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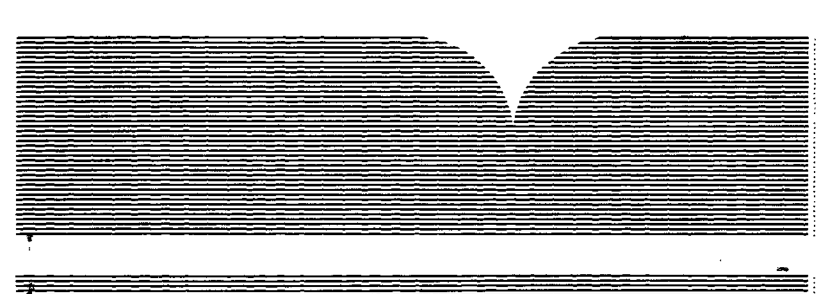
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Fewer semivolatile compounds of interest were found. Polychlorinated naphthalenes, polybrominated biphenyls, chlorinated phenols, and other compounds were specifically sought and not detected (limit of detection about 20-100 ng/mL milk). Polychlorinated biphenyls (PCBs) and DDE were found.

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ACQUISITION AND CHEMICAL ANALYSIS OF MOTHER'S MILK FOR SELECTED TOXIC SUBSTANCES

bу

Mitchell D. Erickson, Benjamin S. H. Harris, III, Edo D. Pellizzari, Kenneth B. Tomer, Richard D. Waddell and Donald A. Whitaker

> Contract No. 68-01-3849 Task 2

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December 1980

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ABSTRACT

Samples of mother's milk were collected from Bayonne, NJ; Jersey City, . NJ; Pittsburgh, PA; Baton Rouge, LA; and Charleston, WV, and analyzed for volatile (purgeables) and semivolatile (extractable) organics using glass capillary gas chromatography/mass spectrometry/computer. In the volatile fraction, 26 halogenated hydrocarbons, 17 aldehydes, 20 ketones, 11 alcohols, 2 acids, 3 ethers, 1 epoxide, 14 furans, 26 other oxygenated compounds, 4 sulfur-containing compounds, 7 nitrogen-containing compounds, 13 alkanes, 12 alkenes, 7 alkynes, 11 cyclic hydrocarbons, and 15 aromatics were found, including major peaks for hexanal, limonene, dichlorobenzene, and some esters. The levels of dichlorobenzene appeared to be significantly higher in the samples from Jersey City and Bayonne than in samples from other sites. Jersey City samples also appeared to have significantly higher levels of tetrachloroethylene. Charleston and Jersey City samples appeared to have significantly higher levels of chloroform; however, chloroform was observed in the blanks at about 20% of that in the samples. Due to the small sample size and lack of control over the solicitation of sample donors, the data cannot be used to extrapolate to the general population.

Fewer semivolatile compounds of interest were found. Polychlorinated naphthalenes, polybrominated biphenyls, chlorinated phenols, and other compounds were specifically sought and not detected (limit of detection about 20-100 ng/mL milk). Polychlorinated biphenyls (PCBs) and DDE were found.

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LIST OF ABBREVIATIONS AND SYMBOLS

ABBREVIATIONS

DDT -- 1,1-Bis(p-chlorophenyl)-2,2-trichloroethane

dpm -- Disintegrations per minute
 ECD -- Electon capture detection

GC -- Gas chromatography

MS -- Mass spectrometry (electron impact ionization)

NICIMS -- Negative ion chemical ionization mass spectrometry

OMB -- Office of Management and Budget

PBBs -- Polybrominated biphenyls
PCBs -- Polychlorinated biphenyls
PCF -- Participant Consent Form
PCN -- Polychlorinated Naphthalene
PLF -- Participant Listing Form

SQ -- Study Questionnaire

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SECTION 1 INTRODUCTION

BACKGROUND

It is becoming increasingly important to correlate ambient environmental pollutant levels with human body burden. Establishment of this correlation ("exposure assessment") may provide a link between pollution and health effects. This correlation is of interest for both scientific research and regulatory risk assessment.

Measurement of pollutant body burden levels generally requires invasive techniques (exceptions are breath and urine sampling) which are undesirable from the subjects' viewpoint. Some invasive techniques are generally regarded as acceptable (e.g., blood samples), while others are generally considered unacceptable from living donors (e.g. adipose tissue, internal organs, etc.). Mother's milk is an attractive medium for several reasons: (1) sample collection is reasonably straightforward; (2) milk contains a high amount of fat (about 3.5 percent, as shown in see Table 1), so fat-soluble pollutants such as DDT and polychlorinated biphenyls (PCBs) are likely to be found in higher concentrations in milk than in blood or urine; (3) large (50-100 mL) volumes are easily collected for analysis, increasing analytical reliability and detection limit; and (4) the population of nursing mothers is large relative to pathology samples such as adipose tissue. In addition, an assessment of pollutant concentrations in mother's milk may be used to predict the pollutant intake by the nursing infant.

The major disadvantages of mother's milk as a human-sampling medium relate to the sampling demography: only young-to-middle-aged females are nursing. Thus, any use of mother's milk in a probability-based sampling framework extrapolated to the general population would be fraught with difficulties, such as locating donors.

Table 1. COMPARISON BETWEEN HUMAN AND COW'S MILK $^{(1)}$

	Human Milk	Cow's Milk			
Water and solid content	Same in both; 87 to 87.5 percent	: is water			
Calories	Same in both; 20 calories per ou	ince			
Protein	l to 1.5 percent; 60 percent of this is lactalbumin and 40 percent casein	3.5 percent; 15 percent of this is lactalbumin and 85 percent casein			
Carbohydrate (in form of lactose)	6.5 to 7.5 percent	4.5 to 5.0 percent			
Fat(s)	Variable, but both have approximately 3.5 percent. (Differs qualitatively)				
	Contains more olein, which is is readily adsorbed	Contains more volatile fatty acids, which are irritat-ing to the gastric mucosa			
	Digestion of fat easy	Digestion of fat sometimes difficult			
Minerals	0.15 to 0.25 percent	0.7 to 0.75 percent. Con- tains more of all minerals with the exception of iron and copper			
	Iron content is low in both mill	ks, approximately:			
	1.5 mg/1	0.5 mg/1			
Vitamins	Varies with maternal intake				

Table 1 (cont'd.)

Parameter	Human Milk	Cow's Milk
Vitamin A	Relative large amounts in both	mi1ks
Vitamin B	Probably adequate in both milks	•
Vitamin C	More is found in human milk	
Thiamine	Higher content in cow's milk	
Riboflavin	Higher content in cow's milk	
Vitamin D	Relatively small amount in both	milks
Vitamin E	Satisfactory level in breath mi	.1k
Digestion	Cow's milk has a higher buffer	content and
	can therefore adsorb much more	
	than breast milk before it reac	hes the
	acidity necessary for digestion	. The large
	amount of casein on cow's milk	make large,
	tough curds in the stomach as c	
	the fine, easily broken down cu	rds of breast
	mi1k	

The purpose of this study was to measure levels of environmental pollutants in human milk by gas chromatography/mass spectrometry (GC/MS) and to evaluate the utility of using this body fluid in specific pollutant studies for populations in the vicinity of chemical manufacturing plants and/or industrial user facilities. All routes of exposure, <u>i.e.</u>, air, water, particulate, clothing and food were of interest. Mother's milk samples were acquired and analyzed for selected industrial chemicals. The chemicals of interest included: polychlorinated naphthalenes (PCNs), tetrachloroethylene, trichloroethane, dichloropropanes, benzene, polybrominated biphenyls (PBBs), chlorinated phenols, toluene, chlorinated benzenes, and chloroform.

Where possible, any other chemicals found in the extracts were identified and quantitated. The levels of selected organic compounds in mother's milk were investigated to assess the possibility of using this medium as an indicator of body burden for a wide range of organic compounds. For this feasibility study, no attempts were made to develop a statistically valid sample; sites were selected as having a high probability of pollutant detection and subjects were selected on a volunteer basis.

LITERATURE REVIEW

A review of the literature concerning pollutants in mother's milk was conducted. A computer search of MEDLARS II and ORBIT--III yielded 108 citations. These citations, plus personal contacts and manual searches yielded the data discussed below.

By far, most of the literature on environmental pollutants in mother's milk deals with chlorinated insecticides (e.g. DDT). PCBs have also been studied. Only a few references discuss the presence of other compounds in milk.

Table 2 lists the levels of pollutants found in mother's milk in the United States. Table 3 summarizes these findings. Table 4 summarizes pollutants found in mother's milk outside the United States. With the exception of one reference (27) regarding 1,2-dichloroethane exposure, all of the compounds found in mother's milk are semivolatile (extractable) halogenated compounds.

Table 2. LEVELS OF ORGANIC COMPOUNDS FOUND IN HUMAN MILK IN THE UNITED STATES

Compound	Sample Matrix	Mean (ppb)	Range (ppb)	Number of Determinations	Locations	References
β-BHC	Mi1k	0.5	T-10	57	AR, MS	2
	Milk		T-28	40	co	2 3
ү-ВНС	Milk Fat	83	30-270	53	PA	4
Total BHC	Mi1k	6.5	<0.1-20,2	14†	us	5
	Milk	7.7	n.d37.0	28	TΧ	6
	Milk	6.2	3.6-9.0	7	Houston, TX	6
p,p'-DDD	Milk	4.7	<0.1-14	14†	US	5
	Milk Fat	10.8	n.d30	53	PA	5 4
	Mi1k		T-5	40	co	3
o,p¹-DDE	Milk	1.0	<0.1-2.8	14†	US	5
p.pDDE	Milk	227	10-1720	57	AS, MS	2
	Milk	29	5.2-981	14†	us	2 5 5
	Mi1k	84.1	13.4-236	28	TX	5
	Milk	92.4	16.7-138	7	Houston, TX	6
	Milk Fat	1766	790-4350	53	PA	4
	Mi 1k		79-386	40	CO	3
DDE	Milk	194	74-314	30*	AZ	7
	Milk	60	20-90	4	Chicago, IL	8
	Milk	30	<10-140	5	Wenatche, WA	8
	Milk	30	_**	. 1**	Phoenix, AZ	8 8 8
	Mi1k	100	70-120		US	8

Ċ

Table 2 (cont'd.)

Compound	Sample Matrix	Mean (ppb)	Range (ppb)	Number of Determinations	Locations	References
o,p'-DDT	Milk	92	10-840	57	AR, MS	2
	Milk	25	<0.1-10.8	14†	บร	5 7
	Mi 1k	10	5-36	30*	ΛZ	7
	Milk		T-13	40	CO	3
p,p'-DDT	Milk	29	7.8-89	14 †	US	5
Mari Ann	Milk	114	9-383	30*	AZ	7
	Milk Fat	513	90-2120	53	PA	4
	Milk		7-109	40	CO	3
DDT (unspeci-	Milk	100	80-130	4	Chicago, IL	8
fied)	Milk	60	<10-220	5	Wenatche, WA	8
	Milk	60	_**	1	Phoenix, AZ	8
	Milk	70	50-90	**	บร	8
	Milk		10-110	40	co	3
	Mi1k	130	n.d770	32	DC	9
Total DDT Equiv.	Mi1k	334	20-2760	57	AR, MS	2
•	Milk	70.5	40.4-156	14	บร	
	Milk	100	SD=100	14	Long Island, NY	10
	Milk	170	SD=130	20	Rochester, NY	10
	Milk	180	SD=100	19	Chicago, IL	10
	Milk	220	SD=170	27	Lexington, KY	10
	Milk	170	SD=150	34	Nashville, TN	10
	Milk	150	SD=80	6	Memphis, TN	10
	Milk	180	SD=120	18	Los Angeles, CA	11
	Milk	447	59-1899	38	MS, AK	11
	Milk	75	15-133	14	Nashville, TN	11
	Milk	323	185~721	7	MS, AK	11
	Milk	130	n.d770	32	Washington, DC	9

Φ

Table 2 (cont'd.)

Compound	Sample Matrix	Mean (ppb)	Range (ppb)	Number of Determinations	Locations	References
Dieldrin	Mi1k	0.4	T-50	57	AR, MS	2
	Milk	6.2	2.9-14.6	14†	บร	2 5 5 5 3
	Milk	3.3	n.d,-21	28	TX	5
	Milk	7.5	1.9-21	7	Houston, TX	S
	Milk		T-11	40	co	3
ileptachlor	Mi1k	4	T-30	57	AR, MS	2
Epoxide	Milk	1.7	<0.1-4.4	14†	US	2 5 4 3
•	Milk Fat	160	40-460	53	PA	4
	Mi 1k		T-5	40	CO	3
t-Nonachior	Milk	1	T-10	57	AR, MS	2
Oxychlordane	Milk	5	T-20	57	AR, MS	2
PCBs	Milk	т	т	57	AR, MS	2
	Milk	∿10	<40-100	39	co	12
	Milk		40-100	40	co	3
Nicotine	Breast Fluid		n.d195	6	CA	13

NOTES: BIIC = benzenehexachloride (hexachlorocyclohexane)

DDD = 2,2-bis(chlorophenyl)-1,1-dichloroethane

DDE = 1,1-dichloro-2,2-bis(chlorophenyl)ethylene

DDT = 1,1,1-trichloro-2,2-bis(chlorophenyi)ethane

Total DDT equiv. = sum of all DDT-related peaks calculated as if all were DDT

PCBs = polychlorinated biphenyls. Quantitation generally based on comparison to an Aroclor mixture

T = trace

n.d. = not detected

SD = standard deviation

† = 5 women. Separate determinations make total of 14 samples.

* = 6 women. Separate samples makes total of 30 samples.

** = unspecified pool of donors in Denver and other US areas, no range given. Missing values indicate no data in original article

.

Table 2 (cont'd.)

NOTES (cont'd.): Mean values were taken from original citation where available; otherwise arithmetic mean was calculated, counting "ND" values as zero and "T" values as 0.5 times the lowest reported value.

Table 3. RANKING OF PESTICIDES AND PCBs BY REPORTED CONCENTRATIONS IN HUMAN MILK²

Compound	Weighted Mean Concentration (ppb) ^b	Number of Samples
DDEC	99	103
DDT ^C	94	100
PCBs ^C	<10	96
Oxychlordane	5	57
Dieldrin	4	92
DDD ^c	4	54
Heptachlor epoxide	4	71
BHCC	3	106
t-Nonachlor	1	57

⁸Whole milk only.

b Mean value calculated from a weighted mean of values in Table 2. Where either the mean or number of samples analyzed were unavailable, the data were excluded from calculation.

CAll isomers summed.

Table 4. LEVELS OF ORGANIC COMPOUNDS FOUND IN HUMAN MILK OUTSIDE THE UNITED STATES

Compound	Sample Matrix	Mean (ppb)	Range (ppb)	Number of Determinations	Number of Positives	Location	Date	Referenç
a-141C	#11k	0.58	0.1-1.9	50	17	Horway	1975	14
p-019C	MAIR	4.69	1.2-17.8	50	49	Norway	1975	14
	Mi 1k	70	NO-900	96	64	Germany	1971	15
	Milk	200	80-910	22	19	Vienaa	1973	16
	Milk	280	10-850	9	,	Rurel Austria	1973	16
	Mijk	4	1-16	50	42	Leiden (Neth.)	1969	17
	MEIR	2	ND-21	100	91	Canada	1975	13
y-MC	Milk	10.91	1.0-35.0	50	17	Hozway	197\$	14
	Hilk	•	Mb	96	0	Germany	1971	15
	Milk Pot	48	26-114	22	19	Vienna	1973	16
	Milk Fat	63	40-100	9	7	Rurel Austria	1973	16
	MI Ik	10.1		29		Israel	1975	18
	Milk	3	<1-35	147		Canada	1967-6	19
e-MC	Milk	1.14	0.3-3.2	50	34	Norway	1975	14
Total BIC	M11k	9.4	1.7-45.5	50	50	Нотнау	1975	14
	Milk	13	7-33	19	19	England	1964	20
P.P*-DD0	MIIK	9.9		29		Israel		1.6
000	Hi 1k	7	3-14	67	12	Australia	1970	21
<u>o,p</u> 1-002	Milk	18.02	1.6-43.6	50	30	Norway	1975	14
	Milk	9.5		29		Isteel	1975	18
₽.£,-006	Mik	65.10	0.9-113.2	50	50	Norway	1975	14
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	Milk	90	MD-600	96	95	Germany	1971	15
	MI 1k	21.7		29		îsrael	1975	I.O
	Milk	97	6-770	147		Canada	1967-8	19
	MLIK	30	***	50	50	Loidon (Meth.)	1969	17
	Hilk	35	17-68	6	6	New Brunswick	1973	23
	Milk	19	9-40	ě		Nova Scotia	1973	23
	361k	35	1-144	100	100	Canada	1975	24
	Milk	73	40-100	19	19	Eng land	1964	20

Table 4 (cont'd.)

Compound	Sample Matrix	Hean (ppb)	Range (ppb)	Humber of Determinations	Number of Positives	Location	Date	Reference
DOE	Milk	105	12-450	67	67	Australia	1970	21
	Milk Fac	3380	1930-7950	22	22	Vienna	1973	16
	Milk Fat	3920	3420-5970	9	9	Aurel Austria	1973	16
	Milk	61	15-112	26	26	W. Australia	1970-1	25
<u>e,p</u> *-DOT	Milk	18.52	1.6-120.9	so	49	Nosway	1975	14
	Mi ik	7.3		28		israel	1975	16
	Mi 1k	S	<1-31	147		Canada	1967-8	19
	Hl1k	3	KD-48	100	32	Canada	1975	24
2. <u>p</u> 1-001	Milk	17.89	2.3-130.3	50	50	Horway	1975	14
	Mijk		3-345	168	167	Portugal	1972	22
	Misk	90	10-250	96	95	Germany	1971	15
	Milk	7.3		29		larael	1975	38
	Milk	32	3-344	147		Consán	1967-8	18 19 17
	Ni 1k	16		50	50	Leiden (Meth.)	1969	17
	MiJk	13	6-30	6	6	New Brunswick	1973	23
	Milk	6	<2-11_	9	9	Nova Scotie	1973	23
	· Milk	6	?-21ª	100	100	Canada	1975	24
	Milk	45	20-75	19	19	England	1964	20
DOT	Mi ik	36	7-160	67	67	Australla	1970	21
•••	Milk Fat	1060	300-2680	22	21	Vienna	1973	16
	Milk Fat	1760	1030-2530	9	9	Rural Austria	1973	16
	MEIK	10	2-25	26	26	W. Austrelia	1970-1	25

Table 4 (cont'd.)

Compound	Sample Matrix	Hean (ppb)	Range (ppb)	Number of Determinations	Number of Positives	Location	Date	Reference
Total DDT	Mi lk	81.74	5.2-349.0	so	50	Horway	1975	14
Equiy.	Mi ik	386	<10-780	160	167	Portugal	1972	22
•	Milk Fet	1390	220-2580	19	19	Ontario	1975-4	26
	Milk Fat	3480	330-18800	34	34	Ontario	1971-2	26
	Hijk Fat	3460	110-11400	46	48	Ontario	1969-70	26
	Milk	320	30-870	96	96	Germany	1971	15
	Milk	141	15-580	67	67	Australia	1970	21
	Mi 1k	139	10-1020	147		Canada	1967-8	19
	Milk	78	19-137	26	26	W. Australia	1970-1	25
	Milk	378	3-5868	290	290	Gustens la	1973-4	27
	Mik	126	75-170	19	19	England	1964	24
Dieldrin	Milk	2.75	0.3-3.6	50	6	Norway	1975	14
	Milk	40	5-31	168	15	Portugal	1972	14
	Hilk Fat	48	<10-60	19		Ontario	1973-4	26
	Hilk Fat	90	<10-170	34		Ontario	1971-2	26
	Milk Fat	90	<10-250	46		Onterio	1969-70	25
	Milk	6	1-29	67	29	Austrolia	1970	21
	Mi 1k	7.0	=	29		Israel	1975	18
	MEIR	s ·	1-60	147		Canada	1967-8	19
	Milk	5	3-11	26	26	W. Australia	1970-1	25 .
	Milk	3	0.1-10.7	50	46	Leiden (Moth.)	1969	17
	Milk	Ž	ND-6	100	84	Canada	1975	24
	M1 1k	6	1-13	19	19	England	1969	20
Aldrin	MLIK	21.8		50	1	Norway	1975	14
Heptschlor	Milk	1.57	D.6-2.6	50	16	Norway	1975	14
Epox i de	Milk	9.1		29		!sreol	1975	14
• -	Mi 1k	3	<1-23	147		Canada	1967-8	19
	Mi 1k	1.2	0.3-3.5	50	50	Leldon (Woth.)	1969	17
	Milk	<u>t</u>	NO-3	100	69	Canada	1975	24

Table 4 (cont'd.)

Compound	Sample Metrix	Hean (ppb)	Rango (ppb)	Number of Determinations	Humber of Positives	Location	Date	Roferença
HCD	Milk	9.1	1.7-60.5	50	\$9	Horway	1975	14
	Milk Fat	100	NO-250	19		Ontario	1973-4	26
	Milk Fat	1240	260-4360	22	22	Vien na	1973	26 16 16 25 24
	Mitk	3670	2140-5110	9	9	Aurel Austria	1975	16
	Mi 1k	25	12-34	26	26	M. Austrašia	1970-1	25
	Mi Ik	2	MD-21	100	61	Canada	1975	24
PCB	Wijk Pat	1200	100-2500	19	19	Ontario	1973-4	26
·	Hilk Fat	1200	200-3000	34	34	Omtario	1971-2	26
	Hilk Fat	1000	700-12000	40	48	Ontario	1969-70	26
	Milk	90		96	64	Germany	1971	15
	Hilk Fat	1540	580-3780	22	22	Vienna	1973	16
	Hilk Fat	1290	950-3570	•	9	•	1973	16
	Milk	22	15-30	6	á	New Brunswick	1973	23
	MLLk	38	12-32	9	ġ	Nova Scotis	1973	23
	Milk	12	ND-64	100	100	Canada	1975	26 15 16 16 23 23 23
Oxychlordono	MELE	1	ND-2	300	77	Canada	1975	24
trans- Nonschler	Misk	1	ND-2	100	77	Cansda	1975	24
1,2-Bichlore- ethane	Hilk	6000		1	i			20

NOTES:

BHC = benzenehexachloride (hexachlorocyclohexane)

DDD = 2,2-bis(chlorophenyl)-1,1-dichloroethane

DDE = 1,1-dichloro-2,2-bis(chlorophenyl)ethylene

DDT = 1,1,1-trichloro-2,2-bis(chlorophenyl)ethane

Total DDT equiv. = sum of all DDT-related peaks calculated as if all were DDT.

PCB = polychlorinated biphenyls. Quantitation generally based on comparison to an Aroclor mixture.

HCB = hexachlorobenzene

ND = not detected.

Mean values were taken from original citation where available; otherwise arithmetic mean was calculated, counting "NO" values as zero and "T" values as 0.5 times the lowest reported value.

 ${}_{a}^{\mbox{\scriptsize Missing values}}$ indicate no data in original article. Lowest value not reported,

The literature shows that mother's milk often contains semivolatile chlorinated organic pollutants (pesticides). Presumably due to lack of analytical techniques and/or sensitivity, the presence of other pollutants has apparently not been investigated.

SECTION 2 SUMMARY AND CONCLUSIONS

The results show that sampling and analysis for organic compounds in mother's milk is feasible. The sample collection technique presented no significant problems. Analysis of the samples was generally satisfactory.

The use of purge and trap with gas chromatography/mass spectrometry/computer (GC/MS/COMP) analysis for volatile organics was successful, although the intrusion of contaminants during analysis presented problems with some compounds. The wide range of volatile compounds found includes common air and water pollutants and possible metabolites. Thus, it may be possible to use mother's milk as an indicator of body burden if a correlation between exposure and mother's milk concentration is established.

The extraction and GC/MS analysis for semivolatile organics was only marginally successful due to limited sensitivity (about 20-100 ppb milk). PCBs and DDE were the only halogenated semivolatiles found. The target semivolatile compounds (PCNs, PBBs, chlorinated phenols, and the higher chlorinated benzenes) were not present in quantities detectable by the survey techniques. The use of more sensitive (generally a factor of 100-1000) and selective methods [GC/electron capture detection (ECD), GC/negative ion chemical ionization mass spectrometry (NICIMS) or GC/single ion monitoring MS] may detect these compounds, but was outside the scope of this project.

SECTION 3

RECOMMENDATIONS

Further studies of the applicability of mother's milk as a matrix for assessing the human body burden of pollutants must directly compare human milk with the other available sample matrices. For example, comparison of the volatiles in breath, blood, urine, and mother's milk would determine which matrices are most suitable for measuring these compounds. It may also be advisable to use animal studies to determine the extent of environmental exposure-body burden correlation.

In addition, the effects of transport of pollutants to a newborn infant should be studied. Infants may be uniquely affected by some pollutants due to their small body weight and different metabolism relative to adults.

The measurement of semivolatile organics in mother's milk requires more sensitive techniques than those used in this study. For example, chlorinated compounds could best be detected using GC/ECD or GC/negative ion chemical ionization mass spectrometry and polynuclear aromatics by GC/photoionization detection.

Improvement in analytical methodology could occur at several points:

- (1) As discussed above, more sensitive, analytical procedures could be used for specific compound classes.
- (2) For volatile organics, background levels could be reduced with an on-line purge and trap/GC system.

Potential improvements in survey and sampling methodology include:

- (1) Addition of questions regarding length of nursing, age of infant, time since last nursing, etc.
- (2) Selection of participants according to a more statistically valid method (e.g. statistically random sampling).
- (3) Closer control over physical collection methodologies (e.g. all respondents gathered at one location).

The 5-month time lag in the study awaiting OMB clearance was seriously detrimental to the project. The personnel and apparatus used for the validation studies had to be reassembled once OMB clearance was obtained. Restarting a project following a long dormant period requires retraining analytical personnel (or training new personnel if original personnel have been reassigned to other research projects), recalibration of instruments, and assembling the necessary laboratory apparatus and supplies, all of which consume government resources. Reducing this time lag is extremely important for execution of programs involving human testing.

SECTION 4 SELECTION OF SAMPLING SITES

Five urban areas were chosen as sampling sites. Each of these cities is a high-probability area for the presence of one or more of the chemicals of interest in mother's milk. Since many of the compounds of interest are probably specific to certain industrial sites, the samples from the other sites were intended to serve as controls for the site-specific compounds. Other compounds are considered ubiquitous and their levels in milk was probably not related to local industrial activity. The rationale for selecting the five sampling sites is discussed below.

BRIDGEVILLE, PENNSYLVANIA

PCNs are manufactured by Koppers Company, Inc., of Pittsburgh, PA, at the Koppers Chemical and Coatings plant in Bridgeville, about 10 km SW of Pittsburgh. (29) Reported production levels were 7 million 1b in 1956 and 5 million 1b in 1972, (29) indicating a potential long-term, relatively constant, exposure level in the surrounding area. Results from environmental monitoring in the area immediately (< 1 km) surrounding the plant indicated higher levels of PCNs in air and soil than those found near five PCN user sites, as shown in Table 5. (30-34) Furthermore, fish and apple samples from the same area were found to contain PCNs, indicating a potential link to the human food chain.

In addition to PCNs, plants in the Bridgeville area have been reported to emit large quantities of phthalic anhydride particulate. (35) At this plant site, Koppers is reported to manufacture chlorinated naphthalenes, phthalic anhydride, maleic anhydride, and alkyd resins. (36)

Table 5. SUMMARY OF PCN CONCENTRATIONS FOUND NEAR MANUFACTURING AND USE SITES (32)

	 1 2 ·	Air, ng/m ³			Water, µg/L		Soil, µg/kg		
Site	Sampling Period	Low	High	Mean	Up- stream	Down- stream	Low	High	Mean
PCN manufacturer (Koppers)	1	25	450	150	0.2	1.4	130	2300	940
	2	120	2900	1400	a				
Capacitor manufacturing A	1	мрр	7.3	3.1	ND	ND	ND	7.3	2.0
	2	ND	3.9	1.2					
Capacitor manufacturing B	1	9.8	31	19	ND	0.6	ND	470	100
	2	9.8	33	17					

No water samples collected for period 2.

b_{Not detected.}

NORTHERN NEW JERSEY - STATEN ISLAND, NEW YORK, AREA (NNJ)

The Northern New Jersey (NNJ) area was selected as a sampling site on two bases: production of PBBs and general chemical industrial activity.

Three facilities are of interest (37) with respect to PBBs: White Chemical Co., E 22nd St., Bayonne, NJ; Marcor, Inc., Standard T. Chemical Co., subsidiary, 2500 Richmond Terrace, Staten Island, NY; and Hexcel Corp., Fine Organics Division, 880 Main St., Sayreville, NJ. White produced 45,000 kg of PBBs (specifically octabromobiphenyl and decabromobiphenyl) between 1970 and 1973. (38) Hexcel is reported (39) to have produced unspecified amounts of decabromobiphenyl [as well as to have produced or used decabromobiphenyl oxide, ethylene dichloride, and 1,2-bis(2,4,6-tribromophenoxy)ethane]. Standard T is thought to have been a PBB user up to about 1974. (39)

Results of environmental sampling in the area surrounding these three companies (40,41) indicated the presence of PBBs, especially the more highly brominated homologs, in sediment, water, soil, human hair, fish, turtle, and plant matter. The findings in human hair oil (18 total samples), which ranged from undetectable to 310 ppm, are especially relevant to this study, since they indicate that the PBB manufacturing in this area and the resultant environmental contamination has resulted in human exposure.

Northern New Jersey has a high concentration of chemical industries, (42) many of which use or produce halogenated hydrocarbons. The list of industries and locations are summarized below. Coastal Industries, Inc. (swimming pool chemicals), Diamond Shamrock (textile processing chemicals), Scientific Chemical Processing (chemical waste disposal) and Tenneco Chemicals (synthetic foam rubbers) are located in Carlstadt. Crompton & Knowles Corp. (dyes, colors and chemicals) are located in Fairlawn. Fisher Scientific (chemicals), Conoco Chemicals are in Saddle Brook. In Bayonne are CIBA-Geigy (dyes and intermediates) and ICI America (organics). In Jersey City are Hallinkrodt (analytical reagents) and Onya Chemical Co. (textile finish compounds, water repellants, germicides, and detergents). In Kearney are Standard Chlorine Chemical Co. (chlorobenzenes), Theobald Industries (bleaches), PPG Industries (paint) and Monsanto (industrial chemicals). In South Kearney is BASF-Wyandotte (dyestuffs and vinylidine chloride). In Newark are American Oil and Supply Co. (surfactants and chemicals), Celanese Plastics (plastics),

DuPont (pigments), Inmont (paint), Maas & Waldstein (paint), Otto B. May (dyes, surfactants), 3M (chemicals), Benjamin Moore (paint), Sherwin-Williams (paint) and Vulcan Materials (chloromethanes). In Elizabeth are Perk (chlorinated solvents) and Speciality Chemicals Division of Allied Chemical Corp. Linden Chlorine Products (chlorine) is in Linden. In Rahway are M & T Chemicals (speciality chemicals) and Merck and Co. (industrial chemicals). In Edison are Cary Page Chemicals (PVC compounds) and Mobile Chemical (paint). In Parlin, Hercules manufactures chloroform. In Passaic are Pantasote Co. of New York (PVC resin film), Stauffer (vinyl sheet and film) and United Wool Piece Dyeing and Finishing (dyes). In Patterson are several dye manufacturers. In Wayne are American Cyanamid (chemicals) and Owens Illinois (plastics). Many of these and other firms in NNJ undoubtedly manufacture or use compounds which are of interest to this study.

The levels of general organic pollutants in NNJ have been found to be high due to intense chemical manufacturing in the area. Environmental monitoring by RTI under separate contracts, (43-46) has found a wide variety of organic pollutants in this area. In addition, preliminary results from ground and surface water samples indicate measurable levels of a number of volatile halogenated hydrocarbons. (44,45) These data, summarized in Table 6, are indicative of environmental levels of organics in the NNJ area to which humans may be exposed and thus are indicative of the types of compounds anticipated in mother's milk. Under a separate research project, (45) the daily intake of some selected organics was roughly estimated. These estimates are given in Tables 7 and 8. Clearly there is ample exposure to pollutants which could potentially partition into milk.

The statistics for cancer in two counties of NNJ are very high. (58,59)
The overall rate for all malignant neoplasms is significantly above the national average. This cancer incidence in New Jersey has been partially linked to the chemical and allied industries located there. (60-64)

Northern New Jersey is a metropolitan area with a relatively static population, a well-established chemical industry, known environmental levels of organics (including PBBs) and abnormally high cancer rates. These factors make this area especially suited to this study of organics in mother's milk.

Table 6. PREVALENT HALOGENATED COMPOUNDS IN AMBIENT AIR AND WATER OF RAHWAY/WOODBRIDGE, BOUNDBROOK AND PASSAIC, NJ (44)

Medium	Occurrence									
	Ubiquitous	Mean Concentration ^a	Area Specific	Mean Concentration ^a						
	tetrachloroethylene trichloroethylene 1,1,1-trichloroethane 1,2-dichloroethane chloroform carbon tetrachloride o,m,p-dichlorobenzenes chlorobenzene	210,000 125,000 62,000 96,000 47,000 29,000 11,000 2,700	1,1,2-trichloroethane vinyl chloride 1,2-dichloroethylene 1,1,2,2-tetrachloroethane	9,000 1,200 1,000 750						
Water	dichlorobenzene trichloroethane chloroform trichloroethylene dichloroethane bromodichloroethane bromodichloromethane tetrachloroethylene dibromochloromethane	209 42 14 7 5 5 3.7 3.6 3.3	chloronitrobenzene methyl trichlorophenoxy acetate methyl dichlorophenoxy acetate bromopropylbenzene bromobenzene tetrachloroethane dichloroethylene	10.7 5 3.5 3 2.5 1.8						

^aConcentrations for air expressed in ng/m³ and for water in µg/L.

Table 7. ESTIMATED DAILY INTAKE OF SELECTED VOLATILE COMPOUNDS AND EXPECTED CONCENTRATIONS IN BLOOD IN NORTHERN NEW JERSEY (45)

Toxic Chemical	Air ^a (ng/day)	Water ^b (ng/day)	Food ^C (ng/day)	Total (ng/day)	Potential Blood Concentration ^d (ppb)
tetrachloroethylene	2,100,000	3,600	4,150	2,108,000	88
trichloroethylene	1,250,000	7,000	18,660	1,276,000	53
1,1,1-trichloroethane	620,000	42,000	5,290	667,000	28
1,2-dichloroethane	960,000	5,000		965,000	40
chloroform	470,000	14,500	14,280	499,000	21
carbon tetrachloride	290,000	1,000	12,070	303,000	13
dichlorobenzene	110,000	209,000		319,000	13
chlorobenzene	27,000	1,000		28,000	1.2
vinyl chloride	12,000			12,000	0.5
bromodichloromethane		3,700		3,700	0.2
benzene	7,500 ^e	300 ^f		7,800	0.2
total				6,188,200	258.2

^aFrom Ref. 44, calculated on basis of 10,000 L/24 h respiration rate.

bFrom Ref. 44, calculated on basis of 1 L/24 h intake.

CFrom Ref. 47, calculated from FDA standard diet (Ref. 48).

dExpected blood concentration is total daily intake divided by blood volume (8.000 mL) assuming 4 half-lives/day.

eFrom Ref. 49, 50.

from Ref. 50.

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Table 8. TOTAL DAILY INTAKE OF TARGET COMPOUNDS, PESTICIDES, PCBs, BaP AND METALS AND CONCENTRATIONS IN BLOOD IN NORTHERN NEW JERSEY $^{(45)}$

Toxic Chemicals	Air (ng/day)	Water (ng/day)	Food (ng/day)	Total (ng/day)	Expected Blood Concentration (ppb)
a-BHC	10	0	1,100	1,110	0.14
lindane	60		586	646	0.08
heptachlor	30	0	52	92	0.01
heptachlor epoxide		7	640	647	0.08
chlordane	20			20	∿0
DDE			3,500	3,500	0.44
DDT/DDD	70	0	2,500	2,570	0.32
HCB	50		73	123	0.02
PCBs	∿200	<60	388	648	0.08
Total Halogenated Compounds	440	 <67	8,849	9,356	1.16
benzo(a)pyrene	21	2	7,800	7,823	1.0
arsenic	2,800	<1,000	31,300	34,100	4.4
cadmium	50	<1,000	32,000	33,000	<10 ^a
lead	7,500	3,200	105,000	115,700	100-500 ^b

^aRef. 56.

b_{Ref. 57.}

Table 8 (cont'd.)

Sources:

Pesticides and PCBs in air	Ref. 51 (US) BaP in wa	iter Ref. 53 (World)
Pesticides in water	Ref. 44 (NJ) BaP in fo	ood Rough estimation
Pesticides and PCBs in food	l Ref. 48 (US)	(from Ref. 53 [World])
PCBs in water	Ref. 51 (US) Metals in	n air Ref. 54 (NJ)
BaP in air	Ref. 52 (US) Metals in	water Ref. 55 (NJ)
•	Metals in	n food Ref. 48 (N.E. NJ)

BATON ROUGE, LOUISIANA

Baton Rouge was selected on the basis of extensive organic chemical production (especially volatile halogenated hydrocarbons) as summarized in Table 9. (43) In addition, RTI has collected and analyzed ambient air samples from this area and established the presence of a number of compounds of interest in ambient air. (43) A summary of the levels of halogenated compounds found in water and air is presented in Table 10.

In addition to the industrial production in Baton Rouge, industries in Plaquemine (15 km SSW), St. Gabriel (20 km SSE) and Geismar (27 km SSE) may emit significant levels of chemicals which may contribute to the levels observed in mother's milk in Baton Rouge. These industries and their production are listed in Table 11. (36)

KANAWHA VALLEY, WEST VIRGINIA

Many manufacturers of organic chemicals are located in the Kanawha Valley, WV. DuPont, near Belle, WV, has a large chemical complex for the synthesis of substances such as methylmethacrylate, methylamines, ammonia, hydrogen cyanide, herbicides, and insecticides. In South Charleston are production and consumption plants (Union Carbide, and FMC). Plastics, PVC, antifreeze, chlorine, halogenated organics, carbon disulfide, peroxides, etc., are the predominant chemicals produced here. The major industrial facility in the town of Institute is Union Carbide, which also processes a broad spectrum of compounds, e.g., viscose rayon and phthalate esters. There is also a large-scale olefin processing complex and a rubber accelerator plant. A major terminal loading facility in South Charleston handles large quantities of a variety of organic compounds. Monsanto, FMC, Allied, and Fike have plants near Nitro for the production of antioxidants, rubber accelerators, industrial chemicals, and other materials. Several other chemical manufacturers, consumers, and transporters are located in the Kanawha Valley, some or all of which may contribute to the presence of organic materials in the ambient air or water and thus contribute to human exposure.

Previous RTI sampling (43,46,65,66) in the Kanawha Valley found a broad range of halogenated, ketone, aldehyde, ester, aromatic, and aliphatic compounds. Quantitative results included high values in air of 11,000 ng/m³

Table 9. POTENTIAL EMISSIONS FROM CHEMICAL INDUSTRY IN BATON ROUGE, LA^{a(43)}

Chemical	Total Production (mmlb/yr)	Raw Material	Company	
chlorodifluoromethane (101)	-	chloroform	ACC ^C	
dichlorodifluoromethane (12)	-	carbon tetrachloride	ACC	
dichlorotetrafluoroethane (114)	NA	perchloroethylene	ACC	
ethylene dichloride	1100	ethylene	ACC, EC	
polyethylene resin	460	ethylene	ACC	
trichlorofluoromethane (11)	-	-	ACC	
1,1,2-trich1oro-1,2,2-trifluoroethane (113)	NA	perchloroethylene	ACC	
vinyl chloride	480	ethylene dichloride	ACC, EC	
ethyl chloride	210	ethylene	EC	
methyl chloride	75	methanol	EC	
perchloroethylene	100	ethylene dichloride	EC	
tetraethyl lead	312	ethyl chloride	EC	
1,1,1-trichloroethane	40	1,1-dichloroethane	EC	
trichloroethylene	32	ethylene	EC	
PVC	144	-	EC	
benzene	440	petroleum	EXCC	
butadiene	428	ethane, etc.	EXCC, CR	
n-butyl alcohol	NA	-	EXCC	

Table 9 (cont'd.)

Chemical	Total Production (mmlb/yr)	Raw Material	Company	
decano1 ^c	NA	nonene	EXCC	
diisodecylphthalate	NA	phthalic anhydride, isodecanol	EXCC	
dodecene	100	propane/propylene	EXCC	
ethy lene	700	ethane, etc.	EXCC	
isobutylene	NΛ	petroleum	EXCC	
isodecano1 ^C	NΛ	nonene	EXCC	
isooctyl alcohoi ^c	NA	neptene	EXCC	
isoprene	10	ethylene by-product	EXCC	
isopropano1	680	propyleme	EXCC	
neopentanoic acid	5.5	isobutylene	EXCC	
nonene	300	propane/propylene	EXCC	
phthalic anhydride	90	o-xylene	EXCC	
propylene resin	320	ethylene	EXCC	
toluene	378	petroleum	EXCC, FGG	
ethylbenzene	900	benzene	FGC	
styrene	800	ethylbenzene	FGC	
vinyl toluene	NA ·	toluene, ethylene	FGC	

^aData provided by the Louisiana State Air Board.

bACC = Allied Chemical Corp., EC = Ethyl Corp., EXCC = Exxon Chem. Corp., FGC = Foster-Grant Co. Inc.

^cInvolves production of other alcohols also, C_6 , C_8 , C_9 , C_{10} , C_{13} , C_{16} .

NA = not available.

Table 10. PREVALENT HALOGENATED COMPOUNDS OCCURRING IN AMBIENT AIR AND WATER OF BATON ROUGE, GEISMAR AND PLAQUEMINE, $\mathsf{LA}^{(44)}$

		0cc	currence	
Medium	Medium Ubiquitous	Mean Concentration ^a	Area Specific	Mean Concentration
Air	chloroform	5,500	1,1,2-trichloroethane	632
	1,2-dichloroethane	1,656	1,2-dichloroethylene	472
	carbon tetrachloride	811	dichlorobutane	409
	1,1,1-trichloroethane	605	1,2-dichloropropane	306
	trichloroethylene	142	vinylidene chloride	78
	tetrachloroethylene	118	1,1,2,2-tetrachloroethane	70
	1,1-dichloroethane	86		
Water	trichloroethylene	96	bromobenzene	13
	chloroform	20	1,2-dichloroethylene	4
	trichloroethane	11	hexachloroethane	1.6
	dichloroethane	7. 7		
	carbon tetrachloride	7.1		
	dichlorobenzene	4.2		
	chlorodibromomethane	3.5		
	tetrachloroethylene	1.9		

^aConcentrations for air expressed in ng/m³ and for water in µg/L.



Table 11. POTENTIAL EMISSIONS FROM CHEMICAL INDUSTRY IN PLAQUEMINE, GEISMAR, AND ST. GABRIEL, LA (36)

City	Chemical	Annual Capacity (million pounds)	Company ⁸
Plaquemine	chloroform	ь	Dow
•	1,2-dichloropropane	10	H
	ethylene dichloride	1325	tt
	methyl chloride	150	Ħ
	methylene chloride	190	**
	tetrachloroethylene	150	11
	vinyl chloride	450	#t
Geismar	chloroform	46	VOM
	ethylene dichloride	330	11
	methylene chloride	80	11
	tetrachloroethylene	150	F F
	1,1,1-trichloroethane	65	**
	phosgene	55	BASF
	phosgene	125	RCC
	vinyl chloride	300	BOR
	vinyl chloride	300	MCJ
St. Gabriel	phosgene	NA	scc

aDow = Dow Chem. USA

VMC = Vulcan Materials Co.

BASF = BASF Wyandotte Corp.

RCC = Rubicon Chems., Inc.

BOR = Borden, Inc.

MCI = Monochem, Inc. SCC = Stauffer Chem Co., Agric. Chem. Div.

b200 million pounds combined capacity in Plaquemine and Freeport, TX plants.

for methylene chloride, 1500 ng/m³ for tetrachloroethylene, and 72,000 ng/m³ for benzene. Compounds identified in the air particulate fraction included long-chain alkanes, polycyclic aromatic hydrocarbons (PAH) from naphthalene through anthanthrene (or an isomer), alkyl-PAH derivatives, and nitrogen-containing heterocycles.

SECTION 5 SAMPLE COLLECTION

At each of the five sites, arrangements were made to work through clinical facilities to recruit a suitable panel of respondents. These facilities included the Bayonne Hospital in Bayonne, NJ; the Medical Center Hospital in Jersey City, NJ; Magee-Women's Hospital in Pittsburgh, PA; Charleston Area Medical Center in Charleston, WV; and the East Baton Rouge Parish Health Clinic in Baton Rouge, LA.

Advance arrangements were made through a contact person at each facility. This person was responsible for recruiting a professional member of the facility's staff to serve as the data collector. The data collector was usually a registered, licensed practical, or public health nurse associated with the facility.

Respondents were paid \$5 for their assistance in providing a milk sample and completing the survey questionnaire.

The data collection effort is discussed in the following sections.

Under the Federal Reports Act, clearance for the study of human subjects must be obtained from the Office of Management and Budget. This clearance was obtained on October 18, 1978. The OMB number is 158-578010. This study was approved with the understanding that: (1) the surveys were conducted as a pretest of the feasibility of information collection procedures; (2) the information collected will not be used to generalize to either local areas or the nation as a whole. These two caveats were invoked since the sample size was small and a nonprobability sampling method (subject selection) was used.

3.0

TRAINING

Before data collection began at a site, a training session was held to acquaint the facility contact person and data collector(s) with the survey. The session addressed the study objectives; use of the data collection instruments; administrative instructions; quality control procedures; and instructions for collecting, packing, and shipping milk samples to RTI. The training was conducted by an RTI survey specialist from the Survey Operations Center. A detailed manual and necessary field reporting forms were developed for use in these sessions. All training was conducted at the participating facility and lasted approximately 4 hours.

SURVEY INSTRUMENTS

Three data collection instruments (see Appendix A) were developed for use by the data collectors. The Participant Consent Form (PCF) was used to introduce the study, explain the study objectives and requirements of participation, present the confidentiality procedures, and obtain consent of participant. This form was signed by the respondent, who retained a copy for her files. The original was attached to the data collection instrument and a second copy was filed in the respondent's hospital record.

The Participant Listing Form (PLF) provided a means of assigning unique numbers to participants at each performance site. The data collector completed this form as each participant was solicited; the form was returned to RTI with the completed questionnaires when work at the site was finished.

The Study Questionnaire (SQ) was the primary data collection instrument. Information concerning participant demographic characteristics, residence information, health data, use of medications, and personal characteristics was obtained through this document. The SQ was administered after patients had been screened and prior to collection of the milk sample.

PARTICIPANT SCREENING

Potential participants (lactating women) were screened by the data collector to determine whether or not they met certain study criteria, which included:

- ability and desire to provide a milk sample of approximately 100 mL.
- permanent residence within the area of interest for at least the preceding 12 months, and
- no travel outside the area of interest for the seven days preceding sample collection.

After potential participants were screened, 10 women who met all the criteria for participation were asked to provide a milk sample and complete the SQ.

When an eligible person agreed to participate, her name was listed on the PLF and she was assigned a unique participant number. The data collector then read the information contained on the PCF to the participant while she followed along using a second copy. After answering questions or handling problems, the data collector asked the participant to sign the PCF prior to administration of the SQ.

The data collector then completed the SQ by asking the questions directly to the participant. Completion time averaged 15 minutes. An adhesive, computer-generated ID label was affixed to the SQ; a duplicate label was provided to be used for identifying the milk sample bottle.

Each participant was a self-respondent unless she was under 18 years of age, in which case the SQ could have been administered in whole or part to the parent or guardian, but in the participant's presence.

SAMPLE COLLECTION PROCEDURES

PLF, PCF, AND SQ COMPLETION PROCEDURES

After completion of the SQ, the data collector made the necessary arrangements for the participant to provide the milk sample. A collection bottle was taken from the shipping box and the adhesive ID label was affixed to the bottle. The milk was manually expressed directly into the bottle; no breast pumps or other devices were allowed. Immediately after the milk was collected, the bottle was capped and the sample frozen until all ten samples were collected and ready for shipment to RTI. A minimum of 60 mL (half-full bottle) was required for each sample. If insufficient milk was collected, the sample was discarded and an additional subject was added to the study.

SHIPPING PROCEDURES

Sample bottles were packed in the shipping container, cooled with dry ice, and sent directly to RTI via Federal Express.

SECTION 6 SAMPLE ANALYSIS METHODS

The milk samples were analyzed using gas chromatography/mass spectrometry/computer. Due to the broad range of volatilities, the samples were partitioned into two general classes of compounds: volatiles (e.g. benzene, chloroform) and semivolatiles (e.g. PCNs, PCBs, pesticides). The analytical protocols developed for the volatile and semivolatile components in mother's milk are reproduced in Appendices B and C, respectively. The experiments conducted which led to these protocols are discussed below.

DEVELOPMENT OF ANALYTICAL PROTOCOL FOR VOLATILES

The headspace purge technique was validated by determining the recovery of four model compounds from raw cow's milk samples. Compounds labeled with carbon-14 were chosen in order to examine both the amounts recovered on Tenax GC and the amounts remaining in purged samples.

Twelve 50 mL cow's milk samples were spiked with methanol solutions of the ¹⁴C-compounds. The analysis for each of the four model compounds was performed in triplicate. In addition, standards were prepared in triplicate by adding the appropriate amount of each compound in solution to a scintillation-counting vial containing 15 mL of Triton X/toluene/Omnifluor scintillation "cocktail." Milk samples were purged as described in Appendix B; Tenax cartridges were stored, and aliquots of the purged samples were retained for oxidation and counting.

Tenax cartridges were desorbed at 270°C and 30 mL/min $\rm N_2$ for 10 minutes into 15 mL of Triton X cocktail in tandem scintillation vials. The vials were capped and refrigerated until scintillation counting. An aliquot (1 mL) of each purged milk sample was oxidized in the Packard Tricarb Sample Oxidizer, which converted all carbon-containing compounds to carbon dioxide and water. The 14 C-carbon dioxide was collected in a trapping solution and

referenced to a quench correction curve. All standards, Tenax samples and oxidized milk samples were counted on a Packard Liquid Scintillation Counter with automatic standardization. Counting data was analyzed by computer to obtain the number of disintegrations per minute (dpm) for each vial. The percent recovery was calculated for each milk sample as shown below:

The second of the tandem scintillation vials contained <2 percent of the radioactivity in every case. The amounts of ¹⁴C compounds retained in the purged sample was calculated:

The data are tabulated in Table 12. The recoveries for the volatile chloroform and carbon tetrachloride were about 90 percent, as expected. The
less-volatile chlorobenzene and bromobenzene exhibited correspondingly
poorer recoveries. These compounds are generally considered only marginally
purgeable from water, so these results from milk are not surprising.

The methodology validation experiment indicated that the proposed method of analyzing human milk for volatile organic compounds was adequate. Sensitivity and detection limits were determined by the capabilities of the GC/MS/COMP system.

DEVELOPMENT OF ANALYTICAL PROTOCOL FOR SEMIVOLATILES

The extraction and cleanup method was validated using six model compounds (2,4-dichlorophenol, pentachlorobenzene, 1,2,3,4-tetrachloronaphthalene, 4,4'-dibromobiphenyl, 2,2',5,5'-tetrabromobiphenyl, and octachloronaphthalene) which were representative of the semivolatile (nonpurgeable) compounds of interest. The compounds were spiked into raw cow's milk at a level of about 1 µg/mL. Raw cow's milk was chosen as the closest readily available analog to mother's milk.

The results are presented in Table 13. The overall mean recovery was about 70 percent and the mean of the relative standard deviations was 22

Table 12. METHOD VALIDATION RECOVERY OF SELECTED VOLATILE STANDARDS FROM MILK

Compound ^a	b.p. (°C)	Percent Recovered	Percent _b Retained	Percent Accounted for
14 _{C-chloroform}	62	88 <u>+</u> 5	6 <u>+</u> 0.3	94 + 2
¹⁴ C-carbon tetrachloride	76	88 <u>+</u> 6	3 <u>+</u> 3	91 <u>+</u> 3
¹⁴ C-chlorobenzene	132	63 <u>+</u> 2	26 <u>+</u> 3	89 <u>+</u> 1
14 _{C-bromobenzene}	156	35 <u>+</u> 3	51 <u>+</u> 13	86 <u>+</u> 10

 $^{^{\}mathbf{a}}$ 80,000-94,000 dpm added to each sample.

bMean + standard deviation of three replicates.

CSum of percent recovered and percent retained.

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Table 13. METHOD VALIDATION RECOVERY OF SEMIVOLATILE COMPOUNDS SPIKED INTO RAW CON'S MILK

Compound	mp (°C)	bp (°C)	Concentration in Milk (ng/mL)	Mean Recovery (%)	Standard Deviation (%)	Relative Standard Deviation (%)
2,4-Dichlorophenol	45	207	1.12	59	12	20
Pentachlorobenzene	85	277	1.24	76	19	24
1,2,3,4-Tetrachloronaphtha- lene	197		1.37	59	15	25
4,4'-Dibromobiphenyl	164	357	1.04	58	19	33
2,2',5,5'-Tetrabromobiphenyl			0.93	94 ^C	10	11
Octachloronaphthalene	198	441	1.08	78 ^C	14	17

^aSeven replicates.

^bStandard deviation divided by mean multiplied by 100.

^cSix replicates.

percent. These results indicated that refinements in the method should be considered prior to a large-scale study.

Two methods were available for removing fat and other nonvolatile components of the milk extract: Florisil column chromatography and gel permeation chromatography (GPC). Evaluation of the two techniques indicated that the Florisil method was more suitable to this project. The Florisil method was faster and had greater sample capacity than the GPC. In addition, the GPC procedure required the use of a pumping system, UV detector, and expensive, fragile GPC columns. Initial tests with both methods revealed interference problems, although those with GPC were more severe. Using GPC, decabromobiphenyl and hexabromobiphenyl eluted with the fat peak. This was judged totally unsatisfactory. Using Florisil, some fat eluted in the fraction with the compounds of interest, but repetition of the procedure yielded samples sufficiently clean for analysis.

DEPARTURES FROM THE ANALYTICAL PROTOCOLS

Emulsions

The formation of an emulsion during the toluene-acetone extraction of semivolatiles (step 6, Appendix C) was an area of concern. Approximately 80 percent of the time an emulsion occurred. To eliminate this, three approaches were taken with reasonable success. The first was to avoid the emulsion formation by swirling rather than shaking the toluene and acetone extracts. The second approach was to break the emulsion by adding Na₂SO₄ and waiting. Both the amounts of Na₂SO₄ and the time required varied. In severe cases emulsions were broken by filtering through glass wool wetted with toluene.

Lipid Removal Using Florisil

Problems were also encountered during the Florisil cleanup. Some samples had a tendency to solidify while concentrating the ether/pentane eluate, apparently due to abnormally high fat content. This usually occurred when the sample volume reached 1-3 mL. The samples to which this happened were diluted with pentane and eluted through another Florisil column. The Florisil cleanup was repeated until the samples remained liquid at small (<1.0 mL) volumes. Three cleanups was the maximum required for any sample.

GC/MS ANALYSIS PROCEDURES

Samples were analyzed by gas chromatography/mass spectrometry using an LKB 2091 EI/CI GC/MS. Operating conditions for the analysis of purgeables is given in Table 14 and the operating conditions for the extractables is given in Table 15. Analysis of the purgeables involved the use of the desorption apparatus described in Appendix B.

Quantitation of the unknowns was accomplished using relative molar responses (RMRs) as discussed in Appendices B and C. The RMRs were calculated from replicate determinations of known amounts of standards and analytes.

Qualitative Analysis

Initial identification of compounds by GC/MS involved comparisons of unknown spectra with data compiled in the Eight Peak Index of Mass Spectra (67). If the peaks present in the unknown spectra clearly matched the peaks of the standard compound in the tables and the intensities were about the same, then a positive identification was usually made. If peak intensities of unknowns varied from those of the standards, and there were isomers of the compounds that were not listed in the Eight Peak Index, then the compound was listed as an "isomer."

When the background peaks interfered with the spectrum of an unknown to an extent that made identification uncertain, the compound identification was labeled as "tentative" (tent.). If no standard spectra similar to those of the unknowns appeared in the mass spectral references, but fragments characteristic of a certain class of compounds were identified, tentative identifications were made on the basis of the characteristic fragments and apparent molecular weights. These identifications were also labeled "tent". Usually tentative identifications involved alkyl derivatives or homologs of classes of compounds that were positively identified in the same sample.

Positive identifications, as well as some tentative identifications, often required more detailed investigations of standard spectra in the Registry of Mass Spectral Data (68) or standard spectra found in other literature such as scientific journals. The Registry of Mass Spectral Data presents data in the form of histograms rather than as a list of peaks and their intensities. This type of format allowed more subtle differences in mass spectra to be considered when several similar standard spectra in the

Table 14. OPERATING CONDITIONS FOR GC/MS ANALYSIS OF PURGEABLES

Instrument	LKB 2091
Column	80m - SE-30 WCOT Capillary Column
Flow	1.7 mL/mdn He
Desorption Temperature	270°C
Desorption Time	8 min
Desorption Flow	15 mL/min He
Column Temperature	30°C for 2 min programmed to 240°C at 4°C/min
Scan Range	5 → 490 Dalton
Scan Speed	0 + 670 in 2 sec
Scan Cycle	1.7 sec
Injector Temperature	250°C
Accelerating Voltage	3500 V
Ionizing Energy	70 eV
Trap Current	50 μ A
Source Temperature	210°C

Table 15. OPERATING CONDITIONS FOR THE GC/MS ANALYSIS OF SEMIVOLATILES

Instrument	LKB 2091
GC Column	25m SE-52 WCOT capillary column
Flow	1.5 mL/min with 15:1 split
Column Temperature	80°C for 3 min then 8°C/min to 265°C
Scan Range	5 + 530 Dalton
Scan Speed	2 sec 0 + 670 Dalton
Scan Cycle	2.4 sec
Injector Temperature	240°C
Accelerating Voltage	3500 V
Ionizing Energy	70 eV
Trap Current	50 µA
Source Temperature	210°C

Eight Peak Index appeared to represent possible candidates for unknown identifications.

A large number of sample components remained unidentified. These unidentified components were labeled "unknown."

In order to quantify the degree of certainty with which a compound has been identified, a "level" heirarchy has been established. The compound identification criteria are listed below:

- Level I Computer Interpretation. The raw data generated from the analysis of samples are subjected to computerized deconvolution/library search. Compounds identified using this approach have the lowest level of confidence. In general Level I is reserved for only those cases where compound verification is the primary intent of the qualitative analysis.
- Level II Manual Interpretation. The plotted mass spectra are manually interpreted and compared to those spectra compiled in a data compendium by a skilled interpreter. In general a minimum of five masses and intensities (±5 percent) should match between the unknown and the library spectrum. This level does not utilize any further information such as retention time since the authentic compound may not be available for establishing retention times.
- Level III Manual Interpretation Plus Retention Time/Boiling Point
 of Compound. In addition to the effort described under
 Level II, the retention time of the compound is compared to
 the retention time that has been derived from previous chromatographic analysis. Also the boiling point of the identified
 component is compared to the boiling points of other compounds
 in the near vicinity of the one in question when a capillary
 coated with a nonpolar phase has been used.
- Level IV Manual Interpretation Plus Retention Time of Authentic Compounds.

 Under this Level, the authentic compound has been chromatographed on the same capillary column using identical operating conditions and the mass spectrum of the authentic compound is compared to that of the unknown.
- Level V Level IV Plus Independent Confirmation Techniques. This Level utilizes other physical methods of analysis such as GC/Fourier transform infrared spectrometry, GC/high resolution mass spectrometry, or nmr analysis. This Level constitutes the highest degree of confidence in the identification of organic compounds.

Unless otherwise stated, all identifications in this report were Level II.

SECTION 7 RESULTS

VOLATILES

All 42 of the purged samples were analyzed by thermal desorption/GC/MS. The mass spectra from selected samples were interpreted manually to determine which compounds should be quantitated. From these data, selected compounds were quantitated in all samples. All data were stored on magnetic tape for subsequent processing and are routinely archived for at least 5 years.

Qualitative Identifications

Eight samples were interpreted. The results are presented in Appendix D. Samples were selected according to the following criteria. At least two samples were required from each collection site (Jersey City and Bayonne, NJ, were counted as two separate sites). The total ion current chromatograms were inspected and the samples with the greatest number of peaks or those containing very intense unique peaks (not observed in other samples) were selected. For those samples selected, all of the mass spectra were printed and interpreted manually by experienced spectroscopists.

Table 16 summarizes the compounds found and their frequency of occurrence. It is interesting to note that some compounds (e.g. 1,1,1-trichloroethane and hydrocarbons) are common air pollutants, others (e.g., dibromochloromethane) are common water pollutants, others (dimethyldisulfide, furans, aldehydes) appear to be metabolites, others (chlorofluorocarbons, siloxanes) are known background interferents, and others (iodopentane) are of unknown source.

Quantitation

Based upon the qualitative identifications summarized above, nine compounds were selected for quantitation in all of the samples. The results for four compounds are summarized in Table 17. As discussed below, the

Table 16. SUMMARY OF QUALITATIVE IDENTIFICATIONS OF VOLATILE COMPOUNDS IN MOTHER'S MILK

		•	Sam	ple Numi	per			
Compound	1081	1040	1107	1115	2048	2071	3053	3111
Halogenated Compounds								
chlorodif.uoromethane	-	•	+	-	-	-	-	-
chlorotrifluoromethane	+	+	-	-	+	-	+	-
dichlorodifluoromethane	•	-	+	-	-	+	-	-
chloromethane	-	-	-	+	-	-	+	-
chloroethane	-	-	+	-	-	+	-	-
trichlorofluoromethane	+	+	+	+	+	+	-	+
dichloroethylene	-	+	-	-		-	-	-
Freon 113	+	+	+	+	+	+	+	+
methylene chloride	+	+	+	+	+	+	+	+
chloroform	+	+	+	+	+	+	-	+
1,1,1-trichloroethane	+	+	+	+	+	+	+	+
carbon tetrachloride	-	+	+	+	-	+	_	+
trichloroethylene	+	+	+	+	+	+	+	+
chloropentane	+	+	-	-	-	-	•	_
dibromochloromethane	-	*	-	-	-	+	_	_
tetrachloroethylene	+	+	+	+	+	+	-	+
dichloropropene	-	_	_	+	_	_	_	-
chlorobenzene	4	-	•	+	+	•	-	_
chlorohexane		.		_		_	_	_
iodopentane	_	_	_	1	_	_	_	_
3-methyl-1-iodobutane	_	_	_	_	_	_	_	_
chloroethylbenzene	-		•	+	_		_	_
dibromodichloromethane	-	_	-	+	-	-	•	•
dichlorobenzene	-	-	-	·	-	_	-	-
chlorodecane	*	+	+	+	+	*	+	+
	•	•	-	•	-	-	-	-
trichlorobenzene	•	-	-	-	-	*	•	-
Aldehydes								
acetaldehyde	+	-	+	-	+	+	-	-
methylpropanal	-	+	+	-	-	•	-	-
n-butanal	+	-	+	+	-	+	+	+
methylbutanal	-	+	-	+	-	-	-	-
crotonaldehyde	-	-	-	+	-	-	•	-
n-pentanal	+	-	+	+	+	+	+	+
n-hexanal	+	+	+	+	+	+	+	+
f uraldehyde	-	+	-	+	-	-	+	-
n-heptanal	+	+	+	+	+	+	+	-
benzaldehyde	+	+	+	+	+	+	+	+
n-octanal	+	-	+	+	-	-	+	-
phenyl acetaldehyde	-	-	_	+	-	-	-	-

Table 16 (cont'd.)

			Sa	mple Nu	mber			
Compound	1081	1040	1107	1115	2048	2071	3053	311
n-nonanal	+	+	+	+	+	-	+	
methyl furaldehyde	-	-	•	-	•	•	+	-
n-decanal	-	-	-	+	-	•	+	-
n-undecanal	-	-	-	+	-	-	+	-
n-dodecanal	-	-	-	-		-	+	-
(etones								
acetone	+	+	+	+	+	+	+	+
methyl ethyl ketone	+	+	•	-	+	+	+	-
methyl isopropyl ketone	-	-	-	+	-	+	-	-
methyl vinyl ketone	-	-	+	-	-	-	-	-
ethyl vinyl ketone	+	+	+	+	-	-	+	-
2-pentanone	+	+	+	+	-	-	-	-
methyl pentanone	-	-	+	+	-	-	-	~
methyl hydrofuranone	-	-	•	+	-	-	-	-
2-methy1-3-hexanone	-	-	-	+	-	-	-	-
4-heptanone	-	•	+	-	•	-	-	-
3-hept anone	+	-	+	-	+	+	-	-
2-heptanone	+	+	+	+	+	+	-	-
methyl heptanone	-	•.	-	+	-	+	-	•
furyl methyl ketone	-	-	-	+	•	-	-	-
octanone	+	-	-	+	-	-	-	-
acetophenone	+	+	+	+	+	+	+	+
2-nonanone	+	-	+		-	+	•	•
2-decanone	-	-	-	+	-	-	-	-
alkylated lactone	-	-	- +	+	-	-	-	-
phthalide	-	-	*	-	-	-	-	-
ther Oxygenated Isomers								
C ₄ H ₆ O	-	-	-	-	-	-	+	-
C ₄ H ₈ O	-	-	-	•	-	+	+	-
C ₅ H ₁₀ O	-	-	+	•	+	+	+	+
CeHaO	-	-	-	•	-	•	+	-
C ₆ H ₁₀ O C ₄ H ₆ O ₂	-	-	+	-	•	-	+	-
C ₄ H ₆ O ₂	+	-	-	-	-	*	-	-
C6H12U	+	-	-	-	+	~	-	-
C7H12O	-	•	+	+	-	•	+	+
C7H100	-	-	+	. +	-	-	•	-
C ₂ H ₁ H ₂ D	-	-	-	-	+	-	•	•
CéHéO2	-	-	-	-	-	•	*	•
C H1402	-	-	+	-	-	•	-	•
С _в И ₁₆ О	+	-	-	-	+	-	•	•
C7H 02	-	-	*	*	-	-	+	•
C7H1002	-	•	-	•	-	-	₹	~

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Table 16 (cont'd.)

	180	Sample Number ^b							
		···	Sa	mple Nu	mber -				
Compound	1081	1040	1107	1115	2048	2071	3053	3111	
Other Oxygenated Isomers (continued)									
C9H18O C8H6O2	+	-	-	-	+	-	+	-	
C ₁₀ H ₁₂ O	-	-	+	-	-	-	-	-	
C ₁₀ H ₁₄ O	-	-	-	-	+	-	-	+	
C ₁₀ H ₁₆ O	-	-	-	+	-	-	+	-	
C ₁₀ H ₁₈ O	-	-	+	+	-	+	-	-	
C ₁₀ H ₂₀ O	+	-	-	-	-	-	+	-	
C ₁₀ H ₂₂ O	-	-	-	-	+	-	-	-	
c ₉ H ₈ O ₂	-	-	-	-	_	-	+	-	
C ₁₁ H ₂₀ O	-	-	-	-	_	-	+	-	
c ₁₀ H ₁₀ o ₂	-	-	-	-	-	-	+	-	
Alcohols									
methanol	-	₩-	_	•-	-	+	_	-	
isopropanol	+	+	+	+	+	+	+	+	
2-methyl-2-propanol	-	-	-	-	-	+	-	-	
n-propanol	-	-	-	-	-	+	-	-	
1-butanol	-	-	+	+	+	-	-	-	
1-pentanol	-	+	-	+	+	+	-	-	
o-furfuryl alcohol	-	-	-	+	-	-	+	-	
2-ethyl-1-hexanol	-	-	-	-	+	-	-	*	
phenol	-	-	+	+	-	+	_	-	
2,2,4-trimethylpentyl- 1,3-diol	-	-	_	7	-	-	_	-	
α-terpineol	_	-	*	-	+	-	-	-	
Acids									
acetic acid	-	*	_	+	_	_	+	_	
decanoic acid	-	-	-	-	+	-	-	-	
Sulfur Compounds									
sulfur dioxide	-	-	-	-	-	-	-	+	
carbon disulfide	+	+	+	+	+	+	+ +	. +	
dimethyl disulfide	-	+	+ +	+	+	+	+	+	
carbonyl sulfide	-	-	-	+	-	_	-	-	

Table 16 (cont'd.)

			Samp	le Numb	er ^b		_	
Compound	1081	1040	1107	1115	2048	2071	3053	3111
Nitrogen Compounds								
nitromethane	-	-	+	-	-	-	-	-
C ₅ H ₆ N ₂	•	-	-	+	-	-	-	-
с ₅ н ₈ и ₂	-	-	-	+	-	-	-	-
C4H4N2O	-	-	-	+	-	-	-	-
methyl acetamide	-	_	+	~	-	-	-	-
benzonitrile	-	-	+	+	-	+	-	-
methyl cinnoline	-	-	**	+	-	-	-	-
Esters								
vinyl propionate	_	+	+	+		-	_	-
ethyl acetate	-	_	-	-	-	+	-	-
ethyl-n-caproate	-	-	-	-	-	+	-	-
methyl caprylate	_	-	-	-	-	+	-	-
ethyl caprylate	-	-	-	-	•	+	-	-
iscamyl formate	-	+	-	-	-	-	-	-
methyl decanoate	~	-	-	-	•	+	-	~
ethyl decanoate	-	•	-	-	-	+	-	-
Ethers								
dimethyl ether	-	+		-	-	-	_	_
<u>p</u> -dioxane	-	-	+	-	-	-	-	-
dihydropyran	-	-	+	+	-	•	-	-
poxide								
1,8-cineole	•	-	-	+	~	-	-	-
urans								
furan	-	-	-	-	-	-	+	-
tetrahydrofuran	_	-	+	-	-	-	-	-
methyl furan	-	~	-	+	-	-	+	-
methyl tetrahydrofuran	-	+	-	-	•	-	•	-
ethylfuran	-	-	+	+	-	-	-	-
dimethylfuran	-	-	-	+	-	-	-	-
2-vinylfuran	-	-	-	-	-	-	+	-
furaldehyde	-	-	-	+	-	-	+	-
2-n-butylfuran	-	+	-		-	-	-	***
2-pentylfuran	+	+	+	+	+	+	+	-
methylfuraldehyde	-	-	-	-	-	-	+	-
furyl methyl ketone	-	-	-	+	-	-	-	-
q-furfuryl alcohol	-	-	•	+	-	-	+	-
benzofuran	-	-	+	+	-	+	-	-

Table 16 (cont'd.)

			Sam	ple Num	ber ^b			
Compound	1081	1040	1107	1115	2048	2071	3053	3111
Alkanes								
с ₃ н ₈	-	-	+	-	-		~	-
C4H10	+	+	+	-	+	+	+	-
c ₅ H ₁₂	+	+	+	+	+	+	+	+
^C 6 ^H 14	+	+	+	+	+	+	+	+
^C 7 ^H 16	+	+	+	+	-	+	+	+
C8H18	+	+	+	+	+	-	+	+
C9H20	+	+	+	+	+	+	+	+
C ₁₀ H ₂₂	+	-	+	+	+	+	+	+
C ₁₁ H ₂₄	+	-	+	+	+	+	+	+
C ₁₂ H ₂₆	+	-	+	+	+	+	+	+
с ₁₃ н ₂₈	-	+	~	***	+	-	+	-
с ₁₄ н ₃₀	-	-	-	+	+	-	+	-
C ₁₅ H ₃₂	•	-	-	+	-	-	+	-
Alkenes								
с ₃ н ₆	+	-	-	~	-	+	-	_
C4H8	+	-	+	-	+	+	-	+
C5H10	-	-	+	-	+	-	-	+
C6H12	+	+	+	+	+	+	+	+
C7H14	+	+	+	+	+	+	+	+
C8H16	+	+	+	+	+	+	+	+
C9H18	+	+	+	+	_	+	+	+
C ₁₀ H ₂₀	-	+	+	+	+	+	+	-
C ₁₁ H ₂₂	+	+	+	+	-		+	-
C ₁₂ H ₂₄	-	-	+	-	-	-	-	-
C ₁₃ H ₂₆	-		-	-	-		+	-
isoprene	-	+	-		••	-	~	-
Alkynes								
с ₅ н ₈	-	-	-	~	-	+	-	+
C6H10	•	-	-	-	+	-	-	-
C7H12	+	•	-	-	+	•	+	-
		50				(c	ontinue	d)

Table 16 (cont'd.)

			Sam	ple Num	berb			
Compound	1081	1040	1107	1115	2048	2071	3053	311:
Alkynes (continued)								
C ₈ H ₁₄	-	+	-	+	-	-	+	
C9H16	+	-	-	+	+	-	+	-
C ₁₀ H ₁₈	-	-	+	+	-		-	-
c ₁₂ H ₂₂	-	-	-	+	-	-	-	-
Cyclic Hydrocarbons								
cyclopentane	+	+	+	+	-	+	_	+
methylcyclopentane	+	-	+	•	+	+	+	+
cyclohexane	+	+	+	-	+	+	-	-
ethylmethylcyclohexane	-	-	+	-	-	•	-	-
C ₁₀ H ₁₄ isomers	+	-	-	-	-	-	-	-
C ₁₀ H ₁₆ isomers (other)	+	+	-	-	+	+	-	-
limonene	+	+	+	+	+	+	+	+
methyldecalin	-	-	+	-	_	-	-	_
a-pinene	-	-	+	-	-	-	•	_
camphene	-	-	-	-	-	+	•	-
camphor	•	-	-	-	•	+	~	-
Aromatics								
benzene	+	+	+	+	+	+	+	+
toluene	+	+	+	+	+	+	+	+
ethylbenzene	+	+	+	+	+	+	+	+
xylene	+	+	+	+	+	+	+	+
phenylacetylene	-	-	-	+	-	-	-	-
styrene	+	+	+	÷	+	+	+	+
benzaldehyde	+	+	+	+	+	+	+	+
C3-alkylbenzene isomers	+	+	+	+	+	+	+	+
C2-alkylbenzene isomers	-	+	+	+	+	+	+	-
methylstyrene	-	-	+	-	-	+	-	-
dimethylstyrene	+	+	•	+	+	+	-	*
C5-alkylbenzene isomers	-	-	+	+	-	-	-	-
naphthalene	+	+	_	+	+	+	+	-
C6-alkylbenzene isomers	-	-	-	+	-	-	-	-

Arrenged by class in approximate elution order. See Appendix D for sampleby-sample identifications. + * present; - = not identified in sample.

b Participant code number.

Table 17. VOLATILES QUANTITATED IN MOTHER'S MILK SAMPLES (ng/ml.)

Site	Sample Number	Ch loroform ^b	Tetrachloro- ethylene	Chlorobenzene	Dichloro- benzene ^C
Bayonne, NJ	1016	_d	1.5	0.2	6.7
,,	1032	0.3	1.5	0.1	9.1
	1040	0,1	1.1	0.1	66
	1057	0.7	0.9	0.1	0.2
	1073	0.7	3.8	0.1	2.2
	1081	1.3	6.3	0.1	32
Jersey City, NJ	1024	13	43	0.1	2.8
•	1107	17	7.4	0.2	68
	1115	1.7	8.1	0.3	49
	1123	20	17	0.1	2.2
	1164	65	4.0	0.1	0.9
Pittsburgh, PA	2014	0.9	0.8	0.2	0.2
	2022	1.5	1.8	0.1	1,1
	2048	0.6	1.8	0.1	8.9
	2055	0.8	1.0	0.05	0.7
	2063	0.6	1.6	0.1	3.1
	2071	1.2	1.0	0.1	1.4
	2089	0.7	26	0.2	0.5
	2097	6.7	1.8	-	0.3
	2105	2.8	1.3	0.4	1.1
	2113	1.2	0.7	0.1	0.4
	2121	0.8	2.4	TRe	2.0
	2139	0.6	0.7	0.1	0.9
Baton Rouge, LA	3012	2.9	0.1	0.3	4.2
-	3020	0.7	0.5	0.1	0.6
	3038	0.8	1.7	0.2	1.3
	3046	21	2.5	0.1	2.2

Table 17 (cont'd)

Site	Sample Number	Chloroform ^b	Tetrachloro- ethylene	Chlorobenzene	Dichloro- benzene [©]
	3053	0.3	0.4	0.2	1.8
	3079	0.8	0.6	0.1	0.2
	3087	0.7	0.4	0.2	5,2
	3095	1.3	1.0	0.3	4.2
	3103	0.6	0.2	0.1	>22
	3111	1.8	0.5	~	44
Charleston, WV ^g	4010	5.0	1,2	0.1	0.7
·	4028	7.2	1.4	0.2	1.9
	4036	7.5	3.9	10	0.2
	4051	8.2	0.6	0.2	1.1
	4069	_	0.4	0.1	3.6
	4085	5.3	0.4	-	3.8
	4093	12	1.0	0.1	0.04
	4101	8.7	1.0	0.1	26
	4119	11	>19f	0.04	1.4

^aParticipant code number.

bSee text for caveats with respect to chloroform.

^cAll isomers summed.

d_{Not detected.}

e_{Trace.}

finstrument saturated.

 $g_{\text{Sample 4044 lost due to instrumental malfunction.}}$

quantitation of the other five compounds is not reported, since the levels in milk were not judged sufficiently greater than background to be reliable.

Upon inspection, it is obvious that most values are low relative to only a few high "outliers." These high values suggest that there is a range of levels of these compounds which may correlate with exposure. These results were analyzed statistically to determine if any of the values correlated significantly. As can be seen in Table 18, the arithmetic mean and median values generally are quite different. The arithmetic mean is skewed toward the high end, generally due to one or two relatively high values. A more realistic representation of the central data is the geometric mean. These geometric mean values were tested for their significance (i.e., are the geometric means significantly different from site to site?). Table 19 summarizes this data. From this table, it appears that samples from Jersey City have significantly higher levels of chloroform, tetrachloroethylene, and dichlorobenzene than the other study samples. Charleston samples appear to have significantly higher levels of chloroform, and Bayonne samples appear to have significantly higher levels of dichlorobenzene.

To test if any of the compound levels were related, the Spearman correlation coefficients (nonparametric correlation based on the sample, designed to lessen the weight of a single high outlier) were determined as shown in Table 20. There does not appear to be any compound-to-compound correlation among the subjects.

In interpreting these data, it must be remembered that this is a very small data set. Therefore these data should not be used to extrapolate to the city or area from which the samples were collected.

Quality Control

Table 21 presents the quality control results for chloroform, tetrachloroethylene, chlorobenzene, and dichlorobenzene. The very high recovery of chloroform from the controls indicates either a miscalculation of the amount actually spiked or contamination of the samples used as controls. Since the procedural blanks contained about 15 times less chloroform, the former explanation is most reasonable. However, the chloroform values reported in Table 17 must be interpreted subject to the following

Table 18. SUMMARY STATISTICS FOR VOLATILE COMPOUNDS BY SITE a

Site	Chloroform	Tetrachloro- ethylene	Chloro- benzene	Dichloro- benzene
Bayonne, NJ				
Maximum	1.3	6.3	0.2	66
Meanb	0.52	2.52	0.12	19.37
Median	0.5	1.5	0.004	7.9
S.D.	0.48	2.13	0.1	25.54
n	6	6	6	6
Jersey City, NJ				
Maximum	65	43	0.3	68
Mean ^b	23.34	15.9	0.16	24.48
Median	17	8.1	0.1	2.8
S.D.	24.3	15.9	0.089	31.69
n	5	5	5	5
Pittsburgh, PA				
Maximum	6.7	26	0.4	8.9
Mean ^b	1.53	3.41	0.12	1.71
Median	0.85	1.45	0.1	1
S.D.	1.74	7.13	0.11	2.41
n	12	12	12	12
Baton Rouge, LA				
Maximum	21	2.5	0.3	44
Mean ^b	3.09	0.79	0.16	8
Median	0.8	0.5	0.15	3.2
S.D.	6.34	0.75	0.096	13.98
n	10	10	10	10
Charleston, WV				
Maxipum	12	>19	10	26
Mean ^b	7.21	3.21	1.20	4.30
Median	7.5	1	0.1	1.4
S.D.	3.55	6.02	3.30	8.25
n	9	9	9	9
Overall				
Maximum	65	43	10	68
Mean ^D	5.57	4.10	0.37	9.15
Median ·	1.25	1.25	0.1	1.95
S.D.	10.9	8.15	1.53	17.3
n	42	42	42	42

aMaximum, mean and median values are ng/mL.

^bArithmetic mean.

Table 19. SIGNIFICANCE OF THE DIFFERENCES IN THE GEOMETRIC MEANS BY SITE

		Geometric Mean (ng/mL)							
Site	Chloroform	Tetrachloroethylene	Chlorobenzene	Dichlorobenzene					
Bayonne	0.45	2.09	0.12	8.33					
Jorsey City	14.7	11.5	0.16	8.55					
Pittsburgh	1.23	1.82	0.12	1.21					
Baton Rouge	1.53	0.67	0.15	3.83					
Charleston	5.92	1.65	0.42	1.98					
Significance ^a	0.01	0.01	N.S.b	0.05					

 $^{^{8}0.01}$ implies 99 percent confidence that the numbers are statistically different, while 0.05 implies 95 percent confidence.

^bNot significant.

Table 20. SPEARMAN CORRELATION COEFFICIENTS FOR VOLATILE ORGANICS FOUND IN MOTHER'S MILK

	Chloroform	Tetrachloro- ethylene	Chlorobenzene	Dichloro- benzene
Chloroform	1.0	0.37 ^a	-0.02 ^b	-0.13 ^b
Tetrachloro- ethylene		1.0	0.007 ^b	0.05 ^b
Chlorobenzene			1.0	0.03 ^b
Dichloro- benzene				1.0

aSignificant at 0.05 level (95 percent confidence).

b Not significant
Sample size = 42

5

Table 21. QUALITY CONTROL RESULTS FOR VOLATILES IN MILK

Type of Sample	Chloroform	Tetrachloroethylene	Chlorobenzene	Dichlorobenzene
Blanks ^a				
n	7	7	7	17
Mean (ng/mL) ^b	1.2	0.22	0.03	0.12
S.D.	1.3	0.11	0.025	0.19
RSD (Z)	108	49	84	159
Controls ^C				
n	8 -	8	8	o d
Mean Recovery e	14.02 ¹	1.12	0.62	_
S.D.	8.20	0.41	0.34	_
RSD (%)	58	37	55	

a Blanks consisted of two field water blanks and five water blanks purged with the milk samples to monitor procedural background. No difference between the two types of blanks was observed.

D Arithmetic mean.

Controls consisted of two spiked raw cow's milk samples carried to the field and returned, two spiked raw cow's milk samples stored in the laboratory, two spiked water samples carried to the field and returned, and two spiked water samples stored in the laboratory. No major differences were observed between the four types of samples. Samples were spiked at 30-90 ng/volume purged (or about 1 ng/mL).

d Not included in control spiking solution.

^{1.0 = 100} percent recovery.

Extremely high recovery probably a result of improper loading of controls.

considerations: the mean reported levels in the samples were only 4.9 times the blank levels; the recovery from controls was about 1400 percent, invalidating the recovery study; and chloroform is known to be a laboratory atmospheric contaminant.

The compounds presented in Table 17 represented significant levels above the background in blanks. Several other compounds were quantitated that did not exhibit substantial concentrations. These compounds, with the ratio of the mean in the samples to the mean in the background given in parenthesis, were: 1,1,1-trichloroethane (1:1), benzene (2:1), toluene (2:4), trichloroethylene (1:2) and carbon tetrachloride (1:4). These levels in the samples cannot be reliably assigned to either the milk sample or to laboratory contamination. If these compounds are present in milk, they are very low and cannot be regarded as significant, given the limitations of the technique employed. Apparently, mother's milk does not represent a bioconcentration matrix for these compounds.

SEMIVOLATILES

Three samples were fully interpreted, as presented in Appendix E. As can be seen from the data, few compounds of interest were observed in the mass spectra. The data were searched on the GC/MS data system for target compounds (PCNs, PBBs and PCBs) using single ion plots called up from the full data set. No evidence for any of these compounds was observed at a detection limit of about 20 ppb. DDE was quantitated in five samples as shown in Table 22. These values were in the range generally reported by previous investigators (see Tables 2 - 4). Since none of the target compounds were present in detectable quantities, no further identification or quantitation was attempted.

Table 22. DDE AND TETRACHLOROBIPHENYL LEVELS IN SELECTED MOTHER'S MILK SAMPLES

		ng/ml Milk		
Site	Sample Number	DDE	Tetrachlorobiphenyl	
Pittsburgh	2105	45	ирь	
Pittsburgh	2121	73	Ic	
Charleston, WV	4069	107	ND	
Charleston, WV	4085	38	ND .	
Charleston, WV	4093	91	nd	
	Mean d	71	-	
	S.D.	29		
	RSD (%)	42		
	Median	73		

a Samples selected as having the most intense total ion current chromatograms.

b Not detected.

c Trace.

d Arithmetic mean.

REFERENCES

- Ziegel, E. and C. C. Van Blarcom, <u>Obstetric Nursing</u>, 6th ed., Macmillan, New York, 651 (1972).
- Strassman, S. C. and F. W. Kutz, "Insecticide Residues in Human Milk from Arkansas and Mississippi, 1973-74," Pest. Mon. J., <u>10</u>, 130-133 (1977).
- 3. Savage, E. P., et al., "Organochlorine Pesticide Residues and Polychlorinated Biphenyls in Human Milk 1971-72," Pest. Mon. J., 7, 1-3 (1973).
- Kroger, M., "Insecticide Residues in Human Milk," J. Pediat., 80, 401-405 (1972).
- 5. Curley, A. and R. Kimbrough, "Chlorinated Hydrocarbon Insecticides in Plasma and Milk of Pregnant and Lactating Women," Arch. Environ. Health, 18, 156-164 (1969).
- Dyment, P. G., et al., "Relationship Between Levels of Chlorinated Hydrocarbon Insecticides in Human Milk and Serum," Bull. Environ. Contamin. Toxicol., 6, 449-452 (1971).
- Hagyard, S. B., W. H. Brown, J. W. Stull, F. M. Whiting, and S. R. Kemberling, "DDT and DDE Content of Human Milk in Arizona," Bull. Environ. Contamin. Toxicol., 9, 169-172 (1973).
- Quinby, G. E., J. F. Armstrong and W. F. Burham, "DDT in Human Milk," Nature, 207, 726-728 (1965).
- 9. Laug, E. P., F. M. Kunze and E. S. Prickett, "Occurrence of DDT in Human Fat and Milk," Arch. Indust. Hyg., 3, 245-246 (1951).
- Wilson, D. J., et al., "DDT Concentrations in Human Milk," Am. J. Dis. Child, 125, 814-817 (1973).
- 11. Woodard, B. T., B. B. Ferguson and D. J. Wilson, "DDT Levels in Milk of Rural Indigent Blacks," EPA-600/1-76-032 (1976).

- Savage, E. P., et al., "A Search for Polychlorinated Bipbenyls in Human Milk in Rural Colorado," Bull. Environ. Contamin. Toxicol., 9, 222-226 (1973).
- Petrakis, N. L., L. D. Gruenka, T. C. Beelen, N. Castagnoli, Jr., and J. C. Craig, "Nicotine in Breast Fluid of Nonlactating Women," Science, 199, 303-305 (1978).
- 14. Bakken, A. F. and M. Seip, "Insecticides in Human Breast Milk," Acta Paediatr. Scan., 65, 525-529 (1976).
- 15. Knoll, W. and S. Jayaraman, "Zur Kontamination Von Humanmilch mit chlorierten Kohlenwasserstoffer," Die Nahrung, 17, 599-615 (1973).
- 16. Psendorfer, Von H., "Rüchstände von Organochlorpestiziden (DDT u.a.) und polychlorierten Biphenylen (PCBs) in der Muttermilch," Wiener Klinische Wochenschrift, 87, 731-736 (1976).
- 17. Th. Tuinstra, L. G. M., "Organochlorine Insecticide Residues in Human Milk in the Leiden Region," Neth. Milk Diary J., 25, 24-32 (1971).
- Polishuk, Z. W., M. Ron, M. Wasserman, S. Cucas, O. Wasserman, and C. Lemesch, "Organochlorine Compounds in Human Blood Plasma and Milk," Pest. Mon. J., <u>10</u>, 121-129 (1977).
- 19. Ritchy, W. R., G. Savary and K. A. McCulley, "Organochlorine Insecticide Residues in Human Milk, Evaporated Milk, and Some Milk Substitutes in Canada," Can. Publ. Health J., 63, 125-132 (1972).
- 20. Egan, H., R. Goulding, J. Roburn and J. O'G. Tatton, "Organo-chlorine Pesticide Residues in Human Fat and Human Milk," Brit. Med. J., 2, 66-69 (1965).
- 21. Newton, K. G. and N. C. Greene, "Organochlorine Pesticide Residue Levels in Human Milk -- Victoria, Australia 1970," Pest. Mon. J., 6, 4-8 (1972).
- 22. Graca, I., A. M. S. Silva Fernandes and H. C. Mourso, "Organochlorine Insecticide Residues in Human Milk in Portugal," Pest. Mon. J., 8, 148-156 (1974).
- 23. Musial, C. J., O. Hutzinger, V. Zitko and J. Crocker, "Presence of PCB, DDE, and DDT in Human Milk in the Providences of New Brunswick and Nova Scotia, Canada," Bull. Environ. Contamin. Toxicol., 12, 258-267 (1974).

- 24. Mes. J. and D. J. Davies, "Presence of Polychlorinated Biphenyl and Organochlorine Pesticide Residues and the Absence of Polychlorinated Terphenyls in Canadian Human Milk Samples," Bull. Environ. Contam. Toxicol., 21, 381-387 (1979).
- 25. Stacey, C. I. and B. W. Thomas, "Organochlorine Pesticide Residues in Human Milk, Western Australia -- 1970-71," Pest. Mon. J., 9, 64-66 (1975).
- 26. Van House Holdrinet, M., H. E. Braun, R. Frank, G. J. Stopps, M. S. Smout, and J. W. McWade, "Organochlorine Residues in Human Adipose Tissue and Milk from Ontario Residents," Can. J. Pub. Health, 68, 74-80 (1977).
- 27. Winter, M., M. Thomas, S. Wernick, S. Levin and M. T. Farver, "Analysis of Pesticide Residues in 290 Samples of Guatemalan Mother's Milk,"
 Bull. Environ. Contamin. Toxicol., 16, 652-657 (1976).
- 28. "Criteria for a Recommended Standard...Occupational Exposure to Ethylene Dichloride (1,2-dichloroethane)," HEW Publ. No. (NIOSH) 76-139 (March 1976).
- 29. Kover, F. D., Environmental Hazard Assessment Report. Chlorinated Naphthalenes. EPA 560/8-75-001 (December 1975).
- 30. Erickson, M. D., R. A. Zweidinger, L. C. Michael and E. D. Pellizzari, "Environmental Monitoring Near Industrial Sites: Polychloronaphthalenes," EPA-560/6-77-019 (1977).
- 31. Erickson, M. D., L. C. Michael, R. A. Zweidinger, and E. D. Pellizzari, "Development of Methods for Sampling and Analysis of Polychlorinated Naphthalenes in Ambient Air," Environ. Sci. Technol., 12, 927-931 (1978).
- 32. Erickson, M. D., L. C. Michael, R. A. Zweidinger and E. D. Pellizzari, "Sampling and Analysis for Polychlorinated Naphthalenes in the Environment," JAOAC, 61, 1335-1346 (1978).
- 33. Erickson, M. D., L. C. Michael, R. A. Zweidinger, and E. D. Pellizzari, "Development of Methods for Sampling and Analysis of Polychlorinated Naphthalenes in Ambient Air," 1977 Annual Meeting, American Chemical Society, Chicago, IL (August 31, 1977).

- 34. Erickson, M. D., L. C. Michael, R. A. Zweidinger, and E. D. Pellizzari, "Sampling and Analysis for Polychlorinated Naphthalenes in the Environment," 1977 Annual Meeting AOAC, Washington, DC (October 20, 1977).
- 35. Unpublished data, E. Roessler, Borough of Bridgeville, PA (1976).
- 36. 1977 Directory of Chemical Producers-USA, Chemical Information Services, Stanford Research Inst., Menlo Park, CA (1977).
- 37. Environmental Sciences and Engineering, "Trip Report for Sampling of Polybrominated Biphenyls (PBBs)," submitted to OTS, EPA, Washington, DC, Contract No. 68-01-3248 (April 1977)).
- 38. Mumma, C. E. and D. D. Wallace, "Survey of Industrial Processing Data.

 Task I Pollution Potential of Polybrominated Biphenyls," EPA-560/3-75-004 (June 1975).
- 39. Unpublished data, E. J. Londres, New Jersey Dept. of Environmental Protection via G. E. Parris, OTS, EPA, Washington, DC (1977).
- 40. Erickson, M. D., R. A. Zweidinger, and E. D. Pellizzari, "Analysis of a Series of Samples for Polybrominated Biphenyls (PBBs)," EPA-560/6-77-020 (August 1977).
- 41. Environmental Science and Engineering, "Data Report for Polybrominated Biphenyl Near Manufacture (sic) in the Northeast," submitted to OTS, EPA, Washington, DC Contract No. 68-01-3248 (June 16, 1977).
- 42. 1974 New Jersey State Industrial Directory, New Jersey State Industrial Directory, 2 Penn Plaza, NY, 10001 (1974).
- 43. Pellizzari, E. D., "The Measurement of Carcinogenic Vapors in Ambient Atmospheres," EPA-600/7-77-055 (June 1977).
- 44. Pellizzari, E. D., M. D. Erickson, and R. A. Zweidinger, "Formulation of a Preliminary Assessment of Halogenated Organic Compounds in Man and Environmental Media," EPA-560/13-79-006 (July 1979).
- 45. Pellizzari, E. D., M. D. Erickson, T. D. Hartwell, S. R. Williams, C. M. Sparacino and R. D. Waddell, "Preliminary Study on Toxic Chemicals in Environmental and Human Samples. Part I: Formulation of an Exposure and Body Burden Monitoring Program," submitted to U. S. Environmental Protection Agency, Washington, DC, Contract No. 68-01-3849 (June 1980).

- 46. Pellizzari, E. D., "Analysis of Organic Air Pollutants by Gas Chromatography and Mass Spectroscopy," Publication No. EPA-600/2-77-100, Contract No. 68-02-2262, (June 1977).
- 47. McDonnell, G., D. M. Ferguson and C. R. Pearson, "Chlorinated Hydrocarbons and the Environment," Endeavour, 34, 13-18 (1975).
- 48. FDA Compliance Program, Evaluation, "FY 74 Total Diet Studies (7320.08)," Date accepted: January 21, 1977.
- 49. State of New Jersey Department of Environmental Protection, "Initial Report on the Findings of the State Air Monitoring Program for Selected Volatile Organic Substances in Air," (October 1979).
- 50. Zweidinger, R. A., A. Sherdon, B. S. Harris, III, H. Zelon, T. Hartwell, and E. D. Pellizzari, "Measurement of Benzene Body Burden of Potentially Environmentally Exposed Individuals," Final Report, EPA Contract No. 68-01-3849, Task 1 (May 1980).
- 51. Hartwell, T., P. Piserchia, S. White, N. Gustafson, A. Sherdon, R. Lucas, D. Lucas, D. Myers, J. Batts, R. Handy, and S. Williams, "Analysis of E?A Pesticide Monitoring Networks," Office of Toxic Substances, Washington, DC. Draft Report (1979).
- 52. U.S. Environmental Protection Agency, Office of Research and Development, "Health Assessment Document for Polycyclic Organic Matter," (May 1978).
- 53. Stanford Research Institute, "The Environmental Fate of Selected Polynuclear Aromatic Hydrocarbons," Prepared for U. S. Environmental Protection Agency (February 1976).
- 54. State of New Jersey Department of Environmental Protection, "Initial Report on the Findings of the State Air Monitoring Program for Selected Heavy Metals in Air," (October 1979).
- ..55. Unpublished data, William J. Librizzi, U.S. Environmental Protection Agency, Region II (October 1977).
 - 56. Fribers, L., M. Piscator, G. F. Nandberg and T. Kjellstrom, "Cadmium in the Environment," CRC Press, Cleveland, OH (1974).

- 57. National Academy of Sciences, "Lead," Washington, DC (1972).
- 58. Mason, T. J., F. W. McKay, "U.S. Cancer Mortality by County: 1950-69,"

 DHEW Publ. No. (NIH), 74-615, Washington, DC, U.S. Govt. Printing Office (1974).
- 59. Mason, T. J., F. W. McKay, J. R. Hoover, W. Blot and J. F. Fraumeni, Jr., "Atlas of Cancer Mortality for U.S. Counties: 1950~69," DHEW Publ. No. (NIH) 75-780, Washington, DC, U.S. Govt. Printing Office (1975).
- 60. Greenberg, Michael R., "The Spacial Distribution of Cancer Mortality and of High and Low Risk Factors in the New Jersey-New York-Philadelphia Metropolitan Regions, 1950-1969, Part I," New Jersey Dept. of Environmental Protection, Program on Environmental Cancer and Toxic Substances (January 1979).
- 61. Greenberg, M., F. McKay, and P. White, "A Time-Series Comparison of Cancer Mortality Rates in the New Jersey-New York-Philadelphia Metropolitan Region and the Remainder of the United States, 1950-1969," Am. Jour. of Epidemiology, 111, 166 (1980).
- 62. Greenberg, M. R., P. W. Preuss, and R. Anderson, "Clues for Case Control Studies of Cancer in the Northeast Urban Corridor," Soc, Sci. & Med., 14D, 37-43 (1980).
- 63. Greenberg, M. R., J. Caruana, B. Holcomb, G. Greenberg, R. Parker, J. Louis, and P. White, "High Cancer Mortality Rates from Childhood Leukemia and Young Adult Hodgkin's Disease and Lymphoma in the New Jersey-New York-Philadelphia Metropolitan Corridor, 1950-1969," Cancer Research, 40, 439-443 (1980).
- 64. Cross, J. and G. B. Wiersma, "Preliminary Analysis of Cancer Rates in Organic Chemical-Producing Counties," EPA-600/1-79-022 (June 1979).
- 65. Pellizzari, E. D., and M. D. Erickson, "Analysis of Organic Air Pollutants in the Kanawha Valley, WV and the Shenandoah Valley, VA," Publication No. EPA-903/9-78-007, Contract No. BOA 68-02-2543 (June 1978).
- 66. Erickson, M. D., S. P. Parks, D. Smith and E. D. Pellizzari, "Sampling and Analysis of Organic Air Pollutants in Two Industrialized Valleys," FACSS V, Boston (October 30 November 3, 1978.
- 67. McLafferty, F. W., E. Stenhagen, and S. Abrahammson, Ed., Registry of Mass Spectral Data, John Wiley and Sons, New York (1974).

68. <u>Eight Peak Index of Mass Spectra</u>. Vol. I (Tables 1 and 2) and II (Table 3), Mass Spectrometry Data Centre, AWRE, Aldermaston, Reading, RG74PR, UK (1970).

APPENDIX A DATA COLLECTION INSTRUMENTS

STUDY OF ORGANIC COMPOUNDS IN HUMAN MILK

EPA Contract No. 68-01-3849 RTI Project No. 31U-1521-22

DATA COLLECTION INSTRUCTIONS

Performed for

Office of Toxic Substances
Environmental Protection Agency
Washington, DC 20460

1.0 Introduction

Under contract to the Office of Toxic Substances, Environmental
Protection Agency (EPA), the Research Triangle Institute (RTI) is
conducting a limited study designed to measure environmental pollutant
levels in human milk and to evaluate the utility of using this body
fluid in specific pollutant studies for populations in the vicinity of
manufacturing plants and/or industrial user facilities. RTI is responsible
for all phases of the study, including study design, subject recruitment,
chemical analysis of milk samples, and report writing. RTI is a not-for-profit
contract research organization located in North Carolina's Research Triangle
Park between Raleigh, Durham, and Chapel Hill. The Institute was incorporated
as a separate operating entity in 1958 by the University of North Carolina .

(UNC) at Chapel Hill, Duke University at Durham, and North Carolina State
University at Raleigh, and is still closely affiliated with the three
universities.

2.0 Overview

Four urban areas have been chosen as performance sites; they are Bridgeville, Pennsylvania; the area which includes Linden and Bayonne, New Jersey and western Staten Island, New York; Baton Rouge, Louisiana; and South Charleston and Nitro, West Virginia. These sites represent high-probability areas for the presence of one or more of the chemicals of interest in human milk. The selected industrial chemicals of interest include polychlorinated naphthalenes, tetrachlorethylene, trichloroethane, dichloropropane, benzene, polybrominated biphenyls, chlorinated phenols, toluene, chlorinated benzenes, and chloroform.

At each of the four sites, arrangements will be made to work through clinical facilities such as hospitals, clinics, or physician's offices, in order to recruit a panel of respondents. At each site ten participants will be recruited, for a total of 40. Potential participants (lactating females) will be screened to determine that they live in one of the areas of interest and are willing and able to provide the milk sample.

A questionnaire will be administered for each participant to obtain information on demographic variables, residence histories, and potential exposure situations; for each participant, a sample of milk will be collected and analyzed for the compounds of interest by gas chromatography/mass spectrometry or high pressure liquid chromatography. A professional member of the facility's staff, such as a registered nurse, will be trained in the proper procedures to administer the questionnaire and obtain the milk sample. To try to reduce the non-participation rate due to refusals, and to reimburse the subject for the time spent on the study, volunteers will be offered a \$5.00 incentive for participating.

3.0 Data Collection

3.1 General Remarks

Data collection for this research effort consists of the following steps:

1. Screening of potential participants (lactating women) to determine that they live in one of the areas of interest (see below), that they have resided in that area for at least the preceding 12 months, that they have remained in that area continuously for the preceding week, and that they are willing and able to provide a milk sample.

- When an eligible person is encountered, the nature and purpose of the study will be explained and their participation solicited.
- 3. When an eligible person agrees to participate, the person will be required to sign a Participant Consent Form (PCF) in order to participate in the study.
- 4. Once the participant has signed the PCF, the person should be listed on the Participant Listing Form (PLF), a Patient Number assigned, and the data collector will proceed to administer the Study Questionnaire (SQ) and collect the milk sample.
- 5. Once the SQ has been administered and the milk sample collected, the participant will be offered a \$5.00 incentive for participating.
- Milk samples and completed data collection instruments will be returned to RTI.

3.2 Survey Instruments

As indicated in the preceding section, there are 3 data collection instruments for this research effort, the PCF, the PLF, and the SQ; subsequent sections contain instructions for the use of each instrument as well as item-by-tiem explanations for their completion, and general descriptions are provided below. The survey instruments have been designed hopefully to provide an efficient means of collecting and recording the requisite data for the study. It is imperative that all survey instruments be completed accurately. The success and reliability of the study and its results are dependent upon the quality of data collected, which will be fully dependent

on the accuracy of your execution of your assignment. As you complete a form, conduct a thorough edit to verify that required data have been entered and entered correctly. Copies of the data collection instruments appear in Attachment 1.

3.2.1 Participant Consent Form (PCF)

- Purpose: The purposes of the PCF are to introduce the study; explain its objectives, sponsorship (the relationship and roles of RTI and EPA), and requirements of and risks, burdens, and benefits to participants; and stress that participation is completely voluntary and that all data collected will be kept confidential.
- . General Description: The PCF is a single page form printed on special paper which makes three copies from a single impression. The survey title appears at the top, along with the name of RTI; spaces for necessary identifying information appear at the bottom.
 - Administration: The PCF will be signed by the participant and contains an agreement to provide the necessary information and milk sample. Participants may freely withdraw from the study at any time; however, in order to encourage participation RTI offers an incentive of five dollars to each participant to be paid after each data set (PCF, SQ, and milk sample) is obtained. Again, confidentiality of data is stressed, including steps

taken to disassociate the name of the participant from the data once collected; for example, the PCF is the only data collection instrument which bears the name of the participant and allows its association to study identification numbers, but will be maintained in hard copy only and stored in a restricted area. To further emphasize this disassociation, the incentive will be paid in cash rather than by check or money order, although the participant will sign the PCF indicating that the incentive was received. A signed PCF must be obtained for each participant before proceeding with Study Questionnaire (SQ) administration and collection of the milk sample.

<u>Disposition</u>: The top (white) copy will be attached to the appropriate SQ until it is received at RTI and verified; the yellow copy will be provided to the participant; the pink copy will be retained by the data collector.

3.2.2 Participant Listing Form (PLF)

- Purpose: The purpose of the PLF is to provide a means of assigning unique numbers to participants at each performance site.
 - General Description: The PLF is a single page form printed on pink paper; space for Comments is provided on the reverse side. The survey title appears at the top, along with the names and addresses of RTI and EPA/OTS and a confidentiality statement.

- . Administration: As each participant is enlisted up to the required number (10), that participant should be listed on the PLF.
- . <u>Disposition</u>: When data collection at a site or facility is completed, the PLF (or a copy) should be sent to RTI.

3.2.3 Study Questionnaire (SQ)

- Purpose: The purpose of the SQ is to obtain information on participants, including demographic characteristics such as age, sex, race, and occupation; residence information; health information such as current health status and prescription medications; and personal characteristics such as hobbies.
- . General Description: The SQ is divided into six sections, dealing respectively with demographic characteristics, occupation, health and personal habits, residence and household information, information on the interviewer and respondent, and information regarding the milk sample, including an indication as to whether or not the milk sample was obtained, the date and time of acquisition of the sample, and the date the sample was shipped to RTI. Participants will be identified by a unique study number used to correlate and cross-identify the questionnaires and samples by way of pre-printed self-adhesive labels. The SQ is 5 pages long, with space provided for comments.

- . Administration: An SQ is to be completed for each participant for whom a signed PCF is obtained.
- . Disposition: The SQ's are to be sent to RTI as instructed.

3.3 Screening

As indicated in section 3.1, potential participants (lactating women) should be screened to determine that they meet certain study criteria for participation:

- That they are willing and able to provide a milk sample of sufficient quantity (approximately 100 ml.),
- 2. That they live in one of the areas of interest (see below),
- That they have resided in that area for at least the preceding
 months, and
- 4. That they have remained in that area continuously for the praceding 7 days.

As indicated in section 2.0, four areas have been chosen as performance sites, with a specific Site Number assigned to each which will remain constant for each site and is to be entered where appropriate on data collection instruments as follows:

<u>S1te</u>	Site Number	
Northern New Jersey/Staten Island, New York	1	
Bridgeville, Permsylvania	2	
Baton Rouge, Louisiana	3	
Nitro/South Charleston, West Virginia	4	

With the exception of Bridgeville, Pennsylvania, participants residing in some areas at each site are of considerably more interest to the study than those living in others, as discussed in the following sections.

3.3.1 Northern New Jersey/Staten Island, New York

Within the Northern New Jersey/Staten Island area, potential participants residing in some communities are of more interest than those residing in others, more or less in the order listed below:

- 1. Bayonne, NJ
- 2. Northern Staten Island (Port Richmond), NY
- 3. Linden, NJ
- 4. Carlstadt, NJ
- 5. Saddle Brook, NJ
- 6. Jersey City, NJ
- 7. Kearney, NJ
- 8. Newark, NJ

- 9. Elizabeth, NJ
- 10. Sayreville, NJ
- 11. Rahway, NJ
- 12. Edison, NJ
- 13. Parlin, NJ
- 14. Passaic. NJ
- 15. Patterson, NJ
- 16. Wayne, NJ

3.3.2 Baton Rouge, Louisiana

Potential participants residing in Baton Rouge are of primary interest to this study; other communities in the Baton Rouge area of interest are Placquemine, St. Gabriel, and Geismar.

3.3.3 Nitro/South Charleston, West Virginia

Potential participants residing in Nitro and South Charleston are of primary interest to this study; other communities of interest in the area are Belle and Institute.

3.4 Participant Listing Form

When an eligible person is encountered who agrees to participate, that person should be listed on a PLF in order to be assigned a unique Participant Number. The PLF is completed by entering the appropriate Site Number (see section 3.3 above); then, each time that an eligible participant is encountered who agrees to participate, up to the number required, enter the Participant's Name (Last, First, Middle) on the PLF and assign a Participant Number in the left-hand column, beginning with 0001 at each site unless other-

wise instructed. Assign Participant Numbers consecutively for all study participants. Where appropriate, enter the participant's Medical Record Number in the right-hand column. When making numerical entries, right-adjust and enter leading zeros.

3.5 Participant Consent Form

Potential participants must understand exactly what is involved in participation in the study and what benefits may be realized by participation; this understanding and agreement must be documented by a signed PCF. In the event that the potential participant is under the age of 18 years, the person's parent or other legal guardian must sign the PCF in order for the designated eligible to participate.

More specifically, the potential participant and/or that person's parent, guardian or other spokesman, must understand that full participation in the study consists of providing answers to a questionnaire related to environmental exposure, part of which relates to the individual's household in general and part of which is related to the individual participant (be prepared to show the person the SQ), and providing a milk sample of approximately 100 ml. (be prepared to show the person one of the collection bottles.)

The individual must further understand that she will only enjoy certain limited benefits in return for her time and inconvenience, primarily a \$5.00 incentive to be disbursed after administration of the questionnaire and collection of the milk sample. The individual must understand that participation in the study is completely voluntary and that she may withdraw at any time, but that payment of the incentive is dependent on full participation. The individual must also understand that all data collected in the study will be held strictly confidential, and that names will not be disclosed.

If the participant or that perons's parent, guardian or other spokesman agrees to participate, read through the PCF with them and make entries where appropriate. At the bottom, record the Data (month, day, and year) that the PCF is signed and print the Participant's full Name (First, Middle or Maiden, Last - do not abbreviate); record the appropriate Site Number (see section 3.3 above) and Participant Number (from the PLF); have the participant (or other appropriate person) sign the PCF; enter your signature as witness; and record the participant's home Address (Street Number and Name, City, State, and Zip Code) in the spaces provided.

After data collection (administration of SQ and collection of milk sample) is completed, the participant (or that person's parent or guardian) should be given \$5.00. The recipient must sign in the space provided at the bottom of the PCF to indicate receipt of the incentive. Should the signatures on the PCF for Participant and Recipient be other than the participant's, please explain in the Comments section of the SQ.

Finally, as indicated in section 3.2.1, the top (white) copy of the PCF is to be attached to the appropriate SQ; the yellow copy is to be provided to the participant or her guardian; and the pink copy is to be retained by the data collector.

3.6 Study Questionnaire

Before proceeding with administration of the SQ, read the justification and confidentiality statement in the box on the cover. Enter the appropriate Site (see section 3.3 above) and Participant (from the FLF) Mambers. Stapled inside the SQ you will find a set of pre-printed, self-achesive labels which are necessary to identify corresponding SQs and samples. Each label contains a unique Study Number, which should be the same on all

labels in a set, and an indication of what the label is for. You should also have some labels that have cally a Study Number and a few that are completely blank; these are for your use in the event that a label is damaged or missing. If you use a label that has a Study Number only, you will have to write on the label what it is intended for, such as MILK; if you use a blank label, you must write on the label the Study Number and what it is intended for. Check to be sure that all the labels in a given SQ contain the same Study Number; if not, do not use the SQ and return it to RTI. If the Study Number is the same on all labels, remove the one for the QUESTIONNAIRE and place it on the cover of the SQ over the spaces provided for the Study Number. Space for Comments is provided on page 5.

If the participant is under 18 years of age, the SQ may have to be administered in whole or part to the parent or guardian, and must be administered in that person's presence. If the participant suffers from a speech or hearing deficit, or is otherwise incapacitated, the SQ may have to be administered to the spouse or some other spokesman.

- Item 1 Race: Indicate the participant's race by placing an X in the appropriate box. This question may be answered by observation; however, if there is any doubt whatsoever, ask.
- Item 2 Age: Determine and enter the participant's age in years as of the last birthday.
- Item 3 Birthdate: Determine and enter the participant's exact birthdate (month, day and year). Again, remember to rightadjust and enter leading zeros. A note on dates: accept and record partial dates, if that is all that the respondent can provide; in that case, indicate missing elements of the date

- with a dash (-) -- for example, April 1977 would be recorded as 014 -1 717.
- Item 4 Weight: Determine and enter the participant's approximate weight in pounds (to the nearest pound—no fractions!)
 or kilograms, in which case observe the decimal.
- Item 5 Height: Determine and enter the participant's approximate height in inches or centimeters.
- Item 6 Current Employment: Determine if the participant is currently employed in any capacity and place an X in the appropriate box.
 If the answer is Yes, continue to Item 7; if the answer is
 No, skip to Item 10.
- Item 7 Length of Present Employment: Determine and record the length of time that the participant has been employed by her present employer; enter the units in the spaces provided and then place an X in the appropriate box to indicate whether the units represent days, months, or years.
- Item 8 Occupation Away From Home: Determine if the participant's occupation usually takes her away from home and place an X in the appropriate box. If Yes, continue to Item 9; if No, skip to Item 11. This question, and Item 9 below, are aimed at eliciting information regarding the location of the participant's various exposure to the environment.
- Item 9 Location of Present Employment: If the participant is
 currently employed, determine the nature (not the name) and
 location (street address, city, state, and Zip Code, if known)

- of the employer. By nature, we mean the type of business, such as service station, school, hospital, grocery store, doctor's office, hotel, restaurant, etc.
- Item 10 Employment Status: If the participant is not presently employed, determine which of the provided categories best describes the participant's status and place an X in the appropriate box. If the response is choice 1 or 2, skip to Item 15; if the response is choice 3-5, continue to Item 11.
- Item 11 Usual Occupation: Determine and record the participant's

 usual (or most common) occupation (when employed); be succinct
 e.g., high school coach, waitress, hotel desk clerk, taxi driver.
- Item 12 Present Occupation: Determine if the participant is presently employed in her usual occupation (indicated in Item 11) and place an X in the appropriate box. Items 12 and 13 may be skipped for unemployed, retired and disabled persons.
- Item 13: If the response to Item 12 was positive, determine how long the participant has been employed in her usual occupation (recorded in Item 11) and record; enter the units in the spaces provided and then place an X in the appropriate box to indicate whether the units represent days, months or years.
- Item 14: Determine if the participant presently works at or in any of the listed occupations or establishments and place an X in each appropriate box.
- Item 15 Present Smoking Status: Ascertain if the participant currently smokes cigarettes, and place an X in the appropriate box. If YES, continue to Item 16; if NO, skip to Item 18.

- "Item 16 Age at First Smoke: If the participant is a smoker

 (a positive response to Item 15), ascertain the age (in years)

 at which the participant started smoking and record in the
 spaces provided.
 - Item 17 Smoking Frequency: Ascertain how many cigarettes the participant smokes per day, on the average, and place an X in the appropriate box. If the participant uses tobacco in some form other than cigarettes, such as snuff, record in the space provided.
 - Item 18 Time Outdoors: Ascertain the average number of hours

 that the participant spends out of doors each day and record
 in the spaces provided -- another indication of environmental
 exposure.
 - Item 19 Time Away From Home: Determine how many hours of the day on the average the participant normally spends more than 2 miles away from home, and record in the spaces provided. This determination should be done separately for weekdays and weekends.
 - Item 20 General Health Status: Using the four qualifiers provided,
 ascertain the participant's general current health status and
 place an X in the appropriate box.
 - Item 21 Prescription Medications: Inquire as to whether the
 participant is currently taking any prescription medication(s)
 on a regular daily basis and place an X in the appropriate
 box; if YES, determine and record the drug name e.g., penicillin,
 oral contraceptives, Valium, phenobarbital, etc.

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- Item 22 Non-prescription Medications: Inquire as to whether the participant has taken any non-prescription medications in the past 24 hours, and place an X in the appropriate box; If YES, determine and record the drug name -e.g., aspirin, vitamins, Dristan, Bufferin, Alka-Seltzer, etc.
- Item 23 Gasoline: Inquire as to whether the participant pumps her own gasoline, for example at self-service pumps, and place and X in the appropriate box.
- Item 24 Egg Consumption: Determine and record the approximate number of eggs that the participant has eaten in the past 48 hours. Again, in recording numerical entries, remember to right-adjust and enter leading zeros.
- Item 25 Hobbies: Determine if the participant pursues any of the listed avocations and place an X in each appropriate box.
- Item 26: Determine if the participant pursues any activity that includes regular use of solvent glue or model airplane cement, and place an X in the appropriate box.
- Item 27 Length of Residence in Area: Determine how many years the participant has lived in the area of interest, and record in the spaces provided. Round to the nearest year, except that if the response is less than one year record as <1 and terminate the interview; the individual is ineligible to participate further in the study. This situation should be detected during the screening process.

- Item 28 Length of Residence at Current Address: Determine how long the participant has lived at her current address; record the units in the spaces provided and place an X in the appropriate box to indicated whether the units represent days, months, or years. Use the most appropriate units and round to the nearest appropriate unit. For example, more than 28 days should be expressed in months and more than 11 months should be expressed in years. If the participant has resided at her current address for less than 12 months, but has lived in the area of interest for at least 12 months, record any pravious addresses during the preceding 12 months (city and state is sufficient) in the Comments section.
- Item 29 Cooling Appliances: Determine whether any of the indicated appliances or others, in which case specify, are used to cool the participant's home and place an X in the appropriate box(es) for all that apply.
- Item 30 Home Garden: Determine if the participant's household consumes food grown in a home garden and indicate the response by placing an X in the appropriate box. If a positive response is obtained, determine the location of the garden and record. Location could be participant's backyard, or another community, in which case specify city and state; be as specific as possible.
- Item 31 Commercial Food Source: Determine where the participant's household usually obtains fruit and/or vegetables and record.

Again, be as specific as possible. For example, if
the city or town has more than one store by the same name,
the store name alone would not be an adeuqate answer; as a
matter of course, record the name and location of the store,
market, or vendor.

- Items 32-34 Water Sources: In Item 32, try to determine the primary source of drinking water for the participant's household and place an X in the appropriate box. In Item 33, determine if the same primary drinking water source indicated in Item 32 is used for drink mixes such as coffee and tea; if it differs, indicate how. In Item 24, try to determine the primary source of water for cooking in the participant's household and place an X in the appropriate box. For example, some households in some areas of the country use bottled water for drinking and drink mixes but tap water (from whatever source) in cooking.
- Item 35 Other Household Tobacco Use: Inquire as to whether

 other members of the participant's household smoke, and place
 an X in the appropriate box; if YES, determine if the other

 members smoke cigarettes, cigars, a pipe, etc. and place an

 X in each appropriate box.
- Item 36 Occupation of Other Household Members: Determine if any other members of the participant's household work at any of the listed occupations or businesses, and place an X in each appropriate box.

Item 37 - Hobbies of Other Rousehold Members: Determine if any other members of the participant's household pursue any of the listed avocations, and place an X in each appropriate box.

Respondent/Interviewer Information

- Item 38 Respondent: Indicate, by placing an X in the appropriate box, whether the person who served as the primary respondent was the participant or some other person, in which case specify in the space provided.
- Item 39 Interviewer Number: Enter your assigned 3-digit

 Interviewer identification Number.
- Item 40 Date of Interview: Enter the date (month, day and year)
 that the interview was conducted and the questionnaire completed.
- Item 41 Interviewer Name: The name of the person administering the questionnaire should be printed in the space provided.

Sample Information

- Item 42: Indicate, by placing an X in the appropriate box, whether or not a milk sample was collected; if not, explain in the Comments section below.
- Item 43 Date and Time of Milk Sample Collection: If a milk sample is collected, record the date (month, day and year) and approximate time (using a 24-hour clock) of such collection.

 The time should correspond to the time that collection was completed; on a 24-hour clock, add 12 to the p.m. hours e.g., 1:00 p.m. would be 13:00, 5:30 would be 17:30, etc.

Item 44 - Date Shipped to RTI: Record the date (month, day and year) that the respective milk sample was shipped to RTI, or turned over to an RTI representative.

3.7 Collection of the Milk Sample

3.7.1 General Remarks

As indicated in section 1.0 above, the milk samples are being collected for chemical analysis by RTI as part of an EPA study to measure pollutant levels in human milk and evaluate the utility of using this body fluid in specific pollutant studies. The chemical compounds for which the samples will be analyzed are present in extremely low levels, so the utmost care and cleanliness must be used to prevent either contamination or loss. The instructions below are designed to preserve the integrity of the sample and should be followed precisely.

3.7.2 Sample Collection Instructions

- The bottles provided have been thoroughly cleaned and should be kept tightly closed, except during sampling;
 do not wash or otherwise clean them.
- Remove the MILK SAMPLE label from the sheet of labels
 in the appropriate SQ and place on one of the collection
 bottles.
- 3. The milk should be manually expressed directly into the the bottle; do not use breast pumps or other devices as the plastics in such devices would contaminate the sample. Hands should be cleaned and thoroughly rinsed to remove any residual soap; do not use rubber gloves.

- 4. Collect as much milk as possible. Unless the mother has recently nursed her infant, at least half a bottle should be easily obtainable. Less than half a bottle is unuseable and does not constitute a sample. The ability of the participant to provide an adequate sample should be determined during the screening process.
- Immediately cap the bottle and double check to see that the study numbers on the bottle and questionnaire match.
- 6. The milk sample should be immediately frozen following collection and remain so until shipping.
- Note any deviations from this procedure in the Comments
 section of the appropriate SQ.

3.7.3 Shipping Instructions

- 1. Pack the container as it was received.
- 2. Fill the can with dry ice.
- 3. Make sure that there is adequate padding to prevent breakage, that all excess space is filled with packing material.
- 4. Fill out enclosed Federal Express forms, attach to the outside of the box, and seal the box.
- 5. Call Federal Express and have them pick up the package.
- 6. When Federal Express picks up the package, call Dr. Mitch Erickson at RTI (see below) to notify him that Federal Express has picked up the package; if Dr. Erickson is out, leave an appropriate message with his secretary.

- Mail the corresponding questionnaires to RTI in one of the envelopes provided.
- 8. When the questionnaires are in the mail, call Ben Harris at RTI (see below) to notify him that the questionnaires are in the mail; if Mr. Harris is out, leave an appropriate message with his secretary.

4.0 Confidentiality

All survey research conducted by RTI is based on highest ethical standards, including those related to confidentiality. These standards are applied from the earliest steps of deciding whether or not RTI should participate in a proposed survey to the final steps of analyzing and reporting the information obtained. Strict precautions must be observed at all times to protect the rights of those whom we interview or about whom we collect data. Such precautions are built into the study design, so that promises of confidentiality and anonymity will be upheld during all phases of data handling and analysis.

No amount of effort to insure confidentiality will be successful, however, unless those responsible for data collection in the field maintain equally rigid standards, treating with utmost confidence all information offered or observed during data collection. Successful and meaningful survey research is dependent on the establishment of trust between individuals engaged in data collection and sources of information, and maintaining this sense of responsibility to the public throughout all survey activities.

Each data collector will be required to sign in duplicate a contractual agreement which includes provisions on confidential treatment of data. This agreement is designed to protect you as well as RTI and participating institutions and individuals. A copy of this agreement appears in Attachment 2.

The importance of total confidentiality cannot be over-emphasized. Any breach of confidence could result in litigation.

5.0 Contacts with Project Staff

During the data collection period it will be necessary for data collectors to maintain regular contact with RTI project staff by telephone. While you are collecting data, problems or confusing issues may arise that are not addressed in these instructions. You are encouraged to telephone RTI whenever you experience a problem or encounter a situation which you feel you cannot adequately handle.

All supplies required for data collection will be furnished by RTI.

Should you require additional supplies during the conduct of data collection, inform your RTI contact so that proper arrangements can be made. Need for additional supplies should be anticipated so that your work will not be delayed while you await recaipt of needed items. All study-related items that are in your possession at the conclusion of data collection are to be returned to RTI or disposed of according to instructions from your RTI contact.

Calls to RTI should be made between the hours of 8:30 a.m. and 5:00 p.m. (Eastern Time), Monday through Friday, to RTI's toll-free number, 800-334-8571. Request to speak to the appropriate project staff member listed below:

- Dr. Mitch Erickson Extension 6505 (regarding milk sample collection)
- Mr. Ben Harris Extension 6055

 (regarding participant selection and questionnaire administration)

If the problem is particularly acute, and you have trouble getting through on the toll-free line, call collect 919-541-6505 (Dr. Erickson) or 919-541-6055 (Mr. Harris). After 6:00 p.m. Eastern Time you may call Mr. Harris collect at work (919-541-6055) or person-to-person at home (919-942-6988).

Attachment 1
Data Collection Instruments

RESEARCH TRIANGLE INSTITUTE STUDY OF ORGANIC COMPOUNDS IN HUMAN MILK

PARTICIPANT CONSENT FORM

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	[Name of Loc	al Agency)		
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	: S. H. Harriz, III, Sun 17709, telephone numbe		earch Triangle Institu	ute. Research Triangle Park.
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STUDY OF ORGANIC COMPOUNDS IN HUMAN MILK

Spendored by:

Office of Toxic Submences Environmental President Agency Washington, D.G. 30460 Conducted by:

Remarch Triumpe Institute
P.O. Box 12194
Remarch Triangle Park, Rorth Caroline 27709

PARTICIPANT LISTING FORM

NOTICE: All information recorded on the document which would bennit identification of an instruction or an entablement will be held in strict confidence, will be used only by serable endaged in and for the purpose stated for the study, and will not be disclosed or released to other persons or used for any other surpose.

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COMMENTS



STUDY OF ORGANIC COMPOUNDS IN HUMAN MILK

Sectional by:

Office of Texts Setstances Environmental Protection Agency Washington, D.C. 20460 Conducted by:

Research Triengle Institute
P.O. Sex: 12184
Research Triengle Park, North Caroline 27709

QUESTIONNAIRE

THE RESEARCH TRIANGLE INSTITUTE OF RESEARCH TRIANGLE PARK, NORTH CAROLINA, IS UNDERTAKING A RESEARCH STUDY FOR THE U.S. ENVIRONMENTAL PROTECTION AGENCY OF LEVELS OF VARIOUS ORGANIC COMPOUNDS IN HUMAN MILK. THE INFORMATION RECORDED IN THIS QUESTIONNAIRE WILL SE HELD IN STRICT CONFIDENCE AND WILL BE USED SOLELY FOR RESEARCH INTO THE EFFECTS OF ENVIRONMENTAL FACTORS ON PUBLIC HEALTH. ALL RESULTS WILL BE SUMMARIZED FOR GROUPS OF PEOPLE; NO INFORMATION ABOUT INDIVIDUAL PERSONS WILL BE RELEASED WITHOUT THE CONSENT OF THE INDIVIDUAL. THIS QUESTIONNAIRE IS AUTHORIZED BY LAW (P.L. 94-469). WHILE YOU ARE NOT RECUIRED TO RESPOND, YOUR COOPERATION IS NEEDED TO MAKE THE RESULTS OF THIS SURVEY COMPREHENSIVE, ACCURATE, AND TIMELY.

Study number:	Site number:	Partitipant number:	
	رب		ليستنسا

First, I would like to ask some general questions about you.
1. Rese: 1 Historia 2 American Indian/ 3. When is year birthdoon!
3 Black, not of 4 Anion/Pasting (Month) (Day) (Year) Historia origin
White and of Section Other
2. What was your age in years at last birthday? Years 4. What is your approximate weight?
E. What is your height?irehusorns.
Next, I would like to ask some questions about your corupation.
6. Are you presently employed in any expectey? 1 Yes (Continue) 2 No (Co to C. 10)
7. How long have you been employed by your present employer? Units 1 Days 2 Mondas 2 Years
E. Dost your occupation usually take you away from home? 1 Yes (Contenue) 2 No (Ga so C. 11)
S. When it the nature and localitin intrest address of the company for which you work?
(Specify)
3C. If not presently employed, which of the following best describer your status?
1 Navember)
Go to Q. 181 Astired (Continue)
() Question
11. What is/was your vaud excupation? (Specify)
12. Are you presently employed in this occupation?
13. If yet to above exection, how long have you been employed in thet estuperion?
(Questions 12 and 13 mer be shipped for unemployed, retired, and disabled personal) Units 1 David 2 Months 3 Years
14. Do you work at or in any of the following possipations or establishments? (Check all sharesery.)
1 Pointing 3 Chamical plant 5 Service station/garage/angine report
3 Dry eleaning 4 Petraleum plant 4 Furniture refinishing or repair

Nec	t, it would like to ask some questions requising your health and personal habits.
15.	On you amount 1 Yes (Compleyed 2 No (Co so 4, 18)
16.	How old were you when you first started sthetung?
17.	On the overage, how many eigenesse de you amake per day?
	1 Loss than 3 park (1-4 eigensteed 4 About 1% parks (25-14 eigensteed
	2 About % pack IS-14 algorithms 5 About 2 packs (35-49 cigarothm)
	About 6 pack (18-34 eigensteel)
NCT	E: If the participant uses sebasco in some other form fother then dipersons—e.g., south, second here:
18.	What is the average number of hours that you spond out of depre each day?
18.	Now many hours of the day, on the eventer, do you normally spend away from home? (Average separately for weakness) and withhonds. Hours
20.	What do you consider the current status of your health? (Check one.)
	. I templem I acce I for I for
	Are you currently taking any prescription medication(s) on a requier delty basis?
	#ye. ###y:
	Here you taken any non-prescription medications in the past 48 hours?
	M year openifyr
je,	How many age have you estain in the past 46 hours?
	Be you surse any of the following hebbles? (Check of over apply.) 1 Furniture refinishing: 2 Pointing: 2 Soule models: 4 Gardening
L	Co you pursue any activity that includes regular use of solvent glue or model sirplane coment? 1 Yes 2 No

Les	tly, I would like to ask some quartiers about your residence and household.
27.	Now many years have you shoul in this pres? Years
21.	Months of Days You lived at your current editrant Units 1 Days 2 Months 2 Years
28.	Do you coal your home with any of the telloying appliances? (Cheek all that asply.)
	1 Control of conditioning 4 Window fundal 7 None of these
	2 Window air sondhronstal B Colling achieut fehial Bo Act Eller
	3 Evaporative assisted Circulating Sanial Other (Speedly)
30	Does your household grow any of its own food in a home garden? 1 Yet 2 No 2 De set know
	If you socially institute of greater
31.	Where does your household obtain fresh fruit and/or vegetables? (Seecify)
32	What at the privacy source of your vester for dranking?
	1 Bocked water 2 Top - community well 1 Top - cimem
	2 Tap - municipal supply 4 Tap - private wall 0 Do not brow
	2 Other (Specify)
33.	Is that the same primary square of water for sinuk mixer such as coffee, tas. Kool-Ald, stc?
	1 Yes 2 No If no, how does it differ? (Specify)
34,	What is the primary source of your weter for attaking?
	3 Bootled water 2 Tap - community well 3 Tap - comm
	2 Tap - enumicipal supply 4 Tap - pulvette well 9 Do not torow
	7 One Breit/
	
35.	Does anyone also in your hausehold smoke? 1 Yes 2 No 3 Do not know
	If you sheet all shor apply: 1 Cigarettas 2 Cigars 2 Plac 4 Other (Smecify)
34.	Does anything sine in your household work as any of the following conscitonations/second: (Chast all shar assay)
	1 Painting 3 Chamical plant 5 Service attation/garage/angine repair
	2 Dry stuming 4 Petrolaum plant # Perntyre refinishing or repair
37.	Does snyone also in your household purious any of she following habibles? (Check of sher apply.)
-	7 Princeng 2 Runnique refinishing 3 Seals mediate 4 Gardening
	RESPONDENT/INTERVIEWER INFORMATION
*	Respondent: 1 Participant 2 Other (Specify) (Month) (Ony) (Year)
34.	Interviews autober:
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	. SAMPLE INFORMATION
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43.	If you, down - - and some Hours : Minages
44.	Date shapped to MTI: (Day) + (Your)
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Attachment 2
Research Triangle Institute
Data Collection Agreement

	Triangle Institute	For Project		
	AGREEMENT	Project No.		
. –				
I, employee Research	of Powerforce Company, In Triangle Institute in com	c., field data collection services for nection with the project named above.		
4.	I agree to provide servi tions for project data o Triangle Institute;	ces within the guidelines and specifics- ollection activities provided by Research		
ъ.	I am aware that the rese being performed under co	erch being conducted by the Institute is neractual arrangement with		
c.	I agree to treat as confinterviews or obtained i period I am providing se	idential all information secured during h any project-related way during the rvices to the Institute;		
d. I shall at all times recognize and protect the confidentiality of all information secured while providing my services throughou the conduct of this research project;				
4.	from which all the enaly that all work for which	sy instruments completed form the basis sis will be drawn, and therefore agree I submit invoices will be of high quality roject specifications; and		
f. I fully agree to conduct myself at all times in a manner will obtain the respect and confidence of all individual whom data will be collected and I will not betray this of by divulging information obtained to anyone other than a representatives of Research Triangle Institute.				
Dated at	(City/Town)	(State)		
this		day of 19		
		Employee		
		For Research Triangle Institute		

Disposition: Original to RTI; yellow copy ratained by Employee.

APPENDIX B

SAMPLING AND ANALYSIS OF VOLATILE ORGANICS IN MILK

SAMPLING AND ANALYSIS OF VOLATILE ORGANICS IN MILK

1.0 Principle of the Method

Volatile compounds are recovered from an aqueous or solid sample by warming the sample and purging helium over it. The vapors are then trapped on a Tenax cartridge which can be introduced by thermal desorption directly into the GC/MS for analysis. This protocol is the result of extensive development efforts. (1-9)

2.0 Range and Sensitivity

For a typical organic compound approximately 30 ng is required to obtain mass spectral identification using high resolution gas capillary GC/MS analysis. Based on a 50 g milk sample, a detection limit of about 0.6 µg/kg would be possible. The dynamic range (limit of detection to saturation on the mass spectrometer) for a purged sample is ~10⁴; however, smaller samples may be purged and the upper end of the range increased commensurately.

3.0 Interferences

Two possible types of interferences must be considered: (1) material present in the sample which physically prevents the effective purge of the sample, and (2) material which interferes with the analysis of the purged sample. In the former case, several techniques have been developed to . handle such problems (e.g., foaming) by diluting and stirring the sample. The second case is minimized by the use of GC/MS for the analysis, since unique combinations of m/z and retention time can be selected for most compounds. This permits the evaluation of compounds even though chromatographic resolution is not obtained.

4.0 Precision and Accuracy

The purge and trap technique has been evaluated for a variety of matrices using model compounds which are expected to be typical of volatile halogenated compounds. (1)

The recovery of the purge step was validated using cow's milk samples spiked with ¹⁴C-chloroform, ¹⁴C-carbon tetrachloride, ¹⁴C-chlorobenzene and ¹⁴C-bromobenzene. The average recoveries were 88, 88, 63, and 35 percent, respectively. The recoveries correlate roughly with volatility (inversely with boiling point), so anticipated recovery for other compounds may be interpolated from these data.

5.0 Apparatus

5.1 Purge Apparatus

The purge apparatus is shown in Figure 1.

5.2 Sampling Cartridges

The sampling tubes are prepared by packing a 10-cm long x 1.5-cm i.d. glass tube containing 6 cm of 35/60 mesh Tenax GC with glass wool in the ends to provide support. (2,3) Virgin Tenax is extracted in a Soxhlet extractor for a minimum of 24 h with redistilled methanol and pentane prior to preparation of cartridge samples. (2,3) After purification of the Tenax GC sorbent and drying in a vacuum oven at 100°C for 2-3 h all of the sorbent material is meshed to provide a 35/60 mesh-size range. Sample cartridges are then prepared and conditioned at 270°C with helium flow at 30 mL/min for 30 minutes. The conditioned cartridges are transferred to Kimax (2.5 cm x 150 cm) culture tubes, immediately sealed using Teflon-lined caps, and cooled. This procedure is performed in order to avoid recontamination of the sorbent bed. (2,3)

5.3 GC/MS/COMP

The volatile halogenated hydrocarbons purged from water are analyzed on either an LKB 2091 GC/MS with an LKB 2031 data system or a Varian MAT CH-7 GC/MS with a Varian 620/i data system. The sample, concentrated on a Tenax GC cartridge, is thermally desorbed using an inlet manifold system. (2,4) The operating conditions for the thermal desorption unit and the analysis Tenax GC cartridges are given in Table 1.

6.0 Materials

6.1 Sampling

Clean, 120 mL, wide-mouth glass bottles with Teflon-lined caps are used for the collection of milk samples.

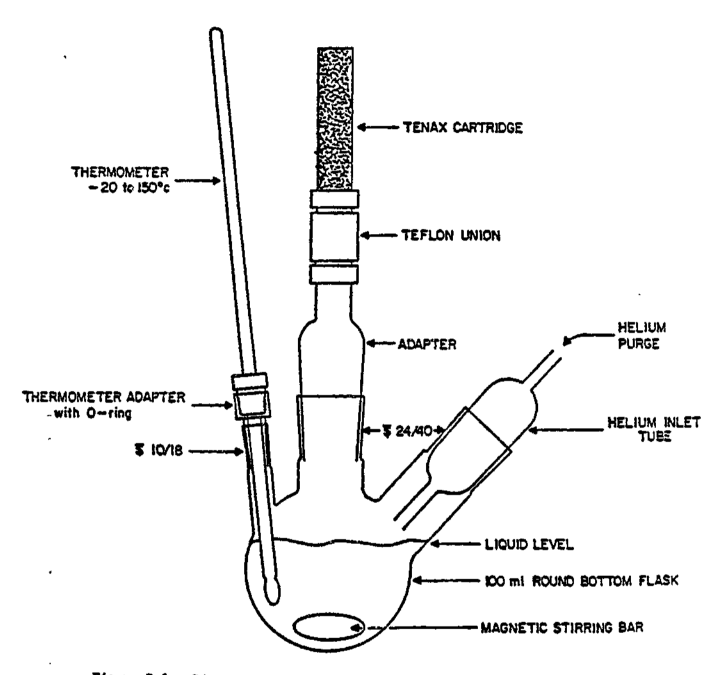


Figure 8-1. Diagram of headspace purge and trap system.

Table B-1. INSTRUMENTAL OPERATING CONDITIONS

	LER 2091	Verien HAT CH-7
Descrition chamber temperature	270	265
Description chamber He flow	15 mL/min	10 el/eio
Description time	9.0 min	9.0 mia
Capillary trap temperature during description	-196°C	-196°C
Temperature of capillary trop during injection onto column	-196°C to 250°C - the	held at 190°C
Time of He flow through capillary trap	12 3/4 min	12 3/4 min
Re flow through column [sweep time]	9.5 min	4 min
Carrier flow	2.0 mL/min	1.0 ol/mia
Capillary column	100 m SR-30 SCOT	20 m St-30 WCOT
Column temperature	30°C for 2 mim, them 4°/mim to 240°	20 + 240° at 4°/mi
Scott range	5-490 dalton	20 - 500 delton
Scan rate	2 sec full scale	1 sec/decade
Scan cycle time	2,4 sec	4.5 sec
Scan mode	perabolic	exponential
Trap current	44	
Filement current	50pA	300µA
Acceleration volstage	3,5 k¥	2kV

6.2 Purge

Tenax cartridges - 16-mm o.d. x 10.5 cm glass tubes filled with 6 cm of Tenax with 1-cm glass-wool plugs in each end.

Charcoal cartridges - 16-mm o.d. x 6 cm filled with 4 cm of charcoal and glass-wool plugs in each end.

Glass culture tubes with Teflon-lined screw caps.

7.0 Procedure

7.1 Collection of Field Samples

Milk (60-120 mL) is expressed directly into the wide-mouth bottle, capped tightly, and frozen for shipment and storage. To preserve the integrity with respect to volatiles, handling and transfer must be minimized.

7.2 Purging of Volatiles

The apparatus is assembled as depicted in Figure 1, including the Tenax GC cartridges (1.5-cm diameter x 6.0-cm length). A carbon cartridge 1.5-cm diameter x 4.0-cm length is connected to the effluent end of the Tenax cartridge to prevent contamination of the cartridge by laboratory vapors. The milk sample is cooled to ~4°C, shaken vigorously and 100 mL diluted with 350 mL distilled water. The pH of the solution is adjusted to 4.0 with sulfuric acid. A glass-wool plug is inserted into the center neck of the flask just above the level of the solution and, with the flask in a heating mantle, the solution is heated to 70°C while it is stirred with a magnetic stirrer. The sample is purged at 15 mL helium/min and 70°C for 90 minutes. The loaded cartridge is removed and stored in a culture tube containing 1-2 g CaSO₄ desiccant for 2-12 h. The desiccant is removed from the culture tube and the dry, loaded cartridge stored at ~20°C.

7.3 Analysis of Sample Purged on Cartridge

The instrumental conditions for the analysis of volatile compounds of the sorbent Tenax GC sampling cartridge are shown in Table 1. (2-9) The thermal desorption chamber and six-port valve are maintained at 270°C and 200°C, respectively. The helium purge gas through the desorption chamber is adjusted to 15-20 mL/min. The nickel capillary trap at the inlet manifold is cooled with liquid nitrogen. In a typical thermal desorption cycle a sampling cartridge is placed in the preheated desorption chamber and helium gas is channeled through the cartridge to purge the vapors into the liquid

nitrogen cooled nickel capillary trap. After desorption the six-port valve is rotated and the temperature on the capillary loop is rapidly raised; the carrier gas then introduces the vapors onto the high resolution GC column. The glass capillary column is temperature programmed from 20°C to 240°C at 4°/min and held at the upper limit for a minimum of 10 minutes. After all of the components have eluted from the capillary column, the analytical column is cooled to ambient temperature and the next sample is processed.

7.4 Quantitation

All data are acquired in the full scan mode. Quantitation of the halogenated compounds of interest is accomplished by utilizing selected ion plots (SIPs), which are plots of the intensity of specific ions (obtained from full scan data) versus time. Using SIPs of ions characteristic of a given compound in conjunction with retention times permits quantitation of components of overlapping peaks. Two external standards, perfluorobenzene and perfluorotoluene, were added to each Tenax GC cartridge in known quantities just prior to analysis. In order to eliminate the need to construct complete calibration curves for each compound quantitated, the method of relative molar response (RMR) is used. In this method the relationship of the RMR of the unknown to the RMR of the standard is determined as follows:

$$RMR_{std} = \frac{A_{unk}/moles_{unk}}{A_{std}/moles_{std}}$$

$$RMR_{unk/std} = \frac{A_{unk}/g_{unk}/GMW_{unk}}{A_{std}/g_{std}/GMW_{std}}$$

where A = peak response of a selected ion,

std = standard

unk = unknown

g = number of grams present, and

GMW = gram molecular weight.

Thus, in the sample analyzed:

$$g_{unk} = \frac{(A_{unk})(GMW_{unk})(g_{std})}{(A_{std})(GMW_{std})(RMR_{unk/std})}$$

The value of an RMR is determined from at least three independent analyses of standards of accurately known concentration prepared using a gas permeation system. (3) The precision of this method has been determined to be generally ±10 percent when replicate sampling cartridges are examined.

8.0 References

- Michael, L. C., M. D. Erickson, S. P. Parks, and E. D. Pellizzari, Anal. Chem., <u>52</u>, 1836-1841 (1980).
- Pellizzari, E. D., "Development of Analytical Techniques for Measuring Ambient Atmospheric Carcinogenic Vapors," Publication No. EPA-600/2-76-076, Contract No. 68-02-1228, 185 (November 1975).
- 3. Pellizzari, E. D., "Development of Analytical Techniques for Measuring Ambient Atmospheric Carcinogenic Vapors," EPA 600/2-75-075, 187, (November 1975).
- Pellizzari, E. D., J. E. Bunch, R. E. Berkley and J. McRae, Anal. Chem., 48, 803 (1976).
- 5. Pellizzari, E. D., J. E. Bunch, B. H. Carpenter and E. Sawicki, Environ. Sci. Tech., 9, 552 (1975).
- Pellizzari, E. D., B. H. Carpenter, J. E. Bunch, and E. Sawicki, Environ. Sci. Tech., 9, 556 (1975).
- 7. Pellizzari, E. D., Quarterly Report No. 1, EPA Contract No. 68-02-2262, February, 1976.
- Pellizzari, E. D., J. E. Bunch, R. E. Berkley and J. McRae, Anal. Lett., 9, 45 (1976).
- Pellizzari, E. D., Analysis of Organic Air Pollutants by Gas Chromatography and Mass Spectroscopy. EPA-600/2-79-057, 243 pp., March, 1979.
 Protocol Prepared, June, 1980

APPENDIX C ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS IN MILK

ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS IN MILK

1.0 Principle of the Method

Milk samples are collected from nursing mothers and frozen until ready for analysis. An aliquot of the thawed sample is then extracted, cleaned up by Florisil column chromatography and analyzed by GC/MS/COMP.

The extraction procedure used here is preferable to that used by the AOAC⁽¹⁾, since both polar and nonpolar compounds are extracted from the milk. The AOAC method is designed for pesticide residues and would not efficiently extract polar and/or acidic compounds.

Open column chromatography is a necessary prerequisite to GC/MS/COMP analysis. Although some loss of sample may occur during the extraction and cleanup, these procedures remove proteins and fats from the sample which would otherwise create overwhelming interferences for GC/MS/COMP analysis.

Since the compounds of interest in these fractions cover such a broad range of volatilities, the GC/MS/COMP analysis can be rather complex. The higher PBBs of interest in the extracted fraction must be chromatographed on a very short column (45 cm x 0.2-cm i.d., 2 percent OV-101 on Gas-Chrom Q) at high temperatures to elute them as sharp peaks which may be identified and quantitated. These chromatographic conditions are not applicable to more volatile compounds since they are not resolved from the solvent. Thus, the extracted fraction is analyzed a second time using a nonpolar SCOT capillary column (either OV-101 or SE-30 liquid phase) to separate and identify semivolatile constituents (e.g. chlorobenzenes, PCNs, pesticides, etc.). The chromatographic conditions are typically 60°C initially, programmed to 240°C (or the column limit) at 6°/min.

The mass spectral data are stored on magnetic tape. The mass spectra of interest will be printed out by the instrument operator for qualitative analysis. Quantitation from this data may be achieved by integrating the area of selected ions and comparing them to the area of the external standard.

The sensitivity of the determination may be significantly improved for quantitative purposes by using the technique of selected ion monitoring (SIM), also known as multiple ion detection (MID). This technique monitors up to 9 ions at a sensitivity 10-100 greater than the normal operating mode. This technique is used for quantitation of compounds in samples where the increased sensitivity is necessary for detection or accurate determination.

2.0 Range and Sensitivity

The detection limit of the GC/MS/COMP system has been determined to be about 5-50 ng/ μ L for pesticides such as γ -BHC, p,p'-DDE, atrazine, trifluralin and heptachlor using a 40 m SE-30 capillary column. When SIM was used, the detection limit was about one order of magnitude less (i.e., 0.5-5 ng/ μ L). The detection limit for tetrabromobiphenyl is about 1 ng/ μ L in the SIM mode using 45 x 0.2-cm i.d. column packed with 2 percent OV-101 coated on Gas-Chrom O.

For an instrumental detection limit of 1 ng/ μ L, the overall sensitivity of the method should be about 6 ng/mL (6 ppb) milk assuming a 50 mL milk sample extracted and extract concentrated to 0.3 mL. This detection limit may be improved by using SIM and may be worsened by background interferences.

3.0 Precision and Accuracy

When electron capture gas chromatography (GC/ECD) was used, the mean recoveries from cow's milk for seven replicates ranged from 57 to 93 percent for six model compounds. Thus, the results obtained may be as little as half the actual amount in the sample. The relative standard deviations (RSD) for the above replicates ranged from 11 to 33 percent, with the average RSD at 21.7 percent. Thus the precision of the method is about ± 20 percent. It is anticipated that accuracy and precision will improve with experience with the method.

4.0 Apparatus

4.1 Gas Chromatograph

A Fisher-Victoreen 4400 gas chromatograph with an ³H electron capture detector, a 10⁻¹³ AFS electrometer, and a 1.0 mV recorder is used.

4.2 Gas Chromatography Column

For most compounds, separation is achieved using a 40 m SCOT glass capillary column coated with 1 percent SE-30 and 0.32 percent Tullanox. For

the compounds of very low volatility (e.g. the higher PBBs) which will not chromatograph on the capillary column, a 45- x 0.2-cm i.d. glass column packed with 2 percent OV-101 on Gas-Chrom Q is used.

4.3 Liquid Chromatography Column

A 24-mm i.d. glass column with a Teflon stopcock is used.

4.4 Gas Chromatography/Mass Spectrometer

An LKB 2091 gas chromatograph/mass spectrometer with 2 PDP 11/4 computer is used. The system is equipped with a glass jet separator and is used with either glass capillary or packed glass column.

5.0 Materials

Kuderna-Danish evaporators:

5 mL receivers

250 mL KD flasks

Snyder columns

500 mL flat-bottom boiling flasks

250 mL separatory funnels

Clean glass wool

Whatman 1 P/S filter paper

Florisil

Sodium sulfate (anhydrous)

Acetone "Distilled in Glass", redistilled

Pentane "Distilled in Glass", redistilled

Toluene "Distilled in Glass", redistilled

Ethyl ether "Distilled in Glass"

6.0 Procedure

6.1 Extraction

- (1) Mix 50 mL (or volume available up to 50 mL) of a milk sample with clean glass wool and 150 mL of acetone to precipitate the proteins.
- (2) Decant and filter the acetone/water layer.
- (3) Repeat steps 1 and 2 with two 50 mL acetone fractions.
- (4) Concentrate to about 20 mL using a Kuderna-Danish evaporator.
- (5) Extract the precipitate with 40 mL of toluene; decant and filter the toluene layer.

- (6) Combine the toluene extract and the acetone extract with shaking.
- (7) Let the layers separate and draw off toluene (top) layer.
- (8) Repeat Steps 5-7 with 40 mL toluene and then with 10-20 mL toluene.
- (9) Discard the lower water layer.
- (10) Dry the organic layer with anhydrous sodium sulfate and concentrate to desired volume using a flat-bottom boiling flask and Snyder column. Quantatively transfer to a vial and concentrate to 5-10 mL under a gentle stream of nitrogen.

6.2 Florisil Column Chromatography (1)

- (1) Prepare Florisil by heating to 130°C for at least 5 hours.
- (2) Prepare a 24-mm i.d. column so that the Florisil is 10 cm high after settling.
- (3) Place about 1 cm of anhydrous sodium sulfate on top of the Florisil.
- (4) Rinse column with 40-50 mL pentane, never allowing the solvent to go below the Na₂SO_L layer, as channeling may result.
- (5) Add up to 10 mL of sample to column.
- (6) Elute with 200 mL of 6 percent ethyl ether/pentane solution at ≤5 mL/min.
- (7) Collect and concentrate in a Kuderna-Danish evaporator.
- (8) Evaporate under nitrogen stream to ~ 1.5 mL. Quantitatively transfer to a vial, store in a freezer.
- (9) If sample solidifies after concentration, repeat the Florisil cleanup (Steps 1-8).

6.3 Standards

Standards are spiked into the sample following the extraction and workup (d_{10} -pyrene was used at 200 ng/mL).

6.4 Analysis

6.4.1 GC/MS/COMP Analysis for Semivolatiles

Inject 0.2 μ L onto a 40 m SE-30 SCOT capillary at 60°C initially, program at 6°/min to 240°C, then hold until no more peaks are observed. Collect mass spectral data at 2 sec/scan from m/z 20-500. Compounds amenable to this analysis include organic compounds with volatility lower than that for purgeable compounds. Only the very low volatile compounds (e.g. higher PBBs) will not elute from the capillary.

6.4.2 GC/MS/COMP Analysis for Low Volatile Compounds

6.4.2.1 Normal Procedure

Inject 1.0 µL onto a 45 x 0.2-cm i.d. glass column packed with 2 percent OV-101 on GasChrom Q at 220°C initially, program to 300° at 12°/min and hold until all peaks have eluted. A helium flow rate of 20 mL/min is used. The mass spectrometer is scanned from m/z 20-1000 at 2 sec/scan.

6.4.2.2 Alternate Procedure

Using the same chromatographic conditions analyze the sample by SIM. Preselect up to 8 ions characteristic of the compound(s) of interest and one ion characteristic of the standard. Retention times provide qualitative identifications. Peak areas may be used for quantification as discussed below. This alternate procedure has 10-100 times better sensitivity than the full scan mode and provides faster quantitative results. The main disadvantage is that only preselected compounds may be identified.

In addition, if specific halogenated compounds are found to be present with little interference in most samples, they may be analyzed by GC/ECD. This procedure improves the sensitivity and reduces the analysis time (since GC/MS/COMP requires an offline data output). If GC/ECD is used, approximately 10 percent of the analyses are verified by GC/MS/COMP.

6.4.3 Qualitative Data Interpretation

Spectra are interpreted by visual comparison with standard spectral reference collections (2,3) where possible. Where standard spectra are not available, tentative identifications are made based upon interpretation of the mass spectrum. Where possible, the GC retention time is also used to assist in the identification procedure.

All identifications and interpretations are checked independently by other experienced chemists or spectroscopists to assure that the interpretations are correct.

6.4.4 Quantitative Analysis

In order to eliminate the need to construct complete calibration curves for each compound to be quantified, the method of relative molar response (RMR) is used. Successful use of this method requires information on the exact amount of standard added and the relationship of RMR (unknown) to the RMR (standards). In general, the RMR for a compound is determined for a



characteristic ion (parent or fragment) in its mass spectrum. The integrated ion current may also be used, but is generally less precise. The value of RMR is determined from at least three independent analyses. The method of calculation is as follows:

(1) RMR_{unknown/standard} =
$$\frac{A_{unk}/moles_{unk}}{A_{std}/moles_{std}}$$

A = peak area, determined by integration or triangulation of the total ion current or for a selected mass of each compound

(2)
$$RMR_{unk/std} = \frac{A_{unk}/g_{unk}/GMW_{unk}}{A_{std}/g_{std}/GMW_{std}}$$

A = peak area, as above g = number of grams present GHW = gram molecular weight

Thus, in the sample analyzed:

(3)
$$g_{unk} = \frac{A_{unk}/GMW_{unk}/g_{std}}{A_{std}/GMW_{std}/RMR_{unk}/std}$$

7.0 References

- Horowitz, W., ed., <u>AOAC Methods of Analysis</u>, 12th ed., Association of Official Analytical Chemists, Washington, DC. (1975).
- McLafferty, F. W., E. Stenhagen, and S. Abrahammson, ed., "Registry of Mass Spectral Data," John Wiley and Sons, New York (1974).
- Eight Peak Index of Mass Spectra. Vol. I (Tables 1 and 2) and II
 (Table 3), Mass Spectrometry Data Centre, AWRE, Aldermaston, Reading,
 RG74PR, UK (1970).

Protocol Prepared, June, 1980

APPENDIX D VOLATILE COMPOUNDS IDENTIFIED IN SELECTED PURGES OF MOTHER'S MILK

Table D-1. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 1081 (Bayonne, NJ)

hromato- rephic lesk Ho.	Eintion Temp. (*C)	Compound	Chromato- graphic Peak No.	Elution Temp. (°C)	Cospound
14	58	carbon dioxide	42	150	<u>e-octana</u>
13	56	chlorotrifluoroum thama	42A	152	tetrechloroethylene
1	61	propylese	423	152	CgH ₁₆ ispasr (tent.)
3	65		43	154	Calling isomer (tent.)
	66	C _a R _B isomer C _a R ₁₀ isomer	1 44	156	eilozasa
5	67		45	159	Call isomer (tent.)
4	73	C _e H _g isomer acetaldahyda	46	161	chlorobeasens
7 <u>4</u>	73	ecetope *	47	163	1-chlorohexane (cent.)
73	74	trichlorefluorem thene	48	166	ethylbensens
*	76		49	168	zylene isomer
	77	p-pentane	50	171	3-heptanone
7	79	1sopropasol	51	171	• •
10	-	methylene chloride	52	173	2-heptanoor
11	80	freen 113	53A	173	Styrane C.B. dames
12	83	carbon disulfide			C _p E ₁₆ isomer
13	93	ē-pat ure j	538	173	C _{pli20} isomer
14	87	cyclopentane	\$3C	174 174	g-beptenal
15	89	C4H6O2 isomer	530		mylene isomer
16	91	methyl sthyl ketone	54	175	C ₁₀ H ₂₂ isomer (tent.)
17	92	C6H12 isomer	55	178	2-acnada
18	94	herafluorobensene (int. atd.)	56	179	C ₁₀ E ₂₂ leoner
19	95	₽-passue	57	181	3-mathyl-1-iodobutane
20	96	chloreform	58A	163	isopropylbenzane
21	97	C ₇ E ₁₄ impear	583	184	Close 100mer
22	99	C ₆ R ₁₂ isomer	39	188	Cligat teomer
23A	102	perfluorocolumns (int. std.)	60A	189	C ₁₀ K ₁₆ isomer
23B	102	te thyloyclopensame	603	189	CgB ₁₆ O isomer (tent.)
24	104	1,1,1-trichlorosthame	624	191	bensaldehyde
25	105	CyB _{2A} teomer	613	191	n-propyl bensene
26	108	pensane	62	293	C ₃ -alkyl bensens
27	1.12	cyclohezane	63	194	C ₉ E ₂₀ isomer (tent.)
28A	213	ethyl winyl ketone	64	195	t _g B ₁₈ isomr
281	234	2-pentanope	63	196	C ₁₁ R ₂₄ isomer
29	715	C ₆ H ₁₂ O (teat.)	66	197	octamose isomer
3 0	216	n-pentanal	67	199	C ₁₁ E ₂₄ isomer
31 A	19	trichloroethylane	64	200	2-pentyliuren
31B	119	CyB, or CaB,O isomer	69A	201	C ₁₁ H ₂₄ isomer
32	122	h-haptene	693	202	g-octanel
33	126	CgR ₁₆ isomer	70	203	eilozana
34	129	Cylin isomer	71A	204	C ₁₀ H ₂₂ isomer
35	134	1-chloropensame	713	205	dichlorobensune
36	135	unknown	72	206	C ₂₁ B ₂₄ tecner
37	138	toluese	73A	210	C ₁₀ E ₁₄ iscent
38	143	C ₆ H ₁₂ O iscuss (tent.)	733	210	CgElé isomet (tent.)
39	145	g-berseal	73C	210	ant. hydrocarbon
40	147	Calle termer	24	211	eat, hydrocarbon

⁻ Continued -

Table D-1 (cont'd.)

Chromato- graphic Peak No.	Elucion Temp. (°C)	Compound	Garonato- graphic Peak No.	Tiution Temp. (*C)	Compound
75	212	limeese	85	240	unsat. hydrocarbon
76	215	sat. hydrocarbon	87	240	siloTene
77A	215	unsat. bydrocarboa	88	240	naphthelena
773	216	C11H24 isomer (tent.)	89	240	C ₁₀ 2 ₂₀ 0 feomer (tent.)
78	218	monochlorodecane (cent.)	90	240	g-dedecane
79A	219	c _o u _{la} o	91	240	pokaova
793	219	ecetophenone	92	240	wasst. hydrocarbon
50	223	eat. kyd ro carbou	93	240	eilozene
81	222	set. hydrocarbon	94	240	C ₁₁ E ₂₂ iscuer
82	224	2-20042026	95	240	allowene
83	225	dimethyletyrone	96	240	unknown
84	227	g-oreanal	97	240	silorane
85	230	<u>p-undacana</u>	į		

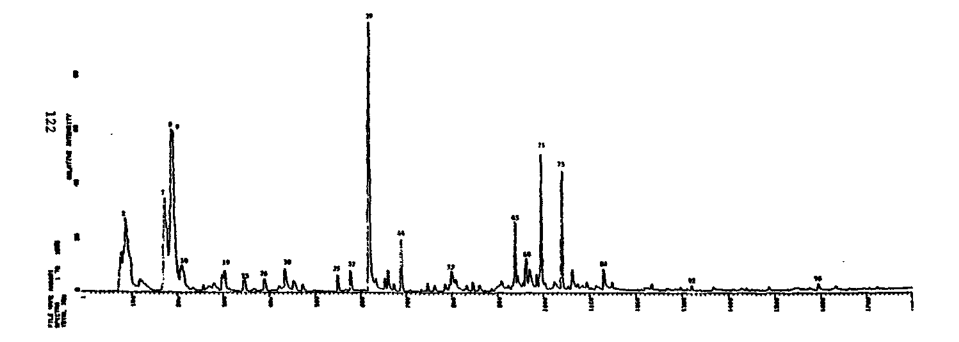


Figure D-1. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 1081 (Bayonne, NJ).

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Table D-2. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 1040 (Bayonne, NJ)

Chromaco- graphic feek No.	Elution Temp. (°C)	Compound	Chromato- graphic Peak No.	Slution Temp. (°C)	Compound
1	38	Carbon disuide	34	140	coluene
2	59	chiorotrifiuorometheme	35A	141	1-pentanol
3.	60	directly) other	350	142	wkaova
4	67	C, N,O isomer	36	145	Cyll ₁₆ incorr
5a	74	isopentane	37	146	p-berenel
50	74	trichleroflyoromethene	38	149	C ₂ B ₁₆ Asomer
SC	75	Acatona	39A	150	unknown
50	75	C _S X ₁₀ isomer	398	151	Call't recovat
4	77	g-pentana	404	152	C _E S ₁₆ isomer
65	78	Leoprese	408	153	trans-4-octans
6C	78	isopropenol	41	153	tetrachloroethylene
SD	79	C ₆ E ₁₂ isomer	42A	154	C ₃ H _{2O} isomer
6 Z	79	vicylidine chloride	428	154	set. bydrocerbon
7	81	methylene chioride	42C	154	unnet. hydrocarbon
8	82	Press 113	43	155	*
•	84	carbon disulfide	444	157	C _S H ₁₆ isomer C _S H ₁₄ isomer
10	85	2-mthylpropanal	448	157	silomene
ш	87	tyclopentane	45	161	unset. bydrocarbon
12	90	whitness	46A	162	est. hydrocarbon
13	92	eathyl ethyl ketone	468	162	unsec. hydrocarbon
14	94	Cally isomer	47	163	unkaceva
15	96	bezafluorobensene (inc. std.)	48	165	chlorobezana
16	97	S-parate (180: 200:)	49	167	etbylbinsene
17	98	chloroform	30	169	sylaps isomer
18	101	Cally locust	\$1	173	2-beptenooe
 19	104	perfluorotolusse (int. std.)	52A	174	\$177404
20A	106	1,1,1-trichlorouthage	528	175	2-g-butylfuran (tent.)
20B	107	3-eathylbutanel (tent.)	53A	175	n-haptanal
21	109	2-testylbutessl	338	176	Tylene isomer
 22	110	benzene	54	177	·
 23	111	carbon tetrachlorida	55	179	C _p E ₁₆ isomer
24 <u>4</u>	113	Cycloberane	36	181	C _y B ₂₀ isomer mat. hydrocarbon
24B	113	wethyltetrahydrofuran (tent.)	57	181	
25A	113	• • •	58A	181	C _p B ₁₈ isomer 3-methyl-1-iodobutane
25B	115	CyElia ethyl vinyl ketone	563	183	•
 26	115	2-pensanone	59A	184	C _p H _{LS} isomer isopropylbenzene
27A	117	Visyl propionets (cent.)	59B	185	est. Nydrocerbon
283	121	trichistoethylene	60	189	bydrocarbon
28A	123	C,H12 or C,H20	61	190	C ₁₀ H ₁₆ isomer
:43 :43	124	7 ² 12 4 4 4 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	62	190	"10"16 "THE CATEGORIAN CONTRACTOR
t y	127		63	191	benneldebyde
30	130	C _y H ₁₄ isomer C _y H ₁₄ isomer	4	192	2-propylbensens (tent.)
n n	132	dimerkyl disulfide	43	194	g-propylomename (cont)
12	136	1-chlotopeacane	1 #	196	isomyl formers (tent.)
33	130		,	***	

Table D-2 (cont'd.)

Chrometo- rephic Peak No.	Elucise Temp. (°C)	Compound	Chromato- graphic Peak No.	Elution Temp. (°C)	Compound
673	197	est. hydrocarbon	84	220	usktions.
68A	198	CgE ₂₀ isomer	#5	222	ace to phonone
683	199	Cy-alkyl banzana	86	223	sat. byátosárbon
69	200	est. bydrocarbon	67	225	C ₁₀ E ₂₂ income
70	201	2-pentyl furan	84	226	disethyletyrens
71	203	C ₃ -alkyl banzana	•	226	g-scassal
72	203	¢16820	904	230	eilongue
73	204	ailmanne	902	231	cilorene
74	206	dichlorobeaseae	91	234	tetramethylbenzame (tent.)
75	207	C ₃ -alkyl bensene (tent.)	92	239	&ilokane
76	209	Cat ₁₄	93	240	#110x404
77	211	disechylschylbensene isomer	94	240	naphthelene
78	212	menthese (tent.)	95	240	C ₁₂ R ₂₆ isomer
79	213	limopens	96	240	(m)coops
80	216	C ₁₁ E ₂₂ isomet	97	240	#11opana
61	216	unsat- hydrocarbon	98	240	2-thdecanons
82	217	sat. bydrocarbon	99	240	C _{2.5} 11 ₂₈
23	215	· unkoom	100	240	silozane

Figure D-2. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 1040 (Bayonne, NJ).

Table D-3. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 1107 (Jersey City, NJ)

Chronato-	flution		Chronato	Liution	
graphic	Temp.	Compound	graphic	Temp.	Compound
Peak No.	(°C)		Peak No.	(°C)	·
1	64	Tesos	203	113	vinyl propiomate
2	65	carbon dioxide	21	114	<u>e-pentanal</u>
14	67	freen 22	22A	116	C ₇ E ₇₄ isomer
31 ·	67	dichlorodifluoromethene	223	117	crichloroethylene
4	69	n-propage	22C	118	p-diprane
SA	70	butens iscent	220	116	ethyl furan (test.)
56	71	g-buteos .	23	120	n-bepcape
SC	72	acetaldehyde	24	123	2,2,4-trimethyl-1-pentens
30	73	butana isonar	25	124	Ingeredoni
6A	74	chloroethaoa	26A	125	C _e E ₁₀ 0 isomer
7	75	tetramethylailane	263	127	4-methyl-2-pentanone
BA	76	trichlorofluoromethane	26C	127	CgB ₁₆ isomer
83	78	1-pentage	27	128	dimethyl disulfide
BC	78	achtone	28	1.29	dibydropyran
PA	79	icopropanel	29	131	chloropentane
9B	79	g-yestane	30A	234	toluene
10A	87	mathylens chloride	303	137	C _a H _{lg} isomer
103	83	Press 113 .	31	139	C _c E ₁₂ 0 isomer
10C	85	carbon disulfide (trace)	324	141	g-bezanal
100	86	mathyl winyl ketone (trace)	323	143	CgH ₁₆ isomer
101	86	methyl propanol	334	146	p-octane
10F	86	nitromethane (tent.)	333	147	CgE ₁₆ isomer
214	B B	cyclopen tane	34	148	tetrachloroethylene
113	89	2-methyl pentane	35	149	CgR ₁₆ isomer
12A	90	winyl acetate	36	151	silozme
123	91	g-butsoal	37	154	white CVD
134	92	3-methyl pentane	364	156	Collin issuer
133	93	C ₆ H ₁₂ isomer	363	156	chlorobenrene
141	94	perfluorobensene (int. std.)	36C	154	2-beranel (tent.)
143	97	2-pezane	394	254	chlorobezane
140	98	chleroform	39%	159	C ₇ B ₁₂ O isomer
25	100	43bydrofuran	404	160	ethyl bengene
164	101	tetrabydrofuran	402	161	C _g R _{lB} isomer
163	102	perfluorotuluene (int. std.)	4DC	161	4-heptasone
160	102	methylcyclopentane	434	162	zylene isome:
17A	104	g-methyl scetamide	413	163	phenylacetylene
178	105	1,1,1-trichloreethans	42A	164	3-heptenope
170	106	3,3-disethylogetan (tent.)	423	165	2-heptenooe
284	106	bequepe	43	166	C ₇ R ₁₂ 0 (tent.)
163	109	carbon tetrachlorida	444	1.67	Styrene
19A	110	1-bucanol	443	168	F-pahcape;
191	110	cyclohezane	44C	168	mylane isomer
19¢	111	C3W100 1somer	434	169	sec. hydrocarbos
19 0	112	ethyl winyl ketoes (tent.)	453	170	CyR ₁₈ Secont
30A	112	2-pentacene	44	172	B-208VD4
		2-pentacone	**	114	L-sorres

Continued -

Table D-3 (cont'd.)

Chromaco- graphic reak Wo.	Elution Temp. (°C)	Coupound	Chromato- graphic Peak No.	Elucion Tamp. (°C)	Compound
47	173	sec. hydrocarbos	694	204	Calkylbensane
48	174	C ₁₀ E ₂₀ isomer	698	209	C ₁₁ H ₂₂ isomer
494	175	sat. bydrocathon	70	210	C ₁₂ H ₂₂ isomer
493	175	athyl mathyl cycloberana	72A	210	C ₁₁ R ₂₂ isomet
49C	176	unknown	713	211	phthalide (cent.)
49D	176	Cyligo leaser	72A	211	sat. bydrocarbos
SQA	176	isopropyl bentene .	723	212	decalin (tent.)
503	177	C ₁₀ F ₂₂ isomer	73A	212	ast. bydrocarbon
\$1	176	Call o isomer (tent.)	733	212	C ₁₁ E ₂₄ 1somer
S2A	181	crans-2-heptenel	73C	213	Galkylbensens isomet
523	182	d-pisese	74	213	2-20048504
52C	182	banzaldobyde	75A	214	C ₁₁ H ₂₂ 1somer
53	184	a-propylheazane	752	215	C,-alkyl bessess isomer
54	186	mylene isomer	76A	215	sac. bydrocarbon
55A	187	set. hydrocerbon	763	216	g-smenal
558	187	C ₁₀ E ₂₂ isomer	77	227	-
5 6	187	bensonitrile (trace)	78A	218	C ₁₀ H ₁₂ O isomer
57	188	sat. hydrocarbon	783	219	n-madecene
5 8 A	190	phenol	79A	220	silozane
583	190	trinethylbensene			
59	192	peacyl furan	798	220	C ₁₁ H ₂₂ isomer
60A	193		80	221	C ₁₀ H ₁₈ isomer
	193	b-octanal	#1	222	C ₄ -elkylbengene isomer
608		bensofuran	82A	223	Cl2 ^H 26 isomer
61A	194	trimethylbenzene isomere	823	224	ClaRas isomer
613	194	C ₁₀ 20 isomer	83	224	2-mechyldecalin (tent.)
62	195	SiloSane	84A	225	C12826
63A	196	c ⁷³ 300	1443	226	Cy-alkylbanzana isomer
633	196	g-decase	BAC	226	C4-alkylbensene isomer
63C	197	dicklorobenzene	85	226	silomas (tent.)
63D	196	C ₁₃ H ₂₂ isomer	BEY	228	Class inches
64A	200	telepto	963	228	C ₃₃ E ₂₀ isomer (trace)
643	20 0	trimethyl bensene isomer	86C	229	C ₁₂ H ₂₄ isomer
eac .	201	uakaova	86D	229	C ₁₀ E ₁₂ O isomer
64D	201	C ₄ -elkylbensene	962	229	Closis temes
64E	201	sat. hydrocarban	867	230	unknown
65	202	C ₁₁ E ₂₅ isomer	86G	230	C ₁₁ E ₁₆ isomer
66A	203	aat. bydrocarbon	87	230	eilozana
463	203	14monenie	[#4	230	eat. hydrocarbon
66C	204	C ₁₁ E ₂₂ isomer	89	230	sat. hydrocarbon
66D	204	sethyl styrune	90	230	est. bydrocarbon
67A	205	sat. hydrocarbon	91	230	eat. bydrocarbon
473	206	C ₁₁ E ₂₂ incust	92	230	asphthelene
67C	206	disthylbenzens isomer	93	230	wasst. kydrocarbon
68A	207	set. hydrocerbon	94	230	<u>p</u> -dodecane
68B	267	acetophenone	95 '	230	sat. bydrocarbon

⁻ Continued -

Table D-3 (cont'd.)

Chromato- graphia Mak Ho.	flution feep. (°C)	Compound	Chromato- graphic Peak No.	Elution Temp. (*C)	Compound
96	230	cilozane	105	230	unkaowa
97	230	2-vodscanone	106	230	2-tridecenone
98	230	sat. hydrocarbon	107	230	est. hydrocarbon
99	230	anknove	108	230	silozana
100	230	ellomane	109	230	phthalate
101	230	est. hydrocarbon	110	230	lactone impmer (tent.)
102	230	gakh@Ws *	111	230	diisobutyrate isomer
103	230	diphenyl ether	112	230	C ₁₄ H ₂₂ C isomer
104	230	sat. hydrocerbon			47 44

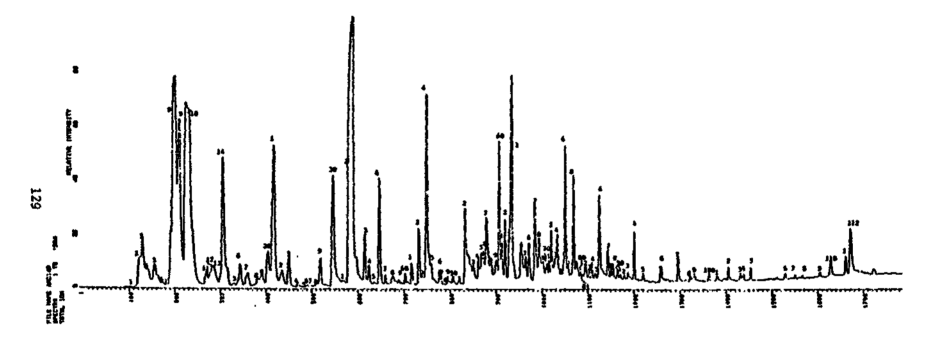


Figure D-3. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 1107. (Jersey City, NJ).

Table D-4. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 1115 (Jersey City, NJ)

Chrometo- graphic Peak Mo.	Elucios Temp. (°C)	Compound	Chromate- graphic reak No.	Klutton Youp. (°C)	Compound
34	62	carbon dioxide	24	120	C,K, isomer
18	63	Espon (trace)	25	122	dimethyldiculfide
2	65	carbonyl sulfide (teat.)	26	122	dibydropyron
34	67	chloromethene	27	124	chioropeo tana
33	68	unkaoun	28	126	anksows
44	76	Trichlorofluoromechame	29A	128	toluene
48	76	acatome	29B	129	1-pentanol
34	77	imprepriane	1 30	131	4-actby1-2-pencanone
SB	78	14opropagal	n	134	g-batena)
6A	80	mothylene chloride	324	136	CgH ₁₆ isomer
61	81	Praon 113	323	137	furaldahyde (tent.) (trace)
6C	82	carbon disulfida (crane)	33	136	proctane
6D	82	unknown	344	140	tettachlorosthylens
7	83	white	343	140	dichleropropage (trace)
8.A	86	cyclopentane	346	341	unknown
83	87	• · • · · · · · · · · · · · · · · · · ·	354	142	
8C	35	methyl isopropy) katoma	353	142	C ₅ H ₆ N ₂
9 9		g-buteral	× ×		C _g E ₁₆ isomer
-	90	1-berges (tent.)	374	143	silotane
104	92	hexafluorobensane (int. std.)	372	146	2-beansl
103	92	<u>p-bezane</u>	ş	147	chlorobensens
114	94	chloroform (trace)	384	148	CgE ₁₄ isomer
113	94	mechyl futen	388	149	5-methyl-3-hydrofuran-2-one (text
1.2	96	wasat. bydrocarbos	39	151	o-furfuryl elcobol
13	98	perfluorateluene (int. std.)	40	151	ethylbensene
144	99	crotonaldebyde (tent.)	414	132	C _g H ₁₈ isomer
141	100	1,1,1-trichloroschens	418	152	C4R4H2O (tent.)
14C	100	3-sethylburenal	424	153	mylane isomer
15	102	2-methylbutanal (cent.)	423	153	bpenklacechiese
164	104	Pensahe	42C	155	5-methyl-3-bensoone
163	105	carbon tetrachioride (trace)	43A	155	2-beptenose
L6C	105	l-butanol (tent.)	433	156	c ₇ 8 ₁₂ 0
17	106	NEW PORTS	444	157	C _p E ₂₀ (trace)
LBA	107	ethyl Viayl hetone	443	158	SETTEDS
183	107	2-pentanone	44C	158	n-beptacal
19	108	winyl propionata	44D	159	zyleme isomer
AO.	109	p-pentanel	45	159	C _p R ₁₈ incomer
103	110	sat. hydrocarbon	46	160	2-furyl methyl hetone (tent.)
:OC	110	machylberese (tent.) (trace)	47	162	g-contox
1.4	111	1-hexene	48	165	iodopentane
13	112	trichloroethylene	49	166	unknown
120	112	ethylfuren (tent.)	50	170	trans-2-heptenal
2≜	114	2,5-dimethylfuren	51A	171	banzaldshyds
23	114	g-beptane	51%	172	5-mathyl-2-ferfural
20	115	C.E. Mount	SIC	172	without
42	116	tingsports	510	173	n-propylbensens
33	117	C ₅ E ₆ E ₂ (tent.) (trace)	52A	174	mylene isomer

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Table D-4 (cont'd.)

Chromaco- graphic Peak No.	Elution Temp. (°C)	Compound	Chromaco- graphic Peak No.	Elution Temp. (°C)	Compound
529	175	bensoniertle	72A	199	2-100280000
52C	175	octanona	713	200	dimethylstyreme (trace)
52D	175	C ₁₀ H ₂₂	nc	200	C_a-elkylbensene (trace)
52E	176	C ₃ -alkylbenzene	720	200	CloH160 isomer
53A	176	1-chloro-3-sthylbanzame (tent.)	72	202	3-conenal
53B	176	dibromodichloromethane (tent.)	73	204	undecape
53C	176	phenol	74	212	unent. bydrocerbon
530	177	sat. hydrocathon	75	213	C ₁₀ H ₁₈ O isceer
53E	177	5-methyl-3-heptamone (tent.)	76A	214	g-pentylbensene
53F	177	wakaowa	763	215	silosAne
54	178	6-methyl-2-heptanone	77	216	sat. bydrocarbon
55	180	pentyl furan	76	218	2-decanone
\$6	180	g-octanal	79A	229	amphthalane
57A	181	bensofuran (trace)	79%	220	C ₁₂ B ₂₂ isomet
578	182	C3-alkylbentene	80	221	g-decanal.
57C	162	Clost leaser	81	223	g-dodecane
57D	182	Cylino incomer	#2	225	est. bydrocarbon
58	182	silomane	83A	226	Suppose
59	184	<u>g-decase</u>	833	227	methyl cianoline (tent.) (trace)
60	184	dichlorobenzene	84	228	lactone isomer (tent.)
61	187	C9H16 -	85	231	oxygenated bydrocarbon
62A	168	C ₆ -alkylbensens	86	233	phenyl hexane
628	166	phenylacesaldehyde	87	237	CloH160 (tent.)
52C	186	C ₁₀ H ₂₀ isomer	68	238	unknove
63A		Linonene	89	239	undecane
63B	190	1,6-citeols	90	240	C ₁₀ E ₁₆ O (tent.)
63C	191	C ₁₀ H ₁₈ (trace)	91A	240	unknown
64		unset. bydrocerbos	913	240	silozane
65A	192	est. bydrocarbon	92	240	unest, hydrocarbon
633	193	acacophanose	93	240	sac. hydrocarbon
66A	194	g-butylbensene (tent.)	94	240	2,2,4-trimethylpenta-1,3-diol
66B	195	C7E802 (tent.)			di-isobutywaza (BEG)
67		C ₁₁ E ₂₂ isomer	95	240	set. bydrocarbog
68		walknows	96	240	C ₁₄ 8 ₃₀ isomer
69	197	unimova	97	240	unset, hydrocarbon
7QA	198	CloHis isomer	98	240	set. hydrocarbon
706	196	sat. hydrocarbon	99	240	C ₁₅ H ₃₂ isomer
			100	240	set. hydrocarbon

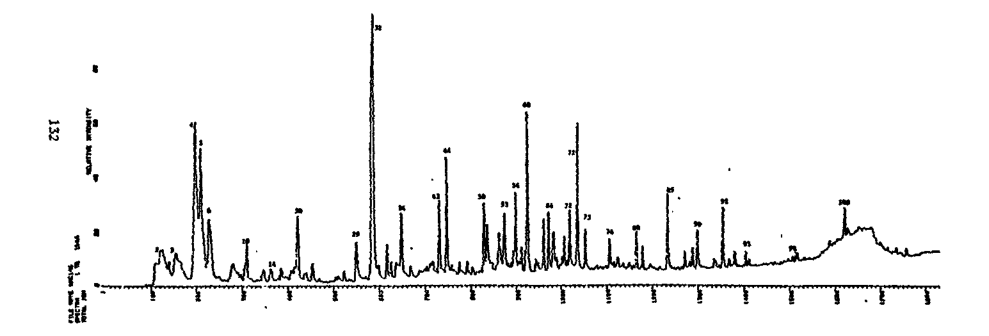


Figure D-4. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 1115 (Jersey City, NJ).

Table D-5. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 2048 (Pittsburgh, PA)

tabhte prometo-	flution Temp.	Compound	Chronato- graphic	Elucion Temp.	Compound
eek No.	(†c)		Peak No.	<u>्ट्</u>	
1A	58	carbos dioxida	334	145	tetrachlorouthylana
19	58	chloretrifluoromethana	332	146	Call ₁₆ isomer
2	64	C ₄ Z ₃ isomer	34	147	C ₇ E ₁₄ O isomer
3	66	C _i E ₁₀ isomer	35	149	silcutos
44	70	acecaldebydo	36A	153	C ₆ H ₁₂ O isomer
43	70	C ₅ H ₁₂ isomer	343	154	oproperses (tiets)
SA	72	trichlorofluoromethene	37	156	chlorohexane (trace)
52	72	ace code	38	159	erhylbensene
64	73	g-pentane	394	161	set. bydrocatbon
63	74	isopropenol	393	161	zylene inconer
7A	77	Frece 113	390	162	wksove
73 73	77	anthylene chloride	390	162	C _p B ₂₀ isomer
8	79	carbos disulfide	40	164	3-beptanone
94	83	C ₅ % ₁₀ isomer '	41	165	2-haptanona
93	63	CgR ₁₄ Leonar	424	166	etytene
10	84	C ₅ H ₂₀ O isomer (tent.)	423	167	•
11A	87	methi sthi perses	426	167	G _p E ₁₆ isomer (tent.) ent. bydrocarbon
113	87		43A	168	g-teptensl
12A	89	, C ₆ H ₁₂ isomer bezefluorobensene (int. std.)	438	168	_
128	89	g-bettens		169	sylena isomer
13	91	chloroform	44	170	g-00404
14A	96	perfluerotoluege (int. std.)			C ₁₀ E ₂₀ isomer
148	96	methyleyelopescame 'Lat. sta./	46	173	C ₁₀ H ₂₂ isomer
15A	94	1,1,1-trichlorocthese	47	175	G ₁₀ H ₂₂ isomer (tent.)
158	98	1-butanol (tent.)	444	177	teopropylbenzens
16 16	LO2	bearens	483	177	C ⁷⁰ g ⁵⁵ teomez
17	104		49	161	C ₁₃ E ₂₄ isomer
17 18a	104	eyelohezase	304	162	Close isomer
183	107	C ₆ H ₁₂ isomer	500	103	CBE160 decmer
	-	C ₅ H ₁₀ O isomer	51A	184	umset. hydrocarbon
18C	109	C ₆ H ₁₀ isomer	513	184	heazaldebyde
19 20a	109	gryentene)	310	184	a-propylbensame
20A 20B	112	trichloroethylene	52A	186	Close teomer
	112	CyH ₁₂ isomer	528	186	Gy-elkyl bensene isomet
21	115	mknorn	53	187	eat. bydrocarbon
22 	119	C7 ^H 14	ļ <u>s</u>	189	west. bydrocarbon
23	126	C ₆ E ₁₂ O issuer	55	190	C ₁₁ E ₂₄ isomer
24 <u>4</u>	126	unset. hydrocarbon	36A	190	Callió isomer
243	127	chloropentane	368	192	C ₁₀ E ₂₂ isoter
25	130	meet. hydrocarbon (tent.)	57	192	C ₁₁ 2 ₂₄ isomet
26 	131	teluene	58	194	2-yeutylfuran
27	133	1-pentanol	59	194	C _{ll} E ₂₄ isomer (cenc.)
28	134	Calla louner	60A	195	Cy-alkylbensone isomer
29	136	C ₆ H ₁₂ O isomer	603	195	C ₁₀ E ₂₀ isomer
30	138	S-pezzori	61	197	silozaca
31	140	C _p E ₁₆ incorr	62A	196	set. hydrocarbon
32	143	g-octane	623	198	dichlorobenzene .

-Continued-

Table D-5 (continued)

Chromato- graphic Peak No.	Elution Temp. (°C)	Compound	Chromato- graphic Peak No.	Elution Temp. (°C)	Compound
63A	200	unest. bydrocarbon	82	231	unsat. hydrocarbon
63B	200	est. hydrocarbon (tent.)	83	232	C ₁₁ k ₂₀ tenner
64	202	unest. hydrocarbon	84	233	C ₁₀ N ₁₈ D isomer
65	203	2-ethyl-1-hexanol	B5A	235	silomane
66A	206	limonana	8SB	236	C ₁₀ H ₁₈ O deceer
668	206	C ₁₀ H ₁₈ O isomer	#5¢	236	Closico incomer
67	208	sat. bydrocarbos (tent.)	86	236	unsat. bydrocarbon
68	209	ast. hydrocarbon	87	240	sat, bydrocathon
69	211	C _z -alkylbensene	884	240	asphthalens
70	712	acetophenons	488	240	C ₁₀ E ₂₂ O isomer (rent.)
71	213	sat. hydrocarbon	B9A	240	u-tespineol (text.)
72	214	sat. hydrocarbon	891	240	unsat. hydrocarbon
73	215	eat. hydrocarbon	90	240	esesabob- <u>e</u>
74A	216	C _L -alkylbenzené	91	240	silchane
748	217	C _g E ₁₆ O iscent	92	240	unsat, hydrocarbon
75A	218	dimethyletyrane	93	240	silowane
75B	219	est. bydrocarbon	94	240	2-undecanone
76	220	n-conanal	95	240	silomane
77	222	<u>p~mdecane</u>	96	240	C ₁₃ H ₂₅ isomer
78	223	siloxane	97	240	silowape
79	226	C ₄ -alkylbenzene	98	240	decempic sold (tent.)
BOA	226	C _Z -alkylbensens	99	240	C ₁₄ E ₃₀ isomer
808	227	unknown	100	240	unsat. hydrocarbon
81	229	est. Bydrocarbon	101	240	afloxane

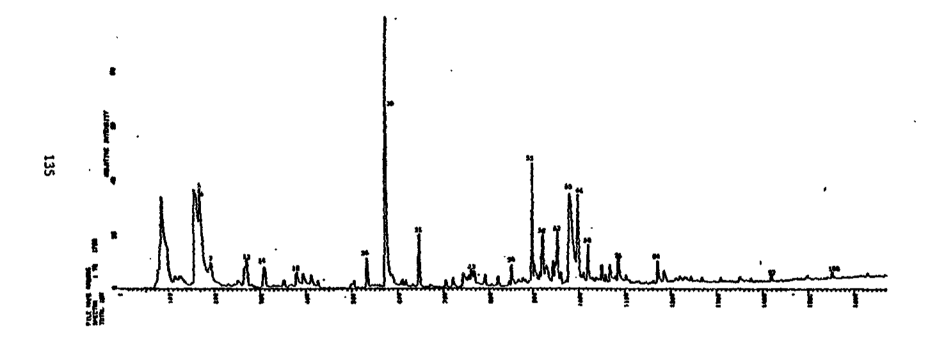


Figure D-5. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 2048 (Pittsburgh, PA).

Table D-6. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 2071 (Pittsburgh, PA)

hromato- raphic ask Ma.	Ilution Imp. (°C)	Compound	Chromato- graphic Peak No.	flution funp. (°C)	Compensed.
1		carbon dioxide	33	116	trichloroethyleng
2A	60	propylane (traca)	344	110	<u>p-haptana</u>
 23	61	dichlorodifluoromethene (space)	343	119	C ₇ E ₁₄ Security
M	62	dimethyldifluorosilaps	35	122	Call isomer
33	63	isobutane	»	124	C ₇ R ₁₄ isomer
44	64		37	226	dimethyl disulfide
	65	C ₄ H ₀ incomer	38	127	myswo
·-	66	E-parame (trace)	39	129	
	68	Chicrosthese (trace)	40	133	C ₇ E ₁₄ imomer (cent.) tolumne
,	71	methanol	41	138	
	73		42	139	dibromochloromethane (trace)
.	-	Acetone			g-barres)
35	73	trichlorofluorame thans	43	141	Cagle toomer
)A	75	isopropenol	44.	144	D-octube
73	75	E-pagtane	454	145	tetrachloroetbylene
ec •	76	C ₅ H ₈ isouer	453	346	CgE ₁₆ isomer (rent.)
LO	77	C ₆ B ₁₂ isomer	46	347	takeom
.14	78	methylene chicyide	47A	149	ansat. bydrocerbon
110	79	2-methyl-2-propanol	473	149	ailozape
TC	80	Press 213	1 44	152	C ₅ E ₁₈ isomer
2	61	c ₆ 14	49	153	chlorobensene .
31	82	terbos disulfida	50A	158	achylbensone
.3B	83	C ⁴ B ⁶ O	503	159	Comer 18 190mer
4	85	p-propanol (tent.)	514	160	zylene isomer
SA.	86	Cyclopestane	51B	160	phenylacocylene
53	87	C ₆ 2 ₁₂ isomer	S2A	162	3-heptanone
6	87	C ₆ H ₁₄ isomer	528	163	2-keptanone
7	88	Visyl scetate ·	53	164	OT/TODA
•	89	h-butenal	54	166	kylone isoser
9	90	methyl ethyl hatone	55	167	<u>p-beptanel</u>
9	91	CgE12 isomer	56	169	2-eccess
1	93	hexafluorobensene (int. acd.)	57	170	Clows isomer
2	94	F-pezene	58A	173	imopropylbenzene
34	94	athyl scetate	389	174	C ₁₀ E ₂₂ isomer
33	95	chloroform	59	176	C ₁₀ E ₁₆ tecmer
4	96	C ₇ E ₃₄ isomer	60	177	C ₁₀ E ₂₀ isomer
3A	100	perfluorocoluene (int. std.)	61	179	n-piasus
55	100	mathylcyclopentens	62A	180	bensaldebyde
5	101	C_P_A isomer	623	180	g-propylbensens
7.6.	102	1,1,1-srichloresthene	634	182	C ₁₀ H ₁₆ isomer
13	103	C ₅ k ₂₀ 0 isomer (tent.)	633	182	C'-eykAlpenzene
: ,	106	bensene	44	184	trimethylbenzene isomer
, '	107	carbon tetrachloride (trace)	65	183	C ₁₀ E ₂₂ irener
DA	108	g-butanol (tent.)	66	185	bentouitrile
DB.	106	eyeloberane	67A	184	methylheptanoon isomer
ı	111	mathyl propyl hacone	673	186	0-sethylutyrese
ž	113	p-pentanal	664	187	triunthylbengame isomer

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Table D-6 (continued)

Chromeco- graphic Peak No.	Elution Yearp. (*C)	Compound	Chromato- graphic Peak No.	Flution Temp. (*C)	Compound
463	(87	set. hydrocarbon	43	211	dimethylszyrene
692	188	ethyl g-caproate	1 44	211	sat. bydrocarbon
693	166	pencylfuran (tent.)	85	212	camohena (tent.)
70A	190	bensofuran (tent.)	86	214	eilowane
703	190	Calkylbenzene	87	215	sat. hydrocarbon
70C	190	trimethylbensese isomer	68	216	methyl captylece
700	191	phenol (trace)	89	222	silomane
71	192	#110140B	90	223	camphor
72A	192	C ₁₀ H ₂₂ isomer	91	225	C ₁₀ E ₁₈ O (trace) (tent.)
728	193	dichiorobensens	92	227	silozane
72C	193	unknown	93	230	trichlorobenzane (trace)
72D	194	C _{lo} H ₁₆ isomer	94A	231	ethyl capsylate
73	194	sat. hydrocarbon	943	232	naphthalane
74	196	C ₁₀ H ₁₆ isomer (tent.)	95	235	<u>g-dodecane</u>
75A	196	C ₁₀ H ₁₆ isomer	96	239	unsar. hydrocarbon (renc.)
758	197	C _L -elkylbenzene	97	240	#110xAD#
76	199	limonene	984	240	2-undecapone
77	201	onkaowa	983	240	set. hydrocerbon
78	203	sec. hydrocarbon	99	240	sat. hydrocarbon
79A	205	scetophenone	100	240	methyl decements
79B	205 .	C ₁₀ E ₁₆ isomer	101	240	silomane
80	207	eac. hydrocarbon	102	240	C ₁₄ E ₃₀ (tent.)
81	206	unknova	103	240	ethyl decamente
82	210	2-00040000	104	240	unset. hydrocarbon
			,		

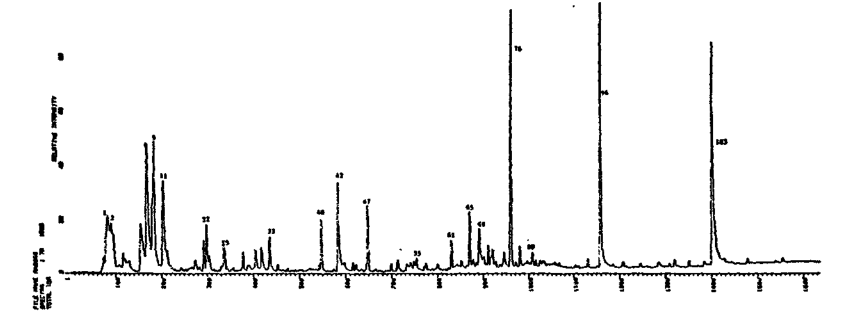


Figure D-6. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 2071 (Pittsburgh, PA).

Table D-7. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 3053 (Baton Rouge, LA)

Chromato- graphic Peak No.	Elucion Trup. (°C)	Compound	grephic Pask No.	Elector Temp. (°C)	Compound
14	57	carbon dioxide	34	136	C T A terrer
13	58	chloro trifluoromechane	35	139	C ₄ E ₁₂ O leoner g-hexanel
2.	62	chloromethane	36	141	
3	43		37	144	C _B E ₁₆ isomer
	58	C ₄ H ₁₀ isomer dimerkyldifluoromilama	38		g-octone
SA	70	ectylaldehyde	Į i	145	C ₆ H ₁₀ O isomer
53	71	·	39	147	fureldehyde inomer
3C	72	scrtone	40	149	Calle isomer
<i>.</i>	73	furen	41	150	ellozene
•	74	g-pentane	42A	153	Call isomer
7		1-propasol	429	153	C ₆ B ₁₀ O isomer (tent.)
8A 45	76	methylene obloride	434	155	C ₈ H ₁₄ facmor
41	77	Preon 113	438	156	Callino isomer
,	79	carbon disulfide (trace)	44	158	enimores
10	80	C'H ⁸ O Troner	45A	159	ethylbeasene
11	85	C ₅ II ₁₀ 0 isomer	45%	159	C ₇ H ₁₂ isomer (tent.)
11	#6	g-botanal	4SC	160	G-furfutyl elechol
134	47	methyl ethyl hetone	45Þ	161	Zylane isome:
138	86	C ₆ H ₁₄ isomer	4SE	161	C ₈ B ₁₈ isomer
144	90	hexafluorobensena (inc. etd.)	46	164	C ₇ H ₁₄ O Secret
143	90	2-methylfurau	47	165	C _g E ₁₆ isomer
14C	90	E-persus	- 48A	166	acyzens
15A	92	angulous.	483	166	C ₇ B ₁₀ O ₂ leouer
153	93	3-methylfuran	48C	167	g-hoptmanl
16	94	C ₆ H ₁₂ isomer	494	169	C ₆ H ₆ O ₂ isomer
17A	97	perfluoreteluene (int. std.)	498	170	wakanya
173	97	methyleyelopentans	30	172	<u> </u>
16	98	CARGO isomer .	51A	173	C _a H ₁₆ isomer
19	100	1,1,1-trichlorosthene	31 3	173	C _g R ₁₄ isomer
20	3.04	bensene	52	175	makaowa
21	3.06	C ₆ S ₁₂ iscorr	53	176	Cyll ₁₈ isoner
224	300	ethyl winyl ketege	54	178	C _p B ₁₈ isomer
223	106	C ₅ E ₁₀ G isomer	55	180	C ¹⁰ B ²⁰ isomer
23	209	C ₆ H ₁₂ O isomer	56	181	C _p E ₁₈ incomer (cent.)
24	1.10	g-peatunel	57A	182	merbylfuraldebyde fecuer
25A	113	C,B, isomer	573	182	benseldshyde
25B	123	trichloroschylene	58	184	mechylfuraldebyde isomer
25¢	134	C, NO	59A	186	5-btohijpergere
26A	316	P-pobtwe	593	187	Clog teomet
263	317	acetic acid	59C	188	Close isomer
17	120	2-vieylfuren	40	189	-10-22
2 8	122	CySia facent (tent.)	- u	190	C ₃₁ 9 ₂₄ fauner
29	123	Cylia Sacurer	624	190	Cylly O-isomer
50	125	dimethyl disulfide	623		C. B. Commit (seek.)
ii	126	dihydropyran (tage.)	1	191	C ₁₁ B ₂₂ isomer (tent.)
32	133	Columns (come.)	634	192	C ₁₁ H ₂₄ isomer
 33	134		638	193	2-pentylfuren
		C ₇ E _{3A} termer	64 tisued -	194	g-octanal

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Table D-7 (continued)

Chromato- graphic Peak No.	Elution Temp. (°C)	Сопромой	Chromato- graphic Pask No.	Elution Temp. ('C)	Compound
65A	194	C3-alkyl benzene isomer	878	231	C ₁₂ R ₂₄ isomer
653	194	unknova	BAL	234	Siloxese
66	196	silomas	888	234	C ₁₀ 2 ₁₀ 0 ₂ (tent.)
67A	197	<u>p-decape</u>	89	235	est. hydrocarbon
673	198	dichlorobensens	90	236	C ₁₂ H ₂₆ isomer
68	199	unant. bydrocarbon	91	237	C ₁₂ B ₂₆ isomer
69	201	C ₉ H ₁₆ isomer	92A	238	C ₁₀ R ₂₀ O isomer
70	202	Calkylbenzene (tent.)	923	239	uman. bydracarbon
71	204	CgE ₆ O ₂ isomer	934	240	naphthalana (trace)
72A	204	lisonene	938	240	C ₁₂ H ₂₂ isomer
723	204	sat. hydrocarbon	944	240	n-decever
73A	207	unsat. hydrocarbon	948	240	C ₁₂ R ₂₄ isomer
735	207	C ₁₁ E ₂₄ isomer	95	240	n-dodecane
74A	206	set. hydrocarbon	96	240	C ₁₃ R ₂₈ isomer
748	209	scetophenons	97	240	sat. hydrocarbon
75	210	Calkylbehrene	98A	240	C ₁₃ R ₂₆ isomer
76	211	C ₁₁ E ₂₄ isomer	983	240	C ₁₁ E ₂₀ O isomer
77A	212	C ₁₁ E ₂₄ isomer	99	240	C ₁₃ R ₂₈ isomer
773	212	unsat. hydrocarbon	100	240	C ₁₃ E ₂₈ isomer
77C	213	eat, hydrocathon	101	240	C ₁₃ E ₂₈ isomer
770	213	C _c H _c O ₂ isomer	102	240	C ₁₀ R ₁₆ O isomer
784	214	CyEgO, teomer (tent.)	103	240	C ₁₃ E ₂₄ isomer
783	215	C ₁₁ E ₂₄ isomer	104	240	<u>s-undecenal</u>
79	217	CloH160 isomer	105	240	p-tridecane
80	218	<u>b</u> -mosen4l	106	240	CloBlet isomer
61	221	n-undecane	107	240	siloxane
82	222	unset. hydrocarbon	108	240	unsat. bydrocatbon
83	224	est. hydrocarbon	109	240	unsat. bydrocarbon
84	226	C ₁₂ E ₂₆ isomer	110	240	n-dodecanal
8 5	227	mat. hydrocarbon	111	240	n-tetradecane
86	228	C ₁₂ H ₂₆ isomer	112	240	unsec. hydrocarbon
B7A	229	elloxans	113	240	D-beutagacava

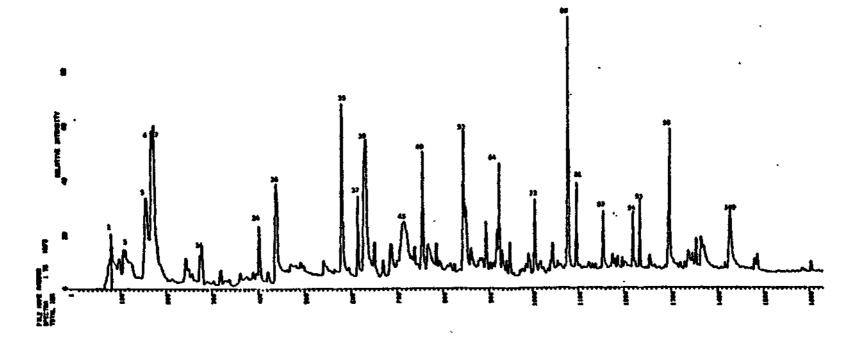


Figure D-7. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 3053 (Baton Rouge, LA).

Table D-8. VOLATILE COMPOUNDS IDENTIFIED IN PURGE OF SAMPLE NO. 3111 (Baton Rouge, LA)

Chromato- graphic Pank No.	Elution Temp. (°C)	Compound	Chromate- graphic Peak No.	Elution Temp. (°C)	Compound
1	39	carbon diexida	333	148	unsat. bydrocarbos
2	61	dichloredifluoremethane	34	150	CgE ₁₆ isomer (tent.)
34 .	65	sulfur dioxida	35	152	ellozene
38	65	C_H_ incom	36A	155	C _p E ₁₈ isomer
4	71	C ₅ R ₂₀ isomer	363	155	C _g E ₂₀ isomer (tent.)
SA	73	trichlorofluoromethane	37	161	ethylbensene .
57	74	acetosa	384	143	mylene isomet
64	76	isopropanol	363	164	C _g E ₂₀ isomer
63	76	g-pentane	39A	168	styrene
60	77	C ₅ H _g incor	393	16\$	C _g B ₂₀ isomer
74	80	methylene chloride	40	169	mylene isomer
78	81	Preon 113	41	170	Cg ^E 20 isomer
	82	carbon disulfids	42	173	C _g ll ₂₀ leoner
3	84	p-bureas)	434	177	aat. hydrocarbon
10A	87	cyclopentane	438	177	Calkyl bensens (tent.)
103	88	C ₆ N ₁₄ isomer	[44	178	C ₁₀ E ₂₂ isomer
11	69	C ₅ E ₁₀ O isomer	43	179	C ₁₀ u ₂₂ isomer
124	91	C ₅ B ₁₀ C isomer	46	181	eat. hydrocarbon
128	92	C ₆ 2 ₁₂ isomer	47	183	ailozape
13	94	bexafluorobensene (int. etd.)	46	186	bangaldehyda
14	95	B-pezebe	49	1.69	unknown
15	96	chloroform	50	189	C ₁₁ E ₂₄ isomer
164	101	perfluorotoluene (int. ecd.)	51	191	C3-alkyl benzent
163	101	me thyleyclopentane	52	192	C _{II} E ₂₄ Asomer
174	104	1,1,1-trichlorosthess	53	193	C ₁₁ E ₂₄ former
173	104	C ₅ E ₁₀ O iscour (test.)	544	194	C ₁₁ R ₂₄ former
18	106	C ₆ B ₁₂ O isomer,	543	193	Cy-alkyl bensene
19	108	beniehe	\$5A	196	ellerane
20	109	carbon tetrachloride	55B	197 198	C ₁₁ H ₂₄ isomer dichlorobensene
21 22	110 111	C_6R_{12} incomer C_6R_{12} 0 isomer (tent.)	56 57	202	C ₃ -slkyl benzene
23	112	C ₆ E ₁₂ O isomer (test.)	58	204	linosene
21	114	n-pentanel	59	206	mat. hydrocarbon
25	117	grichloroethylene	60	208	sat. hydrocarbon
26	120	p-heptene	61A	212	acatophenons
27	123	C _g E ₁₆ isomer	613	213	sat. bydrocarbon
26	126	C ₇ E ₁₄ isomer	62	214	eat. hydrocarbon
29	128	dimethyl disulfide	63	217	sat. hydrocarbos
30	135	toluens	4	221	g-undecese
314	142	n-bersasl	65	233	cilozes
31.3	164	C _a E ₁₆ isomer	66	240	g-dodecase
32	146	Successes	67	240	mast. hydrocarbon
33A	148	tetrachlorosthylene	65	240	ailogane

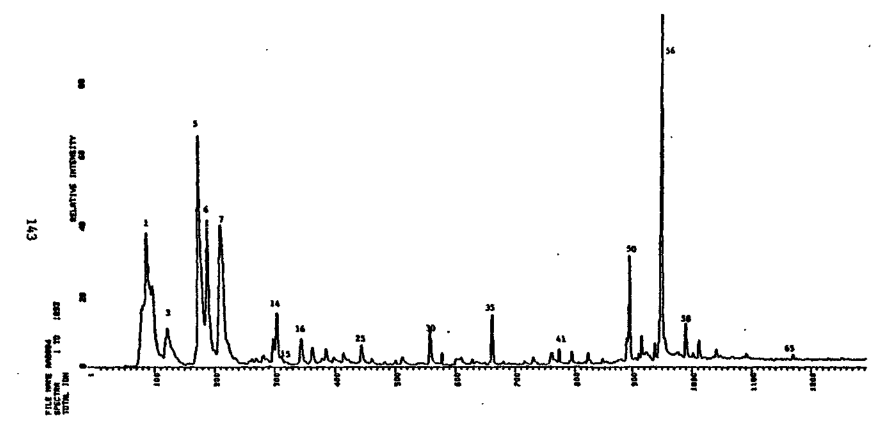


Figure D-8. Total ion current chromatogram from GC/MS analysis for volatiles in sample no. 3111 (Baton Rouge, LA).

APPENDIX E SEMIVOLATILE COMPOUNDS IDENTIFIED IN SELECTED EXTRACTS OF MOTHER'S MILK

Table E-1. SEMIVOLATILE COMPOUNDS IDENTIFIED IN EXTRACT OF SAMPLE 1032 (Bayonne, NJ)

Chromato- graphic Meak No.	Election Temp. Compound (°C)	Chromato- graphic Peak No.	llution Imp. (°C)	Compound
34	tolutes	25		wakaawa
13	zylane isomer	26		wakaowa
2	eilekan e	27		#11gEade
3	#1lexact	28		eilexans
4	stlowes	29		d ₁₀ -pyrene (std.)
\$	ailouses	30		set. and unset. bydrocarbons
6	eilensee .	31		ellozane
7	#110240e	32		200
8	atlemme	33		saknowa
•	dimethylbiphamyl (tent.)	344		silezane
10	ellouse	348		ankaova
11A	silozane	35		unknown
112	anikoone	36		set. and unset. hydroterbous
12	silozana	37		silozane
13	aat. hydrocarbon	30		sat. and unset. hydrocarbons
14	eilozana	39		set. and unset. bydrocarbons
15	ailonma	40		silouse
16	est. bydrocarbon	41		elloume '
17	est. and waset. hydrocarbons	42		silozme
18	ailoxane	43		silomana
19	ellemee	44		silozane
20	eilogane	45		silogene
21	est. bydrocarboa	46		Тусорегоеза
22	phthelate (tent.)	47		cholesteryl acatate
23	silomane	44		eilowane
24	est. and wast. hydrocarbons	ì		

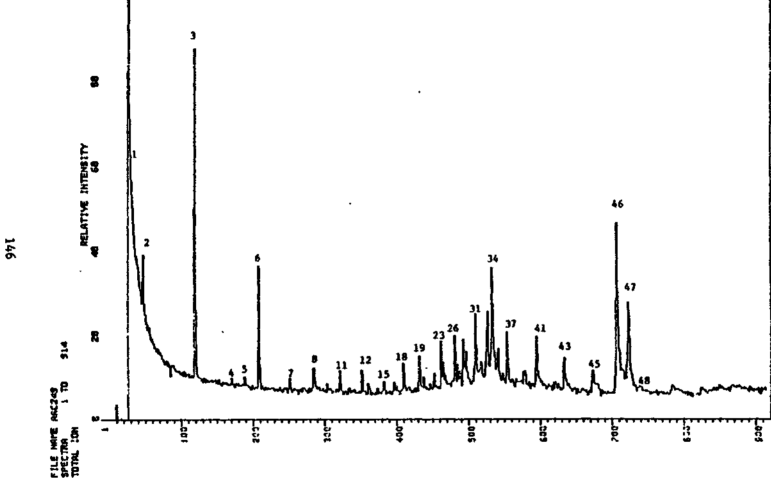


Figure E-1. Total ion current chromatogram from GC/MS analysis for Semivolatiles in sample 1032 (Bayonne, NJ).

Table E-2. SEMIVOLATILE COMPOUNDS IDENTIFIED IN EXTRACT OF SAMPLE 2121 (Pittsburgh, PA)

Chromato- graphic Pask No.	Elucion Temp. ("C)	Compound	Chromato- graphic Pask No.	Elution Yeap. (°C)	Compound
1		toluene	28		unsat. hydrocarbon
2		eilorane	29A		unsat. hydrocarbos
3		#iloxane	292		TOE
4		cilorace	30		eat. and unsat. hydrocarbons
5		eiloxane	31		eilozene
6		2,4-di-cert-busyl-4-methylphenol	32		pentachlorobiphenyl
7		methyl dodecanosts	23		agt. and uneat, bydrocarbons
•		ethyl butyrate (tent.)	34		silozana
9		eilozane	35		sat. and unsat. hydrocarbons
10		sat, hydrocarbon	36		hexachlorobiphenyl
11		edlozene	37		silozane
12		eilousce	38		est. bydrocarbon
13		aat, hydrocarbon	39		silomes
14		gilousse	40		sat. and unsat. bydrocarboos
15		ellozane	434		est. and unset. hydrocarbons
16		ellemane	418		heptachlorobiphenyl
17		sat. and unsat. bydrocarbons	42		eflowene
18		sat. bydrocarbon	43		sat. and unsat. hydrocarbons
19		wekeowe.	44		silomana
20		ailozane	45		silozane
21		set. and unset. hydrocarbons	46		ensolte
22		unknown.	47		eilowane
23		unkasen.	46		eilezane
24		ailorene	49		lycopersons
25		silozane	50		eilozace
26		d ₁₀ -pyrens (int. std.)	51		cholesteryl scetate
27		silomane			

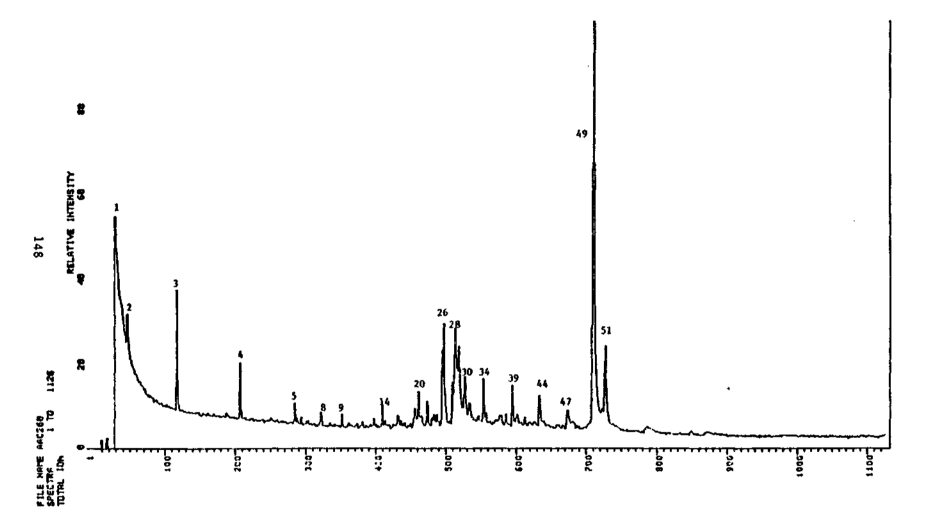


Figure E-2. Total ion current chromatogram from GC/MS analysis for semivolatiles in sample 2121 (Pittsburgh, PA).

Table E-3. SEMIVOLATILE COMPOUNDS IDENTIFIED IN EXTRACT OF SAMPLE 3095 (Baton Rouge, LA)

Chromato- graphic Peak Mo.	Elucion Temp. (*C)	Compound	Chrosato- graphic feak No.	Riution Temp. (°C)	Compound
1		methylene chloride	12		d ₁₀ -pyrese
2		toluene	334		ast. hydrocarbos
3		silozane	332		unsat. hydrocarbon
4 "		set. hydrocatbon	34		ellorane
5		sat. hydrocathon (tent.)	35		DDZ
6		silexana	36A		unknown
7		sac. bydrocurbos (test.)	363		unset. bydrocarbon
4		silorane	37A		silomane
9		sat. hydrocarbon (tent.)	378		tak potra
10		ailorage	38		set. hydrocarbon (tent.)
11		sat. hydrocarbon	39		silozane
12		sat. hydrocarboo	40		unsat. bydrocarbon (tent.)
13		myrano	41		#1loxan4
14		unkaowa	42		sat. bydrocurbon (tant.)
15		sac. hydrocarbon	43		cilorane
16		silozene	44		sat. bydrocarbon
17		est. bydrocarbon	45		set. bydrocerbon
18		silozane	464		eet. hydrodathou
19		silozens	463		silozane
20		sat. hydrocarbon	47		silomene
21		ser- phqiacaipos.	48		sat. hydrocarboc
22		silomnot	49		silozane (tent.)
23		silozane	30A		wilerane
24		#1loxage	503		sat. hydrocarbon
25		set. hydrocarbon	51		sac. hydrocarbon
26		silogane	52		lycopetaene
27A		sat. hydrocarbon	53A		Bilozane
273		unsat. hydrocarbon	532		cholesteryl scetate
28		unknown	54		ailgrane
29		wakneva	55		est. bydrocarbon
30		siloxane	56		unknown
31		siloxene	57		#110xana

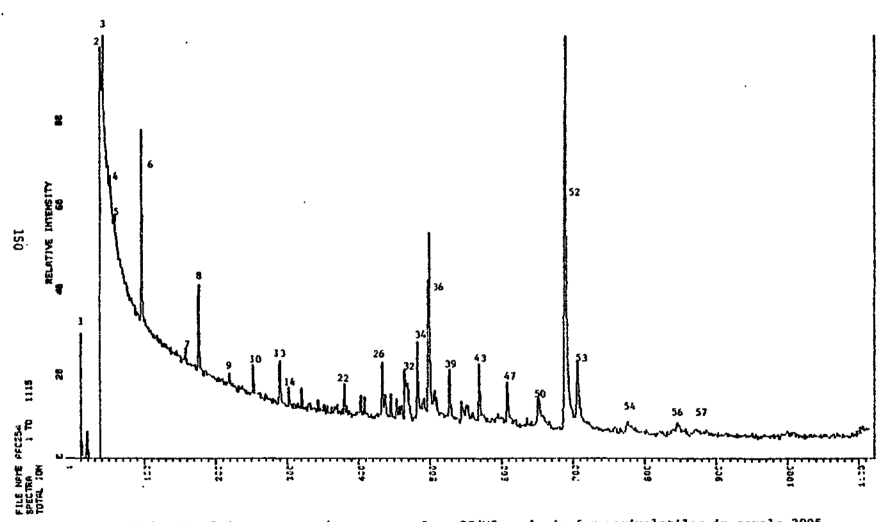


Figure E-3. Total ion current chromatogram from GC/MS analysis for semivolatiles in sample 3095 (Baton Rouge, LA).

Table E-4. SEMIVOLATILE COMPOUNDS IDENTIFIED IN EXTRACT OF SAMPLE 4093 (Charleston, WV)

Chromaco- graphic Paak No.	Tiution Temp- ("C)	Compound	Chroneto- graphia Peak No.	Elution Temp. (°C)	Coupound
1		toluses	30		eilorene
2		silenma	31		eilozase
3		eilomane	32		d ₁₆ -pyrane (int. std.)
44.		silomana	33		eat. and uneat, bydrocarbons
42		sat. hydrocarbon) ¥		eilogene
5		dilozane	35		est. and usest. hydrocarbons
4		elloxene ·	36		est. and unest. hydrocarbons
7		butyric erhydride (tent.)	37A		est. and unset. hydrocathons
•		sac. bydrocarbon	37%		DOE
94		C _g R ₂₀ isomer	38		est. and unset. hydrocarbons
93		unknown	39		eiloxene
10		#1lozane	40		eilemme
11		sat. hydrocarbon	41		set. and unset. hydrocarbons
12		sat. hydrocarbon	42A		eilozane
13		silomene	423		methyl dehydrosbietsta (tent.)
14		silomene	43		eilezace
15		set. hydrocarbon	44		est. hydrocarbon
16		sat. bydrocarbon	45		ailozana '
17		sac. hydrocarbon	46		est. and usset. bydrocarbone
18		eat. hydrocarbon	47		ailonne
19		minora	48		phthelate
20		Silozene	49 .		Silonane
21		est. bydrocarbon	50		ecknown .
22		eat. and monat. hydrocarbon	51		silozane
23A		eiloxane	52		#ilexace
233		sat. and wase. hydrocarbons	53		eilexace
24		eilozene	54		lycoperment
25		est. and unest. hydrocarbose	55		bilozane
26		silemene	36		cholesteryl acetate
27		telarite.	57		est, and unset, bydrocerboos
26		set. and usent. hydrocarbons	36		dilogane
29		set. and unset. hydrocarbons	59		Q-sacopherol (vitamia)

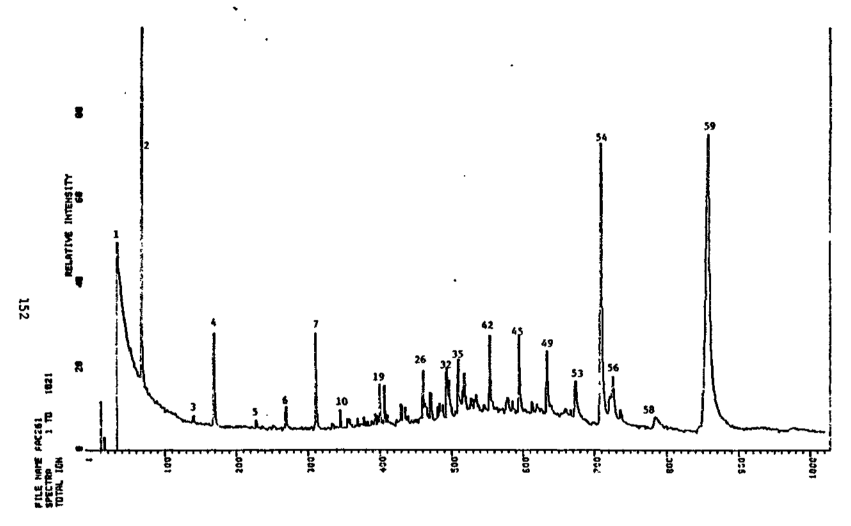


Figure E-4. Total ion current chromatogram from GC/MS analysis for semivolatiles in sample 4093 (Charleston, WV).



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