CONCENTRATION TECHNOLOGIES FOR HAZARDOUS AQUEOUS WASTE-TREATMENT

by

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FOREWORD

CONCENTRATION TECHNOLOGIES FOR HAZARDOUS AQUEOUS WASTE TREATMENT

ABSTRACT

This report describes an ongoing program to evaluate and verify several selected concentration techniques for hazardous constitutents of aqueous waste streams. In the first phase of the project, data was obtained regarding the performance of unit processes for concentrating the hazardous constituents. Applications are expected in the treatment of ground and surface waters affected by the disposal of hazardous wastes.

In conjunction with gathering data on the unit processes, data were obtained on the composition of the waste streams to which the processes could be applied.

The second phase involved a stepwise evaluation of the potential applicability of the candidate technologies to the types of wastes identified earlier. Technology profiles describing the pertinent unit processes and current applications were prepared. These technology profiles formed the basis for an initial screening of the applicability of individual technologies to concentration of hazardous constituents of aqueous wastes. At this point, certain technologies were eliminated from further consideration for reasons discussed in the individual technology profiles. Remaining technologies were carried forward for more detailed review. Compounds identified in the waste streams fell into one of thirteen chemical classes: alcohol, aliphatic, amine, aromatic, halocarbon, metal, miscellaneous, PCB, pesticide, phenol, phthalate, or polynuclear aromatic.

The next step in the evaluation process was an extensive literature review which focused on the technologies which survived the initial screening and upon chemical compounds in the classes given above.

Since it was evident that in most cases no single unit process would be sufficient in itself to adequately treat the diverse waste streams in question, five candidate process trains were formulated as being most broadly applicable to the types of waste streams identified. A desktop analysis then was performed to assess the ability of each process train to treat each of three waste streams. Results of these evaluations provide a basis for making an initial judgment on the applicability of a given concentration technology to specific situations in the absence of experimental data. Results also were used to select

ABSTRACT (continued)

and arrange technologies in priority order for experimental study in the ongoing third phase.

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CONTENTS

Forewor	-A						_	_	_		_		_	_	_	_	_	_	_	_				iii
Abstrac	+	•	•		•	•	•	•	•	•		•	•	•	-	•						•		iv
Figures		•	•		•	•	•	•	•		•	•	•	•	_	•	-	•	•					хi
Tables	• • •	•	•	•	•	•	•	•	•		•	•	•	•	-	•		•	•	_				xii
Acknow]	edome	nte	•	• •	•	•	•	•	•	• •	•	•	•	•	•	•	•	•	•	•	-	•	٠.	xiii
ACKITOW	Leagine	1163	•	• •	•	•	•	•	•	• •	•	•	•	•	•	•	•	•	•	•	•	•		
1.	Intro	duc	+ i	on.		_		_	_				_	_	_	_				_	_		_	1
2.	Concl	nai	0n	C11.	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	Ī	
۷.		ast																						
	W	onc	e i	- ~ =	.ea	UD UIID	• ™≏	ch	· no	100	٠,	•	•	•	•	•	•	•	•	•	•	•	•	
3.	Chemi	0110	m	CIC	1	Dii Ni	16 +1	47 41	C11	mm a	у. тт	•	•	•	•	•	•	•	•	•	•	•	•	
3. 4.	Waste	Car		760	ıca Tə	22.	r r	y i c	ou.	iana	.+ y	٠.	Ch	•	•		·		• • + •	•	•	•	•	35
4.	waste	 	re	alii		en:	LLL	TC	ac	TOU	. a.	u	C1	ıaı	. a.	- 4-0	3 L J	L 22 C	26.	LOI	1 •	•	•	35
	7	TOD	те	M 1	УP	es 	· ~	•	•			•	•	•	•	•	•	•	•	•	•	•	•	35 37
	W	ast	.e	Str	ea	m (JOM	po	91	<u> </u>	n.	•	•	•	•	•	•	•	•	•	•	•	•	
_		ast																						
5.	Techn	ото	дy	EV	7a_	uai	510	n	AÞ	pro	acı	1.	•	•	•	•	•	•	•	•	•	•	•	
6.	Techn B	отō	gy	.Pr	or	 7 T 6	es _.	•	• .	• •	•	٠	•	•	•	•	•	•	•	٠	•	•	•	49
	В	101	og	1Ca	11	Tre	eat	me	nt	: •	•	•	•	•	•	•	•	•	•	٠	•	•	•	49
			Pr	oce	ess	De	esc	ŗı	pt	ion	•	•	•	•	•	•	•	•	•	•	•	•	•	
			Pr	oce	ess	A	5DT	10	at	ion	s.	•	•	•	٠	•	•	•	•	•	•	•	•	52
										1.														52
	С	arb	on	Ac	lso	rp	tio	n	•		•	•	•	•	•	•	•	•	•	•	٠	•	•	53
			Pr	oce	SS	De	esc	ri	pt	ion		•	•	•	•	•	•	•	•	•	•	٠	•	53
			Pr	oce	ess	A	pp1	ic	at	ion	s.	•	•	•	•	•	•	•	•	•	•	•	•	54
										ı.														55
	С	ata	1y	sis							•	•	•		•		•	•	•			•	•	55
			Pr	oce	ess	De	esc	ri	pt	ion		٠		•	•				•	•		•	•	55
			Pr	oce	ess	Αį	opl	ic	at	ion	s.					•	•	•	•				•	56
			Pr	oce	ess	Po	ote	nt	ia	1.													•	56
	С	ent	ri	fuc	rat	io	a .																•	57
			Pr	oce	SS	De	esc	ri	pt	ion														57
			Pr	oce	SS	A	1ac	ic	at	ion	s.													57
										1.														
	C	hem	ic.	a 1	P۳	ec:	ini	t.a	+ i	on.	_													
	Ŭ	* * C * * *	Dr	೧೭೯	255	De	-g-c	ri	nt	ion	_	•	•	_	_	_	_				•		_	
										ion														
										1.														
	~	rys																						
	C	- Y S	D~	~~~	.20	D.	26.W	· ~i	• n+	ion	•	•	•	•	•	•	•	•	•	•	•	•	•	
			D~			7.	55C	<u> </u>	- 2	ion	٠.	•	•	•	•	•	•	•	•	•	•	•	•	60
			D~	006	222	D.	ンナン	n+	1 =	1 - U11	٠.	•	•	•	•	•	•	•	•	•	•	•	•	61
			P F"	uce		(ノレビ	11 i.	. <u></u>		•	-		-	•	-	•	•	•	•	•	•	•	U

CONTENTS (continued)

6.	Technology Profiles (cont.)
	Density Separation
	Process Description 6
	Sedimentation 6
	Flotation 6 Process Applications
	Process Applications 6
	Sedimentation 6
	Flotation 6 Process Potential 6
	Process Potential 6
	Sedimentation 6
	Flotation 6
	Dialucic and Floctrodialucic
	Process Description
	Process Applications 6
	Process Potential 6
	Dictilation 6
	Process Description 6
	Process Description
	Process Applications 6
	Francostion 6
	Process Description 6
	Process Description
	Process Applications
	Process Potential
	Process Description
	1100coo bedetaperen
	Granular Media
	Flexible Media
	Process Applications
	Process Potential
	Flocculation
	Process Description
	Process Applications
	Process Potential
	Ion Exchange
	Process Description
	Process Applications
	Process rotelitat
	recorn recorporation to the second se
	Process Description
	Process Applications
	Process Potential
	Reverse Osmosis
	Process Description
	Process Applications
	Process Potential
	Solvent Extraction
	Process Description
	Process Applications
	Process Potential 7

CONTENTS (continued)

6.	Technology Profiles (cont.)			
	Stripping	•	•	77
	Process Description	•		77
	Process Applications	•	•	78
	Process Potential	•	•	78
	Ultrafiltration	•	•	79
	Process Description			79
	Process Applications			80
	Process Potential	٠.		80
7.	Literature Review			81
, .	Description			81
	Literature Summary			83
	Biological Treatment			83
	Alcohols	•	•	83
	Aliphatics	•	•	83
	Amines			84
	Anomatics			84
				84
	Ethers			84
	Halocarbons			85
	Metals			85
	Pesticides			
	Phenols			85
	Phthalates	•	•	85
	Polynuclear Aromatics	•	•	86
	Chemical Coagulation			86
	Membrane Process - Reverse Osmosis			86
	Membrane Process - Ultrafiltration			87
	Stripping			88
	Solvent Extraction	•	•	88
	Sorption Process - Carbon Adsorption	•	•	89
	Alcohols			89
	Aliphatics			90
	Amines			90
	Aromatics			90
	Ethers			91
	Halocarbons			91
	Metals			91
	PCBs			0.1
	Pesticides			92
	Phenols			92
	Phthalates		_	92
	Polynuclear Aromatics	•	•	92
	Sorption Process - Resin Adsorption	•	•	92
	Alcohols	•	•	93
	Aliphatics	•	•	93
	Ariphacius	•	•	93
	Amines	•	•	94
	Aromatics	•	•	
	Halocarbons	•	•	94 94
	DCRe	_	_	94

CONTENTS (continued)

/ •	Tite	rati	ire i	Kevi	ew																		
		Lite	erati	ure S	Sun	mar	У																
			Sor	ption	n F	roc	es	s -	- F	Res	in	1 <i>P</i>	ds	or	:pt	ic	n	((cor	nt.	.)		
			_	Pest																			94
				Phei	nol	.s .																	95
				Phtl	hal	ate	s.		٠	۰													95
				Poly																			95
			Sor	ption																			95
			•	Meta																			95
				PCB:																			96
8.	Proc	ess	Tra:	ins.																			97
				ion (
				mary																			
		Disc		ion																			
				logi																			
				mica.																			
				ption																			
			Memi	brane	I	roc	65	200	•	•	•	٠	•	•	•	•	•	•		•	•	•	106
				ippi																			
		Fort	nulai	tion	9 f	Dr	000	200	. T	· Tra	ni	ne	•	•	•	•	Ť	•	•	•	•	•	107
		r OLI		cess																			
				cess																			
				cess																			
				cess																			
				cess																			
		Der o		ion (
		Eva.	Luat.	TOIL	OL	PIC	Ces	55	T T	. a.ı	.112	•	•	•	•	•	•	•	•	•	•	•	110
Refere																							126
Append:		• •		• •	•	• •	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	120
whheild:	LCES																						
Α.	Enti	+100	= C01	ntact	+=:	ī																	139
В.	C4+~	してしょ	araci	teri	フコナ	ion		• = + =	•	•	•	•	•	•	•	•	۰	•	•	۰	•	•	141
C.				eatal																	•	•	161
L a		LLCa.	T TT 6	=a Lai	レユゴ	ユエレソ	•	•	•	•	•	•		•	•	•	•	•	•	•		•	

FIGURES

Number		Page
1	Schematic of biological/carbon sorption process train	. 109
2	Schematic of carbon sorption/biological process train	. 111
3	Schematic of biophysical process train	. 112
4	Schematic of membrane/biological process train	. 114
5	Schematic of stripping/carbon sorption process train	. 115
6	Waste stream categorization matrix	. 117

TABLES

Number		Page
1	Chemical Treatability Summary	8
2	Contaminant Classification System	39
3	Summary List of Contaminants Reported	40
4	Wastewater Characterization - Site 010	120
5	Wastewater Characterization - Site 026	122
6	Wastewater Characterization - Synthetic Leachate	124
A-1	Entities Contacted	139
B-1	Summary of Reported Water Contamination Problems	142
C-1	Chemical Treatability	163

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SECTION 1

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INTRODUCTION

Indiscriminate past disposal practices - the placement of waste chemicals in nonsecure ponds, lagoons, and landfills - have created serious environmental and public health problems. Indeed, it has become evident that contamination from unsecured industrial waste storage and disposal sites is a widespread problem. Often, this contamination manifests itself in the form of hazardous leachates, and contaminated ground and surface waters. These contaminant streams are diverse in terms of composition and concentration - varying from site to site, from location to location within a site, and often over time at any given location. Some contaminant streams contain a broad spectrum of organic and inorganic constituents, while others have only a few compounds of concern.

Regardless of whether contaminant streams are associated with active or abandoned sites, the need to detoxify/decontaminate these hazardous aqueous wastes sometimes arises. Moreover, since contaminant streams often are relatively dilute, a preprocessing or concentration step prior to detoxification or disposal may be necessary. However, hazardous aqueous waste treatment for this application is not a routine operation. Little information on and/or experience with concentration technology applied to hazardous leachate or contaminated groundwater exists.

This report describes portions of an ongoing project to evaluate and verify several selected concentration techniques for hazardous constituents of aqueous waste streams. The three year project entails literature search/data acquisition, desktop technology evaluations, and experimental investigations to evaluate and adapt appropriate technologies for the applications of interest. Literature search and desktop evaluations have been completed and are reported herein. At the time of this writing, experimental evaluations of selected concentration technologies are underway.

The major thrust of the initial efforts were twofold:

1) to obtain and compile data on the composition of actual contaminant streams which may require or could benefit from treatment by the concentration technologies; and 2) to collect and compile existing data on candidate concentration technologies.

Subsequent efforts involved assessing the ability of various technologies to concentrate hazardous constituents present in aqueous contaminant streams previously identified. This assessment was based upon characteristics of both the technologies and the contaminant streams. As a result of this evaluation/assessment, several process trains judged to have broad applicability were conceptualized for subsequent experimental study.

Succeeding sections of this report discuss the data gathering efforts, stepwise technology evaluations, and process train formulation efforts.

Because of the large quantity of information involved, detailed data on waste stream composition and on the treatability of 505 chemical compounds are contained in the appendices. To provide quick reference on the potential applicability of a technology to a particular compound, a summary table on chemical treatability is contained in the main body of the report.

SECTION 2

CONCLUSIONS

WASTE STREAMS

- The most widespread hazardous waste problem faced by the public sector is contamination from unsecured waste disposal sites - generally in the form of leachates and contaminated ground and surface waters.
- There is no such thing as a "typical" hazardous waste problem each site is unique.
- Wastes encountered are diverse in terms of composition and concentration - varying from site to site and often varying over time at any given site.
- Some waste streams contain a broad spectrum of organic and inorganic compounds, while others have only a few constituents of concern.
- Waste streams identified in this study primarily fell into one of two composition categories: high organiclow inorganic or low organic-high inorganic.
- Twenty-seven problem sites were identified in this study. The number of different problem sites where various contaminant classes were reported is as follows:

	-
Alcohol	2
Aliphatic	4
Amine	2
Aromatic	8
Halocarbon	9
Metal	15
Miscellaneous*	11
PCB	2
Pesticide	7
Phenol	7
Phthalate	2
Polynuclear Aromatic	5

- Actual or threatened legal proceedings almost invariably
- * See Table 2 for definition of this category

restrict the availability of data on the nature of the problem and effectiveness of cleanup operations.

CONCENTRATION TECHNOLOGY

- Only a limited number and range of unit operations have been applied in the treatment of hazardous aqueous wastes, even though concentration technologies have been used for other applications.
- Activated carbon has been used almost exclusively for concentration of organics in the limited number of larger scale hazardous waste treatment operations.
- Concentration technology performance and operating data for industrial process wastes containing a variety of pollutants usually are reported using a surrogate parameter such as TOC or COD. Specific compound removal data are available only for a very limited number of materials.
- Limited specific information is available through vendors because much of their work is considered proprietary and/or confidential.
- Most available data on specific compound removal has been generated in laboratory and pilot scale experimental studies.
- Much of the experimental data on chemical treatability has been generated from pure compound systems. Removal from multicomponent systems may differ substantially.
- High analytical costs associated with specific compound identification will continue to restrict the data base.
- Several concentration processes are promising for treatment of hazardous aqueous wastes. However, for the application of interest, it is unlikely that any single unit process will be sufficient. In most instances, process trains must be utilized.
- Concentration technologies judged to have the greatest broad spectrum potential are chemical precipitation, flocculation, sedimentation, filtration, biological treatment, carbon adsorption, and resin adsorption.
- Reverse osmosis, stripping, and ultrafiltration are believed to have more limited and specialized applicability.

- Ion exchange for removal of inorganic species also may have potential but usually, competing processes such as chemical precipitation are more economical.
- Since hazardous waste contamination problems differ substantially from place-to-place, treatability studies in some form are almost always a prerequisite to selection of an optimum treatment approach.

SECTION 3

CHEMICAL TREATABILITY SUMMARY

An extensive amount of information on the treatability of hundreds of chemical compounds by various concentration technologies was collected. This information has been assembled in Appendix C which is organized primarily by concentration technology with the treatability of individual compounds organized according to chemical compound classification. The following concentration technologies are addressed in Appendix C:

Process	Process Code No.
Biological	I
Coagulation/Precipitation	II
Reverse Osmosis	III
Ultrafiltration	IV
Stripping	V
Solvent Extraction	VII
Carbon Adsorption	IX
Resin Adsorption	X
Miscellaneous Sorbents	XII

The chemical classification system used is described in detail later in this report; the following chemical classes are addressed in Appendix C:

Chemical Classification	Classification Code No.
Alcohol	A
Aliphatic	В
Amine	С
Aromatic	D
Ether	E
Halocarbon	F
Metal	G
PCB	I
Pesticide	J
Phenol	K
Phthalate	L
Polynuclear Aromatic	$\dot{ extbf{M}}$

A total of 505 different chemical compounds are addressed in Appendix C.

To provide a quick reference on the treatability of each of these 505 compounds, a concise summary of information contained in Appendix C has been prepared and is presented in Table 1. Compounds are arranged in Table 1 in alphabetical order according to their chemical classification. Process and chemical classification code numbers are identical to those in Appendix C. For each compound, a summary statement describing its treatability is given with information on treatability by more than one concentration technology provided for the majority of compounds. Many compounds are known by several names. Attempts were made to use preferred or generic names according to The Merck Index. However, in some cases it was necessary to use the names which were used in the reference documents. Users of Table 1 are advised to check for compounds under several potential alphabetic listings.

An example of a typical entry in the table is that for decanol which reads "IX 100% reduction @ 100 μ g/l." This should be interpreted to mean that in the referenced study, carbon adsorption effected complete removal of decanol which was initially present at a concentration of 100 μ g/l.

TABLE 1 CHEMICAL TREATABILITY SUMMARY

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
A. ALCOHOLS	†					,	
Ally Alcohol	IX	22% reduction @ 100 mg/1	35	Ethanol (con't)	i .	7% reduction @ 286 mg/l	1
. →	1	toxic @>350 mg/l	99		IX	10% reduction @1000mg/1	20
(1-Pentanol)	IX	72% reduction @ 1000mg/1	. 35	Ethylbutanol	I	75-100% reduction	56,100,
Borneol	I	90% reduction	81				101
Butanol	I	70-100% reduction	56,81,99	2-Ethylbutanol	IX	86% reduction @1000mg/]	35
			100,101	2-Ethylhexanol	I	75-85% reduction	56
	IX	53-100% reduction @ 0.1 to 1000 mg/l	20,35,72		IX	98% reduction @ 700mg/l	35
	х	100% reduction @ 100µg/l	20	2-Ethyl-1-Hexanol	IX X	100% reduction @100µg/1 100% reduction @100µg/1	
Sec-Butanol	I	98% reduction	81	Furfuryl Alcohol	I	97% reduction	81
Tert-Butanol	I	98% reduction 0 1000mg/1	81,101 35	m-Heptanol	IX	100% reduction @100µg/1	1
1,4-Butanedio	1 1	99% reduction	81	1-Hexanol	I	70-100% reduction	56,100
Cyclohexanol	I	96% reduction 100% reduction @ 100%/1	8 1 20	m-Hexanol	IX		'
	х	100% reduction @ 100μg/l	20	Isobutanol	IX	42% reduction @1000µg/l	35
Decanol	l l	100% reduction @ 100µg/l 100% reduction @ 100µg/l	20 20	Isopropanol	I	70-100% reduction	56,81, 100,101
Dimethylcyclo	_ I	92% reduction	81	•	IX	13% reduction @1000µg/l	35
hexanol				Methanol	I	30-85% reduction	56,65,
1,2-Ethanedio	1 1	depressed performance @ 484 mg/l	103				100,101 103
Ethanal	_	- ·	300 303		III	0-40% reduction @ 1000	18,30
Ethanol	I	70-100% reduction @ up to 1000 mg/l	100,101 103			mg/l - dependent upon membrane	
	III	<20-100% reduction	18,30		IX	4-33% reduction @15-	35,72
		@ 1000 mg/l - dependent upon membrane				1000 mg/l (conti	' .nued)

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CHEMICAL]	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
4-Methylcyclo- hexanol	I	94% reduction	81	B. ALIPHATICS			
Octanol	I IX X	30-75% reduction 100% reduction @ 100µg/1 100% reduction @ 100µg/1		Acetaldehyde Acetic Acid	IX	30-95% reduction 12% reduction @1000mg/1 <20-80% reduction @1000	
Pentanol	IX X	100% reduction @ 100µg/l 100% reduction @ 100µg/l				mg/l - dependent upon membrane	
Pentarythritol	I	No toxic effect	104		IX	24% reduction @1000mg/1	35
Phenyl methyl carbinol	I	85-95% reduction	101	Acetone	I	50-100% reduction	100,102 103
Propanol	IX	100% reduction @ 100µg/l 19% reduction @ 1000mg/l	: ' 1		III	15-100% reduction @1000 mg/l - dependent upon membrane	18,30
	х	100% reduction @ 100µg/1	20		IX	22% reduction @1000mg/l	35
i-Propanol	III	20-100% reduction @ 1000 mg/l - dependent upon	18,30	Acetone Cyanohydrin	IX	30-60% reduction @100- 1000 mg/1	72
D	_	membrane		Acetonitrile	I	Inhibitory @ 500 mg/l	103,106
m-Propanol	I	99% reduction	81	Acetylglycine	I	Readily oxidized @ 500 mg/l	107
				Acrolein	VII IX	Extractable w/xylene 30% reduction @1000mg/l	90 35,90
				Acrylic Acid	I	50-95% reduction	56,100, 101
					IX	64% reduction @1000mg/l	35,90
				Acrylonitrile	I	70-100% reduction	56,90 107
					V	Could be flash evaporated (conti	90 .nued)

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TABLE 1 (continued)

ether Readi @ 100 Readi @ 500 92% r	ly oxidized 0 mg/1 ly degraded	90 106 103 81 35	Caproic Acid Caprolactam Citric Acid Crotonaldehyde	IX X I	90-98% reduction @ 0.1-1000 mg/1 50% reduction @1000mg/1 94% reduction Biodegradable; de- pressed 0 ₂ consumption 90-100% reduction	81 103 56,100
@ 100 Readi @ 500 92% r X 88% r	0 mg/l ly degraded mg/l eduction	103	Citric Acid	ı	94% reduction Biodegradable; de- pressed 0 ₂ consumption	81 103 56,100
Readi @ 500 92% r X 88% r	ly degraded mg/l eduction	81	Citric Acid	I	Biodegradable; de- pressed 0 ₂ consumption	103 56,100
@ 500 92% r X 88% r	mg/l eduction	81			pressed 02 consumption	56,100
X 88% r			Crotonaldehyde	I	90-100% reduction	, ,
Toxic	eduction @ 985 mg/l	25				101
		20		IX	46% reduction@1000mg/1	35
	@ 500 mg/l; also	106,107	Cyclohexanolone	I	92% reduction	81
	ted to be readily lowly oxidized		Cyclohexanone	IX	96% reduction 67% reduction@1000mg/l	81 35
		106,107	Cyclopentanone	I	96% reduction	81
_	- ;		Cystine	I	Completed inhibited 0 ₂ consumption @ 1000 mg/1	103
X 85% r	eduction @ 1000mg/1	35	L-Cystine	I	Slowly oxidized	107
X 96% r	eduction @ 1000mg/1	35	_		@ 1000 mg/1	
Degra	ded very slowly	107	Decanoic Acid	IX	100% reduction @100µg/1	
x 53% r	eduction @ 1000mg/l	35			100% reduction @100µg/1	
50-95	% reduction; rapid-	56,100	Dicyclopentadiene	IX	Found to vaporize	86
-		106,107	Diethylene Glycol		95% reduction	81
100%	reduction @ 100µg/1		Diisobutyl Ketone		100% reduction @300mg/l	
		103	Disopropyl Methylphosphonate	IX	98% reduction @ 2680 μg/l	86
x	Toxic report but s. 85% re 96% re 53% re 60% re 100% :	96% reduction @ 1000mg/l Degraded very slowly 53% reduction @ 1000mg/l 50-95% reduction; rapid- idly oxidized 60% reduction @ 1000mg/l 100% reduction @ 100µg/l	Toxic @ 500 mg/l; also reported to be readily but slowly oxidized 85% reduction @ 1000mg/l 35 96% reduction @ 1000mg/l 35 Degraded very slowly 107 53% reduction @ 1000mg/l 35 50-95% reduction; rapid-idly oxidized 56,100 idly oxidized 56,100 106,107 20,35 100% reduction @ 1000mg/l 20,35	Toxic @ 500 mg/l; also reported to be readily but slowly oxidized 85% reduction @ 1000mg/l 96% reduction @ 1000mg/l 50-95% reduction; rapididly oxidized 60% reduction @ 1000mg/l 100% reduction @ 100mg/l	Toxic @ 500 mg/l; also reported to be readily but slowly oxidized 85% reduction @ 1000mg/l 35 Degraded very slowly 107 53% reduction @ 1000mg/l 35 50-95% reduction; rapididly oxidized 100% reduction @ 1000mg/l 20,35 100% reduction @ 1000mg/l 20 Rapidly oxidized 100% reduction @ 100µg/l 100% reduction @ 100µg/l 20 Rapidly oxidized 103 Disopropyl Methyl- IX	Toxic @ 500 mg/l; also reported to be readily but slowly oxidized 85% reduction @ 1000mg/l 96% reduction @ 1000mg/l Degraded very slowly 53% reduction @ 1000mg/l 50-95% reduction; rapid-idly oxidized 106,107 100% reduction @ 1000mg/l 100% reduction @ 1000mg/l

CHEMICAL	E	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	CESS - TREATABILITY	REF
Dimethyl Sulfoxide	III	63-88% reduction @ 250-1000 mg/1	18	Formamide	I	Slowly oxidized @ 500 mg/l	107
Dipropylene Glycol	IX	16% reduction @ 1000mg/1	35	Formic Acid	I	Rapidly oxidized @ 720 mg/l	107
2,3-Dithia-	I	100% reduction @ 120µg/1	65		IX	24% reduction @1000mg/l	35
butane				Glutamic Acid	1	Readily oxidized	103
Dodecane	X X	100% reduction @ 100μg/l 25% reduction @ 100 μg/l	•	Glycerol	III	20-100% reduction @ 1000 mg/l dependent	18,30
Dulcitol	I	Slightly inhibitory @ 1700 mg/l	109	Glycerine	I	upon membrane Readily oxidized	103
Erucic Acid	I	Oxidized @ 500 mg/l	107	_		@ 720 mg/l	
Ethyl Acetate	I	90-100% reduction	56,100 101	Glycine	I	Rapidly oxidized @ 720 mg/l	103
	IX	50% reduction@ 1000mg/1	1	Heptane	I	90-100% reduction	56,100
Ethyl	I	90-100% reduction	56,100				101,106
Acrylate	IX	78% reduction @ 1015mg/1	101 35	Heptanoic Acid	IX X	10% reduction @100µg/l 50% reduction @100µg/l	20 20
Ethylene Glycol	IX	97% reduction 7% reduction @ 1000mg/l	81 35	Hexadecane	IX X	100% reduction @100µg/l 25% reduction @100µg/l	20 20
2-Ethylhexyl-	I	90-100% reduction	56,100	Hexylene Glycol	IX	61% reduction @1000mg/1	35
acrylate			101	Hydracrylonitrile	1	0-10% reduction	100
Formaldehyde	I	Conflicting data; removable & inhibitory @ 720-		Isobutyl Acetate	ıx	82% reduction @1000mg/l	35
		3000 mg/l		Isophorone	I	93% reduction	81
	III	<20-80% reduction @1000 mg/l dependent upon	18,30		VII	Extractable w/ethyl ether	90
	IX	membrane 9% reduction @ 1000 mg/I	35	Isoprene	IX	86% reduction @ 500- 1000 mg/l (conti	72 .nued)

TABLE 1 (continued)

CHEMICAL	F	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
Isopropyl Acetate	IX	68% reduction @ 1000mg/1	35	Methyl Isoamyl Ketone	IX	85% reduction @ 986 mg/l	35
Lactic Acid	I	Rapidily oxidized @ 720 mg/l	7	Methyl Isobutyl Ketone	IX	85% reduction @1000mg/1	35
Lauric Acid	IX	Slowly oxidized @500mg/l 100% reduction @ 100µg/l	107 20	Methyl Octadeca-	IX X	100% reduction @100µg/1 100% reduction @100µg/1	20 20
L-Malic Acid	X	100% reduction @ 100µg/l Rapidly oxidized @ 500 mg/l	20 107	Methyl Propyl Ketone	IX	70% reduction @1000mg/1	35
DL-Malic Acid	I	Oxidized after 10-16 hr lag period	107	Myristic Acid	IX X	100% reduction @100µg/1 100% reduction @100µg/1	20 20
	Ι	Inhibitory @ 500 mg/l	107	Nitrilotriacetate	I	>90% reduction @500mg/1	111
Methyl Acetate	IX	4-80% reduction @ 1000 mg/l dependent upon membrane 26% reduction @ 1030 mg/l	18,30 35	Octadecane	IХ	after acclimation 100% reduction @100µg/l 25% reduction @100µg/l	20 20
Methyl Butyl Ketone	IX	81% reduction @ 988 mg/1	35	Octanoic Acid	X	50% reduction @ 100 μg/l 90% reduction @ 100 μg/l	20 20
Methyl	IX	100% reduction @ 100 µg/l	20	Oleic Acid	I	Inhibitory	109
Decanoate '	х	100% reduction @ 100 µg/1	1	Oxalic Acid	I	Inhibitory@ 250 mg/l	103
Methyl	IX	100% reduction @ 100 μg/1	1	Pentane	I	Inhibitory @ 500 mg/l	106
Dodecanoate Methyl Ethyl	X VII	100% reduction @100 μg/1 69-88% reduction	20 27	Pentanedinitrile	I	Slowly oxidized or toxic @ 500 mg/l	106,107
Ketone	IX	@ 12,200 mg/l 47% reduction @ 1000 mg/l	35	Pentanitrile	I	Toxic @ 500 mg/l	106
Methyl Hexade	IX	100% reduction @ 100 μg/1		Propanedinitrile	I	Toxic @ 500 mg/l	106
canoate	х	100% reduction @ 100 μg/1	1	Propanenitrile	I	Toxic @ 500 mg/l	106
				B-Propriolactone	I	Inhibitory @ 500 mg/l (conti	108 nued)

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13

TABLE 1 (continued)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
-	IX	28% reduction @ 1000 mg/l	35	Thioglycollic Acid	I	Inhibitory	103
hyde Propionic Acid	IX		20,35	Thiouracil	Ι	Very slowly oxidized @ 500 mg/l	108
	X	33% reduction @ 1000 mg/l 100% reduction @ 100 μg/l	20	Thiourea	I	Inhibitory @ 500 mg/l	109
Propyl Acetate	IX	75% reduction@1000 mg/1	35	Triethylene Glycol	I IX	98% reduction 52% reduction@1000mg/1	81 35
Propylene Glycol	IX	12% reduction@1000 mg/1	35	Urea	I	Inhibitory @ 1200 mg/1	103
Propylene	IX	26% reduction @ 1000 mg/1	35	Urethane	I	Inhibitory	108
Oxide		, , , , , , , , , , , , , , , , , , ,	_	Valeric Acid	IX	80-100% reduction	20,35
Pyruvic Acid	IX X	100% reduction@100 µg/1	20 20		x	@ 0.1-1000 mg/l 50% reduction@100 μg/l	20
Sodium Alkyl Sulfate	I	Readily degraded	112	Vinyl Acetate	IX	64% reduction @1000μg/1	35
Sodium Lauryl Sulfate	I	Rapidly oxidized	112				
Sodium-N-	I	Readily oxidized	112	C. AMINES			
Oleyl-N- Methyl Taurate				Acetanilide	I	94% reduction	81
Sodium a Sulfo	1	Readily oxidized	112	Allylamine	IX	31% reduction@1000mg/1	35
Methyl		noutry onlargou		p-Aminoacetanilide	I	93% reduction	81
Myristate				m-Aminobenzoic	I	98% reduction	81
Tannic Acid	I	Inhibitory	109	Acid	Ì		
Tetradecane	X X	100% reduction@100 μg/l 50% reduction@100 μg/l	:	o-Aminobenzoic Acid	I	98% reduction	81
Tetraethylene Glycol	IX	58% reduction@1000 mg/1	35	p-Aminobenzoic Acid	I	96% reduction	81
, -	1		1	П	I	(conti	nued)

TABLE 1 (continued)

CHEMICAL	:	PROCESS - TREATABILITY	REF	CHEMICAL	PR	OCESS - TREATABILITY	REF
m-Aminotoluen	I	98% reduction	81	Cyclohexylamine	IX	100% reduction @100µg/l	20
o-Aminotoluene	I	98% reduction	81		X	100% reduction @100µg/1	20
p-Aminotoluene	I	98% reduction	81	Dibutylamine	IX X	100% reduction @100µg/l	20 20
Aniline	I	Inconsistant data; 100% reduction & inhibitory	81,92, 108	Di-N-Butylamine	IX	87% reduction @ 1000mg/1	35
	 T T T	reported @ 500 mg/l 3-100% reduction @ 1000	18,30	Diethanolamine	I	97% reduction 28% reduction @ 996mg/l	81 35
		mg/l dependent on	10,50	Diethylenetriamine		29% reduction @ 1000mg/l	35
	IX	membrane 75-100% reduction	20,35	Dihexylamine	IX X	100% reduction@100µg/l	20 20
	x	@ 0.1-1000 mg/l 100% reduction@ 100 µg/l	20	Diisopropanolamine	IX	46% reduction @ 1000mg/1	35
Benzamide	I	Initially inhibitory Slowly degraded @ 500 mg/l	107	Dimethylamine	IX X	100% reduction@100µg/l 100% reduction@100µg/l	20 20
Benzidine	I	Inhibitory @ 500 mg/l; not reduced @ 1.6 µg/l	108,81	2,3-Dimethylani- line	ı	96% reduction	81
Benzylamine	IX	Adsorbed Inhibitory @ 500 mg/l	108	2,5-Dimethylani- line	I	96% reduction	81
Butanamide	ļr	Slowly oxidized @ 500 mg/l	107	3,4-Dimethylani-	I	76% reduction	81
Butylamine	ТX	52-100% reduction	20,35	line			
	×	@ 0.1-1000 mg/1 100% reduction@100 μg/1	20	Dimethylnitros- amine	IX	Not adsorbed	31
m-Chloroani- line	I	97% reduction	81	Di-N-Propylamine	IХ	80% reduction@1000mg/1	35
o-Chloroani- line	I	97% reduction	81	Ethylenediamine	IX	98% reduction 11% reduction@1000mg/1	81 35
p-Chloroani- line	Į.	96% reduction	81	N-Ethylmorpholine	IX	47% reduction@1000mg/1	35

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CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRC	CESS - TREATABILITY	REF
2-Fluorenamine	I	Slowly biodegraded @ 500 mg/1	108	m-Phenylenedia- mine	I	60% reduction	81
Hexylamine	IX X	100% reduction@100 µg/l 100% reduction@100 µg/l	20 20	o-Phenylenedia- mine	I	33% reduction	81
2-Methyl-5- Ethylpyridine	IX	89% reduction @ 1000 mg/1	35	p-Phenylenedia- mine	I	80% reduction	81
N-Methyl Morpholine	IХ	42% reduction @ 1000 mg/1	35	Piperidine	IX X	100% reduction @100μg/l 100% reduction @100μg/l	20 20
Monoethanol-	ТX	7% reduction@ 1012 mg/1	35	Pyridine	IX	53% reduction @1000mg/1	35
amine Monoisopro-	тх	20% reduction@1000 mg/1	35	Pyrrole	X	100% reduction @100µg/1 100% reduction @100µg/1	20 20
panolamine		•		Thiocetamide	I	Inhibitory@100 mg/l	103
Morpholine	X	100% reduction @ 100 µg/l 100% reduction @ 100 µg/l	20 20	Tributylamine	IX X	100% reduction @100µg/1 100% reduction @100µg/1	20 20
B-Napthylamine	įχ	Adsorbed	31	2,4,6-Trichloro-	I	Readily degraded	92,113
o-Nitroaniline	e T	<99.9% reduction @ 18.5 mg/1	58	aniline		@ 500 mg/l	
p-Nitroaniline	≱I.	<pre><99.9% reduction @ 6.7 mg/l</pre>	58	Triethanolamine	IX	33% reduction@1000mg/1	35
Octylamine	X X	100% reduction@100 µg/1 100% reduction@100 µg/1	20 20				
Pentanamide	ı	Slowly oxidized @ 50.0mg/1	107	D. AROMATICS			
p-(Phenylazo) aniline	I	Inhibitory @ 500 mg/l	108	Acetophenone	IX	50-92% reduction @ 0.1-1000 mg/l	20,35
Phenylenedia-	I	Toxic @ 500 mg/l	113		х	100% reduction @100µg/1	20
mine				11		(conti	inued)

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TABLE 1 (continued)

CHEMICAL	I	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	CESS - TREATABILITY	REF
sec-Amylben- zene	Ι	Toxic @ 500 mg/l	113	Benzoic Acid	I IX	95-100% reduction 91-100% reduction	56,81 20,35
tert-Amylben- zene	Ι	Toxic @ 500 mg/l	113		х	@ 0.1-1000 mg/l 100% reduction @100μg/l	20
Benzaldehyde	I	Conflicting data; re-	81,108,	Benzanitrile	1	Inhibitory@500 mg/l	106
_		ported to be toxic	109	3,4-Benzpyrene	I	Inhibitory @ 500 mg/l	106
	IX	also 99% reduction 50-99% reduction	20,35,72	sec-Butylbenzene	I	Toxic @ 500 mg/l	113
		@ 0.1-1000 mg/1		tert-Butylbenzene	I	Toxic @ 500 mg/l	113
:	Х	100% reduction@100 µg/1	i i	Chloranil	I	Inhibitory @ 10 mg/l	101
Benzene	V	90-100% reduction @ up to 500 mg/l 95-99% reduction	56,100 101,114 13,90	Chlorobenzene	III	100% reduction @200mg/l 97-100% reduction	66 90
	l		27 6,21,31, 35,38,72, 90			<pre>@ 360 mg/l Steam strippable 99% reduction w/chloro- form solvent 50-95% reduction</pre>	64,90 90 21,64,
	I	Slowly oxidized @500mg/l	108			@ 1-416 mg/l	90
	х	99% reduction	32	1-Chloro-2- Nitrobenzene	ΙX	Adsorbed	21
luene, Xylene (BTX)		@ 20-30 mg/1		Cumene	X	100% reduction @100µg/l 100% reduction @100µg/l	20 20
Benzenethiol	I	Inhibitory@ 500 mg/l	108	1,2,4,5-Dibenz- pyrene	I	Inhibitory @ 500 mg/l	108
Benzil	IX X	50% reduction@ 100 µg/l 100% reduction@100 µg/l		m-Dichlorobenzene	VII	100% reduction @200mg/1 Air & steam strippable Extractable 95-100% reduction @ 0.1-416 mg/1 (conti	66,92 90 90 20,90

16

TABLE 1 (continued)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
m-Dichloro- benzene(cont)	х	100% reduction @ 100 μg/1	20	2,4-Dichloropro- pionic Acid	I	No reduction @ 186 mg/1	115
o-Dichloro- benzene	I V	100% reduction@200 mg/l Air & steam strippable	66 90	Dimethylaniline (Xylidine)	IX	94% reduction @ 380 μg/1	6
	IX	Extractable 95-100% reduction @ 0.1-1000 mg/1	90 20,90	7,9-Dimethyl- benzacridine	I	Inhibitory @ 500 mg/1	108
p-Dichloro-	x	100% reduction @ 100 µg/1 100% reduction @ 200 mg/1		7,10-Dimethyl- benzacridine	I	Inhibitory @ 500 mg/l	108
benzene	V	Steam strippable Extractable 95-100% reduction	90 90 20,90	Dinitrobenzene	III	7-81% reduction@ 30 mg/l dependent upon membrane	8
	x	@ 0.1-416 mg/l 100% reduction @ 100 μg/l	20	3,5-Dinitro- benzoic Acid	I	50% reduction	81
1,2-Dichloro- benzene	v	70% reduction	64	2,4-Dinitro- phenylhydrazine	III	3-91% reduction @ 30mg/ldependent upon membrane	18
1,3-Dichloro- benzene	V	80% reduction	64	2,4-Dinitro-	I	90-100% reduction @ 0.39-188 mg/l	81,90
	V IX	90% reduction 60% reduction	64 64	toldene	VII	Extractable 95% reduction @ 416 mg/l	90 90
3,3'-Dichloro- benzidine	IX	Adsorbed	31	2,6-Dinitro- toluene	VII	Extractable 95% reduction@416 mg/l	90 90
2,4-Dichloro- phenoxyacetic Acid	I	No reduction @ 174 mg/l	115	Ethylbenzene	I	90-100% reduction @ 0.192-105 mg/l	21,56, 100,101 114
2,6-Dichloro- phenoxyacetic Acid	I	No reduction @ 178 mg/l	115		II V	56% reduction @ 153 mg/lw/alum 80-93% reduction	
MCIU						(conti	90

TABLE 1 (continued)

CHEMICAL]	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
Ethylbenzene (cont)	VII	97% reduction 50-84% reduction @ 1-115 mg/1	27,90 21,35	m-Nitrobenzalde- hyde	I	94% reduction	81
Hexachloro-	I	No reduction @ 200 mg/l	64,90 66,92	o-Nitrobenzalde- hyde	I	97% reduction	81
benzene	III V	52% reduction @ 638 mg/l Steam strippable Extractable	90 64 90	p-Nitrobenzalde- hyde	I	97% reduction	81
Hydroquinone	III	95% reduction @ 416 mg/l 2-80% reduction	90 18,30	Nitrobenzene	I	Reported to be toxic @ 500 mg/1; 96-100% reduction @ 58-530 µg/1	21,58 81,108
	ıx	@ 1000 mg/l 83% reduction@1000 mg/l	35		II	34% reduction @ 160 μg/l w/alum	21
Hydroxyben- zenecarbonit-	I	Toxic @ 500 mg/1	106		v	Steam strippable @ 450-2160 mg/l	64
rile	i				VII	Extractable	90
Isophorone	IX	97% reduction @ 1000 mg/1	35,90		IX		21,35,
2-Methylben-	I	Toxic @ 500 mg/l	106			@ 1-1023 mg/1	90
zenecarbonit- rile	1	10x10 @ 300 mg/1	100	m-Nitrobenzoic Acid	I	93% reduction	81
3-Methylben- zenecarbonit-	I	Toxic @ 500 mg/1	106	o-Nitrobenzoic Acid	I	93% reduction	81
rile 4-Methylben-	I	Toxic @ 500 mg/1	106	p-Nitrobenzoic Acid	I	92% reduction	81
zenecarbonit- rile	_	TOATE & JOU Mg/1	100	Nitrofluorine	I	Slowly oxidized @ 500 mg/l	108
4,4'-Methylene	IX	Adsorbed	31	m-Nitrotoluene	I	98% reduction	81
Bis-(2-Chlo-roaniline)		,		o-Nitrotoluene	I	98% reduction	81
	I	10-30% reduction	100	p-Nitrotoluene	I	98% reduction	81
pyridine	1	'	l	1	1	(conti	nued)

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CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL]	PROCESS - TREATABILITY	REF
Paraldehyde	I	30-50% reduction 74% reduction @ 1000 mg/1	100		V VTT	73-92% reduction 94-96% reduction	13,90 27,90
Pentamethyl- benzene	I	Inhibitory @ 500 mg/l	113		IX	@ 41-44 mg/l 79-98% reduction	6,35
m-Propyl- benzene	I	Very slowly oxidized @ 37.5 mg/1	114	m-Toluidine	I	@ 0.12-317 mg/l 100% reduction	90
Pyridine	IX	47-86% reduction	35,72	Toxaphene	IX	>99% reduction@ 155µg/1	
Sodium Alkyl- benzene	I	@ 500-1000 mg/l Slowly oxidized	112	1,2,3-Trichloro- benzene	V	100% reduction @ 200mg/1 50% reduction Extractable	66 64,90 90
Sulfonate					IX	70-100% reduction @ 0.1-416 mg/l	20,64 90
Styrene	I	70-100% reduction	100,101		х	100% reduction @ 100µg/1	
		98-99% reduction >93% reduction 55-97% reduction	13 27 21,35	1,2,4-Trichloro- benzene	Ι	100% reduction @ 200mg/l	66
		@ 20-200 mg/l	72	1,3,5-Trichloro- benzene	I	100% reduction @ 200mg/1	66,92
Styrene Oxide 1,2,3,4-Tetra- chlorobenzene		95% reduction @ 1000 mg/l 74% reduction @ 200 mg/l	35 66	2,4,6-Trichloro- phenoxyacetic	I	50% reduction @ 53 mg/l	115
1,2,3,5-Tetra- chlorobenzene	Ι	80% reduction @ 200 mg/l	66	2,4,5-Trichloro- phenoxypropionic Acid	I	99% reduction @ 107.5 mg/l	115
l,2,4,5-Tetra- chlorobenzene	Ι	@ 200 mg/l, 80% reduction @ 500 mg/l very slowly oxidized	66,113	2,4,6-Trinitro- toluene (TNT)	IV	80-93% TOC reduction @ 200 mg/l TOC	10
Toluene	I	48-100% reduction @ 8μg/l to 500 mg/l; 500 mg/l	100,101,	1	X	Adsorbed 99% reduction @ 81-116 mg/l	2,40
		was inhibitory	106,108, 114	1		(continu	ued)

TABLE 1 (continued)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL]	PROCESS - TREATABILITY	REF
2,6,6-Tri- nitrotoluene	Ι	50-84% reduction @100mg/1		Diethyl Ether	III	9.5-90% reduction @ 1000 mg/l dependent upon membrane	18
Xylene	VII	92-95 reduction @ 20-200 µg/l >97% reduction	65 27	Diethylene Glycol Monobutyl Ether	IX	83% reduction@1000 mg/l	35
	IX	68-99% reduction @ 0.14-200 mg/l	6,72	Diethylene Glycol Monoethyl Ether	IX	44% reduction @ 1010 mg/l	35
m-Xylene	Ι	Inhibitory @ 500 mg/l	113	Ethoxytriglycol	IX	70% reduction @ 1000 mg/1	35
o-Xylene p-Xylene	I	Inhibitory @ 500 mg/l Inhibitory @ 500 mg/l	113 113	Ethylene Glycol Monobutyl Ether	IX	56% reduction@1000 mg/l	35
				Ethylene Glycol Monethyl Ether	IX	31% reduction @ 1022 mg/1	35
E. ETHERS				Ethylene Glycol Monohexyl Ether Acetate	IX	66% reduction@100 mg/1	35
bis(2-Chloro- isopropy1) Ether	III	47-94% reduction @ 250 mg/l dependent upon membrane	18	Ethylene Glycol Monohexyl Ether	IX	87% reduction @ 975 mg/l	35
bis(Chloro-		100% reduction Extractable	90 90	Ethylene Glycol Monomethyl Ether	IX	14% reduction @ 1024 mg/1	35
ethyl) Ether bis (Chlorois- opropyl)Ether	VII	50% reduction @ 94 µg/l Extractable w/ethyl ether & benzene	90 90	Ethyl Ether	III	<20-100% reduction @ 1000 mg/l dependent upon membrane	30
Butyl Ether	IX	100% reduction @ 197 mg/1	35	Isopropyl Ether	I	~	56,10
Dichloroiso- propyl Ether	IX	100% reduction @ 1008 mg/1	35		IX	80% reduction @ 1023 mg/1	101 35
						(continu	ıed)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRO	CESS - TREATABILITY	REF
F. HALOCARBONS	†		•				
Bromochloro- methane	IX	Adsorbed	21	Chloroethylene	VII	Air strippable Soluble in most	90 90
Bromodichloro- methane	ı	Air & steam strippable Soluble in most organics	` 90 90		ıx	organics Adsorbed	90
	x x	Adsorbed @ 2 mg/l	21,46 46	Chloroform	V	Steam strippable @ 140 mg/l	95
Bromoform	I	100% reduction @ 0.4-1.9 ug/l	65		X	Adsorbed Adsorbed	21,46 32,46
	x x	100% reduction @ 100µg/l 100% reduction @ 100µg/l	20,46 20,46	Chloromethane	V	Air strippable Soluble in most	90 90
Bromomethane		Air strippable Soluble in most organics Adsorbed	90 90 90	Dibromochloro- methane	VII	organics Air & steam strippable Extractable w/organics	90 90
Carbon Tetra- chloride	I	100% reduction@177 µg/l 51% reduction@144 µg/l w/alum	21 21		IX	ether, & alcohols Adsorbed	21,46 90
	х	Adsorbed Adsorbed	6,21,90 32	Dichlorodifluoro-	X	Adsorbed Extractable w/organics,	46 90
Chloral	V	Steam strippable @693mg/l		methane		ethers, & alcohols	30
Chloral Hydrate	VII	49% reduction @15,200mg/1	27	Dichloroethane	IX V	Adsorbed @ 12 µg/l 90% reduction w/air	6,21 90
Chloroethane	v	90% reduction by air	90	ethane		stripping Extractable w/alcohols	90
	VII	<pre>stripping Extractable w/alcohol & aromatics</pre>	90		1	& aromatics Adsorbed	46,90
	IX	Adsorbed	90		х	Adsorbed (conti	46

TABLE 1 (continued)

CHEMICAL	PROCESS - TRE	ATABILITY RE	F CHEMICAL	PROCESS - TREATABILITY	REF
l,2-Dichloro- ethane (also see	I Reduced V Air & steam VII Extractable		,95 propane (cont)	VII Soluble in most organics IX 93% reduction	90
Ethylene Dichloride)	& aromatics IX 81% reduction	on @1000mg/1 90	2-Dichloro-	@ 1000 mg/l V Air & steam strippable	
Dichloroethy-	X Adsorbed VII >99% reducti	.on @1500ppm; 27,	propytene	VII Soluble in most	90
lene	Kerosene & C	10-C12 ef-	95	organics IX Adsorbed	90
l,1-Dichloro- ethylene	V Air & steam VII Extractable	strippable 90, w/alcohols, 90	[]	vII Kerosene and C ₁₀ -C ₁₂ organics effective solvents	95
	aromatics, & IX Adsorbed	ethers 90	Ethylene Chloro- hydrin	VII 21% reduction @ 1640 mg/1	27
l,2-Dichloro- ethylene	IX Adsorbed X Adsorbed	46 46	HEthylene Dichlo-	V 99% reduction @ 8700 mg/l	66,95
l,2-trans-Di- chloroethy- lene	V Air & steam VII Soluble in m organics IX Adsorbed		1,2-Dichloro- ethane)	VII 94-100% reduction @ 23-1804 mg/1 w/kero- sene & C ₁₀ -C ₁₂ organics	95
Dichlorofluo-	IX Adsorbed	90		IX 81% reduction @ 1000 mg/1	35,95
rometnane Dichloro-	V 90% reductio	n w/air 90,	05	X Adsorbed	32
nethane	stripping. S pable @ 800	team strip-	Hexachlorobuta- diene	V Air & steam strippable VII Soluble in most	90 90
	VII Soluble in m organics			organics IX 100% reduction	20
	IX Adsorbed	90		@ 100 μg/1 X 100% reduction	20
l,2-Dichloro- propane	V Air & steam	strippable 90		@ 100 μg/l (conti	m., - 31

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TABLE 1 (continued)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	1	PROCESS - TREATABILITY	REF
Hexachlorocy- clopentadiene	V	Polymerizes w/heat	90	Tetrachloro- ethylene	v	90% reduction by air & steam stripping	90
Hexachloro- ethane		Extractable w/aromatics, alcohols, & ethers 100% reduction @ 100 µg/1	90 20	,	VII X		90 46,90 46
	X	100% reduction @ 100 µg/1	20	Tetrachloro-	v	90% reduction by air	90
Methylene Chloride	Ι	80-88% reduction @ 10-430 µg/1	65	methane	VII	& steam stripping Soluble in most organics	90
Pentachloro- ethane		73% reduction @ 190 µg/l 100% reduction w/kero- sene solvent @ 10 mg/l	6 95	Tribromomethane	V VII V	Air & steam strippable Soluble in most organics Adsorbed	90 90 21,90
Perchloro- ethylene	VII V	Steam strippable @ 15 mg/l Extractable w/kerosene	95 95	Trichloroacetic Acid	III	25-49% reduction @ 250 mg/l dependent upon membrane	18
Propylene Dichloride	IX	& C ₁₀ -C ₁₂ solvents 93% reduction@1000 mg/1	35	Trichloroethane	VII	97-99% reduction w/kerosene & C ₁₀ -C ₁₂ solvents	95
Tetrachloro- ethane		Kerosene & C ₁₀ -C ₁₂ organics provided 95% 100% reduction @ 100 µg/1	95 20	1,1,1-Trichloro- ethane	I V	>90% reduction @ 8-79 µg/l Air & steam strippable	65 90,95
	х	100% reduction @ 100 µg/1	20		VII	Extractable w/alcohols & aromatics	90
1,1,1,2-Tetra- chloroethane	V	Steam strippable @ 513 mg/l	95		X X	Adsorbed Adsorbed @ 551 μg/l	90 46
1,1,2,2-Tetra- chloroethane	VII	Difficult to steam strip Extractable w/aromatics, alcohols, & ethers Adsorbed	95 90	1,1,2-Trichloro- ethane	I	<99% reduction @ 1305 µg/l	58
			90		v	Air & steam strippable	90,95
						(continu	ied)

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TABLE 1 (continued)

CHEMICAL	AL PROCESS - TREATABILITY		CHEMICAL	PROCESS - TREATABILITY	REF
1,1,2-Trichlo- roethane(cont)	VII Extractable w/aromatic methanol, & ether IX Adsorbed	s 90 90	G. METALS Antimony	II 28,62,65% reduction	39
Frichloro- ethylene	I 99% reduction@ 78-214 μg/1II 40% reduction @ 103μg/	21,65		@ 600 μg/l w/alum, lime ferric chloride coagu- lants	
	w/alum V Air & steam strippable VII 75% reduction w/kero- sene & C ₁₀ -C ₁₂ solvent	90,95	Arsenic	II 76-90% reduction@5mg/l w/ferric sulfate & lime coagulants IX No reduction@1.1 µg/l	63,64
Trichlorofluo-	<pre>IX 99% reduction @ 21 μg/ VII Extractable w/alcohols & ethers</pre>			XII 96% reduction @ 25 mg/l w/silicon alloy adsor- bent	
Frichloro-	<pre>IX Adsorbed V Air & steam strippable</pre>	90 90	Arsenic (As ⁺⁵)	II 94-97% reduction @ 21-25 mg/l w/alum & lime coagulant	90
methane	VII Soluble in most organics	90	Barium	I Inhibitory @ >100 mg/l II 36-99% reduction	109 39,63
1,2,3-Tri- chloropropane	IX 100% reduction@100µg/X 100% reduction@100µg/	1 20		@ 0.08-5 mg/l w/lime, alum, ferric sulfate	64
Vinyl Chloride Vinylidene Chloride	VII 92% reduction w/keroser	1		III 87-99% reduction @ 0.8-9.2 mg/l IX No reduction @ 32 µg/l	18 64
	& C_{10} - C_{12} solvents @ 13 mg/l		Beryllium	II 98-99% reduction @ 100 μg/l w/alum, lime & ferric chloride	39,90
			Bismuth	II 94-96% reduction @ 600 µg/l w/alum, lime & ferric chloride	39
		1	11	(continu	ued)

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TABLE 1 (continued)

CHEMICAL	3	PROCESS - TREATABILITY	REF	CHEMIÇAL	PRC	CESS - TREATABILITY	REF
Cadmium	I	Inhibitory @ 1-10 mg/1	65,90, 109	Chromium (Cr ⁺³)	II	98-99% reduction @ 0.7-5 mg/1 w/ferric	39,63
	II	45-98% reduction @ 9µg/1-	39,63,			sulfate, lime, & fer-	
		5 mg/l w/lime, ferric	64			ric chloride	
		chloride & ferric sulfate			IX	5-48% reduction	72
	III	90-99% reduction	18	_		@ 100 mg/l	
		@ 0.1-1.0 mg/1		Chromium (Cr ⁺⁶)	I	Inhibitory @ 100 mg/l	109
	VI	Foam fractionation	90	,	II		39,63
	Ì	w/sodium dodecylbenzene				@ 0.7-5mg/l w/ferric	,
		sulfonate	64.00			sulfate, lime & fer-	
	TX	6-37% reduction	64,82			ric chloride	
	VIII	@ 1.8-29 μg/l	00		IX	16-36% reduction	72
	XII	96% reduction @ 25 mg/l	90			@ 100 mg/l	
		w/silicon alloy adsorbent		Cobalt	_	Tubibit 0.0 00 /1	3.0.4
		adsorbent		Cobalt	I II	Inhibitory @ 0.08 mg/l 18-91% reduction	124
Chromic Acid	III	85% reduction @ 200 mg/l	24		1 + +	0 500-800 mg/l	39
Chromium	I	27-78% reduction	122			J.	
		@ 0.8-4 mg/l		Copper	I		118,12
	II	27-54% reduction	16,64			@ 0.2-10 mg/l; reported	124,12
		@ 0.1-5 mg/l w/lime	•		1	to be inhibitory	
	III	85-98% reduction	18			@ >0.5 mg/1	
		@ 1-12 mg/1			111	67-98% reduction	16,37
	VI	Reduction possible using	90			@ 0.2-15 mg/l w/alum,	63,64
		quartenary ammonium salts				lime, ferric sulfate	90
	ΙX	37-43% reduction	64		 T T T	coagulants 95-100% reduction	18
		@ 41-84 μg/l			1	0.6-12 mg/l	18
	XII	100% reduction @ 300mg/l	90		TV	82% reduction @0.44mg/1	59
		w/high clay soil	}		VI	Foam fractionation	90
		adsorbent			" "	w/sodium dodecylbenzene	
Chromium	la la	Complete removal	123			sulfonate	
(Cr +3		TT			IX		64.72
, ,	I		l	H	1	100 mg/l (conti	

TABLE 1 (continued)

CHEMICAL	PROCESS - TREATABILITY	REF	CHEMICAL	PROCESS - TREATABILITY REF
Copper (cont)	XII 96-100% reduction @ 300 mg/l w/silicon alloy & high clay soil adsorbents	90	Manganese	I Conflicting data; 109,124 >10 mg/l inhibited while 12-50 mg/l also reported to stimulate II 18-98% reduction 39,63
Iron	I 62% reduction @ 0.6mg/1 soluble iron II 26-99% reduction @0.2-10 mg/1 w/lime & ferric chloride	126 16,63, 64		@ 0.04-5 mg/l w/lime & ferric sulfate coagu-lants IV 89% reduction @ 4.9mg/l IX 1-50% reduction @ 64,72
	coagulants III 100% reduction@12 mg/			@ 0.002-100 mg/1
	IV 85% reduction@6.8 mg/I IX 45-68% reduction @ 40-207 µg/1	59 64	Mercury	I Conflicting data; 51- 127,13 58% reduction @5-10mg/1 & inhibitory @ any
Iron (Fe ⁺²) Iron (Fe ⁺³)	<pre>I Inhibitory@>100 mg/l I Inhibitory@>100 mg/l</pre>	109 109		concentration II 25-98% reduction 39,63 @ 0.001-5 mg/l w/lime 64
Lead	I Inhibitory @ >10 mg/l II 43-99% reduction @ 0.02-5 mg/l w/lime, ferric sulfate, & alum coagulants	109,124 39,63, 64,90		& ferric chloride coagulants VII 99% reduction@2 mg/l 90 w/high molecular weight amines & quartenary salts
	III 98-100% reduction @ 0.9-12 mg/1 VI Foamfractionation w/soci	90		IX 80-99% reduction 64,72 @ 0.001-100 mg/l w/GAC 87,90 & PAC plus chelating
	ium dodecylbenzene sulfate IX 13-93% reduction @ 100 mg/l; no reduction	64,72		agent XII >99% reduction using 90 silicon alloy adsorbent
	@ 5-22 µg/l XII 96% reduction w/silicor alloy adsorbent; red- wood bark also tried		Molybdenum	II No reduction w/alum & 39 lime; 68% reduction w/ferric chloride @ 600 µg/l (continued)

CHEMICAL	P	ROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
Nickel	I	0-42% reduction @ 0.3-10 mg/1 10-100% reduction @ 0.9-5 mg/1 w/alum,	118,122, 125,128, 129 16,39, 63,90	Titanium	II	96-98% reduction @ 500 µg/l w/lime,alum, & ferric chloride coagulants	39 .
	1	lime, & ferric sulfate 93-97% reduction @ 12 mg/l 4-52% reduction @100mg/l	18 72	Vanadium	II	57-97% reduction @ 500 μg/l w/lime, alum & ferric chloride coagulants	39
Selenium		0-80% reduction @ 0.002-100 mg/l w/lime, alum, & ferric chloride coagulants 96% reduction @ 500 mg/l after GAC & lime precipitation		Zinc	II		90,109 118,122 124,128 131 16,39, 63,64
Silver	II	38-98% reduction @ 0.006-500 mg/l w/lime, alum, & ferric chloride coagulants	39,64 90			ferric chloride coagulants 97-100% reduction @ 9-32 mg/l	18
Strontium	I	No affect @ 5-50 μg/l	124			79% reduction @ 1.8 mg/l 61-81% reduction	59 64
Thallium ,	II	30-60% reduction @ 500 µg/l w/lime, alum, & ferric chloride coagulants				@ 0.4-0.6 mg/l 99% reduction @ 10 mg/l w/silicon oxide & cal- cium oxide slags as	
	IX	84% reduction after GAC & lime precipitation	90			absorbents	
Tin	II	92-98% reduction @ 500 µg/l w/lime, alum & ferric chloride coagulants	39				

TABLE 1 (continued)

CHEMICAL	Pl	ROCESS - TREATABILITY	REF	CHEMICAL	PRO	CESS - TREATABILITY	REF
I. PCB's				Atrazine (cont)	х	100% reduction@100µg/1	20
Arochlor 1242	IX	98-99% reduction @45μg/l	8,22,38, 66	Captan	III	99-100% reduction @ 689 µg/l	18
Arochlor 1254	IX X	94-99% reduction @ 11-160 µg/1 100% reduction@100µg/1	8,20,22, 38,66 20,22	Chlordane	I	Slightly degraded 97-100% reduction @ 13-1430 µg/l	121
Arochlor 1254 and 1260		23-60% reduction @ 1-25 μg/l 73% reduction w/PVC	57 57	Chlorinated Pesticides (unspecified)	х	79% reduction @ 33-118 mg/l	49
		chips; 37% reduction w/polyurethane foam adsorbent		2,4-D Butyl ester	IX X	100% reduction @ 100µg/l 100% reduction @ 100µg/l	
PCB's (unspecified)		100% reduction @ 1-400 µg/l	6	2,4-D & related herbicides	x	>95% reduction @ 20-1500 µg/l	32
				2,4-D-Isoctyl- ester	I	Biodegradable	121
				DDD	IX	99.8% reduction @ 56µg/1	8,38,66
J. PESTICIDES				DDE	I.	100% reduction >97% reduction @ 38 µg/l	18 8,38,66
Aldrin	I	Not significantly degraded	121	DDT	I	Not significantly degraded	121
	1	100% reduction 98-100% reduction 08-100 µg/1	18 6,8,20, 38			98% reduction@10 µg/l w/alum coagulant	6
	х	100% reduction@100µg/1				100% reduction >99% reduction	18 6,8,20
Aminotriazole	I	Not significantly degraded	121		x	@ 10-100 µg/1 100% reduction @ 100µg/1	38,66
Atrazine	III	84-98% reduction	18			(conti	nued)

TABLE 1 (continued)

CHEMICAL	I	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
DDVP	I	Degraded	92	Herbicide Orange	I	77% reduction @1380mg/1	81
Diazinon		Not significantly degraded	92,121	Kepone	IX	100% reduction @ 4000 µg/l	6
 Dieldrin	III	88-98% reduction Not significantly	18 121	Lindane	I	Not significantly degraded	121
preform		degraded			II	<10% reduction @ 10 μ g/1	6
	II	55% reduction@10 µg/l w/alum coagulant	6		III	<pre>w/alum coagulant >99% reduction</pre>	18
	ł	100% reduction 75-100% reduction	18 6,8,38,		IX	30->99% reduction @ 10 μg/l	6
		@ 19 , 60 μg/1	66	Malathion	Ι	_	92,121
Endrin	I	Not significantly degraded	121		III	degraded >99% reduction	18
	II	35% reduction@10 µg/l w/alum coagulant	6	Maneb	I	Biodegradable	121
	X	80-99% reduction @ 10-62 μg/l	6,8,38, 66	Methyl Parathion	I	Not significantly degraded	92,121
Endrin &	x	>97% reduction	32			>99% reduction	18
Heptachlor Ferbam	T	@ 0.1-2 mg/l Biodegradable	121	Parathion	I	Not significantly degraded	92,121
Heptachlor	ī	Slightly degraded	121		II	5% reduction @ 10 μg/l w/alum	6
	TTT	@ 500 mg/l 100% reduction	18			>99% reduction >99% reduction @ 10µg/1	18
		>99% reduction @ 6-80µg/1	1	 Pentachlorophenol]	Not significantly	121
Heptachlor- epoxide	III	99.8% reduction	18	(Also see phenols)		degraded @ 75-150 mg/l	
Herbicides	IX	90-99% TOC reduction	38	Propoxur	I	Biodegradable 72-99% reduction	92
(unspecified)				Indiaox	111		inued)

TABLE 1 (continued)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	F	PROCESS - TREATABILITY	REF
Tetraethyl Pyrophosphate	I	Not significantly degraded	121	K. PHENOLS Bisphenol A	х	>94% @ 900 mg/l when pH	. 23
Thanite	I	Biodegradable	121			adjusted	
Toxaphene	IX	97-99% reduction @ 36-155 μg/l	6,8,38	Brine phenol	x	99% reduction of phenol @ 10-400 mg/l	33
į	Х	>99% reduction @ 70-2600 µg/l	32	Butyl Phenol	ıx	95% reduction @ 300 μg/l	6
2,4,5-T ester	II	65% reduction @ 10 µg/l w/alum coagulant	6	4-Chloro-3- Methylphenol	I	Toxic @ 50-100 mg/l Inhibitory but slowly	90, 102
	IX	80-95% reduction @ 10 µg/1	6		VII	degradable @ <50 mg/l Extractable w/benzene, alcohol, & nitrobenzene	90
2,4,5-Tri- chlorophenoxy-	I	Slightly degraded @ 150 mg/l-99% reduction	115		IX X	100% reduction @ 100 µg/l 100% reduction @ 100 µg/l)
acetic Acid		after 7.5 days aeration		2-Chloro-4-	I	72% reduction	81
Trifluralin	III	100% reduction	18	Nitrophenol			ļ
Ziram	I	Slightly degraded	121	Chlorophenol	V	Steam strippable	90
Zireb	I	Slightly degraded	121	m-Chlorophenol	x	100% reduction@200 mg/l Adsorbed	66 66
				2-Chlorophenol	I	90-95% reduction @ 150-200 mg/l	90
					VII	66% reduction	90
,				o-Chlorophenol	I	96-100% reduction @ 200 mg/1	66,8
	1			11		(continu	aed)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
p-Chloropheno	l I	96-100% reduction @ 200 mg/l	66,81	Dimethylphenol	IX	>99% reduction @ 1220 µg/1	6
Cresol	IX	96% reduction @ 230 μg/l	6	2,3-Dimethyl-	I	96% reduction	81
m-Cresol	I	96% reduction 91% reduction @ 291 mg/1	81 27	phenol 2,4-Dimethyl-	ı	0.49	
o-Cresol	Ι	95% reduction 90-99% reduction	81 27	phenol	ı —	94% reduction Extractable w/benzene & alcohol	8 1 90
p-Cresol	п	@ 307-890 mg/l 96% reduction	81	2,5-Dimethyl-phenol	I	94% reduction	81
•	VII	91% reduction @ 291 mg/1	27	2,6-Dimethyl-phenol	I	94% reduction	81
2,4-Diamino- phenol	I	83% reduction	81	3,4-Dimethyl-	I	98% reduction	81
2,4-Dibromo- phenol	x	Adsorbed	33	phenol 3,5-Dimethyl-	I	89% reduction	
Dichlorophenol	LX	Adsorbed	33	phenol	IX	100% reduction @ 100µg/l	81 20
2,3-Dichloro- phenol	IX X	100% reduction @ 100 µg/l 100% reduction @ 100 µg/l	20 20	4,6-Dinitro-2- Methylphenol	VII	Extractable w/benzene & acetone	90
2,4-Dichloro- phenol	I	98-100% reduction @ 60-200 μg/l	81,90, 115	2,4-Dinitrophenol	1	Extractable w/benzene	81,113 90
	XII	100% reduction @ 430 mg/l Extractable w/benzene,	66 90			& alcohol Adsorbed	21
2,5-Dichloro-	ļ_	alcohol, & nitrobenzene 100% reduction@ 200 mg/l	6.6	B-Napthol	X	100% reduction @ 100μg/1	20
phenol	ľ	1000 reduction e 200 mg/1	66	m-Nitrophenol	I	95% reduction	81
2,6-Dichloro-	I	99% reduction@64 mg/l	115	o-Nitrophenol	I	97-98% reduction	58,8
phenol				2-Nitrophenol	VII	Extractable w/benzene & alcohol	90
				_		·	nue

TABLE 1 (continued)

CHEMICAL		PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
p-Nitrophenol	I	95-99% reduction	58,81	p-Phenylazophenol	I	Inhibitory @ 500 mg/l	108
	X	>99% reduction @ 700-1800 mg/l	23,33	Resorcinol	IX X	100% reduction @100µg/1 100% reduction @100µg/1	
4-Nitrophenol	1	Removable Extractable w/benzene & alcohol	90 90	Sodium Pentachlo- rophenol		No reduction @ 15 mg/l	120
Nonylphenol	ıx	Adsorbed	21	2,3,5-Trichloro- phenol	I	100% reduction @200mg/1	66,92
Pentachloro- phenol	VII	26% reduction @ 200 mg/l Extractable w/benzene, alcohol & nitrobenzene	66,92 90	2,4,5-Trichloro- phenol	I	99% reduction@19 mg/1	115
	IX X	100% reduction@10 mg/l 100% reduction@100 μg/l	6,21 20	2,4,6-Trichloro- phenol	1	100% reduction @ 20- 200 mg/l; reported to	66,90, 102,115
Phenol	I	62-100% reduction @ 5-500 mg/1; reported to be inhibitory @ 500 mg/1	58,66, 88,90, 92,106, 108,118, 119		IX	be inhibitory @ 50- 200 mg/l Extractable w/benzene, alcohol, nitrobenzene 100% reduction @100µg/l	90 20
	III	-6 - 100% reduction @ 1-1000 mg/l dependent	18,30 54,90		Х	100% reduction @ 0.1-510 mg/1	20,60
		upon membrane	_	Trimethylphenol	IX	92% reduction@130 μg/l	6
	v	75% reduction @ 1-100mg/1 Steam strippable 4-98% reduction @ 67-8800 mg/1	54 90 27,90	Xylenol	VII	96% reduction @ 227 mg/l	27
	IX	80-100% reduction @ 0.1-1200 mg/l	6,20,21 35,38, 72,90,				
	х	>99% reduction @ 500-5000 mg/l	23,33				
						(conti	nued)

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CHEMICAL	-	PROCESS - TREATABILITY	REF	CHEMICAL	PRC	CESS - TREATABILITY	REF
L. PHTHALATES Bis(2-Ethyl-	т	70-78% reduction @ 5mg/1	90	Dimethyl Phthalate		Degradable; 100% reduction @ 215 µg/l	21,90
1	ΪΙ	80-90% reduction @ 0.5- 3.5 µg/l w/aluminum	90			15% reduction@183µg/1 w/alum Extractable w/ethyl	90
		sulfate coagulant Extractable w/ethyl ether & benzene	90		IX X	ether & benzene 100% reduction@100µg/1 100% reduction@100µg/1	
Butylbenzyl Phthalate	I	>98% reduction@1300µg/l Biodegradable Extractable w/ethyl ether & benzene	5,90 90 90	Di-N-Octyl Phthalate	VII	Biodegradable @ 63 mg/l Extractable w/ethyl ether & benzene	90 90
-	тх х	100% reduction @ 100 μg/l 100% reduction @ 100 μg/l	20 · 20	Isophthalic Acid Phthalimide	I	95% reduction 96% reduction	81 81
Di-N-Butyl Phthalate		Biodegradable @ 200 mg/l 60-70% reduction @ 2.5-4.5 µg/l w/aluminum sulfate Extractable w/ethyl ether & benzene	90 90 90	Phthalic Acid	I	97% reduction	81
Diethyl Phthalate	,	Biodegradable Extractable w/ethyl ether & benzen	90 90	M. POLYNUCLEAR AR	омат	ICS	
Diethylhexyl	x	100% reduction @ 100 µg/1	20	Acenaphthalene	х	100% reduction@100µg/	20
Phthalate		50 500 1 11		Acenaphthene	II	Precipitated w/alum	90
Di(2-ethyl- hexyl)Phtha-	I	50-70% reduction	100	Acenaphthylene	II	Precipitated w/alum	90
late				Anthracene	VII	Toxic @ 500 mg/l Extractable w/toluene	108 90
l	1		1	H	l	(conti	inued)

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TABLE 1 (continued)

CHEMICAL	T	PROGRAGA TRANSPORT			1		
CHEMICAL	-	PROCESS - TREATABILITY	REF	CHEMICAL	PRO	OCESS - TREATABILITY	REF
Benzanthracen	II	Slowly oxidized@500mg/l Separable by gravity or sand filtration	108 90	Dimethylnaptha- lene	IX X	80% reduction @ 100μg/l 100% reduction @100μg/l	20 20
11,12-Benzo- fluoranthene	II	Separable by gravity or sand filtration	90	1,1,-Diphenyl- hydrazine	IX	Adsorbed	31
Benzoperylene	I	Biodegradable	90	1,2-Diphenyl- hydrazine	I	28% reduction @ 341µg/1	81
1,12-Benzo- perylene	II	Separable by gravity or sand filtration	90	Fluoranthrene	IX X	80% reduction@100µg/1100% reduction@100µg/1	20 20
Benzo(a)- pyrene	II	Separable by gravity or sand filtration	90	7-Methyl-1,1- benzanthracene	I	Inhibitory @ 500 mg/l	108
Biphenyl	x	100% reduction@100 μg/l 100% reduction@100 μg/l	20 20	20-Methylchol- anthrene	I	Toxic or inhibitory; able to undergo slow	108
D-Chloram- phenicol	I	86% reduction	81			biological oxidation @ 500 mg/l	
2-Chloro- napthalene	II	Precipitated w/alum	90	Napthalene	I	85-95% reduction; Inhibitory @ 500 mg/l	56,101, 108
Chrysene	II	Separable by gravity & sand filtration	90		V	Separable by gravity or sand filtration Air strippable by 50:1	90 90
Cumene	IX X	100% reduction @ 100 μg/l 100% reduction @ 100 μg/l	20 20	Phenanthrene	IX	volume of air 70% reduction 80% reduction @ 100µg/l	31 ₂₆ 64
α, α -Diethyl-	ı	Inhibitory	108		Х	100% reduction@100µg/1	20
stilbenediol 9,10-Dimethyl	-I	Degradable@500 mg/l	108	2,3-o-Phenylene Pyrene	II	Separable by gravity or sand filtration	90
anthracene				Pyrene	II	Separable by gravity or sand filtration	90
9,10-Dimethyl 1,2-benzan- thracene]I	Slowly oxidized @ 500 mg/l	108		IX X	80% reduction @ 100µg/l 100% reduction @100µg/l	20 20

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SECTION 4

WASTE STREAM IDENTIFICATION AND CHARACTERIZATION

PROBLEM TYPES

One of the early activities undertaken in this project was an effort to identify actual hazardous aqueous waste problems faced by the public sector which might benefit from the application of concentration technology. This effort was accomplished primarily through personal contacts with governmental entities and companies involved in hazardous waste management since little published information existed. Appendix A contains a list of entities contacted. In many cases there were several contacts within the entity listed.

Individuals contacted were queried about major problems known to them in terms of hazardous materials in aqueous solutions and specifically, priority pollutants. The predominant response was that discharge from waste storage and disposal sites were the biggest problem.

Responders indicated that these discharges, generally leachate, were becoming more numerous and severe, and will become more prevalent as wastewater pretreatment regulations are enforced and greater volumes of residues containing concentrated hazardous materials are produced. Because of the current deficiency in the number of controlled landfills, many sludges and hazardous materials will not receive adequate disposal, and additional discharge and leachate problems can be expected.

Even though there is no such thing as a typical hazardous waste problem, and each site is unique, problems generally can be grouped into three broad categories: 1) land disposal sites; 2) container storage and disposal sites; and 3) lagoons.

Land disposal sites range from simple dumps to fully secured chemical landfills, and can be actively operated or abandoned and inactive. Although there are landfills devoted exclusively to industrial wastes, many co-dispose municipal and industrial liquids, sludges and solid wastes together. Responsibilities and assignment of potential liabilities for active landfills are fairly clear, but for inactive or abandoned sites responsibility

usually devolves to some governmental entity.

Container storage and disposal sites represent a considerable problem. Only recently has the magnitude and potential danger been recognized. In some cases, containers have been breached and concentrated wastes flowed into surface and ground waters. This has been a cause for grave concern, because some container disposal sites encompass many acres and thousands of barrels, drums, tanks, etc. Many containers are in various stages of progressive failure, thus constituting potential problems of enormous magnitude and complexity. For example, conjunctive disposal of containers of corrosive, reactive, flammable, and toxic materials could result in breaching in "domino" fashion if there is failure and leakage from very few. Examples of such situations have been uncovered during the interviews. In one case, excavation of buried drums ceased after several underground detonations.

Most lagoons which cause problems are unlined. Evidence that the integrity of a disposal lagoon has been breached is found as ground or surface water contamination. Such contamination occurs by: 1) vertical percolation, 2) overland flow, or 3) flood flushing. In the case of vertical percolation, wastes are transported through the porous lagoon bottom, through the soil vadose zone, and into the ground water table. Overland flow involves a combination of horizontal percolation and chronic lagoon overflows to surface water. In contrast, flood flushing entails acute release of lagoon contents because of an extreme rainfall event or dike failure.

Occasional discharges of hazardous wastes to municipal sewerage systems have been reported. These generally resulted from spill incidents, either accidental or intentional. While some such discharges have been porblems, those interviewed regarded the leachate and discharge problem as being far more important.

Three other potentially significant problems were considered for inclusion in this project: 1) drum and container contents, 2) waste-contaminated lagoon contents, and 3) sludges. In the first two instances, although neither is dilute, often the wastes can be concentrated further. On the other hand, sludges were deemed important to this project only in terms of the degree to which they are leached or their liquid component drains to ground and/or surface water.

Upon consideration of all of the available information, it was decided to focus primarily on leachate and contaminated ground and surface waters associated with hazardous waste disposal sites. This decision largely was based upon the fact that leachate contamination is believed by knowledgeable individuals to be the largest and most pressing of the cited problems. Moreover, little available data exists on leachate

treatment for hazardous waste repositories and industrial disposal sites. Thus, this project can contribute to plugging the information gap.

WASTE STREAM COMPOSITION

Having set the focus of the project on leachates, and contaminated ground and surface waters, an effort was launched to obtain composition data on known problem sites. This effort was complicated by several factors: 1) little published information exists, 2) record-keeping and reporting procedures for hazardous waste problems are sketchy, 3) actual or potential litigation causes data to be restricted, 4) because of lack of funding only the most severe problems have received attention, and 5) high analytical costs associated with specific organic compound identification often causes measurement and reporting of surrogate parameters such as TOC, COD, and BOD.

Despite the above cited problems, it was possible to obtain composition data on leachates, and contaminated ground and surface waters in the proximity of 27 sites containing hazardous wastes. Much of the obtained data is unpublished.

Because of the large quantity of data, this information is summarized in Appendix B, Table B-1. In addition to data on the 27 sites, this table contains summary data on 43 industrial disposal sites which were surveyed in a previous study (127). There is a wide variation from site to site in the detail and completeness of the data contained in Table B-1 since relatively few waste streams have been well characterized. Nevertheless, this data compilation represents the best available information and is believed to be one of the most complete available at this time.

Study of the compiled data shows that wastes encountered are diverse in terms of composition and concentration. Some contain a broad spectrum of organic and inorganic materials, while others may have only a limited number of compounds. A wide variability in waste composition is observed from site to site. Moreover, waste composition often is highly variable at any given site with respect to both time and location.

WASTE CONSTITUENT CLASSIFICATION

Because of the large number of chemicals and possible combinations and permutations of constituents in hazardous waste streams, it would be desirable to employ predictive techniques to forecast the behavior of chemicals present in such waste streams. Unfortunately, no proven method exists to accurately predict the removability of all of the potential chemical

constituents of hazardous aqueous waste streams.

Nevertheless, some grouping or classification of waste stream constituents was deemed desirable to extend the usefulness of the data and facilitate the evaluation of concentration technology. Therefore, a contaminant classification system was formulated as given in Table 2. This classification system was based upon the twelve groups of compounds that were used to classify the 129 priority pollutants that resulted from the 1976 Flannery Consent Decree (NRDC vs. Train, June 1976). slightly modified categories given in Table 2 were considered a better reflection of compounds actually detected at identified hazardous waste contamination sites. All of the identified constituents of the actual hazardous waste streams given in Table B-l have been classified according to this system. The results of this classification effort together with an indication of the frequency of identification of each constituent is given in Table 3. The number of different sites where compounds in each classification were identified is given below:

Alcohol	2
Aliphatic	4
Amine	2
Aromatic	8
Halocarbon	9
Metal	15
Miscellaneous	11
PCB	2
Pesticide	7
Phenol	7
Phthalate	2
Polynuclear Aromatic	5

This classification system was used to aid in the technology screening effort as described in a subsequent section of this report.

TABLE 2

CONTAMINANT CLASSIFICATION SYSTEM

- 1. Alcohol
- 2. Aliphatic
- 3. Amine
- 4. Aromatic nonhalogenated and halogenated aromatic compounds
- 5. Ether
- 6. Halocarbon halogenated aliphatic compounds
- 7. Metal
- 8. Miscellaneous including selected priority pollutants, pH, BOD, TOC, COD, chloride, sulfate, phosphate, and other parameters generally used to characterize wastewaters
- 9. PCB
- 10. Pesticide
- 11. Phenol including chloro and nitro phenols
- 12. Phthalate
- 13. Polynuclear Aromatic

TABLE 3
SUMMARY LIST OF CONTAMINANTS REPORTED

CONTAMINANT CLASSIFICATION	CONTAMINANT	CONCENTRATION RANGE REPORTED	NO. OF SITES REPORTED
Alcohol	Chlorobenzyl alcohol	P	1
	Ethanol	56.4 mg/l	1
	2-ethylhexanol	$19,000 - 23,000 \mu g/1$	1
	Isopropanol	< 0.1 mg/l	1
	Methanol	42.4 mg/l	1
Aliphatic	Acetone	50.3 mg/l	1
-	Dicyclopentadiene	80 - 1200 μg/l	1
_	Diisopropylmethylphosphonate	400 - 3600 μg/l	1
	2-ethylhexanol	ND - $4500 \mu g/l$	1
	3-heptanone	ND - 1300 $\mu g/1$	1
	Hexachlorocylohexane*		
	alpha isomer*	ND - 600 μg/l	1
	beta isomer*	ND - $70 \mu \text{g}/1$	1
	gamma isomer*	ND - $600 \mu g/1$	1
	delta isomer*	$ND - 120 \mu g/1$	1
	Methyl isobutyl ketone	2000 μg/l	1
	Paraffins	P	1
	Pinene	P	11
Amine	benzylamine or o-toluidine	<10 - 471 μg/l	1
	n-nitrosodiphenylamine	190 μg/1	ī
Aromatic	m-acetonylanisola	<3 - 1357 μg/1	1
	Aniline	140 - 870 μg/l	l ī
	Benzaldehyde	P	1 1
	Benzene*	6 - 7370 μg/l	4
	Benzene hexachloride	P	i
	Benzoic acid	<3 - 12,311 μg/1	$\overline{1}$
	Camphene	P	l ī
	Camphor	<10 - 7571 μg/1	$\bar{1}$
	Chloraniline	<10 - 86 µg/1	l ī
	o-chloroaniline	ND - 360 μg/1	l ī
	1	1	(continued)

TABLE 3 (continued)

CONTAMINANT CLASSIFICATION	CONTAMINANT	CONCENTRATION RANGE REPORTED	NO. OF SITES REPORTED
Aromatic	Chlorobenzaldehyde	P	1
(continued)	Chlorobenzene*	4.6 - 4620 μg/l	· 4
	4-chloro-3-nitro benzamide	440 - 8700 μg/l	1
	p-chloronitrobenzene	460 - 940 μg/l	1
	Chloronitrotoluene	ND - $460 \mu g/1$	1
	p-chlorophenyl methyl		
	sulfide	<10 - 68 μg/l	1
	p-chlorophenyl methyl		
	sulfone	$<10 - 40 \mu g/1$	1
	p-chlorophenyl methyl	1	
	sulfoxide	<10 - 53 μg/1	1
	2,6-dichlorobenzamide	890 - 30,000 μg/l	1
	Dichlorobenzene*	<10 - 517 μg/1	2
	Dimethyl aniline	<10 - 6940 μg/1	1
	m-ethylaniline	<10 - 7640 μg/1	1
	Ethyl benzene*	$3.0 - 470 \mu g/1$	2
	Hexachlorobenzene*	$32 - <100 \mu g/1$	2
	p-isobutylamisol ^a or		
	p-acetonylanisola	<3 - 86 μg/l	1
	Limonene	P	1
	Nicotinic acid	P	1
	o-nitroaniline	$170,000 - 180,000 \mu g/1$	1
	p-nitroaniline	$32,000 - 47,000 \mu g/1$	1
	Nitrobenzene*	$ND - 740 \mu g/1$	1
	Styrene	P	1
	Toluene*	$< 5 - 31,000 \mu g/1$	4
	1,2,4-trichlorobenzene*	$<10 - 28 \mu q/1$	2
	Trimethylbenzene	P 20 19/1	$\bar{1}$
	Xylene	P - 3300 μg/l	ī
			(continued)

TABLE 3 (continued)

C ₄ alkyl cyclopentadiene Bromodichloromethane* Chloroform* C ₅ substituted cyclopentadiene Dibromochloromethane* 1,1-dichloroethane* 1,2-dichloroethane* 1,1-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloroethylene* 1,2-dichloropropene* Hexachlorocyclopentadiene* Octachlorocyclopentadiene* Octachlorocyclopentadiene* Octachlorocyclopentadiene* 1,1,2,2-tetrachloroethane* 1,1,2,2-tetrachloroethane* Tetrachloromethane* Tetrachloromethane* Trichloroethane* 1,1,1-trichloroethane* 1,1,1-trichloroethane* 1,1,1-trichloroethane* 1,1,2-trichloroethylene* Trichloroethylene* Trichloro	CONTAMINANT ASSIFICATION	CONTAMINANT	CONCENTRATION RANGE REPORTED	NO. OF SITES REPORTED
Octachlorocyclopentene Perchloroethylene* 1,1,2,2-tetrachloroethane* Tetrachloroethylene* Tetrachloromethane* Tribromomethane* 1,1,1-trichloroethane* 1,1,2-trichloroethane* Trichloroethene Trichloroethene Trichloroethene Trichloroethene Trichloroethene Trichloroethene Trichloroethylene* Octachlorocyclopentene ND - 1000 µg/l <1 - >50,000 µg/l <1 - 25,000 µg/l <1 - 25,000 µg/l P - 490 µg/l 1.6 - 532 µg/l <5 - 870 µg/l <3 - 10,000 µg/l Trichloroethylene* Trichloroethylene*		C4 alkyl cyclopentadiene Bromodichloromethane* Chloroform* C5 substituted cyclopenta- diene Dibromochloromethane* 1,1-dichloroethane* 1,2-dichloroethane* trans-1,2-dichloroethane* 1,1-dichloroethylene* 1,2-dichloroethylene* Dichloromethane* Dichloromethane* Hexachlorobutadiene*	P ND - 35 μg/1 0.02 - 4550 μg/1 P 3.9 μg/1 <5 - 14,280 μg/1 2.3 - 330 μg/1 25 - 8150 μg/1 28 - 19,850 μg/1 0.2 μg/1 3.1 - 6570 μg/1 P <20 - 109 μg/1	1 1 3 1 1 4 2 4 1 4 1 2
Trichloroethylene* 760 - 260,000 μg/l		Hexachlorocyclopentadiene* Octachlorocyclopentene Perchloroethylene* 1,1,2,2-tetrachloroethane* Tetrachloroethene Tetrachloroethylene* Tetrachloromethane* Tribromomethane* Trichloroethane* 1,1,1-trichloroethane* 1,1,2-trichloroethane*	<pre><100 µg/1 <100 µg/1 ND - 1000 µg/1 <5 - 1590 µg/1 <1 - >50,000 µg/1 23 - 590 µg/1 <1 - 25,000 µg/1 0.2 µg/1 P - 490 µg/1 1.6 - 532 µg/1 <5 - 870 µg/1</pre>	1 1 1 1 3 2 1 1 4 2
Trichloromethane*		Trichloroethylene* Trichlorofluoromethane* Trichloromethane*	760 - 260,000 µg/l <5 - 18 µg/l <1 - <10,000 µg/l	3 3 1 1

(continued)

TABLE 3 (continued)

CONTAMINANT	1	CONCENTRATION	NO. OF SITES
CLASSIFICATION	CONTAMINANT	RANGE REPORTED	REPORTED
Metal	Ag*	1 - 10 μg/1	2
	Al	0.124 mg/l	1
	As*	0.011 - >10,000 mg/1	6
	Ba	0.1 - 2000 mg/l	5
	Be*	0.007 mg/1	1
	Во	0.624 mg/1	1
	Ca	164 - 2500 mg/l	4
	Cd*	0.005 - 8.2 mg/l	6
•	Co	0.01 - 0.22 mg/l	1
	Cr*	< 0.001 - 208 mg/l	7
	Cu*	0.001 - 16 mg/1	9
	F	0.14 - 1.3 mg/1	1
	Fe	0.090 - 678 mg/l	6
	Hg*	0.0005 - 0.007 mg/1	7
	K	6.83 - 961 mg/l	3
	Mg	25 - 453 mg/1	3
	Mn	0.01 - 550 mg/l	4
	Mo	0.1 - 0.24 mg/l	3
	Na	4.6 - 1350 mg/1	5
	Ni*	0.02 - 48 mg/l	4
•	Pb*	0.001 - 19 mg/l	6
	Sb*	2 mg/1	1
	Se*	0.003 - 0.59 mg/l	4
	Zn*	0.024 - 240 mg/l	10
Miscellaneous	Alkalinity, as CaCo ₃	20.6 - 5400 mg/l	3
	BOD ₅	42 - 10,900 mg/l	3
	Cl	3.65 - 9920 mg/l	6
	CN	0.0005 - 14 mg/l	2
	COD	24.6 - 18,600 mg/1	5
	color	50 - 4000	1
	Halogenated Organics	0.002 - 15.9 mg/l	1
	1	1	(continued)

TABLE 3 (continued)

CONTAMINANT CLASSIFICATION	CONTAMINANT	CONCENTRATION RANGE REPORTED	NO. OF SITES
Miscellaneous	Hardness, as CaCo ₃	700 - 4650 mg/l	2
(continued)	Heavy Organics	0.01 - 0.59 mg/l	i
	Light Organics	1.0 - 1000 mg/1	i
	MBAS	0.24 mg/1	ī
	NH ₃ -N	<0.010 - 1000 mg/1	3
	NH 4 -N	0.65 mg/l	1 1
	NO ₂ -N	<0.010 - <.1 mg/1	$\frac{1}{2}$
	NO ₃ -N	0.010 - < .1 mg/1	2 3
	Oil & Grease	90 mg/1	li
	рН	\sigma_3 - 7.9	7
	PO ₄	< 0.010 - 2.74 mg/1	4
	SO ₄	1.2 - 505 mg/l	4
	soc	4200 mg/l	1
	Specific Conductance	80 - 2000 mg/l	2
	SS	< 3 - 1040 mg/l	4
	Sulfide	<0.1 mg/l	1
	TDS	1455 - 15,700 mg/1	4
	temperature	$58 - 63^{\circ} F$	1
	TKN	<1 - 984 mg/l	4
	TOC	10.9 - 4300 mg/l	7
	Total Inogranic Carbon	71 mg/1	1
	Total P	<0.1 - 3.2 mg/1	2
	Total Solids	159 - 1730 mg/l	1
PCB's	Aroclor 1016*/1242*	110 - 1900 μg/l	1
	Aroclor 1016*/1242*/1254*	$66 \mu g/1 - 1.8 g/1$	(1
	Aroclor 1242*/1254*/1260*	$0.56 - 7.7 \mu g/l$	1
	Aroclor 1254*	70 μg/l	1
Pesticide	Aldrin*	<2 - <10 μg/1	2
	Carbofuran	P	1
	DDT*	4.28 - 14.26 μg/l	1
	Dieldrin*	<2 - 4.5 μg/l	1
	ı	1	(continued)

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TABLE 3 (continued)

CONTAMINANT CLASSIFICATION	CONTAMINANT	CONCENTRATION RANGE REPORTED	NO. OF SITES REPORTED
Pesticide	Endrin*	<2 - 9 μg/l	1
(continued)	Heptachlor*	573 μg/l	1
	Kepone	2 mg/l	1
	Nemagon	<1 - 8 µg/l	1
Phenol	p-2-oxo-n-butylphenol	<3 - 1546 μg/l	1
	o-sec-butylphenola	<3 - 83 μg/l	1
	p-sec-butylphenol ^a	<3 - 48 μg/l	1
	2-chlorophenol*	3 μg/l - 20 μg/l	2
	Dimethylphenol	<3 μg/l	1
	2,4-Dinitrophenol*	10 - 99 μg/l	2
	1-ethylpropylphenol ^a	<3 μg/l	1
	Isoprophylphenol ^a	<3 - 8 μg/l	1
	o-nitrophenol*	8600 - 12,000 μg/l	1
	Pentachlorophenol*	2.4 mg/l	1
	Phenol*	<3 - 17,000 μg/l	4
	Phenols*	0.008 - 54.17 μg/l	1
	2,4,5-trichlorophenol	P	1
Phthalate	Phthalate esters	P	1
	Phthalates*	P	1
Polynuclear	Biphenyl napthalene	P	1
Aromatic	Methyl napthalene	<10 - 290 μg/l	1
	Napthalene*	<10 - 66 μg/l	1
	Petroleum oil	P	1
	Phenanthrene* or anthracene		1
	Polynuclear aromatics	3400 μg/l	1
ND - not detect	ed		
	out not quantified		
a - structure	not validated by actual compour	da	
* - Priority P			
<i>1</i> -			l l

SECTION 5

TECHNOLOGY EVALUATION APPROACH

An iterative approach was deemed to be the most effective means of evaluating technologies with potential application to concentration of hazardous constituents of aqueous waste streams. Moreover, although it was recognized that, ultimately, process trains must be evaluated, it was considered most reasonable to begin with an examination of unit processes. Thus, unit processes were screened in increasing levels of detail until there was sufficient justification to either reject or carry forward the process. Technologies which survived the screening then were incorporated in process trains which were subjected to desktop analysis of their ability to treat actual waste streams. Waste streams were selected from those identified in the previously described waste stream characterization portion of this study.

The initial step in the evaluation consisted of identifying technologies with potential application to concentration of hazardous constituents of aqueous wastes. Thus, early in the project, the following list of candidate technologies was developed:

- Biological Treatment
- Carbon Adsorption
- Catalysis
- Centrifugation
- Chemical Precipitation
- Crystallization
- Density Separation
- Dialysis/Electrodialysis
- Distillation
- Evaporation
- Filtration
- Flocculation
- Ion Exchange
- · Resin Adsorption
- Reverse Osmosis
- Solvent Extraction
- Stripping
- Ultrafiltration

Technology profiles then were prepared for each of the

candidate unit processes. The focus of this step was on the characteristics of the technology without regard to specific waste streams to be treated. Factors considered in development of the technology profiles included:

- state of development,
- range of technology application,
- process flexibility,
- process reliability,
- economic and engineering constraints in technology modification and application
- start-up requirements,
- · efficiency,
- specific limitations,
- energy requirements,
- form of concentrated material, and
- environmental acceptability.

Technology profiles formed the basis for the initial screening of the applicability of individual technologies to concentration of hazardous constituents of aqueous wastes. At this point, certain technologies were eliminated from further consideration for reasons discussed in the individual technology profiles. Remaining technologies were carried forward for more detailed review.

The next step in the evaluation process was an extensive literature review which focused on the technologies which survived the initial screening and upon chemical compounds in the classes identified in the preceding section of this report as having been identified as constituents of actual hazardous aqueous waste streams.

Since it was evident that no single unit process would be sufficient in itself to adequately treat the diverse waste streams in question, five candidate process trains were formulated as being most broadly applicable to the types of waste streams identified in Table B-1. In addition, two actual waste stream compositions were selected from this table for use in the next step in the technology screening. A third waste stream composition was hypothesized subsequent to examination of all of the available composition data. It is believed that the three selected waste stream compositions cover a range of constituents and concentrations representative of actual problems likely to be encountered.

A desktop analysis then was performed to assess the ability of each of the five process trains to treat each of the three waste streams.

Simultaneously, selected vendors were requested to evaluate the ability of their technology to adequately treat the three waste streams in question. The results of these evaluations provide a basis for making an initial judgment on the applicability of a given concentration technology to specific situations in the absence of expermental data. In addition, these evaluations were utilized to select and arrange technologies in priority order for experimental study in the next phase of this project.

Subsequent sections of this report discuss each of the steps in the technology evaluation in detail.

SECTION 6

TECHNOLOGY PROFILES

This section contains brief descriptions of each of the candidate technologies together with an initial assessment of the potential applicability of each technology to concentration of hazardous constituents of aqueous waste streams. The focus here is on the characteristics of the technology without regard to specific waste streams to be treated.

Each technology is described and past applications are indicated. No attempt has been made to provide detailed information on the theory, design or operation of the technologies since such information is readily available in standard texts and design manuals. Rather, the basic features of the technologies are highlighted and the potential for the application of interest is assessed.

BIOLOGICAL TREATMENT

Process Description

Biological treatment involves the utilization of microorganisms to decompose organic matter present in wastewater. microorganisms metabolize the organic matter to yield energy for snythesis, motility and respiration. Biological utilization of organic compounds involves a series of enzyme-catalyzed reactions. Simple dissolved or soluble organic compounds are readily incorporated into the cells of microorganisms and oxidized. When microbial cells come into contact with complex organics, extracellular enzymes are released outside the cells to hydrolyze high molecular weight materials into diffusible fractions, enabling their transport through the cell wall for assimilation. Thus the larger, more complex organic compounds are metabolized at a much slower rate. Although microorganisms may be adapted and grown on many types of organic materials, there are some complex organic compounds that will not be removed by biological oxidation; and these are called "refractory" organic compounds.

Inorganics may be partially removed from the liquid phase and concentrated in the biomass during biological treatment through the mechanism of adsorption. However, inorganics are not destroyed by biological treatment and, in fact, at higher

concentrations may be inhibitory to biological processes.

Biological systems can take a variety of forms. A primary distinction is the mode of respiration and synthesis - aerobic or anaerobic. Aerobic biological processes employ microorganisms which require oxygen for their existence while anaerobic processes are carried out in the absence of oxygen. The former processes proceed more rapidly and produce larger quantities of biomass residual than the latter.

Three types of aerobic systems are of primary concern in the current context: trickling filter, activated sludge, and lagoons.

- l. Trickling Filter Process The trickling filter process consists of a fixed bed of supporting media (e.g., crushed rock, plastic medica, redwood slats) upon which a biological slime layer is grown. Wastewater is intermittently or continuously applied to the top of the filter and flows downward through the filter, passing over the layer of microorganisms. Dissolved organic material and nutrients in the wastewater are taken up by the zoogleal film layer for utilization by the microbial population. Oxidized end products are released back to the liquid. A trickling filter will operate properly as long as the void spaces are not clogged by solids or excessive growth of the biological film layer. The biological film layer grows and gradually increases in thickness to the point that hydraulic shear force from the downward flow of wastewater causes portions of the film layer to slough off the filter media.
- Activated Sludge Process The activated sludge process, in one of its several modifications, is probably the most commonly used aerobic biological waste treatment process. It is dependent upon the maintenance of a flocculant suspension of microorganisms which is dispersed in intimate contact with the waste to be treated. In operation of the activated sludge process, wastewater containing soluble organic compounds is fed to an aerobic reactor (aeration tank) which furnishes (1) air required by microorganisms to biochemically oxidize the waste organics, and (2) mixing to insure intimate contact of microorganisms with the organic waste. The aerobic reactor contents are referred to as mixed liquor. In the vigorously mixed aerobic reactor, the organic wastes are metabolized to provide energy and growth factors for the production of more microorganisms with the release of carbon dioxide and water as metabolic end products. Organic waste compounds are thus degraded to innocuous end products and microorganisms. Mixed liquor flows from the aeration tank to a sedimentation tank, which provides quiescent settling to allow separation of the biological solids from the treated wastewater. The treated and clarified water is collected and discharged as process effluent. Most of the settled biological solids are recycled as return activated sludge back to the aerobic reactor

to provide an activated mass of microorganisms for continuous treatment of incoming wastewater. Some of the settled biological solids are wasted to maintain a proper balance in the population of microorganisms in the mixed liquor of the aerobic reactor. Recycling and wasting of biological solids (microorganisms) from the reactor assures a proper ratio of incoming waste to the population of microorganisms (food to microorganisms, or F/M ratio), which is critical to efficient biodegradation of soluble organic waste compounds.

3. Lagoons - The use of lagoons as a biological treatment technique provides an attractive option where land costs are relatively low. Several types of lagoons are possible, but they all share some basic features.

A long residence time for the incoming wastewater (in excess of 7-10 days) provides sufficient time for sedimentation of solids to occur. The long residence time provides an opportunity for biological decomposition of degradable organic material. In some lagoons, mechanical aeration may be provided to enhance oxygen concentrations; other lagoons may provide no aeration, but may rely solely upon natural processes such as air-water transport, and photosynthetic oxygen production by algae. In the latter types of lagoons, especially in deeper situations, the lagoon bottom may become anaerobic, and some of the properties described for anaerobic processes may pertain.

Climatic conditions may limit the application of lagoons to areas or seasons where icing conditions do not prevail.

The capability of anaerobic biological processes to degrade many organic compounds is less than that of aerobic processes, as is the rate of degradation. However, anaerobic biological processes are attractive for the more readily degraded compounds in concentrated form, inasmuch as the relative energy costs may be less for these processes, as compared with aerobic biological processes, and they offer the possibility for energy recovery in the form of methane gas production. Furthermore, rather than producing excess sludge, as in the aerobic processes, the anaerobic processes generally may be operated at levels of negligible solids production.

Anaerobic degradation typically has been used for treatment of sludges. However, more recently attention has been given to treatment of aqueous organic wastes of widely varying strengths by anaerobic processes. Instead of stirred, sealed reactors as used for sludge digestion, upflow anaerobic filters generally are used. Filters may be packed with a support medium for anaerobic microorganisms to become attached or use a configuration which encourages formation of a high density floating sludge blanket. During the residence in the reactor, solids and complex organic materials in the waste are broken down to organic acids

and alcohols. These acids are then biologically converted to methane and carbon dioxide, which may be withdrawn as a gas. The methane may be used on-site, or sold as fuel. A portion of the methane may be used to maintain the reactor at elevated temperatures.

Process Applications

Biological treatment has been applied successfully to a wide variety of aqueous waste streams with organic contaminants.

Trickling filters have been used by many municipalities for the treatment of domestic wastewater. In addition they are regarded as especially suitable for the treatment of high strength wastes prior to other biological or physical-chemical processes. They have been used extensively in the treatment of cannery, pharmaceutical, and petrochemical wastes. Treatment of refinery wastewaters containing oil, phenol and sulfide is a common application.

The activated sludge process has been used extensively in municipal wastewater treatment. Industrial applications include treatment of wastes from canneries, breweries, pulp and paper mills, petrochemical plants, refineries, textile mills, steel mills, and pharmauceutical plants.

Lagoons have been utilized to treat the same categories of waste streams and organic species as the activated sludge process.

Anaerobic processes have been used in the treatment of high-strength organic wastes, municipal wastewater sludges, and agricultural and municipal solid wastes. The broadest application has been for the treatment of sludges generated in the treatment of municipal sewage. Recently the anaerobic filter process or modifications of the process have been used to treat pharmaceutical, petrochemical, coal gasification, and other organic wastes. Full and laboratory scale studies using industrial wastes have examined a broad range of operating conditions; organic loading rates have ranged from 240 to 4000 kg COD/M³/day (15 to 250 lb COD/1000 ft³/day) and empty bed liquid retention times have ranged from 0.33 to 14 days. While anaerobic digestors commonly are heated to 35°C, or above, to increase the rate of biological decomposition, researchers have reported minimal temperature influence on anaerobic filter performance in the 10° to 30°C range.

Process Potential

Biological processes are, in general, the most cost-effective techniques for treating aqueous waste streams containing organic constituents. Moreover, biological processes have been

applied successfully at full scale to a wide variety of industrial wastes. Environmental impacts associated with biological processes are limited. Probably of greatest concern in this regard is the potential release of volatile organic compounds to the atmosphere as a result of aeration.

For biological decomposition of organic materials of a hazardous nature, many of which are toxic to microbial flora at high concentrations, it is necessary that the system be allowed to acclimate to the waste to be treated prior to routine operation of the process.

The activated sludge process, in one of its modifications, appears to have the greatest potential for the application of interest because it can be controlled to the greatest extent and best lends itself to the development of an acclimated culture. However, anaerobic filtration because of ease of operation, minimal sludge production, and energy efficiencies merits consideration in many situations. Thus, biological treatment is judged to be a viable technology which must be considered for treatment of hazardous aqueous wastes containing organic constituents.

CARBON ADSORPTION

Process Description

Activated carbon removes materials from water by the process of adsorption. Since adsorption is a surface phenomenon, the very large surface area associated with activated carbon, typically $500-1400 \text{ m}^2/\text{g}$, makes it a very effective adsorbent.

Pores, created during the activation process, exist throughout the carbon particles and account for the very high surface-to-size ratio. The greatest portion of this surface area is contributed by pores of molecular dimensions. Thus, pore structure in addition to surface area is a major factor in the adsorption process. Pore size distribution defines the size distribution of molecules which can enter the carbon particle to be adsorbed. Therefore, the carbon adsorption process is dependent upon the physical characteristics of the carbon and the molecular size of the adsorbate.

For the most part, activated carbon surfaces are non-polar in nature. Thus, activated carbon will sorb most organic compounds to some extent but is most effective for the least polar and least soluble organic compounds. Inorganic electrolytes are not sorbed effectively.

Other factors which affect the adsorption process include the characteristics of the liquid phase (e.g., pH and temperature) and the contact time between the liquids and the carbon adsorbent. The adsorption process is reversible which makes possible regeneration and reuse of activated carbons in many situations. Thermal regeneration is the most commonly used approach.

Activated carbon can be employed either in a granular or powdered state to effect treatment of wastewaters. Powdered carbon treatment usually involves suspension of the carbon in the wastewater in a stirred container and subsequent separation of the carbon-wastewater via a sedimentation process. Potential advantages associated with powdered activated carbon include:

- the cost of powdered carbon on a per pound basis is less than that of granular carbon,
- powdered carbon will equilibrate with the wastewater in a fraction of the time required by granular carbon,
- powdered carbon is easily slurried and transported, and can be supplied on demand by metering pumps,
- powdered carbon dosage can be rapidly changed to accomodate varying feed organic strength, and
- powdered carbon system requires a fraction of the carbon inventory required by granular carbon systems.

Development of powdered activated carbon technology has lagged behind that of granular carbon primarily as a result of lack of efficient regeneration systems. In addition, powdered activated carbon is sometimes difficult to separate from suspension and larger doses may be required than for granular systems achieving the same level of treatment.

Granular carbon applications are by far the more common. In this mode, the carbon is contained in a column or bed and the wastewater is passed through the contactor. After the capacity of the carbon bed is exhausted, the carbon may be removed and regenerated. Commonly, regeneration is accomplished by dewatering the carbon and then heating to a temperature of 815-925°C to volatilize and combust the adsorbed material.

One other treatment method involving the use of activated carbon exists. This technique involves the addition of powdered activated carbon to the mixed liquor in an activated sludge aeration basin to effect improvement in pollutant removal. Thus, this approach is a combined biological-carbon process. Regeneration of the carbon may be accomplished by thermal or wet oxidation techniques.

Process Applications

Activated carbon technology has been used for municipal

water purification, municipal wastewater treatment, industrial wastewater treatment, sugar decolorization, and purification of fats, oils, foods, beverages and pharmaceuticals. Recently, activated carbon has been used successfully in several emergency hazardous material spill response operations.

Contaminants removed in municipal and industrial wastewater treatment operations include BOD, COD, TOC, color, chlorophenols, cresol, cyanide, insecticides, phenol, polyethers, polynitrophenol, p-nitrophenol, p-chlorobenzene, resorcinol, TNT, toluene, xylene, and other organic chemicals.

Process Potential

Activated carbon adsorption is a well developed technology which has a wide range of potential waste treatment applications. It is especially well suited for the removal of mixed organic contaminants from aqueous wastes. Numerous examples of full scale waste treatment applications exist.

No serious environmental impacts are associated with carbon systems employing regeneration. If regeneration is not carried out, impacts could result from the disposal of carbon contaminated with hazardous materials.

Energy requirements for systems employing thermal reactivation could be significant - approximately 14,000-18,600 kJ/kg of carbon (6,000-8,000 Btu per pound).

Unit costs for carbon adsorption can vary widely depending upon the waste to be treated, the adsorption system, and the regeneration technique. It has been shown to be an economical approach in numerous instances.

In the current context, carbon adsorption must be considered a viable candidate for treatment of hazardous aqueous wastes containing organic contaminants. Granular activated carbon is the most well developed approach. However, combined biological-carbon systems appear promising for this application.

CATALYSIS

Process Description

A catalyst is an agent which accelerates the rate of a chemical reaction without itself being chemically altered at the end of the reaction. Catalysis, therefore, is not a process but rather is a means of enhancing any process which relies upon chemical transformations. In the current context, it is most applicable in improving the rate of chemical detoxification and degradation reactions. Thus, catalysis does not represent a means of concentrating wastewater constituents and is not

considered a concentration technology.

Process Applications

Catalysts are used in a number of chemical reactions including oxidation, reduction, polymerization, hydration, dehydration, hydrolysis, isomerization, dehydrogenation, cracking, and others. Waste treatment applications have included:

- cyanide destruction with activated carbon and copper catalysts
- chlorinated organic pesticide destruction using metallic couples such as zinc/copper, iron/copper, and aluminum/ copper; and pesticide dechlorination using nickel catalysts
- catalytic oxidation of domestic wastewater with proprietary catalysts and aqueous organic wastes with copper chromate catalyst
- oxidation of sulfides with iron and copper catalysts
- oxidation of aqueous phenolic wastes with Raney nickel catalysts and ferrous iron catalysts
- decomposition of sodium hypochlorite solutions with cobalt hydroxide catalyst
- isomerization of maleic acid into less water soluble fumaric acid with a hydrogen chloride or sulfuric acid catalyst.

Generally, catalysts are applied selectively based upon processes and pollutants of concern.

Costs of the catalyst are only a small part of the overall waste treatment process. Generally, the catalytic process permits lower temperature or pressure operation, therefore, capital and operating costs may actually be lower than the non-catalytic process. Costs are dependent upon the application with the only valid cost comparisons being between the catalyzed and non-catalyzed process.

Process Potential

Several potential applications of catalysis to waste treatment have been identified but commercial practicality has not been demonstrated.

Catalysts generally are very selective and, while potentially applicable to destruction or detoxification of a given

component of a complex waste stream, do not have broad spectrum applicability.

In view of the above and the fact that catalysis is not a concentration technology, it was dropped from further consideration in this study.

CENTRIFUGATION

Process Description

Centrifugation involves the application of centrifugal force to effect mechanical solid-liquid or liquid-liquid separation via sedimentation or filtration within the centrifuge vessel. Several types of centrifuges are available - - tubular, disc, conveyor bowl, vertical basket, conical screen, and pusher to name a few. Raw wastewater or sludge characteristics such as particle size and solids concentration as well as desired product consistency should be considered when selecting the appropriate centrifuge.

Process Applications

Centrifugation as a solid/liquid separation process generally is used to process dilute sludges consisting of 2-5% solids. Typically, a dewatered sludge of 15-50% solids can be produced, although drier cakes are possible. Applications include:

- removal of particles and pigment from lacquers, enamels, and dye pastes
- separation of microorganisms from fermentation broths and solvent extracts from antibiotic broths
- recovery of metal particles from film soap and spent catalysts, and deoiling of metal chips
- recovery of crystalline solids from brine solutions, and ethylenediamine liquors and acrylonitrile wastewaters.
- dewatering of waste sludges e.g., domestic wastewater and scrubber sludges, separation of acid sludges from acid treatment of petroleum stocks
- removal of meat tissue from animal fats and pulp skins, and seeds in food processing
- dewatering of oil/water separator bottoms

Centrifugation also has been applied to separate liquids of different densities. Typical applications include:

- separation of oil and water mixtures;
- clarification of oils, extracts, and food products; and
- separation of wash water from fats and oils in vegetable and fish oil refining and purifying.

Centrifugation has greatest applicability for the dewatering of sludges and slurries. It cannot provide solids removal from aqueous waste sufficient for direct discharge of the centrate. The process has several advantages:

- demonstrated operation;
- versatile;
- compact, self-contained process;
- reasonable cost, low operating labor requirements;
- capable of dewatering problem sludges with minimal chemical modification;
- minimal secondary air pollution effects; and
- compatible with waste recovery

Disadvantages include:

- incomplete treatment of aqueous wastes;
- treatment of centrate may be difficult;
- a non-selective and non-destructive physical process;
 and
- possibility of high maintenance requirement when abrasive materials are processed.

Costs are comparable with other sludge dewatering alternatives such as vacuum and pressure filtration and in typical instances range from \$22-50/tonne of dry solids (\$20-45/ton).

Process Potential

Centrifugation is a viable ancillary process for sludge dewatering in an overall wastewater processing train. It may also have limited application for separating liquids of different densities. Because its chief application would be as an ancillary process to support some primary concentration technique, it was not evaluated in detail.

CHEMICAL PRECIPITATION

Process Description

Chemical precipitation is a process whereby some or all of a dissolved substance is transformed into a solid phase as the result of a chemical reaction and is thereby removed from solution. Most common precipitation reactions involve the removal of inorganic ionic species from aqueous solution.

Precipitation is accomplished in wastewater treatment by adding appropriate chemicals to the solution and mixing rapidly. Once the chemicals are dispersed throughout the solution, precipitation reactions generally are very rapid. However, the particles formed may remain very small in which case additional treatment will be necessary to promote particle growth (flocculation) prior to separation of the solid and liquid phases.

Typically precipitation is accomplished by the addition of lime, sodium hydroxide, aluminum salts, iron salts, carbonates, or soluble sulfides. In some instances, oxidation of the waste may result in the precipitation of the oxidized species (e.g., iron). Choice of the chemical to be used is dependent upon the nature of the waste stream and the material to be removed.

Process Applications

Precipitation techniques primarily have been used to remove metals and certain anionic species such as phosphates, sulfates, and fluorides. Numerous industrial applications exist. Examples include treatment of wastes from iron and steel mills, aluminum manufacturing, copper smelting and refining, metal finishing, and inorganic chemicals industry. Species reported to be removed by precipitation reactions include arsenic, cadmium, chromium, copper, fluoride, lead, manganese, mercury, and nickel.

Process Potential

Precipitation processes have been in full scale operation for many years. The technique can be applied to almost any liquid waste stream containing a precipitable hazardous constituent. Required equipment is commercially available. Associated costs are relatively low and thus, precipitation can be applied to relatively large volumes of liquid wastes. Energy consumption also is relatively low.

Precipitation processes result in the production of a wet sludge which must be further processed prior to ultimate disposal. In some instances, the potential for material recovery from this sludge exists. However, very often, non-target materials are precipitated together with the material of interest

thus complicating or eliminating the feasibility of material recovery.

Usually, simple treatability studies must be carried out prior to applying the process to a waste stream to determine the chemical of choice, the degree of removal, and the required chemical dose.

In most instances, precipitation is considered to be the technique of choice for removal of heavy metals from aqueous hazardous wastes.

CRYSTALLIZATION

Process Description

Freeze crystallization is a technique which involves freezing an aqueous solution containing dissolved salts. Relatively pure ice crystals form and the salts are concentrated in the remaining brine solution. Ice crystals are mechanically separated from the brine, washed, and melted to yield fresh water. The remaining brine must be further treated or disposed of in some acceptable manner.

Basically, the process consists of: 1) heat exchange to cool the waste stream, 2) freezing using vacuum flash/vapor compression or secondary refrigerant freezing, 3) washing of the salts from the ice crystal, 4) melting of the ice to yield clean water, and 5) energy recovery to cool the incoming water and recover refrigerant. Major problems relate to the crystal/brine separation step and washing salt from the crystals. Also, because freeze point is influenced by waste stream composition, the process is very sensitive to fluctuations in waste stream composition. Difficulty has been experienced in making rapid operating adjustments to waste stream composition changes.

There are several claimed advantages to the process:

- because freezing is by direct contact with the refrigerant there is no heat transfer surface or membrane to be fouled
- at low temperatures, corrosion problems are minimized and less expensive materials can be used in construction
- volatiles can be separated from product water and condensed in the melting phase.

Process Applications

Demonstration scale testing of freeze crystallization has been carried out for desalination of seawater. However, only

limited laboratory scale testing of the process on industrial wastes has been accomplished. There are no commercial applications of the process. Since AVCO Corporation's efforts in the mid-1970's there has been little or no research conducted.

Industrial wastes which have been tested include:

- cooling tower blowdown
- electronics plant waste
- ammonium nitrate wastes
- weak sulfuric acid wastes
- pulp mill hot caustic extract
- solutions of acetic acid, methanol, and aromatic acids
- metal plating rinsewaters
- arsenal redwater

Unsuccessful attempts also have been made to treat sludges at eutectic temperatures.

Experimental studies have utilized waste streams ranging from 0.003% to 10% TDS. Dissolved metal ions, cyanides, and organics theoretically are treatable provided that the waste stream has a component that freezes. Work has not been attempted on fully organic waste streams.

Energy requirements for desalination are high when compared to membrane processes but lower than evaporation processes, the two major competing desalination techniques.

Process Potential

AVCO Corporation has stated that the inability of the process to respond to changing wastewater characteristics and its operational complexity were primary reasons for abandoning its research efforts.

Since this process has not been reduced to practice, there is no ongoing research and past efforts have not been successful, this process was judged to have little potential for the application of interest and thus was dropped from further consideration.

DENSITY SEPARATION

In the current context, density separation is construed to include the process of sedimentation and flotation because they are the most commonly used techniques for solids/liquids separation in wastewater treatment.

Process Description

Sedimentation

Sedimentation is a physical process whereby suspended solids are separated from the liquid phase as a result of gravitational and inertial forces. Essentially, the technique consists of providing sufficient time and space for solid particles to settle out of a liquid stream. Usually, this is accomplished in special tanks, chambers or ponds designed to provide the necessary time and quiescent conditions to allow solids to settle. A means for physically removing the settled solids as a slurry or sludge usually is provided.

Although sedimentation processes may be batch or continuous, continuous processes are the most common in wastewater treatment applications.

Sedimentation frequently is used in conjunction with chemical precipitation, coagulation, and flocculation processes.

Flotation

The term flotation describes the process of converting suspended, colloidal or emulsified substances to floating matter. This may be brought about by the introduction of minute air bubbles into the liquid phase. These air bubbles attach to the solid particles and the buoyant force of the combination is sufficient to cause the particles to rise to the surface where they form a floating layer which is removed by skimming.

Air bubbles may be formed either by injecting air into the liquid waste under pressure with a subsequent release of pressure to atmospheric, or by saturating the waste with air at atmospheric pressure and then subjecting the waste to a vacuum which causes the release of dissolved air in the form of fine bubbles. Coagulant aids are sometimes added to the waste to assist the agglomeration of solids.

As a solids removal method flotation has several advantages:

- both light solids and greases, as well as heavy solids may be removed in the same apparatus,
- the sludge formed is usually more easily handled, and
- the presence of relatively high concentrations of oxygen in the waste helps promote the oxidation of organic components of the waste.

On the other hand, there are several attendant disadvantages

of the process:

- both investment costs and operating costs are relatively high,
- because of the complexity of the equipment, maintenance costs are somewhat high, and
- the pressure type has high power requirements.

Process Applications

Sedimentation

Sedimentation has a long history of use in many applications. It is widely used in municipal and industrial water purification and wastewater treatment operations. Sedimentation is used in conjunction with chemical precipitation in all of the applications discussed under that topic.

Flotation

Flotation has been used successfully in the treatment of refinery wastes, food processing wastes, meat packing wastes, and paper manufacturing wastes. In general, its greatest application is to wastes containing oil or grease.

Process Potential

Sedimentation

Sedimentation processes have been in use for many years, are easy to operate, are low cost, and consume little energy. Required equipment is relatively simple and commercially available. The process can be applied to almost any liquid waste stream containing settleable material. It is considered to have high potential for the application of interest. However, it is an ancillary process which will be utilized primarily in conjunction with some other concentration technique such as chemical precipitation. Alternatively, it may be used as a pretreatment technique prior to another process such as carbon or resin adsorption.

Flotation

Flotation is a proven solids/liquids separation technique for certain industrial applications. It is characterized by higher operating costs and more skilled maintenance requirements than gravity sedimentation. Power requirements also are higher. This technique is judged to be potentially applicable but probably only in situations where the wastewater contains high concentrations of oil and grease.

DIALYSIS AND ELECTRODIALYSIS

Process Description

Dialysis is the transfer of small solute molecules in a waste stream through a semipermeable membrane into a wash stream flowing along the opposite side of the membrane. The transfer is driven by the concentration gradient between feed stream and wash stream. Factors controlling diffusion include membrane characteristics, membrane area, concentration gradient, and temperature. Membranes are capable of passing salts and small organic species while retaining colloids and higher molecular weight compounds. Dialysis treatment produces two output streams both being more dilute than the feed stream. The dialysate (treated feed stream) generally still will contain a higher concentration of solute than the diffusate (resulting wash stream). Thus, the process is of value in wastewater treatment if the dialysate or diffusate can be recovered and reused. process does not provide volume reduction.

Membranes may be tubular, flat sheets, or hollow fiber configurations of cellulosic or synthetic resin materials. Membrane evaluation factors include transfer rate, mechanical strength, durability, resistance to chemical degradation, thermal stability and cost.

Electrodialysis is similar to dialysis, however, a direct electric current is the driving force causing charged ions to pass through or be rejected by membranes which are either anion or cation permeable. Staging or alternate stacking of anion and cation permeable membranes separated by spacers results in feed stream separations into dilute and concentrated streams. By concentrating salts in the brine stream, the process provides volume reduction. Membranes are formulated of synthetic ion exchange resins cast or copolymerized in sheet form.

Process Applications

For dialysis to work, the concentration gradient must be large; therefore, the process is applicable only to waste streams with high concentrations of low molecular weight dissolved species. Caustics and mineral acids dialyze readily; however, to minimize potential membrane degradation, the membrane must be carefully selected. Industrial waste treatment applications have included separation of caustic soda from hemicellulose waste, separation of soluble impurities from spent acid electrolyte in electrolytic copper refining, recovery of sulfuric acid in several industries, and separation of salts from proteins and other biocolloids in pharmaceutical manufacturing.

The process has several disadvantages in hazardous waste

treatment including the need for pretreatment to minimize plugging, membrane erosion, and film or sludge formation on membrane surfaces; a low transfer rate; applicability only to concentrated waste streams; and generation of two dilute output streams.

Electrodialysis is applicable to aqueous wastes containing moderately high concentrations of inorganic salts (1000-5000 mg/l). The process can yield a brine stream containing up to 10,000 mg/l salt and a product stream of 100-500 mg/l salt. The most frequent application has been production of potable water from brackish water. It also has been used to concentrate seawater for salt production; to remove mineral constituents from organic process streams, e.g., the desalting of whey, de-ashing of sugars, and washing of photographic emulsions. Laboratory and pilot scale applications include treatment of secondary sewage effluent, acid mine drainage, demineralization of cooling waters, and treatment of plating liquors and rinses to salvage metals and acids.

An advantage of electrodialysis is that costs are moderate, but they are heavily dependent upon volume treated and amount of salt removed because of the fixed removal capabilities of a given stack of membranes. The process, however, has a limited range of applicability in terms of wastewater salts concentration and types of solutes which can be concentrated.

Process Potential

Neither process has been judged to have much applicability to aqueous hazardous waste treatment in the current context. They are not well suited to mixed constituent waste streams and both rely heavily on recovery and reuse of at least one product stream to offset costs. Dialysis should not be considered to be a concentration technology. Neither process was evaluated further.

DISTILLATION

Process Description

Distillation may be carried out in a variety of ways but usually involves boiling a mixture of liquids to produce a vapor that is rich in the lower boiling point components of the original mixture. The vapor may be condensed and recovered or recycled in part to the distillation system. Distillation can be carried out in a series of stages which in the limit can approach a complete separation of the components.

Distillation is expensive and energy intensive. It probably can be justified only in cases where valuable product recovery is feasible.

Process Applications

Distillation has wide industrial application in petroleum refining, organic chemical manufacture and purification, and solvent recovery. Treatment of waste by distillation is limited. The only hazardous waste materials which feasibly can be treated are liquid organics such as organic solvents and halogenated organics. Some specialized applications include:

- . waste oil re-refining,
- methylene chloride recovery,
- . ethylbenzene separation from styrene, and
- . waste solvent recovery.

Process Potential

Distillation is judged to have limited applicability to treatment of dilute aqueous hazardous wastes because of its high cost and energy requirements. Therefore, it has been dropped from further consideration.

EVAPORATION

Process Description

Evaporation is the process of vaporization of a liquid from a solution or slurry as the result of application of heat energy. It is applied in situations where one of the components of the system is not appreciably volatile. Products of evaporation are a relatively pure condensed solvent and a concentrate rich in the nonvolatile component.

Evaporation differs from distillation in that the vapor usually is a single component and even when it contains more than one component, no attempt is made to fractionate the vapor.

Usually, heat is supplied by condensing steam in a heat exchanger that is an integral part of the evaporation unit. Commonly, evaporation units are operated under some degree of vacuum to reduce the boiling temperature. Evaporation often is carried out in a series of stages or effects. Since large quantities of vapor are produced, it often is economically advantageous to use the vapor produced in one stage as the heating medium for a subsequent stage. Thus, multiple effect evaporation often is practiced.

Evaporation, usually, is not econmically feasible for solutions having a low solids content. Equipment costs are high and operating costs may become excessive for the concentration of very dilute solutions. Potential operational problems include salt buildup on heat exchange surfaces, foaming, and solids decomposition.

Process Applications

Evaporation is a proven, well-developed process which is utilized in some form by virtually every industry. Waste treatment applications include radioactive wastes, TNT wastes, photographic chemical dye wastes, paper mill wastes, molasses distillery wastes, and metal plating wastes. Often product recovery is associated with industrial waste treatment schemes which employ evaporation.

Process Potential

Evaporation is not expected to have broad application to the treatment of aqueous hazardous wastes containing moderately volatile organic constituents (BP 100°C-300°C). These organics cannot be easily separated in a pretreatment stripper and will appear in the condensate from the evaporator to some extent depending on their volatility. Therefore, good clean separation of these organics is not possible without post-treatment of the condensate.

The major disadvantages of evaporation are high capital and operating costs, and high energy requirements. This process is more adaptable to wastewaters with high concentrations of pollutants than to wastewaters with low concentrations of pollutants.

In view of the above, evaporation was dropped from further consideration for the application of interest.

FILTRATION

Process Description

Filtration is a process for separation of solids from fluids by passage of the fluids through a porous medium. The solids are retained by the filtering medium itself and/or by solids already trapped on the medium. The fluid may be gaseous or liquid but, in the current context, only liquid/solids separations are of interest.

Many types of commercially available filtration systems exist. An important factor in selection of the type of filtration system is the desired objective. If the intent is to produce a purified liquid stream, a different type of filtration system would be selected than if the objective was to concentrate the solids prior to subsequent processing or disposal.

Filtration systems may be classified according to the porous medium used. Generally used filter media fall into one of two classes: (1) granular media, and (2) flexible media.

Granular Media

Granular media filters usually consist of beds of sand or sand and coal within a basin or tank and supported by an underdrain system. Filtration is accomplished by passage of the waste stream through the bed. Through a complex process that may involve several mechanisms, particles are trapped on top of and within the bed. As this occurs, the porous nature of the bed is reduced thus, either reducing the filtration rate at constant pressure or increasing the amount of pressure required to maintain the filtration rate. At some point the filter must be removed from service and backwashed to scour the solids from the media. The spent backwash water containing the suspended solids must be collected and further treated or disposed in some manner.

Granular media filters primarily are used to produce a high quality water low in suspended solids. These systems cannot effectively filter liquids having high suspended solids concentrations because backwash frequently becomes excessive.

Flexible Media

Flexible media filters are characterized by the flow of a waste stream through a fine medium such as cloth or close mesh screen. Solids build up on the medium as a cake which then serves as the true medium for further filtration.

The flow through filters of this type is induced by a pressure difference across the filter media. One type of filter commonly used in the chemical industry is the plate-and-frame filter which consists of alternating hollow frames that serve to contain the retained filter cake. Pressed against this frame are plates which support the filter cloth and which are provided with drainage channels for carrying off the liquid filtrate. When the frames are completely filled with cake, the plates and frames are separated and the cake removed.

Leaf filters consist of cloth supported on thin hollow grids stacked in a cylindrical pressure vessel. Liquid filtrate passes through the cloth and is discharged through passages in the leaf units.

Several types of continuous filters are available which have the advantage of requiring much less labor for operation. Basically they involve the use of a rotating hollow drum covered with filter cloth supported by a screen backing. As the drum slowly rotates on its horizontal axis, the lower segments of the drum dip into a tank containing the slurry to be filtered. The piping is arranged so that a vacuum can be applied in the immersed section of the drum from the inside pulling the filtrate into the section and leaving cake on the outer surface of the drum. The vacuum will produce a partial dewatering of the cake.

At a suitable point in the drum rotation, the cake is scraped from the drum. Filter aids are sometimes added to the slurry to improve the filterability, provided that recovery of uncontaminated solids is not the prime objective of the operation.

These types of filters primarily are used for dewatering sludges rather than for producing a purified liquid.

Process Applications

Filtration is a process with a long history of use in numerous industrial processes, and municipal and industrial water and wastewater treatment operations. Often filtration is used in conjunction with precipitation, flocculation, and sedimentation processes to effect improved solids removal efficiency.

Filtration has been used as a polishing step following precipitation and sedimentation to remove arsenic, cadmium, chromium, lead, nickel, and zinc.

As a dewatering technique, filtration has been utilized extensively to dewater biological and chemical wastewater treatment sludges.

Process Potential

Filtration is a well developed process currently being used in a wide variety of applications. A wide spectrum of filtration systems are commercially available. The economics of filtration are reasonable for many applications. Energy requirements are relatively low and operational parameters are well defined. Therefore, filtration is judged to be a good candidate for the application of interest. However, it is not a primary treatment process but rather will be used to support other processes either as a polishing step subsequent to precipitation and sedimentation or as a dewatering process for sludges generated in other processes.

FLOCCULATION

Process Description

Flocculation, as used herein, is the process by which small particles suspended in a liquid are made to aggregate into larger particles which are more readily settled. Generally, flocculation is accomplished by the addition of chemicals to the suspension under a high degree of turbulence to effect rapid and thorough mixing. This rapid mixing is followed by a period of gentle stirring to promote particle growth.

Flocculating chemicals include alum, lime, iron salts, and organic polymers (polyelectrolytes). The inorganic flocculants

react to form large, fluffy precipitates or floc particles which act to enmesh small particles creating larger, more settleable particles.

Flocculation may be employed in situations where it is desired to remove suspended solids originally present in the wastewater or solids formed in a preceding precipitation process. For example, sulfide precipitation of some metals results in the formation of a relatively stable colloidal suspension. Alum and/or polyelectrolytes can be used to effect flocculation of the metal sulfide precipitates.

Flocculation usually is used in conjunction with precipitation and sedimentation. Indeed, many of the inorganic flocculants make use of precipitation reactions. Once the precipitate has been formed and the suspended particles have been flocculated, they can be separated from the liquid by sedimentation.

Process Applications

Flocculation has a long history of use in numerous municipal and industrial water and wastewater treatment applications. It has been used in conjunction with precipitation to remove arsenic, cadmium, calcium, chromium, copper, lead, magnesium, mercury, and nickel. In addition, it is used in in many water and wastewater treatment systems to remove suspended solids. Inasmuch as many pollutants such as pesticides and PCBs are often adsorbed to particulate matter in suspension, flocculation in conjunction with sedimentation can result in the removal of the associated pollutants.

Process Potential

Flocculation is a relatively simple process to operate and has been in use for many years. Necessary equipment is commercially available. Both costs and energy consumption are relatively low. The process can be applied to almost any aqueous waste stream containing precipitable and/or suspended material.

Flocculation must be carried out in conjunction with a solid/liquid separation process, ususally sedimentation. Often, flocculation is preceded by precipitation.

Flocculation is judged to be a viable candidate process for hazardous aqueous waste treatment, particularly where suspended solids and/or heavy metal removal is an objective. It may be used in conjunction with sedimentation as a pretreatment step prior to a subsequent process such as activated carbon adsorption.

In most instances, the applicability of the technique, the flocculating chemicals to be used, and the chemical dose can be judged based upon experience and simple laboratory treatability

tests.

ION EXCHANGE

Process Description

Ion exchange involves a reversible interchange of ions between an insoluble, solid salt (ion exchanger) and a solution of electrolyte in contact with the ion exchanger. Thus, in an ion exchange process, certain ionic species are removed from solution and replaced by ions of the same sign which are released from the exchange matrix.

Ion exchange materials may be natural minerals or zeolites, or may be snythetic substances specially prepared for specific properties. They generally contain a large number of soluble ionic functional groups at the surface. At these locations, the ion exchange reaction occurs. It is possible to alter selectivity of these materials towards inorganic and organic materials by altering the physical and chemical characteristics of the exchangers.

Commonly, ion exchange media are contained in columns or beds. Liquid which is relatively free of suspended solids is passed through the beds until the effluent concentration of the material which is being removed exceeds a desired value. At that point the exchanger must be regenerated. This is accomplished by passing a regenerant solution containing a high concentration of the ion originally associated with the exchanger through the bed. The exchanger thus is converted back to its original form and the pollutant, at elevated concentrations, is transferred to the regenerant solution. Used regenerant must be recovered for reuse by additional processing or disposed of in an acceptable manner. Usually, the bed is rinsed with a small volume of water to remove excess regenerant prior to the next service cycle.

Process Applications

Ion exchange can be used to remove both cations and anions. Because organic species frequently interact with the exchangers and cause operational problems, most applications of interest in the current context have involved inorganic species.

The ion exchange process has been used for many years to soften water. It also has a long history of use in industrial water purification.

Ion exchange is used extensively in the electroplating industry for treatment of rinse waters containing chromium, cyanide, and nickel. It also has been used as a polishing step in processes designed to treat aqueous metal finishing wastes. Other applications include:

- removal of iron, aluminum, and chromium III from chromic acid plating bath liquors,
- removal of aluminum from strong phosphoric acid/nitric acid solution,
- removal of various species from radioactive wastes, and
- removal of ammonia from biologically treated municipal wastewater.

Process Potential

Ion exchange is a proven process with a long history of use. It will remove dissolved salts, primarily inorganics, from aqueous solutions. For many applications, particularly where product recovery is possible, ion exchange is a relatively economical process. Also, it is characterized by low energy requirements.

Ion exchange is judged to have some potential for the application of interest in situations where it is necessary to remove dissolved inorganic species. However, other competing processes - precipitation, flocculation, and sedimentation - are more broadly applicable to mixed waste streams containing suspended solids, and a spectrum of organic and inorganic species. These competing processes also usually are more economical. Thus, the use of ion exchange probably would be limited to situations where a polishing step was required to remove an inorganic constituent which could not be reduced to satisfactory levels by preceding treatment processes. Therefore, while ion exchange is believed to have some potential for the current application, it is not a process which should receive primary consideration.

RESIN ADSORPTION

Process Description

Resin adsorption functions according to the same principles associated with carbon adsorption. That is, physical and chemical forces cause sorption of the solute onto the resin's surface. A major difference between resin and carbon adsorption is that because adsorption forces are weaker, resins can be chemically rather than thermally regenerated. This provides an opportunity to recover sorbed materials. Another difference is that, while activated carbon sorbs nonpolar compounds most readily, resin surfaces can be produced to be either hydrophobic or hydrophilic and thus be applicable to nonpolar or polar molecules.

Two basic types of synthetic resin adsorbents are available, polymeric and the newer carbonaceous. The polymeric adsorbents

are nonpolar with an affinity for nonpolar solutes in polar solvents or of intermediate polarity capable of sorbing nonpolar solutes from polar solvents and polar solutes from nonpolar solvents. Carbonaceous resins have a chemical composition which is intermediate between polymeric adsorbents and activated carbon and are available in a range of surface polarities. As with activated carbon, pore size distribution and surface area affect the sorption process. These characteristics vary for the different resins. Surface areas range from 100-700 m²/g, generally less than activated carbon.

Resins are used in manner comparable to granular activated carbon, i.e. in beds or columns with wastewater passed through the contactor. After sorbent capacity has been exhausted, spent sorbents generally are regenerated by steam, acid, caustic, or organic solvent (methanol, ethanol, acetone - - although it is highly flammable, isopropanol, and others) washing. Subsequent separation of the desorbed solute from the wash stream permits recovery of the solute. Credit for solute recovery may offset the severalfold higher initial cost of resins relative to carbon. Chemical regeneration also minimizes scale problems when waste streams high in inorganic solids are treated. It is claimed that resins, especially carbonaceous resins, have longer service lives than carbon because of greater resistance to attrition.

Process Applications

Resin sorption technology is not as well developed as carbon sorption and therefore, process applications are more limited. One application which has been examined is the treatment of munitions wastewaters primarily because solvent rather than thermal regeneration was more desirable. Other applications have included color removal from paper mill bleach effluents, dyestuff production plants, water supplies, and in the food and pharmaceutical industries; phenol removal and recovery; pesticide manufacture wastewater treatment; removal of organics in the production of ultra-pure water; removal of chlorinated hydrocarbons in vinyl chloride manufacturing; and removal of chlorinated hydrocarbons from contaminated groundwater. Laboratory studies have shown that phthalate esters, aldehydes and ketones, alcohols, chlorinated aromatics, aromatics, esters, amines, chlorinated alkanes and alkenes, and pesticides are adsorbable by resins. ins adsorbed certain amines and aromatics better than activated carbon did.

Resin adsorption has greatest applicability when:

- color due to organic molecules must be removed
- when solute recovery is practical or thermal regeneration is not practical

- · where selective adsorption is desired
- where low leakages are required
- where wastewaters contain high levels of dissolved inorganics

Process Potential

Because of selectivity, rapid adsorption kinetics, and chemical regenerability, resins have a wide range of potential applications. The primary disadvantage is high initial cost; although, this may be offset if recovery of the solute is practical. Costs for resins recently have been quoted to be \$11-33 per kg (\$5-15 per pound). While not economically competitive with carbon for high volume, high concentration, mixed constituent wastes, benefits may be gained by sequential resin and carbon adsorption.

Energy requirements are heavily dependent upon whether solute recovery from the wash media is practiced. Without solute recovery, energy costs account for 5% of operating costs; however, with solute recovery using distillation, energy costs could account for 50% of operating costs.

As with activated carbon, the only major environmental impacts relate to the regeneration process. If not reused, spent regenerant requires disposal, frequently by incineration or land disposal.

Resin sorption has been judged to be a viable candidate for treatment of hazardous aqueous organic wastes. The technology, however, has not been as well defined as carbon adsorption.

REVERSE OSMOSIS

Process Description

Reverse osmosis (RO) is a salt removal process which has been intensively developed over the past 15 years for treatment of both brackish water supplies and wastewaters.

A natural phenomenon known as osmosis occurs when solutions of two different concentrations are separated by a semi-permeable membrane such as cellophane. Water tends to pass through a semi-permeable membrane from the more dilute side to the more concentrated side, thus producing equal dissolved solids concentrations on both sides of the membrane. The ideal osmotic membrane permits passage of water molecules but prevents passage of ions such as sodium and chloride. For example, if a solution of sodium chloride in water is separated from pure water by means of a semi-permeable membrane, water will pass through the membrane in both directions, but it will pass more rapidly in the direction

of the salt solution. At equilibrium, the quantity of water passing in either direction is equal, and the pressure is defined as the osmotic pressure of the solution having that particular concentration of dissolved solids.

The magnitude of the osmotic pressure depends on the concentration of the salt solution and its temperature. By exerting pressure on the salt solution, the osmosis process can be reversed. When the pressure on the salt solution is greater than the osmotic pressure, fresh waster diffuses through the membrane in the opposite direction to normal osmotic flow--hence the name for the process, reverse osmosis.

Many materials have been studied for possible use as membranes for water and wastewater purification and related separation and concentration procedures. The most widely used membrane developed to date is simply a modified cellulose acetate film. Polyamide materials and polyarylsulfones are more recent developments.

The semi-permeable membrane acts to retain the ions such as sodium and chloride on the brackish water side, while permitting pure or nearly pure water to pass through the membrane. The properties of a membrane that permit water molecules to pass through but will not permit the flow of salt ions are not clearly understood. It is believed not to be simply a molecular filtering action even though individual water molecules are smaller than most of the ions of concern.

The water flux through the membrane is dependent upon the applied pressure, while the salt flux is not. As the pressure of the feed water is increased, the flow of water through the membrane should increase while the flow of salt remains essentially constant. It follows that both the quantity and the quality of the product water should increase with increased driving pressure.

Operating plants carry out the reverse osmosis principle in several different process designs and types of membrane configurations. There are four types of membrane systems which have been used:

- spiral wound,
- 2. hollow fine fiber,
- 3. tubular, and
- plate and frame module.

The first three types are in commercial production and are currently in use in operating plants. The plate and frame approach is not an efficient use of membrane surface area.

Membranes are susceptible to chemical attack and fouling,

and the flow systems are susceptible to plugging and erosion. Therefore, it is common to preprocess feed water to remove oxidizing materials, oils, greases, and particulates. Typical pretreatments include carbon adsorption, chlorination, pH control, and filtration as dictated by the nature of the feed water. Cellulose acetate membranes are typically operated at pressures of $2760-4140\ kN/m^2\ (400-600\ psi)$ to produce flux rates of 0.204-0.815 m³/m².d (5-20 gal/day/ft²).

Process Applications

Probably the most extensive use of reverse osmosis to date has been in the production of purified water from brackish or seawater. Other applications include preparation of rinse water for use in semiconductor and electronic manufacturing, and reclamation of chemicals and water from electroplating rinse waters. To a limited extent reverse osmosis has been used in treatment of sulfite pulping wastes, textile dying wastes, and pharmaceutical wastes.

Process Potential

Reverse osmosis is a relatively new process which has been reduced to practice for some applications. A number of competitive suppliers of reverse osmosis systems exist. Energy requirements for commercially available systems are about 7.61 x 10^6 – 9.51 x 10^6 J/m³ of product water (8-10 kwh/1000 gal). Reverse osmosis is a relatively costly process but it is capable of producing high purity water. The principal application is to concentration of dilute solutions of inorganic and some organic solutes.

The state of development of the process is such that it is necessary to conduct extensive bench and pilot scale testing prior to almost any potential application to ascertain feasibility.

Reverse osmosis in its present state of development is judged to have limited potential for the application of interest. Its use probably will be limited to polishing operations subsequent to other more conventional processes.

SOLVENT EXTRACTION

Process Description

Solvent extraction as used herein is the separation of constituents of a liquid solution by contact with a solvent that is immiscible with the liquid. Components of the original solution are transferred to the solvent for subsequent recovery or removal. Recovery and reuse of the solvent usually is dictated by economics. Unless the solvent has very low solubility in the original liquid, there will be solvent loss which, in addition to

increasing process cost, may cause unacceptable contamination.

A solvent extraction process usually involves effecting intimate contact between the feed and solvent phases by forced mixing or by countercurrent flow. Subsequent to mixing the two phases are separated and the solute is removed from the solvent by distillation, a second solvent extraction step, or some other technique. Solvent recovery from the treated feed stream also may be dictated by economics or discharge requirements.

Process Applications

The use of solvent extraction is limited. Commercial applications include manufacture of lubrication oil from crude oil, upgrading of gasoline, extraction of sulfur compounds from gasoline, refining vegetable oils and fats, and dehydration of acetic acid. The principal wastewater treatment application is removal of phenol and related compounds from petroleum refinery wastes, coke-oven liquors, and phenol resin plant effluents.

Process Potential

Solvent extraction is judged to have minimal potential for the application of interest. Broad spectrum sorbents such as activated carbon are expected to be more effective in treating dilute waste streams containing a diversity of organic compounds. Carbon adsorption also will be more economical unless a valuable product can be recovered which is unlikely in most cases expected to be encountered. Therefore, solvent extraction was dropped from further consideration.

STRIPPING

Process Description

Two types of stripping are possible: air and steam. Air stripping involves the passage of air through an aqueous stream to remove a volatile component. Steam stripping essentially is a fractional distillation of volatile compounds from a wastewater stream.

Although air stripping from tanks and ponds is possible, usually this process is carried out in packed towers. Typically, the liquid stream is introduced at the top of a packed tower and air is forced through the tower countercurrent to the liquid flow. Depending upon the component to be removed, both temperature and pH may be important variables in determining process effectiveness and efficiency. Air pollution control devices will be required unless it can be shown that direct emission of the air stream to the atmosphere has no adverse environmental impact.

Steam stripping usually is carried out in a packed tower or

conventional fractional distillation column with more than one stage of vapor/liquid contact. Preheated wastewater is introduced near the top of the column and flows countercurrent to the steam rising from the bottom of the column. The concentration of the volatile component in the liquid progressively decreases as the liquid passes down through the column. Wastewater at the bottom of the column is heated by the incoming steam. Heat recovered from the wastewater discharged from the bottom of the column is used to preheat the incoming feed.

Steam exiting the column is condensed and must be further processed for product recovery or disposed in an acceptable manner. Recycle of a portion of the condensed vapor to the stripping tower may or may not be practiced.

Process Applications

The only major application of air stripping is the removal of ammonia from domestic wastewater.

Steam stripping has been used primarily for ammonia, hydrogen sulfide and phenol removal from aqueous streams. Ammonia is removed by steam stripping for product recovery from coke oven gas scrubber water. Other recovery operations involving steam stripping include sulfur from refinery sour water and phenol from phenol production process water. Industrial waste treatment applications which have been reported include:

- . phenol removal from phenol plant effluent
- . removal of vinyl chloride monomer from suspension resins of polyvinyl chloride
- removal of methanol and sulfur compounds from Kraft mill condensates

Process Potential

Air stripping is judged to have minimal potential for the application of interest. The process would be difficult to optimize for hazardous aqueous waste streams containing a spectrum of volatile and non-volatile compounds. Air stripping does have appeal as a pretreatment prior to another process such as adsorption to extend the life of the sorbent by removing sorbable organic constituents. However, air pollution control requirements are likely to be severe thus making the economics less attractive. It should be noted that some air stripping of volatile components will occur during the course of any treatment process and may result in safety hazards or air quality problems. This is expected to be most severe in the case of biological treatment processes using aeration devices.

Steam stripping has merit for wastes containing high concentrations of highly volatile compounds. It is a proven process for some applications but will require laboratory and bench scale investigations prior to application to waste streams containing multiple organic compounds. Both energy requirements and costs are relatively high. By-product recovery to offset costs from the types of hazardous waste streams under consideration is unlikely.

For the application of interest, steam stripping is judged to have greatest potential as a pretreatment step to reduce the load of volatile compounds to a subsequent treatment process.

ULTRAFILTRATION

Process Description

Ultrafiltration as a method for removal of contaminants in wastewater is one of a number of processes employing semi-permeable membranes. Ultrafiltration differs from reverse osmosis in that ultrafiltration is not impeded by osmotic pressure and can be effected at low pressure differences of 34.5 to 1380 kN/m 2 (5 to 200 psi).

Ultrafiltration usually is applicable for separation of higher molecular weight (7500) organic materials ranging in size from about 100 angstroms upwards. The upper molecular weight limit for ultrafiltration is usually near 500,000. Above that molecular weight size, separation occurs by conventional microporous filtration.

The predominant mechanism in membrane ultrafiltration is selective sieving through pores of the membrane. Membrane rejection of a certain substance depends upon its molecular shape, size and flexibility as well as the operating conditions. A useful membrane must be able to effect separation distinctly at an economical flow rate.

Polycarbonate resins, substituted olefins and polyelectrolytic complexes have been employed among other polymers to form ultrafiltration membranes. Most ultrafiltration membranes on the market today are cellulose acetate or derivatives therefrom. This imposes some limitations on use. The pH range of the liquid must be between 4 and 9 and operating temperatures are restricted to less than 43°C to avoid hydrolysis of the cellulose acetate. Polyarysulfones and inorganic materials have been introduced to deal with high temperatures and pH values.

Typical membranes used in wastewater treatment are composed of an extremely thin surface layer or skin covering a porous substructure of the same material. The porous substrate is required for mechanical strength. Many times the membranes are reinforced with a nonwoven material such as paper to give added mechanical

strength.

A variety of configurations are available for use of these membranes in the ultrafiltration of wastewater. These include tubular units, plate and frame units, and spiral wound units. Most ultrafiltration systems are designed with similar flow patterns. A series-parallel layout is employed in which the dilute waste stream passes through several parallel membrane blocks or housings. This concept was developed to avoid the fouling inherent in direct onflowing systems. The typical design involves flow across the membrane face instead of directly onto it.

Ultrafiltration generally operates at lower total throughputs and considerably higher solute concentrations than reverse osmosis.

Process Applications

Ultrafiltration has been used primarily in small laboratory and industrial applications for product recovery or production of a highly purified solvent. Primary commercial applications of ultrafiltration include:

- · electropaint paint rejuvenation and rinse water recovery,
- protein recovery from cheese whey,
- metal machining oil emulsion treatment,
- textile sizing (PVA) waste treatment, and
- sterile water production for pharmaceuticals manufacturing.

Potential applications under development include dye waste treatment, pulp mill waste treatment, industrial laundry waste treatment, protein recovery from soy whey, and hot alkaline cleaner treatment.

Process Potential

Ultrafiltration is a commercially used process with several industrial applications. It is characterized by high capital and operating costs. Energy costs could run as high as 30% of direct operating costs.

Ultrafiltration is judged to have limited potential for the application of interest. Its use probably would be limited to relatively low volume streams containing substantial quantities of high molecular weight solutes or suspended materials. Pilot testing is a prerequisite to use.

SECTION 7

LITERATURE REVIEW

DESCRIPTION

This section describes and summarizes an extensive literature review which was undertaken as the second step in the technology screening process. In order to provide a consistent and coherent basis for comparing and evaluating various processes, a standard format was used to record data and observations gleaned from the literature:

- (1) Keywords
- (2) Reference (Literature Citation)
- (3) General Description
- (4) Organization and Location
- (5) State of Development

 - (A) Type of operation(B) Size of operation(c) Duration and frequency of operation
- (6) Influent Waste Characteristics
- (7) Process Ranges
 - (A) Application
 - (B) Operation
 - (C) Constraints
 - (D) Other limitations
- (8) Operations
 - (A) Performance data for major parameters(B) Equipment and supply requirements

 - (C) Energy requirements(D) Flexibility

 - (E) Reliability
- (9) Effectiveness
 - (A) Effluent quality and efficiency
 - (B) Form of Material
- (10) Process Economics and Costs
- (11) Environmental Acceptability
- (12) Pilot Plant Operations

Special effort was made to adhere to this format as closely as prudent engineering and scientific judgment allowed. However, it became apparent early in the effort that limited information on full or even pilot scale application of many of these

processes to hazardous aqueous streams existed. Rarely was information covering all the above items presented. This general absence of data was compounded by frequent reporting of technology performance using gross pollutant indicators such as COD or TOC rather than removal of specific pollutants.

Despite these problems, an extensive amount of pertinent literature was reviewed and summarized. In order to maximize the usefulness to the reader of the large quantity and wide diversity of information extracted from the literature, it is presented in several degrees of detail herein. The most detailed data summary is contained in Appendix C. Information in this Appendix is presented in a standarized tabular format arranged according to each candidate concentration technology. Data is further subdivided within each technology group on the basis of the previously described chemical classification system.

The second level of detail is presented in the form of a narrative literature summary and is contained in a subsequent portion of this report section. The organization of this summary description is similar to that of the tabular presentation contained in Appendix C. Primary emphasis has been placed on the ability of the several processes to treat chemical compounds in the various classes of interest. General descriptions of individual technologies are contained in the technology profiles and can be found in numerous standard texts. Rather than reiterate much of this basic information, this study instead builds upon it, expanding the discussion of technology application, especially treatment of specific chemical compounds.

The most concise summary of the literature information has been presented earlier in Table 1 which is arranged according to unit process and individual chemical. This is intended to provide quick reference on the treatability of a chemical by the various candidate processes. It also serves to illustrate information gaps. This table can serve as a tool in the decision making process to match a treatment process or processes with the waste stream of interest. Efforts to identify all potential processes thus would be greatly reduced. Evaluations and treatability studies, although probably still necessary, could be conducted in a less costly and time consuming manner. This is of special importance when actions must be taken rapidly to mitigate imminent hazards.

LITERATURE SUMMARY

The following summary has been prepared on the basis of information gained from the literature review and is arranged by concentration process. Additional details are contained in Appendix C.

Biological Treatment

A variety of biological processes are used for wastewater treatment. This review generally was limited to configurations of the activated sludge process, i.e., conventional, extended aeration, contact stabilization along with aerated lagoon treatment. Although not solely a concentration technology because pollutant degradation and transformation occur, chemicals are concentrated in and on the biomass via adsorption or metabolic processes.

Biological processes appear to be capable of treating numerous organic and inorganic pollutants, although only limited data on removal of hazardous compounds in full scale applications (56,81,100,101) are available. A report by Pajak, et al.(71) presents an extensive review of the effect of hazardous materials on biological treatment processes. Much of these data, however, reflect laboratory scale studies.

Alcohols

Removal of various alcohols by biological treatment generally was high even at concentrations up to 1000 mg/l. Several references on the activated sludge process reported reductions of 75-100% (56,81,101,133). Aerated lagoon treatment of alcohols achieved 38-85% reductions (100). Placak and Ruchhoft (103) stated that 24-38% of the removal resulted from oxidation and 52-66% by conversion into protoplasm. Several toxicity thresholds to sensitive aquatic organisms were presented by Lund (99).

Aliphatics

Biodegradation-efficiency of aliphatics spanned a wide range. Bess and Conway (100) observed zero to complete removals for various aliphatics by aerated lagoon treatment. Several references reporting on the activated sludge process cited generally high performances (56,81,90,101). Biodegradation of many of the aliphatics was based upon respirometer tests, theoretical

oxygen demands and toxicity thresholds (103,106,107,108,109,112).

Amines

Reported removal of amines was variable. Pitter (133) as reported by SCS Engineers (81) described several amines as readily biodegradable using acclimated activated sludge inocula. Melaney, et al. (107,108), however showed that many of these compounds, e.g., benzamide, benzidine, benzylamine, 2-fluorenamine and others, inhibited oxygen consumption. Alternative systems utilizing mutant bacteria were reported to completely degrade aniline and trichloroaniline (92), although Melaney (108) indicated that aniline inhibited oxygen uptake. Leipzig and Hakenburg (58) reported 99.9% reductions of nitroaniline using powdered activated carbon in an activated sludge system.

Aromatics

Wide variation in the treatability of aromatics has been reported. Bess and Conway (100) reported 10-100% reductions by aerated lagoon treatment. Some aromatics, e.g., mono, di, tri, tetra, and hexachlorobenzenes, were completely degraded by pseudomonas bacteria (66,92). Leipzig and Hakenburg (58) reported up to 96% reductions of nitrobenzene using powdered activated carbon in an activated sludge process. Pure activated sludge performances ranged from 50-100% (56,81,90,101). Dryden, et al. (90), however, stated that the compounds of this group are fairly biorefractory. This is supported by reports on general toxicity or inhibitory effects (102,106,108,109). Dryden, et al. (90) further suggested that achievable reductions attributed to biodegradation may be attained by air stripping or adsorption on the biomass.

Ethers

References relating to ethers all pertained to isopropyl ether. Activated sludge processes achieved 85-95% reductions (56,101). Bess and Conway (100) reported 70-90% removals by aerated lagoon treatment.

Halocarbons

Halocarbons generally are reported as biorefractory and inhibitory to biological growth (90). Several references, however, reported effective removals by biological treatment at concentrations up to 1.3 mg/l (21,58,65). Although not stated, these reductions may be attributable to the volatile nature of these compounds. Dryden, et al. (90) reported that halocarbons may not be detrimental to an activated sludge system since they tend to air strip readily.

Metals

Metals frequently were reported to inhibit biological activity (71,109,124,127). A review by Pajak, et al. (71) reported that bimetallic mixtures often were more toxic than the individual metals. Toxicity thresholds, however, appear to exist for many metals, e.g., barium, cadmium, chromium, manganese, and zinc. Toxicity thresholds varied from metal to metal ranging from 1 to 100 mg/l. At concentrations less than these thresholds, biological activity occasionally was reported to be stimulated (109,124). Reductions of 30-80% often were reported at concentrations ranging from 0.006-10 mg/l (118,122,128,132).

Pesticides

Except for herbicide orange (81) and 2,4,5-trichlorophenoxy-acetic acid (115), only slight biodegradation of pesticides was demonstrated (121). Wilkinson, et al. (92) presented half-life information for several of the pesticides using mutant pseudomonas bacteria.

Phenols

At concentrations of up to 500 mg/l almost complete reductions were demonstrated for most phenolic compounds, especially at contact times of 50 or more hours in acclimated systems. Several references reported greater than 70% reductions utilizing activated sludge processes (81,88,90,118). Leipzig and Hakenburg (58) showed 98.1% removals of nitrophenols using powdered activated carbon in an activated sludge system. Nathan (66) reported complete removals for several of the phenols employing mutant pseudomonas bacteria. Although toxic and inhibitory effects were noted for some compounds in the literature (109,124, 127), it appears that biological treatment can reduce even toxic compounds, e.g., 2,4,6-trichlorophenol, under suitable conditions (66,90,102,115).

Phthalates

Biological treatment was demonstrated to be effective in removing phthalate compounds. Removal efficiencies ranging from 50-100% were reported (21,81,90,100). Dryden, et al. (90), however, noted that a portion of these reductions may be attributable to absorption into cell tissue or air stripping.

Polynuclear Aromatics

All but two of the polynuclear aromatics were reported to be biorefractory or inhibited biological activity (108). Greater than 70% removal of napthalene at up to 500 mg/l and D-chloramphenicol were reported (56,81,100,101).

Chemical Coagulation

For purposes of this review, the category of chemical coagulation has been defined to include coagulation, flocculation, and precipitation. Additionally, filtration, sand or multimedia, often is included as an ancillary process. Typically, chemical coagulation has been used to remove inorganics, primarily heavy metals. Although most of the data examined pertained to metals removal, several documents report removal of organics with moderate success (6,21,90). Alum, aluminum sulfate, lime, and ferric chloride are the coagulants used most frequently.

Cohen (21) reports 15-56% removals of several aromatics, halocarbons, and phthalates at concentrations of 140-183 ppb using alum and dual media filtration. Dryden and Mayes (90) reported 60-90% reduction of phthalates at low ppb levels using aluminum sulfate. Becker and Wilson (6) reported 5-98% removal of several pesticides at low ppb levels using alum followed by sand filtration. Although reduction estimates were not provided, many polynuclear aromatics were reported to be removable by alum coagulation and gravity separation or sand filtration (90).

With regard to removal of metals by coagulation and filtration, reported reductions ranged from 0-100%. However, for each of the 22 metals for which quantitative reductions were reported, at least 30% removal was achieved with one of the coagulants enumerated earlier (16,34,63,64,90). Generally, arsenic, barium, beryllium, bismuth, cadmium, trivalent chromium, copper, iron, lead, manganese, mercury, nickel, silver, tin, titanium, vanadium, and zinc could be reduced by at least 90%.

Membrane Process - Reverse Osmosis

Reverse osmosis was shown to be less effective for separation of low molecular weight, polar organic compounds than for separation of inorganic salts. Two key criteria controlling separation are membrane characteristics and chemical nature of the molecule. Generally, separation of compounds with the same functional groups increased with increasing molecule size and branching. The following discussion illustrates the effectiveness of the various membranes.

Separation of alcohols ranged from 0-90%; cross-linked polyethylenimine (C-PEI) and aromatic polyamide (AP) membrane materials performed better than cellulose acetate (CA) (18,30).

Alcohols in order of decreasing percentage separation were i-propanol, ethanol, and methanol.

For aliphatics, separation varied considerably ranging from 0-90% (18,30). With the exception of dimethyl sulfoxide, aromatic polyamide and cross-linked polyethylenimine membranes performed better than cellulose acetate. Cellulose acetate membranes actually concentrated methyl acetate in the permeate. Trichloracetic acid demonstrated better separation than acetic acid by CA membranes but poorer separation by C-PEI membranes.

Similar results were reported for aniline. Using CA membranes aniline was concentrated in the permeate while C-PEI membranes achieved up to 80% removal (30).

Only limited data were available for aromatic compounds, however, C-PEI membranes again were superior providing 80-90% separation versus 3-7% for CA (18). Similar removals were observed for several ethers (18).

Separation of metal ions generally ranged from 85-100% at metal concentrations of 0.8 to 200 ppm and pH values of 8 to 11. Both CA and C-PEI membranes performed within this range (18).

Both CA and C-PEI membranes were capable of achieving 98-100% separations of numerous pesticides at concentrations of 42 to 1,579 ppb (18).

With CA membranes, concentration of phenol in the permeate was reported. However, 60-80% separation was reported for C-PEI membranes (18,30,90).

Membrane Process - Ultrafiltration

Very little information on the use of ultrafiltration for concentration of hazardous constituents in aqueous waste streams is available. The process has been applied efficiently in electropaint recovery, oil-emulsion waste treatment, and cheese whey separation. Molecules generally larger than 10^{-3} to 10^{-2} μ are retained in the concentrate stream (55).

In waste streams characterized by low suspended solids and high total dissolved solids, significant rejection of organic solutes was reported; e.g., 75% rejection of phenols at 100 mg/l (54) and 80-93% rejection of TOC in a 20 to 200 mg/l TNT containing wastewater (10). For the phenolic wastewater, rejection increased as pH increased with optimum rejection at pH 10, indicating that ionic state of the solute influenced rejection rate. Removal of metals in high suspended solids (125 to 1,550 mg/l) wastewater ranged from 79-89% at metal concentrations of 0.44 to 6.8 mg/l (59).

Stripping

Results of air and steam stripping experiments have been published for numerous organic compounds, particularly halocarbons. In other cases, certain compounds have been reported to be air or steam strippable even in the absence of experimental data because they possess relatively low boiling points.

As noted above, the majority of stripping data pertains to removal of halocarbons. A report by Coco, et al. (95) describes an extensive investigation on steam stripping of wastewaters from the petrochemical industry. Study conditions involved stripper feed flow rates of 250-325 ml/min, pollutant concentrations of 15 to 8,500 mg/l, overhead flows of 2-5% of feed flows, and various reflux flow to overhead flow ratios. Reductions of 75-99% between feed and bottoms were achieved at overhead flows of $\leq 5\%$ of feed flow. In some cases refluxing with a reflux to overhead flow ratio of 0.9:1 enhanced concentration of the pollutant in the overhead with lower levels in the bottoms. Greater than 99% reductions of 1,1,2-trichloroethane, 1,1,2,2-tetrachloroethane, 1,1,1,2-tetrachloroethane, perchloroethylene, chloroform, 1,1-dichloroethylene, and 1,2-dichloroethylene were reported. Up to 75% TOC removals were reported concurrently. Residual TOC was in the form of chloral (trichloroacetaldehyde).

Numerous chlorinated and nonhalogenated aromatics have been studied or reported to be strippable. Results indicate reductions of 50-99.9% (13,64,90). Moreover, phenol and chlorophenols were reported to be steam strippable while napthalene and acrylonitrile were reported to be air strippable (90).

Solvent Extraction

Solvent extraction has been shown to be a viable alternative to stripping and adsorption processes when recovery of a valuable product is possible. Advantages claimed for extraction are that less energy is required than for stripping and that, as opposed to adsorption processes, feed stream concentration has little effect on equipment size. Generally, the C_4 and C_5 hydrocarbons are the best volatile solvents with iso forms preferred because of lower water solubility. For phenolic compounds, dual-solvent extraction (polar solvent and volatile solvent in series or in mixed extractor) is most appropriate. To select an appropriate solvent for the solute in the wastewater, equilibrium distribution coefficients (K_D) , for solute/solvent pairs should be compared. The following criteria have been suggested by Earhart, et al. (27):

 If K_D values for important pollutants are >10, simple extraction with volatile hydrocarbon solvent is preferred.

- If K_D is <2, volatile solvent extraction is not recommended.
- Dual-solvent extraction is favored when K_D for both dual-solvent steps is ≥ 20 while K_D for direct volatile solvent is < 5.
- If best polar solvent gives a K_{D} not more than twice the K_{D} for a volatile solvent, dual-solvent extraction probably is not warranted.

Although removals achieved by extraction are dependent on the solute-solvent pair being tested, results indicate that for many organic pollutants, especially aromatics, halocarbons, and phenols 90-99% removals can be achieved (27,95). Many organics in the other pollutant classifications also are reported to be extractable although quantitative results are not available (90). In addition, 99% reduction of mercury by extraction with high molecular weight amines and quartenary salts was reported (90).

Sorption Process - Carbon Adsorption

Activated carbon adsorption has been the most extensively applied concentration technology. Yet, much of the published data reflects pure compound or synthetic wastewater laboratory testing. Full scale process applications, especially for industrial wastewaters, are numerous; however, for a variety of reasons treatment data are not available.

Generally, carbon adsorption is most effective for materials of high molecular weight and low water solubility, polarity, and degree of ionization. It is difficult, however, to accurately predict performance of a carbon sorption system based solely on properties of the solutes which are present. For example, multicomponent system studies have shown that preferential or competitive adsorption can reduce removals of some compounds to 50-60% of values predicted from pure compound studies (35,40).

Various approaches to regeneration of spent carbon are being investigated. Although thermal regeneration is practiced most frequently, regeneration by solvent desorption has been reported to have varying success dependent upon solvent used and solute being desorbed (20). This provides the potential extraction and recovery of the solute from the solvent.

The following sections describe treatability of compounds in each of the 13 chemical classifications (except for the miscellaneous pollutants class) by carbon sorption.

Alcohols

Sorption of alcohols varies substantially ranging from

about 3% to complete removal. Initial alcohol concentration greatly influenced removal; for example, at 100 $\mu g/l$ propanol was completely removed (20) while at 1000 ppm about 19% reduction was reported (35). Adsorbability was found to increase with molecular weight (35). For compounds of less than four carbons the order of decreasing adsorption was undissociated organic acids, aldehydes, esters, ketones, alcohols (when more than four carbons, alcohols moved ahead of esters), and glycols (35). Aromatics had greater adsorption than aliphatics (35). Desorption of alcohols from carbon by elutriating with various solvents ranged from 4-110% (20).

Aliphatics

Sorption of aliphatics varied widely ranging from complete removal to less than 10% removal. Giusti, et al. (35) reported that results of two component isotherm tests could be predicted from single compound tests; however, in four component tests, only about 60% of predicted adsorption occurred. Continuous flow column studies produced 60-80% of theoretical isotherm capacity. Chriswell, et al. (20) reported that carbon was found to be more efficient for sorption of alkanes and chlorinated alkanes and alkenes, while resin sorption was more efficient for esters, alcohols, phthalate esters, phenols, chlorinated aromatic compounds, aromatics, amines, and pesticides. Neither were efficient for carboxylic acids. Using several solvents, Chriswell, et al. (20) reported generally less than 10% desorption of compounds from carbon with the exception of esters (insoluble in water and soluble in alcohols and ethers) where 35-71% desorption was observed.

Amines

Complete removal of all amines at 100 μ g/l concentrations was reported (20). At 1000 ppm concentrations, removal ranged from 7.2-80.2% (35). Chriswell, et al. (20) reported that carbon was found to be more efficient for alkanes and chlorinated alkanes, while a resin was more efficient for esters, alcohols, phthalate esters, phenols, chlorinated aromatic compounds, aromatics, amines, and pesticides. Using several solvents, Chriswell, et al. (20) reported widely varying efficiencies in desorbing amine compounds from carbon. Desorption ranged from 0-82%, although, for 9 of 12 amines desorption was less than 38%.

Aromatics

Aromatics were reported to be sorbed better than undissociated organic acids, aldehydes, esters, ketones, alcohols, or glycols (35). Resins, however, were reported to sorb several aromatics more efficiently than did activated carbon (20). At concentrations of 0.1 to 6000 ppm, greater than 50% sorption was achieved for all aromatics reported with greater than 90% removal

for many compounds. Desorption from carbon with solvent elutriation was poor, never exceeding 15% (20). Preferential adsorption was reported in several studies (35,40) with only 50-60% of the adsorption predicted from single compound studies occurring in multi-compound tests.

Ethers

At concentrations of approximately 1000 ppm adsorption varied from 13.5-100% generally increasing with increased molecular weight and branching (35). Carbon adsorptive capacity ranged from 0.039 to 0.200 g. compound/g carbon at a sorbent dose of 5 g/l.

Halocarbons

Halocarbons in the concentration range 0.001-1000 mg/l were reported to be readily adsorbed by carbon. Removals of 75-100% frequently were reported. In several instances, halocarbon spills were treated successfully by an EPA mobile treatment trailer using carbon sorption technology (6). As with other chemical classes, sorption of halocarbons increased with molecular size (35). For many compounds, carbon sorption capacity was less in multi-component mixtures than in single compound solutions (21). Elutriation with solvents yielded 9-59% desorption of solutes from the carbon (20).

Metals

Carbon adsorption is not typically used for treatment of inorganics. McCarty, et al. (64) reported little or no reduction in arsenic, barium, cadmium, lead, manganese, and mercury and 35-80% removal of chromium, copper, iron, and zinc when initial metal concentrations were near analytical detection limits. At 100 mg/l concentrations, carbon doses of up to 10,000 mg/l yielded the following removals (72):

hexavalent chrome	36%
copper	96%
lead	93%
manganese	50%
mercury	99%
nickel	52%

PCBs

Activated carbon exhibits a strong affinity for PCBs. Contos, et al. (22) reported that concentrations of 1-160 ppb of Arochlor $\overline{1242}$ and 1254 were reduced to <1.0 ppb with carbon dosages of 4-100 mg/l. An EPA mobile activated carbon treatment trailer reduced PCB levels by 92.5-99.9% in wastewaters initially containing 1-400 ppb PCB levels (6).

Pesticides

Organic pesticides were removed effectively by carbon sorption. A publication by Becker and Wilson (6) cited several references of pesticide treatment with carbon. Reductions of greater than 80% were indicated with reductions frequently exceeding 99% at concentrations up to 4000 ppb. When TOC was used as an indicator of pesticide removal, TOC reductions of greater than 99% were reported at TOC values up to 10,000 mg/l (38). Sorption capacities for several pesticides were reported by Hager (38) and Bernardin, et al. (8).

Phenols

Carbon sorption is efficient for reduction of phenolic compounds. Removals of 92-100% at concentrations of 0.13-10 mg/l were attained by EPA's mobile treatment trailer (6). Chriswell, et al. (20) achieved virtually complete removal of various phenolic compounds at a concentration of 0.1 mg/l. Desorption from the carbon by elutriation with solvents proved ineffective. Illustrating several methods of pretreatment prior to carbon sorption, five full scale adsorption systems treating phenolic wastewater reported 83-99% TOC removals at TOC concentrations of 80-1,200 mg/l (38). Isotherm data including sorption capacities for several phenolic compounds were provided by Cohen (21).

Phthalates

Data on treatment of phthalate compounds is limited. Greater than 98% removal of bis(2-ethylhexyl) phthalate at 1,300 ppb was stated (5). Chriswell, et al. (20) reported complete removals of dibutylphthalate and dimethylphthalate at 100 ppb concentrations. Desorption by elutriation with solvents was poor. Activated carbon pretreatment improved phthalate removal by subsequent aluminum sulfate flocculation (90).

Polynuclear Aromatics

Adsorption of polynuclear aromatics is generally high. Chriswell, et al. (20) reported 80-100% reductions at 100 ppb concentrations. Poor desorption by elutriation with several solvents was indicated. Carbon used to further treat biologically and chemically treated wastewater achieved a 70% reduction of napthalene (64). Fochtman and Dobbs (31) provided isotherm kinetics for several polynuclear aromatics.

Sorption Process - Resin Adsorption

Generally, the principles which apply to carbon adsorption also apply to resin sorption. Major differences exist in initial cost of the sorbents and methods of regeneration. Carbonaceous and polymeric resins are severalfold more expensive than

carbon. However, for some compounds, e.g., trihalomethanes, greater adsorption capacity by the resins has been demonstrated (46), thus smaller quantities of sorbents are needed. Also, it is claimed that the carbonaceous synthetic resins offer greater attrition resistance and regeneration flexibility (135). Solvent desorption rather than thermal regeneration (often used for carbon regeneration) provides the potential for recovery and reuse of sorbed solutes. This could offset the higher initial cost of the sorbents.

Resin sorption technology for the application of interest is not as well developed as carbon sorption technology. Therefore, less information exists on the former technology. Treatability of compounds in 10 of the 13 chemical classifications is described below.

Alcohols

Polymeric resin Amberlite XAD-2 provided complete removal of several alcohols at 100 $\mu g/l$ concentrations (20). Desorption by elutriation with solvent varied from complete desorption to no desorption (20).

Aliphatics

Using a polymeric resin (Amberlite XAD-2) Chriswell, et al. (20) reported the adsorption of several chemical groups in order of decreasing sorbability, to be phthalate esters, aldehydes and ketones, alcohols, chlorinated aromatics, aromatics, esters, amines, chlorinated alkanes and alkenes, and pesticides. Sorption of aliphatics ranged from 25-100%. All but the chlorinated alkanes, chlorinated alkenes, and alkanes were removed better by the resin than by activated carbon. Acidic compounds were not sorbed well by either resin or carbon.

Desorption of aliphatics from resin by solvents ranged from little or no desorption to 50-72% for the esters (20).

Amines

As noted earlier, the adsorption of several chemical groups, in order of decreasing sorbability, was reported to be phthalate esters, aldehydes, and ketones, alcohols, chlorinated aromatics, aromatics, esters, amines, chlorinated alkanes and alkenes, and pesticides (20). Complete sorption of amines at 100 $\mu g/l$ concentrations was reported. Amines were removed better by the resin than by activated carbon.

Desorption of amines from resin by solvents ranged from little or no desorption to 100%. Six compounds demonstrated less than 50% desorption while six others showed greater than 50%.

Aromatics

Using a polymeric resin, Amberlite XAD-2, Chriswell, et al. (20) reported complete sorption of nine aromatics with no desorption to 80% desorption from resin by elutriation with solvent. The order of decreasing sorbability was reported to be phthalate esters, aldehydes and ketones, alcohols, chlorinated aromatics, aromatics, esters, amines, chlorinated alkanes and alkenes, and pesticides. Resins sorbed aromatics more efficiently than did activated carbon. Because of the ability to regenerate with solvents, resins were reported to be less costly for treatment of wastewaters containing TNT (2). However, for munitions wastewaters carbon was reported to have a greater absorption capacity (1b. nitrobodies adsorbed per lb. sorbent) than did Amberlite XAD-4 (40).

Halocarbons

Some of the resin adsorbents demonstrated a strong affinity for halocarbons. Using Amberlite XAD-2, Chriswell, et al. (20) showed complete removals of several halocarbons. Desorption of these compounds from XAD-2 by elutriation with a solvent ranged from 28-100% (20). Physical properties, equilibrium capacities, and results of a column study comparing carbonaceous resins to activated carbon were described by Tsacoff and Bittner (46). Their study indicates that per cubic foot, the resin treated approximately twice as much groundwater as did carbon before comparable breakthrough occurred.

PCBs

Arochlor 1254 at a concentration of 100 ppb was completely sorbed by Amberlite XAD-2 (20). Chriswell, et al. (20) also stated that 76.6% could be desorbed using the proper solvent. Lawrence and Tosine (57) studied the adsorption of PCBs from synthetic aqueous solution and raw sewage. They reported 60% reduction of Arochlor 1254 using Amberlite XAD-4 and about 23% reduction of Arochlor 1260 using Amberlite XAD-2 at PCB concentrations of 1-25 ppb.

Pesticides

Several case studies of resin adsorption of pesticides cited by Fox (32) reported at least 94% removal at pesticide concentrations ranging from 0.07 to 1,500 mg/l. Kennedy (49) showed Amberlite XAD-4 was more effective than activated carbon in treating a wastewater effluent from a pesticide manufacturer. At chlorinated pesticide concentrations ranging from 33-118 mg/l XAD-4 processed about four times more wastewater than carbon before comparable leakages occurred. Leakage could be maintained at <1 mg/l for at least 120 BV. While resin was readily regenerated with isopropanol, carbon was very poorly regenerated. It

was noted that acetone was a more effective regenerant, but that it is highly flammable. Amberlite XAD-2 achieved complete removals of several pesticides at 100 ppb concentrations with 10-49% desorption by solvent elutriation (20).

Phenols

Chriswell, et al. (20) reported complete reductions of several chlorinated phenolic compounds at 100 ppb levels using Amberlite XAD-2. Washing of the resin with various solvents yielded 35-76% desorption of the solutes. Crook, et al. (23) cited several case studies of industrial wastewater treatment by resin sorption. For initial phenolic compound concentrations of 280-6,700 ppm, sorption capacities of 16-87 g of solute/1 of resin were reported for Amberlite XAD-4 and XAD-7. Less than 1 mg/l phenol leakages were attained. Elutriation with methanol, ethanol, and acetone provided effective regeneration. The application of macroreticular ion exchange resins and polymeric adsorbents to waters containing 10-1,800 mg/l concentrations of phenols, dichlorophenol, and nitrophenol yielded greater than 99% removal (33).

Phthalates

Dibutyl phthalate, diethylhexyl phthalate and dimethyl phthalate were completely adsorbed at 100 ppb concentrations by Amberlite XAD-2 (20). Desorption of the solutes from XAD-2 by solvent elutriation ranged from 62-100%.

Polynuclear Aromatics

Amberlite XAD-2 resin completely removed several polynuclear aromatic compounds at initial concentrations of 100 ppb (20). Solute desorption from the resin by solvent elutriation ranged from 41-63%.

Sorption Process - Miscellaneous Adsorbents

In addition to carbon and the synthetic resins, other natural and synthetic sorbents have been studied. Limited data for two chemical classifications, metals and PCB's, have been reported. These are summarized below.

Metals

Dryden, et al. (90) reported on a variety of materials including silicon alloy, high clay soil, ground redwood bark, silicon oxide and calcium oxide slags, for sorption of metals. Copper and chromium at concentrations of 300 mg/l were completely sorbed on a high clay soil. Silicon alloy adsorption reduced 10-25 mg/l arsenic, cadmium, copper, lead, mercury, and zinc concentrations by >96%.

PCBs

Lawrence and Tosine (57) studied adsorption of PCBs from synthetic aqueous solutions and raw sewage. At concentrations of 1-25 ppb, Arochlor 1254 and 1260 were reduced 73% using PVC chips and 35% using polyurethane foams.

SECTION 8

PROCESS TRAINS

Since hazardous aqueous waste streams vary widely in composition and often contain a diversity of constituents, in general, no single unit process is capable of providing optimum treatment. Rather, arrangement of individual processes into process trains is necessary to achieve high levels of treatment in the most cost-effective manner. In this section the formulation of several process trains is discussed. Although not necessarily universally applicable, these process trains have been judged to be broadly applicable to many of the leachate, groundwater, and surface water quality problems identified.

Both selection of the unit processes based upon literature review results and formulation of process trains with broad applicability are described. Performance potential of each train was examined using three wastewaters of differing quality. This "desk-top" evaluation was conducted both independently and with input from representatives of companies marketing the technologies. Based upon these evaluations, priorities were established for subsequent laboratory bench scale evaluations using actual wastewater.

EVALUATION OF UNIT PROCESSES

Summary

Evaluation of candidate technologies led to the conclusion that the following unit processes have greatest broad range applicability to concentration of hazardous constituents of aqueous waste streams:

- biological treatment
- chemical coagulation
- carbon adsorption
- membrane processes
- resin adsorption
- stripping

These, however, must be supplemented with ancillary processes such as sedimentation and filtration.

Conclusions on all of the candidate technologies which led

to the selection of the above unit processes are summarized below.

- 1. Biological Treatment This process was found to be applicable to the treatment of a wide variety of waste streams. Concentration as well as degradation and stripping may occur during biological treatment. For several of the chemical classifications presented earlier, the following performances were observed:
 - a. alcohols generally removals of 75%-100% reported.
 - b. aliphatics wide range of efficiencies reported.
 - c. amines some amines were readily degradable with acclimated cultures while others were shown to inhibit oxygen consumption.
 - d. aromatics generally high removal reported; however, many compounds are biorefractory and removal may be due to air stripping or adsorption to biomass.
 - e. halocarbons generally reported to be biorefractory; removals attributed to biological treatment may be due to air stripping.
 - f. metals at below toxicity thresholds metal removals were reported; however, at higher concentrations toxic and inhibitory effects were noted.
 - g. pesticides no significant degradation reported.
 - h. phenols greater than 70% removals frequently were reported; toxicity effects also were noted.
 - phthalates high removals reported may be attributed to absorption into cell tissue or air stripping.
 - j. polynuclear aromatics generally reported to be inhibitory or biorefractory.
- 2. Carbon Adsorption Of the processes evaluated, the greatest amount of information on hazardous waste applications existed for carbon adsorption. Continuous flow systems using granular carbon in contact columns and powdered carbon in biological treatment systems as well as batch treatment for spill incidents have been reported. Generally, it was found that adsorbability increased with increasing molecular weight. For compounds of less than four carbons, the order of decreasing adsorption was undissociated organic acids, aldehydes, esters, ketones, alcohols (when greater than four carbons, alcohols moved ahead of esters), and gylcols. Aromatics had greater adsorption than aliphatics. Adsorption capacity for a specific compound is affected by other compounds present in the waste stream. Because of this

competitive adsorption, caution must be exercised in basing system design on case history results.

- 3. Catalysis Deemed to be a destruction or detoxification rather than a concentration process; found to be highly compound-specific and poor for mixed streams.
- 4. Centrifugation An ancillary process for concentration of high suspended solid waste streams.
- 5. Chemical Coagulation Numerous reports exist on the removal of heavy metals by chemical coagulation with lime, alum, iron and sulfides. At ppb levels, moderate removals (30%-65%) were reported for several aromatics, halocarbons, pesticides, phthalates, and polynuclear aromatics using alum.
- 6. Crystallization Process considered to be inapplicable. It is reported to be complex to operate and cannot readily handle variations in wastewater composition. There are no commerical operations and there has been very little research since the mid-1970's.
- 7. Density Separation an ancillary process applicable primarily to particulate or insoluble species; may be used with other chemical processes.
- 8. Dialysis/Electrodialysis Dialysis is most effective on feed streams with high concentrations of low molecular weight dissolved species. It is a low flux rate process with both output streams more dilute than the feed stream. Electrodialysis does not affect undissociated species, is complex to operate, can be fouled by high concentrations of organic compounds, and is applicable on streams with TDS levels of less than 5000 mg/l. Neither of these processes were deemed to have a high potential for the application of interest.
- 9. Dissolved Air Flotation An ancillary separation process which can be used in conjunction with chemical coagulation. The process frequently has been used for concentration of biological sludges and separation of oils in water.
- 10. Distillation Distillation is not expected to have broad application to mixed waste streams. Only when credits for recovered materials are considered does the process compete economically with other concentration techniques.
- ll. Evaporation Not expected to have broad application because the moderately volatile organics (boiling point of $100^{\circ}-300^{\circ}$ C) will appear in evaporator condensate. Good clean separations may not be possible without post-treatment. Energy usage is high and both capital, and operating and maintenance costs are high.

- 12. Filtration Ancillary process to remove particulates.
- 13. Ion Exchange Primarily for treatment of inorganic ions; however, heavy metals usually can be removed less expensively by other chemical-physical processes. Not considered to have a high potential.
- 14. Resin Adsorption Comparable in principle to carbon adsorption; however, resins usually are solvent regenerated. Using polymeric and carbonaceous resins, it is possible to adsorb a broader range of compounds than generally reported for carbon. It has been reported that resins were more efficient than carbon for removal of esters, alcohols, phthalate esters, phenols, chlorinated aromatics, aromatics, amines, and pesticides. However, results depend on the resins used. Experience with resin adsorption is much more limited than experience with activated carbon.
- 15. Reverse Osmosis Reverse osmosis (RO) is applicable to treatment of waste streams low in dissolved and suspended solids. It may be necessary to employ suspended solids removal processes prior to RO to remove particles of >25 μ . Performance is heavily dependent on membrane material and configuration. While typically applied to inorganics, up to 90% removal of a variety of organics has been reported. However, some membranes tend to concentrate some organics, e.g., aniline and phenol, in the permeate rather than concentrate stream. For the application of interest, RO probably would have to be paired with biological treatment or stripping for further treatment of the permeate stream. The concentrate stream also would need additional handling.
- 16. Solvent Extraction Potentially applicable when a single or a few reuseable compounds are present. Generally not suitable for waste streams containing a variety of organics at low part per million or part per billion concentrations.
- 17. Stripping Air and steam stripping have been used to remove numerous volatile, low molecular weight organics. Because stripping probably will remove predominantly biodegradable rather than refractory organic compounds, bottoms will require additional treatment possibly using an adsorption process. Although only limited data could be obtained, removal of aromatics, halocarbons, phenols, and polynuclear aromatics were reported to range from about 50% to 90%. Considered to be potentially applicable.
- 18. Ultrafiltration Whereas reverse osmosis can remove dissolved ionic species, UF basically is a filtration process capable of removing insoluble materials and organics of >1000 molecular weight. To date, applications have been largely in the area of waste paint recovery, protein recovery from cheese whey, and treating oil emulsions. Further processing of the permeate would be necessary. Judged to be of limited potential.

Discussion of Selected Processes

Biological Treatment

Biological treatment is expected to offer the most costeffective approach to removal of organic matter particularly biodegradable substances which are not amenable to sorption process-The major problem confronting the use of biological treatment is the potential presence of toxic organics and heavy metals which may interfere with metabolic processes and render this treatment approach ineffective. There are several categories of biological treatment processes including variations within these categories which overcome toxicity problems to some extent. addition, pretreatment or the addition of powdered activated carbon often can be applied successfully to overcome toxicity problems. For example, toxic heavy metal concentrations may be reduced below limiting concentrations by chemical coagulation, such as lime, alum, or iron precipitation, prior to biological Carbon sorption either by packed bed pretreatment or PAC addition to the biological treatment unit can be quite effective in dealing with toxic substances. Nutrient addition (e.g., phosphorous) will probably be required in many instances.

Biological treatment processes which can be used include activated sludge, trickling filters, aerated lagoons, and anaerobic filters. Each is discussed below.

Of the various activated sludge processes, completely mixed, extended aeration, and contact stabilization are used most often. The complete mixed configurations are more tolerant of toxic substances than plug flow schemes. The impact of toxic substances in the wastewater is reduced because complete mixing in the aeration unit reduces constituent concentration by dilution and distributes the load to a greater quantity of biomass. Non-biodegradable substances may pose more of a problem than biodegradable toxics especially if sorbed by the biological sludge where they may concentrate over a period of time and interfere with cell metabolism.

Sludge produced may be a hazardous waste due to the sorption and concentration of toxic substances contained in the wastewater. The quantity of sludge produced is normally governed by hydraulic detention time and sludge age. The conventional approach focuses on maximum sludge production consistent with the desired effluent quality. On the other hand, extended aeration aims to minimize sludge production at the expense of long detention times. Extended aeration typically is used in small operations since the small sludge handling requirements minimize the amount of manpower needed for operation (manpower costs are more significant than aeration costs for small units).

It is doubtful that activated sludge treatment alone will

suffice to meet discharge objectives in most instances. Pretreatment is expected to be needed not only to meet discharge requirements but also to remove toxic materials which would interfere with optimum performance of the biological system. Post-treatment normally serves to polish the effluent by removing refractory substances. These generally are expected to be in much lower concentrations than biodegradable substances. Listed below are potentially useful pretreatment steps:

- Chemical coagulation which can consist of lime, alum, or iron salt addition to form precipitates which scavenge toxic substances such as heavy metals from the wastewater.
- Carbon sorption which may either be accomplished through PAC addition with or without chemical coagulation or by packed beds of granular carbon. The objective is reduction of toxicants to facilitate biological treatment; therefore, large throughputs for packed beds or small PAC additions may be all that is required to achieve this reduction if the toxicants are strongly sorbed by the carbon.
- 3. Ultrafiltration or reverse osmosis are potential pretreatment candidates. These would be aimed at removing large molecular species which typically include the toxic and refractory species while smaller species which are generally biodegradable (e.g. ethanol, acetone) carry through and are removed in the biological unit.
- 4. Steam stripping may be useful in some instances but is more likely to remove a large fraction of biodegradable TOC than the refractory TOC.
- 5. Aeration, sedimentation and filtration may also be useful in some instances. For example, ferrous iron may be oxidized and precipitated to scavenge other heavy metals. Sedimentation with or without filtration could then remove the precipitated ferric hydroxide and reduce toxic heavy metals to acceptable levels.
- 6. Chemical oxidation, with ozone for example, may serve to detoxify certain materials; however, ozone consumption may be high due to oxidation of materials, such as ethanol, which are more appropriately biodegraded at much less cost.
- Wet air oxidation also may detoxify some organic substances but is expected to be a costly pretreatment step.
- 3. Ion exchange can remove toxic metal ions but is probably more expensive than chemical coagulation.

 Electrochemical treatment may be useful in some instances, e.g. it may be preferable to chlorination for reduction of high cyanide concentrations.

Candidate post-treatment steps include:

- 1. Carbon sorption has strong potential when teamed with biological. Biological treatment can substantially reduce the load to a carbon column and thereby minimize the cost.
- Resin sorption is an alternative to carbon sorption and may be less costly if steam and/or solvent regeneration are effective.
- 3. Chemical coagulation sedimentation filtration would be useful for removing residual heavy metals. Some PAC addition may also be performed to clean up low residuals of toxic organics.

Other steps, such as ion exchange, membrane processes, steam stripping, oxidation, are not considered to be good post-treatment candidate processes.

Trickling filters will not produce as high a quality effluent as activated sludge, but may be less troublesome from an operation standpoint. Pre- and post-treatment comments on activated sludge also apply to trickling filters.

Although generally effective, because of their large surface area, containing and collecting off-gases from aerated lagoons would pose a problem. Removal of hazardous sludge from the lagoon also may be a problem.

Anaerobic treatment may have advantages over aerobic treatment because of less off-gas and sludge production. These processes are less susceptible to upsets by many toxic substances such as heavy metals. Possibly, methane produced in the process could be used as fuel. Disadvantages include low quality and effluent necessitating further treatment and generally greater operational difficulty. Successful application of anaerobic treatment of leachate from municipal landfills has been reported on a bench scale level. Pre- and post-treatment considerations discussed for activated sludge also apply.

Chemical Coagulation

The term chemical coagulation as used here includes the processes of chemical addition, precipitation, flocculation, and sedimentation. Typically, it is a process used for the removal of particulate matter and inorganic ions, primarily heavy metals. Generally, precipitation is accomplished by adding alum, lime,

iron salts (ferric chloride, ferrous sulfate), or hydrogen or sodium sulfide. Organic polyelectrolytes also are used as floc-culants or to aid flocculation. A primary variable in determining coagulation chemical doses and removal efficiences is pH because of its effect on pollutant solubility in the wastewater solution. Although removals equal to solubility limits are theoretically possible, the formation of organometallic complexes and the incomplete removal of precipitated particles limits actual removal efficiencies.

When organics are present, post-treatment for organics removal will be required. This could take several forms including biological, sorption, or stripping. Reports indicate, however, that coagulation followed by efficient solids removal, e.g. dual media filtration can provide moderate removals (30-60%) of numerous organic compounds; even when these compounds are present at the low mg/l or ppb levels. Provisions also are required to manage sludges generated by the coagulation process.

Sorption Processes

Activated carbon sorption with packed beds is considered to be a prime candidate for leachate treatment. However, it is anticipated that activated carbon will be used in conjunction with other processes since it is quite expensive to treat moderate to high TOC loads with carbon alone. Furthermore, carbon is not effective for removing many highly soluble low-molecular weight organics. Although most of the low-molecular weight organics are not highly toxic, they will contribute substantially to the COD and BOD of the effluent. Carbon sorption is best suited for removal of refractory organics following biological treatment. These organics generally are adsorbed most strongly by the carbon and at the low concentrations typically found, the carbon sorption cycle can be lengthened. Consequently, the cost of carbon replacement or regeneration is lowered.

If the sorption unit is small, it is unlikely that on-site thermal regeneration of activated carbon will be performed. Instead, commercial replacement services probably would be used. Adsorption by synthetic polymeric and carbonaceous resins is an alternative to activated carbon sorption in some situations. There are, however, several major differences between the two types of sorbents:

- Acids, caustics, hot water, steam, and solvents (acetone, methanol, chloroform, methylene chloride, and mixtures) are used to regenerate spent resins. This permits recovery of desorbed solutes provided that:
 - there is a solute reuse potential,
 - costs for recovery of solute (and credits for recovery) and regeneration of solvent do not

exceed costs for disposal and replacement of spent solvent.

It also is useful where thermal regeneration is not possible, e.g., when nitrobodies from munitions waste are adsorbed; or high inorganic dissolved solids would result in scale formation during thermal regeneration;

- 2. Resin sorption kinetics are more rapid;
- 3. Resins generally have lower adsorption capacities;
- 4. Resins are more resistant than carbon to attrition losses;
- 5. Selective adsorption is possible by applying the proper resins in the correct sequence; and
- 6. Costs for resins range from \$11-33/kg (\$5-15 per pound) as compared to \$1.1/kg (\$0.50 per pound) for carbon.

At this time, there are limited full scale applications of the resin process. Phenol, pesticide, munition wastes, and contaminated groundwater have been successfully treated using various resins.

Alternative pretreatment steps for the sorption process include the following:

- Biological treatment (discussed earlier);
- 2. Solids removed by filtration;
- 3. Chemical coagulation for suspended solids and heavy metals removal followed by sedimentation alone or filtration alone, or a combination of sedimentation and filtration;
- 4. Aeration followed by sedimentation/filtration for oxidation and precipitation of dissolved iron which removes heavy metals as well as suspended solids. Aeration also may remove volatile organics to relieve loading on activated carbon;
- Ozonation to render organics more sorbable by carbon; and
- 6. Steam stripping may be effectively used for removing relatively high concentrations of volatile sorbable organics to reduce loading on carbon. It may be possible also to reduce concentrations of nonsorbable volatile

species such as lower alcohols, aldehydes, ketones, and perhaps acetic acid. The cost may be high however.

Processes such as ultrafiltration and reverse osmosis do not complement sorption and are not considered good pretreatment candidates. Ion exchange possibly may serve to remove ionic substances such as heavy metals, organic acids, amines, or cyanide; but it is likely that alternative processes will be less expensive.

Post-treatment processes which may be useful include the following:

- 1. Precipitation scavenging for removal of residual heavy metals.
- 3. Biological for removing biodegradable residuals.

Membrane Processes

Reverse osmosis and ultrafiltration are considered to be possible candidates for treatment of leachate and other contaminated waters. Reverse osmosis should be used only for waste streams relatively low in dissolved solids because treatment of highly mineralized water would result in a high volume concentrate stream. Ultrafiltration may be used on wastes high in dissolved solids because high molecular weight species are separated while dissolved salts pass into the permeate stream. This would be permissible in instances where the presence of dissolved salts in the process effluent is deemed acceptable.

It is anticipated that a suspended solids removal pretreatment step will be required in most instances to produce a clear feedwater to the membrane process. The membrane should remove the toxic and refractory species leaving biodegradable organics for post-treatment by a biological process. It is possible that steam stripping could serve to remove low molecular species prior to the membrane process to eliminate the need for biological processing.

Treatment of the brine or concentrate streams from the membrane unit must also be considered. Evaporation and incineration are potential treatment processes for these concentrates. Solidification also is an alternative.

Stripping Processes

Two types of stripping processes, air and steam stripping, are possible. Although both can be conducted in packed towers, steam stripping actually is a fractional distillation process with significantly greater energy demands. Typically, air stripping has been used for removal of ammonia from domestic wastewater. Steam stripping has been used for ammonia and hydrogen

sulfide removal from industrial wastes; soluble, low molecular weight volatile organics removal (e.g. methanol) from high BOD waste streams; and more recently, removal of water immiscible organics (e.g. chlorinated hydrocarbons) from process wastewaters. Because of potential for air pollution caused by removal of volatile organics, air stripping is deemed to have more limited utility. Steam stripping, however, may be more attractive, especially if:

- recovery of the pollutant in the condensate stream from a binary pollutant/water mixture is possible (practiced frequently in process wastewater treatment), or
- 2. pollutant load on downstream treatment processes can be reduced.

The efficiency of steam stripping is influenced by feedwater temperature and pH. Pretreatment steps would include pre-heating of feed concurrently with condensate or bottoms cooling and pH adjustment. The extent of pH adjustment necessary will be dependent on the pH of the waste stream and the pollutants present. Chemical coagulation and sedimentation or filtration for metals and suspended solids removal also may be required.

If recovery of a pollutant(s) in the condensate stream is not possible, two streams (condensate and bottoms), both needing additional treatment, are generated. If this is the case, steam stripping could be of marginal utility since cost-effectiveness is dramatically affected if there is no credit for recovered materials. Frequently, incineration will be the best disposal approach for organics-rich condensates. Condensate volumes ranging from 2-5% of feed flow have been reported. By refluxing a portion of the total condensate stream or the water phase if an organics-water separation occurs, the condensate can be further concentrated.

Bottom streams generally will be better suited to treatment by adsorption processes because stripping will remove the less sorbable soluble, volatile, low molecular weight organics. Refractory and the more biologically toxic organics probably will remain. Therefore, biological treatment will have less applicability. Treatment of bottoms by membrane processes also may be feasible although less attractive than sorption.

FORMULATION OF PROCESS TRAINS

Having identified the most promising unit concentration technologies, the next step was to formulate process trains which combined technologies in a fashion which would provide broad spectrum treatment capability. The objective was to identify process trains which would produce high quality effluents when applied to the wide range of waste stream compositions likely to

be encountered. Five such process trains incorporating the selected concentration technologies were formulated and are illustrated in Figures 1-5. Each of these process trains has particular strengths and weaknesses as discussed subsequently. One or more of these process trains should be applicable to almost any situation dictating concentration of a hazardous aqueous waste.

Process Train 1

Figure 1 illustrates a sequence of biological treatment followed by granular carbon sorption. This train is applicable to treatment of wastewaters high in TOC, low in toxic (to a biomass) organics, and containing refractory organics. Chemical coagulation and pH adjustment are provided for heavy metal removal and protection of the subsequent biological system. This may not be necessary if heavy metal concentrations are below toxicity thresholds and if the moderate removal efficiencies typical of activated sludge (20-60% although both poorer and better performances have been reported for some metals) are sufficient. When combined with additional metals removal by activated carbon or resin sorption sufficient removal may be achieved without chemical coagulation. Biological treatment such as activated sludge, rotating biological contactors, or anaerobic filters is included to reduce BOD as well as biodegradable toxic organics. This reduces the organics load to subsequent sorption processes. To prevent rapid head losses caused by accumulation of solids in the sorption columns, clarification and multi-media filtration are The intent is to reduce suspended solids to 25-50 mg/l. provided. Granular carbon adsorption is included to remove refractory organic residuals and toxic organics. Activated carbon rather than polymeric or carbonaceous resins has been specified because more full scale experience exists and performance as well as design and operating criteria have been reported. As previously noted, some concomitant removal of heavy metals also can be expected to occur. This process train is expected to be highly effective and the least costly. Its success, however, is dependent on biological system performance. Moreover, the presence of high concentration of volatile organic constituents may create a potential air contamination problem. Three by-product wastes are produced: chemical sludge, biological sludge, and spent carbon. Spent carbon can be regenerated but the sludge must be disposed.

Because the process is intended to handle multi-component waste streams, pollutant recovery for reuse is unlikely. The only potential for such recovery is during carbon regeneration if materials can be desorbed by steam or solvent washing. This would be reasonable only if a small number of separable compounds were sorbed on the carbon.

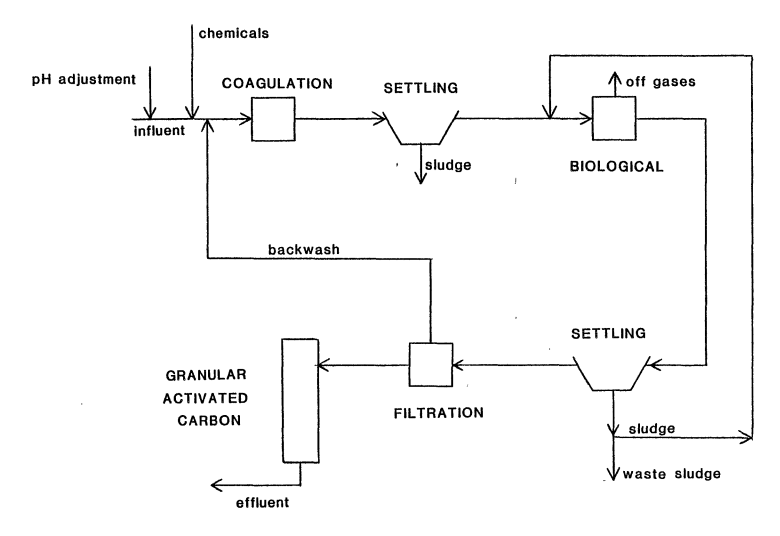


Figure 1. Schematic of biological/carbon sorption process train.

Process Train 2

The flowsheet depicted in Figure 2 employs the same unit processes as in Figure 1, but granular carbon is positioned ahead of biological treatment. This process train which also is applicable to high TOC wastewaters, was designed to respond to situations where waste stream components may be toxic to biological The rationale is to utilize the activated carbon to cultures. protect the biological system from toxicity problems. Therefore, the carbon would be allowed to "leak" relatively high concentrations of TOC (organics) rather than be operated to achieve maximum reduction of organic compounds. Allowable leakage would be based upon determination of the point which the carbon treated effluent becomes toxic to the subsequent biological process. Thus, the selection of the allowable TOC or organics leakage (i.e., breakthrough) from the carbon contactors is crucial to the performance and cost effectiveness of this process train. If biologically toxic organics are present, treatability studies must be conducted for several reasons, one of the primary being to establish the acceptable breakthrough level. Higher organic loads handled by the biological system result in greater service life of the granular carbon and consequently, lower costs related to the carbon treatment phase.

In this configuration, the chemical coagulation step (including settling and filtration) plays a role both in soluble inorganics removal and in particulate removal to minimize head losses in contact columns.

As with the process train in Figure 1, there is little potential for recovery of pollutants.

Process Train 3

The third process train, illustrated in Figure 3, utilizes biophysical treatment which is a combination of biological and powdered activated carbon treatments conducted simultaneously. This approach is simpler than the previously described sequential carbon-biological treatments and has the potential of achieving comparable effluent quality. Potential advantages include the use of less costly carbon (powdered vs. granular) and minimization of physical facilities required. Spent carbon-biological sludge can be regenerated or dewatered and disposed directly. However, if the latter approach is considered, it is necessary to include cost for disposal of toxics-laden carbon when making economic comparisons.

Complete mix activated sludge or contact stabilization are the two biological processes most frequently used. Recent reports suggest operating at long solids retention times (i.e. sludge ages of 100-150 days) and mixed liquor suspended solids concentrations of 20,000-25,000 mg/l with 60% being PAC and 40%

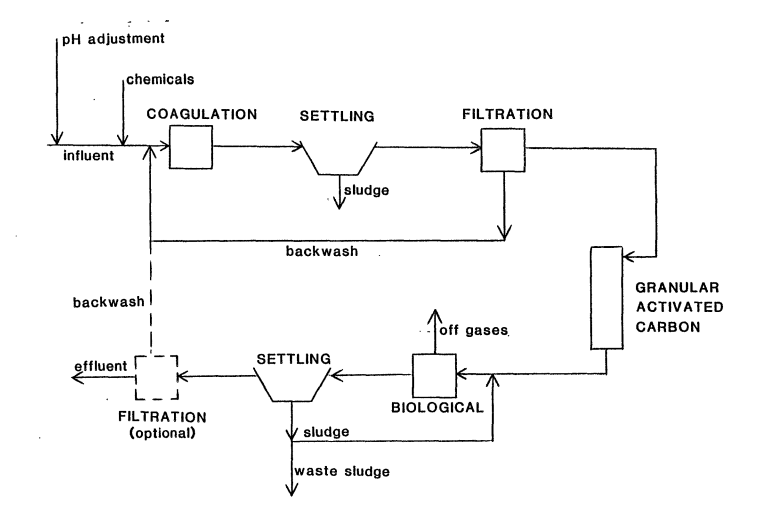


Figure 2. Schematic of carbon sorption/biological process train.

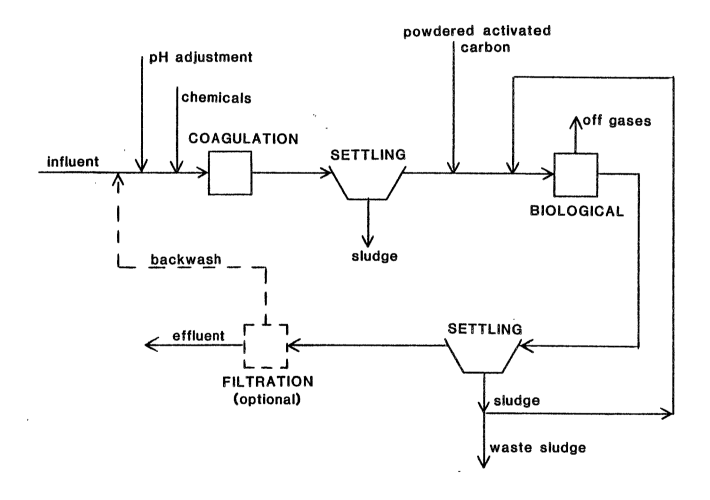


Figure 3. Schematic of biophysical process train.

being biomass.

Process Train 4

Figure 4 illustrates the use of a membrane process preceding biological treatment. This configuration would be applicable to wastewaters containing organic and inorganic pollutants. tion of the appropriate membrane process, ultrafiltration and/or reverse osmosis, would depend upon wastewater composition and treatment goals. Ultrafiltration is a membrane process capable of separating high molecular weight (mw > ~1000) species from a liquid stream on the basis of size. Reverse osmosis utilizes a semipermeable membrane to concentrate numerous dissolved species both organic and inorganic. Salinity is an important factor to be considered since UF will allow dissolved salts to enter the permeate stream while RO will not. Therefore, use of RO on high salinity waste streams is questionable because large volumes of concentrate are generated. Numerous RO membrane materials and configurations are available. Aromatic polyamide and crosslinked polyethylenimine materials have performed better than cellulose acetate. Membrane module configurations include hollow fiber, spiral wound, tubular, and flat sheet. Different configurations provide different surface areas, flux rates, flow velocities, and other process variables. Care must be exercised in selecting membrane materials and configurations. Organic removals of 20-70% have been reported for RO, although some membranes, e.g. cellulose acetate, tend to concentrate some organics, e.g. phenol and aniline, in the permeate stream.

A biological process was paired with the membrane process to address low molecular weight organics. Alternatively, stripping processes could be paired with membranes. Sorption processes were not considered in conjunction with membranes because of the likelihood that the lower molecular weight, readily soluble organics would pass through the system.

A major disadvantage of the process train depicted in Figure 4 is that membrane processes generate concentrate streams which require additional handling and disposal. The concentrate stream flow may be 10-20% of the feed flow.

Process Train 5

A processing system consisting of stripping and carbon adsorption is illustrated in Figure 5. This configuration will be applicable primarily to organic wastewaters although chemical coagulation for inorganics and particulate removal is provided. This flowsheet is suited to situations involving volatile and refractory or toxic organics. It is especially pertinent if a single or limited number of volatile compounds which can be recovered from the overhead condensate stream (if steam stripping is used) are present. Even though the wastewater may contain

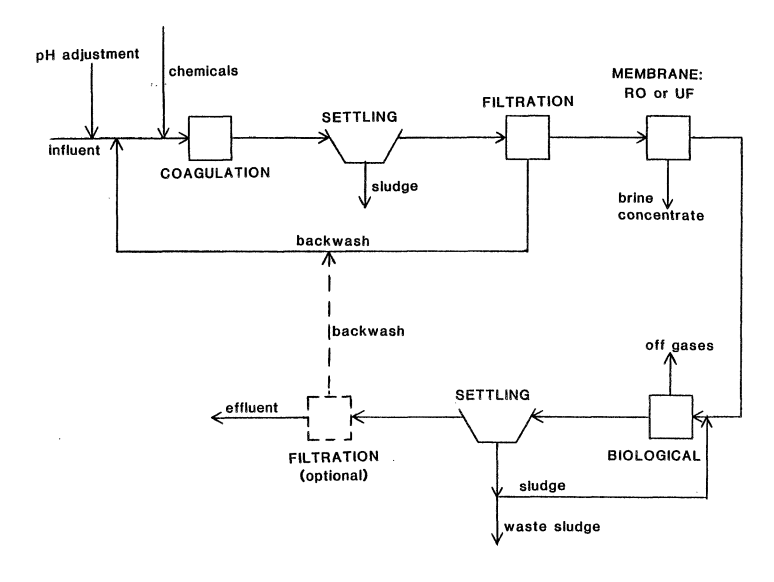


Figure 4. Schematic of membrane/biological process train.

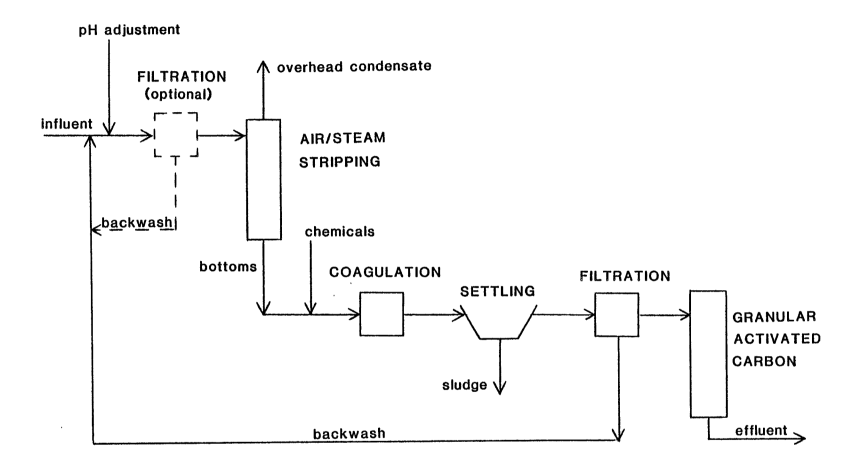


Figure 5. Schematic of stripping/carbon sorption process train.

air-strippable compounds, air stripping may not be the best selection if air pollution is of potential concern; unless off-gases can be contained and collected.

As previously discussed, stripping probably will remove biodegradable rather than refractory TOC. Therefore, it has been paired with activated carbon adsorption rather than a biological process.

Aside from pH adjustment prior to stripping, little pretreatment is necessary. If the wastewater contains readily settleable suspended solids, removal before packed column or tray tower steam stripping will prevent solids build-up in the stripping unit.

In addition to carbon treated effluent, this process train generates three waste streams: overhead condensate, chemical sludge, and spent carbon. Assuming that carbon will be regenerated, either on-site or by a commerical service, the two remaining streams require additional treatment and/or disposal. Preferably, the organic phase of the overhead condensate can be recovered and reused, with the water phase returned to the treatment system. However, if this is not possible, incineration is the best method for condensate disposal. Chemical sludge should be dewatered and disposed by a method commensurate with the materials contained in the sludge.

Process trains illustrated in Figures 1 through 5 do not represent the only possible configurations. They do, however, encompass the concentration technologies which are expected to have greatest broad range applicability and effectiveness. They also are the processes which have been demonstrated to some degree for treatment of hazardous aqueous wastewaters.

EVALUATION OF PROCESS TRAINS

Prior to initiating experimental studies, it was decided to perform a desktop evaluation of the five processes in an attempt to predict performance potential on actual hazardous waste streams. In order to select waste streams for this evaluation, a matrix was devised to group waste streams identified in Table B-l according to the concentration of inorganic and organic constituents. This matrix shown in Figure 6 describes the concentration of the inorganic and organic constituents as high, medium, and low. In general, the working definitions of "high", "medium" and "low" are as follows:

		INORGANICS CONCENTRATION		
		High	Medium	Low
ORGANICS CONCENTRATION	High	Sites 006 011	Site 010	Sites 001 002 003 005 021 023 024 025 026 027
	Medium	Site 022		Sites 008 009 013
	Low	Sites 004 012 014 015 016 018		

Figure 6. Waste stream categorization matrix.

	Hazardous Inorganic Constituent	Hazardous Organic Constituent
High	greater than 5 times water quality criteria*	greater than 400 ppb
Medium	<pre>from 2 to 5 times water quality criteria*</pre>	from 5 to 400 ppb
Low	<pre>less than water quality criteria*</pre>	less than 5 ppb

In addition, if a gross parameter such as BOD or TOC was reported in significant concentration (BOD >20 mg/l; TOC >10 mg/l), the waste stream was considered to fall in the high organic category. Although this system is not rigorous, it does permit a useful grouping of the actual waste streams. Inspection of the matrix revealed that most of the actual waste streams identfied fell into one of two categories: high organic-low inorganic or low organic-high inorganic. With regard to the latter category, concentration technology is essentially state-of-the-art. Therefore, the low organic-high inorganic category was not considered further.

Waste stream data from Site 026 in the high organic-low inorganic category was selected for the evaluation for several reasons: the data set was one of the most comprehensive available; ongoing activity at the site foretold future supplemental data availability; the state had assumed responsibility for mitigation of contaminated groundwater problems; no litigation was involved; the state regulatory agency was cooperative; and a strong possibility existed for use of the actual waste in subsequent laboratory studies.

The second waste stream composition selected for the analysis was that of Site 010 in the high organic-medium inorganic category. Reasons for selection were similar to those given for Site 026. In addition, heavy metals were present. Thus, this waste stream is sufficiently different than that of Site 026 to provide a second case.

The third waste stream utilized is a hypothetical leachate postulated on the basis of data contained in another report (137). Frequency of occurrence of the various classes of chemicals given previously also was considered in formulating the hypothetical leachate. The postulated leachate composition represents the high organic-high inorganic case. Reasons for selecting a hypothetical leachate include: (1) it provides a common basis

^{*} water quality criteria derived from Quality Criteria for Water, U.S. E.P.A., Washington, DC, July, 1976

for testing the appropriateness of various technologies, (2) it represents a reproducable "waste" composition for potential use in laboratory studies, (3) it contains a limited number of constituents representative of the broad range of materials found at actual sites, and (4) it is representative of "average" conditions at numerous sites.

Having selected waste streams for the evaluation, the next step entailed establishing effluent quality objectives for discharge to a receiving stream. Since established effluent limitation guidelines did not exist for the wastes of concern, the following procedure was utilized to define treatment objectives:

- Where published, industrial effluent limitation guideline documents specified a numerical criteria for constituents present in the waste of concern, these criteria were applied. Criteria generally were available for pH, BOD, COD, SS, oil and grease, phenol, cyanide, and several heavy metals.
- 2. When only Interim Primary Drinking Water Standards or numerical water quality based criteria were available, these were used, but the effluent objective was set at an order of magnitude greater than the water quality criteria. This allows for the impact of dilution and is consistent with a tenfold factor originally proposed in the RCRA regulations related to leachate quality. For parameters where this tenfold multiplier was applied, a maximum effluent limitation of 1 mg/l was established on the basis that existing technologies could achieve this level. Utilizing this approach, limits were developed for certain metals and several pesticides.
- 3. Subsequent to the application of items 1 and 2 above, only the priority and non-priority specific organic constituents remained. The following two approaches were used for these:
 - a. For non-priority organics, no effluent limitation was specified; the TOC limitation was the overriding limit.
 - b. For organic priority pollutants, 99.9% reduction was deemed to be achievable and desirable. If achieving 99.9% reduction required removal to less than current analytical limits of detection, the detection limit was specified as the effluent objective.

Quantitative data on the three waste streams of interest together with the defined effluent objectives are given in Tables 4-6.

TABLE 4 WASTEWATER CHARACTERIZATION - SITE 010

Parameter	Raw Wastewater Composition Range **	Effluent Quality Objective **
pH TOC SOC COD Oil & Grease SS TDS SO ₄ Sulfide Total P as P PO ₄ as P TKN NH ₄ -N NO ₃ -N NO ₂ -N	5.6 - 6.9 units 1800 - 4300 mg/1 4200 mg/1 5900 - 11,500 mg/1 90 mg/1 200 - 430 mg/1 15,700 mg/1 240 mg/1 <0.1 mg/1 <0.1 mg/1 5.4 mg/1 0.65 mg/1 <0.1 mg/1 <0.1 mg/1 <0.1 mg/1	5 - 9 units 20 mg/l 20 mg/l 50 mg/l 10 mg/l No increase 250 mg/l 0.3 mg/l 0.1 mg/l No limit 0.5 mg/l
Na Ca Cl Fe Hg* Pb* Sb* As* Cd*	1000 mg/l 2500 mg/l 9500 mg/l 31 - 330 mg/l <0.5 - <1 0.3 - 0.4 mg/l 2 130 11	No limit No limit No increase 1 mg/l 20 0.50 mg/l 200 500
Cr* Cu Ni* Se* Ag* Zn* CN* Hexachlorobutadiene* 1,2,4-Trichlorobenzene*	270 540 240 9 1 480 <0.01 mg/1 109 23	200 250 250 100 20 2 mg/l 0.25 mg/l 10 ³ reduction <0.09
Aldrin* Heptachlor* Phenol* Phenols (total)* 2,4-Dichlorophenols* Methylchloride* 1,1-Dichloroethylene* Chloroform* Trichloroethylene* Dibromochloromethane* 1,1,2,2-Tetrachloro-	<10 573 30 3.5 mg/l 10 180 28 ND - 4550 ND - 760 ND - 35	<1 500 NS <0.1 <0.4 <2.0 10 ³ reduction 10 ³ reduction <0.3
ethylene	ND - 1000	see TOC

TABLE 4 WASTEWATER CHARACTERIZATION - SITE 010 (cont.)

Parameter	Raw Wastewater Composition Range **	Effluent Quality Objective **
Chlorobenzene*	1200	10 ³ reduction
Methanol	42.4 mg/l	see TOC limitation
Ethanol	56.4 mg/l	see TOC limitation
Acetone	50.3 mg/l	see TOC limitation
Isopropyl alcohol	< 0.1 mg/1	see TOC limitation
Benzene*	ND - 3300	10 ³ reduction
Toluene*	ND - 31,000	10 ³ reduction
l,l,l-Trichloroethane*	ND - 225	< 2
Carbon tetrachloride*	92	< 4
Hexachlorocyclohexane-*		see TOC limitation
Alpha isomer	ND - 600	
Beta isomer	ND - 700	
Gamma isomer	ND - 600	
Delta isomer	ND - 120	

Footnotes:

^{* -} A priority pollutant

** - All concentrations in µg/l, except as noted

ND - Not Detected

TABLE 5 WASTEWATER CHARACTERIZATION - SITE 026

Parameter	Raw Wastewater Composition Range **	Effluent Quality Objective **
pH COD TOC NH ₃ -N Organic N Chloride Conductivity SS TDS	11.5 5400 mg/l 1500 mg/l 64 mg/l 110 mg/l 3800 mg/l 18,060 µmhos/CM 100 mg/l 12,000 mg/l	5 - 9 50 mg/l 20 mg/l 0.5 mg/l NL No increase NL 10 mg/l No increase
Volatile Organics:		
Vinyl chloride* Methylene chloride* 1,1-Dichloroethylene* 1,2-Dichloroethane* Benzene* 1,1,2-Trichloroethane* 1,1,2,2-Tetrachloroethar Toluene* Ethyl benzene* Chlorobenzene* Trichlorofluoromethane*	ne*<5 - 1590 <5 - 5850 <5 - 470 <5 - 78	10 ³ reduction 10 ³ reduction 20 ³ reduction
Acid Extractable Organic	cs:	
o-Chlorophenol* Phenol* o-sec-Butylphenol*** p-Isobutylanisol*** p-Acetonylanisol*** p-sec-Butylphenol*** p-2-oxo-n-Butylphenol m-Acetonylanisol*** Isoprophylphenol*** l-Ethylpropylphenol Dimethylphenol* Benzoic acid	<3 - 20 <3 - 33 <3 - 83 <3 - 86 <3 - 48 <3 - 1357 <3 - 1546 <3 - 8 <3 - 3 <3 - 8	0.09 0.5 mg/l see TOC limitation o.01 see TOC limitation

TABLE 5 WASTEWATER CHARACTERIZATION - SITE 026 (cont.)

Parameter	Raw Wastewater Composition Range	** Objective **
Base Extractable Organic	cs:	
Dichlorobenzene*	<10 - 172	10 ³ reduction
Dimethylaniline	<10 - 6940	see TOC limitation
m-Ethylaniline	<10 - 7640	see TOC limitation
1,2,4-Trichlorobenzene*	<10 - 28	0.09
Naphthalene*	<10 - 66	10 ³ reduction
Methylnapthalene	<10 - 290	see TOC limitation
Camphor	<10 - 7571	see TOC limitation
Chloroaniline	<10 - 86	see TOC limitation
Benzylamine or o-Toluid	ine<10 - 471	see TOC limitation
Phenanthrene* or	.7.0	303 3 !
Anthracene*	<10 - 670	10 ³ reduction

Footnotes:

* - A priority pollutant

** - All concentrations in µg/l except as noted

*** - Structure not validated by actual compound

NL - No effluent limitation

TABLE 6 WASTEWATER CHARACTERIZATION - SYNTHETIC LEACHATE

Parameter	Raw Wastewater Composition Range (mg/	Effluent Quality 1) Objective (mg/l)
TOC	500	20
BOD	1000	30
COD	1400	50
pн	5.0	5 - 9
Cl	285	No increase
NH ₄ ⁺	50	0.5
SS	50	10
TDS	350	No increase
Na	113	NL
Ca	110	NL
Mg	50	NL
K	10	NL
Fe ⁺²	10	1.0
Mn	1.0	1.0
As+ ⁵ *	20	0.5
Ва	2.0	1.0
-		
		_
Ca Mg K Fe ⁺² Mn As ⁺⁵ *	110 50 10 10 1.0 20	NL NL 1.0 1.0 0.5

Footnotes:
 * - A priority pollutant

Based upon the unit process performance data compiled from the literature, the performance potential of each of the five process trains was calculated for each of the waste streams. These calculations indicated that all of the process trains were potentially capable of meeting the established effluent quality objectives for stream discharge. However, because much of the available data were generated from single compound, laboratory scale studies, actual treatability of a multi-component wastewater cannot be accurately stated without conducting treatability studies using the actual wastewaters. This point was stressed by various company representatives marketing concentration technology treatment equipment/products. In general, vendors would not provide either performance estimates or process sizing and cost estimates (at +30% levels) without conducting treatability studies even though it would be expected that they would possess the best data for making these estimates. Thus, while the most promising concentration technologies and process trains can and have been identified, subsequent treatability studies are necessary to verify performance expectations, and to select the optimum process train for a particular situation.

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APPENDIX A

This appendix identifies entities queried with regard to data on hazardous aqueous waste problems, waste stream composition, and concentration technology applications and effectiveness.

TABLE A-1

ENTITIES CONTACTED

Location

Environmental Protection Agency

Region I
Region II
Region IV
Region IV
Region V
Region VII
Region IX
Region X
IERL
MERL
Office of Solid Wastes
Oil and Hazardous Materials Spills
Branch
National Enforcement Investigations
Center

Boston, MA
New York, NY
Philadelphia, PA
Atlanta, GA
Chicago, IL
Kansas City, MO
San Francisco, CA
Seattle, WA
Cincinnati, OH
Cincinnati, OH
Washington, DC

Edison, NJ

Denver, CO

Federal Agencies

U.S. Army Toxic and Hazardous
Materials Agency
Rocky Mountain Arsenal
Redstone Arsenal
Bureau of Reclamation
U.S. Geological Survey

Aberdeen, MD Commerce City, CO Huntsville, AL Boulder City, NV Menlo Park, CA

State Agencies

California Connecticut Georgia Illinois

Sacramento, CA
Hartford, CT
Atlanta, GA
Champaign, IL
(continued)

TABLE A-1 (continued)

State Agencies

Location

Kentucky
Louisiana
Maine
Massachusetts
Michigan
Minnesota
Nevada
New Jersey
New York
Ohio
Pennsylvania

Frankfort, KY Baton Rouge, LA Bangor, ME Boston, MA Lansing, MI St.Paul, MN Carson City, NV Trenton, NJ Albany, NY Columbus, OH Harrisburg, PA Pittsburgh, PA Norristown, PA Reading, PA Austin, TX Richmond, VA Charleston, WV

Texas Virginia West Virginia

Others

City of Niagara Falls
Gloucester County Planning Commission

Niagara Falls, NY Clayton, NJ

Companies

ABCOR
AMOCO
Calgon Corp.
Carborundum Co.
Chem-Bac Environmental Systems
FMC
ICI Americas Inc.
Ionics Inc.
Matlack Trucking Company
O & H Materials Inc.
Osmonics Inc.
Permutit Co.
Resources Conservation Company
Rohm & Haas
Westvaco

Wilmington, MA
Chicago, IL
Pittsburgh, PA
Niagara Falls, NY
Pittsburgh, PA
Princeton, NJ
Wilmington, DE
Watertown, MA
Swedesboro, NJ
Findlay, OH
Hopkins, MN
Paramus, NJ
Seattle, WA
Philadelphia, PA
Covington, VA

APPENDIX B

Appendix Table B-l contains data on identified hazardous waste problems and to the extent possible data on waste composition. A reference list which indicates data sources and pertains only to this table follows the main body of the table.

Problem sites are identified by a code number in Table B-1. The code numbers and associated problem sites are listed below.

Site Number	Site Description
001	Helevia Landfill adjacent to West Omerod water supply (near Allentown, PA)
002	Haverford, PA
003	Centre County, PA (near State College, PA)
004	Stringfellow Landfill, Riverside, CA
005	Rocky Mountain Arsenal, Commerce City, CO
006	Geological Reclamation Operations and Waste Systems, Inc. (GROWS) landfill, Falls Township, PA
007	Wade Site, Chester, PA
800	Bridgeport Quarry, Montgomery County, PA
009	Redstone Arsenal, Huntsville, AL
010	Love Canal, Niagara Falls, NY
011	LaBounty Dump Site, Charles City, IA
012	Saco Landfill, Saco, ME
013	Whitehouse, FL
014	near Myerstown, PA
015	Undisclosed
016	Necco Park, Niagara Falls, NY
017	FMC, Middleport, NY
018	Frontier Chemical Waste Process Inc., Pendleton, NY
019	102nd Street, Niagara Falls, NY
020	Pfohl Brothers, Buffalo, NY
021	Reilly Tar & Chemical Co., St.Louis Park, MN
022	Windham Landfill, Windham, CT
023	LiPari Landfill, Gloucester County, NJ
024	Kin-Buc Landfill, Middlesex County, NJ
025	South Brunswick, NJ
026	Ott/Story site, Muskegon County, MI
027	Hooker Chemical Co., Montague, MI

TABLE B-1
SUMMARY OF REPORTED WATER CONTAMINATION PROBLEMS

CONTAMINANT CLASSIFICATION				
Halocarbons	dustrial wastes at rate of 3,000 gal/wk; about 25 to 30% trichloroethylene (TCE)*. Materials percolated from excavated basin which now is under 50 to 60 ft of fill. Other wastes included ethyl acetate and phenols.			
	ļ	TCE* in ground water within plume - 191 to 260 mg/l		
		TCE* in ground water, ¹ / ₄ mi downgradient of site - 15 to 20 mg/l		
Phenols	002	Pentachlorophenol (PCP)* laden oil was deep well injected and later appeared in ground water and streams. EPA carbon trailer used to treat limited amount of contaminated ground water. PCP* in ground water a few hundred feet down gradient of	2,3	
		injection point - 2.4 mg/l		
Pesticides	003	Industrial waste containing Kepone and Mirex both spray irrigated and "Chemfixed" and placed in impoundments. Fixing held metals but promoted release of pesticides.	4	
Kepone in stream - 2 mg/l				
Metals Pesticides	004	Site`included impoundments for liquid industrial wastes and storage of solid industrial wastes. Acids, plating wastes,	5	
Misc.		and DDT were major materials disposed of although wide	(continued)	

TABLE B-1 (continued)

CONTAMINANT CLASSIFICATION	SITE CODE	PROBLEM DESCRIPTION AND WATER QUALITY	REFERENCE
Metals Pesticides Misc. (continued)		variety of materials went to site. Leachate known to exist. Soil and down stream surface water affected; area of ground-water contamination plume unknown. Surface water quality downstream of site (range): Cd* - 4.8 - 8.2 mg/l Cr* - 52 - 205 mg/l Cu* - 7 - 16 mg/l Mn - 340 - 550 mg/l Ni* - 28 - 48 mg/l Zn* - 77 - 115 mg/l pH - ~3	
Aliphatics Halocarbons Pesticides Polynuclear Aromatics Metals	005		
		A1 - 0.124 Ba - 0.1 Be* - 0.007 As* - 0.011 Bo - 0.624 Ca - 164	(continued)

TABLE B-1 (continued)

CLASSIFICATION	SITE	PROBLEM DESCRI	PTION AND WATER	QUALITY	REFERENCE
Aliphatics Halocarbons Pesticides Polynuclear Aromatics Metals (continued)	Co - 0.1 Cr* - 0.012 Cu* - 0.001 Fe - 0.090 Pb* - 0.001 Mg - 49.4 Mn - 1.04 Mo - 0.114 Ni* - 0.032 K - 6.83	Na - 378 Zn* - 0.024 Hg* - 0.0002	рн - 7.6 COD - 24.6		
Metals Misc.	006	Landfill accepts municipal ate with following average BOD - 10,900 COD - 18,600 SS - 1,040 TDS - 13,000 pH - 6.85 Alkalinity, as CaCO ₃ - 5, Hardness, as CaCO ₃ - 4, Ca - 818 Mg - 453 PO ₄ - 2.74 NH ₃ -N - 1000	ge quality is produ		7

144

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION AND WATER QUALITY		REFERENCE
Pesticides	009	Isomers of DDT present in surricide production facility. Effwaters. DDT* - ranged from 4.28 to 11.36 µg/l (over 3 months)	10	
Aromatics Halocarbons Metals Misc. Phenols	010	tive disposal site used by a contions in mg/l, except as noted pH - 5.6 - 6.9	Na - 1000	12 22 27 28
		COD - 5900 - 11,500 Oil & Grease - 90 SS - 200 - 430 TDS - 15,700 SO4 - 240	C1 - 9500 Fe - 31 - 330 Hg* - <0.0005 - <0.001 Pb* - 0.3 - 0.4 Sb* - 2 \mug/1** As* - 130 \mug/1** Cd* - 11 \mug/1** Cr* - 270 \mug/1** Cu* - 540 \mug/1** Ni* - 240 \mug/1** Se* - 9 \mug/1**	
		hexachlorobutadiene* 1,2,4-trichlorobenzene* aldrin* heptachlor*		(continued)

TABLE B-1 (continued)

Phenol*	CONTAMINANT CLASSIFICATION	SITE CODE	PROBLEM DESCRIPTION	AND WATER QUALITY	REFERENCE
sand filtration processes and prior to granular carbon adsorption	Halocarbons Metals Misc.	*	phenols (total)* 2,4-dichlorophenols* methyl chloride* 1,1-dichloroethylene chloroform* trichloroethylene* dibromochloromethane* 1,1,2,2-tetrachloroethyler chlorobenzene* methanol ethanol acetone isopropyl alcohol benzene* toluene* 1,1,1-trichloroethane* carbon tetrachloride* hexachlorocyclohexane alpha isomer beta isomer gamma isomer delta isomer ** denotes concentration follow sand filtration processes an	- 4.5** - 10 μg/1** - 180 μg/1** - 28 μg/1 - ND - 4550 μg/1 - ND - 760 μg/1 - ND - 35 μg/1 - ND - 1000 μg/1 - 1200 μg/1** - 42.4** - 56.4** - 50.3** - <0.1** - ND - 3300 μg/1 - ND - 225 μg/1 - ND - 225 μg/1 - 92 μg/1** - ND - 600 μg/1 - ND - 700 μg/1 - ND - 600 μg/1 - ND - 120 μg/1 - ND - 120 μg/1	

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION AND WATER QUALITY		REFERENCE	
Metals Aromatics Halocarbons Misc. Phenols Polynuclear Aromatics	011	lutants from municipal landfill manufacturer for disposal of pr data represents groundwater qualandfill and river which is dow	coundwater reported to be contaminated by migration of pol- tants from municipal landfill utilized by pharmaceutical nufacturer for disposal of production residues. Following ta represents groundwater quality at well located between ndfill and river which is downgradient. Other wells in ea and downstream also report contamination (concentrations μg/l, except as noted):		
		BOD - 2000 mg/l COD - 7100 mg/l TOC - 2300 mg/l TSS - <3 mg/l Total Phenols - 18 mg/l NH-N - 130 mg/l	Ba - Cu* - Hg* -	- 590 mg/l - 0.60 mg/l - 0.02 mg/l - 0.0048 mg/l - 0.17 mg/l	
		Volatile Organics:	range	average	
		benzene* chlorobenzene* 1,2-dichloroethene* trans-1,2-dichloroethene* dichloromethane* ethyl benzene* toluene* 1,1,1-trichloroethane* trichloromethane* trichloromethane* trichloroethane*	29 - 130 3.0 - 5.2 24 - 34	190 5.5 310 28 82 3.9 28 5.0 600 250 43 23	
		Neutral Extractible Organics:			
		aniline	140 - 870	410	(continued

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION	AND WATER QUAI	LITY	REFERENCE
Metals		Neutral Extractible Organics (c	continued):		
Aromatics			range	average	
Halocarbons		o-chloroaniline	ND - 360	140	
Misc.		p-chloronitrobenzene	460 - 940	720	
Phenols		chloronitrotoluene	ND - 460	120	
Polynuclear Aromatics		4-chloro-3-nitrobenzamide	440 - 8700	4200	
(continued)			890 - 30,000	8800	1
		2-ethylhexanal	ND - 4500	2600	
		1	9,000 - 23,000	22,000	
		3-heptanone	ND - 1300	640	-
			2,000 - 17,000	14,000	
ì		•	0,000 - 180,000	180,000	
		p-nitroaniline 32	2,000 - 47,000	37,000	1
		nitrobenzene*	ND - 740	250	1
		o-nitrophenol*	3,600 - 12,000	11,000	
		2-chlorophenol*	-	3	
		2,4-dinitrophenol*	-	99	
		n-nitrosodiphenylamine*			
		as diphenylamine		190	
		1,1-dichloroethylene*	-	P	
Metals Misc.	012	Following contaminants detected tannery sludge disposal area:	d in groundwater a	t well near	15
		Cr* - 1 mg/l average; 5 mg/l average; 6 mg/l average; 6 pH - 6.35 average; 6.0 mg/l	4.9 mg/l maximum		
				nggaga garaginan a mananan a mananan a ^{man}	(continue

TABLE B-1 (continued)

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION AND WATER QUALITY	REFERENCE
PCB's	013	Impoundments containing PCB contaminated oil and water were dewatered to eliminate threat of stream and groundwater pollution. Influent to powdered activated carbon treatment facility contained: Aroclor 1242* Aroclor 1254* ranged from 0.56 to 7.7 µg/l Aroclor 1260*	16
Metal	014	Groundwater contamination resulted from land disposal of arsenic compounds by pharmaceutical manufacturer. Prior to installation of groundwater purging and treatment system, arsenic* concentrations were 10,000 mg/l; after several years of purging concentrations of 10-30 mg/l remain.	17
Metal	015	Waste aresenic was disposed of in dump. Arsenic* concentrations found in groundwater were 175 mg/l.	17
Metals	016	Following contaminants found in groundwater near inactive chemical waste disposal site: Ba - 2000 mg/l Other inorganics and organics anticipated to be present.	18
Metals Pesticides	017	Arsenic* and Carbofuran found in surface runoff and in lagoon used by chemical manufacturer.	18
			(continued)

CONTAMINANT CLASSIFICATION	SITE CODE	PROBLEM	DESCRIPTION AND WATE	R QUALITY	REFERENCE
Metals Misc.	018	Following contaminants found in impoundment used by chemical waste processor (conc. in mg/l): Cd* - 1 Cu* - 9 Fe - 60 Ni* - 3			18
Metals Aromatics	019		ercury* and benzene hexachloride believed to be in ground- ater in vicinity of chemical manufacturing and waste dispos- l operations.		
Aromatics	020	Chlorinated benzenes found in leachate and groundwater in vicinity of waste disposal operation used by several chemical producers.			18
Phenol Polynuclear Aromatics	021	Following contaminants found in shallow groundwater in vicinity of chemical production facility: phenol* - 50 µg/l polynuclear aromatics - 3400 µg/l		. 19	
Metals Misc.	022	Following range of contaminants were found in ground and surface waters (ponds) in vicinity of municipal landfill which also accepted industrial wastes (conc. in mg/l): 3 worst case 2 worst case			20
		Pollutant Alkalinity PH	wells 20.6 - 300 6.27 - 6.5	81 - 156 6.22 - 6.3	(continued)

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESC	CRIPTION AND WATE	R QUALITY	REFERENCE
Metals Misc. (continued)		Pollutant TS TOC TKN Cl Na - mixed/settled Mn - mixed/settled Fe - mixed/settled Zn*- mixed/settled Cu*-mixed/settled Cu*-mixed/settled Cr*-mixed/settled Cr*-mixed/settled Cr*-mixed/settled Npecific conductance a - results reported NR - not reported	3 worst case wells 840 - 1730 12 - 39 <1 - 8.7 31.0 - 125 4.6 - 34.1/26.9 0.41 - 4.16/3.70 21.1 - 196/162 0.32 - 0.54/0.21 0.082 - 0.365/0.07 0.196 - 0.393/0.27 0.123 - 0.55/0.28 80 - 1200 ed for mixed sample and sample	1.03 / NR 3.38 / NR 0.07 / NR 6 0.006 / NR 1 0.003 / NR <0.001 / NR NR	
Metals Phenols Misc.	023	Following contaminants gradient of landfill w pharmaceutical wastes. poorest quality wells pH specific conductance temperature (°F) color sulfate total hardness Ca	hich accepted large Data represents qu	quantities of ality range at 3	21

CONTAMINANT CLASSIFICATION	SITE CODE	PROBLEM DESCRIPTION AND WATER	R QUALITY	REFERENCE
Metals Phenols Misc. (continued)		Na 13 - 130 CN* Se * 0.01 - 0.02 Pb* COD 168 - 9920 Cu* BOD5 42 - 4040 Hg* MBAS 0.24 Zn* Phenols* 0.008 - 54.17 Ag*	0.005 0.10 0.10 - 0.71 0.0005 0.36 - 26.8 0.01	
Aromatics Halocarbons PCB's Polynuclear Aromatics Phthalates	024	Following range of contaminants were detective from landfill accepting major proportions tion wastes (conc. in µg/l, except as noted aroclor 1254* Aroclor 1016*/1242* Aroclor 1016*/1242*/1254* benzene* biphenyl napthalene chlorobenzenes* camphene C4 alkyl cyclopentadiene C5 substituted cyclopentadiene dichlorobenzene* dichloroethane* dichloroethylene limonene methyl chloride* methyl napthalene parafins petroleum oil phthalates phthalate esters pinene	of chemical produc-	(continued)

TABLE B-1 (continued)

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION AND WATER QUALITY	REFERENCE
Aromatics Halocarbons PCB's Polynuclear Aromatics Phthalates (continued)		styrene tetrachloroethylene* toluene* trichloroethane* trichloroethylene* trichloroethylene* trimethylbenzenes MIBK xylene P to 590 P to 16,200 P to 490 P to 7700 P to 7700 P to 7700 P to 3300	
Halocarbons Misc.	025	Following contaminants were detected in groundwater in vicinity of municipal landfill due to "industrial waste seepage from landfill" (conc. in µg/l): 1,1,1-trichloroethane* 532 tetrachloroethylene* 187 1,1-dichloroethane* 2.3 1,2-dichloroethylene* 0.2 chloroform* 1.1 1,2-dichloroethane* 2.1 dibromochloromethane* 3.9 bromoform* 0.2 TOC 500	24
Halocarbons Aromatics Phenols Polynuclear Aromatics	026	Ground and surface waters were polluted by migration of contaminants from waste disposal lagoons and direct discharge practices attributed to chemical production facility. Following data describe groundwater quality range at four wells located within the groundwater contamination plums (conc. in $\mu g/1$):	25

TABLE B-1 (continued)

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION AND WATE	R QUA	LIT	Y	REFERENCE
Halocarbons		Volatile Organics:	······································		anning and a second	
Aromatics		vinyl chloride*	140	to	32,500	
Phenols		methylene chloride			6570	
Polynuclear Aromatics		1,1-dichloroethylene*			19,850	
(continued)		1,1-dichlorethane*			14,280	
		1,2-dichlorethane*			8150	
		benzene*			7370	
		1,1,2-trichloroethane*			790	
		1,1,2,2-tetrachloroethane*			1590	
		toluene*			5850	
[ethylbenzene*			470	
		chlorobenzene*		to		
		trichlorofluoromethane*		to		
	l i	Acid Extractable Organics:				
		o-chlorophenol*	<3	to	20	
		pheno1*	<3	to	33	
		o-sec-butylphenol	<3	to	83	
		n-isobutylanisol~	<3	to	86	
		or p-acetonylanisol ^b				
		p-sec-butylphenolb	<3	to	48	
		p-2-oxo-n-butylphenol	<3	to	1357	
		m-acetonylanisol ^b	<3	to	1546	
		isoprophylphenol ^b	<3	to	8	
		1-ethylpropylphenol	<3			
		dimethylphenol*	<3			
		benzoic acid	<3	to	12,311	
		Base Extractible Organics:				
		dichlorobenzene*	<10	to	172	
		dimethylaniline			6940	(continue

CONTAMINANT CLASSIFICATION	SITE	PROBLEM DESCRIPTION AND WATER	QUALITY	REFERENCE
Halocarbons Aromatics Phenols Polynuclear Aromatics (continued)		Base Extractible Organics (continued): m-ethylaniline 1,2,4-trichlorobenzene* napthalene* methylnapthalene camphor chloroaniline benzylamine or o-toluidine phenanthrene* or anthracene* b - structure not validated by actual or	<10 to 7640 <10 to 28 <10 to 66 <10 to 290 <10 to 7571 <10 to 86 <10 to 471 <10 to 670 compound	
Halocarbons Aromatics Misc.	027	Groundwater pollution caused by the product storage of chemicals and waste residues in ical production facility (conc. in µg/l, exchloride tetrachloromethane* trichloromethane* trichloroethene tetrachloroethene hexachlorobutadiene* (C46) hexachlorocyclopentadiene* (C56) octachlorocyclopentene (C58) hexachlorobenzene* (C66)	vicinity of chem-	26
Metals Misc.	Compi- lation of sites	Pollutants found to be present in leachates nation of 43 landfills which accept industr	_	11

TABLE B-1 (continued)

CONTAMINANT LASSIFICATION	CODE	PROF	BLEM DES	CRIPTION	AND WATER Q	QUALITY	REFERENC
Metals					Typical	No. of Sites	
Misc. (continued)		Pollutant	Conc. Ra	nge (mg/l)	Conc. $(mg/1)$	Where Detected	
		As*	0.03	- 5.8	0.2	5	
	ļ	Ba	0.01	- 3.8	0.25	24	
		Cr*	0.01	- 4.20	0.02	10	1.
		Co	0.01	- 0.22	0.03	11	
		Cu*	0.01	- 2.8	0.04	15	
	1	CN*	0.005		0.008	14	
	1	Pb*	0.3	- 19	-	3	
		Hg*	0.0005	- 0.0008	0.0006	5	
		Мо	0.15	- 0.24	-	2	
	1	Ni*	0.02	- 0.67	0.15	16	
		Se*	0.01	- 0.59	0.04	21	
		Zn*	0.01	- 240	3.0	9	1
	{	Light					
		Organics	1.0	- 1000	80	10	
		Halogenated			•		1
	1	Organics	0.002	- 15.9	0.005	5]
		Heavy					
		Organics	0.01	- 0.59	0.1	8	
		* - A nr	iority po	llutant			
	}	_		Liucuiic			
		ND - Not	Detected				1
		P - Pres	ent.				
		1103					
	}						}

TABLE B-1 (continued)

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APPENDIX C

CHEMICAL TREATABILITY

Appendix Table C-l presents information on the treatability of individual chemical compounds by various concentration technologies. Primary organization of the table is by concentration process. For each concentration process, the treatability of individual chemical compounds is given with the compounds arranged in alphabetical order within chemical calssifications. The following concentration processes are included:

Process	Process Code No.
Biological	Ī
Coagulation/Precipitation	II
Reverse Osmosis	III
Ultrafiltration	IV
Stripping	V
Solvent Extraction	VII
Carbon Adsorption	IX
Resin Adsorption	X
Miscellaneous Sorbents	XII

The chemical classification system used is described in the body of this report; the following chemical classes are used in Appendix C:

Chemical Classification	Classification Code No.
Alcohols	A
Aliphatics	В
Amines	С
Aromatics	D
Ethers	E
Halocarbons	F
Metals	G
PCBs	I
Pesticides	J
Phenols	K
Phthalates	L
Polynuclear Aromatics	M

In order to present the large quantity of information in a concise manner, it was necessary to code some of the information. The coding system is explained in footnotes at the end of

Table C-1.

Many chemical compounds are known by several names. Attempts were made to use preferred or generic names according to The Merck Index. However, in some cases it was necessary to use the names which were used in the reference documents. Users of Table C-1 are advised to check for compounds under several potential alphabetic listings.

Concentration Process: Biological Treatment (I) Chemical Classification: Alcohols(A)

Chemical b Amyl Alcohol -Pentanol) rneal tanol tanol	R U F R	Waste Type d	Influent Char.	Results of Study Toxic threshold to sensitive aquatic organisms (approx) >350 mg/l. 90.3% reduction based on COD; rate of biodegradation 8.9 mg COD/g hr. 70-90% reduction. 98% reduction w/80% BOD reduction.	Activated sludge process. Aerated lagoon treatment. Completely mixed acti-	99 81 100
-Pentanol) rneal tanol tanol	U F	I		aquatic organisms (approx) >350 mg/l. 90.3% reduction based on COD; rate of biodegradation 8.9 mg COD/g hr. 70-90% reduction. 98% reduction w/80% BOD	process. Aerated lagoon treatment. Completely mixed acti-	81
tanol tanol	F	I		COD; rate of biodegradation 8.9 mg COD/g hr. 70-90% reduction.	process. Aerated lagoon treatment. Completely mixed acti-	100
tanol	F			98% reduction w/80% BOD	treatment. Completely mixed acti-	
tanol		I				101
	R			1	tivated sludge process.	
4 1				Toxic threshold to sensitive aquatic organisms (approx) <250 ppm.		99
tanol	F	I	BOD load of 42 lb/day/ 1000 ft ³	95-100% reduction.	Activated sludge process.	56
tanol	U	P		98.8% reduction based on COD; rate of biodegradation 84 mg COD/g hr.	Activated sludge process.	81
c-Butanol	ט	P		rate of biodegradation 55 mg COD/g hr.	Activated sludge process.	81
	U	P		95.5% reduction based on COD; rate of biodegradation 30 mg COD/g hr.	Activated sludge process.	81
rt-Butanol	L	S		Substrate partially degraded.	Acclimated aerobic culture.	102
r	t-Butanol	t-Butanol U	t-Butanol U P	t-Butanol U P	-Butanol U P 98.5% reduction based on COD; rate of biodegradation 55 mg COD/g hr. t-Butanol U P 95.5% reduction based on COD; rate of biodegradation 30 mg COD/g hr.	-Butanol U P 98.5% reduction based on COD; Activated sludge rate of biodegradation process. 55 mg COD/g hr. t-Butanol U P 95.5% reduction based on COD; Activated sludge rate of biodegradation process. 30 mg COD/g hr. t-Butanol L S Substrate partially degraded. Acclimated aerobic

Concentration Process: Biological Treatment (I) Chemical Classification: Alcohols (A)

ا ا	h	Descr	iption o	of Study			I
No.	Chemical b	Study Type ^C	Waste Type d	Influent Char.	Results of Study	Comments	Ref.
1A- 11	1,4-Butanediol	ט	P		98.7% reduction based on COD; rate of biodegradation 40 mg COD/g hr.	Activated sludge process.	81
1A- 12	Cyclohexanol	U	P		96% reduction based on COD; rate of biodegradation 28 mg COD/g hr.	Activated sludge process.	81
1A- 13	Cyclopentanol	υ	P		97% reduction based on COD; rate of biodegradation 55 mg COD/g hr.	Activated sludge process.	81
1A- 14	Dimethylcyclo- hexanol	U	P		92.3% reduction based on COD; rate of biodegradation 21.6 mg COD/g hr.	Activated sludge process.	81
1A- 15	1,2-Ethanediol	L	S	484 ppm	74-76% reduction of BOD in 24 hrs. 7.5% of TOD exerted in 24 hrs.	Pure aerobic culture.	103
IA- 16	Ethanol	F	I		70-90% reduction.	Treated by aerated lagoon.	100
1A- 17	Ethanol	L	Ü	1000 ppm	>99% reduction of BOD in 24 hrs. 24% of TOD exerted in 24 hrs.	Pure aerobic culture.	103
IA- 18	Ethanol	F	I		95-100% reduction w/80% BOD reduction.	Completely mixed acti- vated sludge process.	101
IA- 19	Ethyl Butanol	F	I		30-50% reduction.	Treated by aerated lagoon.	100
IA- 20	Ethyl Butanol	F	I		95-100% reduction w/80% BOD reduction.	Completely mixed acti- vated sludge process.	101
IA- 21	Ethyl Butanol	F	I	42 lb/day/ 1000 ft ³	75-85% reduction.	Activated sludge process.	56
IA- 22	2-Ethylhexanol	F	I	42 lb/day/ 1000 ft ³	75-85% reduction.	Activated sludge process.	56
	•					(continue	ed)

TABLE C-1 (continued)

Concentration Process: Biological Treatment (I) Chemical Classification: Alcohols (A)

r		T					Γ
No.	Chemical	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IA- 23	Furfuryl Alcohol	υ	P		97.3% reduction based on COD; rate of biodegradation 41 mg COD/g hr.	Activated sludge process.	81
IA- 24	Furfuryl Alcohol	Ū	P		96.1% reduction based on COD; rate of biodegradation 40 mg COD/g hr.	Activated sludge process.	81
IA- 25	Hexanol	Ū	P		95-100% reduction.	Activated sludge process.	56
IA- 26	1-Hexanol	F	I		70-90% reduction.	Treated by aerated lagoon.	100
IA- 27	1-Hexanol	F	I		100% reduction w/80% BOD reduction.	Completely mixed acti- vated sludge process.	101
IA- 28	Isopropanol	F	I		70-90% reduction.	Treated by aerated lagoon.	100
IA- 29	Isopropanol	F	I		96% reduction w/80% BOD reduction.	Completely mixed acti- vated sludge.	101
IA- 30	Isopropanol	L	S		100% reduction; acetone was intermediate where upon 50% reduced by bio-oxidation & 50% removed by air stripping.	Acclimated aerobic culture.	102
IA- 31	Isopropanol	U	р		99% reduction based on COD; rate of biodegradation 52 mg COD/g hr.	Activated sludge process.	81
32	Isopropanol	Ü	P	BOD load of 42 lb/day/ 1000 ft ³	95-100% reduction.	Activated sludge process.	56
				-		(continu	led)

TABLE C-1 (continued)

Concentration Process: Biological Treatment (I) Chemical Classification: Alcohols (A

a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IA- 33	Methanol	F	I	BOD load of 42 lb/day/ 1000 ft ³	75-85% reduction.	Activated sludge process.	56
IA- 34	Methanol	F	I		30-50% reduction.	Treated by aerated lagoon.	100
IA- 35	Methanol	L	U	997 ppm	2.4-5.7% reduction of BOD 24 hrs. 36 to 41 mg 0_2 used in 24 hrs. 2.4 -1.7% of TOD exerted in 24 hrs.	Pure aerobic culture.	103
IA- 36	Methanol	L	Ü	500 ppm	110 mg 0 ₂ used in 24 hrs. 14.6% of TOD exerted in 24 hrs.	Pure aerobic culture.	103
IA- 37	Methanol	F	I		84% reduction w/80% BOD reduction.	Completely mixed acti- vated sludge.	101
1A- 38	Methanol	F,C	I	170-2550 ppb	Effluent conc. of 150-510ppb achieved.	Survey of 2 municipal wastewater treatment plants.	65
IA- 39	4-Methylcyclo- hexanol	U	P		94% reduction based on COD; rate of biodegradation 40 mg COD/g hr.	Activated sludge process.	81
IA- 40	Octanol	F	I		75% reduction w/80% BOD reduction.	Completely mixed acti- vated sludge.	101
IA- 41	Octanol	F	I		30-50% reduction.	Treated by aerated lagoon.	100
IA- 42	Pentarythritol	L	I		No toxic effect.	Aerobic culture.	104
IA- 43	Phenyl Methyl Carbinol	F	I		85-95% reduction	Completely mixed acti- vated sludge.	101
						(continu	ed)

TABLE C-1 (continued)

Concentration Process: Biological Treatment (I) Chemical Classification: Alcohols (A)

No.	Chemical b	Descr Study Type ^C	iption c Waste Type d	f Study Influent Char.	Results of Study	Comments	Ref.
IA- 44	n-Propanol	U	P		98.8% reduction based on COD; rate of biodegradation 71 mg COD/g hr.	Activated sludge process.	81
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				!			
						(continu	ed)

TABLE C-1 (continued)

Concentration Process: Biological Treatment (I) Chemical Classification: Aliphatics (B)

No.	Chemical b	Descr Study Type ^C	Waste	Influent Char.	Results of Study	Comments	Ref.
IB- 1	Acetaldehyde	F	I		70-90% reduction.	Treated by aerated lagoon.	100
IB- 2	Acetaldehyde	F	I	BOD load of 42 lb day/1000 ft ³	85-95% reduction.	Activated sludge process.	56
IB- 3	Acetaldehyde	F,C	I	120-900 ppb	Effluent conc. of 90-1350ppb achieved.	Survey of 2 municipal wastewater treatment plants.	65
IB- 4	Acetone	F,C	Ι	100-600 ppb	Effluent conc. of 50-300 ppb achieved.	See IB-3 for comments.	65
IB- 5	Acetone	F	Ι		70-90% reduction.	Treated by aerated lagoon.	100
IB- 6	Acetone	В	S		Completely degraded or lost by stripping.	No identifiable degra- dation product.	102
IB- 7	Acetonitrile	В	U	490 ppm	Oxygen consumption was to- tally inhibited for 24 hrs.		103
8 8	Acetonitrile	В	S	500 ppm	Toxic or inhibitory during oxidation periods up to 72 hrs. 1.4% TOD was exerted in 72 hrs.	,	106
IB- 9	Acetylglycine	0	D	500 ppm	Readily oxidized w/9.3% of TOD exerted after 6 hr & 18.5% after 24 hr of oxidation.		106
IB- 10	Acrolein	F,C	I	50-150 ppb	Effluent conc. of 20-200 ppb achieved.	Survey of 2 municipal wastewater treatment plants.	65
						(continue	ed)

		Descr	iption o	of Study			T
No.	Chemical b	Study Type ^C		Influent	Results of Study	Comments	Ref.
IB- 11	Acrylic Acid	F	I	BOD load of 42 lb/day/ 1000 ft ³	85-95% reduction.	Activated sludge process.	56
IB- 12	Acrylic Acid	F	I		50-70% reduction.	Treated by aerated lagoon.	100
IB- 13	Acrylic Acid	F	I		85-95% reduction.	Completely mixed acti- vated sludge process.	101
IB- 14	Acrylonitrile	F	I		70-90% reduction.	Treated by aerated lagoon.	100
IB- 15	Acrylonitrile	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
IB- 16	Acrylonitrile	F	I	BOD load of 42 lb/day/ 1000 ft ³	95-100% reduction.	Activated sludge process.	56
IB- 17	Acrylonitrile	F	I	140 ppm	100% reduction.	Activated sludge process.	90
IB- 18	Adipic Acid	I	D	500 ppm	Readily oxidized w/7.1% of TOD exerted after 24 hr of oxidation.	Oxidation improved greatly after 12 hrs.	107
IB- 19	Alanine	В	U	500 ppm	Up to 39% of TOD exerted in 24 hrs.	Oxygen consumption showed no lag period. Material was readily degraded.	103
1B- 20	Ammonium Oxalate	U	P		92.5% reduction based on COD; rate of biodegradation 40 mg COD/g hr.	Activated sludge process.	81
IB- 21	Butanedinitrile	0	D	500 ppm	Toxic at oxidation periods up to 72 hrs.		106
,						(continu	ed)

a No.	Chemical b	Descr Study		of Study Influent	Results of Study	Comments	Ref.
		Type C	Type d	Char.	nours of beaut		ICI.
IB- 22	Butanedinitrile	0	D	500 ppm	Readily, but slowly, oxi- dized, 3.8% of TOD exerted after 24 hr of oxidation.	Oxygen uptake showed plateau effect after 12 hrs.	107
IB- 23	Butanenitrile	0	D	500 ppm	Inhibited oxidation for up to 24 hrs; after 72 hrs,up to 10.5% of TOD was exerted.		106
1B- 24	Butanenitrile	0	D		Readily, but slowly oxidized. Most rapid oxidation occurred in first 6 hrs, 1.7% of TOD exerted after 24 hrs.	for comments.	107
IB- 25	Butyleneoxide	0	D	500 ppm	9.6% of TOD exerted after 144 hrs of oxidation.	Degraded very slowly.	108
IB- 26	Butyric Acid	F	I	BOD load of 42 lb/day/ 1000 ft ³	85-95% reduction.		56
IB- 27	Butyric Acid	0	D	500 ppm	Up to 43% of TOD exerted after 72 hrs of oxidation.		106
IB- 28	Butyric Acid	F	I		50-70% reduction.	Treated by aerated lagoon.	100
IB- 29	Butyric Acid	0	D		Rapidly oxidized for first 6 hrs; after 24 hrs of oxidation up to 27.9% of TOD was exerted.		107
IB- 30	Calcium Gluconate	L	U	250 ppm	13.6% of TOD exerted in 24 hrs.		103
IB- 31	Caprolactam	U	P		94.3% reduction based on COD; rate of biodegradation 16 mg COD/g hr.	-	
		1				(continue	d)

TABLE C-1 (continued)

a No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IB- 32	Citric Acid	L	U	550 ppm	35 mg of 0_2 used in 24 hrs.	Biodegradable, depressed 02 consumption.	
IB- 33	Crotonaldehyde	F	I	BOD load of 421b/day/ 1000 ft ³	95-100% reduction.		56
IB- 34	Crotonaldehyde	F	I		90-100% reduction.	Treated by aerated lagoon.	100
IB- 35	Crotonaldehyde	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
IB- 36	Cystine	L	U	1000 ppm	Completely inhibited any consumption of 0_2 .		103
IB- 37	L-Cystine	0	D	500 ppm	Slowly oxidized w/4.7% of TOD exerted after 24 hrs of oxidation.		107
IB- 38	Cyclohexa- nolone	Ū	P		92.4% reduction based on COD; rate of biodegradation 51.5 mg COD/g hr.	Activated sludge process.	81
IB- 39	Cyclohexanone	Ū	P		96% reduction based on COD; rate of biodegradation 30 mg COD/g hr.	Activated sludge process.	81
IB- 40 ·	Cyclopentanone	Ü	P		95.4% reduction based on COD; rate of biodegradation 57 mg COD/g hr.	Activated sludge process.	81
IB- 41	Diethylene Glycol	Ü	P		95% reduction based on COD; rate of biodegradation 13.7 mg COD/g hr.		81
IB- 42	2,3-Dithiabu- tane	F,C	I	10-120ppb		See IB-3 for comments.	65
						(continu	ed)

a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IB- _43	Dulcitol	0	υ	1700 ppm	Slightly inhibitory		109
IB- 44	Erucic Acid	0	D	500 ppm	11% of TOD exerted after 24 hrs of oxidation.		107
IB- 45	Ethyl Acetate	F	I		90-100% reduction.	Treated by aerobic lagoon.	100
IB- 46	Ethyl Acetate	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
IB- 47	Ethyl Acetate	F	I	BOD load of 42 lb/day, 1000 ft ³	95-100% reduction.	Activated sludge process.	56
IB- 48	Ethyl Acrylate	F	I	BOD load of 42 lb/day 1000 ft ³	95-100% reduction.	Activated sludge process	56
IB- 49	Ethyl Acrylate	F	I		90-100% reduction.	Treated by aerobic lagoon	100
IB- 50	Ethyl Acrylate	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
IB- 51	Ethylene Glycol	U	P		96.8% reduction based on COD; rate of biodegradation 41.7 mg COD/g hr.	Activated sludge process.	81
IB- 52	2-Ethylhexyl- acrylate	F	I	BOD load of 42 lb/day 1000 ft ³	95-100% reduction.	Activated sludge process.	56
IB- 53	2-Ethylhexyl- acrylate	F	I		90-100% reduction	Treated by aerobic lagoon.	100
						(continue	ed)

TABLE C-1 (continued)

r		7	ar cras.	STLICACION	: Aliphatics (B)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
IB- 54	2-Ethylhexyl- acrylate	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
IB- 55	Formaldehyde	L	Ü	720 ppm	Chemical inhibited 0 ₂ consumption.	resea studge process.	103
IB- 56	Formaldehyde	0	D	3000 ppm	<99% reduction after 24 hrs of aeration.	pH held at 7.2.	104
IB- 57	Formamide	0	D	500 ppm	Slowly oxidized for first 12 hrs; 11.8% of TOD exerted after 24 hrs of oxidation.		107
IB- 58	Formic Acid	L		720 ppm	70% of TOD exerted after 24 hrs of oxidation.	No lag period during oxidation.	167
IB- 59	Glutamic Acid	L			31% of TOD exerted after 24 hrs of oxidation.	ontduction.	103
IB- 60	Glycerine	L		720 ppm	248 mg of 0_2 used in 24 hrs.		103
IB- 61	Glycine	L		720 ppm	58% of TOD exerted after 24 hrs.		103
IB- 62	Heptane	F	I	BOD load of 42 lb/day/ 1000 ft ³	95-100% reduction.	Activated sludge process.	56
63	Heptane .	0	D	500 ppm	38.7% of TOD exerted after 72 hrs.		106
64	Heptane	F	I		90-100% reduction.	Treated by aerated lagoon.	100
65	Heptane	F	I		95-100% reduction.	Completely mixed acti-	101
	Hydracrylo- nitrile	F	I		0-10% reduction.	vated sludge process. Treated by aerated lagoon.	100
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No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IB- 67	Isophorone	F,C	D .		93% reduction.	21 day maximum retention time in a series of lagoons.	81
IB- 68	Lactic Acid	L		720 ppm	78% of TOD exerted after 24 hrs.		7
IB- 69	Lauric Acid	0	D	500 ppm	6.1% of TOD exerted after 24 hrs.		107
IB- 70	L-Malic Acid	0	D	500 ppm	44.8% of TOD exerted after 24 hrs.		107
IB- 71	DL-Malic Acid	0	D	500 ppm	20.8% of TOD exerted after 24 hrs.	A 10-16 hr lag period was indicated.	107
IB- 72	Malonic Acid	0	D	500 ppm	Chemical inhibited 0 ₂ uptake.		107
IB- 73	Nitrilotri- acetate	L	S	20 to 500 ppm	>90% reduction after acclimation.		111
IB- 74	Oleic Acid	0			02 uptake inhibited.		109
IB- 75	Oxalic Acid	L		250 ppm	⁰ 2 uptake inhibited.		103
IB- 76	Pentane	0	D	500 ppm	0 ₂ uptake inhibited.		106
IB- 77	Pentanedini- trile	0	D	500 ppm	Toxic at oxidation periods of up to 72 hrs.		106
IB- 78	Pentanedini- trile	0	D	500 ppm	Slowly oxidized with 2.9% of TOD exerted after 24 hrs of oxidation.		106
IB- 79	Pentanenitrile	0	D	500 ppm	Toxic to 2 sludges at oxi- dation periods up to 24 hrs.		106
IB- 80	Propanedini- trile	0	D	500 ppm	Toxic for oxidation periods up to 72 hrs.		106
						(continu	ed)

No.	Chemical b	Descr Study Type ^C		Influent Char.	Results of Study	Comments	Ref.
IB- _81	Propanenitrile	0	D,	500 ppm	Toxic for oxidation periods up to 72 hrs.	,	106
IB- 82	β-Propiolactone	0	D	500 ppm	0 ₂ uptake inhibited.		108
IB- 83	Sodium Alkyl Sulfonate	0			22% of TOD exerted after 5 days.		112
IB- 84	Sodium Lauryl Sulfate	0			65% of TOD exerted after 5 days.		112
IB- 85	Sodium N- Oleyl-N-Methyl Taurate	0			47-52% of TOD exerted in 5 days.		112
IB- 86	Sodium α Sulfo Methyl Myristate	0			33% of TOD exerted after 5 days.		112
IB- 87	Tannic Acid	0			0 ₂ uptake inhibited.		109
IB- 88	Thioglycollic Acid	L			0 ₂ uptake inhibited within 24 hrs.		103
IB- 89	Thiouracil	0	D	500 ppm	Chemical was oxidized but very slowly. 12.8% of TOD exerted after 144 hrs of oxidation.		108
IB- 90	Thiourea	0	D	500 ppm	0, uptake was inhibited by chemical for up to 144 hrs of oxidation.		103
	Triethylene Glycol	U	P		97.7% reduction based on COD; rate of biodegradation was 27 mg COD/g hr.	Activated sludge process	81
IB- 92	Urea	L		1200 ppm	O ₂ uptake inhibited.		103
						(continue	ed)

TABLE C-1 (continued)

a No.	Chemical b	Descr Study Type ^C	iption c Waste Type d	f Study Influent Char.	Results of Study	Comments	Ref.
IB- 93	Urethane	0	D		02 uptake inhibited.		108
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				1		(continue	ed) '

a	h	Descr	iption o	of Study			
No.	Chemical b	Study Type ^C	Waste Type d	Influent	Results of Study	Comments	Ref.
IC- 1	Acetanilide	ט	P		94.5% reduction based on COD; rate of biodegradation 19 mg COD/g hr.	Activated sludge process.	81
IC- 2	p-Aminoacetan- ilide	υ	P		93% reduction based on COD; rate of biodegradation 11.3 mg COD/g hr.	Activated sludge process.	81
IC- 3	m-Aminobenzoic Acid	U	P		97.5% reduction based on COD; rate of biodegradation 27.1 mg COD/g hr.	Activated sludge process.	81
IC- 4	o-Aminobenzoic Acid	Ü	P		97.5% reduction based on COD; rate of biodegradation 7.0 mg COD/g hr.	Activated sludge process.	81
IC- 5	p-Aminobenzoic Acid	Ü	P		96.2% reduction based on COD; rate of biodegradation 12.5 mg COD/g hr.	Activated sludge process.	81
IC- 6	m-Aminotoluene	Ü	P		97.7% reduction based on COD; rate of biodegradation 30 mg COD/g hr.	Activated sludge process.	81
1C- 7	o-Aminotoluene	υ	P		97.7% reduction based on COD; rate of biodegradation 15.1 mg COD/g hr.	Activated sludge process.	81
IC- 8	p-Aminotoluene	Ū	P		97.7% reduction based on COD; rate of biodegradation 20 mg COD/g hr.	Activated sludge process.	81
IC- 9	Aniline	Ū	P		94.5% reduction based on COD; rate of biodegradation 19 mg COD/g hr.	Activated sludge	81
IC- 10	Aniline	Ü	I	500 ppm 30°C	100% reduction in 15 hrs.	Biodegradation by mu- tant pseudomonas.	92
						(continu	ed)

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type C		of Study Influent Char.	Results of Study	Comments	Ref.
IC-	Aniline	0	0	500 ppm	0_2 uptake inhibited for up to 72 hrs.		108
1C- 12	Benzamide	0	0	500 ppm	O ₂ uptake inhibited for first 6 hrs. 63% of TOD exerted after 144 hrs of oxidation.		108
IC- 13	Benzidine	0	D	500 ppm	0 ₂ uptake inhibited.		108
IC- 14	Benzidine	F,C	D	1.6 ppb	0% reduction.	Activated sludge process.	81
IC- 15	Benzylamine	0	D	500 ppm	0_2 uptake inhibited.		108
IC- 16	Butanamide	0	D	500 ppm	Slowly oxidized w/6.4% of TOD exerted after 24 hrs of oxidation.		107
IC- 17	m-Chloroani- line	Ü	P		97.2% reduction based on COD; rate of biodegradation 6.2 mg COD/g hr.	Activated sludge process.	81
IC- 18	o-Chloroani- line	Ü	P		97.2% reduction based on COD; rate of biodegradation 16.7 mg COD/g hr.	Activated sludge process.	81
IC- 19	p-Chloroani- line	U	P		96.5% reduction based on COD; rate of biodegradation 5.7 mg COD/g hr.	Activated sludge process.	81
IC- 20	Diethanolamine	. U	P		97% reduction based on COD; rate of biodegradation 19.5 mg COD/g hr.	Activated sludge process.	81
IC- 21	2,3-Dimethyl- aniline	U	P		96.5% reduction based on COD rate of biodegradation 12.7 mg COD/g hr.	Activated sludge process.	81
						(continue	ed)

a	h	Descr	iption o	of Study			
No.	Chemical b	Study Type ^C	Waste Type d	Influent Char.	Results of Study	Comments	Ref
IC- 22	2,5-Dimethyl- aniline	υ	₽.		96.5% reduction based on COD; rate of biodegradation 3.6 mg COD/g hr.	Activated sludge process.	81
IC- 23	3,4-Dimethyl- aniline	U	P		76% reduction based on COD; rate of biodegradation 30 mg COD/g hr.	Activated sludge process.	81
IC- 24	Ethylene- diamine	U	P	·	97.5% reduction based on COD; rate of biodegradation 9.8 mg COD/g hr.	Activated sludge process.	81
IC- 25	2-Fluorenamine	0	D	500 ppm	<pre>02 uptake showed inhibitory effect but was slowly bio- logically oxidized.</pre>		108
IC- 26	o-Nitroaniline	Ū	ī	18.5 ppm	<99.9% reduction.	Powder activated carbon & activated sludge treatment.	58
IC- 27	p-Nitroaniline	U	I	6.7 ppm	<99.9% reduction.	See IC-26 for comments.	58
IC- 28	P-(Phenylazo) aniline	0	D	500 ppm	<pre>02 uptake inhibited after 72 hrs of oxidation.</pre>		108
IC- 29	Pentanamide	0	D	500 ppm	Slowly oxidized w/13.6% of TOD exerted after 24 hrs of oxidation.	·	107
IC- 30	Phenylene- diamine	0	D	500 ppm	Toxic during 24 hrs of aeration		113
IC- 31	m-Phenylene- diamine	U	P		60% reduction based on COD.	Activated sludge process.	81
IC- 32	o-Phenylene- diamine	Ū	P		33% reduction based on COD.	Activated sludge process.	81
IC- 33	p-Phenylene diamine	U	P		80% reduction based on COD.	Activated sludge process.	81
						(continue	ed)

a No.	Chemical b	Study	Waste	of Study Influent	Results of Study	Comments	Ref.
		Type c	Type d	Char.			
IC- 34	Thioacetamide	L	ָט	100 ppm	0 ₂ uptake inhibited.		103
IC-	2,4,6-Trichlo- roaniline	U	Ī	500 ppm	100% reduction in 30 hrs.	See IC-10 for comments.	92
IC- 36	2,4,6-Trichlo- roaniline	0	S	10 ppm	0 ₂ uptake not inhibited.		113
						(c	ontinued)

TABLE C-1 (continued)

Concentration Process: Biological Treatment (I) Chemical Classification: Aromatics (D)

	····	T	· · · · · · · · · · · · · · · · · · ·				,
a	b			of Study			
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref
		Type ^C	Type d	Char.			
ID- 1	sec-Amyl- benzene	0	D	500 ppm	Toxic for 24 hrs of aeration.		113
ID- 2	tert-Amyl- benzene	0	D	500 ppm	Toxic for 24 hrs of aeration.		113
ID- 3	Benzaldehyde	0			0 ₂ uptake inhibited.		109
ID- 4	Benzaldehyde	U	P		99% reduction based on COD; rate of biodegradation 119 mg COD/g hr.	Activated sludge process.	81
ID- 5	Benzaldehyde	0	D	500 ppm	61.3% of TOD exerted after 144 hrs of oxidation.		108
ID- 6	Benzene	F	I		90-100% reduction.	Treated by aerated lagoon.	100
ID- 7	Benzene	F	I		95-100% reduction.	Completely mixed activated sludge process.	101
ID-	Benzene	0	D	125 ppm	1.44-1.45g of oxygen uti- lized per gram of substrate added after 72 hrs of oxidation.		114
ID- 9	Benzene	0	D	50-500 ppm	02 uptake of 34 ppm 02/hr for 50 ppm chemical & 37 ppm 02/hr for 500 ppm chemical.	•	114
ID- 10	Benzene	F	I		95-100% reduction.	Activated sludge process.	56
ID- 11	Benzene Sulfonate	0	D	500 ppm	Slowly oxidized for first 6 hrs; 62% of TOD exerted after 144 hrs.		108
ID- 12	Benzenethiol	0	D	500 ppm	0_2 uptake inhibited for up to 144 hrs of oxidation.		108
1				,		(continu	led)

181

Concentration Process: Biological Treatment (I)

Chemical Classification: Aromatics (D)

No	a Chemical b	<u>Descr</u> Study			Results of Study	Comments	Ref.
		Type C	Type d	Char.	•		
1	Benzoic Acid	U	₽,		99% reduction based on COD; rate of biodegradation 88.5 mg COD/g hr.		81
11		F	I	BOD load of 42 lb/day, 1000 ft ³	95-100% reduction	Activated sludge process.	56
1	- Benzonitrile 5	0	D	500 ppm	0_2 uptake inhibited for up to 72 hrs of oxidation.		106
II	- 3,4-Benzpyrene	0	D	500 ppm	O ₂ uptake inhibited for up to 144 hrs of oxidation.		106
	sec-Butyl- benzene	0	D	500 ppm	Toxic for 24 hrs of aeration.		113
1	- tert-Butyl- 8 benzene	0	D	500 ppm	Toxic for 24 hrs of aeration.		113
11	Chloranil	0	S	10 ppm	0 ₂ uptake inhibited.		102
11	Chlorobenzene	L	P	200 ppm	100% reduction in 14 hrs.	Biodegradation by mu- tant pseudomonas species.	66
11	1,2,4,5-Dibenz pyrene	0	D	500 ppm	02 uptake inhibited for up to 144 hrs of oxidation.		108
II	D- m-Dichloro- 2 benzene	L	Р	200 ppm	100% reduction in 28 hrs.	See ID-20 for comments.	66
11	D- m-Dichloro- 3 benzene	U	I	200 ppm	100% reduction in 30 hrs.	See ID-20 for comments.	92
II	o-Dichloro- de benzene	L	P	200 ppm	100% reduction in 20 hrs.	See ID-20 for comments.	66
II	p-Dichloro- benzene	L	P	200 ppm	100% reduction in 25 hrs.	See ID-20 for comments.	66
	.]					(continu	ed)

182

a No.	Chemical	Descr Study	iption o Waste	of Study Influent	Results of Study	Comments	Ref.
	OH CHILDREN	Type C					
ID-	2,4-Dichloro-	L	D.	174 ppm	No reduction until after 5	Subjected to continuous	115
26	phenoxyacetic Acid				days.	aeration.	
ID-	2,6-Dichloro-	L	D	178 ppm	No reduction until after 3	See ID-26	115
27	phenoxyacetic Acid				days.	for comments.	
ID-	2,4-Dichloro-	L	D	186 ppm	No reduction after 7 days.	See ID-26	115
28	phenoxypro- pionic Acid				•	for comments.	
ID-	7,9-Dimethyl-	0	D	500 ppm	02 uptake inhibited after		108
29	benzacridine			1	144 hrs of oxidation.		
ID-	7,10-Dimethyl-	0	D	500 ppm	02 uptake inhibited after		108
30	benzacridine				after 144 hrs of oxidation.		
ID-	3,5-Dinitro-	U	P		50% reduction based on COD.	Activated sludge	81
31	benzoic Acid					process.	
ID-	2,4-Dinitro-	F,C	D	390 ppb	Not detectable in effluent.	Activated sludge	81
32	toluene		<u> </u>	<u> </u>		process.	90
ID-	2,4-Dinitro-	R	U	146-188	90% reduction.	Activated sludge	90
33	toluene			ppm		process.	56
ID-	Ethylbenzene	F	I	BOD load	95-100% reduction	Activated sludge	1 30
34				of	,	process.	ļ
.		}		42 lb/day		•	
	T11-11	U	s	192 ppb	100% reduction.		21
ID-	Ethylbenzene	١	3	132 bbp	100% Leadecton.		
35 ID-	Ethylbenzene	F	I	+	90-100% reduction.	Treated by aerated	100
36	Eculythenzene	, r	-		20 200 200002011	lagoon.	
ID-	Ethylbenzene	$+_{\overline{L}}$	I	 	95-100% reduction	Completely mixed acti-	101
37	20, 22020	-				vated sludge.	
T-5-			1				1
						(continu	ed)

a	h			of Study			
No.	Chemical b	Study Type ^C	Waste Type d	Influent Char.	Results of Study	Comments	Ref
38	Ethylbenzene	0	D ,	105 ppm	After 72 hrs of oxidation $1.7g$ of 0_2 was used per g chemical added.		114
ID- 39	Hexachloro- benzene	L	P	200 ppm	0% reduction in 120 hrs.	See ID-20 for comments.	66
ID- 40	Hexachloro- benzene	U	I	200 ppm	0% reduction in 120 hrs.	See ID-20 for comments.	92
ID- 41	4-Hydroxy- benzenecarbo- nitrile	0	Ď	500 ppm	Toxic after 72 hrs of oxidation.		106
ID- 42	2-Methylben- zenecarbo- nitrile	0	D	500 ppm	Toxic after 72 hrs of oxidation.		106
ID- 43	3-Methylben- zenecarbo- nitrile	0	D	500 ppm	Toxic after 72 hrs of oxidation.		106
ID- 44	4-Methylben- zenecarbo- nitrile	0	D	500 ppm	Toxic after 72 hrs of oxidation.		106
ID- 45	Methylethyl- pyridine	F	I		10-30% reduction.	Treated by aerated lagoon.	100
ID- 46	m-Nitrobenz- aldehyde	Ü	P		94% reduction based on COD; rate of biodegradation 10 mg COD/g hr.	Activated sludge process.	81
ID- 47	o-Nitrobenzal- dehyde, p-Ni÷ trobenzáldehyde	Ü	P		97% reduction based on COD; rate of biodegradation 13.8 mg COD/g hr.	Activated sludge	81
ID- 48	Nitrobenzene	υ	P		98% reduction based on COD; rate of biodegradation 14 mg COD/g hr.	Activated sludge process.	81
.]	-					(contin	ued)

TABLE C-1(continued)

a	h	Descr	iption o	f Study			
No.	Chemical b	Study Type ^C		Influent Char.	Results of Study	Comments	Ref.
ID- 49	Nitrobenzene	U	s .	175 ppb	100% reduction.		21
ID- 50	Nitrobenzene	Ü	I	530 ppb	< 96.0% reduction.	Powder activated car- bon & activated sludge treatment.	58
ID- 51	Nitrobenzene	F,C	D	58 ppb	>0.1 ppb effluent conc.	21 day maximum reten- tion time in a series of lagoons.	81
ID- 52	Nitrobenzene	0	D	500 ppm	0 ₂ uptake inhibited for up to 144 hrs of oxidation.		108
ID- 53	m-Nitrobenzoic Acid	U	P		93.4% reduction based on COD; rate of biodegradation 7 mg COD/g hr.	Activated sludge process.	81
ID- 54	o-Nitrobenzoic Acid	U	P		93.4% reduction based on COD; rate of biodegradation 20 mg COD/g hr.	Activated sludge process.	81
ID- 55	p-Nitrobenzoic Acid	Ü	P		92% reduction based on COD; rate of biodegradation 19.7 mg COD/g hr.	Activated sludge process.	81
ID- 56	m-Nitrotoluene	Ū	P		98.5% reduction based on COD; rate of biodegradation 21 mg COD/g hr.	Activated sludge process.	81
ID- 57	o-Nitrotoluene p-Nitrotoluene	ı	P		98% reduction based on COD; rate of biodegradation 32.5 mg COD/g hr.	Activated sludge process.	81
ID- 58	Nitrofluorine	0	D	500 ppm	Slowly oxidized w/13.7% of TOD exerted after 144 hrs.		108
ID- 59	Paraldehyde	F	I		30-50% reduction	Treated by aerated lagoon.	100
						(continu	l ed)

No.	Chemical b	Descr Study Type ^C	Waste	Influent Char.	Results of Study	Comments	Ref.
ID- 60	Pentamethyl- benzene	0	D.	500 ppm	0 ₂ uptake inhibited during first 24 hrs of aeration.		113
ID- 61	n-Propylben- zene	0	D	37.5 ppm	After 72 hrs of oxidation 0.67g of 02 were utilized per g of substrate added.	·	114
ID- 62	Sodium Alkyl- benzene Sul- fonate	0			26% of TOD exerted after 5 days.	·	112
ID- 63	Styrene	F	I		70-90% reduction.	Treated by aerated lagoon.	100
ID- 64	Styrene	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
ID- 65	1,2,3,4-Tetra- chlorobenzene	L	P	200 ppm	74% reduction in 120 hrs.	See ID-20 for comments.	66
ID- 66	1,2,3,5-Tetra- chlorobenzene	L	P	200 ppm	80% reduction in 120 hrs.	See ID-20 for comments.	66
ID- 67	1,2,4,5-Tetra- chlorobenzene	Ū	I	200 ppm	80% reduction in 120 hrs.	See ID-20 for comments.	66
ID- 68	1,2,4,5-Tetra- chlorobenzene	0	0	500 ppm	No 0_2 consumed during first 3 hrs; very slight uptake thereafter for first 24 hrs of aeration.		113
ID- 69	Toluene	F	I		70-90% reduction.	Treated by aerated lagoon.	100
ID- 70	Toluene	F	I		95-100% reduction.	Completely mixed acti- vated sludge process.	101
ID- 71	Toluene	0	D	500 ppm	O ₂ uptake inhibited or very slightly oxidized for first 24 hrs of oxidation.		108
						(continue	ed)

TABLE C-1 (continued)

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a No.	Chemical b	Descr Study		of Study Influent	Results of Study	Comments	Ref.
		Type ^C	Type d	Influent Char.	Results of beday	Commence	wer.
ID- 72	Toluene	0	D	100 ppm	0.53-0.65g of 0 ₂ used per g of substrate added after 72 hrs of oxidation.		114
ID- 73	Toluene	0	D	500 ppm	48.3% of TOD exerted after 72 hrs of oxidation.		106
ID- 74	Toluene	F,C	I		1.0-10.0 ppb effluent conc.	Survey of 2 municipal wastewater treatment plants.	65
1D- 75	Toluene	ħ	I	BOD load of 42 lb day/1000 ft ³	95-100% reduction.	Activated sludge process.	56
76	m-Toluidine	U	I	500 ppm	100% reduction in 10 hrs.	See ID-20 for comments.	92
77	1,2,3-Trichlo- robenzene	L	P	200 ppm	100% reduction in 43 hrs.	See ID-20 for comments.	66
	1,2,4-Trichlo- robenzene	L	P	200 ppm	100% reduction in 46 hrs.	See ID-20 for comments.	66
) 1	1,3,5-Trichlo- robenzene	U	I	200 ppm	100% reduction in 50 hrs.	See ID-20 for comments.	92
i 1	1,3,5-Trichlo- robenzene	L	P	200 ppm	100% reduction in 50 hrs.	See ID-20 for comments.	66
, ,	2,4,5-Trichlo- rophenoxypro- pionic Acid	L	0	107.5 ppm	99% reduction in 16.5 days.		115
	2,4,6-Trichlo- rophenoxy- acetic Acid	L	D	53 ppm	50% reduction in 14 days.	Subjected to continuous aeration.	115
1 1	2,6,6-Trini- trotoluene	L	I	100 ppm	50-84% reduction in 3-14 hrs.		116
						(continue	d)

TABLE C-1 (continued)

a No.	Chemical b	<u>Descr</u> Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
ID- 84	m-xylene o-xylene p-xylene	0	D [']	500 ppm	O ₂ uptake inhibited after 24 hrs of oxidation.		113
ID- 85	Xylene	F,C	I	20-200pph	1.0-15.0 ppb effluent conc.	See ID-74 for comments.	65
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Concentration Process: Biological Treatment (I) Chemical Classification: Ethers (E)

·		,			, Edicio (D)		
No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
IE-	Isopropyl Ether	F	I.	BOD load of 42 lb/day/ 1000 ft ³	85-95% reduction.	Activated sludge process.	56
IE-	Isopropyl Ether	F	I		70-90% reduction.	Treated by aerated lagoon.	100
IE- 3	Isopropyl Ether	F	I		85-95% reduction.	Completely mixed activated sludge process.	101
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TABLE C-1(continued)

a No.	Chemical b	Descr Study		of Study Influent	Results of Study	Comments	Ref.
	-	Type c	Type d	Char.	results of study	ConunciiCs	Rel.
IF- 1	Bromoform	F,C	I,	0.4-1.9 ppb	100% reduction.	Survey of 2 municipal wastewater treatment plants.	65
IF- 2	Carbon Tetrachloride	U	S	177 ppb	100% reduction.		21
1F- 3	Chloroform	F,C	I	13 ppb	100% reduction.	See IF-1 for comments.	65
IF- 4	1,2-Dichloro- ethane	F,C	I	0.4-260 ppb	1.4 ppb effluent conc.	See IF-1 for comments.	65
IF- 5	Methylene Chloride	F,C	I	10-430ppb	2.0-50 ppb effluent conc.	See IF-1 for comments.	65
IF- 6	1,1,1-Trichlo- roethane	F,C	I	8.0-790 ppb	1.0-20.0 ppb effluent conc.	See IF-1 for comments.	65
1F- 7	1,1,2-Trichlo- roethane	υ	I	1305 ppb	<pre>< 99.7% reduction</pre>	Powder activated carbon & activated sludge treatment.	58
IF- 8	Trichloro- ethylene	F,C	I	78 ppb	100% reduction.	See IF-1 for comments.	65
IF- 9	Trichloro- ethylene	F,C	I	214 ppb	99% reduction		21
IF- 10	Vinyl Chloride	F,C	I	8 ppb	100% reduction	See IF-1 for comments.	65
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TABLE C-1(continued)

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Study	Waste	Influent	Results of Study	Comments	Ref
Type ^C	Type d	Char.	·		
0	U.	1-100,000	O ₂ uptake inhibited at conc.		109
		ppm	greater than 100 ppm.		
R	Ū	6 ppb	1.0 ppb effluent conc.	Activated sludge process.	90
F,C	I	27 ppb	16 ppb effluent conc.	Survey of 2 municipal wastewater treatment plants.	65
0	Ü	1-100,000 ppm	Conc. of 1-10 ppm inhibited 02 uptake.		109
F	D	ranged from	22-78% reductions achieved.	Survey of municipal wastewater treatment	122
		0.8-3.6ppm		plants.	
C,P	D	15 ppm	0.2 ppb effluent conc.		123
0	Ü	1-100,000 ppm	02 uptake inhibited at conc. greater than 100 ppm.		109
L	S	0.08-0.5 ppm	Inhibited biological growth.	Study of <u>Nitrosomas</u> bacteria.	124
R	U	10 ppm	75% reduction.	Activated sludge process.	118
F	D	ranged from 0.2-1.5ppm	7-77% reductions achieved.	See IG-5 for comments.	122
L	S	5-30 ppb 50-560ppb	Stimulated biological growth Inhibited biological growth.		124
C,P	D	10 ppm	75% reduction.	Activated sludge process.	125
0	U	10-1000 ppm	0_2 uptake inhibited at conc. greater than 100 ppm.		109
			0 U 10-1000	O U 10-1000 O2 uptake inhibited at conc.	process. O U 10-1000 O ₂ uptake inhibited at conc.

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IG- 14	Iron (Fe ⁺³)	О	U _.	0.01- 100,000 ppm	0_2 uptake inhibited at conc. greater than 100 ppm.		109
IG- 15	Iron	C,F	D	7.17 ppm total iron 0.6 ppm soluble iron	83% reduction. 62% reduction.	,	126
IG- 16	Lead	0		10-100ppm	0 ₂ uptake inhibited		109
IG- 17	Lead	L	S	5-50 ppb	No stimulation or inhibition of biological growth.	See IG-8 for comments.	124
1G- 18	Manganese	L	S	12.5-50 ppm 50-100ppm	Stimulated biological growth Inhibited biological growth.	See IG-8 for comments.	124
IG- 19	Manganese	L	S	10 ppm	02 uptake inhibited.		109
IG- 20	Mercury	0	S	0-200 ppm	0 ₂ uptake inhibited.		127
IG- 21	Mercury	L	S	5-10 ppm	51-58% reduction.		132
IG- 22	Nickel	R	U	10 ppm	28% reduction.	Activated sludge process.	118
IG- 23	Nickel	F	D	ranged from 0.03-2.0 ppm	0-33% reduction achieved.	See IG-5 for comments.	122
IG- 24	Nickel	C,P	D	1-10 ppm	28-42% reduction.	Activated sludge process.	128
						(continue	 d)

Concentration Process: Biological Treatment (I) Chemical Classification: Metals (G)

No.	Chemical b	<u>Descr</u> Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IG- 25	Nickel	C,F	D.	270 ppb	30% reduction.	Activated sludge process.	129
IG- 26	Nickel	P	D	10 ppm	28% reduction.	Activated sludge process.	125
IG- 27	Strontium	L	S	5-50 ppb	No stimulation or inhibition of biological growth.	See IG-8 for comments.	124
IG- 28	Zinc	R	U	10 ppm	89% reduction.	Activated sludge process.	118
IG- 29	Zinc	F	D	ranged from 0.3-2.2ppm	20-91% reduction achieved.	See IG-5 for comments.	122
IG- 30	Zinc	C,P	D	2.5 ppm 10 ppm	13% reduction in primary treatment. 14% reduction in primary treatment.		128
IG- 31	Zinc	L	S	0.08-0.5 ppm	Biological growth inhibited.	See IG-8 for comments.	124
IG- 32	Zinc	C,F	D	0.91 ppm	60% reduction.	Activated sludge process.	131
IG- 33	Zinc	L	S	l ppm	0 ₂ uptake inhibited.		109
IG- 34	Zinc	R	Ü	3.57 ppm	57% reduction.	Activated sludge process.	90
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193

TABLE C-1 (continued)

	1_	Descr	iption o	of Study			
No.	Chemical D	Study Type ^C	Waste Type d	Influent Char.	Results of Study	Comments	Ref.
IJ- 1	Aldrin	0	, U		Not significantly degraded.		121
IJ- 2	Aminotriazole	0	U		Not significantly degraded.	,	121
IJ- 3	Chlordane	0	U		Slightly degraded.		121
IJ~ 4	2,4-D-Isoctyl- ester	0	Ū		Biodegradable.		121
IJ- 5	DDT	0	บ		Not significantly degraded.		121
IJ- 6	DDVP	L	U	37.5°С, 8.0 рН	462 min half-life.	Biodegradation by mutant pseudomonas species.	92
IJ- 7	Diazinon	L	Ü	20 ⁰ С, 10.4 рн	144 hr half-life.	See IJ-6 for comments.	92
IJ- 8	Diazinon	0	ប		Not significantly degraded.		121
IJ- 9	Dieldrin	0	U		Not significantly degraded.		121
IJ- 10	Endrin	0	U		Not significantly degraded.		121
IJ- 11	Ferbam	0	Ü		Biodegradable.		121
IJ- 12	Heptachlor	0	U	500 ppm	Slightly degraded.		121
IJ- 13	Herbicide Orange	F	1	1380 ppm	77% reduction.	Pure 0 ₂ & biological seeding provided.	81
IJ- 14	Lindane	0	Ū		Not significantly degraded.		121
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	1.	Descr	iption o	of Study			·1
No.	Chemical b	Study Type ^C	Waste	Influent	Results of Study	Comments	Ref.
IJ- 15	Malathion	0	U		Not significantly degraded.		121
IJ- 16	Malathion	L	U	25 ^O C, 10.03 pH	28 min half-life.	See IJ-6 for comments.	92
IJ- 17	Maneb	0	Ü		Biodegradable		121
IJ- 18	Methyl Parathion	L	Ü	15 ⁰ C	7.5 min half-life.	See IJ-6 for comments.	92
IJ-		0	Ü		Not significantly degraded.		121
IJ- 20	Parathion	L	U	15°C	32 min half-life.	See IJ-6 for comments.	92
IJ- 21	Parathion	0	Ū		Not significantly degraded.		121
IJ-	Pentachloro- phenol	0	Ü	75-150ppm	Not significantly degraded.		121
	Propoxur	Ö	บ	20 ⁰ С, 10.0 рн	40 min half-life.	See IJ-6 for comments.	92
IJ-	Tetraethyl Pyrophosphate	0	Ü		Not significantly degraded.		121
IJ- 25	Thanite	0	Ü		Biodegradable		121
IJ-	2,4,5-Trichlo- rophenoxyace- tic Acid	0	Ü	150 ppm	Slightly degraded.		121
IJ- 27	i ''	0			99% reduction in 7.5 days.	Subjected to continuous aeration.	115
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Concentration Process: Biological Treatment (I)

Chemical Classification: Pesticides (J)

No.	Chemical b	Descr Study Type C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
IJ- 28	Ziram	0	บ '		Slightly degraded.		121
IJ- 29		Ö	U		Slightly degraded.		121
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Concentration Process: Biological Treatment (I) Chemical Classification: Phenols (K)

		7	ar crab.	3111CGC1OI	: Phenois (K)		
No.	Chemical b	Descr Study Type C	Waste	Influent Char.	Results of Study	Comments	Ref.
IK- 1	4-Chloro-3- Methylphenol	О	S.	10 ppm 50 ppm 100 ppm	O ₂ uptake mildly inhibited. O ₂ uptake strongly inhibited. Toxic		102
IK- 2	4-Chloro-3- Methylphenol	R	U	25 ppm	Biodegradable in 5 days.		90
IK- 3	2-Chloro-4- Nitrophenol	U	P		71.5% reduction based on COD; rate of biodegradation 5.3 mg COD/g hr.	Activated sludge process.	81
IK- 4	2-Chlorophenol	R	U	150-200 ppm	90-95% reduction.	Activated sludge process.	90
IK- 5	m-Chlorophenol	L	P	200 ppm	100% reduction in 28 hrs.	Biodegradation by mu- tant pseudomonas species.	66
IK- 6	o-Chlorophenol	L	P	200 ppm	100% reduction in 26 hrs.	See IK-5 for comments.	66
IK- 7	o-Chlorophenol	Ü	P		95.6% reduction based on COD; rate of biodegradation 25 mg COD/g hr.	Activated sludge process.	81
1K- 8	p-Chlorophenol	U	P		96% reduction based on COD; rate of biodegradation 11 mg COD/g hr.	Activated sludge process.	81
IK- 9	p-Chlorophenol	L	P	200 ppm	100% reduction in 33 hrs.	See IK-5 for comments.	66
IK- 10	m-Cresol	Ü	P		96% reduction based on COD; rate of biodegradation 55 mg COD/g hr.	Activated sludge process.	81
1K- 11	o-Cresol	VU	P		95% reduction based on COD; rate of biodegradation 54 mg COD/g hr.	Activated sludge process.	81
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197

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IK- 12	p-Cresol	ט	P		95.5% reduction based on COD; rate of biodegradation 55 mg COD/g hr.	Activated sludge process.	81
1K-	2,4-Diamino- phenol	υ	р		83% reduction based on COD; rate of biodegradation 12 mg COD/g hr.	Activated sludge process.	81
1K- 14	2,4-Dichloro- phenol	U	P		98% reduction based on COD; rate of biodegradation 10.5 mg COD/g hr.	Activated sludge process.	81
IK- 15	2,4-Dichloro- phenol	R	Ū	60 ppm	Biodegradable in 5 days.		90
IK- 16	2,4-Dichloro- phenol	U	I	200 ppm	100% reduction in 35 hrs.	See IK-5 for comments.	90
IK-	2,4-Dichloro- phenol	L	P	200 ppm	100% reduction in 33 hrs.	See IK-5 for comments.	90
IK- 18	2,4-Dichloro- phenol	L	I	64 ррт	98% reduction in 5 days	Subjected to continuous aeration.	115
IK- 19	2,5-Dichloro- phenol	L	P	200 ppm	100% reduction in 38 hrs.	See IK-5 for comments.	66
IK- 20	2,6-Dichloro- phenol	L	I	64 ppm	99% reduction in 5 days.	See IK-18 for comments.	115
IK- 21	2,3-Dimethy1- phenol	U	P		95.5% reduction based on COD; rate of biodegradation 35 mg COD/g hr.	Activated sludge process.	81
IK- 22	2,4-Dimethyl- phenol	Ü	P		94.5% reduction based on COD; rate of biodegradation 28.2mg COD/g hr.	Activated sludge process.	81
1K- 23	2,5-Dimethy1- phenol	บ	P		94.5% reduction based on COD; rate of biodegradation 10.6 mg COD/g hr.	Activated sludge process.	81
						(continue	d)

Concentration Process: Biological Treatment (I)

Chemical Classification: Phenols (K)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref
IK- 24	2,6~Dimethyl- phenol	U	P		94.3% reduction based on COD; rate of biodegradation 9 mg COD/g hr.	Activated sludge process.	81
1K- 25	3,4-Dimethyl- phenol	U	P		97.5% reduction based on COD; rate of biodegradation 13.4 mg COD/g hr.	Activated sludge process.	81
1K- 26	3,5-Dimethyl- phenol	U	P		89.3% reduction based on COD; rate of biodegradation 11.1 mg COD/g hr.	Activated sludge process.	81
IK- 27	2,4-Dinitro- phenol	0	S	1 ppm	Maximum 0_2 uptake was 27.7ppm 0_2 /hr after 120 hrs of aeration		117
				5 ppm	Maximum 0_2 uptake was 21.3 ppm 0_2 /hr after 120 hrs of aeration.		
IK- 28	2,4-Dinitro- phenol	U	P		85% reduction based on COD; rate of biodegradation 6 mg COD/g hr.	Activated sludge process.	81
1K- 29	m-Nitrophenol p-	U	P		95% reduction based on COD; rate of biodegradation 17.5 mg COD/g hr.	Activated sludge process.	81
30	o-Nitrophenol	U	P		97% reduction based on COD; rate of biodegradation 14 mg COD/g hr.	Activated sludge process.	81
31	o-Nitrophenol	U	I	1275 ppb	<pre>< 98.1% reduction.</pre>	Powder activated carbon & activated sludge treatment.	58
1K- 32	p-Nitrophenol	U	I .	725 ppb	<pre>< 99.5% reduction.</pre>	See IK- 31 for comments.	58
						(continue	ed)

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No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IK- 33	Pentachloro- phenol	L	P	200 ppm	26% reduction in 120 hrs.	See IK-5 for comments.	66
IK- 34	Pentachloro- phenol	L	P	200 ppm	26% reduction in 120 hrs.	See IK-5 for comments.	92
IK- 35	Phenol	R	บ	150-200 ppm	90-95% reduction.	Activated sludge process.	90
1K-	Phenol	บ	I	19 ppm	< 99.9% reduction.	See IK-31 for comments.	58
IK- 37	Phenol	F	I	200 ppm	95% reduction.	Activated sludge process.	118
IK-	Phenol	F	I	5 ppm 18 ppm	71% reduction. 62% reduction.	Acclimated aerobic culture.	119
IK- 39	Pheno1	0	Ď	500 ppm	11.6% of TOD exerted after72 hrs of oxidation.		106
IK- 40	Phenol	0	D	500 ppm	0 ₂ uptake inhibited for first 24 hrs of oxidation. 41.2% TOD exerted in 144 hrs.		108
IK- 41	Phenol	В,С	I	120 ppm @ 500 gpm	< 200 ppb effluent conc.	Activated sludge process.	88
IK- 42	Pheno1	L	P	200 ppm	100% reduction in 8 hrs.	See IK- 5 for comments.	66
IK- 43	Phenol	Ü	I	500 ppm	100% reduction in 10 hrs.	See IK- 5 for comments.	92
IK- 44	p-Phenylazo- phenol	0	D	500 ppm	0 ₂ uptake inhibited.		108
IK- 45	Sodium Penta- chlorophenol	L	D	15 ppm	0% reduction.		120
IK- 46	2,3,5-Trichlo- rophenol	Ū	I	200 ppm	100% reduction in 55 hrs.	See IK- 5 for comments.	92
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TABLE C-1(continued)

Concentration Process: Biological Treatment (I)

Chemical Classification: Phenols (K)

		,			: Phenois (k)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
	2,3,5-Trichlo-rophenol	L	P·	200 ppm	100% reduction in 52 hrs.	See IK-5 for comments.	66
	2,4,5-Trichlo-rophenol	L	D	18.8 ppm	99% reduction in 6.5 days.	See IK-18 for comments.	115
IK- 49	2,4,6-Trichlo-rophenol	R	Ū	20 ppm	Biodegradable in 5 days.		90
IK-	2,4,6-Trichlo-rophenol	L	P	200 ppm	100% reduction in 50 hrs.	See IK- 5 for comments.	66
IK- 51	2,4,6-Trichlo- rophenol	0	S	1-10 ppm	0 ₂ uptake showed no inhibi- tory effect. 0 ₂ uptake inhibited.		102
IK- 52	2,4,6-Trichlo-rophenol	L	D	100ppm	99% reduction in 5 days.	See IK- 18 for comments.	115
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201

Concentration Process: Biological Treatment (I)

Chemical Classification: Phthalates (L)

No.	Chemical b	Descr Study Type ^C	Waste	Influent Char.	Results of Study	Comments	Ref
	Bis(2-ethylhex- yl) Phthalate		U	5 ppm	70-78% reduction.	Activated sludge process.	90
IL-	Butylbenzyl Phthalate	R	Ū		Biodegradable.		90
	Di-N-Butyl Phthalate	R	Ū		Biodegradable in an environ- mental system at a level of 200 ppm.		90
Į.	Diethyl Phthalate	R	U		Biodegradable.		90
	Di(2-ethylhex- yl) Phthalate	F	I		50-70% reduction.	Treated by aerated lagoon.	100
IL-	Dimethyl Phthalate	R	U		Biodegradable, no inhibition of bacteria at levels of 1000 ppm.		90
	Dimethyl Phthalate	U	S	215 ppb	100% reduction.		21
	Di-N-Octyl Phthalate	R	U		Biodegradable in an environ- mental system at a level of 63 ppm.		90
	Isophthalic Acid	U	P		95% reduction based on COD; rate of biodegradation 78.4 mg COD/g hr.	Activated sludge process.	81
IL- 10	Phthalimide	Ū	P		96.2% reduction based on COD; rate of biodegradation 20.8 mg COD/g hr.	Activated sludge process.	81
IL- 11	Phthalic Acid	U	P		96.8% reduction based on COD; rate of biodegradation 78.4 mg COD/g hr.	Activated sludge process.	81

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IM-	Anthracene	0	, D	500 ppm	Toxic or inhibitory for up to 24 hrs.		108
IM- 2	Benzanthracene	0	D	500 ppm	Slowly oxidized; 2.1% of TOD exerted in 144 hrs of oxidation.		108
IM- 3	Benzoperylene	R	Ü		Biodegradable from a conc. of 4×10^{-7} mg/1.		90
IM- 4	D-Chloramphe- nicol	U	P		86.2% reduction based on COD; rate of biodegradation 3.3 mg COD/g hr.	Activated sludge process.	81
IM- 5	α,α'-Diethyl- stilbenediol	0	D		O ₂ uptake inhibited.		108
IM- 6	9,10-Dimethyl- anthracene	0	D	500 ppm	O ₂ uptake was not inhibited. Up to 19.5% of TOD was exerted after 144 hr of oxidation.		108
IM- 7	9,10-Dimethyl- 1,2-benzan- thracene	0	D	500 ppm	Slowly oxidized; 12.7% of TOD exerted after 144 hr of oxidation.		
IM- 8	l,2-Diphenyl- hydrazine	F,C	D	341 ppb @ 45 MGD	28% reduction.	Activated sludge process.	81
IM- 9	7-Methyl-1,2- benzanthracene	0	D	500 ppm	O ₂ uptake inhibited at least 24 hrs.		108
1M- 10	20-Methyl- cholanthrene	0	D	500 ppm	Chemical showed both toxic or inhibitory effect & the ability to undergo slow biological oxidation.		108
IM- 11	Naphthalene	F	Ī		70-90% reduction.	Treated by aerated lagoon.	100
						(contin	ued)

TABLE C-1(continued)

			 				
a	. b		iption o	of Study			i
No.	Chemical b	Study Type ^C	Waste Type d	Influent Char.	Results of Study	Comments	Ref
IM- 12	Naphthalene	F	I		85-95% reduction.	Completely mixed aerated lagoon	101
IM- 13	Naphthalene	0	D	500 ppm	O ₂ uptake inhibited for 24 hrs.		108
IM- 14	Naphthalene	F	I	BOD load of 42 lb/day/ 1000 ft ³	85-95% reduction.	Activated sludge process.	56
						(conti	nued)

Concentration Process: Chemical Precipitation (II)

Chemical Classification: Aromatics (D)

_					14Omacrco (D)	•	
No.	Chemical b	<u>Descr</u> Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
II D- 1	Ethyl Benzene	R	D+P	153 ppb	56% reduction w/alum.	Chemical coagulation was followed by dual media filtration.	21
II D- 2	Nitrobenzene	R	D+P	160 ppb	34% reduction w/alum.	See IID-1 for comments.	21
						(contin	ned)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Halocarbons (F)

1	a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
- 1	I F- 1	Carbon Tetra- chloride	R	D+P	140 ppb	51% reduction w/alum.	Chemical coagulation was followed by dual media filtration.	21
	I F- 2	Trichloro- ethylene	R	D+P	103 ppb	40% reduction w/alum.	See IIF-1 for comments.	21
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				4				
- 1				1			(continu	ed)

Concentration Process:

Chemical Precipitation (II)

Chemical Classification: Metals (G)

No.	Chemical b	Descr Study Type C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
II G- 1	Antimony	P	s	600 ppb	62% reduction w/alum; 28% reduction w/lime. 65% reduction w/ferric chloride.	3 coagulants used: 220 ppm of alum @ pH=6.4. 40 ppm of ferric chloride @ pH=6.2; 415 ppm of lime @ pH=11.5; Chemical coagulation was followed by dual media filtration.	39
II G- 2	Arsenic	P	D+P	5 ppm @ 4 gpm @ pH=7.0	Iron system- 90% reduction; Low lime system- 80% reduction; High lime system- 76% reduction.	3 coagulant systems were used: Iron 'system used 45 ppm as Fe of Fe ₂ (SO ₄) ₃ @pH=6.0. Low lime system used 20 ppm as Fe of Fe ₂ (SO ₄) ₃ & 260 ppm of CaO @ pH=10.0. High lime system used 600 ppm of CaO @ pH=11.5. Chemical coagulation was followed by multimedia filtration.	63
II G- 3	Arsenic	F,C	D	2.5 ppb 3.3 ppb	56% reduction w/lime. 24% reduction w/lime.	Lime dose of 350-400ppm as calcium oxide @ pH=11.3.	64
II G- 4	Arsenic (As ⁺⁵)	R	U .	25 ppm 21 ppm	97% reduction by lime soften- ing. 94% reduction by precipita- tion w/alum.		90
						(continu	ed)

Concentration Process: Chemical Precipitation (II)
Chemical Classification: Metals (G)

	a No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
	II G-	Barium	F,C	D	81 ppb 81 ppb	49% reduction w/lime. 36% reduction w/lime.	See IIG-3 for comments.	64
	II G- 6	Barium	P	D+P	5 ppm 0 4 gpm 0 pH=7.0	Iron system- 94% reduction; Low lime sytem-99% reduction; High lime system-78% reduc- tion.	See IIG-2 for comments.	63
İ	II G- 7	Barium	P	S	500 ppb	79% reduction w/alum.	See IIG-1 for comments.	39
	11 G- 8	Beryllium	R	U	100 ppb	97.8% reduction by lime softening.		90
	II G- 9	Beryllium	P	S	100 ppb	98.1% reduction w/alum; 94% reduction w/ferric chlo- ride; 99.4% reduction w/lime.	See IIG-1 for comments.	39
	II G- 10	Bismuth	P	S	600 ppb 500 ppb	95.5% reduction w/ alum. 95.3% reduction w/lime. 94% reduction w/ferric chloride.	See IIG-1 for comments.	39
	II G- 11	Cadmium	P	S	700 ppb	45% reduction by ferric chloride.	See IIG-1 for comments.	39
•	II G- 12	Cadmium	P	D+P	5 ppm @ 4 gpm @ pH=7.0	Iron system- 93% reduction; Low lime system-95% reduction High lime system-98% reduc- tion.	See IIG-2 for comments.	63
•	II G- 13	Cadmium	F,C	D	29 ppb 9 ppb	92% reduction w/lime. 68% reduction w/lime.	See IIG-3 for comments.	64
					1			(continued)

TABLE C-1 (continued)

Concentration Process: Chemical Precipitation (II)

Chemical Classification: Metals (G)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
II G- 14	Chromium	L,C	Ĺ	5.2 ppm	26.9% reduction w/lime.	Lime dose of 50 ppm added.	16
II G- 15	Chromium	F,C	D	154 ppb 192 ppb	37% reduction w/lime. 54% reduction w/lime.	See IIG-3 for comments.	64
II G- 16	Chromium (Cr ⁺³)	P	S	700 ppb	97.6% reduction w/ferric chloride.	See IIG-1 for comments.	39
II G- 17	Chromium (Cr ⁺³)	P	D+P	5 ppm @ 4 gpm @ pH=7.0	Iron system - 99% reduction; Low lime system - 98% reduc- tion; High lime system - 98% reduction.	See IIG-2 for comments.	63
II G- 18	Chromium (Cr ⁺⁶)	P	S	700 ppb	64% reduction w/ferric chloride.	See IIG-1 for comments.	39
II G- 19	Chromium (Cr ⁺⁶)	P	D+P	5 ppm @ gpm @ pH=7.0	Iron system - 65% reduction; Low lime system - 40% reduc- tion; High lime system - 22% reduction.	See IIG-2 for comments.	63
II G- 20	Cobalt	P	S	500 ppb 800 ppb	18% reduction w/ferric chloride; 91% reduction w/lime. 49% reduction w/alum.	See IIG-1 for comments.	39
II G- 21	Copper	P	S	700 ppb	67% reduction w/alum.	See IIG-1 for comments.	39
II G- 22	Copper	L,C	S	4.6 ppm	97.8% reduction w/lime.	See IIG-14 for comments. (conti	16

TABLE C-1(continued)

Concentration Process: Chemical Precipitation (II)

Chemical Classification: Metals (G)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Re
II G- 23	Copper	P	D+P	5 ppm @ 4 gpm @ pH=7.0	Iron system- 95.6% reduction Low lime system-92.8% reduc- tion; High lime system- 84% reduction.	See IIG-2 for comments.	63
II G- 24	Copper	F,C	D	266 ppb 285 ppb	73% reduction w/lime. 93% reduction w/lime.	See IIG-3 for comments.	64
II G- 25	Copper	R	Ü	15 ppm	96% reduction.		90
II G- 26	Iron	L,C	s	10 ррт	99% reduction w/lime.	See IIG-14 for comments.	16
II G- 27	Iron	P	D+P	5 ppm @ 4 gpm @ pH=7.0	Iron system- 26% reduction; Low lime system-94% reduction	See IIG- 2 for comments.	63
II G- 28	Iron	F,C	D	179 ppb 325 ppb	91% reduction w/lime. 88% reduction w/lime.	See IIG- 3 for comments.	64
II G- 29	Lead .	L,C	S	4.9 ppm	100% reduction w/lime.	See IIG-14 for comments.	16
II G- 30	Lead	P	D+P	5 ppm @ 4 gpm @ pH=7.0	Iron system- 99% reduction; Low lime system-99% reduction High lime system-98% reduc- tion.	See IIG- 2 for comments.	63
II G- 31	Lead	F,C	D	40 ppb 19 ppb	43% reduction w/lime. 81% reduction w/lime.	See IIG- 3 for comments.	64
						`	(continued)

TABLE C-1 (continued)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Metals (G)

							T
No.	Chemical b	Descr Study	iption o	f Study Influent Char.	Results of Study	Comments	Ref.
		Type ^C	Type ^d	Char.	•		
II G- 32	Lead	R	υ	330 ppb	94.4% reduction w/lime.	Lime dose of 400 ppm added.	90
32 II	Lead	P	S	600 ppb	95.5% reduction w/alum.	See IIG-1	39
G-	read	P	ည	add poo	93.3% reduction w/alum.	for comments.	39
33						TOT COMMENCES.	1
II	Manganese	P	S	700 ppb	30% reduction w/alum.	See IIG-1	39
G-	•				·	for comments.	
34		<u> </u>					
II	Manganese	P	D+P	5 ppm @	Iron system- 18% reduction;	See IIG-2	63
G-		ł		4 gpm @	Low lime system-93% reduc-	for comments.	
35				pH=7.0	tion; High lime system-98%		1
II	Managaga	F,C	D	25 mmb	reduction.	See IIG-3	64
G-	Manganese	F,C	l D	35 ppb 38 ppb	87% reduction w/lime. 96% reduction w/lime.	for comments.	04
36				30 PPD	30% reduction wyrime.	TOT COMMENCS.	
II	Mercury	P	D+P	0.5 ppm	High lime system-70% reduc-	See IIG-2	63
G-	•			@ 4 gpm	tion.	for comments.	
37				@ pH=7.0		·	
II	Mercury	F,C	D	9 ppb	71% reduction w/lime.	See IIG-3	64
G- 38				1.2 ppb	25% reduction w/lime.	for comments.	
II	Mercury	P	S	500 ppb	70% reduction w/lime.	See IIG-1	39
G-	receuty	1		60 ppb	94% reduction w/alum.	for comments.	
39				50 ppb	98% reduction w/ferric		
1 1			1		chloride.		ł
II	Molybdenum	P	S	600 ppb	68% reduction w/ferric chlo-	See IIG-1	39
G-					ride; 0% reduction w/alum.	for comments.	
40			<u> </u>	500 ppb	0% reduction w/lime.		
						(continu	ıed)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Metals (G)

a	b			of Study			
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref.
		Type C	Type d	Char.			
II	Nickel	P	s	900 ppb	25% reduction w/alum.	See IIG-1	39
G-				**	· ,	for comments.	
41							
II	Nickel	L,C	S	4.8 ppm	100% reduction w/lime.	See IIG-14	16
G-						for comments.	
42				١			
II	Nickel	P	D+P	5 ppm @	Iron system- 10% reduction;	See IIG-2	63
G-				4 gpm @	Low lime system-94% reduc-	for comments.	1
43		İ		рн=7.0	tion; High lime system-97%		
					reduction.		
II	Nickel	R	U		52.4% reduction w/lime.	Lime dose of 400 ppm	90
G-		1				added.	
44	- n t			1001	750 2	See IIG-1	39
II	Selenium	P	S	100 ppb	75% reduction w/ferric chlo- ride.	for comments.	39
G- 45				500 ppb	35% reduction w/lime; 48%	TOT COMMENCS.	
45				300 ppb	reduction w/lime, 40%		
II	Selenium	F,C	D	<2.5 ppb	0% reduction w/lime.	See IIG-3	64
G-	Defenie	1		6.5 ppb	0% reduction w/lime.	for comments.	
46		1		FF	,		1
II	Selenium	R	U	100 ppm	80% reduction w/ferric	Ferric sulfate dose	90
G-		}		1	sulfate.	of 100 ppm.	1
47							
II	Silver	P	S	500 ppb	98.2% reduction w/ferric	See IIG-1	39
G-					chloride; 97.1% reduction	for comments.	
48					w/lime.		
				600 ppb	96.9% reduction w/alum.		
II	Silver	F,C	D	5.5 ppb	85% reduction w/lime.	See IIG-3	64
G-				13 ppb	38% reduction w/lime.	for comments.	
49	<u></u>		<u> </u>			(continu	ied)
	1	Ì		1) (contains	100,

Concentration Process: Chemical Precipitation (II)

Chemical Classification: Metals (G)

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a	b	1		of Study			
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref
		Type C	Type d	Char.			
II	Silver	R	Ų	500 ppm	96% reduction w/lime.		90
G-					·		ļ
50							
II	Thallium	R	U	500 ppb	54% reduction w/lime.		90
G-		İ					
51							
II	Thallium	P	s	600 ppb	30% reduction w/ferric chlo-	See IIG-1	39
G-		1			ride; 31% reduction w/alum.	for comments.	
52				500 ppb	60% reduction w/lime.		
II	Tin	P	S	500 ppb	98% reduction w/ferric chlo-	See IIG-1	39
G-					ride; 92% reduction w/lime.	for comments.	
53				600 ppb	95.3% reduction w/alum.		
II	Titanium	P	S	500 ppb	98% reduction w/ferric chlo-	See IIG-1	39
G-					ride; 95.5% reduction w/lime	for comments.	ļ
54				600 ppb	95.8% reduction w/alum.	,	
II	Vanadium	P	S	500 ppb	97.2% reduction w/ferric	See IIG-1	39
G-				l	chloride; 94% reduction w/	for comments.	
55					alum; 57% reduction w/lime.		
II	Zinc	P	S	2.5 ppm	1% reduction w/alum.	See IIG-1	39
G-						for comments.	
56							
II	Zinc	P	D+P	5 ppm @	Iron system- 63% reduction;	See IIG-2	63
G-]		4 gpm @	Low lime system-85% reduc-	for comments.	1
57				рн=7.0	tion; High lime system-76%		
					reduction.		
II	Zinc	L,C	S	6.4 ppm	100% reduction w/lime.	See IIG-14	16
G-						for comments.	
58				ļ			
					1		(continued)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Metals (G)

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	h		iption o	of Study			
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref.
		Type C	Type d	Influent Char.			
1		 					 -
]]	. 6						
II	Zinc	F,C	D	300 ppb	90% reduction w/lime.	See IIG-3	64
G-		ļ		380 ppb	37% reduction w/lime.	for comments.	
59 II	Zinc	R	บ	<u> </u>	40.60		ļ
G-	Zinc	R	U		40.6% reduction by sedimentation.		90
1 1	 				sedimentation.		
_60 II	Zinc	R	U		91.4% reduction w/lime.	Time dose of ACC	90
G-	Dino				J1:48 leddccion w/lime.	Lime dose of 400 ppm added.	90
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TABLE C-1(continued)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Pesticides (J)

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No.	Chemical b	<u>Descr</u> Study Type ^C	Waste .	of Study Influent Char.	Results of Study	Comments	Ref.
II J- 1	DDT	L,C	R+P	10 ppb	98% reduction w/alum.	Chemical coagulation was followed by sand filtration.	6
J- 2	Dieldrin	L,C	R+P	10 ppb	55% reduction w/alum.	See IIJ-1 for comments.	6
J- 3	Endrin	L,C	R+P	10 ppb	35% reduction w/alum.	See IIJ-1 for comments.	6
II J- 4	Lindane	L,C	R+P	10 ppb	<10% reduction w/alum.	See IIJ-1 for comments.	6
II J- 5	Parathion	L,C	R+P	10 ppb	5% reduction w/alum.	See IIJ-1 for comments.	6
II J- 6	2,4,5-T ester	L,C	R+P	10 ppb	65% reduction w/alum.	See IIJ-1 for comments.	6
			,				
						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Phthalates (L)

	a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
	II L- 1	Bis(2-ethyl- hexyl)Phtha- late	R	υ	0.5-3.5 ppb @ pH=10.0	80-90% reduction w/Al2(SO4)3		90
	II L- 2	Di-n-Butyl Phthalate	R	U	2.5-4.5 ppb @ pH=10.0	60-70% reduction w/Al ₂ (SO ₄) ₃		90
	T- 3	Dimethyl Phthalate	R	D+P	183 ppb	15% reduction w/alum.	Chemical coagulation was followed by dual media filtration.	21
2			CA sinterwork of the latter account for an					
							(continue	d)

Concentration Process: Chemical Precipitation (II)
Chemical Classification: Polynuclear Aromatics (M)

No.	b			of Study			Ť
No.	Chemical b	Study Type C	Waste Type d	Influent Char.	Results of Study	Comments	Rei
II M- 1	Acenaphthene	R	Ų	0.1-0.9 ppm	Precipitation w/alum.		90
II M- 2	Acenaphthylene	R	U	0.1-0.9 ppm	Precipitation w/alum.		90
II M- 3	Benzanthracene	R	U		Separable by gravity or sand filtration.		90
II M- 4	11,12-Benzo- fluoranthene	R	U		Separable by gravity or sand filtration.		90
II M- 5	1,12-Benzo- perylene	R	U		Separable by gravity or sand filtration.		90
II M- 6	Benzo(a)- pyrene	R	U		Separable by gravity or sand filtration.		90
II M- 7	2-Chloro- Napthalene	R	U	0.1-0.9 ppm	Precipitation w/alum.	,	90
M- 8	Chrysene	R	Ü		Separable by gravity or sand filtration.		90
M- 9	Naphthalene	R	Ū		Separable by gravity or sand filtration.		90
'	1	1		, ((continue	ed)

Concentration Process: Chemical Precipitation (II) Chemical Classification: Polynuclear Aromatics (M)

			ux oxuse	SIL LCGCION	Polynuclear Aromatics (M)		
No.	Chemical b	Descr Study Type C	Waste	f Study Influent Char.	Results of Study	Comments	Ref.
	2,3-o-Phenylene Pyrene	R	U		Separable by gravity or sand filtration.		90
II M- 11	Pyrehe	R	U	-	Separable by gravity or sand filtration.		90
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						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Reverse Osmosis (III) Chemical Classification: Alcohols (A)

	h	Descr	iption o	of Study			
No.	Chemical b	Study Type C	Waste Type d	Influent Char.	Results of Study	Comments	Ref
III A- 1	Ethanol	В	P	1000 ppm @ 150 mls	21.4% reduction w/CA membrane 70.3% reduction w/C-PEI membrane.	CA and C-PEl membranes operated at 600 psig and room temperature.	18
111 A- 2	Ethanol	L	Р	1000 ppm	80-100% reduction w/NS-200 membrane; 60-80% reduction w/NS-100-T membrane; 40-60% reduction w/AP & NS-100 membranes; 20-40% reduction w/CA3 & B-9 membranes; <20% reduction w/CA, CA-T, CAB, PBI, SPPO & B-10 membranes.		30
III A- 3	Methanol	В	P	1000 ppm @ 150 mls		See IIIA-1 for comments.	18
III A- 4	Methanol	L	P	1000 ppm	20-40% reduction w/B-9, NS- 200 & NS-100T membranes; <20% reduction w/B-10, AP, SPPO, PBI, NS-100 membranes; 0% reduction w/CA, CA-T, CAB & CA3 membranes.		30
III A- 5	i-Propanol	В	P	1000 ppm @ 150 mls	40.9% reduction w/CA membrane 88.1% reduction w/C-PEI mem- brane.	See IIIA-1 for comments.	18
111 A- 6	i-Propanol	L	P	1000 ppm	80-100% reduction w/NS-100, NS-100T, NS-200, AP, B-9 & B-10 membranes; 40-60% reduction w/CA-T, CA & CA3 membranes; 20-40% reduction w/SPPO, PBI & CAB membranes.	·	30
		•				(continu	ed)

Concentration Process: Reverse Osmosis (III) Chemical Classification: Aliphatics (B)

No.	Chemical b	Study	Waste	of Study Influent	Results of Study	Comments	Ref
		Type ^C	Туре а	Char.			
III B- 1	Acetic Acid	В	P	1000 ppm @ 150 ml	•	CA and C-PE1 membranes operated at 600 psig & room temperature.	18
III B- 2	Acetic Acid	L	P	1000 ppm	60-80% reduction w/AP, NS-200 & NS-100T membranes; 40-60% reduction w/NS-100 membrane; 20-40% reduction w/SPPO, B-9 & B-10 membranes; <20% reduction w/PBI, CA3, CAB, CA-T & CA membranes.		30
B- 3	Acetone	В	P	1000 ppm @ 150 ml	14.9% reduction w/CA membrane 81.8% reduction w/C-PEI membrane.	See IIIB-1 for comments.	18
III B- 4	Acetone	L	P	1000 ppm	80-100% reduction w/NS-200 & NS-100-T membrances; 60-80% reduction w/AP & NS-100 membranes; 40-60% reduction w/B-9 & B-10 membranes; 20-40% reduction w/CA3 membrane; <20% reduction w/SPPO, PBI, CAB, CA-T & CA membranes.		30
III B- 5	Dimethyl Sulf- oxide	В	P	250 ppm	88.2% reduction w/CA mem- brane; 63.3% reduction w/C-PEI membrane.	See IIIB-l for comments.	18
III B- 6	Formaldehyde	В	P	1000 ppm	21.9% reduction w/CA mem- brane; 56.7% reduction w/ C-PEI membrane.	See IIIB-1 for comments.	18
III B- 7	Formaldehyde	L	P	1000 ppm	60-80% reduction w/NS-200 membrane; 40-60% reduction w/AP, NS-100, CAB & NS-100-T	(continue	30

Concentration Process: Reverse Osmosis (III)

Chemical Classification: Aliphatics (B)

		Descr	intian	A C43			`
No.	Chemical b	Study		of Study	Describe of Gloci		
	CHEMICAL	Type C	Type d	Influent Char.	Results of Study	Comments	Ref.
		TAbe	туре	cnar.			
III					membranes; 20-40% reduction		
В-			٠		w/B-9, CA3 & CA-T membranes;		
7					<20% reduction w/CA, PBI,		
cont					SPPO & B-10 membranes.		
III	Glycerol	В	P	1000 ppm	89.9% reduction w/CA mem-	See IIIB-1	18
В-				@ 150 m1	brane; 97.8% reduction	for comments.	
8					w/C-PEI membrane.	•	
III	Glycerol	L	P	1000 ppm	80-100% reduction w/CA-T,		30
В-					CAB, CA3, NS-100, NS-100T,		}
9					NS-200, AP, B-9 & B-10 mem-		
					branes; 60-80% reduction		
					w/CA membrane; 40-60% re-		
					duction w/PBI membrane; 20-		}
					40% reduction w/SPPO membrane		
III	Methyl Acetate	В	P	1000 ppm	4.6% reduction w/CA membrane	See IIIB-1	18
B- 10				@ 150 ml	76.1% reduction w/C-PEI	for comments.	1
	W-417 2				membrane.		
III	Methyl Acetate	L	P	1000 ppm	60-80% reduction w/NS-200,		30
B- 11					NS-100-T & NS-100 membranes;		
					40-60% reduction w/B-9 mem-		
					brane; 20-40% reduction		
					w/B-10, AP & CA-T membranes;		
1					<20% reduction w/SPPO,PBI &		
					CA3 membranes; 0% reduction		
				<u> </u>	w/CA & CAB membranes.		
- 1							
,						(continue	ed)

Concentration Process: Reverse Osmosis (III)

Chemical Classification: Amines (C)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
III C- 1	Aniline	В	P	1000 ppm @ 150 ml		CA & C-PEI membranes operated at 600 psig & room temperature.	18
C- 2	Aniline	L	Þ	1000 ppm	80-100% reduction w/NS-100-T membrane; 60-80% reduction w/B-10, NS-200 & NS-100 membranes; 40-60% reduction w/B-9 membrane; 20-40% reduction w/AP, CA3 & CAB membranes; <20% reduction w/SPPO & PBI membranes; 0% reduction w/CA & CA-T membranes.	•	30
	,					·	
						(continue	d)

Concentration Process: Reverse Osmosis (III) Chemical Classification: Aromatics (D)

							'
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
III D- 1	Chlorobenzene	R	Ŭ _.	<360 ppm	97-100% reduction @ 50-100 kg/cm ² .		90
III D- 2	Dinitrobenzene	В	P	30 ppm @ 150 ml	7.2% reduction w/CA membrane 81.4% reduction w/C-PEI membrane.	CA & C-PEI membranes operated @ 600 psig & room temperature.	18
III D- 3	2,4-Dinitro- phenylhydra- zine	В	P	30 ppm @ 150 ml	3.2% reduction w/CA membrane 91.1% reduction w/C-PEI membrane.	See IIID-2 for comments.	18
III D- 4	Hexachloro- benzene	R	Ü	638 ppm	52% reduction.		90
III D- 5	Hydroquinone	В	P	1000 ppm	-2.5% reduction w/CA membrane 79.7% reduction w/C-PEI membrane.	CA & C-PEI membranes operated @ 600 psig & room temperature.	18
III D- 6	Hydroquinone	L	P	1000 ppm	80-100% reduction w/AP & NS-200 membranes; 60-80% reduction w/B-10, NS-100-T & NS-100 membranes; 40-60% reduction w/B-9 membrane; 20-40% reduction w/SPPO & CAB membranes; <20% reduction w/PBI & CA3 membranes; 0% reduction w/CA & CA-T membranes	i e	30
						(continu	ued)

Concentration Process: Reverse Osmosis (III)

Chemical Classification: Ethers (E)

a	. b			of Study			ĺ
No.	Chemical b	Study		Influent	Results of Study	Comments	Ref.
		Type c	Туре а	Char.			
III	bis(2-Chloro-	В	P	250 ppm	37.3% reduction w/CA mem-	CA & C-PEI membrane	18
E-	isopropyl)		· ·	@ 150 m1	brane; 94% reduction w/C-PEI	operated at 600 psig	
1	Ether				membrane.	& room temperature.	
III	Diethyl Ether	В	P	1000 ppm	9.5% reduction w/CA membrane	See IIIE-1	18
E-				@ 150 ml		for comments.	
2					membrane.		
III	Ethyl Ether	L	P	1000 ppm	80-100% reduction W/AP,		30
E-				:	NS-200, NS-100-T & NS-100		
3					membranes; 60-80% reduction		
					w/B-10 membrane; 40-60% re-		1
					duction w/B-9, SPPO & PBI		ļ
1					membranes; 20-40% reduction		1
					CAB & CA3 membranes; <20%		
					reduction w/CA-T & CA membranes.		
 					membranes.		
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TABLE C-1(continued)

Concentration Process: Reverse Osmosis (III) Chemical Classification: Halocarbons (F)

					Haiocarbons (1)		
No.	Chemical b	Descr Study Type ^C	Waste	f Study Influent Char.	Results of Study	Comments	Ref.
III F- 1	Trichloroace- tic Acid	В	P '	250 ppm @ 150 ml	49.3% reduction w/CA mem- brane; 25% reduction w/C-PEI membrane.	CA & C-PEI membrane operated at 600 psig & room temperature.	18
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	·					•	
						(continu	ed)

Concentration Process: Reverse Osmosis (III)
Chemical Classification: Metals (G)

		CHemic	ai Ciass	silication	: Metals (G)		
No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
III G- 1	Barium,	В	P	0.85 ppm 9.15 ppm	>86.7% reduction w/CA membrane >88.2% reduction w/CA membrane 97.8% reduction w/CA membrane >98.6% reduction w/CA membrane	CA membrane operated at 400 psig & 16-22°C.	18
III G- 2	Cadmium	В	P	0.10 ppm 0.96 ppm	90% reduction w/CA membrane 90% reduction w/CA membrane 99% reduction w/CA membrane 98.7% reduction w/CA membrane	See IIIG-1 for comments.	18
III G- 3	Chromic Acid	L,C	I		85% rejection over 200 hrs w/polybenzimidazole membrane.	Polybenzimidazole mem- brane operated at 1500 psl.	24
III G- 4	Chromium	В	Þ		97.6% reduction W/C-PEI mem- brane @ pH=8.0. 91.3% reduction w/C-PEI mem- brane @ pH=11.0.	C-PEI membrane operated at 600 psig & room temperature.	18
III G- 5	Chromium	В	P	1.01 ppm 8.65 ppm	96.9% reduction w/CA membrane 95.0% reduction w/CA membrane 93.2% reduction w/CA membrane 85.1% reduction w/CA membrane	See IIIG-1 for comments.	18
III G- 6	Copper	В	P		99.9% reduction w/C-PEI mem- brane @ pH=8.0 & 11.0.	See IIIG- 4 for comments.	18
III G- 7	Copper	В	P	0.7 ppm 6.25 ppm	97% reduction w/CA membrane 94.8% reduction w/CA membrane 99.6% reduction w/CA membrane 99.2% reduction w/CA membrane	See IIIG- l for comments.	18
III G- 8	Iron	В	P		100% reduction w/C-PEI mem- brane @ pH=8.0 & 11.0.	See IIIG-4 for comments.	18
						(continue	d)

Concentration Process: Reverse Osmosis (III)

Chemical Classification: Metals (G)

	3-	Descr	iption o	of Study			į
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref.
ĺ		Type C	Type d	Char.			
III	Lead	В	P	12.5 ppm	100% reduction w/C-PEI mem-	See IIIG-4	18
G-1					brane @ pH=8.0 & 11.0.	for comments.	
III	Lead	В	P	0.95 ppm	99.5% reduction w/CA membrane	See IIIG-1	18
G-		1		1.1 ppm	97.8% reduction w/CA membrane	for comments.	l
2				4.75 ppm	99.9% reduction w/CA membrane		l l
				9.3 ppm	97.8% reduction w/CA membrane		
III	Nickel	В	P		92.8% reduction w/C-PEI mem-	See IIIG-4	18
G-		1		1	brane @ pH=8.0.	for comments.	
3		1		12.5 ppm	97.6% reduction w/C-PEI mem-		
		1			brane @ pH=11.0.		1
III	Zinc	В	P	12.5 ppm	96.6% reduction w/C-PEI mem-	See IIIG-1	18
G-					brane @ pH=8.0.	for comments.	
4				12.5 ppm	100% reduction w/C-PEI mem-	Ī	1
					brane @ pH=11.0.		
III	Zinc	В	P	9.4 ppm	96.9% reduction w/CA membrane	See IIIG-1	18
G-				10.0 ppm	98.6% reduction w/CA membrane	for comments.	
5			ļ	31.4 ppm	98.8% reduction w/CA membrane		
				32.8 ppm	99.5% reduction w/CA membrane		
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Concentration Process: Reverse Osmosis (III) Chemical Classification: Pesticides (J)

No.	Chemical b	Descr Study Type C		of Study Influent Char.	Results of Study	Comments	Ref.
J- 1	Aldrin	В	P	142 µg	100% reduction w/CA membrane 100% reduction w/C-PEI mem- brane.	CA & C-PEI membranes operated at 600 psig & room temperature.	18
III J- 2	Atrazine	В	P	1102 µg	84% reduction w/CA membrane 97.8% reduction w/C-PEI mem- brane.	See IIIJ-1 for comments.	18
J- 3	Captan	В	P	689 µg	98.8% reduction w/CA membrane 100% reduction w/C-PEI mem-brane.	See IIIJ-1 for comments.	18
III J- 4	DDE	В	P	69 µg	100% reduction w/CA membrane 100% reduction w/C-PEI mem- brane.	See IIIJ-1 for comments.	18
J- 5	DDT	В	P	42 μg	100% reduction w/CA membrane 100% reduction w/C-PEI mem-brane.	See IIIJ-1 for comments.	18
III J- 6	Diazinon	В	P	474 μg	98.3% reduction w/CA membrane 88.1% reduction w/C-PEI mem- brane.	See IIIJ-1 for comments.	18
J- 7	Dieldrin	В	P	321 µg	99.9% reduction w/CA membrane 100% reduction w/C-PEI mem-brane.	See IIIJ-1 for comments.	18
J- 8	Heptachlor	В	P	145 µg	100% reduction w/CA & C-PEI membranes.	See IIIJ-1 for comments.	18
J- 9	Heptachlor- epoxide	В	P		99.8% reduction w/CA & C-PEI membranes.	See IIIJ-1 for comments.	18
III J- 10	Lindane	В	P		99.5% reduction w/CA membrane 99.0% reduction w/C-PEI mem-brane.	See IIIJ-1 for comments.	18
						(continue	- <u></u>

Concentration Process: Reverse Osmosis (III) Chemical Classification: Pesticides (J)

	h		iption c	of Study			
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref.
		Type C	Type d	Char.			
III	Malathion	В	P	1058 μg	99.2% reduction w/CA membrane	See IIIJ-1	18
J-	natachion		Ι,	1050 μg	99.7% reduction w/C-PEI mem-	for comments.	
11					brane.		
III	Methyl	В	P	913 μg	99.6% reduction w/CA & C-PEI	See IIIJ-1	18
J- 12	Parathion				membranes.	for comments.	
III	Parathion	В	P	747 μg	99.9% reduction w/CA membrane	See IIIJ-1	18
J-			_		99.8% reduction w/C-PEI mem-	for comments.	İ
13					brane:		
III	Randox	В	P	327 μg	72% reduction w/CA membrane	See IIIJ-1	18
J- 14		ļ			98.6% reduction w/C-PEI mem-	for comments.	
	Trifluralin	В	P	1579 μα	brane. 99.7% reduction w/CA membrane	See IIIJ-1	18
J-	ILLLIULALIM		P	13/9 μg	100% reduction w/C-PEI mem-	for comments.	1 10
15					brane.	zoz commerces	Ì
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TABLE C-1 (continued)

Concentration Process: Reverse Osmosis (III) Chemical Classification: Phenols (K)

a No.	Chemical b	Descr Study Type C		of Study Influent Char.	Results of Study	Comments	Ref.
III K- 1	2-Chlorophenol	R	บ		66.3% reduction.		90
III K- 2	4-Nitrophenol	R	υ		Removable by reverse osmosis.	·	90
III K- 3	Phenol	R	U		17.8% reduction.		90
III K- 4	Phenol	В	P	1000 ppm	-5.7% reduction w/CA membrane 76.5% reduction w/C-PEI membrane.		18
K- 5	Pheno1	P	S	1-100mg/1 each of phenol, resorcin- ol, o- cresol, catechol	In excess of 90% separation at pH 8-10 w/optimum at pH 9 at flux rate of about 70 gpd/ft ² . Results indicate that hyperfiltration (reverse osmosis) produced higher rejection & flux rates than ultrafiltration. Increasing pressure improves rejection slightly & flux rate greatly. Increasing pH increased rejection w/little effect on flux rate. Conc. had little effect on either rejection or flux rate.	Size: 60-130 gpd/ft ² flux. Duration: 0-60hrs Pressure: 250-950 psig. Velocity: 15 fps. Mem- branes: Hydrous Zr (IV) oxide-PAA membrane on carbon stainless steel & selas support.	54
						(continue	ed)

Concentration Process: Ultrafiltration (IV)

Chemical Classification: Aromatics (D)

	VIIII OLABOLLIOADI.						<u> </u>
No.	Chemical b	<u>Descr</u> Study Type ^C	iption o Waste Type d	f Study Influent Char.	Results of Study	Comments	Ref.
IV D-	TNT (accounted for 90% of TOC)	L,C	I+P	20 ppm TOC @ pH=11.0 200 ppm TOC @ pH=11.0	80% TOC reduction by PSAL (Millipore) noncellulose membrane. 93% TOC reduction by PSAL (Millipore) noncellulose membrane.	TDS conc. was 1200 ppm. Average pressure: 25-60 psi. Estimated cost for full scale operation was \$1.85/1000 gal	
						(continu	ed)

Concentration Process: Ultrafiltration (IV)

Chemical Classification: Metals (G)

	· · · · · · · · · · · · · · · · · · ·	Official	ar cras.	errreacton	: Metais (G)		
No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref
IV G- 1	Copper	C,P	I.	0.44 ppm	0.08 ppm effluent conc.		59
IV G- 2	Iron	C,P	I	6.8 ppm	1.0 ppm effluent conc.		59
IV G- 3	Manganese	C,P	I	4.9 ppm	0.52 ppm effluent conc.		59
IV G- 4	Zinc	C,P	Ι	1.8 ppm	0.38 ppm effluent conc.		59
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	,						
						(cont.	inued)

Concentration Process: Ultrafiltration (IV)

Chemical Classification: Phenols (G)

_						Flichors (g)		
'	No.	Chemical b	Descr Study Type ^C		f Study Influent Char.	Results of Study	Comments	Ref.
233	IV G- 1	Phenols	P	S.	1-100 ppm each of phenol, resorcin- ol, o- cresol, catechol	pH 10; rejection increased as pH increased. Ionic state of solute rather than membrane material controlled rejection rate. Increased	Velocity: 15 fps	54
							(continu	ed)

TABLE C-1 (continued)

Concentration Process: Stripping (V)
Chemical Classification: Aliphatics (B)

		Docar	intion o	f Study			
No.	Chemical b	Study Type C	Waste Type d	Influent Char.	Results of Study	Comments	Ref.
VB- 1	Acrylonitrile	R	Ų		Flash vaporization from water by high pressure discharge.		90
		And the state of t					
						,	
						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Stripping (V)
Chemical Classification: Aromatics (D)

						•
cal b	Study Waste Influent Type C Type C Char.		Influent	Results of Study	Comments	Ref.
9	R	U.		Air & steam strippable.		90
9 :	C,P	s	0.13 gpm flow	95-99% reduction by steam stripping.	Estimated cost of \$3.35/1000 gal based on 0.03 MGD	13
benzene	R	U		Steam strippable.		90
benzene	F,C	D	$0.66 \text{ M}^3/\text{s}$ flow	60% reduction by air stripping.		64
loro- e	R	U		Air & steam strippable.	:	90
loro-	R	U		Steam strippable.		90
chloro-	F,C	D	$0.66 \text{ M}^3/\text{s}$	70% reduction by air strip- ping.		64
chloro- e	F,C	D		80% reduction by air strip- ping.		64
chloro- e	F,C	D		90% reduction by air strip- ping.		64
enzene	F,C	D		80% reduction by air strip- ping.		64
enzene	R	U		Air & steam strippable.		90
enzene	P,C	s	0.13 gpm flow	stripping.	See VD- 2 for comments.	13
loro- e	R	U		Steam strippable.		64
loro-				flow	flow stripping.	flow stripping. for comments.

TABLE C-1 (continued)

Concentration Process: Stripping (V)
Chemical Classification: Aromatics (D)

No.	Chemical b	Descr Study Type ^C	iption o Waste Type d	of Study Influent Char.	Results of Study	Comments	Ref.
VD- 14	Nitrobenzene	R	ָט	450-2160 ppm	Steam strippable.		64
VD- 15	Styrene	P,C	S	0.13 gpm	98-99% reduction by steam stripping.	See VD-2 for comments.	13
VD- 16		P,C	S	0.13 gpm flow	73-92% reduction	See VD-2 for comments.	13
VD- 17		R	ប		Air & steam strippable.		90
	1,2,4-Trichlo- robenzene	F,C	D	0.66 M ³ /s	50% reduction by air strip- ping.		64
VD- 19	1,2,4-Trichlo- robenzene	R	U		Steam strippable.		90
					× .		
					(continu	 :ed)	

TABLE C-1 (continued)

		·				
No.	Chemical b	Descr Study Type C	Waste	of Study Influent Char.	Results of Study Comments	Ref
VF- 1	Bromodichlo- romethane	R	Ū		Air & steam strippable.	90
VF- 2	Bromomethane	R	Ū		Air strippable. Gas at STP	90
VF- 3	Chloral	P,C	I	0	Overhead Overhead Bottom flow (% Conc. Conc. of feed) (ppm) (ppm) (ppm) COD - 15100 ppm pH - 0.1 acidity - 102312 ppm C1-116,127 ppm Numerous other halogores overhead ratio 2.5 with 2301.6 434.4 0.9:1 reflux to overhead ratio	
/F- 4 /F-	Chloroethane	R	Ü		90% evaporation from H ₂ 0-79 min with air stripping.	90
5	Chloroethy- lene	R	U		Air strippable Gas at STP	90
/F- 6	Chloroform	P,C		140.3 ppm @ 250ml/min feed rate	2.3 1185.1 0 2.8 882.4 0	95

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type ^C	Waste Type	f Study Influent Char.	Results of Study	Comments	Ref.
VF- 6 cont	,				Overhead Overhead Bottom flow (% Conc. Conc. of feed) (ppm) (ppm) 2.3 with 412.3 0 1.4:1 reflux to overhead ratio 2.5 with 1124.3 64.7 1.4:1 re-		
					flux to overhead ratio		
VF- 7	Chloromethane	R	Ū		Air strippable.	Gas at STP	90
VF-	Dibromochloro- methane	R	Ü		Air & steam strippable.		90
VF- 9	1,1-Dichloro- ethane	R	Ü		90% evaporation from H_20 - 109 min with air stripping.		90
VF- 10	1,2-Dichloro- ethane	R	U		Air & steam strippable.		90
						(continu	ed)

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Resu	lts of Stu	dy	Comments	Ref.
	1,2-Dichloro- ethane	P,C	Ţ	1583.3ppm @ 250 ml/ min feed rate		Overhead Conc. (ppm) 350.8 269.7 465.0 1320.9	Bottom Conc. (ppm) 373.7 1255.4 14.8 16.1	See VF-3 for comments.	95
	1,1-Dichloro- ethylene	R	Ü			m strippab	le.		90
VF-	1,2-trans-Di- chloroethylene	R	Ü			ation from h air stri			90
VF-	1,1-Dichloro- ethylene	P,C	I	61.5 ppm @ 250 ml/ min feed rate	Overhead	Overhead Conc. (ppm) 124.4 111.2 179.9	Bottom Conc. (ppm) 32.8 0	See VF-3 for comments.	95

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type ^C	iption o Waste Type d	Influent	Resu	lts of Stu	dy	Comments		Ref.
VF- 15	Dichloromethane	P,C		800.9 ppm @ 250 ml/ min feed rate	flow (%	Overhead Conc. (ppm) 3511.8 3277.0 2736.5 1183.0	Bottom Conc. (ppm) 114.1 89.5 175.6 296.3	See VF-3 for	comments.	95
								,	(continue	ed)

TABLE C-1(continued)

Concentration Process:

Stripping (V)
Halocarbons (F) Chemical Classification:

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study Comments	Ref.
VF- 16	Dichlorometh- ane	R	υ,		90% evaporation from H ₂ 0-60 min with air stripping.	90
VF- 17	l,2-Dichloro- propane	R	U		Air & steam strippable.	90
VF- 18	1,2-Dichloro- propylene	R	U		Air & steam strippable.	90
19 VF-	Ethylene Dichloride	P,C	I	1593 ppm @ 250ml/min feed rate	Overhead Overhead Bottom flow (% Conc. Conc. of feed) (ppm) (ppm) 2.3 4383.5 42.2 2.8 4105.5 64.5 5.1 4731.5 43.1 2.3 with 3654.5 38.6 1.4:1 re-flux to overhead flow 2.5 with 5541.3 436.4 0.9:1 re-flux to overhead ratio	95
					(continue	 ad)

TABLE C-1 (continued)

a No.	Chemical b	Descr Study Type C		f Study Influent Char.	Resu	ılts of Stı	udy	Comments	Ref.
VF- 20	Ethylene Dichloride	P,C	I .	Average conc. of 4512 ppm e ave. feed rate of 325ml/min	Overhead flow (ml/min) 20.8	Average Overhead Conc. (ppm) 21.6	Bottom Conc. (ppm)	Wastewater quality: COD - 615 ppm TC - 1703 ppm pH - 11.2 Alkalinity - 4840 ppm C1 - 6564 ppm	95
VF- 21	Ethylene Dichloride	P,C	I	8700 ppm @ 10 gpm flowrate	99% reduct				66
VF- 22	Hexachloro- butadiene	R	Ū		Air & stea	am strippa	ble.		90
VF- 23	Hexachloro- cyclopenta- diene	R	U		Polymerize	es with he	at.		90
VF- 24	Perchloro- ethylene	P,C	I	feed rate	flow (% of feed) 2.3 No 2.8 2.5 with 0.9:1 reflux to overhead ratio	Overhead Conc. (ppm) of reported 50.2 9.6	0	See VF-3 for comments.	95
VF- 25	1,1,1,2-Tetra- chloroethane	P,C	I	512.8ppm @ 250ml/min feed rate	Overhead flow (% of feed) 2.3 2.8	Overhead Conc. (ppm) 189.8 393.8	Bottom Conc. (ppm) 0 0.84	See VF-3 for comments. (continu	95 ed)

TABLE C-1 (continued)

Т		_					······································		i	
a	_, _ b			f Study			_		ļ	_
No.	Chemical b	Study	Waste	Influent Char.	Resi	ılts of Stı	ıdy	Comments		Ref.
		Туре ^{°С}	Туре ч	Char.						
VF-					Overhead	Overhead	Bottom			
25			,		flow (%	Conc.	Conc.		1	1
cont	•		,		of feed)	(ppm)	(ppm)			1
					5.1	22.7	0			1
					2.3 with	25.8	0.5			1
					1.4:1 re-					l
					flux to				ĺ	1
					overhead				İ	1
					ratio			ļ	ļ	1
		1			2.5 with	392.5	1.6			
					0.9:1 re-					
					flux to			İ	1	
1 1					overhead					
					ratio					
VF- 26	1,1,2,2-Tetra-	P,C	I	14.9 ppm	Overhead	Overhead	Bottom	See VF-3		95
26	chloroethane	'		e	flow (%	Conc.	·Conc.	for comments.	1	
				250ml/min	of feed)	(ppm)	(ppm)			
		1		feed rate		14.9	32.7			
					2.8	121.7	49.5			
					5.1	444.4	78.4			
	•				2.3 with	8.7	0	1		
		ļ	1		1.4:1 re-					
		ļ			flux to					
		-			overhead					
				ļ.	ratio					
					2.5 with	24.2	0.1			
					0.9:1 re-		0.1			
					flux to	•				
					overhead			,		
					ratio					l .
		1	 						(continue	d)

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study Comments	Ref.
27	Tetrachloro- ethylene	R	υ,		Air & steam strippable, 90% evaporation from H ₂ O - 72 min	90
	Tetrachloro-	R	ט		Air & steam strippable, 90%	90
	methane				evaporation from H ₂ O - 97 min	
VF- 29	Tribromomethane	R	υ		Air & steam strippable.	90
	1,1,1-Trichlo- roethane	R	υ		Air & steam strippable.	90
1 1	1,1,1-Trichlo- roethane	P,C	I	50.92 ppm @ 250 ml/ min feed rate	of feed) (ppm) (ppm) 2.5 with 173.4 41.6 0.9:1 re- flux to overhead	95
	1,1,2-Trichlo-	R	U		ratio Air & steam strippable, 90%	90
	roethane		ļ		evaporation from H ₂ O-102 min	
l l	1,1,2-Trichlo- roethane	P,C	I	14.14 ppm @ 250 ml/ min feed rate	Overhead Overhead Bottom flow (% Conc. Conc. of feed) (ppm) (ppm) 2.3 24.6 0.19 2.8 34.0 0 5.1 76.5 0 2.3 with 42.4 0 1.4:1 reflux to overhead ratio	95
			l		(continue	d)

TABLE C-1 (continued)

	a No.	Chemical b	<u>Descr</u> Study Type ^C		of Study Influent Char.	Resul	ts of Stu	dy	Comments		Ref.
	VF- 33 cont						Overhead Conc. (ppm) 66.1	Bottom Conc. (ppm) 0			
	VF-	Trichloro-	R	Ū		Air & steam					90
245	34 VF- 35	ethylene Trichloro- ethylene	P,C	I	250ml/min feed rate	flow (% of feed) 2.3 2.8 5.1 2.3 with 1.4:1 reflux to overhead ratio 2.5 with 0.9:1 reflux to	Overhead Conc. (ppm) 640.8 567.0 627.4 640.8		See VF-3 for comments.		95
	VF- 36	Trichloro- methane	R	U		overhead ratio Air & steam evaporation				(continu	90 ed)

Concentration Process: Stripping (V) Chemical Classification: Phenols (K)

		·		IIICacion		nois (K)		•	
a No.	Chemical	Descr	iption o	f Study Influent Char.		Results of Study	Comments		Ref.
	Onemiz da i	Study Type C	Type d	Char.		Acsults of Study	Commencs		Rer.
VK-	Phenol	R	U		Steam	strippable.			90
VK- 2	Chlorophenol	R	Ü		Steam	strippable.			90
							•		
					1				
	,						,		
		1	Į į		1			(continued	1)

TABLE C-1 (continued)

Concentration Process: Stripping (V)
Chemical Classification: Polynuclear Aromatic (M)

			rolyndcieal Alomatic (H)		
No. Chemical b	Description of Study Waste Type Type	of Study Influent Char.	Results of Study	Comments	Ref.
M- Naphthalene			Air stripping by 50:1 volumes of air.		90
,			·		
			·		
			`		
				(continu	ıed)

TABLE C-1 (continued)

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Alcohols (A)

	a No.	Chemical b	Descri Study Type C	iption o Waste Type d	f Study Influent Char.	Results of Study	Comments	Ref.
	VII A- 1	Ethanol	L,C	I	286 ppm	7% reduction.	Extraction of neutral- ized oxychlorination wastewater using 2-ethyl- hexanol (S/W=0.106); RDC extractor used.	27
248								
					,	S 196 S		
		·						
							(continue	ed)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Aliphatics (B)

اد	h	Descr	iption o	of Study	•		
No.	Chemical b	Study		Influent	Results of Study	Comments	Ref
		Type ^C	Type d	Char.			
VII	Acrolein	R	U		Extractable w/xylene. Sol-		90
в-					vent recovery by azeotropic		
1					distillation.		
VII	Acrylonitrile	R	Ū		Extractable w/ethyl ether.		90
В-							
2		ļ					
VII	Isophorone	R	ט		Extractable w/ethyl ether.		90
B- 3		ļ				•	İ
VII	Methyl Ethyl	L,C	I	12200ppm	69% reduction.	Sequential extraction of	27
В-	Ketone	-/-	1 -	@ 3.21	ost reduction.	waste water from lube-	1 2 '
4				gal/hr		oil refining using butyl	
ĺ						acetate (S/W=0.10) &	
						isobutylene (S/W=0.101);	
						RDC extractor used.	L
VII	Methyl Ethyl	L,C	I	12200ppm	88% reduction.	Sequential extraction of	27
B-	Ketone			@ 3.21		waste water from lube-	
5				gal/hr		oil refining using butyl	
						acetate (S/W=0.10) & isobutylene (S/W=0.101);	ļ
	•					RDC extractor used.	
						The extractor used.	-
			1			·	1
							1
					,	,	1
						(continue	ed)
	ı	1	1	1	•	1	4

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Aromatics (D)

No.	Chemical b	<u>Descr</u> Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
VII D- 1	Benzene	R	U		Extractable w/suitable solvent.		90
VII D- 2	Benzene	L,C	I	290 ppm @ 3 gal/hr	97% reduction.	Extraction of waste- water from styrene man- ufacture using isobuty- lne (S/W=0.107); RDC extractor used.	
VII D- 3	Benzene	L,C	I	71 ppm @ 4.6 gal/ hr	96% reduction.	Extraction of ethylene quench wastewater using isobutylene (S/W=0.101) RDC extractor used.	27
VII D- 4	Benzene	L,C	I	81 ppm @ 4.6 gal/ hr	97% reduction.	Extraction of ethylene quench wastewater using isobutane (S/W=0.097); RDC extractor used.	27
VII D- 5	Chlorobenzene	R	Ü	600 ppm	3 ppm effluent conc. using chloroform solvent.		90
VII D- 6	o-Dichloro- benzene m- p-	R	Ü		Extractable w/suitable solvent.	^	90
VII D- 7	2,4-Dinitro- toluene	R	U		Extractable w/suitable solvent.		90
VII 8	2,6-Dinitro- toluene	R	U		Extractable w/suitable solvent.		90
						(continue	d)

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Aromatics (D)

Chemical b Ethylbenzene Ethylbenzene	Study Type ^C	Waste	of Study Influent Char.	Results of Study 97% reduction.	Comments See VIID-2 for comments.	Ref
Ethylbenzene Ethylbenzene	Type ^C	Type d	Char.	-	See VIID-2	
Ethylbenzene				97% reduction.	1	27
_	R	Ü			1	
Hexachloro-		1		Extractable w/suitable solvent.	·	90
benzene	R	U		Extractable w/suitable solvent.		90
Nitrobenzene	R	U		Extractable w/suitable solvent.		90
Styrene	L,C	· I		>93% reduction.	See VIID-2 for comments.	27
roluene	R	ט		Extractable w/suitable solvent.		90
•	L,C	I	41-44ppm @ 4.6 gal/hr	94%-96% reduction.	See VIID-3 & 4 for comments.	27
l,2,4-Tri- hlorobenzene	R	ט		Extractable w/suitable solvent.	·	90
	L,C	I		>97% reduction.	See VIID-3 for comments.	27
Xylene	L,C	I		>97% reduction.	See VIID-4 for comments.	27
N I	Styrene Styrene Coluene Coluene And Andrew Coluene Coluene Coluene Coluene Coluene Coluene Coluene Coluene Coluene	Styrene L,C Coluene R Coluene R Coluene R Coluene L,C Coluene L,C Coluene L,C Coluene L,C	Styrene L,C I Coluene R U Coluene R U Coluene L,C I L,2,4-Tri-R hlorobenzene L,C I	Styrene L,C I Coluene R U Coluene L,C I 41-44ppm @ 4.6 gal/hr hlorobenzene Sylene L,C I	Extractable w/suitable solvent. Coluene Coluene L,C I Extractable w/suitable solvent. Coluene L,C I 41-44ppm (e) 4.6 (gal/hr L,2,4-Tri- hlorobenzene L,C I Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent.	Extractable w/suitable solvent. Styrene L,C I >93% reduction. See VIID-2 for comments. Coluene R U Extractable w/suitable solvent. Coluene L,C I 41-44ppm 94%-96% reduction. See VIID-3 & 4 for comments. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent. Extractable w/suitable solvent.

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Ethers (E)

		Descr	iption o	f Study			
No.	Chemical b	Study Type C	Waste	Influent Char.	Results of Study	Comments	Ref.
VII E- 1	bis-Chloro- ethyl Ether	R	Ų		Extractable w/ethyl ether & benzene.		90
VII E- 2	bis-Chloro- isopropyl Ether	R	υ		Extractable w/ethyl ether & benzene.		90
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						(continu	ed)

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Halocarbons (F)

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a No.	Chemical	Descr Study		of Study Influent	Results of Study	Comments	Ref
NO.	CHEMICUI	Type C		Char.	Results of Study	Commence	NC I
VII Fī	Bromodichlo- romethane	R	Ŭ		Soluble in most organics.		90
VII F- 2	Bromomethane	R	U		Soluble in most organics.		90
VII F- 3	Chloral Hydrate	L,C	Ι	15200 ppm	49% reduction.	Extraction of neutral- ized oxychlorination wastewater using 2- ethylhexanol (S/W=0.106) RDC extractor used.	27
VII F- 4	Chloroethane	R	ΰ		Extractable w/alcohols and aromatics.		90
VII F- 5	Chloroethylene	R	Ü		Soluble in most organics.		90
VII F- 6	Chloromethane	R	Ü		Soluble in most organics.	·	90
VII F7	Dibromochloro- methane	R	Ü		Extractable w/organics, ethers and alcohols.		90
VII F- 8	Dichlorodi- fluoromethane	R	Ū		Extractable w/organics, ethers and alcohols.		90
VII F- 9	1,1-Dichloro- ethane	R	Ü		Extractable w/alcohols and aromatics.		90
						(continue	ed)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Halocarbons (F)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
1 .	1,2-Dichloro- ethane	R	ָ ט		Extractable w/alcohols and aromatics.		90
1 1	Dichloro- ethylene	L,B	I	49 ppm	Kerosene effluent conc 2 ppm; C ₁₀ -C ₁₂ effluent conc 1 + ppm	Solvent extraction used separatory funnel w/kerosene & C ₁₀ -C ₁₂ hydrocarbon solvents at 7:1 solvent to wastewater ratio.	95
	Dichloro- ethylene	L,C	I	1500 ppm	>99% reduction.	See VIIF-3 for comments.	27
1	l,1-Dichloro- ethylene	R	Ŭ		Extractable w/alcohols, aromatics and ethers.		90
	1,2-trans-Di- chloroethylene	R	U		Soluble in most organics.		90
VII F-	Dichloromethane	R	U		Soluble in most organics.		90
	1,2-Dichloro- propane	R	Ū		Soluble in most organics.		90
	1,2-Dichloro- propylene	R	Ü		Soluble in most organics.		90
	_					(continue	ed)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Halocarbons (F)

a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
VII F- 9	Ethyl Chloride	L,B	I	3 ppm	Kerosene effluent conc 1 ppm; C_{10} - C_{12} hydrocarbon effuent - 1 + ppm.	Solvent extraction used separatory funnel w/ kerosene & C ₁₀ -C ₁₂ hydrocarbon solvents at 7:1 solvent to waste-water ratio.	95
F- 10		L,C	I	1640 ppm	21% reduction.	See VIIF-3 for comments.	27
F- 11		L,B	I	320 ppm	No detectable conc. in kerosene effluent; C_{10} - C_{12} hydrocarbon effluent - 1 + ppm.	See VIIF-9 for comments.	95
F- 12	·	P,C	I	23-1804 ppm @ 2.76-3.76 1/min	A 5.5:1 water to solvent ratio gave 94-96% reduction. C ₁₀ -C ₁₂ paraffin solvent at 5:1 to 16.5:1 water to solvent ratio showed 94-99% reduction	other halocarbons in- cluding 30-350 ppm l,1,2-trichloroethane	
i	Hexachloro- butadiene	R	υ		Soluble in most organics.		90
	Hexachloro- ethane	R	ט		Extractable w/aromatics, alcohols and ethers	(continue	90

Concentration Process: Solvent Extraction (VII) Chemical Classification: Halocarbons (F)

					<u> </u>		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
	Pentachloro- ethane	L,B	, I	10 ppm	Kerosene effluent conc 2 ppm; No detectable conc. in C10-C12 hydrocarbon effluent.	See VIIF-9 for comments.	95
F- 16		L,B	Ι	14 ppm	Kerosene effluent conc 2 ppm; C ₁₀ -C ₁₂ hydrocarbon effluent conc 1 ppm.	See VIIF-9 for comments.	95
F- 17		L,B	I	148 ppm	Kerosene effluent conc 7 ppm; C ₁₀ -C ₁₂ hydrocarbon effluent conc 6 ppm.	See VIIF-9 for comments.	95
F- 18		R	Ü		Extractable w/aromatics, alcohols and ethers.		90
F- 19		R	บ		Soluble in most organics.		90
F- 20	I	R	Ū		Soluble in most organics.		90
F- 21	Tribromomethane		U		Soluble in most organics.		90
F- 22		L,B	Ι	75 ppm	Kerosene effluent conc 2 ppm; C ₁₀ -C ₁₂ hydrocarbon effluent conc 1 ppm.	See VIIF-9 for comments.	95
F- 23	<u> </u>	R	Ü		Extractable w/alcohols and aromatics.		90
	1,1,2-Trichlo- roethane	R	U		Extractable w/aromatics, methanol and ethers.	,	90
						(c	ontinued)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Halocarbons (F)

				STITCACTON.	natocarbons (r)		
No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref
VII F- 25	Trichloro- ethylene	L,B	I	24 ppm	Kerosene effluent conc 6 ppm; C ₁₀ -C ₁₂ hydrocarbon effluent conc 5 ppm.	See VIIF- 9 for comments.	95
VII F- 26	Trichloro- ethylene	R	Ü		Soluble in most organics.		90
VII F- 27	Trichloro- fluoromethane	R	Ü		Extractable w/alcohol, ether and organics.		90
VII F- 28	Trichloro- methane	R	ΰ		Soluble in most organics.		90
VII F- 29	Vinylidene Chloride	L,B	I	13 ppm	Kerosene effluent conc l ppm; C ₁₀ -C ₁₂ effluent conc l ppm.	See VIIF-9 for comments.	95
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					·	r	
				1		(contir	nued)

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Metals (G)

No. Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref
VII Mercury G- 1	R	Ü	2 ppm	99% reduction w/high molec- ular weight amines & quartenary salts.		90
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				1		
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					(continue	ed)

Concentration Process: Solvent Extraction (VII)

Chemical Classification: Phenols (K)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
VII K- 1	4-Chloro- 3-Methylphenol		υ		Extractable w/benzene, alcohol and nitrobenzene		90
VII K- 2	2-Chlorophenol	R	U		Extractable w/Diisopropyl- ether, benzene, butylacetate, and nitrobenzene		90
VII K- 3	m-Cresol p-	L,C	I	291 ppm	91% reduction.	Extraction of evaporator condensate from spent caustic processing using isobutylene (S/W=1.8); spray extractor used.	27
VII K- 4	o-Cresol	L,C	I	307 ppm	90% reduction.	See VIIK-3 for comments.	27
VII K- 5	o-Cresol	L,C	I	890 ppm @ 3.21 gal/ hr		Sequential extraction of wastewater from lube-oil refining using butyl acetate (S/W=0.100)& isobutylene (S/W=0.101); RDC extractor used.	27
VII K- 6	o-Cresol	L,C	I	890 ppm 6 3.21 gal/ hr	99.9% reduction.	Sequential extraction of wastewater from lube-oil refining using butyl acetate (S/W=0.30) & isobutylene (S/W=0.101): RDC extractor used.	27
						(continu	ed)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Phenols (K)

No.	Chemical b	Study	Waste	of Study Influent	Results of Study	Comments	Ref.
		Type C	Type d	Char.			
VII K- 7	2,4-Dichloro- phenol	R	. U		Extractable w/benzene, alcohol and nitrobenzene.		90
VII 8	2,4-Dimethyl- phenol	R	U		Extractable w/benzene and alcohol.		90
VII K- 9	4,6-Dinitro-2- Methylphenol	R	U		Extractable w/benzene and acetone.		90
VII K- 10	2,4-Dinitro- phenol	R	U		Extractable w/benzene and alcohol.		90
VII K- 11	2-Nitrophenol	R	U		Extractable w/benzene and alcohol.		90
VII K- 12	4-Nitrophenol	R	U		Extractable w/benzene and alcohol.		90
VII K- 13	Pentachloro- phenol	R	U		Extractable w/benzene and alcohol and nitrobenzene.		90
VII K- 14	Phenol	R	Ü		Extractable w/diisopropyl- ether, benzene, butylacetate and nitrobenzene.		90
VII K- 15	Phenol	L,C	Ι	67 ppm @ 4.6 gal/ hr	6% reduction.	Extraction of ethylene quench wastewater using isobutylene (S/W=0.101); RDC extractor used.	
						(continue	d)

TABLE C-1 (continued)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Phenols (K)

						INCHOLD (K)		
	No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
	VII K- 16	Phenol	L,C	I	69 ppm @ 4.6 gal/ hr	4% reduction.	Extraction of ethylene quench wastewater using isobutane (S/W=0.097); RDC extractor used.	27
	VII K- 17	Phenol	L,C	1	579 ppm	72% reduction.	See VIIK-3 for comments.	27
	VII K- 18	Phenol	L,C	I	8800 ppm @ 3.21 gal/hr	97% reduction	See VIIK-5 for comments.	27
1	VII K- 19	Phenol	L,C	I	8800 ppm @ 3.21 gal/hr	98% reduction.	See VIIK-7 for comments.	27
	VII K- 20	2,4,6-Trichlo- rophenol	R	Ü		Extractable w/benzene, alcohol and nitrobenzene.		90
	VII K- 21	Xylenols	L,C	Ι	227 ppm	96% reduction.	See VIIK-3 for comments.	27
		·						
							(continu	ed)

Concentration Process: Solvent Extraction (VII) Chemical Classification: Phthalates (L)

No.	Chemical b	<u>Descr</u> Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
VII L- 1	Bis (2-ethyl- hexyl) Phtha- late	R	U ·		Extractable w/ethyl ether & benzene.		90
VII L- 2	Butylbenzyl Phthalate	R	U		Extractable w/ethyl ether & benzene.		90
VII L- 3	Phthalate	R	U		Extractable w/ethyl ether & benzene.		90
VII L- 4	Diethyl Phthalate	R	U		Extractable w/ethyl ether & benzene.		90
VII L- 5	Dimethyl Phthalate	R	U		Extractable w/ethyl ether & benzene.		90
VII L- 6	DiN-Octyl Phthalate	R	Ū		Extractable w/ethyl ether & benzene.		90
							e-ventramon ventramon
			l			(continu	ed)

Concentration Process: Solvent Extraction (VII)
Chemical Classification: Polynuclear Aromatics (M)

	*	Dogg	intion o	£ Ctudu			<u> </u>
No.	Chemical b	Study	iption o	Influent	Results of Study	Comments	Ref.
		Study Type ^C	Type d	Influent Char.			1
 							
VII	Anthracene	R	U		Extractable w/toluene.		90
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TABLE C-1CHEMICAL TREATABILITY

Concentration Process: Activated Carbon (IX) Chemical Classification: Alcohols (A)

No.	Chemical b	<u>Descr</u> Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
1X A 1	Allyl Alcohol	I			21.9% reduction; final conc. of 789 ppm; capacity was 0.024 gm/gm of carbon. Adsorbability found to increase with molecular weight. For compounds of <4 carbons order of decreasing adsorption was: undissociated organic acids, aldehydes, esters, ketones, alcohols (when >4 carbons, alcohols moved ahead of esters), glycols. Aromatics had greatest adsorption. Results of two component isotherm tests could be predicted from single compound tests; however, in four-component tests, only about 60% of predicted adsorption occurred. Continuous columns produced 60-80% of theoretical isotherm capacity.		35
IX A- 2	n-Amyl Alcohol (1-Pentanol)	I	P	1000 ppm	71.8% reduction; 282 ppm final conc., 0.155 gm/gm carbon capacity.	See IXA-1 for additional results.	35
IX A- 3	Butanol	B,L	Р	100 ug/1		Filtrasorb 300 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, methyl (continue)	20

TABLE C-1 (continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Alcohols (A)

	1.	Descr	iption o	of Study			· · · · · · · · · · · · · · · · · · ·
No.	Chemical b	Study		Influent	Results of Study	Comments	Ref.
1		ТуреС	Type d	Char.	•		
		 					
		1				chloride-acetone, and	
'				İ		acetone.	
IX	Butanol	I	P	1000 ppm	53.4% reduction; 466 ppm final	See IXA-1 for additional	35
A-					conc., 0.107 gm/gm carbon	results.	1
4					capacity.		
IX	Butanol	I	Р	1000 ppm	75% reduction	24 hr. contact time;	72
A-				500 ppm	67% reduction	carbon does was 10 times	1
5				100 ppm	78% reduction	chemical conc.	
IX	t-Butanol	I	P	1000 ppm	29.5% reduction; 705 ppm fi-	See IXA-1 for additional	35
A-					nal conc., 0.059 gm/gm carbon	results.	
6					capacity.		<u> </u>
IX	Cyclohexanol	B,L	P	100 /ug/1	Complete removal.	See IXA-3 for additional	20
A-		1				results.	
7							
IX	Decanol	B,L	P	100 /ug/1	Complete removal.	See IXA-3 for additional	20
A-						results.	
8							
IX	Ethanol	I	P	1000 ppm	10% reduction; 901 ppm final	See IXA-1 for additional	35
A-		1			conc., 0.020 gm/gm carbon	results.	1
9					capacity.		
IX	2-Ethyl-	I	P	1000 ppm	[See IXA-1 for additional	35
A-	Butanol				nal conc., 0.170 gm/gm carbon	results.	
10			<u> </u>		capacity.		<u> </u>
IX	2-Ethyl-	I	P	700 ppm	98.5% reduction; 10 ppm final		35
A-	Hexanol		1	1	conc., 0.138 gm/gm carbon	results.	1
11					capacity.		
IX	2-Ethyl-1-	B,L	P	100 /ug/l	Complete removal.	See IXA-3 for additional	. 20
A-	Hexanol					results.	
12							
							l .
			ļ	1		(continu	ed)

Concentration Process: Activated Carbon (IX)
Chemical Classification: Alcohols (A)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX A- 13	m-Heptanol	B,L	P	100 µg/l	Complete removal.	See IXA-3 for addi- tional results.	20
IX A- 14	m-Hexanol	I	P	1000 ppm	95.5% reduction; 45 ppm final conc., 0.155 gm/gm carbon capacity.	See IXA-l for addi- tional results.	35
IX A- 15	Isobutanol	I	P	1000 ppm	41.9% reduction; 581 ppm final conc., 0.084 gm/gm carbon capacity.	See IXA-l for addi- tional results.	35
IX A- 16	Isopropanol	I	P	1000 ppm	12.6% reduction; 874 ppm final conc., 0.025 gm/gm carbon capacity.	See IXA-l for addi- tional results.	35
IX A- 17	Methanol	I	P	1000 ppm	3.6% reduction; 964 ppm final conc., 0.007 gm/gm carbon capacity.	See IXA-l for addi- tional results.	35
IX A- 18	Methanol	I	P	1000 ppm 200 ppm 15 ppm	17% reduction 33% reduction 33% reduction	24 hr. contact time; carbon dose was 10 times chemical conc.	72
IX A- 19	Octanol	B,L	P	100 /ug/l	Complete removal.	See IXA-3 for addi- tional results.	20
IX A- 20	Pentanol	B,L	P	100 дід/1	Complete removal.	See IXA-3 for addi- tional results.	20
IX A- 21	Propanol	B,L	P	100 /ug/1	Complete removal.	See IXA-3 for addi- tional results.	20
IX A- 22	Propanol	I	P	1000 ppm	18.9% reduction; 811 ppm final conc., 0.038 gm/gm carbon capacity.	See IXA-l for addi- tional results.	35
						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

_						-		
1	vo.	Chemical b	Study	Waste	of Study Influent	Results of Study	Comments	Ref.
	ı		Type ^C	Type d	Char.	•		
	IX B- 1	Acetaldehyde	I	P	1000 ppm	11.9% reduction; 881 ppm final conc., 0.022 gm/gm carbon capacity. Adsorbability found to increase with molecular weight. For compounds of <4 carbons order of decreasing adsorption was: undissociated organic acids, aldehydes, esters, ketones,	Carbon dose was 5 g/l Westvaco Nuchar.	35
						alcohols (when >4 carbons, alcohols moved ahead of esters), gylcols. Aromatics had greatest adsorption. Results of two-component isotherm tests could be predicted from single compound tests however, in four-component tests, only about 60% of predicted adsorption occurred. Continuous columns produced 60-80% of theoretical isotherm capacity.		
	IX B- 2	Acetic Acid	I	P	1000 ppm	24% reduction; 760 ppm final conc., 0.048 gm/gm carbon capacity.	See IXB-1 for additional results.	35
	IX B- 3	Acetone	I	P	1000 ppm	21.8% reduction; 782 ppm final conc., 0.043 gm/gm carbon capacity.	See IXB-l for additional results.	35
	IX B- 4	Acetone Cyanohydrin	I	P	1000 ppm 200 ppm 100 ppm	60% reduction 45% reduction 30% reduction	24 hr. contact time; carbon dose was 10 times chemical conc.	1
-		1					Continue	:u)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

a No.	Chemical b			of Study			Ì
No.	Chemical	Study		Influent	Results of Study	Comments	Ref.
		Type ^C	Туре а	Char.			
IX	Acrolein	I	P	1000 ppm	30.6% reduction; 694 ppm	See IXB-1 for	35
В-					final conc., 0.061 gm/gm	additional results.	
5					carbon capacity.		
IX	Acrolein	R	Ū	1000 ppm	30% reduction at 0.5% carbon		90
В-					dose.		
6							
IX	Acrylic Acid	Ι	P	1000 ppm		See IXB-1 for	35
B-					final conc., 0.129 gm/gm	additional results.	
7					carbon capacity.		
IX	Acrylonitrile	·I	P	1000 ppm	51% reduction	24 hr. contact time;	72
B- 8				100 ppm	28% reduction	carbon dose was 10	
1	Amyl Acetate	Ī	P	005	000	times chemical conc.	
IX B-	(primary)	1	P	985 ppm		See IXB-1 for	35
9	(bringry)				final conc., 0.175 gm/gm carbon capacity.	additional results.	l [
IX	Butyl Acetate	I	P	1000 ppm	84.6% reduction; 154 ppm	See IXB-1 for	35
B-	bucyr neceute	1	r	1000 ppm	final conc., 0.169 gm/gm	additional results.	33
10					carbon capacity.	additional lesuits.	
IX	Butyl Acrylate	I	P	1000 ppm		See IXB-1 for	35
В~	· · ·			1	final conc., 0.193 qm/qm	additional results.	
11					carbon capacity.		
IX	Butyraldehyde	I	P	1000 ppm	52.8% reduction; 472 ppm	See IXB-1 for	35
В-					final conc., 0.106 gm/gm	additional results.	
12					carbon capacity.		
IX	Butyric Acid	I	P	1000 ppm	·	See IXB-1 for	35
B-					final conc., 0.119 gm/gm	additional results.	
13					carbon capacity.		
IX	Butyric Acid	B,L	P	100 ug/l	•	Filtrasorb 300 used.	20
B					sorption from carbon by	Solvents included pen-	
14					elutriating with solvent.	tane-acetone, diethyl	
						ether, (continue	d)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

					MIIPINCIOS (D)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX B- 14 cont						methylene chloride- acetone, methyl chlo- ride-acetone, and acetone.	
IX B- 15	Caproic Acid	B,L	P	100 ug/l	90% reduction; 3% desorbed from carbon by elutriating with solvent.	See IXB-14 for additional results	20
	Caproic Acid	I	P	1000 ppm	97% reduction; 30 ppm final conc., 0.194 gm/gm carbon capacity.	See IXB-l for additional results.	35
IX B-		I	P	1000 ppm	45.6% reduction; 544 ppm final conc., 0.092 gm/gm carbon capacity.	See IXB-l for additional results.	35
	Cyclohexanone	I	P	1000 ppm	66.8% reduction; 332 ppm final conc., 0.134 gm/gm carbon capacity.	See IXB-l for additional results.	35
1X B- 19	Decanoic Acid	B,L	P	100 ug/1	Complete reduction; 2% desorbed from carbon by elutriating with solvent.	See IXB-14 for additional results.	20
B	Dicyclo- pentadiene (DCPC)	P,C	I	82 to 1000 ppb	Diisopropyl methylphosphonate (DIMP) and TOC used to measure performance. DCPC found to vaporize.	Contaminated ground- water. See IXB-23 for remarks.	86
IX B- 21	Diethylene Glycol	I	P	1000 ppm	26.2% reduction; 738 ppm final conc., 0.053 gm/gm carbon capacity.	See IXB- 1 for additional results.	35
IX B- 22	Diisobutyl Ketone	I	P	300 ppm	100% reduction; 0.060 gm/gm carbon capacity.	See IXB-1 for additional results.	35
						(continu	led)

TABLE C-1 (continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

a No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
В-	Diisoproply Methyl- phosphonate (DIMP)	P,C	I (Bog Water)	210 to 430 ppb DIMP; TOC about 40 ppm; pH 7.6 to 8.0	Average DIMP removal was 99.75% (<1.9 ppb in effluent)	Test 1- Influent flow 7 gpm; carbon feed rate 1649 µg/l, anionic polymer Herufloc 836.2 at 0.556 gm/l conc. and 1000 cc/min flow added; cationic polymer Catfloc at 4 µg/l conc. and 26.5 cc/min flow added; duration of test 4 weeks; 28,600 gal. throughput.	86
			I (Bog Water)	290 to 470 ppb	Average DIMP removal was 98.77% (6.4 ppb in effluent)	Test 2- Carbon feed 1000 ug/l duration of test 3 weeks; other con- ditions similar to Test 1.	
					DIMP removal averaged 99% at 350 Ag/1 carbon dose and 96.33% at 250 ug/1 carbon dose. Optimum anionic/cationic mixture was found to be anionic-0.13 gm/1 and 120 cc/min, cationic - 1.59 gm/1 & 25 cc/min.	Test 3- Influent flow rate 5 gpm; anionic conc. and flow-0.13 gm/l & 120 cc/min; cationic conc. and flow-1.59 gm/l & 25 cc/min; carbon feed at 350 ug/l & 250 Aug/l for 1 week each.	
					DIMP removal ranged from 92.5 to 97.5% at 175 Aug/l carbon dose and 98.7% at 220 ug/l carbon dose.		(a)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

							<u>`</u>
No.	Chemical b	Descr Study Type ^C	Waste	f Study Influent Char.	Results of Study	Comments	Ref.
IX B- 23 cont			I (Bog Water)	400 ppb	DIMP removal steadily decreased to about 40% at carbon dose of 100 Aug/l. DIMP conc. reduced to 50 ppb, reactivated carbon tested 17000 gal before breakthrough, virgin carbon treated 9600 gal; reactivated carbon capacity-3.8 ug DIMP/gm carbon (0.9 lb carbon/1000 gal); virgin carbon capacity 2.3 aug DIMP/gm carbon (1.41b carbon/1000 gal.)	Filtrasorb 300 carbon was used.	
			I (Ground Water)	2680 ppb	98% removal at carbon dose of 252 ug/l	Hydrodarco C carbon; duration of test- 13100 gal.	
				2400 ppb	94 to 97% removal at carbon dose of 200 µg/l	Hydrodarco C carbon; duration of test - 9000 gal.	
				2564 ppb	Could not achieve steady state performance at carbon dose of 252 ug/l & flow rate of 225 gal/hr.	Aqua Nuchar carbon; duration of test - 15200 gal (2 weeks).	
IX B-	1 -	I	P	1000 ppm	16.5% reduction; 835 ppm fi- nal conc., 0.033 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 25	Dodecane	B,L	P	100 ug/1	<u> </u>	See IXB-14 for additional results.	20
						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
ІХ В- 26	Ethyl Acetate	I	P	1000 ppm	50.5% reduction; 495 ppm fi- nal conc., 0.100 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 27	Ethyl Acrylate	I	P	1015 ppm	77.7% reduction; 226 ppm fi- nal conc., 0.157 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 28	Ethylene Glycol	I	Р	1000 ppm	6.8% reduction; 932 ppm fi- nal conc., 0.014 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 29	Formaldehyde	I	P	1000 ppm	9.2% reduction; 908 ppm fi- nal conc., 0.018 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 30	Formic Acid	I	P	1000 ppm	23.5% reduction; 765 ppm fi- nal conc., 0.047 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 31	Heptanoic Acid	B,L	P	100 ug/1	10% reduction; 1% desorbed from carbon by elutriating with solvent.	See IXB-14 for additional results.	20
IX B- 32	Hexadecane	B,L	P	100 ug/1	Complete reduction; 12% de- sorbed from carbon by elutriating with solvent.	See IXB- 14 for additional results.	20
IX B- 33	Hexylene Glycol	Ι	P	1000 ppm	61.4% reduction; 386 ppm fi- nal conc., 0.122 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 34	Isobutyl Acetate	I	P	1000 ppm	82% reduction; 180 ppm fi- nal conc., 164 gm/gm carbon capacity.	See IXB-1 for additional results.	35
1						(continue	:d)

Concentration Process: Activated Carbon (IX)
Chemical Classification: Aliphatics (B)

					withurtes (D)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX B- 35	Isoprene	I	P	1000 ppm 500 ppm	86% reduction 86% reduction	See IXA-5	72
IX B- 36	Isopropyl Acetate	Ι	P	1000 ppm	68.1% reduction; 319 ppm final conc., 0.137 gm/gm carbon capacity.	See IXB-1 for additional results	35
IX B- 37	Lauric Acid	B,L	P	100 /ug/1	Complete removal; No desorp- tion from carbon by elutria- tion with solvent.	See IXB-14 for additional results.	20
IX B- 38	Methyl Acetate	I	P	1030 ppm	26.2% reduction; 760 ppm final conc., 0.054 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 39	Methyl Butyl Ketone	I	P	988 ppm	80.7% reduction; 191 ppm final conc., 0.159 gm/gm carbon capacity.	See IXB-l for additional results.	35
IX B- 40	Methyl Decanoate	B,L	P	100 /ug/l	Complete removal; 71% de- sorbed from carbon by elutriation with solvent.	See IXB-14 for additional results.	20
IX B- 41	Methyl Dodecanoate	B,L	P	100 /ug/l	Complete removal; 50% de- sorbed from carbon by elutriation with solvent.	See IXB-14 for additional results.	20
IX B- 42	Methyl Ethyl Ketone	I	P	1000 ppm	46.8% reduction; 532 ppm final conc., 0.094 gm/gm carbon capacity.	See IXB- 1 for additional results.	35
IX B- 43	Hexadecanoate	B,L	P	100 /ug/1	Complete removal; 35% de- sorbed from carbon by elutriation with solvent.	See IXB- 14 for additional results.	20
IX B- 44	Methyl Isoamyl Ketone	I	P	986 ppm	85.2% reduction; 146 ppm final conc., 0.169 gm/gm carbon capacity.	See IXB- 1 for additional results.	35
						(continue	ed)

Concentration Process: Activated Carbon (IX)

Chemical Classification: Aliphatics (B)

اء	h	Descr		of Study			
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref.
	•	Type ^C	Type d	Char.	_		
IX	Methyl	B,L	P	100 /ug/1	Complete removal; 40% de-	See IXB-14	20
B-	Octadecanoate	בונט	•	100 /49/ 1	sorbed from carbon by	for additional results.	-
45	octaaccanoacc				elutriation with solvent.	ior addressed about 55.	
IX	Methyl Propyl	ī	P	1000 ppm	69.5% reduction; 305 ppm	See IXB-1	35
В-	Ketone		_		final conc., 0.139 gm/gm	for additional results.	
46					carbon capacity.	:	
IX	Myristic Acid	B,L	P	100 /ug/1	Complete removal; no de-	See IXB-14	20
В-	_				sorption from carbon by	for additional results.	
47					elutriation with solvent.		
IX	Octadecane	B,L	P	100 /ug/l	Complete removal; no desorp-	See IXB- 14	20
В-					tion from carbon by	for additional results.	
48					elutriation w/solvent.		L
IX	Octanoic Acid	B,L	P	100 /ug/1	50% removal; 1% desorbed	See IXB- 14	20
B-					from carbon by elutriation	for additional results.	
49					w/solvent.		
IX	Propional-	L	P	1000 ppm	27.7% reduction; 723 ppm	See IXB- 1	35
B-	dehyde				final conc., 0.057 gm/gm	for additional results.	
50					carbon capacity.		
IX	Propionic Acid	B,L	P	100 µg/l	Complete removal, no desorp-	See IXB- l	20
В-	'				tion from carbon by	for additional results.	
51					elutriation with solvent.		
IX	Propionic Acid	I	P	1000 ppm	32.6% reduction; 674 ppm	See IXB- 1	35
В-					final conc., 0.065 gm/gm	for additional results.	
52					carbon capacity.		
IX	Propyl Acetate	I	P	1000 ppm	75.2% reduction; 248 ppm	See IXB-1	35
В-					final conc., 0.149 gm/gm	for additional results.	
53					carbon capacity.		
IX	Propylene	I	P	1000 ppm	11.6% reduction; 884 ppm	See IXB-l	35
B-	Glycol]			final conc., 0.024 gm/gm	for additional results.] [
54		<u> </u>			carbon capacity.		<u> </u>
						(continue	ed)

Concentration Process: Activated Carbon (IX) Chemical Classification: Aliphatics (B)

					: Allphacies (B)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX B- 55	Propylene Oxide	I	P	1000 ppm	26.1% reduction; 739 ppm final conc., 0.052 gm/gm carbon capacity.	See IXB- l for additional results.	35
IX B- 56	Pyruvic Acid	B,L	P	100 AIG/1	Complete removal; no desorp- tion from carbon using organic solvent.	See IXB-14 for additional results.	20
IX B- 57		B,L	P	100 µg/l	Complete removal; 25% de- sorbed from carbon by elutriation with solvent.	See IXB-14 for additional results.	20
IX B- 58	Tetraethylene Glycol	I	Р	1000 ppm	58.1% reduction; 419 ppm final conc., 0.116 gm/gm carbon capacity.	See IXB-l for additional results.	35
IX B- 59	Triethylene Glycol	I	P	1000 ppm	52.3% reduction; 477 ppm final conc., 0.105 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 60	Valeric Acid	B,L	P	100 /ug/l	Complete removal; 10% de- sorbed from carbon by elutriation with solvent.	See IXB-14 for additional results.	20
IX B- 61	Valeric Acid	Ι	P	1000 ppm	79.7% reduction; 203 ppm final conc., 0.159 gm/gm carbon capacity.	See IXB-1 for additional results.	35
IX B- 62	Vinyl Acetate	I	P	1000 ppm	64.3% reduction; 357 ppm final conc., 0.129 gm/gm carbon capacity.	See IXB-1 for additional results.	35
	·						
				1		(continue	e g)

Concentration Process: Activated Carbon (IX)

Chemical Classification: Amines (C)

	No.	Chemical b	Study Type ^C	Waste	Influent	Results of Study	Comments	Ref.
-	TV		Type C	a				wer.
	TY		**	Туре d	Char.			
	C-1	Allyamine	I	Þ	1000 ppm	31.4% reduction; 686 ppm final conc., 0.063 gm/gm carbon capacity. Adsorbability found to increase with molecular weight. For compounds of <4 carbons order of decreasing adsorption was: undissociated organic acids, aldehydes, esters, ketones, alcohols (when >4 carbons, alcohols moved ahead of esters), glycols. Aromatics had greatest adsorption. Results of two component isotherm tests could be predicted from single compound tests; however, in four component tests only 60% of predicted adsorption occurred. Continuous columns produced 60-80% of theoretical isotherm capacity.	Carbon dose was 5 g/l Westvaco Nuchar.	35
	IX C- 2	Aniline	B,L	P	100 μg/l	cal isotherm capacity. 100% reduction; No desorption from carbon by elutriation with solvents.	Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, methyl chloride-acetone, and acetone.	20
	IX C-		I	P	1000 ppm	74.9% removal; 251 ppm final conc.; 0.15 gm/gm carbon capacity.	See IXC-1 for addition- al results. (continue	35 d)

TABLE C-1(continued)

		Descr	iption o	of Study			,
No.	Chemical ^b	Study Type ^C		Influent	Results of Study	Comments	Ref
IX C- 4	Butylamine	B,L	P	100 /ug/l	100% removal; no desorption from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
C- 5	Butylamine	I	P	1000 ppm	52% reduction; 480 ppm final conc., 0.103 gm/gm carbon capacity.	See IXC-1 for additional results.	35
C- 6	Cyclohexyl- amine	B,L	P	100 Aug/1	100% removal; 38% desorption from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
IX C- 7	Dibutylamine	B,L	P	100 Aug/l	100% removal; No desorption from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
C- 8	Di-N- Butylamine	I	P	1000 ppm	87% removal; 130 ppm final conc., 0.174 gm/gm carbon capacity.	See IXC-1 for additional results.	35
C- 9	Diethanolamine	I	P	996 ppm	27.5% removal; 722 ppm final conc., 0.057 gm/gm carbon capacity.	See IXC-1 for additional results.	35
C- 10	Diethylene- triamine	I	P	1000 ppm	29.4% removal; 706 ppm final conc., 0.062 gm/gm carbon capacity.	See IXC-1 for additional results.	35
C- 11	Dihexylamine	B,L	P	100 µg/l	100% removal; 24% desorption from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
C- 12	Diisopropan- olamine	I	P	1000 ppm	45.7% removal; 543 ppm final conc., 0.091 gm/gm carbon capacity.	See IXC-1 for additional results.	35
IX C- 13	Dimethylamine	B,L	P	100 /ug/1	100% removal; 82% desorption from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
1						(continue	ed)

Concentration Process: Activated Carbon (IX)
Chemical Classification: Amines (C)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IX C- 14	Dimethyl- nitrosamine	Ι	P		Not adsorbed.		31
IX C- 15	Di-N- Propylamine	I	P	1000 ppm	80.2% removal; 198 ppm final conc., 0.174 gm/gm carbon capacity.	See IXC-1 for additional results.	35
IX C- 16	Ethylene- diamine	I	P	1000 ppm	10.7% removal; 893 ppm final conc., 0.021 gm/gm carbon capacity.	See IXC-l for additional results.	35
IX C- 17	N-Ethyl- morpholine	I	P	1000 ppm	47.3% removal; 527 ppm final conc., 0.095 gm/gm carbon capacity.	See IXC-l for additional results.	35
IX C- 18	Hexylamine	B,L	P	100 Aug/1	100% removal; 24% desorbed from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
IX C- 19	2-Methyl-5- Ethylpyridine	I	P	1000 ppm	89.3% removal; 107 ppm final conc., 0.179 gm/gm carbon capacity.	See IXC-l for additional results.	35
IX C- 20	N-Methyl Morpholine	I	P	1000 ppm	42.5% removal; 575 ppm final conc., 0.085 gm/gm carbon capacity.	See IXC-1 for additional results.	35
IX C- 21	Monoethan- olamine	I	Р	1012 ppm	7.2% removal; 939 ppm final conc., 0.015 gm/gm carbon capacity.	See IXC-1 for additional results.	35
IX C- 22	Monoisopro- panolamine	I	P	1000 ppm	20% removal; 800 ppm final conc., 0.04 gm/gm carbon capacity.	See IXC-l for additional results.	35
IX C- 23	Morpholine	B,L	P	100 Aug/l	100% removal; 67% desorbed from carbon by elutriation with solvent.	See IXC-2 for additional results.	20
						(continue	d)

Concentration Process: Activated Carbon (IX)

Chemical Classification: Amines (C)

Type C Type d Char. IX C-24 B-Napthylamine I P I P Isotherm kinetics were as follows: Carbon K 1/n Darco 77.4 0.361 Filtrasorb 166.0 0.234 Carbon dose (mg/l) required to reduce 1 mg/l to 0.1 mg/l: Darco - 27 Filtrasorb - 10 IX Octylamine B,L P 100 Aug/l 100% removal; no desorption from carbon by elutriation with solvent. IX Piperidine B,L P 100 Aug/l 100% removal; 73% desorbed from carbon by elutriation with solvent. IX Pyridine I P 1000 ppm 53.3% removal; 467 ppm final conc., 0.107 gm/gm carbon capacity. IX Pyrrole B,L P 100 Aug/l 100% removal; 16% desorbed for additional results. IX Pyrrole B,L P 100 Aug/l 100% removal; 16% desorbed for additional results. IX Pyrrole B,L P 100 Aug/l 100% removal; 16% desorbed for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results.								
follows: Carbon K 1/n Darco 77.4 0.361 Filtrasorb 166.0 0.234 Carbon dose (mg/l) required to reduce 1 mg/l to 0.1 mg/l: Darco - 27 Filtrasorb - 10 IX C- C- C- 25 Piperidine B,L P 100 ug/l 100% removal; no desorption from carbon by elutriation with solvent. IX Pyridine IX Pyridine IX Pyridine IX Pyridine IX Pyridine IX Pyridine IX C- 26 IX Pyridine IX Pyridine IX Pyridine IX Pyridine IX C- C- 27 IX Tributylamine B,L P 100 ug/l 100% removal; 16% desorbed from carbon by elutriation with solvent. IX Pyrrole B,L P 100 ug/l 100% removal; 16% desorbed from additional results. Carbon Machine I 100 ug/l 100% removal; 16% desorbed from additional results. IX Pyrrole B,L P 100 ug/l 100% removal; 16% desorbed from additional results. C- C- 28 IX Tributylamine B,L P 100 ug/l 100% removal; no desorption from carbon by elutriation with solvent. IX Tributylamine B,L P 100 ug/l 100% removal; no desorption from carbon by elutriation with solvent. IX Triethanol- 29 IX Triethanol- 35 IX Triethanol- 36 IX Triethanol- 37 IX Triethanol- 38 IX Triethanol- 39 IX Triethanol- 30 IX Triethanol- 30 IX Triethanol- 31 I P 1000 ppm 33% removal; 670 ppm final conc., 0.067 gm/gm carbon for additional results.	No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref.
to reduce 1 mg/l to 0.1 mg/l: Darco - 27 Filtrasorb - 10 IX Octylamine C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-	c-	B-Napthylamine	I	Р		follows: Carbon K 1/n Darco 77.4 0.361 Filtrasorb 166.0 0.234		31
from carbon by elutriation with solvent. Piperidine B,L P 100 Aug/l 100% removal; 73% desorbed from carbon by elutriation with solvent. Pyridine I P 1000 ppm 53.3% removal; 467 ppm final conc., 0.107 gm/gm carbon for additional results. Pyrrole B,L P 100 Aug/l 100% removal; 16% desorbed from additional results. Pyrrole B,L P 100 Aug/l 100% removal; 16% desorbed from additional results. Pyrrole B,L P 100 Aug/l 100% removal; 16% desorbed from additional results. Pyrrole B,L P 100 Aug/l 100% removal; 10% desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results. Pyrrole B,L P 100 Aug/l 100% removal; no desorption for additional results.	•					to reduce 1 mg/l to 0.1 mg/l: Darco - 27		
from carbon by elutriation with solvent. IX Pyridine C-27 IX Pyrrole C-28 IX Pyrrole C-28 IX Tributylamine C-29 IX Triethanol-29 IX Triethanol-29 IX Triethanol-30 IX	C-	Octylamine	B,L	P	100 Mg/1	from carbon by elutriation	for additional results.	
C- C- C- C- C- C- C- C- C- C- C- C- C- C	C-	Piperidine	B,L	P	100 /ug/1	from carbon by elutriation		i
from carbon by elutriation with solvent. IX Tributylamine B,L P 100 Aug/l 100% removal; no desorption for additional results. IX Triethanol- I P 1000 ppm 33% removal; 670 ppm final conc., 0.067 gm/gm carbon for additional results.	C-	Pyridine	I	P	1000 ppm	conc., 0.107 gm/gm carbon		1
from carbon by elutriation with solvent. IX Triethanol- I P 1000 ppm 33% removal; 670 ppm final conc., 0.067 gm/gm carbon for additional results.	C-	-	B,L	P	100 /ug/1	100% removal; 16% desorbed from carbon by elutriation		1
C- amine conc., 0.067 gm/gm carbon for additional results.	C-	_	B,L	P	100 дз/1	from carbon by elutriation		•
	C-	amine	I	P	1000 ppm	conc., 0.067 gm/gm carbon		

Concentration Process: Activated Carbon (IX) Chemical Classification: Aromatics (D)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX D- 1	Acetophenone	B,L	P	100 Alg/1	50% reduction; 2% desorbed from carbon by elutriation with solvent.	Filtrasorb 300 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, methey chloride-acetone, and acetone.	20
IX D- 2	Acetophenone	I	P	1000 ppm	97.2% removal; 28 ppm final conc., 0.194 gm/gm carbon capacity. Adsorbability found to increase with molecular weight. For compounds of <4 carbons order of decreasing adsorption was: undissociated organic acids, aldehydes, esters, ketones, alcohols (when > 4 carbons, alcohols moved ahead of esters), glycols. Aromatics had greatest adsorption. Results of two component isotherm tests could be predicted from single compound tests; however, in four-component tests, only about 60% of predicted adsorption occurred. Continuous columns produced 60-80% of theoretical isotherm capacity.	Carbon dose was 5 g/l Westvaco Nuchar.	35
			l			(continue	ed)

TABLE C-1(continued)

r					· MI OMA CIOD (D)		
a	Ъ	Descr	iption o	of Study			
No.	Chemical b	Study	Waste		Results of Study	Comments	Ref.
		Type C	Type d	Influent Char.	,	COMMUNICIPED .	Wer.
							
[[•		
						·	
IX	Benzaldehyde	B,L	P	100 Aug/1	50% reduction; 2% desorbed	See IXD-1	20
D-	•				from carbon by elutriation	for additional results.	
3					with solvent.		
IX	Benzaldehyde	T	P	1000 ppm		See IXD-2	35
D-			_		conc., 0.188 gm/gm carbon	for additional results.	
4					capacity.		
IX	Benzaldehyde	ī	P	1000 ppm	99% removal	24 hr. contact time;	72
D-	•				99% removal	carbon dose was 10	
5					98% removal	times chemical conc.	
IX	Benzene	P,C	H	1 ppb	90% removal (to 0.1 ppb ef-	Spilled material treat-	6
D-		,			fluent conc.) achieved in	ed using EPA's mobile]
6					8.5 min. contact time.	treatment trailer.	
IX	Benzene	I	P	1 ppm	0.7 mg/gm carbon capacity.		21
D-					3, 3		
7							
IX	Benzene	I	P		Isotherm kinetics were as		31
D-	•				follows:		
8					i i		
					$\begin{array}{ccc} Carbon & \underline{K} & \underline{1/n} \\ Darco & \underline{26.8} & \underline{1.305} \end{array}$		
j					Filtrasorb 18.5 1.158		
							1
		•			Carbon dose (mg/l) required		İ
		İ			to reduce 1 mg/1 to 0.1 mg/1:		İ
					Darco - 678		
		<u> </u>			Filtrasorb - 705		L
IX	Benzene	I	P	416 ppm	95% reduction; 21 ppm final	See IXD- 2	35
D-					conc., 0.080 gm/gm carbon	for additional results.)
9		<u> </u>			capacity.		1
, 1		1	ţ	1	((continue	a)

TABLE C-1(continued)

No.	Chemical b	Descr Study Type C	Waste	Influent Char.	Results of Study	Comments	Ref.
IX D-	Benzene	R	I	1500 ppm TOC	Effluent conc. of 30 ppm TOC achieved (98% removal)	At contact time of 55 min.; 0.15 MGD flow; pretreatment included pH adjustment.	38
IX D- 11	Benzene	I	P	250 ppm 50 ppm	95% removal 91% removal 60% removal	24 hr. contact time; carbon dose was 10 times chemical conc.	72
IX D- 12	Benzene	R	ט	416 ppm	95% removal at 0.5% carbon dose.		90
1X D- 13	Benzidine	I	P		Isotherm kinetics were as as follows: Carbon K 1/n Darco 85.4 0.253 Filtrasorb 173 0.288 Carbon dose (mg/l) required to reduce 1 mg/l to 0.1 mg/l: Darco - 19 Filtrasorb - 10		31
IX D- 14	Benzil	B,L	P	100 ug/l	carbon by elutriation with solvent.	for additional results.	20
IX D- 15	Benzoic Acid	B,L	P	100 ug/1	from carbon by elutriation with solvent.	See IXD-l for additional results.	20
1X D- 16	Benzoic Acid	I	P	1000 ppm	91.1% removal; 89 ppm final conc., 0.183 gm/gm carbon capacity.	See IXD-2 for additional results.	35
	-					(continue	d)

Concentration Process: Activated Carbon (IX)

Chemical Classification: Aromatics (D)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IX D- 17	Chlorinated Aromatics	R	I	6000 ppm TOC	Effluent conc. of 3000 ppm TOC achieved (50% reduction). High effluent conc. because activated carbon served as pretreatment before biological system.	At contact time of 1375 min; flow of 6000 gpd; pretreatment included chemical reduction.	38
IX D- 18	Chlorobenzene	I	P	1 mg/1	93 mg/gm carbon capacity.	•	21
IX D- 19	Chlorobenzene	F,C	D		50% reduction.	Treatment of effluent from 0.66 m ³ /sec biological system.	64
IX D- 20	Chlorobenzene	R	Ü	416 ppm	95% removal at 0.5% carbon dose.	*	90
IX D- 21	1-Chloro-2- Nitrobenzene	I	P	1 ppm	103 mg/gm adsorption capacity.		21
IX D- 22	Cumene	B,L	P	100 µg/l	Complete removal; 8% desorbed from carbon by elutriation with solvent.	See IXD-l for additional results.	20
IX D- 23	o-Dichloro- benzene	B,L	P	100 μg/1	Complete removal; 5% desorbed from carbon by elutriation with solvent.	See IXD-1 for additonal results.	20
1X D- 24	o-Dichloro- benzene	R	Ü	416 ppm	95% removal at 0.5% carbon dose.		90
						(continue	ed)

					- IIIOMGCICD (D)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
IX D- 25	m-Dichloro- benzene	B,L	P	100 ug/l	Complete removal; 15% described from carbon by elutriation with solvent.	See IXD-l for additional results.	20
IX D- 26	m-Dichloro- benzene	R	U	416 ppm	95% removal at 0.5% carbon dose.		90
IX D- 27	1,4-Dichloro- benzene	F,C	D		60% removal	Treatment of effluent from 0.66 m ³ /sec biological system.	64
IX D- 28		B,L	P	100 Aug/1	100% removal; 2% desorbed from carbon by elutriation with solvent.	See IXD-1 for additional results.	20
IX D- 29	p-Dichloro- benzene	R	Ū	416 ppm	95% removal at 0.5% carbon dose.		90
IX D- 30	3,3'-Dichloro- benzidine	I	P		Isotherm kinetics were as follows: Carbon K 1/n Darco 126 0.253 Filtrasorb 240 0.194 Carbon dose (mg/l) to reduce 1 mg/l to 0.1 mg/l: Darco - 12.8 Filtrasorb - 5.7		31
IX D- 31	Dimethylaniline (Xylidine)	P,C	Н	380 ppb	94% removal (23 ppb in effluent) achieved in 85 min. contact time.	250,000 gal. spilled materials treated with EPA mobile treatment trailer.	6
:						(continue	d)

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
IX D- 32	2,4-Dinitro- toluene	R	ט	416 ppm	95% removal at 0.5% carbon dose.	Not thermally regenerable.	90
IX D- 33	2,6-Dinitro- toluene	R	Ū	416 ppm	95% removal at 0.5% carbon dose.	Not thermally regenerable.	90
IX D- 34	Ethylbenzene	I	P	1 mg/l	53 mg/gm carbon capacity.		21
IX D- 35	Ethylbenzene	I	L	115 ppm	84.3% reduction; 21 ppm final conc., 0.08 gm/gm carbon capacity.	See IXD-2 for additional results.	35
IX D- 36	Ethylbenzene	F,C	D		50% removal	Treatment of effluent from 0.66 m ³ /sec bio-logical system.	64
IX D- 37.	Ethylbenzene	R	Ü	115 ppm	84.3% removal at 0.5% carbon dose.		90
IX D- 38	Hexachloro- benzene	R	U	416 ppm	95% removal at 0.5% carbon dose.		90 ;
IX D- 39	Hydroquinone	I	P	1000 ppm	83.3% removal; 167 ppm final conc., 0.167 gm/gm carbon capacity.	See IXD-2 for additional results.	
IX D- 40	Isophrone	I	P	1000 ppm	96.6% removal; 34 ppm final conc., 0.193 gm/gm carbon capacity.	See IXD-2 for additional results	
IX D- 41	Isophrone	R	U	1000 ppm	96.6% removal at 0.5% carbon dose.		90
						(continu	ed)

Description of Study ${\tt Chemical}^{\,b}$ No. Waste Influent Study Results of Study Comments Ref. Type d Char. Type C 4,4'-Methylene Ι р Isotherm kinetics were as IX 31 Bis-(2-Chlorofollows: aniline Carbon 1/n 120 0.96 Darco 0.982 Filtrasorb 240 Carbon dose (mg/l) to reduce 1 mg/l to 0.1 mg/l: Darco - 27 Filtrasorb - 15 Nitrobenzene T P 68 mg/gm adsorption capacity 21 IX 1 ppm D-43 95.6% removal; 44 ppm final See IXD-2 IX Nitrobenzene Ι P 1023 ppm 35 conc., 0.196 gm/gm carbon Dfor additional results. capacity. 44 U 416 ppm 95% removal at 0.5% carbon Nitrobenzene R 90 IX Ddose. Paraldehyde 73.9% removal; 261 ppm final P See IXD-2 Ι 1000 ppm 35 conc., 0.148 gm/gm carbon Dfor additional results. capacity. 46 Pyridine I P 1000 ppm 47.3% removal; 527 ppm final See IXD-2 35 conc., 0.095 gm/gm carbon Dfor additional results. 47 capacity. 86% removal; 145 ppm final P 24 hr. contact time; IX Pyridine I 1000 ppm 72 D-500 ppm conc., 86% removal; 71 ppm carbon dose was 10 48 times chemical conc. final conc. 120 mg/gm adsorption Styrene P 1 ppm Ι 21 capacity. D-(continued)

28

TABLE C-1 (continued)

a	b	***************************************		of Study			
No.	Chemical b	Study	Waste ,	Influent	Results of Study	Comments	Ref.
		туре с	Type d	Char.			
IX	Styrene	I	P	180 ppm	88.8% removal; 44 ppm final	See IXD-2 for additional	35
D-					conc., 0.196 gm/gm carbon	results.	
50					capacity.		
IX	Styrene	I	P	200 ppm	97% removal	24 hr contact time;	72
D-				100 ppm	93% removal	carbon dose was 10	
51				20 ppm	55% removal	times chemical conc.	
IX	Styrene Oxide	I	P	1000 ppm	95.3% removal; 47 ppm final	See IXD-2	35
D-					con., 0.19 gm/gm carbon	for additional results.	
52					capacity.		
IX	Toluene	P,C	Н	120 ppb	99.8% removal (0.3 ppb in	250,000 gal spilled	6
D-		1			effluent achieved in 8.5 min	materials treated with	
53		Ì			contact time.	EPA mobile treatment	
1						trailer.	
IX	Toluene	I	P	317 ppm	79.2% removal; 66 ppm final	See IXD-2	35
D-					conc., 0.05 gm/gm carbon	for additional results.	
54					capacity.		<u> </u>
IX	Toluene	R	Ū	317 ppm	79% removal at 0.5% carbon		90
D-		ł			dose.	•	
55		<u> </u>					
IX	Toxaphene	I	I	155 ppb	>99% removal; <1 ppb final		66
D-				рн 7.0	conc., 42 mg/gm carbon		
56		<u> </u>			capacity.		
IX	1,2,4-Tri-	B,L	P	100 μg/l	100% removal; no desorption	See IXD-1	20
D	chlorobenzene				from carbon by elutriation	for additional results.	
57					with solvent.		
IX	1,2,4-Tri-	F,C	D		70% reduction.	Treatment of effluent	64
D-	chlorobenzene					from 0.66 m ³ /sec bio-	1
_58	· · · · · · · · · · · · · · · · · · ·	<u> </u>				logical system.	
IX	1,2,4-Tri-	R	U	416 ppm	95% removal at 0.5% carbon		90
D-	chlorobenzene				dose.		
59				<u> </u>		(continue	ed) —
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TABLEC-1 (continued)

 			***************************************		<u> </u>	
No.	Chemical b	Descr Study Type C		Influent Char.	Results of Study Comments	Ref.
IX D- 60	2,4,6-Trinitro- toluene (TNT)	P,C	I	108 ppm	Carbon adsorption capacity was 0.125 gm/gm at 1 ppm Thermal regeneration not possible because of volume (B.V.) explosion potential.	2
	2,4,6-Trinitro- toluene (TNT) and other muni- tions plant wastewaters: Cyclonite(RDX), Nitramine (Tetry1), and cyclotetrameth- ylene tetrani- tramine (HMX).		I	Not reported	Adsorption capacities TNT is preferentially	40
D- 62	Xylene	P,C	H	140 ppb	> 99.9% removal (0.1 ppb in effluent) achieved in 8.5 min. contact time. 250,000 gal. spilled materials treated with EPA mobile treatment trailer.	6
					(continu	ed)

288

Concentration Process: Activated Carbon (IX) Chemical Classification: Aromatics (D)

					III Oliu Caco (D)		
No.	Chemical b	Descr Study Type ^C	iption o Waste Type d	f Study Influent Char.	Results of Study	Comments	Ref.
IX D- 63		I	P	200 ppm 100 ppm	86% removal 68% removal	24 hr. contact time; carbon dose was 10 times chemical conc.	72
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	,						
						(continu	ed)

289

Concentration Process: Activated Carbon (IX)
Chemical Classification: Ethers (E)

	, , , , , , , , , , , , , , , , , , , 				Ethers (E)		
No.	Chemical b	<u>Descr</u> Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
•	Bis(2-chlorogisopropyl) Ether	- R	υ	Not re- ported	100% removal at 0.5% carbon dose.		90
IX E- 2	Bis(Chlether	R	U	94 ppb	50% removal		90
IX Е- З	Butyl Ether	I	P	197 ppm	100% removal; 0.039gm/gm carbon capacity. Adsorbality found to increase with molecular weight. For compounds of <4 carbons, order of decreasing adsorption was: undissociated organic acids, aldehydes, esters, ketones, alcohols (when>4 carbons, alcohols moved ahead of esters), glycols. Aromatics had greatest adsorption. Results of two-component isotherm tests could be predicted from single compound tests; however, in four-component tests, only about 60% of predicted adsorption occurred. Continuous columns produced 60-80% of theoretical isotherm capacity.		35

Concentration Process: Activated Carbon (IX) Chemical Classification: Ethers (E)

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a No.	Chemical b	Study		of Study Influent	Results of Study	Comments	Ref.
		Type C	Type d	Influent Char.			
IX	Dichloroiso-	I	P	1008 ppm	100% removal; 0.20 qm/qm	See IXE-3	35
E-	propyl Ether				carbon capacity.	for additional results.	
4							
IX	Diethylene	I	P	1000 ppm	82.7% removal; 173 ppm final	See IXE-3	35
E-	Glycol Mono-				conc., 0.166 gm/gm carbon	for additional results.	
5	butyl Ether	!			capacity.		
IX	Diethylene	I	P	1010 ppm	43.6% removal; 570 ppm final	See IXE-3	35
E-	Glycol Mono-				conc., 0.087 gm/gm carbon	for additional results.	
6	ethyl Ether				capacity.		
IX	Ethoxytri-	I	P	1000 ppm	69.7% removal; 303 ppm final	See IXE-3	35
E-	glycol				conc., 0.139 gm/gm carbon	for additional results.]
7				i i	capacity.		ŀ
IX	Ethylene	I	P	1000 ppm	55.9% removal; 441 ppm final	See IXE-3	35
E-	Glycol Mono-				conc., 0.112 gm/gm carbon	for additional results.	•
8	butyl Ether				capacity.		
IX	Ethylene	I	P	1022 ppm	31% removal; 705 ppm final	See IXE-3	35
E-	Glycol Mono-				conc., 0.063 gm/gm carbon	for additional results.	
9	ethyl Ether				capacity.		
IX	Ethylene	I	P	1000 ppm	65.8% removal; 324 ppm final	See IXE-	35
E-	Glycol Mono-				conc., 0.132 gm/gm carbon	for additional results.	
10	ethyl Ether				capacity.		
	Acetate						
IX	Ethylene	I	P	975 ppm	87.1% removal; 126 ppm final	See IXE-3	35
E-	Glycol Mono-				conc., 0.170 gm/gm carbon	for additional results.	1
_11	hexyl Ether				capacity.		
IX	Ethylene	I	P	1024 ppm	13.5% removal; 886 ppm final	See IXE-3	35
E-	Glycol Mono-				conc., 0.028 gm/gm carbon	for additional results.	
12	methyl Ether				capacity.		
IX	Isopropyl	I	P	1023 ppm	80% removal; 203 ppm final	See IXE-3	35
E-	Ether				conc., 0.162 gm/gm carbon	for additional results.	
13					capacity.	(continue	(be
,			1			1 Concinue	1

Concentration Process: Activated Carbon (IX)

Chemical Classification: Halocarbons (F)

No.	Chemical b	Descr Study Type ^C	Waste	f Study Influent Char.	Results of Study	Comments	Ref.
IX F- 1	Bromochloro- methane	Ι	P S S,M	Not re- ported	Sorptive capacity x/m at residual conc (C _f) of 100 ppb was 3.37 mg/g in pure compound studies, 2.56 in a mixture and 0.875 in secondary effluent.		21
IX F- 2	Bromodi- chloro- methane	R	Ū		Reported to be adsorbed		90
IX F- 3	Bromoform	L	W	0.2ppb		See IXF-44 for results.	46
IX F- 4	Bromoform	B,L	P	100 ppb	100% removal; 10% de- sorbed from carbon by elutriation with solvent	Filtrasorb 300 used Solvent included pentane-acetone, diethylether, methylene chloride-acetone, methyl chloride-acetone, and acetone.	20
IX F- 5	Bromomethane	R	Ū		Reported to be adsorbed.		90
IX F- 6	Carbon Tetrachlo- ride	P,C	Н	1.1 ppb	Not detected in effluent after 8.5 min contact time.	250,000 gal spilled materials treated with EPA mobile treatment trailer.	6
F - 7	Carbon Tetrachlo- ride	I	P	Not re- ported	Sorptive capacity (x/m) at residual conc.(C _f) of 100 ppb was 4.66 mg/g	(continue	21 d)

Concentration Process: Activated Carbon (IX) Chemical Classification: Halocarbons (F)

No.	Chemical b	Descr Study Type ^C		f Study Influent Char.	Results of Study	Comments	Ref.
IX F- 8	Carbon Tetrachloride	R	ប		Reported to be adsorbed.		90
IX F- 9	Chloroethane	R	Ü		Reported to be adsorbed.		90
IX F- 10	Chloroethy- lene	R	U		Reported to be adsorbed.		90
IX F- 11	Chloroform	I	P S S,M	Not re- ported	Sorptive capacity (x/m) at residual conc.(C _f) of 100 ppb was 1.58 mg/g in pure compound studies, 0.93 in a mixture, and 0.365 in secondary effluent.		21
IX F- 12	Chloroform	L	W		At 2 ppm chloroform, equilibrium capacity was 12 mg/g.	See IXF-44 for results.	46
IX F- 13	Dibromochlo- romethane	L	W	3.9 ppb		See IXF-44 for results.	46
1X F- 14	Dibromochlo- romethane	I	P S S,M	Not re- ported	Sorptive capacity (x/m) at residual conc. (C_f) of 100 ppb was 7.52 mg/g in pure compound studies, 4.54 in a mixture, and 0.885 in secondary effluent.		21
						(continu	ed)

Concentration Process: Activated Carbon (IX)

Chemical Classification: Halocarbons(F)

4 7					Q		
No.	Chemical b	<u>Descr</u> Study Type ^C	Waste	Influent Char.	Results of Study	Comments	Ref.
IX F- 15	Dibromochlo- romethane	R	υ		Reported to be adsorbed.		90
IX F- 16	Dichloro- ethane	P,C	Н	12 ppb	Not detected in effluent after 8.5 min contact time.	250,000 gal spilled materials treated with EPA mobile treatment trailer.	6
IX F- 17	Dichloro- ethane	I	P S S,M	Not re- ported	Sorptive capacity (x/m) at residual conc.(C _f) of 100 ppb was 1.07 mg/g in pure compound studies, 0.44 in a mixture, and 0.52 in secondary effluent.	Mixture of 6 halo- carbons added to	21
IX F- 18	1,1-Dichloro ethane	L	W	2.3 ppb		See IXF-44 for results.	46
IX F- 19	1,1-Dichloro ethane	R	U		Reported to be adsorbed.		90
IX F- 20	1,2-Dichloroethane	L	W	2.1 ppb		See IXF-44 for results.	46
IX F- 21	1,2-Dichloroethane	R	Ū	1000ppm	81.1% removal at 0.5% carbon dose.		90
IX F- 22	1,1-Dichloroethylene	R	U		Reported to be adsorbed.		90
						(continue	d)

				JIL ICUCION	· Halocarbons (F)		
No.	Chemical b	Descr Study Type ^C	Waste	Influent Char.	Results of Study	Comments	Ref.
IX F- 23	1,2-Dichloro- ethylene	L	W	0.2 ppb		See IXF-44 for results.	46
IX F- 24	1,2-trans- Dichloro- ethylene	R	Ū		Reported to be adsorbed.		90
IX F- 25	Dichloro- fluoromethane	R	Ü		Reported to be adsorbed.		90
IX F- 26	Chlorinated Hydrocarbons	R	บ	4 ppm TOC at 1 MGD	Effluent conc. of 0.05 ppm TOC achievable at contact time of 8 min.	Flow equalization used as pretreatment.	38
IX F- 27	Dichloro- methane	R	Ŭ		Reported to be adsorbed.		90
IX F- 28	1,2-Dichloro- propane	R	ט	1000 ppm	92.9% removal at 0.5% carbon dose.		90
IX F- 29	1,2-Dichloro- propylene	R	Ū		Reported to be adsorbed.		90
1X F- 30	Ethylene Dichloride (EDC)	I	F	1000 ppm	81.1% reduction, 189 ppm final conc., 0.163 g/g carbon capacity. Adsorbability found to increase with molecular weight. For compounds of <4 carbons, order of decreasing adsorption was: undissociated organic acids, aldehydes, esters, ketones, alcohols (when	Carbon dose was 5 g/l Westvaco Nuchar.	35

TABLE C-1 (continued)

Concentration Process: Activated Carbon (IX)
Chemical Classification: Halocarbons(F)

No.	Chemical b	Descr Study Type ^C	- 1	Influent	Results of Study	Comments	Ref.
IX F- 30 cont	Ethylene Dichloride	I	I	waters contain- ing num- erous halocar- bons with predomi- nately EDC at up to	>4 carbons, alcohols moved ahead of esters), glycols. Aromatics had greatest adsorption. Re- sults of two-component isotherm tests could be predicted from single compound tests; however, in four-component tests, only about 60% of pre- dicted adsorption oc- curred. Continuous col- umns produced 60-80% of theoretical isotherm capacity. Carbon adsorption capaci- ty to achieve 10 ppm EDC residual ranged from 0.47 to 1.25 gm EDC/gm carbon Capacity to achieve 0.1 ppm EDC residual ranged from 0.0145 to 0.13 gm EDC/gm carbon. To obtain 0.5 ppm TOC residual, capacity ranged from 0.052 to 0.7 gm TOC/gm carbon. Capacity to achieve 50 ppm TOC resid- ual ranged from 7.0 to 150 gm TOC/gm carbon.	400), Westvaco(WVG) WITCO, and Barneby- Cheney (BCNB-9377) carbons were used. Capacity was depend- ent on wastewater being tested and the carbon.	
	†	+	 	FF.		(Concinue	-4,

TABLE C-1(continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Halocarbons(F)

No.	Chemical b	Descr Study Type ^C	Waste	f Study Influent Char.	Results of Study	Comments	Ref.
1X F- 32		L,C (3 col ums in series 20 mm ID by 450mm length	I	ing numerous halo-carbons with predom-inately EDC. TC 1200ppm EDC-6400 to 6800ppm pH->11 total chlorinated hydro-carbons -8000ppm		carbon was regenerated with 1 atm of steam for 5 min; after 5 regenerations carbon capacity was 0.186 gm EDC/gm carbon or 93% of fresh carbon.	
IX F-	Hexachloro- butadiene	B,L	P	100 ppb	100% removal; 31% de- sorbed from carbon by elutriation with solvent	See IXF-4 for additional comments.	20
1X F- 34	Hexachloro- ethane	B,L	Þ	100 ppb	100% removal; 98% de- sorbed from carbon by elutriation with solvent	See IXF-4 for additional	20 ed)

Concentration Process: Activated Carbon Chemical Classification: Halocarbons (F)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX F- 35	Hexachloro- ethane	R	Ū		Reported to be adsorbed.	,	90
IX F- 36	Methylene Chloride	P,C	Н	190 ppb	73% removal with 51 ppb de- tected in effluent after 8.5 min contact time.	250,000 gal spilled materials treated with EPA mobile treatment trailer.	6
IX F- 37	Propylene Dichloride	I	L	1000 ppm	92.9% reduction, 71 ppm fi- nal conc., 0.183 g/g carbon capacity.	See IXF-32 for additional results.	35
IX F- 38	Tetrachloro- ethane	B,L	P	100 ppb	100% removal; 70% desorbed from carbon by elutriation with solvent.	See IXF-4 for additional comments	20
IX F- 39	1,1,2,2-Tetra- chloroethane	R	Ü		Reported to be adsorbed.		90
IX F- 40	Tetrachloro- ethylene	L	W	179 ppb		See IXF-44 for results.	46
IX F- 41	Tetrachloro- ethylene	R	Ü		Reported to be adsorbed.		90
IX F- 42	Tribromo- methane	R	U		Reported to be adsorbed.		90
						(continue	ed)

TABLE C-1(continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Halocarbons(F)

a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX F- 43	Tribromo- methane	I	P S S,M	Not re- ported	Sorptive capacity(x/m) at residual conc. (C _f) of 100 ppb was 28.7 mg/g in pure compound studies, 10.8 in a mixture, and 1.53 in secondary effluent.		21
IX F- 44	1,1,1-Tri- chloroethane	L	W	551 ppb	Performance for treatment of water containing several halogens. Virgin Regenerated BV to 33ppb 5100 4000 com- pound leak- age Days 13.3 10.4 Gal 38,250 30.000 treat- ed/cu ft sor- bent	height 4" (15 cu cm adsorbent) Flow-2 gpm/cu ft (16 BV/hr) Regenerated at 37 lb steam/cu ft @5 psig	
IX F- 45		R	Ŭ		Reported to be adsorbed.		90
IX F- 46	1,1,2-Tri chloroethane	R	Ū		Reported to be adsorbed.	(continu	90 ed)

Concentration Process: Activated Carbon (IX) Chemical Classification: Halocarbons(F)

,					. narocarpons(r)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX F- 47	<u>1</u>	P,C	н	21 ppb	98.6% removal with 0.3 ppb detected in effluent after 8.5 min contact time.	250,000 gal spilled materials treated with EPA mobile treatment trailer.	6
IX F- 48	-	R	Ŭ		Reported to be adsorbed.		90
IX F- 49		R	บ		Reported to be adsorbed.		90
IX F- 50		B,L	P	100 ppb	100% reduction; 35% de- sorbed from carbon by elutriation with solvent	See IXF-4 for additional comments.	20
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						(continue	d)

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Concentration Process: Activated Carbon (IX)

				SILICACION	Metals (G)		
No.	Chemical b	Descr Study Type ^C		f Study Influent Char.	Results of Study	Comments	Ref.
IX G- 1	Arsenic	F,C	м		No reduction. Increase to 2.4 ppb.	Carbon used as advanced treatment of biologically & chemically treated waste water. Plant capacity 0.66 cu m/sec. Data presented for two time periods.	
IX G- 2	Barium	F,C	М	32 ppb 31 ppb	No reduction. No reduction.	See IXG-1 for comments.	64
IX G- 3	Cadmium	F,C	М		12% reduction; 2.2 ppb effluent conc. 6% reduction; 1.7 ppb effluent conc.	See IXG-1 for comments.	64
IX G- 4	Cadmium	P,C	R	0.029-p.ლ	With virgin Filtrasorb 200 average removal was 19%; w/exhausted FS 200 average removal was 37%.	Study consisted of 8 tests of about 100 hr duration each.	82
IX G-	Chromium	F,C	М	84.0ppb	43% reduction; 48.0 ppb effluent conc.	See IXG-1 for comments.	64
IX G-	Chromium	F,C	М	41.0ppb	37% reduction; 26.0 ppb effluent conc.	See IXG-1 for comments.	64
IX G- 7	Chromium ⁺³	L,I	P	100 ppm	Carbon dose % Removal (ppm) 0 0 0 500 5 1,000 7.5	Test chemical used was Cr Cl ₃ with 24 hr carbon contact time.	72 ed)

Concentration Process: Activated Carbon (IX)

a No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of	Study	Comments	Ref.
IX G- 7					Carbon dose (ppm) 5,000	% Removal		
IX G- 8	Chromium ⁺⁶	L,I	P	100 ppm	10,000 Carbon dose (ppm) 0 500 1,000 5,000	47.5 % Removal 0 16 26 34	24 hr contact time, test chemical was K ₂ Cr ₂ O ₇	72
I X G - 9	Copper	F,C	М	-,	10,000 69% reduction; effluent conc.	36 27 ppb	See IXG-1 for comments.	64
IX G- 10	Copper	F,C	М	49 ppb	35% reduction; effluent conc.	32 ppb	See IXG-1 for comments.	64
IX G- 11	Copper	L,I	P	100 ppm	Carbon Dose (ppm) 0 500 1,000 5,000 10,000	% Removal 0 8 10 73 96.4	24 hr contact time, test chemical was Cu SO ₄	72
I X G- 12	Iron	F,C	М	207 ppb	68% reduction; effluent conc.		See IXG-1 for comments.	64
IX G- 13	Iron	F,C	М	40 ppb	Conc increased in effluent.	to 45 ppb	See IXG-1 for comments. (continu	64 ed)

Concentration Process: Activated Carbon (IX)

a No.	Chemical b	Descr Study		of Study	Results of Study	Comments	Ref.
NO.	Chemical	Type C	Type d	Influent Char.	Results of Study	Comments	Rel.
IX	Lead	F,C	М	22 ppb	Conc. increased to 26 ppb	See IXG-1 for comments.	64
G- 14							
IX G-	Lead	F,C	М	4.7 ppb	Conc. increased to 5.3 ppb.	See IXG-1 for comments.	64
15							7.0
IX G-	Lead	L,I	P	100 ppm	Carbon dose % Removal	24 hr contact time, test chemical used	72
16					0 0	Pb(NO ₃) ₂	
					500 - 13 1,000 17.7		
					5,000 84.0 10,000 93.0		
IX	Manganese	F,C	М	6.2 ppb	21% reduction; 4.9 ppb	See IXG-1	64
G- 17	-				effluent conc.	for comments.	
IX	Manganese	F,C	M	2.3 ppb	Conc. increased to 4.1	See IXG-1	64
G- 18					ppb.	for comments.	
IX	Manganese	L,I	P	100 ppb	Carbon dose % Removal	24 hr contact time, test chemical used	72
G-	•				(ppm) 0 0	was MnCl ₂ .	
					500 1 1,000 3		
					5,000 25		
IX	Mercury	F,C	M	3 6 nnh	10,000 50 Conc. increased to 6.7	See IXG-1	64
G-	Mercury	1,0	1.1	J.o ppb	ppb.	for comments.	
20			 				
						(continu	l ed)

Concentration Process: Activated Carbon (IX)

No.	Chemical b	Descr Study Type C		Influent Char.	Results of Study	Comments	Ref.
I X G- 21	Mercury	F,C	M	1.2 ppb	Conc. increased to 4.9 ppb.	See IXG-1 for comments.	64
IX G- 22	Mercury	L,I	P	100 ppm	Carbon Dose % Remova (ppm) 0 0 500 99 1,000 99 5,000 99 10,000 99	1 24 hr contact time, test chemical used was Hg Cl ₂ .	72
IX G- 23	Mercury	U	U	10 ppb	80% reduction achieved with carbon dose of 100 Mg/l. PAC + chelating agent.	Efficiency of reduction was dependent on pH. Optimum pH was 7.0. Tannic Acid and Citric Acid were ineffective as chelating agents.	
I X G - 24	Mercury	R	U		80% reduction by PAC & Alum Coagulation.	GAC reduction of Hg enhanced by use of chelating agent.	90
1X G- 25	Nickel Selenium	L,I	P	100 ppm	Carbon dose % Remova (ppm) 0 0 0 500 4 1,000 5 5,000 10.5 10,000 52 GAC treatment after Lim	1 24 hr contact time, test chemical used was NiCl ₂ .	72
G- 26		K	U	200 bbm	ppt. yielded 96% reduction.	•	

Concentration Process: Activated Carbon (IX)

No.	Chemical b	Descr Study Type ^C	iption o Waste Type d	of Study Influent Char.	Results of Study	Comments	Ref.
IX G- 27	Thallium	R	ט		GAC treatment after Lime ppt. yielded 84% reduction.		90
IX G- 28	Zinc	F,C	М	670 ppb	81% reduction; 124 ppb effluent conc.	See IXG-1 for comments.	64
IX G- 29	Zinc	F,C	М	412 ppb	61% reduction; 162 ppb effluent conc.	See IXG-1 for comments.	64
						·	
			İ			(continu	ed)

Concentration Process: Activated Carbon (IX)

Chemical Classification: Polychlorinated Biphenyls (I)

						·	
No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
	PCB's (Unspecified)	C,P	Н	19 ppb	Not detectable in efflu- ent with 60 min contact	Treatment by EPA trailer.	6
	PCB's (Unspecified)	C,P	н	1 '	time. Not detectable in effluent with 30-40 min contact time.	See IXI-1 for comments.	6
	PCB's (Unspecified)	C,P	н		Not detectable in efflu- ent with 8.5 min contact		6
IX	Arochlor 1242	L,B,I	P	45 ppb	<0.5 ppb final conc. carbon capacity was 25 mg/g.	Pulverized FS-300	8
	Arochlor 1242	I	P	45 ppb	4.3 mg/g capacity for a 1.1 ppb final conc.		22
	Arochlor 1242	I	S	45 ppb	25 mg/g capacity for a <0.5 ppb final conc.	·	38
1	Arochlor 1242	I	I	45 ppb	25 mg/g capacity for a <0.5 ppb final conc.		66
	Arochlor 1254	L,B,I	P	49 ppb	72 mg/g of carbon capac- ity for a final conc. of <0.5 ppb		8
ΙX	Arochlor 1254	I	P	160 ppb	15.75 mg/g capacity for 98.5% reduction.		22
	Arochlor 1254	I	P	11.15ppb and 37.5 ppb	0.37 mg/g capacity for 99% reduction.		22
						(continu	ed)

306

Concentration Process: Activated Carbon (IX)

Chemical Classification: Polychlorinated Biphenyls (I)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
	Arochlor 1254	C,L	P	1	<0.05 ppb final conc. for 240 BV.	Experiment lasted 5 days.	22
IX	Arochlor 1254	F,C	P	50 ppb	<1.0 ppb final effluent at 0.006 lb/lb capacity.	Cost estimate for full scale columns are \$0.65/100 gal at 0.25 Mgd.	22
	Arochlor 1254	I	P	49 ppb	1.0 mg/g capacity for 1.2 ppb effluent.		22
IX	Arochlor 1254	I	S	49 ppb pH=7.0	7.2 mg/g capacity for final conc. of 0.5 ppb.		38
IX	Arochlor 1254	I	I	49 ppb	See IXI-13 results		66
IX	Arochlor 1254	B,L	Р	100 ppb	94.4% average reduction; 14% desorbed from carbon by elutriation w/solvent		20
						(continue	d)

TABLE C-1 (continued)

No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX J- 1	Aldrin	B,L	P	100 ppb	100% reduction; 2% desorbed by elutriation with solvent.	Calgon FS-300 used.	20
IX J- 2	Aldrin	I	S	48 ppb	30 mg/g of carbon capacity for a final conc. of <1.0 ppb.	рH = 7.0	38
IX J- 3	Aldrin	L,B,I	P	48 ppb	30 mg/g of carbon capacity for a final conc. of <1.0 ppb.	Pulverized FS-300	8
IX J- 4	Aldrin	C,P	Н	8.5 ppb @ 0.1 MG treated	98% reduction w/17 min contact time.	Treated by EPA mobile trailer.	6
IX J- 5	Aldrin	C,P	Н	60.5 ppb @ 3000 gal treated.	99.8% reduction w/240 min contact time.	See IXJ-4 for comments.	6
IX J- 6	2,4-D butyl ester	L,B	P	100 ppb	100% reduction; 10% desorbed from carbon by elutriation w/solvent.	Calgon FS-300 used.	20
IX J- 7	Chlordane	C,P	Н	13 ppb @ 1.0 MG treated	97.3 reduction; w/17 min contact time.	See IXJ-4 for comments.	6
IX J- 8	Chlordane	C,P	Н	1430 ppb @ 3000 gal treated	99.99% reduction; w/240 min contact time.	See IXJ-4 for comments.	6
J- 9	DDD	I	S	56 ppb pH = 7.0	130 mg/g carbon capacity for a final conc. of 0.1 ppb.		38
IX J- 10	DDD	I	P	56 ppb	See IXJ-9 results.	Pulverized FS-300 used. (continue	•

Concentration Process: Activated Carbon (IX) Chemical Classification: Pesticides (J)

					· Pesticides (b)		
No.	Chamian, b			of Study			
No.	Chemical b	Study Type ^C	Waste Type d	Influent Char.	Results of Study	Comments	Ref.
IX J-	DDD	I	I	56 ppb pH=7.0	See IXJ-9 results.		66
11							
1	DDE	I	I	38 ppb	9.4 mg/g carbon capacity		66
J-				ph=7.0	for a final conc. of		
12 IX	DDE	I	P	38 ppb	<1.0 ppb. See IXJ-12 results.	Pulverized FS-300	8
J-	DDE	-	r	be bbn	see ixu-iz resuits.	used.	В
13				,			
ΙX	DDE	I	S	38 ppb	See IXJ-12 results.		38
J-				pH=7.0			
14							
	DDT	I	S	41 ppb	11 mg/g of carbon capac-		38
J- 15				рн=7	ity for a final conc. of 0.1 ppb		
	DDT	L,B,I	P	41 ppb	11 mg/g of carbon capac-	Pulverized FS-300	8
J-			-		ity for a final conc. of		
16					0.15 ppb.		
1	DDT	C,L,R	P,R	10 ppb	Greater than 99% reduc-	Cumulative removal	6
J-	,				tion achieved.	following prechlo-]
17						rination and coagu-	•
ΙX	DDT	B,L	P	100 ppb	100% reduction; 51% de-	lation-filtration Calgon FS-300	20
J-		"	•	L C C PPD	so rbed from carbon by	Cargon ra-300	[]
18					elutriation w/solvent.		
IX	DDT	I	I	41 ppb	See IXJ-15 results.		66
J-				pH=7			
19							
IX J-	Dieldrin	I	S	19 ppb	15 mg/g carbon capacity		38
20					for a final conc. of 0.05 ppb.	/	1
	L		ļ	 	0.00 ppn.	(continue	a)

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no.	b			of Study			T
No.	Chemical b	Study	Waste	Influent	Results of Study	Comments	Ref
		Type ^C	Type C	Char.	-		
IX.	Dieldrin	L,B,I	P	19 ppb	15 mg/g carbon capacity	Pulverized FS-300	8
J -					for a final conc. of		
21					0.08 ppb.		
IX	Dieldrin	C,P	Н	11 ppb	No detectable level in	Treated by EPA	6
J-				@ 0.1MG	effluent w/17 min con-	mobile trailer.	
22				treated	tact time.		
IX	Dieldrin	C,P	Н	60.5ppb	No detectable level in	See IXJ-22	6
J –				@ 3000	effluent w/240 min con-	for comments.	
23				gal	tact time.		
				treated.			
IX	Dieldrin	B,L,R	P,R	10 ppb	Carbon Conc. % Removal	Cumulative removal	6
J-					5 mg/1 75	following prechlo-	
24	,				10 mg/1 85	rination & coagula-	1
					20 mg/l 92	tion-sedimentation.	
IX	Dieldrin	C,L,R	P,R	10 ppb	Greater than 99% reduc-	See IXJ- 24	6
J-				@ 0.5	tion achieved.	for comments.	
25				gpm/ft ³			
IX	Dieldrin	I	I	19 ppb	See IXJ-10 results.		66
J-		1 1		pH=7.0			
26							
IX	Endrin	I	I	62 ppb	100 mg/g carbon capacity		66
J-]]		pH=7.0	for a final conc. of		
_27					0.05 ppb.		
IX	Endrin	L,B,I	P	62 ppb	100 mg/g carbon capacity	Pulverized FS-300	8
J-		1 1			for a final conc. of		1
28	L				0.07 ppb		
IX J-	Endrin	B,L,R	P,R	10 ppb	Carbon Conc. % Removal	See IXJ-24	6
-					5 mg/l 80	for comments.	
29					10 mg/1 90		
					20 mg/1 94		<u></u>
	•	1 1		1		(continue	3.

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N	a o.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
	1-	Endrin	C,L,R	P,R	10 ppb @ 0.5 gpm/ft ³	Greater than 99% reduction achieved.	See IXJ-24 for comments.	6
1 -	г- 31	Endrin	I	S	62 ppb pH = 7.0	See IXJ-27 results.		38
- 1	Γ- 12	Heptachlor	C,P	Н	6.1 ppb @ 0.1 MG treated	99% reduction w/17 min contact time.	Treated by EPA mobile trailer.	6
	(- 3	Heptachlor	C,P	Н	80 ppb @ 3000 gal treated	99.9% reduction w/240 min contact time.	Treated by EPA mobile trailer.	6
3	4	Herbicides (unspecified)	R	Ū	10,000 ppm TOC @ 0.02 MGD	99% TOC reduction achieved w/412 min contact time.	Pretreatment included pH adjustment.	38
	f	Herbicides (unspecified)	R	Ü	1500 ppm TOC @ 0.02 MGD	90% TOC reduction achieved w/412 min contact time.	Pretreatment included settling and filtration	38
3	6	Kepone	C,P	H	4000 ppb @ 0.225MG treated	No detectable levels in effluent w/45 min contact time.	Treated by EPA mobile trailer.	6
1 -	7	Lindane	B,L,R	P,R	10 ppb	Carbon Conc. % Removal 5 mg/l 30 10 mg/l 55 20 mg/l 80	See IXJ-24 for comments.	6
							(continue	đ)

Chemical Classification: Pesticides (J)

Concentration Process: Activated Carbon (IX)

a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX J- 38	Lindane	C,L,R	P,R	10 ppb @ 0.5 gpm/ft ³	Greater than 99% reduction achieved.	See IXJ-24 for comments.	6
IX J- 39	Parathion	B,L,R	P,R	10 ppb	Carbon Conc. % Removal 5 mg/l >99 10 mg/l >99 20 mg/l >99	See IXJ-24 for comments.	6
IX J- 40	Parathion	C,L,R	P,R	10 ppb	Greater than 99% reduction achieved.	See IXJ-24 for comments.	6
IX J- 41	2,4,5-T ester	B,L,R	P,R	10 ppb	Carbon Conc. % Removal 5 mg/l 80 10 mg/l 90 20 mg/l 95	See IXJ-24 for comments.	6
IX J- 42	2,4,5-T ester	C,L,R	P,R	10 ppb @ 0.5 gpm/ft ³	Greater than 99% reduction achieved.	See IXJ-24 for comments.	6
IX J- 43	Toxaphene	C,P	P	36 ppb @ 0.25 MG treated	97% reduction w/26 min contact time.	Treated by EPA mobile trailer.	6
IX J- 44	Toxaphene	L,B,I	P	155 ppb	42 mg/g carbon capacity for a final conc. of <1.0 ppb.	Pulverized FS-300.	8
IX J- 45	Toxaphene	I	S	155 ppb	See IXJ-44 results.		38
						(continu	ed)

Concentration Process:

Activated Carbon (IX)

Chemical Classification:

Phenols(K)

						· PHEHOTS(K)		
No.	Chemical b	Descr Study Type ^C	iption o Waste Type d	Influ	ent	Results of Study	Comments	Ref
IX K- 1	Butyl Phenol	C,P	Н	300	ppb	95% reduction w/8.5 min contact time.	250,000 gal spill treated by EPA mo- bile treatement trailer.	6
IX K- 2	4-Chloro- 3-Methyl- phenol	B,L	P	100	ppb	100% reduction; 10% de- sorbed from carbon by elutriation w/solvent.	Calgon FS-300 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, chloroformacetone and acetone.	20
IX K- 3	Cresol	C,P	Н	230	ppb	96.5% reduction w/8.5 min contact time.	250,000 gal spill treated by EPA mobile treatment trailer.	6
I X K- 4	2,3-Dichloro phenol	B,L	P	100	ppb	100% reduction; 14% de- sorbed from carbon by elutriation w/solvent.	See IXK-2 for comments.	20
IX K- 5	Dimethyl- phenol	C,P	Н	1220	ppb	99.6% reduction w/8.5 min contact time.	See IXK-3 for comments.	6
IX K- 6	3,5-Dimethyl phenol	B,L	P	100	ppb	100% reduction; 5% de- sorbed from carbon by elutriation w/solvent.	See IXK-2 for comments.	20
IX K- 7	2,4-Dinitro- phenol	I	P			For pH=3.0: Carbon capacity=405mg/g		21
			1				(continue	ed)

Concentration Process: Activated Carbon (IX)
Chemical Classification: Phenols(K)

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No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
IX K- 7 cont					For pH=7.0: Carbon capacity=160mg/g		
					1/n =0.25 r =0.87		
8 K-	Nonylphenol .	Ī	P		For pH=3.0: Carbon capacity=570mg/g		21
IX K- 9	Pentachloro- phenol	I	P		r =0.98 For pH=3.0: Carbon capacity=635mg/g K =260 1/n =0.4 r =0.98	(continue	21 21

Concentration Process: Activated Carbon (IX)

Chemical Classification: Phenols(K)

					111011024 (117		•
No.	Chemical b	Descr Study Type ^C	Waste	Influent Char.	Results of Study	Comments	Ref.
IX K- 9 cont					For pH=7.0: Carbon capacity=385mg/g K =145 1/n =0.42 r =0.98		
					For pH=9.0: Carbon capacity=260mg/g K =100 1/n =0.41 r =0.98		
I X K- 10	Pentachloro- phenol	C,P	Н	10 ppm	Not detectable in efflu- ent after 26 min contact time.		
IX K- 11	Phenol	B,L	P	100 ppb	100% reduction; 6% de- sorbed from carbon by elutriation w/solvent.	See IXK-2 for comments.	20
ІХ К- 12	Phenol	I	P		For pH=3.0: Carbon capacity=85 mg/g		21
					For pH=7.0: Carbon capacity=80 mg/g		
					Carbon capacity=70 mg/g K =22	(continu	ed)

Concentration Process:

Activated CArbon (IX)
Phenols(K)

Chemical Classification:

	No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
	IX K- 12					1/n =0.49 r =0.94		
	cont IX K- 13	Phenol	I	p	1.0 ppm	Adsorption capacity 21 mg/g		21
	IX K- 14	Phenol	C,P	Н	140 ppb	100% reduction w/8.5 min contact time.	See IXK-3 for comments.	6
)	IX K- 15	Phenol	L,I	P	500 ppm	99% reduction 99% reduction 99% reduction	24 hr contact time time w/carbon dose of 10x phenol conc.	72
•	IX K- 16	Phenol	I	S		80% reduction; 194 ppm final conc., 161 mg/g carbon capacity.		35
	IX K- 17	Phenol	R	Ū	200 ppm @ 0.05 MGD	effluent conc. of 0.01 ppm achievable at contact time of 165 min.	Settling, equaliza- tion & mixed media filtration used as pretreatment.	38
	IX K- 18	Phenol	R	U		Effluent conc. of 100ppm achievable at contact time of 41 min.		38
	IX K- 19	Phenol	R	Ū		Effluent con. of 0.05ppm achievable at contact time of 24 min.	Biological & mixed media filtration used as pretreatment	38
	IX K- 20	Phenol	R	U	1200 ppm @0.15MGD	Effluent conc. of 1.0ppm achievable at contact time of 55 min.	Sand filtration & settling used as pretreatment.	38
							(continue	d)

Concentration Process: Activated Carbon (IX)

Chemical Classification:

Phenols (K)

			ar oraci	STLICACION	: Phenois (K)		
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
IX K- 21	Phenol	R	ט	80 ppm @ 0.3MGD	Effluent conc. of 1.0ppm achievable at contact time of 33 min.	Biological, set- tling & multi media filtration used as pretreatment.	38
IX K- 22	Phenol	R	Ü	1000 ppm	80.6% reduction achieved		90
IX K- 23	Phenol	B,L	P	100 ppb	100% reduction; 6% de- sorbed from carbon by elutriation w/solvent.	See IXK-2 for comments.	20
I X K- 24	Resorcinol	B,L	Þ	100 ppb	100% reduction; 0% de- sorbed from carbon by elutriation w/solvent.	See IXK-2 for comments.	20
IX K- 25	2,4,6-Tri- chlorophenol	B,L	Þ	100 ppb	sorbed from carbon by elutriation w/solvent.	See IXK-2 for comments.	20
IX K- 26	Trimethyl- phenol	C,P	Н	130 ppb	92% reduction w/8.5 min contact time.	See IXK-3 for comments.	6
	,						
						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Activated Carbon (IX) Chemical Classification: Phthalates (L)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref
IX L- 1	Bis(2-ethyl- hexyl)Phthalate	В	Ι	1300 ppb @ 1.0gpm/ft2	Final conc. of <22 ppb achiev able at 90 min EBCT.	TOC conc. of influent was 15000 ppm; estimated cost excluding pretreatment was \$27.00/1000 gal	
IX L- 2	Bis(2-Ethyl- hexyl)Phthalate		Ü		Reduction by flocculation w/Al ₂ (SO ₄) improved w/granu-lar activated carbon pretreatment.		90
IX L- 3	Dibutyl Phthalate	B,L	Þ	100 ppb	100% reduction; 38% desorbed from carbon by elutriation w/solvent.	Calgon FS-300 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, chloroform-acetone and acetone.	20
IX L- 4	Dimethyl Phthalate	B,L	P	100 ppb	100% reduction; 13% desorbed from carbon by elutriation w/solvent.	See IXL-3 for comments.	20
						(continue	d)

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Concentration Process: Activated Carbon (IX)

Chemical Classification: Polynuclear Aromatics (M)

		CHEMIL	ar Clas.	silication	: Polynuclear Aromatics (M)		
No.	Chemical b	Descr Study Type ^C	iption d Waste Type	of Study Influent Char.	Results of Study	Comments	Ref
IX M- 1	Biphenyl	B,L	Р	100 ppb	100% reduction; 2% desorbed from carbon by elutriation w/solvent.	Calgon FS-300 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, chloroform-acetone and acetone.	
IX M- 2	Cumene	B,L	P	100 ppb	100% reduction; 8% desorbed from carbon by elutriation w/solvent.	See IXM-1 for comments.	20
IX M- 3	Dimethyl- Naphthalene	B,L	P .	100 ppb	80% reduction; ll% desorbed from carbon by elutriation w/solvent.	See IXM-1 for comments.	20
	1,1-Diphenyl- hydrazine	I	P	рн=7.5	Isotherm kinetics were as follows: Carbon K 1/n Darco 94.8 0.279 Filtrasorb 149.0 0.232 Carbon dose (mg/l) required to reduce 1.0 mg/l to 0.lmg/l: Darco - 18.0 Filtrasorb - 10.0	,	31
IX M- 5	Fluoranthrene	B,L	P	100 ppb	80% reduction; 5% desorbed from carbon by elutriation w/solvent.	See IXM-1 for comments.	20
IX M- 6	Napthalene	I	P		Isotherm kinetics were as follows: Carbon K 1/n Darco 62.8 0.30 Filtrasorb 1.69 0.56		31
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Concentration Process: Activated Carbon (IX)

Chemical Classification: Polynuclear Aromatics (M)

					· Polynuclear Aromatics (M)		
No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
IX M- 6 cont					Carbon dose (mg/l) required to reduce 1.0mg/l to 0.1mg/l: Darco - 29.0 Filtrasorb - 19.0		
IX M- 7	Napthalene	F,C	М	Conc. not re- ported	70% reduction achieved in carbon treatment phase.	Carbon used as advanced treatment of biologically & chemically treated wastewater. Plant capacity 0.66 M ³ /sec.	64
IX M- 8	Phenanthrene	B,L	P	100 ppb	80% reduction; 6% desorbed from carbon by elutriation w/solvent.	See IXM- 1 for comments.	20
1X M 9	Pyrene	B,L	Р	100 ppb	80% reduction; 5% desorbed from carbon by elutriation w/solvent.	See IXM- 1 for comments.	20
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-						(continue	(d)

TABLE C-1 (continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Alcohols (A)

					: Alconois (A)		
No.	Chemical b	Descr Study Type C	iption d Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
XA- 1	Butanol	B,L	P	100 µg/1	Complete removal. 38% desorption of butanol by elutriation with solvent was achieved.	Resin was Amberlite XAD-2. Resin found to be more effective than carbon for phthalate esters, most aromatics, and pesticides. Carbon was more efficient for alkanes; neither effec- tive for acidic com- pounds.	20
XA- 2	Cyclohexanol	B,L	P	100 Aug/1	Complete removal. 81% description of cyclohexanol by elutriation with solvent was achieved.	See XA-1 for additional results.	20
XA- 3	Decanol	B,L	P	100 Aug/1	Complete removal. 89% de- sorption of decanol by elutriation with solvent was achieved.	See XA-1 for additional results.	20
XA- 4	2-Ethyl-l- Hexanol	B,L	р	100 /ug/1	Complete removal. 100% de- sorption of 2-Ethyl-1-Hexa- nol by elutriation with sol- vent was achieved.	See XA-l for additional results.	20
XA- 5	m-Heptanol	B,L	P	100 Aug/1	Complete removal. 100% de- sorption of n-Heptanol by elutriation with solvent was achieved.	See XA-l for additional results.	20
XA- 6	Octanol	B,L	P	100 Alg/1	Complete removal. Greater than 100% desorption of Octanol by elutriation with solvent was reported.	See XA-1 for additional results.	20
						(continue	ed)

TABLE C-1(continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Alcohols (A)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
XA- 7	Pentanol	B,L	P	100 J ug/1	Complete removal. 67% de- sorption of pentanol by elutriation with solvent was achieved.	See XA-1 for additional results.	20
-AX	Propanol	B,L	P	100 J ug/1	Complete removal. Propanol could not be desorbed by elutriation with solvent.	See XA-1 for additional results.	20
						(continue	(d)

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TABLE C-1 (continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Aliphatics (B)

					· Aliphacies (B)		
No.	Chemical	Descr Study Type ^C	iption o Waste Type d	Influent Char.	Results of Study	Comments	Ref.
XB- 1	•	B,L	P	100 ag/1	tion from resin by elutria- tion with solvent.	Resin was Amberlite XAD-2. Resin found to be more effective than carbon for phthalate esters, most aromatics, and pesticides; carbon more efficient for alkanes; neither effective for acidic compounds.	20
XB- 2	•	B,L	р	100 Aug/1	tion from resin by elutria- tion with solvent.	See XB-1 for additional results.	20
3 XB-	Decanoic Acid	B,L	P	100 /ug/1	100% reduction; No desorption from resin by elutriation with solvent.	See XB-1 for additional results.	20
XB- 4		B,L	P	100 /ug/1	25% reduction; No desorption from resin by elutriation with solvent.	See XB-1 for additional results.	20
XB- 5	•	·	P	100 µug/1	50% reduction; 4% desorption from resin by elutriation with solvent.	See XB-1 for additional results.	20
XB- 6	Hexadecane	B,L	Р	100 /ug/l	25% reduction; No desorption from resin by elutriation with solvent.	See XB-1 for additional results.	20
XB- 7	•	B,L	Р	100 /ug/1	100% reduction; No desorption from resin by elutriation with solvent.	See XB-1 for additional results.	20
8 XB-	Methyl Decanoate	B,L	P	100 µg/1	100% reduction; 50% desorption from resin by elutriation with solvent.	See XB-1 for additional results. (continue	

Concentration Process: Resin Adsorption (X)
Chemical Classification: Aliphatics (B)

No.	Chemical b	<u>Descr</u> Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
XB- .9	Methyl Dodecanoate	B,L	Þ	100 Aug/1	100% reduction; 72% desorption from resin by elutriation with solvent.	See XB-l for additional results.	20
XB- 10	decanoate	B,L	P	100 Aug/1	100% reduction; 67% desorp- tion from resin by elutria- tion with solvent.	See XB-1 for additional results.	20
XB- 11	decanoate	B,L	P	100 Aug/l	100% reduction; 54% desorp- tion from resin by elutria- tion with solvent.	See XB-1 for additional results.	20
XB- 12	Myristic Acid	B,L	P	100 Aug/1	100% reduction; No desorp- tion from resin by elutria- tion with solvent.	See XB-l for additional results.	20
XB- 13	Octadecane	B,L	P	100 /ug/1	25% reduction; No desorp- tion from resin by elutria- tion with solvent.	See XB-1 ' for additional results.	20
XB- 14	Octanoic Acid	B,L	P	100 <u>/</u> ug/1	90% reduction; No desorp- tion from resin by elutria- tion with solvent.	See XB-l for additional results.	20
XB- 15	Propionic Acid	B,L	P	100 /ug/1	100% reduction; No desorp- tion from resin by elutria- tion with solvent.	See XB-1 for additional results.	20
XB- 16	Pyruvic Acid	B,L	P	100 /ug/1	100% reduction; No desorp- tion from resin by elutria- tion with solvent.	See XB-1 for additional results.	20
XB- 17	Tetradecane	B,L	P	100 /ug/l	50% reduction; 23% desorp- tion from resin by elutria- tion with solvent.	See XB-l for additional results.	20
XB- 18	Valeric Acid	B,L	P	100 Aug/1	50% reduction; 2% desorp- tion from resin by elutria- tion with solvent.	See XB-1 for additional results.	20
					•	(continue	:d)

TABLE C-1 (continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Amines (C)

No.	Chemical b	Descr Study Type ^C		f Study Influent Char.	Results of Study	Comments	Ref.
XC- 1	Aniline	B,L	P	100 /ug/1	Complete removal; No desorp- tion from resin by elutria- tion with solvent.	Resin was Amberlite XAD-2. Resin found to be more effective than carbon for phthalate esters, most aromatics, and pesticides; carbon was more efficient for alkanes; neither effective for acidic compounds.	
XC- 2	Butylamine	B,L	P	100 Aug/1	Complete removal; 74% desorption from resin by elutriation with solvent.	for additional results.	
XC-	Cyclohexyl- amine	B,L	P	100 /ug/1	Complete removal; 94% desorption from resin by elutriation with solvent.	See XC-1 for additional results.	20
XC-	Dibutylamine	B,L	P	100 Alg/1	Complete removal; 62% desorption from resin by elutriation with solvent.	See XC-1 for additional results.	20
XC-	Dihexylamine	B,L	P	100 Alg/1	Complete removal; 11% desorption from resin by elutriation with solvent.	See XC-1 for additional results.	
XC-	Dimethylamine	B,L	P	100 /ug/l	100% removal; 50% desorption from resin by elutriation with solvent.	See XC-1 for additional results.	
7	Hexylamine	B,L	P	100 /ug/1	100% removal; 110% desorption from resin by elutriation with solvent.	See XC-1 for additional results.	20
						(continu	 .ed)

Concentration Process: Resin Adsorption (X) Chemical Classification: Amines (C)

	·							
No.	Chemical b	Descr Study Type ^C		Influent Char.	Results of Study Comments	Ref.		
8 XC-	Morpholine	B,L	P	100 /ug/1	100% removal; 28% desorption See XC-1 from resin by elutriation for additional results. with solvent.	20		
XC- 9	Octylamine	B,L	P	100 /ug/1	100% removal; 15% desorption See XC-1 from resin by elutriation for additional results. with solvent.	20		
XC- 10	Piperidine	B,L	P	100 ug/1	100% removal; 42% desorption See XC-1 from resin by elutriation for additional results. with solvent.	20		
XC- 11	Pyrrole	B,L	P	100 µg/1	100% removal; 5% desorption See XC-1 from resin by elutriation for additional results. with solvent.	20		
XC- 12	Tributylamine	B,L	P	100 µg/1	100% removal; 108% desorption See XC-1 from resin by elutriation for additional results. with solvent.	20		
	Ī				(continued	1)		

TABLE C-1 (continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Aromatics (D)

no.	Chemical b	<u>Descr</u> Study Type ^C	iption o Waste Type d	of Study Influent Char.	Results of Study	Comments	Ref.
XD- 1	Acetophenone	B,L	P	100 µg/1	100% reduction; 80% desorption from resin by elutriation with solvent.	Resin was Amberlite XAD-2. Resin found to be more effective than carbon for phthalate esters, most aromatics, and pesticides; carbon more efficient for alkanes; neither effective for acid compounds.	20
XD- 2	Benzaldehyde	B,L	P	100 pg/1	100% reduction; 79% desorp- tion from resin by elutria- tion with solvent.	See XD-1 for additional results.	20
J	Benzil	B,L	P	100 µg/1	100% reduction; 63% desorp- tion from resin by elutria- tion with solvent.	See XD-1 ' for additional results.	20
XD-4	Benzoic Acid	B,L	P		100% reduction; No desorp- tion from resin by elutria- tion with solvent.	See XD-1 for additional results.	20
	Benzene, Toluene, Xylene (BTX)	P	I	20 to 300 ppm	Effluent (leakage) is 0.2ppm	Costs estimated to be \$3.36/1000 gal. at 250 gpm and 300 ppm BTX. Resin regenerant is steam. Recovery of BTX reduces costs to \$1.09/1000 gal.	32
XD-	Cumene	B,L	P	100 jng/1	100% removal; 63% desorption from resin by elutriation with solvent.	See XD-1 for additional results.	20
						(continue	d)

TABLE C-1(continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Aromatics (D)

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No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study Comments	Ref.
	m-Dichloro- benzene	B,L	P	100 µg/1	1 100% removal; 52% desorption from resin by elutriation for additional results.	20
8	o-Dichloro- benzene	B,L	P	100 µg/1	1 100% removal; 61% desorption See XD-1 from resin by elutriation for additional results. with solvent.	20
	p-Dichloro- benzene	B,L	P	100 µg/1	100% removal; 35% desorption See XD-1 from resin by elutriation for additional results. with solvent.	20
10	1,2,4-Trichloro- benzene		P	100 µg/1	100% removal; 67% desorption See XD-1 from resin by elutriation for additional results. with solvent.	20
	2,4,6-Trinitro- toluene (TNT)	P,C	I	81 to 116 ppm	Resin adsorption capacity was 0.116 to 0.154 gm/gm at 1 ppm breakthrough. No loss in capacity after 15 regeneration cycles. 1 ppm breakthrough occurred after 633 to 1193 B.V. Amberlite XAD-4 used; acetone regenerant. Less costly than carbon due to regenerability.	2
12	2,4,6-Trinitro- toluene (TNT) and other muni- tions plant wastewaters: Cyclonite(RDX), Nitramine (Tetryl) and cyclotetrameth- ylene tetrani- tramine (HMX).	R	I	Not reported	Adsorption capacities (Lb/Lb For 80 gpm facility Amberlite XAD-4 resin): costs estimated to be Contami	40
7	a - {	• •	ţ	•	(continued	1)

TABLE C-1(continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Aromatics (D)

	_	Descr	iption o	f Study			
No.	Chemical b	Study Type C	Waste Type d	Influent Char.	Results of Study	Comments	Ref
XD-					(Note: breakthrough conc. not		
12					defined.)		
cont.					Typical conc. of contaminants in wastewaters:		
					TNT - 0-400 ppm		
					RDX - 50-100 ppm pH - 3.5-7.0		
					Flow - 0.02-1.0 MGD Temp - 60-160 ^O F		
					Tomp of 200		
	·						
	·						
						(continu	l ed)

77

Concentration Process: Resin Adsorption (X) Chemical Classification: Halocarbons (F)

							
No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
XF- 1	Bromoform	L	W	0.2 ppb		See XF-16 for results.	46
XF – 2	Bromoform	B,L	P	100 ppb	100% removal; 28% de- sorption from resin by elutriation w/solvent.	Amberlite XAD-2 used Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, methyl chloride-acetone, and acetone.	
XF- 3	Bromodichlo- methane	L	W		At 2 ppm, equilibriu m capacity was 48 mg/g.	See XF-16 for results.	46
XF – 4	Carbon Tetrachlo- ride	P		100 to 7000 ppm chlori- nated hydro- carbons	Effluent of <pre><lppm pre="" total<=""></lppm></pre>	Steam used as regen	
XF- 5	Chloroform	Þ		100 to 7000 ppm chlori- nated hydro- carbons	Effluent of <pre><pre>chlorinated hydrocarbons</pre> could be achieved.</pre>	See XF- 4	32
XF - 6	Chloroform	L	W	l.l ppb	At 2ppm, equilibrium capacity was 50 mg/g.	See XF-16 for results. (continue	46 d)

TABLE C-1(continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Halocarbons (F)

		CHEMILO	ar Clas	silication	: Halocarbons (F)		
No.	Chemical b	<u>Descr</u> Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref
XF- 7	Dibromochlo- romethane	L	W	3.9 ppb		See XF-16 for results.	46
	1,1-Dichlo- roethane	L	W	2.3 ppb		See XF-16 for results.	46
XF- 9	roethane	L	W	2.1 ppb		See XF-16 for results.	46
	roethylene	L	W	0.2 ppb		See XF-16 for results.	46
XF-	Dichloride	₽	Ι	100 to 7000 ppm chlori- nated hydro- carbons	Effluent of <1ppm total chlorinated hydrocarbons could be achieved.	See XF-4 for comments.	32
XF- 12	Hexachloro- butadiene	B,L	P	100 ppb	100% removal; 73% de- sorption from resin by elutriation with solvent	See XF-2 for comments.	20
XF- 13	Hexachloro- ethane	B,L	P	100 ppb	100% removal; 55% de- sorption from resin by elutriation with solvent	See XF-2 for comments.	20
XF- 14	Tetrachloro- ethane	B,L	P	100 ppb	100% removal; 53% de- sorption from resin by elutriation with solvent	See XF-2 for comments.	20
XF- 15	Tetrachloro- ethylene	L	W	179 ppb		See XF-16 for results.	46
XF-	1,1,1-Tri- chloroethane	L	W	551 ppb	Performance for treat- ment of water containing several halogens	Column studies: 14 mm	

TABLE C-1 (continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Halocarbons (F)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study Comments	Ref.
XF- 16 cont					Virgin Regenerated BV to 9000 8500 33 ppb com- pound leakage Virgin Regenerated (16 BV/hr) Regener- ated at 37 lb steam/ cu ft @ 5 psig	
					Days 23.4 22.1 Gal treated/ cu ft 67500 63750 sorbent	
XF- 17	1,2,3-Tri- chloropro- pane	B,L	P	100 ppb		20
	,					
					(continued	1)

TABLE C-1(continued)

Concentration Process: Resin Adsorption (X)
Chemical Classification: Polychlorinated Biphenyls (I)

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No.	Chemical b	Descr Study Type ^C	Waste	f Study Influent Char.	Results of Study	Comments	Ref.
xI- 1	Arochlor 1254	B,L	Þ	100 ppb	100% reduction; 76.6% de- sorbed from carbon by elutriation w/solvent.	Amberlite XAD-2 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, chloroform-acetone & acetone.	20
XI- 2	Arochlor 1254	C,L	P		Final effluent conc. was 0-0.25 ppb for 192 B.V.	5 day study.	22
XI- 3	Arochlor 1254 & 1260	С	М		60% reduction w/Amberlite XAD-4. 23% ± 2% reduction w/Amberlite XAD-2.	In continuous flow system reduction de-creased greatly w/time.	57
						(continu	ed)
		1		1		(continue	ed)

Concentration Process: E Chemical Classification:

Resin Adsorption (X): Pesticides (J)

Description of Study Chemical b No. Waste Influent Study Results of Study Comments Ref. Type d Char. Type C XJ-Aldrin B,L P 100 ppb 100% reduction; 39% desorbed Amberlite XAD-2 used. 20 from resin by elutriation Solvents included penw/solvent. tane-acetone, diethyl ether, methylene chloride-acetone, chloroform-acetone and acetone XJ-Atrazine B,L P 100% reduction; 38% desorbed 100 ppb See XJ-1 20 from resin by elutriation for comments. w/solvent. Chlorinated XJ-L Ι Column studies indicatd that 33 to Solvents ranking in 49 Pesticides Amberlite XAD-4 could pro-118 ppm terms of decreasing ef-(Unspecified) cess about four times more fectiveness were acetone, throughput before experiencisopropanol, and methaing some leakage as carbon nol; however, acetone column. Leakages of <1 ppm is very flammable. Colmaintained at longer than umn study conditions: 120 BV. Resin could be ef-50-150 BV passed, 4 BV/hr fectively regenerated w/2 BV flow, 12.5-125 hr duraof isopropanol whereas even tion. Costs estimated 8 BV did not effectively to be \$0.83 for resin generate carbon. sorption and \$1.33/1000 gal for carbon. XJ-2,4-D Butyl B,L P 100 ppb 100% reduction; 10% desorbed See XJ-1 20 ester from resin by elutriation for comments. w/solvent. 2,4-D and re-XJ-U I 20-1500 Effluent conc. reduced to Amberlite XAD-4 resin 20 lated herbippm @70-<1.0 ppm. used. cides map 08

(continued)

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Concentration Process: Resin Adsorption (X)
Chemical Classification: Pesticides (J)

	1	Descr	iption o	of Study			1
No.	Chemical b	Study Type ^C		Influent Char.	Results of Study	Comments	Ref
ХЈ- 6	DDT	B,L	P	100 ppb	100% reduction; 49% desorbed from resin by elutriation w/solvent.	See XJ-1 for comments.	20
XJ- 7	Endrin and Heptachlor	F	I	0.1-2.0 ppm @ 100 gpm	Effluent conc. reduced to <3.0 ppb.	Amberlite XAD-4 used.	32
XJ- 8	Toxaphene	Ü	I	70-2600 ppb	Effluent conc. reduced to 0.1-4.2 ppb.	Amberlite XAD-4 used.	32
		1				(continu	ied)

Concentration Process: Resin Adsorption (X)
Chemical Classification: Phenols (K)

No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref
1 1	Bisphenol-A	C,L	I	900 ppm 2 BV/hr	At pH 11.4, poor adsorption achieved on either XAD-4 or XAD-7. At pH 10.0, XAD-4 treated 33.5 B.V.'s to 50ppm breakthrough. XAD-7 treated 16 B.V. to 50 ppm breakthrough.	95% regeneration achieved w/l B.V. of 4% NaOH & 4 B.V. deionized water.	23
XK- 2	Bisphenol-A	C,L	I	280 ppm 2 BV/ r	At pH 6.9, XAD-4 capacity was 34 g/l and XAD-7 capa-city was 16 g/l.	See XK-1 for comments.	23
XK- 3	Brine Phenol	U	I	20% brine w/10-150 ppm phenol		Wastewater of brine purification process 5 B.V. of 4% NaOH required for regeneration	33
XK- 4	Brine Phenol	υ	I	10% brine w/10-400 ppm phenol		Wastewater from a	33
XK- 5	4-Chloro-3- methylphenol	B,L	P	100 ppb	100% reduction; 70% de- sorbed from resin by elutriation w/solvent.	Amberlite XAD-2 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, chloroform-acetone and acetone.	20
ХК- 6	m-Chlorophenol w/13% NaCl	U	I	350 ppm @ 0.5 gpm/ft ³	At zero leakage sorption capacity was 0.07 lb/lb.	15 min contact time Amberlite XAD-4 used.	66
						(continue	d)

Concentration Process:

Resin Adsorption (X)

Chemical Classification: Phenols (K)

No.	Chemical b	Descr Study Type C		of Study Influent Char.	Results of Study	Comments	Ref.
XK- 7	2,4-Dibromo- phenol	B,L	P	100 ppb	100% reduction; 44% desorbed from resin by elutriation w/solvent.	See XK-5 for comments.	20
8 XK-	1 *	U	I	1500 ppm w/15% brine, pH = 2-3	Resin capacity was 5.6 lb phenols/ft ³ @ 5 ppm break-through.	Amberlite XAD-2 used. 2% caustic soda heated to 80°-85°C used as regenerant.	33
ХК- 9	1	B,L	P	100 ppb	100% reduction; 54% desorbed from resin by elutriation w/solvent.	See XK-5 for comments.	20
XK- 10	1 '	U	I	430 ppm @ 0.5 gpm/ft ³	At zero leakage sorption capacity was 0.116 lb/lb.	15 min contact time. Amberlite XAD-4 used.	66
XK-		B,L	P	100 ppb	100% reduction; 76% desorbed by elutriation w/solvent.	See XK-5 for comments.	20
12	1	C,L	Ι	700-1300 ppm @ 50°C	Effluent conc. reduced to 5.0-6.0 ppm for 32 B.V. Resin capacity was about 40 g/l. Efficient ethanol regeneration.	Amberlite XAD-7 used. 20 ml columns used w/experimental runs of up to 40 B.V.	23
13		U	I	1000- 1800 ppm @ pH=2.0	Effluent conc. reduced to 1-5 ppm by cross-linked polystyrene adsorbent resin.	Effluent from parathion manufacturer. 4% aque- ous caustic soda (2B.V.) followed by water rinse used as regnerant.	
14	1	B,L	P	100 ppb	100% reduction; 60% desorbed from resin by elutriation w/solvent.	See XK-5 for comments.	20
						(continue	ed)

TABLE C-1 (continued)

Concentration Process: Resin Adsorption (X) Chemical Classification: Phenols (K)

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No.	Chemical b	Descr Study Type ^C		of Study Influent Char.	Results of Study	Comments	Ref.
XK- 15	Phenol	C,L	P	6700 ppm	Effluent conc. of <1.0 ppm achieved.	Amberlite XAD-4 used. Acetone & methanol used as regenerants.	23
хк- 16	Phenol	U	I	500-1500 ppm	Effluent conc. of 1.0-3.0ppm achieved.	Amberlite XAD-4 used. Wastewater from Bisphenol A manufacturer containing 0.5-1.5% phenol, 0.5-1.0% NaCl, 100-1000 ppm acetone @ pH=0.2-1.5. Acetone & methanol used as regenerant.	
XK- 17	Phenol	Ū	Ī	5000 ppm	Effluent conc. reduced to <25 ppm.	Wastewater from phenolic resin manufacturer. Warm 44% formaldehyde used as regenerant.	33
XK- 18	Regordinol	B,L	P	100 ppb	100% reduction; 35% desorbed from resin by elutriation w/solvent.	See XK-5 for comments.	20
XK- 19	2,4,6-Trichlo- rophenol	B,L	P	100 ppb	100% reduction; 60% desorbed from resin by elutriation w/solvent.	See XK-5 for comments.	20
XK- 20	2,4,6-Trichlo- rophenol	υ	I	510 ppm @ 0.5 gpm/ft ³	At zero leakage sorption capacity was 0.272 lb/lb.	15 min contact time. Amberlite XAD-4 used.	66
					,	(continue	d)

Concentration Process: Resin Adsorption (X) Chemical Classification: Phthalates (L)

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No.	Chemical b	Descr Study Type ^C	iption o Waste Type	of Study Influent Char.	Results of Study	Comments	Ref.
XL- 1	Dibutyl Phthalate	B,L	Þ	100 ppb	100% reduction; 108% desorbed from resin by elutriation w/solvent.	Amberlite XAD-2 used. Solvents included pentane-acetone, diethylether, methylene chloride-acetone, chloroform-acetore & acetone.	20
XL- 2	Diethylhexyl Phthalate	B,L	P	100 ppb	100% reduction; 76% desorbed from resin by elutriation w/solvent.	See XL-1 for comments.	20
3 	Dimethyl Phthalate	B,L		100 ppb	100% reduction; 62% desorbed from resin by elutriation w/solvent.	See XL-1 for comments.	20
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Concentration Process: Resin Adsorption (X)
Chemical Classification: Polynuclear Aromatics (M)

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a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
XM- 1	Acenapththa- lene	B,L	P	100 ppb	100% reduction; 78% desorbed from resin by elutriation w/solvent.	Amberlite XAD-2 used. Solvents included pentane-acetone, diethyl ether, methylene chloride-acetone, chloroform-acetone & acetone.	20
XM- 2	Biphenyl	B,L	P	100 ppb	100% reduction; 73% desorbed from resin by elutriation w/solvent.	See XM-1 for comments.	20
XM- 3	Cumene	B,L	P	100 ppb	100% reduction; 63% desorbed from resin by elutriation w/solvent.	See XM-1 for comments.	20
XM- 4	Dimethyl- naphthalene	B,L	P	100 ppb	100% reduction; 90% desorbed from resin by elutriation w/solvent.	See XM-1 for comments.	20
ХМ- 5	Fluoranthrene	B,L	P	100 ppb	100% reduction; 66% desorbed from resin by elutriation w/solvent.	See XM-1 for comments.	20
XM- 6	Phenanthrene	B,L	P	100 ppb	100% reduction; 41% desorbed from resin by elutriation w/solvent.	See XM-1 for comments.	20
ХМ 7	Pyrene	B,L	P	100 ppb	100% reduction; 63% desorbed from resin by elutriation w/solvent.	See XM-1 for comments.	20
						(continue	ed)

TABLE (continued)

Concentration Process: Miscellaneous Sorbents (XII) Chemical Classification: Metals (G)

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a No.	Chemical b	Descr Study Type ^C	Waste	of Study Influent Char.	Results of Study	Comments	Ref.
XII G- 1	Arsenic	R	ប	25 ppm	Effluent conc. of 1.0ppm achieved.	Silicon alloy used.	90
XII G- 2	Cadmium	R	บ	25 ppm	Effluent conc. of 1.0ppm achieved.	Silicon alloy used.	90
XII G- 3	Chromium	R	Ü	300 ppm	100% removal.	High clay soil used	90
XII G- 4	Copper	R	Ŭ	300 ppm	100% removal.	High clay soil used	90
XII G- 5	Copper	R	Ü	25 ppm	Effluent conc. of 1.0ppm achieved.	Silicon alloy used.	90
XII G- 6	Lead	R	Ü		Residual of <5.0 mg/l achieved.	Ground redwood bark used.	90
XII G- 7	Lead	R	Ü	25 ppm	Effluent conc. of 1.0ppm achieved.	Silicon alloy used.	90
XII G- 8	Mercury	R	Ü	25 ppm	Final conc. of 10 ppb achieved.	Silicon alloy used.	90
XII G- 9	Zinc	R	U	10 ppm	Final conc. reduced to 0.1 ppb.	SiO ₂ & CaO slags used.	90
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147

Concentration Process: Miscellaneous Sorbents (XII)
Chemical Classification: Polychlorinated Biphenyls (I)

No.	Chemical b		Description of Study tudy Waste Influent ype C Type d Char.		Results of Study	Comments	Ref.
XII I- 1		С	М		73% reduction in raw sewage w/PVC chips. Polyurethane foam adsorbed 35% ± 3%.	In continuous flow system reduction decreased greatly w/time.	57

Footnotes:

- a. Three part code number assigned to each individual chemical compound. First part is a Roman numeral which corresponds to the concentration process code number. Second part is a capital letter corresponding to the chemical classification code number. Third part is unique number for each individual compound.
- b. Chemicals are presented in alphabetical order generally according to The Merck Index preferred or generic name. However, it is recommended to check for a compound under several potential names.
- c. Describes the scale of the referenced study:

B - Batch Flow

C - Continuous Glow

F - Full Scale

I - Isotherm Test

L - Laboratory Scale

N - Flow Not Controlled

O - Respirometer Study

P - Pilot Scale

R - Literature Review

S - Slug Dose Chemical Addition

U - Unknown

Footnotes (continued):

- d. Describes the type of wastewater used in the referenced study:
 - D Domestic wastewater
 - H Hazardous material spill
 - I Industrial wastewater
 - P Pure Compound (one solute in a solvent)
 - R River water
 - S Synthetic wastewater
 - U Unknown
 - W Well water