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## Source Assessment: Textile Plant Wastewater Toxics Study, Phase I

Monsanto Research Corp, Dayton, Ohio

Prepared for

Industrial Environmental Research Lab, Research Triangle Park, N C

Mar 78

## SOURCE ASSESSMENT: TEXTILE PLANT WASTEWATER TOXICS STUDY PHASE I



Industrial Environmental Research Laboratory
Office of Research and Development
U.S. Environmental Protection Agency
Research Triangle Park, North Carolina 27711

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and toxicological baseline data on wastewater samples collected from textile plants in the U.S. Raw waste and secondary effluent wastewater samples were analyzed for 129 consent decree priority pollutants, effluent guidelines criteria pollutants, and nutrients. Level 1 chemical analyses were also performed. Secondary effluent samples from the 23 plants selected for study in the EPA/ATMI BATEA Study (American Textile Manufacturers Institute/best available technology economically achievable) (EPA Grant 804329) were submitted for the following bioassays: mutagenicity, cytotoxicity, clonal assay, freshwater ecology series (fathead minnows, Daphnia, and algae), marine ecology series (sheepshead minnows, grass shrimp, and algae), 14-day rat acute toxicity, and soil microcosm. The bioassay results indicated that 10 of the 23 textile plants have secondary effluents sufficiently toxic to proceed to a second phase of the study. In the second phase, samples will be collected from these 10 plants to determine the level of toxicity removal attained by selected tertiary treatment technologies.

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Waste Water		Toxic Materials			
Toxicity		Source Assessment	06T		
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# SOURCE ASSESSMENT: TEXTILE PLANT WASTEWATER TOXICS STUDY PHASE I

by

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### PREFACE

The Industrial Environmental Research Laboratory (IERL) of the U.S. Environmental Protection Agency (EPA) has the responsibility for insuring that pollution control technology is available for stationary sources to meet the requirements of the Clean Air Act, the Federal Water Pollution Control Act, and solid waste legislation. If control technology is unavailable, inadequate, or uneconomical, then financial support is provided for the development of the needed control techniques for industrial and extractive process industries. The Chemical Processes Branch of the Industrial Processes Division of IERL has the responsibility for investing tax dollars in programs to develop control technology for a large number of operations (more than 500) in the chemical industries.

Monsanto Research Corporation (MRC) has contracted with EPA to investigate the environmental impact of various industries which represent sources of pollution in accordance with EPA's responsibility as outlined above. Dr. Robert C. Binning serves as MRC Program Manager in this overall program entitled "Source Assessment," which includes the investigation of sources in each of four categories: combustion, organic materials, inorganic materials, and open sources. Dr. Dale A. Denny of the Industrial Processes Division at Research Triangle Park serves as EPA Project Officer. Reports prepared in this program are of three types: Source Assessment Documents, State-of-the-Art Reports, and Special Project Reports.

Source Assessment Documents contain data on emissions from specific industries. Such data are gathered from the literature, government agencies, and cooperating companies. Sampling and analysis are also performed by the contractor when the available information does not adequately characterize the source emissions. These documents contain all of the information necessary for IERL to decide whether emissions reduction is required.

State-of-the-Art Reports include data on emissions from specific industries which are also gathered from the literature, government agencies, and cooperating companies. However, no extensive sampling is conducted by the contractor for such industries. Results from such studies are published as State-of-the-Art Reports for potential utility by the government, industry, and others having specific needs and interests.

Special projects provide specific information or services which are applicable to a number of source types or have special utility to EPA but are not part of a particular source assessment study. This special project report, "Source Assessment: Textile Plant Wastewater Toxics Study, Phase 1," was prepared to provide chemical and toxicological data on wastewater samples collected from selected textile plants in the United States. Dr. Max Samfield of the Chemical Processes Branch at IERL-RTP served as EPA Task Officer.

A second phase of this project is underway to collect samples of secondary effluents from 10 textile plants to determine the level of toxicity removal attained by selected tertiary treatment technologies.

#### **ABSTRACT**

The purpose of this study was to provide chemical and toxicological baseline data on wastewater samples collected from textile plants in the United States. Raw waste and secondary effluent wastewater samples were analyzed for 129 consent decree priority pollutants, effluent quidelines criteria pollutants, and nutrients; Level 1 chemical analyses were also performed. Secondary effluent samples from the 23 plants selected for study in the EPA/ATMI BATEA Study (American Textile Manufacturers Institute/ best available technology economically achievable) (Grant No. 804329) were submitted for the following bioassays: mutagenicity, cytotoxicity, clonal assay, freshwater ecology series (fathead minnows, Daphnia, and algae), marine ecology series (sheepshead minnows, grass shrimp, and algae), 14-day rat acute toxicity, and soil microcosm. Since this was a screening study, samples of the textile plant intake water were not collected for chemical analysis.

Based on the bioassay results, 10 of the 23 textile plants were found to have secondary effluents sufficiently toxic to proceed to a second phase of the study. In the second phase, samples will be collected from these 10 plants to determine the level of toxicity removal attained by selected tertiary treatment technologies.

This report was submitted in partial fulfillment of Contract 68-02-1874 by Monsanto Research Corporation under the sponsorship of the U.S. Environmental Protection Agency. This report covers a period from January 1977 to December 1977.

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#### ABBREVIATIONS

ATMI		American Textile Manufacturers Institute
ATP		adenosine triphosphate
BATEA		best available technology economically achievable
BOD <sub>5</sub>		5-day biochemical oxygen demand
COD		chemical oxygen demand
DO		dissolved oxygen
EC		electron capture detector on a gas chromatograph
EC <sub>50</sub>		effective concentration at which 50% of the test
		species reach the desired effect
GC		gas chromatograph
GCMA		gas chromatography mass analysis
ICAP		inductively coupled argon plasma
IR		infrared analyzer
LC		liquid chromatography
LC <sub>50</sub>		lethal concentration which causes 50% mortality
		in the test species
LD <sub>50</sub>		lethal dosage which causes 50% mortality in the test species
LRMS		low resolution mass spectrometer
MS		mass spectrometer
PCB		polychlorinated biphenyls
ppt		parts per thousand
RAM	_	rabbit alvelor macraphage
SSMS		spark source mass spectrometer
T/C		ratio of population density in treated samples
•		to population density in controls
TDS		total dissolved solids
TKN		total Kjeldahl nitrogen
TOC		total organic carbon
TSS		total suspended solids
v/v		volume-to-volume ratio
w/v		weight-to-volume ratio
•		<del></del>

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#### SECTION 1

#### INTRODUCTION

The Industrial Environmental Research Laboratory - RTP (IERL-RTP) of the U.S. Environmental Protection Agency (EPA) is currently engaged in a joint study with the American Textile Manufacturers Institute (ATMI) (EPA Grant No. 804329) to determine the best available technology economically achievable (BATEA) for textile plant wastewaters. A total of 23 textile mills representing eight textile processing categories and having well-operated secondary wastewater treatment facilities were selected by EPA and ATMI for the BATEA study. For that study, two mobile pilot plants containing four tertiary wastewater treatment technologies were constructed to gather technical data to identify the best available technology applicable to the 23 plants. Two additional tertiary treatment technologies were tested in the laboratory. The grant study focused on only a limited number of so-called criteria pollutants; i.e., 5-day biochemical oxygen demand, chemical oxygen demand, color, sulfides, total suspended solids, phenol, and pH.

However, on 7 June 1976 the U.S. District Court of Washington, D.C., issued a consent decree (resulting from Natural Resources Defense Council et al. v. Train) requiring EPA to enhance development of effluent standards. The court mandate focused federal water pollution control efforts on potentially toxic and hazardous pollutants. In response to the consent decree EPA developed a list of 129 specific compounds (known as priority pollutants) that the agency agreed to consider during the standards setting process. Based on the consent decree, EPA-IERL/RTP decided to conduct a study parallel to the ATMI/EPA Grant Study of the textile industry. The objective of the IERL/RTP study was to determine both the removal efficiencies for the 129 consent decree priority pollutants and the reduction in toxicity by the six tertiary treatment technologies being investigated under the original grant study.

The overall wastewater toxicity study is divided into two phases. The first, covered by this report, establishes a baseline data base concerning toxicity and level of priority pollutants present in raw wastewater and secondary effluents at 23 textile plants. These data are used to screen the 23 plants and to select those plants with secondary effluents of highest toxicity for further study. Toxicity tests were designed to evaluate

only the reduction in wastewater toxicity by control technologies, not the potential environmental impacts on receiving waters.

The second phase of the effort, to be covered in a subsequent report, will determine the reduction in priority pollutant concentrations and in toxicity by the mobile pilot plant tertiary treatment systems. Only those plants selected in the first phase of the study will be investigated.

Covering the first phase of the toxics study, this report describes sampling, chemical analysis, and bioassay procedures used to establish baseline data. Chemical analyses of raw waste and secondary effluents are presented for the 23 basic plants and for 9 additional textile plants. Bioassay data are presented for secondary effluents from the basic 23 plants.

The plants are ranked according to relative secondary effluent toxicity, and a number of plants are selected for study in the second phase of the overall program. Modifications and recommendations for improvements to the sampling, chemical analysis, and bioassay protocols are also discussed.

#### SECTION 2

#### SUMMARY

The purpose of this Phase I study was to provide chemical and toxicological baseline data on wastewater samples collected from selected textile plants in the United States. Raw waste (untreated wastewater) and secondary effluent (wastewater from secondary wastewater treatment facilities) samples were collected from the 23 textile plants selected for the ATMI/EPA Grant Study (Grant No. 804329) and from 9 additional textile plants. Since this was a screening study, samples of the textile plant intake water were not collected for chemical analysis.

Samples were analyzed for 129 consent decree priority pollutants and for effluent guidelines criteria pollutants. The Level 1 chemical analytical scheme developed by EPA-IERL/RTP was employed to detect other possible pollutants. Nutrient levels were measured at 23 of the 32 plants to supplement interpretation of algal bioassays. The following bioassays were performed on secondary effluent samples from the 23 plants chosen for the ATMI/EPA Grant Study: mutagenicity, cytotoxicity, clonal assay, freshwater ecology series (fathead minnows, Daphnia, and algae), marine ecology series (sheepshead minnows, grass shrimp, and algae), 14-day rat acute toxicity, and soil microcosm.

Grab samples and 8-hr continuous samples were collected both before and after the wastewater treatment system at each of the 32 plants. Samples were stored in ice at 4°C and shipped by air freight to the laboratories for analysis. Chemical analyses and bioassays were performed at eight EPA and commercial laboratories

Analysis for the 129 priority pollutants in raw waste and secondary effluent samples (totaling 64 samples) was performed by Monsanto Research Corporation (MRC). Analytical procedures followed those recommended by EPA. However, the recommended analytical protocol for priority pollutant analysis is still in the developmental stage and requires further verification and validation. Consequently, the analytical results of textile wastewater samples must be looked upon as good estimates of which priority pollutants are present, with concentrations accurate to within an order of magnitude.

The EPA analytical protocol divided the 129 priority pollutants into 5 fractions for analysis: volatile compounds, base/neutral compounds, acid compounds, pesticides and polychlorinated

biphenyls (PCB), and metals. EPA recommended that laboratories not acquire analytical standards for 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) because of its extreme toxicity. Asbestos was not analyzed due to the presence of interfering fibrous materials in textile wastewaters.

A summary of the organic compounds found in the 32 raw waste and 32 secondary effluent samples is given in Tables 1 and 2. Of the 114 organic compounds on the priority pollutant list, a total of 45 different compounds were found, 39 in raw waste samples and 34 in secondary effluent samples. On an individual plant basis the greatest number of organic compounds found in a raw waste and in a secondary effluent sample were 14 and 8, respectively, with an average number per plant of 7 in the raw waste and 5 in the secondary effluent. The predominant compounds were bis(2-ethylhexyl) phthalate in 54 samples (0.5 mg/m³ to 300 mg/m³), toluene in 44 samples (0.4 mg/m³ to 300 mg/m³), and ethylbenzene in 30 samples (0.7 mg/m³ to 3,000 mg/m³).

A summary of the 13 priority pollutant metals and cyanide concentrations in raw waste and secondary effluent samples is given in Table 3, which also summarizes the criteria pollutant and nutrient concentrations for secondary effluent samples. Nutrient analyses were performed to support freshwater algae bioassays.

On an individual plant basis it was frequently observed, especially for the metals data, that the concentration of a specific pollutant was greater in the secondary effluent sample than in the raw waste sample. This phenomenon is due, in part, to the hydraulic retention time of the wastewater treatment facility. Since raw waste and secondary effluent samples were collected simultaneously, concentrations in the secondary effluent were due to raw waste loads that entered the treatment system 1 day to 30 days prior to sampling. The average retention time for the 32 plants was about 5 days.

Level 1 chemical analyses were performed on secondary effluent samples from 15 of the 23 basic textile plants. Level 1 protocol identifies classes of compounds present in environmental samples and measures the general concentration range. Results indicate that total concentration of methylene chloride extractable organics ranges from  $3 \text{ g/m}^3$  to  $64 \text{ g/m}^3$ . This value is 5 to 10 times lower than the range for total organic carbon (Table 3).

In the Level 1 procedure each sample was fractionated by a liquid chromatography column into eight fractions based on polarity. Infrared analysis of each fraction indicated the presence of aliphatic hydrocarbons, esters and acids, aromatic compounds, phthalate esters, and fatty acid groups. Low resolution mass spectrophotometric analysis of the eight fractions of each sample detected the following types of compounds: paraffinic/olefinic,

TABLE 1. SUMMARY OF PRIORITY POLLUTANTS FOUND IN RAW WASTE SAMPLES, FROM 129 TOTAL, SHOWING CONCENTRATION RANGES AND NUMBER OF PLANTS WHERE THE SPECIES WERE IDENTIFIED

Volatile o	rganic		Base/neuti	al organic	
	Number			Number	
	of times	Concentration		of times	Concentration
Compound	found	range, mg/m <sup>3</sup>	Compound	found	range, mg/m
Toluene	22	2 to 300	Naphthalene	20	0.03 to 300
Benzene	4	5 to 200	Dimethyl phthalate	5	3 to 110
Chloroform	12	2 to 500	Diethyl phthalate	12	0.2 to 70
Chlorobenzene	6	1 to 300	Bis(2-ethylhexyl)	27	0.5 to 300
Ethylbenzene	20	0.7 to 2,800	phthalate		
Trichlorofluoromethane	2	<b>3</b> 0 to 50	1,4-Dichlorobenzene	5	1 to 210
1,1,1-Trichloroethane	5	2 to 300	1,2,4-Trichlorobenzene	8	30 to 440
Trichloroethylene	8	2 to 200	1,2-Dichlorobenzene	8	0.1 to 300
1,1,2,2-Tetrachloroethylene	8	15 to 1,100	Anthracene	1	0.1
Trans-1,2,-dichloroethylene	1	2	Pyrene	1	0.9
l,l-Dichloroethane	2	0.6 to 4	Acenaphthene	7	9 to 270
1,2-Dichloropropane	1	2	Di-n-butyl phthalate	6	2 to 23
Cis-1,3-dichloropropene	1	2	Fluorene	2	5 to 15
			Hexachlorobenzene	2	0.5 to 2
			N-Nitrosodiphenylamine	1	11
			2,6-Dinitrotoluene	1	50
			Indeno(1,2,3-cd)pyrene	1	2
Acid org	anic		Pes	ticide	
Phenol	19	0.5 to 100	β−ВНС	1	0.4
Phentachlorophenol	8	2 to 70	Heptachlor	1	6
2-Nitrophenol	1	70		_	-
p-Chloro-m-cresol	ī	5	•		
4-Nitrophenol	ī	70			
2,4,6-Trichlorophenol	2	0.7 to 20			
2-Chlorophenol	1	130	•		

G

TABLE 2. SUMMARY OF PRIORITY POLLUTANTS FOUND IN EFFLUENT SAMPLES, FROM 129 TOTAL, SHOWING CONCENTRATION RANGES AND NUMBER OF PLANTS WHERE THE SPECIES WERE IDENTIFIED

	rganic				Base/neutral	organic	
	Number			Number			
	of times	Concer				of times	Concentration
Compound	found	range	2, 1	ng/m <sup>3</sup>	Compound	found	range, mg/m
Toluene	22	0.4	to	110	Naphthalene	5	0.5 to 250
Benzene	2	0.5	to	60	Dimethyl phthalate	3	0.2 to 1
Chloroform	5	5	to	60	Diethyl phthalate	9	0.5 to 10
Chlorobenzene	2	4	to	30 '	Bis(2-ethylhexyl)phthalate	27	<b>2</b> to 230
Ethylbenzene	10	0.7	to	3,000	1,4-Dichlorobenzene	3	0.05 to 2
Trichlorofluoromethane	6 -	2	to	2,100	1,2,4-Trichlorobenzene	6	2 to 920
Trichloroethylene	2	5	to	80	1,2-Dichlorobenzene	5	0.2 to 25
1,1,2,2-Tetrachloroethylene	3	0.4	to	40	Anthracene	1	4
Cis-1,3-dichloropropene	1		6		N-Nitroso-di-n-propylamine	2	2 to 20
Trans-1,3-dichloropropene	2	0.9	to	4	Pyrene	4	0.1 to 0.3
Bromodichloromethane	1		2		Acenaphthene	2	0.5 to 2
					Di-n-butyl phthalate	3	4 to 60
					Hexachlorobenzene	3	0.3 to 0.8
					Butylbenzyl phthalate	1	<b>70</b>
Acid org	anic				Pesti	cides	
Phenol	2	2	to	3	α-ВНС	1	0.3
2,4-Dimethylphenol	2	8	to	9	Heptachlor	1	2
p-chloro-m-cresol	1		2		-		
2,4,6-Trichlorophenol	1		20				
Chloro cresol	1		30				
2-Chlorophenol	1		10				

SUMMARY OF METAL, CRITERIA POLLUTANT, AND NUTRIENT ANALYSES TABLE 3.

Metal				- a	2		
_	Concentration range, g/m <sup>3</sup>		Criteria p	ollutant"	Nutrient		
Element	Raw waste sample	Secondary effluent sample	Pollutant	Concentration		Concentration	
DIEMENC	Sample	erritent sample	FOITUCANC	range, g/m <sup>3</sup>	Parameter	range, g/m <sup>3</sup>	
Antimony	0.0005 to 0.06	0.0005 to 0.07	BOD <sub>5</sub> b	<5 to 170	Nitrite	0 to 17	
Arsenic	0.005 to 0.2	0.005 to 0.02	CODC	45 to 1,600	Nitrate	0.002 to 40	
Beryllium	<0.0001	<0.0001	Color (APHA)	10 to 2,500	Ammonia	0.02 to 14	
Cadmium	0.0005 to 0.05	0.0005 to 0.01	Sulfide	0.01 to 6	TKN <sup>e</sup>	2 to 40	
Chromium	0.0002 to 0.9	0.0002 to 2.0	Phenol	0.01 to 0.2	o-Phosphate	0.02 to 11	
Copper	0.0002 to 2.4	0.0002 to 0.3	TSS <sup>†</sup>	0.02 to 580	Total phosphorus	0.4 to 15	
Cyanide	0.004 to 0.2	0.004 to 0.2	pН	5.8 to 10	TOC	19 to 260	
Lead	0.001 to 0.2	0.001 to 0.2			•		
Mercury	0.0005 to 0.004	0.0005 to 0.0009					
Nickel	0.01 to 0.2	0.01 to 0.2					
Selenium	<0.005	<0.005					
Silver	0.005 to 0.1	0.005 to 0.1					
Thallium	<0.005	<0.005				•	
Zinc	0.03 to 8.0	0.07 to 38					

<sup>&</sup>lt;sup>a</sup>For secondary effluent samples.

Total Kjehldehl nitrogen.

f
Total suspended solids.

 $\mathbf{g}_{\mathtt{Total}}$  organic carbon.

b<sub>5-day</sub> biochemical oxygen demand.

Chemical oxygen demand.

 $<sup>{\</sup>sf d}_{\sf American}$  Public Health Association color standards.

bis (hydroxy-t-butyl phenol) propane, tri-t-butyl benzene, alkyl phenols, dichloroaniline, toluene-sulfonyl groups, vinyl stearate and azo compounds.

Bioassays used were selected by EPA and include tests for assessment of both health and ecological effects. Health effects tests estimate the potential mutagenicity, potential or presumptive carcinogenicity, and potential toxicity of the samples to mammalian organisms. Ecological effects tests focus on the potential toxicity of the samples to vertebrates (fish), invertebrates (daphnids and shrimp), and plants (algae) in freshwater, marine and terrestrial ecosystems.

A total of 8 bioassay systems were tested using 21 different tester organisms to evaluate the toxicity of secondary effluents. Table 4 lists the bioassays used and the purpose of each test.

TABLE 4. PURPOSE OF SELECTED BIOASSAY TESTS IN EVALUATING THE POTENTIAL TOXICITY OF SECONDARY EFFLUENTS

Bioassay system	Test organism	Purpose of test
Microbial mutagenicity	Nine different strains of bacteria and one of yeast.	To determine if a chemical mutagen (possibly a carcinogen) is present. These microbial strains were selected because of their sensitivity to various classes of chemical compounds.
Cytotoxicity	Rabbit alveolar cells and Chinese hamster ovary cells.	To measure metabolic impairment and death in mammalian cells. These primary cell cultures have some degree of metabolic repair capability.
Freshwater and marine static bioassay	Fathead minnow, daph- nids, sheepshead minnows, and grass shrimp.	To detect potential toxicity to organisms present in aquatic environments.
Freshwater and marine algal assay	Freshwater and marine algae.	To detect potential growth inhibition and stimulation effects on aquatic plants.
Range finding acute toxicity	Young adult rats.	Whole animal test to detect potential toxic effects to mammals. These live animals were selected because of the extensive data base on their response to known chemicals and because they have several metabolic systems closely approximating those in humans.
Terrestrial ecology	Soil microorganisms.	To determine potential inhibition and stimulation effects on soil microorganisms. These data are useful if the effluent is used for crop irrigation.

A summary of the bioassay results is presented in Table 5. Toxicity is expressed as the percent of a secondary effluent solution that will cause the effect specified for each bioassay over the testing period. For the cytotoxicity, Daphnia and algal bioassays an Effective Concentration 20 or 50 (EC<sub>20</sub> or EC<sub>50</sub>) was calculated. EC<sub>20</sub> for the cytotoxicity test means the concentration of secondary effluent which impairs metabolic processes in 20% of the test cells.

The viability test is a measure of the cells' ability to survive exposure to the sample, and the adenosine triphosphate (ATP) test measures the quantity of the coenzyme ATP produced, indirectly measuring cellular metabolic activity.

 $EC_{50}$  for the algal tests means the concentration of secondary effluent which causes a 50% reduction in algal growth as compared to a control sample. The freshwater algae test was performed over a 14-day period and the marine algae test over a 96-hr period.

For the fathead minnow, sheepshead minnow, and grass shrimp bioassays, death was used to measure toxicity, which was expressed as Lethal Concentration 50 ( $LC_{50}$ ).  $LC_{50}$  indicates the calculated concentration of secondary effluent that is expected to cause the death of 50% of the test species. Since rats were given a specific quantity of secondary effluent, toxicity was expressed as Lethal Dose 50 ( $LD_{50}$ ).  $LD_{50}$  indicates the quantity of material fed to the rats that resulted in the death of 50% of the test animals.

The measure of toxicity to a soil microcosm was the quantity of carbon dioxide  $(CO_2)$  produced after sample exposure as compared to a control sample. The quantity of  $CO_2$  produced over a 3-wk period after subtracting the quantity produced by the control was plotted on graph paper. The slope of the curve then represented the rate of increase or decrease in  $CO_2$  production due to addition to the sample.

Based on the bioassay results, the EPA Bioassay Subcommittee ranked the 23 plants according to their overall secondary effluent toxicity. The following nine textile plants were selected for further study based on their relatively high ranking: N, A, L, T, C, P, S, V, and W. Note the low toxicity response for Plant Y where the effluent samples were collected after the polishing pond.

During the second phase of this program the secondary effluents from these plants will be treated using the mobile pilot plants constructed under the ATMI/EPA Grant Study. Since effluent samples were inadvertently collected between the aeration lagoon and settling basin at Plant R, it will also be included in the Phase II program. While each of these 10 plants is being tested

TABLE 5. SUMMARY OF BIOTEST DATA FOR SECONDARY EFFLUENT WASTEWATER SAMPLES a, b

			Freshwater ecology series			Marine ecology series			
lant	Cytoto Viability (24-hr EC <sub>20</sub> ), secondary effluent	ATP (24-hr EC <sub>20</sub> ), % secondary effluent	Fathead minnow (96-hr LC <sub>50</sub> ), % secondary effluent	Daphnia (48-hr EC <sub>50</sub> ), * secondary effluent	Algae C (14-day EC20), * secondary effluent	Sheepshead minnow (96-hr LC <sub>50</sub> ), secondary effluent	Grass shrimp (96-hr LC <sub>50</sub> ), secondary effluent	Algae <sup>C</sup> (96-hr EC <sub>50</sub> ), • secondary effluent	Soil microsm, normalized relative CO <sub>2</sub> rate change
	NAT							f	~0.032
A		NAT	19.0	9.0	76	62.0	21.2	<u>_</u> 9	
В	NAT h	NATh	NAT	nat	30	NAT	NAT		0.020
С	16.8 <sup>n</sup>	6.1	46.5	41.0	0]	69 <sub>‡</sub> 5	12,8	9 <u>9</u>	-0.005
D	NAT	NAT	NAT	nat	0}	-'	-'	_	-0.099
E	NAT	NAT	NAT	7.8	27	NAT	nat	10 to 50	-0.048
F	· TAN	9.4	NAT	81.7	0;	NAT	NAT	85	-0.039
G	NAT	NAT	64,7	62.4	o'	NAT	NAT	59 _f	0.017
н	NAT	NAT	_k	40% dead at 100%	96	_T	_f	_T	-0.083
	NAT	NAT	*** m	concentration	,i	f	f	f	-0.163
J "			TAN	NAT	o;	-	-	- 77	-0.004
Κ.	NAT	NAT	NAT	TAN	0'	NAT	NAT		-0.020
T-	NAT	4.0	23.5	28.0	42 <sub>1</sub>	naţ	naţ	1.7	-0.059
M	NAT	NAT	NAT	60.0	o'k	<b></b>	<b></b> '	-1	0.059
N	13.3	3.8	48.8	100% dead at all	2	47.5	26.3	2.3	0.059
				dilutions		f	f		0.000
Pl	NAT	NAT	NAT	NAT	43	-f	-F	9≱0	0.022
R'	naţ	naŢ	16.5	8.0	934	-	<b>-</b> '	_g′	-0.062
S	_1	~'	nat	nsa <sup>m</sup>	o;	nat	NAT		-0.017
T	TAN	2.5	46.5	NAT	o;	68.0	34.5	70	0.020
U	NAT	nat	NAT	12.1	o;	naŢ	nat	_9	0.055
v	nat	nat	36.0	9.4	٥'	_f	_T	94	-0.066
W	NAT	13.7	55.2	6.3	94,	37.5	19.6	50	0.031
X	NAT	4.8	nat	NAT	o¦	nat	nat	-ģ	0.047
Y	NAT	NAT	NAT	nat	o'	- <u>Ţ</u>	- <del>T</del>	- <u>₹</u>	-0.172
Z	TAN	NAT	NAT	42.6	18	_1	_1	_1	-0.112

 $<sup>^{\</sup>mathbf{a}}$ No chemical mutagen was detected by the 10 microbial strains.

f Analysis not performed on this sample.

b. No rat mortality after 14 days due to maximum dosage of  $10^{-5}~\text{m}^3/\text{kg}$  body weight (LD<sub>50</sub>). However, six samples (B, F, L, N, and S) showed potential body weight effects, and sample R resulted in eye irritation.

Effect was algal growth inhibition.

d Negative sign indicates inhibition in  ${\rm CO_2}$  generation rate compared to a control sample; positive number indicates  ${\rm CO_2}$  stimulation.

e<sub>No acute toxicity.</sub>

Growth inhibition <50% in 100% solution of secondary effluent.

h pH = 9.1 not adjusted before testing.

Sample stimulated algal growth.

 $<sup>\</sup>mathbf{j}_{95\$}$  growth inhibition in 2\$ solution of secondary effluent.

 $k_{\mbox{\scriptsize Diseased}}$  batch of fish nullified this analysis.

Sample inadvertently collected prior to settling pond.

Mo statistical analysis because heavy solids concentration obscured the analysis; the sample did not appear to be acutely toxic.

to determine BATEA, samples will also be collected from the "best" treatment system to evaluate the reduction in acute toxicity. Samples will also be collected to measure removal efficiencies for the 129 priority pollutants. Textile plant intake water samples will also be collected and analyzed for the priority pollutants.

#### SECTION 3

#### RESULTS AND RECOMMENDATIONS

Several results can be noted from the data presented with respect to textile plant secondary effluents.

- 1. None of the secondary effluent samples resulted in a positive mutagenic response or indicated acute toxicity to rats.
- 2. Even though the series of bioassays are unrelated in terms of toxicity mechanisms, the data did provide sufficient information to relatively rank secondary effluents in terms of toxicity and to select those plants for further study under the tertiary treatment technology assessment.
- 3. The effluent sample collected from Plant R was inadvertently collected between the aeration lagoon and settling basin. Therefore the sample tested was that of the aeration lagoon slurry. Note that for bioassays where the sample was first filtered (cytotoxicity and freshwater algae), the toxicity was slight; for unfiltered samples (fathead minnow, daphnia, and soil microcosm) the toxic effects were significant. This result tends to imply that the toxicity for Plant R is associated with the filterable solids. Plant R will be resampled in Phase II of the program to further evaluate the toxicity response.
- 4. In terms of priority pollutants, only 56 of the 114 organic species were detected in either raw waste or secondary effluent sample, with 49 found in the raw waste samples and 46 in the secondary effluent samples. The dominant organic species present in secondary effluent samples include common plasticizer species, such as phthalates, and common raw materials used by the textile manufacturing industry, such as chlorobenzenes.

- 5. Of the 13 priority pollutant metals, beryllium, selenium, and thallium were below analytical detection limits in all raw waste and secondary effluent samples. The dominant metal species detected were arsenic, chromium, copper, and zinc.
- 6. The data indicates that secondary treatment by aeration lagoons and clarifiers produce a significant decrease in phenolic compounds and toluene and a moderate reduction in chlorobenzenes.
- 7. It is difficult to accurately evaluate the treatment efficiencies because for this program raw waste and secondary effluent samples were collected simultaneously. Due to the 1-day to 30-day hydraulic retention time in the treatment plants, secondary effluent samples reflect the waste loading 1-day to 30-day in the past. Also, data are not available to indicate the quantity of organic compounds stripped from the wastewater due to the action of surface aerators.

As a result of performing newly developing bioassay tests, priority pollutant analyses, and Level 1 chemical analyses on environmental samples, several recommendations can be made with respect to improvements. The major recommendations are discussed below:

- For bioassay screening purposes, it proved to be more economical to conduct mutagenicity and rat acute toxicity tests using the maximum dose. If no toxicity was indicated, then the test was completed; when toxicity was detected, then the dose response testing scheme was used.
- 2. Toxicity testing should be performed on filtered and unfiltered samples. Also, due to the detection limits of the mutagenicity, cytotoxicity, and rat acute toxicity tests, consideration should be given to testing concentrated samples.
- 3. Whenever extractions are performed on environmental samples, the percent of extractable organics should be determined. For the Level 1 chemical analysis scheme, potentially more organics can be extracted in methylene chloride if the sample pH is adjusted to pH 10 or 11 as opposed to use of neutral pH. Since the Level 1 chemical analysis scheme requires field extractions, a presurvey and preliminary tests should be performed to determine the extractability of the sample and to identify potential problems.

- 4. For priority pollutant volatiles analysis, MRC obtained better results by cryogenic trapping at -40°C than at room temperature as recommended by EPA. In addition, to reduce interference effects, three internal standards were used as opposed to the one recommended.
- 5. To identify the source of priority pollutants in wastewater samples, samples of the intake water to the industry complex should be analyzed.

#### SECTION 4

#### SCOPE OF WORK

#### BACKGROUND

To understand the nature and purpose of the textile wastewater toxics program it is first necessary to briefly review the events which formed the study's foundation. The principal event occurred on 1 October 1974 when the American Textile Manufacturers Institute (ATMI) filed a petition with the U.S. Fourth Circuit Court of Appeals asking for review of the 1983 effluent guidelines proposed for the textile industry. ATMI's grounds for the suit were that the best available technology economically achievable (BATEA) had not been demonstrated for the textile industry. As a result, ATMI and EPA filed a joint motion for delay of the petition, stating that additional information would be developed through a cooperative study by ATMI and EPA (IERL/RTP).

The objective of this ATMI/EPA Grant Study was to gather enough technical and economic data to determine what is the BATEA for removing criteria pollutants from textile wastewaters. Criteria pollutants for the textile industry include 5-day biochemical oxygen demand (BOD $_5$ ), chemical oxygen demand (COD), color, sulfide, pH, chromium, phenol, and total suspended solids (TSS). On 3 January 1975 the court instructed ATMI and EPA to proceed as promptly as feasible to a completion and review of the study.

The ATMI/EPA Grant Study was divided into two phases: Phase I, to determine the best available technology, and Phase II, to determine which technology(s) was economically achievable. A generalized program outline of Phase I is shown in Figure 1. To evaluate the best available technology, two mobile pilot plants were constructed by ATMI. This strategy allowed for real-time, flowthrough treatment studies. Each pilot plant contained four tertiary wastewater treatment unit operations; one was scheduled to visit 12 textile plants and the other to visit 11. Two additional tertiary treatment technologies were laboratory tested.

Treatment operations in each mobile unit include a reactor/clarifier (using combinations of alum, lime, ferric chloride, and anionic and cationic polyelectrolytes), two multimedia filters, three granular activated carbon columns, ozonation and dissolved air flotation. Powdered activated carbon treatability tests

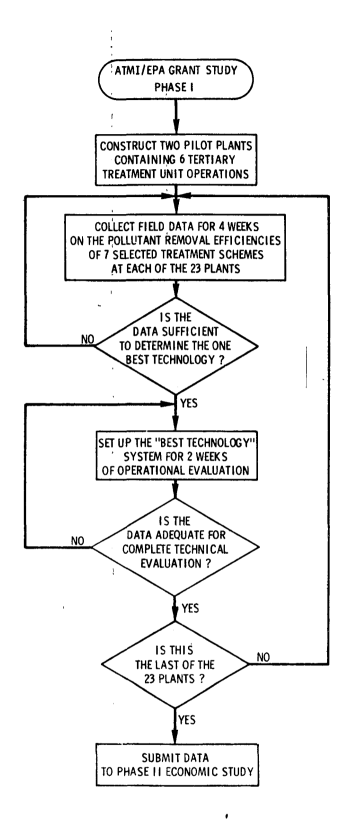


Figure 1. Program outline for Phase I: technology assessment for the ATMI/EPA Grant Study.

were performed in the laboratory instead of in the field with the pilot plant. Using these six unit operations ATMI and EPA selected seven treatment systems for evaluation (Figure 2).

MODE A: REACTOR / CLARIFIER --- MULTIMEDIA FILTER

MODE B: MULTIMEDIA FILTER --- GRANULAR ACTIVATED CARBON COLUMNS

MODE C: MULTIMEDIA FILTER --- OZONATOR

MODE D: OZONATOR

MODE E: REACTOR / CLARIFIER --- MULTIMEDIA FILTER --- GRANULAR ACTIVATED

(OPTIONAL) CARBON → OZONATOR

MODE F: COAGULATION -- MULTIMEDIA FILTER

MODE G: DISSOLVED AIR FLOTATION

Figure 2. Seven tertiary treatment modes for "best available technology" evaluation.

Each of the seven treatability systems was to be set up, and operational and pollutant data were to be collected over a 2-day to 3-day period. Based on that data, the "best" system was to be selected and set up for 2 weeks of operational evaluation. These data were then to be forwarded to Phase II for economic evaluation.

The second event that formed the foundation for this project occurred when a group of environmental action organizations filed a class action suit against EPA (Natural Resources Defense Council et al. v. Train, U.S. District Court of Washington, D.C.) for not developing and promulgating regulations establishing effluent limitations and guidelines and new source performance standards for 21 industrial point sources, including the textile industry. As a result, on 7 June 1976 the court issued a consent decree requiring EPA to enhance development of effluent standards.

The most significant result from the court mandate was that it focused federal water pollution control efforts on potentially toxic and hazardous pollutants. The original consent decree required that 38 classes of chemical compounds (Table 6) be analyzed in wastewater samples. Recognizing the difficulty of analyzing for all chemical species present in each category of compounds, EPA developed a list of 129 specific compounds (Appendix A) representative of the classes of compounds listed in the consent decree. These compounds are referred to as the consent decree priority pollutants, or priority pollutants for short.

Acenaphthene Acrolein Acrylonitrile Aldrin/Dieldrin Antimony and compounds Arsenic and compounds Asbestos Benzene Benzidine Beryllium and compounds Cadmium and compounds Carbon tetrachloride Chlordane (technical mixture and metabolites) Chlorinated benzenes (other than dichlorobenzenes) Chlorinated ethanes (including 1,2-dichloroethane, 1,1,1-trichloroethane, and hexachloroethane) Chloroalkyl ethers (chloromethyl, chloroethyl, and mixed ethers) Chlorinated naphthalene Chlorinated phenols (other than those listed elsewhere; includes trichlorophenols and chlorinated cresols) Chloroform

2-Chlorophenol Chromium and compounds Copper and compounds Cyanides DDT and metabolites Dichlorobenzenes (1,2-,1,3-, and 1,4-dichlorobenzenes) Dichlorobenzidine Polychlorinated biphenyls (PCB) Polynuclear aromatic hydrocarbons (including benzanthracenes, benzopyrenes, benzofluoranthene, chrysenes, dibenzanthracenes, and indenopyrenes) Selenium and compounds Silver and compounds 2,3,7,8-Tetrachlorodibenzo-pdioxin (TCDD) Tetrachloroethylene Thallium and compounds Toluene Toxaphene Trichloroethylene Vinyl chloride Zinc and compounds

EPA also developed a sampling and analytical procedures manual to be used as a laboratory guide for the analysis of priority pollutants (1). The analytical methods recommended by EPA are still in the developmental phase and require further verification and validation.

Therefore, in addition to evaluating the removal of criteria pollutants by tertiary treatment technologies, EPA was charged with the task of evaluating the removals of toxicity and priority pollutants by the treatment systems.

<sup>(1)</sup> Draft Final Report: Sampling and Analysis Procedures for Screening of Industrial Effluents for Priority Pollutants. U.S. Environmental Protection Agency, Cincinnati, Ohio, March 1977. 145 pp.

The final event which influenced the formation of the present program was the three-phase sampling and analytical strategy for environmental assessment developed by EPA, Process Measurements Branch, IERL/RTP. The purpose of the assessment procedure was to determine in a stepwise and cost-effective manner all chemical species being discharged to the environment from a point source. Level 1, the first part of the three-phase approach, is designed to focus available resources on emissions that have a high potential for causing measurable health or ecological effects.

The second phase, Level 2, has as its goal the identification and quantification of specific compounds. Level 3 is designed to continuously monitor indicator compounds as surrogates for a large number of specific pollutants. At the start of this textile project, only Level 1 analytical and biological procedures were available (2).

In addition to chemical analyses, the Level 1 recommended protocol included bioassay testing procedures for evaluating toxicity removal by control technologies (3). Bioassays are required to provide direct evidence of complex biological effects such as synergism, antagonism, and bioavailability.

#### PROGRAM OBJECTIVE

The fundamental objective of the textile wastewaters program conducted by MRC in conjunction with the EPA is to determine the reduction in toxicity and priority pollutant concentrations achieved by the tertiary treatment technologies under investigation in the ATMI/EPA Grant Study. The latter study focuses directly on the treatability of criteria pollutants. Thus, the overall EPA-IERL/RTP textile program consists of two separate projects, each with different activities, running parallel in time, but converging towards the same goal: determination of the best available technology economically achievable for removing textile wastewaters (Figure 3).

To evaluate the reduction in toxicity in a cost-effective manner for the MRC/EPA project, a two-phase approach was developed. Phase I was designed to collect baseline toxicity data on

<sup>(2)</sup> Hamersma, J. W., S. L. Reynolds, and R. F. Maddalone. IERL-RTP Procedures Manual: Level 1 Environmental Assessment. EPA-600/2-76-160a (PB 257 850), U.S. Environmental Protection Agency, Research Triangle Park, North Carolina, June 1976. 147 pp.

<sup>(3)</sup> Duke, K. M., M. E. Davis, and A. J. Dennis. IERL-RTP Procedures Manual: Level 1 Environmental Assessment Biological Tests for Pilot Studies. EPA-600/7-77-043 (PB 268 484), U.S. Environmental Protection Agency, Research Triangle Park, North Carolina, April 1977. 114 pp.

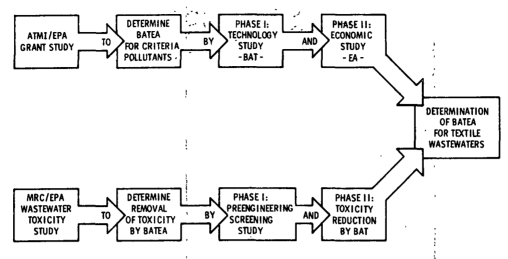


Figure 3. Overall program approach to determine BATEA.

secondary effluents from 23 selected textile plants and to rank the plants in descending order of toxicity (Figure 4). Phase II was designed to determine the level of toxicity removal attained by the tertiary treatment systems in the ATMI/EPA Grant Study at only those plants with relatively high secondary effluent toxicity (Figure 5). Sampling locations for Phase II of the study are shown in Figure 6, and the strategy used for evaluating control technologies in terms of toxicity removal is illustrated in Figure 7.

#### PROJECT ORGANIZATION

The major effort of the Phase I MRC/EPA screening study was devoted to the collection, chemical analysis, and biological toxicity testing of single, 8-hr composited wastewater samples from the 23 textile plants scheduled for testing in the ATMI/EPA Grant Study. In addition, samples were collected from nine other textile plants for chemical analyses only. Wastewater characterization data were therefore assembled for a total of 32 plants.

The scope of work for Phase I was divided into three separate task areas, each based on different EPA data requirements, as shown in Figure 8. CPB (Chemical Processes Branch, IERL/RTP, Project Officer, M. Samfield) requested chemical and bioassay data on secondary effluent samples from the 23 textile plants scheduled to be studied in the ATMI/EPA Grant Study. These data were used to characterize and compare the relative toxicities of the plant effluents tested. EGD (Effluent Guidelines Division, EPA, Washington, J. D. Gallup) requested chemical analyses of the raw waste streams entering the 23 wastewater treatment plants, as well as chemical analyses of the raw waste and secondary effluent streams at 9 additional textile plants. Raw waste and effluent data were needed to evaluate the pollutant removal efficiencies

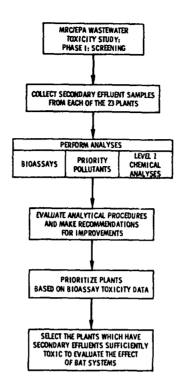


Figure 4. Program outline for Phase I of the MRC wastewater toxicity study.

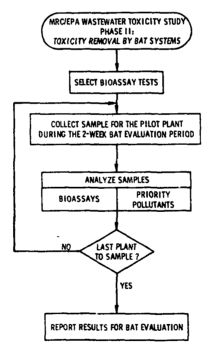


Figure 5. Program outline for Phase II of the MRC wastewater toxicity study.

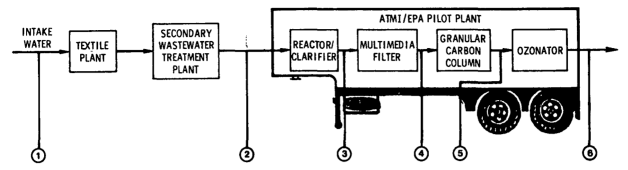


Figure 6. Sampling locations for Phase II of the MRC wastewater toxicity study.

# INTERPRETATION OF BIOASSAY TEST RESULTS

Bioassay Results		Toxic Substance Interpretation
Inlet	Outlet	
+	+	Control Technology Is Not Effective
+	-	Control Technology Is Effective
-	+	Control Technology Is Deterimental
-	-	Control Technology Is Not Deterimental

Figure 7. Interretation of bioassay test results.

of current state-of-the-art secondary treatment systems. In order to detect other possible pollutant species, PMB (Process Measurements Branch, IERL/RTP, L. D. Johnson) requested that Level 1 chemical characterization also be performed on the effluent samples at the basic 23 textile plants (2). Since these data were requested after the program began, only 15 textile plants were sampled for Level 1 chemical characterization.

Chemical characterization of wastewater samples involves four categories of analysis:

- 129 consent decree priority pollutants analysis (1)
- nutrient analysis (4, 5)
- effluent quidelines criteria pollutants analysis (4, 5)
- Level 1 chemical characterization (2)

<sup>(4)</sup> Manual of Methods for Chemical Analysis of Water and Wastes. EPA-625/6-76-003a (PB 259 973), U.S. Environmental Protection Agency, Cincinnati, Ohio, 1976. 317 pp.

<sup>(5)</sup> Standard Methods for the Examination of Water and Wastewater, Fourteenth Edition. American Public Health Association, Washington, D.C., 1976. 874 pp.

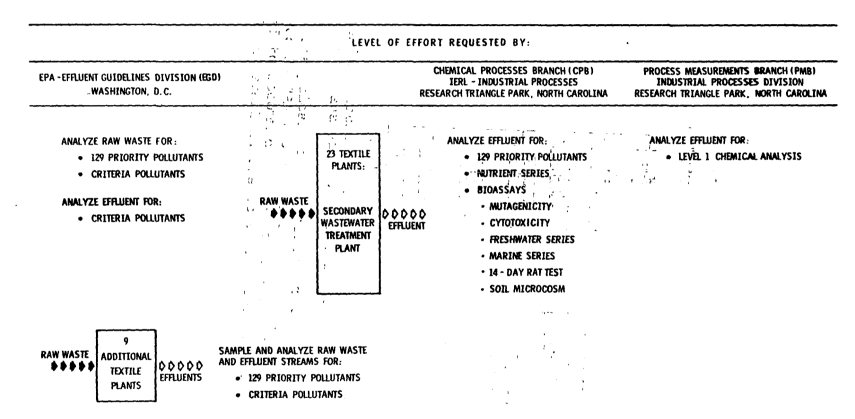


Figure 8. Scope of work for the analysis of textile plant wastewaters.

The 129 consent decree priority pollutants as listed in Appendix A are divided into volatile compounds, nonvolatile compounds, and metals (see Appendix B). The nutrient series required to support algal tests includes analysis of nitrite, nitrate, ammonia, total Kjeldahl nitrogen (TKN), o-phosphate, phosphorus, and total organic carbon (TOC). Effluent guidelines criteria pollutants include 5-day biochemical oxygen demand (BOD<sub>5</sub>), chemical oxygen demand (COD), sulfide, color, pH, total suspended solids (TSS), total dissolved solids (TDS), and total phenol. A detailed description of Level 1 chemical characterization is given in Section 6.

The bioassay scheme established by EPA for evaluating the reduction in toxicity of water samples by control technologies is shown in Figure 9. All the tests shown were used in this project. The marine ecology series and the soil microcosm tests were requested after the project began, therefore data were obtained from 15 textile plants as opposed to the basic 23 plants.

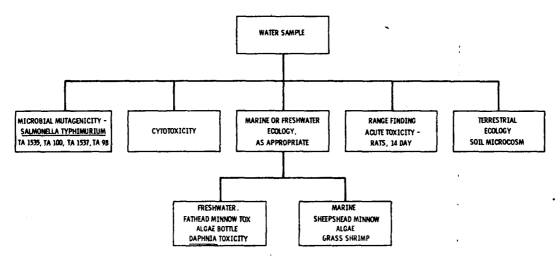


Figure 9. EPA-recommended bioassay testing scheme for toxicity analysis of water samples.

Figure 10 illustrates the distribution of samples among the eight EPA and private laboratories that performed the chemical analyses and bioassay tests. Appendix C lists the names and addresses of all persons involved in the textile project.

MRC collected raw wastes and effluent samples at 23 of the ATMI/EPA-designated plants. Wastewater samples were collected by EPA-Environmental Research Laboratory (ERL) (Athens, Georgia) at two of the additional textile plants and sent to MRC for chemical analysis. Sverdrup and Parcel and Associates, Inc., (St. Louis, Missouri) collected the remaining samples at the additional seven plants and sent them to MRC for chemical analysis.

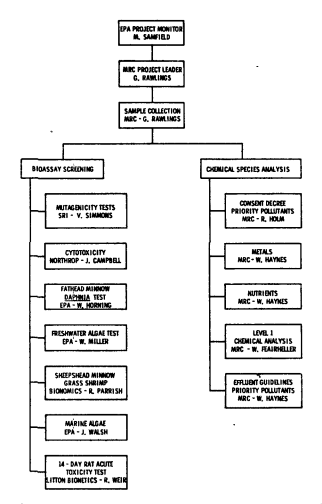


Figure 10. Laboratories and persons involved in sample analysis of textile plant effluents.

Sampling, analytical, and bioassay procedures followed those recommended in EPA reports (1-5). All procedural modifications instituted to accommodate the textile wastewater samples are discussed in detail in the remaining sections of this report.

#### SECTION 5

#### SAMPLING PROCEDURES

#### COLLECTION TECHNIQUE

Wastewater was collected by composite and grab sampling techniques. Composite samplers (Isco Model 1680) were used to collect raw waste samples for analysis of nonvolatile organics and metals.

Tygon® sample tubing used was washed with detergent, rinsed thoroughly, and given a final washing with organic-free water. A 0.001-m³ sample blank was then collected and analyzed for organic leachates. Organic-free water was prepared by passing water, distilled in glass, through a 0.6-m-long activated carbon column. The blank was collected in glass, sealed with a Teflon®-lined cap, and stored in ice at 4°C until analyzed.

Grab sampling techniques were used to collect raw waste samples for other analyses, and for all secondary effluent samples, Figure 11. Eight individual grab samples were collected at equally spaced time intervals during the normal working day. To insure that each of the eight laboratories received a sufficient portion of the same sample, grab samples were collected in a Teflon-lined, 0.01-m³ stainless steel bucket. A specified aliquot was transferred to each of the sample bottles from this container. Care was taken to insure that the sample remained homogeneous throughout each of the 10-min pouring sessions. Containers for volatile organics analysis were collected and sealed first to minimize possible evaporation losses.

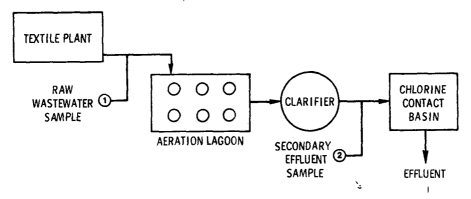


Figure 11. Phase I sampling locations.

TABLE 7	COLLECTION	AND HAN	NDLING REQUIREMENTS (	4)
Analysis	Total sample of vol required,			storage emper- ature,
		61 2		
Consent decree pollutants Volatile organics	320 x 10-6	Glass	_c ()	4 <sup>d</sup>
Nonvolatile organics	4'000 X TO .	GIASS		4
Metals C	1,000 x 10 %	Plastic	Nitric acid $5 \times 10^{-3} \text{ m}^3/\text{m}^3$	4
Nutrient analysis	to hay it is	3	ថ្ងៃ គ្រឹ	
Total organic carbon	25 x 10-6	Glass	Sulfuric acid to pH < 2	4
Ammonia C	400 x 10 <sup>-6</sup>	Glass g	Sulfuric acid to pH <, 2	4
Total Kjeldahl nitrogen	500 x 10 <sup>-6</sup>	,		4
Nitrate	100 x 10 <sup>-6</sup>	Glass g	Sulfuric acid to pH < 2	4
Nitrite (*) Orthophosphate	50 x 10-6		ė –	
Total phosphorus	50 x 10-6	Glass		
्ट्री अ । ७	, ,		· 在音音文化的	
Criteria pollutants	"			
Biochemical 5-day	$1,000 \times 10^{-6}$	Glass		Δ.
oxygen; demand Chemical Coxygen demand	50 x 10 <sup>-6</sup>	Glass 4	Sulfuric acid to pH <2	4
Color	50 x 10 <sup>-6</sup>	Glass 4		4
Sulfide .	500 x 10 <sup>-6</sup>	Glass	$2 \times 10^{-6} \text{ m}^3$ zinc acetate	4
Total suspended solids	100 x 10 <sup>-6</sup>			
Phenol	500 x 10 <sup>-6</sup>	Glass	Phosphoric acid to pH' < 4, 10 <sup>3</sup> g copper sulfate/m <sup>3</sup>	4
Cyanide 및 기를 위 기	500 x 10-6	Plastic	Sodium hydroxide to pH > 12	4
Cyanide Cyanide	- C	[P. 13]		
Freshwater ecology		iš,	į	
Fathead minnow and Daphnia	60,000-x-10-6	. Glass		A
Algae	$9,600 \times 10^{-6}$	' Plastic	•	4
3 138	, , , , , , , , , , , , , , , , , , , ,	1 C		
Marine ecology		17 28		
Sheepshead minnow	60 000 - 10-6	03		
and grass shrimp	60,000 x 10 <sup>-6</sup> 2,000 x 10 <sup>-6</sup>	Plastic	to the contract of the contract of	4. A
Riyae	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1., 1 145020	· ·	•
Soil microcosm	100-x 10-6	Plastic	1 2 2 2 2 2	4
Ames test	1250 x 10-6	Glass	h dage a	4
Cytotoxicity tests 14-day Rat test	250 x 10-6 1,000 x 10-6		<b>1</b> :	4
O F	1,000 x 10	Glass	1	4
a	6 3	<del>                                     </del>		
Eight individual 40 x 10	m³ samples		0 0 0 0 0 0	
blo-6 m3 equals 1 ml	10 miles		0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	
CBlanks indicate no chemic	al preservation	required.		•
			3 -3 -5 200 -2 24 200 -	
sulfate is added to each	of the 4 x 10-5	.e, 3 X IU	<sup>3</sup> m <sup>3</sup> of 10% solution of sodium vials to neutralize the resident	n thio-
chlorine; otherwise, abno	rmally high tri	halomethan	e_values may result.	Jual
	1,0		de d	
	. '	,		

An additional  $0.02~\text{m}^3$  were required to properly apportion samples into the six  $0.02\text{-m}^3$  bottles for the marine and freshwater ecology tests. Samples were poured into these bottles at the end of each of the eight grab sampling sessions. There was no visible change in flow rate during each of the 15-min sample collection periods. Fluctuations in effluent and raw waste stream flow rates were usually on a 45-min to 1-hr time schedule.

# SAMPLE CONTAINER PREPARATION

All glass containers, except the 0.02-m³ bottles, were thoroughly cleaned with strong acid (50% sulfuric acid + 50% nitric acid), rinsed, and heated in a glass annealing oven at 400°C for at least 30 min. The 0.02-m³ bottles were detergent washed and thoroughly rinsed. The glass containers used to store samples for mutagenicity testing were rinsed with acetone. The rest of the glass containers were rinsed with methylene chloride and dried in the oven at 100°C. All glass bottles had Teflon-lined caps.

Plastic sample containers were thoroughly cleaned before use. Each bottle was washed with detergent and tap water, then rinsed with 1:1 nitric acid/tap water, 1:1 hydrochloric acid/tap water, and, finally, deionized distilled water (1).

#### SAMPLING LOGISTICS

The type and volume of sample container varied, depending on the analysis to be made. Some samples required the addition of chemical preservatives in the field to prevent deterioration during shipment to the laboratory. The volume of sample required, the container used to hold the sample, and the preservation steps used in this project are shown in Table 7 (4).

A field sampling instructional worksheet was designed to facilitate the arduous task of filling 37 glass and plastic bottles of different sizes requiring different sample volumes and preservatives at each plant. Table 8 shows part of this worksheet.

Each sampling day, before sampling, bottle labels were filled out and affixed to the appropriate sample bottles. Figure 12 shows the bottle label that MRC has designed for sample identification.

TABLE 8. PORTION OF THE FIELD INSTRUCTIONAL FORM USED BY MRC
TO ASSURE ACCURATE SAMPLE COLLECTION AND PRESERVATION

Am				Marine	TOC, NH3, TKN		anics		
te	Sulfide	Phenol	Cyanide	algae	COD, nitrate	Metals	Nonvolatile	Volatile	Type of test
				Gulf					Sample
S	MRC	MRC	MRC_	Breeze	MRC	MRC	MRC	MRC	destination
									Bottle
	8	7	6	5		3	2	,	dentification number
	<del></del>				<del></del>				Number of
	1	1	1	1	1	1	1	8	bottles
	<del></del>	<del></del>			<del></del>				DOCCIES
	1	1	<b>1</b> .						
									ype of bottle
2 -	62	62	240	240	240	240	3,785	40	Sample size 10 <sup>-6</sup> m <sup>3</sup>
			preservation	for sample	Chemicals added				
· No	+2 x 10 <sup>-6</sup> m <sup>3</sup> Zinc acetate	pH <4 with phosphoric acid	+4 x 10 <sup>-6</sup> m <sup>3</sup> Sodium hydroxide, check with potassium iodide paper	None	+Sulfuric acid, pH <2	+5 x 10 <sup>-6</sup> m <sup>3</sup> Nitric acid	None	+3 x 10 <sup>-6</sup> m <sup>3</sup> Sodium thiosulfate, seal	ample number 1
No.	None	0.5 g copper sulfate	None	None	Same as No. 1	None	None	Same as No. 1	2
	None	None	None	None	Same as No. 1	None	None	Same as No. 1	3
Sa	None	Same as No. 1	None	None	Same as No. 1	None	None	Same as No. 1	4
No	None	None	None	None	Same as No. 1	None	None	Same as No. 1	5
No	None	None	None	None	Same as No. 1	None	None	Same as No. 1	6
No	None	None	None	None	Same as No. 1	None	None	Same as No. 1	7
	Seal	Same as No. 1	pH > 12, check seal	None	Same as No. 1	Seal	Seal	Same as No. 1	8

7

Job		
Sample Location		
Type of Sample		
Analyze for		
Preservation	·	
Comments		
	·	
	/	
Log No	Date	
Initials		

Figure 12. MRC sample bottle label.

#### SAMPLE SHIPPING PROCEDURES

Each bottle was capped and sealed with tape to prevent leakage. Glass bottles were individually wrapped to prevent breakage.

Sample bottles were then packed in one-piece, molded, styrene foam shipping cartons with 3.8-cm walls and fitted tops. Each such unit was then placed in a corrugated cardboard box.

Each carton was half-filled with sample bottles, filled with ice, sealed with cellophane tape, and reinforced with 0.05-m duct tape.

Address labels were affixed to box tops. Warning labels--"This carton contains glass and ice"--"Hold at airport and call \_\_\_\_" messages were also put on the box tops.

All samples were shipped by conventional air freight on the day that they were collected. The airlines selected offered the most direct route without carrier changes.

#### SECTION 6

#### WASTEWATER CHEMICAL ANALYSES

#### EFFLUENT GUIDELINES CRITERIA POLLUTANTS

Parameters determined under the category of effluent guidelines criteria pollutants were: 5-day biochemical oxygen demand (BOD $_5$ ), chemical oxygen demand (COD), color, sulfides, total suspended solids (TSS), pH, and total phenol. As sample shipments arrived at MRC, they were logged in and distributed to the designated technicians for analysis.

Analytical and supporting procedures followed those described in References 4 and 5.

Criteria pollutants were determined on the raw waste and secondary effluent streams from the basic 23 plants and the additional 9 plants.

Results of the chemical analyses are given in Table 9. The first row of numbers for each plant represents data obtained on the raw waste stream, and the second row of numbers corresponds to the wastewater treatment plant effluent stream. All values except color and pH are given in  $g/m^3$  (ppm). Color was measured using the American Public Health Association (APHA) system.

Effluent values obtained from wastewater treatment facilities in some plants were greater than those of the influent raw waste. This occurred, in part, because the wastewater entered the treatment system 1 day to 5 days prior to leaving the treatment plant. The hydraulic retention time in textile wastewater treatment plants ranged from 1 day to 30 days, with an average value of 5 days.

All of the textile plants sampled had a secondary wastewater treatment facility that included a lagoon with several surface aerators, followed by a clarifier. Several plants used equalization basins prior to the aerated lagoons. Effluent samples were collected between the clarifier and the finishing pond in plants that had both. The two exceptions were plant Y, where the sample was taken after the finishing pond, and plant R, where the effluent sample was inadvertently collected between the aerated lagoon and the settling basin. All other plant effluent samples were collected after the clarifiers.

TABLE 9. ANALYSIS OF WASTEWATER SAMPLES FOR EFFLUENT GUIDELINES CRITERIA POLLUTANTS

	Criteria pollutant, g/m³							
Plant code	5-Day biochemical oxygen demand	Chemical oxygen demand	Color, APHA	Sulfide	Phenol	Total suspended solids	рн	Cyanide
/raw waste	459	1,735	2,000	6.0	0.092	165	10.7	<0.004
√effluent	168	1,652	2,000	4.0	0.065	228	7.3	0.015
3/raw waste	1,050	1,264	1,400	1.4	0.042	32	10.5	0.017
3/effluent	<5	99	90	0.2	0.015	8	7.5	<0.004
C/raw waste	445	802	2,600	5.2	0.074	49	11.2	0.007
c/effluent	25	396	1,920	5.0	0.088	300	10	0.013
/raw waste	71	224	1,875	<0.02	0.024	16	10 ,	0.21
/effluent	6.6	64	1,625	2.8	0.018	154	7.2	0.21
/raw waste	18	2,660	250	<1	0.069	52	10	<0.004
E/effluent	<5	78	30	<1	0.014	19	7.2	<0.004
/raw waste	194	583	150	2.1	0.74	23	9.2	<0.004
effluent	69	276	80	0.1	0.028	44 '	7.4	<0.004
/raw waste	203	1,340	300	<1	0.028	37	11	<0.004
/effluent	42	502	300	<sup> </sup> <1	0.054	6	7.5	0.006
i/raw waste	288	320	1,250	<0.02	0.047	39	10	<0.004
i/effluent	14	300	500	<0.02	0.019	43 🕴	7.6	<0.004
/raw.waste	210	810	1,875	1.8	0.063	0.01	11	< 0.004
/effluent	25	376	1,375	1.8	0.024	0.023	7.8	< 0.004
/raw waste	564	1,725	40,000	<1	0.067	69	10	< 0.004
K/effluent	<5	131	150	<1	0.018	21	7.2	< 0.004
/raw waste	379	1,117	1,300	4.5	0.038	19	7.4	< 0.004
/effluent	13	234	370	3.0	0.026	78	5.8	0.172
l/raw waste	830	2,265	1,000	<1	0.037	210	11	< 0.004
M/effluent	< 5	255	500	<1	0.025	21	7.5	< 0.004
/raw waste	334	1,140	1,050	1.1	0.156	68	9.2	< 0.004
/effluent	36	286	90	0.1	0.068	. 77	7.0	< 0.004
/raw waste	680	172	300	6.19	0.228	6	10	0.19
/effluent	28	45	250	6.09	0.032	45	7.1	0.14
/raw wastę	450	1,692	1,500	<1	0.282	87	10	< 0.004
/effluent <sup>a</sup>	70	830	2,000	<1	0.162	225	8.1	< 0.004
/raw waste	219	559	250	9.2	0.107	<b>25</b> '	10	0.007
effluent	59	1,035	75	<1	0.029	581	7.8	< 0.004

TABLE 9 (continued)

•	5-Day			eria pollu				
Plant code	biochemical oxygen demand	Chemical oxygen demand	Color,	Sulfide	Phenol	Total suspended solids	рΗ	Cyanide
Flant Code	demand	demand	AFAR	Bulline	FILEHOL	SULLUB	<u>pn</u>	Cyanius
T/raw waste	501	500	1,345	7.6	0.073	28 🗸		<0.004
T/effluent	32	414	350	6.0	0.041	35	7.4	<0.004
U/raw waste	400	1,464	3,200	5.6	0.057	111	10	<0.004
U/effluent	24	748	2,480	3.5	0.007	92	7.3	0.212
V/raw waste	53	b	500	<1	0.018	54	9.0	0.006
V/effluent	< 5	128	500	<1	0.016	26	7.1	0.018
W/raw waste	1,920	6,124	2,200	0.5	0.670	2,300	10.4	0.015
W/effluent	84	837	1,900	0.1	0.232	300	8.1	0.020
X/raw waste	237	786	1.200	0.75	0.940	24	10.2	<0.004
X/effluent	15	258	10	0.01	0.035	18	7.2	0.101
Y/raw waste	122	457	10,000	<1	0.064	33	10.5	<0.004
Y/effluent	< 5	115	250	<1	0.022	17	8.0	<0.004
2/raw waste	351	812	500	2.48	0.56	20	10	<0.004
Z/effluent	< 5	105	750	<1	0.023	13 💡	8	<0.004
JJ/raw waste	b	1,545	b	<1	0.144	ъ	b	0.005
JJ/effluent	b	510	p	<1	0.055	b	b	0.028
KK/raw waste	ь	1,955	ь	<1	0.150	ь	b	<0.004
KK/effluent	ъ	447	p	<1	0.052	b	b	<0.004
LL/raw waste	b	727	ь	<1	0.001	ъ	b	0.008
LL/effluent	þ	155	þ	<1	0.094	þ	þ	0.006
MM-1	ь	ь	b	<1	0.033	ь	ь	<0.004
MM-2	b	þ	b	<1	0.031	b	b	<0.004
MM-3	ъ	р	þ	<1	0.036	b	ь	<0.004
MM-4	b	b	Ъ	<1	0.039	ь	р	<0.004
NN/raw waste	b	938	b	<1	0.043	b	b	0.04
NN/effluent	b	236	ь	<1	0.014	b	Þ	<0.004
00/raw waste	ь	1,889	b	<1	0.082	ъ	b	<0.004
00/effluent	þ	635	ь	<1	0.026	ь	ь	<0.004
PPC	p	339	b	<1	0.044	þ	ь	<0.004
Y-001/raw waste	ъ	b	b	b	b	þ	b	< 0.004
Y-001/effluent	ь	Þ	b	b	þ	Þ	b	0.029
C-001 <sup>C</sup>	b	b	ь	ь	b	b	b	< 0.004

Secondary effluent sample was inadvertently collected between the aerated lagoon and settling ponds.

b Analysis not performed on sample.

C Secondary effluent sample only.

### ANALYSIS PROTOCOL FOR THE 129 CONSENT DECREE PRIORITY POLLUTANTS

Recommended analytical procedures (1) developed by EPA were used throughout this project. It is important to realize that these EPA procedures are still under development and require further verification and validation. Therefore, the data presented in this section only serve to identify which of the 129 chemical species are present and to indicate the general concentration ranges within an order of magnitude.

Adaptations of these procedures to accommodate the special requirements of textile wastewaters and/or any ambiguities in analytical techniques are discussed below. Three chemical species were not determined in this project: endrin aldehyde, 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), and asbestos. EPA-Environmental Monitoring and Support Laboratory (EMSL) recommended that TCDD should be omitted because of its extreme toxicity, and the potential health hazard involved in preparing standard solutions from the pure compound. Pure endrin aldehyde could not be obtained in time to prepare standard solutions. Asbestos was eliminated, as recommended by EPA-IERL-RTP and EPA-EGD, due to the presence of other fibrous materials in textile wastewaters.

The analytical protocol (2) divides the 129 chemical species into three basic categories: volatile organics, nonvolatile organics, and metals. Appendix B lists the chemical species which belong to each category. The following sections outline the analytical procedures and MRC modifications for each category.

## Volatile Organics

The recommended analytical method was designed to determine those chemical species which are amenable to the Bellar purge and trap method. Eight 40 x  $10^{-6}$ -m<sup>3</sup>, hermetically sealed glass vials, stored in ice, were sent to the laboratory from each sampling The vials were composited within 1 day of receipt at the laboratory. Two vials of composite solution were sealed and retained at  $4^{\circ}$ C as reserve samples. Volatiles from 5 x  $10^{-6}$ -m<sup>3</sup> samples of composite solution were sparged with helium onto two Tenax GC-silica-packed sample tubes. (Internal standards were added to the solutions in the later stages of the program. The majority of the samples had been sparged and stored before the protocol (1) was received and appropriate internal standard could be obtained.) The second Tenax tube was used as a backup sample. Tenax tubes were sealed under a nitrogen atmosphere in glass tubes and stored in a freezer at -18°C until analyzed.

Analyses were carried out using a Hewlett Packard 5981 GC-Mass Spectrometer with 5934 Data System. Sample tubes were heated to 180°C over a 1-min period and held at that temperature for 4 min to desorb the compounds onto a Carbowax 1500 column held at -40°C. Cryogenic trapping at -40°C (liquid nitrogen cooling) gave better

reproducibility of retention time than using the suggested temperature of 30°C, for compounds with boiling points below room temperature. After desorption, the GC column temperature was raised 8°C/min to 170°C.

The mass spectrometric analysis method involves fragmentation of molecules using electron bombardment (70 eV). Masses and relative intensities of the most characteristic molecular fragments for each compound are listed in the protocol (1). The population of ion fragments covering the mass range from 35 atomic mass units to 500 atomic mass units was measured every 6 s, and the data were stored on magnetic tape.

These data allow the operator to reconstruct chromatograms of observed intensity for an individual mass during the course of the scanning. Specific molecules may be detected in the presence of other compounds by examining the reconstructed intensity time plots of their characteristic masses.

Qualitative identification of a compound was made using the three criteria listed in the protocol: 1) retention time must coincide with known retention times, 2) the three characteristic masses must elute simultaneously, and 3) intensities of the characteristic masses must stand in the known proper proportions.

Quantitation of volatile organics was initially made using peak area counts and concentration calibration curves. Later in the program, response ratios using the 1,4-dichlorobutane internal standard were used in quantifying the concentrations. Base/neutral and acid organic compounds were quantified using deuterated anthracene and response ratios as prescribed in the protocol (1).

Figure 13 is a simplified diagram of the analytical scheme for volatile organic analysis.

# Nonvolatile Organics

This method determined the nonvolatile solvent-extractable organic compounds that could be analyzed by gas chromatographic methods. The 129 consent decree priority pollutants contain 81 organic compounds classified as nonvolatile organics.

Nonvolatile organics are divided into three groups: base/neutral fraction, acid fraction (phenols), and pesticides and polychlorinated biphenyls (PCB). A list of compounds that are classified as nonvolatile organics is given in Appendix B.

The analytical procedure is described in Reference 1. Figure 14 depicts the sample processing scheme for the base/neutral and acid fractions. The sample solution,  $2 \times 10^{-3}$ -m<sup>3</sup>, was made alkaline (pH greater than 11) with sodium hydroxide, and then

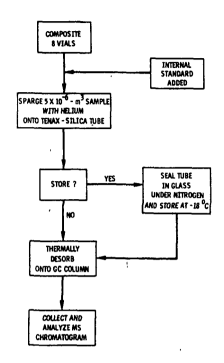


Figure 13. Analytical scheme for volatile organics analysis.

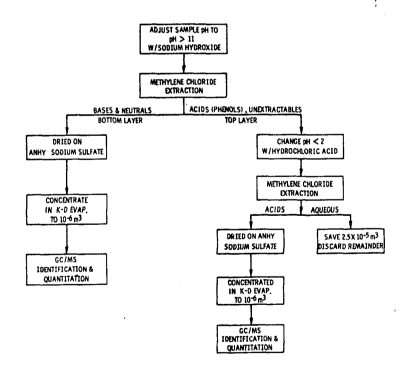


Figure 14. Sample processing scheme for nonvolatile organics analysis.

extracted three times with methylene chloride. Textile raw waste and effluent samples formed strong emulsions upon extraction with methylene chloride. The problem was resolved by drawing off small amounts of separated solvent and pouring the extract through the sample in the separatory funnel. Separation was also enhanced by slowly dripping the emulsion onto the wall of a slightly tilted flask. This approach gives better separation by providing a greater surface area for the solvent and water fractions. Some samples required centrifugation at 1,500 rpm for 1 hr to break the emulsion.

Extracts were dried on a column of anhydrous sodium sulfate, concentrated to  $10^{-6}$  m<sup>3</sup> in a Kuderna-Danish (K-D) evaporator with a Snyder column spiked with deuterated anthracene, sealed in septum capped vials, and stored at 4°C until analyzed. Analyses were performed on the GC/MS system using SP 2250 and Tenax GC columns for base/neutral and acid samples, respectively (1).

A separate 0.001-m<sup>3</sup> sample was used for analysis of the pesticides and PCB (Aroclor® fluids). The basic sample processing scheme is shown in Figure 15. These compounds were extracted with a 15% methylene chloride and 85% hexane solvent mixture. The aqueous phase was discarded, and the organic phase was analyzed by GC with an electron capture detector. Where necessary, acetonitrile partitioning and a Florisil® chromatography column were used for further celanup of the sample. In 85% of the samples, additional cleanup was not required.

Confirmation of identify and quantitation were made using two different GC columns: SP-2550 and Dexil 410. Compound verification was made with the MS when the concentration was greater than 0.01 g/m³. Concentrations of potential pesticides ranged from 0.0001 g/m³ to 0.01 g/m³; therefore, MS verification was not possible in this study. Pesticide species identified only by GC below 0.01 g/m³ were reported only if they met the following two criteria: 1) the retention time window between standards and unknown peaks correlated within  $\pm 3$  s, and 2) concentrations calculated from both GC columns had to agree within  $\pm 20$ %. Unknown peaks not meeting these criteria were assumed not to be the pesticide species.

#### Metals

In addition to the volatile and nonvolatile organics, the 129 chemical species include 13 metals, asbestos, and cyanide. Each metal is measured as the total metal. Asbestos was not determined in this study; cyanide was measured by conventional wet chemistry techniques (5).

Eight metals were analyzed by the inductively coupled argon plasma (ICAP) excitation technique: antimony, cadmium, chromium, copper, lead, nickel, silver, and zinc. Five metals which can

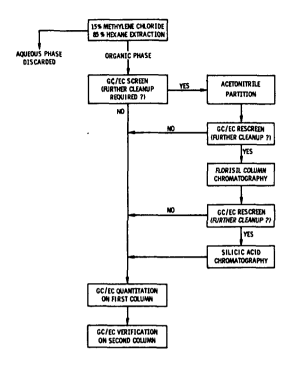


Figure 15. Sample processing scheme for pesticide and PCB analysis.

not be quantified by ICAP analysis were measured by conventional atomic absorption techniques: arsenic, beryllium, mercury, selenium, and thallium (4, 5).

ICAP forms an analytical system for simultaneous multielement determinations of trace metals at the sub-ppm level in solutions. The basis of this method is atomic emission. Excitation energy is supplied by coupling a nebulized sample with high temperature argon gas which has been passed through a powerful radio-frequency field. Emitted light is simultaneously monitored at 22 wavelengths corresponding to 22 different elements.

All samples for metals analysis were acidified in the field by adding 5 x  $10^{-6}$  m<sup>3</sup> of redistilled nitric acid to each  $10^{-3}$  m<sup>3</sup> of sample. Nitric acid blanks were also analyzed for metals.

## Analytical Results

Raw waste samples were collected with continuous samplers using a peristaltic pump which pulled the sample through Tygon tubing to the sample bottle. Sample blanks were collected by drawing laboratory-prepared organic-free water through the sampler prior to sample collection to determine the presence of base/neutral, acid, and pesticide organic priority pollutants. Results of these analyses are given in Table 10.

TABLE 10. SUMMARY OF CONTINUOUS SAMPLER AND VOLATILE ORGANIC BLANK ANALYSES

Fraction	Compound found	Concentration range, mg/m <sup>3a</sup>
Base/neutral	Naphthalene	2
<b>3430</b> , 3304 32 32	Dimethylphthalate	16
	Diethylphthalate	0.5 to 10.2
	Bis (2-ethylhexyl) phthalate	1.5 to 46
	Di-n-butylphthalate	1.3 to 1.7
Acids	Phenol	0.6 to 1.1
Volatiles	Toluene	2.6 to 55
	Trans-1,2-dichloroethylene	3.2
	Trichloroethylene	2.4
	Ethylbenzene	8.3
Pesticides and PCB	None	-

 $<sup>^{</sup>a}$ l mg/m<sup>3</sup> equals 1 µg/l.

To determine if any volatile organic species were absorbed from the air to the samples, EPA recommended that laboratory-prepared organic-free water be carried to the plant site, poured from the container into a sample vial, sealed, and shipped back to the laboratory for analysis (1). These results are included in Table 10.

All secondary effluent samples were collected by grab sampling techniques. Therefore, these samples were not passed through Tygon tubing. Samples were shipped and stored in ice at 4°C until extracted. A special effort was made to initiate methylene chloride extraction as rapidly as possible. The data on which each sample arrived at MRC and the date of its solvent extraction are shown in Table 11. Only 5 out of 64 samples were not extracted within 24-hr of receipt at MRC.

Results of GC/MS analysis of textile raw waste and secondary effluent samples for base/neutral organic compounds are presented in Table 12. GC/MS analyses for the volatile, acid, and pesticide/PCB organic compounds are presented in Table 13.

Of the 114 organic compounds in the priority pollutant list, a total of 45 different compounds were identified in textile wastewaters, 39 in raw waste samples and 34 in secondary effluent samples. The number of compounds found at each plant is summarized in Table 14. On an individual plant basis, the greatest number of organic compounds detected in a raw waste and secondary effluent sample were 14 and 8, respectively, with an average number per plant of 7 in the raw waste and 5 in the secondary effluent.

TABLE 11. TEXTILE SAMPLE EXTRACTION DATES

<del></del>		1 1
Plant	Date received at MRC	Date of initial extraction
		:
A	5-6-77	5-6-77
В	4-28-77	4-29-77
С	4-13-77	4-14-77
D	3-3-77	3-3-77
E	3-31-77	3-31-77
F	4-19-77	4-20-77
G-1-2	4-1-77	4-1-77
G-2-2	4-1-77	4-4-77
H .	3-10-77	3-10-77
J	3-9-77	3-9-77
K	4-5-77	4-6-77
L	4-12-77	4-12-77
M	3-16-77	3-17-77
N	4-27-77	4-28-77
P	3-2-77	3-2-77
' R	3-15-77	3-16-77
S	4-6-77	4-7-77
T	4-26-77	4-27-77
U	5-4-77	5-4-77
V	3-31-77	3-31-77
W	4-14-77	4-14-77
X	4-21-77	4-21-77
Y	3-11-77	3-11-77
Z	3-17-77	3-18-77
C-001	5-20-77	5-20-77
Y-001	5-20-77	5-20-77
JJ	6-23-77	6-23-77
KK	6-24-77	6-27-77
LL	6-27-77	6-29-77
MM	7-5-77	7-8-77
NN	6-30-77	7-1-77
00	7-5-77	7-6-77
PP	7-5-77	7-8-77

TABLE 12. GC/MS ANALYSES FOR BASE/NEUTRAL ORGANIC COMPOUNDS IN RAW WASTE AND EFFLUENT SAMPLES

Plant/source	<10 mg/m <sup>3</sup>	unds identified and concentrations of 10 to 100 mg/m <sup>3</sup>	>100 mg/m <sup>3</sup>
A/raw waste	Naphthalene 0.1 Dimethyl phthalate 3	1,4-Dichlorobenzene 11 1,2,4-Trichlorobenzene 90	мъьор
	Diethyl phthalate 1 Bis(2-ethylhexyl) phthalate 0.5		
<b>Veffluent</b>	1,2-Dichlorobenzene 1	1,2,4-Trichlorobenzene 46	NPPO
	1,4-Dichlorobensene 0.05 Bis(2-ethylhexyl) phthalate 6		
/raw waste	Diethyl phthalate 3.3	Naphthalene 41	nppo
	Bis(2-ethylhexyl) phthalate 5.7 Anthracene 0.1		
effluent	N-nitroso-di-n-propylamine 2	nppo	nppo
	Bis(2-ethylhexyl) phthalate 3 Pyrene 0.3		
C/raw waste	1,2-Dichlorobenzene 1.1	NPPO	Bis(2-ethylhexyl) phthalate 13
/effluent	Diethyl phthalate 4.1 1,2-Dichlorobenzene 0.3	1,2,4-Trichlorobenzene 10.2	ирро
/errruenc	Acenaphthene 0.5	1,2,4-111Ch1olobenzene 10.2	RFFO
	Bis(2-ethylhexyl) phthalate 3.0		
/raw waste	Anthracene 4.4 Naphthalene 0.3	Di-n-butyl phthalate 16.2	NPPO
	Bis(2-ethylhexyl) phthalate 8.9	NDDO	NDDO
/effluent	Diethyl phthalate 1 Bis(2-ethylhexyl) phthalate 5	NPPO	NPPO
/raw waste	1,4-Dichlorobenzene 2	NPPO	NPPO .
	Napthalene 1 Bis(2-ethylhexyl) phthalate 5		
effluent/	1,4-Dichlorobenzene 0.2	Bis(2-ethylhexyl) phthalate 18	NPPO
	1,2-Dichlorobenzene 0.2 Dimethyl phthalate 1		•
	Diethyl phthalate 0.5		ſ
/raw waste	Pyrene 0.1 1,4-Dichlorobenzene 6.5	1,2-Dichlorobenzene 34.6	1,2,4-Trichlorobenzene 120
/Idw waste	1,4-bicinolobenzene v.s	Acenaphthene 12.0 Fluorene 14.6	1,2,4-XIICHIOIODENZENE 120
/effluent	1,2,4-Trichlorobenzene 6.3	Diethyl phthalate 33.6 Bis(2-ethylhexyl) phthalate 23	NPPO
/raw waste	Pluorene 5	Naphthalene 95	Acenaphthene 273
/effluent	Acenaphthene 2.0	Bis(2-ethylhexyl) phthalate 19 Diethyl phthalate 11.1	NPPO
	Hexachlorobenzene 0.8	Bis(2-ethylhexyl) phthalate 10.3	*****
/raw waste	1,2-Dichlorobenzene 0.5 Naphthalene 3	Acenaphthene 27 Bis(2-ethylhexyl) phthalate 14	NPPO
	Di-n-butyl phthalate 2	-	
/effluent /raw waste	NPPO Diethyl phthalate 6.5	NPPO Naphthalene 79.7	Bis(2-ethylhexyl) phthalate 23 Bis(2-ethylhexyl) phthalate 16
/14# #4BCC	breenji phenatace v.s	Di-n-butyl phthalate 23.2	bid(z-echylhekyi) phohatace it
/effluent	Di-n-butyl phthalate 3.6 Pyrene 0.1	Bis(2-ethylhexyl) phthalate 35.2	NPPO
/raw waste	Naphthalene 0.03	NPPO	NPPC
/- 6 6 1 m h	Diethyl phthalate 0.2	tinno	
/effluent	Naphthalene 0.5 Bis(2-ethylhexyl) phthalate 8	NPPO	NPPO
/raw waste	1,4-Dichlorobenzene 1	Acenaphthene 30	Dimethyl phthalate lll
/effluent	Bis(2-ethylhexyl) phthalate 3 Bis(2-ethylhexyl) phthalate 2	NPPO	NPPO
/raw waste	NPPO	Naphthalene 92.9	1,2,4-Trichlorobenzene 156
/effluent	1,2,4-Trichlorobenzene 1.8	Di-n-butyl phthalate 58.4	Bis(2-ethylhexyl) phthalate 30 NPPO
/raw waste	Diethyi phthalate 5.9	Naphthalene 17	1,4-Dichlorobenzene 215
/off1	1 A-Drahlamahamana 1 E	Bis(2-ethylhexyl) phthalate 10.1	1,2-Dichlorobenzene 287
/effluent	1,4-Dichlorobenzene 1.5 1,2-Dichlorobenzene 6.0	Bis(2-ethylhexyl) phthalate 15.7	NPPO
,	Diethyl phthalate 9.4	<b>-1</b>	
/raw waste	Naphthalene 1.9 Diethyl phthalate 1.7	Dimethyl phthalate 11.6 Bis(2-ethylhexyl) phthalate 30.2	NPPO
	Di-n-butyl phthalate 9.8		
/effluent	NPPO	Bis (2-ethylhexyl) phthalate 72 N-nitroso-di-n-propylamine 18.9	NPPO
/raw waste	Di-n-butyl phthalate 7.3	NPPO	Bis(2-ethylhexyl) phthalate 12
/effluent <sup>C</sup> /raw waste	Diethyl phthalate 2 NPPO	Bis(2-ethylhexyl) phthalate 12 NPPO	NPPO 1.2.4-Trichlorobenzene 190
, Law waste	NEFO	MEFU ,	Naphthalene 143
/offluc=t	NDBO	Dig (2-oahu) howell ships at a 43	Bis(2-ethylhexyl) phthalate 13
/effluent	NPPO	Bis(2-ethylhexyl) phthalate 41	1,2,4-Trichlorobenzene 916 Naphthalene 255

al mg/m³ equals µg/].
b
No priority pollutants observed.

cSample inadvertently collected prior to the settling pond.

TABLE 12 (continued)

11 ans /an	<10 mg/m³	المعاني وسيد	is identified and concentration 10 to 100 mg/m <sup>3</sup>		>100 mg/m <sup>3</sup>
Plant/source	<10 mg/m²		10 to 100 mg/m		>100 mg/m²
T/raw waste	nppo <sup>b</sup>	•	N-nitrosodiphenylamine 11.3		Bis(2-ethylhexyl) phthalate 13
r/effluent	NPPO		Bis(2-ethylhexyl) phthalate 2:	3	NPPO
J/raw waste	1,2-Dichlorobenzene 2.0		Bis(2-ethylhexyl) phthalate 1		NPPO
,, 144 -4554	Naphthalene 1.5			•••	
	Diethyl phthalate 6.1				· · · · · · · · · · · · · · · · · · ·
/effluent	NPPO		Naphthalene 22		Bis(2-ethylhexyl) phthalate 14
//raw waste	Acenaphthene 8.7		1,2,4-Trichlorobensene 27.9		NPPO
/ Luw wabtu	Bis(2-ethylhexyl) phthalate 5		Dimethyl phthalate 12.9		
	Hexachlorobenzene 2.0		Present burneran serv		
//effluent	Bis (2-ethylhexyl) phthalate 9	1.5	NPPO		NPPO
/raw waste	Hexachlorobenzene 0.5		Bis(2-ethylhexyl) phthalate 1	8.1	NPPO
/effluent	NPPO		Bis(2-ethylhexyl) phthalate 1		NPPO
/raw waste	Naphthalene 1		Acenaphthene 53		NPPO
/Idw wasta	Bis(2-ethylhexyl) phthalate 1	ı	neutraphonome 33		W110
/offluent	Diethyl phthalate 3.2	•	NPPO		NPPO
/effluent			arro		NPFO
	Bis(2-ethylhexyl) phthalate 2				
	Hexachlorobenzene 0.5 1,2-Dichlorobenzene 0.1		2 fullitations 52 f		WDDA
/raw waste			2,6-Dinitrotoluene 53.5		NPPO
	Naphthalene 0.4		Bis(2-ethylhexyl) phthalate 8	/.3	****
effluent/	1,2-Dichlorobenzene 0.6		Bis(2-ethylhexyl) phthalate 2	5.2	NPPO
	Naphthalene 0.6				
	Diethyl phthalate 2.6				
	Di-n-butyl phthalate 6.7				
_	Hexachlorobenzene 0.3				
raw waste	NPPO		1,2,4-Trichlorobenzene 45		Naphthalene 309
					Bis(2-ethylhexyl) phthalate 21
/effluent	Bis(2-ethylhexyl) phthalate 2	ł .	NPPO		NPPO
-001/raw					
waste	NPPO		Naphthalene 17.0		NPPO
			Diethyl phthalate 69.0		
			Bis(2-ethylhexyl) phthalate 2:	3.3	
-001/raw			• • •		
waste	Naphthalene 4		Acenaphthene 13		NPPO
	Indeno(1,2,3-cd) pyrene 2		Diethyl phthalate 15		
	Bis(2-ethylhexyl) phthalate 3	3			
-001/		-			
effluent	Naphthalene 4.5		Diethyl phthalate 11.7		Bis(2-ethylhexyl) phthalata 13
J/raw waste	NPPO		1,2-Dichlorobenzene 11.4		1,2,4-Trichlorobenzene 435
J/effluent	NPPO		1,2,4-Trichlorobenzene 32.4		NPPO
K/raw waste	Diethyl phthalate 2.5		Dimethyl phthalate 11.6		NPPO
Wam mable	Bis (2-ethylhexyl) phthalate 9		name sult businesses 41.0		*****
	Pyrene 0.9				
¥ /a f f } u a n b	Bis (2-ethylhexyl) phthalate 4	. 1	NPPO		NPPO
K/effluent	Pyrene 0.2		MFFO		MPPO L
			Wambabalana 81 2		3 0 4 0-4-53
L/raw waste	1,2-Dichlorobenzene 0.6		Naphthalene 51.3		1,2,4-Trichlorobenzene 315
L/effluent	Dimethyl phthalate 0.2		NPPO		NPPO
	Bis(2-ethylhexyl) phthalate 5	.2			
M/raw waste	NPPO		Bis(2-ethylhexyl) phthalate 1	5.4	NPPO
i/effluent	Dimethyl phthalate 0.2		(MM-2-1) NPPO		NPPO
	Diethyl phthalate 1.2				
	Bis(2-ethylhexyl) phthalate 6				
	Bis(2-ethylhexyl) phthalate 2		(MM-3-1) NPPO		NPPO
	Bis(2-ethylhexyl) phthalate 3	1.0	(MM-4-1) NPPO		NPPO j
N/raw waste	NPPO		Bis(2-ethylhexyl) phthalate 2:		NPPO
N/effluent	NPPO		Bis(2-ethylhexyl) phthalate 2		NPPO
O/raw waste	NPPO		Bis(2-ethylhexyl) phthalate 2	6.0	NPPO
			Di-n-butyl phthalate 61.4		
O/effluent	Bis(2-ethylhexyl) phthalate 3	3.2	NPPO		NPPO
P/raw waste	NPPO		Butyl benzyl phthalate 72.8		NPPO
			Naphthalene 44.3		

al mg/m³ equals 1 µg/l.
bNo priority pollutant observed.

TABLE 13. GC/MS ANALYSES FOR VOLATILE, ACID, PESTICIDE, AND PCB ORGANIC COMPOUNDS IN RAW WASTE AND EFFLUENT SAMPLES

Toluene 61.1 Pentachlorophenol 30.1 Ethylbenzene 20.7 Chloroform 21.5 Trans-1,2-dichloroethylene 1.8 1,1,1-trichloroethane 16.7 Trichloroethylene 2.0 Chlorobenzene 1.0  E/effluent Toluene 5.5 NPPO NPPO  F/raw waste Trichloroethane 45 1,1-Dichloroethane 0.59 Pentachlorophenol 2.4 NPPO 1,2-Dichloropropane 1.50 1,1,1-Trichloroethane 11.26 Cis-1,3-dichloropropene 2.08 Toluene 12.28	Plant/source	Compounds iden Volatiles, mg/m³	atified and concentrations of Acids, mg/m <sup>3</sup>	pbserveda Pesticides and PCB, mg/m
### B/effluent	A/raw waste	NPPO		Heptachlor 6.37
Toluene 3.74  B/effluent Trichlorofluoromethane 2.60 NPPO NPPO  C/raw waste Toluene 236  Trichloroethylene 17.8 Toluene 236  Ethylbenzene 112 1,1,2,2-Tetrachloroethylene 26.4  C/effluent Toluene 2.6  D/raw waste Chloroform 3.3 Toluene 2.3  Ethylbenzene 57.3  D/effluent Toluene 1.67  E/raw waste Benzene 5.4 Toluene 61.1  E/raw waste Benzene 5.4 Toluene 61.5  Trichloroethylene 2.0  Chlorobenzene 1.0  E/effluent Toluene 5.5  F/raw location 2.7 Trichloroethylene 2.0  Chlorobenzene 1.0  E/effluent Toluene 5.5  F/raw waste Trichloroethane 1.73 Trichloroethylene 2.08 Toluene 12.28  F/effluent Trichloroethane 1.73 Trichloropopene 2.08 Toluene 0.8  F/effluent Toluene 0.9  F/effl	A/effluent	Toluene 8.4	NPPO	Heptachlor 1.55
C/raw waste C/raw waste C/raw waste Trichloroethylene 17.8 Toluene 236 Ethylbenzene 112 1,1,2,2-Tetrachloroethylene 26.4  C/efffluent Toluene 2.6 Ethylbenzene 2.0  D/raw waste Chloroform 3.3 Toluene 2.7 Ethylbenzene 57.3  D/effluent Toluene 6.1 E/raw waste  D/effluent Toluene 5.5  Frans-1,2-dichloroethylene 1.8 1,1,1-trichloroethane 16.7 Trichloroethylene 2.0 Chloroform 21.5 Frans-1,2-dichloroptopene 2.0 Chlorobenzene 1.0  E/effluent Toluene 5.5  F/raw waste  Trichlorofluoromethane 45 1,1-bichloroethylene 1.8 Columnia-1,3-dichloroptopene 2.08 Toluene 12.28  F/effluent Trichlorofluoromethane 1.73 Trichlorofluoromethane 1.73 Trichlorofluoromethane 1.73 Trichlorofluoromethane 3.90 Columnia-1,3-dichloroptopene 5.61 Toluene 0.8  F/raw waste Chloroform 5.19 Phenol 0.8 Phenol 0.8 Phenol 0.8 Phenol 12 Phenol 63 Z-wittrophenol 60 p-chloror-m-cresol 4.5 A-wittrophenol 65 A-wittrophenol 65 A-wittrophenol 65 A-wittrophenol 65 Toluene 36.1 NPPO NPPO NPPO NPPO NPPO NPPO NPPO NPP	B/raw waste		NPPO	NPPO
Toluene 236 Ethylbenzene 112 1,1,2,2-Tetrachloroethylene 26.4  C/effluent Toluene 2.0  D/raw waste Chloroform 3.3 Toluene 2.3 Ethylbenzene 57.3  D/effluent Toluene 1.67  E/raw waste Benzene 5.4 Toluene 61.1 Ethylbenzene 20.7 Chloroform 21.5 Frans-1,2-dichloroethylene 1.8 1,1,1-trichloroethylene 2.0 Chlorobenzene 1.0 Chlorobenzene 1.0 Chlorobenzene 1.0 Chlorobenzene 1.0 Chlorobenzene 1.0  E/effluent Toluene 5.5  F/raw waste Trichlorofuromethane 45 1,1-Dichloroethane 0.59 1,2-Dichloropropane 1.50 1,1,1-Trichloroethane 1.66 Cis-1,3-dichloropropene 2.08 Toluene 12.28  F/effluent Trichloroflucromethane 1.73 Frans-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 5.61 Toluene 0.8  E/effluent Toluene 0.8  E/raw waste Chloroform 5.19  Phenol 0.8  Phenol 0.8  Phenol 1.8  Phenol 2  Phenol 63 2-Nitrophenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65  Anitrophenol 65  NPPO NPPO NPPO NPPO NPPO NPPO NPPO NP	B/effluent	Trichlorofluoromethane 2.60	NPPO	NPPO
Ethylbenzene 2.0	C/raw waste	Toluene 236 Ethylbenzene 112	Phenol 0.5	NPPO `
Toluene 2.3   Ethylbenzene 57.3   D/effluent   Toluene 1.67   NPPO   N	C/effluent		NPPO	NPPO
E/raw waste  Benzene 5.4 Toluene 61.1 Ethylbenzene 20.7 Chloroform 21.5 Trans-1,2-dichloroethylene 1.8 1,1,1-trichloroethylene 2.0 Chlorobenzene 1.0  E/effluent  Toluene 5.5  MPPO  NPPO  Additional service of the service of t	D/raw waste	Toluene 2.3	Pentachlorophenol 22	NPPO .
Toluene 61.1 Ethylbenzene 20.7 Chloroform 21.5 Trans-1,2-dichloroethylene 1.8 1,1,1-trichloroethylene 2.0 Chlorobenzene 1.0  E/effluent Toluene 5.5  MPPO NPPO NPPO NPPO Praw waste Trichloroethane 0.59 1,2-Dichloropropane 1.50 1,1,1-Trichloroethane 11.26 Cis-1,3-dichloropropane 2.08 Toluene 12.28  F/effluent Trichlorofluoromethane 1.73 Trans-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 5.61 Toluene 0.85 Ethylbenzene 2.66  G/raw waste Chloroform 5.19 Phenol 0.8 Phenol 0.8 Phenol 2 NPPO Phenol 63 2-Nitrophenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65 4-Nitrophenol 65 MPPO Trichlorofluoromethane 2,138  J/raw waste Toluene 3.6.1 NPPO NPPO NPPO NPPO NPPO NPPO NPPO NPP	D/effluent	Toluene 1.67	NPPO	NPPO
F/raw waste  Trichlorofluoromethane 45 1,1-Dichloropropane 1.50 1,2-Dichloropropane 1.50 1,1,1-Trichloroethane 11.26 Cis-1,3-dichloropropene 2.08 Toluene 12.28  F/effluent  Trichlorofluoromethane 1.73 Trans-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 5.61 Toluene 0.85 Ethylbenzene 2.66  G/raw waste  Chloroform 5.19  Phenol 0.8  Phenol 0.8  Phenol 2  NPPO  MPPO  MPPO  H/raw waste  Toluene 0.8  Toluene 25.7 Ethylbenzene 5.7  Phenol 63 2-Nitrophenol 60 p-Chlorom-cresol 4.5 4-Nitrophenol 65  MPPO  MPPO  MPPO  MPPO  MPPO  NPPO  NPPO  NPPO  MPPO  NPPO	E/raw waste	Toluene 61.1 Ethylbenzene 20.7 Chloroform 21.5 Trans-1,2-dichloroethylene 1.8 1,1,1-trichloroethane 16.7 Trichloroethylene 2.0		NPPO
1,1-Dichloroethane 0.59 1,2-Dichloropropane 1.50 1,1,1-Trichloroethane 11.26 Cis-1,3-dichloropropene 2.08 Toluene 12.28  F/effluent  Trichlorofluoromethane 1.73 Trans-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 5.61 Toluene 0.85 Ethylbenzene 2.66  G/raw waste Chloroform 5.19  Phenol 0.8  Phenol 0.8  Phenol 2  Phenol 2  Phenol 3  2-Nitrophenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65  M/effluent  Toluene 36.1  NPPO	E/effluent	Toluene 5.5	NPPO	NPPO
Trans-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 5.61 Toluene 0.85 Ethylbenzene 2.66  G/raw waste Chloroform 5.19 Phenol 0.8 NPPO  G/effluent Toluene 0.8 Phenol 2 NPPO  H/raw waste Toluene 25.7 Ethylbenzene 5.7 Phenol 63 2-Nitrophenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65  H/effluent Toluene 11.9 Trichlorofluoromethane 2,138  J/raw waste Toluene 36.1 NPPO NPPO  D/effluent Toluene 8.0 NPPO NPPO  NPPO NPPO	F/raw waste	1,1-Dichloroethane 0.59 1,2-Dichloropropane 1.50 1,1,1-Trichloroethane 11.26 Cis-1,3-dichloropropene 2.08		NPPO
G/effluent Toluene 0.8 Phenol 2 NPPO  H/raw waste Toluene 25.7 Phenol 63 NPPO  Ethylbenzene 5.7 Phenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65  H/effluent Toluene 11.9 NPPO  Trichlorofluoromethane 2,138  J/raw waste Toluene 36.1 NPPO NPPO  J/effluent Toluene 8.0 NPPO NPPO  J/effluent Toluene 8.0 NPPO NPPO	F/effluent	Trans-1,3-dichloropropene 3.90 Cis-1,3-dichloropropene 5.61 Toluene 0.85	2,4-Dimethylphenol 9	прро
H/raw waste Toluene 25.7 Ethylbenzene 5.7 Phenol 63 2-Nitrophenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65  NPPO Trichlorofluoromethane 2,138  J/raw waste Toluene 36.1 NPPO NPPO NPPO NPPO NPPO NPPO NPPO NPP	G/raw waste :	Chloroform 5.19	Phenol 0.8	NPPO
Ethylbenzene 5.7  2-Nitrophenol 60 p-Chloro-m-cresol 4.5 4-Nitrophenol 65  NPPO	G/effluent	Toluene 0.8	Phenol 2	NPPO
Trichlorofluoromethane 2,138  I/raw waste Toluene 36.1 NPPO NPPO I/effluent Toluene 8.0 NPPO NPPO I/effluent Toluene 8.0 NPPO NPPO	H/raw waste		2-Nitrophenol 60 p-Chloro-m-cresol 4.5	NPPO
7/effluent Toluene 8.0 NPPO NPPO D/effluent Toluene 8.0 NPPO NPPO	H/effluent		NPPO <sup>b</sup>	NPPO
J/effluent Toluene 8.0 NPPO NPPO	J/raw waste	Toluene 36.1	NPPO	NPPO
	J/effluent	Toluene 8.0	NPPO	NPPO
	J/effluent		NPPO	NPPO

a1 mg/m<sup>3</sup> equals 1  $\mu$ g/1.

bNo priority pollutant observed.

TABLE 13 (continued)

Plant/source	Volatiles, mg/m <sup>3</sup>	ntifled and concentrations ob Acids, mg/m <sup>3</sup>	Pesticides and PCB, mg/m
K/raw waste	Chloroform 4.8 Toluene 29.3 Ethylbenzene 63.8	2,4,6-Trichlorophenol 0.7 Pentachlorophenol 3.9	NPPO .
K/effluent	Chloroform 58.1 Trichloroethylene 4.6 Toluene 24.0 Ethylbenzene 0.7	NPPO	ү-ВНС 0.31
L/raw waste	Chloroform 2.5 Toluene 5.2 Ethylbenzene 2.0	NPPO	NPPO
L/effluent	Benzene 0.5	NPPO	NPPO
M/raw waste	ирро <sup>р</sup>	Phenol 12.4 Pentachlorophenol 6.9	NPPO
M/effluent	Toluene 0.4	NPPO	NPPO
N/raw waste	Trichloroethylene 20.8 Toluene 43.8 Ethylbenzene 1,770	Phenol 11	NPPO
N/effluent	Toluene 16.6 Ethylbenzene 75.0	2,4-Dimethylphenol 8	NPPO
P/raw waste	Chloroform 17.3 Toluene 36.1 Ethylbenzene 1,209 Chlorobenzene 24.8	Phenol 6.6	NPPO
P/effluent	Chloroform 6.9 Toluene 22.4 Ethylbenzene 278	NPPO	NPPO
R/raw waste	Chloroform 33.2 Benzene 31.0 Toluene 281 Ethylbenzene 2,835 Chlorobenzene 296 1,1,2,2-Tetrachloroethylene 15.1	NPPO	NPPO
R/effluent <sup>C</sup>	Toluene 16.8 Ethylbenzene 28.7	Chloro cresol 32 Pentachlorophenol 56	NPPO
S/raw waste	Chloroform 71.1 1,1,2,2-Tetrachloroethylene 38.7 Chlorobenzene 13.6 Toluene 60.7 Ethylbenzene 851.7	NPPO	β <b>-В</b> НС 0.35
3/effluent	Toluene 21.4 Ethylbenzene 109 1,1,2,2-Tetrachloroethylene 0.4	NPPO	NPPO
C/raw waste	Toluene 303 Ethylbenzene 18.4 1,1,2,2-Tetrachloroethylene 6.4	NPPO	NPPO

bno priority pollutant observed.

Sample inadvertently collected prior to the settling pond.

TABLE 13 (continued)

Plant/source	Compounds identities, mg/m <sup>3</sup>	tified and concentrations of Acids, mg/m3	bserved <sup>a</sup> Pesticides and PCB, mg/m <sup>3</sup>
T/effluent	Toluene 33.1 1,1,2,2-Tetrachloroethylene 2.9	nppo <sup>b</sup>	NPPO
U/raw waste	1,1-Dichloroethane 3.67 1,1,1-Trichloroethane 306	Phenol 0.7 Pentachlorophenol 1.6	NPPO
U/effluent	Chloroform 18.05 Bromodichloromethane 1.54 Trans-1,3,-dichloropropene 0.89 Toluene 13.03	NPPO	NPPO .
V/raw waste	Toluene 8.4 Ethylbenzene 4.9	NPPO	NPPO
V/effluent	Toluene 1,401	NPPO	NPPO
W/raw waste	Trichloroethylene 13.1 Benzene 19.4 Toluene 62.2 Ethylbenzene 1.1	Phenol 100	NPPO .
W/effluent	Toluene 1.7	NPPO	NPPO
X/raw waste	1,1,1-Trichloroethane 8.2 Toluene 63.5 Ethylbenzene 369 1,1,2,2-Tetrachlorethylene 414.2	Phenol 3.8	NPPO
X/effluent	Toluene 39.6 Trichlorofluoromethane 35.0 1,1,2,2-Tetrachloroethylene 40.5	NPPO	NPPO
Y/raw waste	NPPO	Phenol 10.0	NPPO
Y/effluent	Chloroform 4.8 Trichlorofluoromethane 10.1	NPPO	NPPO
2/raw waste	Toluene 5.5 Ethylbenzene 0.7 1,1,2,2-Tetrachloroethylene 12.0	Phenol 34	NPPO
Z/effluent	Toluene 110.6 Ethylbenzene 3,018 Chlorobenzene 3.5 Trichlorofluoromethane 89.3	NPPO	NPPO
C-001/raw waste	Toluene 5.7 Trichlorofluoromethane 26.8	NPPO	NPPO
Y-001/raw waste	Chloroform 14.3 Chlorobenzene 1.6 Chlorobenzene 1.6 Toluene 11.6 Ethylbenzene 1.9	Phenol 19	NPPO
Y-001 effluent	Toluene 15.1	Phenol 2.9 p-Chloro-m-cresol 1.6	NPPO
JJ/raw waste	Trichloroethylene 187 1,1,2,2-Tetrachloroethylene 1,126 Ethylbenzene 14	Phenol 41.4	Not analyzed

 $<sup>\</sup>overline{^{3}}$ l mg/m $^{3}$  equals 1 µg/l.  $^{b}$ No priority pollutant observed.

TABLE 13 (continued)

	Compounds identified and concentrations observed												
Plant/source	Volatiles, mg/m <sup>3</sup>	Acids, mg/m <sup>3</sup>	Pesticides and PCB, mg/m										
JJ/effluent	Trichloroethylene 84	NPPO <sup>b</sup>	Not analyzed										
KK/raw waste	Trichloroethylene 52 Toluene 28 Chlorobenzene 42 Ethylbenzene 26	2-Chlorophenol 131 2,4,6-Trichlorophenol 20.2 Pentachlorophenol 20.4	Not analyzed										
KK/effluent	Benzene 64 Chlorobenzene 26	2-Chlorophenol 10 2,4,6-Trichlorophenol 21.1	Not analyzed										
LL/raw waste	Chloroform 498 Trichloroethylene 121 1,1,2,2-Tetrachloroethylene 1,108 Ethylbenzene 484	Phenol 16.1	Not analyzed										
LL/effluent	NPPO	NPPO .	Not analyzed										
MM-1/raw waste	NPPO	NPPO	Not analyzed										
MM-2/effluent	NPPO	NPPO	Not analyzed										
MM-3/effluent	NPPO	NPPO	Not analyzed										
MM-4/effluent	Toluene 2	NPPO	Not analyzed										
NN/raw waste	NPPO	Phenol 10.1	Not analyzed										
NN/effluent	NPPO	NPPO	Not analyzed										
00/raw waste	Chloroform 48 Trichloroethylene 42	Phenol 22.9	Not analyzed										
00/effluent	Chloroform 10 Toluene 3	NPPO	Not analyzed										
PP/raw waste	Benzene 200 Toluene 83 Ethylbenzene 42	NPPO	Not analyzed										

al mg/m³ equals 1 µg/l.
bNo priority pollutant observed.

TABLE 14. NUMBER OF PRIORITY ORGANIC POLLUTANTS FOUND IN THE RAW WASTE AND SECONDARY EFFLUENT STREAMS

	Number	of organi	c compounds	detecteda effluent					
Plant	In raw	waste I	n secondary	effluent					
<b>À</b>		)	6	÷.					
<b>B</b> .	ē	5	4						
B C D E F G H	8	3							
Ď		7	7 3						
E	13	3	7 8						
F	14	l	8						
G	11	5	6 3 5 7 2 3 7 5 8 3 6 2 2 7 8 5						
H	11	L	3						
J	5	5	5						
K	5	5	7						
L	7	7 .	2						
M	5	5	3						
N '	- 10	)	7						
P	10	)	5 <sub>k</sub>						
Ŕ	8	3	3n						
;S	9	9	8						
T			3	,					
Ü		3	6						
<b>.V</b>	-	<u>′</u>	2						
W		/	2						
X	č	<u>.</u>	/						
Y Z		)	8						
C-001	\$ 8 5 8 8 8	) :	N <sub>V</sub> C						
V-001	1 1	)	NA ·						
77d X-001	11	L 2	2						
KKď	11	,	2						
$\frac{n}{n}d$	7.1	L ·	2						
LLd	11 8 . 2 5	<b>,</b>	6 2 6 2 4						
NN d		- )	1						
OOd	2		1 3						
PPd		<b>,</b>							
PPu	5	•	NA						

a In the list of priority pollutants there are 114 organic compounds.

bSample inadvertently collected prior to the setting pond.

CNot applicable; these plants discharge their wastewater to a municipal treatment system.

dPesticides were not analyzed at this plant.

The frequency of occurrence of 45 organic species in 64 waste-water samples is given in Table 15. Dominant compounds were bis(2-ethylhexyl)phthalate found in 54 samples, toluene found in 44 samples, and ethylbenzene found in 30 samples.

Results of metal analyses by inductively coupled argon plasma (ICAP) and atomic absorption (AA) analysis are presented in Tables 16 and 17.

Table 16 lists concentrations for the metals included in the 129 consent decree priority pollutants list (Appendix A). Table 17 lists data for the additional elements automatically measured in ICAP analysis. The upper and lower rows of numbers for each plant correspond to metal concentrations in the raw wastewater and secondary effluent, respectively. All metal concentrations are reported as  $g/m^3$  (ppm).

The lower detection limits for routine ICAP and AA analyses, referred to in footnote "a" of Tables 16 and 17, are given in Table 18.

#### LEVEL 1 CHEMICAL ANALYSIS

# Analysis Procedures

The EPA-Process Measurements Branch, IERL/RTP, has developed a phased sampling and analytical strategy for environmental assessment programs (2). Level 1 is the first part of a three-phase approach for performing the assessments. The Level 1 chemical analysis procedure, including modifications to accommodate the special requirements of textile wastewaters, are discussed in this section.

Level 1 chemical analyses were performed on samples from only 15 of the 23 basic textile plants because this task was implemented by EPA after the program began. Eight-hour composited grab samples were collected from the secondary effluent at the 15 plants.

Figure 16 is a schematic diagram for field handling of wastewater samples as recommended by EPA (2). The procedure specifies collection of  $0.02~\text{m}^2$  of composite sample, which is divided into two  $0.01~\text{m}^3$  portions of organic and inorganic chemical analysis. Part of the inorganic composite  $(0.001~\text{m}^3)$  is set aside for determination of  $BOD_5$  and COD. The remainder is filtered in the field with Gelman Spectro-Grade glass fiber filters (or equivalent). Tared filters are sent to the laboratory for analysis of inorganic elements, leachable anions, and total suspended solids.

Filtrate is extracted in the field with methylene chloride to separate organic from inorganic chemical species. The aqueous portion is divided into three parts for analysis.

TABLE 15. OCCURRENCE OF PRIORITY ORGANIC POLLUTANTS COMBINED FROM RAW WASTE AND SECONDARY EFFLUENT SAMPLES

		mber of Sam ollutant wa	ples in which	Observed
		Raw waste	Secondary	concentration
Priority pollutant	Total	samples	effluent samples	
Bis(2-ethylhexyl)phthalate	54	27	27	0.5 to 300
Toluene	44	22	22	0.4 to 300
Ethylbenzene	-30	20	10	0.7 to 3,000
Naphthalene	25	20	5	0.03 to 300
Diethyl phthalate	21	12	9	0.2 to 70
Phenol	21	19	2	0.5 to 100
Chloroform	17	12	5	2 to 500
1,2,4-Trichlorobenzene	14	8	6	2 to 900
1,2-Dichlorobenzene	13	8	5	0.1 to 300
1,1,2,2-Tetrachloroethylene	11	8	3	0.4 to 2,100
Trichloroethylene	10	8	2	2 to 200
Acenaphthene	9	7	2	0.5 to 270
Di-n-butyl phthalate	9	6	3	2 to 60
Pentachlorophenol	8	8	0	2 to 70 .
Dimethyl phthalate	8	5	3	0.2 to 110
1,4-Dichlorobenzene	8	5	3	0.05 to 200
Chlorobenzene	8	6	2	1 to 300
Trichlorofluoromethane	8	2	6	2 to 2,100
Benzene	6	4	2	0.5 to 200
1,1,1-Trichloroethane	5	5	0	2 to 300
Pyrene	5	1	4	0.1 to 0.9
Hexachlorobenzene	5	2	3	0.3 to 2
2,4,6-Trichlorophenol	3	2	1	0.7 to 20
N-nitroso-di-n-propylamine	3	1	2	2 to 20
N-nitrosodiphenylamine	3	1	2	2 to 20
Heptachlor	2	1	. 1	2 to 6
Anthracene	2	1	1	0.1 to 4
Fluorene	2	2	0	5 to 15
1,1-Dichloroethane	2	2	0	0.6 to 4
Cis-1,3-dichloropropene	2	1	1	2 to 6
Trans-1,3-dichloropropene	2	0	2	0.9 to 4
2,4-Dimethylphenol	2	0	2	8 to 9
2-Chlorophenol	2	i	1	10 to 130
α-BHC	1	Ö	ī	0.3
β-BHC	1	i	Ö	0.4
2,6-Dinitrotoluene	1	i	Ö	50
Indeno(1,2,3-cd)pyrene	1	1	0	2
Butylbenzyl phthalate	ī	ō	ì	70
Trans-1,2-dichloroethylene	ī	i	ō	2
1,2-Dichloropropane	ī	ĩ	Ö	2
2-Nitrophenol	ī	ĩ	Ö	70
4-Nitrophenol	ī	ā	ŏ	70
Chloro cresol	ī	ō	i	30
Bromodichloromethane	ī	Ŏ	ī	2

Out of a total of 64 samples.

b Rounded to one significant figure.

 $<sup>^{\</sup>text{C}}$ l mg/m³ equals 1 µg/l.

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TABLE 16. CONSENT DECREE METALS CONCENTRATIONS IN WASTEWATER SAMPLES

		Metals concentration - raw waste, $g/m^3$ - secondary effluent, $g/m^3$													
Plant Gode	Silver	Arsenic	Beryllium	Cadmium	Chromium	Copper	Mercury	Nickel	Lead	Antimony	Selenium	Thallium	Zinc		
A	a a	a a	b b	a a	0.19 0.18	0.021 0.027	0.004 a	0.009 0.14	a a	a 0.03	a a	a	1.3		
В	a a	a a	b b	0.0007	0.012 0.004	0.074 0.03	0.0009	a a	a a	a a	a	a a	0.3 0.17		
С	a	a	a	0.005	0.035	0.008	a	0.15	0.12	0.007	a	a	0.074		
	a	a	a	0.006	0.031	0.020	0.0007	0.14	0.12	0.004	a	a	0.12		
D	0.011 a	0.017 0.006	a a	a a	a a	0.031 a	a a	0.03 a	a	0.003 0.002	a a	a a	0.21 0.21		
E	0.007	<b>a</b>	a	0.006	0.011	0.84	a	0.04	0.008	0.008	a	a	7.9		
	a	a	a	0.001	0.004	0.03	a	0.04	&	0.0008	a	a	5.1		
F	0.10 0.08	a a	b b	0.01	0.006 0.004	0.59 0.13	a 0.0009	0.10 0.06	0.08 0.0006	0.001 0.0003	a a	a a	0.26 0.57		
G	0.0085	ā	a	a	0.004	0.063	a	0.028	0.006	0.052	a	a	0.45		
	a	a	a	a	0.003	0.028	a	0.013	a	0.011	a	a	0.26		
н	0.041 a	a a	a <b>a</b>	a a	0.004 a	0.022 a	a a	0.014 a	a	0.004 0.006	a a	a a	3.9 0.96		
J	0.06	a	a	a	0.048	2.4	a	0.097	0.029	0.0007	a	a	2.1		
	a	a	a	a	0.025	0.1	a	0.09	a	a	a	a	0.8		
ĸ	0.13	0.006	a	0.004	0.019	0.026	a	0.1	0.03	0.003	a	a	0.15		
	a	a	a	a	0.004	0.015	a	a	a	0.0008	a	a	0.11		
L	a	a	a	a	0.003	0.30	a	0.054	0.036	0.005	a	a	1.0		
	a	a	a	a	0.03	0.096	a	0.035	a	0.003	a	a	0.72		
м	a a	a a	a a	a a	a a	0.009 0.005	a a	a	a a	0.0008 0.004	a a	a a	1.2 0.41		
N	a	a	b	0.046	0.88	0.020	0.0004	a	a	0.0002	a	a	7.5		
	a	a	b	a	1.8	0.008	a	0.03	a	0.002	a	a	38.4		
P	0.03	· a	a	a	0.003	a	a	0.10	0.013	a	a	a	0.20		
	0.008	a	a	a	a	a	a	0.04	a	a	a	a	0.14		
R <sup>C</sup>	a	a	a	a	0.067	0.51	a	a	a	a	a	a _	0.24		
	a	a	a	a	<b>0.14</b>	0.29	a	a	a	a	a	a	0.21		
s	a a	0.005 a	a	a a	0.0007 a	0.04 0.06	a a	а	a a.	0.057 0.074	a a	a a	0.12 0.084		

a Metals concentration below instrument detection limit - see Table 18 for detection limit.

Analysis not performed.

<sup>&</sup>lt;sup>C</sup>Secondary effluent sample inadvertently collected prior to the settling pond.

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TABLE 16 (continued)

Plant				Metals	concentrat				- /- 3				
code	Silver	Arsenic	Beryllıum	Cadmium	Chromium	Copper	condary e	Nickel	Lead	Antimony	Selenium	Thallium	Zinc
T	a	а	b	a	a	0.12	0.0007	0.05	0.025		a	a	0.29
	a	a	b	, a	a	0.06	а	0.004	a	• -	a	, a	0.08
ប	a	a	ь	a	0.027	0.040	0.0004	0.008	a	0.007	a	. <b>a</b>	0.26
	a	a	b	a	0.014	0.023	a		ā	0.001	ă	ā	0.19
v	a	a	a	0.005	0.004	0.23	a	a	a	a	a	a	0.46
	a	a	a	a	0.003	0.17	a	a	a	0.004	8	a	0.34
W	0.065	· a	а	0.009	0.012	0.023	a	0.054	0.018	a	a	a	0.19
	0.095	0.004	а	0.013	0.003	0.002	0.0005	0.060	0.057	a	a	a	0.09
x	0.017	а	ь	0.005	0.024	0.084	a	0.11	0.032	0.0003	a	a	0.034
	0.033	a	b	0.007	0.039	0.11	0.0009	0.072	0.026	0.0009	a	a	0.07
Y	a	a	а	a	0.026	0.096	a	0.012	a	0.016	ā	a	0.24
	a	a	a	a	0.001	0.11	a	a	a	0.003	<b>a</b>	a	0.09
2	a	a	а	a	a	0.097	a	0.011	a	0.011	a	a	0.11
	a	a	a	а	a	0.050	a	A	a	0.012	a	a	0.37
Y-001	0.068	a	ь	0.006	0.65	0.041	ь	0.20	0.16	a	ь	b	0.13
	0.057	a	b	0.007	0.29	a	ь	0.16	0.16	a	b	ъ	0.10
C-001	0.033	a	b	0.003	0.024	a	b	0.099	0.073	þ	b	Þ	0.056
JJ	0.047	0.20	b	0.005	0.16	0.032	ь	0.10	0.084	Þ	b	b	0.13
	0.049	0.16	ь	0.005	0.08	0.031	Þ	0.12	0.085	p	þ	þ	0.32
KK	0.022	0.12	ь	0.002	0.016	0.086	ь	0.077	0.049	b	b	ь	1.08
	0.044	a	ь	0.004	0.013	0.037	b	0.11	0.044	Þ	þ	b	0.39
LL	0.058	0.10	þ	0.004	0.011	0.038	þ	0.13	0.060	ь	b	b	0.06
	0.056	0.07	ь	0.002	0.020	0.092	ь	0.15	0.048	þ	þ	b	0.06
<b>1</b>	0.016	0.055	b	0.002	0.111	0.036	þ	0.044	0.011	ь	b	b	0.24
-2	0.025	0.003	þ	0.002	0.058	0.059	ь	0.072	0.037	ь	ь	þ	0.19
-3 -4	0.028 0.032	0.007 0.006	b b	a 0.002	0.11 0.13	0.028 0.042	b b	0.067 0.081	0.031 0.035	b b	b b	b b	0.28 0.37
NN	0.042	а	ь	0.002	0.023	0.047	b	0.098	0.033	ь	ь	b	0.08
F444	0.033	ā	Ď	0.004	0.17	0.046	Ď	0.079	0.025	Ď	Ď	b	0.13
00	0.046	a	b	0.004	0.011	0.039	ь	0.11	0.043	ь	ь	ь	0.12
	0.050	a	b	0.005	0.012	0.037	b	0.12	0.084	ь	þ	b	2.3
PP	0.048	a	b	0.005	0.010	0.041	ď	0.12	0.078	ь	å	ь	0.07

<sup>&</sup>lt;sup>a</sup>Metals concentration below instrument detection limit - see Table 18 for detection limit.

b Analysis not performed.

<sup>&</sup>lt;sup>C</sup>Secondary effluent sample inadvertently collected prior to the settling pond.

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TABLE 17. OTHER METALS GIVEN BY ICAP ANALYSIS

Plant	Metals concentration - raw waste, $g/m^3$ - secondary effluent, $g/m^3$														
code	Aluminum	Barlum	Boron	Calcium	Cobalt	Iron	Magnesium		Molybdenum		Bilicon	Tin	'Phosphorus	Titanium	Vanadius
A	0.24 0.23	0.064	0.15 0.18	18 28	0.0042 0 013	0.72 2.1	3.6 4.0	0.05 0.07	a a	>100 >100	3.3 1.4	0.05	1.2 0.50	0.012 0.008	0.060 0.59
В	0.16	0.037	0.07	20	0.0056	0.32	9.5	0.093	a	>100	4.4	0.018	12	0.0039	0.058
	0.024	0.008	0.043	12	0.0052	0.27	4.67	0.059	a	>100	2.3	0.041	6.5	a	0.030
С	0.30 0.19	0.083	0.39 0.15	5.0 4.5	0.0081 0.0056	1.0 0.22	3.9 0.73	0.029 0.017	0.019 0.024	>100 >100	16 15	0.07	4.0 4.1	0.020 0.012	0.29 0.40
D	0.38 0.15	a A	1.4 0.90	3.6 6.6	0.001	0.39	3.1 4.6	0.03 0.07	a a	>100 >100	6.B 7.3	0.02	1.6 1.0	0.008 0.0001	0.03
		_							a						
E	0.028 0.038	0.015 0.012	2.36 1.06	5.4 39	a a	0.12 0.62	2.5 3.1	0.027 0.10	a 0.019	>100 54	11 8.6	0.03	1.9 1.4	0.018 0.012	0.019 0.021
F	0.11	0.03	0.84	4.1	0.0041	0.33	3.1	0.007	0.025	>100	10	0.07	24 9.5	0.0028 0.0039	0.014 0.013
	0.06	0.03	0.68	4.2	0.012	0.28	3.2	0.009	0.023	>100	13			_	
G	0.52 0.23	0.008 0.015	0.69 0.22	4.2 3.1	0.0008 0.011	0.17 0.39	0.52 0.49	0.05 0.038	0.0028 •	97 55	2.4 3.2	0.03	- 6.4 6.1	0.011	0.020
H	0.093 0.055	ā A	a a	1.3 6.2	0.0054 a	0.16 0.40	2.6 3.2	0.013 0.009	0.011 a	>100 >100	17 15	0.019 0.055	0.99 0.20	0.009 a	0.035 0.032
J	0.95 0.01	0.12 0.024	a a	4.9 5.2	0.0084	0.70 0.52	5.8 6.9	0.05 0.05	0.0043 a	>100 >100	18 17	0.03 0.05	3.3 0.6	1.5 0.06	0.13 0.11
K	0.28 a	0.028 0.018	13	5.8 3.6	0.009	0.67 0.088	4.2 3.7	0.038 0.011	0.020 0.0006	>100 >100	23 15	0.05 0.04	1.9 0.93	0.0035 0.0104	0.012 0.035
L	0.095 0.064	0.008	6.1 8.9	3.4 6.3	0.0012 0.0012		1.3	0.022 0.021	<b>a</b> a	>100 >100	6.0 6.9	0.019	2.2 1.6	0.010 0.0013	0.012 0.032
м	0.33 0.009	0.013 a	0.91 0.53	9.7 8 8	0.0077 0.0045	0.18	6.7 6.4	0.034 0.14	a	>100 >100	15 14	4.0 3.5	3.99 3.46	0.0092 a	0.042 0.037
N	0.29 1.2	0.018 0.16	0.041 0.058	10 26	0.007 0.030	1.4	1.68 4.2	0.46 1.1	a 0.008	>100 >100	11 6.9	0.012 a	0.43 5.2	0.010	0.013 0.037
P	0.62 0.14	a a	0.20 0.52	1.6 9.5	0.007 0.003	0.72 0.12	0.48 1.8	0.042 0.02	0.002 a	76 >100	5.7 4.8	0.06 0.004	5.7 2.2	0.006	0.02 0.02
R <sup>C</sup>	0.28 0.07 -	0.003	0.21	7.5 . 5.1.	0.028 0.026	0.30 0.13	3.6 _ 3.0	0.018 0.013	a a	>100 >100	13 11	0.003 a	3.9 .0.66	0.071	0.021 0.016
s	0.068 0.91	0.005 0.028	1.6	2.8 7.3	0.001 0.001	0.12	1.8	0.021 0.03	a a	>100 >100	14 15	0.007	1.6 5.0	0.0005	0.008

Metals concentration below instrument detection limit - see Table 18 for detection limits.

b Secondary effluent sample only.

C Secondary effluent samples inadvertently collected prior to settling pond.

TABLE 17 (continued)

Plant			• • • • •			Meta	ls concent	ation - was	ite, g/m <sup>3</sup>		3				
code	Aluminum	Barium	Boron	Calcium	Cobalt	Iron	Magnesium	- sec	condary efflu Molybdenum	Sodium	Silicon	Tin	Phosphorus	Titanium	Vanadium
_		•													
T	0.20 0.075	0.013	0.73	12	0.008	0.14	4.2	0.18	a	>100	15	0.042	12	0.0009	0.045
	0.075	0.008	0.34	9.9	د0.00	0.43	3.5	0.17	a	>100	6.6	0.028	17	0.0021	0.037
U	0.90	0.01	0.032	5.8	0.005	0.68	2.5	0.076	a	>100	3.7	0.028	3.5	0.006	0.023
	0.17	0.006	0.28	8.0	a	0.30	2.5	0.016	a	>100	3.0	a	3.7	0.003	0.019
v	0.20	0.014	3.4	3.2	a	0.37	1.4	0.063	a	94	4.5	0.013	0.75	0.003	0.019
	0.16	0.013	0.49	3.6	0.004	0.47	1.7	0.072	ā	>100	4.5	0.005	0.78	a	0.013
W	6.0	0.33	0.44	93	0.14	5.7	14	0.89	a	65	20	0.030	5.1	0.035	0.14
	0.77	0.15	0.14	36	0.005	1.7	5.0	0.38	0.021	51	10	0.030	0.15	0.012	0.036
x	0.14	0.028	0.28	8.3	0.034	0.22	1.4	0.020	0.008	82	7.9	0.006	4.6	0.011	0.022
	0.94	0.005	18.0	9.3	0.024	<b>0.35</b>	1.9	0.010	0.012	>100	8.4	0.002	5.4	0.001	0.029
<b>y</b> .	0.076	0.13	1.4	2.1	0.026	0.14	3.2	0.044	a	>100	9.8	0.001	16	0.001	0.019
_	0.018	0.029	2.3	8.9	0.004	0.17	5.8	0.046	ā	>100	8.2	0.002	15	a	0.037
_															
Z	0.023	a	0.12	2.5	0.009	0.16	0.65	0.014	<b>a</b>	>100 >100	8.8		1.1 0.5	0.002	0.089
	, 4	a	0.36	2.1	0.006	0.075	1.2	0.010	a	>100	6.2	a	0.5	•	0.086
Y-001	0.57	0.015	0.045	68	0.32	1.2	3.1	0.12	0.031	65	7.8	0.068	11.7	0.020	0.039
	6.3	0.011	0.68	61	0.27	1.1	3.0	0.08	0.042	59	7.7	0.058	6.8	0.008	0.088
C-001	0.29	0.070	0.11	22	0.019	2.9	8.1	0.45	0.011	>100	12	0.040	2.7	0.010	0.072
JJ	2.7	0.048	0.23	7.3	0.012	2.3	2.2	0.14	0.026	88	9.8	0.057	3.5	0.096	0.044
	1.8	0.042	0.24	16.5	0.010	1.5	4.0	0.19	0.024	>100	7.8	0.042	2.3	0.056	0.049
KK	0.38	0.060	1.0	12.4	0.004	0.42	14.4	0.054	0.010	>100	31.9	0.077	6.3	0.042	0.091
	0.26	0.051	0.85	11.8	0.009	0.46	11.6	0.036	0.015	>100	27.8	0.064	6.4	0.017	0.077
		_													
ᄔ	0.22	0.008	0.043	4.5	0.004	0.16	2.0	0.040	0.015	>100	5.7	0.033	18.8	0.014	0.032
	0.17	0.004	0.091	3.7	0.004	0.10	2.0	0.021	0.016	>100	7.1	0.031	28.8	0.008	0.086
KM-1	0.44	0.010	1.1	5.7	0.004	0.39	3.7	0.030	0.001	>100	7.7	0.028	1.9	0.006	0.036
-2	3.7	0.005	1.0	5.1	0.009	0.30	3.5	0.025	0.010	>100	5.9	0.021	0.78	0.011	0.065
-3	0.48	0.020	1.1	4.3	0.005	0.25	2.9	0.017	0.004	>100	7.6	0.009	1.9	0.009	+ 0.034
-4	0.50	0.011	1.1	8.9	0.006	0.40	3.8	0.032	0.004	>100	7.7	0.003	2.1	0.008	0.039
NN	1.0	0.002	0.043	0.85	0.010	0.28	0.26	0.12	0.005	>100	2.3	0.052	48.8	0.006	0.019
	1.3	0.001	0.800	5.7	0.008	0.75	0.55	0.008	0.006	>100	1.6	0.051	46.8	0.007	0.025
00	0.24	0.018	1.1	3.3	0.004	0.37	1.3	0.020	0.001	52	5.2	0.053	4.6	0.004	0.020
•	0.15	0.018	1.1	3.5	0.004	0.10	0.88	0.020	0.012	66	6.3	0.036	0.66	0.004	0.020
h															
PPp	0.052	0.004	0.017	0.021	0.007	0.009	0.0004	0.002	0.001	0.196	0.059	0.002	0.08	0.0001	0.006

Metals concentration below instrument detection limit - see Table 18 for detection limits.

<sup>&</sup>lt;sup>b</sup>Secondary effluent sample only.

<sup>&</sup>lt;sup>C</sup>Secondary effluent samples inadvertently collected prior to settling pond.

TABLE 18. LOWER DETECTION LIMIT OF METALS ANALYSIS SYSTEMS

		lyzed by ICAPa	
	Detection		Detection
Metal	limit, g/m <sup>3</sup>	Metal	limit, g/m <sup>3</sup>
Aluminum	0.050	Manganese	0.005
Arsenic	0.005	Molybdenum	0.0006
Barium	0.0002	Nickel	0.01
Boron	0.0001	Phosphorus	0.01
Cadmium	0.0005	Sodium	0.050
Calcium	0.0002	Silicon	0.003
Chromium	0.0002	Silver	0.0050
Cobalt	0.0005	Tin	0.001
Copper	0.0002	Titanium	0.001
Iron	0.005	Vanadium	0.002
Lead	0.001	Zinc	0.025
Magnesium	0.001		
Metals	analyzed by	atomic absorp	tion
Antimony	0.0005	Selenium	0.005
Beryllium	0.0001	Thallium	0.005
Mercury	0.0005		2.000

al g/m<sup>3</sup> equals 1 mg/l.

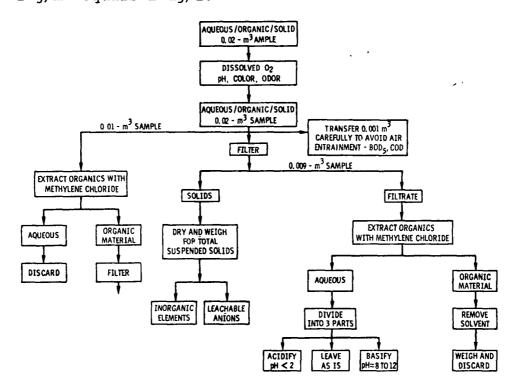


Figure 16. EPA-recommended field handling scheme for liquid/slurry samples (1).

The 0.01 m<sup>3</sup> portion for organic analysis is extracted in the field with methylene chloride. The aqueous portion is discarded; the methylene chloride extract is filtered and sent to the laboratory for organic chemical analysis.

Because textile wastewaters from stable emulsions with methylene chloride, the field handling procedure was modified, with the approval of the EPA-Process Measurements Branch, IERL/RTP. It was not feasible to conduct methylene chloride extractions in the field because previous experience has shown that it requires from 1 hr to 3 hr of concerted effort to break these emulsions. Centrifugation was necessary occasionally. The modified field handling scheme for textile wastewater is shown in Figure 17. The fundamental difference between the two schemes is that the field analysis of some of the common wastewater chemical species is performed on unextracted water samples. Cyanide samples were not extracted before analysis.

The filter obtained from the field was dried and the concentration of total suspended solids determined. Leachate analyses were not performed because each textile plant sampled was meeting its EPA effluent standard for TSS at the time samples were collected. The filter paper was asked by means of low temperature plasma, digested and analyzed for metals by spark source mass spectrometry (SSMS) and conventional atomic absorption (AA). Level 1 chemical analysis protocoal requires SSMS for metals, analysis and AA for those metals not accurately detected by SSMS.

SSMS can be used for analysis of nonvolatile compounds, such as inorganic solids and trace elements. The spark produces ions from the sample by high voltage breakdown across two electrodes. One electrode usually consists of or contains the sample material. In the spark ion source, the spark is sustained between ions from the rf spark source are accelerated through a potential field and focused with dual collimating slits. The ion beam passes into the mass spectrometer where the ions are separated. Photographic plates are used to record the emissions spectra for various mass fractions that correspond to specific elements or compounds.

A detailed flow diagram for Level 1 organic analysis is shown in Figure 18 (2). First, 0.01 m<sup>3</sup> of the sample is extracted with methylene chloride. The aqueous phase is saved and the organic phase separated for analysis.

The modification to the Level 1 organic analysis scheme, as recommended by the EPA, was employed in addition to the basic Level 1 chemical analysis procedure.

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Figure 17. MRC-modified Level 1 field sampling and analysis scheme.

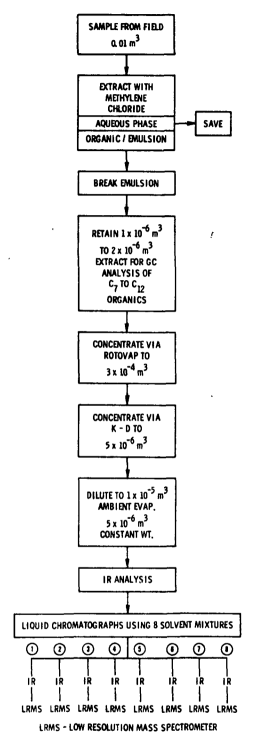


Figure 18. Level 1 organic analysis scheme (2).

Funds to perform the additional methods development required for the modification were supplied under another EPA contract.<sup>a</sup>

The principal feature of this modification is that the organic extract is not evaporated to dryness prior to liquid chromatography (LC) and/or low resolution mass spectrometry (LRMS). Two new sample handling steps are required:

- A quantitative assay procedure using GC/MS to complement gravimetric analysis and give quantitative data on  $C_7-C_{1\,2}$  hydrocarbons.
- A solvent exchange step to transfer the sample from methylene chloride extract to nonpolar solvent and allow lower boiling material (less than C<sub>7</sub>) to pass through the chromatograph and subsequent steps.

The new flow diagram incorporating these changes is shown in Figure 19.

## Results of Inorganic Analyses

The Level 1 chemical analysis scheme is divided into inorganic and organic analysis. Inorganic analysis includes:  $BOD_5$ , COD, TSS, TDS, metals analysis by spark source mass spectrometer (SSMS) of filtered solids and the filtrate, field analysis of selected species, and total nonvolatile organic concentration.

Table 19 shows the concentrations of the following parameters in 15 plant effluents:  $BOD_5$ , COD, TSS, TDS, and total organic concentration. Data from the field analysis of effluent samples are found in Table 20. Metals concentrations of the suspended solids collected on the filter paper are listed in Table 21. Results of SSMS analyses of the filtrate are given in Table 22. Concentrations are reported as  $g/m^3$  (ppm) of textile effluent sample. All elements for which values are not entered have concentrations below the detection limit.

### Results of Organic Analyses

As illustrated in Figures 17 and 19, there are four points within the Level 1 chemical analysis scheme where organic analysis takes place. At each plant one portion of the sample was filtered and a part of this volume was extracted with methylene chloride. The aqueous phase was used for metals analysis by SSMS, and the organic phase was dried and weighed to determine the concentration of total methylene chloride extractable organics. Results of these analyses are shown in Table 23.

<sup>&</sup>lt;sup>a</sup>Contract No. 68-02-1411, Task 19, "Analysis Support of Textile Environmental Assessments."

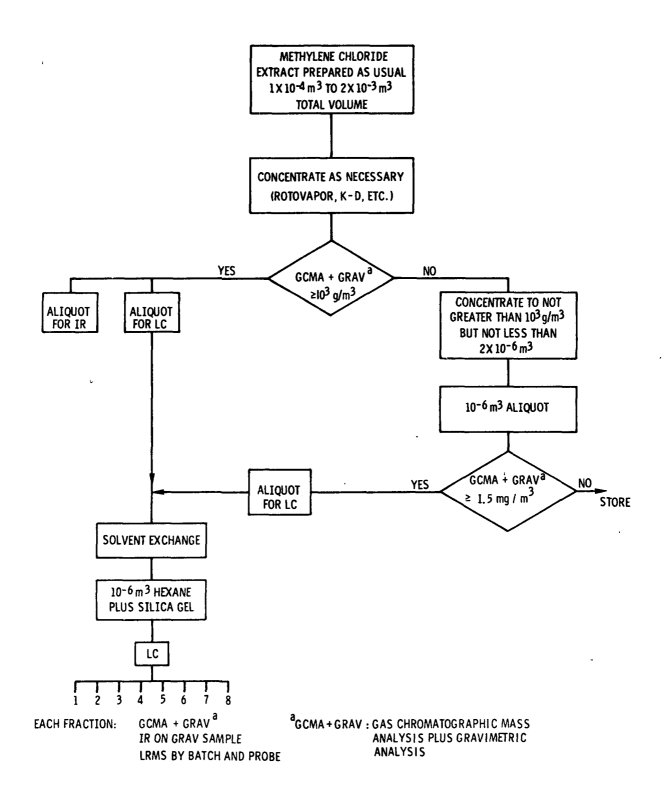


Figure 19. EPA, IERL/RTP modification to Level 1 organics analysis procedure.

TABLE 19. SELECTED PARAMETERS OF TEXTILE EFFLUENT SAMPLES FROM THE INORGANIC SEGMENT OF LEVEL 1 CHEMICAL ANALYSIS PROTOCOL

Plant code	5-day biochemical oxygen demand, g/m <sup>3ª</sup>	Chemical oxygen demand g/m <sup>3</sup>	Total suspended solids, g/m <sup>3</sup>	Total dissolved solids, g/m <sup>3</sup>	Total organic concentration, g/m <sup>3</sup>
A	168	1652	234	1,725	63.7
В	b	99	7	1,681	3.18
С	25	396	24	2,924	28.2
E	<5	78	10	13,120	3.60
F	69	276	10	2,006	16.0
G	42	502	5	276	27.2
K	<5	131	14	1,256	2.73
L	13	234	42	725	18.3
N	36	286	13	1,352	9.24
s	59	1035	349	692	5.40
T	32	414	44	660	17.8
U	24	748	111	1,331	14.6
v	<5	128	b	b	; <b>b</b>
W	84	837	217	1,648	15.0
x	15	, 258	1.3	437	13.5

b d g/m³ equals 1 mg/l.
bAnalysis not performed.

Portions (1 to 2 x  $10^{-6}$  m<sup>3</sup>) of each sample were then analyzed by gas chromatography (GC) to determine the concentration of C<sub>7</sub> to C<sub>12</sub> hydrocarbons. The Following GC columsn were used: a) 1.8 m x 3.2 mm stainless steel column packed with 10% VC, W98 on 80 to 100S, and b) 1.8 m x 3.2 mm stainless steel column packed with 10% SP-2100. Each column was held at 50°C for 4 min, then programmed at  $16^{\circ}$ C/min to 250°C and held at 250°C for 4 min.

Results of these analyses indicated that in 13 of the 14 samples the  $C_7$  to  $C_{12}$  hydrocarbon concentrations were below the threshold detection limit of 1.0 g/m³ (ppm). The secondary effluent sample from plant X contained a total concentration of about 3.0 g/m³ of  $C_7$  to  $C_{12}$  hydrocarbons.

TABLE 20. FIELD ANALYSIS OF SELECTED SECONDARY WASTEWATER PARAMETERS ON ILTERED, UNEXTRACTED SAMPLES AS PER LEVEL 1 ANALYSIS PROTOCOL (2).

					Wate	r paramete:	rs, g/m36						
Plant code	Color, APHA	Specific conductivity b	Nitrate	Nitrite	Hydrogen sulfide	Sulfate	Methyl orange acidity	Dissolved oxygen	Ammonia	o-phosphate	Total alkalinity	Chromium	
A	2,000	1,500	1.9	0.06	4	8.5	0	5.5	12.8	1.0	100	0.18	7.
В	90	1,200	0.002	<0.005	0.20	368	o	7	2.5	7.3	5.5	0.004	7.
С	1,920	2,400	23.3	4.64	5	40	0	6	3.4	1.08	8.3	0.031	10
E	0	310	79.2	0.016	0.1	12	0	8.5	3.4	2.0	35	0.004	7.
P	80	1,900	0	0.043	0.1	10	0	5	1.54	0.56	2.4	0.004	7.4
G	500	155	1.32	0.076	<2	56	o	8	72.5	4	30	0.003	7.
ĸ	270	875	4.4	0.056	2	>1	0	5	3.1	2.5	710	0.004	7.
L	370	555	13.5	0.864	3	460	0	4	0.5	0.88	30	0.03	5.8
N	90	990	5.5	0.003	0.1	640	20	9	12.8	11.2	ο .	1.8	7.0
s	240	640	4.4	0.033	0.3	150	0	7	72.5	72	130	0	7.8
T	350	460	0.8	0.04	6	100	0	8	13.6	6.4	300	0	7.4
υ	2,480	770	0.8	<0.005	3.5	0	0	9	5.44	2.96	120	0.014	7.
٧	500	360	0.88	0.264	0.5	. 57	0	9	2.5	1.7	0.4	0.003	7.1
W	1,900	1,250	12.3	0.145	0.1	0	0	5	0.38	0.075	950	0.003	8.
x	>10	285	0.033	0.44	0.01	ı	0	7.2	0.05	0	140	0.039	7.2

 $<sup>\</sup>frac{a}{1}$  g/m<sup>3</sup> equals 1 mg/1.

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Units umhos at 25°C.

TABLE 21. LEVEL 1 SPARK SOURCE MASS SPECTROMETER METALS ANALYSIS OF THE SUSPENDED SOLIDS COLLECTED ON THE FILTER PAPER DURING FIELD FILTRATION

Ple	ant A	Concentration: 1	ng/m³	Detection lim	it: 0.01	ma∕w <sub>3</sub>	
Uranium	<0.04	Terbium	<0.03	Ruthenium		Vanadium	318
Thorium	<0.04	Gadolinium	<0.04	Molybdenum	17	Titanium	9.5
Bismuth	3.0	Europium	<0.04	Niobium	0.03	Scandium	0.0
Lead	42	Samarium	<0.07	Zirconium	1.9	Calcium	1,220
Thallium	<0.04 a	Neodymium	<0.09	Yttrium	0.08	Potassium	265
Mercury	_°	Praseodymium	<0.05	Strontium	6.0	Chlorine	15
Gold		Cerium	1.3	Rubidium	1.0	Sulphur	1,987
Platinum		Lanthanum	0.91	Bromine	3.0	Phosphorus	1,590
Iridium		Barium	15	Selenium	0.25	Silicon	450
Osmium	b	Cesium		Arsenic	3.4	Aluminum	874
Rhenium	-	Iodine	0.33	Germanium	0.03	Magnesium	702
Tungsten	<0.08	Tellurium		Gallium	0.13	Sodium	4,636
Pantalum	<0.03	Antimony	1.6	Zinc	300	Pluorine	28
lafnium	<0.11	Tin	13 _b	Copper	172	Oxygen	-
Lutecium		Indium		Nickel	26	Nitrogen	-
(tterbium	<0.04	Cadmium	2.5	Cobalt	2.0	Carbon	_
Chullium	<0.03	Silver	1.1	Iron	582	Boron	
Erbium	<0.03	Palladium		Manganese	11	Beryllium	<0.0
lolmium		Rhodium		Chromium	56	Lithium	1.3
ysprosium	<0.01						
	Plant B	Concentration	n: mg/m³	Detection	limit: 0.	01 g/m <sup>3</sup>	
<b>Jranium</b>		Terbium		Ruthenium		Vanadium	0.0
Thorium		Gadolinium		Molybdenum		Titanium	0.3
3ismuth	0.05	Europium		Niobium	0.04	Scandium	
ead	0.2	Samarium		Zirconium		Calcium	30
hallium		Neodymium	0.1	Yttrium		Potassium	11
lercury		Praseodymium		Strontium	0.1	Chlorine	0.3
old	•	Cerium	0.02	Rubidium	0.08	Sulphur	24
latinum		Lanthanum	0.01	Bromine	0.02	Phosphorus	120
Cridium		Barıum	1.8	Selenium		Silicon	340
Osmium	_	Cesium		Arsenic		Aluminum	25
Rhenium	_b	Iodine		Germanium		Magnesium	130
Cungsten		Tellurium		Gallium		Sodium	240
'antalum	0.003 <sup>C</sup>	Antimony		Zinc	1.8	Fluorine	6.8
lafnıum	<0.02	Tin	0.01	Copper	0.3	Oxygen	_
utecium		Indium	_ь	Nickel	0.07	Nitrogen	_
tterbium'		Cadmium		Cobalt	•	Carbon	_
hullium		Silver		Iron	13	Boron	_
rbium		Palladium		Manganese	0.3	Beryllium	
olmium		Rhodium		Chromium	0.5	Lithium	0.0
ysprosium							
Pla	int C	Concentration: m	g/m³	Detection li	mit: 0.05	mg/m³	
Jranium		Terbium		Ruthenium		Vanadium	1.1
horium		Gadolinium	0.07	Molybdenum	0.1	Titanium	0.8
ismuth	0.7	Europium		Niobium		Scandium	0.1
ead	0.6	Samarium	0.07	Zirconium	0.2	Calcium	83
hallıum	0.07	Neodymium		Yttrium	0.03	Potassium	330
ercury	_a	Praseodymium		Strontium	1.6	Chlorine	43
old		Cerium	0.1	Rubidium	2.8	Sulphur	110
latinum		Lanthanum	0.07	Bromine	1.1	Phosphorus	700
ridium		Barium	57	Selenium		Silicon	1,100
		Cesium	0.03	Arsenic	0.07	Aluminum	530
smium	_ь	Iodine	0.03	Germanium		Magnesium	160
	0.07	Tellurium		Gallium		Sodium	2.800
henium	0.07		0.1	Zinc	29	Fluorine	33
smium henium ungsten antalum	0.01c	Antimony		~			33
henium ungsten antalum	0.07 0.01 0.1	Antimony Tin	0.2.	Copper	4	Ovuden	
henium ungsten antalum afnium	0.01 <sup>c</sup> 0.1	Tin	0.2 <sub>b</sub>	Copper	4 0 9	Oxygen	•
henium ungsten antalum afnium utecium	0.1	Tin Indium	0.2 <sub>b</sub>	Nickel	0.9	Nitrogen	
henium ungsten antalum afnium utecium tterbium		Tin Indium Cadmium	0.2 <sub>b</sub> 0.07	Nickel Cobalt	0.9 0.07	Nitrogen Carbon	
henium ungsten antalum afnium utecium tterbium hullium	0.1	Tin Indium Cadmium Silver	0.2 <sub>b</sub>	Nickel Cobalt Iron	0.9 0.07 20	Nitrogen Carbon Boron	•
henium ungsten antalum afnium utecium tterbium hullium	0.1	Tin Indium Cadmium Silver Palladium	0.2 <sub>b</sub>	Nickel Cobalt Iron Manganese	0.9 0.07 20 0.9	Nitrogen Carbon Boron Beryllium	
nenium ungsten antalum afnium utecium tterbium nullium	0.1	Tin Indium Cadmium Silver	0.2 <sub>b</sub>	Nickel Cobalt Iron	0.9 0.07 20	Nitrogen Carbon Boron	0.:

aNot reported. bInternal standard. CInstrument source.

TABLE 21 (continued)

		oncentration: mg/	m <sup>3</sup>	Detection lim	it: <0.02 r	ng/m³	
Pla	nt B C	OHOBICE GEORGE		Dukhan i		Vanadium	
ranium		Terbium		Ruthenium Molybdenum	0.04	Titanium	0.05
horium		Gadolinium		Niobium	••••	Scandium	<0.04
smuth	0.07	Europium		Zirconium	0.02	Calcium	3.9
ad	0.09	Samarium		Yttrium	• • • •	Potassium	110
nallium	• • • • • • • • • • • • • • • • • • • •	Neodymium	0.03	Strontium	0.32	Chlorine	18
ercury	_a	Praseodymium	0.02	Rubidium	0.49	Sulphur	0.74
old	-	Cerium	0.02 0.02	Bromine	2.8	Phosphorus	0.04
Latinum		Lanthanum	0.02	Selenium		Silicon	3.5
cidium		Barium	0.05	Arsenic	<0.07	Aluminum	18
mium		Cesium	0.16	Germanium		Magnesium	5.8
nenium	_b	Iodine	0.10	Gallium		Sodium	972
ngsten	0.09	Tellurium	0.02	Zinc	0.12	Fluorine	11
intalum		Antimony	0.04	Copper	0.04	Oxygen	_ <u>a</u>
fnium	<0.05	Tin	0.04 b	Nickel	0.05	Nitrogen	_a
itecium		Indium	0.02	Cobalt		Carbon	_a_
terbium	<0.04	Cadmium	0.02	Iron	0.94	Boron	_a
hullium	• • • • • • • • • • • • • • • • • • • •	Silver		Manganese	0.05	Beryllium	
rbium		Palladium		Chromium	0.07	Lithium	0.02
olmium		Rhodium		Curomiam	••••		
ysprosium			_3	Detection 1	imit: 0.05	mq/m³	<del></del>
Plan	t P C	oncentration: mg/	m			Vanadium	0.1
		Terbium		Ruthenium		Titanium	1.4
ranium		Gadolinium		Molybdenum	0.1	Scandium	<0.1
horium		Europium		Niobium			120
ismuth				Zirconium	0.4	Calcium	87
ead	4.0	Samarium		Yttrium		Potassium	34
hallium	_a	Neodymium		Strontium	0.6	Chlorine	106
lercury		Praseodymium	0.1	Rubidium	0.1	Sulphur	290
old		Cerium	0.05	Bromine	1.6	Phosphorus	820
Platinum		Lanthanum	4.3	Selenium		Silicon	42
Cridium		Barium	4.3	Arsenic	0.5	Aluminum	30
Osmium		Cesium		Germanium		Magnesium	
Rhenium	_ь	Iodine		Gallium		Sodium	1,100
rungsten	_	Tellurium		Zinc	39	Fluorine	82
rantalum	0.002 <sup>C</sup>	Antimony	0.4	Copper	· 29	Oxygen	-
Hafnium	<0.1	Tin	1.3 <sub>b</sub>	Nickel	1.2	Nitrogen	-
Lutecium		Indium		Cobalt	0.1	Carbon	-
Ytterbium		Cadmium	0.2	Iron	67	Boron	-
Thullium		Silver		Manganese	1.1	Beryllium	_
Erbium		Palladium		Chromium	2.0	Lithium	-
Holmium		Rhodium		Curomram			
Dysprosium					0.03	ma/m³	
_		oncentration: mg/	m <sup>3</sup>	Detection 1	mit: U.UI	ma m	
Plan	nt K C	Oncentration				Vanadium	0.00
		Terbium		Ruthenium	0.06	Titanium	0.0
ranium	•	Gadolinium		Molybdenum		Scandium	<0.0
horium		Europium		Niobium	0.06	Calcium	100
ismuth		Samarium		Zirconium	0.00	Potassium	23
ead	0.51	Neodymium		Yttrium	0.06	Chlorine	1.2
challium	0.01	Praseodymium	0.01	strontium	0.3	Sulphur	18
dercury	-		0.03	Rubidium		Phosphorus	9.7
Gold		Cerium	0.04	Bromine	0.03	Silicon	23
Platinum		Lanthanum	0.84	Selenium	.0.04	Aluminum	45
Iridium		Barium		Arsenic	<0.04	Magnesium	7.2
Osmium		Cesium	0.03	Germanium		Sodium	308
Rhenium	۵_	Iodine	0.05	Gallium		Fluorine	1.
Tungsten	0.06	Tellurium	0.01	Zinc	0.75	Oxygen	1.
Tantalum		Antimony		Copper	0.97	Nitrogen	
Hafnium	<0.03	Tin	0.04	Nickel	0.32	Carbon	
		Indium		Cobalt			
Lutecium		Cadmium		Iron	3.2	Boron	
Ytterbium		Silver		Manganese	0.1	Beryllium	0.
Thullium		Palladium		Chromium	0.81	Lithium	•
Erbium		Rhodium		CITE OME			
							ntinued)
Holmium Dysprosium						I CC	III C TII G C C \

TABLE 21 (continued)

Pla	nt L Co	ncentration: m	q/m³	Detection	limit: 0.05	mg/m <sup>3</sup>	
Jranium		Terbium		Ruthenium		Vanadium	1.0
Thorium		Gadolinium		Molybdenum	0.1	Titanium	3.2
Bismuth	0.05	Europium		Niobium		Scandium	<0.1
Lead	9	Samarium	0.3	Zirconium	0.3	Calcium	1,200
Phallium	_	Neodymium	0.05	Yttrium		Potassium	95
dercury	<sub>M</sub> a	Praseodymium	0.05	Strontium ·	1.0	Chlorine	37
Gold		Cerium	0.2	Rubidium	0.3	Sulphur	360
Platinum		Lanthanum	0.1	Bromine	0.4	Phosphorus	1,200
Iridium		Barium	23	Selenium		Silicon	4,600
Demium		Cesium		Arsenic		Aluminum	440
Rhenium	طـ	Iodine		Germanium		Magnesium	650
<b>Tungsten</b>	_	Tellurium		Gallium		Sodium	5,000
antalum	0.001 <sup>c</sup>	Antimony	0.2	Zinc	1,900	Fluorine	33
lafnium	<0.1	Tim	٥٠٤	Copper	31	Oxygen	-
Lutecium		Indium		Nickel	3.6	Nitrogen	-
tterbium		Cadmium	0.05	Cobalt	0.05	Carbon	-
hullium		Silver	,	Iron	380	Boron	-
Erbium	Palladiu	m Palladium		Manganese	2.2	Beryllium	
lolmium <sub>.</sub>		Rhodium		Chromium	13	Lithium	0.3
ysprosium							
Pla	nt N Co	ncentration: m	g/m³	Detection	limit: 0.04	mg/m³	
		~		Double and some		**	
Jranium		Terbium	0.00	Ruthenium		Vanadium	0.4
Chorium	0.04	Gadolinium	0.08	Molybdenum	0.2	Titanium Scandium	5.6
Sismuth	0.04	Europium	<0.04	Niobium	0.6		<0.2
ead	4.0	Samarium	0.1 0.1	Zirconium		Calcium	280
hallium	ھ	Neodymium	0.1	Yttrium	0.4	Potassium	44
lercury		Praseodymium	0.5	Strontium	0.8 0.08	Chlorine	3.8 1,240
old		Cerium Lanthanum		Rubidium		Sulphur	2,000
latinum			0.1 14	Bromine	0.1 0.04	Phosphorus	2,000
iridium		Barium		Selenium		Silicon	
Smium	ەلـ	Cesium		Arsenic	0.8	Aluminum Magnesium	200 110
Chenium		Iodine	0.4	Germanium	0.04	Sodium	520
Cungsten	0.04 0.003 <sup>c</sup>	Tellurium :	0.2	Gallium Zinc	150	Fluorine	76
antalum afnıum	<0.08	Tin		Copper	12	Oxygen	76
utecium	10.08	Indium	0.3 <sub>b</sub>	Nickel	2.9	Nitrogen	
tterbium	<0.08	Cadmium	0.08	Cobalt	1.7	Carbon	-
hullium	10.00	Silver	V. 50	Iron	600	Boron	
rbium		Palladium		Manganese	12	Beryllium	
lolmium		Rhodium		Chromium	680	Lithium	0.2
ysprosium		KNOUTUM		CITOMIAN	000	DICHIUM	0.2
				<del></del>			
Pla	nt S Co	ncentration: m	g/m <sup>3</sup>	Detection	n limit: 0.0	1 mg/m <sup>3</sup>	
ranium	0.08	Terbium		Ruthenium	1	Vanadium	6.1
horium	0.21	Gadolinium	0.04	Molybdenum	6.5	Titanium	40
ısmuth	2.4	Duropium	0.05	Niobium	0.1	Scandium	0.1
ead	7.1	Samarium	0.09	Zirconium	1.6	Calcium	4,353
hallıum	<0.02 <sub>a</sub>	Neodymium	1.0	Yttrium	0.33	Potassium	85 <b>9</b>
ercury		Praseodymium	0.36	Strontium	6.8	Chlorine	129
old		Cerium	3.4	Rubidium	11	Sulphur	1,882
latinum		Lanthanum	4.0	Bromine	14	Phosphorus	4,824
rıdıum		Barıum	49	Selenium	<0.22	Silicon	1,106
sm1um	_b	Cesium	0.99	Arsenic	7.0	Aluminum	2,000
henium		Iodine	0.38	Germanium	0.16	Magnesium	1,035
ungsten	<0.11 <sub>c</sub>	Tellurium		Gallium	0.56	Sodium	694
antalum	0.55°	Antimony	188	Zinc	13	Fluorine	56
afnium	<0.13	Tin	31 _b	Copper	306	Oxygen	•
utecium	<0.02	Indium		Nickel	16	Nitrogen	-
tterbium	<0.07	Cadmium	1.1	Cobalt	0.44	Carbon	-
	<0.02	Silver	0.25	Iron	2,588	Boron	_ •
hullium				Manganese	56	Beryllium	<0.0
hullium rbium	<0.04	Palladium					
nullium	<0.04 0.01 0.01	Rhodium		Chromium	26	Lithium	0.9

anot reported. bInternal standard. CInstrument source.

TABLE 21 (continued)

Pla	nt T	Concentration: mo	/m³	Detection	limit: 0.01	mg/m³	
Uranium		Terbium		Ruthenium		Vanadium	0.4
Thorium		Gadolinium		Molybdenum	0.2	Titanium	7,3
Bismuth		Europium		Niobium	<b></b>	Scandium	,,,,
Lead	53	Samarium		Zirconium	0.9	Calcium	1,500
Thallium		Neodymium		Yttrium	0.2	Potassium	330
Mercury	_8	Praseodymium		Strontium	1.6	Chlorine	11
Gold		Cerium	0.4	Rubidium	1.1	Sulphur	140
Platinum		Lanthanum	0.4	Bromine	1.8		1,300
Iridium		Barium	150	Selenium	1.0	Phosphorus Silicon	1.040
Osmium		Cesium	150	Arsenic		Aluminum	550
Rhenium	_b	Iodine		Germanium		Magnesium	530
		Tellurium		Gallium			2,700
Tungsten			0.7		98	Sodium	890
Tantalum		Antimony	0.7	Zinc	29	Fluorine	a ve
Hafnium		Tin	1.1 <sub>b</sub>	Copper		Oxygen	_a
Lutecium		Indium		Nickel	1.6	Nitrogen	_a
Ytterbium		Cadmium		Cobalt	0.2	Carbon	- <u>-</u> -
Thullium		Silver		Iron	320	Boron	
Erbium		Palladium		Manganese	4.9	Beryllium	
Holmium		Rhodium		Chromium	5.1	Lithium	0.7
Dysprosium							
	-4 11	Concontration	_/_1	Dot	Timino -0 1	11 == (= 1	· · · · · · · · · · · · · · · · · · ·
Pla	int U	Concentration: m	g/m³	Detection	11m1t: <0.0	ol mg/m <sup>3</sup>	<del></del>
Uranium	<0.02	Terbium		Ruthenium		Vanadium	0.49
Thorium	<0.04	Gadolinium		Molybdenum	5.2	Titanium	18
Bismuth	<0.05	Europium	0.02	Niobium	٥٠٠	Scandium	0.1
Lead	7.4	Samarium	0.02	Zirconium	2.2	Calcium	756
Thallium	<0.02	Neodymium	0.02	Yttrium	0.16	Potassium	402
Mercury	`a	Praseodymium	0.01	Strontium	6.4	Chlorine	12
Gold	-	Cerium	0.62	Rubidium	3.3		2,680
		Lanthanum	0.38	Bromine		Sulphur	
Platinum		Barium			5.5	Phosphorus	597
Iridium			12 0.01	Selenium	0.09	Silicon	378
Osmium	_b	Cesium		Arsenic	<0.13	Aluminum	1,220
Rhenium	<del>-</del>	Iodine	0.28	Germanium		Magnesium	116
Tungsten	<0.07	Tellurium		Gallium	0.18	Sodium	1,950
Tantalum	<0.23°	Antimony	1.7	Zinc	60	Fluorine	96 a
Hafnium	<0.09	Tin	4.3 <sub>b</sub>	Copper	29	Oxygen	-ā
Lutecium		Indium		Nickel	11	Nitrogen	-,"
Ytterbium	<0.06	Cadmium	0.44	Cobalt	0.20	Carbon	-"
Thullium		Silver	0.23	Iron	560	Boron	~~
Erbium	<0.04	Palladium		Manganese	12	Beryllium	
Holmium		Rhodium		Chromium	23	Lithium	0.40
Dysprosium							•
Pla	nt W	Concentration: mg	/m <sup>3</sup>	Detection	limit: 0.05	ma /m 3	
		COCOCIGOTO	<del>/</del>	Detection	111111111111111111111111111111111111111	ш9/ ш-	
Uranıum		Terbium	0.1	Ruthenium		Vanadium	15
Thorium	0.5	Gadolinium	0.5	Molybdenum	0.5	Titanium	160
Bismuth	<0.2	Europium	0.2	Niobium	0.2	Scandium	0.4
Lead	14	Samarium	<0.9	Zirconium	1.9	Calcium	15,000
Thallium	<0.1	Neodymium	1.5	Yttrium	1,1	Potassium	1,500
Mercury	_a	Praseodymium	1.3	Strontium	19	Chlorine	290
Gold		Cerium	8.0	Rubidium	3.1	Sulphur	330
Platinum		Lanthanum	2.3	Bromine	2.5	Phorphorus	1,700
Iridium		Barium	230	Selenium	-,,	Silicon	2,600
Osmium		Cesium	0.1	Arsenic	13	Aluminum	6,700
Rhenium	_b	Iodine	0.07	Germanium		Magnesium	3,700
Tungsten	<b>-0</b> 2	Tellurium	0.07	Gallium	2.3		
Tantalum	0.006 <sup>C</sup>		0.4			Sodium	3,300
	0.000	Antimony	2 - 5	Zinc	66	Fluorine	390 a
Hafnium	<0.3	Tin	1.3 <sub>b</sub>	Copper	23	Oxygen	- <u>-</u> a
Lutecium	-0.0	Indium	<u>-</u>	Nickel	17	Nitrogen	_a _a
Ytterbium	<0.2	Cadmium	0.3	Cobalt	8.0	Carbon	_a _a
Thullium		Silver	0.07	Iron	4,500	Boron	
Erbium	<0.1	Palladium		Manganese	370	Beryllium	0.2
Holmium		Rhodium		Chromium	10	Lithium	13
Dysprosium							
3							

a Not reported. bInternal standard. CInstrument source.

TABLE 21 (continued)

Pla	nt X	Concentration: mg	/m <sup>3</sup>	Detection	limit: 0.03	mg/m <sup>3</sup>	
Uranium		Terbium		Ruthenium		Vanadium	0.4
Thorium		Gadolinium	<0.06	Molybdenum	0.1	Titanium	4.7
Bismuth	<0.09	Europium		Niobium	0.06	Scandium	0.06
Lead	5.0	Samarium	<0.06	Zirconium	0.4	Calcium	1,200
Thallium	<0.06	Neodymium	0.06	Yttrium	0.08	Potassium	53
Mercury	_a	Praseodymium	0.06	Strontium	1.3	Chlorine	17
Gold		Cerium -	0.2	Rubidium	0.4	Sulphur	110
Platinum		Lanthanum	0.09	Bromine	17	Phosphorus	2,400
Irıdium		Barium	22	Selenium		Silicon	880
Osmium		Cesium		Arsenic	<0.06	Aluminum	640
Rhenium	-р	Iodine	0.09	Germanium		Magnesium	
Tungsten	0.06	Tellurium		Gallium	0.06	Sodium	2,200
Tantalum	0.08 <sup>€</sup>	Antimony	2.4	Zinc	410	Fluorine	38 _
Hafnium	<0.1	Tin	0.44	Copper	27	Oxygen	9
Lutecium		Indium	_b	Nickel	7	Nitrogen	
Ytterbium	<0.09	Cadmium	0.06	Cobalt	2.7	Carbon	-3
Thullium		Sılver		Iron	1,300	Boron	_4
Erbium	<0.06	Palladium		Manganese	2.5	Beryllium	
Holmium Dysprosium		Rhodium		Chromium	12	Lithium	0.1

a<sub>Not</sub> reported. bInternal standard. CInstrument source.

TABLE 22. LEVEL 1 SPARK SOURCE MASS SPECTROMETERS METALS ANALYSIS OF FILTERED SECONDARD EFFLUENT (detection limit of 0.001 g/m³)

			Plant A,	g/m <sup>3</sup>			
Uranium	<0.003	Terbium		Ruthenium		Vanadium	2.5
Thorium	<0.003	Gadolinium	<0.003	Molybdenum	0.043	Titanium	0.087
Bismuth	<0.006	Europium	<0.002	Niobium		Scandium	<0.006
Lead	0.38	Samarium	<0.002	Zirconium	0.031	Calcium	170
Thallium	<0.004 <sub>a</sub>	Neodymium	<0.004	Yttrium	0.003	Potassium	8.8
Mercury	_a	Praseodymium	0.009	Strontium	0.70	Chlorine	11
Gold		Cerium	0.002	Rubidium	0.19	Sulphur	130
Platinum		Lanthanum	0.004	Bromine	0.47	Phosphorus	6
Iridium		Barium	0.51	Selenium	0.003	Silicon	18
Osmium	. р	Cesium	0.003	Arsenic	<0.015	Aluminum	6.4
Rhenium	-	Iodine	0.011	Garmanium		Magnesium	10
<b>Tungsten</b>	<0.008_	Tellurium		Gallium		Sodium	180
<b>Fantalum</b>	0.008	Antimony	0.095	Zinc	13	Fluorine	1.1
Hafnium	<0.010	Tin	0.10 <sub>b</sub>	Copper	0.14	Oxygen	~~
Lutecium	<0.002	Indium		Nickel	1.0	Nitrogen	_8
Ytterbium	<0.007	Cadmium	0.005	Cobalt	0.021	Carbon	_a
Thullium	<0.002	Silver	0.001	Iron	6.1	Boron	
Erbium	<0.007	Palladium		Manganese	0.48	Beryllium	<0.003
Holmium		Rhodium		Chromium	1.4	Lithium	0.12
ysprosium							
		<del>, , , , , , , , , , , , , , , , , , , </del>	Plant B, g	/m³			
				/ <del></del>			
Jranium	<0.003	Terbium		Ruthenium		Vanadium	0.005
Chorium	<0.003	Gadolinium	<0.003	Molybdenum	<0.007	Titanium	0.079
Bismuth	<0.005	Europium	<0.002	Niobium	_	Scandium	<0.003
Lead	0.17	Samarium	<0.002	Zirconium	0.004	Calcium	21
Thallium	<0.002	Neodymium	<0.003	Yttrium	0.002	Potassium	79
dercury	_4	Praseodymium	<u>&lt;0.002</u>	Strontium	0.22	Chlorine	2.1
Gold		Cerium	<b>₹0.</b> 003	Rubidium	0.25	Sulphur	300
Platinum		Lanthanum	₹0.002	Bromine	0.26	Phosphorus	45
Iridium	•	Barium	0.20	Selenium	0.006	Silicon	8.0
Osmium	_b	Cesium	0.001	Arsenic	0.28	Aluminum	0.96
Rhenium	_	Iodine	0.004	Germanium		Magnesium	14
<b>lungsten</b>	<0.007	Tellurium		Gallium		Sodium	190
<b>Tantalum</b>	0.10 <sup>C</sup>	Antimony	0.019	Zinc	1.2	Fluorine	3.7
Hafnium	<0.009	Tin	0.017	Copper	0.16	Oxygen	ړ
Lutecium		Indium	_	Nickel	0.027	Nitrogen	_6 _8
/tterbium	<0.006	Cadmium	0.006	Cobalt	0.008	Carbon	
Chullium	<0.002	Silver	0.001	Iron	2.4	Boron	
Erbium	<0.003	Palladium		Manganese	0.41	Beryllium	<0.003
Iolmium		Rhodium		Chromium	0.24	Lithium	0.032
Dysprosium							
			Plant C, g	/m 3			
ranium	<0.005	Terbium	<0.002	Ruthenium		Vanadium	4.4
Chorium	<0.005	Gadolinium	<0.004	Molvbdenum	0.011	Titanium	0.71
ismuth	<0.010	Europium	<0.003	Niobium	0.004	Scandium	<0.01
Lead	c0 25	Samarium	<0.006	Zirconium	0.009	Calcium	41
Challium	<0.007 <sub>a</sub>	Neodymium	<0.005	Yttrium	0.001	Potassium	77.7
lercury	a	Praseodymium	0.004	Strontium	1.1	Chlorine	2.3
old		Cerium	<0.003	Rubidium	0.38	Sulphur	340
latinum		Lanthanum	<0.004	Bromine	13	Phosphorus	20
ridium		Barium	1.1	Selenium	0.027	Silicon	27
smium		Cesium	0.001	Arsenic	<0.025	Aluminum	0.99
thenium	_6	Iodine	0.024	Germanium	0.004	Magnesium	1.8
ungsten		Tellurium	<0.002	Gallium		Sodium	370
Cantalum	<0.013 <0.045	Antimony	0.62	Zinc	1.2	Fluorine	1.2
	<0017	Tin	0.082	Copper	0.63	Oxygen	
lafnıum	<0.003	Indium	0.005	Nickel	0.45	Nitrogen	12 
lafnıum .utecıum		Cadmium	0.030	Cobalt	0.018	Carbon	
utecium	<0.011						
utecium !tterbium	<0.011 <0.003		10	Iron	2.1	Boron	`
utecium tterbium hullium	<0.003	Silver	10	Iron Manganese	2.1 0.18	Boron Bervllium	
utecium tterbium			10	Iron Manganese Chromium	0.18 0.053	Boron Beryllium Lithium	<0.005 0.044

anot reported. bInternal standard. CInstrument source.

TABLE 22 (continued)

			Plant E, g	/m³			
Uranium		Terbium		Ruthenium		Vanadium	0.018
Thorium		Gađolinium		Molybdenum	0.004	Titanium	0.13
Bismuth	<0.002	Europium		Niobium		Scandium	
Lead	0.063	Samarium	<0.002	Zirconium	0.009	Calcium	36
Thallium	<0.002	Neodymium ·	0.007	Yttrium		Potassium	29
Mercury	_a	Praseodymium	0.004	Strontium	0.094	Chlorine	1.3
Gold		Cerium	0.022	Rubidium	0.071	Sulphur	290 🐞
Platinum		Lanthanum	0.019	Bromine	0.92	Phosphorus	9.2
Iridium		Barium	0.29	Selenium	0.002	Silicon	2.9
Osmium	_b	Cesium		Arsenic	0.035	Aluminum	0.39
Rhenium	_	Iodine	0.011	Germanium		Magnesium	2.1
Tungsten	<0.003	Tellurium		Gallium		Sodium	70
<b>Tantalum</b>	0.001 <sup>c</sup>	Antimony	0.13	Zinc	0.76	Fluorine	13 .
Hafnium	<0.003	Tin	0.006	Copper	0.10	Oxygen	_ 25 _ 6 _ 6
Lutecıum		Indium	_0	Nickel	0.038	Nitrogen	-:
Ytterbium	<0.002	Cadmium	0.004	Cobalt	0.001	Carbon	-9
Thullium		Silver	0.001	Iron	0.86	Boron	-٩
Erbium	<0.002	Palladium		Manganese	0.035	Beryllium	
Holmium		Rhodium		Chromium	0.043	Lithium	0.014
Dysprosium		,					
			Plant F, g	/m3			
Jranium	<0.003	Terbium		Ruthenium		Vanadium	0.022
Thorium	<0.004	Gadolinium	<0.003	Molybdenum	0.005	Titanium	0.10
Bismuth	<0.007	Europium	<0.002	Niobium	0.003	Scandium	<0.01
Lead	0.033	Samarium	<0.002	Zirconium	0.011	Calcium	27
Thallium	<0.005	Neodymium	<0.004	Yttrium	0.011	Potassium	10
	\v. 003a	Praseodymium	0.002	Strontium	0:28	Chlorine	
Mercury Gold	_	Cerium		Rubidium	0.054		0.83
		Lanthanum	0.003			Sulphur	56
Platinum Iridium		Barium	0.002	Bromine	0.16	Phosphorus	38
			0.16	Selenium	0.006	Silicon	48
Osmium	_b	Cesium		Arsenic	<0.017	Aluminum	4.6
Rhenium	<0.009	Iodine	0.002	Germanium	0 001	Magnesium	15
Tungsten	<0.003°	Tellurium	<0.002	Gallium	0.001	Sodium	490
Tantalum	<0.003	Antimony (	0.19	Zinc	1.5	Fluorine	2.9
Hafnium			0.004	Copper	0.49	Oxygen	
Lutecium	<0.002	Indium		Nickel	0.024	Nitrogen	_;
Ytterbium	<0.007	Cadmium	0.004	Cobalt	0.021	Carbon	-;
Thullium	<0.002	Silver		Iron	2.7	Boron	
Erbium	<0.005	Palladium		Manganese	0.13	Beryllium	<0.00
Holmium		Rhodium		Chromium	0.015	Lithium	0.004
Dysprosium	<0.002						
			Plant G	, g/m <sup>3</sup>			
Uranıum		Terbium		Ruthenium	0.006	Vanadium	0.12
Thorium		Gadolinium		Molybdenum Niobium	0.006	Titanium	0.11
Bismuth	0.10	Europium		Zirconium	0 011	Scandium	16
Lead	0.10	Samarium		Yttrium	0.011	Calcium	16
Thallium	_a	Neodymium			0.089	Potassium	2.0
Mercury	_	Praseodymium Cerium	0.004	Strontium Rubidium	0.089	Chlorine	1.1 43
Gold Platinum		Lanthanum	0.004	Bromine	0.017	Sulphur Phosphorus	16
Iridium		Barium	0.23	Selenium	0.001	Silicon	8.7
osmium Osmium		Cesium	0.23	Arsenic	0.001	Aluminum	2.2
Rhenium	_b	Iodine	0.007	Germanium	0.014	Magnesium	1.4
	_	Tellurium	0.007	Gallium		Sodium	48
Cungsten			1.2	Zinc	0.04		40
lantalum		Antimony	1.4		0.84	Fluorine	3.2 <sub>a</sub>
Hafnium		Tin	0.032	Copper	0.11	Oxygen	-a
Lutecium		Indium	_	Nickel	0.038	Nitrogen	_a _a _a _a
tterbium		Cadmium	0.002	Cobalt	0.13	Carbon	-a
Chullium		Silver	0.003	Iron	1.6	Boron	-"
		Palladium		Manganese	0.17	Beryllium	
Erbium Holmium Dysprosium		Rhodium		Chromium	0.018	Lithium	0.32

a<sub>Not</sub> reported. b<sub>Internal</sub> standard. c<sub>Instrument</sub> source.

TABLE 22 (continued)

			Plant 1	K, g/m <sup>3</sup>			7-11-17
Uranium	<0.002	Terbium		Ruthenium		Vanadium	0.01
Thorium	<0.002	Gadolinium	<0.002	Molybdenum	0.006	Titanium	0.01
Bismuth	<0.004	Europium	<0.002	Niobium	•	Scandium	<0.00
Lead	0.017	Samarium	₹0.002	Zirconium	0.005	Calcium	24
Thallium		Neodymium	<0.002	Yttrium		Potassium	26
Mercury	_a	Praseodymium		Strontium	0.19	Chlorine	650
Gold		Cerium		Rubidium	0.027	Sulphur	105
Platinum		Lanthanum	<0.002	Bromine	2.9	Phosphorus	6.8
Iridium		Barium	0.19	Selenium		Silicon	120
Osmium		Cesium		Arsenic	0.32	Aluminum	0.71
Rhenium	_b	Iodine	23	Germanium	0.010	Magnesium	13
Tungsten	<0.005 <0.015	Tellurium		Gallium	0.001	Sodium	120
Tantalum		Antimony	0.048	Zinc	0.77	Fluorine	87
Hafnium	<0.006	Tin	0.011 _b	Copper	0.11	Oxygen	- - -
Lutecium		Indium		Nickel	0.014	Nitrogen	-
Ytterbium	<0.004	Cadmium	0.003	Cobalt	0.012	Carbon	-
Thullium		Silver		Iron	0.72	Boron	
Erbium	<0.003	Palladium		Manganese	0.008	Beryllium	<0.00
Holmium		Rhodium		Chromium	0.090	Lithium	0.00
Dysprosium							
		· · · · · · · · · · · · · · · · · · ·	Plant L	, g/m <sup>3</sup>			
Jranium	•	Terbium		Ruthenium		Vanadium	0.23
Thorium		Gadolinium		Molybdenum	0.015	Titanium	0.24
Sismuth	<0.002	Europium		Niobium	0.005	Scandium	****
Lead	0.14	Samarium	<0.007	Zirconium	0.026	Calcium	110
Challium		Neodymium		Yttrium	0.001	Potassium	5.3
dercury	_a	Praseodymium		Strontium	0.99	Chlorine	2.1
Gold		Cerium	0.011	Rubidium	0.20	Sulphur	330
Platinum		Lanthanum	0.005	Bromine	0.51	Phosphorus	10
Cridium		Barium	0.37	Selenium	0.002	Silicon	15
Osmium		Cesium	0.007	Arsenic	0.031	Aluminum	0.44
Rhenium	_b	Iodine,	0.005	Germanium		Magnesium	4.6
<b>Tungsten</b>	<0.003	Tellurium		Gallium ·		Sodium	78
Cantalum	0.003°	Antimony	0.30	Zinc	2.4	Fluorine	2.5 <sub>a</sub>
<b>lafnium</b>	<0.004	Tin	0.046	Copper	0.54	Oxygen	- <u>°</u>
Lutecium		Indium	_D	Nickel	0.25	Nitrogne	- <u>°</u>
/tterbium	<0.003	Cadmium	0.003	Cobalt	0.033	Carbon	_a _a _a _a
Chullium		Silver	0.001	Iron	4.5	Boron	-°
Erbium	0.001	Palladium		Manganese	0.27	Beryllium	
Rolmium		Rhodium		Chromium	0.26	Lithium	2.1
Dysprosium							
	···		Plant N	, g/m <sup>3</sup>			
Jranium	<0.004	Terbium		Ruthenium		Vanadium	0.033
horium	<0.004	Gadolinium	<0.003	Molybdenum	0.030	Titanium	0.089
ismuth	<0.008	Europium	<0.003	Niobium	0.001	Scandium	<0.004
ead	0.95	Samarium	<0.005	Zirconium	0.054	Calcium	570
hallıum	<0.002	Neodymium	<0.004	Yttrium	0.017	Potassium	58
lercury	<u>-</u> ā	Praseodymium	0.002	Strontium	2.1	Chlorine	1.1
old		Cerium	0.002	Rubidium	0.51		L,400
latinum		Lanthanum	0.008	Bromine	0.19	Phosphorus	110
ridium		Barıum	1.3	Selenium	0.063	Silicon	54
smium		Cesium	0.001	Arsenic	0.40	Aluminum	110
henium	_b	Iodine	0.12	Germanium	0.005	Magnesium	42
ungsten	<0.010	Tellurium	V. 42	Gallium	0.002	Sodium	150
antalum	<0.011	Antimony	0.12	Zinc	580	Fluorine	4.7
lafnium	<0.012	Tin	0.008	Copper	0.11	Oxygen	
utecium	<0.002	Indium	0.00g	Nickel	0.39	Nitrogen	
tterblum	<0.002	Cadmium	0.004	Cobalt	0.46	Carbon	_
hullium	<0.003	Silver	0.002	Iron	80	Boron	
rpram	<0.003	Palladium	0.002	Manganese	27	Beryllium	<0.00
olmium	.0.003	Rhodium		Chromium	44	Lithium	0.03
ysprosium	<0.002	MICALUM		CHIOMITUM	11	DA CHAUM	0.03

<sup>&</sup>lt;sup>8</sup>Not reported. <sup>b</sup>Internal standard. <sup>c</sup>Instrument source.

TABLE 22 (continued)

			Plant S	3, g/m <sup>3</sup>			
Jranium	<0.002	Terbium	<0.002	Ruthenium		Vanadium	0.0
Thorium	<0.002	Gadolinium	_	Molybdenum	0.021	Titanium	0.0
3ismuth	0.012	Europlum		Niobium	0.002	Scandium	<0.0
Lead	0.085	Samarium		Zirconium	0.016	Calcium	11
Thallium	_a	Neodymium	≤0.004	Yttrium		Potassium	73
dercury	_"	Praseodymium	₹0.005	Strontium	0.027	Chlorine	2.5
Gold		Cerium	₹0.005	Rubidium	0.81	Sulphur	24
Platinum		Lanthanum	₹0.007	Bromine	13	Phosphorus	15
[ridium		Barium	0.097	Selenium	0.001	Silicon	18
Osmium	b	Cesium	0.002	Arsenic	0.064	Aluminum	18
Rhenium	-	Iodine	0.017	Germanium		Magnesium	2.8
<b>Tungsten</b>	<0.005	Tellurium		Gallium		Sodium	95
lantalum	<0.012	Antimony	0.84	Zinc	0.29	Fluorine	1.1
lafnium	<0.004	Tin	0.024 _b	Copper	0.28	Oxygen	
Lutecium		Indium		Nickel	0.005	Nitrogen	
<b>(tterbium</b>	<0.003	Cadmium	0.008	Cobalt		Carbon	
Chullium	<0.002	Silver	0.001	Iron	1.0	Boron	
Erbium	<b>≧0.003</b>	Palladium		Manganese	0.10	Beryllium	<0.0
<b>Holmium</b>	_	Rhodium		Chromium	0.016	Lithium	0.0
ysprosium							
,	<del></del> -		Plant T	, g/m <sup>3</sup>			
Jranium	0.002	Terbium		Ruthenium		Vanadium	0.0
horium	4.444	Gadolinium		Molybdenum	0.005	Titanium	0.0
ismuth		Europium		Niobium	*****	Scandium	0.0
ead	0.042	Samarium		Zirconium	0.006	Calcium	5.2
hallium		Neodymium		Yttrium	0.002	Potassium	36
ercury	_a	Praseodymium	0.002	Strontium	0.030	Chlorine	0.5
old		Cerium	0.002	Rubidium	0.13	Sulphur	0.7
latinum		Lanthanum	0.006	Bromine	0.13	Phosphorus	0.7
ridium		Barium	0.022	Selenium	0.005	Silicon	1.9
Smium		Cesium	0.001	Arsenic	<0.003	Aluminum	3.3
Chenium	_b	Iodine	0.002	Germanium	.0.005	Magnesium	1 4
ungsten	<0.002	Tellurium	0.002	Gallium		Sodium	40
antalum		Antimony	0.009	Zinc	0.29	Fluorine	0.9
afnium	<0.002	Tin	0.005	Copper	0.040	Oxygen	0.7
utecium		Indium	2.072	Nickel	0.045	Nitrogen	
tterblum	<0.002	Cadmium	0.001	Cobalt	0.002	Carbon	
hullium		Silver	0.002	Iron	0.57	Boron	
rbium	<0.002	Palladium	0.002	Manganese	0.059	Beryllium	
olmium	<0.002	Rhodium		Chromium	0.058	Lithium	0.0
ysprosium	20.002	111002011		,	0.030	2101110111	0.0
			Plant U	, q/m <sup>3</sup>			
		m 1 !		<u> </u>	<del></del>	***	
tanium		Terbium		Ruthenium	0 007	Vanadium	0.0
horium	40.000	Gadolinium		Molybdenum	0.007	Titanium	0.0
ismuth	<0.002	Europium		Niobium	0.000	Scandium	0.0
ead	0.006	Samarıum		Zirconium	0.002	Calcium	180
hallium	_a	Neodymium	.0 000	Yttrium	0.001	Potassium	37
ercury	-	Praseodymium	<0.026	Strontium	0.32	Chlorine	170
old		Cerium	<0.001	Rubidium	0.043	Sulphur	16
latinum		Lanthanum	≤0.081	Bromine	0.55	Phosphorus	6.4
rıdium		Barium	0.16	Selenium	0.018	Silicon	14
smium	_b	Cesium	0.002	Arsenic	0.14	Aluminum	0.2
henium		Iodine	0.076	Germanium		Magnesium	11
ungsten	<0.002	Tellurium	0.10	Gallium		Sodium	83
antalum	40.000	Antimony	0.19	Zinc	16	Fluorine	2.4
afnıum	<0.002	Tin	0.003	Copper	0.099	Oxygen	
utecium		Indium	_b	Nickel	0.058	Nitrogen	
tterblum		Cadmium	0.003	Cobalt	0.10	Carbon	
hullium		Silver		Iron	0.12	Boron	
rbium		Palladium		Manganese	0.53	Beryllıum	_
		Rhodium		Chromium	0.005	Lithium	0.0
olmıum ysprosium							

TABLE 22 (continued)

			Plant V	, g/m <sup>3</sup>			
Uranium		Terbium		Ruthenium		Vanadium	0.012
Thorium d		Gadolinium		Molybdenum	0.005	Titanium	0.12
Bismuth	0.003	Europium		Niobium		Scandium	• • • • •
Lead	0.089	Samarium		Zirconium	0.002	Calcium	43
Thallium		Neodymium	<0.002	Yttrium	0.001	Potassium	2.0
Mercury	_a	Praseodymium	₹0.002	Strontium	0.19	Chlorine	1.0
Gold		Cerium	<b>~0.005</b>	Rubidium	0.022	Sulphur	45
Platinum		Lanthanum	0.003	Bromine	1.1	Phosphorus	4.2
Iridium		Barium	0.36	Selenium		Silicon	9.5
Osmium	_b	Cesium		Arsenic	0.012	Aluminum	4.0
Rhenium	-	Iodine	≤0.003	Germanium		Magnesium	7.4
Tungsten	<0.002	Tellurium		Gallium		Sodium	42
Tantalum		Antimony	0.010	Zinc	3.1	Fluorine	1.4
Hafnium	<0.002	Tin	0.002	Copper	2.0	Oxygen	-
Lutecium		Indium	_	Nickel	0.010	Nitrogen	-
Ytterbium		Cadmium	0.011	Cobalt	0.073	Carbon	-
Thullium	<0.001	Silver		Iron	4.7	Boron	_
Erbium	₹0.002	Palladium		Manganese	0.31	Beryllium	
Holmium		Rhodium		Chromium	0.066	Lithium	0.041
Dysprosium						•	
			Plant W	, g/m <sup>3</sup>			
Uranium	<0.003	Terbium		Ruthenium		Vanadium	0.011
Thorium	<0.003	Gadolinium	<0.002	Molybdenum	<0.007	Titanium	0.079
Bismuth	₹0.005	Europium	<0.004	Niobium		Scandium	<0.002
Lead	0.17	Samarium	₹0.003	Zirconium	0.023	Calcium	94
Thallium	<0.002	Neodymium	<0.004	Yttrium	0.006	Potassium	660
Mercury	a	Praseodymium	₹0.004	Strontium	0.31	Chlorine	1.2
Gold		Cerium	₹0.010	Rubidium	2.0	Sulphur	16
Platinum		Lanthanum	₹0.006	Bromine	0.55	Phosphorus	0.12
Iridium		Barium		Selenium	0.018	Silicon	30
Osmium	•	Cesium	0.003	Arsenic	0.033	Aluminum	7.3
Rhenium	_b	Iodine	<0.013	Germanium	<0.004	Magnesium	12
Tungsten	<0.007	Tellurium		Gallium	0.004	Sodium	8.3
Tantalum	<0.003 <sup>C</sup>	Antimony	<0.013	Zinc	0.060	Fluorine	11
Hafnium	<0.009	Tin	0.011	Copper	0.072	Oxygen	a
Lutecium	₹0.001	Indium	_6	Nickel	0.024	Nitrogen	_a _a
Ytterbium	<0.006	Cadmium	0.006	Cobalt	0.048	Carbon	_a
Thullium	<0.002	Silver	0.003	Iron	0.67	Boron	_a
Erbium	<0.003	Palladium		Manganese	0.22	Beryllium	<0.003
Holmium		Rhodium		Chromium	0.028	Lithium	0.054
Dysprosium	<0.001				0.022		••••
		<del></del>	Plant X	(, a/m³			
Uranium		Terbium		Ruthenium		Vanadium	0.026
Thorium		Gadolinium		Molybdenum	0.005	Titanium	0.018
Bismuth		Europium		Niobium		Scandium	<0.002
Lead	0.030	Samarium	•	Zirconium		Calcium	23
Thallium	· _a	Neodymium		Yttrium	0.007	Potassium	9.0
Mercury	•	Praseodymium		Strontium	0.14	Chlorine	130
Gold		Cerium		Rubidium	0.023	Sulphur	29
Platinum		Lanthanum	0.001	Bromine	2.5	Phosphorus	29
Iridium		Barium	0.13	Selenium		Silicon	18
Osmium	_b	Cesium	0.000	Arsenic	0.022	Aluminum	8.2
Rhenium		Iodine	0.037	Germanium	0.005	Magnesium	4.6
Tungsten	<0.002	Tellurium Antimony	1.9	Gallium	0.001	Sodium	33
Tantalum	40.000			Zinc	0.14	Fluorine	11 _a
Hafnium	<0.002	Tin	0.004	Copper	0.87	Oxygen	- a
Lutecium	.0.000	Indium	_0	Nickel	0.006	Nitrogen	_a _a _a
Ytterbium	<0.002	Cadmium		Cobalt	0.12	Carbon	
Thullium		Silver		Iron	2.5	Boron	-u
Erbium		Palladium		Manganese	0.028	Beryllium	
Holmium		Rhodium		Chromium	0.027	Lithium	0.052
Dysprosium							

a<sub>Not</sub> reported. b<sub>Internal</sub> standard. c<sub>Instrument</sub> source.

TABLE 23. CONCENTRATION OF METHYLENE CHLORIDE EXTRACTABLE ORGANICS IN FILTERED SECONDARY EFFLUENTS

Plant_	Organic concentration, g/m <sup>3</sup>	Plant	Organic concentration g/m <sup>3</sup>
	CO 7		0.04
A	63.7	N	9.24
В	3.18	S	5.40
С	28.2	${f T}$	17.8
E	3.60	ប	14.6
F	16.0	V	a
G	. 27.2	W	15.0
K	2.73	X	13.5
L	18.3		T.

aAnalysis not performed.

The methylene chloride extract then went through a solvent exchange step to transfer the sample to a nonpolar solvent. This extract was then passed through a liquid chromatography column that divided the sample into eight fractions. Each of the eight fractions was analyzed by an infrared (IR) spectrophotometer and then by a low resolution mass spectrometer (LRMS). The classes of organic compounds and their relative intensities found in each fraction are presented in Tables 24 and 25.

IR analyses indicated the presence of aliphatic hydrocarbons, C=O esters and acids, aromatics, phthalate esters, and fatty acid groups. LRMS analyses identified the following major classes of compounds: paraffinic/olefinic, alkyl benzenes, alcoholic ethers, di-n-octyl phthalate, bis(hydroxy-t-butyl phenol) propane, tri-t-butyl benzene, alkyl phenols, dichloroaniline, toluene-sulfonyl groups, vinyl stearate, and azo compounds.

TABLE 24. LEVEL 1 INFRARED ANALYSIS OF THE ORGANIC EXTRACTS

	Plant A								
Before extraction	Bonded OH, aliphatic CH, C=O ester and acid, ketone or aldehyde, conjugated C=C, possible aromatic C=C, ether groups $(CH_2)_n$ , where $n=24$								
1	All aliphatic hydrocarbons								
2	Aliphatic and aromatic hydrocarbons, aromatic C=C								
3	Aliphatic C-H, ester or aldehyde C=O, conjugated C=C, various CH2 groups, or aromatic substitution bonds								
4	Similar to fraction 3								
5	Similar to fraction 3								
6	Bonded OH; aliphatic C-H; acid, ketone, or aldehyde C=O; conjugated and aromatic C=C; possible phthalate ester; ether group or S								
7	Bonded OH, aliphatic CH, ester or aldehyde C-O, water in material, ether or Si-O groups								
8	Bonded OH and 1,630 cm <sup>-1</sup> absorption-water, aliphatic CH trace of C=0, SiO <sub>2</sub> , poorly defined organic								
	Plant B								
Before extraction	Bonded OH, aliphatic CH, ether, (CH <sub>2</sub> ) <sub>4</sub> , C=O, or C=C?								
ı	Aliphatic CN, trace aromatic CH, Si-CH <sub>3</sub> (?), methylene CH <sub>2</sub> groups >4								
2	Very strong background adsorptiononly aliphatic CH visible								
3	Bonded OH, aliphatic CH, ester C=O, acid, aldehyde or ketone C=O (CH2) or >4.								
4	Bonded OH, trace aromatic CH, aliphatic CH, ester C=O acid, aldehyde or ketone C=O, various CH <sub>2</sub> groups.								
5	Similar to 4, but stronger bonded OH, less ester C=O, more acid, aldehyde or ketone C=O, various CH2 groups complex spectrum.								
6	Bonded OH, aliphatic CH, some ester C=O, acid, aldehyde or ketone C=O, ether groups, may contain glycol ether type compounds.								
7	Bonded OH, aliphatic CH, ester C=O, acid, aldehyde or ketone C=O, strong ether group, glycol ether type of compound.								
. 8	Strong bonded OH, weak C-H (aliphatic) trace C=O, nonconjugated C=C, SiO2 present.								
	Plant E								
Before extraction	Aliphatic CH, diffuse C=O region, carboxylate ion, ether group complex spectrum								
1	Aliphatic hydrocarbons, no indication of number of CH <sub>2</sub> groups.								
2	Poor spectrumaliphatic CH, CO2, and water vapor in spectrum.								
3	Poorly defined spectrumaliphatic CH, numerous ill-defined bonds. No C=O or C=C.								
4	Aliphatic CH, ester C=O, strong background adsorption.								
5	Aliphatic CH, ester=0, most likely aliphatic ester, possibly acetatemay be single compound.								
6	Spectrum too strongbonded OH, aliphatic CH, ester C=O, acid, ketone or aldehyde C=O, large portion of compound in No. 5 plus additional carbonyl compounds.								
6 (repeat)	Mixture of several comps. Ester C=O, and acid, ketone or aldehyde C=O. Ether group.								
7	Bonded OH, aliphatic CH, acid, ketone or aldehyde C=O, spectrum too strong for good identification.								
7 (repeat)	Bonded CH, aliphatic CH, acid, aldehyde or ketone C=O, evidence of both acid and acid salt (carboxylate ion) CH(CH <sub>3</sub> ) <sub>2</sub> group possible, no long chain (CH <sub>2</sub> ) <sub>n</sub> groups.								
8	Bonded OHevidence of water (3,350 cm <sup>-1</sup> and 1,635 cm <sup>-1</sup> ) SiO <sub>2</sub> present aliphatic CH.								
	(continue								

# TABLE 24 (continued)

Fraction No.	· Interpretation						
<del></del>	Plant F						
Before extraction	Bonded OH, allphatic CH, diffuse C=O, and C=C regions Si-O possible, trace CH <sub>2</sub> Cl <sub>2</sub> .						
1	Long chain aliphatic hydrocarbons. Unknown 1,265 cm <sup>-1</sup> bond. Si-(CH <sub>3</sub> )-?						
2	Aliphatic CH, ester C=O, conjugated C=C, trace 1,265 cm <sup>-1</sup> .						
3	Weak bonded OH, aliphatic CH, aromatic CH-?, ester C=O, conjugated C=C, progression of CH substitution bonds, series of CH₂ bond.						
4	Similar to No. 3, series of ill-defined bonds below 1,400 cm <sup>-1</sup> .						
5	Bonded OH, trace aromatic CH, aliphatic CH, ester or aldehyde C=O, conjugated C=C, series of CH2 groups aromatic substitution bonds						
6	Bonded OH, allphatic CH, ester C=O, nonconjugated C=C or amides, ether or SiO groups, secondary amide possible, CH2 groups n >4.						
7	Similar to No. 6.						
8	Strong bonded OH, aliphatic CH, weak ester or aldehyde C=O, H2O present (1,640 cm <sup>-1</sup> ) SiO <sub>2</sub> present.						
	Plant G						
Before extraction	Aromatic and aliphatic CH, residual CH <sub>2</sub> Cl <sub>2</sub> in spectrum.						
1	Aliphatic hydrocarbonchain length >C4, possible C(CH3)3 group.						
2	Aliphatic CH, ester C=O, phthalate bonds, various chain lengths of CH2.						
3	Aliphatic CH, ester C=O, some C=C, various CH <sub>2</sub> groupings.  Bonded O-H, aliphatic CH, ester C=O, some acid, aldehyde or ketone C=O, C=C, possible fatty acid group, various CH <sub>2</sub> groups.						
4							
5	Identical to No. 4.						
6	Considerable bonded OH, aliphatic CH, ester C=O conjugated C=C, Si=O or ether group.						
7	Similar to No. 6.						
8	Considerable O-H, aliphatic C-H, ester C=O, SiO or ether groups.						
	Plant K						
Before extraction	Bonded OH, aliphatic CH, diffuse C=O region Si(CH3) group? Diffuse spectrum.						
1	Appears to contain water, aliphatic C-H, Si-CH3, mainly hydrocarbon compounds.						
2 Aliphatic hydrocarbons, silicones.							
3 Mainly silicone type materials.							
4. Bonded CH, some aromatic CN, aliphatic CH, ester C=O, acid, ketone or aldehyde C=O, carboxylate ion, conjugated C=C, material, (CH <sub>2</sub> ) <sub>n</sub> where n ≥4.							
5	Strong background adsorption, aliphatic CH, ester C=O various (CH <sub>2</sub> ) groups, some silicone adsorption.						
6	Bonded OH, aliphatic CH, trace aromatic CH, ester C=O, aromatic C=C, silicone adsorption.						
7	Similar to No. 6, not as strong a spectrum.						
8	Strong OH adsorption, very weak C-H, trace of C=O (may be H2D background) SiO2 adsorption, low organic content.						
	(continued)						

# TABLE 24 (continued)

Fraction No.	Interpretation								
	Plant L								
Before extraction	Bonded OH, aliphatic CH, C=0, aromatic or conjugated C=C, ether group. Complex spectrum CEN, SiO2-?								
1	Aliphatic CH, trace C=0, C=C, CH2 group >4, possible CH(CH3)2 groupmainly aliphatic hydrocarbons.								
2	Aliphatic hydrocarbonsbranched chain, trace of C=O, C=C no long (CR <sub>2</sub> ) <sub>n</sub> groups.								
3	Bonded OH, aliphatic CH, ester C=O, conjugated C=C strong ether group, possible glycol ether.								
4	Bonded OH, aliphatic CH, ester C=O, nitrite (CEN) group, strong ether group, various streight and branched CH2 groups.								
5	Bonded OH, some aromatic CH, aliphatic CH, CEN nitrite group, ester C=O, conjugated or aromatic C=C, ether group, very complex mixture spectrums.								
	Similar to No. 5 but less CEN, more bonded CH, ester C=O, conjugated or aromatic C=C, some aromatic C-H, aliphatic CH ether groupingcomplex spectra.								
7	Some bonded OH, aliphatic CH, trace CEN, ester C=O, carboxylate ion, possible C-Cl group.								
8	Contain water and SiO2, plus some of materials found in No. 7. Carboxylate ion.								
	Plant E								
Before extraction	Bonded OH, aliphatic CH, diffuse C=O, C=C region, silicone (CH <sub>2</sub> ) <sub>4</sub> ether?								
1	Long chain aliphatic hydrocarbon.								
2	Aliphatic CH, ester C=O, possible (CH(CH <sub>3</sub> ) <sub>2</sub> group, spectrum not very distinct.								
3	Weak bonded OH, aliphatic CH, ester C=O, conjugated C=C, CH(CH <sub>3</sub> ) <sub>2</sub> group, various CH <sub>2</sub> groups.								
4	Weak bonded OH, aliphatic CH, ester C=O, possible fatty acid groups.								
5	Bonded OH, aliphatic CR, medium ester C=O,-nonconjugated C=C, spectrum not very distinct.								
6	Bonded CH, aliphatic CH, ester C=O, acid, aldehyde or ketone C=O, conjugated C=C, aliphatic ketone or ester group, possible ether group, various CH <sub>2</sub> chain lengths.								
7	Bonded OH, aliphatic CH, ester C=O, conjugated C=C, possible fatty acid gropus, ether group.								
8	Strong OH, weak aliphatic CH, ketone, acid or aldehyde C=O, ester C=O, strong C=C, ether group.								
	, Plant S								
Before extraction	Bonded OH, aliphatic CH, ester C=O, (CH <sub>2</sub> ) groups, broad diffuse spectrum, ether groups possible.								
1	Aliphatic hydrocarbon, chain length <cu 1,265="" at="" bond="" cm<sup="" unidentified="">-1.</cu>								
2	Aliphatic hydrocarbons, trace of ester C=O, trace of C=C, not well defined below 1,300 cm <sup>-1</sup> :								
, 3	Aliphatic CH, trace of aromatic CH, very strong ester C=O or aldehyde C=O. Ester group may be acetate; if aldehyde, long chain aldehyde.								
4	Strong background absorption 4,000 cm <sup>-1</sup> to 900 cm <sup>-1</sup> . Aliphatic CH, ester or aldehyde C=O. Poor spectrum for interpretation.								
5	Nearly identical to fraction No. 4.								
6	Very complex spectra. Bonded OH, aliphatic CH, ester and acid, ketone or aldehyde C∞O, ether or SiO groups. Several types of CH groupings.								
7	Weak and diffuse spectra. Aliphatic CH, ester C=O serve C=C, ether or SiO group.								
8	Strong OH group, weak alighatic CH, weak acid; aldehyde or ketone C=O, strong conjugated C=C contains some SiO2, diffuse spectrum								
	•								

# TABLE 24 (continued)

Fraction No.	Interpretation						
	Plant T						
Before extraction	Bonded OH, aliphatic CH, C=O, C=C, ether of SiO groups, (CH <sub>2</sub> ) groups. Very diffuse spectrum.						
1	Bonded OH, aliphatic CH, ester C=O, some C=C, CH <sub>2</sub> groups >C <sub>4</sub> .						
2	Bonded OH, allphatic CH, medium ester C=O, possible fatty acid groups, no definite CH2 groupings.						
3	Borded OH, allphatic CH, strong ester C=P, nonconjugated C=C, ether or SiO groups. Various chain lengths of (CH2) n.						
4	Bonded OH, medium aliphatic CH, medium ester C=O, nonconjugated C=C, ether or SiO group, various (CH <sub>2</sub> ) groups, possible acid salts.						
5	Similar to No. 3.						
6	Bonded OH, aliphatic CH, medium ester C=O, series of 5 unknown bonds medium intensity 1,510 cm <sup>-1</sup> to 1,610 cm <sup>-1</sup> , ether or SiO grovarious (CH <sub>2</sub> ) groups.						
7	Similar to No. 6, but weaker OH, C=CO, conjugated C=C, fatty acid groups? ether or SiO group. (CH2) groups.						
8	Very strong bonded OH group, medium aliphatic CH, weak ester C=O, strong nonconjugated C=C, possible fatty acid group, ether or SiO group, no CH2 >C4.						
	Plant U						
Before extraction	Trace bonded CH, aliphatic CH, acid, ketone or aldehyde C=O, silicones, some (CH <sub>2</sub> ) groups.						
1	Aliphatic hydrocarbon, conjugated or aromatic C=C, weak, (CH <sub>2</sub> ),no >4.						
2	Aliphatic hydrocarbon, conjugated C=C (alkene?), diffuse CH <sub>2</sub> groups.						
3	Aliphatic C-H, ester C=O, conjugated or aromatic C=C, possible unsaturated ester-fumarate, maleate, etc.						
4	Bonded OH, trace sec N-H or oxcetone C=O, aliphatic C-H possible trace C=N, ester C=O, conjugated C=C, unsaturated ester group.  Complex spectra below 1,500 cm <sup>-1</sup> .						
5	Aliphatic C-H, ester C=O, diffuse spectra below 1,100 cm <sup>-1</sup> .						
6	Complex spectra, bonded OH, NH or C=O overtone, aliphatic C-H, ester C=O, acid, aldehyde or ketone C=O (weak), ether group, complex bond pattern below 1,500 cm <sup>-1</sup> .						
7	Bonded OH, aliphatic C-H, ester C=O, conjugated C=C, possible ether group, miscellaneous (CH2) groups.						
8	Bonded OH strong, aliphatic CH, strong C=C, ether group, some SiO2, possible glycol ethers.						
	Plant V						
Sefore extraction	Trace bonded OH, aliphatic CH, ester C=O, acid, aldehyde or ketone C=O, silicone adsorption, some (CH2) groups.						
1	Aliphatic CH, (CH <sub>2</sub> ) where n >4, hydrocarbons plus possible Si-CH <sub>3</sub> .						
2	Aliphatic and aromatic CH, some ester C=O, silicones.						
3	Poor spectrumlow organic content? aliphatic CH strong background adsorption.						
4	Bonded OH, aromatic and aliphatic CH, ester C=O, conjugated C=C, silicones.						
5	Identical to No. 4.						
6	Aliphatic CH, ester C=0, silicones, CH <sub>2</sub> various groups ester stronger, silicones weaker than in No. 4 or No. 5.						
7	Bonded OH, aliphatic CH, ester C=O, conjugated C=C ether grouppossible glycol ether-type compounds.						
8	Strong bonded OH, weak aliphatic CH, ester C=O, strong C=C, some ether under SiO <sub>2</sub> adsorption.						

Fraction No.	Interpretation						
	Plant W						
Before extraction	Bonded OH, aliphatic CH, ester C=O, acid, ketone or aldehyde C=O. (CH <sub>2</sub> ) -n where n >4, diffuse spectrum 1,300 cm-1 to 900 cm						
1	Bonded OH, aliphatic CH, ester C=O, possible ether group (CH <sub>2</sub> ) where n >4. Not typical fraction 1.						
2	Aliphatic CH groups, ester C=0, weak spectrum.						
3	Aliphatic ester compounds, ester C=O, no aromatic CH.						
4	Trace OH, aliphatic CH, ester C=P, weak diffuse spectrum.						
5	Weak diffuse spectrum, poor background, OH (water?), aliphatic CH, weak ester C=O, nonconjugated C=C.						
6	Bonded OH, aliphatic CH, ester C=O, long chain CH <sub>2</sub> groups, unsaturated acid, aldehyde or ketone, conjugated C=C, possible amic groups.						
7	Poor spectrum. Bonded OH, aliphatic CH, numerous C=O types, C=C, amide possible, very diffuse below 1,400 cm-1.						
8	Weak spectrum, bonded OHlikely H2O, very weak aliphatic CH, low organic content.						
	Plant X						
Before extraction	Bonded OH, aliphatic CH, C=O, ether or SiO, (CH <sub>2</sub> ), numerous broad diffuse bonds.						
1	Aliphatic hydrocarbons, possible Si(CH <sub>3</sub> ).						
2	Aliphatic CH, ester C=O, conjugated C=C, hydrocarbon.						
3	Bonded OH, aliphatic ch, ester C=O, phthalate plus other types of ester materials.						
4	Similar to fraction No. 3.						
5	5 Bonded OH, trace aromatic CH, aliphatic CH, ester C=O, acid, aldehyde or ketone C=O, conjugated C=C, various straight and branched CH <sub>2</sub> chains.						
6	Bonded OH, aliphatic CH, ester C=O, conjugated and aromatic C=C, ether group, long CH2 chains.						
7	Very similar to No. 6.						
8	Strong OH adsorption, similar to No. 6 and No. 7 but more OH and presence of SiO2.						

TABLE 25. LEVEL 1 LOW RESOLUTION MASS SPECTROMETER ANALYSIS OF ORGANIC FRACTIONS

Plant and	Fraction	Cat	egories present		Subcategories present	
organic fraction	weight,	Relative intensity	Category	Relative intensity	Specific compounds	Other unknown compounds present
1	_b	_b	ь	_b	ь	_b
2	16.1	100 100	Aliphatics <sup>C</sup> Aromatics	100 100	Peraffinic/olefinic (or cyclic-paraffinic, etc.). d.e Alkyl benzenes (91, 105, 114, 133 ions present.	No masses above 498.
3	5.3	100 100	Aliphetics Aromatics	100 100	Paraffinio/olefinio (or cyclic-paraffizio, etc.) d.e Alkyl benzenes (91, 105, 114, 133 ions present).d.e	10:368(100), 369(45), 353(20) 1:345(100), 396(35), 411(30)
4	_b	_b	_b	_6	_"	_ь
5	_b	_6	_b	_6	<u>,</u>	_b
6	10.2	100 10 10	Aliphatics Alcohols/ethers Phenols	100 10 10	Paraffinic/olefinic (or cyclic-paraffinic, etc.). d,e Alcoholic ethers (45 ions to 89 ions). d,e Bis(hydroxy-t-butyl phenyl) propane (C29H2CO2) 386 340.	No masses above 414.
		10	Esters	10	Di-n-octyl phthalata (C2+N2gO4) MH390.	
7	_b	_ <b>p</b>	_p	_6	_ <b>6</b>	_b
6	21.0	10 100	Aliphatics Alcohola/ethers	10 100	Paraffinic/olefinic (or cyclic-paraffinic, etc.). $^{d}_{*}e$ Alcoholic-ethers (45 ions to 89 ions). $^{d}_{*}e$ Ethers: Di-n-octyl pathalate ( $C_{24}H_{36}O_{4}$ ) MW390.	No masses above 368. 100:254(100), 126(20), 127(15)
					Plant B	
1	_b	_b	_b	_b	_b	_b
2	10.3	100	Aliphatics	100	Summas Plant A, Fraction 2.	No masses above 354,
	b	_p 100	Esters b	100	Same as esters, Plant A, Praction 6.	_b
3	_	_	-	-p	•	_
4	3.8	10	Aliphatic	10	Same as Plant A, Fraction 2.	No masses above 483. 10:279(100), 294(28), 280(12) 100:341(100), 356(36), 342(27) 100:381(100), 356(36), 392(11) 100:410(100), 151(37), 411(31) 100:429(100), 444(23), 445(15)
5	_b	_b	_ь	-p	_b	_b
6	13.9	1	Aliphatics	1	Same as Plant A, Fraction 2.	No masses above 437.
7	_b	_p _p	Phenols _b	100 _b	Bis (hydroxy-t-butyl phenol) propane (C23H32O2) 188390. b	_b
8	15.0	10	Aliphatics	10	Same as Plant & Praction 2.	No masses above 414.
		10 10	Alcohols/ethers Esters	10 10	Alcoholiseethers (48 idea to 89 ions). d.e Same as esters, Plant A, Praction 6.	100:294(100), 127(20), 128(12)
					Plant F	
1	9.4	100	Aliphatics	100	Primarily paraffinic.d.e	No masses above 446.
2	0.3	100 100	Aliphatics Esters	100 100	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	No masses above 354.
3	1.9	1 ¶00 10	Aliphatics Aromatics Esters	1 100 10	Aliphatics. d.e  Alkyl benzenes: Tri-t-butyl benzene (C18H30) MM246.  Same as esters, Plant A, Fraction 6.	No masses above 354.
4	3.3	100 10 10	Aliphatics Aromatics Esters	100 10 10	Same as Plant A, Fraction 2. Tri-t-butyl bensens (CleMag) MM246. Same as esters, Plant A, Fraction 6.	No masses above 381.
5	1.4	10	Aliphatics Aromatics	10 10 1	Same as Plant A, Fraction 2,	No masses above 340. 10:239(100), 240(20), 254(25)
		10/100	Phenols	10 100	Alkyl bensenes: Tri-t-butyl bensene (C <sub>18</sub> H <sub>30</sub> ) M6246. Alkyl phenols (135, 107, 121, 148 ions).0,e Ais(hydroxy-t-butyl phenyl) propene (C <sub>2</sub> H <sub>32</sub> O <sub>2</sub> ) M6390.	
			Esters	100	Same as esters, Plant A, Praction 6.	
6	18.6	10/100	Aromatics Phenols	1 10 100	Tri-t-butyl bensene (CleHso) NW246. Alkyl phenols (135, 107, 121, 149 ions).d,e Bis(hydroxy-t-butyl phenyl) propene (C23H32O2) NW 390.	No masses above 354, 100:45(100), 42(90) (2 compound
			Esters	10	Same as esters, Plant A, Fraction 6.	100:59(100) (4 compounds)
7	51.7	100	Aliphatic Phenols	100 10	Primarily paraffinio. d,e	No masses above 354.
		100	Esters	100	Bis(hydroxy-t-butyl phenyl) propane ( $C_{23}H_{32}O_2$ ) MW390. Same as esters, Plant A, Praction 6.	
8	42.7	10	Aliphatics	10	Same as Plant A, Fraction 2.	No masses above 354.
		100	Esters	100	Same as esters, Plant A, Fraction 6.	
	42.7		Esters			No masses above

Relative intensity of the mass-to-charge ratio (intensity relative to dominant ion).

f Molecular weight.

borganic weight of fraction below gravimetric threshold of 0.1 mg; therefore, no analysis was performed.

Generally all ions up through 498 present in aliphatic-type pattern; however, all mass >100 are abnormally strong for typical aliphatics.

d No molecular weight range determination possible.

<sup>&</sup>lt;sup>e</sup>No composition determination possible.

TABLE 25 (continued)

Plant and	Praction		Categories present		Subcategories present	
organic fraction	weight,	Relative intensity	Category	Relative	Specific compounds	Other unknown compounds present
					Plant G	•
1	31.8	100	Aliphatics	100	Same as Plant P, Fraction 2.	No masses above 410.
2	7.2	1 100	Aliphatics Esters	1 100	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	
3	2.3	100 1	Aliphatics Esters	100 1	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	No masses above 378.
4	0.9	100 1	Aliphatics . Esters	100 1	Same as Plant A, Fraction 2. Same as estern, Plant A, Fraction 6.	No masses above 381.
5	0.3	10 1 100	Aliphatics Phonols Esters	10 1 100	Same as Plant A, Praction 2. Same as phenols, Plant A, Praction 6. Same as esters, Plant A, Praction 6.	No masses above 325.
6	29.0	100 100	Aliphatics Alcohols/ethers	100 100	Same as Plant A, Praction 2. Alcoholic ethers with 41, 43, 45 ions and 55, 57, 59 ions. 6,6	No masses above 354. 1:69(100), 41(87), 43(78)
,		1	Phenols Esters	1	Same as phenols, Plant A, Fraction 6. Same as esters, Plant A, Praction 6.	
*	13.9	100 100 1	Aliphatics Alcohols/ethers Phenols Esters	100 100 1	Same as Plant A, Fraction 2. Same as Praction 6. Same as phenols, Plant A, Fraction 6. Same as estars, Plant A, Fraction 6.	No masses above 354, 10:69(100), 41(80), 43(78)
8	15.5	100 10 1	Aliphatics Alcohols/ethers Esters	100 10 1	Same as Plant A, Fraction 2. Same as Fraction 6. Same as esters, Plant A, Fraction 6.	No masses above 354.
						1.
					Plant L	U
1	12.7	100	Aliphatics	100	Seme as Plant P, Fraction 1.	No masses above 367.
2	2,2	100 100	Alighatics Arcmetics	100 100	Sees as Plant F, Fraction 1. Sees as Plant A, Fraction 3.	No masses above 367.
3	8.6	100 100	Phenola Estare	700 700	Alkyl phenols (135, 107, 121, 149 ions). d.e Same as esters, Plant A, Fraction 6.	No masses above 354. 10:69(100), 41(80), 43(78)
4	8.6	100 100 100 100	Alighetics Arometics Phenols Esters	100 100 100 100	Same as Plant F, Fraction 1. Same as Plant F, Fraction 3. Di-t-butyl phenol (O <sub>10</sub> Hg/20) MM/206. Same as estars, Plant A, Fraction 6.	No masses above 429.
5	9.7	100 100 100	Aliphatics Phenole Esters	100 100 100	Same as Plant F, Fraction 1. Alkyl phenol (135, 107, 121, 149 ions) d,e Same as estars, Plant A, Fraction 6.	No masses above 429. 10:69(100), 41(80), 43(78)
6	55.1	100 100	Phonola Esters	100 100	Alkyl phenols (135, 107, 121, 149 ions). d,e Phthalate, probably di-Cs alkyl but with a new series of ion odded (223, 237, 251, 265, 279).	No masses above 340.
7	11.7	100/10	Aliphatics Phemola	100 100 10	Same as Plant A, Fraction 2. Alkyl phanols (135, 107, 121, 144 ions).d.e Ris(Rydroxy-t-butyl phanyl) propane (C23H32O2) MM340.	
		100	Esters	100	Same as Plant A, Fraction 6.	1:200(100), 201(86) 10:280(100), 279(98)
8	19.1	100 100	Aliphatics Esters	100 100	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	No masses above 279. 1:Cesium iodide from infrared plates.

Relative intensity of the mass-to-charge ratio (intensity relative to dominant ion).

No molecular weight range determination possible.

<sup>&</sup>lt;sup>e</sup>No composition determination possible.

TABLE 25 (continued)

Plant	Praction	on <u>Categories present</u>			Subcategories present	
organic fraction	weight,	Relative intensity	Category	Relative intensity	Specific compounds	Other unknown compounds present
=					Plant H	
1	44.5	100 10	Aliphatics Esters	100 10	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	No masses above 428.
2	1.0	100 100 100	Aliphatics Aromatics Enters	100 100 100	Same as Plant A, Fraction 2. Alkyl bensons (91, 105, 119 ions).d.e Same as esters, Plant A, Fraction 6.	No masses above 394.
3	3.9	100	Aliphatics	100	Same as Plant A, Fraction 2.	No masses above 498.
		100 10	Aromatics Esters	100 10	Same as Plant A, Fraction 3. Same as estars, Plant A, Fraction 6	10:368(100), 369(30), 353(20) 10:395(100), 396(30) 1:410(100), 411(30)
4	2.1	100 10	Aliphatics Esters	100 10	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	No masses above 499. 10:368(100), 369(30), 353(20) 10:395(100), 396(30) 10:410(100), 411(30)
5	1.0	100 10 100	Aliphatics Phenols Esters	100 10 100	Same as Plant A, Fraction 2. Same as Plant F, Fraction 5. Same as esters, Plant A, Fraction 6.	No maskes above 279.
6	20.5	100 10 10 10 10/10	Aliphatics Arometics Amines Phenols	100 10 10 10	Same as Plant A, Fraction 2. t-Butyl dichlorobenseme (C10H12C12) MM202 Dichloroanilme (C6H3MC12) MM 161 Same as phenole, Plant F, Fraction 5. Same as phenole, Plant A, Fraction 6. Phthalate, probably di-C6 alkyl but with a new series of ione added: 223, 237, 251, 265, 279 (C2H35O4)	No masses above 490. 1:155
7	8.9	100 10/1	Aliphatics Phenols Esters	100 10 1	MM 390 (probably).  Same as Plant A, Fraction 2.  Same as phenols, Plant F, Fraction 5.  Same as phenols, Plant A, Fraction 6.  Same as esters, Plant A, Fraction 6.	No masses above 340.
8	16.6	100 10 100	Aliphatics Phenols Esters	100 10 100	Same as Plant A, Fraction 2. Same as phemols, Plant F, Fraction 5. Same as esters, Plant A, Fraction 6.	No masses above 279. 1:Cesium iodide.
					Plant 8	
1	12.3	100	Aliphatics	100	Same as Plant A, Fraction 2.	No masses above 446.
2	4.7	100 100	Aliphatics Aromatics	100 100	Same as Plant A, Fraction 2. Same as Plant A, Fraction 2.	No masses above 446.
3	19.7	100 10 1	Alighatics Aromatics Enters	100 10 1	Same as Plant A, Fraction 2. Same as Plant F, Fraction 3. Same as Plant A, Fraction 6.	No masses above 452.
4	11.1	100/10	Aliphatics Aromatics	10 100	Seme as Plant A, Fraction 2. Toluene-sulfonyl-group (91, 155 ions). Best identity is p-toluene sulfonemide.	No masses above 477. 1:381(100), 382(27), 396(17)
		10 10	Phenols Phenols	10 10	Same as Plant P, Fraction 3. Di-t-butyl phenol (C14H22O) MH2O6.	
5	6.6	100 100	Aliphatics Aromatics	100 100	Base as Plant A, Fraction 2. Toluene-sulfomyl group (91, 155 ions). Best identity p-toluene sulfonemide.	No masses above 354. 100:98(100), 97(75)
6	<b>26.8</b> .	1 100 10 10	Aliphatics Aromatics Phenols Esters	1 100 10 10	Same as Plant A, Fraction 2. Toluene-sulfonyl group (91, 155 ions) d.e Same as phenols, Plant A, Fraction 6. Same as esters, Plant A, Fraction 6.	No masses above 446. 10:90(100), 91(68), 106(58)
7	6.5		Aliphatics Aromatics Esters	1 100	Same as Plant A, Fraction 2. Same as aromatics, Plant A, Fraction 4. Same as esters, Plant A, Fraction 6.	No masses above 446. 10:90(100), 91(68), 106(58)
8	13.6	100 10	Aliphatics Esters	100 10 1		No masses above 354.

Relative intensity of the mass-to-charge ratio (intensity relative to dominant ion).

No molecular weight range determination possible.

No composition determination possible.

TABLE 25 (continued)

Plant	Praction	Cate	egories present		Subcat	egories present	_
rganic raction	weight,	Relative intensity	Category	Relative intensity	Plant T	Specific compounds	Other unknown compounds presen
1	18.5	100	Aliphatics Esters	100 1	Same as Plant A, Same as esters,	Praction 2. Plant A, Fraction 6.	No masses above 362.
2	2.6	100 100 10	Aliphatics Aromatics Esters	100 100 10	Same as Plant A, Same as Plant A,		Mo masses above 312. 10:69(100), 41(80), 43(78)
3	6.7	100/100	Aliphatics Aromatics	100 100 10	Same as Plant A, Same as Plant F, Same as Plant F,	Praction 2. Praction 1.	No masses above 486. 1:314(100), 315(25) 10:410(100), 411(32)
4	12.9	10 100	Esters Alighatics	10 100		Plant A, Fraction 6.	No masses above 396.
·		10 10 1	Aromatics Phenols Esters	10 10 · 1	Same as Plant P,	Praction 3. 1 (C <sub>14</sub> H <sub>22</sub> O) NW 206.	10:69(100), 41(80), 43(78)
5	6.9	10 100	Aliphatics Esters	10 100	Same as Plant A, Same as esters,	Praction 2. Plant A, Fraction 6.	No masses above 396.
6	17.4	100 100/100	Aliphatics Phenols	100 100 100		Praction 3. Plant A, Fraction 6.	No messes above 340. 100:99
7	29.2	100 100 100 100	Esters Aliphatics Phenols Esters	100 100 100 100	Same as Plant A, Same as phenols, Phthalate, probal of ions added:	Plant P, Praction 5. oly di-C <sub>6</sub> alkyl but with a new serie 223, 237, 251, 265, 279 ions '	No masses above 279.
8	17.1	10 10	Aliphatics Esters	10 10	(C <sub>2</sub> 4H <sub>38</sub> O <sub>4</sub> ) MM : Same as Plant A, Same as esters, 1		to masses above 336. 100:117(100), 59(44) 10:69(100), 41(80), 43(78) Also many organo-silicon ions; e.g., 207, 221, etc.
					Plant U		
1	59.1	100	Aliphatics	100	Same as Plant P,	Fraction 1.	No masses above 404.
2	8.1	100 100	Aliphatics Aromatics	100 100	Same as Plant A, Same as Plant A,		No masses above 410.
3	11.0	100	Aliphatics	100	Same as Plant A,	Praction 2.	No masses above 398 in aliphati type pattern; all masses abov 100 are abnormally strong for typical aliphatics.
4	4.4	100 100/10	Aliphatics Esters	100 100 10	Vinyl stearate (	Plant A, Praction 6.	No masses above 411. 10:410(100), 411(30)
		10	Amines	10	(C16H12OH2) HH	248.	No magges above 444.
5	5.3	100 100 10	Aliphatics Esters Amines	100 100 10		Praction 2. Plant A, Fraction 6. Ps: chloroaniline (C6R6C1) MM127.	No masses above 444. 10:429(100), 444(20)
6	21.9	100 100	Aliphatics Esters	100 100	stearate (C20H;	Plant A, Fraction 6, and vinyl 3802) ##310.	No masses above 496.
,	11.5	10 100	Amines Aliphatics	10 100	Chloroaniline (Co	• •	No masses above 495.
,	11.3	100 10/100	Phenols Esters	100 10 100	Same as phenols,	Plant A, Praction 6. Plant A, Praction 6.	Similar to unusual pattern in Praction 3 through mass 495.
8	23.8	100 10	Aliphatics Esters	100 10	Same as Plant A, Same as esters, 1	Fraction 2. Plant A, Praction 6.	No masses above 340. 100:155 10:254, 127 (diatomic iodine or probably napthyl iodide);

 $<sup>^{\</sup>overline{d}}_{ ext{Relative}}$  intensity of the mass-to-charge ratio (intensity relative to dominant ion).  $^{d}_{ ext{NO}}$  molecular weight range determination possible.  $^{e}_{ ext{NO}}$  composition determination possible.

TABLE 25 (continued)

Plant and	Praction		egories present		Subcategories present	
rganic fraction	weight,	Relative intensity	Category	Relative intensity	Specific compounds	Other unknown compounds present
			······································			· · · · · · · · · · · · · · · · · · ·
1	15.3	100 10	Aliphatics Estera	100 10	Same as Plant P, Fraction 1. Same as esters, Plant A, Fraction 6.	No masses above 451.
2	4.9	100 10	Organo-silicon species Esters	100 10	73(dominant), 147, 207, 221, 355 ions. d,e Same as esters, Plant A, Fraction 6.	No masses above 491.
3	3.8	100 1	Aliphatics Aromatics	100	Same as Plant A, Fraction 2. Same as Plant A, Fraction 2.	No masses above 296. 10:69(100), 41(80), 43(78)
		100	Estere	100	Same as esters, Plant A, Praction 6.	
4	5.5	10 10 100	Aliphatics Organo-silicon species Esters	10 10 100	Same as Plant F, Praction 1. 73(dominat), 147 ions. d. e Same as esters, Plant A, Fraction 6.	No masses above 477. 100:69(100), 41(80), 43(78)
5	11.8	10 100	Phanola Estera	10 100	Same as phenois, Plant A, Fraction 6. Same as esters, Plant A, Fraction 6.	No masses above 340. 10:69(100), 41(80), 43(78)
6	3.9	100	Aliphatics	100	Same as Plant F, Fraction 1.	No masses above 253.
		100	Phenols	100	Same as phenols, Plant A, Fraction 6.	10:156(100), 155(35)
		100	Esters	100	Same as esters, Plant A, Fraction 6.	(Possibly bipyridyl or phenyl cyclohexadiene, HM156 each)
,	22.6	10 100	Alighetics Alcohols/ethers	10 100	Same as Plant F, Fraction 1. Pattern indicates alcoholic ethers: 41, 43, 49 (dominant) and 55, 57, 59 (dominant) ion clusters. 9.0	No masses above 373.
ı		100	Esters	100	Same as esters, Plant A, Praction 6.	
8	19.8	100	Esters	100	Some as esters, Plant A, Fraction 6.	No masses above 279.
					Plant W	
1	20.9	100/100	Aliphatics	100	n-Paraffins.d,e	No masses above 424.
-		1	Saters	100 1	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	1:368(100), 369(45), 353(18)
2	3.4	100	Aliphatics	100	Same as Plant A, Praction 2.	No masses above 409.
		10	Aromatics	10	Same as Plant A, Fraction 2.	100:368(100), 369(45), 353(18)
		100	Esters	100	Same as esters, Plant A, Praction 6.	
3	13.2	10 1	Aliphatics Esters	10 1	Same as Plant A, Praction 2. Same as esters, Plant A, Praction 6.	No masses above 485. 100:368(100), 369(45), 353(18)
4	8.5	100	Aliphatica Esters	100 10	Same as Plant A, Praction 2. Same as esters, Plant A, Fraction 6.	No masses above 495. 10:69(100), 41(80), 43(78) 100:368(100), 369(45), 353(18)
5 '	5.7	100	Aliphatics	100	Semm as Plant A, Fraction 2.	No masses above 480.
		10	Phenol.	10 10	Same as phenol, Plant A, Praction 6.	100:368(100), 369(45), 353(18)
		10	Esters		Same as esters, Plant A, Fraction 6.	
6	4.2	100 100/10	Aliphatics Phenols	100 100	Same as Plant A, Fraction 2. Same as Plant F, Fraction 5.	No masses above 451.
		200,10	7,000,025	10	Same as phenols, Plant A, Fraction 6,	
		10	Esters	10	Same as esters, Plant A, Fraction 6.	
7	14.1	100 10	Aliphatics Esters	100 10	Same as Plant A, Fraction 2. Same as esters, Plant A, Fraction 6.	No masses above 495, 10:69(100), 41(80), 43(78) 100:384(100), 383(98), 368(90) 392(68)
8	11.7	100	Aliphatics	100	Same as Plant A, Fraction 2.	No masses above 499. 10:Cesium iodide or nathyl iod 100:69(100), 41(80), 43(78) 10:383(100), 368(96), 382(72), 369(64)

 $<sup>^{8}</sup>$  Relative intensity of the mass-to-charge ratio (intensity relative to dominant ion).  $^{4}$  No molecular weight range determination possible.  $^{e}$  No composition determination possible.

#### SECTION 7

#### BIOASSAY OF SECONDARY EFFLUENTS

The primary objective of the entire wastewater toxicity study is to determine the level of toxicity removal from secondary wastewater achieved by the tertiary treatment technologies selected for the ATMI/EPA BATEA study. To this end, the purpose of this Phase I screening study was to provide chemical and toxicological base-line data on secondary effluents from the 23 textile plants and to select plants for the toxicity removal study (Phase II).

Bioassays used were selected by EPA and included tests for assessment of both health and ecological effects (3). Health effects tests estimated the potential mutagenicity, potential or presumptive carcinogenicity, and potential toxicity of the secondary effluent wastewater samples to mammalian organisms. Ecological effects tests focused on the potential toxicity of samples to vertebrates (fish), invertebrates (daphnids and shrimp), and plants (algae) in freshwater, marine, and terrestrial ecosystems.

Biological testing, as well as chemical and physical parameters, must be considered when assessing the potential impact of industrial or municipal/industrial wastewaters on the aquatic environment. Biological testing involves determination of toxicity for samples of treated effluents. In a toxicity test, aquatic organisms will integrate the synergistic and antagonistic effects of all the effluent components over the duration of exposure.

Although toxicity tests with aquatic organisms can be conducted by applying wastewater samples directly to the test organisms, or by injection or feeding, most tests are conducted by exposing the test organisms to test solutions containing various concentrations of effluent samples. One or more controls are used to provide a measure of test acceptability by giving some indication of test organism health and the suitability of dilution water, test conditions, handling procedures, etc. A control test is an exposure of the organisms to dilution water with no effluent sample added. Bioassay tests are exposures of test organisms to dilution water with effluent samples added. Generally the most important data obtained from a toxicity test are the percentages of test organisms that are affected in a specified way by each concentration of wastewater sample added. The result derived

from these data is a measure of the toxicity of the effluent sample to the test organisms under the test conditions.

Acute toxicity tests are used to determine the level of toxic agent that produces an adverse effect on a specified percentage of test organisms in a short period of time. The most common acute toxicity test is the acute mortality test. Experimentally, 50% effect is the most reproducible measure of the toxicity of a toxic agent to a group of test organisms, and 96 hr is often a convenient, reasonably useful exposure duration. The 96-hr median lethal concentration (96-hr  $LC_{50}$ ) is most often used with fish and macroinvertebrates. Thus the acute mortality test is a statistical estimate of the  $LC_{50}$ , which is the concentration of toxicant in dilution water that is lethal to 50% of the test organisms during continuous exposure for a specified period of However, the 48-hr median effective concentration (48-hr  $EC_{50}$ ), based on immobilization, is most often used with daphnids. The terms median lethal concentration (LC<sub>50</sub>) and median effective concentration (EC<sub>50</sub>) are consistent with the widely used terms median lethal dose (LD<sub>50</sub>) and median effective dose (ED<sub>50</sub>), "Concentration" refers to the amount of toxicant respectively. per unit volume of test solution; "dose" refers to the measured amount of toxicant given to the test organism.

A total of 8 biological systems were used for wastewater toxicity evaluation, utilizing 21 different tester organisms. Specific tests used and the purpose of each bioassay are summarized in Table 26. The tests, testing conditions, and toxicity results for 23 secondary effluent samples are described in this section.

Under guidance of appropriate EPA Technical Advisors, four of the eight bioassays were performed at commercial laboratories experienced with the bioassays. The remaining four bioassays were performed by the EPA Technical Advisor, as shown in Figure 20. Bioassay results were sent to MRC and are included in the following sections.

### MICROBIOLOGICAL MUTAGENICITY

#### Introduction

The purpose of the mutagenicity bioassay was to determine if a chemical mutagen was present in secondary effluents. Nine different bacteria strains and one yeast strain were used in the test because of their individual sensitivities to various classes of chemical compounds. Secondary effluent samples were shipped to Stanford Research Institute (SRI) for mutagenicity testing by in vitro microbiological assays with Salmonella typhimurium (TA1535, TA1537, TA98, and TA100), Escherichia coli WP2, repairdeficient and proficient strains of Bacillus subtilis H17 and M45, and E. coli Pol A, and the yeast Saccharomyces cerevisiae

TABLE 26. BIOASSAY TESTS USED TO EVALUATE THE TOXICITY OF SECONDARY EFFLUENTS

Bioassay test system	Indicator organisms	Purpose of test
Microbial mutagenicity	Salmonella typhimurium (Ames test) (Strains TA1535, TA1537, TA98, TA100) Escherichia coli (Strains WP2, W3110, p3478) Bacillus subtilis (Strains H17 and M45) Saccharomyces cerevisial (Strain D3)	To determine if a chemical mutagen (possibly a carcinogen) is present. These microbial strains were selected because of their sensitivity to various classes of chemical compounds.
Cytotoxicity	Rabbit alveolar macrophage (RAM) (viability and ATP determinations) Chinese hamster ovary cells	To measure metabolic impairment and death in mammalian cells. These primary cell cultures have some degree of metabolic repair capability.
Freshwater static bioassay	Pimephales promelus (fathead minnow) Daphnia pulex (daphnid)	To detect potential toxicity to organisms in aquation environments.
Freshwater algal assay	Selenastrum capricornutum	To detect potential toxicity to aquatic plants.
Marine static bioassay	Cyprinodon variegatus (sheepshead minnow) Palaemonetes pugio (grass shrimp)	To detect potential toxicity to organisms in a marine environment.
Marine algal assay	Skeletonema costatum	To detect potential toxicity to marine plants.
Range finding acute toxicity	Rats (Charles River CD strain)	To detect potential toxicity to whole animals.
Terrestrial ecology	Soil microorganisms	To determine potential effects on soil ecosystems.

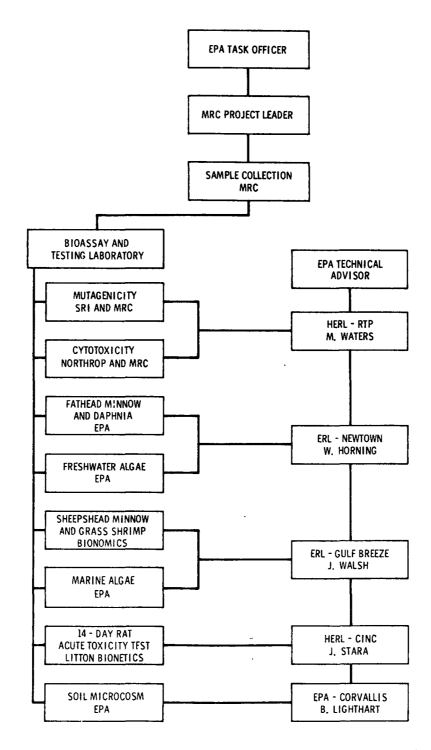


Figure 20. Laboratories and EPA technical advisors involved in biotesting of effluent samples.

D3. An Aroclor 1254-stimulated, rat-liver homogenate metabolic activation system was included in each procedure.

The assay procedure with *S. typhimurium* has proven to be 85% to 90% accurate in detecting mutagens, and it has about the same accuracy in identifying chemicals that are not carcinogenic (6). The assay procedure with *S. cerevisiae* is about 50% accurate in detecting carcinogens as agents that increase mitotic recombination. The *E. coli* WP2 assay and the microbial sensitivity assay are two additional methods of detecting mutagens. The combination of these assay procedures significantly enhances the probability of detecting potentially hazardous substances.

To date the most sensitive assay for deoxyribonucleic acid (DNA) damage is the induction of mutations in bacteria. The Ames test, the most highly developed of the bacterial mutagenesis tests, used mutant strains of S. typhimurium which were specially selected because of their abilities to detect specific types of mutations. For example, the TA1535 strain was designed to detect mutations due to base-pair substitutions. This strain responded particularly well to alkylating agents. Similarly, the TA1537 and TA1538 strains were used to detect frameshift mutations. Tester strains also included mutations which greatly increase their overall sensitivity to mutagens. One of these was responsibly for loss of the DNA excision repair system, while the other was responsible for loss of the lipopolysaccharide barrier that coats the surface of the bacteria, thereby enhancing the penetration of large molecules.

Mutant Salmonella tester strains lack the ability to synthesize histidine and are therefore unable to grow unless histidine is supplied. These bacteria are cultured in media containing minimal levels of histidine to sustain growth. Under these conditions only microscopic colonies of bacteria develop during the course of the test. However, if a mutagen is added to the medium, a reversion occurs in a certain number of the bacteria, restoring their ability to synthesize histidine. This reversion (backfuntation) is evidenced by the appearance of visible colonies in the histidine-limited agar, thus indicating the presence of a chemical mutagen.

Many compounds are not directly acting mutagen but are converted to active forms by normal body metabolism. A special microsomal preparation (usually liver) is added in the Salmonella tests to simulate  $in\ vivo$  metabolic actions. In practice, the substance is tested with and without this microsomal preparation to determine whether it requires metabolic transformation or is, itself, mutagenically active.

<sup>(6)</sup> McCann, J., E. Choi, E. Yamasaki, and B. N. Ames. Detection of Carcinogens as Mutagens in the Salmonella/Microsome Test: Assay of 300 Chemicals. Proceedings of the National Academy of Science, 72:5135-5139, 1975.

The testing procedure used for each type of microbe and the bioassay results of each test are described below.

# Bioassay Procedures

Each secondary effluent sample was shipped and stored in the laboratory at 4°C. Preliminary experiments on the first two samples received indicated microbial contamination when an aliquot of the sample was incubated on a culture medium. Therefore, each sample was filtered before it was tested in any microbial system. Nalgene filters (0.45  $\mu m$ ) were used. Approximately 50 x 10<sup>-6</sup> m³ of each sample was filtered; the remaining 200 x 10<sup>-6</sup> m³ was stored for possible future testing.

Four strains of S. typhimurium (TA1535, TA1537, TA98, and TA100) were obtained from Dr. Bruce Ames of the University of California at Berkeley and stored in 10% sterile glycerol at -80°C. New stock cultures were prepared every two months from single colony reisolates that were checked for their genotypic characteristics and for presence of the plasmid. For each experiment, an inoculum from the stock cultures was grown overnight at 37°C in nutrient broth (Oxoid, CM67). After stationary overnight growth, the cultures were shaken for 3 hr to 4 hr to ensure optimal growth.

The metabolic activation mixture used for each experiment consisted of

- 1.0 x  $10^{-6}$  m<sup>3</sup> of S-9 rat liver fraction
- 0.2 x  $10^{-6}$  m<sup>3</sup> of magnesium chloride (0.4 M) and potassium chloride (1.65 M)
- $0.05 \times 10^{-6} \text{ m}^3$  of glucose-6-phosphate (1 M)
- $0.4 \times 10^{-6} \text{ m}^3$  of nicotine adenine dinucleotide phosphate
- 5.0 x  $10^{-6}$  m<sup>3</sup> of sodium phosphate (0.2 M, pH 7.4)
- 3.35 x  $10^{-6}$  m<sup>3</sup> of water

For each experiment, the following solutions, listed in the order of addition, were added to a sterile 13 mm x 100 mm test tube placed in a 43°C heating block:

- 2.00 x  $10^{-6}$  m<sup>3</sup> of 0.6% agar (containing 0.05 mM histidine and 0.05 mM biotin)
- $0.05 \times 10^{-6} \text{ m}^3$  of indicator organisms
- $0.50 \times 10^{-6} \text{ m}^3$  of metabolic activation mixture
- $0.05 \times 10^{-6} \text{ m}^3$  of the secondary effluent sample

Dimethylsulfoxide (DMSO) was added to each sample to improve the water solubility of organic compounds. The resulting mixture was gently stirred and poured onto minimal agar plates. These plates

consisted of 15 kg of agar, 50 kg of glucose, 0.2 kg of magnesium sulfate (MgSO<sub>4</sub>•7 H<sub>2</sub>O), 2 kg of citric acid monohydrate, 10 kg of potassium orthophosphite, and 3.5 kg of sodium ammonium phosphate, per cubic meter. After the top agar had set, the plates were incubated at 37°C for 2 days. Then the number of revertant colonies was counted. Each sample was run in duplicate.

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All samples were run with both positive and negative controls, with and without metabolic activation at five concentrations. Positive controls were run using various combinations of 2-anthramine, 9-aminoacridine,  $\beta$ -propiolacetone, sodium azide, n-methyl-n'-nitro-n-nitrosoguanidine, daunomycin, and 2-(2-furyl)-3-(5-nitro-2-furyl) acrylamide (AF2). Controls were also run on DMSO.

The S. cerevisiae tester strain was stored at -80°C. For each experiment, the tester strain was inoculated in 1% tryptone and 0.5% yeast extract and grown overnight at 37°C with aeration.

The *in vitro* yeast mitotic recombination assay in suspension was conducted as follows. The overnight culture was centrifuged, and cells were resuspended at a concentration of  $10^8$  cells per  $10^{-6}$  m<sup>3</sup> in a 67-mM phosphate buffer (pH 7.4). The following solutions were added to each sterile test tube:

- 1.30 x  $10^{-6}$  m<sup>3</sup> of S. cerevisiae
- 0.50 x  $10^{-6}$  m<sup>3</sup> of either the metabolic activation mixture or buffer
- 0.20 x  $10^{-6}$  m<sup>3</sup> of the secondary effluent sample

Because many organic chemicals are not appreciably water soluble, DMSO was added as the solvent for the secondary effluent sample. Several doses of the chemical [up to 5%, weight-to-volume ratio (w/v) or volume-to-volume ratio (v/v)] were tested in each experiment, and appropriate controls were included.

The suspension mixture was incubated at 30°C for 4 hr on a roller drum. The sample was diluted serially in a sterile physiological saline, and volumes of  $0.2 \times 10^{-6} \text{ m}^3$  of the  $10^{-5}$  and  $10^{-3}$  dilutions were spread on tryptone-yeast agar plates; five plates were used for the  $10^{-3}$  dilution and three plates for the  $10^{-5}$  dilution. These plates were incubated for 2 days at 30°C, followed by 2 days at 4°C to enhance the development of the red pigment. Plates of the  $10^{-3}$  dilution were scanned with a dissection microscope at  $10 \times 10^{-3}$  dilution were scanned with a dissection microscope at  $10 \times 10^{-3}$  dilution, and the number of red colonies or red sectors (mitotic recombinants) was recorded. The surviving fraction of organisms was determined from the number of colonies appearing on the plates of the  $10^{-5}$  dilution.

The number of mitotic recombinants was calculated per  $10^5$  survivors. A positive response in this assay was indicated by a

dose-related increase of more than threefold in the absolute number of mitotic recombinants per  $10^{-6}$  m<sup>3</sup> as well as in the relative number of mitotic recombinants per  $10^{5}$  survivors. Positive, negative, and reagent controls were run at four concentrations with each test, with and without metabolic activation. Positive controls were performed using 1,2,3,4-diepoxybutane.

A procedure similar to the Ames Salmonella assay was used to measure the reversion of  $E.\ coli$  WP2 to tryptophan independence. However, the minimal agar was supplemented with 1.25 g of Oxoid nutrient broth (CM67) per  $10^{-3}$  m³ to provide each plate with the trace of tryptophan required for enhancement of any mutagenic effect of the test chemical. No additional tryptophan was added to the top agar. The positive controls used for the Ames test were also used for the  $E.\ coli$  WP2 test.

As an alternative to reversion of the mutated tryptophan gene; WP2 may undergo a forward suppressor mutation in a tryptophan transfer ribonucleic acid (RNA) gene to obtain tryptophan independence. The test did not distinguish experimentally between true revertants and suppressor mutants (although the latter tend to form smaller colonies).

E. coli strains W3110 and p3478 and B. subtilis strains H17 and M45 were stored in the laboratory at -80°C. Inoculums from the frozen stocks were grown overnight at 37°C with shaking in a nutrient broth. The broth contained 1% of tryptone and 0.5% yeast extract, and was supplemented with 5  $\mu g$  of thymine per  $10^{-6}$  m³. To 2 x  $10^{-6}$  m³ of top agar containing 0.6% agar was added 0.1 x  $10^{-6}$  m³ of the test culture. This suspension was mixed and poured onto plates containing nutrient broth and 2% agar.

After the soft agar solidified, a sterile filter disc impregnated with a secondary effluent sample was placed in the center of the plate. Plates were incubated at 37°C for 16 hr, and the width of the zone of toxicity or inhibition of growth was measured. Several concentrations of chemical were tested to accurately detect differences in the zones of growth inhibition, because higher initial concentrations lead to steep concentration gradients that may reduce the difference in growth inhibition of the test strains.

The positive control for this assay was 2 mg of 1-phenyl-3,3-dimethyltriazene placed on the disc. Larger zones of inhibition were observed in the DNA repair-deficient strains (p3478 and M45). The negative control was 20  $\mu g$  of chloramphenicol. Equal zones of inhibition were observed in all four strains since the toxicity of chloramphenicol did not depend on a mechanism that leads to DNA damage.

### Results

SRI tested secondary effluent wastewater samples from 22 of the 23 basic textile plants for bacterial mutagenicity using 10 tester microorganisms and 3 different assay systems. The secondary effluent sample from Plant E was lost in shipment.

The voluminous amount of mutagenicity raw data generated by SRI are given in Reference 7 and a summary of the results is given in the following paragraphs.

All 22 of the samples were tested twice in the standard Ames Salmonella microsome procedure using four test strains: TAl535, TAl537, TA98, and TAl00. A metabolic activation mixture was included in each experiment. Each sample was tested to a maximum dose of 1 x  $10^{-6}$  m³ of sample per plate (the maximum amount possible in 2 x  $10^{-6}$  m³ of top agar). The second experiment was a confirming experiment. None of the samples caused an increase in the number of histidine-independent revertants above the normal background, thus no chemical mutagen was detected.

Twenty-two samples were tested in the *E. coli* WP2 strain. None of the samples caused an increase in the number of tryptophan-independent revertants above the normal background. Twenty-two samples were tested in the *S. cerevisiae* D3 suspension assay, with and without a metabolic activation system. The maximum concentration tested was 50% v/v. None of these samples caused an increase in the number of mitotic recombinants above the normal background.

The samples were also tested in the microbial inhibition assay using DNA repair-deficient and -proficient strains of  $E.\ coli$  and  $B.\ subtilis$ . The maximum dose tested was 20 x  $10^{-9}$  m<sup>3</sup>. Each sample was applied to a filter disc on the plate. None of the samples was toxic to any of the strains of the organisms used.

All 22 secondary effluent samples were tested in three assay systems. In each assay, the maximum possible dose was tested. None of the samples gave a mutagenic or toxic response in any strain in any assay or experiment.

<sup>(7)</sup> Poole, D. C. and V. F. Simmon. Final Report of in Vitro Microbiological Studies of Twenty-two Wastewater Effluent Samples. Contract 68-01-2458, U.S. Environmental Protection Agency, Biomedical Research Branch, Research Triangle Park, North Carolina, November 1977. 111 pp.

## MRC Ames Testing

To provide backup duplicate results to the Ames tests performed by SRI, MRC performed the Ames test on eight randomly selected secondary effluent samples from Plants D, H, J, M, P, R, Y, and Z.

Biotest procedures used by MRC were the same as those used by SRI. Final test results indicated no positive responses on filtered, unconcentrated effluent samples. Samples from Plants D, P, and R were rerun on selected strains with no indication of a positive response.

#### CYTOTOXICITY ASSAY

### Introduction

Cytotoxicity (cell toxicity) assays were performed to measure quantitatively any cellular metabolic impairment and death resulting from exposure in vitro to secondary effluent samples. Primary cell cultures, such as the rabbit alveolar macrophage (RAM) used in this study, exhibit many of the metabolic and functional attributes of the in vitro state. These cells can therefore combat, to some degree, the effects of chemical mutagens on mammalian cells.

Recently this system has been applied in evaluating the relative cellular toxicity of hazardous metallic salts and industrial air particulates (8). As compared to conventional whole animal tests for acute toxicity, these cytotoxicity assays are more rapid, less costly, and require less sample. These tests provide useful information about the relative toxicity of unknown samples. However, it should be understood that because the assays employ isolated cells and not intact animals, they can provide only preliminary information about the ultimate human health hazards of toxic chemicals.

Two tests were used to measure the toxic effects of secondary effluent samples on rabbit alvaolar macrophage: viability and adenosine triphosphate (ATP) production. Viability refers to the ability of cells to survive, and it was measured by the trypan blue dye exclusion method. Living (viable) cells do not absorb trypan blue dye. Therefore, the measure of cell mortality is the number of blue (dead) cells counted after exposure to the sample.

Adenosine triphosphate (ATP) is a coenzyme in mammalian cells that plays an important role in energy metabolism. Living cells synthesize ATP. Therefore, another measure of cell mortality or

<sup>(8)</sup> Waters, M. D., D. E. Gardner, C. Aranyi, and D. L. Coffin. Metal Toxicity of Rabbit Alveolar Macrophages in Vitro. Environmental Research, 9(1):32-47, 1975.

inhibition was the quantity of ATP produced by a cell culture exposed to the secondary effluent sample compared to the amount produced by an identical culture not exposed to the solution.

Both of these methods were used to evaluate secondary effluent toxicity to RAM. The measure of toxicity was expressed as  $EC_{20}$  or  $EC_{50}$ ; i.e., the concentration of secondary effluent that inhibits RAM metabolism by 20% or 50% over a specified time period (20 hr).

The following section discusses the test procedure used to evaluate secondary effluent toxicity and the bioassay results.

## Bioassay Procedure

Rabbit alveolar macrophage primary cell cultures were obtained from New Zealand white rabbits of both sexes. Each of the 23 secondary effluent samples was filtered through a 0.45- $\mu$ m filter. There was no concentrating of the samples prior to testing. Each sample was prepared in five dilutions: 6, 20, 60, 200, and 600 x  $10^{-6}$  m³ sample per  $10^{-3}$  m³ solution. To each concentration, fetal calf serum (heat inactivated) was added to give a final serum concentration of 10%. Antibiotics were added to give  $10^{-6}$  units per  $10^{-6}$  m³ penicillin, and 100  $\mu$ g combined streptomycin and kanamycin per  $10^{-6}$  m³. The pH of each concentration was recorded, but no adjustments were made.

Samples were then added to Falcon cluster dishes,  $1.5 \times 10^{-6} \text{ m}^3$  per well. A volume of  $0.5 \times 10^{-6} \text{ m}^3$  of complete 1X Medium 199 (with 10% fetal calf serum and antibiotics) containing approximately  $2 \times 10^6$  rabbit alveolar macrophages per  $10^{-6} \text{ m}^3$  was added to each well and gently mixed. Dishes were then incubated for 20 hr at 37°C in a humidified atmosphere of 5% CO<sub>2</sub> in air on a Belico rocker platform.

At the end of the 20-hr exposure, the medium containing unattached cells was removed from each well and transferred to a separate test tube. A volume of  $10^{-6}$  m<sup>3</sup> of 0.25% trypsin was added to each well and left until the cells were removed from the dish. This was then combined with the original pouroff and mixed to inactivate the trypsin.

Cell counts, viabilities, and ATP determinations were then performed.

Viability was determined by trypan blue dye exclusion. Filtered samples were counted in the Cytograf; unfiltered samples, because of the particulate matter present, were counted in a hemocytometer. For hemocytometer counts, 1 part of 0.4% trypan blue was added to 5 parts cell suspension and counted after a 5-min exposure. For Cytograf counts, dilutions, usually fourfold, were made with cold

0.85% saline to yield a suspension of no more than  $2 \times 10^5$  cells/ $10^{-6}$  m<sup>3</sup>. Trypan blue was diluted (immediately before use) with 0.85% saline to a final concentration of 0.01% and added to an equal volume of cell suspension. Simultaneous determinations of cell viability and cell numbers per  $10^{-6}$  m<sup>3</sup> of cell suspension were made. The numbers of viable cells were expressed as a percentage of the number of cells in control cultures; viability was expressed as the concentration of secondary effluent which inhibited 20% and 50% of the test cells (EC<sub>20</sub> and EC<sub>50</sub>).

ATP was determined according to a procedure supplied with the Du Pont Model 760 Luminescence Biometer. Dimethyl sulfoxide  $(0.4 \times 10^{-6} \text{ m}^3)$  was used to extract ATP from a 0.1  $\times 10^{-6} \text{ m}^3$  aliquot of trypsinized cell suspension containing 0.3 to 0.4  $\times 10^{5}$  cells. After 2 min at room temperature, 2.5  $\times 10^{-6}$  m³ of cold 0.1 M morpholinopropane sulfonic acid (MOPS) at pH 7.4 was added to buffer the extracted sample. The tube containing the buffered sample was then placed in an ice bath. Aliquots of  $10^{-8}$  m³ were injected into the luminescence meter's reaction cuvette containing 0.7 mM luciferin (crystalline), 100 units luciferase (purified and stabilized), and 0.01 M magnesium sulfate in a total volume of  $10^{-7}$  m³ of 0.01 M MOPS buffer, pH 7.4 at 25°C. Light emitted from the reaction cuvette was measured photometrically in the luminescence meter and was proportional to the ATP concentration of the sample. ATP values were expressed per  $10^6$  cells and as a percent of the control cells.

Nutrient agar plates were streaked with  $0.5 \times 10^{-6}$  m<sup>3</sup> from each sample (unfiltered) and antibiotic sensitivity discs were added for a 24-hr incubation period. This was done to ascertain what antibiotics were capable of suppressing growth of any bacteria present in the samples. Antibiotics present in the culture medium were found to be capable of inhibiting bacterial growth so that the samples could be tested unfiltered.

### Results

All determinations were performed in duplicate or triplicate. The voluminous amount of cytotoxicity raw data generated by Northrop are given in Reference 9 and a summary of the results are given below.

Because cell viability could be considered a binomial response, the arc-sine transformation was employed in the regression

<sup>(9)</sup> Campbell, J. A., H. F. Stack, and P. R. Williams. Cyto-toxicity Screening of Twenty-three Textile Mill Effluent Water Samples Utilizing the Rabbit Alveolar Macrophage Assay. Contract 68-02-2566, U.S. Environmental Protection Agency, Biomedical Research Branch, Research Triangle Park, North Carolina, December 1977. 86 pp.

analysis. Samples listed as nondeterminable were such high extrapolations that they could not be considered significant estimates.

Table 27 shows the estimated  $EC_{20}$  and  $EC_{50}$  concentrations for the filtered effluents. The EC values are the concentrations expected to cause a decrease in viability and ATP by 20% and 50%, respectively. Control values were routinely 92% to 100%.

TABLE 27. ESTIMATED EC<sub>20</sub> AND EC<sub>50</sub> VALUES FOR CYTOTOXICITY SCREENING OF FILTERED SECONDARY EFFLUENT SAMPLES

	Pe	ercent	effluen	t		Pe	ercent	effluen	t
	Viab	ility	AT	a P		Viab	ility	AT	a
Plant	EC <sub>20</sub>	EC <sub>50</sub>	EC <sub>20</sub>	EC <sub>50</sub>	Plant	EC <sub>20</sub>	EC <sub>50</sub>	EC <sub>20</sub>	EC <sub>50</sub>
A					M				
Bb					N	13.3		3.8	12.8
c c c	16.8		6.1	33.5	P				
cc			d	_d	R				•
D			-	~	s	ď	ď	d	d
E					T	-	-	2.5	-
F			9.4		ช				
G					v				
H					W			13.7	
J					x			4.8	
ĸ					Y				
L			4.0	35.1	Z			•	

Note.—Blanks indicate data not determinable.

Samples from Plant N and C caused the greatest response by viability and ATP determinations. The response of Sample C was largely due to its high (9.1) pH. When testing was repeated with the pH adjusted to 7.2, there was much less response.

Because the antibiotic sensitivity testing showed that samples could be tested without prior filtration, five samples were retested without prior filtration. Much of the solid material removed by filtration appeared to be biological material (e.g., algae) by microscopic observation. In each instance, the unfiltered sample caused greater response than the filtered one.

# MRC Clonal Assay

MRC performed clonal assay acute toxicity tests using Chinese hamster ovary cells (CHO-Kl) on selected secondary effluent

Adenosine triphosphate. bpH equals 9.1 not adjusted before testing.

C pH adjusted to 7.2. Test not performed.

samples. The purpose of these tests was to evaluate the response of another test system to complex environmental samples. MRC has developed this in vitro clonal assay for measuring acute toxicity of compounds using CHO-Kl cells. This test is a modification of a clonal assay described by Malcolm (10). Preliminary studies indicate that the sensitivity of the CHO-Kl clonal assay appears to be two orders of magnitude greater than that of the WI-38 assay.

Secondary effluent samples from eight textile plants (Plants D, H, J, M, P, R, Y, and Z) were selected at random and analyzed according to the following procedure.

#### Test Procedure --

Effluent samples for this test were filtered through  $0.45-\mu m$  and  $0.22-\mu m$  filters. Samples were run at 5 to 7 concentration levels using approximately 300 to 500 cells that had been plated on the previous day. After incubation at 37°C for 6 days to 7 days, the media and sample were removed and the cells were fixed, stained, and counted. Results are reported as experimental versus controls or percent survival. A detailed test procedure is given in Table 28.

#### Results--

Screening tests were first performed on the eight samples to determine whether any of them were toxic and in what concentration range. Results showed that secondary effluent samples from Plants P, R, Y, and Z had no acute toxicity to the CHO-Kl cells, but samples from Plants D, H, J, and M did exhibit toxicity. The latter four samples were therefore rerun using narrower sample concentrations.

Test results are presented in Figure 21. The range bar associated with each data point corresponds to the standard deviation of that value.

Graphical interpolation of Figure 21 yields the following  $LC_{50}$  values (in percent of secondary effluent sample) for Plants D, H, J, and M: 2.4%, 13.3%, 18.5%, and 3.0%, respectively. These values indicate that the secondary effluent from Plants D and M contain chemical species more acutely toxic to CHO-Kl cells than do samples from Plants H and J.

<sup>(10)</sup> Malcolm, A. R., B. H. Pringle, and H. W. Fisher. Chemical Toxicity Studies with Cultured Mammalian Cells. In: Bioassay Techniques and Environmental Chemistry, G. E. Glass, ed. Ann Arbor Science Publishers, Inc., Ann Arbor, Michigan, 1974. pp. 217-230.

### TABLE 28. CHO-K1 CLONAL CYTOTOXICITY TEST

Cell line: Chinese hamster ovary epithelial cells ATCC No. CCL 61

Medium:  $F-12 \text{ GIBCO No. H}-17 \ 10.8 \times 10^3 \text{ g/m}^3$ 

Sodium hydrogen carbonate

10% Fetal calf serum, virus, mycoplasma screened

GIBCO No. 629

Incubation: 37°C, 5% CO2, Saturated humidity

Samples: 6 Controls (blank)

5 to 7 Concentrations of test compound in triplicate

5 Concentrations of a positive toxic control in triplicate

### Test procedure

To stock CHO-Kl, add 5 x  $10^{-6}$  m<sup>3</sup> 0.25% trypsin at 37°C for 5 min.

Shake cells and add to centrifuge tube.

Add 5 x  $10^{-6}$  m<sup>3</sup> media to flask, shake, and add to centrifuge tube.

Centrifuge 5 min at 1,200 g, pour off liquid, retaining cells.

Add 10 x  $10^{-6}$  m<sup>3</sup> medium, shake, centrifuge 5 min, pour off medium.

Add 10 x  $10^{-6}$  m<sup>3</sup> medium, shake.

Make hemocytometer count of trypsinized cells.

Dilute so that  $5 \times 10^{-6} \text{ m}^3$  media contain 300 to 500 cells.

Add 5 x  $10^{-6}$  m<sup>3</sup> media and cells to T-25 flasks.

Incubate 12 hr to 18 hr to allow attachment using normal media.

Replace  $5 \times 10^{-6} \text{ m}^3$  of media and sample.

Incubate 6 days to 7 days total.

Fix with 10% formaldehyde/0.5% sodium chloride/4% methanol for 30 min.

Stain with crystal violet (0.04% for 15 min).

Count clonal colonies of remaining cells macroscopically using Fisher Count-All Model 600.

Score with respect to experimental vs. controls as percent survival.

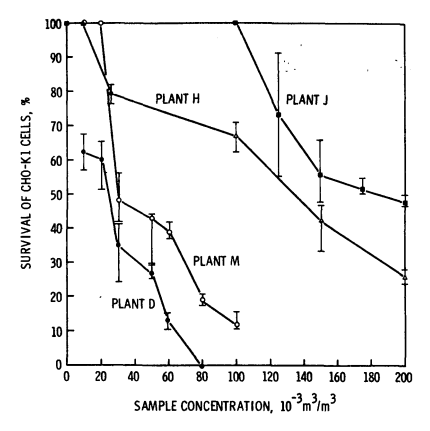


Figure 21. Results of CHO-K1 clonal assay.

## FRESHWATER ECOLOGY TOXICITY

### Algal Assay Bottle Test

Algal assay was performed to estimate the potential toxicity of secondary effluents on aquatic plants. The algal assay was based on the principle that growth is limited by the essential nutrients that are in shortest supply. The test was designed to quantify the biological response (algal growth) to changes in concentrations of nutrients, and to determine whether various effluents were stimulatory or inhibitory to algae. These measurements were made by adding a selected test alga to the effluent and determining its growth response.

The freshwater algae testing series was performed at the Environmental Research Laboratory (ERL, Corvallis, Oregon) under the direction of W. E. Miller. Each effluent sample (0.01 m³ from each of the 23 plants) as apportioned in three autoclavable, 0.004-m³ plastic bottles, was packed in ice and flown air freight to Corvallis, Oregon. Most samples arrived at the laboratory within 36 hr from the time they were shipped. The following paragraphs summarize the sample handling and analysis procedures used at ERL. An EPA manual (3) gives a detailed description of the procedure.

Each 0.01-m<sup>3</sup> textile effluent sample was thoroughly composited in the laboratory in a 0.02-m<sup>3</sup> cubitainer, then redistributed into bottles. A 0.001-m<sup>3</sup> aliquot was taken for soil microcosm studies. Samples were stored in the dark at 5°C.

Well water from the Western Fish Toxicology Station was used for dilution water and for control samples. The water was filtered with a  $0.45-\mu m$  porosity membrane filter and pad to remove any particulate matter before mixing with raw effluent. Dilutions, Table 29, were made in a  $0.02-m^3$  cubitainer for a total of  $0.016~m^3$  of diluted effluent.

TABLE 29. DILUTIONS USED FOR FRESHWATER ALGAL TESTS

Dilution, %	Effluent volume, x 10 <sup>-4</sup> m <sup>3</sup>	Well water volume, x 10 <sup>-3</sup> m <sup>3</sup>
2	3.2	15.68
· 5	8.0	15.20
10	16.0	14.40
20	32.0	12.80
Total volume used	5,9.2	58.08

Each dilution of 0.016 m<sup>3</sup> was divided into four parts of 0.004 m<sup>3</sup> each for four different treatments: One part was autoclaved and filtered, one was autoclaved only, one was filtered and autoclaved, and one was filtered only.

Diluted samples were autoclaved in polypropylene bottles (washed with 10% hydrochloric acid and autoclaved before use) at a pressure of 108 kPa at 10 min/ $10^{-3}$  m<sup>3</sup>. Diluted samples were filtered with a 0.45- $\mu$ m porosity membrane filter and pad. During and after treatment, diluted samples were stored in the dark at 5°C.

The pH values were taken on raw effluent and on diluted effluent before and after treatment to determine changes in pH that can affect growth. Each secondary effluent sample was analyzed by MRC for the following nutrient indicators: o-phosphate, ammonia, nitrite, nitrate, total Kjeldahl nitrogen, total phosphorus, and total organic carbon, Table 30 (4, 5).

The algal assay bottle test procedure used Selenastrum capricornutum Printz as the test alga and was used to assess algal
growth response to 23 secondary effluent wastewater samples.
Growth response was measured as net biomass produced (gram dry
weight per cubic meter of sample), i.e., the total biomass in
the sample minus the biomass produced in a control sample using
well water. Algal response was expressed as percent stimulation
or inhibition at the 20% wastewater concentration as compared
to the control sample. Tests were performed on two types of

TABLE 30. NUTRIENT ANALYSIS OF SECONDARY EFFLUENT SAMPLES

				Nutrient,	$g/m^3$		
_				Total			Total
Plant				Kjehldahl		Total	organic
<u>code</u>	Nitrite	Nitrate	Ammonia	nitrogen	o-Phosphate	phosphorus	carbon
A	0.06	1.9	12.8	21.3	1.0	0:39	
В	0	0.002	2.5	4.2	7.3	6.0	29.0
С	4.64	23.3	3.4	1.85	1.08	0.50	219
D	7.5	0.08	0.20	4.5	1.6	2.48	36
E	<0.02	40	1.13	6.75	1.11	1.19	31.8
F	0.04	0	1:54	14.8	0.56	9.7	57.5
G	<0.02	1.3	0.40	12	5.02	6.29	
H	16.8	0.69	0.02	5 <b>.5</b>	0.06	0.39	84
J	11.9	0.24	1.25	2.3	0.28	1.68	150
K	<0.02	0.16	0.40	5.25	0.82	1.05	19.1
L	0.86	13.5	0.5	1.9	0.88	1.98	42.9
M	0.27	0.30	0.84	6.15	0.55	3.03	-84
N	<0.02	5.5	12.8	17.0	11.2	2.05	90.4
P	7.3	0.08	0.20	2.9	0.02	3.12	54
R	1.82	0.015	2.6	19.8	0.06	1.82	244
S	<0.02	0.23	1.65	40	3.1	5.23	261
T	0.04	0.80	13.6	30.2	6.4	14.9	142
U	<0.02	0.8	5.44	11.3	2.96	2.43	
V	0.07	1.0	1.26	6.3	0.11	0.76	
W	0.145	12.3	0.38	7.4	0.075	0.50	199
X	0.44	0.033	0.65	5.25	0	4.87	94
Y	8.5	0.05	1.9	5.75	9.9	14.3	41.8
Z	<0.02	0.94	0.85	4.75	0.40	0.54	40

Note.—Blanks indicate analysis not performed.

pretreated samples: 1) filtered followed by autoclaving, and 2) filtered only. Results are given in Table 31.

TABLE 31. RESULTS OF FRESHWATER ALGAE BIOASSAY TESTS

1	-Day growth as com	response in 200 pared to contro	secondary e	fluent	
	Pilter	ed only	Filtered-autoclaved		
	Percent	Percent	Percent	Percent	
Plant	inhibition	stimulation	inhibition	stimulation	
A	53	0	33	0	
В	Ö	83	Ö	44	
č	ŏ	187	Ó	146	
Ď	Ŏ	100	0	125	
Ē	958	0	95ª	0	
P	Ö	598	0	866	
Ğ	0	390	0	578	
H	92	0	95	0	
J	0	76	0	217	
K	0	57	0	243	
L	81	0	0	98	
M	0	149	0	291	
N	95ª	0	958	0	
P	0.	38	5.	0	
R	95b	0	93b	0	
S	0	382	0	365	
T	0	1,911	0	2,362	
Ū	0	377	0	639	
v	0	232	D	503	
W	95	0	95	0	
x	0	163	0	348	
Y	0 ;	261	0	365	
Z	84	0	17	0	

<sup>395%</sup> growth inhibition in 2% solution of secondary effluent.

Five distinct growth patterns were discerned in the initial screening; they are described in the following sample subsets.

Sample inadvertently collected prior to the settling pond.

Two samples (Plants E and N) failed to support growth of the test alga in all waste concentration levels (2%, 5%, 10%, and 20%). This response has been attributed to toxicity since nitrogen (N) and phosphorus (P) concentrations (2% level) were adequate to support up to 8.0 g dry wt/m $^3$  of S. capricornutum.

Seven samples (Plants B, D, J, K, P, X, and Z) supported similar growth at all dilutions ranging from 5.5 to 16.0 g dry wt/m³ of S. capricornutum, depending on nutrient bioavailability within waste samples. Failure of the growth response to increase at a rate proportional to the analyzed incremental concentration of nitrogen and phosphorus in these samples suggests that some constituent other than nitrogen and phosphorus is limiting the maximum yield. However, the suboptimal yield obtained is considered to be stimulatory for the support of test alga.

Four samples (Plants H, L, R, and W) supported growth at the 2% and 5% levels but inhibited growth at the 10% and 20% concentrations.

Seven samples (Plants C, F, G, M, S, V, and Y) supported increasing growth of test alga with similar increase in waste concentration, proportional to the bioavailable (but not chemically analyzed) inorganic nitrogen content. The growth thus obtained indicates that these wastes are highly stimulatory for the support of test alga.

Two samples (Plants T and U) were extremely stimulatory, 124.5 and 29.1 g dry  $\text{wt/m}^3$ , respectively, for these wastes at the 20% concentration level. Growth obtained in these samples was directly proportional to their chemical nitrogen and phosphorus content, indicating its complete availability for support of algal growth.

Using test results from the filtered-and-autoclaved and filtered-only samples, ERL was able to rank the secondary effluent samples in terms of inhibition and stimulation; results are shown in Table 32. In those samples categorized as nontoxic, the primary limiting factor regulating growth response of the test alga was bioavailability and utilization of the total soluble inorganic nitrogen (TSIN equals nitrite, nitrate, and ammonia). Shiroyama, Miller, and Greene (11) demonstrated that maximum yield for Selenastrum capricornutum Printz is predictable provided the TSIN is known, other essential nutrients are in adequate supply, and toxicants are absent. Under these conditions 0.001 g/m³ of TSIN can yield 0.038 g/m³ dry weight of the alga. Based on this

<sup>(11)</sup> Shiroyama, T., W. E. Miller, and J. C. Greene. The Efforts of Nitrogen and Phosphorus on the Growth of Selenastrum Capricornutum Printz. EPA-606/3-75-034, U.S. Environmental Protection Agency, Corvallis, Oregon, March 1975. pp. 132-142.

TABLE 32. RELATIVE RANKING OF TEXTILE PLANTS BY TOXIC AND STIMULATORY EFFECTS OF SECONDARY WASTEWATER ON S. CAPRICORNUTUM

Fi.	tered and autoclaved	samples	Fi.	ltered only	samples
Toxic rating	Plant rank	Stimulatory rating	Toxic rating	Plant rank	Stimulatory rating
rating	Talix	racing	racing	Lank	rating
lost toxic	E, N	Nonstimulatory	Most toxic	E, N	Nonstimulatory
	W, aH	P		W _	•
east toxic	R, a P, L, B, Z, A	•		R, a H	•
ontoxic	D	Least stimulatory	Least toxic	A, Z, L	
	С	<b>.</b>	Nontoxic	P	Least stimulator
•	Y	•	я	J	
	K		•	В	
*	J		•	ĸ	•
	M	•	•	M	₩
	X	•		D	•
	Š	•	•	x	•
	P	-		Ÿ	•
	v	•		č	•
H	Ġ	•		ii.	•
#	ii	•		Ğ	
	Ť	Most stimulatory		r P	
	•	nost stimulatory		v	
			#	Tr.	Most stimulatory

<sup>&</sup>lt;sup>a</sup>Sample inadvertently collected prior to the setting pond.

information a linear regression between biomass produced in the textile waste samples and that predicted from the TSIN content of the test water showed a correlation coefficient in the filtered followed by autoclaving and filtered only samples of 0.980 and 0.989, respectively, resulting in dry weight yield per unit concentration of TSIN relationship of 0.910 and 0.997, respectively. These relationships indicate that the complexity of the textile water samples under the two pretreated conditions does not affect the algal from utilizing the essential nutrients in obtaining its maximum yield.

### Acute Static Bioassays with Freshwater Fish and Daphnia

The acute static bioassay technique with freshwater animals provided an easy measure of toxicity and was recommended by EPA for the wastewater assessment (3). Fathead minnow (Pimephales promelus) and Daphnia pulex were the selected test animals because they are a readily available, hardy species, and they can be conveniently and economically maintained in a laboratory (3).

Primary objections to the following procedure are that the recommended dilution water may not closely simulate receiving water characteristics, and the fathead minnow may not be representative of the most sensitive species in a given geographical area. However, the procedure does adequately serve to develop relative toxicity data for the purpose of ranking industries based on the toxicity of their effluents.

This series of tests was performed at the EPA Fish Toxicology Station (Newtown, Ohio) under the direction of Mr. W. Horning. Each of the three 0.02-m<sup>3</sup> glass bottles of effluent from 23 plants was packed in ice and shipped by air freight.

The fathead minnow test utilized 16 x 10<sup>-3</sup> m<sup>3</sup>, wide mouthed jars. A control dilution and effluent dilutions of 100%, 60%, 36%, 21.6%, 13%, 7.8%, and 4.7% effluent were set up. A total volume of 0.015 m<sup>3</sup> was used in each test jar. Ten fish were randomly distributed to each jar. Duration of the test was 96 hr, and temperature ranged from 20.5°C to 21.6°C. Aeration was not used during the test. At the end of the test, the fish length and weight were determined. Length ranged from 28 mm to 44 mm, with an average length of 33 mm. Weight ranged from 0.18 g to 0.80 g, with an average weight of 0.29 g.

Dissolved oxygen, pH, specific conductivity, temperature, and turbidity were determined in each test jar at the beginning of the test and every 24 hr through the end of the test. At the end of each 24-hr period, the number of fish surviving was recorded, and dead fish were removed.

The same test procedures and dilutions of effluent used for the fathead minnow test with D. pulex. The D. pulex were from a laboratory culture maintained at the Newtown Fish Toxicology Station. Estimated  $LC_{50}$  and  $EC_{50}$  values and 95% confidence limits were reported where possible and are shown in Table 33. Data were evaluated with Probit Analysis whenever possible, with Moving Average Angle where Probit was not applicable.

TABLE 33. ACUTE TOXICITY DATA FOR FATHEAD MINNOW AND DAPHNIA PULEX

	percent	Pathead minnow, effluent concentration		baphnia pulex, percent effluent concentration		
Plant code	96-Hr LCso	95% Confidence limit	Statistical analysis	48-Hr EC50	95% Confidence limit	Statistical analysis
A B	19.0 NATC	15.5 to 24.0	MA	9.0 NAT	6.8 to 11.6	Pp
B C D E P	46.5 NAT	37.6 to 57.5	MA	41.0 NAT	32.4 to 50.2	MA
E P	NAT NAT			7.8 81.7	5.6 to 9.8 66.7 to 101.6	P P
G H J K	64.7 d NÃT	57.0 to 74.8	AM	62.4 _e	54.3 to 72.7	MA
	NAT			nāt Nat		
L M	23.5 NAT	18.3 to 28.7	MA.	28.0 60.0	40.7 to 89.0	MA MA
P g	49.8 NAT	38.8 to 61.8	P	nāt		
N P R S T U	16.5 NAT 46.5	12.4 to 21.7 37.6 to 57.5	P MA	8.0 Ngah	6.1 to 8.0	MA
Ů V	NAT			NAT 12.1	8.7 to 16.3	P
W X	36.0 55.2 NAT	27.4 to 43.9 45.2 to 70.7	MA MA	9.4 6.3 NAT	7.1 to 12.2 3.7 to 8.4	P P
Y Z	NAT NAT			NAT 42.6	30.8 to 64.1	P

amoving average angle. Probit. CNO acute toxicity. A bad batch of fish nullified this test.

e40% Dead at 100% concentration. f100% Dead at all dilutions. gsample inadvertently collected prior to the settling pond.

hNo statistical analysis; heavy solids concentration bbscured the analysis; the sample did not appear to be acutely toxic.

Generally, the toxicity of textile mill effluent samples was exerted within the first 24 hr for both the fathead minnow and D. pulex. The relative sensitivity of the animals cannot be differentiated on the basis of these samples. In 5 out of the 15 samples, D. pulex appeared to be somewhat more sensitive than the fathead minnow. In one instance, the effluent was not acutely toxic to D. pulex but was acutely toxic to fathead minnow, with an estimated  $LC_{50}$  value of 46.5% effluent dilution. It should be noted that 8 samples were acutely toxic to fathead minnow and 10 were acutely toxic to D. pulex. Thus, it is desirable to use, when feasible, more than 1 organism to evaluate the toxicity of an effluent.

Reactions of test organisms to each effluent sample are briefly described in Appendix D.

MARINE ECOLOGY TOXICITY

## Bioassay with Unicellular Marine Algae

Unicellular algae are important constituents of marine ecosystems. They are comprised of a variety of species that have different growth rates, photosynthetic rates, nutrient requirements, and other functions that regulate species composition and diversity in the community in relation to environmental parameters. The algal community, through photosynthesis, produces most of the food and oxygen used in the marine ecosystem.

It is well known that algal species and communities are sensitive to environmental changes. Species may be either inhibited or stimulated by pollutants. In a community, a pollutant may affect some species but not others, thereby causing changes in species diversity and composition. This can be followed by changes in composition of the animal community and altered routes of flow of energy and materials. Often, the altered ecosystem is undesirable from the human standpoint. On this basis, a bioassay program designed to study effects of suspected pollutants should include research on unicellular algae.

Marine algae tests were performed on 15 textile effluent samples at ERL (Sabine Island, Gulf Breeze, Florida) under the direction of Dr. J. Walsh. Fifteen wastewater samples were subjected to this testing series instead of 23 samples because this bioassay was integrated into the program after sampling began. EPA test procedures (3) used for this analysis were modified as follows:

- Continuous lighting was used instead of a 14-hr-light, 10-hr-dark cycle.
- Salinity was 30 parts per thousand (ppt) instead of 10 ppt.
- · Wastes were not sterilized.

- Optical density of cultures was determined on days 3, 4, and 5.
- · Final biomass was not estimated.
- Only the 96-hr EC<sub>50</sub> is reported.

The relationship between optical density as a measure of population density and cell counts was determined by spot checks. In all cases, it was found that optical density and cell counts using a hemocytometer were closely correlated.

In order to estimate stimulation of growth, the ratio of population density in treated samples to population density in controls (T/C) was calculated, and the highest value for each waste is reported.

Effects of the textile wastes on population density of Skeletonema costatum are given in Table 34. There was a wide distribution of toxicity. Wastes L and N were by far the most toxic, whereas B, S, U, and X were not toxic.

TABLE 34. RESULTS OF MARINE ALGAE ACUTE TOXICITY TESTS

		Stimulat	ion
		Percent growth	Percent
	Inhibition, EC <sub>50</sub> ,	over control,	secondary
Plant	percent secondary effluent	T/C	effluent <sup>d</sup>
_	' <b>b</b>	222	100
В	<b>-</b> "	200	100
С	90	130	80
E	10 to 50 <sup>c</sup>	0	0
${f F}$	85	230	10
G	59	130	10
K	77	180	10 to 30
L	1.7	110	0.5
N	2.3	120	0.1 to 0.5
P	9.0	110	0.1
S	<b>_</b> b	410	90
T	70,	310	35
U	_b	180	70
V	94	120	10 to 70
W	50	230	30
Х	b	170	95

<sup>&</sup>lt;sup>a</sup>Percent of secondary effluent solution corresponding to highest stimulation growth rate.

b Inhibition less than 50% in 100% secondary effluent.

<sup>&</sup>lt;sup>C</sup>Small volume of sample received, not enough to complete tests.

Most wastes stimulated growth. Although there were not enough data for proper statistical analysis, wastes with T/C ratios of 1.1 to 1.3 were not significantly different from the controls.

The importance of growth stimulation by waste must not be underestimated. Of the 15 samples, 8 definitely stimulated growth. Five wastes caused population densities two to four times those of the controls.

Another way to look at this is to estimate the concentrations that doubled the population density (T/C equals 2):

Plant	Percent waste
${f T}$	<10
S	10
F	30
W	30
В	100

Fewer than 10% of the wastes from Plants U, X, and K causes an increase of 50% in population density (T/C equals 1.5).

Growth stimulation could have a substantial impact on natural bodies of water by causing 1) eutrophication and/or 2) changes in relative numbers of important phytoplankton.

### Marine Animal Series

Marine animal bioassay testing was performed to ascertain the concentration of secondary wastewater sample that was acutely toxic to juvenile sheepshead minnows and to grass shrimp. Although none of the textile plants discharge directly into a marine environment, this biotest was performed to provide general information about the toxicity of textile plant wastewaters and to evaluate this new bioassay testing procedure. Since this testing series was integrated into the program after sampling began, only 14 textile plant wastewater samples were subjected to this test.

The EPA static bioassay procedure incorporates several methods (5, 12) and is the simplest, most economic marine animal assay test available. Juvenile sheepshead minnows (Cyprinodon variegatus) and adult grass shrimp (Palaemonetes pugio or P. vulgaris) were used as test animals because they adapt easily to a wide range of salinity and temperature in static bioassays.

<sup>(12)</sup> Sprague, J. B. The ABC's of Pollutant Bioassay Using Fish. In: Biological Methods for the Assessment of Water Quality, J. Cairns, Jr., and K. L. Dickson, eds. ASTM Special Technical Publication 528, American Society for Testing and Materials, Philadelphia, Pennsylvania, 1973. pp. 6-30.

Additionally, various phases of their life cycles may be studied due to their short life span.

There are two objections to this procedure: Receiving water characteristics are not closely simulated, and the test organism may not be representative of the most sensitive species in a given geographical area. However, the method is adequate for ranking industries according to their effluent toxicity.

Tests on 14 textile plant wastewater samples were performed at EG&G-Bionomics Marine Research Laboratory (BMRL; Pensacola, Florida) under the direction of Dr. R. Parrish. Dr. J. Walsh was the EPA Technical Advisor associated with the marine ecology biotests.

A total of 0.02  $\rm m^3$  of secondary effluent sample was collected from each of the 14 plants for sheepshead minnow and grass shrimp acute toxicity analysis. Samples were shipped via air freight in glass bottles, packed in ice, and were stored at BMRL in a room with the temperature controlled at 15  $\pm$  1°C until testing. A description of the samples as they arrived at BMRL is given in Table 35.

TABLE 35. PHYSICAL DESCRIPTION OF EFFLUENT SAMPLES AS THEY ARRIVED AT BMRL FOR SHEEPSHEAD MINNOW AND GRASS SHRIMP ACUTE TOXICITY ANALYSIS

Plant		Description
A	Chlorinated; pH particulate.	6.2; cloudy, gray with considerable amount of fine
В	Nonchlorinated;	pH 7.6; clear, light yellow.
С	Nonchlorinated; particulate.	pH 10.2; clear, blue black with moderate amount of
E	Nonchlorinated;	pH 6.8; clear with small amount of particulate.
F	Nonchlorinated;	pH 7.5; clear, light salmon.
G	Chlorinated; pH	6.0; light olive with fair amount of particulate.
K	Chlorinated; pH	7.8; clear, light gray with small amount of particulate
L	Nonchlorinated;	pH 7.3. <sup>a</sup>
N	Nonchlorinated; particulate.	pH 3.7; clear, light gray with moderate amount of
s	Nonchlorinated; particulate.	pH 7.7; clear, light champagne with small amount of
T	Nonchlorinated; particulate.	pH 7.4; clear, blue green with moderate amount of
υ	Chlorinated; pH particulate.	9.0; clear, dark amber with moderate amount of
W	Nonchlorinated; particulate.	pH 8.0; cloudy, orange with moderate amount of
x	Chlorinated; pH particulate.	7.1; clear, light gold with moderate amount of

a Incomplete description because subsample was lost and no other material remained.

Grass shrimp, 15 mm to 26 mm rostrum-telson length, were collected from Big Lagoon, near BMRL. Shrimp were held in flowing water and acclimated to a salinity of 10 ppt for a minimum of 2 days before testing. Mortality was less than 3% during the acclimation.

Methods for the 96-hr, static tests followed EPA test procedures (3) as modified by the EPA Technical Advisor (ERL--Gulf Breeze). Sheepshead minnow and grass shrimp tests were conducted in  $0.004\text{-m}^3$  uncovered glass jars which contained  $0.003~\text{m}^3$  of test solution. Five fish or shrimp were tested per jar, and all test concentrations and controls were duplicated, except as noted. There was no aeration; temperature was maintained at  $20~\pm~1^\circ\text{C}$  during the tests.

Test concentrations were prepared by mixing appropriate volumes of effluent and dilution water directly in test containers. Dilution water was glass-distilled water adjusted to 10 ppt salinity with Rila Marine Mix (Rila Products, Teaneck, NJ). Batches of dilution water were aged for at least 24 hr with aeration and then filtered (5  $\mu m$ ) before mixing test concentrations. Salinity of the effluents was also adjusted to 10 ppt with Rila Marine Mix before preparing the test concentrations. Control jars received 0.003 m³ of 10-ppt dilution water, but no effluent.

Range-finding tests (48-hr) were conducted with all effluents to determine appropriate test concentrations for 96-hr definitive tests. Range-finding tests in which no mortality had occurred after 48 hr of exposure to 100% effluent were extended to 96 hr of exposure. If mortality remained less than or equal to 50%, no further tests were conducted with the effluent. In range-finding tests, only five fish or shrimp were tested, and test concentrations were not duplicated.

In cases where mortality was greater than 50% during rangefinding tests, 96-hr definitive tests were conducted with five fish or shrimp per jar, and all test concentrations and controls were duplicated.

Based on the test results, 24-hr, 48-hr, and 96-hr  $LC_{50}$ 's were graphically interpolated. Two points, representing death in concentrations that were lethal to more than one-half and less than one-half of the fish at the specified times, were plotted on semilogarithmic coordinate paper (test concentrations on the logarithmic axis and corresponding percentages of dead fish on the arithmetic axis, Figure 22). The concentration at which a straight line drawn between the two points crossed the 50% mortality line was the estimated  $LC_{50}$ .

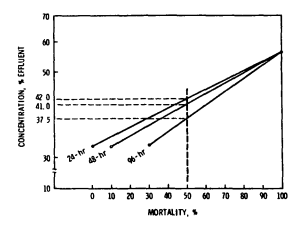


Figure 22. Example of graphically interpolated 24-hr, 48-hr, 96-hr, LC<sub>50</sub>'s for juvenile sheepshead minnow (Cyprinodon variegatus) exposed to textile effluent W.

Sheepshead Minnow Results--

Five of the 14 textile effluents caused greater than 50% mortality of minnows after 96-hr of exposure to effluent concentrations less than or equal to 100% in static, artificial seawater (Table 36). The most toxic effluent (lowest  $LC_{50}$  value) was from Plant W, with a graphically interpolated 96-hr  $LC_{50}$  of 37.5% (solution containing 37.5% effluent sample). Plant N was less toxic, with a 96-hr  $LC_{50}$  of 47.5%. The 96-hr  $LC_{50}$ 's for effluents from Plants A, C, and T were 62.0%, 69.5%, and 68.0%, respectively. Four effluents, from Plants E, G, L, and S, caused less than 50% mortality when sheepshead minnows were exposed to 100% effluent for 96-hr. Therefore, no  $LC_{50}$ 's were determined for these effluents and values are reported as no acute toxicity. Five effluents, from Plants B, F, K, U, and X, caused no deaths when minnows were exposed to 100% effluent for 96-hr, and 96-hr  $LC_{50}$ 's are reported as no acute toxicity effluent.

TABLE 36. ACUTE TOXICITY OF 14 TEXTILE EFFLUENTS TO JUVENILE SHEEPSHEAD MINNOWS (C. VARIEGATUS)

D1	24-Hr	40 11-	lary effluent
Plant	24-HT	48-Hr	96-Hr
A	75.0	69.5	62.0
В			
С		72.0	69.5
E			
F			
G			
K			
L			
N	75.0	75.0	47.5
S			
T			68.0
Ü			
W	42.0	41.0	37.5
Х			

Note. -- Blanks indicate no acute toxicity.

Measured concentrations of dissolved oxygen (DO) in most tests remained greater than 40% of saturation after 96 hr of testing. However, DO was low in test concentrations of Plant T effluent (particularly in 100% effluent) early in the test. Apparently, low DO concentrations in this effluent were due more to the nature of the effluent than to the oxygen demand of test animals.

### Grass Shrimp Results--

Five of the 14 textile effluents caused more than 50% mortality of grass shrimp after 96 hr of exposure to effluent concentrations less than or equal to 100% in static, artificial seawater (Table 37). The most toxic effluent was from Plant C, with a graphically interpolated 96-hr  $LC_{50}$  of 12.8%. Plant W and A effluents were less toxic, with 96-hr  $LC_{50}$ 's of 19.6% and 21.2%; and the 96-hr  $LC_{50}$ 's for effluents from Plants N and T were 26.3% and 34.5%, respectively. Three effluents, from Plants E, G, and S, caused less than 50% mortality when grass shrimp were exposed to 100% effluent for 96 hr. Therefore, no  $LC_{50}$ 's were determined, and values are reported as no acute toxicity. Six effluents, from Plants B, F, K, L, U, and X, caused no deaths when grass shrimp were exposed to 100% effluent for 96 hr, and 96-hr  $LC_{50}$ 's are reported as no acute toxicity.

TABLE 37. ACUTE TOXICITY OF 14 TEXTILE EFFLUENTS TO GRASS SHRIMP (P. pugio)

	LC <sub>50</sub> , percen	nt second	dary effluent
Plant	24-Hr	48-Hr	96-Hr
A	37.6	24.0	21.2
В			
С	17 <b>.7</b>	15.5	12.8
E			
F			
G			
K			
L			
N	43.5	37.4	26.3
S			
${f T}$			34.5
U			
W	37 <b>.5</b>	24.8	19.6
X	•		

Note. -- Blanks indicate no acute toxicity.

Measured concentrations of dissolved oxygen (DO) in most tests remained greater than 40% of saturation after 96 hr of testing. However, DO was low in test concentrations of Plant T effluent (particularly in 100% effluent) early in the test. Apparently, low DO concentrations in this effluent were due more to the nature of the effluent than to the oxygen demand of test animals.

#### RANGE-FINDING ACUTE TOXICITY 14-DAY RAT TEST

The major objective of any biological testing procedure is the identification of toxicological problems at minimal cost. Therefore, a two-step approach was used to evalute the acute in vivo toxicity of samples containing unknown compounds. The first approach is based on the quantal (all-or-none) response; the second is based on the quantitative (graded) response. Normally, the quantal test is used to determine whether or not the quantitative assay is necessary.

#### The Quantal Test

Five male and five female young adult rats (weighing approximately 250 g each) were purchased from the supplier and conditioned at the laboratory for a minumum of 5 days. A single  $10^{-5}$ m<sup>3</sup>/kg dose of undiluted sample was administered by gavage to each animal. Immediately following administration of the test substance and at frequent intervals during the first day, observations were recorded on all toxic signs or pharmacological effects as described in Table 38 (3). The frequency and severity of the signs were scored. Particular attention was paid to time of onset and disappearance of signs. Daily observations were made on all animals through a 14-day observation period. Effluent samples which produced harmful effects in vivo and did not result in deaths were further investigated. At termination of the observation period, all surviving animals were killed and necropsies were performed. Similarly, necropsies were performed on all animals that died during the course of the study.

If mortality did not occur in the quantal study, no further work was done on the test substance, and the  $LD_{50}$  was reported as greater than 10 g/kg.

## The Quantitative Assay

If a single animal in the quantal study died in the 14-day observation period, a quantitative study was performed. Eighty animals equally divided by sex were maintained for 7 days in quarantine to determine good health in the study population. From these, 40 animals then were randomly divided into 4 groups of 5 male and 5 female animals per group. The test substance, treated as in the quantal test, was administered in graded dosages according to the following schedule: 3.0, 1.0, 0.3, and 0.1 g/kg. Dosage was related to the numbers of animals that died and to the severities and types of signs. Observations,

TABLE 38. PHYSICAL EXAMINATIONS IN ACUTE TOXICITY TESTS IN RODENTS (3)

Organ system	Observation and examination	Common signs of toxicity
Central nervous system and somatomotor	Behavior	Change in attitude to observer, unusual vocalization, restlessness, sedation.
	Movements	Twitch, tremor, ataxia, catatonia, paralysis, convulsion, forced movements.
	Reactivity to various stimuli	<pre>Irritability, passivity, anaesthesia,   hyperanaesthesia.</pre>
	Cerebral and spinal reflexes	Sluggishness, absence.
	Muscle tone	Rigidity, flaccidity.
Autonomic nervous system	Pupil size	Myosis, mydriasis.
	Secretion	Salivation, lacrimation.
Respiratory	Nostrils	Discharge.
	Character and rate of breathing	Bradypnoea, dyspnoea, Cheyne-Stokes breathing Kussmaul breathing.
Cardiovascular	Palpataion of cardiac region	Thrill, bradycardia, arrhythmia, stronger or weaker beat.
Gastrointestinal	Events	Diarrhea, constipation.
	Abdominal shape	Flatulence, contraction.
	Feces consistency and color	Unformed, black or clay colored.
Genitourinary	Vulva, mammary glands	Swelling.
	Penis	Prolapse.
	Perineal region	Soiled.
Skin and fur	Color, turgor, integrity	Reddening, flaccid skinfold, eruptions, piloerection.
Mucous membranes	Conjunctiva, mouth	Discharge, congestion, hemorrhage cyanosis, jaundice.
Eye	Eyeball	Exophthalmus, nystagmus.
	Transparency	Opacities.
Others	Rectal or pay skin temperature	Subnormal, increased.
	Injection site	Swelling.
	General condition	Abnormal posture, emaciation.

duration of study, and necropsies were carried out as indicated above. The  $LD_{5\,0}$  was calculated by the method described in Reference 3.

The range-finding tests were conducted at Litton Bionetics under the direction of Dr. R. Beliles. Dr. J. Stara served as the EPA Technical Advisor.

Actual experimental design parameters used in this study were as follows. Young adult rats of the Charles River CD strain (CRL: COBS CD (SD) BR) were obtained from the Charles River Breeding Laboratories, Inc., Portage, Michigan. Animals were individually housed in wire bottom cages in temperature-controlled quarters under artificial illumination controlled to provide a 12-hr light cycle. Water and Purina Laboratory Chow were provided ad libitum with the exception of the night before treatment when food was removed from cages.

Effluent samples were kept refrigerated until used. A single undiluted dose of  $10^{-5}$  m<sup>3</sup>/kg of test material was administered by gastric intubation to five rats of each sex. If any rats died at this dose, an LD<sub>50</sub> value was to be determined by giving additional doses of the test material.

The rats were observed frequently on the day of treatment and daily thereafter. Animals were weighed on the day of treatment, and on days 7 and 14 following treatment. All surviving animals were killed 14 days after treatment and necropsies were performed.

In summary, no rat deaths occurred following the oral administration of  $10^{-5}$  m³ of effluent sample per kilogram of rat body weight for any of the 23 textile plant effluent samples. Residual effects of treatment were not evident from necropsy findings. Twelve samples (from Plants A, C, E, G, K, M, T, U, V, W, X, and Z) showed no effects that could be related to treatment. Reduced activity was observed immediately after dosing with some of the effluent samples (those from Plants D, H, J, P, and Y). Rats treated with sample from Plant R showed signs of eye irritation which were related to treatment. Effluent samples were collected between the aeration lagoon and the settling pond at Plant R. Reduced body weights or weight gains were noted after administration of five samples (those from Plants B, F, L, N, and S).

A more detailed description of the reactions of the test rats to the effluent samples is given in Appendix E.

## SOIL MICROCOSM TEST

Decomposition of dead organic matter by microorganisms is essential for maintaining ecological balance. It is becoming increasingly apparent that as more and more toxic materials from anthropogenic sources are introduced into the soil, they will

progressively inhibit these organisms, ultimately creating a critical imbalance.

Soil decomposition is a complex matrix of biological, physical, and chemical processes, dependent on a myriad of variables. Any test designed to detect and ultimately predict the effects of toxic materials on soils must control as many variables as practicable. In addition, disposal of toxic test substances must be simple, convenient, and practical. To meet these needs, a soil/litter microcosm test contained in a "life support" system was designed, constructed, and used to determine the soil response to toxic materials (3).

A soil microcosm is, by design, a miniature model of the natural system—in this case, the site of decomposition in the upper (50 mm) soil layers. In the test microcosm, soil and litter are separately homogenized and layered in an array of airtight containers in which carbon dioxide generation is monitored with time. Carbon dioxide generation is an accepted measure of soil respiration and, in this case, is assumed to be a measure of the microbial activity in the microcosm during the decomposition of organic matter.

Carbon dioxide generation rates were measured from replicates of each waste solution after 2 wk to 3 wk of incubation. For data analysis, rates were compared with three sets of waste solutions and controls using a linear analysis of variance. Treatments showing statistically significant differences in the stimulation of respiration indicated inhibition or stimulation of respiration.

The soil microcosm test was performed at EPA--Corvallis Environ-mental Research Laboratory, Corvallis, Oregon under the direction of Dr. B. Lighthart. The soil microcosm test used 0.001 m³ of the 0.01-m³ freshwater algae samples. The bioassay procedure and apparatus were those described by the EPA (3). A brief description of the materials and methods used is given in Appendix F.

Instead of reporting all data on carbon dioxide ( $CO_2$ ) generation rates, only the mean qualities of  $CO_2$  produced over the 3-wk incubation period for replicate samples were reported and are presented in Table 39. Samples were analyzed in three batches with one set of replicate controls (using organic-free water) run per batch. The quantity of  $CO_2$  produced each day over the test period was plotted on graph paper and normalized with the control sample. The slope of the resulting curve is the normaalized relative  $CO_2$  rate change (Table 39). If the wastewater sample stimulated the microorganisms, the slope of the curve was positive; if the sample was inhibitory or toxic to the microorganisms, the slope was negative. Of the 23 secondary effluent wastewater samples tested, 15 inhibited the production of  $CO_2$  and 8 stimulated the production of  $CO_2$ .

TABLE 39. RESULTS OF SOIL MICROCOSM TESTS ON SECONDARY WASTEWATER SAMPLES

		Mean total	Normalized	······································
	Plant	CO2 produced,	relative CO <sub>2a</sub>	, k
Run	code	m <sup>3</sup>	rate change	F-value <sup>b</sup>
1	D	$221.9 \times 10^{-6}$	-0.099	789
•	H	$220.2 \times 10^{-6}$	-0.083	242
t	J	$211.6 \times 10^{-6}$	-0.163	442
•	M	$229.9 \times 10^{-6}$	-0.059	224
		$238.7 \times 10^{-6}$	0.022	11.1
ı	Р R	253.3 x 10 <sup>-6</sup>	-0.062	10.9
	Ÿ	$^{204.3} \times 10^{-6}$	-0.172	611
	ż	$222.2 \times 10^{-6}$	-0.112	465
,	Control	246.8 x 10 <sup>-6</sup>	******	
2	E	$275.9 \times 10^{-6}$	-0.048	245
2	Ğ	$302.0 \times 10^{-6}$	0.017	13.5
	ĸ	291.9 x 10 <sup>-6</sup>	-0.004	33.6
	r v	$285.5 \times 10^{-6}$	-0.020	312
	S.	$288.4 \times 10^{-6}$	-0.017	12.4
	v	$275.3 \times 10^{-6}$	-0.066	247
	W	$305.5 \times 10^{-6}$	0.031	12.4
	Control	298.7 x 10 <sup>-6</sup>	0.031	72.4
_		c	0.022	0.14
3	A	246.7 x 10 <sup>-6</sup>	-0.032	55.4
	В	$270.4 \times 10^{-6}$	0.020	
	<u>c</u>	$285.7 \times 10^{-6}$	-0.005	119
	F	236.8 x 10 <sup>-6</sup>	-0.039	6.15
	N	$269.7 \times 10^{-6}$	0.059	25.2
	T	$276.6 \times 10^{-6}$	0.020	62.4
	υ	$277.3 \times 10^{-6}$	0.055	55.0
	X	$274.6 \times 10^{-6}$	0.047	59.5
	Control	$254.2 \times 10^{-6}$		
			•	

Note. -- Blanks indicate information not applicable.

The final column in Table 39 as reported by the EPA Technical Advisor is a measure of the statistical significance of the data. A standard "Student t" test was employed using the F-table and all the individual CO<sub>2</sub> generation rate data. Based on those data, if the value of F is greater than 5.99, then the probability of a Type I error is 5%. If the value of F is greater than 13.7, then the probability of a Type I error is 1%. A Type I error means that one rejects the hypothesis when it is in fact true.

From the data in Table 39, all secondary effluent samples have a statistically significant effect at the 5% confidence level. Data for Plant A have a greater than 5% probability of a Type I error. Of the 23 samples, 17 have a statistically significant effect at the 1% confidence level.

<sup>&</sup>lt;sup>a</sup> Negative sign indicates inhibition of  ${\rm CO}_2$  generation rate compared to a control sample; positive sign indicates  ${\rm CO}_2$  stimulation.

bResults are significant at a 95% confidence level for F > 5.99 and at a 99% confidence level for F > 13.7.

<sup>&</sup>lt;sup>C</sup>Samples inadvertently collected prior to the settling pond.

#### SECTION 8

### DISCUSSION OF RESULTS

### PLANT RANKING BY RELATIVE WASTEWATER TOXICITY

The primary objective of the Phase I screening study was to rank textile plants according to the toxicity of their secondary wastewater and to select plants for detailed toxicity evaluation in Phase II. To accomplish this objective, members of the EPA Bioassay Subcommittee met to evaluate the bioassay data. Members of the Subcommittee are illustrated as EPA Technical Advisors in Figure 20 (Section 7). A summary of all the bioassay results is given in Table 5 (Section 3).

Data evaluation began with ranking of the plants in each set of bioassays. Results are discussed in the following sections.

# Freshwater Ecology Series

Results from these tests showed sufficient variation to permit relative ranking of the toxicity of effluent samples. A composite ranking based on the responses of fathead minnows and Daphnia is shown in Table 40. No general rule can be made concerning the relative response between fathead minnows and Daphnia. For example, Plant E's effluent was significantly toxic to Daphnia but not toxic at all to fathead minnows; at Plant T, the reverse was true.

TABLE 40. RELATIVE TOXICITY RANKING BY BIOASSAY TEST

Relative toxicity	Cytotoxicity (RAM)	Composite freshwater ecology (fathead minnow and Daphn(a)	Composite marine ecology (sheepshead minnows, grass shrimp, and marine algae)	Clonal toxicity (CHO-K1)
Most toxic	C,N	W	L,N,T	D
		A,L		
		C,V		
Intermediate		W		
Toxicity	A,P,L,T,W,X	£	A,C,B,W	M,H,J
		U		
		T		
		G, Z		
Least toxic	м	P, H, M	B,E,P,G,K,U,X	R,P,Y,2
Nontoxic	B,E,G;K,U,D, H,J,P,R,V, Y,E	B, K, S, X, D, J, P, Y		
Not analyz <b>eć</b>	5	·	D,H,J,M,P,R,V,Y,E	A,B,C,E,F G,R,L,N S,T,U,V W,X

At the time of the evaluation, freshwater algae results were not available, but as seen in Table 32 (Section 7), ranking of plants by algal inhibition is similar to that for fathead minnow and Daphnia. Because the focus of the program is toxicity removal by treatment technologies and not ecological impacts, algal stimulation effects were not considered in the ranking.

## Marine Ecology Series

Based on toxicity data for sheepshead minnows, grass shrimp, and marine algae, ranking of effluents by toxicity was accomplished and is shown in Table 40. In all samples, grass shrimp were more sensitive than sheepshead minnows. Also, the fathead minnows were more sensitive in the majority of the samples than sheepshead minnows. No general correlation was seen between the response of Daphnia and grass shrimp.

## Cytotoxicity

Rabbit alveolar macrophage tests indicated that none of the samples was highly toxic. Two samples, N and C, were moderately toxic and the following seven samples were slightly toxic: L, W, T, X, A, F, and J.

Only eight samples were tested by MRC using the clonal toxicity test: D, H, J, M, P, R, Y, and Z. Of the eight samples, four showed significant toxicity: D, M, H, and J.

# Mutagenicity

None of the 23 effluent samples produced a positive response in any of the bacterial tester strains. The Bioassay Subcommittee expressed concern that the detection limits for this bioassay series were not sensitive enough to detect the presence of significant concentrations (0.001 to 0.1  $g/m^3$ ) of chemical mutagens.

# Rat Acute Toxicity Tests

No acute toxicity was observed from the maximum dose  $(10^{-5} \text{ m}^3/\text{kg})$  of rat body weight) ingested by the rats. However, six effluent samples showed potential body weight effects: F, N, C, L, S, and B. The subcommittee expressed concern about the detection limits of this test also.

### Plant Ranking

Based on all of the above analyses, the subcommittee ranked the 23 textile plants in descending order of secondary effluent toxicity, and results are shown in Table 41.

TABLE 41. PRIORITIZATION OF TEXTILE PLANTS BY TOXICITY OF SECONDARY EFFLUENT

Toxicity ranking	Plant
Most toxic	N,A W C,T V,L
Least toxic	S,P
Nontoxic	B,D,E,F,G,H,J,K M,U,X,Y,Z

From the above list, the subcommittee recommended that the following nine textile plants be tested to determine the removal of toxicity achieved by the tertiary treatment technologies being tested in the ATMI/EPA Grant Study: N, A, L, T, C, P, S, W, and V. (Plant R was also recommended for study under Phase II because its secondary effluent samples were inadvertently collected prior to the settling pond.) In addition, they recommended that the freshwater ecology series be used to measure reduction in wastewater toxicity by the treatment technologies. The marine ecology series was not selected because none of the textile plants discharge wastewater into a marine environment.

#### PROGRAM OUTLINE FOR PHASE II STUDY

The objective of the second part of the textile wastewater toxicity study is to determine reduction in priority pollutant concentrations and in acute toxicity as a result of applying the ATMI/EPA BATEA tertiary treatment technologies to the secondary effluent at the 10 textile plants.

Pilot plants are scheduled to be at each (10) textile plant for from 6 wk to 8 wk. For the first 4 wk, seven tertiary treatment systems will be tested to determine which one provides the best removal of criteria pollutants. A treatment system consists of one or more of the six tertiary treatment technologies. From the data collected, the "best" system will be identified. This system will then be set up and operated at steady-state conditions for a final period of 2 wk.

For toxicity and priority pollutant analyses at each plant, 24-hr composited samples will be collected during the 2 wk of steady-state operations from the one system identified as the "best available technology." Since the tertiary treatment system will be composed of several of the six treatment technologies, samples will be collected before and after each unit operation in the system, resulting in approximately four samples.

In order to evaluate the reduction in toxicity and priority pollutant concentrations, 24-hr composited secondary effluent samples will also be collected. Due to hydraulic retention time through the pilot plant, secondary effluent sampling will lead the tertiary treatment sampling by the appropriate time for the tertiary treatment system selected.

A 24-hr composited sample of the intake water to the textile plant will be collected at each of the 10 plants to understand the presence of priority pollutants in wastewater samples. Either continuous or grab samples will be collected depending upon the sampling conditions around the intake water facilities. Samples will be collected for volatile organics, nonvolatile organics, and metals analyses. Therefore, a total of approximately 6 samples will be collected at each of the 10 plants as illustrated in Table 42.

TABLE 42. SAMPLE SCHEDULE AT EACH OF THE 10 TEXTILE PLANTS

Sample site	No. of samples	Analyze for
Plant intake water	1	129 priority pollutants
Secondary effluent	ı	<pre>129 priority pollutants   and freshwater ecology   series</pre>
Best tertiary treatment system	4	<pre>129 priority pollutants   and freshwater ecology   series</pre>

The freshwater ecology series consists of bioassay tests on the following three test organisms: fathead minnows, Daphnia, and freshwater algae. Five sample fractions are collected for priority pollutant analysis: volatile organics, nonvolatile organics, metals, cyanide, and phenol. Criteria pollutant analyses will not be performed since these analyses will be routinely performed under the ATMI/EPA Grant Study.

#### REFERENCES

- 1. Draft Final Report: Sampling and Analysis Procedures for Screening of Industrial Effluents for Priority Pollutants. U.S. Environmental Protection Agency, Cincinnati, Ohio, March 1977. 145 pp.
- Hamersma, J. W., S. L. Reynolds, and R. F. Maddalone. IERL-RTP Procedures Manual: Level 1 Environmental Assessment. EPA-600/2-76-160a (PB 257 850), U.S. Environmental Protection Agency, Research Triangle Park, North Carolina, June 1976. 147 pp.
- 3. Duke, K. M., M. E. Davis, and A. J. Dennis. IERL-RTP Procedures Manual: Level 1 Environmental Asssessment Biological Tests for Pilot Studies. EPA-600/7-77-043 (PB 268 484), U.S. Environmental Protection Agency, Research Triangle Park, North Carolina, April 1977. 114 pp.
- 4. Manual of Methods for Chemical Analysis of Water and Wastes. EPA-625/6-76-003a (PB 259 973), U.S. Environmental Protection Agency, Cincinnati, Ohio, 1976. 317 pp.
- 5. Standard Methods for the Examination of Water and Wastewater, Fourteenth Edition. American Public Health Association, Washington, D.C., 1976. 874 pp.
- 6. McCann, J., E. Choi, E. Yamasaki, and B. N. Ames. Detection of Carcinogens as Mutagens in the Salmonella/Microsome Test: Assay of 300 Chemicals. Proceedings of the National Academy of Science, 72:5135-5139, 1975.
- 7. Poole, D. C. and V. F. Simmon. Final Report of in Vitro Microbiological Studies of Twenty-two Wastewater Effluent Samples. Contract 68-01-2458, U.S. Environmental Protection Agency, Biomedical Research Branch, Research Triangle Park, North Carolina, November 1977. 111 pp.
- 8. Waters, M. D., D. E. Gardner, C. Aranyi, and D. L. Coffin. Metal Toxicity of Rabbit Alveolar Macrophages in Vitro. Environmental Research, 9(1):32-47, 1975.

- 9. Campbell, J. A., H. F. Stack, and P. R. Williams. Cytotoxicity Screening of Twenty-three Textile Mill Effleunt Water Samples Utilizing the Rabbit Alveolar Macrophage Assay. Contract 68-02-2566, U.S. Environmental Protection Agency, Biomedical Research Branch, Research Triangle Park, North Carolina, December 1977. 86 pp.
- 10. Malcolm, A. R., B. H. Pringle, and H. W. Fisher. Chemical Toxicity Studies with Cultured Mammalian Cells. In: Bioassay Techniques and Environmental Chemistry, G. E. Glass, ed. Ann Arbor Science Publishers, Inc., Ann Arbor, Michigan, 1974. pp. 217-230.
- 11. Shiroyama, T., W. E. Miller, and J. C. Greene. The Efforts of Nitrogen and Phosphorus on the Growth of Selenastrum Capriocornutum Printz. EPA-606/3-75-034, U.S. Environmental Protection Agency, Corvallis, Oregon, March 1975. pp. 132-142.
- 12. Sprague, J. B. The ABC's of Pollutant Bioassay Using Fish. In: Biological Methods for the Assessment of Water Quality, J. Cairns, Jr. and K. L. Dickson, eds. ASTM Special Technical Publication 528, American Society for Testing and Materials, Philadelphia, Pennsylvania, 1973. pp. 6-30.
- 13. Standard for Metric Practice. ANSI/ASTM Designation: E 380-76<sup>c</sup>, IEEE Std 268-1976, American Society for Testing and Materials, Philadelphia, Pennsylvania, February 1976. 37 pp.

#### APPENDIX A

### RECOMMENDED LIST OF PRIORITY POLLUTANTS

## TABLE A-1. RECOMMENDED LIST OF PRIORITY POLLUTANTS

# Compound name Acenaphthene Acrolein Acrylonitrile Benzene Benzidine Carbon tetrachloride (tetrachloromethane) Chlorinated benzenes (other than dichlorobenzenes) Chlorobenzene 1,2,4-Trichlorobenzene Hexachlorobenzene Chlorinated ethanes (including 1,2-dichloroethane, 1,1,1-trichloroethane and hexachloroethane) 1,2-Dichloroethane 1,1,1-Trichloroethane Hexachloroethane 1,1-Dichloroethane 1,1,2-Trichloroethane 1,1,2,2-Tetrachloroethane Chloroethane Chloroalkyl ethers (chloromethyl, chloroethyl and mixed ethers) Bis(chloromethyl) ether Bis(2-chloroethyl) ether 2-Chloroethyl vinyl ether (mixed)

(continued)

Chlorinated naphthalene

2-Chloronaphthalene

## Compound name

```
Chlorinated phenols (other than those listed elsewhere; includes trichlorophenols and chlorinated cresols)
```

```
2,4,6-Trichlorophenol
p-Chloro-m-cresol (4-chloro-3-methylphenol)
```

Chloroform (trichloromethane)

2-Chlorophenol

Dichlorobenzenes

- 1,2-Dichlorobenzene
- 1,3-Dichlorobenzene
- 1,4-Dichlorobenzene

Dichlorobenzidine

3,3'-Dichlorobenzidine

Dichloroethylenes (1,1-dichloroethylene and 1,2-dichloroethylene)

- 1,1-Dichloroethylene (vinylidine chloride)
- 1,2-Trans-dichloroethylene
- 2,4-Dichlorophenol

Dichloropropane and dichloropropene

- 1,2-Dichloropropane
- 1,3-Dichloropropylene (cis and trans-1,3-dichloropropene)
- 2,4-Dimethylphenol

Dinitrotoluene

- 2,4-Dinitrotoluene
- 2,6-Dinitrotoluene
- 1,2-Diphenylhydrazine

Ethylbenzene

Fluoranthene

(continued)

## TABLE A-1 (continued).

## Compound name

```
Haloethers (other than those listed elsewhere)
```

4-Chlorophenyl phenyl ether

4-Bromophenyl phenyl ether

Bis(2-chloroisopropyl) ether

Bis (2-chloroethoxy) methane

Halomethanes (other than those listed elsewhere)

Methylene chloride (dichloromethane)

Methyl chloride (chloromethane)

Methyl bromide (bromomethane)

Bromoform (tribromomethane)

Dichlorobromomethane

Trichlorofluoromethane

Dichlorodifluoromethane

Chlorodibromomethane

#### Hexachlorobutadiene

Hexachlorocyclopentadiene

Isophorone (3,5,5-trimethyl-2-cyclohexen-1-one)

Naphthalene

Nitrobenzene

Nitrophenols (including 2,4-dinitrophenol and dinitrocresol)

2-Nitrophenol

4-Nitrophenol

2,4-Dinitrophenol

4,6-Dinitro-o-cresol

#### Nitrosoamines

N-nitrosodimethylamine

N-nitrosodiphenylamine

N-nitroso-di-n-propylamine

Penta chlorophenol

Phenol

(continued)

## Compound name

```
Phthalate esters
  Bis(2-ethylhexyl) phthalate
  Butyl benzyl phthalate
  Di-n-butyl phthalate
  Diethyl phthalate
  Dimethyl phthalate
  Di-n-octyl phthalate
Polynuclear aromatic hydrocarbons
  Benz(a)anthracene (1,2-benzanthracene)
  Benzo(a) pyrene (3,4-benzopyrene)
  3,4-Benzofluoranthene
  Benzo(k) fluoranthene
   (11,12-benzofluoranthene)
  Chrysene
  Acenaphthylene
  Anthracene
  Benzo(ghi)perylene (1,12-benzoperylene)
  Fluorene
  Phenanthrene
  Dibenz (ah) anthracene
    (1,2,5,6-dibenzanthracene)
  Indeno(1,2,3-cd)pyrene
    (2,3-o-phenylenepyrene)
  Pyrene
Tetrachloroethylene
Toluene
Trichloroethylene
Vinyl chloride (chloroethylene)
Pesticides and metabolites
  Aldrin
  Dieldrin
  Chlorodane (technical mixture and metabolites)
DDT and metabolites
  4,4'-DDT
  4,4'-DDE (p,p'-DDX)
  4,4'-DDD (p,p'-TDE)
                                     (continued)
```

## TABLE A-1 (continued).

### Compound name

```
Endosulfan and metabolites
```

α-Endosulfan

β-Endosulfan

Endosulfan sulfate

#### Endrin and metabolites

Endrin

Endrin aldehyde

# Heptachlor and metabolites

Heptachlor

Heptachlor epoxide

# Hexachlorocyclohexane

α-BHC

β-BHC

 $\lambda$ -BHC (lindane)

δ-BHC

# Polychlorinated biphenyls (PCB)

PCB-1242 (Arochlor 1242)

PCB-1254 (Arochlor 1254)

PCB-1221 (Arochlor 1221)

PCB-1232 (Arochlor 1232)

PCB-1248 (Arochlor 1248) PCB-1260 (Arochlor 1260)

PCB-1016 (Arochlor 1016)

# Toxaphene

# Elements

Antimony (Total)

Arsenic (Total)

Asbestos (Fibrous)

Beryllium (Total)

Cadmium (Total)

Chromium (Total)

Copper (Total)

Cyanide (Total)

Lead (Total)

(continued)

# TABLE A-1 (continued).

# Compound name

# Elements (continued)

Mercury (Total) Nickel (Total) Selenium (Total) Silver (Total) Thallium (Total) Zinc (Total)

2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD)

## APPENDIX B

## PRIORITY POLLUTANT ANALYSIS FRACTIONS

## TABLE B-1. VOLATILE COMPOUNDS

Compound	Compound
Chloromethane Dichlorodifluoromethane Bromomethane Vinyl chloride Chloroethane Methylene chloride Trichlorofluoromethane 1,1,-Dichloroethylene 1,1-Dichloroethane trans-1,2,-dichloroethane Chloroform 1,2-Dichloroethane 1,1,1-Trichloroethane	1,2-Dichloropropane trans-1,3-dichloropropene Trichloroethylene Dibromochloromethane Cis-1,3-dichloropropene 1,1,2-Trichloroethane Benzene 2-Chloroethyl vinyl ether Bromoform 1,1,2,2-Tetrachloroethylene 1,1,2,2-Tetrachloroethane Toluene Chlorobenzene
Carbon tetrachloride Bromodichloromethane Bis(chloromethyl) ether	Ethylbenzene Acrolein Acrylonitrile

TABLE B-2. BASE NEUTRAL EXTRACTABLE COMPOUNDS

Compound	Compound
1,3-Dichlorobenzene 1,4-Dichlorobenzene Hexachloroethane 1,2-Dichlorobenzene Bis (2-chloroisopropyl) ether Hexachlorobutadiene 1,2,4-Trichlorobenzene Naphthalene Bis (2-chloroethyl) ether Hexachlorocyclopentadiene Nitrobenzene Bis (2-chloroethoxy) methane 2-Chloronaphthalene Acenaphthylene Acenaphthene Isophorone Fluorene 2,6-Dinitrotoluene 1,2-Diphenylhydrazine 2,4-Dinitrotoluene N-nitrosodiphenylamine Hexachlorobenzene 4-Bromophenyl phenyl ether Phenanthrene	Anthracene Diethyl phthalate Dimethyl phthalate Fluoranthene Pyrene Di-n-butyl phthalate Benzidine Butyl benzyl phthalate Chrysene Bis(2-ethylhexyl) phthalate Benz(a) anthracene Benzo(b) fluoranthene Benzo(k) fluoranthene Benzo(a) pyrene Indeno(1,2,3-cd) pyrene Dibenz(a,h) anthracene Benzo(g,h,i) perylene N-nitrosodimethylamine N-nitrosodimethylamine 4-Chlorophenyl phenyl ether 3,3'-Dichlorobenzidine 2,3,7,8-Tetrachlorodibenzo- p-dioxina Bis-(chloromethyl) ether

This compound was specifically listed in the consent decree.

Because of TCDD's extreme toxicity, EPA recommends that laboratories not acquire analytical standards for this compound.

TABLE B-3. ACID EXTRACTABLE COMPOUNDS

2-Chlorophenol
Phenol
2,4-Dichlorophenol
2-Nitrophenol
p-Chloro-m-cresol
2,4,6-Trichlorophenol
2,4-Dimethylphenol
2,4-Dinitrophenol
4,6-Dinitro-o-cresol
4-Nitrophenol
Pentachlorophenol

TABLE B-4. PESTICIDES AND PCB

Co	mpound	
β-Endosul	fan	
α-BHC		
γ-BHC		
•		
β-BHC		
Aldrin		
Heptachlo		
Heptachlo		<b>.</b>
α-Endosul	.fan	
Dieldrin		
4,4'-DDE		
4,4'-DDD		
4,4'-DDT		
Endrin		
Endosulfa	n sulfate	3
δ-BHC		
Chlordane	•	
Toxaphene	<b>:</b>	
PCB-1242	(Aroclor	1242)
PCB-1254	(Aroclor	
PCB-1221	(Aroclor	•
PCB-1232	(Aroclor	
PCB-1248	(Aroclor	
PCB-1260	(Aroclor	•
PCB-1016	(Aroclor	
	, · · · · · ·	

TABLE B-5. METALS AND OTHER COMPOUNDS

Metals,	
total	Others
Antimony	Asbestos
Antimony	
Arsenic	Cyanide
Beryllium .	
Cadmium	
Chromium	
Copper	
Lead	
Mercury	
Nickel	
Selenium	
Silver	
Thallium	
Zinc	

## APPENDIX C

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#### APPENDIX D

# REACTION OF FATHEAD MINNOWS AND DAPHNIA TO TEXTILE SECONDARY EFFLUENTS

## Plant A

This sample was acutely toxic to both the fathead minnow and D. pulex, resulting in estimated 96-hr  $LC_{50}$  and 48-hr  $EC_{50}$  values of 19.0% and 9.0% effluent dilution, respectively. The 2-day delay from collection of the sample to the beginning of the test probably did not affect sample toxicity.

Control fish ranged in length 30 mm to 39 mm, with the average being 34.1 mm. Their weight ranged from 0.2 g to 0.6 g, with the average being 0.37 g.

## Plant B

This sample was not acutely toxic to either the fathead minnow or D. pulex.

Control fish ranged in length from 25 mm to 51 mm, with the average being 36.5 mm. Their weight ranged from 0.2 g to 1.25 g, with the average being 0.56 g.

#### Plant C

This sample was acutely toxic to both fathead minnow and D. pulex, resulting in estimated 96-hr LC<sub>50</sub> and 48-hr EC<sub>50</sub> values of 46.5% and 41.0% effluent dilution, respectively. All fish died in 100% effluent, and half of the fish died in 60% effluent dilution within the first 1/2 hr. The remaining fish in the 60% dilution were dead by the next morning.

Control fish ranged in length from 28 mm to 38 mm, with the average being 33.5 mm. Their weight ranged from 0.2 g to 0.5 g, with the average being 0.30 g.

## Plant D

At the end of the test, fish from the control were measured and weighed. Fish ranged in length from 32 mm to 48 mm, with an average 38.6 mm. Fish ranged in weight from 0.15 g to 0.73 g, with an average weight of 0.34 g. At the end of the test, two

fish had died in the 60% dilution and one in the 4.7% dilution. In the D. pulex test, no animals died during the 48-hr period.

Results indicate that this sample was not acutely toxic to fathead minnows or D. pulex.

## Plant E

At the end of the test, fathead minnows from the control jar were measured and weighed. Fish ranged in length from 21 mm to 37 mm, with an average length of 31 mm. Their weight ranged from 0.1 g to 0.4 g, with the average being 0.29 g.

Data indicate that this textile mill sample was not acutely toxic to fathead minnows over a 96-hr period. However, the sample was acutely toxic to the  $D.\ pulex$ , with an estimated 48-hr EC<sub>50</sub> of 7.8 % waste.

## Plant F

This sample evidenced no acute 96-hr toxicity to fathead minnows. However, the estimated 48-hr  $EC_{50}$  value of 81.7% indicated the sample was somewhat toxic to D. pulex.

Fish ranged in length from 28 mm to 38 mm, with the average being 32.1 mm. Their weight ranged from 0.2 g to 0.5 g, with the average being 0.28 g.

## Plant G

This sample indicated acute toxicity to both fathead minnow and D. pulex with estimated 96-hr  $LC_{50}$  and 48-hr  $EC_{50}$  values of 64.7% and 62.4% effluent dilution, respectively. In this instance, toxicity was essentially the same for both species.

Fish from the control jar ranged in length from 26 mm to 35 mm, with an average length of 29.6 mm. Their weight ranged from 0.15 g to 0.40 g, with the average being 0.26 g.

## Plant H

The fathead minnow bioassay is inconclusive for this sample. It appears that the fish were diseased. There was considerable mortality in all of the jars including the control. This was not a reliable test run.

The bioassay run using the D. pulex, however, was meaningful. Mortality in the 100% effluent was 40%. In the other concentrations, 10 D. pulex were alive at the end of the test. The 100% concentration of effluent showed some indication of toxicity, although an EC<sub>50</sub> value could not be determined.

## Plant J

Fish from the control jar were weighed and measured at the end of the test. Fish ranged in length from 32 mm to 44 mm, with an average of 37 mm. They ranged in weight from 0.3 g to 0.75 g, with an average weight of 0.46 g. At the end of the test, one fish had died in the 100%, one in the 60%, two in the 7.8%, and three in the 13% dilutions. Three of the D. pulex died in 100% effluent; one died in the 60% dilution; and one died in the control jar.

Based on the results from these test, this sample was not acutely toxic to the fathead minnows or the D. pulex.

## Plant K

This sample indicated no acute toxicity to fathead minnows and little toxicity to D. pulex. Four D. pulex died in 100% effluent during the 48-hr period of the test.

Control fish ranged in length from 28 mm to 35 mm, with an average length of 30 mm. Their weight ranged from 0.15 g to 0.4 g, with the average being 0.26 g.

## Plant L

This sample was quite acutely toxic to both fathead minnows and  $D.\ pulex$ , resulting in estimated 96-hr LC<sub>50</sub> and 48-hr EC<sub>50</sub> values of 23.5% and 28.0% effluent dilution, respectively. The two statistical procedures used could not be utilized to calculate the toxicity values with 95% confidence limits for the  $D.\ pulex$ .

Control fish ranged in length from 26 mm to 40 mm, with the average being 30.6 mm. Their weight ranged from 0.2 g to 0.6 g, with the average being 0.33 g.

## Plant M

At the end of the test, fathead minnows in the control jar weighed an average 0.29 g, with a range of 0.15 g to 0.50 g. Average length of the fish was 34 mm, with a range of 28 mm to 48 mm.

Data indicate that this sample was not acutely toxic to the fathead minnow, but it was acutely toxic to D. pulex. A 48-hr  $EC_{50}$  value of 60.0% effluent, with 95% confidence limits of 40.72% to 88.95%, was determined with the moving average angle procedure.

## Plant N

This sample was acutely toxic to both fathead minnow and D. pulex. The estimated 96-hr LC50 value for the fish was 48.8% effluent dilution. All fish in 100% effluent were dead within 19 hr after the beginning of the test. The fish showed evidence of hemorrhaging aroung the mouth and tail. The pH of the 100% effluent was 4.0 at the beginning of the test and only 4.5 24 hr later. Thus, low pH was responsible for this mortality. The sample was extremely toxic to D. pulex, with all animals being killed within 24 hr in as low as 13% effluent dilution, and all were dead in every dilution at the end of 48 hr. The 2-day delay from the time of collection to the beginning of the test probably did not affect the toxicity. The temperature of the sample was 2°C when it reached the Newtown Fish Toxicology Station. Biological degradation would be minimal under these conditions.

Control fish ranged in length from 28 mm to 45 mm, with the average being 36.2 mm. Their weight ranged from 0.2 g to 1.0 g, with the average being 0.55 g.

 $_{\rm H}$ 

## Plant P

At the end of the test (96 hr), there were nine fish surviving in each test jar. None of the fish in the control jar died. The sample did not indicate 96-hr acute toxicity.

At the end of the test there wer 10 Daphnia alive in each container.

Results indicate that this sample was not acutely toxic to either fathead minnows or D. pulex.

#### Plant R

Based on appearance, this sample, was both high in suspended solids and highly turbid (149 units). Turbidity of previous samples ranged from a low of 6 units to a high of 54 units. Considerable sample agitation achieved only 2.8 g/m $^3$  dissolved oxygen in the 100% effluent at the beginning of the test.

At the end of the test, fish from the control jar averaged 0.28 g in weight, with a range of 0.15 g to 0.65 g. Average length was 35.4 mm, with a range of 28 mm to 47 mm.

Mortality in both the fathead minnow bioassay test and the D. pulex test usually was observed within the first 24 hr. BOD appears to have been a contributing factor. Dissolved oxygen dropped to low levels in the four high effluent volumes. However, after 24 hr, dissolved oxygen was still at a level (0.3  $g/m^3$  to 1.8  $g/m^3$ ) wherein fathead minnows can survive for a

considerable period of time. In 18 hr there was a complete kill of the fish in 100% and 60% effluent. In 36% and 21.6% effluent there was 80% and 30% mortality, respectively, in 24 hr.

Probit analysis of the data at the end of the fathead bioassay (96 hr) indicates an  $LC_{50}$  value of 16.45% effluent, with 95% confidence limits of 12.39% to 21.72%. A 48-hr  $LC_{50}$  value for D. pulex could not be determined with probit analysis. The moving average angle procedure indicated the 48-hr  $EC_{50}$  value to be 7.96% effluent, with 95% confidence limits of 6.14% to 7.96%. Thus, the data show this particular textile mill effluent sample to be acutely toxic to both fathead minnows and D. pulex. The D. pulex were more sensitive than the fathead minnows.

## Plant S

This sample exhibited no 96-hr acute toxicity to fathead minnows and little toxicity to D. pulex. There were three, two, and one D. pulex deaths in 100%, 60%, and 36% effluent dilution, respectively.

Unlike all other samples, this sample contained a heavy, flocculent, fibrous material that settled to the bottom of the test containers (to a depth approximately 5.1 cm from the 100% effluent container). Fish hid in this material throughout the test and were not adversely affected.

Control fish ranged in length from 27 mm to 34 mm, with the average being 29.6 mm. Their weight ranged from 0.20 g to 0.35 g, with the average being 0.25 g.

## Plant T

This sample was acutely toxic to fathead minnow, with an estimated 96-hr  $LC_{50}$  value of 46.5% effluent dilution. All of the fish were dead in 100% effluent within 18 hr after the test was started. Slight evidence of toxicity to D. pulex was indicated with three animals dying in 100% effluent and one in the 60% dilution. The 2-day delay from the time of collection to the beginning of the test probably did not affect the toxicity of the sample.

Control fish ranged in length from 26 mm to 38 mm, with the average being 31.3 mm. Their weight ranged from 0.2 g to 0.5 g, with the average being 0.33 g.

## Plant U

This sample was not acutely toxic in 96 hr to fathead minnow. However, it was toxic to D. pulex, having an estimated 48-hr  $EC_{50}$  value of 12.1% effluent dilution. The 2-day delay from

sample collection to the beginning of the test probably did not affect its toxicity.

The length of the control fish ranged from 29 mm to 46 mm, with the average being 37.7 mm. Their weight ranged from 0.2 g to 1.0 g, with the average being 0.51 g.

## Plant V

This sample indicated acute toxicity to both fathead minnows and D. pulex. All fish died in 100% effluent within 1 hr; all died in the 60% dilution within 2.5 hr. The estimated 96-hr LC<sub>50</sub> was 36% effluent dilution. D. pulex were much more sensitive with an estimated EC<sub>50</sub> value of 9.4% effluent.

Fathead minnows in the control jar at the end of the test ranged in length from 27 mm to 38 mm, with an average length of 29 mm. Their weight ranged from 0.1 g to 0.45 g, with an average weight of 0.23 g.

## Plant W

This sample was acutely toxic to fathead minnow and D. pulex, resulting in estimated 96-hr  $LC_{50}$  and 48-hr  $EC_{50}$  values of 55.2% and 6.3% effluent dilution, respectively. The sample was much more toxic to D. pulex than to fathead minnow.

Control fish ranged in length from 28 mm to 41 mm, with the average being 34 mm. Their weight ranged from 0.2 g to 0.7 g with the average being 0.37 g.

#### Plant X

This sample was not acutely toxic to either fathead minnows or  $D.\ pulex$ . In contrast with a 2-day delay for most samples, it should be noted that there was a delay of 2 day from the time of sample collection until the test was begun. It is possible that the toxic components may have decomposed during the extra day of storage, although this seems unlikely when compared to other samples.

Control fish ranged in length from 28 mm to 42 mm, with the average being 35.3 mm. Their weight ranged from 0.2 g to 0.85 g, with the average being 0.48 g.

#### Plant Y

The fathead minnow bioassay was not significant with this sample because a disease caused the loss of 30% of the control fish. Despite the disease problem, there was 70% to 90% survival of the fish throughout the range of effluent dilutions. In the

D. pulex test, only one animal died in 100% effluent and one in the 21.6% dilution.

Based on the results from these tests, this sample was not acutely toxic to fathead minnows or D. pulex.

## Plant Z

At the end of the test, fathead minnows in the control jar averaged 0.5 g in weight with a range of 0.2 g to 1.1 g. Average length of the fish was 35.8 mm, with a range of 28 mm to 50 mm.

Data indicate that this particular sample was not acutely toxic to fathead minnow. However, the sample was acutely toxic to  $D.\ pulex.$  A 48-hr EC<sub>50</sub> value of 42.57% with 95% confidence limits of 30.79% to 64.07% was determined with probit analysis.

#### APPENDIX E

# REACTIONS OF RATS TO TEXTILE PLANT SECONDARY EFFLUENT

## Plant A

Although reduced activity was observed immediately after treatment, there were no deaths. Necropsy showed mottled kidneys; however, this has been observed frequently in untreated rats in the Litton Bionetics laboratory and was not considered treatment related. Mean body weights (grams) are tabulated below.

		Day		
Dose	<u>Sex</u>	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	M	240	312	340
• •	F	240	250	266

Data did not suggest any treatment effect.

## Plant B

There were no deaths following treatment. One male had soft stools during several days of the observation period. Necropsy findings were limited to kidney changes described above. Mean body weights (grams) are tabulated below.

		Day		
Dose	Sex	0 7 14		
$10^{-5} \text{ m}^3/\text{kg}$	M	249 248 300		
, ,	F	202 233 241		

Body weight loss of males between 0 and 7 was not normal.

## Plant C

There were no deaths and no signs of toxicity following treatment. Necropsy findings were limited to kidney changes previously described. Mean body weights (grams) are tabulated below.

			Day	
Dose	<u>Sex</u>	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	М	297	328	365
	F	214	221	223

Data did not suggest any treatment-related effect.

## Plant D

There were no deaths following treatment. Reduced activity immediately after treatment among all male rats and a soft stool in one male rat 1 day after treatment were observed. Necropsy showed mottled kidneys; however, this was not considered treatment related. Mean body weights (grams) are tabulated below.

			Day	
Dose	Sex	0	7	14
$10^{-5} \text{m}^3/\text{kg}$	M	164	235	274
. 3	$\mathbf{F}$	171	208	216

Body weight data did not suggest any adverse effect.

## Plant E

There were no deaths following treatment. Signs of eye irritation appeared in 6 of 10 rats near the end of the 14-day observation period, but were not considered treatment related. Necropsy findings, limited to changes in heart surface and kidneys, were observed only among male rats; they also were not considered treatment related. Mean body weights (grams) are tabulated below.

			Day	
Dose	Sex	0	_7_	14
$10^{-5} \text{ m}^3/\text{kg}$	M	176	225	283
	F	157	175	214

Body weight data did not suggest any treatment effect.

## Plant F

There were no deaths and no signs of toxicity following treatment. Necropsy findings were limited to kidney changes previously described. Mean body weights (grams) are tabulated below.

			Day	
Dose	Sex	0	7_	14
$10^{-5} \text{ m}^3/\text{kg}$	M	187	244	291
_	F	210	235	229

Decreased body weights were observed in four of five females during the last 7 days of the observation period.

## Plant G

There were no deaths following treatment. Signs of eye irritation were observed in a few rats near the end of the observation period. Because of the time elapsing between treatment and the onset of changes, no treatment relationship was judged to be present. Necropsy findings were limited to heart and kidney changes previously described. Mean body weights (grams) are tabulated below.

			Day	ì
Dose	<u>Sex</u>	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	М	171	212	263
	F	166	173	214

Body weight data did not suggest any treatment effect.

## Plant H

Although reduced activity was observed among the females immediately following treatment, there were no deaths. Mean body weights (grams) are tabulated below.

		Day
Dose	<u>Sex</u>	0 7 14
$10^{-5} \text{ m}^3/\text{kg}$	M	198 251 268
, •	F	180 224 229

Body weight data did not suggest any treatment related effect.

#### Plant J

Although reduced activity was observed immediately after treatment, there were no deaths. Mottled kidneys were observed at necropsy, but were not considered treatment related. Mean body weights (grams) are tabulated below.

	l.	Day
Dose	<u>Sex</u>	0 7 14
$10^{-5} \text{ m}^3/\text{kg}$	M	201 275 210
	F	187 221 232

Body weight data did not suggest any adverse effect.

## Plant K

There were no deaths following treatment. Signs were limited to redness around the eyes of one female rat. Necropsy findings were limited to mottled kidneys. Mean body weights (grams) are tabulated below.

	1			Day	
Dose	1	<u>Sex</u>	0	_7_	.14
$10^{-5} \text{ m}^3/\text{kg}$		М	209	277	313
• •	1	F	192	233	248

Body weight data did not suggest any treatment effect.

## Plant L

There were no deaths and no signs of toxicity following treatment. Necropsy findings were limited to kidney changes previously described among the male rats. Mean body weights (grams) are tabulated below.

	1		Day	
Dose	Sex	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	M	335	354	380
	F	238	240	251

Weight gain among female rats was less than normal.

## Plant M

There were no deaths and no signs of toxicity except for soft stool in one male rat on day 11 after treatment. Necropsy findings were limited to mottled kidneys. Mean body weights (grams) are tabulated below.

	] - 	Day
Dose	<u>Sex</u>	0 7 14
$10^{-5} \text{ m}^3/\text{kg}$	M	262 320 332
	F	196 227 233

Data did not suggest any treatment-related effect.

## Plant N

There were no deaths and no signs of toxicity following treatment. Except for kidney changes previously described, Necropsy findings were limited to a distended cecum in one male rat. Mean body weights (grams) are tabulated below.

			Day	
Dose	Sex	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	M	166	267	284
• -	F	193	237	213

Loss of body weight among females between day 7 and 14 was unusual.

## Plant P

There are no deaths following treatment. Soft stools were observed in one male rat following treatment. Necropsy findings were limited to kidney changes previously described. Mean body weights (grams) are tabulated below.

		. Day	
Dose	<u>Sex</u>	0 7 1	4
$10^{-5} \text{ m}^3/\text{kg}$	М	164 239 28	4
	F	170 204 21	2

Body weight data did not suggest any adverse effect.

## Plant R

There were no deaths following treatment. Dark red material was observed around the nose or eyes of male rats the day following treatment. No other signs were observed. Necropsy findings consisted of several instances of mottled kidneys. Mean body weights (grams) are tabulated below.

		Day
Dose	Sex	0 7 14
$10^{-5} \text{ m}^3/\text{kg}$	M	236 292 320
. •	F	196 225 231

Body weight data did not suggest any treatment related effect.

## Plant S

There were no deaths following treatment. Redness around the eye of one male rat seen near the end of the observation period was not judged to be related to the treatment. Necropsy findings were limited to kidney changes previously described. Mean body weights (grams) are tabulated below.

			Day	
Dose	<u>Sex</u>	0	7_	14
$10^{-5} \text{ m}^3/\text{kg}$	M	312	341	364
	F	237	239	248

Weight gain of females in the first seven days after dosing was judged to be below normal.

## Plant T

There were no deaths and no signs of toxicity following treatment. Necropsy showed mottled kidneys in only one rat. Mean body weights (grams) are tabulated below.

			Day	
Dose	. <u>Sex</u>	0	7_	14
$10^{-5} \text{ m}^3/\text{kg}$	M	172	268	289
	F	208	234	249

Data did not suggest any treatment-related effect.

## Plant U

There were no deaths and no signs of toxicity following treatment. Necropsy findings were limited to kidney changes previously described. Mean body weights (grams) are tabulated below.

			Day	
Dose	Sex	0	_7_	14
$10^{-5} \text{ m}^3/\text{kg}$	М	360	403	435
	F	206	231	239

Data did not suggest any treatment-related effect.

## Plant V

There were no deaths following treatment. Signs suggestive of eye irritation occurred in 4 of 10 treated rats. These appeared 4 to 5 days after administration of test material and were not

judged to be related to treatment. Except for mottled kidneys, necropsy signs were limited to rough appearance of heart ventricles. Mean body weights (grams) are tabulated below.

			Day	
Dose	<u>Sex</u>	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	M	174	233	270
	F	158	171	208

Body weight data did not suggest any adverse effect.

## Plant W

There were no deaths and no signs of toxicity following treatment. Necropsy findings were limited to kidney changes previously described among male rats. Mean body weights (grams) are tabulated below.

	•			Day	ŧ
Dose	<u>Sex</u>	•	0	7_	14
$10^{-5} \text{ m}^3/\text{kg}$	M		350	378	394
, ,	F			234	244

Data did not suggest any treatment-related effect.

## Plant X

There were no deaths and no signs of toxicity following treatment. Necropsy findings were limited to kidney changes previously described. Mean body weights (grams) are tabulated below.

		Day		
Dose	<u>Sex</u>	0 7 14		
$10^{-5} \text{ m}^3/\text{kg}$	M	291 359 374		
• •	F	203 235 247		

Data did not suggest any treatment-related effect.

## Plant Y

Although reduced activity was observed immediately following treatment, there were no deaths. One female rat developed an ulceration on the ventral thorax 9 days after treatment, but this was not considered treatment related. Mean body weights (grams) are tabulated below.

		Day		
Dose	Sex	0	_7_	14
$10^{-5} \text{ m}^3/\text{kg}$	M	220	277	318
	F	191	214	233

Body weight data did not suggest any treatment-related effect.

## Plant Z

There were no deaths and no signs of toxicity following treatment. Except for mottled kidney changes, Necropsy findings were limited to an unusual rough appearance of the right ventricle in one male rat. Mean body weights (grams) are tabulated below.

		_ Day		
Dose	Sex	0	7	14
$10^{-5} \text{ m}^3/\text{kg}$	M	249		
	F	198	228	236

Data did not suggest any adverse effect.

#### APPENDIX F

PROTOCOL TO TEST EFFECTS OF WASTE MATERIALS ON MICROBIAL RESPIRATION (CARBON DIOXIDE REDUCTION) IN A SIMPLE SOIL MICROCOSM

## Materials:

- Homogenized soil from site in question
- 12 9.46 x  $10^{-4}$  m<sup>3</sup> Mason jars with airtight lids 12 3 x  $10^{-5}$  m<sup>3</sup> carbon dioxide trap bottles with airtight lids

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- 20°C incubator
- Approximately 0.5 N NaOH solution
- Approximately 0.6 N HCl solution
- 8 bent glass rods
- 10 x 10<sup>-6</sup> m<sup>3</sup> burette or titrometric device
- Trizma (to prepare standard)

#### Methods:

- (A) Soil preparation:
  - · Air dry soil and grind to pass a 1- to 2-mesh screen. Ball milling or crushing may be required to break larger particles.
- (B) Microcosm preparation:
  - Weigh 100 g (air dried) sieved and homogenized soil into each of eight Mason jars. One set of four replicates each for test and control treatments.
  - Optional soil inoculum solution:
    - Thoroughly mix approximately 200 g of fresh nondried soil with 10-3 m3 water.
    - Separate microorganisms from sediment by filtration or light centrifugation.
  - Moisten soil in each Mason jar from 60% to 80% of field water holding capacity (FWHC) by uniformly pipetting dropwise over the surface in each of four replicates,  $25 \times 10^{-6} \text{ m}^3$  of either test or control solution, plus 5 x  $10^{-6}$  m<sup>3</sup> of inoculum and/or water to bring soil to desired FWHC.

#### **GLOSSARY**

- acute toxicity: Toxic effects to an organism due to a shortterm exposure.
- concentration: Amount of sample (or toxicant) by weight or volume per unit volume of test solution.
- criteria pollutants: Pollutant species identified by EPA Effluent Guidelines Division which require effluent standards and include BOD<sub>5</sub> COD, TSS, chrome, phenol, color, sulfide, and pH for the textile industry.
- cytotoxicity: Toxicity to mammalian cells.
- dose: Measured weight or volume of sample (or toxicant) fed to
   test organism.
- EC $_{50}$ : Effective concentration at which 50% of the test organisms reach the desired effect. The "effect", for example, can be growth inhibition or stimulation.
- gastrointubation: Insertion of a tube into the intestinal tract to feed effluent sample to test animal.
- gavage: Forced feeding of an animal through a tube.
- hemocytometer: Microscope slide with square rulings used for counting blood corpuscles or other cells.
- in vitro: Describing a biological reaction which can be performed outside the living organism, such as in a test tube.
- in vivo: Describing a biological reaction which takes place
   within the living organism.
- $L\dot{C}_{50}$ : Lethal concentration fifty calculated concentration of substance which is expected to cause death in 50% of the test organism population, as determined from their exposure to the substance.
- LD<sub>50</sub>: Lethal dose fifty calculated dose of chemical substance which is expected to cause death in 50% of the test organism population, as determined from their exposure to the substance.

- moving average angle: Iterative computer model designed to estimate the median of a tolerance distribution using number of test species used and number that dried due to exposure to the sample.
- necropsy: Sacrificing the test organism to perform an autopsy.
- priority pollutants: The 129 chemical species identified by EPA as a result of the consent decree.
- probit analysis: Iterative computer model designed to calculate  $LC_{50}$  values from dose response tests using dosage values, number of test species in control and those exposed to the effluent sample, and probability values of response.
- raw waste: Untreated wastewater as it leaves the textile plant and enters the wastewater treatment facility.
- secondary effluent: Textile wastewater treated by aerated lagoons and clarified.
- soil microcosm: Miniature model of the natural system; in this case, the site of decomposition in the upper (5 cm) soil layer.
- viability: Capacity of an organism to live and grow.

## CONVERSION FACTORS AND METRIC PREFIXES (13)

## CONVERSION FACTORS

To convert from	to	Multiply by
Grams/meter <sup>3</sup> (g/m <sup>3</sup> ) Kilogram (kg) Meter (m) Meter <sup>3</sup> (m <sup>3</sup> ) Meter <sup>3</sup> (m <sup>3</sup> )	Milligrams/liter Pound-mass (avoirdupois) Inch Gallon (U.S. liquid) Liter	1.0 2.205 3.937 x 10 <sup>1</sup> 2.642 x 10 <sup>2</sup> 1.0 x 10 <sup>3</sup>

## METRIC PREFIXES

Prefix	Symbol	Multiplication factor	Example	_
Kilo Milli Micro	k m	10 <sup>3</sup> 10 <sup>-3</sup> 10 <sup>-6</sup>	5 kg = 5 x $10^3$ grams 5 mg = 5 x $10^{-3}$ gram 5 $\mu$ g = 5 x $10^{-3}$ gram	;

<sup>(13)</sup> Standard for Metric Practice. ANSI/ASTM Designation: E 380-76°, IEEE Std 268-1976, American Society for Testing and Materials, Philadelphia, Pennsylvania, February 1976. 37 pp.