#### METHOD 8270D

# <u>SEMIVOLATILE ORGANIC COMPOUNDS</u> BY GAS CHROMATOGRAPHY/MASS SPECTROMETRY (GC/MS)

SW-846 is not intended to be an analytical training manual. Therefore, method procedures are written based on the assumption that they will be performed by analysts who are formally trained in at least the basic principles of chemical analysis and in the use of the subject technology.

In addition, SW-846 methods, with the exception of required method use for the analysis of method-defined parameters, are intended to be guidance methods which contain general information on how to perform an analytical procedure or technique which a laboratory can use as a basic starting point for generating its own detailed Standard Operating Procedure (SOP), either for its own general use or for a specific project application. The performance data included in this method are for guidance purposes only, and are not intended to be and must not be used as absolute QC acceptance criteria for purposes of laboratory accreditation.

## 1.0 SCOPE AND APPLICATION

1.1 This method is used to determine the concentration of semivolatile organic compounds in extracts prepared from many types of solid waste matrices, soils, air sampling media and water samples. Direct injection of a sample may be used in limited applications. The following RCRA analytes have been determined by this method:

		Approp	riate Prepa	aration T	echnique	es <sup>b</sup>
				3540/		
Compounds	CAS No <sup>a</sup>	3510	3520	3541	3550	3580
Acenaphthene	83-32-9	1 (X	$11 \times 0$	) [X]	S X (	X
Acenaphthylene	208-96-8	Χ	Χ	X	Χ	Χ
Acetophenone	98-86-2	Χ	ND	ND	ND	X
2-Acetylaminofluorene	53-96-3	Χ	ND	ND	ND	Χ
1-Acetyl-2-thiourea	591-08-2	LR	ND	ND	ND	LR
Aldrin	309-00-2	Χ	Χ	Χ	Χ	Χ
2-Aminoanthraquinone	117-79-3	Χ	ND	ND	ND	X
Aminoazobenzene	60-09-3	Χ	ND	ND	ND	Χ
4-Aminobiphenyl	92-67-1	Χ	ND	ND	ND	Χ
3-Amino-9-ethylcarbazole	132-32-1	Χ	Χ	ND	ND	ND
Anilazine	101-05-3	Χ	ND	ND	ND	Χ
Aniline	62-53-3	Χ	Χ	ND	Χ	Χ
o-Anisidine	90-04-0	Χ	ND	ND	ND	Χ
Anthracene	120-12-7	Χ	Χ	Χ	Χ	Χ
Aramite	140-57-8	HS	ND	ND	ND	Χ
Aroclor 1016	12674-11-2	Χ	Χ	Χ	Χ	Χ
Aroclor 1221	11104-28-2	Χ	Χ	Χ	Χ	Χ
Aroclor 1232	11141-16-5	Χ	Χ	Χ	X	X
Aroclor 1242	53469-21-9	Χ	Χ	Χ	X	X
Aroclor 1248	12672-29-6	Χ	Χ	Χ	Χ	Χ

		Appropriate Preparation Technique			echnique	es <sup>b</sup>
				3540/		
Compounds	CAS No <sup>a</sup>	3510	3520	3541	3550	3580
Aroclor 1254	11097-69-1	X	X	X	X	Χ
Aroclor 1260	11096-82-5	Х	Х	Χ	Χ	Χ
Azinphos-methyl	86-50-0	HS	ND	ND	ND	Χ
Barban	101-27-9	LR	ND	ND	ND	LR
Benzidine	92-87-5	CP	CP	CP	CP	CP
Benzoic acid	65-85-0	Χ	Χ	ND	Χ	Χ
Benz(a)anthracene	56-55-3	Χ	Χ	Χ	Χ	Χ
Benzo(b)fluoranthene	205-99-2	Χ	Х	Χ	Χ	Χ
Benzo(k)fluoranthene	207-08-9	Χ	Χ	Χ	Χ	Χ
Benzo(g,h,i)perylene	191-24-2	Χ	Χ	X	X	Χ
Benzo(a)pyrene	50-32-8	Х	X	X	X	Χ
<i>p</i> -Benzoquinone	106-51-4	OE	ND	ND	ND	Χ
Benzyl alcohol	100-51-6	Χ	X	ND	X	Χ
α-BHC	319-84-6	Χ	X	X	Х	Χ
β-ВНС	319-85-7	Х	Х	X	Х	Χ
δ-BHC	319-86-8	Х	Х	Χ	Х	Χ
γ-BHC (Lindane)	58-89-9	Х	Х	X	Х	Χ
Bis(2-chloroethoxy)methane	111-91-1	Х	Х	Χ	Х	Χ
Bis(2-chloroethyl) ether	111-44-4	X	X	X	X	Χ
Bis(2-chloroisopropyl) ether	39638-32-9	X	X	X	X	χ
Bis(2-ethylhexyl) phthalate	117-81-7	X	$\perp$ x	X	X	X
4-Bromophenyl phenyl ether	101-55-3	X	X	X	X	Χ
Bromoxynil	1689-84-5	Х	ND	ND	ND	Χ
Butyl benzyl phthalate	85-68-7	X	X	X	X	Χ
Captafol	2425-06-1	III HS	111	ND	O <sub>ND</sub>	$A(\mathbf{x}, 0)$
Captan	133-06-2	HS	ND	ND	ND	X
Carbaryl	63-25-2	X	ND	ND	ND	X
Carbofuran	1563-66-2	X	ND	ND	ND	X
Carbophenothion	786-19-6	X	ND	ND	ND	X
Chlordane (NOS)	57-74-9	X	X	X	Х	X
Chlorfenvinphos	470-90-6	X	ND	ND	ND	X
4-Chloroaniline	106-47-8	X	ND	ND	ND	X
Chlorobenzilate	510-15-6	X	ND	ND	ND	X
5-Chloro-2-methylaniline	95-79-4	X	ND	ND	ND	X
4-Chloro-3-methylphenol	59-50-7	X	X	X	X	X
3-(Chloromethyl)pyridine	6959-48-4	X	ND	ND	ND	X
hydrochloride					ND	
1-Chloronaphthalene	90-13-1	X	X	X	X	Χ
2-Chloronaphthalene	91-58-7	Х	Х	X	Χ	Χ
2-Chlorophenol	95-57-8	Х	Х	X	Χ	Χ
4-Chloro-1,2-phenylenediamine	95-83-0	Χ	Х	ND	ND	ND
4-Chloro-1,3-phenylenediamine	5131-60-2	Χ	Х	ND	ND	ND
4-Chlorophenyl phenyl ether	7005-72-3	Χ	Х	Χ	Χ	Χ

	<u>_</u>	Appropr	iate Prepa	aration T	echnique	es <sup>b</sup>
Compounds	CAS Noª	3510	3520	3540/ 3541	3550	3580
Chrysene	218-01-9	X	Χ	Χ	Χ	Χ
Coumaphos	56-72-4	X	ND	ND	ND	Χ
p-Cresidine	120-71-8	X	ND	ND	ND	Χ
Crotoxyphos	7700-17-6	X	ND	ND	ND	Χ
2-Cyclohexyl-4,6-dinitro-phenol	131-89-5	X	ND	ND	ND	LR
4,4'-DDD	72-54-8	X	Χ	Χ	Χ	Χ
4,4'-DDE	72-55-9	X	Χ	Χ	Χ	Χ
4,4'-DDT	50-29-3	X	Χ	Χ	Χ	Χ
Demeton-O	298-03-3	HS	ND	ND	ND	Χ
Demeton-S	126-75-0	X	ND	ND	ND	Χ
Diallate (cis or trans)	2303-16-4	X	ND	ND	ND	Χ
2,4-Diaminotoluene	95-80-7	DC, OE	ND	ND	ND	Χ
Dibenz(a,j)acridine	224-42-0	Χ	ND	ND	ND	X
Dibenz(a,h)anthracene	53-70-3	X	X	X	X	X
Dibenzofuran	132-64-9	X	X	ND	X	X
Dibenzo(a,e)pyrene	192-65-4	ND	ND	ND	ND	X
1,2-Dibromo-3-chloropropane	96-12-8	X	X	ND	ND	ND
Di-n-butyl phthalate	84-74-2	X	X	X	X	Х
Dichlone	117-80-6	OE OE	ND	ND	ND	X
1,2-Dichlorobenzene	95-50-1	X	X	X	X	X
1,3-Dichlorobenzene	541-73-1	$\begin{pmatrix} x \\ x \end{pmatrix}$	X	X	X	x
1,4-Dichlorobenzene	106-46-7	X	X	X	X	X
3,3'-Dichlorobenzidine	91-94-1	X	X	X	X	X
2,4-Dichlorophenol	120-83-2		X	X	_ X	X
2,6-Dichlorophenol	87-65-0	fix I	n 🖟 C		OND 8	$11^{\circ}_{X}$
Dichlorovos	62-73-7	X	ND	ND	ND	X
	141-66-2	X	ND	ND	ND	X
Dicrotophos						
Dieldrin  Diethyl phthalata	60-57-1	X	X	X	X	X
Diethyl phthalate	84-66-2	X AM, OS	X	X	X	X
Diethylstilbestrol	56-53-1	AW, OS	ND	ND	ND	X
Diethyl sulfate	64-67-5	LR UE UC	ND	ND	ND	LR
Dimethoate	60-51-5	HE, HS	ND	ND	ND	Х
3,3'-Dimethoxybenzidine	119-90-4	X	ND	ND	ND	LR
Dimethylaminoazobenzene	60-11-7	X	ND	ND	ND	Х
7,12-Dimethylbenz(a)-anthracene	57-97-6	CP	ND	ND	ND	CP
3,3'-Dimethylbenzidine	119-93-7	X	ND	ND	ND	X
α,α-Dimethylphenethylamine	122-09-8	ND	ND	ND	ND	X
2,4-Dimethylphenol	105-67-9	X	X	X	X	X
Dimethyl phthalate	131-11-3	X	X	X	Х	Х
1,2-Dinitrobenzene	528-29-0	Χ	ND	ND	ND	Χ
1,3-Dinitrobenzene	99-65-0	Χ	ND	ND	ND	Χ
1,4-Dinitrobenzene	100-25-4	HE	ND	ND	ND	Χ
4,6-Dinitro-2-methylphenol	534-52-1	X	Χ	Χ	Χ	Χ

		Appropr	iate Prep	aration T	echnique	es <sup>b</sup>
				3540/		
Compounds	CAS No <sup>a</sup>	3510	3520	3541	3550	3580
2,4-Dinitrophenol	51-28-5	X	Χ	Χ	Χ	Χ
2,4-Dinitrotoluene	121-14-2	X	X	X	Χ	Χ
2,6-Dinitrotoluene	606-20-2	X	X	X	Χ	Χ
Dinocap	39300-45-3	CP, HS	ND	ND	ND	CP
Dinoseb	88-85-7	X	ND	ND	ND	Χ
Diphenylamine	122-39-4	X	X	Χ	Χ	Χ
5,5-Diphenylhydantoin	57-41-0	X	ND	ND	ND	Χ
1,2-Diphenylhydrazine	122-66-7	X	Χ	Χ	Χ	Χ
Di-n-octyl phthalate	117-84-0	X	X	X	Χ	Χ
Disulfoton	298-04-4	X	ND	ND	ND	X
Endosulfan I	959-98-8	X	X	Χ	Χ	X
Endosulfan II	33213-65-9	X	Х	Х	X	X
Endosulfan sulfate	1031-07-8	X	Х	Х	Х	X
Endrin	72-20-8	X	X	X	X	X
Endrin aldehyde	7421-93-4	X	X	X	X	X
Endrin ketone	53494-70-5	X	X	ND	X	X
EPN	2104-64-5	X	ND	ND	ND	X
Ethion	563-12-2	X	ND	ND	ND	X
Ethyl carbamate	51-79-6	DC	ND	ND	ND	X
Ethyl methanesulfonate	62-50-0	X	ND	ND	ND .	X
Famphur	52-85-7	X = X	ND	ND	ND	X
Fensulfothion	115-90-2	X	ND	ND	ND	X
Fenthion	55-38-9	X	ND	ND	ND	X
Fluchloralin	33245-39-5		ND	ND	ND	X
Fluoranthene	206-44-0	fig I	$n_{\mathbf{x}}^{\mathbf{z}}$	$\mathcal{I}_{X}^{X}$	$O_{X}^{-2}$	a t <u>x</u> LO
Fluorene	86-73-7	X	X	X	X	X
2-Fluorobiphenyl (surr)	321-60-8	X	X	X	X	X
2-Fluorophenol (surr)	367-12-4	X	X	X	X	X
Heptachlor	76-44-8	X	X	X	X	X
Heptachlor epoxide	1024-57-3	X	X	X	X	X
Hexachlorobenzene	118-74-1	X	X	X	X	X
Hexachlorobutadiene	87-68-3	X	X	X	X	X
Hexachlorocyclopentadiene	77-47-4	X	X	X	X	X
Hexachloroethane	67-72-1	X	X	X	X	X
Hexachlorophene	70-30-4	AW, CP	ND	ND	A ND	CP
Hexachloropropene	1888-71-7	X X	ND ND	ND ND	ND	X
Hexamethylphosphoramide	680-31-9	X	ND ND	ND ND	ND ND	X
• • •		ND				
Hydroquinone	123-31-9		ND ~	ND V	ND ×	X
Indeno(1,2,3-cd)pyrene	193-39-5	X	X	X	X	X
Isodrin	465-73-6	X	ND	ND	ND	X
Isophorone	78-59-1	X	X	X	X	X
Isosafrole	120-58-1	DC	ND	ND	ND	X
_Kepone	143-50-0	X	ND	ND	ND	X

Compounds         CAS No®         3510         3520         3541         3550         3580           Leptophos         21609-90-5         X         ND         ND         ND         X           Malathion         121-75-5         HS         ND         ND         ND         X           Maleic anhydride         108-31-6         HE         ND         ND         ND         X           Mestranol         72-33-3         X         ND         ND         ND         X           Methapyrilene         91-80-5         X         ND         ND         ND         X           Methoxychlor         72-43-5         X         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         ND           4,4'-Methylenebis (N,N-dimethyl-aniline)         101-61-1         X         X         ND         X         X         ND
Leptophos         21609-90-5         X         ND         ND         X           Malathion         121-75-5         HS         ND         ND         ND         X           Maleic anhydride         108-31-6         HE         ND         ND         ND         X           Mestranol         72-33-3         X         ND         ND         ND         ND         X           Methapyrilene         91-80-5         X         ND         ND         ND         X           Methoxychlor         72-43-5         X         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         X           4,4'-Methylenebis (N,N-dimethyl- aniline)         101-61-1         X         X         ND         X         X         ND         ND         ND         X         X         ND         ND         ND         X
Malathion         121-75-5         HS         ND         ND         ND         X           Maleic anhydride         108-31-6         HE         ND         ND         ND         X           Mestranol         72-33-3         X         ND         ND         ND         X           Methapyrilene         91-80-5         X         ND         ND         ND         X           Methoxychlor         72-43-5         X         ND         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-61-1         X         X         ND         ND         ND         ND         X           4,4'-Methylenebis (N,N-dimethyl-aniline)         101-61-1         X         X         ND         X         X         ND         ND         ND         X         X         ND         ND         ND         X         X
Maleic anhydride         108-31-6         HE         ND         ND         ND         X           Mestranol         72-33-3         X         ND         ND         ND         X           Methapyrilene         91-80-5         X         ND         ND         ND         X           Methoxychlor         72-43-5         X         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         LR           4,4'-Methylenebis (N,N-dimethyl-aniline)         101-61-1         X         X         ND         X         X         ND         ND         ND         X         X         X         ND         ND         ND         X         X         X         X         X         X         ND         ND         ND <t< td=""></t<>
Mestranol         72-33-3         X         ND         ND         ND         X           Methapyrilene         91-80-5         X         ND         ND         ND         X           Methoxychlor         72-43-5         X         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         LR           4,4'-Methylenebis (N,N-dimethyl-aniline)         101-61-1         X         X         ND         X         X         ND         ND         ND
Methapyrilene         91-80-5         X         ND         ND         ND         X           Methoxychlor         72-43-5         X         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         LR           4,4'-Methylenebis (N,N-dimethyl-aniline)         101-61-1         X         X         ND         X         X         A
Methoxychlor         72-43-5         X         ND         ND         ND         X           3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         LR           4,4'-Methylenebis (N,N-dimethylaniline)         101-61-1         X         X         ND         X         X         ND         ND         ND         ND         X         X         MD         ND         ND         ND         X         X         MD         ND         ND         ND         X         X         ND         ND         ND         ND         ND         X         X         ND         ND         ND         ND         X         X         ND         ND
3-Methylcholanthrene         56-49-5         X         ND         ND         ND         X           4,4'-Methylenebis (2-chloroaniline)         101-14-4         OE, OS         ND         ND         ND         ND         LR           4,4'-Methylenebis (N,N-dimethyl-aniline)         101-61-1         X         X         ND         X         X         MD         ND         ND         ND         X         X         MD         ND         ND         ND         X         X         MD         ND         ND         X         X         MD         ND         ND         X         X         ND         ND         ND         X         X         MD         ND         ND         X         X         MD         ND         ND         X         X         X         ND         ND         ND         ND         X         X         X         ND         ND         ND         ND         ND         X         X         X         ND         ND         ND         ND
4,4'-Methylenebis (2-chloroaniline)       101-14-4       OE, OS       ND       ND       ND       LR         4,4'-Methylenebis(N,N-dimethyl-aniline)       101-61-1       X       X       ND       ND       ND       ND         Methyl methanesulfonate       66-27-3       X       ND       ND       ND       X         2-Methylnaphthalene       91-57-6       X       X       ND       ND       ND       X         Methyl parathion       298-00-0       X       ND       ND       ND       ND       X         2-Methylphenol       95-48-7       X       ND       ND       ND       ND       X         3-Methylphenol       108-39-4       X       ND       ND       ND       ND       X         4-Methylphenol       106-44-5       X       ND       ND       ND       X         Mevinphos       7786-34-7       X       ND       ND       ND       X         Mexacarbate       315-18-4       HE, HS       ND       ND       ND       X         Mirex       2385-85-5       X       ND       ND       ND       X         Monocrotophos       6923-22-4       HE       ND       ND
4,4'-Methylenebis(N,N-dimethyl-aniline)       101-61-1       X       X       ND       ND       ND         Methyl methanesulfonate       66-27-3       X       ND       ND       ND       X         2-Methylnaphthalene       91-57-6       X       X       ND       ND       ND       X         Methyl parathion       298-00-0       X       ND       ND       ND       ND       X         2-Methylphenol       95-48-7       X       ND       ND       ND       ND       X         3-Methylphenol       108-39-4       X       ND       ND       ND       X         4-Methylphenol       106-44-5       X       ND       ND       ND       X         Mevinphos       7786-34-7       X       ND       ND       ND       X         Mexacarbate       315-18-4       HE, HS       ND       ND       ND       X         Mirex       2385-85-5       X       ND       ND       ND       X         Monocrotophos       6923-22-4       HE       ND       ND       ND       X         Naled       300-76-5       X       ND       ND       ND       ND       ND       ND
aniline)         Methyl methanesulfonate         66-27-3         X         ND         ND         ND         X           2-Methylnaphthalene         91-57-6         X         X         ND         ND         X           Methyl parathion         298-00-0         X         ND         ND         ND         X           2-Methylphenol         95-48-7         X         ND         ND         ND         X           3-Methylphenol         108-39-4         X         ND         ND         ND         X           4-Methylphenol         106-44-5         X         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X
2-Methylnaphthalene         91-57-6         X         X         ND         X         X           Methyl parathion         298-00-0         X         ND         ND         ND         X           2-Methylphenol         95-48-7         X         ND         ND         ND         X           3-Methylphenol         108-39-4         X         ND         ND         ND         X           4-Methylphenol         106-44-5         X         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X         X         X
Methyl parathion         298-00-0         X         ND         ND         X           2-Methylphenol         95-48-7         X         ND         ND         ND         X           3-Methylphenol         108-39-4         X         ND         ND         ND         X           4-Methylphenol         106-44-5         X         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X         X
2-Methylphenol         95-48-7         X         ND         ND         ND         X           3-Methylphenol         108-39-4         X         ND         ND         ND         X           4-Methylphenol         106-44-5         X         ND         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
3-Methylphenol         108-39-4         X         ND         ND         X           4-Methylphenol         106-44-5         X         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
4-Methylphenol         106-44-5         X         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
4-Methylphenol         106-44-5         X         ND         ND         ND         X           Mevinphos         7786-34-7         X         ND         ND         ND         ND         X           Mexacarbate         315-18-4         HE, HS         ND         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
Mexacarbate         315-18-4         HE, HS         ND         ND         ND         X           Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
Mirex         2385-85-5         X         ND         ND         ND         X           Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
Monocrotophos         6923-22-4         HE         ND         ND         ND         X           Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
Naled         300-76-5         X         ND         ND         ND         X           Naphthalene         91-20-3         X         X         X         X         X
1-Naphthylamine 134-32-7 OS ND ND ND X
2-Naphthylamine 91-59-8 X ND ND ND X
Nicotine 54-11-5 DC ND ND ND X
5-Nitroacenaphthene 602-87-9 X ND ND ND X
2-Nitroaniline 88-74-4 X X ND X X
3-Nitroaniline 99-09-2 X X ND X X
4-Nitroaniline 100-01-6 X X ND X X
5-Nitro- <i>o</i> -anisidine 99-59-2 X ND ND ND X
Nitrobenzene 98-95-3 X X X X
4-Nitrobiphenyl 92-93-3 X ND ND ND X
Nitrofen 1836-75-5 X ND ND ND X
2-Nitrophenol 88-75-5 X X X X
4-Nitrophenol 100-02-7 X X X X X
5-Nitro-o-toluidine 99-55-8 X X ND ND X
Nitroquinoline-1-oxide 56-57-5 X ND ND ND X
<i>N</i> -Nitrosodi- <i>n</i> -butylamine 924-16-3 X ND ND ND X
N-Nitrosodiethylamine 55-18-5 X ND ND ND X
N-Nitrosodimethylamine 62-75-9 X X X X X
N-Nitrosodiphenylamine 86-30-6 X X X X X
<i>N</i> -Nitrosodi- <i>n</i> -propylamine 621-64-7 X X X X X

	Appropriate Preparation Techniques <sup>b</sup>			es <sup>b</sup>		
				3540/		
Compounds	CAS No <sup>a</sup>	3510	3520	3541	3550	3580
N-Nitrosomethylethylamine	10595-95-6	Х	ND	ND	ND	Χ
N-Nitrosomorpholine	59-89-2	ND	ND	ND	ND	Χ
<i>N</i> -Nitrosopiperidine	100-75-4	X	ND	ND	ND	Χ
<i>N</i> -Nitrosopyrrolidine	930-55-2	X	ND	ND	ND	Χ
Octamethyl pyrophosphoramide	152-16-9	LR	ND	ND	ND	LR
4,4'-Oxydianiline	101-80-4	X	ND	ND	ND	Χ
Parathion	56-38-2	X	Χ	ND	ND	Χ
Pentachlorobenzene	608-93-5	X	ND	ND	ND	Χ
Pentachloronitrobenzene	82-68-8	X	ND	ND	ND	Χ
Pentachlorophenol	87-86-5	X	X	X	Χ	Χ
Phenacetin	62-44-2	X	ND	ND	ND	Χ
Phenanthrene	85-01-8	Χ	X	Χ	X	Χ
Phenobarbital	50-06-6	Χ	ND	ND	ND	X
Phenol	108-95-2	DC	Х	X	X	X
1,4-Phenylenediamine	106-50-3	X	ND	ND	ND	X
Phorate	298-02-2	Χ	ND	ND	ND	X
Phosalone	2310-17-0	HS	ND	ND	ND	X
Phosmet	732-11-6	HS	ND	ND	ND	X
Phosphamidon	13171-21-6	HE	ND	ND	ND	X
Phthalic anhydride	85-44-9	CP, HE	ND	ND	ND .	CP
2-Picoline (2-Methylpyridine)	109-06-8	X	X	ND	ND	ND
Piperonyl sulfoxide	120-62-7	X	ND	ND	ND	X
Pronamide	23950-58-5	X	ND	ND	ND	X
Propylthiouracil	51-52-5	C LR T	ND	ND	ND	LR
Pyrene	129-00-0	$\Pi_{X}^{C_{X}}$	$n_{\mathbf{x}}^{\mathbf{x}}$		OX	
Resorcinol	108-46-3	DC, OE	ND	ND	ND	X
Safrole	94-59-7	X	ND	ND	ND	X
Strychnine	57-24-9	AW, OS	ND	ND	ND	X
Sulfallate	95-06-7	X	ND	ND	ND	X
Terbufos	13071-79-9	X	ND	ND	ND	X
1,2,4,5-Tetrachlorobenzene	95-94-3	X	ND	ND	ND	X
2,3,4,6-Tetrachlorophenol	58-90-2	X	ND	ND	ND	X
Tetrachlorvinphos	961-11-5	X	ND	ND	ND	X
Tetraethyl dithiopyrophosphate	3689-24-5	X	Х	ND	ND	ND
Tetraethyl pyrophosphate	107-49-3	Χ	ND	ND	ND	X
Thionazine	297-97-2	Χ	ND	ND	ND	X
Thiophenol (Benzenethiol)	108-98-5	Χ	ND	ND	ND	X
Toluene diisocyanate	584-84-9	HE	ND	ND	ND	X
o-Toluidine	95-53-4	Χ	ND	ND	ND	X
Toxaphene	8001-35-2	Χ	Χ	Χ	Χ	Χ
1,2,4-Trichlorobenzene	120-82-1	Χ	Χ	Χ	X	X
2,4,5-Trichlorophenol	95-95-4	Χ	Χ	ND	Χ	X
2,4,6-Trichlorophenol	88-06-2	X	X	X	Χ	Χ

		Approp	riate Prep	aration T	echnique	es <sup>b</sup>
Compounds	CAS No <sup>a</sup>	3510	3520	3540/ 3541	3550	3580
Trifluralin	1582-09-8	Χ	ND	ND	ND	Х
2,4,5-Trimethylaniline	137-17-7	Χ	ND	ND	ND	Χ
Trimethyl phosphate	512-56-1	HE	ND	ND	ND	X
1,3,5-Trinitrobenzene	99-35-4	Χ	ND	ND	ND	Χ
Tris(2,3-dibromopropyl) phosphate	126-72-7	Χ	ND	ND	ND	LR
Tri-p-tolyl phosphate	78-32-0	Χ	ND	ND	ND	X
O, O, O-Triethyl phosphorothioate	126-68-1	X	ND	ND	ND	Χ

<sup>&</sup>lt;sup>a</sup> Chemical Abstract Service Registry Number

## **KEY TO ANALYTE LIST**

- AW = Adsorption to walls of glassware during extraction and storage.
- CP = Nonreproducible chromatographic performance.
- DC = Unfavorable distribution coefficient.
- HE = Hydrolysis during extraction accelerated by acidic or basic conditions.
- HS = Hydrolysis during storage potential.
- LR = Low response.
- ND = Not determined.
- OE = Oxidation during extraction accelerated by basic conditions.
- OS = Oxidation during storage potential.
- X = Historically, adequate recovery can be obtained by this technique. However, actual recoveries may vary depending on the extraction efficiency, the number of constituents being analyzed concurrently, and the analytical instrumentation.
- 1.2 In addition to the sample preparation methods listed in the above analyte list, Method 3535 describes a solid-phase extraction procedure that may be applied to the extraction of semivolatiles from TCLP leachates (see Tables 16 and 17 of this method for performance data). Method 3542 describes sample preparation for semivolatile organic compounds in air sampled by Method 0010 (see Table 11 of this method for surrogate performance data), Method 3545 describes an automated solvent extraction device for semivolatiles in solids (see Table 12 of this method for performance data), Method 3561 describes a supercritical fluid device for the extraction of PAHs from solids (see Tables 13, 14, and 15 of this method for performance data), and Method 3546 provides an extraction procedure employing commercially available microwave equipment to extract semivolatiles while using less solvent and taking less time than procedures such as a Soxhlet extraction (see Tables 19 through 23 of this method for the applicable performance data). (The tabulated data are provided for guidance purposes only.)
- 1.3 This method can be used to quantitate most neutral, acidic, and basic organic compounds that are soluble in methylene chloride (or other suitable solvents provided that the desired performance data can be generated) and are capable of being eluted, without derivatization, as sharp peaks from a gas chromatographic fused-silica capillary column coated with a slightly polar silicone. Such compounds include polynuclear aromatic hydrocarbons, chlorinated hydrocarbons and pesticides, phthalate esters, organophosphate esters, nitrosamines, haloethers, aldehydes, ethers, ketones, anilines, pyridines, quinolines, aromatic

<sup>&</sup>lt;sup>b</sup> See Sec. 1.2 for other acceptable preparation methods.

nitro compounds, and phenols, including nitrophenols. See Table 1 for a list of compounds and their characteristic ions that have been evaluated.

In most cases, this method is not appropriate for the quantitation of multicomponent analytes, e.g., Aroclors, Toxaphene, Chlordane, etc., because of limited sensitivity for those analytes. When these analytes have been identified by another technique, Method 8270 may be appropriate for confirmation of the identification of these analytes when concentration in the extract permits. Refer to Methods 8081 and 8082 for guidance on calibration and quantitation of multicomponent analytes such as the Aroclors, Toxaphene, and Chlordane.

- 1.4 The following compounds may require special treatment when being determined by this method:
  - 1.4.1 Benzidine may be subject to oxidative losses during solvent concentration and its chromatographic behavior is poor.
  - 1.4.2 Under the alkaline conditions of the extraction step from aqueous matrices,  $\alpha$ -BHC,  $\gamma$ -BHC, Endosulfan I and II, and Endrin are subject to decomposition. Neutral extraction should be performed if these compounds are expected to be present.
  - 1.4.3 Hexachlorocyclopentadiene is subject to thermal decomposition in the inlet of the gas chromatograph, chemical reaction in acetone solution, and photochemical decomposition.
  - 1.4.4 N-nitrosodimethylamine is difficult to separate from the solvent under the chromatographic conditions described.
  - 1.4.5 N-nitrosodiphenylamine decomposes in the gas chromatographic inlet and cannot be separated from diphenylamine. For this reason, it is acceptable to report the combined result for n-nitrosodiphenylamine and diphenylamine for either of these compounds as a combined concentration.
  - 1.4.6 1,2-Diphenylhydrazine is unstable even at room temperature and readily converts to azobenzene. Given the stability problems, it would be acceptable to calibrate for 1,2-diphenylhydrazine using azobenzene. Under these poor compound separation circumstances the results for either of these compounds should be reported as a combined concentration.
  - 1.4.7 Pentachlorophenol, 2,4-dinitrophenol, 4-nitrophenol, benzoic acid, 4,6-dinitro-2-methylphenol, 4-chloro-3-methylphenol, 2-nitroaniline, 3-nitroaniline, 4-nitroaniline, and benzyl alcohol are subject to erratic chromatographic behavior, especially if the GC system is contaminated with high boiling material.
  - 1.4.8 Pyridine may perform poorly at the GC injection port temperatures listed in this method. Lowering the injection port temperature may reduce the amount of degradation. However, the analyst must use caution in modifying the injection port temperature, as the performance of other analytes may be adversely affected. Therefore, if pyridine is to be determined in addition to other target analytes, it may be necessary to perform separate analyses. In addition, pyridine may be lost during the evaporative concentration of the sample extract. As a result, many of the extraction methods listed above may yield low recoveries unless great care is exercised during the concentration steps. For this reason, analysts may wish to consider the use of extraction techniques such as pressurized fluid extraction (Method 3545), microwave extraction (Method 3546),

or supercritical fluid extraction, which involve smaller extract volumes, thereby reducing or eliminating the need for evaporative concentration techniques for many applications.

- 1.4.9 Toluene diisocyanate rapidly hydrolyzes in water (half-life of less than 30 min). Therefore, recoveries of this compound from aqueous matrices should not be expected. In addition, in solid matrices, toluene diisocyanate often reacts with alcohols and amines to produce urethane and ureas and consequently cannot usually coexist in a solution containing these materials.
- 1.4.10 In addition, analytes in the list provided above are flagged when there are limitations caused by sample preparation and/or chromatographic problems.
- 1.5 The lower limits of quantitation for this method when determining an individual compound are approximately 660  $\mu$ g/kg (wet weight) for soil/sediment samples, 1-200 mg/kg for wastes (dependent on matrix and method of preparation), and 10  $\mu$ g/L for ground water samples (see Table 2). Lower limits of quantitation will be proportionately higher for sample extracts that require dilution to avoid saturation of the detector. The lower limits of quantitation listed in Table 2 are provided for guidance and may not always be achievable.
- 1.6 Prior to employing this method, analysts are advised to consult the base method for each type of procedure that may be employed in the overall analysis (e.g., Methods 3500, 3600, 5000, and 8000) for additional information on quality control procedures, development of QC acceptance criteria, calculations, and general guidance. Analysts also should consult the disclaimer statement at the front of the manual and the information in Chapter Two for guidance on the intended flexibility in the choice of methods, apparatus, materials, reagents, and supplies, and on the responsibilities of the analyst for demonstrating that the techniques employed are appropriate for the analytes of interest, in the matrix of interest, and at the levels of concern.

In addition, analysts and data users are advised that, except where explicitly specified in a regulation, the use of SW-846 methods is *not* mandatory in response to Federal testing requirements. The information contained in this method is provided by EPA as guidance to be used by the analyst and the regulated community in making judgments necessary to generate results that meet the data quality objectives for the intended application.

1.7 Use of this method is restricted to use by, or under supervision of, personnel appropriately experienced and trained in the use of gas chromatograph/mass spectrometers and skilled in the interpretation of mass spectra. Each analyst must demonstrate the ability to generate acceptable results with this method.

## 2.0 SUMMARY OF METHOD

- 2.1 The samples are prepared for analysis by gas chromatography/mass spectrometry (GC/MS) using the appropriate sample preparation (refer to Method 3500) and, if necessary, sample cleanup procedures (refer to Method 3600).
- 2.2 The semivolatile compounds are introduced into the GC/MS by injecting the sample extract into a gas chromatograph (GC) equipped with a narrow-bore fused-silica capillary column. The GC column is temperature-programmed to separate the analytes, which are then detected with a mass spectrometer (MS) connected to the gas chromatograph.
- 2.3 Analytes eluted from the capillary column are introduced into the mass spectrometer via a jet separator or a direct connection. Identification of target analytes is

accomplished by comparing their mass spectra with the electron impact (or electron impact-like) spectra of authentic standards. Quantitation is accomplished by comparing the response of a major (quantitation) ion relative to an internal standard using an appropriate calibration curve for the intended application.

2.4 This method includes specific calibration and quality control steps that supersede the general recommendations provided in Method 8000.

#### 3.0 DEFINITIONS

Refer to Chapter One and the manufacturer's instructions for definitions that may be relevant to this procedure.

# 4.0 INTERFERENCES

- 4.1 Solvents, reagents, glassware, and other sample processing hardware may yield artifacts and/or interferences to sample analysis. All of these materials must be demonstrated to be free from interferences under the conditions of the analysis by analyzing method blanks. Specific selection of reagents and purification of solvents by distillation in all-glass systems may be necessary. Refer to each method to be used for specific guidance on quality control procedures and to Chapter Four for general guidance on the cleaning of glassware. Also refer to Method 8000 for a discussion of interferences.
- 4.2 Raw GC/MS data from all blanks, samples, and spikes must be evaluated for interferences. Determine if the source of interference is in the preparation and/or cleanup of the samples and take corrective action to eliminate the problem.
- 4.3 Contamination by carryover can occur whenever high-concentration and low-concentration samples are sequentially analyzed. To reduce carryover, the sample syringe must be rinsed with solvent between sample injections. Whenever an unusually concentrated sample is encountered, it should be followed by the analysis of solvent to check for cross-contamination.

### 5.0 SAFETY

This method does not address all safety issues associated with its use. The laboratory is responsible for maintaining a safe work environment and a current awareness file of OSHA regulations regarding the safe handling of the chemicals listed in this method. A reference file of material safety data sheets (MSDSs) should be available to all personnel involved in these analyses.

## 6.0 EQUIPMENT AND SUPPLIES

The mention of trade names or commercial products in this manual is for illustrative purposes only, and does not constitute an EPA endorsement or exclusive recommendation for use. The products and instrument settings cited in SW-846 methods represent those products and settings used during method development or subsequently evaluated by the Agency. Glassware, reagents, supplies, equipment, and settings other than those listed in this manual may be employed provided that method performance appropriate for the intended application has been demonstrated and documented.

This section does not list common laboratory glassware (e.g., beakers and flasks).

- 6.1 Gas chromatograph/mass spectrometer system
- 6.1.1 Gas chromatograph -- An analytical system equipped with a temperature-programmable gas chromatograph suitable for splitless injection and all required accessories, including syringes, analytical columns, and gases. The capillary column should be directly coupled to the source.
- 6.1.2 Column -- 30-m x 0.25-mm ID (or 0.32-mm ID) 0.25, 0.5, or 1-µm film thickness silicone-coated fused-silica capillary column (J&W Scientific DB-5 or equivalent). The columns listed in this section were the columns used in developing the method. The listing of these columns in this method is not intended to exclude the use of other columns that may be developed. Laboratories may use these columns or other capillary columns provided that the laboratories document method performance data (e.g., chromatographic resolution, analyte breakdown, and sensitivity) that are appropriate for the intended application.

# 6.1.3 Mass spectrometer

- 6.1.3.1 Capable of scanning from 35 to 500 amu every 1 sec or less, using 70 volts (nominal) electron energy in the electron impact ionization mode. The mass spectrometer must be capable of producing a mass spectrum for decafluorotriphenylphosphine (DFTPP) which meets the criteria as outlined in Sec. 11.3.1.
- 6.1.3.2 An ion trap mass spectrometer may be used if it is capable of axial modulation to reduce ion-molecule reactions and can produce electron impact-like spectra that match those in the EPA/NIST Library. The mass spectrometer must be capable of producing a mass spectrum for DFTPP which meets the criteria as outlined in Sec. 11.3.1
- 6.1.4 GC/MS interface -- Any GC-to-MS interface may be used that gives acceptable calibration points for each compound of interest and achieves acceptable tuning performance criteria. For a narrow-bore capillary column, the interface is usually capillary-direct into the mass spectrometer source.
- 6.1.5 Data system -- A computer system should be interfaced to the mass spectrometer. The system must allow the continuous acquisition and storage on machine-readable media of all mass spectra obtained throughout the duration of the chromatographic program. The computer should have software that can search any GC/MS data file for ions of a specific mass and that can plot such ion abundances versus time or scan number. This type of plot is defined as an Extracted Ion Current Profile (EICP). Software should also be available that allows integrating the abundances in any EICP between specified time or scan-number limits. The most recent version of the EPA/NIST Mass Spectral Library should also be available.
- 6.1.6 Guard column (optional) -- (J&W deactivated fused-silica, 0.25-mm ID x 6-m, or equivalent) between the injection port and the analytical column joined with column connectors (Agilent Catalog No. 5062-3556, or equivalent).
- 6.2 Syringe -- 10-μL.

- 6.3 Volumetric flasks, Class A -- Appropriate sizes equipped with ground-glass stoppers.
  - 6.4 Balance -- Analytical, capable of weighing 0.0001 g.
- 6.5 Bottles -- Glass equipped with polytetrafluoroethylene (PTFE)-lined screw caps or crimp tops.

#### 7.0 REAGENTS AND STANDARDS

- 7.1 Reagent-grade chemicals must be used in all tests. Unless otherwise indicated, it is intended that all reagents conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society, where such specifications are available. Other grades may be used, provided it is first ascertained that the reagent is of sufficiently high purity to permit its use without lessening the accuracy of the determination. Reagents should be stored in glass to prevent the leaching of contaminants from plastic containers.
- 7.2 Organic-free reagent water -- All references to water in this method refer to organic-free reagent water.

### 7.3 Standard solutions

The following sections describe the preparation of stock, intermediate, and working standards for the compounds of interest. This discussion is provided as an example, and other approaches and concentrations of the target compounds may be used, as appropriate for the intended application. See Method 8000 for additional information on the preparation of calibration standards.

- 7.4 Stock standard solutions (1000 mg/L) -- Standard solutions can be prepared from pure standard materials or purchased as certified solutions.
  - 7.4.1 Prepare stock standard solutions by accurately weighing about 0.0100 g of pure material. Dissolve the material in pesticide quality acetone or other suitable solvent and dilute to volume in a 10-mL volumetric flask. Larger volumes can be used at the convenience of the analyst. When compound purity is assayed to be 96% or greater, the weight may be used without correction to calculate the concentration of the stock standard. Commercially-prepared stock standards may be used at any concentration if they are certified by the manufacturer or by an independent source.
  - 7.4.2 Transfer the stock standard solutions into bottles equipped with PTFE-lined screw-caps. Store, protected from light, at #6 EC or as recommended by the standard manufacturer. Stock standard solutions should be checked frequently for signs of degradation or evaporation, especially just prior to preparing calibration standards from them.
  - 7.4.3 Stock standard solutions must be replaced after 1 year or sooner if comparison with quality control check samples indicates a problem.
  - 7.4.4 It is recommended that nitrosamine compounds be placed together in a separate calibration mix and not combined with other calibration mixes. When using a premixed certified standard, consult the manufacturer's instructions for additional guidance.

- 7.4.5 Mixes with hydrochloride salts may contain hydrochloric acid, which can cause analytical difficulties. When using a premixed certified standard, consult the manufacturer's instructions for additional guidance.
- 7.5 Internal standard solutions -- The internal standards recommended are 1,4-dichlorobenzene- $d_4$ , naphthalene- $d_8$ , acenaphthene- $d_{10}$ , phenanthrene- $d_{10}$ , chrysene- $d_{12}$ , and perylene- $d_{12}$  (see Table 5). Other compounds may be used as internal standards as long as the criteria in Sec. 11.3.2 are met.
  - 7.5.1 Dissolve 0.200 g of each compound with a small volume of carbon disulfide. Transfer to a 50-mL volumetric flask and dilute to volume with methylene chloride so that the final solvent is approximately 20% carbon disulfide. Most of the compounds are also soluble in small volumes of methanol, acetone, or toluene, except for perylene- $d_{12}$ . The resulting solution will contain each standard at a concentration of 4,000 ng/µL. Each 1-mL sample extract undergoing analysis should be spiked with 10 µL of the internal standard solution, resulting in a concentration of 40 ng/µL of each internal standard. Store away from any light source at #6 EC when not in use (-10 EC is recommended). When using premixed certified solutions, store according to the manufacturer's documented holding time and storage temperature recommendations.
  - 7.5.2 If a more sensitive mass spectrometer is employed to achieve lower quantitation levels, a more dilute internal standard solution may be required. Area counts of the internal standard peaks should be between 50-200% of the area of the target analytes in the mid-point calibration analysis.
- 7.6 GC/MS tuning standard -- A methylene chloride solution containing 50 ng/μL of decafluorotriphenylphosphine (DFTPP) should be prepared. The standard should also contain 50 ng/μL each of 4,4'-DDT, pentachlorophenol, and benzidine to verify injection port inertness and GC column performance. Alternate concentrations may be used to compensate for different injection volumes if the total amount injected is 50 ng or less. Store away from any light source at #6 EC when not in use (-10 EC is recommended). If a more sensitive mass spectrometer is employed to achieve lower quantitation levels, a more dilute tuning solution may be necessary. When using premixed certified solutions, store according to the manufacturer's documented holding time and storage temperature recommendations.
- 7.7 Calibration standards -- A minimum of five calibration standards should be prepared at different concentrations. At least one of the calibration standards should correspond to a sample concentration at or below that necessary to meet the data quality objectives of the project. The remaining standards should correspond to the range of concentrations found in actual samples but should not exceed the working range of the GC/MS system. Each standard and/or series of calibration standards prepared at a given concentration should contain all the desired project-specific target analytes for which quantitation and quantitative results are to be reported by this method.
  - 7.7.1 It is the intent of EPA that all target analytes for a particular analysis be included in the calibration standard(s). These target analytes may not include the entire list of analytes (Sec. 1.1) for which the method has been demonstrated. However, the laboratory shall not report a quantitative result for a target analyte that was not included in the calibration standard(s).
  - 7.7.2 Each 1-mL aliquot of calibration standard should be spiked with 10  $\mu$ L of the internal standard solution prior to analysis. All standards should be stored away from any light source at #6 EC when not in use (-10 EC is recommended), and should be freshly prepared once a year, or sooner if check standards indicate a problem. The calibration

verification standard should be prepared, as necessary, and stored at #6 EC. When using premixed certified solutions, store according to the manufacturer's documented holding time and storage temperature recommendations.

7.8 Surrogate standards -- The recommended surrogates are phenol- $d_6$ , 2-fluorophenol, 2,4,6-tribromophenol, nitrobenzene- $d_5$ , 2-fluorobiphenyl, and p-terphenyl- $d_{14}$ . See Method 3500 for instructions on preparing the surrogate solutions.

NOTE: In the presence of samples containing residual chlorine, phenol- $d_6$  has been known to react to form chlorinated phenolic compounds that are not detected as the original spiked surrogate. Sample preservation precautions outlined in Chapter Four should be used when residual chlorine is known to be present in order to minimize degradation of deuterated phenols or any other susceptible target analyte.

7.8.1 Surrogate standard check -- Determine what the appropriate concentration should be for the blank extracts after all extraction, cleanup, and concentration steps. Inject this concentration into the GC/MS to determine recovery of surrogate standards. It is recommended that this check be done whenever a new surrogate spiking solution is prepared.

NOTE: Method 3561 (SFE Extraction of PAHs) recommends the use of bromobenzene and p-quaterphenyl to better cover the range of PAHs listed in the method.

- 7.8.2 If a more sensitive mass spectrometer is employed to achieve lower quantitation levels, a more dilute surrogate solution may be necessary.
- 7.9 Matrix spike and laboratory control standards -- See Method 3500 for instructions on preparing the matrix spike standard. The same standard may be used as the laboratory control standard (LCS) and the spiking solution should be the same source as used for the initial calibration standards to restrict the influence of standard accuracy on the determination of recovery through preparation and analysis.
  - 7.9.1 Matrix spike check -- Determine what concentration should be in the blank extracts after all extraction, cleanup, and concentration steps. Inject this concentration into the GC/MS to determine recovery. It is recommended that this check be done whenever a new matrix spiking solution is prepared.
  - 7.9.2 If a more sensitive mass spectrometer is employed to achieve lower quantitation levels, a more dilute matrix and LCS spiking solution may be necessary.
  - 7.9.3 Some projects may require the spiking of the specific compounds of interest, since the spiking compounds listed in Method 3500 would not be representative of the compounds of interest required for the project. When this occurs, the matrix and LCS spiking standards should be prepared in methanol, with each compound present at a concentration appropriate for the project.
- 7.10 Solvents -- Acetone, hexane, methylene chloride, isooctane, carbon disulfide, toluene, and other appropriate solvents. All solvents should be pesticide quality or equivalent. Solvents may be degassed prior to use.
- 8.0 SAMPLE COLLECTION, PRESERVATION, AND STORAGE
  - 8.1 See the introductory material to Chapter Four, "Organic Analytes."

8.2 Store the sample extracts at #6 EC, protected from light, in sealed vials (e.g., screw-cap vials or crimp-capped vials) equipped with unpierced PTFE-lined septa.

## 9.0 QUALITY CONTROL

- 9.1 Refer to Chapter One for guidance on quality assurance (QA) and quality control (QC) protocols. When inconsistencies exist between QC guidelines, method-specific QC criteria take precedence over both technique-specific criteria and those criteria given in Chapter One, and technique-specific QC criteria take precedence over the criteria in Chapter One. Any effort involving the collection of analytical data should include development of a structured and systematic planning document, such as a Quality Assurance Project Plan (QAPP) or a Sampling and Analysis Plan (SAP), which translates project objectives and specifications into directions for those that will implement the project and assess the results. Each laboratory should maintain a formal quality assurance program. The laboratory should also maintain records to document the quality of the data generated. All data sheets and quality control data should be maintained for reference or inspection.
- 9.2 Refer to Method 8000 for specific determinative method QC procedures. Refer to Method 3500 or 5000 for QC procedures to ensure the proper operation of the various sample preparation techniques. If an extract cleanup procedure is performed, refer to Method 3600 for the appropriate QC procedures. Any more specific QC procedures provided in this method will supersede those noted in Methods 8000, 5000, 3500, or 3600.
- 9.3 Quality control procedures necessary to evaluate the GC system operation are found in Method 8000 and include evaluation of retention time windows, calibration verification and chromatographic analysis of samples. In addition, discussions regarding the instrument QC requirements listed below can be found in the referenced sections of this method:
  - The GC/MS must be tuned to meet the recommended DFTPP criteria prior to the initial calibration and for each 12-hr period during which analyses are performed. See Secs. 11.3.1 and 11.4.1 for further details.
  - There must be an initial calibration of the GC/MS system as described in Sec. 11.3. In addition, the initial calibration curve should be verified immediately after performing the standard analyses using a second source standard (prepared using standards different from the calibration standards). The suggested acceptance limits for this initial calibration verification analysis are 70 130%. Alternative acceptance limits may be appropriate based on the desired project-specific data quality objectives. Quantitative sample analyses should not proceed for those analytes that fail the second source standard initial calibration verification. However, analyses may continue for those analytes that fail the criteria with an understanding these results could be used for screening purposes and would be considered estimated values.
  - The GC/MS system must meet the calibration verification acceptance criteria in Sec. 11.4, each 12 hrs.
  - The RRT of the sample component must fall within the RRT window of the standard component provided in Sec. 11.6.1.

## 9.4 Initial demonstration of proficiency

Each laboratory must demonstrate initial proficiency with each sample preparation and determinative method combination it utilizes by generating data of acceptable accuracy and precision for target analytes in a clean matrix. The laboratory must also repeat the following operations whenever new staff members are trained or significant changes in instrumentation are made. See Method 8000 for information on how to accomplish a demonstration of proficiency.

9.5 Initially, before processing any samples, the analyst should demonstrate that all parts of the equipment in contact with the sample and reagents are interference-free. This is accomplished through the analysis of a method blank. As a continuing check, each time samples are extracted, cleaned up, and analyzed, a method blank must be prepared and analyzed for the compounds of interest as a safeguard against chronic laboratory contamination. If a peak is observed within the retention time window of any analyte that would prevent the determination of that analyte, determine the source and eliminate it, if possible, before processing the samples. The blanks should be carried through all stages of sample preparation and analysis. When new reagents or chemicals are received, the lab should monitor the preparation and/or analysis blanks associated with samples for any signs of contamination. It is not necessary to test every new batch of reagents or chemicals prior to sample preparation if the source shows no prior problems. However, if reagents are changed during a preparation batch, separate blanks need to be prepared for each set of reagents.

# 9.6 Sample quality control for preparation and analysis

The laboratory must also have procedures for documenting the effect of the matrix on method performance (precision, accuracy, method sensitivity). At a minimum, this should include the analysis of QC samples including a method blank, a matrix spike, a duplicate, and a laboratory control sample (LCS) in each analytical batch and the addition of surrogates to each field sample and QC sample when surrogates are used. Any method blanks, matrix spike samples, and replicate samples should be subjected to the same analytical procedures (Sec. 11.0) as those used on actual samples.

- 9.6.1 Documenting the effect of the matrix should include the analysis of at least one matrix spike and one duplicate unspiked sample or one matrix spike/matrix spike duplicate pair. The decision on whether to prepare and analyze duplicate samples or a matrix spike/matrix spike duplicate must be based on a knowledge of the samples in the sample batch. If samples are expected to contain target analytes, laboratories may use a matrix spike and a duplicate analysis of an unspiked field sample. If samples are not expected to contain target analytes, then laboratories should use a matrix spike and matrix spike duplicate pair. Consult Method 8000 for information on developing acceptance criteria for the MS/MSD.
- 9.6.2 A laboratory control sample (LCS) should be included with each analytical batch. The LCS consists of an aliquot of a clean (control) matrix similar to the sample matrix and of the same weight or volume. The LCS is spiked with the same analytes at the same concentrations as the matrix spike, when appropriate. When the results of the matrix spike analysis indicate a potential problem due to the sample matrix itself, the LCS results are used to verify that the laboratory can perform the analysis in a clean matrix. Consult Method 8000 for information on developing acceptance criteria for the LCS.
- 9.6.3 Also see Method 8000 for the details on carrying out sample quality control procedures for preparation and analysis. In-house method performance criteria for

evaluating method performance should be developed using the guidance found in Method 8000.

## 9.7 Surrogate recoveries

If surrogates are used, the laboratory should evaluate surrogate recovery data from individual samples versus the surrogate control limits developed by the laboratory. See Method 8000 for information on evaluating surrogate data and developing and updating surrogate limits. Procedures for evaluating the recoveries of multiple surrogates and the associated corrective actions should be defined in an approved project plan.

- 9.8 The experience of the analyst performing GC/MS analyses is invaluable to the success of the methods. Each day that analysis is performed, the calibration verification standard should be evaluated to determine if the chromatographic system is operating properly. Questions that should be asked are: Do the peaks look normal? Is the response obtained comparable to the response from previous calibrations? Careful examination of the standard chromatogram can indicate whether the column is still performing acceptably, the injector is leaking, the injector septum needs replacing, etc. When any changes are made to the system (e.g., the column is changed, a septum is changed), see the guidance in Method 8000 regarding whether recalibration of the system must take place.
- 9.9 It is recommended that the laboratory adopt additional quality assurance practices for use with this method. The specific practices that are most productive depend upon the needs of the laboratory and the nature of the samples. Whenever possible, the laboratory should analyze standard reference materials and participate in relevant performance evaluation studies.

### 10.0 CALIBRATION AND STANDARDIZATION

See Sec 11.3 for information on calibration and standardization.

### 11.0 PROCEDURE

# 11.1 Sample preparation

11.1.1 Samples are normally prepared by one of the following methods prior to GC/MS analysis.

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<u>Matrix</u>	<u>Methods</u>
Air (particulates and sorbent resin) Water (including TCLP leachates) Soil/sediment Waste	3542 3510, 3520, 3535 3540, 3541, 3545, 3546, 3550, 3560, 3561 3540, 3541, 3545, 3546, 3550, 3560, 3561, 3580

11.1.2 In very limited applications, direct injection of the sample into the GC/MS system with a 10- $\mu$ L syringe may be appropriate. The quantitation limit is very high (approximately 10,000  $\mu$ g/L). Therefore, it is only appropriate where concentrations in excess of 10,000  $\mu$ g/L are expected.

11.2 Extract cleanup -- Cleanup procedures may not be necessary for a relatively clean sample matrix, but most extracts from environmental and waste samples will require additional preparation before analysis. The specific cleanup procedure used will depend on the nature of the sample to be analyzed and the data quality objectives for the measurements. General guidance for sample extract cleanup is provided in this section and in Method 3600.

Extracts may be cleaned up by any of the following methods prior to GC/MS analysis.

Analytes of Interest	<u>Methods</u>
Aniline and aniline derivatives Phenols Phthalate esters	3620 3630, 3640, 8041 <sup>a</sup> 3610, 3620, 3640
Nitrosamines	3610, 3620, 3640
Organochlorine pesticides PCBs	3610, 3620, 3630, 3640, 3660 3620, 3630, 3660, 3665
Nitroaromatics and cyclic ketones Polynuclear aromatic hydrocarbons	3620, 3640 3611, 3630, 3640
Haloethers	3620, 3640
Chlorinated hydrocarbons	3620, 3640
Organophosphorus pesticides	3620
Petroleum waste	3611, 3650
All base, neutral, and acid	
priority pollutants	3640

<sup>&</sup>lt;sup>a</sup> Method 8041 includes a derivatization technique and a GC/ECD analysis, if interferences are encountered on GC/FID.

### 11.3 Initial calibration

Establish the GC/MS operating conditions, using the following recommendations as guidance.

Mass range: 35-500 amu Scan time: #1 sec/scan

Initial temperature: 40 EC, hold for 4 min Temperature program: 40-320 EC at 10 EC/min

Final temperature: 320 EC, hold until 2 min after benzo[q,h,i]perylene elutes

Injector temperature: 250-300 EC Transfer line temperature: 250-300 EC

Source temperature: According to manufacturer's specifications

Injector: Grob-type, splitless

Injection volume: 1-2 µL

Carrier gas: Hydrogen at 50 cm/sec or helium at 30 cm/sec

lon trap only: Set axial modulation, manifold temperature, and emission

current to manufacturer's recommendations

Split injection is allowed if the sensitivity of the mass spectrometer is sufficient.

11.3.1 The GC/MS system must be hardware-tuned such that injecting 50 ng or less of DFTPP meets the manufacturer's specified acceptance criteria or as listed in Table 3. The tuning criteria as outlined in Table 3 were developed using quadrupole mass spectrometer instrumentation and it is recognized that other tuning criteria may be more effective depending on the type of instrumentation, e.g., Time-of-Flight, Ion Trap, etc. In

these cases it would be appropriate to follow the manufacturer's tuning instructions or some other consistent tuning criteria. However, no matter which tuning criteria is selected, the system calibration must not begin until the tuning acceptance criteria are met with the sample analyses performed under the same conditions as the calibration standards.

- 11.3.1.1 In the absence of specific recommendations on how to acquire the mass spectrum of DFTPP from the instrument manufacturer, the following approach should be used: Three scans (the peak apex scan and the scans immediately preceding and following the apex) are acquired and averaged. Background subtraction is required, and must be accomplished using a single scan acquired within 20 scans of the elution of DFTPP. The background subtraction should be designed only to eliminate column bleed or instrument background ions. Do not subtract part of the DFTPP peak or any other discrete peak that does not coelute with DFTPP.
- 11.3.1.2 Use the DFTPP mass intensity criteria in the manufacturer's instructions as primary tuning acceptance criteria or those in Table 3 as default tuning acceptance criteria if the primary tuning criteria are not available. Alternatively, other documented tuning criteria may be used (e.g. CLP, or Method 625), provided that method performance is not adversely affected. The analyst is always free to choose criteria that are tighter than those included in this method or to use other documented criteria provided they are used consistently throughout the initial calibration, calibration verification, and sample analyses.

NOTE: All subsequent standards, samples, MS/MSDs, and blanks associated with a DFTPP analysis must use the identical mass spectrometer instrument conditions.

11.3.1.3 The GC/MS tuning standard solution should also be used to assess GC column performance and injection port inertness. Degradation of DDT to DDE and DDD should not exceed 20%. (See Method 8081 for the percent breakdown calculation.) Benzidine and pentachlorophenol should be present at their normal responses, and should not exceed a tailing factor of 2 given by the following equation:

TailingFactor = 
$$\frac{BC}{AB}$$

Where the peak is defined as follows: AC is the width at 10% height; DE is the height of peak and B is the height at 10% of DE. This equation compares the width of the back half of the peak to the width of the front half of the peak at 10% of the height. (See Figure 1 for an example tailing factor calculation.)

11.3.1.4 If degradation is excessive and/or poor chromatography is noted, the injection port may require cleaning. It may also be necessary to break off the first 6 to12 in. of the capillary column. The use of a guard column (Sec. 6.1.6) between the injection port and the analytical column may help prolong analytical column performance life.

- 11.3.2 The internal standards selected in Sec. 7.5 should permit most of the components of interest in a chromatogram to have retention times of 0.80-1.20 relative to one of the internal standards. Use the base peak ion from the specific internal standard as the primary ion for quantitation (see Table 1). If interferences are noted, use the next most intense ion as the quantitation ion (e.g., for 1,4-dichlorobenzene- $d_4$ , use m/z 150 for quantitation).
- 11.3.3 Analyze 1-2  $\mu$ L of each calibration standard (containing the compounds for quantitation and the appropriate surrogates and internal standards) and tabulate the area of the primary ion against concentration for each target analyte (as indicated in Table 1). A set of at least five calibration standards is necessary (see Sec. 7.7 and Method 8000). Alternate injection volumes may be used if the applicable quality control requirements for using this method are met. The injection volume must be the same for all standards and sample extracts. Figure 2 shows a chromatogram of a calibration standard containing base/neutral and acid analytes.

## 11.3.4 Initial calibration calculations

Calculate response factors (RFs) for each target analyte relative to one of the internal standards (see Table 5) as follows:

 $RF \cdot \frac{A_s \times C_{is}}{A_{is} \times C_s}$ 

where:

A<sub>s</sub> = Peak area (or height) of the analyte or surrogate.

 $A_{is}$  = Peak area (or height) of the internal standard.

C<sub>s</sub> = Concentration of the analyte or surrogate, in μg/L.

 $C_{is}$  = Concentration of the internal standard, in  $\mu g/L$ .

11.3.4.1 Calculate the mean response factor and the relative standard deviation (RSD) of the response factors for each target analyte using the following equations. The RSD should be less than or equal to 20% for each target analyte. It is also recommended that a minimum response factor for the most common target analytes, as noted in Table 4, be demonstrated for each individual calibration level as a means to ensure that these compounds are behaving as expected. In addition, meeting the minimum response factor criteria for the lowest calibration standard is critical in establishing and demonstrating the desired sensitivity. Due to the large number of compounds that may be analyzed by this method, some compounds will fail to meet this criteria. For these occasions, it is acknowledged that the failing compounds may not be critical to the specific project and therefore they may be used as qualified data or estimated values for screening purposes. The analyst should also strive to place more emphasis on meeting the calibration criteria for those compounds that are critical project compounds, rather than meeting the criteria for those less important compounds.

mean RF ' 
$$\frac{1}{RF}$$
 '  $\frac{1}{n}$  RF<sub>i</sub>

SD ' 
$$\sqrt{\frac{\int_{1/1}^{n} (RF_i \& \overline{RF})^2}{n \& 1}}$$

RSD ' 
$$\frac{SD}{RE}$$
 × 100

where:

RF<sub>i</sub> = RF for each of the calibration standards

**RR** = mean RF for each compound from the initial calibration

n = Number of calibration standards, e.g., 5

11.3.4.2 If more than 10% of the compounds included with the initial calibration exceed the 20% RSD limit and do not meet the minimum correlation coefficient (0.99) for alternate curve fits, then the chromatographic system is considered too reactive for analysis to begin. Clean or replace the injector liner and/or capillary column, then repeat the calibration procedure beginning with Sec. 11.3.

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11.3.5 Evaluation of retention times -- The relative retention time (RRT) of each target analyte in each calibration standard should agree within 0.06 RRT units. Late-eluting target analytes usually have much better agreement.

RRT 
$$\frac{}{}$$
 Retention time of the analyte Retention time of the internal standard

- 11.3.6 Linearity of target analytes -- If the RSD of any target analyte is 20% or less, then the relative response factor is assumed to be constant over the calibration range, and the average relative response factor may be used for quantitation (Sec. 11.7.2).
  - 11.3.6.1 If the RSD of any target analyte is greater than 20%, refer to Method 8000 for additional calibration options. One of the options must be applied to GC/MS calibration in this situation, or a new initial calibration must be performed. The average RF should not be used for compounds that have an RSD greater than 20% unless the concentration is reported as estimated.
  - 11.3.6.2 When the RSD exceeds 20%, the plotting and visual inspection of a calibration curve can be a useful diagnostic tool. The inspection may indicate analytical problems, including errors in standard preparation, the presence of active sites in the chromatographic system, analytes that exhibit poor chromatographic behavior, etc.

- 11.3.6.3 Due to the large number of compounds that may be analyzed by this method, some compounds may fail to meet either the 20% RSD, minimum correlation coefficient criteria (0.99), or the acceptance criteria for alternative calibration procedures in Method 8000. Any calibration method described in Method 8000 may be used, but it should be used consistently. It is considered inappropriate once the calibration analyses are completed to select an alternative calibration procedure in order to pass the recommended criteria on a case-by-case basis. If compounds fail to meet these criteria, the associated concentrations may still be determined but they must be reported as estimated. In order to report non-detects, it must be demonstrated that there is adequate sensitivity to detect the failed compounds at the applicable lower quantitation limit.
- 11.4 GC/MS calibration verification -- Calibration verification consists of three steps that are performed at the beginning of each 12-hr analytical shift.
  - 11.4.1 Prior to the analysis of samples or calibration standards, inject 50 ng or less of the DFTPP standard into the GC/MS system. The resultant mass spectrum for DFTPP must meet the criteria as outlined in Sec. 11.3.1 before sample analysis begins. These criteria must be demonstrated each 12-hr shift during which samples are analyzed.
  - 11.4.2 The initial calibration function for each target analyte should be checked immediately after the first occurrence in the region of the middle of the calibration range with a standard from a source different from that used for the initial calibration. The value determined from the second source check should be within 30% of the expected concentration. An alternative recovery limit may be appropriate based on the desired project-specific data quality objectives. Quantitative sample analyses should not proceed for those analytes that fail the second source standard initial calibration verification. However, analyses may continue for those analytes that fail the criteria with an understanding these results could be used for screening purposes and would be considered estimated values.
  - 11.4.3 The initial calibration (Sec. 11.3) for each compound of interest should be verified once every 12 hrs prior to sample analysis, using the introduction technique and conditions used for samples. This is accomplished by analyzing a calibration standard (containing all the compounds for quantitation) at a concentration either near the midpoint concentration for the calibrating range of the GC/MS or near the action level for the project. The results must be compared against the most recent initial calibration curve and should meet the verification acceptance criteria provided in Secs. 11.4.5 through 11.4.7.
  - NOTE: The DFTPP and calibration verification standard may be combined into a single standard as long as both tuning and calibration verification acceptance criteria for the project can be met without interferences.
- 11.4.4 A method blank should be analyzed prior to sample analyses in order to ensure that the total system (introduction device, transfer lines and GC/MS system) is free of contaminants. If the method blank indicates contamination, then it may be appropriate to analyze a solvent blank to demonstrate that the contamination is not a result of carryover from standards or samples. See Method 8000 for information regarding method blank performance criteria.

- 11.4.5.1 Each of the most common target analytes in the calibration verification standard should meet the minimum response factors as noted in Table 4. This criteria is particularly important when the common target analytes are also critical project-required compounds. This is the same check that is applied during the initial calibration.
- 11.4.5.2 If the minimum response factors are not met, the system should be evaluated, and corrective action should be taken before sample analysis begins. Possible problems include standard mixture degradation, injection port inlet contamination, contamination at the front end of the analytical column, and active sites in the column or chromatographic system.
- 11.4.5.3 All target compounds of interest must be evaluated using a 20% criterion. Use percent difference when performing the average response factor model calibration. Use percent drift when calibrating using a regression fit model. Refer to Method 8000 for guidance on calculating percent difference and drift.
- 11.4.5.4 If the percent difference or percent drift for a compound is less than or equal to 20%, then the initial calibration for that compound is assumed to be valid. Due to the large numbers of compounds that may be analyzed by this method, it is expected that some compounds will fail to meet the criterion. If the criterion is not met (i.e., greater than 20% difference or drift) for more than 20% of the compounds included in the initial calibration, then corrective action must be taken prior to the analysis of samples. In cases where compounds fail, they may still be reported as non-detects if it can be demonstrated that there was adequate sensitivity to detect the compound at the applicable quantitation limit. For situations when the failed compound is present, the concentrations must be reported as estimated values.
- 11.4.5.5 Problems similar to those listed under initial calibration could affect the ability to pass the calibration verification standard analysis. If the problem cannot be corrected by other measures, a new initial calibration must be generated. The calibration verification criteria must be met before sample analysis begins.
- 11.4.5.6 The method of linear regression analysis has the potential for a significant bias to the lower portion of a calibration curve, while the relative percent difference and quadratic methods of calibration do not have this potential bias. When calculating the calibration curves using the linear regression model, a minimum quantitation check on the viability of the lowest calibration point should be performed by re-fitting the response from the low concentration calibration standard back into the curve (see Method 8000 for additional details). It is not necessary to re-analyze a low concentration standard, rather the data system can recalculate the concentrations as if it were an unknown sample. The recalculated concentration of the low calibration point should be within ± 30% of the standard's true concentration. Other recovery criteria may be applicable depending on the project's data quality objectives and for those situations the minimum quantitation check criteria should be outlined in a laboratory standard operating procedure, or a project-specific Quality Assurance Project Plan. Analytes which do not meet the minimum quantitation calibration re-fitting criteria should be considered "out of control" and corrective action such as redefining the lower limit of quantitation

and/or reporting those "out of control" target analytes as estimated when the concentration is at or near the lowest calibration point may be appropriate.

- 11.4.6 Internal standard retention time -- The retention times of the internal standards in the calibration verification standard must be evaluated immediately after or during data acquisition. If the retention time for any internal standard changes by more than 30 sec from that in the mid-point standard level of the most recent initial calibration sequence, then the chromatographic system must be inspected for malfunctions and corrections must be made, as required. When corrections are made, reanalysis of samples analyzed while the system was malfunctioning is required.
- 11.4.7 Internal standard response -- If the EICP area for any of the internal standards in the calibration verification standard changes by a factor of two (-50% to +100%) from that in the mid-point standard level of the most recent initial calibration sequence, the mass spectrometer must be inspected for malfunctions and corrections must be made, as appropriate. When corrections are made, reanalysis of samples analyzed while the system was malfunctioning is required.

# 11.5 GC/MS analysis of samples

- 11.5.1 It is highly recommended that sample extracts be screened on a GC/FID or GC/PID using the same type of capillary column used in the GC/MS system. This will minimize contamination of the GC/MS system from unexpectedly high concentrations of organic compounds.
- 11.5.2 Allow the sample extract to warm to room temperature. Just prior to analysis, add 10  $\mu$ L of the internal standard solution to the 1 mL of concentrated sample extract obtained from sample preparation.
- 11.5.3 Inject an aliquot of the sample extract into the GC/MS system, using the same operating conditions that were used for the calibration (Sec. 11.3). The volume to be injected should include an appropriate concentration that is within the calibration range of base/neutral and acid surrogates using the surrogate solution as noted in Sec. 7.8. The injection volume must be the same volume that was used for the calibration standards.
- 11.5.4 If the response for any quantitation ion exceeds the initial calibration range of the GC/MS system, the sample extract must be diluted and reanalyzed. Additional internal standard solution must be added to the diluted extract to maintain the same concentration as in the calibration standards (usually 40 ng/ $\mu$ L, or other concentrations as appropriate, if a more sensitive GC/MS system is being used). Secondary ion quantitation should be used only when there are sample interferences with the primary ion.
- NOTE: It may be a useful diagnostic tool to monitor internal standard retention times in all samples, spikes, blanks, and standards to effectively check drifting, method performance, poor injection execution, and anticipate the need for system inspection and/or maintenance. Internal standard responses (area counts) must be monitored in all samples, spikes, blanks for similar reasons. If the EICP area for any of the internal standards in samples, spikes and blanks changes by a factor of two (-50% to +100%) from the areas determined in the continuing calibration analyzed that day, corrective action must be taken. The samples, spikes or blanks should be reanalyzed or the data should be qualified.

- 11.5.4.1 When ions from a compound in the sample saturate the detector, this analysis should be followed by the analysis of an instrument blank consisting of clean solvent. If the blank analysis is not free of interferences, then the system must be decontaminated. Sample analysis may not resume until the blank analysis is demonstrated to be free of interferences. Contamination from one sample to the next on the instrument usually takes place in the syringe. If adequate syringe washes are employed, then carryover from high concentration samples can usually be avoided.
- 11.5.4.2 All dilutions should keep the response of the major constituents (previously saturated peaks) in the upper half of the linear range of the curve.
- 11.5.5 The use of selected ion monitoring (SIM) is acceptable for applications requiring quantitation limits below the normal range of electron impact mass spectrometry. However, SIM may provide a lesser degree of confidence in the compound identification, since less mass spectral information is available. Using the primary ion for quantitation and the secondary ions for confirmation set up the collection groups based on their retention times. The selected ions are nominal ions and most compounds have small mass defect, usually less than 0.2 amu, in their spectra. These mass defects should be used in the acquisition table. The dwell time may be automatically calculated by the laboratory's GC/MS software or manually calculated using the following formula. The total scan time should be less than 1,000 msec and produce at least 5 to 10 scans per chromatographic peak. The start and stop times for the SIM groups are determined from the full scan analysis using the formula below:



Additional guidance for performing SIM analyses, in particular for PAHs and phenol target analyte compounds, can be found in the most recent CLP semivolatile organic methods statement of work (SOW). See the SIM sections from the following CLP SOW for further details: EPA CLP Organics SOW. (Reference 14)

# 11.6 Analyte identification

- 11.6.1 The qualitative identification of compounds determined by this method is based on retention time and on comparison of the sample mass spectrum, after background correction, with characteristic ions in a reference mass spectrum. The reference mass spectrum must be generated by the laboratory using the conditions of this method. The characteristic ions from the reference mass spectrum are defined as the three ions of greatest relative intensity, or any ions over 30% relative intensity, if less than three such ions occur in the reference spectrum. Compounds are identified when the following criteria are met.
  - 11.6.1.1 The intensities of the characteristic ions of a compound must maximize in the same scan or within one scan of each other. Selection of a peak by a data system target compound search routine where the search is based on the presence of a target chromatographic peak containing ions specific for the

target compound at a compound-specific retention time will be accepted as meeting this criterion.

- 11.6.1.2 The RRT of the sample component is within  $\pm$  0.06 RRT units of the RRT of the standard component.
- 11.6.1.3 The relative intensities of the characteristic ions agree within 30% of the relative intensities of these ions in the reference spectrum. (Example: For an ion with an abundance of 50% in the reference spectrum, the corresponding abundance in a sample spectrum can range between 20% and 80%.) Use professional judgement in interpretation where interferences are observed.
- 11.6.1.4 Structural isomers that produce very similar mass spectra should be identified as individual isomers if they have sufficiently different GC retention times. Sufficient GC resolution is achieved if the height of the valley between two isomer peaks is less than 50% of the average of the two peak heights. Otherwise, structural isomers are identified as isomeric pairs. The resolution should be verified on the mid-point concentration of the initial calibration as well as the laboratory designated continuing calibration verification level if closely eluting isomers are to be reported (e.g., benzo(b)fluoranthene and benzo(k)fluoranthene).
- 11.6.1.5 Identification is hampered when sample components are not resolved chromatographically and produce mass spectra containing ions contributed by more than one analyte. When gas chromatographic peaks obviously represent more than one sample component (i.e., a broadened peak with shoulder(s) or a valley between two or more maxima), appropriate selection of analyte spectra and background spectra is important.
- 11.6.1.6 Examination of extracted ion current profiles of appropriate ions can aid in the selection of spectra and in qualitative identification of compounds. When analytes coelute (i.e., only one chromatographic peak is apparent), the identification criteria may be met, but each analyte spectrum will contain extraneous ions contributed by the coeluting compound.
- 11.6.2 For samples containing components not associated with the calibration standards, a library search may be made for the purpose of tentative identification. The necessity to perform this type of identification will be determined by the purpose of the analyses being conducted. Data system library search routines should not use normalization routines that would misrepresent the library or unknown spectra when compared to each other.

For example, the RCRA permit or waste delisting requirements may require the reporting of non-target analytes. Only after visual comparison of sample spectra with the nearest library searches may the analyst assign a tentative identification. Guidelines for tentative identification are:

- (1) Relative intensities of major ions in the reference spectrum (ions > 10% of the most abundant ion) should be present in the sample spectrum.
- (2) The relative intensities of the major ions should agree within ± 30%. (Example: For an ion with an abundance of 50% in the standard spectrum, the corresponding sample ion abundance must be between 20 and 80%.)

- (3) Molecular ions present in the reference spectrum should be present in the sample spectrum.
- (4) Ions present in the sample spectrum but not in the reference spectrum should be reviewed for possible background contamination or presence of coeluting compounds.
- (5) Ions present in the reference spectrum but not in the sample spectrum should be reviewed for possible subtraction from the sample spectrum because of background contamination or coeluting peaks. Data system library reduction programs can sometimes create these discrepancies.

### 11.7 Quantitation

- 11.7.1 Once a target compound has been identified, the quantitation of that compound will be based on the integrated abundance of the primary characteristic ion from the EICP.
  - 11.7.1.1 It is highly recommended to use the integration produced by the software if the integration is correct because the software should produce more consistent integrations. However, manual integrations may be necessary when the software does not produce proper integrations because baseline selection is improper; the correct peak is missed; a coelution is integrated; the peak is partially integrated; etc. The analyst is responsible for ensuring that the integration is correct whether performed by the software or done manually.
  - 11.7.1.2 Manual integrations should not be substituted for proper maintenance of the instrument or setup of the method (e.g. retention time updates, integration parameter files, etc). The analyst should seek to minimize manual integration by properly maintaining the instrument, updating retention times, and configuring peak integration parameters.
- 11.7.2 If the RSD of a compound's response factor is 20% or less, then the concentration in the extract may be determined using the average response factor (RR) from initial calibration data (Sec. 11.3.4). See Method 8000 for the equations describing internal standard calibration and either linear or non-linear calibrations.
- 11.7.3 Where applicable, the concentration of any non-target analytes identified in the sample (Sec. 11.6.2) should be estimated. The same formula as in Sec. 11.3.4 should be used with the following modifications: The areas  $A_x$  and  $A_{is}$  should be from the total ion chromatograms, and the RF for the compound should be assumed to be 1.
- 11.7.4 The resulting concentration should be reported indicating that the value is an estimate. Use the nearest internal standard free of interferences.
- 11.7.5 Quantitation of multicomponent compounds (e.g., Toxaphene, Aroclors, etc.) is beyond the scope of Method 8270. Normally, quantitation is performed using a GC/ECD, for example by using Methods 8081 or 8082. However, this method (8270) may be used to confirm the identification of these compounds, when the concentrations are at least 10 ng/µL in the concentrated sample extract.
- 11.7.6 Quantitation of multicomponent parameters such as diesel range organics (DROs) and total petroleum hydrocarbons (TPH) using the Method 8270 recommended internal standard quantitation technique is beyond the scope of this method. Typically.

analyses for these parameters are performed using GC/FID or GC with a MS detector capability that is available with Method 8015.

11.7.7 Structural isomers that produce very similar mass spectra should be quantitated as individual isomers if they have sufficiently different GC retention times. Sufficient GC resolution is achieved if the height of the valley between two isomer peaks is less than 50% of the average of the two peak heights. Otherwise, structural isomers are identified as isomeric pairs. The resolution should be verified on the mid-point concentration of the initial calibration as well as the laboratory designated continuing calibration verification level if closely eluting isomers are to be reported (e.g., benzo(b)fluoranthene and benzo(k)fluoranthene).

### 12.0 DATA ANALYSIS AND CALCULATIONS

See Sec. 11.7 and Method 8000 for information on data analysis and calculations.

## 13.0 METHOD PERFORMANCE

- 13.1 Performance data and related information are provided in SW-846 methods only as examples and guidance. The data do not represent required performance criteria for users of the methods. Instead, performance criteria should be developed on a project-specific basis, and the laboratory should establish in-house QC performance criteria for the application of this method. These performance data are not intended to be and must not be used as absolute QC acceptance criteria for purposes of laboratory accreditation.
- 13.2 Single laboratory initial demonstration of capability data were generated from five replicate measurements using a modified continuous liquid-liquid extractor (Method 3520) with hydrophobic membrane. In this case only a single acid pH extraction was performed using the CLP calibration criteria and the applicable CLP target analytes. These data are presented in Table 6. Laboratories should generate their own acceptance criteria depending on the extraction and instrument conditions. (See Method 8000.)
- 13.3 Chromatograms from calibration standards analyzed with Day 0 and Day 7 samples were compared to detect possible deterioration of GC performance. These recoveries (using Method 3510 extraction) are presented in Table 7. These data are provided for guidance purposes only.
- 13.4 Method performance data using Method 3541 (automated Soxhlet extraction) are presented in Tables 8 and 9. Single laboratory accuracy and precision data were obtained for semivolatile organics in a clay soil by spiking at a concentration of 6 mg/kg for each compound. The spiking solution was mixed into the soil during addition and then allowed to equilibrate for approximately 1 hour prior to extraction. The spiked samples were then extracted by Method 3541 (Automated Soxhlet). Three extractions were performed and each extract was analyzed by gas chromatography/mass spectrometry following Method 8270. The low recovery of the more volatile compounds is probably due to volatilization losses during equilibration. These data as listed were taken from Reference 7 and are provided for guidance purposes only.
- 13.5 Surrogate precision and accuracy data are presented in Table 10 from a field dynamic spiking study based on air sampling by Method 0010. The trapping media were prepared for analysis by Method 3542 and subsequently analyzed by this method (8270). These data are provided for guidance purposes only.

- 13.6 Single laboratory precision and bias data using Method 3545 (pressurized fluid extraction) for semivolatile organic compounds are presented in Table 11. The samples were conditioned spiked samples prepared and certified by a commercial supplier that contained 57 semivolatile organics at three concentrations (250, 2500, and 12,500 µg/kg) on three types of soil (clay, loam and sand). Spiked samples were extracted both by the Dionex Accelerated Solvent Extraction system and by the Perstorp Environmental Soxtec™ (automated Soxhlet). The data in Table 11 represent seven replicate extractions and analyses for each individual sample and were taken from Reference 9. The average recoveries from the three matrices for all analytes and all replicates relative to the automated Soxhlet data are as follows: clay 96.8%, loam 98.7% and sand 102.1%. The average recoveries from the three concentrations also relative to the automated Soxhlet data are as follows: low 101.2%, mid 97.2% and high 99.2%. These data are provided for guidance purposes only.
- 13.7 Single laboratory precision and bias data using Method 3561 (SFE extraction of PAHs with a variable restrictor and solid trapping material) were obtained for the method analytes by the extraction of two certified reference materials (EC-1, a lake sediment from Environment Canada and HS-3, a marine sediment from the National Science and Engineering Research Council of Canada, both naturally-contaminated with PAHs). The SFE instrument used for these extractions was a Hewlett-Packard Model 7680. Analysis was by GC/MS. Average recoveries from six replicate extractions range from 85 to 148% (overall average of 100%) based on the certified value (or a Soxhlet value if a certified value was unavailable for a specific analyte) for the lake sediment. Average recoveries from three replicate extractions range from 73 to 133% (overall average of 92%) based on the certified value for the marine sediment. The data are found in Tables 12 and 13 and were taken from Reference 10. These data are provided for guidance purposes only.
- 13.8 Single laboratory precision and accuracy data using Method 3561 (SFE extraction of PAHs with a fixed restrictor and liquid trapping) were obtained for twelve of the method analytes by the extraction of a certified reference material (a soil naturally contaminated with PAHs). The SFE instrument used for these extractions was a Dionex Model 703-M. Analysis was by GC/MS. Average recoveries from four replicate extractions range from 60 to 122% (overall average of 89%) based on the certified value. The instrument conditions that were utilized to extract a 3.4 g sample were as follows: Pressure -- 300 atm; time -- 60 min.; extraction fluid -- CO<sub>2</sub>; modifier -- 10% 1:1 (v/v) methanol/methylene chloride; Oven temperature -- 80 EC; Restrictor temperature -- 120 EC; and, trapping fluid -- chloroform (methylene chloride has also been used). The data are found in Table 14 and were taken from Reference 11. These data are provided for guidance purposes only.
- 13.9 Tables 15 and 16 contain single-laboratory precision and accuracy data for solidphase extraction of TCLP buffer solutions spiked at two levels and extracted using Method 3535. These data are provided for guidance purposes only.
- 13.10 Table 17 contains multiple-laboratory data for solid-phase extraction of spiked TCLP soil leachates extracted using Method 3535. These data are provided for guidance purposes only.
- 13.11 Tables 18 through 22 contain single-laboratory PAH recovery data for microwave extraction of contaminated soils and standard reference materials using Method 3546. These data are provided for guidance purposes only.

#### 14.0 POLLUTION PREVENTION

- 14.1 Pollution prevention encompasses any technique that reduces or eliminates the quantity and/or toxicity of waste at the point of generation. Numerous opportunities for pollution prevention exist in laboratory operations. The EPA has established a preferred hierarchy of environmental management techniques that places pollution prevention as the management option of first choice. Whenever feasible, laboratory personnel should use pollution prevention techniques to address their waste generation. When wastes cannot be feasibly reduced at the source, the Agency recommends recycling as the next best option.
- 14.2 For information about pollution prevention that may be applicable to laboratories and research institutions consult *Less is Better: Laboratory Chemical Management for Waste Reduction* available from the American Chemical Society's Department of Government Relations and Science Policy, 1155 16th St., N.W. Washington, D.C. 20036, <a href="http://www.acs.org">http://www.acs.org</a>.

### 15.0 WASTE MANAGEMENT

The Environmental Protection Agency requires that laboratory waste management practices be conducted consistent with all applicable rules and regulations. The Agency urges laboratories to protect the air, water, and land by minimizing and controlling all releases from hoods and bench operations, complying with the letter and spirit of any sewer discharge permits and regulations, and by complying with all solid and hazardous waste regulations, particularly the hazardous waste identification rules and land disposal restrictions. For further information on waste management, consult *The Waste Management Manual for Laboratory Personnel* available from the American Chemical Society at the address listed in Sec. 14.2.

## 16.0 REFERENCES

- 1. J. W. Eichelberger, L. E. Harris, and W. L. Budde, W.L., "Reference Compound to Calibrate Ion Abundance Measurement in Gas Chromatography-Mass Spectrometry Systems," Analytical Chemistry, 47, 995-1000, 1975.
- 2. P. Olynyk, W. L. Budde, and J. W. Eichelberger, "Method Detection Limit for Methods 624 and 625," unpublished report, October 1980.
- 3. "Interlaboratory Method Study for EPA Method 625-Base/Neutrals, Acids, and Pesticides," Final Report for EPA Contract 68-03-3102.
- 4. J. A. Burke, "Gas Chromatography for Pesticide Residue Analysis: Some Practical Aspects," Journal of the Association of Official Analytical Chemists (AOAC), 48, 1037, 1965.
- 5. S. V. Lucas, R. A. Kornfeld, "GC-MS Suitability Testing of RCRA Appendix VIII and Michigan List Analytes," U.S. Environmental Protection Agency, Environmental Monitoring and Support Laboratory, Cincinnati, OH 45268, February 20, 1987, Contract No. 68-03-3224.
- T. M. Engel, R. A. Kornfeld, J. S. Warner, and K. D. Andrews, "Screening of Semivolatile Organic Compounds for Extractability and Aqueous Stability by SW-846, Method 3510," U.S. Environmental Protection Agency, Environmental Monitoring and Support Laboratory, Cincinnati, OH 45268, June 5, 1987, Contract 68-03-3224.

- V. Lopez-Avila (W. Beckert, Project Officer); "Development of a Soxtec Extraction Procedure for Extraction of Organic Compounds from Soils and Sediments;" U.S. Environmental Protection Agency, Environmental Monitoring and Support Laboratory, Las Vegas, NV, October 1991; EPA 600/X-91/140.
- 8. J. Bursey, R. Merrill, R. McAllister, and J. McGaughey, "Laboratory Validation of VOST and SemiVOST for Halogenated Hydrocarbons from the Clean Air Act Amendments List," Vol. 1 and 2, U.S. Environmental Protection Agency, EPA 600/R-93/123a and b, (NTIS PB 93-227163 and 93-27171), Research Triangle Park, NC, July 1993.
- 9. B. Richter, J. Ezzell, and D. Felix, "Single Laboratory Method Validation Report: Extraction of Target Compound List/Priority Pollutant List BNAs and Pesticides using Accelerated Solvent Extraction (ASE) with Analytical Validation by GC/MS and GC/ECD," Document 101124, Dionex Corporation, Salt Lake City, UT, June 16, 1994.
- 10. H. B. Lee, T. E. Peart, R. L. Hong-You, and D. R. Gere, "Supercritical Carbon Dioxide Extraction of Polycyclic Aromatic Hydrocarbons from Sediments," J. Chromatography, A 653 83-91 (1993).
- 11. S. Warner, "SFE Extraction of PNAs from Solid Matrices Using the Dionex 703M SFE Extractor and a Liquid Trap," EPA Region III, Central Regional Laboratory, 839 Bestgate Road, Annapolis, MD 21401, December 12, 1994.
- 12. C. Markell, "3M Data Submission to EPA," letter to B. Lesnik, June 27, 1995.
- 13. USEPA Method 525.2, "Determination of Organic Compounds in Drinking Water by Liquid-Solid Extraction and Capillary Column Gas Chromatography/Mass Spectrometry," Environmental Monitoring Systems Laboratory, Office of Research and Development, US EPA, Cincinnati, OH, Revision 2.0, March 1995.
- 14. USEPA, Superfund Analytical Services/Contract Laboratory Program (CLP), Multi-Media, Multi-Concentration Organics Analysis, SOM01.X, Exhibit D Analytical Methods, "Analytical Method for the Analysis of Semivolatile Organic Compounds," November, 2003

### 17.0 TABLES, DIAGRAMS, FLOW CHARTS, AND VALIDATION DATA

The following pages contain the tables and figures referenced by this method.

TABLE 1

CHARACTERISTIC IONS FOR SEMIVOLATILE COMPOUNDS IN APPROXIMATE RETENTION TIME ORDER <sup>a</sup>

	D :	0 1 1 ()
Compound	Primary Ion	Secondary Ion(s)
2-Picoline	93	66,92
Aniline	93	66,65
Phenol	94	65,66
Bis(2-chloroethyl) ether	93	63,95
2-Chlorophenol	128	64,130
1,3-Dichlorobenzene	146	148,111
1,4-Dichlorobenzene-d <sub>4</sub> (IS)	152	150,115
1,4-Dichlorobenzene	146	148,111
Benzyl alcohol	108	79,77
1,2-Dichlorobenzene	146	148,111
N-Nitrosomethylethylamine	88	42,43,56
Bis(2-chloroisopropyl) ether	45	77,121
Ethyl carbamate	62	44,45,74
Thiophenol (Benzenethiol)	110	66,109,84
Methyl methanesulfonate	80	79,65,95
N-Nitrosodi-n-propylamine	70	42,101,130
Hexachloroethane	117 -	201,199
Maleic anhydride	54	98,53,44
Nitrobenzene	77	123,65
Isophorone	82	95,138
N-Nitrosodiethylamine	102	42,57,44,56
2-Nitrophenol	139	109,65
2,4-Dimethylphenol	122	107,121
p-Benzoquinone	108	54,82,80
Bis(2-chloroethoxy)methane	93	95,123
Benzoic acid	122	105,77
2,4-Dichlorophenol	162	164,98
Trimethyl phosphate	110	79,95,109,140
Ethyl methanesulfonate	79	109,97,45,65
1,2,4-Trichlorobenzene	180	182,145
Naphthalene-d <sub>8</sub> (IS)	136	68
Naphthalene	128	129,127
Hexachlorobutadiene	225	223,227
Tetraethyl pyrophosphate	99	155,127,81,109
Diethyl sulfate	139	45,59,99,111,125
4-Chloro-3-methylphenol	107	144,142
2-Methylnaphthalene	142	141
2-Methylphenol	107	108,77,79,90
Hexachloropropene	213	211,215,117,106,141
Hexachlorocyclopentadiene	237	235,272
N-Nitrosopyrrolidine	100	41,42,68,69
Acetophenone	105	71,51,120
3/4-Methylphenol <sup>b</sup>	107	108,77,79,90

TABLE 1 (continued)

	Primary	Secondary Ion(s)
Compound	lon	occordary form(s)
2,4,6-Trichlorophenol	196	198,200
o-Toluidine	106	107,77,51,79
2-Chloronaphthalene	162	127,164
N-Nitrosopiperidine	114	42,55,56,41
1,4-Phenylenediamine	108	80,53,54,52
1-Chloronaphthalene	162	127,164
2-Nitroaniline	65	92,138
5-Chloro-2-methylaniline	106	141,140,77,89
Dimethyl phthalate	163	194,164
Acenaphthylene	152	151,153
2,6-Dinitrotoluene	165	63,89
Phthalic anhydride	103	76,50,148
o-Anisidine	104	80,123,52
3-Nitroaniline	138	108,92
Acenaphthene-d <sub>10</sub> (IS)	164	162,160
	154	153,152
Acenaphthene	184	63,154
2,4-Dinitrophenol 2,6-Dinitrophenol	16 <del>4</del> 162	164,126,98,63
4-Chloroaniline	102	129,65,92
Isosafrole	162	~
Dibenzofuran	168	131,104,77,51 139
	121	
2,4-Diaminotoluene	165	122,94,77,104
2,4-Dinitrotoluene	139	63,89
4-Nitrophenol	143	109,65 115,116
2-Naphthylamine		104,102,76,50,130
1,4-Naphthoquinone	158	
p-Cresidine	122	94,137,77,93
Dichlorovos	109	185,79,145
Diethyl phthalate	149	177,150
Fluorene	166	165,167
2,4,5-Trimethylaniline	120	135,134,91,77
N-Nitrosodi-n-butylamine	84	57,41,116,158
4-Chlorophenyl phenyl ether	204	206,141
Hydroquinone	110	81,53,55
4,6-Dinitro-2-methylphenol	198	51,105
Resorcinol	110	81,82,53,69
N-Nitrosodiphenylamine	169	168,167
Safrole	162	104,77,103,135
Hexamethyl phosphoramide	135	44,179,92,42
3-(Chloromethyl)pyridine hydrochloride	92	127,129,65,39
Diphenylamine	169	168,167
1,2,4,5-Tetrachlorobenzene	216	214,179,108,143,218
1-Naphthylamine	143	115,89,63
1-Acetyl-2-thiourea	118	43,42,76
4-Bromophenyl phenyl ether	248	250,141

TABLE 1 (continued)

Compound	Primary Ion	Secondary Ion(s)
Toluene diisocyanate	174	145,173,146,132,91
2,4,5-Trichlorophenol	196	198,97,132,99
Hexachlorobenzene	284	142,249
Nicotine	84	133,161,162
Pentachlorophenol	266	264,268
5-Nitro-o-toluidine	152	77,79,106,94
Thionazine	107	96,97,143,79,68
4-Nitroaniline	138	65,108,92,80,39
Phenanthrene-d <sub>10</sub> (IS)	188	94,80
Phenanthrene a <sub>10</sub> (13)	178	179,176
Anthracene	178	176,179
1,4-Dinitrobenzene	168	75,50,76,92,122
Mevinphos	127	192,109,67,164
Naled	109	145,147,301,79,189
1,3-Dinitrobenzene	168	76,50,75,92,122
Diallate (cis or trans)	86	234,43,70
1,2-Dinitrobenzene	168	50,63,74
Diallate (trans or cis)	86	234,43,70
Pentachlorobenzene	250	252,108,248,215,254
5-Nitro-o-anisidine	168	79,52,138,153,77
Pentachloronitrobenzene	237	142,214,249,295,265
4-Nitroquinoline-1-oxide	174	101,128,75,116
Di-n-butyl phthalate	149	150,104
2,3,4,6-Tetrachlorophenol	232	131,230,166,234,168
Dihydrosaffrole	135	64,77
Demeton-O	88	89,60,61,115,171
Fluoranthene	202	101,203
		74,213,120,91,63
1,3,5-Trinitrobenzene	75 127	
Dicrotophos  Ponzidino	127 194	67,72,109,193,237
Benzidine Trifluralia	184	92,185
Trifluralin	306 277	43,264,41,290
Bromoxynil	277	279,88,275,168
Pyrene	202	200,203
Monocrotophos	127	192,67,97,109
Phorate	75 100	121,97,93,260
Sulfallate	188	88,72,60,44
Demeton-S	88	60,81,89,114,115
Phenacetin	108	180,179,109,137,80
Dimethoate	87	93,125,143,229
Phenobarbital	204	117,232,146,161
Carbofuran	164	149,131,122
Octamethyl pyrophosphoramide	135	44,199,286,153,243
4-Aminobiphenyl	169	168,170,115
Dioxathion	97	125,270,153
Terbufos	231	<u>57,97,153,103</u>

TABLE 1 (continued)

Compound	Primary	Secondary Ion(s)
Compound	lon	
$\alpha,\alpha$ -Dimethylphenylamine	58	91,65,134,42
Pronamide	173	175,145,109,147
Aminoazobenzene	197	92,120,65,77
Dichlone	191	163,226,228,135,193
Dinoseb	211	163,147,117,240
Disulfoton	88	97,89,142,186
Fluchloralin	306	63,326,328,264,65
Mexacarbate	165	150,134,164,222
4,4'-Oxydianiline	200	108,171,80,65
Butyl benzyl phthalate	149	91,206
4-Nitrobiphenyl	199	152,141,169,151
Phosphamidon	127	264,72,109,138
2-Cyclohexyl-4,6-Dinitrophenol	231	185,41,193,266
Methyl parathion	109	125,263,79,93
Carbaryl	144	115,116,201
•	225	120,77,105,148,42
Dimethylaminoazobenzene		
Propylthiouracil	170	142,114,83
Benz(a)anthracene	228	229,226
Chrysene-d <sub>12</sub> (IS)	240	120,236
3,3'-Dichlorobenzidine	252	254,126
Chrysene	228	226,229
Malathion	173	125,127,93,158
Kepone	272	274,237,178,143,270
Fenthion	278	125,109,169,153
Parathion	109	97,291,139,155
Anilazine	239	241,143,178,89
Bis(2-ethylhexyl) phthalate	149	167,279
3,3'-Dimethylbenzidine	212	106,196,180
Carbophenothion	157	97,121,342,159,199
5-Nitroacenaphthene	199	152,169,141,115
Methapyrilene	97	50,191,71
Isodrin	193	66,195,263,265,147
Captan	79	149,77,119,117
Chlorfenvinphos	267	269,323,325,295
Crotoxyphos	127	105,193,166
Phosmet	160	77,93,317,76
EPN	157	169,185,141,323
Tetrachlorvinphos	329	109,331,79,333
Di-n-octyl phthalate	149	167,43
* •	223	167,43 167,195
2-Aminoanthraquinone		•
Barban	222	51,87,224,257,153
Aramite	185	191,319,334,197,321
Benzo(b)fluoranthene	252	253,125
Nitrofen	283	285,202,139,253
Benzo(k)fluoranthene	252	253,125

TABLE 1 (continued)

Primary   Secondary Ion(s)   Compound   Chlorobenzilate   251   139,253,111,141   139,253,111,141   141   151		Drimony	Secondary Ion(s)
Chlorobenzilate	Compound	•	Secondary IOH(S)
Ensulfothion 293 97,308,125,292 Ethion 231 97,153,125,121 10 10 10 10 10 10 10 10 10 10 10 10 10	•		139.253.111.141
Ethion 231 97,153,125,121 Diethylstilbestrol 268 145,107,239,121,159 Famphur 218 125,93,109,217 Tri-p-tolyl phosphate* 368 367,107,165,198 Benzo(a)pyrene 252 253,125 Perylene-d <sub>12</sub> (IS) 264 260,265 T,12-Dimethylbenz(a)anthracene 256 241,239,120 Dinocap 180 104,252,223,209 Dinocap 69 41,39 Methoxychlor 227 228,152,114,274,212 P-Acetylaminofluorene 181 180,223,152 P-Acetylaminofluorene 181 180,223,152 P-Methylcholanthrene 268 252,253,126,134,113 Phosalone 182 184,367,121,379 Dinocap 184 184,367,121,379 Dinocap 185 252,253,126,134,113 Phosalone 182 184,367,121,379 Dinocap 185 277 Dinocap 187 1377,375,77,155,379 Dinocap 187 1377,375,77,155,379 Dinocap 188 189,227,250 Dinocap 189 280,277,250 Dinocap 189 299 262,210,364,97,109 Dinocap 189 299,211,406,408 Dinocap 189 299,211,406,408 Dinocap 189 292 362,326 Dinocap 292 362,326			
Diethylstilbestrol         268         145,107,239,121,159           Famphur         218         125,93,109,217           Fri-p-tolyl phosphate <sup>c</sup> 368         367,107,165,198           Benzo(a)pyrene         252         253,125           Perylene-d <sub>12</sub> (IS)         264         260,265           7,12-Dimethylbenz(a)anthracene         256         241,239,120           3,5-Diphenylhydantoin         180         104,252,223,209           Captafol         79         77,80,107           Dinocap         69         41,39           Methoxychlor         227         228,152,114,274,212           2-Acetylaminofluorene         181         180,223,152           3,3-Dimethoxybenzidine         244         201,229           3-Methylcholanthrene         268         252,253,126,134,113           2-Nosalone         182         184,367,121,379           3-Inex         272         237,274,270,239,235           Airex         272         237,74,270,239,235           Airex         272         237,77,250           Alestranol         277         310,174,147,242           Coumaphos         362         226,210,364,97,109           Dibenz(a,h)anthracene         276			
Famphur         218         125,93,109,217           Fri-p-tolyl phosphate <sup>6</sup> 368         367,107,165,198           Benzo(a)pyrene         252         253,125           Perylene-d <sub>12</sub> (IS)         264         260,265           7,12-Dimethylbenz(a)anthracene         256         241,239,120           5,5-Diphenylhydantoin         180         104,252,223,209           Captafol         79         77,80,107           Dinocap         69         41,39           Methoxychlor         227         228,152,114,274,212           2-Acetylaminofluorene         181         180,223,152           4,4-Methylenebis(2-chloroaniline)         231         266,268,140,195           3,3-Dimethoxybenzidine         244         201,229           3-Methylcholanthrene         268         252,253,126,134,113           2-Nosalone         182         184,367,121,379           Azinphos-methyl         160         132,93,104,105           2-Tris(2,3-dibromopropyl) phosphate         201         137,119,217,219,199           Dibenz(a,j)acridine         279         280,277,250           Mestranol         277         310,174,147,242           Doumaphos         362         226,210,364,97,109			
Tri-p-tolyl phosphate <sup>c</sup> Benzo(a)pyrene Benzo(a)pyrene Berylene-d <sub>12</sub> (IS) Captafol Dinocap Methoxychlor Be-devlene-discording by the toleration of the tole	•		
Benzo(a)pyrene       252       253,125         Perylene-d <sub>12</sub> (IS)       264       260,265         7,12-Dimethylbenz(a)anthracene       256       241,239,120         5,5-Diphenylhydantoin       180       104,252,223,209         Captafol       79       77,80,107         Dinocap       69       41,39         Methoxychlor       227       228,152,114,274,212         2-Acetylaminofluorene       181       180,223,152         3,3-Dimethoxybenzidine       244       201,229         3-Methylcholanthrene       268       252,253,126,134,113         2-Nosalone       182       184,367,121,379         3-Discopende       182       184,367,121,379         3-Phosalone       182       184,367,121,379         3-Inicx       272       237,274,270,239,235         3-Tris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Meneno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Bernzo(g,h,i)perylene       276       138,277         3-Cy-bibenzopyrene       302 </td <td>•</td> <td></td> <td></td>	•		
Perylene-d <sub>12</sub> (IS) 264 260,265 7,12-Dimethylbenz(a) anthracene 256 241,239,120 6,5-Diphenylhydantoin 180 104,252,223,209 7,80,107 Dinocap 69 41,39 Methoxychlor 227 228,152,114,274,212 7,2-Acetylaminofluorene 181 180,223,152 7,4-Methylenebis(2-chloroaniline) 231 266,268,140,195 7,3-Dimethoxybenzidine 244 201,229 7,3-Dimethoxybenzidine 256 252,253,126,134,113 7,3-Diosalone 182 184,367,121,379 7,3-Diosalone 182 184,367,121,379 7,3-Diosalone 171 377,375,77,155,379 7,3-Directory 171 377,375,77,155,379 7,3-Dibenz(a,j)acridine 279 280,277,250 7,3-Dibenz(a,j)acridine 279 280,277,250 7,3-Dibenz(a,j)acridine 279 280,277,250 7,3-Dibenz(a,j)acridine 279 280,277,250 7,3-Dibenz(a,j)acridine 279 130,174,147,242 7,3-Dibenz(a,j)aridine 279 280,277,250 7,3-Dibenz(a,j)aridin			
7,12-Dimethylbenz(a)anthracene 256 241,239,120 5,5-Diphenylhydantoin 180 104,252,223,209 77,80,107 77,80,107 79 77,80,107 190 227 228,152,114,274,212 227 228,152,114,274,212 228,152,114,274,212 228,152,114,274,212 231 266,268,140,195 231 266,268,140,195 231 266,268,140,195 244 201,229 242,101,201 201,202 201,202 201,203 201,			-
5,5-Diphenylhydantoin Captafol	12 1		•
Captafol       79       77,80,107         Dinocap       69       41,39         Methoxychlor       227       228,152,114,274,212         2-Acetylaminofluorene       181       180,223,152         4,4-Methylenebis(2-chloroaniline)       231       266,268,140,195         3,3'-Dimethoxybenzidine       244       201,229         8-Methylcholanthrene       268       252,253,126,134,113         Phosalone       182       184,367,121,379         Azinphos-methyl       160       132,93,104,105         Leptophos       171       377,375,77,155,379         Airex       272       237,274,270,239,235         Tris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         ndeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Berzo(g,h,i)perylene       276       138,277         3,2-2,4,5-Dibenzopyrene       302       151,150,300         347       334,335,333       334         Piperonyl sulfoxide       162 </td <td></td> <td></td> <td></td>			
Dinocap       69       41,39         Methoxychlor       227       228,152,114,274,212         2-Acetylaminofluorene       181       180,223,152         4,4'-Methylenebis(2-chloroaniline)       231       266,268,140,195         3,3'-Dimethoxybenzidine       244       201,229         3-Methylcholanthrene       268       252,253,126,134,113         Phosalone       182       184,367,121,379         Azinphos-methyl       160       132,93,104,105         Jeptophos       171       377,375,77,155,379         Mirex       272       237,274,270,239,235         Tris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         Jerophyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1221			
Methoxychlor       227       228,152,114,274,212         2-Acetylaminofluorene       181       180,223,152         4,4'-Methylenebis(2-chloroaniline)       231       266,268,140,195         3,3'-Dimethoxybenzidine       244       201,229         3-Methylcholanthrene       268       252,253,126,134,113         2-Nosalone       182       184,367,121,379         Azinphos-methyl       160       132,93,104,105         3-ptophos       171       377,375,77,155,379         Mirex       272       237,274,270,239,235         Tris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Brychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         dexachlorophene       196       198,209,211,406,408         <	•		
2-Acetylaminofluorene 3,4'-Methylenebis(2-chloroaniline) 3,3'-Dimethoxybenzidine 244 201,229 3-Methylcholanthrene	·		,
4,4'-Methylenebis(2-chloroaniline)       231       266,268,140,195         3,3'-Dimethoxybenzidine       244       201,229         3-Methylcholanthrene       268       252,253,126,134,113         Phosalone       182       184,367,121,379         Azinphos-methyl       160       132,93,104,105         Jeptophos       171       377,375,77,155,379         Airex       272       237,274,270,239,235         Fris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Olibenz(a,h)anthracene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         J,2:4,5-Dibenzopyrene       302       151,150,300         Brychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1221       190       224,260         Aroclor 1242       22			
3,3'-Dimethoxybenzidine 3-Methylcholanthrene 3-Methylcholanthrene 3-Methylcholanthrene 3-Azinphos-methyl 3-Azinphos-methyl 3-Phosalone 3-Azinphos-methyl 3-Phosalone 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos-methyl 3-Azinphos			• •
## Process of Section 1985 (1985)			
Phosalone 182 184,367,121,379 Azinphos-methyl 160 132,93,104,105 Leptophos 171 377,375,77,155,379 Mirex 272 237,274,270,239,235 Tris(2,3-dibromopropyl) phosphate 201 137,119,217,219,199 Dibenz(a,j)acridine 279 280,277,250 Mestranol 277 310,174,147,242 Coumaphos 362 226,210,364,97,109 Indeno(1,2,3-cd)pyrene 276 138,277 Dibenz(a,h)anthracene 278 139,279 Benzo(g,h,i)perylene 276 138,277 L,2:4,5-Dibenzopyrene 302 151,150,300 Strychnine 334 334,335,333 Piperonyl sulfoxide 162 135,105,77 Hexachlorophene 196 198,209,211,406,408 Addrin 66 263,220 Aroclor 1016 222 260,292 Aroclor 1221 190 224,260 Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394	•		•
Azinphos-methyl Leptophos Leptophos Mirex  272 237,274,270,239,235  Tris(2,3-dibromopropyl) phosphate Dibenz(a,j)acridine  Mestranol Coumaphos Indeno(1,2,3-cd)pyrene Dibenz(a,h)anthracene Dibenz(a,b)anthracene Dibenz(a,b			
Leptophos       171       377,375,77,155,379         Mirex       272       237,274,270,239,235         Cris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Strychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
Mirex       272       237,274,270,239,235         Tris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Strychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
Tris(2,3-dibromopropyl) phosphate       201       137,119,217,219,199         Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Strychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
Dibenz(a,j)acridine       279       280,277,250         Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Birychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
Mestranol       277       310,174,147,242         Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         J,2:4,5-Dibenzopyrene       302       151,150,300         Brychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394	1 1 1 1 1 1		
Coumaphos       362       226,210,364,97,109         Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Strychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394	/lestranol		
Indeno(1,2,3-cd)pyrene       276       138,277         Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Brychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394	010111		
Dibenz(a,h)anthracene       278       139,279         Benzo(g,h,i)perylene       276       138,277         1,2:4,5-Dibenzopyrene       302       151,150,300         Brychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394	•		
Benzo(g,h,i)perylene       276       138,277         I,2:4,5-Dibenzopyrene       302       151,150,300         Brychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
1,2:4,5-Dibenzopyrene       302       151,150,300         Strychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
Strychnine       334       334,335,333         Piperonyl sulfoxide       162       135,105,77         Hexachlorophene       196       198,209,211,406,408         Aldrin       66       263,220         Aroclor 1016       222       260,292         Aroclor 1221       190       224,260         Aroclor 1232       190       224,260         Aroclor 1242       222       256,292         Aroclor 1248       292       362,326         Aroclor 1254       292       362,326         Aroclor 1260       360       362,394			
Piperonyl sulfoxide 162 135,105,77 Hexachlorophene 196 198,209,211,406,408 Aldrin 66 263,220 Aroclor 1016 222 260,292 Aroclor 1221 190 224,260 Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394	• •		
Hexachlorophene 196 198,209,211,406,408 Aldrin 66 263,220 Aroclor 1016 222 260,292 Aroclor 1221 190 224,260 Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394	•		
Aldrin 66 263,220 Aroclor 1016 222 260,292 Aroclor 1221 190 224,260 Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394	•		
Aroclor 1016 222 260,292 Aroclor 1221 190 224,260 Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394	•		
Aroclor 1221 190 224,260 Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394			•
Aroclor 1232 190 224,260 Aroclor 1242 222 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394			
Aroclor 1242 22 256,292 Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394			
Aroclor 1248 292 362,326 Aroclor 1254 292 362,326 Aroclor 1260 360 362,394			
Aroclor 1254 292 362,326 Aroclor 1260 360 362,394			
Aroclor 1260 360 362,394			
•			
			•
3-BHC 181 183,109			
5-BHC 183 181,109			
r-BHC (Lindane) 183 181,109			
1,4'-DDD 235 237,165	· · · · · · · · · · · · · · · · · · ·		

TABLE 1 (continued)

	Primary	Secondary Ion(s)
Compound	lon	• ( )
4,4'-DDE	246	248,176
4,4'-DDT	235	237,165
Dieldrin	79	263,279
1,2-Diphenylhydrazine	77	105,182
Endosulfan I	195	339,341
Endosulfan II	337	339,341
Endosulfan sulfate	272	387,422
Endrin	263	82,81
Endrin aldehyde	67	345,250
Endrin ketone	317	67,319
2-Fluorobiphenyl (surr)	172	171
2-Fluorophenol (surr)	112	64
Heptachlor	100	272,274
Heptachlor epoxide	353	355,351
Nitrobenzene-d <sub>5</sub> (surr)	82	128,54
N-Nitrosodimethylamine	42	74,44
Phenol-d <sub>6</sub> (surr)	99	42,71
Terphenyl-d <sub>14</sub> (surr)	244	122,212
2,4,6-Tribromophenol (surr)	330	332,141
Toxaphene	159	231,233

IS = internal standard

surr = surrogate

a The data presented are representative of DB-5 type analytical columns
b Compounds cannot be separated for quantitation
c Substitute for the non-specific mixture, tricresyl phosphate

TABLE 2

EXAMPLE LOWER LIMITS OF QUANTITATION FOR SEMIVOLATILE ORGANICS

Compound         (μg/L)         (μg/kg)           Acenaphthene         10         Acenaphthylene         10           Acetophenone         10         2           2-Acetylaminofluorene         20         1-Acetyl-2-thiourea         1000           2-Aminoarthraquinone         20         Aminoazobenzene         10           4-Aminobiphenyl         20         Amilazine         100           o-Anisidine         10         Anthracene         10           Aramite         20         Azinphos-methyl         100           Barban         200         Benz(a)anthracene         10           Benzo(b)fluoranthene         10         Benzo(b)fluoranthene         10           Benzo(g,h,i)perylene         10         Benzo(a)pyrene         10           Benzo(a)pyrene         10         Benzo(a)pyrene         10           p-Benzoquinone         10         10           Benzo(a)p		Lower Limits o	f Quantitation <sup>a</sup>
Acenaphthylene	ound		Low Soil/Sediment <sup>b</sup> (μg/kg)
Acetophenone 10 2-Acetyl-aminofluorene 20 1-Acetyl-2-thiourea 1000 2-Aminoanthraquinone 20 Aminoazobenzene 10 4-Aminobiphenyl 20 Anilazine 100 o-Anisidine 100 Aramite 20 Azinphos-methyl 100 Barban 200 Benzo(a),pirene 10 Benzo(a),pirene 10 Benzoquinone 10 Bis(2-chloroethoxy)methane 10 Bis(2-chlorosopropyl) ether 10 Bis(2-chlorosopropyl) ether 10 Captafol 20 Captan 50 Carbaryl 10 Carbofuran 10 Carbofuran 10 Carbofuroniline 20 11	phthene	10	660
2-Acetylaminofluorene       20         1-Acetyl-2-thiourea       1000         2-Aminoanthraquinone       20         Aminoazobenzene       10         4-Aminobiphenyl       20         Anilazine       100         o-Anisidine       10         Anthracene       10         Aramite       20         Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         P-Benzoquinone       10         Benzo(a)pyrene       10         P-Benzoquinone       10         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroethyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Captafol       20	phthylene	10	660
1-Acetyl-2-thiourea       1000         2-Aminoanthraquinone       20         Aminoazobenzene       10         4-Aminobiphenyl       20         Anilazine       100         o-Anisidine       10         Anthracene       10         Aramite       20         Azinphos-methyl       100         Barban       200         Benza(a)anthracene       10         Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(g,h,i)perylene       10         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Beis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroethyl) ether       10         Gradian       50	phenone	10	ND
2-Aminoarthraquinone       20         Aminoazobenzene       10         4-Aminobiphenyl       20         Anilazine       100         o-Anisidine       10         Anthracene       10         Aramite       20         Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(acid       50         Benzo(acid       10         Benzo(acid       10         Benzo(acid       10         Benz	tylaminofluorene	20	ND
Aminoazobenzene       10         4-Aminobiphenyl       20         Anilazine       100         o-Anisidine       10         Anthracene       10         Aramite       20         Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(k)fluoranthene       10         Benzo(g,h,i)perylene       10         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethoxy)methane       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bortonoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captar       50         Carbaryl       10         Carboyluna       10         Carbophenothion       10         Chloroanilline	tyl-2-thiourea	1000	ND
4-Aminobiphenyl       20         Anilazine       100         o-Anisidine       10         Anthracene       10         Aramite       20         Azinphos-methyl       100         Barban       200         Benza(a)anthracene       10         Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzoic acid       50         Benzoic acid       50         Benzo(g,h,i)perylene       10         Benzo(a)apyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carboylenothion       10         Chlordenvinphos       20         4-Chloroaniline       20	noanthraquinone	20	ND
Anilazine	pazobenzene	10	ND
o-Anisidine       10         Anthracene       10         Aramite       20         Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzoic acid       50         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroethyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captar       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chloroaniline       20         4-Chloroaniline       20	nobiphenyl	20	ND
Anthracene 10 Aramite 20 Azinphos-methyl 100 Barban 200 Benz(a)anthracene 10 Benzo(b)fluoranthene 10 Benzoic acid 50 3 Benzo(a,h,i)perylene 10 Benzo(a)pyrene 10 Benzoquinone 10 Benzoquinone 10 Bis(2-chloroethoxy)methane 10 Bis(2-chloroisopropyl) ether 10 Bis(2-chloroisopropyl) ether 10 Bis(benzoyil benzyl phthalate 10 Bromoxynil 10 Butyl benzyl phthalate 10 Captafol 20 Captan 50 Carbaryl 10 Carbophenothion 10 Chlorfenvinphos 20 4-Chloroaniline 20 11 Cavo	•	100	ND
Aramite       20         Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzoic acid       50         Benzoic acid       50         Benzo(a)pyrene       10         Benzoqaijnone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captaryl       10         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chloroaniline       20         4-Chloroaniline       20	sidine	10	ND
Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzoic acid       50         Benzoic acid       50         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chloroaniline       20         4-Chloroaniline       20	acene	10	660
Azinphos-methyl       100         Barban       200         Benz(a)anthracene       10         Benzo(b)fluoranthene       10         Benzoic k)fluoranthene       10         Benzoic acid       50         Benzoic acid       50         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captaryl       10         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chloroaniline       20         4-Chloroaniline       20	te	20	ND
Barban         200           Benz(a)anthracene         10           Benzo(b)fluoranthene         10           Benzo(k)fluoranthene         10           Benzoic acid         50           Benzo(g,h,i)perylene         10           Benzo(a)pyrene         10           p-Benzoquinone         10           Benzyl alcohol         20         1           Bis(2-chloroethoxy)methane         10           Bis(2-chloroethyl) ether         10           Bis(2-chloroisopropyl) ether         10           4-Bromophenyl phenyl ether         10           Bromoxynil         10           Butyl benzyl phthalate         10           Captafol         20           Captaryl         10           Carbofuran         10           Carbophenothion         10           Chlorfenvinphos         20           4-Chloroaniline         20           1         10	hos-methyl	100	ND
Benzo(b)fluoranthene       10         Benzo(c)(k)fluoranthene       10         Benzoic acid       50       3         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captarn       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20	•	200	ND
Benzo(b)fluoranthene       10         Benzo(k)fluoranthene       10         Benzoic acid       50       3         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captarn       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20       1	a)anthracene	10	660
Benzo(k)fluoranthene       10         Benzoic acid       50       3         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10       9         p-Benzoquinone       10       9         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10       10         Bis(2-chloroethyl) ether       10       10         4-Bromophenyl penyl ether       10       10         Bromoxynil       10       10         Butyl benzyl phthalate       10       20         Captafol       20       20         Carbaryl       10       20         Carbofuran       10       20         Carbophenothion       10       20         Chlorfenvinphos       20       4-Chloroaniline       20			660
Benzoic acid       50       3         Benzo(g,h,i)perylene       10         Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20       1			660
Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20		50	3300
Benzo(a)pyrene       10         p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20	o(g,h,i)perylene	10	660
p-Benzoquinone       10         Benzyl alcohol       20       1         Bis(2-chloroethoxy)methane       10         Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20	p(a)pyrene	tific line	01100126601
Bis(2-chloroethoxy)methane10Bis(2-chloroethyl) ether10Bis(2-chloroisopropyl) ether104-Bromophenyl phenyl ether10Bromoxynil10Butyl benzyl phthalate10Captafol20Captan50Carbaryl10Carbofuran10Carbophenothion10Chlorfenvinphos204-Chloroaniline20	zoquinone		ND
Bis(2-chloroethoxy)methane10Bis(2-chloroethyl) ether10Bis(2-chloroisopropyl) ether104-Bromophenyl phenyl ether10Bromoxynil10Butyl benzyl phthalate10Captafol20Captan50Carbaryl10Carbofuran10Carbophenothion10Chlorfenvinphos204-Chloroaniline20	rl alcohol	20	1300
Bis(2-chloroethyl) ether       10         Bis(2-chloroisopropyl) ether       10         4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20		10	660
4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20	chloroethyl) ether	10	660
4-Bromophenyl phenyl ether       10         Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20	• /	10	660
Bromoxynil       10         Butyl benzyl phthalate       10         Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20		10	660
Butyl benzyl phthalate 10 Captafol 20 Captan 50 Carbaryl 10 Carbofuran 10 Carbophenothion 10 Chlorfenvinphos 20 4-Chloroaniline 20 1	. , , ,	10	ND
Captafol       20         Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20	•	10	660
Captan       50         Carbaryl       10         Carbofuran       10         Carbophenothion       10         Chlorfenvinphos       20         4-Chloroaniline       20		20	ND
Carbaryl 10 Carbofuran 10 Carbophenothion 10 Chlorfenvinphos 20 4-Chloroaniline 20 1		50	ND
Carbofuran10Carbophenothion10Chlorfenvinphos204-Chloroaniline20			ND
Carbophenothion10Chlorfenvinphos204-Chloroaniline20			ND
Chlorfenvinphos 20 4-Chloroaniline 20 1			ND
4-Chloroaniline 20 1	•		ND
	•		1300
CHOIODEHZHALE	obenzilate	10	ND
5-Chloro-2-methylaniline 10			ND
•	•		1300

TABLE 2 (continued)

	Lower Limits	of Quantitation <sup>a</sup>
Compound	Ground water (μg/L)	Low Soil/Sediment <sup>b</sup> (µg/kg)
3-(Chloromethyl)pyridine hydrochloride	100	ND
2-Chloronaphthalene	10	660
2-Chlorophenol	10	660
4-Chlorophenyl phenyl ether	10	660
Chrysene	10	660
Coumaphos	40	ND
p-Cresidine	10	ND
Crotoxyphos	20	ND
2-Cyclohexyl-4,6-dinitrophenol	100	ND
Demeton-O	10	ND
Demeton-S	10	ND
Diallate (cis or trans)	10	ND
Diallate (trans or cis)	10	ND
2,4-Diaminotoluene	20	ND
Dibenz(a,j)acridine	10	ND
Dibenz(a,h)anthracene	10	660
Dibenzofuran Diben	10	660
Dibenzo(a,e)pyrene	10	ND
Di-n-butyl phthalate	10	ND
Dichlone	_NA	ND
1,2-Dichlorobenzene	T1T1C 1101	201001660101
1,3-Dichlorobenzene	10	660
1,4-Dichlorobenzene	10	660
3,3'-Dichlorobenzidine	20	1300
2,4-Dichlorophenol	10	660
2,6-Dichlorophenol	10	ND
Dichlorovos	10	ND
Dicrotophos	10	ND
Diethyl phthalate	10	660
Diethylstilbestrol	20	ND
Diethyl sulfate	100	ND
Dimethoate	20	ND
3,3'-Dimethoxybenzidine	100	ND
Dimethylaminoazobenzene	10	ND
7,12-Dimethylbenz(a)anthracene	10	ND
3,3'-Dimethylbenzidine	10	ND
2,4-Dimethylphenol	10	660
Dimethyl phthalate	10	660
1,2-Dinitrobenzene	40	ND

TABLE 2 (continued)

	Lower Limits	of Quantitation <sup>a</sup>
Compound	Ground water (μg/L)	Low Soil/Sediment <sup>b</sup> (µg/kg)
1,3-Dinitrobenzene	20	ND
1,4-Dinitrobenzene	40	ND
4,6-Dinitro-2-methylphenol	50	3300
2,4-Dinitrophenol	50	3300
2,4-Dinitrotoluene	10	660
2,6-Dinitrotoluene	10	660
Dinocap	100	ND
Dinoseb	20	ND
5,5-Diphenylhydantoin	20	ND
Di-n-octyl phthalate	10	660
Disulfoton	10	ND
EPN	10	ND
Ethion	10	ND
Ethyl carbamate	50	ND
Bis(2-ethylhexyl) phthalate	10	660
Ethyl methanesulfonate	20	ND
Famphur	20	ND
Fensulfothion	40	ND
Fenthion	10	ND
Fluchloralin	20	ND
Fluoranthene	t1110 1100	
Fluorene	10	660
Hexachlorobenzene	10	660
Hexachlorobutadiene	10	660
Hexachlorocyclopentadiene	10	660
Hexachloroethane	10	660
Hexachlorophene	50	ND
Hexachloropropene	10	ND
Hexamethylphosphoramide	20	ND
Indeno(1,2,3-cd)pyrene	10	660
Isodrin	20	ND
Isophorone	10	660
Isosafrole	10	ND
Kepone	20	ND
Leptophos	10	ND
Malathion	50	ND
Mestranol	20	ND
Methapyrilene	100	ND
Methoxychlor	10	ND

TABLE 2 (continued)

	Lower Limits	of Quantitation <sup>a</sup>
Compound	Ground water (μg/L)	Low Soil/Sediment <sup>b</sup> (µg/kg)
3-Methylcholanthrene	10	ND
Methyl methanesulfonate	10	ND
2-Methylnaphthalene	10	660
Methyl parathion	10	ND
2-Methylphenol	10	660
3-Methylphenol	10	ND
4-Methylphenol	10	660
Mevinphos	10	ND
Mexacarbate	20	ND
Mirex	10	ND
Monocrotophos	40	ND
Naled	20	ND
Naphthalene	10	660
1,4-Naphthoquinone	10	ND
1-Naphthylamine	10	ND
2-Naphthylamine	10	ND
Nicotine	20	ND
5-Nitroacenaphthene	10	ND
2-Nitroaniline	50	3300
3-Nitroaniline	50	3300
4-Nitroaniline	1111C 20	COLDOLANDIOI
5-Nitro-o-anisidine	10	ND
Nitrobenzene	10	660
4-Nitrobiphenyl	10	ND
Nitrofen	20	ND
2-Nitrophenol	10	660
4-Nitrophenol	50	3300
5-Nitro-o-toluidine	10	ND
4-Nitroquinoline-1-oxide	40	ND
N-Nitrosodi-n-butylamine	10	ND
N-Nitrosodiethylamine	20	ND
N-Nitrosodiphenylamine	10	660
N-Nitroso-di-n-propylamine	10	660
N-Nitrosopiperidine	20	ND
N-Nitrosopyrrolidine	40	ND
Octamethyl pyrophosphoramide	200	ND
4,4'-Oxydianiline	20	ND
Parathion	10	ND
Pentachlorobenzene	10	ND

TABLE 2 (continued)

_		Lower Limits	of Quantitation <sup>a</sup>
Compound		Ground water (μg/L)	Low Soil/Sediment <sup>b</sup> (µg/kg)
Pentachloronitrobe	enzene	20	ND
Pentachloropheno	I	50	3300
Phenacetin		20	ND
Phenanthrene		10	660
Phenobarbital		10	ND
Phenol		10	660
1,4-Phenylenediar	mine	10	ND
Phorate		10	ND
Phosalone		100	ND
Phosmet		40	ND
Phosphamidon		100	ND
Phthalic anhydride	<b>)</b>	100	ND
2-Picoline		ND	ND
Piperonyl sulfoxide	Э	100	ND
Pronamide		10	ND
Propylthiouracil Propylthiouracil		100	ND
Pyrene		10	660
Resorcinol		100	ND
Safrole		10	ND
Strychnine	~ .	40	ND
Sulfallate	Scien	ntific Inc	corporandlo
Terbufos		20	ND
1,2,4,5-Tetrachlor	obenzene	10	ND
2,3,4,6-Tetrachlor	ophenol	10	ND
Tetrachlorvinphos		20	ND
Tetraethyl pyropho	sphate	40	ND
Thionazine		20	ND
Thiophenol (Benze	enethiol)	20	ND
o-Toluidine		10	ND
1,2,4-Trichloroben	zene	10	660
2,4,5-Trichlorophe	nol	10	660
2,4,6-Trichlorophe	nol	10	660
Trifluralin		10	ND
2,4,5-Trimethylani	line	10	ND
Trimethyl phospha	te	10	ND
1,3,5-Trinitrobenze	ene	10	ND
Tris(2,3-dibromopr	opyl) phosphate	200	ND
	nte(h)	10	ND

### TABLE 2 (continued)

- <sup>a</sup> Sample lower limits of quantitation are highly matrix-dependent and those listed here are provided for guidance and may not always be achievable.
- b Lower limits of quantitation listed for soil/sediment are based on wet weight. When data are reported on a dry weight basis, the lower limits will be higher based on the % dry weight of each sample. These lower limits are based on a 30-g sample and gel permeation chromatography cleanup.

ND = Not Determined

NA = Not Applicable

Other Matrices	<u>Factor</u> <sup>c</sup>
High-concentration soil and sludges by ultrasonic extractor	7.5
Non-water miscible waste	75

<sup>c</sup>Lower limit of quantitation = (Lower limit of quantitation for low soil/sediment given above in Table 2) x (Factor)



SpectraLab

Scientific Incorporation

TABLE 3

DFTPP KEY IONS AND ION ABUNDANCE CRITERIA<sup>a,b</sup>

Mass	Ion Abundance Criteria
51	10-80% of Base Peak
68	< 2% of mass 69
70	< 2% of mass 69
127	10-80% of Base Peak
197	< 2% of mass 198
198	Base peak, or > 50% of Mass 442
199	5-9% of mass 198
275	10-60% of Base Peak
365	> 1% of mass 198
441	present but < 24% of mass 442
442	Base Peak, or > 50% of mass 198
443	15-24% of mass 442



<sup>a</sup> The majority of the data are taken from Reference 13 (Method 525.2).

The criteria in this table are intended to be used as default criteria for quadrupole instrumentation if optimized manufacturer's operating conditions are not available. Alternate tuning criteria may be employed (e.g., CLP or Method 625), provided that method performance is not adversely affected. See Sec. 11.3.1

TABLE 4

### RECOMMENDED MINIMUM RESPONSE FACTOR CRITERIA FOR INITIAL AND CONTINUING CALIBRATION VERIFICATION USING THE SUGGESTED IONS FROM TABLE 1

Semivolatile Compounds	Minimum Response Factor (RF)
Benzaldehyde	0.010
Phenol	0.800
Bis(2-chloroethyl)ether	0.700
2-Chlorophenol	0.800
2-Methylphenol	0.700
2,2'-Oxybis-(1-chloropropane)	0.010
Acetophenone	0.010
4-Methylphenol	0.600
N-Nitroso-di-n-propylamine	0.500
Hexachloroethane	0.300
Nitrobenzene	0.200
Isophorone	0.400
2-Nitrophenol	0.100
2,4-Dimethylphenol	0.200 corporation
Bis(2-chloroethoxy)methane	0.300
2,4-Dichlorophenol	0.200
Naphthalene	0.700
4-Chloroaniline	0.010
Hexachlorobutadiene	0.010
Caprolactam	0.010
4-Chloro-3-methylphenol	0.200
2-Methylnaphthalene	0.400
Hexachlorocyclopentadiene	0.050
2,4,6-Trichlorophenol	0.200
2,4,5-Trichlorophenol	0.200
1,1'-Biphenyl	0.010

0.800

2-Chloronaphthalene

TABLE 4 (continued)

٠	Semivolatile Compounds	Minimum Response Factor (RF)
•	2-Nitroaniline	0.010
	Dimethyl phthalate	0.010
	2,6-Dinitrotoluene	0.200
	Acenaphthylene	0.900
	3-Nitroaniline	0.010
	Acenaphthene	0.900
	2,4-Dinitrophenol	0.010
	4-Nitrophenol	0.010
	Dibenzofuran	0.800
	2,4-Dinitrotoluene	0.200
	Diethyl phthalate	0.010
	1,2,4,5-Tetrachlorobenzene	0.010
	4-Chlorophenyl-phenyl ether	0.400
	Fluorene	0.900
	4-Nitroaniline	0.010
	4,6-Dinitro-2-methylphenol	0.010
	4-Bromophenyl-phenyl ether	0.1001 Corporation
	N-Nitrosodiphenylamine	0.010
	Hexachlorobenzene	0.100
	Atrazine	0.010
	Pentachlorophenol	0.050
	Phenanthrene	0.700
	Anthracene	0.700
	Carbazole	0.010
	Di-n-butyl phthalate	0.010
	Fluoranthene	0.600
	Pyrene	0.600
	Butyl benzyl phthalate	0.010
	3,3'-Dichlorobenzidine	0.010
	Benzo(a)anthracene	0.800

TABLE 4 (continued)

Semivolatile Compounds	Minimum Response Factor (RF)
Chrysene	0.700
Bis-(2-ethylhexyl)phthalate	0.010
Di-n-octyl phthalate	0.010
Benzo(b)fluoranthene	0.700
Benzo(k)fluoranthene	0.700
Benzo(a)pyrene	0.700
Indeno(1,2,3-cd)pyrene	0.500
Dibenz(a,h)anthracene	0.400
Benzo(g,h,i)perylene	0.500
2,3,4,6-Tetrachlorophenol	0.010



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TABLE 5

SEMIVOLATILE INTERNAL STANDARDS WITH CORRESPONDING ANALYTES ASSIGNED FOR QUANTITATION

1,4-Dichlorobenzene-d <sub>4</sub>	Naphthalene-d <sub>8</sub>	Acenaphthene-d <sub>10</sub>
Aniline	Acetophenone	Acenaphthene
Benzyl alcohol	Benzoic acid	Acenaphthylene
Bis(2-chloroethyl) ether	Bis(2-chloroethoxy)methane	1-Chloronaphthalene
Bis(2-chloroisopropyl) ether	4-Chloroaniline	2-Chloronaphthalene
2-Chlorophenol	4-Chloro-3-methylphenol	4-Chlorophenyl phenyl ether
1,3-Dichlorobenzene	2,4-Dichlorophenol	Dibenzofuran
1,4-Dichlorobenzene	2,6-Dichlorophenol	Diethyl phthalate
1,2-Dichlorobenzene	$\alpha,\alpha$ -Dimethyl-	Dimethyl phthalate
Ethyl methanesulfonate	phenethylamine	2,4-Dinitrophenol
2-Fluorophenol (surr)	2,4-Dimethylphenol	2,4-Dinitrotoluene
Hexachloroethane	Hexachlorobutadiene	2,6-Dinitrotoluene
Methyl methanesulfonate	Isophorone	Fluorene
2-Methylphenol	2-Methylnaphthalene	2-Fluorobiphenyl (surr)
4-Methylphenol	Naphthalene	Hexachlorocyclopentadiene
N-Nitrosodimethylamine	Nitrobenzene	1-Naphthylamine
N-Nitroso-di-n-propylamine	Nitrobenzene-d <sub>8</sub> (surr)	2-Naphthylamine
Phenol	2-Nitrophenol	2-Nitroaniline
Phenol-d <sub>6</sub> (surr)	N-Nitrosodi-n-butylamine	3-Nitroaniline
2-Picoline	N-Nitrosopiperidine	4-Nitroaniline
	1,2,4-Trichlorobenzene	4-Nitrophenol
		Pentachlorobenzene
		1,2,4,5-Tetrachlorobenzene
		2,3,4,6-Tetrachlorophenol
		2,4,6-Tribromophenol (surr)
		2,4,6-Trichlorophenol
		2,4,5-Trichlorophenol

(surr) = surrogate

TABLE 5 (continued)

Phenanthrene-d <sub>10</sub>	Chrysene-d <sub>12</sub>	Perylene-d <sub>12</sub>
4-Aminobiphenyl	Benzidine	Benzo(b)fluoranthene
Anthracene	Benzo(a)anthracene	Benzo(k)fluoranthene
4-Bromophenyl phenyl ether	Bis(2-ethylhexyl) phthalate	Benzo(g,h,i)perylene
Di-n-butyl phthalate	Butyl benzyl phthalate	Benzo(a)pyrene
4,6-Dinitro-2-methylphenol	Chrysene	Dibenz(a,j)acridine
Diphenylamine	3,3'-Dichlorobenzidine	Dibenz(a,h)anthracene
Fluoranthene	p-Dimethyl aminoazobenzene	7,12-Dimethylbenz(a) anthracene
Hexachlorobenzene	Pyrene	Di-n-octyl phthalate
N-Nitrosodiphenylamine	Terphenyl-d <sub>14</sub> (surr)	Indeno(1,2,3-cd) pyrene
Pentachlorophenol		3-Methylcholanthrene
Pentachloronitrobenzene	4	
Phenacetin Phenacetin	nect	ralah
Phenanthrene	Pool	

Pronamide

(surr) = surrogate

TABLE 6

EXAMPLE SINGLE LABORATORY PERFORMANCE DATA<sup>a</sup>

Compound	Test conc. (μg/L)	♣of 5 replicates (µg/L)	% Recovery of Avg.	
Acenaphthene	50	46.7	93.4	
Acenaphthylene	50	46.1	92.2	
Aniline	50	8.3	16.7	
Anthracene	50	48.4	96.8	
Benzoic acid	50	43.7	87.4	
Benz(a)anthracene	50	49.6	99.2	
Benzo(b)fluoranthene	50	49.8	99.6	
Benzo(k)fluoranthene	50	50.6	101	
Benzo(a)pyrene	50	47.7	95.5	
Benzo(g,h,i)perylene	50	52.6	105	
Benzyl alcohol	50	44.4	88.8	
Bis(2-chloroethyl) ether	50	44.2	88.4	
Bis(2-chloroethoxy)methane	50	46.6	93.1	
Bis(2-chloroisopropyl) ether	50	43.4	86.8	
Bis(2-ethylhexyl) phthalate	50	50.2	100	
4-Bromophenyl phenyl ether	50	48.6	97.2	21
Butyl benzyl phthalate	50	49.6	99.3	
Carbazole	50	52.1	104	
2-Chloroaniline	50	38.9	77.7	
4-Chloro-3-methylphenol	50 11	47.3	nc 94.6 nora	tion
2-Chloronaphthalene	50	45.3	90.8	
2-Chlorophenol	50	43.1	86.2	
4-Chlorophenyl phenyl ether	50	47.3	94.6	
Chrysene	50	50.3	101	
Dibenzofuran	50	47.4	94.7	
Dibenz(a,h)anthracene	50	51.6	103	
Di-n-butyl phthalate	50	50.5	101	
1,2-Dichlorobenzene	50	35.8	71.6	
1,3-Dichlorobenzene	50	33.3	66.7	
1,4-Dichlorobenzene	50	34.4	68.7	
3,3'-Dichlorobenzidine	50	32.0	64.0	
2,4-Dichlorophenol	50	47.4	94.8	
Diethyl phthalate	50	50.0	99.9	
Dimethyl phthalate	50	48.5	97.0	
2,4-Dimethylphenol	50	31.2	62.3	
4,6-Dinitro-2-methylphenol	50	57.6	115	
2,4-Dinitrophenol	50	58.7	117	
2,4-Dinitrotoluene	50	51.3	103	

TABLE 6 (continued)

Compound	Test conc. (μg/L)	<b>&amp;</b> of 5 replicates	% Recovery of Avg.	:
P	(1-3-)	(µg/L)	3	
2,6-Dinitrotoluene	50	50.2	100	
Di-n-octyl phthalate	50	51.1	102	
Fluoranthene	50	51.0	102	
Fluorene	50	48.5	97.0	
Hexachlorobenzene	50	49.0	97.9	
Hexachlorobutadiene	50	34.7	69.5	
Hexachlorocyclopentadiene	50	1.9	3.8	
Hexachloroethane	50	29.9	58.8	
Indeno(1,2,3-cd)pyrene	50	51.7	103	
Isophorone	50	47.1	94.3	
2-Methylnaphthalene	50	44.7	89.4	
2-Methylphenol	50	41.7	83.4	
4-Methylphenol	50	42.6	85.2	
Naphthalene	50	43.4	86.8	
2-Nitroaniline	50	48.4	96.7	
3-Nitroaniline	50	46.8	93.6	
4-Nitroaniline	50	56.1	112	
Nitrobenzene	50	47.1	94.1	1 a 10
2-Nitrophenol	50	47.3	94.6	
4-Nitrophenol	50	55.4	111	
N-Nitrosodiphenylamine	50	46.7	93.4	matian
N-Nitroso-di-propylamine	50	44.6	11 C 89.3	ration
Pentachlorophenol	50	56.9	114	
Phenanthrene	50	49.7	99.4	
Phenol	50	40.9	81.8	
Pyrene	50	49.2	98.4	
1,2,4-Trichlorobenzene	50	39.1	78.2	
2,4,5-Trichlorophenol	50	47.7	95.4	
2,4,6-Trichlorophenol	50	49.2	98.4	

Average recovery for five initial demonstration of capability measurements, in μg/L

<sup>&</sup>lt;sup>a</sup> Extraction using acidic pH only with a modified continuous liquid-liquid extractor with hydrophobic membrane according to Method 3520. These values are for guidance only. Appropriate derivation of acceptance criteria for similar extraction conditions may result in much different recovery ranges. See Method 8000 for information on developing and updating acceptance criteria for method performance.

TABLE 7

EXTRACTION EFFICIENCY AND AQUEOUS STABILITY RESULTS

	Percent Rec	overy, Day 0	Percent Rec	overy, Day 7
Compound	Mean	RSD	Mean	RSD
3-Amino-9-ethylcarbazole	80	8	73	3
4-Chloro-1,2-phenylenediamine	91	1	108	4
4-Chloro-1,3-phenylenediamine	84	3	70	3
1,2-Dibromo-3-chloropropane	97	2	98	5
Dinoseb	99	3	97	6
Parathion	100	2	103	4
4,4'-Methylenebis(N,N-dimethylaniline)	108	4	90	4
5-Nitro-o-toluidine	99	10	93	4
2-Picoline	80	4	83	4
Tetraethyl dithiopyrophosphate	92	7	70	Jal

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Data taken from Reference 6.

MEAN PERCENT RECOVERIES AND PERCENT RSD VALUES FOR SEMIVOLATILE ORGANIC FROM SPIKED CLAY SOIL AND TOPSOIL BY AUTOMATED SOXHLET EXTRACTION (METHOD 3541) WITH HEXANE-ACETONE (1:1)<sup>a</sup>

TABLE 8

·	Clay	Soil	Тор	soil
Compound	Mean Recovery	RSD	Mean Recovery	RSD
1,3-Dichlorobenzene	0		0	
1,2-Dichlorobenzene	0		0	
Nitrobenzene	0		0	
Benzal chloride	0		0	
Benzotrichloride	0		0	
4-Chloro-2-nitrotoluene	0		0	
Hexachlorocyclopentadiene	4.1	15	7.8	23
2,4-Dichloronitrobenzene	35.2	7.6	21.2	15
3,4-Dichloronitrobenzene	34.9	15	20.4	11
Pentachlorobenzene	13.7	7.3	14.8	13
2,3,4,5-Tetrachloronitrobenzene	55.9	6.7	50.4	6.0
Benefin	62.6	4.8	62.7	2.9
alpha-BHC	58.2	7.3	54.8	4.8
Hexachlorobenzene	26.9	13	25.1	5.7
delta-BHC	95.8	4.6	99.2	1.3
Heptachlor Properties of the Heptachlor	46.9	9.2	49.1	6.3
Aldrin	97.7	12	102	7.4
Isopropalin	102	4.3	105	2.3
Heptachlor epoxide	90.4	4.4	93.6	ra <sup>2.4</sup> or
trans-Chlordane	90.1	4.5	95.0	2.3
Endosulfan I	96.3	4.4	101	2.2
Dieldrin	129	4.7	104	1.9
2,5-Dichlorophenyl-4-nitrophenyl ether	110	4.1	112	2.1
Endrin	102	4.5	106	3.7
Endosulfan II	104	4.1	105	0.4
p,p'-DDT	134	2.1	111	2.0
2,3,6-Trichlorophenyl-4'-nitrophenyl ether	110	4.8	110	2.8
2,3,4-Trichlorophenyl-4'-nitrophenyl ether	112	4.4	112	3.3
Mirex	104	5.3	108	2.2

The operating conditions for the Soxtec apparatus were as follows: immersion time 45 min; extraction time 45 min; the sample size was 10 g; the spiking concentration was 500 ng/g, except for the surrogate compounds at 1000 ng/g, 2,5-Dichlorophenyl-4-nitrophenyl ether, 2,3,6-Trichlorophenyl-4-nitrophenyl ether, and 2,3,4-Trichlorophenyl-4-nitrophenyl ether at 1500 ng/g, Nitrobenzene at 2000 ng/g, and 1,3-Dichlorobenzene and 1,2-Dichlorobenzene at 5000 ng/g.

TABLE 9

SINGLE LABORATORY ACCURACY AND PRECISION DATA FOR THE EXTRACTION
OF SEMIVOLATILE ORGANICS FROM SPIKED CLAY BY
AUTOMATED SOXHLET (METHOD 3541)<sup>a</sup>

Compound	Mean Recovery	RSD	=
Phenol	47.8	5.6	_
Bis(2-chloroethyl)ether	25.4	13	
2-Chlorophenol	42.7	4.3	
Benzyl alcohol	55.9	7.2	
2-Methylphenol	17.6	6.6	
Bis(2-chloroisopropyl)ether	15.0	15	
4-Methylphenol	23.4	6.7	
N-Nitroso-di-n-propylamine	41.4	6.2	
Nitrobenzene	28.2	7.7	
Isophorone	56.1	4.2	
2-Nitrophenol	36.0	6.5	
2,4-Dimethylphenol	50.1	5.7	
Benzoic acid	40.6	7.7	
Bis(2-chloroethoxy)methane	44.1	3.0	
2,4-Dichlorophenol	55.6	4.6	
1,2,4-Trichlorobenzene	18.1	31	
Naphthalene	26.2	15	
4-Chloroaniline	55.7	11 <sup>12</sup> 1	ation
4-Chloro-3-methylphenol	65.1	5.1	ation
2-Methylnaphthalene	47.0	8.6	
Hexachlorocyclopentadiene	19.3	19	
2,4,6-Trichlorophenol	70.2	6.3	
2,4,5-Trichlorophenol	26.8	2.9	
2-Chloronaphthalene	61.2	6.0	
2-Nitroaniline	73.8	6.0	
Dimethyl phthalate	74.6	5.2	
Acenaphthylene	71.6	5.7	
3-Nitroaniline	77.6	5.3	
Acenaphthene	79.2	4.0	
2,4-Dinitrophenol	91.9	8.9	
4-Nitrophenol	62.9	16	
Dibenzofuran	82.1	5.9	
2,4-Dinitrotoluene	84.2	5.4	
2,6-Dinitrotoluene	68.3	5.8	_

Compound	Mean Recovery	RSD	=
Diethyl phthalate	74.9	5.4	_
4-Chlorophenyl-phenyl ether	67.2	3.2	
Fluorene	82.1	3.4	
4-Nitroaniline	79.0	7.9	
4,6-Dinitro-2-methylphenol	63.4	6.8	
N-Nitrosodiphenylamine	77.0	3.4	
4-Bromophenyl-phenyl ether	62.4	3.0	
Hexachlorobenzene	72.6	3.7	
Pentachlorophenol	62.7	6.1	
Phenanthrene	83.9	5.4	
Anthracene	96.3	3.9	
Di-n-butyl phthalate	78.3	40	
Fluoranthene	87.7	6.9	
Pyrene	102	0.8	
Butyl benzyl phthalate	66.3	5.2	
3,3'-Dichlorobenzidine	25.2	11	
Benzo(a)anthracene	73.4	3.8	
Bis(2-ethylhexyl) phthalate	77.2	4.8	
Chrysene	76.2	4.4	
Di-n-octyl phthalate	83.1	4.8	. 91 r
Benzo(b)fluoranthene	82.7	5.0	
Benzo(k)fluoranthene	71.7	4.1	
Benzo(a)pyrene	71.7	4.1	
Indeno(1,2,3-cd)pyrene	72.2	4.3	otion
Dibenz(a,h)anthracene	66.7	6.3	ation
Benzo(g,h,i)perylene	63.9	8.0	
1,2-Dichlorobenzene	0		
1,3-Dichlorobenzene	0		
1,4-Dichlorobenzene	0		
Hexachloroethane	0		
Hexachlorobutadiene	0		

Number of determinations was three. The operating conditions for the Soxtec apparatus were as follows: immersion time 45 min; extraction time 45 min; the sample size was 10 g clay soil; the spike concentration was 6 mg/kg per compound. The sample was allowed to equilibrate 1 hour after spiking.

Data taken from Reference 7.

TABLE 10

PRECISION AND BIAS VALUES FOR METHOD 3542<sup>1</sup>

Compound	Mean Recovery	Standard Deviation	% RSD
2-Fluorophenol	74.6	28.6	38.3
Phenol-d <sub>5</sub>	77.8	27.7	35.6
Nitrobenzene-d <sub>5</sub>	65.6	32.5	49.6
2-Fluorobiphenyl	75.9	30.3	39.9
2,4,6-Tribromophenol	67.0	34.0	50.7
Terphenyl-d <sub>14</sub>	78.6	32.4	41.3

<sup>&</sup>lt;sup>1</sup> The surrogate values shown in Table 10 represent mean recoveries for surrogates in all Method 0010 matrices in a field dynamic spiking study.



## SpectraLab

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TABLE 11

PRESSURIZED FLUID EXTRACTION (METHOD 3545) RECOVERY VALUES
AS PERCENT OF SOXTEC™

		Clay			Loam			Sand		Mean
Compound	Low	Mid	High	Low	Mid	High	Low	Mid	High	Rec.
Phenol	93.3	78.7	135.9	73.9	82.8	124.6	108.8	130.6	89.7	102.0
Bis(2-chloroethyl) ether	102.1	85.1	109.1	96.0	88.0	103.6	122.3	119.9	90.8	101.9
2-Chlorophenol	100.8	82.6	115.0	93.8	88.9	111.1	115.0	115.3	91.9	101.6
1,3-Dichlorobenzene	127.7	129.7	110.0	*364.2	129.9	119.0	*241.3	*163.7	107.1	120.6
1,4-Dichlorobenzene	127.9	127.0	110.5	*365.9	127.8	116.4	*309.6	*164.1	105.8	119.2
1,2-Dichlorobenzene	116.8	115.8	101.3	*159.2	113.4	105.5	*189.3	134.0	100.4	112.5
2-Methylphenol	98.9	82.1	119.7	87.6	89.4	111.0	133.2	128.0	92.1	104.7
Bis(2-chloroisopropyl)ether	109.4	71.5	108.0	81.8	81.0	88.6	118.1	148.3	94.8	100.2
o-Toluidine	100.0	89.7	117.2	100.0	*152.5	120.3	100.0	*199.5	102.7	110.3
N-Nitroso-di-n-propylamine	103.0	79.1	107.7	83.9	88.1	96.2	109.9	123.3	91.4	98.1
Hexachloroethane	97.1	125.1	111.0	*245.4	117.1	128.1	*566.7	147.9	103.7	118.6
Nitrobenzene	104.8	82.4	106.6	86.8	84.6	101.7	119.7	122.1	93.3	100.2
Isophorone	100.0	86.4	98.2	87.1	87.5	109.7	135.5	118.4	92.7	101.7
2,4-Dimethylphenol	100.0	104.5	140.0	100.0	114.4	123.1	100.0	*180.6	96.3	109.8
2-Nitrophenol	80.7	80.5	107.9	91.4	86.7	103.2	122.1	107.1	87.0	96.3
Bis(chloroethoxy)methane	94.4	80.6	94.7	86.5	84.4	99.6	130.6	110.7	93.2	97.2
2,4-Dichlorophenol	88.9	87.8	111.4	85.9	87.6	103.5	123.3	107.0	92.1	98.6
1,2 <mark>,4-T</mark> richl <mark>or</mark> obenzene	98.0	97.8	98.8	123.0	93.7	94.5	137.0	99.4	95.3	104.2
Naphthalene	101.7	97.2	123.6	113.2	102.9	129.5	*174.5	114.0	89.8	106.1
4-Chloroaniline	100.0	*150.2	*162.4	100.0	125.5	*263.6	100.0	*250.8	114.9	108.1
Hexachlorobutadiene	101.1	98.7	102.2	124.1	90.3	98.0	134.9	96.1	96.8	104.7
4-Chloro-3-methylphenol	90.4	80.2	114.7	79.0	85.2	109.8	131.6	116.2	90.1	99.7
2-Methylnaphthalene	93.2	89.9	94.6	104.1	92.2	105.9	146.2	99.1	93.3	102.1
Hexachlorocyclopentadiene	100.0	100.0	0.0	100.0	100.0	6.8	100.0	100.0	*238.3	75.8
2,4,6-Trichlorophenol	94.6	90.0	112.0	84.2	91.2	103.6	101.6	95.9	89.8	95.9
2,4,5-Trichlorophenol	84.4	91.9	109.6	96.1	80.7	103.6	108.9	83.9	87.9	94.1
2-Chloronaphthalene	100.0	91.3	93.6	97.6	93.4	98.3	106.8	93.0	92.0	96.2
2-Nitroaniline	90.0	83.4	97.4	71.3	88.4	89.9	112.1	113.3	87.7	92.6
2,6-Dinitrotoluene	83.1	90.6	91.6	86.4	90.6	90.3	104.3	84.7	90.9	90.3
Acenaphthylene	104.9	95.9	100.5	99.0	97.9	108.8	118.5	97.8	92.0	101.7
3-Nitroaniline	*224.0	115.6	97.6	100.0	111.8	107.8	0.0	111.7	99.0	92.9
Acenaphthene	102.1	92.6	97.6	97.2	96.9	104.4	114.2	92.0	89.0	98.4
4-Nitrophenol	0.0	93.2	121.5	18.1	87.1	116.6	69.1	90.5	84.5	75.6
2,4-Dinitrotoluene	73.9	91.9	100.2	84.7	93.8	98.9	100.9	84.3	87.3	90.7

TABLE 11 (continued)

		Clay			Loam			Sand		Mean
Compound	Low	Mid	High	Low	Mid	High	Low	Mid	High	Rec.
Dibenzofuran	89.5	91.7	109.3	98.5	92.2	111.4	113.8	92.7	90.4	98.8
4-Chlorophenyl phenyl ether	83.0	94.5	98.7	95.7	94.3	94.2	111.4	87.7	90.3	94.4
Fluorene	85.2	94.9	89.2	102.0	95.5	93.8	121.3	85.7	90.9	95.4
4-Nitroaniline	77.8	114.8	94.5	129.6	103.6	95.4	*154.1	89.3	87.5	99.1
N-Nitrosodiphenylamine	82.6	96.7	93.8	92.9	93.4	116.4	97.5	110.9	86.7	96.8
4-Bromophenyl phenyl ether	85.6	92.9	92.8	91.1	107.6	89.4	118.0	97.5	87.1	95.8
Hexachlorobenzene	95.4	91.7	92.3	95.4	93.6	83.7	106.8	94.3	90.0	93.7
Pentachlorophenol	68.2	85.9	107.7	53.2	89.8	88.1	96.6	59.8	81.3	81.2
Phenanthrene	92.1	93.7	93.3	100.0	97.8	113.3	124.4	101.0	89.9	100.6
Anthracene	101.6	95.0	93.5	92.5	101.8	118.4	123.0	94.5	90.6	101.2
Carbazole	94.4	99.3	96.6	105.5	96.7	111.4	115.7	83.2	88.9	99.1
Fluoranthene	109.9	101.4	94.3	111.6	96.6	109.6	123.2	85.4	92.7	102.7
Pyrene	106.5	105.8	107.6	116.7	90.7	127.5	103.4	95.5	93.2	105.2
3,3'-Dichlorobenzidine	100.0	*492.3	131.4	100.0	*217.6	*167.6	100.0	*748.8	100.0	116.5
Benzo(a)anthracene	98.1	107.0	98.4	119.3	98.6	104.0	105.0	93.4	89.3	101.5
Chrysene	100.0	108.5	100.2	116.8	93.0	117.0	106.7	93.6	90.2	102.9
Benzo(b)fluoranthene	106.6	109.9	75.6	121.7	100.7	93.9	106.9	81.9	93.6	99.0
Benzo(k)fluoranthene	102.4	105.2	88.4	125.5	99.4	95.1	144.7	89.2	78.1	103.1
Be <mark>nzo(</mark> a)py <mark>re</mark> ne	107.9	105.5	80.8	122.3	97.7	104.6	101.7	86.2	92.0	99.9
Indeno(1,2,3-cd)pyrene	95.1	105.7	93.8	126.0	105.2	90.4	133.6	82.6	91.9	102.7
Dibenz(a,h)anthracene	85.0	102.6	82.0	118.8	100.7	91.9	142.3	71.0	93.1	98.6
Benzo(g,h,i)perylene	98.0	0.0	81.2	0.0	33.6	78.6	128.7	83.0	94.2	66.4
Mean	95.1	94.3	101.0	95.5	96.5	104.1	113.0	100.9	92.5	

<sup>\*</sup> Values greater than 150% were not used to determine the averages, but the 0% values were used.

TABLE 12

SINGLE LABORATORY ACCURACY AND PRECISION FOR THE EXTRACTION OF PAHS FROM A CERTIFIED REFERENCE SEDIMENT EC-1, USING METHOD 3561 (SFE - SOLID TRAP)

Compound	Certified Value (mg/kg)	SFE Value <sup>a</sup> (mg/kg)	Percent of Certified Value	SFE RSD
Naphthalene	(27.9) <sup>b</sup>	41.3 ± 3.6	(148)	8.7
Acenaphthylene	(0.8)	$0.9 \pm 0.1$	(112)	11.1
Acenaphthene	(0.2)	0.2 ± 0.01	(100)	0.05
Fluorene	(15.3)	15.6 ± 1.8	(102)	11.5
Phenanthrene	15.8 ± 1.2	16.1 ± 1.8	102	11.2
Anthracene	(1.3)	1.1 ± 0.2	(88)	18.2
Fluoranthene	23.2 ± 2.0	24.1 ± 2.1	104	8.7
Pyrene	16.7 ± 2.0	17.2 ± 1.9	103	11.0
Benz(a)anthracene	8.7 ± 0.8	8.8 ± 1.0	101	11.4
Chrysene	(9.2)	7.9 ± 0.9	(86)	11.4
Benzo(b)fluoranthene	$7.9 \pm 0.9$	8.5 ± 1.1	108	12.9
Benzo(k)fluoranthene	4.4 ± 0.5	4.1 ± 0.5	91	12.2
Benzo(a)pyrene	5.3 ± 0.7	5.1 ± 0.6	96	11.8
Indeno(1,2,3-cd)pyrene	5.7 ± 0.6	5.2 ± 0.6	91	11.5
Benzo(g,h,i)perylene	$4.9 \pm 0.7$	$4.3 \pm 0.5$	88	11.6
Dibenz(a,h)anthracene	(1.3)	1.1 ± 0.2	(85)	18.2

<sup>&</sup>lt;sup>a</sup> Relative standard deviations for the SFE values are based on six replicate extractions.

Data are taken from Reference 10.

<sup>&</sup>lt;sup>b</sup> Values in parentheses were obtained from, or compared to, Soxhlet extraction results which were not certified.

TABLE 13

SINGLE LABORATORY ACCURACY AND PRECISION FOR THE EXTRACTION OF PAHS FROM A CERTIFIED REFERENCE SEDIMENT HS-3, USING METHOD 3561 (SFE - SOLID TRAP)

Compound	Certified Value (mg/kg)	SFE Value <sup>a</sup> (mg/kg)	Percent of Certified Value	SFE RSD
Naphthalene	$9.0 \pm 0.7$	$7.4 \pm 0.6$	82	8.1
Acenaphthylene	$0.3 \pm 0.1$	$0.4 \pm 0.1$	133	25.0
Acenaphthene	4.5 ± 1.5	$3.3 \pm 0.3$	73	9.0
Fluorene	13.6 ± 3.1	10.4 ± 1.3	77	12.5
Phenanthrene	85.0 ± 20.0	86.2 ± 9.5	101	11.0
Anthracene	$13.4 \pm 0.5$	12.1 ± 1.5	90	12.4
Fluoranthene	$60.0 \pm 9.0$	54.0 ± 6.1	90	11.3
Pyrene	$39.0 \pm 9.0$	$32.7 \pm 3.7$	84	11.3
Benz(a)anthracene	14.6 ± 2.0	12.1 ± 1.3	83	10.7
Chrysene	14.1 ± 2.0	12.0 ± 1.3	85	10.8
Benzo(b)fluoranthene	7.7 ± 1.2	$8.4 \pm 0.9$	109	10.7
Benzo(k)fluoranthene	2.8 ± 2.0	$3.2 \pm 0.5$	114	15.6
Benzo(a)pyrene	7.4 ± 3.6	$6.6 \pm 0.8$	CO 189	12.1
Indeno(1,2,3-cd)pyrene	$5.0 \pm 2.0$	$4.5 \pm 0.6$	90	13.3
Benzo(g,h,i)perylene	$5.4 \pm 1.3$	$4.4 \pm 0.6$	82	13.6
Dibenz(a,h)anthracene	$1.3 \pm 0.5$	1.1 ± 0.3	85	27.3

Relative standard deviations for the SFE values are based on three replicate extractions.
 Data are taken from Reference 10.

TABLE 14

SINGLE LABORATORY ACCURACY AND PRECISION FOR THE EXTRACTION OF PAHS FROM A CERTIFIED REFERENCE SOIL SRS103-100, USING METHOD 3561 (SFE - LIQUID TRAP)

	Certified Value	SFE Value <sup>a</sup>	Percent of	SFE
Compound	(mg/kg)	(mg/kg)	Certified Value	RSD
Naphthalene	32.4 ± 8.2	29.55	91	10.5
2-Methylnaphthalene	62.1 ± 11.5	76.13	122	2.0
Acenaphthene	632 ± 105	577.28	91	2.9
Dibenzofuran	$307 \pm 49$	302.25	98	4.1
Fluorene	492 ± 78	427.15	87	3.0
Phenanthrene	1618 ± 340	1278.03	79	3.4
Anthracene	422 ± 49	400.80	95	2.6
Fluoranthene	1280 ± 220	1019.13	80	4.5
Pyrene	1033 ± 285	911.82	88	3.1
Benz(a)anthracene	252 ± 8	225.50	89	4.8
Chrysene	297 ± 26	283.00	95	3.8
Benzo(a)pyrene	97.2 ± 17.1	58.28	60	6.5
Benzo(b)fluoranthene + Benzo(k)fluoranthene	153 ± 22	130.88	co1 <sup>86</sup> po1	10.7

Relative standard deviations for the SFE values are based on four replicate extractions.
 Data are taken from Reference 11.

TABLE 15

SINGLE LABORATORY RECOVERY DATA FOR SOLID-PHASE EXTRACTION (METHOD 3535) OF BASE/NEUTRAL/ACID EXTRACTABLES FROM SPIKED TCLP BUFFERS LOW SPIKE LEVEL

Spike	Buffer 1 (pH = 2	2.886)	Buffer 2 (pH =	4.937)
Level (µg/L)	Recovery (%)	RSD	Recovery (%)	RSD
3,750	63	10	63	9
1,500	55	6	77	4
1,000	82	10	100	5
250	65	3	56	4
65	89	4	101	5
65	98	5	95	6
100,000	83	10	85	5
100,000	86	8	85	3
100,000	<b>A</b> *C1	*	*	*
1,000	84	12	95	12
200,000	83	11	88	3
50,000	en 82 fic	9	CO1 <sup>78</sup> 2O1	ratio
	(μg/L) 3,750 1,500 1,000 250 65 65 100,000 100,000 1,000 200,000	Spike Level (μg/L)         Recovery (%)           3,750         63           1,500         55           1,000         82           250         65           65         89           65         98           100,000         83           100,000         *           1,000         84           200,000         83	Level (μg/L)       Recovery (%)       RSD         3,750       63       10         1,500       55       6         1,000       82       10         250       65       3         65       89       4         65       98       5         100,000       83       10         100,000       *       *         1,000       84       12         200,000       83       11	Spike Level (μg/L)         Recovery (%)         RSD         Recovery (%)           3,750         63         10         63           1,500         55         6         77           1,000         82         10         100           250         65         3         56           65         89         4         101           65         98         5         95           100,000         83         10         85           100,000         86         8         85           100,000         *         *         *           1,000         84         12         95           200,000         83         11         88

Results from seven replicate spiked buffer samples.

Data from Reference 12.

<sup>\*</sup> In this study, m-cresol and p-cresol co-eluted and were quantitated as a mixture of both isomers.

TABLE 16

SINGLE LABORATORY RECOVERY DATA FOR SOLID-PHASE EXTRACTION (METHOD 3535) OF BASE/NEUTRAL/ACID EXTRACTABLES FROM SPIKED TCLP BUFFERS HIGH SPIKE LEVEL

	Spike	Buffer 1 (pH = 2	2.886)	Buffer 2 (pH =	4.937)
Analyte	Level (µg/L)	Recovery (%)	RSD	Recovery (%)	RSD
1,4-Dichlorobenzene	15,000	63	10	63	9
Hexachloroethane	6,000	54	7	46	7
Nitrobenzene	4,000	81	4	81	13
Hexachlorobutadiene	1,000	81	5	70	11
2,4-Dinitrotoluene	260	99	8	98	3
Hexachlorobenzene	260	89	8	91	9
o-Cresol*	400,000	92	15	90	4
m-Cresol*	400,000	95	8	82	6
p-Cresol*	400,000	82	14	84	7
2,4,6-Trichlorophenol	4,000	93	12	104	12
2,4,5-Trichlorophenol	800,000	93	14	97	23
Pentachlorophenol	200,000	en&fic	9	CO1 <sup>73</sup> 2O1	atic

Results from seven replicate spiked buffer samples.

Data from Reference 12.

<sup>\*</sup> In this study, recoveries of these compounds were determined from triplicate spikes of the individual compounds into separate buffer solutions.

TABLE 17

RECOVERY DATA FROM THREE LABORATORIES FOR SOLID-PHASE EXTRACTION (METHOD 3535)
OF BASE/NEUTRAL/ACID EXTRACTABLES FROM SPIKED TCLP LEACHATES FROM SOIL SAMPLES

Buffer 1 pH = 2.886	, <u>-</u>		Lab 1			Lab 2			Lab 3	
Analyte	Spike Level (µg/L)*	%R	RSD	n	%R	RSD	n	%R	RSD	n
o-Cresol	200,000	86	8	7	35.3	0.7	3	7.6	6	3
m-Cresol**		77	8	7						
p-Cresol**		4 =						7.7	11	3
2,4,6-Trichlorophenol	2,000	106	6	7	96.3	3.9	3	44.8	5	3
2,4,5-Trichlorophenol	400,000	93	3	7	80.5	4.5	3	63.3	11	3
Pentachlorophenol	100,000	79	2	7	33.8	12.2	3	29.2	13	3
1,4-Dichlorobenzene	7,500	51	5	7	81.3	5.3	3	19.2	7	3
Hexachloroethane	3,000	50	5	7	66.2	2.1	3	12.6	11	3
Nitrobenzene	2,000	80	8	7	76.3	5.3	3	63.9	12	3
Hexachlorobutadiene	500	53	8	7	63.3	4.8	3	9.6	9	3
2,4-Dinitrotoluene	130	89	8	7	35.7	2.6	3	58.2	17	3
Hexachlorobenzene	130	84	21	7	92.3	1.6	3	71.7	9	3

(continued)

TABLE 17 (continued)

Buffer 2 pH = 4.937	_		Lab 1			Lab 2			Lab 3	
Analyte	Spike Level (µg/L)*	%R	RSD	n	%R	RSD	n	%R	RSD	n
o-Cresol	200,00	97	13	7	37.8	4.5	3	6.1	24	3
m-Cresol**		83	4	7				6.0	25	3
p-Cresol**										
2,4,6-Trichlorophenol	2,000	104	4	7	91.7	8.0	3	37.7	25	3
2,4,5-Trichlorophenol	400,000	94	4	7	85.2	0.4	3	64.4	10	3
Pen <mark>tachl</mark> orophenol	100,000	109	11	7	41.9	28.2	3	36.6	32	3
1,4-Dichlorobenzene	7,500	50	5	7	79.7	1.0	3	26.5	68	3
He <mark>xachl</mark> oro <mark>eth</mark> ane	3,000	51	3	7	64.9	2.0	3	20.3	90	3
Nitrobenzene	2,000	80	4	7	79.0	2.3	3	59.4	6	3
Hexachlorobutadiene	500	57	5	1-7-	60	3.3	3	16.6	107	3
2,4-Dinitrotoluene	130	86	6	7	38.5	5.2	3	62.2	6	3
Hexachlorobenzene	130	86	7	7	91.3	0.9	3	75.5	5	3

<sup>\* 250-</sup>mL aliquots of leachate were spiked. Lab 1 spiked at one-half these levels.

Data from Reference 12.

<sup>\*\*</sup> m-Cresol and p-Cresol coelute. Lab 1 and Lab 3 reported o-Cresol and the sum of — and p-Cresol. Lab 2 reported the sum of all three isomers of Cresol.

TABLE 18

SINGLE-LABORATORY PAH ANALYSIS DATA FROM A REAL SOIL CONTAMINATED WITH

CREOSOTE, USING METHOD 3546

(MICROWAVE EXTRACTION)

Compound	Concentration (µg/kg)	RSD (%)	REAC values (µg/kg)
Naphthalene	2,170	12.4	710,000
2-Methylnaphthalene	28,710	3.1	N/R
1-Methylnaphthalene	33,180	2.4	N/R
Biphenyl	13,440	6.0	N/R
2,6-Dimethylnaphthalene	52,990	3.8	N/R
Acenaphthylene	16,320	3.1	21,000
Acenaphthene	801,210	6.0	1,700,000
Fluorene	789,980	3.4	990,000
Phenanthrene	1,627,480	0.7	3,300,000
Anthracene	346,010	4.0	360,000
Benzo(a)anthracene	300,380	2.7	310,000
Fluoranthene	1,331,690	1.6	1,600,000
Pyrene	1,037,710	3.0	1,100,000
Chrys <mark>en</mark> e	293,200	3.4	320,000
Benzo(b)fluoranthene	152,000	3.8	140,000
Benzo(k)fluoranthene	127,740	3.6	130,000
Benzo(e)pyrene	87,610	3.9	N/R
Benzo(a)pyrene	128,330	3.9	110,000
Perylene	35,260	4.3	N/R
Indeno(123-cd)pyrene	63,900	5.0	25,000
Dibenz(a,h)anthracene	17,290	6.9	N/R
Benzo(ghi)perylene	42,720	6.9	20,000

<sup>\*</sup>n = 4

Soil samples obtained from US EPA Emergency Response Center archive bank through their contract laboratory REAC (Edison, NJ). The standard Soxhlet extraction procedures were performed by REAC three years earlier; this long storage period is believed to account for the low naphthalene recovery data in the present study

REAC data labeled N/R = not reported

TABLE 19
SINGLE-LABORATORY PAH RECOVERY DATA FROM HS-5 MARINE SEDIMENT
MATERIALS, USING METHOD 3546 (MICROWAVE EXTRACTION)

Compound	Certified Value (µg/kg)	Confidence Interval (µg/kg)	Recovery (%)	
Naphthalene	250	180 - 320	76	
Acenaphthylene	150	*	107	
Acenaphthene	230	130 - 330	61	
Fluorene	400	300 - 500	63	
Phenanthrene	5,200	4,200 - 6,200	72	
Anthracene	380	230 - 530	84	
Fluoranthene	8,400	5,800 - 10,000	81	
Pyrene	5,800	4,000 - 7,600	69	
Benzo(a)anthracene	2,900	1,700 - 4,100	53	1
Chrysene	2,800	1,900 - 3,700	76	
Benzo(b)fluoranthene	2,000	1,000 - 3,000	84	
Benzo(k)fluoranthene	1,000	600 - 1,400	137	
Benzo(a)pyrene	1,700	900 - 2,500	rp Gratioi	1
Indeno(123-cd) pyrene	1,300	600 - 2,000	63	
Dibenz(a,h)anthracene	200	100 - 300	125	
Benzo(ghi)perylene	1,300	1000 - 1600	64	

n = 3

The uncertainties represent 90% confidence intervals

<sup>\*</sup> values not certified

TABLE 20
SINGLE-LABORATORY PAH RECOVERY DATA FROM HS-4 MARINE SEDIMENT MATERIALS, USING METHOD 3546 (MICROWAVE EXTRACTION)

Compound	Certified Value	Confidence Interval	Recovery (%)	
Compound	(µg/kg)	(µg/kg)		
Naphthalene	150	*	54	
Acenaphthylene	150	*	82	
Acenaphthene	150	*	63	
Fluorene	150	*	81	
Phenanthrene	680	600 - 760	81	
Anthracene	140	70 - 210	108	
Fluoranthene	1250	1,150 - 1,350	84	
Pyrene	940	820 - 1,060	85	
Benzo(a)anthracene	530	470 - 580	78	
Chrysene	650	570 - 730	84	
Benzo(b)fluoranthene	700	550 - 850	84	
Benzo(k)fluoranthene	360	310 - 410	156	
Benzo(a)pyrene	650	570 - 730	73	7
Indeno(123-cd) pyrene	510	360 - 660	88	
Dibenz(a,h)anthracene	120	70 - 170	117	
Benzo(ghi)perylene	580	360 - 800	91	

n = 3

The uncertainties represent 90% confidence intervals

<sup>\*</sup> values not certified

TABLE 21

SINGLE-LABORATORY PAH RECOVERY DATA FROM HS-3 MARINE SEDIMENT MATERIALS, USING METHOD 3546 (MICROWAVE EXTRACTION)

Compound	Certified Value (µg/kg)	Confidence Interval (µg/kg)	Recovery (%)	
Naphthalene	9,000	8300 - 9,700	61	
Acenaphthylene	300	200 - 400	199	
Acenaphthene	4,500	3,000 - 6,000	80	
Fluorene	13,300	10,200 -16,400	58	
Phenanthrene	85,000	65000 -105,000	87	
Anthracene	13,400	12,900 -13,900	48	
Fluoranthene	60,000	51,000-69,000	91	
Pyrene	39,000	30,000-48,000	86	
Benzo(a)anthracene	14,600	12,600-16,600	78	
Chrysene	14,100	12,100-16,100	91	9
Benzo(b)fluoranthene	7,700	6,500-8,900	101	all
Benzo(k)fluoranthene	2,800	800-4,800	275	
Benzo(a)pyrene	7,400	3,000-7,000	74.	tion
Indeno(123-cd)pyrene	5,400	4,100-6,700	100	
Dibenz(a,h)anthracene	1,300	800-1,800	118	
Benzo(ghi)perylene	5,000	3,000-7,000	99	

n = 3

The uncertainties represent 90% confidence intervals

<sup>\*</sup> values not certified

TABLE 22
SINGLE-LABORATORY PAH RECOVERY DATA FROM SRM 1941 MARINE SEDIMENT,
USING METHOD 3546 (MICROWAVE EXTRACTION)

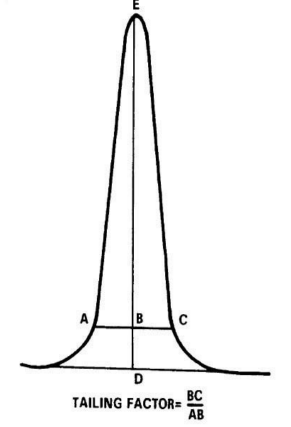
Compound	Certified Value (µg/kg)	Recovery (%)
Naphthalene	1010	97.4
Fluorene	100	100.0
Phenanthrene	490	102.0
Fluoranthene	980	116.7
Pyrene	810	97.3
Benz(a)anthracene	430	89.8
Chrysene	380	130.3
Benzo(b)fluoranthene	740	95.8
Benzo(k)fluoranthene	360	130.2
Benz(e)pyrene	550	81.0
Benzo(a)pyrene	630	76.0
Perylene	450	72.4
Indeno(123-cd)pyrene	500	126.0
Dibenz(a,h)anthracene	entificato	11 C O 1 78.7
Benz(ghi)perylene	530	85.2

n = 3

All RSDs < 10%

tion

### FIGURE 1 TAILING FACTOR CALCULATION



**ab** 

Example calculation: Peak Height = DE = 100 mm

10% Peak Height = BD = 10 mm

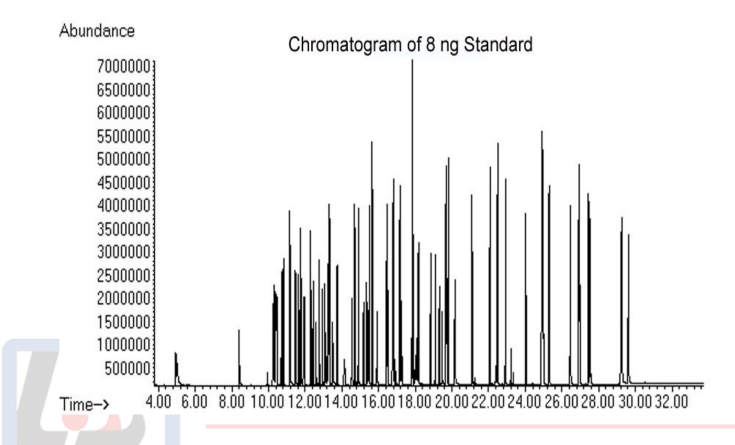
Peak Width at 10% Peak Height = AC = 23 mm

AB = 11 mm

BC = 12 mm

Therefore: Tailing Factor =  $\frac{12}{11}$  = 1.1

### FIGURE 2 GAS CHROMATOGRAM OF BASE/NEUTRAL AND ACID CALIBRATION STANDARD



Scientific Incorporation