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## Final Report

# Hot Mix Asphalt Plants Truck Loading and Silo Filling Manual Methods Testing

## Asphalt Plant C Los Angeles, California

Volume 6 of 8



**FINAL REPORT**

**HOT MIX ASPHALT PLANTS  
TRUCK LOADING AND SILO FILLING  
MANUAL METHODS TESTING  
ASPHALT PLANT C, LOS ANGELES, CALIFORNIA**

**VOLUME 6 OF 8  
APPENDIX G.3 (CONCLUDED)**

**EPA Contract No. 68-D-98-004  
Work Assignment No. 3-02**

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**May 2000**

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## **DISCLAIMER**

The information in this document has been funded wholly or in part by the Office of Air Quality Planning and Standards, U.S. Environmental Protection Agency (EPA) under contract to Pacific Environmental Services, Inc. (PES). PES performed the work presented in this document under three EPA contracts and seven Work Assignments; EPA Contract No. 68-D-98-004, Work Assignment Nos. 1-08, 2-07, 3-02, and 3-05, EPA Contract No. 68-D-70002, Work Assignment Nos. 0-05 and 1-07, and EPA Contract No. 68-D-70069, Work Assignment No. 2-16. This document has been prepared by PES, reviewed following PES' internal quality assurance procedures, and approved by PES for distribution. This document has been subjected to the Agency's review, and has been approved by EPA for publication as an EPA document. Mention of trade names does not constitute endorsement by the EPA or PES.

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## GLOSSARY OF TERMS

ASTM – American Society for Testing and Materials  
CEMS – Continuous Emissions Monitoring System  
CTS – Calibration Transfer Standard  
EMC – Emissions Measurement Center  
EMAD – Emission Monitoring and Analysis Division  
ESP – Electrostatic Precipitator  
FID – Flame Ionization Detector  
FTIR – Fourier Transform Infrared Spectroscopy  
HAP – Hazardous Air Pollutant  
MCEM – Methylene Chloride Extractable Matter  
MRI – Midwest Research Institute  
PES – Pacific Environmental Services  
PM – Particulate Matter  
PTE – Permanent Total Enclosure  
RAP – Recycled Asphalt  
RTFOT – Rolling Thin Film Oven Test  
SED – Silo Exhaust Duct

## GLOSSARY OF TERMS (CONTINUED)

SMTG – Source Measurement Technology Group  
SVOHAP – Semi-Volatile Organic Hazardous Air Pollutant  
TED – Tunnel Emissions Duct  
TFOT – Thin Film Oven Test  
THC – Total Hydrocarbons  
VOHAP – Volatile Organic Hazardous Air Pollutant  
VOST – Volatile Organic Sampling Train



VOLUME 5B

APPENDIX G

ANALYTICAL DATA (CONTINUED)

G.3 SVOHAPS DATA (CONCLUDED)



Semivolatile Organics  
Method 0010/8270

Client Name: Pacific Environmental Services  
Client ID: S-MM5-RB-F, FH, XAD, COND, BH  
LAB ID: 300681-0009-SA  
Matrix: AIRTRAIN  
Authorized: 30 JUL 98

Sampled: 25 JUL 98  
Prepared: 31 JUL 98

Received: 30 JUL 98  
Analyzed: 31 AUG 98

Dilution Factor: 1.0

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
Phenol	ND	ug/Sample	30	R
bis(2-Chloroethyl)ether	ND	ug/Sample	30	
2-Chlorophenol	ND	ug/Sample	30	
1,3-Dichlorobenzene	ND	ug/Sample	30	
1,4-Dichlorobenzene	ND	ug/Sample	30	
Benzyl alcohol	ND	ug/Sample	30	
1,2-Dichlorobenzene	ND	ug/Sample	30	
2-Methylphenol	ND	ug/Sample	30	
2,2'-Oxybis(1-chloropropane)	ND	ug/Sample	30	
3/4-Methylphenol	ND	ug/Sample	30	
N-Nitroso-di-n-propylamine	ND	ug/Sample	30	
Hexachloroethane	ND	ug/Sample	30	
Nitrobenzene	ND	ug/Sample	30	
Isophorone	ND	ug/Sample	30	
2-Nitrophenol	ND	ug/Sample	30	
2,4-Dimethylphenol	ND	ug/Sample	30	
Benzoic acid	ND	ug/Sample	150	
bis(2-Chloroethoxy)-methane	ND	ug/Sample	30	
2,4-Dichlorophenol	ND	ug/Sample	30	
1,2,4-Trichlorobenzene	ND	ug/Sample	30	
Naphthalene	ND	ug/Sample	30	
4-Chloroaniline	ND	ug/Sample	30	
Hexachlorobutadiene	ND	ug/Sample	30	
4-Chloro-3-methylphenol	ND	ug/Sample	30	
2-Methylnaphthalene	ND	ug/Sample	30	
Hexachlorocyclopentadiene	ND	ug/Sample	30	
2,4,6-Trichlorophenol	ND	ug/Sample	30	
2,4,5-Trichlorophenol	ND	ug/Sample	150	
2-Chloronaphthalene	ND	ug/Sample	30	
2-Nitroaniline	ND	ug/Sample	30	
Dimethyl phthalate	ND	ug/Sample	30	
Acenaphthylene	ND	ug/Sample	30	
3-Nitroaniline	ND	ug/Sample	150	
Acenaphthene	ND	ug/Sample	30	
2,4-Dinitrophenol	ND	ug/Sample	150	
4-Nitrophenol	ND	ug/Sample	150	
Dibenzofuran	ND	ug/Sample	30	

Note R = Reporting limit(s) raised due to sample volume limitations.  
ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Karin Yee

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Semivolatile Organics  
Method 0010/8270

(cont.)

Client Name: Pacific Environmental Services  
Client ID: S-MM5-RB-F, FH, XAD, COND, BH  
LAB ID: 300681-0009-SA  
Matrix: AIRTRAIN  
Authorized: 30 JUL 98

Sampled: 25 JUL 98  
Prepared: 31 JUL 98

Received: 30 JUL 98  
Analyzed: 31 AUG 98

Dilution Factor: 1.0

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
2,4-Dinitrotoluene	ND	ug/Sample	30	
2,6-Dinitrotoluene	ND	ug/Sample	30	
Diethyl phthalate	ND	ug/Sample	30	
4-Chlorophenyl phenyl ether	ND	ug/Sample	30	
Fluorene	ND	ug/Sample	30	
4-Nitroaniline	ND	ug/Sample	150	
4,6-Dinitro-2-methylphenol	ND	ug/Sample	150	
N-Nitrosodiphenylamine	ND	ug/Sample	30	
4-Bromophenyl phenyl ether	ND	ug/Sample	30	
Hexachlorobenzene	ND	ug/Sample	30	
Pentachlorophenol	ND	ug/Sample	150	
Phenanthrene	ND	ug/Sample	30	
Anthracene	ND	ug/Sample	30	
Di-n-butyl phthalate	ND	ug/Sample	30	
Fluoranthene	ND	ug/Sample	30	
Pyrene	ND	ug/Sample	30	
Butyl benzyl phthalate	ND	ug/Sample	30	
3,3'-Dichlorobenzidine	ND	ug/Sample	60	
Benzo(a)anthracene	ND	ug/Sample	30	
bis(2-Ethylhexyl)-phthalate	ND	ug/Sample	30	
Chrysene	ND	ug/Sample	30	
Di-n-octyl phthalate	ND	ug/Sample	30	
Benzo(b)fluoranthene	ND	ug/Sample	30	
Benzo(k)fluoranthene	ND	ug/Sample	30	
Benzo(a)pyrene	ND	ug/Sample	30	
Indeno(1,2,3-cd)pyrene	ND	ug/Sample	30	
Dibenz(a,h)anthracene	ND	ug/Sample	30	
Benzo(g,h,i)perylene	ND	ug/Sample	30	
Acetophenone	ND	ug/Sample	30	
4-Aminobiphenyl	ND	ug/Sample	150	
Aniline	ND	ug/Sample	30	
Benzidine	ND	ug/Sample	300	
3,3'-Dimethylbenzidine	ND	ug/Sample	60	
N-Nitrosodimethylamine	ND	ug/Sample	30	
N-Nitrosomorpholine	ND	ug/Sample	30	
Pentachloronitrobenzene (PCNB)	ND	ug/Sample	150	
o-Toluidine	ND	ug/Sample	60	

ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Karin Yee

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Semivolatile Organics  
Method 0010/8270

(cont.)

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-RB-F, FH, XAD, COND, BH  
 LAB ID: 300681-0009-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98  
 Sampled: 25 JUL 98  
 Prepared: 31 JUL 98  
 Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 1.0

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
2-Methoxybenzenamine	ND	ug/Sample	--	K
Biphenyl	ND	ug/Sample	--	K
Chloroacetophenone	ND	ug/Sample	--	K
Cumene	ND	ug/Sample	--	K
DBCP (1,2-Dibromo-3-chloropropane)	ND	ug/Sample	--	K
Benzo(e)pyrene	ND	ug/Sample	--	K
N-N-Diethylaniline	ND	ug/Sample	--	K
Dimethylaniline	ND	ug/Sample	--	K
3,3'-Dimethoxybenzidine	ND	ug/Sample	--	K
Hydroquinone	ND	ug/Sample	--	K
4,4'-Methyl-bis(2-chloroaniline)	ND	ug/Sample	--	K
4-Nitrodiphenyl	ND	ug/Sample	--	K
Trifluralin	ND	ug/Sample	--	K

Surrogate	Recovery	Acceptable Range
Nitrobenzene-d5	61 %	45 - 107
2-Fluorobiphenyl	86 %	62 - 110
Terphenyl-d14	97 %	58 - 135
Phenol-d5	67 %	43 - 130
2-Fluorophenol	62 %	36 - 111
2,4,6-Tribromophenol	84 %	58 - 131

Note K = Identified by mass spectrum only; quantitation based on 1:1 response with internal standard.

ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Karin Yee

The cover letter is an integral part of this report.

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Semivolatiles Library Search (20 Compound TID)  
Method 8270

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-RB-F, FH, XAD, COND, BH  
 LAB ID: 300681-0009-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98  
 Sampled: 25 JUL 98  
 Prepared: NA  
 Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 1.0

Parameter	Result	Units	Reporting Limit	Qualifier
n-Nonane	400	ug/Sample	--	0
Unknown	51	ug/Sample	--	
Unknown	77	ug/Sample	--	
Unknown	71	ug/Sample	--	
Unknown	42	ug/Sample	--	
Unknown	63	ug/Sample	--	
Unknown	99	ug/Sample	--	
5-Eiconsene, (E) -	250	ug/Sample	--	0
Unknown	94	ug/Sample	--	
Unknown	63	ug/Sample	--	
Unknown alkane	27	ug/Sample	--	
Unknown	35	ug/Sample	--	
Unknown	140	ug/Sample	--	
4-Hydroxy-4-methyl-2-pentanone	270	ug/Sample	--	0
Unknown	110	ug/Sample	--	
Unknown	81	ug/Sample	--	
Unknown	26	ug/Sample	--	
9-Eicosene (E)-	57	ug/Sample	--	0
Unknown	61	ug/Sample	--	
Unknown	72	ug/Sample	--	

Note 0 = Or structurally similar compound (isomer).  
 NA = Not Applicable

Reported By: Emily Uebelhoer

Approved By: Karin Yee

The cover letter is an integral part of this report.  
 Rev 230787

1251

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068109  
Std Id: ST16980831

Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 22:58  
Analyst: DAT Instrument ID: F16 Run Factor: 3.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	30.60	61.2	45 107
CS25 2-FLUOROBIPHENY	50.00	43.00	86.0	62 110
CS30 TERPHENYL-D14	50.00	48.60	97.2	58 135
CS45 PHENOL-D5	100.0	66.90	66.9	43 130
CS50 2-FLUOROPHENOL	100.0	62.40	62.4	36 111
CS55 2,4,6-TRIBROMOP	100.0	84.30	84.3	58 131

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
HEXACHLOROBENZENE-C13		ND	UG/A	30.0
C310 N-NITROSODIMETHYLAMINE		ND	UG/A	30.0
PYRIDINE		ND	UG/A	60.0
2-PICOLINE		ND	UG/A	30.0
N-NITROSOMETHYLETHYLAMINE		ND	UG/A	30.0
METHYLMETHANESULFONATE		ND	UG/A	30.0
N-NITROSODIETHYLAMINE		ND	UG/A	30.0
ETHYLMETHANE SULFONATE		ND	UG/A	30.0
PENTACHLOROETHANE		ND	UG/A	30.0
C320 ANILINE		ND	UG/A	30.0
C315 PHENOL		ND	UG/A	30.0
C325 BIS(2-CHLOROETHYL)ETHER		ND	UG/A	30.0
C330 2-CHLOROPHENOL		ND	UG/A	30.0
C335 1,3-DICHLOROBENZENE		ND	UG/A	30.0
C340 1,4-DICHLOROBENZENE		ND	UG/A	30.0
C345 BENZYL ALCOHOL		ND	UG/A	30.0
C350 1,2-DICHLOROBENZENE		ND	UG/A	30.0
C355 2-METHYLPHENOL		ND	UG/A	30.0
C360 2,2'-OXYBIS(1-CLPROPAN)		ND	UG/A	30.0
C361 ACETOPHENONE		ND	UG/A	30.0
N-NITROSPYRROLIDINE		ND	UG/A	30.0
N-NITROSOMORPHOLINE		ND	UG/A	30.0
3-METHYL PHENOL		ND	UG/A	30.0
C365 4-METHYLPHENOL		ND	UG/A	30.0
C370 N-NITROSO-DI-N-PROPYLAM		ND	UG/A	30.0
O-TOLUIDINE		ND	UG/A	30.0
C375 HEXACHLOROETHANE		ND	UG/A	30.0
C410 NITROBENZENE		ND	UG/A	30.0
N-NITROSOPIPERIDINE		ND	UG/A	30.0

Reviewed by: 149/17/ab

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QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068109  
Std Id: ST16980831

Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 22:58  
Analyst: DAT Instrument ID: F16 Run Factor: 3.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C415 ISOPHORONE		ND	UG/A	30.0
C420 2-NITROPHENOL		ND	UG/A	30.0
C425 2,4-DIMETHYLPHENOL		ND	UG/A	30.0
C435 BIS(2-CHLOROETHOXY)METH		ND	UG/A	30.0
C440 2,4-DICHLOROPHENOL		ND	UG/A	30.0
C430 BENZOIC ACID		ND	UG/A	150.0
AA-DIMETHYLPHENETHYLAMINE		ND	UG/A	30.0
C445 1,2,4-TRICHLOROBENZENE		ND	UG/A	30.0
C450 NAPHTHALENE		ND	UG/A	30.0
C455 4-CHLOROANALINE		ND	UG/A	30.0
2,6-DICHLOROPHENOL		ND	UG/A	30.0
HEXACHLOROPROPENE		ND	UG/A	30.0
C460 HEXACHLOROBUTADIENE		ND	UG/A	30.0
P-PHENYLENE DIAMINE		ND	UG/A	30.0
N-NITROSODI-N-BUTYLAMINE		ND	UG/A	30.0
C465 4-CHLORO-3-METHYLPHENO		ND	UG/A	30.0
SAFROLE		ND	UG/A	30.0
C470 2-METHYLNAPHTHALENE		ND	UG/A	30.0
1,2,4,5-TETRACHLOROBENZENE		ND	UG/A	30.0
ISOSAFROLE (#1)		ND	UG/A	60.0
C510 HEXACHLOROCYCLOPENTADI		ND	UG/A	30.0
C515 2,4,6-TRICHLOROPHENOL		ND	UG/A	30.0
C520 2,4,5-TRICHLOROPHENOL		ND	UG/A	30.0
ISOSAFROLE (#2)		ND	UG/A	60.0
C525 2-CHLORONAPHTHALENE		ND	UG/A	30.0
1-CHLORONAPHTHALENE		ND	UG/A	15.0
C530 2-NITROANALINE		ND	UG/A	150.0
1,4-NAPHTHOQUINONE		ND	UG/A	30.0
C535 DIMETHYLPHTHALATE		ND	UG/A	30.0
1,3-DINITROBENZENE		ND	UG/A	30.0
C540 ACENAPHTHYLENE		ND	UG/A	30.0
C543 2,6-DINITROTOLUENE		ND	UG/A	30.0
C545 3-NITROANILINE		ND	UG/A	150.0
C550 ACENAPHTHENE		ND	UG/A	30.0
C555 2,4-DINITROPHENOL		ND	UG/A	150.0
C565 DIBENZOFURAN		ND	UG/A	30.0
C560 4-NITROPHENOL		ND	UG/A	150.0
PENTACHLOROBENZENE		ND	UG/A	30.0
C570 2,4-DINITROTOLUENE		ND	UG/A	30.0
1-NAPHTHYLAMINE		ND	UG/A	30.0
2-NAPHTHYLAMINE		ND	UG/A	30.0
2,3,4,6-TETRACHLOROPHENOL		ND	UG/A	60.0
C580 DIETHYLPHTHALATE		ND	UG/A	30.0
C590 FLUORENE		ND	UG/A	30.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068109  
Std Id: ST16980831

Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 22:58  
Analyst: DAT Instrument ID: F16 Run Factor: 3.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C585 4-CHLOROPHENYL-PHENYLE		ND	UG/A	30.0
5-NITRO-O-TOLUIDINE		ND	UG/A	30.0
C595 4-NITROANILINE		ND	UG/A	150.0
C610 4,6-DINITRO-2-METHYLPH		ND	UG/A	150.0
C615 N-NITROSODIPHENYLAMINE		ND	UG/A	30.0
C620 AZOBENZENE		ND	UG/A	30.0
SYM-TRINITROBENZENE		ND	UG/A	30.0
C625 4-BROMOPHENYL-PHENYLET		ND	UG/A	30.0
PHENACETIN		ND	UG/A	30.0
DIALATE	234	ND	UG/A	30.0
C630 HEXACHLOROBENZENE		ND	UG/A	30.0
4-AMINOBIIPHENYL		ND	UG/A	30.0
C635 PENTACHLOROPHENOL		ND	UG/A	150.0
PRONAMIDE		ND	UG/A	30.0
PENTACHLORONITROBENZENE		ND	UG/A	150.0
C640 PHENANTHRENE		ND	UG/A	30.0
C645 ANTHRACENE		ND	UG/A	30.0
2SECBUTYL-4,6-DINITROPHENOL		ND	UG/A	30.0
C647 CARBAZOLE		ND	UG/A	30.0
C650 DI-N-BUTYLPHTHALATE		ND	UG/A	30.0
4-NITROQUINOLINE-1-OXIDE		ND	UG/A	30.0
METHAPYRILENE		ND	UG/A	30.0
ISODRIN		ND	UG/A	30.0
C655 FLUORANTHENE		ND	UG/A	30.0
CHLOROBENZILATE		ND	UG/A	30.0
C710 BENZIDINE		ND	UG/A	300.0
C715 PYRENE		ND	UG/A	30.0
ARAMITE (#1)		ND	UG/A	30.0
ARAMITE (#2)		ND	UG/A	30.0
P-DIMETHYLAMINOAZOBENZENE		ND	UG/A	30.0
3,3'-DIMETHYLBENZIDINE		ND	UG/A	30.0
KEPONE		ND	UG/A	150.0
C720 BUTYLBENZYLPHTHALATE		ND	UG/A	30.0
2-ACETYLAMINOFLUORENE		ND	UG/A	30.0
C730 BENZO(A)ANTHRACENE		ND	UG/A	30.0
C725 3,3'-DICHLOROBENZIDINE		ND	UG/A	60.0
C740 CHRYSENE		ND	UG/A	30.0
C745 BIS(2-ETHYLHEXYL)PHTHA		ND	UG/A	30.0
3-METHYLCHOLANTHRENE		ND	UG/A	30.0
C760 DI-N-OCTYL PHTHALATE		ND	UG/A	30.0
C765 BENZO(B)FLUORANTHENE		ND	UG/A	30.0
7,12-DIMETHYLBENZANTHRACENE		ND	UG/A	30.0
C770 BENZO(K)FLUORANTHENE		ND	UG/A	30.0
HEXACHLOROPHENE		ND	UG/A	30.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

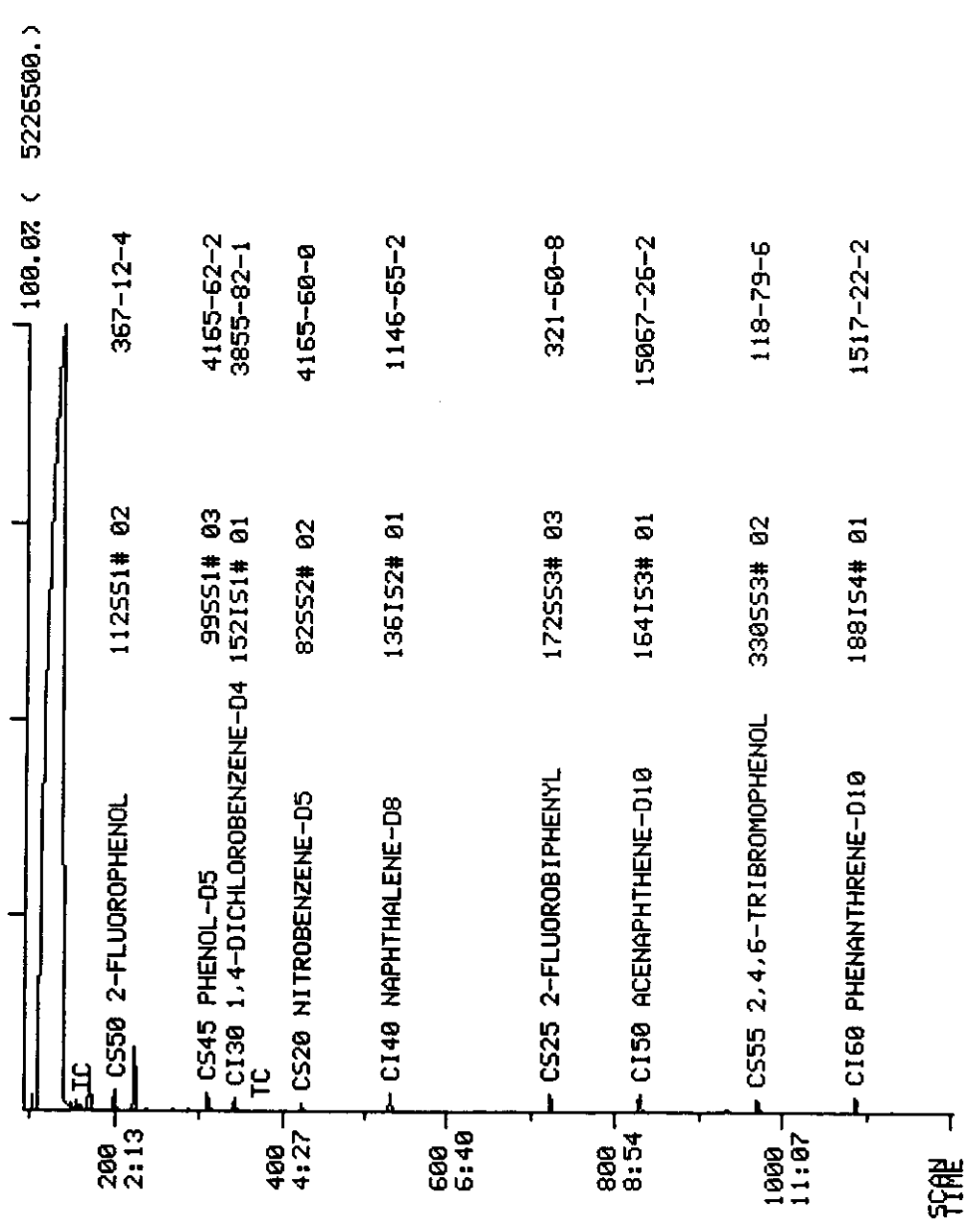
Data File: 30068109  
Std Id: ST16980831

Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 22:58  
Analyst: DAT Instrument ID: F16 Run Factor: 3.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Target Compounds: SAP9

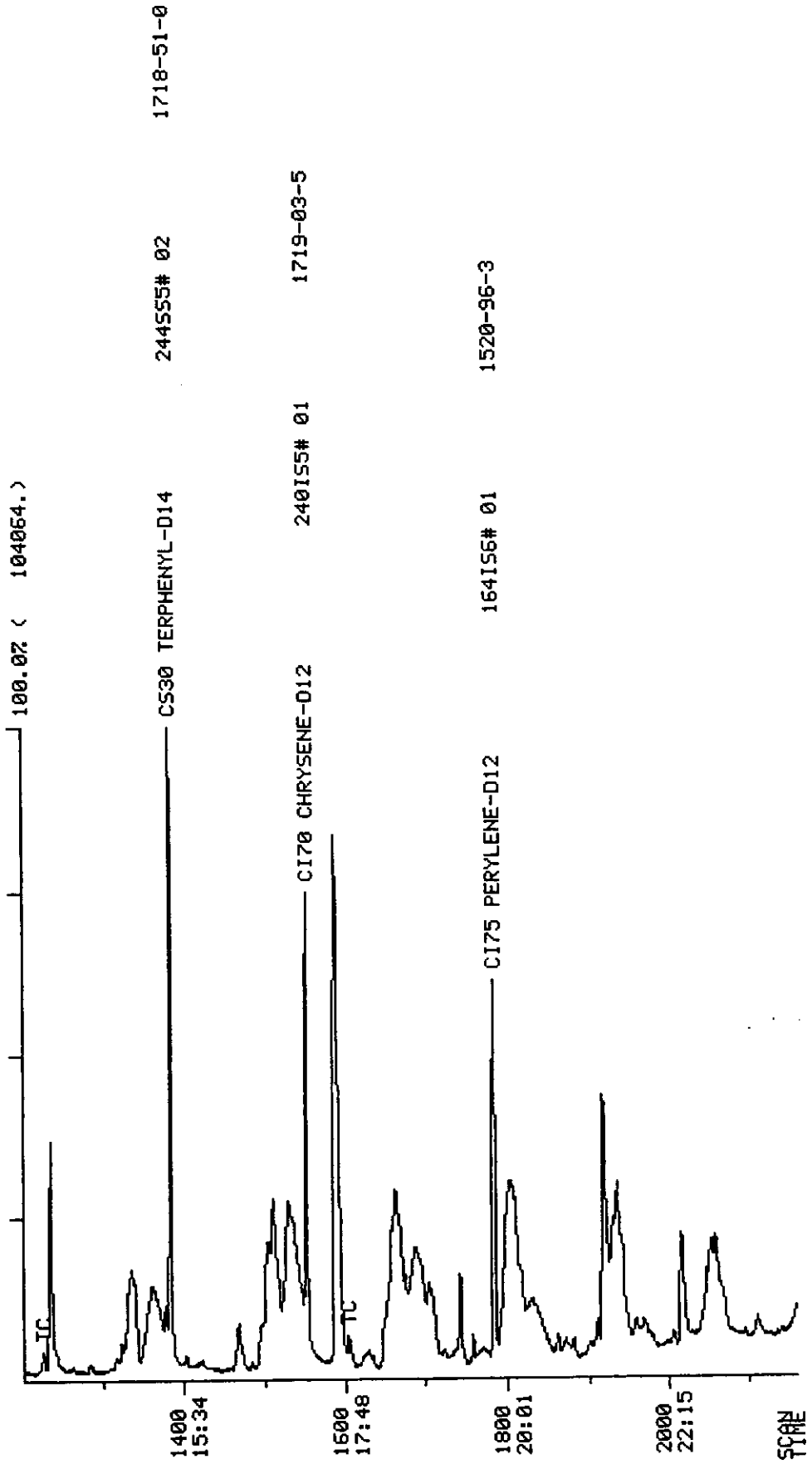
Parameter	Scan	Result	Units	Reporting Limit
C775 BENZO(A)PYRENE		ND	UG/A	30.0
C780 INDENO(1,2,3-CD)PYRENE		ND	UG/A	30.0
C785 DIBENZ(A,H)ANTHRACENE		ND	UG/A	30.0
C790 BENZO(G,H,I)PERYLENE		ND	UG/A	30.0

DATA FROM FILE: 30068109    SCANS    93 TO 1203    ACQUIRED: 08/31/98 22:58:00  
 SAMPLE: 5-MMS-R8    1/35A/1ML    CALI: 30068109 #3  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M    INST. ID: F16





DATA FROM FILE: 30068109      SCANS 1203 TO 2158    ACQUIRED: 08/31/98 22:58:00  
 CALI: 30068109 #3  
 SAMPLE: S-MMS-RB    1/35A/1ML    INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA >/1/35A NA M



ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 30068109

Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 22:58

Compounds with amounts less than 0.20 reported as NOT FOUND

Library							Units: UG/ML		
No	Entry	Name	Mass	Meth	Scan	Ref	Area	RRF(L)	Amount
1	S1#	1 C130 1,4-DICHLOROBENZENE	152	A	BB	340 1	36331.	1.000	40.000
2	S2#	1 C140 NAPHTHALENE-D8	136	A	BB	529 2	137829.	1.000	40.000
3	S3#	1 C150 ACENAPHTHENE-D10	164	A	BB	827 3	74485.	1.000	40.000
4	S4#	1 C160 PHENANTHRENE-D10	188	A	BB	1085 4	122676.	1.000	40.000
5	S5#	1 C170 CHRYSENE-D12	240	A	BB	1551 5	71172.	1.000	40.000
6	S6#	1 C175 PERYLENE-D12	264	A	BB	1782 6	56685.	1.000	40.000
7	S2#	2 CS20 NITROBENZENE-D5	82	A	BB	421 2	49520.	0.470	30.606
8	S3#	3 CS25 2-FLUOROBIPHENYL	172	A	BB	722 3	98741.	1.234	42.961
9	S5#	2 CS30 TERPHENYL-D14	244	A	BB	1383 5	86030.	0.995	48.582
10	S1#	3 CS45 PHENOL-D5	99	A	BB	308 1	123023.	2.025	66.871
11	S1#	2 CS50 2-FLUOROPHENOL	112	A	BB	198 1	77514.	1.368	62.374
12	S3#	2 CS55 2,4,6-TRIBROMOPHENO	330	A	BB	968 3	36995.	0.236	84.305
13	S4#	4 HEXACHLOROBENZENE-C13	294			4 NOT FOUND			
14	S1#	15 C310 N-NITROSODIMETHYLAM	74			1 NOT FOUND			
15	S1#	20 PYRIDINE	79			1 NOT FOUND			
16	S1#	30 2-PICOLINE	93	A	BB	146 1	449.	1.565	0.316
17	S1#	40 N-NITROSOMETHYLETHYLAMIN	42	A	VB	159 1	4136.	0.811	5.614
18	S1#	60 METHYLMETHANESULFONATE	80			1 NOT FOUND			
19	S1#	70 N-NITROSODIETHYLAMINE	102			1 NOT FOUND			
20	S1#	85 ETHYLMETHANE SULFONATE	79			1 NOT FOUND			
21	S1#	95 PENTACHLOROETHANE	117			1 NOT FOUND			
22	S1#	100 C320 ANILINE	93			1 NOT FOUND			
23	S1#	105 C315 PHENOL	94	A	BB	309 1	486.	1.968	0.272
24	S1#	110 C325 BIS(2-CHLOROETHYL)E	93			1 NOT FOUND			
25	S1#	115 C330 2-CHLOROPHENOL	128			1 NOT FOUND			
26	S1#	125 C335 1,3-DICHLOROBENZENE	146			1 NOT FOUND			
27	S1#	130 C340 1,4-DICHLOROBENZENE	146			1 NOT FOUND			
28	S1#	145 C345 BENZYL ALCOHOL	108	A	BB	372 1	2329.	0.854	3.001
29	S1#	150 C350 1,2-DICHLOROBENZENE	146			1 NOT FOUND			
30	S1#	160 C355 2-METHYLPHENOL	108			1 NOT FOUND			
31	S1#	165 C360 2,2'-OXYBIS(1-CLPRO	45			1 NOT FOUND			
32	S1#	170 C361 ACETOPHENONE	105			1 NOT FOUND			
33	S1#	175 N-NITROSOPYRROLIDINE	100			1 NOT FOUND			
34	S1#	180 N-NITROSOMORPHOLINE	56			1 NOT FOUND			
35	S1#	182 3-METHYL PHENOL	108	A	BB	421 1	568.	2.252	0.278
36	S1#	185 C365 4-METHYLPHENOL	108			1 NOT FOUND			
37	S1#	190 C370 N-NITROSO-DI-N-PROP	70			1 NOT FOUND			
38	S1#	195 O-TOLUIDINE	106			1 NOT FOUND			
39	S1#	200 C375 HEXACHLOROETHANE	117			1 NOT FOUND			
40	S2#	10 C410 NITROBENZENE	77			2 NOT FOUND			
41	S2#	15 N-NITROSOPIPERIDINE	42			2 NOT FOUND			
42	S2#	20 C415 ISOPHORONE	82			2 NOT FOUND			
43	S2#	25 C420 2-NITROPHENOL	139			2 NOT FOUND			
44	S2#	30 C425 2,4-DIMETHYLPHENOL	107			2 NOT FOUND			
45	S2#	35 C435 BIS(2-CHLOROETHOXY)	93			2 NOT FOUND			

46 S2# 40 C440 2,4-DICHLOROPHENOL	162	2	NOT FOUND
47 S2# 45 C430 BENZOIC ACID	122	2	NOT FOUND
48 S2# 50 AA-DIMETHYLPHENETHYLAMIN	58	2	NOT FOUND
49 S2# 55 C445 1,2,4-TRICHLOROBENZ	180	2	NOT FOUND

50	S2# 60	C450	NAPHTHALENE	128	2	NOT FOUND
51	S2# 80	C455	4-CHLOROANALINE	127	2	NOT FOUND
52	S2# 85		2,6-DICHLOROPHENOL	162	2	NOT FOUND
53	S2# 90		HEXACHLOROPROPENE	213	2	NOT FOUND
54	S2# 95	C460	HEXACHLOROBUTADIENE	225	2	NOT FOUND
55	S2#115		P-PHENYLENE DIAMINE	108	2	NOT FOUND
56	S2#120		N-NITROSODI-N-BUTYLAMINE	84	2	NOT FOUND
57	S2#130	C465	4-CHLORO-3-METHYLPH	107	2	NOT FOUND
58	S2#140		SAFROLE	162	2	NOT FOUND
59	S2#145	C470	2-METHYLNAPHTHALENE	142	2	NOT FOUND
60	S3# 10		1,2,4,5-TETRACHLOROBEZNE	214	3	NOT FOUND
61	S3# 15		ISOSAFROLE (#1)	162	3	NOT FOUND
62	S3# 20	C510	HEXACHLOROCYCLOPENT	237	3	NOT FOUND
63	S3# 25	C515	2,4,6-TRICHLOROPHEN	196	3	NOT FOUND
64	S3# 30	C520	2,4,5-TRICHLOROPHEN	196	3	NOT FOUND
65	S3# 35		ISOSAFROLE (#2)	104	3	NOT FOUND
66	S3# 40	C525	2-CHLORONAPHTHALENE	162	3	NOT FOUND
67	S3# 42		1-CHLORONAPHTHALEN	162	3	NOT FOUND
68	S3# 45	C530	2-NITROANALINE	65	3	NOT FOUND
69	S3# 50		1,4-NAPHTHOQUINONE	158	3	NOT FOUND
70	S3# 55	C535	DIMETHYLPHTHALATE	163	3	NOT FOUND
71	S3# 60		1,3-DINITROBENZENE	168	3	NOT FOUND
72	S3# 65	C540	ACENAPHTHYLENE	152	3	NOT FOUND
73	S3# 70	C543	2,6-DINITROTOLUENE	165	3	NOT FOUND
74	S3# 75	C545	3-NITROANILINE	138	3	NOT FOUND
75	S3# 80	C550	ACENAPHTHENE	153	3	NOT FOUND
76	S3# 85	C555	2,4-DINITROPHENOL	184	3	NOT FOUND
77	S3# 90	C565	DIBENZOFURAN	168	3	NOT FOUND
78	S3# 95	C560	4-NITROPHENOL	109	3	NOT FOUND
79	S3#100		PENTACHLOROBEZNE	250	3	NOT FOUND
80	S3#105	C570	2,4-DINITROTOLUENE	165	3	NOT FOUND
81	S3#110		1-NAPHTHYLAMINE	143	3	NOT FOUND
82	S3#115		2-NAPHTHYLAMINE	143	3	NOT FOUND
83	S3#120		2,3,4,6-TETRACHLOROPHENO	232	3	NOT FOUND
84	S3#130	C580	DIETHYLPHTHALATE	149	3	NOT FOUND
85	S3#135	C590	FLUORENE	166	3	NOT FOUND
86	S3#140	C585	4-CHLOROPHENYL-PHEN	204	3	NOT FOUND
87	S3#145		5-NITRO-O-TOLUIDINE	152	3	NOT FOUND
88	S3#150	C595	4-NITROANALINE	138	3	NOT FOUND
89	S4# 10	C610	4,6-DINITRO-2-METHY	198	4	NOT FOUND
90	S4# 15	C615	N-NITROSODIPHENYLAM	169	4	NOT FOUND
91	S4# 20	C620	AZOBENZENE	77	4	NOT FOUND
92	S4# 25		SYM-TRINITROBENZENE	75	4	NOT FOUND
93	S4# 30	C625	4-BROMOPHENYL-PHENY	248	4	NOT FOUND
94	S4# 35		PHENACETIN	108	4	NOT FOUND
95	S4# 37		DIALLATE	234	4	NOT FOUND
96	S4# 40	C630	HEXACHLOROBEZNE	284	4	NOT FOUND
97	S4# 45		4-AMINOBIIPHENYL	169	4	NOT FOUND
98	S4# 50	C635	PENTACHLOROPHENOL	266	4	NOT FOUND
99	S4# 55		PRONAMIDE	173	4	NOT FOUND
100	S4# 60		PENTACHLORONITROBENZENE	237	4	NOT FOUND
101	S4# 65	C640	PHENANTHRENE	178	4	NOT FOUND
102	S4# 70	C645	ANTHRACENE	178	4	NOT FOUND
103	S4# 75		2SECBUTYL-4,6-DINITROPHE	211	4	NOT FOUND
104	S4# 80	C647	CARBAZOLE	167	4	NOT FOUND
105	S4# 85	C650	DI-N-BUTYLPHTHALATE	149	A BB 1226 4 4688. 1.247 1.226	
106	S4#100		4-NITROQUINOLINE-1-OXIDE	190	4	NOT FOUND
107	S4#105		METHAPYRILENE	58	4	NOT FOUND
108	S4#106		ISODRIN	193	4	NOT FOUND
109	S4#110	C655	FLUORANTHENE	202	4	NOT FOUND
110	S4#120		CHLOROBEZNYLATE	139	4	NOT FOUND

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111	S5#	10	C710	BENZIDINE	184	5	NOT	FOUND				
112	S5#	15	C715	PYRENE	202	5	NOT	FOUND				
113	S5#	20	ARAMITE	(#1)	185	5	NOT	FOUND				
114	S5#	25	ARAMITE	(#2)	185	5	NOT	FOUND				
115	S5#	30	P-DIMETHYLAMINOAZOBENZEN		120	5	NOT	FOUND				
116	S5#	35	3,3'-DIMETHYLBENZIDINE		212	5	NOT	FOUND				
117	S5#	37	KEPONE		272	5	NOT	FOUND				
118	S5#	40	C720	BUTYLBENZYLPHTHALAT	149	5	NOT	FOUND				
119	S5#	45	2-ACETYLAMINOFLUORENE		181	5	NOT	FOUND				
120	S5#	50	C730	BENZO(A)ANTHRACENE	228	5	NOT	FOUND				
121	S5#	55	C725	3,3'-DICHLOROBENZID	252	5	NOT	FOUND				
122	S5#	60	C740	CHRYSENE	228	5	NOT	FOUND				
123	S5#	65	C745	BIS(2-ETHYLHEXYL)PH	149	A	BB	1603	5	3310.	0.980	1.899
124	S5#	85	3-METHYLCHOLANTHRENE		268	5	NOT	FOUND				
125	S6#	10	C760	DI-N-OCTYL PHTHALAT	149	6	NOT	FOUND				
126	S6#	15	C765	BENZO(B)FLUORANTHEN	252	6	NOT	FOUND				
127	S6#	20	7,12-DIMETHYLBENZANTHRAC		256	6	NOT	FOUND				
128	S6#	25	C770	BENZO(K)FLUORANTHEN	252	6	NOT	FOUND				
129	S6#	30	HEXACHLOROPHENE		196	6	NOT	FOUND				
130	S6#	35	C775	BENZO(A)PYRENE	252	6	NOT	FOUND				
131	S6#	55	C780	INDENO(1,2,3-CD)PYR	276	6	NOT	FOUND				
132	S6#	60	C785	DIBENZ(A,H)ANTHRACE	278	6	NOT	FOUND				
133	S6#	65	C790	BENZO(G,H,I)PERYLEN	276	6	NOT	FOUND				

## QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 30068109	36331.= 126		3.78	137829.= 137		5.88	74485.= 137		9.20

IS# 1 = C130 1,4-DICHLOROBENZENE-D4

IS# 2 = C140 NAPHTHALENE-D8

IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 30068109	122676.= 124		12.07	71172.= 114		17.25	56685.= 111		19.82

IS# 4 = C160 PHENANTHRENE-D10

IS# 5 = C170 CHRYSENE-D12

IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

Data Reduced by: 2/1/98 Date: 9/11/96  
Data Reviewed by: 2/1/98 Date: 9/11/96

Data File: 30068109

QUANTERRA GC/MS TIC REPORT ( Part 1 )

Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
Analyst: DAT Date Analyzed: 08/31/98 22:58  
Run Factor: 3.00

Concentration

in Sample

*UOA 1/27*  
# SCAN (UG/A) CAS #  
169 270. 123-42-2  
2-PENTANONE, 4-HYDROXY-4-METHYL-

2 222 400. 111-84-2  
NONANE *or isomer*

~~3 285 25. 100-52-7  
BENZALDEHYDE~~



~~4 932 17. 74381-40-1  
PROPANOIC ACID, 2-METHYL-, 1-(1,1-DIMETHYLETHYL)-2-METHYL-1,3-PROPANEI~~

*unknown 5700*

5 1236 51. ~~57-10-3~~  
~~HEXADECANOIC ACID~~

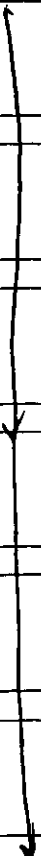
6 1333 77. 00-00-0  
UNKNOWN

7 1358 71. 00-00-0  
UNKNOWN

~~8 1466 23. 2432-89-5  
DECANOIC ACID, DIDECYL ESTER~~

9 1502 42. 00-00-0  
UNKNOWN

10 1509 63. 00-00-0  
UNKNOWN



11 1529 99. 00-00-0  
UNKNOWN 5700

12 1587 250. 74685-30-6  
5-EICOSENE, (E)- or *isomer*

13 1661 94. 00-00-0  
UNKNOWN 5700

14 1687 63. 00-00-0  
UNKNOWN

15 1706 27. 295-17-0  
CYCLOTETRADECANE *alkane 5746400*

16 1741 35. 00-00-0  
UNKNOWN 5700

17 1801 140. 00-00-0  
UNKNOWN

18 1834 14. 00-00-0  
UNKNOWN

19 1916 110. 1653-33-4  
~~1-TETRADECANOL~~

20 1934 81. 00-00-0  
UNKNOWN

*Use*  
21 1940 26. ✓ 00-00-0  
UNKNOWN

~~22~~ 1966 21. 00-00-0  
UNKNOWN



23 2014 57. 74685-29-3  
9-EICOSENE, (E)- *or Rosmar*

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---

24 2051 61. 00-00-0  
UNKNOWN

---

---

5700

25 2057 72. 00-00-0  
UNKNOWN

---

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## QUANTERRA GC/MS TIC REPORT ( Part 2 )

CONCENTRATION = AREA(TIC)\*CONC(1S)/AREA(1S)

#	FIT	PURITY	INT.			AMOUNT					
			STD.	RT	RRT	AREA	HEIGHT	(UG/ML )	LIB	LIB #	
1	957	921	1	1:52	0.319	648508.	251136.	90.815	NB	3241.	
2	986	940	1	2:28	0.420	946332.	411648.	132.522	NB	5159.	
3	966	921	1	3:10	0.539	59934.	18879.	8.393	NB	2017.	
4	918	803	2	10:21	1.127	49207.	16122.	5.755	NB	40501.	
5	976	775	3	13:44	1.139	139728.	35968.	16.915	NB	35182.	
6	891	556	4	14:49	0.859	135216.	12976.	25.735	UK	1.	
7	763	578	4	15:06	0.876	124360.	9168.	23.669	UK	1.	
8	957	689	4	16:18	0.945	39434.	6486.	7.505	NB	58380.	
9	760	383	4	16:42	0.968	73824.	12016.	14.050	UK	1.	
10	796	590	4	16:47	0.973	110512.	17856.	21.033	UK	1.	
11	931	516	4	17:00	0.986	174176.	15312.	33.150	UK	1.	
12	986	810	4	17:39	1.023	437632.	79488.	83.292	NB	39516.	
13	925	476	4	18:28	1.071	164160.	15600.	31.244	UK	1.	
14	800	573	5	18:45	0.947	97552.	7632.	20.916	UK	1.	
15	871	684	5	18:58	0.957	41976.	5648.	9.000	NB	21960.	
16	866	561	5	19:21	0.977	55048.	13486.	11.803	UK	1.	
17	876	591	5	20:01	1.011	221408.	17472.	47.472	UK	1.	
18	924	556	5	20:23	1.029	21952.	2816.	4.707	UK	1.	
19	980	781	5	21:18	1.075	175464.	34566.	37.621	NB	26416.	
20	775	478	5	21:30	1.085	125664.	16368.	26.943	UK	1.	
21	695	441	5	21:34	1.089	41056.	9312.	8.803	UK	1.	
22	789	391	5	21:52	1.103	32442.	3297.	6.956	UK	1.	
23	949	729	5	22:24	1.130	89076.	16549.	19.099	NB	39515.	
24	922	471	5	22:48	1.151	94560.	12016.	20.274	UK	1.	
25	749	477	5	22:52	1.154	111664.	13104.	23.942	UK	1.	

Library Search Data: 30068109 # 169 Base m/z: 43  
 08/31/98 22:58:00 + 1:53 Cali: 30068109 # 3 RIC: 229120.  
 Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

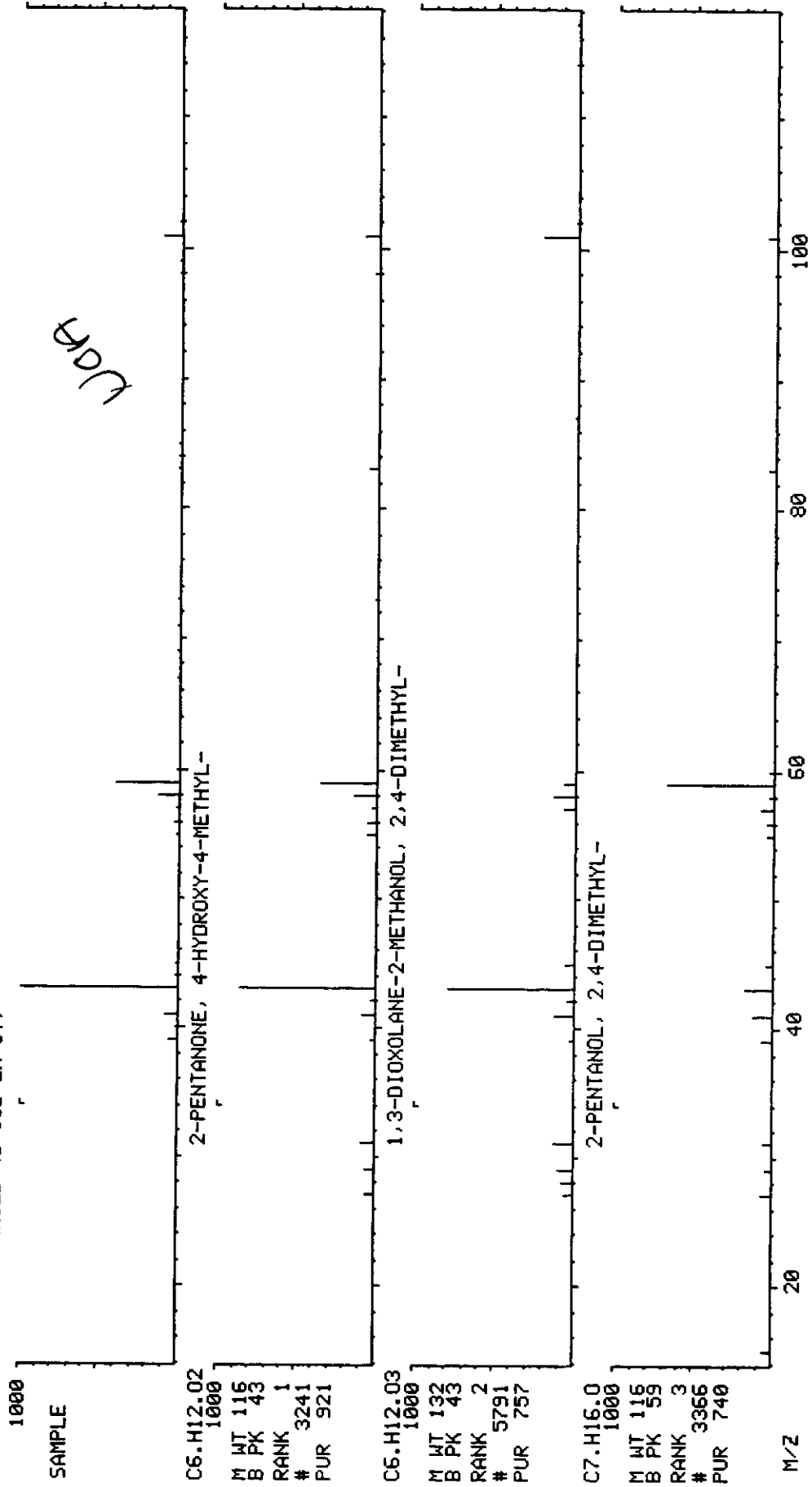
62231 spectra in LIBRARYNB searched for maximum PURITY  
 630 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	3241 2-PENTANONE, 4-HYDROXY-4-METHYL-
2	5791 1,3-DIOXOLANE-2-METHANOL, 2,4-DIMETHYL-
3	3366 2-PENTANOL, 2,4-DIMETHYL-
4	3288 ACETIC ACID, 1,1-DIMETHYLETHYL ESTER
5	919 TERT-BUTYL HYDROPEROXIDE
6	5594 2-PROPANOL, 2-NITROSO-, ACETATE (ESTER)
7	3349 2-HEXANOL, 2-METHYL-
8	8432 2-HEXANONE, 3-HYDROXY-3,5-DIMETHYL-
9	3234 OXIRANE, [(1-METHYLETHOXY)METHYL]-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H12.O2	116	43	921	957	934
2	C6.H12.O3	132	43	757	825	800
3	C7.H16.O	116	59	740	760	740
4	C6.H12.O2	116	43	722	769	735
5	C4.H10.O2	90	59	719	861	741
6	C5.H9.O3.N	131	43	706	818	796
7	C7.H16.O	116	59	701	732	701
8	C8.H16.O2	144	59	666	722	693
9	C6.H12.O2	116	43	658	703	676

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	123-42-2
2	---	---	---	---	53951-43-2
3	---	---	---	---	625-06-9
4	---	---	---	---	540-88-5
5	---	---	---	---	75-91-2
6	---	---	---	---	6931-04-0
7	---	---	---	---	625-23-0
8	---	---	---	---	6321-14-8
9	---	---	---	---	4016-14-2

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 22:58:00 + 1:53  
 SAMPLE: S-MM5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 # 169 BASE M/Z: 43  
 CALI: 30068109 # 3 RIC: 229120.



Library Search                    Data: 30068109 # 222            Base m/z: 43  
 08/31/98 22:58:00 + 2:28        Cali: 30068109 # 3            RIC: 312320.  
 Sample: S-MMS-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

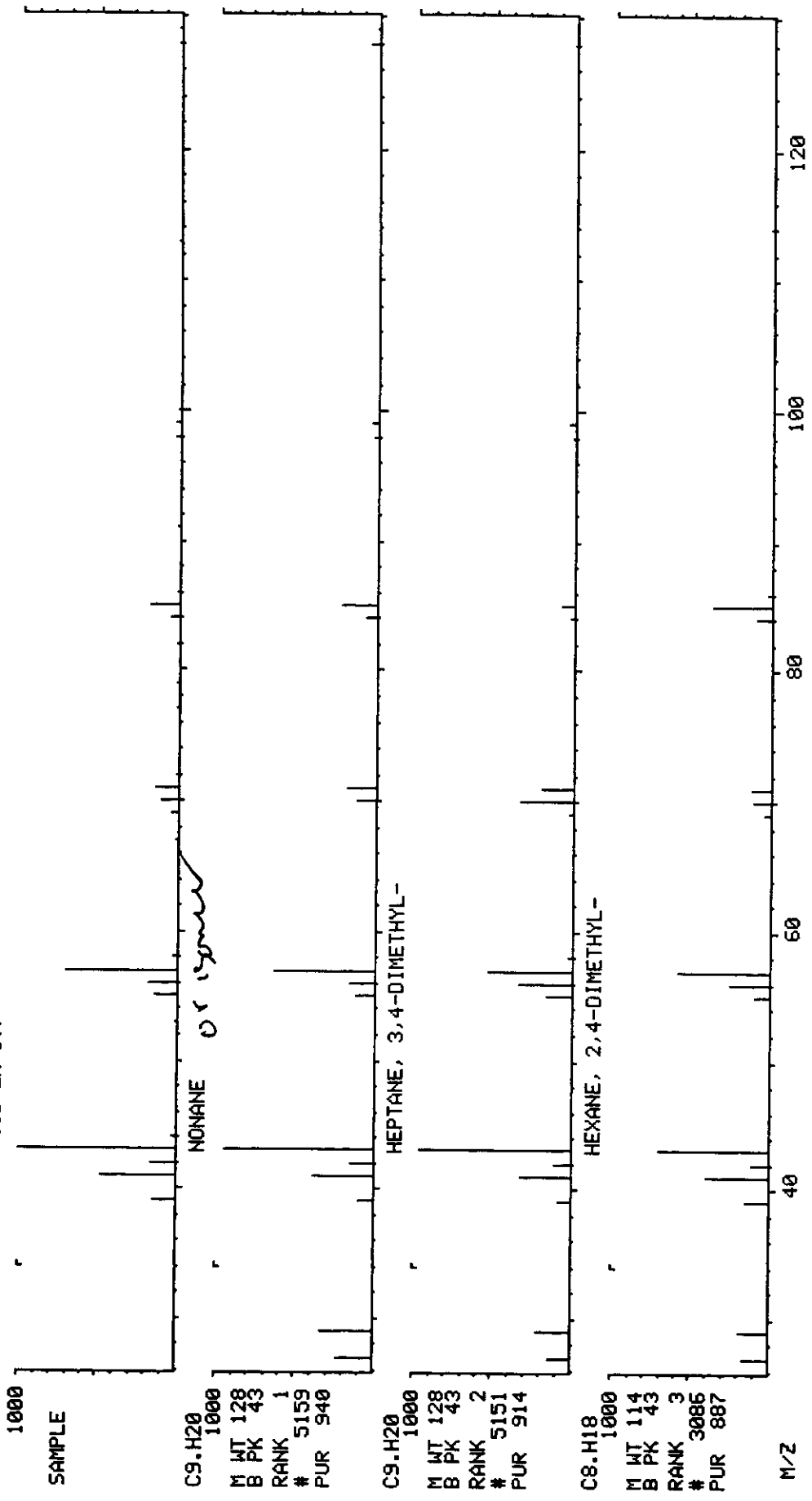
62231 spectra in LIBRARYNB searched for maximum PURITY  
 921 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	5159 NONANE
2	5151 HEPTANE, 3,4-DIMETHYL-
3	3086 HEXANE, 2,4-DIMETHYL-
4	19015 DECANE, 2,5,6-TRIMETHYL-
5	5141 HEPTANE, 2,4-DIMETHYL-
6	5154 HEXANE, 4-ETHYL-2-METHYL-
7	11607 UNDECANE
8	5160 PENTANE, 2,2,3,4-TETRAMETHYL-
9	5144 HEPTANE, 4-ETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C9.H20	128	43	940	986	940
2	C9.H20	128	43	914	939	914
3	C8.H18	114	43	887	957	898
4	C13.H28	184	57	885	921	895
5	C9.H20	128	43	883	932	884
6	C9.H20	128	57	875	923	875
7	C11.H24	156	43	869	925	913
8	C9.H20	128	57	866	924	866
9	C9.H20	128	43	865	909	872

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	111-84-2
2	---	---	---	---	922-28-1
3	---	---	---	---	589-43-5
4	---	---	---	---	62108-23-0
5	---	---	---	---	2213-23-2
6	---	---	---	---	3074-75-7
7	---	---	---	---	1120-21-4
8	---	---	---	---	1186-53-4
9	---	---	---	---	2216-32-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 2:28  
 SAMPLE: S-MM5-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 # 222  
 CALI: 30068109 # 3  
 BASE M/Z: 43  
 RIC: 312320.



1270

Library Search Data: 30068109 # 285 Base m/z: 77  
 08/31/98 22:58:00 + 3:10 Cali: 30068109 # 3 RIC: 15136.  
 Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

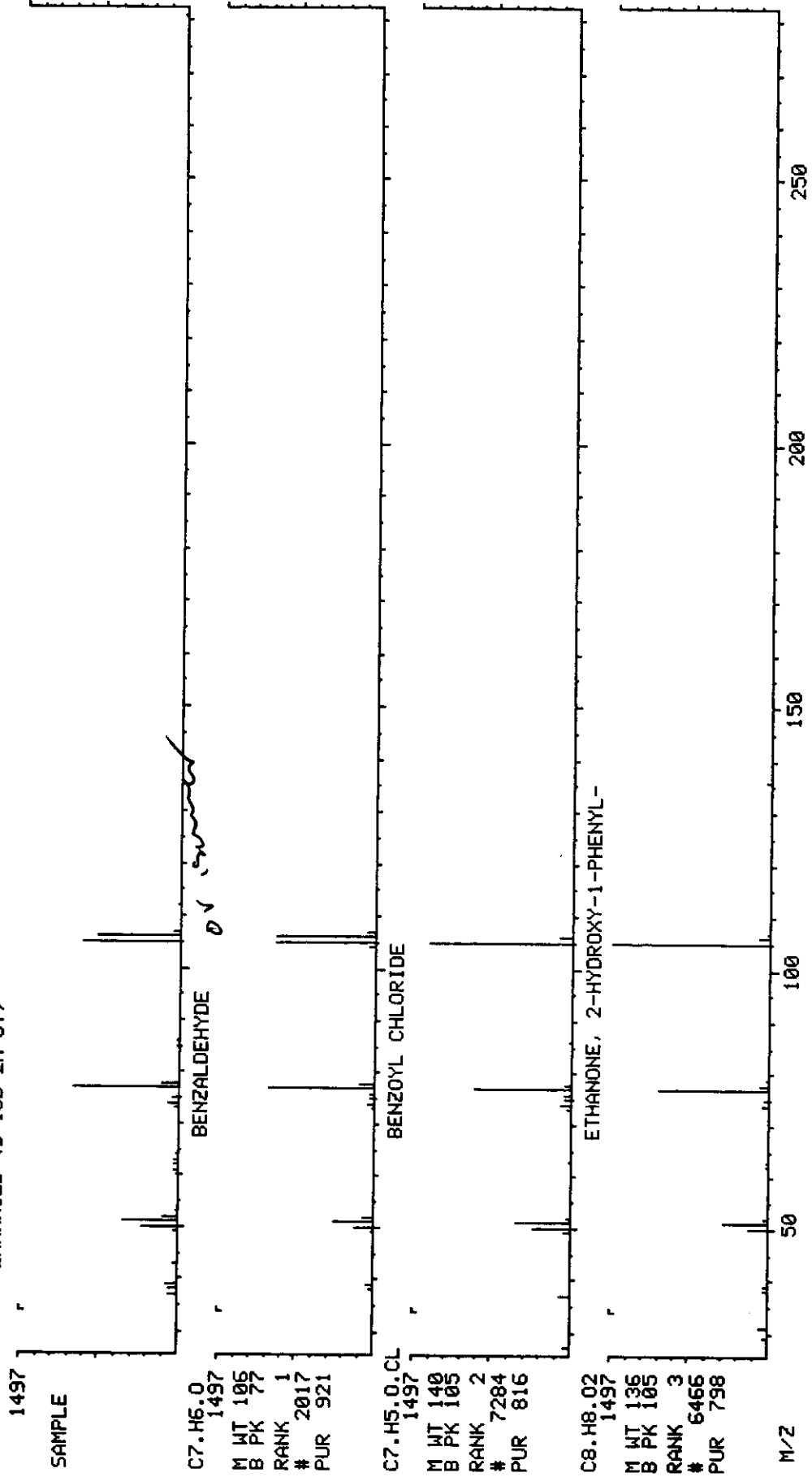
62231 spectra in LIBRARYNB searched for maximum PURITY  
 352 matched at least 5 of the 16 largest peaks in the unknown

Rank In. Name  
 1 2017 BENZALDEHYDE  
 2 7284 BENZOYL CHLORIDE  
 3 6466 ETHANONE, 2-HYDROXY-1-PHENYL-  
 4 2014 2,4-HEPTADIEN-6-YNAL, (E,E)-  
 5 9645 BENZOYLFORMIC ACID  
 6 9261 1,2-PROPANEDIONE, 1-PHENYL-  
 7 13112 BENZOYL ISOTHIOCYANATE  
 8 6857 BENZENECARBOTHIOIC ACID  
 9 13359 BENZENEACETIC ACID, .ALPHA.-OXO-, METHYL ESTER

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H6.O	106	77	921	966	942
2	C7.H5.O.CL	140	105	816	876	829
3	C8.H8.O2	136	105	798	839	819
4	C7.H6.O	106	105	793	838	803
5	C8.H6.O3	150	105	781	848	806
6	C9.H8.O2	148	105	775	822	801
7	C8.H5.O.N.S	163	105	773	821	789
8	C7.H6.O.S	138	77	763	846	809
9	C9.H8.O3	164	105	761	835	788

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	100-52-7
2	---	---	---	---	98-88-4
3	---	---	---	---	582-24-1
4	---	---	---	---	7200-04-6
5	---	---	---	---	611-73-4
6	---	---	---	---	579-07-7
7	---	---	---	---	532-55-8
8	---	---	---	---	98-91-9
9	---	---	---	---	15206-55-0

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 3:10  
 SAMPLE: S-MM5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 # 285  
 CALI: 30068109 # 3  
 BASE M/Z: 77  
 RIC: 15136.



1272



Library Search Data: 30068109 # 932 Base m/z: 71  
 08/31/98 22:58:00 + 10:22 Cali: 30068109 # 3 RIC: 14704.  
 Sample: S-MM5-R8 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

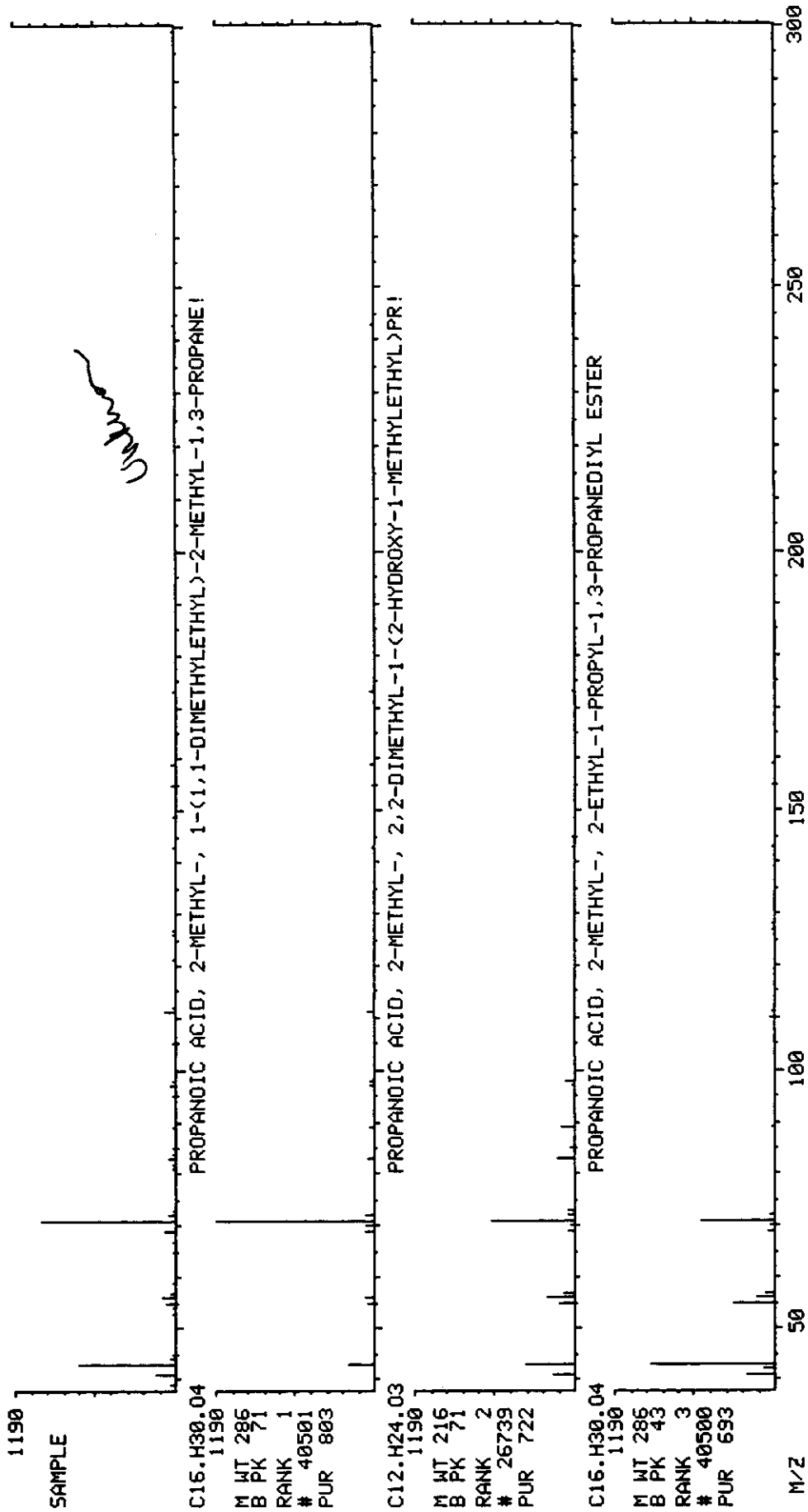
62231 spectra in LIBRARYNB searched for maximum PURITY  
 552 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	40501 PROPANOIC ACID, 2-METHYL-, 1-(1,1-DIMETHYLETHYL)-2-METHYL-1,3-PROPA*
2	26739 PROPANOIC ACID, 2-METHYL-, 2,2-DIMETHYL-1-(2-HYDROXY-1-METHYLETHYL)*
3	40500 PROPANOIC ACID, 2-METHYL-, 2-ETHYL-1-PROPYL-1,3-PROPANEDIYL ESTER
4	4625 1-HEXENE, 3,4,5-TRIMETHYL-
5	4556 ETHANONE, 1-(3-ETHYLCYCLOBUTYL)-
6	2720 4,4-DIMETHYL-1-HEXENE
7	4677 2,4,4-TRIMETHYL-1-HEXENE
8	15779 1,2-CYCLOHEXANEDIOL, 1-METHYL-4-(1-METHYLETHYL)-
9	14787 1-NONENE, 4,6,8-TRIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H30.O4	286	71	803	918	825
2	C12.H24.O3	216	71	722	883	741
3	C16.H30.O4	286	43	693	841	748
4	C9.H18	126	43	608	810	687
5	C8.H14.O	126	43	593	745	622
6	C8.H16	112	71	592	789	645
7	C9.H18	126	71	556	716	656
8	C10.H20.O2	172	71	528	696	642
9	C12.H24	168	43	519	694	556

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	74381-40-1
2	---	---	---	---	74367-33-2
3	---	---	---	---	74367-30-9
4	---	---	---	---	56728-10-0
5	---	---	---	---	56335-71-8
6	---	---	---	---	1647-08-1
7	---	---	---	---	51174-12-0
8	---	---	---	---	33669-76-0
9	---	---	---	---	54410-98-9

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 22:58:00 + 10:22  
 SAMPLE: 5-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA) /1/35A NA M  
 ENHANCED (S 158 2N 0T)  
 DATA: 30068109 # 932 BASE M/Z: 71  
 CALI: 30068109 # 3 RIC: 14704.



Library Search                      Data: 30068109 #1236                      Base m/z: 73  
 08/31/98 22:58:00 + 13:45                      Cali: 30068109 # 3                      RIC: 27392.  
 Sample: S-MM5-RB 1/3SA/1ML                      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 750 matched at least 6 of the 16 largest peaks in the unknown

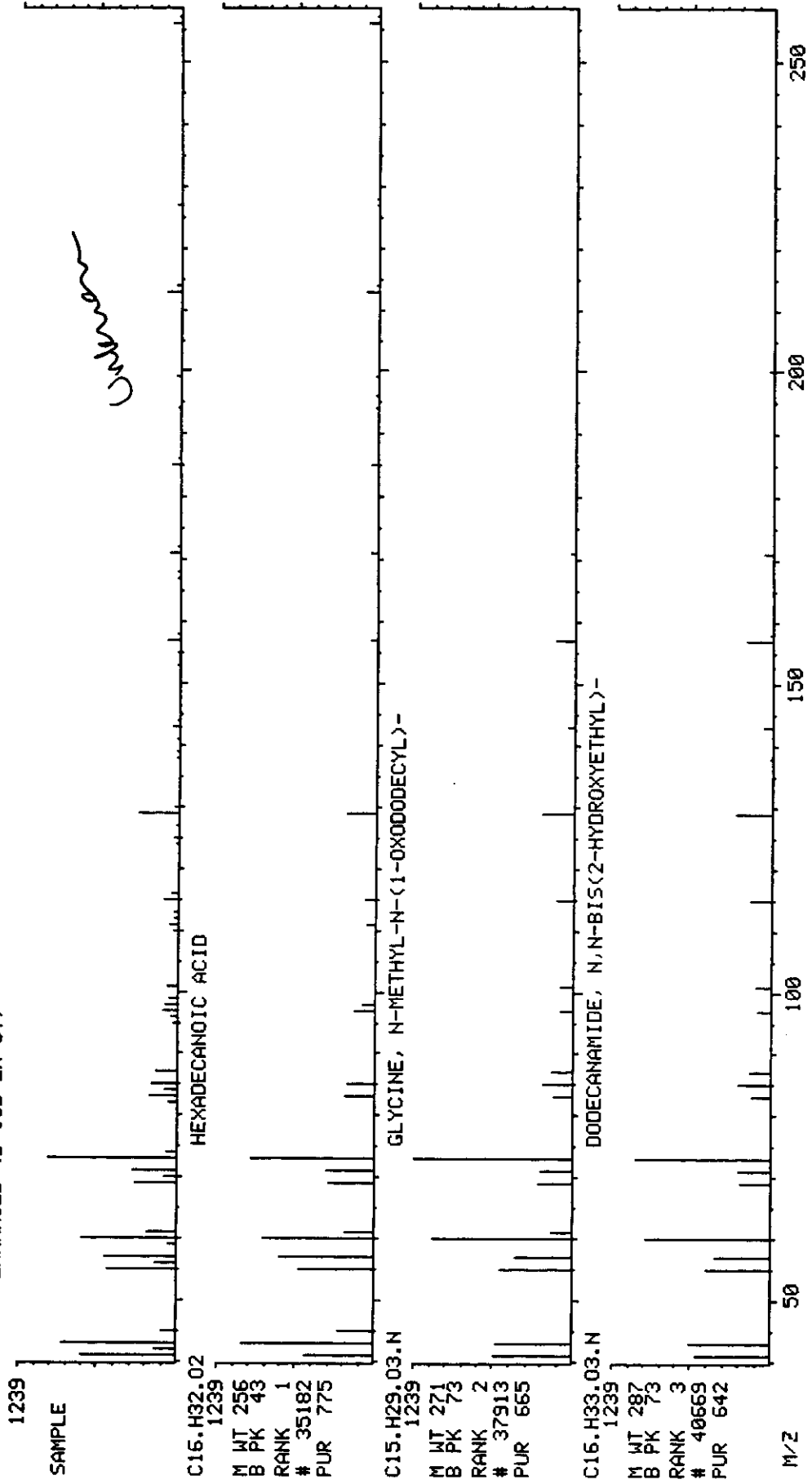
- Rank In.                      Name  
 1 35182 HEXADECANOIC ACID  
 2 37913 GLYCINE, N-METHYL-N-(1-OXODODECYL)-  
 3 40669 DODECANAMIDE, N,N-BIS(2-HYDROXYETHYL)-  
 4 29642 TETRADECANOIC ACID  
 5 26351 TRIDECANOIC ACID  
 6 40184 OCTADECANOIC ACID  
 7 22949 DODECANOIC ACID  
 8 19469 UNDECANOIC ACID  
 9 15784 DECANOIC ACID

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H32.O2	256	43	775	976	778
2	C15.H29.O3.N	271	73	665	979	665
3	C16.H33.O3.N	287	73	642	968	650
4	C14.H28.O2	228	73	623	900	671
5	C13.H26.O2	214	73	607	892	627
6	C18.H36.O2	284	43	604	768	766
7	C12.H24.O2	200	60	604	907	655
8	C11.H22.O2	186	60	597	949	603
9	C10.H20.O2	172	60	573	955	577

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	57-10-3
2	—	—	—	—	97-78-9
3	—	—	—	—	120-40-1
4	—	—	—	—	544-63-8
5	—	—	—	—	638-53-9
6	—	—	—	—	57-11-4
7	—	—	—	—	143-07-7
8	—	—	—	—	112-37-8
9	—	—	—	—	334-48-5

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 13:45  
 SAMPLE: S-MM5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #1236  
 CALI: 30068109 # 3  
 BASE M/Z: 73  
 RIC: 27392.



*Unknown*

11 20 01

Library Search                    Data: 30068109 #1333            Base m/z: 59  
 08/31/98 22:58:00 + 14:50      Cali: 30068109 # 3            RIC: 7896.  
 Sample: S-MMS-RB 1/3SA/1ML      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

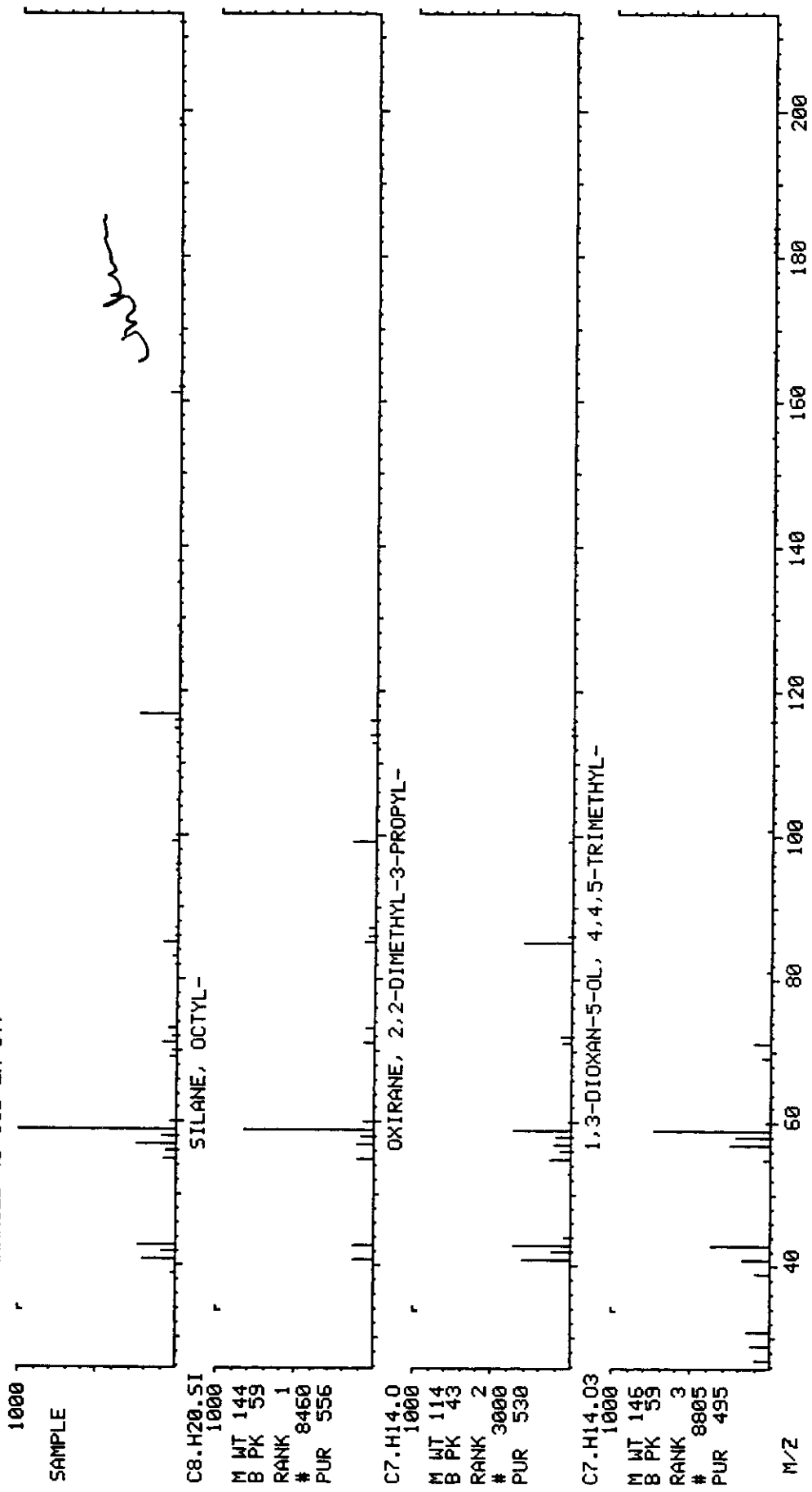
62231 spectra in LIBRARYNB searched for maximum PURITY  
 289 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	8460 SILANE, OCTYL-
2	3000 OXIRANE, 2,2-DIMETHYL-3-PROPYL-
3	8805 1,3-DIOXAN-5-OL, 4,4,5-TRIMETHYL-
4	1513 OXIRANE, 3-ETHYL-2,2-DIMETHYL-
5	19877 BUTANE, 1,1'-[(1-METHYLETHYLIDENE)BIS(OXY)]BIS-
6	8544 2-OCTANOL, 2-METHYL-
7	1576 OXIRANE, TETRAMETHYL-
8	1556 OXIRANE, 2,3-DIETHYL-
9	12073 2-METHYL-2-NONANOL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C8.H20.SI	144	59	556	891	577
2	C7.H14.O	114	43	530	828	546
3	C7.H14.O3	146	59	495	896	530
4	C6.H12.O	100	59	495	853	506
5	C11.H24.O2	188	59	488	690	504
6	C9.H20.O	144	59	483	827	495
7	C6.H12.O	100	59	483	853	491
8	C6.H12.O	100	58	464	749	477
9	C10.H22.O	158	59	454	839	497

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	871-92-1
2	---	---	---	---	17612-35-0
3	---	---	---	---	54063-14-8
4	---	---	---	---	1192-22-9
5	---	---	---	---	141-72-0
6	---	---	---	---	628-44-4
7	---	---	---	---	5076-20-0
8	---	---	---	---	4468-66-0
9	---	---	---	---	10297-57-1

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 14:50  
 SAMPLE: S-MMS-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)  
 DATA: 30068109 #1333  
 CALI: 30068109 # 3  
 BASE M/Z: 59  
 RIC: 7896.



Library Search                    Data: 30068109 #1358            Base m/z: 59  
 08/31/98 22:58:00 + 15:06      Cali: 30068109 # 3            RIC: 5376.  
 Sample: S-MM5-RB 1/3SA/1ML      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

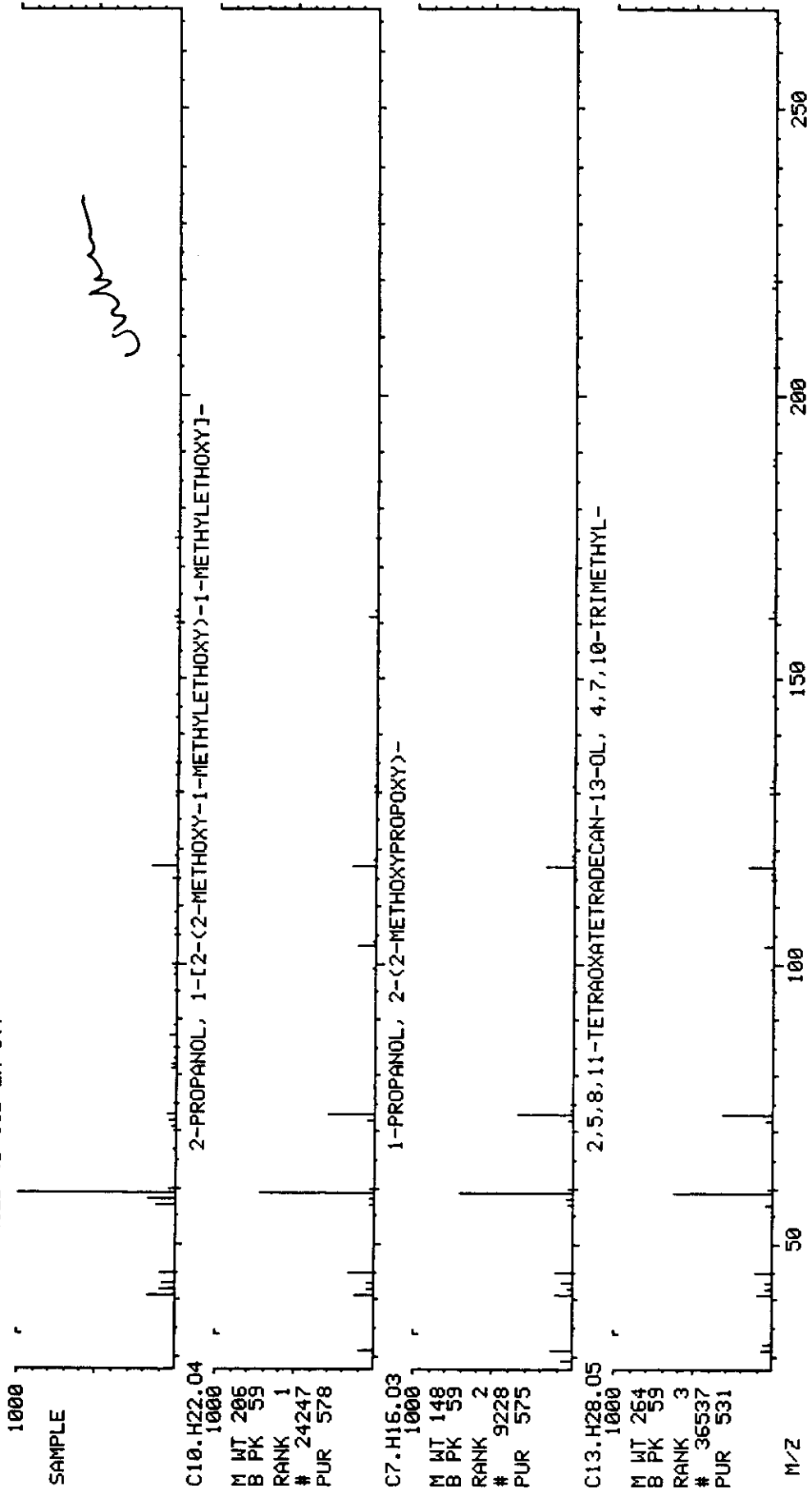
62231 spectra in LIBRARYNB searched for maximum PURITY  
 875 matched at least 5 of the 16 largest peaks in the unknown

Rank In.	Name
1	24247 2-PROPANOL, 1-(2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY)-
2	9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-
3	36537 2,5,8,11-TETRAOXATETRADECAN-13-OL, 4,7,10-TRIMETHYL-
4	5826 2-PROPANOL, 1-ISOPROPOXY-2-METHYL-
5	1775 PENTANE, 2-METHOXY-
6	5786 PROPANOIC ACID, 2-HYDROXY-2-METHYL-, ETHYL ESTER
7	1905 2-BUTANOL, 3-METHOXY-
8	9229 2-PROPANOL, 1-(2-METHOXYPROPOXY)-
9	848 3-PENTANOL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H22.O4	206	59	578	763	646
2	C7.H16.O3	148	59	575	824	595
3	C13.H28.O5	264	59	531	753	624
4	C7.H16.O2	132	59	517	745	540
5	C6.H14.O	102	59	511	903	530
6	C6.H12.O3	132	59	510	817	539
7	C5.H12.O2	104	59	504	904	521
8	C7.H16.O3	148	59	503	841	558
9	C5.H12.O	88	59	499	916	537

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	20324-33-8
2	---	---	---	---	13588-28-8
3	---	---	---	---	20324-34-9
4	---	---	---	---	3587-75-5
5	---	---	---	---	6795-88-6
6	---	---	---	---	80-55-7
7	---	---	---	---	53778-72-6
8	---	---	---	---	13429-07-7
9	---	---	---	---	584-02-1

MID LIBRARY SEARCH (LIBRARYNB) DATA: 30068109 #1358 BASE M/Z: 59  
 08/31/98 22:58:00 + 15:06 CALI: 30068109 # 3 RIC: 5376.  
 SAMPLE: S-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML #100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)



12800



Library Search                      Data: 30068109 #1466                      Base m/z: 57  
 08/31/98 22:58:00 + 16:18                      Cali: 30068109 # 3                      RIC: 5872.  
 Sample: S-MM5-RB 1/3SA/1ML                      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

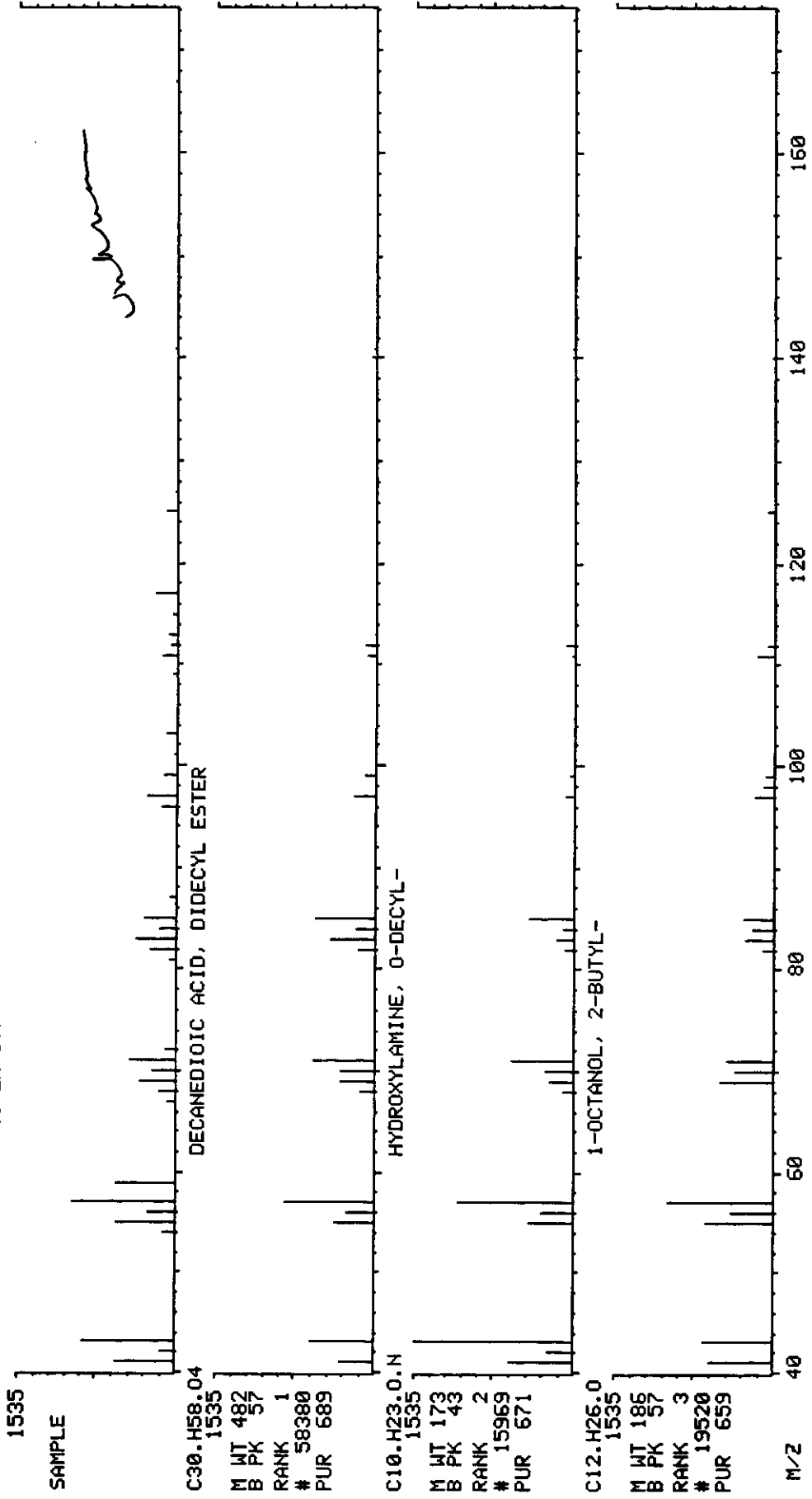
62231 spectra in LIBRARYNB searched for maximum PURITY  
 920 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.                      Name  
 1 58380 DECANEDIOIC ACID, DIDECYL ESTER  
 2 15969 HYDROXYLAMINE, O-DECYL-  
 3 19520 1-OCTANOL, 2-BUTYL-  
 4 19525 2-METHYL-1-UNDECANOL  
 5 42180 OCTADECANE, 1-(ETHENYLOXY)-  
 6 60775 HEXADECANE, 1,1-BIS(DODECYLOXY)-  
 7 60679 PHOSPHONIC ACID, DIOCTADECYL ESTER  
 8 19523 1-DECANOL, 2-ETHYL-  
 9 40233 NONADECANOL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C30.H58.O4	482	57	689	957	699
2	C10.H23.O.N	173	43	671	918	681
3	C12.H26.O	186	57	659	918	701
4	C12.H26.O	186	57	652	935	681
5	C20.H40.O	296	43	652	925	690
6	C40.H82.O2	594	43	647	873	730
7	C36.H75.O3.P	586	55	646	917	677
8	C12.H26.O	186	57	645	928	669
9	C19.H40.O	284	43	639	915	680

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	2432-89-5
2	—	—	—	—	29812-79-1
3	—	—	—	—	3913-02-8
4	—	—	—	—	10522-26-6
5	—	—	—	—	930-02-9
6	—	—	—	—	56554-64-4
7	—	—	—	—	19047-85-9
8	—	—	—	—	21078-65-9
9	—	—	—	—	52783-43-4

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 16:18  
 SAMPLE: 5-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*1002/1002 \*(NA/NA) /1/35A NA M  
 ENHANCED (S 158 2N 0T)  
 DATA: 30068109 #1455  
 CALI: 30068109 # 3  
 BASE M/Z: 57  
 RIC: 5872.



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Library Search                      Data: 30068109 #1502                      Base m/z: 45  
 08/31/98 22:58:00 + 16:42                      Cali: 30068109 # 3                      RIC: 3992.  
 Sample: S-MM5-RB                      1/3SA/1ML                      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 325 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	3365 2-PENTANOL, 4,4-DIMETHYL-
2	3362 4-METHYL-2-HEXANOL
3	1789 2-HEXANOL
4	6417 2-PENTANOL, 3-CHLORO-4-METHYL-, (R@,R@)-(./-.)-
5	8510 2-NONANOL
6	3188 N-NITROSO-2-METHYL-OXAZOLIDINE
7	3060 OXIRANE, 2-BUTYL-3-METHYL-, CIS-
8	23326 2-PROPANONE, 1,1-DIBUTOXY-
9	15842 2-UNDECANOL

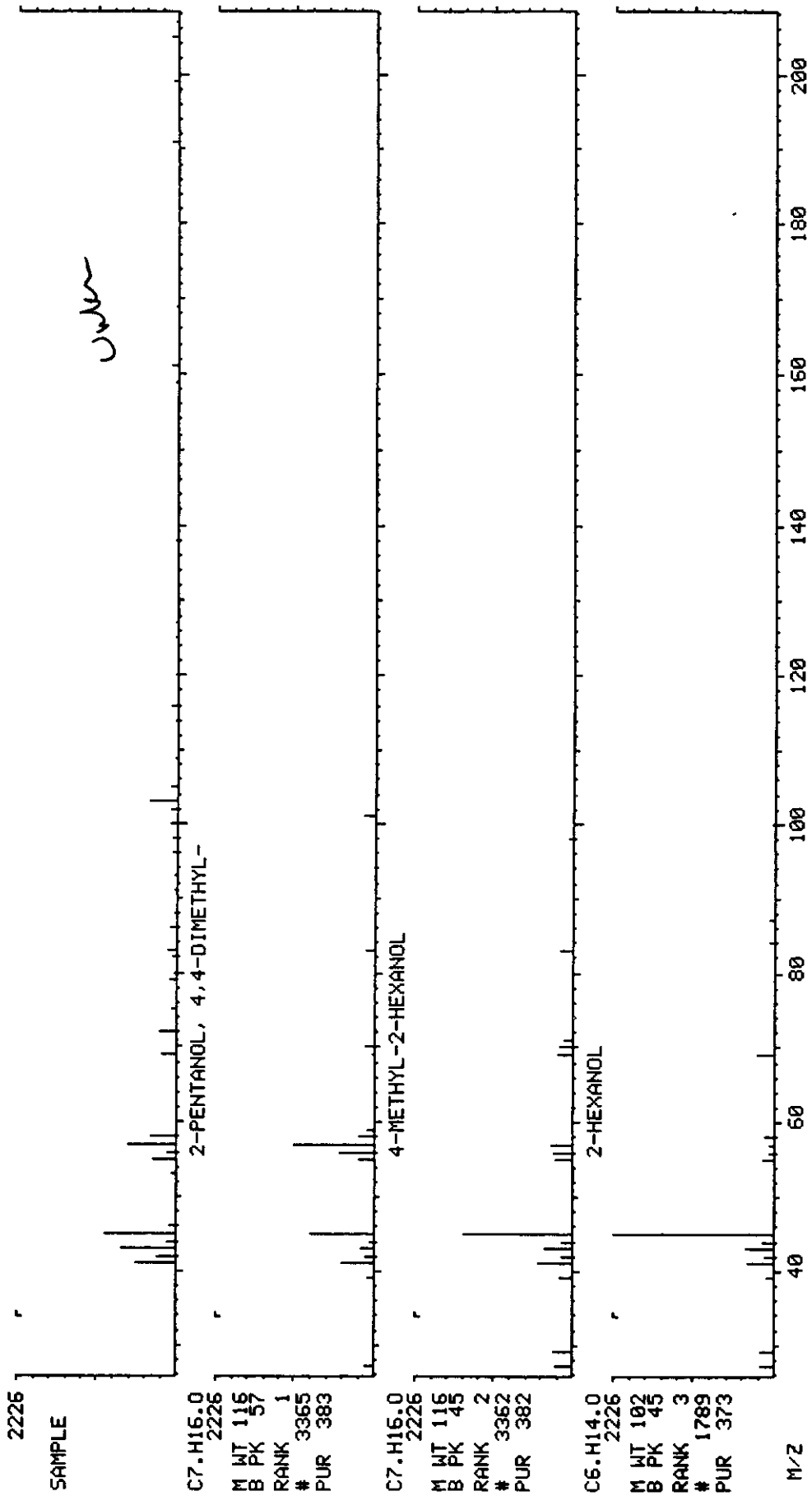
Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C7.H16.O	116	57	383	760	443
2	C7.H16.O	116	45	382	818	437
3	C6.H14.O	102	45	373	741	405
4	C6.H13.O.CL	136	45	369	792	412
5	C9.H20.O	144	45	367	786	431
6	C4.H8.O2.N2	116	43	362	703	406
7	C7.H14.O	114	43	358	722	465
8	C11.H22.O3	202	57	354	655	383
9	C11.H24.O	172	45	345	739	443

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	6144-93-0
2	---	---	---	---	2313-61-3
3	---	---	---	---	626-93-7
4	---	---	---	---	74685-47-5
5	---	---	---	---	628-99-9
6	---	---	---	---	39884-53-2
7	---	---	---	---	56052-93-8
8	---	---	---	---	19255-82-4
9	---	---	---	---	1653-30-1

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 16:42  
 SAMPLE: 5-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M  
 ENHANCED (S 158 2H 0T)

DATA: 30068109 #1502  
 CALI: 30068109 # 3

BASE M/Z: 45  
 RIC: 3992.



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Library Search Data: 30068109 #1509 Base m/z: 59  
 08/31/98 22:58:00 + 16:47 Cali: 30068109 # 3 RIC: 12128.  
 Sample: S-MM5-RB 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 329 matched at least 6 of the 16 largest peaks in the unknown

Rank In. Name  
 1 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-  
 2 16192 2-PROPANOL, 1-[1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-  
 3 20626 2-PROPANOL, 1,1'-[[1-METHYL-1,2-ETHANEDIYL)BIS(OXY)]BIS-  
 4 3522 2,3-BUTANEDIOL, 2,3-DIMETHYL-  
 5 9235 1-PROPANOL, 2-(2-METHOXY-1-METHYLETHOXY)-  
 6 5778 BUTANOIC ACID, 3-HYDROXY-3-METHYL-, METHYL ESTER  
 7 12385 BUTYRIC ACID, 4-ISOPROPOXY-, METHYL ESTER  
 8 8432 2-HEXANONE, 3-HYDROXY-3,5-DIMETHYL-  
 9 8822 BUTANOIC ACID, 4-ETHOXY-, METHYL ESTER

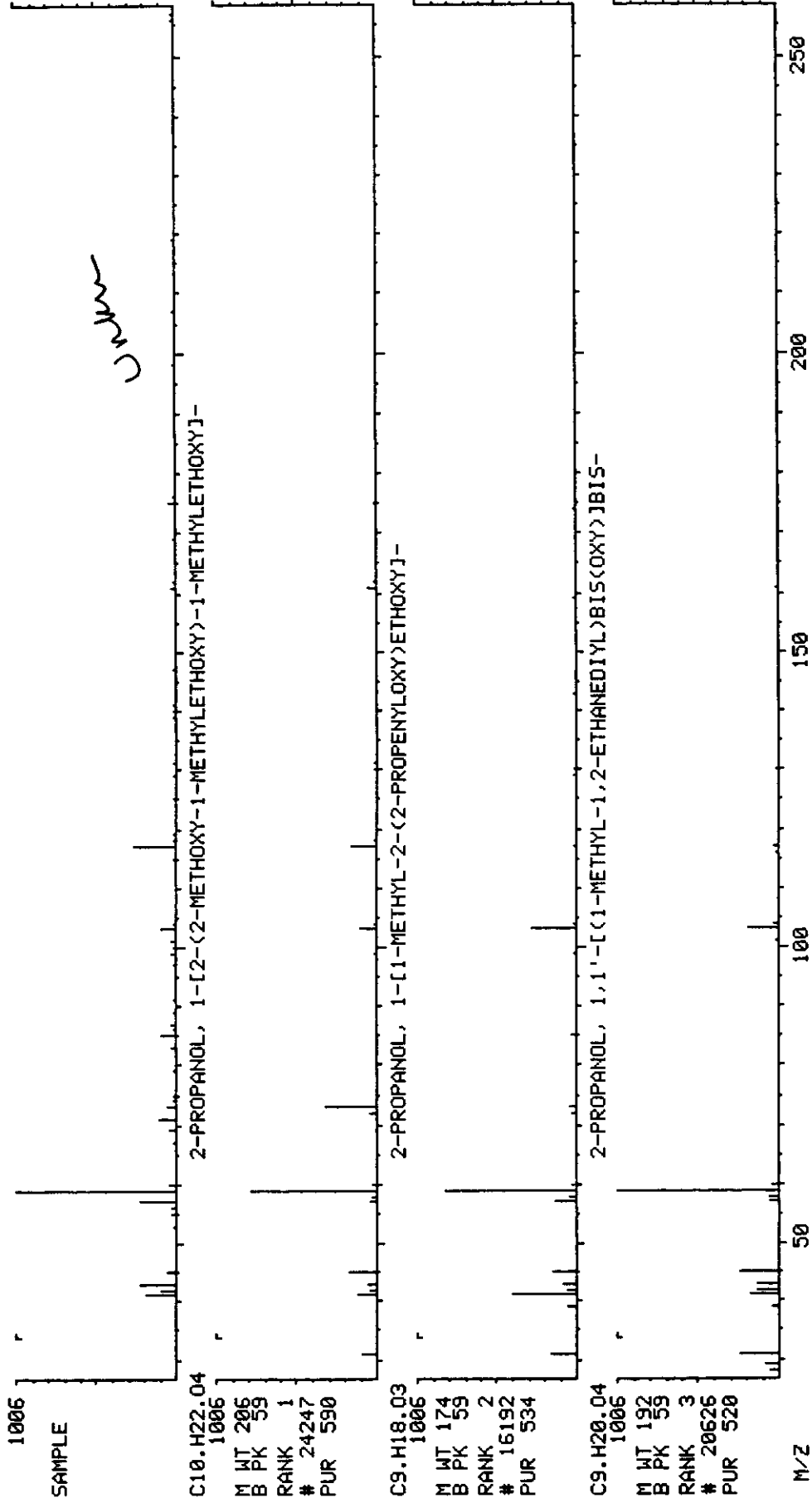
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H22.O4	206	59	590	796	614
2	C9.H18.O3	174	59	534	748	564
3	C9.H20.O4	192	59	520	780	546
4	C6.H14.O2	118	59	509	919	522
5	C7.H16.O3	148	59	485	727	488
6	C6.H12.O3	132	43	479	827	496
7	C8.H16.O3	160	85	467	682	496
8	C8.H16.O2	144	59	466	933	475
9	C7.H14.O3	146	59	457	685	483

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	20324-33-8
2	---	---	---	---	55956-25-7
3	---	---	---	---	1638-16-0
4	---	---	---	---	76-09-5
5	---	---	---	---	55956-21-3
6	---	---	---	---	6149-45-7
7	---	---	---	---	29006-05-1
8	---	---	---	---	6321-14-8
9	---	---	---	---	29006-04-0

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 16:47  
 SAMPLE: S-MMS-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (<S 15B 2N 0T)

DATA: 30068109 #1509  
 CALI: 30068109 # 3

BASE M/Z: 53  
 RIC: 12128.



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Library Search                    Data: 30068109 #1529            Base m/z: 59  
 08/31/98 22:58:00 + 17:00        Cali: 30068109 # 3            RIC: 10864.  
 Sample: S-MM5-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 691 matched at least 6 of the 16 largest peaks in the unknown

- Rank In.            Name  
 1 3522 2,3-BUTANEDIOL, 2,3-DIMETHYL-  
 2 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-  
 3 16192 2-PROPANOL, 1-[1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-  
 4 5497 3-ETHYL-2-METHYL-2-PENTANOL  
 5 12073 2-METHYL-2-NONANOL  
 6 8432 2-HEXANONE, 3-HYDROXY-3,5-DIMETHYL-  
 7 8460 SILANE, OCTYL-  
 8 20626 2-PROPANOL, 1,1'-[[1-METHYL-1,2-ETHANEDIYL]BIS(OXY)]BIS-  
 9 6419 2-PENTANOL, 3-CHLORO-2-METHYL-

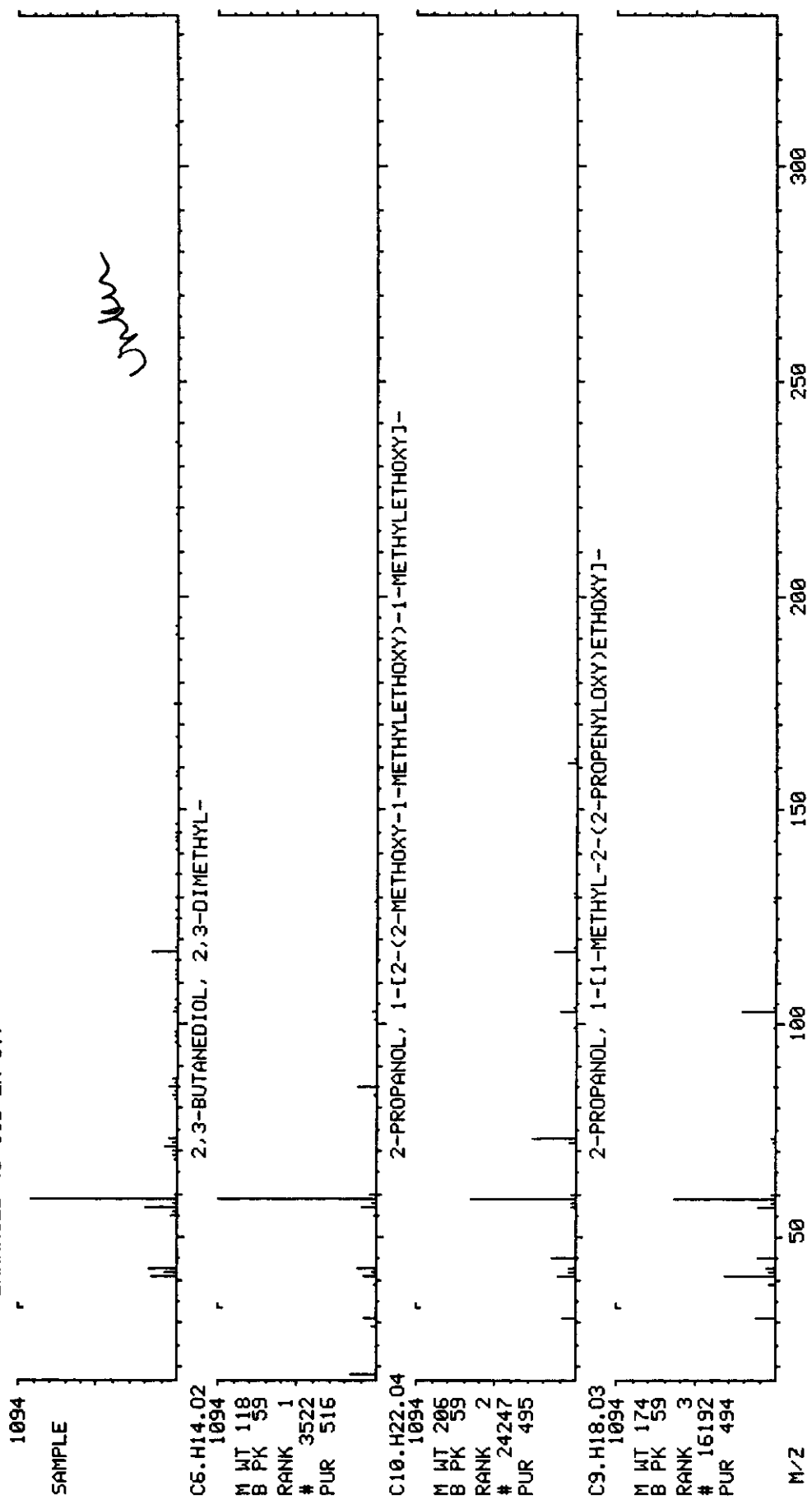
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H14.O2	118	59	516	931	528
2	C10.H22.O4	206	59	495	719	580
3	C9.H18.O3	174	59	494	710	537
4	C8.H18.O	130	59	472	919	494
5	C10.H22.O	158	59	471	846	491
6	C8.H16.O2	144	59	468	852	537
7	C8.H20.SI	144	59	461	781	564
8	C9.H20.O4	192	59	459	734	519
9	C6.H13.O.CL	136	59	457	844	495

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	76-09-5
2	---	---	---	---	20324-33-8
3	---	---	---	---	55956-25-7
4	---	---	---	---	-
5	---	---	---	---	10297-57-1
6	---	---	---	---	6321-14-8
7	---	---	---	---	871-92-1
8	---	---	---	---	1638-16-0
9	---	---	---	---	74685-49-7

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 17:00  
 SAMPLE: S-MMS-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100Z \*(NA/NA) /1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068109 #1529  
 CALI: 30068109 # 3

BASE M/Z: 59  
 RIC: 10854.



10854



Library Search                    Data: 30068109 #1587            Base m/z: 57  
 08/31/98 22:58:00 + 17:39        Cali: 30068109 # 3            RIC: 68224.  
 Sample: S-MM5-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 950 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 39516 5-EICOSENE, (E)-  
 2 34410 3-OCTADECENE, (E)-  
 3 39515 9-EICOSENE, (E)-  
 4 34411 5-OCTADECENE, (E)-  
 5 34418 9-OCTADECENE, (E)-  
 6 31653 1-HEPTADECENE  
 7 39517 3-EICOSENE, (E)-  
 8 37062 1-NONADECENE  
 9 26416 4-TETRADECANOL

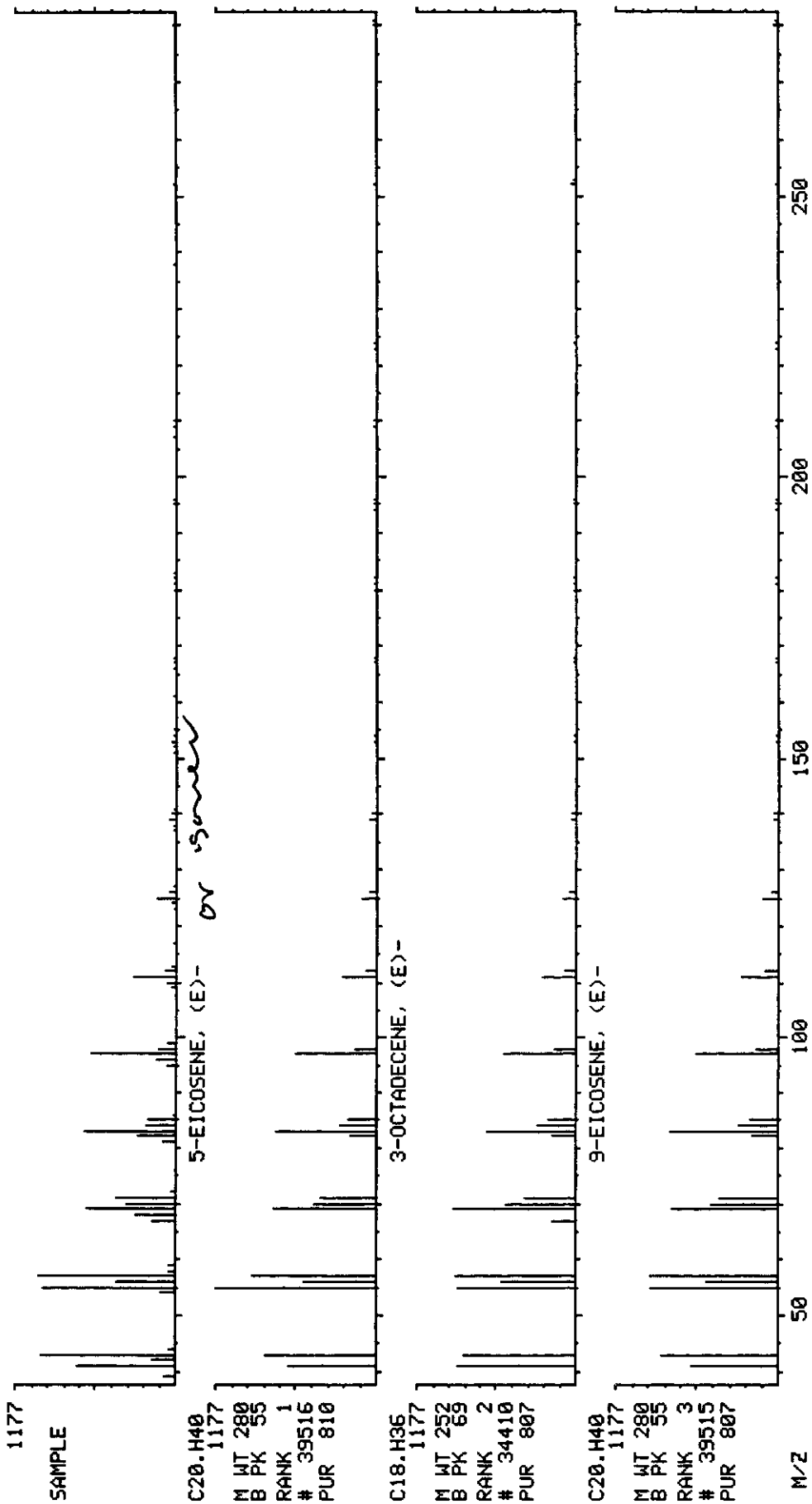
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C20.H40	280	55	810	986	813
2	C18.H36	252	69	807	973	811
3	C20.H40	280	55	807	985	810
4	C18.H36	252	55	804	971	806
5	C18.H36	252	55	803	970	806
6	C17.H34	238	55	800	981	801
7	C20.H40	280	57	797	988	799
8	C19.H38	266	97	794	971	802
9	C14.H30.O	214	43	793	990	795

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	74685-30-6
2	---	---	---	---	7206-19-1
3	---	---	---	---	74685-29-3
4	---	---	---	---	7206-21-5
5	---	---	---	---	7206-25-9
6	---	---	---	---	6765-39-5
7	---	---	---	---	74685-33-9
8	---	---	---	---	18435-45-5
9	---	---	---	---	1653-33-4

MID LIBRARY SEARCH (LIBRARY.MB)  
 08/31/98 22:58:00 + 17:39  
 SAMPLE: S-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #1587  
 CALI: 30068109 # 3

BASE M/Z: 57  
 RIC: 58224.



Library Search                      Data: 30068109 #1661                      Base m/z: 59  
 08/31/98 22:58:00 + 18:29                      Cali: 30068109 # 3                      RIC: 9856.  
 Sample: S-MM5-RB 1/3SA/1ML                      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

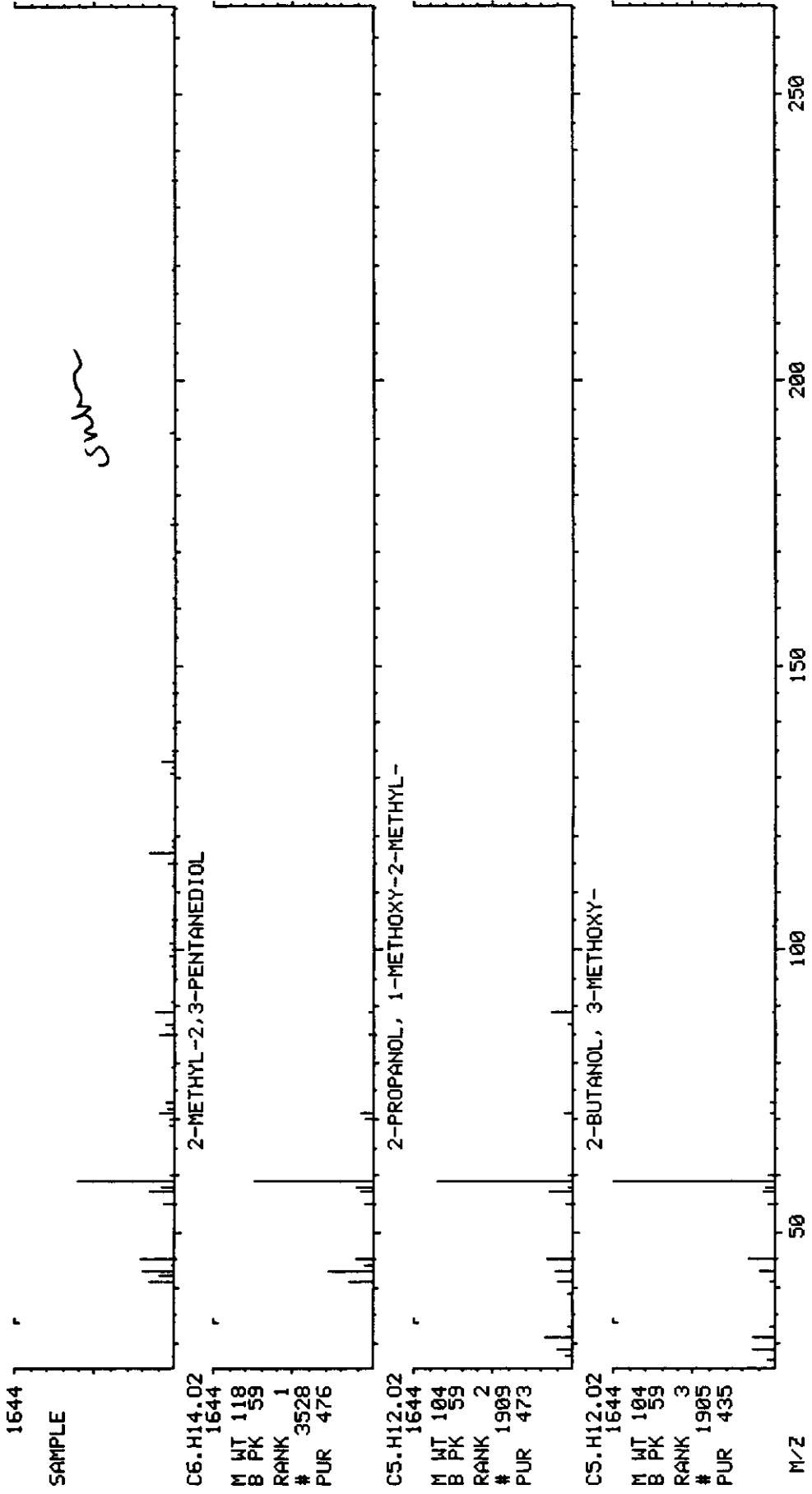
62231 spectra in LIBRARYNB searched for maximum PURITY  
 217 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	3528 2-METHYL-2,3-PENTANEDIOL
2	1909 2-PROPANOL, 1-METHOXY-2-METHYL-
3	1905 2-BUTANOL, 3-METHOXY-
4	3502 BUTANOIC ACID, 3-HYDROXY-3-METHYL-
5	9227 1,3,3-TRIMETHOXYBUTANE
6	8822 BUTANOIC ACID, 4-ETHOXY-, METHYL ESTER
7	5826 2-PROPANOL, 1-ISOPROPOXY-2-METHYL-
8	1919 1-BUTANOL, 3-METHOXY-
9	36510 1,4,7,10,13,16-HEXAOXACYCLOOCTADECANE

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H14.O2	118	59	476	925	476
2	C5.H12.O2	104	59	473	946	477
3	C5.H12.O2	104	59	435	870	435
4	C5.H10.O3	118	59	432	815	489
5	C7.H16.O3	148	45	429	674	442
6	C7.H14.O3	146	59	427	743	515
7	C7.H16.O2	132	59	426	677	490
8	C5.H12.O2	104	59	424	893	437
9	C12.H24.O6	264	45	423	678	455

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	7795-80-4
2	---	---	---	---	3587-64-2
3	---	---	---	---	53778-72-6
4	---	---	---	---	625-08-1
5	---	---	---	---	6607-66-5
6	---	---	---	---	29006-04-0
7	---	---	---	---	3587-75-5
8	---	---	---	---	2517-43-3
9	---	---	---	---	17455-13-9

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 18:29  
 SAMPLE: S-MM5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 #1661  
 CALI: 30068109 # 3  
 BASE M/Z: 59  
 RIC: 9856.



11202

Library Search                    Data: 30068109 #1687            Base m/z: 59  
 08/31/98 22:58:00 + 18:46       Cali: 30068109 # 3            RIC: 6712.  
 Sample: S-MM5-RB 1/3SA/1ML       INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2M 0T)

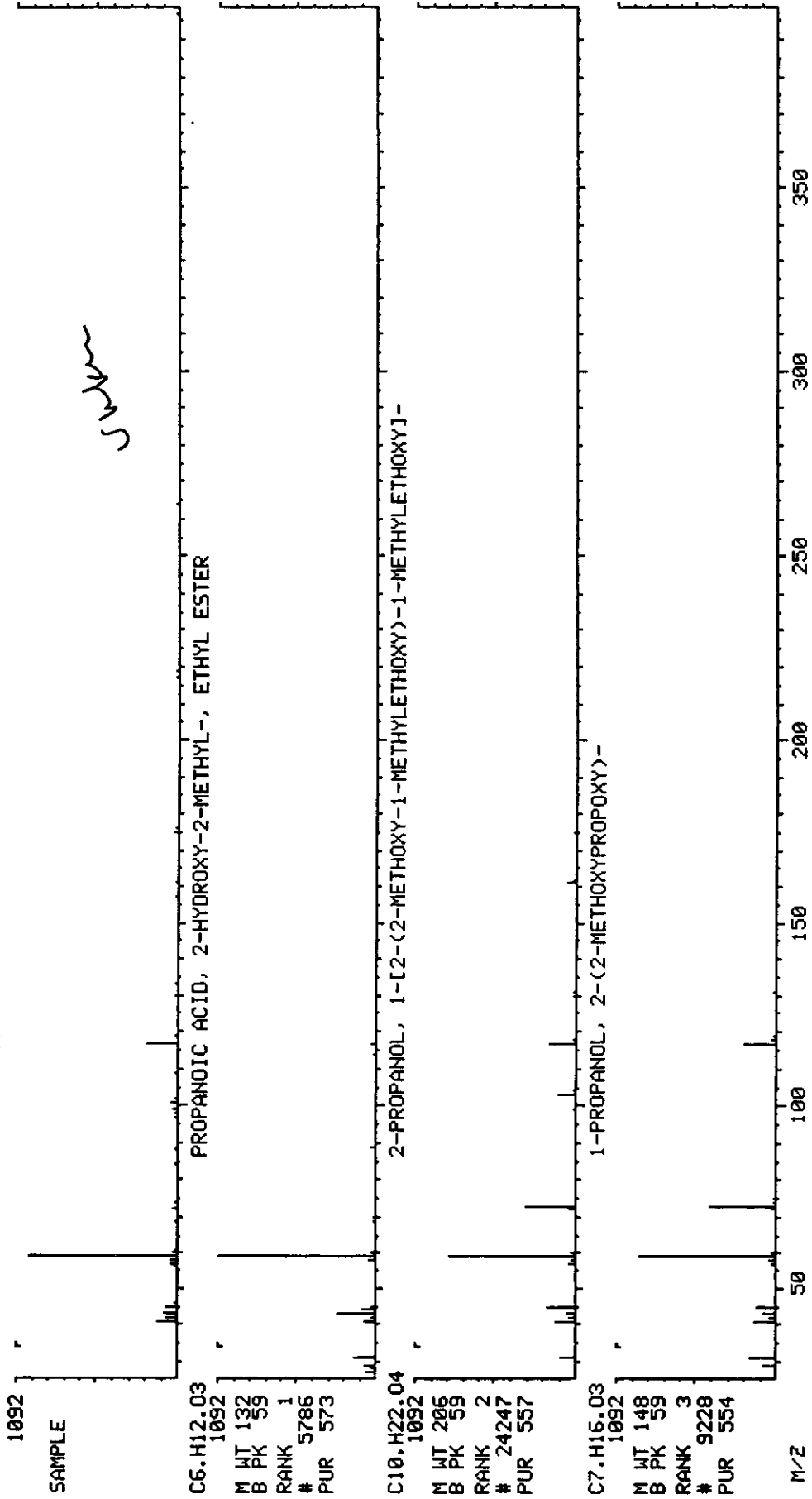
62231 spectra in LIBRARYNB searched for maximum PURITY  
 613 matched at least 5 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 5786 PROPANOIC ACID, 2-HYDROXY-2-METHYL-, ETHYL ESTER  
 2 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-  
 3 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-  
 4 20626 2-PROPANOL, 1,1'-[(1-METHYL-1,2-ETHANEDIYL)BIS(OXY)]BIS-  
 5 36537 2,5,8,11-TETRAOXATETRADECAN-13-OL, 4,7,10-TRIMETHYL-  
 6 926 1-PROPANOL, 2-METHOXY-  
 7 334 2-PROPANOL, 2-METHYL-  
 8 8432 2-HEXANONE, 3-HYDROXY-3,5-DIMETHYL-  
 9 848 3-PENTANOL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H12.O3	132	59	573	800	620
2	C10.H22.O4	206	59	557	664	767
3	C7.H16.O3	148	59	554	737	744
4	C9.H20.O4	192	59	546	698	681
5	C13.H28.O5	264	59	536	667	770
6	C4.H10.O2	90	59	534	944	551
7	C4.H10.O	74	59	524	915	548
8	C8.H16.O2	144	59	517	867	552
9	C5.H12.O	88	59	514	907	553

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	80-55-7
2	---	---	---	---	20324-33-8
3	---	---	---	---	13588-28-8
4	---	---	---	---	1638-16-0
5	---	---	---	---	20324-34-9
6	---	---	---	---	1589-47-5
7	---	---	---	---	75-65-0
8	---	---	---	---	6321-14-8
9	---	---	---	---	584-02-1

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 18:46  
 SAMPLE: 5-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 #1687  
 CALI: 30068109 # 3  
 BASE M/Z: 59  
 RIC: 6712.



1294

Library Search                    Data: 30068109 #1706            Base m/z: 43  
 08/31/98 22:58:00 + 18:59       Cali: 30068109 # 3            RIC: 5680.  
 Sample: S-MM5-RB 1/3SA/1ML       INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

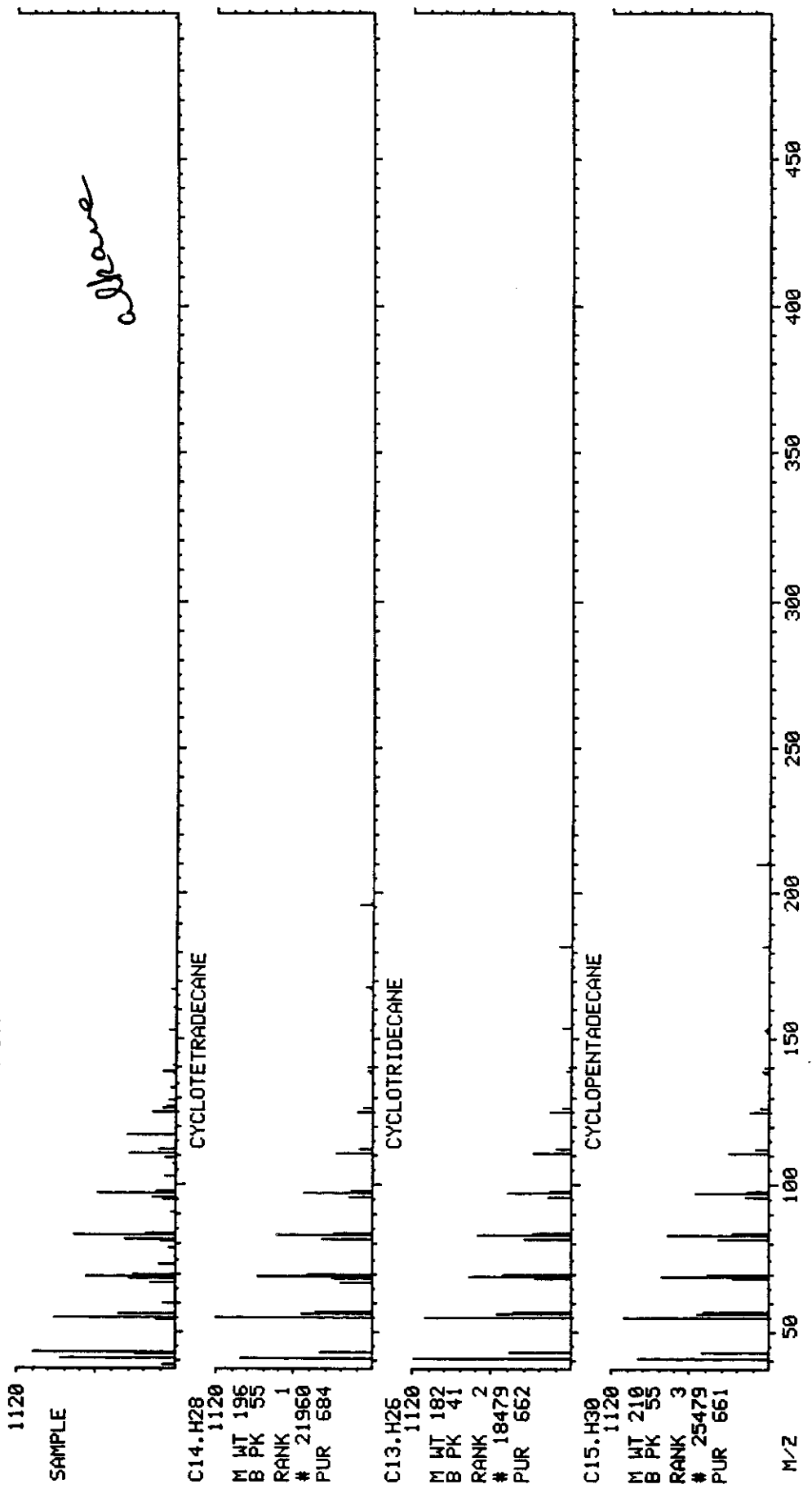
62231 spectra in LIBRARYNB searched for maximum PURITY  
 482 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 21960 CYCLOTETRADECANE  
 2 18479 CYCLOTRIDECANE  
 3 25479 CYCLOPENTADECANE  
 4 14780 CYCLOODECANE  
 5 15841 1-UNDECANOL  
 6 32420 1-HEXADECANOL  
 7 11075 CYCLODECANE, METHYL-  
 8 25482 CYCLOPENTANE, DECYL-  
 9 14765 CYCLOPROPANE, NONYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C14.H28	196	55	684	871	760
2	C13.H26	182	41	662	885	729
3	C15.H30	210	55	661	868	748
4	C12.H24	168	55	661	882	717
5	C11.H24.O	172	55	632	895	670
6	C16.H34.O	242	55	628	855	719
7	C11.H22	154	55	628	904	662
8	C15.H30	210	41	628	827	720
9	C12.H24	168	55	620	835	673

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	295-17-0
2	---	---	---	---	295-02-3
3	---	---	---	---	295-48-7
4	---	---	---	---	294-62-2
5	---	---	---	---	112-42-5
6	---	---	---	---	36653-82-4
7	---	---	---	---	13151-43-4
8	---	---	---	---	1795-21-7
9	---	---	---	---	74663-85-7

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 22:58:00 + 18:59  
 SAMPLE: 5-MM5-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 #1706  
 CALI: 30068109 # 3  
 BASE M/Z: 43  
 RIC: 5680.



1120

SAMPLE

C14.H28  
1120

M WT 196  
 B PK 55  
 RANK 1  
 # 21960  
 PUR 684

C13.H26  
1120

M WT 182  
 B PK 41  
 RANK 2  
 # 18479  
 PUR 662

C15.H30  
1120

M WT 210  
 B PK 55  
 RANK 3  
 # 25479  
 PUR 661

M/Z

1120  
 5680  
 (C)  
 (S)



Library Search                    Data: 30068109 #1741            Base m/z: 59  
 08/31/98 22:58:00 + 19:22        Cali: 30068109 # 3            RIC: 11328.  
 Sample: S-MM5-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 615 matched at least 6 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 39622 9-OCTADECENAMIDE, (Z)-  
 2 11721 NONANAMIDE  
 3 3133 PENTANAMIDE, 4-METHYL-  
 4 15482 HEPTANAMIDE, 4-ETHYL-5-METHYL-  
 5 22656 DODECANAMIDE  
 6 19516 1-DECANOL, 5,9-DIMETHYL-  
 7 7998 TRANS-3,4-EPOXYNONANE  
 8 3132 HEXANAMIDE  
 9 1617 PENTANAMIDE

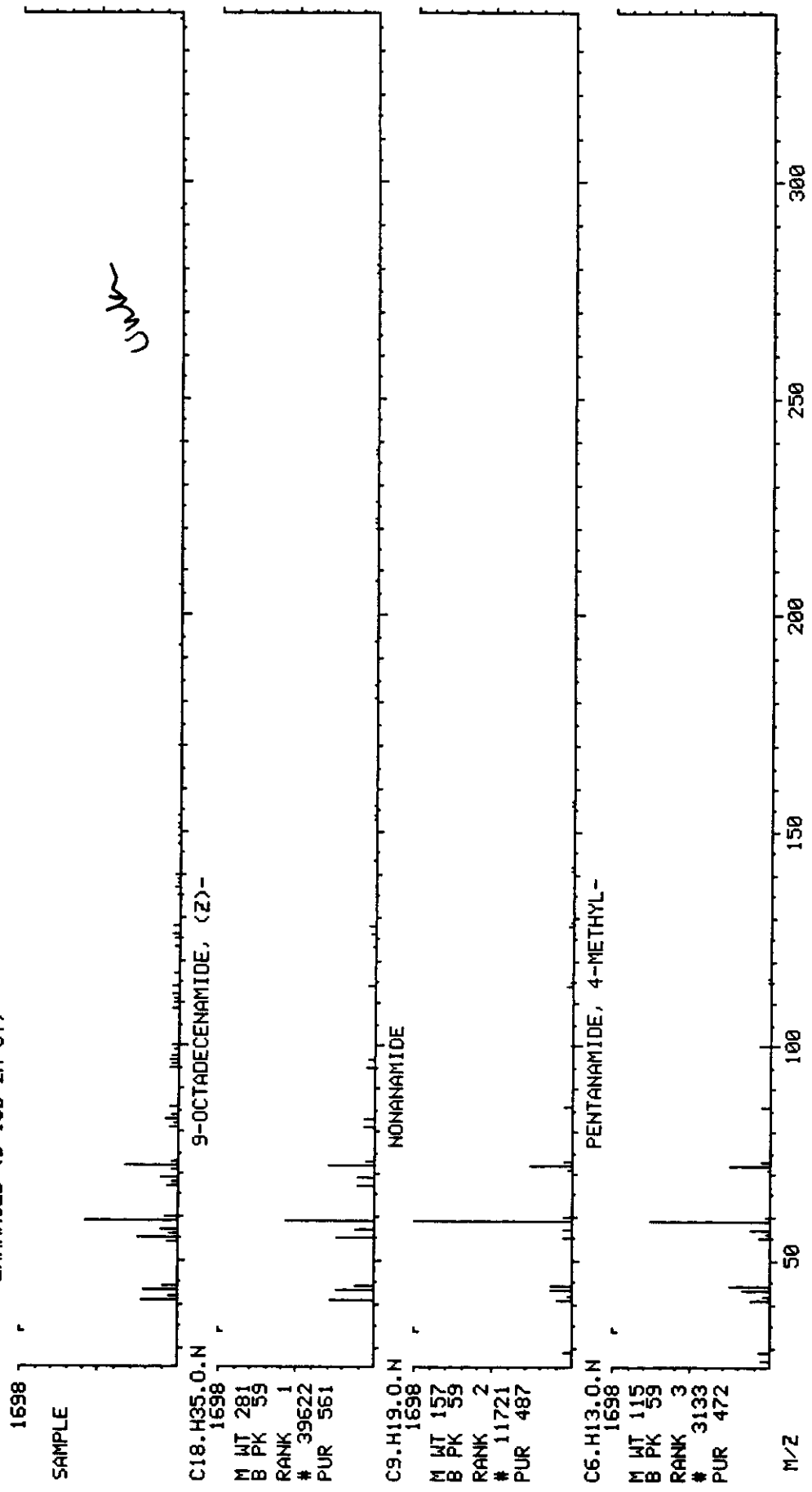
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C18.H35.O.N	281	59	561	866	644
2	C9.H19.O.N	157	59	487	894	498
3	C6.H13.O.N	115	59	472	913	477
4	C10.H21.O.N	171	59	461	860	485
5	C12.H25.O.N	199	59	458	852	515
6	C12.H26.O	186	41	454	767	497
7	C9.H18.O	142	41	438	769	482
8	C6.H13.O.N	115	59	432	866	437
9	C5.H11.O.N	101	59	419	858	428

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	301-02-0
2	---	---	---	---	1120-07-6
3	---	---	---	---	1119-29-5
4	---	---	---	---	54789-40-1
5	---	---	---	---	1120-16-7
6	---	---	---	---	91482-38-1
7	---	---	---	---	- -
8	---	---	---	---	628-02-4
9	---	---	---	---	626-97-1

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 19:22  
 SAMPLE: S-MMS-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #1741  
 CALI: 30068109 # 3

BASE M/Z: 59  
 RIC: 11328.



1202

Library Search                      Data: 30068109 #1801                      Base m/z: 59  
 08/31/98 22:58:00 + 20:02        Cali: 30068109 # 3                      RIC: 8960.  
 Sample: S-MM5-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

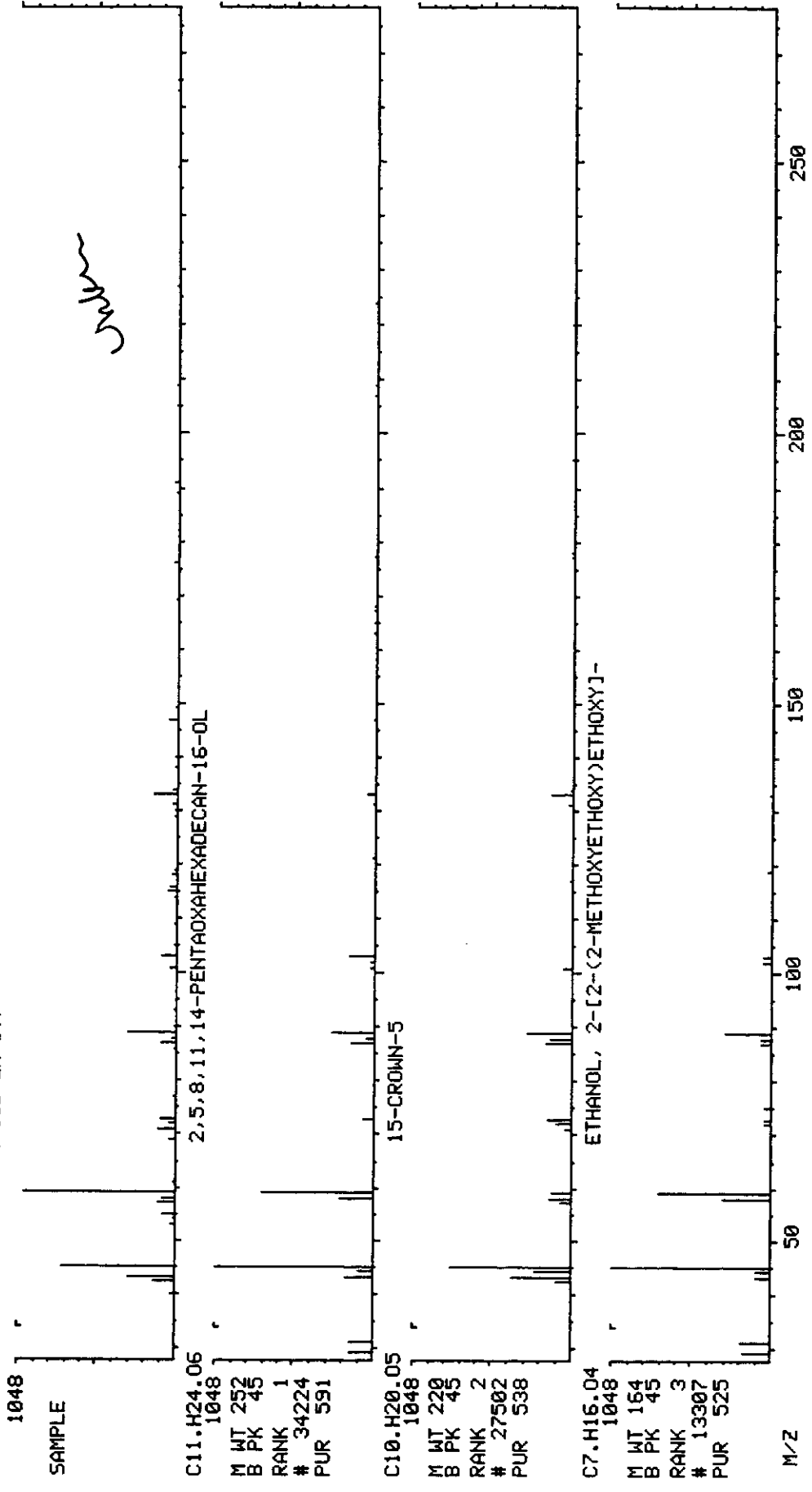
62231 spectra in LIBRARYNB searched for maximum PURITY  
 752 matched at least 5 of the 16 largest peaks in the unknown

Rank In.            Name  
 1 34224 2,5,8,11,14-PENTAOXAHEXADECAN-16-OL  
 2 27502 15-CROWN-5  
 3 13307 ETHANOL, 2-[2-(2-METHOXYETHOXY)ETHOXY]-  
 4 17070 2,5,8,11-TETRAOXADODECANE  
 5 28000 2,5,8,11,14-PENTAOXAPENTADECANE  
 6 36510 1,4,7,10,13,16-HEXAOXACYCLOOCTADECANE  
 7 8813 ETHENE, (2-ETHOXY-1-METHOXYETHOXY)-  
 8 36865 2,5,8,11,14,17-HEXAOXAOCTADECANE  
 9 1909 2-PROPANOL, 1-METHOXY-2-METHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C11.H24.O6	252	45	591	876	632
2	C10.H20.O5	220	45	538	772	590
3	C7.H16.O4	164	45	525	827	565
4	C8.H18.O4	178	59	511	814	532
5	C10.H22.O5	222	59	506	742	533
6	C12.H24.O6	264	45	498	767	555
7	C7.H14.O3	146	59	488	838	504
8	C12.H26.O6	266	59	484	737	520
9	C5.H12.O2	104	59	476	864	510

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	23778-52-1
2	---	---	---	---	33100-27-5
3	---	---	---	---	112-35-6
4	---	---	---	---	112-49-2
5	---	---	---	---	143-24-8
6	---	---	---	---	17455-13-9
7	---	---	---	---	54063-18-2
8	---	---	---	---	1191-87-3
9	---	---	---	---	3587-64-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 20:02  
 SAMPLE: S-MM5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UC/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 #1801 BASE M/Z: 59  
 CALI: 30068109 # 3 RIC: 8960.



Library Search                    Data: 30068109 #1834            Base m/z: 59  
 08/31/98 22:58:00 + 20:24        Cali: 30068109 # 3            RIC: 2840.  
 Sample: S-MMS-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

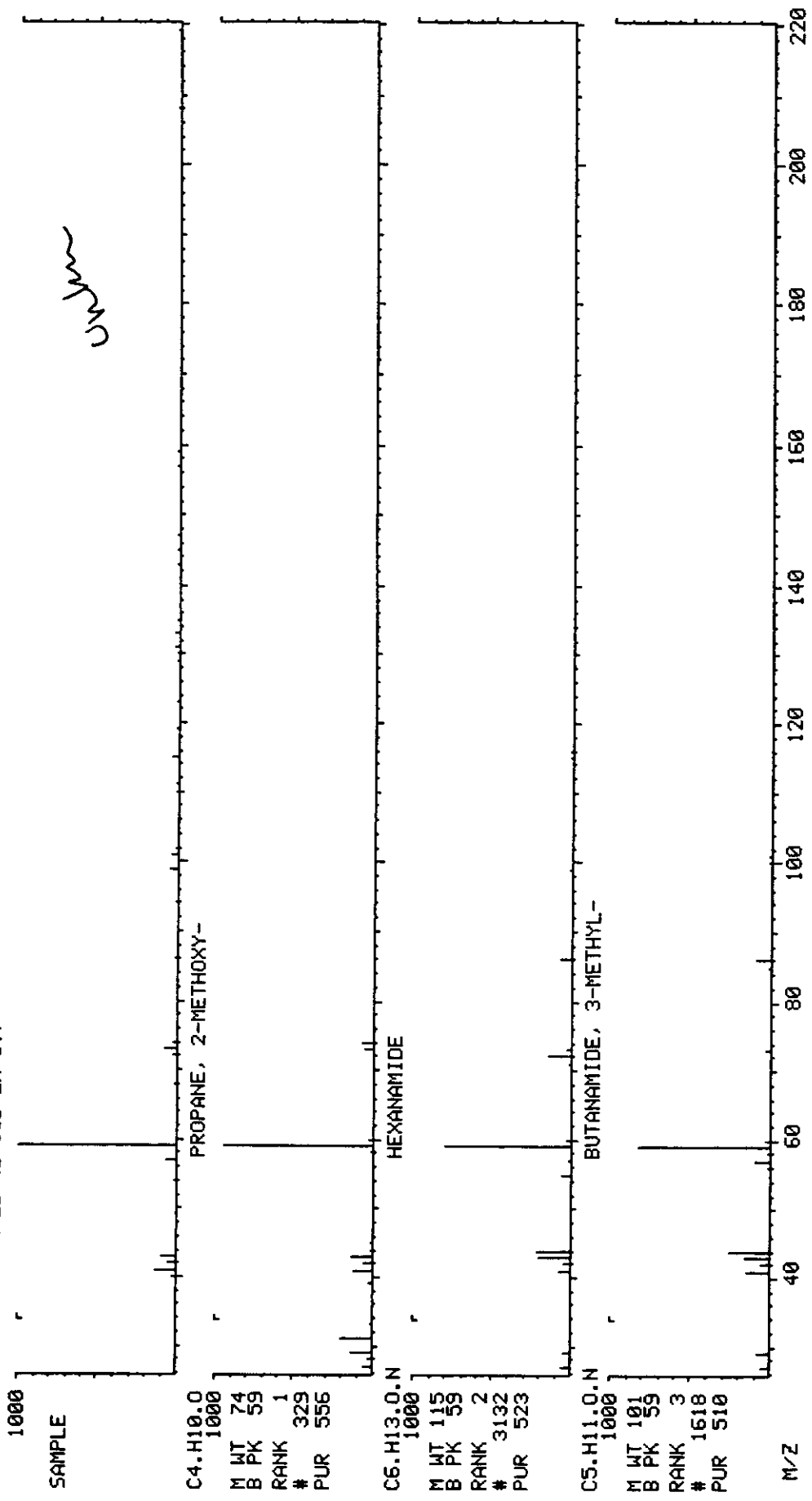
62231 spectra in LIBRARYNB searched for maximum PURITY  
 435 matched at least 5 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 329 PROPANE, 2-METHOXY-  
 2 3132 HEXANAMIDE  
 3 1618 BUTANAMIDE, 3-METHYL-  
 4 11721 NONANAMIDE  
 5 8432 2-HEXANONE, 3-HYDROXY-3,5-DIMETHYL-  
 6 1694 3-HYDROXY-3-METHYL-2-BUTANONE  
 7 1617 PENTANAMIDE  
 8 8871 METHANE, TERT-BUTOXYISOPROPOXY-  
 9 22656 DODECANAMIDE

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C4.H10.O	74	59	556	924	590
2	C6.H13.O.N	115	59	523	726	621
3	C5.H11.O.N	101	59	510	780	614
4	C9.H19.O.N	157	59	502	713	610
5	C8.H16.O2	144	59	499	818	574
6	C5.H10.O2	102	59	474	834	541
7	C5.H11.O.N	101	59	470	734	595
8	C8.H18.O2	146	57	467	690	527
9	C12.H25.O.N	199	59	463	640	633

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	598-53-8
2	---	---	---	---	628-02-4
3	---	---	---	---	541-46-8
4	---	---	---	---	1120-07-6
5	---	---	---	---	6321-14-8
6	---	---	---	---	115-22-0
7	---	---	---	---	626-97-1
8	---	---	---	---	4346-01-4
9	---	---	---	---	1120-16-7

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 20:24  
 SAMPLE: S-MM5-R8 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)  
 DATA: 30068109 #1834  
 CALI: 30068109 # 3  
 BASE M/Z: 59  
 RIC: 2840.



Library Search                    Data: 30068109 #1916            Base m/z: 43  
 08/31/98 22:58:00 + 21:19       Cali: 30068109 # 3            RIC: 29632.  
 Sample: S-MM5-RB 1/3SA/1ML       INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

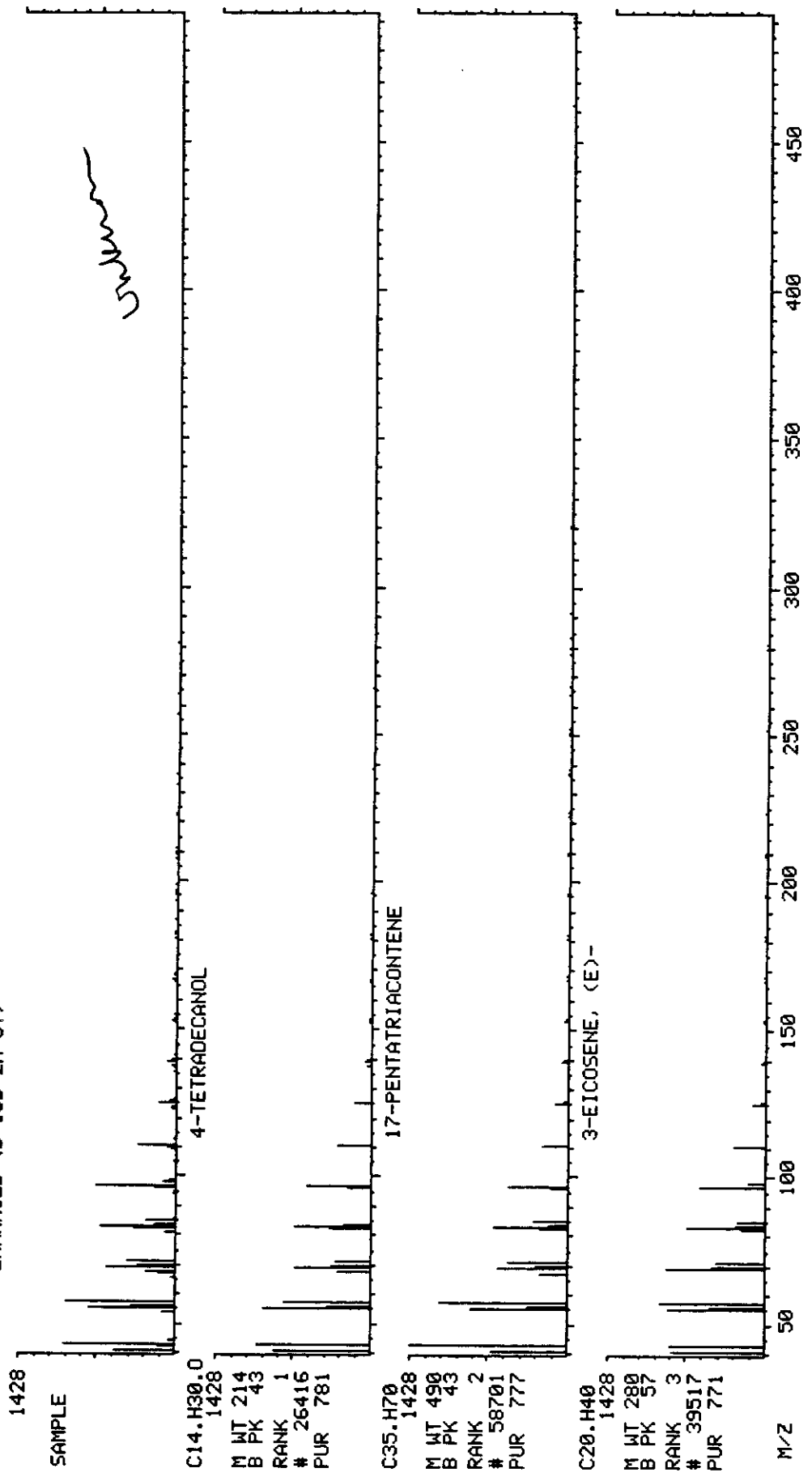
62231 spectra in LIBRARYNB searched for maximum PURITY  
 818 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 26416 4-TETRADECANOL  
 2 58701 17-PENTATRIACONTENE  
 3 39517 3-EICOSENE, (E)-  
 4 31653 1-HEPTADECENE  
 5 35206 1-HEPTADECANOL  
 6 40232 1-NONADECANOL  
 7 39516 5-EICOSENE, (E)-  
 8 39515 9-EICOSENE, (E)-  
 9 32420 1-HEXADECANOL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C14.H30.O	214	43	781	980	787
2	C35.H70	490	43	777	941	812
3	C20.H40	280	57	771	962	780
4	C17.H34	238	55	769	955	778
5	C17.H36.O	256	55	769	978	773
6	C19.H40.O	284	55	758	982	767
7	C20.H40	280	55	757	943	779
8	C20.H40	280	57	757	944	780
9	C16.H34.O	242	55	756	967	760

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	1653-33-4
2	---	---	---	---	6971-40-0
3	---	---	---	---	74685-33-9
4	---	---	---	---	6765-39-5
5	---	---	---	---	1454-85-9
6	---	---	---	---	1454-84-8
7	---	---	---	---	74685-30-6
8	---	---	---	---	74685-29-3
9	---	---	---	---	36653-82-4

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 21:19  
 SAMPLE: S-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068109 #1916  
 CALI: 30068109 # 3  
 BASE M/Z: 43  
 RIC: 29632.





Library Search                      Data: 30068109 #1934                      Base m/z: 59  
 08/31/98 22:58:00 + 21:31                      Cali: 30068109 # 3                      RIC: 11856.  
 Sample: S-MM5-RB 1/3SA/1ML                      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 265 matched at least 6 of the 16 largest peaks in the unknown

Rank In.                      Name  
 1 41712 3,6,9,12,15-PENTAOXANONADECAN-1-OL  
 2 36510 1,4,7,10,13,16-HEXAOXACYCLOOCTADECANE  
 3 1909 2-PROPANOL, 1-METHOXY-2-METHYL-  
 4 34224 2,5,8,11,14-PENTAOXAHEXADECAN-16-OL  
 5 9227 1,3,3-TRIMETHOXYBUTANE  
 6 5825 2-PROPANOL, 1-(2-METHYLPROPOXY)-  
 7 5832 2-PROPANOL, 1-(1-METHYLPROPOXY)-  
 8 3528 2-METHYL-2,3-PENTANEDIOL  
 9 8813 ETHENE, (2-ETHOXY-1-METHOXYETHOXY)-

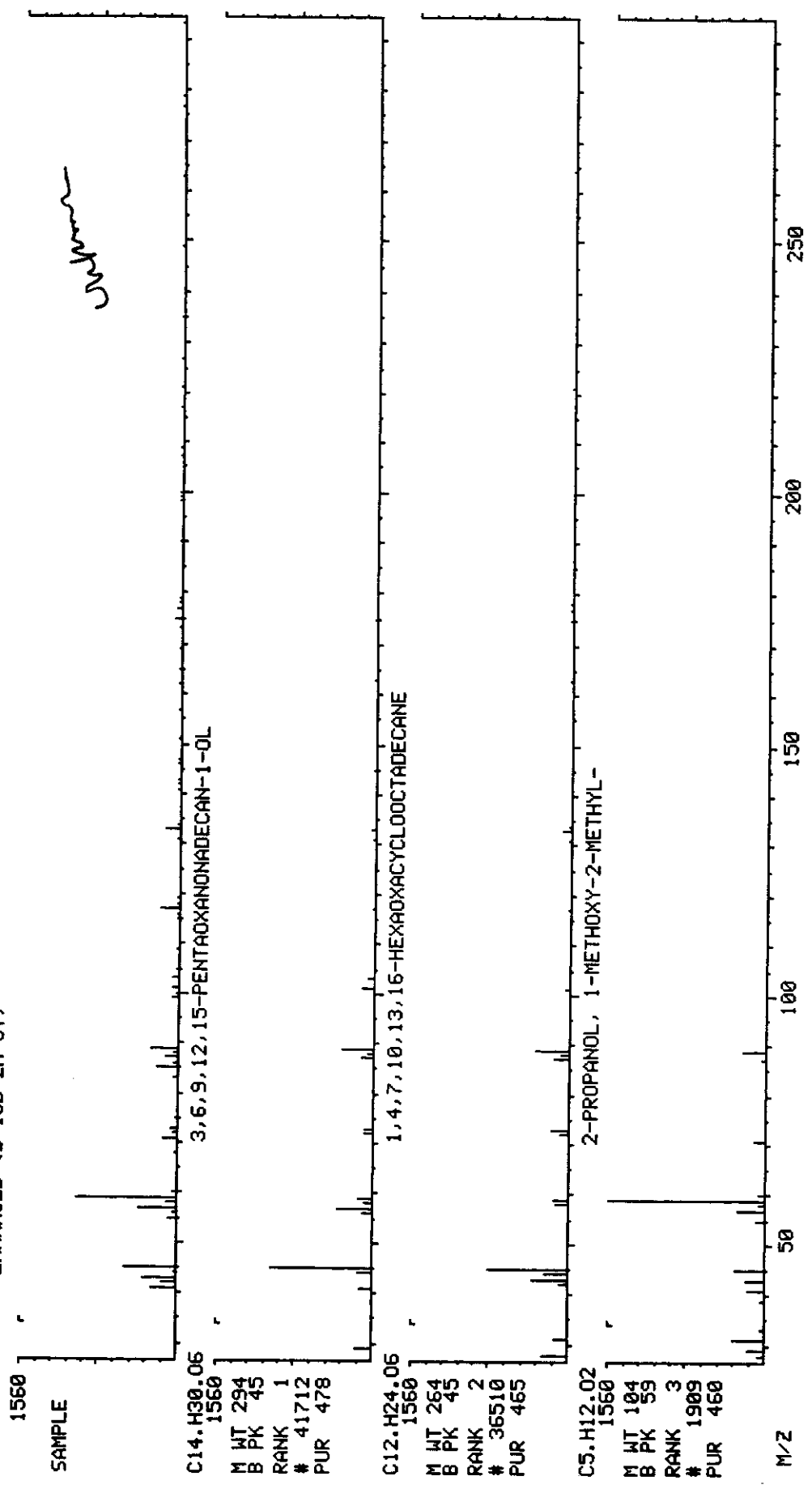
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C14.H30.O6	294	45	478	775	489
2	C12.H24.O6	264	45	465	776	466
3	C5.H12.O2	104	59	460	937	464
4	C11.H24.O6	252	45	447	872	464
5	C7.H16.O3	148	45	444	718	454
6	C7.H16.O2	132	57	443	823	457
7	C7.H16.O2	132	43	433	827	450
8	C6.H14.O2	118	59	426	846	443
9	C7.H14.O3	146	59	422	853	422

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	1786-94-3
2	---	---	---	---	17455-13-9
3	---	---	---	---	3587-64-2
4	---	---	---	---	23778-52-1
5	---	---	---	---	6607-66-5
6	---	---	---	---	23436-19-3
7	---	---	---	---	53907-95-2
8	---	---	---	---	7795-80-4
9	---	---	---	---	54063-18-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 21:31  
 SAMPLE: S-MMS-RB 1/3SA/1ML INST. ID: F15  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #1934  
 CALI: 30068109 # 3

BASE M/Z: 59  
 RIC: 11856.



1808

Library Search                    Data: 30068109 #1940            Base m/z: 59  
 08/31/98 22:58:00 + 21:35       Cali: 30068109 # 3            RIC: 8400.  
 Sample: S-MM5-RB 1/3SA/1ML       INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 400 matched at least 5 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 9227 1,3,3-TRIMETHOXYBUTANE  
 2 27502 15-CROWN-5  
 3 1909 2-PROPANOL, 1-METHOXY-2-METHYL-  
 4 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-  
 5 1919 1-BUTANOL, 3-METHOXY-  
 6 36510 1,4,7,10,13,16-HEXAOXACYCLOCTADECANE  
 7 41712 3,6,9,12,15-PENTAOXANONADECAN-1-OL  
 8 9235 1-PROPANOL, 2-(2-METHOXY-1-METHYLETHOXY)-  
 9 3528 2-METHYL-2,3-PENTANEDIOL

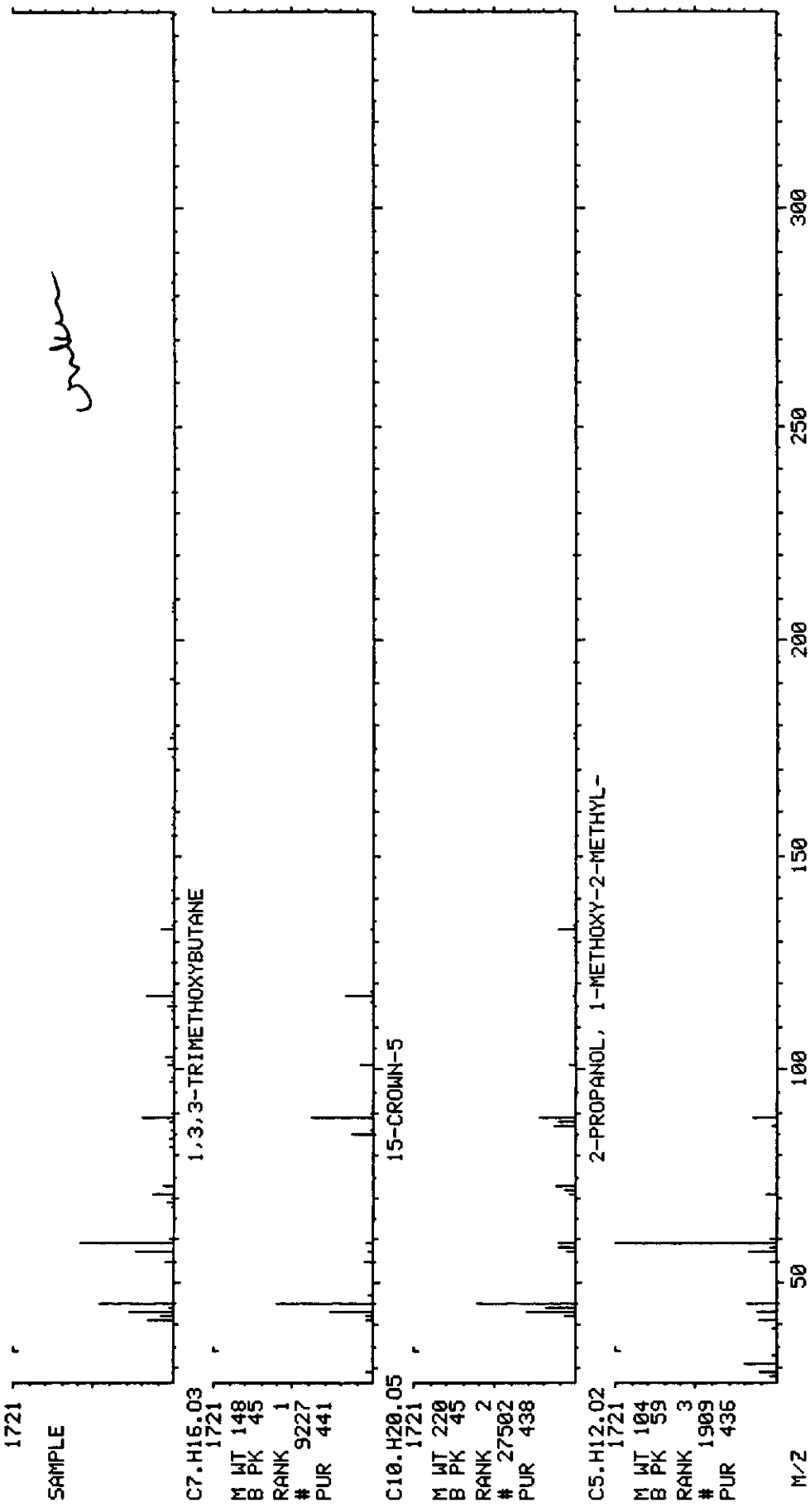
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H16.O3	148	45	441	695	488
2	C10.H20.O5	220	45	438	648	528
3	C5.H12.O2	104	59	436	882	457
4	C10.H22.O4	206	59	418	738	448
5	C5.H12.O2	104	59	418	811	431
6	C12.H24.O6	264	45	406	661	487
7	C14.H30.O6	294	45	404	648	496
8	C7.H16.O3	148	59	398	763	408
9	C6.H14.O2	118	59	394	793	432

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	6607-66-5
2	---	---	---	---	33100-27-5
3	---	---	---	---	3587-64-2
4	---	---	---	---	20324-33-8
5	---	---	---	---	2517-43-3
6	---	---	---	---	17455-13-9
7	---	---	---	---	1786-94-3
8	---	---	---	---	55956-21-3
9	---	---	---	---	7795-80-4

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 21:35  
 SAMPLE: S-MM5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #1940  
 CALI: 30068109 # 3

BASE M/Z: 59  
 RIC: 8400.



100  
 100  
 100

Library Search                    Data: 30068109 #1966            Base m/z: 59  
 08/31/98 22:58:00 + 21:52       Cali: 30068109 # 3            RIC: 3004.  
 Sample: S-MM5-RB 1/3SA/1ML       INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 783 matched at least 5 of the 16 largest peaks in the unknown

- Rank In.       Name  
 1 19528 3-DODECANOL  
 2 12059 4-ETHYL-3-OCTANOL  
 3 12057 3-DECANOL  
 4 3237 4-HYDROXY-3-HEXANONE  
 5 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-  
 6 12062 3-OCTANOL, 6-ETHYL-  
 7 15851 2-METHYL-2-DECANOL  
 8 15839 5-ETHYL-3-NONANOL  
 9 8506 4,5-DIMETHYL-3-HEPTANOL

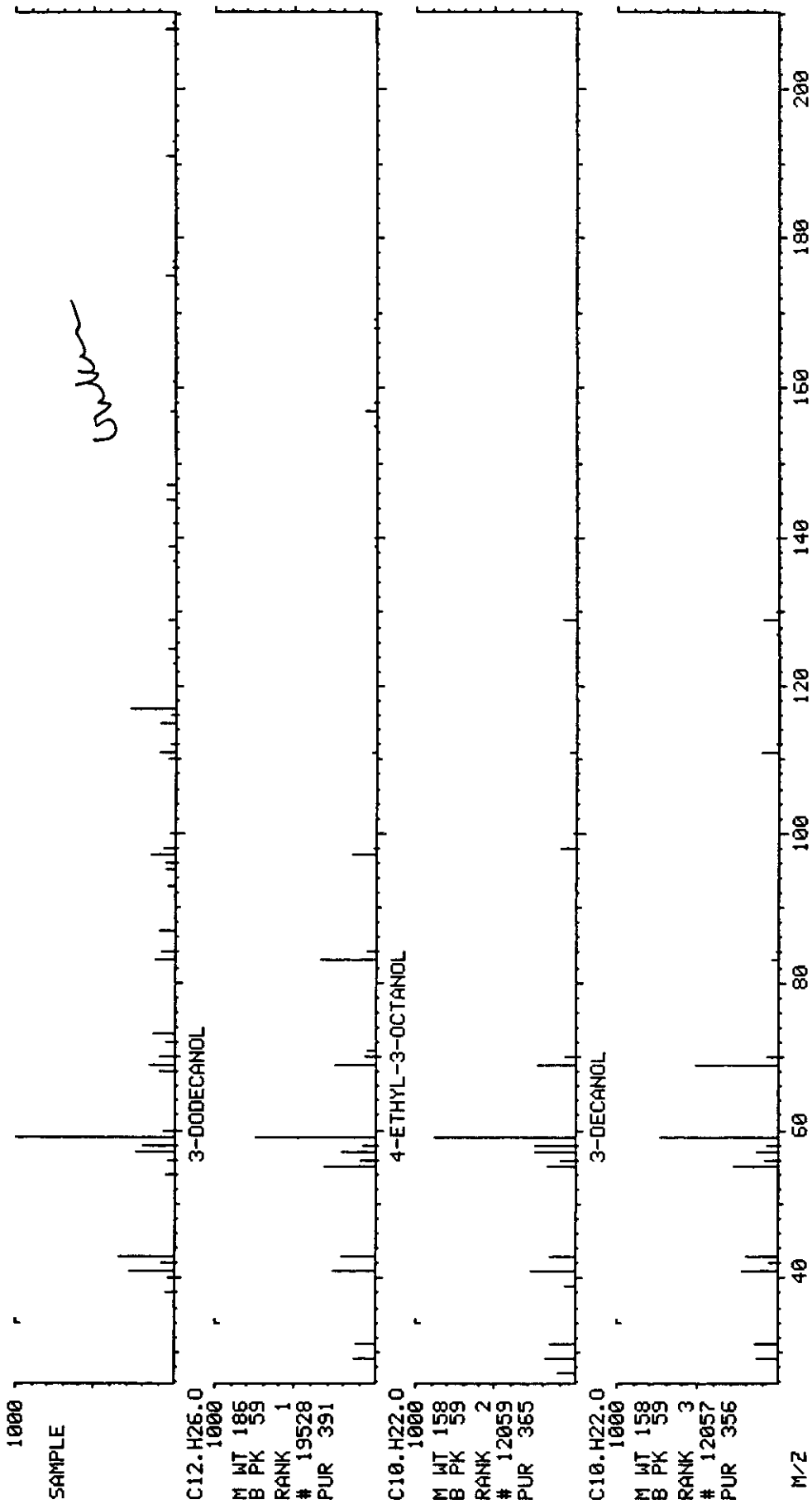
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C12.H26.O	186	59	391	789	450
2	C10.H22.O	158	59	365	887	397
3	C10.H22.O	158	59	356	792	401
4	C6.H12.O2	116	59	355	770	366
5	C7.H16.O3	148	59	348	792	380
6	C10.H22.O	158	59	341	755	401
7	C11.H24.O	172	59	339	731	400
8	C11.H24.O	172	57	338	719	398
9	C9.H20.O	144	59	338	824	403

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	10203-30-2
2	---	---	---	---	19781-26-1
3	---	---	---	---	1565-81-7
4	---	---	---	---	4984-85-4
5	---	---	---	---	13588-28-8
6	---	---	---	---	19781-27-2
7	---	---	---	---	3396-02-9
8	---	---	---	---	- -
9	---	---	---	---	- -

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 21:52  
 SAMPLE: S-MN5-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA >)/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #1966  
 CALI: 30068109 # 3

BASE M/2: 59  
 RIC: 3004.



Library Search                    Data: 30068109 #2014            Base m/z: 43  
 08/31/98 22:58:00 + 22:24       Cali: 30068109 # 3            RIC: 14592.  
 Sample: S-MM5-RB 1/3SA/1ML       INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 950 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 39515 9-EICOSENE, (E)-  
 2 39516 5-EICOSENE, (E)-  
 3 34410 3-OCTADECENE, (E)-  
 4 37062 1-NONADECENE  
 5 34418 9-OCTADECENE, (E)-  
 6 34411 5-OCTADECENE, (E)-  
 7 31653 1-HEPTADECENE  
 8 39517 3-EICOSENE, (E)-  
 9 26416 4-TETRADECANOL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C20.H40	280	55	729	949	741
2	C20.H40	280	55	728	947	741
3	C18.H36	252	69	718	938	734
4	C19.H38	266	97	717	948	737
5	C18.H36	252	55	716	935	730
6	C18.H36	252	55	716	934	730
7	C17.H34	238	55	715	942	728
8	C20.H40	280	57	713	951	725
9	C14.H30.O	214	43	711	970	721

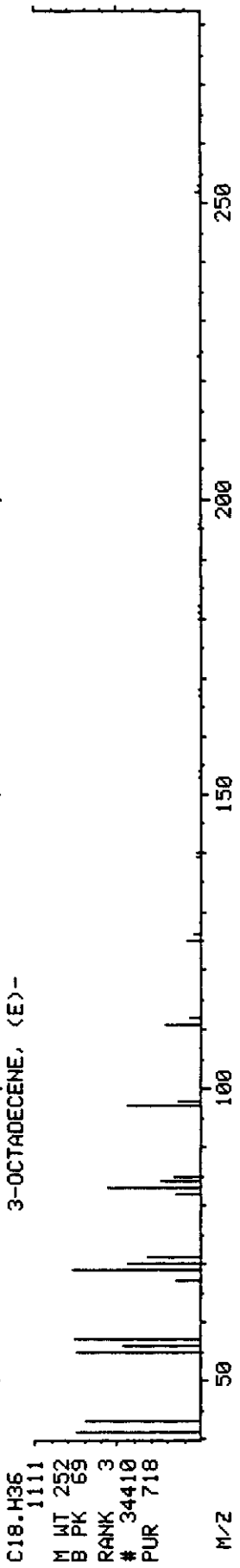
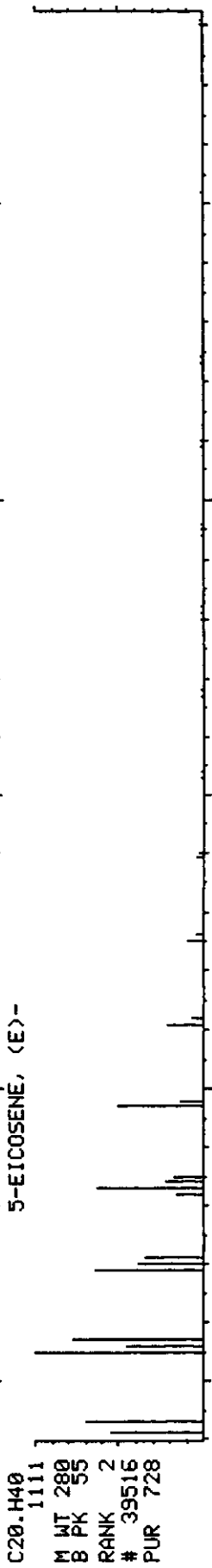
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	74685-29-3
2	---	---	---	---	74685-30-6
3	---	---	---	---	7206-19-1
4	---	---	---	---	18435-45-5
5	---	---	---	---	7206-25-9
6	---	---	---	---	7206-21-5
7	---	---	---	---	6765-39-5
8	---	---	---	---	74685-33-9
9	---	---	---	---	1653-33-4

BASE M/Z: 43  
RIC: 14592.

DATA: 30068109 #2014  
CALI: 30068109 # 3

MID LIBRARY SEARCH <LIBRARYNB>  
08/31/98 22:58:00 + 22:24  
SAMPLE: S-MMS-RB 1/35A/1ML INST. ID: F16  
CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M  
ENHANCED (S 15B 2N 0T)

~~000000~~  
29119159



1111  
SAMPLE

C20.H40  
1111  
M WT 280  
B PK 55  
RANK 1  
# 39515  
PUR 729

C20.H40  
1111  
M WT 280  
B PK 55  
RANK 2  
# 39516  
PUR 728

C18.H36  
1111  
M WT 252  
B PK 69  
RANK 3  
# 34410  
PUR 718

M/Z



Library Search                      Data: 30068109 #2051                      Base m/z: 59  
 08/31/98 22:58:00 + 22:49                      Cali: 30068109 # 3                      RIC: 5984.  
 Sample: S-MM5-RB 1/3SA/1ML                      INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYN8 searched for maximum PURITY  
 363 matched at least 6 of the 16 largest peaks in the unknown

Rank In.                      Name  
 1 1909 2-PROPANOL, 1-METHOXY-2-METHYL-  
 2 27502 15-CROWN-5  
 3 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-  
 4 9227 1,3,3-TRIMETHOXYBUTANE  
 5 8822 BUTANOIC ACID, 4-ETHOXY-, METHYL ESTER  
 6 8871 METHANE, TERT-BUTOXYISOPROPOXY-  
 7 36510 1,4,7,10,13,16-HEXAOXACYCLOOCTADECANE  
 8 12385 BUTYRIC ACID, 4-ISOPROPOXY-, METHYL ESTER  
 9 5826 2-PROPANOL, 1-ISOPROPOXY-2-METHYL-

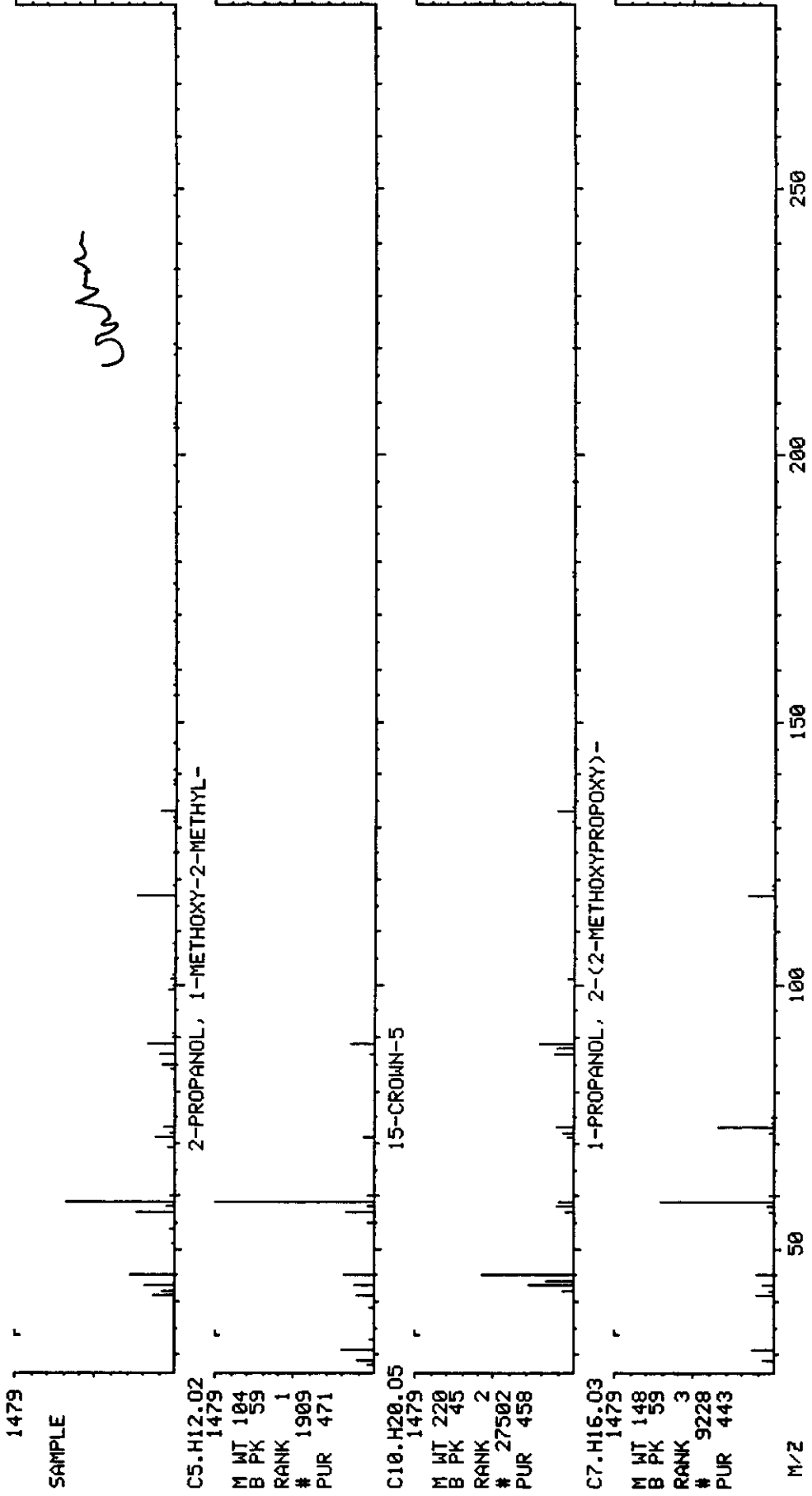
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C5.H12.O2	104	59	471	922	486
2	C10.H20.O5	220	45	458	634	535
3	C7.H16.O3	148	59	443	798	461
4	C7.H16.O3	148	45	430	661	457
5	C7.H14.O3	146	59	425	719	522
6	C8.H18.O2	146	57	419	791	435
7	C12.H24.O6	264	45	413	654	477
8	C8.H16.O3	160	101	413	694	459
9	C7.H16.O2	132	59	412	641	484

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	3587-64-2
2	---	---	---	---	33100-27-5
3	---	---	---	---	13588-28-8
4	---	---	---	---	6607-66-5
5	---	---	---	---	29006-04-0
6	---	---	---	---	4346-01-4
7	---	---	---	---	17455-13-9
8	---	---	---	---	29006-05-1
9	---	---	---	---	3587-75-5

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 22:49  
 SAMPLE: S-MMS-RB 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA >)/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #2051  
 CALI: 30068109 # 3

BASE M/Z: 59  
 RIC: 5984.



1479  
 SAMPLE

C5.H12.02  
 1479  
 M WT 104  
 B PK 59  
 RANK 1  
 # 1909  
 PUR 471

C10.H20.05  
 1479  
 M WT 220  
 B PK 45  
 RANK 2  
 # 27502  
 PUR 458

C7.H16.03  
 1479  
 M WT 148  
 B PK 59  
 RANK 3  
 # 9228  
 PUR 443

M/Z

Library Search                      Data: 30068109 #2057                      Base m/z: 59  
 08/31/98 22:58:00 + 22:53        Cali: 30068109 # 3                      RIC: 7840.  
 Sample: S-MMS-RB 1/3SA/1ML        INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 323 matched at least 6 of the 16 largest peaks in the unknown

- Rank In.                      Name  
 1 9227 1,3,3-TRIMETHOXYBUTANE  
 2 41712 3,6,9,12,15-PENTAOXANONADECAN-1-OL  
 3 5809 1-ETHOXPENTAN-3-OL  
 4 3528 2-METHYL-2,3-PENTANEDIOL  
 5 1909 2-PROPANOL, 1-METHOXY-2-METHYL-  
 6 16192 2-PROPANOL, 1- [1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-  
 7 3000 OXIRANE, 2,2-DIMETHYL-3-PROPYL-  
 8 34224 2,5,8,11,14-PENTAOXAHEXADECAN-16-OL  
 9 16199 BUTYRIC ACID, 4-BUTOXY-, METHYL ESTER

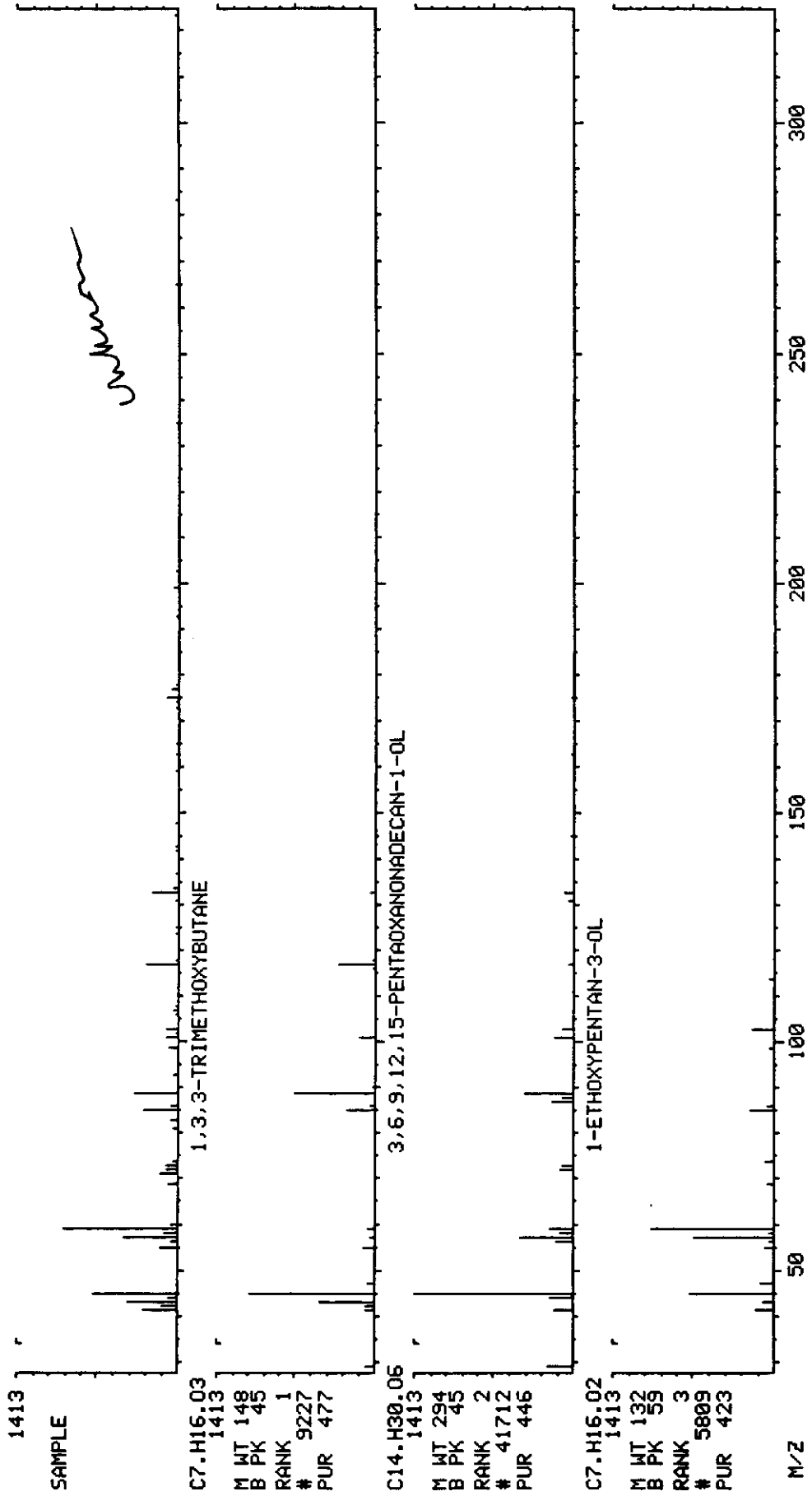
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H16.O3	148	45	477	749	491
2	C14.H30.O6	294	45	446	723	499
3	C7.H16.O2	132	59	423	903	442
4	C6.H14.O2	118	59	407	812	422
5	C5.H12.O2	104	59	404	903	416
6	C9.H18.O3	174	59	396	715	410
7	C7.H14.O	114	43	388	917	395
8	C11.H24.O6	252	45	383	798	432
9	C9.H18.O3	174	57	376	719	435

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	6607-66-5
2	---	---	---	---	1786-94-3
3	---	---	---	---	-
4	---	---	---	---	7795-80-4
5	---	---	---	---	3587-64-2
6	---	---	---	---	55956-25-7
7	---	---	---	---	17612-35-0
8	---	---	---	---	23778-52-1
9	---	---	---	---	29006-06-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 22:58:00 + 22:53  
 SAMPLE: S-MMS-RB 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068109 #2057  
 CALI: 30068109 # 3

BASE M/Z: 59  
 RIC: 7840.



TIC SELECTION REPORT

DATA FILE: 30068109

THE FOLLOWING PEAKS WERE REJECTED BECAUSE  
AT LEAST 40 % OF THEIR SIZE WAS ACCOUNTED FOR BY  
TARGET COMPOUNDS ELUTING WITHIN 4 SCANS OF THE  
PEAK TOP.

SCAN	SIZE	AMOUNT
198	273224.	38.262
308	328440.	45.994
340	229200.	32.097
421	136284.	19.085
529	285638.	40.000
722	317824.	37.173
827	341992.	40.000
968	294902.	35.699
1085	330432.	40.000
1383	308168.	58.652
1383	308168.	58.652
1551	210168.	40.000
1782	186560.	40.000

TOTAL NUMBER OF UNIDENTIFIED PEAKS WITH SIZE  
GREATER THAN 10 % OF THE CLOSEST INTERNAL STANDARD  
THAT DOES NOT HAVE INTERFERENCES = 25

INTERNAL STANDARDS THAT HAVE RIC SIZE LESS THAN  
50 % OR GREATER THAN 200 % OF THE ESTIMATED RIC SIZE  
ARE CONSIDERED TO HAVE INTERFERENCES AND WILL NOT BE USED  
FOR QUANTITATION.

#	INTERNAL STANDARD	RIC SIZE	PERCENT OF ESTIMATED	
			RIC SIZE	SCAN
1	C140 NAPHTHALENE-D8	285638.	130	529
2	C150 ACENAPHTHENE-D10	341992.	127	827
3	C160 PHENANTHRENE-D10	330432.	118	1085
4	C170 CHRYSENE-D12	210168.	120	1551
5	C175 PERYLENE-D12	186560.	122	1782

\* INDICATES INTERFERENCE

SIZE = AREA

Semivolatile Organics  
Method 0010/8270

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-4-F, FH, XAD, COND, BH  
 LAB ID: 300681-0010-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98

Sampled: 25 JUL 98  
 Prepared: 31 JUL 98

Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
Phenol	ND	ug/Sample	3000	GR
bis(2-Chloroethyl)ether	ND	ug/Sample	3000	
2-Chlorophenol	ND	ug/Sample	3000	
1,3-Dichlorobenzene	ND	ug/Sample	3000	
1,4-Dichlorobenzene	ND	ug/Sample	3000	
Benzyl alcohol	ND	ug/Sample	3000	
1,2-Dichlorobenzene	ND	ug/Sample	3000	
2-Methylphenol	ND	ug/Sample	3000	
2,2'-Oxybis(1-chloropropane)	ND	ug/Sample	3000	
3/4-Methylphenol	ND	ug/Sample	3000	
N-Nitroso-di-n-propylamine	ND	ug/Sample	3000	
Hexachloroethane	ND	ug/Sample	3000	
Nitrobenzene	ND	ug/Sample	3000	
Isophorone	ND	ug/Sample	3000	
2-Nitrophenol	ND	ug/Sample	3000	
2,4-Dimethylphenol	ND	ug/Sample	3000	
Benzoic acid	ND	ug/Sample	15000	
bis(2-Chloroethoxy)-methane	ND	ug/Sample	3000	
2,4-Dichlorophenol	ND	ug/Sample	3000	
1,2,4-Trichlorobenzene	ND	ug/Sample	3000	
Naphthalene	1500	ug/Sample	3000	J
4-Chloroaniline	ND	ug/Sample	3000	
Hexachlorobutadiene	ND	ug/Sample	3000	
4-Chloro-3-methylphenol	ND	ug/Sample	3000	
2-Methylnaphthalene	2300	ug/Sample	3000	J
Hexachlorocyclopentadiene	ND	ug/Sample	3000	
2,4,6-Trichlorophenol	ND	ug/Sample	3000	
2,4,5-Trichlorophenol	ND	ug/Sample	15000	
2-Chloronaphthalene	ND	ug/Sample	3000	
2-Nitroaniline	ND	ug/Sample	3000	
Dimethyl phthalate	ND	ug/Sample	3000	
Acenaphthylene	ND	ug/Sample	3000	
3-Nitroaniline	ND	ug/Sample	15000	
Acenaphthene	ND	ug/Sample	3000	
2,4-Dinitrophenol	ND	ug/Sample	15000	

Note G = Reporting limit(s) raised due to matrix interference.

Note J = Result is detected below the reporting limit or is an estimated concentration.

Note R = Reporting limit(s) raised due to sample volume limitations.

ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.

Rev 230787

Semivolatile Organics  
Method 0010/8270

(cont.)

Client Name: Pacific Environmental Services  
Client ID: S-MM5-4-F,FH,XAD,COND,BH  
LAB ID: 300681-0010-SA  
Matrix: AIRTRAIN  
Authorized: 30 JUL 98

Sampled: 25 JUL 98  
Prepared: 31 JUL 98

Received: 30 JUL 98  
Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
4-Nitrophenol	ND	ug/Sample	15000	
Dibenzofuran	ND	ug/Sample	3000	
2,4-Dinitrotoluene	ND	ug/Sample	3000	
2,6-Dinitrotoluene	ND	ug/Sample	3000	
Diethyl phthalate	ND	ug/Sample	3000	
4-Chlorophenyl phenyl ether	ND	ug/Sample	3000	
Fluorene	ND	ug/Sample	3000	
4-Nitroaniline	ND	ug/Sample	15000	
4,6-Dinitro-2-methylphenol	ND	ug/Sample	15000	
N-Nitrosodiphenylamine	ND	ug/Sample	3000	
4-Bromophenyl phenyl ether	ND	ug/Sample	3000	
Hexachlorobenzene	ND	ug/Sample	3000	
Pentachlorophenol	ND	ug/Sample	15000	
Phenanthrene	590	ug/Sample	3000	J
Anthracene	ND	ug/Sample	3000	
Di-n-butyl phthalate	ND	ug/Sample	3000	
Fluoranthene	ND	ug/Sample	3000	
Pyrene	ND	ug/Sample	3000	
Butyl benzyl phthalate	ND	ug/Sample	3000	
3,3'-Dichlorobenzidine	ND	ug/Sample	6000	
Benzo(a)anthracene	ND	ug/Sample	3000	
bis(2-Ethylhexyl)-phthalate	ND	ug/Sample	3000	
Chrysene	ND	ug/Sample	3000	
Di-n-octyl phthalate	ND	ug/Sample	3000	
Benzo(b)fluoranthene	ND	ug/Sample	3000	
Benzo(k)fluoranthene	ND	ug/Sample	3000	
Benzo(a)pyrene	ND	ug/Sample	3000	
Indeno(1,2,3-cd)pyrene	ND	ug/Sample	3000	
Dibenz(a,h)anthracene	ND	ug/Sample	3000	
Benzo(g,h,i)perylene	ND	ug/Sample	3000	
Acetophenone	ND	ug/Sample	3000	
4-Aminobiphenyl	ND	ug/Sample	15000	
Aniline	ND	ug/Sample	3000	
Benzidine	ND	ug/Sample	30000	
3,3'-Dimethylbenzidine	ND	ug/Sample	6000	
N-Nitrosodimethylamine	ND	ug/Sample	3000	
N-Nitrosomorpholine	ND	ug/Sample	3000	

Note J = Result is detected below the reporting limit or is an estimated concentration.  
ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.  
Rev 230787

1315

Semivolatile Organics  
Method 0010/8270

(cont.)

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-4-F, FH, XAD, COND, BH  
 LAB ID: 300681-0010-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98  
 Sampled: 25 JUL 98  
 Prepared: 31 JUL 98  
 Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
Pentachloronitrobenzene (PCNB)	ND	ug/Sample	15000	
o-Toluidine	ND	ug/Sample	6000	
2-Methoxybenzenamine	ND	ug/Sample	--	
Biphenyl	ND	ug/Sample	--	
Chloroacetophenone	ND	ug/Sample	--	
Cumene	ND	ug/Sample	--	
DBCP (1,2-Dibromo-3-chloropropane)	ND	ug/Sample	--	
Benzo(e)pyrene	ND	ug/Sample	--	
N-N-Diethylaniline	ND	ug/Sample	--	
Dimethylaniline	ND	ug/Sample	--	
3,3'-Dimethoxybenzidine	ND	ug/Sample	--	
Hydroquinone	ND	ug/Sample	--	
4,4'-Methyl-bis(2-chloroaniline)	ND	ug/Sample	--	
4-Nitrodiphenyl	ND	ug/Sample	--	
Trifluralin	ND	ug/Sample	--	

Surrogate	Recovery	Acceptable Range	
Nitrobenzene-d5	ND %	45 - 107	H
2-Fluorobiphenyl	ND %	62 - 110	
Terphenyl-d14	ND %	58 - 135	
Phenol-d5	ND %	43 - 130	
2-Fluorophenol	ND %	36 - 111	
2,4,6-Tribromophenol	ND %	58 - 131	

Note H = Spiked analyte not detected because of required sample dilution.  
 ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.  
 Rev 230787

1820



Semivolatiles Library Search (20 Compound TID)  
Method 8270

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-4-F, FH, XAD, COND, BH  
 LAB ID: 300681-0010-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98

Sampled: 25 JUL 98  
 Prepared: NA

Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Result	Units	Reporting Limit	Qualifier
Undecane	7600	ug/Sample	--	0
Unknown	7500	ug/Sample	--	
Unknown alkane	7100	ug/Sample	--	
Decane, 2,5,9-Trimethyl-	6200	ug/Sample	--	0
Unknown	12000	ug/Sample	--	
Unknown alkane	12000	ug/Sample	--	
Unknown hydrocarbon	6900	ug/Sample	--	
Octane, 3,5-dimethyl-	9800	ug/Sample	--	0
Unknown	6400	ug/Sample	--	
Heptadecane, 2,6-dimethyl-	13000	ug/Sample	--	0
Tetradecane	18000	ug/Sample	--	0
Unknown	8600	ug/Sample	--	
Heptadecane, 2,6,10,14 -tetramethyl-	12000	ug/Sample	--	0
Pentadecane	14000	ug/Sample	--	0
Unknown	6400	ug/Sample	--	
Nonadecane	16000	ug/Sample	--	
Undecane, 2,6-dimethyl-	13000	ug/Sample	--	0
Heptadecane, 2,6-dimethyl-	28000	ug/Sample	--	0
Unknown alkane	7600	ug/Sample	--	
Heptadecane, 2,6-dimethyl-	9800	ug/Sample	--	0

Note 0 = Or structurally similar compound (isomer).  
 NA = Not Applicable

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.  
 Rev 230787

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068110  
Std Id: ST16980831

Sample: S-MM5-4 1/3SA/100M ✓ INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:28  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.0	BDL	0* ↑	45 107
CS25 2-FLUOROBIPHENY	50.0	BDL	0*	62 110
CS30 TERPHENYL-D14	50.00	122.0	244. ↓	* 58 135
CS45 PHENOL-D5	100.	BDL	0*	43 130
CS50 2-FLUOROPHENOL	100.0	116.0	116. ↓	* 36 111
CS55 2,4,6-TRIBROMOP	100.0	127.0	127. ↓	58 131

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
HEXACHLOROENZENE-C13		ND	UG/A	3000.0
C310 N-NITROSODIMETHYLAMINE		ND	UG/A	3000.0
PYRIDINE		ND	UG/A	6000.0
2-PICOLINE		ND	UG/A	3000.0
N-NITROSOMETHYLETHYLAMINE		ND	UG/A	3000.0
METHYLMETHANESULFONATE		ND	UG/A	3000.0
N-NITROSODIETHYLAMINE		ND	UG/A	3000.0
ETHYLMETHANE SULFONATE		ND	UG/A	3000.0
PENTACHLOROETHANE		ND	UG/A	3000.0
C320 ANILINE		ND	UG/A	3000.0
C315 PHENOL		ND	UG/A	3000.0
C325 BIS(2-CHLOROETHYL)ETHER		ND	UG/A	3000.0
C330 2-CHLOROPHENOL		ND	UG/A	3000.0
C335 1,3-DICHLOROBENZENE		ND	UG/A	3000.0
C340 1,4-DICHLOROBENZENE		ND	UG/A	3000.0
C345 BENZYL ALCOHOL		ND	UG/A	3000.0
C350 1,2-DICHLOROBENZENE		ND	UG/A	3000.0
C355 2-METHYLPHENOL		ND	UG/A	3000.0
C360 2,2'-OXYBIS(1-CLPROPAN)		ND	UG/A	3000.0
C361 ACETOPHENONE		ND	UG/A	3000.0
N-NITROSPYRROLIDINE		ND	UG/A	3000.0
N-NITROSOMORPHOLINE		ND	UG/A	3000.0
3-METHYL PHENOL		ND	UG/A	3000.0
C365 4-METHYLPHENOL		ND	UG/A	3000.0
C370 N-NITROSO-DI-N-PROPYLAM		ND	UG/A	3000.0
O-TOLUIDINE		ND	UG/A	3000.0
C375 HEXACHLOROETHANE		ND	UG/A	3000.0
C410 NITROBENZENE		ND	UG/A	3000.0
N-NITROSOPIPERIDINE		ND	UG/A	3000.0

Reviewed by:

*[Handwritten Signature]* 9.18.98  
DGL 10/6/98

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068110  
Std Id: ST16980831

Sample: S-MM5-4 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:28  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C415 ISOPHORONE		ND	UG/A	3000.0
C420 2-NITROPHENOL		ND	UG/A	3000.0
C425 2,4-DIMETHYLPHENOL		ND	UG/A	3000.0
C435 BIS(2-CHLOROETHOXY)METH		ND	UG/A	3000.0
C440 2,4-DICHLOROPHENOL		ND	UG/A	3000.0
C430 BENZOIC ACID		ND	UG/A	15000.0
AA-DIMETHYLPHENETHYLAMINE		ND	UG/A	3000.0
C445 1,2,4-TRICHLOROBENZENE		ND	UG/A	3000.0
C450 NAPHTHALENE		ND 1500	UG/A	3000.0
C455 4-CHLOROANALINE		ND	UG/A	3000.0
2,6-DICHLOROPHENOL		ND	UG/A	3000.0
HEXACHLOROPROPENE		ND	UG/A	3000.0
C460 HEXACHLOROBUTADIENE		ND	UG/A	3000.0
P-PHENYLENE DIAMINE		ND	UG/A	3000.0
N-NITROSODI-N-BUTYLAMINE		ND	UG/A	3000.0
C465 4-CHLORO-3-METHYLPHENO		ND	UG/A	3000.0
SAFROLE		ND	UG/A	3000.0
C470 2-METHYLNAPHTHALENE		ND 2300	UG/A	3000.0
1,2,4,5-TETRACHLOROBENZENE		ND	UG/A	3000.0
ISOSAFROLE (#1)		ND	UG/A	6000.0
C510 HEXACHLOROCYCLOPENTADI		ND	UG/A	3000.0
C515 2,4,6-TRICHLOROPHENOL		ND	UG/A	3000.0
C520 2,4,5-TRICHLOROPHENOL		ND	UG/A	3000.0
ISOSAFROLE (#2)		ND	UG/A	6000.0
C525 2-CHLORONAPHTHALENE		ND	UG/A	3000.0
1-CHLORONAPHTHALENE		ND	UG/A	1500.0
C530 2-NITROANILINE		ND	UG/A	15000.0
1,4-NAPHTHOQUINONE		ND	UG/A	3000.0
C535 DIMETHYLPHTHALATE		ND	UG/A	3000.0
1,3-DINITROBENZENE		ND	UG/A	3000.0
C540 ACENAPHTHYLENE		ND	UG/A	3000.0
C543 2,6-DINITROTOLUENE		ND	UG/A	3000.0
C545 3-NITROANILINE		ND	UG/A	15000.0
C550 ACENAPHTHENE		ND	UG/A	3000.0
C555 2,4-DINITROPHENOL		ND	UG/A	15000.0
C565 DIBENZOFURAN		ND	UG/A	3000.0
C560 4-NITROPHENOL		ND	UG/A	15000.0
PENTACHLOROBENZENE		ND	UG/A	3000.0
C570 2,4-DINITROTOLUENE		ND	UG/A	3000.0
1-NAPHTHYLAMINE		ND	UG/A	3000.0
2-NAPHTHYLAMINE		ND	UG/A	3000.0
2,3,4,6-TETRACHLOROPHENOL		ND	UG/A	6000.0
C580 DIETHYLPHTHALATE		ND	UG/A	3000.0
C590 FLUORENE		ND	UG/A	3000.0

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QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068110  
Std Id: ST16980831

Sample: S-MM5-4 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:28  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C585 4-CHLOROPHENYL-PHENYLE		ND	UG/A	3000.0
5-NITRO-O-TOLUIDINE		ND	UG/A	3000.0
C595 4-NITROANALINE		ND	UG/A	15000.0
C610 4,6-DINITRO-2-METHYLPH		ND	UG/A	15000.0
C615 N-NITROSODIPHENYLAMINE		ND	UG/A	3000.0
C620 AZOBENZENE		ND	UG/A	3000.0
SYM-TRINITROBENZENE		ND	UG/A	3000.0
C625 4-BROMOPHENYL-PHENYLET		ND	UG/A	3000.0
PHENACETIN		ND	UG/A	3000.0
DIALLATE	234	ND	UG/A	3000.0
C630 HEXACHLOROBENZENE		ND	UG/A	3000.0
4-AMINOBIPHENYL		ND	UG/A	3000.0
C635 PENTACHLOROPHENOL		ND	UG/A	15000.0
PRONAMIDE		ND	UG/A	3000.0
PENTACHLORONITROBENZENE		ND	UG/A	15000.0
C640 PHENANTHRENE		ND	UG/A	3000.0
C645 ANTHRACENE		ND	UG/A	3000.0
2SECBUTYL-4,6-DINITROPHENOL		ND	UG/A	3000.0
C647 CARBAZOLE		ND	UG/A	3000.0
C650 DI-N-BUTYLPHTHALATE		ND	UG/A	3000.0
4-NITROQUINOLINE-1-OXIDE		ND	UG/A	3000.0
METHAPYRILENE		ND	UG/A	3000.0
ISODRIN		ND	UG/A	3000.0
C655 FLUORANTHENE		ND	UG/A	3000.0
CHLOROBENZILATE		ND	UG/A	3000.0
C710 BENZIDINE		ND	UG/A	30000.0
C715 PYRENE		ND	UG/A	3000.0
ARAMITE (#1)		ND	UG/A	3000.0
ARAMITE (#2)		ND	UG/A	3000.0
P-DIMETHYLAMINOAZOBENZENE		ND	UG/A	3000.0
3,3'-DIMETHYLBENZIDINE		ND	UG/A	3000.0
KEPONE		ND	UG/A	15000.0
C720 BUTYLBENZYLPHTHALATE		ND	UG/A	3000.0
2-ACETYLAMINOFLUORENE		ND	UG/A	3000.0
C730 BENZO(A)ANTHRACENE		ND	UG/A	3000.0
C725 3,3'-DICHLOROBENZIDINE		ND	UG/A	6000.0
C740 CHRYSENE		ND	UG/A	3000.0
C745 BIS(2-ETHYLHEXYL)PHTHA		ND	UG/A	3000.0
3-METHYLCHOLANTHRENE		ND	UG/A	3000.0
C760 DI-N-OCTYL PHTHALATE		ND	UG/A	3000.0
C765 BENZO(B)FLUORANTHENE		ND	UG/A	3000.0
7,12-DIMETHYLBENZANTHRACENE		ND	UG/A	3000.0
C770 BENZO(K)FLUORANTHENE		ND	UG/A	3000.0
HEXACHLOROPHENE		ND	UG/A	3000.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

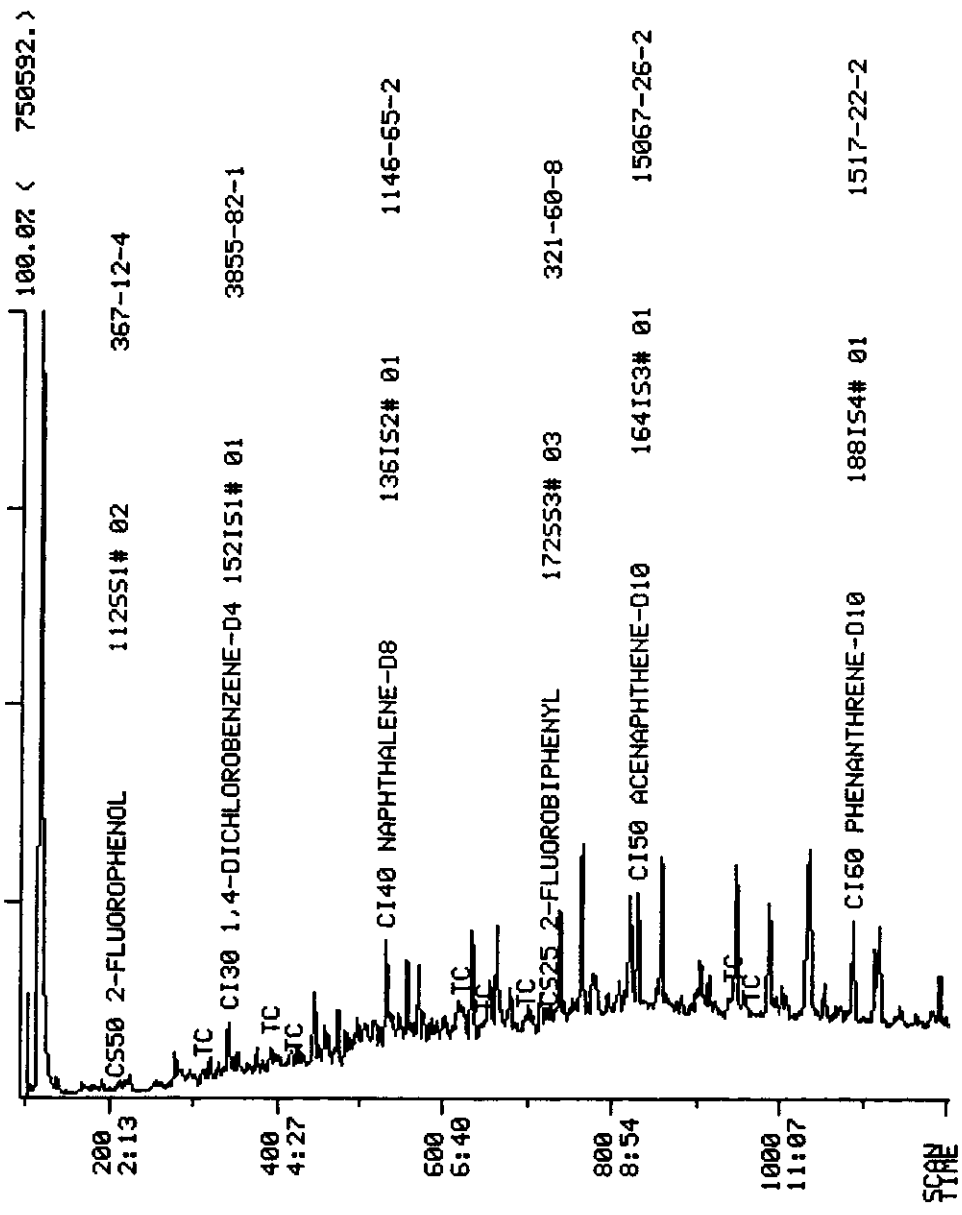
Data File: 30068110  
Std Id: ST16980831

Sample: S-MM5-4 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:28  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

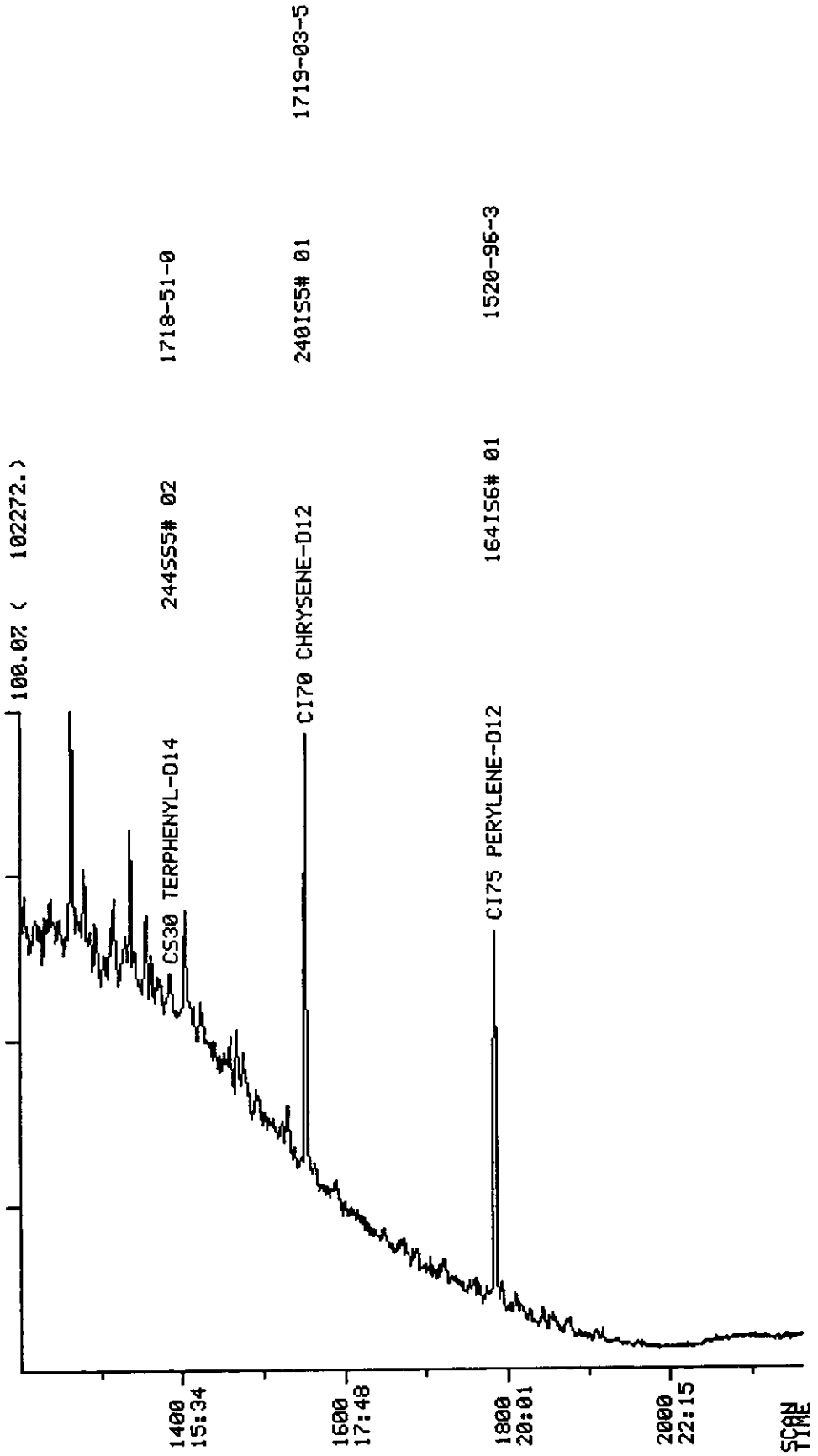
Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C775 BENZO(A)PYRENE		ND	UG/A	3000.0
C780 INDENO(1,2,3-CD)PYRENE		ND	UG/A	3000.0
C785 DIBENZ(A,H)ANTHRACENE		ND	UG/A	3000.0
C790 BENZO(G,H,I)PERYLENE		ND	UG/A	3000.0

DATA FROM FILE: 30068110 SCANS 93 TO 1203 ACQUIRED: 08/31/98 23:28:00  
 SAMPLE: S-NMS-4 1/35A/100M CALI: 30068110 #3  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M INST. ID: F16



DATA FROM FILE: 30068110 SCANS 1203 TO 2158 ACQUIRED: 08/31/98 23:28:00  
CALI: 30068110 #3  
SAMPLE: S-MM5-4 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100ML \*100%100% \*(NA/NA >1/35A NA M



ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 30068110

Sample: S-MM5-4 1/3SA/100M INST. ID: F16  
Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 23:28

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML						
No Entry	Name	Mass Meth	Scan	Ref	Area	RRF(L)	Amount	
1 S1#	1 C130 1,4-DICHLOROBENZENE	152 A VB	340	1	31781.	1.000	40.000	
2 S2#	1 C140 NAPHTHALENE-DB	136 A BB	530	2	115218.	1.000	40.000	
3 S3#	1 C150 ACENAPHTHENE-D10	164 A BB	829	3	63489.	1.000	40.000	
4 S4#	1 C160 PHENANTHRENE-D10	188 A BB	1087	4	109667.	1.000	40.000	
5 S5#	1 C170 CHRYSENE-D12	240 A BB	1552	5	68491.	1.000	40.000	
6 S6#	1 C175 PERYLENE-D12	264 A BV	1783	6	54962.	1.000	40.000	
7 S2#	2 CS20 NITROBENZENE-D5	82 A BB	424	2	535.	0.470	0.396	
8 S3#	3 CS25 2-FLUOROBIPHENYL	172 A BB	723	3	1682.	1.234	0.859	
9 S5#	2 CS30 TERPHENYL-D14	244 A VB	1384	5	2079.	0.995	1.220	
10 S1#	3 CS45 PHENOL-D5	99	1	NOT FOUND				
11 S1#	2 CS50 2-FLUOROPHENOL	112 A VV	206	1	1264.	1.368	1.163	
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330 A BB	970	3	474.	0.236	1.267	
13 S4#	4 HEXACHLOROBENZENE-C13	294	4	NOT FOUND				
14 S1#	15 C310 N-NITROSODIMETHYLAM	74	1	NOT FOUND				
15 S1#	20 PYRIDINE	79	1	NOT FOUND				
16 S1#	30 2-PICOLINE	93	1	NOT FOUND				
17 S1#	40 N-NITROSOMETHYLETHYLAMIN	42	1	NOT FOUND				
18 S1#	60 METHYLMETHANESULFONATE	80	1	NOT FOUND				
19 S1#	70 N-NITROSODIETHYLAMINE	102	1	NOT FOUND				
20 S1#	85 ETHYLMETHANE SULFONATE	79	1	NOT FOUND				
21 S1#	95 PENTACHLOROETHANE	117 A BB	306	1	252.	0.602	0.527	
22 S1#	100 C320 ANILINE	93	1	NOT FOUND				
23 S1#	105 C315 PHENOL	94 A BV	311	1	1812.	1.968	1.159	
24 S1#	110 C325 BIS(2-CHLOROETHYL)E	93 A BB	318	1	221.	1.368	0.203	
25 S1#	115 C330 2-CHLOROPHENOL	128	1	NOT FOUND				
26 S1#	125 C335 1,3-DICHLOROBENZENE	146	1	NOT FOUND				
27 S1#	130 C340 1,4-DICHLOROBENZENE	146	1	NOT FOUND				
28 S1#	145 C345 BENZYL ALCOHOL	108	1	NOT FOUND				
29 S1#	150 C350 1,2-DICHLOROBENZENE	146	1	NOT FOUND				
30 S1#	160 C355 2-METHYLPHENOL	108 A BV	392	1	1358.	1.175	1.455	
31 S1#	165 C360 2,2'-OXYBIS(1-CLPRO	45 A BB	394	1	2002.	2.539	0.992	
32 S1#	170 C361 ACETOPHENONE	105	1	NOT FOUND				
33 S1#	175 N-NITROSPYRROLIDINE	100	1	NOT FOUND				
34 S1#	180 N-NITROSOMORPHOLINE	56	1	NOT FOUND				
35 S1#	182 3-METHYL PHENOL	108	1	NOT FOUND				
36 S1#	185 C365 4-METHYLPHENOL	108 A VB	419	1	1696.	1.276	1.673	
37 S1#	190 C370 N-NITROSO-DI-N-PROP	70 A BB	412	1	677.	0.906	0.941	
38 S1#	195 O-TOLUIDINE	106	1	NOT FOUND				
39 S1#	200 C375 HEXACHLOROETHANE	117	1	NOT FOUND				
40 S2#	10 C410 NITROBENZENE	77	2	NOT FOUND				
41 S2#	15 N-NITROSOPIPERIDINE	42 A BB	449	2	763.	0.320	0.827	
42 S2#	20 C415 ISOPHORONE	82	2	NOT FOUND				
43 S2#	25 C420 2-NITROPHENOL	139	2	NOT FOUND				
44 S2#	30 C425 2,4-DIMETHYLPHENOL	107 A BB	499	2	791.	0.330	0.833	
45 S2#	35 C435 BIS(2-CHLOROETHOXY)	93	2	NOT FOUND				

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10/6/98



46	S2# 40	C440	2,4-DICHLOROPHENOL	162	2	NOT FOUND			
47	S2# 45	C430	BENZOIC ACID	122	2	NOT FOUND			
48	S2# 50	AA-DIMETHYLPHENETHYLAMIN	58	2	NOT FOUND				
49	S2# 55	C445	1,2,4-TRICHLOROBENZ	180 A VV	523	2	334.	0.331	0.350

50	S2# 60	C450 NAPHTHALENE	128	A	BV	533	2	14749.	1.022	5.009
51	S2# 80	C455 4-CHLOROANALINE	127	A	BB	555	2	464.	0.454	0.355
52	S2# 85	2,6-DICHLOROPHENOL	162							
53	S2# 90	HEXACHLOROPROPENE	213							
54	S2# 95	C460 HEXACHLOROBUTADIENE	225							
55	S2#115	P-PHENYLENE DIAMINE	108							
56	S2#120	N-NITROSODI-N-BUTYLAMINE	84	A	VB	619	2	1201.	0.213	1.957
57	S2#130	C465 4-CHLORO-3-METHYLPH	107							
58	S2#140	SAFROLE	162	A	BB	647	2	942.	0.248	1.320
59	S2#145	C470 2-METHYLNAPHTHALENE	142	A	BB	652	2	16754.	0.751	7.742
60	S3# 10	1,2,4,5-TETRACHLOROBENZE	214							
61	S3# 15	ISOSAFROLE (#1)	162	A	BB	697	3	518.	0.044	7.477
62	S3# 20	C510 HEXACHLOROCYCLOPENT	237							
63	S3# 25	C515 2,4,6-TRICHLOROPHEN	196							
64	S3# 30	C520 2,4,5-TRICHLOROPHEN	196	A	BV	715	3	406.	0.412	0.621
65	S3# 35	ISOSAFROLE (#2)	104	A	BB	738	3	283.	0.195	0.915
66	S3# 40	C525 2-CHLORONAPHTHALENE	162							
67	S3# 42	1-CHLORONAPHTHALENE	162							
68	S3# 45	C530 2-NITROANALINE	65	A	BB	763	3	579.	0.502	0.727
69	S3# 50	1,4-NAPHTHOQUINONE	158							
70	S3# 55	C535 DIMETHYLPHTHALATE	163							
71	S3# 60	1,3-DINITROBENZENE	168	A	BB	806	3	283.	0.197	0.904
72	S3# 65	C540 ACENAPHTHYLENE	152							
73	S3# 70	C543 2,6-DINITROTOLUENE	165	A	BB	816	3	221.	0.324	0.430
74	S3# 75	C545 3-NITROANILINE	138							
75	S3# 80	C550 ACENAPHTHENE	153	A	BV	834	3	1111.	1.159	0.604
76	S3# 85	C555 2,4-DINITROPHENOL	184							
77	S3# 90	C565 DIBENZOFURAN	168	A	VB	866	3	1273.	1.669	0.480
78	S3# 95	C560 4-NITROPHENOL	109							
79	S3#100	PENTACHLOROBENZENE	250							
80	S3#105	C570 2,4-DINITROTOLUENE	165	A	BB	881	3	292.	0.397	0.464
81	S3#110	1-NAPHTHYLAMINE	143							
82	S3#115	2-NAPHTHYLAMINE	143	A	BB	901	3	357.	0.966	0.233
83	S3#120	2,3,4,6-TETRACHLOROPHENO	232							
84	S3#130	C580 DIETHYLPHTHALATE	149							
85	S3#135	C590 FLUORENE	166	A	BB	926	3	1855.	1.298	0.901
86	S3#140	C585 4-CHLOROPHENYL-PHEN	204							
87	S3#145	5-NITRO-O-TOLUIDINE	152	A	VB	942	3	1344.	0.356	2.382
88	S3#150	C595 4-NITROANALINE	138	A	BB	947	3	162.	0.307	0.333
89	S4# 10	C610 4,6-DINITRO-2-METHY	198	A	BV	950	4	318.	0.132	0.879
90	S4# 15	C615 N-NITROSODIPHENYLAM	169	A	BB	963	4	1849.	0.551	1.225
91	S4# 20	C620 AZOBENZENE	77							
92	S4# 25	SYM-TRINITROBENZENE	75							
93	S4# 30	C625 4-BROMOPHENYL-PHENY	248	A	VB	1021	4	184.	0.222	0.302
94	S4# 35	PHENACETIN	108							
95	S4# 37	DIALATE	234							
96	S4# 40	C630 HEXACHLOROBENZENE	284							
97	S4# 45	4-AMINOBIIPHENYL	169							
98	S4# 50	C635 PENTACHLOROPHENOL	266	A	VB	1063	4	332.	0.155	0.781
99	S4# 55	PRONAMIDE	173							
100	S4# 60	PENTACHLORONITROBENZENE	237							
101	S4# 65	C640 PHENANTHRENE	178	A	BB	1090	4	5600.	1.033	1.978
102	S4# 70	C645 ANTHRACENE	178							
103	S4# 75	2SECBUTYL-4,6-DINITROPHE	211							
104	S4# 80	C647 CARBAZOLE	167							
105	S4# 85	C650 DI-N-BUTYLPHTHALATE	149							
106	S4#100	4-NITROQUINOLINE-1-OXIDE	190							
107	S4#105	METHAPYRILENE	58							
108	S4#106	ISODRIN	193							
109	S4#110	C655 FLUORANTHENE	202							
110	S4#120	CHLOROBENZILATE	139							

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593.4

111	S5# 10	C710 BENZIDINE	184	5	NOT FOUND			
112	S5# 15	C715 PYRENE	202 A BB	1335	5	1384.	1.323	0.611
113	S5# 20	ARAMITE (#1)	185	5	NOT FOUND			
114	S5# 25	ARAMITE (#2)	185	5	NOT FOUND			
115	S5# 30	P-DIMETHYLAMINOAZOBENZEN	120	5	NOT FOUND			
116	S5# 35	3,3'-DIMETHYLBENZIDINE	212	5	NOT FOUND			
117	S5# 37	KEPONE	272	5	NOT FOUND			
118	S5# 40	C720 BUTYLBENZYLPHTHALAT	149	5	NOT FOUND			
119	S5# 45	2-ACETYLAMINOFLUORENE	181	5	NOT FOUND			
120	S5# 50	C730 BENZO(A)ANTHRACENE	228 A BB	1554	5	667.	1.125	0.346
121	S5# 55	C725 3,3'-DICHLOROBENZID	252 A VB	1564	5	342.	0.408	0.490
122	S5# 60	C740 CHRYSENE	228 A BB	1554	5	667.	1.014	0.384
123	S5# 65	C745 BIS(2-ETHYLHEXYL)PH	149	5	NOT FOUND			
124	S5# 85	3-METHYLCHOLANTHRENE	268	5	NOT FOUND			
125	S6# 10	C760 DI-N-OCTYL PHTHALAT	149	6	NOT FOUND			
126	S6# 15	C765 BENZO(B)FLUORANTHEN	252	6	NOT FOUND			
127	S6# 20	7,12-DIMETHYLBENZANTHRAC	256 A BB	1740	6	194.	0.610	0.231
128	S6# 25	C770 BENZO(K)FLUORANTHEN	252	6	NOT FOUND			
129	S6# 30	HEXACHLOROPHENE	196	6	NOT FOUND			
130	S6# 35	C775 BENZO(A)PYRENE	252 A BB	1783	6	522.	1.182	0.321
131	S6# 55	C780 INDENO(1,2,3-CD)PYR	276	6	NOT FOUND			
132	S6# 60	C785 DIBENZ(A,H)ANTHRACE	278	6	NOT FOUND			
133	S6# 65	C790 BENZO(G,H,I)PERYLEN	276	6	NOT FOUND			

QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.==+200		4.28	200864.==+200		6.40	108695.==+200		9.71
Lower Limit	14324.==+ 50		3.28	50216.==+ 50		5.40	27174.==+ 50		8.71
-----									
Filename									
=====									
1 30068110	31781.= 110		3.78	115218.= 114		5.90	63489.= 116		9.22
-----									

IS# 1 = C130 1,4-DICHLOROBENZENE-D4  
 IS# 2 = C140 NAPHTHALENE-D8  
 IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.==+200		12.58	124578.==+200		17.76	101500.==+200		20.33
Lower Limit	49260.==+ 50		11.58	31145.==+ 50		16.76	25375.==+ 50		19.33
-----									
Filename									
=====									
1 30068110	109667.= 111		12.09	68491.= 109		17.26	54962.= 108		19.83
-----									

IS# 4 = C160 PHENANTHRENE-D10  
 IS# 5 = C170 CHRYSENE-D12  
 IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

TARGET COMPOUND COMPARISON

COMPOUND: C450 NAPHTHALENE

128 S2# 60

91-20-3

RAW DATA: 30058110 #533

08/31/98 23:28

SAMPLE: S-MM5-4 1/35A/100M

CONDS.: UG/ML \*100%/100% \*(NA/NA )/1/35A NA M

*Handwritten:* "S" and a signature

BASE M/Z: 41

RIC: 80128.

4720.

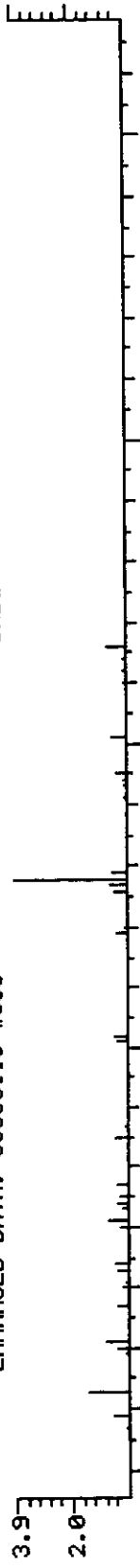


ENHANCED DATA: 30058110 #533

BASE M/Z: 128

RIC: 12508.

3108.



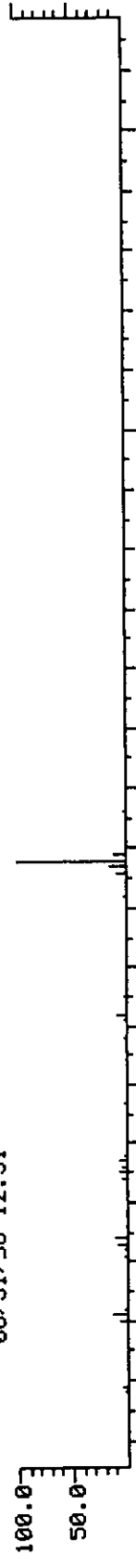
STANDARD FILE: ST16980831 #533

08/31/98 12:31

BASE M/Z: 128

RIC: 195840.

79232.



\*\*OUT\*\*

TARGET COMPOUND COMPARISON

COMPOUND: C470 2-METHYLNAPHTHALENE 142 S2#145 91-57-6

RAW DATA: 30068110 #652 BASE M/Z: 41 RIC: 99840.

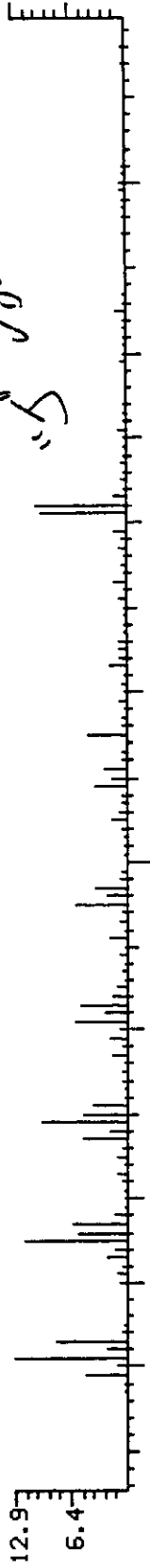
08/31/98 23:28

SAMPLE: S-MMS-4 1/35A/100M INST. ID: F16

CONDS.: UG/ML \*100Z/100Z \*(NA/NA) >1/35A NA M

6416.

*Handwritten initials: "S" and "J" with a checkmark.*



ENHANCED DATA: 30068110 #652

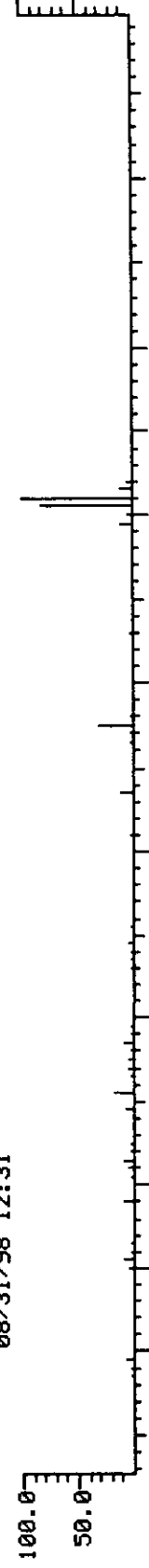
4328.



STANDARD FILE: ST16980831 #651

08/31/98 12:31

49792.



M/Z 100.0 50.0 0.0 -50.0 -100.0



\*\*OUT\*\*

TARGET COMPOUND COMPARISON

COMPOUND: C640 PHENANTHRENE

178 S4# 65

85-01-8

RAW DATA: 30058110 #1090

08/31/98 23:28

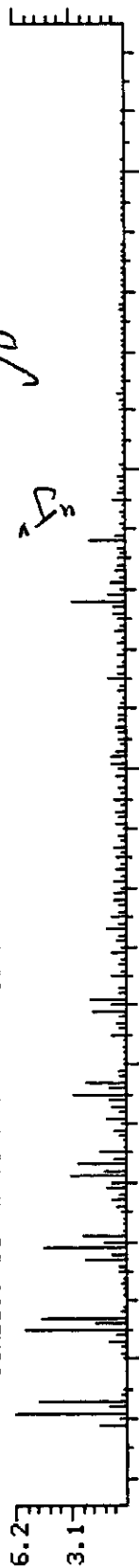
SAMPLE: S-MM5-4 1/3SA/100M

COND.: UG/ML \*100%/100Z \*(NA/NA )/1/3SA NA M

BASE M/Z: 41

RIC: 91008.

4512.

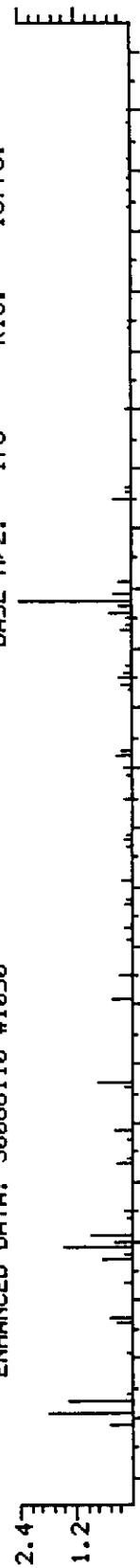


ENHANCED DATA: 30058110 #1090

BASE M/Z: 178

RIC: 13776.

1724.

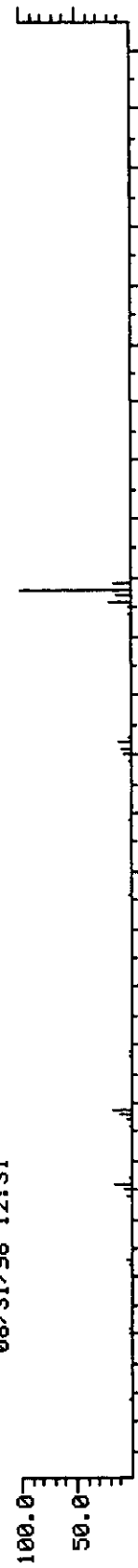


STANDARD FILE: ST16980831 #1091

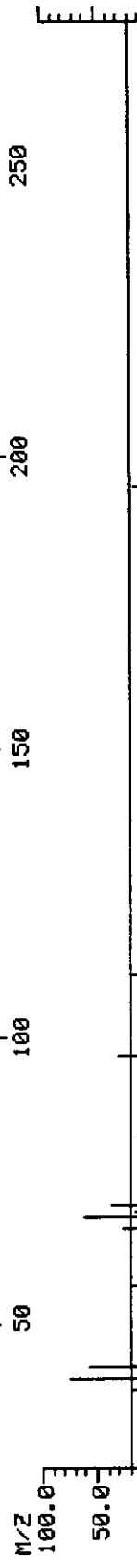
BASE M/Z: 178

RIC: 204544.

73344.



\*\*OUT\*\*



Data Reduced by: *EJ* Date: *9/18/98*  
Data Reviewed by: *JW* Date: *4.11.98*

Data File: 30068110

QUANTERRA GC/MS TIC REPORT ( Part 1 )

Sample: S-MM5-4 1/3SA/100M INST. ID: F16  
Analyst: DAT Date Analyzed: 08/31/98 23:28  
Run Factor: 300.

# SCAN	Concentration in Sample (UG/A)	CAS #	
1 444	7600.	1120-21-4	
			UNDECANE or isomer
<hr/>			
2 473	7500.	<del>74962-98-4</del>	unknown 5700
			<del>2 TRIDECEN-1-OL, (E)</del>
<hr/>			
3 555	7100.	<del>62016-37-9</del>	alkane 5746400
			<del>OCTANE, 2,4,6-TRIMETHYL</del>
<hr/>			
4 569	6200.	62108-22-9	
			DECANE, 2,5,9-TRIMETHYL- or isomer
<hr/>			
5 616	12000.	00-00-0	5700
			UNKNOWN
<hr/>			
6 632	12000.	<del>54105-67-8</del>	alkane
			<del>HEPTADECANE, 2,6-DIMETHYL</del>
<hr/>			
7 654	6900.	<del>36653-82-4</del>	unkn. hydrocarbon 57904300
			<del>1-HEXADECANOL</del>
<hr/>			
8 662	9800.	15869-93-9	
			OCTANE, 3,5-DIMETHYL- or isomer
<hr/>			
9 699	6400.	00-00-0	5700
			UNKNOWN
<hr/>			
X 714	4800.	00-00-0	
			UNKNOWN





11	737	13000.	54105-67-8	
HEPTADECANE, 2,6-DIMETHYL- <i>or isomer</i>				
12	763	18000.	629-59-4	
TETRADECANE				
13	776	8600.	00-00-0	5700
UNKNOWN				
14	820	12000.	18344-37-1	
HEPTADECANE, 2,6,10,14-TETRAMETHYL- <i>or isomer</i>				
15	857	14000.	629-62-9	
PENTADECANE <i>or isomer</i>				
16	903	6400.	5234-28-4	5700
<del>OXIRANE, DODECYL-</del> <i>unknown</i>				
17	947	16000.	629-92-5	
NONADECANE <i>or isomer</i>				
18	987	13000.	17301-23-4	
UNDECANE, 2,6-DIMETHYL- <i>or isomer</i>				
19	1035	28000.	54105-67-8	
HEPTADECANE, 2,6-DIMETHYL-				
<del>20</del>	<del>1053</del>	<del>5300.</del>	<del>7206-19-1</del>	<i>unknown 5700</i>
<del>3-OCTADECENE, (E)-</del>				
21	1113	7600.	29812-79-1	
<del>HYDROXYLAMINE, O-DECYL-</del> <i>alkane</i>				
22	1118	9800.	54105-67-8	
HEPTADECANE, 2,6-DIMETHYL- <i>or isomer</i>				

X 1190 6400. 629-92-5  
NONADECANE or isomer

X 1264 6400. ~~34105-67-8~~  
~~HEPTADECANE, 2,6-DIMETHYL-~~

alkane



X 1335 5200. ~~00-00-0~~  
UNKNOWN

5700 29/18/98

QUANTERRA GC/MS TIC REPORT ( Part 2 )

CONCENTRATION = AREA(TIC)\*CONC(IS)/AREA(IS)

#	FIT	PURITY	INT.		AREA	HEIGHT	AMOUNT		LIB	LIB #
			STD.	RT			RRT	(UG/ML )		
1	984	701	2	4:56	0.536	231686.	64883.	25.245	NB	11607.
2	883	667	2	5:15	0.571	230400.	53289.	25.105	NB	22482.
3	966	816	2	6:10	0.669	218304.	73600.	23.787	NB	11602.
4	964	831	2	6:19	0.686	190400.	68571.	20.746	NB	19013.
5	866	467	2	6:51	0.743	356160.	34752.	38.808	UK	1.
6	931	784	2	7:01	0.762	370112.	103303.	40.328	NB	37462.
7	922	681	2	7:16	0.789	211506.	41371.	23.046	NB	32420.
8	939	825	2	7:21	0.799	300032.	94196.	32.692	NB	8104.
9	786	462	2	7:46	0.843	194560.	23981.	21.199	UK	1.
10	865	422	2	7:56	0.861	146432.	35328.	15.955	UK	1.
11	953	832	2	8:11	0.889	403712.	103936.	43.989	NB	37462.
12	960	734	2	8:29	0.920	555008.	162304.	60.474	NB	22530.
13	902	440	2	8:37	0.936	263296.	38272.	28.689	UK	1.
14	957	792	2	9:07	0.989	380416.	103424.	41.450	NB	42196.
15	955	770	2	9:31	1.034	436116.	137255.	47.520	NB	25997.
16	918	641	2	10:02	1.089	197120.	44544.	21.478	NB	25971.
17	958	842	2	10:32	1.142	479616.	139008.	52.259	NB	37465.
18	949	827	3	10:58	0.908	400256.	105216.	44.229	NB	19054.
19	971	867	3	11:30	0.952	844160.	153344.	93.281	NB	37462.
20	901	704	3	11:42	0.969	161024.	34944.	17.793	NB	34410.
21	948	794	3	12:22	1.024	228224.	70016.	25.219	NB	15969.
22	941	834	3	12:26	1.029	295680.	89856.	32.673	NB	37462.
23	940	728	3	13:14	1.095	191872.	48640.	21.202	NB	37465.
24	921	706	3	14:03	1.163	194304.	38592.	21.471	NB	37462.
25	855	571	4	14:50	0.860	110720.	22639.	17.169	UK	1.

Library Search                    Data: 30068110 # 444            Base m/z: 43  
 08/31/98 23:28:00 + 4:56        Cali: 30068110 # 3            RIC: 54464.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 892 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 11607 UNDECANE  
 2 19015 DECANE, 2,5,6-TRIMETHYL-  
 3 11565 ISOCTANE, (ETHENYLOXY)-  
 4 8104 OCTANE, 3,5-DIMETHYL-  
 5 19523 1-DECANOL, 2-ETHYL-  
 6 14793 1-UNDECENE, 4-METHYL-  
 7 5151 HEPTANE, 3,4-DIMETHYL-  
 8 15969 HYDROXYLAMINE, O-DECYL-  
 9 12050 1-OCTANOL, 2,7-DIMETHYL-

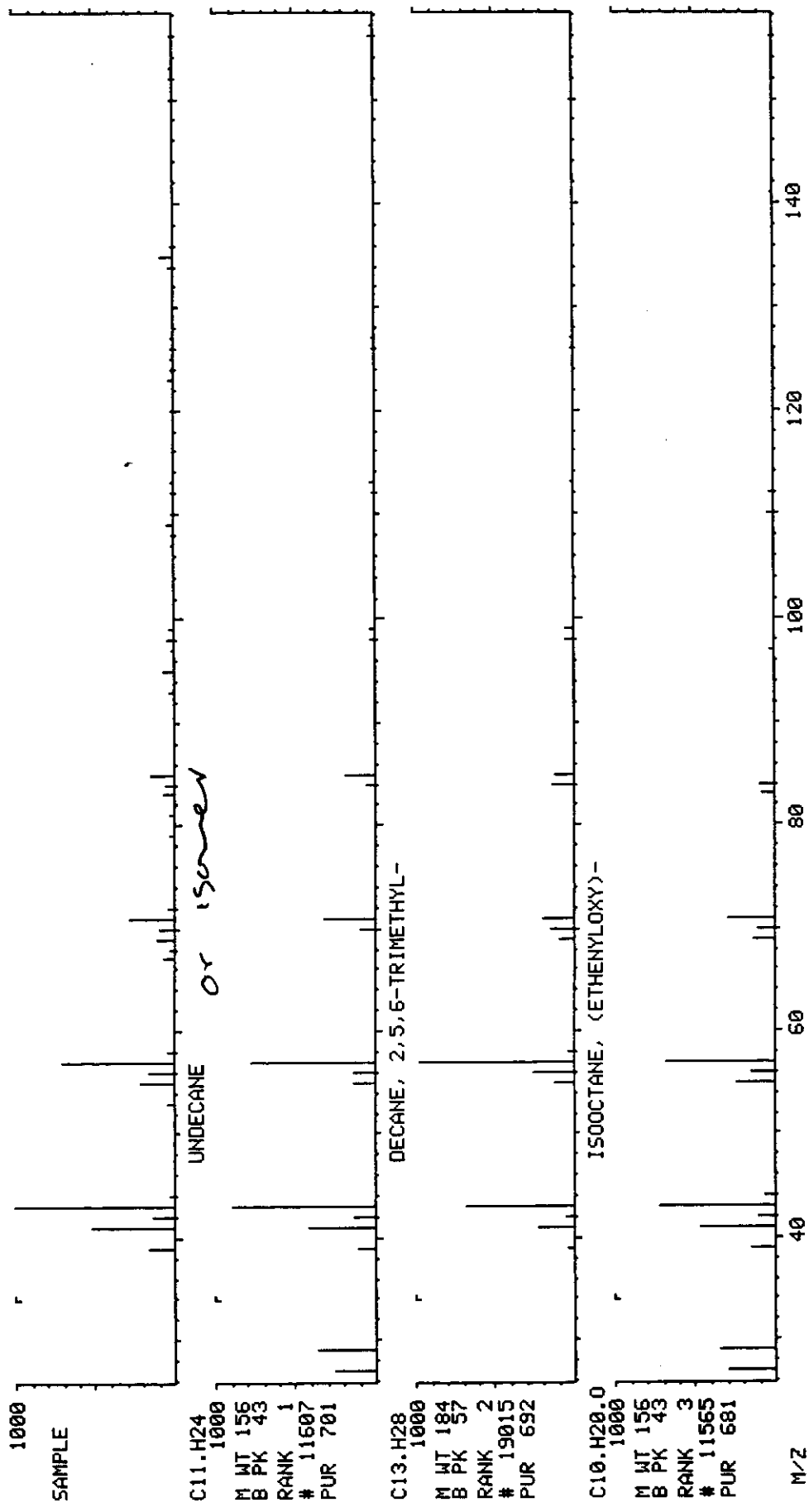
Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C11.H24	156	43	701	984	703
2	C13.H28	184	57	692	934	692
3	C10.H20.O	156	43	681	975	686
4	C10.H22	142	57	677	932	683
5	C12.H26.O	186	57	677	922	697
6	C12.H24	168	43	670	925	673
7	C9.H20	128	43	668	922	670
8	C10.H23.O.N	173	43	665	930	698
9	C10.H22.O	158	43	665	935	665

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	1120-21-4
2	---	---	---	---	62108-23-0
3	---	---	---	---	37769-62-3
4	---	---	---	---	15869-93-9
5	---	---	---	---	21078-65-9
6	---	---	---	---	74630-39-0
7	---	---	---	---	922-28-1
8	---	---	---	---	29812-79-1
9	---	---	---	---	15250-22-3

DATA: 30068110 # 444  
CALI: 30068110 # 3  
BASE M/Z: 43  
RIC: 54464.

DATA: 30068110 # 444  
CALI: 30068110 # 3

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:28:00 + 4:56  
SAMPLE: 5-NM5-4 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 15B 2N 0T)



Library Search                    Data: 30068110 # 473            Base m/z: 43  
 08/31/98 23:28:00 + 5:16        Cali: 30068110 # 3            RIC: 45440.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 367 matched at least 7 of the 16 largest peaks in the unknown

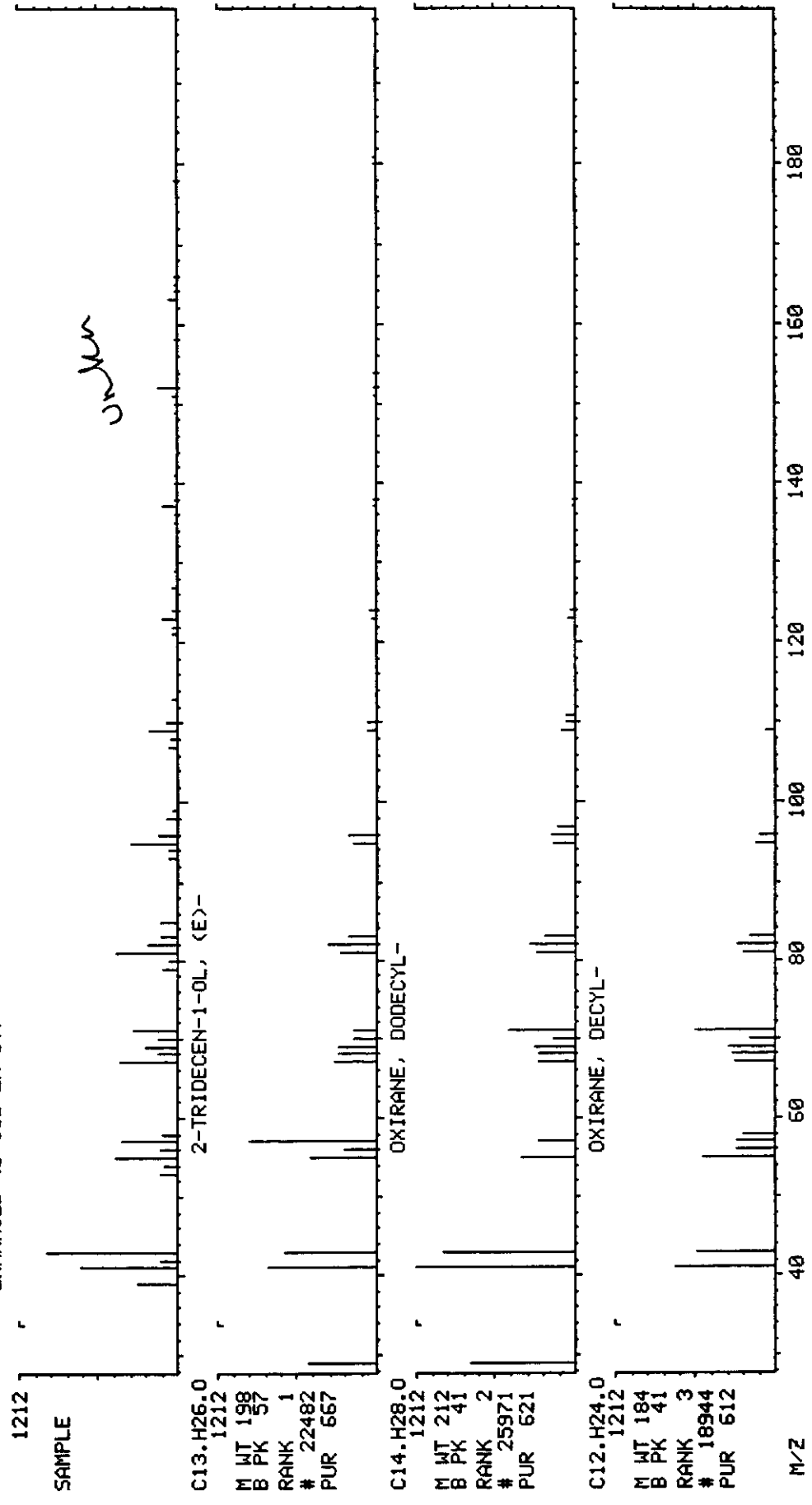
Rank In.        Name  
 1 22482 2-TRIDECEN-1-OL, (E)-  
 2 25971 OXIRANE, DODECYL-  
 3 18944 OXIRANE, DECYL-  
 4 39230 9-EICOSYNE  
 5 15294 TRANS-2-UNDECEN-1-OL  
 6 37449 OCTADECANAL  
 7 28267 3-HEXADECYNE  
 8 28261 7-HEXADECYNE  
 9 34010 3-OCTADECYNE

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C13.H26.O	198	57	667	883	674
2	C14.H28.O	212	41	621	892	654
3	C12.H24.O	184	41	612	919	612
4	C20.H38	278	81	609	876	642
5	C11.H22.O	170	57	609	894	609
6	C18.H36.O	268	43	599	811	671
7	C16.H30	222	67	598	916	603
8	C16.H30	222	67	597	900	603
9	C18.H34	250	67	597	908	602

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	74962-98-4
2	---	---	---	---	3234-28-4
3	---	---	---	---	2855-19-8
4	---	---	---	---	71899-38-2
5	---	---	---	---	- -
6	---	---	---	---	638-66-4
7	---	---	---	---	61886-62-2
8	---	---	---	---	74685-28-2
9	---	---	---	---	61886-64-4

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 5:16  
 SAMPLE: S-MM5-4 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068110 # 473  
 CALI: 30068110 # 3  
 BASE M/Z: 43  
 RIC: 45440



1320

Library Search                    Data: 30068110 # 555            Base m/z: 43  
 08/31/98 23:28:00 + 6:10        Cali: 30068110 # 3            RIC: 71808.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 988 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 11602 OCTANE, 2,4,6-TRIMETHYL-  
 2 15969 HYDROXYLAMINE, O-DECYL-  
 3 11607 UNDECANE  
 4 19016 UNDECANE, 4,7-DIMETHYL-  
 5 19006 DECANE, 6-ETHYL-2-METHYL-  
 6 19015 DECANE, 2,5,6-TRIMETHYL-  
 7 14793 1-UNDECENE, 4-METHYL-  
 8 22530 TETRADECANE  
 9 19028 DECANE, 2,4,6-TRIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C11.H24	156	57	816	966	816
2	C10.H23.O.N	173	43	816	940	834
3	C11.H24	156	43	814	952	851
4	C13.H28	184	43	813	946	821
5	C13.H28	184	57	807	940	826
6	C13.H28	184	57	807	926	817
7	C12.H24	168	43	800	932	812
8	C14.H30	198	43	798	910	866
9	C13.H28	184	43	796	924	812

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	62016-37-9
2	---	---	---	---	29812-79-1
3	---	---	---	---	1120-21-4
4	---	---	---	---	17301-32-5
5	---	---	---	---	62108-21-8
6	---	---	---	---	62108-23-0
7	---	---	---	---	74630-39-0
8	---	---	---	---	629-59-4
9	---	---	---	---	62108-27-4

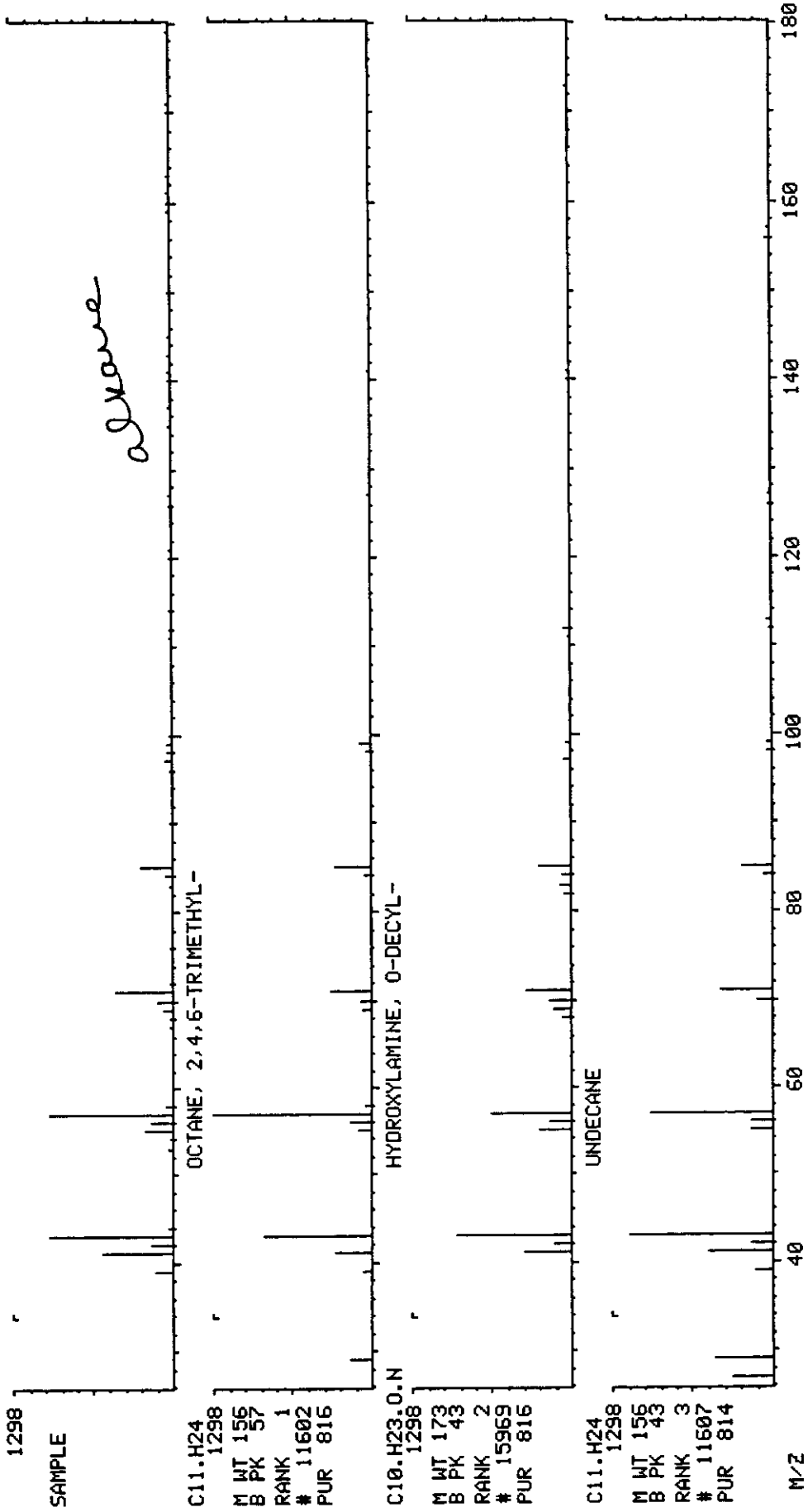


DATA: 30068110 # 555  
CALI: 30068110 # 3

BASE M/Z: 43  
RIC: 71808.

MID LIBRARY SEARCH (LIBRARYNB)

08/31/98 23:28:00 + 6:10  
SAMPLE: S-MM5-4 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100%/100% \*(NA/NA) >1/35A NA M  
ENHANCED (S 15B 2N 0T)



Library Search                    Data: 30068110 # 569            Base m/z: 57  
 08/31/98 23:28:00 + 6:20        Cali: 30068110 # 3            RIC: 62144.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 259 matched at least 8 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 19013 DECANE, 2,5,9-TRIMETHYL-  
 2 19054 UNDECANE, 2,6-DIMETHYL-  
 3 8104 OCTANE, 3,5-DIMETHYL-  
 4 19026 DECANE, 2,6,8-TRIMETHYL-  
 5 8077 HEPTANE, 2,3,5-TRIMETHYL-  
 6 8092 HEPTANE, 2,3,6-TRIMETHYL-  
 7 5154 HEXANE, 4-ETHYL-2-METHYL-  
 8 8081 OCTANE, 2,3-DIMETHYL-  
 9 11602 OCTANE, 2,4,6-TRIMETHYL-

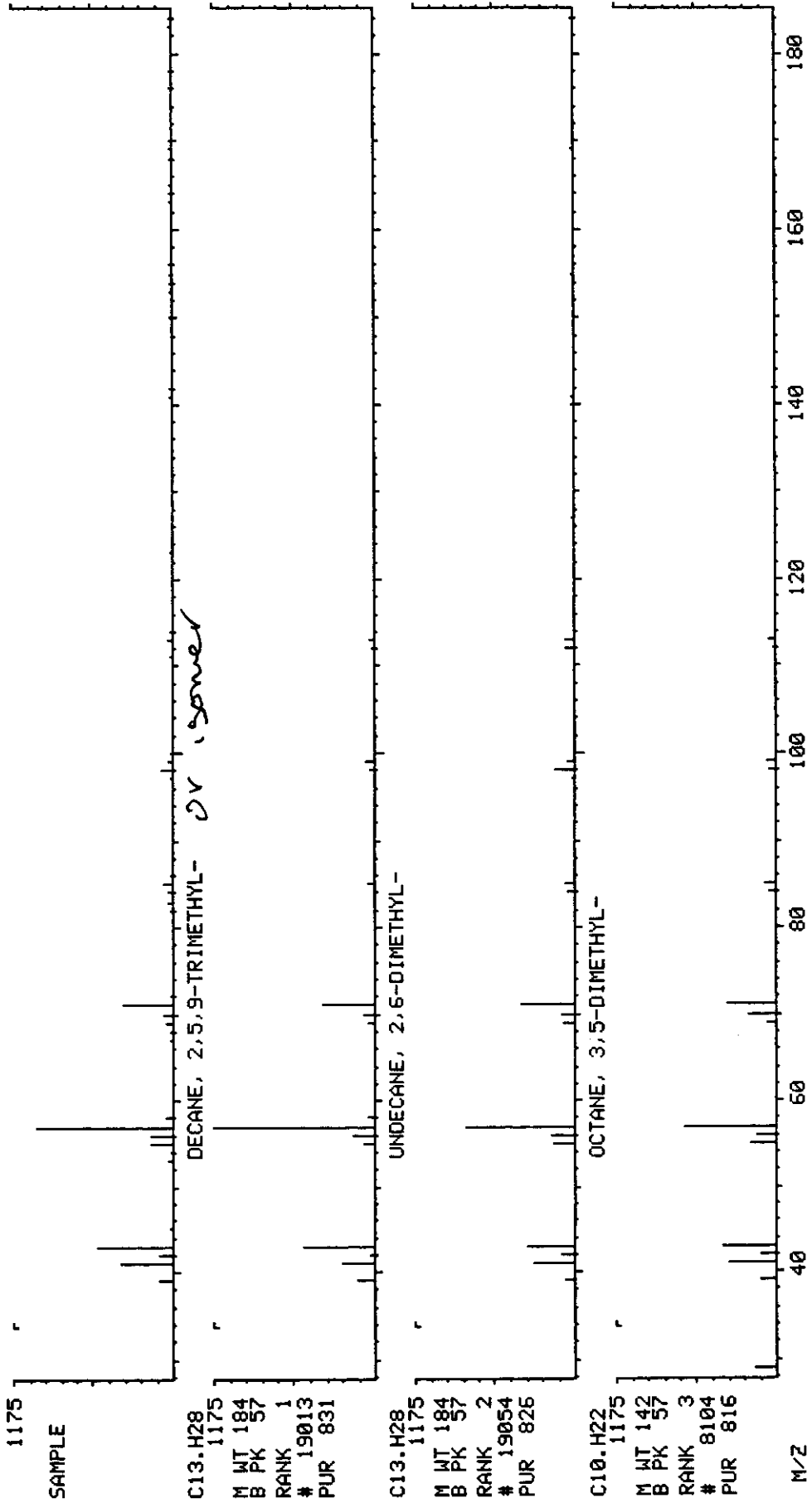
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C13.H28	184	57	831	964	841
2	C13.H28	184	57	826	935	863
3	C10.H22	142	57	816	949	824
4	C13.H28	184	57	815	949	836
5	C10.H22	142	57	810	954	810
6	C10.H22	142	57	809	955	809
7	C9.H20	128	57	801	947	806
8	C10.H22	142	57	796	940	798
9	C11.H24	156	57	788	933	798

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	62108-22-9
2	---	---	---	---	17301-23-4
3	---	---	---	---	15869-93-9
4	---	---	---	---	62108-26-3
5	---	---	---	---	20278-85-7
6	---	---	---	---	4032-93-3
7	---	---	---	---	3074-75-7
8	---	---	---	---	7146-60-3
9	---	---	---	---	62016-37-9

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:28:00 + 6:20  
 SAMPLE: S-MMS-4 1/35A/100M  
 COND5.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068110 # 569  
 CALI: 30068110 # 3

BASE M/Z: 57  
 RIC: 62144.



Library Search                    Data: 30068110 # 616            Base m/z: 55  
 08/31/98 23:28:00 + 6:51        Cali: 30068110 # 3            RIC: 31296.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

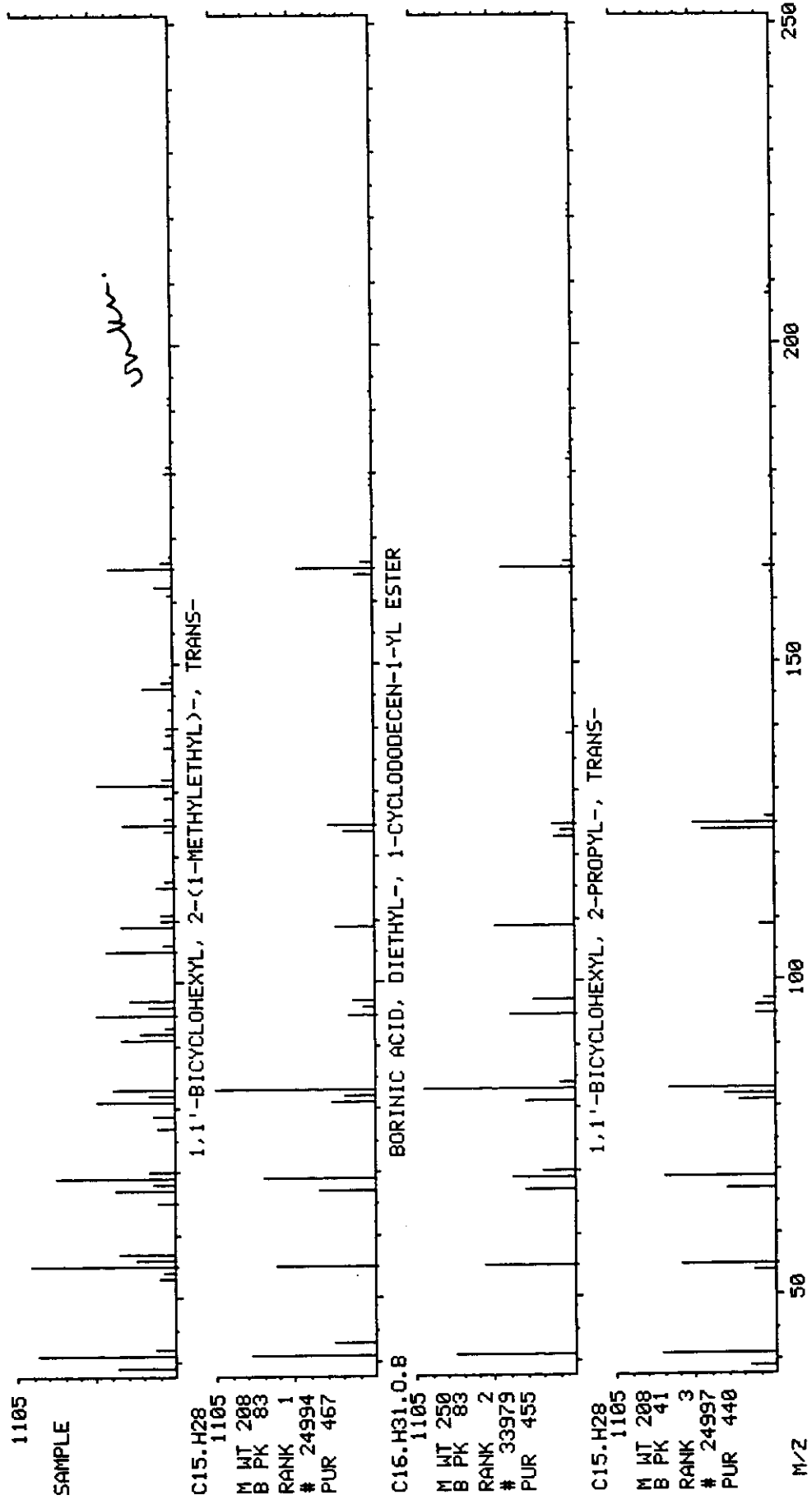
62231 spectra in LIBRARYNB searched for maximum PURITY  
 457 matched at least 6 of the 16 largest peaks in the unknown

- Rank In.            Name  
 1 24994 1,1'-BICYCLOHEXYL, 2-(1-METHYLETHYL)-, TRANS-  
 2 33979 BORINIC ACID, DIETHYL-, 1-CYCLODODECEN-1-YL ESTER  
 3 24997 1,1'-BICYCLOHEXYL, 2-PROPYL-, TRANS-  
 4 24993 1,1'-BICYCLOHEXYL, 2-(1-METHYLETHYL)-, CIS-  
 5 24996 1,1'-BICYCLOHEXYL, 2-PROPYL-, CIS-  
 6 39230 9-EICOSYNE  
 7 32770 1,15-PENTADECANEDIOL  
 8 29229 CYCLOPENTADECANOL-  
 9 14159 1,11-DODECADIENE

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C15.H28	208	83	467	866	496
2	C16.H31.O.B	250	83	455	875	492
3	C15.H28	208	41	440	789	448
4	C15.H28	208	83	430	834	477
5	C15.H28	208	69	416	787	435
6	C20.H38	278	81	413	753	464
7	C15.H32.O2	244	55	410	890	426
8	C15.H30.O	226	55	408	743	451
9	C12.H22	166	55	405	838	421

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	50991-16-7
2	---	---	---	---	61142-73-2
3	---	---	---	---	54934-89-3
4	---	---	---	---	50991-15-6
5	---	---	---	---	54934-88-2
6	---	---	---	---	71899-38-2
7	---	---	---	---	- -
8	---	---	---	---	4727-17-7
9	---	---	---	---	5876-87-9

MID LIBRARY SEARCH (LIBRARYNB) DATA: 30068110 # 616 BASE M/Z: 55  
 08/31/98 23:28:00 + 6:51 CALI: 30068110 # 3 RIC: 31296.  
 SAMPLE: S-MMS-4 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)



Library Search                    Data: 30068110 # 632            Base m/z: 57  
 08/31/98 23:28:00 + 7:02        Cali: 30068110 # 3            RIC: 90240.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 312 matched at least 8 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 37462 HEPTADECANE, 2,6-DIMETHYL-  
 2 14799 1-DECENE, 3,4-DIMETHYL-  
 3 19523 1-DECANOL, 2-ETHYL-  
 4 14751 2-UNDECENE, 5-METHYL-  
 5 22534 TRIDECANE, 7-METHYL-  
 6 8539 1-PENTANOL, 4-METHYL-2-PROPYL-  
 7 15353 2,6-DIMETHYLDECANE  
 8 25994 DODECANE, 2,6,11-TRIMETHYL-  
 9 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-

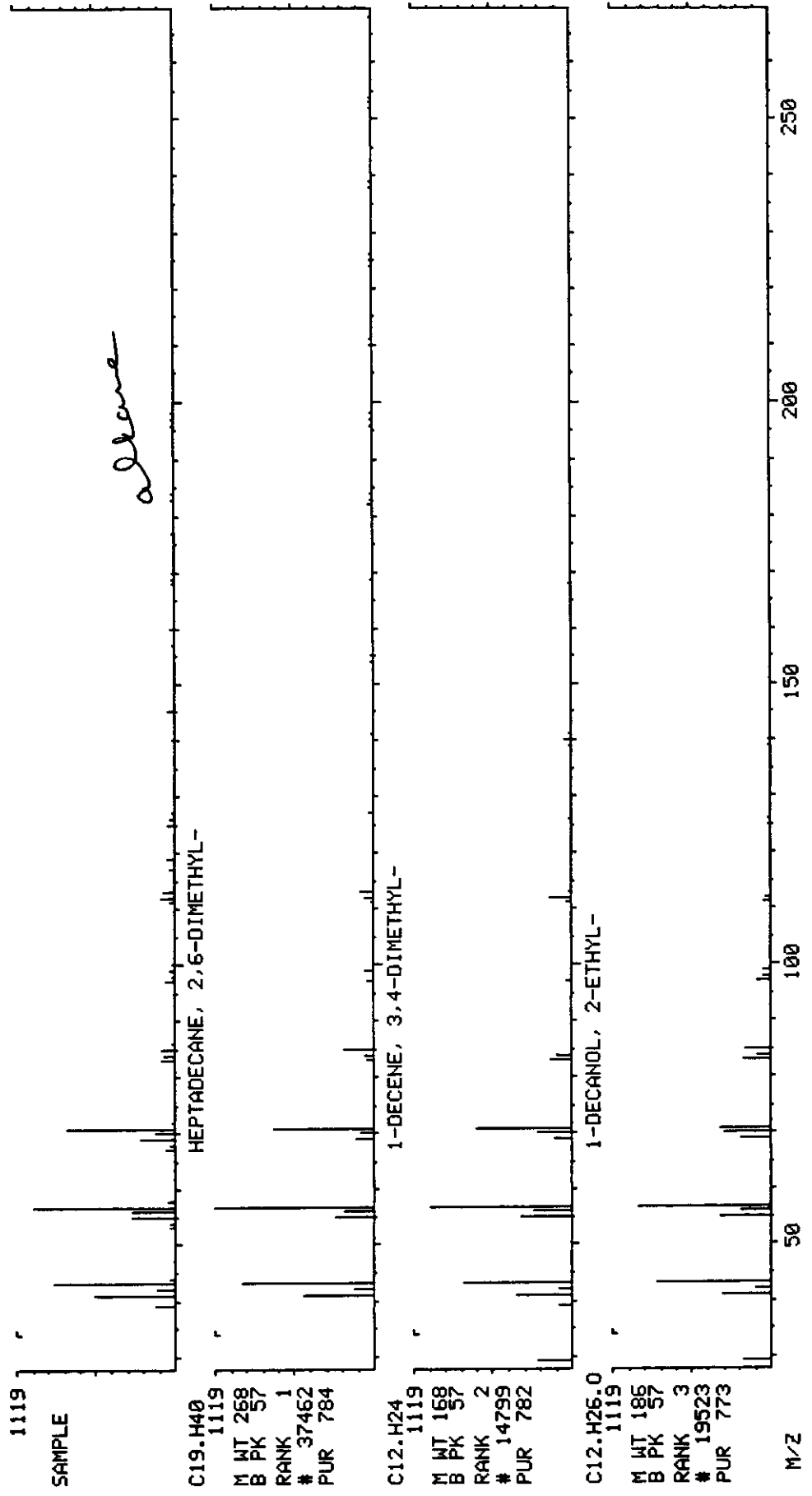
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	784	931	815
2	C12.H24	168	57	782	953	785
3	C12.H26.O	186	57	773	934	779
4	C12.H24	168	57	768	966	768
5	C14.H30	198	57	766	921	780
6	C9.H20.O	144	57	756	974	756
7	C12.H26	170	43	753	915	762
8	C15.H32	212	57	751	947	756
9	C21.H44	296	57	748	908	788

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	54105-67-8
2	---	---	---	---	50871-03-9
3	---	---	---	---	21078-65-9
4	---	---	---	---	56851-34-4
5	---	---	---	---	26730-14-3
6	---	---	---	---	54004-41-0
7	---	---	---	---	13150-81-7
8	---	---	---	---	31295-56-4
9	---	---	---	---	18344-37-1

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 7:02  
 SAMPLE: S-MMS-4 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068110 # 632  
 CALI: 30068110 # 3

BASE M/Z: 57  
 RIC: 90240.



*allene*

Library Search                    Data: 30068110 # 654            Base m/z: 41  
 08/31/98 23:28:00 + 7:16        Cali: 30068110 # 3            RIC: 43584.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 820 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	32420 1-HEXADECANOL
2	21962 3-TETRADECENE, (Z)-
3	19526 1-DODECANOL
4	21963 3-TETRADECENE, (E)-
5	28772 7-HEXADECENE, (Z)-
6	31653 1-HEPTADECENE
7	28768 3-HEXADECENE, (Z)-
8	40193 CIS-9,10-EPOXYOCTADECAN-1-OL
9	34411 5-OCTADECENE, (E)-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H34.O	242	55	681	922	692
2	C14.H28	196	41	680	948	689
3	C12.H26.O	186	43	678	950	687
4	C14.H28	196	41	677	944	687
5	C16.H32	224	55	677	912	707
6	C17.H34	238	55	676	922	699
7	C16.H32	224	55	674	915	707
8	C18.H36.O2	284	55	674	903	706
9	C18.H36	252	55	672	905	704

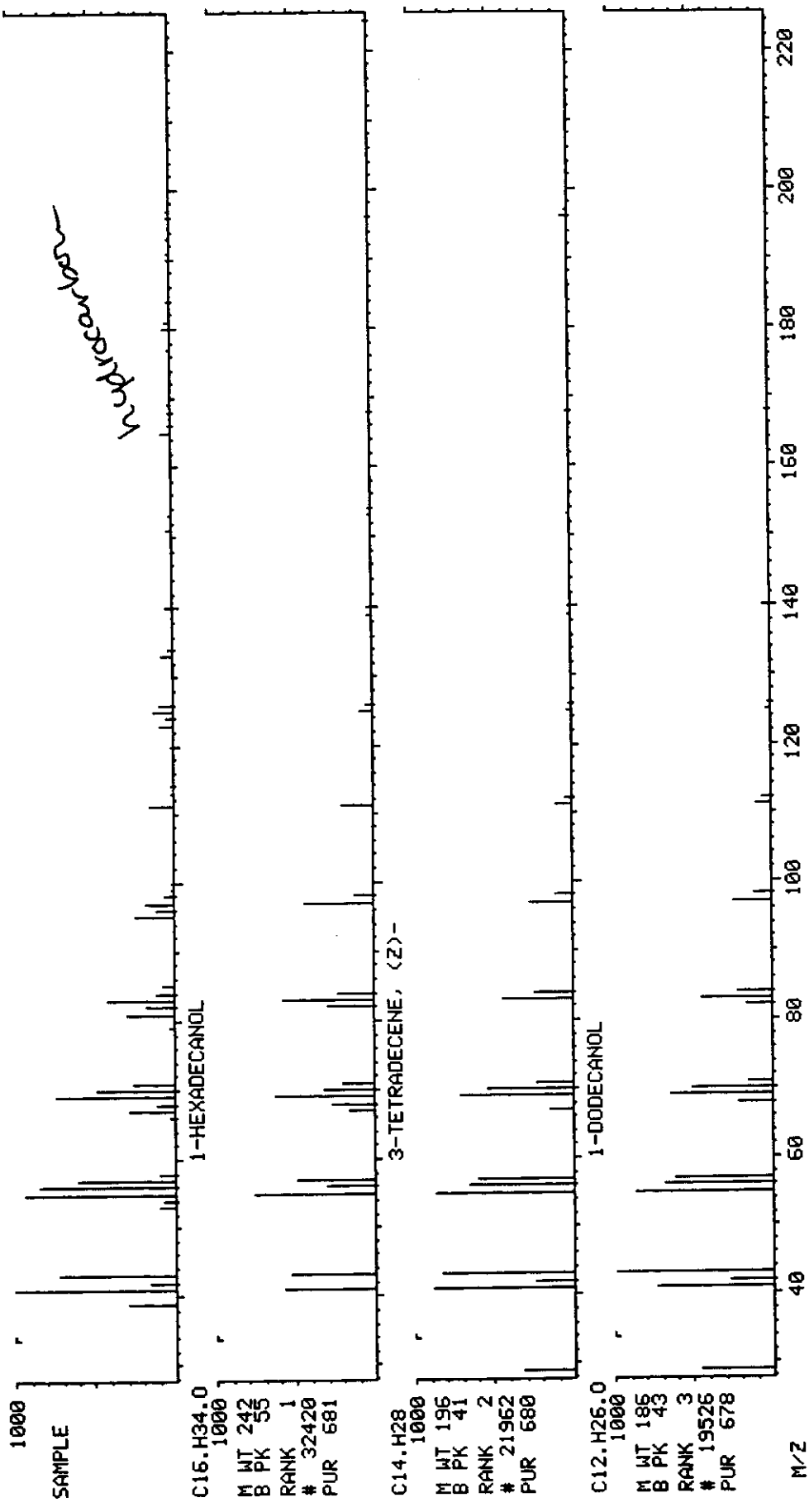
Rank	Ret.Time	B.P.int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	36653-82-4
2	---	---	---	---	41446-67-7
3	---	---	---	---	112-53-8
4	---	---	---	---	41446-68-8
5	---	---	---	---	35507-09-6
6	---	---	---	---	6765-39-5
7	---	---	---	---	34303-81-6
8	---	---	---	---	13980-12-6
9	---	---	---	---	7206-21-5



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 7:16  
 SAMPLE: S-MM5-4 1-35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1-35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068110 # 654  
 CALI: 30068110 # 3

BASE M/Z: 41  
 RIC: 43584.



Library Search                      Data: 30068110 # 662                      Base m/z: 43  
 08/31/98 23:28:00 + 7:22            Cali: 30068110 # 3                      RIC: 88832.  
 Sample: S-MM5-4 1/3SA/100M            INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 346 matched at least 8 of the 16 largest peaks in the unknown

Rank In.	Name
1	8104 OCTANE, 3,5-DIMETHYL-
2	19015 DECANE, 2,5,6-TRIMETHYL-
3	11602 OCTANE, 2,4,6-TRIMETHYL-
4	19016 UNDECANE, 4,7-DIMETHYL-
5	19013 DECANE, 2,5,9-TRIMETHYL-
6	19026 DECANE, 2,6,8-TRIMETHYL-
7	18985 TRIDECANE
8	8102 HEXANE, 2,2,3,3-TETRAMETHYL-
9	19054 UNDECANE, 2,6-DIMETHYL-

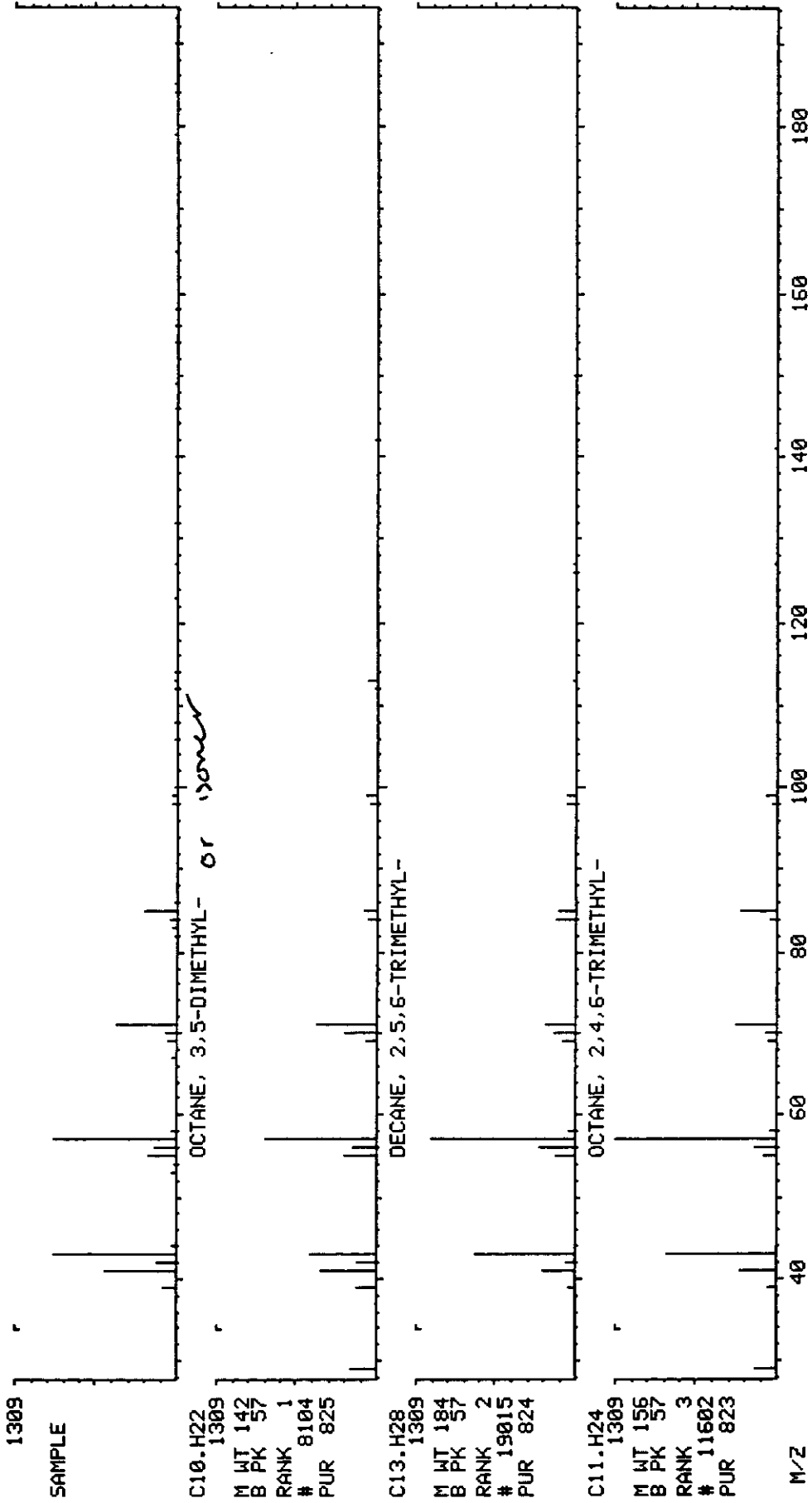
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H22	142	57	825	939	832
2	C13.H28	184	57	824	931	828
3	C11.H24	156	57	823	968	825
4	C13.H28	184	43	822	934	843
5	C13.H28	184	57	815	935	815
6	C13.H28	184	57	811	928	828
7	C13.H28	184	57	808	939	830
8	C10.H22	142	57	800	923	800
9	C13.H28	184	57	797	889	822

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	15869-93-9
2	---	---	---	---	62108-23-0
3	---	---	---	---	62016-37-9
4	---	---	---	---	17301-32-5
5	---	---	---	---	62108-22-9
6	---	---	---	---	62108-26-3
7	---	---	---	---	629-50-5
8	---	---	---	---	13475-81-5
9	---	---	---	---	17301-23-4

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:28:00 + 7:22  
 SAMPLE: S-MM5-4 1/35A/100M  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068110 # 662  
 CALI: 30068110 # 3

BASE M/Z: 43  
 RIC: 88832.



1355

Library Search                    Data: 30068110 # 699            Base m/z: 81  
 08/31/98 23:28:00 + 7:46        Cali: 30068110 # 3            RIC: 23040.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 366 matched at least 6 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 28273 8-HEXADECYNE  
 2 36677 7-OCTADECYNE, 2-METHYL-  
 3 28267 3-HEXADECYNE  
 4 34010 3-OCTADECYNE  
 5 36681 1H-INDENE, 2-BUTYL-5-HEXYLOCTAHYDRO-  
 6 10423 3-UNDECYNE  
 7 25001 CYCLOHEXANE, 1-(CYCLOHEXYLMETHYL)-2-ETHYL-, TRANS-  
 8 25003 CYCLOHEXANE, 1-(CYCLOHEXYLMETHYL)-2-ETHYL-, CIS-  
 9 21386 2(1H)-BENZOCYCLOOCTENONE, DECAHYDRO-10A-METHYL-, TRANS-

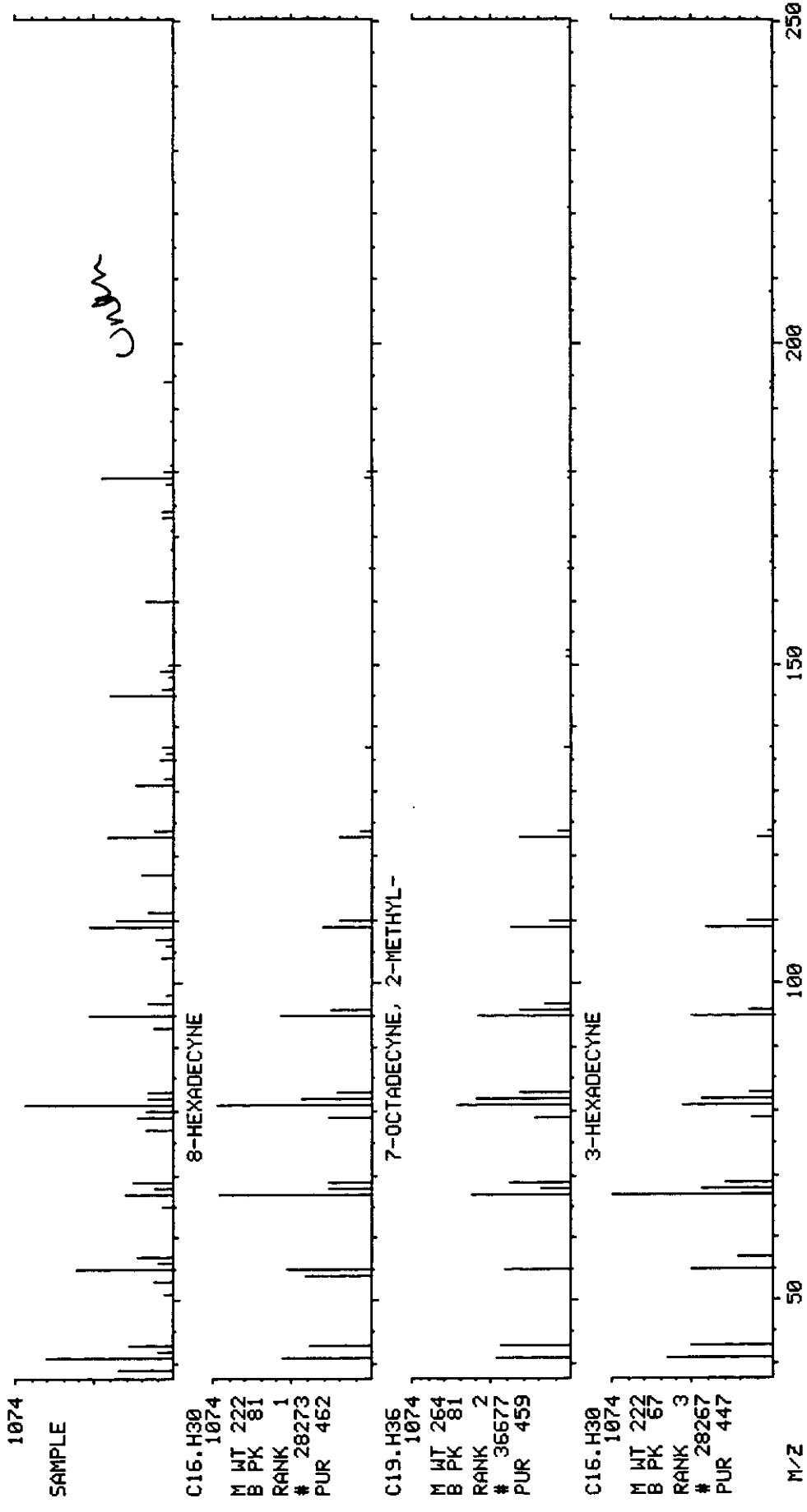
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H30	222	81	462	786	509
2	C19.H36	264	81	459	760	506
3	C16.H30	222	67	447	731	464
4	C18.H34	250	67	434	821	455
5	C19.H36	264	95	433	747	544
6	C11.H20	152	67	425	894	425
7	C15.H28	208	55	424	722	452
8	C15.H28	208	55	420	714	451
9	C13.H22.O	194	55	418	766	459

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	19781-86-3
2	---	---	---	---	35354-38-2
3	---	---	---	---	61886-62-2
4	---	---	---	---	61886-64-4
5	---	---	---	---	55044-33-2
6	---	---	---	---	60212-30-8
7	---	---	---	---	54934-92-8
8	---	---	---	---	54934-93-9
9	---	---	---	---	55103-68-9

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 7:46  
 SAMPLE: 5-MMS-4 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2H 0T)

DATA: 30068110 # 699  
 CALI: 30068110 # 3

BASE M/Z: 81  
 RIC: 23040.



Library Search                    Data: 30068110 # 714            Base m/z: 41  
 08/31/98 23:28:00 + 7:57        Cali: 30068110 # 3            RIC: 36480.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 212 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 21927 TRANS-2-TRIDECENAL  
 2 32052 OXIRANE, TETRADECYL-  
 3 31486 ACETAMIDE, N-METHYL-N-[4-[4-METHOXY-1-HEXAHYDROPYRIDYL]-2-BUTYNYL]-  
 4 35931 HEXADECANE, 1-CHLORO-  
 5 40193 CIS-9,10-EPOXYOCTADECAN-1-OL  
 6 46251 DODECANE, 1,2-DIBROMO-  
 7 40233 NONADECANOL  
 8 42521 1-EICOSANOL  
 9 56499 9-OCTADECENOIC ACID (Z)-, 2-(ACETYLOXY)-1-[(ACETYLOXY)METHYL]ETHYL \*

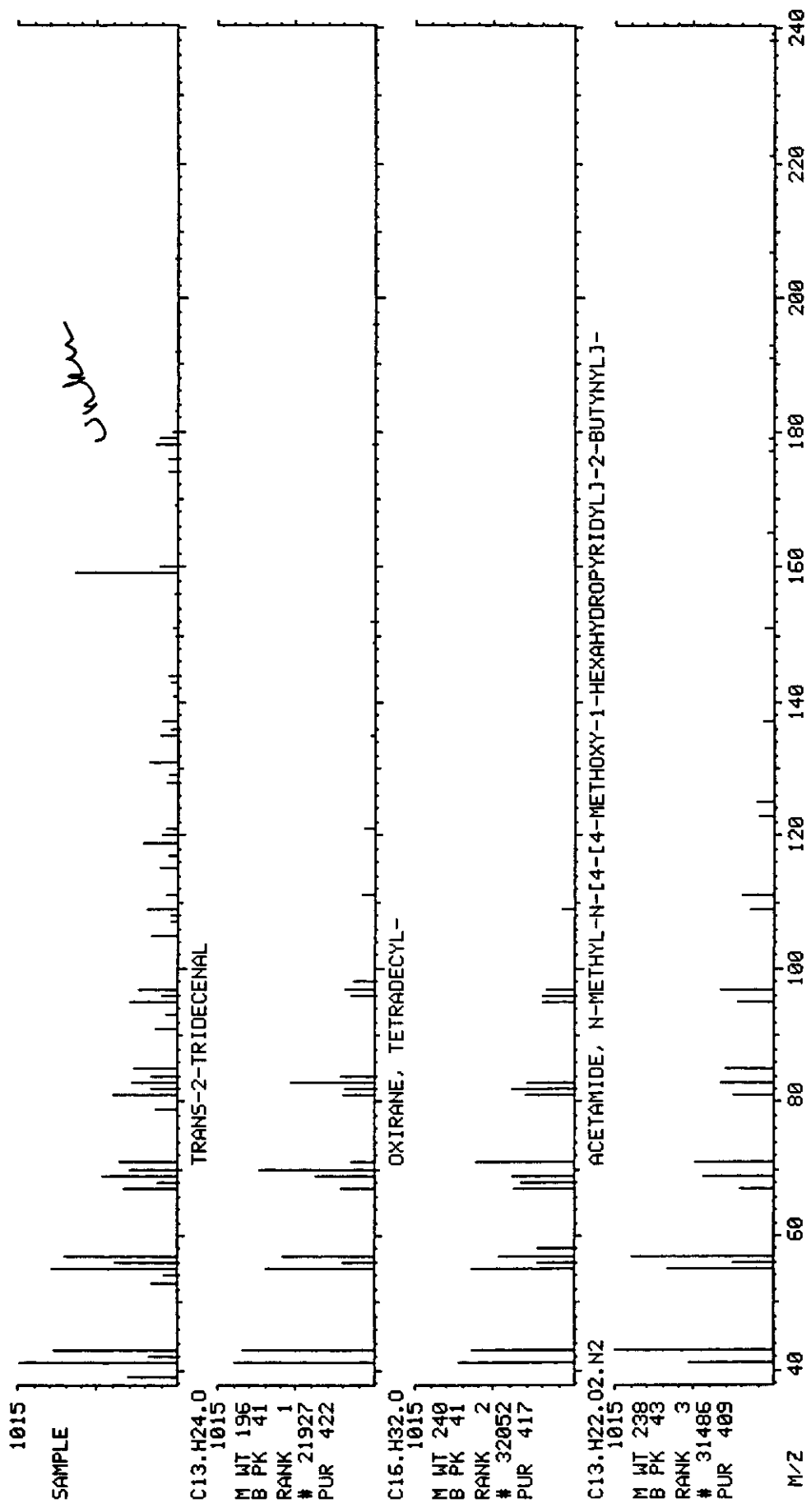
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C13.H24.O	196	41	422	865	444
2	C16.H32.O	240	41	417	918	431
3	C13.H22.O2.N2	238	43	409	829	472
4	C16.H33.CL	260	57	408	921	419
5	C18.H36.O2	284	55	406	883	439
6	C12.H24.BR2	326	41	403	802	482
7	C19.H40.O	284	43	401	898	431
8	C20.H42.O	298	43	398	892	430
9	C25.H44.O6	440	43	394	731	497

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	- - -
2	---	---	---	---	7320-37-8
3	---	---	---	---	- - -
4	---	---	---	---	4860-03-1
5	---	---	---	---	13980-12-6
6	---	---	---	---	55334-42-4
7	---	---	---	---	52783-43-4
8	---	---	---	---	629-96-9
9	---	---	---	---	55401-63-3

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 7:57  
 SAMPLE: S-MM5-4 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA) /1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068110 # 714  
 CALI: 30068110 # 3

BASE M/Z: 41  
 RIC: 36480.



14  
 C2  
 C1  
 C0

Library Search                    Data: 30068110 # 737            Base m/z: 57  
 08/31/98 23:28:00 + 8:12        Cali: 30068110 # 3            RIC: 99328.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 712 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 37462 HEPTADECANE, 2,6-DIMETHYL-  
 2 25991 DODECANE, 2,6,10-TRIMETHYL-  
 3 22535 DODECANE, 4,6-DIMETHYL-  
 4 26001 DODECANE, 2,7,10-TRIMETHYL-  
 5 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 6 25994 DODECANE, 2,6,11-TRIMETHYL-  
 7 15352 UNDECANE, 2-METHYL-  
 8 37465 NONADECANE  
 9 18998 UNDECANE, 3,7-DIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	832	953	849
2	C15.H32	212	57	825	976	830
3	C14.H30	198	57	820	959	821
4	C15.H32	212	57	816	981	823
5	C21.H44	296	57	814	932	853
6	C15.H32	212	57	805	969	814
7	C12.H26	170	43	797	947	801
8	C19.H40	268	57	795	917	837
9	C13.H28	184	43	794	940	796

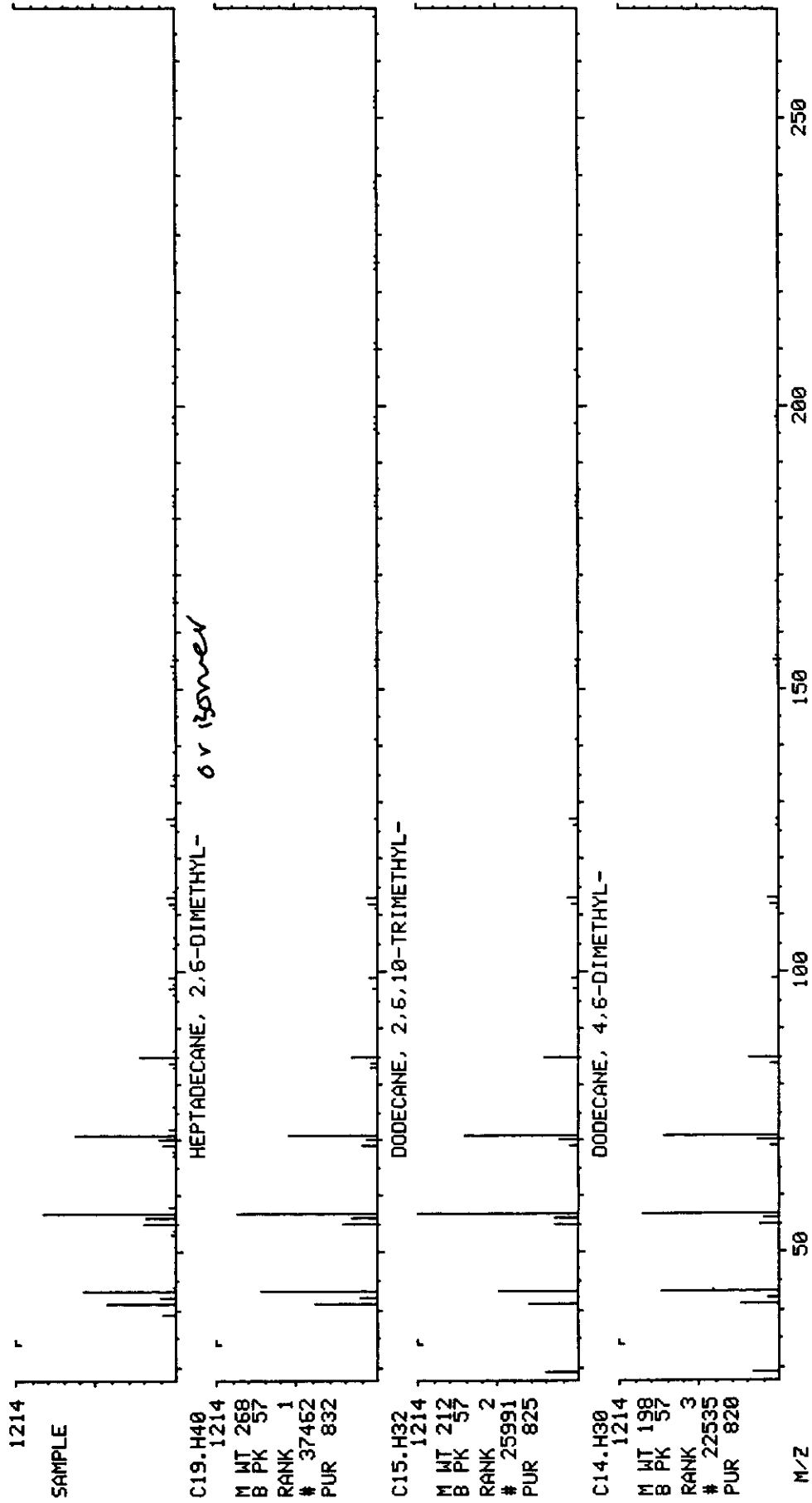
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	54105-67-8
2	---	---	---	---	3891-98-3
3	---	---	---	---	61141-72-8
4	---	---	---	---	74645-98-0
5	---	---	---	---	18344-37-1
6	---	---	---	---	31295-56-4
7	---	---	---	---	7045-71-8
8	---	---	---	---	629-92-5
9	---	---	---	---	17301-29-0



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 8:12  
 SAMPLE: S-MM5-4 1/35A/100M  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068110 # 737  
 CALI: 30068110 # 3

BASE M/Z: 57  
 RIC: 99328.



Library Search                    Data: 30068110 # 763            Base m/z: 43  
 08/31/98 23:28:00 + 8:29        Cali: 30068110 # 3            RIC: 140800.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

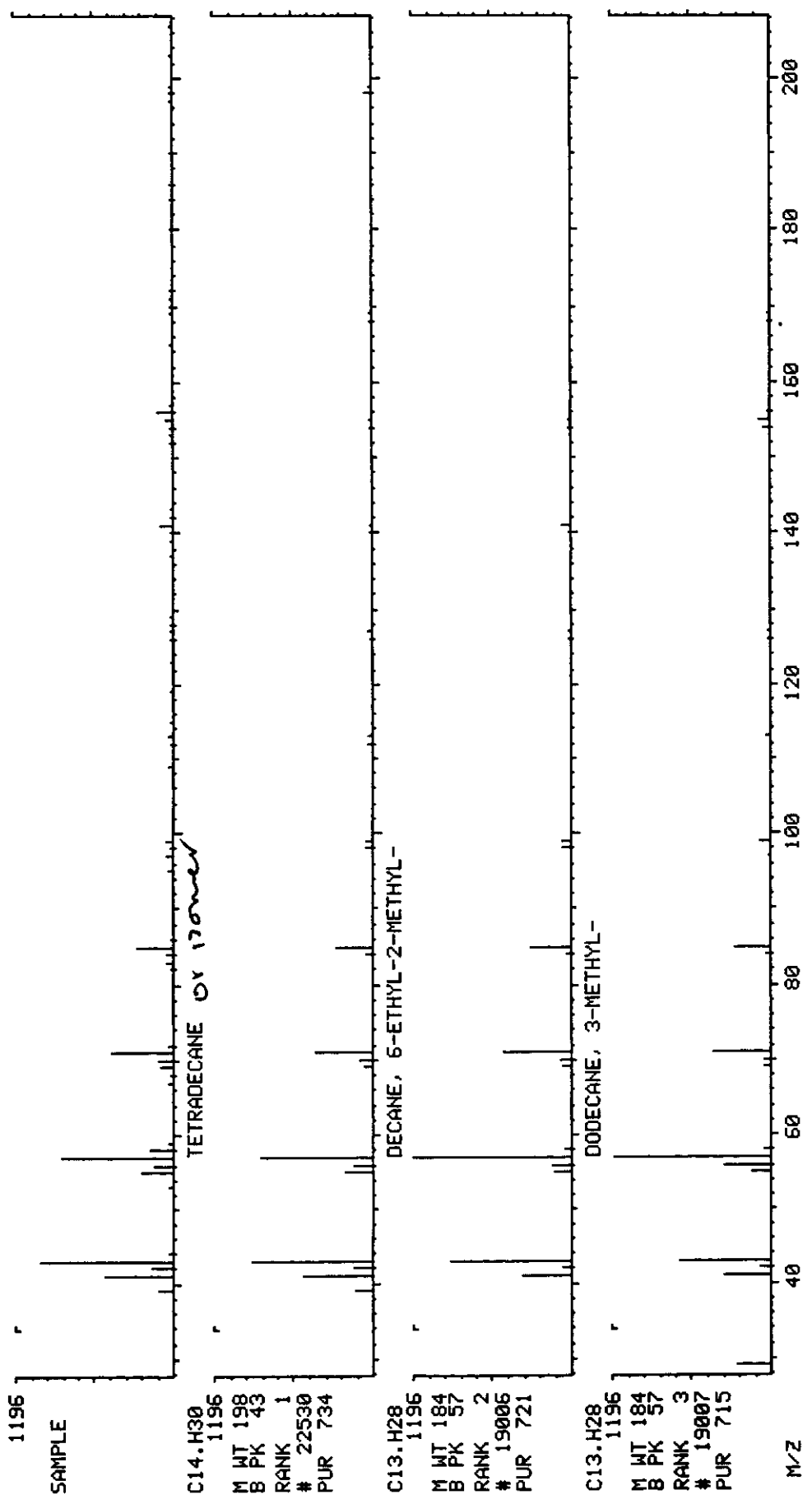
62231 spectra in LIBRARYNB searched for maximum PURITY  
 611 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.            Name  
 1 22530 TETRADECANE  
 2 19006 DECANE, 6-ETHYL-2-METHYL-  
 3 19007 DODECANE, 3-METHYL-  
 4 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 5 15358 UNDECANE, 3-METHYL-  
 6 11607 UNDECANE  
 7 39681 1-iodoundecane  
 8 11612 NONANE, 2,5-DIMETHYL-  
 9 37252 DECANE, 1-iodo-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C14.H30	198	43	734	960	736
2	C13.H28	184	57	721	951	721
3	C13.H28	184	57	715	908	716
4	C21.H44	296	57	709	928	736
5	C12.H26	170	57	706	919	709
6	C11.H24	156	43	705	977	705
7	C11.H23.I	282	57	704	889	710
8	C11.H24	156	57	701	871	701
9	C10.H21.I	268	57	694	911	700

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-59-4
2	---	---	---	---	62108-21-8
3	---	---	---	---	17312-57-1
4	---	---	---	---	18344-37-1
5	---	---	---	---	1002-43-3
6	---	---	---	---	1120-21-4
7	---	---	---	---	4282-44-4
8	---	---	---	---	17302-27-1
9	---	---	---	---	2050-77-3

MID LIBRARY SEARCH <LIBRARYNB> DATA: 30068110 # 763 BASE M/Z: 43  
 08/31/98 23:28:00 + 8:29 CALI: 30068110 # 3 RIC: 140800.  
 SAMPLE: S-MM5-4 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*KNA/NA >1/35A NA M  
 ENHANCED <S 15B 2N 0T>



1300

Library Search                      Data: 30068110 # 776                      Base m/z: 41  
 08/31/98 23:28:00 + 8:38            Cali: 30068110 # 3                      RIC: 31936.  
 Sample: S-MM5-4 1/3SA/100M            INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 677 matched at least 6 of the 16 largest peaks in the unknown

- Rank In.            Name  
 1 40193 CIS-9,10-EPOXYOCTADECAN-1-OL  
 2 32052 OXIRANE, TETRADECYL-  
 3 25971 OXIRANE, DODECYL-  
 4 42521 1-EICOSANOL  
 5 37449 OCTADECANAL  
 6 40233 NONADECANOL  
 7 37444 OXIRANE, HEXADECYL-  
 8 29226 PENTADECANAL-  
 9 29187 BUTANOIC ACID, 3,7-DIMETHYL-6-OCTENYL ESTER

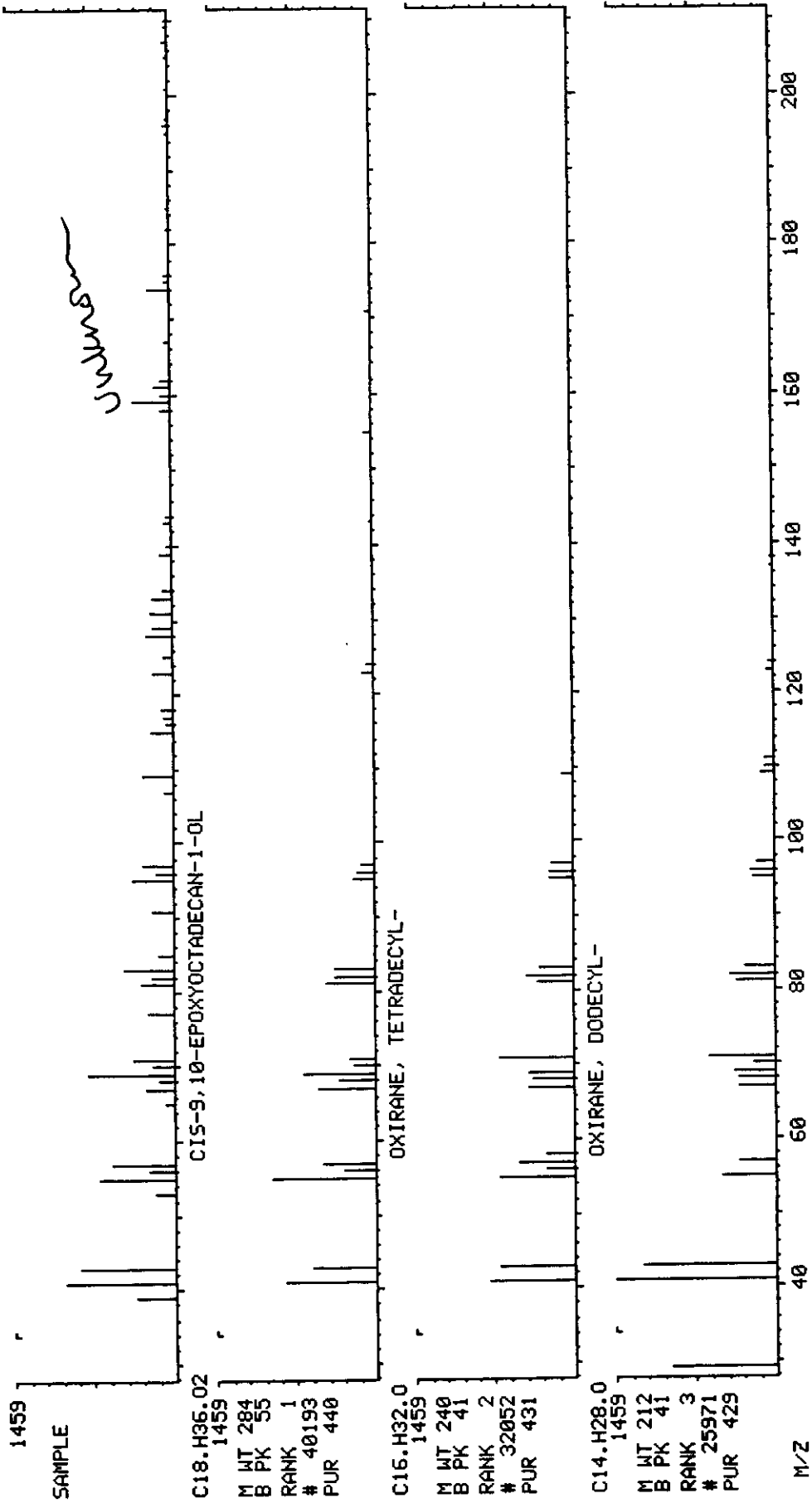
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C18.H36.O2	284	55	440	902	464
2	C16.H32.O	240	41	431	915	446
3	C14.H28.O	212	41	429	870	460
4	C20.H42.O	298	43	409	886	440
5	C18.H36.O	268	43	404	813	465
6	C19.H40.O	284	43	401	897	432
7	C18.H36.O	268	57	398	793	467
8	C15.H30.O	226	82	393	860	415
9	C14.H26.O2	226	41	389	865	410

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	13980-12-6
2	---	---	---	---	7320-37-8
3	---	---	---	---	3234-28-4
4	---	---	---	---	629-96-9
5	---	---	---	---	638-66-4
6	---	---	---	---	52783-43-4
7	---	---	---	---	7390-81-0
8	---	---	---	---	2765-11-9
9	---	---	---	---	141-16-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 8:38  
 SAMPLE: 5-NMS-4 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068110 # 776  
 CALI: 30068110 # 3

BASE M/Z: 41  
 RIC: 31936.



13  
 05  
 01

Library Search                    Data: 30068110 # 820            Base m/z: 57  
 08/31/98 23:28:00 + 9:07       Cali: 30068110 # 3            RIC: 91136.  
 Sample: S-MM5-4 1/3SA/100M       INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 760 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 2 37462 HEPTADECANE, 2,6-DIMETHYL-  
 3 18987 UNDECANE, 2,8-DIMETHYL-  
 4 15344 DECANE, 3,6-DIMETHYL-  
 5 8104 OCTANE, 3,5-DIMETHYL-  
 6 19004 UNDECANE, 4,6-DIMETHYL-  
 7 19023 DECANE, 2,6,7-TRIMETHYL-  
 8 19039 DECANE, 2,3,5-TRIMETHYL-  
 9 15358 UNDECANE, 3-METHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C21.H44	296	57	792	957	816
2	C19.H40	268	57	776	932	805
3	C13.H28	184	43	762	955	768
4	C12.H26	170	57	757	952	764
5	C10.H22	142	57	755	963	755
6	C13.H28	184	57	755	955	757
7	C13.H28	184	57	752	937	752
8	C13.H28	184	57	751	942	752
9	C12.H26	170	57	751	938	756

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	18344-37-1
2	---	---	---	---	54105-67-8
3	---	---	---	---	17301-25-6
4	---	---	---	---	17312-53-7
5	---	---	---	---	15869-93-9
6	---	---	---	---	17312-82-2
7	---	---	---	---	62108-25-2
8	---	---	---	---	62238-11-3
9	---	---	---	---	1002-43-3

DATA: 30058110 # 820  
CALI: 30058110 # 3

BASE M/Z: 57  
RIC: 91136.

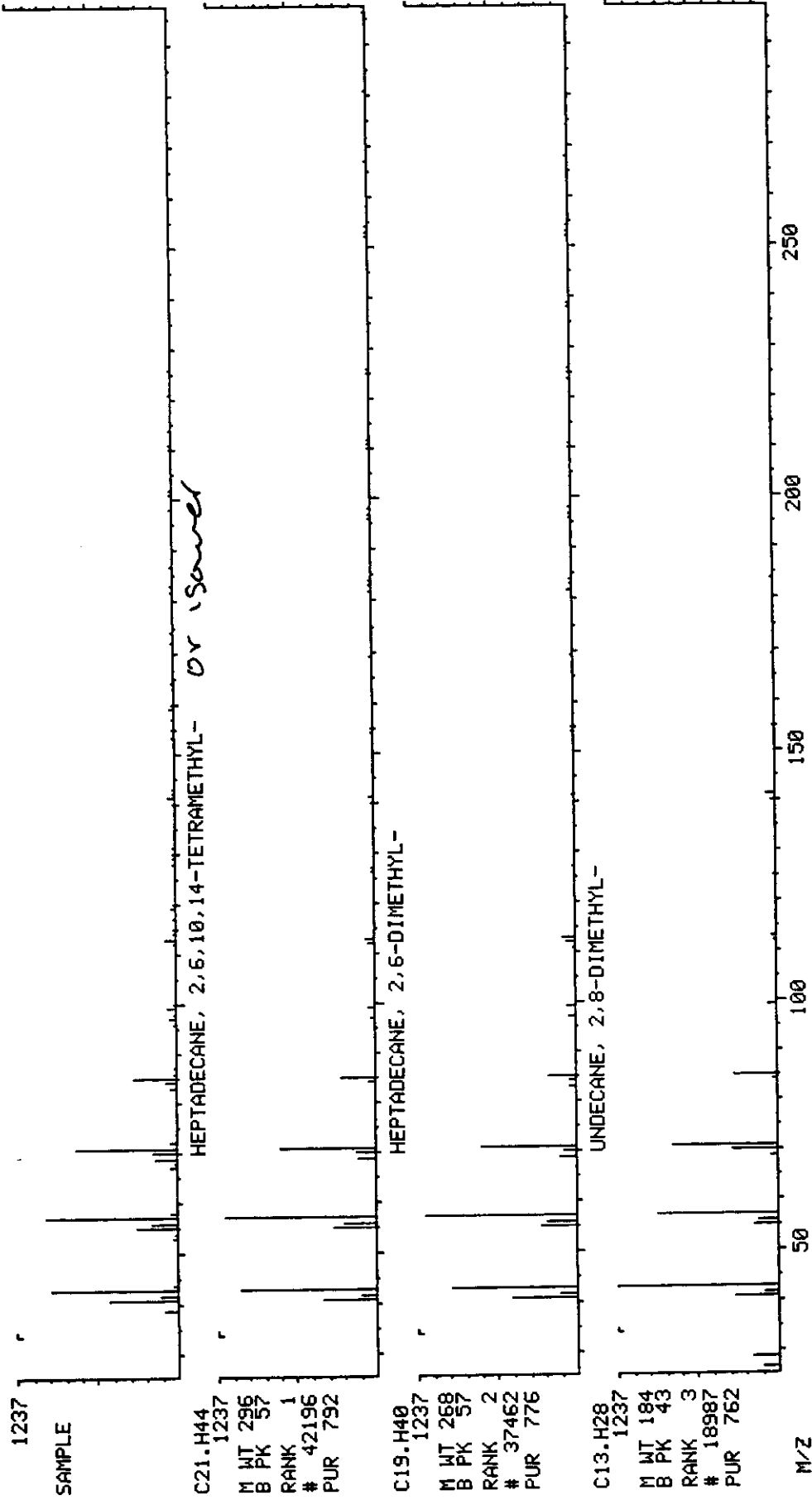
MID LIBRARY SEARCH (LIBRARY#)

08/31/98 23:28:00 + 9:07

SAMPLE: S-MM5-4 1/35A/100M

COND5.: UG/ML \*100ML \*100% /100% \*(NA/NA) /1/35A NA M  
ENHANCED (S 15B 2N 0T)

INST. ID: F16



1237

SAMPLE

C21.H44

1237

M WT 296

B PK 57

RANK 1

# 42196

PUR 792

C19.H40

1237

M WT 268

B PK 57

RANK 2

# 37462

PUR 776

C13.H28

1237

M WT 184

B PK 43

RANK 3

# 18987

PUR 762

M/Z

Library Search                    Data: 30068110 # 857            Base m/z: 57  
 08/31/98 23:28:00 + 9:32        Cali: 30068110 # 3            RIC: 132352.  
 Sample: S-MMS-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N DT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 242 matched at least 8 of the 16 largest peaks in the unknown

- | Rank In. | Name                                      |
|----------|---|
| 1        | 25997 PENTADECANE                         |
| 2        | 19007 DODECANE, 3-METHYL-                 |
| 3        | 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL- |
| 4        | 37465 NONADECANE                          |
| 5        | 39681 1-IODOUNDECANE                      |
| 6        | 18990 UNDECANE, 2,9-DIMETHYL-             |
| 7        | 19016 UNDECANE, 4,7-DIMETHYL-             |
| 8        | 19026 DECANE, 2,6,8-TRIMETHYL-            |
| 9        | 37252 DECANE, 1-IODO-                     |

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C15.H32	212	57	770	955	773
2	C13.H28	184	57	767	961	769
3	C21.H44	296	57	761	941	778
4	C19.H40	268	57	753	926	783
5	C11.H23.I	282	57	748	970	754
6	C13.H28	184	57	747	954	749
7	C13.H28	184	43	744	936	744
8	C13.H28	184	57	741	939	744
9	C10.H21.I	268	57	738	972	743

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-62-9
2	---	---	---	---	17312-57-1
3	---	---	---	---	18344-37-1
4	---	---	---	---	629-92-5
5	---	---	---	---	4282-44-4
6	---	---	---	---	17301-26-7
7	---	---	---	---	17301-32-5
8	---	---	---	---	62108-26-3
9	---	---	---	---	2050-77-3

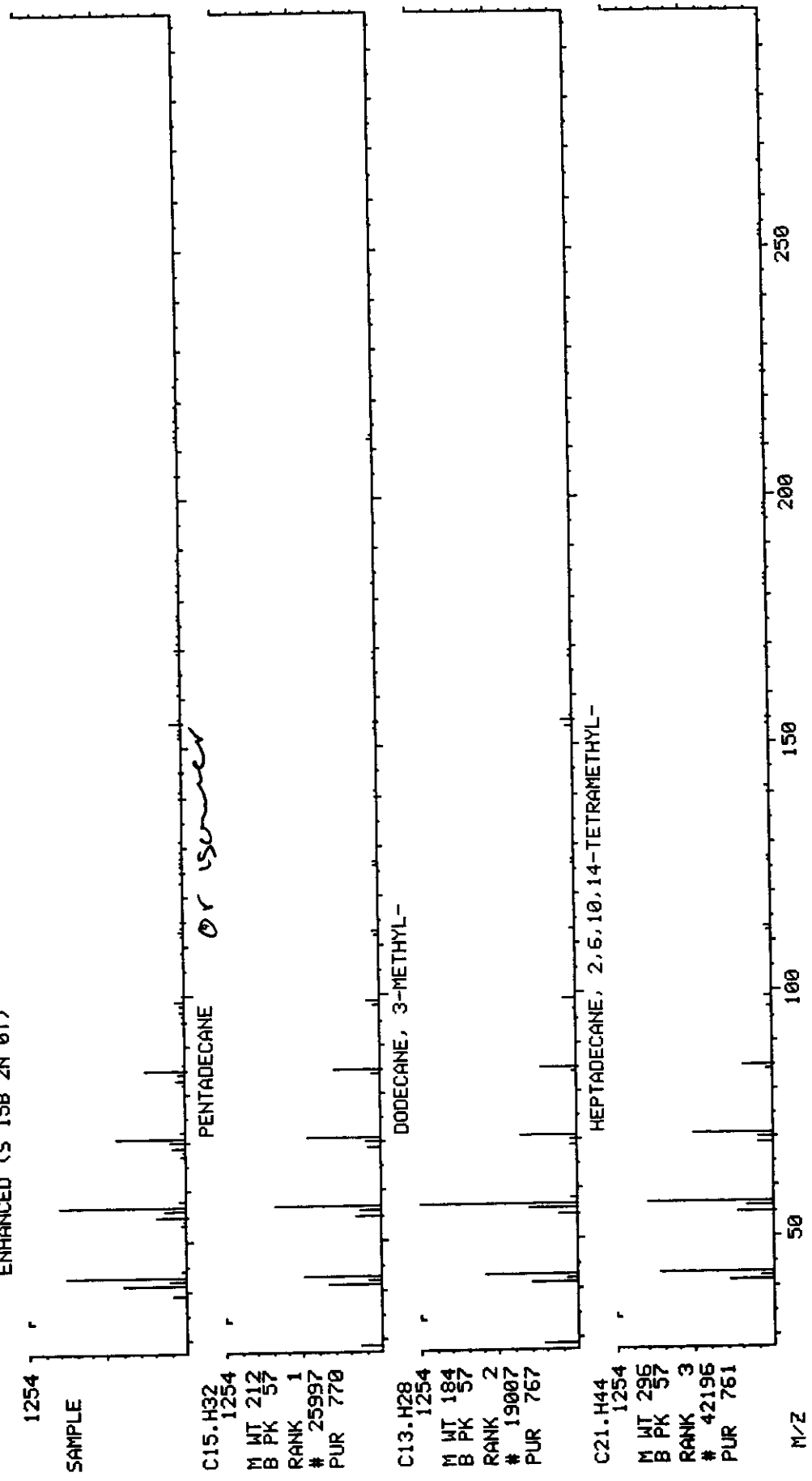


BASE M/Z: 57  
RIC: 132352.

DATA: 30068110 # 857  
CALI: 30068110 # 3

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:28:00 + 9:32  
SAMPLE: 5-MM5-4 1/35A/100M  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 15B 2N 0T)

INST. ID: F16



13000

Library Search                      Data: 30068110 # 903                      Base m/z: 41  
 08/31/98 23:28:00 + 10:03                      Cali: 30068110 # 3                      RIC: 46592.  
 Sample: S-MM5-4 1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

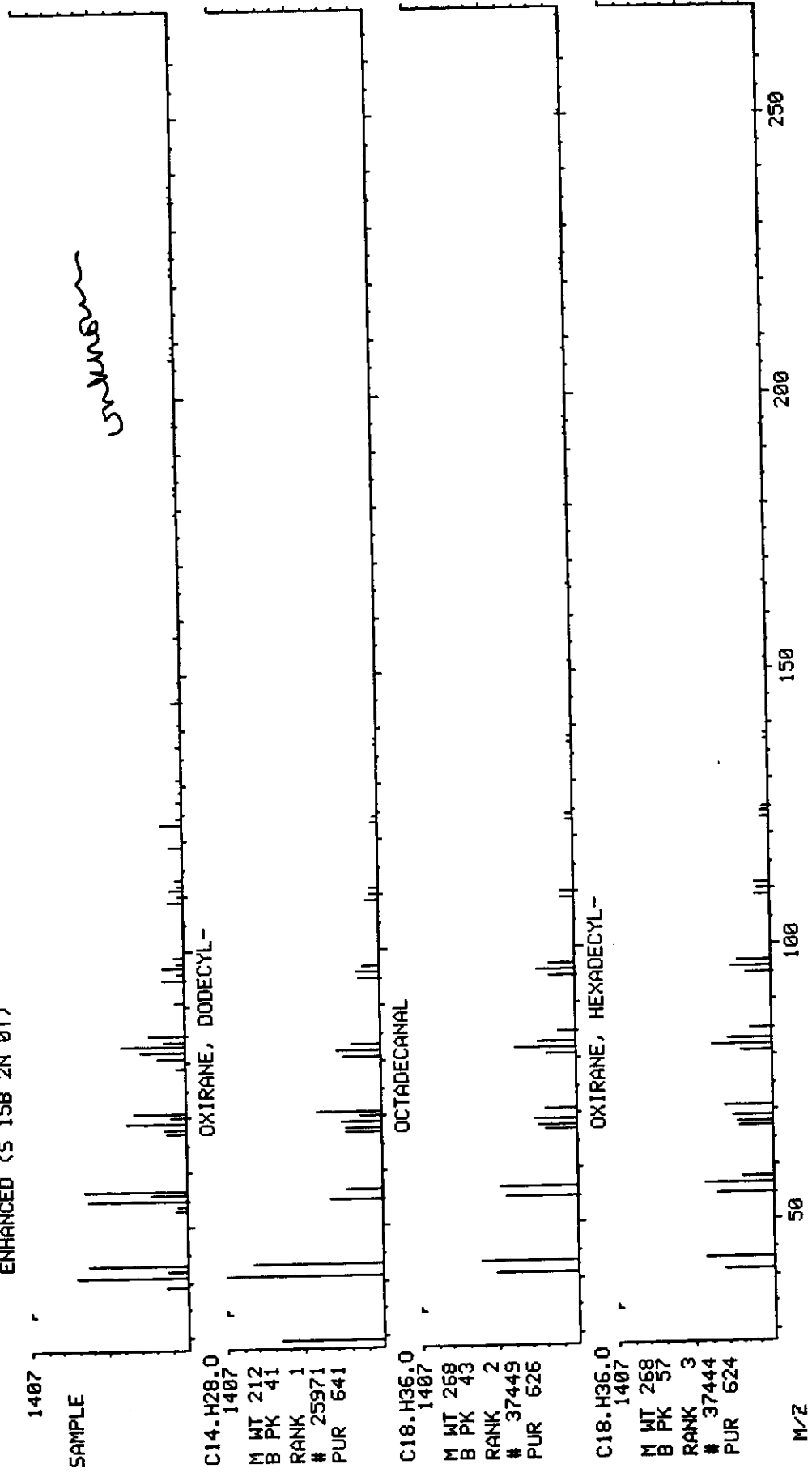
62231 spectra in LIBRARYNB searched for maximum PURITY  
 606 matched at least 7 of the 16 largest peaks in the unknown

Rank In.                      Name  
 1 25971 OXIRANE, DODECYL-  
 2 37449 OCTADECANAL  
 3 37444 OXIRANE, HEXADECYL-  
 4 40233 NONADECANOL  
 5 42521 1-EICOSANOL  
 6 58380 DECANEDIOIC ACID, DIDECYL ESTER  
 7 40193 CIS-9,10-EPOXYOCTADECAN-1-OL  
 8 42180 OCTADECANE, 1-(ETHENYLOXY)-  
 9 15969 HYDROXYLAMINE, O-DECYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C14.H28.O	212	41	641	918	646
2	C18.H36.O	268	43	626	872	669
3	C18.H36.O	268	57	624	855	675
4	C19.H40.O	284	43	612	926	651
5	C20.H42.O	298	43	609	920	653
6	C30.H58.O4	482	57	603	923	612
7	C18.H36.O2	284	55	602	898	631
8	C20.H40.O	296	43	599	930	633
9	C10.H23.O.N	173	43	592	891	600

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	3234-28-4
2	---	---	---	---	638-66-4
3	---	---	---	---	7390-81-0
4	---	---	---	---	52783-43-4
5	---	---	---	---	629-96-9
6	---	---	---	---	2432-89-5
7	---	---	---	---	13980-12-6
8	---	---	---	---	930-02-9
9	---	---	---	---	29812-79-1

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:28:00 + 10:03  
 SAMPLE: S-MM5-4 1/35A/100M  
 CONDS.: UC/ML \*100ML \*100%/100% \*(NA/NA) >/1/35A NA M  
 ENHANCED (S 15B 2N 0T)  
 DATA: 30068110 # 903  
 CALI: 30068110 # 3  
 BASE M/Z: 41  
 RIC: 46592.



1371

Library Search                    Data: 30068110 # 947            Base m/z: 57  
 08/31/98 23:28:00 + 10:32        Cali: 30068110 # 3            RIC: 123136.  
 Sample: S-MM5-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 343 matched at least 8 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 37465 NONADECANE  
 2 37462 HEPTADECANE, 2,6-DIMETHYL-  
 3 25997 PENTADECANE  
 4 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 5 15969 HYDROXYLAMINE, O-DECYL-  
 6 19016 UNDECANE, 4,7-DIMETHYL-  
 7 25994 DODECANE, 2,6,11-TRIMETHYL-  
 8 19523 1-DECANOL, 2-ETHYL-  
 9 26001 DODECANE, 2,7,10-TRIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	842	958	864
2	C19.H40	268	57	827	940	854
3	C15.H32	212	57	813	957	826
4	C21.H44	296	57	813	944	838
5	C10.H23.O.N	173	43	802	950	820
6	C13.H28	184	43	802	964	804
7	C15.H32	212	57	796	963	800
8	C12.H26.O	186	57	791	928	804
9	C15.H32	212	57	789	959	794

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-92-5
2	---	---	---	---	54105-67-8
3	---	---	---	---	629-62-9
4	---	---	---	---	18344-37-1
5	---	---	---	---	29812-79-1
6	---	---	---	---	17301-32-5
7	---	---	---	---	31295-56-4
8	---	---	---	---	21078-65-9
9	---	---	---	---	74645-98-0

DATA: 30068110 # 947  
CALI: 30068110 # 3

BASE M/Z: 57  
RIC: 123135.

MID LIBRARY SEARCH <LIBRARYNB>

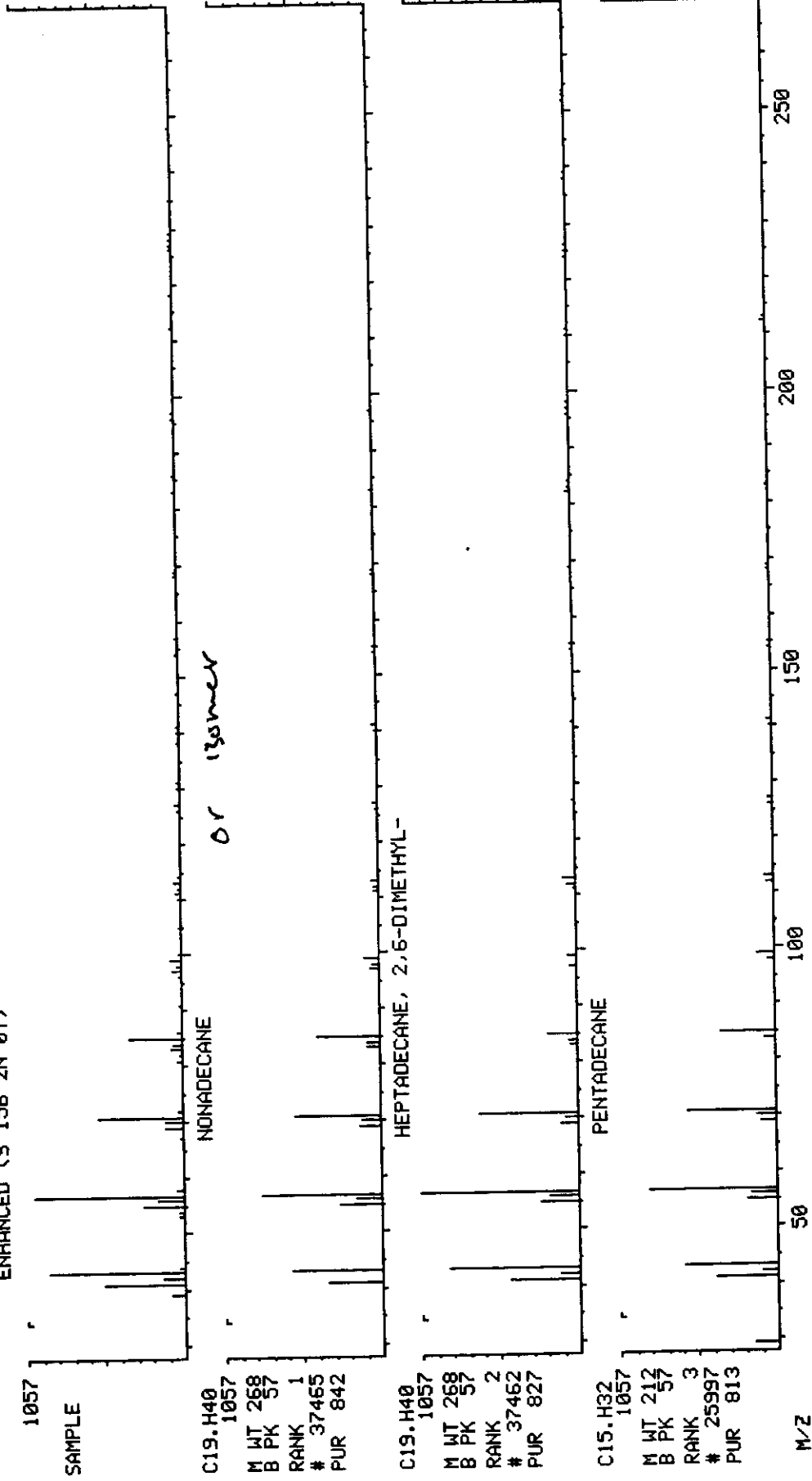
08/31/98 23:28:00 + 10:32

SAMPLE: S-MM5-4 1/35A/100M

COND.S.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M

INST. ID: F16

ENHANCED (S 15B 2N 0T)



Library Search                    Data: 30068110 # 987            Base m/z: 57  
 08/31/98 23:28:00 + 10:59      Cali: 30068110 # 3            RIC: 93568.  
 Sample: S-MM5-4 1/3SA/100M      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 644 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.            Name  
 1 19054 UNDECANE, 2,6-DIMETHYL-  
 2 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 3 37462 HEPTADECANE, 2,6-DIMETHYL-  
 4 25996 TRIDECANE, 4,8-DIMETHYL-  
 5 25997 PENTADECANE  
 6 29263 HEXADECANE  
 7 22530 TETRADECANE  
 8 32059 HEPTADECANE  
 9 19026 DECANE, 2,6,8-TRIMETHYL-

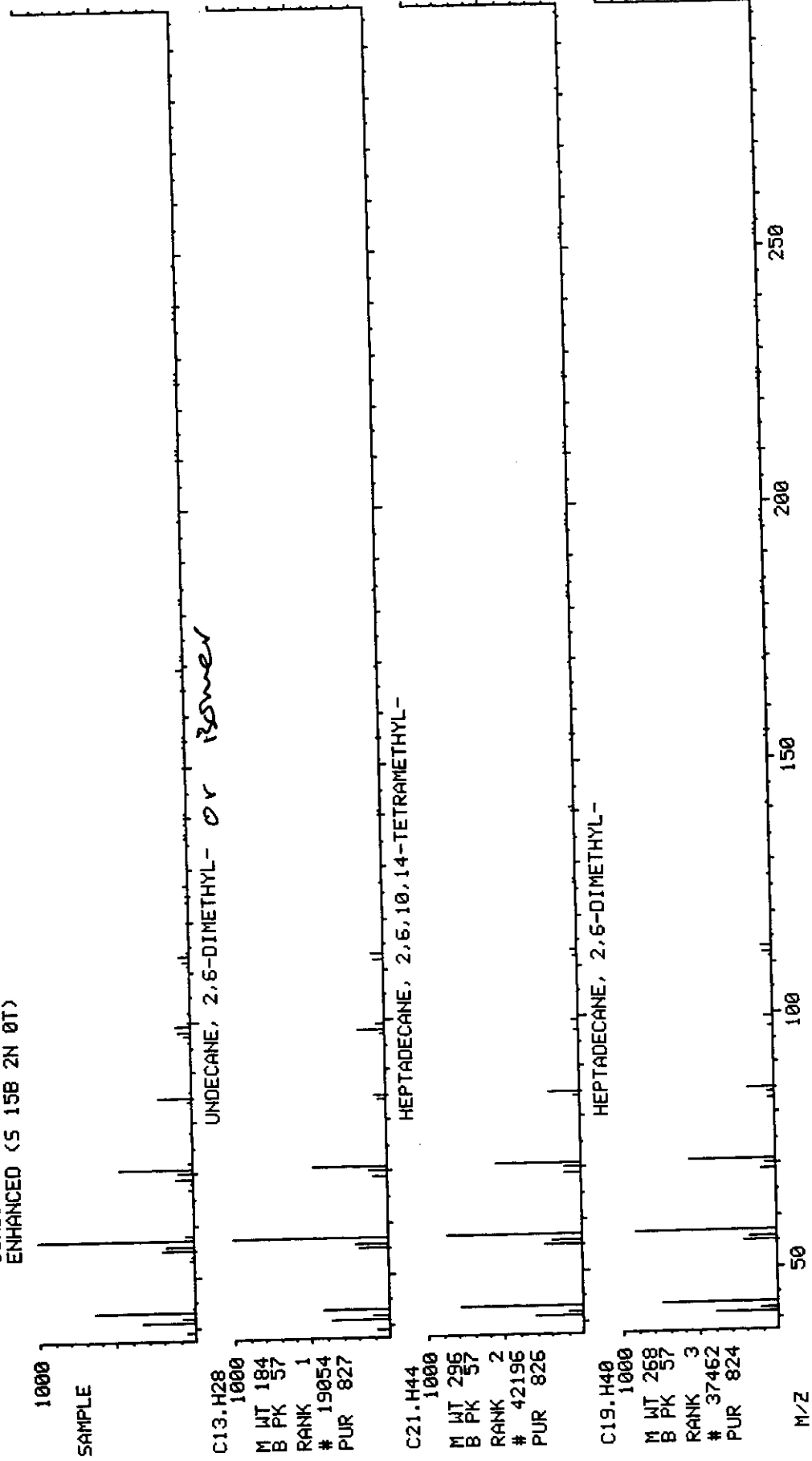
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C13.H28	184	57	827	949	834
2	C21.H44	296	57	826	957	837
3	C19.H40	268	57	824	943	845
4	C15.H32	212	57	819	936	834
5	C15.H32	212	57	819	945	833
6	C16.H34	226	57	816	931	834
7	C14.H30	198	43	812	936	826
8	C17.H36	240	57	810	923	831
9	C13.H28	184	57	805	953	813

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	17301-23-4
2	---	---	---	---	18344-37-1
3	---	---	---	---	54105-67-8
4	---	---	---	---	55030-62-1
5	---	---	---	---	629-62-9
6	---	---	---	---	544-76-3
7	---	---	---	---	629-59-4
8	---	---	---	---	629-78-7
9	---	---	---	---	62108-26-3

BASE M/Z: 57  
RIC: 93568.

DATA: 30068110 # 987  
CALI: 30068110 # 3

MID LIBRARY SEARCH <LIBRARYNB>  
08/31/98 23:28:00 + 10:59  
SAMPLE: S-MMS-4 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 158 2N 0T)



11  
CS  
21  
C1

Library Search                      Data: 30068110 #1035                      Base m/z: 57  
 08/31/98 23:28:00 + 11:31                      Cali: 30068110 # 3                      RIC: 132608.  
 Sample: S-MMS-4 1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 840 matched at least 7 of the 16 largest peaks in the unknown

Rank In.                      Name  
 1 37462 HEPTADECANE, 2,6-DIMETHYL-  
 2 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 3 37465 NONADECANE  
 4 25991 DODECANE, 2,6,10-TRIMETHYL-  
 5 26001 DODECANE, 2,7,10-TRIMETHYL-  
 6 37466 PENTADECANE, 2,6,10,14-TETRAMETHYL-  
 7 22535 DODECANE, 4,6-DIMETHYL-  
 8 25994 DODECANE, 2,6,11-TRIMETHYL-  
 9 29264 TRIDECANE, 5-PROPYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	867	971	875
2	C21.H44	296	57	823	946	848
3	C19.H40	268	57	819	938	837
4	C15.H32	212	57	815	970	817
5	C15.H32	212	57	814	966	818
6	C19.H40	268	71	811	944	830
7	C14.H30	198	57	810	959	811
8	C15.H32	212	57	792	941	801
9	C16.H34	226	57	790	937	831

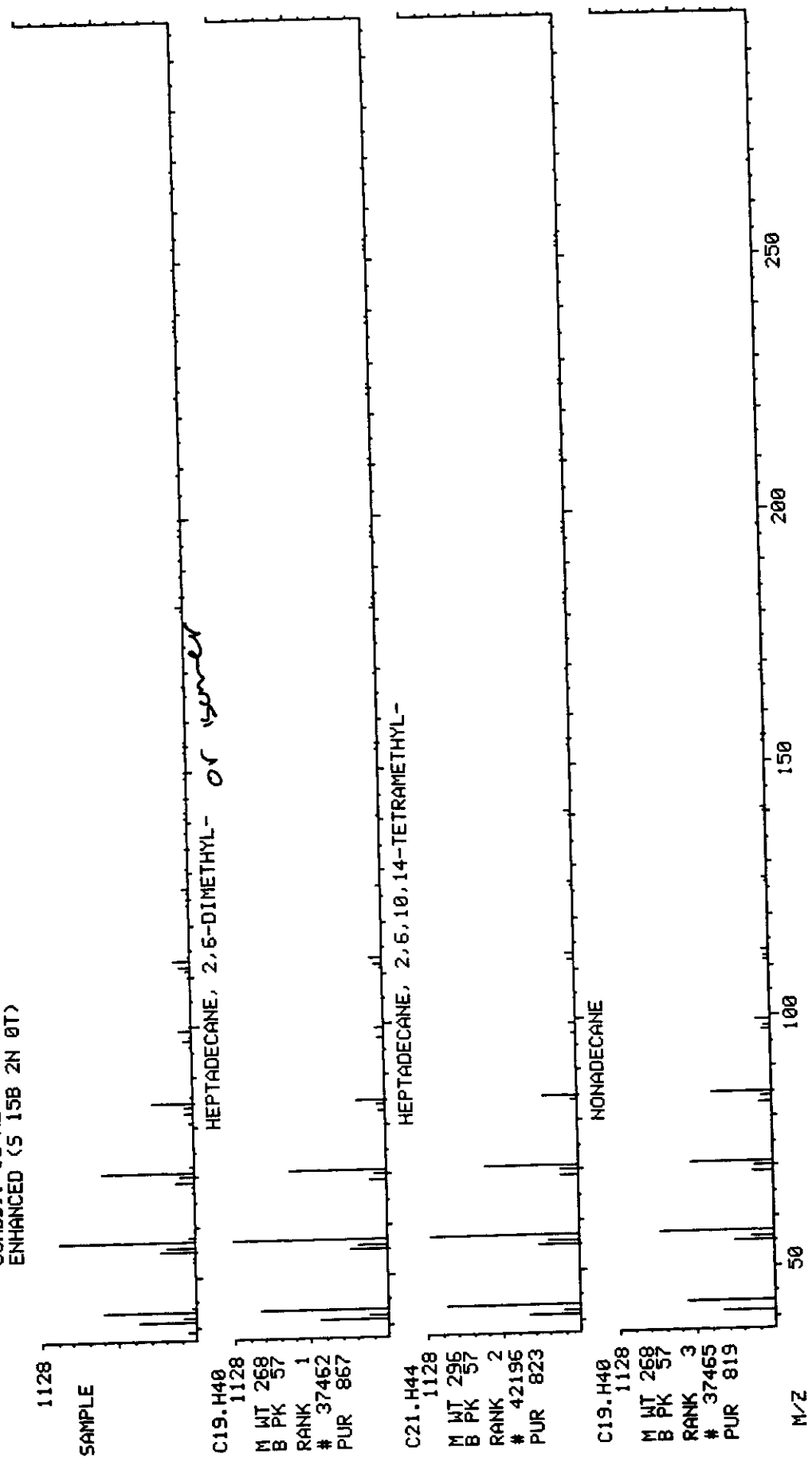
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	54105-67-8
2	---	---	---	---	18344-37-1
3	---	---	---	---	629-92-5
4	---	---	---	---	3891-98-3
5	---	---	---	---	74645-98-0
6	---	---	---	---	1921-70-6
7	---	---	---	---	61141-72-8
8	---	---	---	---	31295-56-4
9	---	---	---	---	55045-11-9



BASE M/Z: 57  
RIC: 132608.

DATA: 30068110 #1035  
CALI: 30068110 # 3

MID LIBRARY SEARCH <LIBRARYNB>  
08/31/98 23:28:00 + 11:31  
SAMPLE: 5-MM5-4 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100ML \*100Z \*(NA/NA) /1/35A NA M  
ENHANCED (S 15B 2N 0T)



1128

SAMPLE

C19.H40  
1128

M WT 268  
B PK 57  
RANK 1  
# 37462  
PUR 867

HEPTADECANE, 2,6-DIMETHYL- or ymer

C21.H44  
1128

M WT 296  
B PK 57  
RANK 2  
# 42196  
PUR 823

HEPTADECANE, 2,6,10,14-TETRAMETHYL-

C19.H40  
1128

M WT 268  
B PK 57  
RANK 3  
# 37465  
PUR 819

NONADECANE

M/Z

1377

Library Search Data: 30068110 #1053 Base m/z: 69  
 08/31/98 23:28:00 + 11:43 Cali: 30068110 # 3 RIC: 38336.  
 Sample: S-MM5-4 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 725 matched at least 7 of the 16 largest peaks in the unknown

Rank	In.	Name
1	34410	3-OCTADECENE, (E)-
2	40233	NONADECANOL
3	34418	9-OCTADECENE, (E)-
4	28768	3-HEXADECENE, (Z)-
5	34411	5-OCTADECENE, (E)-
6	26409	2-HEXYL-1-OCTANOL
7	28772	7-HEXADECENE, (Z)-
8	21963	3-TETRADECENE, (E)-
9	21962	3-TETRADECENE, (Z)-

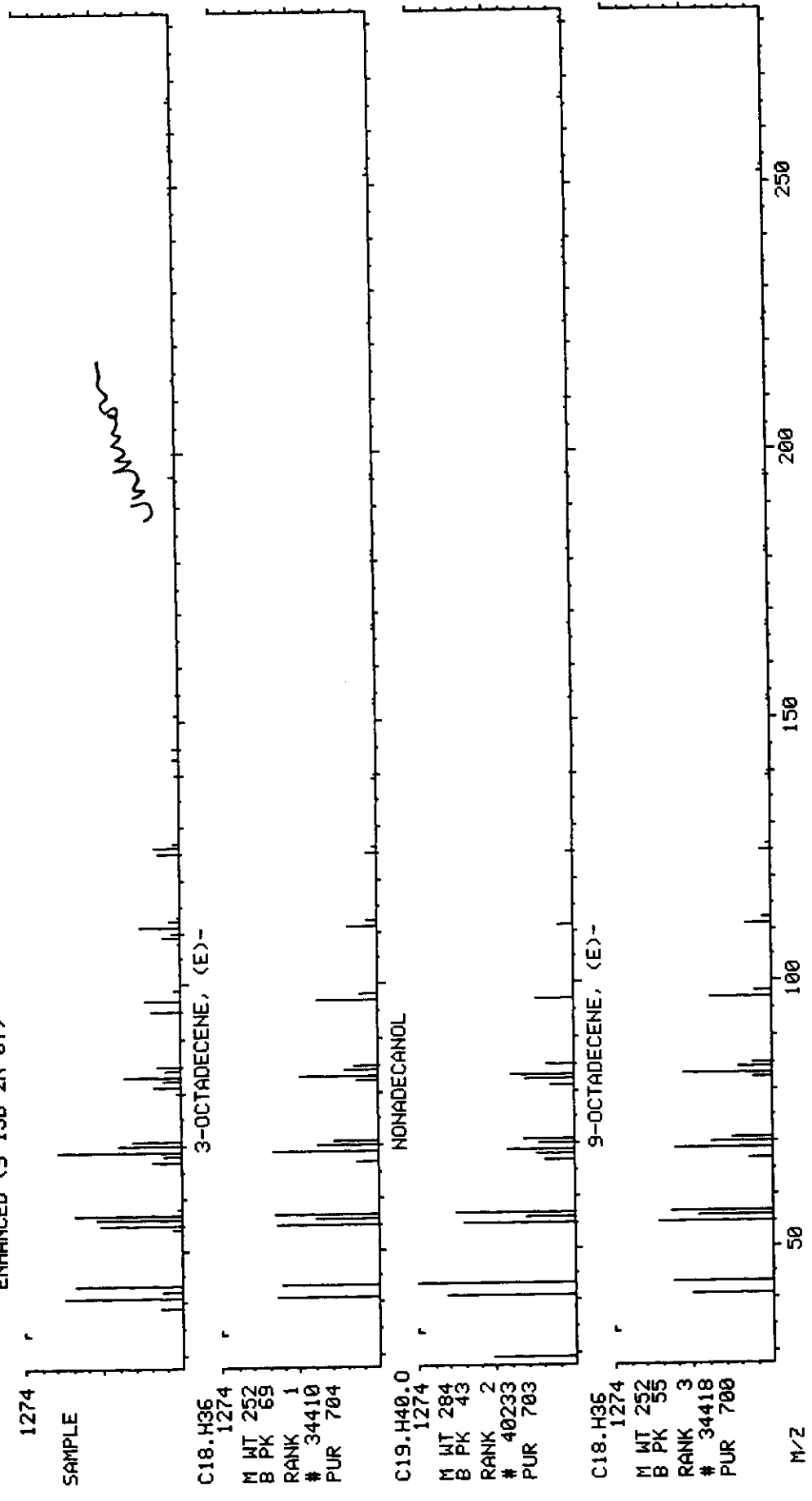
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C18.H36	252	69	704	901	742
2	C19.H40.O	284	43	703	908	726
3	C18.H36	252	55	700	895	734
4	C16.H32	224	55	699	896	746
5	C18.H36	252	55	698	893	733
6	C14.H30.O	214	57	695	912	725
7	C16.H32	224	55	691	885	734
8	C14.H28	196	41	687	911	712
9	C14.H28	196	41	685	909	707

Rank	Ret.Time	B.P.int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	7206-19-1
2	—	—	—	—	52783-43-4
3	—	—	—	—	7206-25-9
4	—	—	—	—	34303-81-6
5	—	—	—	—	7206-21-5
6	—	—	—	—	- -
7	—	—	—	—	35507-09-6
8	—	—	—	—	41446-68-8
9	—	—	—	—	41446-67-7

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:28:00 + 11:43  
 SAMPLE: S-MM5-4 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100% \*100% \*(NA/NA) >1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068110 #1053  
 CALI: 30068110 # 3

BASE M/Z: 69  
 RIC: 36336.



Library Search                      Data: 30068110 #1113                      Base m/z: 57  
 08/31/98 23:28:00 + 12:23                      Cali: 30068110 # 3                      RIC: 55040.  
 Sample: S-MM5-4 1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 938 matched at least 7 of the 16 largest peaks in the unknown

- | Rank In. | Name                                      |
|----------|---|
| 1        | 15969 HYDROXYLAMINE, O-DECYL-             |
| 2        | 15358 UNDECANE, 3-METHYL-                 |
| 3        | 37462 HEPTADECANE, 2,6-DIMETHYL-          |
| 4        | 22536 TRIDECANE, 3-METHYL-                |
| 5        | 37465 NONADECANE                          |
| 6        | 22530 TETRADECANE                         |
| 7        | 19016 UNDECANE, 4,7-DIMETHYL-             |
| 8        | 25997 PENTADECANE                         |
| 9        | 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL- |

Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C10.H23.O.N	173	43	794	948	813
2	C12.H26	170	57	789	943	792
3	C19.H40	268	57	782	913	827
4	C14.H30	198	57	778	932	791
5	C19.H40	268	57	775	919	829
6	C14.H30	198	43	772	930	821
7	C13.H28	184	43	771	954	778
8	C15.H32	212	57	766	933	809
9	C21.H44	296	57	762	929	802

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	29812-79-1
2	---	---	---	---	1002-43-3
3	---	---	---	---	54105-67-8
4	---	---	---	---	6418-41-3
5	---	---	---	---	629-92-5
6	---	---	---	---	629-59-4
7	---	---	---	---	17301-32-5
8	---	---	---	---	629-62-9
9	---	---	---	---	18344-37-1

1380

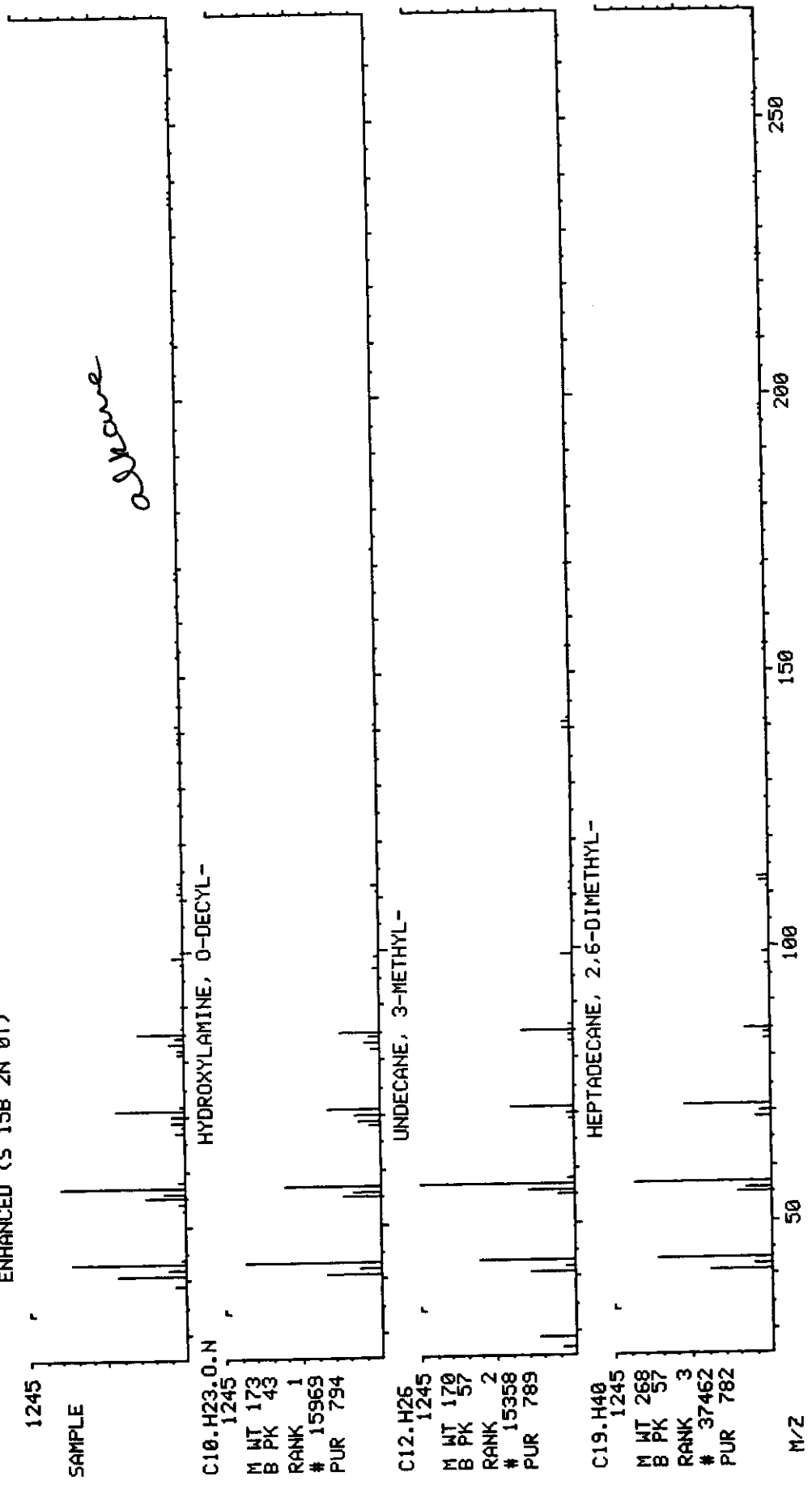
*[Handwritten signature]*  
 11/17/98

BASE M/Z: 57  
RIC: 55040.

DATA: 30068110 #1113  
CALI: 30068110 # 3

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:28:00 + 12:23  
SAMPLE: S-MMS-4 1/35A/100M  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 15B 2N 0T)

INST. ID: F16



Library Search                    Data: 30068110 #1118            Base m/z: 57  
 08/31/98 23:28:00 + 12:26        Cali: 30068110 # 3            RIC: 73216.  
 Sample: S-MMS-4 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 590 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 37462 HEPTADECANE, 2,6-DIMETHYL-  
 2 37465 NONADECANE  
 3 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 4 46161 TRICOSANE  
 5 32059 HEPTADECANE  
 6 25991 DODECANE, 2,6,10-TRIMETHYL-  
 7 37466 PENTADECANE, 2,6,10,14-TETRAMETHYL-  
 8 42192 HEPTADECANE, 2,6,10,15-TETRAMETHYL-  
 9 26001 DODECANE, 2,7,10-TRIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	834	941	842
2	C19.H40	268	57	833	957	853
3	C21.H44	296	57	823	935	838
4	C23.H48	324	43	813	915	860
5	C17.H36	240	57	802	927	836
6	C15.H32	212	57	802	962	804
7	C19.H40	268	71	801	930	814
8	C21.H44	296	57	801	952	828
9	C15.H32	212	57	800	963	800

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	54105-67-8
2	—	—	—	—	629-92-5
3	—	—	—	—	18344-37-1
4	—	—	—	—	638-67-5
5	—	—	—	—	629-78-7
6	—	—	—	—	3891-98-3
7	—	—	—	—	1921-70-6
8	—	—	—	—	54833-48-6
9	—	—	—	—	74645-98-0

BASE M/Z: 57  
RIC: 73216.

DATA: 30068110 #1118  
CALI: 30068110 # 3

MID LIBRARY SEARCH <LIBRARYNB>

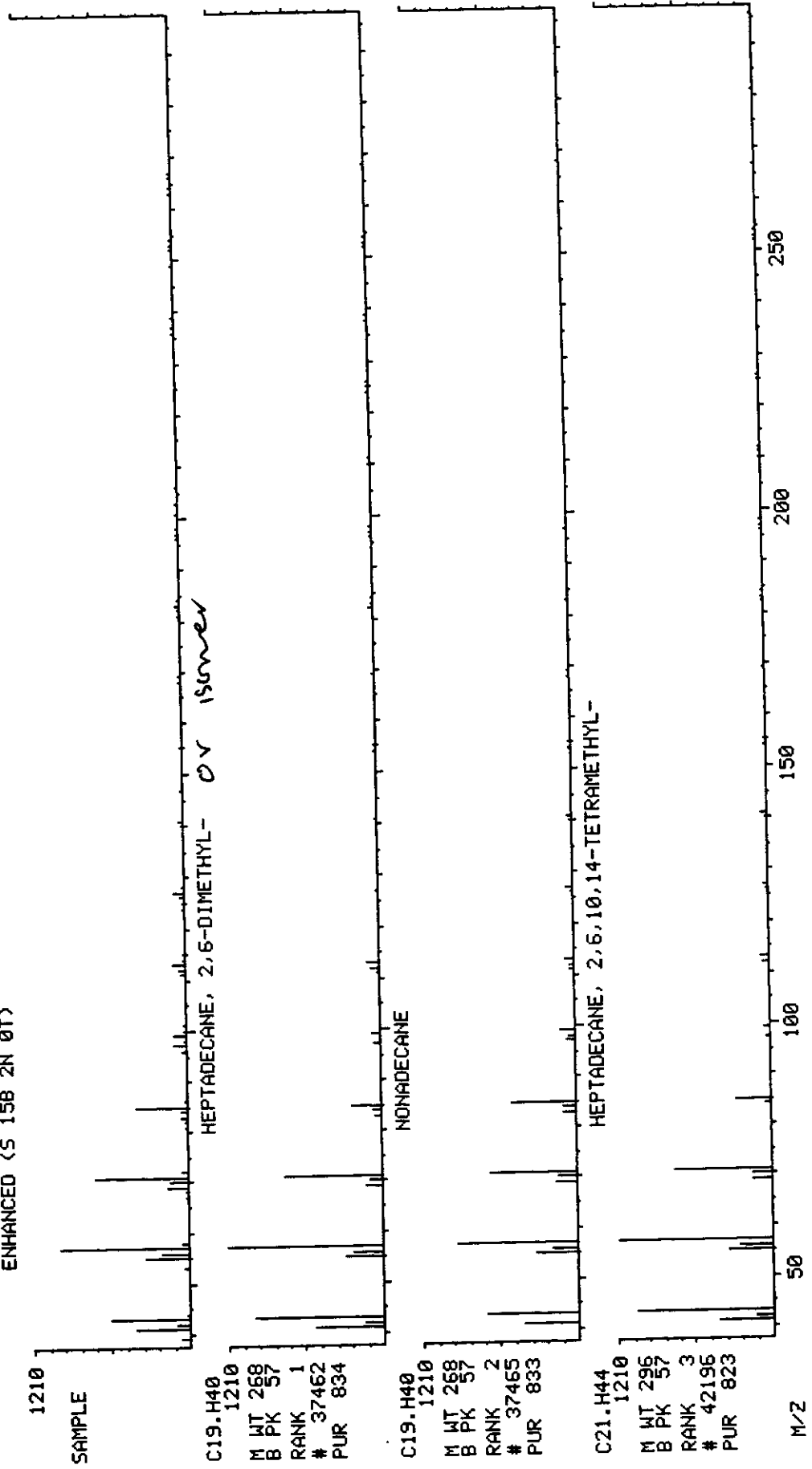
08/31/98 23:28:00 + 12:26

SAMPLE: 5-MMS-4 1/35A/100M

CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M

ENHANCED <S 158 2N 0T>

INST. ID: F16



1210

SAMPLE

C19.H40

1210

M WT 268

B PK 57

RANK 1

# 37462

PUR 834

C19.H40

1210

M WT 268

B PK 57

RANK 2

# 37465

PUR 833

C21.H44

1210

M WT 296

B PK 57

RANK 3

# 42196

PUR 823

M/Z

Library Search                    Data: 30068110 #1190            Base m/z: 57  
 08/31/98 23:28:00 + 13:14       Cali: 30068110 # 3            RIC: 45056.  
 Sample: S-MM5-4 1/3SA/100M       INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 294 matched at least 8 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 37465 NONADECANE  
 2 37462 HEPTADECANE, 2,6-DIMETHYL-  
 3 15969 HYDROXYLAMINE, O-DECYL-  
 4 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 5 19016 UNDECANE, 4,7-DIMETHYL-  
 6 15353 2,6-DIMETHYLDECANE  
 7 19005 UNDECANE, 3,8-DIMETHYL-  
 8 19523 1-DECANOL, 2-ETHYL-  
 9 46161 TRICOSANE

Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C19.H40	268	57	728	940	761
2	C19.H40	268	57	727	920	759
3	C10.H23.O.N	173	43	727	962	727
4	C21.H44	296	57	707	931	738
5	C13.H28	184	43	698	959	704
6	C12.H26	170	43	693	926	697
7	C13.H28	184	57	689	970	689
8	C12.H26.O	186	57	687	925	696
9	C23.H48	324	43	686	877	772

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-92-5
2	---	---	---	---	54105-67-8
3	---	---	---	---	29812-79-1
4	---	---	---	---	18344-37-1
5	---	---	---	---	17301-32-5
6	---	---	---	---	13150-81-7
7	---	---	---	---	17301-30-3
8	---	---	---	---	21078-65-9
9	---	---	---	---	638-67-5

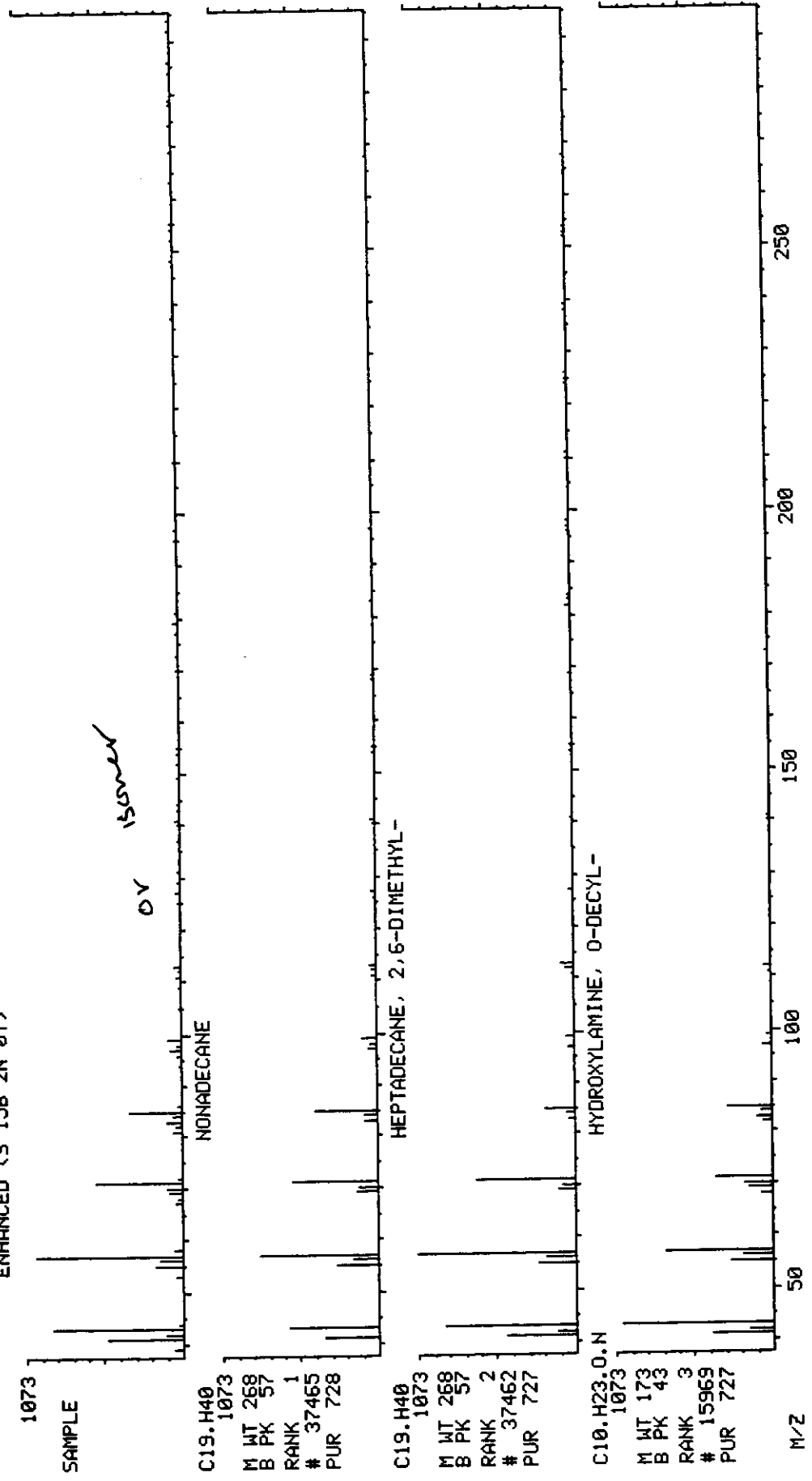


BASE M/Z: 57  
RIC: 45056.

DATA: 30068110 #1190  
CALI: 30068110 # 3

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:28:00 + 13:14  
SAMPLE: S-MMS-4 1/35A/100M  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 15B 2N 0T)

INST. ID: F16



130501

Library Search                      Data: 30068110 #1264                      Base m/z: 57  
 08/31/98 23:28:00 + 14:04                      Cali: 30068110 # 3                      RIC: 34496.  
 Sample: S-MM5-4 1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 819 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.                      Name  
 1 37462 HEPTADECANE, 2,6-DIMETHYL-  
 2 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 3 32058 HEXADECANE, 3-METHYL-  
 4 15969 HYDROXYLAMINE, O-DECYL-  
 5 37465 NONADECANE  
 6 25997 PENTADECANE  
 7 34811 HEPTADECANE, 2-METHYL-  
 8 34816 HEPTADECANE, 4-METHYL-  
 9 42192 HEPTADECANE, 2,6,10,15-TETRAMETHYL-

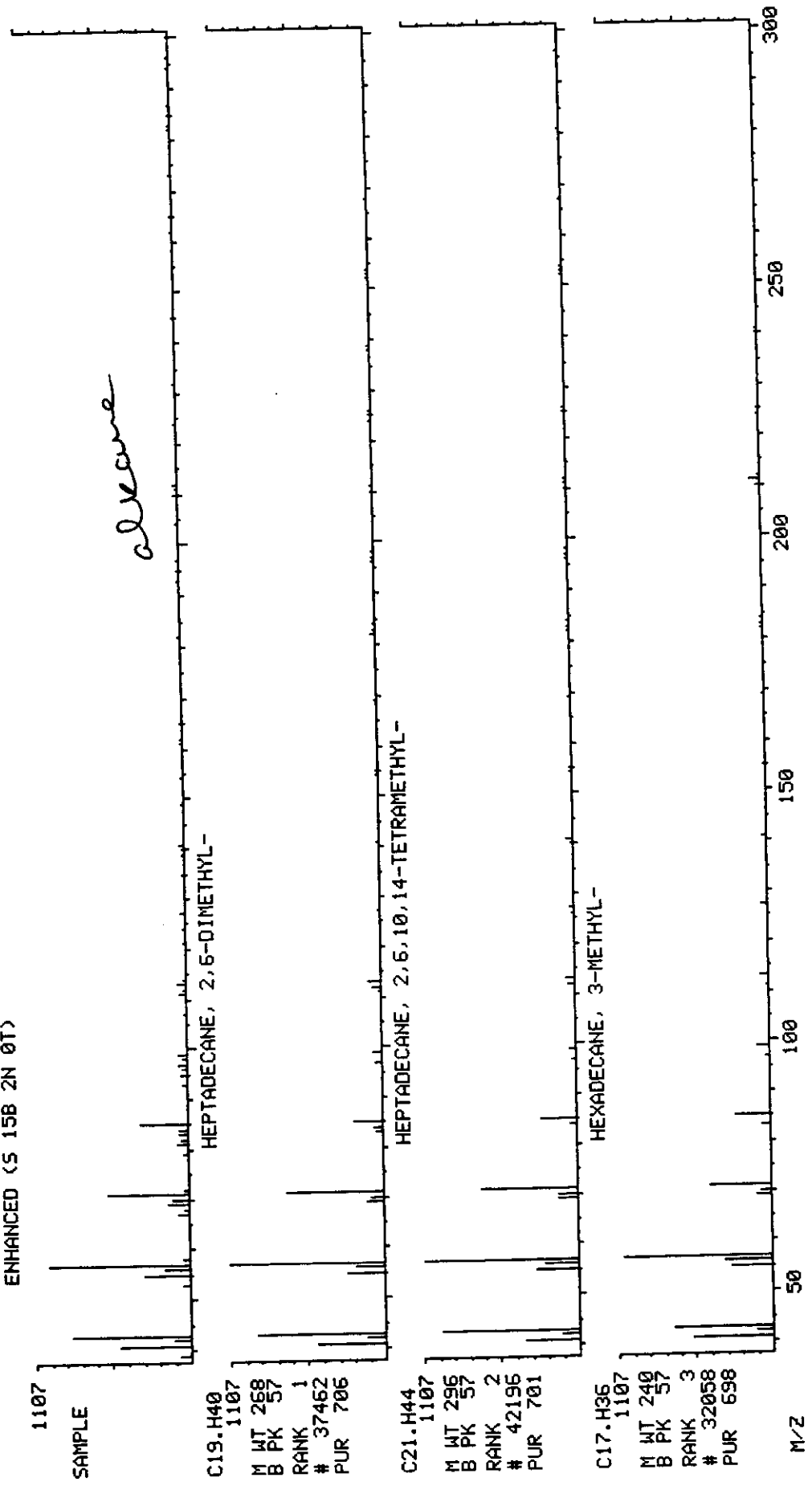
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	706	921	728
2	C21.H44	296	57	701	933	725
3	C17.H36	240	57	698	928	717
4	C10.H23.O.N	173	43	687	968	690
5	C19.H40	268	57	685	936	723
6	C15.H32	212	57	674	947	704
7	C18.H38	254	43	673	885	725
8	C18.H38	254	43	669	894	704
9	C21.H44	296	57	669	904	706

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	54105-67-8
2	---	---	---	---	18344-37-1
3	---	---	---	---	6418-43-5
4	---	---	---	---	29812-79-1
5	---	---	---	---	629-92-5
6	---	---	---	---	629-62-9
7	---	---	---	---	1560-89-0
8	---	---	---	---	26429-11-8
9	---	---	---	---	54833-48-6

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:28:00 + 14:04  
 SAMPLE: 5-MMS-4 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068110 #1264  
 CALI: 30068110 # 3

BASE M/Z: 57  
 RIC: 34496.



Library Search                      Data: 30068110 #1335                      Base m/z: 57  
 08/31/98 23:28:00 + 14:51                      Cali: 30068110 # 3                      RIC: 22112.  
 Sample: S-MM5-4 1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 470 matched at least 7 of the 16 largest peaks in the unknown

Rank In.                      Name  
 1 37456 2-METHYLOCTADECANE  
 2 37462 HEPTADECANE, 2,6-DIMETHYL-  
 3 19523 1-DECANOL, 2-ETHYL-  
 4 29263 HEXADECANE  
 5 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 6 32058 HEXADECANE, 3-METHYL-  
 7 37458 OCTADECANE, 2-METHYL-  
 8 37465 NONADECANE  
 9 15969 HYDROXYLAMINE, O-DECYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	43	571	855	615
2	C19.H40	268	57	568	831	599
3	C12.H26.O	186	57	548	935	557
4	C16.H34	226	57	547	863	599
5	C21.H44	296	57	542	823	582
6	C17.H36	240	57	541	885	584
7	C19.H40	268	43	534	819	607
8	C19.H40	268	57	533	900	578
9	C10.H23.O.N	173	43	532	926	554

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	- -
2	---	---	---	---	54105-67-8
3	---	---	---	---	21078-65-9
4	---	---	---	---	544-76-3
5	---	---	---	---	18344-37-1
6	---	---	---	---	6418-43-5
7	---	---	---	---	1560-88-9
8	---	---	---	---	629-92-5
9	---	---	---	---	29812-79-1

BASE M/Z: 57  
RIC: 22112.

DATA: 30068110 #1335  
CALI: 30068110 # 3

MID LIBRARY SEARCH (LIBRARYB)

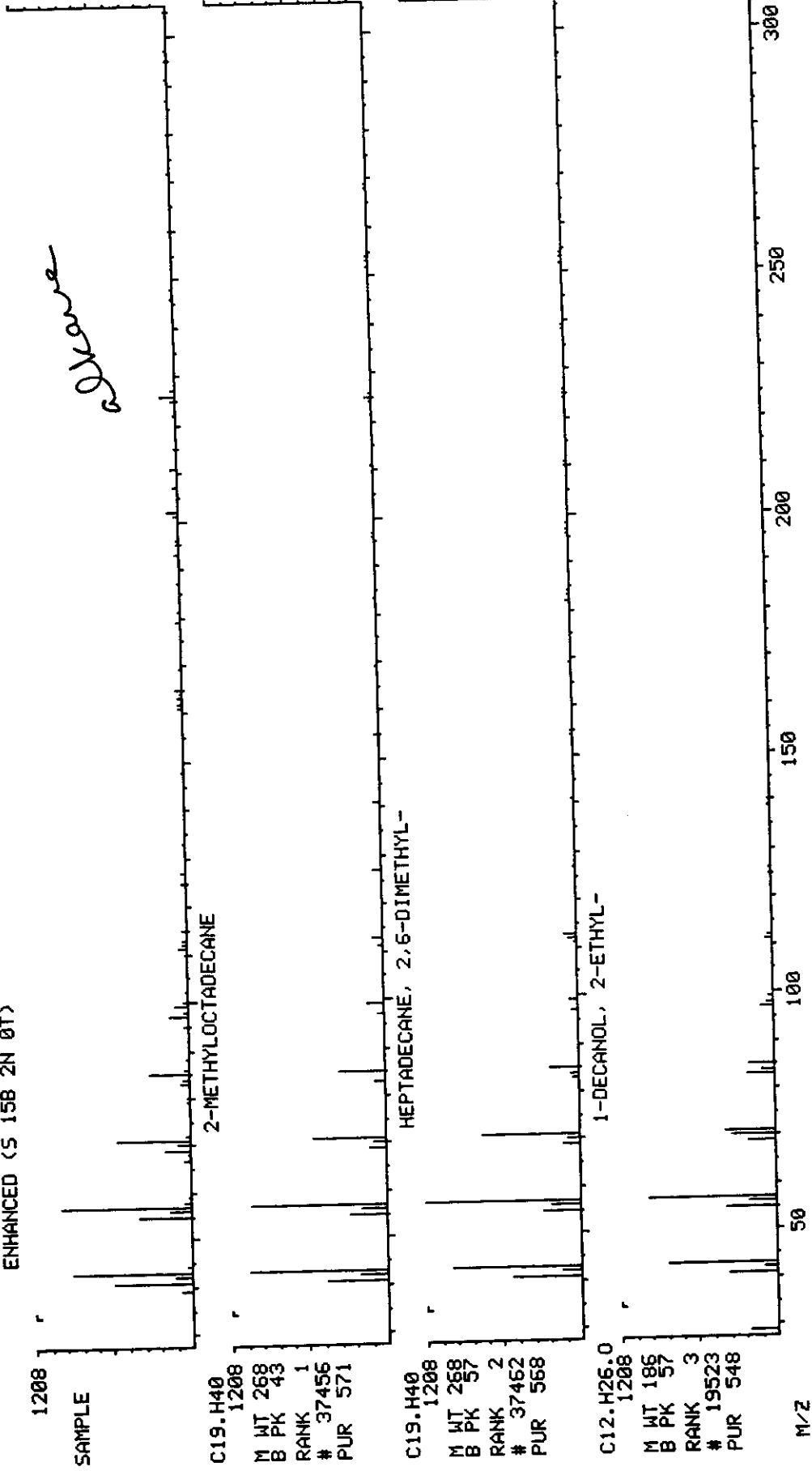
08/31/98 23:28:00 + 14:51

SAMPLE: 5-MMS-4 1/35A/100M

COND.S.: UG/ML \*100ML \*100Z \*(NA/NA )/1/35A NA M

ENHANCED (S 15B 2N 0T)

INST. ID: F16



H  
CO  
CO  
CO

TIC SELECTION REPORT

DATA FILE: 30068110

THE FOLLOWING PEAKS WERE REJECTED BECAUSE AT LEAST 40 % OF THEIR SIZE WAS ACCOUNTED FOR BY TARGET COMPOUNDS ELUTING WITHIN 4 SCANS OF THE PEAK TOP.

SCAN	SIZE	AMOUNT
340	197830.	21.556
530	455680.	49.651
829	367104.	40.000
1087	361984.	40.000
1552	257952.	40.000
1783	238432.	40.000

TOTAL NUMBER OF UNIDENTIFIED PEAKS WITH SIZE GREATER THAN 10 % OF THE CLOSEST INTERNAL STANDARD THAT DOES NOT HAVE INTERFERENCES = 84

INTERNAL STANDARDS THAT HAVE RIC SIZE LESS THAN 50 % OR GREATER THAN 200 % OF THE ESTIMATED RIC SIZE ARE CONSIDERED TO HAVE INTERFERENCES AND WILL NOT BE USED FOR QUANTITATION.

#	INTERNAL STANDARD	RIC SIZE	PERCENT OF ESTIMATED	
			RIC SIZE	SCAN
1	C140 NAPHTHALENE-D8	455680.	248 *	530
2	C150 ACENAPHTHENE-D10	367104.	161	829
3	C160 PHENANTHRENE-D10	361984.	145	1087
4	C170 CHRYSENE-D12	257952.	153	1552
5	C175 PERYLENE-D12	238432.	162	1783

\* INDICATES INTERFERENCE

SIZE = AREA

Semivolatile Organics  
Method 0010/8270

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-5-F, FH, XAD, COND, BH  
 LAB ID: 300681-0011-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98

Sampled: 28 JUL 98  
 Prepared: 31 JUL 98

Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
Phenol	ND	ug/Sample	3000	GR
bis(2-Chloroethyl)ether	ND	ug/Sample	3000	
2-Chlorophenol	ND	ug/Sample	3000	
1,3-Dichlorobenzene	ND	ug/Sample	3000	
1,4-Dichlorobenzene	ND	ug/Sample	3000	
Benzyl alcohol	ND	ug/Sample	3000	
1,2-Dichlorobenzene	ND	ug/Sample	3000	
2-Methylphenol	ND	ug/Sample	3000	
2,2'-Oxybis(1-chloropropane)	ND	ug/Sample	3000	
3/4-Methylphenol	ND	ug/Sample	3000	
N-Nitroso-di-n-propylamine	ND	ug/Sample	3000	
Hexachloroethane	ND	ug/Sample	3000	
Nitrobenzene	ND	ug/Sample	3000	
Isophorone	ND	ug/Sample	3000	
2-Nitrophenol	ND	ug/Sample	3000	
2,4-Dimethylphenol	ND	ug/Sample	3000	
Benzoic acid	ND	ug/Sample	15000	
bis(2-Chloroethoxy)-methane	ND	ug/Sample	3000	
2,4-Dichlorophenol	ND	ug/Sample	3000	
1,2,4-Trichlorobenzene	ND	ug/Sample	3000	
Naphthalene	2600	ug/Sample	3000	J
4-Chloroaniline	ND	ug/Sample	3000	
Hexachlorobutadiene	ND	ug/Sample	3000	
4-Chloro-3-methylphenol	ND	ug/Sample	3000	
2-Methylnaphthalene	4400	ug/Sample	3000	
Hexachlorocyclopentadiene	ND	ug/Sample	3000	
2,4,6-Trichlorophenol	ND	ug/Sample	3000	
2,4,5-Trichlorophenol	ND	ug/Sample	15000	
2-Chloronaphthalene	ND	ug/Sample	3000	
2-Nitroaniline	ND	ug/Sample	3000	
Dimethyl phthalate	ND	ug/Sample	3000	
Acenaphthylene	ND	ug/Sample	3000	
3-Nitroaniline	ND	ug/Sample	15000	
Acenaphthene	ND	ug/Sample	3000	
2,4-Dinitrophenol	ND	ug/Sample	15000	

Note G = Reporting limit(s) raised due to matrix interference.  
 Note J = Result is detected below the reporting limit or is an estimated concentration.  
 Note R = Reporting limit(s) raised due to sample volume limitations.  
 ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.  
Rev 230787

1391

Semivolatile Organics  
Method 0010/8270

(cont.)

Client Name: Pacific Environmental Services  
Client ID: S-MM5-5-F,FH,XAD,COND,BH  
LAB ID: 300681-0011-SA  
Matrix: AIRTRAIN  
Authorized: 30 JUL 98

Sampled: 28 JUL 98  
Prepared: 31 JUL 98

Received: 30 JUL 98  
Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
4-Nitrophenol	ND	ug/Sample	15000	
Dibenzofuran	ND	ug/Sample	3000	
2,4-Dinitrotoluene	ND	ug/Sample	3000	
2,6-Dinitrotoluene	ND	ug/Sample	3000	
Diethyl phthalate	ND	ug/Sample	3000	
4-Chlorophenyl phenyl ether	ND	ug/Sample	3000	
Fluorene	560	ug/Sample	3000	
4-Nitroaniline	ND	ug/Sample	15000	J
4,6-Dinitro-2-methylphenol	ND	ug/Sample	15000	
N-Nitrosodiphenylamine	ND	ug/Sample	3000	
4-Bromophenyl phenyl ether	ND	ug/Sample	3000	
Hexachlorobenzene	ND	ug/Sample	3000	
Pentachlorophenol	ND	ug/Sample	15000	
Phenanthrene	1200	ug/Sample	3000	J
Anthracene	ND	ug/Sample	3000	
Di-n-butyl phthalate	ND	ug/Sample	3000	
Fluoranthene	ND	ug/Sample	3000	
Pyrene	ND	ug/Sample	3000	
Butyl benzyl phthalate	ND	ug/Sample	3000	
3,3'-Dichlorobenzidine	ND	ug/Sample	6000	
Benzo(a)anthracene	ND	ug/Sample	3000	
bis(2-Ethylhexyl)-phthalate	ND	ug/Sample	3000	
Chrysene	ND	ug/Sample	3000	
Di-n-octyl phthalate	ND	ug/Sample	3000	
Benzo(b)fluoranthene	ND	ug/Sample	3000	
Benzo(k)fluoranthene	ND	ug/Sample	3000	
Benzo(a)pyrene	ND	ug/Sample	3000	
Indeno(1,2,3-cd)pyrene	ND	ug/Sample	3000	
Dibenz(a,h)anthracene	ND	ug/Sample	3000	
Benzo(g,h,i)perylene	ND	ug/Sample	3000	
Acetophenone	ND	ug/Sample	3000	
4-Aminobiphenyl	ND	ug/Sample	15000	
Aniline	ND	ug/Sample	3000	
Benzidine	ND	ug/Sample	30000	
3,3'-Dimethylbenzidine	ND	ug/Sample	6000	
N-Nitrosodimethylamine	ND	ug/Sample	3000	
N-Nitrosomorpholine	ND	ug/Sample	3000	

Note J = Result is detected below the reporting limit or is an estimated concentration.  
ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.  
Rev 230787

1392



Semivolatile Organics  
Method 0010/8270

(cont.)

Client Name: Pacific Environmental Services  
Client ID: S-MM5-5-F,FH,XAD,COND,BH  
LAB ID: 300681-0011-SA  
Matrix: AIRTRAIN  
Authorized: 30 JUL 98

Sampled: 28 JUL 98  
Prepared: 31 JUL 98

Received: 30 JUL 98  
Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Wet wt. Result	Units	Reporting Limit	Qualifier
Pentachloronitrobenzene (PCNB)	ND	ug/Sample	15000	
o-Toluidine	ND	ug/Sample	6000	
2-Methoxybenzenamine	ND	ug/Sample	--	
Biphenyl	ND	ug/Sample	--	
Chloroacetophenone	ND	ug/Sample	--	
Cumene	ND	ug/Sample	--	
DBCP (1,2-Dibromo-3-chloropropane)	ND	ug/Sample	--	
Benzo(e)pyrene	ND	ug/Sample	--	
N-N-Diethylaniline	ND	ug/Sample	--	
Dimethylaniline	ND	ug/Sample	--	
3,3'-Dimethoxybenzidine	ND	ug/Sample	--	
Hydroquinone	ND	ug/Sample	--	
4,4'-Methyl-bis(2-chloroaniline)	ND	ug/Sample	--	
4-Nitrodiphenyl	ND	ug/Sample	--	
Trifluralin	ND	ug/Sample	--	

Surrogate	Recovery	Acceptable Range	
Nitrobenzene-d5	ND %	45 - 107	H
2-Fluorobiphenyl	ND %	62 - 110	
Terphenyl-d14	ND %	58 - 135	
Phenol-d5	ND %	43 - 130	
2-Fluorophenol	ND %	36 - 111	
2,4,6-Tribromophenol	ND %	58 - 131	

Note H = Spiked analyte not detected because of required sample dilution.  
ND = Not Detected

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.  
Rev 230787

1396

Semivolatiles Library Search (20 Compound TID)  
Method 8270

Client Name: Pacific Environmental Services  
 Client ID: S-MM5-5-F,FH,XAD,COND,BH  
 LAB ID: 300681-0011-SA  
 Matrix: AIRTRAIN  
 Authorized: 30 JUL 98  
 Sampled: 28 JUL 98  
 Prepared: NA  
 Received: 30 JUL 98  
 Analyzed: 31 AUG 98

Dilution Factor: 100

Parameter	Result	Units	Reporting Limit	Qualifier
Unknown	11000	ug/Sample	--	
Unknown	13000	ug/Sample	--	
Decane, 2,9-dimethyl-	9200	ug/Sample	--	0
Undecane, 2,6-dimethyl-	9500	ug/Sample	--	0
Unknown	15000	ug/Sample	--	
Unknown alkane	20000	ug/Sample	--	
Unknown	12000	ug/Sample	--	
Unknown alkane	11000	ug/Sample	--	
Unknown	10000	ug/Sample	--	
Unknown	12000	ug/Sample	--	
Dodecane, 2,6,10-trimethyl-	19000	ug/Sample	--	0
Tetradecane	25000	ug/Sample	--	0
Unknown	15000	ug/Sample	--	
Heptadecane, 2,6,10,14 -tetramethyl-	22000	ug/Sample	--	0
Dodecane, 3-methyl-	25000	ug/Sample	--	0
Oxirane, hexadecyl-	13000	ug/Sample	--	0
Unknown alkane	27000	ug/Sample	--	
Unknown alkane	14000	ug/Sample	--	
Heptadecane, 2,6-dimethyl-	28000	ug/Sample	--	0
Nonadecane	11000	ug/Sample	--	0

Note 0 = Or structurally similar compound (isomer).  
 NA = Not Applicable

Reported By: Emily Uebelhoer

Approved By: Mike Orbanosky

The cover letter is an integral part of this report.

Rev 230787

1394

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068111  
Std Id: ST16980831

Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:58  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC Limits
CS20 NITROBENZENE-D5	50.0	BDL	0*	45 107
CS25 2-FLUOROBIPHENY	50.00	108.0	216.	* 62 110
CS30 TERPHENYL-D14	50.0	BDL	0*	58 135
CS45 PHENOL-D5	100.	BDL	0*	43 130
CS50 2-FLUOROPHENOL	100.	BDL	0*	36 111
CS55 2,4,6-TRIBROMOP	100.	BDL	0*	58 131

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
HEXACHLOROENZENE-C13		ND	UG/A	3000.0
C310 N-NITROSODIMETHYLAMINE		ND	UG/A	3000.0
PYRIDINE		ND	UG/A	6000.0
2-PICOLINE		ND	UG/A	3000.0
N-NITROSOMETHYLETHYLAMINE		ND	UG/A	3000.0
METHYLMETHANESULFONATE		ND	UG/A	3000.0
N-NITROSODIETHYLAMINE		ND	UG/A	3000.0
ETHYLMETHANE SULFONATE		ND	UG/A	3000.0
PENTACHLOROETHANE		ND	UG/A	3000.0
C320 ANILINE		ND	UG/A	3000.0
C315 PHENOL		ND	UG/A	3000.0
C325 BIS(2-CHLOROETHYL)ETHER		ND	UG/A	3000.0
C330 2-CHLOROPHENOL		ND	UG/A	3000.0
C335 1,3-DICHLOROBENZENE		ND	UG/A	3000.0
C340 1,4-DICHLOROBENZENE		ND	UG/A	3000.0
C345 BENZYL ALCOHOL		ND	UG/A	3000.0
C350 1,2-DICHLOROBENZENE		ND	UG/A	3000.0
C355 2-METHYLPHENOL		ND	UG/A	3000.0
C360 2,2'-OXYBIS(1-CLPROPAN)		ND	UG/A	3000.0
C361 ACETOPHENONE		ND	UG/A	3000.0
N-NITROSOPYRROLIDINE		ND	UG/A	3000.0
N-NITROSOMORPHOLINE		ND	UG/A	3000.0
3-METHYL PHENOL		ND	UG/A	3000.0
C365 4-METHYLPHENOL		ND	UG/A	3000.0
C370 N-NITROSO-DI-N-PROPYLAM		ND	UG/A	3000.0
O-TOLUIDINE		ND	UG/A	3000.0
C375 HEXACHLOROETHANE		ND	UG/A	3000.0
C410 NITROBENZENE		ND	UG/A	3000.0
N-NITROSOPIPERIDINE		ND	UG/A	3000.0

Reviewed by:

*Mund* 9.18.98  
*DSC* 10/16/98

1385

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068111  
Std Id: ST16980831

Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:58  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C415 ISOPHORONE		ND	UG/A	3000.0
C420 2-NITROPHENOL		ND	UG/A	3000.0
C425 2,4-DIMETHYLPHENOL		ND	UG/A	3000.0
C435 BIS(2-CHLOROETHOXY)METH		ND	UG/A	3000.0
C440 2,4-DICHLOROPHENOL		ND	UG/A	3000.0
C430 BENZOIC ACID		ND	UG/A	15000.0
AA-DIMETHYLPHENETHYLAMINE		ND	UG/A	3000.0
C445 1,2,4-TRICHLOROBENZENE		ND	UG/A	3000.0
C450 NAPHTHALENE		ND 2600	UG/A	3000.0
C455 4-CHLOROANILINE		ND	UG/A	3000.0
2,6-DICHLOROPHENOL		ND	UG/A	3000.0
HEXACHLOROPROPENE		ND	UG/A	3000.0
C460 HEXACHLOROBUTADIENE		ND	UG/A	3000.0
P-PHENYLENE DIAMINE		ND	UG/A	3000.0
N-NITROSODI-N-BUTYLAMINE		ND	UG/A	3000.0
C465 4-CHLORO-3-METHYLPHENO		ND	UG/A	3000.0
SAFROLE		ND	UG/A	3000.0
C470 2-METHYLNAPHTHALENE	653	4400.	UG/A	3000.0
1,2,4,5-TETRACHLOROBENZENE		ND	UG/A	3000.0
ISOSAFROLE (#1)		ND	UG/A	6000.0
C510 HEXACHLOROCYCLOPENTADI		ND	UG/A	3000.0
C515 2,4,6-TRICHLOROPHENOL		ND	UG/A	3000.0
C520 2,4,5-TRICHLOROPHENOL		ND	UG/A	3000.0
ISOSAFROLE (#2)		ND	UG/A	6000.0
C525 2-CHLORONAPHTHALENE		ND	UG/A	3000.0
1-CHLORONAPHTHALENE		ND	UG/A	1500.0
C530 2-NITROANILINE		ND	UG/A	15000.0
1,4-NAPHTHOQUINONE		ND	UG/A	3000.0
C535 DIMETHYLPHTHALATE		ND	UG/A	3000.0
1,3-DINITROBENZENE		ND	UG/A	3000.0
C540 ACENAPHTHYLENE		ND	UG/A	3000.0
C543 2,6-DINITROTOLUENE		ND	UG/A	3000.0
C545 3-NITROANILINE		ND	UG/A	15000.0
C550 ACENAPHTHENE		ND	UG/A	3000.0
C555 2,4-DINITROPHENOL		ND	UG/A	15000.0
C565 DIBENZOFURAN		ND	UG/A	3000.0
C560 4-NITROPHENOL		ND	UG/A	15000.0
PENTACHLOROBENZENE		ND	UG/A	3000.0
C570 2,4-DINITROTOLUENE		ND	UG/A	3000.0
1-NAPHTHYLAMINE		ND	UG/A	3000.0
2-NAPHTHYLAMINE		ND	UG/A	3000.0
2,3,4,6-TETRACHLOROPHENOL		ND	UG/A	6000.0
C580 DIETHYLPHTHALATE		ND	UG/A	3000.0
C590 FLUORENE		ND 560	UG/A	3000.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 30068111  
Std Id: ST16980831

Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:58  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C585 4-CHLOROPHENYL-PHENYLE		ND	UG/A	3000.0
5-NITRO-O-TOLUIDINE		ND	UG/A	3000.0
C595 4-NITROANALINE		ND	UG/A	15000.0
C610 4,6-DINITRO-2-METHYLPH		ND	UG/A	15000.0
C615 N-NITROSODIPHENYLAMINE		ND	UG/A	3000.0
C620 AZOBENZENE		ND	UG/A	3000.0
SYM-TRINITROBENZENE		ND	UG/A	3000.0
C625 4-BROMOPHENYL-PHENYLET		ND	UG/A	3000.0
PHENACETIN		ND	UG/A	3000.0
DIALLATE	234	ND	UG/A	3000.0
C630 HEXACHLOROBENZENE		ND	UG/A	3000.0
4-AMINOBIIPHENYL		ND	UG/A	3000.0
C635 PENTACHLOROPHENOL		ND	UG/A	15000.0
PRONAMIDE		ND	UG/A	3000.0
PENTACHLORONITROBENZENE		ND	UG/A	15000.0
C640 PHENANTHRENE		ND (1200)	UG/A	3000.0
C645 ANTHRACENE		ND	UG/A	3000.0
2SECBUTYL-4,6-DINITROPHENOL		ND	UG/A	3000.0
C647 CARBAZOLE		ND	UG/A	3000.0
C650 DI-N-BUTYLPHTHALATE		ND	UG/A	3000.0
4-NITROQUINOLINE-1-OXIDE		ND	UG/A	3000.0
METHAPYRILENE		ND	UG/A	3000.0
ISODRIN		ND	UG/A	3000.0
C655 FLUORANTHENE		ND	UG/A	3000.0
CHLOROBENZILATE		ND	UG/A	3000.0
C710 BENZIDINE		ND	UG/A	30000.0
C715 PYRENE		ND	UG/A	3000.0
ARAMITE (#1)		ND	UG/A	3000.0
ARAMITE (#2)		ND	UG/A	3000.0
P-DIMETHYLAMINOAZOBENZENE		ND	UG/A	3000.0
3,3'-DIMETHYLBENZIDINE		ND	UG/A	3000.0
KEPONE		ND	UG/A	15000.0
C720 BUTYLBENZYLPHTHALATE		ND	UG/A	3000.0
2-ACETYLAMINOFLUORENE		ND	UG/A	3000.0
C730 BENZO(A)ANTHRACENE		ND	UG/A	3000.0
C725 3,3'-DICHLOROBENZIDINE		ND	UG/A	6000.0
C740 CHRYSENE		ND	UG/A	3000.0
C745 BIS(2-ETHYLHEXYL)PHTHA		ND	UG/A	3000.0
3-METHYLCHOLANTHRENE		ND	UG/A	3000.0
C760 DI-N-OCTYL PHTHALATE		ND	UG/A	3000.0
C765 BENZO(B)FLUORANTHENE		ND	UG/A	3000.0
7,12-DIMETHYLBENZANTHRACENE		ND	UG/A	3000.0
C770 BENZO(K)FLUORANTHENE		ND	UG/A	3000.0
HEXACHLOROPHENE	1761	ND (5300)	UG/A	3000.0

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ND (1200)

ND (5300)

2 9/16/98

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

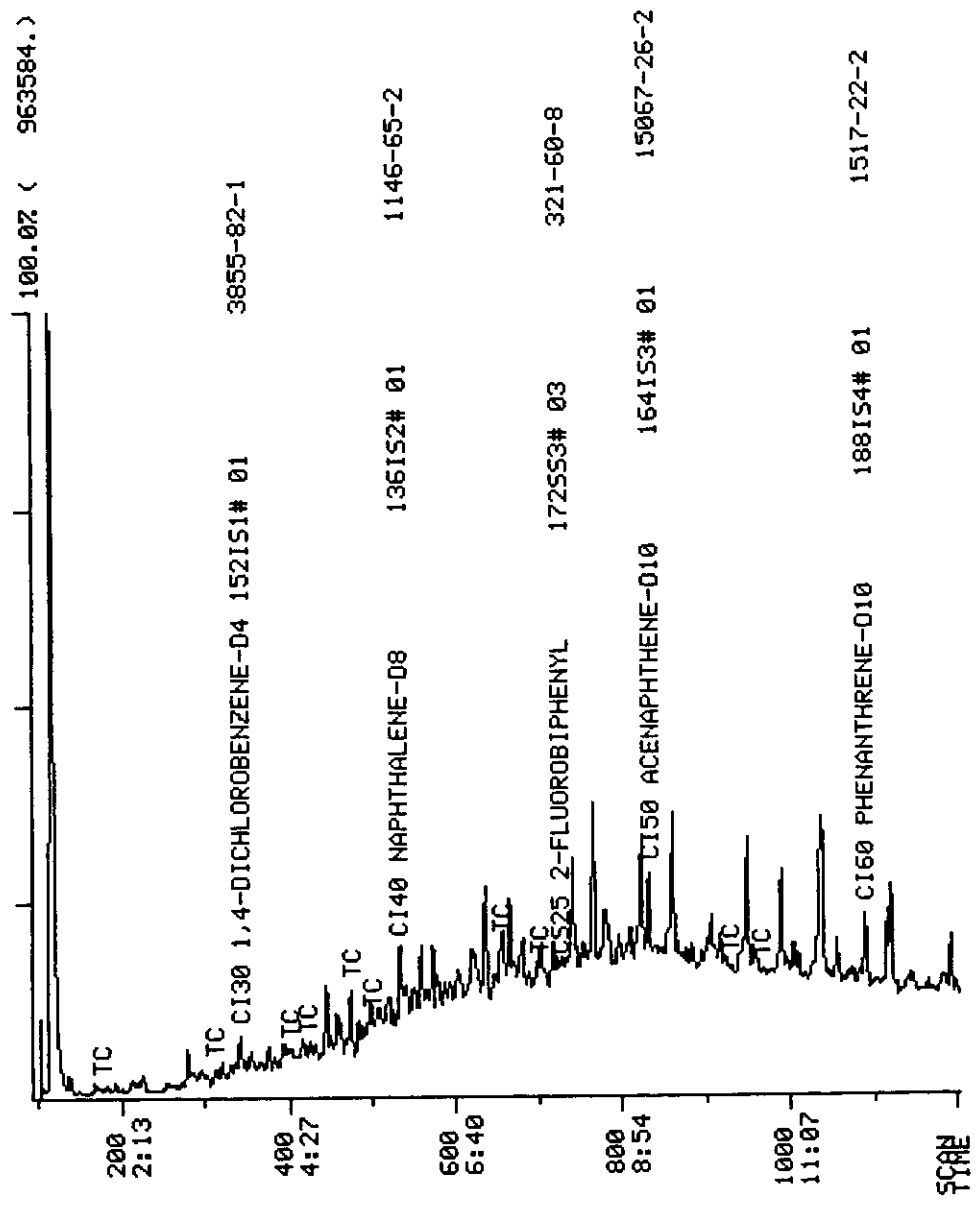
Data File: 30068111  
Std Id: ST16980831

Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 23:58  
Analyst: DAT Instrument ID: F16 Run Factor: 300.  
Quan List Threshold: 0.95 Surrogate Vol.: 100.

Target Compounds: SAP9

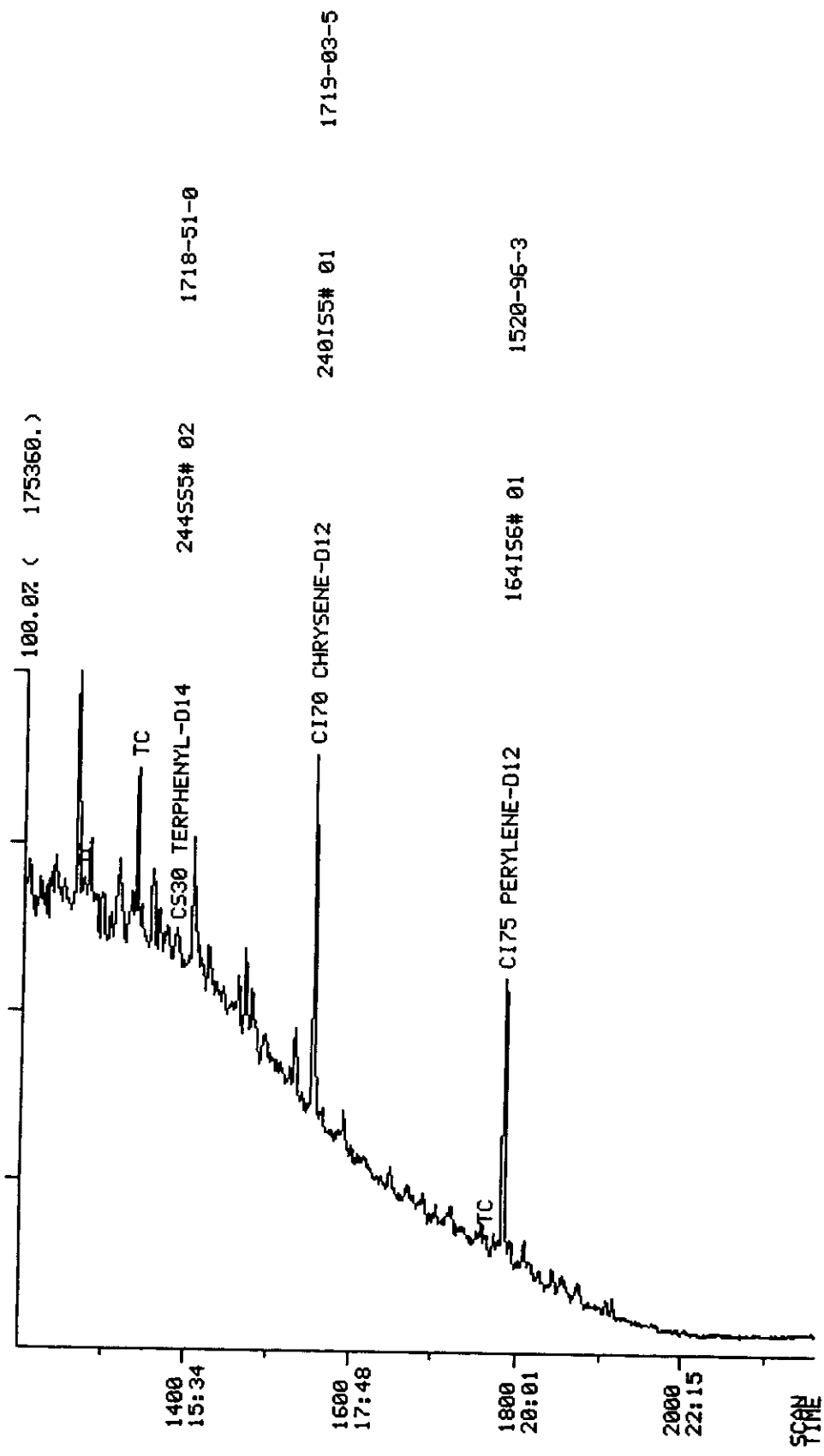
Parameter	Scan	Result	Units	Reporting Limit
C775 BENZO(A)PYRENE		ND	UG/A	3000.0
C780 INDENO(1,2,3-CD)PYRENE		ND	UG/A	3000.0
C785 DIBENZ(A,H)ANTHRACENE		ND	UG/A	3000.0
C790 BENZO(G,H,I)PERYLENE		ND	UG/A	3000.0

DATA FROM FILE: 30068111 SCANS 93 TO 1203 ACQUIRED: 08/31/98 23:58:00  
 CALLI: 30068111 #3  
 SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M



1300

DATA FROM FILE: 30068111 SCANS 1203 TO 2158 ACQUIRED: 08/31/98 23:58:00  
 CALI: 30068111 #3  
 SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA )/1/35A NA M





ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 30068111

Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 23:58

Compounds with amounts less than 0.20 reported as NOT FOUND

Library						Units: UG/ML	
No Entry	Name	Mass Meth	Scan Ref	Area RRF(L)	Amount		
1 S1#	1 C130 1,4-DICHLOROBENZENE	152 A BV	340 1	30555. 1.000	40.000		
2 S2#	1 C140 NAPHTHALENE-D8	136 A VB	530 2	109200. 1.000	40.000		
3 S3#	1 C150 ACENAPHTHENE-D10	164 A BB	830 3	62156. 1.000	40.000		
4 S4#	1 C160 PHENANTHRENE-D10	188 A BB	1088 4	110919. 1.000	40.000		
5 S5#	1 C170 CHRYSENE-D12	240 A BB	1553 5	89347. 1.000	40.000		
6 S6#	1 C175 PERYLENE-D12	264 A BB	1783 6	73934. 1.000	40.000		
7 S2#	2 CS20 NITROBENZENE-D5	82	2 NOT FOUND				
8 S3#	3 CS25 2-FLUOROBIPHENYL	172 A BB	724 3	2070. 1.234	1.079		
9 S5#	2 CS30 TERPHENYL-D14	244 A BB	1385 5	1349. 0.995	0.607		
10 S1#	3 CS45 PHENOL-D5	99	1 NOT FOUND				
11 S1#	2 CS50 2-FLUOROPHENOL	112	1 NOT FOUND				
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330 A BB	972 3	276. 0.236	0.754		
13 S4#	4 HEXACHLOROBENZENE-C13	294	4 NOT FOUND				
14 S1#	15 C310 N-NITROSODIMETHYLAM	74	1 NOT FOUND				
15 S1#	20 PYRIDINE	79	1 NOT FOUND				
16 S1#	30 2-PICOLINE	93	1 NOT FOUND				
17 S1#	40 N-NITROSOMETHYLETHYLAMIN	42 A VV	174 1	2680. 0.811	4.325		
18 S1#	60 METHYLMETHANESULFONATE	80	1 NOT FOUND				
19 S1#	70 N-NITROSODIETHYLAMINE	102	1 NOT FOUND				
20 S1#	85 ETHYLMETHANE SULFONATE	79	1 NOT FOUND				
21 S1#	95 PENTACHLOROETHANE	117 A BB	306 1	296. 0.602	0.644		
22 S1#	100 C320 ANILINE	93	1 NOT FOUND				
23 S1#	105 C315 PHENOL	94 A BB	311 1	1764. 1.968	1.174		
24 S1#	110 C325 BIS(2-CHLOROETHYL)E	93 A BB	319 1	232. 1.368	0.222		
25 S1#	115 C330 2-CHLOROPHENOL	128	1 NOT FOUND				
26 S1#	125 C335 1,3-DICHLOROBENZENE	146	1 NOT FOUND				
27 S1#	130 C340 1,4-DICHLOROBENZENE	146	1 NOT FOUND				
28 S1#	145 C345 BENZYL ALCOHOL	108 A BB	373 1	515. 0.854	0.789		
29 S1#	150 C350 1,2-DICHLOROBENZENE	146	1 NOT FOUND				
30 S1#	160 C355 2-METHYLPHENOL	108	1 NOT FOUND				
31 S1#	165 C360 2,2'-OXYBIS(1-CLPRO	45	1 NOT FOUND				
32 S1#	170 C361 ACETOPHENONE	105 A BB	400 1	5565. 1.934	3.767		
33 S1#	175 N-NITROSPYRROLIDINE	100	1 NOT FOUND				
34 S1#	180 N-NITROSOMORPHOLINE	56 A BB	410 1	3449. 0.821	5.503		
35 S1#	182 3-METHYL PHENOL	108 A BB	419 1	1130. 2.252	0.657		
36 S1#	185 C365 4-METHYLPHENOL	108	1 NOT FOUND				
37 S1#	190 C370 N-NITROSO-DI-N-PROP	70 A VB	403 1	584. 0.906	0.844		
38 S1#	195 O-TOLUIDINE	106	1 NOT FOUND				
39 S1#	200 C375 HEXACHLOROETHANE	117	1 NOT FOUND				
40 S2#	10 C410 NITROBENZENE	77 A VB	422 2	1803. 0.445	1.484		
41 S2#	15 N-NITROSOPIPERIDINE	42	2 NOT FOUND				
42 S2#	20 C415 ISOPHORONE	82 A BB	473 2	8202. 0.741	4.053		
43 S2#	25 C420 2-NITROPHENOL	139	2 NOT FOUND				
44 S2#	30 C425 2,4-DIMETHYLPHENOL	107 A BB	500 2	1569. 0.330	1.742		
45 S2#	35 C435 BIS(2-CHLOROETHOXY)	93	2 NOT FOUND				

*DGC*  
*10/6/98*

1401

46	S2#	40	C440	2,4-DICHLOROPHENOL	162	A	BB	520	2	312.	0.308	0.372
47	S2#	45	C430	BENZOIC ACID	122			2	NOT FOUND			
48	S2#	50	AA-	DIMETHYLPHENETHYLAMIN	58			2	NOT FOUND			
49	S2#	55	C445	1,2,4-TRICHLOROBENZ	180	A	VV	524	2	454.	0.331	0.503

50	S2# 60	C480 NAPHTHALENE	128 A BV	533 2	24195. 1.022	8.670	
51	S2# 80	C455 4-CHLOROANILINE	127 A BB	556 2	439. 0.454	0.354	
52	S2# 85	2,6-DICHLOROPHENOL	162	2	NOT FOUND		
53	S2# 90	HEXACHLOROPROPENE	213	2	NOT FOUND		
54	S2# 95	C460 HEXACHLOROBUTADIENE	225	2	NOT FOUND		
55	S2#115	P-PHENYLENE DIAMINE	108	2	NOT FOUND		
56	S2#120	N-NITROSODI-N-BUTYLAMINE	84	2	NOT FOUND		
57	S2#130	C465 4-CHLORO-3-METHYLPH	107	2	NOT FOUND		
58	S2#140	SAFROLE	162 A BB	652 2	1274. 0.248	1.884	
59	S2#145	C470 2-METHYLNAPHTHALENE	142 A BB	653 2	30297. 0.751	14.772	
60	S3# 10	1,2,4,5-TETRACHLOROBENZE	214	3	NOT FOUND		
61	S3# 15	ISOSAFROLE (#1)	162 A BB	698 3	1034. 0.044	15.245	
62	S3# 20	C510 HEXACHLOROCYCLOPENT	237	3	NOT FOUND		
63	S3# 25	C515 2,4,6-TRICHLOROPHEN	196	3	NOT FOUND		
64	S3# 30	C520 2,4,5-TRICHLOROPHEN	196	3	NOT FOUND		
65	S3# 35	ISOSAFROLE (#2)	104 A BB	739 3	420. 0.195	1.388	
66	S3# 40	C525 2-CHLORONAPHTHALENE	162	3	NOT FOUND		
67	S3# 42	1-CHLORONAPHTHALENE	162	3	NOT FOUND		
68	S3# 45	C530 2-NITROANILINE	65 A BB	764 3	476. 0.502	0.611	
69	S3# 50	1,4-NAPHTHOQUINONE	158	3	NOT FOUND		
70	S3# 55	C535 DIMETHYLPHTHALATE	163	3	NOT FOUND		
71	S3# 60	1,3-DINITROBENZENE	168	3	NOT FOUND		
72	S3# 65	C540 ACENAPHTHYLENE	152	3	NOT FOUND		
73	S3# 70	C543 2,6-DINITROTOLUENE	165	3	NOT FOUND		
74	S3# 75	C545 3-NITROANILINE	138	3	NOT FOUND		
75	S3# 80	C550 ACENAPHTHENE	153 A BV	835 3	1969. 1.159	1.094	
76	S3# 85	C555 2,4-DINITROPHENOL	184 A BV	859 3	206. 0.189	0.700	
77	S3# 90	C565 DIBENZOFURAN	168 A BB	867 3	1131. 1.669	0.436	
78	S3# 95	C560 4-NITROPHENOL	109	3	NOT FOUND		
79	S3#100	PENTACHLOROBENZENE	250	3	NOT FOUND		
80	S3#105	C570 2,4-DINITROTOLUENE	165 A BB	882 3	391. 0.397	0.634	
81	S3#110	1-NAPHTHYLAMINE	143	3	NOT FOUND		
82	S3#115	2-NAPHTHYLAMINE	143 A BB	902 3	472. 0.966	0.314	
83	S3#120	2,3,4,6-TETRACHLOROPHENO	232 A BB	898 3	155. 0.297	0.336	
84	S3#130	C580 DIETHYLPHTHALATE	149	3	NOT FOUND		
85	S3#135	C590 FLUORENE	166 A BB	927 3	3769. 1.298	1.869	
86	S3#140	C585 4-CHLOROPHENYL-PHEN	204	3	NOT FOUND		
87	S3#145	5-NITRO-O-TOLUIDINE	152 A VB	942 3	1772. 0.356	3.209	
88	S3#150	C595 4-NITROANILINE	138 A BB	949 3	396. 0.307	0.830	
89	S4# 10	C610 4,6-DINITRO-2-METHY	198	4	NOT FOUND		
90	S4# 15	C615 N-NITROSODIPHENYLAM	169 A VB	964 4	3698. 0.551	2.422	
91	S4# 20	C620 AZOBENZENE	77	4	NOT FOUND		
92	S4# 25	SYM-TRINITROBENZENE	75	4	NOT FOUND		
93	S4# 30	C625 4-BROMOPHENYL-PHENY	248	4	NOT FOUND		
94	S4# 35	PHENACETIN	108	4	NOT FOUND		
95	S4# 37	DIALLATE	234	4	NOT FOUND		
96	S4# 40	C630 HEXACHLOROBENZENE	284	4	NOT FOUND		
97	S4# 45	4-AMINOBIIPHENYL	169	4	NOT FOUND		
98	S4# 50	C635 PENTACHLOROPHENOL	266	4	NOT FOUND		
99	S4# 55	PRONAMIDE	173	4	NOT FOUND		
100	S4# 60	PENTACHLORONITROBENZENE	237 A BB	1064 4	72. 0.080	0.323	
101	S4# 65	C640 PHENANTHRENE	178 A BB	1091 4	11634. 1.033	4.063	
102	S4# 70	C645 ANTHRACENE	178	4	NOT FOUND		
103	S4# 75	2SEC BUTYL-4,6-DINITROPHE	211	4	NOT FOUND		
104	S4# 80	C647 CARBAZOLE	167 A BB	1149 4	472. 0.764	0.223	
105	S4# 85	C650 DI-N-BUTYLPHTHALATE	149	4	NOT FOUND		
106	S4#100	4-NITROQUINOLINE-1-OXIDE	190 A BB	1240 4	183. 0.073	0.910	
107	S4#105	METHAPYRILENE	58	4	NOT FOUND		
108	S4#106	ISODRIN	193 A VV	1272 4	2153. 0.127	6.130	
109	S4#110	C655 FLUORANTHENE	202 A BB	1300 4	591. 0.900	0.237	
110	S4#120	CHLOROBENZILATE	139	4	NOT FOUND		

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111	S5# 10	C710 BENZIDINE	184	5	NOT FOUND			
112	S5# 15	C715 PYRENE	202	A BB 1336	5	2966.	1.323	1.004
113	S5# 20	ARAMITE (#1)	185	5	NOT FOUND			
114	S5# 25	ARAMITE (#2)	185	5	NOT FOUND			
115	S5# 30	P-DIMETHYLAMINOAZOBENZEN	120	5	NOT FOUND			
116	S5# 35	3,3'-DIMETHYLBENZIDINE	212	5	NOT FOUND			
117	S5# 37	KEPONE	272	5	NOT FOUND			
118	S5# 40	C720 BUTYLBENZYLPHTHALAT	149	5	NOT FOUND			
119	S5# 45	2-ACETYLAMINOFLUORENE	181	5	NOT FOUND			
120	S5# 50	C730 BENZO(A)ANTHRACENE	228	A BB 1556	5	1877.	1.125	0.747
121	S5# 55	C725 3,3'-DICHLOROBENZID	252	5	NOT FOUND			
122	S5# 60	C740 CHRYSENE	228	A BB 1556	5	1877.	1.014	0.828
123	S5# 65	C745 BIS(2-ETHYLHEXYL)PH	149	5	NOT FOUND			
124	S5# 85	3-METHYLCHOLANTHRENE	268	5	NOT FOUND			
125	S6# 10	C760 DI-N-OCTYL PHTHALAT	149	6	NOT FOUND			
126	S6# 15	C765 BENZO(B)FLUORANTHEN	252	6	NOT FOUND			
127	S6# 20	7,12-DIMETHYLBENZANTHRAC	256	6	NOT FOUND			
128	S6# 25	C770 BENZO(K)FLUORANTHEN	252	6	NOT FOUND			
129	S6# 30	HEXACHLOROPHENE	196	A VB 1761	6	130.	0.004	17.527
130	S6# 35	C775 BENZO(A)PYRENE	252	6	NOT FOUND			
131	S6# 55	C780 INDENO(1,2,3-CD)PYR	276	6	NOT FOUND			
132	S6# 60	C785 DIBENZ(A,H)ANTHRACE	278	6	NOT FOUND			
133	S6# 65	C790 BENZO(G,H,I)PERYLEN	276	6	NOT FOUND			

QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297. =+200		4.28	200864. =+200		6.40	108695. =+200		9.71
Lower Limit	14324. =+ 50		3.28	50216. =+ 50		5.40	27174. =+ 50		8.71
Filename									
1 30068111	30555. = 106		3.78	109200. = 108		5.90	62156. = 114		9.23

IS# 1 = C130 1,4-DICHLOROBENZENE-D4  
 IS# 2 = C140 NAPHTHALENE-D8  
 IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039. =+200		12.58	124578. =+200		17.76	101500. =+200		20.33
Lower Limit	49260. =+ 50		11.58	31145. =+ 50		16.76	25375. =+ 50		19.33
Filename									
1 30068111	110919. = 112		12.10	89347. = 143		17.27	73934. = 145		19.83

IS# 4 = C160 PHENANTHRENE-D10  
 IS# 5 = C170 CHRYSENE-D12  
 IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

TARGET COMPOUND COMPARISON

COMPOUND: C450 NAPHTHALENE 128 52# 60 91-20-3

RAW DATA: 30068111 #533

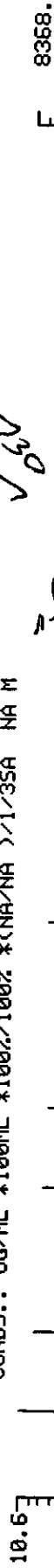
08/31/98 23:58

SAMPLE: S-NMS-5 1/35A/100M

CONDS.: UG/ML \*100Z/100Z \*(NA/NA >1/35A NA M

BASE M/Z: 41 RIC: 134400.

*✓ 30*



ENHANCED DATA: 30068111 #533



STANDARD FILE: ST16980831 #533

08/31/98 12:31

BASE M/Z: 128 RIC: 195840.



\*\*OUT\*\*

TARGET COMPOUND COMPARISON

COMPOUND: C470 2-METHYLNAPHTHALENE 142 S2#145 91-57-6

RAW DATA: 30068111 #653

08/31/98 23:58

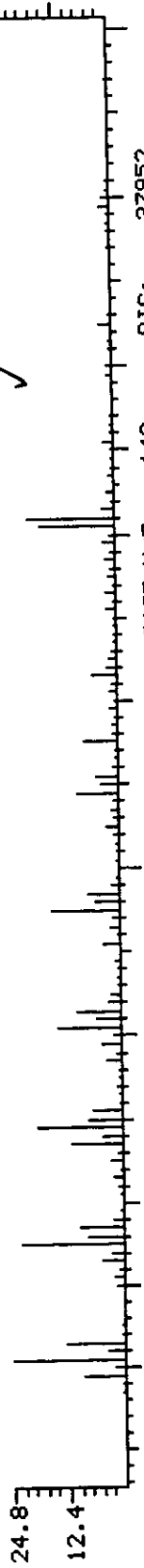
SAMPLE: S-MMS-5 1/3SA/100M

CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA) /1/3SA NA M

INST. ID: F16

BASE M/Z: 41 RIC: 194304.

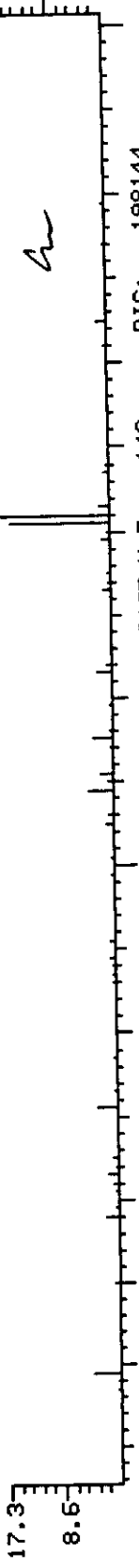
12368.



ENHANCED DATA: 30068111 #653

BASE M/Z: 142 RIC: 37952.

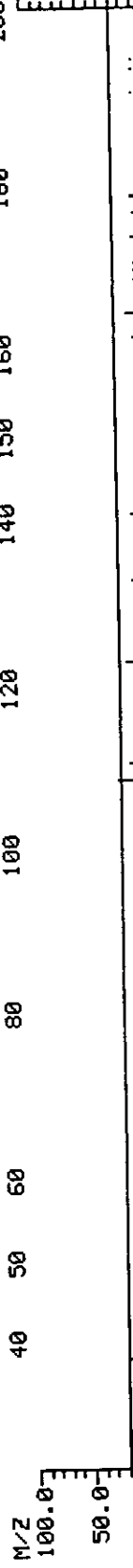
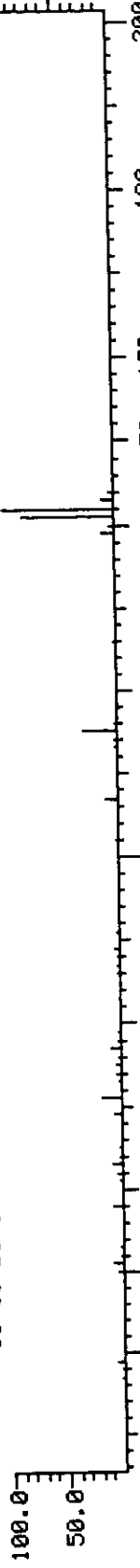
8608.



STANDARD FILE: ST16980831 #651

BASE M/Z: 142 RIC: 198144.

49792.



\*\*OUT\*\*

TARGET COMPOUND COMPARISON

COMPOUND: C590 FLUORENE

155 S3#135

86-73-7

RAW DATA: 30068111 #927

08/31/98 23:58

SAMPLE: S-MM5-5 1/35A/100M

INST. ID: F16

CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M

BASE M/Z: 41 RIC: 156160.

*125*  
*spk*



ENHANCED DATA: 30068111 #927



STANDARD FILE: ST16980831 #926

08/31/98 12:31

BASE M/Z: 81 RIC: 14336.



M/Z 100.0 50.0 0.0 -50.0 -100.0

BASE M/Z: 166 RIC: 203520.



\*\*OUT\*\*



TARGET COMPOUND COMPARISON

COMPOUND: C640 PHENANTHRENE 178 S4# 65 85-01-8

RAW DATA: 30068111 #1091

08/31/98 23:58

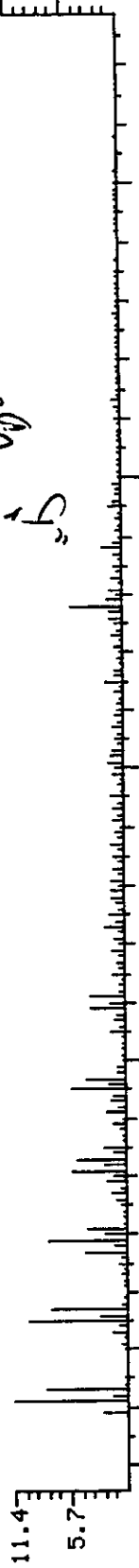
SAMPLE: S-MM5-5 1/35A/100M

CONDS.: UG/ML \*100Z/100Z \*(NA/NA) /1/35A NA M

*Handwritten:* b  
V  
b

BASE M/Z: 41 RIC: 155392.

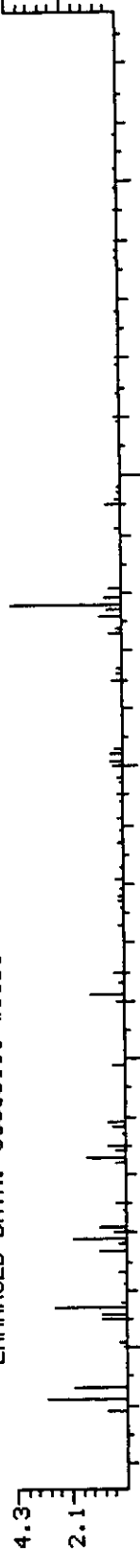
8352.



ENHANCED DATA: 30068111 #1091

BASE M/Z: 178 RIC: 27648.

3148.

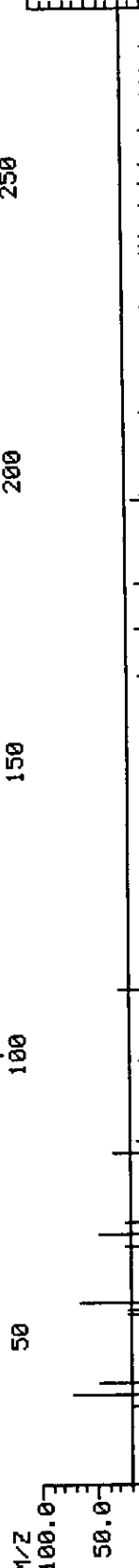
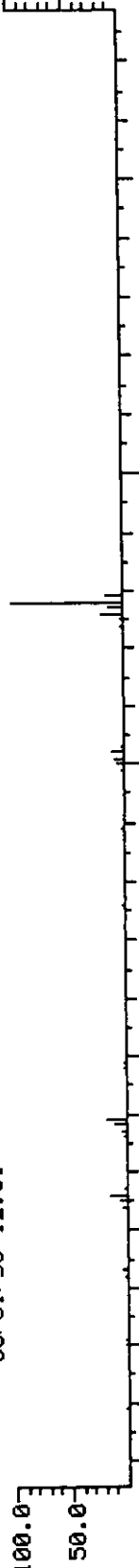


STANDARD FILE: ST16980831 #1091

08/31/98 12:31

BASE M/Z: 178 RIC: 204544.

73344.



\*\*OUT\*\*

Data Reduced by: 2 Date: 9/18/98  
Data Reviewed by:      Date:     

Data File: 30068111

QUANTERRA GC/MS TIC REPORT ( Part 1 )

Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
Analyst: DAT Date Analyzed: 08/31/98 23:58  
Run Factor: 300.

# SCAN	Concentration in Sample (UG/A)	CAS #	
1 444	11000.	<del>29812-79-1</del>	<del>HYDROXYLAMINE, O-DECYL-</del> unknown 5700
X 456	7100.	<del>2958-76-1</del>	<del>NAPHTHALENE, DECAHYDRO-2-METHYL-</del>
3 473	13000.	71899-38-2	9-EICOSYNE
4 557	9200.	1002-17-1	DECANE, 2,9-DIMETHYL- or isomer
5 570	9500.	17301-23-4	UNDECANE, 2,6-DIMETHYL-
X 600	7600.	00-00-0	UNKNOWN 5700
7 618	15000.	<del>90991-16-7</del>	<del>1,1'-BICYCLOHEXYL, 2-(1-METHYLETHYL), TRANS</del> unknown 5700
8 633	20000.	<del>26730-14-3</del>	<del>IRIDECANE, 7-METHYL</del> alkane 5746400
9 655	12000.	<del>36653-82-4</del>	<del>1-HEXADECANOL</del> unknown 5700
10 663	11000.	<del>1120-21-4</del>	<del>UNDECANE</del> alkane

11 679 10000. 00-00-0  
UNKNOWN

5700

12 701 12000. 00-00-0  
UNKNOWN

~~13~~ 715 8100. 00-00-0  
UNKNOWN

14 739 19000. 3891-98-3  
DODECANE, 2,6,10-TRIMETHYL- *or isomer*

15 764 25000. 629-59-4  
TETRADECANE

16 778 15000. 00-00-0  
UNKNOWN

5700

~~17~~ 809 7500. 00-00-0  
UNKNOWN

18 821 22000. 18344-37-1  
HEPTADECANE, 2,6,10,14-TETRAMETHYL- *or isomer*

19 859 25000. 17312-57-1  
DODECANE, 3-METHYL-

20 904 13000. 7390-81-0  
OXIRANE, HEXADECYL-

21 948 27000. ~~629-92-5~~  
NONADECANE

*alkane*

*5746400*

22 988 14000. ~~55030-62-1~~  
TRIDECANE, 4,8-DIMETHYL-

23 1036 28000. 54105-67-8  
HEPTADECANE, 2,6-DIMETHYL- *or isomer*

---

---

~~24~~ 1114 7500. ~~29812-79-1~~  
~~HYDROXYLAMINE, O-DECYL~~ *alkane*

---

---

25 1119 11000. 629-92-5  
NONADECANE *or isomer*

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## QUANTERRA GC/MS TIC REPORT ( Part 2 )

CONCENTRATION = AREA(TIC)\*CONC(IS)/AREA(IS)

#	FIT	PURITY	INT.	STD.	RT	RRT	AREA	HEIGHT	AMOUNT		LIB	LIB #
									AS ANALYZED			
									(UG/ML )			
1	934	591	2	4:56	0.535	320640.	84352.	37.754	NB	15969.		
2	971	771	2	5:04	0.549	201408.	45952.	23.715	NB	10432.		
3	906	673	2	5:15	0.570	358080.	80768.	42.163	NB	39230.		
4	958	860	2	6:11	0.671	259712.	77696.	30.580	NB	15357.		
5	941	828	2	6:20	0.687	268160.	77952.	31.575	NB	19054.		
6	827	659	2	6:40	0.723	216448.	37376.	25.486	UK	1.		
7	923	673	2	6:52	0.745	413696.	49810.	48.711	NB	24994.		
8	923	768	2	7:02	0.763	569856.	136550.	67.099	NB	22534.		
9	923	655	2	7:17	0.789	345108.	61049.	40.635	NB	32420.		
10	959	854	2	7:22	0.799	298496.	96768.	35.147	NB	11607.		
11	874	355	2	7:33	0.818	289024.	57344.	34.032	UK	1.		
12	887	427	2	7:47	0.845	350208.	52736.	41.236	UK	1.		
13	904	453	2	7:57	0.861	228352.	50432.	26.888	UK	1.		
14	956	817	2	8:13	0.890	537600.	135936.	63.301	NB	25991.		
15	950	710	2	8:29	0.920	719104.	193280.	84.672	NB	22530.		
16	837	424	2	8:39	0.937	431872.	61440.	50.852	UK	1.		
17	949	562	2	8:59	0.975	212992.	42240.	25.079	UK	1.		
18	967	804	2	9:07	0.989	613120.	137984.	72.193	NB	42196.		
19	953	759	2	9:33	1.035	706304.	171008.	83.165	NB	19007.		
20	880	641	2	10:03	1.089	359680.	56576.	42.351	NB	37444.		
21	958	828	2	10:32	1.142	754944.	160256.	88.892	NB	37465.		
22	923	808	3	10:59	0.908	519680.	129024.	45.516	NB	25996.		
23	973	865	3	11:31	0.952	1063170.	190464.	93.117	NB	37462.		
24	960	765	3	12:23	1.024	286976.	90112.	25.135	NB	15969.		
25	958	834	3	12:26	1.028	421376.	126720.	36.906	NB	37465.		

Library Search Data: 30068111 # 444 Base m/z: 43  
 08/31/98 23:58:00 + 4:56 Cali: 30068111 # 3 RIC: 73216.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYN8 searched for maximum PURITY  
 797 matched at least 7 of the 16 largest peaks in the unknown

- | Rank In. | Name                               |
|----------|------------------------------------|
| 1        | 15969 HYDROXYLAMINE, O-DECYL-      |
| 2        | 19523 1-DECANOL, 2-ETHYL-          |
| 3        | 5049 OXIRANE, (3,3-DIMETHYLBUTYL)- |
| 4        | 19015 DECANE, 2,5,6-TRIMETHYL-     |
| 5        | 12050 1-OCTANOL, 2,7-DIMETHYL-     |
| 6        | 11565 ISOCTANE, (ETHENYLOXY)-      |
| 7        | 14793 1-UNDECENE, 4-METHYL-        |
| 8        | 12074 1-HEPTANOL, 2-PROPYL-        |
| 9        | 12492 DECANE, 1-FLUORO-            |

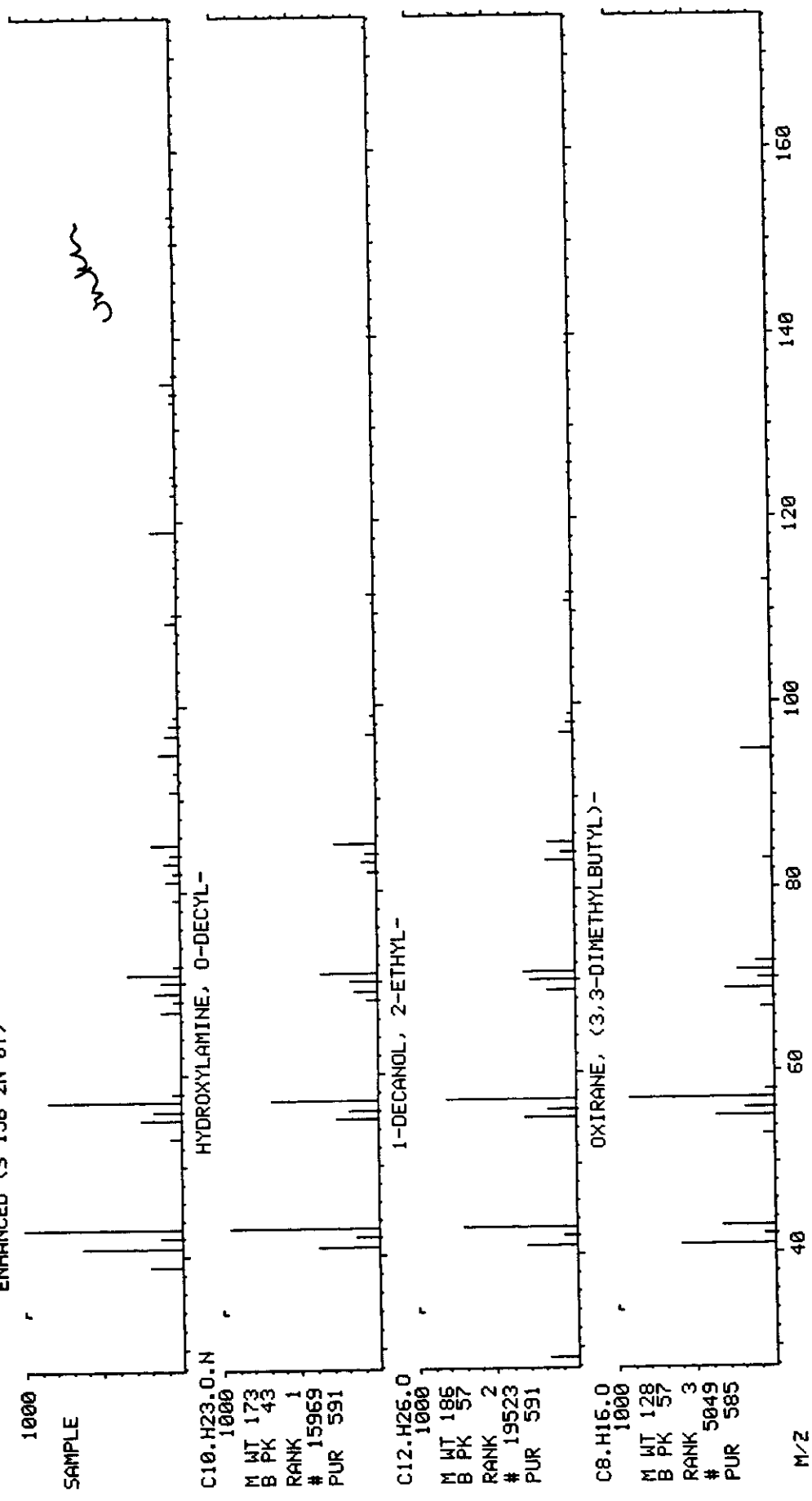
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H23.O.N	173	43	591	934	623
2	C12.H26.O	186	57	591	921	621
3	C8.H16.O	128	57	585	939	585
4	C13.H28	184	57	583	926	589
5	C10.H22.O	158	43	579	931	584
6	C10.H20.O	156	43	579	951	601
7	C12.H24	168	43	573	901	610
8	C10.H22.O	158	43	573	915	600
9	C10.H21.F	160	43	572	914	592

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	29812-79-1
2	—	—	—	—	21078-65-9
3	—	—	—	—	53907-77-0
4	—	—	—	—	62108-23-0
5	—	—	—	—	15250-22-3
6	—	—	—	—	37769-62-3
7	—	—	—	—	74630-39-0
8	—	—	—	—	10042-59-8
9	—	—	—	—	334-56-5

BASE M/Z: 43  
RIC: 73216.

DATA: 30068111 # 444  
CALI: 30068111 # 3

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 4:56  
SAMPLE: S-MMS-5 1/35A/100M INST. ID: F15  
CONDS.: UG/ML \*100ML \*100Z \*NA/NA >/1/35A NA M  
ENHANCED (S 158 2N 0T)



14101

Library Search Data: 30068111 # 456 Base m/z: 81  
 08/31/98 23:58:00 + 5:04 Cali: 30068111 # 3 RIC: 40192.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 475 matched at least 7 of the 16 largest peaks in the unknown

Rank In. Name  
 1 10432 NAPHTHALENE, DECAHYDRO-2-METHYL-  
 2 10376 BICYCLO[4.1.0]HEPTAN-3-ONE, 4,7,7-TRIMETHYL-, [1R-(1.ALPHA.,4.ALPHA.\*  
 3 10357 BICYCLO[4.1.0]HEPTAN-3-ONE, 4,7,7-TRIMETHYL-, [1R-(1.ALPHA.,4.BETA.\*  
 4 10304 CYCLOHEXANONE, 2-METHYL-5-(1-METHYLETHENYL)-  
 5 10349 2H-INDEN-2-ONE, OCTAHYDRO-3A-METHYL-, CIS-  
 6 10358 CYCLOHEXANONE, 2-METHYL-5-(1-METHYLETHENYL)-, TRANS-  
 7 10352 CYCLOHEXANONE, 5-METHYL-2-(1-METHYLETHYLIDENE)-  
 8 17938 1,12-TRIDECADIENE  
 9 10355 CYCLOHEXANONE, 5-METHYL-2-(1-METHYLETHENYL)-, TRANS-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C11.H20	152	95	771	971	771
2	C10.H16.O	152	81	757	948	757
3	C10.H16.O	152	81	752	941	752
4	C10.H16.O	152	67	715	938	715
5	C10.H16.O	152	95	714	914	714
6	C10.H16.O	152	95	709	930	709
7	C10.H16.O	152	81	704	934	704
8	C13.H24	180	55	703	851	713
9	C10.H16.O	152	81	695	925	695

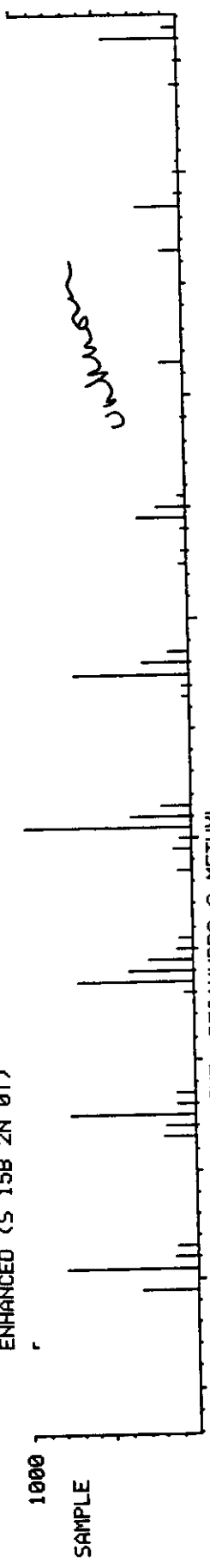
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	2958-76-1
2	---	---	---	---	4176-04-9
3	---	---	---	---	4176-01-6
4	---	---	---	---	7764-50-3
5	---	---	---	---	13351-29-6
6	---	---	---	---	5948-04-9
7	---	---	---	---	15932-80-6
8	---	---	---	---	21964-48-7
9	---	---	---	---	29606-79-9



DATA: 30068111 # 456  
CALI: 30068111 # 3

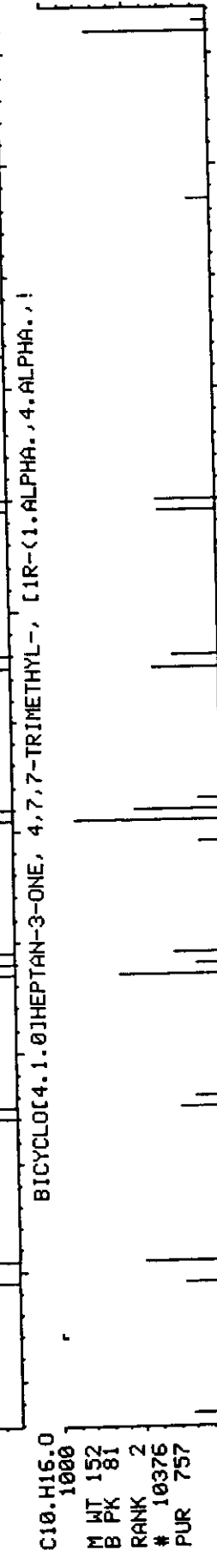
BASE M/Z: 81  
RIC: 40192.

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 5:04  
SAMPLE: S-MMS-5 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100ML \*100Z \*(NA/NA) /1/35A NA M  
ENHANCED (S 15B 2N 0T)



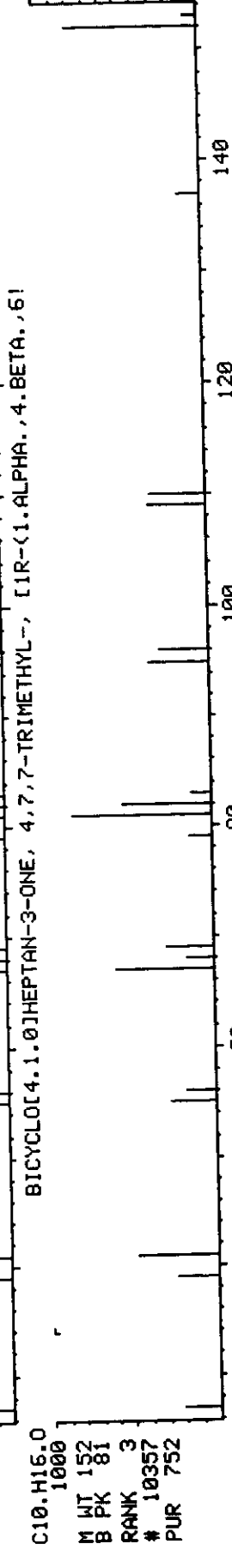
NAPHTHALENE, DECAHYDRO-2-METHYL-

C11.H20  
1000  
M WT 152  
B PK 95  
RANK 1  
# 10432  
PUR 771



BICYCLO[4.1.0]HEPTAN-3-ONE, 4,7,7-TRIMETHYL-, [1R-(1.ALPHA.,4.ALPHA.,6)]

C10.H16.O  
1000  
M WT 152  
B PK 81  
RANK 2  
# 10376  
PUR 757



BICYCLO[4.1.0]HEPTAN-3-ONE, 4,7,7-TRIMETHYL-, [1R-(1.ALPHA.,4.BETA.,6)]

C10.H16.O  
1000  
M WT 152  
B PK 81  
RANK 3  
# 10357  
PUR 752

M/Z

1417

Library Search                    Data: 30068111 # 473            Base m/z: 41  
 08/31/98 23:58:00 + 5:16        Cali: 30068111 # 3            RIC: 70016.  
 Sample: S-MMS-5 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 459 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	39230 9-EICOSYNE
2	34800 (R)-(-)-(Z)-14-METHYL-8-HEXADECEN-1-OL
3	22482 2-TRIDECEN-1-OL, (E)-
4	25971 OXIRANE, DODECYL-
5	36677 7-OCTADECYNE, 2-METHYL-
6	29222 (Z)6-PENTADECEN-1-OL
7	34010 3-OCTADECYNE
8	37449 OCTADECANAL
9	28267 3-HEXADECYNE

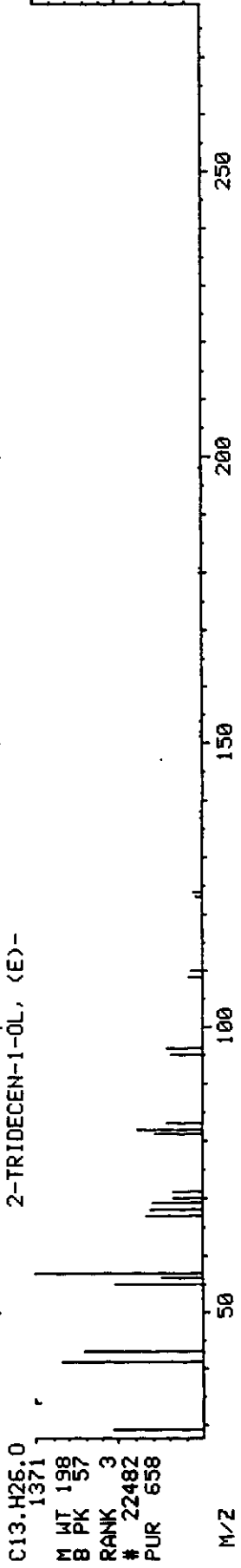
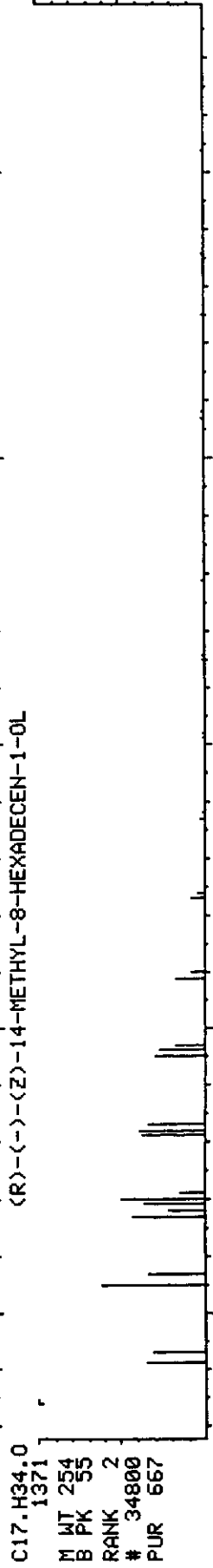
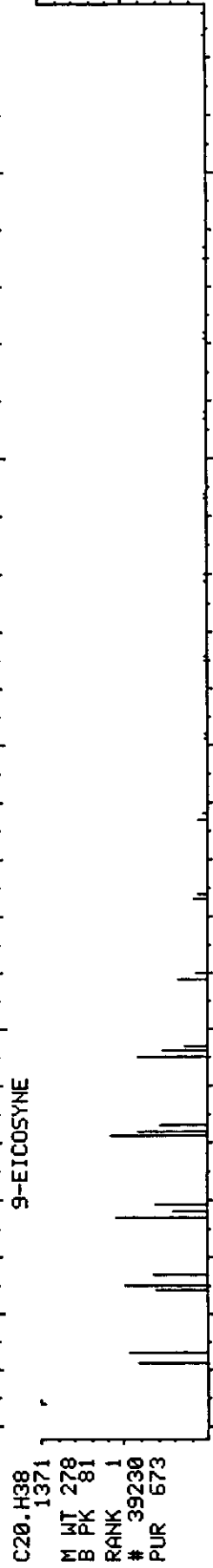
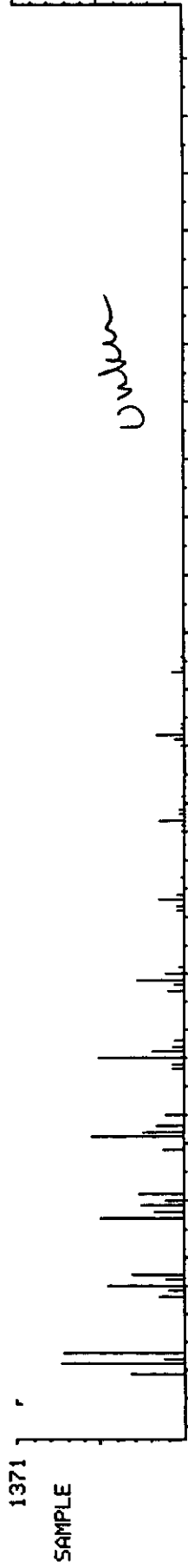
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C20.H38	278	81	673	906	690
2	C17.H34.O	254	55	667	857	676
3	C13.H26.O	198	57	658	850	664
4	C14.H28.O	212	41	658	906	658
5	C19.H36	264	81	649	906	666
6	C15.H30.O	226	67	645	869	664
7	C18.H34	250	67	638	926	644
8	C18.H36.O	268	43	637	828	680
9	C16.H30	222	67	637	934	642

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	71899-38-2
2	---	---	---	---	30689-78-2
3	---	---	---	---	74962-98-4
4	---	---	---	---	3234-28-4
5	---	---	---	---	35354-38-2
6	---	---	---	---	68797-95-5
7	---	---	---	---	61886-64-4
8	---	---	---	---	638-66-4
9	---	---	---	---	61886-62-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 5:16  
 SAMPLE: S-MMS-5 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 473  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 70016.



Library Search Data: 30068111 # 557 Base m/z: 43  
 08/31/98 23:58:00 + 6:12 Cali: 30068111 # 3 RIC: 75008.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 731 matched at least 7 of the 16 largest peaks in the unknown

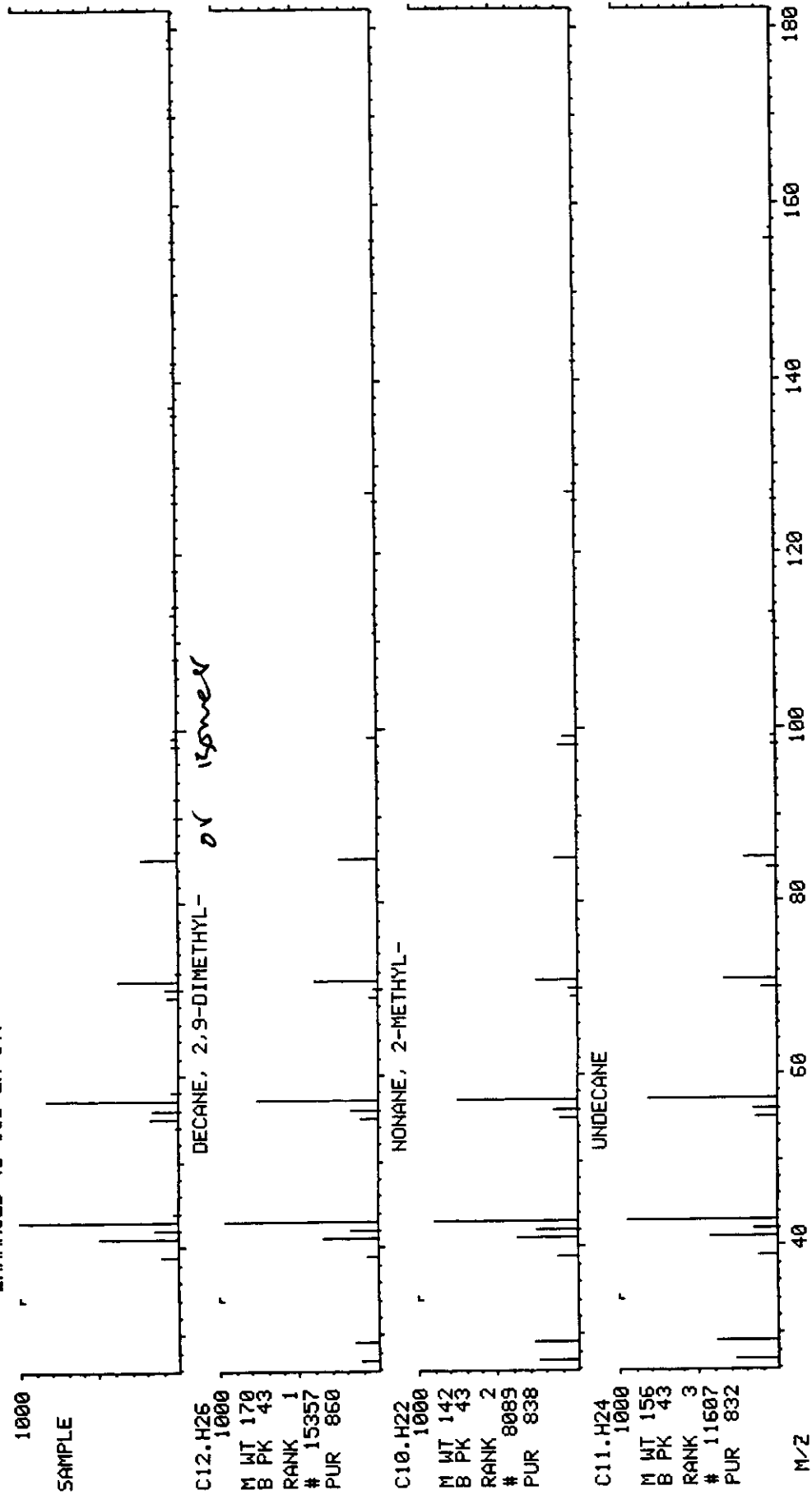
Rank In. Name  
 1 15357 DECANE, 2,9-DIMETHYL-  
 2 8089 NONANE, 2-METHYL-  
 3 11607 UNDECANE  
 4 19013 DECANE, 2,5,9-TRIMETHYL-  
 5 15343 DODECANE  
 6 8074 OCTANE, 2,7-DIMETHYL-  
 7 11602 OCTANE, 2,4,6-TRIMETHYL-  
 8 19006 DECANE, 6-ETHYL-2-METHYL-  
 9 19035 DODECANE, 2-METHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C12.H26	170	43	860	958	870
2	C10.H22	142	43	838	941	842
3	C11.H24	156	43	832	941	860
4	C13.H28	184	57	830	916	840
5	C12.H26	170	57	829	923	868
6	C10.H22	142	43	828	917	828
7	C11.H24	156	57	825	944	844
8	C13.H28	184	57	824	930	847
9	C13.H28	184	43	819	932	848

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	1002-17-1
2	---	---	---	---	871-83-0
3	---	---	---	---	1120-21-4
4	---	---	---	---	62108-22-9
5	---	---	---	---	112-40-3
6	---	---	---	---	1072-16-8
7	---	---	---	---	62016-37-9
8	---	---	---	---	62108-21-8
9	---	---	---	---	1560-97-0

DATA: 30068111 # 557  
CALI: 30068111 # 3  
BASE M/Z: 43  
RIC: 75008.

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 6:12  
SAMPLE: 5-MM5-5 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/35A NA M  
ENHANCED (S 15B 2N 0T)



Library Search                      Data: 30068111 # 570                      Base m/z: 57  
 08/31/98 23:58:00 + 6:20            Cali: 30068111 # 3                      RIC: 75520.  
 Sample: S-MM5-5 1/3SA/100M            INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 718 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	19054 UNDECANE, 2,6-DIMETHYL-
2	19013 DECANE, 2,5,9-TRIMETHYL-
3	8104 OCTANE, 3,5-DIMETHYL-
4	19026 DECANE, 2,6,8-TRIMETHYL-
5	8077 HEPTANE, 2,3,5-TRIMETHYL-
6	19015 DECANE, 2,5,6-TRIMETHYL-
7	8092 HEPTANE, 2,3,6-TRIMETHYL-
8	19006 DECANE, 6-ETHYL-2-METHYL-
9	5154 HEXANE, 4-ETHYL-2-METHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C13.H28	184	57	828	941	861
2	C13.H28	184	57	818	962	828
3	C10.H22	142	57	808	953	816
4	C13.H28	184	57	802	948	823
5	C10.H22	142	57	797	950	797
6	C13.H28	184	57	797	928	806
7	C10.H22	142	57	791	946	791
8	C13.H28	184	57	787	935	796
9	C9.H20	128	57	786	941	792

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	17301-23-4
2	---	---	---	---	62108-22-9
3	---	---	---	---	15869-93-9
4	---	---	---	---	62108-26-3
5	---	---	---	---	20278-85-7
6	---	---	---	---	62108-23-0
7	---	---	---	---	4032-93-3
8	---	---	---	---	62108-21-8
9	---	---	---	---	3074-75-7

BASE M/Z: 57  
RIC: 75520.

DATA: 30068111 # 570  
CALI: 30068111 # 3

MID LIBRARY SEARCH <LIBRARYNB>

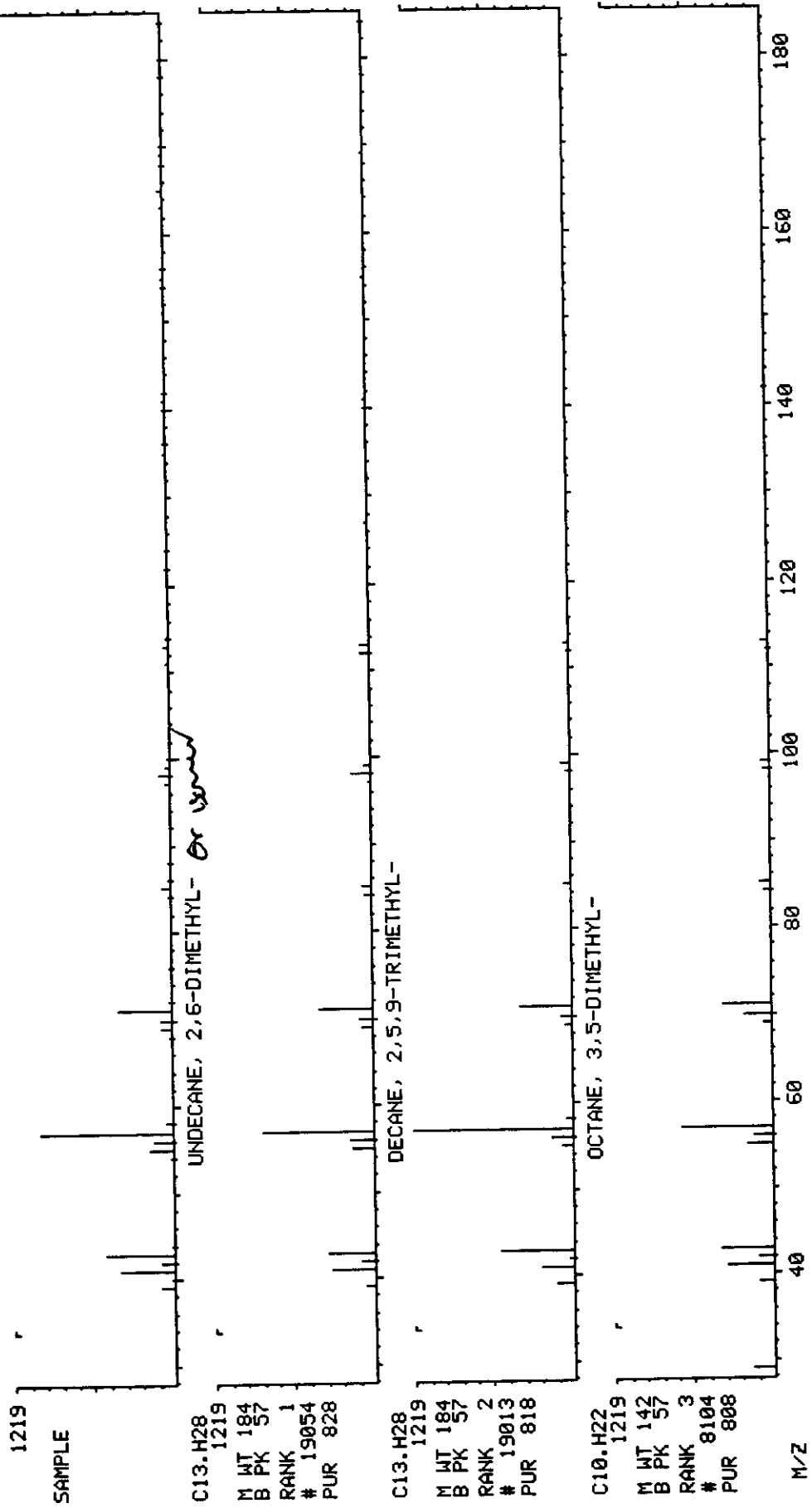
08/31/98 23:58:00 + 6:20

SAMPLE: S-MM5-5 1/35A/100M

INST. ID: F16

CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA) /1/35A NA M

ENHANCED (S 15B 2N 0T)



1219  
184  
57  
8104

Library Search                      Data: 30068111 # 600                      Base m/z: 41  
 08/31/98 23:58:00 + 6:40            Cali: 30068111 # 3                      RIC: 53760.  
 Sample: S-MM5-5 1/3SA/100M            INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 340 matched at least 7 of the 16 largest peaks in the unknown

Rank In.            Name  
 1 39230 9-EICOSYNE  
 2 17938 1,12-TRIDECADIENE  
 3 21421 1,13-TETRADECADIENE  
 4 25960 CIS-9-TETRADECEN-1-OL  
 5 44549 ETHANOL, 2-(9-OCTADECENYLOXY)-, (Z)-  
 6 25963 CIS-11-TETRADECEN-1-OL  
 7 22481 CYCLODODECANEMETHANOL  
 8 29222 (Z)6-PENTADECEN-1-OL  
 9 29229 CYCLOPENTADECANOL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C20.H38	278	81	659	827	685
2	C13.H24	180	55	626	872	636
3	C14.H26	194	55	616	810	636
4	C14.H28.O	212	55	614	860	628
5	C20.H40.O2	312	55	612	758	683
6	C14.H28.O	212	41	609	805	630
7	C13.H26.O	198	55	606	771	635
8	C15.H30.O	226	67	606	824	636
9	C15.H30.O	226	55	606	776	658

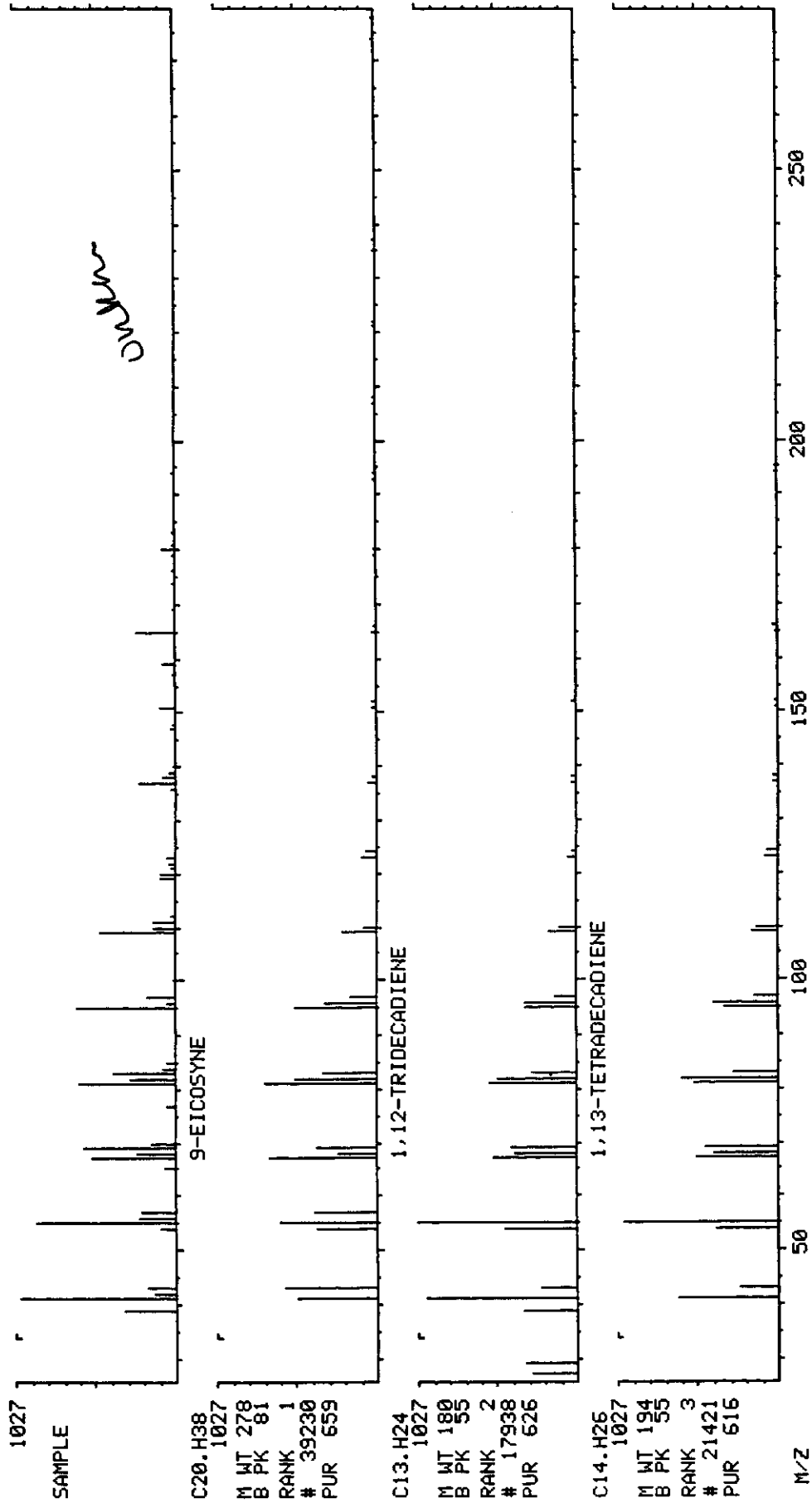
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	71899-38-2
2	---	---	---	---	21964-48-7
3	---	---	---	---	21964-49-8
4	---	---	---	---	35153-15-2
5	---	---	---	---	5353-25-3
6	---	---	---	---	34010-15-6
7	---	---	---	---	1892-12-2
8	---	---	---	---	68797-95-5
9	---	---	---	---	4727-17-7



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 6:40  
 SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068111 # 600  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 53750.



Library Search                    Data: 30068111 # 618            Base m/z: 41  
 08/31/98 23:58:00 + 6:52        Cali: 30068111 # 3            RIC: 54528.  
 Sample: S-MM5-5 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 375 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1 24994	1,1'-BICYCLOHEXYL, 2-(1-METHYLETHYL)-, TRANS-
2 24993	1,1'-BICYCLOHEXYL, 2-(1-METHYLETHYL)-, CIS-
3 33979	BORINIC ACID, DIETHYL-, 1-CYCLODODECEN-1-YL ESTER
4 24996	1,1'-BICYCLOHEXYL, 2-PROPYL-, CIS-
5 24997	1,1'-BICYCLOHEXYL, 2-PROPYL-, TRANS-
6 18965	11-DODECENOL
7 22482	2-TRIDECEN-1-OL, (E)-
8 18944	OXIRANE, DECYL-
9 24990	MUROLANE-B

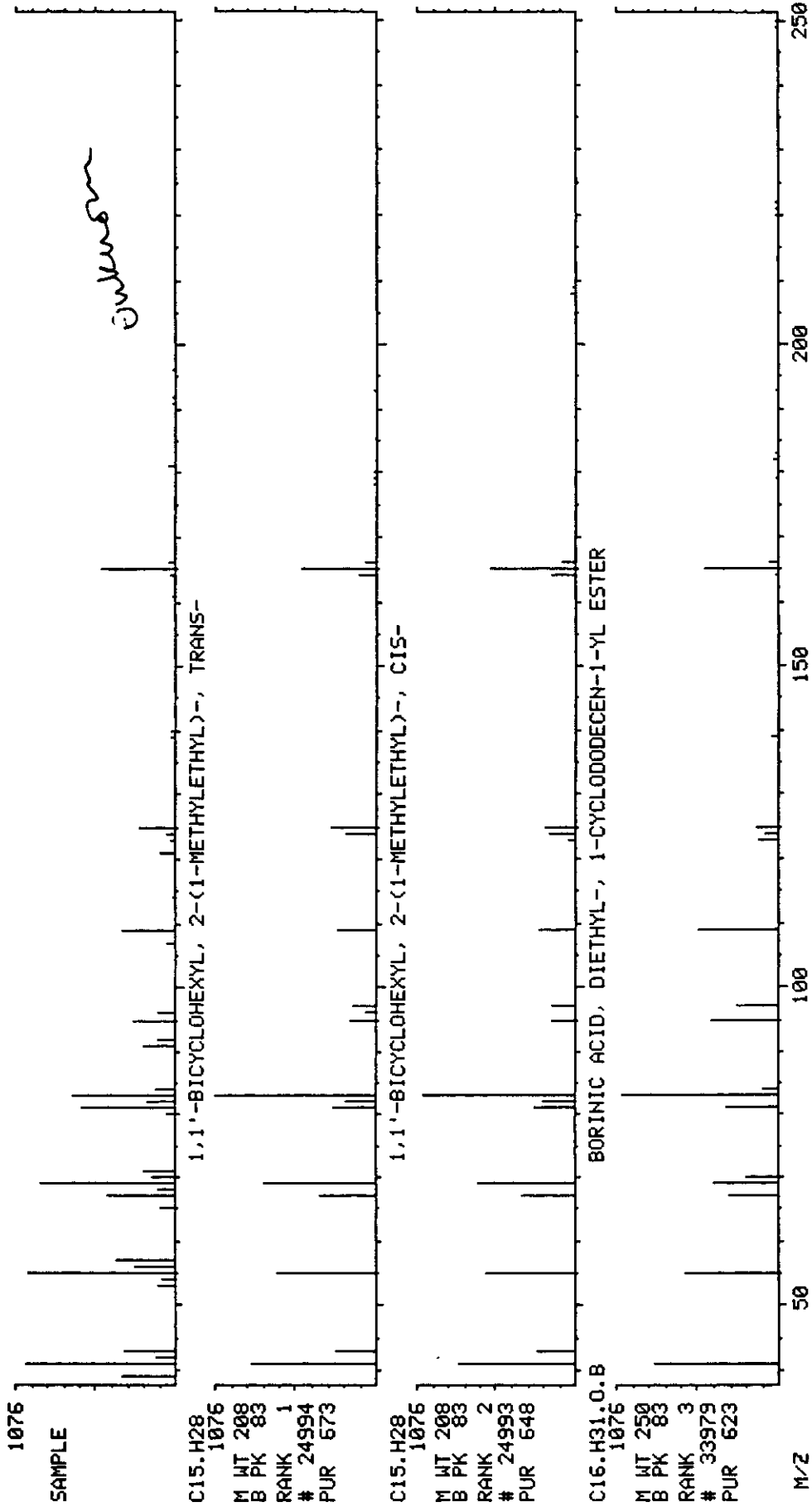
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C15.H28	208	83	673	923	693
2	C15.H28	208	83	648	900	676
3	C16.H31.O.B	250	83	623	875	675
4	C15.H28	208	69	576	786	609
5	C15.H28	208	41	568	774	595
6	C12.H24.O	184	55	558	879	558
7	C13.H26.O	198	57	553	823	581
8	C12.H24.O	184	41	547	841	570
9	C15.H28	208	109	546	812	582

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	50991-16-7
2	---	---	---	---	50991-15-6
3	---	---	---	---	61142-73-2
4	---	---	---	---	54934-88-2
5	---	---	---	---	54934-89-3
6	---	---	---	---	35289-31-7
7	---	---	---	---	74962-98-4
8	---	---	---	---	2855-19-8
9	---	---	---	---	- -

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:58:00 + 6:52  
 SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 618  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 54528.



Library Search                    Data: 30068111 # 633            Base m/z: 57  
 08/31/98 23:58:00 + 7:02        Cali: 30068111 # 3            RIC: 127872.  
 Sample: S-MM5-5 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 781 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 22534 TRIDECANE, 7-METHYL-  
 2 14799 1-DECENE, 3,4-DIMETHYL-  
 3 37462 HEPTADECANE, 2,6-DIMETHYL-  
 4 19523 1-DECANOL, 2-ETHYL-  
 5 8539 1-PENTANOL, 4-METHYL-2-PROPYL-  
 6 14751 2-UNDECENE, 5-METHYL-  
 7 26408 2-ETHYL-1-DODECANOL  
 8 14793 1-UNDECENE, 4-METHYL-  
 9 8071 NONANE, 3-METHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C14.H30	198	57	768	923	774
2	C12.H24	168	57	763	949	767
3	C19.H40	268	57	754	922	787
4	C12.H26.O	186	57	749	928	764
5	C9.H20.O	144	57	746	981	746
6	C12.H24	168	57	746	958	746
7	C14.H30.O	214	57	734	906	740
8	C12.H24	168	43	728	936	728
9	C10.H22	142	57	728	949	734

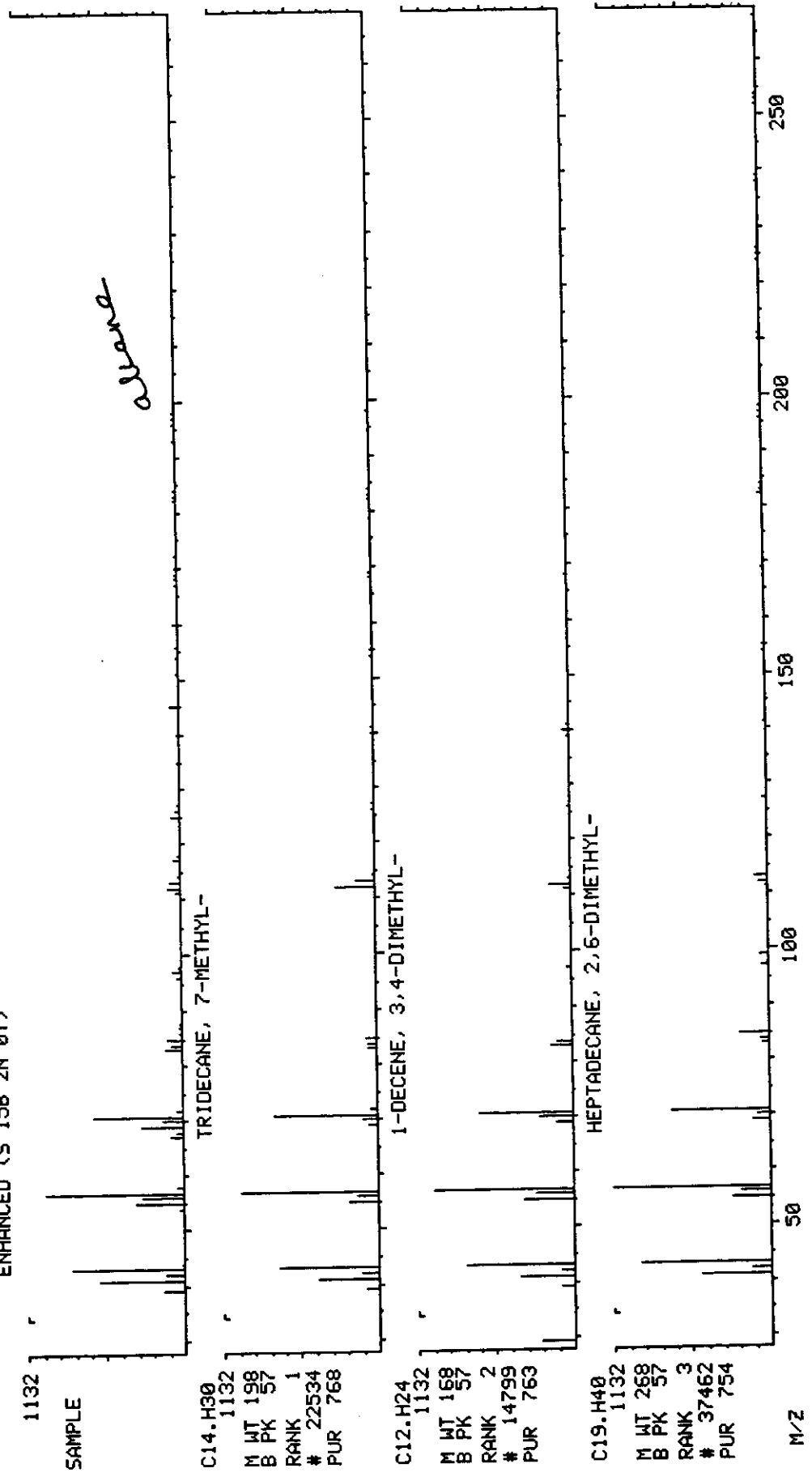
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	26730-14-3
2	---	---	---	---	50871-03-9
3	---	---	---	---	54105-67-8
4	---	---	---	---	21078-65-9
5	---	---	---	---	54004-41-0
6	---	---	---	---	56851-34-4
7	---	---	---	---	- -
8	---	---	---	---	74630-39-0
9	---	---	---	---	5911-04-6

DATA: 30068111 # 633  
CALI: 30068111 # 3

BASE M/Z: 57  
RIC: 127872.

MID LIBRARY SEARCH <LIBRARYNB>  
08/31/98 23:58:00 + 7:02  
SAMPLE: S-MMS-5 1/35A/100M  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 158 2N 0T)

INST. ID: F16



Library Search                    Data: 30068111 # 655            Base m/z: 41  
 08/31/98 23:58:00 + 7:17        Cali: 30068111 # 3            RIC: 70912.  
 Sample: S-MM5-5 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 744 matched at least 7 of the 16 largest peaks in the unknown

Rank In.        Name  
 1 32420 1-HEXADECANOL  
 2 28772 7-HEXADECENE, (Z)-  
 3 34410 3-OCTADECENE, (E)-  
 4 21960 CYCLOTETRADECANE  
 5 34411 5-OCTADECENE, (E)-  
 6 40233 NONADECANOL  
 7 28768 3-HEXADECENE, (Z)-  
 8 7515 2-NONENAL, (E)-  
 9 14780 CYCLODODECANE

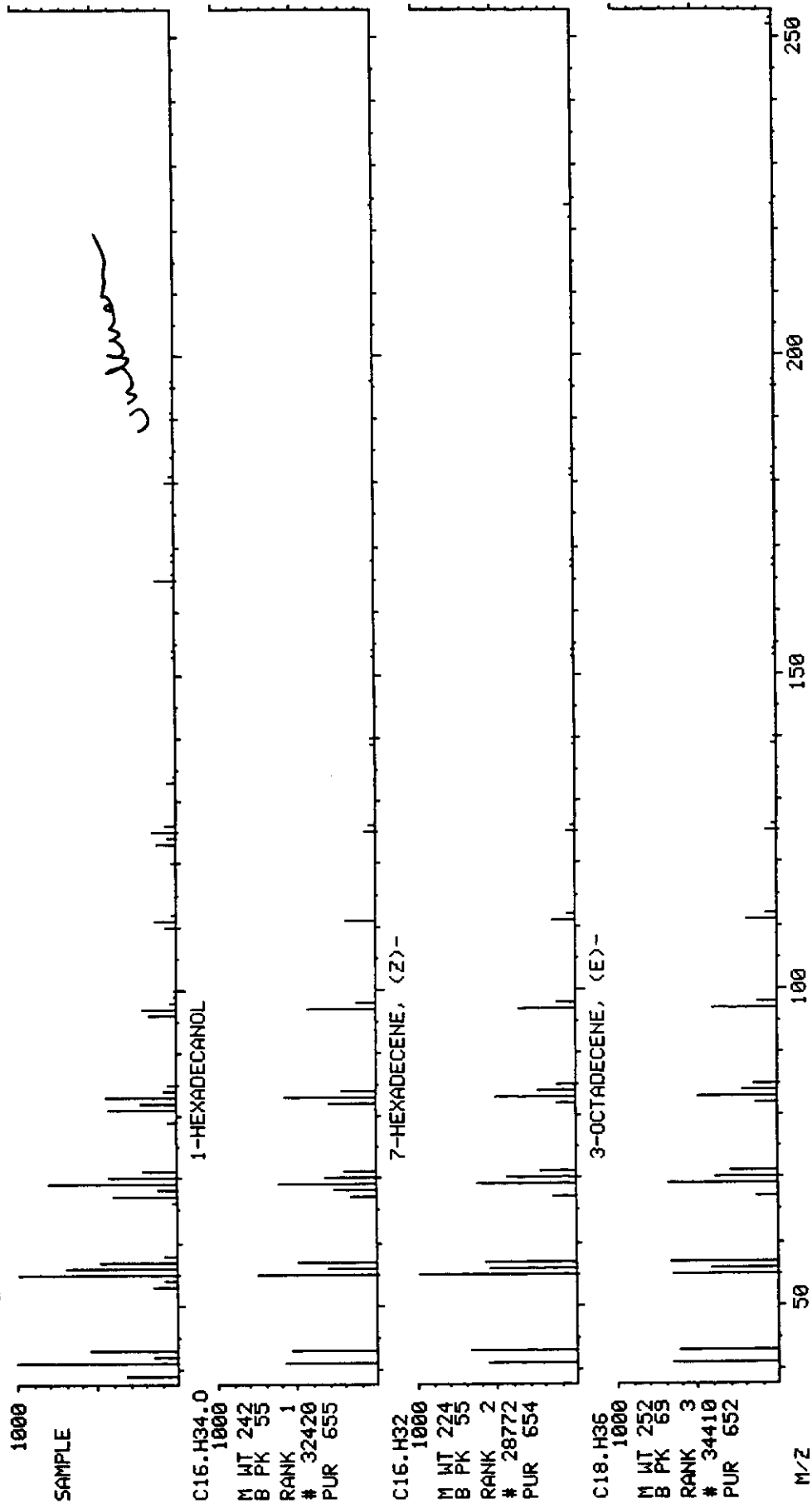
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H34.O	242	55	655	923	667
2	C16.H32	224	55	654	911	675
3	C18.H36	252	69	652	891	674
4	C14.H28	196	55	652	911	671
5	C18.H36	252	55	652	908	673
6	C19.H40.O	284	43	650	881	682
7	C16.H32	224	55	650	917	673
8	C9.H16.O	140	41	649	927	649
9	C12.H24	168	55	649	933	662

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	36653-82-4
2	---	---	---	---	35507-09-6
3	---	---	---	---	7206-19-1
4	---	---	---	---	295-17-0
5	---	---	---	---	7206-21-5
6	---	---	---	---	52783-43-4
7	---	---	---	---	34303-81-6
8	---	---	---	---	18829-56-6
9	---	---	---	---	294-62-2

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 7:17  
 SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UC/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 655  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 70912.



Library Search Data: 30068111 # 663 Base m/z: 57  
 08/31/98 23:58:00 + 7:22 Cali: 30068111 # 3 RIC: 107264.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 560 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	11607 UNDECANE
2	19013 DECANE, 2,5,9-TRIMETHYL-
3	19026 DECANE, 2,6,8-TRIMETHYL-
4	15357 DECANE, 2,9-DIMETHYL-
5	11602 OCTANE, 2,4,6-TRIMETHYL-
6	8104 OCTANE, 3,5-DIMETHYL-
7	8089 NONANE, 2-METHYL-
8	5159 NONANE
9	8074 OCTANE, 2,7-DIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C11.H24	156	43	854	959	859
2	C13.H28	184	57	815	914	829
3	C13.H28	184	57	813	911	839
4	C12.H26	170	43	811	932	841
5	C11.H24	156	57	801	933	822
6	C10.H22	142	57	798	902	826
7	C10.H22	142	43	797	913	827
8	C9.H20	128	43	795	914	834
9	C10.H22	142	43	794	901	807

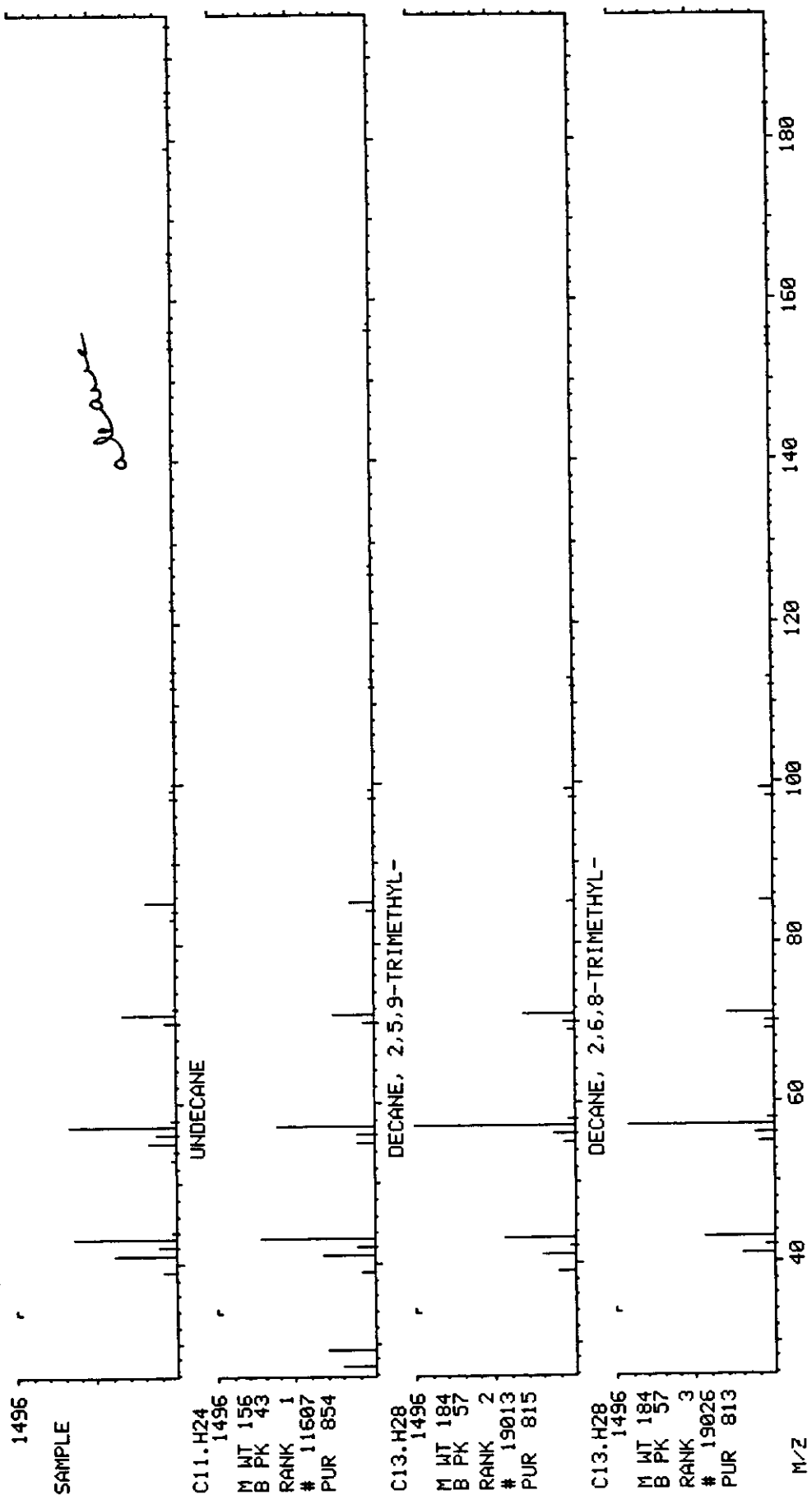
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	1120-21-4
2	---	---	---	---	62108-22-9
3	---	---	---	---	62108-26-3
4	---	---	---	---	1002-17-1
5	---	---	---	---	62016-37-9
6	---	---	---	---	15869-93-9
7	---	---	---	---	871-83-0
8	---	---	---	---	111-84-2
9	---	---	---	---	1072-16-8



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 7:22  
 SAMPLE: S-MMS-5 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100% /100% \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 663  
 CALI: 30068111 # 3

BASE M/Z: 57  
 RIC: 107264.



1496

SAMPLE

C11.H24  
 1496  
 M WT 156  
 B PK 43  
 RANK 1  
 # 11607  
 PUR 854

C13.H28  
 1496  
 M WT 184  
 B PK 57  
 RANK 2  
 # 19013  
 PUR 815

C13.H28  
 1496  
 M WT 184  
 B PK 57  
 RANK 3  
 # 19026  
 PUR 813

M/Z

1433

Library Search                    Data: 30068111 # 679            Base m/z: 57  
 08/31/98 23:58:00 + 7:33        Cali: 30068111 # 3            RIC: 62656.  
 Sample: S-MM5-5    1/3SA/100M            INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 324 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	5159 NONANE
2	37251 1-IOOO-2-METHYLNONANE
3	40866 OCTADECANE, 1-CHLORO-
4	5151 HEPTANE, 3,4-DIMETHYL-
5	19826 OCTANETHIOIC ACID, S-ETHYL ESTER
6	30843 4,4,6-TRIMETHYL-6-PHENYLTETRAHYDRO-1,3-OXAZINE-2-THIONE
7	11607 UNDECANE
8	35931 HEXADECANE, 1-CHLORO-
9	5154 HEXANE, 4-ETHYL-2-METHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C9.H20	128	43	355	874	375
2	C10.H21.I	268	43	353	871	381
3	C18.H37.CL	288	57	339	782	416
4	C9.H20	128	43	334	836	349
5	C10.H20.O.S	188	57	334	744	345
6	C13.H17.O.N.S	235	118	334	676	441
7	C11.H24	156	43	333	852	367
8	C16.H33.CL	260	57	333	774	414
9	C9.H20	128	57	332	831	364

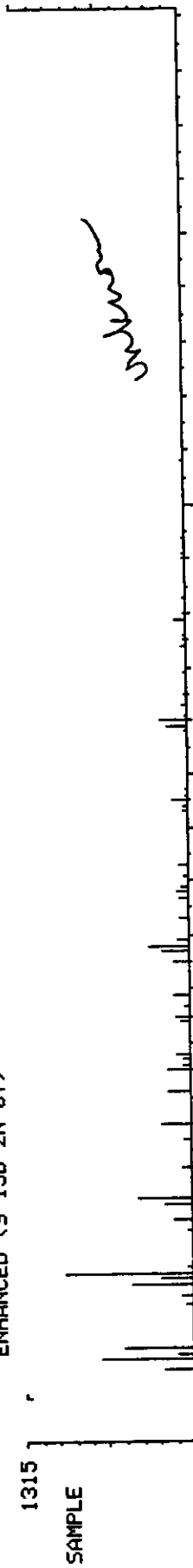
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	111-84-2
2	---	---	---	---	- -
3	---	---	---	---	3386-33-2
4	---	---	---	---	922-28-1
5	---	---	---	---	2432-84-0
6	---	---	---	---	- -
7	---	---	---	---	1120-21-4
8	---	---	---	---	4860-03-1
9	---	---	---	---	3074-75-7

DATA: 30068111 # 679  
CALI: 30068111 # 3

BASE M/Z: 57  
RIC: 62656.

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 7:33  
SAMPLE: 5-MM5-5 1/35A/100M  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA) >/1/35A NA M  
ENHANCED (S 15B 2N 0T)

INST. ID: F16



NONANE

C9.H20  
1315  
M WT 128  
B PK 43  
RANK 1  
# 5159  
PUR 355

1-IODO-2-METHYLNONANE

C10.H21.I  
1315  
M WT 268  
B PK 43  
RANK 2  
# 37251  
PUR 353

OCTADECANE, 1-CHLORO-

C18.H37.CL  
1315  
M WT 288  
B PK 57  
RANK 3  
# 40866  
PUR 339

M/Z

Library Search Data: 30068111 # 701 Base m/z: 41  
 08/31/98 23:58:00 + 7:48 Cali: 30068111 # 3 RIC: 54272.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYN8 searched for maximum PURITY  
 976 matched at least 5 of the 16 largest peaks in the unknown

Rank In. Name  
 1 10423 3-UNDECYNE  
 2 14158 3-DODECYNE  
 3 28273 8-HEXADECYNE  
 4 6996 1-ETHYNYL-1-CYCLOHEPTANOL  
 5 7028 1,3-HEXADIENE, 3-ETHYL-2,5-DIMETHYL-  
 6 28267 3-HEXADECYNE  
 7 10438 1,4-UNDECADIENE, (Z)-  
 8 10408 THUJONE  
 9 18371 5,7-DIMETHYLOCTAHYDROCOUMARIN

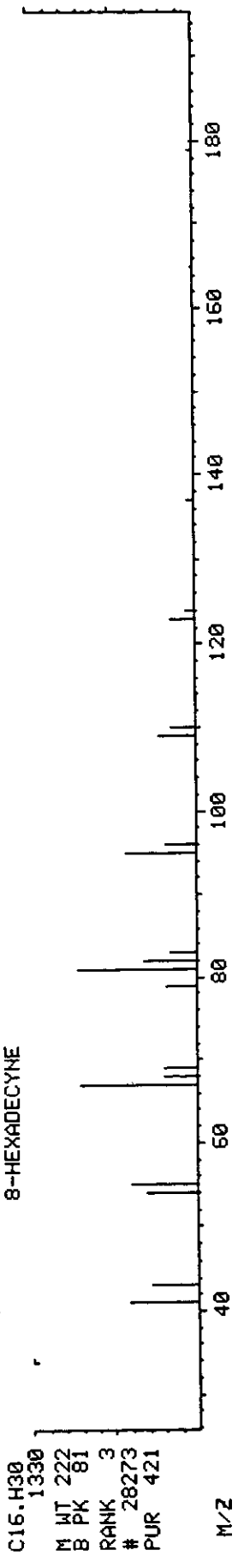
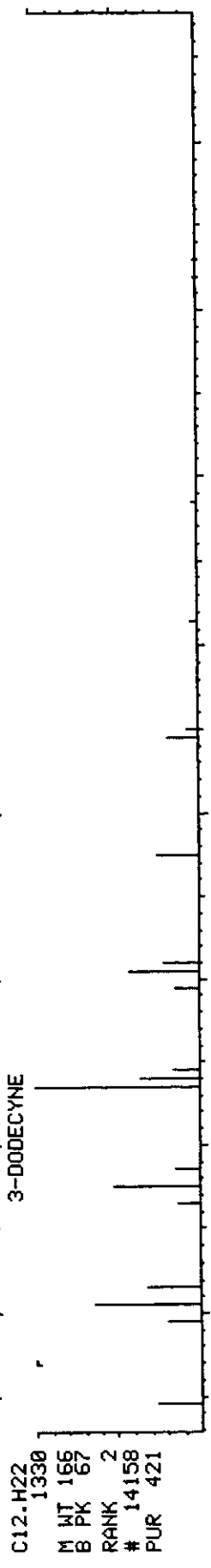
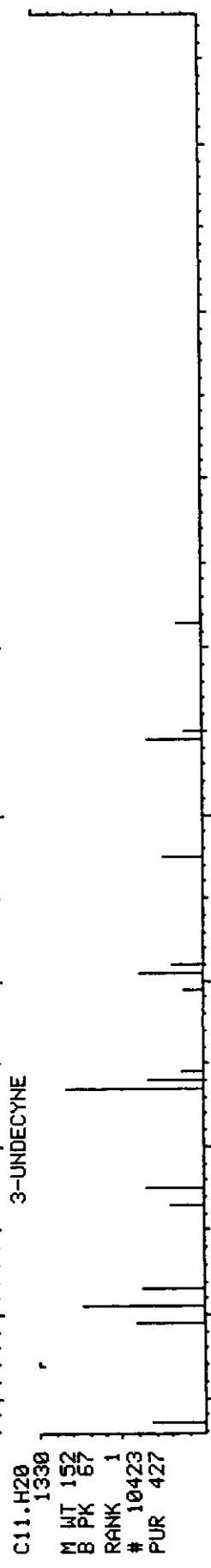
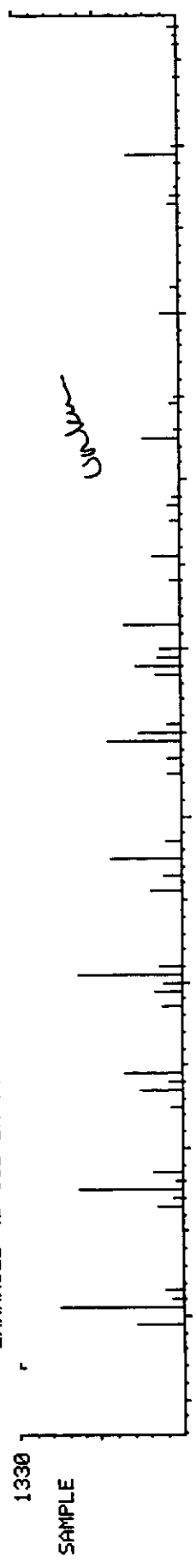
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C11.H20	152	67	427	887	427
2	C12.H22	166	67	421	831	422
3	C16.H30	222	81	421	764	467
4	C9.H14.O	138	81	417	836	438
5	C10.H18	138	67	414	859	414
6	C16.H30	222	67	408	713	437
7	C11.H20	152	67	407	889	431
8	C10.H16.O	152	81	404	896	416
9	C11.H18.O2	182	95	397	827	432

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	60212-30-8
2	---	---	---	---	6790-27-8
3	---	---	---	---	19781-86-3
4	---	---	---	---	2809-78-1
5	---	---	---	---	62338-07-2
6	---	---	---	---	61886-62-2
7	---	---	---	---	55976-14-2
8	---	---	---	---	546-80-5
9	---	---	---	---	-

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 7:48  
 SAMPLE: S-NMS-5 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068111 # 701  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 54272.



Library Search                    Data: 30068111 # 715            Base m/z: 41  
 08/31/98 23:58:00 + 7:57        Cali: 30068111 # 3            RIC: 58944.  
 Sample: S-MM5-5 1/3SA/100M       INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 309 matched at least 7 of the 16 largest peaks in the unknown

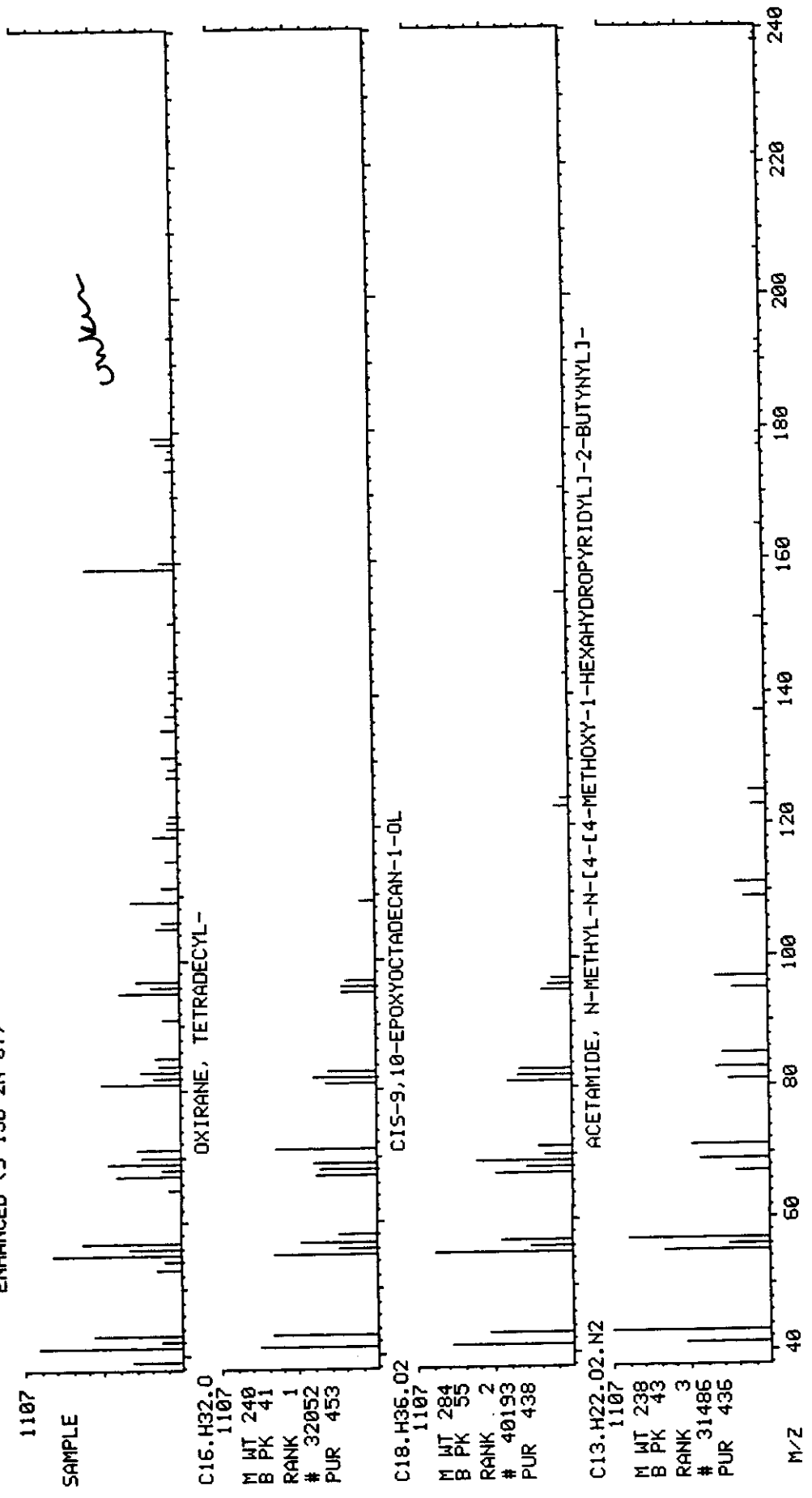
- Rank In.        Name  
 1 32052 OXIRANE, TETRADECYL-  
 2 40193 CIS-9,10-EPOXYOCTADECAN-1-OL  
 3 31486 ACETAMIDE, N-METHYL-N-[4-[4-METHOXY-1-HEXAHYDROPIRIDYL]-2-BUTYNYL]-  
 4 21927 TRANS-2-TRIDECENAL  
 5 50599 PENTALENE, OCTAHYDRO-1-(2-OCTYLDECYL)-  
 6 46251 DODECANE, 1,2-DIBROMO-  
 7 56499 9-OCTADECENOIC ACID (Z)-, 2-(ACETYLOXY)-1-[(ACETYLOXY)METHYL]ETHYL \*  
 8 37035 9-OCTADECENAL, (Z)-  
 9 18445 2-DODECENAL

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H32.O	240	41	453	904	468
2	C18.H36.O2	284	55	438	896	466
3	C13.H22.O2.N2	238	43	436	817	497
4	C13.H24.O	196	41	433	863	455
5	C26.H50	362	43	428	833	487
6	C12.H24.BR2	326	41	414	790	490
7	C25.H44.O6	440	43	414	709	515
8	C18.H34.O	266	55	407	848	467
9	C12.H22.O	182	70	403	874	425

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	7320-37-8
2	---	---	---	---	13980-12-6
3	---	---	---	---	- -
4	---	---	---	---	- -
5	---	---	---	---	55401-65-5
6	---	---	---	---	55334-42-4
7	---	---	---	---	55401-63-3
8	---	---	---	---	2423-10-1
9	---	---	---	---	4826-62-4

DATA: 30068111 # 715  
CALI: 30068111 # 3  
BASE M/Z: 41  
RIC: 58944.

MID LIBRARY SEARCH <LIBRARYNB>  
08/31/98 23:58:00 + 7:57  
SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA >)/1/35A NA M  
ENHANCED <S 158 2N 0T>



Library Search Data: 30068111 # 739 Base m/z: 57  
 08/31/98 23:58:00 + 8:13 Cali: 30068111 # 3 RIC: 131072.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 748 matched at least 7 of the 16 largest peaks in the unknown

Rank In. Name  
 1 25991 DODECANE, 2,6,10-TRIMETHYL-  
 2 37462 HEPTADECANE, 2,6-DIMETHYL-  
 3 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 4 22535 DODECANE, 4,6-DIMETHYL-  
 5 26001 DODECANE, 2,7,10-TRIMETHYL-  
 6 25997 PENTADECANE  
 7 19026 DECANE, 2,6,8-TRIMETHYL-  
 8 25994 DODECANE, 2,6,11-TRIMETHYL-  
 9 18998 UNDECANE, 3,7-DIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	Rfit
1	C15.H32	212	57	817	956	834
2	C19.H40	268	57	816	931	852
3	C21.H44	296	57	815	927	858
4	C14.H30	198	57	803	940	815
5	C15.H32	212	57	794	960	815
6	C15.H32	212	57	789	918	825
7	C13.H28	184	57	787	919	795
8	C15.H32	212	57	782	944	808
9	C13.H28	184	43	781	919	790

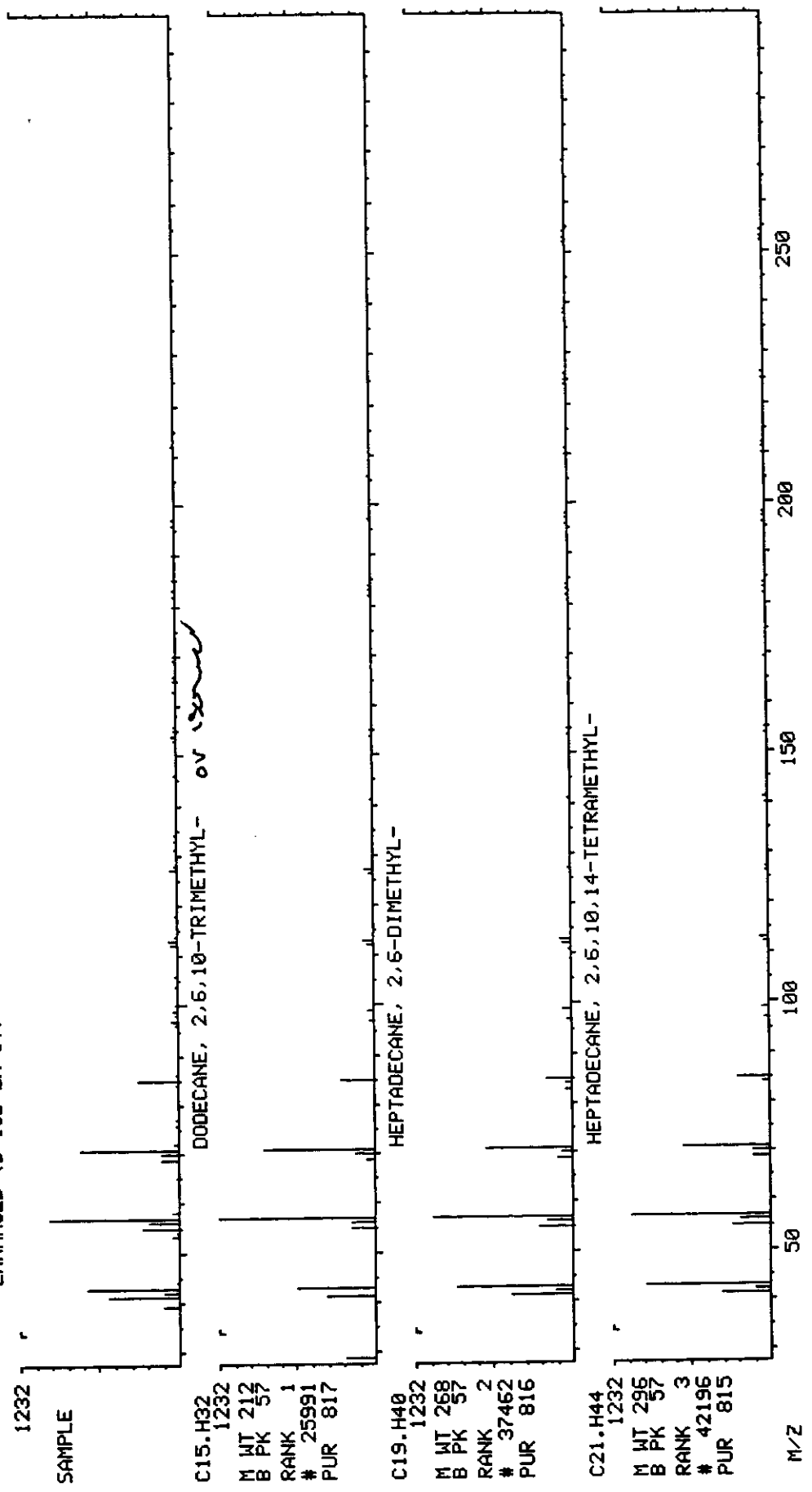
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	3891-98-3
2	---	---	---	---	54105-67-8
3	---	---	---	---	18344-37-1
4	---	---	---	---	61141-72-8
5	---	---	---	---	74645-98-0
6	---	---	---	---	629-62-9
7	---	---	---	---	62108-26-3
8	---	---	---	---	31295-56-4
9	---	---	---	---	17301-29-0



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 8:13  
 SAMPLE: 5-MMS-5 1/3SA/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA) /1/3SA NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 739  
 CALI: 30068111 # 3

BASE M/Z: 57  
 RIC: 131072.



Library Search Data: 30068111 # 764 Base m/z: 43  
 08/31/98 23:58:00 + 8:30 Cali: 30068111 # 3 RIC: 187392.  
 Sample: S-MMS-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 634 matched at least 7 of the 16 largest peaks in the unknown

Rank In.	Name
1	22530 TETRADECANE
2	19006 DECANE, 6-ETHYL-2-METHYL-
3	11612 NONANE, 2,5-DIMETHYL-
4	11607 UNDECANE
5	19007 DODECANE, 3-METHYL-
6	15358 UNDECANE, 3-METHYL-
7	22532 TRIDECANE, 2-METHYL-
8	42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-
9	39681 1-iodoundecane

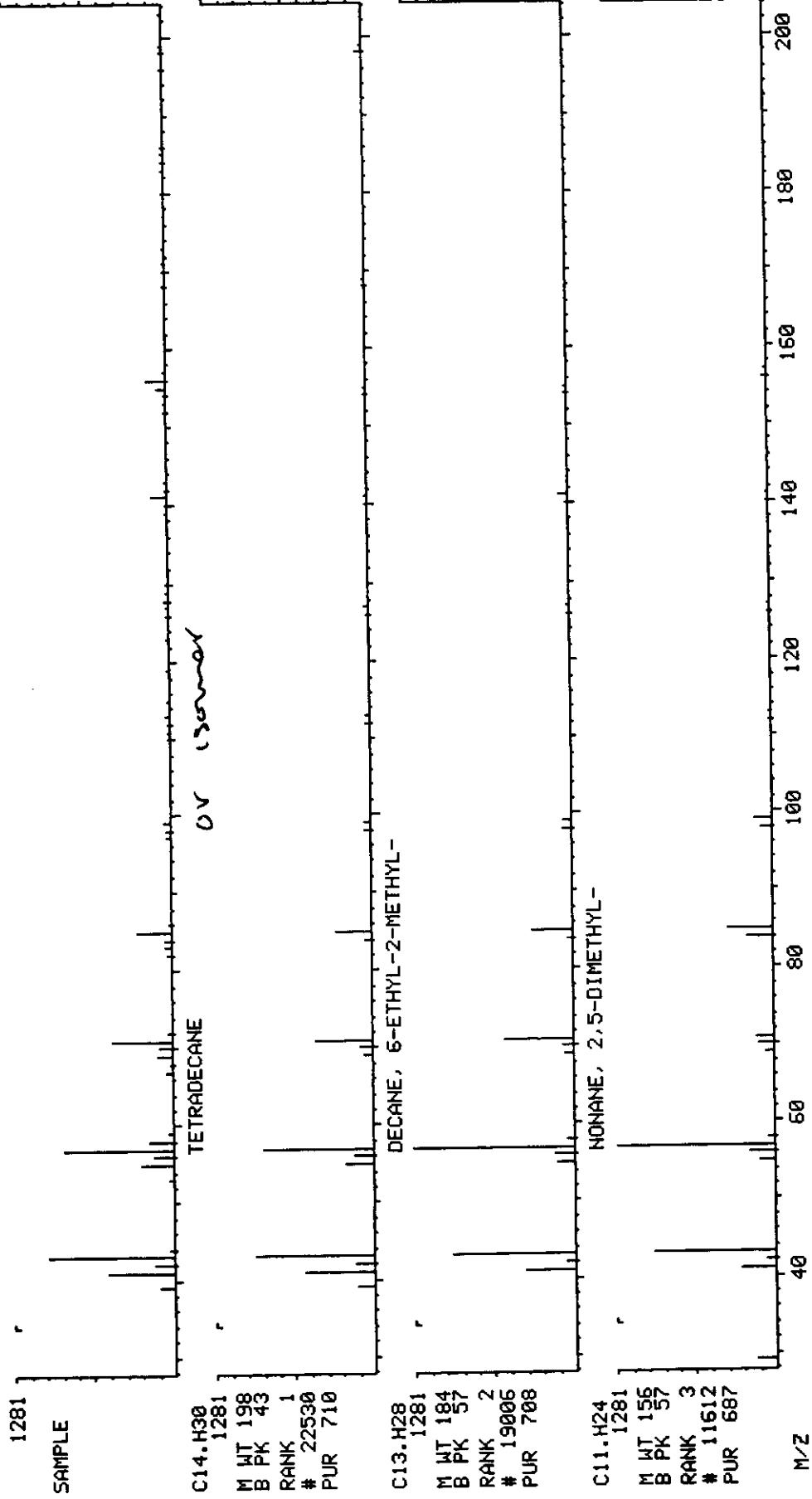
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C14.H30	198	43	710	950	715
2	C13.H28	184	57	708	954	708
3	C11.H24	156	57	687	865	687
4	C11.H24	156	43	684	970	684
5	C13.H28	184	57	684	896	686
6	C12.H26	170	57	682	911	682
7	C14.H30	198	57	676	878	679
8	C21.H44	296	57	676	921	696
9	C11.H23.I	282	57	675	864	682

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-59-4
2	---	---	---	---	62108-21-8
3	---	---	---	---	17302-27-1
4	---	---	---	---	1120-21-4
5	---	---	---	---	17312-57-1
6	---	---	---	---	1002-43-3
7	---	---	---	---	1560-96-9
8	---	---	---	---	18344-37-1
9	---	---	---	---	4282-44-4

DATA: 30058111 # 764  
CALI: 30058111 # 3

BASE M/Z: 43  
RIC: 187392.

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 8:30  
SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100% \*100% \*100% \*(NA/NA) >1/35A NA M  
ENHANCED (S 15B 2N 0T)



Library Search                      Data: 30068111 # 778                      Base m/z: 41  
 08/31/98 23:58:00 + 8:39            Cali: 30068111 # 3                      RIC: 59520.  
 Sample: S-MM5-5 1/3SA/100M            INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 618 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	31174 1-HEPTADECYNE
2	40193 CIS-9,10-EPOXYOCTADECAN-1-OL
3	56499 9-OCTADECENOIC ACID (Z)-, 2-(ACETYLOXY)-1-[(ACETYLOXY)METHYL]ETHYL *
4	37444 OXIRANE, HEXADECYL-
5	34009 1-OCTADECYNE
6	31486 ACETAMIDE, N-METHYL-N-[4-[4-METHOXY-1-HEXAHYDROPIRIDYL]-2-BUTYNYL]-
7	22481 CYCLODODECANEMETHANOL
8	28275 1-HEXADECYNE
9	25010 1-PENTADECYNE

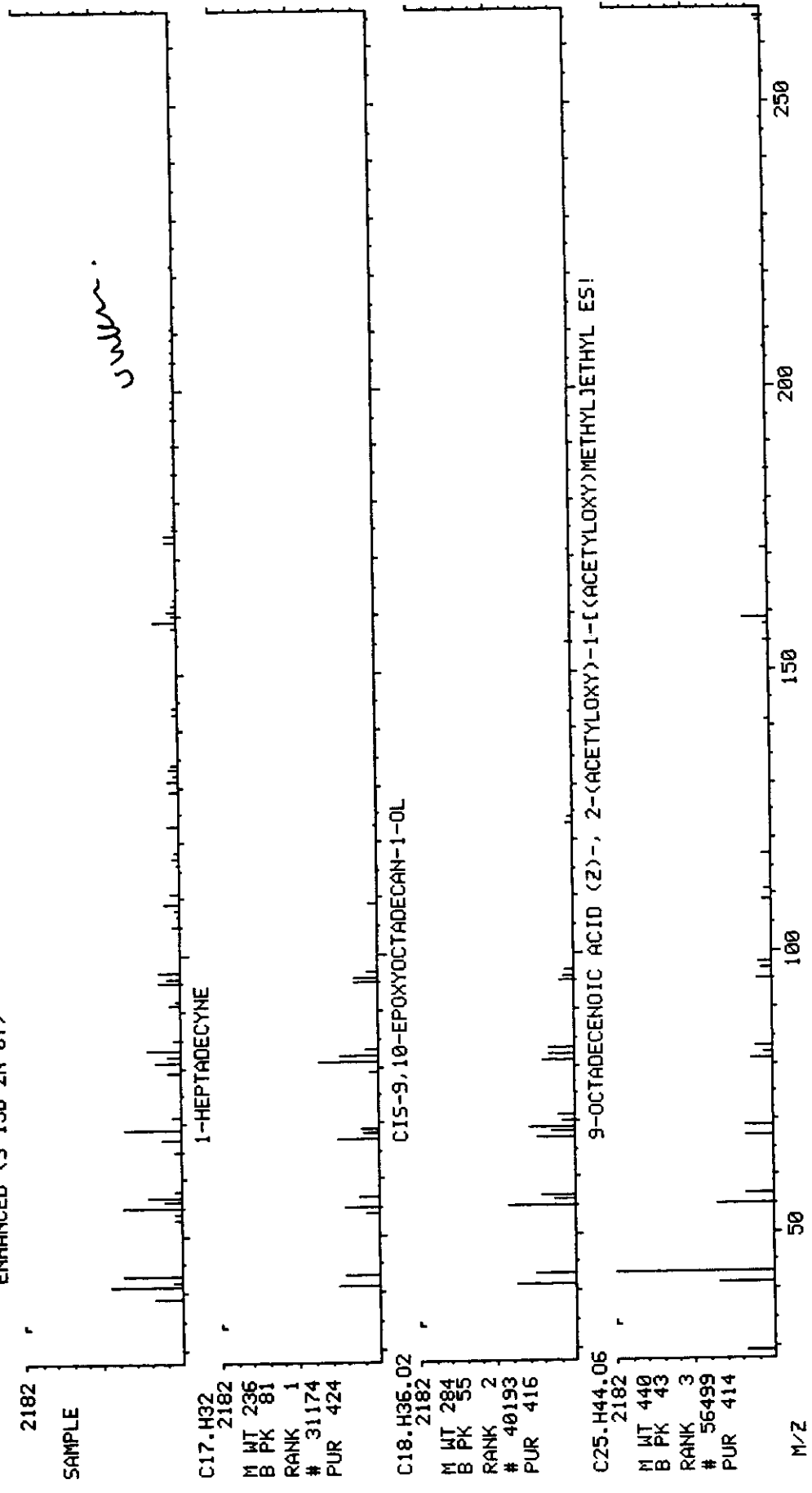
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C17.H32	236	81	424	837	439
2	C18.H36.O2	284	55	416	839	476
3	C25.H44.O6	440	43	414	744	520
4	C18.H36.O	268	57	412	783	479
5	C18.H34	250	81	410	837	435
6	C13.H22.O2.N2	238	43	405	798	472
7	C13.H26.O	198	55	404	808	463
8	C16.H30	222	81	404	830	420
9	C15.H28	208	81	402	825	418

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	26186-00-5
2	---	---	---	---	13980-12-6
3	---	---	---	---	55401-63-3
4	---	---	---	---	7390-81-0
5	---	---	---	---	629-89-0
6	---	---	---	---	- -
7	---	---	---	---	1892-12-2
8	---	---	---	---	629-74-3
9	---	---	---	---	765-13-9

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 8:39  
 SAMPLE: S-MMS-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100% \*100% \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 778  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 59520.



H  
 W  
 O  
 O

Library Search Data: 30068111 # 809 Base m/z: 41  
 08/31/98 23:58:00 + 9:00 Cali: 30068111 # 3 RIC: 56000.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 345 matched at least 7 of the 16 largest peaks in the unknown

- Rank In. Name
- 1 40193 CIS-9,10-EPOXYOCTADECAN-1-OL
- 2 25971 OXIRANE, DODECYL-
- 3 37444 OXIRANE, HEXADECYL-
- 4 37449 OCTADECANAL
- 5 22482 2-TRIDECEN-1-OL, (E)-
- 6 39230 9-EICOSYNE
- 7 31486 ACETAMIDE, N-METHYL-N-[4-[4-METHOXY-1-HEXAHYDROPYRIDYL]-2-BUTYNYL]-
- 8 11590 6-OCTEN-1-OL, 3,7-DIMETHYL-
- 9 44549 ETHANOL, 2-(9-OCTADECENYLOXY)-, (Z)-

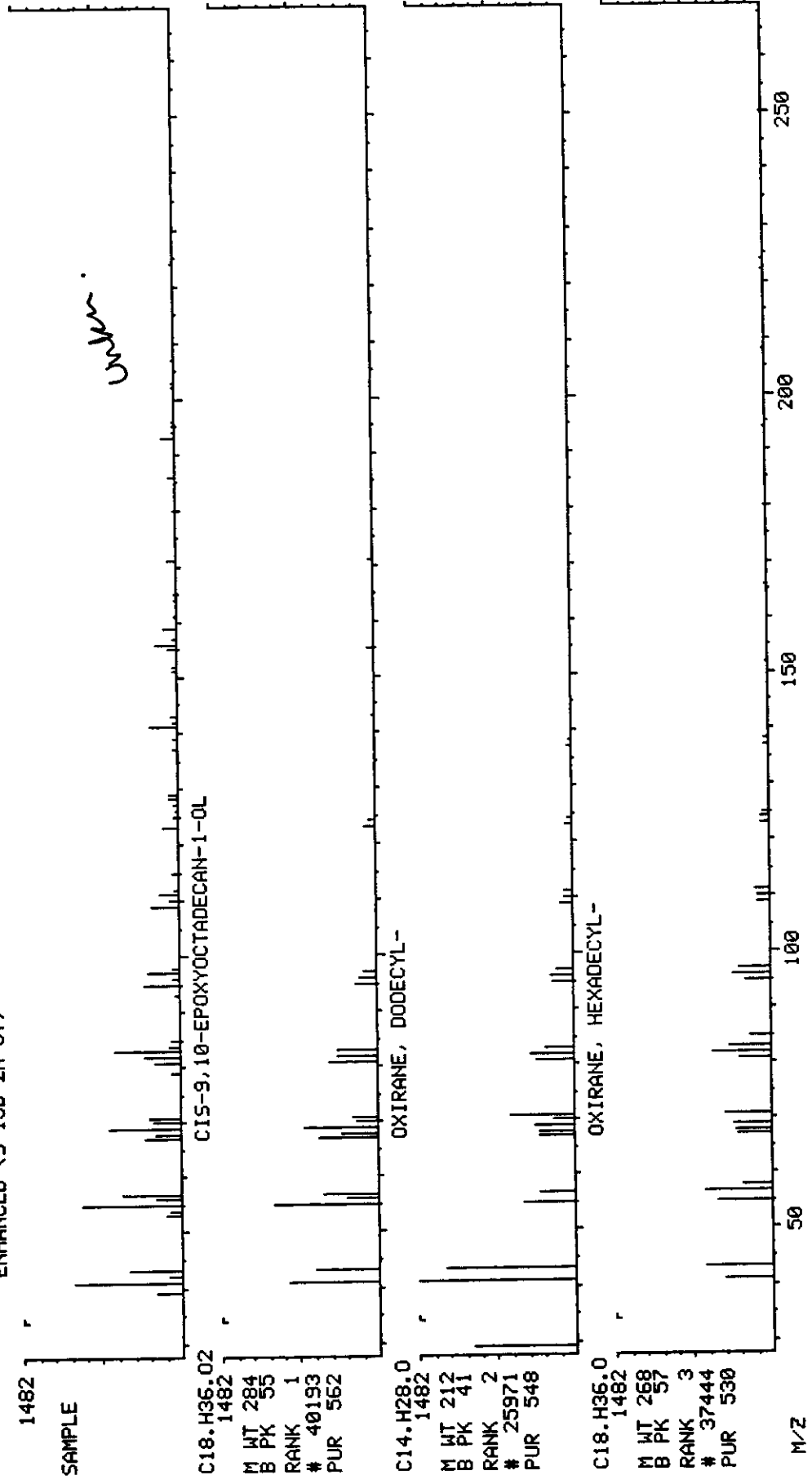
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C18.H36.O2	284	55	562	949	571
2	C14.H28.O	212	41	548	898	560
3	C18.H36.O	268	57	530	861	573
4	C18.H36.O	268	43	526	850	565
5	C13.H26.O	198	57	524	891	537
6	C20.H38	278	81	513	856	533
7	C13.H22.O2.N2	238	43	509	865	542
8	C10.H20.O	156	41	507	897	515
9	C20.H40.O2	312	55	503	806	562

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	13980-12-6
2	---	---	---	---	3234-28-4
3	---	---	---	---	7390-81-0
4	---	---	---	---	638-66-4
5	---	---	---	---	74962-98-4
6	---	---	---	---	71899-38-2
7	---	---	---	---	- -
8	---	---	---	---	106-22-9
9	---	---	---	---	5353-25-3

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 9:00  
 SAMPLE: S-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z \*(NA/NA >1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 809  
 CALI: 30068111 # 3

BASE M/Z: 41  
 RIC: 56000.



Library Search                    Data: 30068111 # 821            Base m/z: 43  
 08/31/98 23:58:00 + 9:08        Cali: 30068111 # 3            RIC: 133632.  
 Sample: S-MMS-5 1/3SA/100M        INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 855 matched at Least 7 of the 16 largest peaks in the unknown

- | Rank In. | Name                                      |
|----------|---|
| 1        | 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL- |
| 2        | 37462 HEPTADECANE, 2,6-DIMETHYL-          |
| 3        | 19016 UNDECANE, 4,7-DIMETHYL-             |
| 4        | 18987 UNDECANE, 2,8-DIMETHYL-             |
| 5        | 19023 DECANE, 2,6,7-TRIMETHYL-            |
| 6        | 8104 OCTANE, 3,5-DIMETHYL-                |
| 7        | 19054 UNDECANE, 2,6-DIMETHYL-             |
| 8        | 15353 2,6-DIMETHYLDECANE                  |
| 9        | 19004 UNDECANE, 4,6-DIMETHYL-             |

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C21.H44	296	57	804	967	822
2	C19.H40	268	57	790	949	811
3	C13.H28	184	43	756	940	762
4	C13.H28	184	43	756	955	760
5	C13.H28	184	57	755	951	755
6	C10.H22	142	57	754	966	754
7	C13.H28	184	57	754	909	763
8	C12.H26	170	43	754	916	774
9	C13.H28	184	57	750	960	750

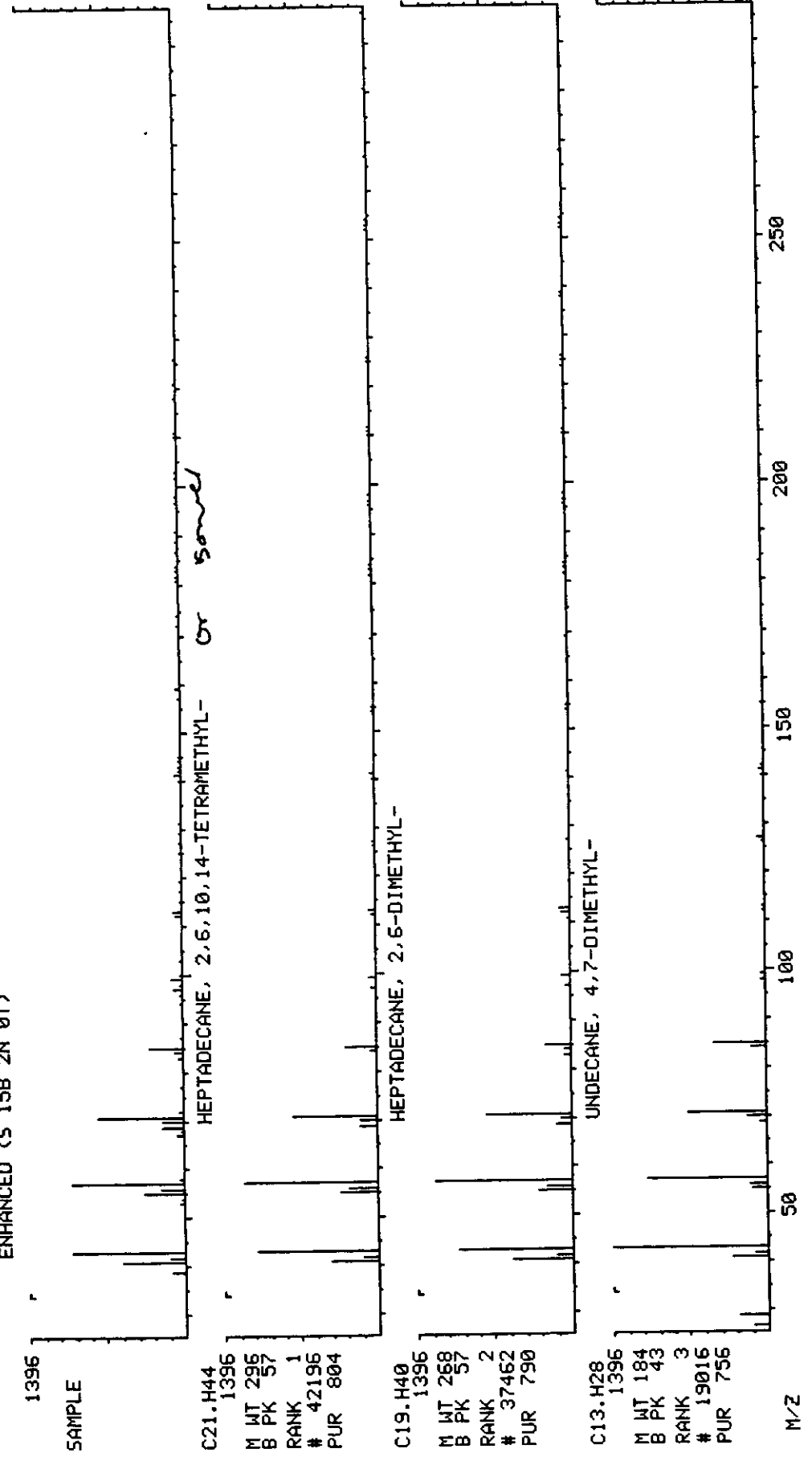
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	18344-37-1
2	—	—	—	—	54105-67-8
3	—	—	—	—	17301-32-5
4	—	—	—	—	17301-25-6
5	—	—	—	—	62108-25-2
6	—	—	—	—	15869-93-9
7	—	—	—	—	17301-23-4
8	—	—	—	—	13150-81-7
9	—	—	—	—	17312-82-2



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 9:08  
 SAMPLE: S-MMS-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/(NA/NA) >1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 821  
 CALI: 30068111 # 3

BASE M/Z: 43  
 RIC: 133632.



SAMPLE

C21.H44  
 1396  
 M WT 296  
 B PK 57  
 RANK 1  
 # 42196  
 PUR 804

C19.H40  
 1396  
 M WT 258  
 B PK 57  
 RANK 2  
 # 37462  
 PUR 790

C13.H28  
 1396  
 M WT 184  
 B PK 43  
 RANK 3  
 # 19016  
 PUR 756

M/Z

1396

Library Search                      Data: 30068111 # 859                      Base m/z: 57  
 08/31/98 23:58:00 + 9:33                      Cali: 30068111 # 3                      RIC: 151040.  
 Sample: S-MM5-5    1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 663 matched at least 7 of the 16 largest peaks in the unknown

Rank In.                      Name  
 1 19007 DODECANE, 3-METHYL-  
 2 18990 UNDECANE, 2,9-DIMETHYL-  
 3 15352 UNDECANE, 2-METHYL-  
 4 39681 1- IODOUNDECANE  
 5 22536 TRIDECANE, 3-METHYL-  
 6 46040 TETRADECANE, 1-IODO-  
 7 37252 DECANE, 1-IODO-  
 8 19006 DECANE, 6-ETHYL-2-METHYL-  
 9 19032 UNDECANE, 5-ETHYL-

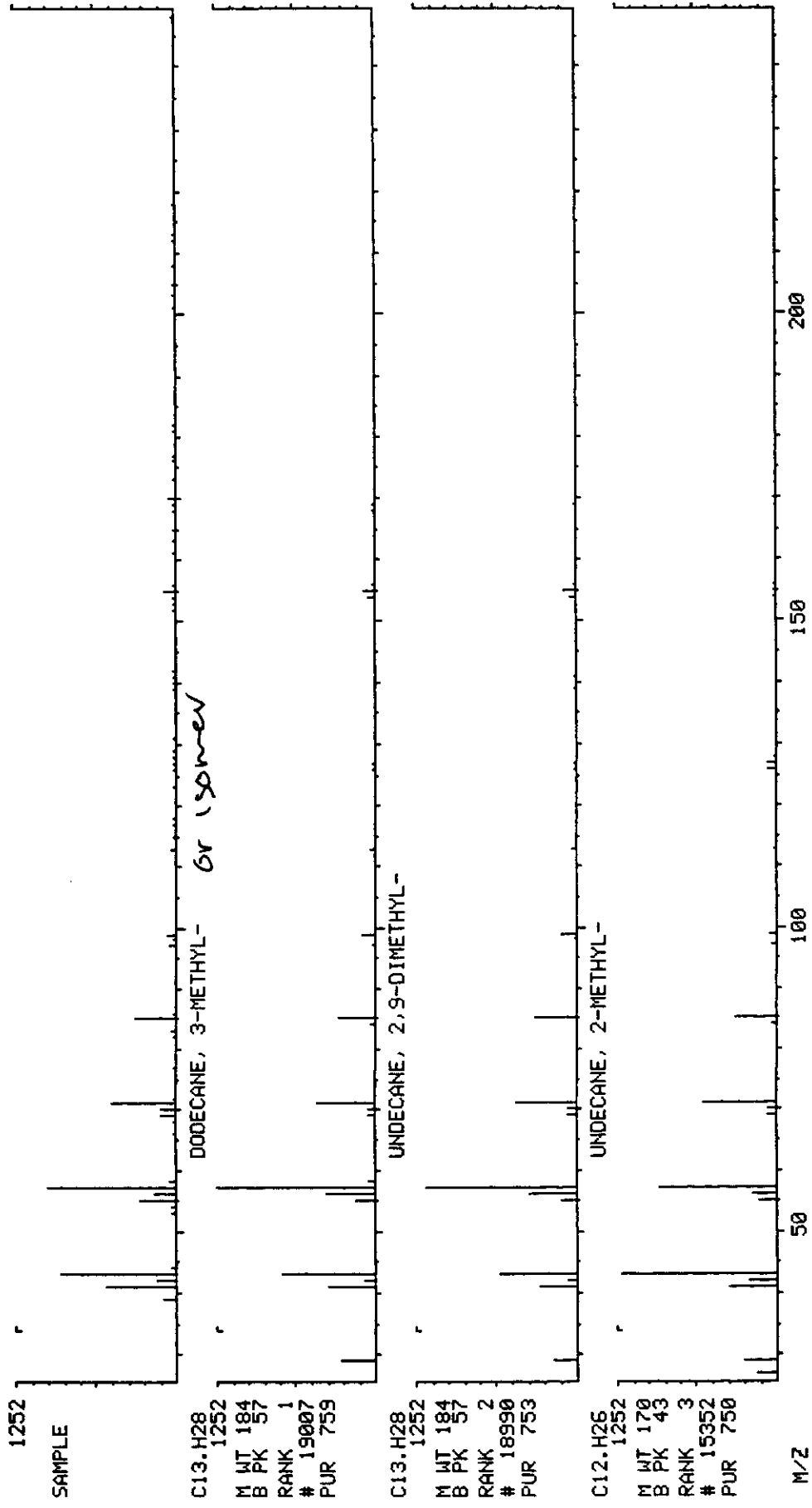
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C13.H28	184	57	759	953	766
2	C13.H28	184	57	753	956	753
3	C12.H26	170	43	750	925	760
4	C11.H23.I	282	57	747	966	755
5	C14.H30	198	57	743	940	748
6	C14.H29.I	324	57	742	983	746
7	C10.H21.I	268	57	741	971	747
8	C13.H28	184	57	737	949	745
9	C13.H28	184	57	736	963	736

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	17312-57-1
2	---	---	---	---	17301-26-7
3	---	---	---	---	7045-71-8
4	---	---	---	---	4282-44-4
5	---	---	---	---	6418-41-3
6	---	---	---	---	19218-94-1
7	---	---	---	---	2050-77-3
8	---	---	---	---	62108-21-8
9	---	---	---	---	17453-94-0

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 9:33  
 SAMPLE: 5-MM5-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100% \*100% \*1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30058111 # 859  
 CALI: 30058111 # 3

BASE M/Z: 57  
 RIC: 151040.



Library Search Data: 30068111 # 904 Base m/z: 41  
 08/31/98 23:58:00 + 10:03 Cali: 30068111 # 3 RIC: 68864.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 551 matched at least 7 of the 16 largest peaks in the unknown

Rank In. Name  
 1 37444 OXIRANE, HEXADECYL-  
 2 37449 OCTADECANAL  
 3 40233 NONADECANOL  
 4 42521 1-EICOSANOL  
 5 42180 OCTADECANE, 1-(ETHENYLOXY)-  
 6 40193 CIS-9,10-EPOXYOCTADECAN-1-OL  
 7 60775 HEXADECANE, 1,1-BIS(DODECYLOXY)-  
 8 48775 OCTADECANAL, 2-BROMO-  
 9 31486 ACETAMIDE, N-METHYL-N-[4-[4-METHOXY-1-HEXAHYDROPIRIDYL]-2-BUTYNYL]-

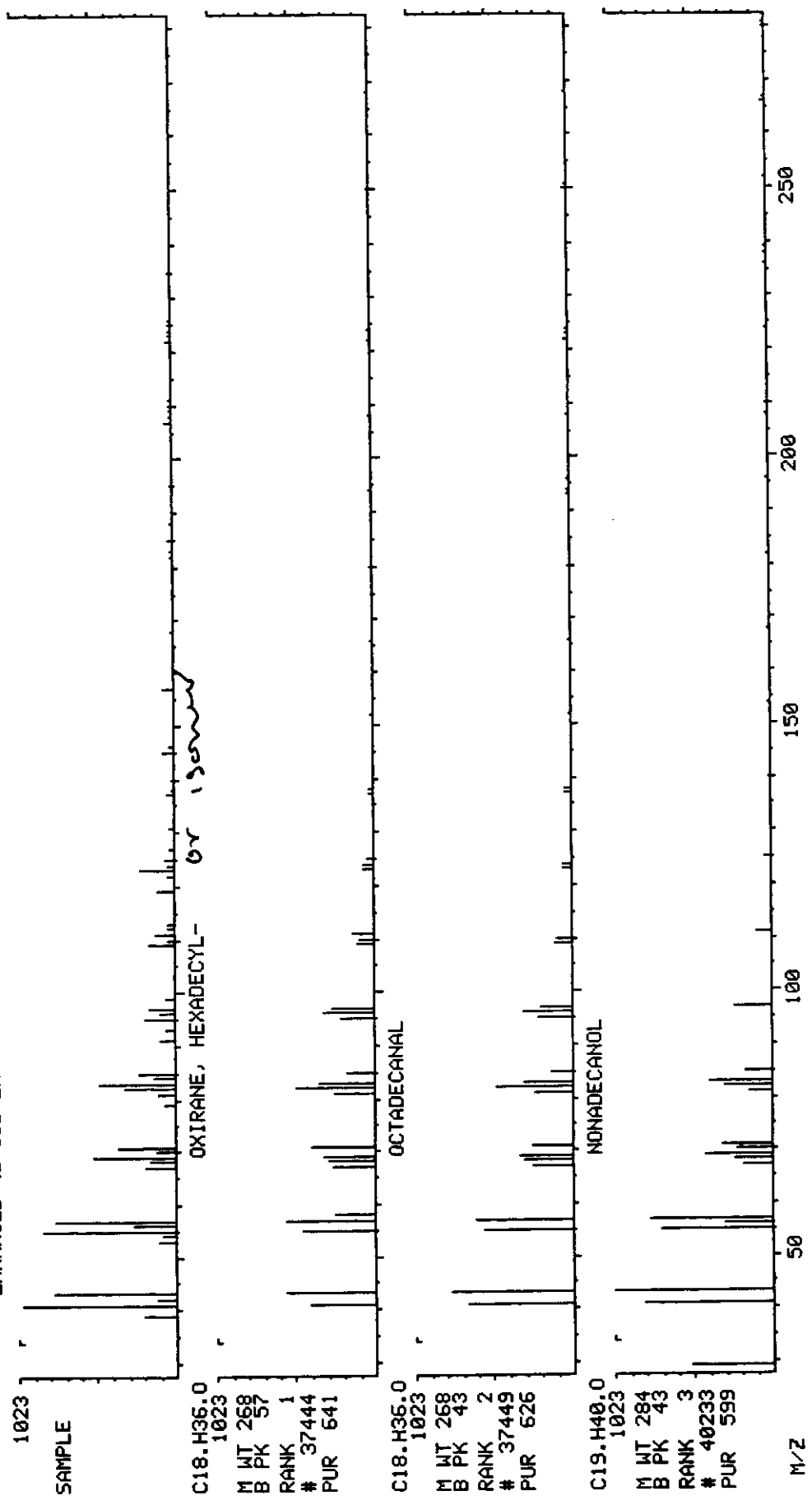
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C18.H36.O	268	57	641	880	675
2	C18.H36.O	268	43	626	892	657
3	C19.H40.O	284	43	599	945	623
4	C20.H42.O	298	43	595	941	623
5	C20.H40.O	296	43	587	942	610
6	C18.H36.O2	284	55	580	898	607
7	C40.H82.O2	594	43	575	900	612
8	C18.H35.O.BR	346	43	575	821	663
9	C13.H22.O2.N2	238	43	571	865	633

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	7390-81-0
2	---	---	---	---	638-66-4
3	---	---	---	---	52783-43-4
4	---	---	---	---	629-96-9
5	---	---	---	---	930-02-9
6	---	---	---	---	13980-12-6
7	---	---	---	---	56554-64-4
8	---	---	---	---	56599-95-2
9	---	---	---	---	

MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:58:00 + 10:03  
 SAMPLE: S-MM5-5 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100%/100% \*(NA/NA >)/1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068111 # 904  
 CALLI: 30068111 # 3

BASE M/Z: 41  
 RIC: 68864.



Library Search                    Data: 30068111 # 948            Base m/z: 57  
 08/31/98 23:58:00 + 10:33       Cali: 30068111 # 3            RIC: 154880.  
 Sample: S-MM5-5 1/3SA/100M       INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 338 matched at least 8 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 37465 NONADECANE  
 2 15969 HYDROXYLAMINE, O-DECYL-  
 3 37462 HEPTADECANE, 2,6-DIMETHYL-  
 4 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 5 25997 PENTADECANE  
 6 19016 UNDECANE, 4,7-DIMETHYL-  
 7 19523 1-DECANOL, 2-ETHYL-  
 8 46161 TRICOSANE  
 9 15353 2,6-DIMETHYLDECANE

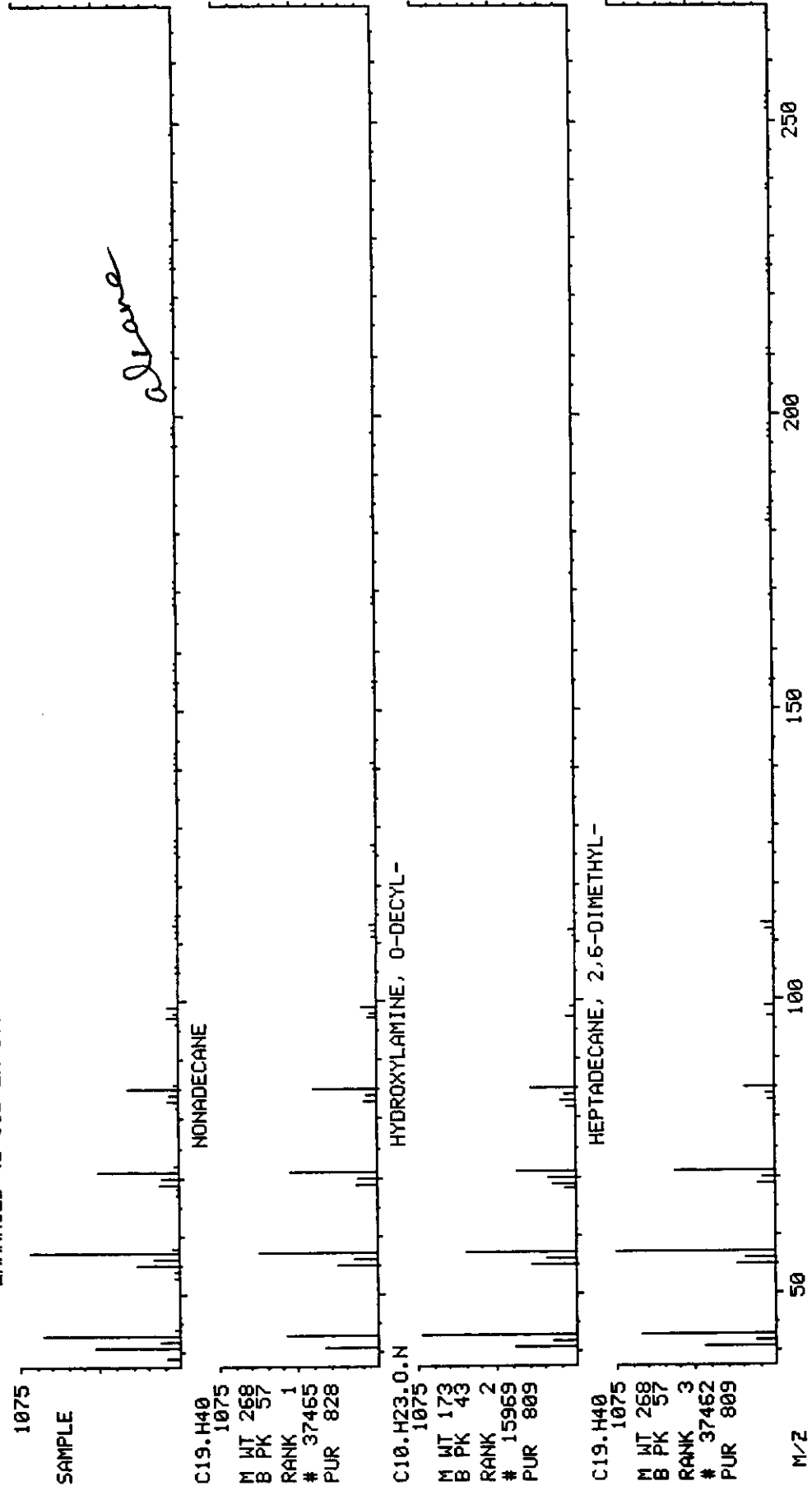
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	828	958	848
2	C10.H23.O.N	173	43	809	967	811
3	C19.H40	268	57	809	935	842
4	C21.H44	296	57	799	941	824
5	C15.H32	212	57	793	952	817
6	C13.H28	184	43	786	958	787
7	C12.H26.O	186	57	783	933	796
8	C23.H48	324	43	778	898	845
9	C12.H26	170	43	775	922	784

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-92-5
2	---	---	---	---	29812-79-1
3	---	---	---	---	54105-67-8
4	---	---	---	---	18344-37-1
5	---	---	---	---	629-62-9
6	---	---	---	---	17301-32-5
7	---	---	---	---	21078-65-9
8	---	---	---	---	638-67-5
9	---	---	---	---	13150-81-7

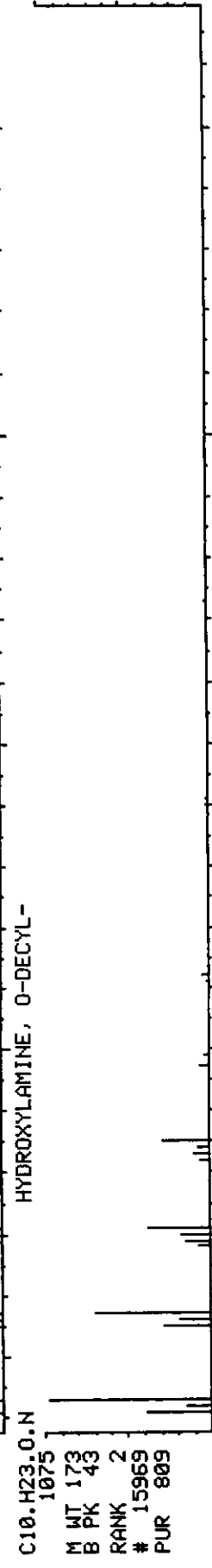
MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 23:58:00 + 10:33  
 SAMPLE: S-MMS-5 1/35A/100M INST. ID: F16  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 158 2N 0T)

DATA: 30068111 # 948  
 CALI: 30068111 # 3

BASE M/Z: 57  
 RIC: 154880.



*allane*



Library Search                      Data: 30068111 # 988                      Base m/z: 57  
 08/31/98 23:58:00 + 10:59                      Cali: 30068111 # 3                      RIC: 118144.  
 Sample: S-MM5-5 1/3SA/100M                      INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 446 matched at least 7 of the 16 largest peaks in the unknown

- Rank In.                      Name  
 1 25996 TRIDECANE, 4,8-DIMETHYL-  
 2 22530 TETRADECANE  
 3 19054 UNDECANE, 2,6-DIMETHYL-  
 4 25997 PENTADECANE  
 5 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 6 32059 HEPTADECANE  
 7 37462 HEPTADECANE, 2,6-DIMETHYL-  
 8 29263 HEXADECANE  
 9 22527 DODECANE, 2,5-DIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C15.H32	212	57	808	923	835
2	C14.H30	198	43	805	927	828
3	C13.H28	184	57	804	919	833
4	C15.H32	212	57	803	932	829
5	C21.H44	296	57	802	920	838
6	C17.H36	240	57	796	908	829
7	C19.H40	268	57	795	914	834
8	C16.H34	226	57	786	898	846
9	C14.H30	198	57	782	913	800

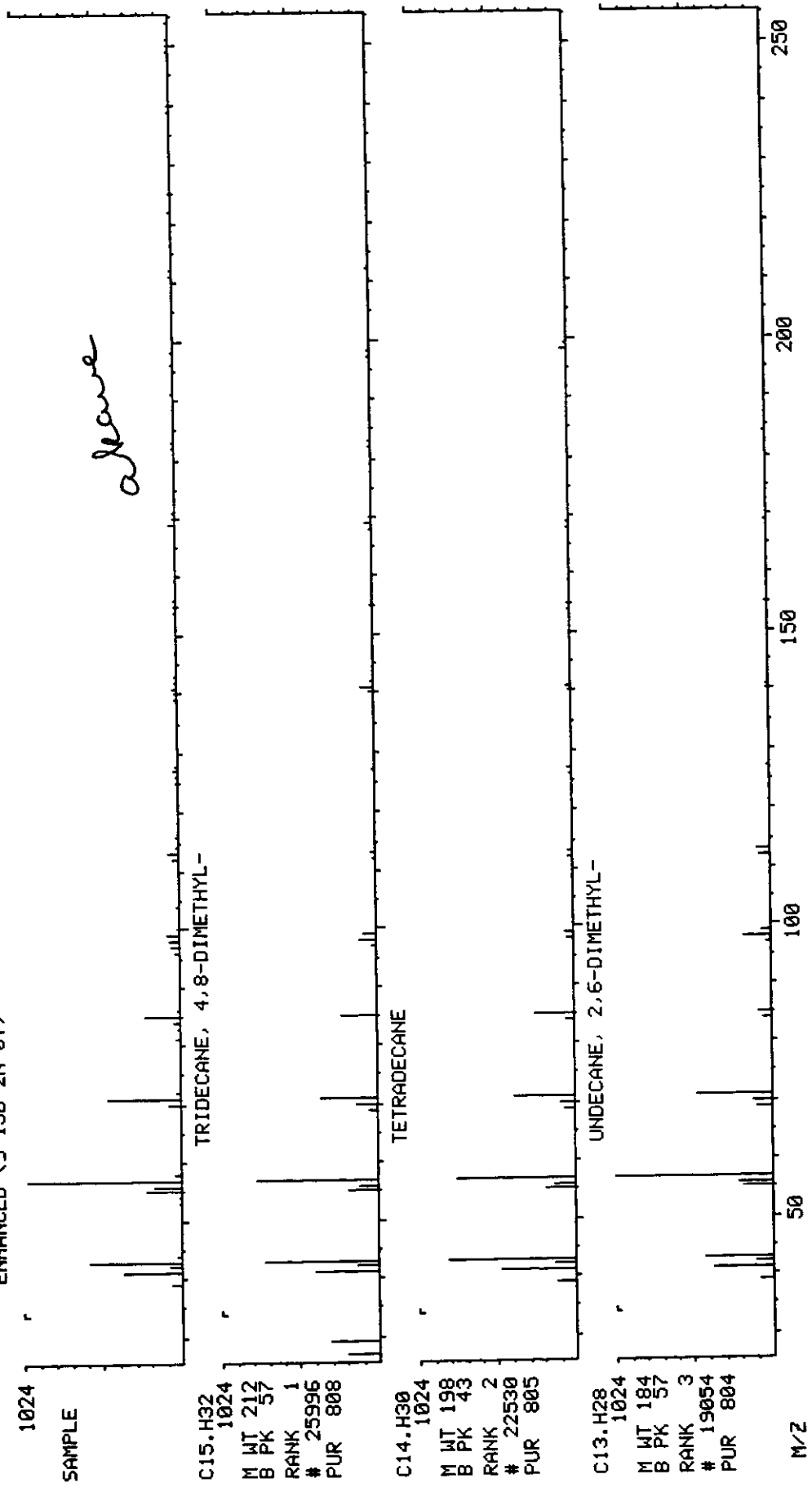
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	—	—	—	—	55030-62-1
2	—	—	—	—	629-59-4
3	—	—	—	—	17301-23-4
4	—	—	—	—	629-62-9
5	—	—	—	—	18344-37-1
6	—	—	—	—	629-78-7
7	—	—	—	—	54105-67-8
8	—	—	—	—	544-76-3
9	—	—	—	—	56292-65-0



MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 23:58:00 + 10:59  
 SAMPLE: S-RMS-5 1/35A/100M  
 CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA >)/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 30068111 # 988  
 CALI: 30068111 # 3

BASE M/Z: 57  
 RIC: 118144.



Library Search Data: 30068111 #1036 Base m/z: 57  
 08/31/98 23:58:00 + 11:31 Cali: 30068111 # 3 RIC: 180992.  
 Sample: S-MMS-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (\$ 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 840 matched at least 7 of the 16 largest peaks in the unknown

Rank In. Name  
 1 37462 HEPTADECANE, 2,6-DIMETHYL-  
 2 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 3 37465 NONADECANE  
 4 26001 DODECANE, 2,7,10-TRIMETHYL-  
 5 25991 DODECANE, 2,6,10-TRIMETHYL-  
 6 22535 DODECANE, 4,6-DIMETHYL-  
 7 37466 PENTADECANE, 2,6,10,14-TETRAMETHYL-  
 8 25994 DODECANE, 2,6,11-TRIMETHYL-  
 9 42192 HEPTADECANE, 2,6,10,15-TETRAMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	865	973	873
2	C21.H44	296	57	829	955	841
3	C19.H40	268	57	822	943	833
4	C15.H32	212	57	811	967	815
5	C15.H32	212	57	809	971	811
6	C14.H30	198	57	808	963	809
7	C19.H40	268	71	805	944	820
8	C15.H32	212	57	791	944	799
9	C21.H44	296	57	788	932	820

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	54105-67-8
2	---	---	---	---	18344-37-1
3	---	---	---	---	629-92-5
4	---	---	---	---	74645-98-0
5	---	---	---	---	3891-98-3
6	---	---	---	---	61141-72-8
7	---	---	---	---	1921-70-6
8	---	---	---	---	31295-56-4
9	---	---	---	---	54833-48-6

BASE M/Z: 57  
RIC: 180992.

DATA: 30058111 #1036  
CALI: 30058111 # 3

MID LIBRARY SEARCH <LIBRARYNB>

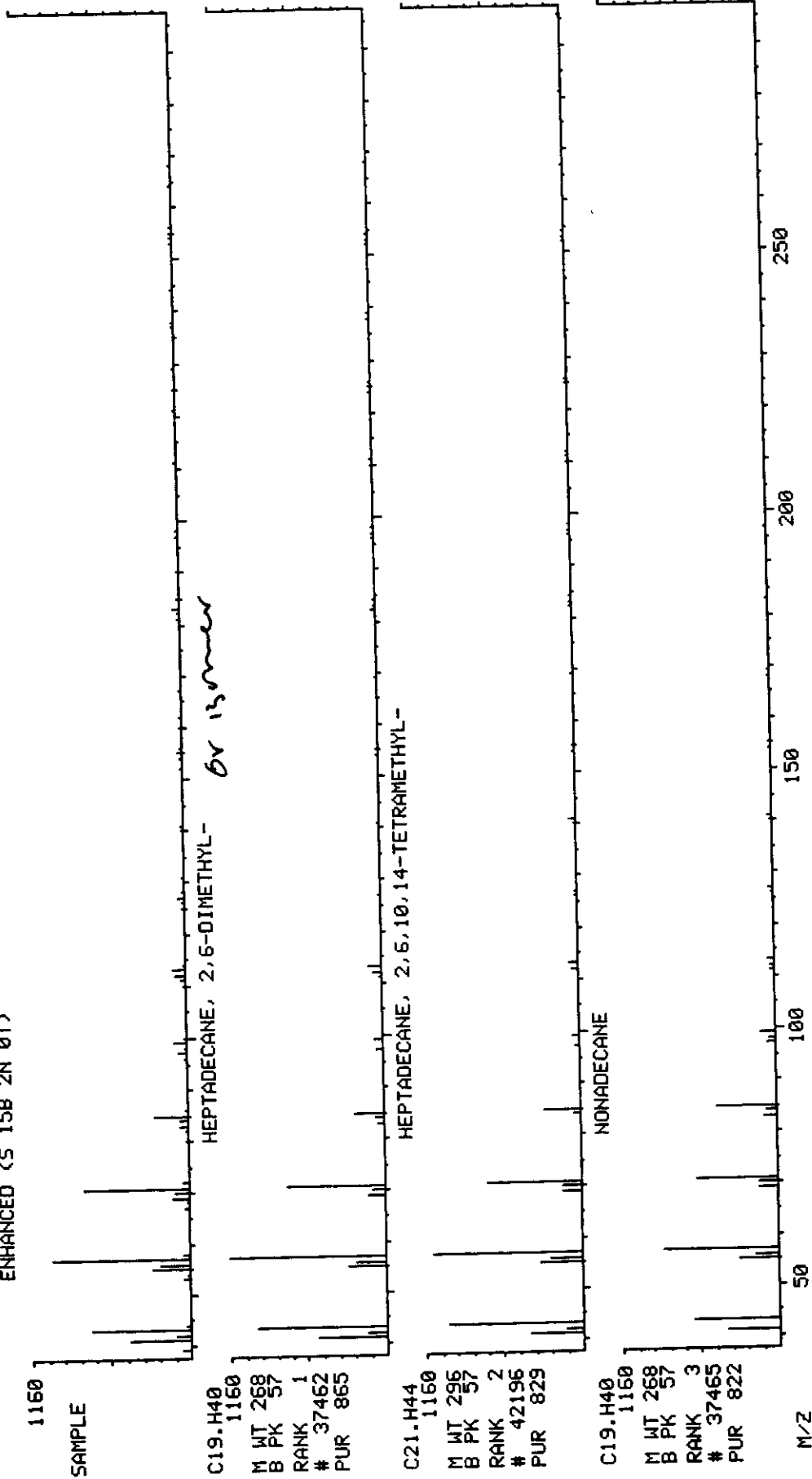
08/31/98 23:58:00 + 11:31

SAMPLE: 5-NMS-5 1/35A/100M

COND5.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M

ENHANCED <S 158 2N 0T>

INST. ID: F16



14  
CM  
CC

Library Search Data: 30068111 #1114 Base m/z: 57  
 08/31/98 23:58:00 + 12:23 Cali: 30068111 # 3 RIC: 76160.  
 Sample: S-MMS-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 317 matched at least 8 of the 16 largest peaks in the unknown

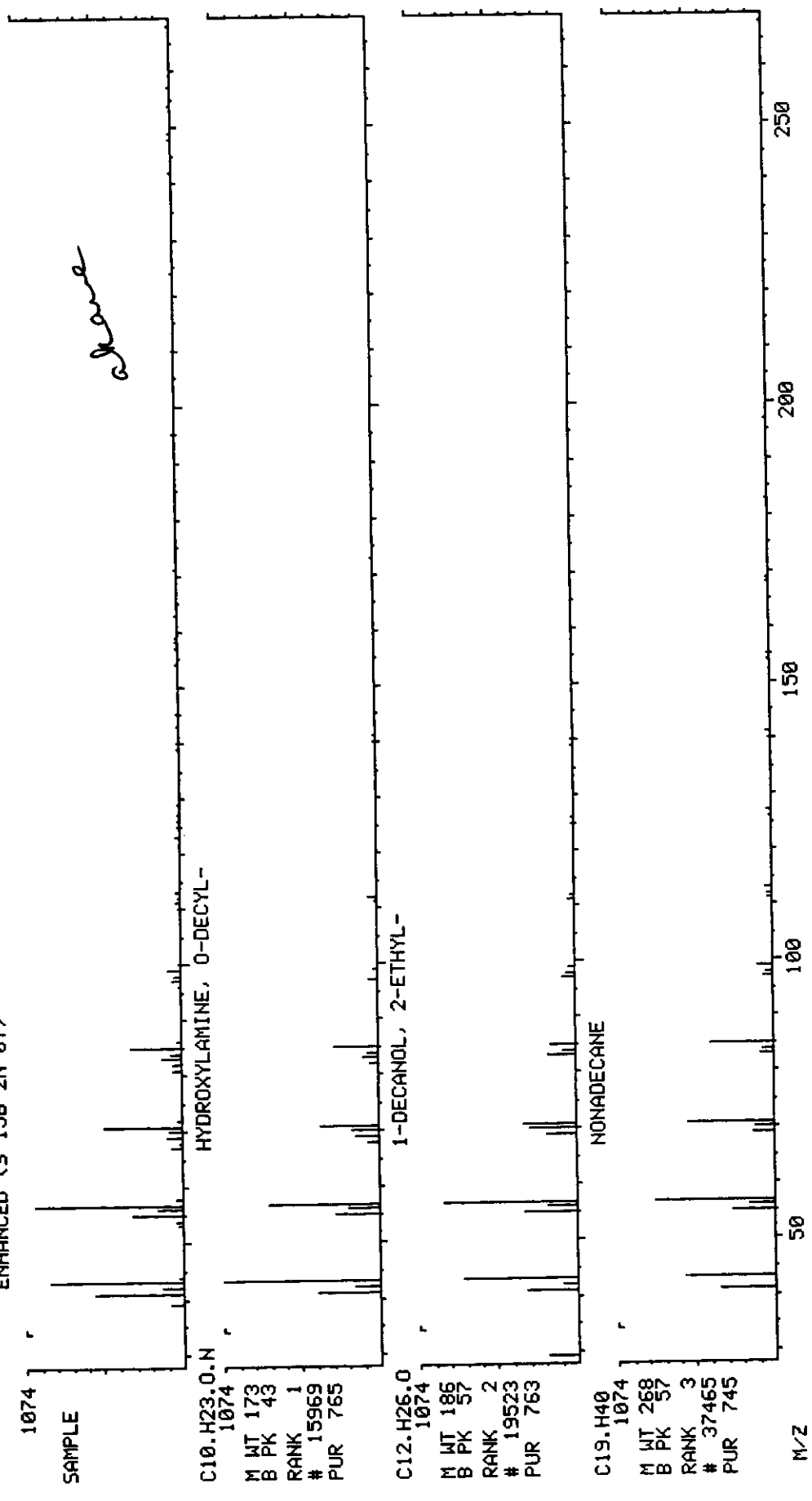
Rank In.	Name
1	15969 HYDROXYLAMINE, O-DECYL-
2	19523 1-DECANOL, 2-ETHYL-
3	37465 NONADECANE
4	12074 1-HEPTANOL, 2-PROPYL-
5	19015 DECANE, 2,5,6-TRIMETHYL-
6	19016 UNDECANE, 4,7-DIMETHYL-
7	42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-
8	8104 OCTANE, 3,5-DIMETHYL-
9	19520 1-OCTANOL, 2-BUTYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H23.O.N	173	43	765	960	773
2	C12.H26.O	186	57	763	947	763
3	C19.H40	268	57	745	928	780
4	C10.H22.O	158	43	733	953	733
5	C13.H28	184	57	713	929	713
6	C13.H28	184	43	711	937	724
7	C21.H44	296	57	711	911	760
8	C10.H22	142	57	710	942	717
9	C12.H26.O	186	57	710	890	723

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	29812-79-1
2	---	---	---	---	21078-65-9
3	---	---	---	---	629-92-5
4	---	---	---	---	10042-59-8
5	---	---	---	---	62108-23-0
6	---	---	---	---	17301-32-5
7	---	---	---	---	18344-37-1
8	---	---	---	---	15869-93-9
9	---	---	---	---	3913-02-8

DATA: 30068111 #1114  
CALI: 30068111 # 3  
BASE M/Z: 57  
RIC: 76160.

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 12:23  
SAMPLE: 5-MMS-5 1/35A/100M INST. ID: F16  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
ENHANCED (S 158 2N 0T)



Library Search Data: 30068111 #1119 Base m/z: 57  
 08/31/98 23:58:00 + 12:27 Cali: 30068111 # 3 RIC: 107648.  
 Sample: S-MM5-5 1/3SA/100M INST. ID: F16  
 Conds.: UG/ML \*100ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 840 matched at least 7 of the 16 largest peaks in the unknown

Rank In. Name  
 1 37465 NONADECANE  
 2 37462 HEPTADECANE, 2,6-DIMETHYL-  
 3 42196 HEPTADECANE, 2,6,10,14-TETRAMETHYL-  
 4 46161 TRICOSANE  
 5 32059 HEPTADECANE  
 6 37466 PENTADECANE, 2,6,10,14-TETRAMETHYL-  
 7 25991 DODECANE, 2,6,10-TRIMETHYL-  
 8 29263 HEXADECANE  
 9 26001 DODECANE, 2,7,10-TRIMETHYL-

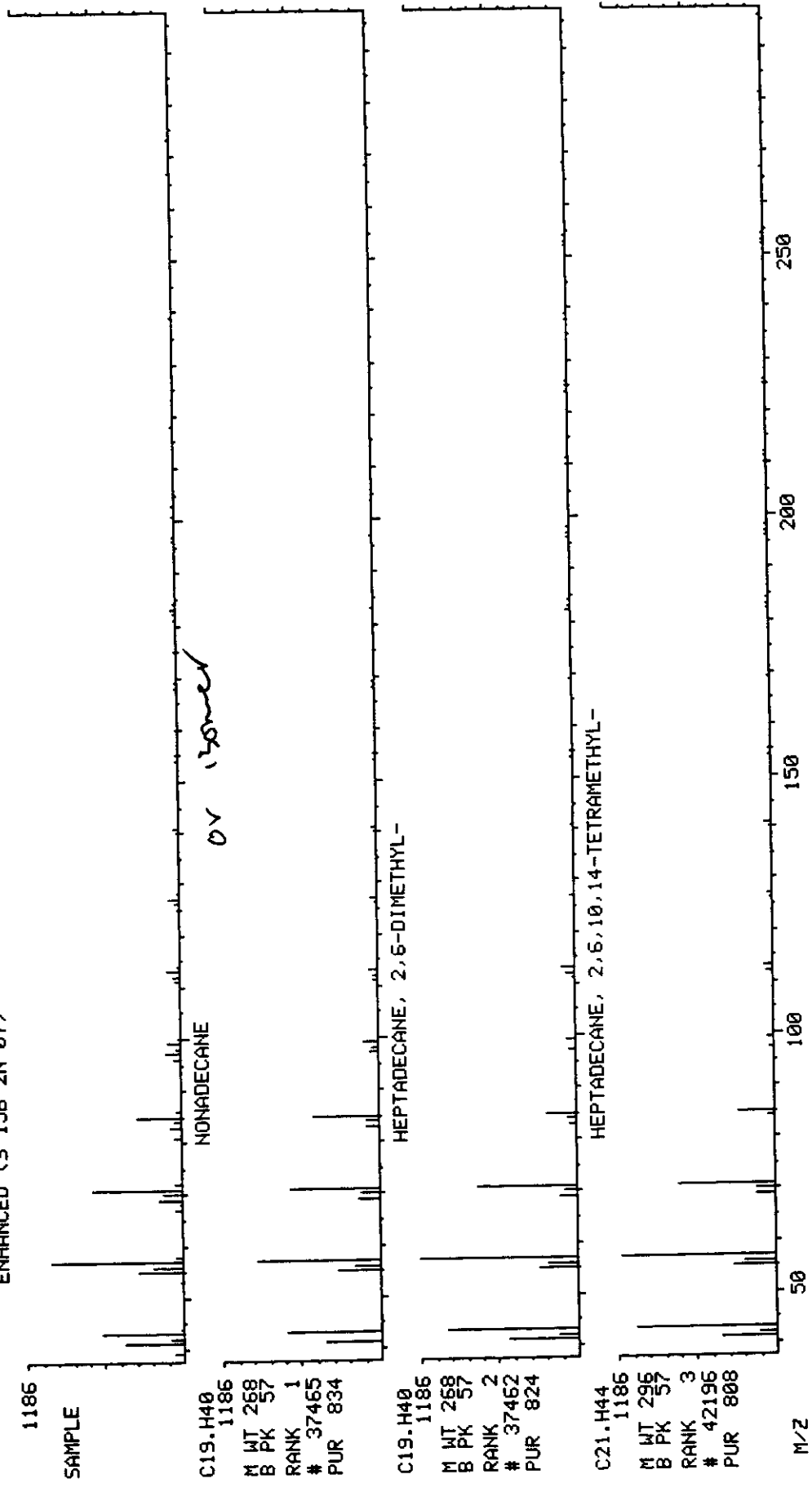
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C19.H40	268	57	834	958	848
2	C19.H40	268	57	824	941	834
3	C21.H44	296	57	808	935	824
4	C23.H48	324	43	800	907	854
5	C17.H36	240	57	794	925	819
6	C19.H40	268	71	793	931	804
7	C15.H32	212	57	792	961	793
8	C16.H34	226	57	789	920	822
9	C15.H32	212	57	787	962	787

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	629-92-5
2	---	---	---	---	54105-67-8
3	---	---	---	---	18344-37-1
4	---	---	---	---	638-67-5
5	---	---	---	---	629-78-7
6	---	---	---	---	1921-70-6
7	---	---	---	---	3891-98-3
8	---	---	---	---	544-76-3
9	---	---	---	---	74645-98-0

BASE M/Z: 57  
RIC: 107648.

DATA: 30068111 #1119  
CALI: 30068111 # 3

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 23:58:00 + 12:27  
SAMPLE: S-MMS-5 1/35A/100M INST. ID: F15  
CONDS.: UG/ML \*100ML \*100Z/100Z \*(NA/NA >/1/35A NA M  
ENHANCED (S 158 2N 0T)



TIC SELECTION REPORT

DATA FILE: 30068111

THE FOLLOWING PEAKS WERE REJECTED BECAUSE AT LEAST 40 % OF THEIR SIZE WAS ACCOUNTED FOR BY TARGET COMPOUNDS ELUTING WITHIN 4 SCANS OF THE PEAK TOP.

SCAN	SIZE	AMOUNT
340	249568.	29.386
531	398976.	46.978
830	339712.	40.000
1088	456704.	40.000
1553	344352.	40.000
1783	292571.	40.000

TOTAL NUMBER OF UNIDENTIFIED PEAKS WITH SIZE GREATER THAN 10 % OF THE CLOSEST INTERNAL STANDARD THAT DOES NOT HAVE INTERFERENCES = 97

INTERNAL STANDARDS THAT HAVE RIC SIZE LESS THAN 50 % OR GREATER THAN 200 % OF THE ESTIMATED RIC SIZE ARE CONSIDERED TO HAVE INTERFERENCES AND WILL NOT BE USED FOR QUANTITATION.

#	INTERNAL STANDARD	PERCENT OF ESTIMATED		SCAN
		RIC SIZE	RIC SIZE	
1	C140 NAPHTHALENE-D8	398976.	229 *	531
2	C150 ACENAPHTHENE-D10	339712.	152	830
3	C160 PHENANTHRENE-D10	456704.	181	1088
4	C170 CHRYSENE-D12	344352.	157	1553
5	C175 PERYLENE-D12	292571.	147	1783

\* INDICATES INTERFERENCE

SIZE = AREA



Initial Calibration

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

Library Entry Compound	Initial Calibration			Continuing Calibration	
	Response Factor Avg	% RSD Min	% RSD Max	Response Factor Min	% Diff Max
S1 15 C310 N-NITROSODIMETH	1.019		4.4		
S1 100 C320 ANILINE	1.846		6.2		
S1 105 C315 PHENOL	1.968		7.8	30.0	
S1 110 C325 BIS(2-CHLOROETH	1.368		6.1		
S1 115 C330 2-CHLOROPHENOL	1.497		5.5		
S1 125 C335 1,3-DICHLOROBEN	1.586		4.5		
S1 130 C340 1,4-DICHLOROBEN	1.618		4.8	30.0	
S1 145 C345 BENZYL ALCOHOL	0.854		9.3		
S1 150 C350 1,2-DICHLOROBEN	1.500		5.0		
S1 160 C355 2-METHYLPHENOL	1.174		7.4		
S1 165 C360 2,2'-OXYBIS(1-C	2.539		3.9		
S1 185 C365 4-METHYLPHENOL	1.276		9.0		
S1 190 C370 N-NITROSO-DI-N-	0.906	0.050	5.1		
S1 200 C375 HEXACHLOROETHAN	0.708		3.5		
S2 10 C410 NITROBENZENE	0.445		7.2		
S2 20 C415 ISOPHORONE	0.742		5.5		
S2 25 C420 2-NITROPHENOL	0.213		7.9	30.0	
S2 30 C425 2,4-DIMETHYLPHE	0.330		16.9		
S2 45 C430 BENZOIC ACID	0.276		4.7		
S2 35 C435 BIS(2-CHLOROETH	0.511		4.1		
S2 40 C440 2,4-DICHLOROPHE	0.308		6.6	30.0	
S2 55 C445 1,2,4-TRICHLORO	0.331		5.0		
S2 60 C450 NAPHTHALENE	1.022		6.6		
S2 80 C455 4-CHLOROANILINE	0.454		4.5		
S2 95 C460 HEXACHLOROBUTAD	0.168		5.1	30.0	
S2 130 C465 4-CHLORO-3-METH	0.338		9.0	30.0	
S2 145 C470 2-METHYLNAPHTHA	0.751		10.8		
S3 20 C510 HEXACHLOROCYCLO	0.372	0.050	10.1		
S3 25 C515 2,4,6-TRICHLORO	0.404		10.6	30.0	
S3 30 C520 2,4,5-TRICHLORO	0.412		7.1		
S3 40 C525 2-CHLORONAPHTHA	1.193		6.6		
S3 45 C530 2-NITROANILINE	0.502		2.9		
S3 55 C535 DIMETHYLPHTHALA	1.341		6.1		
S3 65 C540 ACENAPHTHYLENE	1.839		8.2		
S3 70 C543 2,6-DINITROTOLU	0.324		8.5		
S3 75 C545 3-NITROANILINE	0.383		1.4		
S3 80 C550 ACENAPHTHENE	1.158		6.9	30.0	
S3 85 C555 2,4-DINITROPHEN	0.189	0.050	10.3		

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

Library Entry Compound	Initial Calibration			Continuing Calibration	
	Response Factor Avg	% RSD Min	% RSD Max	Response Factor Min	% Diff Max
S3 90 C565 DIBENZOFURAN	1.669		6.2		
S3 105 C570 2,4-DINITROTOLU	0.397		8.8		
S3 130 C580 DIETHYLPHTHALAT	1.369		7.7		
S3 140 C585 4-CHLOROPHENYL-	0.613		5.4		
S3 135 C590 FLUORENE	1.298		6.8		
S3 150 C595 4-NITROANALINE	0.307		8.4		
S4 10 C610 4,6-DINITRO-2-M	0.132		7.6		
S4 15 C615 N-NITROSODIPHEN	0.551		4.2	30.0	
S4 20 C620 AZOBENZENE	1.171		3.9		
S4 30 C625 4-BROMOPHENYL-P	0.223		3.9		
S4 40 C630 HEXACHLOROBENZE	0.260		6.7		
S4 50 C635 PENTACHLOROPHEN	0.155		5.9	30.0	
S4 65 C640 PHENANTHRENE	1.033		5.2		
S4 70 C645 ANTHRACENE	0.987		11.0		
S4 80 C647 CARBAZOLE	0.764		10.3		
S4 85 C650 DI-N-BUTYLPHTHA	1.247		9.4		
S4 110 C655 FLUORANTHENE	0.900		13.5	30.0	
S5 10 C710 BENZIDINE	0.129		23.7		
S5 15 C715 PYRENE	1.323		6.5		
S5 40 C720 BUTYLBENZYLPHTH	0.685		7.7		
S5 55 C725 3,3'-DICHLORBE	0.408		5.7		
S5 50 C730 BENZO(A)ANTHRAC	1.125		5.4		
S5 60 C740 CHRYSENE	1.014		6.7		
S5 65 C745 BIS(2-ETHYLHEXY	0.980		9.4		
S6 10 C760 DI-N-OCTYL PHTH	1.925		7.4	30.0	
S6 15 C765 BENZO(B)FLUORAN	1.444		5.0		
S6 25 C770 BENZO(K)FLUORAN	1.159		9.8		
S6 35 C775 BENZO(A)PYRENE	1.182		5.3	30.0	
S6 55 C780 INDENO(1,2,3-CD	1.279		6.6		
S6 60 C785 DIBENZ(A,H)ANTH	1.267		8.1		
S6 65 C790 BENZO(G,H,I)PER	1.331		8.2		
S1 6 CS75 1,2-DICHLOROBENZ	0.920		6.5		
S2 2 CS20 NITROBENZENE-D5	0.470		5.2		
S3 3 CS25 2-FLUOROBIPHENY	1.234		6.2		
S5 2 CS30 TERPHENYL-D14	0.995		5.3		
S1 3 CS45 PHENOL-D5	2.026		2.0		
S1 2 CS50 2-FLUOROPHENOL	1.368		4.4		
S1 5 CS70 2-CHLOROPHENOL-	1.482		4.1		
S3 2 CS55 2,4,6-TRIBROMOP	0.236		6.9		

0 CCC and SPCC compounds are out

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

	ST16980827A		ST16980827B		ST16980827C		ST16980827		ST16980827D		ST16980827E	
Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S1 15	10	0.952	20	1.010	50	1.016	80	1.087	120	1.008	160	1.042
S1 100	10	1.947	20	1.988	50	1.814	80	1.876	120	1.778	160	1.674
S1 105	10	1.699	20	2.037	50	2.035	80	2.148	120	1.907	160	1.982
S1 110	10	1.336	20	1.465	50	1.438	80	1.414	120	1.291	160	1.261
S1 115	10	1.386	20	1.569	50	1.522	80	1.603	120	1.450	160	1.451
S1 125	10	1.515	20	1.668	50	1.618	80	1.659	120	1.546	160	1.510
S1 130	10	1.580	20	1.695	50	1.666	80	1.697	120	1.549	160	1.518
S1 145	10	0.702	20	0.851	50	0.872	80	0.913	120	0.866	160	0.922
S1 150	10	1.475	20	1.606	50	1.482	80	1.573	120	1.454	160	1.409
S1 160	10	0.998	20	1.206	50	1.221	80	1.209	120	1.199	160	1.213
S1 165	10	2.393	20	2.676	50	2.597	80	2.582	120	2.502	160	2.486
S1 185	10	1.049	20	1.298	50	1.355	80	1.287	120	1.316	160	1.351
S1 190	10	0.819	20	0.934	50	0.938	80	0.940	120	0.913	160	0.890
S1 200	10	0.665	20	0.740	50	0.713	80	0.720	120	0.706	160	0.706
S2 10	10	0.423	20	0.477	50	0.455	80	0.486	120	0.419	160	0.410
S2 20	10	0.685	20	0.778	50	0.774	80	0.778	120	0.726	160	0.708
S2 25	10	0.182	20	0.212	50	0.218	80	0.232	120	0.219	160	0.217
S2 30	10	0.234	20	0.309	50	0.334	80	0.400	120	0.349	160	0.353
S2 45			20	0.257	50	0.279	80	0.270	120	0.280	160	0.292
S2 35	10	0.481	20	0.529	50	0.517	80	0.538	120	0.502	160	0.498
S2 40	10	0.270	20	0.313	50	0.314	80	0.332	120	0.311	160	0.306
S2 55	10	0.319	20	0.350	50	0.334	80	0.350	120	0.322	160	0.310
S2 60	10	1.008	20	1.090	50	1.038	80	1.102	120	0.960	160	0.935
S2 80	10	0.419	20	0.466	50	0.454	80	0.480	120	0.449	160	0.455
S2 95	10	0.160	20	0.178	50	0.170	80	0.177	120	0.167	160	0.157

*[Handwritten signature]*

S2	130	10, 0.279	20, 0.337	50, 0.353	80, 0.365	120, 0.347	160, 0.346
S2	145	10, 0.773	20, 0.840	50, 0.827	80, 0.757	120, 0.671	160, 0.639
S3	20	10, 0.329	20, 0.387	50, 0.376	80, 0.428	120, 0.379	160, 0.330
S3	25	10, 0.327	20, 0.389	50, 0.405	80, 0.435	120, 0.417	160, 0.448
S3	30	10, 0.397	20, 0.437	50, 0.429	80, 0.443	120, 0.401	160, 0.366
S3	40	10, 1.172	20, 1.275	50, 1.208	80, 1.283	120, 1.128	160, 1.089
S3	45		20, 0.491	50, 0.510	80, 0.523	120, 0.487	160, 0.497
S3	55	10, 1.257	20, 1.423	50, 1.392	80, 1.426	120, 1.297	160, 1.251
S3	65	10, 1.751	20, 1.959	50, 1.896	80, 2.047	120, 1.707	160, 1.671
S3	70	10, 0.274	20, 0.339	50, 0.341	80, 0.351	120, 0.321	160, 0.317
S3	75		20, 0.382	50, 0.392	80, 0.382	120, 0.379	160, 0.379
S3	80	10, 1.140	20, 1.227	50, 1.182	80, 1.261	120, 1.075	160, 1.066
S3	85		20, 0.156	50, 0.190	80, 0.202	120, 0.203	160, 0.196
S3	95		20, 0.164	50, 0.178	80, 0.184	120, 0.168	160, 0.169

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

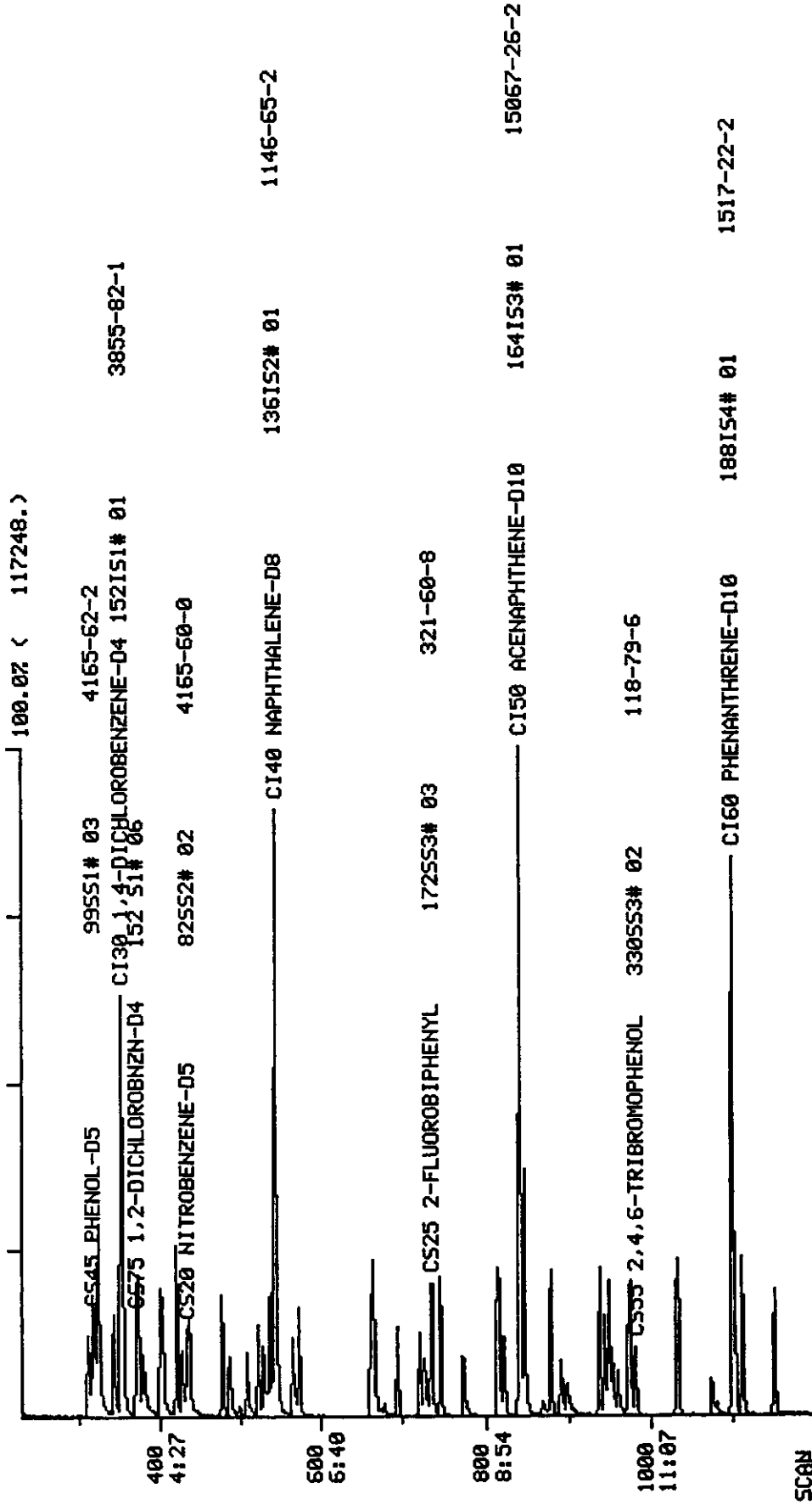
	ST16980827A		ST16980827B		ST16980827C		ST16980827		ST16980827D		ST16980827E	
Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S3 90	10, 1.617	20, 1.782	50, 1.736	80, 1.762	120, 1.556	160, 1.561						
S3 105	10, 0.339	20, 0.412	50, 0.420	80, 0.437	120, 0.396	160, 0.378						
S3 130	10, 1.300	20, 1.462	50, 1.452	80, 1.462	120, 1.328	160, 1.209						
S3 140	10, 0.597	20, 0.647	50, 0.631	80, 0.646	120, 0.579	160, 0.575						
S3 135	10, 1.251	20, 1.398	50, 1.340	80, 1.386	120, 1.224	160, 1.188						
S3 150		20, 0.326	50, 0.320	80, 0.306	120, 0.263	160, 0.321						
S4 10		20, 0.121	50, 0.139	80, 0.145	120, 0.124	160, 0.130						
S4 15	10, 0.568	20, 0.582	50, 0.562	80, 0.539	120, 0.530	160, 0.523						
S4 20	10, 1.150	20, 1.222	50, 1.186	80, 1.212	120, 1.098	160, 1.158						
S4 30	10, 0.224	20, 0.231	50, 0.223	80, 0.232	120, 0.215	160, 0.210						
S4 40	10, 0.273	20, 0.281	50, 0.255	80, 0.268	120, 0.238	160, 0.242						
S4 50		20, 0.148	50, 0.156	80, 0.170	120, 0.155	160, 0.147						
S4 65	10, 1.017	20, 1.081	50, 1.044	80, 1.096	120, 0.948	160, 1.010						
S4 70	10, 1.019	20, 1.100	50, 1.026	80, 1.059	120, 0.908	160, 0.809						
S4 80	10, 0.914	20, 0.763	50, 0.729	80, 0.769	120, 0.707	160, 0.701						
S4 85	10, 1.233	20, 1.366	50, 1.332	80, 1.333	120, 1.137	160, 1.080						
S4 110	10, 1.013	20, 1.008	50, 0.925	80, 0.950	120, 0.788	160, 0.715						
S5 10		20, 0.155	50, 0.143	80, 0.077	120, 0.141	160, 0.131						
S5 15	10, 1.209	20, 1.310	50, 1.295	80, 1.468	120, 1.295	160, 1.358						
S5 40	10, 0.587	20, 0.673	50, 0.692	80, 0.722	120, 0.700	160, 0.734						
S5 55		20, 0.441	50, 0.421	80, 0.383	120, 0.405	160, 0.391						
S5 50	10, 1.058	20, 1.195	50, 1.116	80, 1.203	120, 1.086	160, 1.092						
S5 60	10, 1.034	20, 1.068	50, 1.005	80, 1.102	120, 0.959	160, 0.919						

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S6	15	10, 1.322	20, 1.536	50, 1.448	80, 1.453	120, 1.490	160, 1.412
S6	25	10, 1.185	20, 1.223	50, 1.171	80, 1.264	120, 1.171	160, 0.940
S6	35	10, 1.168	20, 1.257	50, 1.206	80, 1.229	120, 1.147	160, 1.084
S6	55	10, 1.253	20, 1.355	50, 1.312	80, 1.368	120, 1.241	160, 1.144
S6	60	10, 1.254	20, 1.386	50, 1.288	80, 1.360	120, 1.212	160, 1.105
S6	65	10, 1.339	20, 1.436	50, 1.356	80, 1.426	120, 1.289	160, 1.139
S1	6	10, 0.832	20, 0.999	50, 0.946	80, 0.959	120, 0.902	160, 0.882
S2	2	10, 0.446	20, 0.494	50, 0.469	80, 0.504	120, 0.452	160, 0.452
S3	3	10, 1.252	20, 1.315	50, 1.253	80, 1.299	120, 1.171	160, 1.116
S5	2	10, 0.932	20, 1.002	50, 0.976	80, 1.082	120, 0.960	160, 1.019
S1	3		20, 2.057	50, 2.059	80, 2.046	120, 1.995	160, 1.971
S1	2		20, 1.291	50, 1.332	80, 1.448	120, 1.376	160, 1.394
S1	5		20, 1.561	50, 1.501	80, 1.503	120, 1.437	160, 1.407
S3	2		20, 0.261	50, 0.239	80, 0.233	120, 0.228	160, 0.217

1471  
 I-~~1471~~  
 [Signature]

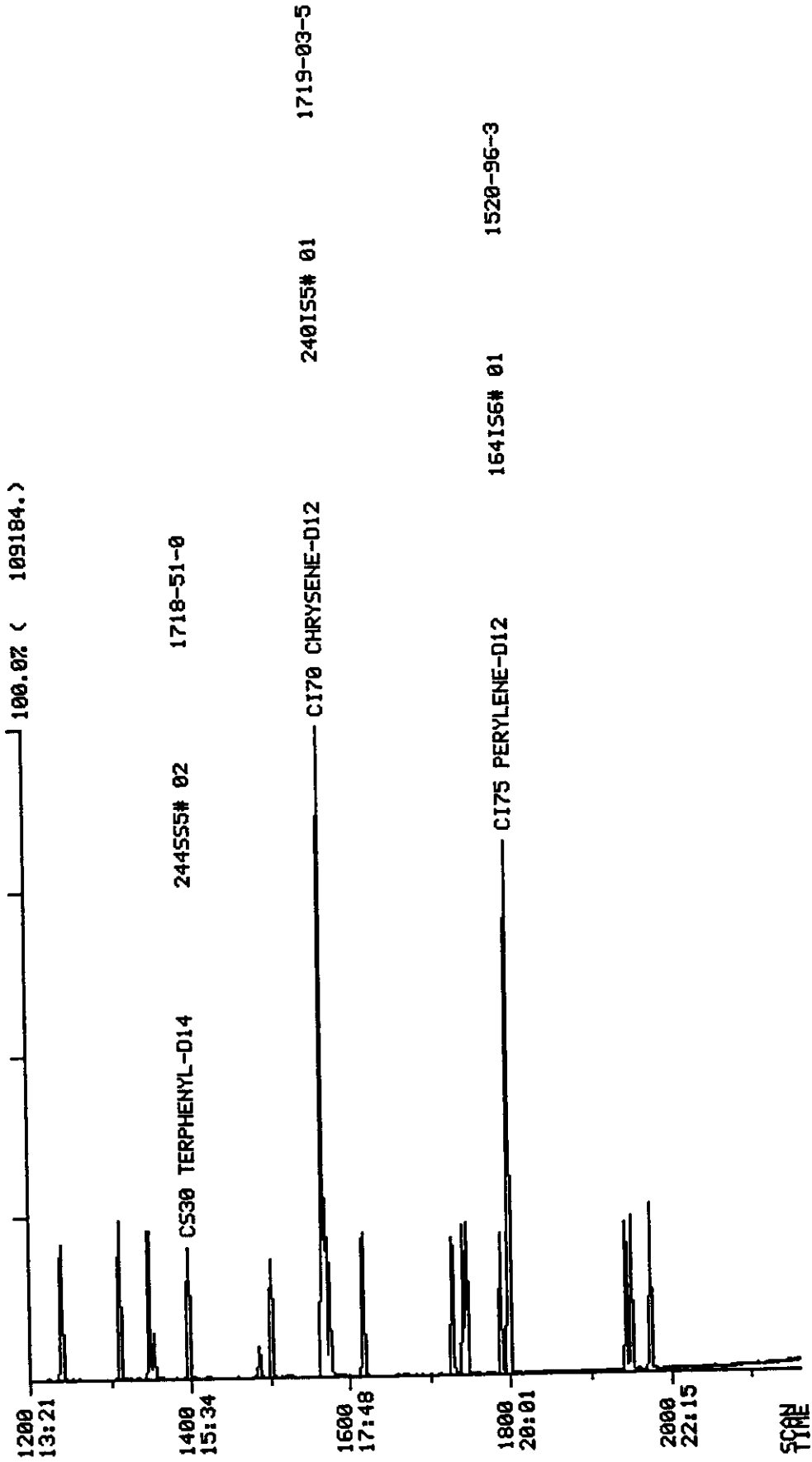
DATA FROM FILE: ST16980827A SCANS 230 TO 1200 ACQUIRED: 08/27/98 9:56:00  
 CALI: ST16980827A #3  
 SAMPLE: SST0010 10UG/ML CALI VI 070698D  
 CONDS.: INST. ID: F16



SCAN 1172



DATA FROM FILE: ST16980827A SCANS 1200 TO 2158 ACQUIRED: 08/27/98 9:56:00  
CALI: ST16980827A #3  
SAMPLE: SST0010 10UG/ML CALI VI 070698D  
CONDS.: INST. ID: F16



Data: ST16980827A.T1

08/27/98 9:56:00

Sample: SSTD010 10UG/ML CALI VI 070698D

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	C170 CHRYSENE-D12	240IS5# 01	1719-03-5
6	C175 PERYLENE-D12	164IS6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS75 1,2-DICHLOROBENZ-D4	152 S1# 06	
11	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
12	C320 ANILINE	93 S1#100	62-53-3
13	C315 PHENOL	94 S1#105CC	108-95-2
14	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
15	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
16	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
17	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
18	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
19	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
20	C355 2-METHYLPHENOL	108 S1#160	95-48-7
21	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
22	C365 4-METHYLPHENOL	108 S1#185	106-44-5
23	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
24	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
25	C410 NITROBENZENE	77 S2# 10	98-95-3
26	C415 ISOPHORONE	82 S2# 20	75-59-1
27	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
28	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
29	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
30	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
31	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
32	C450 NAPHTHALENE	128 S2# 60	91-20-3
33	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
34	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
35	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
36	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
37	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
38	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
39	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
40	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
41	C535 DIMETHYLPHTHALATE	163 S3# 55	131-11-3
42	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
43	C543 2,6-DINITROTOLUENE	165 S3# 70	606-20-2
44	C550 ACENAPHTHENE	153 S3# 80CC	83-32-9
45	C565 DIBENZOFURAN	168 S3# 90	132-64-9
46	C570 2,4-DINITROTOLUENE	165 S3#105	121-14-2
47	C580 DIETHYLPHTHALATE	149 S3#130	84-66-2

No	Name			
48	C590 FLUORENE	166	S3#135	86-73-7
49	C585 4-CHLOROPHENYL-PHENYLE	204	S3#140	7005-72-3
50	C615 N-NITROSDIPHENYLAMINE	169	S4# 15CC	87-30-6

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	350	3:54	1	1.000	A BB	33779.	40.000 UG/ML	4.55
2	136	541	6:01	2	1.000	A BB	121305.	40.000 UG/ML	4.55
3	164	840	9:21	3	1.000	A BB	64350.	40.000 UG/ML	4.55
4	188	1098	12:13	4	1.000	A BB	105698.	40.000 UG/ML	4.55
5	240	1565	17:24	5	1.000	A BV	92782.	40.000 UG/ML	4.55
6	264	1797	19:59	6	1.000	A BB	77868.	40.000 UG/ML	4.55
7	82	431	4:48	2	0.797	A BB	13522.	10.000 UG/ML	1.14
8	172	733	8:09	3	0.873	A BB	20134.	10.000 UG/ML	1.14
9	244	1395	15:31	5	0.891	A BB	21622.	10.000 UG/ML	1.14
10	152	369	4:06	1	1.054	A BB	7028.	10.000 UG/ML	1.14
11	74	109	1:13	1	0.311	A VB	8036.	10.000 UG/ML	1.14
12	93	310	3:27	1	0.886	A BV	16443.	10.000 UG/ML	1.14
13	94	316	3:31	1	0.903	A VB	14346.	10.000 UG/ML	1.14
14	93	323	3:36	1	0.923	A VB	11282.	10.000 UG/ML	1.14
15	128	323	3:36	1	0.923	A BB	11708.	10.000 UG/ML	1.14
16	146	341	3:48	1	0.974	A BV	12793.	10.000 UG/ML	1.14
17	146	352	3:55	1	1.006	A VB	13344.	10.000 UG/ML	1.14
18	108	377	4:12	1	1.077	A BB	5932.	10.000 UG/ML	1.14
19	146	371	4:08	1	1.060	A BB	12454.	10.000 UG/ML	1.14
20	108	399	4:26	1	1.140	A BB	8432.	10.000 UG/ML	1.14
21	45	399	4:26	1	1.140	A BB	20206.	10.000 UG/ML	1.14
22	108	425	4:44	1	1.214	A BB	8860.	10.000 UG/ML	1.14
23	70	418	4:39	1	1.194	A BV	6919.	10.000 UG/ML	1.14
24	117	418	4:39	1	1.194	A BB	5614.	10.000 UG/ML	1.14
25	77	434	4:50	2	0.802	A VB	12839.	10.000 UG/ML	1.14
26	82	475	5:17	2	0.878	A BV	20768.	10.000 UG/ML	1.14
27	139	484	5:23	2	0.895	A BV	5516.	10.000 UG/ML	1.14
28	107	506	5:38	2	0.935	A BB	7096.	10.000 UG/ML	1.14
29	93	520	5:47	2	0.961	A BV	14580.	10.000 UG/ML	1.14
30	162	526	5:51	2	0.972	A BB	8180.	10.000 UG/ML	1.14
31	180	535	5:57	2	0.989	A BB	9662.	10.000 UG/ML	1.14
32	128	544	6:03	2	1.006	A BB	30558.	10.000 UG/ML	1.14
33	127	563	6:16	2	1.041	A BB	12708.	10.000 UG/ML	1.14
34	225	571	6:21	2	1.055	A BB	4840.	10.000 UG/ML	1.14
35	107	659	7:20	2	1.218	A BB	8456.	10.000 UG/ML	1.14
36	142	662	7:22	2	1.224	qedt (2)	23441.	10.000 UG/ML	1.14
37	237	692	7:42	3	0.824	A BB	5288.	10.000 UG/ML	1.14
38	196	718	7:59	3	0.855	A BV 29/92	5258.	10.000 UG/ML	1.14
39	196	724	8:03	3	0.862	A VB	6382.	10.000 UG/ML	1.14
40	162	744	8:17	3	0.886	A BB	18852.	10.000 UG/ML	1.14
41	163	815	9:04	3	0.970	A BB	20216.	10.000 UG/ML	1.14
42	152	813	9:03	3	0.968	A BB	28168.	10.000 UG/ML	1.14
43	165	820	9:07	3	0.976	A BB	4408.	10.000 UG/ML	1.14
44	153	845	9:24	3	1.006	A BB	18346.	10.000 UG/ML	1.14
45	168	877	9:45	3	1.044	A BB	26008.	10.000 UG/ML	1.14
46	165	889	9:53	3	1.058	A BB	5460.	10.000 UG/ML	1.14
47	149	943	10:29	3	1.123	A BB	20908.	10.000 UG/ML	1.14
48	166	936	10:25	3	1.114	A BB	20124.	10.000 UG/ML	1.14
49	204	948	10:33	3	1.129	A BB	9610.	10.000 UG/ML	1.14
50	169	971	10:48	4	0.884	A BV	15016.	10.000 UG/ML	1.14

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	6:01	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:21	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:25	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:59	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:49	1.00	0.800	1.00	10.00	10.00	0.446	0.446	1.00
8	8:11	1.00	0.875	1.00	10.00	10.00	1.252	1.252	1.00
9	15:32	1.00	0.891	1.00	10.00	10.00	0.932	0.932	1.00
10	4:07	1.00	1.057	1.00	10.00	10.00	0.832	0.832	1.00
11	1:12	1.01	0.309	1.01	10.00	10.00	0.952	0.952	1.00
12	3:27	1.00	0.886	1.00	10.00	10.00	1.947	1.947	1.00
13	3:33	0.99	0.911	0.99	10.00	10.00	1.699	1.699	1.00
14	3:36	1.00	0.926	1.00	10.00	10.00	1.336	1.336	1.00
15	3:36	1.00	0.926	1.00	10.00	10.00	1.386	1.386	1.00
16	3:48	1.00	0.974	1.00	10.00	10.00	1.515	1.515	1.00
17	3:55	1.00	1.006	1.00	10.00	10.00	1.580	1.580	1.00
18	4:13	0.99	1.083	0.99	10.00	10.00	0.702	0.702	1.00
19	4:08	1.00	1.063	1.00	10.00	10.00	1.475	1.475	1.00
20	4:27	1.00	1.143	1.00	10.00	10.00	0.998	0.998	1.00
21	4:26	1.00	1.140	1.00	10.00	10.00	2.393	2.393	1.00
22	4:44	1.00	1.217	1.00	10.00	10.00	1.049	1.049	1.00
23	4:40	1.00	1.200	1.00	10.00	10.00	0.819	0.819	1.00
24	4:39	1.00	1.194	1.00	10.00	10.00	0.665	0.665	1.00
25	4:51	1.00	0.806	1.00	10.00	10.00	0.423	0.423	1.00
26	5:19	0.99	0.884	0.99	10.00	10.00	0.685	0.685	1.00
27	5:24	1.00	0.896	1.00	10.00	10.00	0.182	0.182	1.00
28	5:39	1.00	0.939	1.00	10.00	10.00	0.234	0.234	1.00
29	5:48	1.00	0.965	1.00	10.00	10.00	0.481	0.481	1.00
30	5:52	1.00	0.974	1.00	10.00	10.00	0.270	0.270	1.00
31	5:57	1.00	0.989	1.00	10.00	10.00	0.319	0.319	1.00
32	6:04	1.00	1.007	1.00	10.00	10.00	1.008	1.008	1.00
33	6:16	1.00	1.043	1.00	10.00	10.00	0.419	0.419	1.00
34	6:22	1.00	1.057	1.00	10.00	10.00	0.160	0.160	1.00
35	7:20	1.00	1.220	1.00	10.00	10.00	0.279	0.279	1.00
36	7:22	1.00	1.226	1.00	10.00	10.00	0.773	0.773	1.00
37	7:42	1.00	0.824	1.00	10.00	10.00	0.329	0.329	1.00
38	8:00	1.00	0.856	1.00	10.00	10.00	0.327	0.327	1.00
39	8:03	1.00	0.862	1.00	10.00	10.00	0.397	0.397	1.00
40	8:17	1.00	0.887	1.00	10.00	10.00	1.172	1.172	1.00
41	9:06	1.00	0.974	1.00	10.00	10.00	1.257	1.257	1.00
42	9:03	1.00	0.969	1.00	10.00	10.00	1.751	1.751	1.00
43	9:09	1.00	0.980	1.00	10.00	10.00	0.274	0.274	1.00
44	9:25	1.00	1.007	1.00	10.00	10.00	1.140	1.140	1.00
45	9:46	1.00	1.045	1.00	10.00	10.00	1.617	1.617	1.00
46	9:55	1.00	1.061	1.00	10.00	10.00	0.339	0.339	1.00
47	10:31	1.00	1.126	1.00	10.00	10.00	1.300	1.300	1.00
48	10:26	1.00	1.117	1.00	10.00	10.00	1.251	1.251	1.00
49	10:33	1.00	1.130	1.00	10.00	10.00	0.597	0.597	1.00
50	10:50	1.00	0.886	1.00	10.00	10.00	0.568	0.568	1.00

Quantitation Report File: ST16980827A

Data: ST16980827A.TI

08/27/98 9:56:00

Sample: SSTD010 10UG/ML CALI VI 070698D

Conds.: INST. ID: F16

Formula: 1UL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
51	C620 AZOBENZENE	77 S4# 20	103-33-3
52	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30	101-55-3
53	C630 HEXACHLOROBENZENE	284 S4# 40	118-74-1
54	C640 PHENANTHRENE	178 S4# 65	85-01-8
55	C645 ANTHRACENE	178 S4# 70	120-12-7
56	C647 CARBAZOLE	167 S4# 80	86-74-8
57	C650 DI-N-BUTYLPHTHALATE	149 S4# 85	84-74-2
58	C655 FLUORANTHENE	202 S4#110CC	206-44-0
59	C715 PYRENE	202 S5# 15	129-00-0
60	C720 BUTYLBENZYLPHTHALATE	149 S5# 40	85-68-7
61	C730 BENZO(A)ANTHRACENE	228 S5# 50	56-55-3
62	C740 CHRYSENE	228 S5# 60	218-01-9
63	C745 BIS(2-ETHYLNEXYL)PHTHA	149 S5# 65	117-81-7
64	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC	117-84-0
65	C765 BENZO(B)FLUORANTHENE	252 S6# 15	205-99-2
66	C770 BENZO(K)FLUORANTHENE	252 S6# 25	207-08-9
67	C775 BENZO(A)PYRENE	252 S6# 35	50-32-8
68	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55	193-39-5
69	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60	53-70-3
70	C790 BENZO(G,H,I)PERYLENE	276 S6# 65	191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	77	974	10:50	4	0.887	A BB	30384.	10.000 UG/ML	1.14
52	248	1032	11:29	4	0.940	A BB	5922.	10.000 UG/ML	1.14
53	284	1030	11:27	4	0.938	A BB	7204.	10.000 UG/ML	1.14
54	178	1102	12:15	4	1.004	A BV	26864.	10.000 UG/ML	1.14
55	178	1111	12:21	4	1.012	A VB	26934.	10.000 UG/ML	1.14
56	167	1149	12:47	4	1.046	A BB	24164.	10.000 UG/ML	1.14
57	149	1238	13:46	4	1.128	A BB	32570.	10.000 UG/ML	1.14
58	202	1310	14:34	4	1.193	A BB	26774.	10.000 UG/ML	1.14
59	202	1346	14:58	5	0.860	A BB	28050.	10.000 UG/ML	1.14
60	149	1499	16:40	5	0.958	A BB	13618.	10.000 UG/ML	1.14
61	228	1564	17:24	5	0.999	A BV	24540.	10.000 UG/ML	1.14
62	228	1569	17:27	5	1.003	A VB	23974.	10.000 UG/ML	1.14
63	149	1616	17:58	5	1.033	A BB	18824.	10.000 UG/ML	1.14
64	149	1726	19:12	6	0.960	A BB	31870.	10.000 UG/ML	1.14
65	252	1740	19:21	6	0.968	A BV	25726.	10.000 UG/ML	1.14
66	252	1744	19:24	6	0.971	A VB	23064.	10.000 UG/ML	1.14
67	252	1787	19:53	6	0.994	A BV	22740.	10.000 UG/ML	1.14
68	276	1942	21:36	6	1.081	A BV	24388.	10.000 UG/ML	1.14
69	278	1949	21:41	6	1.085	A VB	24406.	10.000 UG/ML	1.14
70	276	1973	21:57	6	1.098	A BB	26073.	10.000 UG/ML	1.14

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	10:51	1.00	0.888	1.00	10.00	10.00	1.150	1.150	1.00
52	11:29	1.00	0.940	1.00	10.00	10.00	0.224	0.224	1.00
53	11:29	1.00	0.939	1.00	10.00	10.00	0.273	0.273	1.00
54	12:17	1.00	1.005	1.00	10.00	10.00	1.017	1.017	1.00
55	12:22	1.00	1.012	1.00	10.00	10.00	1.019	1.019	1.00
56	12:47	1.00	1.046	1.00	10.00	10.00	0.914	0.914	1.00
57	13:47	1.00	1.127	1.00	10.00	10.00	1.233	1.233	1.00
58	14:36	1.00	1.194	1.00	10.00	10.00	1.013	1.013	1.00
59	15:00	1.00	0.861	1.00	10.00	10.00	1.209	1.209	1.00
60	16:41	1.00	0.958	1.00	10.00	10.00	0.587	0.587	1.00
61	17:24	1.00	0.999	1.00	10.00	10.00	1.058	1.058	1.00
62	17:28	1.00	1.003	1.00	10.00	10.00	1.034	1.034	1.00
63	17:58	1.00	1.032	1.00	10.00	10.00	0.812	0.812	1.00
64	19:13	1.00	0.961	1.00	10.00	10.00	1.637	1.637	1.00
65	19:23	1.00	0.969	1.00	10.00	10.00	1.322	1.322	1.00
66	19:26	1.00	0.972	1.00	10.00	10.00	1.185	1.185	1.00
67	19:54	1.00	0.996	1.00	10.00	10.00	1.168	1.168	1.00
68	21:38	1.00	1.082	1.00	10.00	10.00	1.253	1.253	1.00
69	21:43	1.00	1.086	1.00	10.00	10.00	1.254	1.254	1.00
70	21:59	1.00	1.100	1.00	10.00	10.00	1.339	1.339	1.00

**MANUAL EDIT CODES**

1. PEAK NOT FOUND
2. POOR CHROMATOGRAPHY
3. WRONG ISOMER

**ALL MANUAL EDITS MUST BE  
INITIALED, DATED, AND SIGNED**

SCANS 640 TO 679

DATA: ST16980827A #1  
CALI: ST16980827A #3

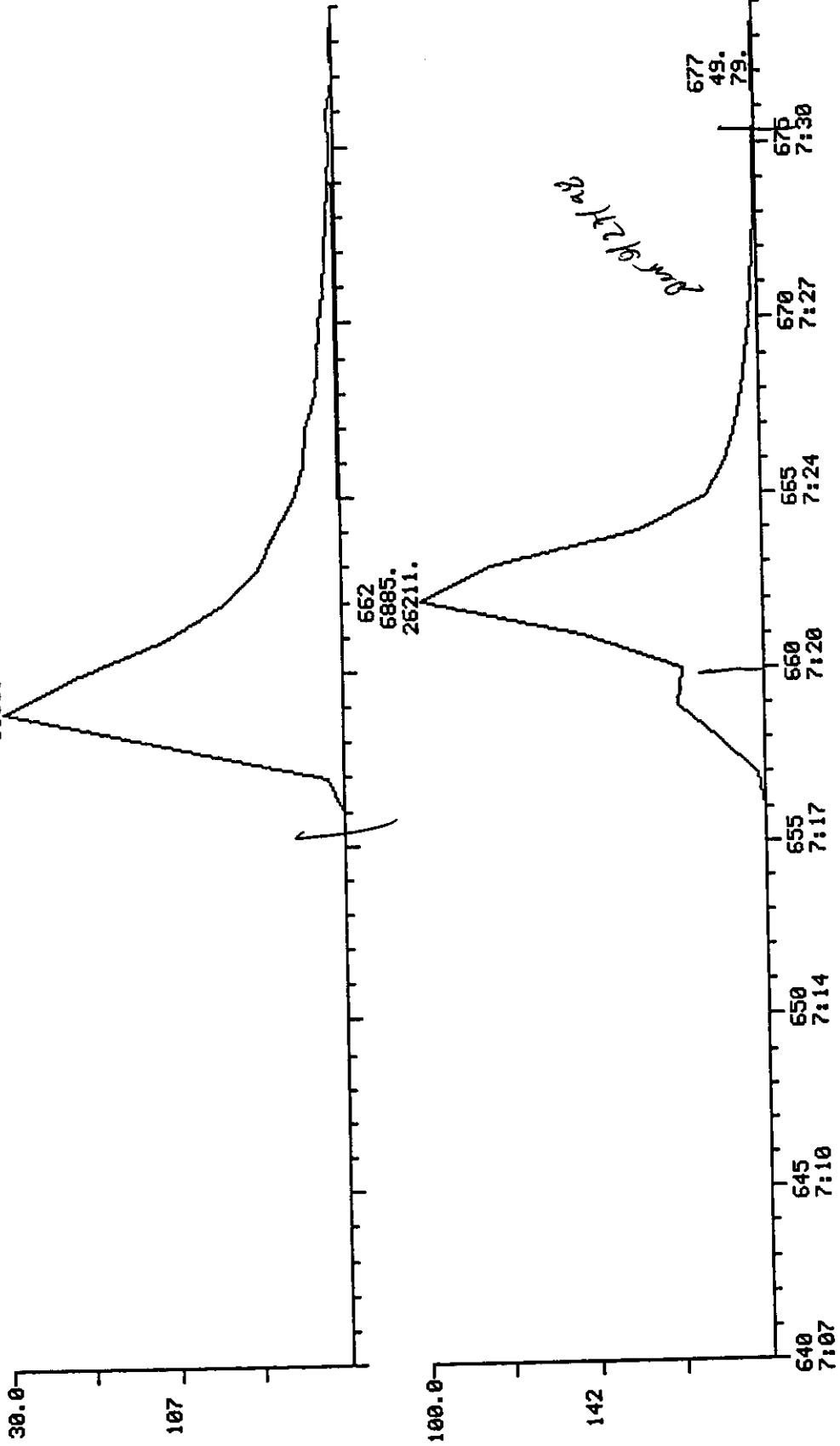
MASS CHROMATOGRAMS  
08/27/98 9:56:00  
SAMPLE: SST0010 10UG/ML CALI VI 0706980  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 2. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20. 3

2064.

107.032  
± 0.500

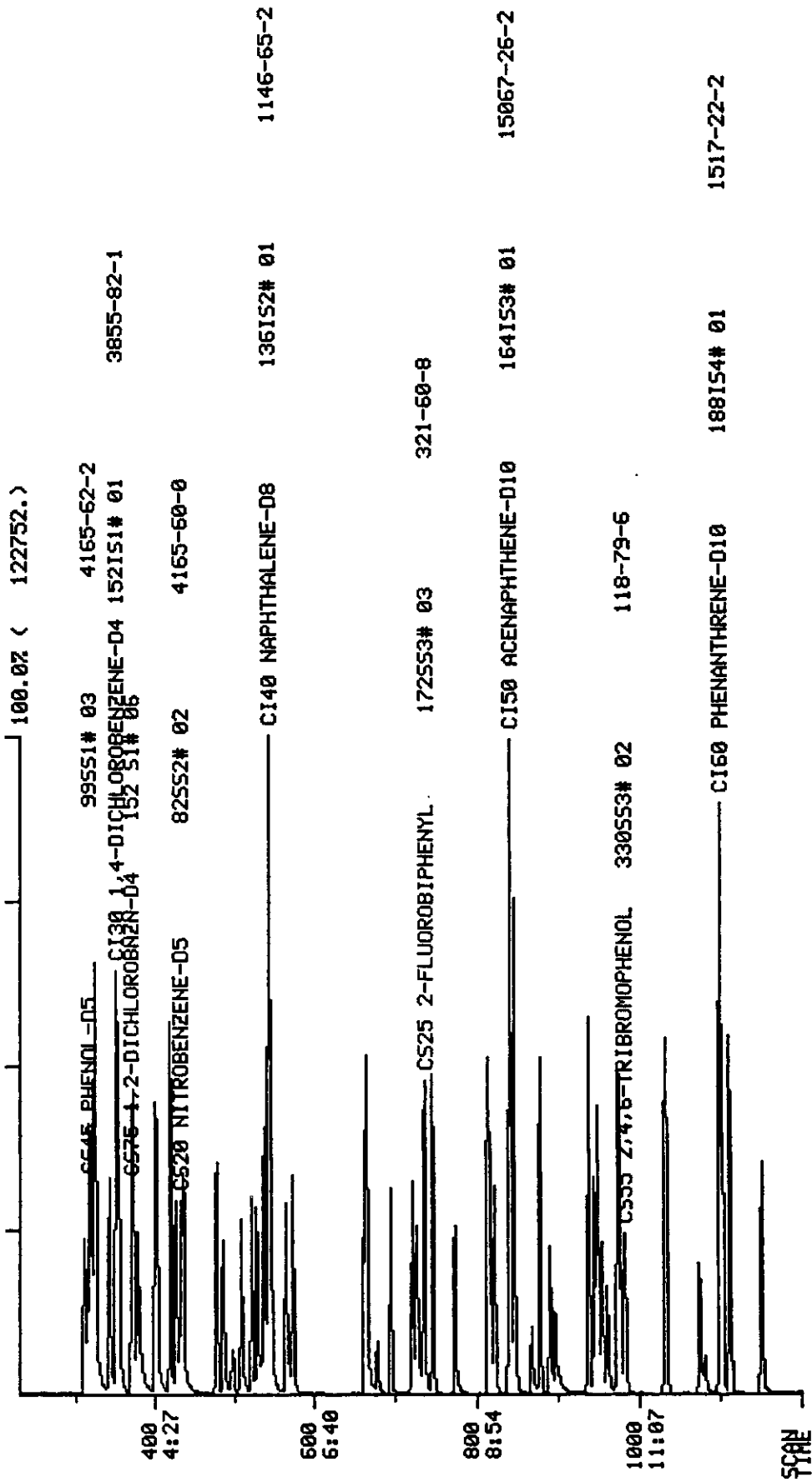
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142.042  
± 0.500



142.0

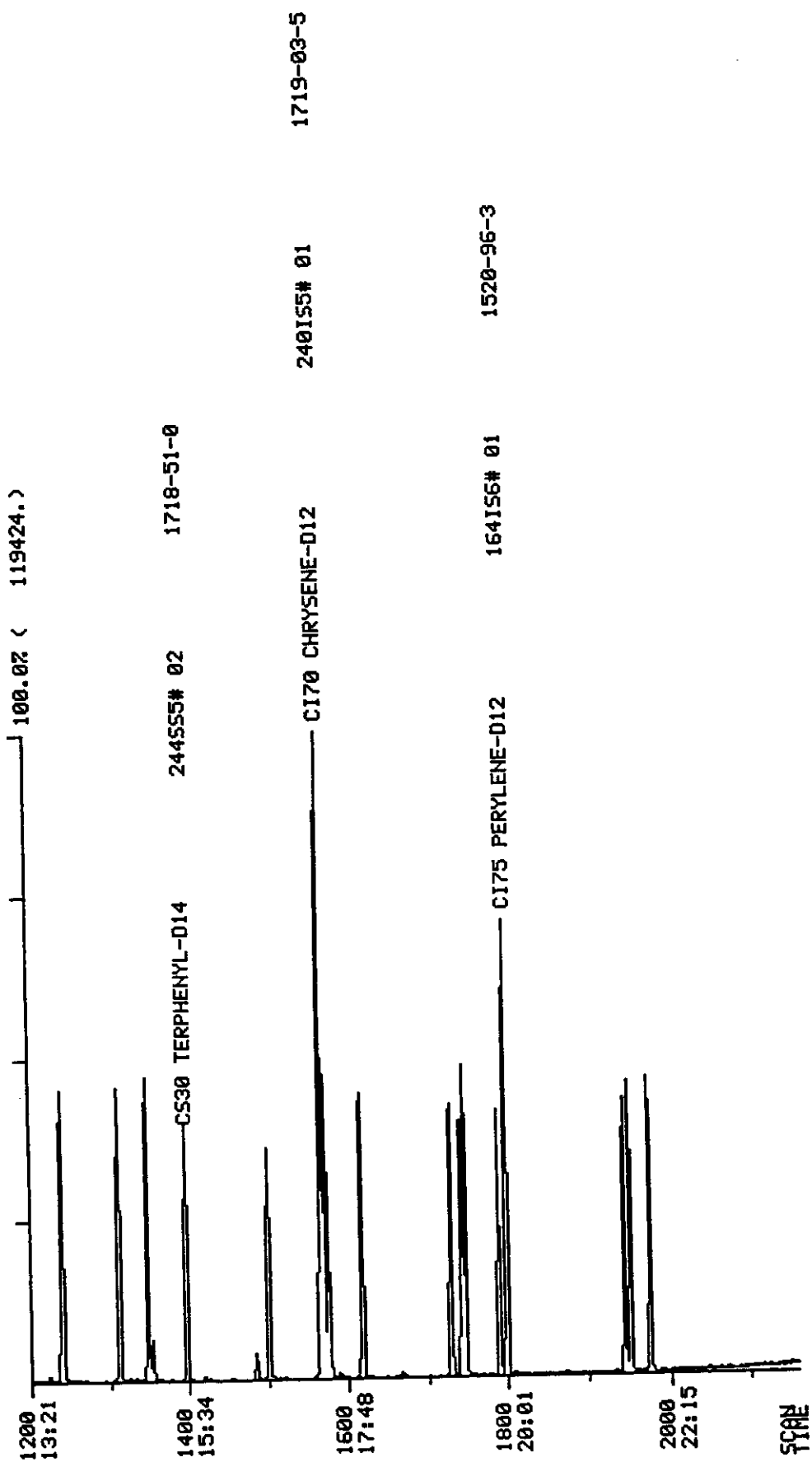
DATA FROM FILE: ST16980827B SCANS 230 TO 1200 ACQUIRED: 08/27/98 10:21:00  
 CALI: ST16980827B #3  
 SAMPLE: SSTD020 20UG/ML CALI U 070698E  
 CONDS.: INST. ID: F15



1480



DATA FROM FILE: ST16980827B SCANS 1200 TO 2158 ACQUIRED: 08/27/98 10:21:00  
 SAMPLE: SST020 20UG/ML CALI U 070698E CALI: ST16980827B #3  
 CONDS.: INST. ID: F16



Quantitation Report File: ST16980827B

Data: ST16980827B.TI

08/27/98 10:21:00

Sample: SST020 20UG/ML CALI V 070698E

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
1	CI30 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	CI40 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	CI50 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	CI60 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	CI70 CHRYSENE-D12	240IS5# 01	1719-03-5
6	CI75 PERYLENE-D12	164IS6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBENZENE-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANILINE	65 S3# 45	88-74-4

No	Name		
48	C535 DIMETHYLPHthalate	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROToluene	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	350	3:54	1	1.000	A BB	34646.	40.000 UG/ML	2.22
2	136	541	6:01	2	1.000	A BB	125585.	40.000 UG/ML	2.22
3	164	840	9:21	3	1.000	A BB	68486.	40.000 UG/ML	2.22
4	188	1099	12:13	4	1.000	A BV	117544.	40.000 UG/ML	2.22
5	240	1565	17:24	5	1.000	A BB	91058.	40.000 UG/ML	2.22
6	264	1797	19:59	6	1.000	A BB	75628.	40.000 UG/ML	2.22
7	82	432	4:48	2	0.799	A BV	31049.	20.000 UG/ML	1.11
8	172	734	8:10	3	0.874	A BB	45046.	20.000 UG/ML	1.11
9	244	1395	15:31	5	0.891	A BV	45608.	20.000 UG/ML	1.11
10	99	315	3:30	1	0.900	A BB	35638.	20.000 UG/ML	1.11
11	112	204	2:16	1	0.583	A BB	22364.	20.000 UG/ML	1.11
12	330	980	10:54	3	1.167	A BB	8930.	20.000 UG/ML	1.11
13	132	321	3:34	1	0.917	A BV	27038.	20.000 UG/ML	1.11
14	152	369	4:06	1	1.054	A BB	17300.	20.000 UG/ML	1.11
15	NOT FOUND								
16	74	109	1:13	1	0.311	A BB	17503.	20.000 UG/ML	1.11
17	93	310	3:27	1	0.886	A BV	34446.	20.000 UG/ML	1.11
18	94	317	3:32	1	0.906	A VB	35279.	20.000 UG/ML	1.11
19	93	323	3:36	1	0.923	A VB	25381.	20.000 UG/ML	1.11
20	128	323	3:36	1	0.923	A BB	27177.	20.000 UG/ML	1.11
21	146	341	3:48	1	0.974	A BV	28889.	20.000 UG/ML	1.11
22	146	352	3:55	1	1.006	A VB	29367.	20.000 UG/ML	1.11
23	108	377	4:12	1	1.077	A BB	14744.	20.000 UG/ML	1.11
24	146	371	4:08	1	1.060	A BB	27815.	20.000 UG/ML	1.11
25	108	399	4:26	1	1.140	A BB	20889.	20.000 UG/ML	1.11
26	45	399	4:26	1	1.140	A BB	46351.	20.000 UG/ML	1.11
27	108	425	4:44	1	1.214	A BB	22478.	20.000 UG/ML	1.11
28	70	418	4:39	1	1.194	A BV	16174.	20.000 UG/ML	1.11
29	117	418	4:39	1	1.194	A BB	12820.	20.000 UG/ML	1.11
30	77	434	4:50	2	0.802	A VB	29943.	20.000 UG/ML	1.11
31	82	476	5:18	2	0.880	A BV	48830.	20.000 UG/ML	1.11
32	139	484	5:23	2	0.895	A BB	13326.	20.000 UG/ML	1.11
33	107	506	5:38	2	0.935	A BB	19411.	20.000 UG/ML	1.11
34	93	520	5:47	2	0.961	A BV	33193.	20.000 UG/ML	1.11
35	162	526	5:51	2	0.972	A BB	19655.	20.000 UG/ML	1.11
36	122	542	6:02	2	1.002	qedt ②	16146.	20.000 UG/ML	1.11
37	180	535	5:57	2	0.989	A BB	21964.	20.000 UG/ML	1.11
38	128	544	6:03	2	1.006	A BB	68435.	20.000 UG/ML	1.11
39	127	563	6:16	2	1.041	A BB	29283.	20.000 UG/ML	1.11
40	225	571	6:21	2	1.055	A BB	11206.	20.000 UG/ML	1.11
41	107	659	7:20	2	1.218	A BB	21143.	20.000 UG/ML	1.11
42	142	662	7:22	2	1.224	qedt ②	52750.	20.000 UG/ML	1.11
43	237	692	7:42	3	0.824	A BB	13238.	20.000 UG/ML	1.11
44	196	718	7:59	3	0.855	A BV	13337.	20.000 UG/ML	1.11
45	196	724	8:03	3	0.862	A VB	14963.	20.000 UG/ML	1.11
46	162	744	8:17	3	0.886	A BB	43660.	20.000 UG/ML	1.11
47	65	772	8:35	3	0.919	A BV	16816.	20.000 UG/ML	1.11
48	163	816	9:05	3	0.971	A BB	48724.	20.000 UG/ML	1.11
49	152	813	9:03	3	0.968	A BB	67074.	20.000 UG/ML	1.11
50	165	821	9:08	3	0.977	A BB	11592.	20.000 UG/ML	1.11

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	6:01	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:21	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:25	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:59	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:49	1.00	0.800	1.00	20.00	20.00	0.494	0.494	1.00
8	8:11	1.00	0.875	1.00	20.00	20.00	1.315	1.315	1.00
9	15:32	1.00	0.891	1.00	20.00	20.00	1.002	1.002	1.00
10	3:32	0.99	0.906	0.99	20.00	20.00	2.057	2.057	1.00
11	2:15	1.00	0.580	1.00	20.00	20.00	1.291	1.291	1.00
12	10:55	1.00	1.168	1.00	20.00	20.00	0.261	0.261	1.00
13	3:34	1.00	0.917	1.00	20.00	20.00	1.561	1.561	1.00
14	4:07	1.00	1.057	1.00	20.00	20.00	0.999	0.999	1.00
15	12:39		0.941						
16	1:12	1.01	0.309	1.01	20.00	20.00	1.010	1.010	1.00
17	3:27	1.00	0.886	1.00	20.00	20.00	1.988	1.988	1.00
18	3:33	0.99	0.911	0.99	20.00	20.00	2.037	2.037	1.00
19	3:36	1.00	0.926	1.00	20.00	20.00	1.465	1.465	1.00
20	3:36	1.00	0.926	1.00	20.00	20.00	1.569	1.569	1.00
21	3:48	1.00	0.974	1.00	20.00	20.00	1.668	1.668	1.00
22	3:55	1.00	1.006	1.00	20.00	20.00	1.695	1.695	1.00
23	4:13	0.99	1.083	0.99	20.00	20.00	0.851	0.851	1.00
24	4:08	1.00	1.063	1.00	20.00	20.00	1.606	1.606	1.00
25	4:27	1.00	1.143	1.00	20.00	20.00	1.206	1.206	1.00
26	4:26	1.00	1.140	1.00	20.00	20.00	2.676	2.676	1.00
27	4:44	1.00	1.217	1.00	20.00	20.00	1.298	1.298	1.00
28	4:40	1.00	1.200	1.00	20.00	20.00	0.934	0.934	1.00
29	4:39	1.00	1.194	1.00	20.00	20.00	0.740	0.740	1.00
30	4:51	1.00	0.806	1.00	20.00	20.00	0.477	0.477	1.00
31	5:19	1.00	0.884	1.00	20.00	20.00	0.778	0.778	1.00
32	5:24	1.00	0.896	1.00	20.00	20.00	0.212	0.212	1.00
33	5:39	1.00	0.939	1.00	20.00	20.00	0.309	0.309	1.00
34	5:48	1.00	0.965	1.00	20.00	20.00	0.529	0.529	1.00
35	5:52	1.00	0.974	1.00	20.00	20.00	0.313	0.313	1.00
36	6:10	0.98	1.024	0.98	20.00	20.00	0.257	0.257	1.00
37	5:57	1.00	0.989	1.00	20.00	20.00	0.350	0.350	1.00
38	6:04	1.00	1.007	1.00	20.00	20.00	1.090	1.090	1.00
39	6:16	1.00	1.043	1.00	20.00	20.00	0.466	0.466	1.00
40	6:22	1.00	1.057	1.00	20.00	20.00	0.178	0.178	1.00
41	7:20	1.00	1.220	1.00	20.00	20.00	0.337	0.337	1.00
42	7:22	1.00	1.226	1.00	20.00	20.00	0.840	0.840	1.00
43	7:42	1.00	0.824	1.00	20.00	20.00	0.387	0.387	1.00
44	8:00	1.00	0.856	1.00	20.00	20.00	0.389	0.389	1.00
45	8:03	1.00	0.862	1.00	20.00	20.00	0.437	0.437	1.00
46	8:17	1.00	0.887	1.00	20.00	20.00	1.275	1.275	1.00
47	8:37	1.00	0.921	1.00	20.00	20.00	0.491	0.491	1.00
48	9:06	1.00	0.974	1.00	20.00	20.00	1.423	1.423	1.00
49	9:03	1.00	0.969	1.00	20.00	20.00	1.959	1.959	1.00
50	9:09	1.00	0.980	1.00	20.00	20.00	0.339	0.339	1.00

Quantitation Report File: ST16980827B

Data: ST16980827B.TI

08/27/98 10:21:00

Sample: SSTD020 20UG/ML CALI V 070698E

Conds.: INST. ID: F16

Formula: 1UL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name			
51	C545 3-NITROANILINE	138 S3# 75		99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC		83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP		51-28-5
54	C565 DIBENZOFURAN	168 S3# 90		132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP		100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105		121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130		84-66-2
58	C590 FLUORENE	166 S3#135		86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140		7005-72-3
60	C595 4-NITROANALINE	138 S3#150		100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10		534-52-1
62	C615 N-NITROSODIPHENYLAMINE	169 S4# 15CC		87-30-6
63	C620 AZOBENZENE	77 S4# 20		103-33-3
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30		101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40		118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50		87-86-5
67	C640 PHENANTHRENE	178 S4# 65		85-01-8
68	C645 ANTHRACENE	178 S4# 70		120-12-7
69	C647 CARBAZOLE	167 S4# 80		86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85		84-74-2
71	C655 FLUORANTHENE	202 S4#110CC		206-44-0
72	C710 BENZIDINE	184 S5# 10		92-81-5
73	C715 PYRENE	202 S5# 15		129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40		85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50		56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55		91-94-1
77	C740 CHRYSENE	228 S5# 60		218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65		117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC		117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15		205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25		207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35		50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55		193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60		53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65		191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	138	846	9:25	3	1.007	A BB	13084.	20.000 UG/ML	1.11
52	153	845	9:24	3	1.006	A BB	42016.	20.000 UG/ML	1.11
53	184	866	9:38	3	1.031	A BV	5334.	20.000 UG/ML	1.11
54	168	877	9:45	3	1.044	A BB	61028.	20.000 UG/ML	1.11
55	109	895	9:57	3	1.065	A BB	5627.	20.000 UG/ML	1.11
56	165	889	9:53	3	1.058	A BB	14108.	20.000 UG/ML	1.11
57	149	944	10:30	3	1.124	A BB	50078.	20.000 UG/ML	1.11
58	166	937	10:25	3	1.115	A BV	47880.	20.000 UG/ML	1.11

1485

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	XTot
59	204	948	10:33	3	1.129	A BB	22148.	20.000 UG/ML	1.11
60	138	953	10:36	3	1.135	A BB	11167.	20.000 UG/ML	1.11
61	198	959	10:40	4	0.873	A BV	7140.	20.000 UG/ML	1.11
62	169	972	10:49	4	0.884	A BV	34182.	20.000 UG/ML	1.11
63	77	975	10:51	4	0.887	A BB	71792.	20.000 UG/ML	1.11
64	248	1032	11:29	4	0.939	A BB	13566.	20.000 UG/ML	1.11
65	284	1031	11:28	4	0.938	A BB	16534.	20.000 UG/ML	1.11
66	266	1073	11:56	4	0.976	A BB	8694.	20.000 UG/ML	1.11
67	178	1102	12:15	4	1.003	A BV	63550.	20.000 UG/ML	1.11
68	178	1111	12:21	4	1.011	A VB	64657.	20.000 UG/ML	1.11
69	167	1149	12:47	4	1.045	A BB	44833.	20.000 UG/ML	1.11
70	149	1238	13:46	4	1.126	A BB	80264.	20.000 UG/ML	1.11
71	202	1310	14:34	4	1.192	A BB	59268.	20.000 UG/ML	1.11
72	184	1352	15:02	5	0.864	A BB	7046.	20.000 UG/ML	1.11
73	202	1347	14:59	5	0.861	A BB	59638.	20.000 UG/ML	1.11
74	149	1499	16:40	5	0.958	A BB	30622.	20.000 UG/ML	1.11
75	228	1564	17:24	5	0.999	A BV	54385.	20.000 UG/ML	1.11
76	252	1574	17:30	5	1.006	A BB	20062.	20.000 UG/ML	1.11
77	228	1569	17:27	5	1.003	A VB	48636.	20.000 UG/ML	1.11
78	149	1615	17:58	5	1.032	A BB	44542.	20.000 UG/ML	1.11
79	149	1726	19:12	6	0.960	A BB	74084.	20.000 UG/ML	1.11
80	252	1740	19:21	6	0.968	A BV	58084.	20.000 UG/ML	1.11
81	252	1745	19:25	6	0.971	A VB	46263.	20.000 UG/ML	1.11
82	252	1787	19:53	6	0.994	A BV	47524.	20.000 UG/ML	1.11
83	276	1943	21:37	6	1.081	A BV	51246.	20.000 UG/ML	1.11
84	278	1950	21:41	6	1.085	A BB	52402.	20.000 UG/ML	1.11
85	276	1974	21:57	6	1.098	A BB	54312.	20.000 UG/ML	1.11

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:26	1.00	1.010	1.00	20.00	20.00	0.382	0.382	1.00
52	9:25	1.00	1.007	1.00	20.00	20.00	1.227	1.227	1.00
53	9:39	1.00	1.032	1.00	20.00	20.00	0.156	0.156	1.00
54	9:46	1.00	1.045	1.00	20.00	20.00	1.782	1.782	1.00
55	9:59	1.00	1.068	1.00	20.00	20.00	0.164	0.164	1.00
56	9:55	1.00	1.061	1.00	20.00	20.00	0.412	0.412	1.00
57	10:31	1.00	1.126	1.00	20.00	20.00	1.462	1.462	1.00
58	10:26	1.00	1.117	1.00	20.00	20.00	1.398	1.398	1.00
59	10:33	1.00	1.130	1.00	20.00	20.00	0.647	0.647	1.00
60	10:39	1.00	1.139	1.00	20.00	20.00	0.326	0.326	1.00
61	10:42	1.00	0.875	1.00	20.00	20.00	0.121	0.121	1.00
62	10:50	1.00	0.886	1.00	20.00	20.00	0.582	0.582	1.00
63	10:51	1.00	0.888	1.00	20.00	20.00	1.222	1.222	1.00
64	11:29	1.00	0.940	1.00	20.00	20.00	0.231	0.231	1.00
65	11:29	1.00	0.939	1.00	20.00	20.00	0.281	0.281	1.00
66	11:57	1.00	0.977	1.00	20.00	20.00	0.148	0.148	1.00
67	12:17	1.00	1.005	1.00	20.00	20.00	1.081	1.081	1.00
68	12:22	1.00	1.012	1.00	20.00	20.00	1.100	1.100	1.00
69	12:47	1.00	1.046	1.00	20.00	20.00	0.763	0.763	1.00
70	13:47	1.00	1.127	1.00	20.00	20.00	1.366	1.366	1.00
71	14:36	1.00	1.194	1.00	20.00	20.00	1.008	1.008	1.00
72	15:03	1.00	0.864	1.00	20.00	20.00	0.155	0.155	1.00
73	15:00	1.00	0.861	1.00	20.00	20.00	1.310	1.310	1.00
74	16:41	1.00	0.958	1.00	20.00	20.00	0.673	0.673	1.00
75	17:24	1.00	0.999	1.00	20.00	20.00	1.195	1.195	1.00
76	17:31	1.00	1.006	1.00	20.00	20.00	0.441	0.441	1.00
77	17:28	1.00	1.003	1.00	20.00	20.00	1.068	1.068	1.00

1486

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:58	1.00	1.032	1.00	20.00	20.00	0.978	0.978	1.00
79	19:13	1.00	0.961	1.00	20.00	20.00	1.959	1.959	1.00
80	19:23	1.00	0.969	1.00	20.00	20.00	1.536	1.536	1.00
81	19:26	1.00	0.972	1.00	20.00	20.00	1.223	1.223	1.00
82	19:54	1.00	0.996	1.00	20.00	20.00	1.257	1.257	1.00
83	21:38	1.00	1.082	1.00	20.00	20.00	1.355	1.355	1.00
84	21:43	1.00	1.086	1.00	20.00	20.00	1.386	1.386	1.00
85	21:59	1.00	1.100	1.00	20.00	20.00	1.436	1.436	1.00

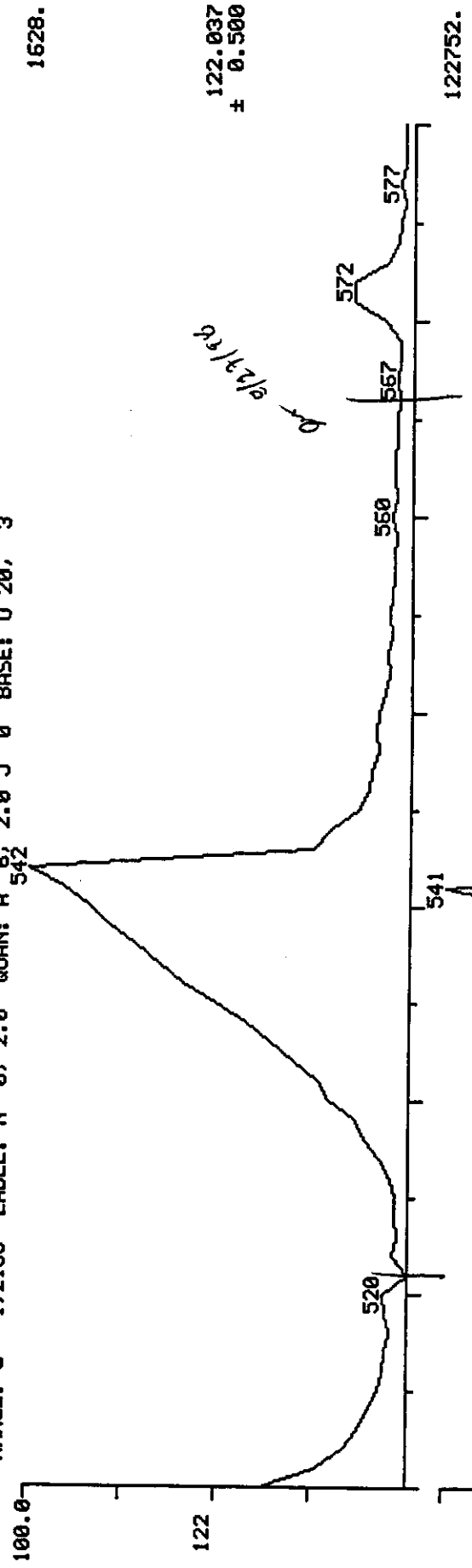
**MANUAL EDIT CODES**

1. PEAK NOT FOUND
2. POOR CHROMATOGRAPH.
3. WRONG ISOMER

**ALL MANUAL EDITS MUST BE  
INITIALED, DATED, AND COINITIALED**

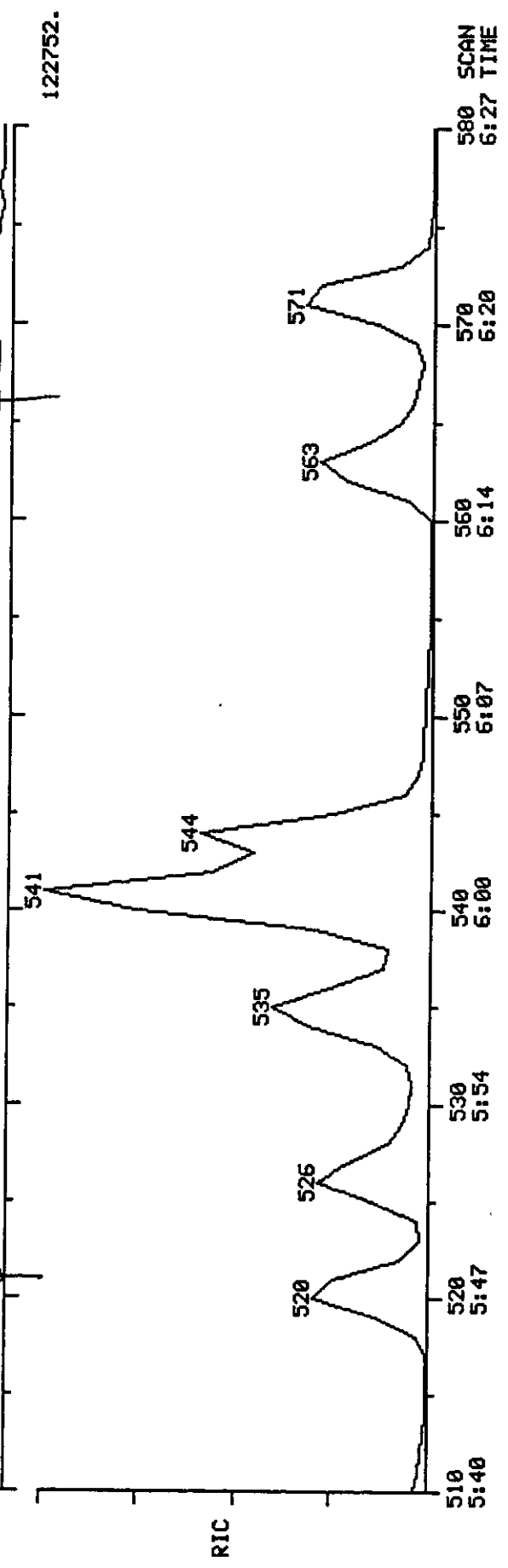
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 08/27/98 10:21:00 DATA: ST169808278 #1 SCANS 510 TO 580  
 CALI: ST169808278 #3  
 SAMPLE: SSTD020 20UG/ML CALI U 070698E  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

1628.



122.037  
± 0.500

122752.

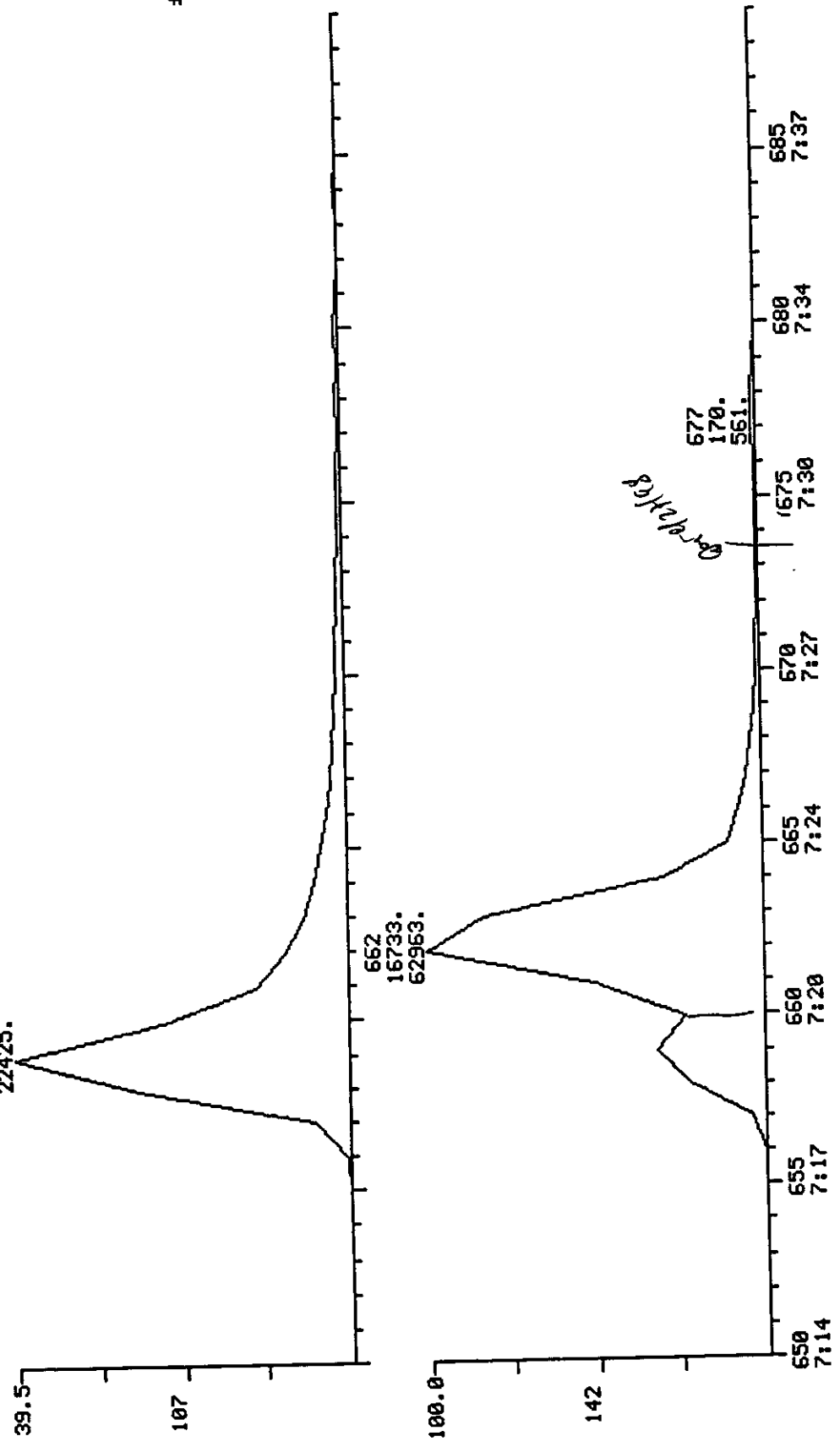


3241



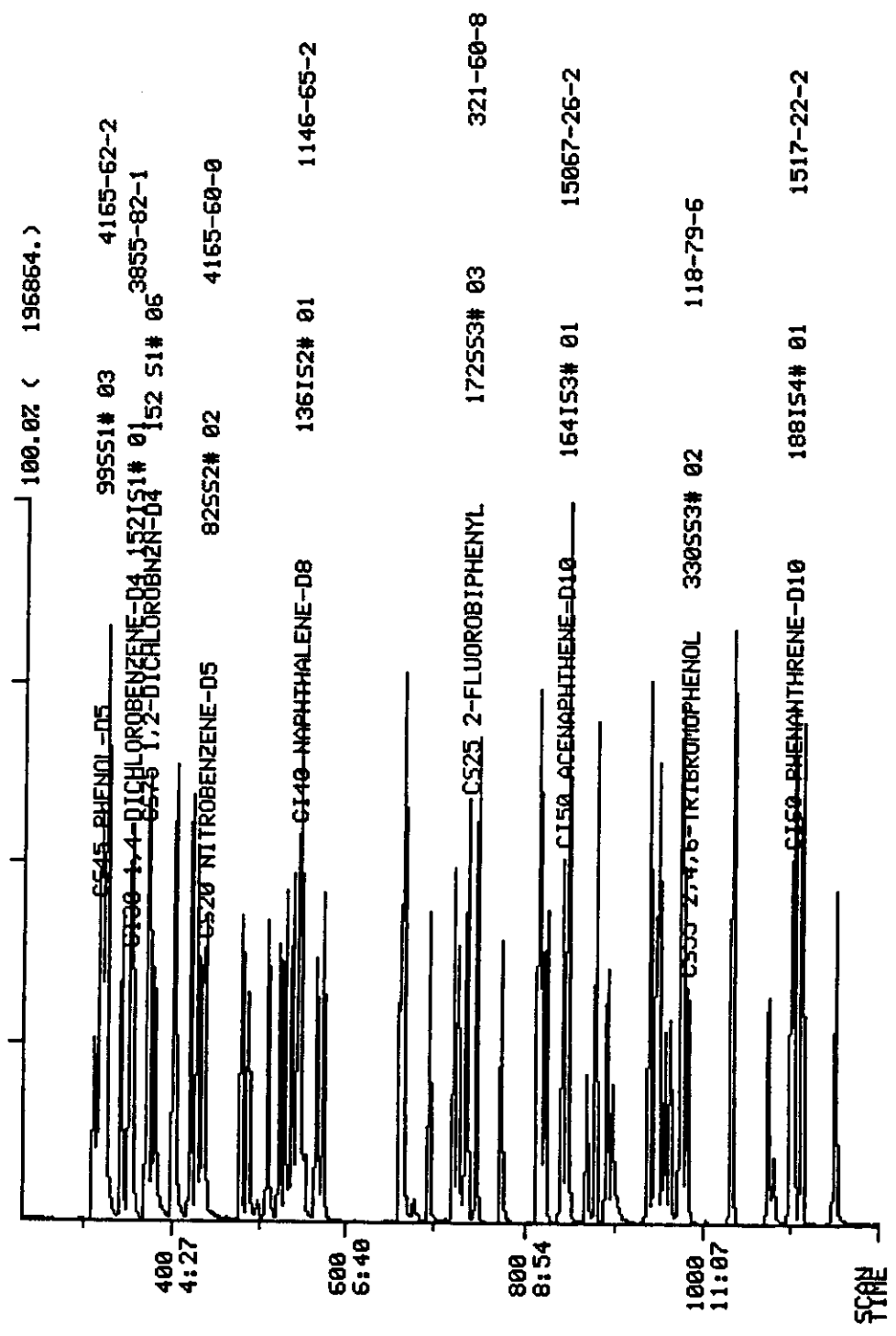
MASS CHROMATOGRAMS  
 08/27/98 10:21:00  
 SAMPLE: SSTD020 20UG/ML CALI U 070698E  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3  
 DATA: ST16980827B #1  
 CALI: ST16980827B #3  
 SCANS 650 TO 689

5608.  
 107.032  
 ± 0.500

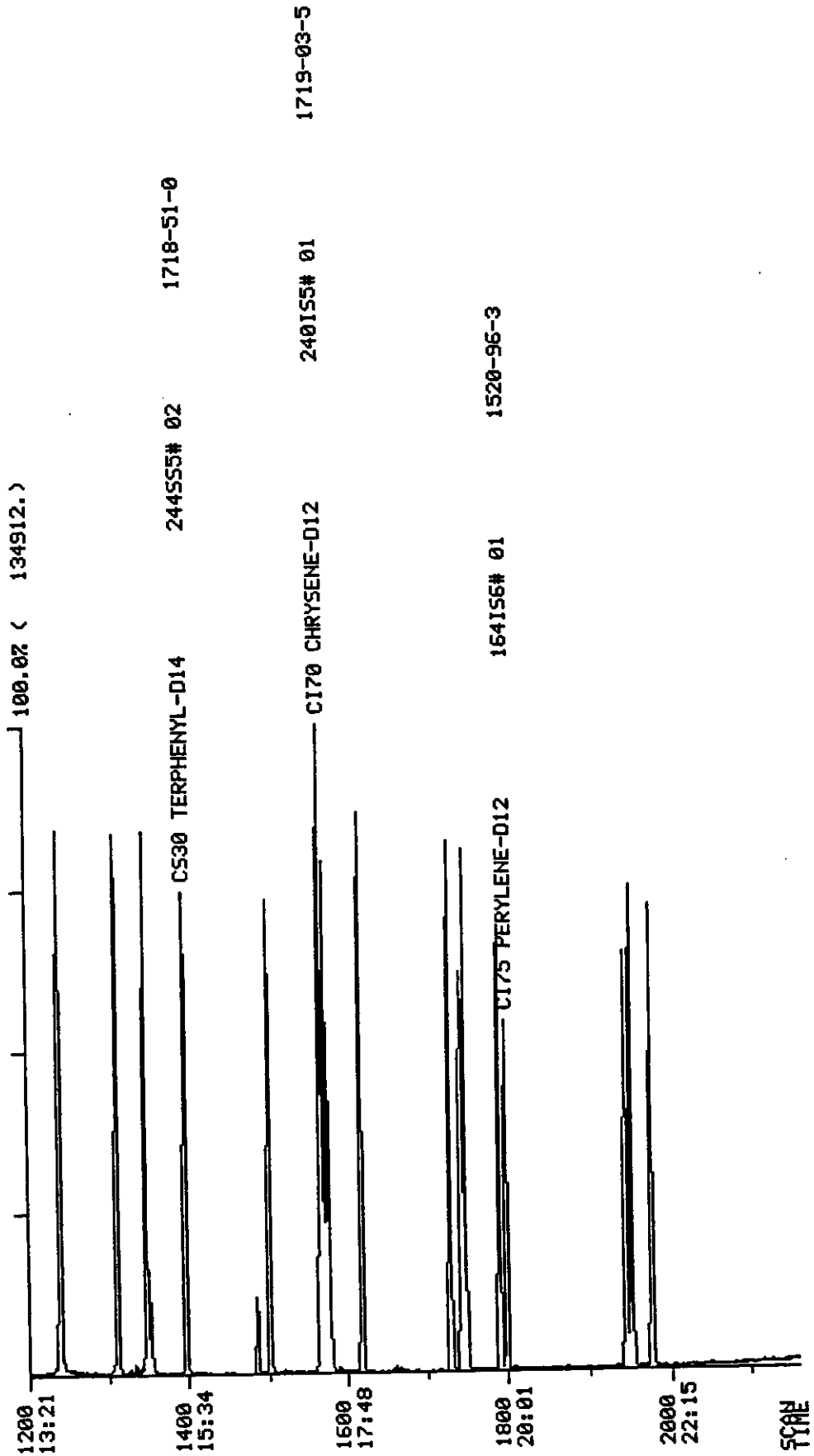


1489

DATA FROM FILE: ST16980827C SCANS 230 TO 1200 ACQUIRED: 08/27/98 10:51:00  
 CALI: ST16980827C #3  
 SAMPLE: SST0050 50UG/ML CALI IV 0706598F  
 CONDS.: INST. ID: F16



DATA FROM FILE: ST16980827C SCANS 1200 TO 2158 ACQUIRED: 08/27/98 10:51:00  
CALI: ST16980827C #3  
SAMPLE: SSTD050 50UG/ML CALI IV 070698F  
CONDS.: INST. ID: F16



Data: ST16980827C.TI

08/27/98 10:51:00

Sample: SSTD050 50UG/ML CALI IV 070698F

Conds.: INST. ID: F16

Formula: IUL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
1	CI30 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	CI40 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	CI50 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	CI60 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	CI70 CHRYSENE-D12	240IS5# 01	1719-03-5
6	CI75 PERYLENE-D12	164IS6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBENZENE-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANILINE	65 S3# 45	88-74-4

No	Name		
48	C535 DIMETHYLPHTHALATE	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROTOLUENE	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	350	3:54	1	1.000	A BB	31156.	40.000 UG/ML	0.97
2	136	541	6:01	2	1.000	A BB	114584.	40.000 UG/ML	0.97
3	164	840	9:21	3	1.000	A BB	63340.	40.000 UG/ML	0.97
4	188	1099	12:13	4	1.000	A BV	109036.	40.000 UG/ML	0.97
5	240	1566	17:25	5	1.000	A VV	77798.	40.000 UG/ML	0.97
6	264	1797	19:59	6	1.000	A BB	63787.	40.000 UG/ML	0.97
7	82	433	4:49	2	0.800	A BB	67191.	50.000 UG/ML	1.21
8	172	734	8:10	3	0.874	A BB	99214.	50.000 UG/ML	1.21
9	244	1396	15:32	5	0.891	A BB	94942.	50.000 UG/ML	1.21
10	99	316	3:31	1	0.903	A BB	80179.	50.000 UG/ML	1.21
11	112	204	2:16	1	0.583	A BB	51862.	50.000 UG/ML	1.21
12	330	981	10:55	3	1.168	A BB	18938.	50.000 UG/ML	1.21
13	132	321	3:34	1	0.917	A BB	58440.	50.000 UG/ML	1.21
14	152	370	4:07	1	1.057	A BB	36852.	50.000 UG/ML	1.21
15	NOT FOUND								
16	74	109	1:13	1	0.311	A VB	39558.	50.000 UG/ML	1.21
17	93	310	3:27	1	0.886	A BV	70664.	50.000 UG/ML	1.21
18	94	318	3:32	1	0.909	A VB	79234.	50.000 UG/ML	1.21
19	93	324	3:36	1	0.926	A VB	56008.	50.000 UG/ML	1.21
20	128	324	3:36	1	0.926	A BB	59290.	50.000 UG/ML	1.21
21	146	341	3:48	1	0.974	A BV	63015.	50.000 UG/ML	1.21
22	146	352	3:55	1	1.006	A VB	64864.	50.000 UG/ML	1.21
23	108	378	4:12	1	1.080	A BB	33967.	50.000 UG/ML	1.21
24	146	372	4:08	1	1.063	A BB	57718.	50.000 UG/ML	1.21
25	108	400	4:27	1	1.143	A BB	47563.	50.000 UG/ML	1.21
26	45	399	4:26	1	1.140	A BB	101138.	50.000 UG/ML	1.21
27	108	426	4:44	1	1.217	A BB	52789.	50.000 UG/ML	1.21
28	70	420	4:40	1	1.200	A BV	36545.	50.000 UG/ML	1.21
29	117	418	4:39	1	1.194	A BB	27750.	50.000 UG/ML	1.21
30	77	436	4:51	2	0.806	A VB	65237.	50.000 UG/ML	1.21
31	82	477	5:18	2	0.882	A BB	110794.	50.000 UG/ML	1.21
32	139	484	5:23	2	0.895	A BB	31214.	50.000 UG/ML	1.21
33	107	507	5:38	2	0.937	A BB	47890.	50.000 UG/ML	1.21
34	93	521	5:48	2	0.963	A BV	74000.	50.000 UG/ML	1.21
35	162	527	5:52	2	0.974	A BB	44945.	50.000 UG/ML	1.21
36	122	552	6:08	2	1.020	qedt(2)	39970.	50.000 UG/ML	1.21
37	180	535	5:57	2	0.989	A BB	47850.	50.000 UG/ML	1.21
38	128	545	6:04	2	1.007	A BB	148742.	50.000 UG/ML	1.21
39	127	564	6:16	2	1.043	A BB	65035.	50.000 UG/ML	1.21
40	225	572	6:22	2	1.057	A BB	24296.	50.000 UG/ML	1.21
41	107	659	7:20	2	1.218	A BB	50489.	50.000 UG/ML	1.21
42	142	663	7:22	2	1.226	qedt(2)	118484.	50.000 UG/ML	1.21
43	237	692	7:42	3	0.824	A BB	29806.	50.000 UG/ML	1.21
44	196	719	8:00	3	0.856	A BV	32070.	50.000 UG/ML	1.21
45	196	724	8:03	3	0.862	A VB	33958.	50.000 UG/ML	1.21
46	162	745	8:17	3	0.887	A BB	95618.	50.000 UG/ML	1.21
47	65	773	8:36	3	0.920	A BB	40406.	50.000 UG/ML	1.21
48	163	817	9:05	3	0.973	A BV	110200.	50.000 UG/ML	1.21
49	152	814	9:03	3	0.969	A BB	150122.	50.000 UG/ML	1.21
50	165	823	9:09	3	0.980	A BB	27004.	50.000 UG/ML	1.21

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	6:01	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:21	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:25	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:59	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:49	1.00	0.800	1.00	50.00	50.00	0.469	0.469	1.00
8	8:11	1.00	0.875	1.00	50.00	50.00	1.253	1.253	1.00
9	15:32	1.00	0.891	1.00	50.00	50.00	0.976	0.976	1.00
10	3:32	1.00	0.906	1.00	50.00	50.00	2.059	2.059	1.00
11	2:15	1.00	0.580	1.00	50.00	50.00	1.332	1.332	1.00
12	10:55	1.00	1.168	1.00	50.00	50.00	0.239	0.239	1.00
13	3:34	1.00	0.917	1.00	50.00	50.00	1.501	1.501	1.00
14	4:07	1.00	1.057	1.00	50.00	50.00	0.946	0.946	1.00
15	12:39		0.941						
16	1:12	1.01	0.309	1.01	50.00	50.00	1.016	1.016	1.00
17	3:27	1.00	0.886	1.00	50.00	50.00	1.814	1.814	1.00
18	3:33	1.00	0.911	1.00	50.00	50.00	2.035	2.035	1.00
19	3:36	1.00	0.926	1.00	50.00	50.00	1.438	1.438	1.00
20	3:36	1.00	0.926	1.00	50.00	50.00	1.522	1.522	1.00
21	3:48	1.00	0.974	1.00	50.00	50.00	1.618	1.618	1.00
22	3:55	1.00	1.006	1.00	50.00	50.00	1.666	1.666	1.00
23	4:13	1.00	1.083	1.00	50.00	50.00	0.872	0.872	1.00
24	4:08	1.00	1.063	1.00	50.00	50.00	1.482	1.482	1.00
25	4:27	1.00	1.143	1.00	50.00	50.00	1.221	1.221	1.00
26	4:26	1.00	1.140	1.00	50.00	50.00	2.597	2.597	1.00
27	4:44	1.00	1.217	1.00	50.00	50.00	1.355	1.355	1.00
28	4:40	1.00	1.200	1.00	50.00	50.00	0.938	0.938	1.00
29	4:39	1.00	1.194	1.00	50.00	50.00	0.713	0.713	1.00
30	4:51	1.00	0.806	1.00	50.00	50.00	0.455	0.455	1.00
31	5:19	1.00	0.884	1.00	50.00	50.00	0.774	0.774	1.00
32	5:24	1.00	0.896	1.00	50.00	50.00	0.218	0.218	1.00
33	5:39	1.00	0.939	1.00	50.00	50.00	0.334	0.334	1.00
34	5:48	1.00	0.965	1.00	50.00	50.00	0.517	0.517	1.00
35	5:52	1.00	0.974	1.00	50.00	50.00	0.314	0.314	1.00
36	6:10	1.00	1.024	1.00	50.00	50.00	0.279	0.279	1.00
37	5:57	1.00	0.989	1.00	50.00	50.00	0.334	0.334	1.00
38	6:04	1.00	1.007	1.00	50.00	50.00	1.038	1.038	1.00
39	6:16	1.00	1.043	1.00	50.00	50.00	0.454	0.454	1.00
40	6:22	1.00	1.057	1.00	50.00	50.00	0.170	0.170	1.00
41	7:20	1.00	1.220	1.00	50.00	50.00	0.353	0.353	1.00
42	7:22	1.00	1.226	1.00	50.00	50.00	0.827	0.827	1.00
43	7:42	1.00	0.824	1.00	50.00	50.00	0.376	0.376	1.00
44	8:00	1.00	0.856	1.00	50.00	50.00	0.405	0.405	1.00
45	8:03	1.00	0.862	1.00	50.00	50.00	0.429	0.429	1.00
46	8:17	1.00	0.887	1.00	50.00	50.00	1.208	1.208	1.00
47	8:37	1.00	0.921	1.00	50.00	50.00	0.510	0.510	1.00
48	9:06	1.00	0.974	1.00	50.00	50.00	1.392	1.392	1.00
49	9:03	1.00	0.969	1.00	50.00	50.00	1.896	1.896	1.00
50	9:09	1.00	0.980	1.00	50.00	50.00	0.341	0.341	1.00

Quantitation Report File: ST16980827C

Data: ST16980827C.TI

08/27/98 10:51:00

Sample: SSTD050 50UG/ML CALI IV 070698F

Conds.: INST. ID: F16

Formula: 1UL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
51	C545 3-NITROANILINE	138 S3# 75	99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC	83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP	51-28-5
54	C565 DIBENZOFURAN	168 S3# 90	132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP	100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105	121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130	84-66-2
58	C590 FLUORENE	166 S3#135	86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140	7005-72-3
60	C595 4-NITROANALINE	138 S3#150	100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10	534-52-1
62	C615 N-NITROSODIPHENYLAMINE	169 S4# 15CC	87-30-6
63	C620 AZOBENZENE	77 S4# 20	103-33-3
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30	101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40	118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50	87-86-5
67	C640 PHENANTHRENE	178 S4# 65	85-01-8
68	C645 ANTHRACENE	178 S4# 70	120-12-7
69	C647 CARBAZOLE	167 S4# 80	86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85	84-74-2
71	C655 FLUORANTHENE	202 S4#110CC	206-44-0
72	C710 BENZIDINE	184 S5# 10	92-81-5
73	C715 PYRENE	202 S5# 15	129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40	85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50	56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55	91-94-1
77	C740 CHRYSENE	228 S5# 60	218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65	117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC	117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15	205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25	207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35	50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55	193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60	53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65	191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	138	847	9:25	3	1.008	A BV	31064.	50.000 UG/ML	1.21
52	153	846	9:25	3	1.007	A BB	93562.	50.000 UG/ML	1.21
53	184	867	9:39	3	1.032	A BB	15019.	50.000 UG/ML	1.21
54	168	878	9:46	3	1.045	A BB	137484.	50.000 UG/ML	1.21
55	109	896	9:58	3	1.067	A BB	14092.	50.000 UG/ML	1.21
56	165	891	9:55	3	1.061	A BB	33286.	50.000 UG/ML	1.21
57	149	945	10:31	3	1.125	A BB	114985.	50.000 UG/ML	1.21
58	166	938	10:26	3	1.117	A BV	106082.	50.000 UG/ML	1.21

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
59	204	949	10:33	3	1.130	A BB	49994.	50.000 UG/ML	1.21
60	138	956	10:38	3	1.138	A BB	25302.	50.000 UG/ML	1.21
61	198	962	10:42	4	0.875	A BV	18974.	50.000 UG/ML	1.21
62	169	973	10:49	4	0.885	A BV	76610.	50.000 UG/ML	1.21
63	77	976	10:51	4	0.888	A BB	161611.	50.000 UG/ML	1.21
64	248	1032	11:29	4	0.939	A BB	30356.	50.000 UG/ML	1.21
65	284	1031	11:28	4	0.938	A BB	34726.	50.000 UG/ML	1.21
66	266	1074	11:57	4	0.977	A BB	21227.	50.000 UG/ML	1.21
67	178	1103	12:16	4	1.004	A BV	142301.	50.000 UG/ML	1.21
68	178	1112	12:22	4	1.012	A VB	139815.	50.000 UG/ML	1.21
69	167	1150	12:47	4	1.046	A BB	99303.	50.000 UG/ML	1.21
70	149	1239	13:47	4	1.127	A BB	181488.	50.000 UG/ML	1.21
71	202	1311	14:35	4	1.193	A BB	126092.	50.000 UG/ML	1.21
72	184	1352	15:02	5	0.863	A BB	13954.	50.000 UG/ML	1.21
73	202	1348	15:00	5	0.861	A BB	125982.	50.000 UG/ML	1.21
74	149	1500	16:41	5	0.958	A BB	67310.	50.000 UG/ML	1.21
75	228	1564	17:24	5	0.999	A BV	108560.	50.000 UG/ML	1.21
76	252	1575	17:31	5	1.006	A BV	40964.	50.000 UG/ML	1.21
77	228	1570	17:28	5	1.003	A VB	97736.	50.000 UG/ML	1.21
78	149	1616	17:58	5	1.032	A BB	95888.	50.000 UG/ML	1.21
79	149	1727	19:13	6	0.961	A BB	158236.	50.000 UG/ML	1.21
80	252	1742	19:23	6	0.969	A BV	115462.	50.000 UG/ML	1.21
81	252	1747	19:26	6	0.972	A VB	93397.	50.000 UG/ML	1.21
82	252	1789	19:54	6	0.996	A BV	96136.	50.000 UG/ML	1.21
83	276	1944	21:37	6	1.082	A BV	104637.	50.000 UG/ML	1.21
84	278	1951	21:42	6	1.086	A BB	102669.	50.000 UG/ML	1.21
85	276	1976	21:59	6	1.100	A BV	108148.	50.000 UG/ML	1.21

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:26	1.00	1.010	1.00	50.00	50.00	0.392	0.392	1.00
52	9:25	1.00	1.007	1.00	50.00	50.00	1.182	1.182	1.00
53	9:39	1.00	1.032	1.00	50.00	50.00	0.190	0.190	1.00
54	9:46	1.00	1.045	1.00	50.00	50.00	1.736	1.736	1.00
55	9:59	1.00	1.068	1.00	50.00	50.00	0.178	0.178	1.00
56	9:55	1.00	1.061	1.00	50.00	50.00	0.420	0.420	1.00
57	10:31	1.00	1.126	1.00	50.00	50.00	1.452	1.452	1.00
58	10:26	1.00	1.117	1.00	50.00	50.00	1.340	1.340	1.00
59	10:33	1.00	1.130	1.00	50.00	50.00	0.631	0.631	1.00
60	10:39	1.00	1.139	1.00	50.00	50.00	0.320	0.320	1.00
61	10:42	1.00	0.875	1.00	50.00	50.00	0.139	0.139	1.00
62	10:50	1.00	0.886	1.00	50.00	50.00	0.562	0.562	1.00
63	10:51	1.00	0.888	1.00	50.00	50.00	1.186	1.186	1.00
64	11:29	1.00	0.940	1.00	50.00	50.00	0.223	0.223	1.00
65	11:29	1.00	0.939	1.00	50.00	50.00	0.255	0.255	1.00
66	11:57	1.00	0.977	1.00	50.00	50.00	0.156	0.156	1.00
67	12:17	1.00	1.005	1.00	50.00	50.00	1.044	1.044	1.00
68	12:22	1.00	1.012	1.00	50.00	50.00	1.026	1.026	1.00
69	12:47	1.00	1.046	1.00	50.00	50.00	0.729	0.729	1.00
70	13:47	1.00	1.127	1.00	50.00	50.00	1.332	1.332	1.00
71	14:36	1.00	1.194	1.00	50.00	50.00	0.925	0.925	1.00
72	15:03	1.00	0.864	1.00	50.00	50.00	0.143	0.143	1.00
73	15:00	1.00	0.861	1.00	50.00	50.00	1.295	1.295	1.00
74	16:41	1.00	0.958	1.00	50.00	50.00	0.692	0.692	1.00
75	17:24	1.00	0.999	1.00	50.00	50.00	1.116	1.116	1.00
76	17:31	1.00	1.006	1.00	50.00	50.00	0.421	0.421	1.00
77	17:28	1.00	1.003	1.00	50.00	50.00	1.005	1.005	1.00



No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:58	1.00	1.032	1.00	50.00	50.00	0.986	0.986	1.00
79	19:13	1.00	0.961	1.00	50.00	50.00	1.985	1.985	1.00
80	19:23	1.00	0.969	1.00	50.00	50.00	1.448	1.448	1.00
81	19:26	1.00	0.972	1.00	50.00	50.00	1.171	1.171	1.00
82	19:54	1.00	0.996	1.00	50.00	50.00	1.206	1.206	1.00
83	21:38	1.00	1.082	1.00	50.00	50.00	1.312	1.312	1.00
84	21:43	1.00	1.086	1.00	50.00	50.00	1.288	1.288	1.00
85	21:59	1.00	1.100	1.00	50.00	50.00	1.356	1.356	1.00

**MANUAL EDIT CODES**

1. PEAK NOT FOUND
2. POOR CHROMATOGRAPH.
3. WRONG ISOMER

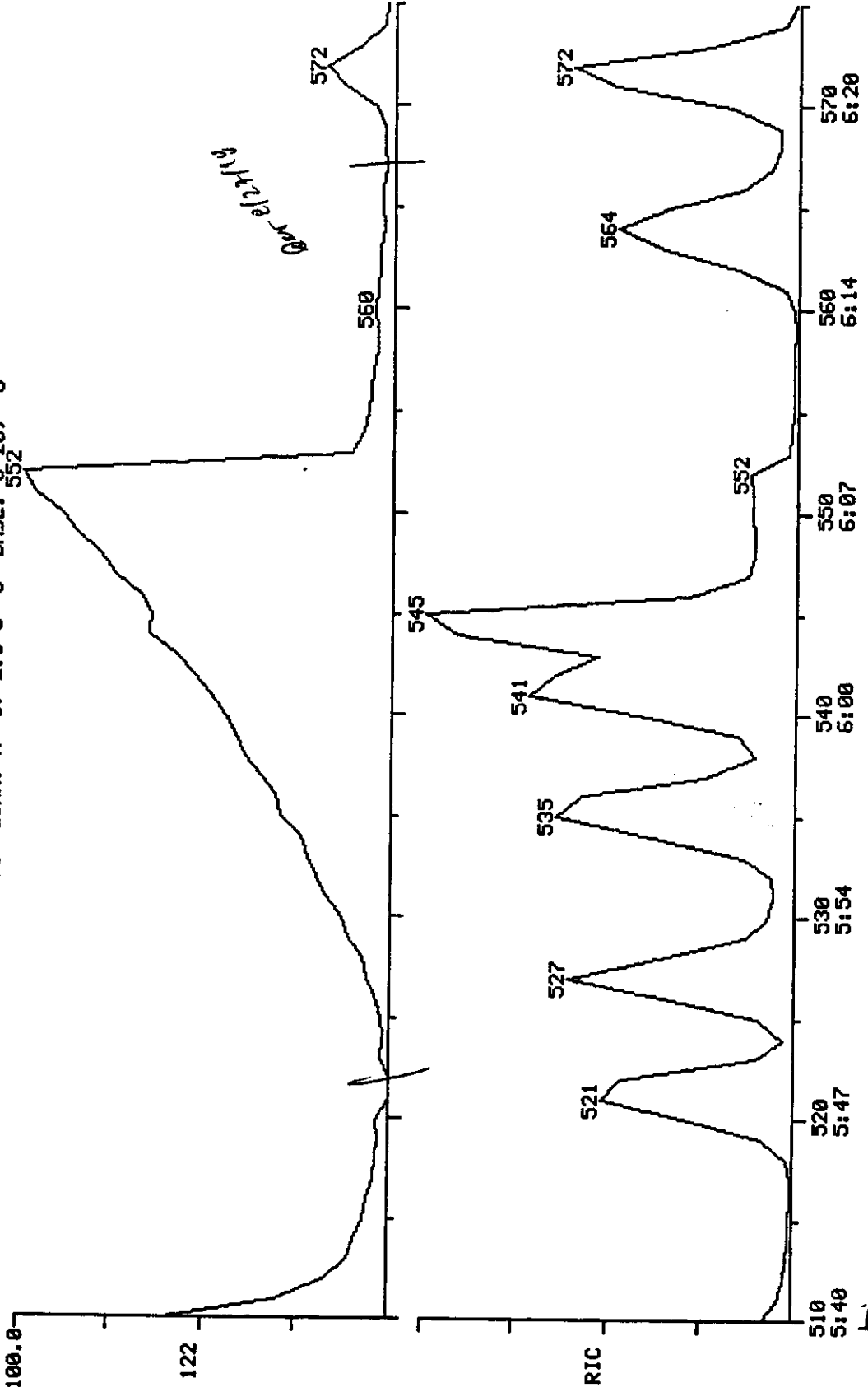
ALL MANUAL EDITS MUST BE  
INITIALED, DATED, AND CODED

RIC+MASS CHROMATOGRAM  
 08/27/98 10:51:00  
 SAMPLE: SST0650 50UG/ML CALI IV 070698F  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3  
 SCANS 510 TO 575

3040.

122.037  
± 0.500

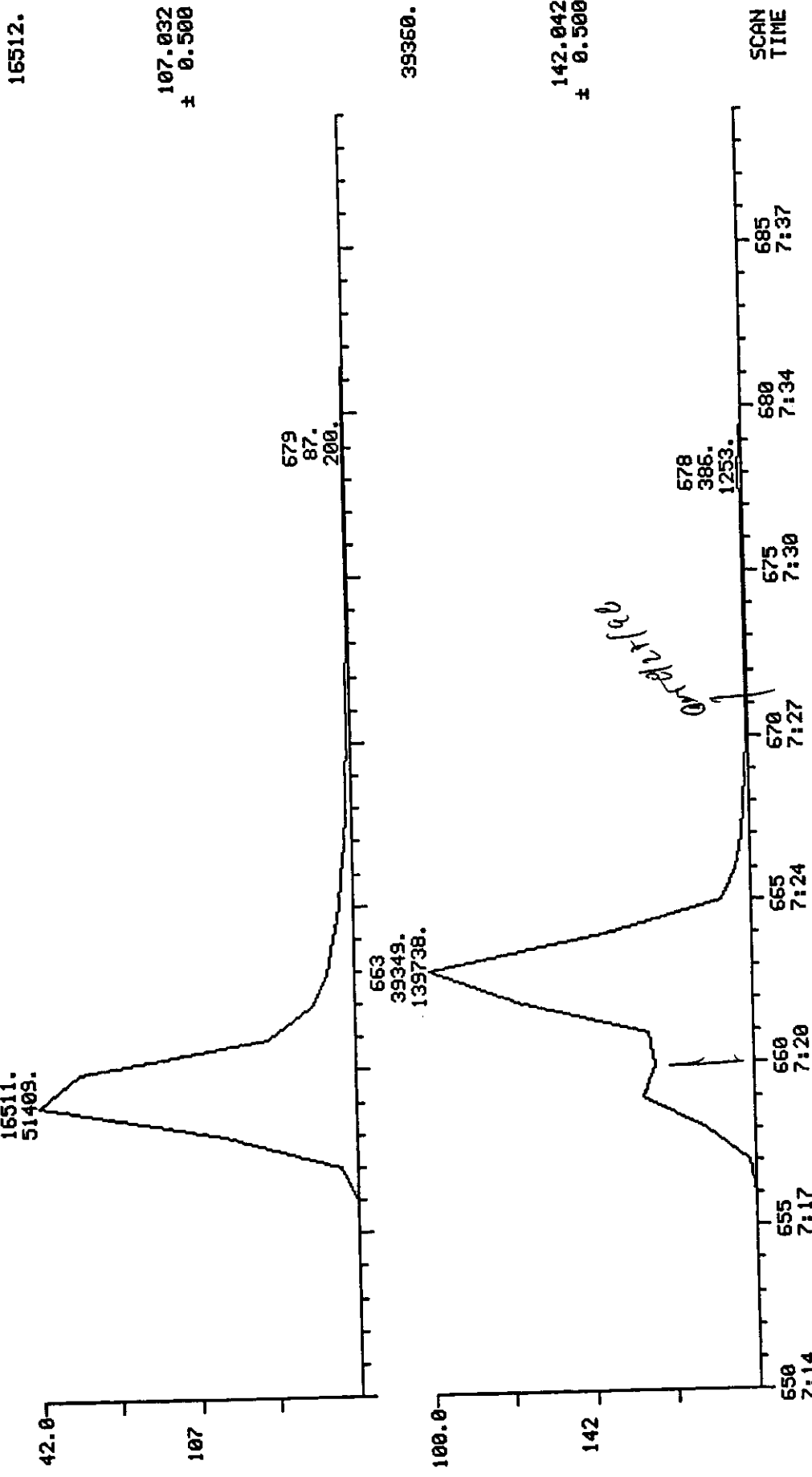
148224.



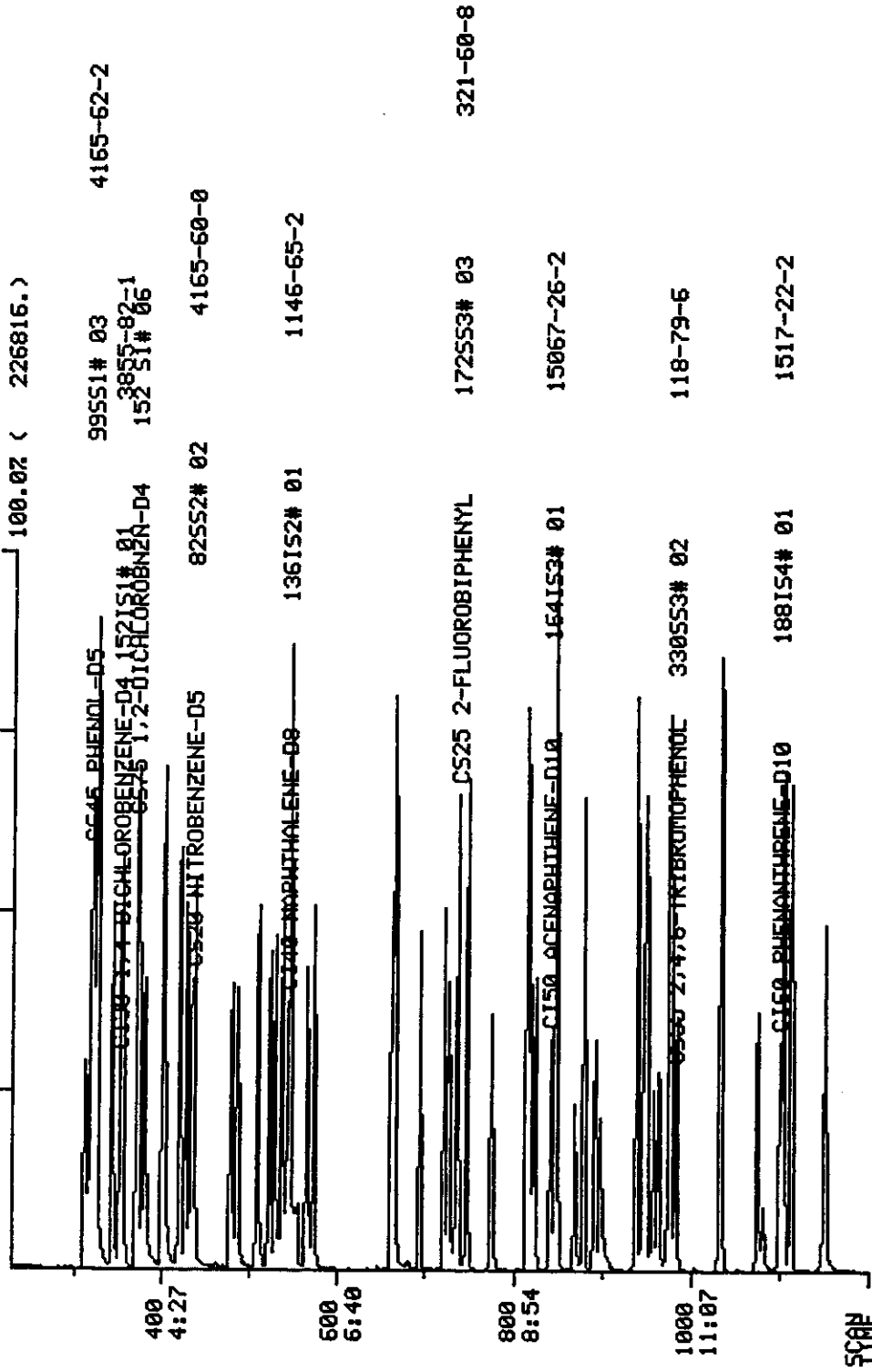
SCAN TIME

510 5:40 1488

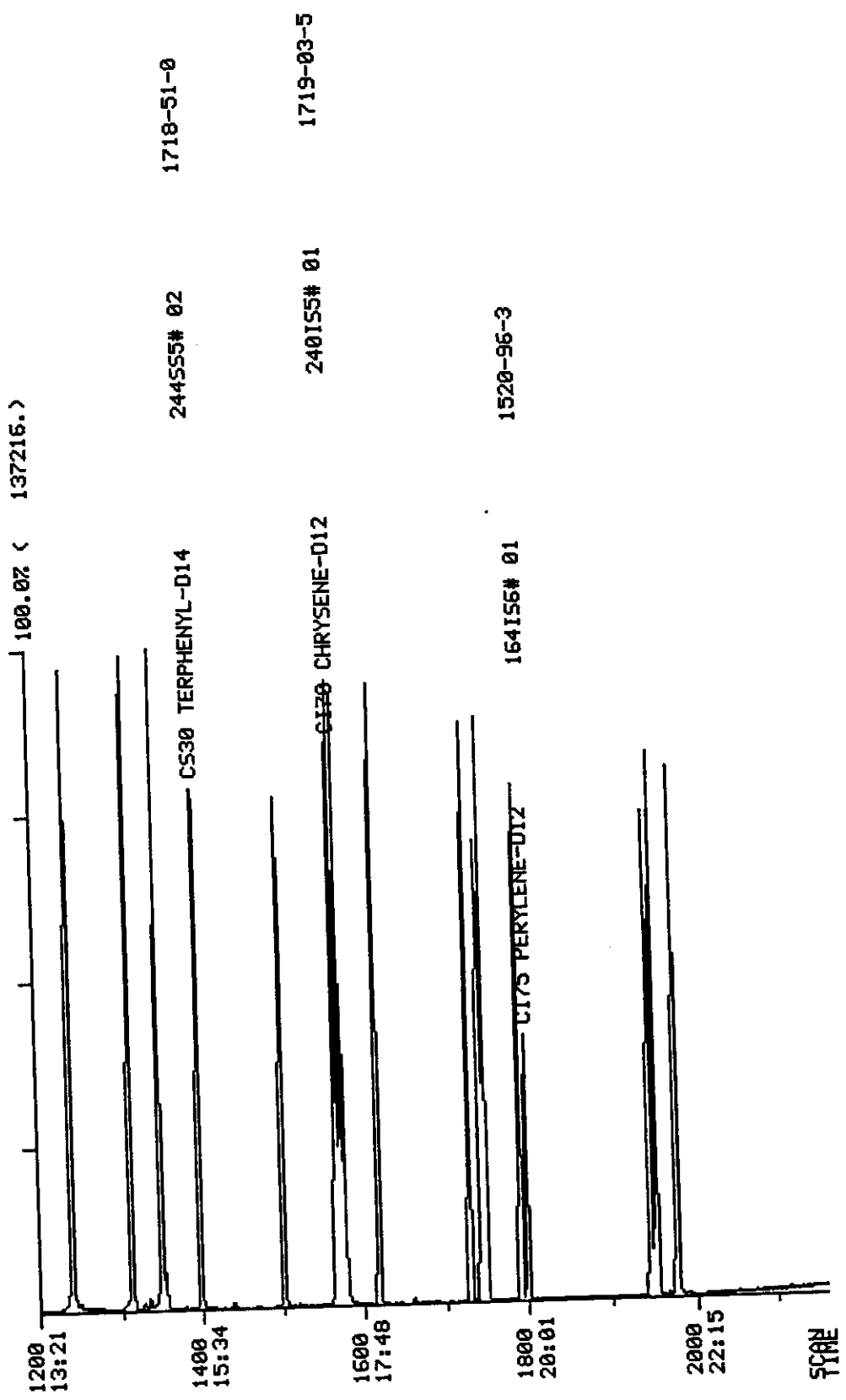
MASS CHROMATOGRAMS  
 08/27/98 10:51:00 DATA: ST16980827C #1 SCANS 650 TO 689  
 CALI: ST16980827C #3  
 SAMPLE: SSTD050 50UG/ML CALI IV 070698F  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20. 3



DATA FROM FILE: ST16980627 SCANS 230 TO 1200 ACQUIRED: 08/27/98 9:22:00  
 SAMPLE: SSTD080 80UG/ML CALI III 082198A  
 COND.: INST. ID: F16  
 CALI: ST16980627 #3



DATA FROM FILE: ST16980827 SCANS 1200 TO 2158 ACQUIRED: 08/27/98 9:22:00  
CALI: ST16980827 #3  
SAMPLE: S5TD080 80UG/ML CALI III 082198A  
CONDS.: INST. ID: F16



Data: ST16980827.T1

08/27/98 9:22:00

Sample: SST0080 80UG/ML CALI III 082198A

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBENZ-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANILINE	65 S3# 45	88-74-4

No	Name		
48	C535 DIMETHYLPHthalate	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROToluene	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	350	3:54	1	1.000	A BB	25488.	40.000 UG/ML	0.62
2	136	541	6:01	2	1.000	A BB	88640.	40.000 UG/ML	0.62
3	164	840	9:21	3	1.000	A BB	47354.	40.000 UG/ML	0.62
4	188	1099	12:13	4	1.000	A BB	80196.	40.000 UG/ML	0.62
5	240	1566	17:25	5	1.000	A BB	51312.	40.000 UG/ML	0.62
6	264	1797	19:59	6	1.000	A BB	44290.	40.000 UG/ML	0.62
7	82	433	4:49	2	0.800	A VB	89306.	80.000 UG/ML	1.23
8	172	735	8:11	3	0.875	A BV	123006.	80.000 UG/ML	1.23
9	244	1396	15:32	5	0.891	A BB	111089.	80.000 UG/ML	1.23
10	99	317	3:32	1	0.906	A BB	104277.	80.000 UG/ML	1.23
11	112	203	2:15	1	0.580	A BB	73823.	80.000 UG/ML	1.23
12	330	981	10:55	3	1.168	A BB	22048.	80.000 UG/ML	1.23
13	132	321	3:34	1	0.917	A BV	76599.	80.000 UG/ML	1.23
14	152	370	4:07	1	1.057	A BB	48888.	80.000 UG/ML	1.23
15	NOT FOUND								
16	74	108	1:12	1	0.309	A BB	55404.	80.000 UG/ML	1.23
17	93	310	3:27	1	0.886	A BV	95626.	80.000 UG/ML	1.23
18	94	319	3:33	1	0.911	A VB	109518.	80.000 UG/ML	1.23
19	93	324	3:36	1	0.926	A VB	72104.	80.000 UG/ML	1.23
20	128	324	3:36	1	0.926	A BB	81692.	80.000 UG/ML	1.23
21	146	341	3:48	1	0.974	A BV	84592.	80.000 UG/ML	1.23
22	146	352	3:55	1	1.006	A VB	86508.	80.000 UG/ML	1.23
23	108	379	4:13	1	1.083	A BB	46538.	80.000 UG/ML	1.23
24	146	372	4:08	1	1.063	A BB	80185.	80.000 UG/ML	1.23
25	108	400	4:27	1	1.143	A BB	61644.	80.000 UG/ML	1.23
26	45	399	4:26	1	1.140	A BB	131645.	80.000 UG/ML	1.23
27	108	426	4:44	1	1.217	A BB	65587.	80.000 UG/ML	1.23
28	70	420	4:40	1	1.200	A BV	47937.	80.000 UG/ML	1.23
29	117	418	4:39	1	1.194	A BB	36710.	80.000 UG/ML	1.23
30	77	436	4:51	2	0.806	A VB	86151.	80.000 UG/ML	1.23
31	82	478	5:19	2	0.884	A BB	137948.	80.000 UG/ML	1.23
32	139	485	5:24	2	0.896	A VB	41152.	80.000 UG/ML	1.23
33	107	508	5:39	2	0.939	A BB	70834.	80.000 UG/ML	1.23
34	93	522	5:48	2	0.965	A BV	95388.	80.000 UG/ML	1.23
35	162	527	5:52	2	0.974	A BB	58887.	80.000 UG/ML	1.23
36	122	554	6:10	2	1.024	qedt (2)	47914.	80.000 UG/ML	1.23
37	180	535	5:57	2	0.989	A BB	62130.	80.000 UG/ML	1.23
38	128	545	6:04	2	1.007	A BB	195384.	80.000 UG/ML	1.23
39	127	564	6:16	2	1.043	A BB	85089.	80.000 UG/ML	1.23
40	225	572	6:22	2	1.057	A BB	31304.	80.000 UG/ML	1.23
41	107	660	7:20	2	1.220	A BB	64626.	80.000 UG/ML	1.23
42	142	663	7:22	2	1.226	qedt (2)	134249.	80.000 UG/ML	1.23
43	237	692	7:42	3	0.824	A BB	40564.	80.000 UG/ML	1.23
44	196	719	8:00	3	0.856	A BV	41178.	80.000 UG/ML	1.23
45	196	724	8:03	3	0.862	A VB	41938.	80.000 UG/ML	1.23
46	162	745	8:17	3	0.887	A BB	121530.	80.000 UG/ML	1.23
47	65	774	8:37	3	0.921	A BB	49570.	80.000 UG/ML	1.23
48	163	818	9:06	3	0.974	A BV	135082.	80.000 UG/ML	1.23
49	152	814	9:03	3	0.969	A BB	193878.	80.000 UG/ML	1.23
50	165	823	9:09	3	0.980	A BB	33232.	80.000 UG/ML	1.23

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	6:01	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:21	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:25	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:59	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:49	1.00	0.800	1.00	80.00	80.00	0.504	0.504	1.00
8	8:11	1.00	0.875	1.00	80.00	80.00	1.299	1.299	1.00
9	15:32	1.00	0.891	1.00	80.00	80.00	1.082	1.082	1.00
10	3:32	1.00	0.906	1.00	80.00	80.00	2.046	2.046	1.00
11	2:15	1.00	0.580	1.00	80.00	80.00	1.448	1.448	1.00
12	10:55	1.00	1.168	1.00	80.00	80.00	0.233	0.233	1.00
13	3:34	1.00	0.917	1.00	80.00	80.00	1.503	1.503	1.00
14	4:07	1.00	1.057	1.00	80.00	80.00	0.959	0.959	1.00
15	12:39		0.941						
16	1:12	1.00	0.309	1.00	80.00	80.00	1.087	1.087	1.00
17	3:27	1.00	0.886	1.00	80.00	80.00	1.876	1.876	1.00
18	3:33	1.00	0.911	1.00	80.00	80.00	2.148	2.148	1.00
19	3:36	1.00	0.926	1.00	80.00	80.00	1.414	1.414	1.00
20	3:36	1.00	0.926	1.00	80.00	80.00	1.603	1.603	1.00
21	3:48	1.00	0.974	1.00	80.00	80.00	1.659	1.659	1.00
22	3:55	1.00	1.006	1.00	80.00	80.00	1.697	1.697	1.00
23	4:13	1.00	1.083	1.00	80.00	80.00	0.913	0.913	1.00
24	4:08	1.00	1.063	1.00	80.00	80.00	1.573	1.573	1.00
25	4:27	1.00	1.143	1.00	80.00	80.00	1.209	1.209	1.00
26	4:26	1.00	1.140	1.00	80.00	80.00	2.582	2.582	1.00
27	4:44	1.00	1.217	1.00	80.00	80.00	1.287	1.287	1.00
28	4:40	1.00	1.200	1.00	80.00	80.00	0.940	0.940	1.00
29	4:39	1.00	1.194	1.00	80.00	80.00	0.720	0.720	1.00
30	4:51	1.00	0.806	1.00	80.00	80.00	0.486	0.486	1.00
31	5:19	1.00	0.884	1.00	80.00	80.00	0.778	0.778	1.00
32	5:24	1.00	0.896	1.00	80.00	80.00	0.232	0.232	1.00
33	5:39	1.00	0.939	1.00	80.00	80.00	0.400	0.400	1.00
34	5:48	1.00	0.965	1.00	80.00	80.00	0.538	0.538	1.00
35	5:52	1.00	0.974	1.00	80.00	80.00	0.332	0.332	1.00
36	6:10	1.00	1.024	1.00	80.00	80.00	0.270	0.270	1.00
37	5:57	1.00	0.989	1.00	80.00	80.00	0.350	0.350	1.00
38	6:04	1.00	1.007	1.00	80.00	80.00	1.102	1.102	1.00
39	6:16	1.00	1.043	1.00	80.00	80.00	0.480	0.480	1.00
40	6:22	1.00	1.057	1.00	80.00	80.00	0.177	0.177	1.00
41	7:20	1.00	1.220	1.00	80.00	80.00	0.365	0.365	1.00
42	7:22	1.00	1.226	1.00	80.00	80.00	0.757	0.757	1.00
43	7:42	1.00	0.824	1.00	80.00	80.00	0.428	0.428	1.00
44	8:00	1.00	0.856	1.00	80.00	80.00	0.435	0.435	1.00
45	8:03	1.00	0.862	1.00	80.00	80.00	0.443	0.443	1.00
46	8:17	1.00	0.887	1.00	80.00	80.00	1.283	1.283	1.00
47	8:37	1.00	0.921	1.00	80.00	80.00	0.523	0.523	1.00
48	9:06	1.00	0.974	1.00	80.00	80.00	1.426	1.426	1.00
49	9:03	1.00	0.969	1.00	80.00	80.00	2.047	2.047	1.00
50	9:09	1.00	0.980	1.00	80.00	80.00	0.351	0.351	1.00



Data: ST16980827.TI

08/27/98 9:22:00

Sample: SSTD080 80UG/ML CALI III 082198A

Conds.: INST. ID: F16

Formula: IUL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name			
51	C545 3-NITROANILINE	138 S3# 75		99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC		83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP		51-28-5
54	C565 DIBENZOFURAN	168 S3# 90		132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP		100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105		121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130		84-66-2
58	C590 FLUORENE	166 S3#135		86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140	7005-72-3	
60	C595 4-NITROANILINE	138 S3#150		100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10		534-52-1
62	C615 N-NITROSODIPHENYLAMINE	169 S4# 15CC		87-30-6
63	C620 AZOBENZENE	77 S4# 20	103-33-3	
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30		101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40		118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50		87-86-5
67	C640 PHENANTHRENE	178 S4# 65		85-01-8
68	C645 ANTHRACENE	178 S4# 70		120-12-7
69	C647 CARBAZOLE	167 S4# 80		86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85		84-74-2
71	C655 FLUORANTHENE	202 S4#110CC		206-44-0
72	C710 BENZIDINE	184 S5# 10		92-81-5
73	C715 PYRENE	202 S5# 15		129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40		85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50		56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55		91-94-1
77	C740 CHRYSENE	228 S5# 60		218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65		117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC		117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15		205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25		207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35		50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55		193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60		53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65		191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	XTot
51	138	848	9:26	3	1.010	A BB	36135.	80.000 UG/ML	1.23
52	153	846	9:25	3	1.007	A BB	119458.	80.000 UG/ML	1.23
53	184	867	9:39	3	1.032	A BB	19158.	80.000 UG/ML	1.23
54	168	878	9:46	3	1.045	A BB	166910.	80.000 UG/ML	1.23
55	109	897	9:59	3	1.068	A BB	17386.	80.000 UG/ML	1.23
56	165	891	9:55	3	1.061	A BB	41432.	80.000 UG/ML	1.23
57	149	946	10:31	3	1.126	A BB	138464.	80.000 UG/ML	1.23
58	166	938	10:26	3	1.117	A BV	131258.	80.000 UG/ML	1.23

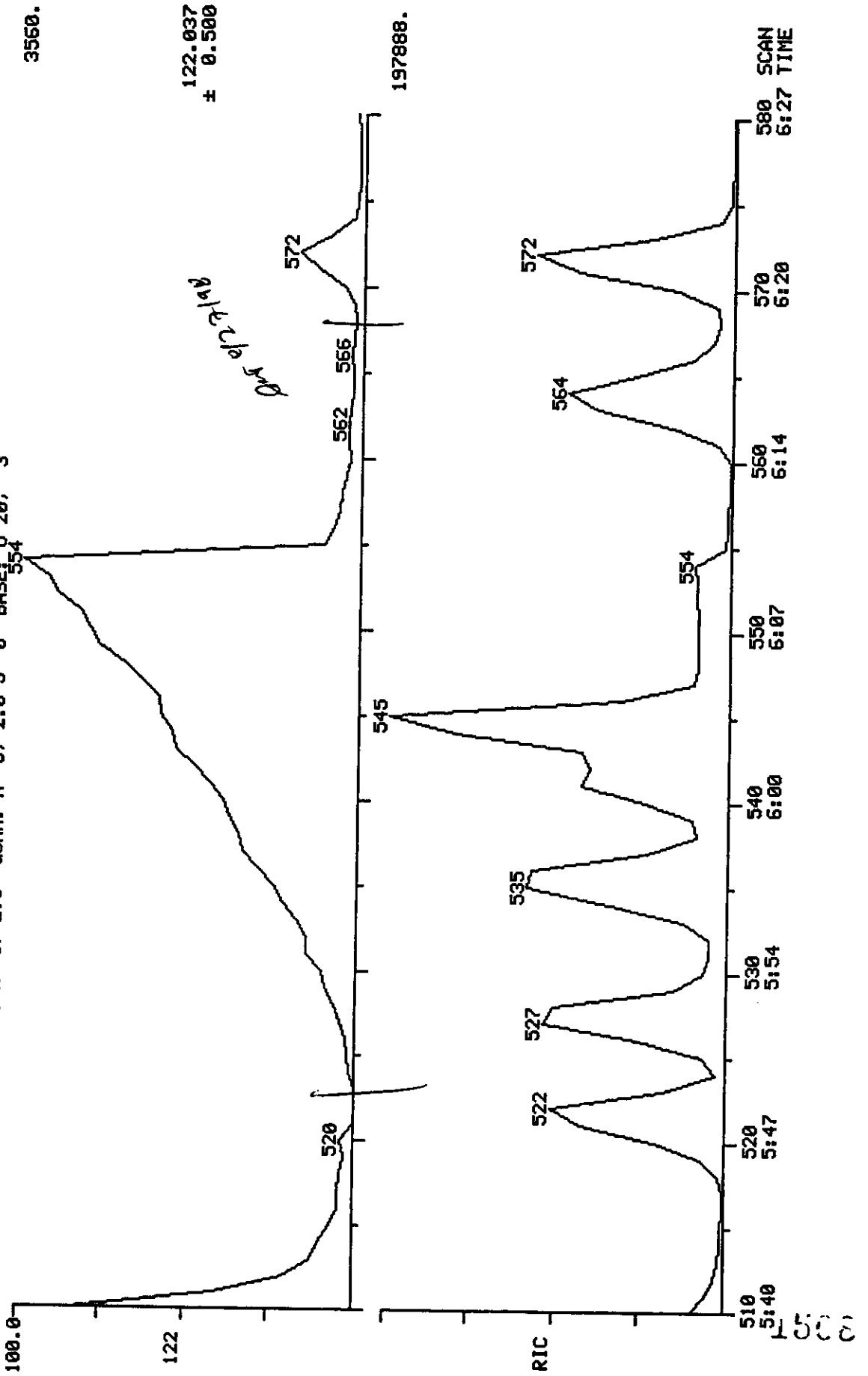
No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	XTot
59	204	949	10:33	3	1.130	A BB	61220.	80.000 UG/ML	1.23
60	138	957	10:39	3	1.139	A BB	28966.	80.000 UG/ML	1.23
61	198	962	10:42	4	0.875	A BB	23256.	80.000 UG/ML	1.23
62	169	974	10:50	4	0.886	A BV	86506.	80.000 UG/ML	1.23
63	77	976	10:51	4	0.888	A BB	194334.	80.000 UG/ML	1.23
64	248	1033	11:29	4	0.940	A BB	37192.	80.000 UG/ML	1.23
65	284	1032	11:29	4	0.939	A BB	43018.	80.000 UG/ML	1.23
66	266	1074	11:57	4	0.977	A BB	27210.	80.000 UG/ML	1.23
67	178	1104	12:17	4	1.005	A BV	175815.	80.000 UG/ML	1.23
68	178	1112	12:22	4	1.012	A VB	169868.	80.000 UG/ML	1.23
69	167	1150	12:47	4	1.046	A BB	123401.	80.000 UG/ML	1.23
70	149	1239	13:47	4	1.127	A BB	213776.	80.000 UG/ML	1.23
71	202	1312	14:36	4	1.194	A BB	152339.	80.000 UG/ML	1.23
72	184	1353	15:03	5	0.864	A BB	7938.	80.000 UG/ML	1.23
73	202	1348	15:00	5	0.861	A BB	150637.	80.000 UG/ML	1.23
74	149	1500	16:41	5	0.958	A BB	74136.	80.000 UG/ML	1.23
75	228	1565	17:24	5	0.999	A BV	123435.	80.000 UG/ML	1.23
76	252	1575	17:31	5	1.006	A BB	39292.	80.000 UG/ML	1.23
77	228	1571	17:28	5	1.003	A VB	113108.	80.000 UG/ML	1.23
78	149	1616	17:58	5	1.032	A BB	105344.	80.000 UG/ML	1.23
79	149	1727	19:13	6	0.961	A BB	175103.	80.000 UG/ML	1.23
80	252	1742	19:23	6	0.969	A BV	128697.	80.000 UG/ML	1.23
81	252	1747	19:26	6	0.972	A VB	111928.	80.000 UG/ML	1.23
82	252	1789	19:54	6	0.996	A BV	108866.	80.000 UG/ML	1.23
83	276	1945	21:38	6	1.082	A BV	121168.	80.000 UG/ML	1.23
84	278	1952	21:43	6	1.086	A BV	120479.	80.000 UG/ML	1.23
85	276	1977	21:59	6	1.100	A BB	126303.	80.000 UG/ML	1.23

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:26	1.00	1.010	1.00	80.00	80.00	0.382	0.382	1.00
52	9:25	1.00	1.007	1.00	80.00	80.00	1.261	1.261	1.00
53	9:39	1.00	1.032	1.00	80.00	80.00	0.202	0.202	1.00
54	9:46	1.00	1.045	1.00	80.00	80.00	1.762	1.762	1.00
55	9:59	1.00	1.068	1.00	80.00	80.00	0.184	0.184	1.00
56	9:55	1.00	1.061	1.00	80.00	80.00	0.437	0.437	1.00
57	10:31	1.00	1.126	1.00	80.00	80.00	1.462	1.462	1.00
58	10:26	1.00	1.117	1.00	80.00	80.00	1.386	1.386	1.00
59	10:33	1.00	1.130	1.00	80.00	80.00	0.646	0.646	1.00
60	10:39	1.00	1.139	1.00	80.00	80.00	0.306	0.306	1.00
61	10:42	1.00	0.875	1.00	80.00	80.00	0.145	0.145	1.00
62	10:50	1.00	0.886	1.00	80.00	80.00	0.539	0.539	1.00
63	10:51	1.00	0.888	1.00	80.00	80.00	1.212	1.212	1.00
64	11:29	1.00	0.940	1.00	80.00	80.00	0.232	0.232	1.00
65	11:29	1.00	0.939	1.00	80.00	80.00	0.268	0.268	1.00
66	11:57	1.00	0.977	1.00	80.00	80.00	0.170	0.170	1.00
67	12:17	1.00	1.005	1.00	80.00	80.00	1.096	1.096	1.00
68	12:22	1.00	1.012	1.00	80.00	80.00	1.059	1.059	1.00
69	12:47	1.00	1.046	1.00	80.00	80.00	0.769	0.769	1.00
70	13:47	1.00	1.127	1.00	80.00	80.00	1.333	1.333	1.00
71	14:36	1.00	1.194	1.00	80.00	80.00	0.950	0.950	1.00
72	15:03	1.00	0.864	1.00	80.00	80.00	0.077	0.077	1.00
73	15:00	1.00	0.861	1.00	80.00	80.00	1.468	1.468	1.00
74	16:41	1.00	0.958	1.00	80.00	80.00	0.722	0.722	1.00
75	17:24	1.00	0.999	1.00	80.00	80.00	1.203	1.203	1.00
76	17:31	1.00	1.006	1.00	80.00	80.00	0.383	0.383	1.00
77	17:28	1.00	1.003	1.00	80.00	80.00	1.102	1.102	1.00

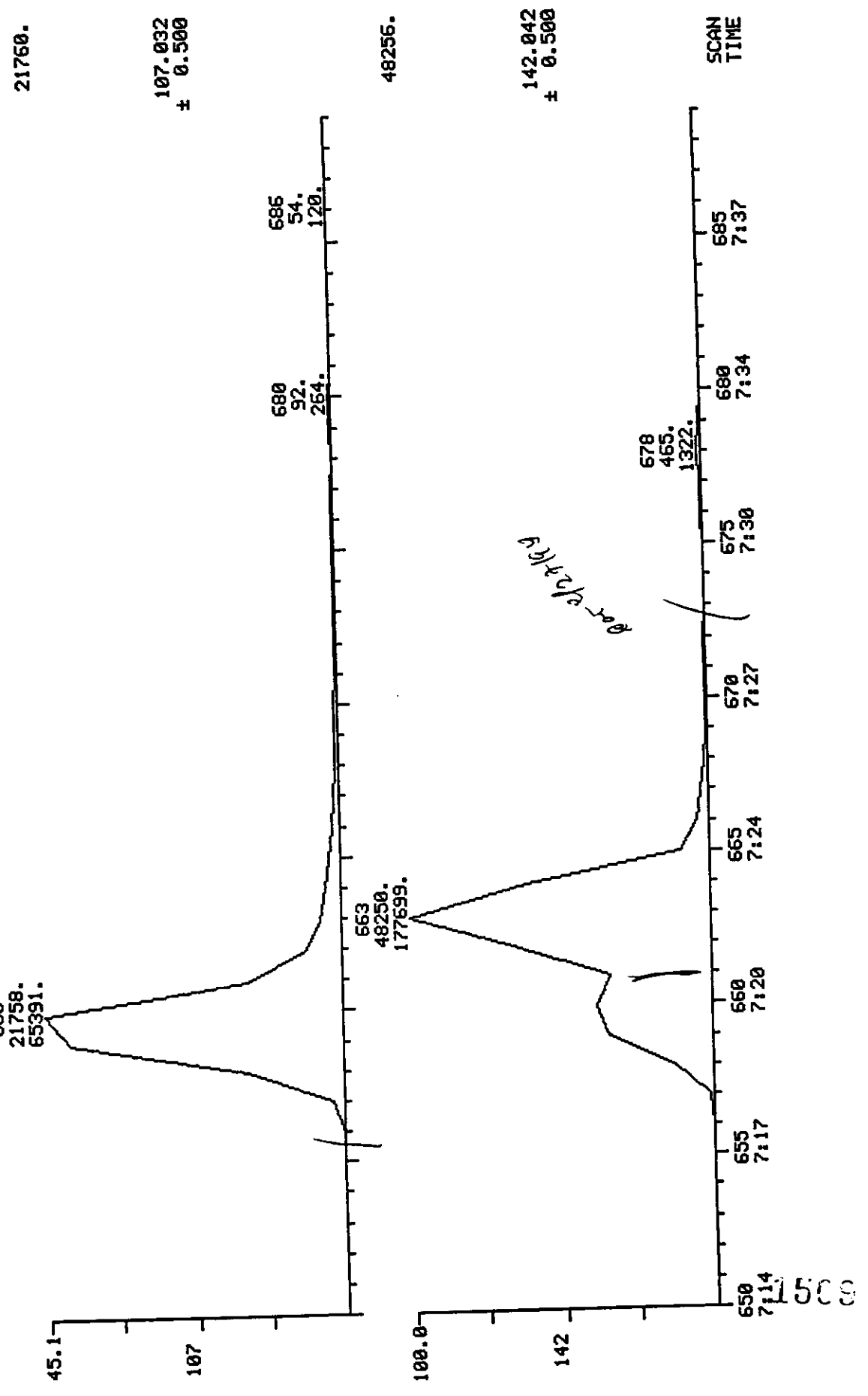
No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:58	1.00	1.032	1.00	80.00	80.00	1.027	1.027	1.00
79	19:13	1.00	0.961	1.00	80.00	80.00	1.977	1.977	1.00
80	19:23	1.00	0.969	1.00	80.00	80.00	1.453	1.453	1.00
81	19:26	1.00	0.972	1.00	80.00	80.00	1.264	1.264	1.00
82	19:54	1.00	0.996	1.00	80.00	80.00	1.229	1.229	1.00
83	21:38	1.00	1.082	1.00	80.00	80.00	1.368	1.368	1.00
84	21:43	1.00	1.086	1.00	80.00	80.00	1.360	1.360	1.00
85	21:59	1.00	1.100	1.00	80.00	80.00	1.426	1.426	1.00

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPHY**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND CODED**

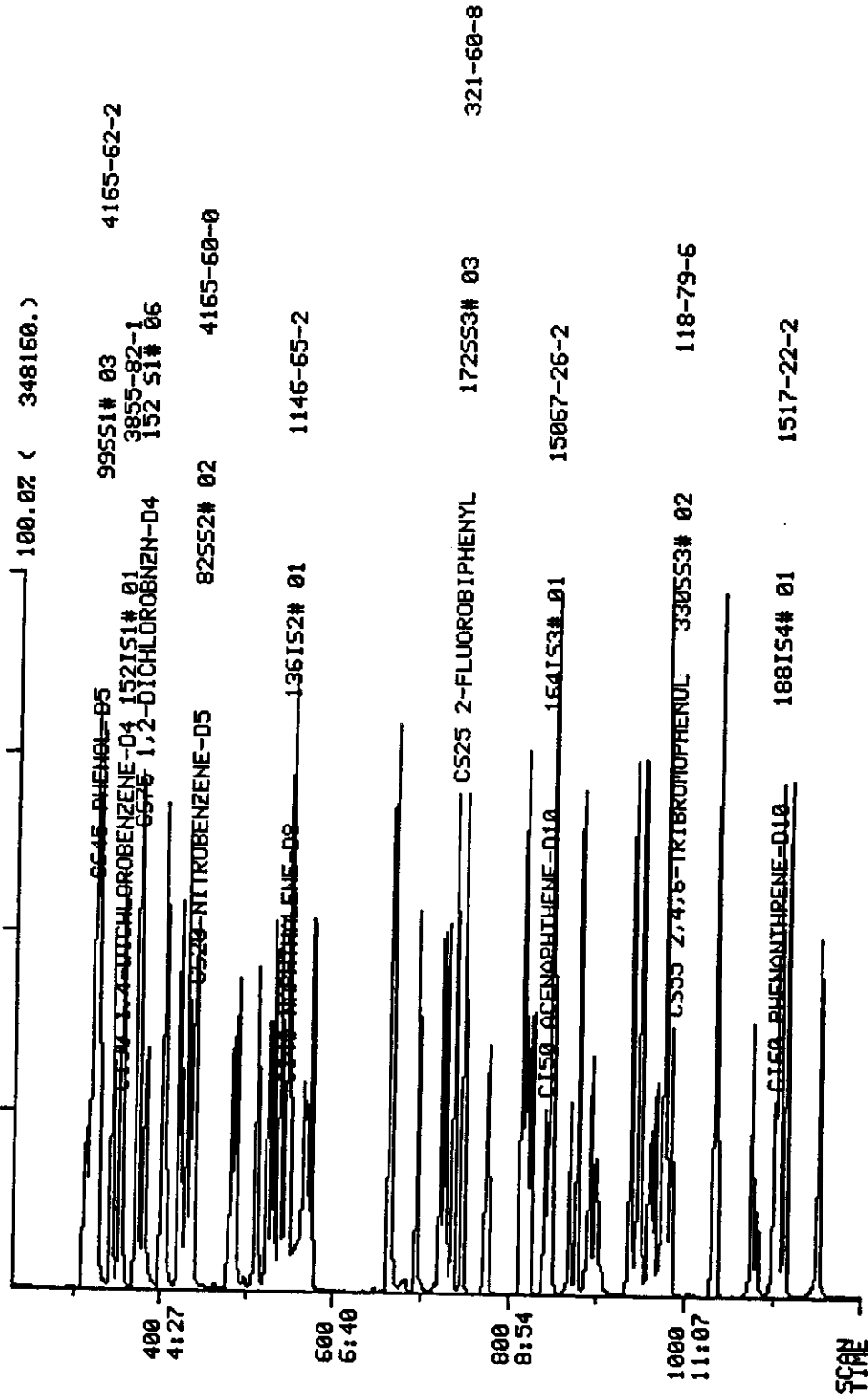
RIC+MASS CHROMATOGRAM  
 08/27/98 9:22:00 DATA: ST16980827 #1  
 CALI: ST16980827 #3  
 SAMPLE: SST080 80UG/ML CALI III 062198A  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3



MASS CHROMATOGRAMS  
 08/27/98 9:22:00 DATA: ST16980827 #1 SCANS 650 TO 689  
 CALI: ST16980827 #3  
 SAMPLE: SST080 80UG/ML CALI III 082198A  
 CONDS.: INST. ID: F16  
 RANGE: G 1,2158 LABEL: N 2, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3

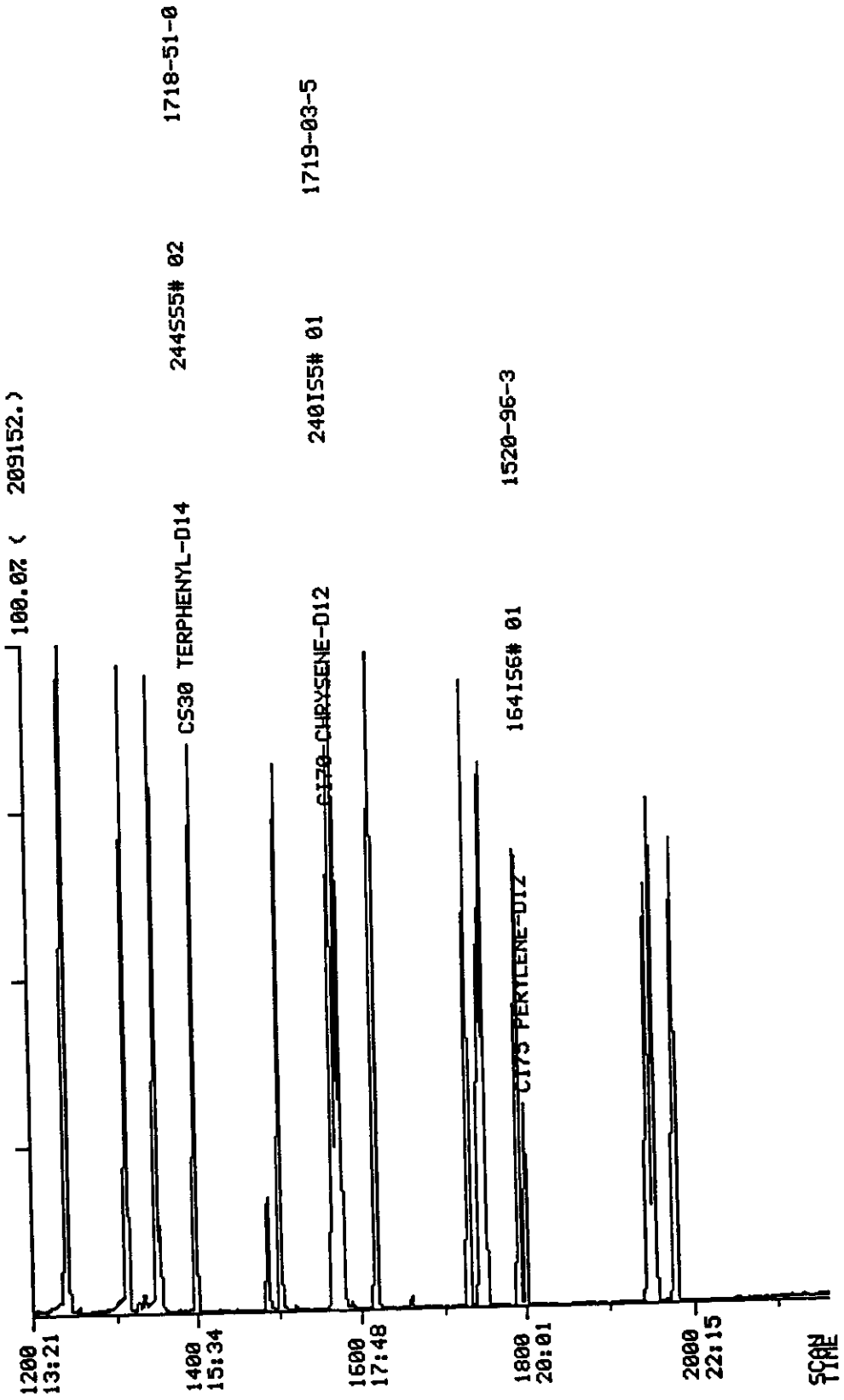


DATA FROM FILE: ST169808270 SCANS 230 TO 1200 ACQUIRED: 08/27/98 11:20:00  
 CALI: ST169808270 #3  
 SAMPLE: SSTD120 120UG/ML CALI II 070698H  
 CONDS.: INST. ID: F16



1510

DATA FROM FILE: ST16980827D SCANS 1200 TO 2158 ACQUIRED: 08/27/98 11:20:00  
CALI: ST16980827D #3  
SAMPLE: SSTD120 120UG/ML CALI II 070698H  
CONDS.: INST. ID: F16



Data: ST16980827D.TI

08/27/98 11:20:00

Sample: SSTD120 120UG/ML CALI II 070698H

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	C170 CHRYSENE-D12	240IS5# 01	1719-03-5
6	C175 PERYLENE-D12	164IS6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBENZENE-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANILINE	65 S3# 45	88-74-4



No	Name		
48	C535 DIMETHYLPHTHALATE	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROTOLUENE	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	XTot
1	152	351	3:54	1	1.000	A BB	28868.	40.000 UG/ML	0.42
2	136	542	6:02	2	1.000	A BB	109146.	40.000 UG/ML	0.42
3	164	841	9:21	3	1.000	A BB	62260.	40.000 UG/ML	0.42
4	188	1100	12:14	4	1.000	A BB	102618.	40.000 UG/ML	0.42
5	240	1567	17:26	5	1.000	A BB	60424.	40.000 UG/ML	0.42
6	264	1798	20:00	6	1.000	A BB	49906.	40.000 UG/ML	0.42
7	82	435	4:50	2	0.803	A BB	148159.	120.000 UG/ML	1.25
8	172	736	8:11	3	0.875	A BB	218708.	120.000 UG/ML	1.25
9	244	1397	15:32	5	0.892	A VB	173968.	120.000 UG/ML	1.25
10	99	319	3:33	1	0.909	A BB	172744.	120.000 UG/ML	1.25
11	112	204	2:16	1	0.581	A BB	119161.	120.000 UG/ML	1.25
12	330	983	10:56	3	1.169	A BB	42668.	120.000 UG/ML	1.25
13	132	323	3:36	1	0.920	A BB	124435.	120.000 UG/ML	1.25
14	152	371	4:08	1	1.057	A BB	78148.	120.000 UG/ML	1.25
15	NOT FOUND								
16	74	109	1:13	1	0.311	A VB	87333.	120.000 UG/ML	1.25
17	93	312	3:28	1	0.889	A BV	154018.	120.000 UG/ML	1.25
18	94	321	3:34	1	0.915	A VB	165127.	120.000 UG/ML	1.25
19	93	326	3:38	1	0.929	A VB	111846.	120.000 UG/ML	1.25
20	128	325	3:37	1	0.926	A BB	125615.	120.000 UG/ML	1.25
21	146	342	3:48	1	0.974	A BV	133856.	120.000 UG/ML	1.25
22	146	353	3:56	1	1.006	A VB	134148.	120.000 UG/ML	1.25
23	108	381	4:14	1	1.085	A BB	74988.	120.000 UG/ML	1.25
24	146	373	4:09	1	1.063	A BB	125957.	120.000 UG/ML	1.25
25	108	402	4:28	1	1.145	A BB	103881.	120.000 UG/ML	1.25
26	45	400	4:27	1	1.140	A BB	216653.	120.000 UG/ML	1.25
27	108	429	4:46	1	1.222	A BB	113933.	120.000 UG/ML	1.25
28	70	423	4:42	1	1.205	A BV	79097.	120.000 UG/ML	1.25
29	117	419	4:40	1	1.194	A BB	61106.	120.000 UG/ML	1.25
30	77	438	4:52	2	0.808	A VB	137055.	120.000 UG/ML	1.25
31	82	481	5:21	2	0.887	A BB	237871.	120.000 UG/ML	1.25
32	139	486	5:24	2	0.897	A BB	71784.	120.000 UG/ML	1.25
33	107	510	5:40	2	0.941	A BB	114342.	120.000 UG/ML	1.25
34	93	523	5:49	2	0.965	A BB	164456.	120.000 UG/ML	1.25
35	162	529	5:53	2	0.976	A BB	101851.	120.000 UG/ML	1.25
36	122	564	6:16	2	1.041	qedt(2)	91808.	120.000 UG/ML	1.25
37	180	537	5:58	2	0.991	A BB	105480.	120.000 UG/ML	1.25
38	128	546	6:04	2	1.007	A BB	314427.	120.000 UG/ML	1.25
39	127	567	6:18	2	1.046	A BB	146940.	120.000 UG/ML	1.25
40	225	572	6:22	2	1.055	A BB	54706.	120.000 UG/ML	1.25
41	107	661	7:21	2	1.220	A BB	113771.	120.000 UG/ML	1.25
42	142	664	7:23	2	1.225	qedt(2)	219628.	120.000 UG/ML	1.25
43	237	693	7:42	3	0.824	A BB	70772.	120.000 UG/ML	1.25
44	196	720	8:01	3	0.856	A BV	77913.	120.000 UG/ML	1.25
45	196	726	8:05	3	0.863	A VB	74944.	120.000 UG/ML	1.25
46	162	747	8:19	3	0.888	A BB	210616.	120.000 UG/ML	1.25
47	65	775	8:37	3	0.922	A BB	90989.	120.000 UG/ML	1.25
48	163	820	9:07	3	0.975	A BB	242196.	120.000 UG/ML	1.25
49	152	815	9:04	3	0.969	A BB	318836.	120.000 UG/ML	1.25
50	165	825	9:11	3	0.981	A BB	59941.	120.000 UG/ML	1.25

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	6:01	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:21	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:25	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:59	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:49	1.00	0.800	1.00	120.00	120.00	0.452	0.452	1.00
8	8:11	1.00	0.875	1.00	120.00	120.00	1.171	1.171	1.00
9	15:32	1.00	0.891	1.00	120.00	120.00	0.960	0.960	1.00
10	3:32	1.01	0.906	1.00	120.00	120.00	1.995	1.995	1.00
11	2:15	1.00	0.580	1.00	120.00	120.00	1.376	1.376	1.00
12	10:55	1.00	1.168	1.00	120.00	120.00	0.228	0.228	1.00
13	3:34	1.01	0.917	1.00	120.00	120.00	1.437	1.437	1.00
14	4:07	1.00	1.057	1.00	120.00	120.00	0.902	0.902	1.00
15	12:39		0.941						
16	1:12	1.01	0.309	1.01	120.00	120.00	1.008	1.008	1.00
17	3:27	1.01	0.886	1.00	120.00	120.00	1.778	1.778	1.00
18	3:33	1.01	0.911	1.00	120.00	120.00	1.907	1.907	1.00
19	3:36	1.01	0.926	1.00	120.00	120.00	1.291	1.291	1.00
20	3:36	1.00	0.926	1.00	120.00	120.00	1.450	1.450	1.00
21	3:48	1.00	0.974	1.00	120.00	120.00	1.546	1.546	1.00
22	3:55	1.00	1.006	1.00	120.00	120.00	1.549	1.549	1.00
23	4:13	1.01	1.083	1.00	120.00	120.00	0.866	0.866	1.00
24	4:08	1.00	1.063	1.00	120.00	120.00	1.454	1.454	1.00
25	4:27	1.01	1.143	1.00	120.00	120.00	1.199	1.199	1.00
26	4:26	1.00	1.140	1.00	120.00	120.00	2.502	2.502	1.00
27	4:44	1.01	1.217	1.00	120.00	120.00	1.316	1.316	1.00
28	4:40	1.01	1.200	1.00	120.00	120.00	0.913	0.913	1.00
29	4:39	1.00	1.194	1.00	120.00	120.00	0.706	0.706	1.00
30	4:51	1.00	0.806	1.00	120.00	120.00	0.419	0.419	1.00
31	5:19	1.01	0.884	1.00	120.00	120.00	0.726	0.726	1.00
32	5:24	1.00	0.896	1.00	120.00	120.00	0.219	0.219	1.00
33	5:39	1.00	0.939	1.00	120.00	120.00	0.349	0.349	1.00
34	5:48	1.00	0.965	1.00	120.00	120.00	0.502	0.502	1.00
35	5:52	1.00	0.974	1.00	120.00	120.00	0.311	0.311	1.00
36	6:10	1.02	1.024	1.02	120.00	120.00	0.280	0.280	1.00
37	5:57	1.00	0.989	1.00	120.00	120.00	0.322	0.322	1.00
38	6:04	1.00	1.007	1.00	120.00	120.00	0.960	0.960	1.00
39	6:16	1.01	1.043	1.00	120.00	120.00	0.449	0.449	1.00
40	6:22	1.00	1.057	1.00	120.00	120.00	0.167	0.167	1.00
41	7:20	1.00	1.220	1.00	120.00	120.00	0.347	0.347	1.00
42	7:22	1.00	1.226	1.00	120.00	120.00	0.671	0.671	1.00
43	7:42	1.00	0.824	1.00	120.00	120.00	0.379	0.379	1.00
44	8:00	1.00	0.856	1.00	120.00	120.00	0.417	0.417	1.00
45	8:03	1.00	0.862	1.00	120.00	120.00	0.401	0.401	1.00
46	8:17	1.00	0.887	1.00	120.00	120.00	1.128	1.128	1.00
47	8:37	1.00	0.921	1.00	120.00	120.00	0.487	0.487	1.00
48	9:06	1.00	0.974	1.00	120.00	120.00	1.297	1.297	1.00
49	9:03	1.00	0.969	1.00	120.00	120.00	1.707	1.707	1.00
50	9:09	1.00	0.980	1.00	120.00	120.00	0.321	0.321	1.00

Quantitation Report File: ST16980827D

Data: ST16980827D.TI

08/27/98 11:20:00

Sample: SSTD120 120UG/ML CALI II 070698H

Conds.: INST. ID: F16

Formula: IUL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name			
51	C545 3-NITROANILINE	138 S3# 75		99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC		83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP		51-28-5
54	C565 DIBENZOFURAN	168 S3# 90		132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP		100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105		121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130		84-66-2
58	C590 FLUORENE	166 S3#135		86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140	7005-72-3	
60	C595 4-NITROANALINE	138 S3#150		100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10		534-52-1
62	C615 N-NITROSOIPHENYLAMINE	169 S4# 15CC		87-30-6
63	C620 AZOBENZENE	77 S4# 20	103-33-3	
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30		101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40		118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50		87-86-5
67	C640 PHENANTHRENE	178 S4# 65		85-01-8
68	C645 ANTHRACENE	178 S4# 70		120-12-7
69	C647 CARBAZOLE	167 S4# 80		86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85		84-74-2
71	C655 FLUORANTHENE	202 S4#110CC		206-44-0
72	C710 BENZIDINE	184 S5# 10		92-81-5
73	C715 PYRENE	202 S5# 15		129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40		85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50		56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55		91-94-1
77	C740 CHRYSENE	228 S5# 60		218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65		117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC		117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15		205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25		207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35		50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55		193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60		53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65		191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	138	850	9:27	3	1.011	A BB	70791.	120.000 UG/ML	1.25
52	153	848	9:26	3	1.008	A BB	200860.	120.000 UG/ML	1.25
53	184	870	9:41	3	1.034	A BB	37857.	120.000 UG/ML	1.25
54	168	880	9:47	3	1.046	A VB	290690.	120.000 UG/ML	1.25
55	109	899	10:00	3	1.069	A BB	31391.	120.000 UG/ML	1.25
56	165	893	9:56	3	1.062	A BB	74054.	120.000 UG/ML	1.25
57	149	948	10:33	3	1.127	A BB	247974.	120.000 UG/ML	1.25
58	166	939	10:27	3	1.117	A BV	228542.	120.000 UG/ML	1.25

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	XTot
59	204	950	10:34	3	1.130	A BB	108212.	120.000 UG/ML	1.25
60	138	961	10:41	3	1.143	A VB	49100.	120.000 UG/ML	1.25
61	198	965	10:44	4	0.877	A VB	38167.	120.000 UG/ML	1.25
62	169	976	10:51	4	0.887	A BV	163243.	120.000 UG/ML	1.25
63	77	978	10:53	4	0.889	A BB	338029.	120.000 UG/ML	1.25
64	248	1034	11:30	4	0.940	A BB	66106.	120.000 UG/ML	1.25
65	284	1033	11:29	4	0.939	A BB	73348.	120.000 UG/ML	1.25
66	266	1075	11:57	4	0.977	A BB	47725.	120.000 UG/ML	1.25
67	178	1105	12:17	4	1.005	A BV	291774.	120.000 UG/ML	1.25
68	178	1114	12:23	4	1.013	A VB	279557.	120.000 UG/ML	1.25
69	167	1151	12:48	4	1.046	A BB	217660.	120.000 UG/ML	1.25
70	149	1240	13:48	4	1.127	A BB	349945.	120.000 UG/ML	1.25
71	202	1313	14:36	4	1.194	A BB	242593.	120.000 UG/ML	1.25
72	184	1353	15:03	5	0.863	A BB	25549.	120.000 UG/ML	1.25
73	202	1349	15:00	5	0.861	A BB	234777.	120.000 UG/ML	1.25
74	149	1501	16:42	5	0.958	A BB	126873.	120.000 UG/ML	1.25
75	228	1565	17:24	5	0.999	A BV	196831.	120.000 UG/ML	1.25
76	252	1576	17:32	5	1.006	A BB	73334.	120.000 UG/ML	1.25
77	228	1572	17:29	5	1.003	A VB	173809.	120.000 UG/ML	1.25
78	149	1616	17:58	5	1.031	A BB	178906.	120.000 UG/ML	1.25
79	149	1728	19:13	6	0.961	A BB	294445.	120.000 UG/ML	1.25
80	252	1744	19:24	6	0.970	qedt	223052.	120.000 UG/ML	1.25
81	252	1748	19:27	6	0.972	qedt	175371.	120.000 UG/ML	1.25
82	252	1790	19:55	6	0.996	A BB	171744.	120.000 UG/ML	1.25
83	276	1946	21:39	6	1.082	A BV	185834.	120.000 UG/ML	1.25
84	278	1953	21:43	6	1.086	A BB	181428.	120.000 UG/ML	1.25
85	276	1978	22:00	6	1.100	A BB	192918.	120.000 UG/ML	1.25

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:26	1.00	1.010	1.00	120.00	120.00	0.379	0.379	1.00
52	9:25	1.00	1.007	1.00	120.00	120.00	1.075	1.075	1.00
53	9:39	1.00	1.032	1.00	120.00	120.00	0.203	0.203	1.00
54	9:46	1.00	1.045	1.00	120.00	120.00	1.556	1.556	1.00
55	9:59	1.00	1.068	1.00	120.00	120.00	0.168	0.168	1.00
56	9:55	1.00	1.061	1.00	120.00	120.00	0.396	0.396	1.00
57	10:31	1.00	1.126	1.00	120.00	120.00	1.328	1.328	1.00
58	10:26	1.00	1.117	1.00	120.00	120.00	1.224	1.224	1.00
59	10:33	1.00	1.130	1.00	120.00	120.00	0.579	0.579	1.00
60	10:39	1.00	1.139	1.00	120.00	120.00	0.263	0.263	1.00
61	10:42	1.00	0.875	1.00	120.00	120.00	0.124	0.124	1.00
62	10:50	1.00	0.886	1.00	120.00	120.00	0.530	0.530	1.00
63	10:51	1.00	0.888	1.00	120.00	120.00	1.098	1.098	1.00
64	11:29	1.00	0.940	1.00	120.00	120.00	0.215	0.215	1.00
65	11:29	1.00	0.939	1.00	120.00	120.00	0.238	0.238	1.00
66	11:57	1.00	0.977	1.00	120.00	120.00	0.155	0.155	1.00
67	12:17	1.00	1.005	1.00	120.00	120.00	0.948	0.948	1.00
68	12:22	1.00	1.012	1.00	120.00	120.00	0.908	0.908	1.00
69	12:47	1.00	1.046	1.00	120.00	120.00	0.707	0.707	1.00
70	13:47	1.00	1.127	1.00	120.00	120.00	1.137	1.137	1.00
71	14:36	1.00	1.194	1.00	120.00	120.00	0.788	0.788	1.00
72	15:03	1.00	0.864	1.00	120.00	120.00	0.141	0.141	1.00
73	15:00	1.00	0.861	1.00	120.00	120.00	1.295	1.295	1.00
74	16:41	1.00	0.958	1.00	120.00	120.00	0.700	0.700	1.00
75	17:24	1.00	0.999	1.00	120.00	120.00	1.086	1.086	1.00
76	17:31	1.00	1.006	1.00	120.00	120.00	0.405	0.405	1.00
77	17:28	1.00	1.003	1.00	120.00	120.00	0.959	0.959	1.00

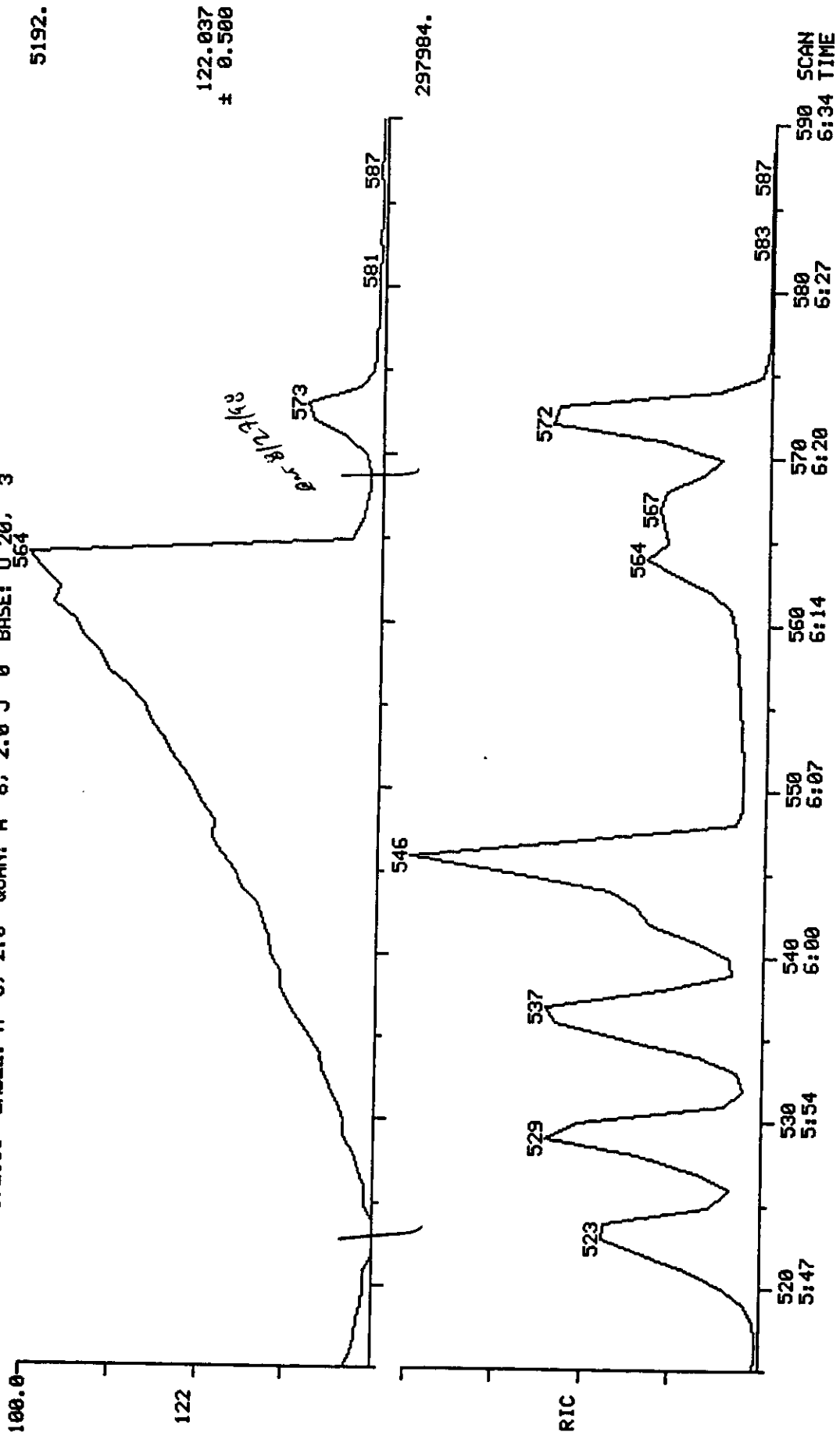
No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:58	1.00	1.032	1.00	120.00	120.00	0.987	0.987	1.00
79	19:13	1.00	0.961	1.00	120.00	120.00	1.967	1.967	1.00
80	19:23	1.00	0.969	1.00	120.00	120.00	1.490	1.490	1.00
81	19:26	1.00	0.972	1.00	120.00	120.00	1.171	1.171	1.00
82	19:54	1.00	0.996	1.00	120.00	120.00	1.147	1.147	1.00
83	21:38	1.00	1.082	1.00	120.00	120.00	1.241	1.241	1.00
84	21:43	1.00	1.086	1.00	120.00	120.00	1.212	1.212	1.00
85	21:59	1.00	1.100	1.00	120.00	120.00	1.289	1.289	1.00

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPHY**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**IDENTIFIED, DATED, AND CODED**

RIC+MASS CHROMATOGRAM

08/27/98 11:20:00 DATA: ST169808270 #1  
CALI: ST169808270 #3  
SAMPLE: SSTD120 120UG/ML CALI II 070698H  
COND.S.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U.20, 3

SCANS 515 TO 590



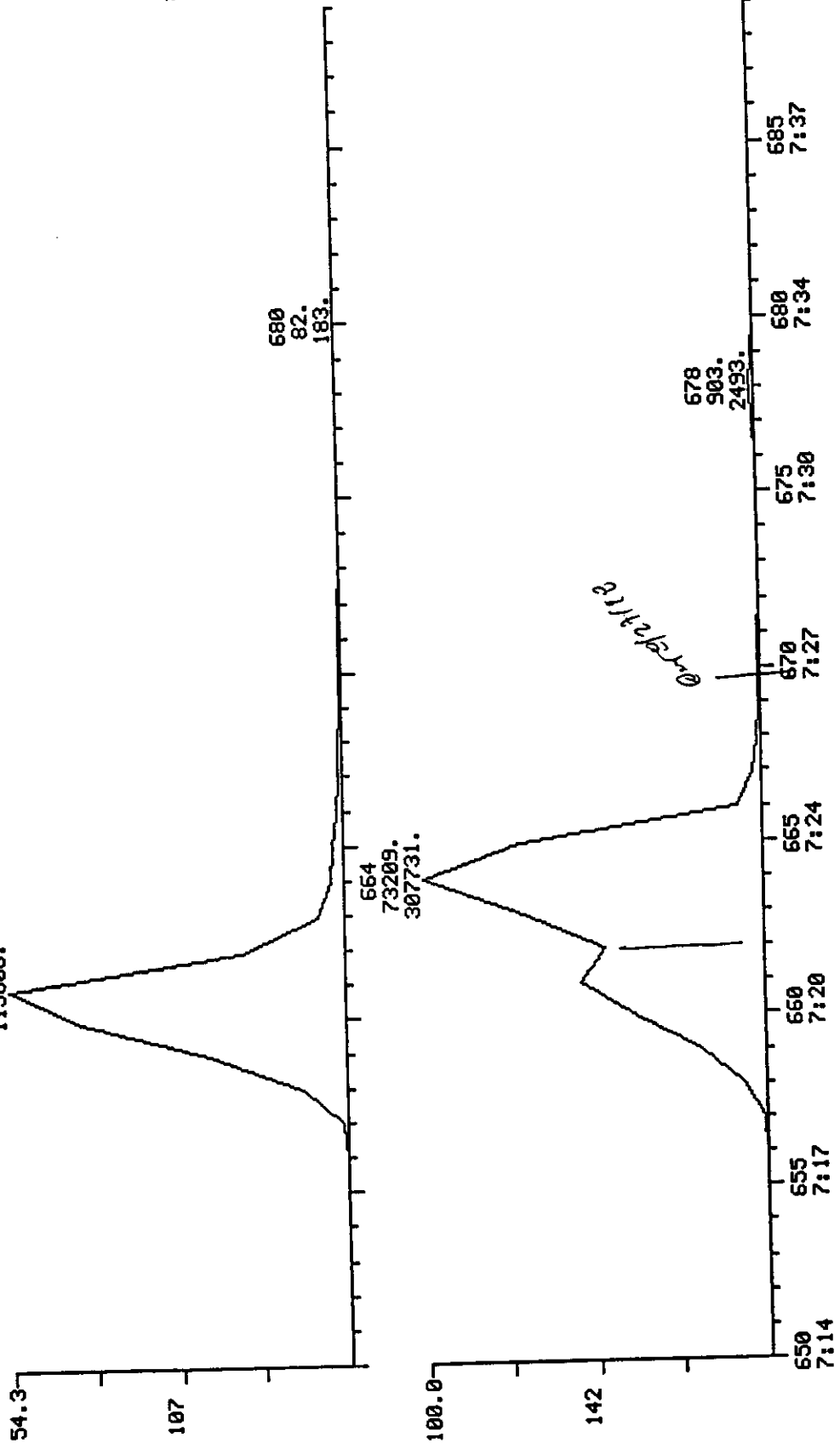
SCANS 650 TO 689

DATA: ST169808270 #1  
CALI: ST169808270 #3

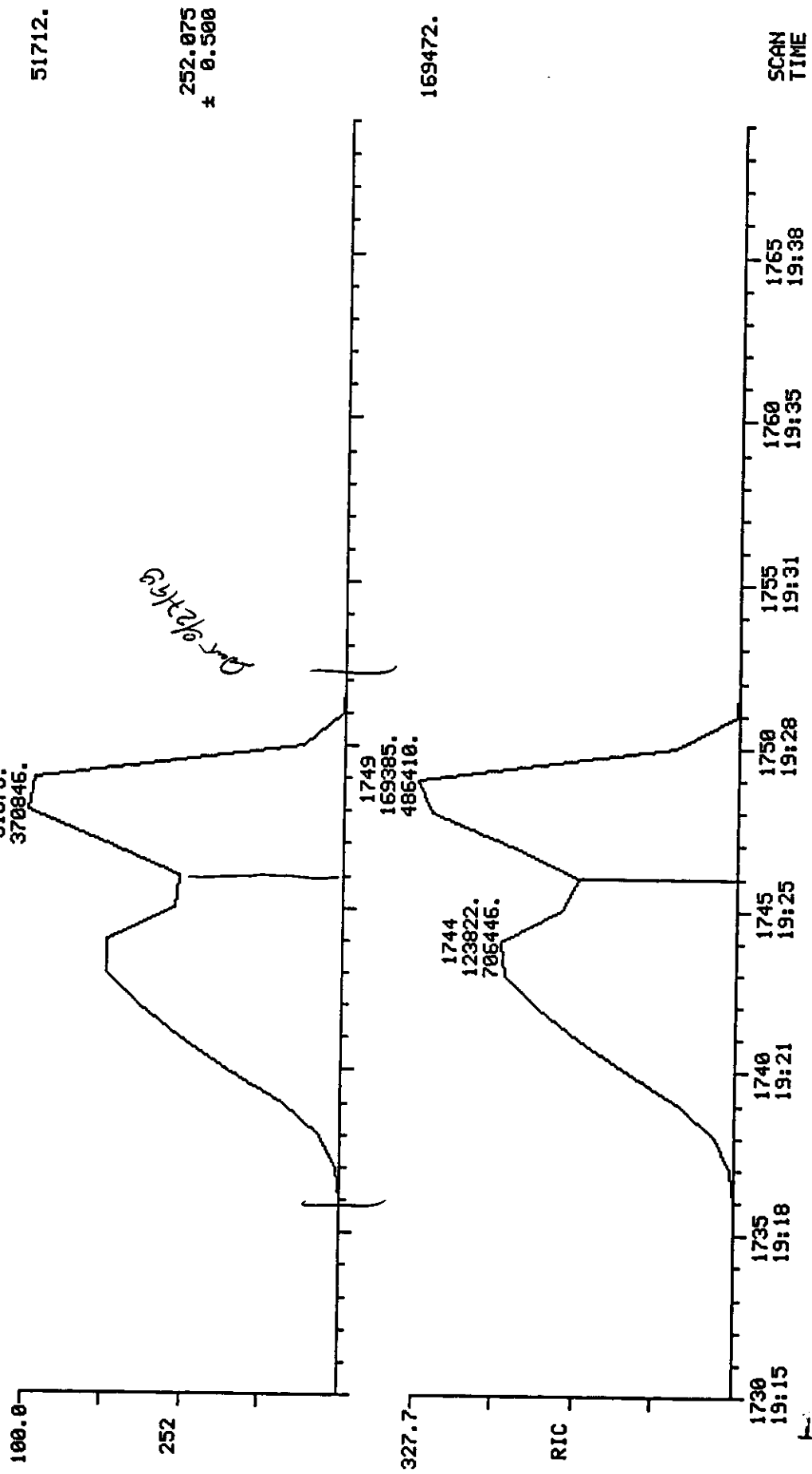
MASS CHROMATOGRAMS

08/27/98 11:20:00  
SAMPLE: SSTD120 120UG/ML CALI II 070698H  
CONDS.: INST. ID: F16  
RANGE: G 1,2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

661 39742. 113806. 54.3 107 107.032 ± 0.500 680 82. 183. 664 73209. 307731. 73216. 678 675 670 685 680 678 903. 2493. 142.042 ± 0.500 650 7:14 655 7:17 660 7:20 665 7:24 670 7:27 675 7:30 680 7:34 685 7:37



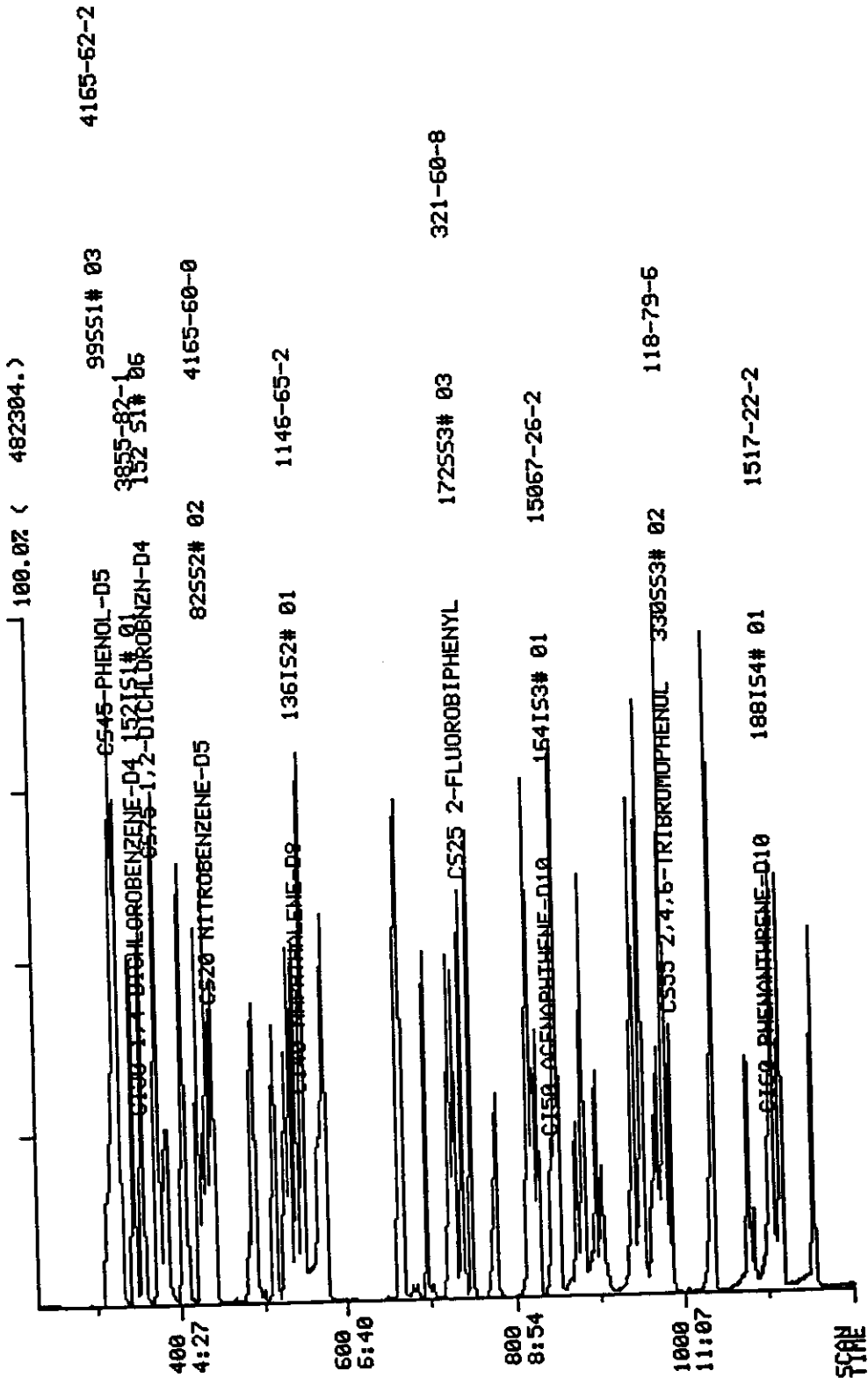
RIC+MASS CHROMATOGRAM  
 08/27/98 11:20:00 DATA: ST169808270 #1 SCANS 1730 TO 1769  
 CALI: ST169808270 #3  
 SAMPLE: SSTD120 120UG/ML CALI II 070698H  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3



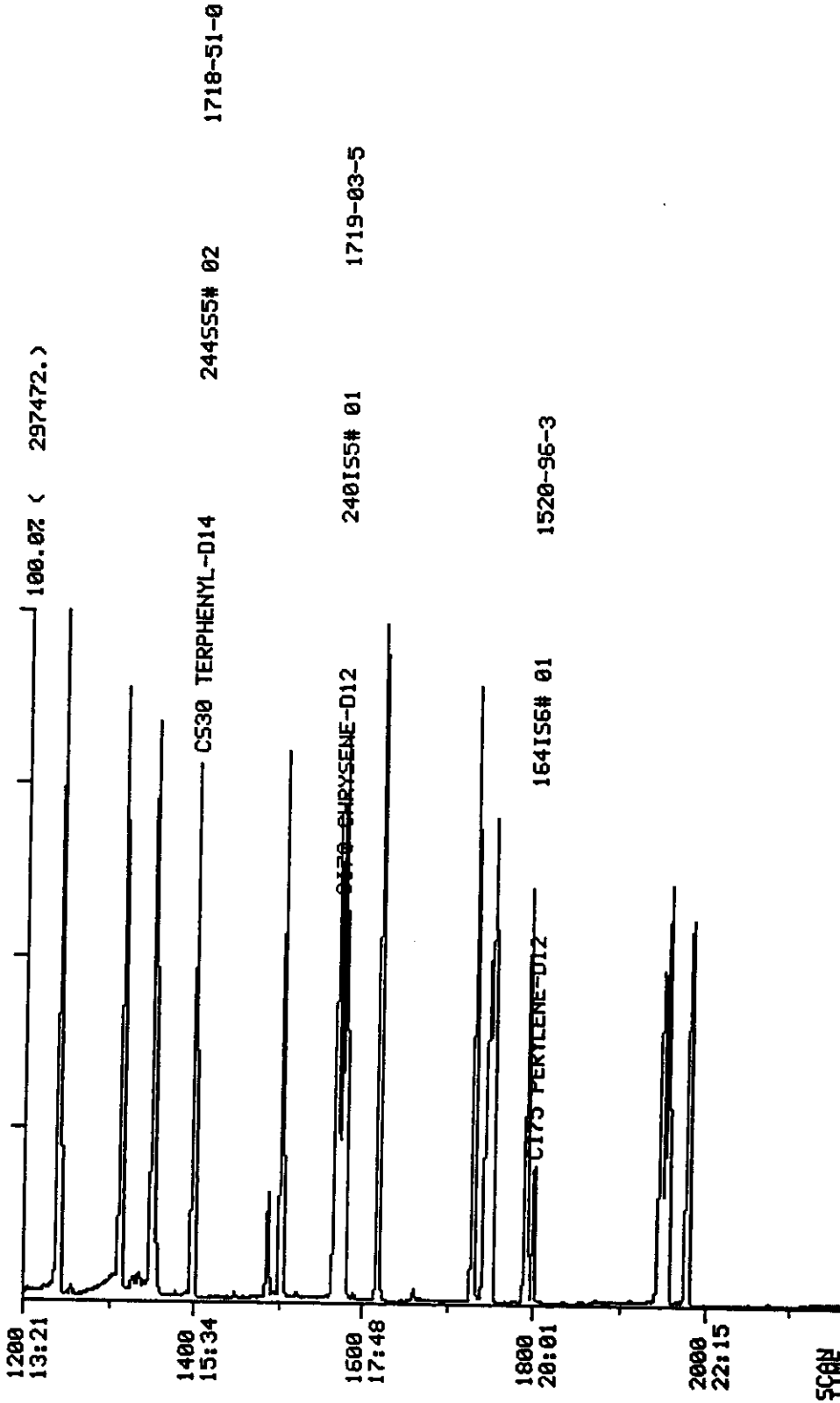
1520



DATA FROM FILE: ST16980827E SCANS 230 TO 1200 ACQUIRED: 08/27/98 11:50:00  
 SAMPLE: SSTD160 160UG/ML CALI I 0706981 CALI: ST16980827E #3  
 CONDS.: INST. ID: F16



DATA FROM FILE: ST16980827E SCANS 1200 TO 2158 ACQUIRED: 08/27/98 11:50:00  
CALI: ST16980827E #3  
SAMPLE: SSTD160 160UG/ML CALI I 0706981  
CONDS.: INST. ID: F16



SCALE

1522

Data: ST16980827E.TI

08/27/98 11:50:00

Sample: SSTD160 160UG/ML CALI I 070698I

Conds.: INST. ID: F16

Formula: IUL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
1	CI30 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	CI40 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	CI50 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	CI60 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	CI70 CHRYSENE-D12	240IS5# 01	1719-03-5
6	CI75 PERYLENE-D12	164IS6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBZN-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANILINE	65 S3# 45	88-74-4

No	Name		
48	C535 DIMETHYLPHTHALATE	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROTOLUENE	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	352	3:55	1	1.000	A BB	32486.	40.000 UG/ML	0.31
2	136	544	6:03	2	1.000	A BB	124189.	40.000 UG/ML	0.31
3	164	842	9:22	3	1.000	A BB	71932.	40.000 UG/ML	0.31
4	188	1101	12:15	4	1.000	A BB	116804.	40.000 UG/ML	0.31
5	240	1568	17:26	5	1.000	A BB	60904.	40.000 UG/ML	0.31
6	264	1798	20:00	6	1.000	A BB	50722.	40.000 UG/ML	0.31
7	82	437	4:52	2	0.803	A BB	224354.	160.000 UG/ML	1.26
8	172	737	8:12	3	0.875	A BB	321084.	160.000 UG/ML	1.26
9	244	1398	15:33	5	0.892	A VB	248225.	160.000 UG/ML	1.26
10	99	322	3:35	1	0.915	A BB	256146.	160.000 UG/ML	1.26
11	112	205	2:17	1	0.582	A BB	181194.	160.000 UG/ML	1.26
12	330	985	10:57	3	1.170	A BB	62460.	160.000 UG/ML	1.26
13	132	325	3:37	1	0.923	A BB	182800.	160.000 UG/ML	1.26
14	152	372	4:08	1	1.057	A BB	114576.	160.000 UG/ML	1.26
15	NOT FOUND								
16	74	109	1:13	1	0.310	A BB	135427.	160.000 UG/ML	1.26
17	93	312	3:28	1	0.886	A BV	217517.	160.000 UG/ML	1.26
18	94	323	3:36	1	0.918	A BB	257557.	160.000 UG/ML	1.26
19	93	328	3:39	1	0.932	A BB	163916.	160.000 UG/ML	1.26
20	128	327	3:38	1	0.929	A BB	188486.	160.000 UG/ML	1.26
21	146	343	3:49	1	0.974	A BV	196267.	160.000 UG/ML	1.26
22	146	354	3:56	1	1.006	A VB	197247.	160.000 UG/ML	1.26
23	108	383	4:16	1	1.088	A BB	119784.	160.000 UG/ML	1.26
24	146	374	4:10	1	1.062	A BB	183117.	160.000 UG/ML	1.26
25	108	403	4:29	1	1.145	A BB	157686.	160.000 UG/ML	1.26
26	45	401	4:28	1	1.139	A BB	323096.	160.000 UG/ML	1.26
27	108	431	4:48	1	1.224	A BB	175498.	160.000 UG/ML	1.26
28	70	426	4:44	1	1.210	A BV	115670.	160.000 UG/ML	1.26
29	117	420	4:40	1	1.193	A BB	91776.	160.000 UG/ML	1.26
30	77	440	4:54	2	0.809	A VB	203653.	160.000 UG/ML	1.26
31	82	484	5:23	2	0.890	qedt(1)	351500.	160.000 UG/ML	1.26
32	139	487	5:25	2	0.895	A BB	108037.	160.000 UG/ML	1.26
33	107	512	5:42	2	0.941	A BB	175563.	160.000 UG/ML	1.26
34	93	525	5:50	2	0.965	A BB 33	247182.	160.000 UG/ML	1.26
35	162	531	5:54	2	0.976	A BB 33	151820.	160.000 UG/ML	1.26
36	122	571	6:21	2	1.050	qedt(2)	145098.	160.000 UG/ML	1.26
37	180	538	5:59	2	0.989	A BB	154240.	160.000 UG/ML	1.26
38	128	548	6:06	2	1.007	A BB 27	464474.	160.000 UG/ML	1.26
39	127	570	6:20	2	1.048	A BB 27	225820.	160.000 UG/ML	1.26
40	225	573	6:22	2	1.053	A BV	78174.	160.000 UG/ML	1.26
41	107	662	7:22	2	1.217	A BB	172070.	160.000 UG/ML	1.26
42	142	666	7:24	2	1.224	qedt(2)	317611.	160.000 UG/ML	1.26
43	237	694	7:43	3	0.824	A BB	94850.	160.000 UG/ML	1.26
44	196	722	8:02	3	0.857	A BV	128899.	160.000 UG/ML	1.26
45	196	727	8:05	3	0.863	A VB	105318.	160.000 UG/ML	1.26
46	162	748	8:19	3	0.888	A BB	313318.	160.000 UG/ML	1.26
47	65	777	8:39	3	0.923	A BB	143092.	160.000 UG/ML	1.26
48	163	822	9:09	3	0.976	A BB	359876.	160.000 UG/ML	1.26
49	152	817	9:05	3	0.970	A BB	480732.	160.000 UG/ML	1.26
50	165	828	9:13	3	0.983	qedt(2)	91334.	160.000 UG/ML	1.26

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:54	1.01	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	6:01	1.01	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:21	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:25	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:59	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:49	1.01	0.800	1.00	160.00	160.00	0.452	0.452	1.00
8	8:11	1.00	0.875	1.00	160.00	160.00	1.116	1.116	1.00
9	15:32	1.00	0.891	1.00	160.00	160.00	1.019	1.019	1.00
10	3:32	1.02	0.906	1.01	160.00	160.00	1.971	1.971	1.00
11	2:15	1.01	0.580	1.00	160.00	160.00	1.394	1.394	1.00
12	10:55	1.00	1.168	1.00	160.00	160.00	0.217	0.217	1.00
13	3:34	1.01	0.917	1.01	160.00	160.00	1.407	1.407	1.00
14	4:07	1.01	1.057	1.00	160.00	160.00	0.882	0.882	1.00
15	12:39		0.941						
16	1:12	1.01	0.309	1.00	160.00	160.00	1.042	1.042	1.00
17	3:27	1.01	0.886	1.00	160.00	160.00	1.674	1.674	1.00
18	3:33	1.01	0.911	1.01	160.00	160.00	1.982	1.982	1.00
19	3:36	1.01	0.926	1.01	160.00	160.00	1.261	1.261	1.00
20	3:36	1.01	0.926	1.00	160.00	160.00	1.451	1.451	1.00
21	3:48	1.01	0.974	1.00	160.00	160.00	1.510	1.510	1.00
22	3:55	1.01	1.006	1.00	160.00	160.00	1.518	1.518	1.00
23	4:13	1.01	1.083	1.00	160.00	160.00	0.922	0.922	1.00
24	4:08	1.01	1.063	1.00	160.00	160.00	1.409	1.409	1.00
25	4:27	1.01	1.143	1.00	160.00	160.00	1.213	1.213	1.00
26	4:26	1.01	1.140	1.00	160.00	160.00	2.486	2.486	1.00
27	4:44	1.01	1.217	1.01	160.00	160.00	1.351	1.351	1.00
28	4:40	1.01	1.200	1.01	160.00	160.00	0.890	0.890	1.00
29	4:39	1.00	1.194	1.00	160.00	160.00	0.706	0.706	1.00
30	4:51	1.01	0.806	1.00	160.00	160.00	0.410	0.410	1.00
31	5:19	1.01	0.884	1.01	160.00	160.00	0.708	0.708	1.00
32	5:24	1.00	0.896	1.00	160.00	160.00	0.217	0.217	1.00
33	5:39	1.01	0.939	1.00	160.00	160.00	0.353	0.353	1.00
34	5:48	1.01	0.965	1.00	160.00	160.00	0.498	0.498	1.00
35	5:52	1.01	0.974	1.00	160.00	160.00	0.306	0.306	1.00
36	6:10	1.03	1.024	1.03	160.00	160.00	0.292	0.292	1.00
37	5:57	1.01	0.989	1.00	160.00	160.00	0.310	0.310	1.00
38	6:04	1.01	1.007	1.00	160.00	160.00	0.935	0.935	1.00
39	6:16	1.01	1.043	1.01	160.00	160.00	0.455	0.455	1.00
40	6:22	1.00	1.057	1.00	160.00	160.00	0.157	0.157	1.00
41	7:20	1.00	1.220	1.00	160.00	160.00	0.346	0.346	1.00
42	7:22	1.00	1.226	1.00	160.00	160.00	0.639	0.639	1.00
43	7:42	1.00	0.824	1.00	160.00	160.00	0.330	0.330	1.00
44	8:00	1.00	0.856	1.00	160.00	160.00	0.448	0.448	1.00
45	8:03	1.00	0.862	1.00	160.00	160.00	0.366	0.366	1.00
46	8:17	1.00	0.887	1.00	160.00	160.00	1.089	1.089	1.00
47	8:37	1.00	0.921	1.00	160.00	160.00	0.497	0.497	1.00
48	9:06	1.00	0.974	1.00	160.00	160.00	1.251	1.251	1.00
49	9:03	1.00	0.969	1.00	160.00	160.00	1.671	1.671	1.00
50	9:09	1.01	0.980	1.00	160.00	160.00	0.317	0.317	1.00

Data: ST16980827E.TI

08/27/98 11:50:00

Sample: SSTD160 160UG/ML CALI I 070698I

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name		
51	C545 3-NITROANILINE	138 S3# 75	99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC	83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP	51-28-5
54	C565 DIBENZOFURAN	168 S3# 90	132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP	100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105	121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130	84-66-2
58	C590 FLUORENE	166 S3#135	86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140	7005-72-3
60	C595 4-NITROANALINE	138 S3#150	100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10	534-52-1
62	C615 N-NITROSODIPHENYLAMINE	169 S4# 15CC	87-30-6
63	C620 AZOBENZENE	77 S4# 20	103-33-3
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30	101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40	118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50	87-86-5
67	C640 PHENANTHRENE	178 S4# 65	85-01-8
68	C645 ANTHRACENE	178 S4# 70	120-12-7
69	C647 CARBAZOLE	167 S4# 80	86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85	84-74-2
71	C655 FLUORANTHENE	202 S4#110CC	206-44-0
72	C710 BENZIDINE	184 S5# 10	92-81-5
73	C715 PYRENE	202 S5# 15	129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40	85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50	56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55	91-94-1
77	C740 CHRYSENE	228 S5# 60	218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65	117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC	117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15	205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25	207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35	50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55	193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60	53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65	191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	XTot
51	138	853	9:29	3	1.013	A BB	108951.	160.000 UG/ML	1.26
52	153	849	9:27	3	1.008	A BB	306695.	160.000 UG/ML	1.26
53	184	872	9:42	3	1.036	A BB	56407.	160.000 UG/ML	1.26
54	168	881	9:48	3	1.046	A BB	449013.	160.000 UG/ML	1.26
55	109	901	10:01	3	1.070	qedt	48707.	160.000 UG/ML	1.26
56	165	896	9:58	3	1.064	A BB	108669.	160.000 UG/ML	1.26
57	149	950	10:34	3	1.128	qedt	347720.	160.000 UG/ML	1.26
58	166	941	10:28	3	1.118	A BV	341839.	160.000 UG/ML	1.26

RATED

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
59	204	951	10:35	3	1.129	A BB	165442.	160.000 UG/ML	1.26
60	138	965	10:44	3	1.146	qedt	92245.	160.000 UG/ML	1.26
61	198	969	10:47	4	0.880	qedt	60946.	160.000 UG/ML	1.26
62	169	978	10:53	4	0.888	A BB	244263.	160.000 UG/ML	1.26
63	77	979	10:53	4	0.889	A BB	541010.	160.000 UG/ML	1.26
64	248	1035	11:31	4	0.940	A BB	98271.	160.000 UG/ML	1.26
65	284	1034	11:30	4	0.939	A BB	112917.	160.000 UG/ML	1.26
66	266	1076	11:58	4	0.977	A BB	68614.	160.000 UG/ML	1.26
67	178	1106	12:18	4	1.005	A BV	471716.	160.000 UG/ML	1.26
68	178	1115	12:24	4	1.013	A VB	378128.	160.000 UG/ML	1.26
69	167	1153	12:49	4	1.047	A BB	327617.	160.000 UG/ML	1.26
70	149	1241	13:48	4	1.127	A BB	504598.	160.000 UG/ML	1.26
71	202	1314	14:37	4	1.193	A BB	334270.	160.000 UG/ML	1.26
72	184	1353	15:03	5	0.863	A BB	32031.	160.000 UG/ML	1.26
73	202	1350	15:01	5	0.861	A BB	330800.	160.000 UG/ML	1.26
74	149	1502	16:42	5	0.958	A BB	178737.	160.000 UG/ML	1.26
75	228	1566	17:25	5	0.999	A VV	265942.	160.000 UG/ML	1.26
76	252	1577	17:32	5	1.006	A BB	95194.	160.000 UG/ML	1.26
77	228	1573	17:30	5	1.003	A VB	223869.	160.000 UG/ML	1.26
78	149	1617	17:59	5	1.031	A BB	265391.	160.000 UG/ML	1.26
79	149	1729	19:14	6	0.962	A BB	410452.	160.000 UG/ML	1.26
80	252	1745	19:25	6	0.971	A BV	286447.	160.000 UG/ML	1.26
81	252	1750	19:28	6	0.973	A VB	190774.	160.000 UG/ML	1.26
82	252	1792	19:56	6	0.997	A BB	219973.	160.000 UG/ML	1.26
83	276	1947	21:39	6	1.083	A BV	232127.	160.000 UG/ML	1.26
84	278	1954	21:44	6	1.087	A BB	224248.	160.000 UG/ML	1.26
85	276	1980	22:01	6	1.101	A BB	231038.	160.000 UG/ML	1.26

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:26	1.01	1.010	1.00	160.00	160.00	0.379	0.379	1.00
52	9:25	1.00	1.007	1.00	160.00	160.00	1.066	1.066	1.00
53	9:39	1.01	1.032	1.00	160.00	160.00	0.196	0.196	1.00
54	9:46	1.00	1.045	1.00	160.00	160.00	1.561	1.561	1.00
55	9:59	1.00	1.068	1.00	160.00	160.00	0.169	0.169	1.00
56	9:55	1.01	1.061	1.00	160.00	160.00	0.378	0.378	1.00
57	10:31	1.00	1.126	1.00	160.00	160.00	1.209	1.209	1.00
58	10:26	1.00	1.117	1.00	160.00	160.00	1.188	1.188	1.00
59	10:33	1.00	1.130	1.00	160.00	160.00	0.575	0.575	1.00
60	10:39	1.01	1.139	1.01	160.00	160.00	0.321	0.321	1.00
61	10:42	1.01	0.875	1.01	160.00	160.00	0.130	0.130	1.00
62	10:50	1.00	0.886	1.00	160.00	160.00	0.523	0.523	1.00
63	10:51	1.00	0.888	1.00	160.00	160.00	1.158	1.158	1.00
64	11:29	1.00	0.940	1.00	160.00	160.00	0.210	0.210	1.00
65	11:29	1.00	0.939	1.00	160.00	160.00	0.242	0.242	1.00
66	11:57	1.00	0.977	1.00	160.00	160.00	0.147	0.147	1.00
67	12:17	1.00	1.005	1.00	160.00	160.00	1.010	1.010	1.00
68	12:22	1.00	1.012	1.00	160.00	160.00	0.809	0.809	1.00
69	12:47	1.00	1.046	1.00	160.00	160.00	0.701	0.701	1.00
70	13:47	1.00	1.127	1.00	160.00	160.00	1.080	1.080	1.00
71	14:36	1.00	1.194	1.00	160.00	160.00	0.715	0.715	1.00
72	15:03	1.00	0.864	1.00	160.00	160.00	0.131	0.131	1.00
73	15:00	1.00	0.861	1.00	160.00	160.00	1.358	1.358	1.00
74	16:41	1.00	0.958	1.00	160.00	160.00	0.734	0.734	1.00
75	17:24	1.00	0.999	1.00	160.00	160.00	1.092	1.092	1.00
76	17:31	1.00	1.006	1.00	160.00	160.00	0.391	0.391	1.00
77	17:28	1.00	1.003	1.00	160.00	160.00	0.919	0.919	1.00

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:58	1.00	1.032	1.00	160.00	160.00	1.089	1.089	1.00
79	19:13	1.00	0.961	1.00	160.00	160.00	2.023	2.023	1.00
80	19:23	1.00	0.969	1.00	160.00	160.00	1.412	1.412	1.00
81	19:26	1.00	0.972	1.00	160.00	160.00	0.940	0.940	1.00
82	19:54	1.00	0.996	1.00	160.00	160.00	1.084	1.084	1.00
83	21:38	1.00	1.082	1.00	160.00	160.00	1.144	1.144	1.00
84	21:43	1.00	1.086	1.00	160.00	160.00	1.105	1.105	1.00
85	21:59	1.00	1.100	1.00	160.00	160.00	1.139	1.139	1.00

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPHY**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND CODED**



SCANS 450 TO 520

DATA: ST16980827E #1  
CALI: ST16980827E #3

RIC+MASS CHROMATOGRAM

08/27/98 11:50:00

SAMPLE: SSTD160 160UG/ML CALI 1 0706981

CONDS.: INST. ID: F16

RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

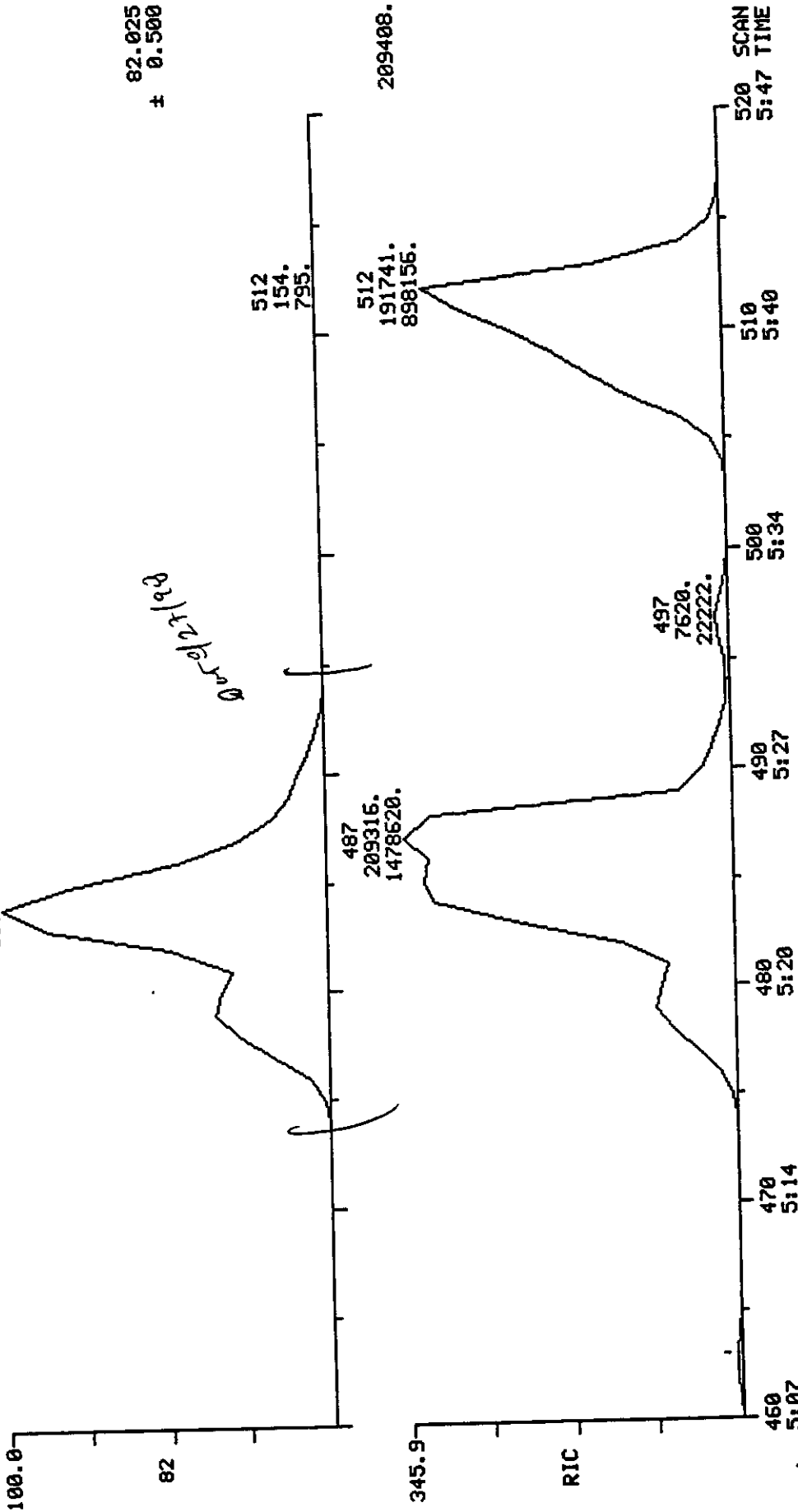
484

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351149.

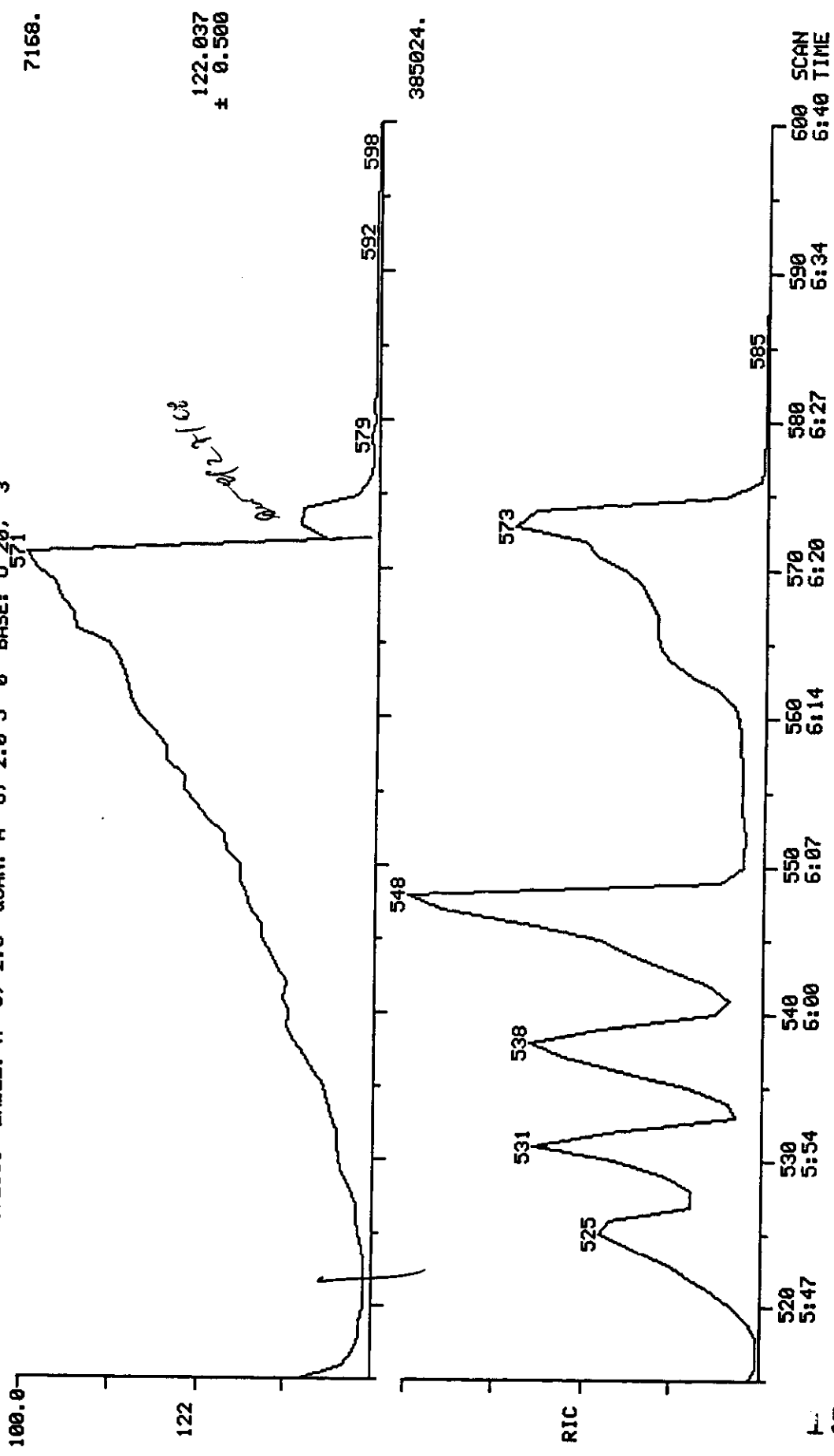
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82.025  
± 0.500



8201

RIC+MASS CHROMATOGRAM  
 08/27/98 11:50:00  
 SAMPLE: SST0160 160UG/ML CALI I 070698I  
 CONDS.: INST. ID: F15  
 RANGE: G 1.2158 LABEL: N 0. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20. 3  
 DATA: ST116980827E #1  
 CALI: ST116980827E #3  
 SCANS 515 TO 600



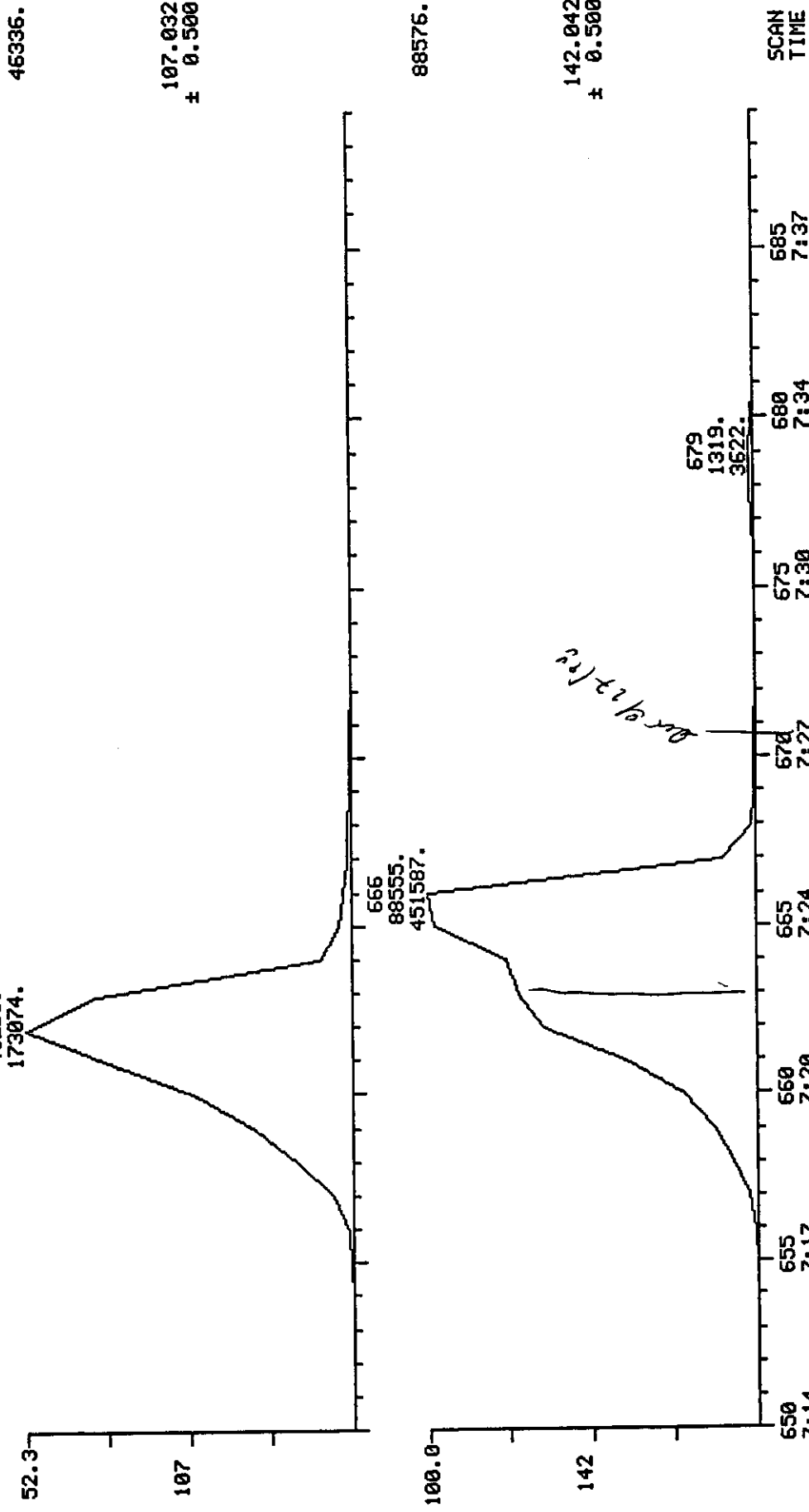
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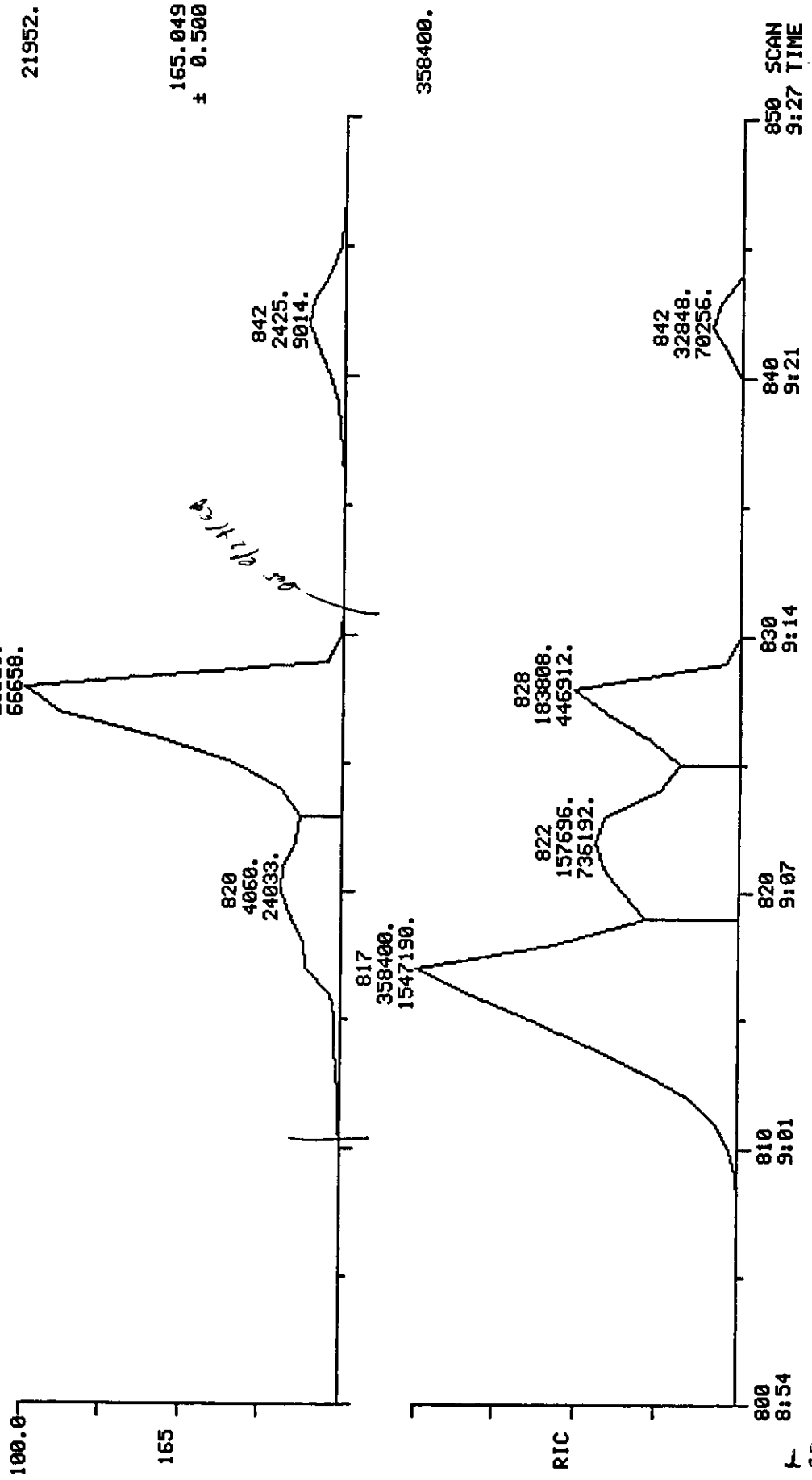
4530

MASS CHROMATOGRAMS  
 08/27/98 11:50:00 DATA: ST16980827E #1 SCANS 650 TO 689  
 CALI: ST16980827E #3  
 SAMPLE: SSTD160 160UG/ML CALI I 0706981  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2.2.0 QUAN: A 6. 2.0 J 0 BASE: U 20, 3



1531

RIC+MASS CHROMATOGRAM  
 08/27/98 11:50:00 DATA: ST16980827E #1 SCANS 800 TO 850  
 CALI: ST16980827E #3  
 SAMPLE: SST0160 160UG/ML CALI I 0706981  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3



SCANS 870 TO 930

DATA: ST16980827E #1  
CALI: ST16980827E #3

RIC+MASS CHROMATOGRAM

08/27/98 11:50:00

SAMPLE: SSTD160 160UG/ML CALI I 070698I

CONDS.: INST. ID: F16

RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

901

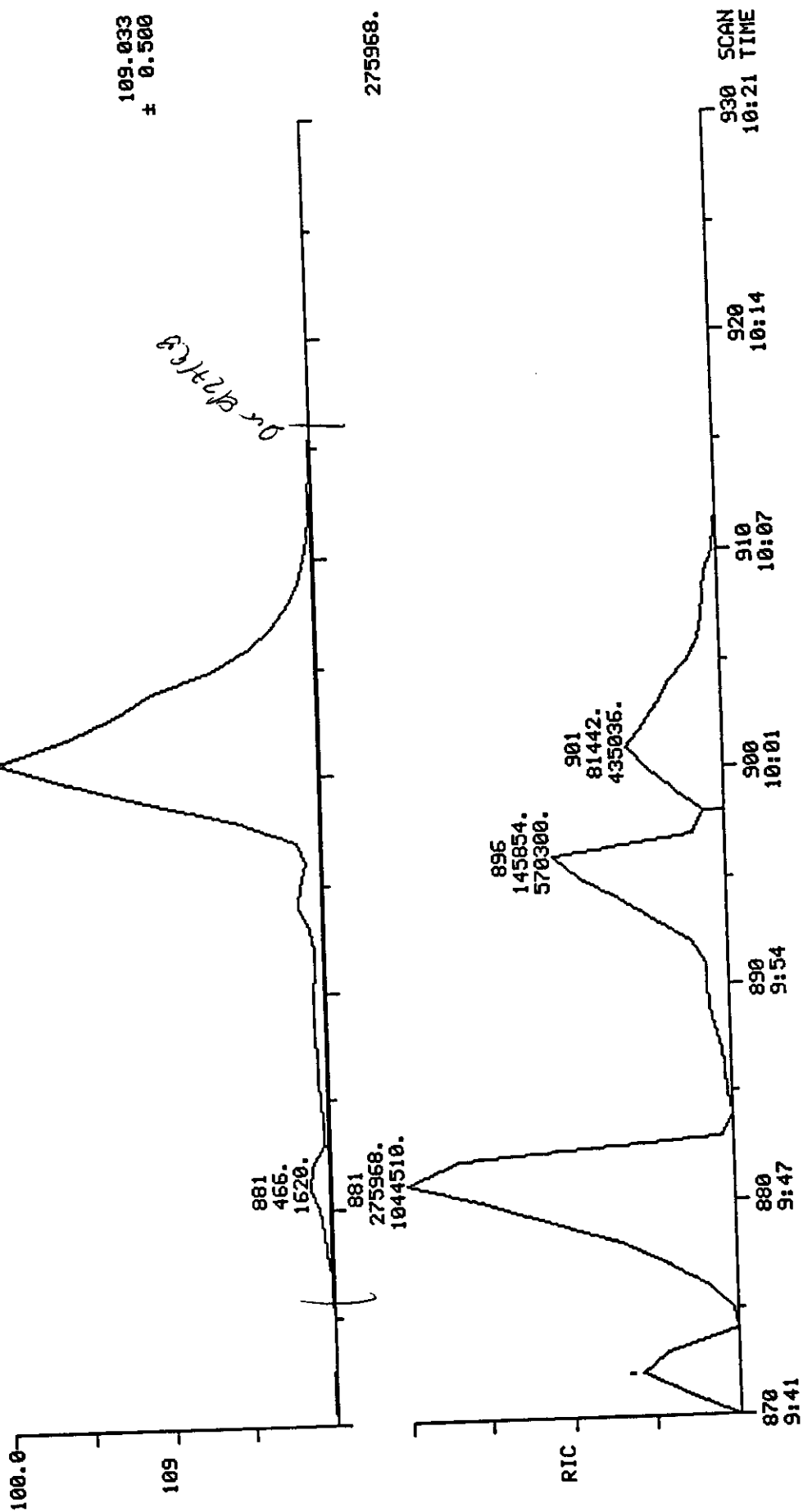
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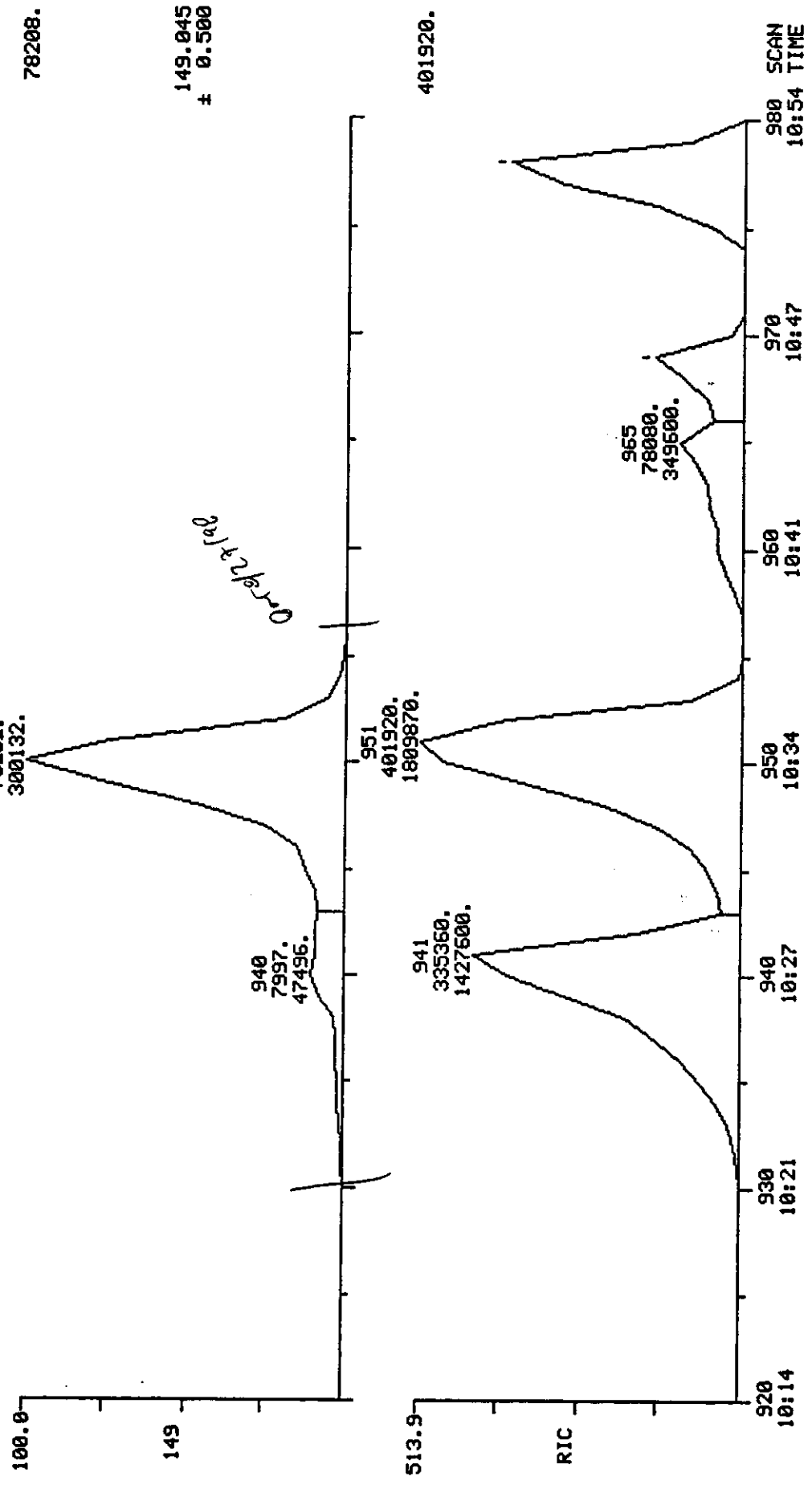
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1505

RIC+MASS CHROMATOGRAM  
 08/27/98 11:50:00 DATA: ST16980827E #1 SCANS 920 TO 980  
 CALI: ST16980827E #3  
 SAMPLE: SSTD160 160UG/ML CALI I 0706981  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3



1584

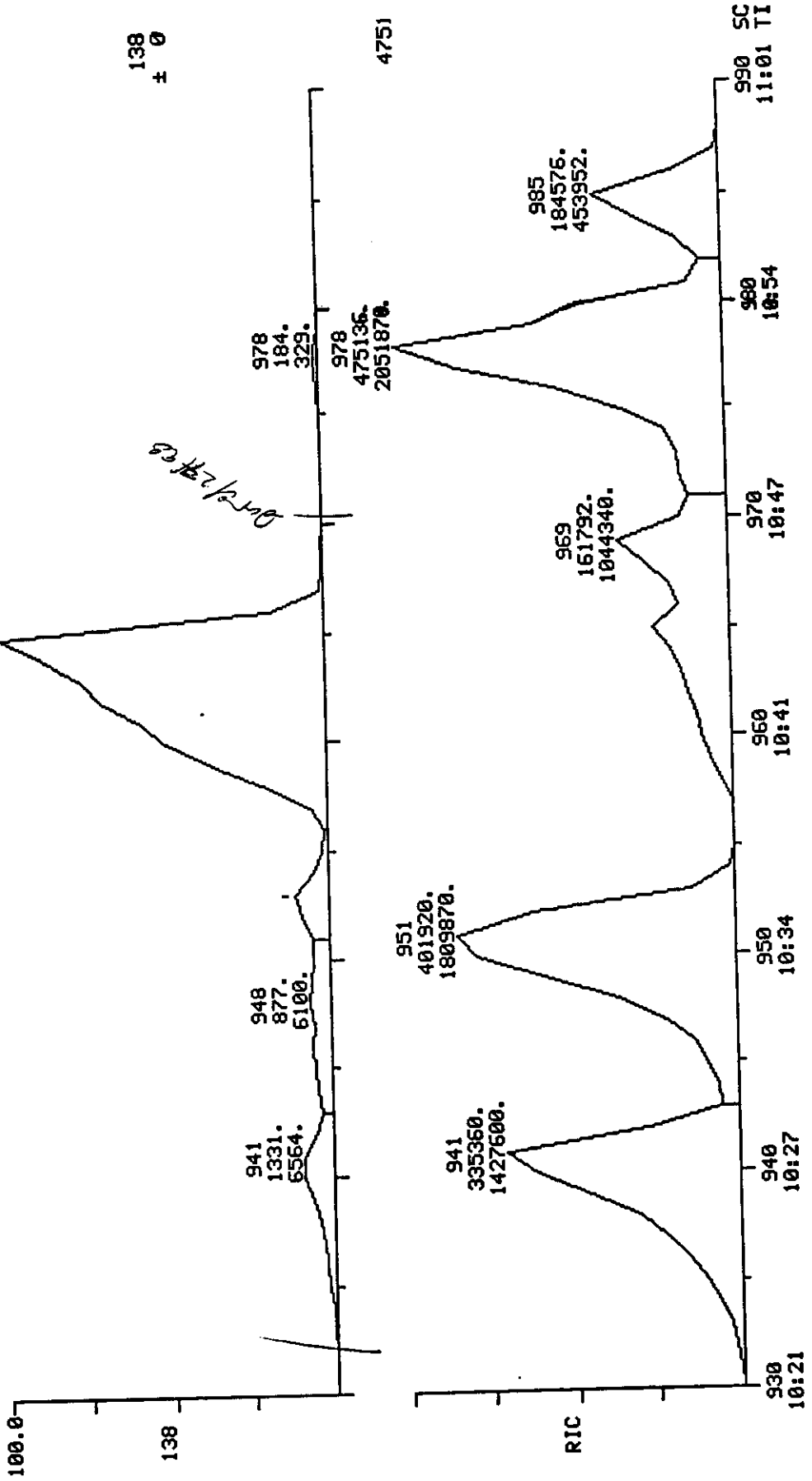
SCANS 930 TO 990

DATA: ST16980827E #1  
CALI: ST16980827E #3

RIC+MASS CHROMATOGRAM  
08/27/98 11:50:00  
SAMPLE: SSTD160 160UG/ML CALI I 0706981

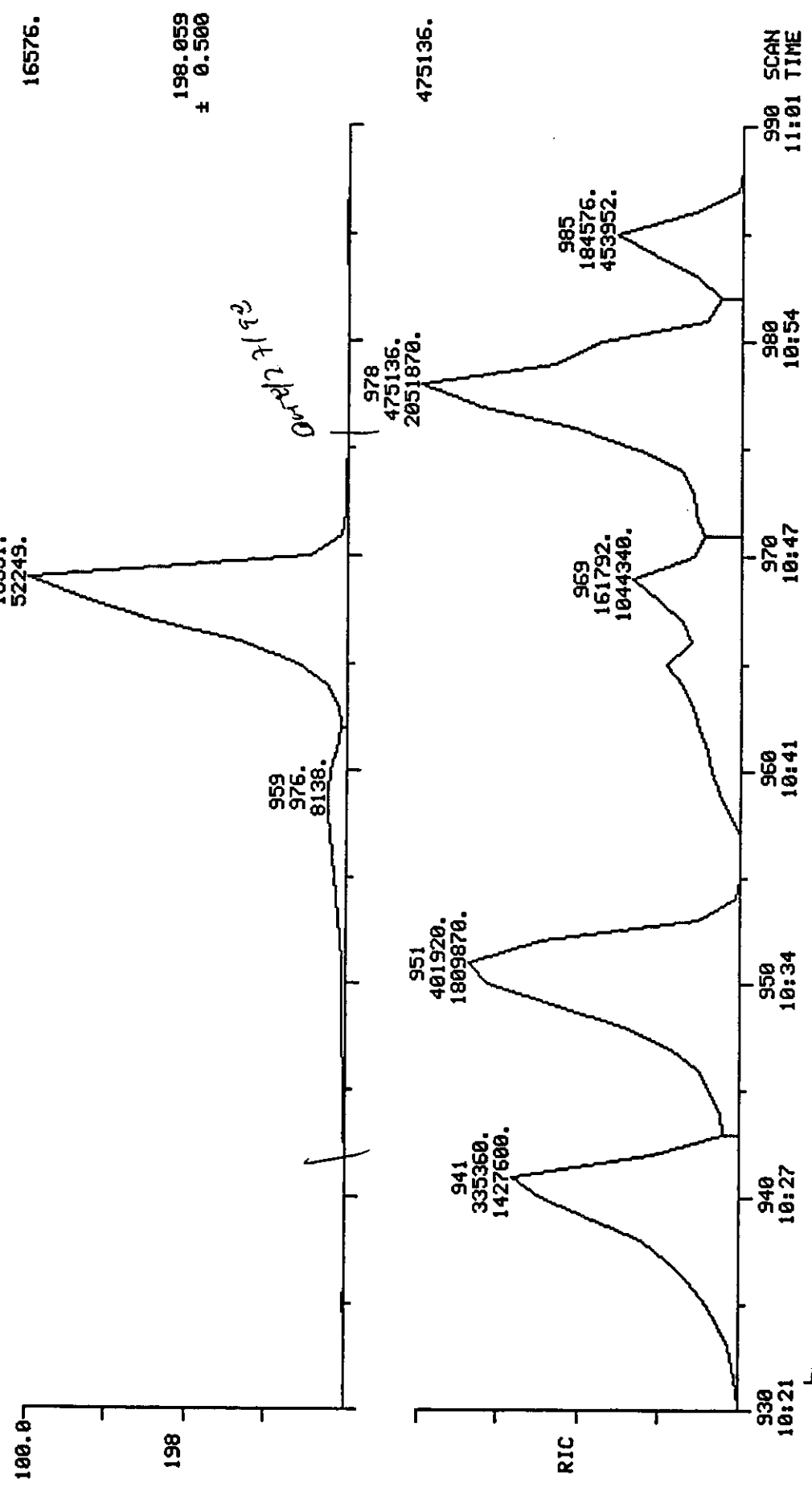
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 2. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20. 3  
965  
14453.  
74867.

14



1535

RIC+MASS CHROMATOGRAM  
 08/27/98 11:50:00 DATA: ST16980827E #1 SCANS 930 TO 990  
 CALI: ST16980827E #3  
 SAMPLE: SSTD160 160UG/ML CALI I 070698I  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3



4500



QUANTERRA GC/MS  
SEMIVOLATILE APPENDIX IX CALIBRATION CHE

INITIAL CALIBRATION: 10 UG/ML APPENDIX ST16980831B 08/31/98 13:35:00  
 INITIAL CALIBRATION: 20 UG/ML APPENDIX ST16980831C 08/31/98 14:02:00  
 INITIAL CALIBRATION: 50 UG/ML APPENDIX ST16980831D 08/31/98 14:32:00  
 INITIAL CALIBRATION: 80 UG/ML APPENDIX ST16980831A 08/31/98 13:07:00  
 INITIAL CALIBRATION: 120 UG/ML APPENDIX ST16980831E 08/31/98 15:02:00  
 INITIAL CALIBRATION: 160 UG/ML APPENDIX ST16980831F 08/31/98 15:31:00

Library Entry	Compound	Initial Calibration			Continuing Calibration	
		Response Factor Avg	% RSD Min	% RSD Max	Response Factor Min	% Diff Max
S1	20 PYRIDINE	1.590		6.5		
S1	30 2-PICOLINE	1.565		1.3		
S1	40 N-NITROSOMETHYLETHYL	0.811		9.2		
S1	60 METHYLMETHANESULFONA	0.636		6.2		
S1	70 N-NITROSODIETHYLAMIN	0.521		7.6		
S1	85 ETHYLMETHANE SULFONA	1.109		11.0		
S1	95 PENTACHLOROETHANE	0.602		1.4		
S1	170 C361 ACETOPHENONE	1.934		2.8		
S1	175 N-NITROSOPYRROLIDINE	0.553		6.5		
S1	180 N-NITROSOMORPHOLINE	0.821		1.4		
S1	182 3-METHYL PHENOL	2.252		9.9		
S1	195 O-TOLUIDINE	2.160		9.2		
S2	15 N-NITROSOPIPERIDINE	0.321		4.9		
S2	50 AA-DIMETHYLPHENETHYL	1.285		15.6		
S2	85 2,6-DICHLOROPHENOL	0.257		12.8		
S2	90 HEXACHLOROPROPENE	0.198		0.9		
S2	115 P-PHENYLENE DIAMINE	0.131		42.2		
S2	120 N-NITROSODI-N-BUTYLA	0.213		7.2		
S2	140 SAFROLE	0.248		0.9		
S3	10 1,2,4,5-TETRACHLOROB	0.433		4.4		
S3	15 ISOSAFROLE (#1)	0.044		3.4		
S3	35 ISOSAFROLE (#2)	0.195		2.2		
S3	42 1-CHLORONAPHTH	1.318		16.5		
S3	50 1,4-NAPHTHOQUINONE	0.414		5.6		
S3	60 1,3-DINITROBENZENE	0.198		8.6		
S3	100 PENTACHLOROBENZENE	0.489		3.8		
S3	110 1-NAPHTHYLAMINE	0.837		6.6		
S3	115 2-NAPHTHYLAMINE	0.966		5.8		
S3	120 2,3,4,6-TETRACHLOROP	0.297		4.5		
S3	145 5-NITRO-O-TOLUIDINE	0.355		7.2		
S4	25 SYM-TRINITROBENZENE	0.162		5.3		
S4	35 PHENACETIN	0.317		3.2		
S4	37 DIALLATE	0.090		4.4		
S4	45 4-AMINOBIIPHENYL	0.609		7.6		
S4	55 PRONAMIDE	0.299		5.2		
S4	60 PENTACHLORONITROBENZ	0.080		1.8		
S4	75 2SECBUTYL-4,6-DINITR	0.153		9.3		
S4	100 4-NITROQUINOLINE-1-O	0.072		11.5		
S4	105 METHAPYRILENE	0.305		26.8		
S4	106 ISODRIN	0.127		1.5		
S4	120 CHLOROBENZILATE	0.323		11.1		
S5	20 ARAMITE (#1)	0.061		4.4		
S5	25 ARAMITE (#2)	0.087		4.4		
S5	30 P-DIMETHYLAMINOAZOBE	0.419		1.1		
S5	35 3,3'-DIMETHYLBENZIDI	0.418		17.0		
S5	37 KEPONE	0.082		34.6		
S5	45 2-ACETYLAMINOFLUOREN	0.432		4.0		
S5	85 3-METHYLCHOLANTHRENE	0.548		6.1		

1537

S6 20 7,12-DIMETHYLBENZANT 0.610 0.5

0 CCC and SPCC compounds are out

1533

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QUANTERRA GC/MS  
SEMIVOLATILE APPENDIX IX CALIBRATION CHE

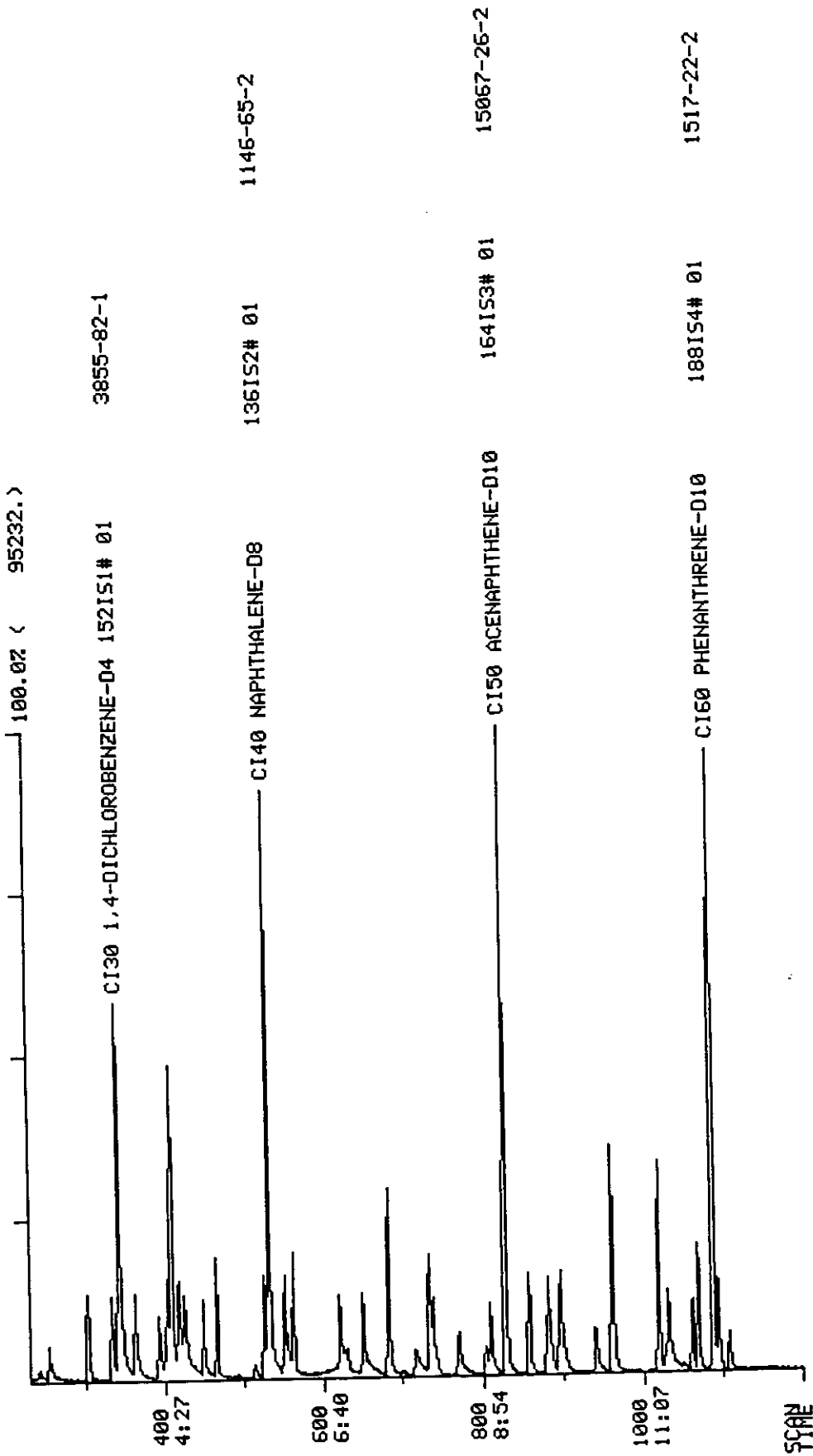
INITIAL CALIBRATION: 10 UG/ML APPENDIX ST16980831B 08/31/98 13:35:00  
 INITIAL CALIBRATION: 20 UG/ML APPENDIX ST16980831C 08/31/98 14:02:00  
 INITIAL CALIBRATION: 50 UG/ML APPENDIX ST16980831D 08/31/98 14:32:00  
 INITIAL CALIBRATION: 80 UG/ML APPENDIX ST16980831A 08/31/98 13:07:00  
 INITIAL CALIBRATION: 120 UG/ML APPENDIX ST16980831E 08/31/98 15:02:00  
 INITIAL CALIBRATION: 160 UG/ML APPENDIX ST16980831F 08/31/98 15:31:00

	ST16980831B		ST16980831C		ST16980831D		ST16980831A		ST16980831E		ST16980831F	
Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S1	20	10, 1.770	20, 1.583	50, 1.578	80, 1.587	120, 1.449	160, 1.576					
S1	30		20, 1.563	50, 1.594	80, 1.577	120, 1.547	160, 1.545					
S1	40	10, 0.700	20, 0.737	50, 0.823	80, 0.876	120, 0.860	160, 0.870					
S1	60	10, 0.619	20, 0.688	50, 0.685	80, 0.602	120, 0.619	160, 0.604					
S1	70	10, 0.450	20, 0.506	50, 0.521	80, 0.549	120, 0.547	160, 0.555					
S1	85	10, 0.921	20, 1.008	50, 1.120	80, 1.168	120, 1.190	160, 1.244					
S1	95		20, 0.612	50, 0.605	80, 0.605	120, 0.595	160, 0.592					
S1	170	10, 1.869	20, 2.024	50, 1.960	80, 1.945	120, 1.903	160, 1.903					
S1	175	10, 0.481	20, 0.555	50, 0.575	80, 0.562	120, 0.569	160, 0.575					
S1	180	10, 0.828	20, 0.838	50, 0.821	80, 0.805	120, 0.817	160, 0.816					
S1	182	10, 1.848	20, 2.130	50, 2.339	80, 2.408	120, 2.391	160, 2.394					
S1	195		20, 2.463	50, 2.228	80, 2.129	120, 2.032	160, 1.949					
S2	15	10, 0.293	20, 0.312	50, 0.326	80, 0.327	120, 0.337	160, 0.329					
S2	50		20, 1.575	50, 1.353	80, 1.036	120, 1.276	160, 1.184					
S2	85	10, 0.202	20, 0.234	50, 0.264	80, 0.280	120, 0.279	160, 0.285					
S2	90		20, 0.196	50, 0.201	80, 0.197	120, 0.198	160, 0.198					
S2	115		20, 0.163	50, 0.167	80, 0.038	120, 0.165	160, 0.121					
S2	120	10, 0.186	20, 0.205	50, 0.213	80, 0.225	120, 0.224	160, 0.224					
S2	140		20, 0.246	50, 0.247	80, 0.250	120, 0.245	160, 0.250					
S3	10	10, 0.469	20, 0.431	50, 0.434	80, 0.430	120, 0.416	160, 0.419					
S3	15		20, 0.046	50, 0.042	80, 0.043	120, 0.044	160, 0.044					
S3	35		20, 0.201	50, 0.195	80, 0.196	120, 0.190	160, 0.192					
S3	42	10, 1.531	20, 1.098	50, 1.129	80, 1.528	120, 1.134	160, 1.488					
S3	50		20, 0.403	50, 0.429	80, 0.447	120, 0.404	160, 0.389					
S3	60	10, 0.166	20, 0.191	50, 0.202	80, 0.210	120, 0.212	160, 0.204					

S3 100	10, 0.516	20, 0.504	50, 0.483	80, 0.493	120, 0.469	160, 0.471
S3 110	10, 0.869	20, 0.859	50, 0.889	80, 0.765	120, 0.872	160, 0.769
S3 115	10, 0.924	20, 0.974	50, 1.041	80, 0.996	120, 0.980	160, 0.881
S3 120		20, 0.277	50, 0.302	80, 0.310	120, 0.306	160, 0.290
S3 145		20, 0.345	50, 0.368	80, 0.381	120, 0.367	160, 0.316
S4 25		20, 0.149	50, 0.162	80, 0.172	120, 0.166	160, 0.159
S4 35		20, 0.305	50, 0.324	80, 0.316	120, 0.330	160, 0.311
S4 37		20, 0.088	50, 0.087	80, 0.089	120, 0.089	160, 0.097
S4 45		20, 0.643	50, 0.664	80, 0.602	120, 0.590	160, 0.546
S4 55		20, 0.312	50, 0.315	80, 0.301	120, 0.291	160, 0.277
S4 60		20, 0.081	50, 0.079	80, 0.079	120, 0.082	160, 0.081
S4 75		20, 0.130	50, 0.165	80, 0.163	120, 0.159	160, 0.150
S4 100		20, 0.058	50, 0.074	80, 0.076	120, 0.079	160, 0.075
S4 105		20, 0.424	50, 0.339	80, 0.205	120, 0.279	160, 0.277
S4 106	10, 0.127	20, 0.125	50, 0.129	80, 0.128	120, 0.127	160, 0.124
S4 120	10, 0.376	20, 0.351	50, 0.329	80, 0.308	120, 0.295	160, 0.281
S5 20		20, 0.057	50, 0.063	80, 0.061	120, 0.064	160, 0.061
S5 25		20, 0.081	50, 0.090	80, 0.090	120, 0.087	160, 0.085
S5 30		20, 0.417	50, 0.425	80, 0.420	120, 0.412	160, 0.420
S5 35		20, 0.522	50, 0.450	80, 0.406	120, 0.364	160, 0.346
S5 37		20, 0.123	50, 0.087	80, 0.089	120, 0.056	160, 0.054
S5 45		20, 0.404	50, 0.450	80, 0.438	120, 0.429	160, 0.440
S5 85		20, 0.586	50, 0.576	80, 0.549	120, 0.515	160, 0.514
S6 20		20, 0.609	50, 0.615	80, 0.610	120, 0.608	160, 0.609

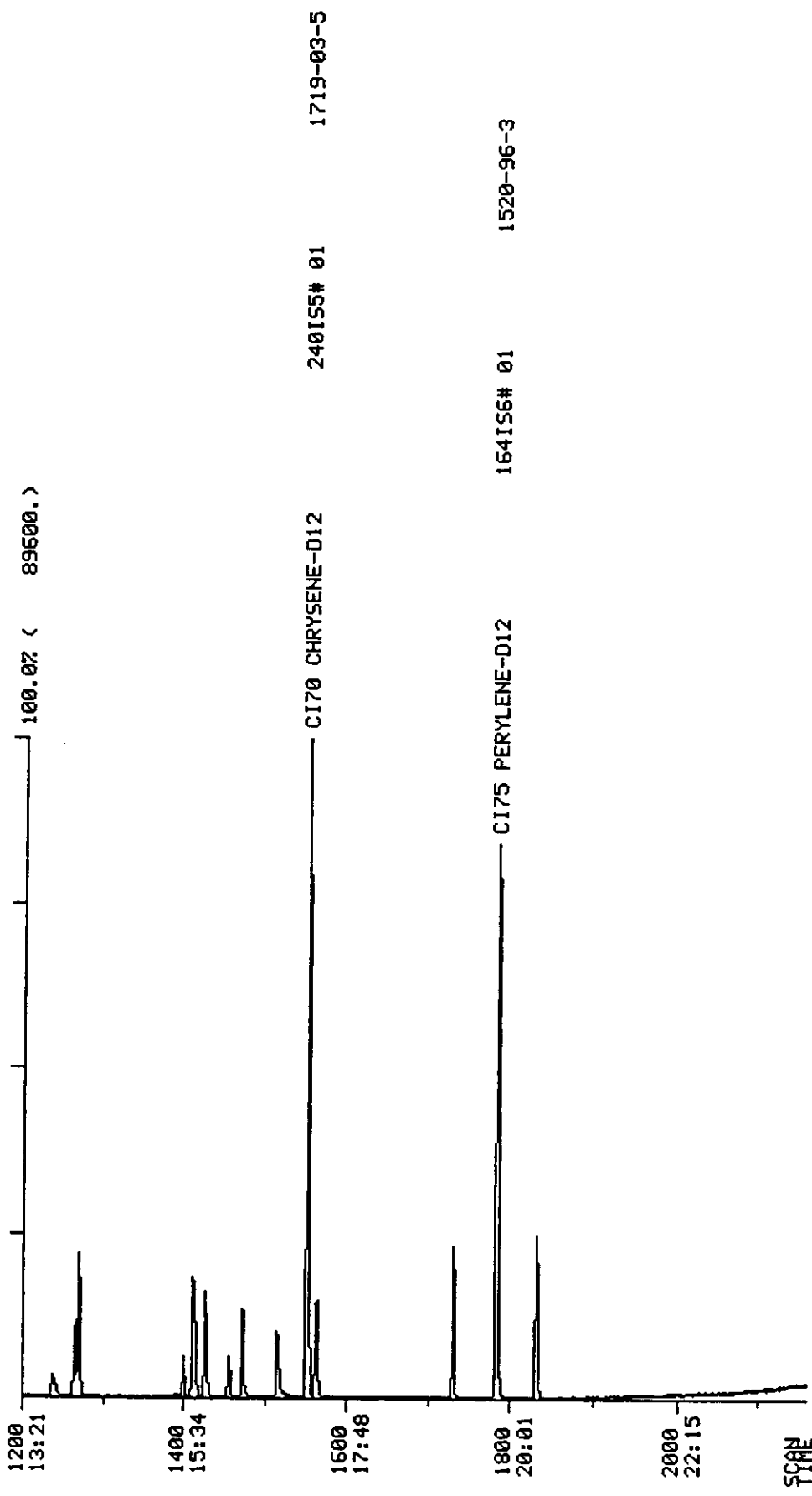
DATA FROM FILE: ST16980831B SCANS 230 TO 1200 ACQUIRED: 08/31/98 13:35:00  
CALL: ST16980831B #3

SAMPLE: 10UG/ML AP9 062598C  
CONDS.: INST. ID: F16



1541

DATA FROM FILE: ST16980831B SCANS 1200 TO 2158 ACQUIRED: 08/31/98 13:35:00  
CALI: ST16980831B #3  
SAMPLE: 10UG/ML AP9 062598C  
CONDS.: INST. ID: F16



Data: ST16980831B.T1

08/31/98 13:35:00

Sample: 10UG/ML AP9 062598C

Conds.: INST. ID: F16

Formula: IUL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
9	METHYLMETHANESULFONATE	80 S1# 60	
10	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
11	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
12	C361 ACETOPHENONE	105 S1#170	
13	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
14	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
15	3-METHYL PHENOL	108 S1#182	
16	N-NITROSOPIPERIDINE	42 S2# 15	100-75-4
17	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
18	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
19	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
20	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
21	1,3-DINITROBENZENE	168 S3# 60	99-65-0
22	PENTACHLOROBENZENE	250 S3#100	708-93-5
23	1-NAPHTHYLAMINE	143 S3#110	134-32-7
24	2-NAPHTHYLAMINE	143 S3#115	91-59-8
25	ISODRIN		
26	CHLOROBENZILATE	139 S4#120	

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	29461.	40.000 UG/ML	9.09
2	136	529	5:53	2	1.000	A BB	109632.	40.000 UG/ML	9.09
3	164	827	9:12	3	1.000	A BB	59743.	40.000 UG/ML	9.09
4	188	1085	12:04	4	1.000	A BB	100292.	40.000 UG/ML	9.09
5	240	1551	17:15	5	1.000	A BB	83642.	40.000 UG/ML	9.09
6	264	1782	19:49	6	1.000	A BB	69904.	40.000 UG/ML	9.09
7	79	103	1:09	1	0.303	A BB	13033.	10.000 UG/ML	2.27
8	42	157	1:45	1	0.462	A BB	5159.	10.000 UG/ML	2.27
9	80	181	2:01	1	0.532	A BB	4559.	10.000 UG/ML	2.27
10	102	216	2:24	1	0.635	A BB	3318.	10.000 UG/ML	2.27
11	79	253	2:49	1	0.744	A BB	6780.	10.000 UG/ML	2.27
12	105	401	4:28	1	1.179	A BB	13763.	10.000 UG/ML	2.27
13	100	399	4:26	1	1.174	qedt	3544.	10.000 UG/ML	2.27
14	56	407	4:32	1	1.197	A BB	6098.	10.000 UG/ML	2.27
15	108	416	4:38	1	1.224	A BB	13612.	10.000 UG/ML	2.27
16	42	447	4:58	2	0.845	A BB	8018.	10.000 UG/ML	2.27
17	162	551	6:08	2	1.042	A BB	5546.	10.000 UG/ML	2.27

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
18	84	618	6:52	2	1.168	A BB	5111.	10.000 UG/ML	2.27
19	214	680	7:34	3	0.822	A BB	6998.	10.000 UG/ML	2.27
20	162	732	8:09	3	0.885	A BB	22862.	10.000 UG/ML	2.27
21	168	802	8:55	3	0.970	A BB	2486.	10.000 UG/ML	2.27
22	250	857	9:32	3	1.036	A BB	7702.	10.000 UG/ML	2.27
23	143	881	9:48	3	1.065	A BB	12978.	10.000 UG/ML	2.27
24	143	896	9:58	3	1.083	A BB	13797.	10.000 UG/ML	2.27
25	193	1269	14:07	4	1.170	A BB	3194.	10.000 UG/ML	2.27
26	139	1426	15:52	4	1.314	A BB	9428.	10.000 UG/ML	2.27

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:16	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:50	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	10.00	10.00	1.770	1.770	1.00
8	1:45	1.00	0.462	1.00	10.00	10.00	0.700	0.700	1.00
9	2:01	0.99	0.535	0.99	10.00	10.00	0.619	0.619	1.00
10	2:25	0.99	0.641	0.99	10.00	10.00	0.450	0.450	1.00
11	2:50	0.99	0.750	0.99	10.00	10.00	0.921	0.921	1.00
12	4:30	0.99	1.188	0.99	10.00	10.00	1.869	1.869	1.00
13	4:30	0.99	1.191	0.99	10.00	10.00	0.481	0.481	1.00
14	4:34	0.99	1.209	0.99	10.00	10.00	0.828	0.828	1.00
15	4:40	0.99	1.232	0.99	10.00	10.00	1.848	1.848	1.00
16	5:01	0.99	0.851	0.99	10.00	10.00	0.293	0.293	1.00
17	6:08	1.00	1.042	1.00	10.00	10.00	0.202	0.202	1.00
18	6:54	1.00	1.170	1.00	10.00	10.00	0.186	0.186	1.00
19	7:34	1.00	0.822	1.00	10.00	10.00	0.469	0.469	1.00
20	8:09	1.00	0.885	1.00	10.00	10.00	1.531	1.531	1.00
21	8:56	1.00	0.970	1.00	10.00	10.00	0.166	0.166	1.00
22	9:33	1.00	1.036	1.00	10.00	10.00	0.516	0.516	1.00
23	9:49	1.00	1.066	1.00	10.00	10.00	0.869	0.869	1.00
24	10:00	1.00	1.086	1.00	10.00	10.00	0.924	0.924	1.00
25	14:08	1.00	1.168	1.00	10.00	10.00	0.127	0.127	1.00
26	15:52	1.00	1.313	1.00	10.00	10.00	0.376	0.376	1.00

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPHY**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND COPIED**



DATA: ST16980831B #1 SCANS 380 TO 440

CALI: ST16980831B #3

RIC+MASS CHROMATOGRAM

08/31/98 13:35:00

SAMPLE: 10UG/ML AP9 062598C

COND5.: INST. ID: F16

RANGE: G 1,2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

399  
856.  
3544.

856.



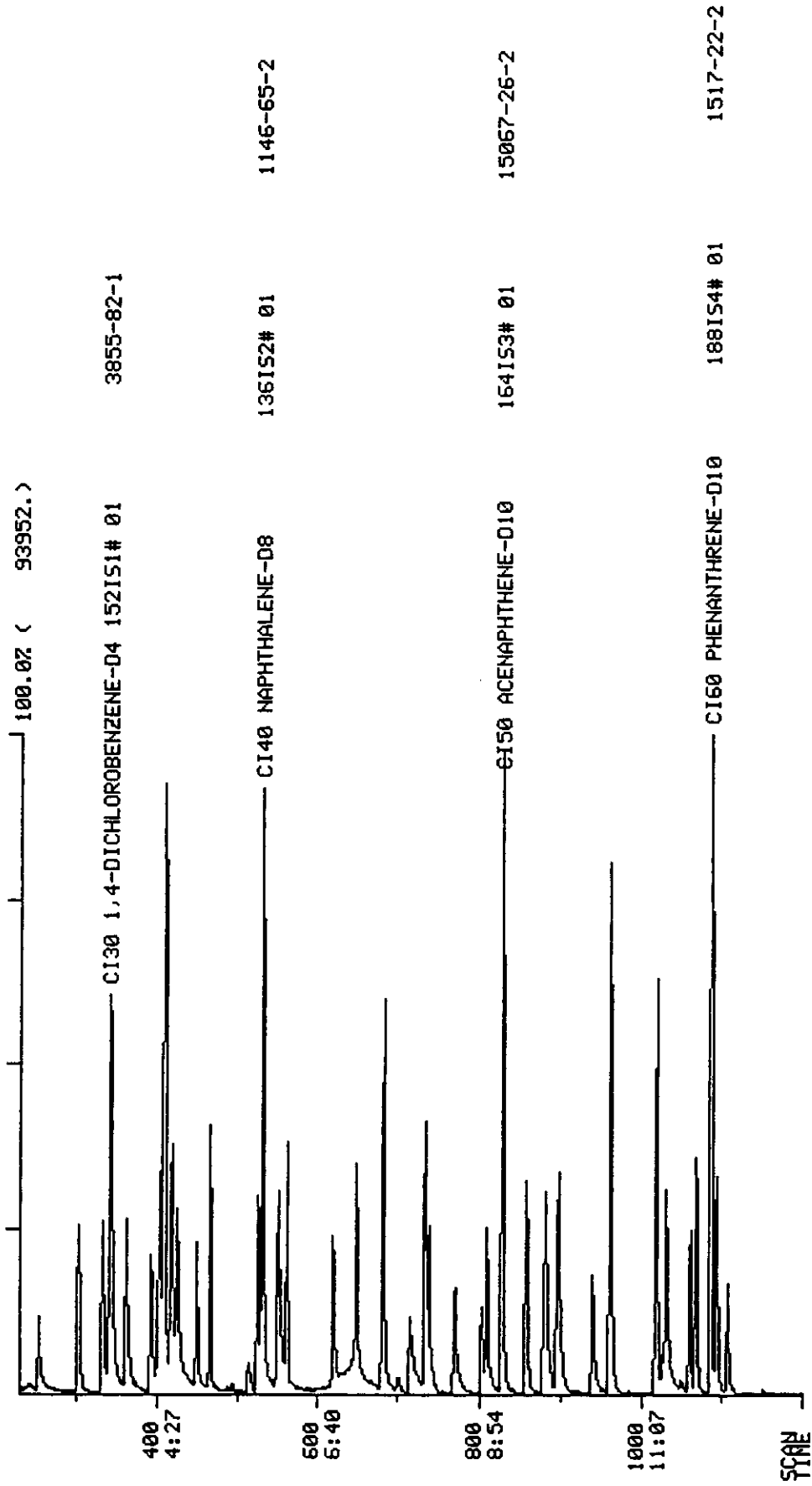
*81110-200*

RIC

1543

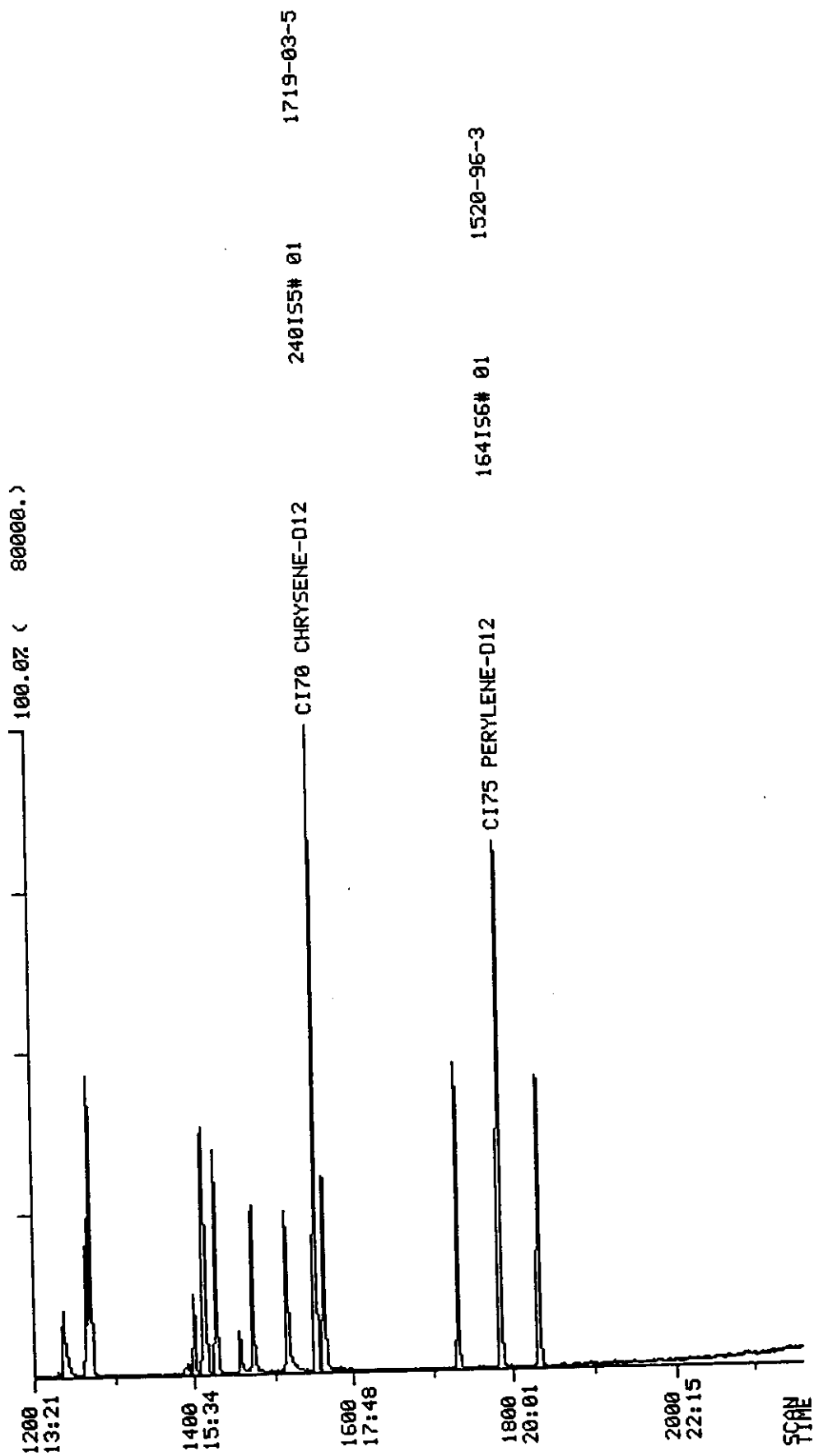
DATA FROM FILE: ST16980831C SCANS 230 TO 1200 ACQUIRED: 08/31/98 14:02:00  
CALL: ST16980831C #3

SAMPLE: 20UG/ML AP9 0625980  
CONDS.: INST. ID: F16



DATA FROM FILE: ST16980831C SCANS 1200 TO 2158 ACQUIRED: 08/31/98 14:02:00  
CALI: ST16980831C #3

SAMPLE: 20UG/ML AP9 0625980  
CONDS.: INST. ID: F16



Data: ST16980831C.TI

08/31/98 14:02:00

Sample: 20UG/ML AP9 062598D

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROSOPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALLATE	234 S4# 37	
40	4-AMINOBIIPHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	2SECBUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPYRILENE	58 S4#105	91-80-5
46	ISODRIN		
47	CHLOROBENZILATE	139 S4#120	

No	Name	185 S5# 20	140-57-8
48	ARAMITE (#1)	185 S5# 25	140-57-8
49	ARAMITE (#2)	120 S5# 30	60-11-7
50	P-DIMETHYLAMINOAZOBENZENE		

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Nght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	28379.	40.000 UG/ML	3.28
2	136	529	5:53	2	1.000	A BB	104398.	40.000 UG/ML	3.28
3	164	827	9:12	3	1.000	A BB	58510.	40.000 UG/ML	3.28
4	188	1086	12:05	4	1.000	A BB	99710.	40.000 UG/ML	3.28
5	240	1551	17:15	5	1.000	A BV	75805.	40.000 UG/ML	3.28
6	264	1782	19:49	6	1.000	A BB	63674.	40.000 UG/ML	3.28
7	79	103	1:09	1	0.303	A BB	22463.	20.000 UG/ML	1.64
8	93	145	1:37	1	0.426	A BB	22172.	20.000 UG/ML	1.64
9	42	156	1:44	1	0.459	A BB	10461.	20.000 UG/ML	1.64
10	80	181	2:01	1	0.532	A BB	9765.	20.000 UG/ML	1.64
11	102	216	2:24	1	0.635	A BB	7176.	20.000 UG/ML	1.64
12	79	253	2:49	1	0.744	A BB	14300.	20.000 UG/ML	1.64
13	117	301	3:21	1	0.885	A BB	8682.	20.000 UG/ML	1.64
14	105	402	4:28	1	1.182	A BB	28725.	20.000 UG/ML	1.64
15	100	400	4:27	1	1.176	qedt (1)	7882.	20.000 UG/ML	1.64
16	56	407	4:32	1	1.197	A BV	11884.	20.000 UG/ML	1.64
17	108	416	4:38	1	1.224	A BB	30227.	20.000 UG/ML	1.64
18	106	406	4:31	1	1.194	A BB	34950.	20.000 UG/ML	1.64
19	42	447	4:58	2	0.845	A BB	16262.	20.000 UG/ML	1.64
20	58	646	7:11	2	1.221	qedt (1)	82208.	20.000 UG/ML	1.64
21	162	551	6:08	2	1.042	A BB	12238.	20.000 UG/ML	1.64
22	213	548	6:06	2	1.036	A BB	10240.	20.000 UG/ML	1.64
23	108	656	7:18	2	1.240	qedt (2)	8496.	20.000 UG/ML	1.64
24	84	618	6:52	2	1.168	A BB	10715.	20.000 UG/ML	1.64
25	162	647	7:12	2	1.223	A BB	12848.	20.000 UG/ML	1.64
26	214	680	7:34	3	0.822	A BB	12617.	20.000 UG/ML	1.64
27	162	698	7:46	3	0.844	A BB	1336.	20.000 UG/ML	1.64
28	104	737	8:12	3	0.891	A BB	5880.	20.000 UG/ML	1.64
29	162	732	8:09	3	0.885	A BV	32113.	20.000 UG/ML	1.64
30	158	768	8:33	3	0.929	A BB	11800.	20.000 UG/ML	1.64
31	168	802	8:55	3	0.970	A BB	5576.	20.000 UG/ML	1.64
32	250	857	9:32	3	1.036	A BB	14750.	20.000 UG/ML	1.64
33	143	881	9:48	3	1.065	A BB	25121.	20.000 UG/ML	1.64
34	143	896	9:58	3	1.083	A VB	28484.	20.000 UG/ML	1.64
35	232	894	9:57	3	1.081	A BB	8117.	20.000 UG/ML	1.64
36	152	938	10:26	3	1.134	A BB	10100.	20.000 UG/ML	1.64
37	75	1025	11:24	4	0.944	A BB	7446.	20.000 UG/ML	1.64
38	108	1030	11:27	4	0.948	A BB	15198.	20.000 UG/ML	1.64
39	234	1017	11:19	4	0.936	A BB	4406.	20.000 UG/ML	1.64
40	169	1067	11:52	4	0.983	A BB	32077.	20.000 UG/ML	1.64
41	173	1093	12:09	4	1.006	A BB	15570.	20.000 UG/ML	1.64
42	237	1060	11:47	4	0.976	A BB	4036.	20.000 UG/ML	1.64
43	211	1107	12:19	4	1.019	A BB	6506.	20.000 UG/ML	1.64
44	190	1235	13:44	4	1.137	A BB	2912.	20.000 UG/ML	1.64
45	58	1264	14:04	4	1.164	A BB	21120.	20.000 UG/ML	1.64
46	193	1269	14:07	4	1.169	A BB	6234.	20.000 UG/ML	1.64
47	139	1426	15:52	4	1.313	A BB	17520.	20.000 UG/ML	1.64
48	185	1398	15:33	5	0.901	A BB	2154.	20.000 UG/ML	1.64
49	185	1413	15:43	5	0.911	A BB	3074.	20.000 UG/ML	1.64
50	120	1410	15:41	5	0.909	A BB	15810.	20.000 UG/ML	1.64

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:15	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:49	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	20.00	20.00	1.583	1.583	1.00
8	1:37	1.00	0.426	1.00	20.00	20.00	1.563	1.563	1.00
9	1:45	0.99	0.462	0.99	20.00	20.00	0.737	0.737	1.00
10	2:01	0.99	0.535	0.99	20.00	20.00	0.688	0.688	1.00
11	2:25	0.99	0.641	0.99	20.00	20.00	0.506	0.506	1.00
12	2:50	0.99	0.750	0.99	20.00	20.00	1.008	1.008	1.00
13	3:22	1.00	0.888	1.00	20.00	20.00	0.612	0.612	1.00
14	4:30	1.00	1.188	1.00	20.00	20.00	2.024	2.024	1.00
15	4:30	0.99	1.191	0.99	20.00	20.00	0.555	0.555	1.00
16	4:34	0.99	1.209	0.99	20.00	20.00	0.838	0.838	1.00
17	4:40	0.99	1.232	0.99	20.00	20.00	2.130	2.130	1.00
18	4:32	1.00	1.200	1.00	20.00	20.00	2.463	2.463	1.00
19	5:01	0.99	0.851	0.99	20.00	20.00	0.312	0.312	1.00
20	7:54	0.91	1.340	0.91	20.00	20.00	1.575	1.575	1.00
21	6:08	1.00	1.042	1.00	20.00	20.00	0.234	0.234	1.00
22	6:06	1.00	1.036	1.00	20.00	20.00	0.196	0.196	1.00
23	7:25	0.98	1.258	0.99	20.00	20.00	0.163	0.163	1.00
24	6:54	1.00	1.170	1.00	20.00	20.00	0.205	0.205	1.00
25	7:12	1.00	1.223	1.00	20.00	20.00	0.246	0.246	1.00
26	7:34	1.00	0.822	1.00	20.00	20.00	0.431	0.431	1.00
27	7:46	1.00	0.843	1.00	20.00	20.00	0.046	0.046	1.00
28	8:12	1.00	0.890	1.00	20.00	20.00	0.201	0.201	1.00
29	8:09	1.00	0.885	1.00	20.00	20.00	1.098	1.098	1.00
30	8:34	1.00	0.930	1.00	20.00	20.00	0.403	0.403	1.00
31	8:56	1.00	0.970	1.00	20.00	20.00	0.191	0.191	1.00
32	9:33	1.00	1.036	1.00	20.00	20.00	0.504	0.504	1.00
33	9:49	1.00	1.066	1.00	20.00	20.00	0.859	0.859	1.00
34	10:00	1.00	1.086	1.00	20.00	20.00	0.974	0.974	1.00
35	9:57	1.00	1.081	1.00	20.00	20.00	0.277	0.277	1.00
36	10:27	1.00	1.135	1.00	20.00	20.00	0.345	0.345	1.00
37	11:24	1.00	0.943	1.00	20.00	20.00	0.149	0.149	1.00
38	11:30	1.00	0.951	1.00	20.00	20.00	0.305	0.305	1.00
39	11:19	1.00	0.937	1.00	20.00	20.00	0.088	0.088	1.00
40	11:53	1.00	0.983	1.00	20.00	20.00	0.643	0.643	1.00
41	12:11	1.00	1.008	1.00	20.00	20.00	0.312	0.312	1.00
42	11:48	1.00	0.976	1.00	20.00	20.00	0.081	0.081	1.00
43	12:19	1.00	1.019	1.00	20.00	20.00	0.130	0.130	1.00
44	13:45	1.00	1.137	1.00	20.00	20.00	0.058	0.058	1.00
45	14:05	1.00	1.165	1.00	20.00	20.00	0.424	0.424	1.00
46	14:08	1.00	1.168	1.00	20.00	20.00	0.125	0.125	1.00
47	15:52	1.00	1.313	1.00	20.00	20.00	0.351	0.351	1.00
48	15:34	1.00	0.902	1.00	20.00	20.00	0.057	0.057	1.00
49	15:44	1.00	0.912	1.00	20.00	20.00	0.081	0.081	1.00
50	15:42	1.00	0.910	1.00	20.00	20.00	0.417	0.417	1.00

Quantitation Report File: ST16980831C

Data: ST16980831C.T1  
08/31/98 14:02:00  
Sample: 20UG/ML AP9 0625980  
Conds.: INST. ID: F16

Formula: 1UL INJ. Instrument: F16 Weight: 0.000  
Submitted by: QES Analyst: DAT Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
Resp. fac. from Library Entry

No	Name		
51	3,3'-DIMETHYLBENZIDINE	212 S5# 35	
52	KEPONE	272 S5# 37	
53	2-ACETYLAMINOFLUORENE	181 S5# 45	53-96-3
54	3-METHYLCHOLANTHRENE	268 S5# 85	56-49-5
55	7,12-DIMETHYLBENZANTHRACENE	256 S6# 20	75-97-6
56	HEXACHLOROPHENE	196 S6# 30	70-30-4

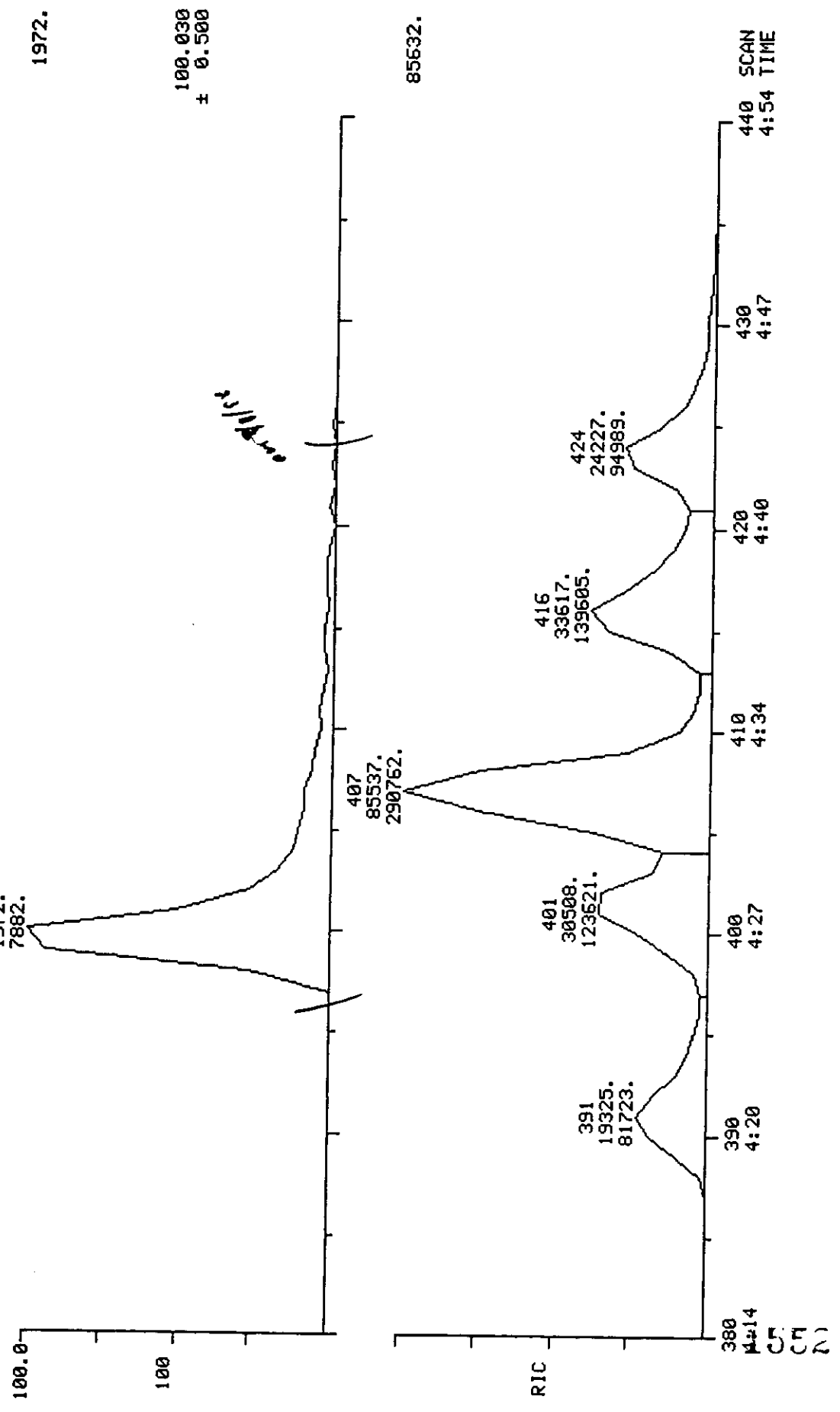
No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1472	16:22	5	0.949	A BB	19802.	20.000 UG/ML	1.64
52	272	1455	16:11	5	0.938	qedt	4676.	20.000 UG/ML	1.64
53	181	1514	16:50	5	0.976	A BB	15324.	20.000 UG/ML	1.64
54	268	1830	20:21	5	1.180	A BB	22226.	20.000 UG/ML	1.64
55	256	1729	19:14	6	0.970	A BB	19376.	20.000 UG/ML	1.64
56	NOT FOUND								

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:22	1.00	0.949	1.00	20.00	20.00	0.522	0.522	1.00
52	16:13	1.00	0.940	1.00	20.00	20.00	0.123	0.123	1.00
53	16:52	1.00	0.977	1.00	20.00	20.00	0.404	0.404	1.00
54	20:23	1.00	1.181	1.00	20.00	20.00	0.586	0.586	1.00
55	19:15	1.00	0.971	1.00	20.00	20.00	0.609	0.609	1.00
56	21:11		0.992						

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPHY**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND CODED.**

RIC+MASS CHROMATOGRAM  
 08/31/98 14:02:00  
 SAMPLE: 20UG/ML AP9 0625980  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 2, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

DATA: ST16980831C #1  
 CALI: ST16980831C #3





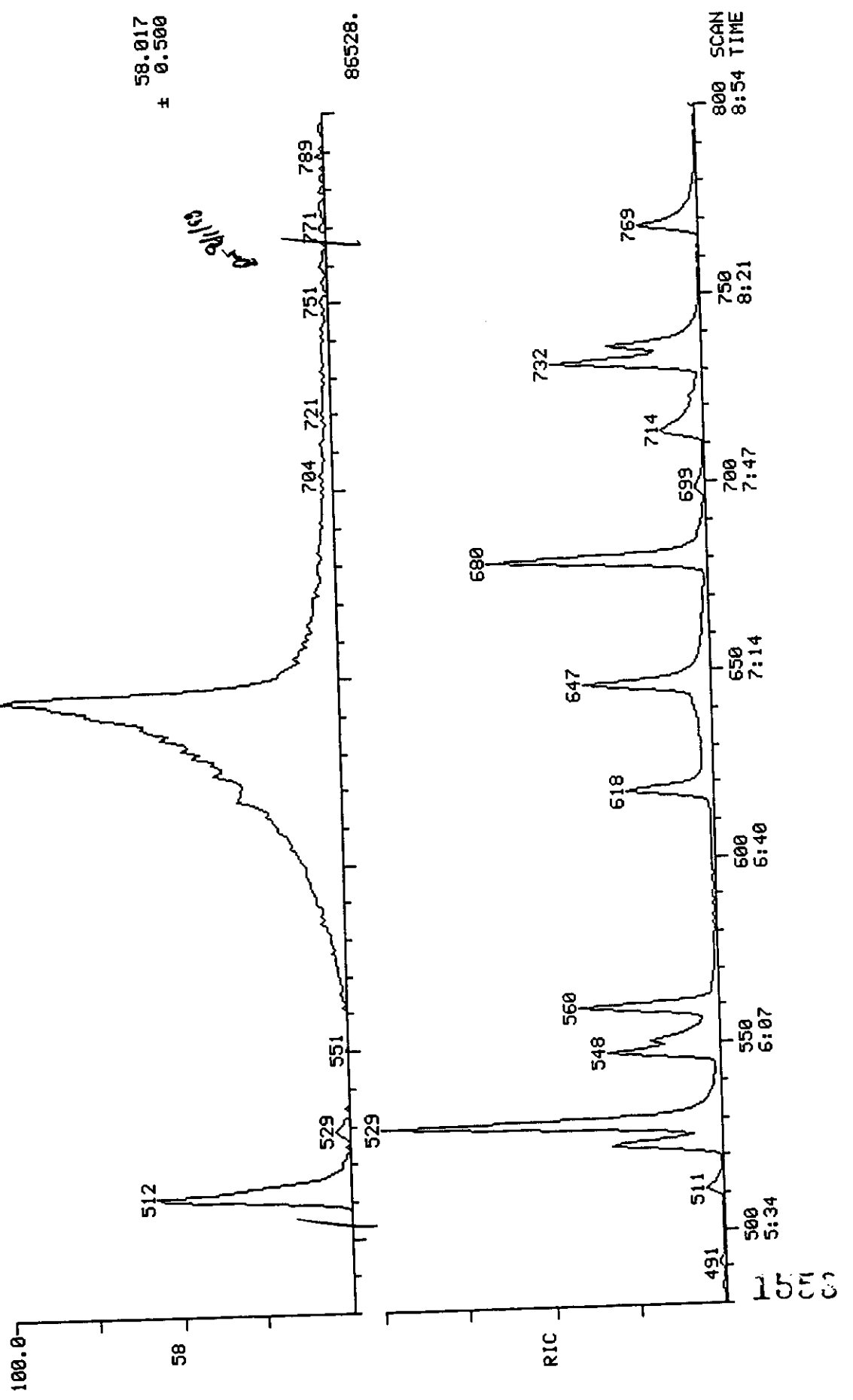
SCANS 480 TO 800

DATA: ST16980831C #1  
CALI: ST16980831C #3

RIC+MASS CHROMATOGRAM

08/31/98 14:02:00  
SAMPLE: 20UG/ML AP9 062598D  
CONDS.: INST. ID: F16  
RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

2796.



RIC+MASS CHROMATOGRAM  
08/31/98 14:02:00  
SAMPLE: 20UG/ML AP9 062598D  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

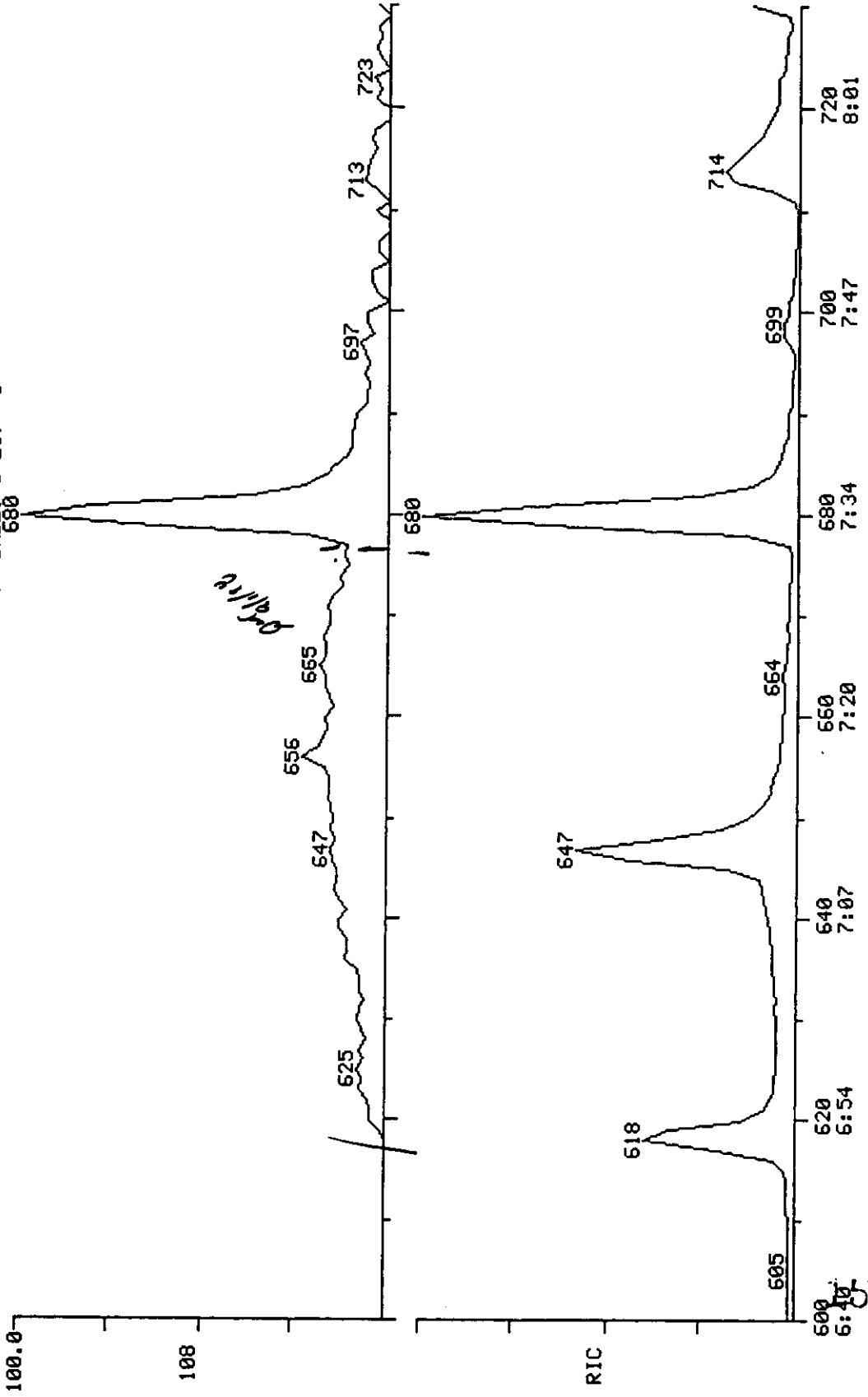
SCANS 600 TO 730

DATA: ST16980831C #1  
CALI: ST16980831C #3

1230.

108.032  
± 0.500

56384.



SCAN  
TIME

720  
8:01

700  
7:47

680  
7:34

660  
7:20

640  
7:07

620  
6:54

600  
6:40

CTI  
MS

SCANS 1430 TO 1550

DATA: ST16980831C #1  
CALI: ST16980831C #3

RIC+MASS CHROMATOGRAM

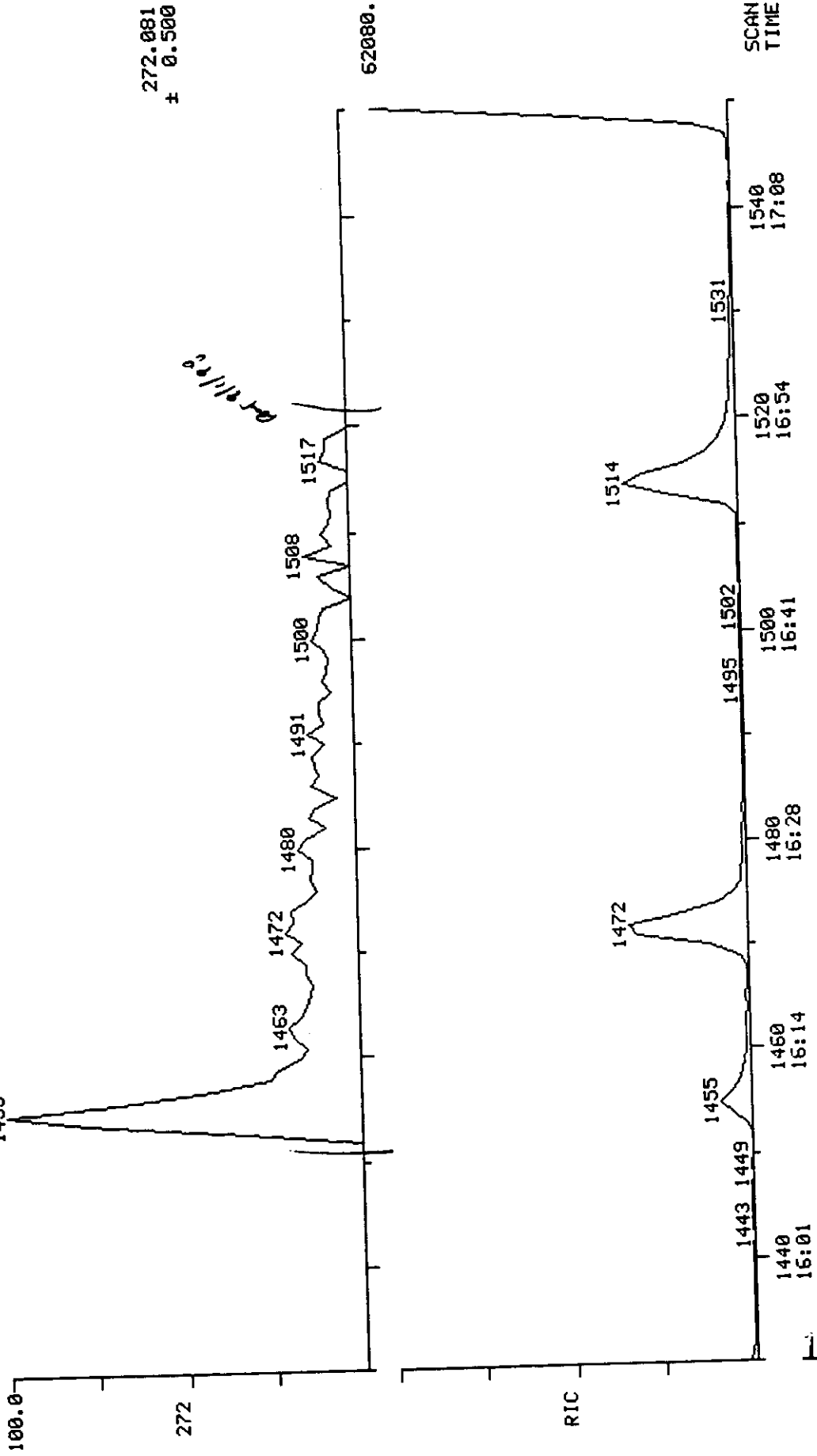
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SAMPLE: 20UG/ML AP9 062598D

CONDS.: INST. ID: F16

RANGE: G 1.2158 LABEL: N 0. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20, 3

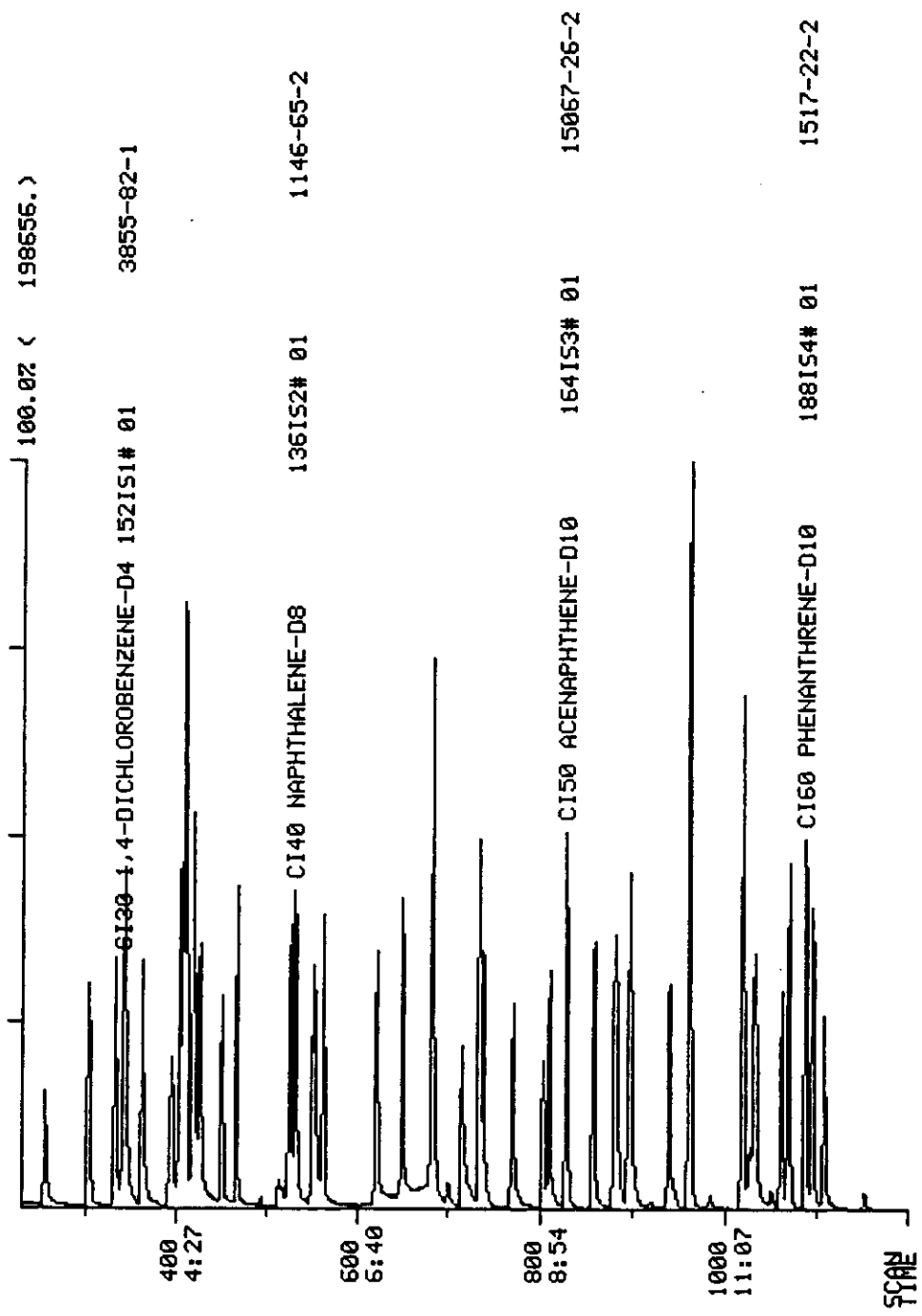
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1550

DATA FROM FILE: ST16980831D SCANS 230 TO 1200 ACQUIRED: 08/31/98 14:32:00  
CALI: ST16980831D #3

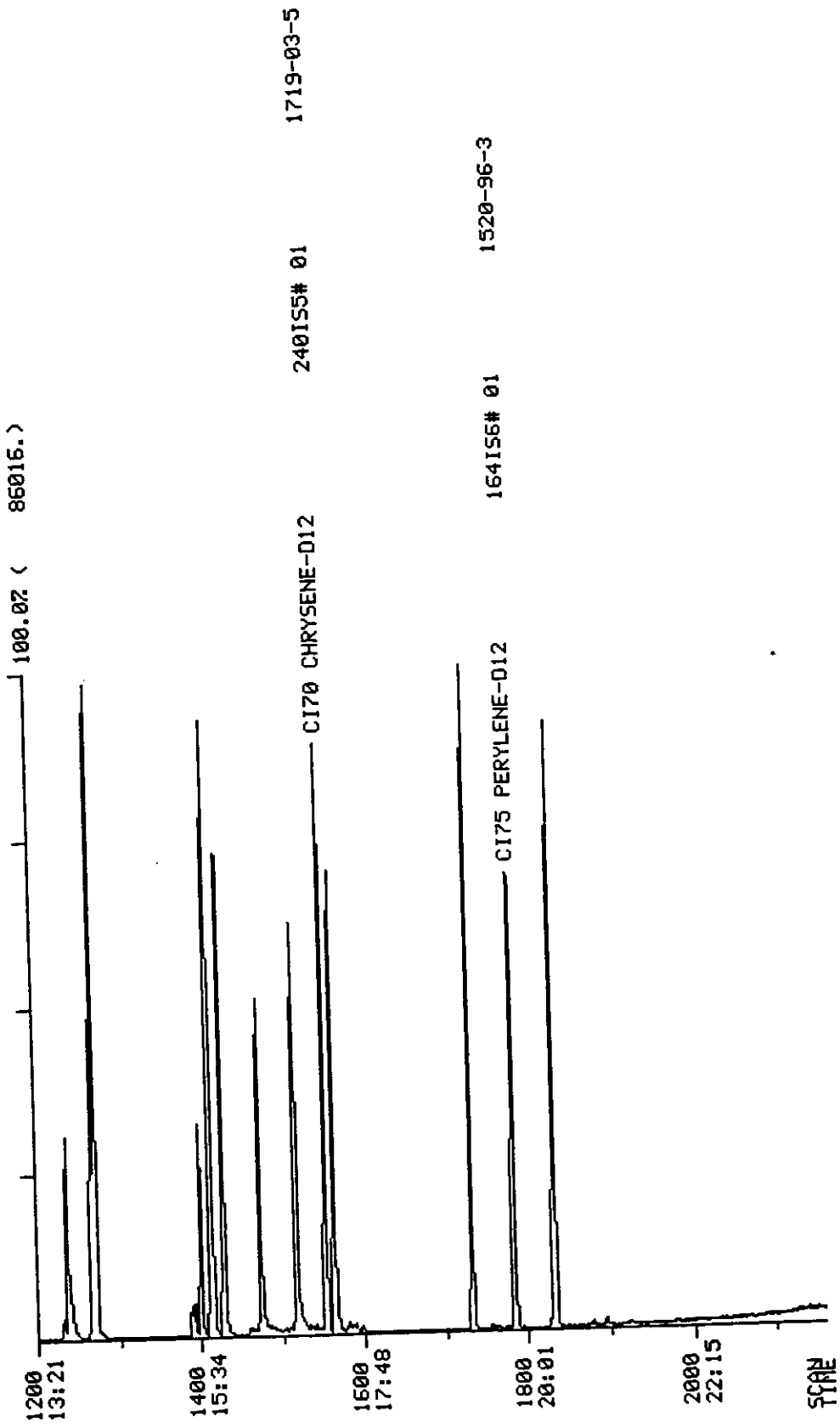
SAMPLE: 50UG/ML AP9 062598E  
CONDS.: INST. ID: F16



1558

DATA FROM FILE: ST16980831D SCANS 1200 TO 2158 ACQUIRED: 08/31/98 14:32:00  
CALI: ST16980831D #3

SAMPLE: 50UG/ML AP9 062598E  
CONDS.: INST. ID: F16



Data: ST16980831D.T1

08/31/98 14:32:00

Sample: 50UG/ML AP9 062598E

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
1	CI30 1,4-DICHLORO BENZENE-D4	152I S1# 01	3855-82-1
2	CI40 NAPHTHALENE-D8	136I S2# 01	1146-65-2
3	CI50 ACENAPHTHENE-D10	164I S3# 01	15067-26-2
4	CI60 PHENANTHRENE-D10	188I S4# 01	1517-22-2
5	CI70 CHRYSENE-D12	240I S5# 01	1719-03-5
6	CI75 PERYLENE-D12	164I S6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROSOPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSOI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALATE	234 S4# 37	
40	4-AMINOBI PHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	2SEC BUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPYRILENE	58 S4#105	91-80-5
46	ISODRIN		
47	CHLOROBENZILATE	139 S4#120	

No	Name		
48	ARAMITE (#1)	185 S5# 20	140-57-8
49	ARAMITE (#2)	185 S5# 25	140-57-8
50	P-DIMETHYLAMINOAZOBENZENE	120 S5# 30	60-11-7

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	28687.	40.000 UG/ML	1.49
2	136	529	5:53	2	1.000	A BB	104087.	40.000 UG/ML	1.49
3	164	828	9:13	3	1.000	A BV	58806.	40.000 UG/ML	1.49
4	188	1086	12:05	4	1.000	A BB	103078.	40.000 UG/ML	1.49
5	240	1551	17:15	5	1.000	A VB	69225.	40.000 UG/ML	1.49
6	264	1782	19:49	6	1.000	A BB	57648.	40.000 UG/ML	1.49
7	79	103	1:09	1	0.303	A BB	56577.	50.000 UG/ML	1.86
8	93	145	1:37	1	0.426	A BB	57154.	50.000 UG/ML	1.86
9	42	157	1:45	1	0.462	A BB	29520.	50.000 UG/ML	1.86
10	80	182	2:01	1	0.535	A BB	24557.	50.000 UG/ML	1.86
11	102	217	2:25	1	0.638	A BB	18688.	50.000 UG/ML	1.86
12	79	254	2:50	1	0.747	A BB	40170.	50.000 UG/ML	1.86
13	117	302	3:22	1	0.888	A BB	21710.	50.000 UG/ML	1.86
14	105	403	4:29	1	1.185	A BV	70275.	50.000 UG/ML	1.86
15	100	402	4:28	1	1.182	A BB	20603.	50.000 UG/ML	1.86
16	56	409	4:33	1	1.203	A BB	29437.	50.000 UG/ML	1.86
17	108	417	4:38	1	1.226	A BB	83872.	50.000 UG/ML	1.86
18	106	407	4:32	1	1.197	A BB	79912.	50.000 UG/ML	1.86
19	42	449	5:00	2	0.849	A BB	42444.	50.000 UG/ML	1.86
20	58	685	7:37	2	1.295	qedt (1)	176012.	50.000 UG/ML	1.86
21	162	551	6:08	2	1.042	A BB	34375.	50.000 UG/ML	1.86
22	213	548	6:06	2	1.036	A BB	26126.	50.000 UG/ML	1.86
23	108	618	6:52	2	1.168	qedt (1)	21776.	50.000 UG/ML	1.86
24	84	619	6:53	2	1.170	A BB	27751.	50.000 UG/ML	1.86
25	162	647	7:12	2	1.223	A BB	32194.	50.000 UG/ML	1.86
26	214	680	7:34	3	0.821	A BB	31894.	50.000 UG/ML	1.86
27	162	698	7:46	3	0.843	A BB	3066.	50.000 UG/ML	1.86
28	104	737	8:12	3	0.890	A BB	14360.	50.000 UG/ML	1.86
29	162	732	8:09	3	0.884	A BV	82983.	50.000 UG/ML	1.86
30	158	769	8:33	3	0.929	A BB	31543.	50.000 UG/ML	1.86
31	168	802	8:55	3	0.969	A BB	14862.	50.000 UG/ML	1.86
32	250	857	9:32	3	1.035	A BB	35528.	50.000 UG/ML	1.86
33	143	882	9:49	3	1.065	A BB	65362.	50.000 UG/ML	1.86
34	143	897	9:59	3	1.083	A BB	76505.	50.000 UG/ML	1.86
35	232	895	9:57	3	1.081	A BB	22189.	50.000 UG/ML	1.86
36	152	938	10:26	3	1.133	A BB	27016.	50.000 UG/ML	1.86
37	75	1025	11:24	4	0.944	A BB	20893.	50.000 UG/ML	1.86
38	108	1032	11:29	4	0.950	A BB	41692.	50.000 UG/ML	1.86
39	234	1017	11:19	4	0.936	A BB	11272.	50.000 UG/ML	1.86
40	169	1068	11:53	4	0.983	A VB	85529.	50.000 UG/ML	1.86
41	173	1094	12:10	4	1.007	A BB	40637.	50.000 UG/ML	1.86
42	237	1060	11:47	4	0.976	A BB	10186.	50.000 UG/ML	1.86
43	211	1107	12:19	4	1.019	A BV	21220.	50.000 UG/ML	1.86
44	190	1235	13:44	4	1.137	A BB	9477.	50.000 UG/ML	1.86
45	58	1265	14:04	4	1.165	A BB	43644.	50.000 UG/ML	1.86
46	193	1270	14:08	4	1.169	A BB	16648.	50.000 UG/ML	1.86
47	139	1427	15:52	4	1.314	A BB	42346.	50.000 UG/ML	1.86
48	185	1399	15:34	5	0.902	A VB	5450.	50.000 UG/ML	1.86
49	185	1413	15:43	5	0.911	A BB	7782.	50.000 UG/ML	1.86
50	120	1411	15:42	5	0.910	A BB	36805.	50.000 UG/ML	1.86

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:15	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:49	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	50.00	50.00	1.578	1.578	1.00
8	1:37	1.00	0.426	1.00	50.00	50.00	1.594	1.594	1.00
9	1:45	1.00	0.462	1.00	50.00	50.00	0.823	0.823	1.00
10	2:01	1.00	0.535	1.00	50.00	50.00	0.685	0.685	1.00
11	2:25	1.00	0.641	1.00	50.00	50.00	0.521	0.521	1.00
12	2:50	1.00	0.750	1.00	50.00	50.00	1.120	1.120	1.00
13	3:22	1.00	0.888	1.00	50.00	50.00	0.605	0.605	1.00
14	4:30	1.00	1.188	1.00	50.00	50.00	1.960	1.960	1.00
15	4:30	0.99	1.191	0.99	50.00	50.00	0.575	0.575	1.00
16	4:34	1.00	1.209	1.00	50.00	50.00	0.821	0.821	1.00
17	4:40	1.00	1.232	1.00	50.00	50.00	2.339	2.339	1.00
18	4:32	1.00	1.200	1.00	50.00	50.00	2.228	2.228	1.00
19	5:01	1.00	0.851	1.00	50.00	50.00	0.326	0.326	1.00
20	7:54	0.96	1.340	0.97	50.00	50.00	1.353	1.353	1.00
21	6:08	1.00	1.042	1.00	50.00	50.00	0.264	0.264	1.00
22	6:06	1.00	1.036	1.00	50.00	50.00	0.201	0.201	1.00
23	7:25	0.93	1.258	0.93	50.00	50.00	0.167	0.167	1.00
24	6:54	1.00	1.170	1.00	50.00	50.00	0.213	0.213	1.00
25	7:12	1.00	1.223	1.00	50.00	50.00	0.247	0.247	1.00
26	7:34	1.00	0.822	1.00	50.00	50.00	0.434	0.434	1.00
27	7:46	1.00	0.843	1.00	50.00	50.00	0.042	0.042	1.00
28	8:12	1.00	0.890	1.00	50.00	50.00	0.195	0.195	1.00
29	8:09	1.00	0.885	1.00	50.00	50.00	1.129	1.129	1.00
30	8:34	1.00	0.930	1.00	50.00	50.00	0.429	0.429	1.00
31	8:56	1.00	0.970	1.00	50.00	50.00	0.202	0.202	1.00
32	9:33	1.00	1.036	1.00	50.00	50.00	0.483	0.483	1.00
33	9:49	1.00	1.066	1.00	50.00	50.00	0.889	0.889	1.00
34	10:00	1.00	1.086	1.00	50.00	50.00	1.041	1.041	1.00
35	9:57	1.00	1.081	1.00	50.00	50.00	0.302	0.302	1.00
36	10:27	1.00	1.135	1.00	50.00	50.00	0.368	0.368	1.00
37	11:24	1.00	0.943	1.00	50.00	50.00	0.162	0.162	1.00
38	11:30	1.00	0.951	1.00	50.00	50.00	0.324	0.324	1.00
39	11:19	1.00	0.937	1.00	50.00	50.00	0.087	0.087	1.00
40	11:53	1.00	0.983	1.00	50.00	50.00	0.664	0.664	1.00
41	12:11	1.00	1.008	1.00	50.00	50.00	0.315	0.315	1.00
42	11:48	1.00	0.976	1.00	50.00	50.00	0.079	0.079	1.00
43	12:19	1.00	1.019	1.00	50.00	50.00	0.165	0.165	1.00
44	13:45	1.00	1.137	1.00	50.00	50.00	0.074	0.074	1.00
45	14:05	1.00	1.165	1.00	50.00	50.00	0.339	0.339	1.00
46	14:08	1.00	1.168	1.00	50.00	50.00	0.129	0.129	1.00
47	15:52	1.00	1.313	1.00	50.00	50.00	0.329	0.329	1.00
48	15:34	1.00	0.902	1.00	50.00	50.00	0.063	0.063	1.00
49	15:44	1.00	0.912	1.00	50.00	50.00	0.090	0.090	1.00
50	15:42	1.00	0.910	1.00	50.00	50.00	0.425	0.425	1.00



Quantitation Report File: ST16980831D

Data: ST16980831D.T1

08/31/98 14:32:00

Sample: 50UG/ML AP9 062598E

Conds.: INST. ID: F16

Formula: 1UL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name		
51	3,3'-DIMETHYLBENZIDINE	212 S5# 35	
52	KEPONE	272 S5# 37	
53	2-ACETYLAMINOFLUORENE	181 S5# 45	53-96-3
54	3-METHYLCHOLANTHRENE	268 S5# 85	56-49-5
55	7,12-DIMETHYLBENZANTHRACENE	256 S6# 20	75-97-6
56	HEXACHLOROPHENE	196 S6# 30	70-30-4

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1472	16:22	5	0.949	A BB	38968.	50.000 UG/ML	1.86
52	272	1472	16:22	5	0.949	qedt(2)	7520.	50.000 UG/ML	1.86
53	181	1515	16:51	5	0.977	A BB	38918.	50.000 UG/ML	1.86
54	268	1831	20:22	5	1.181	A BB	49854.	50.000 UG/ML	1.86
55	256	1730	19:15	6	0.971	A BB	44288.	50.000 UG/ML	1.86
56									

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:22	1.00	0.949	1.00	50.00	50.00	0.450	0.450	1.00
52	16:13	1.01	0.940	1.01	50.00	50.00	0.087	0.087	1.00
53	16:52	1.00	0.977	1.00	50.00	50.00	0.450	0.450	1.00
54	20:23	1.00	1.181	1.00	50.00	50.00	0.576	0.576	1.00
55	19:15	1.00	0.971	1.00	50.00	50.00	0.615	0.615	1.00
56	21:11		0.992						

MANUAL EDIT CODES  
1. PEAK NOT FOUND  
2. POOR CHROMATOGRAPHY  
3. WRONG ISOMER  
ALL MANUAL EDITS MUST BE  
INITIALED, DATED, AND CODED

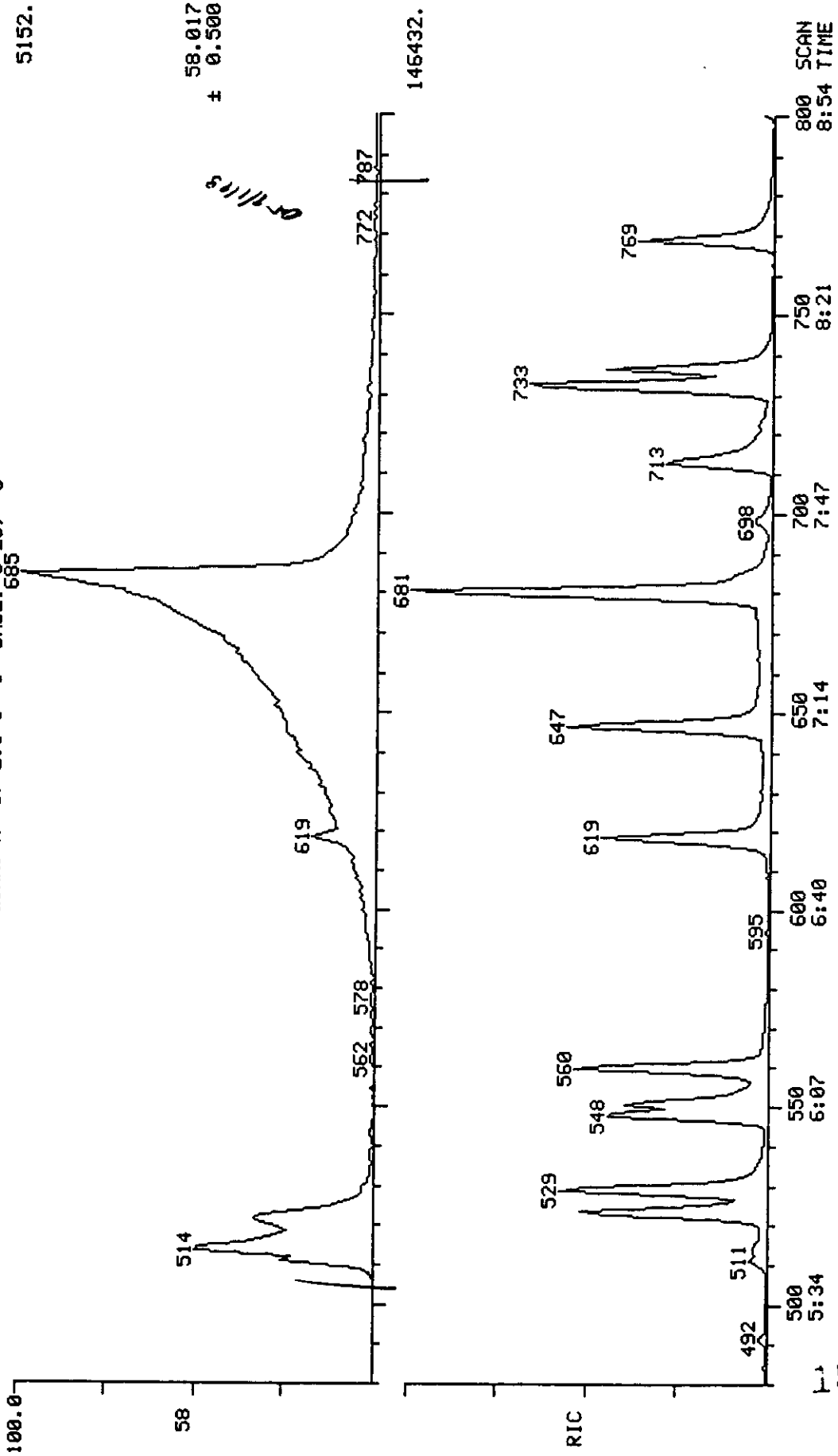
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1581  
1581

RIC+MASS CHROMATOGRAM

08/31/98 14:32:00  
SAMPLE: 50UG/ML AP9 062598E  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20. 3

DATA: ST16980831D #1  
CALI: ST16980831D #3  
SCANS 480 TO 800

5152.



1562

RIC+MASS CHROMATOGRAM DATA: ST16980831D #1 SCANS 590 TO 720

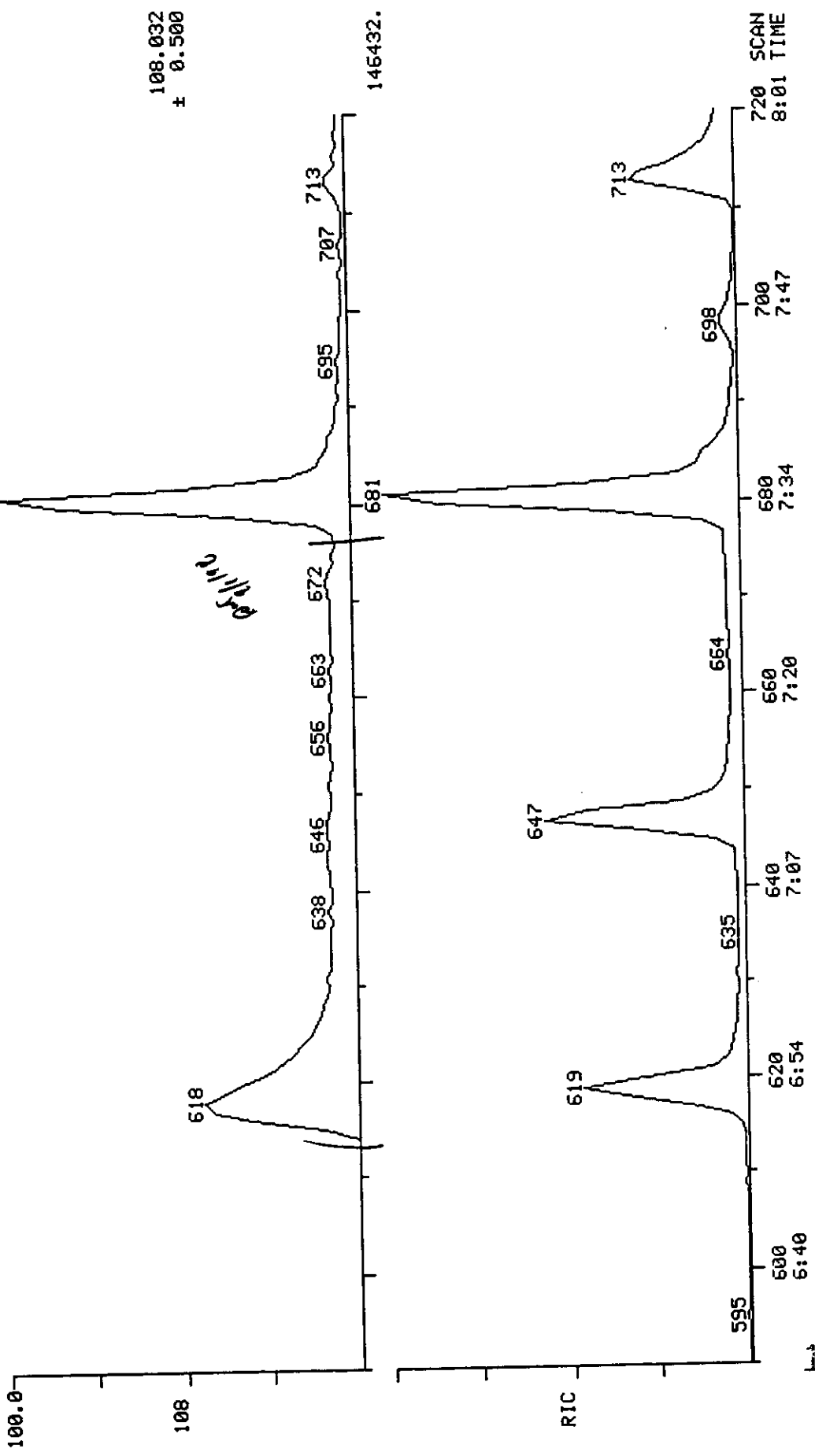
08/31/98 14:32:00 CALLI: ST16980831D #3

SAMPLE: 50UG/ML AP9 062598E

CONDS.: INST. ID: F15

RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 681

3304.



1503

RIC+MASS CHROMATOGRAM

08/31/98 14:32:00

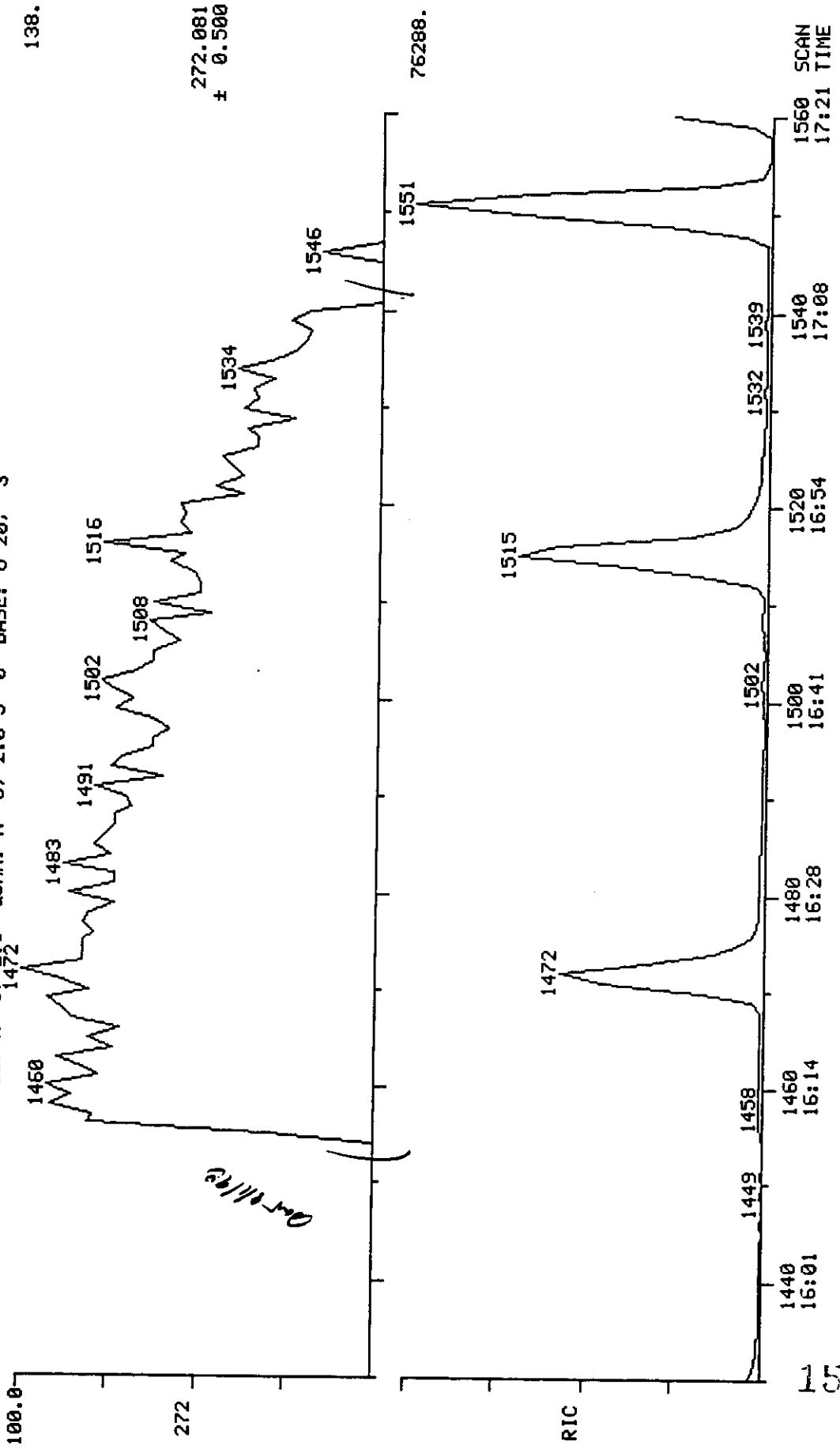
SAMPLE: 50UG/ML AP9 062598E

CONDS.: INST. ID: F16

RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

DATA: ST16980831D #1  
CALI: ST16980831D #3

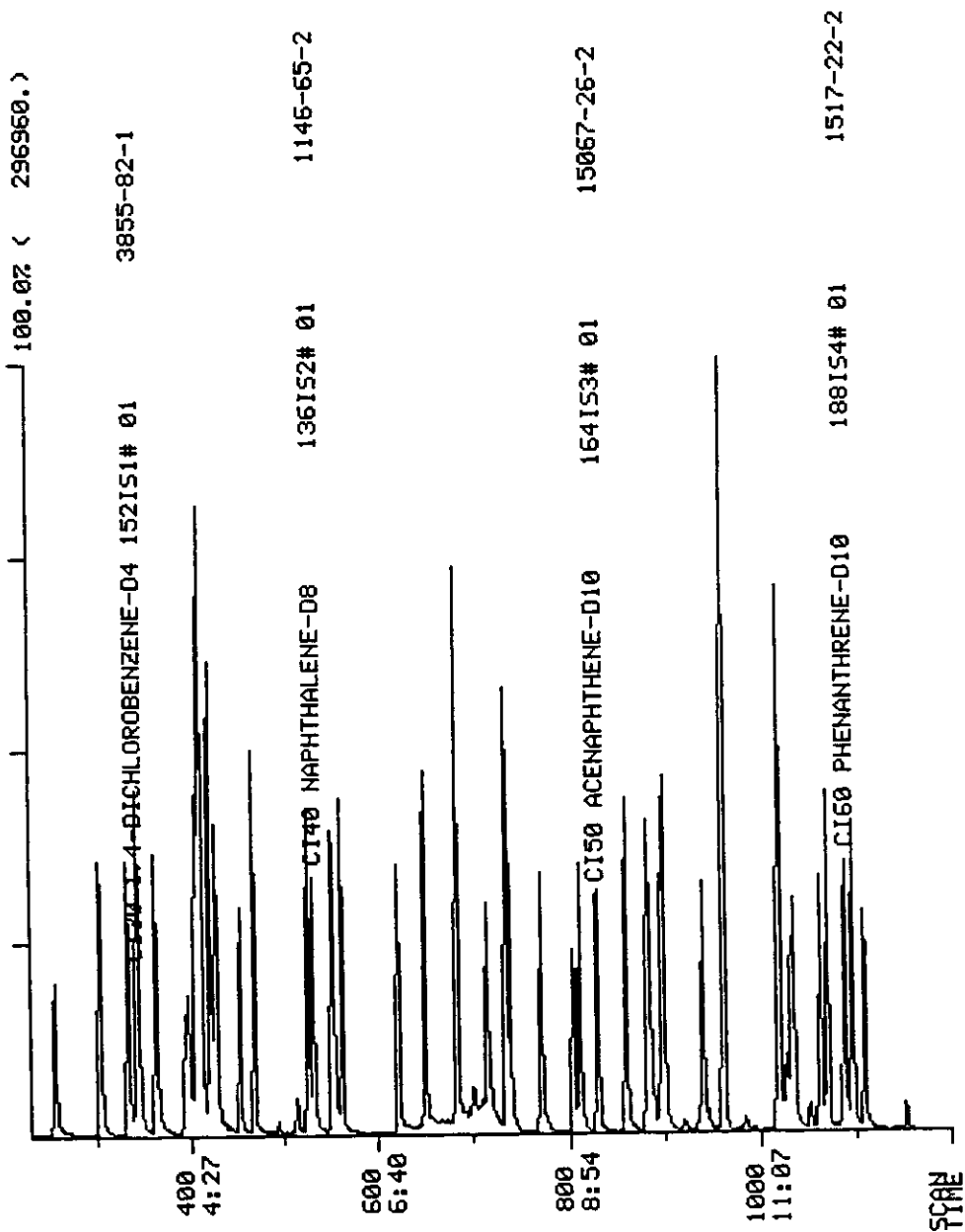
SCANS 1430 TO 1560



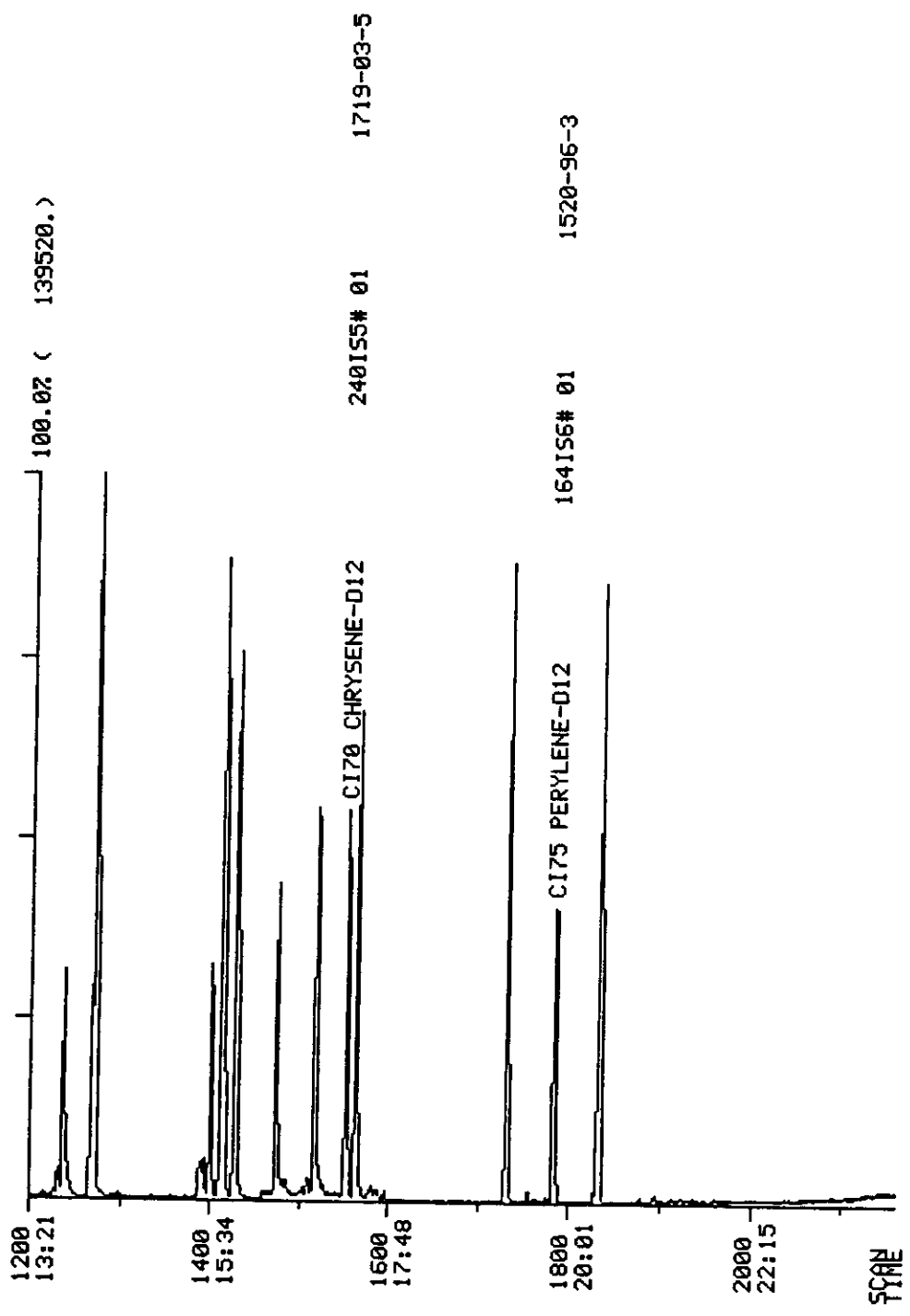
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DATA FROM FILE: ST16980831A SCANS 230 TO 1200 ACQUIRED: 08/31/98 13:07:00  
CALI: ST16980831A #3

SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16



DATA FROM FILE: ST16980831A SCANS 1200 TO 2158 ACQUIRED: 08/31/98 13:07:00  
CALI: ST16980831A #3  
SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16



1550

Data: ST16980831A.TI

08/31/98 13:07:00

Sample: 80UG/ML AP9 062598F

Conds.: INST. ID: F16

Formula: 1UL INJ.

Submitted by: QES

Instrument: F16

Analyst: DAT

Weight: 0.000

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROSOPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALATE	234 S4# 37	
40	4-AMINOBIPHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	2SECBUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPRYLENE	58 S4#105	91-80-5
46	ISODRIN		
47	CHLOROBENZILATE	139 S4#120	

No	Name		
48	ARAMITE (#1)	185 S5# 20	140-57-8
49	ARAMITE (#2)	185 S5# 25	140-57-8
50	P-DIMETHYLAMINOAZOBENZENE	120 S5# 30	60-11-7

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	29680.	40.000 UG/ML	0.96
2	136	530	5:54	2	1.000	A BB	108356.	40.000 UG/ML	0.96
3	164	828	9:13	3	1.000	A BB	60104.	40.000 UG/ML	0.96
4	188	1087	12:05	4	1.000	A BB	107988.	40.000 UG/ML	0.96
5	240	1551	17:15	5	1.000	A BV	69686.	40.000 UG/ML	0.96
6	264	1782	19:49	6	1.000	A BB	54786.	40.000 UG/ML	0.96
7	79	103	1:09	1	0.303	A BB	94198.	80.000 UG/ML	1.92
8	93	145	1:37	1	0.426	A BB	93595.	80.000 UG/ML	1.92
9	42	157	1:45	1	0.462	A BB	51975.	80.000 UG/ML	1.92
10	80	182	2:01	1	0.535	A BB	35745.	80.000 UG/ML	1.92
11	102	218	2:25	1	0.641	A BB	32568.	80.000 UG/ML	1.92
12	79	255	2:50	1	0.750	A BB	69355.	80.000 UG/ML	1.92
13	117	302	3:22	1	0.888	A BB	35884.	80.000 UG/ML	1.92
14	105	404	4:30	1	1.188	A BB	115427.	80.000 UG/ML	1.92
15	100	405	4:30	1	1.191	A BB	33375.	80.000 UG/ML	1.92
16	56	411	4:34	1	1.209	A BV	47803.	80.000 UG/ML	1.92
17	108	419	4:40	1	1.232	A BB	142918.	80.000 UG/ML	1.92
18	106	408	4:32	1	1.200	A BB	126404.	80.000 UG/ML	1.92
19	42	451	5:01	2	0.851	A BV	70838.	80.000 UG/ML	1.92
20	58	710	7:54	2	1.340	qedt	224618.	80.000 UG/ML	1.92
21	162	552	6:08	2	1.042	A BB	60603.	80.000 UG/ML	1.92
22	213	549	6:06	2	1.036	A BB	42724.	80.000 UG/ML	1.92
23	108	667	7:25	2	1.258	qedt	8144.	80.000 UG/ML	1.92
24	84	620	6:54	2	1.170	A BB	48711.	80.000 UG/ML	1.92
25	162	648	7:12	2	1.223	A BB	54270.	80.000 UG/ML	1.92
26	214	681	7:34	3	0.822	A BB	51722.	80.000 UG/ML	1.92
27	162	698	7:46	3	0.843	A BB	5166.	80.000 UG/ML	1.92
28	104	737	8:12	3	0.890	A BB	23560.	80.000 UG/ML	1.92
29	162	733	8:09	3	0.885	A BB	183709.	80.000 UG/ML	1.92
30	158	770	8:34	3	0.930	A BB	53790.	80.000 UG/ML	1.92
31	168	803	8:56	3	0.970	A BB	25287.	80.000 UG/ML	1.92
32	250	858	9:33	3	1.036	A BB	59261.	80.000 UG/ML	1.92
33	143	883	9:49	3	1.066	A BB	91990.	80.000 UG/ML	1.92
34	143	899	10:00	3	1.086	A BB	119758.	80.000 UG/ML	1.92
35	232	895	9:57	3	1.081	A BB	37212.	80.000 UG/ML	1.92
36	152	940	10:27	3	1.135	A BB	45826.	80.000 UG/ML	1.92
37	75	1025	11:24	4	0.943	A VB	37064.	80.000 UG/ML	1.92
38	108	1034	11:30	4	0.951	A BB	68243.	80.000 UG/ML	1.92
39	234	1018	11:19	4	0.937	A BV	19166.	80.000 UG/ML	1.92
40	169	1069	11:53	4	0.983	A BB	130115.	80.000 UG/ML	1.92
41	173	1096	12:11	4	1.008	A BB	65067.	80.000 UG/ML	1.92
42	237	1061	11:48	4	0.976	A BB	17138.	80.000 UG/ML	1.92
43	211	1108	12:19	4	1.019	A VB	35264.	80.000 UG/ML	1.92
44	190	1236	13:45	4	1.137	A BB	16496.	80.000 UG/ML	1.92
45	58	1266	14:05	4	1.165	A BB	44262.	80.000 UG/ML	1.92
46	193	1270	14:08	4	1.168	A BV	27606.	80.000 UG/ML	1.92
47	139	1427	15:52	4	1.313	A BB	66594.	80.000 UG/ML	1.92
48	185	1399	15:34	5	0.902	A BB	8471.	80.000 UG/ML	1.92
49	185	1414	15:44	5	0.912	A BB	12567.	80.000 UG/ML	1.92
50	120	1412	15:42	5	0.910	A VB	58473.	80.000 UG/ML	1.92



No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:15	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:49	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	80.00	80.00	1.587	1.587	1.00
8	1:37	1.00	0.426	1.00	80.00	80.00	1.577	1.577	1.00
9	1:45	1.00	0.462	1.00	80.00	80.00	0.876	0.876	1.00
10	2:01	1.00	0.535	1.00	80.00	80.00	0.602	0.602	1.00
11	2:25	1.00	0.641	1.00	80.00	80.00	0.549	0.549	1.00
12	2:50	1.00	0.750	1.00	80.00	80.00	1.168	1.168	1.00
13	3:22	1.00	0.888	1.00	80.00	80.00	0.605	0.605	1.00
14	4:30	1.00	1.188	1.00	80.00	80.00	1.945	1.945	1.00
15	4:30	1.00	1.191	1.00	80.00	80.00	0.562	0.562	1.00
16	4:34	1.00	1.209	1.00	80.00	80.00	0.805	0.805	1.00
17	4:40	1.00	1.232	1.00	80.00	80.00	2.408	2.408	1.00
18	4:32	1.00	1.200	1.00	80.00	80.00	2.129	2.129	1.00
19	5:01	1.00	0.851	1.00	80.00	80.00	0.327	0.327	1.00
20	7:54	1.00	1.340	1.00	80.00	80.00	1.036	1.036	1.00
21	6:08	1.00	1.042	1.00	80.00	80.00	0.280	0.280	1.00
22	6:06	1.00	1.036	1.00	80.00	80.00	0.197	0.197	1.00
23	7:25	1.00	1.258	1.00	80.00	80.00	0.038	0.038	1.00
24	6:54	1.00	1.170	1.00	80.00	80.00	0.225	0.225	1.00
25	7:12	1.00	1.223	1.00	80.00	80.00	0.250	0.250	1.00
26	7:34	1.00	0.822	1.00	80.00	80.00	0.430	0.430	1.00
27	7:46	1.00	0.843	1.00	80.00	80.00	0.043	0.043	1.00
28	8:12	1.00	0.890	1.00	80.00	80.00	0.196	0.196	1.00
29	8:09	1.00	0.885	1.00	80.00	80.00	1.528	1.528	1.00
30	8:34	1.00	0.930	1.00	80.00	80.00	0.447	0.447	1.00
31	8:56	1.00	0.970	1.00	80.00	80.00	0.210	0.210	1.00
32	9:33	1.00	1.036	1.00	80.00	80.00	0.493	0.493	1.00
33	9:49	1.00	1.066	1.00	80.00	80.00	0.765	0.765	1.00
34	10:00	1.00	1.086	1.00	80.00	80.00	0.996	0.996	1.00
35	9:57	1.00	1.081	1.00	80.00	80.00	0.310	0.310	1.00
36	10:27	1.00	1.135	1.00	80.00	80.00	0.381	0.381	1.00
37	11:24	1.00	0.943	1.00	80.00	80.00	0.172	0.172	1.00
38	11:30	1.00	0.951	1.00	80.00	80.00	0.316	0.316	1.00
39	11:19	1.00	0.937	1.00	80.00	80.00	0.089	0.089	1.00
40	11:53	1.00	0.983	1.00	80.00	80.00	0.602	0.602	1.00
41	12:11	1.00	1.008	1.00	80.00	80.00	0.301	0.301	1.00
42	11:48	1.00	0.976	1.00	80.00	80.00	0.079	0.079	1.00
43	12:19	1.00	1.019	1.00	80.00	80.00	0.163	0.163	1.00
44	13:45	1.00	1.137	1.00	80.00	80.00	0.076	0.076	1.00
45	14:05	1.00	1.165	1.00	80.00	80.00	0.205	0.205	1.00
46	14:08	1.00	1.168	1.00	80.00	80.00	0.128	0.128	1.00
47	15:52	1.00	1.313	1.00	80.00	80.00	0.308	0.308	1.00
48	15:34	1.00	0.902	1.00	80.00	80.00	0.061	0.061	1.00
49	15:44	1.00	0.912	1.00	80.00	80.00	0.090	0.090	1.00
50	15:42	1.00	0.910	1.00	80.00	80.00	0.420	0.420	1.00

Quantitation Report File: ST16980831A

Data: ST16980831A.TI  
08/31/98 13:07:00  
Sample: 80UG/ML AP9 062598F  
Conds.: INST. ID: F16

Formula: 1UL INJ. Instrument: F16 Weight: 0.000  
Submitted by: QES Analyst: DAT Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
Resp. fac. from Library Entry

No	Name			
51	3,3'-DIMETHYLBENZIDINE	212	S5# 35	
52	KEPONE	272	S5# 37	
53	2-ACETYLAMINOFLUORENE	181	S5# 45	53-96-3
54	3-METHYLCHOLANTHRENE	268	S5# 85	56-49-5
55	7,12-DIMETHYLBENZANTHRACENE	256	S6# 20	75-97-6
56	HEXACHLOROPHENE	196	S6# 30	70-30-4

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1472	16:22	5	0.949	A BB	56612.	80.000 UG/ML	1.92
52	272	1458	16:13	5	0.940	qedt (h)	12434.	80.000 UG/ML	1.92
53	181	1516	16:52	5	0.977	A BB	61022.	80.000 UG/ML	1.92
54	268	1832	20:23	5	1.181	A BB	76474.	80.000 UG/ML	1.92
55	256	1731	19:15	6	0.971	A BB	66838.	80.000 UG/ML	1.92
56	NOT FOUND								

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:22	1.00	0.949	1.00	80.00	80.00	0.406	0.406	1.00
52	16:13	1.00	0.940	1.00	80.00	80.00	0.089	0.089	1.00
53	16:52	1.00	0.977	1.00	80.00	80.00	0.438	0.438	1.00
54	20:23	1.00	1.181	1.00	80.00	80.00	0.549	0.549	1.00
55	19:15	1.00	0.971	1.00	80.00	80.00	0.610	0.610	1.00
56	21:11		0.992						

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPH.**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND CODED.**

1570

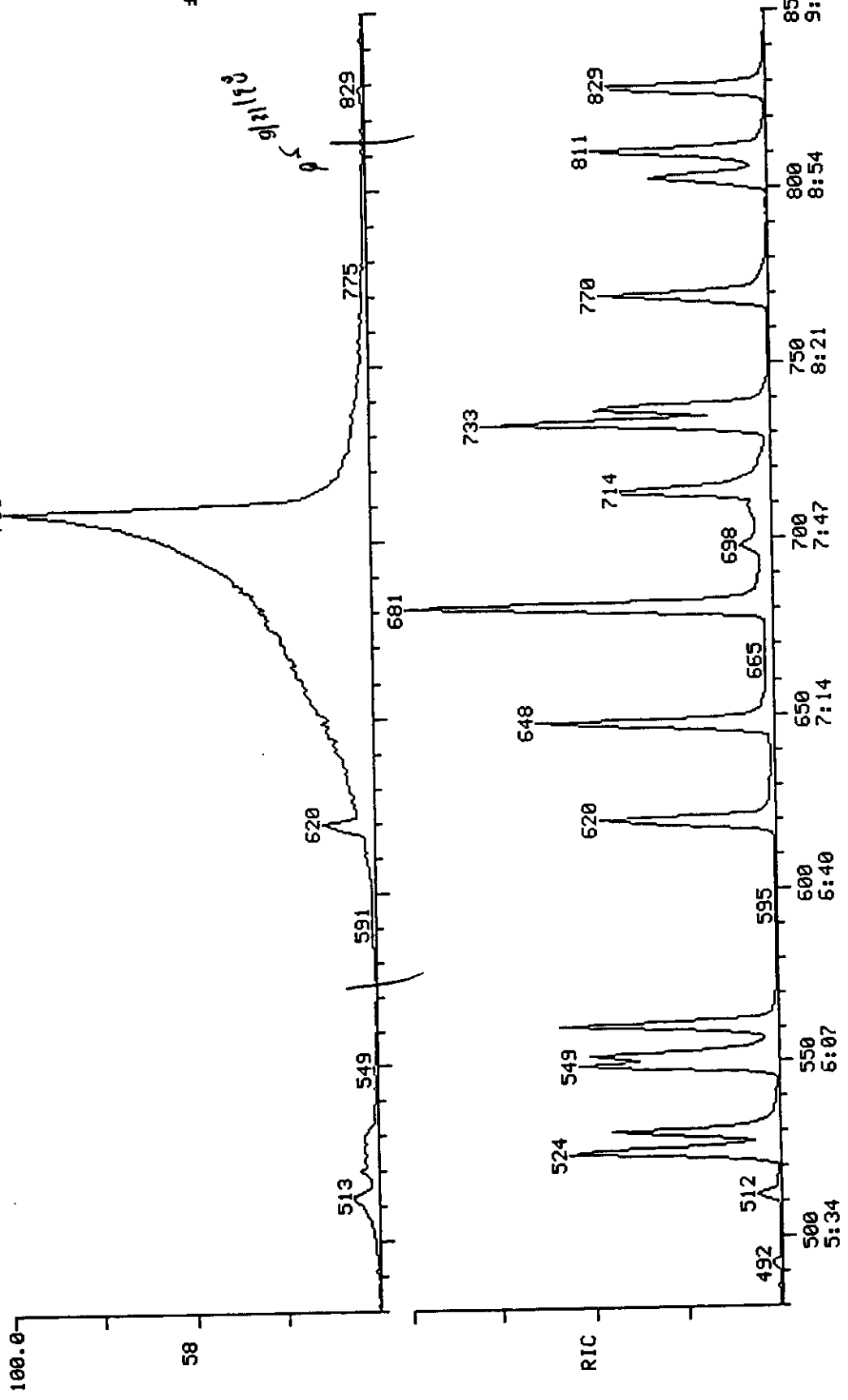
RIC+MASS CHROMATOGRAM  
08/31/98 13:07:00  
SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

DATA: ST16980831A #1  
CALI: ST16980831A #3

7528.

58.017  
± 0.500

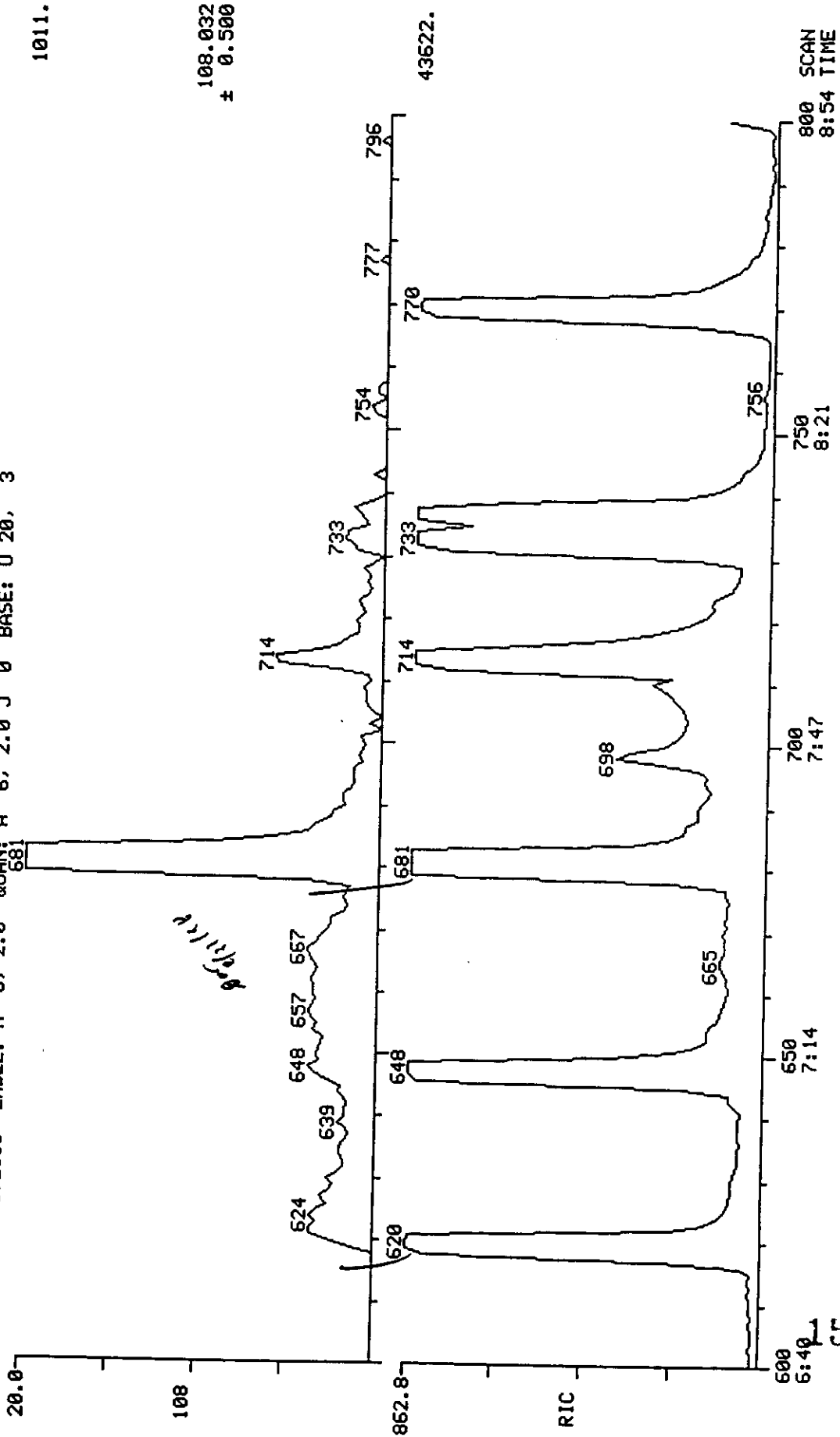
218112.



1571

RIC+MASS CHROMATOGRAM

08/31/98 13:07:00  
SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3  
SCANS 600 TO 800



1011.

108.032  
± 0.500

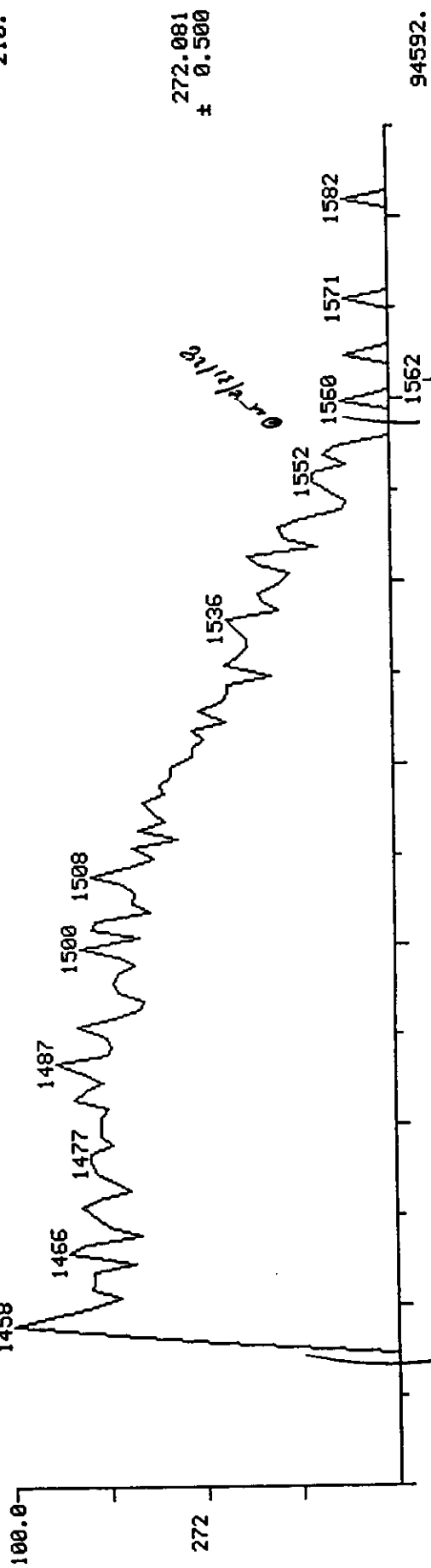
43622.

1572

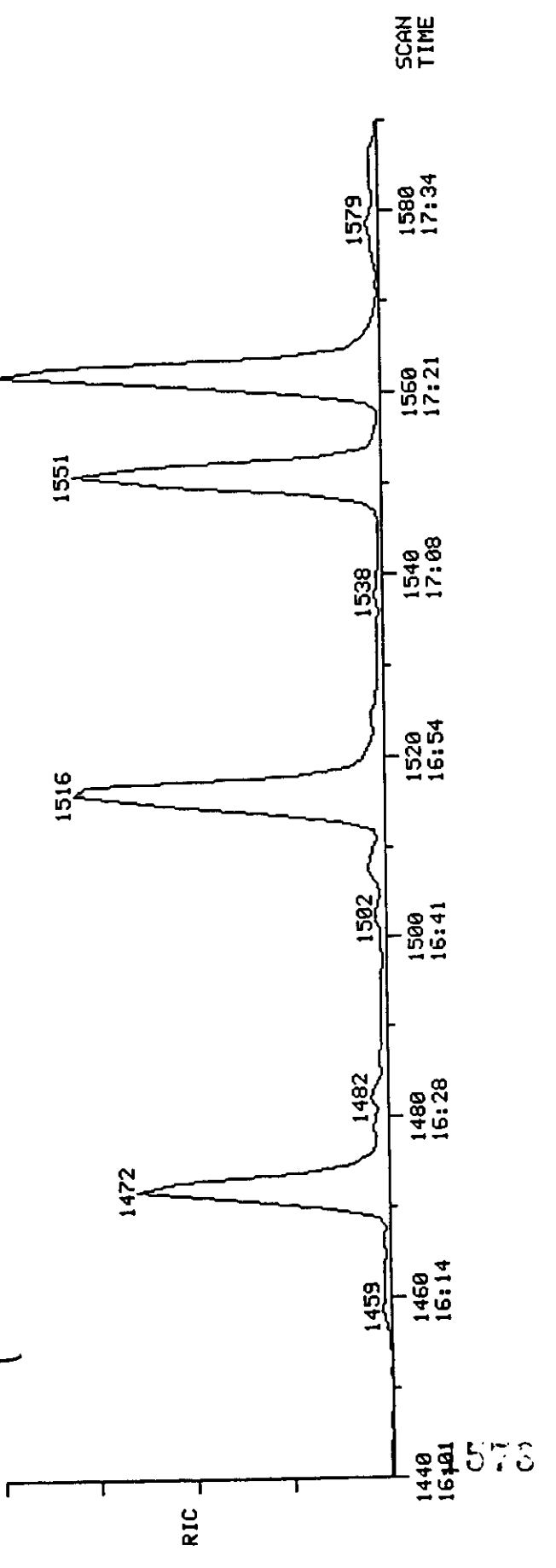
RIC+MASS CHROMATOGRAM  
 DATA: ST16980831A #1  
 CALLI: ST16980831A #3  
 SCANS 1440 TO 1590

08/31/98 13:07:00  
 SAMPLE: 80UG/ML AP9 062598F  
 CONDS.: INST. ID: F16  
 RANGE: G 1458 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

210.



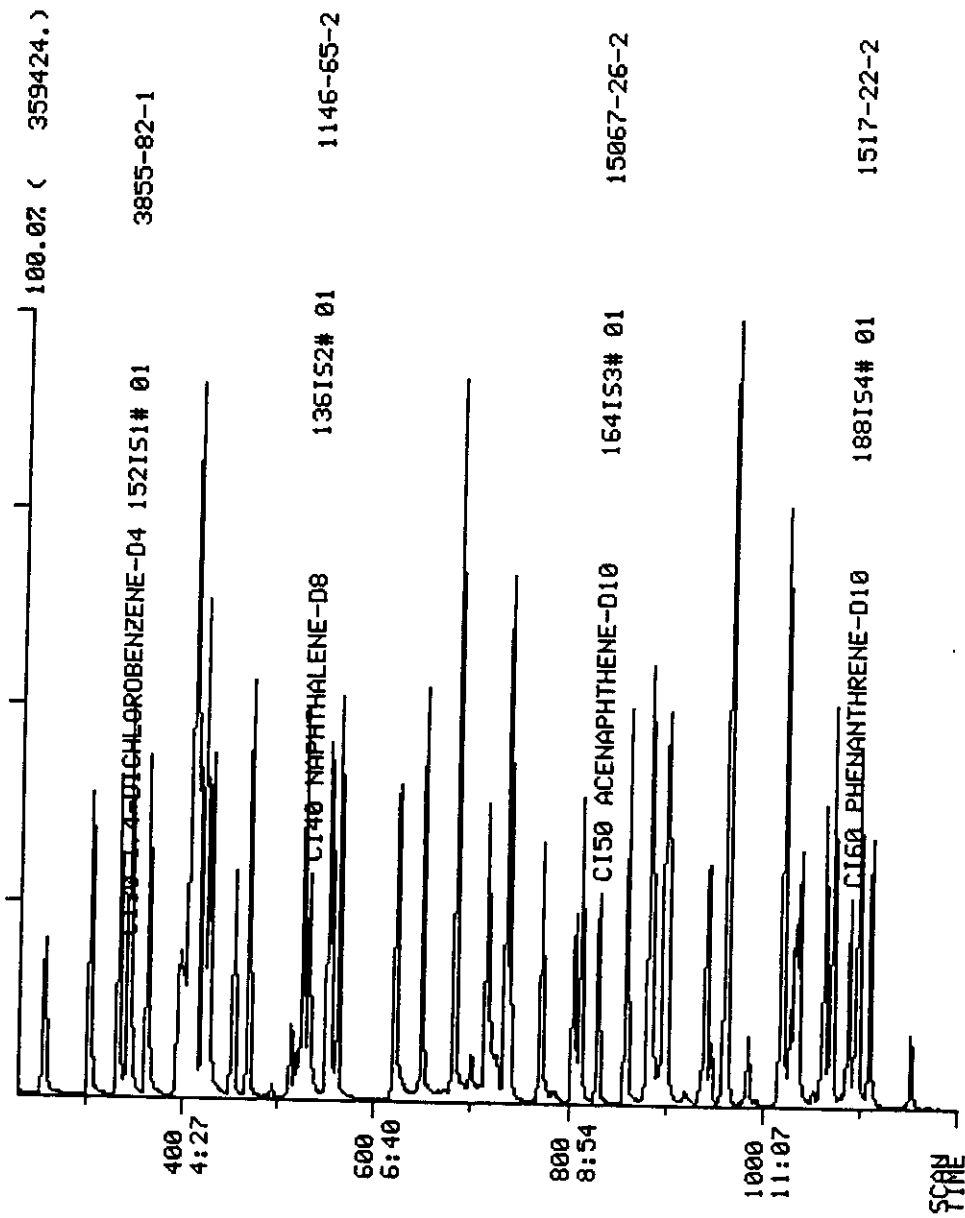
94592.



SCAN TIME

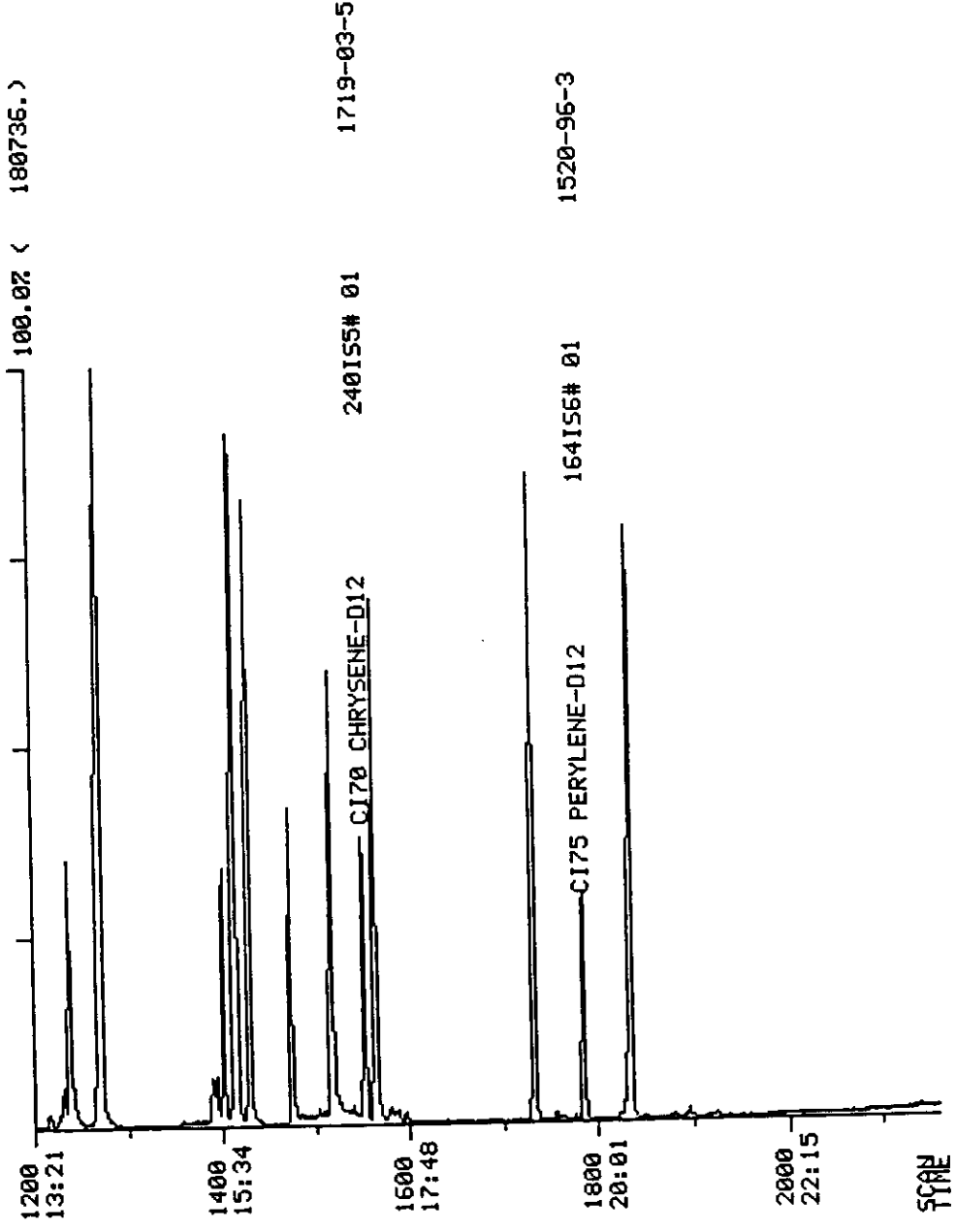
1440	1460	1480	1500	1520	1540	1560	1580
16:14	16:28	16:41	16:54	17:08	17:21	17:34	17:34

DATA FROM FILE: ST16980831E SCANS 230 TO 1200 ACQUIRED: 08/31/98 15:02:00  
CALI: ST16980831E #3  
SAMPLE: 120UG/ML AP9 062598C  
CONDS.: INST. ID: F16



DATA FROM FILE: ST16980831E SCANS 1200 TO 2158 ACQUIRED: 08/31/98 15:02:00  
CALI: ST16980831E #3

SAMPLE: 120UG/ML AP9 062598G  
CONDS.: INST. ID: F16



1575

Data: ST16980831E.T1

08/31/98 15:02:00

Sample: 120UG/ML AP9 062598G

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSOPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROSOPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALLATE	234 S4# 37	
40	4-AMINOBIIPHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	ZSECBUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPYRILENE	58 S4#105	91-80-5
46	ISORIN		
47	CHLOROBENZILATE	139 S4#120	

1578



No	Name			
48	ARAMITE (#1)	185	S5# 20	140-57-8
49	ARAMITE (#2)	185	S5# 25	140-57-8
50	P-DIMETHYLAMINOAZOBENZENE	120	S5# 30	60-11-7

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	28744.	40.000 UG/ML	0.65
2	136	530	5:54	2	1.000	A BB	103999.	40.000 UG/ML	0.65
3	164	829	9:13	3	1.000	A BB	59122.	40.000 UG/ML	0.65
4	188	1087	12:05	4	1.000	A BB	105381.	40.000 UG/ML	0.65
5	240	1551	17:15	5	1.000	A BB	65508.	40.000 UG/ML	0.65
6	264	1782	19:49	6	1.000	A BB	50966.	40.000 UG/ML	0.65
7	79	103	1:09	1	0.303	A BB	124986.	120.000 UG/ML	1.96
8	93	145	1:37	1	0.426	A BB	133361.	120.000 UG/ML	1.96
9	42	157	1:45	1	0.462	A BB	74200.	120.000 UG/ML	1.96
10	80	183	2:02	1	0.538	A BB	53349.	120.000 UG/ML	1.96
11	102	218	2:25	1	0.641	A BB	47209.	120.000 UG/ML	1.96
12	79	255	2:50	1	0.750	A BB	102579.	120.000 UG/ML	1.96
13	117	302	3:22	1	0.888	A BB	51302.	120.000 UG/ML	1.96
14	105	404	4:30	1	1.188	A BB	164114.	120.000 UG/ML	1.96
15	100	408	4:32	1	1.200	A BB	49105.	120.000 UG/ML	1.96
16	56	413	4:36	1	1.215	A BV	70419.	120.000 UG/ML	1.96
17	108	420	4:40	1	1.235	A BB	206207.	120.000 UG/ML	1.96
18	106	409	4:33	1	1.203	A BB	175232.	120.000 UG/ML	1.96
19	42	453	5:02	2	0.855	A BV	105075.	120.000 UG/ML	1.96
20	58	518	5:46	2	0.977	qedt	398258.	120.000 UG/ML	1.96
21	162	553	6:09	2	1.043	A BB	87089.	120.000 UG/ML	1.96
22	213	549	6:06	2	1.036	A BB	61908.	120.000 UG/ML	1.96
23	108	620	6:54	2	1.170	qedt	51610.	120.000 UG/ML	1.96
24	84	621	6:54	2	1.172	A BB	69976.	120.000 UG/ML	1.96
25	162	648	7:12	2	1.223	A BB	76372.	120.000 UG/ML	1.96
26	214	682	7:35	3	0.823	A BB	73776.	120.000 UG/ML	1.96
27	162	698	7:46	3	0.842	A BB	7754.	120.000 UG/ML	1.96
28	104	738	8:13	3	0.890	A BB	33662.	120.000 UG/ML	1.96
29	162	734	8:10	3	0.885	A BV	201116.	120.000 UG/ML	1.96
30	158	770	8:34	3	0.929	A BB	71569.	120.000 UG/ML	1.96
31	168	804	8:57	3	0.970	A BB	37561.	120.000 UG/ML	1.96
32	250	859	9:33	3	1.036	A BB	83210.	120.000 UG/ML	1.96
33	143	884	9:50	3	1.066	A BB	154679.	120.000 UG/ML	1.96
34	143	900	10:01	3	1.086	A BB	173906.	120.000 UG/ML	1.96
35	232	897	9:59	3	1.082	A BB	54245.	120.000 UG/ML	1.96
36	152	941	10:28	3	1.135	A BB	65110.	120.000 UG/ML	1.96
37	75	1026	11:25	4	0.944	A VB	52333.	120.000 UG/ML	1.96
38	108	1037	11:32	4	0.954	A BB	104180.	120.000 UG/ML	1.96
39	234	1018	11:19	4	0.937	A BV	27989.	120.000 UG/ML	1.96
40	169	1070	11:54	4	0.984	A BB	186405.	120.000 UG/ML	1.96
41	173	1097	12:12	4	1.009	A BB	91956.	120.000 UG/ML	1.96
42	237	1062	11:49	4	0.977	A BB	25788.	120.000 UG/ML	1.96
43	211	1109	12:20	4	1.020	A VB	50397.	120.000 UG/ML	1.96
44	190	1236	13:45	4	1.137	A BB	25074.	120.000 UG/ML	1.96
45	58	1267	14:06	4	1.166	A BB	88136.	120.000 UG/ML	1.96
46	193	1270	14:08	4	1.168	A BB	40006.	120.000 UG/ML	1.96
47	139	1428	15:53	4	1.314	A BB	93332.	120.000 UG/ML	1.96
48	185	1400	15:34	5	0.903	A BB	12488.	120.000 UG/ML	1.96
49	185	1414	15:44	5	0.912	A BB	17022.	120.000 UG/ML	1.96
50	120	1412	15:42	5	0.910	A VB	81049.	120.000 UG/ML	1.96

1577

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:15	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:49	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	120.00	120.00	1.449	1.449	1.00
8	1:37	1.00	0.426	1.00	120.00	120.00	1.547	1.547	1.00
9	1:45	1.00	0.462	1.00	120.00	120.00	0.860	0.860	1.00
10	2:01	1.01	0.535	1.01	120.00	120.00	0.619	0.619	1.00
11	2:25	1.00	0.641	1.00	120.00	120.00	0.547	0.547	1.00
12	2:50	1.00	0.750	1.00	120.00	120.00	1.190	1.190	1.00
13	3:22	1.00	0.888	1.00	120.00	120.00	0.595	0.595	1.00
14	4:30	1.00	1.188	1.00	120.00	120.00	1.903	1.903	1.00
15	4:30	1.01	1.191	1.01	120.00	120.00	0.569	0.569	1.00
16	4:34	1.00	1.209	1.00	120.00	120.00	0.817	0.817	1.00
17	4:40	1.00	1.232	1.00	120.00	120.00	2.391	2.391	1.00
18	4:32	1.00	1.200	1.00	120.00	120.00	2.032	2.032	1.00
19	5:01	1.00	0.851	1.00	120.00	120.00	0.337	0.337	1.00
20	7:54	0.73	1.340	0.73	120.00	120.00	1.276	1.276	1.00
21	6:08	1.00	1.042	1.00	120.00	120.00	0.279	0.279	1.00
22	6:06	1.00	1.036	1.00	120.00	120.00	0.198	0.198	1.00
23	7:25	0.93	1.258	0.93	120.00	120.00	0.165	0.165	1.00
24	6:54	1.00	1.170	1.00	120.00	120.00	0.224	0.224	1.00
25	7:12	1.00	1.223	1.00	120.00	120.00	0.245	0.245	1.00
26	7:34	1.00	0.822	1.00	120.00	120.00	0.416	0.416	1.00
27	7:46	1.00	0.843	1.00	120.00	120.00	0.044	0.044	1.00
28	8:12	1.00	0.890	1.00	120.00	120.00	0.190	0.190	1.00
29	8:09	1.00	0.885	1.00	120.00	120.00	1.134	1.134	1.00
30	8:34	1.00	0.930	1.00	120.00	120.00	0.404	0.404	1.00
31	8:56	1.00	0.970	1.00	120.00	120.00	0.212	0.212	1.00
32	9:33	1.00	1.036	1.00	120.00	120.00	0.469	0.469	1.00
33	9:49	1.00	1.066	1.00	120.00	120.00	0.872	0.872	1.00
34	10:00	1.00	1.086	1.00	120.00	120.00	0.980	0.980	1.00
35	9:57	1.00	1.081	1.00	120.00	120.00	0.306	0.306	1.00
36	10:27	1.00	1.135	1.00	120.00	120.00	0.367	0.367	1.00
37	11:24	1.00	0.943	1.00	120.00	120.00	0.166	0.166	1.00
38	11:30	1.00	0.951	1.00	120.00	120.00	0.330	0.330	1.00
39	11:19	1.00	0.937	1.00	120.00	120.00	0.089	0.089	1.00
40	11:53	1.00	0.983	1.00	120.00	120.00	0.590	0.590	1.00
41	12:11	1.00	1.008	1.00	120.00	120.00	0.291	0.291	1.00
42	11:48	1.00	0.976	1.00	120.00	120.00	0.082	0.082	1.00
43	12:19	1.00	1.019	1.00	120.00	120.00	0.159	0.159	1.00
44	13:45	1.00	1.137	1.00	120.00	120.00	0.079	0.079	1.00
45	14:05	1.00	1.165	1.00	120.00	120.00	0.279	0.279	1.00
46	14:08	1.00	1.168	1.00	120.00	120.00	0.127	0.127	1.00
47	15:52	1.00	1.313	1.00	120.00	120.00	0.295	0.295	1.00
48	15:34	1.00	0.902	1.00	120.00	120.00	0.064	0.064	1.00
49	15:44	1.00	0.912	1.00	120.00	120.00	0.087	0.087	1.00
50	15:42	1.00	0.910	1.00	120.00	120.00	0.412	0.412	1.00

Data: ST16980831E.T1  
 08/31/98 15:02:00  
 Sample: 120UG/ML AP9 062598G  
 Conds.: INST. ID: F16  
 Formula: IUL INJ.  
 Submitted by: QES

Instrument: F16 Weight: 0.000  
 Analyst: DAT Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name			
51	3,3'-DIMETHYLBENZIDINE	212 S5# 35		
52	KEPONE	272 S5# 37		
53	2-ACETYLAMINOFLUORENE	181 S5# 45	53-96-3	
54	3-METHYLCHOLANTHRENE	268 S5# 85	56-49-5	
55	7,12-DIMETHYLBENZANTHRACENE	256 S6# 20	75-97-6	
56	HEXACHLOROPHENE	196 S6# 30	70-30-4	

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1473	16:23	5	0.950	A BBQ	71459.	120.000 UG/ML	1.96
52	272	1510	16:48	5	0.974	qedt (1)	11032.	120.000 UG/ML	1.96
53	181	1518	16:53	5	0.979	A BB	84234.	120.000 UG/ML	1.96
54	268	1833	20:23	5	1.182	A BB	101188.	120.000 UG/ML	1.96
55	256	1731	19:15	6	0.971	A BV	92986.	120.000 UG/ML	1.96
56	NOT FOUND								

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:22	1.00	0.949	1.00	120.00	120.00	0.364	0.364	1.00
52	16:13	1.04	0.940	1.04	120.00	120.00	0.056	0.056	1.00
53	16:52	1.00	0.977	1.00	120.00	120.00	0.429	0.429	1.00
54	20:23	1.00	1.181	1.00	120.00	120.00	0.515	0.515	1.00
55	19:15	1.00	0.971	1.00	120.00	120.00	0.608	0.608	1.00
56	21:11		0.992						

MANUAL EDIT CODES  
 1. PEAK NOT FOUND  
 2. POOR CHROMATOGRAPHY  
 3. WRONG ISOMER  
 ALL MANUAL EDITS MUST BE  
 INITIALED, DATED, AND CORRECTED

SCANS 480 TO 800

DATA: ST16980831E #1  
CALI: ST16980831E #3

RIC+MASS CHROMATOGRAM

08/31/98 15:02:00

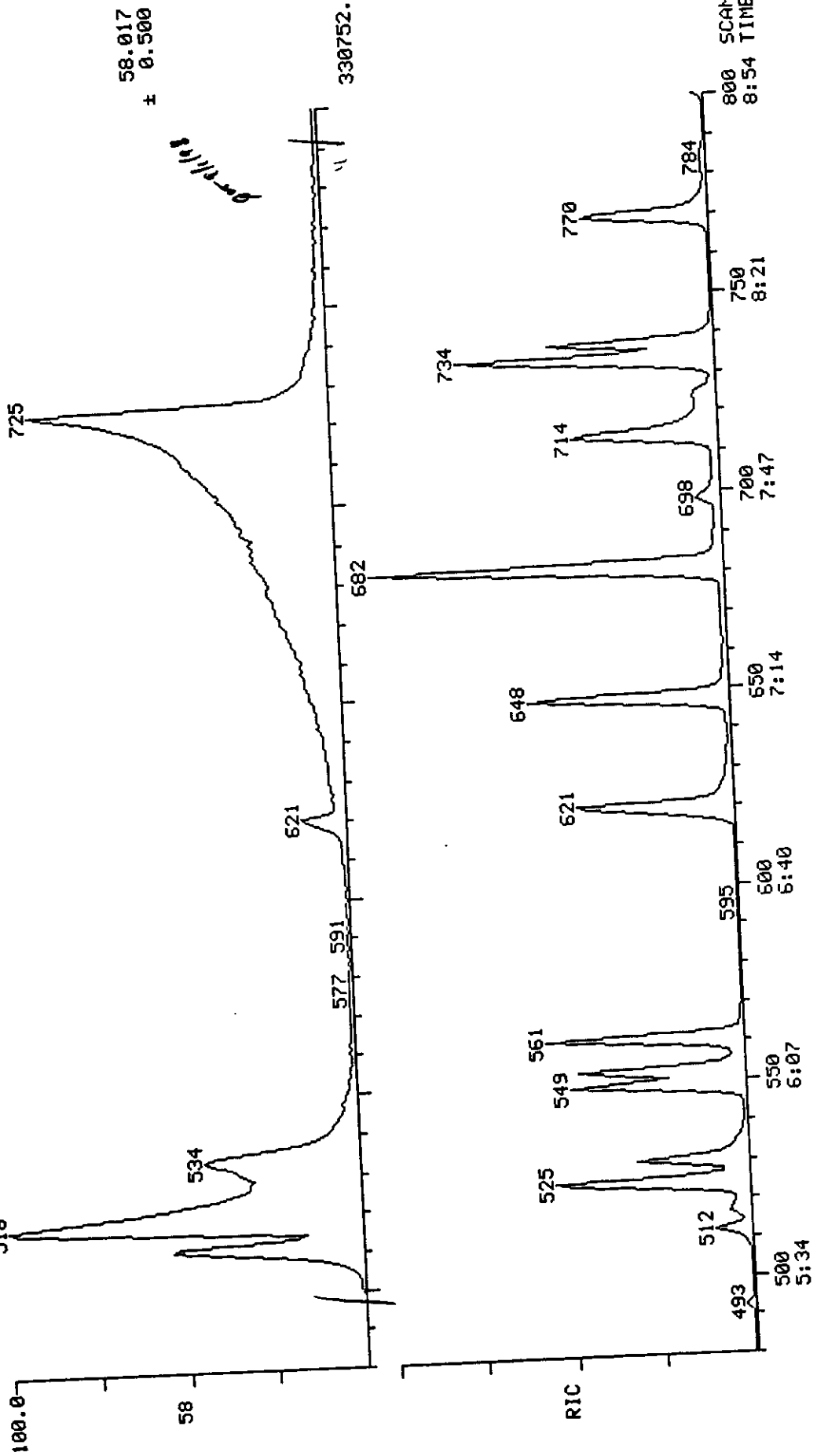
SAMPLE: 120UG/ML AP9 062598G

CONDS.: INST. ID: F16

RANGE: G 518

LABEL: N 0, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3

9664.



1580

SCANS 600 TO 700

DATA: ST16980831E #1  
CALI: ST16980831E #3

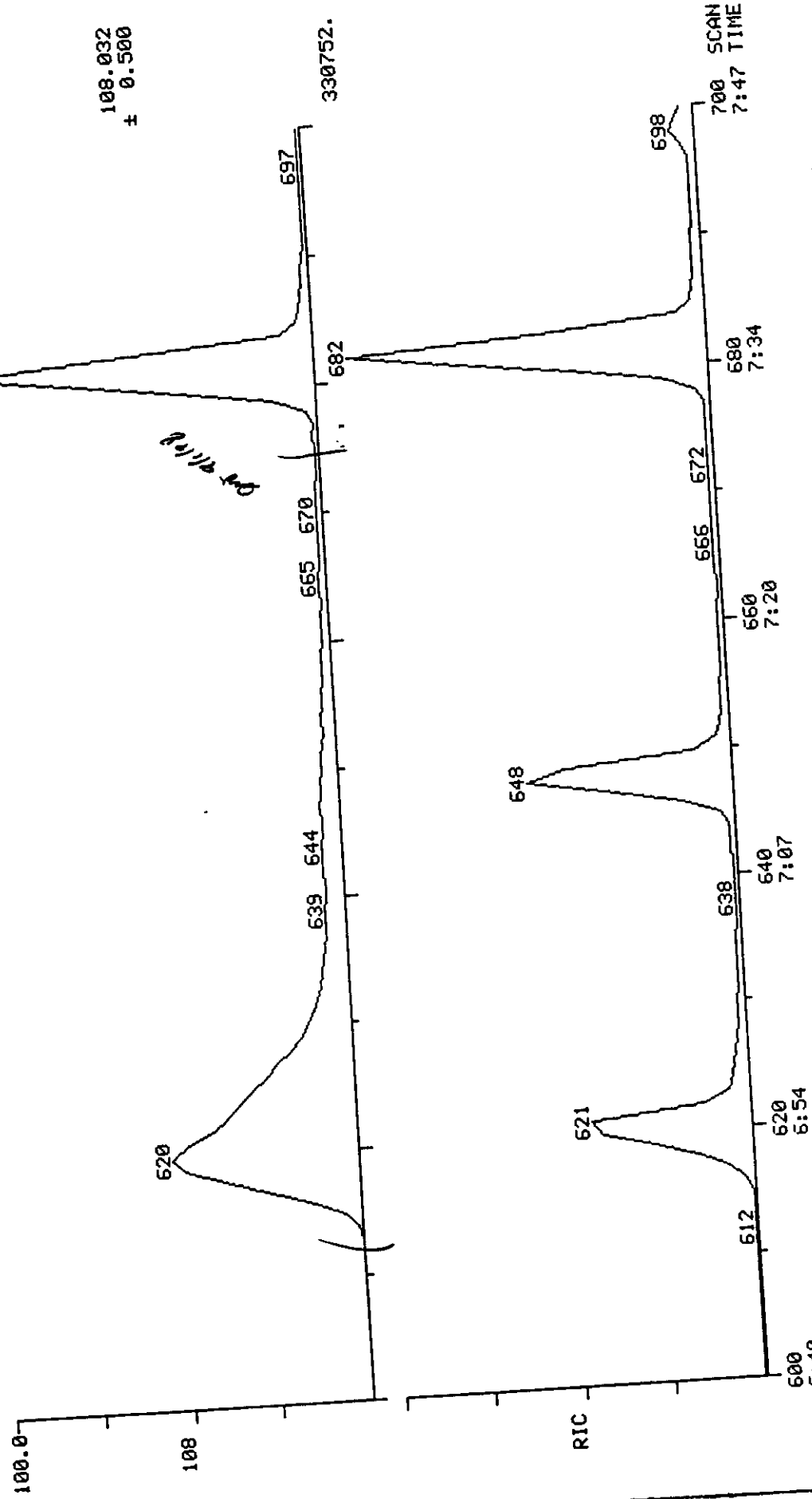
RIC+MASS CHROMATOGRAM

08/31/98 15:02:00  
SAMPLE: 120UG/ML AP9 062598G  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

8024.

108.032  
± 0.500

330752.

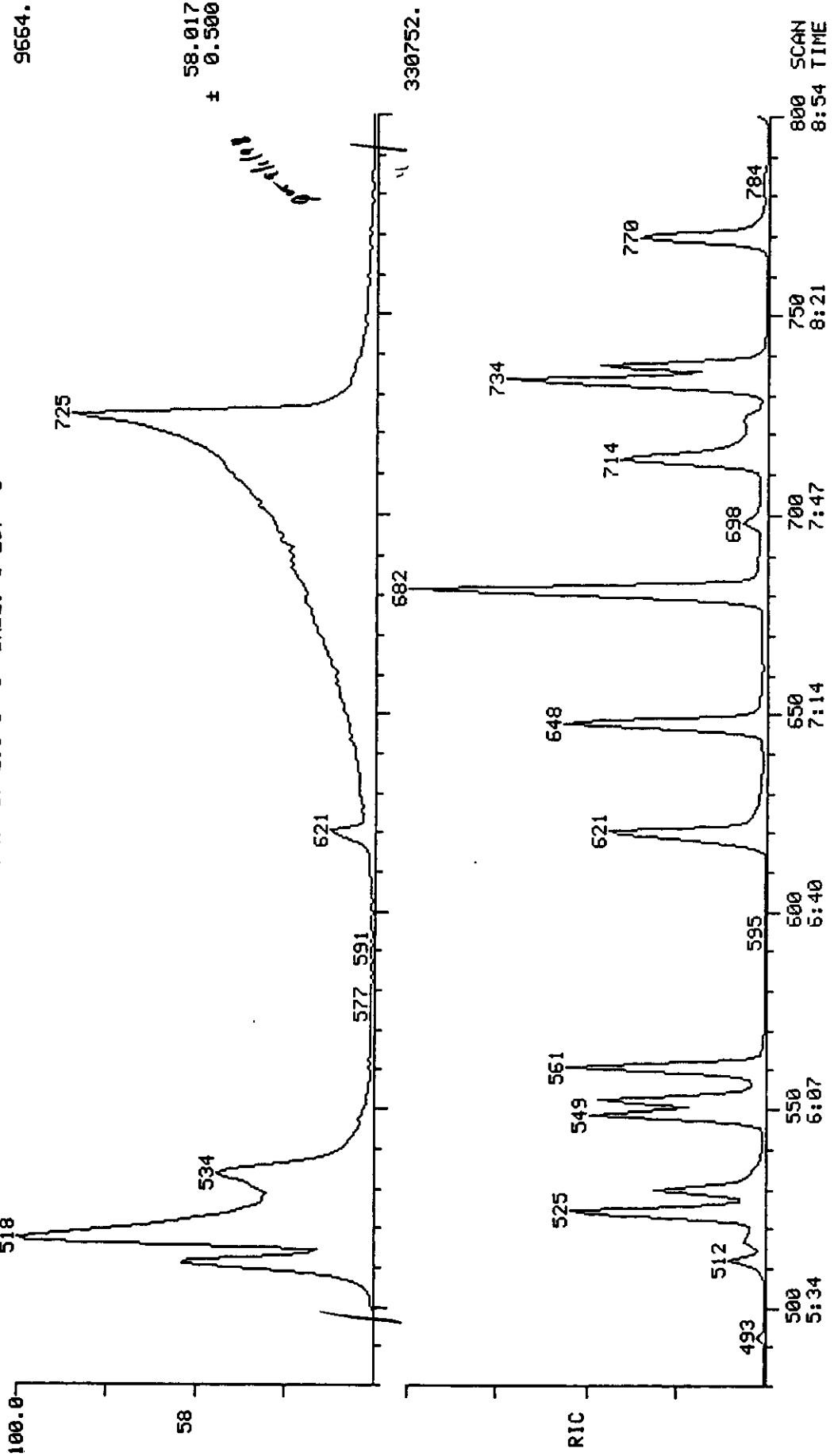


600  
6:40  
1581

RIC+MASS CHROMATOGRAM  
08/31/98 15:02:00  
SAMPLE: 120UG/ML AP9 062598G  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0. 2.0 QUAN: A 6. 2.0 J 0 BASE: U 20. 3

DATA: ST16980831E #1  
CALI: ST16980831E #3  
SCANS 480 TO 800

9664.



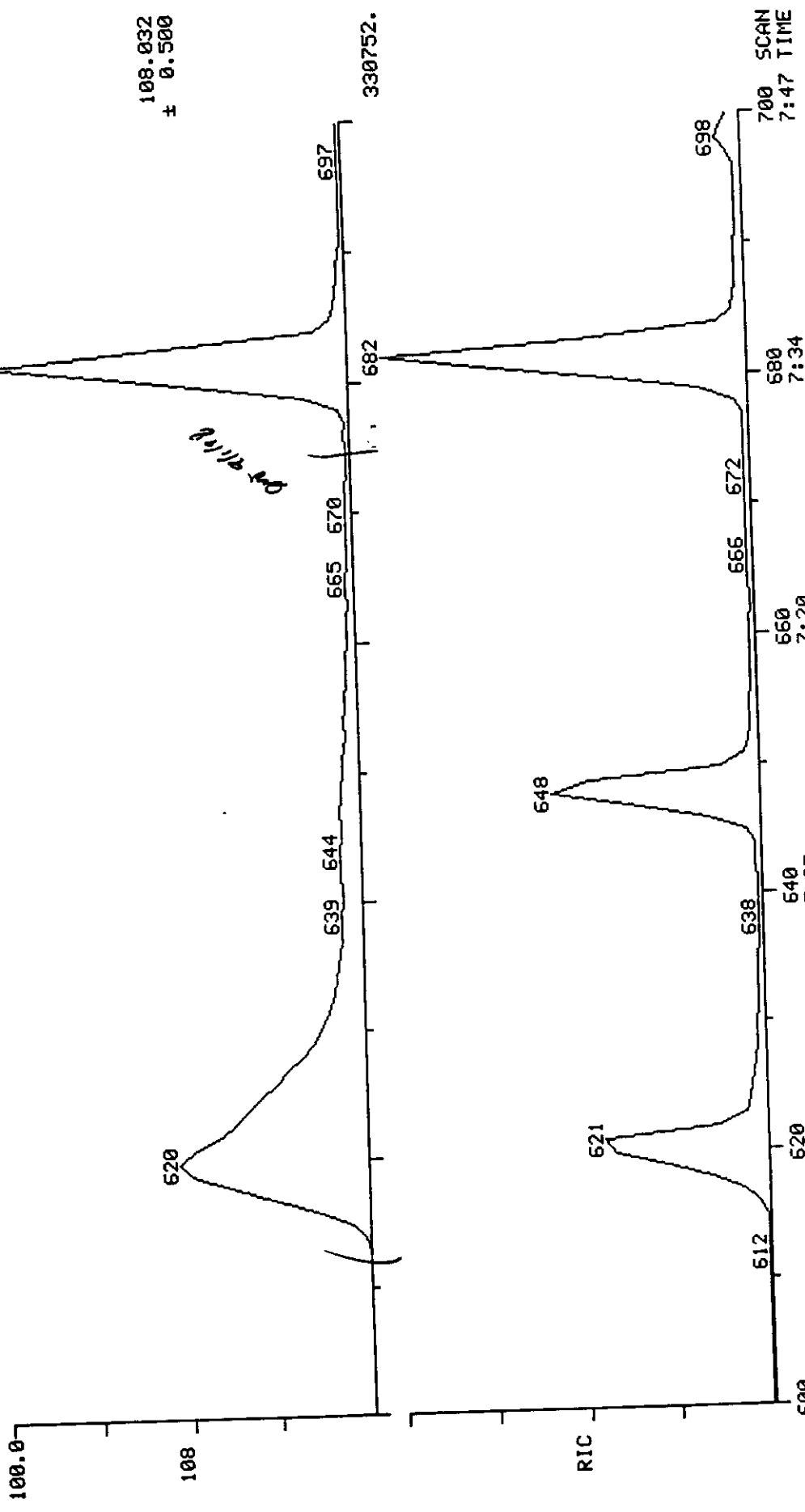
0221

SCANS 600 TO 700

DATA: ST16980831E #1  
CALI: ST16980831E #3

RIC+MASS CHROMATOGRAM  
08/31/98 15:02:00  
SAMPLE: 120UG/ML AP9 062598G  
CONDS.: INST. ID: F16  
RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3

8024.

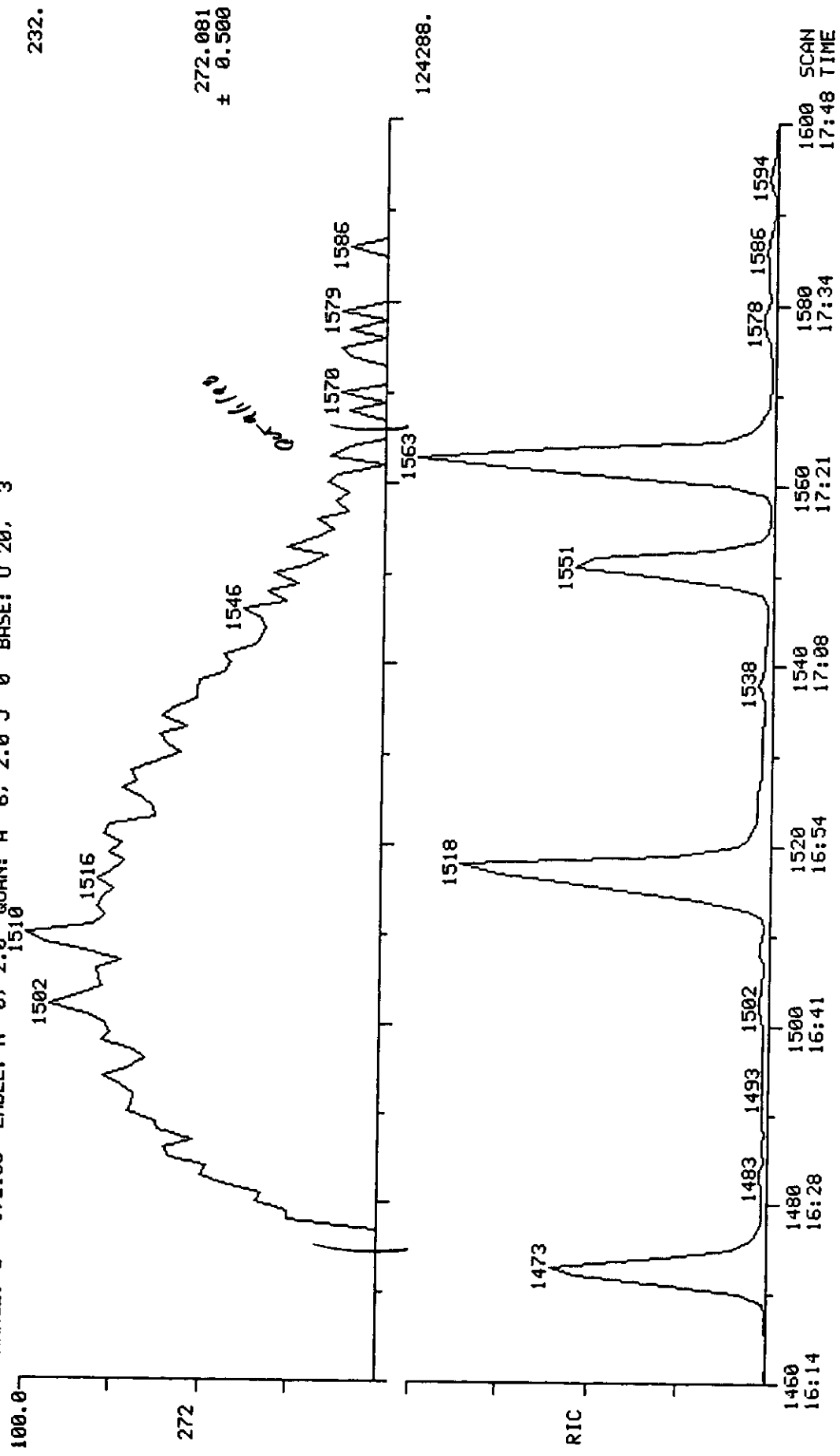


600 6:40  
620 6:54  
640 7:07  
660 7:20  
680 7:34  
700 7:47  
SCAN TIME

1281

RIC+MASS CHROMATOGRAM  
08/31/98 15:02:00  
SAMPLE: 120UG/ML AP9 062598G  
CONDS.: INST. ID: F16  
RANGE: G 1.2158 LABEL: N 0, 2.0, QUAN: A 6, 2.0 J 0 BASE: U 20, 3

DATA: ST16980831E #1  
CALI: ST16980831E #3  
SCANS 1460 TO 1600



232.

272.081  
± 0.500

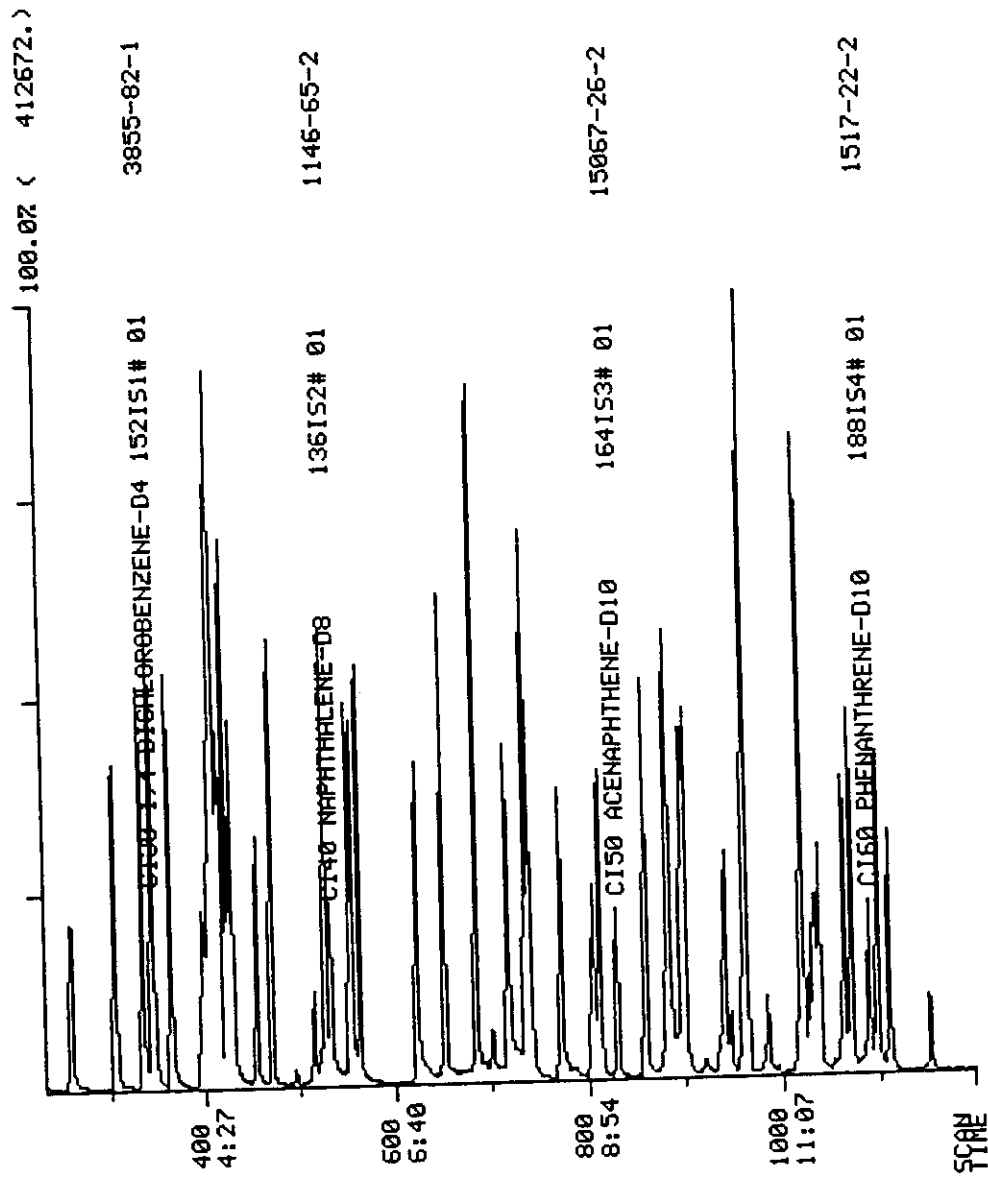
124288.

1582

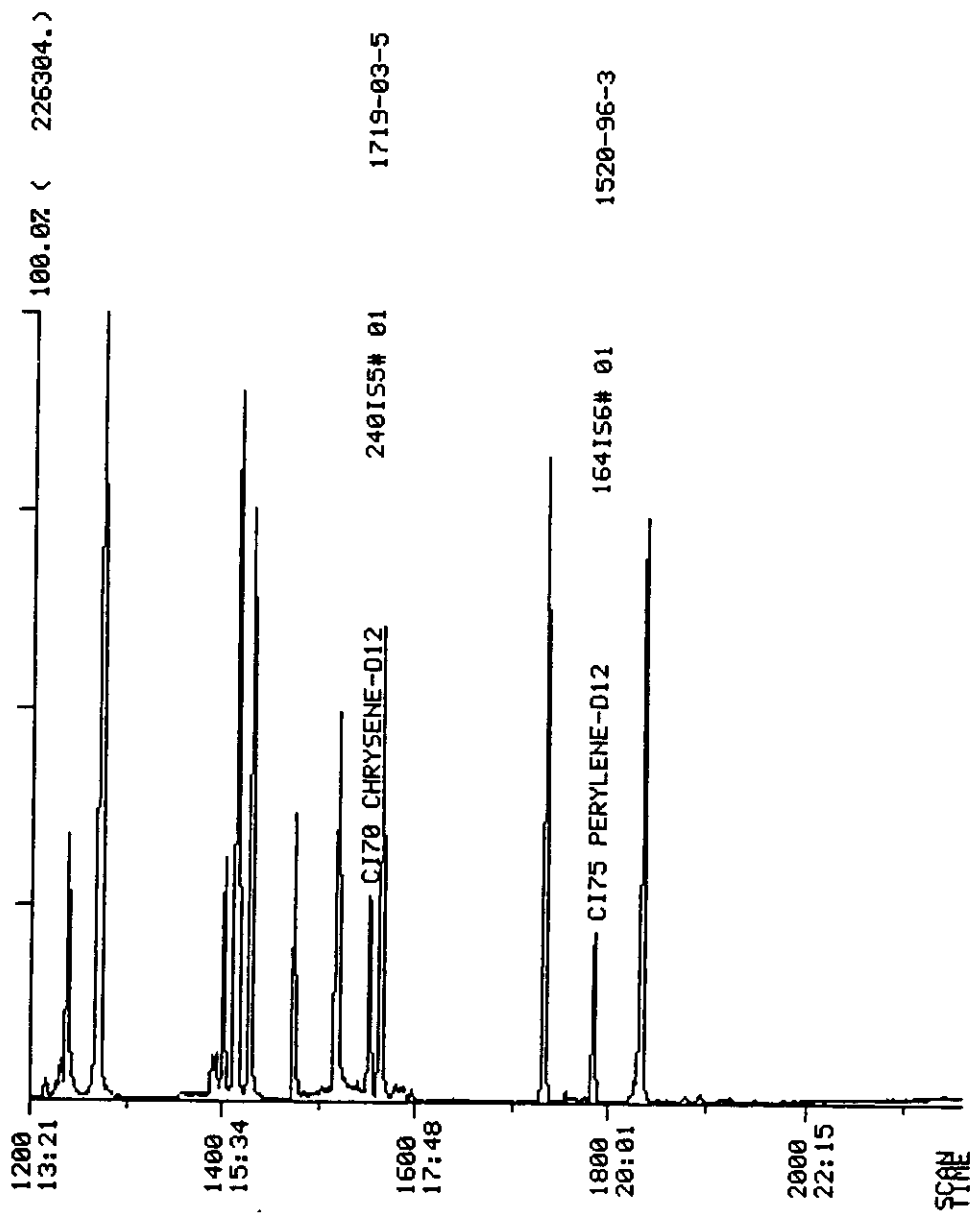


DATA FROM FILE: ST16980831F SCANS 230 TO 1200 ACQUIRED: 08/31/98 15:31:00  
CALI: ST16980831F #3

SAMPLE: 160UG/ML AP9 062598H  
CONDS.: INST. ID: F16



DATA FROM FILE: ST16980831F SCANS 1200 TO 2158 ACQUIRED: 08/31/98 15:31:00  
SAMPLE: 160UG/ML AP9 062598H CALI: ST16980831F #3  
CONDS.: INST. ID: F16



1584

Data: ST16980831F.TI  
 08/31/98 15:31:00  
 Sample: 160UG/ML AP9 062598H  
 Conds.: INST. ID: F16  
 Formula: 1UL INJ.  
 Submitted by: QES

Instrument: F16  
 Analyst: DAT

Weight: 0.000  
 Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from Library Entry

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROSOPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALATE	234 S4# 37	
40	4-AMINOBIIPHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	2SEC BUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPYRILENE	58 S4#105	91-80-5
46	ISODRIN		
47	CHLOROBENZILATE	139 S4#120	

No	Name		
48	ARAMITE (#1)	185 S5# 20	140-57-8
49	ARAMITE (#2)	185 S5# 25	140-57-8
50	P-DIMETHYLAMINOAZOBENZENE	120 S5# 30	60-11-7

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	341	3:48	1	1.000	A BB	28546.	40.000 UG/ML	0.50
2	136	530	5:54	2	1.000	A BB	103260.	40.000 UG/ML	0.50
3	164	829	9:13	3	1.000	A BB	58634.	40.000 UG/ML	0.50
4	188	1088	12:06	4	1.000	A BV	95521.	40.000 UG/ML	0.50
5	240	1551	17:15	5	1.000	A BB	56500.	40.000 UG/ML	0.50
6	264	1783	19:50	6	1.000	A BV	44220.	40.000 UG/ML	0.50
7	79	103	1:09	1	0.302	A BB	179951.	160.000 UG/ML	1.98
8	93	145	1:37	1	0.425	A BB	176388.	160.000 UG/ML	1.98
9	42	157	1:45	1	0.460	A BB	99310.	160.000 UG/ML	1.98
10	80	183	2:02	1	0.537	A BB	68989.	160.000 UG/ML	1.98
11	102	219	2:26	1	0.642	A BB	63322.	160.000 UG/ML	1.98
12	79	256	2:51	1	0.751	A BB	142082.	160.000 UG/ML	1.98
13	117	303	3:22	1	0.889	A BB	67568.	160.000 UG/ML	1.98
14	105	405	4:30	1	1.188	A BB	217271.	160.000 UG/ML	1.98
15	100	410	4:34	1	1.202	A BB	65650.	160.000 UG/ML	1.98
16	56	414	4:36	1	1.214	A BV	93134.	160.000 UG/ML	1.98
17	108	422	4:42	1	1.238	A BB	273406.	160.000 UG/ML	1.98
18	106	410	4:34	1	1.202	A BB	222560.	160.000 UG/ML	1.98
19	42	454	5:03	2	0.857	A BB	135876.	160.000 UG/ML	1.98
20	58	742	8:15	2	1.400	qedt	489048.	160.000 UG/ML	1.98
21	162	553	6:09	2	1.043	A BB	117693.	160.000 UG/ML	1.98
22	213	549	6:06	2	1.036	A BB	81868.	160.000 UG/ML	1.98
23	108	621	6:54	2	1.172	qedt	49954.	160.000 UG/ML	1.98
24	84	621	6:54	2	1.172	A BB	92687.	160.000 UG/ML	1.98
25	162	649	7:13	2	1.225	A BB	103119.	160.000 UG/ML	1.98
26	214	683	7:36	3	0.824	A BB	98348.	160.000 UG/ML	1.98
27	162	698	7:46	3	0.842	A BB	10361.	160.000 UG/ML	1.98
28	104	738	8:13	3	0.890	A BB	44958.	160.000 UG/ML	1.98
29	162	734	8:10	3	0.885	A BB	348940.	160.000 UG/ML	1.98
30	158	771	8:35	3	0.930	A BB	91288.	160.000 UG/ML	1.98
31	168	805	8:57	3	0.971	A BB	47740.	160.000 UG/ML	1.98
32	250	859	9:33	3	1.036	A BB	110577.	160.000 UG/ML	1.98
33	143	884	9:50	3	1.066	A BB	180448.	160.000 UG/ML	1.98
34	143	901	10:01	3	1.087	A BB	206685.	160.000 UG/ML	1.98
35	232	897	9:59	3	1.082	A BB	68020.	160.000 UG/ML	1.98
36	152	942	10:29	3	1.136	A BB	74228.	160.000 UG/ML	1.98
37	75	1026	11:25	4	0.943	A VB	60745.	160.000 UG/ML	1.98
38	108	1038	11:33	4	0.954	A BB	118791.	160.000 UG/ML	1.98
39	234	1019	11:20	4	0.937	A BV	36960.	160.000 UG/ML	1.98
40	169	1070	11:54	4	0.983	A BB	208505.	160.000 UG/ML	1.98
41	173	1097	12:12	4	1.008	A BB	105932.	160.000 UG/ML	1.98
42	237	1062	11:49	4	0.976	A BB	30912.	160.000 UG/ML	1.98
43	211	1109	12:20	4	1.019	A VB	57279.	160.000 UG/ML	1.98
44	190	1237	13:46	4	1.137	A BB	28601.	160.000 UG/ML	1.98
45	58	1267	14:06	4	1.165	A BB	106019.	160.000 UG/ML	1.98
46	193	1271	14:08	4	1.168	A BV	47357.	160.000 UG/ML	1.98
47	139	1428	15:53	4	1.312	A BB	107482.	160.000 UG/ML	1.98
48	185	1400	15:34	5	0.903	A BB	13837.	160.000 UG/ML	1.98
49	185	1414	15:44	5	0.912	A BV	19178.	160.000 UG/ML	1.98
50	120	1413	15:43	5	0.911	A BB	94845.	160.000 UG/ML	1.98

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:15	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:49	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	160.00	160.00	1.576	1.576	1.00
8	1:37	1.00	0.426	1.00	160.00	160.00	1.545	1.545	1.00
9	1:45	1.00	0.462	1.00	160.00	160.00	0.870	0.870	1.00
10	2:01	1.01	0.535	1.00	160.00	160.00	0.604	0.604	1.00
11	2:25	1.00	0.641	1.00	160.00	160.00	0.555	0.555	1.00
12	2:50	1.00	0.750	1.00	160.00	160.00	1.244	1.244	1.00
13	3:22	1.00	0.888	1.00	160.00	160.00	0.592	0.592	1.00
14	4:30	1.00	1.188	1.00	160.00	160.00	1.903	1.903	1.00
15	4:30	1.01	1.191	1.01	160.00	160.00	0.575	0.575	1.00
16	4:34	1.01	1.209	1.00	160.00	160.00	0.816	0.816	1.00
17	4:40	1.01	1.232	1.00	160.00	160.00	2.394	2.394	1.00
18	4:32	1.00	1.200	1.00	160.00	160.00	1.949	1.949	1.00
19	5:01	1.01	0.851	1.01	160.00	160.00	0.329	0.329	1.00
20	7:54	1.05	1.340	1.05	160.00	160.00	1.184	1.184	1.00
21	6:08	1.00	1.042	1.00	160.00	160.00	0.285	0.285	1.00
22	6:06	1.00	1.036	1.00	160.00	160.00	0.198	0.198	1.00
23	7:25	0.93	1.258	0.93	160.00	160.00	0.121	0.121	1.00
24	6:54	1.00	1.170	1.00	160.00	160.00	0.224	0.224	1.00
25	7:12	1.00	1.223	1.00	160.00	160.00	0.250	0.250	1.00
26	7:34	1.00	0.822	1.00	160.00	160.00	0.419	0.419	1.00
27	7:46	1.00	0.843	1.00	160.00	160.00	0.044	0.044	1.00
28	8:12	1.00	0.890	1.00	160.00	160.00	0.192	0.192	1.00
29	8:09	1.00	0.885	1.00	160.00	160.00	1.488	1.488	1.00
30	8:34	1.00	0.930	1.00	160.00	160.00	0.389	0.389	1.00
31	8:56	1.00	0.970	1.00	160.00	160.00	0.204	0.204	1.00
32	9:33	1.00	1.036	1.00	160.00	160.00	0.471	0.471	1.00
33	9:49	1.00	1.066	1.00	160.00	160.00	0.769	0.769	1.00
34	10:00	1.00	1.086	1.00	160.00	160.00	0.881	0.881	1.00
35	9:57	1.00	1.081	1.00	160.00	160.00	0.290	0.290	1.00
36	10:27	1.00	1.135	1.00	160.00	160.00	0.316	0.316	1.00
37	11:24	1.00	0.943	1.00	160.00	160.00	0.159	0.159	1.00
38	11:30	1.00	0.951	1.00	160.00	160.00	0.311	0.311	1.00
39	11:19	1.00	0.937	1.00	160.00	160.00	0.097	0.097	1.00
40	11:53	1.00	0.983	1.00	160.00	160.00	0.546	0.546	1.00
41	12:11	1.00	1.008	1.00	160.00	160.00	0.277	0.277	1.00
42	11:48	1.00	0.976	1.00	160.00	160.00	0.081	0.081	1.00
43	12:19	1.00	1.019	1.00	160.00	160.00	0.150	0.150	1.00
44	13:45	1.00	1.137	1.00	160.00	160.00	0.075	0.075	1.00
45	14:05	1.00	1.165	1.00	160.00	160.00	0.277	0.277	1.00
46	14:08	1.00	1.168	1.00	160.00	160.00	0.124	0.124	1.00
47	15:52	1.00	1.313	1.00	160.00	160.00	0.281	0.281	1.00
48	15:34	1.00	0.902	1.00	160.00	160.00	0.061	0.061	1.00
49	15:44	1.00	0.912	1.00	160.00	160.00	0.085	0.085	1.00
50	15:42	1.00	0.910	1.00	160.00	160.00	0.420	0.420	1.00

Data: ST16980831F.TI

08/31/98 15:31:00

Sample: 160UG/ML AP9 062598H

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from Library Entry

No	Name			
51	3,3'-DIMETHYLBENZIDINE	212 S5# 35		
52	KEPONE	272 S5# 37		
53	2-ACETYLAMINOFLUORENE	181 S5# 45	53-96-3	
54	3-METHYLCHOLANTHRENE	268 S5# 85	56-49-5	
55	7,12-DIMETHYLBENZANTHRACENE	256 S6# 20	75-97-6	
56	HEXACHLOROPHENE	196 S6# 30	70-30-4	

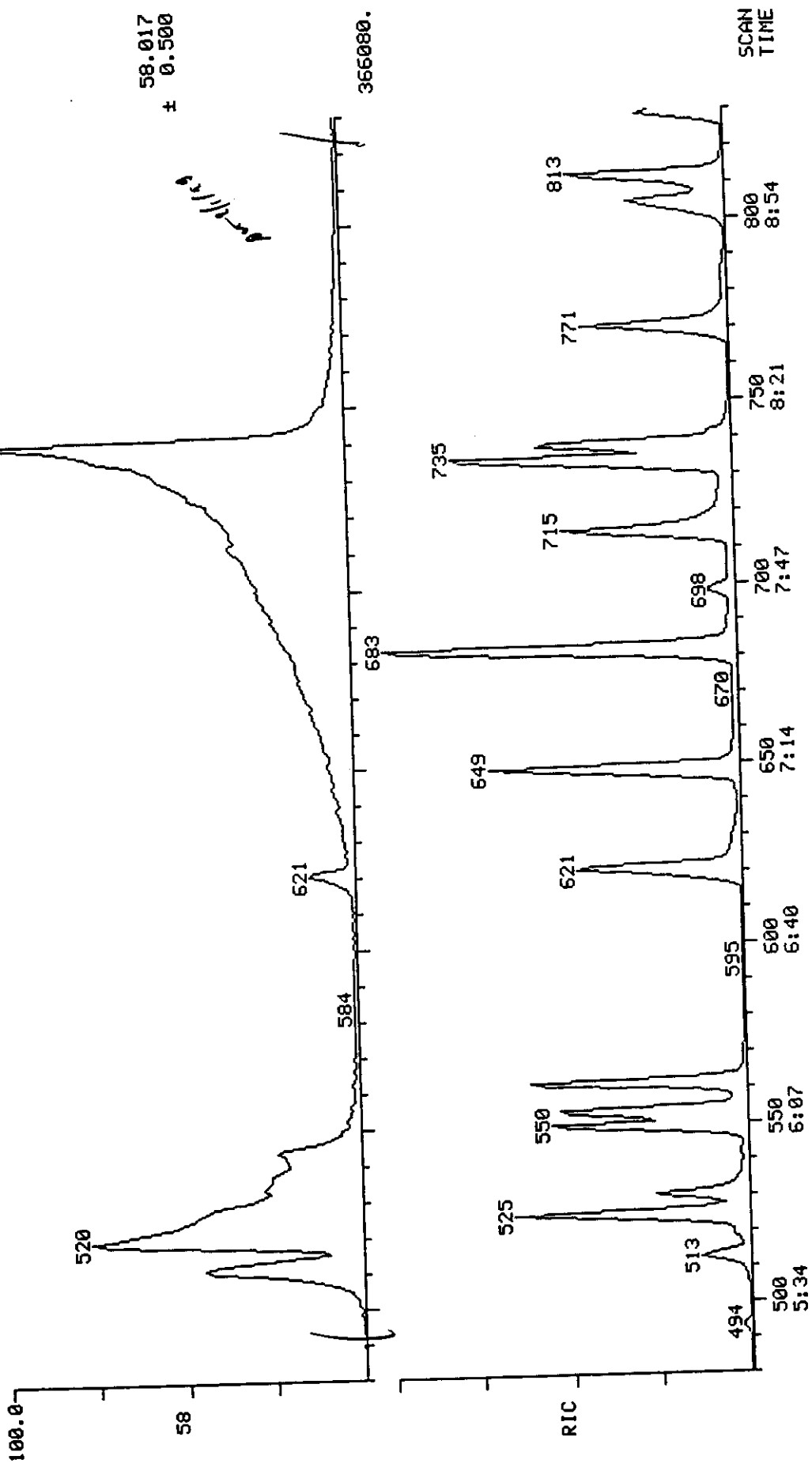
No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1473	16:23	5	0.950	A BV	78087.	160.000 UG/ML	1.98
52	272	1508	16:46	5	0.972	qedt	12248.	160.000 UG/ML	1.98
53	181	1518	16:53	5	0.979	A BB	99486.	160.000 UG/ML	1.98
54	268	1834	20:24	5	1.182	A BB	116191.	160.000 UG/ML	1.98
55	256	1732	19:16	6	0.971	A BB	107710.	160.000 UG/ML	1.98
56	NOT FOUND								

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:22	1.00	0.949	1.00	160.00	160.00	0.346	0.346	1.00
52	16:13	1.03	0.940	1.03	160.00	160.00	0.054	0.054	1.00
53	16:52	1.00	0.977	1.00	160.00	160.00	0.440	0.440	1.00
54	20:23	1.00	1.181	1.00	160.00	160.00	0.514	0.514	1.00
55	19:15	1.00	0.971	1.00	160.00	160.00	0.609	0.609	1.00
56	21:11		0.992						

MANUAL EDIT CODES  
 1. PEAK NOT FOUND  
 2. POOR CHROMATOGRAPHY  
 3. WRONG ISOMER  
 ALL MANUAL EDITS MUST BE  
 INITIALED, DATED, AND CODED

RIC+MASS CHROMATOGRAM  
08/31/98 15:31:00  
SAMPLE: 160UG/ML AP9 062598H  
CONDS.: INST. ID: F16  
RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3 742

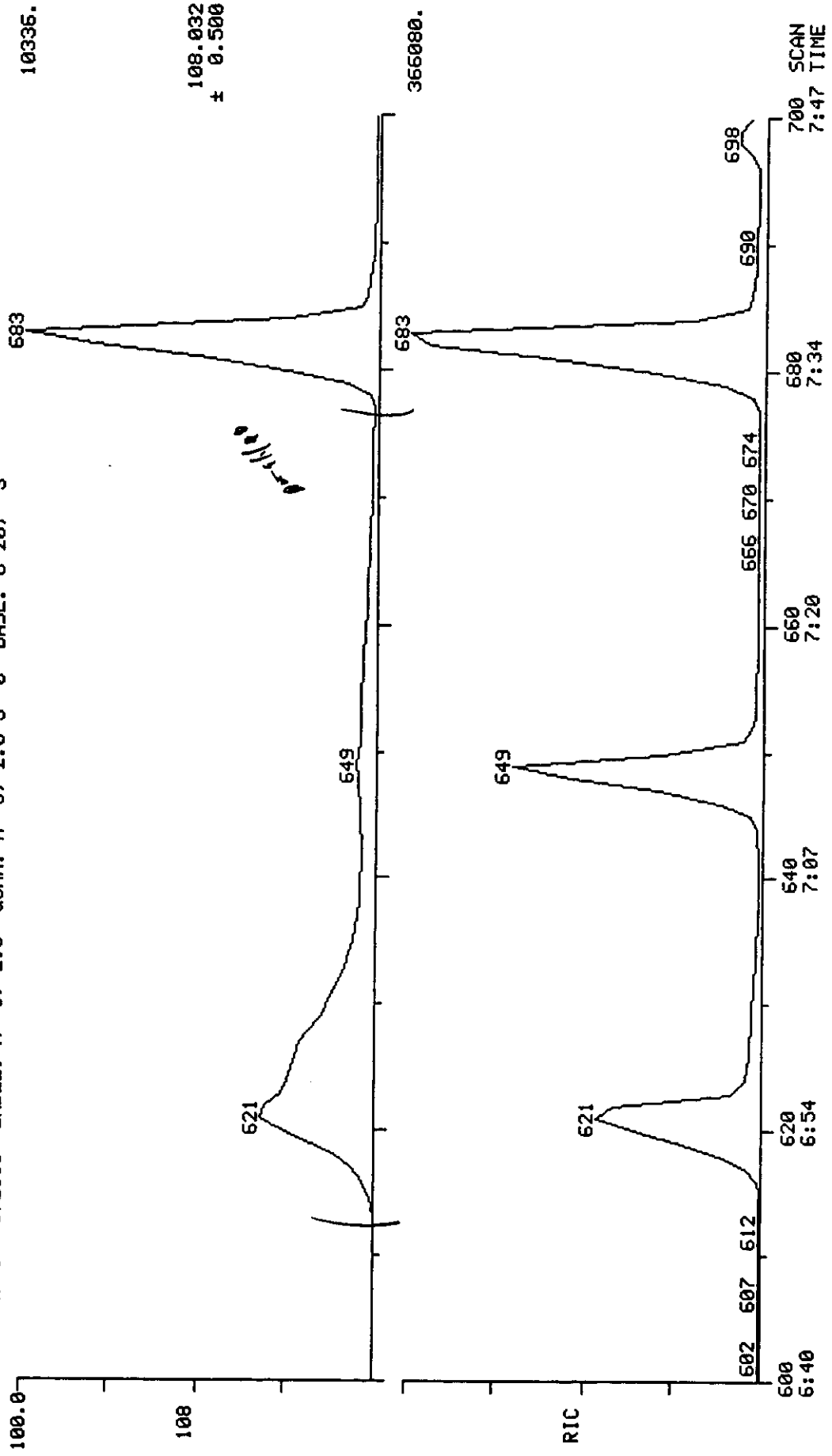
10544.



6301

RIC+MASS CHROMATOGRAM  
08/31/98 15:31:00  
SAMPLE: 160UG/ML AP9 062598H  
CONDS.: INST. ID: F16  
RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3

DATA: ST16980831F #1  
CALI: ST16980831F #3  
SCANS 500 TO 700



1590



SCANS 1450 TO 1600

DATA: ST16980831F #1  
CALI: ST16980831F #3

RIC+MASS CHROMATOGRAM

08/31/98 15:31:00

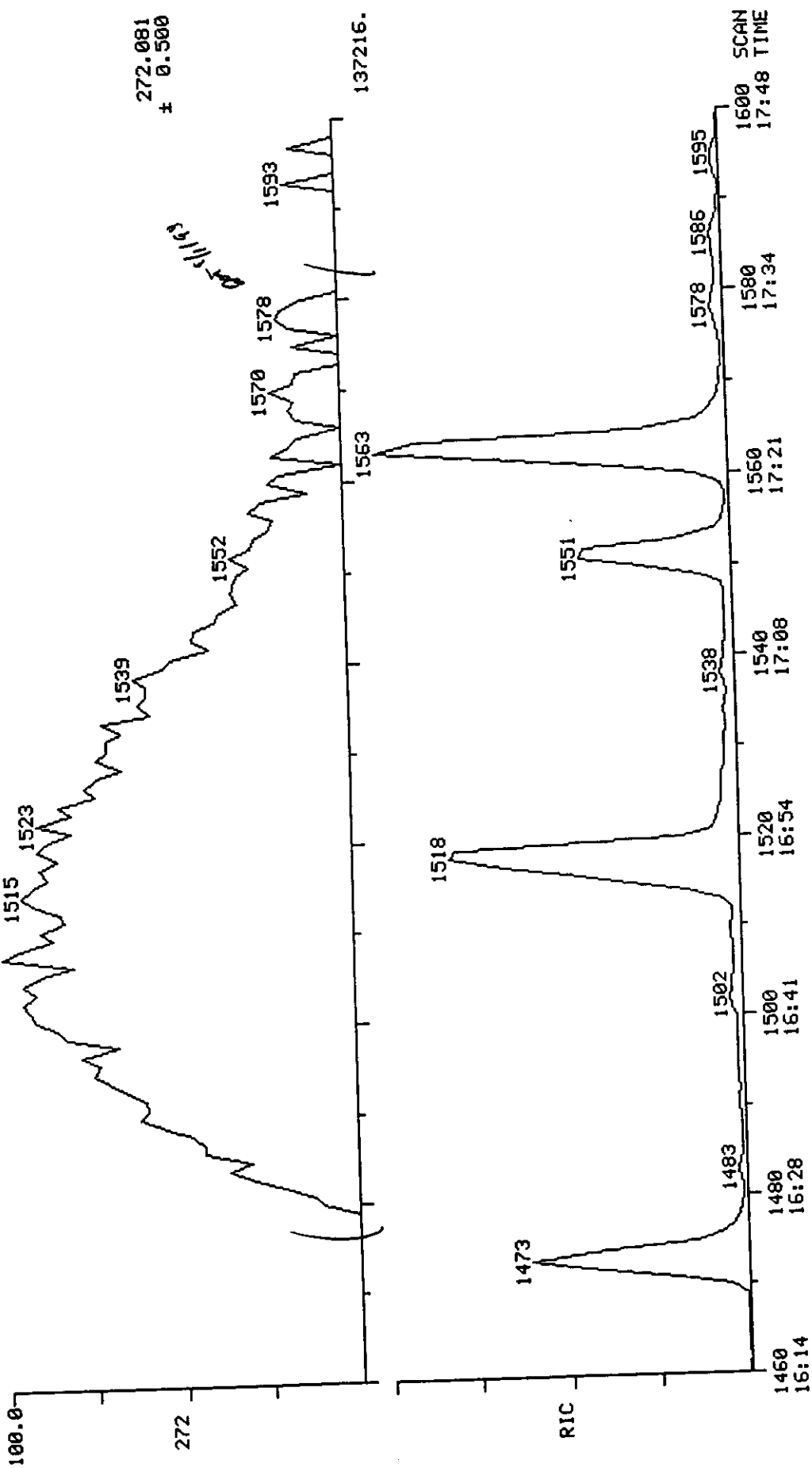
SAMPLE: 160UG/ML AP9 062598H

CONDS.: INST. ID: F16

RANGE: G 1.2158 LABEL: N 0, 2.0

QUAN: A 6, 2.0 J 0 BASE: U 20, 3

238.



Continuing Calibration

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980831 08/31/98 12:31:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

Library Entry	Compound	Initial Calibration			Continuing Calibration			
		Response Factor		% RSD	Amount		% Diff	
		Avg	Min	Max	Calc	Exp	Max	
S1 15	C310 N-NITROSODIMETH	1.019		4.4	69.65	80.00	12.9	50.0
S1 100	C320 ANILINE	1.846		6.2	81.93	80.00	-2.4	50.0
S1 105	C315 PHENOL	1.968		7.8	84.53	80.00	-5.7	20.0
S1 110	C325 BIS(2-CHLOROETH	1.368		6.1	83.20	80.00	-4.0	50.0
S1 115	C330 2-CHLOROPHENOL	1.497		5.5	85.59	80.00	-7.0	50.0
S1 125	C335 1,3-DICHLOROBEN	1.586		4.5	83.03	80.00	-3.8	50.0
S1 130	C340 1,4-DICHLOROBEN	1.618		4.8	83.73	80.00	-4.7	20.0
S1 145	C345 BENZYL ALCOHOL	0.854		9.3	78.77	80.00	1.5	50.0
S1 150	C350 1,2-DICHLOROBEN	1.500		5.0	82.86	80.00	-3.6	50.0
S1 160	C355 2-METHYLPHENOL	1.174		7.4	79.35	80.00	0.8	50.0
S1 165	C360 2,2'-OXYBIS(1-C	2.539		3.9	79.64	80.00	0.4	50.0
S1 185	C365 4-METHYLPHENOL	1.276		9.0	77.91	80.00	2.6	50.0
S1 190	C370 N-NITROSO-D1-N-	0.906	0.050	5.1	79.52	80.00	0.6	50.0
S1 200	C375 HEXACHLOROETHAN	0.708		3.5	81.62	80.00	-2.0	50.0
S2 10	C410 NITROBENZENE	0.445		7.2	85.86	80.00	-7.3	50.0
S2 20	C415 ISOPHORONE	0.742		5.5	79.94	80.00	0.1	50.0
S2 25	C420 2-NITROPHENOL	0.213		7.9	84.24	80.00	-5.3	20.0
S2 30	C425 2,4-DIMETHYLPHE	0.330		16.9	92.67	80.00	-15.8	50.0
S2 45	C430 BENZOIC ACID	0.276		4.7	48.36	80.00	39.5	50.0
S2 35	C435 BIS(2-CHLOROETH	0.511		4.1	81.27	80.00	-1.6	50.0
S2 40	C440 2,4-DICHLOROPHE	0.308		6.6	83.17	80.00	-4.0	20.0
S2 55	C445 1,2,4-TRICHLORO	0.331		5.0	84.22	80.00	-5.3	50.0
S2 60	C450 NAPHTHALENE	1.022		6.6	84.81	80.00	-6.0	50.0
S2 80	C455 4-CHLOROANILINE	0.454		4.5	82.23	80.00	-2.8	50.0
S2 95	C460 HEXACHLOROBUTAD	0.168		5.1	84.88	80.00	-6.1	20.0
S2 130	C465 4-CHLORO-3-METH	0.338		9.0	82.44	80.00	-3.0	20.0
S2 145	C470 2-METHYLNAPHTHA	0.751		10.8	104.45	80.00	-30.6	50.0
S3 20	C510 HEXACHLOROCYCLO	0.372	0.050	10.1	88.84	80.00	-11.0	50.0
S3 25	C515 2,4,6-TRICHLORO	0.404		10.6	84.50	80.00	-5.6	20.0
S3 30	C520 2,4,5-TRICHLORO	0.412		7.1	80.97	80.00	-1.2	50.0
S3 40	C525 2-CHLORONAPHTHA	1.193		6.6	86.65	80.00	-8.3	50.0
S3 45	C530 2-NITROANILINE	0.502		2.9	83.54	80.00	-4.4	50.0
S3 55	C535 DIMETHYLPHTHALA	1.341		6.1	85.06	80.00	-6.3	50.0
S3 65	C540 ACENAPHTHYLENE	1.839		8.2	87.87	80.00	-9.8	50.0
S3 70	C543 2,6-DINITROTOLU	0.324		8.5	90.03	80.00	-12.5	50.0
S3 75	C545 3-NITROANILINE	0.383		1.4	83.67	80.00	-4.6	50.0
S3 80	C550 ACENAPHTHENE	1.158		6.9	86.87	80.00	-8.6	20.0
S3 85	C555 2,4-DINITROPHEN	0.189	0.050	10.3	84.84	80.00	-6.0	50.0

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980831 08/31/98 12:31:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

Library Entry Compound	Initial Calibration			Continuing Calibration		
	Response Factor		% RSD	Amount		% Diff
	Avg	Min	Max	Calc	Exp	Max
S3 90 C565 DIBENZOFURAN	1.669		6.2	86.67	80.00	-8.3 50.0
S3 105 C570 2,4-DINITROTOLU	0.397		8.8	95.11	80.00	-18.9 50.0
S3 130 C580 DIETHYLPHTHALAT	1.369		7.7	88.14	80.00	-10.2 50.0
S3 140 C585 4-CHLOROPHENYL-	0.613		5.4	86.06	80.00	-7.6 50.0
S3 135 C590 FLUORENE	1.298		6.8	88.29	80.00	-10.4 50.0
S3 150 C595 4-NITROANALINE	0.307		8.4	83.02	80.00	-3.8 50.0
S4 10 C610 4,6-DINITRO-2-M	0.132		7.6	83.52	80.00	-4.4 50.0
S4 15 C615 N-NITROSODIPHEN	0.551		4.2 30.0	77.22	80.00	3.5 20.0
S4 20 C620 AZOBENZENE	1.171		3.9	78.58	80.00	1.8 50.0
S4 30 C625 4-BROMOPHENYL-P	0.223		3.9	82.03	80.00	-2.5 50.0
S4 40 C630 HEXACHLOROBENZE	0.260		6.7	81.39	80.00	-1.7 50.0
S4 50 C635 PENTACHLOROPHEN	0.155		5.9 30.0	80.24	80.00	-0.3 20.0
S4 65 C640 PHENANTHRENE	1.033		5.2	83.59	80.00	-4.5 50.0
S4 70 C645 ANTHRACENE	0.987		11.0	84.41	80.00	-5.5 50.0
S4 80 C647 CARBAZOLE	0.764		10.3	80.70	80.00	-0.9 50.0
S4 85 C650 DI-N-BUTYLPHTHA	1.247		9.4	85.76	80.00	-7.2 50.0
S4 110 C655 FLUORANTHENE	0.900		13.5 30.0	82.37	80.00	-3.0 20.0
S5 10 C710 BENZIDINE	0.129		23.7	34.62	80.00	56.7* 50.0
S5 15 C715 PYRENE	1.323		6.5	87.30	80.00	-9.1 50.0
S5 40 C720 BUTYLBENZYLPHTH	0.685		7.7	82.69	80.00	-3.4 50.0
S5 55 C725 3,3'-DICHLORBE	0.408		5.7	73.23	80.00	8.5 50.0
S5 50 C730 BENZO(A)ANTHRAC	1.125		5.4	86.19	80.00	-7.7 50.0
S5 60 C740 CHRYSENE	1.014		6.7	82.50	80.00	-3.1 50.0
S5 65 C745 BIS(2-ETHYLHEXY	0.980		9.4	83.00	80.00	-3.8 50.0
S6 10 C760 DI-N-OCTYL PHTH	1.925		7.4 30.0	85.61	80.00	-7.0 20.0
S6 15 C765 BENZO(B)FLUORAN	1.444		5.0	78.88	80.00	1.4 50.0
S6 25 C770 BENZO(K)FLUORAN	1.159		9.8	90.18	80.00	-12.7 50.0
S6 35 C775 BENZO(A)PYRENE	1.182		5.3 30.0	83.29	80.00	-4.1 20.0
S6 55 C780 INDENO(1,2,3-CD	1.279		6.6	85.08	80.00	-6.3 50.0
S6 60 C785 DIBENZ(A,H)ANTH	1.267		8.1	84.76	80.00	-5.9 50.0
S6 65 C790 BENZO(G,H,I)PER	1.331		8.2	85.64	80.00	-7.1 50.0
S1 6 CS75 1,2-DICHLOROBNZ	0.920		6.5	81.28	80.00	-1.6 50.0
S2 2 CS20 NITROBENZENE-D5	0.470		5.2	82.78	80.00	-3.5 50.0
S3 3 CS25 2-FLUOROBIPHENY	1.234		6.2	84.73	80.00	-5.9 50.0
S5 2 CS30 TERPHENYL-D14	0.995		5.3	86.97	80.00	-8.7 50.0
S1 3 CS45 PHENOL-D5	2.026		2.0	79.53	80.00	0.6 50.0
S1 2 CS50 2-FLUOROPHENOL	1.368		4.4	76.35	80.00	4.6 50.0
S1 5 CS70 2-CHLOROPHENOL-	1.482		4.1	80.04	80.00	-0.1 50.0
S3 2 CS55 2,4,6-TRIBROMOP	0.236		6.9	81.97	80.00	-2.5 50.0

2.6% of the compounds exceed the %RSD value  
of 15.0. 2 of 78 compounds are out.

0 CCC cmpds out  
1 of nonCCC exceed 50 %

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980831 08/31/98 12:31:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
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 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

ST16980831 ST16980827A ST16980827B ST16980827C ST16980827 ST16980827D  
 ST16980827E

Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S1 15	80,	0.887	10,	0.952	20,	1.010	50,	1.016	80,	1.087
S1 15	160,	1.042							120,	1.008
S1 100	80,	1.891	10,	1.947	20,	1.988	50,	1.814	80,	1.876
S1 100	160,	1.674							120,	1.778
S1 105	80,	2.079	10,	1.699	20,	2.037	50,	2.035	80,	2.148
S1 105	160,	1.982							120,	1.907
S1 110	80,	1.422	10,	1.336	20,	1.465	50,	1.438	80,	1.414
S1 110	160,	1.261							120,	1.291
S1 115	80,	1.601	10,	1.386	20,	1.569	50,	1.522	80,	1.603
S1 115	160,	1.451							120,	1.450
S1 125	80,	1.646	10,	1.515	20,	1.668	50,	1.618	80,	1.659
S1 125	160,	1.510							120,	1.546
S1 130	80,	1.693	10,	1.580	20,	1.695	50,	1.666	80,	1.697
S1 130	160,	1.518							120,	1.549
S1 145	80,	0.841	10,	0.702	20,	0.851	50,	0.872	80,	0.913
S1 145	160,	0.922							120,	0.866
S1 150	80,	1.553	10,	1.475	20,	1.606	50,	1.482	80,	1.573
S1 150	160,	1.409							120,	1.454
S1 160	80,	1.165	10,	0.998	20,	1.206	50,	1.221	80,	1.209
S1 160	160,	1.213							120,	1.199
S1 165	80,	2.528	10,	2.393	20,	2.676	50,	2.597	80,	2.582
S1 165	160,	2.486							120,	2.502
S1 185	80,	1.243	10,	1.049	20,	1.298	50,	1.355	80,	1.287
S1 185	160,	1.351							120,	1.316
S1 190	80,	0.900	10,	0.819	20,	0.934	50,	0.938	80,	0.940
S1 190	160,	0.890							120,	0.913
S1 200	80,	0.723	10,	0.665	20,	0.740	50,	0.713	80,	0.720
S1 200	160,	0.706							120,	0.706
S2 10	80,	0.478	10,	0.423	20,	0.477	50,	0.455	80,	0.486
S2 10	160,	0.410							120,	0.419
S2 20	80,	0.741	10,	0.685	20,	0.778	50,	0.774	80,	0.778
S2 20	160,	0.708							120,	0.726

S2	25	80, 0.225	10, 0.182	20, 0.212	50, 0.218	80, 0.232	120, 0.219
S2	25	160, 0.217					
S2	30	80, 0.382	10, 0.234	20, 0.309	50, 0.334	80, 0.400	120, 0.349
S2	30	160, 0.353					
S2	45	80, 0.167		20, 0.257	50, 0.279	80, 0.270	120, 0.280
S2	45	160, 0.292					
S2	35	80, 0.519	10, 0.481	20, 0.529	50, 0.517	80, 0.538	120, 0.502
S2	35	160, 0.498					
S2	40	80, 0.320	10, 0.270	20, 0.313	50, 0.314	80, 0.332	120, 0.311
S2	40	160, 0.306					
S2	55	80, 0.348	10, 0.319	20, 0.350	50, 0.334	80, 0.350	120, 0.322
S2	55	160, 0.310					
S2	60	80, 1.084	10, 1.008	20, 1.090	50, 1.038	80, 1.102	120, 0.960
S2	60	160, 0.935					
S2	80	80, 0.466	10, 0.419	20, 0.466	50, 0.454	80, 0.480	120, 0.449
S2	80	160, 0.455					
S2	95	80, 0.178	10, 0.160	20, 0.178	50, 0.170	80, 0.177	120, 0.167
S2	95	160, 0.157					
S2	130	80, 0.348	10, 0.279	20, 0.337	50, 0.353	80, 0.365	120, 0.347
S2	130	160, 0.346					
S2	145	80, 0.981	10, 0.773	20, 0.840	50, 0.827	80, 0.757	120, 0.671
S2	145	160, 0.639					
S3	20	80, 0.413	10, 0.329	20, 0.387	50, 0.376	80, 0.428	120, 0.379
S3	20	160, 0.330					
S3	25	80, 0.426	10, 0.327	20, 0.389	50, 0.405	80, 0.435	120, 0.417
S3	25	160, 0.448					
S3	30	80, 0.417	10, 0.397	20, 0.437	50, 0.429	80, 0.443	120, 0.401
S3	30	160, 0.366					
S3	40	80, 1.292	10, 1.172	20, 1.275	50, 1.208	80, 1.283	120, 1.128
S3	40	160, 1.089					
S3	45	80, 0.524		20, 0.491	50, 0.510	80, 0.523	120, 0.487
S3	45	160, 0.497					
S3	55	80, 1.426	10, 1.257	20, 1.423	50, 1.392	80, 1.426	120, 1.297
S3	55	160, 1.251					
S3	65	80, 2.019	10, 1.751	20, 1.959	50, 1.896	80, 2.047	120, 1.707
S3	65	160, 1.671					
S3	70	80, 0.364	10, 0.274	20, 0.339	50, 0.341	80, 0.351	120, 0.321
S3	70	160, 0.317					
S3	75	80, 0.400		20, 0.382	50, 0.392	80, 0.382	120, 0.379
S3	75	160, 0.379					
S3	80	80, 1.258	10, 1.140	20, 1.227	50, 1.182	80, 1.261	120, 1.075

S3 80 160, 1.066

S3 85 80, 0.201

S3 85 160, 0.196

S3 95 80, 0.174

S3 95 160, 0.169

20, 0.156 50, 0.190 80, 0.202 120, 0.203

20, 0.164 50, 0.178 80, 0.184 120, 0.168

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980831 08/31/98 12:31:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

ST16980831 ST16980827A ST16980827B ST16980827C ST16980827 ST16980827D  
 ST16980827E

Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S3 90	80,	1.808	10,	1.617	20,	1.782	50,	1.736	80,	1.762	120,	1.556
S3 90	160,	1.561										
S3 105	80,	0.472	10,	0.339	20,	0.412	50,	0.420	80,	0.437	120,	0.396
S3 105	160,	0.378										
S3 130	80,	1.508	10,	1.300	20,	1.462	50,	1.452	80,	1.462	120,	1.328
S3 130	160,	1.209										
S3 140	80,	0.659	10,	0.597	20,	0.647	50,	0.631	80,	0.646	120,	0.579
S3 140	160,	0.575										
S3 135	80,	1.432	10,	1.251	20,	1.398	50,	1.340	80,	1.386	120,	1.224
S3 135	160,	1.188										
S3 150	80,	0.319			20,	0.326	50,	0.320	80,	0.306	120,	0.263
S3 150	160,	0.321										
S4 10	80,	0.138			20,	0.121	50,	0.139	80,	0.145	120,	0.124
S4 10	160,	0.130										
S4 15	80,	0.532	10,	0.568	20,	0.582	50,	0.562	80,	0.539	120,	0.530
S4 15	160,	0.523										
S4 20	80,	1.150	10,	1.150	20,	1.222	50,	1.186	80,	1.212	120,	1.098
S4 20	160,	1.158										
S4 30	80,	0.228	10,	0.224	20,	0.231	50,	0.223	80,	0.232	120,	0.215
S4 30	160,	0.210										
S4 40	80,	0.264	10,	0.273	20,	0.281	50,	0.255	80,	0.268	120,	0.238
S4 40	160,	0.242										
S4 50	80,	0.156			20,	0.148	50,	0.156	80,	0.170	120,	0.155
S4 50	160,	0.147										
S4 65	80,	1.079	10,	1.017	20,	1.081	50,	1.044	80,	1.096	120,	0.948
S4 65	160,	1.010										
S4 70	80,	1.041	10,	1.019	20,	1.100	50,	1.026	80,	1.059	120,	0.908
S4 70	160,	0.809										
S4 80	80,	0.771	10,	0.914	20,	0.763	50,	0.729	80,	0.769	120,	0.707
S4 80	160,	0.701										
S4 85	80,	1.337	10,	1.233	20,	1.366	50,	1.332	80,	1.333	120,	1.137
S4 85	160,	1.080										



S4	110	80, 0.927	10, 1.013	20, 1.008	50, 0.925	80, 0.950	120, 0.788
S4	110	160, 0.715					
S5	10	80, 0.056		20, 0.155	50, 0.143	80, 0.077	120, 0.141
S5	10	160, 0.131					
S5	15	80, 1.443	10, 1.209	20, 1.310	50, 1.295	80, 1.468	120, 1.295
S5	15	160, 1.358					
S5	40	80, 0.708	10, 0.587	20, 0.673	50, 0.692	80, 0.722	120, 0.700
S5	40	160, 0.734					
S5	55	80, 0.374		20, 0.441	50, 0.421	80, 0.383	120, 0.405
S5	55	160, 0.391					
S5	50	80, 1.212	10, 1.058	20, 1.195	50, 1.116	80, 1.203	120, 1.086
S5	50	160, 1.092					
S5	60	80, 1.046	10, 1.034	20, 1.068	50, 1.005	80, 1.102	120, 0.959
S5	60	160, 0.919					
S5	65	80, 1.017	10, 0.812	20, 0.978	50, 0.986	80, 1.027	120, 0.987
S5	65	160, 1.089					
S6	10	80, 2.060	10, 1.637	20, 1.959	50, 1.985	80, 1.977	120, 1.967
S6	10	160, 2.023					
S6	15	80, 1.423	10, 1.322	20, 1.536	50, 1.448	80, 1.453	120, 1.490
S6	15	160, 1.412					
S6	25	80, 1.306	10, 1.185	20, 1.223	50, 1.171	80, 1.264	120, 1.171
S6	25	160, 0.940					
S6	35	80, 1.230	10, 1.168	20, 1.257	50, 1.206	80, 1.229	120, 1.147
S6	35	160, 1.084					
S6	55	80, 1.360	10, 1.253	20, 1.355	50, 1.312	80, 1.368	120, 1.241
S6	55	160, 1.144					
S6	60	80, 1.343	10, 1.254	20, 1.386	50, 1.288	80, 1.360	120, 1.212
S6	60	160, 1.105					
S6	65	80, 1.425	10, 1.339	20, 1.436	50, 1.356	80, 1.426	120, 1.289
S6	65	160, 1.139					
S1	6	80, 0.935	10, 0.832	20, 0.999	50, 0.946	80, 0.959	120, 0.902
S1	6	160, 0.882					
S2	2	80, 0.486	10, 0.446	20, 0.494	50, 0.469	80, 0.504	120, 0.452
S2	2	160, 0.452					
S3	3	80, 1.307	10, 1.252	20, 1.315	50, 1.253	80, 1.299	120, 1.171
S3	3	160, 1.116					
S5	2	80, 1.082	10, 0.932	20, 1.002	50, 0.976	80, 1.082	120, 0.960
S5	2	160, 1.019					
S1	3	80, 2.014		20, 2.057	50, 2.059	80, 2.046	120, 1.995
S1	3	160, 1.971					
S1	2	80, 1.306		20, 1.291	50, 1.332	80, 1.448	120, 1.376

S1 2 160, 1.394

S1 5 80, 1.483

S1 5 160, 1.407

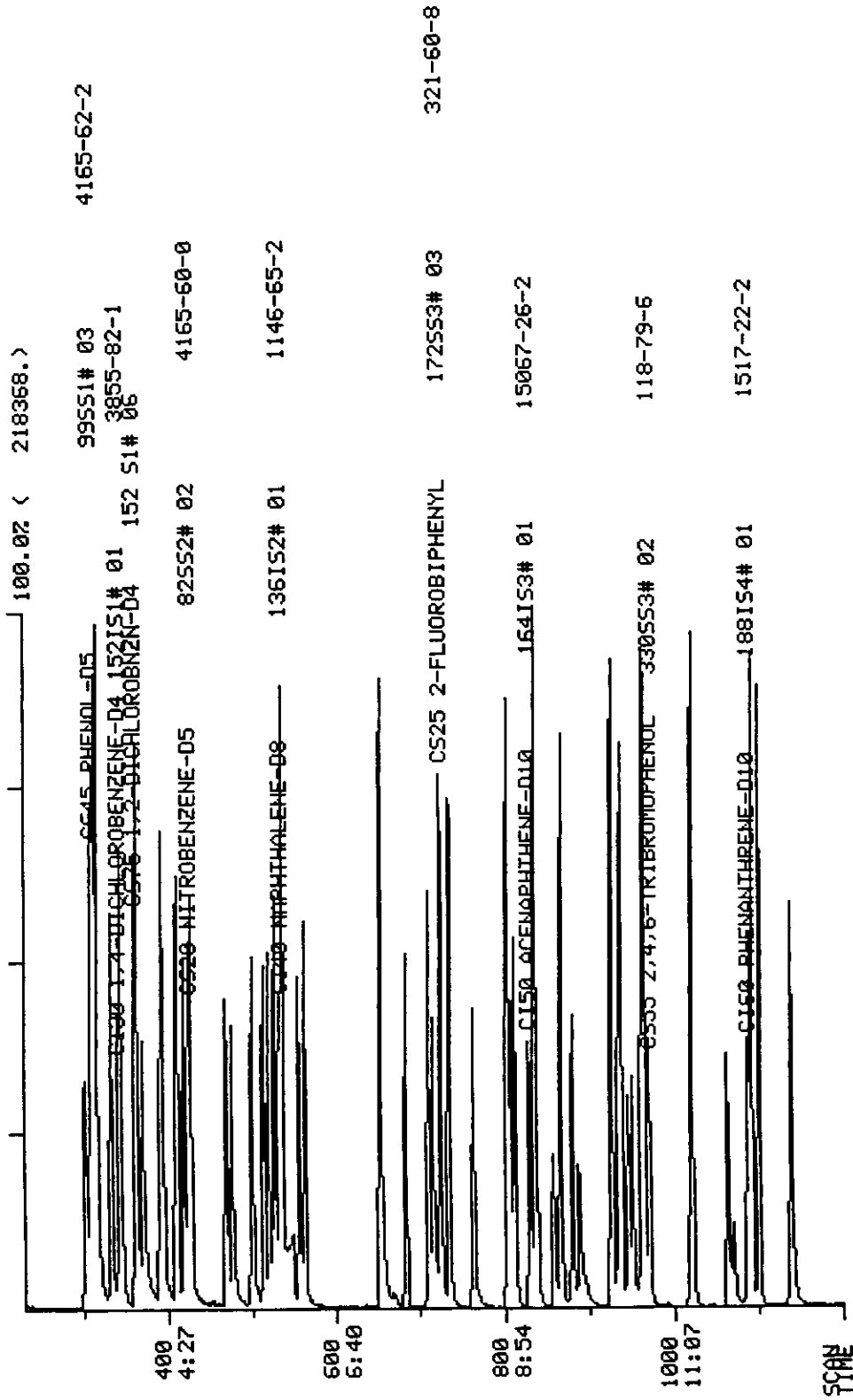
20, 1.561 50, 1.501 80, 1.503 120, 1.437

S3 2 80, 0.241

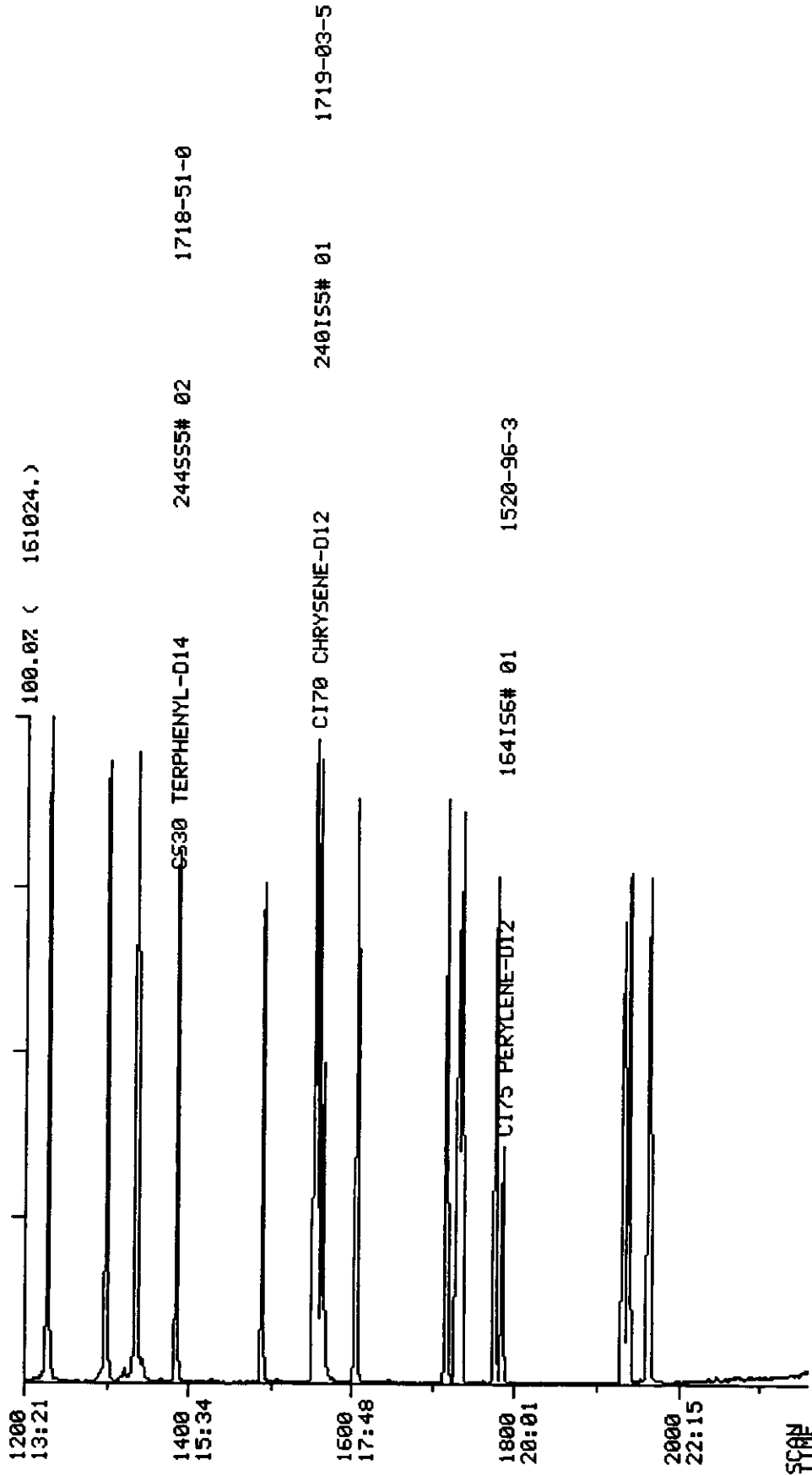
S3 2 160, 0.217

20, 0.261 50, 0.239 80, 0.233 120, 0.228

DATA FROM FILE: ST16980831 SCANS 230 TO 1200 ACQUIRED: 08/31/98 12:31:00  
 CALI: ST16980831 #3  
 SAMPLE: S5TD080 80UG/ML CALI III 082198A  
 CONDS.: INST. ID: F16



DATA FROM FILE: ST16980831 SCANS 1200 TO 2158 ACQUIRED: 08/31/98 12:31:00  
CALI: ST16980831 #3  
SAMPLE: SSTD080 80UG/ML CALI III 082198A  
COND.: INST. ID: F15



Data: ST16980831.TI

08/31/98 12:31:00

Sample: SSTD080 80UG/ML CALI III 082198A

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from average of whole .RL

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	C170 CHRYSENE-D12	240IS5# 01	1719-03-5
6	C175 PERYLENE-D12	164IS6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBENZENE-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANILINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANILINE	65 S3# 45	88-74-4

No	Name		
48	C535 DIMETHYLPHthalate	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROToluene	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	28649.	40.000 UG/ML	0.60
2	136	530	5:54	2	1.000	A BB	100432.	40.000 UG/ML	0.60
3	164	828	9:13	3	1.000	A BB	54347.	40.000 UG/ML	0.60
4	188	1086	12:05	4	1.000	A BB	98520.	40.000 UG/ML	0.60
5	240	1552	17:16	5	1.000	A BB	62289.	40.000 UG/ML	0.60
6	264	1783	19:50	6	1.000	A BB	50750.	40.000 UG/ML	0.60
7	82	422	4:42	2	0.796	A BB	97588.	82.774 UG/ML	1.24
8	172	723	8:03	3	0.873	A BB	142092.	84.729 UG/ML	1.27
9	244	1383	15:23	5	0.891	A BB	134779.	86.967 UG/ML	1.30
10	99	308	3:26	1	0.906	A BB	115381.	79.535 UG/ML	1.19
11	112	196	2:11	1	0.576	A BB	74816.	76.346 UG/ML	1.14
12	330	969	10:47	3	1.170	A BB	26238.	81.946 UG/ML	1.23
13	132	312	3:28	1	0.918	A BB	84951.	80.059 UG/ML	1.20
14	152	359	4:00	1	1.056	A BB	53560.	81.280 UG/ML	1.22
15	NOT FOUND								
16	74	103	1:09	1	0.303	A BB	50841.	69.648 UG/ML	1.04
17	93	301	3:21	1	0.885	A BV	108330.	81.918 UG/ML	1.23
18	94	309	3:26	1	0.909	A VB	119147.	84.537 UG/ML	1.27
19	93	315	3:30	1	0.926	A VB	81492.	83.187 UG/ML	1.25
20	128	314	3:30	1	0.924	A BB	91760.	85.590 UG/ML	1.28
21	146	331	3:41	1	0.974	A BV	94316.	83.030 UG/ML	1.24
22	146	342	3:48	1	1.006	A VB	97001.	83.731 UG/ML	1.25
23	108	369	4:06	1	1.085	A BB	48199.	78.765 UG/ML	1.18
24	146	361	4:01	1	1.062	A BB	89006.	82.857 UG/ML	1.24
25	108	390	4:20	1	1.147	A BB	66743.	79.332 UG/ML	1.19
26	45	389	4:20	1	1.144	A BB	144852.	79.645 UG/ML	1.19
27	108	417	4:38	1	1.226	A BB	71198.	77.917 UG/ML	1.17
28	70	410	4:34	1	1.206	A BV	51578.	79.498 UG/ML	1.19
29	117	407	4:32	1	1.197	A BB	41410.	81.636 UG/ML	1.22
30	77	425	4:44	2	0.802	A VB	95928.	85.850 UG/ML	1.29
31	82	467	5:12	2	0.881	A BB	148832.	79.956 UG/ML	1.20
32	139	474	5:16	2	0.894	A VB	45121.	84.179 UG/ML	1.26
33	107	497	5:32	2	0.938	A BB	76740.	92.634 UG/ML	1.39
34	93	511	5:41	2	0.964	A BB	104239.	81.300 UG/ML	1.22
35	162	517	5:45	2	0.975	A BB	64246.	83.195 UG/ML	1.25
36	122	546	6:04	2	1.030	A BB	33466.	48.330 UG/ML	0.72
37	180	524	5:50	2	0.989	A BB	69958.	84.196 UG/ML	1.26
38	128	533	5:56	2	1.006	A BB	217673.	84.809 UG/ML	1.27
39	127	553	6:09	2	1.043	A BB	93696.	82.234 UG/ML	1.23
40	225	560	6:14	2	1.057	A BB	35840.	84.907 UG/ML	1.27
41	107	649	7:13	2	1.225	A BB	69925.	82.458 UG/ML	1.23
42	142	651	7:14	2	1.228	A BV	197003.	104.439 UG/ML	1.56
43	237	680	7:34	3	0.821	A BB	44842.	88.855 UG/ML	1.33
44	196	708	7:53	3	0.855	A BV	46323.	84.486 UG/ML	1.27
45	196	713	7:56	3	0.861	A VB	45345.	80.983 UG/ML	1.21
46	162	733	8:09	3	0.885	A BB	140394.	86.659 UG/ML	1.30
47	65	762	8:29	3	0.920	A BB	56935.	83.499 UG/ML	1.25
48	163	806	8:58	3	0.973	A BB	154975.	85.067 UG/ML	1.27
49	152	802	8:55	3	0.969	A BB	219486.	87.869 UG/ML	1.32
50	165	812	9:02	3	0.981	A BB	39614.	90.042 UG/ML	1.35

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:16	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:50	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:42	1.00	0.796	1.00	82.77	50.00	0.777	0.470	1.66
8	8:03	1.00	0.873	1.00	84.73	50.00	2.092	1.234	1.69
9	15:23	1.00	0.891	1.00	86.97	50.00	1.731	0.995	1.74
10	3:26	1.00	0.906	1.00	79.53	50.00	3.222	2.025	1.59
11	2:11	1.00	0.576	1.00	76.35	50.00	2.089	1.368	1.53
12	10:47	1.00	1.170	1.00	81.95	50.00	0.386	0.236	1.64
13	3:28	1.00	0.918	1.00	80.06	50.00	2.372	1.482	1.60
14	4:00	1.00	1.056	1.00	81.28	50.00	1.496	0.920	1.63
15	12:39		0.941						
16	1:09	1.00	0.303	1.00	69.65	50.00	1.420	1.019	1.39
17	3:21	1.00	0.885	1.00	81.92	50.00	3.025	1.846	1.64
18	3:26	1.00	0.909	1.00	84.54	50.00	3.327	1.968	1.69
19	3:30	1.00	0.926	1.00	83.19	50.00	2.276	1.368	1.66
20	3:30	1.00	0.924	1.00	85.59	50.00	2.562	1.497	1.71
21	3:41	1.00	0.974	1.00	83.03	50.00	2.634	1.586	1.66
22	3:48	1.00	1.006	1.00	83.73	50.00	2.709	1.617	1.67
23	4:06	1.00	1.085	1.00	78.77	50.00	1.346	0.854	1.58
24	4:01	1.00	1.062	1.00	82.86	50.00	2.485	1.500	1.66
25	4:20	1.00	1.147	1.00	79.33	50.00	1.864	1.175	1.59
26	4:20	1.00	1.144	1.00	79.65	50.00	4.045	2.539	1.59
27	4:38	1.00	1.226	1.00	77.92	50.00	1.988	1.276	1.56
28	4:34	1.00	1.206	1.00	79.50	50.00	1.440	0.906	1.59
29	4:32	1.00	1.197	1.00	81.64	50.00	1.156	0.708	1.63
30	4:44	1.00	0.802	1.00	85.85	50.00	0.764	0.445	1.72
31	5:12	1.00	0.881	1.00	79.96	50.00	1.186	0.741	1.60
32	5:16	1.00	0.894	1.00	84.18	50.00	0.359	0.213	1.68
33	5:32	1.00	0.938	1.00	92.63	50.00	0.611	0.330	1.85
34	5:41	1.00	0.964	1.00	81.30	50.00	0.830	0.511	1.63
35	5:45	1.00	0.975	1.00	83.19	50.00	0.512	0.308	1.66
36	6:04	1.00	1.030	1.00	48.33	50.00	0.267	0.276	0.97
37	5:50	1.00	0.989	1.00	84.20	50.00	0.557	0.331	1.68
38	5:56	1.00	1.006	1.00	84.81	50.00	1.734	1.022	1.70
39	6:09	1.00	1.043	1.00	82.23	50.00	0.746	0.454	1.64
40	6:14	1.00	1.057	1.00	84.91	50.00	0.285	0.168	1.70
41	7:13	1.00	1.225	1.00	82.46	50.00	0.557	0.338	1.65
42	7:14	1.00	1.228	1.00	104.44	50.00	1.569	0.751	2.09
43	7:34	1.00	0.821	1.00	88.86	50.00	0.660	0.371	1.78
44	7:53	1.00	0.855	1.00	84.49	50.00	0.682	0.404	1.69
45	7:56	1.00	0.861	1.00	80.98	50.00	0.667	0.412	1.62
46	8:09	1.00	0.885	1.00	86.66	50.00	2.067	1.192	1.73
47	8:29	1.00	0.920	1.00	83.50	50.00	0.838	0.502	1.67
48	8:58	1.00	0.973	1.00	85.07	50.00	2.281	1.341	1.70
49	8:55	1.00	0.969	1.00	87.87	50.00	3.231	1.838	1.76
50	9:02	1.00	0.981	1.00	90.04	50.00	0.583	0.324	1.80

Data: ST16980831.TI

08/31/98 12:31:00

Sample: SSTD080 80UG/ML CALI III 082198A

Conds.: INST. ID: F16

Formula: IUL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from average of whole .RL

No	Name		
51	C545 3-NITROANILINE	138 S3# 75	99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC	83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP	51-28-5
54	C565 DIBENZOFURAN	168 S3# 90	132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP	100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105	121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130	84-66-2
58	C590 FLUORENE	166 S3#135	86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140	7005-72-3
60	C595 4-NITROANILINE	138 S3#150	100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10	534-52-1
62	C615 N-NITROSODIPHENYLAMINE	169 S4# 15CC	87-30-6
63	C620 AZOBENZENE	77 S4# 20	103-33-3
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30	101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40	118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50	87-86-5
67	C640 PHENANTHRENE	178 S4# 65	85-01-8
68	C645 ANTHRACENE	178 S4# 70	120-12-7
69	C647 CARBAZOLE	167 S4# 80	86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85	84-74-2
71	C655 FLUORANTHENE	202 S4#110CC	206-44-0
72	C710 BENZIDINE	184 S5# 10	92-81-5
73	C715 PYRENE	202 S5# 15	129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40	85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50	56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55	91-94-1
77	C740 CHRYSENE	228 S5# 60	218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65	117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC	117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15	205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25	207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35	50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55	193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60	53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65	191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	138	837	9:19	3	1.011	A BB	43517.	83.685 UG/ML	1.25
52	153	834	9:17	3	1.007	A BB	136730.	86.856 UG/ML	1.30
53	184	856	9:31	3	1.034	A BB	21832.	84.885 UG/ML	1.27
54	168	866	9:38	3	1.046	A VB	196548.	86.670 UG/ML	1.30
55	109	887	9:52	3	1.071	A BB	18892.	80.538 UG/ML	1.21
56	165	880	9:47	3	1.063	A BB	51300.	95.049 UG/ML	1.42
57	149	935	10:24	3	1.129	A BB	163921.	88.144 UG/ML	1.32
58	166	926	10:18	3	1.118	A BB	155682.	88.292 UG/ML	1.32



No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
59	204	937	10:25	3	1.132	A BB	71622.	86.032 UG/ML	1.29
60	138	946	10:31	3	1.143	A BB	34652.	83.074 UG/ML	1.24
61	198	951	10:35	4	0.876	A BB	27112.	83.379 UG/ML	1.25
62	169	962	10:42	4	0.886	A BV	104728.	77.208 UG/ML	1.16
63	77	964	10:43	4	0.888	A VB	226631.	78.592 UG/ML	1.18
64	248	1020	11:21	4	0.939	A BB	44954.	82.055 UG/ML	1.23
65	284	1019	11:20	4	0.938	A BB	52022.	81.399 UG/ML	1.22
66	266	1062	11:49	4	0.978	A BB	30674.	80.327 UG/ML	1.20
67	178	1091	12:08	4	1.005	A BV	212616.	83.599 UG/ML	1.25
68	178	1100	12:14	4	1.013	A VB	205171.	84.402 UG/ML	1.26
69	167	1138	12:39	4	1.048	A BB	151821.	80.691 UG/ML	1.21
70	149	1227	13:39	4	1.130	A BB	263375.	85.782 UG/ML	1.28
71	202	1298	14:26	4	1.195	A BB	182566.	82.359 UG/ML	1.23
72	184	1341	14:55	5	0.864	A BB	6976.	34.563 UG/ML	0.52
73	202	1335	14:51	5	0.860	A BB	179798.	87.298 UG/ML	1.31
74	149	1488	16:33	5	0.959	A BV	88160.	82.691 UG/ML	1.24
75	228	1551	17:15	5	0.999	A BV	150997.	86.203 UG/ML	1.29
76	252	1562	17:22	5	1.006	A BB	46552.	73.268 UG/ML	1.10
77	228	1557	17:19	5	1.003	A VB	130339.	82.506 UG/ML	1.24
78	149	1603	17:50	5	1.033	A BB	126644.	83.005 UG/ML	1.24
79	149	1715	19:05	6	0.962	A BV	209065.	85.620 UG/ML	1.28
80	252	1728	19:13	6	0.969	A BV	144473.	78.892 UG/ML	1.18
81	252	1733	19:17	6	0.972	A VB	132603.	90.166 UG/ML	1.35
82	252	1775	19:45	6	0.996	A BB	124893.	83.293 UG/ML	1.25
83	276	1930	21:28	6	1.082	A BV	138037.	85.069 UG/ML	1.27
84	278	1937	21:33	6	1.086	A BB	136299.	84.763 UG/ML	1.27
85	276	1962	21:49	6	1.100	A BB	144611.	85.643 UG/ML	1.28

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:19	1.00	1.011	1.00	83.68	50.00	0.641	0.383	1.67
52	9:17	1.00	1.007	1.00	86.86	50.00	2.013	1.159	1.74
53	9:31	1.00	1.034	1.00	84.89	50.00	0.321	0.189	1.70
54	9:38	1.00	1.046	1.00	86.67	50.00	2.893	1.669	1.73
55	9:52	1.00	1.071	1.00	80.54	50.00	0.278	0.173	1.61
56	9:47	1.00	1.063	1.00	95.05	50.00	0.755	0.397	1.90
57	10:24	1.00	1.129	1.00	88.14	50.00	2.413	1.369	1.76
58	10:18	1.00	1.118	1.00	88.29	50.00	2.292	1.298	1.77
59	10:25	1.00	1.132	1.00	86.03	50.00	1.054	0.613	1.72
60	10:31	1.00	1.143	1.00	83.07	50.00	0.510	0.307	1.66
61	10:35	1.00	0.876	1.00	83.38	50.00	0.220	0.132	1.67
62	10:42	1.00	0.886	1.00	77.21	50.00	0.850	0.551	1.54
63	10:43	1.00	0.888	1.00	78.59	50.00	1.840	1.171	1.57
64	11:21	1.00	0.939	1.00	82.05	50.00	0.365	0.222	1.64
65	11:20	1.00	0.938	1.00	81.40	50.00	0.422	0.259	1.63
66	11:49	1.00	0.978	1.00	80.33	50.00	0.249	0.155	1.61
67	12:08	1.00	1.005	1.00	83.60	50.00	1.726	1.033	1.67
68	12:14	1.00	1.013	1.00	84.40	50.00	1.666	0.987	1.69
69	12:39	1.00	1.048	1.00	80.69	50.00	1.233	0.764	1.61
70	13:39	1.00	1.130	1.00	85.78	50.00	2.139	1.247	1.72
71	14:26	1.00	1.195	1.00	82.36	50.00	1.482	0.900	1.65
72	14:55	1.00	0.864	1.00	34.56	50.00	0.090	0.130	0.69
73	14:51	1.00	0.860	1.00	87.30	50.00	2.309	1.323	1.75
74	16:33	1.00	0.959	1.00	82.69	50.00	1.132	0.685	1.65
75	17:15	1.00	0.999	1.00	86.20	50.00	1.939	1.125	1.72
76	17:22	1.00	1.006	1.00	73.27	50.00	0.598	0.408	1.47
77	17:19	1.00	1.003	1.00	82.51	50.00	1.674	1.014	1.65

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:50	1.00	1.033	1.00	83.00	50.00	1.627	0.980	1.66
79	19:05	1.00	0.962	1.00	85.62	50.00	3.296	1.925	1.71
80	19:13	1.00	0.969	1.00	78.89	50.00	2.277	1.443	1.58
81	19:17	1.00	0.972	1.00	90.17	50.00	2.090	1.159	1.80
82	19:45	1.00	0.996	1.00	83.29	50.00	1.969	1.182	1.67
83	21:28	1.00	1.082	1.00	85.07	50.00	2.176	1.279	1.70
84	21:33	1.00	1.086	1.00	84.76	50.00	2.149	1.267	1.70
85	21:49	1.00	1.100	1.00	85.64	50.00	2.280	1.331	1.71

QUANTERRA GC/MS  
SEMIVOLATILE APPENDIX IX CALIBRATION CHE

CONTIN. CALIBRATION: 80 UG/ML APPENDIX ST16980831A 08/31/98 13:07:00  
 INITIAL CALIBRATION: 10 UG/ML APPENDIX ST16980831B 08/31/98 13:35:00  
 INITIAL CALIBRATION: 20 UG/ML APPENDIX ST16980831C 08/31/98 14:02:00  
 INITIAL CALIBRATION: 50 UG/ML APPENDIX ST16980831D 08/31/98 14:32:00  
 INITIAL CALIBRATION: 80 UG/ML APPENDIX ST16980831A 08/31/98 13:07:00  
 INITIAL CALIBRATION: 120 UG/ML APPENDIX ST16980831E 08/31/98 15:02:00  
 INITIAL CALIBRATION: 160 UG/ML APPENDIX ST16980831F 08/31/98 15:31:00

Library Entry	Compound	Initial Calibration			Continuing Calibration		
		Response Factor	% RSD		Amount		% Diff
		Avg	Min	Max	Calc	Exp	Max
S1	20 PYRIDINE	1.590		6.5	79.82	80.00	0.2 50.0
S1	30 2-PICOLINE	1.565		1.3	80.59	80.00	-0.7 50.0
S1	40 N-NITROSOMETHYLETHYL	0.811		9.2	86.37	80.00	-8.0 50.0
S1	60 METHYLMETHANESULFONA	0.636		6.2	75.73	80.00	5.3 50.0
S1	70 N-NITROSODIETHYLAMIN	0.521		7.6	84.19	80.00	-5.2 50.0
S1	85 ETHYLMETHANE SULFONA	1.109		11.0	84.32	80.00	-5.4 50.0
S1	95 PENTACHLOROETHANE	0.602		1.4	80.36	80.00	-0.5 50.0
S1	170 C361 ACETOPHENONE	1.934		2.8	80.44	80.00	-0.5 50.0
S1	175 N-NITROSPYRROLIDINE	0.553		6.5	81.36	80.00	-1.7 50.0
S1	180 N-NITROSOMORPHOLINE	0.821		1.4	78.49	80.00	1.9 50.0
S1	182 3-METHYL PHENOL	2.252		9.9	85.54	80.00	-6.9 50.0
S1	195 O-TOLUIDINE	2.160		9.2	78.86	80.00	1.4 50.0
S2	15 N-NITROSOPIPERIDINE	0.321		4.9	81.55	80.00	-1.9 50.0
S2	50 AA-DIMETHYLPHENETHYL	1.285		15.6	64.54	80.00	19.3 50.0
S2	85 2,6-DICHLOROPHENOL	0.257		12.8	86.94	80.00	-8.7 50.0
S2	90 HEXACHLOROPROPENE	0.198		0.9	79.65	80.00	0.4 50.0
S2	115 P-PHENYLENE DIAMINE	0.131		42.2	22.98	80.00	71.3* 50.0
S2	120 N-NITROSODI-N-BUTYLA	0.213		7.2	84.49	80.00	-5.6 50.0
S2	140 SAFROLE	0.248		0.9	80.91	80.00	-1.1 50.0
S3	10 1,2,4,5-TETRACHLOROB	0.433		4.4	79.47	80.00	0.7 50.0
S3	15 ISOSAFROLE (#1)	0.044		3.4	78.49	80.00	1.9 50.0
S3	35 ISOSAFROLE (#2)	0.195		2.2	80.49	80.00	-0.6 50.0
S3	42 1-CHLORONAPHTH	1.318		16.5	92.76	80.00	-16.0 50.0
S3	50 1,4-NAPHTHOQUINONE	0.414		5.6	86.39	80.00	-8.0 50.0
S3	60 1,3-DINITROBENZENE	0.198		8.6	85.21	80.00	-6.5 50.0
S3	100 PENTACHLOROBENZENE	0.489		3.8	80.60	80.00	-0.7 50.0
S3	110 1-NAPHTHYLAMINE	0.837		6.6	73.13	80.00	8.6 50.0
S3	115 2-NAPHTHYLAMINE	0.966		5.8	82.51	80.00	-3.1 50.0
S3	120 2,3,4,6-TETRACHLOROP	0.297		4.5	83.39	80.00	-4.2 50.0
S3	145 5-NITRO-O-TOLUIDINE	0.355		7.2	85.81	80.00	-7.3 50.0
S4	25 SYM-TRINITROBENZENE	0.162		5.3	84.96	80.00	-6.2 50.0
S4	35 PHENACETIN	0.317		3.2	79.69	80.00	0.4 50.0
S4	37 DIALLATE	0.090		4.4	78.88	80.00	1.4 50.0
S4	45 4-AMINOBIIPHENYL	0.609		7.6	79.14	80.00	1.1 50.0
S4	55 PRONAMIDE	0.299		5.2	80.55	80.00	-0.7 50.0
S4	60 PENTACHLORONITROBENZ	0.080		1.8	78.96	80.00	1.3 50.0
S4	75 2SECBUTYL-4,6-DINITR	0.153		9.3	85.15	80.00	-6.4 50.0
S4	100 4-NITROQUINOLINE-1-O	0.072		11.5	84.40	80.00	-5.5 50.0
S4	105 METHAPYRILENE	0.305		26.8	53.79	80.00	32.8 50.0
S4	106 ISODRIN	0.127		1.5	80.73	80.00	-0.9 50.0
S4	120 CHLOROBENZILATE	0.323		11.1	76.29	80.00	4.6 50.0
S5	20 ARAMITE (#1)	0.061		4.4	79.45	80.00	0.7 50.0
S5	25 ARAMITE (#2)	0.087		4.4	83.30	80.00	-4.1 50.0
S5	30 P-DIMETHYLAMINOAZOBE	0.419		1.1	80.14	80.00	-0.2 50.0
S5	35 3,3'-DIMETHYLBENZIDI	0.418		17.0	77.82	80.00	2.7 50.0
S5	37 KEPONE	0.082		34.6	87.25	80.00	-9.1 50.0
S5	45 2-ACETYLAMINOFLUOREN	0.432		4.0	81.04	80.00	-1.3 50.0

S5	85	3-METHYLCHOLANTHRENE	0.548	6.1	80.10	80.00	-0.1	50.0
S6	20	7,12-DIMETHYLBENZANT	0.610	0.5	79.97	80.00	0.0	50.0

12.2% of the compounds exceed the %RSD value  
of 15.0. 6 of 49 compounds are out.

0 CCC cmpds out  
1 of nonCCC exceed 50 %D

QUANTERRA GC/MS  
SEMIVOLATILE APPENDIX IX CALIBRATION CHE

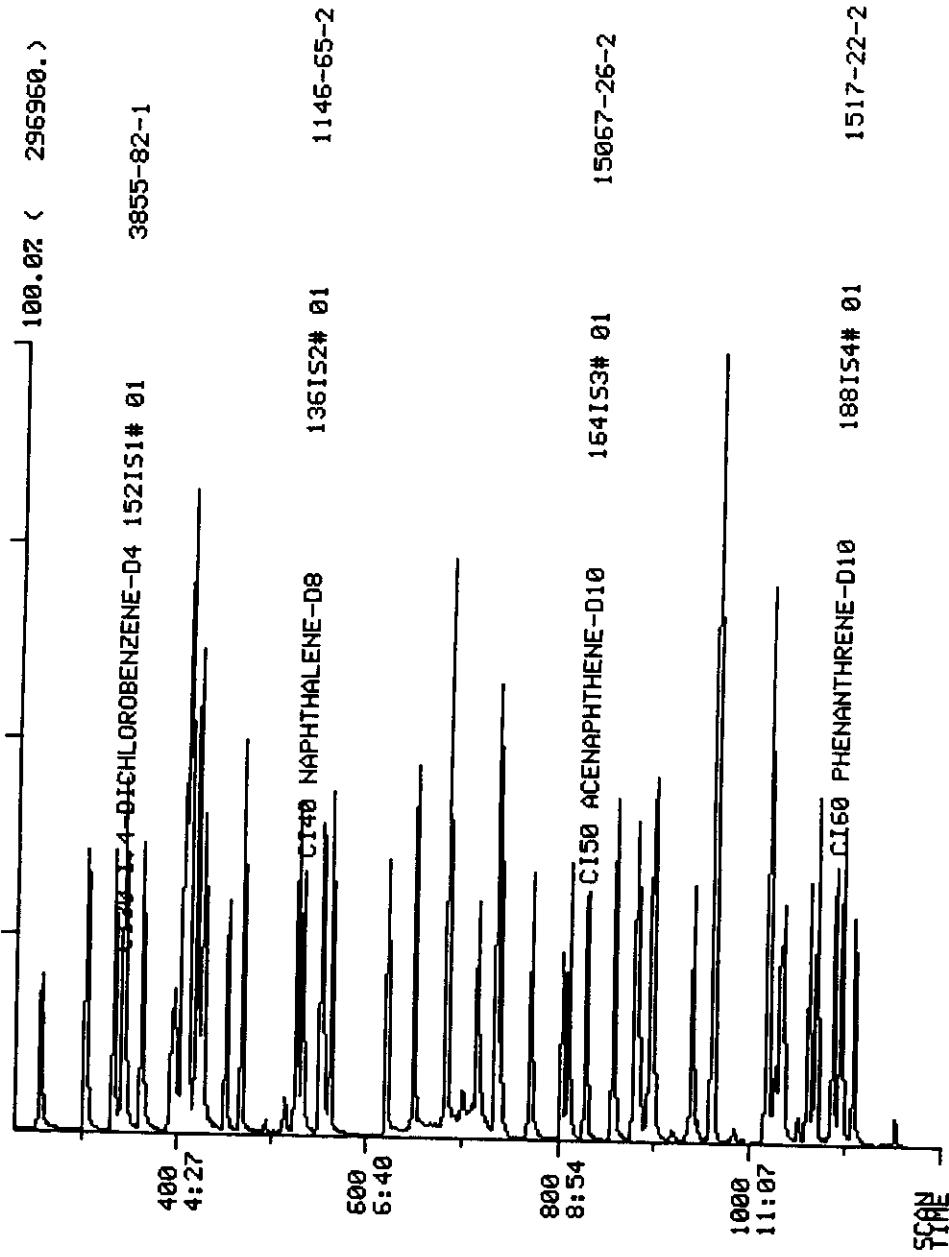
CONTIN. CALIBRATION: 80 UG/ML APPENDIX ST16980831A 08/31/98 13:07:00  
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		ST16980831A		ST16980831B		ST16980831C		ST16980831D		ST16980831A		ST16980831E	
		ST16980831F											
Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	
S1	20	80, 1.587	10, 1.770	20, 1.583	50, 1.578	80, 1.587	120, 1.449						
S1	20	160, 1.576											
S1	30	80, 1.577		20, 1.563	50, 1.594	80, 1.577	120, 1.547						
S1	30	160, 1.545											
S1	40	80, 0.876	10, 0.700	20, 0.737	50, 0.823	80, 0.876	120, 0.860						
S1	40	160, 0.870											
S1	60	80, 0.602	10, 0.619	20, 0.688	50, 0.685	80, 0.602	120, 0.619						
S1	60	160, 0.604											
S1	70	80, 0.549	10, 0.450	20, 0.506	50, 0.521	80, 0.549	120, 0.547						
S1	70	160, 0.555											
S1	85	80, 1.168	10, 0.921	20, 1.008	50, 1.120	80, 1.168	120, 1.190						
S1	85	160, 1.244											
S1	95	80, 0.605		20, 0.612	50, 0.605	80, 0.605	120, 0.595						
S1	95	160, 0.592											
S1	170	80, 1.945	10, 1.869	20, 2.024	50, 1.960	80, 1.945	120, 1.903						
S1	170	160, 1.903											
S1	175	80, 0.562	10, 0.481	20, 0.555	50, 0.575	80, 0.562	120, 0.569						
S1	175	160, 0.575											
S1	180	80, 0.805	10, 0.828	20, 0.838	50, 0.821	80, 0.805	120, 0.817						
S1	180	160, 0.816											
S1	182	80, 2.408	10, 1.848	20, 2.130	50, 2.339	80, 2.408	120, 2.391						
S1	182	160, 2.394											
S1	195	80, 2.129		20, 2.463	50, 2.228	80, 2.129	120, 2.032						
S1	195	160, 1.949											
S2	15	80, 0.327	10, 0.293	20, 0.312	50, 0.326	80, 0.327	120, 0.337						
S2	15	160, 0.329											
S2	50	80, 1.036		20, 1.575	50, 1.353	80, 1.036	120, 1.276						
S2	50	160, 1.184											
S2	85	80, 0.280	10, 0.202	20, 0.234	50, 0.264	80, 0.280	120, 0.279						
S2	85	160, 0.285											
S2	90	80, 0.197		20, 0.196	50, 0.201	80, 0.197	120, 0.198						
S2	90	160, 0.198											

S2 115 80, 0.038		20, 0.163	50, 0.167	80, 0.038	120, 0.165
S2 115 160, 0.121					
S2 120 80, 0.225	10, 0.186	20, 0.205	50, 0.213	80, 0.225	120, 0.224
S2 120 160, 0.224					
S2 140 80, 0.250		20, 0.246	50, 0.247	80, 0.250	120, 0.245
S2 140 160, 0.250					
S3 10 80, 0.430	10, 0.469	20, 0.431	50, 0.434	80, 0.430	120, 0.416
S3 10 160, 0.419					
S3 15 80, 0.043		20, 0.046	50, 0.042	80, 0.043	120, 0.044
S3 15 160, 0.044					
S3 35 80, 0.196		20, 0.201	50, 0.195	80, 0.196	120, 0.190
S3 35 160, 0.192					
S3 42 80, 1.528	10, 1.531	20, 1.098	50, 1.129	80, 1.528	120, 1.134
S3 42 160, 1.488					
S3 50 80, 0.447		20, 0.403	50, 0.429	80, 0.447	120, 0.404
S3 50 160, 0.389					
S3 60 80, 0.210	10, 0.166	20, 0.191	50, 0.202	80, 0.210	120, 0.212
S3 60 160, 0.204					
S3 100 80, 0.493	10, 0.516	20, 0.504	50, 0.483	80, 0.493	120, 0.469
S3 100 160, 0.471					
S3 110 80, 0.765	10, 0.869	20, 0.859	50, 0.889	80, 0.765	120, 0.872
S3 110 160, 0.769					
S3 115 80, 0.996	10, 0.924	20, 0.974	50, 1.041	80, 0.996	120, 0.980
S3 115 160, 0.881					
S3 120 80, 0.310		20, 0.277	50, 0.302	80, 0.310	120, 0.306
S3 120 160, 0.290					
S3 145 80, 0.381		20, 0.345	50, 0.368	80, 0.381	120, 0.367
S3 145 160, 0.316					
S4 25 80, 0.172		20, 0.149	50, 0.162	80, 0.172	120, 0.166
S4 25 160, 0.159					
S4 35 80, 0.316		20, 0.305	50, 0.324	80, 0.316	120, 0.330
S4 35 160, 0.311					
S4 37 80, 0.089		20, 0.088	50, 0.087	80, 0.089	120, 0.089
S4 37 160, 0.097					
S4 45 80, 0.602		20, 0.643	50, 0.664	80, 0.602	120, 0.590
S4 45 160, 0.546					
S4 55 80, 0.301		20, 0.312	50, 0.315	80, 0.301	120, 0.291
S4 55 160, 0.277					
S4 60 80, 0.079		20, 0.081	50, 0.079	80, 0.079	120, 0.082
S4 60 160, 0.081					
S4 75 80, 0.163		20, 0.130	50, 0.165	80, 0.163	120, 0.159

S4	75	160,	0.150										
S4	100	80,	0.076		20,	0.058	50,	0.074	80,	0.076	120,	0.079	
S4	100	160,	0.075										
S4	105	80,	0.205		20,	0.424	50,	0.339	80,	0.205	120,	0.279	
S4	105	160,	0.277										
S4	106	80,	0.128	10,	0.127	20,	0.125	50,	0.129	80,	0.128	120,	0.127
S4	106	160,	0.124										
S4	120	80,	0.308	10,	0.376	20,	0.351	50,	0.329	80,	0.308	120,	0.295
S4	120	160,	0.281										
S5	20	80,	0.061		20,	0.057	50,	0.063	80,	0.061	120,	0.064	
S5	20	160,	0.061										
S5	25	80,	0.090		20,	0.081	50,	0.090	80,	0.090	120,	0.087	
S5	25	160,	0.085										
S5	30	80,	0.420		20,	0.417	50,	0.425	80,	0.420	120,	0.412	
S5	30	160,	0.420										
S5	35	80,	0.406		20,	0.522	50,	0.450	80,	0.406	120,	0.364	
S5	35	160,	0.346										
S5	37	80,	0.089		20,	0.123	50,	0.087	80,	0.089	120,	0.056	
S5	37	160,	0.054										
S5	45	80,	0.438		20,	0.404	50,	0.450	80,	0.438	120,	0.429	
S5	45	160,	0.440										
S5	85	80,	0.549		20,	0.586	50,	0.576	80,	0.549	120,	0.515	
S5	85	160,	0.514										
S6	20	80,	0.610		20,	0.609	50,	0.615	80,	0.610	120,	0.608	
S6	20	160,	0.609										

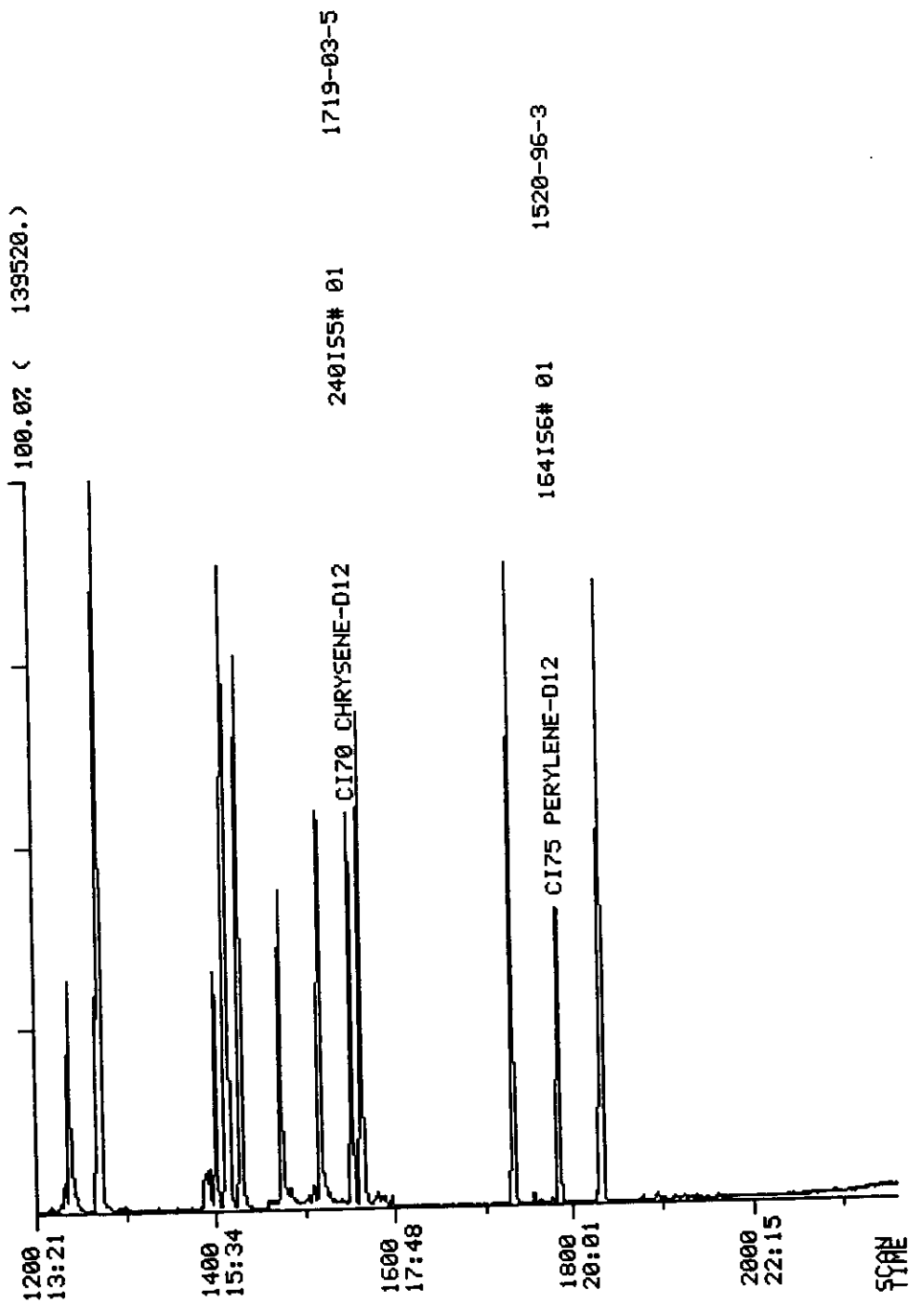
DATA FROM FILE: ST16980831A SCANS 230 TO 1200 ACQUIRED: 08/31/98 13:07:00  
 SAMPLE: 80UG/ML AP9 062598F CALI: ST16980831A #3  
 CONDS.: INST. ID: F16





DATA FROM FILE: ST16980831A SCANS 1200 TO 2158 ACQUIRED: 08/31/98 13:07:00  
CALI: ST16980831A #3

SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16



PC 11

Data: ST16980831A.T1

08/31/98 13:07:00

Sample: 80UG/ML AP9 062598F

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from average of whole .RL

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	C170 CHRYSENE-D12	240IS5# 01	1719-03-5
6	C175 PERYLENE-D12	164IS6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALATE	234 S4# 37	
40	4-AMINOBIPHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	2SEC BUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPYRILENE	58 S4#105	91-80-5
46	ISODRIN		
47	CHLOROBENZILATE	139 S4#120	

No	Name			
48	ARAMITE (#1)	185 S5# 20	140-57-8	
49	ARAMITE (#2)	185 S5# 25	140-57-8	
50	P-DIMETHYLAMINOAZOBENZENE	120 S5# 30	60-11-7	

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	29680.	40.000 UG/ML	0.97
2	136	530	5:54	2	1.000	A BB	108356.	40.000 UG/ML	0.97
3	164	828	9:13	3	1.000	A BB	60104.	40.000 UG/ML	0.97
4	188	1087	12:05	4	1.000	A BB	107988.	40.000 UG/ML	0.97
5	240	1551	17:15	5	1.000	A BV	69686.	40.000 UG/ML	0.97
6	264	1782	19:49	6	1.000	A BB	54786.	40.000 UG/ML	0.97
7	79	103	1:09	1	0.303	A BB	94198.	79.822 UG/ML	1.93
8	93	145	1:37	1	0.426	A BB	93595.	80.606 UG/ML	1.95
9	42	157	1:45	1	0.462	A BB	51975.	86.359 UG/ML	2.09
10	80	182	2:01	1	0.535	A BB	35745.	75.724 UG/ML	1.83
11	102	218	2:25	1	0.641	A BB	32568.	84.191 UG/ML	2.04
12	79	255	2:50	1	0.750	A BB	69355.	84.324 UG/ML	2.04
13	117	302	3:22	1	0.888	A BB	35884.	80.375 UG/ML	1.94
14	105	404	4:30	1	1.188	A BB	115427.	80.441 UG/ML	1.94
15	100	405	4:30	1	1.191	A BB	33375.	81.342 UG/ML	1.97
16	56	411	4:34	1	1.209	A BV	47803.	78.504 UG/ML	1.90
17	108	419	4:40	1	1.232	A BB	142918.	85.538 UG/ML	2.07
18	106	408	4:32	1	1.200	A BB	126404.	78.852 UG/ML	1.91
19	42	451	5:01	2	0.851	A BV	70838.	81.594 UG/ML	1.97
20	58	710	7:54	2	1.340	qedt	224618.	64.531 UG/ML	1.56
21	162	552	6:08	2	1.042	A BB	60603.	86.895 UG/ML	2.10
22	213	549	6:06	2	1.036	A BB	42724.	79.594 UG/ML	1.92
23	108	667	7:25	2	1.258	qedt	8144.	22.982 UG/ML	0.56
24	84	620	6:54	2	1.170	A BB	48711.	84.388 UG/ML	2.04
25	162	648	7:12	2	1.223	A BB	54270.	80.884 UG/ML	1.96
26	214	681	7:34	3	0.822	A BB	51722.	79.457 UG/ML	1.92
27	162	698	7:46	3	0.843	A BB	5166.	78.765 UG/ML	1.90
28	104	737	8:12	3	0.890	A BB	23560.	80.505 UG/ML	1.95
29	162	733	8:09	3	0.885	A BB	183709.	92.771 UG/ML	2.24
30	158	770	8:34	3	0.930	A BB	53790.	86.357 UG/ML	2.09
31	168	803	8:56	3	0.970	A BB	25287.	85.215 UG/ML	2.06
32	250	858	9:33	3	1.036	A BB	59261.	80.576 UG/ML	1.95
33	143	883	9:49	3	1.066	A BB	91990.	73.120 UG/ML	1.77
34	143	899	10:00	3	1.086	A BB	119758.	82.503 UG/ML	1.99
35	232	895	9:57	3	1.081	A BB	37212.	83.399 UG/ML	2.02
36	152	940	10:27	3	1.135	A BB	45826.	85.785 UG/ML	2.07
37	75	1025	11:24	4	0.943	A VB	37064.	84.994 UG/ML	2.05
38	108	1034	11:30	4	0.951	A BB	68243.	79.750 UG/ML	1.93
39	234	1018	11:19	4	0.937	A BV	19166.	78.905 UG/ML	1.91
40	169	1069	11:53	4	0.983	A BB	130115.	79.141 UG/ML	1.91
41	173	1096	12:11	4	1.008	A BB	65067.	80.495 UG/ML	1.95
42	237	1061	11:48	4	0.976	A BB	17138.	78.989 UG/ML	1.91
43	211	1108	12:19	4	1.019	A VB	35264.	85.064 UG/ML	2.06
44	190	1236	13:45	4	1.137	A BB	16496.	84.279 UG/ML	2.04
45	58	1266	14:05	4	1.165	A BB	44262.	53.806 UG/ML	1.30
46	193	1270	14:08	4	1.168	A BV	27606.	80.734 UG/ML	1.95
47	139	1427	15:52	4	1.313	A BB	66594.	76.253 UG/ML	1.84
48	185	1399	15:34	5	0.902	A BB	8471.	79.616 UG/ML	1.92
49	185	1414	15:44	5	0.912	A BB	12567.	83.361 UG/ML	2.02
50	120	1412	15:42	5	0.910	A VB	58473.	80.139 UG/ML	1.94

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:15	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:49	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.303	1.00	79.82	50.00	2.539	1.590	1.60
8	1:37	1.00	0.426	1.00	80.61	50.00	2.523	1.565	1.61
9	1:45	1.00	0.462	1.00	86.36	50.00	1.401	0.811	1.73
10	2:01	1.00	0.535	1.00	75.72	50.00	0.963	0.636	1.51
11	2:25	1.00	0.641	1.00	84.19	50.00	0.878	0.521	1.68
12	2:50	1.00	0.750	1.00	84.32	50.00	1.869	1.108	1.69
13	3:22	1.00	0.888	1.00	80.38	50.00	0.967	0.602	1.61
14	4:30	1.00	1.188	1.00	80.44	50.00	3.111	1.934	1.61
15	4:30	1.00	1.191	1.00	81.34	50.00	0.900	0.553	1.63
16	4:34	1.00	1.209	1.00	78.50	50.00	1.288	0.821	1.57
17	4:40	1.00	1.232	1.00	85.54	50.00	3.852	2.252	1.71
18	4:32	1.00	1.200	1.00	78.85	50.00	3.407	2.160	1.58
19	5:01	1.00	0.851	1.00	81.59	50.00	0.523	0.320	1.63
20	7:54	1.00	1.340	1.00	64.53	50.00	1.658	1.285	1.29
21	6:08	1.00	1.042	1.00	86.89	50.00	0.447	0.257	1.74
22	6:06	1.00	1.036	1.00	79.59	50.00	0.315	0.198	1.59
23	7:25	1.00	1.258	1.00	22.98	50.00	0.060	0.131	0.46
24	6:54	1.00	1.170	1.00	84.39	50.00	0.360	0.213	1.69
25	7:12	1.00	1.223	1.00	80.88	50.00	0.401	0.248	1.62
26	7:34	1.00	0.822	1.00	79.46	50.00	0.688	0.433	1.59
27	7:46	1.00	0.843	1.00	78.76	50.00	0.069	0.044	1.58
28	8:12	1.00	0.890	1.00	80.51	50.00	0.314	0.195	1.61
29	8:09	1.00	0.885	1.00	92.77	50.00	2.445	1.318	1.86
30	8:34	1.00	0.930	1.00	86.36	50.00	0.716	0.415	1.73
31	8:56	1.00	0.970	1.00	85.22	50.00	0.337	0.197	1.70
32	9:33	1.00	1.036	1.00	80.58	50.00	0.789	0.489	1.61
33	9:49	1.00	1.066	1.00	73.12	50.00	1.224	0.837	1.46
34	10:00	1.00	1.086	1.00	82.50	50.00	1.594	0.966	1.65
35	9:57	1.00	1.081	1.00	83.40	50.00	0.495	0.297	1.67
36	10:27	1.00	1.135	1.00	85.79	50.00	0.610	0.356	1.72
37	11:24	1.00	0.943	1.00	84.99	50.00	0.275	0.162	1.70
38	11:30	1.00	0.951	1.00	79.75	50.00	0.506	0.317	1.59
39	11:19	1.00	0.937	1.00	78.90	50.00	0.142	0.090	1.58
40	11:53	1.00	0.983	1.00	79.14	50.00	0.964	0.609	1.58
41	12:11	1.00	1.008	1.00	80.50	50.00	0.482	0.299	1.61
42	11:48	1.00	0.976	1.00	78.99	50.00	0.127	0.080	1.58
43	12:19	1.00	1.019	1.00	85.06	50.00	0.261	0.154	1.70
44	13:45	1.00	1.137	1.00	84.28	50.00	0.122	0.073	1.69
45	14:05	1.00	1.165	1.00	53.81	50.00	0.328	0.305	1.08
46	14:08	1.00	1.168	1.00	80.73	50.00	0.205	0.127	1.61
47	15:52	1.00	1.313	1.00	76.25	50.00	0.493	0.323	1.53
48	15:34	1.00	0.902	1.00	79.62	50.00	0.097	0.061	1.59
49	15:44	1.00	0.912	1.00	83.36	50.00	0.144	0.087	1.67
50	15:42	1.00	0.910	1.00	80.14	50.00	0.671	0.419	1.60

1018

Data: ST16980831A.TI  
08/31/98 13:07:00  
Sample: 80UG/ML AP9 062598F  
Conds.: INST. ID: F16  
Formula: IUL INJ.  
Submitted by: QES

Instrument: F16 Weight: 0.000  
Analyst: DAT Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
Resp. fac. from average of whole .RL

No	Name	212 S5# 35	272 S5# 37	181 S5# 45	268 S5# 85	256 S6# 20	196 S6# 30
51	3,3'-DIMETHYLBENZIDINE						
52	KEPONE						
53	2-ACETYLAMINOFLUORENE			53-96-3			
54	3-METHYLCHOLANTHRENE			56-49-5			
55	7,12-DIMETHYLBENZANTHRACENE			75-97-6			
56	HEXACHLOROPHENE			70-30-4			

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1472	16:22	5	0.949	A BB	56612.	77.811 UG/ML	1.88
52	272	1458	16:13	5	0.940	qedt (2)	12434.	87.077 UG/ML	2.10
53	181	1516	16:52	5	0.977	A BB	61022.	81.053 UG/ML	1.96
54	268	1832	20:23	5	1.181	A BB	76474.	80.095 UG/ML	1.94
55	256	1731	19:15	6	0.971	A BB	66838.	79.991 UG/ML	1.93
56	NOT FOUND								

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:22	1.00	0.949	1.00	77.81	50.00	0.650	0.418	1.56
52	16:13	1.00	0.940	1.00	87.08	50.00	0.143	0.082	1.74
53	16:52	1.00	0.977	1.00	81.05	50.00	0.701	0.432	1.62
54	20:23	1.00	1.181	1.00	80.10	50.00	0.878	0.548	1.60
55	19:15	1.00	0.971	1.00	79.99	50.00	0.976	0.610	1.60
56	21:11		0.992						

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPHY**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND CODED**

RIC-MASS CHROMATOGRAM

08/31/98 13:07:00

SAMPLE: 80UG/ML AP9 062598F

CONDS.: INST. ID: F16

RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 5, 2.0 J 0

DATA: ST16980831A #1

CALI: ST16980831A #3

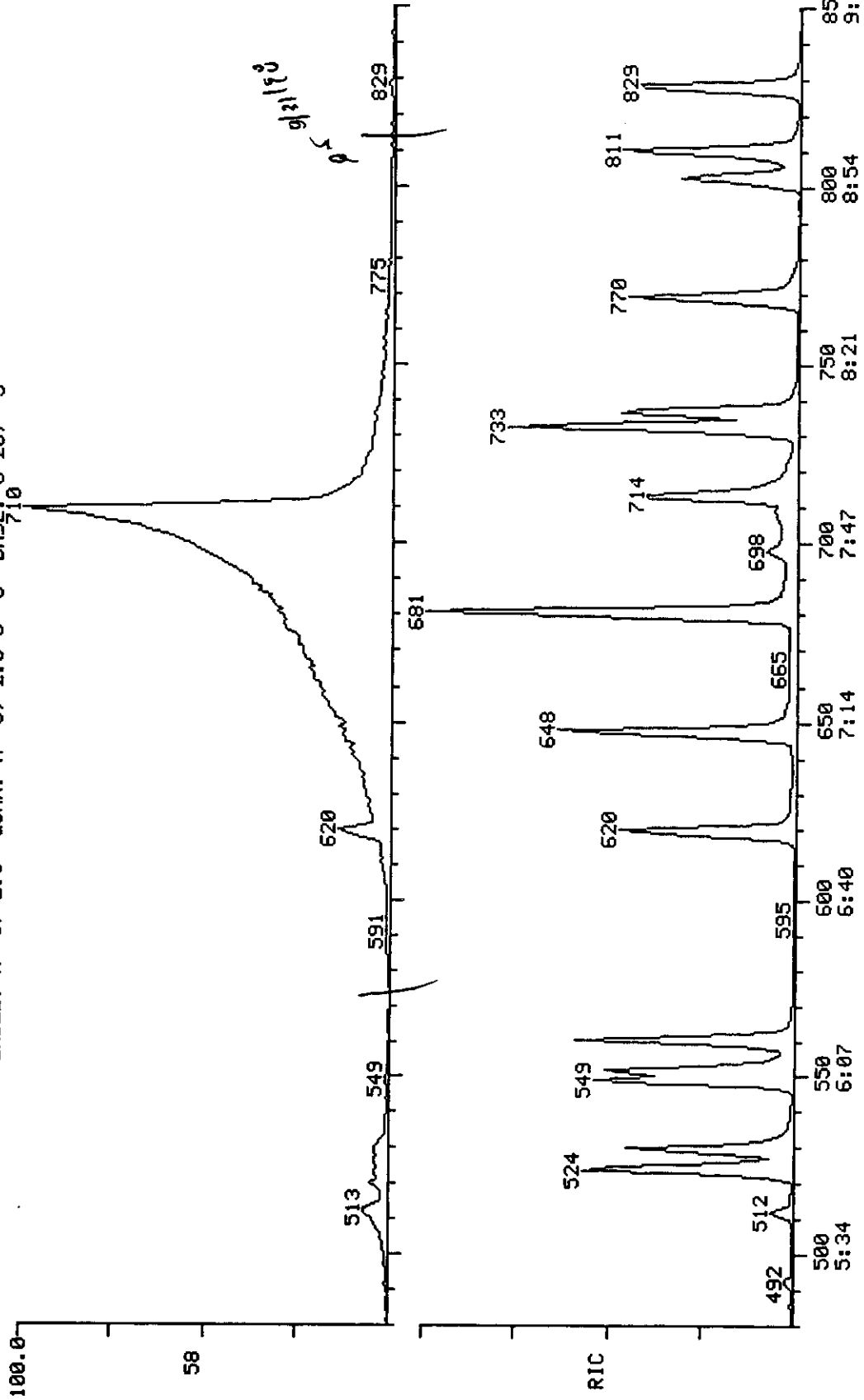
SCANS 480 TO 850

BASE: U 20, 3

100.0 58 7528.

513 549 591 620 681 710 775 829 58.017 ± 0.500 218112.

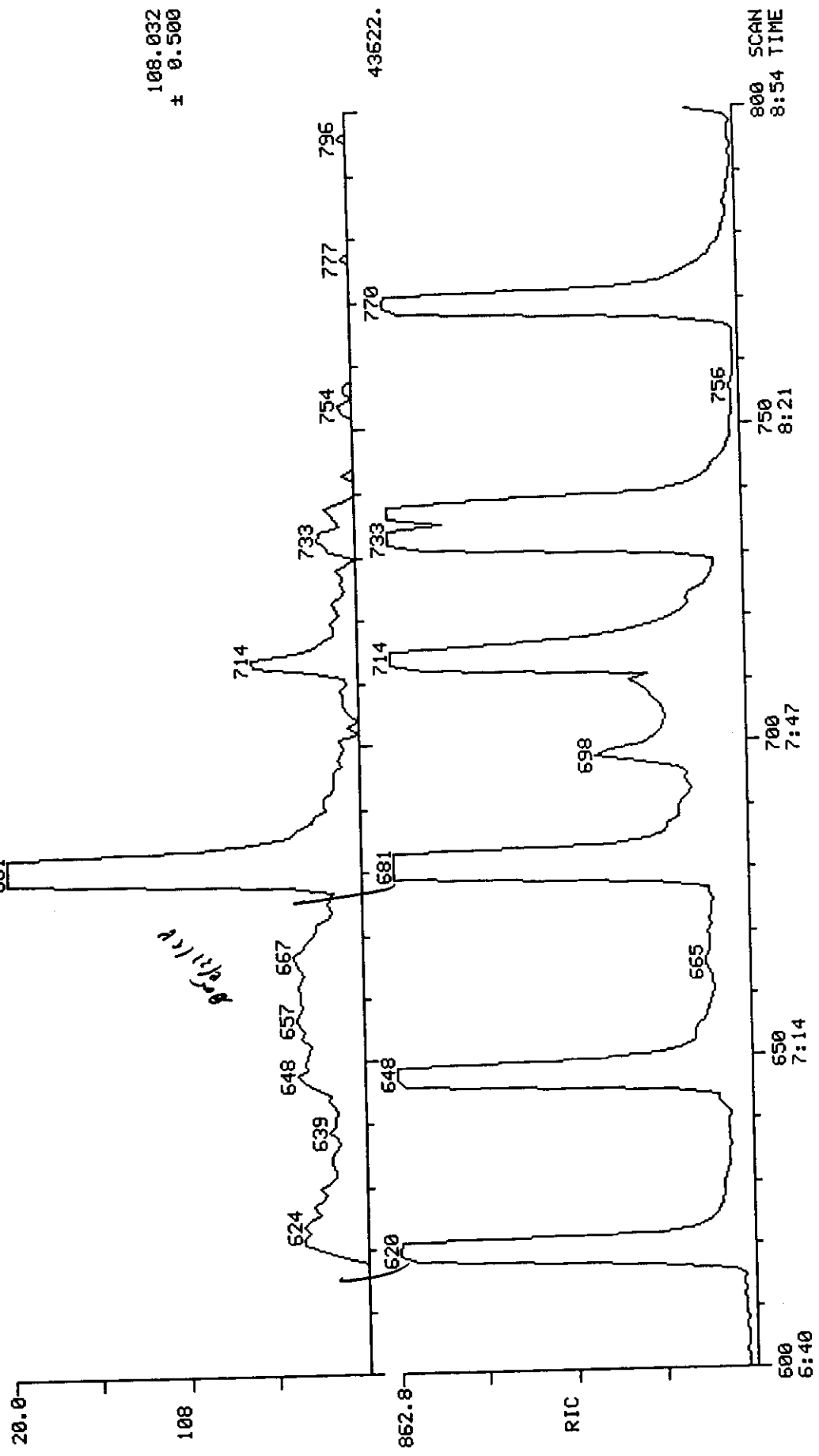
6/21/98



1020

RIC+MASS CHROMATOGRAM  
08/31/98 13:07:00  
SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16  
RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3  
DATA: ST16980831A #1  
CALI: ST16980831A #3  
SCANS 600 TO 800

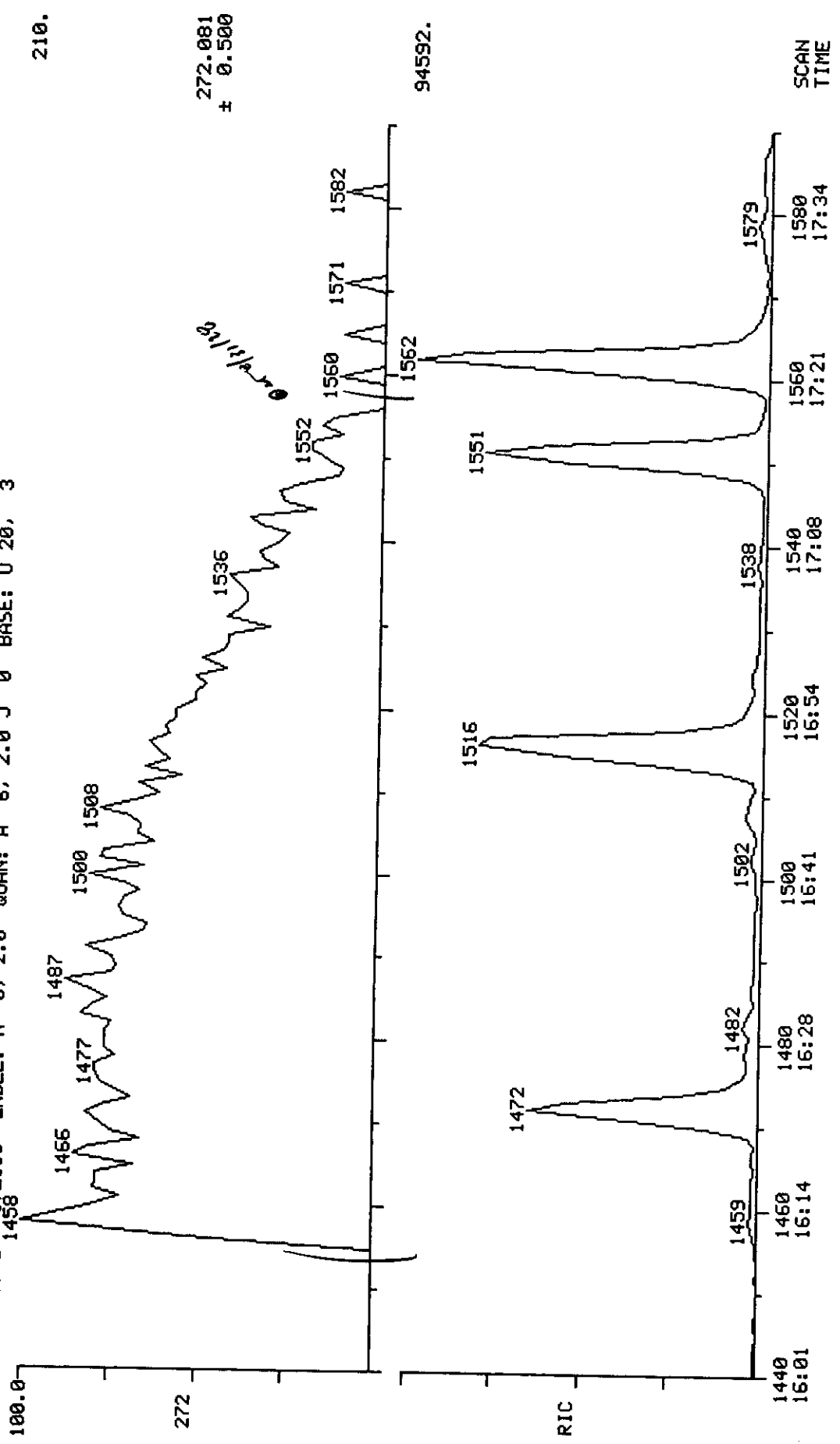
1011.



108.032  
± 0.500

43522.

RIC+MASS CHROMATOGRAM  
 08/31/98 13:07:00 DATA: ST16980831A #1 SCANS 1440 TO 1590  
 CALLI: ST16980831A #3  
 SAMPLE: 80UG/ML AP9 062598F  
 CONDS.: INST. ID: F16  
 RANGE: G 1, 2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3



272.081  
± 0.500

94592.



QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980902 09/02/98 14:53:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

Library Entry	Compound	Initial Calibration			Continuing Calibration		
		Response Factor	% RSD		Amount		% Diff
		Avg	Min	Max	Calc	Exp	Max
S1 15	C310 N-NITROSODIMETH	1.019		4.4	77.17	80.00	3.5 50.0
S1 100	C320 ANILINE	1.846		6.2	72.66	80.00	9.2 50.0
S1 105	C315 PHENOL	1.968		7.8 30.0	78.81	80.00	1.5 20.0
S1 110	C325 BIS(2-CHLOROETH	1.368		6.1	84.70	80.00	-5.9 50.0
S1 115	C330 2-CHLOROPHENOL	1.497		5.5	82.69	80.00	-3.4 50.0
S1 125	C335 1,3-DICHLOROBEN	1.586		4.5	81.00	80.00	-1.3 50.0
S1 130	C340 1,4-DICHLOROBEN	1.618		4.8 30.0	80.31	80.00	-0.4 20.0
S1 145	C345 BENZYL ALCOHOL	0.854		9.3	71.16	80.00	11.1 50.0
S1 150	C350 1,2-DICHLOROBEN	1.500		5.0	80.48	80.00	-0.6 50.0
S1 160	C355 2-METHYLPHENOL	1.174		7.4	74.75	80.00	6.6 50.0
S1 165	C360 2,2'-OXYBIS(1-C	2.539		3.9	77.25	80.00	3.4 50.0
S1 185	C365 4-METHYLPHENOL	1.276		9.0	71.19	80.00	11.0 50.0
S1 190	C370 N-NITROSO-DI-N-	0.906	0.050	5.1	75.97	80.00	5.0 50.0
S1 200	C375 HEXACHLOROETHAN	0.708		3.5	79.54	80.00	0.6 50.0
S2 10	C410 NITROBENZENE	0.445		7.2	84.18	80.00	-5.2 50.0
S2 20	C415 ISOPHORONE	0.742		5.5	80.59	80.00	-0.7 50.0
S2 25	C420 2-NITROPHENOL	0.213		7.9 30.0	82.30	80.00	-2.9 20.0
S2 30	C425 2,4-DIMETHYLPHE	0.330		16.9	91.51	80.00	-14.4 50.0
S2 45	C430 BENZOIC ACID	0.276		4.7	44.76	80.00	44.1 50.0
S2 35	C435 BIS(2-CHLOROETH	0.511		4.1	80.57	80.00	-0.7 50.0
S2 40	C440 2,4-DICHLOROPHE	0.308		6.6 30.0	80.28	80.00	-0.3 20.0
S2 55	C445 1,2,4-TRICHLORO	0.331		5.0	85.14	80.00	-6.4 50.0
S2 60	C450 NAPHTHALENE	1.022		6.6	85.54	80.00	-6.9 50.0
S2 80	C455 4-CHLOROANILINE	0.454		4.5	82.80	80.00	-3.5 50.0
S2 95	C460 HEXACHLOROBUTAD	0.168		5.1 30.0	86.60	80.00	-8.2 20.0
S2 130	C465 4-CHLORO-3-METH	0.338		9.0 30.0	80.18	80.00	-0.2 20.0
S2 145	C470 2-METHYLNAPHTHA	0.751		10.8	103.64	80.00	-29.5 50.0
S3 20	C510 HEXACHLOROCYCLO	0.372	0.050	10.1	83.99	80.00	-5.0 50.0
S3 25	C515 2,4,6-TRICHLORO	0.404		10.6 30.0	79.06	80.00	1.2 20.0
S3 30	C520 2,4,5-TRICHLORO	0.412		7.1	80.17	80.00	-0.2 50.0
S3 40	C525 2-CHLORONAPHTHA	1.193		6.6	85.88	80.00	-7.4 50.0
S3 45	C530 2-NITROANILINE	0.502		2.9	79.87	80.00	0.2 50.0
S3 55	C535 DIMETHYLPHTHALA	1.341		6.1	84.51	80.00	-5.6 50.0
S3 65	C540 ACENAPHTHYLENE	1.839		8.2	86.94	80.00	-8.7 50.0
S3 70	C543 2,6-DINITROTOLU	0.324		8.5	87.18	80.00	-9.0 50.0
S3 75	C545 3-NITROANILINE	0.383		1.4	81.46	80.00	-1.8 50.0
S3 80	C550 ACENAPHTHENE	1.158		6.9 30.0	86.26	80.00	-7.8 20.0
S3 85	C555 2,4-DINITROPHEN	0.189	0.050	10.3	68.85	80.00	13.9 50.0

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980902 09/02/98 14:53:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
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 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

Library Entry Compound	Initial Calibration			Continuing Calibration		
	Response Factor	% RSD		Amount		% Diff
	Avg	Min	Max	Calc	Exp	Max
S3 90 C565 DIBENZOFURAN	1.669		6.2	85.82	80.00	-7.3 50.0
S3 105 C570 2,4-DINITROTOLU	0.397		8.8	93.46	80.00	-16.8 50.0
S3 130 C580 DIETHYLPHTHALAT	1.369		7.7	86.28	80.00	-7.9 50.0
S3 140 C585 4-CHLOROPHENYL-	0.613		5.4	88.61	80.00	-10.8 50.0
S3 135 C590 FLUORENE	1.298		6.8	88.71	80.00	-10.9 50.0
S3 150 C595 4-NITROANALINE	0.307		8.4	78.22	80.00	2.2 50.0
S4 10 C610 4,6-DINITRO-2-M	0.132		7.6	82.12	80.00	-2.7 50.0
S4 15 C615 N-NITROSODIPHEN	0.551		4.2 30.0	79.08	80.00	1.2 20.0
S4 20 C620 AZOBENZENE	1.171		3.9	80.34	80.00	-0.4 50.0
S4 30 C625 4-BROMOPHENYL-P	0.223		3.9	87.69	80.00	-9.6 50.0
S4 40 C630 HEXACHLOROBENZE	0.260		6.7	87.45	80.00	-9.3 50.0
S4 50 C635 PENTACHLOROPHEN	0.155		5.9 30.0	75.84	80.00	5.2 20.0
S4 65 C640 PHENANTHRENE	1.033		5.2	84.48	80.00	-5.6 50.0
S4 70 C645 ANTHRACENE	0.987		11.0	85.24	80.00	-6.6 50.0
S4 80 C647 CARBAZOLE	0.764		10.3	84.72	80.00	-5.9 50.0
S4 85 C650 DI-N-BUTYLPHTHA	1.247		9.4	86.27	80.00	-7.8 50.0
S4 110 C655 FLUORANTHENE	0.900		13.5 30.0	84.07	80.00	-5.1 20.0
S5 10 C710 BENZIDINE	0.129		23.7	12.00	80.00	85.0* 50.0
S5 15 C715 PYRENE	1.323		6.5	82.90	80.00	-3.6 50.0
S5 40 C720 BUTYLBENZYLPHTH	0.685		7.7	82.18	80.00	-2.7 50.0
S5 55 C725 3,3'-DICHLOROBE	0.408		5.7	76.65	80.00	4.2 50.0
S5 50 C730 BENZO(A)ANTHRAC	1.125		5.4	87.03	80.00	-8.8 50.0
S5 60 C740 CHRYSENE	1.014		6.7	82.83	80.00	-3.5 50.0
S5 65 C745 BIS(2-ETHYLHEXY	0.980		9.4	80.96	80.00	-1.2 50.0
S6 10 C760 DI-N-OCTYL PHTH	1.925		7.4 30.0	80.21	80.00	-0.3 20.0
S6 15 C765 BENZO(B)FLUORAN	1.444		5.0	77.01	80.00	3.7 50.0
S6 25 C770 BENZO(K)FLUORAN	1.159		9.8	86.88	80.00	-8.6 50.0
S6 35 C775 BENZO(A)PYRENE	1.182		5.3 30.0	82.06	80.00	-2.6 20.0
S6 55 C780 INDENO(1,2,3-CD	1.279		6.6	80.36	80.00	-0.4 50.0
S6 60 C785 DIBENZ(A,H)ANTH	1.267		8.1	81.54	80.00	-1.9 50.0
S6 65 C790 BENZO(G,H,I)PER	1.331		8.2	81.13	80.00	-1.4 50.0
S1 6 CS75 1,2-DICHLOROBNZ	0.920		6.5	79.99	80.00	0.0 50.0
S2 2 CS20 NITROBENZENE-D5	0.470		5.2	82.78	80.00	-3.5 50.0
S3 3 CS25 2-FLUOROBIPHENY	1.234		6.2	84.45	80.00	-5.6 50.0
S5 2 CS30 TERPHENYL-D14	0.995		5.3	84.17	80.00	-5.2 50.0
S1 3 CS45 PHENOL-D5	2.026		2.0	75.06	80.00	6.2 50.0
S1 2 CS50 2-FLUOROPHENOL	1.368		4.4	71.18	80.00	11.0 50.0
S1 5 CS70 2-CHLOROPHENOL-	1.482		4.1	76.79	80.00	4.0 50.0
S3 2 CS55 2,4,6-TRIBROMOP	0.236		6.9	88.86	80.00	-11.1 50.0

2.6% of the compounds exceed the %RSD value  
of 15.0. 2 of 78 compounds are out.

0 CCC cmpds out  
1 of nonCCC exceed 50 %D

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980902 09/02/98 14:53:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
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 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

ST16980902 ST16980827A ST16980827B ST16980827C ST16980827 ST16980827D  
 ST16980827E

Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S1 15	80,	0.983	10,	0.952	20,	1.010	50,	1.016	80,	1.087
S1 15	160,	1.042							120,	1.008
S1 100	80,	1.677	10,	1.947	20,	1.988	50,	1.814	80,	1.876
S1 100	160,	1.674							120,	1.778
S1 105	80,	1.939	10,	1.699	20,	2.037	50,	2.035	80,	2.148
S1 105	160,	1.982							120,	1.907
S1 110	80,	1.448	10,	1.336	20,	1.465	50,	1.438	80,	1.414
S1 110	160,	1.261							120,	1.291
S1 115	80,	1.547	10,	1.386	20,	1.569	50,	1.522	80,	1.603
S1 115	160,	1.451							120,	1.450
S1 125	80,	1.606	10,	1.515	20,	1.668	50,	1.618	80,	1.659
S1 125	160,	1.510							120,	1.546
S1 130	80,	1.624	10,	1.580	20,	1.695	50,	1.666	80,	1.697
S1 130	160,	1.518							120,	1.549
S1 145	80,	0.760	10,	0.702	20,	0.851	50,	0.872	80,	0.913
S1 145	160,	0.922							120,	0.866
S1 150	80,	1.509	10,	1.475	20,	1.606	50,	1.482	80,	1.573
S1 150	160,	1.409							120,	1.454
S1 160	80,	1.097	10,	0.998	20,	1.206	50,	1.221	80,	1.209
S1 160	160,	1.213							120,	1.199
S1 165	80,	2.452	10,	2.393	20,	2.676	50,	2.597	80,	2.582
S1 165	160,	2.486							120,	2.502
S1 185	80,	1.135	10,	1.049	20,	1.298	50,	1.355	80,	1.287
S1 185	160,	1.351							120,	1.316
S1 190	80,	0.860	10,	0.819	20,	0.934	50,	0.938	80,	0.940
S1 190	160,	0.890							120,	0.913
S1 200	80,	0.704	10,	0.665	20,	0.740	50,	0.713	80,	0.720
S1 200	160,	0.706							120,	0.706
S2 10	80,	0.468	10,	0.423	20,	0.477	50,	0.455	80,	0.486
S2 10	160,	0.410							120,	0.419
S2 20	80,	0.747	10,	0.685	20,	0.778	50,	0.774	80,	0.778
S2 20	160,	0.708							120,	0.726

1625

S2 25	80, 0.219	10, 0.182	20, 0.212	50, 0.218	80, 0.232	120, 0.219
S2 25	160, 0.217					
S2 30	80, 0.377	10, 0.234	20, 0.309	50, 0.334	80, 0.400	120, 0.349
S2 30	160, 0.353					
S2 45	80, 0.154		20, 0.257	50, 0.279	80, 0.270	120, 0.280
S2 45	160, 0.292					
S2 35	80, 0.514	10, 0.481	20, 0.529	50, 0.517	80, 0.538	120, 0.502
S2 35	160, 0.498					
S2 40	80, 0.309	10, 0.270	20, 0.313	50, 0.314	80, 0.332	120, 0.311
S2 40	160, 0.306					
S2 55	80, 0.352	10, 0.319	20, 0.350	50, 0.334	80, 0.350	120, 0.322
S2 55	160, 0.310					
S2 60	80, 1.093	10, 1.008	20, 1.090	50, 1.038	80, 1.102	120, 0.960
S2 60	160, 0.935					
S2 80	80, 0.470	10, 0.419	20, 0.466	50, 0.454	80, 0.480	120, 0.449
S2 80	160, 0.455					
S2 95	80, 0.182	10, 0.160	20, 0.178	50, 0.170	80, 0.177	120, 0.167
S2 95	160, 0.157					
S2 130	80, 0.339	10, 0.279	20, 0.337	50, 0.353	80, 0.365	120, 0.347
S2 130	160, 0.346					
S2 145	80, 0.973	10, 0.773	20, 0.840	50, 0.827	80, 0.757	120, 0.671
S2 145	160, 0.639					
S3 20	80, 0.390	10, 0.329	20, 0.387	50, 0.376	80, 0.428	120, 0.379
S3 20	160, 0.330					
S3 25	80, 0.399	10, 0.327	20, 0.389	50, 0.405	80, 0.435	120, 0.417
S3 25	160, 0.448					
S3 30	80, 0.413	10, 0.397	20, 0.437	50, 0.429	80, 0.443	120, 0.401
S3 30	160, 0.366					
S3 40	80, 1.280	10, 1.172	20, 1.275	50, 1.208	80, 1.283	120, 1.128
S3 40	160, 1.089					
S3 45	80, 0.501		20, 0.491	50, 0.510	80, 0.523	120, 0.487
S3 45	160, 0.497					
S3 55	80, 1.417	10, 1.257	20, 1.423	50, 1.392	80, 1.426	120, 1.297
S3 55	160, 1.251					
S3 65	80, 1.998	10, 1.751	20, 1.959	50, 1.896	80, 2.047	120, 1.707
S3 65	160, 1.671					
S3 70	80, 0.353	10, 0.274	20, 0.339	50, 0.341	80, 0.351	120, 0.321
S3 70	160, 0.317					
S3 75	80, 0.390		20, 0.382	50, 0.392	80, 0.382	120, 0.379
S3 75	160, 0.379					
S3 80	80, 1.249	10, 1.140	20, 1.227	50, 1.182	80, 1.261	120, 1.075

S3 80 160, 1.066

S3 85 80, 0.163

S3 85 160, 0.196

S3 95 80, 0.131

S3 95 160, 0.169

20, 0.156 50, 0.190 80, 0.202 120, 0.203

20, 0.164 50, 0.178 80, 0.184 120, 0.168

QUANTERRA GC/MS  
SEMIVOLATILE HSL CALIBRATION CHECK

CONTIN. CALIBRATION: 80 UG/ML HSL STD ST16980902 09/02/98 14:53:00  
 INITIAL CALIBRATION: 10 UG/ML HSL STD ST16980827A 08/27/98 9:56:00  
 INITIAL CALIBRATION: 20 UG/ML HSL STD ST16980827B 08/27/98 10:21:00  
 INITIAL CALIBRATION: 50 UG/ML HSL STD ST16980827C 08/27/98 10:51:00  
 INITIAL CALIBRATION: 80 UG/ML HSL STD ST16980827 08/27/98 9:22:00  
 INITIAL CALIBRATION: 120 UG/ML HSL STD ST16980827D 08/27/98 11:20:00  
 INITIAL CALIBRATION: 160 UG/ML HSL STD ST16980827E 08/27/98 11:50:00

ST16980902 ST16980827A ST16980827B ST16980827C ST16980827 ST16980827D  
 ST16980827E

Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S3 90	80,	1.790	10,	1.617	20,	1.782	50,	1.736	80,	1.762
S3 90	160,	1.561							120,	1.556
S3 105	80,	0.464	10,	0.339	20,	0.412	50,	0.420	80,	0.437
S3 105	160,	0.378							120,	0.396
S3 130	80,	1.476	10,	1.300	20,	1.462	50,	1.452	80,	1.462
S3 130	160,	1.209							120,	1.328
S3 140	80,	0.678	10,	0.597	20,	0.647	50,	0.631	80,	0.646
S3 140	160,	0.575							120,	0.579
S3 135	80,	1.439	10,	1.251	20,	1.398	50,	1.340	80,	1.386
S3 135	160,	1.188							120,	1.224
S3 150	80,	0.300			20,	0.326	50,	0.320	80,	0.306
S3 150	160,	0.321							120,	0.263
S4 10	80,	0.135			20,	0.121	50,	0.139	80,	0.145
S4 10	160,	0.130							120,	0.124
S4 15	80,	0.544	10,	0.568	20,	0.582	50,	0.562	80,	0.539
S4 15	160,	0.523							120,	0.530
S4 20	80,	1.176	10,	1.150	20,	1.222	50,	1.186	80,	1.212
S4 20	160,	1.158							120,	1.098
S4 30	80,	0.244	10,	0.224	20,	0.231	50,	0.223	80,	0.232
S4 30	160,	0.210							120,	0.215
S4 40	80,	0.284	10,	0.273	20,	0.281	50,	0.255	80,	0.268
S4 40	160,	0.242							120,	0.238
S4 50	80,	0.147			20,	0.148	50,	0.156	80,	0.170
S4 50	160,	0.147							120,	0.155
S4 65	80,	1.090	10,	1.017	20,	1.081	50,	1.044	80,	1.096
S4 65	160,	1.010							120,	0.948
S4 70	80,	1.051	10,	1.019	20,	1.100	50,	1.026	80,	1.059
S4 70	160,	0.809							120,	0.908
S4 80	80,	0.809	10,	0.914	20,	0.763	50,	0.729	80,	0.769
S4 80	160,	0.701							120,	0.707
S4 85	80,	1.345	10,	1.233	20,	1.366	50,	1.332	80,	1.333
S4 85	160,	1.080							120,	1.137

1622

S4	110	80, 0.946	10, 1.013	20, 1.008	50, 0.925	80, 0.950	120, 0.788
S4	110	160, 0.715					
S5	10	80, 0.019		20, 0.155	50, 0.143	80, 0.077	120, 0.141
S5	10	160, 0.131					
S5	15	80, 1.370	10, 1.209	20, 1.310	50, 1.295	80, 1.468	120, 1.295
S5	15	160, 1.358					
S5	40	80, 0.703	10, 0.587	20, 0.673	50, 0.692	80, 0.722	120, 0.700
S5	40	160, 0.734					
S5	55	80, 0.391		20, 0.441	50, 0.421	80, 0.383	120, 0.405
S5	55	160, 0.391					
S5	50	80, 1.224	10, 1.058	20, 1.195	50, 1.116	80, 1.203	120, 1.086
S5	50	160, 1.092					
S5	60	80, 1.050	10, 1.034	20, 1.068	50, 1.005	80, 1.102	120, 0.959
S5	60	160, 0.919					
S5	65	80, 0.992	10, 0.812	20, 0.978	50, 0.986	80, 1.027	120, 0.987
S5	65	160, 1.089					
S6	10	80, 1.930	10, 1.637	20, 1.959	50, 1.985	80, 1.977	120, 1.967
S6	10	160, 2.023					
S6	15	80, 1.390	10, 1.322	20, 1.536	50, 1.448	80, 1.453	120, 1.490
S6	15	160, 1.412					
S6	25	80, 1.259	10, 1.185	20, 1.223	50, 1.171	80, 1.264	120, 1.171
S6	25	160, 0.940					
S6	35	80, 1.212	10, 1.168	20, 1.257	50, 1.206	80, 1.229	120, 1.147
S6	35	160, 1.084					
S6	55	80, 1.285	10, 1.253	20, 1.355	50, 1.312	80, 1.368	120, 1.241
S6	55	160, 1.144					
S6	60	80, 1.292	10, 1.254	20, 1.386	50, 1.288	80, 1.360	120, 1.212
S6	60	160, 1.105					
S6	65	80, 1.350	10, 1.339	20, 1.436	50, 1.356	80, 1.426	120, 1.289
S6	65	160, 1.139					
S1	6	80, 0.920	10, 0.832	20, 0.999	50, 0.946	80, 0.959	120, 0.902
S1	6	160, 0.882					
S2	2	80, 0.486	10, 0.446	20, 0.494	50, 0.469	80, 0.504	120, 0.452
S2	2	160, 0.452					
S3	3	80, 1.303	10, 1.252	20, 1.315	50, 1.253	80, 1.299	120, 1.171
S3	3	160, 1.116					
S5	2	80, 1.047	10, 0.932	20, 1.002	50, 0.976	80, 1.082	120, 0.960
S5	2	160, 1.019					
S1	3	80, 1.900		20, 2.057	50, 2.059	80, 2.046	120, 1.995
S1	3	160, 1.971					
S1	2	80, 1.217		20, 1.291	50, 1.332	80, 1.448	120, 1.376

S1 2 160, 1.394

S1 5 80, 1.422

S1 5 160, 1.407

20, 1.561 50, 1.501 80, 1.503 120, 1.437

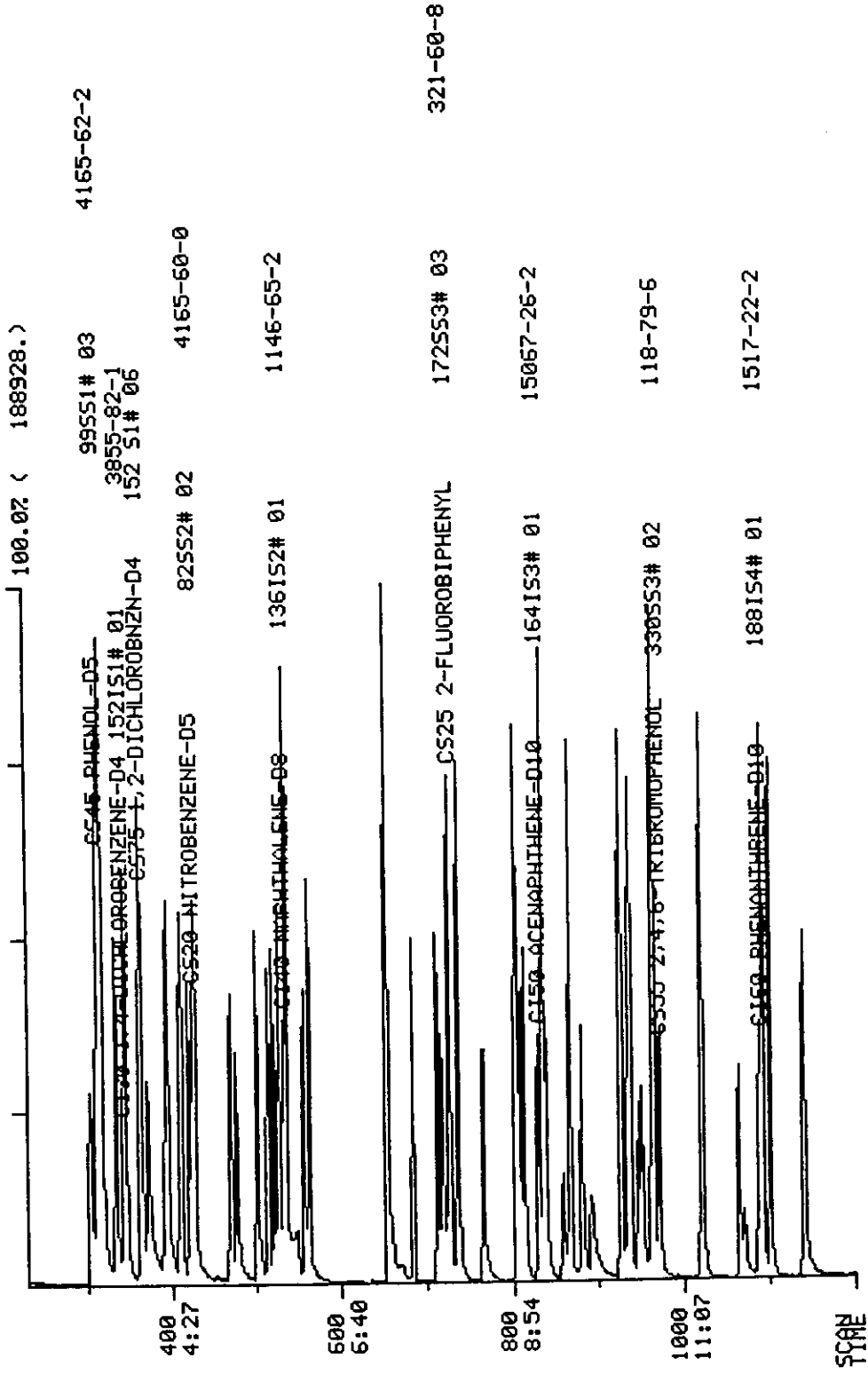
S3 2 80, 0.262

S3 2 160, 0.217

20, 0.261 50, 0.239 80, 0.233 120, 0.228

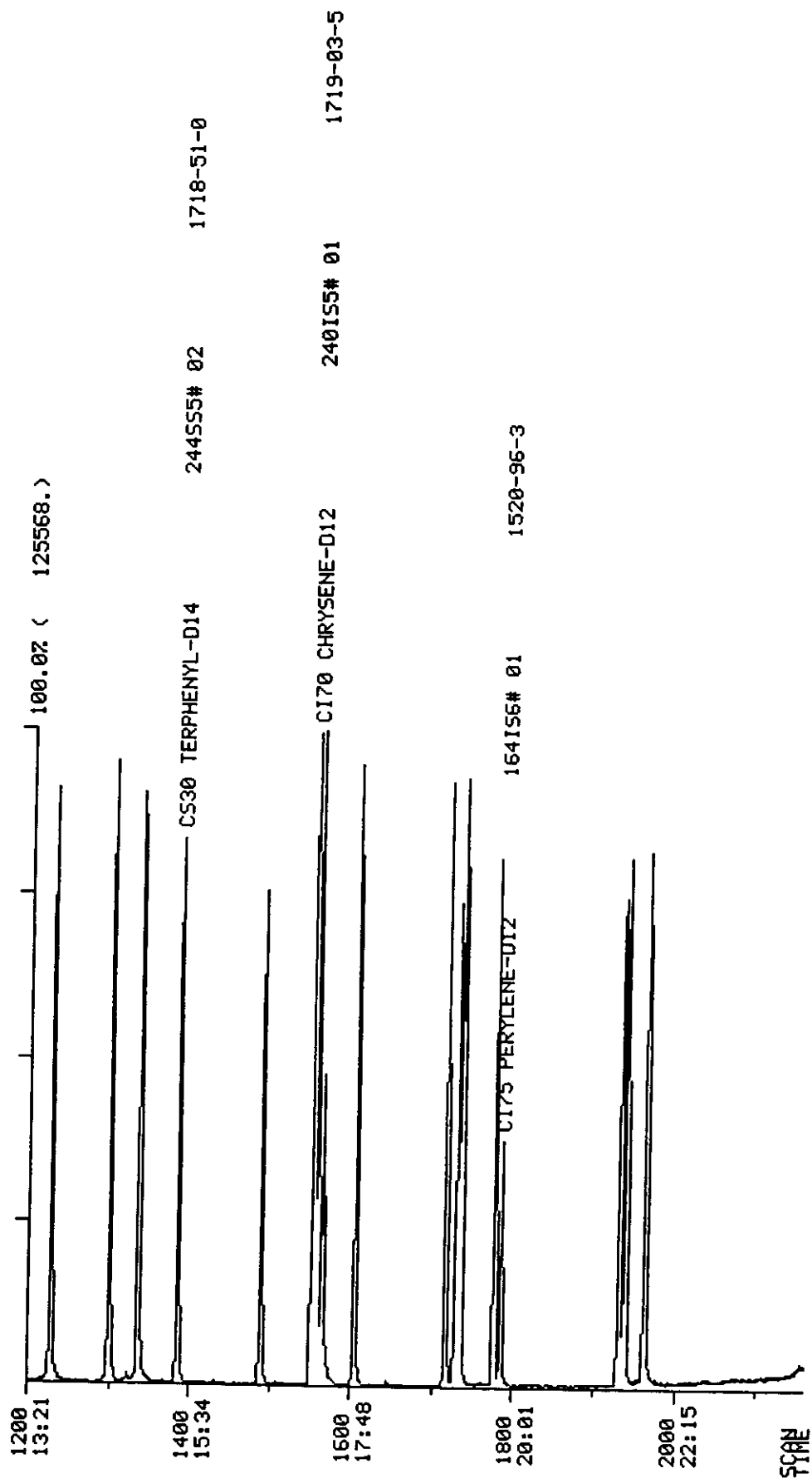


DATA FROM FILE: ST16980902 SCANS 230 TO 1200 ACQUIRED: 09/02/98 14:53:00  
 CALI: ST16980902 #3  
 SAMPLE: SST080 80UG/ML CALI III 082198A  
 CONDS.: INST. ID: F16



1631

DATA FROM FILE: ST16980902 SCANS 1200 TO 2158 ACQUIRED: 09/02/98 14:53:00  
 CALI: ST16980902 #3  
 SAMPLE: SSTD080 80UG/ML CALI III 082198A  
 CONDS.: INST. ID: F16



Data: ST16980902.T1

09/02/98 14:53:00

Sample: SST0080 80UG/ML CALI III 082198A

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from average of whole .RL

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	1521S1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	1361S2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	1641S3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	1881S4# 01	1517-22-2
5	C170 CHRYSENE-D12	2401S5# 01	1719-03-5
6	C175 PERYLENE-D12	1641S6# 01	1520-96-3
7	CS20 NITROBENZENE-D5	82SS2# 02	4165-60-0
8	CS25 2-FLUOROBIPHENYL	172SS3# 03	321-60-8
9	CS30 TERPHENYL-D14	244SS5# 02	1718-51-0
10	CS45 PHENOL-D5	99SS1# 03	4165-62-2
11	CS50 2-FLUOROPHENOL	112SS1# 02	367-12-4
12	CS55 2,4,6-TRIBROMOPHENOL	330SS3# 02	118-79-6
13	CS70 2-CHLOROPHENOL-D4	132 S1# 05	
14	CS75 1,2-DICHLOROBNZN-D4	152 S1# 06	
15	HEXACHLOROBENZENE-C13		
16	C310 N-NITROSODIMETHYLAMINE	74 S1# 15	62-75-9
17	C320 ANILINE	93 S1#100	62-53-3
18	C315 PHENOL	94 S1#105CC	108-95-2
19	C325 BIS(2-CHLOROETHYL)ETHER	93 S1#110	111-44-4
20	C330 2-CHLOROPHENOL	128 S2#115	95-57-8
21	C335 1,3-DICHLOROBENZENE	146 S1#125	541-73-1
22	C340 1,4-DICHLOROBENZENE	146 S1#130CC	106-46-7
23	C345 BENZYL ALCOHOL	108 S1#145	100-51-6
24	C350 1,2-DICHLOROBENZENE	146 S1#150	95-50-1
25	C355 2-METHYLPHENOL	108 S1#160	95-48-7
26	C360 2,2'-OXYBIS(1-CLPROPAN)	45 S1#165	108-60-1
27	C365 4-METHYLPHENOL	108 S1#185	106-44-5
28	C370 N-NITROSO-DI-N-PROPYLAM	70 S1#190SP	621-64-7
29	C375 HEXACHLOROETHANE	117 S1#200	67-72-1
30	C410 NITROBENZENE	77 S2# 10	98-95-3
31	C415 ISOPHORONE	82 S2# 20	75-59-1
32	C420 2-NITROPHENOL	139 S2# 25CC	88-75-5
33	C425 2,4-DIMETHYLPHENOL	107 S2# 30	105-67-9
34	C435 BIS(2-CHLOROETHOXY)METH	93 S2# 35	111-91-1
35	C440 2,4-DICHLOROPHENOL	162 S2# 40	120-83-2
36	C430 BENZOIC ACID	122 S2# 45	65-85-0
37	C445 1,2,4-TRICHLOROBENZENE	180 S2# 55	120-82-1
38	C450 NAPHTHALENE	128 S2# 60	91-20-3
39	C455 4-CHLOROANALINE	127 S2# 80	106-47-8
40	C460 HEXACHLOROBUTADIENE	225 S2# 95CC	87-68-3
41	C465 4-CHLORO-3-METHYLPHENO	107 S2#130CC	59-50-7
42	C470 2-METHYLNAPHTHALENE	142 S2#145	91-57-6
43	C510 HEXACHLOROCYCLOPENTADI	237 S3# 20	77-47-4
44	C515 2,4,6-TRICHLOROPHENOL	196 S3# 25CC	88-06-2
45	C520 2,4,5-TRICHLOROPHENOL	196 S3# 30	95-95-4
46	C525 2-CHLORONAPHTHALENE	162 S3# 40	91-58-7
47	C530 2-NITROANALINE	65 S3# 45	88-74-4

No	Name	S3#	
48	C535 DIMETHYLPHthalate	163 S3# 55	131-11-3
49	C540 ACENAPHTHYLENE	152 S3# 65	208-96-8
50	C543 2,6-DINITROToluene	165 S3# 70	606-20-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	340	3:47	1	1.000	A BB	23319.	40.000 UG/ML	0.61
2	136	531	5:54	2	1.000	A BB	79260.	40.000 UG/ML	0.61
3	164	829	9:13	3	1.000	A BB	43605.	40.000 UG/ML	0.61
4	188	1087	12:05	4	1.000	A BV	76218.	40.000 UG/ML	0.61
5	240	1553	17:16	5	1.000	A BB	51424.	40.000 UG/ML	0.61
6	264	1784	19:51	6	1.000	A BB	44306.	40.000 UG/ML	0.61
7	82	423	4:42	2	0.797	A BB	77010.	82.769 UG/ML	1.27
8	172	724	8:03	3	0.873	A BB	113630.	84.449 UG/ML	1.30
9	244	1384	15:24	5	0.891	A BB	107689.	84.168 UG/ML	1.29
10	99	310	3:27	1	0.912	A BB	88631.	75.060 UG/ML	1.15
11	112	197	2:11	1	0.579	A BB	56775.	71.178 UG/ML	1.09
12	330	970	10:47	3	1.170	A BB	22822.	88.836 UG/ML	1.36
13	132	313	3:29	1	0.921	A BB	66336.	76.805 UG/ML	1.18
14	152	360	4:00	1	1.059	A BB	42899.	79.981 UG/ML	1.23
15	NOT FOUND								
16	74	104	1:09	1	0.306	A BB	45851.	77.168 UG/ML	1.18
17	93	302	3:22	1	0.888	A BV	78198.	72.648 UG/ML	1.11
18	94	311	3:28	1	0.915	A VB	90423.	78.820 UG/ML	1.21
19	93	316	3:31	1	0.929	A VB	67521.	84.679 UG/ML	1.30
20	128	315	3:30	1	0.926	A BB	72156.	82.688 UG/ML	1.27
21	146	332	3:42	1	0.976	A BV	74894.	81.001 UG/ML	1.24
22	146	343	3:49	1	1.009	A VB	75730.	80.312 UG/ML	1.23
23	108	370	4:07	1	1.088	A BB	35440.	71.152 UG/ML	1.09
24	146	362	4:02	1	1.065	A BB	70366.	80.476 UG/ML	1.24
25	108	392	4:22	1	1.153	A BB	51172.	74.726 UG/ML	1.15
26	45	390	4:20	1	1.147	A BB	114359.	77.250 UG/ML	1.19
27	108	418	4:39	1	1.229	A BB	52957.	71.200 UG/ML	1.09
28	70	411	4:34	1	1.209	A BV	40110.	75.951 UG/ML	1.17
29	117	408	4:32	1	1.200	A BB	32844.	79.548 UG/ML	1.22
30	77	426	4:44	2	0.802	A VB	74224.	84.170 UG/ML	1.29
31	82	468	5:12	2	0.881	A BB	118401.	80.599 UG/ML	1.24
32	139	475	5:17	2	0.895	A VB	34790.	82.243 UG/ML	1.26
33	107	499	5:33	2	0.940	A BB	59805.	91.475 UG/ML	1.40
34	93	512	5:42	2	0.964	A BB	81550.	80.594 UG/ML	1.24
35	162	518	5:46	2	0.976	A BB	48941.	80.305 UG/ML	1.23
36	122	546	6:04	2	1.028	qedt	24441.	44.726 UG/ML	0.69
37	180	525	5:50	2	0.989	A BB	55810.	85.111 UG/ML	1.31
38	128	534	5:56	2	1.006	A BB	173260.	85.538 UG/ML	1.31
39	127	554	6:10	2	1.043	A BB	74460.	82.809 UG/ML	1.27
40	225	561	6:14	2	1.056	A BB	28856.	86.622 UG/ML	1.33
41	107	651	7:14	2	1.226	A BB	53671.	80.198 UG/ML	1.23
42	142	652	7:15	2	1.228	A BV	154259.	103.624 UG/ML	1.59
43	237	681	7:34	3	0.821	A BB	34014.	84.003 UG/ML	1.29
44	196	709	7:53	3	0.855	A BV	34776.	79.050 UG/ML	1.21
45	196	715	7:57	3	0.862	A VB	36021.	80.179 UG/ML	1.23
46	162	734	8:10	3	0.885	A BB	111646.	85.891 UG/ML	1.32
47	65	764	8:30	3	0.922	A BB	43674.	79.829 UG/ML	1.23
48	163	807	8:59	3	0.973	A BB	123549.	84.524 UG/ML	1.30
49	152	803	8:56	3	0.969	A BB	174241.	86.940 UG/ML	1.33
50	165	813	9:03	3	0.981	A BB	30775.	87.184 UG/ML	1.34

M/S 1/14

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:47	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:16	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:51	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	4:42	1.00	0.797	1.00	82.77	50.00	0.777	0.470	1.66
8	8:03	1.00	0.873	1.00	84.45	50.00	2.085	1.234	1.69
9	15:24	1.00	0.891	1.00	84.17	50.00	1.675	0.995	1.68
10	3:27	1.00	0.912	1.00	75.06	50.00	3.041	2.025	1.50
11	2:11	1.00	0.579	1.00	71.18	50.00	1.948	1.368	1.42
12	10:47	1.00	1.170	1.00	88.84	50.00	0.419	0.236	1.78
13	3:29	1.00	0.921	1.00	76.81	50.00	2.276	1.482	1.54
14	4:00	1.00	1.059	1.00	79.98	50.00	1.472	0.920	1.60
15	12:39		0.941						
16	1:09	1.00	0.306	1.00	77.17	50.00	1.573	1.019	1.54
17	3:22	1.00	0.888	1.00	72.65	50.00	2.683	1.846	1.45
18	3:28	1.00	0.915	1.00	78.82	50.00	3.102	1.968	1.58
19	3:31	1.00	0.929	1.00	84.68	50.00	2.316	1.368	1.69
20	3:30	1.00	0.926	1.00	82.69	50.00	2.475	1.497	1.65
21	3:42	1.00	0.976	1.00	81.00	50.00	2.569	1.586	1.62
22	3:49	1.00	1.009	1.00	80.31	50.00	2.598	1.617	1.61
23	4:07	1.00	1.088	1.00	71.15	50.00	1.216	0.854	1.42
24	4:02	1.00	1.065	1.00	80.48	50.00	2.414	1.500	1.61
25	4:22	1.00	1.153	1.00	74.73	50.00	1.756	1.175	1.49
26	4:20	1.00	1.147	1.00	77.25	50.00	3.923	2.539	1.55
27	4:39	1.00	1.229	1.00	71.20	50.00	1.817	1.276	1.42
28	4:34	1.00	1.209	1.00	75.95	50.00	1.376	0.906	1.52
29	4:32	1.00	1.200	1.00	79.55	50.00	1.127	0.708	1.59
30	4:44	1.00	0.802	1.00	84.17	50.00	0.749	0.445	1.68
31	5:12	1.00	0.881	1.00	80.60	50.00	1.195	0.741	1.61
32	5:17	1.00	0.895	1.00	82.24	50.00	0.351	0.213	1.64
33	5:33	1.00	0.940	1.00	91.48	50.00	0.604	0.330	1.83
34	5:42	1.00	0.964	1.00	80.59	50.00	0.823	0.511	1.61
35	5:46	1.00	0.976	1.00	80.31	50.00	0.494	0.308	1.61
36	6:04	1.00	1.028	1.00	44.73	50.00	0.247	0.276	0.89
37	5:50	1.00	0.989	1.00	85.11	50.00	0.563	0.331	1.70
38	5:56	1.00	1.006	1.00	85.54	50.00	1.749	1.022	1.71
39	6:10	1.00	1.043	1.00	82.81	50.00	0.752	0.454	1.66
40	6:14	1.00	1.056	1.00	86.62	50.00	0.291	0.168	1.73
41	7:14	1.00	1.226	1.00	80.20	50.00	0.542	0.338	1.60
42	7:15	1.00	1.228	1.00	103.62	50.00	1.557	0.751	2.07
43	7:34	1.00	0.821	1.00	84.00	50.00	0.624	0.371	1.68
44	7:53	1.00	0.855	1.00	79.05	50.00	0.638	0.404	1.58
45	7:57	1.00	0.862	1.00	80.18	50.00	0.661	0.412	1.60
46	8:10	1.00	0.885	1.00	85.89	50.00	2.048	1.192	1.72
47	8:30	1.00	0.922	1.00	79.83	50.00	0.801	0.502	1.60
48	8:59	1.00	0.973	1.00	84.52	50.00	2.267	1.341	1.69
49	8:56	1.00	0.969	1.00	86.94	50.00	3.197	1.838	1.74
50	9:03	1.00	0.981	1.00	87.18	50.00	0.565	0.324	1.74

Quantitation Report File: ST16980902

Data: ST16980902.T1

09/02/98 14:53:00

Sample: SSTD080 80UG/ML CALI III 082198A

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)  
 Resp. fac. from average of whole .RL

No	Name		
51	C545 3-NITROANILINE	138 S3# 75	99-09-2
52	C550 ACENAPHTHENE	153 S3# 80CC	83-32-9
53	C555 2,4-DINITROPHENOL	184 S3# 85SP	51-28-5
54	C565 DIBENZOFURAN	168 S3# 90	132-64-9
55	C560 4-NITROPHENOL	109 S3# 95SP	100-02-7
56	C570 2,4-DINITROTOLUENE	165 S3#105	121-14-2
57	C580 DIETHYLPHTHALATE	149 S3#130	84-66-2
58	C590 FLUORENE	166 S3#135	86-73-7
59	C585 4-CHLOROPHENYL-PHENYLE	204 S3#140	7005-72-3
60	C595 4-NITROANILINE	138 S3#150	100-01-6
61	C610 4,6-DINITRO-2-METHYLPH	198 S4# 10	534-52-1
62	C615 N-NITROSODIPHENYLAMINE	169 S4# 15CC	87-30-6
63	C620 AZOBENZENE	77 S4# 20	103-33-3
64	C625 4-BROMOPHENYL-PHENYLET	248 S4# 30	101-55-3
65	C630 HEXACHLOROBENZENE	284 S4# 40	118-74-1
66	C635 PENTACHLOROPHENOL	266 S4# 50	87-86-5
67	C640 PHENANTHRENE	178 S4# 65	85-01-8
68	C645 ANTHRACENE	178 S4# 70	120-12-7
69	C647 CARBAZOLE	167 S4# 80	86-74-8
70	C650 DI-N-BUTYLPHTHALATE	149 S4# 85	84-74-2
71	C655 FLUORANTHENE	202 S4#110CC	206-44-0
72	C710 BENZIDINE	184 S5# 10	92-81-5
73	C715 PYRENE	202 S5# 15	129-00-0
74	C720 BUTYLBENZYLPHTHALATE	149 S5# 40	85-68-7
75	C730 BENZO(A)ANTHRACENE	228 S5# 50	56-55-3
76	C725 3,3'-DICHLOROBENZIDINE	252 S5# 55	91-94-1
77	C740 CHRYSENE	228 S5# 60	218-01-9
78	C745 BIS(2-ETHYLHEXYL)PHTHA	149 S5# 65	117-81-7
79	C760 DI-N-OCTYL PHTHALATE	149 S6# 10CC	117-84-0
80	C765 BENZO(B)FLUORANTHENE	252 S6# 15	205-99-2
81	C770 BENZO(K)FLUORANTHENE	252 S6# 25	207-08-9
82	C775 BENZO(A)PYRENE	252 S6# 35	50-32-8
83	C780 INDENO(1,2,3-CD)PYRENE	276 S6# 55	193-39-5
84	C785 DIBENZ(A,H)ANTHRACENE	278 S6# 60	53-70-3
85	C790 BENZO(G,H,I)PERYLENE	276 S6# 65	191-24-2

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	138	838	9:19	3	1.011	A BB	33994.	81.477 UG/ML	1.25
52	153	835	9:17	3	1.007	A BV	108934.	86.246 UG/ML	1.32
53	184	858	9:33	3	1.035	A BB	14215.	68.887 UG/ML	1.06
54	168	867	9:39	3	1.046	A VB	156137.	85.811 UG/ML	1.32
55	109	892	9:55	3	1.076	A BB	11396.	60.551 UG/ML	0.93
56	165	881	9:48	3	1.063	A BB	40446.	93.400 UG/ML	1.43
57	149	935	10:24	3	1.128	A BB	128756.	86.290 UG/ML	1.32
58	166	926	10:18	3	1.117	A BB	125506.	88.713 UG/ML	1.36

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
59	204	938	10:26	3	1.131	A BB	59164.	88.575 UG/ML	1.36
60	138	947	10:32	3	1.142	A BB	26197.	78.275 UG/ML	1.20
61	198	951	10:35	4	0.875	A BB	20624.	81.985 UG/ML	1.26
62	169	963	10:43	4	0.886	A BV	82971.	79.067 UG/ML	1.21
63	77	965	10:44	4	0.888	A BB	179267.	80.357 UG/ML	1.23
64	248	1021	11:21	4	0.939	A BB	37178.	87.718 UG/ML	1.35
65	284	1020	11:21	4	0.938	A BB	43240.	87.455 UG/ML	1.34
66	266	1063	11:49	4	0.978	A BB	22429.	75.922 UG/ML	1.17
67	178	1092	12:09	4	1.005	A BV	166229.	84.485 UG/ML	1.30
68	178	1101	12:15	4	1.013	A VB	160282.	85.229 UG/ML	1.31
69	167	1139	12:40	4	1.048	A BB	123311.	84.715 UG/ML	1.30
70	149	1227	13:39	4	1.129	A BB	204955.	86.287 UG/ML	1.32
71	202	1299	14:27	4	1.195	A BB	144144.	84.053 UG/ML	1.29
72	184	1346	14:58	5	0.867	A BB	1996.	11.979 UG/ML	0.18
73	202	1335	14:51	5	0.860	A BB	140939.	82.890 UG/ML	1.27
74	149	1488	16:33	5	0.958	A BV	72332.	82.180 UG/ML	1.26
75	228	1552	17:16	5	0.999	A BV	125866.	87.038 UG/ML	1.34
76	252	1563	17:23	5	1.006	A BB	40227.	76.690 UG/ML	1.18
77	228	1558	17:20	5	1.003	A VB	108028.	82.832 UG/ML	1.27
78	149	1604	17:50	5	1.033	A BV	101978.	80.960 UG/ML	1.24
79	149	1715	19:05	6	0.961	A BV	171003.	80.218 UG/ML	1.23
80	252	1729	19:14	6	0.969	A BV	123130.	77.017 UG/ML	1.18
81	252	1734	19:17	6	0.972	A VB	111530.	86.867 UG/ML	1.33
82	252	1776	19:45	6	0.996	A BB	107418.	82.057 UG/ML	1.26
83	276	1932	21:29	6	1.083	A BV	113827.	80.352 UG/ML	1.23
84	278	1938	21:33	6	1.086	A BB	114484.	81.551 UG/ML	1.25
85	276	1963	21:50	6	1.100	A BB	119594.	81.129 UG/ML	1.25

No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	9:19	1.00	1.011	1.00	81.48	50.00	0.624	0.383	1.63
52	9:17	1.00	1.007	1.00	86.25	50.00	1.999	1.159	1.72
53	9:33	1.00	1.035	1.00	68.89	50.00	0.261	0.189	1.38
54	9:39	1.00	1.046	1.00	85.81	50.00	2.865	1.669	1.72
55	9:55	1.00	1.076	1.00	60.55	50.00	0.209	0.173	1.21
56	9:48	1.00	1.063	1.00	93.40	50.00	0.742	0.397	1.87
57	10:24	1.00	1.128	1.00	86.29	50.00	2.362	1.369	1.73
58	10:18	1.00	1.117	1.00	88.71	50.00	2.303	1.298	1.77
59	10:26	1.00	1.131	1.00	88.58	50.00	1.085	0.613	1.77
60	10:32	1.00	1.142	1.00	78.27	50.00	0.481	0.307	1.57
61	10:35	1.00	0.875	1.00	81.99	50.00	0.216	0.132	1.64
62	10:43	1.00	0.886	1.00	79.07	50.00	0.871	0.551	1.58
63	10:44	1.00	0.888	1.00	80.36	50.00	1.882	1.171	1.61
64	11:21	1.00	0.939	1.00	87.72	50.00	0.390	0.222	1.75
65	11:21	1.00	0.938	1.00	87.46	50.00	0.454	0.259	1.75
66	11:49	1.00	0.978	1.00	75.92	50.00	0.235	0.155	1.52
67	12:09	1.00	1.005	1.00	84.49	50.00	1.745	1.033	1.69
68	12:15	1.00	1.013	1.00	85.23	50.00	1.682	0.987	1.70
69	12:40	1.00	1.048	1.00	84.72	50.00	1.294	0.764	1.69
70	13:39	1.00	1.129	1.00	86.29	50.00	2.151	1.247	1.73
71	14:27	1.00	1.195	1.00	84.05	50.00	1.513	0.900	1.68
72	14:58	1.00	0.867	1.00	11.98	50.00	0.031	0.130	0.24
73	14:51	1.00	0.860	1.00	82.89	50.00	2.193	1.323	1.66
74	16:33	1.00	0.958	1.00	82.18	50.00	1.125	0.685	1.64
75	17:16	1.00	0.999	1.00	87.04	50.00	1.958	1.125	1.74
76	17:23	1.00	1.006	1.00	76.69	50.00	0.626	0.408	1.53
77	17:20	1.00	1.003	1.00	82.83	50.00	1.681	1.014	1.66

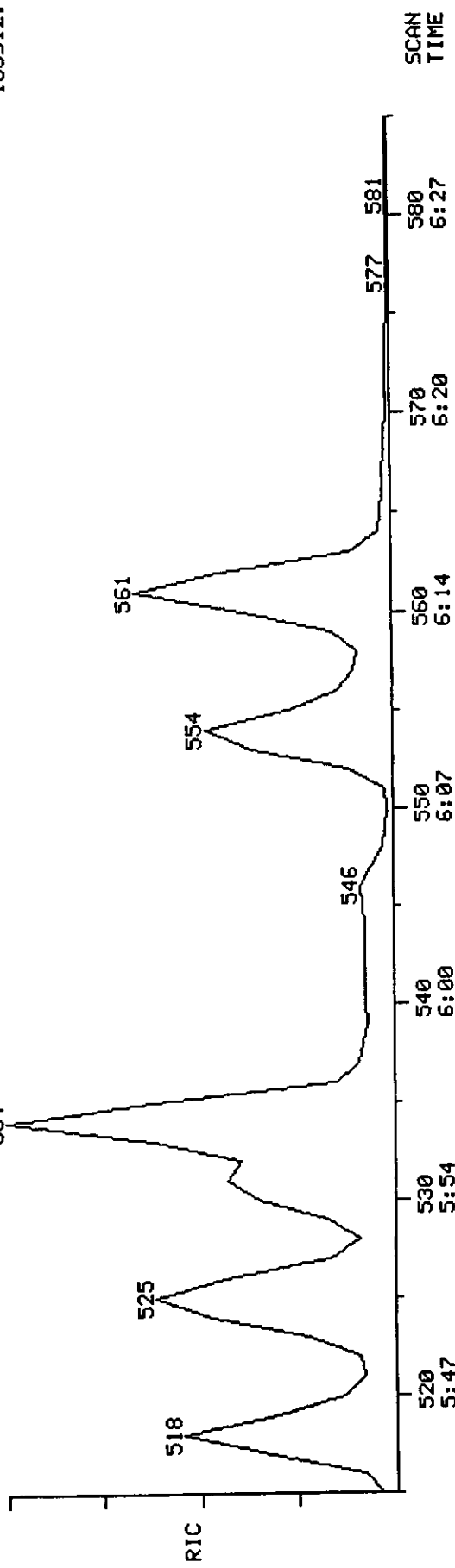
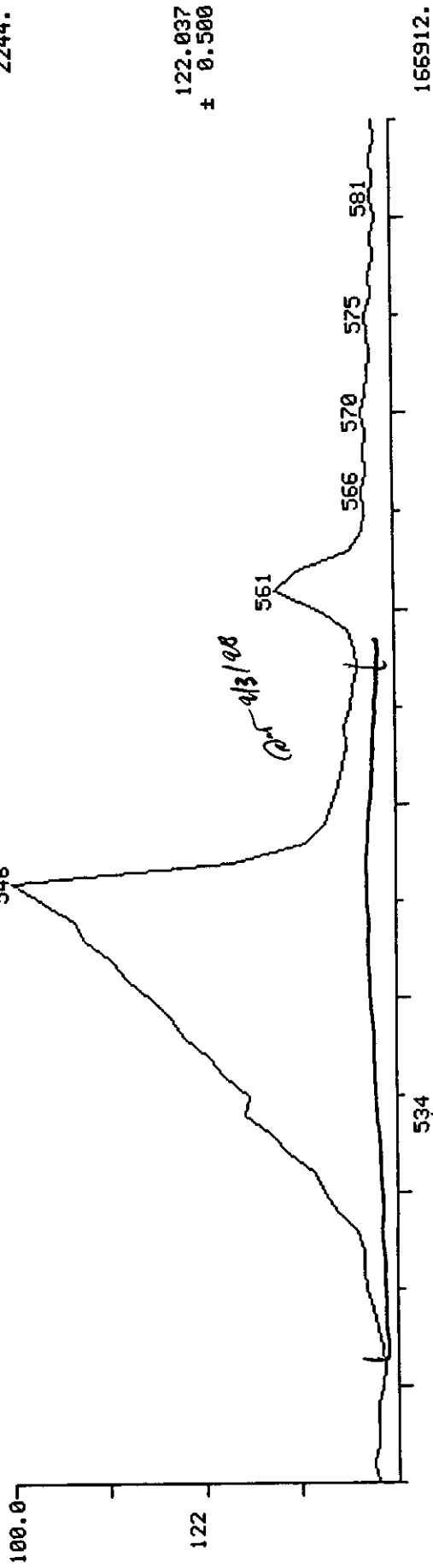
No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
78	17:50	1.00	1.033	1.00	80.96	50.00	1.586	0.980	1.62
79	19:05	1.00	0.961	1.00	80.22	50.00	3.088	1.925	1.60
80	19:14	1.00	0.969	1.00	77.02	50.00	2.223	1.443	1.54
81	19:17	1.00	0.972	1.00	86.87	50.00	2.014	1.159	1.74
82	19:45	1.00	0.996	1.00	82.06	50.00	1.940	1.182	1.64
83	21:29	1.00	1.083	1.00	80.35	50.00	2.055	1.279	1.61
84	21:33	1.00	1.086	1.00	81.55	50.00	2.067	1.267	1.63
85	21:50	1.00	1.100	1.00	81.13	50.00	2.159	1.331	1.62

**MANUAL EDIT CODES**  
**1. PEAK NOT FOUND**  
**2. POOR CHROMATOGRAPH**  
**3. WRONG ISOMER**  
**ALL MANUAL EDITS MUST BE**  
**INITIALED, DATED, AND CODED**



RIC+MASS CHROMATOGRAM  
 09/02/98 14:53:00 DATA: ST16980902 #1  
 CALI: ST16980902 #3  
 SAMPLE: SSTD080 80UG/ML CALI III 082198A  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3  
 SCANS 515 TO 585

2244.



10391

QUANTERRA GC/MS  
SEMIVOLATILE APPENDIX IX CALIBRATION CHE

CONTIN. CALIBRATION: 80 UG/ML APPENDIX ST16980902A 09/02/98 15:24:00  
 INITIAL CALIBRATION: 10 UG/ML APPENDIX ST16980831B 08/31/98 13:35:00  
 INITIAL CALIBRATION: 20 UG/ML APPENDIX ST16980831C 08/31/98 14:02:00  
 INITIAL CALIBRATION: 50 UG/ML APPENDIX ST16980831D 08/31/98 14:32:00  
 INITIAL CALIBRATION: 80 UG/ML APPENDIX ST16980831A 08/31/98 13:07:00  
 INITIAL CALIBRATION: 120 UG/ML APPENDIX ST16980831E 08/31/98 15:02:00  
 INITIAL CALIBRATION: 160 UG/ML APPENDIX ST16980831F 08/31/98 15:31:00

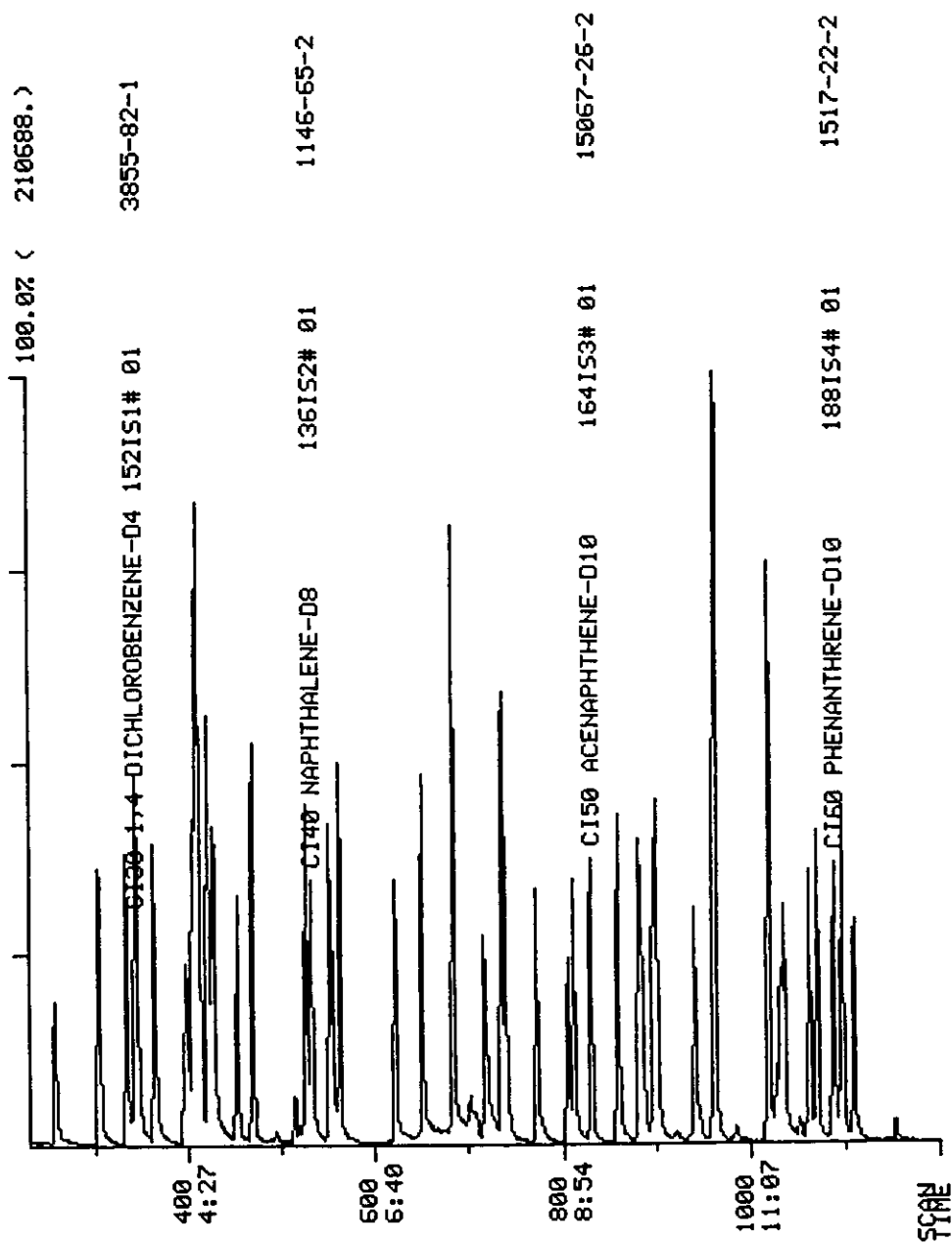
ST16980902A ST16980831B ST16980831C ST16980831D ST16980831A ST16980831E  
 ST16980831F

Lib #	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
S1 20	80,	1.559	20,	1.583	50,	1.578	80,	1.587	120,	1.449
S1 20	160,	1.576								
S1 30	80,	1.497	20,	1.563	50,	1.594	80,	1.577	120,	1.547
S1 30	160,	1.545								
S1 40	80,	0.792	20,	0.737	50,	0.823	80,	0.876	120,	0.860
S1 40	160,	0.870								
S1 60	80,	0.561	20,	0.688	50,	0.685	80,	0.602	120,	0.619
S1 60	160,	0.604								
S1 70	80,	0.512	20,	0.506	50,	0.521	80,	0.549	120,	0.547
S1 70	160,	0.555								
S1 85	80,	1.091	20,	1.008	50,	1.120	80,	1.168	120,	1.190
S1 85	160,	1.244								
S1 95	80,	0.582	20,	0.612	50,	0.605	80,	0.605	120,	0.595
S1 95	160,	0.592								
S1 170	80,	1.856	20,	2.024	50,	1.960	80,	1.945	120,	1.903
S1 170	160,	1.903								
S1 175	80,	0.540	20,	0.555	50,	0.575	80,	0.562	120,	0.569
S1 175	160,	0.575								
S1 180	80,	0.773	20,	0.838	50,	0.821	80,	0.805	120,	0.817
S1 180	160,	0.816								
S1 182	80,	2.218	20,	2.130	50,	2.339	80,	2.408	120,	2.391
S1 182	160,	2.394								
S1 195	80,	2.043	20,	2.463	50,	2.228	80,	2.129	120,	2.032
S1 195	160,	1.949								
S2 15	80,	0.324	20,	0.312	50,	0.326	80,	0.327	120,	0.337
S2 15	160,	0.329								
S2 50	80,	1.060	20,	1.575	50,	1.353	80,	1.036	120,	1.276
S2 50	160,	1.184								
S2 85	80,	0.272	20,	0.234	50,	0.264	80,	0.280	120,	0.279
S2 85	160,	0.285								
S2 90	80,	0.203	20,	0.196	50,	0.201	80,	0.197	120,	0.198
S2 90	160,	0.198								

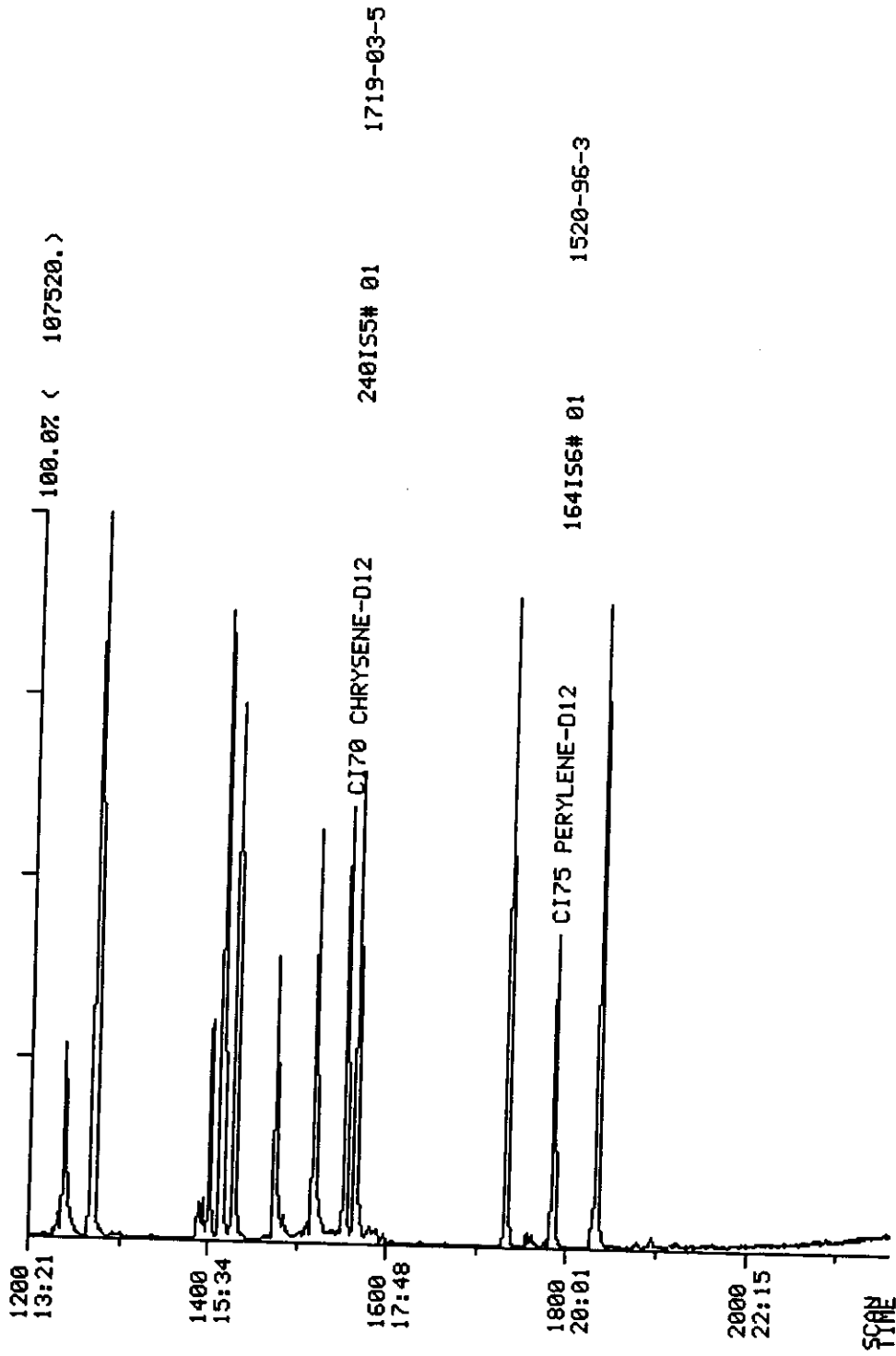
S2 115 80, 0.040	20, 0.163	50, 0.167	80, 0.038	120, 0.165
S2 115 160, 0.121				
S2 120 80, 0.215	20, 0.205	50, 0.213	80, 0.225	120, 0.224
S2 120 160, 0.224				
S2 140 80, 0.248	20, 0.246	50, 0.247	80, 0.250	120, 0.245
S2 140 160, 0.250				
S3 10 80, 0.427	20, 0.431	50, 0.434	80, 0.430	120, 0.416
S3 10 160, 0.419				
S3 15 80, 0.045	20, 0.046	50, 0.042	80, 0.043	120, 0.044
S3 15 160, 0.044				
S3 35 80, 0.191	20, 0.201	50, 0.195	80, 0.196	120, 0.190
S3 35 160, 0.192				
S3 42 80, 1.119	20, 1.098	50, 1.129	80, 1.528	120, 1.134
S3 42 160, 1.488				
S3 50 80, 0.432	20, 0.403	50, 0.429	80, 0.447	120, 0.404
S3 50 160, 0.389				
S3 60 80, 0.202	20, 0.191	50, 0.202	80, 0.210	120, 0.212
S3 60 160, 0.204				
S3 100 80, 0.498	20, 0.504	50, 0.483	80, 0.493	120, 0.469
S3 100 160, 0.471				
S3 110 80, 0.746	20, 0.859	50, 0.889	80, 0.765	120, 0.872
S3 110 160, 0.769				
S3 115 80, 0.900	20, 0.974	50, 1.041	80, 0.996	120, 0.980
S3 115 160, 0.881				
S3 120 80, 0.296	20, 0.277	50, 0.302	80, 0.310	120, 0.306
S3 120 160, 0.290				
S3 145 80, 0.338	20, 0.345	50, 0.368	80, 0.381	120, 0.367
S3 145 160, 0.316				
S4 25 80, 0.164	20, 0.149	50, 0.162	80, 0.172	120, 0.166
S4 25 160, 0.159				
S4 35 80, 0.293	20, 0.305	50, 0.324	80, 0.316	120, 0.330
S4 35 160, 0.311				
S4 37 80, 0.094	20, 0.088	50, 0.087	80, 0.089	120, 0.089
S4 37 160, 0.097				
S4 45 80, 0.577	20, 0.643	50, 0.664	80, 0.602	120, 0.590
S4 45 160, 0.546				
S4 55 80, 0.300	20, 0.312	50, 0.315	80, 0.301	120, 0.291
S4 55 160, 0.277				
S4 60 80, 0.086	20, 0.081	50, 0.079	80, 0.079	120, 0.082
S4 60 160, 0.081				
S4 75 80, 0.165	20, 0.130	50, 0.165	80, 0.163	120, 0.159

S4 75 160, 0.150					
S4 100 80, 0.074		20, 0.058	50, 0.074	80, 0.076	120, 0.079
S4 100 160, 0.075					
S4 105 80, 0.237		20, 0.424	50, 0.339	80, 0.205	120, 0.279
S4 105 160, 0.277					
S4 106 80, 0.129		20, 0.125	50, 0.129	80, 0.128	120, 0.127
S4 106 160, 0.124					
S4 120 80, 0.321	10, 0.376	20, 0.351	50, 0.329	80, 0.308	120, 0.295
S4 120 160, 0.281					
S5 20 80, 0.064		20, 0.057	50, 0.063	80, 0.061	120, 0.064
S5 20 160, 0.061					
S5 25 80, 0.085		20, 0.081	50, 0.090	80, 0.090	120, 0.087
S5 25 160, 0.085					
S5 30 80, 0.400		20, 0.417	50, 0.425	80, 0.420	120, 0.412
S5 30 160, 0.420					
S5 35 80, 0.352		20, 0.522	50, 0.450	80, 0.406	120, 0.364
S5 35 160, 0.346					
S5 37 80, 0.103		20, 0.123	50, 0.087	80, 0.089	120, 0.056
S5 37 160, 0.054					
S5 45 80, 0.435		20, 0.404	50, 0.450	80, 0.438	120, 0.429
S5 45 160, 0.440					
S5 85 80, 0.554		20, 0.586	50, 0.576	80, 0.549	120, 0.515
S5 85 160, 0.514					
S6 20 80, 0.599		20, 0.609	50, 0.615	80, 0.610	120, 0.608
S6 20 160, 0.609					

DATA FROM FILE: ST16980902A SCANS 230 TO 1200 ACQUIRED: 09/02/98 15:24:00  
CALI: ST16980902A #3  
SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16



DATA FROM FILE: ST16980902A SCANS 1200 TO 2158 ACQUIRED: 09/02/98 15:24:00  
CALI: ST16980902A #3  
SAMPLE: 80UG/ML AP9 062598F  
CONDS.: INST. ID: F16



Data: ST16980902A.TI

09/02/98 15:24:00

Sample: 80UG/ML AP9 062598F

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from average of whole .RL

No	Name		
1	C130 1,4-DICHLOROBENZENE-D4	152IS1# 01	3855-82-1
2	C140 NAPHTHALENE-D8	136IS2# 01	1146-65-2
3	C150 ACENAPHTHENE-D10	164IS3# 01	15067-26-2
4	C160 PHENANTHRENE-D10	188IS4# 01	1517-22-2
5	C170 CHRYSENE-D12	240IS5# 01	1719-03-5
6	C175 PERYLENE-D12	164IS6# 01	1520-96-3
7	PYRIDINE	79 S1# 20	110-86-1
8	2-PICOLINE	93 S1# 30	109-06-8
9	N-NITROSOMETHYLETHYLAMINE	42 S1# 40	10595-95-6
10	METHYLMETHANESULFONATE	80 S1# 60	
11	N-NITROSODIETHYLAMINE	102 S1# 70	55-18-5
12	ETHYLMETHANE SULFONATE	79 S1# 85	66-27-3
13	PENTACHLOROETHANE	117 S1# 95	76-01-7
14	C361 ACETOPHENONE	105 S1#170	
15	N-NITROSPYRROLIDINE	100 S1#175	930-55-2
16	N-NITROSOMORPHOLINE	56 S1#180	59-89-2
17	3-METHYL PHENOL	108 S1#182	
18	O-TOLUIDINE	106 S1#195	95-53-4
19	N-NITROPIPERIDINE	42 S2# 15	100-75-4
20	AA-DIMETHYLPHENETHYLAMINE	58 S2# 50	122-09-8
21	2,6-DICHLOROPHENOL	162 S2# 85	87-65-0
22	HEXACHLOROPROPENE	213 S2# 90	
23	P-PHENYLENE DIAMINE	108 S2#115	106-50-3
24	N-NITROSODI-N-BUTYLAMINE	84 S2#120	924-16-3
25	SAFROLE	152 S2#140	94-59-7
26	1,2,4,5-TETRACHLOROBENZENE	214 S3# 10	95-94-3
27	ISOSAFROLE (#1)	162 S3# 15	120-58-1
28	ISOSAFROLE (#2)	104 S3# 35	120-58-1
29	1-CHLORONAPHTHALENE	162 S3# 42	90-13-1
30	1,4-NAPHTHOQUINONE	158 S3# 50	130-15-4
31	1,3-DINITROBENZENE	168 S3# 60	99-65-0
32	PENTACHLOROBENZENE	250 S3#100	708-93-5
33	1-NAPHTHYLAMINE	143 S3#110	134-32-7
34	2-NAPHTHYLAMINE	143 S3#115	91-59-8
35	2,3,4,6-TETRACHLOROPHENOL	232 S3#120	58-90-2
36	5-NITRO-O-TOLUIDINE	152 S3#145	99-52-5
37	SYM-TRINITROBENZENE	75 S4# 25	
38	PHENACETIN	108 S4#35	62-44-2
39	DIALATE	234 S4# 37	
40	4-AMINOBIIPHENYL	169 S4# 45	92-67-1
41	PRONAMIDE	173 S4# 55	23950-58-5
42	PENTACHLORONITROBENZENE	237 S4# 60	82-68-8
43	ZSECBUTYL-4,6-DINITROPHENOL	211 S4# 75	88-85-7
44	4-NITROQUINOLINE-1-OXIDE	190 S4#100	
45	METHAPRYLENE	58 S4#105	91-80-5
46	ISODRIN		
47	CHLOROBENZILATE	139 S4#120	

No	Name	185 S5# 20	140-57-8
48	ARAMITE (#1)	185 S5# 25	140-57-8
49	ARAMITE (#2)	120 S5# 30	60-11-7

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
1	152	341	3:48	1	1.000	A BV	23430.	40.000 UG/ML	0.99
2	136	530	5:54	2	1.000	A BB	82177.	40.000 UG/ML	0.99
3	164	829	9:13	3	1.000	A BB	47233.	40.000 UG/ML	0.99
4	188	1087	12:05	4	1.000	A BB	81529.	40.000 UG/ML	0.99
5	240	1552	17:16	5	1.000	A BV	57252.	40.000 UG/ML	0.99
6	264	1783	19:50	6	1.000	A BB	46438.	40.000 UG/ML	0.99
7	79	103	1:09	1	0.302	A BB	73075.	78.441 UG/ML	1.94
8	93	146	1:37	1	0.428	A BB	70141.	76.521 UG/ML	1.90
9	42	157	1:45	1	0.460	A BB	37109.	78.106 UG/ML	1.94
10	80	183	2:02	1	0.537	A BB	26293.	70.558 UG/ML	1.75
11	102	218	2:25	1	0.639	A BB	23982.	78.533 UG/ML	1.95
12	79	255	2:50	1	0.748	A BB	51129.	78.747 UG/ML	1.95
13	117	303	3:22	1	0.889	A BB	27292.	77.437 UG/ML	1.92
14	105	404	4:30	1	1.185	A BV	86991.	76.796 UG/ML	1.90
15	100	406	4:31	1	1.191	A BV	25281.	78.050 UG/ML	1.93
16	56	412	4:35	1	1.208	A BB	36244.	75.399 UG/ML	1.87
17	108	420	4:40	1	1.232	A BB	103957.	78.816 UG/ML	1.95
18	106	409	4:33	1	1.199	A BV	95752.	75.665 UG/ML	1.88
19	42	452	5:02	2	0.853	A BV	53309.	80.966 UG/ML	2.01
20	58	706	7:51	2	1.332	qedt	174290.	66.024 UG/ML	1.64
21	162	553	6:09	2	1.043	A BB	44702.	84.515 UG/ML	2.09
22	213	549	6:06	2	1.036	A BV	33370.	81.974 UG/ML	2.03
23	108	648	7:12	2	1.223	qedt	6560.	24.410 UG/ML	0.60
24	84	620	6:54	2	1.170	A BB	35296.	80.627 UG/ML	2.00
25	162	648	7:12	2	1.223	A BB	40806.	80.192 UG/ML	1.99
26	214	682	7:35	3	0.823	A BB	40306.	78.792 UG/ML	1.95
27	162	699	7:46	3	0.843	A BB	4260.	82.650 UG/ML	2.05
28	104	738	8:13	3	0.890	A BB	18010.	78.310 UG/ML	1.94
29	162	734	8:10	3	0.885	A BV	105725.	67.939 UG/ML	1.68
30	158	770	8:34	3	0.929	A BB	40775.	83.300 UG/ML	2.06
31	168	804	8:57	3	0.970	A BB	19068.	81.768 UG/ML	2.03
32	250	858	9:33	3	1.035	A BB	47076.	81.449 UG/ML	2.02
33	143	883	9:49	3	1.065	A BB	70446.	71.254 UG/ML	1.77
34	143	899	10:00	3	1.084	A BB	85044.	74.553 UG/ML	1.85
35	232	896	9:58	3	1.081	A BB	27957.	79.729 UG/ML	1.98
36	152	941	10:28	3	1.135	A BB	31963.	76.138 UG/ML	1.89
37	75	1026	11:25	4	0.944	A BB	26738.	81.214 UG/ML	2.01
38	108	1035	11:31	4	0.952	A BB	47788.	73.971 UG/ML	1.83
39	234	1018	11:19	4	0.937	A BV	15382.	83.878 UG/ML	2.08
40	169	1069	11:53	4	0.983	A BB	94094.	75.804 UG/ML	1.88
41	173	1096	12:11	4	1.008	A BB	48872.	80.082 UG/ML	1.98
42	237	1061	11:48	4	0.976	A BB	13992.	85.418 UG/ML	2.12
43	211	1108	12:19	4	1.019	A VB	26967.	86.160 UG/ML	2.14
44	190	1237	13:46	4	1.138	A BB	12009.	81.269 UG/ML	2.01
45	58	1267	14:06	4	1.166	A BB	38678.	62.277 UG/ML	1.54
46	193	1270	14:08	4	1.168	A BB	21038.	81.494 UG/ML	2.02
47	139	1428	15:53	4	1.314	A BB	52292.	79.309 UG/ML	1.97
48	185	1400	15:34	5	0.902	A VB	7298.	83.488 UG/ML	2.07
49	185	1414	15:44	5	0.911	A BB	9760.	78.798 UG/ML	1.95
50	120	1412	15:42	5	0.910	A BB	45827.	76.447 UG/ML	1.89



No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
1	3:48	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
2	5:54	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
3	9:13	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
4	12:05	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
5	17:16	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
6	19:50	1.00	1.000	1.00	40.00	40.00	1.000	1.000	1.00
7	1:09	1.00	0.302	1.00	78.44	50.00	2.495	1.590	1.57
8	1:37	1.00	0.428	1.00	76.52	50.00	2.395	1.565	1.53
9	1:45	1.00	0.460	1.00	78.11	50.00	1.267	0.811	1.56
10	2:02	1.00	0.537	1.00	70.56	50.00	0.898	0.636	1.41
11	2:25	1.00	0.639	1.00	78.53	50.00	0.819	0.521	1.57
12	2:50	1.00	0.748	1.00	78.75	50.00	1.746	1.108	1.57
13	3:22	1.00	0.889	1.00	77.44	50.00	0.932	0.602	1.55
14	4:30	1.00	1.185	1.00	76.80	50.00	2.970	1.934	1.54
15	4:31	1.00	1.191	1.00	78.05	50.00	0.863	0.553	1.56
16	4:35	1.00	1.208	1.00	75.40	50.00	1.238	0.821	1.51
17	4:40	1.00	1.232	1.00	78.82	50.00	3.550	2.252	1.58
18	4:33	1.00	1.199	1.00	75.66	50.00	3.269	2.160	1.51
19	5:02	1.00	0.853	1.00	80.97	50.00	0.519	0.320	1.62
20	7:51	1.00	1.332	1.00	66.02	50.00	1.697	1.285	1.32
21	6:09	1.00	1.043	1.00	84.51	50.00	0.435	0.257	1.69
22	6:06	1.00	1.036	1.00	81.97	50.00	0.325	0.198	1.64
23	7:12	1.00	1.223	1.00	24.41	50.00	0.064	0.131	0.49
24	6:54	1.00	1.170	1.00	80.63	50.00	0.344	0.213	1.61
25	7:12	1.00	1.223	1.00	80.19	50.00	0.397	0.248	1.60
26	7:35	1.00	0.823	1.00	78.79	50.00	0.683	0.433	1.58
27	7:46	1.00	0.843	1.00	82.65	50.00	0.072	0.044	1.65
28	8:13	1.00	0.890	1.00	78.31	50.00	0.305	0.195	1.57
29	8:10	1.00	0.885	1.00	67.94	50.00	1.791	1.318	1.36
30	8:34	1.00	0.929	1.00	83.30	50.00	0.691	0.415	1.67
31	8:57	1.00	0.970	1.00	81.77	50.00	0.323	0.197	1.64
32	9:33	1.00	1.035	1.00	81.45	50.00	0.797	0.489	1.63
33	9:49	1.00	1.065	1.00	71.25	50.00	1.193	0.837	1.43
34	10:00	1.00	1.084	1.00	74.55	50.00	1.440	0.966	1.49
35	9:58	1.00	1.081	1.00	79.73	50.00	0.474	0.297	1.59
36	10:28	1.00	1.135	1.00	76.14	50.00	0.541	0.356	1.52
37	11:25	1.00	0.944	1.00	81.21	50.00	0.262	0.162	1.62
38	11:31	1.00	0.952	1.00	73.97	50.00	0.469	0.317	1.48
39	11:19	1.00	0.937	1.00	83.88	50.00	0.151	0.090	1.68
40	11:53	1.00	0.983	1.00	75.80	50.00	0.923	0.609	1.52
41	12:11	1.00	1.008	1.00	80.08	50.00	0.480	0.299	1.60
42	11:48	1.00	0.976	1.00	85.42	50.00	0.137	0.080	1.71
43	12:19	1.00	1.019	1.00	86.16	50.00	0.265	0.154	1.72
44	13:46	1.00	1.138	1.00	81.27	50.00	0.118	0.073	1.63
45	14:06	1.00	1.166	1.00	62.28	50.00	0.380	0.305	1.25
46	14:08	1.00	1.168	1.00	81.49	50.00	0.206	0.127	1.63
47	15:53	1.00	1.314	1.00	79.31	50.00	0.513	0.323	1.59
48	15:34	1.00	0.902	1.00	83.49	50.00	0.102	0.061	1.67
49	15:44	1.00	0.911	1.00	78.80	50.00	0.136	0.087	1.58
50	15:42	1.00	0.910	1.00	76.45	50.00	0.640	0.419	1.53

Quantitation Report File: ST16980902A

Data: ST16980902A.T1

09/02/98 15:24:00

Sample: 80UG/ML AP9 062598F

Conds.: INST. ID: F16

Formula: 1UL INJ.

Instrument: F16

Weight: 0.000

Submitted by: QES

Analyst: DAT

Acct. No.: FIN

AMOUNT=AREA \* REF AMNT/(REF AREA \* RESP FACT)

Resp. fac. from average of whole .RL

No	Name			
51	3,3'-DIMETHYLBENZIDINE	212	S5# 35	
52	KEPONE	272	S5# 37	
53	2-ACETYLAMINOFLUORENE	181	S5# 45	53-96-3
54	3-METHYLCHOLANTHRENE	268	S5# 85	56-49-5
55	7,12-DIMETHYLBENZANTHRACENE	256	S6# 20	75-97-6
56	HEXACHLOROPHENE	196	S6# 30	70-30-4

No	m/z	Scan	Time	Ref	RRT	Meth	Area(Hght)	Amount	%Tot
51	212	1473	16:23	5	0.949	A BB	40357.	67.516 UG/ML	1.67
52	272	1504	16:44	5	0.969	qedt	11784.	100.447 UG/ML	2.49
53	181	1518	16:53	5	0.978	A BB	49800.	80.513 UG/ML	2.00
54	268	1833	20:23	5	1.181	A BB	63472.	80.915 UG/ML	2.01
55	256	1731	19:15	6	0.971	A BB	55616.	78.526 UG/ML	1.95
56	NOT FOUND								

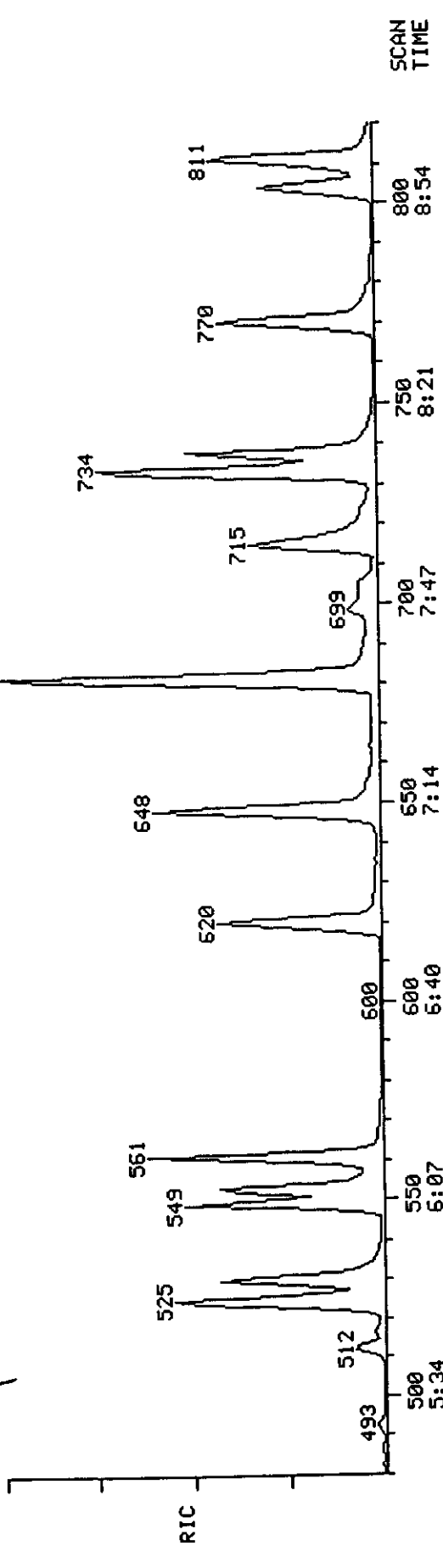
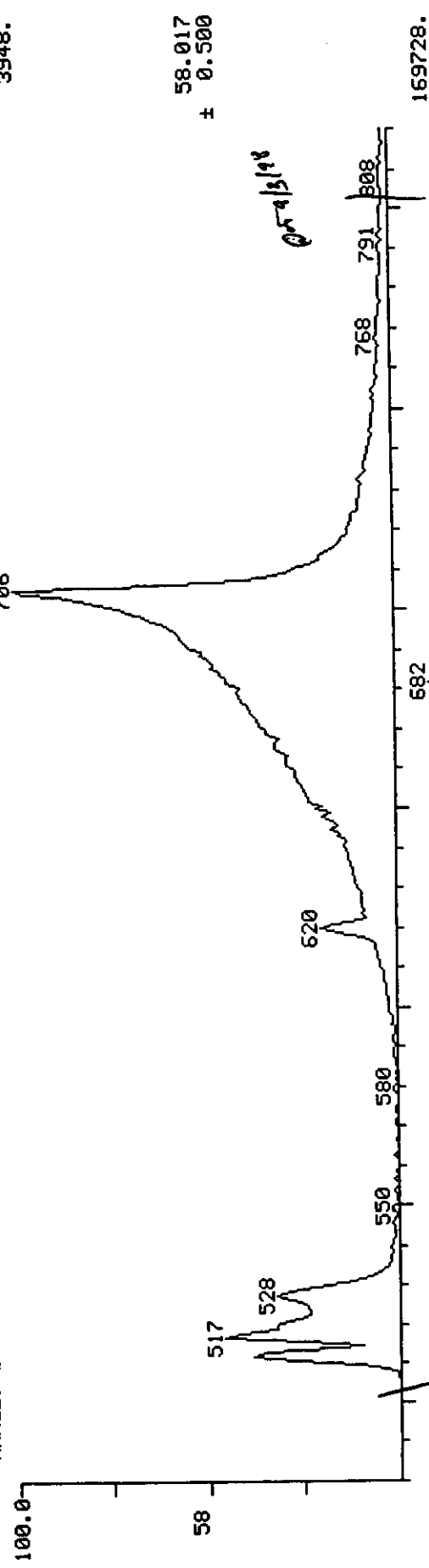
No	Ret(L)	Ratio	RRT(L)	Ratio	Amnt	Amnt(L)	R.Fac	R.Fac(L)	Ratio
51	16:23	1.00	0.949	1.00	67.52	50.00	0.564	0.418	1.35
52	16:44	1.00	0.969	1.00	100.45	50.00	0.165	0.082	2.01
53	16:53	1.00	0.978	1.00	80.51	50.00	0.696	0.432	1.61
54	20:23	1.00	1.181	1.00	80.91	50.00	0.887	0.548	1.62
55	19:15	1.00	0.971	1.00	78.53	50.00	0.958	0.610	1.57
56	21:11		0.992						

MANUAL EDIT CODES  
1. PEAK NOT FOUND  
2. POOR CHROMATOGRAPHY  
3. WRONG ISOMER  
ALL MANUAL EDITS MUST BE  
INITIALED, DATED, AND CODED

1648

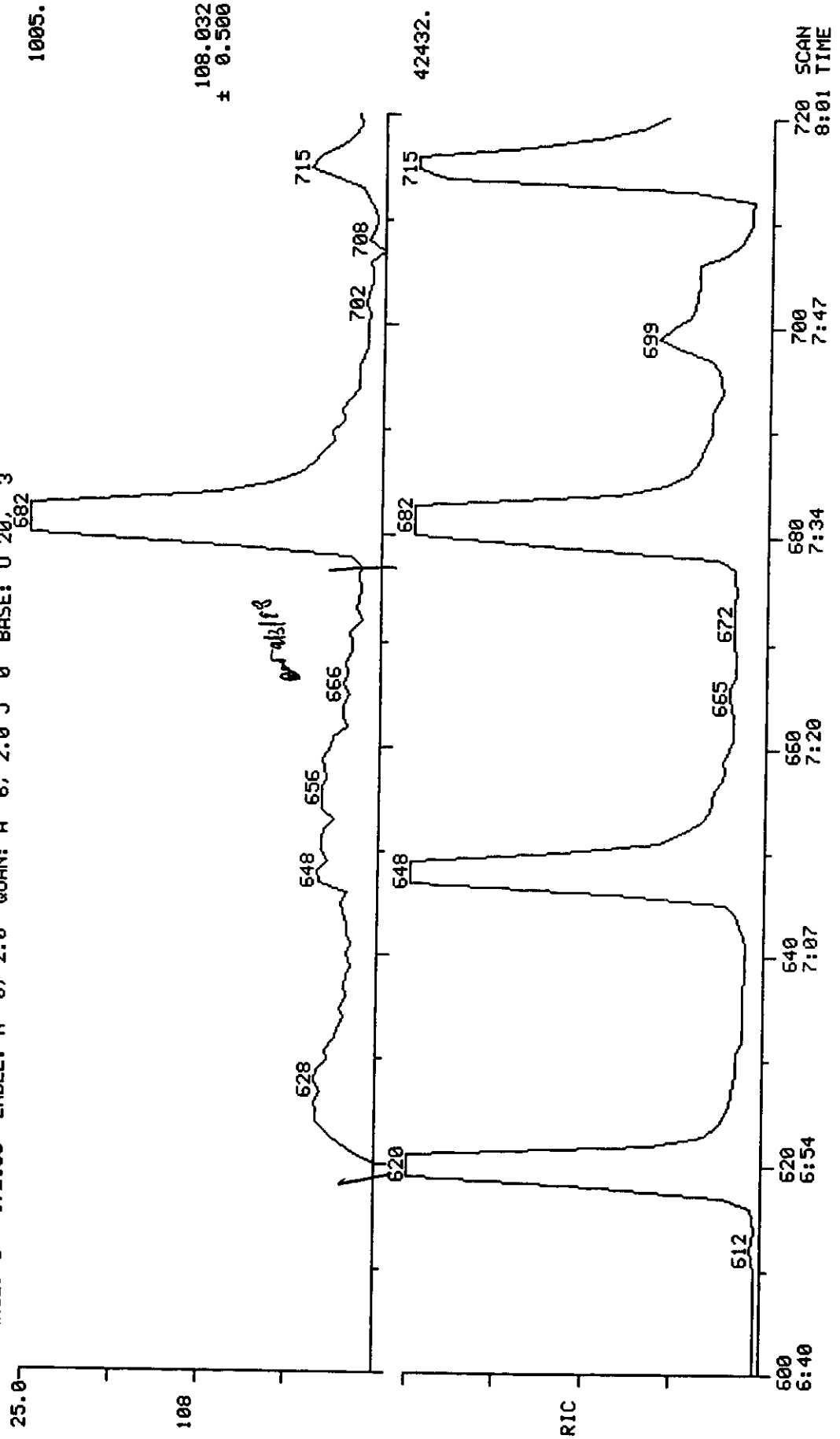
RIC+MASS CHROMATOGRAM  
 09/02/98 15:24:00  
 SAMPLE: 80UG/ML AP9 052598F  
 CONDS.: INST. ID: F16  
 RANGE: G 1.2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3  
 DATA: ST16980902A #1  
 CALI: ST16980902A #3  
 SCANS 480 TO 820

3948.



SCAN TIME

RIC+MASS CHROMATOGRAM  
 09/02/98 15:24:00 DATA: ST16980902A #1 SCANS 600 TO 720  
 SAMPLE: 80UG/ML AP9 062598F CALI: ST16980902A #3  
 CONDS.: INST. ID: F16  
 RANGE: G 1,2158 LABEL: N 0, 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3





Method Blank (w/SCS)

METHOD BLANK REPORT  
 Semivolatile Organics by GC/MS  
 Project: 300681

Test: 8270-TCL-G  
 Method: 0010/8270  
 Matrix: FILTER  
 QC Lot: 31 JUL 98-16A  
 Analyzed: 31 AUG 98

Semivolatile Organics

QC Run: 31 JUL 98-16A  
 Time: 16:02

Analyte	Result	Units	Reporting Limit	Qualifier
Phenol	ND	ug/Sample	10	
bis(2-Chloroethyl)ether	ND	ug/Sample	10	
2-Chlorophenol	ND	ug/Sample	10	
1,3-Dichlorobenzene	ND	ug/Sample	10	
1,4-Dichlorobenzene	ND	ug/Sample	10	
Benzyl alcohol	ND	ug/Sample	10	
1,2-Dichlorobenzene	ND	ug/Sample	10	
2-Methylphenol	ND	ug/Sample	10	
2,2'-Oxybis(1-chloropropane)	ND	ug/Sample	10	
3/4-Methylphenol	ND	ug/Sample	10	
N-Nitroso-di-n-propylamine	ND	ug/Sample	10	
Hexachloroethane	ND	ug/Sample	10	
Nitrobenzene	ND	ug/Sample	10	
Isophorone	ND	ug/Sample	10	
2-Nitrophenol	ND	ug/Sample	10	
2,4-Dimethylphenol	ND	ug/Sample	10	
Benzoic acid	ND	ug/Sample	50	
bis(2-Chloroethoxy)-methane	ND	ug/Sample	10	
2,4-Dichlorophenol	ND	ug/Sample	10	
1,2,4-Trichlorobenzene	ND	ug/Sample	10	
Naphthalene	ND	ug/Sample	10	
4-Chloroaniline	ND	ug/Sample	10	
Hexachlorobutadiene	ND	ug/Sample	10	
4-Chloro-3-methylphenol	ND	ug/Sample	10	
2-Methylnaphthalene	ND	ug/Sample	10	
Hexachlorocyclopentadiene	ND	ug/Sample	10	
2,4,6-Trichlorophenol	ND	ug/Sample	10	
2,4,5-Trichlorophenol	ND	ug/Sample	50	
2-Chloronaphthalene	ND	ug/Sample	10	
2-Nitroaniline	ND	ug/Sample	10	
Dimethyl phthalate	ND	ug/Sample	10	
Acenaphthylene	ND	ug/Sample	10	
3-Nitroaniline	ND	ug/Sample	50	
Acenaphthene	ND	ug/Sample	10	
2,4-Dinitrophenol	ND	ug/Sample	50	
4-Nitrophenol	ND	ug/Sample	50	
Dibenzofuran	ND	ug/Sample	10	
2,4-Dinitrotoluene	ND	ug/Sample	10	
2,6-Dinitrotoluene	ND	ug/Sample	10	
Diethyl phthalate	ND	ug/Sample	10	
4-Chlorophenyl phenyl ether	ND	ug/Sample	10	
Fluorene	ND	ug/Sample	10	
4-Nitroaniline	ND	ug/Sample	50	
4,6-Dinitro-2-methylphenol	ND	ug/Sample	50	
N-Nitrosodiphenylamine	ND	ug/Sample	10	

ND = Not Detected

METHOD BLANK REPORT (cont.)  
 Semivolatile Organics by GC/MS  
 Project: 300681

Test: 8270-TCL-G  
 Method: 0010/8270  
 Matrix: FILTER  
 QC Lot: 31 JUL 98-16A  
 Analyzed: 31 AUG 98

Semivolatile Organics

QC Run: 31 JUL 98-16A  
 Time: 16:02

Analyte	Result	Units	Reporting Limit	Qualifier
4-Bromophenyl phenyl ether	ND	ug/Sample	10	
Hexachlorobenzene	ND	ug/Sample	10	
Pentachlorophenol	ND	ug/Sample	50	
Phenanthrene	ND	ug/Sample	10	
Anthracene	ND	ug/Sample	10	
Di-n-butyl phthalate	ND	ug/Sample	10	
Fluoranthene	ND	ug/Sample	10	
Pyrene	ND	ug/Sample	10	
Butyl benzyl phthalate	ND	ug/Sample	10	
3,3'-Dichlorobenzidine	ND	ug/Sample	20	
Benzo(a)anthracene	ND	ug/Sample	10	
bis(2-Ethylhexyl)-phthalate	ND	ug/Sample	10	
Chrysene	ND	ug/Sample	10	
Di-n-octyl phthalate	ND	ug/Sample	10	
Benzo(b)fluoranthene	ND	ug/Sample	10	
Benzo(k)fluoranthene	ND	ug/Sample	10	
Benzo(a)pyrene	ND	ug/Sample	10	
Indeno(1,2,3-cd)pyrene	ND	ug/Sample	10	
Dibenz(a,h)anthracene	ND	ug/Sample	10	
Benzo(g,h,i)perylene	ND	ug/Sample	10	
Acetophenone	ND	ug/Sample	10	
4-Aminobiphenyl	ND	ug/Sample	50	
Aniline	ND	ug/Sample	10	
Benzidine	ND	ug/Sample	100	
3,3'-Dimethylbenzidine	ND	ug/Sample	20	
N-Nitrosodimethylamine	ND	ug/Sample	10	
N-Nitrosomorpholine	ND	ug/Sample	10	
Pentachloronitrobenzene (PCNB)	ND	ug/Sample	50	
o-Toluidine	ND	ug/Sample	20	
2-Methoxybenzenamine	ND	ug/Sample	--	
Biphenyl	ND	ug/Sample	--	
Chloroacetophenone	ND	ug/Sample	--	
Cumene	ND	ug/Sample	--	
DBCP (1,2-Dibromo-3-chloropropane)	ND	ug/Sample	--	
Benzo(e)pyrene	ND	ug/Sample	--	
N-N-Diethylaniline	ND	ug/Sample	--	
Dimethylaniline	ND	ug/Sample	--	
3,3'-Dimethoxybenzidine	ND	ug/Sample	--	
Hydroquinone	ND	ug/Sample	--	
4,4'-Methyl-bis(2-chloroaniline)	ND	ug/Sample	--	
4-Nitrodiphenyl	ND	ug/Sample	--	
Trifluralin	ND	ug/Sample	--	

ND = Not Detected



METHOD BLANK REPORT (cont.)  
Semivolatile Organics by GC/MS  
Project: 300681

Surrogate	% Recovery	Acceptable Range
Nitrobenzene-d5	86	45 -107
2-Fluorobiphenyl	102	62 -110
Terphenyl-d14	106	58 -135
Phenol-d5	85	43 -130
2-Fluorophenol	82	36 -111
2,4,6-Tribromophenol	94	58 -131

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSX

Std Id: ST16980831

Sample: SBLKSX 31JUL98-16A 1/3SA/1ML

INST. ID: F16

Client: PACIFI

Date Analyzed: 08/31/98 16:02

Analyst: DAT Instrument ID: F16

Run Factor: 3.00

Quan List Threshold: 0.95

Surrogate Vol.: 1.00

Surrogate Spike Recoveries

8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	43.20	86.4	45 107
CS25 2-FLUOROBIPHENY	50.00	51.20	102.	62 110
CS30 TERPHENYL-D14	50.00	52.90	106.	58 135
CS45 PHENOL-D5	100.0	84.60	84.6	43 130
CS50 2-FLUOROPHENOL	100.0	81.80	81.8	36 111
CS55 2,4,6-TRIBROMOP	100.0	94.50	94.5	58 131

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting
				Limit
HEXACHLOROETHANE		ND	UG/A	30.0
C310 N-NITROSODIMETHYLAMINE		ND	UG/A	30.0
PYRIDINE		ND	UG/A	60.0
2-PICOLINE		ND	UG/A	30.0
N-NITROSOMETHYLETHYLAMINE		ND	UG/A	30.0
METHYLMETHANESULFONATE		ND	UG/A	30.0
N-NITROSODIETHYLAMINE		ND	UG/A	30.0
ETHYLMETHANE SULFONATE		ND	UG/A	30.0
PENTACHLOROETHANE		ND	UG/A	30.0
C320 ANILINE		ND	UG/A	30.0
C315 PHENOL		ND	UG/A	30.0
C325 BIS(2-CHLOROETHYL)ETHER		ND	UG/A	30.0
C330 2-CHLOROPHENOL		ND	UG/A	30.0
C335 1,3-DICHLOROBENZENE		ND	UG/A	30.0
C340 1,4-DICHLOROBENZENE		ND	UG/A	30.0
C345 BENZYL ALCOHOL		ND	UG/A	30.0
C350 1,2-DICHLOROBENZENE		ND	UG/A	30.0
C355 2-METHYLPHENOL		ND	UG/A	30.0
C360 2,2'-OXYBIS(1-CLPROPAN)		ND	UG/A	30.0
C361 ACETOPHENONE		ND	UG/A	30.0
N-NITROSOPYRROLIDINE		ND	UG/A	30.0
N-NITROSOMORPHOLINE		ND	UG/A	30.0
3-METHYL PHENOL		ND	UG/A	30.0
C365 4-METHYLPHENOL		ND	UG/A	30.0
C370 N-NITROSO-DI-N-PROPYLAM		ND	UG/A	30.0
O-TOLUIDINE		ND	UG/A	30.0
C375 HEXACHLOROETHANE		ND	UG/A	30.0
C410 NITROBENZENE		ND	UG/A	30.0
N-NITROSOPIPERIDINE		ND	UG/A	30.0

Reviewed by: KG 9/17/98

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QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSX  
Std Id: ST16980831

Sample: SBLKSX 31JUL98-16A 1/3SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 08/31/98 16:02  
Run Factor: 3.00  
Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C415 ISOPHORONE		ND	UG/A	30.0
C420 2-NITROPHENOL		ND	UG/A	30.0
C425 2,4-DIMETHYLPHENOL		ND	UG/A	30.0
C435 BIS(2-CHLOROETHOXY)METH		ND	UG/A	30.0
C440 2,4-DICHLOROPHENOL		ND	UG/A	30.0
C430 BENZOIC ACID		ND	UG/A	150.0
AA-DIMETHYLPHENETHYLAMINE		ND	UG/A	30.0
C445 1,2,4-TRICHLOROBENZENE		ND	UG/A	30.0
C450 NAPHTHALENE		ND	UG/A	30.0
C455 4-CHLOROANILINE		ND	UG/A	30.0
2,6-DICHLOROPHENOL		ND	UG/A	30.0
HEXACHLOROPROPENE		ND	UG/A	30.0
C460 HEXACHLOROBUTADIENE		ND	UG/A	30.0
P-PHENYLENE DIAMINE		ND	UG/A	30.0
N-NITROSODI-N-BUTYLAMINE		ND	UG/A	30.0
C465 4-CHLORO-3-METHYLPHENO		ND	UG/A	30.0
SAFROLE		ND	UG/A	30.0
C470 2-METHYLNAPHTHALENE		ND	UG/A	30.0
1,2,4,5-TETRACHLOROBENZENE		ND	UG/A	30.0
ISOSAFROLE (#1)		ND	UG/A	60.0
C510 HEXACHLOROCYCLOPENTADI		ND	UG/A	30.0
C515 2,4,6-TRICHLOROPHENOL		ND	UG/A	30.0
C520 2,4,5-TRICHLOROPHENOL		ND	UG/A	30.0
ISOSAFROLE (#2)		ND	UG/A	60.0
C525 2-CHLORONAPHTHALENE		ND	UG/A	30.0
1-CHLORONAPHTHALENE		ND	UG/A	15.0
C530 2-NITROANILINE		ND	UG/A	150.0
1,4-NAPHTHOQUINONE		ND	UG/A	30.0
C535 DIMETHYLPHTHALATE		ND	UG/A	30.0
1,3-DINITROBENZENE		ND	UG/A	30.0
C540 ACENAPHTHYLENE		ND	UG/A	30.0
C543 2,6-DINITROTOLUENE		ND	UG/A	30.0
C545 3-NITROANILINE		ND	UG/A	150.0
C550 ACENAPHTHENE		ND	UG/A	30.0
C555 2,4-DINITROPHENOL		ND	UG/A	150.0
C565 DIBENZOFURAN		ND	UG/A	30.0
C560 4-NITROPHENOL		ND	UG/A	150.0
PENTACHLOROBENZENE		ND	UG/A	30.0
C570 2,4-DINITROTOLUENE		ND	UG/A	30.0
1-NAPHTHYLAMINE		ND	UG/A	30.0
2-NAPHTHYLAMINE		ND	UG/A	30.0
2,3,4,6-TETRACHLOROPHENOL		ND	UG/A	60.0
C580 DIETHYLPHTHALATE		ND	UG/A	30.0
C590 FLUORENE		ND	UG/A	30.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSX  
Std Id: ST16980831

Sample: SBLKSX 31JUL98-16A 1/3SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 08/31/98 16:02  
Run Factor: 3.00  
Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C585 4-CHLOROPHENYL-PHENYLE		ND	UG/A	30.0
5-NITRO-O-TOLUIDINE		ND	UG/A	30.0
C595 4-NITROANALINE		ND	UG/A	150.0
C610 4,6-DINITRO-2-METHYLPH		ND	UG/A	150.0
C615 N-NITROSODIPHENYLAMINE		ND	UG/A	30.0
C620 AZOBENZENE		ND	UG/A	30.0
SYM-TRINITROBENZENE		ND	UG/A	30.0
C625 4-BROMOPHENYL-PHENYLET		ND	UG/A	30.0
PHENACETIN		ND	UG/A	30.0
DIALLATE	234	ND	UG/A	30.0
C630 HEXACHLOROENZENE		ND	UG/A	30.0
4-AMINOBIIPHENYL		ND	UG/A	30.0
C635 PENTACHLOROPHENOL		ND	UG/A	150.0
PRONAMIDE		ND	UG/A	30.0
PENTACHLORONITROBENZENE		ND	UG/A	150.0
C640 PHENANTHRENE		ND	UG/A	30.0
C645 ANTHRACENE		ND	UG/A	30.0
2SECBUTYL-4,6-DINITROPHENOL		ND	UG/A	30.0
C647 CARBAZOLE		ND	UG/A	30.0
C650 DI-N-BUTYLPHTHALATE		ND	UG/A	30.0
4-NITROQUINOLINE-1-OXIDE		ND	UG/A	30.0
METHAPYRILENE		ND	UG/A	30.0
ISODRIN		ND	UG/A	30.0
C655 FLUORANTHENE		ND	UG/A	30.0
CHLOROBENZILATE		ND	UG/A	30.0
C710 BENZIDINE		ND	UG/A	300.0
C715 PYRENE		ND	UG/A	30.0
ARAMITE (#1)		ND	UG/A	30.0
ARAMITE (#2)		ND	UG/A	30.0
P-DIMETHYLAMINOAZOBENZENE		ND	UG/A	30.0
3,3'-DIMETHYLBENZIDINE		ND	UG/A	30.0
KEPONE		ND	UG/A	150.0
C720 BUTYLBENZYLPHTHALATE		ND	UG/A	30.0
2-ACETYLAMINOFLUORENE		ND	UG/A	30.0
C730 BENZO(A)ANTHRACENE		ND	UG/A	30.0
C725 3,3'-DICHLOROBENZIDINE		ND	UG/A	60.0
C740 CHRYSENE		ND	UG/A	30.0
C745 BIS(2-ETHYLHEXYL)PHTHA		ND	UG/A	30.0
3-METHYLCHOLANTHRENE		ND	UG/A	30.0
C760 DI-N-OCTYL PHTHALATE		ND	UG/A	30.0
C765 BENZO(B)FLUORANTHENE		ND	UG/A	30.0
7,12-DIMETHYLBENZANTHRACENE		ND	UG/A	30.0
C770 BENZO(K)FLUORANTHENE		ND	UG/A	30.0
HEXACHLOROPHENE		ND	UG/A	30.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSX  
Std Id: ST16980831

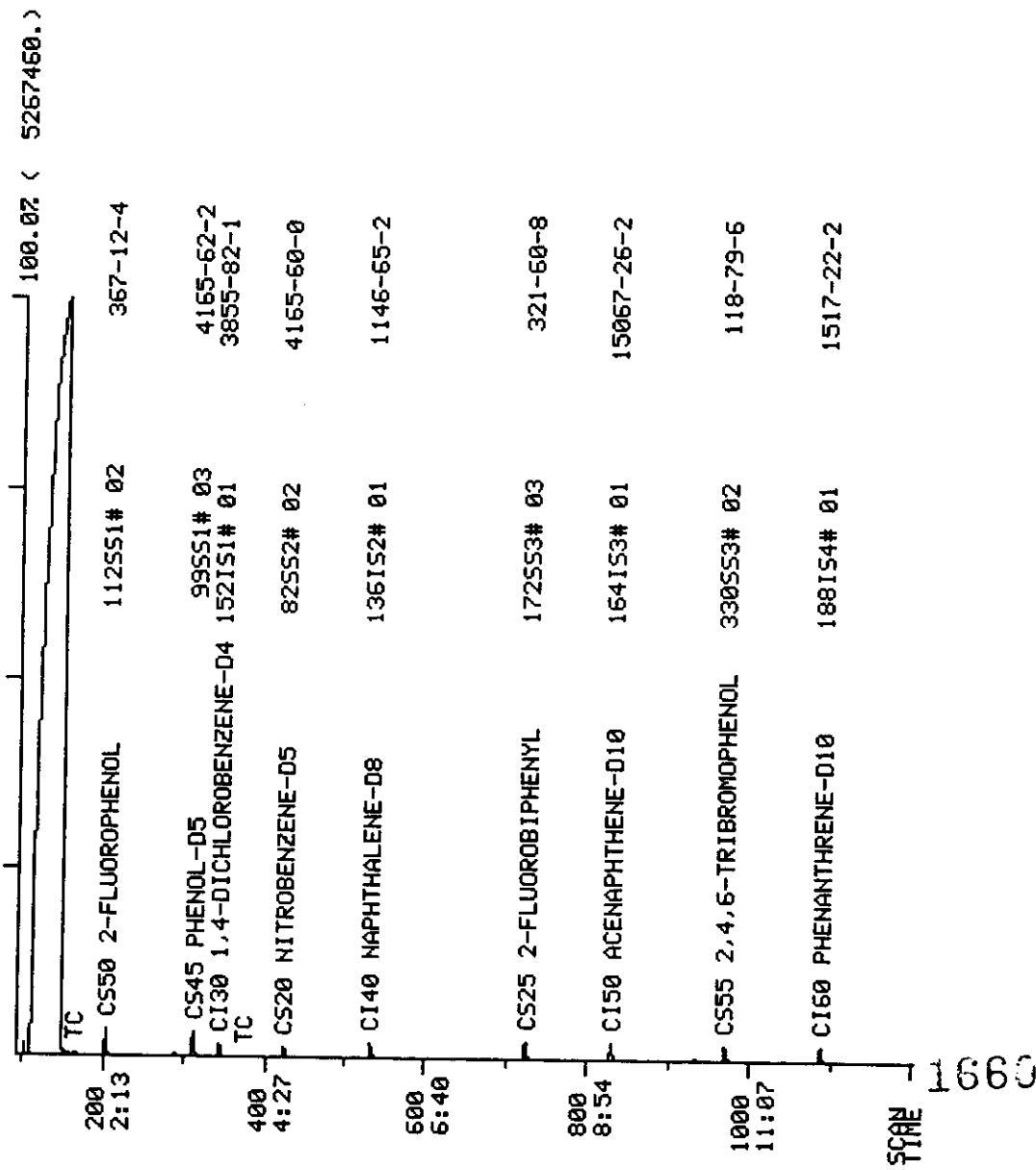
Sample: SBLKSX 31JUL98-16A 1/3SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 08/31/98 16:02  
Run Factor: 3.00  
Surrogate Vol.: 1.00

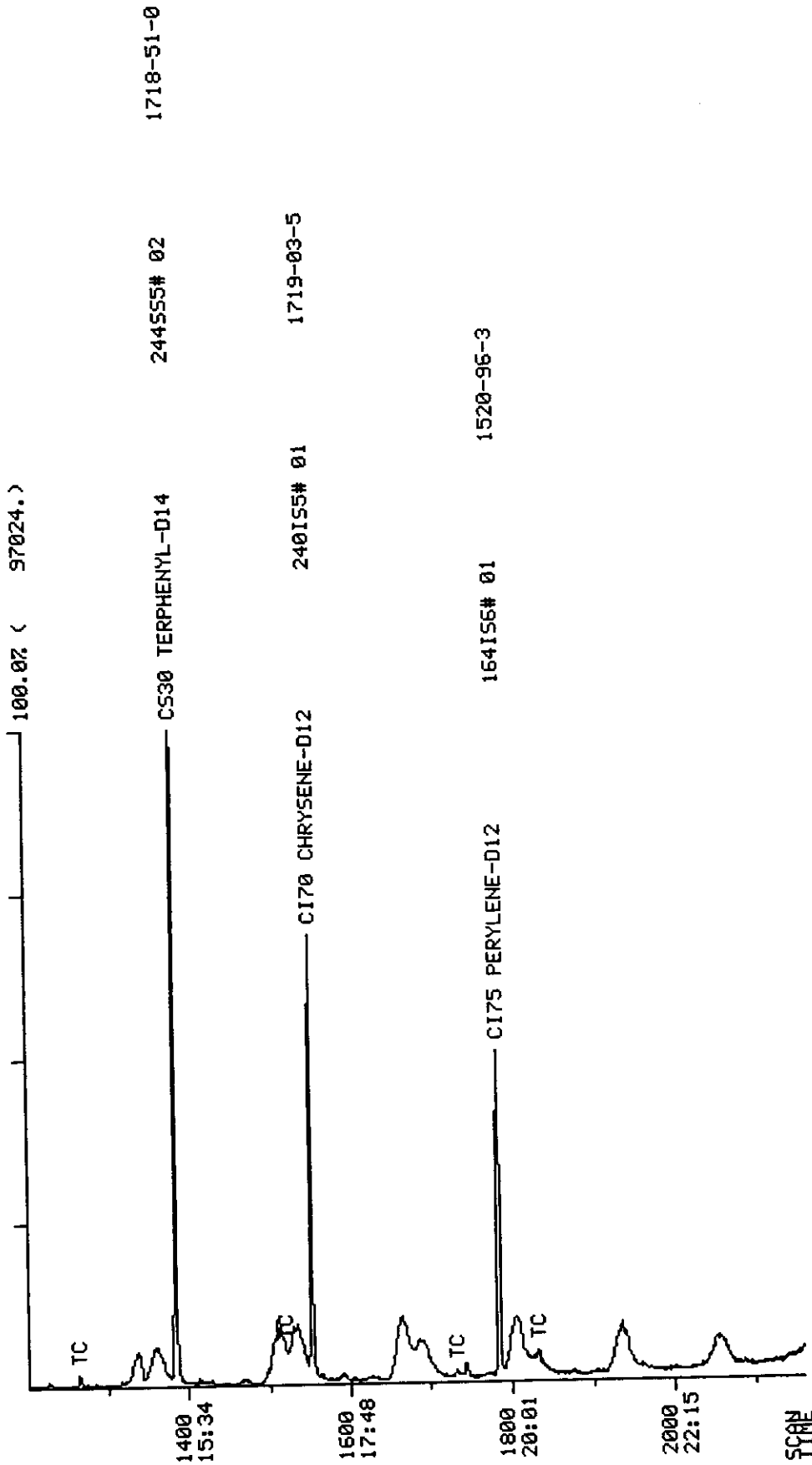
Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C775 BENZO(A)PYRENE		ND	UG/A	30.0
C780 INDENO(1,2,3-CD)PYRENE		ND	UG/A	30.0
C785 DIBENZ(A,H)ANTHRACENE		ND	UG/A	30.0
C790 BENZO(G,H,I)PERYLENE		ND	UG/A	30.0

DATA FROM FILE: 300681MBSX SCANS 93 TO 1203 ACQUIRED: 08/31/98 16:02:00  
 SAMPLE: SBLK5X 31JUL98-16A 1/35A/1ML CALI: 300681MBSX #3  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA) >1/35A NA M INST. ID: F16



DATA FROM FILE: 300681MBSX SCANS 1203 TO 2158 ACQUIRED: 08/31/98 16:02:00  
 CALI: 300681MBSX #3  
 SAMPLE: SBLKSX 31JUL98-16A 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M



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*[Handwritten signature]*

ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBSX

Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 16:02

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML							
No	Entry	Name	Mass	Meth	Scan	Ref	Area	RRF(L)	Amount
1	S1# 1	CI30 1,4-DICHLOROBENZENE	152	A	BB	342 1	36292.	1.000	40.000
2	S2# 1	CI40 NAPHTHALENE-D8	136	A	BB	530 2	131128.	1.000	40.000
3	S3# 1	CI50 ACENAPHTHENE-D10	164	A	BB	828 3	72270.	1.000	40.000
4	S4# 1	CI60 PHENANTHRENE-D10	188	A	BB	1086 4	123798.	1.000	40.000
5	S5# 1	CI70 CHRYSENE-D12	240	A	BB	1551 5	64001.	1.000	40.000
6	S6# 1	CI75 PERYLENE-D12	264	A	BB	1782 6	47432.	1.000	40.000
7	S2# 2	CS20 NITROBENZENE-D5	82	A	BB	422 2	66532.	0.470	43.222
8	S3# 3	CS25 2-FLUOROBIPHENYL	172	A	BB	723 3	114183.	1.234	51.202
9	S5# 2	CS30 TERPHENYL-D14	244	A	BB	1383 5	84273.	0.995	52.923
10	S1# 3	CS45 PHENOL-D5	99	A	BB	309 1	155502.	2.025	84.617
11	S1# 2	CS50 2-FLUOROPHENOL	112	A	BB	201 1	101569.	1.368	81.818
12	S3# 2	CS55 2,4,6-TRIBROMOPHENO	330	A	BB	968 3	40241.	0.236	94.513
13	S4# 4	HEXACHLOROBENZENE-C13	294			4 NOT FOUND			
14	S1# 15	C310 N-NITROSODIMETHYLAM	74			1 NOT FOUND			
15	S1# 20	PYRIDINE	79			1 NOT FOUND			
16	S1# 30	2-PICOLINE	93			1 NOT FOUND			
17	S1# 40	N-NITROSOMETHYLETHYLAMIN	42	A	VV	162 1	2058.	0.811	2.796
18	S1# 60	METHYLMETHANESULFONATE	80			1 NOT FOUND			
19	S1# 70	N-NITROSODIETHYLAMINE	102			1 NOT FOUND			
20	S1# 85	ETHYLMETHANE SULFONATE	79			1 NOT FOUND			
21	S1# 95	PENTACHLOROETHANE	117			1 NOT FOUND			
22	S1#100	C320 ANILINE	93			1 NOT FOUND			
23	S1#105	C315 PHENOL	94			1 NOT FOUND			
24	S1#110	C325 BIS(2-CHLOROETHYL)E	93			1 NOT FOUND			
25	S1#115	C330 2-CHLOROPHENOL	128			1 NOT FOUND			
26	S1#125	C335 1,3-DICHLOROBENZENE	146			1 NOT FOUND			
27	S1#130	C340 1,4-DICHLOROBENZENE	146			1 NOT FOUND			
28	S1#145	C345 BENZYL ALCOHOL	108	A	BB	372 1	2165.	0.854	2.793
29	S1#150	C350 1,2-DICHLOROBENZENE	146			1 NOT FOUND			
30	S1#160	C355 2-METHYLPHENOL	108			1 NOT FOUND			
31	S1#165	C360 2,2'-OXYBIS(1-CLPRO	45			1 NOT FOUND			
32	S1#170	C361 ACETOPHENONE	105			1 NOT FOUND			
33	S1#175	N-NITROSOPYRROLIDINE	100			1 NOT FOUND			
34	S1#180	N-NITROSOMORPHOLINE	56			1 NOT FOUND			
35	S1#182	3-METHYL PHENOL	108	A	BB	421 1	1034.	2.252	0.506
36	S1#185	C365 4-METHYLPHENOL	108			1 NOT FOUND			
37	S1#190	C370 N-NITROSO-D1-N-PROP	70			1 NOT FOUND			
38	S1#195	O-TOLUIDINE	106			1 NOT FOUND			
39	S1#200	C375 HEXACHLOROETHANE	117			1 NOT FOUND			
40	S2# 10	C410 NITROBENZENE	77			2 NOT FOUND			
41	S2# 15	N-NITROSOPIPERIDINE	42			2 NOT FOUND			
42	S2# 20	C415 ISOPHORONE	82			2 NOT FOUND			
43	S2# 25	C420 2-NITROPHENOL	139			2 NOT FOUND			
44	S2# 30	C425 2,4-DIMETHYLPHENOL	107			2 NOT FOUND			
45	S2# 35	C435 BIS(2-CHLOROETHOXY)	93			2 NOT FOUND			

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46 S2# 40 C440 2,4-DICHLOROPHENOL	162	2	NOT FOUND
47 S2# 45 C430 BENZOIC ACID	122	2	NOT FOUND
48 S2# 50 AA-DIMETHYLPHENETHYLAMIN	58	2	NOT FOUND
49 S2# 55 C445 1,2,4-TRICHLOROBENZ	180	2	NOT FOUND

50	S2# 60	C450 NAPHTHALENE	128	2	NOT FOUND			
51	S2# 80	C455 4-CHLOROANALINE	127	2	NOT FOUND			
52	S2# 85	2,6-DICHLOROPHENOL	162	2	NOT FOUND			
53	S2# 90	HEXACHLOROPROPENE	213	2	NOT FOUND			
54	S2# 95	C460 HEXACHLOROBUTADIENE	225	2	NOT FOUND			
55	S2#115	P-PHENYLENE DIAMINE	108	2	NOT FOUND			
56	S2#120	N-NITROSODI-N-BUTYLAMINE	84	2	NOT FOUND			
57	S2#130	C465 4-CHLORO-3-METHYLPH	107	2	NOT FOUND			
58	S2#140	SAFROLE	162	2	NOT FOUND			
59	S2#145	C470 2-METHYLNAPHTHALENE	142	2	NOT FOUND			
60	S3# 10	1,2,4,5-TETRACHLOROBENZE	214	3	NOT FOUND			
61	S3# 15	ISOSAFROLE (#1)	162	3	NOT FOUND			
62	S3# 20	C510 HEXACHLOROCYCLOPENT	237	3	NOT FOUND			
63	S3# 25	C515 2,4,6-TRICHLOROPHEN	196	3	NOT FOUND			
64	S3# 30	C520 2,4,5-TRICHLOROPHEN	196	3	NOT FOUND			
65	S3# 35	ISOSAFROLE (#2)	104	3	NOT FOUND			
66	S3# 40	C525 2-CHLORONAPHTHALENE	162	3	NOT FOUND			
67	S3# 42	1-CHLORONAPHTHALEN	162	3	NOT FOUND			
68	S3# 45	C530 2-NITROANALINE	65	3	NOT FOUND			
69	S3# 50	1,4-NAPHTHOQUINONE	158	3	NOT FOUND			
70	S3# 55	C535 DIMETHYLPHTHALATE	163	3	NOT FOUND			
71	S3# 60	1,3-DINITROBENZENE	168	3	NOT FOUND			
72	S3# 65	C540 ACENAPHTHYLENE	152	3	NOT FOUND			
73	S3# 70	C543 2,6-DINITROTOLUENE	165	3	NOT FOUND			
74	S3# 75	C545 3-NITROANILINE	138	3	NOT FOUND			
75	S3# 80	C550 ACENAPHTHENE	153	3	NOT FOUND			
76	S3# 85	C555 2,4-DINITROPHENOL	184	3	NOT FOUND			
77	S3# 90	C565 DIBENZOFURAN	168	3	NOT FOUND			
78	S3# 95	C560 4-NITROPHENOL	109	3	NOT FOUND			
79	S3#100	PENTACHLOROBENZENE	250	3	NOT FOUND			
80	S3#105	C570 2,4-DINITROTOLUENE	165	3	NOT FOUND			
81	S3#110	1-NAPHTHYLAMINE	143	3	NOT FOUND			
82	S3#115	2-NAPHTHYLAMINE	143	3	NOT FOUND			
83	S3#120	2,3,4,6-TETRACHLOROPHENO	232	3	NOT FOUND			
84	S3#130	C580 DIETHYLPHTHALATE	149	3	NOT FOUND			
85	S3#135	C590 FLUORENE	166	3	NOT FOUND			
86	S3#140	C585 4-CHLOROPHENYL-PHEN	204	3	NOT FOUND			
87	S3#145	5-NITRO-O-TOLUIDINE	152	3	NOT FOUND			
88	S3#150	C595 4-NITROANALINE	138	3	NOT FOUND			
89	S4# 10	C610 4,6-DINITRO-2-METHY	198	4	NOT FOUND			
90	S4# 15	C615 N-NITROSODIPHENYLAM	169	A BV	961 4	792.	0.551	0.465
91	S4# 20	C620 AZOBENZENE	77	4	NOT FOUND			
92	S4# 25	SYM-TRINITROBENZENE	75	4	NOT FOUND			
93	S4# 30	C625 4-BROMOPHENYL-PHENY	248	4	NOT FOUND			
94	S4# 35	PHENACETIN	108	4	NOT FOUND			
95	S4# 37	DIALLATE	234	4	NOT FOUND			
96	S4# 40	C630 HEXACHLOROBENZENE	284	4	NOT FOUND			
97	S4# 45	4-AMINOBIIPHENYL	169	4	NOT FOUND			
98	S4# 50	C635 PENTACHLOROPHENOL	266	4	NOT FOUND			
99	S4# 55	PRONAMIDE	173	4	NOT FOUND			
100	S4# 60	PENTACHLORONITROBENZENE	237	4	NOT FOUND			
101	S4# 65	C640 PHENANTHRENE	178	4	NOT FOUND			
102	S4# 70	C645 ANTHRACENE	178	4	NOT FOUND			
103	S4# 75	2SECBUTYL-4,6-DINITROPHE	211	4	NOT FOUND			
104	S4# 80	C647 CARBAZOLE	167	4	NOT FOUND			
105	S4# 85	C650 DI-N-BUTYLPHTHALATE	149	A BB	1226 4	1220.	1.247	0.316
106	S4#100	4-NITROQUINOLINE-1-OXIDE	190	4	NOT FOUND			
107	S4#105	METHAPYRILENE	58	A BB	1264 4	1726.	0.305	1.830
108	S4#106	ISODRIN	193	4	NOT FOUND			
109	S4#110	C655 FLUORANTHENE	202	4	NOT FOUND			
110	S4#120	CHLOROBENZILATE	139	A BB	1426 4	356.	0.313	0.368

111	S5# 10	C710 BENZIDINE	184	5	NOT FOUND			
112	S5# 15	C715 PYRENE	202	5	NOT FOUND			
113	S5# 20	ARAMITE (#1)	185	5	NOT FOUND			
114	S5# 25	ARAMITE (#2)	185	5	NOT FOUND			
115	S5# 30	P-DIMETHYLAMINOAZOBENZEN	120	A BB	1412	5	590.	0.419 0.880
116	S5# 35	3,3'-DIMETHYLBENZIDINE	212		5	NOT FOUND		
117	S5# 37	KEPONE	272		5	NOT FOUND		
118	S5# 40	C720 BUTYLBENZYLPHTHALAT	149		5	NOT FOUND		
119	S5# 45	2-ACETYLAMINOFLUORENE	181	A BB	1518	5	960.	0.432 1.388
120	S5# 50	C730 BENZO(A)ANTHRACENE	228		5	NOT FOUND		
121	S5# 55	C725 3,3'-DICHLOROBENZID	252	A BB	1563	5	636.	0.408 0.974
122	S5# 60	C740 CHRYSENE	228		5	NOT FOUND		
123	S5# 65	C745 BIS(2-ETHYLHEXYL)PH	149	A BB	1603	5	350.	0.980 0.223
124	S5# 85	3-METHYLCHOLANTHRENE	268	A BB	1831	5	1012.	0.548 1.154
125	S6# 10	C760 DI-N-OCTYL PHTHALAT	149		6	NOT FOUND		
126	S6# 15	C765 BENZO(B)FLUORANTHEN	252		6	NOT FOUND		
127	S6# 20	7,12-DIMETHYLBENZANTHRAC	256	A BB	1730	6	688.	0.610 0.951
128	S6# 25	C770 BENZO(K)FLUORANTHEN	252		6	NOT FOUND		
129	S6# 30	HEXACHLOROPHENE	196		6	NOT FOUND		
130	S6# 35	C775 BENZO(A)PYRENE	252		6	NOT FOUND		
131	S6# 55	C780 INDENO(1,2,3-CD)PYR	276		6	NOT FOUND		
132	S6# 60	C785 DIBENZ(A,H)ANTHRACE	278		6	NOT FOUND		
133	S6# 65	C790 BENZO(G,H,I)PERYLEN	276		6	NOT FOUND		

## QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 300681MBSX	36292.= 126		3.80	131128.= 130		5.90	72270.= 132		9.21

IS# 1 = C130 1,4-DICHLOROBENZENE-D4

IS# 2 = C140 NAPHTHALENE-D8

IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 300681MBSX	123798.= 125		12.08	64001.= 102		17.25	47432.= 93		19.82

IS# 4 = C160 PHENANTHRENE-D10

IS# 5 = C170 CHRYSENE-D12

IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

Data Reduced by: ea Date: 9/16/98  
Data Reviewed by: ✓ Date: 10/9/98

Data File: 300681MBSX

QUANTERRA GC/MS TIC REPORT ( Part 1 )

Sample: SBLKSX 31JUL98-16A 1/3SA/1ML  
Analyst: DAT

INST. ID: F16  
Date Analyzed: 08/31/98 16:02  
Run Factor: 3.00

# SCAN	Concentration in Sample (UG/A)	CAS #
1 288	29.	100-52-7
BENZALDEHYDE Or isomer		
<hr/>		
2 932	22.	<del>74381-40-1</del>
<del>PROPANOIC ACID, 2-METHYL-, 1-(1,1-DIMETHYLETHYL)-2-METHYL-, 3-PROPANE!</del>		
Unknown 5700		
<hr/>		
3 1338	19.	<del>20324-33-8</del>
<del>2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-</del>		
Unknown 5700		
<hr/>		
4 1360	28.	<del>3588-26-8</del>
<del>1-PROPANOL, 2-(2-METHOXYPROPOXY)</del>		
Unknown 5700		
<hr/>		
5 1509	27.	00-00-0
UNKNOWN		
<hr/>		
6 1533	33.	00-00-0
UNKNOWN		
<hr/>		
7 1661	33.	00-00-0
UNKNOWN		
<hr/>		
8 1690	21.	00-00-0
UNKNOWN		
<hr/>		
9 1803	51.	00-00-0
UNKNOWN		
<hr/>		
10 1933	39.	00-00-0
UNKNOWN		
<hr/>		

11/6/98

11 2052            15.            00-00-0  
UNKNOWN

---

---

12 2057            21.            00-00-0  
UNKNOWN

---

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QUANTERRA GC/MS TIC REPORT ( Part 2 )

CONCENTRATION = AREA(TIC)\*CONC(IS)/AREA(IS)

#	FIT	PURITY	INT. STD.	RT	RRT	AREA	HEIGHT	AMOUNT		LIB	LIB #
								AS ANALYZED			
								(UG/ML )			
1	967	920	1	3:12	0.543	63822.	16736.	9.581	NB	2017.	
2	928	789	2	10:21	1.126	58873.	17802.	7.266	NB	40501.	
3	848	680	4	14:52	0.863	32240.	3764.	6.454	NB	24247.	
4	871	757	4	15:07	0.877	45936.	3636.	9.195	NB	9228.	
5	808	625	4	16:47	0.973	44204.	6352.	8.849	UK	1.	
6	812	599	4	17:03	0.988	55272.	4496.	11.064	UK	1.	
7	839	463	4	18:28	1.071	54732.	5216.	10.956	UK	1.	
8	771	524	5	18:47	0.948	27952.	2224.	7.142	UK	1.	
9	651	406	5	20:03	1.012	65928.	5408.	16.845	UK	1.	
10	844	425	5	21:30	1.085	51292.	4584.	13.105	UK	1.	
11	722	310	5	22:49	1.152	20187.	3052.	5.158	UK	1.	
12	748	422	5	22:52	1.154	26821.	3087.	6.853	UK	1.	

Library Search Data: 300681MBSX # 288 Base m/z: 77  
 08/31/98 16:02:00 + 3:12 Cali: 300681MBSX # 3 RIC: 14800.  
 Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 366 matched at least 5 of the 16 largest peaks in the unknown

Rank In.	Name
1	2017 BENZALDEHYDE
2	7284 BENZOYL CHLORIDE
3	6466 ETHANONE, 2-HYDROXY-1-PHENYL-
4	2014 2,4-HEPTADIEN-6-YNAL, (E,E)-
5	9645 BENZOYLFORMIC ACID
6	10676 ETHANONE, 2-CHLORO-1-PHENYL-
7	13112 BENZOYL ISOTHIOCYANATE
8	13709 N-METHOXY-N-METHYLBENZAMIDE
9	6857 BENZENECARBOTHIOIC ACID

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H6.O	106	77	920	967	940
2	C7.H5.O.CL	140	105	817	875	830
3	C8.H8.O2	136	105	804	845	821
4	C7.H6.O	106	105	802	844	808
5	C8.H6.O3	150	105	779	850	804
6	C8.H7.O.CL	154	105	773	841	798
7	C8.H5.O.N.S	163	105	772	823	789
8	C9.H11.O2.N	165	105	763	812	793
9	C7.H6.O.S	138	77	762	847	808

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	100-52-7
2	---	---	---	---	98-88-4
3	---	---	---	---	582-24-1
4	---	---	---	---	7200-04-6
5	---	---	---	---	611-73-4
6	---	---	---	---	532-27-4
7	---	---	---	---	532-55-8
8	---	---	---	---	6919-61-5
9	---	---	---	---	98-91-9



BASE M/Z: 77  
RIC: 14800.

DATA: 300581MBSX # 288  
CALI: 300581MBSX # 3

INST. ID: F16

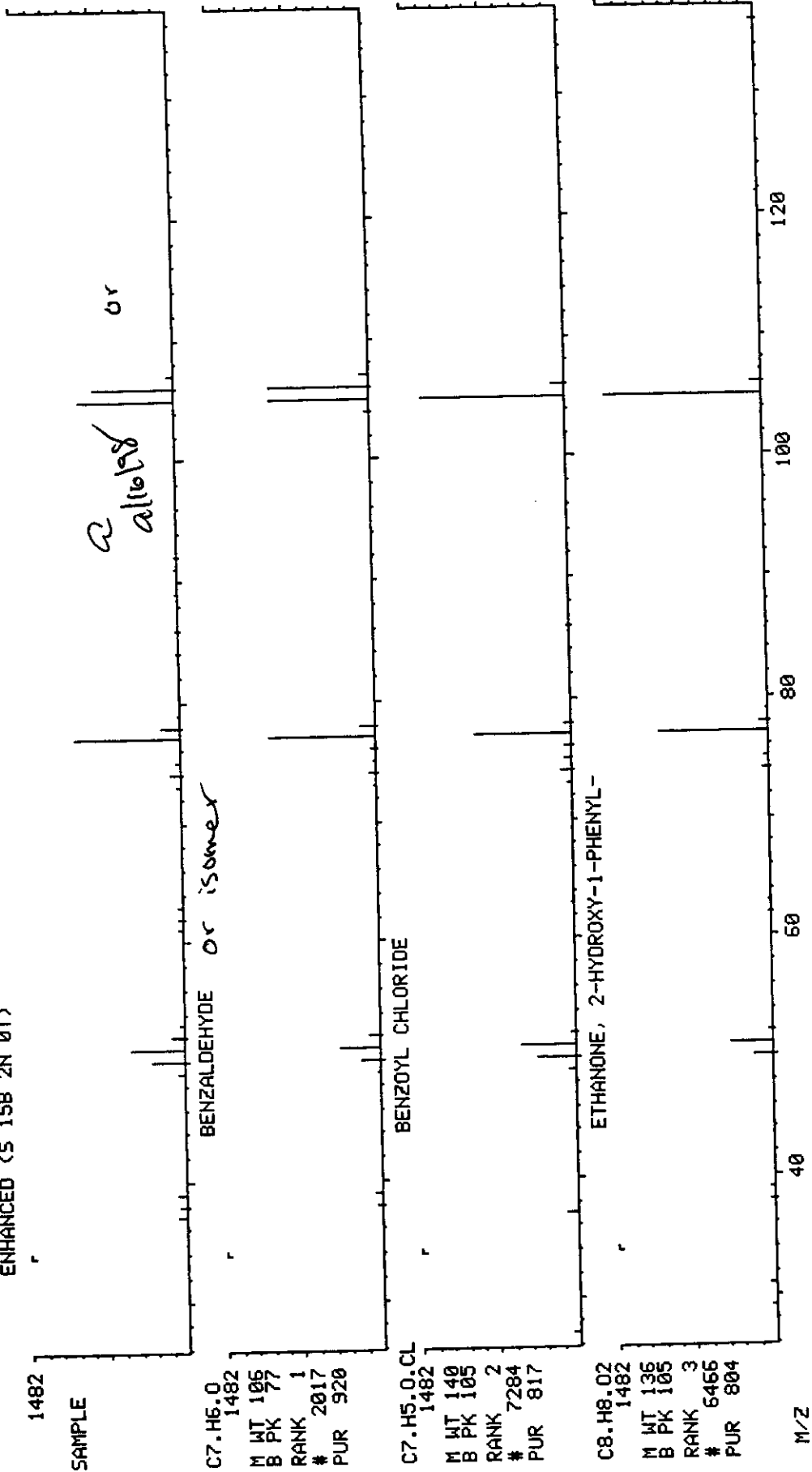
MID LIBRARY SEARCH <LIBRARYNB>

08/31/98 16:02:00 + 3:12

SAMPLE: 5BLK5X 31JUL98-16A 1/35A/1ML

CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) >1/35A NA M

ENHANCED (S 15B 2N 0T)



1671

Library Search Data: 300681MBSX # 932 Base m/z: 71  
 08/31/98 16:02:00 + 10:22 Cali: 300681MBSX # 3 RIC: 16896.  
 Sample: SBLKX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 691 matched at least 6 of the 16 largest peaks in the unknown

Rank In.	Name
1	40501 PROPANOIC ACID, 2-METHYL-, 1-(1,1-DIMETHYLETHYL)-2-METHYL-1,3-PROPA*
2	40500 PROPANOIC ACID, 2-METHYL-, 2-ETHYL-1-PROPYL-1,3-PROPANEDIYL ESTER
3	26739 PROPANOIC ACID, 2-METHYL-, 2,2-DIMETHYL-1-(2-HYDROXY-1-METHYLETHYL)*
4	4625 1-HEXENE, 3,4,5-TRIMETHYL-
5	4556 ETHANONE, 1-(3-ETHYLCYCLOBUTYL)-
6	2720 4,4-DIMETHYL-1-HEXENE
7	4677 2,4,4-TRIMETHYL-1-HEXENE
8	14787 1-NONENE, 4,6,8-TRIMETHYL-
9	15779 1,2-CYCLOHEXANEDIOL, 1-METHYL-4-(1-METHYLETHYL)-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C16.H30.O4	286	71	789	928	811
2	C16.H30.O4	286	43	727	882	737
3	C12.H24.O3	216	71	704	883	723
4	C9.H18	126	43	589	807	665
5	C8.H14.O	126	43	586	752	623
6	C8.H16	112	71	579	794	630
7	C9.H18	126	71	546	720	644
8	C12.H24	168	43	531	715	557
9	C10.H20.O2	172	71	518	701	641

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	74381-40-1
2	---	---	---	---	74367-30-9
3	---	---	---	---	74367-33-2
4	---	---	---	---	56728-10-0
5	---	---	---	---	56335-71-8
6	---	---	---	---	1647-08-1
7	---	---	---	---	51174-12-0
8	---	---	---	---	54410-98-9
9	---	---	---	---	33669-76-0

DATA: 300681MBSX # 932  
CALI: 300681MBSX # 3  
BASE M/Z: 71  
RIC: 16896.

MID LIBRARY SEARCH (LIBRARYNB)

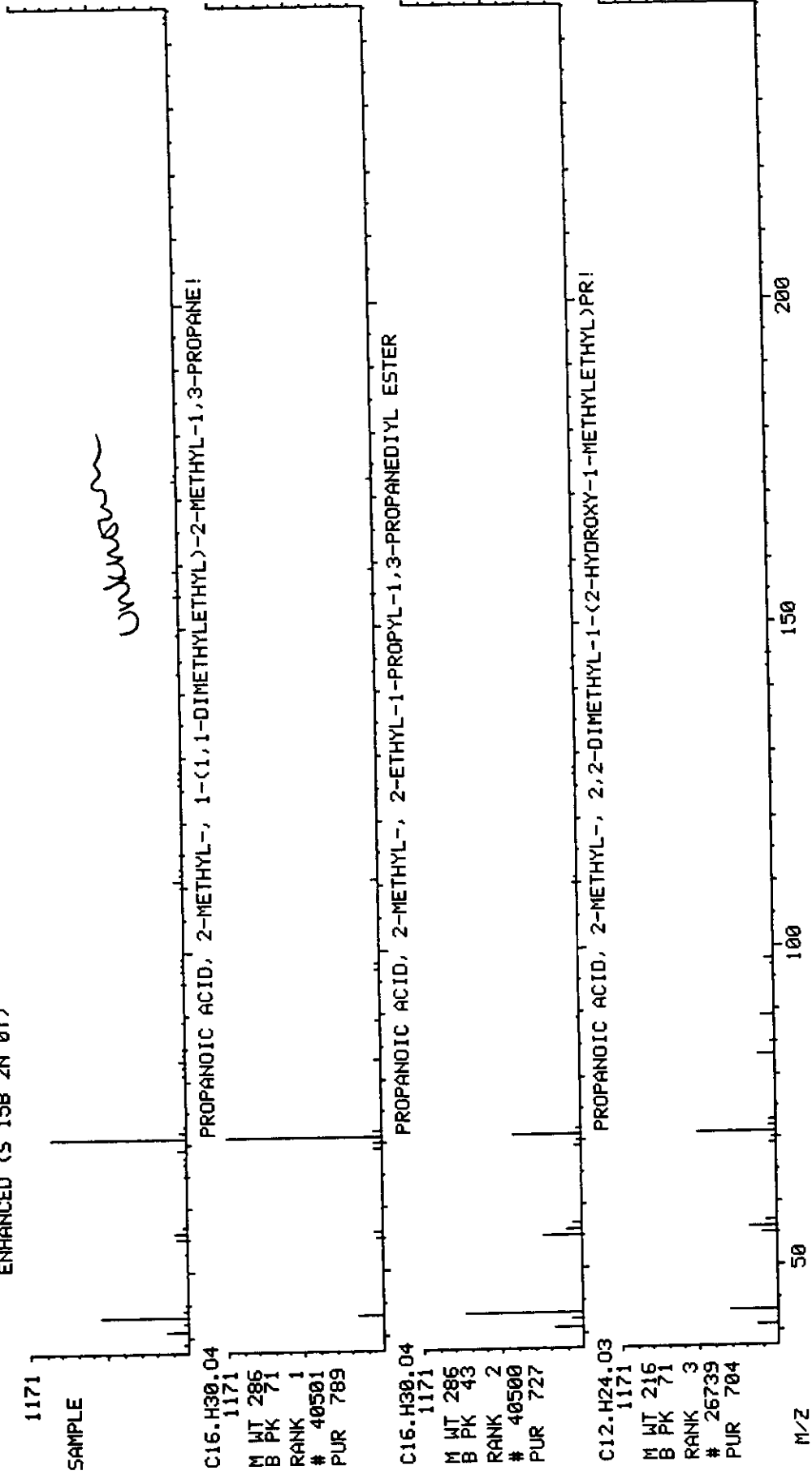
08/31/98 16:02:00 + 10:22

SAMPLE: 5BLKX 31JUL98-16A 1/35A/1ML

CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /1/35A NA M

ENHANCED (S 15B 2N 0T)

INST. ID: F16



Library Search Data: 300681MBSX #1338 Base m/z: 59  
 08/31/98 16:02:00 + 14:53 Cali: 300681MBSX # 3 RIC: 2672.  
 Sample: SBLKXS 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 842 matched at least 6 of the 16 largest peaks in the unknown

- Rank In. Name
- 1 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-
  - 2 16192 2-PROPANOL, 1-[1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-
  - 3 36537 2,5,8,11-TETRAOXATETRADECAN-13-OL, 4,7,10-TRIMETHYL-
  - 4 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-
  - 5 5522 2-HEXANOL, 2,3-DIMETHYL-
  - 6 20626 2-PROPANOL, 1,1'-[(1-METHYL-1,2-ETHANEDIYL)BIS(OXY)]BIS-
  - 7 3000 OXIRANE, 2,2-DIMETHYL-3-PROPYL-
  - 8 5527 2-HEPTANOL, 2-METHYL-
  - 9 9235 1-PROPANOL, 2-(2-METHOXY-1-METHYLETHOXY)-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H22.O4	206	59	680	848	701
2	C9.H18.O3	174	59	611	777	635
3	C13.H28.O5	264	59	609	752	704
4	C7.H16.O3	148	59	575	843	588
5	C8.H18.O	130	59	573	830	614
6	C9.H20.O4	192	59	559	777	597
7	C7.H14.O	114	43	555	811	581
8	C8.H18.O	130	59	539	890	573
9	C7.H16.O3	148	59	533	744	543

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	20324-33-8
2	---	---	---	---	55956-25-7
3	---	---	---	---	20324-34-9
4	---	---	---	---	13588-28-8
5	---	---	---	---	19550-03-9
6	---	---	---	---	1638-16-0
7	---	---	---	---	17612-35-0
8	---	---	---	---	625-25-2
9	---	---	---	---	55956-21-3

DATA: 300681MBSX #1338  
CALI: 300681MBSX # 3  
BASE M/Z: 59  
RIC: 2672.

INST. ID: F16

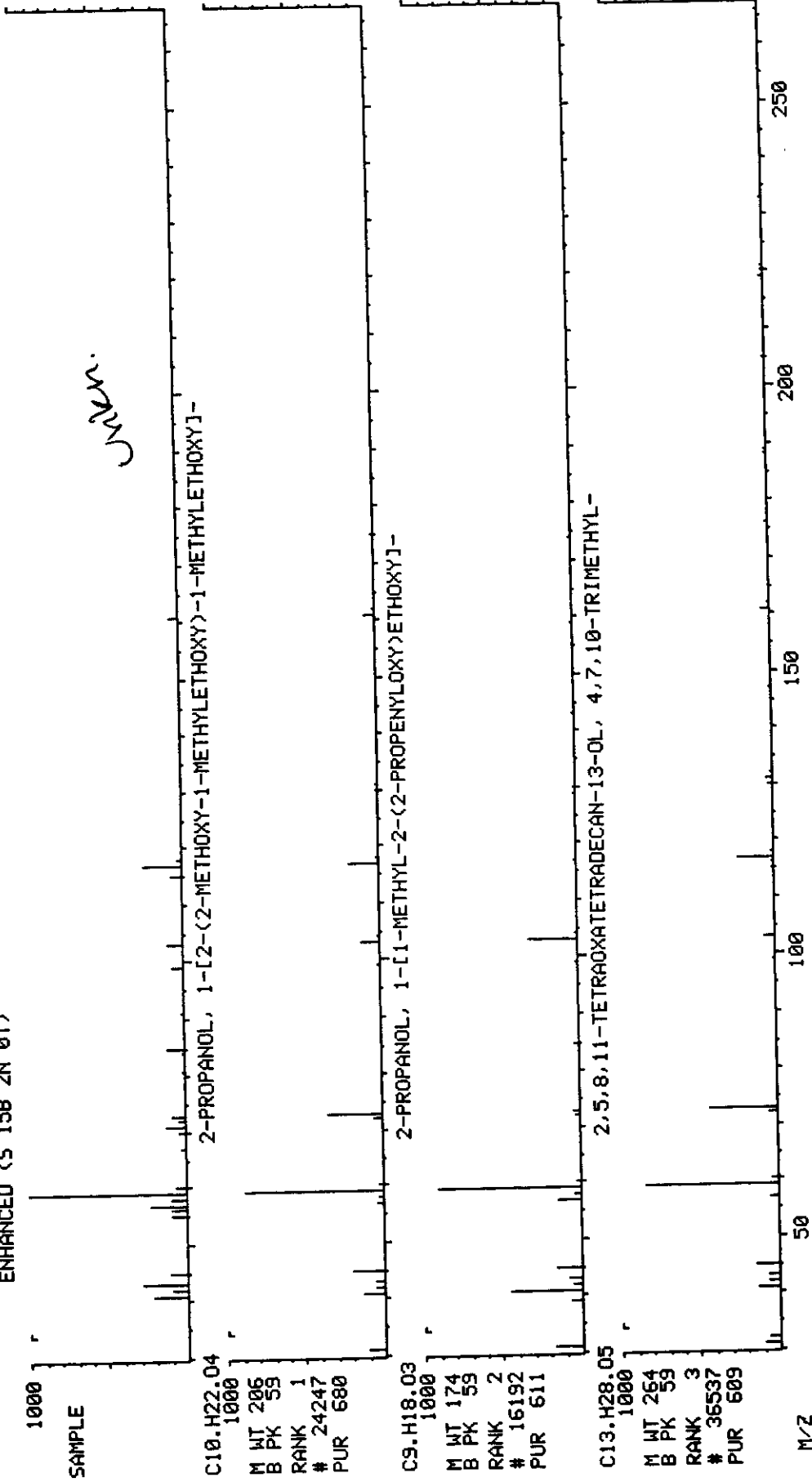
MID LIBRARY SEARCH (LIBRARYNB)

08/31/98 16:02:00 + 14:53

SAMPLE: SBLK5X 31JUL98-16A 1/35A/1ML

CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA) /1/35A NA M

ENHANCED (S 158 2N 0T)



1675

Library Search                      Data: 300681MBSX #1360    Base m/z: 59  
 08/31/98 16:02:00 + 15:08        Cali: 300681MBSX # 3    RIC: 2164.  
 Sample: SBLKSX 31JUL98-16A    1/3SA/1ML            INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 469 matched at least 6 of the 16 largest peaks in the unknown

- | Rank In. | Name   |
|----------|--|
| 1        | 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-                             |
| 2        | 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]- |
| 3        | 1775 PENTANE, 2-METHOXY-   |
| 4        | 5826 2-PROPANOL, 1-ISOPROPOXY-2-METHYL-                            |
| 5        | 1778 3-PENTANOL, 2-METHYL-   |
| 6        | 3331 HYDRAZINE, 2-BUTYL-1,1-DIMETHYL-                              |
| 7        | 915 1,2-BUTANEDIOL   |
| 8        | 1788 BUTANE, 2-METHOXY-3-METHYL-                                   |
| 9        | 3333 HYDRAZINE, 1,1-DIMETHYL-2-(1-METHYLPROPYL)-                   |

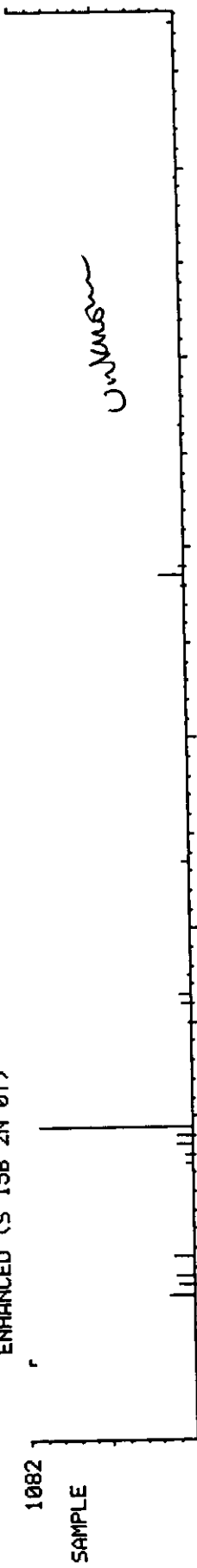
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H16.O3	148	59	757	871	777
2	C10.H22.O4	206	59	644	721	798
3	C6.H14.O	102	59	630	916	658
4	C7.H16.O2	132	59	600	743	669
5	C6.H14.O	102	59	590	851	627
6	C6.H16.N2	116	59	583	742	625
7	C4.H10.O2	90	59	565	814	660
8	C6.H14.O	102	59	555	901	595
9	C6.H16.N2	116	59	555	658	632

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	13588-28-8
2	---	---	---	---	20324-33-8
3	---	---	---	---	6795-88-6
4	---	---	---	---	3587-75-5
5	---	---	---	---	565-67-3
6	---	---	---	---	54007-23-7
7	---	---	---	---	584-03-2
8	---	---	---	---	62016-49-3
9	---	---	---	---	54007-24-8

DATA: 300681MBSX #1350  
CALI: 300681MBSX # 3  
BASE M/Z: 59  
RIC: 2154.

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 16:02:00 + 15:08  
SAMPLE: 5BLK5X 31JUL98-16A 1/35A/1ML  
CONDS.: UG/ML \*1ML \*1002/1002 \*(NA/NA )/1/35A NA M  
ENHANCED (S 15B 2N 0T)

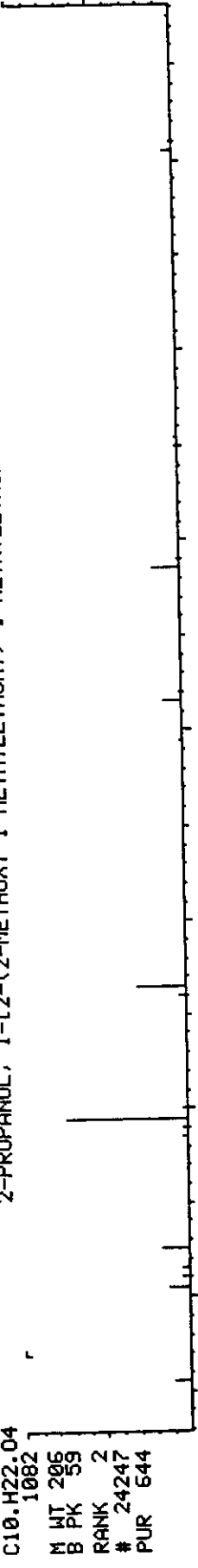
INST. ID: F16  
1-PROPANOL, 2-(2-METHOXYPROPOXY)-



C7.H16.03  
1-PROPANOL, 2-(2-METHOXYPROPOXY)-

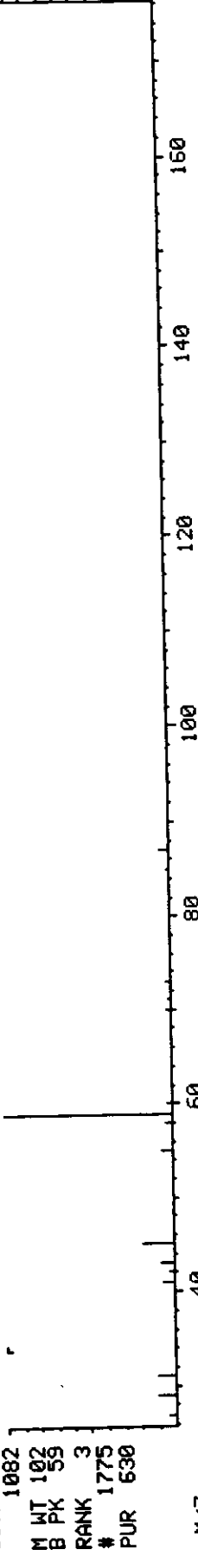
1082  
M WT 148  
B PK 59  
RANK 1  
# 9228  
PUR 757

C10.H22.04  
2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-



1082  
M WT 206  
B PK 59  
RANK 2  
# 24247  
PUR 544

C6.H14.0  
PENTANE, 2-METHOXY-



1082  
M WT 102  
B PK 59  
RANK 3  
# 1775  
PUR 530

M/Z

Library Search Data: 300681MBSX #1509 Base m/z: 59  
 08/31/98 16:02:00 + 16:47 Cali: 300681MBSX # 3 RIC: 4384.  
 Sample: SBLKX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 991 matched at least 5 of the 16 largest peaks in the unknown

- Rank In. Name
- 1 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-
  - 2 36537 2,5,8,11-TETRAOXATETRADECAN-13-OL, 4,7,10-TRIMETHYL-
  - 3 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-
  - 4 3502 BUTANOIC ACID, 3-HYDROXY-3-METHYL-
  - 5 16192 2-PROPANOL, 1-[1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-
  - 6 20626 2-PROPANOL, 1,1'-[(1-METHYL-1,2-ETHANEDIYL)BIS(OXY)]BIS-
  - 7 3522 2,3-BUTANEDIOL, 2,3-DIMETHYL-
  - 8 9235 1-PROPANOL, 2-(2-METHOXY-1-METHYLETHOXY)-
  - 9 5522 2-HEXANOL, 2,3-DIMETHYL-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H22.O4	206	59	625	808	636
2	C13.H28.O5	264	59	618	815	659
3	C7.H16.O3	148	59	552	814	564
4	C5.H10.O3	118	59	534	780	623
5	C9.H18.O3	174	59	511	717	537
6	C9.H20.O4	192	59	492	733	526
7	C6.H14.O2	118	59	483	911	498
8	C7.H16.O3	148	59	471	690	480
9	C8.H18.O	130	59	459	758	512

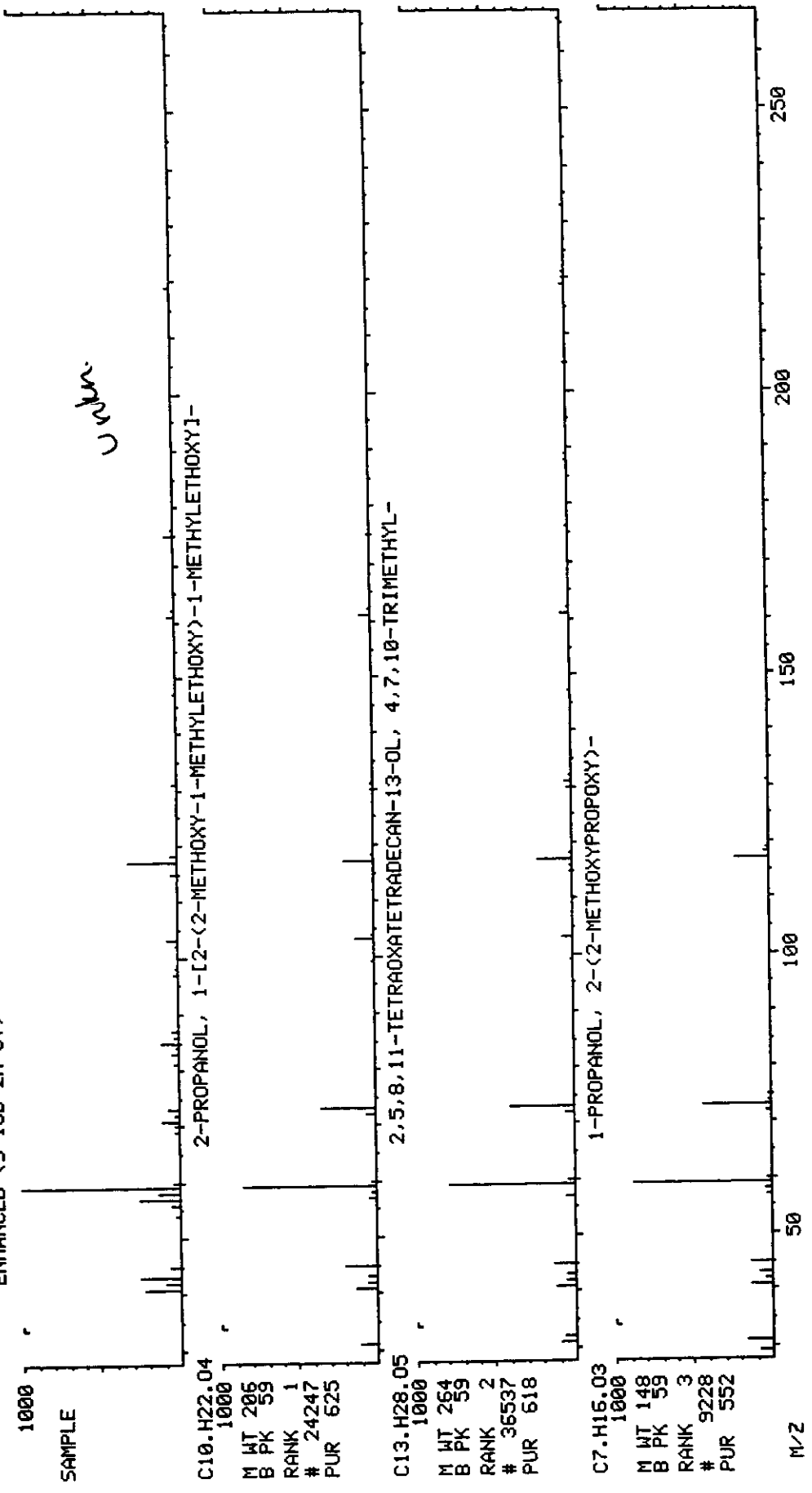
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	20324-33-8
2	---	---	---	---	20324-34-9
3	---	---	---	---	13588-28-8
4	---	---	---	---	625-08-1
5	---	---	---	---	55956-25-7
6	---	---	---	---	1638-16-0
7	---	---	---	---	76-09-5
8	---	---	---	---	55956-21-3
9	---	---	---	---	19550-03-9



MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 16:02:00 + 16:47  
 SAMPLE: SBLKX 31JUL98-16A 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) >1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSX #1509  
 CALI: 300681MBSX # 3  
 INST. ID: F16

BASE M/Z: 59  
 RIC: 4384.



Library Search Data: 300681MBSX #1533 Base m/z: 59  
 08/31/98 16:02:00 + 17:03 Cali: 300681MBSX # 3 RIC: 3384.  
 Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 427 matched at least 5 of the 16 largest peaks in the unknown

Rank	In.	Name
1	9228	1-PROPANOL, 2-(2-METHOXYPROPOXY)-
2	5826	2-PROPANOL, 1-ISOPROPOXY-2-METHYL-
3	24247	2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-
4	5506	2,3,3-TRIMETHYL-2-PENTANOL
5	334	2-PROPANOL, 2-METHYL-
6	11721	NONANAMIDE
7	1618	BUTANAMIDE, 3-METHYL-
8	3366	2-PENTANOL, 2,4-DIMETHYL-
9	329	PROPANE, 2-METHOXY-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H16.O3	148	59	599	812	663
2	C7.H16.O2	132	59	542	721	581
3	C10.H22.O4	206	59	523	673	699
4	C8.H18.O	130	59	513	844	576
5	C4.H10.O	74	59	508	936	536
6	C9.H19.O.N	157	59	492	755	559
7	C5.H11.O.N	101	59	487	804	570
8	C7.H16.O	116	59	475	863	530
9	C4.H10.O	74	59	475	899	525

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	13588-28-8
2	---	---	---	---	3587-75-5
3	---	---	---	---	20324-33-8
4	---	---	---	---	23171-85-9
5	---	---	---	---	75-65-0
6	---	---	---	---	1120-07-6
7	---	---	---	---	541-46-8
8	---	---	---	---	625-06-9
9	---	---	---	---	598-53-8

BASE M/Z: 59  
RIC: 3384.

DATA: 300681MBSX #1533  
CALI: 300681MBSX # 3  
F16

INST. ID:  
NA M

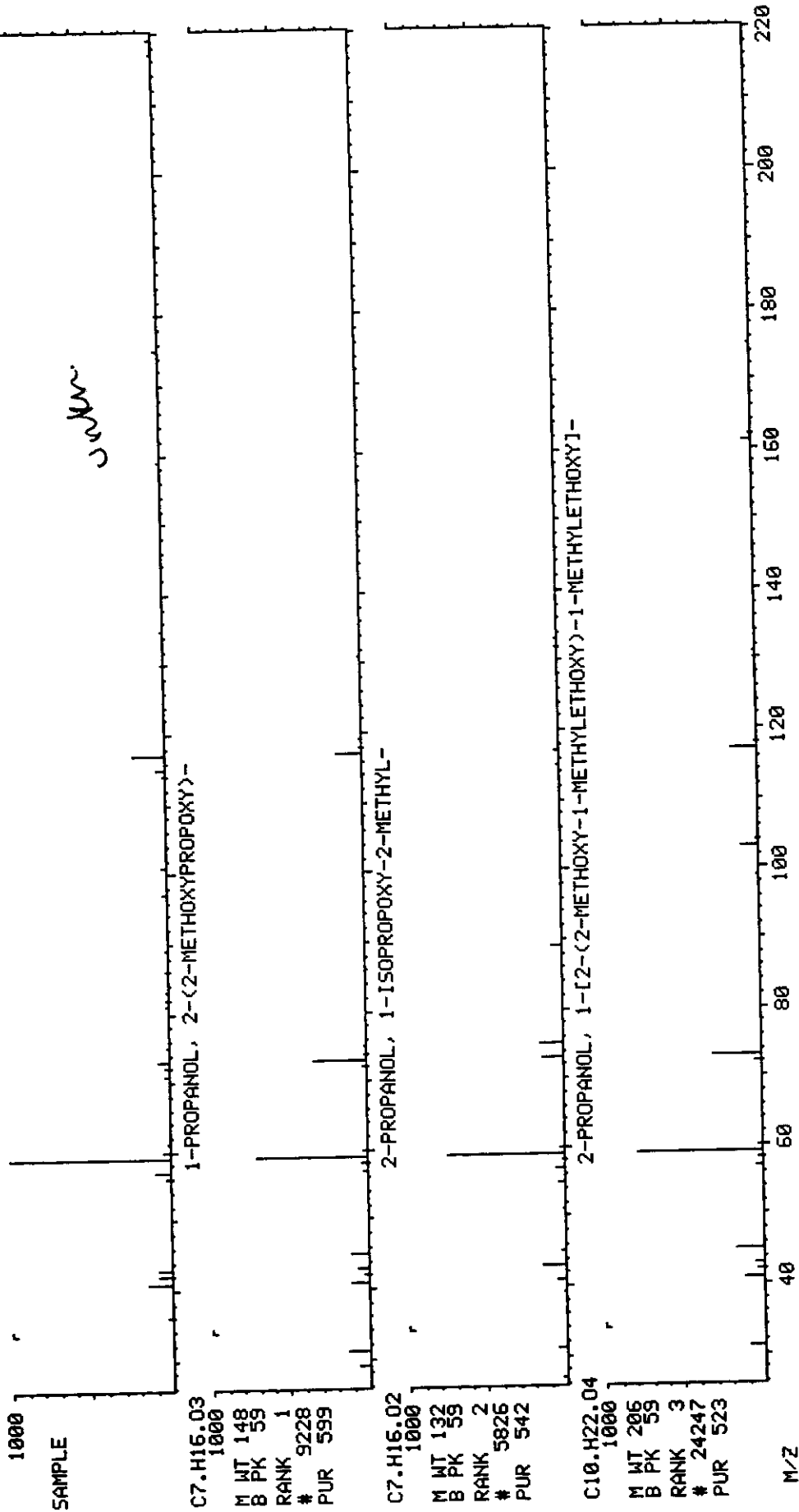
MID LIBRARY SEARCH <LIBRARYNB>

08/31/98 16:02:00 + 17:03

SAMPLE: 5BLKX 31JUL98-16A 1/35A/1ML

CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA )/1/35A

ENHANCED <S 15B 2N 0T>



SAMPLE

C7.H16.03

M WT 148  
B PK 59  
RANK 1  
# 9228  
PUR 599

C7.H16.02

M WT 132  
B PK 59  
RANK 2  
# 5826  
PUR 542

C10.H22.04

M WT 206  
B PK 59  
RANK 3  
# 24247  
PUR 523

M/Z

Library Search Data: 300681MBSX #1661 Base m/z: 59  
 08/31/98 16:02:00 + 18:29 Cali: 300681MBSX # 3 RIC: 3272.  
 Sample: SBLKX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 295 matched at least 6 of the 16 largest peaks in the unknown

- Rank In. Name
- 1 3502 BUTANOIC ACID, 3-HYDROXY-3-METHYL-
  - 2 3247 2-PENTANONE, 5-METHOXY-
  - 3 8871 METHANE, TERT-BUTOXYISOPROPOXY-
  - 4 16192 2-PROPANOL, 1-[1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-
  - 5 23273 ETHANEDIOIC ACID, BIS(1-METHYLPROPYL) ESTER
  - 6 5810 2-PENTANOL, 5-METHOXY-2-METHYL-
  - 7 9227 1,3,3-TRIMETHOXYBUTANE
  - 8 5832 2-PROPANOL, 1-(1-METHYLPROPOXY)-
  - 9 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-

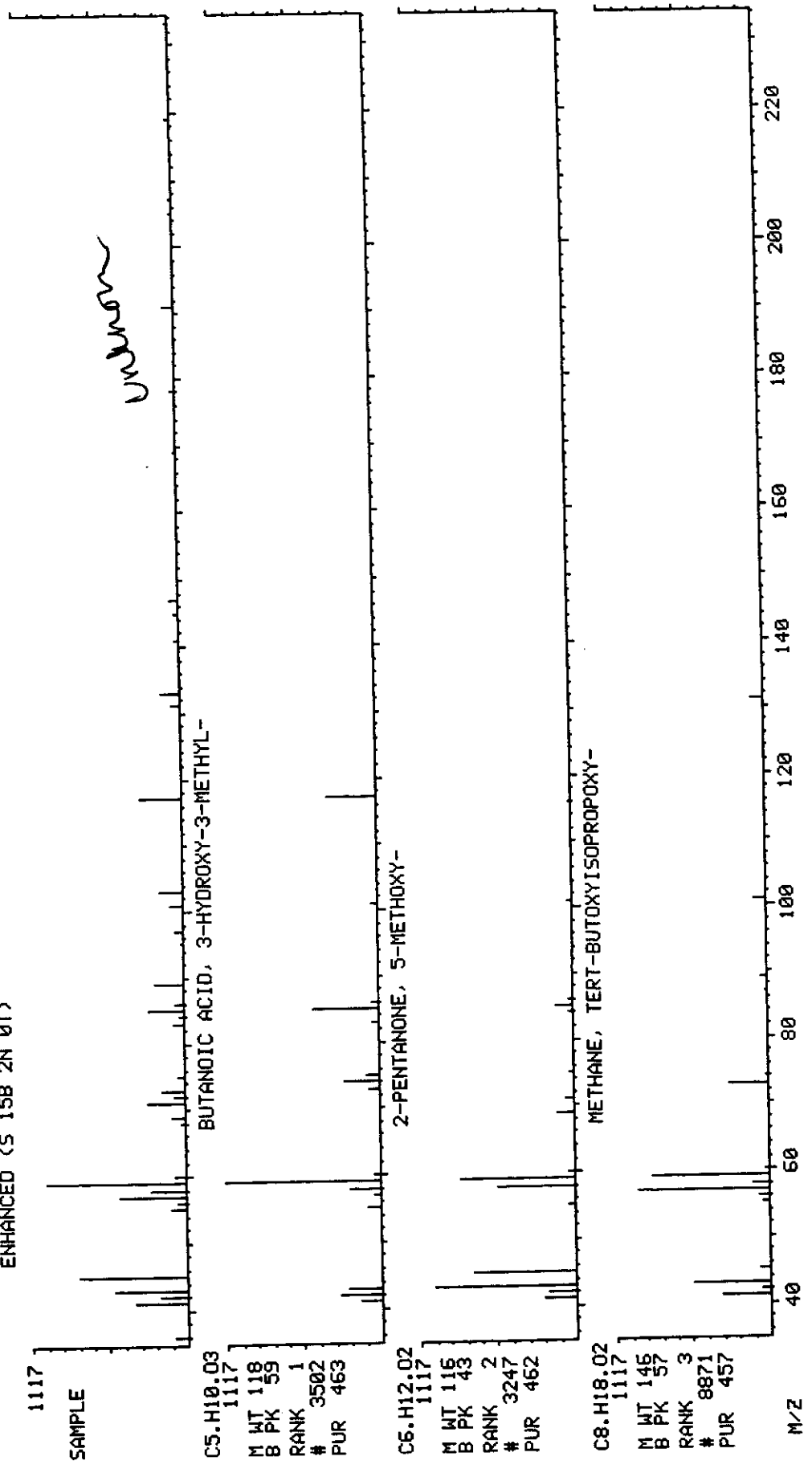
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C5.H10.O3	118	59	463	839	501
2	C6.H12.O2	116	43	462	902	468
3	C8.H18.O2	146	57	457	852	464
4	C9.H18.O3	174	59	453	738	477
5	C10.H18.O4	202	57	448	696	476
6	C7.H16.O2	132	59	447	834	478
7	C7.H16.O3	148	45	446	717	463
8	C7.H16.O2	132	43	445	748	519
9	C10.H22.O4	206	59	441	755	491

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	625-08-1
2	---	---	---	---	17429-04-8
3	---	---	---	---	4346-01-4
4	---	---	---	---	55956-25-7
5	---	---	---	---	13784-89-9
6	---	---	---	---	55724-04-4
7	---	---	---	---	6607-66-5
8	---	---	---	---	53907-95-2
9	---	---	---	---	20324-33-8

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 16:02:00 + 18:29  
 SAMPLE: 5BLK5X 31JUL98-16A 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSX #1661  
 CALI: 300681MBSX # 3  
 F15  
 INST. ID: NA M

BASE M/Z: 59  
 RIC: 3272.



1683

Library Search Data: 300681MBSX #1690 Base m/z: 59  
 08/31/98 16:02:00 + 18:48 Cali: 300681MBSX # 3 RIC: 1908.  
 Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 158 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 360 matched at least 5 of the 16 largest peaks in the unknown

Rank In.	Name
1	5432 2-BUTANONE, 3-ETHOXY-3-METHYL-
2	1775 PENTANE, 2-METHOXY-
3	20626 2-PROPANOL, 1,1'-[[1-METHYL-1,2-ETHANEDIYL]BIS(OXY)]BIS-
4	1694 3-HYDROXY-3-METHYL-2-BUTANONE
5	5527 2-HEPTANOL, 2-METHYL-
6	1760 2-PENTANOL, 2-METHYL-
7	6082 1-PROPANOL, 2-(2-HYDROXYPROPOXY)-
8	8432 2-HEXANONE, 3-HYDROXY-3,5-DIMETHYL-
9	16649 1-PROPANOL, 3-[3-(1-METHYLETHOXY)PROPOXY]-

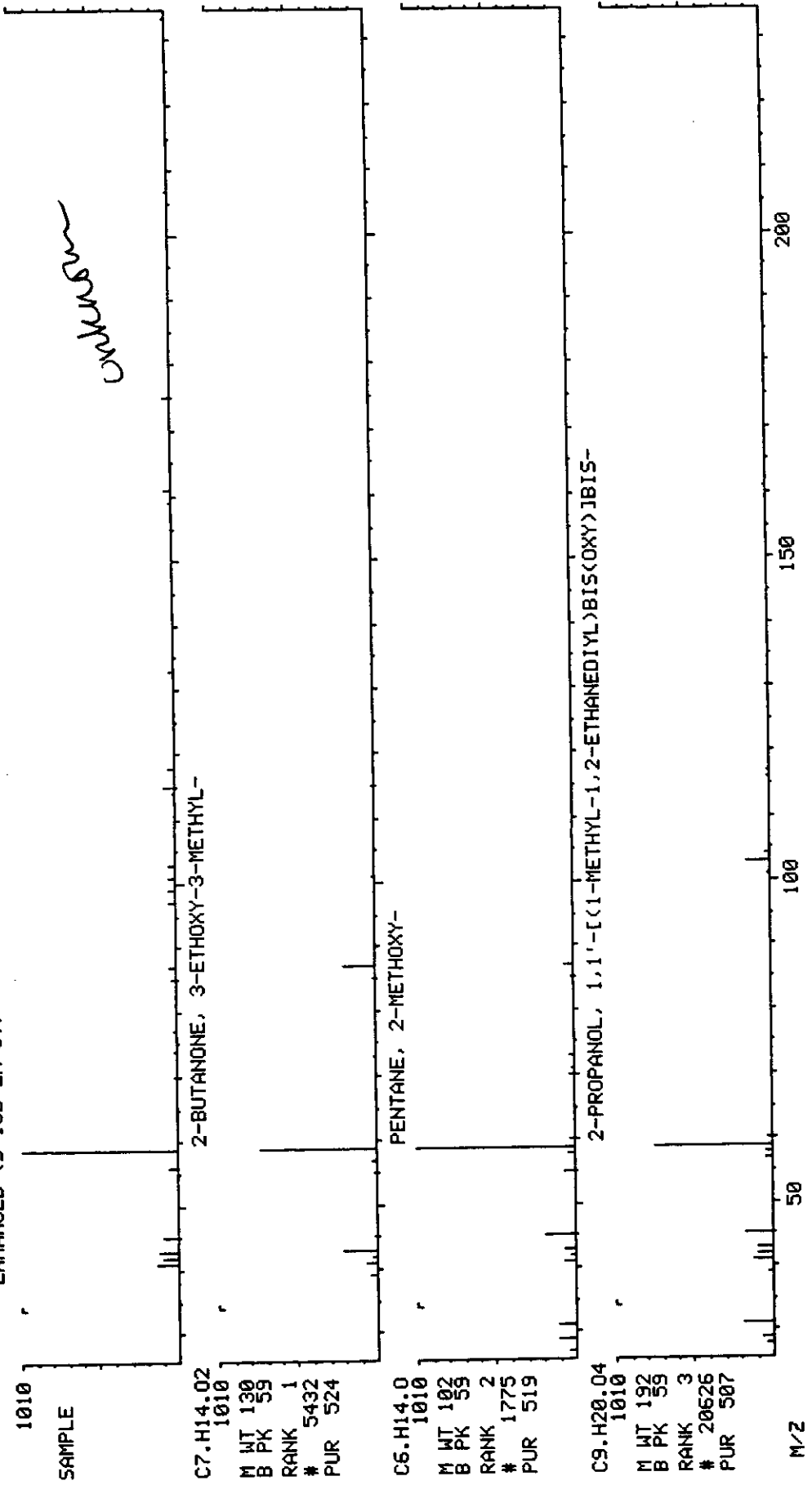
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C7.H14.O2	130	59	524	771	547
2	C6.H14.O	102	59	519	855	587
3	C9.H20.O4	192	59	507	769	587
4	C5.H10.O2	102	59	496	854	550
5	C8.H18.O	130	59	491	784	615
6	C6.H14.O	102	59	490	818	540
7	C6.H14.O3	134	59	480	782	529
8	C8.H16.O2	144	59	478	782	581
9	C9.H20.O3	176	59	477	748	552

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	36687-99-7
2	---	---	---	---	6795-88-6
3	---	---	---	---	1638-16-0
4	---	---	---	---	115-22-0
5	---	---	---	---	625-25-2
6	---	---	---	---	590-36-3
7	---	---	---	---	106-62-7
8	---	---	---	---	6321-14-8
9	---	---	---	---	54518-03-5

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 16:02:00 + 18:48  
 SAMPLE: SBLK5X 31JUL98-16A 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSX #1690  
 CALI: 300681MBSX # 3  
 INST. ID: F16

BASE M/Z: 59  
 RIC: 1908.



1000

Library Search Data: 300681MBSX #1803 Base m/z: 59  
 08/31/98 16:02:00 + 20:03 Cali: 300681MBSX # 3 RIC: 2652.  
 Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 726 matched at least 4 of the 16 largest peaks in the unknown

- | Rank In. | Name   |
|----------|--|
| 1        | 3333 HYDRAZINE, 1,1-DIMETHYL-2-(1-METHYLPROPYL)-               |
| 2        | 9228 1-PROPANOL, 2-(2-METHOXYPROPOXY)-                         |
| 3        | 5815 1-TERT-BUTOXY-2-METHOXYETHANE                             |
| 4        | 1760 2-PENTANOL, 2-METHYL-                                     |
| 5        | 12782 BIS(2-METHOXYETHYL)NITROSAMINE                           |
| 6        | 3364 ETHER, SEC-BUTYL ISOPROPYL                                |
| 7        | 3454 ETHANEDIOIC ACID, DIMETHYL ESTER                          |
| 8        | 20626 2-PROPANOL, 1,1'-[(1-METHYL-1,2-ETHANEDIYL)BIS(OXY)]BIS- |
| 9        | 5482 HYDRAZINE, 1,1-DIMETHYL-2-(1-METHYLBUTYL)-                |

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H16.N2	116	59	406	651	426
2	C7.H16.O3	148	59	358	611	517
3	C7.H16.O2	132	57	356	587	450
4	C6.H14.O	102	59	354	773	411
5	C6.H14.O3.N2	162	45	344	593	482
6	C7.H16.O	116	45	340	720	398
7	C4.H6.O4	118	59	332	897	337
8	C9.H20.O4	192	59	331	586	453
9	C7.H18.N2	130	59	330	635	453

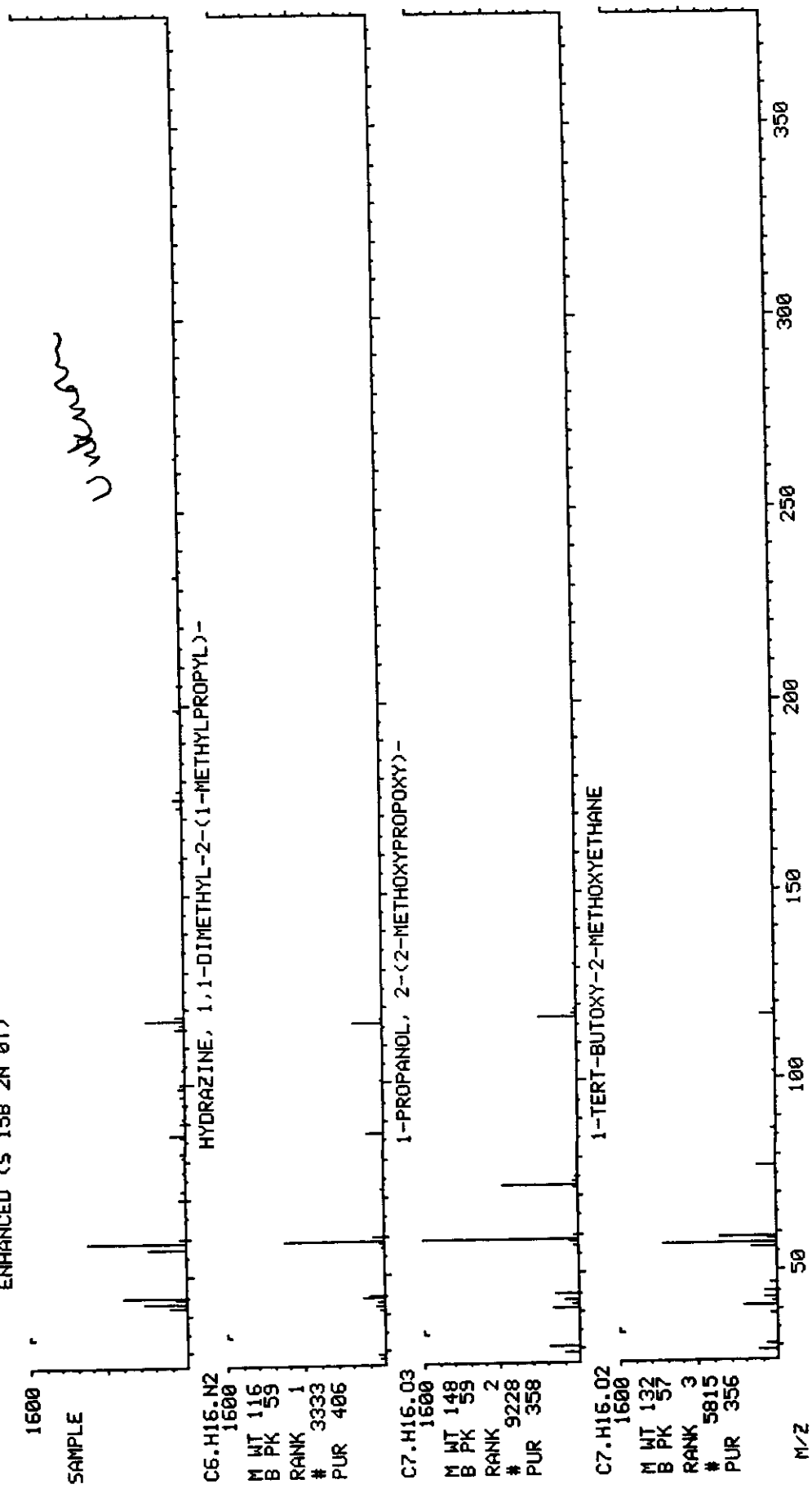
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	54007-24-8
2	---	---	---	---	13588-28-8
3	---	---	---	---	66728-50-5
4	---	---	---	---	590-36-3
5	---	---	---	---	- -
6	---	---	---	---	18641-81-1
7	---	---	---	---	553-90-2
8	---	---	---	---	1638-16-0
9	---	---	---	---	75267-97-9



MID LIBRARY SEARCH <LIBRARYNB>  
 08/31/98 16:02:00 + 20:03  
 SAMPLE: SBLK5X 31JUL98-16A 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSX #1803  
 CALI: 300681MBSX # 3  
 INST. ID: F16

BASE M/Z: 59  
 RIC: 2652.



2891

Library Search Data: 300681MBSX #1933 Base m/z: 59  
 08/31/98 16:02:00 + 21:30 Cali: 300681MBSX # 3 RIC: 3144.  
 Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 580 matched at least 5 of the 16 largest peaks in the unknown

- | Rank In. | Name   |
|----------|--|
| 1        | 3528 2-METHYL-2,3-PENTANEDIOL                                      |
| 2        | 20626 2-PROPANOL, 1,1'-[[1-METHYL-1,2-ETHANEDIYL)BIS(OXY)]BIS-     |
| 3        | 9227 1,3,3-TRIMETHOXYBUTANE  |
| 4        | 1909 2-PROPANOL, 1-METHOXY-2-METHYL-                               |
| 5        | 5809 1-ETHOXPENTAN-3-OL  |
| 6        | 3247 2-PENTANONE, 5-METHOXY-                                       |
| 7        | 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]- |
| 8        | 5810 2-PENTANOL, 5-METHOXY-2-METHYL-                               |
| 9        | 6083 2-PROPANOL, 1,1'-OXYBIS-                                      |

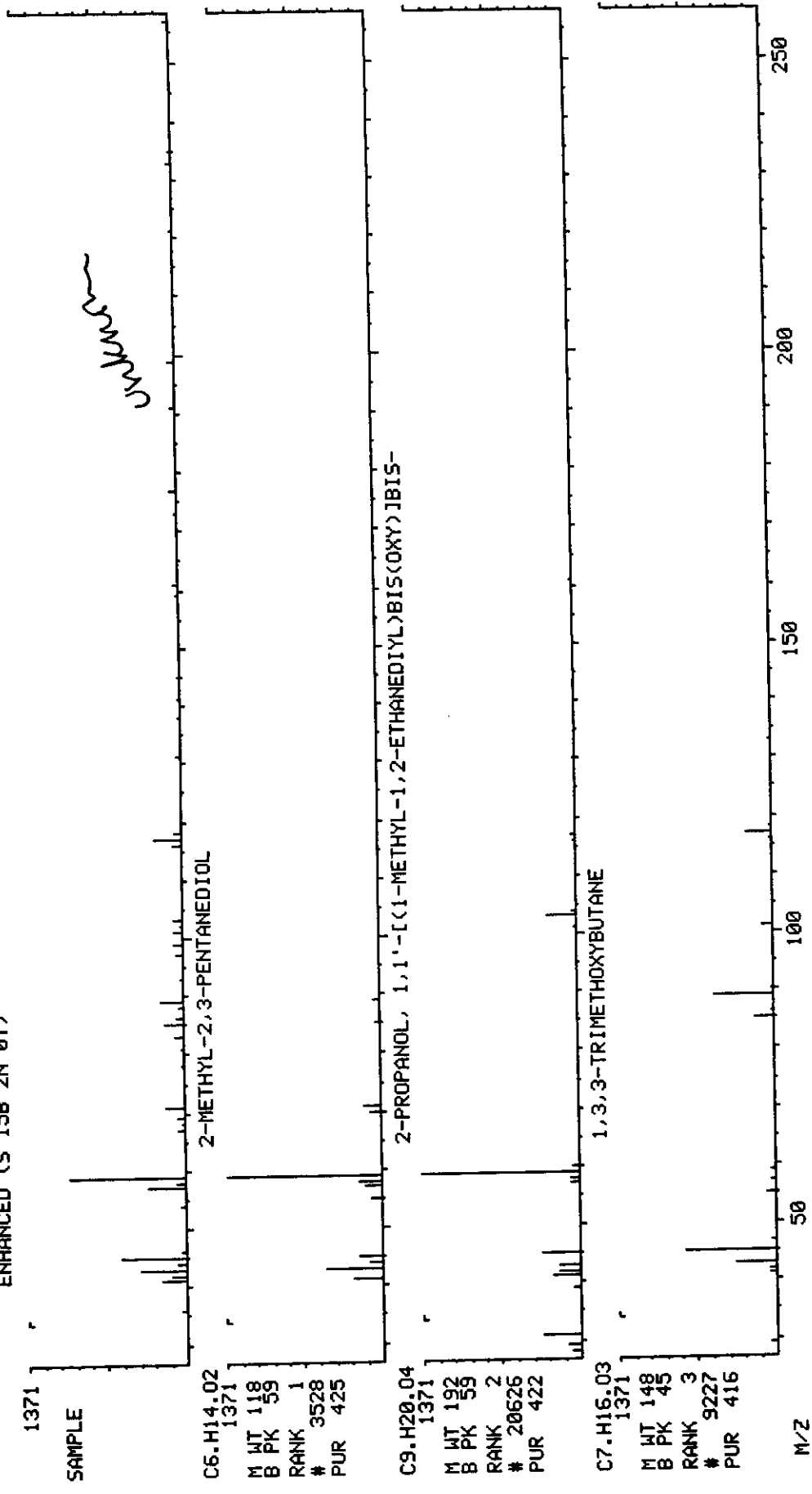
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H14.O2	118	59	425	844	455
2	C9.H20.O4	192	59	422	763	451
3	C7.H16.O3	148	45	416	675	444
4	C5.H12.O2	104	59	403	885	432
5	C7.H16.O2	132	59	401	845	446
6	C6.H12.O2	116	43	391	826	411
7	C10.H22.O4	206	59	390	695	497
8	C7.H16.O2	132	59	389	779	437
9	C6.H14.O3	134	59	386	840	412

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	7795-80-4
2	---	---	---	---	1638-16-0
3	---	---	---	---	6607-66-5
4	---	---	---	---	3587-64-2
5	---	---	---	---	- -
6	---	---	---	---	17429-04-8
7	---	---	---	---	20324-33-8
8	---	---	---	---	55724-04-4
9	---	---	---	---	110-98-5

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 16:02:00 + 21:30  
 SAMPLE: SBLK5X 31JUL98-16A 1/35A/1ML  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) >1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSX #1933  
 CALI: 300681MBSX # 3  
 INST. ID: F16

BASE M/Z: 59  
 RIC: 3144.



Library Search                      Data: 300681MBSX #2052    Base m/z: 59  
 08/31/98 16:02:00 + 22:49        Cali: 300681MBSX # 3    RIC: 1154.  
 Sample: SBLKSX 31JUL98-16A    1/3SA/1ML            INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N OT)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 159 matched at least 5 of the 16 largest peaks in the unknown

- Rank In.        Name  
 1 16192 2-PROPANOL, 1-[1-METHYL-2-(2-PROPENYLOXY)ETHOXY]-  
 2 8153 BUTANAMIDE, N-(1-OXOPROPYL)-  
 3 24247 2-PROPANOL, 1-[2-(2-METHOXY-1-METHYLETHOXY)-1-METHYLETHOXY]-  
 4 17561 GLUCOSE  
 5 8871 METHANE, TERT-BUTOXYISOPROPOXY-  
 6 12047 BUTANE, 1,1'-OXYBIS[3-METHYL-  
 7 16017 BOROXIN, TRIMETHOXY-  
 8 8519 3-HEPTANOL, 3,5-DIMETHYL-  
 9 3461 2-PROPANAMINE, N-METHYL-N-NITRO-

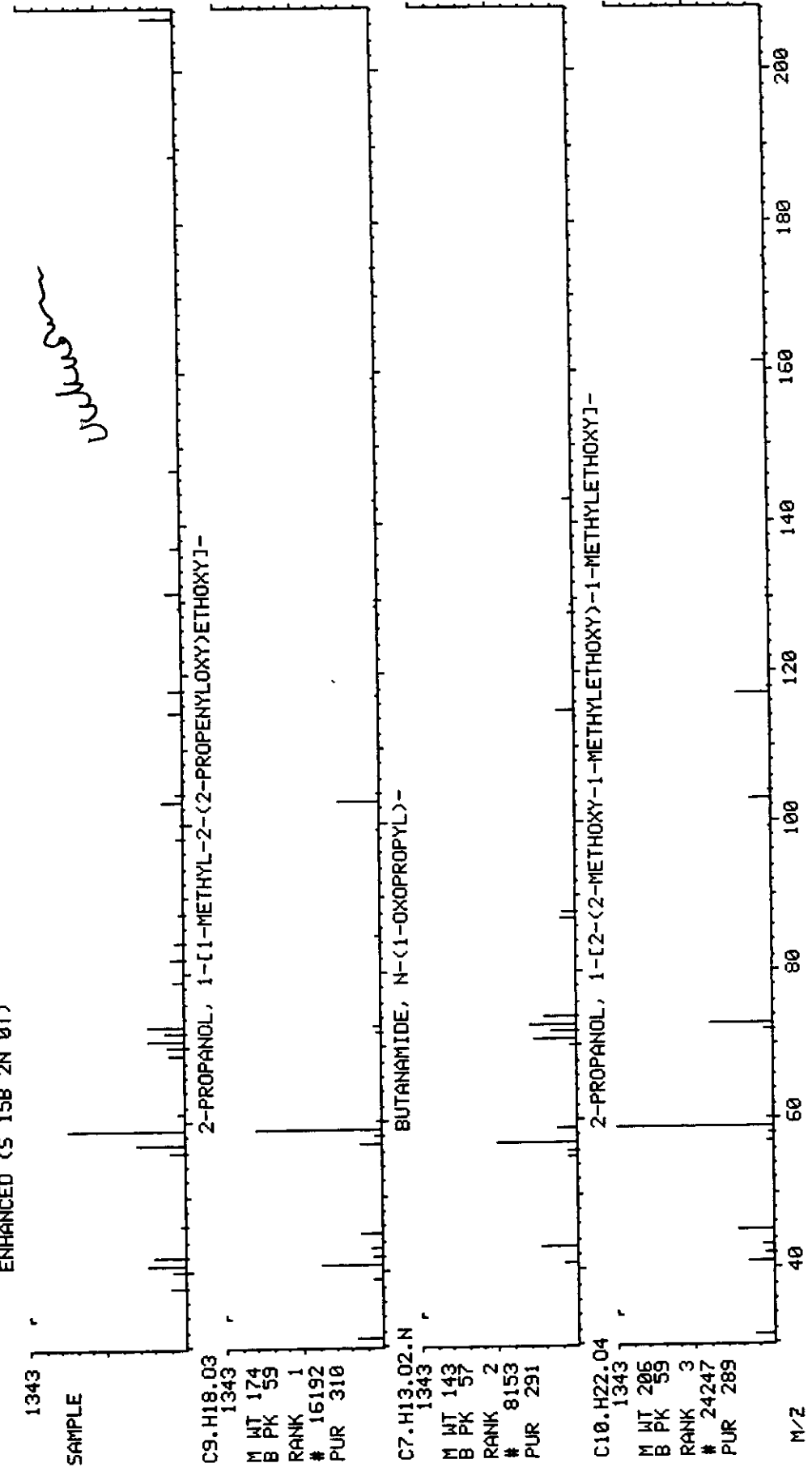
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C9.H18.O3	174	59	310	722	370
2	C7.H13.O2.N	143	57	291	604	404
3	C10.H22.O4	206	59	289	581	424
4	C6.H12.O6	180	72	278	621	361
5	C8.H18.O2	146	57	273	695	366
6	C10.H22.O	158	71	271	607	351
7	C3.H9.O6.B3	174	73	265	562	312
8	C9.H20.O	144	73	264	590	357
9	C4.H10.O2.N2	118	103	253	669	274

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	55956-25-7
2	---	---	---	---	32796-69-3
3	---	---	---	---	20324-33-8
4	---	---	---	---	50-99-7
5	---	---	---	---	4346-01-4
6	---	---	---	---	544-01-4
7	---	---	---	---	102-24-9
8	---	---	---	---	19549-74-7
9	---	---	---	---	30893-20-0

MID LIBRARY SEARCH (LIBRARYNB)  
 08/31/98 16:02:00 + 22:49  
 SAMPLE: 5BLKSX 31JUL98-16A 1/35A/1ML  
 CONDS.: UG/ML \*1007/1002 \*(NA/NA) /1/35A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSX #2052  
 CALL: 300681MBSX # 3  
 INST. ID: F16

BASE M/Z: 59  
 RIC: 1154.



1001

Library Search Data: 300681MBSX #2057 Base m/z: 59  
 08/31/98 16:02:00 + 22:53 Cali: 300681MBSX # 3 RIC: 1700.  
 Sample: SBLKSX 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/3SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 722 matched at least 5 of the 16 largest peaks in the unknown

Rank In. Name  
 1 3528 2-METHYL-2,3-PENTANEDIOL  
 2 1909 2-PROPANOL, 1-METHOXY-2-METHYL-  
 3 1919 1-BUTANOL, 3-METHOXY-  
 4 9227 1,3,3-TRIMETHOXYBUTANE  
 5 3534 1-BUTANOL, 4-ETHOXY-  
 6 5810 2-PENTANOL, 5-METHOXY-2-METHYL-  
 7 5522 2-HEXANOL, 2,3-DIMETHYL-  
 8 36537 2,5,8,11-TETRAOXATETRADECAN-13-OL, 4,7,10-TRIMETHYL-  
 9 9010 ACETAMIDE, 2-METHOXY-N-(2-METHOXYETHYL)-

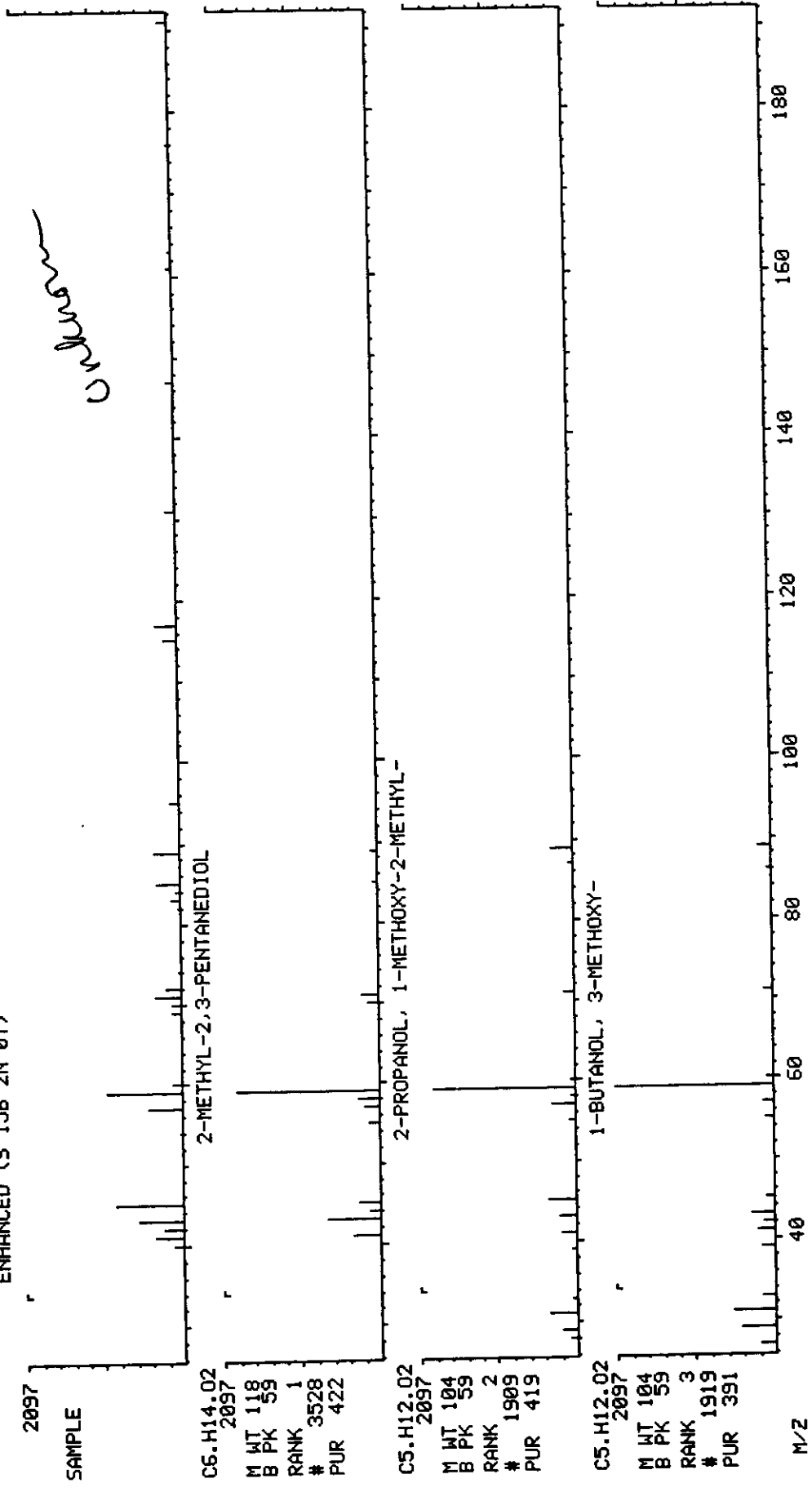
Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C6.H14.O2	118	59	422	748	481
2	C5.H12.O2	104	59	419	836	449
3	C5.H12.O2	104	59	391	746	426
4	C7.H16.O3	148	45	389	663	457
5	C6.H14.O2	118	59	384	692	512
6	C7.H16.O2	132	59	384	733	455
7	C8.H18.O	130	59	381	709	423
8	C13.H28.O5	264	59	370	569	549
9	C6.H13.O3.N	147	45	368	665	479

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	7795-80-4
2	---	---	---	---	3587-64-2
3	---	---	---	---	2517-43-3
4	---	---	---	---	6607-66-5
5	---	---	---	---	111-73-9
6	---	---	---	---	55724-04-4
7	---	---	---	---	19550-03-9
8	---	---	---	---	20324-34-9
9	---	---	---	---	55956-18-8

DATA: 300681MBSX #2057  
CALI: 300681MBSX # 3  
BASE M/Z: 59  
RIC: 1700.

DATA: 300681MBSX #2057  
CALI: 300681MBSX # 3  
INST. ID: F16

MID LIBRARY SEARCH (LIBRARYNB)  
08/31/98 16:02:00 + 22:53  
SAMPLE: 5BLK5X 31JUL98-16A 1/35A/1ML  
CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA >)/1/35A NA M  
ENHANCED (S 15B 2N 0T)



1000

TIC SELECTION REPORT

DATA FILE: 300681MBSX

THE FOLLOWING PEAKS WERE REJECTED BECAUSE AT LEAST 40 % OF THEIR SIZE WAS ACCOUNTED FOR BY TARGET COMPOUNDS ELUTING WITHIN 4 SCANS OF THE PEAK TOP.

SCAN	SIZE	AMOUNT
201	344792.	51.759
309	406704.	61.053
342	223900.	33.611
422	183296.	27.516
530	266460.	40.000
723	358875.	44.289
828	324123.	40.000
968	355248.	42.527
1086	334141.	40.000
1383	308474.	61.749
1383	307034.	61.460
1551	199826.	40.000
1782	156554.	40.000

TOTAL NUMBER OF UNIDENTIFIED PEAKS WITH SIZE GREATER THAN 10 % OF THE CLOSEST INTERNAL STANDARD THAT DOES NOT HAVE INTERFERENCES = 12

INTERNAL STANDARDS THAT HAVE RIC SIZE LESS THAN 50 % OR GREATER THAN 200 % OF THE ESTIMATED RIC SIZE ARE CONSIDERED TO HAVE INTERFERENCES AND WILL NOT BE USED FOR QUANTITATION.

#	INTERNAL STANDARD	PERCENT OF ESTIMATED		SCAN
		RIC SIZE	RIC SIZE	
1	CI40 NAPHTHALENE-D8	266460.	127	530
2	CI50 ACENAPHTHENE-D10	324123.	124	828
3	CI60 PHENANTHRENE-D10	334141.	119	1086
4	CI70 CHRYSENE-D12	199826.	127	1551
5	CI75 PERYLENE-D12	156554.	123	1782

\* INDICATES INTERFERENCE

SIZE = AREA



QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSF  
Std Id: ST16980831

Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 16:31  
Analyst: DAT Instrument ID: F16 Run Factor: 1.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC Limits
CS20 NITROBENZENE-D5	50.00	28.10	56.2	45 107
CS25 2-FLUOROBIPHENY	50.00	27.30	54.6	* 62 110
CS30 TERPHENYL-D14	50.00	38.70	77.4	58 135
CS45 PHENOL-D5	100.0	19.40	19.4	* 43 130
CS50 2-FLUOROPHENOL	100.0	37.00	37.0	36 111
CS55 2,4,6-TRIBROMOP	100.0	71.80	71.8	58 131

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
HEXACHLOROENZENE-C13		ND	UG/A	10.0
C310 N-NITROSODIMETHYLAMINE		ND	UG/A	10.0
PYRIDINE		ND	UG/A	20.0
2-PICOLINE		ND	UG/A	10.0
N-NITROSOMETHYLETHYLAMINE		ND	UG/A	10.0
METHYLMETHANESULFONATE		ND	UG/A	10.0
N-NITROSODIETHYLAMINE		ND	UG/A	10.0
ETHYLMETHANE SULFONATE		ND	UG/A	10.0
PENTACHLOROETHANE		ND	UG/A	10.0
C320 ANILINE		ND	UG/A	10.0
C315 PHENOL		ND	UG/A	10.0
C325 BIS(2-CHLOROETHYL)ETHER		ND	UG/A	10.0
C330 2-CHLOROPHENOL		ND	UG/A	10.0
C335 1,3-DICHLOROBENZENE		ND	UG/A	10.0
C340 1,4-DICHLOROBENZENE		ND	UG/A	10.0
C345 BENZYL ALCOHOL		ND	UG/A	10.0
C350 1,2-DICHLOROBENZENE		ND	UG/A	10.0
C355 2-METHYLPHENOL		ND	UG/A	10.0
C360 2,2'-OXYBIS(1-CLPROPAN)		ND	UG/A	10.0
C361 ACETOPHENONE		ND	UG/A	10.0
N-NITROSPYRROLIDINE		ND	UG/A	10.0
N-NITROSOMORPHOLINE		ND	UG/A	10.0
3-METHYL PHENOL		ND	UG/A	10.0
C365 4-METHYLPHENOL		ND	UG/A	10.0
C370 N-NITROSO-DI-N-PROPYLAM		ND	UG/A	10.0
O-TOLUIDINE		ND	UG/A	10.0
C375 HEXACHLOROETHANE		ND	UG/A	10.0
C410 NITROBENZENE		ND	UG/A	10.0
N-NITROSOPIPERIDINE		ND	UG/A	10.0

Reviewed by: 6/9/17/16/

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSF  
Std Id: ST16980831

Sample: SBLKSF 01AUG98-16A 1SA/1ML

INST. ID: F16

Client: PACIFI

Date Analyzed: 08/31/98 16:31

Analyst: DAT

Instrument ID: F16

Run Factor: 1.00

Quan List Threshold: 0.95

Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C415 ISOPHORONE		ND	UG/A	10.0
C420 2-NITROPHENOL		ND	UG/A	10.0
C425 2,4-DIMETHYLPHENOL		ND	UG/A	10.0
C435 BIS(2-CHLOROETHOXY)METH		ND	UG/A	10.0
C440 2,4-DICHLOROPHENOL		ND	UG/A	10.0
C430 BENZOIC ACID		ND	UG/A	50.0
AA-DIMETHYLPHENETHYLAMINE		ND	UG/A	10.0
C445 1,2,4-TRICHLOROBENZENE		ND	UG/A	10.0
C450 NAPHTHALENE		ND	UG/A	10.0
C455 4-CHLOROANALINE		ND	UG/A	10.0
2,6-DICHLOROPHENOL		ND	UG/A	10.0
HEXACHLOROPROPENE		ND	UG/A	10.0
C460 HEXACHLOROBUTADIENE		ND	UG/A	10.0
P-PHENYLENE DIAMINE		ND	UG/A	10.0
N-NITROSODI-N-BUTYLAMINE		ND	UG/A	10.0
C465 4-CHLORO-3-METHYLPHENO		ND	UG/A	10.0
SAFROLE		ND	UG/A	10.0
C470 2-METHYLNAPHTHALENE		ND	UG/A	10.0
1,2,4,5-TETRACHLOROBENZENE		ND	UG/A	10.0
ISOSAFROLE (#1)		ND	UG/A	20.0
C510 HEXACHLOROCYCLOPENTADI		ND	UG/A	10.0
C515 2,4,6-TRICHLOROPHENOL		ND	UG/A	10.0
C520 2,4,5-TRICHLOROPHENOL		ND	UG/A	10.0
ISOSAFROLE (#2)		ND	UG/A	20.0
C525 2-CHLORONAPHTHALENE		ND	UG/A	10.0
1-CHLORONAPHTHALENE		ND	UG/A	5.0
C530 2-NITROANALINE		ND	UG/A	50.0
1,4-NAPHTHOQUINONE		ND	UG/A	10.0
C535 DIMETHYLPHTHALATE		ND	UG/A	10.0
1,3-DINITROBENZENE		ND	UG/A	10.0
C540 ACENAPHTHYLENE		ND	UG/A	10.0
C543 2,6-DINITROTOLUENE		ND	UG/A	10.0
C545 3-NITROANILINE		ND	UG/A	50.0
C550 ACENAPHTHENE		ND	UG/A	10.0
C555 2,4-DINITROPHENOL		ND	UG/A	50.0
C565 DIBENZOFURAN		ND	UG/A	10.0
C560 4-NITROPHENOL		ND	UG/A	50.0
PENTACHLOROBENZENE		ND	UG/A	10.0
C570 2,4-DINITROTOLUENE		ND	UG/A	10.0
1-NAPHTHYLAMINE		ND	UG/A	10.0
2-NAPHTHYLAMINE		ND	UG/A	10.0
2,3,4,6-TETRACHLOROPHENOL		ND	UG/A	20.0
C580 DIETHYLPHTHALATE		ND	UG/A	10.0
C590 FLUORENE		ND	UG/A	10.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSF  
Std Id: ST16980831

Sample: SBLKSF 01AUG98-16A 1SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 08/31/98 16:31  
Run Factor: 1.00  
Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C585 4-CHLOROPHENYL-PHENYLE		ND	UG/A	10.0
5-NITRO-O-TOLIDINE		ND	UG/A	10.0
C595 4-NITROANALINE		ND	UG/A	50.0
C610 4,6-DINITRO-2-METHYLPH		ND	UG/A	50.0
C615 N-NITROSODIPHENYLAMINE		ND	UG/A	10.0
C620 AZOBENZENE		ND	UG/A	10.0
SYM-TRINITROBENZENE		ND	UG/A	10.0
C625 4-BROMOPHENYL-PHENYLET		ND	UG/A	10.0
PHENACETIN		ND	UG/A	10.0
DIALLATE	234	ND	UG/A	10.0
C630 HEXACHLOROBENZENE		ND	UG/A	10.0
4-AMINOBIIPHENYL		ND	UG/A	10.0
C635 PENTACHLOROPHENOL		ND	UG/A	50.0
PRONAMIDE		ND	UG/A	10.0
PENTACHLORONITROBENZENE		ND	UG/A	50.0
C640 PHENANTHRENE		ND	UG/A	10.0
C645 ANTHRACENE		ND	UG/A	10.0
2SECBUTYL-4,6-DINITROPHENOL		ND	UG/A	10.0
C647 CARBAZOLE		ND	UG/A	10.0
C650 DI-N-BUTYLPHTHALATE		ND	UG/A	10.0
4-NITROQUINOLINE-1-OXIDE		ND	UG/A	10.0
METHAPYRILENE		ND	UG/A	10.0
ISODRIN		ND	UG/A	10.0
C655 FLUORANTHENE		ND	UG/A	10.0
CHLOROBENZILATE		ND	UG/A	10.0
C710 BENZIDINE		ND	UG/A	100.0
C715 PYRENE		ND	UG/A	10.0
ARAMITE (#1)		ND	UG/A	10.0
ARAMITE (#2)		ND	UG/A	10.0
P-DIMETHYLAMINDAZOBENZENE		ND	UG/A	10.0
3,3'-DIMETHYLBENZIDINE		ND	UG/A	10.0
KEPONE		ND	UG/A	50.0
C720 BUTYLBENZYLPHTHALATE		ND	UG/A	10.0
2-ACETYLAMINOFLUORENE		ND	UG/A	10.0
C730 BENZO(A)ANTHRACENE		ND	UG/A	10.0
C725 3,3'-DICHLOROBENZIDINE		ND	UG/A	20.0
C740 CHRYSENE		ND	UG/A	10.0
C745 BIS(2-ETHYLHEXYL)PHTHA		ND	UG/A	10.0
3-METHYLCHOLANTHRENE		ND	UG/A	10.0
C760 DI-N-OCTYL PHTHALATE		ND	UG/A	10.0
C765 BENZO(B)FLUORANTHENE		ND	UG/A	10.0
7,12-DIMETHYLBENZANTHRACENE		ND	UG/A	10.0
C770 BENZO(K)FLUORANTHENE		ND	UG/A	10.0
HEXACHLOROPHENE		ND	UG/A	10.0

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSF  
Std Id: ST16980831

Sample: SBLKSF 01AUG98-16A 1SA/1ML

INST. ID: F16

Client: PACIFI

Date Analyzed: 08/31/98 16:31

Analyst: DAT Instrument ID: F16

Run Factor: 1.00

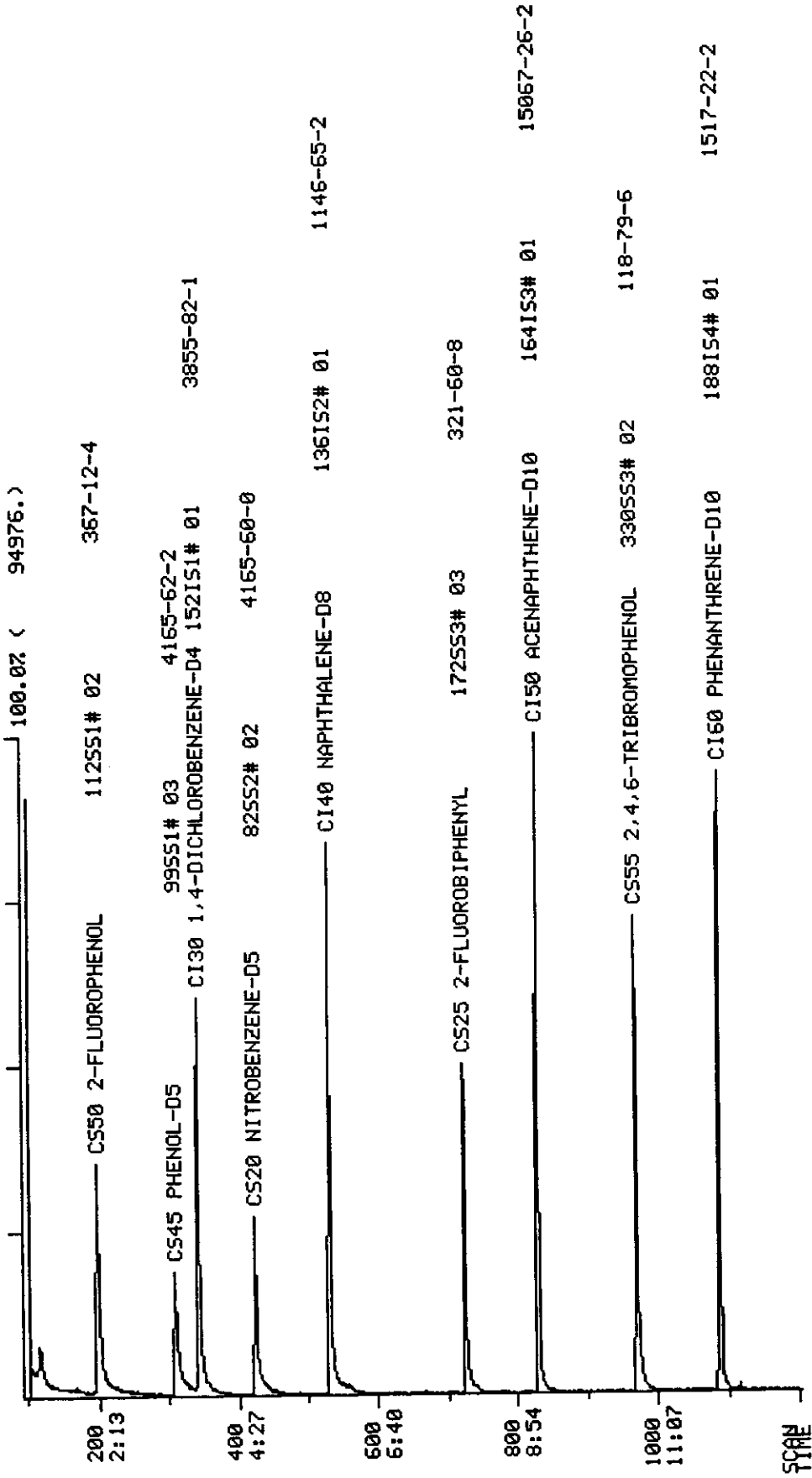
Quan List Threshold: 0.95

Surrogate Vol.: 1.00

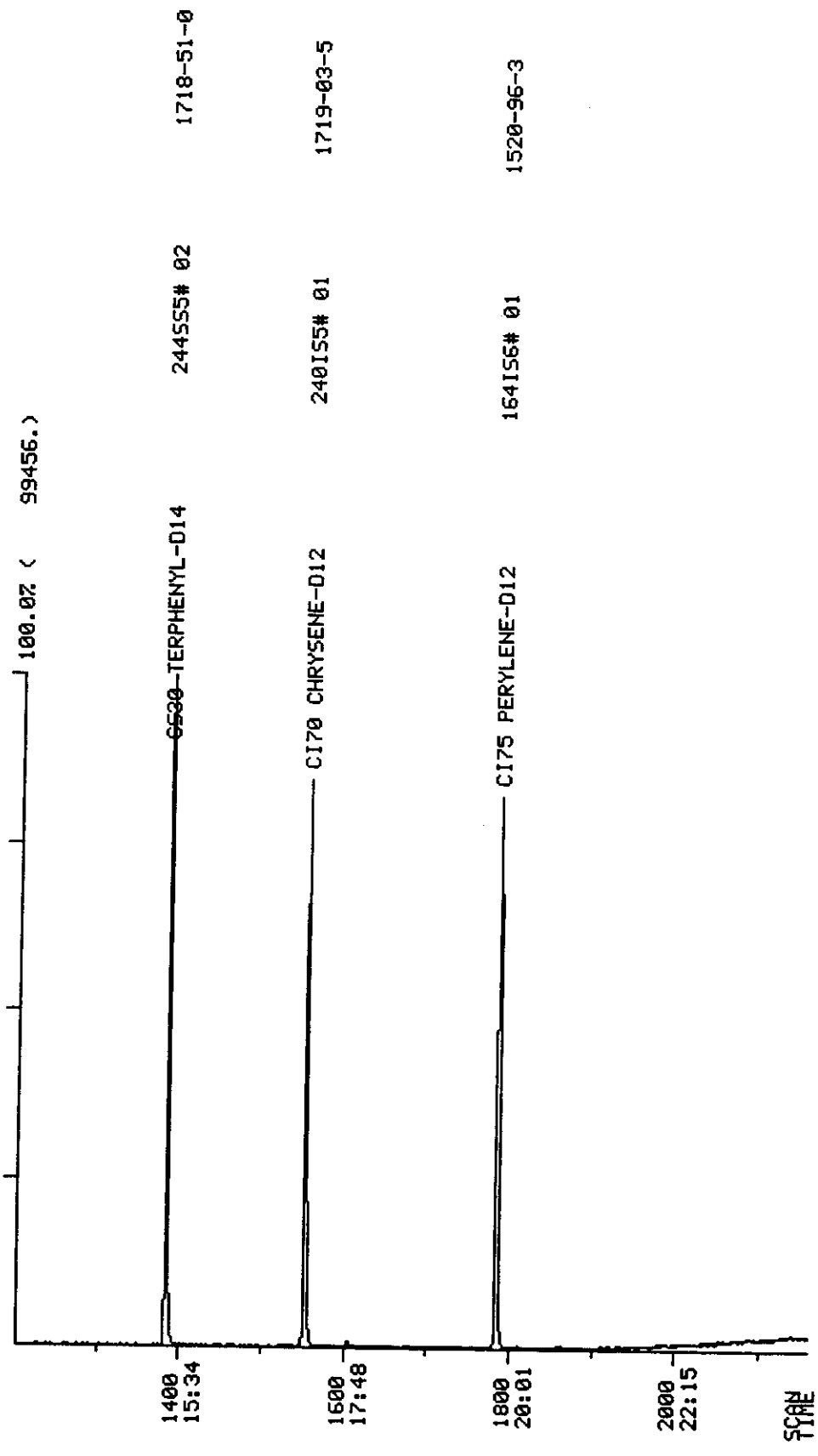
Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C775 BENZO(A)PYRENE		ND	UG/A	10.0
C780 INDENO(1,2,3-CD)PYRENE		ND	UG/A	10.0
C785 DIBENZ(A,H)ANTHRACENE		ND	UG/A	10.0
C790 BENZO(G,H,I)PERYLENE		ND	UG/A	10.0

DATA FROM FILE: 300681MBSF SCANS 93 TO 1203 ACQUIRED: 08/31/98 16:31:00  
 CALI: 300681MBSF #3  
 SAMPLE: SBLKSF 01AUG98-16A 15A/1ML INST. ID: F16  
 COND.: UG/ML \*100%/100% \*(NA/NA )/15A NA M



DATA FROM FILE: 300681MBSF SCANS 1203 TO 2158 ACQUIRED: 08/31/98 16:31:00  
 CALI: 300681MBSF #3  
 SAMPLE: SBLKSF 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA )/15A NA M



1700

ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBSF

Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 16:31

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML							
No	Entry	Name	Mass	Meth	Scan	Ref	Area	RRF(L)	Amount
1	S1# 1	CI30 1,4-DICHLOROBENZENE	152	A	BB	340 1	30721.	1.000	40.000
2	S2# 1	CI40 NAPHTHALENE-D8	136	A	BB	529 2	108221.	1.000	40.000
3	S3# 1	CI50 ACENAPHTHENE-D10	164	A	BB	828 3	59070.	1.000	40.000
4	S4# 1	CI60 PHENANTHRENE-D10	188	A	BB	1086 4	102024.	1.000	40.000
5	S5# 1	CI70 CHRYSENE-D12	240	A	BB	1551 5	87857.	1.000	40.000
6	S6# 1	CI75 PERYLENE-D12	264	A	BB	1783 6	75258.	1.000	40.000
7	S2# 2	CS20 NITROBENZENE-D5	82	A	BB	421 2	35709.	0.470	28.108
8	S3# 3	CS25 2-FLUOROBIPHENYL	172	A	BB	722 3	49743.	1.234	27.290
9	S5# 2	CS30 TERPHENYL-D14	244	A	BB	1383 5	84547.	0.995	38.678
10	S1# 3	CS45 PHENOL-D5	99	A	BB	307 1	30107.	2.025	19.354
11	S1# 2	CS50 2-FLUOROPHENOL	112	A	BB	196 1	38883.	1.368	37.002
12	S3# 2	CS55 2,4,6-TRIBROMOPHENO	330	A	BB	968 3	24984.	0.236	71.791
13	S4# 4	HEXACHLOROBENZENE-C13	294			4 NOT FOUND			
14	S1# 15	C310 N-NITROSODIMETHYLAM	74			1 NOT FOUND			
15	S1# 20	PYRIDINE	79			1 NOT FOUND			
16	S1# 30	2-PICOLINE	93			1 NOT FOUND			
17	S1# 40	N-NITROSOMETHYLETHYLAMIN	42	A	BB	159 1	156.	0.811	0.250
18	S1# 60	METHYLMETHANESULFONATE	80			1 NOT FOUND			
19	S1# 70	N-NITROSODIETHYLAMINE	102			1 NOT FOUND			
20	S1# 85	ETHYLMETHANE SULFONATE	79			1 NOT FOUND			
21	S1# 95	PENTACHLOROETHANE	117			1 NOT FOUND			
22	S1#100	C320 ANILINE	93			1 NOT FOUND			
23	S1#105	C315 PHENOL	94			1 NOT FOUND			
24	S1#110	C325 BIS(2-CHLOROETHYL)E	93			1 NOT FOUND			
25	S1#115	C330 2-CHLOROPHENOL	128			1 NOT FOUND			
26	S1#125	C335 1,3-DICHLOROBENZENE	146			1 NOT FOUND			
27	S1#130	C340 1,4-DICHLOROBENZENE	146			1 NOT FOUND			
28	S1#145	C345 BENZYL ALCOHOL	108			1 NOT FOUND			
29	S1#150	C350 1,2-DICHLOROBENZENE	146			1 NOT FOUND			
30	S1#160	C355 2-METHYLPHENOL	108			1 NOT FOUND			
31	S1#165	C360 2,2'-OXYBIS(1-CLPRO	45			1 NOT FOUND			
32	S1#170	C361 ACETOPHENONE	105			1 NOT FOUND			
33	S1#175	N-NITROSOPYRROLIDINE	100			1 NOT FOUND			
34	S1#180	N-NITROSOMORPHOLINE	56			1 NOT FOUND			
35	S1#182	3-METHYL PHENOL	108			1 NOT FOUND			
36	S1#185	C365 4-METHYLPHENOL	108			1 NOT FOUND			
37	S1#190	C370 N-NITROSO-DI-N-PROP	70			1 NOT FOUND			
38	S1#195	O-TOLUIDINE	106			1 NOT FOUND			
39	S1#200	C375 HEXACHLOROETHANE	117			1 NOT FOUND			
40	S2# 10	C410 NITROBENZENE	77			2 NOT FOUND			
41	S2# 15	N-NITROSOPIPERIDINE	42			2 NOT FOUND			
42	S2# 20	C415 ISOPHORONE	82			2 NOT FOUND			
43	S2# 25	C420 2-NITROPHENOL	139			2 NOT FOUND			
44	S2# 30	C425 2,4-DIMETHYLPHENOL	107			2 NOT FOUND			
45	S2# 35	C435 BIS(2-CHLOROETHOXY)	93			2 NOT FOUND			

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46	S2# 40	C440	2,4-DICHLOROPHENOL	162	2	NOT FOUND
47	S2# 45	C430	BENZOIC ACID	122	2	NOT FOUND
48	S2# 50	AA-	DIMETHYLPHENETHYLAMIN	58	2	NOT FOUND
49	S2# 55	C445	1,2,4-TRICHLOROBENZ	180	2	NOT FOUND



50 S2# 60 C450 NAPHTHALENE	128	2	NOT FOUND
51 S2# 80 C455 4-CHLOROANALINE	127	2	NOT FOUND
52 S2# 85 2,6-DICHLOROPHENOL	162	2	NOT FOUND
53 S2# 90 HEXACHLOROPROPENE	213	2	NOT FOUND
54 S2# 95 C460 HEXACHLOROBUTADIENE	225	2	NOT FOUND
55 S2#115 P-PHENYLENE DIAMINE	108	2	NOT FOUND
56 S2#120 N-NITROSODI-N-BUTYLAMINE	84	2	NOT FOUND
57 S2#130 C465 4-CHLORO-3-METHYLPH	107	2	NOT FOUND
58 S2#140 SAFROLE	162	2	NOT FOUND
59 S2#145 C470 2-METHYLNAPHTHALENE	142	2	NOT FOUND
60 S3# 10 1,2,4,5-TETRACHLOROBENZE	214	3	NOT FOUND
61 S3# 15 ISOSAFROLE (#1)	162	3	NOT FOUND
62 S3# 20 C510 HEXACHLOROCCYCLOPENT	237	3	NOT FOUND
63 S3# 25 C515 2,4,6-TRICHLOROPHEN	196	3	NOT FOUND
64 S3# 30 C520 2,4,5-TRICHLOROPHEN	196	3	NOT FOUND
65 S3# 35 ISOSAFROLE (#2)	104	3	NOT FOUND
66 S3# 40 C525 2-CHLORONAPHTHALENE	162	3	NOT FOUND
67 S3# 42 1-CHLORONAPHTHALEN	162	3	NOT FOUND
68 S3# 45 C530 2-NITROANALINE	65	3	NOT FOUND
69 S3# 50 1,4-NAPHTHOQUINONE	158	3	NOT FOUND
70 S3# 55 C535 DIMETHYLPHTHALATE	163	3	NOT FOUND
71 S3# 60 1,3-DINITROBENZENE	168	3	NOT FOUND
72 S3# 65 C540 ACENAPHTHYLENE	152	3	NOT FOUND
73 S3# 70 C543 2,6-DINITROTOLUENE	165	3	NOT FOUND
74 S3# 75 C545 3-NITROANILINE	138	3	NOT FOUND
75 S3# 80 C550 ACENAPHTHENE	153	3	NOT FOUND
76 S3# 85 C555 2,4-DINITROPHENOL	184	3	NOT FOUND
77 S3# 90 C565 DIBENZOFURAN	168	3	NOT FOUND
78 S3# 95 C560 4-NITROPHENOL	109	3	NOT FOUND
79 S3#100 PENTACHLOROBENZENE	250	3	NOT FOUND
80 S3#105 C570 2,4-DINITROTOLUENE	165	3	NOT FOUND
81 S3#110 1-NAPHTHYLAMINE	143	3	NOT FOUND
82 S3#115 2-NAPHTHYLAMINE	143	3	NOT FOUND
83 S3#120 2,3,4,6-TETRACHLOROPHENO	232	3	NOT FOUND
84 S3#130 C580 DIETHYLPHTHALATE	149	3	NOT FOUND
85 S3#135 C590 FLUORENE	166	3	NOT FOUND
86 S3#140 C585 4-CHLOROPHENYL-PHEN	204	3	NOT FOUND
87 S3#145 5-NITRO-O-TOLUIDINE	152	3	NOT FOUND
88 S3#150 C595 4-NITROANALINE	138	3	NOT FOUND
89 S4# 10 C610 4,6-DINITRO-2-METHY	198	4	NOT FOUND
90 S4# 15 C615 N-NITROSODIPHENYLAM	169	4	NOT FOUND
91 S4# 20 C620 AZOBENZENE	77	4	NOT FOUND
92 S4# 25 SYM-TRINITROBENZENE	75	4	NOT FOUND
93 S4# 30 C625 4-BROMOPHENYL-PHENY	248	4	NOT FOUND
94 S4# 35 PHENACETIN	108	4	NOT FOUND
95 S4# 37 DIALLATE	234	4	NOT FOUND
96 S4# 40 C630 HEXACHLOROBENZENE	284	4	NOT FOUND
97 S4# 45 4-AMINOBIIPHENYL	169	4	NOT FOUND
98 S4# 50 C635 PENTACHLOROPHENOL	266	4	NOT FOUND
99 S4# 55 PRONAMIDE	173	4	NOT FOUND
100 S4# 60 PENTACHLORONITROBENZENE	237	4	NOT FOUND
101 S4# 65 C640 PHENANTHRENE	178	4	NOT FOUND
102 S4# 70 C645 ANTHRACENE	178	4	NOT FOUND
103 S4# 75 2SECBUTYL-4,6-DINITROPHE	211	4	NOT FOUND
104 S4# 80 C647 CARBAZOLE	167	4	NOT FOUND
105 S4# 85 C650 DI-N-BUTYLPHTHALATE	149 A BB 1226 4 796. 1.247 0.250		
106 S4#100 4-NITROQUINOLINE-1-OXIDE	190	4	NOT FOUND
107 S4#105 METHAPYRILENE	58	4	NOT FOUND
108 S4#106 ISODRIN	193	4	NOT FOUND
109 S4#110 C655 FLUORANTHENE	202	4	NOT FOUND
110 S4#120 CHLOROBENZILATE	139	4	NOT FOUND

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111	S5# 10	C710 BENZIDINE	184	5	NOT FOUND		
112	S5# 15	C715 PYRENE	202	5	NOT FOUND		
113	S5# 20	ARAMITE (#1)	185	5	NOT FOUND		
114	S5# 25	ARAMITE (#2)	185	5	NOT FOUND		
115	S5# 30	P-DIMETHYLAMINOAZOBENZEN	120	5	NOT FOUND		
116	S5# 35	3,3'-DIMETHYLBENZIDINE	212	5	NOT FOUND		
117	S5# 37	KEPONE	272	5	NOT FOUND		
118	S5# 40	C720 BUTYLBENZYLPHTHALAT	149	5	NOT FOUND		
119	S5# 45	2-ACETYLAMINOFLUORENE	181	5	NOT FOUND		
120	S5# 50	C730 BENZO(A)ANTHRACENE	228	5	NOT FOUND		
121	S5# 55	C725 3,3'-DICHLOROBENZID	252	5	NOT FOUND		
122	S5# 60	C740 CHRYSENE	228	5	NOT FOUND		
123	S5# 65	C745 BIS(2-ETHYLHEXYL)PH	149	A BB 1603	5	576.0.980	0.268
124	S5# 85	3-METHYLCHOLANTHRENE	268	5	NOT FOUND		
125	S6# 10	C760 DI-N-OCTYL PHTHALAT	149	6	NOT FOUND		
126	S6# 15	C765 BENZO(B)FLUORANTHEN	252	6	NOT FOUND		
127	S6# 20	7,12-DIMETHYLBENZANTHRAC	256	6	NOT FOUND		
128	S6# 25	C770 BENZO(K)FLUORANTHEN	252	6	NOT FOUND		
129	S6# 30	HEXACHLOROPHENE	196	6	NOT FOUND		
130	S6# 35	C775 BENZO(A)PYRENE	252	6	NOT FOUND		
131	S6# 55	C780 INDENO(1,2,3-CD)PYR	276	6	NOT FOUND		
132	S6# 60	C785 DIBENZ(A,H)ANTHRACE	278	6	NOT FOUND		
133	S6# 65	C790 BENZO(G,H,I)PERYLEN	276	6	NOT FOUND		

QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 300681MBSF	30721.= 107		3.78	108221.= 107		5.88	59070.= 108		9.21

IS# 1 = C130 1,4-DICHLOROBENZENE-D4  
 IS# 2 = C140 NAPHTHALENE-D8  
 IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 300681MBSF	102024.= 103		12.08	87857.= 141		17.25	75258.= 148		19.83

IS# 4 = C160 PHENANTHRENE-D10  
 IS# 5 = C170 CHRYSENE-D12  
 IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

Data Reduced by: \_\_\_\_ Date: \_\_\_\_  
Data Reviewed by: \_\_\_\_ Date: \_\_\_\_

Data File: 300681MBSF

QUANTERRA GC/MS TIC REPORT ( Part 1 )

Sample: SBLKSF 01AUG98-16A 1SA/1ML  
Analyst: DAT

INST. ID: F16  
Date Analyzed: 08/31/98 16:31  
Run Factor: 1.00

# SCAN	Concentration in Sample (UG/A)	CAS #
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No peaks to search.

1708

DATA FILE: 300681MBSF

QUANTERRA GC/MS TIC REPORT ( Part 2 )

CONCENTRATION = AREA(TIC)\*CONC(IS)/AREA(IS)

#	FIT PURITY STD.	INT.	RT	RRT	AREA	HEIGHT	AMOUNT AS ANALYZED (UG/ML )	LIB	LIB #
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1707

TIC SELECTION REPORT

DATA FILE: 300681MBSF

THE FOLLOWING PEAKS WERE REJECTED BECAUSE  
AT LEAST 40 % OF THEIR SIZE WAS ACCOUNTED FOR BY  
TARGET COMPOUNDS ELUTING WITHIN 4 SCANS OF THE  
PEAK TOP.

SCAN	SIZE	AMOUNT
196	129024.	22.541
307	75740.	13.232
340	185996.	32.494
421	91232.	15.939
529	228960.	40.000
722	152596.	22.325
828	273414.	40.000
968	231786.	33.574
1086	276153.	40.000
1384	308768.	44.436
1384	308768.	44.436
1551	277946.	40.000
1783	258361.	40.000

TOTAL NUMBER OF UNIDENTIFIED PEAKS WITH SIZE  
GREATER THAN 10 % OF THE CLOSEST INTERNAL STANDARD  
THAT DOES NOT HAVE INTERFERENCES = 0

INTERNAL STANDARDS THAT HAVE RIC SIZE LESS THAN  
50 % OR GREATER THAN 200 % OF THE ESTIMATED RIC SIZE  
ARE CONSIDERED TO HAVE INTERFERENCES AND WILL NOT BE USED  
FOR QUANTITATION.

#	INTERNAL STANDARD	RIC SIZE	PERCENT OF ESTIMATED RIC SIZE	SCAN
1	C140 NAPHTHALENE-D8	228960.	132	529
2	C150 ACENAPHTHENE-D10	273414.	128	828
3	C160 PHENANTHRENE-D10	276153.	119	1086
4	C170 CHRYSENE-D12	277946.	129	1551
5	C175 PERYLENE-D12	258361.	128	1783

\* INDICATES INTERFERENCE

SIZE = AREA

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSFI  
Std Id: ST16980902

Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 09/02/98 16:29  
Analyst: DAT Instrument ID: F16 Run Factor: 1.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

*confirms*

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	27.50	55.0	45 107
CS25 2-FLUOROBIPHENY	50.00	26.00	52.0	* 62 110
CS30 TERPHENYL-D14	50.00	37.90	75.8	58 135
CS45 PHENOL-D5	100.0	16.90	16.9	* 43 130
CS50 2-FLUOROPHENOL	100.0	35.80	35.8	* 36 111
CS55 2,4,6-TRIBROMOP	100.0	72.50	72.5	58 131

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
CS70 2-CHLOROPHENOL-D4		ND	UG/A	10.0
CS75 1,2-DICHLOROBENZ-D4		ND	UG/A	10.0
HEXACHLOROBENZENE-C13		ND	UG/A	10.0
C310 N-NITROSODIMETHYLAMINE		ND	UG/A	10.0
PYRIDINE		ND	UG/A	20.0
2-PICOLINE		ND	UG/A	10.0
N-NITROSOMETHYLETHYLAMINE		ND	UG/A	10.0
METHYLMETHANESULFONATE		ND	UG/A	10.0
N-NITROSODIETHYLAMINE		ND	UG/A	10.0
ETHYLMETHANE SULFONATE		ND	UG/A	10.0
PENTACHLOROETHANE		ND	UG/A	10.0
C320 ANILINE		ND	UG/A	10.0
C315 PHENOL		ND	UG/A	10.0
C325 BIS(2-CHLOROETHYL)ETHER		ND	UG/A	10.0
C330 2-CHLOROPHENOL		ND	UG/A	10.0
C335 1,3-DICHLOROBENZENE		ND	UG/A	10.0
C340 1,4-DICHLOROBENZENE		ND	UG/A	10.0
C345 BENZYL ALCOHOL		ND	UG/A	10.0
C350 1,2-DICHLOROBENZENE		ND	UG/A	10.0
C355 2-METHYLPHENOL		ND	UG/A	10.0
C360 2,2'-OXYBIS(1-CLPROPAN)		ND	UG/A	10.0
C361 ACETOPHENONE		ND	UG/A	10.0
N-NITROSPYRROLIDINE		ND	UG/A	10.0
N-NITROSOMORPHOLINE		ND	UG/A	10.0
3-METHYL PHENOL		ND	UG/A	10.0
C365 4-METHYLPHENOL		ND	UG/A	10.0
C370 N-NITROSO-DI-N-PROPYLAM		ND	UG/A	10.0
O-TOLUIDINE		ND	UG/A	10.0
C375 HEXACHLOROETHANE		ND	UG/A	10.0

1709

Reviewed by: \_\_\_\_\_

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSFI  
Std Id: ST16980902

Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 09/02/98 16:29  
Analyst: DAT Instrument ID: F16 Run Factor: 1.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C410 NITROBENZENE		ND	UG/A	10.0
N-NITROSOPIPERIDINE		ND	UG/A	10.0
C415 ISOPHORONE		ND	UG/A	10.0
C420 2-NITROPHENOL		ND	UG/A	10.0
C425 2,4-DIMETHYLPHENOL		ND	UG/A	10.0
C435 BIS(2-CHLOROETHOXY)METH		ND	UG/A	10.0
C440 2,4-DICHLOROPHENOL		ND	UG/A	10.0
C430 BENZOIC ACID		ND	UG/A	50.0
AA-DIMETHYLPHENETHYLAMINE		ND	UG/A	10.0
C445 1,2,4-TRICHLOROBENZENE		ND	UG/A	10.0
C450 NAPHTHALENE		ND	UG/A	10.0
C455 4-CHLOROANILINE		ND	UG/A	10.0
2,6-DICHLOROPHENOL		ND	UG/A	10.0
HEXACHLOROPROPENE		ND	UG/A	10.0
C460 HEXACHLOROBUTADIENE		ND	UG/A	10.0
P-PHENYLENE DIAMINE		ND	UG/A	10.0
N-NITROSODI-N-BUTYLAMINE		ND	UG/A	10.0
C465 4-CHLORO-3-METHYLPHENO		ND	UG/A	10.0
SAFROLE		ND	UG/A	10.0
C470 2-METHYLNAPHTHALENE		ND	UG/A	10.0
1,2,4,5-TETRACHLOROBENZENE		ND	UG/A	10.0
ISOSAFROLE (#1)		ND	UG/A	20.0
C510 HEXACHLOROCYCLOPENTADI		ND	UG/A	10.0
C515 2,4,6-TRICHLOROPHENOL		ND	UG/A	10.0
C520 2,4,5-TRICHLOROPHENOL		ND	UG/A	10.0
ISOSAFROLE (#2)		ND	UG/A	20.0
C525 2-CHLORONAPHTHALENE		ND	UG/A	10.0
1-CHLORONAPHTHALENE		ND	UG/A	5.0
C530 2-NITROANILINE		ND	UG/A	50.0
1,4-NAPHTHOQUINONE		ND	UG/A	10.0
C535 DIMETHYLPHTHALATE		ND	UG/A	10.0
1,3-DINITROBENZENE		ND	UG/A	10.0
C540 ACENAPHTHYLENE		ND	UG/A	10.0
C543 2,6-DINITROTOLUENE		ND	UG/A	10.0
C545 3-NITROANILINE		ND	UG/A	50.0
C550 ACENAPHTHENE		ND	UG/A	10.0
C555 2,4-DINITROPHENOL		ND	UG/A	50.0
C565 DIBENZOFURAN		ND	UG/A	10.0
C560 4-NITROPHENOL		ND	UG/A	50.0
PENTACHLOROBENZENE		ND	UG/A	10.0
C570 2,4-DINITROTOLUENE		ND	UG/A	10.0
1-NAPHTHYLAMINE		ND	UG/A	10.0
2-NAPHTHYLAMINE		ND	UG/A	10.0
2,3,4,6-TETRACHLOROPHENOL		ND	UG/A	20.0

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QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBSF1  
Std Id: ST16980902

Sample: SBLKSF 01AUG98-16A 1SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 09/02/98 16:29  
Run Factor: 1.00  
Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting Limit
C580 DIETHYLPHthalate		ND	UG/A	10.0
C590 FLUORENE		ND	UG/A	10.0
C585 4-CHLOROPHENYL-PHENYLE		ND	UG/A	10.0
5-NITRO-O-TOLUIDINE		ND	UG/A	10.0
C595 4-NITROANALINE		ND	UG/A	50.0
C610 4,6-DINITRO-2-METHYLPH		ND	UG/A	50.0
C615 N-NITROSODIPHENYLAMINE		ND	UG/A	10.0
C620 AZOBENZENE		ND	UG/A	10.0
SYM-TRINITROBENZENE		ND	UG/A	10.0
C625 4-BROMOPHENYL-PHENYLET		ND	UG/A	10.0
PHENACETIN		ND	UG/A	10.0
DIALlate	234	ND	UG/A	10.0
C630 HEXACHLOROBENZENE		ND	UG/A	10.0
4-AMINOBIphenyl		ND	UG/A	10.0
C635 PENTACHLOROPHENOL		ND	UG/A	50.0
PRONAMIDE		ND	UG/A	10.0
PENTACHLORONITROBENZENE		ND	UG/A	50.0
C640 PHENANTHRENE		ND	UG/A	10.0
C645 ANTHRACENE		ND	UG/A	10.0
2SECBUTYL-4,6-DINITROPHENOL		ND	UG/A	10.0
C647 CARBAZOLE		ND	UG/A	10.0
C650 DI-N-BUTYLPHthalate		ND	UG/A	10.0
4-NITROQUINOLINE-1-OXIDE		ND	UG/A	10.0
METHAPYRILENE		ND	UG/A	10.0
ISODRIN		ND	UG/A	10.0
C655 FLUORANTHENE		ND	UG/A	10.0
CHLOROBENZILATE		ND	UG/A	10.0
C710 BENZIDINE		ND	UG/A	100.0
C715 PYRENE		ND	UG/A	10.0
ARAMITE (#1)		ND	UG/A	10.0
ARAMITE (#2)		ND	UG/A	10.0
P-DIMETHYLAMINOAZOBENZENE		ND	UG/A	10.0
3,3'-DIMETHYLBENZIDINE		ND	UG/A	10.0
KEPONE		ND	UG/A	50.0
C720 BUTYLBENZYLPHthalate		ND	UG/A	10.0
2-ACETYLAMINOFLUORENE		ND	UG/A	10.0
C730 BENZO(A)ANTHRACENE		ND	UG/A	10.0
C725 3,3'-DICHLOROBENZIDINE		ND	UG/A	20.0
C740 CHRYSENE		ND	UG/A	10.0
C745 BIS(2-ETHYLHEXYL)PHTha		ND	UG/A	10.0
3-METHYLCHOLANTHRENE		ND	UG/A	10.0
C760 DI-N-OCTYL PHthalate		ND	UG/A	10.0
C765 BENZO(B)FLUORANTHENE		ND	UG/A	10.0
7,12-DIMETHYLBENZANTHRACENE		ND	UG/A	10.0

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QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681M8SF1  
Std Id: ST16980902

Sample: SBLKSF 01AUG98-16A 1SA/1ML

INST. ID: F16

Client: PACIFI

Date Analyzed: 09/02/98 16:29

Analyst: DAT Instrument ID: F16

Run Factor: 1.00

Quan List Threshold: 0.95

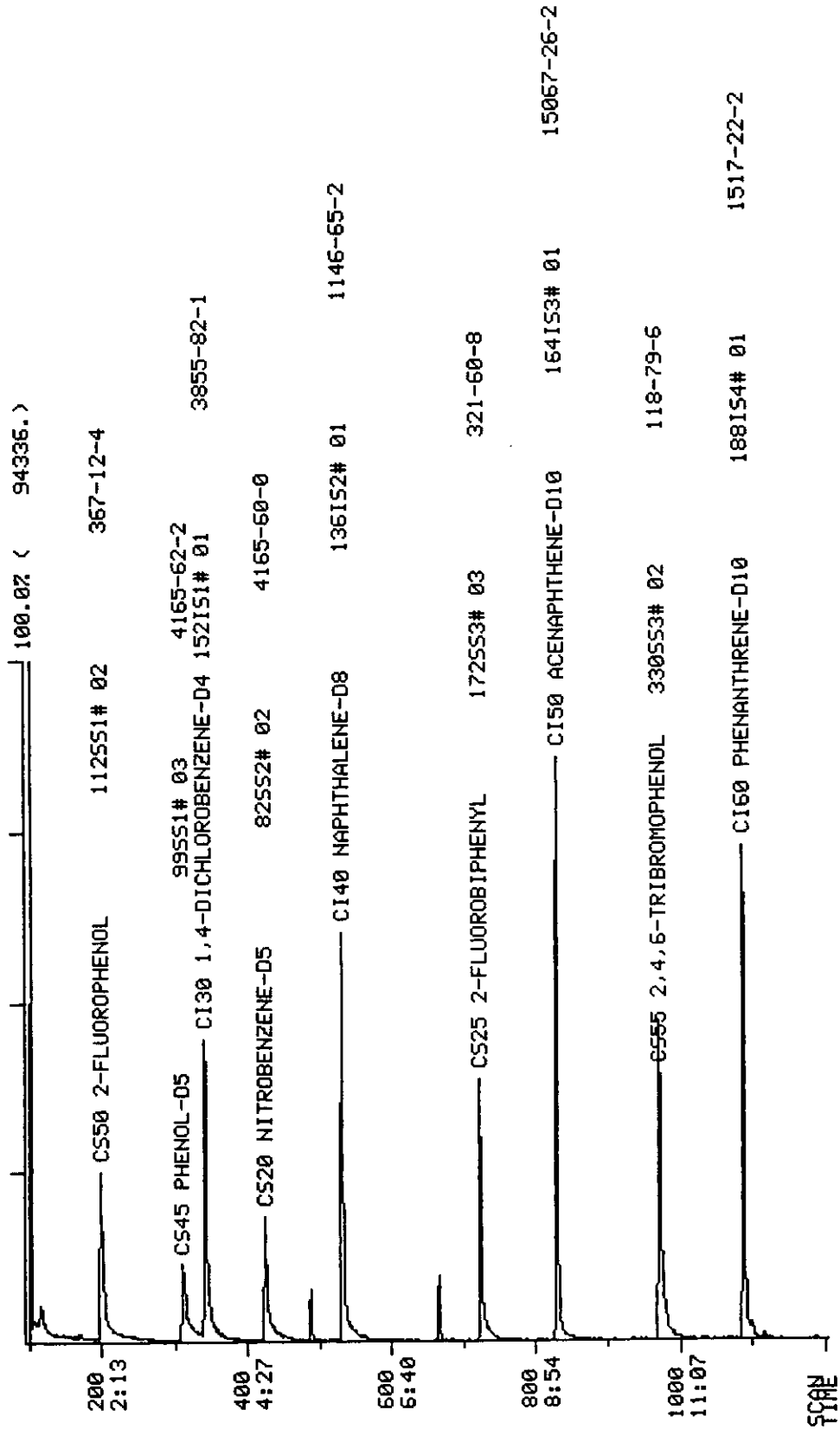
Surrogate Vol.: 1.00

Target Compounds: SAP9

Parameter	Scan	Result	Units	Reporting
				Limit
C770 BENZO(K)FLUORANTHENE		ND	UG/A	10.0
HEXACHLOROPHENE		ND	UG/A	10.0
C775 BENZO(A)PYRENE		ND	UG/A	10.0
C780 INDENO(1,2,3-CD)PYRENE		ND	UG/A	10.0
C785 DIBENZ(A,H)ANTHRACENE		ND	UG/A	10.0
C790 BENZO(G,H,I)PERYLENE		ND	UG/A	10.0

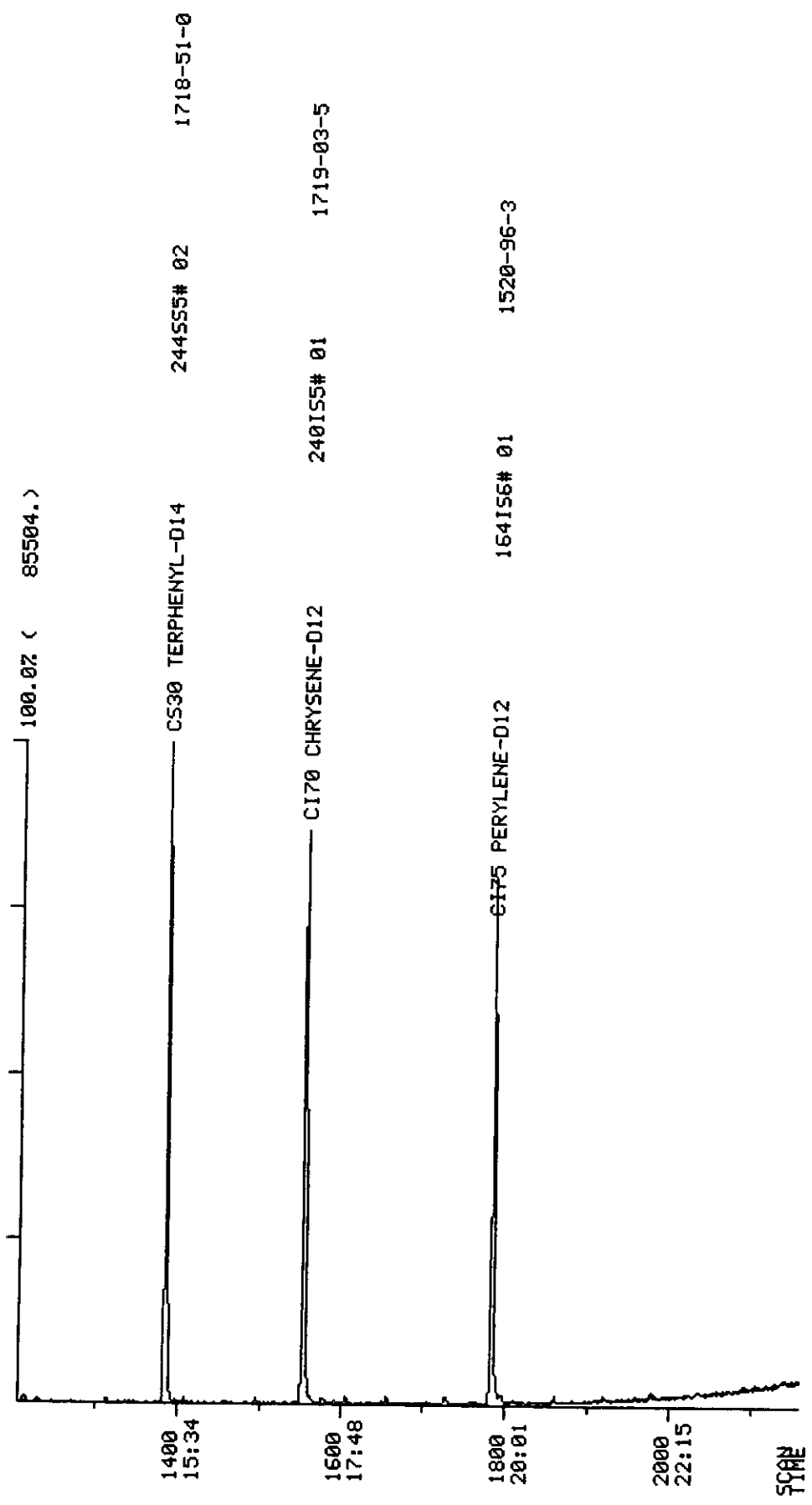
1712

DATA FROM FILE: 300681MBSFI SCANS 94 TO 1204 ACQUIRED: 09/02/98 16:29:00  
 CALI: 300681MBSFI #3  
 SAMPLE: SBLKSF 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA )/15A NA M



1713

DATA FROM FILE: 300681MBSFI SCANS 1204 TO 2158 ACQUIRED: 09/02/98 16:29:00  
 CALI: 300681MBSFI #3  
 SAMPLE: SBLKSF 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA )/15A NA M



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ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980902A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBSFI

Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 09/02/98 16:29

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML						
No Entry	Name	Mass	Meth	Scan	Ref	Area	RRF(L)	Amount
1 S1#	1 CI30 1,4-DICHLOROBENZENE	152	A	BB	340	1	24026.1	40.000
2 S2#	1 CI40 NAPHTHALENE-D8	136	A	BB	530	2	80168.1	40.000
3 S3#	1 CI50 ACENAPHTHENE-D10	164	A	BB	828	3	46995.1	40.000
4 S4#	1 CI60 PHENANTHRENE-D10	188	A	BB	1086	4	81567.1	40.000
5 S5#	1 CI70 CHRYSENE-D12	240	A	BB	1552	5	76304.1	40.000
6 S6#	1 CI75 PERYLENE-D12	264	A	BB	1783	6	67873.1	40.000
7 S2#	2 CS20 NITROBENZENE-D5	82	A	BB	422	2	25918.0	27.541
8 S3#	3 CS25 2-FLUOROBIPHENYL	172	A	BB	723	3	37757.1	26.037
9 S5#	2 CS30 TERPHENYL-D14	244	A	BB	1384	5	71990.0	37.920
10 S1#	3 CS45 PHENOL-D5	99	A	BB	308	1	20583.2	16.918
11 S1#	2 CS50 2-FLUOROPHENOL	112	A	BB	197	1	29385.1	35.755
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330	A	BB	968	3	20063.0	72.466
13 S1#	5 CS70 2-CHLOROPHENOL-D4	132						1 NOT FOUND
14 S1#	6 CS75 1,2-DICHLOROBENZENE-D4	152						1 NOT FOUND
15 S4#	4 HEXACHLOROBENZENE-C13	294						4 NOT FOUND
16 S1#	15 C310 N-NITROSODIMETHYLAM	74						1 NOT FOUND
17 S1#	20 PYRIDINE	79						1 NOT FOUND
18 S1#	30 2-PICOLINE	93						1 NOT FOUND
19 S1#	40 N-NITROSOMETHYLETHYLAMIN	42						1 NOT FOUND
20 S1#	60 METHYLMETHANESULFONATE	80						1 NOT FOUND
21 S1#	70 N-NITROSODIETHYLAMINE	102						1 NOT FOUND
22 S1#	85 ETHYLMETHANE SULFONATE	79						1 NOT FOUND
23 S1#	95 PENTACHLOROETHANE	117						1 NOT FOUND
24 S1#	100 C320 ANILINE	93						1 NOT FOUND
25 S1#	105 C315 PHENOL	94						1 NOT FOUND
26 S1#	110 C325 BIS(2-CHLOROETHYL)E	93						1 NOT FOUND
27 S1#	115 C330 2-CHLOROPHENOL	128						1 NOT FOUND
28 S1#	125 C335 1,3-DICHLOROBENZENE	146						1 NOT FOUND
29 S1#	130 C340 1,4-DICHLOROBENZENE	146						1 NOT FOUND
30 S1#	145 C345 BENZYL ALCOHOL	108						1 NOT FOUND
31 S1#	150 C350 1,2-DICHLOROBENZENE	146						1 NOT FOUND
32 S1#	160 C355 2-METHYLPHENOL	108						1 NOT FOUND
33 S1#	165 C360 2,2'-OXYBIS(1-CLPRO	45						1 NOT FOUND
34 S1#	170 C361 ACETOPHENONE	105						1 NOT FOUND
35 S1#	175 N-NITROSOPYRROLIDINE	100						1 NOT FOUND
36 S1#	180 N-NITROSOMORPHOLINE	56						1 NOT FOUND
37 S1#	182 3-METHYL PHENOL	108						1 NOT FOUND
38 S1#	185 C365 4-METHYLPHENOL	108						1 NOT FOUND
39 S1#	190 C370 N-NITROSO-DI-N-PROP	70						1 NOT FOUND
40 S1#	195 O-TOLUIDINE	106						1 NOT FOUND
41 S1#	200 C375 HEXACHLOROETHANE	117						1 NOT FOUND
42 S2#	10 C410 NITROBENZENE	77						2 NOT FOUND
43 S2#	15 N-NITROSOPIPERIDINE	42						2 NOT FOUND
44 S2#	20 C415 ISOPHORONE	82						2 NOT FOUND
45 S2#	25 C420 2-NITROPHENOL	139						2 NOT FOUND

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46	S2#	30	C425	2,4-DIMETHYLPHENOL	107	2	NOT FOUND
47	S2#	35	C435	BIS(2-CHLOROETHOXY)	93	2	NOT FOUND
48	S2#	40	C440	2,4-DICHLOROPHENOL	162	2	NOT FOUND
49	S2#	45	C430	BENZOIC ACID	122	2	NOT FOUND

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50 S2# 50	AA-DIMETHYLPHENETHYLAMIN	58	2	NOT FOUND
51 S2# 55	C445 1,2,4-TRICHLOROBENZ	180	2	NOT FOUND
52 S2# 60	C450 NAPHTHALENE	128	2	NOT FOUND
53 S2# 80	C455 4-CHLOROANALINE	127	2	NOT FOUND
54 S2# 85	2,6-DICHLOROPHENOL	162	2	NOT FOUND
55 S2# 90	HEXACHLOROPROPENE	213	2	NOT FOUND
56 S2# 95	C460 HEXACHLOROBUTADIENE	225	2	NOT FOUND
57 S2#115	P-PHENYLENE DIAMINE	108	2	NOT FOUND
58 S2#120	N-NITROSODI-N-BUTYLAMINE	84	2	NOT FOUND
59 S2#130	C465 4-CHLORO-3-METHYLPH	107	2	NOT FOUND
60 S2#140	SAFROLE	162	2	NOT FOUND
61 S2#145	C470 2-METHYLNAPHTHALENE	142	2	NOT FOUND
62 S3# 10	1,2,4,5-TETRACHLOROBENZE	214	3	NOT FOUND
63 S3# 15	ISOSAFROLE (#1)	162	3	NOT FOUND
64 S3# 20	C510 HEXACHLOROCYCLOPENT	237	3	NOT FOUND
65 S3# 25	C515 2,4,6-TRICHLOROPHEN	196	3	NOT FOUND
66 S3# 30	C520 2,4,5-TRICHLOROPHEN	196	3	NOT FOUND
67 S3# 35	ISOSAFROLE (#2)	104	3	NOT FOUND
68 S3# 40	C525 2-CHLORONAPHTHALENE	162	3	NOT FOUND
69 S3# 42	1-CHLORONAPHTHALEN	162	3	NOT FOUND
70 S3# 45	C530 2-NITROANALINE	65	3	NOT FOUND
71 S3# 50	1,4-NAPHTHOQUINONE	158	3	NOT FOUND
72 S3# 55	C535 DIMETHYLPHTHALATE	163	3	NOT FOUND
73 S3# 60	1,3-DINITROBENZENE	168	3	NOT FOUND
74 S3# 65	C540 ACENAPHTHYLENE	152	3	NOT FOUND
75 S3# 70	C543 2,6-DINITROTOLUENE	165	3	NOT FOUND
76 S3# 75	C545 3-NITROANILINE	138	3	NOT FOUND
77 S3# 80	C550 ACENAPHTHENE	153	3	NOT FOUND
78 S3# 85	C555 2,4-DINITROPHENOL	184	3	NOT FOUND
79 S3# 90	C565 DIBENZOFURAN	168	3	NOT FOUND
80 S3# 95	C560 4-NITROPHENOL	109	3	NOT FOUND
81 S3#100	PENTACHLOROBENZENE	250	3	NOT FOUND
82 S3#105	C570 2,4-DINITROTOLUENE	165	3	NOT FOUND
83 S3#110	1-NAPHTHYLAMINE	143	3	NOT FOUND
84 S3#115	2-NAPHTHYLAMINE	143	3	NOT FOUND
85 S3#120	2,3,4,6-TETRACHLOROPHENO	232	3	NOT FOUND
86 S3#130	C580 DIETHYLPHTHALATE	149	3	NOT FOUND
87 S3#135	C590 FLUORENE	166	3	NOT FOUND
88 S3#140	C585 4-CHLOROPHENYL-PHEN	204	3	NOT FOUND
89 S3#145	5-NITRO-O-TOLUIDINE	152	3	NOT FOUND
90 S3#150	C595 4-NITROANALINE	138	3	NOT FOUND
91 S4# 10	C610 4,6-DINITRO-2-METHY	198	4	NOT FOUND
92 S4# 15	C615 N-NITROSODIPHENYLAM	169	4	NOT FOUND
93 S4# 20	C620 AZOBENZENE	77	4	NOT FOUND
94 S4# 25	SYM-TRINITROBENZENE	75	4	NOT FOUND
95 S4# 30	C625 4-BROMOPHENYL-PHENY	248	4	NOT FOUND
96 S4# 35	PHENACETIN	108	4	NOT FOUND
97 S4# 37	DIALLATE	234	4	NOT FOUND
98 S4# 40	C630 HEXACHLOROBENZENE	284	4	NOT FOUND
99 S4# 45	4-AMINOBIIPHENYL	169	4	NOT FOUND
100 S4# 50	C635 PENTACHLOROPHENOL	266	4	NOT FOUND
101 S4# 55	PRONAMIDE	173	4	NOT FOUND
102 S4# 60	PENTACHLORONITROBENZENE	237	4	NOT FOUND
103 S4# 65	C640 PHENANTHRENE	178	4	NOT FOUND
104 S4# 70	C645 ANTHRACENE	178	4	NOT FOUND
105 S4# 75	2SECBTYL-4,6-DINITROPHE	211	4	NOT FOUND
106 S4# 80	C647 CARBAZOLE	167	4	NOT FOUND
107 S4# 85	C650 DI-N-BUTYLPHTHALATE	149	A BB 1227 4	732. 1.247 0.288
108 S4#100	4-NITROQUINOLINE-1-OXIDE	190	4	NOT FOUND
109 S4#105	METHAPYRILENE	58	4	NOT FOUND
110 S4#106	ISODRIN	193	4	NOT FOUND

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111	S4#110	C655	FLUORANTHENE	202	4	NOT FOUND			
112	S4#120		CHLOROBENZILATE	139	4	NOT FOUND			
113	S5# 10	C710	BENZIDINE	184	5	NOT FOUND			
114	S5# 15	C715	PYRENE	202	5	NOT FOUND			
115	S5# 20		ARAMITE (#1)	185	5	NOT FOUND			
116	S5# 25		ARAMITE (#2)	185	5	NOT FOUND			
117	S5# 30		P-DIMETHYLAMINOAZOBENZEN	120	5	NOT FOUND			
118	S5# 35		3,3'-DIMETHYLBENZIDINE	212	5	NOT FOUND			
119	S5# 37		KEPONE	272	5	NOT FOUND			
120	S5# 40	C720	BUTYLBENZYLPHTHALAT	149	5	NOT FOUND			
121	S5# 45		2-ACETYLAMINOFUORENE	181	5	NOT FOUND			
122	S5# 50	C730	BENZO(A)ANTHRACENE	228	5	NOT FOUND			
123	S5# 55	C725	3,3'-DICHLOROBENZID	252	5	NOT FOUND			
124	S5# 60	C740	CHRYSENE	228	5	NOT FOUND			
125	S5# 65	C745	BIS(2-ETHYLHEXYL)PH	149	A BB	1604	S	738.	0.980 0.395
126	S5# 85		3-METHYLCHOLANTHRENE	268	5	NOT FOUND			
127	S6# 10	C760	DI-N-OCTYL PHTHALAT	149	6	NOT FOUND			
128	S6# 15	C765	BENZO(B)FLUORANTHEN	252	6	NOT FOUND			
129	S6# 20		7,12-DIMETHYLBENZANTHRAC	256	6	NOT FOUND			
130	S6# 25	C770	BENZO(K)FLUORANTHEN	252	6	NOT FOUND			
131	S6# 30		HEXACHLOROPHENE	196	6	NOT FOUND			
132	S6# 35	C775	BENZO(A)PYRENE	252	6	NOT FOUND			
133	S6# 55	C780	INDENO(1,2,3-CD)PYR	276	6	NOT FOUND			
134	S6# 60	C785	DIBENZ(A,H)ANTHRACE	278	6	NOT FOUND			
135	S6# 65	C790	BENZO(G,H,I)PERYLEN	276	6	NOT FOUND			

1718



QUANTERRA Internal Standard Check

Standard Filename: ST16980902

Analyzed: 09/02/98 14:53

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980902	23319.		3.78	79260.		5.91	43605.		9.22
Upper Limit	46638.=+200		4.28	158519.=+200		6.41	87211.=+200		9.72
Lower Limit	11659.=+ 50		3.28	39630.=+ 50		5.41	21803.=+ 50		8.72
-----									
Filename									
=====									
1 300681MBSFI	24026.= 103		3.78	80168.= 101		5.90	46995.= 107		9.21

IS# 1 = CI30 1,4-DICHLOROBENZENE-D4  
 IS# 2 = CI40 NAPHTHALENE-D8  
 IS# 3 = CI50 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980902	76218.		12.09	51424.		17.27	44306.		19.84
Upper Limit	152436.=+200		12.59	102848.=+200		17.77	88612.=+200		20.34
Lower Limit	38109.=+ 50		11.59	25712.=+ 50		16.77	22153.=+ 50		19.34
-----									
Filename									
=====									
1 300681MBSFI	81567.= 107		12.08	76304.= 148		17.26	67873.= 153		19.83

IS# 4 = CI60 PHENANTHRENE-D10  
 IS# 5 = CI70 CHRYSENE-D12  
 IS# 6 = CI75 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

Data Reduced by: \_\_\_\_\_ Date: \_\_\_\_\_  
Data Reviewed by: \_\_\_\_\_ Date: \_\_\_\_\_

Data File: 300681MBSFI

QUANTERRA GC/MS TIC REPORT ( Part 1 )

Sample: SBLKSF 01AUG98-16A 1SA/1ML  
Analyst: DAT

INST. ID: F16  
Date Analyzed: 09/02/98 16:29  
Run Factor: 1.00

# SCAN	Concentration in Sample (UG/A)	CAS #
1 486	4.5	541-02-6
CYCLOPENTASILOXANE, DECAMETHYL-		

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2 665	6.0	540-97-6
CYCLOHEXASILOXANE, DODECAMETHYL-		

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1720

DATA FILE: 300681MBSF1

QUANTERRA GC/MS TIC REPORT ( Part 2 )

CONCENTRATION = AREA(TIC)\*CONC(IS)/AREA(IS)

#	FIT	PURITY	STD.	INT.	RT	RRT	AREA	HEIGHT	AMOUNT		
									AS ANALYZED	(UG/ML )	LIB
1	945	872	1		5:24	0.917	19664.	7045.	4.476	NB	51262.
2	901	836	1		7:23	1.255	26330.	9065.	5.993	NB	56707.

1721

Library Search Data: 300681MBSFI # 486 Base m/z: 73  
 09/02/98 16:29:00 + 5:24 Cali: 300681MBSFI # 3 RIC: 6464.  
 Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 530 matched at least 3 of the 16 largest peaks in the unknown

- Rank In. Name  
 1 51262 CYCLOPENTASILOXANE, DECAMETHYL-  
 2 51292 BENZOIC ACID, 2,6-BIS[(TRIMETHYLSILYL)OXY]-, TRIMETHYLSILYL ESTER  
 3 55755 3,4-DIHYDROXYMANDELIC ACID, ETHYL ESTER, TRI-TMS  
 4 51291 BENZOIC ACID, 2,5-BIS(TRIMETHYLSILOXY)-, TRIMETHYLSILYL ESTER  
 5 60158 N-(TRIFLUOROACETYL)-N,O,O',O''-TETRAKIS(TRIMETHYLSILYL)NOREPINEPHRI\*  
 6 58322 N-(TRIFLUOROACETYL)-O,O',O''-TRIS(TRIMETHYLSILYL)NOREPINEPHRINE  
 7 58803 N-(TRIFLUOROACETYL)-O,O',O''-TRIS(TRIMETHYLSILYL)EPINEPHRINE  
 8 60335 BENZENEETHANAMINE, N-[(PENTAFLUOROPHENYL)METHYLENE]-.BETA.,3,4-TRIS\*  
 9 49864 3,4-DIHYDROXYBENZYL ALCOHOL, TRIS(TRIMETHYLSILYL)-

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C10.H30.O5.S15	370	73	872	945	891
2	C16.H30.O4.S13	370	73	664	775	728
3	C19.H36.O5.S13	428	355	596	677	660
4	C16.H30.O4.S13	370	355	592	665	660
5	C22.H42.O4.N.F3.S14	553	73	589	749	621
6	C19.H34.O4.N.F3.S13	481	73	577	711	628
7	C20.H36.O4.N.F3.S13	495	73	561	710	616
8	C24.H34.O3.N.F5.S13	563	355	558	699	607
9	C16.H32.O3.S13	356	73	549	792	601

Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	541-02-6
2	---	---	---	---	3782-85-2
3	---	---	---	---	- -
4	---	---	---	---	3618-20-0
5	---	---	---	---	- -
6	---	---	---	---	- -
7	---	---	---	---	- -
8	---	---	---	---	55429-13-5
9	---	---	---	---	68595-79-9

1722

DATA: 300681MBSFI # 486  
CALI: 300681MBSFI # 3  
BASE M/Z: 73  
RIC: 6464.

DATA: 300681MBSFI # 486  
CALI: 300681MBSFI # 3  
INST. ID: F16

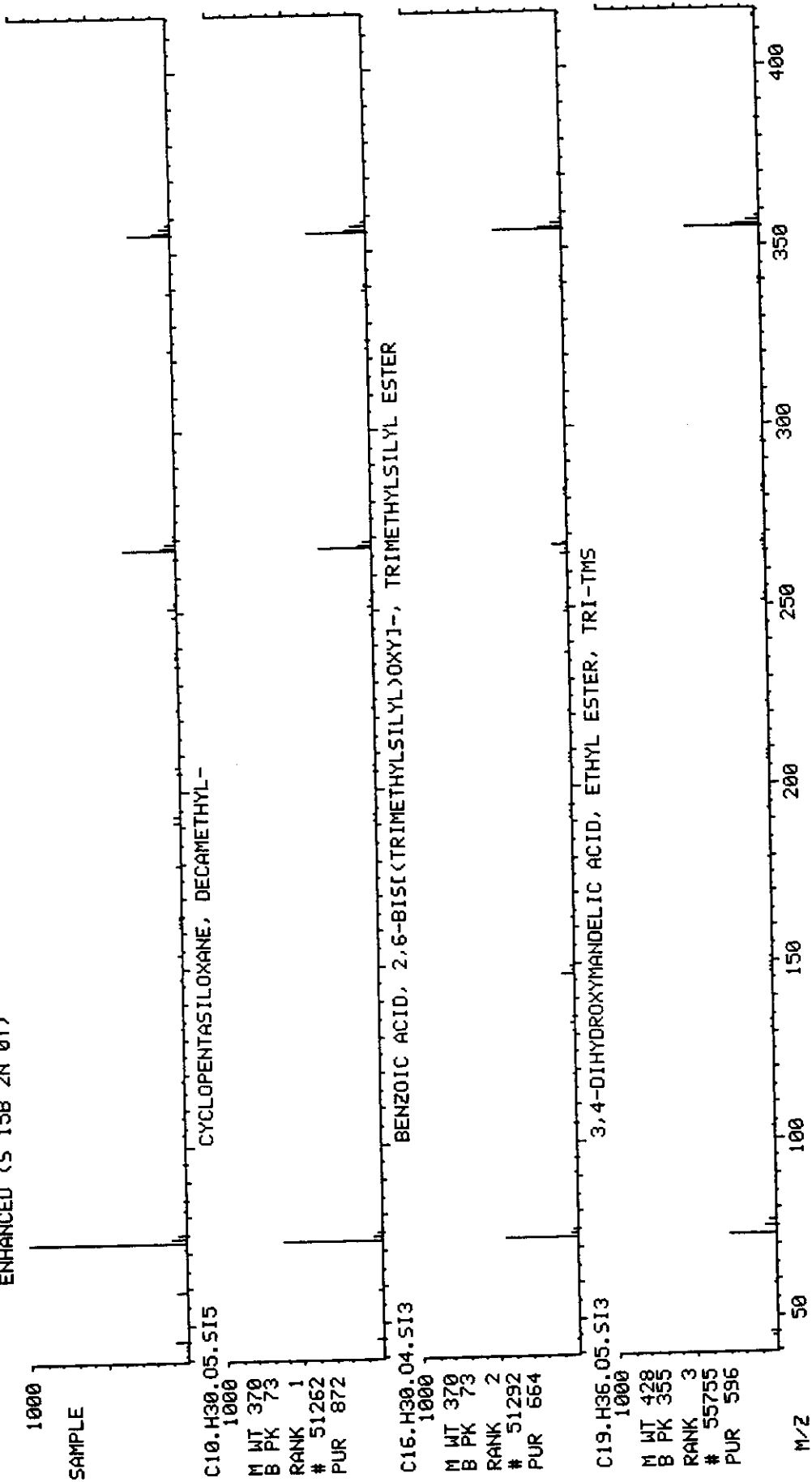
MID LIBRARY SEARCH (LIBRARYNB)

09/02/98 16:29:00 + 5:24

SAMPLE: SBLKSF 01AUG98-16A 15A/1ML

CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA) /15A NA M

ENHANCED (S 15B 2N 0T)



120  
10

Library Search Data: 300681MBSFI # 665 Base m/z: 73  
 09/02/98 16:29:00 + 7:24 Cali: 300681MBSFI # 3 RIC: 8352.  
 Sample: SBLKSF 01AUG98-16A 1SA/1ML INST. ID: F16  
 Conds.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1SA NA M  
 Enhanced (S 15B 2N 0T)

62231 spectra in LIBRARYNB searched for maximum PURITY  
 425 matched at least 3 of the 16 largest peaks in the unknown

Rank In. Name  
 1 56707 CYCLOHEXASILOXANE, DODECAMETHYL-  
 2 59692 1,1,1,5,7,7,7-HEPTAMETHYL-3,3,5-TRIS(TRIMETHYLSILOXY)TETRASILOXANE  
 3 58659 1,1,1,3,5,7,9,11,11,11-DECAMETHYL-5-(TRIMETHYLSILOXY)HEXASILOXANE  
 4 58080 3,3,5-TRIETHOXY-1,1,1,7,7,7-HEXAMETHYL-5-(TRIMETHYLSILOXY)TETRASI\*  
 5 48333 2H-1,4-BENZODIAZEPIN-2-ONE, 7-CHLORO-1,3-DIHYDRO-5-PHENYL-1-(TRIMET\*  
 6 56710 1,1,1,5,7,7,7-HEPTAMETHYL-3,3-BIS(TRIMETHYLSILOXY)TETRASILOXANE  
 7 49835 ACETIC ACID, [BIS[(TRIMETHYLSILYL)OXY]PHOSPHINYL]-, TRIMETHYLSILYL \*  
 8 60248 3,3,5-TRIBUTOXY-1,1,1,7,7,7-HEXAMETHYL-5-(TRIMETHYLSILOXY)TETRASILO\*  
 9 48319 1,3,5-TRIS(TRIMETHYLSILOXY)BENZENE

Rank	Formula	M.Wt	B.Pk	Purity	Fit	RFit
1	C12.H36.O6.SI6	444	73	836	901	855
2	C16.H48.O6.SI7	532	73	546	721	729
3	C13.H42.O6.SI7	490	73	471	678	650
4	C15.H42.O7.SI5	474	73	458	569	614
5	C18.H19.O.N2.CL.SI	342	73	447	776	517
6	C13.H40.O5.SI6	444	73	430	588	600
7	C11.H29.O5.P.SI3	356	341	418	569	660
8	C21.H54.O7.SI5	558	73	399	505	590
9	C15.H30.O3.SI3	342	342	372	540	435

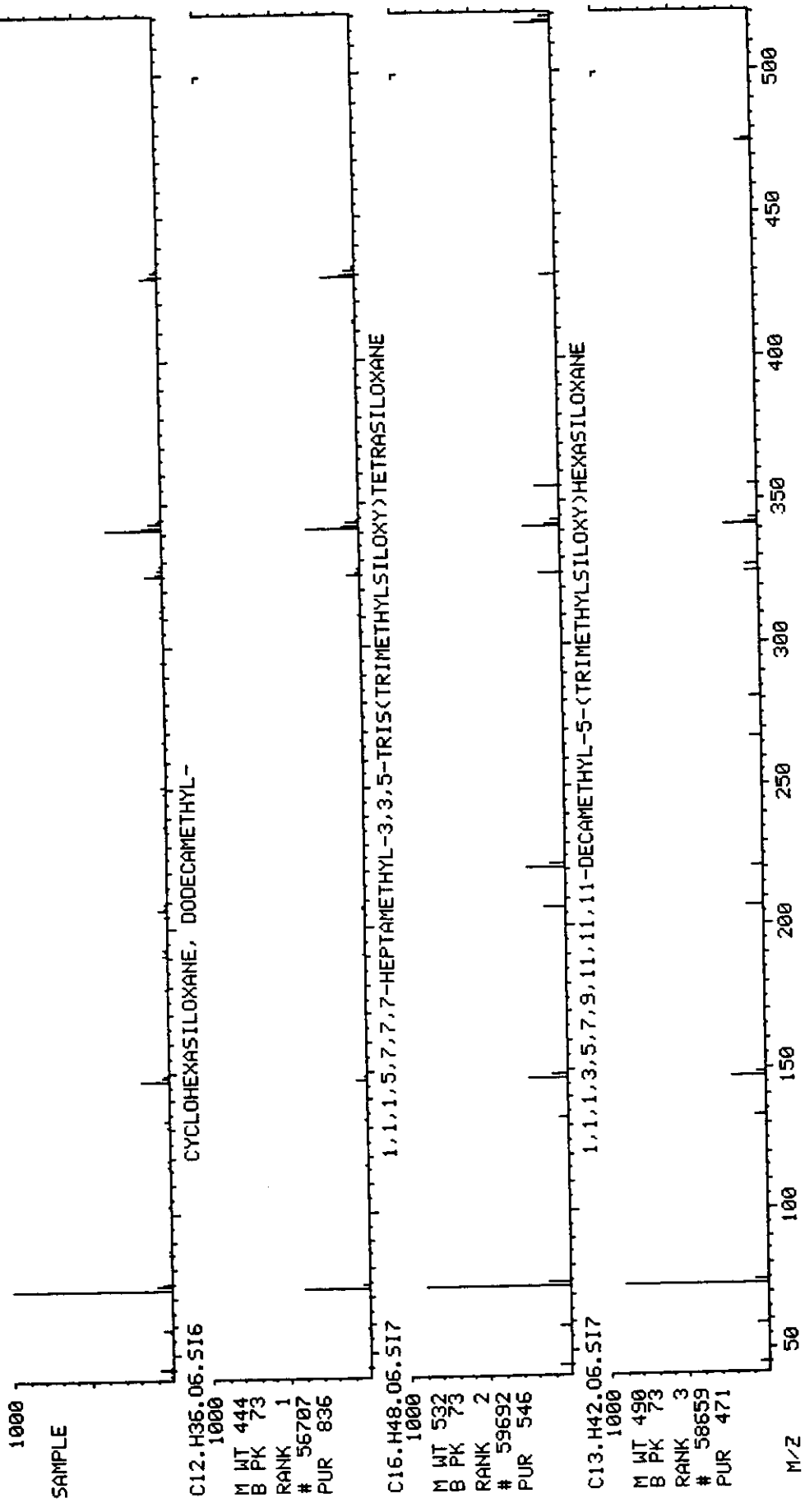
Rank	Ret.Time	B.P.Int.	US.Par.1	US.Par.2	C.A.S. #
1	---	---	---	---	540-97-6
2	---	---	---	---	34915-06-5
3	---	---	---	---	50694-26-3
4	---	---	---	---	-
5	---	---	---	---	55299-24-6
6	---	---	---	---	38147-00-1
7	---	---	---	---	53044-27-2
8	---	---	---	---	72439-86-2
9	---	---	---	---	10586-12-6

1724

MID LIBRARY SEARCH (LIBRARYNB)  
 09/02/98 16:29:00 + 7:24  
 SAMPLE: SBLKSF 01AUG98-16A 15A/1ML  
 CONDS.: UG/ML \*1ML \*100Z/100Z \*(NA/NA )/15A NA M  
 ENHANCED (S 15B 2N 0T)

DATA: 300681MBSFI # 665  
 CALI: 300681MBSFI # 3  
 INST. ID: F16

BASE M/Z: 73  
 RIC: 8352.



C12.H36.06.S16  
 1000  
 M WT 444  
 B PK 73  
 RANK 1  
 # 56707  
 PUR 836

C16.H48.06.S17  
 1000  
 M WT 532  
 B PK 73  
 RANK 2  
 # 59692  
 PUR 546

C13.H42.06.S17  
 1000  
 M WT 490  
 B PK 73  
 RANK 3  
 # 58659  
 PUR 471

M/Z  
 73  
 73  
 73

TIC SELECTION REPORT

DATA FILE: 300681MBSF1

THE FOLLOWING PEAKS WERE REJECTED BECAUSE AT LEAST 40 % OF THEIR SIZE WAS ACCOUNTED FOR BY TARGET COMPOUNDS ELUTING WITHIN 4 SCANS OF THE PEAK TOP.

SCAN	SIZE	AMOUNT
197	98212.	22.354
308	55052.	12.530
340	149548.	34.039
422	65244.	14.850
530	175738.	40.000
723	113212.	19.967
828	226798.	40.000
969	163679.	30.306
1086	216033.	40.000
1384	260393.	44.628
1384	260393.	44.628
1552	233390.	40.000
1784	229389.	40.000

TOTAL NUMBER OF UNIDENTIFIED PEAKS WITH SIZE GREATER THAN 10 % OF THE CLOSEST INTERNAL STANDARD THAT DOES NOT HAVE INTERFERENCES = 2

INTERNAL STANDARDS THAT HAVE RIC SIZE LESS THAN 50 % OR GREATER THAN 200 % OF THE ESTIMATED RIC SIZE ARE CONSIDERED TO HAVE INTERFERENCES AND WILL NOT BE USED FOR QUANTITATION.

#	INTERNAL STANDARD	PERCENT OF ESTIMATED		SCAN
		RIC SIZE	RIC SIZE	
1	C140 NAPHTHALENE-D8	175738.	137	530
2	C150 ACENAPHTHENE-D10	226798.	134	828
3	C160 PHENANTHRENE-D10	216033.	116	1086
4	C170 CHRYSENE-D12	233390.	124	1552
5	C175 PERYLENE-D12	229389.	126	1784

\* INDICATES INTERFERENCE

SIZE = AREA



Laboratory/Duplicate Control Sample

DUPLICATE CONTROL SAMPLE REPORT  
 Semivolatile Organics by GC/MS  
 Project: 300681

Category: 8270-G Acid, Base and Neutrals by GC/MS.  
 Testcode: 8270-TCL-G  
 Matrix: FILTER  
 QC Lot: 31 JUL 98-16A  
 Analyzed Date: 31 AUG 98  
 Method: 0010/8270  
 Concentration Units: ug/sample  
 Time: 18:30

Analyte	Spiked	-----Concentration-----		Accuracy (%)		Limits	Precision (RPD)	
		-----Measured-----		DCS1	DCS2		DCS Limit	DCS Limit
Phenol	100	DCS1	DCS2	DCS1	DCS2	47-108	5.3	18
2-Chlorophenol	100	83.4	87.9	83	88	47-113	4.9	20
1,4-Dichlorobenzene	50.0	83.7	87.9	84	88	42-114	3.4	22
N-Nitroso-di-		43.4	44.9	87	90			
n-propylamine	50.0	38.3	40.9	77	82	46-107	6.6	15
1,2,4-								
Trichlorobenzene	50.0	43.1	45.9	86	92	45-118	6.3	16
4-Chloro-3-								
methylphenol	100	78.6	85.8	79	86	55-118	8.8	13
Acenaphthene	50.0	46.6	50.2	93	100	54-119	7.4	10
4-Nitrophenol	100	78.8	89.6	79	90	43-166	13	17
2,4-Dinitrotoluene	50.0	45.3	49.0	91	98	59-113	7.8	10
Pentachlorophenol	100	83.4	93.6	83	94	59-128	12	10
Pyrene	50.0	48.0	50.7	96	101	45-140	5.5	11

Surrogate	Spiked	-----Concentration-----		Accuracy (%)		Limits
		-----Measured-----		DCS1	DCS2	
Nitrobenzene-d5	50	DCS1	DCS2	DCS1	DCS2	45-107
2-Fluorobiphenyl	50	40	40	79	79	62-110
Terphenyl-d14	50	45	46	90	92	58-135
Phenol-d5	100	40	40	79	80	43-130
2-Fluorophenol	100	80	81	80	81	36-111
2,4,6-Tribromophenol	100	77	74	77	74	58-131
		84	84	84	84	

Note \*: Outside of RPD limits.  
 Calculations are performed before rounding to avoid round-off errors in calculated results.

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBXMS  
Std Id: ST16980831

Sample: DCS-A 31JUL98-16A 1/3SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 08/31/98 18:00  
Run Factor: 3.00  
Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	QC limits
	Spiked	Measured		
CS20 NITROBENZENE-D5	50.00	39.70	79.4	45 107
CS25 2-FLUOROBIPHENY	50.00	44.80	89.6	62 110
CS30 TERPHENYL-D14	50.00	39.60	79.2	58 135
CS45 PHENOL-D5	100.0	80.00	80.0	43 130
CS50 2-FLUOROPHENOL	100.0	77.20	77.2	36 111
CS55 2,4,6-TRIBROMOP	100.0	83.90	83.9	58 131

Target Compounds: MS

Parameter	Scan	Result	Units	Reporting
				Limit (J values)
CS70 2-CHLOROPHENOL-D4		ND	UG/A	30.0
CS75 1,2-DICHLOROBENZ-D4	340	<del>130</del> 130	UG/A	30.0
C315 PHENOL	309	<del>250</del> 250	UG/A	30.0
C330 2-CHLOROPHENOL	314	<del>250</del> 250	UG/A	30.0
C340 1,4-DICHLOROBENZENE	342	<del>130</del> 130	UG/A	30.0
C370 N-NITROSO-DI-N-PROPYLAM	408	<del>110</del> 110	UG/A	30.0
C445 1,2,4-TRICHLOROBENZENE	524	130.	UG/A	30.0
C465 4-CHLORO-3-METHYLPHENO	649	240.	UG/A	30.0
C550 ACENAPHTHENE	833	140.	UG/A	30.0
C560 4-NITROPHENOL	885	240.	UG/A	150.0
C570 2,4-DINITROTOLUENE	878	140.	UG/A	30.0
C635 PENTACHLOROPHENOL	1061	250.	UG/A	150.0
C715 PYRENE	1334	140.	UG/A	30.0

See Quan report for adjusted amounts found.

2/17/98

29/17/98

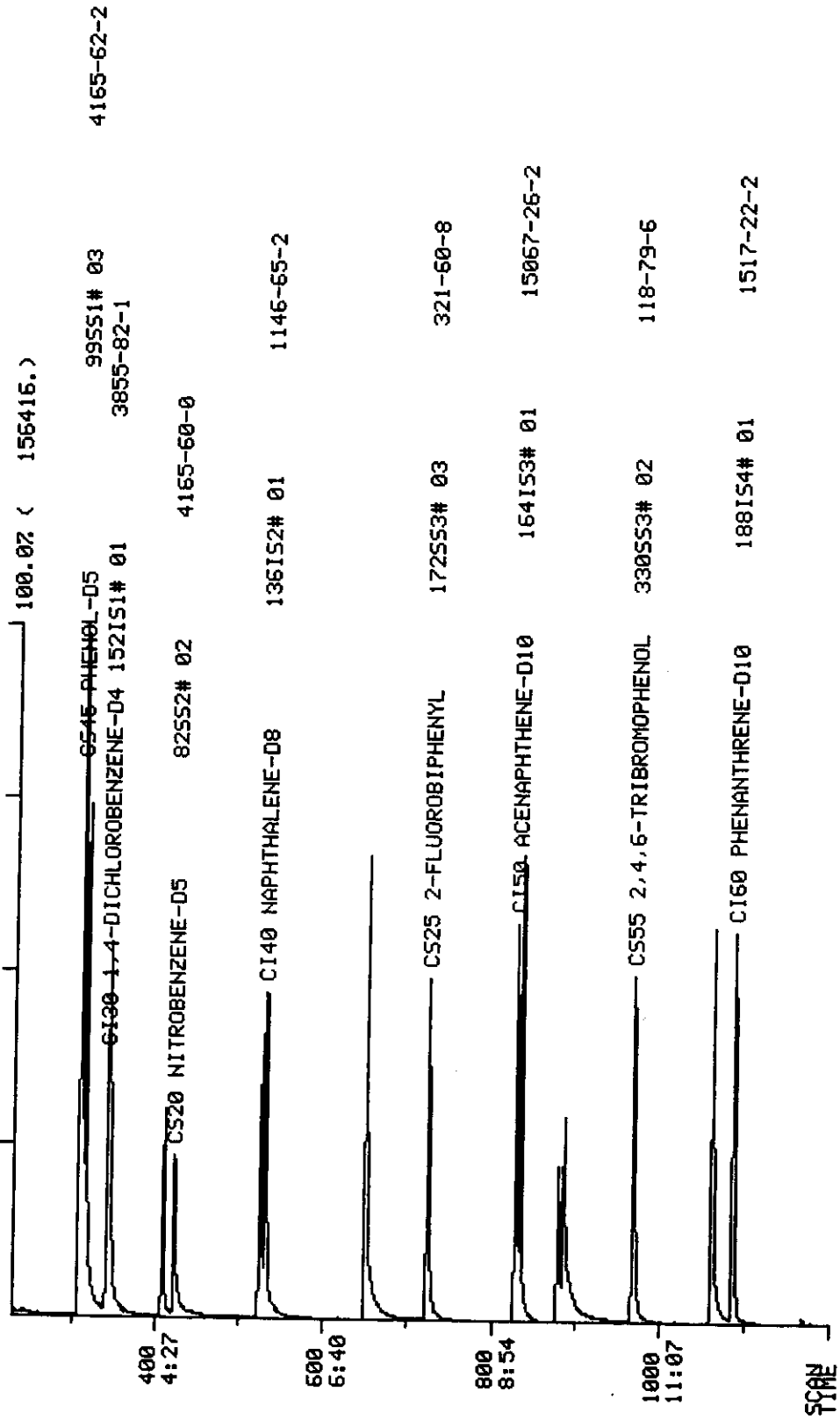
Reviewed by:

*[Handwritten signature]*

1789

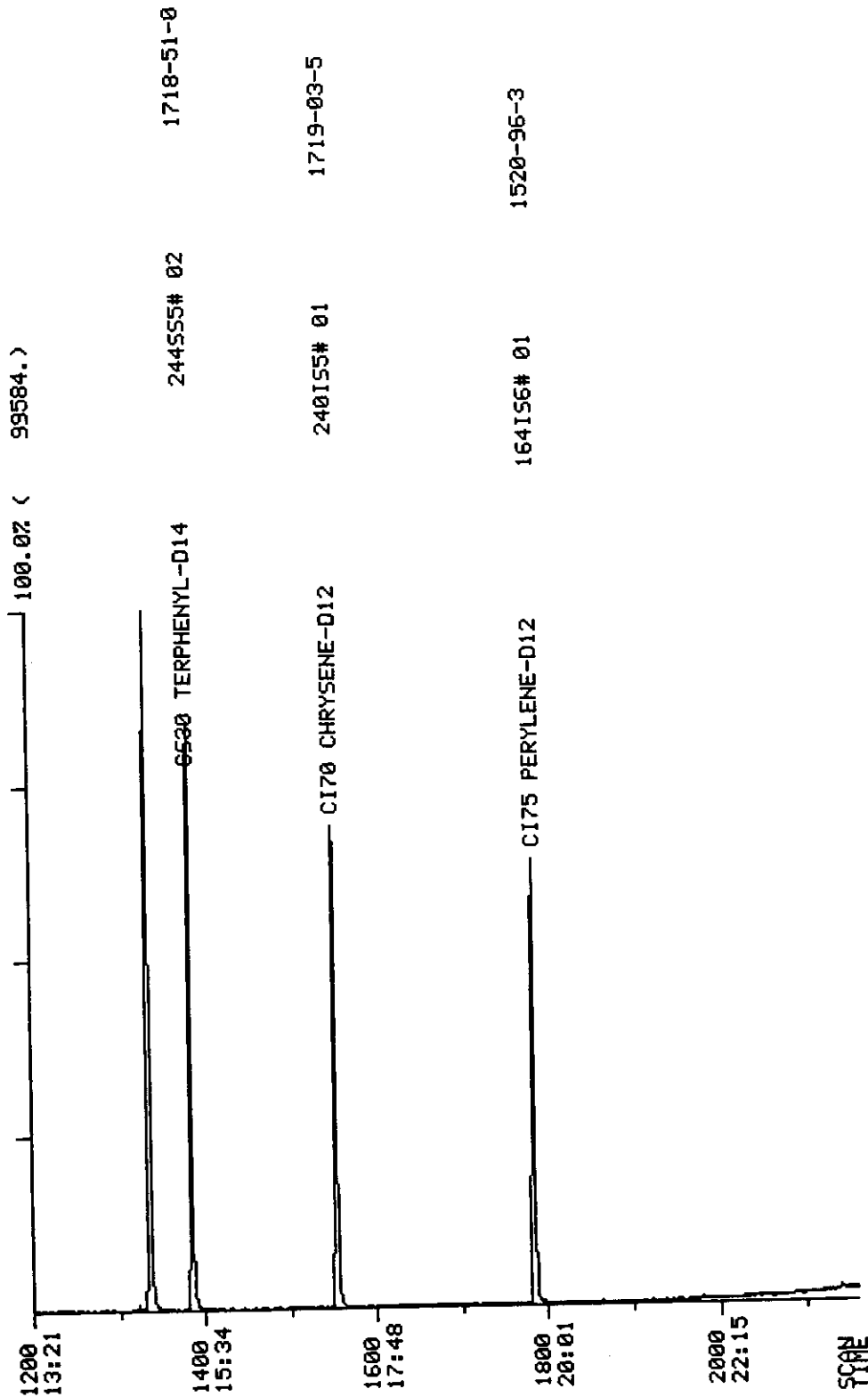
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DATA FROM FILE: 300681MBXMS SCANS 230 TO 1200 ACQUIRED: 08/31/98 18:00:00  
 CALI: 300681MBXMS #3  
 SAMPLE: DCS-A 31JUL98-16A 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M



177 000

DATA FROM FILE: 300681MBXMS SCANS 1200 TO 2158 ACQUIRED: 08/31/98 18:00:00  
 CALI: 300681MBXMS #3  
 SAMPLE: DCS-A 31JUL98-16A 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M



13:21  
 17:48  
 20:01

ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBXMS

Sample: DCS-A 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 18:00

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML					
No Entry	Name	Mass Meth	Scan Ref	Area	RRF(L)	Amount	
1 S1#	1 C130 1,4-DICHLOROBENZENE	152 A BB	340 1	27622.	1.000	40.000	
2 S2#	1 C140 NAPHTHALENE-DB	136 A BB	529 2	96564.	1.000	40.000	
3 S3#	1 C150 ACENAPHTHENE-D10	164 A BB	828 3	53847.	1.000	40.000	
4 S4#	1 C160 PHENANTHRENE-D10	188 A BB	1086 4	91222.	1.000	40.000	
5 S5#	1 C170 CHRYSENE-D12	240 A BB	1551 5	71648.	1.000	40.000	
6 S6#	1 C175 PERYLENE-D12	264 A BB	1783 6	59838.	1.000	40.000	
7 S2#	2 CS20 NITROBENZENE-D5	82 A BB	421 2	45031.	0.470	39.726	
8 S3#	3 CS25 2-FLUOROBIPHENYL	172 A BB	723 3	74360.	1.234	44.752	
9 S5#	2 CS30 TERPHENYL-D14	244 A BB	1383 5	70651.	0.995	39.633	
10 S1#	3 CS45 PHENOL-D5	99 A BB	307 1	111877.	2.025	79.986	
11 S1#	2 CS50 2-FLUOROPHENOL	112 A BB	196 1	72896.	1.368	77.152	
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330 A BB	968 3	26620.	0.236	83.911	
13 S1#	5 CS70 2-CHLOROPHENOL-D4	132	1 NOT FOUND				
14 S1#	6 CS75 1,2-DICHLOROBENZ-N-D4	152 A BB	340 1	27622.	0.920	43.476	
15 S1#105	C315 PHENOL	94 A BB	309 1	113308.	1.968	83.383	
16 S1#115	C330 2-CHLOROPHENOL	128 A BB	314 1	86494.	1.497	83.677	
17 S1#130	C340 1,4-DICHLOROBENZENE	146 A BB	342 1	48430.	1.617	43.359	
18 S1#190	C370 N-NITROSO-DI-N-PROP	70 A BB	408 1	23952.	0.906	38.290	
19 S2# 55	C445 1,2,4-TRICHLOROBENZ	180 A BB	524 2	34453.	0.331	43.126	
20 S2#130	C465 4-CHLORO-3-METHYLPH	107 A BB	649 2	64106.	0.338	78.625	
21 S3# 80	C550 ACENAPHTHENE	153 A BB	833 3	72613.	1.159	46.555	
22 S3# 95	C560 4-NITROPHENOL	109 A BB	885 3	18323.	0.173	78.836	
23 S3#105	C570 2,4-DINITROTOLUENE	165 A BB	878 3	24212.	0.397	45.277	
24 S4# 50	C635 PENTACHLOROPHENOL	266 A BB	1061 4	29475.	0.155	83.363	
25 S5# 15	C715 PYRENE	202 A BB	1334 5	113602.	1.323	47.954	

QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 300681MBXMS	27622.= 96		3.78	96564.= 96		5.88	53847.= 99		9.21

IS# 1 = C130 1,4-DICHLOROBENZENE-D4

IS# 2 = C140 NAPHTHALENE-D8

IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 300681MBXMS	91222.= 92		12.08	71648.= 115		17.25	59838.= 117		19.83

IS# 4 = C160 PHENANTHRENE-D10

IS# 5 = C170 CHRYSENE-D12

IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBXDS

Std Id: ST16980831

Sample: DCS-AD 31JUL98-16A 1/3SA/1ML

INST. ID: F16

Client: PACIFI

Date Analyzed: 08/31/98 18:30

Analyst: DAT Instrument ID: F16

Run Factor: 3.00

Quan List Threshold: 0.95

Surrogate Vol.: 1.00

Surrogate Spike Recoveries

8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	39.50	79.0	45 107
CS25 2-FLUOROBIPHENY	50.00	46.00	92.0	62 110
CS30 TERPHENYL-D14	50.00	40.20	80.4	58 135
CS45 PHENOL-D5	100.0	80.60	80.6	43 130
CS50 2-FLUOROPHENOL	100.0	74.20	74.2	36 111
CS55 2,4,6-TRIBROMOP	100.0	83.90	83.9	58 131

Target Compounds: MS

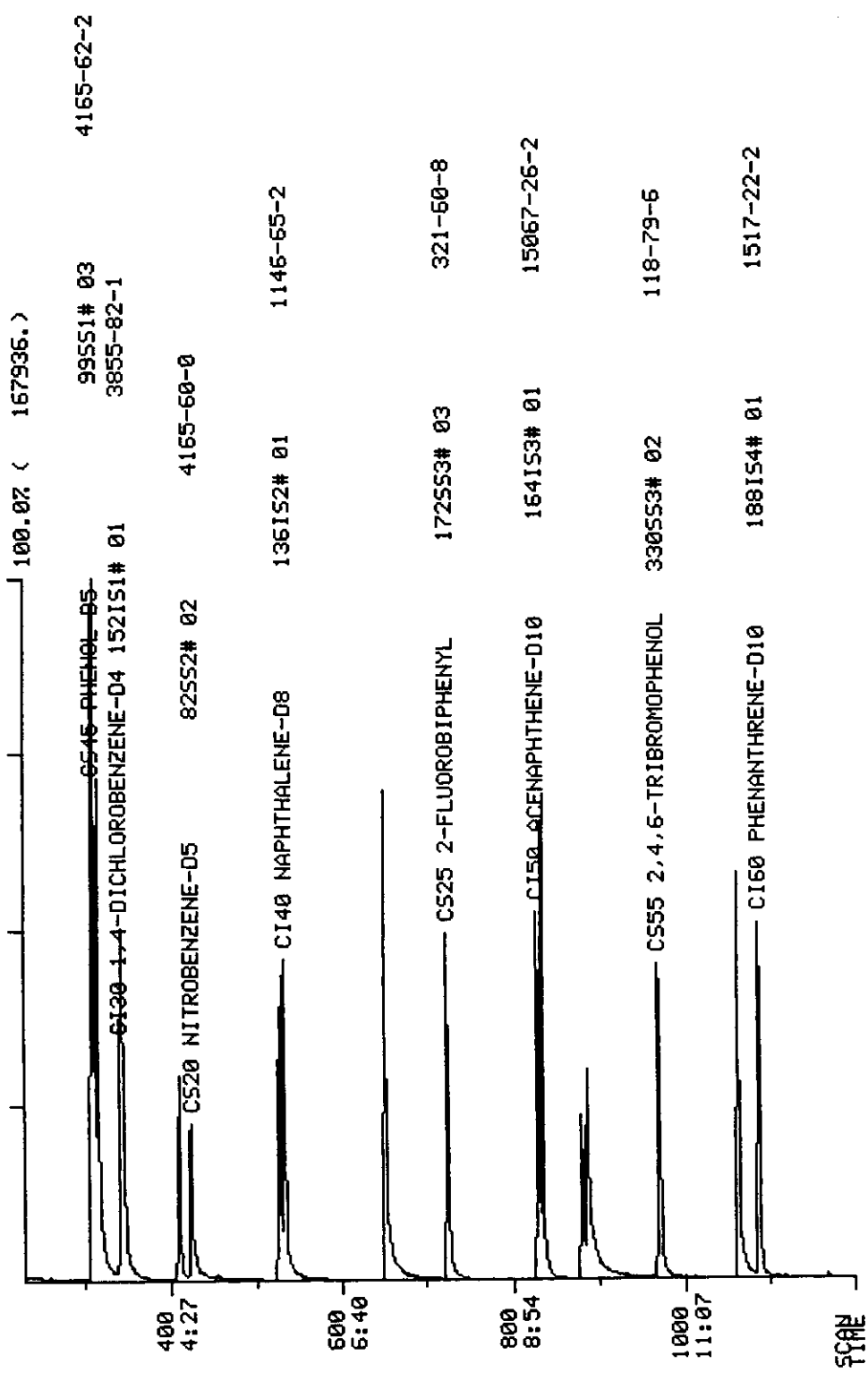
Parameter	Scan	Result	Units	Reporting
				Limit (J values)
CS70 2-CHLOROPHENOL-D4		ND	UG/A	30.0
CS75 1,2-DICHLOROBNZN-D4	340	130.	UG/A	30.0
C315 PHENOL	309	260.	UG/A	30.0
C330 2-CHLOROPHENOL	314	260.	UG/A	30.0
C340 1,4-DICHLOROBENZENE	343	130.	UG/A	30.0
C370 N-NITROSO-DI-N-PROPYLAM	408	120.	UG/A	30.0
C445 1,2,4-TRICHLOROBENZENE	524	140.	UG/A	30.0
C465 4-CHLORO-3-METHYLPHENO	649	260.	UG/A	30.0
C550 ACENAPHTHENE	834	150.	UG/A	30.0
C560 4-NITROPHENOL	885	270.	UG/A	150.0
C570 2,4-DINITROTOLUENE	878	150.	UG/A	30.0
C635 PENTACHLOROPHENOL	1061	280.	UG/A	150.0
C715 PYRENE	1334	150.	UG/A	30.0

Reviewed by:

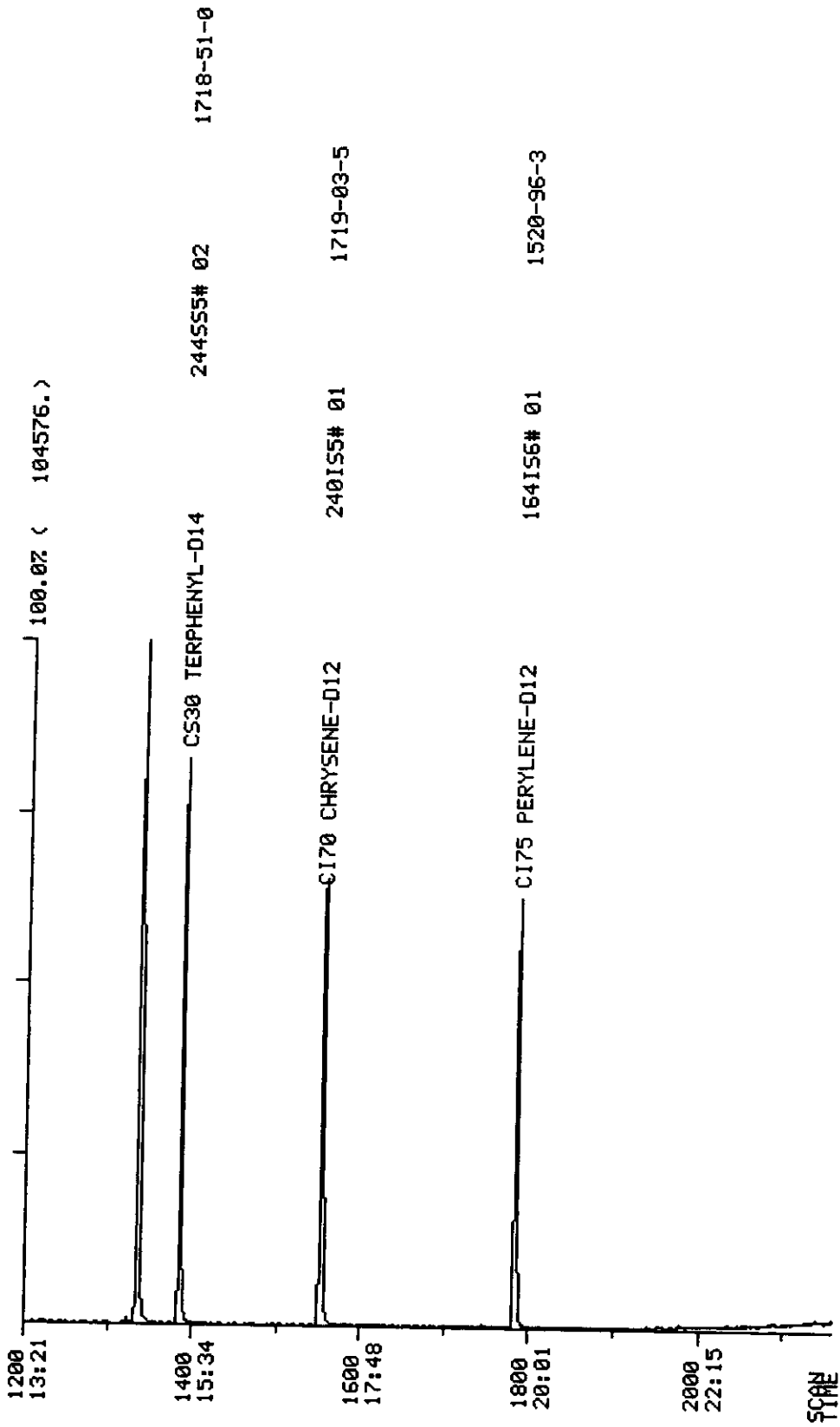
*K/9/17/98*



DATA FROM FILE: 300681MBXD5 SCANS 230 TO 1200 ACQUIRED: 08/31/98 18:30:00  
 CALI: 300681MBXD5 #3  
 SAMPLE: DC5-AD 31JUL98-16A 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100Z/100Z \*(NA/NA )/1/35A NA M



DATA FROM FILE: 300681MBXD5 SCANS 1200 TO 2158 ACQUIRED: 08/31/98 18:30:00  
 CALI: 300681MBXD5 #3  
 SAMPLE: DCS-AD 31JUL98-16A 1/35A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/1/35A NA M



ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBXDS

Sample: DCS-AD 31JUL98-16A 1/3SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 18:30

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML				
No Entry	Name	Mass Meth	Scan Ref	Area RRF(L)	Amount	
1 S1#	1 C130 1,4-DICHLOROBENZENE	152 A BB	340 1	27374. 1.000	40.000	
2 S2#	1 C140 NAPHTHALENE-D8	136 A BB	530 2	96718. 1.000	40.000	
3 S3#	1 C150 ACENAPHTHENE-D10	164 A BB	828 3	53018. 1.000	40.000	
4 S4#	1 C160 PHENANTHRENE-D10	188 A BB	1086 4	87510. 1.000	40.000	
5 S5#	1 C170 CHRYSENE-D12	240 A BB	1551 5	72710. 1.000	40.000	
6 S6#	1 C175 PERYLENE-D12	264 A BB	1783 6	60866. 1.000	40.000	
7 S2#	2 CS20 NITROBENZENE-D5	82 A BB	422 2	44793. 0.470	39.452	
8 S3#	3 CS25 2-FLUOROBIPHENYL	172 A BB	723 3	75190. 1.234	45.960	
9 S5#	2 CS30 TERPHENYL-D14	244 A BB	1384 5	72644. 0.995	40.156	
10 S1#	3 CS45 PHENOL-D5	99 A BB	307 1	111689. 2.025	80.575	
11 S1#	2 CS50 2-FLUOROPHENOL	112 A BB	196 1	69505. 1.368	74.230	
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330 A BB	968 3	26207. 0.236	83.902	
13 S1#	5 CS70 2-CHLOROPHENOL-D4	132	1 NOT FOUND			
14 S1#	6 CS75 1,2-DICHLOROBENZ-N-D4	152 A BB	340 1	27374. 0.920	43.476	
15 S1#105	C315 PHENOL	94 A BB	309 1	118368. 1.968	87.896	
16 S1#115	C330 2-CHLOROPHENOL	128 A BB	314 1	90015. 1.497	87.873	
17 S1#130	C340 1,4-DICHLOROBENZENE	146 A BB	343 1	49710. 1.617	44.908	
18 S1#190	C370 N-NITROSO-DI-N-PROP	70 A BB	408 1	25370. 0.906	40.924	
19 S2#	55 C445 1,2,4-TRICHLOROBENZ	180 A BB	524 2	36709. 0.331	45.877	
20 S2#130	C465 4-CHLORO-3-METHYLPH	107 A BB	649 2	70056. 0.338	85.785	
21 S3#	80 C550 ACENAPHTHENE	153 A BB	834 3	77035. 1.159	50.163	
22 S3#	95 C560 4-NITROPHENOL	109 A BB	885 3	20493. 0.173	89.552	
23 S3#105	C570 2,4-DINITROTOLUENE	165 A BB	878 3	25792. 0.397	48.986	
24 S4#	50 C635 PENTACHLOROPHENOL	266 A BB	1061 4	31741. 0.155	93.581	
25 S5#	15 C715 PYRENE	202 A BB	1334 5	121860. 1.323	50.688	

QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 300681MBXDS	27374.=	95	3.78	96718.=	96	5.90	53018.=	97	9.21

IS# 1 = C130 1,4-DICHLOROBENZENE-D4

IS# 2 = C140 NAPHTHALENE-D8

IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 300681MBXDS	87510.=	88	12.08	72710.=	116	17.25	60866.=	119	19.83

IS# 4 = C160 PHENANTHRENE-D10

IS# 5 = C170 CHRYSENE-D12

IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

QC Lot #: 01AUG98-16A

Units: UG/A

QUANTERRA DCS

QC CATEGORY: 8270-G, LIMS 10/10/96

	Data File	Run Factor	Surr Factor	Calibration Standard
Spike:	300681BFMSI	1.00	1.00	RF from Lib
Spike Dup.:	300681BFDSI	1.00	1.00	RF from Lib

Compound	Limits	Spike Amount	Amount LCS1	%	RPD Limit
			Amount LCS2	%	
C315 PHENOL		100.	27.2	27.2*	4.16
	47-108	100.	26.0	26.0*	18
C330 2-CHLOROPHENOL		100.	64.9	64.9	0.129
	47-113	100.	65.0	65.0	20
C340 1,4-DICHLOROBENZENE		50.0	27.3	54.7	6.96
	42-114	50.0	29.3	58.6	22
C370 N-NITROSO-DI-N-PROP		50.0	35.6	71.1	11.1
	46-107	50.0	31.8	63.6	15
C445 1,2,4-TRICHLOROBENZ		50.0	27.7	55.4	7.59
	45-118	50.0	29.9	59.8	16
C465 4-CHLORO-3-METHYLPH		100.	68.9	68.9	10.0
	55-118	100.	62.3	62.3	13
C550 ACENAPHTHENE		50.0	39.9	79.9	6.04
	54-119	50.0	37.6	75.2	10
C560 4-NITROPHENOL		100.	20.5	20.5*	22.1 *
	43-166	100.	16.4	16.4*	17
C570 2,4-DINITROTOLUENE		50.0	46.4	92.8	16.1 *
	59-113	50.0	39.5	78.9	10
C635 PENTACHLOROPHENOL		100.	77.0	77.0	18.3 *
	59-128	100.	64.1	64.1	10
C715 PYRENE		50.0	48.5	96.9	10.3
	45-140	50.0	43.7	87.4	11

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681BFMSI  
Std Id: ST16980902

Sample: DCS-A 01AUG98-16A 1SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 09/02/98 18:59  
Run Factor: 1.00  
Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	31.00	62.0	45 107
CS25 2-FLUOROBIPHENY	50.00	35.80	71.6	62 110
CS30 TERPHENYL-D14	50.00	41.00	82.0	58 135
CS45 PHENOL-D5	100.0	27.00	27.0	* 43 130
CS50 2-FLUOROPHENOL	100.0	37.60	37.6	36 111
CS55 2,4,6-TRIBROMOP	100.0	88.00	88.0	58 131

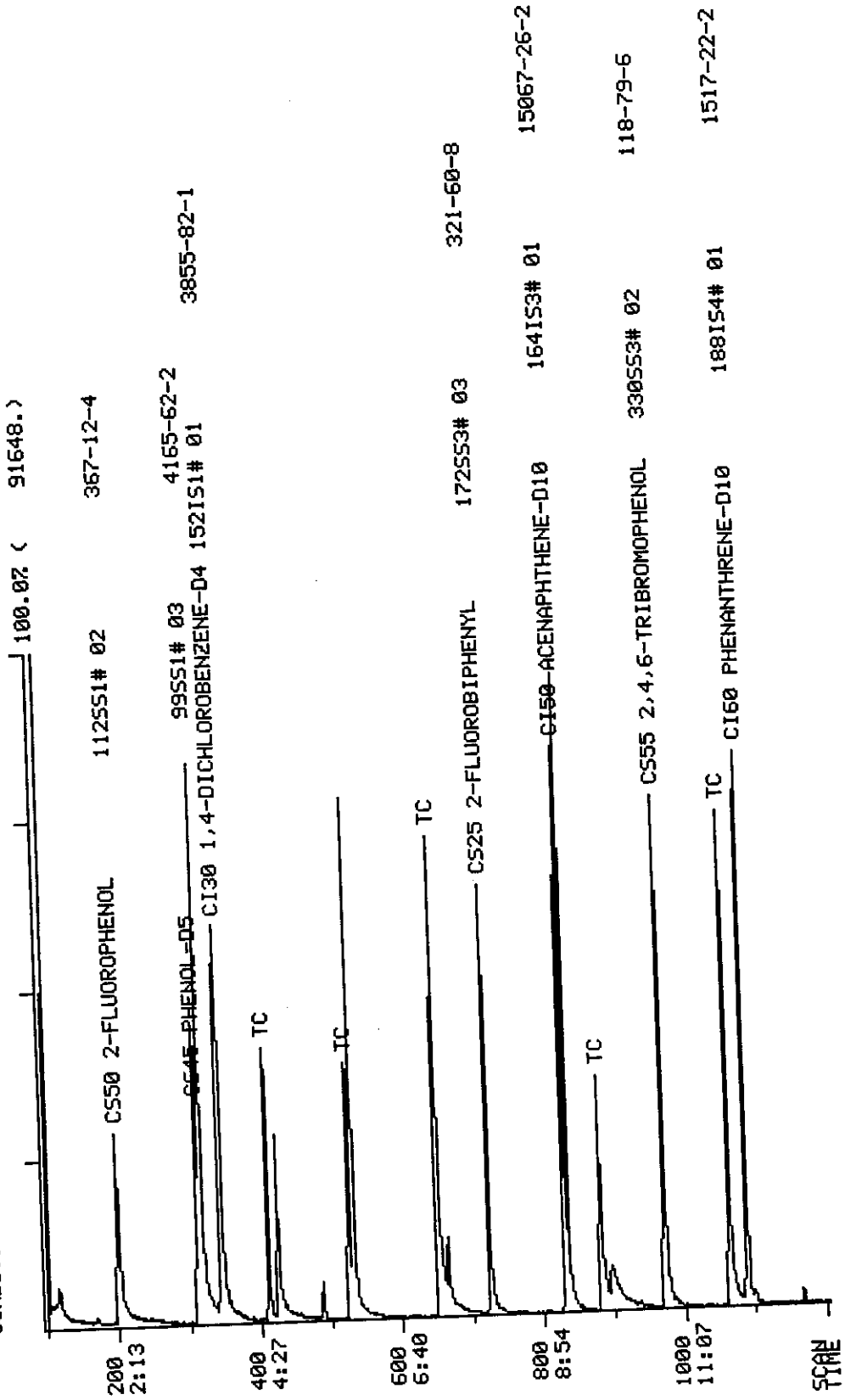
*confrms*

Target Compounds: MS

Parameter	Scan	Result	Units	Reporting Limit (J values)
CS70 2-CHLOROPHENOL-D4		ND	UG/A	10.0
CS75 1,2-DICHLOROBZN-D4		ND	UG/A	10.0
C315 PHENOL	310	27.	UG/A	10.0
C330 2-CHLOROPHENOL	314	65.	UG/A	10.0
C340 1,4-DICHLOROBENZENE	343	27.	UG/A	10.0
C370 N-NITROSO-DI-N-PROPYLAM	408	36.	UG/A	10.0
C445 1,2,4-TRICHLOROBENZENE	524	28.	UG/A	10.0
C465 4-CHLORO-3-METHYLPHENO	650	69.	UG/A	10.0
C550 ACENAPHTHENE	834	40.	UG/A	10.0
C560 4-NITROPHENOL	893	21.	UG/A	50.0 J
C570 2,4-DINITROTOLUENE	879	46.	UG/A	10.0
C635 PENTACHLOROPHENOL	1062	77.	UG/A	50.0
C715 PYRENE	1335	48.	UG/A	10.0

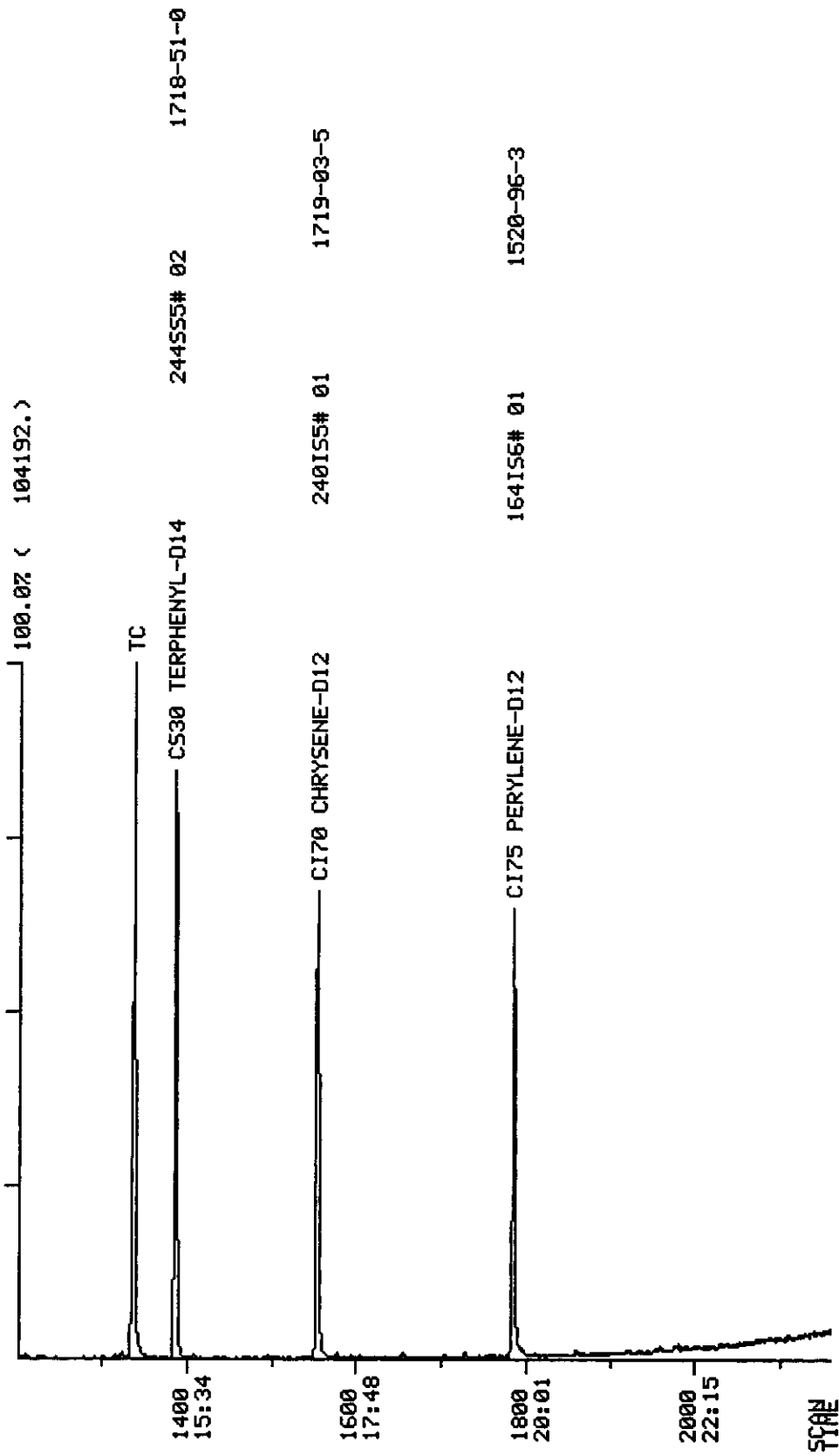
Reviewed by: *[Signature]*

DATA FROM FILE: 300681BFMSI SCANS 94 TO 1204 ACQUIRED: 09/02/98 18:59:00  
 CALI: 300681BFMSI #3 INST. ID: F16  
 SAMPLE: DCS-A 01AUG98-16A 15A/1ML  
 CONDS.: UG/ML \*100%/100% \*(NA/NA >)/15A NA M



1741

DATA FROM FILE: 300681BFMSI SCANS 1204 TO 2158 ACQUIRED: 09/02/98 18:59:00  
CALI: 300681BFMSI #3  
SAMPLE: DCS-A 01AUG98-16A 15A/1ML INST. ID: F16  
CONDS.: UG/ML \*100Z/100Z \*(NA/NA >/15A NA M



1742



ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980902A

## QUANTERRA QUANTITATION SUMMARY

File: 300681BFMSI

Sample: DCS-A 01AUG98-16A 1SA/1ML

INST. ID: F16

Analyst: DAT

Instrument Id: F16

Analyzed: 09/02/98 18:59

Compounds with amounts less than 0.20 reported as NOT FOUND

Library									Units: UG/ML
No Entry	Name	Mass	Meth	Scan	Ref	Area	RRF (L)	Amount	
1 S1#	1 CI30 1,4-DICHLOROBENZENE	152	A BB	341	1	24660.	1.000	40.000	
2 S2#	1 CI40 NAPHTHALENE-D8	136	A BB	530	2	87031.	1.000	40.000	
3 S3#	1 CI50 ACENAPHTHENE-D10	164	A BB	828	3	48399.	1.000	40.000	
4 S4#	1 CI60 PHENANTHRENE-D10	188	A BB	1087	4	83148.	1.000	40.000	
5 S5#	1 CI70 CHRYSENE-D12	240	A BB	1552	5	75605.	1.000	40.000	
6 S6#	1 CI75 PERYLENE-D12	264	A BB	1784	6	66009.	1.000	40.000	
7 S2#	2 CS20 NITROBENZENE-D5	82	A BB	422	2	31682.	0.470	31.011	
8 S3#	3 CS25 2-FLUOROBIPHENYL	172	A BB	723	3	53531.	1.234	35.844	
9 S5#	2 CS30 TERPHENYL-D14	244	A BB	1384	5	77124.	0.995	41.000	
10 S1#	3 CS45 PHENOL-D5	99	A BB	308	1	33750.	2.025	27.028	
11 S1#	2 CS50 2-FLUOROPHENOL	112	A BB	197	1	31682.	1.368	37.559	
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330	A BB	969	3	25098.	0.236	88.020	
13 S1#	5 CS70 2-CHLOROPHENOL-D4	132			1	NOT FOUND			
14 S1#	6 CS75 1,2-DICHLOROBENZEN-D4	152			1	NOT FOUND			
15 S1#105	C315 PHENOL	94	A BB	310	1	32941.	1.968	27.153	
16 S1#115	C330 2-CHLOROPHENOL	128	A BB	314	1	59889.	1.497	64.898	
17 S1#130	C340 1,4-DICHLOROBENZENE	146	A BB	343	1	27254.	1.617	27.331	
18 S1#190	C370 N-NITROSO-DI-N-PROP	70	A BB	408	1	19856.	0.906	35.554	
19 S2# 55	C445 1,2,4-TRICHLOROBENZ	180	A BB	524	2	19952.	0.331	27.711	
20 S2#130	C465 4-CHLORO-3-METHYLPH	107	A BB	650	2	50638.	0.338	68.909	
21 S3# 80	C550 ACENAPHTHENE	153	A BB	834	3	56003.	1.159	39.948	
22 S3# 95	C560 4-NITROPHENOL	109	A BB	893	3	4286.	0.173	20.517	
23 S3#105	C570 2,4-DINITROTOLUENE	165	A BB	879	3	22297.	0.397	46.389	
24 S4# 50	C635 PENTACHLOROPHENOL	266	A BB	1062	4	24829.	0.155	77.041	
25 S5# 15	C715 PYRENE	202	A BB	1335	5	121134.	1.323	48.457	

QUANTERRA Internal Standard Check

Standard Filename: ST16980902

Analyzed: 09/02/98 14:53

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980902	23319.		3.78	79260.		5.91	43605.		9.22
Upper Limit	46638.=+200		4.28	158519.=+200		6.41	87211.=+200		9.72
Lower Limit	11659.=+ 50		3.28	39630.=+ 50		5.41	21803.=+ 50		8.72
Filename									
1 300681BFMSI	24660.= 105		3.79	87031.= 109		5.90	48399.= 110		9.21

IS# 1 = CI30 1,4-DICHLORO BENZENE-D4  
 IS# 2 = CI40 NAPHTHALENE-D8  
 IS# 3 = CI50 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980902	76218.		12.09	51424.		17.27	44306.		19.84
Upper Limit	152436.=+200		12.59	102848.=+200		17.77	88612.=+200		20.34
Lower Limit	38109.=+ 50		11.59	25712.=+ 50		16.77	22153.=+ 50		19.34
Filename									
1 300681BFMSI	83148.= 109		12.09	75605.= 147		17.26	66009.= 148		19.84

IS# 4 = CI60 PHENANTHRENE-D10  
 IS# 5 = CI70 CHRYSENE-D12  
 IS# 6 = CI75 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681BFDSI  
Std Id: ST16980902

Sample: DCS-AD 01AUG98-16A 1SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 09/02/98 19:29  
Run Factor: 1.00  
Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	30.20	60.4	45 107
CS25 2-FLUOROBIPHENY	50.00	33.80	67.6	62 110
CS30 TERPHENYL-D14	50.00	36.50	73.0	58 135
CS45 PHENOL-D5	100.0	26.00	26.0	* 43 130
CS50 2-FLUOROPHENOL	100.0	38.10	38.1	36 111
CS55 2,4,6-TRIBROMOP	100.0	74.60	74.6	58 131

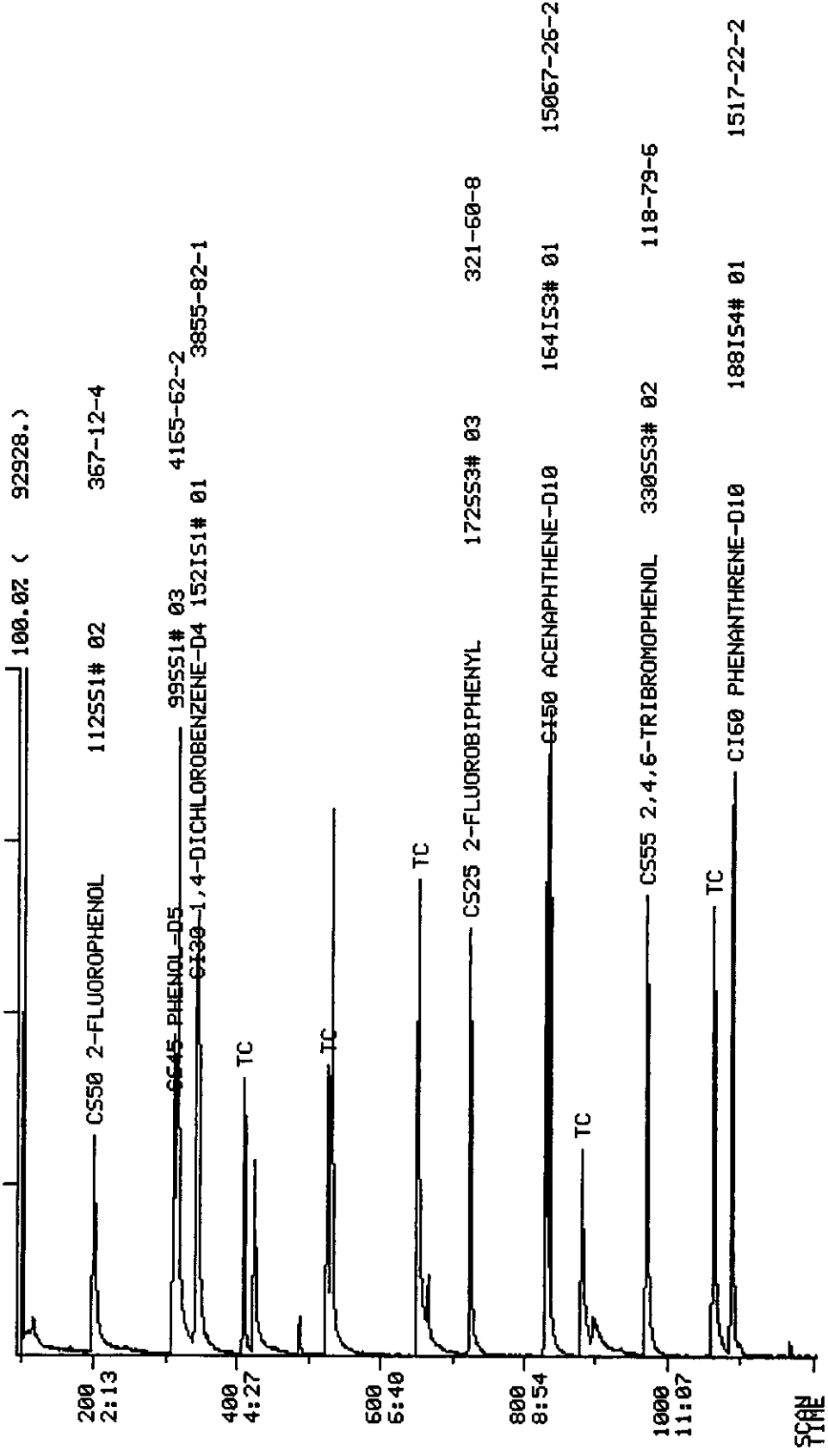
*confirms*

Target Compounds: MS

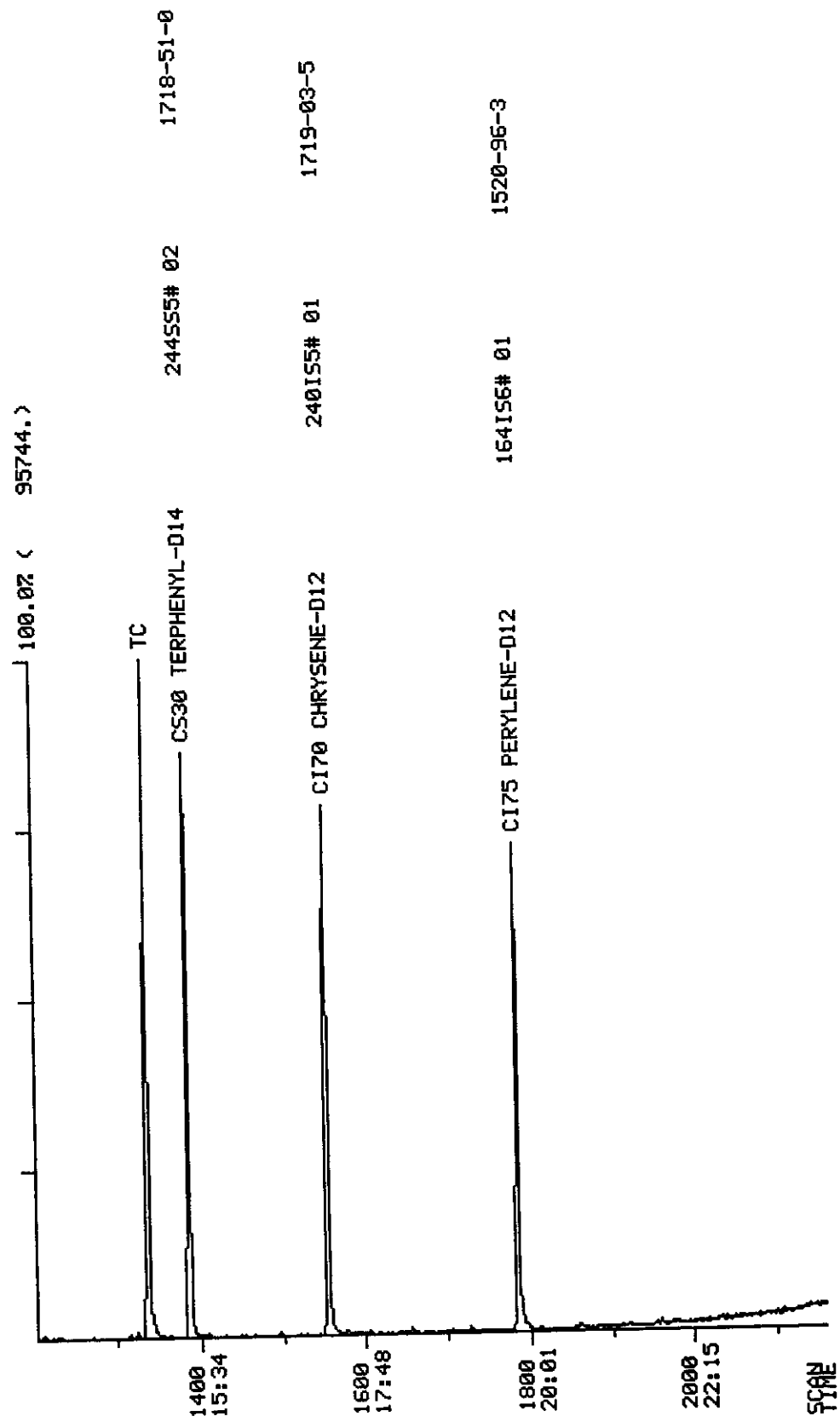
Parameter	Scan	Result	Units	Reporting
				Limit (J values)
CS70 2-CHLOROPHENOL-D4		ND	UG/A	10.0
CS75 1,2-DICHLOROBENZ-N-D4		ND	UG/A	10.0
C315 PHENOL	309	26.	UG/A	10.0
C330 2-CHLOROPHENOL	314	65.	UG/A	10.0
C340 1,4-DICHLOROBENZENE	343	29.	UG/A	10.0
C370 N-NITROSO-DI-N-PROPYLAM	408	32.	UG/A	10.0
C445 1,2,4-TRICHLOROBENZENE	524	30.	UG/A	10.0
C465 4-CHLORO-3-METHYLPHENO	650	62.	UG/A	10.0
C550 ACENAPHTHENE	834	38.	UG/A	10.0
C560 4-NITROPHENOL	895	16.	UG/A	50.0 J
C570 2,4-DINITROTOLUENE	879	39.	UG/A	10.0
C635 PENTACHLOROPHENOL	1062	64.	UG/A	50.0
C715 PYRENE	1335	44.	UG/A	10.0

Reviewed by: *[Signature]*

DATA FROM FILE: 300681BFDSI SCANS 94 TO 1204 ACQUIRED: 09/02/98 19:29:00  
 CALI: 300681BFDSI #3  
 SAMPLE: DCS-AD 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA) >15A NA M



DATA FROM FILE: 300681BFDSI SCANS 1204 TO 2158 ACQUIRED: 09/02/98 19:29:00  
 CALI: 300681BFDSI #3  
 SAMPLE: DCS-AD 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100Z/100Z \*KNA/NA >/15A NA M



1747

ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980902A

## QUANTERRA QUANTITATION SUMMARY

File: 300681BFDSI

Sample: DCS-AD 01AUG98-16A 1SA/1ML

INST. ID: F16

Analyst: DAT

Instrument Id: F16

Analyzed: 09/02/98 19:29

Compounds with amounts less than 0.20 reported as NOT FOUND

Library									Units: UG/ML	
No	Entry	Name	Mass	Meth	Scan	Ref	Area	RRF(L)	Amount	
1	S1#	1 CI30 1,4-DICHLOROBENZENE	152	A	BB	340	1	27115.	1.000	40.000
2	S2#	1 CI40 NAPHTHALENE-D8	136	A	BB	530	2	91722.	1.000	40.000
3	S3#	1 CI50 ACENAPHTHENE-D10	164	A	BB	828	3	52118.	1.000	40.000
4	S4#	1 CI60 PHENANTHRENE-D10	188	A	BB	1087	4	89087.	1.000	40.000
5	S5#	1 CI70 CHRYSENE-D12	240	A	BB	1552	5	79298.	1.000	40.000
6	S6#	1 CI75 PERYLENE-D12	264	A	BB	1784	6	69584.	1.000	40.000
7	S2#	2 CS20 NITROBENZENE-D5	82	A	BB	422	2	32551.	0.470	30.232
8	S3#	3 CS25 2-FLUOROBIPHENYL	172	A	BB	723	3	54422.	1.234	33.840
9	S5#	2 CS30 TERPHENYL-D14	244	A	BB	1384	5	72015.	0.995	36.501
10	S1#	3 CS45 PHENOL-D5	99	A	BB	308	1	35704.	2.025	26.004
11	S1#	2 CS50 2-FLUOROPHENOL	112	A	BB	197	1	35344.	1.368	38.106
12	S3#	2 CS55 2,4,6-TRIBROMOPHENO	330	A	BB	969	3	22909.	0.236	74.610
13	S1#	5 CS70 2-CHLOROPHENOL-D4	132				1			NOT FOUND
14	S1#	6 CS75 1,2-DICHLOROBENZENE-D4	152				1			NOT FOUND
15	S1#105	C315 PHENOL	94	A	BB	309	1	34746.	1.968	26.047
16	S1#115	C330 2-CHLOROPHENOL	128	A	BB	314	1	65937.	1.497	64.982
17	S1#130	C340 1,4-DICHLOROBENZENE	146	A	BB	343	1	32129.	1.617	29.302
18	S1#190	C370 N-NITROSO-DI-N-PROP	70	A	BB	408	1	19533.	0.906	31.809
19	S2#	55 C445 1,2,4-TRICHLOROBENZ	180	A	BB	524	2	22687.	0.331	29.897
20	S2#130	C465 4-CHLORO-3-METHYLPH	107	A	BB	650	2	48270.	0.338	62.327
21	S3#	80 C550 ACENAPHTHENE	153	A	BB	834	3	56768.	1.159	37.604
22	S3#	95 C560 4-NITROPHENOL	109	A	BB	895	3	3696.	0.173	16.429
23	S3#105	C570 2,4-DINITROTOLUENE	165	A	BB	879	3	20426.	0.397	39.464
24	S4#	50 C635 PENTACHLOROPHENOL	266	A	BB	1062	4	22149.	0.155	64.144
25	S5#	15 C715 PYRENE	202	A	BB	1335	5	114565.	1.323	43.694

QUANTERRA Internal Standard Check

Standard Filename: ST16980902

Analyzed: 09/02/98 14:53

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980902	23319.		3.78	79260.		5.91	43605.		9.22
Upper Limit	46638.=+200		4.28	158519.=+200		6.41	87211.=+200		9.72
Lower Limit	11659.=+ 50		3.28	39630.=+ 50		5.41	21803.=+ 50		8.72
Filename									
1 300681BFDSI	27115.= 116		3.78	91722.= 115		5.90	52118.= 119		9.21

IS# 1 = CI30 1,4-DICHLOROBENZENE-D4  
 IS# 2 = CI40 NAPHTHALENE-D8  
 IS# 3 = CI50 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980902	76218.		12.09	51424.		17.27	44306.		19.84
Upper Limit	152436.=+200		12.59	102848.=+200		17.77	88612.=+200		20.34
Lower Limit	38109.=+ 50		11.59	25712.=+ 50		16.77	22153.=+ 50		19.34
Filename									
1 300681BFDSI	89087.= 116		12.09	79298.= 154		17.26	69584.= 157		19.84

IS# 4 = CI60 PHENANTHRENE-D10  
 IS# 5 = CI70 CHRYSENE-D12  
 IS# 6 = CI75 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBFMS  
Std Id: ST16980831

Sample: DCS-A 01AUG98-16A 1SA/1ML INST. ID: F16  
Client: PACIFI Date Analyzed: 08/31/98 17:01  
Analyst: DAT Instrument ID: F16 Run Factor: 1.00  
Quan List Threshold: 0.95 Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC Limits
CS20 NITROBENZENE-D5	50.00	31.30	62.6	45 107
CS25 2-FLUOROBIPHENY	50.00	35.00	70.0	62 110
CS30 TERPHENYL-D14	50.00	41.20	82.4	58 135
CS45 PHENOL-D5	100.0	28.00	28.0	* 43 130
CS50 2-FLUOROPHENOL	100.0	38.90	38.9	36 111
CS55 2,4,6-TRIBROMOP	100.0	83.10	83.1	58 131

Target Compounds: MS

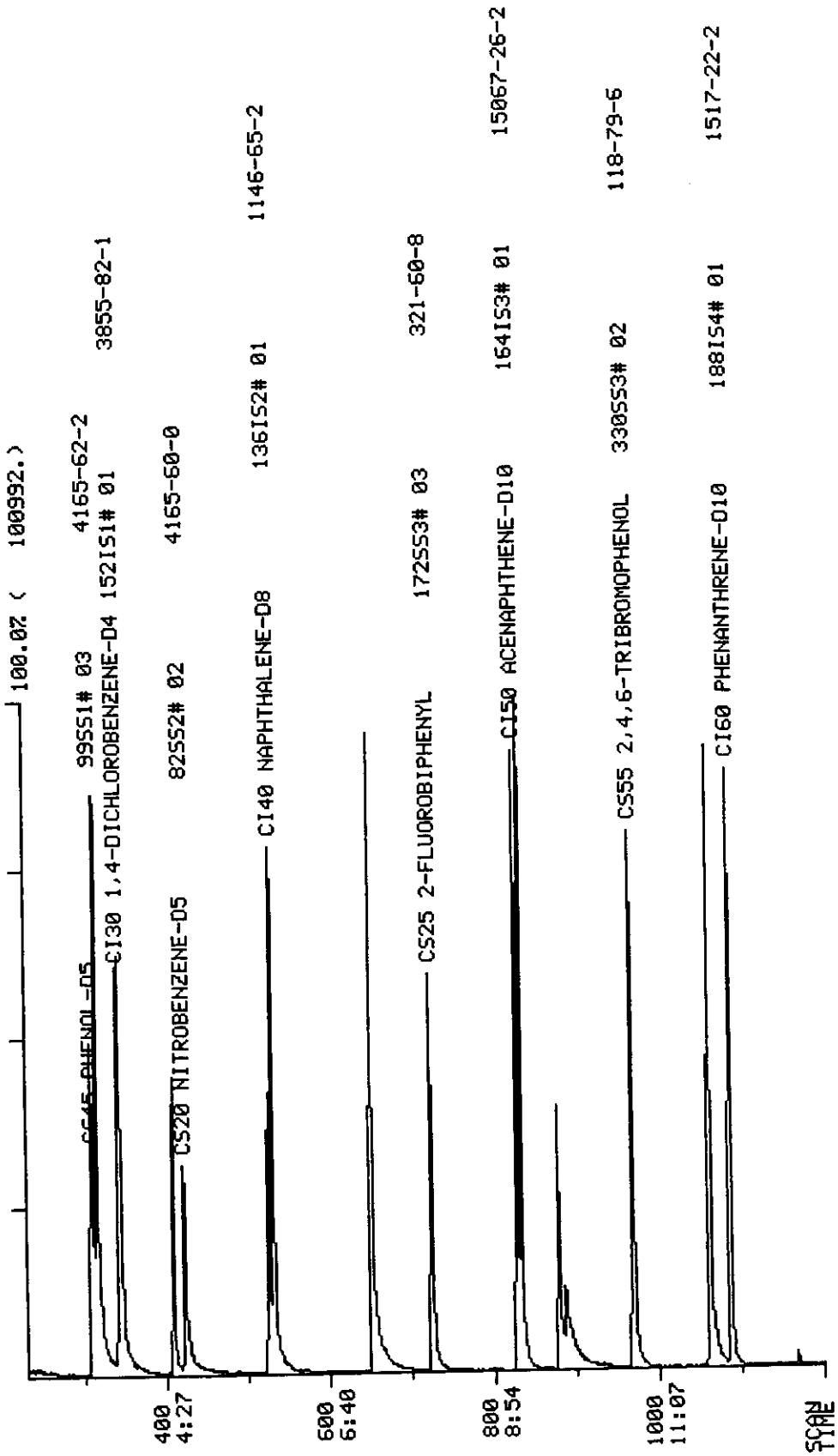
Parameter	Scan	Result	Units	Reporting
				Limit (J values)
CS70 2-CHLOROPHENOL-D4		ND	UG/A	10.0
CS75 1,2-DICHLOROBZN-D4	340	43.	UG/A	10.0
C315 PHENOL	308	29.	UG/A	10.0
C330 2-CHLOROPHENOL	313	66.	UG/A	10.0
C340 1,4-DICHLOROBENZENE	342	28.	UG/A	10.0
C370 N-NITROSO-DI-N-PROPYLAM	408	37.	UG/A	10.0
C445 1,2,4-TRICHLOROBENZENE	524	28.	UG/A	10.0
C465 4-CHLORO-3-METHYLPHENO	649	71.	UG/A	10.0
C550 ACENAPHTHENE	833	40.	UG/A	10.0
C560 4-NITROPHENOL	887	27.	UG/A	50.0 J
C570 2,4-DINITROTOLUENE	878	44.	UG/A	10.0
C635 PENTACHLOROPHENOL	1061	89.	UG/A	50.0
C715 PYRENE	1334	49.	UG/A	10.0

Reviewed by: 

1750

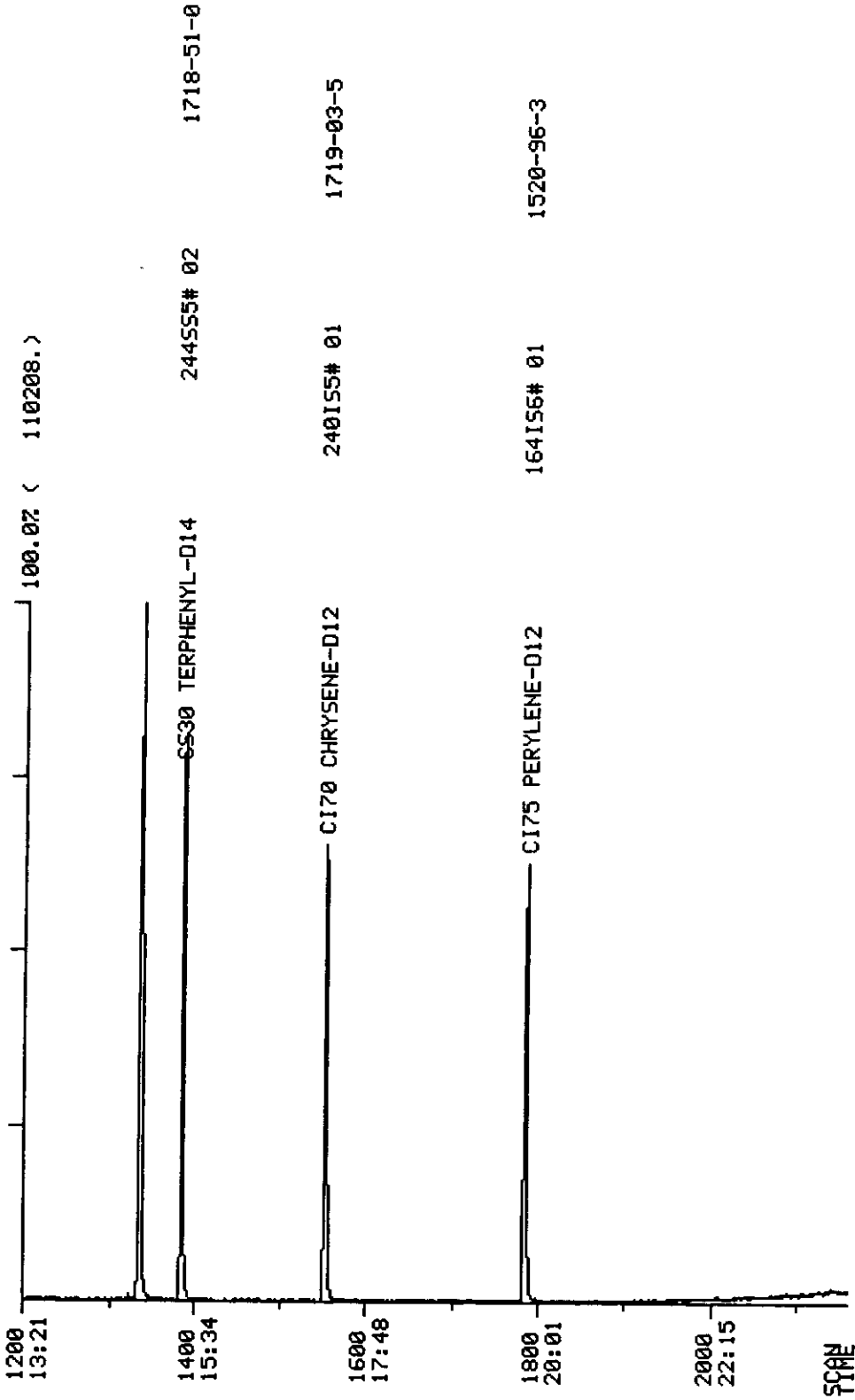


DATA FROM FILE: 300681MBFMS SCANS 230 TO 1200 ACQUIRED: 08/31/98 17:01:00  
 CALI: 300681MBFMS #3  
 SAMPLE: DCS-A 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA )/15A NA M



1701

DATA FROM FILE: 300681MBFMS SCANS 1200 TO 2158 ACQUIRED: 08/31/98 17:01:00  
CALI: 300681MBFMS #3  
SAMPLE: DCS-A 01AUG98-16A 15A/1ML INST. ID: F16  
CONDS.: UG/ML \*100%/100% \*(NA/NA) /15A NA M



1752

ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBFMS

Sample: DCS-A 01AUG98-16A 1SA/1ML INST. ID: F16  
 Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 17:01

Compounds with amounts less than 0.20 reported as NOT FOUND

Library							Units: UG/ML	
No Entry	Name	Mass Meth	Scan	Ref	Area	RRF(L)	Amount	
1 S1#	1 C130 1,4-DICHLOROBENZENE	152 A BB	340	1	28516.	1.000	40.000	
2 S2#	1 C140 NAPHTHALENE-D8	136 A BB	529	2	101945.	1.000	40.000	
3 S3#	1 C150 ACENAPHTHENE-D10	164 A BB	828	3	56455.	1.000	40.000	
4 S4#	1 C160 PHENANTHRENE-D10	188 A BB	1086	4	91166.	1.000	40.000	
5 S5#	1 C170 CHRYSENE-D12	240 A BB	1551	5	75878.	1.000	40.000	
6 S6#	1 C175 PERYLENE-D12	264 A BV	1783	6	63304.	1.000	40.000	
7 S2#	2 CS20 NITROBENZENE-D5	82 A BB	421	2	37439.	0.470	31.284	
8 S3#	3 CS25 2-FLUOROBIPHENYL	172 A BB	722	3	60974.	1.234	35.001	
9 S5#	2 CS30 TERPHENYL-D14	244 A BB	1383	5	77780.	0.995	41.200	
10 S1#	3 CS45 PHENOL-D5	99 A BB	306	1	40389.	2.025	27.971	
11 S1#	2 CS50 2-FLUOROPHENOL	112 A BB	196	1	37991.	1.368	38.949	
12 S3#	2 CS55 2,4,6-TRIBROMOPHENO	330 A BB	968	3	27642.	0.236	83.107	
13 S1#	5 CS70 2-CHLOROPHENOL-D4	132		1	NOT FOUND			
14 S1#	6 CS75 1,2-DICHLOROBENZ-N-D4	152 A BB	340	1	28516.	0.920	43.476	
15 S1#105	C315 PHENOL	94 A BB	308	1	40013.	1.968	28.522	
16 S1#115	C330 2-CHLOROPHENOL	128 A BB	313	1	70308.	1.497	65.887	
17 S1#130	C340 1,4-DICHLOROBENZENE	146 A BB	342	1	31738.	1.617	27.524	
18 S1#190	C370 N-NITROSO-DI-N-PROP	70 A BB	408	1	23823.	0.906	36.890	
19 S2# 55	C445 1,2,4-TRICHLOROBENZ	180 A BB	524	2	23662.	0.331	28.055	
20 S2#130	C465 4-CHLORO-3-METHYLPH	107 A BB	649	2	60821.	0.338	70.658	
21 S3# 80	C550 ACENAPHTHENE	153 A BB	833	3	64768.	1.159	39.607	
22 S3# 95	C560 4-NITROPHENOL	109 A BB	887	3	6606.	0.173	27.110	
23 S3#105	C570 2,4-DINITROTOLUENE	165 A BB	878	3	24726.	0.397	44.102	
24 S4# 50	C635 PENTACHLOROPHENOL	266 A BB	1061	4	31357.	0.155	88.740	
25 S5# 15	C715 PYRENE	202 A BB	1334	5	123841.	1.323	49.361	

1758

QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 300681MBFMS	28516.= 99		3.78	101945.= 101		5.88	56455.= 103		9.21

IS# 1 = C130 1,4-DICHLOROBENZENE-D4  
 IS# 2 = C140 NAPHTHALENE-D8  
 IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 300681MBFMS	91166.= 92		12.08	75878.= 121		17.25	63304.= 124		19.83

IS# 4 = C160 PHENANTHRENE-D10  
 IS# 5 = C170 CHRYSENE-D12  
 IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

QUANTERRA GC/MS  
Target Compound Data Summary Sheet

Data File: 300681MBFDS  
Std Id: ST16980831

Sample: DCS-AD 01AUG98-16A 1SA/1ML  
Client: PACIFI  
Analyst: DAT Instrument ID: F16  
Quan List Threshold: 0.95

INST. ID: F16  
Date Analyzed: 08/31/98 17:30  
Run Factor: 1.00  
Surrogate Vol.: 1.00

Surrogate Spike Recoveries  
8270-G, LIMS 10/10/96

Surrogate	Surrogate Amount (ug)		% Recovery	
	Spiked	Measured	Measured	QC limits
CS20 NITROBENZENE-D5	50.00	32.30	64.6	45 107
CS25 2-FLUOROBIPHENY	50.00	33.90	67.8	62 110
CS30 TERPHENYL-D14	50.00	37.80	75.6	58 135
CS45 PHENOL-D5	100.0	22.40	22.4	* 43 130
CS50 2-FLUOROPHENOL	100.0	39.90	39.9	36 111
CS55 2,4,6-TRIBROMOP	100.0	72.90	72.9	58 131

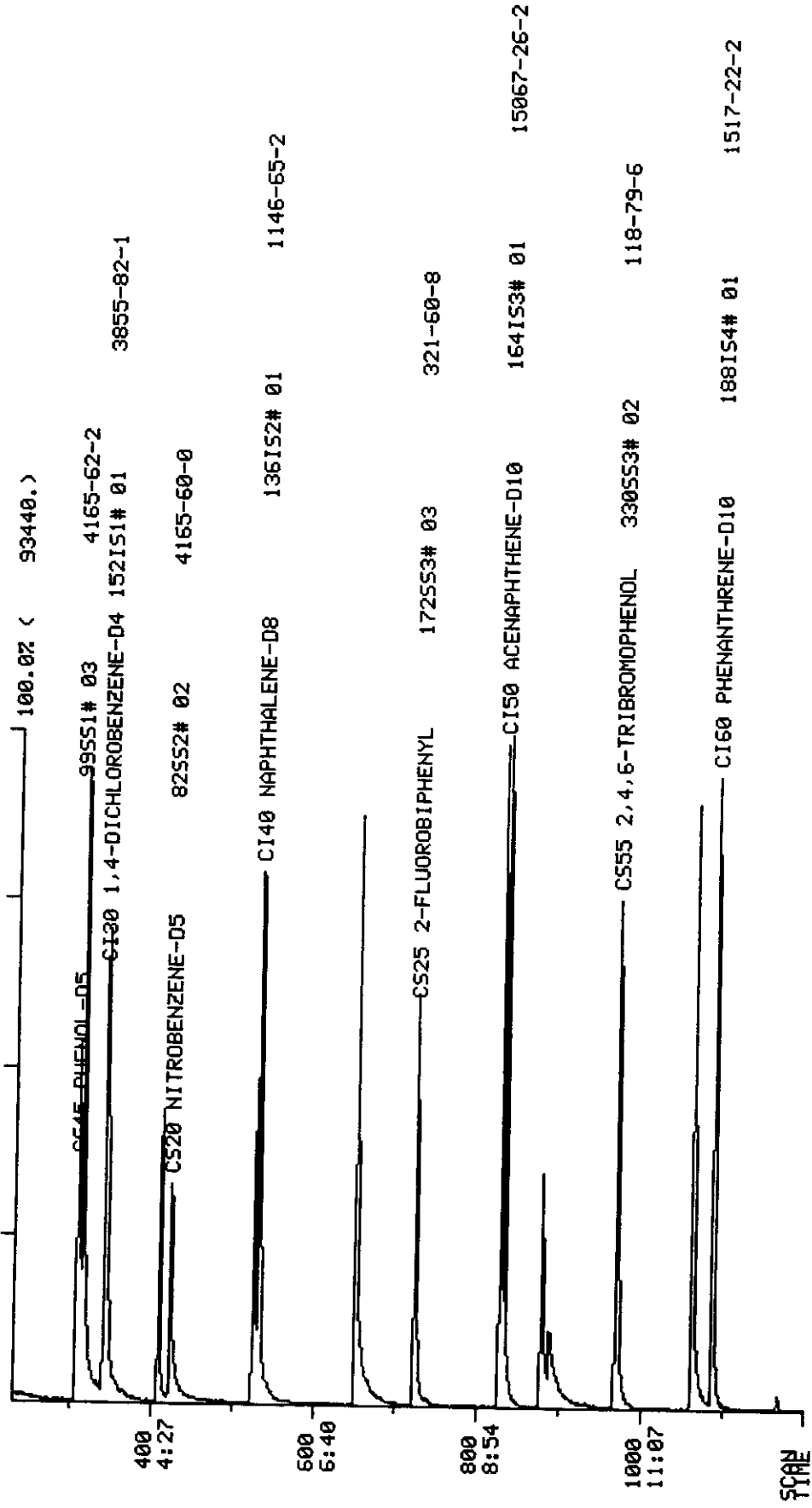
Target Compounds: MS

Parameter	Scan	Result	Units	Reporting
				Limit (J values)
CS70 2-CHLOROPHENOL-D4		ND	UG/A	10.0
CS75 1,2-DICHLOROBENZ-N-D4		ND	UG/A	10.0
C315 PHENOL	308	29.	UG/A	10.0
C330 2-CHLOROPHENOL	313	67.	UG/A	10.0
C340 1,4-DICHLOROBENZENE	342	30.	UG/A	10.0
C370 N-NITROSO-DI-N-PROPYLAM	408	33.	UG/A	10.0
C445 1,2,4-TRICHLOROBENZENE	524	30.	UG/A	10.0
C465 4-CHLORO-3-METHYLPHENO	649	66.	UG/A	10.0
C550 ACENAPHTHENE	833	38.	UG/A	10.0
C560 4-NITROPHENOL	888	24.	UG/A	50.0 J
C570 2,4-DINITROTOLUENE	878	40.	UG/A	10.0
C635 PENTACHLOROPHENOL	1061	76.	UG/A	50.0
C715 PYRENE	1334	45.	UG/A	10.0

Reviewed by: KV 10/19/98

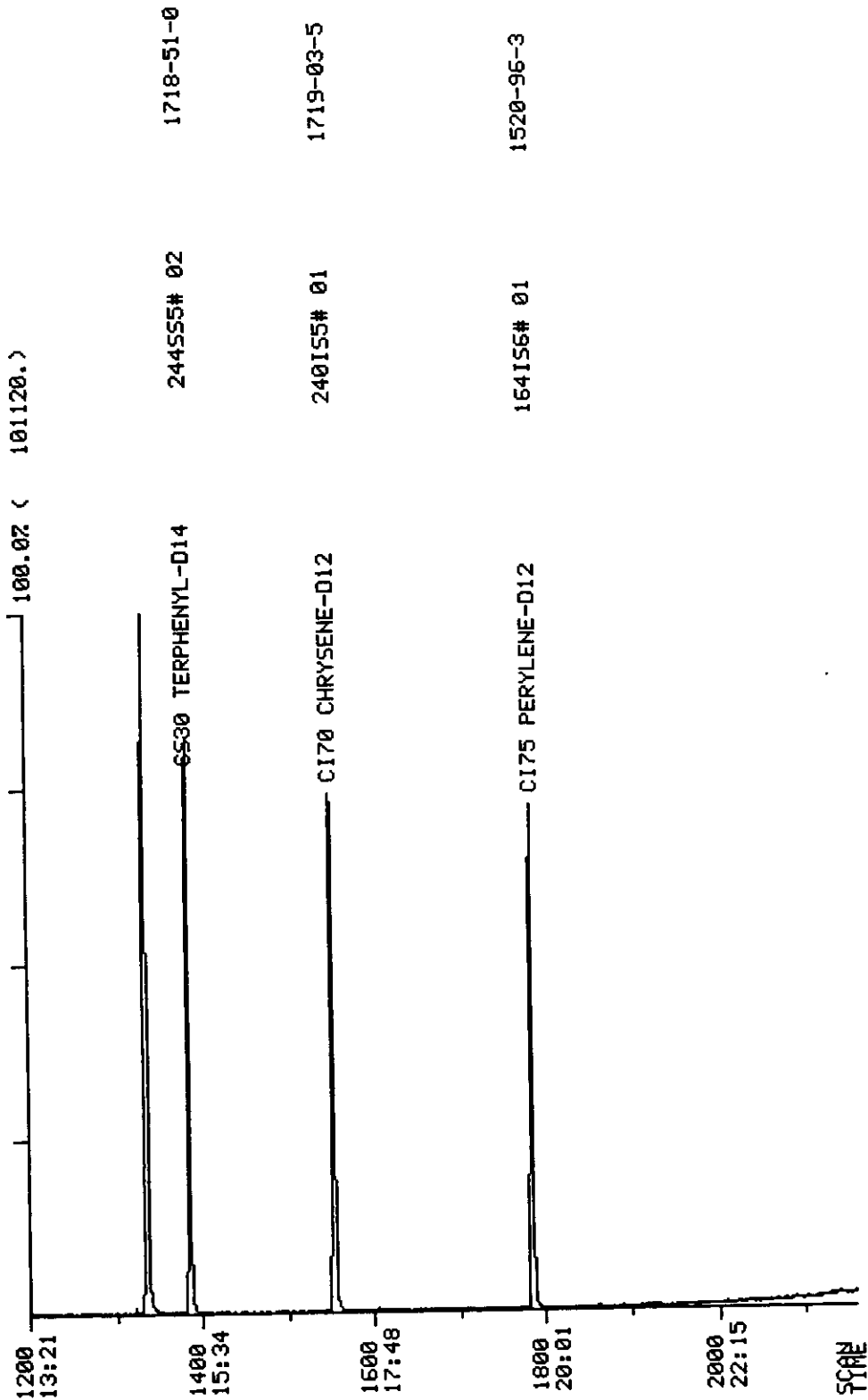
1755

DATA FROM FILE: 300681MBFDS SCANS 230 TO 1200 ACQUIRED: 08/31/98 17:30:00  
 CALI: 300681MBFDS #3  
 SAMPLE: DCS-AD 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*100%/100% \*(NA/NA )/15A NA M



17  
 20  
 20

DATA FROM FILE: 300681MBFDS SCANS 1200 TO 2158 ACQUIRED: 08/31/98 17:30:00  
 CALI: 300681MBFDS #3  
 SAMPLE: DCS-AD 01AUG98-16A 15A/1ML INST. ID: F16  
 CONDS.: UG/ML \*1ML \*100%/100% \*(NA/NA )/15A NA M



1757

ADDITIONAL STANDARD USED FOR THIS REPORT IS: ST16980831A

## QUANTERRA QUANTITATION SUMMARY

File: 300681MBFDS

Sample: DCS-AD 01AUG98-16A 1SA/1ML INST. ID: F16  
Analyst: DAT Instrument Id: F16 Analyzed: 08/31/98 17:30

Compounds with amounts less than 0.20 reported as NOT FOUND

Library		Units: UG/ML								
No	Entry	Name	Mass	Meth	Scan	Ref	Area	RRF(L)	Amount	
1	S1#	1 C130	1,4-DICHLOROBENZENE	152	A	BB	340	1	28626. 1.000	40.000
2	S2#	1 C140	NAPHTHALENE-D8	136	A	BB	529	2	99229. 1.000	40.000
3	S3#	1 C150	ACENAPHTHENE-D10	164	A	BB	828	3	54918. 1.000	40.000
4	S4#	1 C160	PHENANTHRENE-D10	188	A	BB	1086	4	93048. 1.000	40.000
5	S5#	1 C170	CHRYSENE-D12	240	A	BB	1551	5	77501. 1.000	40.000
6	S6#	1 C175	PERYLENE-D12	264	A	BV	1783	6	65820. 1.000	40.000
7	S2#	2 CS20	NITROBENZENE-D5	82	A	BB	421	2	37607. 0.470	32.285
8	S3#	3 CS25	2-FLUOROBIPHENYL	172	A	BB	722	3	57505. 1.234	33.934
9	S5#	2 CS30	TERPHENYL-D14	244	A	BB	1383	5	72798. 0.995	37.753
10	S1#	3 CS45	PHENOL-D5	99	A	BV	306	1	32420. 2.025	22.365
11	S1#	2 CS50	2-FLUOROPHENOL	112	A	BB	196	1	39025. 1.368	39.855
12	S3#	2 CS55	2,4,6-TRIBROMOPHENO	330	A	BV	968	3	23589. 0.236	72.907
13	S1#	5 CS70	2-CHLOROPHENOL-D4	132				1	NOT FOUND	
14	S1#	6 CS75	1,2-DICHLOROBENZ-N-D4	152				1	NOT FOUND	
15	S1#105	C315	PHENOL	94	A	BB	308	1	40618. 1.968	28.842
16	S1#115	C330	2-CHLOROPHENOL	128	A	BB	313	1	71608. 1.497	66.845
17	S1#130	C340	1,4-DICHLOROBENZENE	146	A	BB	342	1	34461. 1.617	29.770
18	S1#190	C370	N-NITROSO-DI-N-PROP	70	A	BB	408	1	21467. 0.906	33.113
19	S2#	55 C445	1,2,4-TRICHLOROBENZ	180	A	BB	524	2	24942. 0.331	30.382
20	S2#130	C465	4-CHLORO-3-METHYLPH	107	A	BB	649	2	55273. 0.338	65.970
21	S3#	80 C550	ACENAPHTHENE	153	A	BB	833	3	60897. 1.159	38.282
22	S3#	95 C560	4-NITROPHENOL	109	A	BB	888	3	5735. 0.173	24.196
23	S3#105	C570	2,4-DINITROTOLUENE	165	A	BB	878	3	21544. 0.397	39.503
24	S4#	50 C635	PENTACHLOROPHENOL	266	A	BB	1061	4	27378. 0.155	75.913
25	S5#	15 C715	PYRENE	202	A	BB	1334	5	114585. 1.323	44.715



## QUANTERRA Internal Standard Check

Standard Filename: ST16980831

Analyzed: 08/31/98 12:31

Standard	IS# 1			IS# 2			IS# 3		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	28649.		3.78	100432.		5.90	54347.		9.21
Upper Limit	57297.=+200		4.28	200864.=+200		6.40	108695.=+200		9.71
Lower Limit	14324.=+ 50		3.28	50216.=+ 50		5.40	27174.=+ 50		8.71
Filename									
1 300681MBFDS	28626.=	99	3.78	99229.=	98	5.88	54918.=	101	9.21

IS# 1 = C130 1,4-DICHLOROBENZENE-D4

IS# 2 = C140 NAPHTHALENE-D8

IS# 3 = C150 ACENAPHTHENE-D10

\* - indicates an I.S. is outside QC limit(s)

Standard	IS# 4			IS# 5			IS# 6		
	Area	%	RT	Area	%	RT	Area	%	RT
ST16980831	98520.		12.08	62289.		17.26	50750.		19.83
Upper Limit	197039.=+200		12.58	124578.=+200		17.76	101500.=+200		20.33
Lower Limit	49260.=+ 50		11.58	31145.=+ 50		16.76	25375.=+ 50		19.33
Filename									
1 300681MBFDS	93048.=	94	12.08	77501.=	124	17.25	65820.=	129	19.83

IS# 4 = C160 PHENANTHRENE-D10

IS# 5 = C170 CHRYSENE-D12

IS# 6 = C175 PERYLENE-D12

\* - indicates an I.S. is outside QC limit(s)

Sample Extraction/Preparation Log Copies

QUANTERRA-CALLAB LABORATORY  
Analytical Bench Sheet

Project Number: 300681

Date: 31 JUL 98

Page 1 of 1

Test Code: P-8270-S

Test Description: Prep - Semivolatile Organics by GC/MS

Turnaround Status: NORMAL

Sample Matrix: AIRTRAIN Date Assigned: 7/31

Prep Completion Due: \_\_\_\_\_

SOP-Section/Page (Revision):

SAC-OP-0008

Extractions:  Separatory Funnel  Continuous Liquid/Liquid  Sonication  Soxhlet  KD  
 Roto Vap  Shaker  Other \_\_\_\_\_

Samp ID	Holding Time	Sample Size	Spiked & in Solvent Date/Initials	1st pH To:	2nd pH To:	Extraction Sonic/Shak Date/Initials	KD/N2 Reduc. ABN.	GPC	KD/N2 Reduc.	Screen Date/Initials	Final Concentn Date/Initials	(X)
0001SA	08 AUG 98	AIR TRAIN				DWT 7/31/98	DWT 8/4/98	NA	DWT 8/5/98	GB 8/5/98	1/3 SA 100ml	300x
0002SA	08 AUG 98	(XAD/FICT)										↓
0003SA	08 AUG 98										1/3 SA ML	3x
0004SA	08 AUG 98											↓
0005SA	09 AUG 98											↓
0006SA	10 AUG 98											↓
0007SA	10 AUG 98					DWT 7/31/98					1/3 SA 100ml	300x
0008SA	09 AUG 98										1/3 SA ML	3x
0009SA	08 AUG 98											↓
0010SA	08 AUG 98										1/3 SA 100ml	300x
0011SA	11 AUG 98										1/3 SA 100ml	↓
MB-A	NA	↓								NA	1/3 SA ML	3x
DCS-A	↓	NA										↓
DCS-AD	↓	↓										↓

Project Due: 19 AUG 98

DCS With Project:

300681

DCS Code:

31 JUL 98 - 16A

LCS With Project:

NA

LCS Code:

NA

Method Blank With Project:

300681

Extraction Includes Project(s):

300681

Notes/Comments:

\*\* PLEASE PUT COMMENTS ON ANOMALY SHEET \*\*

MSQC with Project:

NA

MSQC Code:

NA

1702

Date To Instruments:

8/5/98

By:

DWT

Peer Approval By: \_\_\_\_\_

QUANTERRA-CALLAB LABORATORY  
Quality Assurance Sheet

Project #: 300681 Test: P-8270-S

Quality Assurance/Quality Control:

Sample ID: ALL Spiking Standard Name: ADN SW846 SURROGATE

Spike Code ID: AT031698A Volume: 1500 µl Conc: 100-200 ppm

Spiked By: DJT Witness: GBL

Sample ID: DCS-A/AD Spiking Standard Name: ADN SW846 SPIKE

Spike Code ID: AT040891A Volume: 1500 µl Conc: 100-200 ppm

Spiked By: DJT Witness: GBL

Sample ID: \_\_\_\_\_ Spiking Standard Name: \_\_\_\_\_

Spike Code ID: \_\_\_\_\_ Volume: \_\_\_\_\_ Conc: \_\_\_\_\_

Spiked By: \_\_\_\_\_ Witness: \_\_\_\_\_

Sample ID: \_\_\_\_\_ Spiking Standard Name: \_\_\_\_\_

Spike Code ID: \_\_\_\_\_ Volume: \_\_\_\_\_ Conc: \_\_\_\_\_

Spiked By: \_\_\_\_\_ Witness: \_\_\_\_\_

Solvent: Manufacturer And Lot Number:

Extracting Solvent: DCM (BAK) M17327

Solvent Keeper: \_\_\_\_\_

Transfer Solvent: \_\_\_\_\_

Derivatizing Reagent: \_\_\_\_\_

Final Solvent: \_\_\_\_\_

Other: \_\_\_\_\_

Was an aliquot taken? (Y) / N

If yes, how much was aliquoted? 5 mL / 15 mL

What was the sample concentration before aliquot? SA / 15 mL

Why was the aliquot taken? SPLIT w/HR-PAH, ARCHIVE

QUANTERRA-CALLAB LABORATORY  
Anomaly Report Form

Project #: 300681 Test: P-8270-S

Anomalies:

Sample(s) Affected:

- Sample broken during shipping. \_\_\_\_\_
- Limited sample volume. \_\_\_\_\_
- Matrix required a subsample to facilitate extraction. \_\_\_\_\_
- pH of sample required excess acid/base (please circle). \_\_\_\_\_
- Solvent and sample were miscible. \_\_\_\_\_
- Emulsion problems. \_\_\_\_\_
- Emulsion problem required centrifuging. \_\_\_\_\_
- Sample lost during preparation.  
Re-extraction performed? Yes/No \_\_\_\_\_
- Concentration volume went lower than specified in the SOP. \_\_\_\_\_
- Concentration time longer than normal. \_\_\_\_\_
- Extraction solvent saturated with organic material, therefore sample volume was not concentrated but adjusted based on the screening. \_\_\_\_\_
- Final concentration greater than specified in the SOP due to the screening results. \_\_\_\_\_
- Precipitate observed in the extract. \_\_\_\_\_
- Other, please comment below. 1, 2, 7, 10, 11

Comments:

# 1, 2, 7, 10 + 11 CONTAINED A LOT OF OIL - FINAL CONC BASED ON RESULTS OF SCREEN

Submitted By: DWT

Date: 8/5/98

1785

QUANTERRA-CALLAB LABORATORY  
 Prep (Extraction) Check List

Project #: 300681 Test: P-8270-S

Chemist: DJT Date: 8/5/98

Reviewer: \_\_\_\_\_ Date: \_\_\_\_\_

Number of samples: 11 Number of extracts: 14

Benchsheet:

SOP section and page number filled out properly.  
 Each step of extraction procedure initial'ed and dated.  
 Anomalies or deviations from SOP documented.  
 pH adjustments noted.  
 Solvent manufacturer and lot number completed.  
 Volumes and weights of samples to two sig. figs.

Initiated		Reviewed	
Y <input checked="" type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>

QA/QC:

DCS/MS forms filled out properly.  
 Spike information correct on bench sheet.  
 Spike witness?  
 If not spiked per SOP, were spike amounts verified?

Initiated		Reviewed	
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>

LIMS:

Extraction date (complete date) correct.  
 QC lot number present and correct.  
 QC Run number present and correct.  
 Prep released (date extract goes to Inst.).

Initiated		Reviewed	
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>

Extracts:

Number of extracts agree with benchsheet.  
 Concentrations of extract agree with benchsheet.  
 Testtube rack labelled with project id and test.

Initiated		Reviewed	
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>
Y <input checked="" type="checkbox"/>	N <input type="checkbox"/>	Y <input type="checkbox"/>	N <input type="checkbox"/>

Comments:

VOLUMES ATTACHED

QUANTERRA-CALLAB LABORATORY  
Analytical Bench Sheet

Project Number: 300681

Date: 31 JUL 98

Test Code: P-8270-S

Test Description: Prep - Semivolatile Organics by GC/MS

Turnaround Status: NORMAL

Sample Matrix: AIRTRAIN Date Assigned: \_\_\_\_\_

Prep Completion Due: \_\_\_\_\_

SOP-Section/Page (Revision): \_\_\_\_\_

Extractions:  Separatory Funnel  Continuous Liquid/Liquid  Sonication  Soxhlet  KD  
 Roto Vap  Shaker  Other \_\_\_\_\_

Samp ID	Holding Time	Sample Size	Spiked & in Solvent Date/Initials	1st pH To:	2nd pH To:	Extraction Sonic/Shak Date/Initials	KD/N2 Reduc. ABN	GPC	KD/N2 Reduc.	Screen Date/Initials	Final Concentr Date/Initials
0001SA	08 AUG 98	SEE ATTACHED	3/1/98 8/1	14	1	8/1/98 8/1	ONT 8/5/98				
0002SA	08 AUG 98										
0003SA	08 AUG 98										
0004SA	08 AUG 98										
0005SA	09 AUG 98										
0006SA	10 AUG 98										
0007SA	10 AUG 98										
0008SA	09 AUG 98										
0009SA	08 AUG 98										
0010SA	08 AUG 98										
0011SA	11 AUG 98										
MB	NA	1.0L							ONT 8/5/98	NA	SA ML
A											
AD											

COMBINED WITH SOXHLET FRACTION

Project Due: 19 AUG 98

DCS With Project: 300681

DCS Code: 01 AUG 98-16A

LCS With Project: NA

LCS Code: NA

Method Blank With Project: 300681

Extraction Includes Project(s): 300681

Notes/Comments:

\*\* PLEASE PUT COMMENTS ON ANOMALY SHEET \*\*

MSQC with Project: NA MSQC Code: NA

1705

Date To Instruments: 8/5/98

By: ONT

Peer Approval By: \_\_\_\_\_

QUANTERRA-CALLAB LABORATORY  
Quality Assurance Sheet

Project #: 300681 Test: P-8270-S

Quality Assurance/Quality Control:

Sample ID: MB, A/AD n<sup>y</sup> Spiking Standard Name: ABN SW 846 surrogate  
Spike Code ID: AT031698A Volume: 500  $\mu$ l Conc: 100-200 ppm  
Spiked By: SV Witness: C. Hinton

Sample ID: A/AD Spiking Standard Name: ABN SW 846 Spike  
Spike Code ID: AT04898A Volume: 500  $\mu$ l Conc: 100-200 ppm  
Spiked By: SV Witness: C. Hinton

Sample ID: \_\_\_\_\_ Spiking Standard Name: \_\_\_\_\_  
Spike Code ID: \_\_\_\_\_ Volume: \_\_\_\_\_ Conc: \_\_\_\_\_  
Spiked By: \_\_\_\_\_ Witness: \_\_\_\_\_

Sample ID: \_\_\_\_\_ Spiking Standard Name: \_\_\_\_\_  
Spike Code ID: \_\_\_\_\_ Volume: \_\_\_\_\_ Conc: \_\_\_\_\_  
Spiked By: \_\_\_\_\_ Witness: \_\_\_\_\_

Solvent: Manufacturer And Lot Number:

Extracting Solvent: MeCl<sub>2</sub> Lot # 117327  
Solvent Keeper: \_\_\_\_\_  
Transfer Solvent: \_\_\_\_\_  
Derivatizing Reagent: \_\_\_\_\_  
Final Solvent: \_\_\_\_\_  
Other: \_\_\_\_\_

Was an aliquot taken? Y / N

If yes, how much was aliquoted? \_\_\_\_\_ mL \_\_\_\_\_ mL

What was the sample concentration before aliquot? \_\_\_\_\_

Why was the aliquot taken? 8/5/98

1706



QUANTERRA-CALLAB LABORATORY  
Anomaly Report Form

Project #: 300681 Test: P-8270-S

Anomalies:

Sample(s) Affected:

- Sample broken during shipping. \_\_\_\_\_
- Limited sample volume. \_\_\_\_\_
- Matrix required a subsample to facilitate extraction. \_\_\_\_\_
- pH of sample required excess acid/base (please circle). \_\_\_\_\_
- Solvent and sample were miscible. \_\_\_\_\_
- Emulsion problems. \_\_\_\_\_
- Emulsion problem required centrifuging. \_\_\_\_\_
- Sample lost during preparation.  
Re-extraction performed? Yes/No \_\_\_\_\_
- Concentration volume went lower than specified in the  
SOP. \_\_\_\_\_
- Concentration time longer than normal. \_\_\_\_\_
- Extraction solvent saturated with organic material,  
therefore sample volume was not concentrated but  
adjusted based on the screening. \_\_\_\_\_
- Final concentration greater than specified in the SOP  
due to the screening results. \_\_\_\_\_
- Precipitate observed in the extract. \_\_\_\_\_
- Other, please comment below. MB/DCS

Comments:

*DUE TO LIMITED AMOUNT OF ABNSW846 SURROGATE, MB + DCS  
WERE SPIKED @ 1X, AND TAKEN TO SA/ML (SOXHLETS SPIKED 3X,  
TAKEN TO 1/3 SA/ML).*

Submitted By:   DNT  

Date:   8/5/98  

1707

**QUANTERRA-CALLAB LABORATORY  
Prep (Extraction) Check List**

Project #: 300681 Test: P-8270-S

Chemist: \_\_\_\_\_ Date: \_\_\_\_\_

Reviewer: \_\_\_\_\_ Date: \_\_\_\_\_

Number of samples: ~~11~~ Number of extracts: 3 (MB/DCS)  
*SAMPLES COMBINED  
WITH SOXHLET FRACTIONS*

**Benchsheet:**

SOP section and page number filled out properly.  
Each step of extraction procedure initial'ed and dated.  
Anomalies or deviations from SOP documented.  
pH adjustments noted.  
Solvent manufacturer and lot number completed.  
Volumes and weights of samples to two sig. figs.

<u>Initiated</u>		<u>Reviewed</u>	
Y <input checked="" type="checkbox"/>	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____

**QA/QC:**

DCS/MS forms filled out properly.  
Spike information correct on bench sheet.  
Spike witness?  
If not spiked per SOP, were spike amounts verified?

<u>Initiated</u>		<u>Reviewed</u>	
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____

**LIMS:**

Extraction date (complete date) correct.  
QC lot number present and correct.  
QC Run number present and correct.  
Prep released (date extract goes to Inst.).

<u>Initiated</u>		<u>Reviewed</u>	
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____

**Extracts:**

Number of extracts agree with benchsheet.  
Concentrations of extract agree with benchsheet.  
Testtube rack labelled with project id and test.

<u>Initiated</u>		<u>Reviewed</u>	
Y _____	N _____	Y _____	N _____
Y _____	N _____	Y _____	N _____
Y <input checked="" type="checkbox"/>	N _____	Y _____	N _____

Comments:

Analysis Log Copies

1769



QUANTERRA INCORPORATED  
West Sacramento

GC/MS INSTRUMENT LOG  
SEMI-VOLATILES

DATA UPLOAD/FILE ID

REVIEWED BY/DATE

INITIAL CAL DATE SHSL 8/2/98

AP9 8/11/98

INSTRUMENT ID F16

204-850-822-98A

PAGE 1 OF 1

DATE	TIME	OP	SAMPLE ID	FILE ID	CONC.	UN	TAPE ID	COMMENTS
8/31/98	1220	NAT	S04/48 DEFPF081398A	W16990831	-	2:		
	1231		S5TD080 B04/48 082248A	ST16990831	-			
	1307		B04/48 AP9 062598F	ST16990831A	-			
	1335		10   C		-			
	1402		20   D		-			
	1432		50   E		-			
	1502		120   G		-			
	1531		160   H		-			
	1602		SBLK5X 31 Jul 98-16A	300681MBSX	1/350/1000L			SERIAL ↓
	1631		↓ F 01 Aug 98-	SF	100/100L			SAL ↓
	1701		DCS-A ↓	FMS				SAM ↓
	1730		↓ -AD ↓	FDS				
	1800		↓ -A 31 Jul 98-	XMS	1/3 50/100L			
	1830		↓ -AD ↓	XDS				
	1900		S-MMS-2	01	1/350/1000L			
	1931		↓ -1B	02				SAB ↓
	2001		↓ -2	03	1/350/1000L			
	2030		↓ -FB	04				
	2100		↓ -4	05				SAB ↓
	2130		↓ -3	06				
	2159		S- ↓	07	1/350/1000L			
	2229		↓ -RB	08	1/350/1000L			
	2258		↓ -RB	09				
	2328		↓ -4	10	1/3 50/1000L			
	2358		↓ -5	11				



Instrument Tune

1773

DECAFLUOROTRIPHENYLPHOSPHINE

Tuning Report  
 08/27/98 9:12  
 Instrument: F16  
 # 294  
 Case Number:

Data: U16980827 # 294      Base m/z:  
 Cali: U16980827          RIC:  
 Analyst: DAT              Acct. No.:  
 Laboratory: QUANTERRA      Contract:

m/z	Intensity	% RA	Ion Abundance Criteria			Actual	Status
			Min %	Max %	Mass		
51	15536.	55.2	30.0	60.0	198	55.2	PASS
68	0.	0.0	0.0	2.0	69	0.0	PASS
69	18496.	65.7	0.0	100.0	198	65.7	PASS
70	114.	0.4	0.0	2.0	69	0.6	PASS
127	16080.	57.1	40.0	60.0	198	57.1	PASS
197	0.	0.0	0.0	1.0	198	0.0	PASS
198	28160.	100.0	100.0	100.0	198	100.0	PASS
199	1918.	6.8	5.0	9.0	198	6.8	PASS
275	6992.	24.8	10.0	30.0	198	24.8	PASS
365	800.	2.8	1.0	100.0	198	2.8	PASS
441	3580.	12.7	0.0	100.0	443	79.3	PASS
442	24000.	85.2	40.0	100.0	198	85.2	PASS
443	4512.	16.0	17.0	23.0	442	18.8	PASS

1774



Mass List                      Data: U16980827 # 295      Base m/z: 198  
 08/27/98 9:12:00 + 3:17      Cali: F16980821 # 3      RIC: 264704.  
 Sample: 50UG/ML DFTPP 081398A  
 Conds.: INST. ID: F16  
 #294 to #296 summed - #290

Mass	% RA	Inten.	Minima Maxima Mass	Min Inten: # 0 % RA	Inten.
35	0.00	0.			281.
444					
38?	s 1.13	318.	165	1.10	310.
39?	5.70	1606.	167	4.51	1270.
50?	15.34	4320.	168	2.36	664.
51?	s 55.17	15536.	174	1.06	298.
52?	2.85	802.	175	1.97	554.
56?	2.15	606.	179	3.42	964.
57?	4.20	1184.	180	2.35	662.
63?	2.08	586.	181	1.15	324.
65?	1.34	376.	185	1.30	366.
69	65.68	18496.	186	11.82	3328.
74	5.48	1542.	187	3.52	992.
75	8.17	2300.	192	1.31	370.
76	3.59	1010.	193	1.20	338.
77	57.44	16176.	196	3.66	1030.
78	4.13	1162.	198	100.00	28160.
79	3.93	1106.	199	6.81	1918.
80	3.05	860.	204	2.92	822.
81	3.91	1102.	205	5.36	1508.
82	1.04	294.	206	21.11	5944.
83	1.34	378.	207	3.06	862.
86	1.38	390.	211	1.46	410.
91	1.12	314.	217	5.52	1554.
92	1.14	320.	221	7.40	2084.
93	6.07	1710.	223	1.44	406.
98	4.84	1364.	224	11.39	3208.
99	3.44	970.	225	3.11	876.
101	2.05	578.	227	4.40	1240.
104	1.48	418.	229	1.17	330.
105	1.33	374.	244	9.79	2756.
107	15.37	4328.	245	1.55	436.
108	2.54	714.	246	1.61	452.
110	30.68	8640.	255	45.51	12816.
111	4.28	1206.	256	6.72	1892.
116	1.02	286.	258	2.24	630.
117	10.72	3020.	273	1.63	460.
118	1.02	288.	274	4.53	1276.
122	1.16	328.	275	24.83	6992.
123	1.50	422.	276	3.66	1032.
127	57.10	16080.	277	1.78	502.
128	4.62	1300.	296	5.28	1486.
129	20.06	5648.	323	2.15	606.
130	1.90	536.	334	1.27	358.
135	1.67	470.	365	2.84	800.
141	2.56	722.	372	1.29	362.
142	1.12	316.	423	4.74	1336.
147	1.23	346.	424	1.22	344.
148	2.54	716.	441	12.71	3580.
155	1.41	396.	442	85.23	24000.
156	2.09	588.	443	16.02	4512.
161	1.29	362.	444	1.71	482.

1775

Mass List                      Data: U16980827 # 295      Base m/z: 198  
08/27/98 9:12:00 + 3:17      Cali: F16980821 # 3      RIC: 264704.  
Sample: 50UG/ML DFTPP 081398A  
Conds.: INST. ID: F16  
#294 to #296 summed - #290

Mass	% RA	Inten.	Minima	Min Inten:	0.
65	0.00	0.			
75			Maxima #	0	
65?	1.34	376.			
67?	0.09	24.			
69	65.68	18496.			
70	0.40	114.			
72	0.09	24.			
73	0.82	230.			
74	5.48	1542.			
75	8.17	2300.			

1776

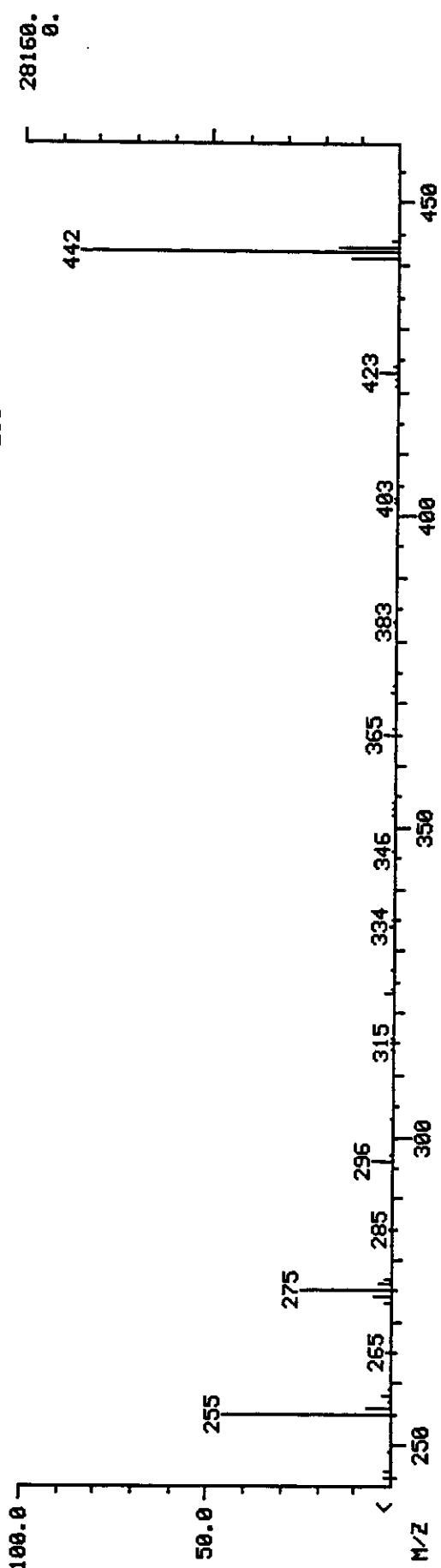
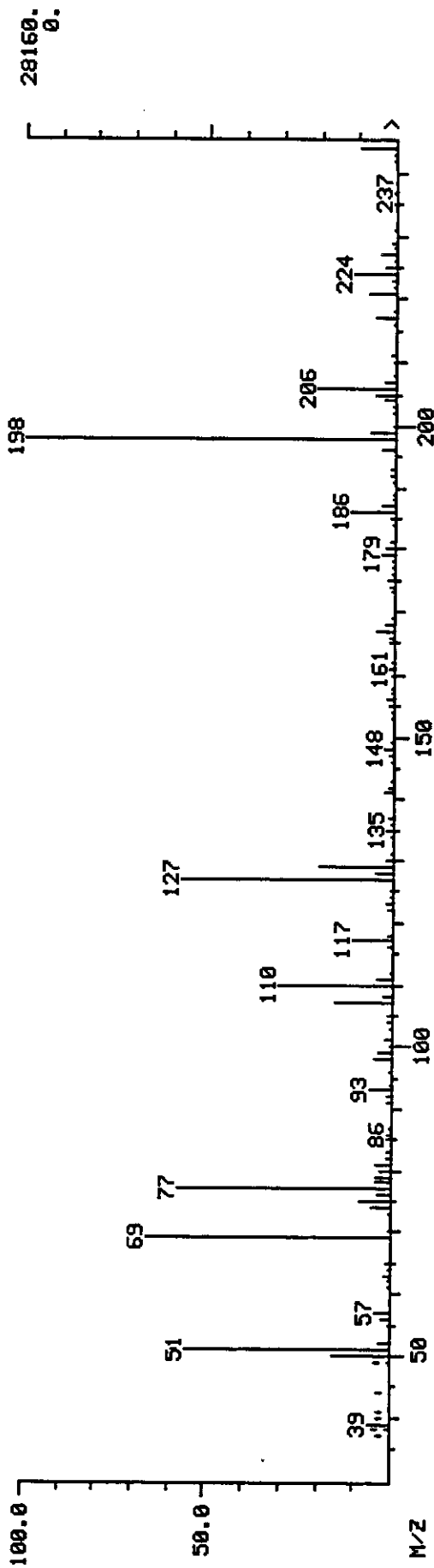
Mass List Data: U16980827 # 295 Base m/z: 198  
08/27/98 9:12:00 + 3:17 Cali: F16980821 # 3 RIC: 264704.  
Sample: SOUG/ML DFTPP 081398A  
Conds.: INST. ID: F16  
#294 to #296 summed - #290

Mass	% RA	Inten.	Minima	Min Inten:	0.
195	0.00	0.			
200			Maxima #	0	
196	3.66	1030.			
198	100.00	28160.			
199	6.81	1918.			
200	0.60	168.			

1777

DATA: U16980827 #295  
CALI: F16980821 #3  
BASE M/Z: 198  
RIC: 254704.

MASS SPECTRUM  
08/27/98 9:12:00 + 3:17  
SAMPLE: 50UG/ML DFTPP 081398A  
CONDS.: INST. ID: F16  
TEMP: 217 DEG. C  
#294 TO #296 SUMMED - #290



1270

SCANS 275 TO 314

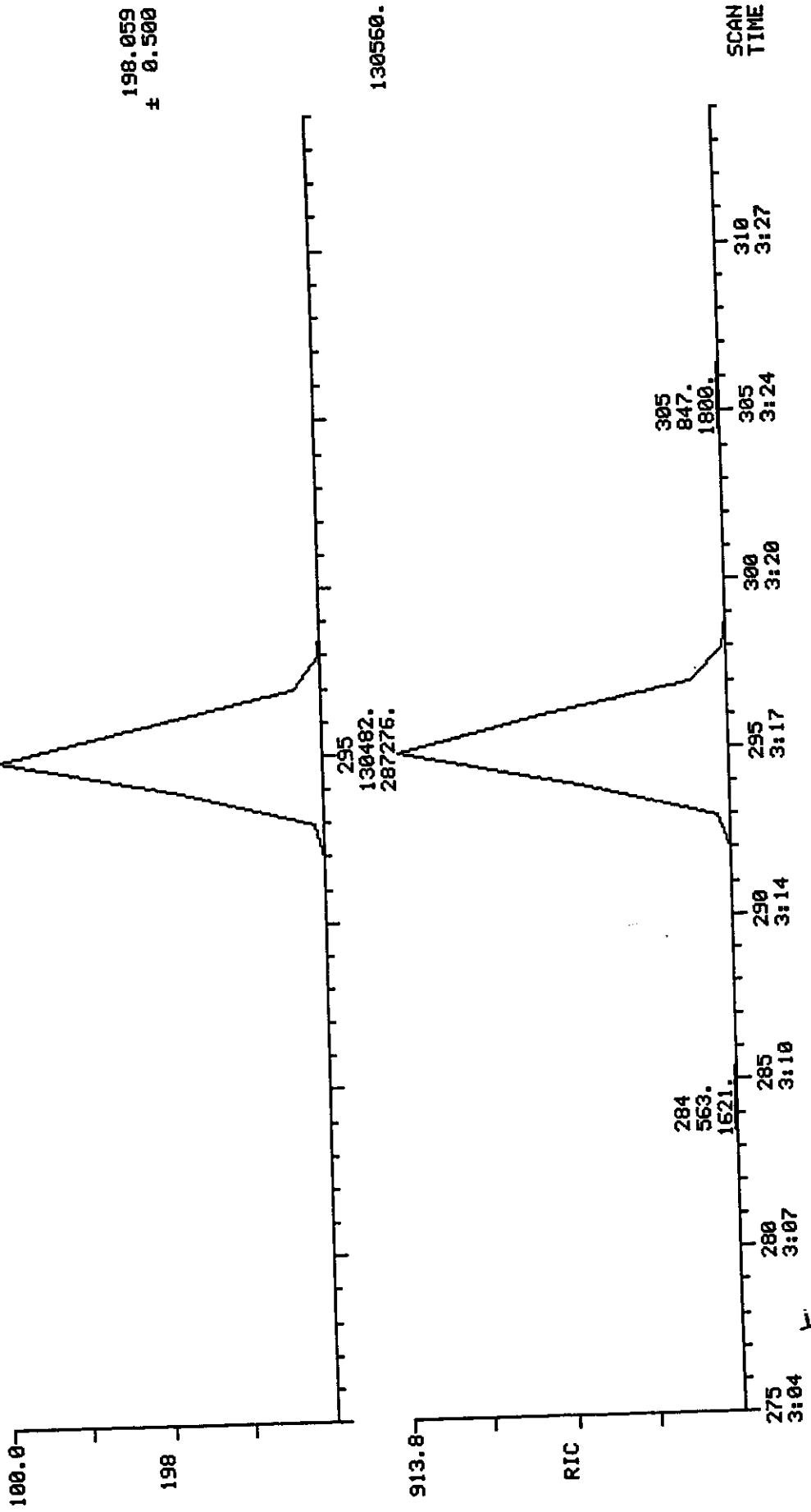
DATA: U16980827 #294  
CALI: F16980821 #3

RIC+MASS CHROMATOGRAM  
08/27/98 9:12:00  
SAMPLE: 50UG/ML DFTPP 081398A  
CONDS.: INST. ID: F16  
RANGE: G 1.3000 LABEL: N 2. 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3

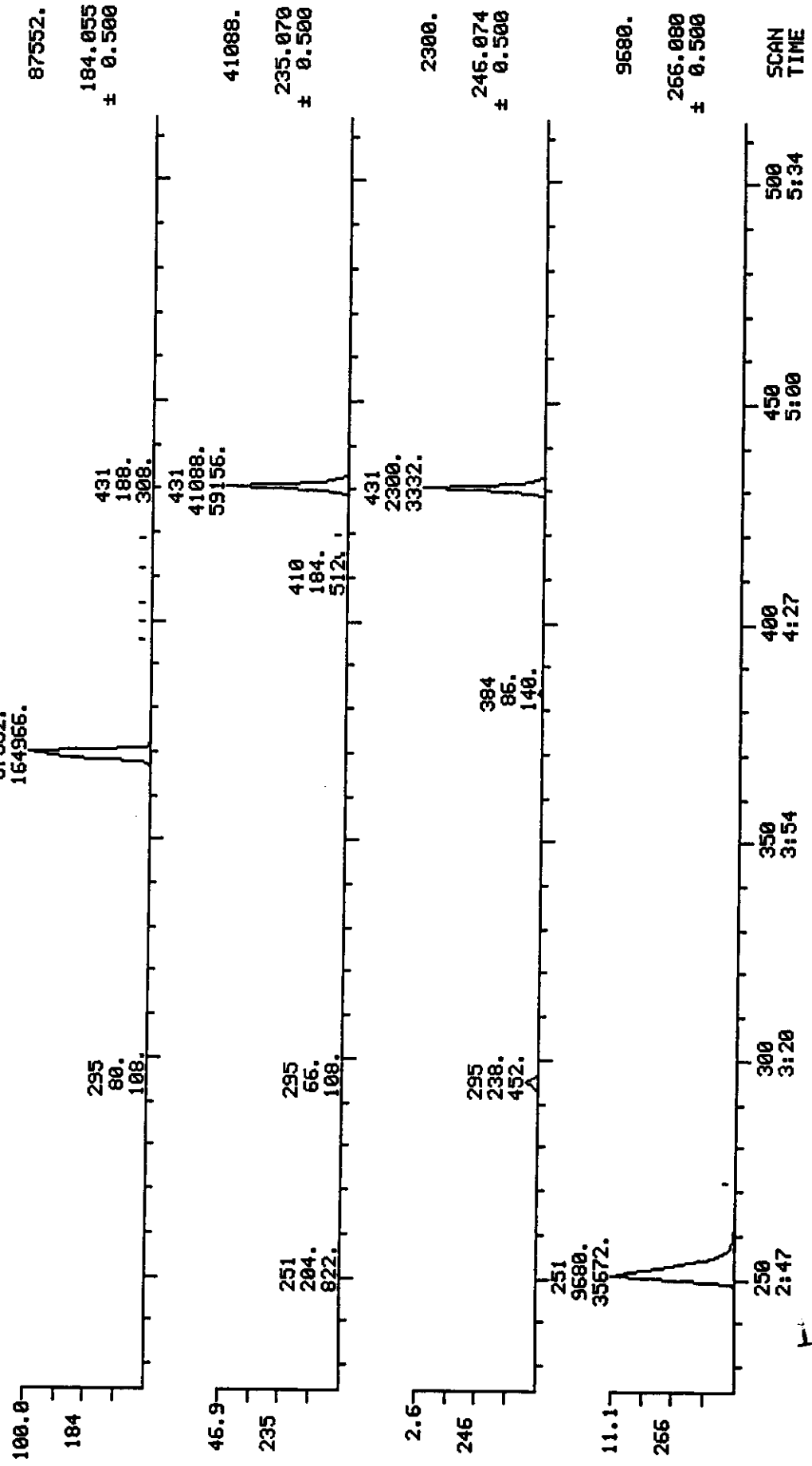
14288.

198.059  
± 0.500

130550.



MASS CHROMATOGRAMS  
 08/27/98 9:12:00  
 SAMPLE: 50UG/ML DFTPP 081398A  
 CONDS.: INST. ID: F15  
 RANGE: G 1.3000 LABEL: N 2, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3  
 DATA: U16980827 #294 SCANS 224 TO 514  
 CALI: F16980821 #3



15  
 00

DECAFLUOROTRIPHENYLPHOSPHINE

Tuning Report  
 08/31/98 12:20  
 Instrument: F16  
 # 288  
 Case Number:

Data: U16980831 # 288  
 Cali: U16980831  
 Analyst: DAT  
 Laboratory: QUANTERRA

Base m/z:  
 RIC:  
 Acct. No.:  
 Contract:

m/z	Intensity	% RA	Ion Abundance Criteria			Actual	Status
			Min %	Max %	Mass		
51	16416.	57.0	30.0	60.0	198	57.0	PASS
68	0.	0.0	0.0	2.0	69	0.0	PASS
69	18912.	65.7	0.0	100.0	198	65.7	PASS
70	176.	0.6	0.0	2.0	69	0.9	PASS
127	16416.	57.0	40.0	60.0	198	57.0	PASS
197	0.	0.0	0.0	1.0	198	0.0	PASS
198	28800.	100.0	100.0	100.0	198	100.0	PASS
199	1914.	6.6	5.0	9.0	198	6.6	PASS
275	6792.	23.6	10.0	30.0	198	23.6	PASS
365	756.	2.6	1.0	100.0	198	2.6	PASS
441	3176.	11.0	0.0	100.0	443	85.8	PASS
442	19264.	66.9	40.0	100.0	198	66.9	PASS
443	3700.	12.8	17.0	23.0	442	19.2	PASS

1181

Mass List                      Data: U16980831 # 289      Base m/z: 198  
 08/31/98 12:20:00 + 3:13      Cali: F16980821 # 3      RIC: 263168.  
 Sample: 50UG/ML DFTPP 081398A  
 Conds.: INST. ID: F16  
 #288 to #290 summed - #284

Mass	% RA	Inten.	Minima Maxima Mass	Min Inten: # 0 % RA	293. 0 Inten.
35	0.00	0.			
444					
38?	1.37	394.	156	1.97	568.
39?	5.71	1644.	161	1.22	352.
50?	15.17	4368.	167	4.35	1254.
51?	57.00	16416.	168	2.35	678.
52?	3.11	896.	174	1.11	320.
56?	2.22	638.	175	1.82	524.
57?	4.37	1258.	179	3.29	948.
62?	1.02	294.	180	2.39	688.
63?	2.19	632.	181	1.20	346.
65?	1.46	420.	185	1.72	494.
69	65.67	18912.	186	11.75	3384.
74	5.44	1566.	187	3.52	1014.
75	8.26	2380.	192	1.24	356.
76	3.68	1060.	193	1.17	338.
77	57.67	16608.	196	3.90	1122.
78	4.07	1172.	198	100.00	28800.
79	4.06	1170.	199	6.65	1914.
80	2.90	834.	204	2.83	816.
81	3.93	1132.	205	5.09	1466.
82	1.15	332.	206	20.83	6000.
83	1.29	372.	207	S 2.76	794.
86	1.51	434.	211	1.19	344.
91	S 1.03	296.	217	5.33	1534.
92	1.47	424.	221	7.61	2192.
93	6.01	1732.	223	1.55	446.
98	4.58	1320.	224	10.94	3152.
99	3.60	1036.	225	2.89	832.
101	2.12	612.	227	4.19	1206.
103	1.03	298.	229	1.15	332.
104	1.49	430.	244	9.39	2704.
105	1.18	340.	245	1.45	418.
107	15.17	4368.	246	1.57	452.
108	2.48	714.	255	44.11	12704.
110	31.33	9024.	256	6.56	1890.
111	4.54	1308.	258	2.19	632.
116	1.25	360.	273	1.52	438.
117	10.15	2924.	274	4.37	1258.
118	1.02	294.	275	23.58	6792.
122	1.18	340.	276	3.47	1000.
123	1.53	440.	277	1.63	470.
127	57.00	16416.	296	5.10	1468.
128	4.61	1328.	323	2.00	576.
129	19.56	5632.	334	1.21	348.
130	1.83	528.	365	2.62	756.
135	1.60	460.	372	1.12	322.
137	1.05	302.	423	4.56	1314.
141	2.56	738.	441	11.03	3176.
147	1.44	414.	442	66.89	19264.
148	2.72	784.	443	12.85	3700.
155	1.37	396.	444	1.40	402.

1782



Mass List Data: U16980831 # 289 Base m/z: 198  
08/31/98 12:20:00 + 3:13 Cali: F16980821 # 3 RIC: 263168.  
Sample: 50UG/ML DFTPP 081398A  
Conds.: INST. ID: F16  
#288 to #290 summed - #284

Mass	% RA	Inten.	Minima	Min Inten:	0.
65	0.00	0.	Maxima	#	0
65?	1.46	420.			
67?	0.11	32.			
69	65.67	18912.			
70	0.61	176.			
73	0.85	244.			
74	5.44	1566.			
75	8.26	2380.			

1783

Mass List                      Data: U16980831 # 289      Base m/z: 198  
08/31/98 12:20:00 + 3:13      Cali: F16980821 # 3      RIC: 263168.  
Sample: 5OUG/ML DFTPP 081398A  
Conds.: INST. ID: F16  
#288 to #290 summed - #284

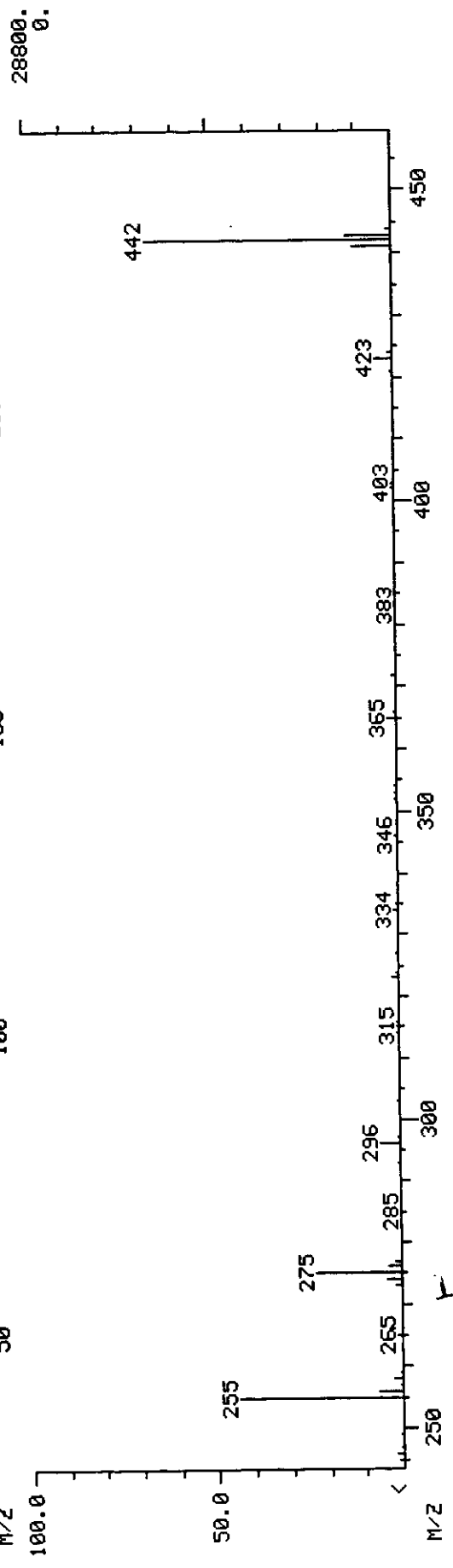
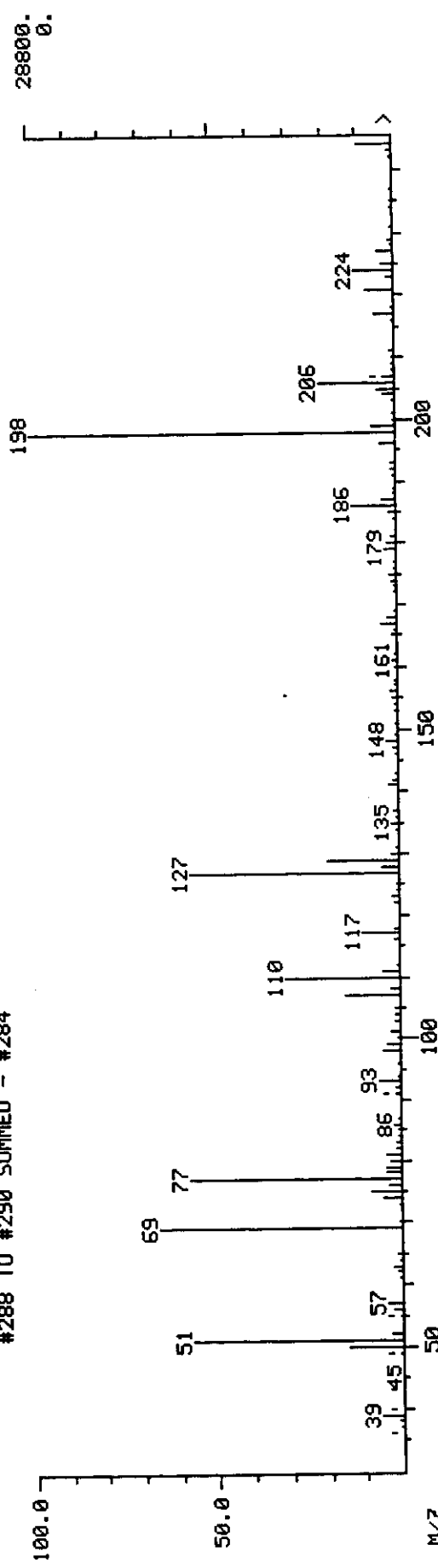
Mass	% RA	Inten.	Minima	Min Inten:	Maxima	#
195	0.00	0.		0.		
200						0
195	0.24	70.				
196	3.90	1122.				
198	100.00	28800.				
199	6.65	1914.				
200	0.72	206.				

1784

DATA: U16980831 #289  
CALI: F16980821 #3

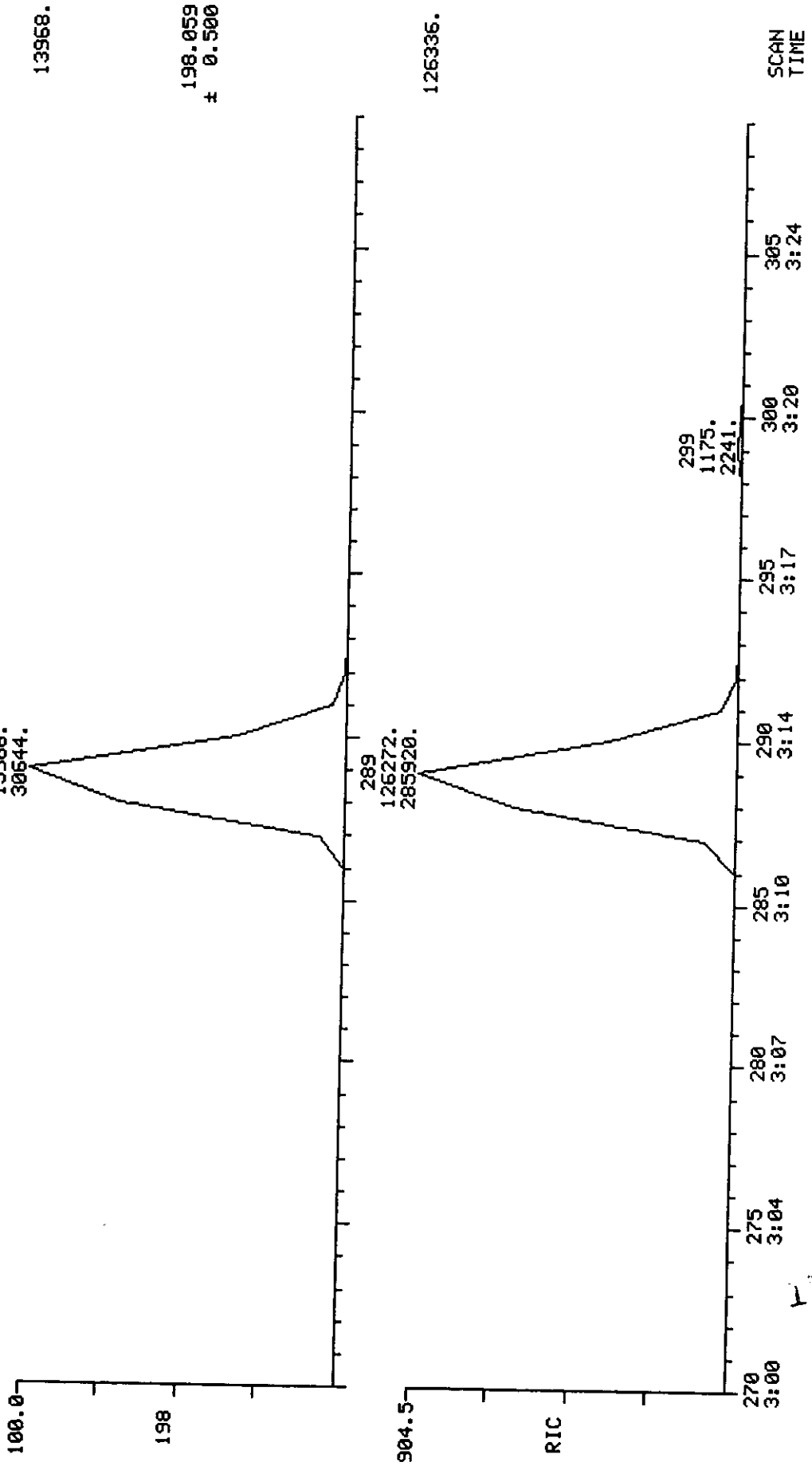
BASE M/Z: 198  
RIC: 263168.

MASS SPECTRUM  
08/31/98 12:20:00 + 3:13  
SAMPLE: 50UG/ML DFTPP 081398A  
CONDS.: INST. ID: F16  
TEMP: 214 DEG. C  
#288 TO #290 SUMMED - #284



F-3  
CO  
CU

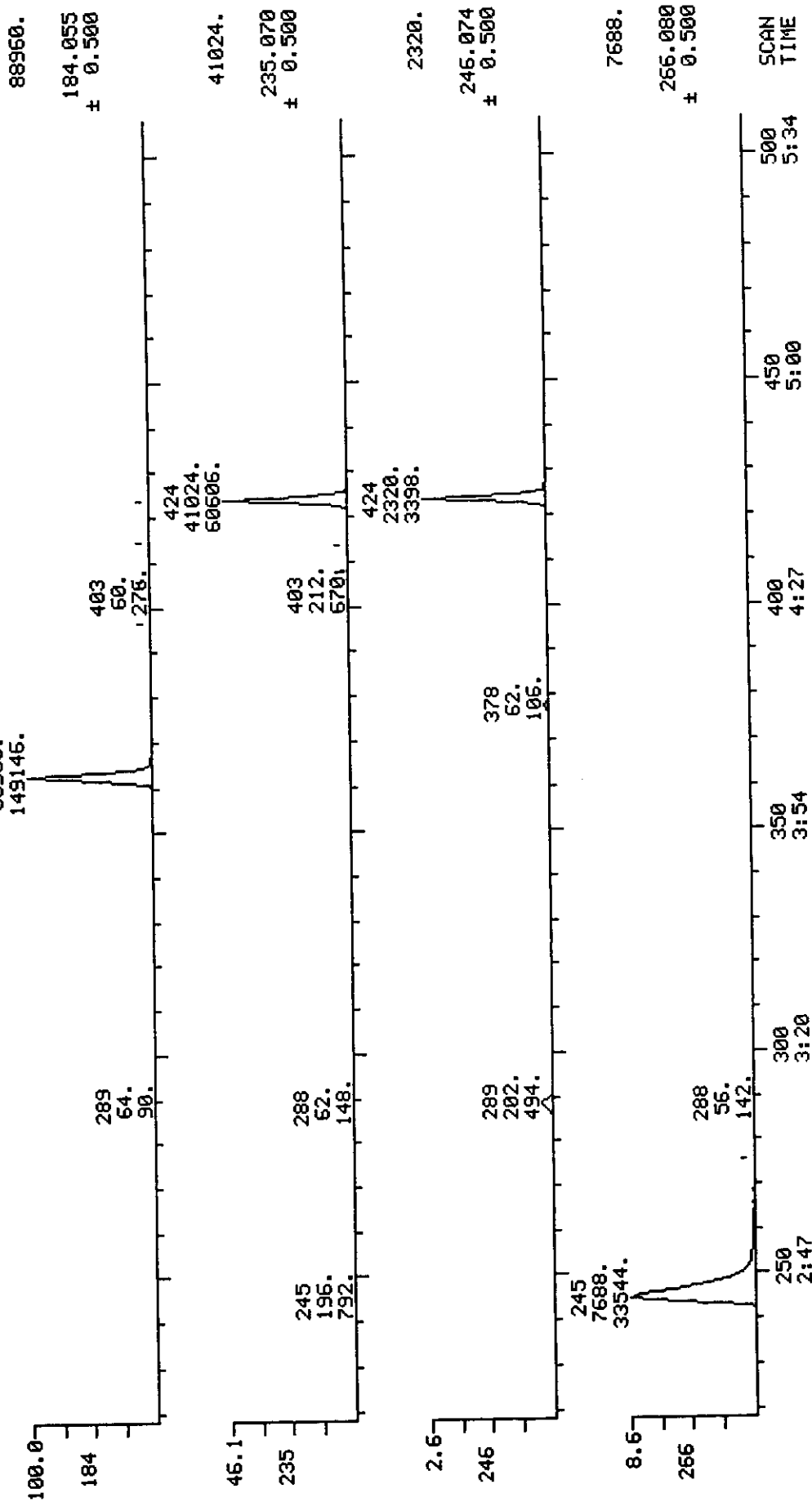
RIC+MASS CHROMATOGRAM  
 08/31/98 12:20:00  
 SAMPLE: 50UG/ML DFTPP 081398A  
 CONDS.: INST. ID: F16  
 RANGE: G 1.3000 LABEL: N 2. 2.0 QUAN: A 5. 2.0 J 0 BASE: U 20. 3  
 DATA: U16980831 #288 SCANS 270 TO 309  
 CALI: F16980821 #3



H-  
 (4)  
 (5)

DATA: U16980831 #288  
 CALL: F16980821 #3

MASS CHROMATOGRAMS  
 08/31/98 12:20:00  
 SAMPLE: 50UG/ML DFTPP 081398A  
 CONDS.: INST. ID: F16  
 RANGE: G 1.3000 LABEL: N 2. 2.0 QUAN: A 6, 2.0 J 0 BASE: U 20, 3



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DECAFLUOROTRIPHENYLPHOSPHINE

Tuning Report  
 09/02/98 14:41  
 Instrument: F16  
 # 288

Data: U16980902 # 288  
 Cali: U16980902  
 Analyst: DAT

Base m/z:  
 RIC:  
 Acct. No.:

Case Number:

Laboratory: QUANTERRA

Contract:

m/z	Intensity	% RA	Ion Abundance Criteria			Actual	Status
			Min %	Max %	Mass		
51	13072.	52.6	30.0	60.0	198	52.6	PASS
68	0.	0.0	0.0	2.0	69	0.0	PASS
69	15232.	61.3	0.0	100.0	198	61.3	PASS
70	126.	0.5	0.0	2.0	69	0.8	PASS
127	13728.	55.3	40.0	60.0	198	55.3	PASS
197	0.	0.0	0.0	1.0	198	0.0	PASS
198	24832.	100.0	100.0	100.0	198	100.0	PASS
199	1688.	6.8	5.0	9.0	198	6.8	PASS
275	6400.	25.8	10.0	30.0	198	25.8	PASS
365	722.	2.9	1.0	100.0	198	2.9	PASS
441	3212.	12.9	0.0	100.0	443	79.8	PASS
442	21024.	84.7	40.0	100.0	198	84.7	PASS
443	4024.	16.2	17.0	23.0	442	19.1	PASS

1728

Mass List                      Data: U16980902 # 289      Base m/z: 198  
 09/02/98 14:41:00 + 3:13      Cali: U16980902 # 3      RIC: 231424.  
 Sample: 50UG/ML DFTPP 081398A  
 Conds.: INST. ID: F16  
 #288 to #290 summed - #284

35	0.00	0.	Minima	Min Inten:	251.
444			Maxima	# 0	
Mass	% RA	Inten.	Mass	% RA	Inten.
39?	S 5.71	1418.	168	2.24	556.
50?	14.30	3552.	174	1.21	300.
51?	52.64	13072.	175	1.90	472.
52?	2.90	720.	177	1.10	272.
56?	2.02	502.	179	3.14	780.
57?	4.16	1032.	180	2.42	600.
63?	1.96	486.	181	1.30	322.
65?	1.36	338.	185	1.78	442.
69	61.34	15232.	186	11.66	2896.
74	4.97	1234.	187	3.70	918.
75	7.72	1918.	192	1.17	290.
76	3.19	792.	193	1.23	306.
77	54.12	13440.	196	3.97	986.
78	3.82	948.	198	100.00	24832.
79	3.93	976.	199	6.80	1688.
80	2.85	708.	204	2.79	692.
81	3.85	956.	205	5.28	1312.
82	1.10	274.	206	21.55	5352.
83	1.06	264.	207	3.09	768.
86	1.33	330.	211	1.68	416.
91	1.19	296.	217	5.56	1380.
92	1.35	334.	221	6.97	1732.
93	6.05	1502.	222	1.29	320.
98	4.46	1108.	223	1.71	424.
99	3.33	826.	224	11.39	2828.
101	2.09	518.	225	3.12	776.
104	1.54	382.	227	4.17	1036.
105	1.36	338.	229	1.15	286.
107	14.11	3504.	244	9.78	2428.
108	2.46	612.	245	1.60	398.
110	28.80	7152.	246	1.81	450.
111	4.19	1040.	255	46.65	11584.
116	1.19	296.	256	7.03	1746.
117	10.07	2500.	258	2.38	590.
123	1.46	362.	273	1.83	454.
125	1.01	252.	274	4.43	1100.
127	55.28	13728.	275	25.77	6400.
128	4.74	1176.	276	3.76	934.
129	18.94	4704.	277	1.79	444.
130	2.00	496.	296	5.38	1336.
135	1.68	418.	323	2.09	520.
137	1.08	268.	334	1.32	328.
141	2.54	630.	365	2.91	722.
147	1.37	340.	372	1.26	314.
148	2.75	684.	423	4.85	1204.
155	1.43	354.	424	1.25	310.
156	1.94	482.	441	12.93	3212.
161	1.31	326.	442	84.66	21024.
165	1.11	276.	443	16.20	4024.
167	4.49	1114.	444	1.80	446.

1783

Mass List                      Data: U16980902 # 289      Base m/z: 198  
09/02/98 14:41:00 + 3:13      Cali: U16980902 # 3      RIC: 231424.  
Sample: 50UG/ML DFTPP 081398A  
Conds.: INST. ID: F16  
#288 to #290 summed - #284

Mass	% RA	Inten.	Minima	Min Inten:	Maxima #
65	0.00	0.		0.	
75					0
65?	1.36	338.			
69	61.34	15232.			
70	0.51	126.			
73	0.74	184.			
74	4.97	1234.			
75	7.72	1918.			

198



Mass List                      Data: U16980902 # 289      Base m/z: 198  
09/02/98 14:41:00 + 3:13      Cali: U16980902 # 3      RIC: 231424.  
Sample: 50UG/ML DFTPP 081398A  
Conds.: INST. ID: F16  
#288 to #290 summed - #284

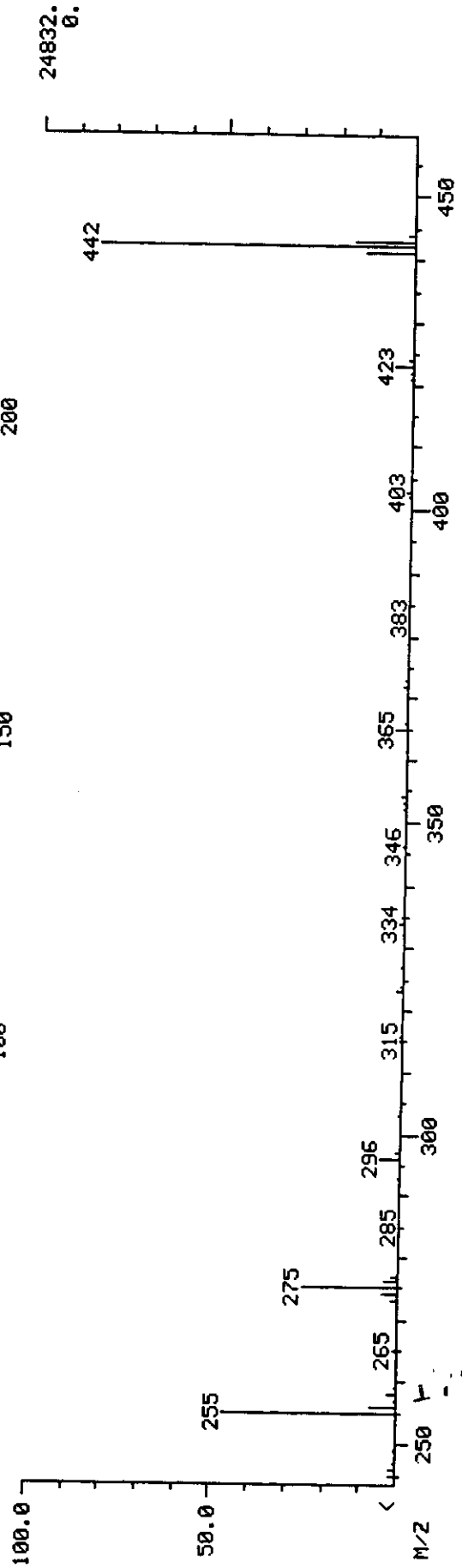
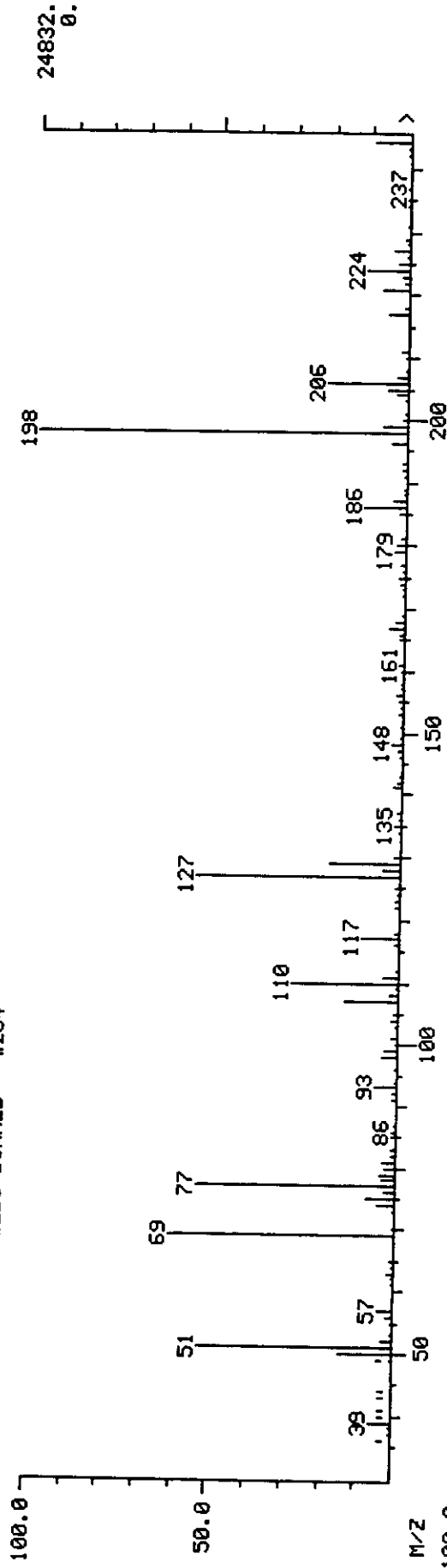
195	0.00	0.	Minima	Min Inten:	0.
200			Maxima	#	0
Mass	% RA	Inten.			
196	3.97	986.			
198	100.00	24832.			
199	6.80	1688.			
200	0.92	228.			

1781

MASS SPECTRUM

09/02/98 14:41:00 + 3:13  
SAMPLE: 50UG/ML DFTPP 081398A  
CONDS.: INST. ID: F16  
TEMP: 215 DEG. C  
#288 TO #290 SUMMED - #284

DATA: U16980902 #289  
CALI: U16980902 #3  
BASE M/Z: 198  
RIC: 231424.

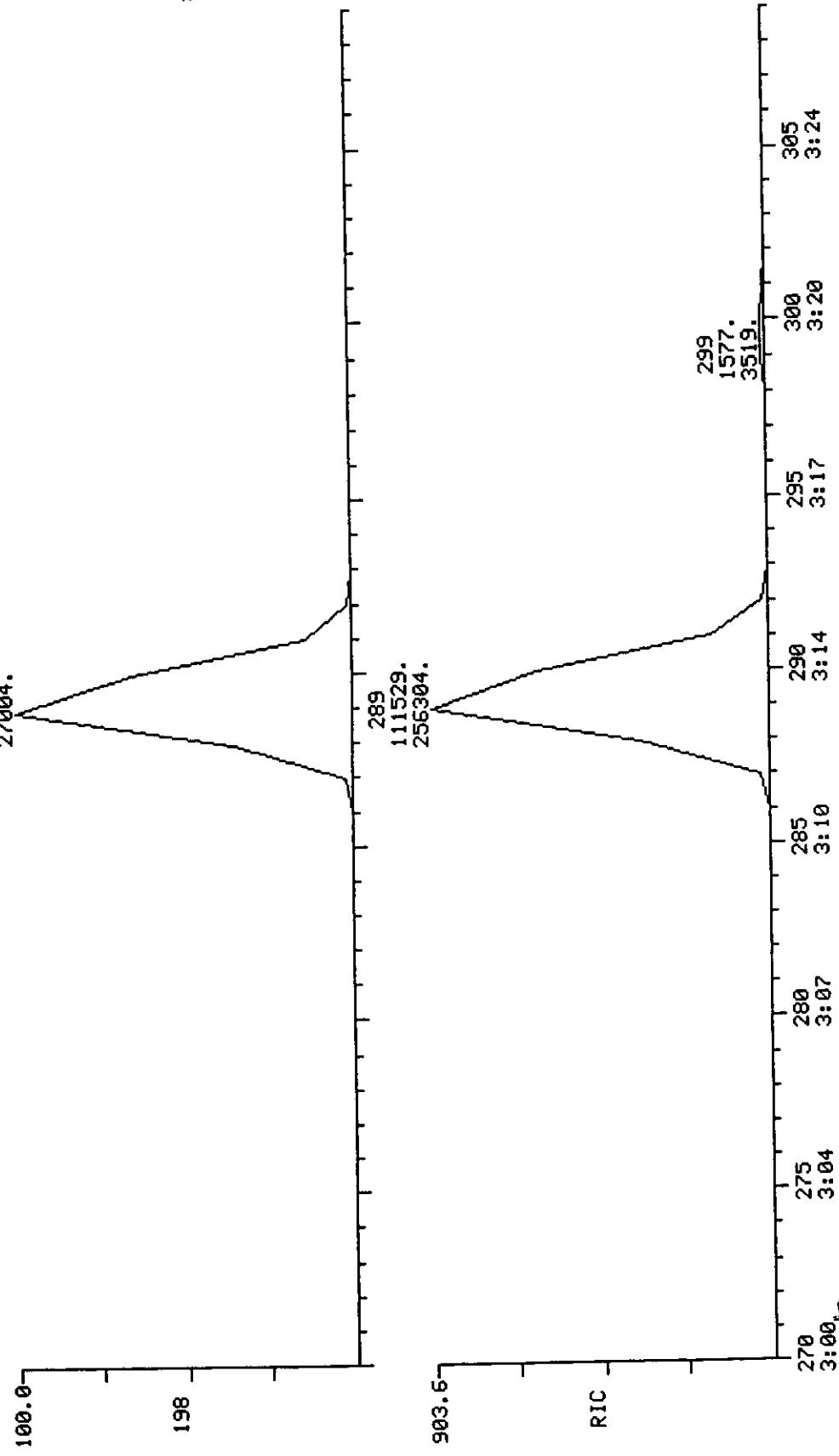


RIC+MASS CHROMATOGRAM  
09/02/98 14:41:00  
SAMPLE: 50UG/ML DFTPP 081398A  
CONDS.: INST. ID: F16  
RANGE: G 1,3000 LABEL: N 2, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3  
DATA: U16980902 #288 SCANS 270 TO 309  
CALI: U16980902 #3

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27004.

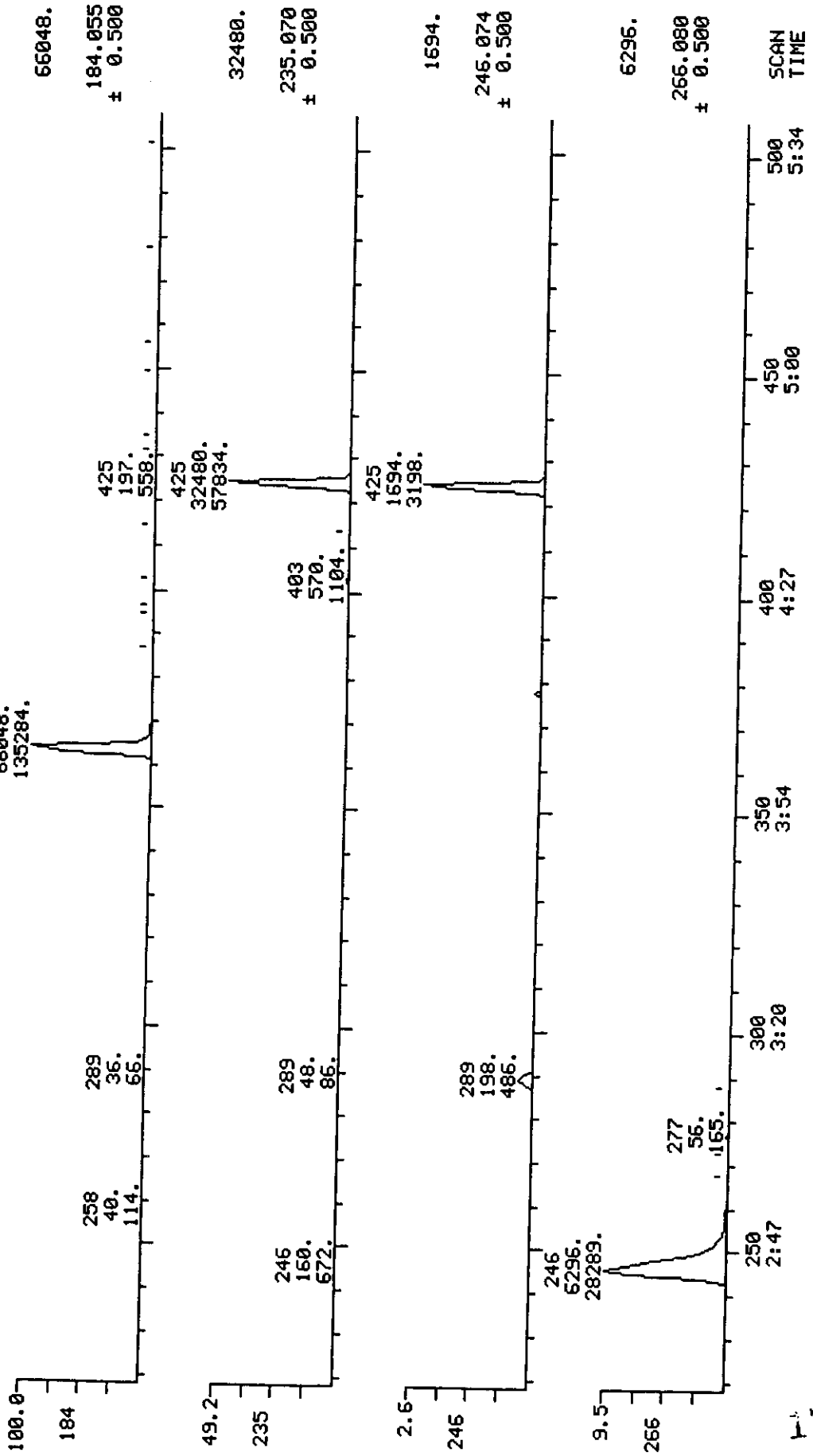
198.059  
± 0.500

111529.  
255304.



270 3:00  
275 3:04  
280 3:07  
285 3:10  
290 3:14  
295 3:17  
300 3:20  
305 3:24

MASS CHROMATOGRAMS  
 09/02/98 14:41:00  
 SAMPLE: 50UG/ML DFTPP 081398A  
 CONDS.: INST. ID: F16  
 RANGE: G 1.3000 LABEL: N 2, 2.0 QUAN: A 5, 2.0 J 0 BASE: U 20, 3  
 DATA: U16980902 #288 SCANS 218 TO 508  
 CALI: U16980902 #3



1104



## TECHNICAL REPORT DATA

Please read instructions on the reverse before completing

1. REPORT NO. EPA-454/R-00-025F	2.	3. RECIPIENT'S ACCESSION NO.
4. TITLE AND SUBTITLE Final Report Hot Mix Asphalt Plants, Truck Loading and Silo Filling, Manual Methods Testing, Asphalt Plant C, Los Angeles, California  Volume 6 of 8	5. REPORT DATE May 2000	
	6. PERFORMING ORGANIZATION CODE	
7. AUTHOR(S) Frank J. Phoenix	8. PERFORMING ORGANIZATION REPORT NO.	
9. PERFORMING ORGANIZATION NAME AND ADDRESS Pacific Environmental Services, Inc. Post Office Box 12077 Research Triangle Park, North Carolina 27709-2077	10. PROGRAM ELEMENT NO.	
	11. CONTRACT/GRANT NO. 68-D-98004	
12. SPONSORING AGENCY NAME AND ADDRESS U.S. Environmental Protection Agency Office of Air Quality Planning and Standards Emissions, Monitoring and Analysis Division Research Triangle Park, North Carolina 27711	13. TYPE OF REPORT AND PERIOD COVERED Final	
	14. SPONSORING AGENCY CODE EPA/200/04	
15. SUPPLEMENTARY NOTES		
16. ABSTRACT  The United States Environmental Protection Agency (EPA) Office of Air Quality Planning and Standards (OAQPS) is investigating hot mix asphalt plants to identify and quantify particulate matter (PM), methylene chloride extractable matter (MCEM), and organic hazardous air pollutant (HAP) emissions during asphalt concrete loading operations. In support of this investigation, the OAQPS issued Pacific Environmental Services, Inc. (PES) a series of work assignments to conduct emissions testing at a hot mix asphalt plant during load-out operations.  The primary objective of the emissions testing was to characterize the uncontrolled emissions of PM, MCEM, polynuclear aromatic hydrocarbons (PAHs), semi-volatile organic hazardous air pollutants (SVOHAPS), and volatile organic hazardous air pollutants (VOHAPS) from a hot mix production plant during loading operations. An asphalt plant south of Los Angeles, California was selected by EPA as the host facility. Testing was performed over five consecutive days beginning on July 24, 1998. Testing was performed under two conditions. Under normal operations, testing was performed to characterize load-out emissions from the tunnel exhaust and load-in emissions from the asphalt concrete storage silo. Under background conditions, testing was performed to characterize emissions from the combustion of diesel fuel in transport trucks.  The entire report consists of eight volumes totaling 4,234 pages, Vol. 1 (388 pages), Vol. 2 (308 pages), Vol. 3 (573 pages), Vol. 4 (694 pages), Vol. 5 (606 pages), Vol. 6 (564 pages), Vol. 7 (570 pages), and Vol. 8 (531 pages).		
17. KEY WORDS AND DOCUMENT ANALYSIS		
a. DESCRIPTIONS	b. IDENTIFIERS/OPEN ENDED TERMS	c. COASTI Field/Group
Hazardous Air Pollutants Methylene Chloride Extractable Matter Particulate Matter Polynuclear Aromatic Hydrocarbons Semi-volatile Organic Hazardous Air Pollutants Volatile Organic Hazardous Air Pollutants		
18. DISTRIBUTION STATEMENT  Unlimited	19. SECURITY CLASS ( <i>This Report</i> ) Unclassified	21. NO. OF PAGES Vol. 6 - 564
	20. SECURITY CLASS ( <i>This page</i> ) Unclassified	22. PRICE