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An Analysis of the Dynamics of DDT in Marine Sediments



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AN ANALYSIS OF THE DYNAMICS OF DDT IN MARINE SEDIMENTS

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ABSTRACT

The concentration of the three chlorinated hydrocarbons, DDT, DDD, and DDE, were measured in sediments at 57 stations in Monterey Bay on the Central California coast during 1970 and 1971. Mean concentration in parts per billion was DDT 3.1, DDD 2.3, and DDE 5.4. Maximum concentrations were DDT 19.3, DDD 8.7, DDE, 20.5 parts per billion. The distribution of the three compounds within South Monterey Bay was charted. During 1973 nineteen of the original stations, representing locations that were low, intermediate, and high concentrations in the original survey, were resampled. The mean concentration approximately three years later were DDT 15.5, DDD 2.3, and DDE 5.4 parts per billion with maximum levels of DDT 83.1, DDD 11.4, and DDE 17.5 parts per billion. A chart of the concentrations in South Monterey Bay revealed essentially the same distribution of chlorinated hydrocarbons.

Two approaches to the estimation of annual system rates for input, I, output, O, decay, D, and internal translocation, T_I and T_O , expressed as decimal fractions of the existing concentration were developed, and Fortran programs that permit rapid estimations were written. The mean annual system rates obtained were for DDT, $I+1.30$, $O-.059$, $D-.036$, T_I and $T_O \pm .80$ with a residence time of 11 years and life time of 29 years. An I of 1.30 means the amount of input is 130% of the existing concentration per year. The mean annual rates obtained for DDD were, $I + 0.25$, $O - 0.11$, $D - 0.025$, T_I and $T_O \pm 0.20$ with residence time of 7 years and life time of 44 years. The rates for DDE were $I + 0.28$, $O - 0.10$, $D - 0.027$, T_O and $T_I \pm 0.22$ with residence time of 8 years and life time of 39 years. The approaches to these estimates are dependent upon variability in net rates of change at the various stations and an approach to evaluation of the standard deviation of the estimated rates relative to distributions of net rates with minimal variance is presented.

Laboratory assays were developed to determine the relative rate of decomposition in sediment placed under conditions selective for various physiologically different kinds of microorganisms. ^{14}C ring labelled substrates were used in all assays. Decay of the three chlorinated hydrocarbons under aerobic conditions without additional nutrients was greater than decay under anaerobic conditions. The addition of accessory energy and carbon sources such as sodium acetate did not increase the rate of decay under anaerobic conditions. There was some decay under anaerobic conditions suggesting mechanisms of ring cleavage not involving incorporation or oxygen prior to ring split. Nitrate as an accessory electron acceptor increased the rate of decomposition under anaerobic conditions. Degradation products formed from the parent compounds included water soluble intermediates as well as carbon dioxide.

The Q_{10} for the decay process as determined by laboratory assays incubated at 10° and $20^\circ C$. is 2.5.

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SECTION I

CONCLUSIONS

Chlorinated hydrocarbons associated with sediment particles tend to concentrate in sedimentation basins which may be at some distance from the input source.

Although the use of chlorinated hydrocarbon pesticides has declined sharply the levels of three materials has continued to increase in marine sediments. The principal source of this additional pollutant load in this instance appears to be more related to translocation of these materials absorbed to sediments of adjacent land drainage systems.

The dynamics of chlorinated hydrocarbons in the coastal marine environment, although complex, are susceptible to study. Approaches to the estimation of rates of input, decay, and translocation can be developed and assessed by continued analysis of environmental samples.

The measurement of decay rate by laboratory assay appears to have its greatest utility in the determination of the effect of environmental conditions on the process of decay. Duplication of conditions existing *in situ* in the laboratory can only be approximated and then only for a limited time. The laboratory work, short term in its execution, serves only as a guide to what is happening in the environment.

SECTION II

RECOMMENDATIONS

The complexities of the dynamics of coastal pollution by chlorinated hydrocarbons necessitates an initial survey of the concentration of these environmental contaminants at a large number of stations. Once basins of accumulation are established and principal translocation paths established a much smaller number of stations require surveillance at later points in time. It doesn't appear to be essential to monitor exactly the same stations in any surveillance program as long as the set of surveillance stations includes established basins and positions along translocation pathways.

It is recommended that initial intensive surveys be carried out in the coastal marine environment adjacent to major agricultural and industrial areas which are known to produce or utilize poorly degraded environmental contaminants such as the chlorinated hydrocarbons.

Monterey Bay is a very useful model coastal marine environment for the establishment and testing of approaches to system rate estimation. Continued surveillance of this area is recommended.

It is also recommended that work be done on extending the approach to estimation of system rates explored with respect to sediments to other environmental systems including populations of organisms. It would appear desirable to concentrate initially upon abundant and useful indicator organisms rather than commercially desirable or affected species.

Finally, it is recommended that additional effort be expended on the study of laboratory assays of decay not only as approximations of the environment but as useful preparations for elucidating the conditions inhibitory and stimulatory to the decay process.

SECTION III

INTRODUCTION

Although the accumulation of chlorinated hydrocarbons in the marine ecosystem has been a matter of concern for some time, methods for assessing the rates of accumulation, decay, and translocation have been lacking. The problem is not unique to the marine environment, and methods for assessment of the dynamics of chemical pollutants in general are needed for meaningful analysis of the residue measurements tabulated in most investigations. Without an assessment of rates such tabulations generally permit only the detection of some general trend of increase or decrease in concentration during the period of study. In many cases, however, the amount of variability is so great that the number of samples required to show such general trends is prohibitive. Yet we have both the data available and a need to use these data for meaningful assessment. In addition, before any feasible monitoring activity geared to control and regulatory strategies are designed and implemented, a means of assessing any new tabulations is required as a determinant in the design of such activities. Whatever systems of assessment may be developed in the future it cannot be expected that they will overcome the variability that plagues environmental sampling. Rather, such systems should be expected to provide an estimate of this variability and a confidence interval for any derived parameter of environmental change.

Several models stressing one or another aspect of the dynamics of pesticides in the environment have been presented (Hamaker 1966, Robinson 1967, Woodwell 1967, Harrison et al. 1970, and Eberhardt et al. 1971), but there still appears to be a need for a general approach that provides a means of estimating rates of input, decay, and translocation from some minimal number of analyses. The study presented here is an attempt to fill this need.

The data used here for these estimations consists of analyses of marine sediment samples for 1,1,2-trichloro-2,2-bis (p-chlorophenyl) ethane, DDT; 1,1-dichloro-2,2-bis (p-chlorophenyl) ethane, DDD; and 1,1-dichloro-2,2-bis (p-chlorophenyl) ethylene, DDE. The rates of decay at a sampling site and translocation away from a sampling site are difficult to separate through the approach to estimation presented. Laboratory measurements of the rate of ^{14}C ring labelled DDT in marine sediments held under a variety of conditions are also presented. These measurements reflect decay to the point of $^{14}\text{CO}_2$ release rather than conversion to any one of a variety of other metabolites including DDD and DDE, but are useful in assessing the method of estimation based upon environmental samples alone.

The analysis of DDT residue levels in marine sediments reported herein is only a part of a larger study correlating the levels of pollutants with density and composition of benthic populations. Other results of this study will be reported elsewhere.

THE STUDY AREA

This study was carried out in Monterey Bay located in the central coastal region of California. Figure 1 shows the study area and the location of the forty-nine Stations from which sediment samples were obtained. The figure also shows several geographical features pertinent to this investigation. The bottom of Monterey Bay is divided by a major submarine canyon over 3800 meters in depth at its deepest point. The sampling effort was concentrated in the southern portion of the bay with no sampling beyond the 200 fathom, 365 meter, line. Residue levels of DDT, DDD, and DDE were first measured in samples from this southern portion of the bay during 1970 and nineteen of these stations were resampled in 1973. A small number of stations were sampled in the northern part of the bay during 1971.

Monterey Bay is the recipient of drainage from a major agricultural area, the Salinas Valley, where DDT was used in large amounts for a period of twenty years. Usage of this pesticide and DDD has decreased sharply since 1969. A tabulation of use was started in 1970 when 33,931 pounds was applied to 19,387 acres in Monterey County. This input level was further reduced in 1971 to 4,697 pounds, and in 1972 to 10 pounds on 20 acres (Calif. Dept. of Agriculture 1970, 1971, 1972). Final tabulations for 1973 will probably show levels of input similar to those of 1972. Although the use of DDT in the area adjacent to Monterey Bay has declined sharply since 1970, the level of DDT in marine sediments appears to be increasing as more of this pesticide finds its way to the sea via the drainage system of the neighboring agricultural area. The decrease in usage on adjacent land and apparent increase in concentration in the marine sediments of the area suggests that continued study of the Monterey area is of particular interest in determining the time lag between terrestrial input and marine accumulation of persistent chemical pollutants.

Although in the past, when DDT was being regularly applied on the adjacent lands, the atmosphere was an important source of input to the bay; at the present time the major source of input appears to be the Salinas River which drains the inland agricultural areas. This river flows directly into the bay only intermittently. Most of the time the mouth of the river is blocked by a bar of sand that is removed only at times of heavy rainfall to prevent flooding. During this investigation this event occurred Jan. 13, 1970, Nov. 30, 1970, Dec. 29, 1971, Nov. 16, 1972, Nov. 17, 1972, and Nov. 20, 1973. Input directly by the river has, therefore, not been continuous.

Analyses of the sediment samples from the river bed along its course in 1972 (State of California, 1974) showed considerable variation in the relative abundance and concentration of the three compounds. Table 1 gives the results of these analyses and the approximate location of the samples relative to the mouth of the river.

During the periods when the mouth of the river is blocked, there is a sluggish flow north to Elkhorn Slough which served as the mouth of the river until 1908. This flow is joined by drainage from Trembladero Slough which receives water and sediments from the Reclamation Canal that flows through the City of Salinas to the east and beyond the right-hand margin of the figures. The Reclamation Canal receives effluents from food processing plants and other industries, and analyses of its sediment in 1972 (State of Calif., 1974) revealed the levels also listed in Table 1.

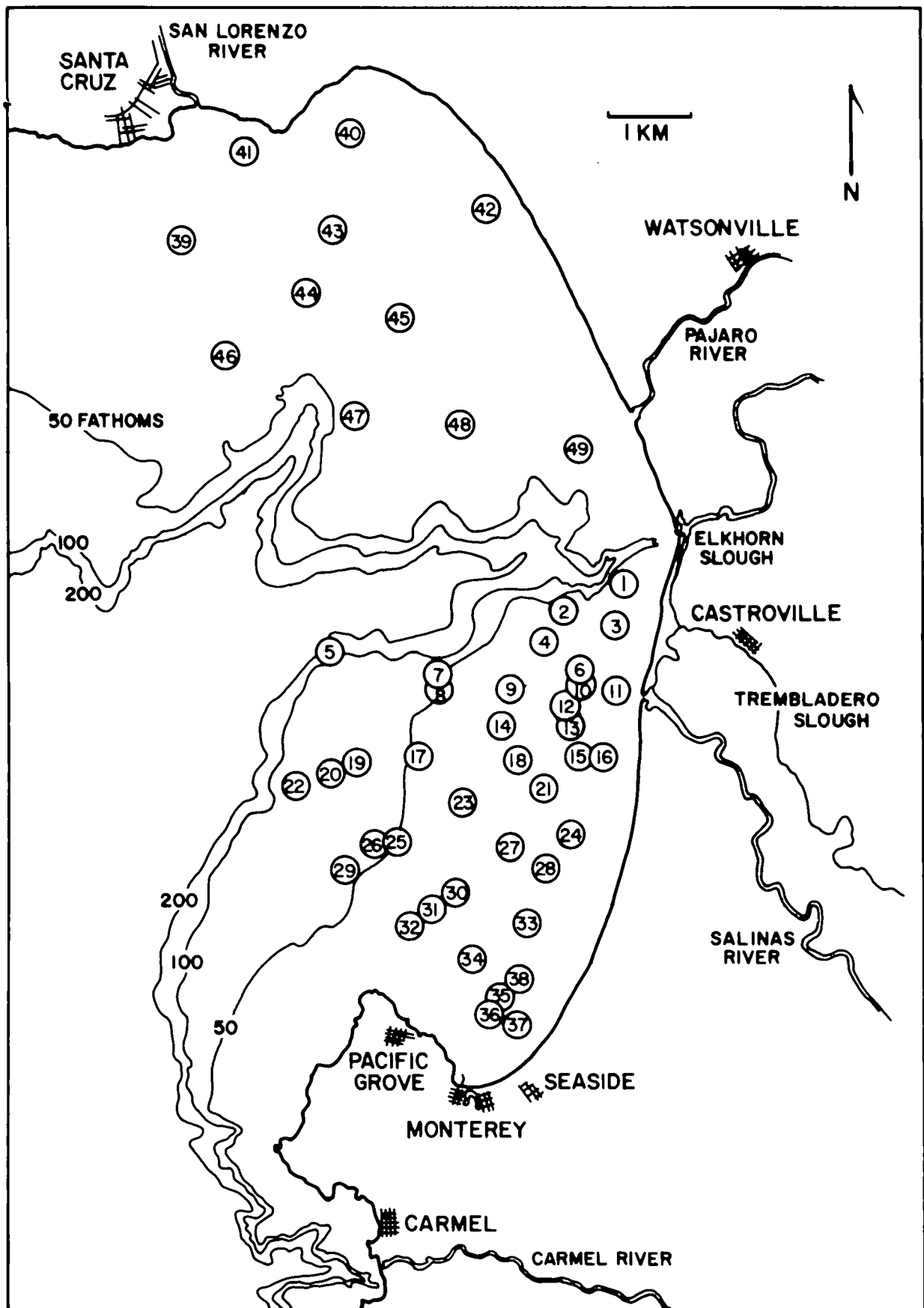


Figure 1. The study area, Monterey Bay. Sampling stations are indicated by number.

Table 1. CONCENTRATION OF DDT, DDD, AND DDE IN SEDIMENTS OF THE MONTEREY AREA LAND DRAINAGE SYSTEM IN 1972 (STATE OF CALIF., 1974)

Salinas River distance from mouth (kilometers)	(ppb)		
	DDT	DDD	DDE
42	1.0	1.3	
25	120.		20.
8	150.	1000.	360.
	16.	620.	
3	0.12	30.	
Reclamation Canal distance from mouth of Elkhorn Slough (kilometers)			
20	7,000.	45,000.	10,000.
	21,000.	150,000.	

RATIONALE OF DESCRIBED WORK

Selection of Study Site and Source of Marine Sediments for Decay Assays—For the estimation of rates governing the dynamics of a chlorinated hydrocarbon pollutant in marine sediments an area with the following characteristics appeared most desirable. (1) The marine area should be adjacent to a land area for which there exists an accounting of input to the environment through normal use. The use of DDT and DDD within the State of California has been subject to such accounting on a square mile section basis since 1970 (Calif. Dept. of Agriculture 1970). Such accounting is available only for normal agricultural and related uses. Therefore, areas which receive or have received less well determined inputs from chlorinated hydrocarbon manufacture, such as the ocean adjacent to Los Angeles, are less desirable for this type of study. (2) In order to assess translocation within the study area it would appear desirable to select a marine area with a limited number of point sources of input rather than one subject to diffuse input by way of the atmosphere. (3) The area should be one open to general oceanic influence rather than a closed system so that translocation of the pollutant out of the system by dilution or dissemination can be assessed. (4) As a source of materials for laboratory assays of decay the area should be one which has had a long exposure to the pollutant, thus insuring the establishment of microbial systems with the capacity for decomposition of the pollutant. (5) The area should be known to be contaminated with the pollutant. (6) The area should be accessible to sampling and close to the required analytical capability.

Monterey Bay, and in particular the southern portion of Monterey Bay, has these characteristics and was selected as the study site and source of materials for the development of laboratory assays for the rate of decay of DDT, DDD, and DDE.

Survey of Residue Levels in Monterey Bay Sediments—In order to assess the variability in concentration and distribution of the three compounds in the sediments of Monterey Bay thirty-seven sample sites were selected for analysis in the southern portion of the bay which receives water and sediments from the agricultural area of Monterey County by way of the Salinas River. An additional eleven sample sites in the northern portion of the bay were selected in order to assess any augmenting effect of additional river input sources such as the San Lorenzo and Pajaro Rivers that drain areas of Santa Cruz and San Benito Counties lying adjacent to Monterey County and Monterey Bay.

Determination of the Amount of Change in Residue Levels with Time—In order to assess the magnitude of change in the concentration of DDT and related compounds a subset of the original survey sampling stations was resampled and analyzed after approximately three years. Nineteen of the original sample stations were selected as this subset. The selection was made on a basis of accessibility and representations of stations showing a broad range of residue concentrations as determined in the original survey.

Determination of the Variance of Sampling—One additional sample station, number 38, which had never before been sampled was added to the resampled subset and sampled three times on the same day. Three aliquots from each of these samples were analyzed for the three compounds to provide an estimate of the variability of sampling.

Approaches to the estimation of rates and Dynamics of the compounds in Sediments— Using the tabulated data obtained from the sampling programs various approaches to the estimation of the rates of input, translocation, and decay were developed for the system of sample sites. Considerable attention was directed to estimation of variance of these derived rates.

Development of Laboratory Assay Methods for the Determination of Decay Rate— Measurement of decay rate based on changes in residue level observed by repeated sampling from the environment are subject to error due to translocation to or away from the sample site. Therefore, a means of estimating decay rate in a closed system not susceptible to such error would be desirable. A variety of preparations using ^{14}C ring labelled compounds were established for such estimations.

Effect of Environmental Variables on Decay Rate—Any closed system preparation is by its very nature selective for one or another metabolic type of microorganism. The initial conditions and conditions which subsequently develop may have a marked effect upon the observed rate of decomposition through the election of particular microbial populations. Therefore, it was necessary to study the process of decay as influenced by a number of environmental variables chosen to encourage one or another of the major metabolic types of microorganisms.

SECTION IV

METHODS

ANALYSIS OF SEDIMENT SAMPLES

Samples of sediment were collected by Shipek grab or shallow dredge. Between 50 and 70 grams of wet sediment were placed in a 250 ml bottle and mixed with 30-50 grams of granular anhydrous sodium sulfate. The sediment was extracted with 50 ml of acetone:hexane, 1:1, by shaking for four hours. The acetone, hexane was decanted and filtered through a fritted glass filter or silicon-treated phase separation paper into a separatory funnel. Three additional 50 ml portions of hexane were used to wash the sediment and added to the original extractant.

The extract was washed with three 200 ml portions of water followed by dehydration of the extract by passage through a 2x5 cm column of anhydrous sodium sulfate and concentration in a Kuderna-Danish concentrator to less than 10 ml. The extract was then cleaned by shaking first with 1 ml of concentrated sulfuric acid and finally with approximately 0.1 ml of mercury. The analysis was performed in a Beckman GC-4 Gas Chromatograph with electron capture detector, using a mixed bed column of Chromosorb W, 80-100 mesh, DMCS treated, and acid washed, containing 5% DC-200 and 5% QF1.

Although the efficiency of extraction is difficult to assess, the effect of concentration and clean-up procedures can be measured by the use of ^{14}C labelled materials added just prior to extraction with acetone, hexane. Recovery was 73.9% for DDT, 94.4% for DDD, and 84.8% for DDE, and these figures were used to correct the results of analyses.

LABORATORY DECAY ASSAYS

A variety of preparations have been investigated for their applicability to decay assay preparations. These preparations have included sealed stationary aliquots of sediment and ^{14}C labelled substrate as well as ones in which the sediment with labelled substrate was subjected to continuous percolation or periodic gas flow. Maintenance of percolating systems for the length of time required to measure the very slow rates of decay is not feasible, and it is difficult to maintain a large number of preparations under conditions whereby they may be subjected to periodic gas flow and trapping of metabolic CO_2 . Therefore, sealed stationary preparations have proved to be the only feasible type of preparation so far developed. The most convenient container for such preparations has been 125 ml Hypovials, Pierce, Rockford, Illinois, No. 12995, fitted with Teflon liners. The preparation of decay assays is as follows. Sediment is collected as for samples for residue analysis, packed in ice, and brought to the laboratory within a few hours. The sediment is rinsed through screen with 16 mesh to the inch to remove macroscopic in-fauna and refrigerated. Aliquots of the slurried sediment are removed for dry weight

determination. A volume of the slurried sediment equivalent to 24 grams dry weight is delivered to a sterile Hypovial and seawater, with or without additional nutrients, is added to give a volume of 98 ml total. One ml each of ^{12}C and ^{14}C substrate adsorbed to sterile sediment is added giving a final volume of 100 ml. The preparation may be gassed with nitrogen to produce an anaerobic environment prior to sealing. All incubators are in the dark for periods of generally twelve weeks. All preparations are set up in quintuplicate. A typical protocol is presented in Table 2.

^{12}C Substrate Preparation—2.4 grams of either, 1,1-bis-(p-chlorophenyl)-2,2,2-trichloro ethane, p-p'DDT 99+% No. 10, 002-1; 2,2-bis-(p-chlorophenyl)-1,1-dichloro ethylene, No. 12, 289-7 (B 3964); or 2,2-bis-(p-chlorophenyl)-1,1-dichloroethane, puriss B 3959 Aldrich Chemical Co. Inc., Milwaukee, Wisconsin, were dissolved in 10 ml of acetone. To 10 grams of dried sterile sediment 1 ml of acetone solution was added and the sediment wet with an additional 3 ml of acetone. The acetone was evaporated off at room temperature and 96 ml of distilled water added to slurry the sediment and its adsorbed substrate. One ml contains 2.4×10^3 ug of substrate on 0.1 gram of sediment per ml. Similar preparations were made giving 2.4×10^2 ug and 21.6 ug of substrate on 0.1 gram of sediment per ml.

^{14}C -DDT Substrate Preparation—Uniformly ring labelled DDT, Amersham/Searle Corp., 63.9 u Ci/mg in benzene was used for preparation of the substrate. The original 250 u Ci preparation was diluted with acetone and 240 ug in 4 ml was added to 10 grams of dried sterile sediment. The acetone was removed by evaporation at room temperature and 96 ml of distilled water added to give 2.4 ug ^{14}C -DDT and 0.1 gram of sediment per ml. A similar preparation was made giving 0.24 ug ^{14}C -DDT and 0.1 gram of sediment per ml.

^{14}C -DDD Substrate Preparation— ^{14}C -DDT was converted to ^{14}C -DDD by the method of Murphy (1970) and purity of the product confirmed by gas chromatography. The resulting material was used to prepare substrate as described above for ^{14}C -DDT.

^{14}C -DDE Substrate Preparation— ^{14}C -DDT was converted to ^{14}C -DDE by the method of Gunther and Blinn (1950) and purity of the product confirmed by gas chromatography. The resulting material was used to prepare substrate as described above for ^{14}C -DDT.

Analysis of Decay Assays—After incubation for generally 12 weeks $^{14}\text{CO}_2$ was trapped by the addition of 1.5 ml of 5 N NaOH to the Hypovial. The base was introduced by syringe and the ampoule resealed with tape. Syringe delivered 5 ml aliquots of the basic slurried sediment were transferred to 25 ml Erlenmeyer flasks containing magnetic stirring bars. The flasks were stoppered with Top stoppers, K-882310, fitted with plastic center wells, K-882320, both from Kontes Glass Co., Vineland, N.J. The center wells contained an accordion pleated Whatman No. 1 filter paper wick, 2.5x5 cm. β -phenyl-ethylamine, 0.15 ml, was delivered to the well and wick by syringe through the stopper. While the sediment in the flask was gently stirred on a magnetic stirrer 0.25 ml of 5 N H_2SO_4 was added to the sediment. The flasks were then held for 24 hours at room temperature after which time the wicks were removed to scintillation vials to which was added 15 ml of Toluene-omnifluor. Appropriate preparations for background

Table 2. TYPICAL DECAY ASSAY PROTOCOL.

Hypovial No.	Slurried (grams)	Sediment (ml)	Seawater plus nutrients (ml)	12C Substrate		14C Substrate		Total Substrate (ppm)	Total volume (ml)
				(ug)	(ml)	(ug)	(ml)		
1-5	24	59	39	2400	1	2.4	1	100	100
6-10	24	59	39	240	1	2.4	1	10	100
11-15	24	59	39	21.6	1	2.4	1	1	100
16-20	24	59	39	0	0	2.4	1	0.1	100
21-25	24	59	39	0	0	0.24	1	0.01	100

measurement were also made. The amount of $^{14}\text{CO}_2$ was determined in a Nuclear Chicago Corp. Unilux II. Diffusion time and trapping volume of β -phenylethylamine were established through tests using a standard preparation of $\text{Na } ^{14}\text{CO}_3$.

DECAY AS AFFECTED BY ENVIRONMENTAL VARIABLES

The effect of temperature was determined by comparing the amount of decomposition at 10° and 20°C , and the effect of oxygen, nitrate, and sulfate as terminal electron acceptors in the presence and absence of cometabolizable sodium acetate and ethanol was determined by appropriate additions to the Hypovials.

SECTION V

RESULTS AND DISCUSSION

SURVEYS OF RESIDUE LEVELS IN MONTEREY BAY SEDIMENTS

The concentration in parts per billion of the three compounds, DDT, DDD, and DDE in sediment samples collected during the three sampling periods are presented in Table 3. Table 4 presents the same set of analyses in terms of the percent of total residues for each of the three compounds.

The variance of sampling at Station 38 can be assessed from the data presented in Table 5. The greatest variation in results can be observed with respect to DDT, the compound also showing the greatest loss during the extraction, concentration, and cleanup procedures as mentioned in the section on methods.

The data obtained in the 1970 and 1971 samplings is presented in Figures 2, 3, and 4, where the distribution of DDT and its two derivatives is displayed in terms of percent of the concentration of total DDT derivatives. Figures 5 and 6 show the distribution in terms of the total concentration of DDT and its two derivatives in parts per billion. Figure 5 shows the distribution in 1970 and 1971, and Figure 6 shows the distribution as indicated by the analyses of the smaller number of samples obtained in 1973.

The small number of sample stations in the northern portion of the bay did not reveal any unusual augmentation in concentrations of the three compounds due to input from the San Lorenzo and Pajaro Rivers although the percent composition of DDT derivatives does indicate differences between the northern and southern portions of the bay.

If particular attention is paid to the southern portion of the bay for which there is the greatest information, the distributions suggest a number of characteristics of the system. After input with sediments from the Salinas River, and perhaps also through Elkhorn Slough, these materials are subjected to considerable translocation due to the currents operating within the south bay. The highest concentration of DDT derivatives is to be found at a considerable distance from the mouth of the river. Close to the mouth of the river, however, the sediments show a high percentage of DDT which is characteristic of some of the sediments within the drainage system. These high DDT percentages are also found at the more distant points where the highest concentrations of derivatives are found as well. Over much of the area in terms of percent, however, DDE represents the major compound.

These plots of distribution reflect input over a considerable period of time. During this time the major routes of input may have changed considerably as has the relative concentrations of the three derivatives in these input sources. Nevertheless, the apparent constancy of location of major basins of deposition is remarkable. Areas with high concentrations in 1970 have become even more heavily contaminated in 1973.

Table 3. CONCENTRATIONS OF DDT, DDD, AND DDE IN MARINE SEDIMENT SAMPLES FROM MONTEREY BAY.

Station	LOCATION		Date	DDT (ppb)	DDD (ppb)	DDE (ppb)	TOTAL
	Latitude	Longitude					
1	36 47.25	121 48.90	8-23-70	8.36	3.67	5.76	17.79
2	36 46.85	121 53.50	11-15-70	1.63	6.76	14.70	23.09
3	36 46.35	121 49.00	2-20-70	5.71	0.71	1.02	7.44
4	36 46.05	121 51.00	11-15-70	4.28	6.61	10.70	21.59
5	36 46.00	121 57.00	5-29-70	2.14	0.93	4.00	7.07
6	36 45.45	121 50.00	11-15-70	2.04	1.17	1.80	5.01
7	36 45.30	121 54.00	5-29-70	0.0	2.50	4.51	7.01
8	36 45.20	121 54.00	5-29-70	2.65	4.26	6.51	13.42
9	36 45.10	121 52.00	5-29-70	4.48	5.14	4.51	14.13
10	36 45.10	121 50.00	5-29-70	6.42	8.67	7.01	22.10
11	36 45.00	121 49.00	2-20-70	3.67	0.40	0.45	4.52
12	36 44.60	121 50.50	2-20-70	0.52	0.18	0.28	0.98
13	36 44.25	121 50.35	11-15-70	0.26	0.19	0.45	0.90
14	36 44.20	121 52.25	8-23-70	5.20	7.50	15.50	28.20
15	36 44.00	121 50.00	5-29-70	0.0	0.19	0.35	0.54
16	36 44.00	121 49.50	2-20-70	0.69	0.14	2.75	3.58
17	36 43.75	121 54.45	11-15-70	1.02	0.38	0.70	2.10
18	36 43.50	121 51.80	2-20-70	1.73	2.64	2.40	6.77
19	36 43.35	121 56.25	8-23-70	1.12	0.25	0.65	2.02
20	36 43.18	121 57.00	2- 8-70	0.0	5.00	20.50	25.50
21	36 43.00	121 51.00	5-29-70	6.12	1.30	6.01	13.43
22	36 42.90	121 58.00	2-20-70	0.0	0.35	1.92	2.27
23	36 42.55	121 53.30	8-23-70	13.20	5.73	13.00	31.93
24	36 42.50	121 50.30	8-23-70	19.30	0.65	2.75	22.70
25	36 41.70	121 55.00	2-20-70	1.22	0.53	2.40	4.15
26	36 41.55	121 55.50	2- 8-70	0.0	2.35	7.01	9.36
27	36 41.50	121 52.00	5-29-70	2.85	2.50	8.01	13.36
28	36 41.00	121 51.00	11-15-70	0.0	1.61	4.26	5.87
29	36 40.90	121 56.40	2-20-70	1.32	1.61	9.02	11.95
30	36 40.50	121 53.50	5-29-70	2.55	1.76	6.76	11.07
31	36 40.08	121 54.05	2- 8-70	0.0	0.82	3.25	4.07
32	36 39.80	121 54.50	5-29-70	2.04	1.91	5.26	9.21
33	36 39.80	121 51.50	2- 9-70	0.0	1.42	8.52	9.94
34	36 39.10	121 53.08	2- 8-70	2.44	0.66	2.40	5.50
35	36 39.10	121 53.08	2- 8-70	8.67	0.66	3.00	12.33
36	36 37.95	121 52.50	2-20-70	2.65	2.79	10.00	15.44
37	36 37.77	121 51.83	2- 8-70	0.49	0.21	0.50	1.20

Table 3. (continued) CONCENTRATIONS OF DDT, DDD, AND DDE IN MARINE SEDIMENT SAMPLES FROM MONTEREY BAY.

Station	LOCATION		Date	DDT (ppb)	DDD (ppb)	DDE (ppb)	TOTAL
	Latitude	Longitude					
39	36 54.80	122 01.00	11-24-71	0.60	1.90	2.00	4.50
40	36 57.10	121 56.20	11-10-71	1.62	8.15	5.54	15.31
41	36 56.70	121 59.20	11-24-71	0.93	2.75	4.48	8.16
42	36 55.50	121 52.60	11-10-71	0.85	1.58	0.66	3.09
43	36 55.10	121 56.70	11-10-71	0.81	3.07	2.59	6.47
44	36 53.60	121 57.50	11-24-71	1.13	2.54	2.47	6.14
45	36 53.00	121 55.00	11-10-71	1.21	2.01	1.88	5.10
46	36 52.30	121 59.80	11-24-71	1.27	3.81	5.06	10.14
47	36 51.00	121 49.80	11-10-71	1.16	1.27	1.13	3.56
48	36 50.80	121 53.60	11-24-71	1.62	5.61	6.72	13.95
49	36 50.20	121 50.20	11-10-71	0.78	1.48	1.29	3.55
1	36 47.25	121 48.90	7- 9-73	1.06	0.53	0.56	2.15
2	36 46.85	121 53.50	7- 9-73	9.50	11.40	17.50	38.40
3	36 46.35	121 49.00	7- 9-73	1.10	0.53	0.63	2.26
4	36 46.05	121 51.00	7- 9-73	3.63	5.43	6.91	15.97
10	36 45.10	121 50.00	7- 2-73	0.92	0.39	0.52	1.83
11	36 45.00	121 49.00	7- 2-73	2.18	0.72	0.83	3.73
14	36 44.20	121 52.25	6-21-73	30.60	6.07	11.20	47.87
16	36 44.00	121 49.50	7- 2-73	0.96	0.06	0.23	1.25
17	36 43.75	121 54.45	8- 9-73	5.41	4.54	17.30	27.25
19	36 43.35	121 56.25	8- 9-73	72.70	3.19	12.00	87.89
20	36 43.18	121 57.00	6-21-73	63.10	0.79	3.48	67.37
22	36 42.90	121 58.00	8- 9-73	0.93	0.90	6.06	7.89
23	36 42.55	121 53.30	6-21-73	29.90	4.32	12.20	46.42
25	36 41.70	121 55.00	7-16-73	1.14	2.74	10.49	14.37
26	36 41.55	121 55.50	7-16-73	0.68	2.20	8.67	11.55
29	36 40.90	121 56.40	7-16-73	0.70	1.11	5.67	7.48
34	36 39.10	121 53.08	8- 9-73	1.18	0.42	2.44	4.04
36	36 37.95	121 52.50	6-21-73	83.10	0.95	3.34	87.39
37	36 37.77	121 51.83	7-16-73	0.54	0.20	0.40	1.14
38	36 38.47	121 51.68	9-21-73	0.62	0.38	2.72	3.72

Table 4. LEVELS OF DDT, DDD, AND DDE AS PERCENT OF TOTAL RESIDUES IN MARINE SEDIMENT SAMPLES FROM MONTEREY BAY.

Station	LOCATION		Date	DDT (%)	DDD (%)	DDE (%)
	Latitude	Longitude				
1	36 47.25	121 48.90	8-23-70	46.99	20.63	32.38
2	36 46.85	121 53.50	11-15-70	7.06	29.28	63.66
3	36 46.35	121 49.00	2-20-70	76.75	9.54	13.71
4	36 46.05	121 51.00	11-15-70	19.82	30.62	49.56
5	36 46.00	121 57.00	5-29-70	30.27	13.15	56.58
6	36 45.45	121 50.00	11-15-70	40.72	23.35	35.93
7	36 45.30	121 54.00	5-29-70	0.0	35.66	64.34
8	36 45.20	121 54.00	5-29-70	19.75	31.74	48.51
9	36 45.10	121 52.00	5-29-70	31.71	36.38	31.92
10	36 45.20	121 50.00	5-29-70	29.05	39.23	31.72
11	36 45.00	121 49.00	2-20-70	81.19	8.85	9.96
12	36 44.60	121 50.50	2-20-70	53.06	18.37	28.57
13	36 44.25	121 50.35	11-15-70	28.89	21.11	50.00
14	36 44.20	121 52.25	8-23-70	18.44	26.60	54.96
15	36 44.00	121 50.00	5-29-70	0.0	35.19	64.81
16	36 44.00	121 49.50	2-20-70	19.27	3.91	76.82
17	36 43.75	121 54.45	11-15-70	48.57	18.10	33.33
18	36 43.50	121 51.80	2-20-70	25.55	39.00	35.45
19	36 43.35	121 56.25	8-23-70	55.45	12.38	32.18
20	36 43.18	121 57.00	2- 8-70	0.0	19.61	80.39
21	36 43.00	121 51.00	5-29-70	45.57	9.68	44.75
22	36 42.90	121 58.00	2-20-70	0.0	15.42	84.58
23	36 42.55	121 53.30	8-23-70	41.34	17.95	40.71
24	36 42.50	121 50.30	8-23-70	85.02	2.86	12.11
25	36 41.70	121 55.00	2-20-70	29.40	12.77	57.83
26	36 41.55	121 55.50	2- 8-70	0.0	25.11	74.89
27	36 41.50	121 52.00	5-29-70	21.33	18.71	59.96
28	36 41.00	121 51.00	11-15-70	0.0	27.43	72.57
29	36 40.90	121 56.40	2-20-70	11.05	13.47	75.48
30	36 40.50	121 53.50	5-29-70	23.04	15.90	61.07
31	36 40.08	121 54.05	2- 8-70	0.0	20.15	79.85
32	36 39.80	121 54.50	5-29-70	22.15	20.74	57.11
33	36 39.80	121 51.50	2- 9-70	0.0	14.29	85.71
34	36 39.10	121 53.08	2- 8-70	44.36	12.00	43.64
35	36 39.10	121 53.08	2- 8-70	70.32	5.35	24.33
36	36 37.95	121 52.50	2-20-70	17.16	18.07	64.77
37	36 37.77	121 51.83	2- 8-70	40.83	17.50	41.67

Table 4. (continued) LEVELS OF DDT, DDD, AND DDE AS PERCENT OF TOTAL RESIDUES IN MARINE SEDIMENT SAMPLES FROM MONTEREY BAY.

Station	LOCATION		Date	DDT (%)	DDD (%)	DDE (%)
	Latitude	Longitude				
39	36 54.80	122 01.00	11-24-71	13.33	42.22	44.44
40	36 57.10	121 56.20	11-10-71	10.58	53.23	36.19
41	36 56.70	121 59.20	11-24-71	11.40	33.70	54.90
42	36 55.50	121 52.60	11-10-71	27.51	51.13	21.36
43	36 55.10	121 56.70	11-10-71	12.52	47.45	40.03
44	36 53.60	121 57.50	11-24-71	18.40	41.37	40.23
45	36 53.00	121 55.00	11-10-71	23.73	39.41	36.86
46	36 52.30	121 59.80	11-24-71	12.52	37.57	49.90
47	36 51.00	121 49.80	11-10-71	32.58	35.67	31.74
48	36 50.80	121 53.60	11-24-71	11.61	40.22	48.17
49	36 50.20	121 50.20	11-10-71	21.97	41.69	36.34
1	36 47.25	121 48.90	7- 9-73	49.30	24.65	26.05
2	36 46.85	121 53.50	7- 9-73	24.74	29.69	45.57
3	36 46.35	121 49.00	7- 9-73	48.67	23.45	27.88
4	36 46.05	121 51.00	7- 9-73	22.73	34.00	43.27
10	36 45.10	121 50.00	7- 2-73	50.27	21.31	28.42
11	36 45.00	121 49.00	7- 2-73	58.45	19.30	22.25
14	36 44.20	121 52.25	6-21-73	63.92	12.68	23.40
16	36 44.00	121 49.50	7- 2-73	76.80	4.80	18.40
17	36 43.75	121 54.45	8- 9-73	19.85	16.66	63.49
19	36 43.35	121 56.25	8- 9-73	82.72	3.63	13.65
20	36 43.18	121 57.00	6-21-73	93.66	1.17	5.17
22	36 42.90	121 58.00	8- 9-73	11.79	11.41	76.81
23	36 42.55	121 53.30	6-21-73	64.41	9.31	26.28
25	36 41.70	121 55.00	7-16-73	7.93	19.07	73.00
26	36 41.55	121 55.50	7-16-73	5.89	19.05	75.06
29	36 40.90	121 56.40	7-16-73	9.36	14.84	75.80
34	36 39.10	121 53.08	8- 9-73	29.21	10.40	60.40
36	36 37.95	121 52.50	6-21-73	95.09	1.09	3.82
37	36 37.77	121 51.83	7-16-73	47.37	17.54	35.09
38	36 38.47	121 51.68	9-21-73	16.67	10.22	73.12

Table 5. VARIANCE OF SAMPLING MEASURED AT STATION 38.

Sample	Subsample	DDT (ppb)	DDD (ppb)	DDE (ppb)	TOTAL (ppb)
1	1	.687	.430	3.01	4.13
	2	.772	.470	2.90	4.14
	3	.550	.370	2.85	3.77
2	1	.561	.345	2.89	3.80
	2	.706	.333	2.38	3.42
	3	.801	.280	2.57	3.65
3	1	.663	.439	2.63	3.73
	2	.398	.315	2.96	3.67
	3	.405	.418	2.32	3.14
Mean		.6159	.3778	2.7233	3.7167
Variance		.02167	.00416	.06574	.09841
Standard Deviation		± .1472	± .0645	± .2564	± .3137
Standard Error		± .0491	± .0215	± .0855	± .1046
95% Confidence Limits		± .1131	± .0495	± .1971	± .2411

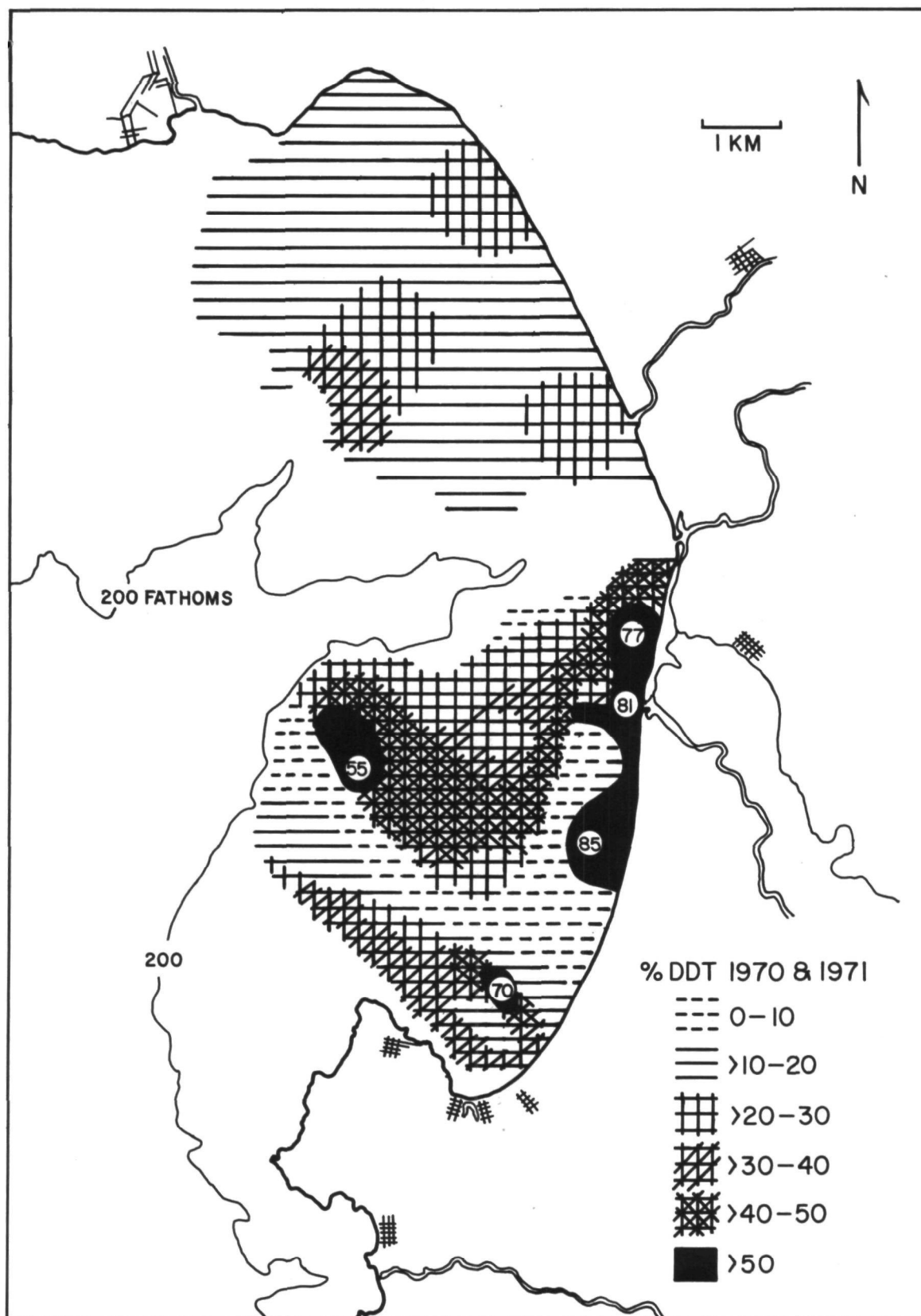


Figure 2. DDT as a percent of the total concentration of DDT, DDD, and DDE plotted for data obtained in 1970 and 1971. Circled numbers indicate actual percents in excess of 50%.

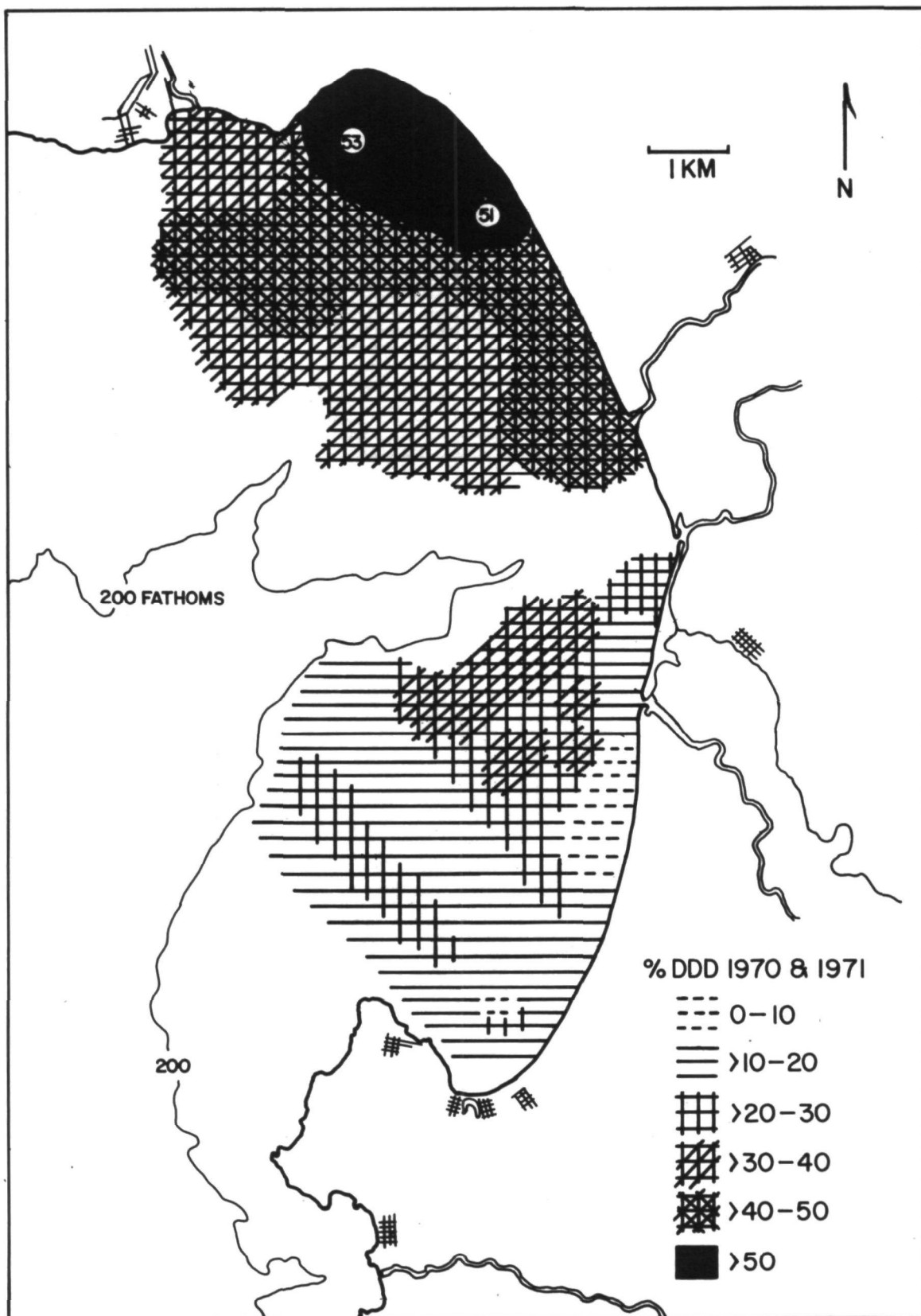


Figure 3. DDD as a percent of the total concentration of DDT, DDD, and DDE plotted for data obtained in 1970 and 1971. Circled numbers indicate actual percents in excess of 50%.

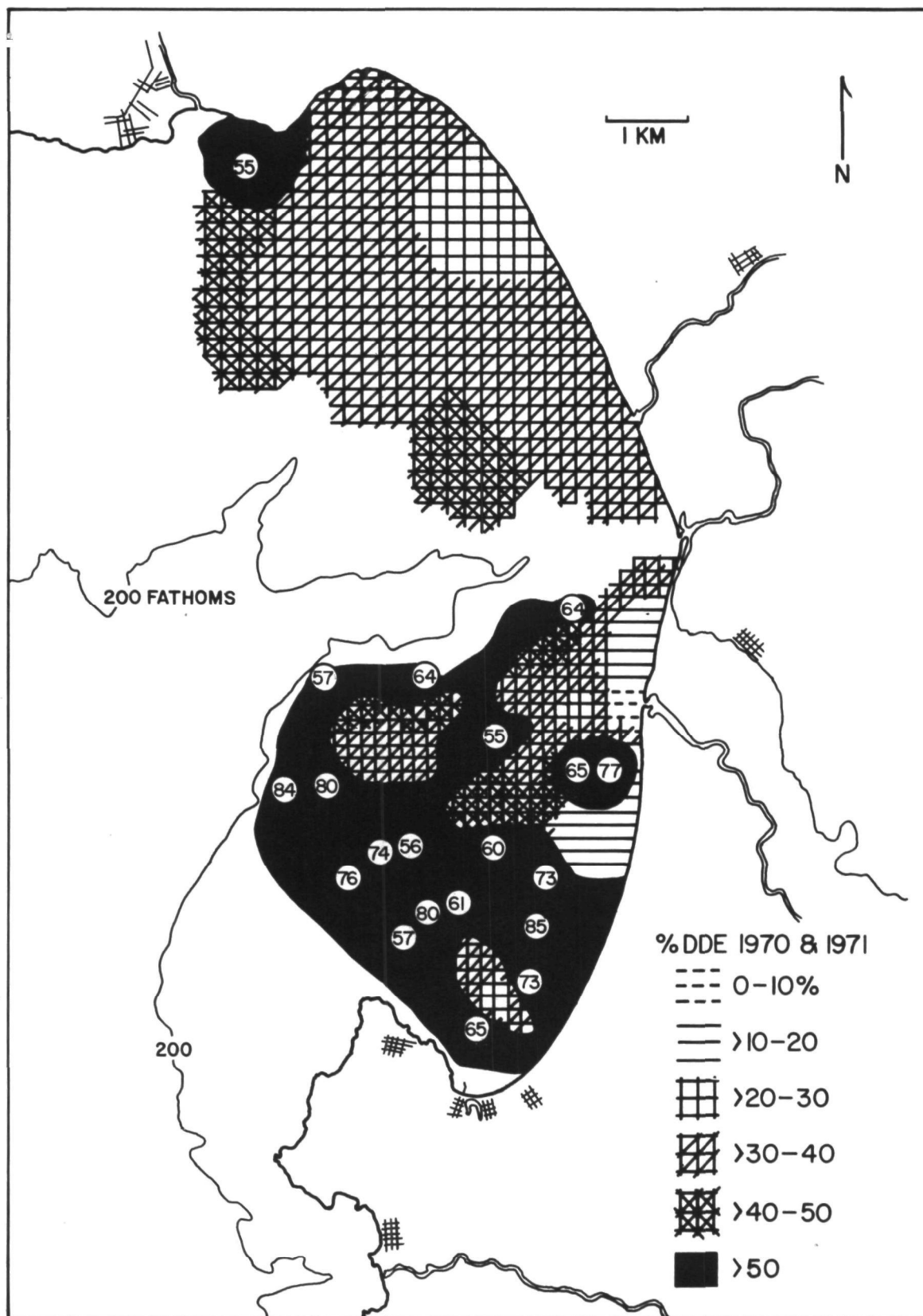


Figure 4. DDE as a percent of the total concentration of DDT, DDD, and DDE plotted for data obtained in 1970 and 1971. Circled numbers indicate actual percents in excess of 50%

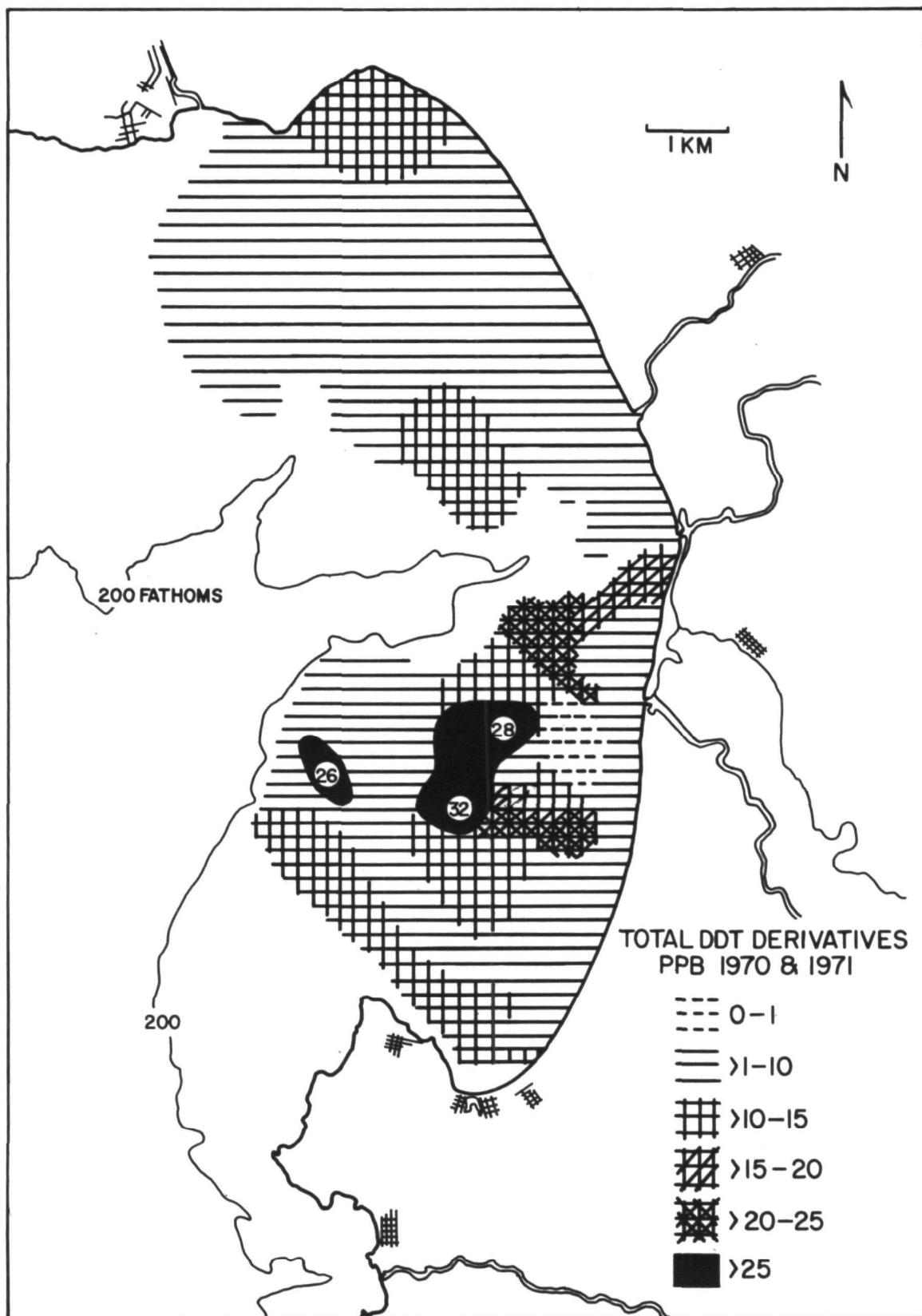


Figure 5. Total concentration in parts per billion of DDT, DDD, and DDE from data obtained in 1970 and 1971. Circled numbers indicate actual concentrations in excess of 50 ppb.

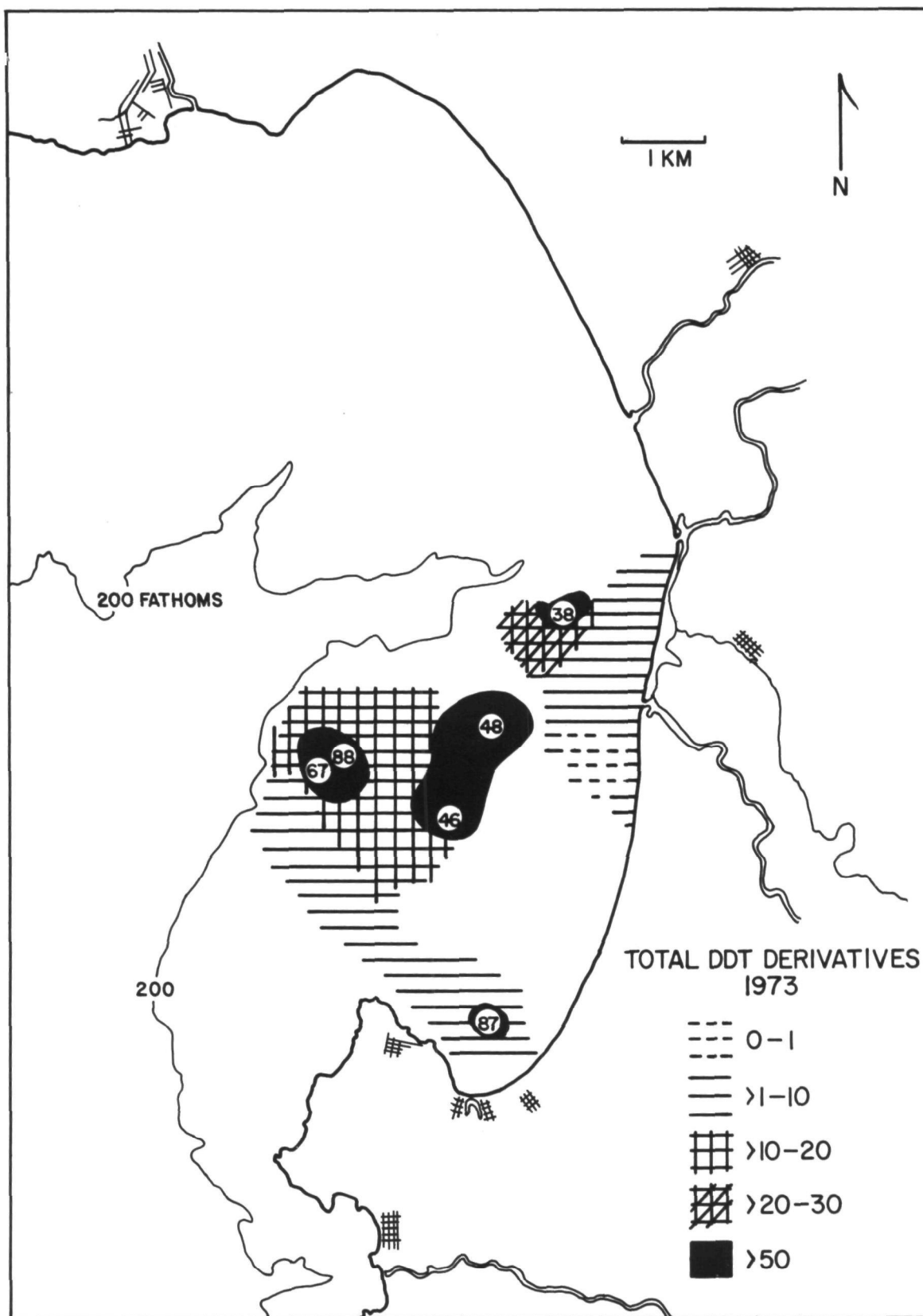


Figure 6. Total concentration in parts per billion of DDT, DDD, and DDE from data obtained in 1973. The blank portions of the area were not sampled. Circled numbers indicate actual concentrations in excess of 50 ppb.

ANALYSIS OF DYNAMICS

An approach to the analysis of the dynamics of sediment systems has been developed and has led to the development of Fortran programs permitting the rapid evaluation of data. The discussion of the approach to analysis will refer to output from these programs. The programs themselves with explanatory documentation are to be found in an appendix at the end of this report.

The first program requires sampling at the same set of stations at two points in time. The residue levels measured in sediments from the 19 stations sampled in both 1970 and 1973 constitute the data set used by this program. These data are presented as the first two pages of output, see Tables 6 and 7, followed by two pages showing the percent composition of total derivatives, see Tables 8 and 9. From the sums and means in Tables 6 and 7 it would appear that while DDT has shown an increase of several-fold the concentrations of DDD and DDE have changed very little. With respect to these latter two compounds input must be rather closely balanced with respect to output and decay. The changes in levels detected at individual stations must be a reflection of the rates of input of new material, output or removal both geographically and into other parts of the ecosystem, decay or decomposition within the sediment, and finally a shifting about of the material from sampling station to sampling station due primarily to the action of currents. The obvious complexity of the effect of these various rates has made the analysis of such a system extremely difficult. The approach presented here has necessitated the making of several simplifying assumptions. The utility of the method and the validity of the assumptions must await further evaluation, and the approach is intended more as a beginning than a final answer to the needs for methods of data analysis.

Figure 7 presents a diagram of the essential features of the system as it is envisaged. The individual stations where sediment samples were obtained are considered as compartments within the system of sediments in the southern portion of Monterey Bay. The diagram indicates that this system has a relationship to all other systems both geographical and of other kinds where the three compounds occur. Systems of different kinds would include the water above the sediment, the atmosphere above the water, organisms, etc. The effect of the rate of input, I , the rate of output, O , the rate of decay, D , and the rates of internal translocation, T_I and T_O , on the concentration within the system and within compartments is indicated.

A comparison of Figures 5 and 6 suggests that with continued input areas with the higher concentrations tend to increase in concentration due to the movement of the compounds within the system to these sinks or basins. Therefore, the amount of increase within any sediment compartment would appear to be related to the concentration already existing in that compartment. A similar relationship between the amount of decrease and concentration is less easily deduced from these Figures. However, the results of laboratory assays to be discussed in a later section have not revealed either a saturation of the decay process nor a stimulation by induction and selection of microbial populations that can be related to the concentration of these compounds. Instead the amount of decomposition appears to be a function of concentration. That the amount of translocation would be similarly related to concentration seems apparent.

Table 6. FIRST PAGE OF COMPUTER OUTPUT SHOWING CONCENTRATION OF POLLUTANT COMPOUNDS IN SEDIMENT FROM SAMPLE STATIONS AT FIRST SAMPLING TIME. C₁ IDENTIFIES AS CONCENTRATIONS AT TIME ONE.

C₁

Station	LOCATION		Date	DDT (ppb)	DDD (ppb)	DDE (ppb)	TOTAL
	Latitude	Longitude					
1	36 47.25	121 48.90	8-23-70	8.36	3.67	5.76	17.79
2	36 46.85	121 53.50	11-15-70	1.63	6.76	14.70	23.09
3	36 46.35	121 49.00	2-20-70	5.71	0.71	1.02	7.44
4	36 46.05	121 51.00	11-15-70	4.28	6.61	10.70	21.59
10	36 45.10	121 50.00	5-29-70	6.42	8.67	7.01	22.10
11	36 45.00	121 49.00	2-20-70	3.67	0.40	0.45	4.52
14	36 44.20	121 52.25	8-23-70	5.20	7.50	15.50	28.20
16	36 44.00	121 49.50	2-20-70	0.69	0.14	2.75	3.58
17	36 43.75	121 54.45	11-15-70	1.02	0.38	0.70	2.10
19	36 43.35	121 56.25	8-23-70	1.12	0.25	0.65	2.02
20	36 43.18	121 57.00	2- 8-70	0.0	5.00	20.50	25.50
22	36 42.90	121 58.00	2-20-70	0.0	0.35	1.92	2.27
23	36 42.55	121 53.30	8-23-70	13.20	5.73	13.00	31.93
25	36 41.70	121 55.00	2-20-70	1.22	0.53	2.40	4.15
26	36 41.55	121 55.50	2- 8-70	0.0	2.35	7.01	9.36
29	36 40.90	121 56.40	2-20-70	1.32	1.61	9.02	11.95
34	36 39.10	121 53.08	2- 8-70	2.44	0.66	2.40	5.50
36	36 37.95	121 52.50	2-20-70	2.65	2.79	10.00	15.44
37	36 37.77	121 51.83	2- 8-70	0.49	0.21	0.50	1.20
TOTALS				59.4199	54.3199	125.9899	239.7298
Mean				3.1274	2.8589	6.6310	12.6174
Standard Deviation				± 3.4385	± 2.9296	± 6.0673	± 10.1773
Standard Error				± 0.7889	± 0.6721	± 1.3919	± 2.3348
95% Confidence Limits				± 1.6574	± 1.4121	± 2.9245	± 4.9055

Table 7. SECOND PAGE OF COMPUTER OUTPUT SHOWING CONCENTRATION OF POLLUTANT COMPOUNDS IN SEDIMENT FROM SAMPLE STATIONS AT THE SECOND SAMPLING TIME. C₂ IDENTIFIES AS CONCENTRATIONS AT TIME TWO.

C₂

Station	LOCATION Latitude Longitude	Date	DDT (ppb)	DDD (ppb)	DDE (ppb)	TOTAL
1	36 47.25 121 48.90	7- 9-73	1.06	0.53	0.56	2.15
2	36 46.85 121 53.50	7- 9-73	9.50	11.40	17.50	38.40
3	36 46.35 121 49.00	7- 9-73	1.10	0.53	0.63	2.26
4	36 46.05 121 51.00	7- 9-73	3.63	5.43	6.91	15.97
10	36 45.10 121 50.00	7- 2-73	0.92	0.39	0.52	1.83
11	36 45.00 121 49.00	7- 2-73	2.18	0.72	0.83	3.73
14	36 44.20 121 52.25	6-21-73	30.60	6.07	11.20	47.87
16	36 44.00 121 49.50	7- 2-73	0.96	0.06	0.23	1.25
17	36 43.75 121 54.45	8- 9-73	5.41	4.54	17.30	27.25
19	36 43.35 121 56.25	8- 9-73	72.70	3.19	12.00	87.89
20	36 43.18 121 57.00	6-21-73	63.10	0.79	3.48	67.37
22	36 42.90 121 58.00	8- 9-73	0.93	0.90	6.06	7.89
23	36 42.55 121 53.30	6-21-73	29.90	4.32	12.20	46.42
25	36 41.70 121 55.00	7-16-73	1.14	2.74	10.49	14.37
26	36 41.55 121 55.50	7-16-73	0.68	2.20	8.67	11.55
29	36 40.90 121 56.40	7-16-73	0.70	1.11	5.67	7.48
34	36 39.10 121 53.08	8- 9-73	1.18	0.42	2.44	4.04
36	36 37.95 121 52.50	6-21-73	83.10	0.95	3.34	87.39
37	36 37.77 121 51.83	7-16-73	0.54	0.20	0.40	1.14
TOTALS			309.3296	46.4899	120.4299	476.2488
Mean			16.2805	2.4468	6.3384	25.0657
Standard Deviation			± 26.9909	± 2.8805	± 5.7417	± 29.2362
Standard Error			± 6.1921	± 0.6608	± 1.3172	± 6.7072
95% Confidence Limits			± 13.0097	± 1.3884	± 2.7675	± 14.0919

Table 8. THIRD PAGE OF COMPUTER OUTPUT SHOWING PERCENT OF TOTAL OF EACH OF THE THREE COMPOUNDS IN SEDIMENTS FROM SAMPLE STATIONS AT THE FIRST SAMPLING TIME. C₁ IDENTIFIES AS DATA FOR TIME ONE.

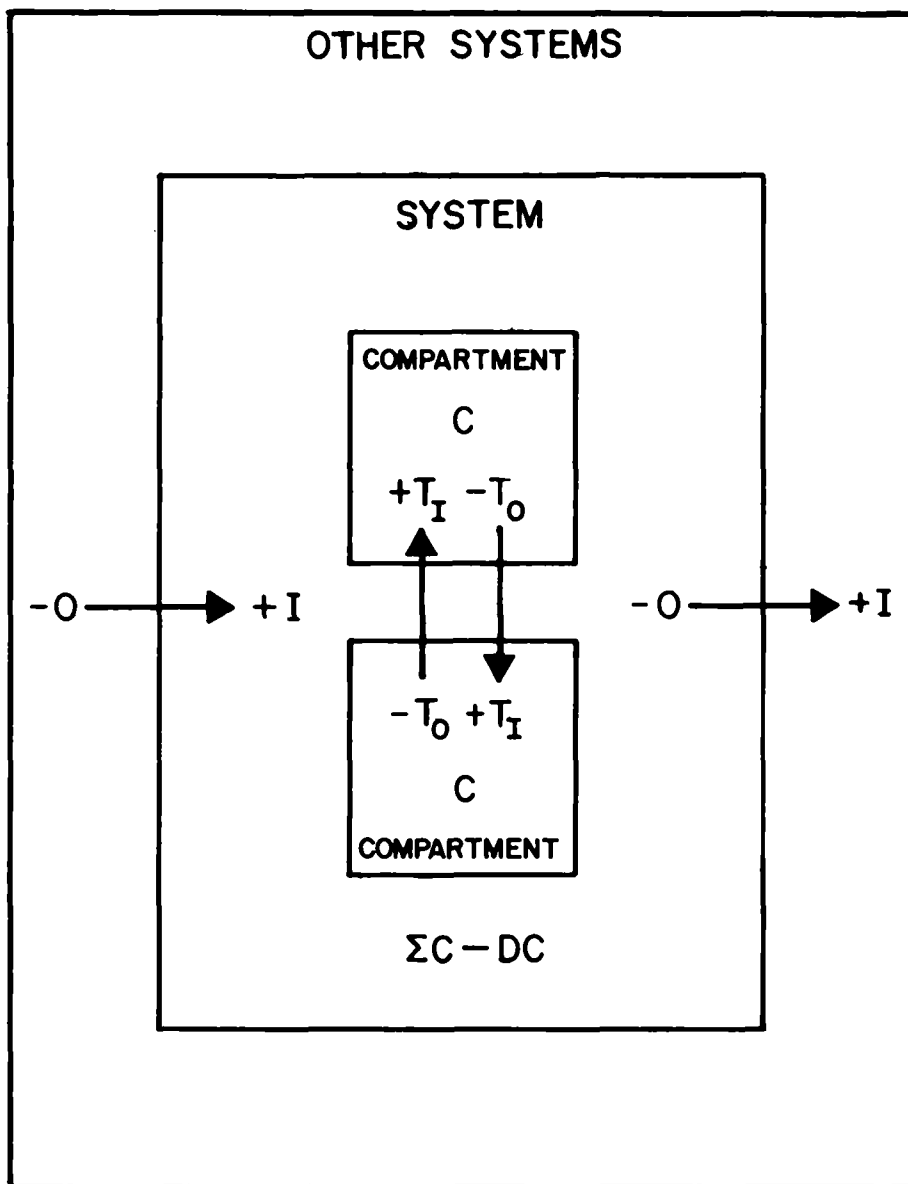
C₁

Station	LOCATION Latitude Longitude	Date	DDT (%)	DDD (%)	DDE (%)
1	36 47.25 121 48.90	8-23-70	46.99	20.63	32.38
2	36 46.85 121 53.50	11-15-70	7.06	29.28	63.66
3	36 46.35 121 49.00	2-20-70	76.75	9.54	13.71
4	36 46.05 121 51.00	11-15-70	19.82	30.62	49.56
10	36 45.10 121 50.00	5-29-70	29.05	39.23	31.72
11	36 45.00 121 49.00	2-20-70	81.19	8.85	9.96
14	36 44.20 121 52.25	8-23-70	18.44	26.60	54.96
16	36 44.00 121 49.50	2-20-70	19.27	3.91	76.82
17	36 43.75 121 54.45	11-15-70	48.57	18.10	33.33
19	36 43.35 121 56.25	8-23-70	55.45	12.38	32.18
20	36 43.18 121 57.00	2- 8-70	0.0	19.61	80.39
22	36 42.90 121 58.00	2-20-70	0.0	15.42	84.58
23	36 42.55 121 53.30	8-23-70	41.34	17.95	40.71
25	36 41.70 121 55.00	2-20-70	29.40	12.77	57.83
26	36 41.55 121 55.50	2- 8-70	0.0	25.11	74.89
29	36 40.90 121 56.40	2-20-70	11.05	13.47	75.48
34	36 39.10 121 53.08	2- 8-70	44.36	12.00	43.64
36	36 37.95 121 52.50	2-20-70	17.16	18.07	64.77
37	36 37.77 121 51.83	2- 8-70	40.83	17.50	41.67
TOTALS			586.7412	351.0149	962.2397
Mean			30.8811	18.4745	50.6442
Standard Deviation			± 24.2998	± 8.6373	± 22.2953
Standard Error			± 5.5748	± 1.9815	± 5.1149
95% Confidence Limits			± 11.7126	± 4.1632	± 10.7464

Table 9. FOURTH PAGE OF COMPUTER OUTPUT SHOWING PERCENT OF TOTAL OF EACH OF THE THREE COMPOUNDS IN SEDIMENT FROM SAMPLE STATIONS AT THE SECOND SAMPLING TIME. C₂ IDENTIFIES AS DATA FOR TIME TWO.

C₂

Station	LOCATION		Date	DDT (%)	DDD (%)	DDE (%)
	Latitude	Longitude				
1	36 47.25	121 48.90	7- 9-73	49.30	24.65	26.05
2	36 46.85	121 53.50	7- 9-73	24.74	29.69	45.57
3	36 46.35	121 49.00	7- 9-73	48.67	23.45	27.88
4	36 46.05	121 51.00	7- 9-73	22.73	34.00	43.27
10	36 45.10	121 50.00	7- 2-73	50.27	21.31	28.42
11	36 45.00	121 49.00	7- 2-73	58.45	19.30	22.25
14	36 44.20	121 52.25	6-21-73	63.92	12.68	23.40
16	36 44.00	121 49.50	7- 2-73	76.80	4.80	18.40
17	36 43.75	121 54.45	8- 9-73	19.85	16.66	63.49
19	36 43.35	121 56.25	8- 9-73	82.72	3.63	13.65
20	36 43.18	121 57.00	6-21-73	93.66	1.17	5.17
22	36 42.90	121 58.00	8- 9-73	11.79	11.41	76.81
23	36 42.55	121 53.30	6-21-73	64.41	9.31	26.28
25	36 41.70	121 55.00	7-16-73	7.93	19.07	73.00
26	36 41.55	121 55.50	7-16-73	5.89	19.05	75.06
29	36 40.90	121 56.40	7-16-73	9.36	14.84	75.80
34	36 39.10	121 53.08	8- 9-73	29.21	10.40	60.40
36	36 37.95	121 52.50	6-21-73	95.09	1.09	3.82
37	36 37.77	121 51.83	7-16-73	47.37	17.54	35.09
TOTALS				862.1616	294.0427	743.7920
Mean				45.3769	15.4759	39.1469
Standard Deviation				± 29.2068	± 9.2122	± 24.6220
Standard Error				± 6.7005	± 2.1134	± 5.6487
95% Confidence Limits				± 14.0777	± 4.4403	± 11.8679



SYSTEMS

$$C_2 = C_1 (1 + I - O - D)^N \quad T_O = T_I$$

COMPARTMENTS

$$C_2 = C_1 (1 + I + T_I - T_O - O - D)^N \quad T_O \neq T_I$$

C_1 = CONCENTRATION OF RESIDUE AT TIME 1

C_2 = CONCENTRATION OF RESIDUE AT TIME 2

I = RATE OF INPUT OF RESIDUE

O = RATE OF OUTPUT OF RESIDUE

D = RATE OF DECAY

T_O = RATE OF TRANSLOCATION OUT OF A COMPARTMENT

T_I = RATE OF TRANSLOCATION INTO A COMPARTMENT

Figure 7. Model of the system of sediment compartments and this system's relation to other systems.

Therefore, for the estimation of the overall rate of change in a compartment, i.e., the resultant of the various rates affecting concentration, the following expression was solved for K,

$$C_2 = C_1 e^{KN} \quad (1)$$

C_1 and C_2 are the concentrations within the compartment at time one and time two, N is the length of the time interval in years, and e is the natural logarithm base. K is a nominal percentage rate in the form of a decimal fraction resulting in continuous compounding, and is converted to an annual rate for the expression,

$$C_2 = C_1 (1+K)^N \quad (2)$$

The results of these calculations for the three compounds are presented as the fifth, sixth, and seventh pages of computer output in Tables 10, 11, and 12. In these tables the values of K are sorted into positive and negative values for purposes discussed below. Compartments which showed a zero concentration at time one were adjusted by substitution of 0.004 ppb, a value generally just below the level of detection in the analyses.

The standard deviation of these estimates was approximated through the use of the expression for the standard deviation of a function of two random variables (Papoulis, 1965),

$$\sigma_{K(C_1, C_2)}^2 \sim \left(\frac{\partial K}{\partial C_1}\right)^2 \sigma_{C_1}^2 + \left(\frac{\partial K}{\partial C_2}\right)^2 \sigma_{C_2}^2 + 2 \frac{\partial K}{\partial C_1} \frac{\partial K}{\partial C_2} \sigma_{C_1 C_2} \quad (3)$$

For ease in computation only two variables at a time were used in developing this approximation to the standard deviation.

If we assume that the rate of change within the system can be approximated by the mean rate of change of its separate compartments, the mean of the K values becomes an estimate of the rate of net change of the system.

$$\text{Net rate of change} = I - (O+D) \quad (4)$$

This net rate of change is unaffected by the rates of internal translocation, T_I and T_O , which are equal in magnitude and opposite in sign. The net rate of change is the sum of two other mean rates. One is the rate of input, I , which can be estimated by the mean of the positive K 's, and the other is obtained as the mean of the negative K 's and may be taken as an estimate of $(O+D)$ in equation 4.

The mean of the differences between each K and the net rate of change, that is the mean deviation from the mean of K , becomes an estimate of T_O and T_I . The results of these calculations are included in Tables 10, 11, and 12.

The separation of the rate O and D is more difficult and several approaches have been attempted. The decimal fraction of the input rate that is translocated within the system, T_I/I , differs from compound to compound: DDT, 0.665; DDD, 0.882; and DDE, 0.860. One explanation for this difference is that they reflect differences in the rates of decomposition within the sediments. Based upon this assumption the rate O and D have been estimated by the following equations,

Table 10. FIFTH PAGE OF COMPUTER OUTPUT SHOWING THE RATE OF CHANGE, K,
FOR DDT IN EACH SEDIMENT COMPARTMENT.

Station	C ₂ DDT	C ₁ DDT	N	+K	-K	+K + -K	+K - Net R	-K - Net R
1	1.06	8.36	2.8795	0.0	-0.5119	-0.5119	0.0	-2.0906
2	9.50	1.63	2.6493	0.9452	0.0	0.9452	0.0	-0.6336
3	1.10	5.71	3.3836	0.0	-0.3854	-0.3854	0.0	-1.9641
4	3.63	4.28	2.6493	0.0	-0.0603	-0.0603	0.0	-1.6390
10	0.92	6.42	3.0959	0.0	-0.4661	-0.4661	0.0	-2.0449
11	2.18	3.67	3.3644	0.0	-0.1434	-0.1434	0.0	-1.7222
14	30.60	5.20	2.8301	0.8706	0.0	0.8706	0.0	-0.7082
16	0.96	0.69	3.3644	0.1031	0.0	0.1031	0.0	-1.4756
17	5.41	1.02	2.7342	0.8408	0.0	0.8408	0.0	-0.7380
19	72.70	1.12	2.9644	3.0866	0.0	3.0866	1.5079	0.0
20	63.10	0.0	3.3671	16.6503	0.0	16.6503	15.0715	0.0
22	0.93	0.0	3.4685	3.8113	0.0	3.8113	2.2325	0.0
23	29.90	13.20	2.8301	0.3350	0.0	0.3350	0.0	-1.2438
25	1.14	1.22	3.4027	0.0	-0.0197	-0.0197	0.0	-1.5985
26	0.68	0.0	3.4356	3.4588	0.0	3.4588	1.8800	0.0
29	0.70	1.32	3.4027	0.0	-0.1701	-0.1701	0.0	-1.7488
34	1.18	2.44	3.5014	0.0	-0.1874	-0.1874	0.0	-1.7661
36	83.10	2.65	3.3342	1.8105	0.0	1.8105	0.2317	0.0
37	0.54	0.49	3.4356	0.0287	0.0	0.0287	0.0	-1.5501
Totals	309.3296	59.4199	60.0930	31.9407	-1.9442	29.9964	20.9236	-20.9235
Mean	16.2805	3.1274	3.1628	1.6811	-0.1023	1.5788	1.1012	-1.1012
S.D.	± 26.9909	± 3.4385	± 0.3100	± 0.9016	± 0.0984	± 1.0000	± 0.8738	± 0.1262
S.E.	± 6.1921	± 0.7889	± 0.0711	± 0.2068	± 0.0226	± 0.2294	± 0.2005	± 0.0289
95% C.L.	± 13.0097	± 1.6574	± 0.1494	± 0.4346	± 0.0474	± 0.4820	± 0.4212	± 0.0608

Table 11. SIXTH PAGE OF COMPUTER OUTPUT SHOWING THE RATE OF CHANGE, K,
FOR DDD IN EACH SEDIMENT COMPARTMENT.

Station	C ₂ DDD	C ₁ DDD	N	+K	-K	+K + -K	+K - Net R	-K - Net R
1	0.53	3.67	2.8795	0.0	-0.4893	-0.4893	0.0	-0.5714
2	11.40	6.76	2.6493	0.2181	0.0	0.2181	0.1360	0.0
3	0.53	0.71	3.3836	0.0	-0.0828	-0.0828	0.0	-0.1648
4	5.43	6.61	2.6493	0.0	-0.0715	-0.0715	0.0	-0.1536
10	0.39	8.67	3.0959	0.0	-0.6328	-0.6328	0.0	-0.7148
11	0.72	0.40	3.3644	0.1909	0.0	0.1909	0.1089	0.0
14	6.07	7.50	2.8301	0.0	-0.0720	-0.0720	0.0	-0.1541
16	0.06	0.14	3.3644	0.0	-0.2226	-0.2226	0.0	-0.3047
17	4.54	0.38	2.7342	1.4774	0.0	1.4774	1.3953	0.0
19	3.19	0.25	2.9644	1.3607	0.0	1.3607	1.2787	0.0
20	0.79	5.00	3.3671	0.0	-0.4219	-0.4219	0.0	-0.5039
22	0.90	0.35	3.4685	0.3130	0.0	0.3130	0.2309	0.0
23	4.32	5.73	2.8301	0.0	-0.0950	-0.0950	0.0	-0.1770
25	2.74	0.53	3.4027	0.6206	0.0	0.6206	0.5386	0.0
26	2.20	2.35	3.4356	0.0	-0.0190	-0.0190	0.0	-0.1011
29	1.11	1.61	3.4027	0.0	-0.1035	-0.1035	0.0	-0.1856
34	0.42	0.66	3.5014	0.0	-0.1211	-0.1211	0.0	-0.2031
36	0.95	2.79	3.3342	0.0	-0.2761	-0.2761	0.0	-0.3582
37	0.20	0.21	3.4356	0.0	-0.0141	-0.0141	0.0	-0.0961
Totals	46.4899	54.3199	60.0930	4.1806	-2.6218	1.5588	3.6884	-3.6884
Mean	2.4468	2.8589	3.1628	0.2200	-0.1380	0.0820	0.1941	-0.1941
S.D.	± 2.8805	± 2.9296	± 0.3100	± 0.7233	± 0.2767	± 1.0000	± 0.7233	± 0.2767
S.E.	± 0.6608	± 0.6721	± 0.0711	± 0.1659	± 0.0635	± 0.2294	± 0.1659	± 0.0635
95% C.L.	± 1.3884	± 1.4121	± 0.1494	± 0.3486	± 0.1334	± 0.4820	± 0.3486	± 0.1334

Table 12. SEVENTH PAGE OF COMPUTER OUTPUT SHOWING THE RATE OF CHANGE, K, FOR DDE IN EACH SEDIMENT COMPARTMENT.

Station	C ₂ DDE	C ₁ DDE	N	+K	-K	+K + -K	+K - Net R	-K - Net R
1	0.56	5.76	2.8795	0.0	-0.5549	-0.5549	0.0	-0.6726
2	17.50	14.70	2.6493	0.0680	0.0	0.0680	0.0	-0.0497
3	0.63	1.02	3.3836	0.0	-0.1327	-0.1327	0.0	-0.2505
4	6.91	10.70	2.6493	0.0	-0.1522	-0.1522	0.0	-0.2699
10	0.52	7.01	3.0959	0.0	-0.5684	-0.5684	0.0	-0.6861
11	0.83	0.45	3.3644	0.1996	0.0	0.1996	0.0818	0.0
14	11.20	15.50	2.8301	0.0	-0.1085	-0.1085	0.0	-0.2262
16	0.23	2.75	3.3644	0.0	-0.5217	-0.5217	0.0	-0.6394
17	17.30	0.70	2.7342	2.2318	0.0	2.2318	2.1141	0.0
19	12.00	0.65	2.9644	1.6740	0.0	1.6740	1.5563	0.0
20	3.48	20.50	3.3671	0.0	-0.4094	-0.4094	0.0	-0.5272
22	6.06	1.92	3.4685	0.3929	0.0	0.3929	0.2752	0.0
23	12.20	13.00	2.8301	0.0	-0.0222	-0.0222	0.0	-0.1399
25	10.49	2.40	3.4027	0.5426	0.0	0.5426	0.4249	0.0
26	8.67	7.01	3.4356	0.0638	0.0	0.0638	0.0	-0.0539
29	5.67	9.02	3.4027	0.0	-0.1275	-0.1275	0.0	-0.2453
34	2.44	2.40	3.5014	0.0047	0.0	0.0047	0.0	-0.1130
36	3.34	10.00	3.3342	0.0	-0.2803	-0.2803	0.0	-0.3980
37	0.40	0.50	3.4356	0.0	-0.0629	-0.0629	0.0	-0.1806
Totals	120.4299	125.9899	60.0930	5.1774	-2.9407	2.2367	4.4522	-4.4522
Mean	6.3384	6.6310	3.1628	0.2725	-0.1548	0.1177	0.2343	-0.2343
S.D.	± 5.7417	± 6.0673	± 0.3100	± 0.7781	± 0.2243	± 1.0024	± 0.7761	± 0.2262
S.E.	± 1.3172	± 1.3919	± 0.0711	± 0.1785	± 0.0515	± 0.2300	± 0.1781	± 0.0519
95% C.L.	± 2.7675	± 2.9245	± 0.1494	± 0.3750	± 0.1081	± 0.4831	± 0.3741	± 0.1091

$$O = \frac{T_I}{I} (O+D) \quad (5)$$

$$D = (1.0 - \frac{T_I}{I}) (O+D) \text{ or } D = (O+D) - O \quad (6)$$

The residence time, T_R , and lifetime, T_L , in years, are calculated as the corresponding reciprocals.

$$T_R = 1.0/(O+D) \quad (7)$$

$$T_L = 1.0/D \quad (8)$$

The last three pages of computer output present a summary of these estimations and are presented in Tables 13, 14, and 15.

The effect of substitution of a minimal value for zero concentrations was investigated by reducing the set of sample stations to sixteen and elimination of all stations showing a zero concentration of DDT at time one. While there was some effect upon the estimates of rates as the system was reduced in size, only the estimates of T_O for DDT were significantly different when tested by the "test of equality of the means of two samples whose variances are assumed to be unequal" (Sokol and Rohlf, 1969). The difference between the other estimates was very small compared to the standard deviation of these estimates. Table 16 presents for comparison the set of rates for the nineteen and sixteen station data sets.

The approach to analysis of the data which provided these estimates of system rates requires sampling at the same stations at two different times. However, as presented in Table 3, there is additional data available with respect to the south bay system at time one. This additional data can not be used by the approach to analysis presented so far. More stations were sampled in the first sampling period than were sampled in the second, and the approach requires pairs of samples identical except for time of sampling. An additional program was written to permit analysis of a system where sampling does not meet the requirements of the first approach. This second program treats all samples as unpaired and evaluates the rate of change, K , at the different sample locations by comparison of the actual measurement at that station at time one or time two with the mean concentrations of the system at either time one or time two. That is, a measurement at time one is paired with the mean concentration at time two and vice versa for the evaluation of K . Further the time interval, N , is evaluated as the interval between the time of actual sample of one sampling time and the mean time of the other sampling period. Equation 1 becomes,

$$\bar{C}_2 = C_1 e^{KN} \quad 9.$$

$$\text{with } N = \bar{T}_2 - T_1$$

Table 13. EIGHTH PAGE OF COMPUTER OUTPUT SHOWING A SUMMARY OF THE ANNUAL SYSTEM RATES EXPRESSED AS DECIMAL FRACTIONS OF THE MEAN CONCENTRATION OF DDT PRESENT IN THE SYSTEM.

System of Rates for DDT				S.D.		S.E.		95% Limit
Net rate of change	= Net = +	1.5788	±	1.0000	±	0.2294	±	0.4820
Translocation into compartments	= T_I = +	1.1012	±	0.8738	±	0.2005	±	0.4212
Translocation out of compartments	= T_O = -	1.1012	±	0.1262	±	0.0289	±	0.0608
Input	= I = +	1.6811	±	0.9016	±	0.2068	±	0.4346
Output and Decay	= O+D = -	0.1023	±	0.0984	±	0.0226	±	0.0474
Output from System	= O = -	0.0670	±	0.0644	±	0.0148	±	0.0311
Decay	= D = -	0.0353	±	0.0339	±	0.0078	±	0.0164
Lifetime in years	= T_L =	28.3322	±	27.2386	±	6.2490	±	13.1291
Residence time in years	= T_R =	9.7724	±	9.3952	±	2.1554	±	4.5285
Summary Equation for the System—								
DDT	Mean C_2	Mean C_1	I	T_I	T_O	O	D	N
	16.2805	= 3.1274	(1.0 + 1.6811 + 1.1012 - 1.1012 - 0.0670 - 0.0353)					3.1628

Table 14. NINTH PAGE OF COMPUTER OUTPUT SHOWING A SUMMARY OF THE ANNUAL SYSTEM RATES EXPRESSED AS DECIMAL FRACTIONS OF THE MEAN CONCENTRATION OF DDD PRESENT IN THE SYSTEM.

System of Rates for DDD				S.D.		S.E.		95% Limit
Net rate of change	= Net = +	0.0820	±	1.0000	±	0.2294	±	0.4820
Translocation into compartments	= T _I = +	0.1941	±	0.7233	±	0.1659	±	0.3486
Translocation out of compartments	= T _O = -	0.1941	±	0.2767	±	0.0635	±	0.1334
Input	= I = +	0.2200	±	0.7233	±	0.1659	±	0.3486
Output and Decay	= O+D = -	0.1380	±	0.2767	±	0.0635	±	0.1334
Output from System	= O = -	0.1217	±	0.2441	±	0.0560	±	0.1177
Decay	= D = -	0.0162	±	0.0326	±	0.0075	±	0.0157
Lifetime in years	= T _L =	61.5459	±	123.4241	±	28.3154	±	59.4907
Residence time in years	= T _R =	7.2469	±	14.5330	±	3.3341	±	7.0049
Summary Equation for the System—								
DDD	Mean C ₂	Mean C ₁	I	T _I	T _O	O	D	N
	2.4468	= 2.8589	(1.0 + 0.2200 + 0.1941 - 0.1941 - 0.1217 - 0.0162)					3.1628

Table 15. TENTH PAGE OF COMPUTER OUTPUT SHOWING A SUMMARY OF THE ANNUAL SYSTEM RATES EXPRESSED AS DECIMAL FRACTIONS OF THE MEAN CONCENTRATION OF DDE PRESENT IN THE SYSTEM.

System of Rates for DDE				S.D.		S.E.		95% Limit
Net rate of change	= Net = +	0.1177	±	1.0024	±	0.2300	±	0.4831
Translocation into compartments	= T_I = +	0.2343	±	0.7761	±	0.1781	±	0.3741
Translocation out of compartments	= T_O = -	0.2343	±	0.2262	±	0.0519	±	0.1091
Input	= I = +	0.2725	±	0.7781	±	0.1785	±	0.3750
Output and Decay	= O+D = -	0.1548	±	0.2243	±	0.0515	±	0.1081
Output from System	= O = -	0.1331	±	0.1929	±	0.0442	±	0.0930
Decay	= D = -	0.0217	±	0.0314	±	0.0072	±	0.0151
Lifetime in years	= T_L =	46.1286	±	66.8453	±	15.3354	±	32.2196
Residence time in years	= T_R =	6.4611	±	9.3629	±	2.1480	±	4.5129
Summary Equation for the System—								
DDE	Mean C_2	Mean C_1	I	T_I	T_O	O	O_D	N
	6.3384	= 6.6310	(1.0 + 0.2725 + 0.2343 - 0.2343 - 0.1331 - 0.0217)					3.1628

Table 16. COMPARISON OF ESTIMATES OBTAINED FROM THE 16 AND 19 STATION DATA SETS AND USING ACTUAL PAIRED SAMPLE ANALYSES STANDARD DEVIATIONS [S.D.] AND COEFFICIENTS OF VARIATION [C.V.] ARE INCLUDED.

	16 STATION DATA SET			19 STATION DATA SET		
	Estimate	S.D.	C.V. (%)	Estimate	S.D.	C.V. (%)
DDT						
\bar{C}_1 (ppb)	3.7137	\pm 3.4446	92.8	3.1274	\pm 3.4385	109.9
\bar{C}_2 (ppb)	15.2887	\pm 26.3645	172.4	16.2805	\pm 26.9909	165.8
Net	+ 0.3798	\pm 1.0000	263.3	+ 1.5788	\pm 1.0000	63.3
I	+ 0.5013	\pm 0.7556	150.7	+ 1.6811	\pm 0.9016	53.6
O + D	- 0.1215	\pm 0.2444	138.2	- 0.1023	\pm 0.0984	96.2
T_O	- 0.3534	\pm 0.2591	73.3	- 1.1012	\pm 0.1262	11.5
T_I	+ 0.3534	\pm 0.7409	209.6	+ 1.1012	\pm 0.8738	79.3
O	- 0.0857	\pm 0.1723	201.1	- 0.0670	\pm 0.0644	96.1
D	- 0.0358	\pm 0.0721	201.4	- 0.0353	\pm 0.0339	96.0
T_L (years)	27.9014	\pm 56.1105	201.1	28.3322	\pm 24.2386	96.1
T_R (years)	8.2294	\pm 16.5496	201.1	9.7724	\pm 9.3952	96.1
DDD						
\bar{C}_1 (ppb)	2.9137	\pm 3.0908	106.1	2.8589	\pm 2.9296	102.5
\bar{C}_2 (ppb)	2.6625	\pm 3.0921	116.1	2.4468	\pm 2.8805	117.7
Net	+ 0.1054	\pm 1.0000	948.8	+ 0.0820	\pm 1.0000	1219.5
I	+ 0.2417	\pm 0.7279	301.2	+ 0.2200	\pm 0.7233	328.8
O + D	- 0.1363	\pm 0.2721	199.6	- 0.1380	\pm 0.2767	200.5
T_O	- 0.2088	\pm 0.2721	130.3	- 0.1941	\pm 0.2767	142.6
T_I	+ 0.2088	\pm 0.7279	348.6	+ 0.1941	\pm 0.7233	5009.4
O	- 0.1177	\pm 0.2350	199.7	+ 0.1217	\pm 0.2441	200.6
D	- 0.0186	\pm 0.0371	199.5	- 0.0162	\pm 0.0326	201.2
T_L (years)	53.8306	\pm 107.4660	199.6	61.5459	\pm 123.4241	200.5
T_R (years)	7.3364	\pm 14.6462	199.6	7.2469	\pm 14.5330	200.5
DDE						
\bar{C}_1 (ppb)	6.0350	\pm 5.4299	90.0	6.6310	\pm 6.0673	91.5
\bar{C}_2 (ppb)	6.3887	\pm 6.2166	97.3	6.3384	\pm 5.7417	90.6
Net	+ 0.1368	\pm 1.0030	733.2	+ 0.1177	\pm 1.0024	851.7
I	+ 0.2950	\pm 0.7843	265.9	+ 0.2725	\pm 0.7781	285.5
O + D	- 0.1582	\pm 0.2186	138.2	- 0.1548	\pm 0.2243	144.9
T_O	- 0.2563	\pm 0.2211	86.3	- 0.2343	\pm 0.2262	96.5
T_I	+ 0.2563	\pm 0.7818	305.0	+ 0.2343	\pm 0.7761	332.1
O	- 0.1374	\pm 0.1899	138.2	- 0.1331	\pm 0.1929	144.9
D	- 0.0208	\pm 0.0287	138.0	- 0.0217	\pm 0.0314	144.7
T_L (years)	48.1189	\pm 66.4924	138.2	46.1286	\pm 66.8543	144.9
T_R (years)	6.3211	\pm 8.7347	138.2	6.4611	\pm 9.3629	144.9

\bar{T}_2 = mean time of second sampling period

T_1 = time of actual sampling in first sampling period

$$\text{and } C_2 = \bar{C}_1 e^{KN} \quad (10)$$

with $N = T_2 - \bar{T}_1$

T_2 = time of actual sampling in second sampling period

\bar{T}_1 = mean time of first sampling period.

Table 17 presents the estimates of the system obtained using this pairing with means approach. Once again the effect of substitution of a minimal value for zero concentrations was explored by eliminating stations with zero concentration thus providing the subset of 49 samples from the complete set of 57. Except for the estimates of T_O for DDT, there was no significant difference between the two sets of estimates once again, nor are these estimates significantly different from either of the sets of estimates based on the 16 and 19 station data sets. The principal effect of inclusion or exclusion of the zero level values with substitution of a minimal value is upon the estimates of the rates of input, I , translocation, T_I and T_O , and the net rate. The stations showing a zero concentration of DDT at time one show high positive rates of change, and therefore, have a particularly marked effect on the positive rate estimates as well as those based to at least some extent upon these positive rate estimates.

The second approach which uses sample values paired to mean values should find use in the analysis of systems where real paired values are impossible to obtain. Animals which are sacrificed at the time of sampling obviously can not be resampled at another point in time. The use of sample values at one sample time paired to the mean value of another permits estimation of system rates for the population. The comparison between the two approaches to these estimates that is presented here indicates that the use of mean values in pairing gives a close approximation of rate estimates obtained with real paired values.

Both of these approaches to the estimation of system rates are dependent upon variability in concentration level and rate of change within compartments. It is essential to these methods of analysis that individual compartments show the effect of the various processes to different degrees. If all the concentration levels and rates of change within compartments were the same, it would be possible to gain an estimate of net rate of change only. Therefore, these approaches to estimation of system rates are dependent upon variability in environmental samples of the system and make use of this variability for estimating the rates of the various processes.

Table 17. COMPARISON OF ESTIMATES OBTAINED FROM THE 49 AND 57 STATION DATA SETS AND USING SAMPLE ANALYSES PAIRED WITH MEAN CONCENTRATION LEVELS. STANDARD DEVIATIONS [S.D.] AND COEFFICIENTS OF VARIATION [C.V.] ARE INCLUDED.

	49 SAMPLE DATA SET			57 SAMPLE DATA SET		
	Estimate	S.D.	C.V. (%)	Estimate	S.D.	C.V. (%)
DDT						
—						
C ₁ (ppb)	3.9576	± 4.1746	105.4	3.1019	± 4.0336	130.0
—						
C ₂ (ppb)	15.4975	± 26.5034	171.0	15.4975	± 26.5034	171.0
Net	+ 0.5905	± 1.0000	169.3	+ 2.2567	± 1.0000	44.3
I	+ 0.6819	± 0.6374	93.5	+ 2.3233	± 0.9204	39.6
O + D	- 0.0913	± 0.3626	397.2	- 0.0667	± 0.0796	119.3
T _O	- 0.3234	± 0.3966	122.6	- 1.4256	± 0.1513	10.6
T _I	+ 0.3234	± 0.6034	186.6	+ 1.4256	± 0.8487	59.5
O	- 0.0433	± 0.1720	397.2	- 0.0409	± 0.0488	119.3
D	- 0.0480	± 0.1906	397.1	- 0.0258	± 0.0307	119.0
T _L (years)	20.8292	± 82.7111	397.1	38.8090	± 46.2947	119.3
T _R (years)	10.9502	± 43.4823	397.1	14.9951	± 17.8875	119.3
DDD						
—						
C ₁ (ppb)	2.4107	± 2.5354	105.2	2.2743	± 2.3532	103.5
—						
C ₂ (ppb)	2.3435	± 2.8415	121.3	2.3435	± 2.8415	121.3
Net	+ 0.1283	± 1.0000	779.4	+ 0.1587	± 1.0000	630.1
I	+ 0.2703	± 0.6357	235.2	+ 0.2813	± 0.6329	225.0
O + D	- 0.1420	± 0.3643	256.5	- 0.1226	± 0.3671	299.4
T _O	- 0.2095	± 0.3653	174.4	- 0.2039	± 0.3698	180.9
T _I	+ 0.2095	± 0.6347	303.0	+ 0.2039	± 0.6311	309.5
O	- 0.1101	± 0.2823	256.4	- 0.0889	± 0.2662	299.4
D	- 0.0319	± 0.0820	257.1	- 0.0337	± 0.1010	299.7
T _L (years)	31.3031	± 80.3119	256.6	29.6518	± 88.7883	299.4
T _R (years)	7.0424	± 18.0682	256.6	8.1558	± 24.4216	299.4
DDE						
—						
C ₁ (ppb)	5.1138	± 4.4111	86.3	5.3681	± 4.8069	89.5
—						
C ₂ (ppb)	6.1575	± 5.6469	91.7	6.1575	± 5.6469	91.7
Net	+ 0.1748	± 1.0010	572.7	+ 0.1793	± 1.0009	558.2
I	+ 0.2802	± 0.6628	236.5	+ 0.2785	± 0.6787	243.7
O + D	- 0.1054	± 0.3382	320.9	- 0.0993	± 0.3222	324.5
T _O	- 0.1946	± 0.3466	178.1	- 0.1906	± 0.3311	173.7
T _I	+ 0.1946	± 0.6544	336.3	+ 0.1906	± 0.6697	351.4
O	- 0.0732	± 0.2348	320.8	- 0.0679	± 0.2204	324.6
D	- 0.0322	± 0.1033	320.8	- 0.0314	± 0.1018	324.2
T _L (years)	31.0400	± 99.5728	320.8	31.8905	± 103.4957	324.5
T _R (years)	9.4853	± 30.4277	320.8	10.0735	± 32.6922	324.5

For any set of estimates of I , $(O+D)$, T_I and T_O , based on a number of samples, n , there is a distribution of K 's with a minimal variance. The members of the distribution can be determined through one of the following sets of equations:

Where the net rate of change, $I + (O+D)$, is positive,

$$j = \frac{nI - nT_I}{I + (O+D)} \quad \text{and } j \text{ is an integer obtained without rounding.} \quad (11)$$

$$I + (O+D) + \frac{nT_I}{j} = K_1, K_2 \dots K_j \quad (12)$$

$$\text{If } \sum_{1}^j K \leq nI$$

$$nI - jK_1 = K_j + 1 \quad (13)$$

$$\frac{n(O+D)}{n-j-1} = K_j + 2, K_j + 3 \dots K_n \quad (14)$$

$$\text{If } \sum_{1}^j K = nI$$

$$\frac{n(O+D)}{n-j} = K_j + 1, K_j + 2 \dots K_n \quad (15)$$

Where the net rate of change, $I + (O+D)$, is zero,

$$j = \frac{n}{2} \quad \text{and } j \text{ is an integer obtained without rounding.} \quad (16)$$

$$\frac{nT_I}{j} = K_1, K_2 \dots K_j \quad (17)$$

$$\frac{nT_O}{j} = K_j + 1, K_j + 2 \dots K_{2j} \quad (18)$$

$$\text{If } 2j \leq n,$$

$$K_n = 0.0 \quad (19)$$

Where the net rate of change, $I + (O+D)$, is negative,

$$j = \frac{n(O+D) - nT_O}{I + (O+D)} \quad (20)$$

$$I + (O+D) + \frac{nT_O}{j} = K_1, K_2, \dots K_j \quad (21)$$

$$\text{If } \sum_{j=1}^j K \leq n(O+D)$$

$$n(O+D) - jK_1 = K_j + 1 \quad (22)$$

$$\frac{nI}{n-j-1} = K_{j+2}, K_{j+3} \dots K_n \quad (23)$$

$$\text{If } \sum_{j=1}^j K = n(O+D)$$

$$\frac{n(I)}{n-j} = K_{j+1}, K_{j+2}, \dots K_n \quad (24)$$

The variances of such distributions are the minimal variances that will permit the estimations of I , T_I and T_O , and $(O+D)$ with a given number of samples. This variance is less affected by the number of samples than it is by the difference between the values of I , T_I and T_O , and $(O+D)$ as can be seen in Table 18. The lowest standard deviations are observed where T_I is low. Where I is increased relative to T_I , the standard deviation is reduced as well but not to the same extent. For example, $I = 2.0$, $T_I = 1.2$ has a ratio of 0.6 as does $I = 1.5$, $T_I = 0.9$, however, the latter has the lower standard deviation. The unavoidable variance related to any series of values of I , T_I and T_O , $O+D$, and n has significance to survey design. The greater the amount of internal translocation due to T_I and T_O the greater the unavoidable variance of the estimation of K . Increasing the number of sampling points has only a minor effect upon the variance although it has a marked effect upon the standard error and 95% confidence limits of the estimates.

The corrected standard deviations with associated standard errors and 95% confidence limits can be calculated using Subroutine FACTOR which will be found in the Appendix. The correction is imposed following the calculation of the standard deviation of K using equation 3, but only with respect to first moment as is true for the other estimations of standard deviations.

The variance is corrected as follows,

$$\left(\frac{s_{K \text{ calc.}}^2 - s_{\text{Min.}}^2}{s_{K \text{ calc.}}^2} \right) s_K^2 = s_{K \text{ corr.}}^2 \quad (25)$$

Where s_K^2 is the variance calculated by equation 3, $s_{\text{min.}}^2$ is the variance of the distribution of K 's with minimal variance, $s_{K \text{ calc.}}^2$ is the variance of the distribution of K 's calculated by equation 3, and $s_{K \text{ corr.}}^2$ is the corrected variance of K . This correction appears to be justified because the variance of interest is that which is related to the variance of a system with particular characteristics as compared to a similar system with minimal unavoidable variance. Table 19 presents a comparison of uncorrected standard deviations from Tables 16 and 17 and the corresponding corrected values. The system estimates for

Table 18. STANDARD DEVIATIONS AND STANDARD ERRORS OF DISTRIBUTIONS OF K WITH MINIMAL VARIANCE FOR GIVEN VALUES OF I, T_I AND T_O , (O+D) AND n.

I	T_I	O+D	Net	n = 5		n = 10		n = 20	
				S.D.	S.E.	S.D.	S.E.	S.D.	S.E.
2.00	1.20	-0.15	1.85	† 2.7524	† 1.2309	† 2.5965	† 0.8211	† 2.5338	† 0.5666
1.75	1.20	-0.15	1.60	† 3.4084	† 1.5243	† 2.7758	† 0.8778	† 2.7107	† 0.6061
1.50	1.20	-0.15	1.35	† 3.3586	† 1.5020	† 3.1663	† 1.0013	† 3.0831	† 0.6894
1.50	1.20	-0.30	1.20	† 3.3719	† 1.5080	† 3.1785	† 1.0051	† 2.8433	† 0.6358
1.50	1.20	-0.60	0.90	† 3.4249	† 1.5317	† 3.2267	† 1.0204	† 2.7077	† 0.6055
1.50	0.90	-0.15	1.35	† 2.0724	† 0.9268	† 1.9558	† 0.6185	† 1.9124	† 0.4276
1.50	0.60	-0.15	1.35	† 1.4335	† 0.6411	† 1.3528	† 0.4278	† 1.3063	† 0.2921

Table 19. COMPARISON OF UNCORRECTED AND CORRECTED STANDARD DEVIATIONS OF SYSTEM ESTIMATES

	16 Sample Set		19 Sample Set		49 Sample Set		57 Sample Set	
	Uncorrected	Corrected	Uncorrected	Corrected	Uncorrected	Corrected	Uncorrected	Corrected
DDT								
Net	± 1.0000	± 0.2751	± 1.0000	± 0.5986	± 1.0000	± 0.3366	± 1.0000	± 0.5379
I	± 0.7556	± 0.2806	± 0.9016	± 0.5397	± 0.6374	± 0.2145	± 0.9204	± 0.4951
O + D	± 0.2444	± 0.0907	± 0.0984	± 0.0589	± 0.3626	± 0.1221	± 0.0796	± 0.0428
T _O	± 0.2591	± 0.0962	± 0.1262	± 0.0755	± 0.3966	± 0.1335	± 0.1513	± 0.0814
T _I	± 0.7409	± 0.2751	± 0.8738	± 0.5231	± 0.6034	± 0.2031	± 0.8487	± 0.4565
O	± 0.1723	± 0.0640	± 0.0644	± 0.0386	± 0.1720	± 0.0579	± 0.0488	± 0.0263
D	± 0.0721	± 0.0268	± 0.0339	± 0.0203	± 0.1906	± 0.0642	± 0.0307	± 0.0165
T _L	± 56.1105	± 20.8365	± 27.2386	± 16.3047	± 82.7111	± 27.8380	± 46.2947	± 24.9713
T _R	± 16.5496	± 6.1457	± 9.3952	± 5.6239	± 43.4823	± 14.6348	± 17.8875	± 9.6215
DDD								
Net	± 1.0000	± 0.3604	± 1.0000	± 0.3860	± 1.0000	± 0.3419	± 1.0000	± 0.3521
I	± 0.7279	± 0.2623	± 0.7233	± 0.2792	± 0.6357	± 0.2174	± 0.6329	± 0.2228
O + D	± 0.2721	± 0.0981	± 0.2767	± 0.1068	± 0.3643	± 0.1246	± 0.3671	± 0.1293
T _O	± 0.2721	± 0.0981	± 0.2767	± 0.1068	± 0.3653	± 0.1249	± 0.3689	± 0.1299
T _I	± 0.7279	± 0.2623	± 0.7233	± 0.2792	± 0.6347	± 0.2170	± 0.6311	± 0.2222
O	± 0.2350	± 0.0847	± 0.2441	± 0.0942	± 0.2823	± 0.0965	± 0.2662	± 0.0937
D	± 0.0371	± 0.0134	± 0.0326	± 0.0126	± 0.0820	± 0.0280	± 0.1010	± 0.0356
T _L	± 107.4660	± 38.7279	± 123.4241	± 47.6463	± 80.3119	± 27.4603	± 88.7883	± 31.2619
T _R	± 14.6462	± 5.2781	± 14.5330	± 5.6103	± 18.0682	± 6.1779	± 24.4216	± 8.5987
DDE								
Net	± 1.0030	± 0.3602	± 1.0024	± 0.4716	± 1.0010	± 0.4379	± 1.0009	± 0.4545
I	± 0.7843	± 0.2817	± 0.7781	± 0.3661	± 0.6628	± 0.2900	± 0.6787	± 0.3082
O + D	± 0.2186	± 0.0785	± 0.2243	± 0.1055	± 0.3382	± 0.1479	± 0.3222	± 0.1463
T _O	± 0.2211	± 0.0794	± 0.2262	± 0.1064	± 0.3466	± 0.1516	± 0.3311	± 0.1504
T _I	± 0.7818	± 0.2808	± 0.7761	± 0.3651	± 0.6544	± 0.2863	± 0.6697	± 0.3041
O	± 0.1899	± 0.0682	± 0.1929	± 0.0907	± 0.2348	± 0.1027	± 0.2204	± 0.1001
D	± 0.0287	± 0.0103	± 0.0314	± 0.0148	± 0.1033	± 0.0452	± 0.1018	± 0.0462
T _L	± 66.4924	± 23.8815	± 66.8543	± 31.4484	± 99.5728	± 43.5593	± 103.4957	± 46.9942
T _R	± 8.7347	± 3.1372	± 9.3629	± 4.4049	± 30.4277	± 13.3109	± 32.6922	± 14.8445

DDT obtained from the four data sets did show some significant differences when compared using these corrected estimates of the standard deviation. The estimates obtained with the 49 and 57 sample sets were significantly different at the .05 level for Net, I, T_O , and T_I . The estimates obtained with the 16 and 57 sample sets were significantly different for Net, I, and T_O , and the estimates of T_O for the 19 and 57 data sets were also significantly different. These differences would appear to be primarily the result of inclusion or exclusion from the system of sites where there are major increases in the concentration of DDT rather than the effect of substitution of a minimal value for the concentration at time one. The estimation of T_O in systems showing a positive Net rate of change are particularly sensitive to significance testing due to their relatively low standard deviations that result from the distribution of variance between T_I and T_O .

If we keep in mind the limitations imposed by the variability of the data, the estimates can be used to gain a picture of the flux of these pollutants in the study area. The area of south Monterey Bay is approximately 280 square kilometers, or 69,190 acres in size. The density of the sediments on a dry weight basis averages 1.32 grams per cm^3 . Table 20 gives the mean of the estimates for system concentrations and rates that were obtained by the two approaches to analysis and the four data sets. Standard deviations, standard errors, 95% confidence limits, and coefficients of variation for these means are included. These latter descriptive statistics refer only to the variation of the estimates and do not include the effect of compartment variability discussed above.

Table 21 uses the mean of the estimates and gives the total amounts of these chlorinated hydrocarbons in the area and the concentration in pounds per acre based upon the mean concentrations at the two times of sampling. These total amounts are estimated as being present in the top 10 cm of sediment, a depth generally sampled with the collecting gear used. Considering that the usual level of application on land is 2 pounds to the acre the total level of these compounds per acre has reached somewhat more than 1/100 of the land applications level.

The estimated annual rates of input, I, as seen in Table 20, average 130% for DDT, 25% for DDD, and 28% for DDE. The corresponding amounts of these materials expected in the next year are indicated in Table 21. Expected loss due to translocation, output, and decay based on the estimated annual rates, O+D, 10% for DDT, 13% for DDD, and 13% for DDE, are also shown. The resulting net effect for the year period following the last sample time in 1973 gives the expected values shown, Table 21. The expected change in the amount of the total chlorinated hydrocarbons derived from DDT amounts to an increase of 182%. The amounts translocated within the system are presented in Table 21 along with a separation of the expected loss into that expected from output and decay. All of the projections, of course, assume that the estimated rates reflecting flux of these materials in the past three years will persist for the next year period.

The K values for the individual compartments can also be used to present a composite view of the translocation of the three compounds within the system and principal points of geographical exit. The stations at their geographical location are connected with arrows

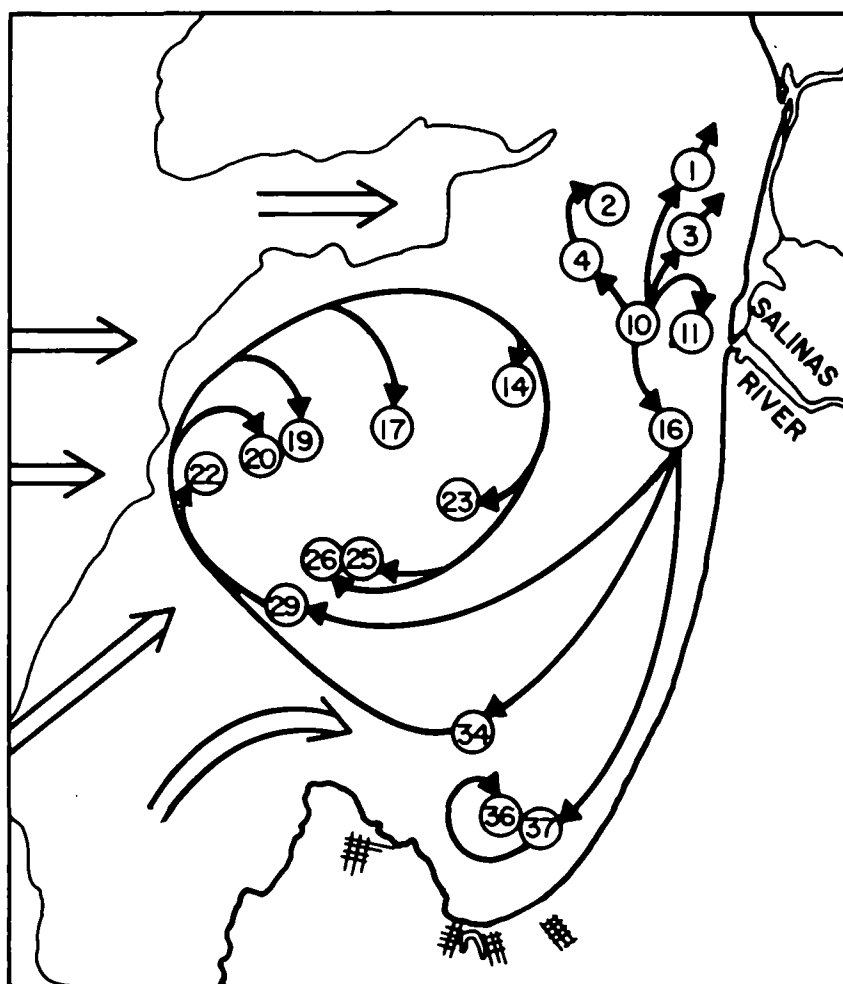
pointing from more negative to less negative K values and ending in basins with positive K values. The result is a kinematic graph representing the movement of these materials within the system. It is composite with respect to the time interval under consideration and would appear to represent the result of several events of translocation. Figure 8 presents such a graph developed for the 19 station data set. The large double arrows indicate the main offshore forces that drive the inshore circulation and correlated with the kinematic expression of circulation within the system.

Table 20. MEAN OF THE ESTIMATES FOR THE SOUTH MONTEREY BAY SYSTEM AND ASSOCIATED DESCRIPTIVE STATISTICS.

			Mean	S.D.	S.E.	95% C.L.	C.V. (%)
C ₁	DDT (ppb)		3.4752	† 0.4281	† 0.2141	† 0.6812	12.3
	DDD (ppb)		2.6144	† 0.3196	† 0.1598	† 0.5086	12.2
	DDE (ppb)		5.7870	† 0.6837	† 0.3419	† 1.0878	11.8
C ₂	DDT (ppb)		15.6411	† 0.4375	† 0.2188	† 0.6961	2.8
	DDD (ppb)		2.4491	† 0.1504	† 0.0752	† 0.2393	6.1
	DDE (ppb)		6.2605	† 0.1207	† 0.0604	† 0.1921	1.9
Net	DDT		+ 1.2015	† 0.8764	† 0.4382	† 1.3944	72.9
	DDD		+ 0.1186	† 0.0327	† 0.0164	† 0.0521	27.6
	DDE		+ 0.1522	† 0.0298	† 0.0149	† 0.0475	19.6
I	DDT		+ 1.2969	† 0.8587	† 0.4294	† 1.3663	66.2
	DDD		+ 0.2533	† 0.0278	† 0.0139	† 0.0442	11.0
	DDE		+ 0.2816	† 0.0096	† 0.0048	† 0.0152	3.4
O + D	DDT		- 0.0955	† 0.0096	† 0.0114	† 0.0364	24.0
	DDD		- 0.1347	† 0.0229	† 0.0042	† 0.0134	6.2
	DDE		- 0.1294	† 0.0084	† 0.0157	† 0.0499	27.3
T _O & T _I	DDT		† 0.8009	† 0.5504	† 0.2752	† 0.8756	68.7
	DDD		† 0.2041	† 0.0071	† 0.0036	† 0.0113	3.5
	DDE		† 0.2190	† 0.0318	† 0.0159	† 0.0505	14.5
O	DDT		- 0.0592	† 0.0212	† 0.0106	† 0.0338	35.8
	DDD		- 0.1096	† 0.0146	† 0.0073	† 0.0233	13.3
	DDE		- 0.1029	† 0.0375	† 0.0187	† 0.0596	36.4
D	DDT		- 0.0362	† 0.0091	† 0.0045	† 0.0145	25.1
	DDD		- 0.0251	† 0.0090	† 0.0143	† 0.0143	35.9
	DDE		- 0.0265	† 0.0061	† 0.0031	† 0.0097	23.0
T _L	DDT (years)		28.9680	† 7.4078	† 3.7039	† 11.7858	25.6
	DDD (years)		44.0829	† 16.0370	† 8.0185	† 25.5148	36.4
	DDE (years)		39.2945	† 9.0835	† 4.5418	† 14.4519	23.1
T _R	DDT (years)		10.9868	† 2.8952	† 1.4476	† 4.6062	26.4
	DDD (years)		7.4454	† 0.4893	† 0.2447	† 0.7785	6.6
	DDE (years)		8.0853	† 1.9717	† 0.9859	† 3.1371	24.4

Table 21. TOTAL AMOUNTS OF DDT, DDD, AND DDE IN THE SOUTH MONTEREY BAY STUDY AREA BASED ON THE MEAN CONCENTRATIONS AT THE TWO SAMPLE TIMES, AND EXPECTED AMOUNTS AFFECTED BY THE MEAN OF THE ESTIMATES OF SYSTEM RATES

		Kilograms	Pounds	Pounds/Acre
Amount at Sample Time 1	DDT	128	284	0.004
	DDD	97	213	0.003
	DDE	214	472	0.007
	TOTAL	439	969	0.014
Amount at Sample Time 2, 3 years later	DDT	579	1276	0.018
	DDD	91	200	0.003
	DDE	232	511	0.007
	TOTAL	932	1987	0.028
Expected input for next year interval	DDT	753	1659	0.024
	DDD	23	50	0.001
	DDE	65	143	0.002
	TOTAL	841	1852	0.027
Expected loss for next year interval	DDT	58	128	0.0018
	DDD	12	26	0.0004
	DDE	30	66	0.0010
	TOTAL	100	220	0.0032
Expected amounts due to Net change for next year interval	DDT	1274	2807	0.041
	DDD	102	224	0.003
	DDE	267	588	0.008
	TOTAL	1643	3619	0.052
Expected amount translocated within the system in next year interval	DDT	463	1020	0.015
	DDD	18	40	0.001
	DDE	51	112	0.002
	TOTAL	532	1172	0.018
Expected amount Output to other systems in next year interval	DDT	35	77	0.0011
	DDD	7	22	0.0003
	DDE	23	51	0.0007
	TOTAL	65	150	0.0021
Expected amount Decayed in next time interval	DDT	21	46	0.0007
	DDD	2	5	0.0001
	DDE	6	14	0.0002
	TOTAL	29	65	0.0010



CIRCULATION OF DDT DERIVATIVES

Figure 8. Composite chart of the translocation of DDT compounds based upon the rates of change, K , at individual stations in the southern portion of Monterey Bay.

DEVELOPMENT OF LABORATORY ASSAY METHODS FOR DETERMINATION OF DECAY RATE

Of the various preparations tested for the assay of decay rate, the sealed hypovial preparations described in the Methods section have best met the following desired criteria.

(1) Preparations must be capable of being sealed to prevent loss of the chlorinated hydrocarbon and its degradation products including CO_2 . (2) The containers must be readily sterilized and of materials that prevent contamination by other chlorinated hydrocarbons. (3) The preparations must be easily manipulated with respect to the establishment of aerobic and anaerobic conditions. (4) The preparation must be susceptible to replication both in terms of individual preparations and aliquots from the same preparation.

The most convenient estimate of decay can be obtained by measurement of the amount of $^{14}\text{CO}_2$ produced from ring labelled substrate after an interval of time. Knowing the initial concentrations of substrate the decay to carbon dioxide can be expressed as a decimal fraction of this initial concentration. The decimal fraction is the D_{CO_2} . Table 22 presents the results of an assay of DDT to CO_2 under aerobic conditions at 10°C . Two aliquots from each of five preparations at four concentrations of DDT were analysed for their $^{14}\text{CO}_2$ content. There is no significant difference between the D_{CO_2} measurements at the four concentrations of DDT. Therefore, over the range from 100 parts per billion to 100 parts per million there was neither a stimulation of the decay process nor a saturation of the decay process by substrate. Table 23 presents the results of assays for D_{CO_2} of DDT, DDD, and DDE. This Table also includes the results of assays in which the effect of environmental variables on the D_{CO_2} was determined.

The Q_{10} for D_{CO_2} of DDT calculated from the aerobic 10° and 20° assays is 2.50. The remaining assays where DDT is the substrate were designed to determine the participation of various physiologically different microbial populations in the decay process. Aerobic conditions without additional nutrients gave the maximum D_{CO_2} . The decay process was inhibited by anaerobiosis, but a rate 27% of the aerobic rate remained. The addition of nitrate as an additional electron acceptor under anaerobic conditions permitted an increase in the anaerobic rate. The three highest concentrations of nitrate, $5 \times 10^{-1}\%$ to $5 \times 10^{-3}\%$ were inhibitory but below these concentrations the anaerobic rate becomes 68% of the aerobic rate at $5 \times 10^{-5}\%$ sodium nitrate.

The addition of a possible cometabolite, sodium acetate, somewhat removes the inhibitory effect of $5 \times 10^{-1}\%$ sodium nitrate probably by its lowering of the nitrate level through denitrification. However, at none of the levels of sodium acetate tested did the anaerobic rate reach the level with $5 \times 10^{-5}\%$ sodium nitrate alone. The effect of the addition of cometabolites on decay in the presence of nitrate reducing systems must be tested at lower concentrations of nitrate.

Sulfate, present in the seawater, was available as an electron acceptor under anaerobic conditions. Attempts to stimulate sulfate reduction systems by the addition of ethanol under anaerobic conditions were successful. However, the anaerobic decay of DDT was not increased over the rate observed with optimum nitrate concentrations and in the absence of added electron donors such as sodium acetate.

Table 22. RESULTS OF A LABORATORY ASSAY OF ANNUAL RATE OF DECAY OF DDT TO CO₂, D_{CO2}, EXPRESSED AS A DECIMAL FRACTION OF THE INITIAL CONCENTRATION OF DDT MAINTAINED AT 10°C. UNDER AEROBIC CONDITIONS.

DDT	Prepar.	D _{CO2}	Means	S.D.	Means	S.D.	Mean	S.D.
100 ppm	1	.0046						
	1	.0045	.00455	± .000071				
	2	.0048						
	2	.0042	.00450	± .000424				
	3	.0059						
	3	.0056	.00575	± .000212				
	4	.0050						
	4	.0045	.00475	± .000354				
	5	.0046						
	5	.0053	.00495	± .000495	.00490	± .000544		
10 ppm	1	.0050						
	1	.0052	.00510	± .000141				
	2	.0058						
	2	.0048	.00530	± .000707				
	3	.0045						
	3	.0056	.00505	± .000778				
	4	.0056						
	4	.0057	.00565	± .000071				
	5	.0051						
	5	.0056	.00535	± .000354	.00529	± .000436		
1 ppm	1	.0050						
	1	.0059	.00545	± .000636				
	2	.0045						
	2	.0046	.00455	± .000071				
	3	.0062						
	3	.0057	.00595	± .000354				
	4	.0058						
	4	.0052	.00550	± .000424				
	5	.0063						
	5	.0058	.00605	± .000354	.00550	± .000638		
100 ppb	1	.0057						
	1	.0057	.00570	± .0000				
	2	.0045						
	2	.0047	.00460	± .000141				
	3	.0051						
	3	.0051	.00510	± .0000				
	4	.0055						
	4	.0058	.00565	± .000212				
	5	.0063						
	5	.0053	.00580	± .000707	.00537	± .000542	.00527	± .000570

Table 23. RESULTS OF LABORATORY ASSAYS OF THE ANNUAL RATES OF DECAY TO CO₂, D_{CO₂}, AND THE EFFECT OF ENVIRONMENTAL VARIABLES ON THE PROCESS.

Conditions	Substrate	D _{CO₂}	Mean	S.D.
Aerobic, 10°C	DDT 100 ppm	.0050	.00529	± .00023
	10 ppm	.0053		
	1 ppm	.0055		
	100 ppb	.0054		
Aerobic, 20°C	DDT 100 ppm	.0100	.01320	± .00335
	10 ppm	.0111		
	1 ppm	.0167		
	100 ppb	.0154		
			Q ₁₀ 2.50	
Anaerobic, 10°C	DDT 100 ppm	.0012	.00145	± .00027
	10 ppm	.0013		
	1 ppm	.0015		
	100 ppb	.0018		
Anaerobic, 10°C 5 x 10 ⁻¹ % NaNO ₃	DDT 10 ppm	.0013	.00150	± .00017
	1 ppm	.0016		
	100 ppb	.0016		
5 x 10 ⁻² % NaNO ₃	DDT 10 ppm	.0017	.00183	± .00015
	1 ppm	.0018		
	100 ppb	.0020		
5 x 10 ⁻³ % NaNO ₃	DDT 10 ppm	.0024	.00250	± .00017
	1 ppm	.0024		
	100 ppb	.0027		
5 x 10 ⁻⁴ % NaNO ₃	DDT 10 ppm	.0030	.00340	± .00035
	1 ppm	.0036		
	100 ppb	.0036		
5 x 10 ⁻⁵ % NaNO ₃	DDT 10 ppm	.0037	.00360	± .00017
	1 ppm	.0034		
	100 ppb	.0037		
5 x 10 ⁻⁶ % NaNO ₃	DDT 10 ppm	.0036	.00310	± .00056
	1 ppm	.0025		
	100 ppb	.0032		
5 x 10 ⁻⁷ % NaNO ₃	DDT 10 ppm	.0031	.00313	± .00006
	1 ppm	.0032		
	100 ppb	.0031		

Table 23 CONTINUED. RESULTS OF LABORATORY ASSAYS OF THE ANNUAL RATES OF DECAY TO CO₂, DCO₂, AND THE EFFECT OF ENVIRONMENTAL VARIABLES ON THE PROCESS.

Conditions	Concentration	DCO ₂	Mean	S.D.
Anaerobic, 10°C, 5 x 10 ⁻¹ % Na NO ₃	5 x 10 ⁻¹ % Na Acetate	DDT 10 ppm	.00090	± .00017
		1 ppm		
		100 ppb		
	5 x 10 ⁻² % Na Acetate	DDT 10 ppm	.00087	± .00012
		1 ppm		
		100 ppb		
	5 x 10 ⁻³ % Na Acetate	DDT 10 ppm	.00223	± .00006
		1 ppm		
		100 ppb		
	5 x 10 ⁻⁴ % Na Acetate	DDT 10 ppm	.00237	± .00015
		1 ppm		
		100 ppb		
	5 x 10 ⁻⁵ % Na Acetate	DDT 10 ppm	.00227	± .00006
		1 ppm		
		100 ppb		
	5 x 10 ⁻⁶ % Na Acetate	DDT 10 ppm	.00213	± .00021
		1 ppm		
		100 ppb		
	5 x 10 ⁻⁷ % Na Acetate	DDT 10 ppm	.00240	± .00000
		1 ppm		
		100 ppb		
Aerobic, 10°C	5 x 10 ⁻¹ % Na Acetate	DDT 10 ppm	.00317	± .00012
		1 ppm		
		100 ppb		
	5 x 10 ⁻² % Na Acetate	DDT 10 ppm	.00307	± .00035
		1 ppm		
		100 ppb		
	5 x 10 ⁻³ % Na Acetate	DDT 10 ppm	.00237	± .00012
		1 ppm		
		100 ppb		

Table 23 CONTINUED. RESULTS OF LABORATORY ASSAYS OF THE ANNUAL RATES OF DECAY TO CO₂, DCO₂, AND THE EFFECT OF ENVIRONMENTAL VARIABLES ON THE PROCESS.

Conditions	Concentration	D _{CO₂}	Mean	S.D.
5 x 10 ⁻⁴ % Na Acetate	DDT 10 ppm	.0028	.00297	± .00015
	1 ppm	.0030		
	100 ppb	.0031		
5 x 10 ⁻⁵ % Na Acetate	DDT 10 ppm	.0027	.00287	± .00015
	1 ppm	.0030		
	100 ppb	.0029		
5 x 10 ⁻⁶ % Na Acetate	DDT 10 ppm	.0025	.00270	± .00020
	1 ppm	.0027		
	100 ppb	.0029		
5 x 10 ⁻⁷ % Na Acetate	DDT 10 ppm	.0028	.00277	± .00025
	1 ppm	.0030		
	100 ppb	.0025		
Anaerobic, 10°C				
5 x 10 ⁻¹ % Ethanol	DDT 10 ppm	.0007	.00043	± .00031
	1 ppm	.0005		
	100 ppb	.0001		
5 x 10 ⁻² % Ethanol	DDT 10 ppm	.0027	.00273	± .00006
	1 ppm	.0028		
	100 ppb	.0027		
5 x 10 ⁻³ % Ethanol	DDT 10 ppm	.0034	.00307	± .00035
	1 ppm	.0031		
	100 ppb	.0027		
5 x 10 ⁻⁴ % Ethanol	DDT 10 ppm	.0029	.00297	± .00006
	1 ppm	.0030		
	100 ppb	.0030		
5 x 10 ⁻⁵ % Ethanol	DDT 10 ppm	.0034	.00320	± .00020
	1 ppm	.0032		
	100 ppb	.0030		
5 x 10 ⁻⁶ % Ethanol	DDT 10 ppm	.0022	.00230	± .00010
	1 ppm	.0023		
	100 ppb	.0024		
5 x 10 ⁻⁷ % Ethanol	DDT 10 ppm	.0023	.00233	± .00015
	1 ppm	.0022		
	100 ppb	.0025		
Aerobic, 10°C	DDD 100 ppm 10 ppm 1 ppm 100 ppb	.0016 .0015 .0015 .0023	.00173	± .00000
Aerobic, 10°C	DDE 100 ppm 10 ppm 1 ppm 100 ppb	.0030 .0028 .0031 .0041	.00325	± .00058

Table 24. RATES OF DECAY TO WATER SOLUBLE COMPOUNDS AND CO₂ DETERMINED BY LABORATORY ASSAYS.

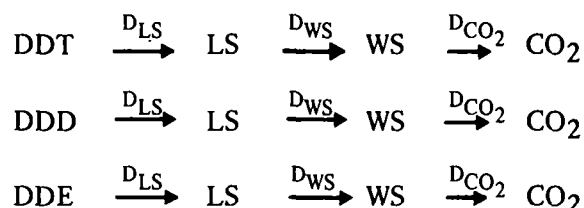
Laboratory Assays	DDT		DDD		DDE	
		S.D.		S.D.		S.D.
D _{CO₂}	.00529	± .00023	.00173	± .00036	.00325	± .00058
D _{WS}	.01539	± .000817	.00309	± .00052	.00459	± .00074
Laboratory Assays Corrected by Q ₁₀						
D _{CO₂}	.00600	± .00026	.00196	± .00041	.00369	± .00066
D _{WS}	.01746	± .00093	.00351	± .00059	.00521	± .00084
Estimations from Field Data						
D	.0362		.0251		.0265	

The addition of sodium acetate as an extra electron donor under aerobic conditions was inhibitory to the aerobic decay process. However, since there was hydrogen sulfate produced in these preparations the inhibition may have been due to the competition for the available oxygen and the production of anaerobic conditions.

In summary, decay to CO_2 appears to be primarily due to the activity of aerobic micro-organisms. The process attains the greatest rate where there is no unusual competition for oxygen. Since the known mechanisms for splitting aromatic rings involve the addition of oxygen to the aromatic nucleus prior to splitting, these observations are not unexpected. However, some considerable activity remains under anaerobic conditions even where an additional oxidizable substrate such as sodium acetate or ethanol is present to remove any traces of residual oxygen. The results also indicate that nitrate and sulfate may be acceptable electron acceptors in the oxidation of aromatic compounds under anaerobic conditions. The mechanisms for anaerobic ring split have not been elucidated. Finally, The Q_{10} for the decay process under aerobic conditions presents no surprise as to its magnitude.

A comparison of the D_{CO_2} for DDT, DDD, and DDE reveals a similar relationship to the total decay rates, D , estimated for South Monterey Bay in that $D_{\text{DDT},\text{CO}_2} > D_{\text{DDE},\text{CO}_2} > D_{\text{DDD},\text{CO}_2}$ just as $D_{\text{DDT}} > D_{\text{DDE}} > D_{\text{DDD}}$. See Table 24.

For purposes of analysis the process of decay can be divided into a series of steps as follows,



where LS represents lipid soluble degradation products of the starting compound and WS represents water soluble degradation products of the starting compound.

Water soluble degradation products were measured as water soluble ^{14}C after high speed centrifugation of samples from the initial preparations followed by acidification to remove $^{14}\text{CO}_2$.

D_{WS} values presented in Table 24 are based on the sum of the ^{14}C present in water soluble form plus that present as $^{14}\text{CO}_2$. Attempts at determining the amount of lipid soluble degradation products were unsuccessful. The high levels of the starting compound still present in the preparations made quantification by gas chromatography difficult. Thin layer chromatography was more successful but revealed that the sodium hydroxide added to stop further biological breakdown and to absorb $^{14}\text{CO}_2$ from the gas phase caused conversion of a considerable amount of the DDT to DDD.

While laboratory assays of decay rate have revealed rates compatible with the field estimation, it has not been possible to use this approach for full appraisal of the method of

estimation of field rates. If we take the difference between the values of D_{WS} obtained from laboratory assays and D obtained from field estimations the rates of decay of the parent compounds to lipid soluble breakdown products, D_{LS} , are .0187 for DDT, .0216 for DDD, and .0213 for DDE under aerobic conditions at 11°C, the mean temperature of the sediments. It should be noted that although every precaution was taken to ensure purity of starting materials in laboratory assays, the amounts of decomposition in three month periods is extremely small and trace contaminants containing label could have a large effect upon the results. In addition it must be emphasized that conditions in laboratory preparations poorly approximate conditions in the field. Therefore, their value is more in terms of results obtained by comparisons between preparations rather than comparisons between laboratory preparation and field observation.

SECTION VI

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APPENDIX A

PROGRAM FOR ESTIMATING SYSTEM RATES BASED ON REAL PAIRED SAMPLE VALUES.

This program for calculation of estimates of rates of input, output, translocation, and decay was written in Fortran IV level G, and was run on an IBM 360/67. In our experience 112k was used and the program required approximately 40 seconds per run. A maximum of 60 stations, 7 chemical compounds, and 2 sample times is permitted with the program as written.

The time interval is calculated in the subroutine, LEAPYR, through use of a calendar table described below. K values are calculated using double precision, and confidence intervals are estimated through use of a table of "t values."

There are eight cards which precede the data deck. Their formats and content are as follows:

First three cards, FORMAT (1X,13F6.3/13F6.3/4F6.3), contain the table of t values.

The following numbers are punched using the indicated format:

First card, 12.706 4.303 3.182 2.776 2.571 2.447 2.365 2.306 2.262 2.228 2.201
2.179 2.160

Second card, 2.145 2.131 2.120 2.110 2.101 2.093 2.086 2.080 2.074 2.069 2.064
2.060 2.056

Third card, 2.052 2.048 2.045 2.042.

Fourth card, FORMAT (12I4), contains numbers for calculation of time intervals.

The following numbers are punched using the indicated format:

0 31 59 90 120 151 181 212 243 273 304 334.

Fifth card, FORMAT (2I5), contains the number of stations followed by the number of chemical compounds in the data set.

Sixth through eighth cards, FORMAT (10A8), contain the names of the chemical compounds entered, left justified, followed by the word TOTAL, followed by the concentration level repeated once for each chemical compound. Any remaining portion of the three cards is left blank. The set of name cards used with the data analyzed in the present case was as follows:

First Card

DDT DDD DDE TOTAL PPB PPB PPB PPB PERCENT PERCENT

Second Card

PERCENT

The third card was left blank.

The data is organized using FORMAT (1X,I2,2(A4,A2),I2,2(1X,I2),7F7.2). The first variable is the station number. The next six fields store the location in terms of latitude and longitude. The next three variables store the month, day, and year, and the remaining fields store the measured concentrations of each chemical compound.

An optional subroutine FACTOR may be called by placing a card before the END card with CALL FACTOR.

```

C      PROGRAM FOR ESTIMATING SYSTEM RATES BASED ON
C      REAL PAIRED SAMPLE VALUES.
C
0001      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8)
1, STE(23,8),CL95(23,8),VAR1(7),VAR2(7),VAR3(7),SUM1(7),SUM2(
27),SUM3(7),SUM4(7),COV1(7),COV2(7)
0002      REAL *4MEAN,MR(7),M(17,8)
0003      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7)
0004      INTEGER CST(2,60),CDATE(2,60,3)
0005      COMMON X, TABLE,IA,I,K,KD,ID
0006      COMMON/BLK1/NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0006      COMMON/BLK2/MR
0007      READ (5,45) TABLE
0008      READ (5,46) MONTH
0009      READ (5,47) IA,ID
C
C      CALCULATE INDEXES.
C      AI          NUMBER OF STATIONS CONVERTED TO A REAL NUMBER
C      IP1          ID + 1
C      IP2          ID + 2
C      I2TP2        2 * ID + 2
C      I2TP3        2 * ID + 3
C      I3TP2        3 * ID + 2
0010      AI=IA
0011      IP1=ID+1
0012      IP2=ID+2
0013      I2TP2=2*ID+2
0014      I2TP3=2*ID+3
0015      I3TP2=3*ID+2
C
C      CLEAR X ARRAY.
C
0016      DO 1 I=1,10
C
0017      DO 1 J=1,IA
C
0018      DO 1 K=1,IP1
0019      1 X(I,J,K)=0.0
C
0020      WRITE (6,50)
C
C      READ IN DERIVATIVE NAMES AND CONCENTRATION LEVEL ON UP TO 3 CAR
0021      READ (5,48) NAME
C
C      READ IN DATA.
C

```

```

0022      DO 2 I=1,2
      C
0023      DO 2 J=1,IA
0024      2 READ (5,49) CST(I,J),(ALOC(I,J,L),L=1,6),(CDATE(I,J,L),L=1,3),(X(I
      1,J,K),K=1,ID)
      C
      C
      C      COMPUTE TOTAL OF EACH STATION.
      C
0025      DO 3 I=1,2
      C
0026      DO 3 J=1,IA
      C
0027      DO 3 L=1,ID
0028      3 X(I,J,IP1)=X(I,J,L)+X(I,J,IP1)
      C
      C
      C      WRITE HEADING OF FIRST TWO PAGES.
      C
0029      DO 5 I=1,2
0030      L=I
0031      WRITE (6,51) I
0032      WRITE (6,53) (NAME(N),N=1,IP1)
0033      WRITE (6,52) (NAME(N),N=IP2,I2TP2)
0034      WRITE (6,54)
      C
0035      DO 4 K=1,IP1
0036      CALL STDEV (TOTAL,MEAN,SD,SE,CL)
0037      TOT(I,K)=TOTAL
0038      M(I,K)=MEAN
0039      STD(I,K)=SD
0040      STE(I,K)=SE
0041      4 CL95(I,K)=CL
      C
      C      L1=NUMBER OF SETS COMPUTED.
      C
      C      WRITE FIRST TWO PAGES.
0042      L1=IP1
0043      CALL PRINT
0044      WRITE (6,53) (NAME(N),N=1,IP1)
0045      WRITE (6,52) (NAME(N),N=IP2,I2TP2)
0046      WRITE (6,54)
0047      CALL PRINT2
0048      5 CONTINUE
      C
      C
      C      COMPUTE PERCENTS.
      C

```

```

0049      DO 8 I=3,4
0050      L1=ID
0051      L=I-2
0052      WRITE (6,51) L
0053      WRITE (6,53) (NAME(N),N=1,ID)
0054      WRITE (6,52) (NAME(N),N=I2TP3,I3TP2)
0055      WRITE (6,54)
      C
0056      DO 6 K=1,ID
      C
0057      DO 6 J=1,IA
0058      6 X(I,J,K)=X(L,J,K)/X(L,J,IP1)*100.
      C
      C
0059      DO 7 K=1,3
0060      CALL STDEV (TOTAL,MEAN,SD,SE,CL)
0061      TOT(I,K)=TOTAL
0062      M(I,K)=MEAN
0063      STD(I,K)=SD
0064      STE(I,K)=SE
0065      7 CL95(I,K)=CL
      C
0066      CALL PRINT
0067      WRITE (6,53) (NAME(N),N=1,ID)
0068      WRITE (6,52) (NAME(N),N=I2TP3,I3TP2)
0069      WRITE (6,54)
0070      CALL PRINT2
0071      8 CONTINUE
      C
      C
0072      DO 10 J=1,IA
      C
0073      DO 10 L=1,IA
0074      IF (CST(1,J).EQ.CST(2,L)) GO TO 9
0075      GO TO 10
0076      9 CALL LEAPYR (J)
      C
0077      DO 10 K=1,ID
0078      X(5,J,K)=YR
0079      10 CONTINUE
      C
      C
      C      CALCULATE TOTAL AND MEAN OF N.
      C

```

```

0080      DO 12 K=1,ID
0081      TOT(5,K)=0.
      C
0082      DO 11 J=1,IA
0083      11 TOT(5,K)=TOT(5,K)+X(5,J,K)
      C
0084      12 M(5,K)=TOT(5,K)/AI
      C
      C
0085      DO 14 K=1,ID
0086      V=0.0
      C
0087      DO 13 J=1,IA
0088      13 V=(M(5,K)-X(5,J,K))**2+V
      C
0089      STD(5,K)=SQRT(V/(AI-1.0))
0090      14 CALL STDEV2 (STD(5,K),STE(5,K),CL95(5,K))
      C
      C
      C      CALCULATE K VALUES.
      C
0091      DO 15 K=1,ID
0092      SUM1(K)=0.0
      C
0093      DO 15 J=1,IA
0094      IF (X(1,J,K).EQ.0) X(1,J,K)=.004
0095      IF (X(2,J,K).EQ.0) X(2,J,K)=.004
0096      V=(DLOG10(X(2,J,K))-DLOG10(X(1,J,K)))/(X(5,J,K))
0097      V2(J,K)=10.**V-1.0
0098      15 SUM1(K)=SUM1(K)+V2(J,K)
      C
      C
      C      SORT K VALUES.
      C
0099      DO 17 K=1,ID
      C
0100      DO 17 J=1,IA
0101      IF (V2(J,K).GT.0) GO TO 16
0102      X(7,J,K)=V2(J,K)
0103      GO TO 17
0104      16 X(6,J,K)+V2(J,K)
0105      17 X(8,J,K)=X(7,J,K)+X(6,J,K)
      C
      C
      C      CALCULATE K-NET.
      C

```



```

0106      DO 19 K=1,ID
          C
0107      DO 19 J=1,IA
0108      V=X(8,J,K)-SUM1(K)/AI
0109      IF (V.GT.0) GO TO 18
0110      X(10,J,K)=V
0111      GO TO 19
0112      18 X(9,J,K)=V
0113      19 CONTINUE
          C
          C
          C
          C      COMPUTE SUM AND MEAN FOR K VALUES.
          C
          C
0114      DO 21 K=1,ID
          C
0115      DO 21 I=6,10
0116      V=0.0
          C
          C
0117      DO 20 J=1,IA
0118      20 V=V+X(I,J,K)
          C
0119      TOT(I,K)=V
0120      21 M(I,K)=V/AI
          C
          C
          C      CALCUALTE STANDARD DEVIATION, STANDARD ERROR, AND 95% CONFIDE
          C      LIMITS OF K VALUES.
          C
0121      DO 22 K=1,ID
0122      SUM 1 (K)=0.0
0123      SUM 2 (K)=0.0
0124      SUM 3 (K)=0.0
0125      22 SUM 4 (K)=0.0
0127      DO 23 J=1,IA
0128      V2(J,K)=DLOG(X(2,J,K))-DLOG(X(1,J,K))
0129      SUM2(K)=V2(J,K)+SUM2(K)
0130      SUM 3 (K)=(DLOG(X(1,J,K))-ALOG(M(1,K)))*2+SUM3(K)

```

```

0131      23 SUM 4 (K)=(DLOG(X(2,J,K))-ALOG(M(2,K)))**2+SUM4(K)
      C
      C
0132      DO 24 K=1,ID
0133      VAR 1(K)=(.43429/M(1,K))**2*SUM3(K)/(AI-1.0)+(-.43429/M(2,K))**2
      1*SUM4 (K)/(AI-1.0)
0134      24 V1(K)=SUM2(K)/AI
0135      DO 25 K=1,ID
0136      VAR2(K)=((1.0/M(5,K))**2*VAR1(K))+(-V1(K)/(M(5,K)**2))**2*STD(5,K)
      1**2
0137      VAR2(K)=10.0**VAR2(K)
0138      STD(8,K)=SQRT(VAR2(K))
0139      25 CALL STDEV2 (STD(8,K),STE(8,K),CL95(8,K))
      C
      C
      C
      C      CALCULATE THE DISTRIBUTION OF VARIANCE BETWEEN +K AND -K
      C
0142      DO 30 K=1,ID
0143      V=0.0
      C
0144      DO 27 J=1,IA
0145      IF (X(6,J,K)) 27,27,26
0146      26 V=(X(6,J,K)-M(8,K))**2+V
0147      27 CONTINUE
      C
      C
0148      V=V/(AI-1.0)
0149      W=0.0
      C
      C
0150      DO 29 J=1,IA
0151      IF (X(7,J,K)) 28,29,29
0152      28 W=(X(7,J,K)-M(8,K))**2+W
0153      29 CONTINUE
      C
0154      W=W/(AI-1.0)
0155      U=V+W
0156      V=STD(8,K)**2*(V/U)**2

```

```

0157         STD(6,K)=SQRT(V)
0158         W=STD(8,K)**2*(W/U)**2
0159         STD(7,K)=SQRT(W)
0160         CALL STDEV2(STD(6,K),STE(6,K),CL95(6,K))
0160     30 CALL STDEV2 (STD(7,K),STE(7,K),CL95(7,K))
        C
        C
        C     CALCULATION OF STANDARD DEVIATION K-NET AND ITS DISTRIBUTION.
        C
        C
        C
0161         DO 35 K=1,ID
0161         V=0.0
0162         W=0.0
0163         DO 34 J=1,IA
0164             IF(X(9,J,K)) 32,32,31
0165         31 V=V+(X(9,J,K)**2)
0166         32 IF(X(10,J,K)) 33,34,34
0167         33 W=W+(X(10,J,K)**2)
0168         34 CONTINUE
        C
        C
0169         V=V/(AI-1.0)
0170         W=W/(AI-1.0)
0171         STD(9,K)=SQRT((V/(V+W))**2*(STD(8,K)**2))
0172         CALL STDEV2(STD(9,K),STE(9,K),CL95(9,K))
0173         STD(10,K)=SQRT((W/(V+W))**2*(STD(8,K)**2))
0174     35 CALL STDEV2(STD(10,K),STE(10,K),CL95(10,K))
        C
        C
        C
0175         CALL PRINT3
        C
        C     CALCULATE 0 AND ITS STANDARD DEVIATION
        C
0176         DO 41 K=1,ID
0177         M(11,K)=(M(9,K)/M(6,K))*M(7,K)
0178         STD(11,K)=SQRT(STD(7,K)**2*((M(9,K)/M(6,K))**2))
0179         CALL STDEV2(STD(11,K),STE(11,K),CL95(11,K))
        C
        C
        C
        C
        C     CALCULATION OF D
        C

```

```

C
C
0192      M(12,K)=M(7,K)-M(11,K)
C
C
C      CALCULATION OF STANDARD DEVIATION OF D
C
0193      STD(12,K)=SQRT(STD(7,K)**2*(1.-M(9,K)/M(6,K))**2)
0194      CALL STDEV2 (STD(12,K),STE(12,K),CL95(12,K))
C
C
C      CALCULATE TL.
C
C
0196      M(13,K)=-1.0*(1.0/M(12,K))
0197      STD(13,K)=SQRT(STD(12,K)**2*(1.0/M(12,K)**2)**2)
0198      41 CALL STDEV2 (STD(13,K),STE(13,K),CL95(13,K))
C
C
C      CALCULATE TR.
C
0199      DO 42 K=1,ID
0200      M(14,K)=-1.0*(1.0/M(7,K))
0201      STD(14,K)=SQRT(STD(7,K)**2*(1.0/M(7,K)**2)**2)
0202      42 CALL STDEV2 (STD(14,K),STE(14,K),CL95(14,K))
C
C
0203      DO 44 K=1,ID
0204      WRITE (6,55) NAME (K)
0205      WRITE (6,56) NAME (K),M(8,K),STD(8,K),STE(8,K),CL95(8,K)
0206      WRITE (6,57) NAME (K),M(9,K),STD(9,K),STE(9,K),CL95(9,K)
0207      WRITE (6,58)
0208      WRITE (6,59) NAME (K),M(10,K),STD(10,K),STE(10,K),CL95(10,K)
0209      WRITE (6,60)
0210      WRITE (6,61) NAME (K),M(6,K),STD(6,K),STE(6,K),CL95(6,K)
0211      WRITE (6,62) NAME (K),M(7,K),STD(7,K),STE(7,K),CL95(7,K)
0212      WRITE (6,63) NAME (K),M(11,K),STD(11,K),STE(11,K),CL95(11,K)
0213      WRITE (6,64) NAME (K),M(12,K),STD(12,K),STE(12,K),CL95(12,K)
0214      WRITE (6,65) NAME (K),M(13,K),STD(13,K),STE(13,K),CL95(13,K)
0215      WRITE (6,66)
0216      WRITE (6,65) NAME (K),M(14,K),STD(14,K),STE(14,K),CL95(14,K)
0217      WRITE (6,67)
C

```

```

0218      DO 43 L=1,3
0219      43 WRITE (6,54)
      C
0220      WRITE (6,68)
0221      WRITE (6,69) M(5,K)
0222      WRITE (6,70) NAME(K),M(2,K),M(1,K),M(6,K),M(9,K),M(10,K),M(11,K),M
1(12,K)
0223      44 CONTINUE
      C
0224      CALL FACTOR
0225      STOP
      C
0226      45 FORMAT (1X,13F6.3/13F6.3/4F6.3)
0227      46 FORMAT (12I4)
0228      47 FORMAT (2I5)
0229      48 FORMAT (10A8)
0230      49 FORMAT (1X,I2,2(2A4,A2),I2,2(1X,I2),7F7.2)
0231      50 FORMAT('1')
0232      51 FORMAT('1','C'/2X,I1,/3X,'STATION',3X,'LATITUDE',3X,'LONGITUDE',
-5X,'DATE')
0233      52 FORMAT(48X,8(3X,A8))
0234      53 FORMAT('+',47X,8(3X,A8))
0235      54 FORMAT(/)
0236      55 FORMAT('1',1X,'RATES OF CHANGE FOR ',A8,30X,'S.D.',7X,'S.E.',4X,
-'95% LIMIT'//)
0237      56 FORMAT(2X,'MEAN OF K',13X,'= NET',3X,A8,'=',3X,4F11.4/)
0238      57 FORMAT(2X,'MEAN OF + ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0239      58 FORMAT(27X,'I'/)
0240      59 FORMAT(2X,'MEAN OF - ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0241      60 FORMAT(27X,'O'/)
0242      61 FORMAT(2X,'MEAN OF + K',11X,'= I',5X,A8,'=',3X,4F11.4//)
0243      62 FORMAT(2X,'MEAN OF - K',11X,'= O + D',1X,A8,'=',3X,4F11.4//)
0244      63 FORMAT(26X,'O',5X,A8,'=',3X,4F11.4/)
0245      64 FORMAT(26X,'D',5X,A8,'=',3X,4F11.4/)
0246      65 FORMAT(26X,'T',5X,A8,'=',3X,4F11.4)
0247      66 FORMAT(27X,'L'/)
0248      67 FORMAT(27X,'R')
0249      68 FORMAT(13X,'MEAN C',6X,'MEAN C',16X,'I',10X,'T',6X,'-',4X,'T',6X,
-','- ',5X,'O',5X,'-',5X,'D',9X,'N'/19X,'2',11X,'1,27X,'I',11X,'O'/)
0250      69 FORMAT(/97X,F11.4)
0251      70 FORMAT(2X,A8,F10.4,'=',F10.4,' ( 1.0 +',F10.4,' +',F10.4,3(F12.4)
-,(') ')
0252      END
      C
      C

```

```

0001      SUBROUTINE PRINT
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8)
1, STE(23,8),CL95(23,8)
0003      REAL *4MEAN,M(17,8)
0004      REAL *8X(10,60,7),NAME(23)
0005      INTEGER CST(2,60),CDATE(2,60,3)
0006      COMMON X,TABLE,IA,I,K,KD,ID
0007      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
C
0008      DO 1 J=1,IA
0009      1 WRITE (6,3) CST(L,J),(ALOC(L,J,K),K=1,6),(CDATE(L,J,K),K=1,3),(X(1,
1,J,K),K=1,L1)
C
C      SKIP TO BOTTOM OF PAGE
0010      N=(68-(IA+6))/2
C
0011      DO 2 J=1,N
0012      2 WRITE (6,4)
C
0013      RETURN
C
0014      3 FORMAT (5X,I2,5X,2A4,A2,2X,2A4,A2,2X,I2,2('-',I2),8F11.2)
0015      4 FORMAT (/)
0016      END
C
C
0001      SUBROUTINE PRINT2
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8)
1, STE(23,8),CL95(23,8)
0003      REAL *4MEAN,M(17,8)
0004      REAL *8X(10,60,7),NAME(23)
0005      INTEGER CST(2,60),CDATE(2,60,3)
0006      COMMON X,TABLE,IA,I,K,KD,ID
0007      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0008      WRITE (6,1) (TOT(I,J),J=1,L1)
0009      WRITE (6,2) (M(I,J),J=1,L1)
0010      WRITE (6,3) (STD(I,J),J=1,L1)
0011      WRITE (6,4) (STE(I,J),J=1,L1)
0012      WRITE (6,5) (CL95(I,J),J=1,L1)
0013      RETURN
C
0014      1 FORMAT (34X,'TOTALS',6X,7F10.4)
0015      2 FORMAT (/34X,'MEAN',8X,7F10.4)
0016      3 FORMAT (/34X,'S.D.',8X,7F10.4)
0017      4 FORMAT (/34X,'S.E.',8X,7F10.4)
0018      5 FORMAT (/34X,'95% CL',6X,7F10.4)
0019      END
C
C

```

```

0001      SUBROUTINE PRINT3
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8)
          1, STE(23,8),CL95(23,8)
0003      REAL *8X(10,60,7)
0004      REAL *8NAME(23)
0005      REAL *4MEAN,M(17,8)
0006      INTEGER CST(2,60),CDATE(2,60,3)
0007      COMMON X,TABLE,IA,I,K,KD,ID
0008      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
C
0009      DO 2 K=1,ID
0010      WRITE (6,3)
0011      WRITE (6,4)
0012      WRITE (6,5) NAME(K),NAME(K)
0013      WRITE (6,6)
0014      WRITE (6,8)
C
0015      DO 1 J=1,IA
0016      1 WRITE (6,7) CST(1,J),X(2,J,K),X(1,J,K),X(5,J,K),(X(IX,J,K),IX=6,10
          1)
C
0017      WRITE (6,8)
0018      WRITE (6,17) TOT(2,K),TOT(1,K),TOT(5,1),(TOT(L,K),L=6,10)
0019      WRITE (6,16)
0020      WRITE (6,14) NAME(K)
0021      WRITE (6,17) M(2,K),M(1,K),M(5,1),(M(L,K),L=6,10)
0022      WRITE (6,9)
0023      WRITE (6,14) NAME(K)
0024      WRITE (6,17) STD(2,K),STD(1,K),STD(5,1),(STD(L,K),L=6,10)
0025      WRITE (6,10)
0026      WRITE (6,13) NAME(K)
0027      WRITE (6,17) STE(2,K),STE(1,K),STE(5,1),(STE(L,K),L=6,10)
0028      WRITE (6,11)
0029      WRITE (6,13) NAME(K)
0030      WRITE (6,17) CL95(2,K),CL95(1,K),CL95(5,1),(CL95(L,K),L=6,10)
0031      WRITE (6,12)
0032      WRITE (6,15) NAME(K)
0033      2 CONTINUE
C
0034      RETURN
C
0035      3 FORMAT ('1',1X,'STATION')
0036      4 FORMAT (12X,'C',9X,'C',11X,'N',8X,'+ K',7X,'- K',6X,'+K - K',
          1 4X,'+K - NET',3X,'-K - NET')
0037      5 FORMAT ('+',15X,A8,2X,A8)
0038      6 FORMAT (13X,'2',9X,'1',52X,'R',10X,'R'/)
0039      7 FORMAT (4X,I2,1X,2F10.2,2X,3F10.4,3F11.4)
0040      8 FORMAT (/)

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```

0041      9 FORMAT ('+',94X,'MEANS')
0042     10 FORMAT ('+',94X,'S.D.')
0043     11 FORMAT ('+',94X,'S.E.')
0044     12 FORMAT ('+',94X,'95% CONFIDENCE LIMITS')
0045     13 FORMAT ('+', 99X,A8)
0046     14 FORMAT ('+',102X,A8)
0047     15 FORMAT ('+',116X,A8)
0048     16 FORMAT ('+',94X,'TOTALS')
0049     17  FORMAT(/9X,5F10.4,3F11.4)
0050      END
      C
      C
0001      SUBROUTINE LEAPYR (J)
0002      DIMENSION TABLE(30), MONTH(12), ALOC(2,60,6), TOT(10,8), STD(23,8)
0003      1, STE(23,8),CL95(23,8)
0004      REAL *4MEAN,M(17,8)
0005      REAL *8X(10,60,7)
0006      REAL *8NAME(23)
0007      INTEGER TOT,YR1,YR2,DA1,DA2,DAYS
0008      INTEGER CST(2,60),CDATE(2,60,3)
0009      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0010      COMMON X,TABLE,IA,I,K,KD,ID
0011      DAYS=0
0012      NT=0
0013      MO1=CDATE(1,J,1)
0014      DA1=CDATE(1,J,2)
0015      YR1=CDATE(1,J,3)
0016      DA2=CDATE(2,L,2)
0017      YR2=CDATE(2,L,3)
0018      MO2=CDATE(2,L,1)
0019      AMO=MO1
      C
0019      DO 4 I=YR1,YR2
0020      A=I
0021      LEAP=0
0022      IZ=A/4.
0023      Z=IZ
0024      Z=Z*4.
0025      IF (I.EQ.YR1) GO TO 1
0026      GO TO 2
0027      1 DAYS=365-(MONTH(MO1)+DA1)
0028      IF (Z.EQ.A.AND.AMO.LT.3.) LEAP=1
0029      GO TO 3
0030      2 IF (Z.EQ.A) LEAP=1
0031      3 NT=DAYS+LEAP+NT
0032      4 DAYS=365
      C

```



```

0033      IF (LEAP.EQ.1) GO TO 5
0034      GO TO 6
0035      5 IF (MO2.LT.3) NT=NT-1
0036      6 YR=NT-365+MONTH(MO2)+DA2
0037      YR=YR/365.
0038      RETURN
0039      END

```

C

C

```

0001      SUBROUTINE TDIST (T)
0002      REAL *8X(10,60,7)
0003      DIMENSION TABLE(30)
0004      COMMON X,TABLE,IA,I,K,KD,ID
0005      I1=IA-1
0006      AI=I1
0007      IF (I1) 1,1,2
0008      1 WRITE (6,I1) I
0009      GO TO 10
0010      2 IF (I1.LT.31) GO TO 9
0011      IF (I1.LT.41) GO TO 3
0012      GO TO 4
0013      3 TINT=((2.042-2.021)/10.)*(AI-30.)
0014      T=TINT+2.042
0015      GO TO 10
0016      4 IF (I1.LT.61) GO TO 5
0017      GO TO 6
0018      5 TINT=((2.021-2.000)/20.)*(AI-40.)
0019      T=TINT+2.021
0020      GO TO 10
0021      6 IF (I1.LT.121) GO TO 7
0022      GO TO 8
0023      7 TINT=((2.000-1.980)/40.)*(AI-60.)
0024      T=TINT+2.000
0025      GO TO 10
0026      8 T=1.960
0027      GO TO 10
0028      9 T=TABLE(I1)
0029      10 RETURN

```

C

```

0030      11 FORMAT ('1','1 IN T TABLE =',I3)
0031      END

```

C

C

```

0001      SUBROUTINE STDEV (SUMX,XBAR,STD,STE,CL$)
0002      REAL *8X(10,60,7)
0003      DIMENSION TABLE(30)
0004      COMMON X,TABLE,IA,I,K,KD,ID
0005      DEV=0.
0006      SUMX=0.

      C
0007      DO 1 J=1,IA
0008      1 SUMX=SUMX+X(I,J,K)

      C
0009      AI=IA
0010      XBAR=SUMX/AI

      C
0011      DO 2 J=1,IA
0012      DEV=(XBAR-X(I,J,K))**2+DEV
0013      2 CONTINUE

      C
0014      STD=SQRT(DEV/(AI-1.))
0015      STE=STD/SQRT(AI)
0016      CALL TDIST (T)
0017      CL$=T*STE
0018      END

      C
      C
0001      SUBROUTINE STDEV2 (STD,STE,CL$)
0002      REAL *8X(10,60,7)
0003      DIMENSION TABLE(30)
0004      COMMON X,TABLE,IA,I,K,KD,ID
0005      AI=IA
0006      STE=STD/SQRT(AI)
0007      CALL TDIST (T)
0008      CL$=T*STE
0009      RETURN
0010      END

```

```

C      SUBROUTINE FOR PROGRAM FOR ESTIMATING SYSTEM RATES
C      BASED ON REAL PAIRED SAMPLE VALUES.
      SUBROUTINE FACTOR
0001      DIMENSION TABLE (30), MONTH(12), ALOC(2,60,6), TOT(10,8), STD(23,8)
      1,STE(23,8), CL95(23,8), VAR1(7), VAR2(7), VAR3(7), SUM1(7), SUM2(
      27), SUM3(7), SUM4(7), COV1(7), COV2(7)
0002      REAL *4MEAN,M(17,8),MR(7)
0003      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7)
0004      INTEGER CST(2,60),CDATE(2,60,3)
0005      COMMON X,TABLE,IA,I,K,KD,ID
0006      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0007      COMMON /BLK2/ MR

C
C
C      CALCULATE CORRECTION FACTOR FOR STANDARD DEVIATION
C
0008      AI=IA
C
0009      DO 14 K=1,ID
C
0010          IF (M(8,K)) 1,4,5
0011      1 JX=(AI*M(7,K)-AI*M(10,K))/M(8,K)
0012          VJ=JX
0013          V=((AI*M(10,K))/VJ)**2)*VJ
0014          IF ((M(8,K)+((AI*M(10,K))/VJ))*VJ-(AI*M(7,K))) 3,3,2
0015      2 V=V+(AI*M(7,K)-VJ*(M(8,K)+((AI*M(10,K))/VJ))-M(8,K))**2
0016          V=V+(((AI*M(6,K))/(AI-VJ-1.0))-M(8,K))**2*(AI-VJ-1.0)
0017          GO TO 8
0018      3 V=V+(((AI*M(6,K))/(AI-VJ))-M(8,K))**2*(AI-VJ)
0019          GO TO 8
0020      4 JX=AI/2.0
0021          VJ=JX
0022          V=((AI*M(6,K)/VJ)**2)*VJ
0023          V=V+((AI*M(7,K)/VJ)**2)*VJ
0024          GO TO 8
0025      5 JX=(AI*M(6,K)-AI*M(9,K))/M(8,K)
0026          VJ=JX
0027          V=((AI*M(9,K))/VJ)**2)*VJ
0028          IF ((M(8,K)+((AI*M(9,K))/VJ))*VJ-(AI*M(6,K))) 6,7,7
0029      6 V=V+(AI*M(6,K)-VJ*(M(8,K)+((AI*M(9,K))/VJ))-M(8,K))**2
0030          V=V+(((AI*M(7,K))/(AI-VJ-1.0))-M(8,K))**2*(AI-VJ-1.0)
0031          GO TO 8
0032      7 V=V+(((AI*M(7,K))/(AI-VJ))-M(8,K))**2*(AI-VJ)
0033      8 V=V/(AI-1.0)
0034          W=0.0
C
C

```

```

0035      DO 9 J=1,IA
0036      9 W=W+(X(8,J,K)-M(8,K))**2
      C
      C
0037      W=W/(AI-1.0)
0038      C=((W-V)/W)**2
      C
      C      CALCULATE CORRECTED STD,6,7,AND 8
      C      STD(15,K) IS CORRECTED STD(6,K)
      C
0039      STD(15,K)=SQRT(C*STD(6,K)**2)
0040      CALL STDEV2 (STD(15,K),STE(15,K),CL95(15,K))
      C
      C      STD(16,K)IS CORRECTED STE(7,K)
      C
0041      STD(16,K)=SQRT(C*STD(7,K)**2)
0042      CALL STDEV2 (STD(16,K),STE(16,K),CL95(16,K))
      C
      C      STD(17,K)IS CORRECTED STD(8,K)
      C
0043      STD(17,K)=SQRT(C*STD(8,K)**2)
0044      CALL STDEV2 (STD(17,K),STE(17,K),CL95(17,K))
      C
      C      CALCULATE CORRECTED STD(9,K) AND STD(10,K)
      C
0045      V=0.0
      C
      C
0046      DO 11 J=1,IA
      C
0047      IF (X(9,J,K)) 11,11,10
0048      10 V=V+X(9,J,K)**2
0049      11 CONTINUE
      C
      C
      C
0050      V=V/(AI-1.0)
0051      W=0.0
      C
      C
0052      DO 13 J=1,IA
      C
0053      IF (X(10,J,K)) 12,13,13
0054      12 W=W+(X(10,J,K))**2
0055      13 CONTINUE
      C
      C
      C

```

```

0056      W=W/(AI-1.0)
      C
      C      STD(18,K) IS CORRECTED STD(9,K)
      C
0057      STD(18,K)=SQRT((V/(V+W))**2*C*(STD(8,K)**2))
0058      CALL STDEV2 (STD(18,K),STE(18,K),CL95(18,K))
      C
      C
      C      STD(19,K) IS CORRECTED STD(10,K)
      C
0059      STD(19,K)=SQRT((W/(V+W))**2*C*(STD(8,K)**2))
0060      CALL STDEV2 (STD(19,K),STE(19,K),CL95(19,K))
      C
      C      CALCULATE CORRECTED STD(20,K) CORRECTED STD(11,K)
      C
0061      STD(20,K)=SQRT(STD(16,K)**2*(M(9,K)/M(6,K))**2)
      C
      C
0062      CALL STDEV2 (STD(20,K),STE(20,K),CL95(20,K))
      C      CALCULATE STD(21,K) CORRECTED STD(12,K)
0063      STD(21,K)=SQRT(STD(16,K)**2*(1.0-M(9,K)/M(6,K))**2)
0064      CALL STDEV2 (STD(21,K),STE(21,K),CL95(21,K))
      C
      C      CALCULATE STD(22,K) CORRECTED STD(13,K)
      C
0065      STD(22,K)=SQRT(STD(21,K)**2*(1.0/M(12,K)**2)**2)
0066      CALL STDEV2 (STD(22,K),STE(22,K),CL95(22,K))
      C
      C      CALCULATE STD(23,K) CORRECTED STD(14,K)
      C
0067      STD(23,K)=SQRT(STD(16,K)**2*(1.0/M(7,K)**2)**2)
0068      CALL STDEV2 (STD(23,K),STE(23,K),CL95(23,K))
      C
      C
0069      14 CONTINUE
      C
      C
0070      DO 15 K=1,ID
      C
0071      WRITE (6,16) NAME(K)
0072      WRITE (6,17) NAME(K),M(8,K),STD(17,K),STE(17,K),CL95(17,K)
0073      WRITE (6,18) NAME(K),M(9,K),STD(18,K),STE(18,K),CL95(18,K)
0074      WRITE (6,19)
0075      WRITE (6,20) NAME(K),M(10,K),STD(19,K),STE(19,K),CL95(19,K)

```

```

0076      WRITE (6,21)
0077      WRITE (6,22) NAME(K),M(6,K),STD(15,K),STE(15,K),CL95(15,K)
0078      WRITE (6,23) NAME(K),M(7,K),STD(16,K),STE(16,K),CL95(16,K)
0079      WRITE (6,24) NAME(K),M(11,K),STD(20,K),STE(20,K),CL95(20,K)
0080      WRITE (6,25) NAME(K),M(12,K),STD(21,K),STE(21,K),CL95(21,K)
0081      WRITE (6,26) NAME(K),M(13,K),STD(22,K),STE(22,K),CL95(22,K)
0082      WRITE (6,27)
0083      WRITE (6,26) NAME(K),M(14,K),STD(23,K),STE(23,K),CL95(23,K)
0084      WRITE (6,28)
0085      15 CONTINUE
      C
      C
0086      16 FORMAT ('1',40X,'CORRECTED STANDARD DEVIATIONS',//,2X,'RATES OF CHA
      INGE FOR ',A8,30X,'S.D.',7X,'S.E.',4X,'95% LIMIT'//)
0087      17 FORMAT (2X,'MEAN OF K',13X,'= NET',3X,A8,'=',3X,4F11.4/)
0088      18 FORMAT (2X,'MEAN OF + ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0089      19 FORMAT (27X,'I'/)
0090      20 FORMAT (2X,'MEAN OF - ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0091      21 FORMAT (27X,'O'/)
0092      22 FORMAT (2X,'MEAN OF + K',11X,'= I',5X,A8,'=',3X,4F11.4//)
0093      23 FORMAT (2X,'MEAN OF - K',11X,'= O + D',1X,A8,'=',3X,4F11.4//)
0094      24 FORMAT (26X,'O',5X,A8,'=',3X,4F11.4/)
0095      25 FORMAT (26X,'D',5X,A8,'=',3X,4F11.4/)
0096      26 FORMAT (26X,'T',5X,A8,'=',3X,4F11.4)
0097      27 FORMAT (27X,'L'/)
0098      28 FORMAT (27X,'R')
      C
0099      RETURN
0100      END

```

PROGRAM FOR ESTIMATING SYSTEM RATES BASED ON SAMPLE VALUES PAIRED TO MEAN VALUES

This program for calculation of estimates of input, output, translocation, and decay was written in Fortran IV level G, and was run on an IBM 360/67. In our experience 112k was used and the program required approximately 40 seconds per run. A maximum of 60 stations, 7 chemical compounds, and 2 sample times is permitted with the program as written.

The time interval is calculated in the subroutine, NCOMP, which calls the subroutine, LEAPYR. K values are calculated using double precision, and confidence intervals are estimated through use of a table of "t values."

There are eight cards which precede the data deck. Their formats and content are as follows:

First four cards, as in preceding program.

Fifth card, Format (3I5), contains the number of stations at time one, followed by the number of stations at time two, followed by the number of chemical compounds in the data set.

Sixth through eighth cards, Format (10A8), as in preceding program.

The data is organized as in the preceding program but is sorted chronologically.

An optional subroutine FACTOR may be called by placing a card before the END card with CALL FACTOR.

```

      C      PROGRAM FOR ESTIMATING SYSTEM RATES BASED ON
      C      SAMPLE VALUES PAIRED TO MEAN VALUES.
      C
      C
0001      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8),
      1, STE(23,8),CL95(23,8),VAR1(7),VAR2(7),VAR3(7),SUM1(7),SUM 2(
      27), SUM3(7), SUM4(7), COV1(7), COV2(7), IA(2), AI(2)
0002      REAL *4MEAN,M(17,8),MR(7)
0003      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7).
0004      INTEGER CST(2,60),CDATE(2,60,3)
0005      COMMON X,TABLE,IA,IB,I,K,KD,ID
0006      COMMON /BLK2/MR
0006      1 FORMAT (1X,13F6.3/13F6.3/4F6.3)
0007      READ (5,1) TABLE
0008      2 FORMAT (12I4)
0009      READ (5,2) MONTH
0010      READ (5,3) IA(1),IA(2),ID
0011      3 FORMAT (3I5)
0012      C

```

	C	CALCULATE INDEXES.
	C	AI NUMBER OF STATIONS CONVERTED TO A REAL NUMBER.
	C	AI3 AI(1) + AI(2)
	C	IA3 IA(1) + IA(2)
	C	IP1 ID + 1
	C	IP2 ID + 2
	C	I2TP2 2 * ID + 2
	C	I2TP3 2 * ID + 3
	C	I3TP2 3 * ID + 2
	C	J2T IA(1) + IA(2)
0013		AI(1)=IA(1)
0014		AI(2)=IA(2)
0015		AI3=AI(1)+AI(2)
0016		IA3=IA(1)+IA(2)
0017		IP1=ID+1
0018		IP2=ID+2
0019		I2TP2=2*ID+2
0020		I2TP3=2*ID+3
0021		I3TP2=3*ID+2
0022		J2T=IA(1)+IA(2)
	C	
	C	CLEAR X ARRAY.
	C	
0023		DO 4 I=1,10
	C	
0024		DO 4 J=1,J2T
	C	
0025		DO 4 K=1,IP1
0026		4 X(I,J,K)=0.0
	C	
0027		WRITE (6,9)
	C	
	C	READ IN DERIVATIVE NAMES AND CONCENTRATION LEVEL ON UP TO 3 CARDS.
0028		READ (5,5) NAME
0029		5 FORMAT (10A8)
	C	
	C	READ IN DATA.
0030		6 FORMAT (1X,I2,2(2A4,A2),I2,2(1X,I2),7F7.2)
	C	
0031		DO 7 I=1,2
0032		IB=IA(I)
	C	
0033		DO 7 J=1,IB
0034		7 READ (5,6) CST(I,J),(ALOC(I,J,L),L=1,6),(CDATE(I,J,L),L=1,3),(X(I, 1J,K),K=1,ID)
	C	
	C	
	C	COMPUTE TOTAL OF EACH STATION.


```

C
0035      DO 8 I=1,2
0036      IB=IA(I)
C
0037      DO 8 J=1,IB
C
0038      DO 8 L=1,ID
0039      8 X(I,J,IP1)=X(I,J,L)+X(I,J,IP1)
C
C
C      WRITE HEADING OF FIRST TWO PAGES.
C
0040      DO 15 I=1,2
0041      IB=IA(I)
0042      L=I
0043      9 FORMAT ('1')
0044      10 FORMAT ('1','C'/2X,I1,/3X,'STATION',3X,'LATITUDE',3X,'LONGITUDE',
1 5X,'DATE')
0045      WRITE (6,10) I
0046      11 FORMAT (48X,8(3X,A8))
0047      12 FORMAT ('+',47X,8(3X,A8))
0048      WRITE (6,12) (NAME(N),N=1,IP1)
0049      13 FORMAT (/)
0050      WRITE (6,11) (NAME(N),N=IP2,I2TP2)
0051      WRITE (6,13)
C
0052      DO 14 K=1,IP1
0053      CALL STDEV (TOTAL,MEAN,SD,SE,CL)
0054      TOT(I,K)=TOTAL
0055      M(I,K)=MEAN
0056      STD(I,K)=SD
0057      STE(I,K)=SE
0058      14 CL95(I,K)=CL
C
C      L1=NUMBER OF SETS COMPUTED.
C
C      WRITE FIRST TWO PAGES.
0059      L1=IP1
0060      CALL PRINT
0061      WRITE (6,12) (NAME(N),N=1,IP1)
0062      WRITE (6,11) (NAME(N),N=IP2,I2TP2)
0063      WRITE (6,13)
0064      CALL PRINT2
0065      15 CONTINUE
C
C
C      COMPUTE PERCENTS.

```

```

C
0066      DO 18 I=3,4
0067      IB=IA(I-2)
0068      L1=ID
0069      L=I-2
0070      WRITE (6,10) L
0071      WRITE (6,12) (NAME(N),N=1,ID)
0072      WRITE (6,11) (NAME(N),N=I2TP3,I3TP2)
0073      WRITE (6,13)

C
0074      DO 16 K=1,ID

C
0075      DO 16 J=1,IB
0076      16 X(I,J,K)=X(L,J,K)/X(L,J,IP1)*100.

C
C
0077      DO 17 K=1,3
0078      CALL STDEV (TOTAL,MEAN,SD,SE,CL)
0079      TOT(I,K)=TOTAL
0080      M(I,K)=MEAN
0081      STD(I,K)=SD
0082      STE(I,K)=SE
0083      17 CL95(I,K)=CL

C
0084      CALL PRINT
0085      WRITE (6,12) (NAME(N),N=1,ID)
0086      WRITE (6,11) (NAME(N),N=I2TP3,I3TP2)
0087      WRITE (6,13)
0088      CALL PRINT2
0089      18 CONTINUE

C
0090      CALL NCOMP

C
C      CALCULATE TOTAL AND MEAN OF N.
C
0091      DO 20 K=1,ID
0092      TOT(5,K)=0.

C
0093      DO 19 J=1,IA3
0094      19 TOT(5,K)=TOT(5,K)+X(5,J,K)

C
0095      20 M(5,K)=TOT(5,K)/AI3

C
C
0096      DO 22 K=1,ID
0097      V=0.0

C

```

```

0098      DO 21 J=1,IA3
0099      21 V=(M(5,K)-X(5,J,K))**2+V
      C
0100      STD(5,K)=SQRT(V/(AI3-1.0))
0101      22 CALL STDEV2 (STD(5,K),STE(5,K),CL95(5,K))
      C
      C
      C      CALCULATE K VALUES.
      C      DATA IN TWO SETS ARRANGED CHRONOLOGICALLY
      C      CALCULATE K VALUES
      C
0102      DO 24 K=1,ID
0103      SUM1(K)=0.0
0104      IB=IA(2)
      C
0105      DO 23 J=1,IB
0106      IF (X(2,J,K).EQ.0.) X(2,J,K)=.004
0107      V=(DLOG10(X(2,J,K))-ALOG10(M(1,K)))/(X(5,J,K))
0108      V2(J,K)=10.**V-1.0
0109      23 SUM1(K)=SUM1(K)+V2(J,K)
      C
0110      IB=IA(1)
      C
0111      DO 24 J=1,IB
0112      IF (X(1,J,K).EQ.0.) X(1,J,K)=.004
0113      V=ALOG10(M(2,K))-DLOG10(X(1,J,K)))/X(5,J+IA(2),K)*.43429)
0114      V2(J+IA(2),K)=10.**V-1.0
0115      24 SUM1(K)=SUM1(K)+V2(J+IA(2),K)
      C
      C
      C      SORT VALUES
      C
0116      DO 26 K=1,ID
      C
0117      DO 26 J=1,J2T
0118      IF (V2(J,K).GT.0.) GO TO 25
0119      X(7,J,K)=V2(J,K)
0120      GO TO 26
0121      25 X(6,J,K)=V2(J,K)
0122      26 X(8,J,K)=X(7,J,K)+X(6,J,K)
      C
      C      CALCULATE K-NET
      C
0123      DO 28 K=1,ID
      C

```

```

0124      DO 28 J=1,J2T
0125      V=X(8,J,K)-SUM1(K)/AI3
0126      IF (V.GT.0) GO TO 27
0127      X(10,J,K)=V
0128      GO TO 28
0129      27 X(9,J,K)=V
0130      28 CONTINUE

C
C      COMPUTE SUM & MEAN FOR K VALUES
C
0131      DO 30 K=1,ID
C
0132      DO 30 I=6,10
0133      V=0.0
C
0134      DO 29 J=1,J2T
0135      29 V=V+X(I,J,K)
C
0136      TOT(I,K)=V
0137      30 M(I,K)=V/AI3
C
0138      DO 31 I=6,10
C
0139      DO 31 K=1,7
0140      STD(I,K)=0.0
0141      STE(I,K)=0.0
0142      31 CL95(I,K)=0.0
C
C      CALCULATE STANDARD DEVIATION, STANDARD ERROR, AND 95% CONFIDENCE
C      LIMITS OF K VALUES.
C
0144      DO 32 K=1,ID
0145      SUM1(K)=0.0
0146      SUM2(K)=0.0
0147      SUM3(K)=0.0
0148      32 SUM4(K)=0.0
0149      IB=IA(2)
C
0150      DO 33 J=1,IB
0151      V2(J,K)=DLOG(X(2,J,K))-DLOG(X(1,J,K))
0152      SUM2(K)=V2(J,K)+SUM2(K)
0153      SUM3(K)=(DLOG(X(1,J,K))-ALOG(M(1,K)))**2+SUM3(K)
0154      33 SUM4(K)=(DLOG(X(2,J,K))-ALOG(M(2,K)))**2+SUM4(K)
C
0155      DO 34 K=1,ID
C

```

```

0156      VAR1(K)=(.43429/M(1,K))**2*SUM3(K)/(AI3-1.0)+(-.43429/M(2,K))**2
          1*SUM4(K)/(AI3-1.0)
0157      34 V1(K)=SUM2(K)/AI3
          C
0158      DO 36 K=1,ID
0159      VAR2(K)=((1.0/M(5,K))**2*VAR1(K))+(-V1(K)/M(5,K)**2))**2*STD(5,K)
          1**2
0160      VAR2(K)=10.0**VAR2(K)
0166      STD(8,K)=SQRT(VAR2(K))
0167      36 CALL STDEV2(STD(8,K),STE(8,K),CL95(8,K))
          C
          C      CALCULATE THE DISTRIBUTION OF VARIANCE BETWEEN +K AND-K.
          C
0168      DO 41 K=1,ID
0169      V=0.0
          C
0170      DO 38 J=1,J2T
0171      IF (X(6,J,K))38,38,37
0172      37 V=(X(6,J,K)-M(8,K))**2+V
0173      38 CONTINUE
          C
0174      V=V/AI3-1.0
0175      W=0.0
          C
0176      DO 40 J=1,J2T
0177      IF (X(7,J,K)) 39,40,40
0178      39 W=(X(7,J,K)-M(8,K))**2+W
0179      40 CONTINUE
          C
0180      W=W/AI3-1.0)
0181      U=V+W
0182      V=STD(8,K)**2*(V/U)**2
0183      STD(6,K)=SQRT(V)
0184      W=STD(8,K)**2*(W/U)**2
0185      STD(7,K)=SQRT(W)
0186      CALL STDEV2(STD(6,K),STE(6,K),CL95(6,K))
0187      41 CALL STDEV2(STD(7,K),STE(7,K),CL95(7,K))
          C
          C      CALCULATION OF STANDARD DEVIATION K-NET AND ITS DISTRIBUTION.
          C
0188      DO 46 K=1,ID
0189      V=0.0
0190      W=0.0
          C
0191      DO 45 J=1,J2T
0192      IF(X(9,J,K))43,43,42
0193      42 V=V+(X(9,J,K))**2)
0194      43 IF(X(10,J,K))44,45,45
0195      44 W=W+(X(10,J,K)**2)
0196      45 CONTINUE

```

```

C
0197      V=V/(AI3-1.0)
0198      W=W/(AI3-1.0)
0199      STD(9,K)=SQRT(((V/(V+W))**2*(STD(8,K)**2))
0200      CALL STDEV2(STD(9,K),STE(9,K),CL95(9,K))
0201      STD(10,K)=SQRT(((W/(V+W))**2*(STD(8,K)**2))
C
0202      46 CALL STDEV2(STD(10,K),STE(10,K),CL95(10,K))
0203      CALL PRINT3
C
C      CALCULATE O AND ITS STANDARD DEVIATION
C
0203      DO 52 K=1,ID
0204      M(11,K)=(M(9,K)/M(6,K))*M(7,K)
0205      STD(11,K)=SQRT(STD(7,K)**2*((M(9,K)/M(6,K))**2))
0206      CALL STDEV2 (STD(11,K),STE(11,K),CL95(11,K))
C
C
C      CALCULATION OF D
0207      M(12,K)=M(7,K),-M(11,K)
C
C      CALCULATION OF STANDARD DEVIATION OF D
C
0208      STD(12,K)=SQRT(STD(7,K)**2*(1.-M(9,K)/M(6,K))**2)
0209      CALL STDEV2 (STD(12,K),STE(12,K),CL95(12,K))
C
C      CALCULATE TL.
C
0222      M(13,K)=-1.0*(1.0/M(12,K))
0223      STD(13,K)=DSQRT(STD(12,K)**2*(1.0/M(12,K)**2)**2)
0224      52 CALL STDEV2(STD(13,K),STE(13,K),CL95(13,K))
C
C      CALCULATE TR.
C
0225      DO 53 K=1,ID
0226      M(14,K)=-1.0*(1.0/M(7,K))
0227      STD(14,K)=SQRT(STD(7,K)**2*(1.0/M(7,K)**2)**2)
0228      53 CALL STDEV2 (STD(14,K),STE(14,K),CL95(14,K))
C
0229      DO 71 K=1,ID
0230      WRITE (6,54) NAME(K)
0231      54 FORMAT ('1',1X,'RATES OF CHANGE FOR',A8,30X,'S.D.',7X,'S.E.',4X,
1 '95%LIMIT'//)
0232      WRITE (6,55) NAME (K),M(8,K),STD(8,K),STE(8,K),CL95(8,K)
0233      55 FORMAT (2X,'MEAN OF K',13X,'=NET',3X,A8,'=',3X,4F11.4/)
0234      WRITE (6,56) NAME(K),M(9,K),STD(9,K),STE(9,K),CL95(9,K)
0235      56 FORMAT (2X, 'MEAN OF + ( K-NET ) = T' ,5X,A8,'=',3X,4F11.4)
0236      WRITE (6,57)

```

```

0237      57 FORMAT (27X,'I'/)
0238      WRITE (6,58) NAME(K),M(10,K),STD(10,K),STE(10,K),CL95(10,K)
0239      58 FORMAT (2X,'MEAN OF - ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0240      WRITE (6,59)
0241      59 FORMAT (27X,'O'/)
0242      WRITE (6,60) NAME(K),M(6,K),STD(6,K),STE(6,K),CL95(6,K)
0243      60 FORMAT (2X,'MEAN OF + K',11X,'= I',5X,A8,'=',3X,4F11.4//)
0244      WRITE (6,61) NAME(K),M(7,K),STD(7,K),STE(7,K),CL95(7,K)
0245      61 FORMAT (2X,'MEAN OF - K',11X,'= O + D',1X,A8,'=',3X,4F11.4//)
0246      WRITE (6,62) NAME(K),M(11,K),STD(11,K),STE(11,K),CL95(11,K)
0247      62 FORMAT (26X,'O',5X,A8,'=',3X,4F11.4/)
0248      WRITE (6,63) NAME(K),M(12,K),STD(12,K),STE(12,K),CL95(12,K)
0249      63 FORMAT (26X,'D',5X,A8,'=',3X,4F11.4/)
0250      WRITE (6,64) NAME(K),M(13,K),STD(13,K),STE(13,K),CL95(13,K)
0251      64 FORMAT (26X,'T',5X,A8,'=',3X,4F11.4)
0252      WRITE(6,65)
0253      65 FORMAT(27X,'L'/)
0254      WRITE(6,64) NAME(K),M(14,K),STD(14,K),STE(14,K),CL95(14,K)
0255      WRITE(6,66)
0256      66 FORMAT(27X,'R')
0257      DO 67 L=1,3
0258      67 WRITE(6,13)
0259      WRITE(6,68)
0260      68 FORMAT(13X,'MEAN C',6X,'MEAN C',16X,'I',10X,'T',6X,'-',4X,'T',6X,
-      '- ',5X,'O',5X,'-',5X,'D',9X,'N'/19X,'2',11X,'1',27X,'I',11X,'O'/)
0261      69 FORMAT(/97X,F11.4)
0262      WRITE(6,69) M(5,K)
0263      WRITE(6,70) NAME(K),M(2,K),M(1,K),M(6,K),M(9,K),M(10,K),M(11,K),
-      M(12,K)
0264      70 FORMAT(2X,A8,F10.4,'=',F10.4,' ( 1.0 +',F10.4,'+',F10.4,3(F12.4)
-      ',')')
0265      71 CONTINUE
0266      CALL FACTOR
0267      STOP
0268      END

C
C

0001      SUBROUTINE PRINT
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8).
1, STE (23,8),CL95(23,8),IA(2),AI(2)
0003      REAL *4MEAN,M(17,8)
0004      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7)
0005      INTEGER CST(2,60),CDATE(2,60,3)
0006      COMMON X, TABLE,IA,IB,I,K,KD,ID
0007      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,I
C

```

```

0008      DO 1 J=1,IB
0009      1 WRITE (6,3) CST(L,J),(ALOC(L,J,K),K=1,6),(CDATE(L,J,K),K=1,3),(X(I
      1,J,K),K=1,L1)
      C
      C      SKIP TO BOTTOM OF PAGE
0010      N=(68-(IB+6))/2
      C
0011      DO 2 J=1,N
0012      2 WRITE (6,4)
      C
0013      RETURN
      C
0014      3 FORMAT (5X,I2,5X,2A4,A2,2X,2A4,A2,2X,I2,2('-',I2),8F11.2)
0015      4 FORMAT (/)
0016      END
      C
      C
0001      SUBROUTINE PRINT2
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8),
      1,STE(23,8),CL95(23,8),IA(2),AI(2)
0003      REAL *MEAN,M(17,8)
0004      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7)
0005      INTEGER CST(2,60),CDATE(2,60,3)
0006      COMMON X,TABLE,IA,IB,I,K,KD,ID
0007      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0008      WRITE (6,1) (TOT(I,J),J=1,L1)
0009      WRITE (6,2) (M(I,J),J=1,L1)
0010      WRITE (6,3) (STD(I,J),J=1,L1)
0011      WRITE (6,4) (STE(I,J),J=1,L1)
0012      WRITE (6,5) (CL95(I,J),J=1,L1)
0013      RETURN
      C
0014      1 FORMAT (34X,'TOTALS',6X,7F10.4)
0015      2 FORMAT (/34X,'MEAN',8X,7F10.4)
0016      3 FORMAT (/34X,'S.D.',8X,7F10.4)
0017      4 FORMAT (/34X,'S.E.',8X,7F10.4)
0018      5 FORMAT (/34X,'95% CL',6X,7F10.4)
0019      END
      C
      C
0001      SUBROUTINE PRINT3
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8),
      1,STE(23,8),CL95(23,8),IA(2),AI(2)
0003      REAL *4MEAN,M(17,8)
0004      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7)
0005      INTEGER CST(2,60),CDATE(2,60,3)
0006      COMMON X,TABLE,IA,IB,I,K,KD,ID

```



```

0007      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0008      DO 19 K=1,ID
0009      WRITE (6,3)
0010      WRITE (6,4)
0011      WRITE (6,5) NAME(K),NAME(K)
0012      WRITE (6,6)
0013      WRITE (6,7)
0014      IB=IA(2)
      C
0015      DO 1 J=1,IB
0016      1 WRITE (6,17) CST(2,J),X(2,J,K),X(5,J,K),(X(IX,J,K),IX=6,10)
      C
0017      JPIA=IA(2)
0018      IB=IA(1)
      C
0019      DO 2 J=1,IB
0020      JPIA=JPIA+1
0021      2 WRITE (6,18) CST(1,J),X(1,J,K),X(5,JPIA,K),(X(IX,JPIA,K),IX=6,10)
      C
0022      WRITE (6,7)
0023      WRITE (6,16) TOT(2,K),TOT(1,K),TOT(5,1),(TOT(N,K),N=6,10)
0024      WRITE (6,15)
0025      WRITE (6,13) NAME(K)
0026      WRITE (6,16) M(2,K),M(1,K),M(5,1),(M(N,K),N=6,10)
0027      WRITE (6,8)
0028      WRITE (6,13) NAME(K)
0029      WRITE (6,16) STD(2,K),STD(1,K),STD(5,1),(STD(N,K),N=6,10)
0030      WRITE (6,9)
0031      WRITE (6,12) NAME(K)
0032      WRITE (6,16) STE(2,K),STE(1,K),STE(5,1),(STE(N,K),N=6,10)
0033      WRITE (6,10)
0034      WRITE (6,12) NAME(K)
0035      WRITE (6,16) CL95(2,K),CL95(1,K),CL95(5,1),(CL95(N,K),N=6,10)
0036      WRITE (6,11)
0037      19 WRITE (6,14) NAME(K)
0038      RETURN
      C
0039      3 FORMAT ('1',1X,'STATION')
0040      4 FORMAT (12X,'C',9X,'C',11X,'N',8X,'+ K',7X,'- K',6X,'+K + -K',
        1 4X,'+K - NET',3X,'-K - NET')
0041      5 FORMAT ('+',15X,A8,2X,A8)
0042      6 FORMAT (13X,'2',9X,'1',52X,'R',10X,'R/')
0043      7 FORMAT (/)
0044      8 FORMAT ('+',94X,'MEANS')
0045      9 FORMAT ('+',94X,'S.D.')
0046      10 FORMAT ('+',94X,'S.E.')

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0047      11 FORMAT ('+',94X,'95% CONFIDENCE LIMITS')
0048      12 FORMAT ('+', 99X,A8)
0049      13 FORMAT ('+',102X,A8)
0050      14 FORMAT ('+',116X,A8)
0051      15 FORMAT ('+',94X,'TOTALS')
0052      16 FORMAT (/9X,5F10.4,3F11.4)
0053      17 FORMAT (4X,I2,1X,F10.2,12X,3F10.4,3F11.4)
0054      18 FORMAT(4X,I2,1X,10X,F10.2,2X,3F10.4,3F11.4)
0055      END

```

C
C

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0001      SUBROUTINE TDIST (KA,T)
0002      REAL *8X(10,60,7)
0003      DIMENSION TABLE(30), IA(2), AI(2)
0004      COMMON X, TABLE, IA, IB, I, K, KD, ID
0005      I1=KA-1
0006      AK=I1
0007      IF (I1) 1,1,2
0008      1 WRITE (6,11) I
0009      GO TO 10
0010      2 IF (I1.LT.31) GO TO 9
0011      IF (I1.LT.41) GO TO 3
0012      GO TO 4
0013      3 TINT=((2.042-2.021)/10.)*(AK-30.)
0014      T=TINT+2.042
0015      GO TO 10
0016      4 IF (I1.Lt.61) GO TO 5
0017      GO TO 6
0018      5 TINT=((2.021-2.000)/20.)*(AK-40.)
0019      T=TINT+2.021
0020      GO TO 10
0021      6 IF (I1.LT.121) GO TO 7
0022      GO TO 8
0023      7 TINT=((2.000-1.980)/40.)*(AK-60.)
0024      T=TINT+2.000
0025      GO TO 10
0026      8 T=1.960
0027      GO TO 10
0028      9 T=TABLE(I1)
0029      10 RETURN

```

C

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0030      11 FORMAT ('1','I IN T TABLE =' ,I3)
0031      END

```

C
C

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0001      SUBROUTINE STDEV (SUMX,XBAR,STD,STE,CL$)
0002      REAL *8X(10,60,7)
0003      DIMENSION TABLE(30), IA(2), AI(2)
0004      COMMON X,TABLE,IA,IB,I,K,KD,ID
0005      DEV=0.
0006      SUMX=0.

      C
0007      DO 1 J=1,IB
0008      1 SUMX=SUMX+X(I,J,K)

      C
0009      AI(I)=IA(I)
0010      XBAR=SUMX/AI(I)

      C
0011      DO 2 J=1,IB
0012      DEV=(XBAR-X(I,J,K))**2+DEV
0013      2 CONTINUE

      C
0014      STD=SQRT(DEV/(AI(I)-1.))
0015      STE=STD/SQRT(AI(I))
0016      KA=IB
0017      CALL TDIST (KA,T)
0018      CL$=T*STE
0019      END

      C
      C
0001      SUBROUTINE STDEV2 (STD,STE,CL$)
0002      REAL *8X(10,60,7)
0003      DIMENSION TABLE(30), IA(2), AI(2).
0004      COMMON X,TABLE,IA,IB,I,K,KD,ID
0005      AI3=IA(1)+IA(2)
0006      STE=STD/SQRT(AI3)
0007      KA=IA(1)+IA(2)
0008      CALL TDIST (KA,T)
0009      CL$=T*STE
0010      RETURN
0011      END

      C
      C
0001      SUBROUTINE NCOMP
0002      DIMENSION TABLE(30),MONTH(12),ALOC(2,60,6),TOT(10,8),STD(23,8)
1, STE(23,8),CL95(23,8),IA(2), AI(2)
0003      DIMENSION IYRVAL(5),ITOTDA(5)
0003      REAL *4M(17,8)
0004      REAL *8X(10,60,7),NAME(23)
0005      INTEGER CST(2,60),CDATE(2,60,3)
0006      INTEGER SUMDA(2)
0007      COMMON X,TABLE,IA,IB,I,K,KD,ID
0008      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1
0009      IJ=0

```

```

C
0010      DO 13 I=1,2
0011      K=0
0012      SUMDA(I)=0
0013      ITOTDA(I)=0
C      STORE INITIAL TIME
0014      MO1=CDATE(I,1,1)
0015      IDA1=CDATE(I,1,2)
0016      IYR1=CDATE(I,1,3)
0017      IYRVAL(1)=365
0018      A=IYR1
0019      IZ=A/4.
0020      IF (IZ*4.EQ.IYR1.AND.MO1.GT.2) IYRVAL(1)=366
0021      INT1=MONTH(MO1)+IDA1
C      FIND TIME INTERVAL OF FIRST DATE TO END OF FIRST YEAR
0022      INT2=IYRVAL(1)-INT1
0023      IB=IA(I)
C
0024      DO 4 J=1,IB
0025      MO2=CDATE(I,J,1)
0026      IDA2=CDATE(I,J,2)
0027      IYR2=CDATE(I,J,3)
C      COMPUTE YEAR VALUES-365 OR 366
0028      IF (IYR1.EQ.IYR2) TO TO 3
C      K STORES NUMBER OF INTERVENING YEARS
0029      K=IYR2-IYR1
C
0030      DO 1 L=1,K
0031      IYRVAL(L+1)=365
0032      A=IYR1+L
0033      IZ=A/4.
0034      IF (IZ*4.EQ.IYR1+L) IYRVAL(L+1)=366
0035      1 CONTINUE
C
C      COMPUTE INTERVAL OF LAST YEAR
0036      LAST1=MONTH(MO2)+IDA2
C      CHECK FOR LEAPYR OF LAST YEAR
0037      IF (IYRVAL(K).EQ.366.AND.MO2.GT.2) LAST=LAST+1
0038      LAST2=IYRVAL(K)-LAST1
C      COMPUTE TOTAL DAYS OF DATA SET
C      INT=FIRST YEAR
0039      K=K+1
C      K= NUMBER OF YEARS
C

```

```

0040      DO 2 L=1,K
0041      2 ITOTDA(I)=ITOTDA(I)+IYRVAL(L)
      C
      C      SUM ALL DAYS OF YEARS INVOLVED
0042      ITOTDA(I)=ITOTDA(I)-INT1-LAST2
0043      SUMDA(I)=SUMDA(I)+ITOTDA(I)
0044      GO TO 4
0045      3 INT2=MONTH(MO2)+IDA2
0046      ITOTDA(I)=INT2-INT1
0047      SUMDA(I)-SUMDA(I)+ITOTDA(I)
0048      4 CONTINUE
      C
      C      COMPUTE MEAN OF TIME
0049      MEANT=SUMDA(I)/IA(I)
      C      SUBTRACT FIRST YEAR
0050      IX=MEANT+INT1
0051      IF (K.EQ.0) GO TO 7
      C
0052      DO 5 L=1,K
0053      IF (IX.LT.IYRVAL(L)) GO TO 6
0054      IF (IX.EQ.IYRVAL(L)) TO TO 6
0055      IX=IX-IYRVAL(L)
0056      5 CONTINUE
      C
      C      COMPUTE YEAR
0057      6 IYR=L-1+IYR1
0058      IF (IYRVAL(L).EQ.366.AND.IX.GT.59) IX=IX-1
0059      GO TO 8
0060      7 IF (IYRVAL(1).EQ.366.AND.IX.GT.59) IX=IX-1
0061      IYR=CDATE(I,1,3)
      C
0062      8 DO 9 N=1,12
      C      LOCATE MONTH
0063      IF (IX.LT.MONTH(N+1)) GO TO 10
0064      IF (IX.EQ.MONTH(N)) GO TO 10
0065      9 CONTINUE
      C
0066      10 IMON=N
0067      IDAY=IX-MONTH(N)
0068      IF (I.EQ.1) IC=IA(2)
0069      IF (I.EQ.2) IC=IA(1)
      C
0070      DO 12 J=1,IC
0071      IJ=IJ+1
0072      CALL LEAPYR (J,IMON,IDAY,IYR)
      C

```

```

0073      DO 11 K=1,ID
0074      X(5,IJ,K)=YR
0075      11 CONTINUE
      C
0076      12 CONTINUE
      C
0077      13 CONTINUE
      C
0078      RETURN
0079      END
      C
      C

0001      SUBROUTINE LEAPYR (J,IMON,IDAY,IYR)
0002      DIMENSION TABLE(30), MONTH(12), ALOC(2,60,6), TOT(10,8), STD(23,8)
0003      1,STE(23,8), CL95(23,8), IA(2), AI(2)
0003      REAL *4MEAN, M(17,8)
0004      REAL *8X(10,60,7),NAME(23)
0005      INTEGER YR1,YR2,DA1,DA2,DAYS
0006      INTEGER CST(2,60),CDATE(2,60,3)
0007      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,L1,L
0008      COMMON X,TABLE,IA,IB,I,K,KD,ID
0009      DAYS=0
0010      NT=0
0011      IF (I.EQ.2) GO TO 1
0012      MO1=IMON
0013      DA1=IDAY
0014      YR1=IYR
0015      MO2=CDATE(2,J,1)
0016      DA2=CDATE(2,J,2)
0017      YR2=CDATE(2,J,3)
0018      GO TO 2
0019      1 MO2=IMON
0020      DA2=IDAY
0021      YR2=IYR
0022      MO1=CDATE(1,J,1)
0023      DA1=CDATE(1,J,2)
0024      YR1=CDATE(1,J,3)
0025      2 AMO=MO1
      C

0026      DO 6 IY=YR1,YR2
0027      A=IY
0028      LEAP=0
0029      IZ=A/4.
0030      Z=IZ
0031      Z=Z*4.
0032      IF (IY.EQ.YR1) GO TO 3
0033      GO TO 4

```

```

0034      3 DAYS=365-(MONTH(MO1)+DA1)
0035      IF (Z.EQ.A.AND.AMO.LT.3.) LEAP=1
0036      GO TO 5
0037      4 IF (Z.EQ.A) LEAP=1
0038      5 NT=DAYS+LEAP+NT
0039      6 DAYS=365

```

C

```

0040      IF (LEAP.EQ.1) GO TO 7
0041      GO TO 8
0042      7 IF (MO2.LT.3) NT=NT-1
0043      8 YR=NT-365+MONTH(MO2)+DA2
0044      YR=YR/365.
0045      RETURN
0046      END

```

```

C      SUBROUTINE FOR PROGRAM FOR ESTIMATING SYSTEM RATES
C      BASED ON SAMPLE VALUES PAIRED TO MEAN VALUES.
      SUBROUTINE FACTOR
0001      DIMENSION TABLE(30), MONTH(12), ALOC(2,60,6), TOT(10,8), STD(23,8)
0002      1, STE(23,8), CL95(23,8), VAR1(7), VAR2(7), VAR3(7), SUM1(7), SUM2(
0003      27), SUM3(7), SUM4(7), COV1(7), COV2(7), IA(2), AI(2)
0002      REAL *4MEAN,M(17,8),MR(7)
0003      REAL *8X(10,60,7),V2(60,7),NAME(23),V1(7)
0004      INTEGER CST(2,60),CDATE(2,60,3)
0005      COMMON X,TABLE,IA,IB,I,K,KD,ID
0006      COMMON /BLK1/ NAME,TOT,M,STD,STE,CL95,ALOC,YR,CST,CDATE,MONTH,LI,L
0007      COMMON /BLK2/ MR
C
C
C      CALCULATE CORRECTION FACTOR FOR STANDARD DEVIATION
C
0008      IA3=IA(1)+IA(2)
0009      AI3=IA3
C
0010      DO 14 K=1, ID
C
0011      IF (M(8,K)) 1,4,5
0012      1 JX=(AI3*M(7,K)-AI3*M(10,K))/M(8,K)
0013      VJ=JX
0014      V=((AI3*M(10,K))/VJ)**2)*VJ
0015      IF ((M(8,K)+((AI3*M(10,K))/VJ))*VJ-(AI3*M(7,K))) 3,3,2
0016      2 V=V+(AI3*M(7,K)-VJ*(M(8,K)+((AI3*M(10,K))/VJ))-M(8,K))**2
0017      V=V+(((AI3*M(6,K))/AI3-VJ-1.0))-M(8,K))**2*(AI3-VJ-1.0)
0018      GO TO 8
0019      3 V=V+(((AI3*M(6,K))/(AI3-VJ))-M(8,K))**2*(AI3-VJ)
0020      GO TO 8
0021      4 JX-AI3/2.0
0022      VJ=JX
0023      V=((AI3*M(6,K)/VJ)**2)*VJ
0024      V=V+((AI3*M(7,K)/VJ)**2)*VJ
0025      GO TO 8
0026      5 JX=(AI3*M(6,K)-AI3*M(9,K))/M(8,K)
0027      VJ=JX
0028      V=(((AI3*M(9,K))/VJ)**2)*VJ
0029      IF ((M(8,K)+((AI3*M(9,K))/VJ))*VJ-(AI3*M(6,K))) 6,7,7
0030      6 V=V+(AI3*M(6,K)-VJ*(M(8,K)+((AI3*M(9,K))/VJ))-M(8,K))**2
0031      V=V+(((AI3*M(7,K))/(AI3-VJ-1.0))-M(8,K))**2*(AI3-VJ-1.0)
0032      GO TO 8
0033      7 V=V+(((AI3*M(7,K))/(AI3-VJ))-M(8,K))**2*(AI3-VJ)
0034      8 V=V/(AI3-1.0)
      W=0.0
C
C

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```

0035      DO 9 J=1,IA3
0036      9 W=W+(X(8,J,K)-M(8,K))**2
      C
      C
0037      W=W/(AI3-1.0)
0038      C=(W-V)/W)**2
      C
      C      CALCULATE CORRECTED STD,6,7,AND 8
      C      STD(15,K) IS CORRECTED STD(6,K)
      C
0039      STD(15,K)=SQRT(C*STD(6,K)**2)
0040      CALL STDEV2 (STD(15,K),STE(15,K),CL95(15,K))
      C
      C      STD(16,K)IS CORRECTED STE(7,K)
      C
0041      STD(16,K)=SQRT(C*STD(7,K)**2)
0042      CALL STDEV2 (STD(16,K),STE(16,K),CL95(16,K))
      C
      C      STD(17,K) IS CORRECTED STD(8,K)
      C
0043      STD(17,K)=SQRT(C*STD(8,K)**2)
0044      CALL STDEV2 (STD(17,K),STE(17,K),CL95(17,K))
      C
      C      CALCULATE CORRECTED STD(9,K) AND STD(10,K)
      C
0045      V=0.0
      C
      C
0046      DO 11 J=1,IA3
      C
0047      IF (X(9,J,K)) 11,11,10
0048      10 V=V+X(9,J,K)**2
0049      11 CONTINUE
      C
      C
      C
0050      V=V/(AI3-1.0)
0051      W=0.0
      C
      C
0052      DO 13 J=1,IA3
      C
0053      IF (X(10,J,K)) 12,13,13
0054      12 W=W+(X(10,J,K))**2
0055      13 CONTINUE
      C
      C
      C

```

```

0056      W=W/(AI3-1.0)
      C
      C      STD(18,K) IS CORRECTED STD (9,K)
      C
0057      STD(18,K)=SQRT((V/(V+W))**2*C*(STD(8,K)**2))
0058      CALL STDEV2 (STD(18,K),STE(18,K),CL95(18,K))
      C
      C
      C      STD(19,K) IS CORRECTED STD(10,K)
      C
0059      STD(19,K)=SQRT((W/(V+W))**2*C*(STD(8,K)**2))
0060      CALL STDEV2 (STD(19,K),STE(19,K),CL95(19,K))
      C
      C      CALCULATE CORRECTED STD(20,K) CORRECTED STD(11,K)
      C
0061      STD(20,K)=SQRT(STD(16,K)**2*(M(9,K)/M(6,K))**2)
      C
      C
0062      CALL STDEV2 (STD(20,K),STE(20,K),CL95(20,K))
      C      CALCULATE STD(21,K) CORRECTED STD(12,K)
0063      STD(21,K)=SQRT(STD(16,K)**2*(1.0-M(9,K)/M(6,K))**2)
0064      CALL STDEV2 (STD(21,K),STE(21,K),CL95(21,K))
      C
      C      CALCULATE STD(22,K) CORRECTED STD(13,K)
      C
0065      STD(22,K)=SQRT(STD(21,K)**2*(1.0/M(12,K)**2)**2)
0066      CALL STDEV2 (STD(22,K),STE(22,K),CL95(22,K))
      C
      C      CALCULATE STD(23,K) CORRECTED STD(14,K)
      C
0067      STD(23,K)=SQRT(STD(16,K)**2*(1.0/M(7,K)**2)**2)
0068      CALL STDEV2 (STD(23,K),STE(23,K),CL95(23,K))
      C
      C
0069      14 CONTINUE
      C
      C
0070      DO 15 K=1,ID
      C
0071      WRITE (6,16) NAME(K)
0072      WRITE (6,17) NAME(K),M(8,K),STD(17,K),STE(17,K),CL95(17,K)
0073      WRITE (6,18) NAME(K),M(9,K),STD(18,K),STE(18,K),CL95(18,K)
0074      WRITE (6,19)
0075      WRITE (6,20) NAME(K),M(10,K),STD(19,K),STE(19,K),CL95(19,K)
0076      WRITE (6,21)
0077      WRITE (6,22) NAME(K),M(6,K),STD(15,K),STE(15,K),CL95(15,K)
0078      WRITE (6,23) NAME(K),M(7,K),STD(16,K),STE(16,K),CL95(16,K)

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0079      WRITE (6,24) NAME(K),M(11,K),STD(20,K),STE(20,K),CL95(20,K)
0080      WRITE (6,25) NAME(K),M(12,K),STD(21,K),STE(21,K),CL95(21,K)
0081      WRITE (6,26) NAME(K),M(13,K),STD(22,K),STE(22,K),CL95(22,K)
0082      WRITE (6,27)
0083      WRITE (6,26) NAME(K),M(14,K),STD(23,K),STE(23,K),CL95(23,K)
0084      WRITE (6,28)
0085      15 CONTINUE
      C
      C
0086      16 FORMAT ('1',40X,'CORRECTED STANDARD DEVIATIONS'//,2X, 'RATES OF CHA
      1NGE FOR,' A8,30X,'S.D.',7X,'S.E.',4X,'95% LIMIT'//)
0087      17 FORMAT (2X,'MEAN OF K',13X,'=NET',3X,A8,'=',3X,4F11.4/)
0088      18 FORMAT (2X,'MEAN OF + ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0089      19 FORMAT (27X,'I'/)
0090      20 FORMAT (2X,'MEAN OF - ( K - NET ) = T',5X,A8,'=',3X,4F11.4)
0091      21 FORMAT (27X,'O'/)
0092      22 FORMAT (2X,'MEAN OF + K',11X,'= I',5X,A8,'=',3X,4F11.4//)
0093      23 FORMAT (2X,'MEAN OF - K',11X,'= 0+D',1X,A8,'=',3X,4F11.4//)
0094      24 FORMAT (26X,'O',5X,A8,'=',3X,4F11.4/)
0095      25 FORMAT (26X,'D',5X,A8,'=',3X,4F11.4/)
0096      26 FORMAT (26X,'T',5X,A8,'=',3X,4F11.4)
0097      27 FORMAT (27X,'L'/)
0098      28 FORMAT (27X,'R')
      C
0099      RETURN
0100      END

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(Please read instructions on the reverse before completing)

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<p>18. ABSTRACT The concentrations of DDT, DDD and DDE were measured in sediments at 57 stations in Monterey Bay on the Central California coast during 1970-1971. Mean concentrations in parts per billion were DDT 3.1, DDD 2.3, and DDE 5.4. During 1973 nineteen of the original stations were sampled. Mean concentrations were DDT 15.5, DDD 2.3, and DDE 5.4 PPB.</p> <p>Two approaches to the estimation of annual system rates for input, I, output, O, decay, D, and internal translocation, T_I and T_O expressed as decimal fractions of existing concentrations were developed, and fraction programs that permit rapid estimations were written. The mean annual rates in South Monterey Bay obtained were for DDT, I +1.30, O -0.059, D -0.036 T_I and T_O +0.80 with a residence time of 11 years and life time of 29 years. An I of 1.30 means the amount of input is 130% of the existing concentration per year. Rates for DDD were, I +0.25, O -0.11, D -0.025, T_I and T_O +0.20 with residence time of 7 years and life time of 44 years. Rates for DDE were I +0.28, O -0.10, D -0.027, T_O and T_I +0.22 with residence time of 8 years and life time of 39 years.</p> <p>Laboratory assays were developed to determine the relative rate of decomposition in sediment under conditions selective for various physiologically different kinds of microorganisms. Decay under aerobic conditions was greater than under anaerobic conditions. Nitrate increased the rate of decomposition under anaerobic conditions. The Q_{10} for decay was 2.5.</p>			
17. KEY WORDS AND DOCUMENT ANALYSIS			
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