferguson and Bingham (1966)
Methyl parathion	Procambarus clarkii	FO	Crowley, La.	(0)	сdер	Experiments were conducted in a flooded rice field. Area was divided into 4 blocks with a fence, restricting craw- fish to desired areas. The rearing of crawfish in rice fields is of considerable commercial importance in Louisiana. No untoward effect on the crawfish occurred. The chemical was applied at the rate of 25 lb/acre.	Hendrick, et al (1966)

Methyl	Puntius	BSA	_	2.1 (T4A)	acdelm	Tap water was used as diluent. Toxicity data are given as	Rao, et al
parathion	puckelli			2.1 (176)	acterm	TL <sub>m</sub> 's in ppm for 24, 48, 96 hr. The pH of the water averaged 8.3. The study was conducted in India.	(1967)
Methyl trithion (50 percent EC)	Gambusia affinis	FL	Ponds— Bakers- field, Cal.	(0)	ас	At 0.5 lb/acre, 14 percent mortality occurred in 24 hours. At 2.0 lb/acre, 76 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
Methyl zimate	Pimephales promelas	BSA		0.25 (T4A)	ac de f	The toxicity of this substance was influenced by the quality of the water (pH, hardness, alkalinity). The chemical was more toxic in soft water.	Pickering and Henderson (196 <b>6</b> )
MGK's Evergreen	Salmo gairdneri	BSA	-	800 (T1A)	а	The experiment was conducted at 55 F. Fish weighed 0.4 g.	Cope (1963)
MGK's 6103	Salmo gairdneri	BSA	-	150 (T1A)	а	The experiment was conducted at 55 F. Fish weighed 0.5 g.	Cope (1963)
MGK's 6243	Salmo gairdneri	BSA	-	750 (T1A)	а	The experiment was conducted at 55 F. Fish weighed 0.8 g.	Cope (1963)
Ortho-MH 30	Channel catfish (fingerlings)	BSA	_	>2.4 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Mirex	Lepomis macrochirus Carassius auratus Salmo clarki	FLCH	-	5.0 (SB) 0.1-1.0 (SB 224) 75.0 (SB 14)	-	No histological lesions or effects on hematocrit or serum protein were observed in the bluegill. Lesions on and fusion of gills occurred with the trout at the indicated concentration. Gill changes and accumulation in gold- fish were found in concentrations of 2.0-1372 ppm in skin, muscle, liver, and gut.	Van Valin, et al (1968)
Mobam (r-benzo- thienyl-N-methyl carbamate)	Australorbis glabratus	BSA and FL	Puerto Rico	(0)	c	Seven of the tested compounds failed to meet accepta- bility criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
Molinate (Ordram)	Daphnia magna Rainbow trout Bluegill	BSA		0.70 (.46- 1.05 (O) 0.29 (O) 0.48 (O)	ac diq	Toxicity, in terms of median immobilization concentra- tion (IC <sub>50</sub> ), is presented for <i>Daphnia</i> ; median lethal concentration (LC <sub>50</sub> ) values for rainbow trout and bluegill are reported.	Crosby and Tucker (1966)
Molinate (tech)	Pteronarcys californica (naiads)	BSA	_	.00034 (T4A)	<u>acdef</u>	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Molucid (isobutyl- triphenyl- methylamine	Bulinus truncatus Biomorpholaria alexandrina Lymnaea caillaudi	FO	Arabia	1.6-2.0 (O)	a b g	Tests were conducted in the Hod el Malaha canal which has a maximum discharge of 11,250 m <sup>3</sup> /day. The mollus- cicide was dispersed by the injection method, with flow regulated by a tap, a concentration of 2 ppm being main- tained during 6 hr of continuous application. The Meyling, Schutte & Pitchford method was used for determining the concentration of molluscicide in the canal. No live organisms were observed for 2, 3, and 4 months after treatment. Egg masses were apparently unaffected.	Dawood and Dazo (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Monuron	Oncorhynchus kisutch	BSA and FL	_	115 (T1) 110 (T2)	a c d e	Concentrations were based on percent active ingredient. Treatment of ponds with 5 and 10 ppm Monuron apparently caused no mortality of frogs, tadpoles, or fishes.	Bond, et al (1960)
Monuron	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	L	-	0.02 (NG) 0.02 (NG) 0.02 (NG) 0.02 (NG) 0.02 (K)	<u>a</u> _	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
Monuron	Crassostrea virginica Penaeus setiferus Mugil cephalus Phytoplankton	BCFA & BSA	. –	2.0 (0, 12%) 0.55 (O) 16.3 (T2A) 94% (O)	_	Seawater was pumped continuously into test aquaria. Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following: Oyster - 96-hr EC50 - Conc. which decreased shell growth. Shrimp - 48-hr EC50 - Conc. which killed or paralyzed 50% of test animals. Fish - 48-hr EC50 - Conc. which killed 50%. Phytoplankton - Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	Butler (1965)
Monuron (25 percent pellet)	Lepomis macrochirus Lepomis macrochirus	BSA	-	40.0 (T4A) 47.0 (T4A)	-	Laboratory bioassays indicated that toxicity of the different formulations evaluated in this varied greatly with the fish used. Mortality data are expressed as $EC_{10}$ , $EC_{50}$ , etc.	Walker (1965)
Monuron (80 percent WP)	lctalurus nebulosis Lepomis macrochirus	BSA	-	57.0 (T4A) 33.0 (T4A)	-	Comment same as above.	Walker (1965)
Monuron	Daphnia magna	BSA	_	106 (O)	acdiq	Toxicity, in terms of median immobilization concentra- tion (IC <sub>50</sub> ), is presented.	Crosby and Tucker (1966)
Monuron (CMV)	Salmon	BSA	-	110.3 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
Monuron-borate (4-percent granular)	Lepomis macrochirus	BSA	-	26.0 (T4A)	-	Laboratory bioassays indicated that toxicity of the different formulations varied greatly with the fish used. Mortality data are expressed as $EC_{10}$ , $EC_{50}$ , etc.	Walker (1965)
Monuron-TCA (3 Ib/gal)	Lepomis macrochirus Micropterus salmoides	BSA	-	1.5-1.8 (T4A) 2.7 (T4A)	-	Comment same as above.	Walker (1965)

Monuron-TCA (22 percent granular)	Lepomis macrochirus	BSA	-	4.8 (T4A)	-	Comment same as above,	Walker (1965)
Monuron-TCA (11 percent granular)	Lepomis macrochirus	BSA	_	3.8 (T4A)	-	Comment same as above.	Walker (1965)
Monuron-TCA (tech)	Lepomis macrochirus Lepomis microlophus Micropterus salmoides Pumpkinseed	BSA	_	4.5 to 5.0 (T4A) 5.4 (T4A) 4.8 (T4A) 3.3 (T4A)	-	Comment same as above.	Wai ker (1965)
Monuron TCA	Lepomis macrochirus (eggs) L. cyanellus (eggs) Microp terus dolomieui (eggs) Erimyzon sucetta (eggs) L. macrochirus (fry)	L	-	10 (NTE) 10/5 (O) 10/4 (O) 10 (NTE) 20 (S)		Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	Hiltibran (1967)
MS-222	Salmo gairdnerii Salmo trutta Salvelinus fontinalis Salvelinus namaycush Esox lucius Lepomis macrochirus Micropterus salmoides Stizostedion vitreum	BSA	_	39.0 (T1A) 39.0 (T2A) 38.5 (T1A) 37.5 (T2A) 50.7 (T1A) 50.0 (T2A) 33.8 (T1A) 33.0 (T2A) 56.0 (T1A) 52.0 (T2A) 45.7 (T1A) 45.7 (T2A) 42.0 (T1A) 42.0 (T1A) 49.0 (T1A) 48.5 (T2A)	<u>a</u> ef	Large specimens of given species were usually more resis- tant to MS-222 than small ones. Trout were more tolerant at lower temperatures. A safety index of concentration is suggested.	Marking (1967)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
MS-222	lctalurus punctatus	BSA	_	12C 58.0 (T1A) 55.0 (T2A) 51.1 (T4A) 17C 60.5 (T1A) 60.0 (T2A) 60.0 (T4A) 22C 59.8 (T1A) 58.8 (T2A) 58.8 (T4A)	<u>a</u> cdef	Anesthesia was induced within 2 minutes by concentra- tions of this chemical of 20 to 40 ppm. Concentrations of 20 to 40 ppm maintained sedation for 6 hours. Safety indices were determined for the anesthesia of channel catfish with MS-222. The toxicity of the chemical is greatly influenced by the size of the fish and also by temperature. TL <sub>m</sub> 's are recorded at 12, 17, and 22 degrees centigrade. It was found that catfish are relatively more resistant when the anesthetic is dissolved in soft water.	Schoettger, et al (1967)
Tricaine methane- sul fonate (MS-222)	Salmo gairdnerii Salmo trutta Salvelinus fontinalis Salvelinus namaycush	BSA	_	100 (SB) 80 (SB) 120 (SB) 135 (SB)	<u>a</u> cf	In this assay the chemical was tested for its efficacy as an anesthetic for the given fish at varied temperatures. Con- centrations of 80-135 ppm of the chemical anesthetized all of these fish within 3 min at 7 to 17 C. 50 to 60 ppm induced a moderate rate of anesthesia which could be maintained for approximately 30 minutes. At 15 to 30 ppm, sedation was produced within 15 minutes and maintained for 5 to 6 hours. Lake trout required larger doses than the other salmonids for complete anesthesia. There was no relation between size of fish and efficacy of MS-222.	Schoettger and Julin (1967)
MS-222 (tricaine methane- sulfonate)	Rainbow trout	BSA	-	(0)	-	MS-222 can be detected in fish tissues by a modified Bratton-Marshall method. Interfering substances were more prevalent in liver and kidney than in blood and muscle. The recovery of spiked samples ranged from 89 to 112 percent. The method was more accurate for measuring MS-222 in blood and muscle than in kidney and liver.	Walker and Schoettger (1967)
MS-222 {tricaine methane- sulfonate}	Rainbow trout Brown trout Brook trout Lake trout	BSA	_	(0)	a f	The residues of MS-222 in selected tissues of fish at 7, 12, and 17 C and in waters of various hardnesses were measured by a modified Bratton-Marshall colorimetric method. The concentrations of drug in the blood, muscle, liver, and kidney of deeply anesthetized rainbow trout dissipated rapidly within 1 to 6 hours. The mean concentrations were 18 to 42 ppm in rainbow trout, 13 to 44 ppm in brown trout, 15 to 28 ppm in brook trout, and 15 to 32 ppm in lake trout.	Walker and Schoettger (1967)
M.S. 222 (tricaine methane- sulfonate)	Salmo gairdneri	BCFA	_	(0)	а	Fish anesthetized in 100 mg/l of M.S. 222 at 12 C excreted the drug in free and acetylated forms via the urine during a 24-hr recovery period in fresh water. Of the M.S. 222 ex- creted, 77-96% was acetylated. Blood and urine were cleared of the two fractions of M.S. 222 in 8 and 24 hr, respectively. Intraperitoneal injections of 10-100 mg/kg of M.S. 222 did not induce anesthesia; however, the 24-hr pattern of drug excretion was similar to that observed after anesthesia by immersion.	Hunn, et al (1968)

	N-2404 EC4	<i>Gambusia affinis</i> Diving	FL	Cal.	0.1 (K1) (O)	-	Toxicity value is in Ib/acre. At the given rates, there was appreciable kill of diving beetle larvae and adults, chirona- mid larvae, and dragonfly naiads.	Mulla (1966)
		beetle (larvae) Chironomid (larvae) Dragonfly (naiads)			(O) (O)			
	N-2788 EC4	Gambusia affinis	FL	Cal.	0.1 (K1)	-	Toxicity value is in Ib/acre.	Mulla (1966)
	N-2790 EC4	Gambusia affinis	FL	Cal.	0.2 (K1)	-	Comment same as above.	Mulla (1966)
	N-2790 EC4	Gambusia affinis Rana catesbeiana	FL	Cal.	1.0 (K1) (O)	-	Toxicity value is in lb/acre. No mortality in tadpoles of <i>R. catesbeiana</i> occurred during an exposure period of one week.	Mulla (1966)
<b>1</b>	Nabam	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum trocornutum Monochrysis lutheri	BSA	-	10.0 (K) 10.0 (K) 1.0 (K) 1.0 (NG) 10.0 (K) 1.0 (K)	a	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
3	Naled (EC8)	Gambusia affinis Rana catesbeiana	FL	Ponds in III.	(0)	_	When applied at 2.0 pounds per acre active ingredient, 20 percent fish mortality occurred in 1 day. No bullfrog mortality occurred at 0.5 pound per acre in 1 day.	Mulla (1963)
	Naled	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	_	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 22 percent.	Jamnback and Frempong- Boadu (1966)
SOMME	Naled	Pteronarcys californica (naiads)	BSA	_	0.008 (T4A)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
ERCIAL CHEMICAL PRODUC	Neburon	Protococcus sp Chlorella sp Duraliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	-	0.20 (NG) 0.20 (K) 0.20 (NG) 0.20 (NG) 0.004 (K)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
PRODUC	Neburon	Leiostomus xanthurus (juvenile)	BSA	-	0.032 (0)	а	Water temperature was 21 C. The figure reported is a 48-hr $EC_{50}$ .	Butler (1965)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Neburon	Crassostrea virginica Penaeus setiferus Leiostomus xanthurus Phytoplankton	BCFA & BSA		0.41 (O) 0.55 (O) 0.22 (T2CFA) 90% (O)	_	Seawater was pumped continuously into test aquaria. Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following: Oyster – 96-hr EC50 – Conc. which decreased shell growth. Shrimp – 48-hr EC50 – Conc. which killed or paralyzed 50% of test animals. Fish – 48-hr EC50 – Conc. which killed 50%. Phytoplankton – Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	Butler (1965)
Neburon, 4 percent granular	Pimephales notatus Notropis umbratilis Lepomis macrochirus Lepomis microlophus	BSA	-	0.6 (T4A) 0.9 (T4A) 0.7 (T4A) 0.8 (T4A)	_	Laboratory bioassays indicated that toxicity of the different formulations evaluated in this varied greatly with the fish used. Mortality data are expressed as $EC_{10}$ , $EC_{50}$ , etc.	Walker (1965)
Nectran	Salmo gairdneri	BSA	-	7,000 (T 18 hr)	а	<ul> <li>The experiment was conducted at 55 F. Fish were 2-3 in. long.</li> </ul>	Cope (1963)
Neguvon	Salmo gairdnerii Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	-	12.2 (T2A) 16.5 (T2A) 16.8 (T2A) 9.0 (T2A) 32.0 (T2A) 71.0 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
Nemagon	Lepomis macrochirus Micropterus salmoides	BSA	-	20 (T2A) 20 (T2A)	<u>a</u> co	The response of bluegill and bass fingerlings to nine agri- cultural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
Neotran	Channel catfish (fingerlings)	BSA	-	146 (K1A)	<u>a</u>	Tap water was used. Considerable addtional data are presented.	Clemens and Sneed (1959)
Nigrosine	Daphnia magna	L	-	(0)	a	Aquaria were treated with 1.2, 2.4, 3.6, and 7.2 ppm nigrosine, corresponding to 10, 20, 30, and 60 pounds per acre. It was shown that <i>Daphnia</i> could survive the 7.2 ppm concentration for at least 5 days and probably for much longer.	Surber (1943)

	N-Serve	Oyster	BCF		0.28 (O)	a	The value reported is a 96-hr	EC50 (decreased shell	Butler (1965)
		Penaeus aztecus	L	-	(0)	8	tures averaging about 28 C.	luated in seawater at tempera- . The values are for 24-hr EC <sub>50</sub> equilibrium or mortelity. No	(1965)
		<i>Leiostomus xanthurus</i> (juvenile)	BSA	_	(0)	a	Water temperature was 16 C exposure to 1.0 ppm.		
	N-Serve	Crassostrea virginica Penaeus aztecus	BCFA & BSA		0.28 (O) 1.0 (NTE)	-	and ambient weather cond	nuously into test aquaria. plankton fluctuated with tide, itions. Some bioassays with was reported for the following:	Butler (1965)
		Leiostomus xanthurus			1.0 (NTE)			EC <sub>50</sub> — Conc. which decreased rowth.	
		Phytoplankton			15% (O)		Shrimp — 48-hr paraly Fish — 48-hr 50%.	EC <sub>50</sub> — Conc. which killed or zed 50% of test animals. EC <sub>50</sub> — Conc. which killed It decrease of CO2 fixation to a	
							4-hr ex	cosure at 1.0 ppm chemical httation.	
B-175	Nonic 218	Semotilus atromaculatus	BSA	-	20 to 60 (CR)	<u>a</u> e	typical water analysis is given the "critical range" (CR), we concentration in ppm belo	aerated Detroit River water. A ren. Toxicity is expressed as which was defined as that w which the 4 test fish lived for test fish died. Additional data	Gillette, et al (1952)
	Noxfish (5.0% rote- none, 10.0% cube extracts emulsifier)	Cyprinus carpio Micropterus salmoides Pimephales promelas	BSA		0.081 (T3A) 0.147 (T3A) 0.159 (T3A)	<u>a</u> c d e i	studied. Toxicity is express concentrations of rotenone conjunction with sulfoxide	re, species, and size of fish were sed as $LD_{50}$ for 72 hr. Smaller were required when used in The data shown are for 70 F. ably more toxic at this tempera- h species.	Hester (1959)
•		Carassius auratus Lepomis macrochirus			0.175 (T3A) 0.179 (T3A)				
COMMER		L. cyanellus Notemigonus crysoleucas lctalurus			0.165 (T3A) 0.470 (T3A) 0.247 (T3A)				
CIAL		nebulosus marmoratus							
COMMERCIAL CHEMICAL PRODUCTS	Noxfish (5% rotenone)	Cyprinus carpio (eggs) (fry) Pimephales promelas (eggs)	BSA	_	0.091 (O) 0.081 (O) 0.142 (L)	<u>a</u>	Toxicity is reported as $LD_{50}$	) in ppm, at 75 F.	Hester (1959)
OUCTS		(fry)			0.159 (L)				

COMMERCIAL CHEMICAL PRODUCTS	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)
AL CHE	Nytron	Penaeus az tecus	L	_
MICAL	Oil, crude	Daphnia magna	BSA	_
PRODU	Omazene	Onchorynchus tshawytscha	BSA	_
JCTS	OMPA (70%)	Pimephales promelas	BSA	_
	OMPA	Fathead minnow	BSA	-
H	ΟΜΡΑ	Pimephales	BSA	_

(larvae)

						different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.
ΟΜΡΑ	Pimephales promelas	BSA	-	121 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.
ΟΜΡΑ	Pimephales promelas	BSA	-	121 (T4A)	<u>a</u>	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.
OMPA (tech, 90 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	₽SA	-	88 (T4A) 120 (T4A) 680 (T4A) 22 (T4A)	<u>a</u> c d <u>e</u>	Soft water primarily was the test medium. TLm's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.
OMS-3 (EC2)	Gambusia affinis	FL	Ponds in III.	(0)	~~	When applied at 1.0 pound per acre active ingredient, 100 percent mortality occurred in 1 day.
OMS 44 (Bayer 37343)	Prosimulum spp Cnephia spp Simulium spp	LCFA	-	0.4 (O)	а	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated

Toxicity.

Active

Ingredient.

ppm(3)

0.0015 (O)

10.000 (T1A)

4.613 (T2A) 752 (T3A)

0.83 (T1A)

0.83 (T2A)

135 (T4A)

135 (T4A)

Experimental

Variables

Controlled

or Noted(4)

а

е

acde

acdef

а

Reference

(Year)

Butler

(1965)

Dowden

(1962)

Bond, et al.

Pickering

(1958) Tarzwell

(1959)

Henderson and

Henderson, et al

Pickering, et al

(1962)

(1960)Tarzweil

(1959)

(1960)

or carrier in most cases.	(1002)
When applied at 1.0 pound per acre active ingredient, 100 percent mortality occurred in 1 day.	Mulla, et al (1963)
Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 89 percent.	Jamnback and Frempong- Boadu (1966)

Comments

or enough to cause loss of equilibrium or mortality. This study is concerned with waste oil emulsifiers.

Concentrations were based on percent active ingredient.

Tests were performed in both hard and soft water. Addi-

It was the authors opinion that pH, alkalinity and hardness,

within the usual range in natural waters, had little effect

on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish". It is interesting that the

tional tolerance limit values are given.

Toxicant chemicals were evaluated in sea water at temperatures averaging about 28 C. The values are for 24-hr ECEn

	OMS-115 (EC2)	Gambusia affinis	FL	Ponds in III.	(0)	-	When applied at 0.5 pound per acre active ingredient, 100 percent mortality occurred in 1 day.	Mulla, et al (1963)
	OMS-144	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	a	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away, they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 8 percent.	Jamnback and Frempong- Boadu (1966)
	OMS-315	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	4.0 (O)	а	Comment same as above except that at that time the number detached amounted to 35 percent.	Jamnback and Frempong- Boadu (1966)
	OMS-437	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	а	Comment same as above except that at that time the number detached amounted to 61 percent.	Jamnback and Frempong- Boadu (1966)
	OMS-595 (SD8447)	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (0)	а	Comment same as above except that at that time the number detached amounted to 9 percent.	Jamnback and Frempong- Boadu (1966)
B-177	OMS-648	<i>Prosimulum</i> spp <i>Cnephia</i> spp <i>Simulium</i> spp (larvae)	LCFA	_	0.4 (O)	a	Comment same as above except that at that time the number detached amounted to 38 percent.	Jamnback and Frempong- Boadu (1966)
	OMS-658	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	4.0 (O)	а	Comment same as above except that at that time the number detached amounted to 35 percent.	Jamnback and Frempong- Boadu (1966)
	OMS-659	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	0.4 (O)	а	Comment same as above except that at that time the number detached amounted to 44 percent.	Jamnback and Frempong- Boadu (1966)
8	OMS-711	<i>Prosimulum</i> spp <i>Cnephia</i> spp <i>Simulium</i> spp (Iarvae)	LCFA	_	0.4 (O)	а	Comment same as above except that at that time the number detached amounted to 10 percent.	Jamnback and Frempong- Boadu (1966)
MERCIAL	OMS-712	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA		0.4 (O)	а	Comment same as above except that at that time the number detached amounted to 9 percent.	Jamnback and Frempong- Boadu (1966)
- CHEMIC	OMS-754	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	4.0 (O)	а	Comment same as above except that at that time the number detached amounted to 15 percent.	Jamnback and Frempong- Boadu (1966)
COMMERCIAL CHEMICAL PRODUCTS	OMS-868	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	_	0.04 (O)	а	Comment same as above except that at that time the number detached amounted to 11 percent.	Jamnback and Frempong- Boadu (1966)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Commen t <del>s</del>	Reference (Year)
OMS-869	Prosimulum spp Cnephia spp Simulium spp (larvae)	LCFA	-	<b>4.0</b> (O)	a	Stones heavily populated with wild larvae were placed in troughs of running water containing the toxicant. When the larvae became detached from the rocks and floated away they were assumed to have undergone lethal intoxi- cation. The larvae were exposed to the toxicant for 5 minutes, then in clean water for 24 hours. At that time the number detached amounted to 13 percent.	Jamnback and Frempong- Boadu (1966)
Organochlorines (pp'-DDT, pp'- DDE, HEOD, Endrin)	Ammodytes lanceolatus Phalacrocorax aristotelis Phalacrocorax carbo Gadus morrhua Mytilus edulis Somateria mollissima Cardium edule Pastella vulgata Homarus vulgata Homarus vulgaris Calcinus maenas Cancer poguras Pleuronectes sp Clupea harengus Gadus merlangus Sula bassana Halichoerus grypus Delphinus	FO	Britain	(0)		Residues of organochlorine insecticides tended to be greater in marine organisms of the higher trophic levels, but the tendency was not found in all food chains.	Robinson, et al (1967)
Organo- phosphorus pesticides	delphis Leiostomus xanthurus Cyprinodon variegatus	FECH	Atlantic and Gulf Coasts	(0)	-	Describes method to detect low level concentration of pollu- tion by measuring the degree of inhibition of acetyl cholin- esterase (AChE). Of 93 samples from 43 stations, 17 showed less than 90% of normal AChE activity.	Holland, et al (1967)
Ortho 5305 EC2	Gambusia affinis	FL	Cal.	0.8 (K1)	-	At a concentration of 0.2 lb/acre, 88% mortality of the fish occurred in 24 hours. At 0.8 lb/acre, 100% mortality occurred.	Mulla (1966)

	Ortho 5353 EC2	Gambusia affinis	FL	Col.	1.0 (K1)	-	Toxicity value is in Ib/acre.	Mulla (1966)			
	Ovex, Tech.	Rainbow trout	BSA	-	0.620 (T4A)	_	The values reported are given as $LC_{50}$ .	Cope (1965)			
	Ovex	Pimephales promelas	BSA		2.5 (T4A)	acdef	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson (1966)			
	Oxydemeton- methyl	Pteronarcys californica (naiads)	BSA	_	0.035 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)			
	Paramar-50	Cyprinus carpio	BSA	-	6.5 (T2A)	acdefp	The test animals were conditioned for 48 hours prior to use.	Sreenivasan and Swanithan			
		Tilapia massambica			4.0-5.0 (T2A)			(1967)			
		Cirrhina mrigala			5.0 (T2A)						
		Labeo fimbriatus			7.5 (T2A)						
		Barbus machecola			2.0 (T2A)						
;	Para-oxon	Pimephales promelas	BSA	_	0.33 (T4A)	<u>acdef</u>	Tests were performed in both hard and soft water. Addi- tional tolerance limit values are given.	Henderson and Pickering (1958)	APPE		
	Para-oxon	Fathead minnow	BSA	-	.25 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959)	APPENDIX B		
	Para-oxon	Pimephales promelas	BSA	-	0.33 (T4A)	adef	Concentrations were based on percent active ingredient.	Henderson, et al (1959)			
COMMERCIAL	Para-oxon	Pimephales promelas	BSA	_	0.33 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not signifi- cantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	Tarzwell (1959)			
CHEN	Paraoxon	Lepomis gibbosus	BSA	-	2/4 (O)	-	The figures given are for mortality in 2 hours when the amount of chemical was 16 mg/kg, given by injection.	Murphy (1966)			
AICA	Paraoxon	Lepomis gibbosus		-	0.120 ± 0.022 (O)	-	This paper is a study of the amounts of organic thiophos- phate and their oxygen analogues which accumulate in liver	Murphy (1966)			
L PR		ictalurus melas	urus		(		0.122±0.005 (O)		slices in an <i>in vitro</i> study of insecticides. The numbers given are for mµm of chemical (in the case of Parathion,	(1900)	
		Pseu dopleu ronectes americanus			0.041 ± 0.006 (O)		Malathion, and Guthion-the oxygen analogue) accumu- lated in 50 mg of liver (wet weight) in 10 minutes.				
CTS		Myxocephalus scorpius			0.076 ± 0.010 (O)						

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Paraquat (cation)	Lepomis macrochirus	BSA	_	400 (T1A)	_	The bioassay methods employed in this experiment were not given in the paper but it was stated that the same pro- cedures were employed as in previous work.	Davis and Hughes (1963)
Paraquat	Lepomis macrochirus	BSA		5 (O)	_	Toxicity to fish was determined as the threshold concen- tration (LD <sub>10</sub> ) in 96 hr at 75 F (65 F for trout). Herbi-	Lawrence, et al (1965)
	Micropterus			5 (O)	-	cidal evaluations are also presented.	(1903)
	salmoides Pimephales			5 (O)	_		
	promelas Ictalurus			5 (0)			
	punctatus Salmo gairdnerii			5 (O)	_		
Paraquat	Spirodela polyrhyza	BSA	-	(O)	a	0.01 ppm caused 82% chlorosis in 7 days.	Blackburn and Weldon
	Lemna					0.01 ppm caused 72% chlorosis in 7 days.	(1965)
	minor Wolffiella					0.01 ppm caused 62% chlorosis in 7 days.	
	floridana Azolla					0.01 ppm caused 40% chlorosis in 7 days.	
	caroliniana Wolffia columbiana					0.01 ppm caused 3% chlorosis in 7 days. Light intensity was kept at 500 foot-candles for 14 hours per day. Light has been shown to increase the rate of kill with paraquat. Test containers were plastic petri dishes.	
Paraquat	Lemna minor Spirodela polyrhyza Wolffia columbiana	۴L	Fla.	(0)	_	Common duckweed and watermeal in small ponds can be controlled with paraquat at rates as low as 0.25 ppmw, but rates greater than 0.5 ppmw are required for control in ponds infested with watermeal.	Blackburn and Weldon (1965)
Paraquat	Fundulus similis (juveniie)	BSA		(O)	а	Water temperature was 19 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Paraquat	Oyster	BCF	_	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
Paraquat	Penaeus az tecus	L		(0)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)

	Paraquat	Crassostrea virginica Penaeus	BCFA & BSA	-	1.0 (NTE) 1.0 (NTE)	-	Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with	Butler (1965)
		aztecus Fundulus similis			1.0 (NTE)		fish were static. Toxicity was reported for the following: Oyster – 96-hr EC50 – Conc. which decreased	
		Phytoplankton			53% (O)		shell growth. Shrimp – 48-hr EC <sub>50</sub> – Conc. which killed or paralyzed 50% of test animals. Fish – 48-hr EC <sub>50</sub> – Conc. which killed 50%. Phytoplankton – Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	
	Paraquat	<i>Pteronarcys</i> sp (nymphs)	BSA	-	>1000 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
	Paraquat	Daphnia magna	BSA	-	11.0 (9.1-12.2) (O)	acdiq	Toxicity, in terms of median immobilization concentration ( $1C_{50}$ ), is presented.	Crosby and Tucker (1966)
	Paraquat	Simocephalus serrulatus	BSA	-	4.0 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr.	Sanders and Cope
_		Daphnia pulex			3.7 (SB)		Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	(1966)
101	Paraquat	Pteronarcys californica (naiads)	BSA		100 (NTE)	<u>acdef</u>	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
	Parathion	Bluegill Rainbow trout Brown trout	BSA	-	(0)	-	A concentration of 0.2 ppm was near the concentration threshold for bluegills. Concentrations of 0.063, 0.189, and 0.378 ppm did not kill 1-inch rainbow and brown trout.	Linduska and Surber (1948)
	Parathion (15 per- cent) 0,0-Diethyl- o-p-nitrophenyl thiophosphate)	Lymnaeid snails	BSA	_	(O)	-	Each test container, 500-ml beaker, was filled with ditch water. 100% mortality occurred in concentrations of 1:400,000 and greater.	Batte, et al (1951)
COMMERCIA	Parathion	<i>Simulium</i> sp (Iarvae)	FR	Streams, S. C. and Fla.	0.5-1.0 (O)	_	In slow-moving streams in Florida, parathion at the indi- cated concentrations eliminated blackfly larvae for dis- tances up to 1.6 miles. In South Carolina, 100 percent reductions for distances of up to 2.8 miles was obtained. Data are presented as percent larval detachment in 1, 2, and 3-days time.	Davis, et al (1957)
IL CHEMIC	Parathion (20% tech, para- thion and 80% triton X-100)	Rainbow trout Eastern brook trout	BCFA	_	0.05-1.0 (K 0%)	<u>a</u>	Spring water (46 F) was used. This flow rate was 10 GPM. The chemical was added by continuous drip dispenser. 0.01 ppm kill mosquito larvae.	Hoffman (1957)

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Parathion	Various	FL	Salt Lake Co., Utah	(0)	_	The chemical was applied at 0.05 lb/acre. Careful application of the chemical at the above concentra- tion controlled mosquito larvae without loss of fish. At the above concentration no adverse effects were observed in mammals, birds, reptiles, and amphibians although some frogs were killed by the application of parathion at several times the normal concentration. Invertebrates were not affected uniformly. Crustaceans were in harmed, nor were larvae of the insect family Ephydridae. Spiders and aquatic insects other than Ephydridae were adverse affected in varying degrees. Aquatic beetles seemed to be af- fected more seriously than other insects excepting mosquito larvae.	
Parathion	Pimephales promelas Lepomis macrochirus	BSA	-	1.4 (T4A) 0.71 (T4A)	acdef	Tests were performed in both hard and soft water. Addi- tional tolerance limit values are given.	Henderson and Pickering (1958)
Parathion	Artemia salina	BSA	-	0.43 (L<1)	ai	Rock salt was used in rearing all cultures employed in bio- assay work. The optimum salt concentration was 3.5%.	Tarpley (1958)
Parathion	Fathead minnow	BSA	-	1.6 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Rela- tive Toxicity of Ten Chlorinated Hydrocarbon Insecticides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) re- port widely different values for the same compounds.	Tarzwell (1959)
Parathion	Daphnia magna	BSA	-	0.0008 (O)	a	The indicated concentration immobilized <i>Daphnia</i> in 50 hours.	Anderson (1960)
Parathion	Pimephales promelas	BSA	-	1.4 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1960)
Parathion	Gambusia affinis	BSA	-	0.004 (K 33%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as avg. percent fish killed in 24 hr.	Lewallen (1959)
Parathion	Pimephales promelas	BSA	-	1.4 (T4A) 0.700 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzweli (1959)
Parathion	Lepomis macrochirus Micropterus salmoides Notemigonus crysoleucas	BSA	-	0.1 (O) 0.1 (O) 0.1 (O)	<u>acdf</u>	This paper reports the effect of insecticides in reducing the anticholinesterase in a fish brain within 2-8 hours. The inhibition of the enzyme was a function of the concentration of the insecticide, extent of exposure, and specific chemical nature of the compound.	Weiss (1959)

		Carassius auratus			0.1 (O)			
	Parathion, (tech, 25% active in xylene)	Acroneuria pacifica Pteronarcys californica Arctopsyche	BSA	-	0.0001 (T4A) 0.0032 (T4A) 0.001 (T4A)	<u>a</u> c <u>e</u> fin	Assays were conducted in hard water.	Gaufin (1961)
	Parathion (tech grade emulsi- fied in exlene)	grandis Hydropsyche californica Acroneuria pacifica Arctopsyche grandis Pteronarcys californica	BSA	-	0.00043 (T4A) 0.001 (T4A) 0.007 (T4A) 0.0086 (T4A)	<u>a</u> cdelm	Test water was obtained from a mountain stream.	Gaufin, et al (1961)
	Palathion (50 percent EC)	Gambusia affinis	FL	Ponds— Bakers- field, Cal.	(0)	a c	At 0.1 lb/acre, 22 percent mortality occurred in 24 hours. At 0.4 lb/acre, 92 percent mortality occurred in 24 hours. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)
B-183	Parathion	Anopheles quadrimaculatus Aedes aegypti A. taeniorhynchus	BSA	-	0.01 (K1) 0.005 (O) 0.005 (O)	-	4th instar larvae of mosquitos were used in this bioassay.At the indicated concentrations, the following mortalitiesoccurred:Anopheles quadrimaculatus94%Aedes aegypti52%A. taeniorhynchus78%Adsorption was determined by use of P <sup>32</sup> labeled parathion.	Schmidt and Weidhaas (1961)
	Parathion	Micropterus salmoides Pimephales promelas	BSA	-	0.5 (O) 0.5 (O)	-	The degree of reaction to the cholinesterase-inhibiting in- secticides is not only a function of time and concentration, but also of chemical and biological species. This paper re- ports many analyses of brain cholinesterase activity which is expressed as percentage of normal. The data are reported as $LT_{50}$ which was the time required for 0.5 ppm of the chemical to kill 50 percent of the fish. For bass the $LT_{50}$ was 24 hr and for the fathead 72 hr.	Weiss (1961)
COMME	Parathion	Salmo gairdnerii (one wk. old sac fry)	BSA	_	0.2 (K 0%) 2.0 (K 0%) 0.2 (K 0%)	<u>a e</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
RCI/		(one mo. old feeding fry)	BSA		2.0 (K 80%)			
COMMERCIAL CHEMICAL PROD	Parathion, (tech, 99 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes	BSA	_	1.3 (T4A) 0.095 (T4A) 2.7 (T4A) 0.056 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)
Õ		reticulatus						

· <u></u> , ,,,	
Chemical	Org
Parathion,	Pimep
emulsible	prom
concentrate	Lepon
(25 percent)	•
(20 pt. 00.11)	Carass
	aurat
	Green
	sunfi
	Larger
	bass
Parathion	Culex
	quad
Parathion	Gamb

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Parathion, emulsible concentrate (25 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus Green sunfish Largemouth bass	BSA	_	3.0 (T4A) Large 0.58 (T4A) Small 0.26 (T4A) 2.6 (T4A) 1.7 (T4A) .0.76 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TLm's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)
Parathion	Culex pipiens quadrimaculatus	BSA	_	(0)	c	Tests were conducted in tap water and artificially polluted tap water. The values reported are the concentration range for an LCg0, 0.034 to 0.1100 ppm for polluted and 0.0072 to 0.0140 ppm for tap water.	Lewallen and Wilder (1963)
Parathion (EC2)	Gambusia affinis Bufo boreas Scaphiopus hammondi	FL	Ponds in III.	(0)	-	When applied at 0.4 pounds per acre active ingredient, 96 percent fish mortality occurred in 1 day. No toad mortality occurred at 0.4 pound per acre in 1 day.	Mulla, et al (1963)
Parathion (tech, 95 percent active in acetone)	Pteronarcys californica (naiad) Acroneuria pacifica (naiad)	BSA	-	0.0320 (T4A) 0.0028 (T4A)	c d e f	A. pacifica was much more sensitive to chlorinated hydro- carbons and to organic phosphate insecticides than P. californica.	Jensen and Gaufin (1964)
Parathion	Gammarus Iacustris	BSA	-	0.0128 (T4A)	<u>a e</u>	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Parathion	Bluegill	BSA	-	0.095 (T4A)	а	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposed to the toxicant.	Weiss (1964)
Parathion	Carassius auratus Lepomis macrochirus Notemigonus crysoleucas	BSCH		10.0 (0)* 1.0 (0)* 0.1 (0)** 1.0 (0)* 1.0 (0)* 0.1? (0)* 0.1? (0)* *response, 15 days **no response, 15 days ***no response, 30 days	<u>a</u> cd <u>e</u>	Toxicity was determined by measuring acetylcholinesterase activity in the brains of fish. Concentrations are given in ppb as either response or no response in 15 or 30 days.	Weiss and Gakstatter (1964)

	Parathion	Acroneuria	BSA	-	0.0028 (T4A)	ac	Additional TL <sub>m</sub> data are given.	Gaufin, et al (1965)
		pacifica Ephemerella grandis			0.003 (T4A)			(1505)
		Gammarus Jacustris			0.0128 (T4A)			
		Pteronarcys californica			0.032 (T4A)			
	Parathion	Arctopsyche grandis	BSA		0.007 (T4A)	<u>a</u>	Unspecified chemical characteristics of assay water were determined by standard methods. General comments	Gaufin, et al (1965)
		Pteronarcys californica			0.03 (T4A)		were made concerning "standardized" conditions, use of "soft" water, and use of emulsifying agents. Additional	
		Acroneuria pacifica			0.003 (T4A)		data are presented.	
		Ephemerella grandis			0.003 (T4A)			
		Hydropsyche californica			0.0004 (T4A)			
		Daphnia magna			0.0008 (T 50 hr A)			
		Gammarus lacustris			0.01 (T4A)			
		Bluegill sunfish			0.06 (T4A)			
		Fathead minnow			1.4 (T4A)			
ň	Parathion	Lepomis gibbosus	BSA	-	0/4 (0)	-	The figures given are for mortality in 2 hours when the amount of chemical was 40 mg/kg, given by injection: number dead/number injected.	Murphy (1966)
	Parathion	Lepomis gibbosus			19.97 ± 7.09 (O)	-	This paper is a study of the amounts of organic thiophos- phate and their oxygen analogues which accumulate in	Murphy (1966)
		ictalurus melas			14.52 $\pm$ 1.56 (O) liver slices in an <i>in vitro</i> study of insecticides. The	liver slices in an <i>in vitro</i> study of insecticides. The numbers given are for m $\mu$ m of chemical (in the case of Para-	(1000)	
		Pseudopleuronectes americanus			5.20±0.81 (O)		thion, Malathion, and Guthion—the oxygen analogue) accumulated in 100 mg (dry weight) of liver in 30	
		Myxocéphalus scorpius			0.4 ± 0.2 (O)		minutes.	
COMMERCIAL CHEMICAL PRODUCTS	Parathion	Sewage organisms	BOD	-	(0)	-	Shock loadings of the chemical as high as 15.0 mg/l were assimilated by microbial systems of 500.0 mg/l with no observable toxic effect. Organophosphate insecticides in low concentration stimulated microbial respiration; however, greater concentrations inhibited the system and eventually destroyed the organisms. The amount of organophosphate insecticide required to inhibit the respira- tion of freshwater microorganisms was shown to be func- tion of the amount of organisms present and not the volume of water in which the organisms are dispersed.	Randall (1966)
AL PRODUCTS	Parathion	Simocephalus serrulatus Daphnia pulex	BSA	-	0.00037 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location <sup>(2)</sup>	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Parathion	Tubifex spp Limnodrilus spp	BSA	-	5-2 (L4A)	асе	Toxicity is reported as the mean lethal dose (LD $_{\rm 50}$ ) for 24, 48, and 96 hours.	Whitten and Goodnight (1966)
Parathion	Leiostromus xanthurus Cyprinodon variegatus	ВСГСН	-	0.01 (O) 0.01 (O)	а	At a concentration of .01 ppm, the following percent acetylcholinesterase activity as compared to controls was found: <i>L. xanthurus</i> – 10 <i>C. variegatus</i> – 26	Butler and Johnson (1967)
Parathion	Micropterus salmoides	BSA	-	2.0 (O) 5.0 (K 3 hr)	a e	At 2.0 ppm, 40 percent mortality occurred in 1 day. Experi- ments were carried out in plastic tubs lined with saran plastic. Fish weights averaged 217 grams.	Mulia, et al (1967)
Parathion	Pteronarcys californica (naiads)	BSA	-	0.0054 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Parathion	Pteronarcys californica (naiads) Pteronarcella badia (naiads) Claasenia sabulosa (naiads)	BSA	-	0.0054 (T4A) 0.0042 (T4A) 0.0015 (T4A)	<u>acdef</u>	Comment same as above.	Sanders and Cope (1968)
Parathion	Lepomis macrochirus	BSSB	-	0.0075, 0.032, and 0.087 (O)	<u>a c f</u>	Critical flicker frequency response in the bluegill was measured by determining this species ability to maintain position relative to continuously rotating stripes. Increas- ing or decreasing the rate of movement of the stripes above or below a certain critical flicker threshold caused the fish to return to random swimming. The effect of different amounts of insecticide was measured. An aberrant response was noted at all three sublethal con- centrations noted.	Scheier and Cairns (1968)
Parathion	Lepomis gibbosus Ictaluras Micropterus dolomieui Myxocephalus scorpius Pseudopleuronectes americanus	BSA	-	(O)	ар	The chemicals were poor inhibitors of brain cholinesterases <i>in vitro;</i> their oxygen analogs were potent inhibitors.	Murphy, et al (1968)
Parzate	Channel catfish (fingerlings)	BSA	-	21.1 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)

	Penicillin G, potassium (crystalline)	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = partially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - T So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
	Perthane	Gambusia affinis	BSA	_	10.4 (L1)* 10.0 (L1)** *Resistant fish **Nonresistant fish	а	This paper deals with the resistance of mosquito fish to chlorinated hydrocarbon compounds. Resistant fish were not always less sensitive to these chemicals.	Boyd and Ferguson (1964)
	Perthane (tech)	Rainbow trout	BSA	·	0.005 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
	Phorate (Thimet)	- Oyster	BCF	-	0.64 (O)	а	The value reported is a 96-hr $\text{EC}_{50}$ (decreased shell growth).	Butler (1965)
	Phosdrin	Rainbow trout Eastern brook trout	BCFA	-	0.05 (K 3 day) 0.1 (K 4 hr) 0.5 (K 80 min) 1.0 (K 30 min) 10.0 (K 15 min)	ar	Spring water (46 F) was used. The flow rate was 10 gpm. The chemical was added by continuous drip dispenser. 0.01 ppm for 180 hr showed toxic effects, but no kill.	Hoffman (1957)
•	Phosdrin (tech)	Rainbow trout Bluegill	BSA	_ ·	0.012 (T4A) 0.023 (T4A)	, <del></del>	The values reported are given as $LC_{50}$ .	Cope (1965)
	Phosdrin	Pteronarcys sp (nymphs)	BSA	—	0.0049 (T4A)	a	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
2	Phosdrin	Simocephalus serrulatus Daphnia pulex	BSA	-	`0.00043 (SB) 0.00016 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
OMMER	Phosdrin	Pteronarcys californica (naiads)	BSA	-	0.00 <u>5</u> (T4A)	acdef	Data reported as $LC_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
	Phusphamidon (tech)	Procambarus clarki	BSA	-	5.5 (T3A)	acdo	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
EMIC	Phosphamidon	Salmo gairdneri	BSA	<b>-</b> .	5,000 (T.1A)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Соре (1963)
	Phosphamidon	Salmo gairdneri	BSA	-	5,000 (T 18 hr)	ä	Comment same as above.	Cope (1963)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Phosphamidon	Simocephalus serrulatus Daphnia pulex	BSA		0.012 (SB) 0.0088 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Daphnia magna	BSA	_	0.0125 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
Phosphamidon	Coho salmon (fry)	FR	Moran Creek, B.C.	7.0 (T2A)	а	The data indicated 1.0 to 3.2 ppm would cause a slight mortality of juvenile coho salmon. The value of 7.0 ppm was taken from a preliminary bioassay.	Schouwenberg and Jackson (1966)
Phosphamidon (in water)	Atlantic salmon Brook trout	FR	New Brunswick	(0)	-	Spraying with this chemical at 1.0 lb per acre had no apparent harmful effects on young salmon or trout.	Kerswill and Edwards (1967)
Phosphamidon	Pteronarcys californica (naiads)	BSA	-	0.15 (T4A)	acdef	Data reported as LC $_{\!$	Sanders and Cope (1968)
Phygon XL	Oncorhynchus kisutch Micropterus sølmoides	BSA	-	0.042 (NTE) 0.08 (T1A) 0.07 (T2A)	a c d e	Concentrations were based on percent active ingredient. No toxicity to <i>O. kisutch</i> occurred at concentrations up to 0.042 ppm.	Bond, et al (1960)
Phygon-XL	Channel catfish (fingerlings)	BSA	-	0.14 (K 29 hr A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Phygon-XL	Salvelinus fontinalis x Salmo trutta Notemigonus crysoleucas	FPA	N.Y.	0.5 (\$23)	a c d	Conventional farm ponds were used having an average surface area of 0.3 acre and a maximum depth of 7-9 ft. Toxicity (in ppm) to fish as maximum safe concentra- tion (S) for 23 days was determined. Concentration of 0.5 ppm was required to control algae.	Eipper (1959)
	lctalurus punctatus Micropterus			0.5 (S23) 0.5 (S23)			
	salmoides Lepomis macrochirus			0.5 (S23)			
Phygon-XL	Salmo gairdneri	BSA	-	0.075 (T1A) 0.075 (T2A)	<u>a</u> cdfg	Hatchery artesian well water was employed for this experiment.	Webb (1961)
Phygon X-L	Richardsonius balteatus hydrofiox	BSA	-	0.13 (T1A) 0.11 (T2A) 0.11 (T4A)	a c d e f	Results given were in soft water. Results in hard water were as follows: 0.15 (T1A), 0.15 (T2A), and 0.14 (T4A).	Webb (1 <b>96</b> 1)

Phygon-XL	Gammarus Iacustris	BSA	-	0.165 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)
Phygon XL (Dichlone)	Salmon	BSA	-	0.043 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
Picloram	Pteronarcys californica (naiads)	BSA	-	0.048 (T4A)	<u>acdef</u>	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
РМА	Salmo gairdneri	BSA	-	3.75 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
	Salmo trutta			6.22 (T2A)			
	Salvelinus fontinalis			10.7 (T2A)			
	Salvelinus namaychush			7.60 (T2A)			
	lctalurus punctatus			2.89 (T2A)			
	Lepomis macrochirus			16.0 (T2A)			
Polyclens	Pandalus montagni	BSA		8.5 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C.	Portmann and Connor
	Crangon crangon			15.7 (T2A)		It was shown that the toxicity of this solvent emulsifier decreased with time due to evaporation of the solvent.	(1968)
	Carcinus maenas			23.2 (T2A)		Polyclens at a concentration of 3.3 ppm killed 100% of Crangon crangon larvae in 3 hr.	
	Cardium edule			70.0 (T2A)			
Polysan	Guppy	BSA	-	100 (K 25 min)	<u>a</u>	Those fish that survived at lower concentrations were still very active several days after they had been taken out and placed in fresh water.	Anonymous (1964)
Prometone [2-methoxy- 4,6-bis(iso- propylamino)- s-triazine (prometone)]	Elodea canadensis Potamogeton	BSA	-	5 (O) 100 (O) 5 (O)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. Injury rating of 0.5. Injury rating of 3.6. Injury rating of 2.3.	Frank, et al (1961)
2	nodosus Potamogeton pectinatus			100 (O) 5 (O) 100 (O)		Injury rating of 7.0. Injury rating of 5.1. Injury rating of 8.3.	
Prometone	Leiostomus xanthurus (juvenile)	BSA	-	(O)	а	Water temperature was 26 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Prometone	Penaeus duorarum	L	_	(O)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
Prometone	Oyster	BCF	-	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Prometone	Crassostrea virginica Penaeus duorarum Leiostomus xanthurus Phytoplankton	BCFA & BSA	_	1.0 (NTE) 1.0 (NTE) 1.0 (NTE) –		<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Prometryne	Oyster <i>Leiostomus xanthurus</i> (juvenile)	BCF BSA	-	(O)	8	Exposure to a concentration of 1 ppm caused a 19.0% decrease in shell growth. Water temperature was 28 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Prometryne	Peneas du orarum	L	-	(0)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr $EC_{50}$ or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
Prometryne	Crassotrea virginica Penaeus duorarum Leiostomus xanthurus Phytoplankton	BCFA & BSA	_	1.0 (0, 19%) 1.0 (NTE) 1.0 (NTE) —	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Pro-noxfish (25% rotenone + 2.5% sulfoxide)	Cyprinus carpio (eggs) (fry) Pimephales promelas (eggs) (fry)	BSA	_	0.178 (K) 0.163 (K) 0.233 (K) 0.191 (K)	<u>a</u>	Toxicity is reported as LD <sub>50</sub> in ppm, at 75 F.	Hester (1959)
Pro-noxfish (2.5% rote- none, 2.5% sulfoxide, 5% cube extracts emulsifier)	Cyprinus carpio Micropterus salmoides Pimephales promelas Carassius auratus	BSA	-	0.163 (K3) 0.081 (K3) 0.191 (K3) 0.242 (K3)	<u>a</u> cdei	Such variables as temperature, species, and size of fish were studied. Toxicity is expressed as $LD_{50}$ for 72 hr. Smaller concentrations of rotenone were required when used in conjunction with sulfoxide. The data shown are for 70 F. The chemical was considerably more toxic at this temperature than at 40 F for all fish species.	Hester (1959)

8:825 (23)

		Lepomis macrochirus L. cyanellus Notemigonus crysoleucas Ictalurus nebulosus marmoratus			0.255 (K3) 0.238 (K3) 0.555 (K3) 0.410 (K3)			
	<b>Pro-noxfish</b>	Oncorhynchus kisutch (eggs)	FR	Ore.	(O)	a	<ul> <li>Pro-noxfish is a formulation containing 2.5% rotenone, 5.0% related rotenoids and cube extractives, 2.5% sulfoxide synergists, and 90% solvent emulsifier.</li> <li>The goal of this experiment was to expose the eggs to the chemical at a concentration of 2 ppm for 24 hr.</li> <li>High survival occurred where the temperatures ranged from 46 to 56 F.</li> <li>High temperatures of 60 and 65 F occurred in Middle Fork and Quartzville Creek and contributed to the mortality rate. No eggs survived.</li> </ul>	Garrison (1968)
<b>B</b> -191	Pro-noxfish	Oncorhynchus tshawytscha (fry, 100-day old) Oncorhynchus kisutch (eggs)	BSA	Corvallis, Ore.	0.15 to 5.0 (K1) (O)	<u>a e</u>	<ul> <li>Pro-noxfish is a formulation containing 2.5% rotenone, 5.0% related rotenoids and cube extractives, 2.5% sulfoxide synergists, and 90% solvent emulsifier.</li> <li>Experiments were conducted in aerated test jars.</li> <li>Temperature was 53 F.</li> <li>Temperature seems to have an influence upon toxicity.</li> <li>Embryos exposed to the chemical for 24 hr showed the follow- ing survival rates. All embryos survived in 1.0 ppm at 53 F, all survived in 3 ppm at 46 F, and 90% survived in 4 ppm at 39 F.</li> </ul>	Garrison (1968)
	Propanil (Stam, Rogue)	Daphnia magna	BSA	_	4.8 (3.8- 6.6) (O)	acdiq	Toxicity, in terms of median immobilization concentration $(IC_{50})$ , is presented.	Crosby and Tucker (1966)
	Pyrethrin	Pteronarcys sp (nymphs)	BSA	-	0.001 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
COMM	Pyrethrins	Simocephalus serrulatus Daphnia pulex	BSA	-	0.042 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
COMMERCIAL CHEMICAL PRODUC	Pyrethrins	Salmo gairdneri Lepomis macrochirus Ictalurus	BSA	_	0.054 (T2A) 0.070 (T2A) 0.082 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on repro- duction and behavior are also discussed. Data presented as EC <sub>50</sub> .	Cope (1966)
MICAL		punctatus Pteronarcys californicus			0.006 (T2A)			
. PRODU		Daphnia pulex Simocephalus serrulatus			0.025 (T2A) 0.042 (T2A)			

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Pyrethrum	Black fiy (larvae) Rainbow trout	FR	Alaskan streams	0.1 (O) 0.1 (O)	-	The chemical was applied to control black flies, and because the acetone solution was most effective, only that data is reported here. The figures reported are for minimum effec- tive dosages for black fly larvae and maximum nonlethal dosage for rainbow trout.	Gjulian, et al (1949)
Pyrethrum	Salmo gairdnerii lctalurus punctatus Lepomis macrochirus Pteronarcys californica	BSA	-	54 (T4A) 80 (T4A) 74 (T4A) 1.0 (T4A)	<u>a</u> cd	Toxicity values reported as median lethan concentration (LC <sub>50</sub> ) for 24, 48, and 96 hr.	Bridges and Cope (1965)
Pyrethrum	Pteronarcys californica (naiads)	BSA	-	0.0010 (T4A)	acdef	Data reported as LC $_{\!$	Sanders and Cope (1968)
Quaternary ammonium salt, commercial	Microcystis aeruginosa	L	-	2.0 (K)	<u>a,</u> etc	The chemical was tested on a 5-day algae culture, $1 \times 10^6$ to $2 \times 10^6$ cells/ml, 75 ml total volume. Chu No. 10 medium was used.	Fitzgerald, et al (1952)
Rivanol	Channel catfish (fingerlings)	BSA	-	2.8 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Roccal	Micropterus salmoides (fry) Ictalurus punctatus (fry) Lepomis macrochirus (fry)	BSA	-	1.0 (SB3) 1.0 (SB3) 0.25 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours.at the concentration listed.	Jones (1965)
Roccal	Guppy	BSA	-	100 (K1)	<u>a</u>	Those fish that survived at lower concentrations were still very active several days after they had been taken out and placed in fresh water.	Anonymous (1964)
Roccal	Salmo gairdneri Salmo trutta Salvelinus fontinalis Salevelinus namaycush Ictalurus punctatus Lapomis macrochirus	BSA	_	2.57 (T2A) 2.05 (T2A) 3.40 (T2A) 1.95 (T2A) 1.12 (T2A) 1.68 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)

	Řonnel	Chaoborus astictopus	BSA	-	0.046 (T1A)	<u>a</u>	Toxicity value given is for the fourth instar larvae.	Hazeltine (1963)
	Ronnel (EC2)	Gambusia affinis Bufo boreas	FL	Ponds in III.	(0)	-	When applied at 0.8 pound per acre active ingredient, no fish mortality occurred in 1 day. At 0.2 pound per acre, 10 percent toad mortality occurred in 1 day. In a duplicate test there was no mortality at 0.8 pound per acre.	Mulla, et al (1963)
	Ronnel	Penaeus aztecus	L	-	0.015 (O)	а	Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
	Ronnel	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	0.32 (O)	а	Water temperature was 13 C. The figure reported is a 48-hr EC <sub>50</sub> .	Butler (1965)
	Ronnel	Oyster	BCF	-	0.17 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
	Rosinamine D acetate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T Ma - T So $- PT$ (7) Cv - T Gp - T Np - T	Palmer and Maloney (1955)
	Rosinamine D acetate	Pimephales promelas	BSA	-	0.23 (T4A)	acdef	Toxicity to 30 species of algae also presented. RADA was algicidal in the range 0.25 to 2.0 ppm.	Maloney and Palmer (1956)
COMMERCIAL (	Rosinamine D sulphate	Cylindrospermum licheniforme (CI) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T Ma - T So $- T$ (14), PT (21) Cv - T Gp - T Np - T	Palmer and Maloney (1955)
CHEMICA	Rosinamine D sulphate (RADS)	Pimephales promelas	BSA	_	0.16 (T4A)	acdef	Toxicity to 30 species of algae also presented. RADS was algicidal in the range 0.25 to 2.0 ppm.	Maloney and Palmer (1956)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Rosinamine D	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 24th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T Ma - T So - T (3), PT (14) Cv - T Gp - T Np - T	Palmer and Maloney (1955)
Rosinamine D pentachloro- phenate	Cylindrospermum licheniforme (CI) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (O)	<u>a</u>	Comment same as above except that: CI — P (7), PT (14) Ma — PT (14) So — NT Cv — T (7), PT (14) Gp — NT Np — T	Palmer and Maloney (1955)
Rosinamine derivative	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	_	2.0 (0)	<u>a</u>	Comment same as above except that: CI – PT Ma – PT (14) So – PT Cv – PT (3) Gp – T Np – T	Palmer and Maloney (1955)
Rotenone (derris or cube with 5% rotenone)	Salvelinus fontinalis (yearling) Couesius plumbeus Catostomus commersonnii Eels Pungitius pungitius Micropterus dolomieu	FL	4 lakes, Nova Scotia	0.20 (K) 0.20 (K) 0.20 (K) 0.25 (K) 0.25 (K)	a c d f	The work was undertaken to test the feasibility of utilizing poisons as a direct means of studying the production of fish in streams and lakes. Caution must be used to prevent irreparable damage by indiscriminate poisoning.	Smith (1939)

		Morone americana			0.25 (K)				
	Rotenone	Carassius auratus	BSA	_	0.100-2.00 (K 8 hr) 0.0600 (O) 0.0400 (T 8 hr) 0.0200 (O)	<u>a</u>	Temperature in test containers was maintained at 27 $\pm$ 0.2 C. Goldfish tested weighed between 2 and 4 g. Rotenone, 0.0600 mg per liter, killed 86% of the fish in 8 hr; 0.0200 mg per liter killed 18% in 8 hr.	Gersdorff and Smith (1940)	
	Rotenone	Perca flavescens Brown trout Rock bass Creek chub Smallmouth bass Common sucker Brown bullhead	BSA	-	0,45 (K) 0,20 (K) 0,32 (K) 0,35 (K) 0,40 (K) 1,7 (K) 2,2 (K)	acdep	A range of concentrations between 0.05 and 0.8 ppm was used in this study and kill occurred in 1 to 4 hr.	Burdick, et al (1956)	
a J	Rotenone (5% cube)	Channel catfish (fingerlings)	BSA	_	0.51 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	•
0.5	Rotenone	Chironomus plumosus (larvae)	BSA FLA	 Lake Erken, Sweden	(O) (O)	<u>a c</u>	Laboratory studies were with and without silt. Without silt 100% kill occurred in 0.3 ppm rotenone, while 50% kill occurred at 3.0 ppm with silt present. Further data were obtained from field studies and from caged animal studies at various depths and sections of the lake. Data on more than 200 species are presented at 0.5 ppm rotenone lake- bottom-dwelling organisms exhibit sensitivity. Use of higher concentrations than this would mean partial or complete disappearance of many species.	Lindgren (1960)	
COMM	Rotenone (2.5 percent, 5 percent cube extractives, and 2.5 percent sulfoxide)	Pimephales promelas	BSA	-	0.066 (T4A)	a c d f g	Test water was spring water diluted with distilled water. Removal of toxic chemicals by carbon adsorption, chlorine and chlorine dioxide treatment, and alum coagulation was studied. The most effective method to remove fish poisons was by use of activated charcoal adsorption.	Cohen, et al (1961)	
ERCIAL C	Rotenone (5 percent and 15 percent toxaphene)	Pimephales promelas	BSA	-	0.066 (T4A)	acdfg	Comment same as above.	Cohen, et al (1961)	
HEMICAL	Rotenone (2.0 percent and 7.0 percent toxaphene)	Pimephales promelas	BSA	-	0.10 (T4A)	acdfg	Comment same as above.	Cohen, et al (1961)	
COMMERCIAL CHEMICAL PRODUCTS	Rotenone (5 percent and 10 percent other extractives	Pimephales promelas	BSA	-	0.10 (T4A)	a c d f g	Comment same as above.	Cohen, et al (1961)	

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Rotenone	Pimephales promelas	BSA	_	0.006 (T4A)	acdfg	Test water was spring water diluted with distilled water. Removal of toxic chemicals by carbon adsorption, chlorine and chlorine dioxide treatment, and alum coagulation was studied. The most effective method to remove fish poisons was by use of activated charcoal adsorption.	Cohen, et al (1961)
Rotenone	Entomostraca Cyclops Diaptomus Ceriodaphnia Bosmina Leptodora Rotaria Filinia Keratella Polyarthra Asplanchna Brachionus Protozoa Ceratium Difflugia	FL	Col.	1.0 (K)	a c d g p	All chemical and physical data were collected and compiled by standard limnological techniques. Chemical analyses were conducted monthly. Biweekly plankton collection showed "reduction to zero" of all organisms studied, but recovery of populations to normal population numbers within several months.	Hoffman and Olive (1961)
Rotenone (2.5 percent)	Fish	FL	Mont.	0.95 (O)	a c d e l	Ponds were treated with the chemical to eradicate fish. The fish population included largemouth bass, bluegills, black crappie, yellow perch, carp, white sucker, and longnose sucker. Counts were made of various fish at various later times. The paper contains little quantitative data.	Wollitz (1963)
Rotenone (5.5 percent, cube extract 11.00 percent)	Fish	FL	Mont.	0.7 (0)	a c d e l	Comment same as above.	Wollitz (1963)
Rotenone	Zooplankton	FL	Fern Lake	0.5 (O)	acdef	Rotenone (5%) was applied at the rate of 0.5 ppm. Samples were taken biweekly. Open water species were completely removed, and remained absent for 3 mo. Organisms along the shore edge resisted the effect of	Kiser, et al (1963)
			Silver Lake	1.0 (O)		rotenone, but eventually disappeared for several weeks. Rotenone (5%) was applied at the rate of 1.0 ppm. After application the greatest reduction, about 70%, occurred within an hour. Two days after application, no zooplankton were found alive in the open-water tows taken at all depths in the lake. The rotenone penetrated to the thermocline at the 30-ft depth in the first 6 hr, killing <i>Cladocera</i> and <i>Copepoda</i> as it sank.	
Rotenone	Gammarus Iacustris	BSA	-	3.52 (T4A)	<u>a</u> e	Emulsible concentrates were prepared from technical grade insecticides with acetone as the solvent. Symptoms prior to death were observed and recorded on graphs.	Nebeker and Gaufin (1964)

	Rotenone	Nereis limnicola	FL	Lake Merced, Cal.	0.025 (K)	-	In a fish killing program, 0.025 ppm of rotenone was used on October 26, 1963. By November 18 the population of the nereid had been reduced from 500/m <sup>2</sup> to no greater than 10/m <sup>2</sup> . How important this organism is in the com- plex food chain is unknown.	Oglesby (1964)
	Rotenone	Salmo gairdnerii Ictalurus punctatus Lepomis macrochirus Pteronarcys	BSA	-	27 (T4A) 28 (T4A) 23 (T4A) 250 (T4A)	<u>a</u> cd	Toxicity values reported as median lethal concentration (LC <sub>50</sub> ) for 24, 48, and 96 hr.	Bridges and Cope (1965)
		californica			200 (144)			
	Rotenone	<i>Pteronarcys</i> sp (nymphs)	BSA	-	0.250 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
	Rotenone	Anax Agrion Siphlonurus Phryganea	BSA	-	2.3 (T2A)	a c c g	Death caused by rotenone is caused by the constriction of the gill capillanes which prevent the passage of blood through the gills.	Claffey and Ruck (1967)
	Rotenone	Simocephalus serrulatus Daphnia pulex	BSA	-	0.190 (SB) 0.100 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Rotenone	Pteronarcys californica (naiads)	BSA	-	0.38 (T4A)	acdef	Data reported as $LC_{\overline{50}}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
COMMERCIAL CHE	Ruelene	Salmo gairdneri Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	-	32.0 (T2A) 25.7 (T2A) 35.0 (T2A) 27.0 (T2A) 34.8 (T2A) 35.0 (T2A)	f	Variance and the 95 percent confidence interval (C.I.) were also determined.	Willford (1966)
JAL CHE	Ryania (Ryanicide 100)	Cyprinus carpio	BSA	-	(O)	a c e	0 percent mortality occurred in 4 days at 0.01 ppm. 0 per- cent mortality occurred in 4 days at 3 ppm.	Hayes (1955)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(</sup> 3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Saponin	Shrimp: Caridina denticulata Penaeus carinadus Fish: Elops saurus Tilapia mossambica	BSA & FL	Taiwan	50.0 (K1) 70.0 (K1) 1.0 (K1) 1.5 (K)	a	Saponin derived from <i>Camellia</i> spp selectively killed fish in bioassays at the concentrations noted while 50-70X higher concentrations were required to kill shrimp. Concentrations of saponin ranging from 2.5-10.0 ppm were similarly effective in pond studies in killing wild fish which prey on or compete with shrimp. This appears to be a good treatment in shrimp culture for shrimp predator control, but the authors caution that further investigation is required.	Tang (1961)
7 Sarín	Pimephales promelas Lepomis cyanellus Carassius auratus	BCFA	_	10-40 (O)	<u>a</u> cde	The time for 50 percent $(T_{50})$ of the fish to die was studied when the toxic material was held constant while dissolved oxygen, temperature and size of fish were varied. Toxic concentrations of sarin were between 10 ppb and 40 ppb oxygen consumption rates are also reported.	Weiss and Botts (1957)
Sərin	Pimephales promelas Lepomis cyananellus Carassius auratus	BSA	_	0.1 to 50.0 (O)	<u>acd</u>	Data are presented as $TL_m/degree$ centigrade with some of the results as follows: 50 ppm at 24 C was lethal in 0.8 minutes and at 1.20 C in 1.3 minutes for fathead minnows. 50 ppm at 24 C was lethal to sunfish in 0.95 minutes, and at 12 C in 1.55 minutes. 50 ppm at 24 was lethal to goldfish 1.5 minutes, and at 12 C, 2.3 minutes. The toxicity of Sarin was shown to be very temperature dependent. Considerable additional data are presented.	Weiss and Botts (1957)
Sarin	Lepomis macrochírus Lepomis cyanellus Pimephales promelas Lebistes reticulatus Carassius auratus	BSA + BCFA	_	(S) 3.2 (T4A) (H) 23.5 (T4A) (S) 4.2 (T4A) (H) 15.2 (T4A) (S) 4.4 (T4A) (H) 31.9 (T4A) (S) 1.4 (T4CF) (H) 4.2 (T4CF) (S) 7.2 (T4A) (H) 13.8 (T4A) (S) 9.8 (T4A) (S) 4.1 (T4CF)	<u>a</u> c d e	Both static and continuous flow bioassays were made in hard (H) and soft (S) waters. 24, 48, and 96-hr $TM_L$ are reported. Sarin was more toxic in hard water.	Pickering and Henderson (1959)
Sarin	Pimephales promelas	BCFA	-	18 ppb (T 2 hr A)	<u>a</u> c <u>e</u> f	Describes a continuous flow method for bioassay of an organo- phosphorus CW agent.	Henderson and Pickering (1963)
Schadran	Channel catfish (fingerlings)	BSA	-	≥8913 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
SD-4294 (EC32)	Gambusia affinis	FL	Ponds in III.	(0)	-	When applied at 0.2 pounds per acre active ingredient, 4 per- cent mortality occurred in 1 day.	Mulla, et al (1963)
SD-4402 (15 percent EC)	Gambusia affinis	FL	Ponds — Bakers- field, Cal.	0.1 (K1) 0.4 (K1)	ac	Toxicity values indicate application rates in Ib/acre. The experiments were conducted in cages placed in the ponds.	Mulla and Isaak (1961)

	SD-7587 (EC2)	Gambusia affinis	FL	Ponds in III,	(O)	-	When applied at 0.40 pounds per acre active ingredient, 32 percent mortality occurred in 1 day.	Mulla, et al (1963)
	SD-8803 (EC2)	Gambusia affinis	FL	Cal.	0.4 (K1)	~	Toxicity value is in Ib/acre.	Mulla (1966)
	SD-9020 (EC2)	Gambusia affinis	FL	Cal.	(O)	~	At a concentration of 0.4 lb/acre, 56% mortality of Gambusia affinis occurred in 24 hours.	Mulla (1966)
	SD-9129 (EC3)	Gambusia affinis	FL	Cal.	(0)	~	At a concentration of 0.8 lb/acre, 16% mortality of the fish occurred in 24 hours.	Mulla (1966)
	Separan (poly- acrylamide)	Rainbow trout	BSCH	-	(0)	a	A concentration of 0.035 and 0.070 ppm of "Separan" for 4 months caused no rainbow trout mortality. No growth retardation was evident in the lot exposed to 0.035 ppm, and only slight retardation occurred at 0.070 ppm.	Olsen and Foster (1958)
	Sernyl	Carassius auratus	BSA	-	36 (T 1.5 hr)	a	Fish reacted sluggishly and remained stationary at all con- centrations evaluated. Median tolerance limits, median lethal concentrations, and the relation of dosage to time were calculated.	Wilber (1965)
	Servin	Leiostomus xanthurus	BCFCH	-	0.1 (SB 90)	а	The toxicity of this chemical to fish was relatively low.	Butler and Johnson (1967)
<b>B-</b> 199	Sevin (50%, and Sevin- tech)	Carassius auratus	BSA	-	25 (K2) 14 (L2) 35 (K2) 28 (L2)	a b	The wettable powder formulation (50% Sevin) was prepared on the basis of active ingredient, and stirred directly into water. As a comparison, results were given for Sevin (tech- nical). The wettable powder appeared to be twice as toxic as the Sevin alone under the conditions of this test.	Haynes, et al (1958) Henderson, et al
	Sevin	Pimephales promelas Lepomis macrochirus	BSA	-	12.0 (T4A) 5.3 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al 🗙 (1960) 🔲
0	Sevin	Pimephales promelas Lepomis macrochirus	BSA	-	12.0 (T4A) 5.3 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)
COMMERCIAL CHEMICAL PRODUCTS	Sevin	Ephemeroptera Plecoptera Coleoptera Trichoptera Diptera Annelida Megaloptera	FR	Oneonta, N. Y.	(0)	-	This chemical was highly toxic to mayflies, stoneflies, and caddieflies at 1/4 lb/acre. The fish food populations of invertebrates in the sprayed sections of the streams were reduced from 50.7 to 97.2 percent.	Burdick, et al (1960)
EMICA	Sevin	Oncorhynchus kisutch	BSA	-	997 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. $TL_{\mathbf{m}}$ expressed in ppb.	Katz (1961)
		Salmo gairdnerii			1,350 (T4A)			
RODUCTS		Ĝasterosteus aculeatus			3,990 (T4A)			

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL PRODUCTS	Sevin	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodactylum tricornutum Monochrysis lutheri	BSA	~	10 (NG) 10 (K) 10 (NG) 0.1 (NG) 10 (K) 1.0 (NG)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were expressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. $NG = no$ growth, but the organisms were viable.	Ukeles (1962)
DUCTS	Sevin (tech)	Procambarus clarki	BSA	-	2.0 (T3A)	acdo	There was no detectable difference in toxicity to male or female crawfish.	Muncy and Oliver (1963)
	Sevin (tech)	Salmo gairdneri	BSA	-	3,500 (T1A) 2,000 (T2A)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Cope (1963)
B-200	Sevin	Aquatic insects: Ephemeroptera Plecoptera Ameletus Iron Heptagenia Brachyptera Alloperla Ephemerella Simulium	FR	Pa.	(0)	_	Insecticide spraying dosage was 1.1 kg/4.21 H <sub>2</sub> O/hectare, covering over 16,000 acres of woodland for control of gypsy moth. It appeared that there was a drastic reduction of the standing crop of aquatic insects as a result of spraying despite precautions taken against direct spraying of open water and washing spray equipment in the streams.	Coutant (1964)
õ	Sevin	Pteronarcys sp (nymphs)	BSA	~	0.0048 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as $LC_{50}$ .	Cope (1965)
	Sevin (tech)	Bluegill	BSA	~	2.0 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
	Sevin (carbaryl)	Brown trout (fingerlings)	BSCFA	~	8,0 (K) 15 to 273 minutes	<u>a</u> c e	No significant different in toxicity was found between flow- through and static evaluations. A wide range of concentra- tions was studied in both hard and soft waters, and a range of sizes of fish were used. The data is given considerable mathematical treatment. The form in which the chemical is used was shown to be important.	Burdick, et al (1965)
	Sevin	Simocephalus serrulatus Daphnia pulex	BSA	-	0.0076 (SB) 0.0064 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
	Sevin (20% active)	Tubifex spp Limnodrilus spp	BSA		750 (L4A)	a c e	Toxicity is reported as the mean lethal dose (LD <sub>50</sub> ) for 24, 48, and 96 hours.	Whitten and Goodnight (1966)

	Sevin (Carbaryl)	Salmo gairdneri	BSA		2.000 (T2A)	а		Cope (1966)
		Lepomis macrochirus			2.500 (T2A)		on reproduction and behavior are also discussed. Data presented as $EC_{EO}$ .	
		lctalurus punctatus			19.000 (T2A)			
		Pteronarcys californicus			0.015 (T2A)			
		Daphnia pulex			0.006 (T2A)			
		Simocephalus serrulatus			0.008 (T2A)			
	Shell 4072	Leiostomus xanthurus (juvenile)	BSA	-	(O)	а		Butler (1965)
	Shell 4072	Oyster	BCF	_	0.60 (O)	а		Sutler (1965)
	Shell SD-7438 (tech)	Rainbow trout Bluegill	BSA	-	0.030 (T4A) 0.250 (T4A)	-	J	Cope (1965)
	Shell	Penaeus	L		0.028 (O)		Toxicant chemicals were evaluated in seawater at tempera-B	lutler
Ψ	SD-7438	aztecus	L	_	0.028 (0)	а		(1965)
B-201	Sheil SD-7438	Oyster	BCF	-	0.10 (O)	а		lutler (1965)
	Shell SD-7961	Crassostrea virginica	BCFA & BSA	-	1.0 (NTE)	-		lutler (1965)
		Penaeus setiferus			1.0 (NTE)		and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:	
		Leiostomus xanthurus			1.0 (NTE)		Oyster – 96-hr EC <sub>50</sub> – Conc. which decreased shell growth.	
		Phytoplankton			(0)		Shrimp – 48-hr EC <sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.	
							Fish 48-hr EC <sub>50</sub> Conc. which killed 50%.	
COM							Phytoplankton — Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	
MERC	Shell SD-7961	Oyster	BCF	-	(0)	а	• • • • • • • • • • •	utler (1965)
COMMERCIAL CHEMICAL PRODUCTS	Shell SD-7961	<i>Leiostomus xanthurus</i> (juvenile)	BSA	_	(O)	а	•	utler (1965)
EMICAL	Shell SD-8447	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	(0)	а	•	utler (1965)
. PRC		Oyster	BCF	-	(O)	а		utler (1965)
DUCTS		Penaeus duorarum	L	-	0.42 (O)	а	Toxicant chemicals were evaluated in sea water at tempera-	utler (1965)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Shell SD-8448	Penaeus duorarum	L	-	0.28 (O)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
Shell SD-8448	Oyster <i>Leiostomus xanthurus</i> (juvenile)	BCF BSA	-	0.40 (O) (O)	а	The value reported is a 96-hr EC <sub>50</sub> (decreased shell growth). Water temperature was 19 C. Lost equilibrium at 1 ppm.	Butler (1965)
Shell SD-9129, EC	Rainbow trout Bluegill	BSA	-	4.90 (T4A) 4.0 (T4A)	-	The values reported are given as $LC_{50}$ .	Соре (1965)
Shell SD-9129	Oyster <i>Fundulus</i> similis (juvenile)	BCF BSA	_	(0) (0)	a a	No effect on exposure to the chemical at 1.0 ppm. Water temperature was 20 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Penaeus aztecus	L	-	0.32 (O)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
Silvex	Aquatic weeds in Georgia including Najas sp Potamogeton sp Myriophyllum heterophyllum Utricularia sp Myriophyllum brasiliense Eleocharis acicularis	FL	Farm ponds in Ga.	(0)	-	<ul> <li>Silvex in concentrations of 0.2 to 3.0 ppm killed 75 to 100 percent of the most prominent and damaging weeds.</li> <li>A slow kill is desirable because there is less chance of a fish kill due to an oxygen depletion resulting from weed decomposition.</li> <li>The results of 2 years' experimentation on control of aquatic weeds in Georgia farm ponds using Silvex indicated that this herbicide has a far wider range of satisfactory control than any other herbicide used in Georgia.</li> </ul>	Thomaston, et al (1959)
Silvex	Bluegill	FP	Okia.	1.5 to 3.0 (O)	-	This paper concerns lack of growth in weight and length of fish. A coefficient of condition $C(TL)$ was derived from fish lengths in inches and weights in grams, and is expressed as a ratio of 100,000 x weight in pounds to the cube of the length in inches. The $C(TL)$ of the fish is reported to be a result of the application of the herbicide over a 2-year period.	Houser (1962)
Silvex (Amchem)	Lepomis macrochirus	BSA	-	700 (T 18 hr) 600 (T 32 hr)	а	The experiment was conducted at 65 F. Fish were 2 in. long.	Cope (1963)
Silvex (K salt)	Lepomis macrochirus	BSA	-	83.0 (T2A) L 100.0 (T2A) G	<u>a</u> c d e g	Toxicity data for 24 and 48 hours are presented for liquid (L) and granular (G) formulations. Various commercial formulations were tested. The liquid formulations were almost invariably more toxic than the granular ones.	Hughes and Davis (1965)

`

COMMERCIAL CHEMICAL PRODUCTS

B-202

	Silvex	Filamentous algae Cladophora Spirogyra Hydrodictyon Submerged plants	FL	N. Y.	(O) (O) 2.0 (K)	ас	2 ppm caused 20% kill. 2 ppm caused 35% kill.	Cowell (1965)
		Chara Potamogeton Emergent plants Alisma			(O) 2.0 (K) (O)		2 ppm did not cause any kill. Complete decomposition occurred in about 3 weeks. 2 ppm did not cause any kill.	
		Sagittaria Floating plants Lemna Zooplankton			(O) 2.0 (K) (O)		2 ppm caused 20% kill. Complete decomposition occurred in about 3 weeks. Applications of 4 ppm produced significant reduction.	
	Silvex (Kuron)	Lepomis macrochirus Pimephales promelas	BSA	_	2.4 (T4A) 7.2 (T4A)	<u>a</u> c <u>e</u>	Bioassay method in Standard Methods for Examination of Water was used. TL <sub>m</sub> values for 24 and 48 hr are also presented.	Surber and Pickering (1962)
	Silvex (pelletized)	Lepomis macrochirus Pimephales promelas	BSA	-	(S) 14 (T4A) (H) 86 (T4A) (S) 13 (T4A) (H) 73 (T4A)	<u>a</u> c <u>e</u>	Comment same as above.	Surber and Pickering (1962)
<b>B-</b> 203	Silvex (butoxyethanol ester)	Lepomis macrochirus	BSA	_	1.2 (T <b>2</b> A)	a e	The various salts of the chemicals showed wide variations in toxicity.	Hughes and Davis (1963)
203	Silvex (isooctyl ester)	Lepomis macrochirus	BSA	_	3.7 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
	Silvex (potassium salt)	Lepomis macrochirus	BSA	_	8.3 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
	Silvex {propylene glycol butylether ester)	Lepomís macrochirus	BSA	_	16.6 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
COMMERCIAL CHEMICAL PRODUCT	Silvex	Benthic community of a farm pond	FL	Boone County, Mo.	2.8 & 4.6 (O)	C d	Many different aquatic plants, insects, molluses, and leeches are listed, 79 organisms in all. Data list populations in treated versus untreated pond areas as well as seasonal variations in numbers. The tests were conducted in a series of plastic enclosures 12 x 18 feet in area, and 4 feet deep. The most conspicuous change in the pond benthos in the enclosures treated with silvex was numerical increase at both treatment concentrations. Tendipedids, oligochaetes, <i>Chaoborus</i> , and libelludids, increased markedly. The densi- ties of damsel flies, leeches, and snails, were unaffected. <i>Chrysops</i> alone decreased. Other groups of organisms were not sufficiently numerous for analysis. The increases may have been caused by the enriching influence of decaying vegetation. The application rate of 2.8 ppm was within the recommended range of concentrations while 4.6 ppm was in excess of recommended rates.	Harp and Campbell (1964)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Silvex (triethyl- amine)	Lepomis macrochirus	BSA		20 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
CHEMICAL P	Silvex (triethy)- amine)	Lepomis macrochirus	BSA	-	16 (T1A)	<u>a</u> b e	Comment same as above.	Hughes and Davis (1967)
PRODUCTS	Silvex (polyglycol butyl ether ester)	Penaeus aztecus	L	_	0.28 (0)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
	Silvex	<i>Leiostomus xanthurus</i> (juvenile)	BSA	-	0.36 (O)	а	Water temperature was 16 C. The figure reported is a 48-hr $\mathrm{EC}_{50}.$	Butler (1965)
	Silvex	Crassostrea	BCFA & BSA	<b>\</b> -	1.0 (O, 20%)	-	Seawater was pumped continuously into test aquaria. Salinity, temperature, and plankton fluctuated with tide,	Butler (1965)
	(polyglycol butyl	virginica Penaeus			0.24 (O)		and ambient weather conditions. Some bioassays with fish	(1903)
	ether ester)	aztecus Leiostomus			0.36 (T2CFA)		were static. Toxicity was reported for the following: Oyster – 96-hr EC <sub>50</sub> – Conc. which decreased	
B-204		xanthurus Phytoplankton			78% (O)		shell growth. Shrimp – 48-hr EC <sub>50</sub> – Conc. which killed or paralyzed 50% of test animals. Fish – 48-hr EC <sub>50</sub> – Conc. which killed 50%. Phytoplankton – Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	
	Silvex	Salmo gairdneri	BSA	-	1.4 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses subacute mortality as well. Effects on repro-	Cope (1966)
		Lepomis			16.6 (T2A)		duction and behavior are also discussed. Data presented	(1300)
		macrochirus Pteronarcys			0.76 (T2A)		as EC <sub>50</sub> .	
		californicus Daphnia			2.40 (T2A)			
		pulex Simocephalus serrulatus			2.0 (T2A)			
	Silvex	Daphnia	BSA	-	100 (O)	acdiq	Toxicity, in terms of median immobilization concentration (IC <sub>50</sub> ), is presented for <i>Daphnia</i> ; median lethal concentra-	Crosby and Tucker
	(K salt)	magna Rainbow trout			21.9 (0)		tion (LC <sub>50</sub> ) for rainbow trout and bluegill are reported.	(1966)
		Bluegill	BSA		14.5 (O) 1.23 (T2A)		Data are given as LC50.	Pohene - t
	Silvex	Salmon Bluegili	BSA	-	0.60 (T2A)	_	Data are given as LU50.	Bohmont (1967)

	Silvex (potassium salt)	Lepomis macrochirus (eggs) L. cyanellus (eggs)	BSA	_	20 (NTE)	_	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest	Hiltibran (1967)
		Micropterus dolomieui (eggs)			20 (NTE)		concentration of chemical which did not cause death in 12 days (S).	
		Erimyzon sucetta (eggs)			20 (NTE)			
		(eggs) L. macrochirus (fry)			50 (S)			
	Silvex (ester)	Lepomis macrochirus (eggs)	BSA	-	10 (NTE)	-	Comment same as above.	Hiltibran (1967)
		L. cyanellus (eggs)			10 (NTE)			
		L. macrochirus (fry)			20 (S)			
	Silvex (ester)	Lepomis macrochirus (eggs)	L	-	2.4/2 (O) 1.0 (NTE)	_	Comment same as above.	Hiltibran (1967)
 		L. cyanellus (eggs)			2.4/4 (O)			-
1		Micropterus dolomieui			1.0/4 (O)			
		Erimyzon sucetta			2.4 (NTE) 1.0 (NTE)			
		<i>L. macrochirus</i> (fry)			2.0 (S)			
	Silvex (potassium	Lepomis macrochirus	BSA	_	30 (S)	-	Comment same as above.	Hiltibran (1967)
	salt)	L. cyanellus	564		10 (NTE)		0	
ი	Silvex (sodium salt)	Lepomis macrochirus (fry)	BSA	_	50 (S)	_	Comment same as above.	Hiltibran (1967)
COMMER	Silvex	Pteronarcys californica (naiads)	BSA	_	0.00034 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
	Simazen (granular)	Althernanthera philoxeroides	FL	Fla.	(O)	-	At 10.0 lb/acre, alligator weed was not affected.	Copeland and Woods (1959)
CHEMICAL PRODUC	Simazine	<i>Nympheae</i> sp <i>Leersia</i> sp <i>Paspalum</i> sp <i>Juncus</i> sp	FL	Farm ponds in Ga.	(O)	_	Although Nympheae sp was killed at a rate of 50 lb/acre, no epinastic effects were noted. The chemical did not translocate and only killed the tops. Treatments on Leersia sp, Paspalum sp, and Juncus sp were unsuccessful and gave no encouraging results. Limited use of simazine has not proven it to be a satisfactory aquatic herbicide in Georgia.	Thomaston, et al (1959)

COMMERCI	nemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Sima	azine	Onchory nchus tshawy tscha	BSA & CF	_	7.0 (T1A) 6.6 (T2A)	a c d e	Concentrations were based on percent active ingredient. Median tolerance limits for 72 and 96 hours estimated from the constant flow experiment were 7.2 ppm and 6.5 ppm, respectively, for this species.	Bond, et al (1960)
Sima (her 196	rboxy-	Phoxinus phoxinus	BSA	-	(0)	<u>a</u> c d e	Two series of aquarium aerated tests were performed, one without plants ( <i>Callitriche</i> , and <i>Elodea</i> ). 20% kill occurred in 3 days. The highest nonlethal concentration was 5 ppm.	Vivier and Nisbet (1965)
Sima (her 196	rboxy-	Phoxinus phoxinus	BSA	-	1.25 (K2A) 1.5 (T2A)	<u>a</u> c d e	Two series of aquarium tests were performed, with and without plants, which lowered the toxicity. 90% kill occurred at 5 ppm in 6 hr.	Vivier and Nisbet (1965)
Sima	izine	Micropterus salmoides (fry) Ictalurus punctatus	BSA	-	25.0 (SB3) 10.0 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
		(fry) Lepomis macrochirus (fry)			10.0 (SB3)			
Sima	zine	Phytoplankton Hydrodictyon reticulatum Zygnema spp	FL	Ala.	2.0 (K1) 2.0 K1)	а	In a series of bass spawnings and rearing ponds, Simazine was used at concentrations of 0.5, 1.0, and 2.0 ppm to control light to medium growths of phytoplankton which interferred with bass production. Success was uniform with control last-	Snow (1963)
K		etc. Zooplank ton Fish			(0)		ing for as long as 85 days. No fish kills occurred and the chemical was apparently not toxic to zooplankton.	
		Micropterus salmoides Lepomis			(O) (O)			
Sima (WP		<i>cyanellus</i> Bluegill Rainbow trout	BSA	-	118 (T4A) 56 (T4A)	а	This is an estimated LC $_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
Sima (WP		Lepomis macrochirus (eggs)	L	-	10/4 (O)	_	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival.	Hiltibran (1967)
		L. cyanellus (eggs)			10/5 (O)		Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration	
		Micropterus dolomieui			10/3 (O)		of chemical which did not cause death in 12 days (S).	
		(eggs) Erimyzon sucetta			10 (NTE)			
		(eggs) L. macrochirus (fry)			0.3 (S)			

	Simazine	Rainbow trout Bluegill	BSA	-	56.0 (T2A) 118.0 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
	Simazine	Lepomis macrochirus (eggs) L. cyanellus (eggs) Micropterus dolomieui (eggs)	L	-	10 (NTE) 10/7 (O) 10 (NTE)	-	Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S)?	Hiltibran (1967)
		Erimyzon sucetta (eggs)			10 (NTE) 10 (S)			
	Sinox	Richardsonius balteatus hydroflox	BSA	-	0.16 (T1A) 0.14 (T2A) 0.13 (T4A)	a c d e f	Results given were in soft water. Results in hard water were as follows: 0.24 (T1A), 0.24 (T2A), and 0.24 (T4A).	Webb (1961)
	Slickgone 1	Pandalus montagni Crangon crangon	BSA	-	5.2 (T2A) 6.6 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C. It was shown that the toxicity of this solvent emulsifier decreased with time due to evaporation of the solvent.	Portmann and Connor (1968)
B-207		Cardium edule Carcinus malmas			32.4 (T2A) 35.0 (T2A)			
7	Slickgone 2	Pandalus montagni Crangon crangon Cardium	BSA	_	4.5 (T2A) 3.5 (T2A) 30.5 (T2A)	<u>a</u> e	Comment same as above.	Portmann and Connor (1968)
		edule Carcinus maenas			21.3 (T2A)			
COMMERCIAL	Slix	Pandalus montagni Crangon carcinus maenas Cardium edule	BSA	-	12.1 (T2A) 114.5 (T2A) 150.0 (T2A) 12.7 (T2A)	<u>a</u> e	Experiments were conducted in tanks holding 10 liters of seawater at 15 C. It was shown that the toxicity of this solvent emulsifier decreased with time due to evaporation of the solvent. Slix at a concentration of 10 ppm killed 100% of <i>Crangon</i> <i>crangon</i> larvae in 3 hr; at 33.3 ppm it killed 70% of <i>Carcinus maenas</i> larvae in 3 hr.	Portmann and Connor (1968)
		Pimephales promelas (juveniles)	BSA	-	(S) 34-39 (T1-4A) (H) 1,470-1,530 (T1-4A)	<u>acdf</u>	Syndets and soaps were of nearly equal toxicity in soft water (S) but syndets were approximately 40X more toxic than soap in hard water (H).	Henderson, et al (1960)
ICAL PE	Sovicide tetra aminol	Phoxinus phoxinus	BSA	-	8 (100%K)	<u>a</u> c d e	The highest concentration nonlethal in 6 hr was 4 ppm.	Vivier and Nisbet (1965)
CHEMICAL PRODUCTS	Stam F-34, tech.	Salmo gairdnerii	BSA	-	4,000 (T2A)	а	The experiment was conducted at 55 F. Fish were 2-3 in. long.	Соре (1963)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Stauffer N2790	Leiostomus xanthurus (juvenile)	BSA	_	0.24 (O)	а	Water temperature was 24 C. The figure reported is a 48-hr EC50.	Butler (1965)
	Oyster	BCF		0.33 (O)		The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
	Penaceus aztecus	L		0.0024 (O)		Toxicant chemicals were evaluated in sea water at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
Stauffer N-2790 (tech)	Raínbow trout Bluegill	BSA	_	0.019 (T4A) 0.0062 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
Stauffer R-1910	Penaeus aztecus	L	-	(0)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
	<i>Leiostomus xanthurus</i> (juvenile)	BSA	_	(O)	а	Water temperature was 25 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Oyster	BCF				No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
Stauffer (R-1910, tech)	Rainbow trout Bluegill	BSA	_	3.6 (T4A) 5.5 (T4A)	а	This is an estimated LC $_{\!$	Cope (1965)
Stauffer R-1910	Crassostrea virginica Penaeus setiferus Leiostomus xanthurus Fundulus similis Mugil cephalus Cyprinodon variegatus Phytoplankton	BCFA	_	1.0 (NTE, all species)	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)
Stauffar R-4461	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	0.45 (O) 1.0 (0, 10%) 0.32 (T4CFA) NTE	-	Comment same as above.	Butler (1965)
Stauffer R-4461	Oyster	BCF	-	0.45 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)

	Stauffer R-4461	Penaeus aztecus	L	-	(O)	a	Toxicant chemicals were evaluated in seawater at temper- atures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
	Stauffer R-4461 (tech)	Rainbow trout Bluegill	BSA	_	0.72 (T4A) 0.81 (T4A)	а	This is an estimated LC $_{\rm 50}$ value at temperatures from 55 to 75 F.	Cope (1965)
	Stauffer R-4461	Leiostomus xanthurus (juvenile)	BSA	-	0.32 (O)	а	Water temperature was 25 C. The figure reported is a 48-hr EC <sub>50</sub> .	Butler (1965)
	Stauffer R-5092	Leiostomus xanthurus (juvenile)	BSA	-	0.02 (O)	а	Water temperature was 26 C. The figure reported is a 48-hr EC <sub>50</sub> .	Butler (1965)
		Oyster	BCF	-	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
		Penaeus aztecus	L	-	0.0032 (O)	а	Toxicant chemicals were evaluated in seawater at temper- atures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality.	(1965) (1965)
	Steramine	Chlorella pyrenoidosa	L	-	20 (AC<1/2	_	Describes a bioassay method to differentiate between an algicide (AC) and an algistat (AS). The treated culture was subcultured as time progressed. Allen's medium was used.	Fitzgerald and Faust (1963)
000 H	Streptomycin sulfate	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T Ma - T So $- T$ Cv - NT Gp - NT Np - T (21)	Palmer and Maloney (1955)
COMMERCIAL CHEMICAL PRODUC	Strobane	Blue crab Marsh fiddler crab Red-jointed fiddler crab Cyprinodon variegatus Mugil curema Leiostomus xanthurus	FE	Bombay Hook Island, Del.	(O) (O) (O) (O) (O)	-	<ul> <li>Strobane was applied at the rate of 0.3 pound per acre.</li> <li>The location under study was a salt marsh bounded by Delaware Bay.</li> <li>Organisms were confined in cages within the test area.</li> <li><i>c. variegatus, M. curema</i>, and <i>L. xanthurus</i> showed 16 percent mortality in 7 days.</li> <li>Blue crabs showed 27 percent mortality when exposed for 7 days in streams and 20 percent mortality in ponds.</li> <li>Marsh fiddler crabs and red-jointed fiddler crabs showed mortalities of 68 and 20 percent, respectively, when exposed for 7 days.</li> </ul>	George, et al (1957)
ICAL PR	Strobane + methyl parathion	Oyster	BCF	-	0.026 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
ODUC	Strobane	Pteronarcys sp (nymphs)	BSA		0.0005 (T4A)	<u>a</u>	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Соре (1965)

Chemical Strobane (tech) Strobane Strobane Strobane	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Strobane	Oyster	BCF	_	0.02-0.059 (O)	а	The value reported is a 96-hr $EC_{50}$ (decreased shell growth).	Butler (1965)
Strobane (tech)	Bluegill	BSA	-	0.0084 (T4A)	-	The values reported are given as $LC_{\overline{50}}.$	Cope (1965)
Strobane	Pteronarcys californica (naiads)	BSA	_	0.0005 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
Sulfotepp	Channet catfish (fingerlings)	BSA	-	<1.0 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1969)
Swep	Lepomis mecrochirus	BSA	-	6.0 (T1A)	abe ~	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
"Synthetic detergent"	Sludge worms	BSA	-	23 (T4A)	<u>a</u> cdi	Data using hard and soft water are presented as well as information on the effect of temperature. Additional TL <sub>m</sub> data are presented.	Wurtz and Bridges (1961)
Systox	Pimephales promelas	BSA		3.9 (T4A)	acdef	Tests were performed in both hard and soft water. Additional tolerance limit values are given.	Henderson and Pickering (1958)
Systox	Fathead minnow	BSA	_	4.2 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish" It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds This experiment was performed in hard water.	Tarzweli (1959)
Systox	Pimephales promelas	BSA	_	3.6 (T4A)	<u>a</u> d e f	Concentrations were based on percent active ingredient.	Henderson, et al (1959)
Systox	Pimephales promelas	BSA	_	3.6 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)
Systox (tech,	Pimephales promeles	BSA		2.9 (T4A)	<u>a</u> cd <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent	Pickering, et al (1962)
92 percent)	Lepomis macrochirus			0.11 (T4A)		or carrier in most cases.	(1302)
	Carassius auratus			12 (T4A)			
	Lebistes reticulatus			0.66 (T4A)			

COMMERCIAL CHEMICAL PRODUCTS

B-210

	2,4,5-T	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - NT Ma - T (3) So - NT Cv - NT Gp - NT Np - NT	Palmer and Maloney (1955)
	2,4,5-T (pellets)	Althernanthera philoxeroides Najas quadalupensis Spatterdock	FL	Fla.	(O)	-	The degree of control was as follows: A. philoxeroides (20 lb/acre) — 95 percent N. quadalupensis (24 lb/acre) — none Spatterdock (21.8 lb/acre) — 3 percent.	Copeland and Woods (1959)
	2,4,5-T	Althernanthera philoxeroides	FL	Fla.	(O)	-	At 0.5 lb/acre, only 1-2 percent control of alligator weed was obtained.	Copeland and Woods (1959)
B-21	2,4,5-T (dimethyl- amine ester)	Lepomis macrochirus	BSA	-	144 (T2A)	ae	The various salts of the chemicals showed wide variations in toxicity.	Hughes and Davis (1963)
211	2,4,5-T (butoxy- ethanol ester)	Lepomis macrochirus	BSA	-	1.4 (T2A)	ae	Comment same as above.	Hughes and Davis (1963)
	2,4,5-T (isooctyl ester)	Lepomis macrochirus	BSA	-	26 (T2A)	ae	Comment same as above.	Hughes and Davis (1963)
	2,4,5-T (propylene glycol butyl ether ester)	Lepomis macrochirus	BSA	-	17 (T2A)	a e	Comment same as above.	Hughes and Davis (1963)
COMMERCIAL	2,4,5-T (oleic-1,3- propylene diamine)	Lepomis macrochirus	BSA	-	2.9 (T1A)	-	The bioassay methods employed in this experiment were not given in the paper but it was stated that the same procedures were employed as in previous work.	Davis and Hughes (1963)
	2,4,5-T (isopropyl ester)	Lepomis macrochirus	BSA	-	1.8 (T1A)	-	Comment same as above.	Davis and Hughes (1963)
MICAL	2,4,5-T (triethyl amine)	Lepomis macrochirus	BSA	_	53.7 (T1A)		Comment same as above.	Davis and Hughes (1963)
CHEMICAL PRODUCTS	2,4,5-T	Lepomis macrochirus	BSA	-	11.0 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)

2,4,6-T (polyglycol butyl ether ester) 2,4,5-T 2,4,5-T (polyglycol butyl ether ester)	Oyster Leiostomus xanthurus (juvenile) Penaeus aztecus Penaeus	BCF BSA L	-	0.14 (O) 0.32 (O)	a	The value reported is a 96-hr EC <sub>50</sub> (decreased shell growth). Water temperature was 16 C. The figure reported is a	Butler (1965) Butler
2,4,5-T (polyglycol butyl ether	xanthurus (juvenile) Penaeus aztecus Penaeus		-		а		
(polyglycol butyl ether	aztecus Penaeus	L	-	1-1		48-hr EC <sub>50</sub> .	(1965)
				(0)	a	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality.	Butler (1965)
2,4,6-T (acid)	aztocus	L	-	(O)	а	Toxicant chemicals were evaluated in seawater at tempera- atures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
2, <b>4</b> ,5-T	Crassostrea	BCFA &	-	0.14 (O)	-	Seawater was pumped continuously into test aquaria.	Butler (1965)
(polyglycol butyl ether	virginica Panaaus	BSA		1.0 (O,20%)		Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with	(1965)
ester)	aztecus Leiostomus xenthurus			0.32 (T2CFA)		fish were static. Toxicity was reported for the following: Oyster – 96-hr EC <sub>50</sub> – Conc. which	
	<i>xanthurus</i> Phytoplankton			89% (O)		decreased shell growth. Shrimp – 48-hr EC <sub>50</sub> – Conc. which killed or paralyzed 50% of test animals. Fish – 48-hr EC <sub>50</sub> – Conc. which killed 50%. Phytoplankton – Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	
2,4,5-T (acid)	Crassotrea virginica	BCFA & BSA	-	20 (NTE)	-	Comment same as above.	Butler (1965)
(8010)	Penaeus	20/1		1.0 (NTE)			(1000)
	aztecus Mugil			50.0 (NTE)			
	<i>caphalus</i> Phytoplankton						
,4,5-T	Crassostrea virginíca	BCFA & BSA		0.14 (O)	_	Comment same as above.	Butler (1965)
(polyglycol butyl ether	Penaeus	DOA		1.0 (0, 20%)			(1500)
ester	aztecus Leiostomus			0.32 (T2CFA)			
	<i>xanthurus</i> Phytopiankton			89% (O)			
2,4,5-T (ester)	Lepomis macrochirus	L	-	10 (NTE)		Fertilized fish eggs of indicated species were placed in 1 liter of test solution and allowed to hatch. Toxicity data are presented as concentration in ppm/number of days queries	Hiltibran (1967)
	(eggs) L. cyanellus L. mecrochirus (fry)			10 (NTE) 10 (S)		presented as concentration in ppm/number of days survival. Maximum length of test was 8 days. No food was added. Small bluegill were tested to find the highest concentration of chemical which did not cause death in 12 days (S).	

1

	2,4,5-T (ester)	Lepomis macrochirus L. cyanellus	L	-	1.0 (NTE) 4/1 (O), 1.0 (NTE)	-	Comment same as above.	Hiltibran (1967)
		Micropterus dolomieui L. macrochirus (fry)			4/0 (O), 1.0 (NTE) 1.0 (S)			
	2,4,5-T (sodium salt)	Lepomis macrochirus (fry)	L	-	50 (S)	-	Comment same as above.	Hiltibran (1967)
	TAG 10%	Channel catfish (fingerlings)	BSA		1.5 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	2,3,5-TBA	Lepomis macrochirus Micropterus salmoides	BSA	-	90 (T2A) 55 (T2A)	aco	The response of bluegill and bass fingerlings to nine agricul- tural chemicals as determined by bioassay using river water is presented in this report. Bluegills were more tolerant of the chemicals tested than bass.	Davis and Hardcastle (1959)
	2,3,6-TBA	Lepomis macrochirus Micropterus salmoides	BSA	_	1750 (T2A) 1250 (T2A)	<u>a</u> co	Comment same as above.	Davis and Hardcastle (1959)
	TCA 90%	Channe) catfish (fingerlings)	BSA	-	>2000 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Sodium TCA	Onchorynchus tshawytscha	BSA	_	870 (NTE)	acde	Concentrations were based on percent active ingredient.	Bond, et al (1960)
	Sodium TCA	Mugil cephalus (juvenile)	BSA	-	(0)	а	Water temperature was 28 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Sodium TCA	Penaeus aztecus	L	-	(O)	a	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
COM	TD 47	Micropterus salmoides (fry)	BSA	-	0.075 (SB3)	acdefp	At least 90 percent of the fry survived for a period of 72 hours at the concentration listed.	Jones (1965)
MERCI		lctalurus punctatus (fry)			0.2 (SB3)			
COMMERCIAL CHEMI		Lepomis macrochirus (fry)			0.2 (SB3)			

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
TD-47	Carp- goldfish hybrid Notropis umbratilis N. lutrensis Pimephales notatus	BSA	_	175 (T4A) 95 (T4A) 105 (T4A) 120 (T4A)	а	In addition to the median tolerance limits, this report also has data on the residue of the chemical in the fish, some of the physiological effects, and degradation curves for the chemical in water.	Walker (1963)
	lctaluris natalis I. melas Lepomis macrochirus L. microlophus Micropteris salmoides			175 (T4A) 180 (T4A) 125 (T4A) 125 (T4A) 120 (T4A)			
TD-72 (EC6)	Gambusia affinis	FL	Ponds in III.	(O)	-	When applied at 0.5 pounds per acre active ingredient, 18 per- cent mortality occurred in 1 day.	Mulla (1963)
TD-282 [di(N,N- dimethyltri- decylamine) salt of Endothall]	Australorbis glabratus	BSA & FL	Puerto Rico	Variable (O)	c	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
TD-283 (mono-N,N- dimethyltri- decylamine) salt of Endothall)	Australorbis glabratus	BSA & FL	Puerto Rico	3.8-6.2 (O)	c	Comment same as above.	Seiffer and Schoof (1967)
TD-440	Lepomis macrochirus	BSA	-	3.0 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
TD-497	Lepomis macrochirus	BSA	_	4.0 (T1A)	<u>a</u> be	Comment same as above.	Hughes and Davis (1967)
Felodrin	Leiostomus xanthurus (juvenile)	BSA	-	0.0003 (O)	а	Water temperature was 13 C. The figure reported is a 48-hr $EC_{50}$ .	Butler (1965)
Telodrin	Leiostomus xanthurus (juvenile)	ВСН	-	(0)	а	A concentration of 0.000025 ppm will kill in 10 days. The fish were able to survive for 5 months in a concentration of 0.00001 ppm.	Butler (1965)
ТЕРР	Fish	BSA	-	0.25 (K)	-	A concentration of 0.25 ppm was lethal in aquarium tests.	Linduska and Surber (1948)

ΤΕΡΡ	Fathead mìnnow	BSA	-	1.0 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in the paper) report widely different values for the same compounds This experiment was performed in hard water.	Tarzwell (1959)	
ТЕРР	Pimephales promelas Lepomis	BSA	-	1.7 (T4A) 0.84 (T4A)	<u>acdef</u>	Tests were performed in both hard and soft water. Additional tolerance limit values are given.	Henderson and Pickering (1958)	
ТЕРР	macrochirus Pimephales promelas	BSA	-	1.7 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ signifi- cantly in different streams.	Tarzwell (1959)	
ТЕРР	Channel catfish (fingerlings)	BSA	-	2.3 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)	APP
TEPP	Pimephales promelas	BSA	-	1.7 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1960)	PENDIX
ТЕРР	Protococcus sp Chlorella sp Dunaliella sp Phaeodactylum tricornutum Monochrysis lutheri	L	-	500 (NG) 500 (NG) 500 (K) 500 (K) 500 (K)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were expressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.		DIX B
TEPP (tech, 40 percent)	Pimephales promelas Lepomis macrochirus Carassius auratus Lebistes reticulatus	BSA	-	2.1 (T4A) 1.3 (T4A) 21 (T4A) 1.8 (T4A)	<u>a</u> c d <u>e</u>	Soft water primarily was the test medium. TL <sub>m</sub> 's reported for 24, 48, and 96 hr. Acetone or alcohol used as solvent or carrier in most cases.	Pickering, et al (1962)	
Terramycin	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (O)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - PT (7) Ma - T So - NT Cv - NT Gp - NT Np - T (3) PT (7)	Palmer and Maloney (1955)	

COMMERCIAL CHEMICAL PRODUCTS

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Fhanite (isobornyl thiocyano- acetate)	Green sunfish	BSA	_	1.0 (K 6 hr) 0.5 (K 6 hr)	aep	The main purpose of this experiment was to determine the repellent characteristics of certain chemicals. The experiments were conducted in a wooden trough.	Summerfelt and Lewis (1967)
etrachloro- phene	Cylindrospermum licheniforme (Cl) Microcystis aeruginosa (Ma) Scenedesmus obliquus (So) Chlorella variegata (Cv) Gomphonema parvulum (Gp) Nitzschia palea (Np)	L	-	2.0 (0)	<u>a</u>	Observations were made on the 3rd, 7th, 14th, and 21st days to give the following (T = toxic, NT = nontoxic, PT = par- tially toxic with number of days in parentheses. No number indicates observation is for entire test period of 21 days): CI - T (3) PT (7) Ma - NT So - PT (7) Cv - PT (3) Gp - NT Np - T (3)	Paimer and Maioney (1955)
	Petromyzon marinus Micropterus salmoides Micropterus dolomieui Lepomis macrochirus Stizostedion v. vitreum Perca flavescens	BSA		3-10 (O) 22-42 (O) 34.5-42 (O) 21.5-44.0 (O) 5.75-11.5 (O) 7.25-20.5 (O)	a c d <u>e</u>	Three types of dilution water used with hardness values of 95.4, 141.7, and 203.3 ppm. As a lamprey larvicide, 3-10 ppm required. Toxicity range (ppm) given as that which kills 25 percent of the test fish.	Applegate and King (1962)
	lctalurus natalis Catostomus commersoni Notropis			5.75-15.5 (O) 5.0-13.0 (O) 13.25-28.0 (O)			
	heterolepis Notemigonus crysoleucas			14.75-33.0 (0)			
	Pimephales promelas Salmo gairdnerii			16.0-35.5 (O) 12.0-25.25 (O)			
FM	Lamprey (larvae) Rainbow trout (fingerlings)	BFR+L	Great Lakes (Mich)	1.5 (K 8 hr) 2.0 (K 2 hr) 3.0 (K 19 hr) 5.0 (K 3 hr)	а	Describes a portable field monitor, using water obtained at the site.	Howell and Marquette (1963)

	TFM	Hydra Turbellarians Eripidelidae Burrowing mayflies Black flies Clams Sea lamprey Rainbow trout	BSA (L)	-	3.0 (K1A) 4.0-8.0 (K1A) 12.0 (K1A) 12.0 (K1A) 8.0 (K1A) 16.0 (K1A) 4.0 (K1A) 13.0 (O)	<u>a e</u>	All numbers cited are for 100% kill in 22-24 hours. The number given for rainbow trout was for a 60% kill in 22-24 hours. Data were given for fourteen other aquatic species, but they are not included here because of very low toxicity or doubtfu data. These included leeches, isopods, scuds, crayfish, stone- flies, dragonflies, waterbugs, water boatmen, mayflies, caddis flies, bloodworms, snipe flies, and snails.	
	Thanite	Gambusia affinis	BSA	-	0.8 (L1)* 0.9 (L1)** *Resistant fish **Nonresistant fish	a	This paper deals with the resistance of mosquito fish to chlorinated hydrocarbon compounds. Resistant fish were not always less sensitive to these chemicals.	Boyd and Ferguson (1964)
	Thimet	Gambusia affinis	BSA	-	0.05 (K 83%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)
B-217	Thimet	Pimephales promelas Lepomis macrochirus	BSA	-	0.25 (T4A) 0.0047 (T4A)	a c d e f	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson (1966)
7	Thimet	Leiostomus xanthurus Cyprinodon variegatus Mugil cephalus	BCFCH	_	0.0005 (O) 0.0005 (O) 0.0005 (O)	а	At a concentration of 0.0005 ppm, the following percent acetylcholinesterase activity as compared to controls was found: <i>L. xanthurus</i> – 84 <i>C. variegatus</i> – 68 <i>M. cephalus</i> – 69.	Butler and Johnson (1967)
8	Thiodan <sup>®</sup> I EC2	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a ç	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in lb/acre, and resulted in a 100 percent mortality for the fish, and a 100 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
COMMERCIAL	Thiodan <sup>®</sup> II EC2	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a C	Comment same as above.	Mulla (1963)
CHEMICA	Thiodan (tech, 96.6 percent)	Pimephales promelas Lebistes reticulatus	BSA	-	0.0033 (T4A) 0.0037 (T4A)	a c d e f	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson (1966)
CHEMICAL PRODUCTS	Thiram	Channel catfish (fingerlings)	BSA	-	>1.0 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Tiguron	Salmo gairdneri Salmo trutta Salvelinus fontinalis Salvelinus namaycush Ictalurus punctatus Lepomis macrochirus	BSA	_	4.35 (T2A) 3.62 (T2A) 5.50 (T2A) 5.30 (T2A) 5.90 (T2A) 8.90 (T2A)	<u>a</u> f	Variance and the 95 percent confidence interval (C.I.) were also determined.	Willford (1966)
Tillam	Crassostrea virginica Penaeus duorarum Penaeus setiferus Leiostomus xanthurus Fundulus similis Mugil cephalus Cyprinodon variegatus Phytoplankton	BCFA & BSA	-	1.0 (O, 20%) 1.0 (NTE) 6.3 (T2A) 24% (O)	_	Seawater was pumped continuously into test aquaria.         Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:         Oyster –       96-hr EC <sub>50</sub> - Conc. which decreased shell growth.         Shrimp –       48-hr EC <sub>50</sub> - Conc. which killed or paralyzed 50% of test animals.         Fish –       48-hr EC <sub>50</sub> - Conc. which killed 50%.         Phytoplankton → Percent decrease of CO <sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	Butler (1965)
TNT	Lythrurus umbratilis Hyborhynchus notatus Cyprinella whippli Helioperca incisor Gambusia affinis Cristivomer n. namaycush Ericymba buccata Cyprinus carpio Ameiurus melas Moxostoma aureolum	BSA	_	(0)	acfio	All sensitive and young stages of fish died in concentrations of TNT red liquor waste greater than approximately 1 to 600 dilution of average samples as described by Mohlman (17 to 18 C). Increase in water temperature decreased survival time of the fish in TNT waste and smaller speci- mens died sooner than larger.	Degani (1943)

		Chaenobryttus gulosus Lepomis cyanellus						
	Tordon	Lepomis macrochirus	BSA	-	43 (T1A)	<u>a</u> be	This report is a simple and straightforward determination of a median tolerance limit for a selected group of herbicides.	Hughes and Davis (1967)
	Torden	Crassostrea virginica Penaeus aztecus Mugil cephalus Phytoplankton	BCFA & BSA	_	1.0 (NTE) 1.0 (NTE) 1.0 (NTE) (O)	~	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Butler (1965)
	Tordon 101	Penaeus aztecus	L	-	(0)	а	concentration. Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC50	Butler (1965)
В		aricos					or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	(1000)
B-219	Tordon 101	Oyster	BCF		(0)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
	Tordon	<i>Mugil cephalus</i> (juvenile)	BSA	~	(O)	а	Water temperature was 28 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Tordon	Rainbow trout Bluegill	BSA	~	2.4 (T2A) 13.1 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
CON	Toxaphene	Lepomis macrochirus Notemigonus crysoleucas Micropterus salmoides	FL	Auburn, Ala.	(O)	_	0.02 ppm killed bluegills and golden shiners. The bass were killed at 0.04 ppm. 0.2 ppm in an earthen pond killed bluegill and bass fingerlings and bait-sized goldfish in 45 hours.	Lawrence (1950)
COMMERCIAL CI	Toxaphene	Carp <i>Perca flavescens</i> Golden shiners	FL	Beckers Lake, Ariz.	0.1 (K)	a b c g	All fish died during an eleven-day period. The lake was successfully stocked about 8 months later with rainbow trout.	Hemphill (1954)
CHEMICAL PRODUCTS	Toxaphene	Carp Gila robusta elegans Largemouth bass Bluegill Brown trout Bullhead catfish	FL	Lyman Reser- voir, Ariz.	0.1 (K)	a b c g	All fish died during a two-day period. The reservoir was successfully stocked 10 months later with rainbow trout.	Hemphill (1954)

COMMERCIAL	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL PRODUCTS		Pimephales promelas	BSA	-	0.036 (T1A) 0.020 (T1A) 0.0057 (T1A) 0.1 (O) 0.060 (T1A) 0.10 (T1A) 1.5 (T1A) 9.5 (T1A)	a c d f	<ul> <li>At 50 F and 6 ppm methyl orange alkalinity.</li> <li>At 50 F and 212 ppm methyl orange alkalinity.</li> <li>At 75 F and 212 ppm methyl orange alkalinity.</li> <li>At 212 ppm methyl orange alkalinity, 75 F, no aeration,</li> <li>6.4-7.0 ppm dissolved O<sub>2</sub> and with light, 100% mortality occurred in 2 days. Toxicant was added immediately before fish.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>At 55 F and 212 ppm methyl orange alkalinity.</li> <li>The chemical becomes detoxified when left standing in water by removal by microorganisms. The chemical is more toxic to fish in hard water than in soft water.</li> <li>0.05 ppm emulsified toxaphene is sufficient for fish eradication. Somewhat lower concentrations can be used in shallow hard water lakes with a higher temperature.</li> </ul>	Hooper and Grzenda (1955)
	Toxaphene	Tendipedae Chaoborus spp	FL	Colo.	0.1 (K3) (O)	a c d e	This is a study of lake bottom fauna. Repopulation of lake was not complete until nine months later.	Cushing and Olive (1957)
B-220	Toxaphene	Bluegill Pumpkinseed Largemouth bass Yellow perch Rock bass	FL	Lakes (Mich.)	0.005 (K)	d	Toxaphene at the indicated concentration killed the majority of small fish while larger fish were not killed. According to the authors, toxaphene at a 5 ppb concentration can be used to reduce the population of small fish without greatly affecting the population of large fish.	Fukano and Hooper (1958)
	Toxaphene	Bullhead Bullhead Carp Bottom fauna	BSA F	_ Iowa	<0.001 (T1A) 0.005 (O) 0.005 (O) -	a —	It was estimated that 25.0 ppb will eradicate an entire fish population in a lake. A concentration of 20.0 ppb seemed sufficient to kill all fish in aquarium tests. In highly turbid water, 200 ppb were required. This suggests that the silt in suspension has a detoxifying effect. The field study reports what is believed to be the first in- stance in which a lake is rid of all fish by chemical means. Bottom fauna declined in volume due to the treatment but recovered rapidly (1 mo).	Rose (1958)
	Toxaphene	Catostomus macrochailus Ptychochailus oregonense Cyprinus carpio Richardsonius baltaatus Mylochailus caurinum	FL	Spectacle Lake, British Columbie	0.07 (O) 0.07 (O) 0.07 (O) 0.07 (O) 0.07 (O)	a c d	<ul> <li>Experiments were conducted in 8 lakes in British Columbia, all of which were alkaline. These lakes had no permanent inlet or outlet streams.</li> <li>Initial results or symptoms were observed in 120 hr. All caged fish were dead except 2 carp which managed to survive for 1 to 2 months.</li> </ul>	Stringer and McMynn (1958)

Toxaphene	Catostomus macrocheilus Ptychocheilus oregonense Cyprinus carpio	FL	Lady King Lake, British Columbia	0.10 (K2)	a c d	Experiments were conducted in 8 lakes in British Columbia, all of which were alkaline. These lakes had no permanent inlet or outlet streams. There was no sign of fish life after 24 hr.	Stringer and McMynn (1958)
	Richardsonius balteatus Perca			0.10 (K2)			
	flavescens Mylocheilus caurinum Couesius plumbeus Coregonus williamsoni Oncorhynhus			0.10 (K2)			
	nerka Salmo			0,10 (K2)			
	gairdneri Lottus asper			0.10 (K2)			
Toxaphene	Catostomus	FL	Gallagher	0.07 (K2)	a c d	Experiments were conducted in 8 lakes in British Columbia,	Stringer and McMynn
	macrocheilus Ptychocheilus		Lake, British	0.07 (K2)		all of which were alkaline. These lakes had no permanent inlet or outlet streams.	(1958)
	oregonense Richardsonius		Columbia	0.07 (K2)			
	balteatus Perca			0.07 (K2)			
	flavescens Mylocheilus			0.07 (K2)			
	caurinum Couesius			0.07 (K2)			
	plumbeus Coregonus			0.07 (K2)			
	williamsoni Oncorhynhus			0.07 (K2)			
	nerka Salmo gairdneri			0.07 (K2)			
Toxaphene	Catostomus	FL	Gladstone	0.03 (O)	acd	Experiments were conducted in 8 lakes in British Columbia,	Stringer and
	macrocheilus Cyprinus		Lake, British	0.03 (O)		all of which were alkaline. These lakes had no permanent inlet or outlet streams.	МсМупл (1958)
£	carpio Richardsonius		Columbia	0.03 (O)		In 12 hours many trout and shiners were dead; all other fish showed signs of distress.	
	balteatus Mylocheilus			0.03 (O)			
Toxaphene Toxaphene CIAL CHEMICAL PB000	caurinum Salmo gairdneri			0.03 (O)			

Chemical Toxaphene	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Toxaphene	Catostomus macrocheilus	FL	Taylor Lake, British	0.01 (K4) 0.01 (K4)	a c d	Experiments were conducted in 8 lakes in British Columbia, all of which were alkaline. These lakes had no permanent inlet or outlet streams.	Stringer and McMynn (1958)
	Ptychocheilus oregonense		Columbia			Rainbow trout and shiners showed definite signs of distress	(1000)
	Cyprinus carpio			0.01 (K4)		in 4 days.	
	Richardsonius balteatus			0.01 (O)			
	Mylocheilus caurinum			0.01 (K4)			
	Salmo gairdneri			0.01 (O)			
Toxaphene	Catostomus macrocheilus	FL	Alleyne Lake,	0.01 (0)	acd	Experiments were conducted in 8 lakes in British Columbia, all of which were alkaline. These lakes had no permanent	Stringer and McMynn
	Salmo gairdneri		British Columbia	0.01 (O)		inlet or outlet streams. In 24-48 hours many fish were dead while others were still in distress.	(1958)
Toxaphene	Cyprinus carpio	FL	Round Lake, British Columbia	0.03 (K3)	a c d	Experiments were conducted in 8 lakes in British Columbia, all of which were alkaline. These lakes had no permanent inlet or outlet streams.	Stringer and McMynn (1958)
Toxaphene	Catostomus macrocheilus	FL	Summit Lake,	0.10 (O)	acol	Experiments were conducted in 8 lakes in British Columbia, all of which were alkaline. These lakes had no permanent	Stringer and McMynn
	Ptychocheilus oregonense		British Columbia	0.10 (O)		inlet or outlet streams. Initial results were recorded in 4 hours. Many dead trout and shiners were observed. All	(1958)
	Cyprinus carpio		Columbia	0.10 (O)		caged fish were dead in 2 days.	
	Richardsonius balteatus			0.10 (O)			
	Mylocheilus			0.10 (O)			
	caurinum Salmo gairdneri			0.10 (O)			
Toxaphene	Fathead minnow	BSA	-	0.0051 (T4A)	<u>a</u>	It was the authors opinion that pH, alkalinity and hardness, within the usual range in natural waters, had little effect on the toxic effect of the compounds studied. The values given are from Henderson, Pickering, and Tarzwell, "The Relative Toxicity of Ten Chlorinated Hydrocarbon Insecti- cides to Four Species of Fish". It is interesting that the different tables from the above book (as reported in this paper) report widely different values for the same compounds. This experiment was performed in hard water.	Tarzwell (1959)
Toxaphene	Fathead minnow Bluegill Goldfish	BSA	-	0.0075 (T4A) 0.0035 (T4A) 0.0056 (T4A)	<u>a</u>	Comment same as above except that this experiment was performed in soft water.	Tarzwell (1959)

	Toxaphene	Pimephales promelas	BSA	-	0.0075 (T4A)	а	Bioassay investigations of the new insecticides indicate that in general the organic phosphorus compounds are not as	Tarzwell (1959)
		Lepomis macrochirus			0.0035 (T4A)		toxic to fish as are the chlorinated hydrocarbons. The toxicity of most of these materials was not significantly influenced by water quality. Therefore it is to be expected that the toxicity of these materials will not differ significantly in different streams.	
	Toxaphene	Channel catfish (fingerlings)	BSA	-	2.5 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	Toxaphene	Pimephales promelas	BSA	-	0.008 (T4A)	<u>a</u> def	Concentrations were based on percent active ingredient.	Henderson, et al (1960)
		Lepomis macrochirus			0.004 (T4A)			
		Carassius auratus			0.006 (T4A)			
		Lebistes reticulatus			0.02 (T4A)			
	Toxaphene (100%)	Pimephales promelas	BSA	-	0.0075 (T4A)	abecdf	Dilution water was usually soft although some studies were conducted with hard water.	Henderson, et al (1959)
		Lepomis macrochirus			0.0035 (T4A)			ş
5		Carassius auratus			0.0056 (T4A)			PE
2		Lebistes reticulatus			0.020 (T4A)			APPENDIX Schoetteer and
	Toxaphene (tech)	Lepomis cyanellus	BSCH	-	0.0036 (SC4)	<u>aep</u>	Toxicity is reported as the sublethal concentration (SC), which is defined as that concentration which produced no	Schoettger and 🗙 Olive 😡
		Onchorhynchus nerka			0.0036 (SC4)		greater mortality among test animals than was sustained by the controls. In fish study, test fish were challenged with	(1961)
		Notropis sp Daphnia			0.01 (SC4) 0.03 (SC7)		solvent extracts of toxaphene-exposed algae and periphyton. Fish tested against the algae extract survived, but fish tested	
		pulex			0.00 (005)		against periphyton extracts died. Various technical grades	
		D. magna Ischnura sp			0.03 (SC5) 0.004 (SC4)		of toxaphene were evaluated.	
		Enallagma sp			0.004 (SC4)			
8		Scenedesmus incrassatulus			0.01 (SC384)			
COMMERCIAL C	Toxaphene	Pimephales promelas	BSA	-	0.013 (T4A)	a c d f g	Test water was spring water diluted with distilled water. Removal of toxic chemicals by carbon adsorption, chlorine and chlorine dioxide treatment, and alum coagulation was studied. The most effective method to remove fish poisons was by use of activated charcoal adsorption.	Cohen, et al (1961)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Chemical Toxaphene	Gizzard shad Bluegill Black crappie Largemouth bass Brown bullhead Shortnose gar Golden shiner Bowfin Gambusia affinis Notropis maculatus Fundulus semínolis	FL	Fla.	(0)	c d	A variety of lake types was employed to discern selective fish-killing properties of Toxaphene. Concentrations ranging from 1 to 85 ppb were placed in fourteen bodies of water varying in size from 0.5 to 2100 acres. Differences in concentrations required to cause total kills of fish popula- tions in treated lakes appeared to be related to bicarbonate alkalinities, bottom types, amounts of plankton, vegetation, and the sizes of fish present. In 4 of the lakes, a total fish kill occurred at 15-36 ppb Toxaphene.	Huish (1961)
Toxaphene	Oncorhynchus kisutch Oncorhynchus tshawytscha Salmo gairdnerii Gasterosteus aculeatus	BSA	_	9.4 (T4A) 2.5 (T4A) 8.4 (T4A) 7.8 (T4A)	<u>a</u> cd <u>e</u>	Chemical dissolved in acetone. TL <sub>m</sub> expressed in ppb.	Katz (1961) /
Toxaphene	Entonostraca Cyclops Diaptomus Ceriodaphnia Bosmina Leptodora Rotaria Filinia Keratella Polyarthia Asplanchna Brachionus Protozoa Ceratium Difflugia	FL	Col.	0.1% (K)	a c d g p	All chemical and physical data were collected and compiled by standard limnological techniques. Chemical analyses were conducted monthly. Biweekly plankton collection showed "reduction to zero" of all organisms studied, but recovery of populations to normal population numbers within several months.	Hoffman and Olive (1961)
Toxaphene	Salmo gairdneri	BSA	-	(O)	<u>a</u> cdfg	Water employed for this experiment was a relatively hard, alkaline-type taken from 3 sources: Mormon Reservoir, Magic Reservoir, and Redfish Lake. The TL <sub>m</sub> given are recorded respectively for each reservoir: 0.0135 (T4), 0.0165 (T4), and 0.0145 (T4).	Webb (1961)

	Toxaphene	Gammarus lacustris lacustris	BSA	-	(0)	аер	The mortality might have been partially due to the suscept- ibility of the organism to higher temperatures, toxicity from extended exposure to copper electrodes (used to shock the organism to determine death), or the increase of CO <sub>2</sub> . Results were expressed as LT50; for example, at 0.5 ppm, 50 percent of the shrimp were killed in 96 ( $\pm$ 11) min.	McDonald (1962)
	Toxaphene	Protococcus sp Chlorella sp Dunaliella euchlora Phaeodacty/um tricornutum Monochrysis lutheri	BSA	-	0.15 (K) 0.15 (K) 0.07 (NG) 0.15 (K) 0.04 (NG) 0.04 (K)	<u>a</u>	This paper concerns the growth of pure cultures of marine plankton in the presence of toxicants. Results were ex- pressed as the ratio of optical density of growth in the presence of toxicants to optical density in the basal medium with no added toxicants. NG = no growth, but the organisms were viable.	Ukeles (1962)
	Toxaphene	Salmo gairdneri Ictalurus melas Lepomis cyanellus Pimephales promelas I. natalis Micropterus dolomieui Catastomus commersoni Potamageton spp Semotilus atromaculatus	FL	Clayton Lake, N. M.	0.01, 0.02, and 0.02 (O)* *treatments on 3 alternate days during a 6-day period	a c d e f g i o	Paper chromatography was the method used to determine toxaphene residues in some of the species listed. Mortality of native fish and others in live cars was 100%. Residues in water and sediments were also determined. Residues were as much as 4.2 ppm in dead trout following first treatment, and as much as 15.2 ppm in dead or dying bull- heads several days after the second treatment. Dead trout in live cars contained up to 3.5 ppm toxaphene up to 8 mos following initial treatment. <i>Potomogeton</i> spp contained up to 18.3 ppm, 9 days after the final treatment. Although the lake was still toxic 9 months after treatment, planting of rain- bow trout was successful 12 months after treatment.	Kallman, et al (1962)
	Toxaphene (EC8)	Gambusia affinis Rana catesbeiana (tadpoles)	FL	Cal.	0.5 (O)	a ç	Mixed populations of the indicated test species contained in cages were exposed to various insecticidal chemicals applied as dilute sprays to ponds 1/16 acre in size. The indicated toxicant concentration is in Ib/acre, and resulted in a 100 percent mortality for the fish, and a 100 percent mortality for the tadpoles in 24 hr.	Mulla (1963)
COMMERC	Toxaphene	Fish	FL	Mont.	0.13 (O)	a c d e l	Ponds were treated with the chemical to eradicate fish. The fish population included largemouth bass, bluegills, black crappie, yellow perch, carp, white sucker, and longnose sucker. Counts were made of various fish at various later times. The paper contains little quantitative data.	Wollitz (1963)
JAL CHEMICAL	Toxaphene	Carassius auratus Gambusia affinis Salmo gairdnerii	BSA	-	0.005-0.066 (O) 0.005-0.059 (O) 0.013-0.054 (O)	<u>a</u> cde	Natural water from various sources were used. Chemical added as either floating or sinking type formulations. Toxicity given as $\rm LC_{50}$ in ppm.	Workman and Newhold (1963)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Toxaphene	Notemigonus crysoleucas Lepomis macrochirus L. cyanellus	BSA	-	(B) 30 (T 1.5) (A) 1200 (T 1.5) (B) 23 (T 1.5) (A) 1600 (T 1.5) (B) 38 (T 1.5) (A) 1500 (T 1.5)	<u>a</u> cf	Chemical was dissolved in acetone. Final concentration of acetone was $\leq 2 \text{ mi/l}$ . Data shows $TL_m$ ppb for insecticide-resistant (A) and insecticide nonresistant (B) strains of the test fish.	Ferguson, et al (1964)
Toxaphene	Gambusia affinis affinis	BSA	-	0.01 to 0.48 (O)	8	The lower value is for fish that had never been exposed to the toxicant, and the higher value was obtained with fish that had been exposed to a sublethal dose in the past. Apparently such an exposure produces a resistance that can be retained when they are later placed in clean watar.	Boyd and Ferguson (1964)
Toxaphene	-	FR	Flint Creek, Ala.	0.210 (K)	-	Conventional treatment in a water purification plant did not reduce the amount of chemical found in the stream. Data are given for 4 years 1959-62, with a range of concentrations. Only the highest value is reported here. Some fish kill is reported, but species are not identified here. Data are also reported for all seasons to show variation; the one listed here is for summer 1960.	Nicholson, et a (1964)
Toxaphene	Bluegill	BSA	-	0.0035 (T4A)	â	Assays were conducted in soft water at 25 C. Decrease in brain cholinesterase was measured in fish exposad to the toxicant.	Weiss (1964)
Toxaphene	Leiostomus xanthurus	BSFCHA	-	0.003 (K)	8 0	Assays were performed in seawater and results are reported simply as number dead. Concentrations above 0.0032 ppm killed all test animals in 24 hr. Fish were also exposed to 0.01 ppb and 0.1 ppb for 5 months, and growth was not different from those in the wild populations.	Lowe (1964)
Toxaphene	<b>Palae</b> monetes kadiakensis	BSA	-	(N) 57.5 (71¼A) (TB) 170.0 (T1½A)	acf	Test organisms were collected from 2 locations, Twin Bayou (TB), Sunflower Co., Miss. (Agricultural area) and Noxubee National Wildlifa Refuge (N), Noxubee Co., Miss. (non- agricultural area) and evaluated in leboratory bioassays.	Ferguson, at al (1965)
Toxaphene	Gambusia affinis Ictalurus malas	BSA	-	0.01-0.04 (T3A) 0.004-0.050 (T3A)	<u>a</u> cde	Test fish were collected from 8 different locations of the Mississippi River. The 3-day TL <sub>m</sub> values were made to detarmina if a resistance gradient existed. The deta indi- cated that there was none.	Ferguson (1965)
Toxaphene	Rainbow trout	BSA	-	5.4 (T4A) 2.7 (T4A) 1.8 (T4A)	<u>a</u>	These experiments were performed to show the effect of temperature on the toxicity. For the toxicant concentrations listed, the temperatures were respectively 45, 55, and 65 F. The fish all were approximately one grain in weight. Toxicant concentrations for one and 2-day times were also listed.	Соре (1965)

	Toxaphene	Fish Chironomus (larvae and pupae)	FL	Wis.	0.1 (K) (O)	_	Elimination of fish population was accomplished with toxaphene at the indicated concentration. This is a popu- lation succession study over a 3-year period with observa- tions on change in populations due to various ecological factors.	Hilsenhoff (1965)
		Chaoborus (larvae and pupae)			(0)			
		Physidae Crustaceans			(O) (O)			
Toxaphene	Toxaphene	Salmo gairdneri Campostoma	BSA	-	0.0084 (T4A) 0.014 (T4A)	acdefim p —	Adult fish were employed in this bioassay. In most cases, concentrations of toxaphene needed to cause 50 percent mortality decreased as the temperature increased from	Mahdi (1966)
		anomalum Carassius			0.094 (T4A)		53 F to 63 F and to 73 F. Data cited are for 53 F.	
		auratus Notemigonus crysoleucas			0.0125 (T4A)			
		Pimephales notatus			0.03 (T4A)			
		lctalurus melas			0.025 (T4A)			
5	Toxaphene	Oncorhynchus garbuscha O. keta O. kisutch Cattus aleuticus Salvelinus malmo Gasterasteus aculeatus Salmo gairdnerii Oncorhynchus nerka Pholis laeta Osmeridae	FR	Big Kitoi Creek, Alaska	(0)	_	<ul> <li>The purpose of this experiment was to determine the extent of predation by sculpins on pink salmon fry, and the effects of toxaphene on the sculpins and bottom fauna.</li> <li>Toxaphene applied to the experimental area was estimated to be an average concentration of 1.5 ppm.</li> <li>At the above concentration insects were completely eradicated, bottom fauna decreased in numbers and weight, but some other invertebrate groups were not completely eliminated.</li> <li>The organisms listed were organisms mentioned as fauna in the experimental area.</li> </ul>	Meehan and Sheridan (1966)
OMMERCIAL CHEMIC	Toxaphene	Fish	FL	Brush and Long Lakes, N. D.	(O)	-	Growth rates for yellow perch that survived a toxaphene treatment in Brush and Long Lakes in North Dakota were calculated. Brush Lake fish exhibited greatly increased growth rates for two growing seasons following the treat- ment. Increased growth rates were not evident for Long Lake fish until the next growing season. The approximate concentration of toxaphene for reducing the density of fish populations is believed to be 25 percent of the rate determined for fish eradication in most N. Dakota waters.	Warnick (1966)

Chemical	Organism	Bioassay or Field Study(1)	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Toxaphene	Polyarthra Keratella Asplanchna Conochiloides Brachionus Trichocera Daphnia Bosmina Ceriodaphnia Cyclops Cyanophyta	FL	Various lakes or reservoirs, N. D.	_	а	Marked reduction of many plankters followed a treatment of 90 ppb toxaphene. The most abundant plant-inhabiting organism and bottom fauna exhibited no marked changes after this treatment. Populations of <i>Gammarus, Physa</i> , and <i>Gyraulus</i> remained constant, while <i>Callibaetis, Caenis,</i> <i>Ischnura</i> , and <i>Tendipes</i> decreased slightly but were again numerous 1 year after treatment.	Needham (1966)
Toxaphene	Lebistes reticulatus	BSA & FL	Canada	0.001 (T2A)	ace	A bioassay method is described for determining the rate of detoxification of lake water after toxaphene treatment during a 1-year period.	Royer (1966)
Toxaphene	Carassius auratus	BCF	-	11.0 mg/i (T4CF) 0.44-1.8 mg/l (S4)	-	This method was developed to detect sublethal effects by observing behavioral aberrations. Detailed description of conditioned avoidance response apparatus is presented.	Warner, et al (1966)
Toxaphene	Simocephalus serrulatus Daphnia pulex	BSA	-	0.019 (SB) 0.015 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
Toxaphene	Fish	F	Cal.	(O)	_	This study was primarily concerned with insecticides found in fish-eating birds. Limited fish studies were also con- ducted. Toxaphene was found in trace to 8.0 ppm concen- trations in whole fish (wet weight).	Keith (1966)
Toxaphene	Salmo gairdneri Lepomis macrochirus Pteronarcys californicus Baetis sp Daphnia pulex Simocephalus serrulatus	BSA	_	0.004 (T2A) 0.004 (T2A) 0.007 (T2A) 0.047 (T2A) 0.015 (T2A) 0.019 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses sub-acute mortality as well. Effects on reproduction and behavior are also discussed. Data presented as $EC_{50}$ .	Cope (1966)
Toxephene	Petromyzon marinus (Iarvae)	FL BSA	East Bay, Alger County, Mich. —	(O) 0.080 (K15-20)	a c e	The amount of toxicant needed to give a concentration of 100 ppb (0.100 ppm) was applied over the surface of the lake. Small fish were observed surfacing and dying the day after treatment was made. Mortality increased daily and reached a peak on the 3rd and 4th days. The first dead larval lempreys were seen on the 4th day after treatment. At the end of 36 days exposure in cages, only 2 of 90 ammocetes were alive.	Gaylord and Smith (1966)

Toxaphene	Carassius auratus Salmo gairdnerii	FL	Big Bear Lake, Cal.	0.3-0.10 (O)	a c	The chemical was sprayed from a plane into the lake to rid it of goldfish. Small fish began dying in 2 hours, and brown bullheads were seen to be in distress. At 0.10 ppm, large goldfish appeared to be in distress. An estimated 95% of the goldfish, and all the other fish were eliminated. The fat and flesh of goldfish, brown bullheads, and some trout were analyzed for the toxicant. In all instances the fat contained the greatest amount. The paper recommends that toxaphene not be used as a fish toxicant, because it detoxifies slowly and is a contaminant for an unknown period of time. Some trout were killed when stocked but no quantitative data are given.	Johnson (1966)
Toxaphene	Salmo gairdneri Esox lucius Cyprinus carpio Notemigonus crysoleucas Pimephales notatus Catastomus commersoni Ictalurus melas Ictalurus nebulosus Lepomis humilis Lepomis humilis Sepomis nacrochirus Pomoxis annularis Pomoxis nigromaculatus Perca flavescens Stizostedion vitreum	FL	Various lakes, N. D.	0.035 (O)		<ul> <li>Minimum levels of toxaphene lethal to fish in prairie lakes and reservoirs were determined.</li> <li>Considering all lakes in general, 0.005 to 0.020 ppm resulted in incomplete mortality, while 0.025 to 0.035 ppm resulted in complete mortality. The minimum lethal concentration for treatment of most North Dakota lakes was 0.025 ppm of toxaphene.</li> </ul>	Henegar (1966)
Toxaphene CO CO CO CO CO CO CO CO CO CO CO CO CO	Gambusia affinis	BSA	_	(O)	<u>a</u>	The effect of combinations of pesticides was studied. In general, the results reflected the extreme levels of Endrin and Toxaphene resistance in the resistant population. The results failed to indicate additive effects wherein the combi- nation mortality exceeded the sum of the mortalities pro- duced by the individual insecticides.	Ferguson and Bingham (1966)

APPENDIX B

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Toxaphene	Pimephales promelas	BSA	-	0.05 (K9) 0.02 (K21) 0.05 (K6) 0.02 (K11)	a b	<ul> <li>At 33 to 39 F a complete kill occurred at concentrations of 0.05 and 0.02 in 9 and 21 days, respectively.</li> <li>At 56 F a complete kill occurred at concentrations of 0.05 and 0.02 ppm in 6 and 11 days, respectively.</li> <li>Toxaphene will kill fish at near-freezing temperatures at concentration as low as 0.02 ppm but the length of time required for a complete kill is longer than at higher temperatures.</li> </ul>	Schaumberg, et al (1967)
Toxaphene	Leiostomus xanthurus	BCF	_	0.0075 (K1A) 0.0056 (K1A) 0.0032 (K1A) 0.0018 (K2A) 0.0001 (SB 5 mo) 0.00001 (SB 5 mo)	a	<ul> <li>Experiments were conducted in salt water.</li> <li>Fish were held in plastic aquaria with a capacity of 25 liters.</li> <li>During the 5-month exposure period there was no significant difference in mortality among control and experimental fish. No symptoms of distress were noted.</li> <li>The total lengths of the fish at the end of 5 months were approximately the same for all groups.</li> <li>After the 5-month test the fish from the experimental and the control groups were exposed 48 hours to concentrations of 0.0005 to 0.0030 ppm. The fish from the experimental group seemed to be more sensitive. Concentrations of 0.0020 ppm caused complete kill whereas 0.0005 ppm did not kill any fish.</li> </ul>	Kaplan and Overpeck (1967)
Toxaphene	Pteronarcys californica (naiads) Pteronarcella badia (naiads) Claasenia sabulosa (naiads)	BSA	-	0.0023 (T4A) 0.003 (T4A) 0.0013 (T4A)	<u>acdef</u>	、Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
2(2,4,5) TP (tech)	Rainbow trout Bluegill	BSA	-	14.8 (T4A) 9.6 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
2, <b>4,5-</b> TP	Rainbow trout Bluegill	BSA	-	1.3 (T2A) 0.50 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)
1(2,4)TP (tech)	Bluegill	BSA	-	8.6 (T4A)	a	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
Freflan	<i>Pteronarcys</i> sp (nymphs)	BSA	-	3.0 (T4A)	a	Experiments were all conducted at 60 F in 1964. The values were listed as LC <sub>50</sub> .	Cope (1965)
Freflan (EC)	Rainbow trout Bluegill	BSA	-	0.010 (T4A) 0.018 (T <b>4</b> A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Соре (1965)
Freflan	Rainbow trout Bluegill	BSA	-	0.011 (T2A)	-	Data are given as LC <sub>50</sub> .	Bohmont (1967)

COMMERCIAL CHEMICAL PRODUCTS

B-230

	Trefmid (WP)	Rainbow trout Bluegill	BSA	_	0.110 (T4A) 0.345 (T4A)	а	This is an estimated LC $_{\rm 50}$ value at temperatures from 55 to 75 F.	Соре (1965)
	Tri-6 (dust No. 30, 3 percent BHC)	Penaeus aztecus P. setiferus	BSA	_	0.035 (T1A) 0.40 (T1A)	<u>a</u> c	<i>P. aztecus</i> and <i>P. setiferus</i> ranged in size from 29 to 50 mm and 11 to 13 mm, respectively. The water was aerated until the end of the assay.	Chin and Allen (1957)
В	Trichlorofon	Pteronarcys californica (naiads) Pteronarcella badia (naiads) Claasenia sabulosa (naiads)	BSA	_	0.035 (T4A) 0.011 (T4A) 0.022 (T4A)	<u>acdef</u>	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
	Tricon (oil spill eradicator)	Roccus saxatilis	BSA	-	0.001 (O) 0.005 (K 1 hr) 2.0 (K 7 min)	а	At 0.001 percent, the fish showed signs of distress in 1-1/2 hours. This compound was toxic at low concen- trations and should not be used to treat oil spills.	Chadwick (1960)
	Tricon oil-spill eradicator	Roccus saxatilis	BSA	_	(O)	<u>a</u>	This chemical is a commercial product designed to emulsify oil spilled on water. At 0.0005% concentration all test fish survived. At 0.001% concentration all fish died within 10 hours. Additional data are presented.	Chadwick (1960)
B-231	Trifluralin	Bluegill Fathead minnow Goldfish	BSA	_	0.0582 (O) 0.0934 (O) 0.585 (O)	<u>a b</u>	In static soil-water tests, 48 and 227 times more Trifluralin was required to produce an $LC_{50}$ to bluegills for two types of soil than was necessary in the static water tests. In a simulated field test using swimming pools, Trifluralin, applied at 1 lb/acre to Brookston soil and then irrigated with 10 inches of water, was not toxic to bluegills. On the basis of these studies, it was concluded that $LC_{50}$ values derived from static water fish tests are unrealistic in predicting the toxicity of Trifluralin to fish under field conditions.	Parka and Worth (1965)
8	Trifluralin	Bluegill	BSA	_	8.4 (T4A)* *ppb	<u>a</u>	The temperature effect is extreme in the case of this com- pound. The T4 listed is for a temperature of 85 F. At 45 F the T4 was 280 ppb. The T1 is even more striking. At 85 F, the value was 10.0 ppb, and at 45 F, 1300 ppb.	Соре (1965)
COMMERCI	Trifluralin (tech)	Bluegill Rainbow trout	BSA	_	0.068 (T4A) 0.086 (T4A)	а	This is an estimated LC $_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
AL CHEMICAL PRODUCTS	Trifluralin	Salmo gairdneri Lepomis macrochirus Pteronarcys californicus Daphnia pulex Simocephalus serrulatus	BSA	-	0.011 (T2A) 0.019 (T2A) 4.200 (T2A) 0.240 (T2A) 0.450 (T2A)	а	This paper reports acute toxicity of a number of compounds, and discusses subacute mortality as well. Effects on re- production and behavior are also discussed. Data presented as EC <sub>50</sub> .	Cope (1966)

COMMERCI	Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm(3)	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
AL CHEMICAL P	Trifluralin	Simocephalus serrulatus Daphnia pulex	BSA	-	0.450 (SB) 0.240 (SB)	_	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. "Water Chemistry" (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
PRODUCTS	Trifluralin	Pteronarcys californica (naiads)	BSA	-	0.003 (T4A)	acdef	Data reported as LC $_{50}$ at 15.5 C in 4 days.	Sanders and Cope (1968)
TS	Tris- buffer	Homarus americanus	BSA	-	2200-4400 (SB10)	ace	Tris-buffer concentrations in the range tested were safe for regulating activity. The lobsters employed weighed 500 grams.	Stewart and Cornick (1964)
	Trithion	Gambusia affinis	BSA	-	0.2 (K 7%)	а	Chemicals were dissolved in acetone, and tests were run in triplicate. Toxicity is given as average percent fish killed in 24 hr.	Lewallen (1959)
	Trithion	<i>Salmo gairdnerii</i> (one wk old sac fry)	BSA	-	0.5 (K 0%) 5.0 (K 0%)	<u>ac</u>	Results are averages of triplicate tests. Toxicity is reported as percent mortality (K %).	Lewallen and Wilder (1962)
,		(one mo old feeding fry)	BSA		0.5 (K 7%) 5.0 (K 93%)			
	Trolene	Salmo gairdneri Salmo trutta Salvelinus	BSA	-	0.74 (T2A) 0.39 (T2A) 0.39 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
		fontinalis Salvelinus namaycush			0.62 (T2A)			
		lctalurus punctatus Lepomis macrochirus			1.26 (T2A) 1.00 (T2A)			
	Trypaflavine (acriflavine neutral)	lctalurus punctatus	BSA	-	17.9 (K2) 11.5 (T2A)	acfi	The experiment was conducted at 66 C.	Clemens and Sneed (1958)
	Trypaflavine (acriflavine hydro- chloride)	Channel catfish (fingerlings)	BSA	-	11.5 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
	TV-1096	Salmo gairdnerii Salvelinus	BSA	-	16.1 (T2A) 19.0 (T2A)	<u>a</u> f	Variance and the 95-percent confidence interval (C.I.) were also determined.	Willford (1966)
		Salvennus fontinalis Salvelinus namaycush			16.5 (T2A)			

		lctalurus punctatus Lepomis macrochirus			20.3 (T2A) 28.2 (T2A)			
	UC-8305 (EC4)	Gambusia affinis	FL	Cal.	0.5 (K1)	_	Toxicity value is in Ib/acre.	Mulla (1966)
	Union Carbide, UC 10854	Rainbow trout Bluegill	BSA	-	0.180 (T4A) 0.110 (T4A)	_	The values reported are given as $LC_{50}$ .	Cope (1965)
	Union Carbide, UC 21149	Rainbow trout Bluegill	BSA	-	0.560 (T4A) 0.050 (T4A)	-	The values reported are given as $LC_{50}$ .	Cope (1965)
	UC-21427 (EC2)	Gambusia affinis Rana catesbeiana	FL	Cal.	0.5 (K1) (O)	-	Toxicity value is in lb/acre. No mortality in tadpoles of <i>Rana catesbeiana</i> occurred during an exposure period of 1 week.	Mulla (1966)
	Urox	Water lettuce	FL	Lakes in Fla.	(O)	_	11.2 to 22.5 lb/acre controlled water lettuce.	Phillippy (1961)
	Vancide 51 salt	Pimephales promelas	BSA	_	0.83 (T4A)	a c d e f	The toxicity of this substance was not influenced by the quality of the water (pH, hardness, alkalinity).	Pickering and Henderson (1966)
	Vancide 51Z	Pimephales promelas Lepomis macrochirus Lebistes reticulatus	BSA	_	0.35 (T4A) 0.85 (T4A) 0.59 (T4A)	a c d e f	Comment same as above.	Pickering and Henderson (1966)
	Vapona (DDVP)	Pimephales promelas Lepomis macrochirus	BSA	-	4.0 (T4A) 0.27 (T4A)	a c d e f	Comment same as above.	Pickering and Henderson (1966)
	Veon 100	Spatterdock	FL	Fla.	(0)	-	At 10.0 lb/acre, 5 percent control of spatterdock was obtained.	Copeland and Woods (1959)
COMMERCIAL CHEMICAL PR	Veon 245 (2,4,5-T)	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	1.0 (NTE) 1.0 (NTE) 1.0 (NTE) 1.0 (NTE)	_	<ul> <li>Seawater was pumped continuously into test aquaria.</li> <li>Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following:</li> <li>Oyster – 96-hr EC<sub>50</sub> – Conc. which decreased with shell growth.</li> <li>Shrimp – 48-hr EC<sub>50</sub> – Conc. which killed or paralyzed 50% of test animals.</li> <li>Fish – 48-hr EC<sub>50</sub> – Conc. which killed 50%.</li> <li>Phytoplankton – Percent decrease of CO<sub>2</sub> fixation to a 4-hr exposure at 1.0 ppm chemical concentration.</li> </ul>	Butler (1965)

# APPENDIX B

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
	Leiostomus xanthurus (juvenile)	BSA		(0)	а	Water temperature was 27 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
	Oyster	BCF	_	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
Vernam	Penaeus aztecus	L	-	(0)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC50 or enough to cause loss of equilibrium or mortality. No effect occurred at 1.0 ppm.	Butler (1965)
Vernam	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	_	1.0 (NTE) 1.0 (O, 20%) 1.0 (NTE) 	_	$\begin{array}{llllllllllllllllllllllllllllllllllll$	Butler (1965)
Vernam	Penaeus aztecus	L	-	(O)	а	Toxicant chemicals were evaluated in seawater at tempera- tures averaging about 28 C. The values are for 24-hr EC <sub>50</sub> or enough to cause loss of equilibrium or mortality. Response irritation occurred at 1.0 ppm.	Butler (1965)
Vernam (tech)	Bluegill	BSA	-	4.0 (T4A)	а	This is an estimated $LC_{50}$ value at temperatures from 55 to 75 F.	Cope (1965)
Vernam	<i>Leiostomus</i> <i>xanthurus</i> (juvenile)	BSA	-	(O)	а	Water temperature was 28 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Vernam	Oyster	BCF	_	(0)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)
Versenol (iron chelate)	Channel catfish (fingerlings)	BSA	-	1.9 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Versene (acid)	Channel catfish (fingerlings)	BSA	-	167 (K1A)	<u>a</u>	Comment same as above.	Clemens and Sneed (1959)
Weeder, MCP	Lepomis macrochirus	BSA	-	(0)	а	No mortality was noted with concentrations of 10,000 mg/l for over 100 hr. The experiment was conducted at 65 F. Fish were 2 in. long.	Cope (1963)
Weedex	Gardonus rutilus Tinca tinca (fry)	FL	France	(NTE) (K)	_	Eight small ponds were studied. The chemical was nontoxic to larger fish, but toxic to the fry. Ponds were emptied after one month of exposure to the chemical.	Vivier and Nisbet (1965)

Weedex (Weedazol, Weedazol T. L.)	Phoxinus phoxinus	BSA	-	(O)	<u>a</u> c <u>d</u> e	The assays were conducted in dual aquaria with aeration. Toxicity was low after 1 month at normally used concen- trations, as follows: weedex — 40-80 ppm; weedazol — 15-30 ppm; weedazol — 20-40 ppm.	Vivier and Nisbet (1965)
Weptachlor	Daphnia magna	BSA	-	0.052 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 64 hr. Data cited are for 78 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
WL 8008 (isobutyl- triphenyl- methylamine)	Bulinus truncatus Biomorpholaria alexandrina Lymnaea caillaudi	FO	Arabia	0.24-0.25 (O)	a b g	Tests were conducted in the Khurshid canal. Flow of molluscicide was discharged directly into the canal and was maintained at a concentration of 0.24 ppm during the 6 hr of treatment. Formulation 1 killed all adult organisms but did not affect eggs. Snail density reached its pretreatment level after 4 months. Formulation 2 was tested in Ganabiet el Sarania canal. Molluscicide was applied for 6 hr by motor-operated dispenser to give a con- centration of 0.25 ppm. Adult organisms were killed while eggs were unaffected.	Dawood and Dazo (1966)
WL 8008 (n-trityl- morpholine)	Australorbis glabratus	BSA & FL	Puerto Rico	Variable	c	Seven of the tested compounds failed to meet acceptability criteria – that is, complete kill after 6-hr exposure to 10 ppm. They were not used in field tests. Field tests showed WL 8008 to be highly effective.	Seiffer and Schoof (1967)
Xylene + 2% nonionic emulsifier	Elodea canadensis Potamogeton nodosus Potamogeton pectinatus	BSA		5 (O) 100 (O) 5 (O) 100 (O) 5 (O) 100 (O)	а	Experiments were conducted in standing water. Results were rated on a scale of 0 to 10, 0 standing for no toxic effect and 10 signifying a complete kill. Evaluation was based on visual observation of the plant response at weekly intervals for 4 weeks. No toxic effect. Injury rating of 9.3. No toxic effect. Injury rating of 7.9. No toxic effect. Injury rating of 8.6.	Frank, et al (1961)
Zinc dimethyl dithio- carbamate (ZDD)	Pimephales promelas	BSA		(0)	a c d e f	Toxicity to 30 species of algae are also presented. ZDD was algicidal in the range 0.25 to 2.0 ppm.	Maloney and Palmer (1956)
Zectran	Penaeus aztecus	L	-	0.0068 (O)	а	Toxicant chemicals were evaluated in sea water at temperatures averaging about 28 C. The values are for 24-hr $EC_{50}$ or enough to cause loss of equilibrium or mortality.	Butler (1965)
Zectran	<i>Cyprinodon variegatus</i> (juvenile)	BSA	-	(O)	а	Water temperature was 12 C. No effect was noticed on exposure to 1.0 ppm.	Butler (1965)
Zectran	Oyster	BCF	-	(O)	а	No effect on exposure to the chemical at 1.0 ppm.	Butler (1965)

APPENDIX B

Chemical	Organism	Bioassay or Field Study <sup>(1)</sup>	Field Location(2)	Toxicity, Active Ingredient, ppm <sup>(3)</sup>	Experimental Variables Controlled or Noted <sup>(4)</sup>	Comments	Reference (Year)
Zectran	Pteronarcys californica (naiads)	BSA	-	0.010 (T4A)	a c d e f	Data reported as LC <sub>50</sub> at 15.5 C in 4 days.	Sanders and Cope (1968)
Zectran	Simocephalus serrulatus Daphnia pulex	BSA	-	0.013 (SB) 0.010 (SB)	-	Concentration reported is for immobilization. Time for immobilization was 48 hr. Data cited are for 60 F, but assays were performed at varied temperatures. Water chemistry (unspecified) was "controlled" during the assay period.	Sanders and Cope (1966)
Zerlate	Channel catfish (fingerlings)	BSA	-	1.0 (K1A)	<u>a</u>	Tap water was used. Considerable additional data are presented.	Clemens and Sneed (1959)
Zinc disodium versenate	Channel catfish (fingerlings)	BSA	-	>500 (K1A)	a	Comment same as above.	Clemens and Sneed (1959)
Zinophos (EC4)	Micropterus salmoides	BSA	-	0.25 (T2A) 0.5 (K2) 1.0 (K1)	a e	Experiments were carried out in fiber glass tubs filled with well water. Fish weights ranged from 2 to 6 pounds.	Mulla, et al (1967)
Zytron	Oyster	BCF	-	0.33 (O)	а	The value reported in a 96-hr $\text{EC}_{50}$ (decreased shell growth).	Butler (1965)
Zytron	Leiostomus xanthurus (juvenile)	BSA	-	0.32 (O)	а	Water temperature was 27 C. The figure reported is a 48-hr $\text{EC}_{\overline{50}}.$	Butler (1965)
Zytron	Crassostrea virginica Penaeus aztecus Leiostomus xanthurus Phytoplankton	BCFA & BSA	-	0.33 (O) 0.0003 (O) 0.32 (T2CFA) 59% (O)	-	Seawater was pumped continuously into test aquaria.Salinity, temperature, and plankton fluctuated with tide, and ambient weather conditions. Some bioassays with fish were static. Toxicity was reported for the following: Oyster – 96-hr EC50 – Conc. which decreased shell growth.Shrimp – timp – 48-hr EC50 – Conc. which killed or paralyzed 50% of test animals.Fish – Phytoplankton – Percent decrease of CO2 fixation to a 4-hr exposure at 1.0 ppm chemical concentration.	Butler (1965)

# APPENDIX C

SPECIES INDEX FOR APPENDICES A AND B

### SPECIES INDEX FOR APPENDICES A AND B

Abramis brama - A-9, A-166 Acartia clausi - A-42, A-64, A-84 Accipiter gentilis - B-53, B-55, B-75, B-96, B-142, B-152 A. nisus - B-55, B-75, B-120, B-142, B-152 Acheilognathous limbata - A-72 Acris crepitans – B-66 A. gryllus B-66 Acroneuria pacifica - B-10, B-11, B-23, B-24, B-63, B-64, B-68, B-70, B-95, B-106, B-107, B-110, B-111, B-112, B-118, B-119, B-120, B-136, B-156, B-158, B-159, B-183, B-184, B-185 Aedes aegypti – B-58, B-64, B-99, B-183 A. tseniorhynchus – B-183 Aeroneuria pacifica – B-94 Agrion spp - B-197 Algae - A-24, A-43, A-44, B-31, B-37, B-41, B-55, B-122 Alisma spp – A-118, B-203 Alloperla spp - B-200 A. pseudoharengus - B-75, B-142, B-143 Althernanthera philoxeroides - B-16, B-17, B-38, B-44, B-45, B-153, B-205, B-211 Alosa chrysochloris – A-158 A. pseudoharengus - B-55, B-79, B-80, B-98 Ambloplites rupestris – A-54, A-87, A-90, A-96, A-154, **B-86** Ameiurus spp – B-218 A. nebulosus - A-80, B-43 Ameletus spp – B-200 Ammodytes lanceolatus – B-178 Amphipnous cuchia – B-38, B-81 Amphipoda - A-126, A-144, A-146, B-74 Anabena spp - A-24, A-48, A-109, A-119 A. circinalis – A-47 A. flos-aquae A-73, A-96 Anacystis spp - A-24, A-48, A-109, A-119 Anarchis canadensis – A-115 Anax spp - B-197 Ancistrodon piscivorus – B-123 Anguilla anguilla – A-134 A. rostrata – B-78 Anguillidae - B-74 Anisoptera spp – B-81 Ankistrodesmus spp - A-24, A-48, A-109, A-119 Annelida - B-56, B-199 Anodonta grandis - B-13, B-78, B-166 Anopheles quadrimaculatus – A-97, B-23, B-49, B-99, B-183 Aphredoderus sayanus - B-139

Aplodinatus grunniens – B-55, B-72, B-79, B-80, B-98, B-142, B-143 Aptenodytes fosteri – B-74 Aquatic insects – B-61 Arctopsyche grandis - B-63, B-64, B-70, B-156, B-159, B-183, B-185 Argia spp - A-130, A-164 Artemia salina – A-42, A-64, A-84, B-61, B-91, B-182 Arthropoda - B-74 Arundo donax - B-56, B-79, B-80 Asellus spp - A-130 A. aquaticus – B-53 A. communis – A-116, A-130, A-164 Asio otus – B-152 Asplanchna spp – B-196, B-224, B-228 Australorbis glabratus – A-151, A-152, A-156, B-6, B-33, B-132, B-143, B-169, B-216, B-235 Azolla caroliniana - B-102, B-180 Bacteria (see sewage organisms) Baetis spp - B-71, B-95, B-120, B-141, B-160, B-228 Balanus balanoides - A-41, A-43, A-83, A-114, A-163 B. eberneus - A-41, A-42 Barbus stigma – B-68, B-93, B-151 B. machecola – B-179 Bass - A-44, A-51, A-69, A-76, A-113, A-115, B-37, B-52, B-55, B-59, B-122, B-134, B-151, B-160, B-184, B-195, B-219, B-220 Belostomidae – B-104 Biomorpholaria alexandrina – A-14, A-24, A-25, A-38, A-65, A-66, A-81, A-103, A-108, A-129, A-137, A-139, A-145, B-169, B-235 Black fish – B-52 Blackfly – B-29, B-30, B-35, B-59, B-192, B-217 Bladderwort - A-117, B-153 Blue crab – B-30, B-61, B-69, B-209 Bluegill – A-35, A-36, A-43, A-48, A-53, A-76, A-113, A-115, A-117, A-118, B-7, B-9-12, B-27, B-30, B-31, B-34, B-35, B-37, B-40, B-42, B-49, B-51, B-52, B-53, B-57, B-59, B-61, B-68, B-70, B-81, B-84, B-87, B-89, B-90, B-91, B-94, B-99, B-101, B-107, B-108, B-111, B-112, B-115, B-118, B-121, B-124, B-128, B-129, B-134, B-136, B-139, B-141, B-144, B-145, B-147-149, B-151, B-157, B-158, B-160, B-164, B-165, B-168, B-169, B-181, B-184, B-185, B-187, B-200-202, B-204, B-206, B-207-210, B-219, B-220, B-222, B-224, B-226, B-230, B-231,

Boleosoma nigrum – A-81 Bosmina - B-196, B-224, B-228 Brachionus spp - B-38, B-196, B-224, B-228 Brachydanio rerio - A-58, A-89, A-105, A-107, A-161, A-165, B-4, B-33, B-39, B-99, B-131 Brasenia schreberi – B-44 Brevoortia patronus – B-119 Buffalo fish - B-12, B-121, B-135 Bufo spp - A-69B. boreas - B-25, B-132, B-136, B-168, B-184, B-193 B. valliceps - A-20, A-38, A-40, A-79, A-83, A-84, A-90, A-114, A-159 Bulinus truncatus – A-14, A-24, A-25, A-38, A-65, A-66, A-81, A-103, A-108, A-129, A-137, A-139, A-145, B-169, B-235 Bulla spp - A-71 Bullhead (see catfish) Buteo buteo – B-53, B-55, B-75, B-96, B-120, B-142, B-152 Cabomba spp - B-58, B-105 Caddisfly - B-29, B-30, B-35, B-59 Caenidae - B-104 Caenis spp - A-116 Calcinus maenas – B-178 Callibaetis spp – A-116 Callinectes sapidus - B-2, B-110 Calliriche spp – A-130 Calothrix spp - A-24, A-48, A-109, A-119 Calico fish - A-117 Cambarus spp - A-154 Campostoma anomalum – A-141, B-4, B-9, B-12, B-16, B-227 Cancer poguras – B-178 Carassius auratus – A-11, A-16, A-18, A-20, A-29–32, A-34, A-38, A-39, A-47, A-49, A-52, A-53, A-55, A-56, A-58-60, A-62-65, A-67, A-69, A-72-75, A-77, A-79, A-80, A-87, A-88, A-91, A-93, A-94, A-96, A-99, A-100, A-107, A-110, A-118, A-134, A-149, A-151, A-152, A-154, A-155, A-157, A-158, A-165, B-8, B-13, B-15, B-16, B-18, B-23, B-30, B-32, B-35, B-36, B-39, B-46, B-42, B-43, B-56-58, B-59, B-62, B-72, B-76, B-83-85, B-87, B-91, B-92, B-97, B-105, B-106, B-110, B-112, B-115, B-116, B-124, B-125, B-135, B-136, B-139, B-140, B-143, B-150, B-153, B-156-158, B-164, B-167-169, B-175, B-176, B-183, B-184, B-190, B-195, B-198, B-199, B-210, B-215, B-223, B-225, B-227-229 C. carassius – A-12, A-13, A-15, A-21, A-22, A-25, A-39, A-43, A-63, A-65, A-80, A-81, A-85, A-96, A-102, A-107, A-108, A-111, A-128, A-131, A-138, A-139, A-148, A-149, B-95 Carp - A-47, B-52, B-121, B-134, B-157, B-160, B-214, B-219, B-220 Carcinus maenas - A-40, A-132, B-19, B-32, B-40,

B-85, B-125, B-134, B-189, B-207

Cardium edule - B-19, B-32, B-40, B-85, B-125, B-133, B-134, B-178, B-189, B-207 Catfish - A-3, A-41, A-49, A-64, A-66, A-68, A-76, A-82, A-83, A-98, A-100, A-109, A-111-113, A-117, A-142, B-12, B-16, B-18, B-19, B-21, B-34-36, B-41, B-52, B-62, B-66, B-84, B-92, B-99, B-121, B-131, B-135, B-138, B-139, B-145, B-149, B-150, B-155, B-160, B-163, B-169, B-174, B-186, B-188, B-192, B-195, B-198, B-210, B-213, B-215, B-217, B-219, B-220, B-223, B-225, B-234, **B-236** Catharcta skua – B-54, B-74 Catostomus commersoni - A-80, A-154, B-9, B-17, B-22, B-85, B-86, B-131, B-159, B-194, B-210, B-225, B-229 C. macrocheilus – B-220–222 Cattails - B-51 Cattus aleuticus - B-227 Ceratium spp - B-196, B-224 Ceratophyllum spp - A-54 C. demersum - B-104 Ceriodaphnia spp – B-38, B-196, B-224, B-228 Chaenobryttus coronarius - B-139 C. gulosus - B-27, B-219 Chalmydomonas spp A-119 Chama punctatus – B-38, B-81 Channel catfish (see catfish) Chaoborus spp - A-47, B-49, B-101, B-220, B-227 C. astictopus - B-24, B-42, B-83, B-89, B-124, B-126, B-167, B-193 Chara spp - A-44, A-115, A-118, B-203 Chelydra serpenti – B-142 Chephia spp – B-82 Chironomidae - A-116, B-9, B-103, B-104, B-168, B-173 Chironomus spp - B-227 C. plumosus - B-195 C. riparius – B-53 Chlamydomonas spp - A-25, A-48, A-109, B-58 Chlorella spp - A-24, A-33, A-48, A-53, A-98, A-109, A-119, B-65, B-101, B-107, B-110, B-130, B-149, B-151, B-170, B-173, B-200, B-215, B-225 C. pyrenoidosa - A-18, A-28, A-47, A-67, B-2,B-4, B-13, B-19, B-43, B-128, B-209 C. variegata – A-3, A-6, A-8, A-17, A-24, A-28, A-35-37, A-42, A-45, A-50, A-54-57, A-60-63, A-73, A-78, A-83, A-86, A-90, A-92, A-93, A-101, A-102, A-112, A-113, A-128, A-129, A-142, A-143, A-151, A-152, A-155, A-157, A-162, A-163, B-7, B-28, B-29, B-44, B-60, B-91, B-187, B-193, B-194, B-209, B-211, B-215, B-216 C. vulgaris - A-40Chlorococcum humicola - A-37

Chroococcaceae - B-58

Chrosomus eos – B-72

Chubs - B-33, B-37, B-55, B-122, B-195 Cirrhina mrigala – B-82, B-154, B-163, B-179 Claassenia sabulosa - B-34, B-63, B-80, B-98, B-110, B-123, B-142, B-156, B-162, B-186, B-231 Cladocera spp - A-47, B-66, B-105 Cladophora spp - A-115, A-118, B-56, B-79, B-80, B-203 C. glomerata - B-13, B-14 C. gracilis - B-78 Clams - B-11, B-37, B-55, B-122, B-127 Closterium spp - A-24, A-48, A-109, A-119 Clupea harengus - B-178 Cnephia spp - B-2, B-14, B-26, B-33, B-75, B-88, B-90, B-99, B-100, B-110, B-129, B-131, B-133, B-144, B-149, B-165, B-173, B-176-178 Coccolithus huxleyi – B-80 Coleoptera – B-56, B-59, B-69, B-74, B-76, B-162, B-199 Coenagrionidae - B-104 Coesius plumbeus - A-154 Colpidium spp – A-116 Coontail - B-113 Copepoda - A-47, B-58, B-66, B-104 Conochiloides spp - B-228 Corbicula spp – B-49 C. manillensis – B-13, B-73, B-97, B-122, B-142, B-153, B-165 Coregonus artedii – B-55, B-79, B-80, B-98, B-142, B-143 C. clupeiformis – B-75 C. williamsoni – B-221 Corixidae - B-104 Corydoras paleatus - A-58 Cottus spp – B-69 Cottus bairdi - B-79, B-98 C. cognatus - A-81 Couesius plumbeus - B-194, B-221 Crabs - B-30, B-61, B-91, B-162, B-166, B-209 Crangon crangon - B-19, B-32, B-40, B-85, B-125, B-134, B-189, B-207 Crappies – A-76, A-113, A-115, B-52, B-59, B-160 Crassostrea virginica - A-5, B-13, B-15, B-20, B-46, B-47, B-50, B-51, B-71, B-73, B-76, B-83, B-97, B-103, B-108, B-122, B-125, B-126, B-130, B-142, B-144, B-146, B-148, B-153, B-154, B-165, B-170, B-174, B-175, B-181, B-190, B-201, B-204, B-208, B-212, B-218, B-219, B-233, B-234, B-236 Crayfish - A-76, B-64, B-66 Cristivomer namaycush - B-86, B-218 Culex spp - A-3, A-110, A-114, A-120, A-121, A-126, A-131, A-146 C. apicalis - B-52, B-58, B-87, B-91, B-109 C. pipiens quadrimaculatus – B-27, B-34, B-66, B-158, **B-184** Cyclops spp - B-31, B-38, B-58, B-196, B-224, B-228 Cylindrospermum spp - A-24, A-48, A-109, A-119

C. licheniforme - A-3, A-6, A-8, A-17, A-24, A-27, A-28, A-35, A-36, A-42, A-45, A-50, A-54-57, A-60-63, A-72, A-73, A-78, A-83, A-86, A-90, A-92, A-93, A-101, A-102, A-112, A-113, A-128, A-129, A-142, A-143, A-151, A-152, A-155, A-157, A-162, A-163, B-7, B-28, B-29, B-44, B-60, B-91, B-187, B-193, B-194, B-209, B-211, B-215, B-216 Cynodon dactylon - B-56, B-79, B-80 Cyprinella whippli – B-218 Cyprinids - A-130, B-77 Cyprinodon spp - B-157, B-235 C. variegatus - B-25, B-26, B-30, B-42, B-46, B-50, B-61, B-69, B-71, B-78, B-87-89, B-111, B-119, B-126, B-135, B-137, B-161, B-178, B-186, B-208, B-209, B-217, B-218 Cyprinus carpio - A-59, A-69, A-72, A-80, A-134, A-158, B-5, B-17, B-21, B-22, B-43, B-72, B-82, B-99, B-112, B-115, B-116, B-127, B-133, B-152, B-154, B-163, B-175, B-179, B-190, B-197, B-218, B-220-222, B-229 Cypris spp - B-38 Danio spp – B-154 Daphnia spp - A-54, A-57, B-38, B-58, B-228 D. carinata - B-12, B-72, B-88, B-96, B-101, B-121, B-127, B-141, B-161 D. magna - A-2-4, A-7, A-8, A-11-16, A-20, A-22, A-23, A-29, A-36, A-38-40, A-50, A-52, A-53, A-63, A-65, A-66, A-68, A-70-73, A-75-81, A-83-85, A-90-92, A-95, A-97, A-99, A-100, A-103, A-106-111, A-112, A-114-116, A-118-129, A-131-141, A-143-150, A-155-157, A-161, A-164, B-8, B-11, B-14-16, B-41, B-47, B-54, B-56, B-63, B-70, B-72, B-87, B-90, B-93, B-95, B-96, B-101, B-104, B-108, B-114, B-115, B-119, B-121, B-123, B-127, B-128, B-134, B-140, B-153-155, B-158, B-159, B-161, B-164, B-165, B-169, B-170, B-173, B-176, B-181, B-182, B-185, B-188, B-191, B-204, B-223, B-235, B-236 D. pulex - A-29, A-50, A-118, A-149, B-12, B-14, B-19, B-27, B-34, B-37, B-48, B-51, B-53, B-71, B-72, B-81, B-82, B-88, B-90, B-96, B-105, B-108, B-120, B-121, B-127-129, B-141, B-145, B-147, B-152, B-161, B-165, B-181, B-185, B-187, B-188, B-191, B-197, B-200, B-201, B-204, B-223, B-228, B-231, B-232 Decapoda – B-74 Delphinus delphis - B-178 Diatoms - A-163, B-58 Diaptomus spp – B-38, B-196, B-224 Difflugia spp - B-196, B-224 Dinoflagellata – B-58 Diptera – B-56, B-59, B-69, B-73, B-74, B-76, B-148, B-162, B-199 Dorosoma cepedianum - A-158, B-16, B-119, B-139 D. petenense – A-158 Dressenia spp – A-130 Drum - B-134

Dugesia spp - A-112, A-126 Dunaliella spp - B-215 D. euchlora - A-33, A-53, A-98, B-65, B-101, B-107, B-110, B-130, B-149, B-151, B-170, B-173, B-200, B-225 Dytiscidae - B-104 Dytiscus spp - B-81 Echinodermata - B-74 Eel - B-162, B-166, B-194 Eichornia crassipes – B-43, B-44 Eleocharis achcularis - B-202 Eleotridae - B-74 Elliptis crassidens – B-48, B-49 Elmidae - B-74 Elminus modestus - A-42, A-64, A-84 Elodea spp - A-54, B-58 E. canadensis - A-5, A-110, A-153, B-102, B-104, B-113 E. canadensis - B-189 Elops saurus - B-198 Enallagma spp – B-223 Enneacanthus gloriosus - B-159 E. chaetodon - B-159 Entosphenus lamottenii - B-86 Ephemerella spp - B-200 E. grandis - B-10, B-24, B-70, B-95, B-107, B-112, B-119, B-136, B-159, B-185 Ephemeroptera – A-47, B-9, B-56, B-59, B-61, B-64, B-65, B-69, B-73, B-74, B-76, B-162, B-199, B-200 Ericymba buccata - A-141, B-4, B-9, B-12, B-218 Erimyzon sucetta - B-16, B-20, B-48, B-49, B-51, B-90, B-106, B-109, B-114, B-129, B-131, B-171, B-205-207 E. oblongus - B-159 Eriocaulon spp - A-44 Erisymba buccata – B-139 Erpobdella punctata – A-130 Eucalia inconstans – B-86 Escherichia coli - A-132 Esomus danrica - B-81 Esox americanus - B-123, B-139 E. lucius – A-84, B-16, B-55, B-72, B-77, B-98, B-105, B-153, B-171, B-229 E. niger - B-78, B-159 Etheostoma blennoides - B-12 E. caeruleum – B-9, B-12 E. exile - B-17 E. flabellare – B-12 E. gracile - B-139 E. nigrum - B-12E. zonale -B-9Eucalia inconstans - B-17, B-72, B-85 Euglena spp – B-58 Eupomotis gibbosus - B-43 Eurhynchium rusciforme – B-13

Falco tinnunculus - B-53, B-55, B-75, B-96, B-120, B-142, B-152 F. peregrinus - B-55, B-75, B-96, B-121, B-143, B-152 Fall fish – B-33, B-57 Filinia spp - B-196, B-224 Fish – A-12, A-27, A-49, A-52, A-70–72, A-101, A-163, B-70, B-75, B-91, B-168, B-196, B-206, B-214 Fundulus spp - B-157 F. chrysotus - B-27, B-139 F. diaphanus – B-159 F. heteroclitus – A-159, B-78 F. notatus - A-141, B-12 F. ocellaris - B-155 F. seminalis - B-46, B-47, B-118, B-224 F. similis - A-5, A-103, B-50, B-51, B-69, B-86, B-99, B-119, B-122, B-148, B-154, B-180, B-181, B-202, B-208, B-218 Gadus merlangus - B-178 G. morrhua – B-178 Galaxiidae - B-74 Gambusia spp – B-157 G. affinis - A-2-4, A-7-12, A-14-18, A-20, A-22-25, A-27, A-46, A-49, A-50, A-52, A-56, A-65-67, A-69, A-70, A-79-82, A-89, A-92, A-97, A-102, A-103, A-105, A-106, A-108, A-109, A-111, A-114, A-120, A-122, A-124, A-126, A-130, A-132, A-135–138, A-140, A-141, A-143-150, A-152, B-2, B-5, B-6, B-9, B-10, B-22-27, B-32, B-34, B-37, B-52, B-53, B-58, B-62, B-64, B-67, B-70, B-71, B-83, B-87, B-88, B-91, B-99, B-109, B-110, B-112, B-117, B-119, B-120, B-129, B-131-133, B-135, B-136, B-139, B-140, B-143, B-145, B-146, B-151, B-155, B-156, B-165, B-168, B-169, B-173, B-176-179, B-182-184, B-187, B-193, B-198, B-199, B-214, B-217, B-218, B-224-226, B-229, B-232, B-233 Gammarus spp - B-54, B-55, B-79, B-98, B-122 G. lacustris – A-47, B-8, B-10, B-11, B-18, B-23, B-24, B-37, B-68, B-70, B-93, B-94, B-107, B-111, B-112, B-117-119, B-136, B-140, B-150, B-158, B-159, B-184, B-185, B-189, B-196, B-224 G. pulex - A-134 Gar – B-134, B-224 Gardonus rutilus - B-234 Gasterosteus aculeatus – A-7, A-12, A-16, A-22, A-25, A-27, A-37-39, A-40, A-42, A-79, A-81, A-83, A-92, A-95, A-108, A-114, A-134, A-138, A-146, A-149, A-159, A-164, B-8, B-36, B-41, B-64, B-78, B-93, B-116, B-117, B-135, B-140, B-150, B-156, B-164, B-199, B-224 Gastropoda - B-74 Gastrotrica spp - B-38 Gerridae - B-104 Gila robusta – B-219 Gleocapsa spp - A-6, A-27, A-63, A-86, A-155 Gleotrichia echinulata - A-47 Gnathepogon gracilis – A-72 Gobio gobio - A-10, A-166, A-167

Goldfish - A-7, A-17, A-24, A-66, A-97, A-125, A-136, A-138, B-7, B-30, B-35, B-52, B-58, B-61, B-76, B-87, B-91, B-109, B-115, B-139, B-149, B-157, B-164, B-222, B-231 Gomphonema spp - A-25, A-48, A-54, A-109, A-119 G. parvulum - A-3, A-6, A-8, A-17, A-24, A-27, A-28, A-35, A-36, A-42, A-45, A-50, A-55-57, A-60-63, A-73, A-78, A-83, A-86, A-90, A-92, A-93, A-101, A-102, A-112, A-113, A-128, A-129, A-142, A-143, A-151, A-152, A-155, A-157, A-162, A-163, B-7, B-28, B-29, B-44, B-60, B-91, B-187, B-193, B-194, B-209, B-211, B-215, B-216 Guppies - B-7, B-30, B-35, B-61, B-91, B-109, B-115, B-139, B-144, B-145, B-149, B-164, B-189, B-192, B-222 Gyraulus circumstriatus - A-46, A-130 Haematopus ostralegus - B-54, B-96, B-121 Haliphidae - B-104 Halichoerus grypus – B-178 Hebridae - B-104 Heleidae spp - B-49 Helioperca incisor – B-218 Helisoma campanulata - A-130, A-165 Helix pomatia - A-48 Heloscidium spp – A-130 Hemigrammus crythrozonus - A-58 Heptagenia spp – B-200 Heteropneustes fossilis - B-31, B-38, B-93, B-151 Hexagenia spp - B-42, B-49, B-112, B-160 Hitch - B-52 Homarus americanus - A-135, B-232 H. vulgaria – B-178 Hurmomya mutabilis – A-87 Huro salmoides - A-54, A-87, A-90, A-96, B-56, B-63, **B-86** Hvalella azteca – B-129 H. knickerbockeri - A-116 Hybopsis bigutta – B-9 Hyborhynchus notatus - A-35, A-53, A-87, A-90, A-96, A-97, B-41, B-57, B-86, B-218 Hydra spp - B-217 Hydracarina spp - A-116, B-58 Hydrodictyon spp - A-115, A-117, B-203, B-206 Hydrophilus spp - B-81 Hydropsyche spp - A-98, A-106, A-107, A-130, B-42, **B-112**, **B-160** H. californica - B-64, B-70, B-159, B-183, B-185 H. stenonema – A-100 Hydropsychidae - A-145 Hypentelium nigricans – B-9. N-12 Hypognathus nuchalis – B-12 Ictalurus melas - A-59, A-69, A-154, B-5, B-10, B-17, B-19, B-22, B-70, B-72, B-94, B-116, B-119, B-120,

B-122, B-137, B-138, B-154, B-161, B-163, B-179,

B-185, B-186, B-214, B-225-227, B-229

I. natalis - A-134, B-3, B-17, B-120, B-214, B-216, B-225 I. nebulosus - A-21, A-74, A-154, A-161, B-18, B-22, B-43, B-108, B-175, B-191, B-229 I. punctatus - A-3, A-68, A-71, A-82, A-92, A-101, A-111, A-112, A-156, B-6, B-7, B-16, B-17, B-19, B-22, B-32, B-42, B-69, B-71, B-72, B-85, B-95, B-102, B-103, B-114, B-120, B-121, B-125, B-137, B-160, B-163, B-166, B-172, B-180, B-188, B-189, B-191, B-192, B-197, B-201, B-206, B-213, B-218, B-232, B-233 Ictiobus cyprinellus – B-17, B-22, B-137 Infusoria spp - B-58 Ischnura spp – B-223 I. verticalis - A-116 Isonychia bicolor - B-3 Jordanella florida – B-4 Juncus spp - A-44, B-205 Justica americana - B-105 J. repens - B-105 Keratella spp - B-196, B-224, B-228 Labea synodontis – B-60 Labeo fimbriatus - B-81, B-154, B-163, B-179 L. rohita - B-38, B-81 Lagodon rhomboides - A-2, A-5, A-6, A-52-54, A-70, A-78, A-153 Lamprey - B-216, B-217 Lampsilis siliquoidae - B-13, B-78, B-166 L. ventricosa - B-13, B-78, B-166 Larus ridibundus - B-54, B-95, B-121 Leander squilla - A-40, A-132 Lebistes spp - B-41 L. reticulatus – A-4, A-6, A-16, A-17, A-20, A-21, A-29, A-34, A-38, A-40, A-47, A-49, A-51, A-52, A-63, A-64, A-72, A-77-80, A-82, A-84, A-87, A-90, A-91, A-95, A-99, A-107, A-114, A-142, A-149, A-152, A-157, A-159, A-164, A-165, B-2, B-5, B-8, B-24, B-30, B-32, B-36, B-40, B-42, B-59, B-62, B-65, B-83, B-92, B-97, B-106, B-111, B-112, B-115–117, B-124, B-126, B-129, B-136, B-139, B-140, B-150, B-157, B-164, B-167, B-176, B-183, B-198, B-210, B-215, B-217, B-223, B-228, B-233 Leersia spp - B-205 Leiostomas xanthurus - B-5, B-6, B-15, B-20, B-25, B-30, B-35, B-46, B-61, B-83, B-89, B-110, B-118, B-119, B-122, B-126, B-130, B-137, B-144, B-146, B-161, B-173-175, B-178, B-186, B-189, B-190, B-193, B-199, B-201, B-202, B-204, B-208, B-209, B-212, B-214, B-218, B-226, B-230, B-233, B-234, B-236 Lemna spp – B-203 Lemna minor - B-102, B-180 L. trisulca – A-130 Lepisosteus osseus – A-134, A-158 L. steinii – A-37 Lepomis spp - A-117, B-126 L. auritus - A-51, A-78, B-159

L. cyanellus - A-15, A-59, A-69, A-133, A-134, B-10, B-12, B-15, B-17, B-20, B-22, B-41, B-48, B-49, B-68, B-70, B-72, B-91, B-94, B-110, B-114, B-118-120, B-131, B-137, B-141, B-171, B-191, B-198, B-205-207, B-212, B-213, B-219, B-223, B-225, B-226 L. gibbosus - A-80, A-154, B-3, B-4, B-17, B-72, B-85, B-86, B-93, B-116, B-137, B-138, B-154, B-161, B-163, B-179, B-185, B-186 L. humilis - A-141, B-229 L. macrochirus - A-2-9, A-11-13, A-15-17, A-19-21, A-23-26, A-29, A-34, A-37-39, A-41, A-43, A-46-49, A-52, A-57, A-59, A-64, A-69, A-70-72, A-78, A-79, A-82, A-83, A-88, A-89, A-91, A-92, A-95, A-98-100, A-102–106, A-108–110, A-112, A-114, A-117–121, A-125, A-126, A-129, A-131, A-134, A-136-141, A-143, A-146, A-148, A-149, A-152, A-156, A-158, A-160-162, A-165, A-166, B-2-6, B-8-11, B-15-17, B-20-24, B-27, B-30-33, B-35, B-36, B-39, B-40, B-42, B-44-52, B-56, B-59, B-62, B-67-69, B-71, B-72, B-76, B-83-85, B-88-95, B-97, B-102-106, B-108, B-109, B-111-118, B-120, B-123-126, B-128-131, B-135, B-136, B-139-144, B-146, B-149, B-150, B-152, B-154-156, B-158, B-160, B-163, B-164, B-166, B-168-171, B-174-176, B-180, B-182-184, B-186, B-188, B-189, B-191, B-192, B-197-199, B-201-207, B-210-219, B-223, B-226, B-228, B-231-234 L. megalotis - B-17, B-72, B-139 L. microlophus - B-27, B-145, B-159, B-174, B-214 L. symmetricus – B-139 Leptodora spp - B-196, B-224 Leptonychotes weddelli - B-54, B-74 Leucichthys spp - B-75 Libellula spp - A-116 Limnaea palustris - A-21, A-39, A-83, A-91, A-106, A-109 Limnephilus rhombicus – B-54, B-79, B-98, B-122 Limnodrilus spp - B-31, B-73, B-96, B-161, B-186, **B-200** L. hoffmeisteri – A-46, A-130, A-164 Lobodon carcinophagus - B-54, B-71 Lota lota - B-55, B-79, B-80, B-98, B-142 Lottus asper – B-221 Lymnaea spp - A-12, A-23, A-29, A-36, A-44, A-57, A-59-62, A-81, A-103, A-115, A-123, A-125, A-126, A-131, A-135, A-138-141, A-143, A-146, B-7, B-18, B-27, B-28, B-91, B-109, B-123, B-163, B-181 L. carillandi - A-24, A-25, A-38, A-108, A-129, A-137, A-145, B-169, B-235 Lythrurus umbratilis - B-218 Macrobdella decora - A-44 Macrognathus aculeatum - B-81 Maia squinado - A-84

Mayfly – B-217

- Mastigophora spp B-58
- Mastocembelus pancalus B-81
- Mayorella palestinensis A-51, B-21, B-38
- Megaloptera B-56, B-59, B-74, B-199
- Megastomatobus cybrinella B-59
- Menidia menidia B-78
- Mercenaria mercenaria B-13, B-73, B-78, B-97, B-122,
  - B-142, B-153, B-162, B-165, B-166
- Mesocyclops obsoletus A-44
- Mesoueliidae B-104
- Microcystis spp A-47
- *M. aeruginosa* A-3, A-5, A-6, A-8, A-15, A-16, A-24, A-27, A-28, A-35, A-36, A-42, A-45, A-47, A-50, A-52, A-54–57, A-59, A-62–64, A-71–73, A-77, A-78, A-82, A-83, A-85, A-86, A-89, A-90, A-92, A-93, A-96, A-100–102, A-104, A-110, A-112, A-113, A-128, A-129, A-142, A-143, A-151, A-152, A-155, A-157, A-162, B-6, B-18, B-19, B-27–29, B-34, B-44, B-60, B-91, B-166, B-167, B-187, B-192–194, B-209, B-211, B-215, B-216
- *Micropterus dolomieu(i)* A-15, B-20, B-22, B-48, B-49, B-51, B-91, B-106, B-114, B-138, B-147, B-163, B-186, B-194, B-205, B-213, B-216, B-225
- *M. salmoides* A-7, A-24, A-46, A-51, A-66, A-82, A-104, A-125, A-134, A-136, A-138, B-2, B-12, B-15, B-17, B-21, B-22, B-31, B-32, B-34, B-35, B-39, B-40, B-44, B-52, B-57, B-59, B-72, B-81, B-84, B-85, B-102, B-103, B-107, B-109, B-112–114, B-123, B-124, B-127, B-129, B-130, B-133, B-135, B-137, B-139, B-142, B-143, B-147, B-154, B-156, B-166, B-170, B-171, B-175, B-180, B-182, B-183, B-186, B-188, B-190, B-192, B-206, B-214, B-216, B-219, B-236
- Minnows A-17, A-34, A-35, A-159, B-30, B-33, B-57, B-61, B-70, B-107, B-111, B-115, B-120, B-123, B-139, B-149, B-155, B-157, B-159, B-164, B-176, B-179, B-182
- Moina macrocopa B-115
- Mollienesia spp B-157
- M. latopinna A-99, A-123, A-126, A-131, A-146
- Mollusca B-56, B-74
- Monochrysis lutherii A-33, A-53, A-98, B-65, B-101, B-107, B-110, B-130, B-149, B-151, B-170, B-173, B-200, B-215, B-225
- Moroco steindachnerii A-72
- Morone americana A-44, B-195
- Moxostoma aureolum B-218
- M. erythrurum B-9
- Mullet B-162, B-166
- Mummichog B-162, B-166
- Mugil cephalus B-25, B-46, B-47, B-89, B-108, B-119, B-125, B-170, B-208, B-209, B-213, B-217–219, B-222
- M. curema B-30, B-61, B-73
- Mya arenaria B-13, B-97, B-122, B-142, B-165
- Mylocheilus caurinum B-220, B-221
- Myriophyllum spp A-54
- M. brasilliensen B-44, B-202

M. exalbescens – B-105 M. heterphyllum – B-44, B-134, B-202 Mystus vittatus – B-38, B-81 Mytilus edulis – A-50, B-178 Mvxocephalus scorpius - B-137, B-138, B-154, B-161, B-163, B-179, B-185, B-186 Nais spp - A-28, A-47, A-130, B-202 *Najas flexilis* – A-115, B-105, B-147 N. quadalupensis – B-19, B-38, B-45, B-211 Nandus nandus - B-38, B-81 Nassarius obsoletus – B-78 Natrix erythrogaster - B-123 N. rhombifera – B-123 Naupluis – B-12, B-38 Navicula spp - A-25, A-48, A-109, A-119 N. seminulum - B-3, B-98 Nemocheilus barbatulus - A-39 Nemertinea - B-74 Nepa spp - B-81 Nereis spp - A-40, A-132N. limnicola - B-197 Neuroptera - B-76, B-162 Nitzchia spp - A-25, A-48, A-109, A-119, B-209 N. linearis - A-4, A-13, A-23, A-26, A-43, A-89, A-100, A-103, A-104, A-108, A-121, A-126, A-131, A-146, A-161, B-3, B-5 N. palea - A-3, A-6, A-8, A-17, A-24, A-27, A-28, A-35, A-36, A-42, A-45, A-50, A-54, A-56, A-57, A-62, A-63, A-73, A-78, A-83, A-86, A-90, A-92, A-93, A-101, A-102, A-112, A-113, A-128, A-129, A-142, A-143, A-151, A-152, A-155, A-157, A-162, B-7, B-28, B-44, B-60, B-91, B-187, B-193, B-194, B-211, B-215, B-216 Nodiflorum spp – A-130 Nostoc spp – A-24, A-48, A-109, A-119 Notemigonus crysoleucas – A-46, A-69, A-73, A-80, A-96, A-117, B-10, B-16, B-23, B-39, B-40, B-42, B-43, B-56, B-59, B-68, B-84-87, B-110, B-118, B-120, B-122, B-124, B-125, B-135-137, B-139, B-143, B-156, B-158, B-168, B-175, B-182, B-184, B-188, B-191, B-216, B-219, B-226, B-227, B-229 Notropis spp - B-223N. ardens - B-4, B-18 N. atherinoides - A-54, A-87, A-90, A-96, B-4, B-5, B-149 N. blennius – B-11 N. chrysocephalus - B-9, B-94 N. cornutus - A-80, B-5, B-11, B-85, B-149 N. heterolepis - B-216 N. hudsonius – A-116 N. lutrensis - B-214 N. maculatus – B-224 N. spilopterus – B-9 N. stramineus - B-4, B-9, B-18 N. umbratilis - A-141, B-9, B-11, B-174, B-214

N. volucellus - B-9 N. whipplii - A-141 Noturus miurus – B-12 Nymphea spp - A-44, A-115, B-44, B-145, B-205 Odonata - B-56, B-59, B-74, B-76, B-104, B-148 Oedogonium spp - A-115 Oenanthe fluviatilis – A-130 Oligochaeta - B-49, B-74, B-103, B-168 Onchorhynchus garbuscha – B-227 O. keta – B-227 O. kisutch - A-59, A-71, A-87, A-125, A-136, A-142, A-146, A-160, B-8, B-19, B-28, B-31, B-33, B-36, B-41, B-62, B-64, B-73, B-89, B-93, B-107, B-109, B-116, B-117, B-135, B-140, B-143, B-147, B-150, B-164, B-170, B-188, B-191, B-199, B-224, B-227 O. nerka - B-221, B-223, B-227 O. tshawytscha - A-59, A-71, A-87, A-125, A-136, A-146, B-6, B-8, B-21, B-36, B-64, B-93, B-101, B-113, B-116, B-117, B-128, B-135, B-140, B-147, B-150, B-156, B-164, B-176, B-191, B-206, B-213, B-224 Oocystis spp – A-24, A-48, A-109, A-119 Ophicephalus punctatus - B-68, B-93, B-151 Opsopoeodus emibiae – B-139 Orconectes rusticus – A-40, B-3 Osio otus - B-55, B-75, B-96, B-121, B-143 Oscillatoria spp - A-24, A-48, A-109, A-119, B-56, B-79 Osmeridae – B-227 Ostracoda - A-47, B-66, B-104 Oyster - A-5, B-5, B-6, B-12, B-13, B-15, B-20, B-27, B-32, B-50, B-55, B-69, B-71, B-76, B-86, B-89, B-91, B-97, B-98, B-103, B-120, B-122, B-142, B-143, B-146, B-147, B-153, B-163, B-175, B-187, B-193, B-201, B-202, B-208-210, B-212, B-234-236 Paleomonetes kadiakensis - B-10, B-94, B-119, B-226 P. paludosus - B-2, B-129 Pandalus montagni - B-32, B-40, B-85, B-125, B-133, B-134, B-189, B-207 Panicum hemitomum – B-15, B-138 Pandorina spp - A-25, A-48, A-109, A-119 Paracheinodon innesi – A-58 Paralichthys dentatus - B-78 Paramecium spp – A-116 Parrot's feather - A-117, B-145, B-153 Paspalum spp - B-205 Pastella vulgata – B-178 Peneas aztecus - A-5, B-15, B-20, B-46, B-50, B-54, B-82, B-83, B-87, B-107, B-126, B-130, B-143, B-144, B-146, B-154, B-175, B-176, B-180, B-181, B-193, B-201, B-202, B-204, B-208, B-209, B-212, B-213, B-219, B-231, B-233-236 P. cardinadus – B-198 P. duorarum - B-46, B-83, B-189, B-190, B-201, B-202, B-218 P. setiferus - B-83, B-103, B-125, B-170, B-174, B-201, **B-218**, **B-231** Perca flavescens - A-44, A-59, A-80, A-154, B-17, B-22,

B-72, B-86, B-159, B-195, B-216, B-221, B-229

P. fluviatilis - A-9, A-10, A-166 Perch – B-157, B-160 Percina caprodes - A-154 P. maculata – B-9 Peridinium trochoideum - B-80 Periphyton – A-165 Pescilia latipinna – A-58 Pestia stradiotes - B-18, B-153 Petromyzon marinus - A-19, A-33, A-55, A-65, A-149, A-153-155, B-101, B-216, B-228 Phaeodactylum tricornutum - A-33, A-53, A-98, B-65, B-107, B-110, B-130, B-149, B-151, B-170, B-173, B-200, B-215, B-225 Phalacrococax spp - B-178 Pholis leata – B-227 Phormodinium inundatum - A-47 Phoxinus phoxinus - A-25, A-49, A-79, A-97, A-115, A-134, A-135, A-143, B-13, B-20, B-206, B-235 Phryganea spp - B-197 Physa heterostropha – A-13, A-46, A-51, A-89, A-100, A-103-105, A-108, A-130, A-161, A-163, A-164, B-3, B-5 Physidae - B-104, B-227 Phytoplankton - see plankton Pimephales notatus - A-141, B-117, B-214, B-227, B-229 P. promelas - A-4, A-6, A-9, A-15-17, A-21, A-22, A-29, A-34, A-35, A-38, A-41, A-45-47, A-49, A-52, A-55, A-64, A-66, A-72, A-77-79, A-87, A-88, A-91, A-95, A-99, A-102, A-104, A-117, A-134, A-142, A-148, A-149, A-152, A-156, A-159, A-160, A-165, A-166, B-4, B-5, B-8, B-9, B-17, B-22, B-24, B-30, B-32, B-36, B-39-41, B-43, B-48-50, B-57, B-59, B-62, B-72, B-81, B-83, B-86, B-87, B-92, B-95, B-100-103, B-106, B-111, B-114-116, B-123, B-124, B-126, B-131, B-135, B-139, B-140, B-143, B-149, B-150, B-155, B-157, B-162, B-164, B-167, B-169, B-174-176, B-179, B-180, B-182-184, B-190, B-193, B-195, B-196, B-198, B-203, B-210, B-215-217, B-220, B-223, B-225, B-230, B-233, B-235 Pithophora spp - A-117, B-84 Planorbidae - B-104 Plankton - A-5, A-43, A-118, B-15, B-20, B-38, B-46, B-50, B-103, B-108, B-125, B-126, B-129, B-130, B-144, B-146, B-148, B-154, B-168, B-174, B-175, B-180, B-196, B-203, B-206, B-212, B-218, B-219, B-233, B-236 Platalea leucorodia - B-54, B-96 Platyhelminths - A-163 Plecoptera - B-73, B-74, B-76, B-148, B-162, B-199, B-200 Plectonema spp - A-24, A-48, A-109, A-119 Pleurobena cordatum – B-47 Pleuronectes spp - B-178 Pleuronectes platessa - A-110

Poecilia reticulata - A-58, B-95, B-97 Poecilichthys exilis - B-86 Polyarthra spp - B-196, B-222, B-228 Polycelis nigra – A-114, A-120, A-124, A-128, A-131, A-135-140, A-143, A-147-149 Pomoxis annularis - A-5, A-51, A-78, B-229 P. nigromaculatus - A-54, A-87, A-90, A-96, B-41, B-59, B-116, B-139 Pondweed - B-113, B-153 Pontederia spp - A-44, B-126 P. cordata - B-45 Pontoporeria affinis - B-75 Potomogeton spp - A-44, A-118, B-202, B-203, B-225 P. crispus – A-115, B-105 P. densus – B-14 P. foliosus - A-115, B-105 P. nodosus - A-5, A-110, A-153, B-102, B-189, B-235 P. pectinatus - A-5, A-110, A-130, A-153, B-14, B-56, B-80, B-102, B-105, B-189, B-235 P. pusillus - B-105 P. annularis -B-17Procambarus clarki - A-87, A-102, A-119, B-10, B-21, B-32, B-66, B-89, B-117, B-168, B-187, B-200 P. simulons – B-24 Prosimulum spp - B-2, B-14, B-26, B-75, B-82, B-88, B-90, B-99, B-100, B-110, B-129, B-131, B-133, B-144, B-149, B-165, B-175, B-176, B-178 Protococcus spp - A-33, A-53, A-98, B-101, B-107, B-110, B-130, B-149, B-151, B-170, B-173, B-200, B-215, B-225 Pseudemys scripta elegans – B-123 Pseudomones pisicida – B-76 Pseudopleuronectes americanus – B-137, B-138, B-154, B-161, B-163, B-186 Psidium idahoense - A-45 Pteronarcella badia - B-34, B-80, B-98, B-110, B-142, B-162, B-186, B-230, B-231 Pteronarcys spp - A-55, A-60, A-118, B-14, B-25, B-26, B-51, B-81, B-82, B-87, B-99, B-111, B-126-128, B-144, B-147, B-151, B-165, B-166, B-181, B-184, B-185, B-187, B-191, B-197, B-200, B-209 P. californica - A-102, A-118, A-119, B-2, B-10, B-11, B-13, B-14, B-22-25, B-32, B-34, B-38, B-48, B-50, B-51, B-54, B-63, B-64, B-68-70, B-80, B-82, B-83, B-86, B-88, B-91, B-94, B-95, B-98, B-99, B-106-108, B-110-112, B-118-120, B-123, B-126, B-128-130, B-136, B-137, B-141, B-142, B-144, B-152, B-156, B-158-160, B-162, B-169, B-173, B-179, B-181, B-183, B-186-189, B-192, B-197, B-201, B-204, B-205, B-210, B-228, B-230-232, **B-236** Ptychocheilus oregonensis - A-86, A-87, B-220-222 Pungitius pungitius – B-194 Puntius javanicus – B-152 P. puckelli - B-5, B-31, B-77, B-122 Pungtungia herzi – A-72 Puntius sophore – B-38, B-81 Pygosciles adeloriae - B-54, B-71

Pygosteus pungitius – A-31, A-44, A-63, A-68, A-83, A-164 Pylodictis olivaris - B-22 Pyramimonas spp - B-80 Rainbow trout - A-10, A-12, A-25, A-27, A-37, A-40, A-99, B-2, B-10, B-14, B-18, B-22, B-24-26, B-28, B-30, B-34, B-37, B-49, B-52, B-59, B-66, B-69, B-82, B-87, B-89, B-90, B-94, B-100, B-111, B-112, B-118, B-127-129, B-141, B-144, B-146, B-151, B-157, B-169, B-172, B-179, B-181, B-216, B-217, B-219, B-226, B-230, B-233 Rana spp - A-16R. catesbeiana - A-69, B-6, B-9, B-24-26, B-34, B-37, B-66, B-117, B-123, B-127, B-132, B-133, B-137, B-141, B-146, B-151, B-158, B-173, B-217, B-225, B-233 R. pipens - A-46, A-69, A-81, B-23 R. temporaria - A-134, B-93 Ranatia filiformis – B-81 Rangia cuneata - B-97, B-122, B-142, B-153, B-165 Ranunculus spp - B-105 R. pseudofluitans – B-13 Rasbora heteromorpha - A-82, A-99, A-106, A-147, B-73, B-97 / Rhigophila dearborni – B-74 Rhinichthys atratulus - A-80, A-104, A-105, B-79 Richardsonius balteatus - B-18, B-109, B-140, B-145, B-156, B-188, B-207, B-220-222 Rita rita - B-81 Roccus saxatilis - B-231 Rotifers - A-47, A-54, A-117, B-2, B-66, B-129 Rutilus rutilus - A-9, A-10, A-166 Saccharomyces ellipsoides - A-132 Sagittaria spp - A-118 Sagittaria latifolia - B-105, B-203 Salmo spp - B-97 S. clarkii - A-71, A-87, A-125, A-136, A-146, B-63, B-68, B-169 S. gairdneri(i) - A-9-13, A-16, A-18-21, A-28-34, A-37, A-41, A-42, A-46, A-48, A-49, A-53, A-55, A-56, A-58-60, A-62, A-63, A-67, A-68, A-74, A-75, A-80, A-82, A-85, A-86, A-88, A-92-96, A-98, A-100, A-101, A-104-106, A-111, A-112, A-118, A-128, A-151, A-153-155, A-160, A-163-166, B-5, B-6, B-8, B-9, B-16, B-17, B-22, B-26, B-32, B-36, B-39, B-41, B-42, B-44, B-48, B-52, B-53, B-60, B-63, B-64, B-66, B-67, B-69, B-71, B-72, B-81, B-86, B-88-90, B-92, B-95, B-100, B-101, B-103, B-104, B-108, B-111, B-116, B-117, B-120, B-125, B-127, B-128, B-131, B-133, B-135, B-140, B-141, B-144, B-145, B-150, B-152, B-157-160, B-163-166, B-169, B-171, B-172, B-174, B-180, B-187-189, B-191, B-192, B-197, B-199-201, B-204, B-216, B-218, B-221, B-222, B-224, B-225, B-227-229, B-231, B-232 S. salar - A-40, A-43, A-47, A-49, A-100, A-158, A-160,

A-165, B-70, B-76, B-79 C-9

S. trutta - A-19, A-33, A-46, A-49, A-68, A-81, A-82, A-92, A-112, A-134, A-153–155, A-165, B-6, B-16, B-17, B-31, B-42, B-57, B-60, B-63, B-65, B-72, B-84-86, B-125, B-131, B-139, B-163, B-166, B-171, B-172, B-174, B-188, B-189, B-192, B-197, B-218, **B-232** Salmon - A-5, A-43, A-160, B-15, B-51, B-54, B-60, B-61, B-63, B-77, B-105, B-108, B-147, B-155, B-157, B-162, B-204 Salvelinus fontinalis - A-46, A-59, A-68, A-82, A-92, A-112, A-154, B-6, B-16, B-22, B-60, B-63, B-79, B-80, B-84-86, B-98, B-104, B-125, B-131, B-132, B-163, B-166, B-171, B-172, B-174, B-188, B-189, B-192, B-194, B-197, B-218, B-227, B-232 S. namaycush - A-59, A-68, A-82, A-92, A-112, B-6, B-16, B-42, B-72, B-125, B-166, B-171, B-172, B-174, B-189, B-192, B-197, B-232 Sarcocheilichthys variegratus - A-72 Sarcodina – B-58 Scardinius erythrophthalmus – A-9 Scenedesmus spp - A-24, A-48, A-109, A-119, B-58 S. incrassulatus – B-223 S. obliquus - A-3, A-6, A-8, A-17, A-24, A-27, A-28, A-35-37, A-42, A-45, A-50, A-54, A-56, A-57, A-62, A-63, A-73, A-78, A-83, A-86, A-90, A-92, A-93, A-101, A-102, A-112, A-113, A-128, A-129, A-142, A-143, A-151, A-152, A-155, A-157, A-162, B-7, B-28, B-29, B-44, B-60, B-91, B-187, B-193, B-194, B-211, B-215, **B-216** Scirpus spp - A-44 S. acutus - B-105 S. validus – A-115 Scophiopus hammondi - B-132, B-168 Sea lamprey - A-19, A-30, A-58, A-67, A-68, A-74, A-75, A-85, A-86 Semotilus atromaculatus - A-3, A-12, A-14, A-17, A-20, A-52, A-53, A-56, A-57, A-62, A-64, A-70, A-76, A-77, A-80, A-85, A-88, A-97, A-109, A-110, A-136, A-139-141, A-152, A-154, A-156, B-9, B-32, B-79, B-85, B-98, B-113, B-175, B-225 Sesarma africanum – B-31 Sewage organisms - A-2, A-4, A-5, A-8, A-10, A-18, A-22, A-32, A-36-39, A-45, A-46, A-49, A-56, A-57, A-61, A-66, A-68, A-69, A-84, A-85, A-88, A-90, A-92, A-93, A-95, A-98, A-102, A-104, A-105, A-107, A-113, A-114, A-116, A-120, A-131–133, A-137, A-139, A-141, A-150, A-152, A-155, A-157, A-160-162, A-164, B-185, B-210 Shiners - A-26, A-76, A-97, B-33, B-57, B-134, B-151, B-160, B-163 Shrimp – B-78, B-133 Sialis spp - B-55, B-79, B-98, B-122 Simocephalus serrulatus - A-29, A-50, A-118, A-149, B-12, B-14, B-19, B-34, B-37, B-48, B-51, B-53, B-71, B-72, B-81, B-82, B-88-90, B-95, B-96, B-101, B-108, B-120, B-121, B-127-129, B-141, B-145, B-147, B-152, B-161, B-165, B-181, B-185, B-187, B-188, B-191, B-197,

B-200, B-201, B-204, B-228, B-231, B-232, B-236

Simulium spp - B-3, B-14, B-27, B-33, B-60, B-75, B-82, B-88, B-90, B-91, B-99, B-100, B-110, B-129, B-131, B-133, B-149, B-161, B-165, B-173, B-177, B-178, B-181 Siphlonurus spp – B-197 Skeletonema costatum – B-80 Snails - A-17, B-91, B-181 Somateria mollissima – B-54, B-96, B-121 Spartina patens – B-78 Spatterdock – B-14, B-38, B-45, B-138, B-147, B-153, B-211, B-237 Sphaerium c.f. tenue – A-130 Sphaerodema annulatum – B-81 Spirodela polyrhyza – B-102, B-180 Spirogyra spp - A-116, A-118 Sterna hirundo - B-121 S. sandvicensis – B-54, B-96, B-121 Stenonema spp – A-98, A-106, A-107, A-130, B-3 S. ares - A-145, B-3 S. heterotarsale – A-145, B-3 S. hirundo - B-54 Stigeoclonium spp - A-24, A-48, A-109, A-119 Stizostedion vitreum – B-105, B-157, B-171, B-216, B-229 Strix aluco - B-53, B-54, B-75, B-96, B-120, B-143, B-152 Stylonichia spp – A-116 Suckers – B-62, B-64, B-67, B-77, B-195 Sula bassana – B-178 Sunfish – A-9, A-21, A-26, A-36, A-52, A-76, A-95, A-134, B-34, B-37, B-57, B-83, B-128, B-140, B-146, B-157, B-160, B-216, B-220 Synedra spp - B-58 Synodontis schall – B-60 Tadorna tadorna – B-54, B-96, B-121 Tadpoles – A-68, A-76, A-141, B-81 Talifridae – B-104 Tench – A-47 Tendipes decorus – A-46, A-164 T. plumosus – A-45 Tendipidae – B-49, B-149, B-220 Tilapia spp - A-69 T. massambica – B-22, B-82, B-131, B-152, B-154, B-163, B-179, B-198 T. melanopleura - B-8, B-92, B-134 Tinca tinca – B-234 Trachelomonas spp – B-58 Trematomus bernacchii – B-74 T. hansoni – B-74 Tribonema spp - A-25, A-48, A-109, A-119 Trichogaster fasciatus - B-81, B-93, B-151 Trichoptera – A-47, B-59, B-61, B-64, B-73, B-74, B-76, B-162, B-199

Trout – A-9, A-26, A-27, A-51, A-53, A-99, A-104, A-111, A-113, A-117, A-118, A-135, A-158, A-164, B-52, B-54, B-61, B-62, B-64, B-65, B-67, B-71, B-77, B-148, B-163, B-172, B-181, B-187, B-192, B-195, B-199-201, B-204, B-206, B-207 Tubifex spp – B-31, B-73, B-96, B-161, B-186, B-200 Tubificids - A-80, A-142, A-166 Turbellaria spp – B-74, B-217 Typha angustifolia – B-105 T. latifolia - B-16, B-45, B-105 Tyto alba - B-53, B-54, B-75, B-96, B-120, B-142, B-152 *Umbra limi* – B-85, B-86 Utricularia spp - B-44, B-134, B-202 Vascular plants - B-37, B-54 Vaucheria spp - B-14 Vellidae – B-104 Venus japonica – A-87 Volvox spp – B-66 Walleye - A-113 Warmouth - B-135 Water hyssop – A-117 Water lettuce -B-131, B-233Water plants – A-17 Whitefish - B-61, B-67 Wolffia columbiana – B-102, B-180 Wolffiella floridana – B-180 Xiphophorus maculatus – A-58 Yellow perch – B-220 Zaccho platypus – A-72 Zebrafish - A-4Zooplankton - see plankton Zygnema spp - A-24, A-48, A-109, A-118, A-119, B-206 APPENDIX D

IDENTIFICATION OF COMMERCIAL CHEMICALS

# APPENDIX D

Trade Name	Chemical Name or Active Ingredient
2,4D	2,4-dichlorophenoxyacetic acid
2,4,5-T	2,4,5-trichlorophenoxyacetic acid
Abate®	0,0,0',0'-tetramethyl 0,0'-thiodi-p-phenylene phosphorothioate
ABS	Alkyl benzene sulfonate
AC 5727	m-isopropylphenyl-N-methylcarbamate
AC 12009	No information available
AC 38023	o,o-dimethyl o,p-(dimethylsulfamoyl) phenyl phosphorothioate
AC 43064	Cyclic ethylene (diethoxyphosphinothioyl) dithiomidocarbonate
AC 43913	See Abate
AC 47031	Cyclic ethylene (diethoxyphosphinyl) dithioimidocarbonate
AC 47921 EC4	No information available
AC 52160	No information available
ACP-M-569	Contains 3-amino-1,2,4-triazole
Acriflavine	A mixture of 2,8-diamino-10-methylacridinium chloride and 2,8-diaminoacridine
Acrolein	Acrylic aldehyde
Acti-dione®	See cycloheximide
Aerosporin	Polymyxin B
Aldrin (Octalene <sup>®</sup> )	1,2,3,4,10,10-hexachloro-1,4,4a,5,8,8a-hexahydro- 1,4-endo-exo-5,8-dimethanonaphthalene
Algeeclear	No information available
Algibiol	No information available
Algimaster	Alkyl quaternary ammonium bromides, organic polyamine, amine hydrobromides
Algimycin	No information available
Alticide	Sodium chlorate
Ametryne (Ametryn <sup>®</sup> )	2-ethylamino-4-isopropylamino-6-merthyl- mercapto-s-triazine
Aminotriazole	See Amitrole
Amitrole (Aminotriazole, Amitrol)	3-amino-1,2,4-triazole
Amitrol T	3-amino-1,2,4-triazole-ammonium thiocyanate mixture
Amiton	o,o-diethyl 5,2-diethylaminoethyl phosphorothioate
Ammate	Ammonium sulfamate
Amopyroquin	No information available
	No information available

Trade Name	Chemical Name or Active Ingredient
Aquaherb	o-dichlorobenzene and aromatic salt
Aqualin	85% acrolein
Aquathol <sup>®</sup>	Disodium salt of endothal (19.2%-H-Pennsalt)
Aramite <sup>®</sup>	2-(p-tert-butylphenoxy) isopropyl-2'-chloroethyl sulfite
Atabrine	6-chloro-9{[4-(diethylamino)-1-methylbutyl] amino} 2-methoxyacridine dihydrochloride
Atlacide-2,4-D	Sodium chlorate-2,3-dichloroxyacetic acid
Atlas 1901	No information available
Atlas A	Sodium arsenate
Atlox	A series of pesticide emulsifiers
Atrazine (Gesaprim <sup>®</sup> )	2-chloro-4-ethylamino-6-isopropylamino-s-triazine
Banvel-D <sup>®</sup>	See Dicamba
Baron®	See Erbon
Barthrin	6-chloropiperonyl chrysanthemumate
Bayer 73	5,2'-dichloro-4'-nitrosalicylanilide
Bayer 4731	No information available
Bayer 9018	No information available
Bayer 22408	o,o-diethyl-o-naphthylamido phosphorothioate
Bayer 25141	o,o-diethyl-o,p-(methylsulfinyl)phenyl phosphorothioate
Bayer 25198	o,o-dimethyl-o-(p-methylsulfinylphenyl) phosphorothioate
Bayer 29492	o,o-diethyl o-(4-methylthio-m-tolyl) phosphorothioate
Bayer 29493 (Baytex <sup>®</sup> & Fenthion)	o,o-dimethyl-o-[4-(methylthio)-m-tolyl] phosphorothioate
Bayer 29952	o-ethyl-o-(p-methylthio) phenyl methyl- phosphonothioate
Bayer 30749	No information available
Bayer 34042	o-ethyl o-(4-methylthio-m-tolyl) methyl phosphoramidothioate
Bayer 37289	No information available
Bayer 37342	o,o-dimethyl o-(3,5-dimethyl-4-methyl-thiophenyl) phosphorothioate
Bayer 37343	No information available
Bayer 37344	4-(methylthio)-3,5-xylyl methylcarbamate
Bayer 38156	o-ethyl-S-p-methylphenyl ethylphosphonodithioate
Bayer 38819	No information available
Bayer 38920	6,7,8,9,10,10-hexachloro-1,5,5a,6,9,9a-hexahydro- 3-methyl-6,8-methano-2,4-benzodioxathiepin
Bayer 41831	o,o-dimethyl-o-4-nitro-m-tolyl) phosphorothioate

Trade Name	Chemical Name or Active Ingredient
Bayer 44646	4-dimethylamino-3-tolyl N-methyl-carbamate
Bayer 46676	o-ethyl-o-2-ethylthio-4-methyl-6-pyrimidyl ethyl- phosphonothioate
Bayer 47940	o,o-dimethyl o-(3-chloro-4-cyanophenyl)- thionophosphate
Bayer 52957	o,o-diethyl o-5-chlorobenzisoxazolyl-3-phos- phorothioate
Baygon®	o-isopropoxyphenyl methyl carbamate
Baytex®	See Bayer 29493
Ben Venue #35	No information available
Ben Venue #3835	No information available
Ben Venue #52	No information available
Ben Venue #54	No information available
BHC (HCK, Hexyclan)	1,2,3,4,5,6-hexachloro-cyclohexane (benzene hexachloride)
Bidrin®	3-(dimethoxyphosphinyloxy)-N,N,dimethyl-cis- crotonamide
Bomyl <sup>®</sup>	Dimethyl-1,3-di(carbomethoxy)-1-propen-2yl phosphate
Borate	Boron trioxide
BP 1002	No information available
Buramine	Crude N-mono-n-butyl urea
C 56	Hexachlorocyclopentadiene
C 2059	n-(3-trifluoro-methylphenyl) n',n'-dimethylurea
C8514	No information available
Camphene	2,2-dimethyl-3-methylenenorbornane
Captan (Orthocide <sup>®</sup> )	n-trichloromethylthio-4-cyclohexene-1,2- dicarboximide
Carbaryl (Sevin <sup>®</sup> )	1-naphthyl-N-methyl-carbamate
Carbophenothion (Trithion <sup>®</sup> )	S- { [(p-chlorophenyl)thio] methyl } o,o-diethyl phosphorodithioate
Casoron®	See Dichlobenil
Catechol	o-dihydroxybenzene
Cela S-1942	o,o-dimethyl o-(2,5-dichloro-4-bromophenyl) thionophosphate
Cela S-2225	o,o-diethyl o-(2,5-dichloro-4-bromophenyl) thionophosphate
Ceresan	Ethylmercuric chloride
Chemagro 4497	No information available
Chem Ban	See Nabam
Chem-Fish Special	Rotenone
Chem Mite	Xylene, p-chlorophenol, p-chlorobenzene sulphonate, and rotenone

Chem Sen Chlordane (Octachlor<sup>®</sup>, Octa-Klor<sup>®</sup>, Chlordan, Velsicol 1068<sup>®</sup>) Chlorea Chloretone Chlorobenzilate Chlorothion Chloroxuron (Tenoran<sup>®</sup>) Chlorox CIPC Cleanosol CMU Conco LCP-12 Co-Ral® Crop Rider Cryolite Cube root Cumate Cyanamid 12009 Cycloheximide (Actidione<sup>®</sup>) Cygon Dacthal® Dalapon DBrDT DDD DDE DDT (Anofex<sup>®</sup>, Dinocide<sup>®</sup>, Gesapon<sup>®</sup>, Cesarex<sup>®</sup>, Gesarol<sup>®</sup>, Guesapon<sup>®</sup>, Guesarol<sup>®</sup>, Gyron<sup>®</sup>, Ixodex<sup>®</sup>, Neocid<sup>®</sup>, Zerdane, DND, GNB, GNB-A DDVP (Dichlorvos, Vapona<sup>®</sup>) Dead X Deet (Delphene<sup>®</sup>, Meta-delphene<sup>®</sup>) DEF<sup>®</sup> Dekafos®

Delrad

Sodium arsenite 1,2,3,5,6,7,8,8-octachloro-2,3,3a,4,7,7a-hexahydro-4,7 methanoindene Sodium chlorate, sodium metholate, and 3-(p-chlorophenyl)-1,1-dimethylurea) Trichloro-tert-butyl alcohol Ethyl 4,4'-dichlorobenzilate o,o-dimethyl o-(3-chloro-4-nitrophenyl) phosphorothioate N'-(4-chlorophenoxy) phenyl N,N-dimethylurea Sodium hypochlorite Isopropyl N-(3-chlorophenyl)-carbamate No information available See Monuron No information available o,o-diethyl o-3-chloro-4-methyl-1-oxo-2H-1-benzopyran-7-yl phosphorothioate No information available Sodium aluminofluoride See Rotenone 50% active copper salt of zimate No information available 3-[2-(3,5-dimethyl-2-oxycyclohexyl)-2-hydroxyethyl] glutarimide See Dimethoate Dimethyl ester of tetrachloroterephthalic acid 2,2-dichloropropionic acid 1,1,1-trichloro-2,2-bis(p-bromophenyl) ethane See TDE Dichlorodiphenyl dichloroethylene  $\alpha$ -bis (p-chlorophenyl) B,B,B-trichloroethane

o,o-dimethyl-o-2,2-dichlorovinyl phosphate No information available N,N-diethyl-m-toluamide S,S,S-Tributylphosphorotrithioate 3-pentadecylphenol o,o-diethylthionophosphate See Dioxathion Dehydroabiethylamine acetate

۰- <sub>م</sub>

Trade Name	Chemical Name or Active Ingredient
Demeton (Systox <sup>®</sup> , Bayer 8173, Isosystov <sup>®</sup> )	Mixture of 0,0-diethyl 0-2-(ethylthio) ethyl phos- phorothioate and 0,0-diethyl S-2 (ethylthio) ethyl phosphorothioate
Dermol	No information available
Derris	See Rotenone
Dexon®	p-dimethylaminobenzenediazo sodium sulfonate
Diazinon (Basudin <sup>®</sup> )	o,o-diethyl o-(2-isopropyl 4-methyl-6-pyrimidyl) phosphorothioate
Dibrom®	See Naled
Dicamba (Banvel D <sup>®</sup> , Velsicol)	3,6-dichloro-o-anisic acid 1
Dicapthon	O-(2-chloro-4-nitrophenyl) 0,0-dimethyl phosphorothioate
Dichlobenil (Casoron <sup>®</sup> )	2,6-dichlorobenzonitrile
Dichlone (Phygon <sup>®</sup> )	2,3-dichloro-1,4-naphthoquinone
Dichlorvos	See DDVP
Dicofol (Kelthane <sup>®</sup> )	4,4'-dichloro-a-trichloromethylbenzydrol
DIDT	DDT analogue
Dieldrin	1,2,3,4,10,10-hexachloro-6,7-epoxy-1,4,4a,5,6,7,8, 8a-octahydro-1,4-endo-exo-5,8-dimethanonaphthalene
Difolatan	n-(1,1,2,2-tetrachloroethylthio)-4-cyclohexene 1,2-dicarboximide
Dilan	(A mixture of Bulan <sup>®</sup> [2-nitro-1,1-bis(p-chlorophenyl) butane] and Prolan <sup>®</sup> [2-nitro-1,1-bis- (p-chlorophenyl) propane]
Dimecron	See Phosphamidon
Dimethoate (Fostion MM <sup>®</sup> , Cygon <sup>®</sup> , Rogar <sup>®</sup> )	o,o-dimethyl S-(N-methylcarbamoyl-methyl) phosphorodithioate
Dimethrin	2,4-dimethylbenzyl 2,2-dimethyl-3-(2-methyl- propenyl) cyclopropane carboxylate
Dimetilan <sup>®</sup>	2-dimethylcarbamyl-3-methylpyrazolyl-(5)- dimethylcarbamate
Dinitrocresol (Sinox <sup>®</sup> , Elgetal <sup>®</sup> )	4,6-dinitro-o-cresol
Dioxathion (Delnav <sup>®</sup> , Navadel <sup>®</sup> , Hercules 528)	2,3-p-dioxane 5, S-bis-(0,0-diethyl- phosphorodithioate)
Diphenamid (Dymid <sup>®</sup> )	n,n-dimethyl 2,2-diphenylacetamide
Dipterex®	See Trichlorofon
Diquat (Reglone <sup>®</sup> , FB/2 <sup>®</sup> )	1,1'-ethylene-2,2'-dipyridinium dibromide
Disulfoton (Di-Systom <sup>®</sup> , Dithiosystox <sup>®</sup> , Frumin Al <sup>®</sup> , Solvirex <sup>®</sup> , Frumin G <sup>®</sup> )	o,o-diethyl S-2-(ethylthio) ethyl phosphorodithioate
Di-Syston	See Disulfoton
Dithane D-14 <sup>®</sup>	See Nabam
Diuron (Karmex <sup>®</sup> , Marmer <sup>®</sup> )	3-(3,4-dichlorophenyl)-1,1-dimethylurea

Chemical Name or Active Ingredient

Dow K-6882	o-ethyl o-(2,4,5-trichlorophenyl) methyl phosphoramidothioate
Dowacide	Sodium 2,2-dichloropropionate
DPT	DDT analogue
Drummer	No information available
Dursban®	o,o-diethyl o-3,5,6-trichloro-2-pyridyl phosphorothioate
DVP-iodine	No information available
Dylox®	See Trichlorofon
Dymid	See diphenamid
EDB	1,2-Dibromoethane
Endosulfan (Thiodan <sup>®</sup> , Malix <sup>®</sup> )	6,7,8,9,10,10-hexachloro-1,5,5a,6,9,9a-hexahydro- 6,9-methano-2,3,4-benzodioxathiepin 3-oxide
Endothall (Endothal <sup>®</sup> )	7-oxabicyclo-[2.2.1]-heptane-2,3-dicarboxylic acid
Endrin	1,2,3,4,10,10-hexachloro-6,7-epoxy-1,4,4a,5,6,7,8, 8a-octahydro-1,4-endo-endo-5,8-dimethano- naphthalene
EPN	o-ethyl o,p-nitrophenyl phenylphosphonothioate
Eptam®	S-ethyl di-N,N-propylthiocarbamate
Erbon (Baron <sup>®</sup> )	2-(2,4,5-trichlorophenoxy) ethyl-2,2-dichloro- propionate
Essolvene	No information available
Esteron 99®	Propylene glycol butyl ether esters of 2,4-D
Ethion (Nialate <sup>®</sup> , Niagaia <sup>®</sup> )	0,0,0',0'-tetraethyl-S,S'-methylene bis- phosphorodithioate
Exalgae	Quaternary ammonium compounds
Fairfield 279	No information available
Fairfield OT 60-6	No information available
F-98	See Acrolein
Fenac	2,3,6-trichlorophenyl-acetic acid
Fenthion	See Bayer 29493
Fenuron TCA (Urab <sup>®</sup> )	3-phenyl-1,1-dimethylurea trichloroacetate
Ferbam (Fermate <sup>®</sup> )	See Ferbam
Flagyl	1-(2-hydroxyethyl)-2-methyl-5-nitroimidazole
Folidol®	See Parathion
Folithion®	o,o-dimethyl o-(4-nitro-m-tolyl) phosphorothioate
Forron	2,3,5-trichlorophenoxyacetic acid propylene glycol butyl ether esters
Fostion	See Dimethoate
Furazolidone	3-(5-nitrofurfurylideneamino)-2-oxazolidinone
Furoxone	N-5-nitro-2-furfurylidene-3-amino-2-oxazolidone

Trade Name	Chemical Name or Active Ingredient	
G 27365	No information available	
G 28029	See Phencapton	
G 30493	o,o-dimethyl S-(3,4-dichlorophenylthio) methyl phosphorodithioate	
G 30494	See Methyl phencapton	
Gamlen CW	No information available	
Gamlen D	No information available	
Gamlen OSR	No information available	
Gammexane®	See Lindane	
Gamosol Solvent D	No information available	
Garlon®	50.8% Dalapon and 7.7% 2-(2,4,5-trichlorophenoxy) propionic acid, propylene glycol butyl esters	
GC-3582	1-(2,5-dichlorophenyl)-2,2-dichlorovinyl diethyl phosphate	
GC-3583	2-chloro-1-(2,5-dichlorophenyl)-vinyl diethyl phosphate	
GC-3707	dimethyl-1,3-di(carbomethoxy)-1-propen-2yl phosphate	
GC-4072	Diethyl-1-(2,4-dichlorophenyl)-2-chlorovinyl phosphate	
GC-9160	δ-(5-hydroxy-1,2,3,4,6,7,8,9,10,10-decachloro- pentacyclo decyl) ethyl levulinate	
GC-9879	$\alpha$ (diethoxyphosphinothioylthio) $\gamma$ -butyrolacetone	
GS-12968	o,o-dimethyl-S [5-ethoxy-1,3,4-thiodiazol-2(3H)- onyl-(3)-methyl] -dithiophosphate	
GS-13005	o,o-dimethyl-S-[2-methoxy-1,3,4-thiodiazol-5(4H)- onyl-(4)-methyl]-dithiophosphate	
Guthion®	o,o-dimethyl S[4-oxo-1,2,3-benzotriazin-3(4H)- ylmethyl] phosphorodithioate	
Gutoxon	See Guthion	
НСК	See BHC	
Hept	Hexaethyl tetraphosphate	
Heptachlor (Velsicol <sup>®</sup> 104)	1,4,5,6,7,8,8-heptachloro-3a,4,7,7a-tetrahydro- 4,7-endo-methanoindene	
Hercules 528	See Dioxathion	
Hercules 3895 G	2,2-bis(ethylthio)-vinyl diethylphosphate	
Hercules 7175	1-(chloro-2-norbornyl)-3,3-dimethylurea	
Hercules 7531	No information available	
Hexyclan	See BHC	
Hyamine 1622	p-diisobutyl phenoxy ethoxy dimethyl benzyl ammonium chloride	

Trade Name	Chemical Name or Active Ingredient	
Hyamine 2389	40% Methyl dodecyl benzyl trimethyl ammonium chloride and 10% methyl xylene bis-trimethyl ammonium chloride	
Hydram <sup>®</sup>	See Molinate	
Hydrothal	Potassium salt of 2,2,4,5 trichlorophenoxy) propionic acid and di-(N,N dimethylalkylamine) salt of 3,6 endoxo-hexahydrophthalic acid	
Ibcol	No information available	
Imidan®	o,o-dimethyl-S-phthalimidomethyl phosphorodithioate	
Inverton <sup>®</sup>	No information available	
lodophor	Iodine formulated with solubilizing agents	
IPC	Isopropyl-N-phenylcarbamate	
Isobenzan (SD-4402)	1,3,4,5,6,7,8,8-octachloro-3a,4,7,7a-tetrahydro- 4,7-methanophthalon	
Isodrin	1,2,3,4,10,10-hexachloro-1,4,4a,5,8,8a-hexahydro- 1,4-endo,endo-5,8-dimethanonaphthalene	
Isolan	Dimethyl 5-(1-isopropyl-3-methyl-pyrazolyl) carbamate	
Isotex	No information available	
K 6882	No information available	
Karmex®	See Diuron	
Kelthane <sup>®</sup>	See Dicofol	
Kepone®	Decachlorooctahydro-1,3,4-methano-2H- cyclobuta-[cd]-pentalen-2-one	
Korlan <sup>®</sup>	See Ronnel	
Kuramine	Amine formulation of 2-(2,4,5-trichlorophenoxy) propionic acid	
Kurosal	No information available	
Kurosal G	No information available	
Kurosal SL	No information available	
Kuron <sup>®</sup>	Propylene glycol butyl ether esters of Silvex	
Куго-Ео	No information available	
Lethane 384	2-(2-butoxyethoxy) ethyl thiocyanate	
Lexone	$\gamma$ -isomer of benzene hexachloride	
Lignasan	Ethylmercury phosphate	
Lindane (Gammexane <sup>®</sup> )	$\gamma$ -isomer of 1,2,3,4,5,6-hexachlorocyclohexane	
M-502	No information available	
M-1499	No information available	
M-1500	2,2,4,5-trichlorophenoxy propionic acid	
M-1845	No information available	
Malaoxon	O <sub>2</sub> analogue of Malathion	

### Trade Name

Malamar-50 Malathion

Manzate<sup>®</sup> Maneb (Manzate<sup>®</sup>, Dithane<sup>®</sup>) MCP MCPA (Agroxone<sup>®</sup>, Methoxone<sup>®</sup>) MCPB Metasystox<sup>®</sup> Methoxychlor Methoyl Demeton (Metasystox<sup>®</sup>)

Methyl parathion (DAEF<sup>®</sup>, Nitrox<sup>®</sup>, Nitrox 80<sup>®</sup>) Methyl phencapton

Methyl trithion<sup>®</sup>

MGA Evergreen MGA 6103 MGA 6243 Mirex

Molinate (Hydram<sup>®</sup>, Ordram<sup>®</sup>) Monuron (Telvar<sup>®</sup>, CMU) Monuron-TCA (Urox<sup>®</sup>)

MS 222 N 2404

N 2788 N 2790 Nabam (Chem Ban, Dithane D-14<sup>®</sup>, Parzate<sup>®</sup>) Naled (Dibrom<sup>®</sup>) Neburon Neguvon<sup>®</sup> Nemagon<sup>®</sup> Neotran<sup>®</sup> Nigrosine Noxfish<sup>®</sup> N-serve 50% Malathion S-[1,2-bis-(ethoxycarbonyl)ethyl] 0,0-dimethyl phosphorodithioate See Maneb Ethylene-bis-dithiocarbamate manganese See MCPA 4-chloro-2-methyl phenoxy acetic acid 4-chloro-2-methyl phenoxy butyric acid See Methyl Demeton 1,1,1-trichloro-2,2-bis-(p-methoxy-phenyl) ethane Mixture of 0.0-dimethyl-o-2-(ethylthio) ethyl phosphorothioate (A) and o,o-dimethyl S-2(ethylthio) ethyl phosphorothioate (B) o,o-dimethyl o,p-nitrophenyl phosphorothioate o,o-dimethyl S-(2,5-dichlorophenylthio) methyl phosphorodithioate o,o-dimethyl s-(p-chlorophenylthio) methyl phosphorodithioate No information available No information available No information available Dodecachlorooctahydro-1,3,4-methano-2Hcyclobuta-[dc]-pentalene S-ethyl hexahydro-1H-azepine-1-carbothioate 3-(p-chlorophenyl)-1,1-dimethylurea [3(p-chlorophenyl)-1,1-dimethylurea trichloracetate] No information available o-isopropyl-o-(2-chloro-4-nitrophenyl)-ethylphosphonothioate o-ethyl-S-p-tolyl-ethylphosphonodithioate o-ethyl-S-phenyl-ethylphosphonodithioate Disodium ethylene bis-dithiocarbamate 1,2-dibromo-2,2-dichloroethyl dimethyl phosphate 3-(3,4-dichlorophenyl)-1-methyl-1-n-butylurea See Trichlorofon 1,2-dibromo-3-chloropropane bis(p-chlorophenoxy) methane Aniline black Rotenone 2-chloro-6-(trichloromethyl) pyridine

Trade Name	Chemical Name or Active Ingredient
Nytron®	25% or more ammonium content
Omazine <sup>®</sup>	Cupric dihydrozinium sulfate
Octachlor	See Chlordane
Octalene	See Aldrin
OMPA	See Schradan
OMS-3	No information available
OMS-44	o-3,5-dichloro-4-methylthiophenyl) o,o-dimethyl phosphorothioate
OMS-115	No information available
OMS-144	No information available
OMS-315	S-p-chlorophenyl 0,0-dimethyl phosphorodithioate
OMS-437	Toluene-α,α-dithiol bis-(0,0-dimethyl phos- phorodithioate)
OMS-595	2-chloro-1-(2,4,5-trichlorophenyl) vinyl dimethyl phosphate
OMS-648	o,o-diethyl-(5-chlorobenzisoxazolyl-3) phos- phorothioate
OMS-658	o-(4-bromo-2,5-dichlorophenyl) o,o-dimethyl phosphorothioate
OMS-659	o-(4-bromo-2,5-dichlorophenyl) o,o-diethyl phosphorothioate
OMS-711	2-chloro-1-(2,5-dichlorophenyl) vinyl dimethyl phosphate
OMS-712	2-chloro-1-(2,4-dichlorophenyl) vinyl dimethyl phosphate
OMS-754	S-(o-chlorophenyl) o,o-dimethyl phosphorodithioate
OMS-868	No information available
Ordram®	See Molinate
Ortho 5305	3-sec-butylphenyl-N-methyl carbamate
Ortho 5353	3-sec-amylphenyl-N-methyl carbama
Ortho 5655	3-sec-butyl 6-chlorophenyl N-methyl carbamate
Ortho MH30	58% diethanolamine salt of 1,2-dihydro-pyridazine- 3,6-dione and 30% maleic hydroxide
Orthocide	See Captan
Ovex (Ovochlor, Ovotran <sup>®</sup> , Estonmite <sup>®</sup> , DOW K-6451 <sup>®</sup> )	p-chlorophenyl, p-chlorobenzene sulfonate
Oxydemetonmethyl (Meta-Systox®)	No information available
Paramar-50	50% Parathion
Para-Oxon (Mintacol®)	o,o-diethyl-o,p-nitrophenyl phosphate
Paraquat	1,1'-dimethyl-4,4'-dipyridylium cation
Parathion (Folidol <sup>®</sup> , Thiophos <sup>®</sup> , Niran <sup>®</sup> , Alkron <sup>®</sup> , Phodiatox <sup>®</sup> )	o,o-diethyl-o,p-nitrophenyl phosphorothioate

Parzate® Pebulate (Tillam®) Perthane® Phencapton (Phenkaptone<sup>®</sup>) Phorate (Thimet<sup>®</sup>) Phosdrin® Phosphamidon (Dimecron<sup>®</sup>) Phygon® Picloram (Tordon<sup>®</sup>) P.M.A. (PMAC, PMAS) Polyclens Polysan Prometone Prometryne Pro-noxfish Propanil (Rogue<sup>®</sup>, Stam F-34<sup>®</sup>) Pyramat Pyrethrin Rivanol Roccal® Rogue® Ronnel (Korlan<sup>®</sup>, Trolene<sup>®</sup>, Viozene<sup>®</sup>, Dow ET-57<sup>®</sup>, Dow ET-14<sup>®</sup>) Rotenone **R**uelene® Ryania (Ryanodine) Sarin Schradan (OMPA, Pestox III<sup>®</sup>, Pestox 3<sup>®</sup>) SD 4402 SD 7727 SD 7772 SD 8211 SD 8447 (OMS-595)

SD 8530

See Nabam S-propylbutylethylthiocarbamate 1,1-dichloro-2,2-bis-(p-ethylphenyl) ethane o,o-diethyl-S-(2,5-dichlorophenylthiomethyl) phosphorodithioate o,o-diethyl-S-(ethylthio) methyl phosphorodithioate 2-carbomethoxy-1-propen-2yl dimethyl phosphate 1-chloro-diethyl-carbamoyl-1-1-propen-2yl dimethyl phosphate See Dichlone 4-amino-3,5,6-trichloro-picolinic acid Phenylmercuric acetate No information available No information available 2-methoxy-4,6-bis(isopropylamino)-2-triazine 2-methylmercapto-4,6-bis(isopropylamino)-3-triazine Rotenone 3',4'-dichloropropion-anilide 2-n-propyl-4-pyridinyl-(6)-dimethyl-carbamate See Barthrin 6,9-diamino-2-ethoxyacridine Alkyl dimethyl benzyl ammonium chloride See Propanil Dimethyl 2,4,5-trichlorophenyl phosphorothioate Decrin 4-tert-butyl-2-chlorophenyl methyl methylphosphoromidite Ground stemwood of Ryania speciosa Isoproporymethyl phosphoryl fluoride Octamethylpyrophosphoramide See Isobenzan 2,4-dichlorophenyl methanesulfonate Phosphoric acid, 2-chloro-1-(2,5-dichlorophenyl) vinyl dimethyl ester Phosphoric acid, 2-chloro-1-(2,5-dichlorophenyl) vinyl dimethyl ester 2-chloro-1-(2,4,5-trichlorophenyl) vinyl dimethyl phosphate Carbamic acid, methyl-3,4,5-trimethyl phenyl ester

Trade Name	Chemical Name or Active Ingredient	
SD 8803	Phosphorothioic acid, o-[2-chloro-1-(2,4-dichloro- phenyl) vinyl]-o,o-diethyl ester	
SD 9129	Dimethyl phosphate of 3-hydroxy-N-methyl-cis- crotonimide	
Separan	No information available	
Sernyl	1-(1-phenylcyclohexyl) piperidine hydrochloride	
Servin	No information available	
Sevin®	See Carbaryl	
Shadran	Octamethylpyrophosphoramide	
Shell 4072	No information available	
Shell SD-7438	Toluene-a, a-dithiol bis-(0,0-dimethyl phos- phorodithioate)	
Shell SD-7961	No information available	
Shell SD-8447	2-chloro-1-(2,4,5-trichlorophenyl) vinyl dimethyl phosphate	
Shell SD-8448	No information available	
Shell SD-9129	Dimethyl phosphate of 3-hydroxy-N-methyl-cis- crotonimide	
Silvex	2-(2,4,5-trichlorophenoxy) propionic acid	
Simazine (Gesatop <sup>®</sup> )	2-chloro-4,6-bis-(ethylamino)-s-triazine	
Sinox General	50% dinitro-o-secondary butyl and 10% dinitro-o- secondary amyl butyl phenol	
Slickgone 1	No information available	
Slickgone 2	No information available	
Slix	No information available	
Sovicide	No information available	
Stam F-34®	See Propanil	
Stauffer N-2790	No information available	
Stauffer R-1910	Ethyl-N,N-diisobutyl thiocarbamate	
Stauffer R-4461	No information available	
Stauffer R-5092	No information available	
Steramine	p-diisobutyl phenoxy ethoxy ethyl dimethyl benzyl ammonium chloride monohydrate	
Strobane <sup>®</sup> (Strobane AC-14 <sup>®</sup> )	Terpene polychlorinates	
Styrene	Phenyl ethylene	
Sulfotepp (Dithione <sup>®</sup> , Bladafume <sup>®</sup> )	0,0,0,0-tetraethyl dithipyrophosphate	
Swep	Methyl 3,4-dichlorocarbanilate	
Systox®	See Demeton	
ТВА	2,3,6-trichlorobenzoic acid	
TCA	Trichloroacetic acid	
TD 47	Di-n,n-dimethylcocoamine salt of 3,6 endoxohexa- hydrophthalic acid	

Trade Name	Chemical Name or Active Ingredient
TD 72	No information available
TD 440	No information available
TD 497	Amine salt of 3,6-endoxohexahydrophthalic acid
TDE (DDD, Rhothane <sup>®</sup> )	2,2-bis-(p-chlorophenyl)-1,1-dichloroethane
Telvar®	See Monuron
Tenoran	See Chloroxuron
Trithion	See Carbophenothion
Telodrin	82% isobornyl thiocyanoacetate
TEPP (Bladan <sup>®</sup> , Tetron <sup>®</sup> , HETP, TEP)	Tetraethyl pyrophosphate
TFM	3-trifluoromethyl-4-nitrophenol
Thanite®	Isobornyl thiocyanoacetate
Tillam®	See Pebulate
Thimet <sup>®</sup>	See Phorate
Thiodan <sup>®</sup>	See Endosulfan
Thionazin (Zinophos)	0,0-diethyl-0-2-pyrazinyl phosphorothioate
Thiram (Nomersan <sup>®</sup> , Pomasol <sup>®</sup> )	Tetramethylthiuram disulfide
Tiguvon	o,o-dimethyl-o-[4-(methylthio)-m-tolyl] phos- phorothioate
Tordon <sup>®</sup>	See Picloram
Tordon 101 <sup>®</sup>	39.6% triisopropanol-amine salt of 2,4-D, and 10.2% picloram triisopropylamine salt
Toxaphene (Phenocide <sup>®</sup> , Phenatox <sup>®</sup> )	Chlorinated camphene
2,4,5TP	No information available
Treflan®	See Trifluralin
Trefmid	No information available
Trichlorofon (Dipterex <sup>®</sup> , Dylox <sup>®</sup> , Neguvon <sup>®</sup> , Tugon <sup>®</sup> )	o,o-dimethyl-(1-hydroxy-2,2,2-trichloroethyl) phosphate
Tricon	No information available
Trifluralin (Treflan®)	a,a,a-trifluoro-2,6-dinitro-N,N-dipropyl-p-toluidine
Trithion <sup>®</sup>	See Carbophenothion
Trolene®	See Ronnel
TV-1096	Lg-threo-2-(5-nitro-2-furyl)-5-(p-nitrophenyl)- 2-oxazoline-4-methanol
UC 8305	p-chloro-2,4-dioxa-5-methyl-p-thiono-3-phosphabi- cyclo-(4.4.0)-decane
UC 10854	m-isopropylphenyl N-methycarbamate
UC 19786	2-sec butyl-4,6-dinitrophenyl isopropyl-carbonate
UC 20047	3-chloro-6-cyano-2-norbornanone o(methyl- carbamoyl) oxime
UC 21149	2-methyl-2-(methylthio) propionaldehyde o-methyl-carbamoyl) oxime

Trade Name	Chemical Name or Active Ingredient
UC 21427	No information available
Urox®	See Monuron-TCA
Vancide 51Z	Mixture of zinc dimethyldithiocarbamate and zinc 2-mercaptobenzothiazole
Vapona®	See DDVP
Velsicol	See Dicamba
Velsicol 1068	See Chlordane
Veon-100	Dimethylamine salt of 2-4 dichlorophenoxyacetic acid-dimethylamine salt-2,4,5-trichlorophenoxy- acetic acid
Vernolate (Vernam <sup>®</sup> )	S-propyl dipropylthiocarbamate
Vernam <sup>®</sup>	See Vernolate
Versene	Sodium acetate
Versenol	A series of chelating agents
Vis-ko Stop-Mold "B"	54% sodium-o-phenylphenate
Weedar MCP	No information available
Weedex	41% sodium metarsenite
Weed Rap	20(2-ethyl hexyl ester of 2,4-dichlorophenoxy- acetic acid)
Zectran®	4-dimethylamino-3,5-xylyl n-methyl-carbamate
Zerlate®	See Ziram
Zinophos	See Thionazin
Ziram (Zerlate <sup>®</sup> , Milbam <sup>®</sup> , Fuklasin <sup>®</sup> )	Zinc dimethyldithiocarbamate
Zytron <sup>®</sup>	o-(2,4-dichlorophenyl)o-methyl isopropylphos- phoramidothioate

## ACKNOWLEDGMENTS

Manufacturing Chemists Association (MCA) recognized for some years the urgent need to summarize information regarding the effect of chemicals on aquatic life as a step toward improved water usage. To this end MCA engaged Battelle to examine the scientific literature and compile pertinent data. Guidance in this program was provided by chemical industry specialists of the MCA Water Resources Committee. Financed by MCA, the planning and execution of the compilation effort and preparation of this report were accomplished by staff members of Battelle's Columbus Laboratories. The authors of this report are H. T. Kemp, J. P. Abrams, and R. C. Overbeck. The Environmental Protection Agency, in supporting publication of this document, is fulfilling its role of making information on water use problems generally available to the scientific community.

1 Accession Number	2 Subject Field & Group 05A and 05C	SELECTED WATER RESOURCE	
A Title	oratories, Columbus, Ohio iteria Data Book - Vo	1. 3	
Effects of Chemicals 10 Author(s) H. T. Kemp, J. P. Abrams, and R. C. Overbeck	16 Project 18 21 Note Co	t Designation 3050GWV5/71 opies available only from GPO. By Battelle's ( aboratories for Manufacturing Chemist Associ	
Data Book, Vol. 3, 528	pp., May, 1971.	ffect of Chemicals on Aquatic Life", Water Q	uality Criteria
23 Descriptors (Starred First) * Toxicity * Bioassay * Industrial wastes * Pesticides * Aquatic organisms * Aquatic animals	* Pest control Pesticide toxicity Bioindicators Agricultural chemicals Fish Chemicals	Chemical wastes Biochemical oxygen demand Fresh water Sea water Bacteria Algae	Aquatic fungi Invertebrates Aquatic insects Oysters Shrimp
25 Identifiers (Starred First)			

# 27 Abstract

Original data from more than 500 technical publications concerning the specific effects of chemicals on individual species of aquatic biota were collected and summarized in uniform format. Alphabetical assembly of the data by chemical allows rapid access to considerable detailed information. A Species Index facilitates search for information on the toxicity of chemicals to individual aquatic species.

The details of major procedures in laboratory bioassay and field assessment of chemical toxicity in water are discussed. Freshwater and marine procedures are included. A total of approximately 1000 references were utilized in preparing this report.

Recommendations include:

- (1) Establishment of an information-analysis center on chemical water pollution based to some extent on the report prepared.
- (2) Preparation of a listing of chemical constituents of effluents and continued up-dating of this list.

(Kemp – Battelle)

Abstractor H. T. Kemp	Institution Battelle's Columbus Laboratories
WR:102 (REV.JULY 1969) WRSIC	SEND, WITH COPY OF DOCUMENT, TO: WATER RESOURCES SCIENTIFIC INFORMATION CENTER U.S. DEPARTMENT OF THE INTERIOR WASHINGTON. D. C. 20240