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**PYROLYSIS GAS CHROMATOGRAPHY**  
**ANALYSIS OF 3 THERMO-LAG**  
**FIRE BARRIER SAMPLES**

**Performed For:**

**Entergy Operations, Inc.**  
**Waterford 3**  
**P.O. Box B**  
**Killona, LA 70066**

**P.O. No. NWC0035**

**30 May 1995**

**Distribution**

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**NEI: Biff Bradley (1)**

**NUCON: 06LP829 Master File (1)**  
**Lab (1)**

**NUCON 06LP829/01**

I. ABSTRACT

Inspection of the pyrograms of 3 Thermo-Lag fire barrier samples indicated that they are all similar in chemical composition. However, sample 0395-37B, a 3-hour rated panel sample had disproportionate amounts of characteristic compounds.

II. OBJECTIVE

Pyrolysis Gas Chromatography (PGC) with Mass Selective Detection (MSD) was used to qualitatively compare three Thermo-Lag fire barrier samples.

III. DESCRIPTION OF METHOD

The samples were compared by pyrolysis gas chromatography using ASTM D3452 as a general guide. A Hewlett-Packard model 5890 series II gas chromatograph equipped with a Hewlett Packard model 5972 mass selective detector was used to generate chromatograms of the pyrolysis products. Pyrolysis of the Thermo-Lag samples were performed with a CDS pyroprobe mounted in an independently heated interface attached to the injection port of the GC. Analysis involved weighing 1-3 mgs. of sample in a quartz tube and placement of the tube in the platinum coil element of the probe. The probe is then placed in the interface and pyrolysed ballistically for 2 seconds. Pyrolytic products are then swept by the carrier gas onto the fused silica capillary column where they are separated and detected with a MSD. Chromatographic and pyrolysis conditions are shown in Table 1. Prior to each analysis, the column is heated to 250°C to elute any volatiles which were not entrained in the polymer.

IV. PRESENTATION OF RESULTS

The three pyrograms for each of the three Thermo-Lag samples are shown in Figures 1, 3 and 5. The extracted ion chromatograms using the acrylate base ion  $m/e$  of 55 common to ethyl acrylate (EA) and base ion  $m/e$  of 69 common to methyl methacrylate (MMA) for each sample are shown in Figures 2, 4 and 6. The EA/MMA area ratios and densities for each sample are shown in Table 2. Following each set of figures a library search is attached which identifies some of the major peaks for each sample followed by a summary area percent report.

## V. DISCUSSION OF RESULTS

The average extracted ion area ratio of  $1.34 \pm 0.1$  ( $\pm \sigma$ ) for EA/MMA shown in Table 2 is consistent with the average ratio of  $1.3 \pm 0.1$  ( $\pm \sigma$ ) obtained from other Thermo-Lag samples.

The extracted ion chromatograms (Figure 2) for sample 0395-37A, a 3 hour rated panel sample, have an EA/MMA ratio of 1.43. Pyridine compounds identified in the pyrogram (Figure 1) are 3-methyl pyridine and 3, 4-dimethyl pyridine. Other key components identified are pentanedioic acid diethyl ester, trimethyl phenyl phosphate, and octicizer.

The extracted ion chromatograms (Figure 4) for sample 0395-37B, a 3 hour rated panel sample, have an EA/MMA ratio of 1.38. Pyridine compounds were not identified in the pyrogram (Figure 3) although visual inspection shows trace amounts present. Other key components identified are pentanedioic acid diethyl ester, octicizer, triphenyl phosphate and trimethylphenyl phosphate. This sample showed a much higher content of organic phosphate compounds than other Thermo-Lag samples.

Sample 0395-37B was retested and the extracted ion chromatograms shown in Figure 7 have an EA/MMA ratio of 1.29. Pyridine compounds identified in the pyrogram (Figure 8) and library search (Figures 9, 10, 11 and 12) are 3-methyl pyridine, 3, 5-dimethyl pyridine and 2, 5-dimethyl pyridine. Other key components identified are pentanedioic acid diethyl ester (Figure 13) 2, 3, 4, 5-tetramethyl 1-H-pyrrole (Figure 14) and octicizer (Figure 15).

The extracted ion chromatograms (Figure 6) for sample 0395-37C, a 3 hour rated panels sample, have an EA/MMA ratio of 1.21. Pyridine compounds identified in the pyrogram (Figure 6) are pyridine, 3-methyl pyridine, 2, 5-dimethyl pyridine, 3-ethenyl pyridine, 3, 5-dimethyl pyridine, 2, 3, 5-trimethyl pyridine and 3-ethyl-5-methyl pyridine. Other key components identified are 2, 3, 4, 5-tetramethyl 1H-pyrrole, pentanedioic acid diethyl ester, octicizer, triphenyl phosphate and trimethyl phenyl phosphate.

In conclusion, inspection of the pyrograms for the Thermo-Lag samples 0395-37A and C indicates that they are consistent with other Thermo-Lag samples in terms of chemical composition. Sample 0395-37C has a much larger quantity of organic phosphate compounds than other Thermo-Lag samples. This may be due to selection of the subsample used for analysis. The heterogeneity of this material is evidenced by the two sets of results for sample 0395-37B, where, in the first test, only trace amounts of pyridine compounds are found but a large amount of octicizer is present, while in the second test of the same sample only a small amount of octicizer is present.

**TABLE 1**

**Chromatographic Conditions:**

30 meter 0.25 mm narrow bore fused silica HP-5 CB capillary column.

Carrier Gas: Helium, 0.9 mL/min, split ratio 35:1

**Column Conditions:**

Initial Temperature: 50°C for 1 minute hold

Temperature Ramp: 8°C/min to 250°C

Final Temperature: Hold at 250°C for 10 minutes

Injector Temperature: 250°C

Detector Temperature: 280°C

Detector was an HP MSD in scan mode (30-550 amu)

**Pyrolysis Conditions:**

Pyrolysis Temperature: 650°C

Interval: 2 seconds

Ramp: 2°C/millisecond

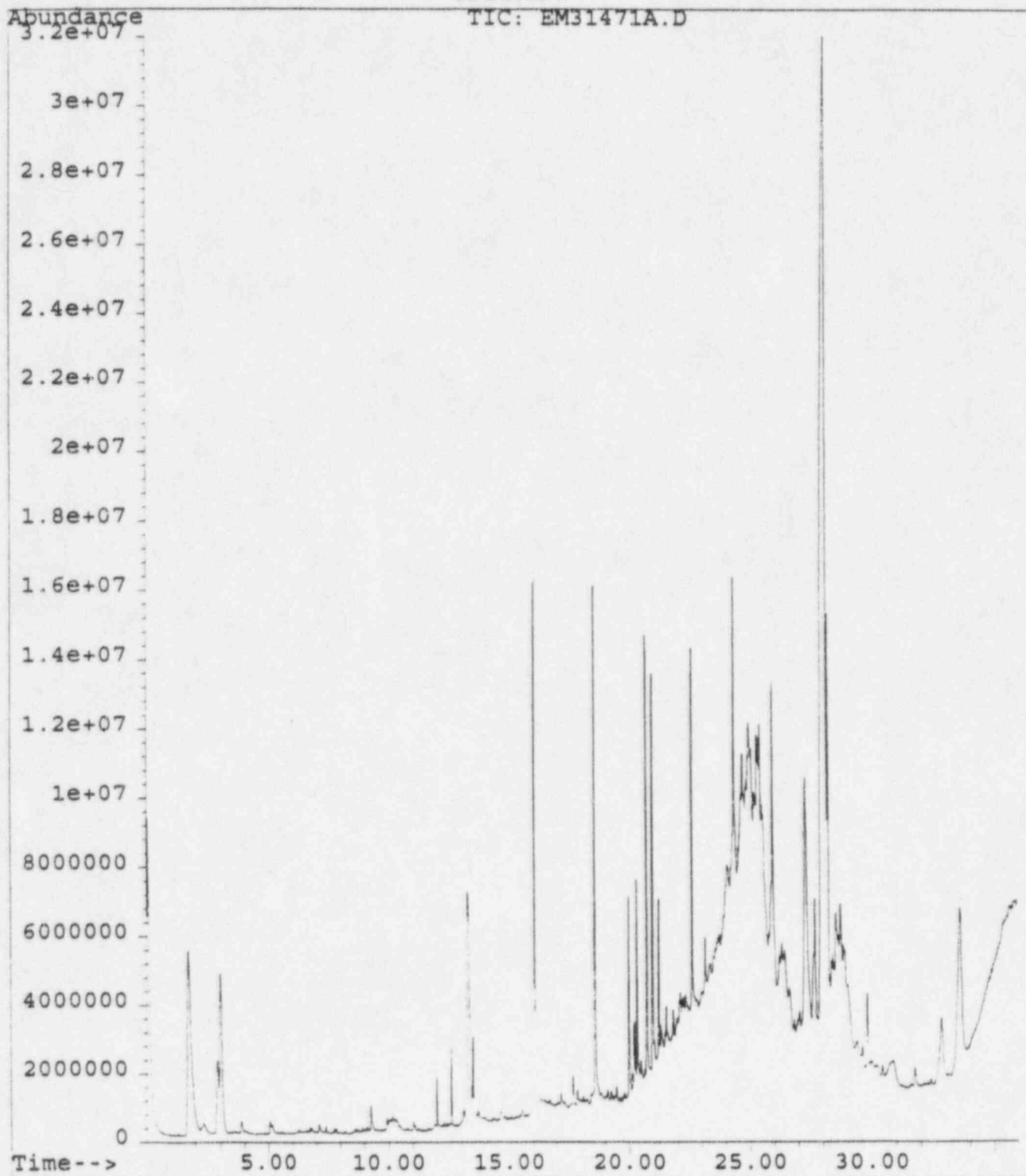
Probe Type: Platinum Coil

Interface Temperature: 205°C

TABLE 2		
Sample	EA/MMA Ratio	Density
NUCON Lab Log # 0395-37A Pre-fabricated, 3 hour rated panel, trowel grade and top coat. Location: RAB + 21 elev. diesel generator Room A FD-77-Fire Area, RAB 16	1.43	1.36 g/mL
NUCON Lab Log # 0395-37B Pre-fabricated, 3 hour rated panel trowel grade and top coat. Location: RAB-4 elev. boric acid concentrator Room A FD-177, Fire Area RAB 31	1.38	1.20 g/mL
NUCON Lab Log # 0395-37C Pre-fabricated 3 hour rated panel, trowel grade and top coat. Location RAB + 21 elev. in front of tool room FD-3HV-B217B- Fire area RAB 25	1.21	1.50 g/mL

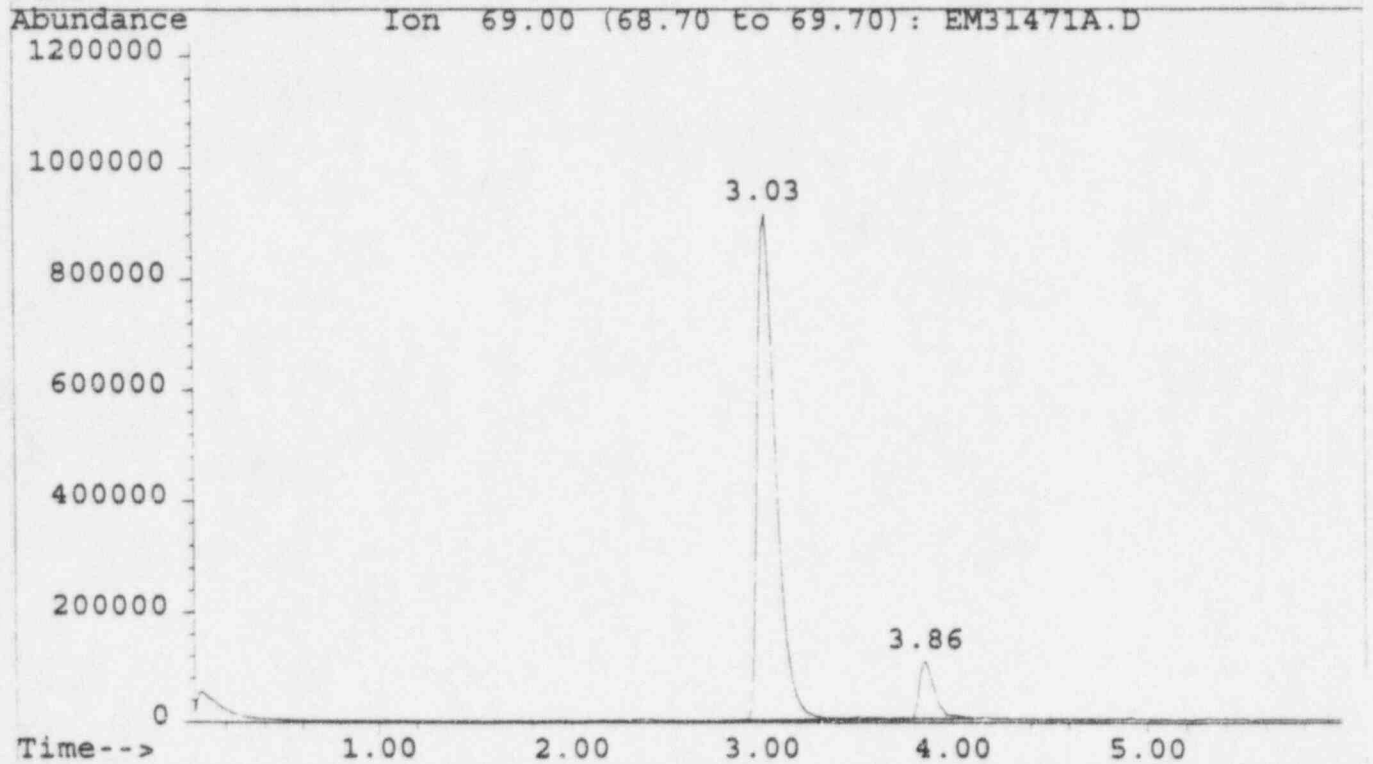
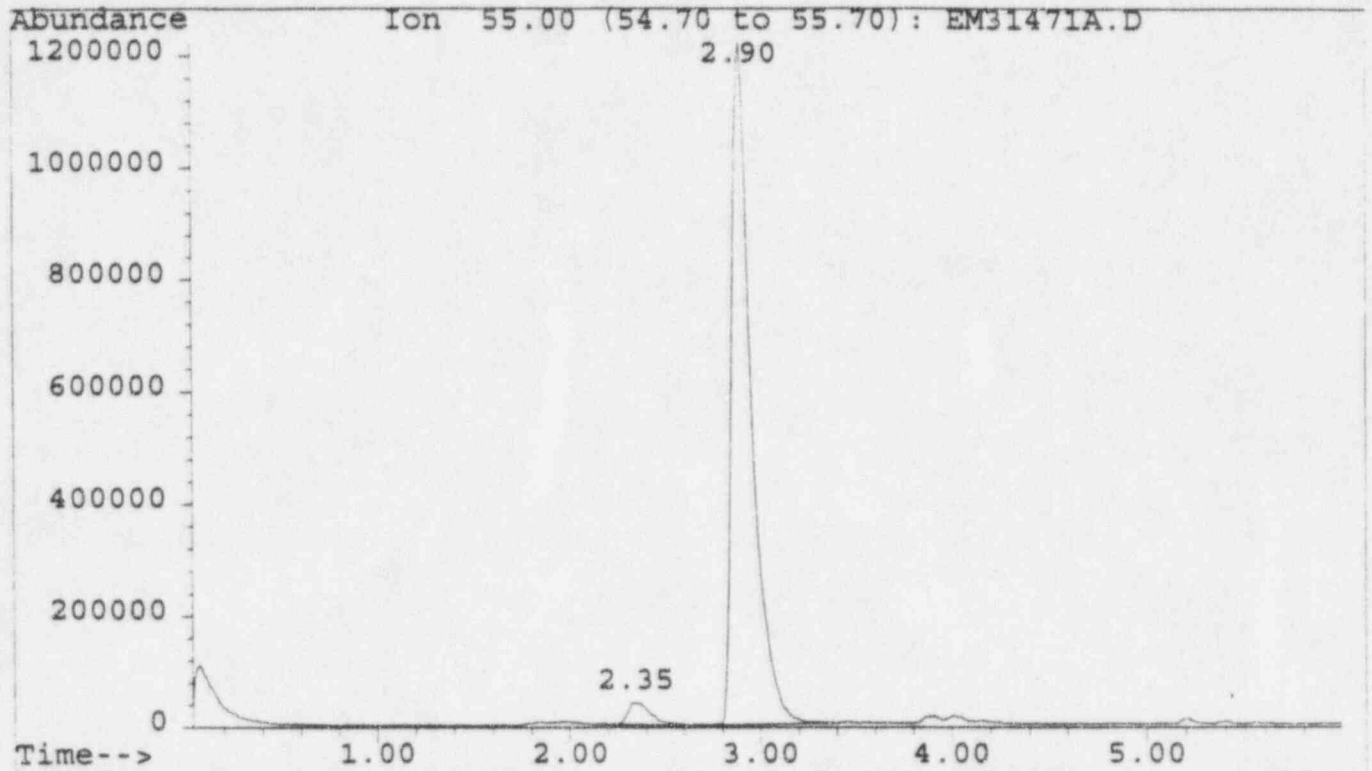
File : C:\HPCHEM\1\DATA\EM31471A.D  
Operator : Marti  
Acquired : 13 Apr 95 12:57 pm using AcqMethod EM31492  
Instrument : 35972 - 33  
Sample Name: 0A95-37a 12 MAR 95 WFA  
Misc Info : Thermolag sample  
Vial Number: 1

FIGURE 1



File : C:\HPCHEM\1\DATA\EM31471A.D  
Operator : Marti  
Acquired : 13 Apr 95 12:57 pm using AcqMethod EM31492  
Instrument : 35972 - 33  
Sample Name: 0295-37a 12/12/95 wpt  
Misc Info : Thermolag sample  
Vial Number: 1

FIGURE 2



Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471A.D  
 Operator : Marti  
 Acquired : 13 Apr 95 12:57 pm using AcqMethod EM31492  
 Sample Name: 0495-37a  
 Misc Info : Thermolag sample  
 Vial Number: 1

Search Libraries: C:\DATABASE\NBS75K.L Minimum Quality: 0

Unknown Spectrum: Apex  
 Integration Params: AutoIntegrate

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
1	1.74	1.89	C:\DATABASE\NBS75K.L No matches found			
2	2.33	0.11	C:\DATABASE\NBS75K.L Xanthatin, 8-[4-[[isopropylamino) Benzeneethanamine, N-methyl- Amphetamine	55737 6311 6317	000000-00-0 000589-08-2 000300-62-9	12 10 10
3	3.03	1.37	C:\DATABASE\NBS75K.L 2-Propenoic acid, 2-methyl-, methy 2-Propenoic acid, 2-methyl-, methy 2-Propenoic acid, 2-methyl-, methy	63329 63330 1484	000080-62-6 000080-62-6 000080-62-6	68 64 64
4	3.87	0.07	C:\DATABASE\NBS75K.L 2-Butenoic acid, 4-nitrophenyl est 2-Propenoic acid, 2-methyl-, ethyl 2-Propenoic acid, 2-methyl-, ethyl	24526 2900 64113	014617-88-0 000097-63-2 000097-63-2	50 45 41
5	5.08	0.08	C:\DATABASE\NBS75K.L Pyridine, 3-methyl- Pyridine, 3-methyl- Pyridine, 3-methyl-	981 63046 63045	000108-99-6 000108-99-6 000108-99-6	92 64 64
6	7.13	0.03	C:\DATABASE\NBS75K.L Pyridine, 3,4-dimethyl- Pyridine, 3,5-dimethyl- Pyridine, 3,4-dimethyl-	63728 2047 2050	000583-58-4 000591-22-0 000583-58-4	90 90 90
7	9.27	0.07	C:\DATABASE\NBS75K.L 2,3-Pyridinediamine Butanamide, N-(4-hydroxyphenyl)- 2(1H)-Pyridinone, 1-methyl-	2194 17435 63813	000452-58-4 000101-91-7 000694-85-9	47 43 43
8	10.17	0.17	C:\DATABASE\NBS75K.L Cyclopentasiloxane, decamethyl- Benzeneacetic acid, .alpha.,3,4-tr Ergoline-8-carboxamide, 9,10-dideh	51266 74564 37179	000541-02-6 037148-65-5 000478-94-4	76 43 27
9	10.29	0.11	C:\DATABASE\NBS75K.L Cyclopentasiloxane, decamethyl- Benzoic acid, 2,6-bis[(trimethylsi Silane, [[4-[1,2-bis[(trimethylsil	51266 51296 57393	000541-02-6 003782-85-2 056114-62-6	76 38 35



Pk#	RT	Area#	Library/ID	Ref#	CAS#	Qual
10	11.03	0.03	C:\DATABASE\NBS75K.L 3-Acetyl-1-methylpyrrole 1H-Pyrrole, 3-ethyl-2,4-dimethyl- Acetamide, N-(4-methoxyphenyl)-	4027 64742 67892	000932-62-7 000517-22-6 000051-66-1	52 52 52
11	12.01	0.12	C:\DATABASE\NBS75K.L Pentanedioic acid, 2-methyl-, dime Pentanedioic acid, diethyl ester 2,4,5-Tri-O-acetyl-3-O-methyl-6-de	68429 69156 42895	014035-94-0 000818-38-2 000000-00-0	38 12 12
12	12.28	0.03	C:\DATABASE\NBS75K.L Benzaldehyde, 2,4,6-trinitro- 5-Bromo-2-iodosobenzoic acid Cyanamide, [2-(1-bromo-6a,7,7a,10a	32082 48296 55409	000606-34-8 000000-00-0 055401-69-9	12 8 7
13	12.60	0.16	C:\DATABASE\NBS75K.L Pentanedioic acid, diethyl ester Pentanedioic acid, diethyl ester Pentanedioic acid, 2,4-dimethyl-,	19778 69156 38087	000818-38-2 000818-38-2 057983-50-3	87 59 37
14	13.32	1.63	C:\DATABASE\NBS75K.L Cyclohexasiloxane, dodecamethyl- 2-Chloro-4-(4-methoxyphenyl)-6-(4- Acetic acid, [bis[(trimethylsilyl)	56711 48253 49839	000540-97-6 063673-76-7 053044-27-2	91 35 34
15	13.50	0.27	C:\DATABASE\NBS75K.L 2-(Diethylamino)-3-methylcycloprop 2H-Quinolizin-3-ol, octahydro-, tr Piperidine, 4-methyl-	11187 11200 1421	091295-98-6 015769-36-5 000626-58-4	32 32 30
16	14.64	0.06	C:\DATABASE\NBS75K.L Octadecanoic acid, 2-[(trimethylsi 4-Amino-5-imidazole carboxamide, t Octadecanoic acid, trimethylsilyl	52732 48317 73766	056196-58-8 000000-00-0 018748-91-9	15 11 10
17	15.55	0.05	C:\DATABASE\NBS75K.L Dimethyl phthalate Dimethyl phthalate Dimethyl phthalate	69376 69379 69380	000131-11-3 000131-11-3 000131-11-3	42 42 38
18	16.11	1.74	C:\DATABASE\NBS75K.L 3-Isopropoxy-1,1,1,7,7,7-hexamethy Tetrasiloxane, 3,5-diethoxy-1,1,1, 1,1,1,3,5,7,9,9,9-Nonamethylpentas	60503 59390 48301	071579-69-6 072439-78-2 084409-41-6	37 27 16
19	16.47	0.12	C:\DATABASE\NBS75K.L Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(4-methylphen	73955 51123 51126	001330-78-5 001330-78-5 000078-32-0	92 58 53
20	17.19	0.10	C:\DATABASE\NBS75K.L 1,3,4,6-Hexanetetrone, 1-(4-chloro Ethanone, 1-[4-(1,1-dimethylethyl) Pyridine, 3-(2-thienyl)-	48347 16773 12681	058330-14-6 000943-27-1 021298-53-3	30 27 27

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
21	17.68	0.11	C:\DATABASE\NBS75K.L Butanethioic acid, S-butyl ester Propanoic acid, 2-methyl-, 1-(1,1- 2-Furanmethanol, tetrahydro-	12349 40505 63508	002432-52-2 074381-40-1 000097-99-4	49 47 47
22	17.86	0.11	C:\DATABASE\NBS75K.L Phenol, 4-(2,2,4-trimethylpentyl)- 2H-Imidazo[4,5-b]pyridin-2-one, 1, Phenol, 4-(1,1-dimethylpropyl)-	24397 6253 13527	000000-00-0 016328-62-4 000080-46-6	64 50 50
23	18.10	0.04	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 51123 73955	000563-04-2 001330-78-5 001330-78-5	95 91 87
24	18.59	1.51	C:\DATABASE\NBS75K.L Silane, [[4-[1,2-bis[(trimethylsil 3,4-Dihydroxymandelic acid, ethyl 1,3,5,7-Tetraethyl-1-ethylbutoxysi	74512 55759 56589	056114-62-6 000000-00-0 073420-30-1	55 49 47
25	19.10	0.12	C:\DATABASE\NBS75K.L Butanoic acid, 2-amino-4,4,4-trifl .delta.2-Tetrazaboroline, 5-ethyl- Phenol, 2,2'-[(1-methyl-1,2-ethane	19090 4392 39799	061141-67-1 020534-01-4 000094-91-7	25 11 11
26	19.26	0.06	C:\DATABASE\NBS75K.L 5-Acetoxymethyl-2-furaldehyde Butanoic acid, 2-amino-4,4,4-trifl Phenol, 2-[(1-methylethyl)thio]-	14397 19090 14457	010551-58-3 061141-67-1 029549-62-0	11 11 11
27	19.48	0.10	C:\DATABASE\NBS75K.L Phenol, 2,2'-[(1-methyl-1,2-ethane Bendazol 1-Benzopyrylium, 2-phenyl-	39799 24958 24651	000094-91-7 000621-72-7 014051-53-7	25 18 14
28	19.80	0.06	C:\DATABASE\NBS75K.L Phosphoric acid, tris(4-methylphen Phenothiaphosphine, 2,8-dimethyl-1 Phosphoric acid, tris(methylphenyl	51126 51105 51123	000078-32-0 023861-49-6 001330-78-5	95 45 42
29	20.01	0.57	C:\DATABASE\NBS75K.L 1,1'-Biphenyl, 3-azido- 3-Cyclobutene-1,2-dicarboxylic aci 7,8-Dihydro-2-methyl(6H)pyrazolo[3	21582 29069 27965	014213-01-5 055673-94-4 000000-00-0	20 18 16
30	20.22	0.38	C:\DATABASE\NBS75K.L cis-2,4,5-Trimethoxy-.beta.-methyl 1,3-Diphenyl-(4H)1,2,4-triazoline- 4-Hexen-2-yn-1-one, 1-phenyl-5-(1-	34481 34507 34549	000000-00-0 000000-00-0 029743-43-9	30 27 25
31	20.34	0.61	C:\DATABASE\NBS75K.L 1H-Purine, 6-(methylthio)- Rheadan-8-ol, 2,3,10,11-tetrametho 7,8-Dihydro-2-methyl(6H)pyrazolo[3	67932 54012 27965	000050-66-8 014028-91-2 000000-00-0	27 25 25

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
32	20.48	0.25	C:\DATABASE\NBS75K.L Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(4-methylphen Phosphoric acid, tris(methylphenyl	51123 51126 73955	001330-78-5 000078-32-0 001330-78-5	97 93 91
33	20.74	1.49	C:\DATABASE\NBS75K.L .beta.-D-Glucopyranosiduronic acid Morphinan, 7,8-didehydro-4,5-epoxy Estra-1,3,5(10)-trien-17-one, 2-[(	62040 55885 55893	052092-53-2 055449-66-6 077883-26-2	43 43 38
34	20.97	1.43	C:\DATABASE\NBS75K.L Benzenamine, N,N,3-trimethyl- Acetic acid, (4-formylphenoxy)-, e 3-Pyridinecarbonitrile, 1,4-dihydr	65619 24812 6098	000121-72-2 051264-69-8 000767-98-6	14 14 14
35	21.24	1.32	C:\DATABASE\NBS75K.L 4-Amino-2-methyl-5,6-trimethylenep 1H-Purin-6-amine, N-methyl- 4-Pyridinecarboxaldehyde, 3-hydrox	9460 66644 14236	076881-49-7 000443-72-1 000066-72-8	25 20 18
36	21.55	1.09	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 51123 73955	000563-04-2 001330-78-5 001330-78-5	95 89 86
37	21.83	0.63	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(4-methylphen Phosphoric acid, tris(methylphenyl	51124 51126 51123	000563-04-2 000078-32-0 001330-78-5	91 86 76
38	22.13	1.34	C:\DATABASE\NBS75K.L Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl Phenothiaphosphine, 2,8-dimethyl-1	51123 73955 51105	001330-78-5 001330-78-5 023861-49-6	90 64 46
39	22.25	0.63	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(4-methylphen	51124 73955 51126	000563-04-2 001330-78-5 000078-32-0	94 87 86
40	22.36	1.20	C:\DATABASE\NBS75K.L Phosphoric acid, tris(4-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51126 73955 51123	000078-32-0 001330-78-5 001330-78-5	95 92 45
41	22.64	2.68	C:\DATABASE\NBS75K.L 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Benzene, 1-phenyl-4-(2-cyano-2-phe 14.alpha.-Morphinan, 7,8-didehydro	56714 39643 39631	038147-00-1 027869-56-3 017939-34-3	27 10 10
42	23.18	2.32	C:\DATABASE\NBS75K.L 2,4,6,8,9,10-Hexathiatricyclo[3.3.	46703	057289-12-0	16
43	23.38	1.30	C:\DATABASE\NBS75K.L 2,4(1H)-Cyclo-3,4-secoakuummilan-1 2-Propanol, 1-[2-(2-methoxy-1-meth Butanamide, 3-methyl-	52355 24251 1621	006472-42-0 020324-33-8 000541-46-8	35 27 25

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
44	23.76	2.96	C:\DATABASE\NBS75K.L Pentane, 2-methoxy- Hydrazine, 1-methyl-1-(2-methylpro Butanamide, 3-methyl-	1778 1757 1621	006795-88-6 020240-63-5 000541-46-8	43 38 38
45	24.11	4.29	C:\DATABASE\NBS75K.L 2,4,6,8,9,10-Hexathiatricyclo[3.3. Butanamide, 3-methyl- 3-Pentanol, 2-methyl-	50037 1621 63561	057274-38-1 000541-46-8 000565-67-3	38 38 38
46	24.40	4.55	C:\DATABASE\NBS75K.L Benzeneacetic acid, .alpha.,3,4-tr 1,3,5,7-Tetraethyl-1-ethylbutoxysi Benzoic acid, 2,4-bis[(trimethylsi	74565 56589 73975	037148-65-5 073420-30-1 010586-16-0	32 30 25
47	24.74	5.03	C:\DATABASE\NBS75K.L 1-Propanol, 2-(2-methoxy-1-methyle Butane, 2-methoxy- 1-Propanol, 2-(2-methoxypropoxy)-	9239 854 9232	055956-21-3 006795-87-5 013588-28-8	43 38 37
48	25.02	7.01	C:\DATABASE\NBS75K.L 1-Propanol, 2-(2-methoxy-1-methyle 2-Naphthalenemethanol, decahydro-. Butanamide, 3-methyl-	9239 28205 1621	055956-21-3 051317-08-9 000541-46-8	47 37 35
49	25.42	8.21	C:\DATABASE\NBS75K.L 3-Pentanol, 2-methyl- Propane, 1-ethoxy-2-methyl- Butanamide, 3-methyl-	63561 63544 1621	000565-67-3 000627-02-1 000541-46-8	43 43 38
50	25.98	3.89	C:\DATABASE\NBS75K.L Morphinan, 7,8-didehydro-4,5-epoxy 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Estra-1,3,5(10)-trien-17-one, 2-[(	55885 56714 55893	055449-66-6 038147-00-1 077883-26-2	46 42 27
51	26.34	1.81	C:\DATABASE\NBS75K.L 2,4(1H)-Cyclo-3,4-secoakuammilan-1 Benzo[1,2-c:4,5-c']dipyrrole-1,3,5 Benzo[1,2-c:4,5-c']dipyrrole-1,3,5	52355 53562 53561	006472-42-0 031663-79-3 031663-75-9	30 11 11
52	26.46	1.28	C:\DATABASE\NBS75K.L Bikaverin 2,4(1H)-Cyclo-3,4-secoakuammilan-1 Sarpagan-16-carboxylic acid, 3,17-	52326 52355 52356	033390-21-5 006472-42-0 053632-75-0	59 35 22
53	26.65	1.35	C:\DATABASE\NBS75K.L 2-Propanol, 1-(2-ethoxypropoxy)- 2-Pentanone, 5-methoxy- Ethane, 1-ethoxy-1-methoxy-	12860 64282 1926	010143-32-5 017429-04-8 010471-14-4	47 47 47
54	27.06	0.89	C:\DATABASE\NBS75K.L Amylene Hydrate 2-Pentanol, 2-methyl- 2-Propanol, 1-(2-ethoxypropoxy)-	62940 1763 12860	000075-85-4 000590-36-3 010143-32-5	47 38 38

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
55	27.33	3.58	C:\DATABASE\NBS75K.L 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Morphinan, 7,8-didehydro-4,5-epoxy .beta.-D-Glucopyranosiduronic acid	56714 55885 62040	038147-00-1 055449-66-6 052092-53-2	45 42 27
56	27.70	1.76	C:\DATABASE\NBS75K.L 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Benzoic acid, 2,4-bis[(trimethylsi Benzoic acid, 2,5-bis(trimethylsil	56714 73975 73978	038147-00-1 010586-16-0 003618-20-0	38 15 10
57	28.16	14.62	C:\DATABASE\NBS75K.L Octicizer Benz[b]acridine, 1,2,3,4,7,8,9,10- Benz[c]acridine, 1,2,3,4,8,9,10,11	50542 34157 34155	001241-94-7 055044-74-1 055044-73-0	38 23 23
58	28.58	2.76	C:\DATABASE\NBS75K.L 1-Propanol, 2-(2-methoxy-1-methyle 1,2-Propanediol, 2-methyl-, 1-meth 2,5,8,11-Tetraoxadodecane	9239 14343 17074	055956-21-3 074792-80-6 000112-49-2	47 38 38
59	28.76	3.59	C:\DATABASE\NBS75K.L 1-Propanol, 2-(2-methoxy-1-methyle Butane, 1-echoxy- 2-Butanol, 2,3-dimethyl-	9239 63550 63570	055956-21-3 000628-81-9 000594-60-5	43 38 35
60	29.41	0.60	C:\DATABASE\NBS75K.L Pentane, 2-methoxy- Ethanedioic acid, dimethyl ester 1-Chloro-2-ethoxyethane	1778 3458 2079	006795-88-6 000553-90-2 000628-34-2	25 25 25
61	29.63	0.29	C:\DATABASE\NBS75K.L 1,2-Benzenedicarboxylic acid, mono 1,2-Benzenedicarboxylic acid, 3-ni Bis(2-ethylhexyl) phthalate	39132 25513 53128	004376-20-9 000603-11-2 000117-81-7	43 43 43
62	29.85	0.50	C:\DATABASE\NBS75K.L Benzoic acid, 2,4-bis[(trimethylsi Benzoic acid, 2,5-bis(trimethylsil Silane, [[4-[1,2-bis[(trimethylsil	73975 51295 74512	010586-16-0 003618-20-0 056114-62-6	20 18 16
63	30.43	0.15	C:\DATABASE\NBS75K.L Tanshinone II-b 4H-1-Benzopyran-2-carboxylic acid, 3-Aminodiftalone	44270 39264 39292	017397-93-2 030192-14-4 067483-30-1	39 22 20
64	30.86	0.63	C:\DATABASE\NBS75K.L Phenol, dimethyl-, phosphate (3:1) Cholesta-5,7-dien-3-one, 4,4-dimet C(14a)-Homo-27-norgammacer-14-ene	54605 54634 54655	025155-23-1 000000-00-0 018046-86-1	51 12 10
65	31.77	0.11	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 73955 51123	000563-04-2 001330-78-5 001330-78-5	90 86 49

Fk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
66	32.41	0.02	C:\DATABASE\NBS75K.L			
			Phosphoric acid, tris(methylphenyl	51123	001330-78-5	91
			1H-12a,5-(Epoxyethano)pyrido[3',4	51134	003329-92-8	25
			Phosphoric acid, tris(4-methylphen	51126	000078-32-0	25
67	32.86	0.59	C:\DATABASE\NBS75K.L			
			1,1,1,5,7,7,7-Heptamethyl-3,3-bis(	56714	038147-00-1	12
			Heptasiloxane, hexadecamethyl-	59695	000541-01-5	12
			Benzoic acid, 2,5-bis(trimethylsil	73978	003618-20-0	11
68	33.13	0.01	C:\DATABASE\NBS75K.L			
			2-Naphthalenemethanol, decahydro-	28205	051317-08-9	47
			Heptane, 1-ethoxy-	8517	001969-43-3	30
			Cyclooctanemethanol, .alpha.,.alph	15325	016624-06-9	27
69	33.62	1.62	C:\DATABASE\NBS75K.L			
			1,1,1,5,7,7,7-Heptamethyl-3,3-bis(	56714	038147-00-1	38
			Heptasiloxane, hexadecamethyl-	59695	000541-01-5	14
			Benzoic acid, 2,5-bis(trimethylsil	73978	003618-20-0	11
70	35.69	0.21	C:\DATABASE\NBS75K.L			
			2-Propanol, 1-(2-methoxypropoxy)-	9233	013429-07-7	38
			Pentane, 2-methoxy-	1778	006795-88-6	38
			Butanamide, 3-methyl-	1621	000541-46-8	38
71	35.88	0.11	C:\DATABASE\NBS75K.L			
			Butanamide, 3-methyl-	1621	000541-46-8	38
			2-Propanol, 1-(2-methoxypropoxy)-	9233	013429-07-7	38
			Propanoic acid, 2-hydroxy-2-methyl	3494	002110-78-3	38

Area Percent Report -- Sorted by Signal

Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471A.D  
 Operator : Marti  
 Acquired : 13 Apr 95 12:57 pm using AcqMethod EM31492  
 Sample Name: 0495-37a  
 Misc Info : Thermolag sample  
 Vial Number: 1  
 CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

Retention Time	Area	Area %	Ratio %
Total Ion Chromatogram			
1.742	601393536	1.891	12.936
2.335	35461107	0.112	0.763
3.027	434519950	1.366	9.347
3.866	22070112	0.069	0.475
5.078	26666286	0.084	0.574
7.130	10865117	0.034	0.234
9.274	22612605	0.071	0.486
10.173	53471448	0.168	1.150
10.290	34352204	0.108	0.739
11.032	10335885	0.032	0.222
12.002	38570794	0.121	0.830
12.280	9140554	0.029	0.197
12.604	51331931	0.161	1.104
13.312	516844108	1.625	11.118
13.494	85651901	0.269	1.842
14.644	19755558	0.062	0.425
15.543	14726153	0.046	0.317
16.107	554815668	1.745	11.934
16.466	39303151	0.124	0.845
17.190	31161994	0.098	0.670
17.677	35868588	0.113	0.772
17.864	34835411	0.110	0.749
18.094	14137921	0.044	0.304
18.587	479371689	1.507	10.312
19.104	36818758	0.116	0.792
19.259	20506049	0.064	0.441
19.478	31975393	0.101	0.688
19.799	20146385	0.063	0.433
20.004	181939663	0.572	3.914
20.225	122243376	0.384	2.630
20.336	193466228	0.608	4.162
20.483	80764082	0.254	1.737
20.733	474225524	1.491	10.201
20.979	454847358	1.430	9.784
21.232	418319486	1.315	8.998

Area Percent Report -- Sorted by Signal

Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471A.D  
 Operator : Marti  
 Acquired : 13 Apr 95 12:57 pm using AcqMethod EM31492  
 Sample Name: 0495-37a  
 Misc Info : Thermolag sample  
 Vial Number: 1  
 CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

Retention Time	Area	Area %	Ratio %
21.545	347601288	1.093	7.477
21.833	199517775	0.627	4.292
22.133	427224089	1.343	9.190
22.254	200848722	0.632	4.320
22.361	381247456	1.199	8.201
22.642	852348617	2.680	18.334
23.180	737361875	2.319	15.861
23.385	414625472	1.304	8.919
23.767	940298835	2.957	20.226
24.113	1363159995	4.286	29.322
24.393	1447207021	4.550	31.130
24.743	1599518140	5.029	34.406
25.016	2228418622	7.007	47.934
25.417	2609952331	8.207	56.141
25.980	1237946489	3.893	26.629
26.340	577189096	1.815	12.416
26.455	406990370	1.280	8.755
26.649	428636001	1.348	9.220
27.058	281595199	0.885	6.057
27.327	1139525626	3.583	24.512
27.703	559735188	1.760	12.040
28.171	4648891930	14.618	100.000
28.587	878228811	2.761	18.891
28.766	1142368693	3.592	24.573
29.410	190822799	0.600	4.105
29.633	90861797	0.286	1.954
29.845	159718577	0.502	3.436
30.429	47831864	0.150	1.029
30.861	201454672	0.633	4.333
31.768	34752574	0.109	0.748
32.415	6300133	0.020	0.136
32.860	187537080	0.590	4.034
33.121	4663432	0.015	0.100
33.620	515022573	1.619	11.078
35.686	67680920	0.213	1.456
35.880	33755208	0.106	0.726



Area Percent Report -- Sorted by Signal

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Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471A.D  
Operator : Marti  
Acquired : 13 Apr 95 12:57 pm using AcqMethod EM31492  
Sample Name: 0495-37a  
Misc Info : Thermolag sample  
Vial Number: 1  
CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

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Retention Time	Area	Area %	Ratio %
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Ion 55.00 (54.70 to 55.70): EM31471A.D  
0495-37a

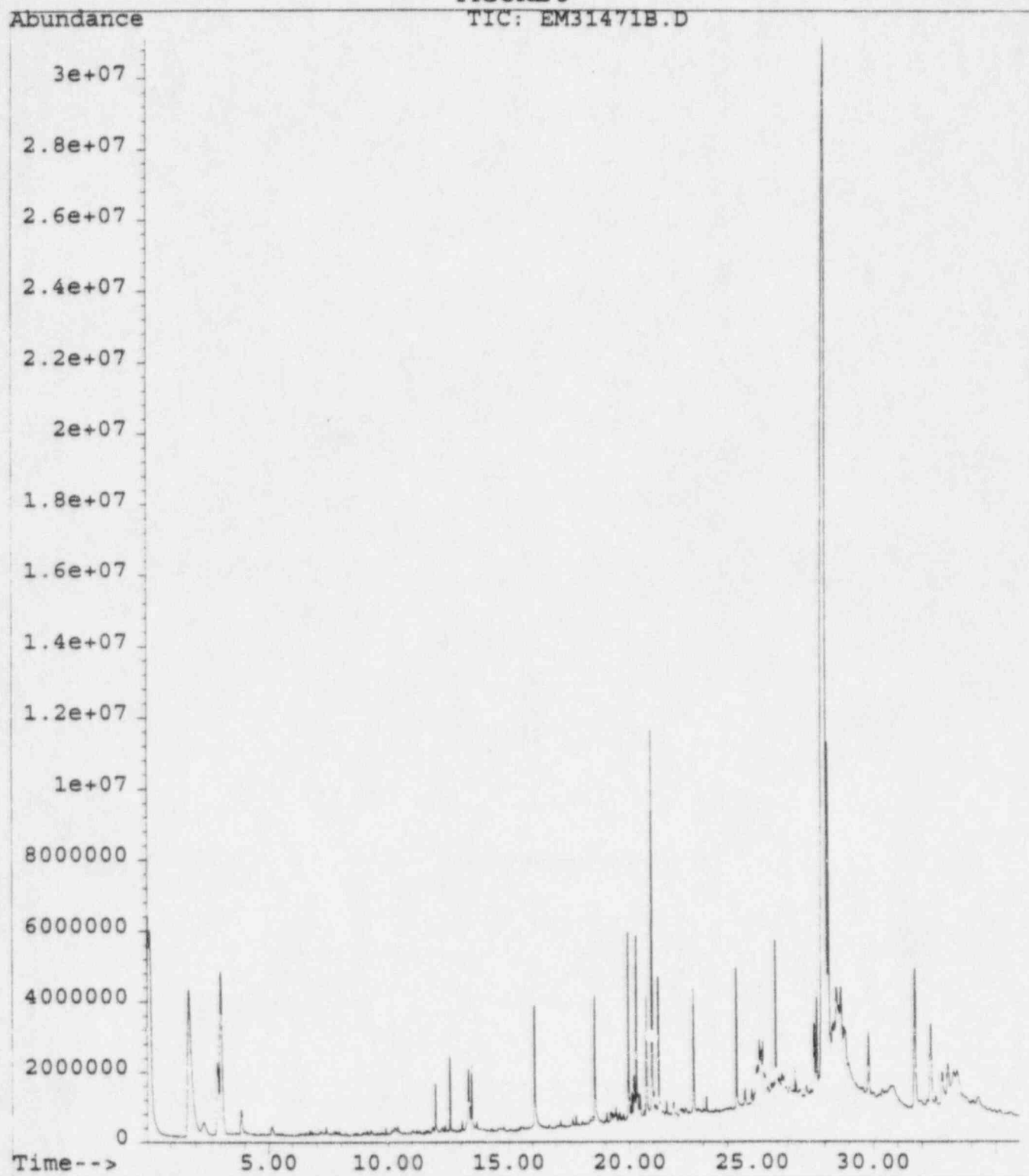
Peak#	Ret Time	Type	Width	Area	Start Time	End Time
1	2.345	BB	0.128	3286432	2.136	2.721
2	2.898	BB	0.109	86746091	2.751	3.386

Ion. 69.00 (68.70 to 69.70): EM31471A.D  
0495-37a

Peak#	Ret Time	Type	Width	Area	Start Time	End Time
1	3.028	BV	0.103	60439623	2.820	3.425
2	3.862	VV	0.086	5842676	3.425	4.100

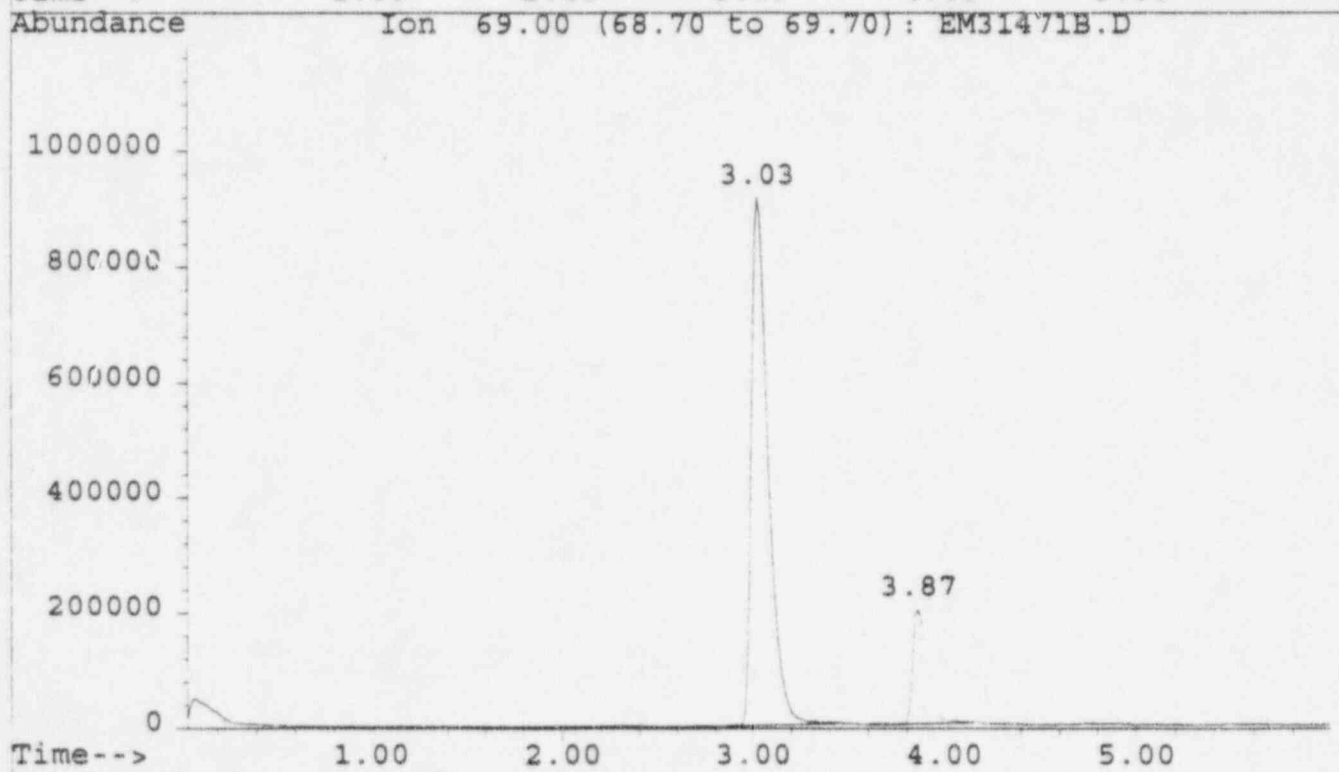
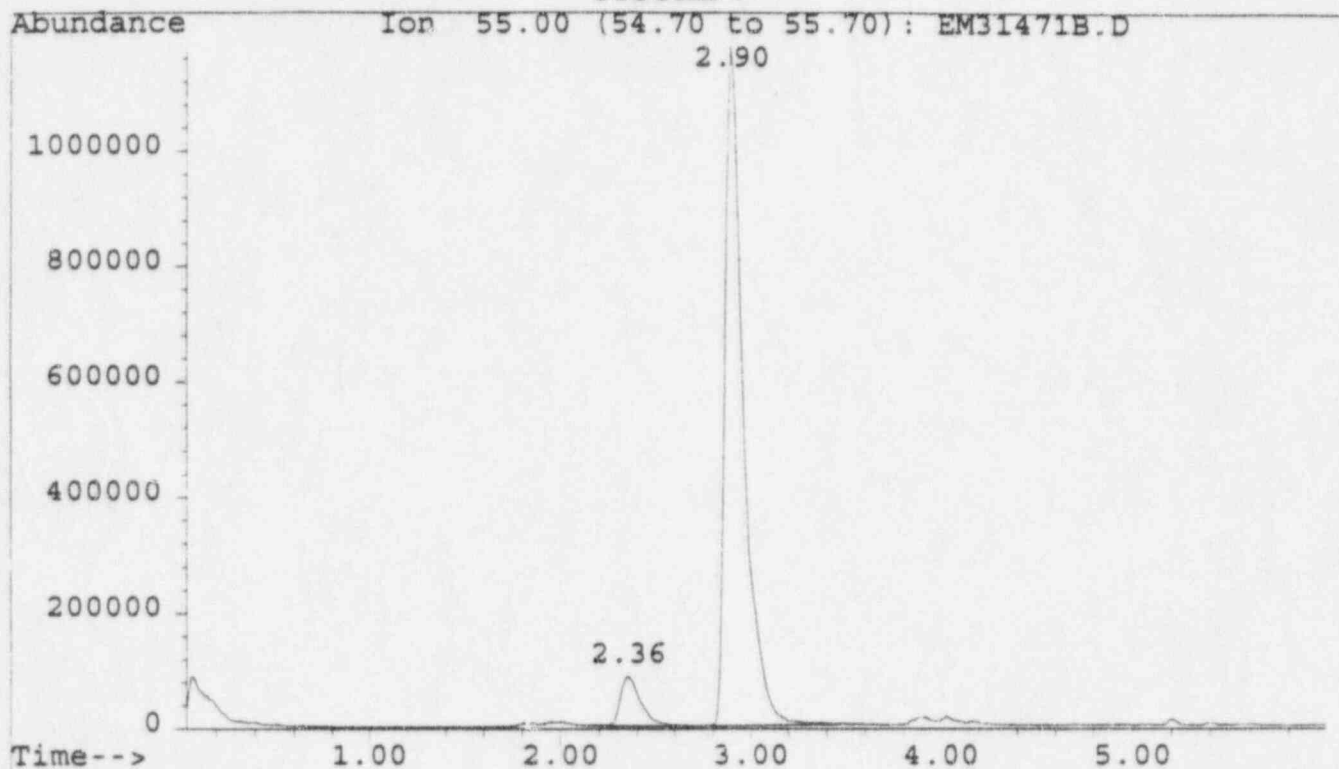
File : C:\HPCHEM\1\DATA\EM31471B.D  
Operator : Marti  
Acquired : 13 Apr 95 2:25 pm using AcqMethod EM31492  
Instrument : 5972 - 33  
Sample Name: 0495-37b WPC 12 MAY 95  
Misc Info : Thermolag sample  
Vial Number: 1

FIGURE 3



File : C:\HPCHEM\1\DATA\EM31471B.D  
Operator : Marti  
Acquired : 13 Apr 95 2:25 pm using AcqMethod EM31492  
Instrument : 5972 - 33  
Sample Name: 0495-37b *WVF 12 MAR 95*  
Misc Info : Thermolag sample  
Vial Number: 1

FIGURE 4



Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471B.D  
 Operator : Marti  
 Acquired : 13 Apr 95 2:25 pm using AcqMethod EM31492  
 Sample Name: 0495-37b W.F. 11/17/95  
 Misc Info : Thermolag sample  
 Vial Number: 1

Search Libraries: C:\DATABASE\NBS75K.L Minimum Quality: 0

Unknown Spectrum: Apex  
 Integration Params: AutoIntegrate

Pk#	RT	Area*	Library/ID	Ref#	CAS#	Qual
1	0.16	-12.73	C:\DATABASE\NBS75K.L			
			2-Hexanol, 2-methyl-	64355	000625-23-0	35
			2-Hexanol, 2-methyl-	64354	000625-23-0	35
			2-Propanol, 1-[2-(2-methoxy-1-meth	24251	020324-33-8	32
2	1.74	6.46	C:\DATABASE\NBS75K.L			
			No matches found			
3	2.33	0.61	C:\DATABASE\NBS75K.L			
			.beta.-D-Allofuranuronic acid, 5-[	59138	019396-06-6	25
			Meprobamate	70443	000057-53-4	25
			Acetamide, N-(aminocarbonyl)-	1671	000591-07-1	25
4	2.91	1.37	C:\DATABASE\NBS75K.L			
			2-Propenoic acid, ethyl ester	63318	000140-88-5	91
			2-Propenoic acid, ethyl ester	1461	000140-88-5	91
			2-Propenoic acid, ethyl ester	63320	000140-88-5	83
5	3.04	4.05	C:\DATABASE\NBS75K.L			
			2-Propenoic acid, 2-methyl-, methy	63329	000080-62-6	68
			2-Butenoic acid, methyl ester, (E)	1458	000623-43-8	64
			2-Propenoic acid, 2-methyl-, methy	63330	000080-62-6	64
6	3.86	0.49	C:\DATABASE\NBS75K.L			
			2-Propenoic acid, 2-methyl-, ethyl	2900	000097-63-2	64
			2-Propenoic acid, 2-methyl-, ethyl	64113	000097-63-2	60
			2-Butenoic acid, 4-nitrophenyl est	24526	014617-88-0	50
7	5.15	0.20	C:\DATABASE\NBS75K.L			
			p-Xylene	63701	000106-42-3	70
			Benzene, 1,2-dimethyl-	63706	000095-47-6	70
			Benzene, 1,3-dimethyl-	63697	000108-38-3	70
8	7.39	0.10	C:\DATABASE\NBS75K.L			
			1,2,4-Trimethylbenzene	3771	000095-36-3	90
			Benzene, 1,2,3-trimethyl-	64576	000526-73-8	86
			Benzene, 1,3,5-trimethyl-	64570	000108-67-8	86
9	11.98	0.49	C:\DATABASE\NBS75K.L			
			Pentanedioic acid, diethyl ester	69155	000818-38-2	27
			Pentanedioic acid, 2-methyl-, mono	12301	072088-36-9	25
			1,3,2-Dioxaborinane, 4,6-dimethyl-	22867	052964-02-0	22

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
10	12.58	0.57	C:\DATABASE\NBS75K.L Pentanedioic acid, diethyl ester	19778	000818-38-2	87
			Pentanedioic acid, diethyl ester	69156	000818-38-2	64
			Pentanedioic acid, diethyl ester	69155	000818-38-2	58
11	13.08	0.09	C:\DATABASE\NBS75K.L 3-Pentanone, dimethylhydrazone	4996	016795-73-6	30
			3-Piperidinol	63447	006859-99-0	30
			Pyrimidine, 5-chloro-2-dimethylami	11664	081568-09-4	25
12	13.34	1.12	C:\DATABASE\NBS75K.L Cyclohexasiloxane, dodecamethyl-	56711	000540-97-6	93
			Acetic acid, [bis[(trimethylsilyl)	49839	053044-27-2	34
			2,4,6(1H,3H,5H)-Pyrimidinetrione,	49865	052988-92-8	32
13	13.47	0.60	C:\DATABASE\NBS75K.L Piperidine, 4-methyl-	63292	000626-58-4	38
			Piperidine, 4-methyl-	1421	000626-58-4	38
			2-(Diethylamino)-3-methylcycloprop	11187	091295-98-6	32
14	16.10	1.53	C:\DATABASE\NBS75K.L 3-Isopropoxy-1,1,1,7,7,7-hexamethy	60503	071579-69-6	25
			Tetrasiloxane, 3,5-diethoxy-1,1,1,	59390	072439-78-2	22
			1-[2,4-Bis(trimethylsilyloxy)pheny	58102	000000-00-0	16
15	17.17	0.15	C:\DATABASE\NBS75K.L 4,5,6,7-Tetramethylphthalide	20315	029002-54-8	42
			Benzene, 1-(1-ethylpropyl)-4-propy	20374	054789-16-1	38
			Benzene, 1-(1,1-dimethylethyl)-3-e	16799	006630-01-9	38
16	17.66	0.09	C:\DATABASE\NBS75K.L Pentane, 3-bromo-	66684	001809-10-5	38
			Borane, diethyl[1-ethyl-2-(methoxy	21917	053670-48-7	38
			Pentane, 2,3,4-trimethyl-	64229	000565-75-3	38
17	17.81	0.09	C:\DATABASE\NBS75K.L Phenol, 4-(1,1,3,3-tetramethylbuty	24424	000140-66-9	64
			Phenol, 4-(2,2,3,3-tetramethylbuty	24409	054932-78-4	52
			Phenol, 4-(1,1,3,3-tetramethylbuty	70047	000140-66-9	52
18	18.58	1.43	C:\DATABASE\NBS75K.L Silane, [[4-[1,2-bis[(trimethylsil	57393	056114-62-6	55
			1,2-Benzenediol, 4-(2-amino-1-hydr	57358	056145-09-6	47
			Silane, [[4-[1,2-bis[(trimethylsil	74512	056114-62-6	46
19	19.08	0.15	C:\DATABASE\NBS75K.L Phenol, 2,2'-[(1-methyl-1,2-ethane	39799	000094-91-7	42
			5H-Dibenzo[a,d]cyclohepten-5-one,	70114	001210-35-1	41
			Naphthalene, 1,2,3,4-tetrahydro-1-	25032	003018-20-0	41
20	19.22	0.20	C:\DATABASE\NBS75K.L Phenol, 2,2'-[(1-methyl-1,2-ethane	39799	000094-91-7	32
			Ethyl N'-isopropylureidoacetate	19738	000000-00-0	30
			Ethanamine, N-methyl-2-[(2-methylp	34965	015301-93-6	27

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
21	19.45	0.23	C:\DATABASE\NBS75K.L 1H-Inden-1-ol, 2,3-dihydro- Dibenzo[b,e]-1,4-diazabicyclo[2.2. 1,2,4-Oxadiazol-5-amine, 3-phenyl-	65500 24947 12634	006351-10-6 000000-00-0 003663-37-4	18 15 14
22	19.60	0.15	C:\DATABASE\NBS75K.L 5-Bromo-2-iodosobenzoic acid N-Desmethyldiphenhydramine Pheno., 2,2'-[(1-methyl-1,2-ethane	48296 32185 39799	000000-00-0 000000-00-0 000094-91-7	25 25 25
23	19.97	1.72	C:\DATABASE\NBS75K.L Benzo[b]thiophene, 2-(butylthio)- 1,1'-Biphenyl, 3-azido- 3-Cyclobutene-1,2-dicarboxylic aci	28074 21582 29069	054965-46-7 014213-01-5 055673-94-4	15 15 11
24	20.12	0.25	C:\DATABASE\NBS75K.L 2-Iodohistidine 1,4-Oxathiin-3-carboxylic acid, 2- Pyridine, 2-ethyl-4,6-dimethyl-	39529 24692 6309	000000-00-0 117238-98-9 001124-35-2	38 15 14
25	20.21	0.66	C:\DATABASE\NBS75K.L 1,3-Diphenyl-(4H)1,2,4-triazoline- 2,5-Cyclohexadien-1-one, 2,5-dimet Triamterene	34507 34546 34464	000000-00-0 054245-93-1 000396-01-0	38 27 18
26	20.31	1.77	C:\DATABASE\NBS75K.L 7,8-Dihydro-2-methyl(6H)pyrazolo[3 Flavone Urea, 1-neopentyl-3-phenyl-2-thio-	27965 28172 28087	000000-00-0 000525-82-6 015093-39-7	25 18 11
27	20.45	0.32	C:\DATABASE\NBS75K.L Propanedioic acid, bis(2-methyl-2- 1H-Indole-3-ethanamine, 6-fluoro- Acetic acid, oxo-	31923 20703 62541	074793-47-8 077590-52-4 000298-12-4	30 22 11
28	20.71	1.28	C:\DATABASE\NBS75K.L 2H-1,4-Benzodiazepin-2-one, 7-chlo Cyclohexasiloxane, dodecamethyl- Heptasiloxane, hexadecamethyl-	55928 56711 59695	055319-93-2 000540-97-6 000541-01-5	35 14 12
29	20.95	3.99	C:\DATABASE\NBS75K.L Benzene, 1-methyl-4-(nitromethyl)- Pyridine, 2-ethyl-4,6-dimethyl- Pyridine, 2,6-diethyl-	9997 6309 6323	029559-27-1 001124-35-2 000935-28-4	14 14 14
30	21.20	1.24	C:\DATABASE\NBS75K.L 3-(3-Pyridyl)propenoic acid 4-Amino-2-methyl-5,6-trimethylenep 7H-Purin-6-amine, 7-methyl-	9438 9460 9421	001126-74-5 076881-49-7 000935-69-3	22 20 18
31	21.52	0.13	C:\DATABASE\NBS75K.L Cobalt, [[3,3'-(1,2-ethanediyldini 8-Methyl-13,14-dioxo-8H,13H,14H-na 12,13-Dihydro-12-methyl-13,14-diox	54324 46504 46505	036466-12-3 059050-31-6 059050-19-0	11 10 10



Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
32	21.81	0.25	C:\DATABASE\NBS75K.L Pyridine, 3,5-dichloro-4-methoxy-2 1,3-Dioxolane, 2-(2,4-dimethylphen 6-Phenylisoquinoline	24013 27648 24117	051050-42-1 074752-98-0 070125-61-0	49 47 46
33	22.63	1.25	C:\DATABASE\NBS75K.L 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Cyclohexasiloxane, dodecamethyl- 4,5-Ethylene-8,9-dimethoxy-6-phena	56714 56711 39611	038147-00-1 000540-97-6 000000-00-0	30 10 10
34	23.16	0.20	C:\DATABASE\NBS75K.L Isoxazole, 5-(4-methoxyphenyl)-3-p Pregn-5-en-20-one, 3,16-bis(acetyl	34124 56086	003672-51-3 023357-24-6	35 7
35	24.37	1.27	C:\DATABASE\NBS75K.L Benzoic acid, 2,4-bis[(trimethylsi Silane, [[4-[1,2-bis[(trimethylsil Silane, [[4-[1,2-bis[(trimethylsil	51292 74512 57393	010586-16-0 056114-62-6 056114-62-6	42 40 40
36	24.72	0.33	C:\DATABASE\NBS75K.L Ethanol, 2-[2-[4-(1,1,3,3-tetramet Octicizer 6-(Methylamino)phenanthren-3-ol	41793 50542 28427	002315-61-9 001241-94-7 098033-23-9	90 42 38
37	24.99	0.26	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-Hydrazino-8-hydroxy-4-phenylquin	50542 34112 34111	001241-94-7 004928-02-3 000000-00-0	46 27 11
38	25.30	4.23	C:\DATABASE\NBS75K.L 2,5,8,11-Tetraoxatetradecan-13-ol, Propane, 1-ethoxy-2-methyl- 2-Propanol, 1-(2-methoxypropoxy)-	36541 63543 9233	020324-34-9 000627-02-1 013429-07-7	43 38 38
39	25.65	0.27	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Octicizer 2-Hydrazino-8-hydroxy-4-phenylquin	34112 50542 34111	004928-02-3 001241-94-7 000000-00-0	52 50 25
40	25.78	0.42	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Octicizer 2-(4-Cyanophenyl)-5-dimethylaminom	34112 50542 34091	004928-02-3 001241-94-7 000000-00-0	53 47 27
41	25.88	0.33	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, 9-(3-Fluorobenzyl)-9-hydroxy-3,6,1 2-Hydrazino-8-hydroxy-4-phenylquin	34112 50407 34111	004928-02-3 000000-00-0 000000-00-0	72 22 22
42	25.97	1.94	C:\DATABASE\NBS75K.L 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Morphinan, 7,8-didehydro-4,5-epoxy Benzoic acid, 2,4-bis[(trimethylsi	56714 55885 51292	038147-00-1 055449-66-6 010586-16-0	40 38 35

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
43	26.10	0.43	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, Heptane, 1-ethoxy-	50542 34112 8517	001241-94-7 004928-02-3 001969-43-3	47 47 18
44	26.22	0.41	C:\DATABASE\NBS75K.L 3-(2-Hydroxy-6-methylphenyl)-2-met 7-Amino-2,3-dihydro-5-phenyl-1H-1, Benzeneacetaldehyde, 3,4-dimethoxy	36963 34112 34066	000000-00-0 004928-02-3 000000-00-0	38 38 38
45	26.31	0.54	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-(4-Cyanophenyl)-5-dimethylaminom	50542 34112 34091	001241-94-7 004928-02-3 000000-00-0	50 50 49
46	26.78	0.14	C:\DATABASE\NBS75K.L Octicizer 2-(4-Cyanophenyl)-5-dimethylaminom 7-Amino-2,3-dihydro-5-phenyl-1H-1,	50542 34091 34112	001241-94-7 000000-00-0 004928-02-3	70 49 45
47	26.83	0.16	C:\DATABASE\NBS75K.L Octicizer 2-(4-Cyanophenyl)-5-dimethylaminom 7-Amino-2,3-dihydro-5-phenyl-1H-1,	50542 34091 34112	001241-94-7 000000-00-0 004928-02-3	78 58 50
48	27.26	0.17	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, Benzenamine, ar,ar-dibromo-	50542 34112 33621	001241-94-7 004928-02-3 050307-05-6	60 59 43
49	27.59	1.08	C:\DATABASE\NBS75K.L Phosphoric acid, triphenyl ester Phosphoric acid, triphenyl ester Phosphoric acid, triphenyl ester	73338 46311 73337	000115-86-6 000115-86-6 000115-86-6	99 95 93
50	27.70	1.18	C:\DATABASE\NBS75K.L 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( Benzoic acid, 2,4-bis[(trimethylsi Benzeneethanamine, N-[(pentafluoro	56714 73975 74706	038147-00-1 010586-16-0 055429-13-5	38 15 10
51	28.12	43.60	C:\DATABASE\NBS75K.L Octicizer Benz[b]acridine, 1,2,3,4,7,8,9,10- Benz[c]acridine, 1,2,3,4,8,9,10,11	50542 34157 34155	001241-94-7 055044-74-1 055044-73-0	38 23 23
52	28.20	4.41	C:\DATABASE\NBS75K.L No matches found			
53	28.52	4.77	C:\DATABASE\NBS75K.L Propane, 2-ethoxy-2-methyl- 2-Butanol, 2,3-dimethyl- Butanamide, 3-methyl-	1790 63571 1621	000637-92-3 000594-60-5 000541-46-8	35 35 35
54	28.70	1.94	C:\DATABASE\NBS75K.L Propanoic acid, 2-hydroxy-2-methyl 3-Pentanol, 2-methyl- 2-Propanol, 1-(2-methoxypropoxy)-	3494 63561 9233	002110-78-3 000565-67-3 013429-07-7	38 38 38



Area Percent Report -- Sorted by Signal

Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471B.D  
 Operator : Marti  
 Acquired : 13 Apr 95 2:25 pm using AcqMethod EM31492  
 Sample Name: 0495-37b  
 Misc Info : <sup>APR 12 MAY '95</sup>Thermolag sample  
 Vial Number: 1  
 CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

Retention Time	Area	Area %	Ratio %
Total Ion Chromatogram			
0.166	-940991759	-12.730	-29.201
1.744	477714650	6.463	14.824
2.336	45118082	0.610	1.400
2.904	100951042	1.366	3.133
3.032	299430828	4.051	9.292
3.866	36385987	0.492	1.129
5.151	14676265	0.199	0.455
7.393	7713126	0.104	0.239
11.981	36416239	0.493	1.130
12.583	42069998	0.569	1.306
13.077	6720577	0.091	0.209
13.341	82553068	1.117	2.562
13.474	44243710	0.599	1.373
16.101	113445396	1.535	3.520
17.170	11435878	0.155	0.355
17.663	6291396	0.085	0.195
17.809	6406232	0.087	0.199
18.573	105712108	1.430	3.280
19.077	11009494	0.149	0.342
19.224	14806605	0.200	0.459
19.448	17014260	0.230	0.528
19.604	10994816	0.149	0.341
19.978	127290902	1.722	3.950
20.116	18277055	0.247	0.567
20.210	48738878	0.659	1.512
20.309	130898074	1.771	4.062
20.455	23328295	0.316	0.724
20.718	94695151	1.281	2.939
20.951	295043716	3.992	9.156
21.203	91827953	1.242	2.850
21.525	9855411	0.133	0.306
21.805	18501093	0.250	0.574
22.626	92106796	1.246	2.858
23.157	14491603	0.196	0.450
24.379	93814414	1.269	2.911

Area Percent Report -- Sorted by Signal

Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471B.D  
 Operator : Marti  
 Acquired : 13 Apr 95 2:25 pm using AcqMethod EM31492  
 Sample Name: 0495-37b  
 Misc Info : Thermolag sample  
 Vial Number: 1  
 CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

Retention Time	Area	Area %	Ratio %
24.714	24091389	0.326	0.748
24.992	19059905	0.258	0.591
25.300	312595369	4.229	9.700
25.648	19622723	0.265	0.609
25.783	31365149	0.424	0.973
25.880	24731593	0.335	0.767
25.968	143739453	1.945	4.460
26.107	31511870	0.426	0.978
26.219	30664316	0.415	0.952
26.305	39590163	0.536	1.229
26.788	10305991	0.139	0.320
26.827	12057232	0.163	0.374
27.263	12793678	0.173	0.397
27.590	79518975	1.076	2.468
27.696	86866396	1.175	2.696
28.114	3222515027	43.596	100.000
28.199	325968746	4.410	10.115
28.523	352471792	4.768	10.938
28.708	143447592	1.941	4.451
28.830	79747261	1.079	2.475
29.825	65838100	0.891	2.043
30.373	12169538	0.165	0.378
30.767	103484203	1.400	3.211
31.743	194030722	2.625	6.021
32.383	115536827	1.563	3.585
32.577	11814532	0.160	0.367
32.843	58374995	0.790	1.811
33.074	67218295	0.909	2.086
33.309	78624229	1.064	2.440
33.448	73011166	0.988	2.266

0A95-37b  
3 12 MAY 1995 WLF

Peak#	Ret Time	Type	Width	Area	Start Time	End Time
1.	2.357	BB	0.116	6332151	2.166	2.702
2	2.904	BV	0.102	78729031	2.781	3.466

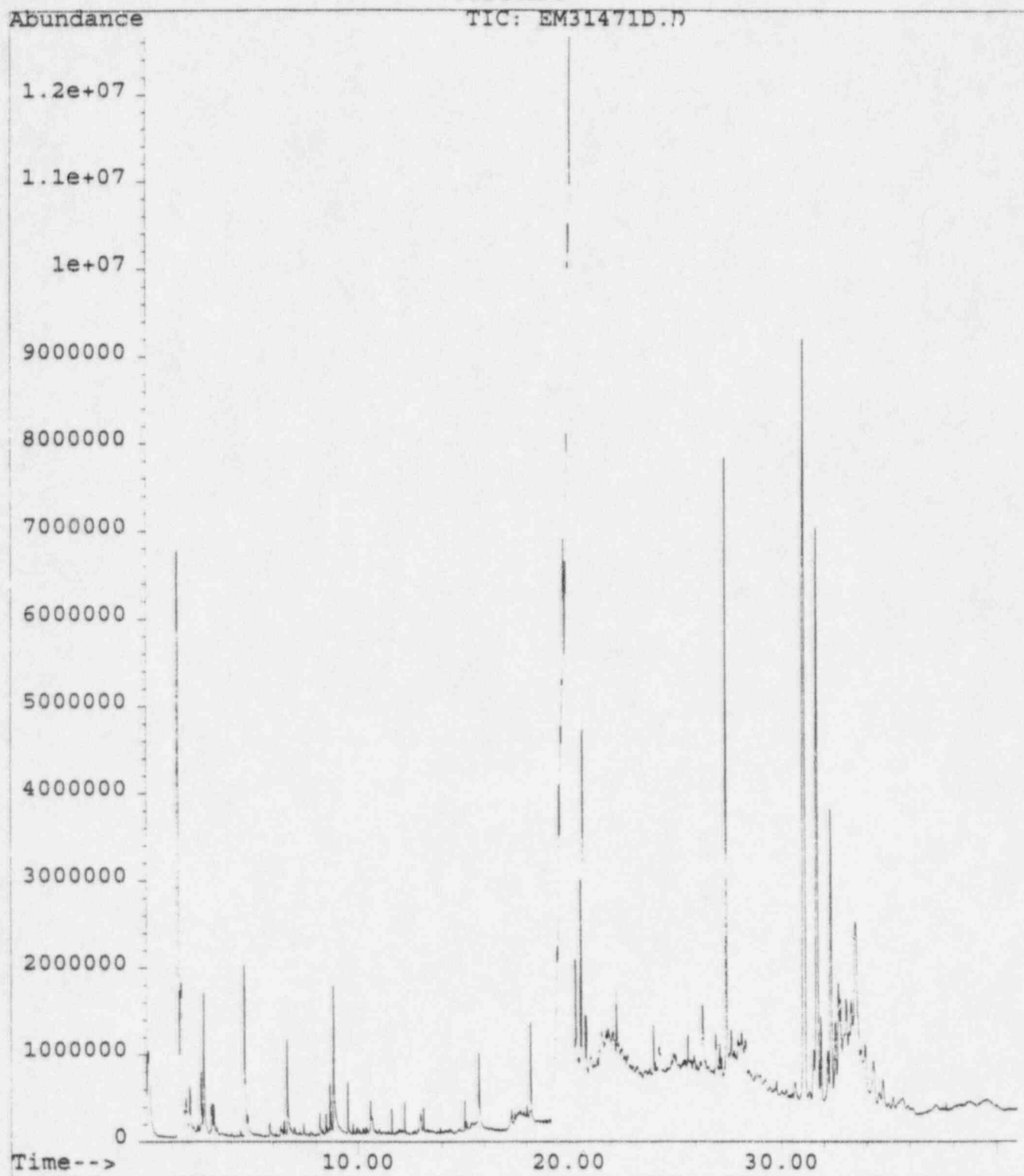
Ion 69.00 (68.70 to 69.70): EM31471B.D

0495-37b  
3 WVF 1 MAY 95

Peak#	Ret Time	Type	Width	Area	Start Time	End Time
1	3.033	PV	0.097	57150189	2.851	3.436
2	3.866	BV	0.077	10004129	3.594	4.150

File : C:\HPCHEM\1\DATA\EM31471D.D  
Operator : Marti  
Acquired : 8 May 95 1:02 pm using AcqMethod EM31492  
Instrument : 5972 - 33  
Sample Name: thermolag 0395-37c  
Misc Info : (2nd run)  
Vial Number: 1

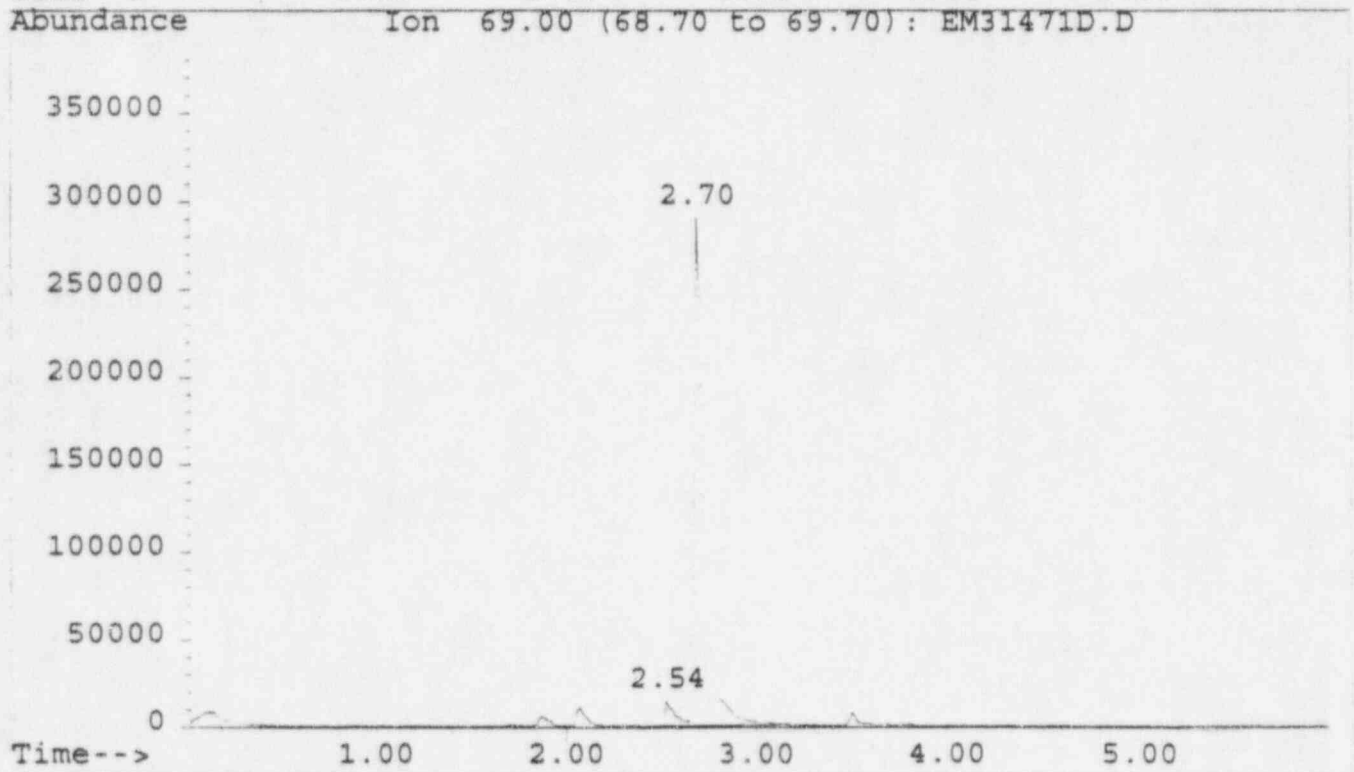
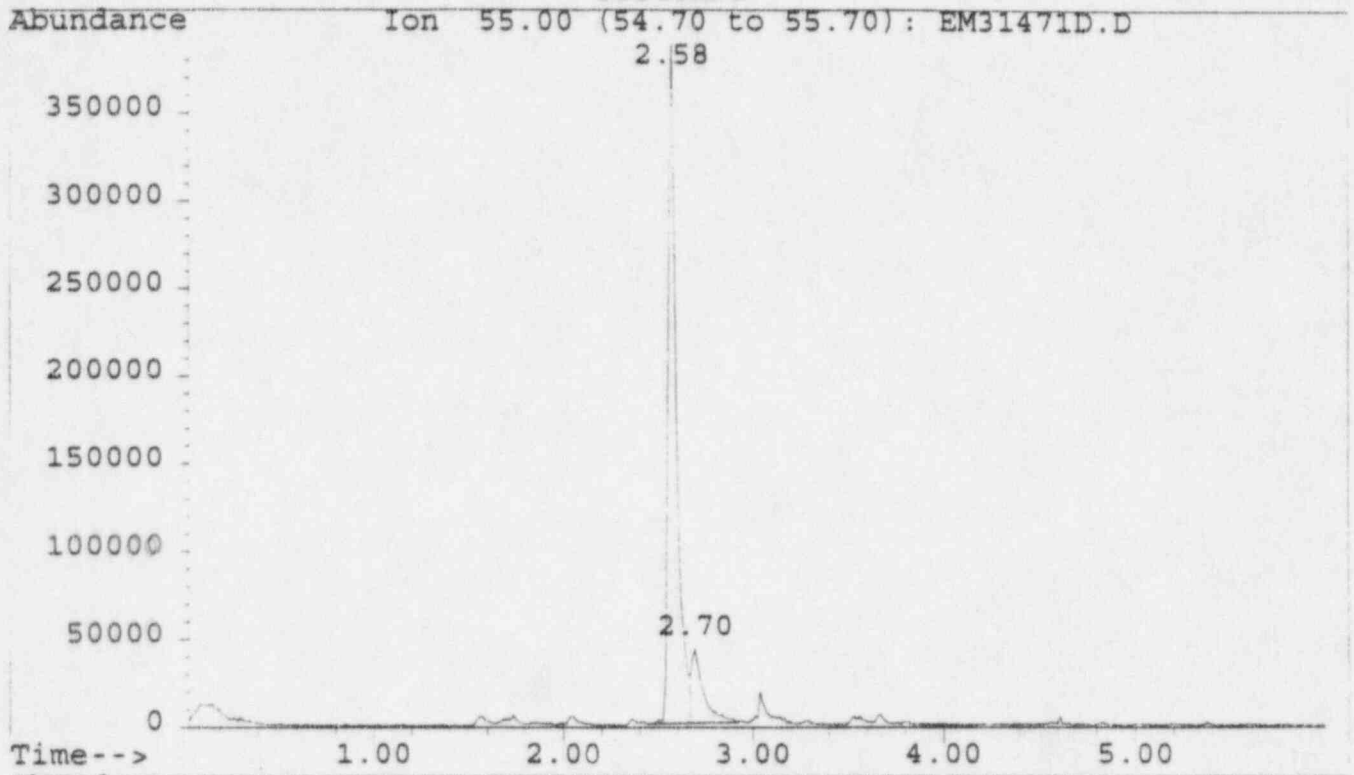
FIGURE 5





File : C:\HPCHEM\1\DATA\EM31471D.D  
Operator : Marti  
Acquired : 8 May 95 1:02 pm using AcqMethod EM31492  
Instrument : 5972 - 33  
Sample Name: thermolag 0395-37c  
Misc Info : (2nd run)  
Vial Number: 1

FIGURE 6



Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471D.D  
 Operator : Marti  
 Acquired : 8 May 95 1:02 pm using AcqMethod EM31492  
 Sample Name: thermolag 0395-37c  
 Misc Info : (2nd run)  
 Vial Number: 1

Search Libraries: C:\DATABASE\NBS75K.L Minimum Quality: 0

Unknown Spectrum: Apex  
 Integration Params: AutoIntegrate

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
1	0.14	1.08	C:\DATABASE\NBS75K.L Hydrazine, (1-methylpropyl)- 2-Propanol, 1-(2-methoxy-1-methyle 4-Ethoxyamphetamine	832 9238 17479	030924-14-2 020324-32-7 000000-00-0	30 27 22
2	1.50	4.40	C:\DATABASE\NBS75K.L No matches found			
3	1.65	1.45	C:\DATABASE\NBS75K.L No matches found			
4	1.87	0.31	C:\DATABASE\NBS75K.L 2-Propenal, 2-methyl- Furan, 2,3-dihydro- Butanal	62430 62438 62486	000078-85-3 001191-99-7 000123-72-8	50 46 43
5	2.05	0.33	C:\DATABASE\NBS75K.L Furan, 2-methyl- Furan, 2-methyl- Furan, 2-methyl-	62666 463 62665	000534-22-5 000534-22-5 000534-22-5	91 76 64
6	2.57	0.28	C:\DATABASE\NBS75K.L 2-Propenoic acid, ethyl ester 2-Propenoic acid, ethyl ester 2-Propenoic acid, ethyl ester	63318 63319 63317	000140-88-5 000140-88-5 000140-88-5	87 78 72
7	2.70	0.70	C:\DATABASE\NBS75K.L 2-Propenoic acid, 2-methyl-, methy 2-Propenoic acid, 2-methyl-, 3-hyd 2-Butenoic acid, methyl ester, (E)	1484 8333 1458	000080-62-6 002761-09-3 000623-43-8	58 50 38
8	3.06	0.15	C:\DATABASE\NBS75K.L Pyridine Pyridine Pyridine	62632 62633 62631	000110-86-1 000110-86-1 000110-86-1	95 76 76
9	3.16	0.12	C:\DATABASE\NBS75K.L 1-Penten-3-yne, 2-methyl- 1H-Pyrrole, 1-methyl- 1H-Pyrrole, 2-methyl-	62647 62656 444	000926-55-6 000096-54-8 000636-41-9	50 46 43

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
10	4.64	0.66	C:\DATABASE\NBS75K.L			
			Pyridine, 3-methyl-	63044	000108-99-6	97
			Pyridine, 3-methyl-	981	000108-99-6	94
			Pyridine, 3-methyl-	63046	000108-99-6	91
11	4.78	0.16	C:\DATABASE\NBS75K.L			
			Benzene, 1,2-dimethyl-	63706	000095-47-6	38
			Benzene, 1,2-dimethyl-	63705	000095-47-6	38
			p-Xylene	2032	000106-42-3	38
12	5.84	0.07	C:\DATABASE\NBS75K.L			
			Pyridine, 2,5-dimethyl-	2046	000589-93-5	93
			Pyridine, 2,5-dimethyl-	63719	000589-93-5	90
			Benzenamine, 3-methyl-	63727	000108-44-1	68
13	6.53	0.06	C:\DATABASE\NBS75K.L			
			Pyridine, 3-ethenyl-	1970	001121-55-7	87
			Pyridine, 2-ethenyl-	1972	000100-69-6	74
			Pyridine, 4-ethenyl-	1971	000100-43-6	64
14	6.69	0.37	C:\DATABASE\NBS75K.L			
			Pyridine, 3,5-dimethyl-	2047	000591-22-0	95
			Pyridine, 3,5-dimethyl-	63720	000591-22-0	91
			Pyridine, 3,4-dimethyl-	2050	000583-58-4	91
15	7.44	0.06	C:\DATABASE\NBS75K.L			
			Pyridine, 2-ethyl-6-methyl-	3834	001122-69-6	43
			6H-Purine, 5,7-dihydro-6-(1-methyl	17628	056247-56-4	35
			1H-Pyrrole, 2,3,5-trimethyl-	2225	002199-41-9	27
16	8.20	0.08	C:\DATABASE\NBS75K.L			
			Pyridine, 2,3,5-trimethyl-	3839	000695-98-7	91
			Pyridine, 2,3,6-trimethyl-	3822	001462-84-6	81
			Pyridine, 2,3,6-trimethyl-	64607	001462-84-6	81
17	8.49	0.09	C:\DATABASE\NBS75K.L			
			Pyridine, 3-ethyl-5-methyl-	3821	003999-78-8	94
			Pyridine, 5-ethyl-2-methyl-	64621	000104-90-5	90
			Benzenamine, 2,6-dimethyl-	64611	000087-62-7	81
18	8.69	0.21	C:\DATABASE\NBS75K.L			
			4-Aminostyrene	3652	001520-21-4	83
			Pyridine, 5-ethenyl-2-methyl-	3656	000140-76-1	76
			Benzoic acid, 2,5-dimethyl-, (2,4-	37433	055000-48-1	59
19	8.86	1.01	C:\DATABASE\NBS75K.L			
			1H-Pyrrole-2-carboxaldehyde, 1-met	63814	001192-58-1	50
			2,6-Pyridinediamine	2190	000141-86-6	47
			3,4-Pyridinediamine	2193	000054-96-6	45
20	9.54	0.16	C:\DATABASE\NBS75K.L			
			1H-Pyrrole, 2,3,4,5-tetramethyl-	4056	001003-90-3	94
			1H-Pyrrole, 3-ethyl-2,4,5-trimethyl-	65853	000520-69-4	37
			Pyrazine, 5-butyl-2,3-dimethyl-	67853	015834-78-3	23

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
21	10.62	0.26	C:\DATABASE\NBS75K.L			
			Ethanone, 1-(1-methyl-1H-pyrrol-2-	64728	000932-16-1	78
			1H-Pyrrole, 3-ethyl-2,4-dimethyl-	64741	000517-22-6	56
			1H-Pyrrole, 2-ethyl-3,5-dimethyl-	4057	032990-59-3	56
22	11.62	0.07	C:\DATABASE\NBS75K.L			
			Pentanedioic acid, 2-methyl-, mono	12301	072088-36-9	32
			Butanedioic acid, 2,2-dimethyl-, d	16117	049827-44-3	27
			Dimethyl ethylbutane-1,4-dioate	16107	000000-00-0	27
23	12.23	0.21	C:\DATABASE\NBS75K.L			
			Pentanedioic acid, diethyl ester	19778	000818-38-2	87
			Pentanedioic acid, diethyl ester	69155	000818-38-2	72
			Pentanedioic acid, diethyl ester	69156	000818-38-2	72
24	12.98	0.15	C:\DATABASE\NBS75K.L			
			Cyclohexasiloxane, dodecamethyl-	56711	000540-97-6	90
			Acetic acid, [bis[(trimethylsilyl)	49839	053044-27-2	53
			1,3,5,7,9-Pentaethylcyclopentasiloxane	51267	017995-44-7	32
25	13.13	0.07	C:\DATABASE\NBS75K.L			
			2H-Quinolizin-3-ol, octahydro-, tr	11200	015769-36-5	38
			2-(Diethylamino)-3-methylcycloprop	11187	091295-98-6	38
			Diethyl isopropylidene malonate	69708	006802-75-1	35
26	15.08	0.09	C:\DATABASE\NBS75K.L			
			Propanamide	62531	000079-05-0	46
			Methanesulfonamide, N,N-dimethyl-	4001	000918-05-8	38
			dl-3-Aminobutyric acid	1815	002835-82-7	35
27	15.76	0.50	C:\DATABASE\NBS75K.L			
			3-Isopropoxy-1,1,1,7,7,7-hexamethy	60503	071579-69-6	37
			1-[2,4-Bis(trimethylsilyloxy)pheny	58102	000000-00-0	27
			Tetrasiloxane, 3,5-diethoxy-1,1,1,	59390	072439-78-2	22
28	17.31	0.05	C:\DATABASE\NBS75K.L			
			Julolidine	16010	000479-59-4	32
			1,2,3,7-Tetramethylindole	15993	000000-00-0	25
			4-(2-Hydroxyphenyl)pyrimidine	15725	068535-55-7	25
29	17.73	0.36	C:\DATABASE\NBS75K.L			
			Cyclopentanone, dimethylhydrazone	4509	014090-60-9	27
			1,3,5-Benzenetriol	64864	000108-73-6	22
			Thiazole, 2,4-dimethyl-5-propyl-	11176	041981-70-8	22
30	17.90	0.12	C:\DATABASE\NBS75K.L			
			.delta.2-Tetrazaboroline, 5-ethyl-	4392	020534-01-4	38
			1-Ethylamino-1-butylcyclohexane	18625	000000-00-0	38
			3H-Pyrazol-3-one, 1,2-dihydro-1,2,	4441	003201-26-1	22
31	18.05	0.12	C:\DATABASE\NBS75K.L			
			N,N-Diethyltrifluoroacetamide	14830	000360-92-9	38
			Benzene, 1-chloro-4-(2,2-dichloro-	27960	054965-40-1	25
			1-Nonanone, 1-(2-thienyl)-	28638	059782-24-0	22

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
32	18.24	0.40	C:\DATABASE\NBS75K.L Benzeneethanamine, N-[(pentafluoro Phenethylamine, N-methyl-.beta.,3, Benzeneacetic acid, .alpha.,3,4-tr	60339 74238 57997	055429-13-5 010538-85-9 037148-65-5	43 43 38
33	19.89	13.30	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-(4-Cyanophenyl)-5-dimethylaminom	50542 34112 34091	001241-94-7 004928-02-3 000000-00-0	93 59 45
34	20.23	22.91	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-(4-Cyanophenyl)-5-dimethylaminom	50542 34112 34091	001241-94-7 004928-02-3 000000-00-0	90 53 42
35	20.39	1.34	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Octicizer Benzenamine, ar,ar-dibromo-	34112 50542 33621	004928-02-3 001241-94-7 050307-05-6	35 20 16
36	20.74	4.62	C:\DATABASE\NBS75K.L No matches found			
37	21.07	0.90	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Octicizer 2-(4-Cyanophenyl)-5-dimethylaminom	34112 50542 34091	004928-02-3 001241-94-7 000000-00-0	49 38 38
38	21.59	2.47	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-Propanol, 1-[2-(2-methoxy-1-meth	50542 34112 24251	001241-94-7 004928-02-3 020324-33-8	66 47 27
39	21.92	1.84	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Butanamide, 3-methyl- Butane, 2-methoxy-	34112 1621 854	004928-02-3 000541-46-8 006795-87-5	35 30 30
40	22.12	1.26	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Methyl 16-methoxyheptadecanoate Octicizer	34112 44818 50542	004928-02-3 000000-00-0 001241-94-7	27 27 25
41	22.30	1.13	C:\DATABASE\NBS75K.L Trisiloxane, 1,1,1,5,5,5-hexamethy 1,1,1,5,7,7,7-Heptamethyl-3,3-bis( 7-Amino-2,3-dihydro-5-phenyl-1H-1,	52459 56714 34112	003555-47-3 038147-00-1 004928-02-3	14 10 10
42	24.05	0.61	C:\DATABASE\NBS75K.L Octicizer 2,4(1H,3H)-Quinolinedione, 3-benzo 7-Amino-2,3-dihydro-5-phenyl-1H-1,	50542 49822 34112	001241-94-7 070611-42-6 004928-02-3	35 32 27
43	24.31	1.10	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Octicizer 2,4,6,8,9,10-Hexathiatricyclo[3.3.	34112 50542 50037	004928-02-3 001241-94-7 057274-38-1	47 25 18

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
44	24.98	1.59	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-Propanol, 1-[2-(2-methoxy-1-meth Octicizer	34112 24251 50542	004928-02-3 020324-33-8 001241-94-7	38 35 35
45	25.46	0.68	C:\DATABASE\NBS75K.L 7-Amino-2,3-dihydro-5-phenyl-1H-1, Octicizer 2-Propanol, 1-[2-(2-methoxy-1-meth	34112 50542 24251	004928-02-3 001241-94-7 020324-33-8	43 42 22
46	25.65	0.66	C:\DATABASE\NBS75K.L 3-(2-Hydroxy-6-methylphenyl)-2-met Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1,	36963 50542 34112	000000-00-0 001241-94-7 004928-02-3	47 45 43
47	25.94	0.53	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-Propanol, 1-[1-methyl-2-(2-prope	50542 34112 68438	001241-94-7 004928-02-3 055956-25-7	70 43 18
48	26.35	1.12	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 51123 73955	000563-04-2 001330-78-5 001330-78-5	99 94 93
49	26.97	0.42	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 73955 51123	000563-04-2 001330-78-5 001330-78-5	97 95 78
50	27.16	0.32	C:\DATABASE\NBS75K.L Phosphoric acid, triphenyl ester Phosphoric acid, triphenyl ester Phosphoric acid, triphenyl ester	73338 73337 46311	000115-86-6 000115-86-6 000115-86-6	99 92 83
51	27.50	2.69	C:\DATABASE\NBS75K.L Octicizer 7-Amino-2,3-dihydro-5-phenyl-1H-1, 2-(4-Cyanophenyl)-5-dimethylaminom	50542 34112 34091	001241-94-7 004928-02-3 000000-00-0	90 59 59
52	27.70	0.76	C:\DATABASE\NBS75K.L Isothiazole, 3-methyl-5-phenyl- 2H-1-Benzopyran-2-one, 7-amino-4-m Oxycarboxin	68481 16440 71900	001732-45-2 026093-31-2 005259-88-1	37 28 25
53	28.20	0.92	C:\DATABASE\NBS75K.L Propanoic acid, 2-hydroxy-2-methyl 2-Butanol, 3-methoxy- Butanamide, 3-methyl-	5790 1908 1621	000080-55-7 053778-72-6 000541-46-8	47 43 38
54	28.37	0.39	C:\DATABASE\NBS75K.L Butane, 2-methoxy- Propanoic acid, 2-hydroxy-2-methyl Pentanamide, 4-methyl-	854 3494 3136	006795-87-5 002110-78-3 001119-29-5	43 38 38

Pk#	RT	Area%	Library/ID	Ref#	CAS#	Qual
.55	29.79	0.09	C:\DATABASE\NBS75K.L Pregna-4,6-diene-3,20-dione, 2.alpha. 10H-Phenoxaphosphine, 8-fluoro-10- 4H-1-Benzopyran-2-carboxylic acid,	56634 39085 39266	005234-56-0 037041-13-7 030095-84-2	27 25 16
56	30.65	0.14	C:\DATABASE\NBS75K.L .alpha.-D-Xylo-Hex-5-enofuranose, Hexane, 1-ethoxy- 8-Hydroxycarvotanacetone	19338 65280 14600	007284-07-3 005756-43-4 007712-46-1	10 10 9
57	31.14	5.94	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 73955 51123	000563-04-2 001330-78-5 001330-78-5	99 93 91
58	31.54	0.37	C:\DATABASE\NBS75K.L Sarpagan-16-carboxylic acid, 3,17- Benzeneethanamine Bikaverin	52356 64604 52326	053632-75-0 000064-04-0 033390-21-5	22 14 11
59	31.73	4.03	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 51123 73955	000563-04-2 001330-78-5 001330-78-5	99 94 93
60	31.89	0.45	C:\DATABASE\NBS75K.L Acetophenone, 2',4'-dimethoxy-3'-m 9H-Fluorene, 9,9-dimethyl- 9H-Fluoren-9-imine	21229 21436 17504	060512-80-3 004569-45-3 004440-33-9	30 22 18
61	32.21	0.37	C:\DATABASE\NBS75K.L Sarpagan-16-carboxylic acid, 3,17- Morphinan, 6,8a-etheno,-3-acetoxy- Acridine	52356 52348 68683	053632-75-0 000000-00-0 000260-94-6	22 18 15
62	32.38	1.90	C:\DATABASE\NBS75K.L Phosphoric acid, tris(3-methylphen Phosphoric acid, tris(methylphenyl Phosphoric acid, tris(methylphenyl	51124 73955 51123	000563-04-2 001330-78-5 001330-78-5	98 96 93
63	32.57	0.62	C:\DATABASE\NBS75K.L Ethanol, 2-[4-(1,1-dimethylethyl)p Acetophenone, 2',4'-dimethoxy-3'-m 2',4'-Dimethoxy-3'-methylpropiope	21346 21229 24862	000713-46-2 060512-80-3 077942-13-3	22 22 14
64	32.72	1.61	C:\DATABASE\NBS75K.L 2,2,6,6-Tetrachloro-1,5-dioxa-2,6- Morphinan, 6,8a-etheno,-3-acetoxy- Sarpagan-16-carboxylic acid, 3,17-	52155 52348 52356	063503-14-0 000000-00-0 053632-75-0	23 22 22
65	33.08	1.62	C:\DATABASE\NBS75K.L Butanamide, 3-methyl- 2,3,3-Trimethyl-2-pentanol Ethane, 1-bromo-2-ethoxy-	1621 5510 10074	000541-46-8 023171-85-9 000592-55-2	22 22 22

Pk#	RT	Area	Library/ID	Ref#	CAS#	Qual
66	33.32	1.50	C:\DATABASE\NBS75K.L			
			2-Propanol, 1-(2-methoxypropoxy)-	9233	013429-07-7	38
			2-Propanol, 1-[1-methyl-2-(2-prope	16196	055956-25-7	38
			Butanamide, 3-methyl-	1621	000541-46-8	38
67	33.52	3.46	C:\DATABASE\NBS75K.L			
			2-Propanol, 1-(2-methoxypropoxy)-	9233	013429-07-7	14
			1-Butanol, 3-methoxy-	1922	002517-43-3	14
			Butanamide, 3-methyl-	1621	000541-46-8	14
68	33.95	0.87	C:\DATABASE\NBS75K.L			
			1-Propanol, 2-(2-methoxy-1-methyle	9239	055956-21-3	43
			2-Propanol, 1-[2-(2-methoxy-1-meth	24251	020324-33-8	37
			Methane, diethoxy-	1915	000462-95-3	27
69	34.33	0.62	C:\DATABASE\NBS75K.L			
			1H-Benzimidazole, 5-chloro-2-(1-me	21182	004886-29-7	20
			Acridine	68684	000260-94-6	11
			Acridine	68685	000260-94-6	11
70	34.76	0.28	C:\DATABASE\NBS75K.L			
			Bikaverin	52326	033390-21-5	38
			p-Quiniquiphenyl	52398	000000-00-0	18
			1,1':3',1'':3'',1''':3''',1''''-Qu	52399	016716-13-5	18
71	35.23	0.08	C:\DATABASE\NBS75K.L			
			Pentane, 2-methoxy-	1778	006795-88-6	30
			2-Propanol, 1-[1-methyl-2-(2-prope	68438	055956-25-7	27
			2,4,6,8,9,10-Hexathiatricyclo[3.3.	50037	057274-38-1	25
72	39.57	0.34	C:\DATABASE\NBS75K.L			
			2-Propanol, 1-(2-methoxy-1-methyle	9238	020324-32-7	38
			Butane, 2-methoxy-	854	006795-87-5	35
			Propanoic acid, 2-hydroxy-2-methyl	3494	002110-78-3	35



Area Percent Report -- Sorted by Signal

Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471D.D  
 Operator : Marti  
 Acquired : 8 May 95 1:02 pm using AcqMethod EM31492  
 Sample Name: thermolag 0395-37c  
 Misc Info : (2nd run)  
 Vial Number: 1  
 CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

Retention Time	Area	Area %	Ratio %
Total Ion Chromatogram			
0.140	92722369	1.082	4.723
1.495	377302691	4.403	19.218
1.657	124284126	1.450	6.330
1.871	26406527	0.308	1.345
2.050	28477981	0.332	1.451
2.578	24074246	0.281	1.226
2.698	60276495	0.703	3.070
3.058	13211834	0.154	0.673
3.154	10119389	0.118	0.515
4.638	56653564	0.661	2.886
4.777	14011580	0.164	0.714
5.844	5834804	0.068	0.297
6.528	5096275	0.059	0.260
6.683	31886156	0.372	1.624
7.443	4983175	0.058	0.254
8.197	7042570	0.082	0.359
8.489	7286336	0.085	0.371
8.695	18230036	0.213	0.929
8.861	86836671	1.013	4.423
9.542	13358226	0.156	0.680
10.624	21907773	0.256	1.116
11.630	5640899	0.066	0.287
12.235	18000000	0.210	0.918
12.987	2278725	0.149	0.649
13.128	5127453	0.074	0.322
15.077	7925161	0.092	0.404
15.760	42928444	0.501	2.187
17.310	4197339	0.049	0.214
17.725	30486288	0.356	1.553
17.893	10339636	0.121	0.527
18.048	9924107	0.116	0.505
18.239	34576125	0.403	1.761
19.885	1140080553	13.305	58.070
20.230	1963278058	22.911	100.000
20.388	114747435	1.339	5.845

Area Percent Report -- Sorted by Signal

Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471D.D  
 Operator : Marti  
 Acquired : 8 May 95 1:02 pm using AcqMethod EM31492  
 Sample Name: thermolag 0395-37c  
 Misc Info : (2nd run)  
 Vial Number: 1  
 CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

Retention Time	Area	Area %	Ratio %
20.735	396014951	4.621	20.171
21.069	76964275	0.898	3.920
21.593	211678932	2.470	10.782
21.910	157992826	1.844	8.047
22.115	107867502	1.259	5.494
22.302	96593679	1.127	4.920
24.056	51858646	0.605	2.641
24.311	94559977	1.104	4.816
24.978	136006512	1.587	6.928
25.455	58452866	0.682	2.977
25.650	56829877	0.663	2.895
25.938	45328226	0.529	2.309
26.358	95901695	1.119	4.885
26.976	35700684	0.417	1.818
27.160	27420393	0.320	1.397
27.496	230192488	2.686	11.725
27.706	65082688	0.760	3.315
28.196	79075966	0.923	4.028
28.369	33750458	0.394	1.719
29.793	7314431	0.085	0.373
30.651	12384558	0.145	0.631
31.134	508619835	5.936	25.907
31.544	31367955	0.366	1.598
31.737	345037129	4.027	17.575
31.888	38957251	0.455	1.984
32.204	31981791	0.373	1.629
32.374	162852128	1.900	8.295
32.566	52874565	0.617	2.693
32.723	138104414	1.612	7.034
33.080	138933784	1.621	7.077
33.312	128192081	1.496	6.529
33.524	296709585	3.463	15.113
33.948	74164746	0.865	3.778
34.326	52854560	0.617	2.692
34.759	24222641	0.283	1.234
35.227	6807583	0.079	0.347

Area Percent Report -- Sorted by Signal

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Information from Data File:

File : C:\HPCHEM\1\DATA\EM31471D.D  
Operator : Marti  
Acquired : 8 May 95 1:02 pm using AcqMethod EM31492  
Sample Name: thermolag 0395-37c  
Misc Info : (2nd run)  
Vial Number: 1  
CurrentMeth: C:\HPCHEM\1\METHODS\EM31492.M

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Retention Time	Area	Area %	Ratio %
39.568	29192820	0.341	1.487

---

Ion 55.00 (54.70 to 55.70): EM31471D.D  
thermolag 0395-37c

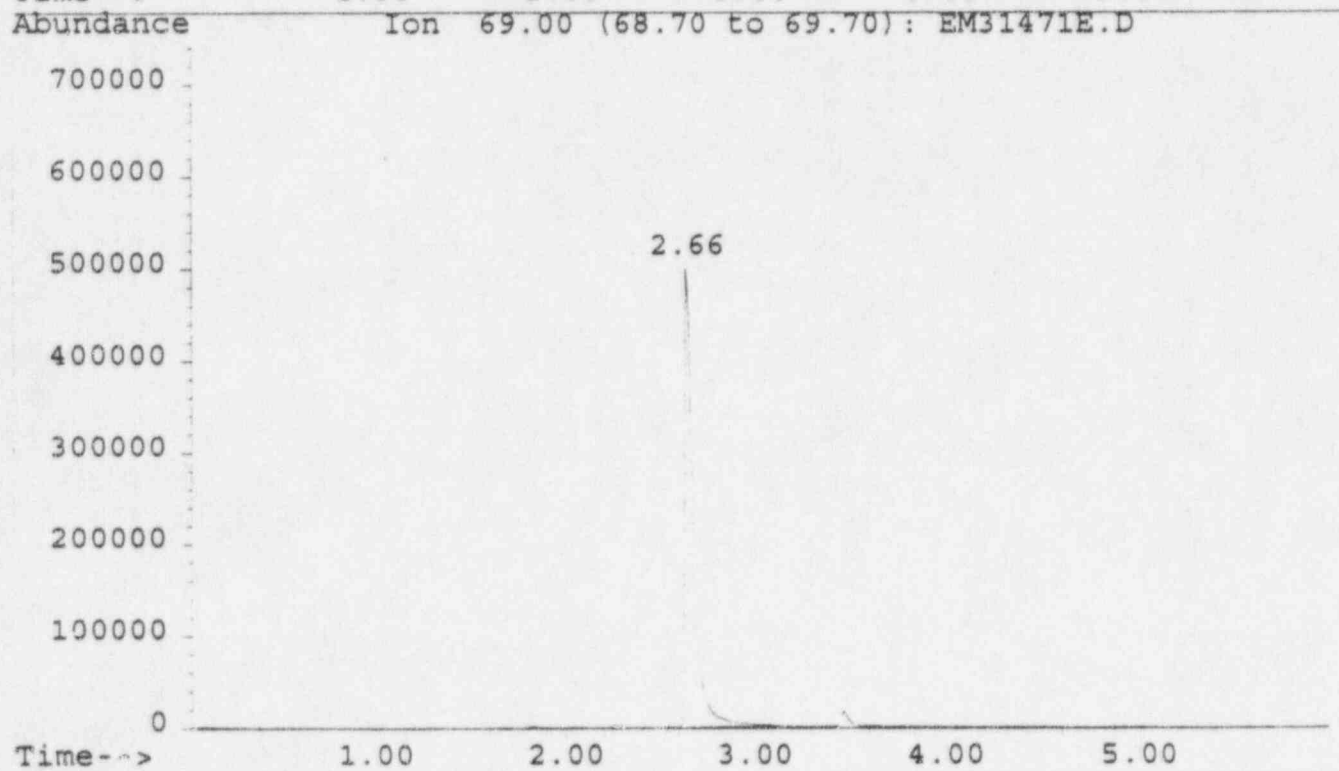
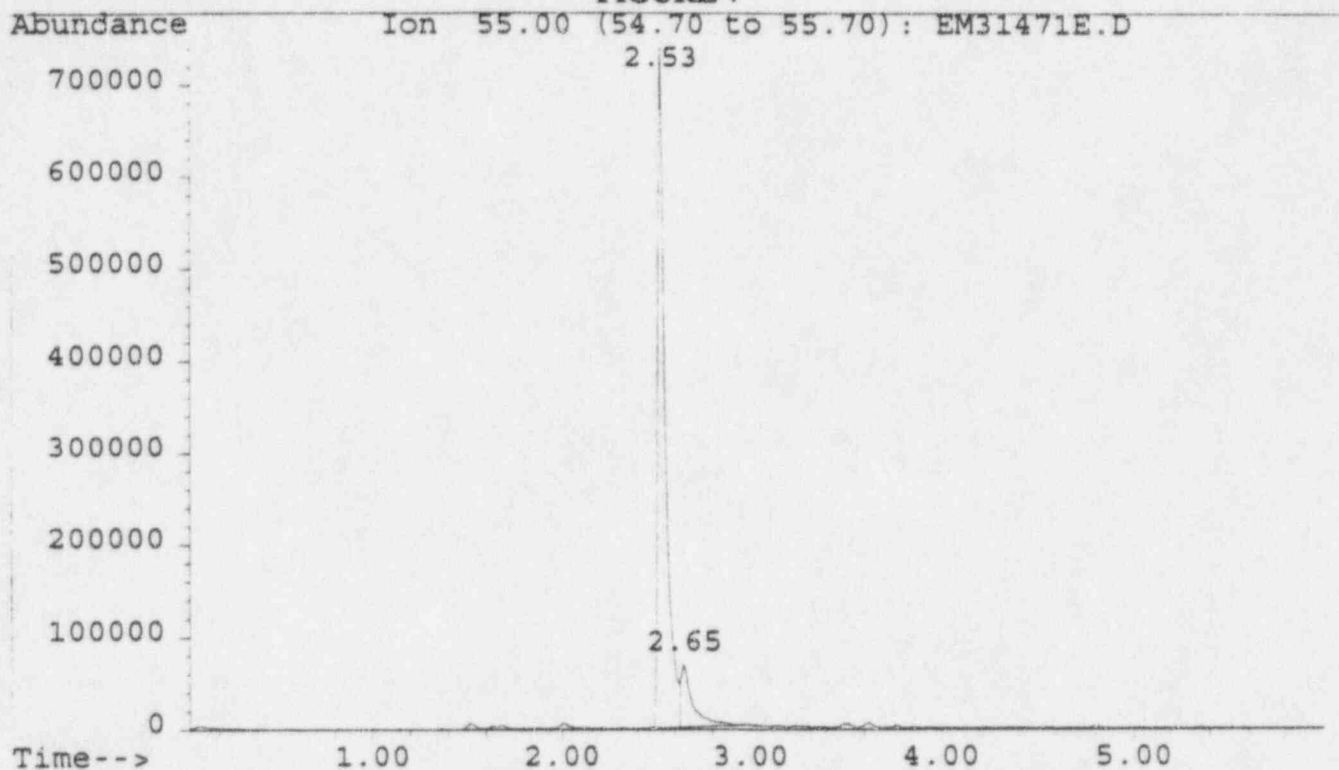
Peak#	Ret Time	Type	Width	Area	Start Time	End Time
1	2.577	BV	0.048	11671429	2.444	2.667
2	2.696	VV	0.079	1914898	2.667	2.920

Ion 69.00 (68.70 to 69.70): EM31471D.D  
thermolag 0395-37c

Peak#	Ret Time	Type	Width	Area	Start Time	End Time
1	2.536	BV	0.058	483214	2.484	2.638
2	2.700	VV	0.052	9629230	2.638	3.109

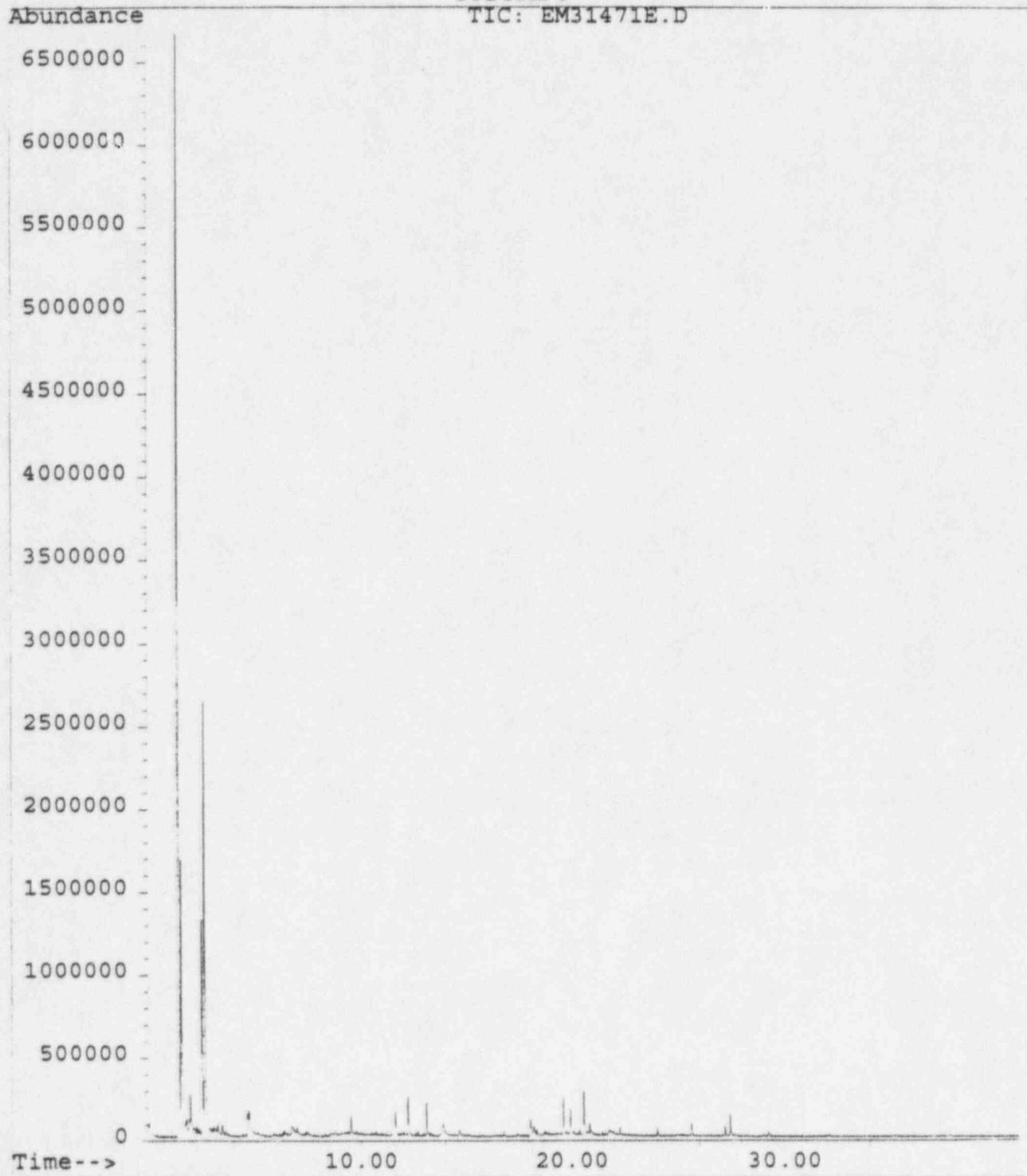
File : C:\HPCHEM\1\DATA\EM31471E.D  
Operator : Marti  
Acquired : 23 May 95 10:14 am using AcqMethod EM31492  
Instrument : 5972 - 33  
Sample Name: 0495-37b  
Misc Info : 3 WFF 30 MAR 95  
Vial Number: 1

FIGURE 7



File : C:\HPCHEM\1\DATA\EM31471E.D  
Operator : Marti  
Acquired : 23 May 95 10:14 am using AcqMethod EM31492  
Instrument : 5972 - 33  
Sample Name: Q495-37b  
Misc Info : 3 *23 May 95*  
Vial Number: 1 *30*  
*w/4*

FIGURE 8



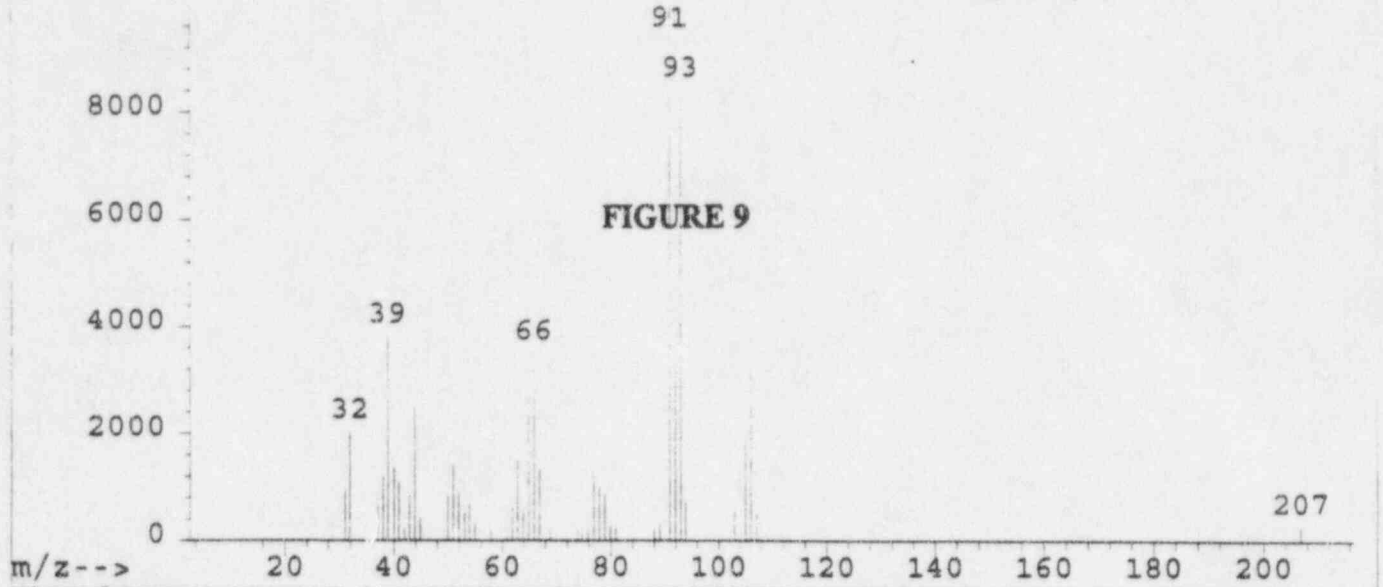
Library Searched : C:\DATABASE\NBS75K.L

Quality : 93

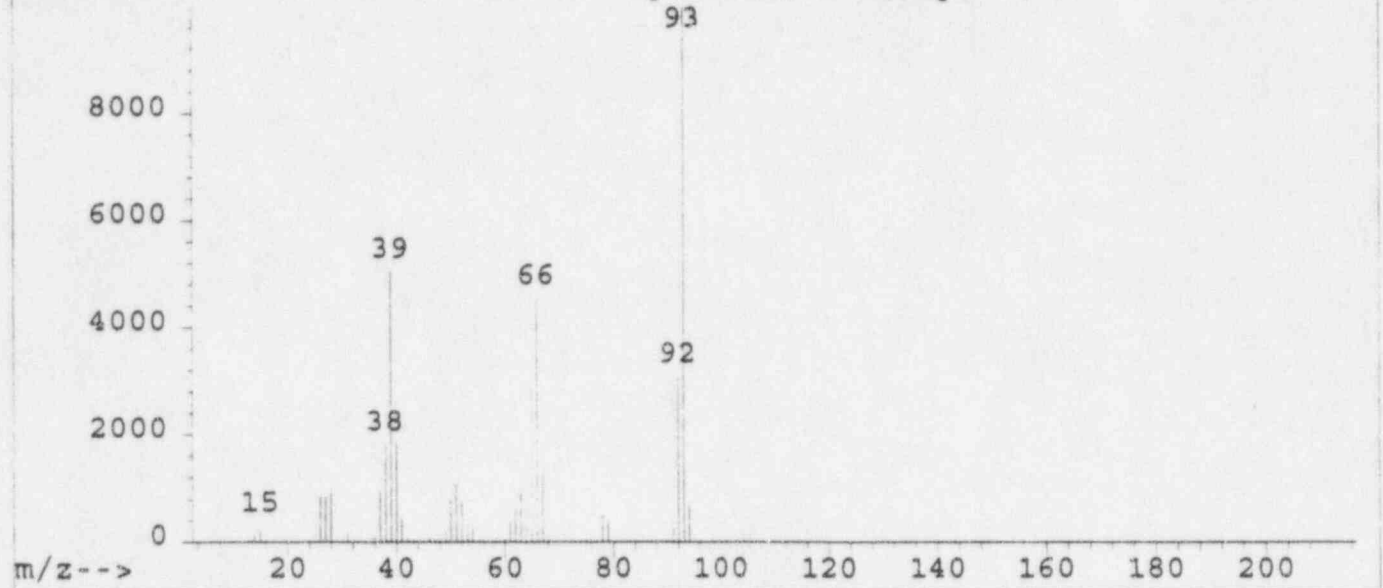
ID : Pyridine, 3-methyl-

SAMPLE 0395-37B

Abundance Scan 394 (4.733 min): EM31471E.D (\*)



Abundance #63045: Pyridine, 3-methyl- (\*)





Library Searched : C:\DATABASE\NBS75K.L

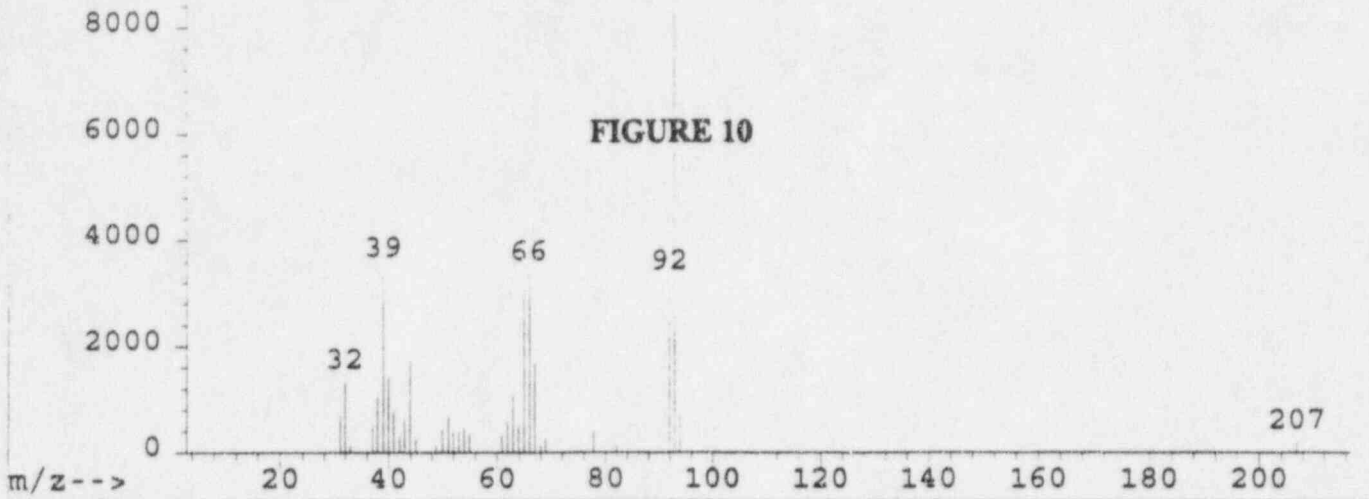
Quality : 96

ID : Pyridine, 3-methyl-

SAMPLE 0395-37B

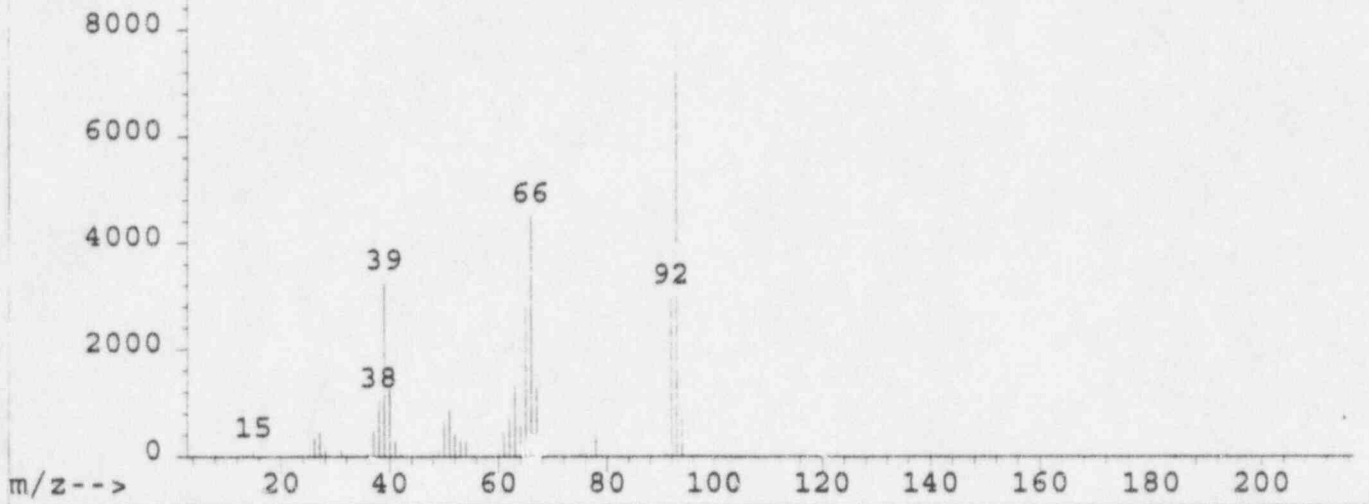
Abundance Scan 389 (4.673 min): EM31471E.D (\*)

93



Abundance #981: Pyridine, 3-methyl- (\*)

93



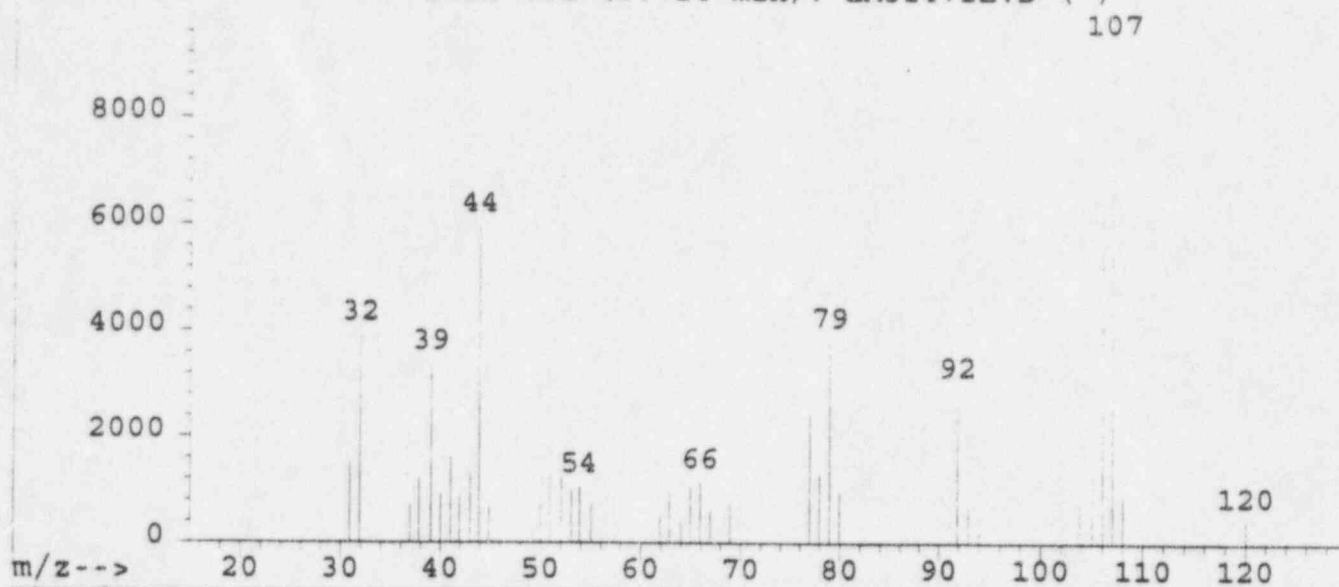
Library Searched : C:\DATABASE\NBS75K.L

Quality : 94

ID : Pyridine, 3,5-dimethyl-

SAMPLE 0395-37B

Abundance Scan 561 (6.724 min): EM31471E.D (\*)



Abundance #2047: Pyridine, 3,5-dimethyl- (\*)

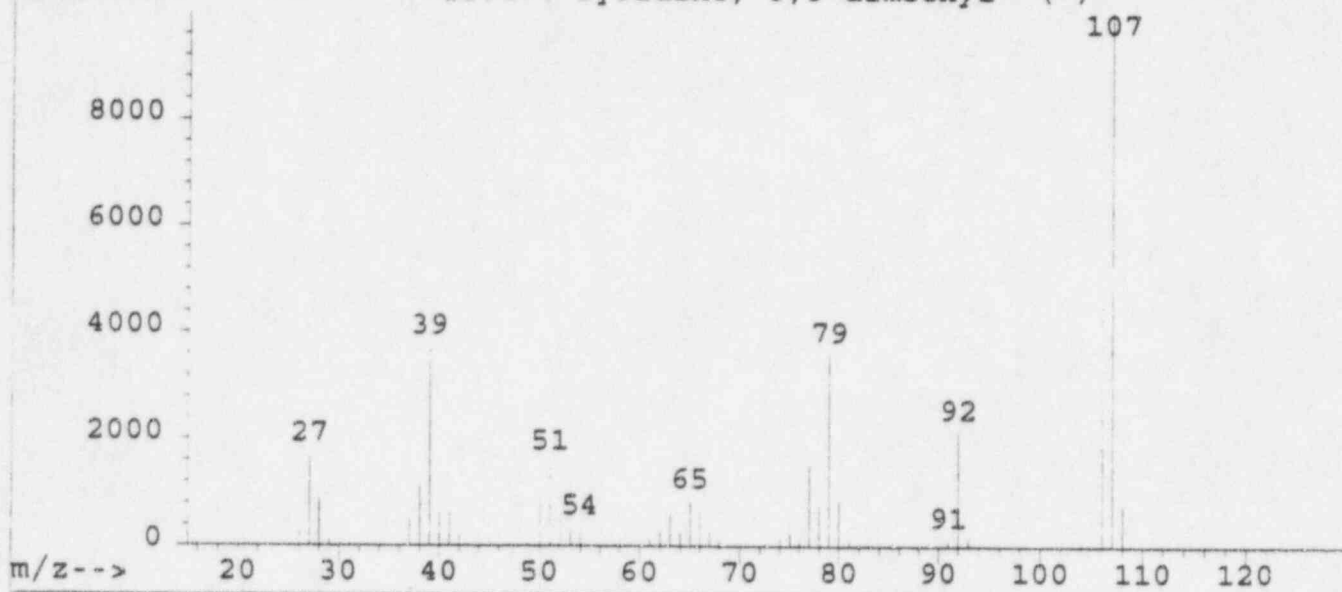


FIGURE 11

Library Searched : C:\DATABASE\NBS75K.L  
Quality : 64  
ID : ~~WIF~~ 7 Pyridine, 2,5-dimethyl-  
SAMPLE 0395-37B

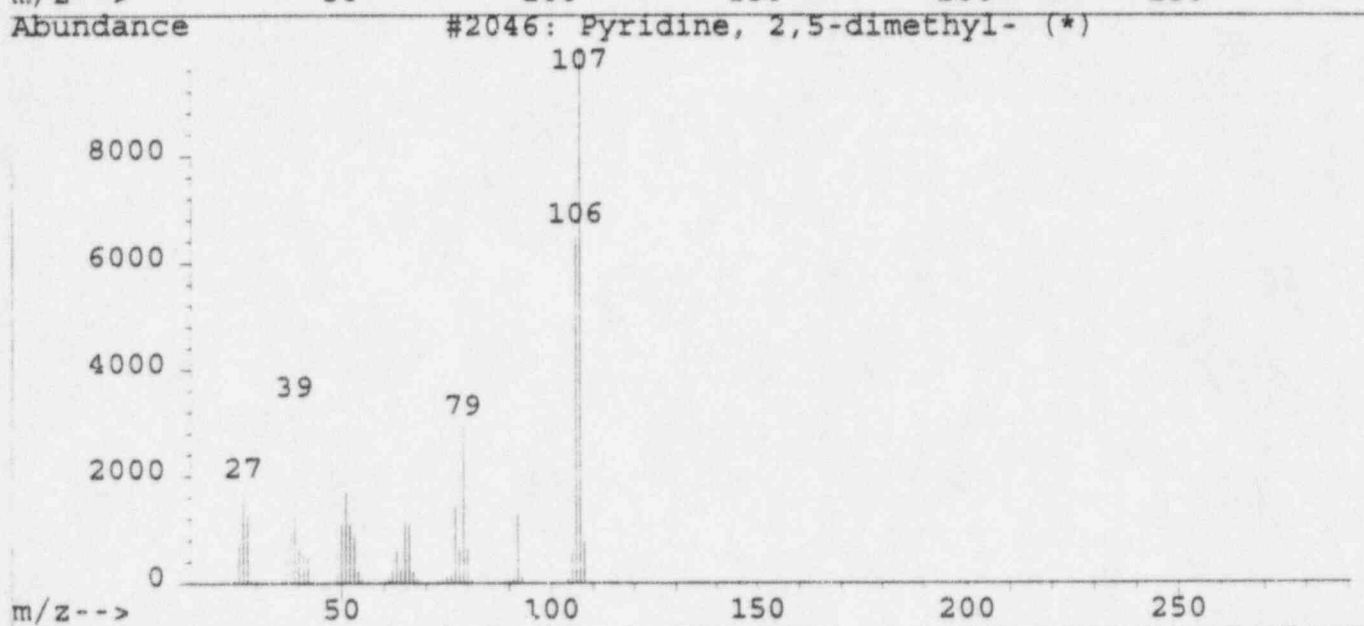
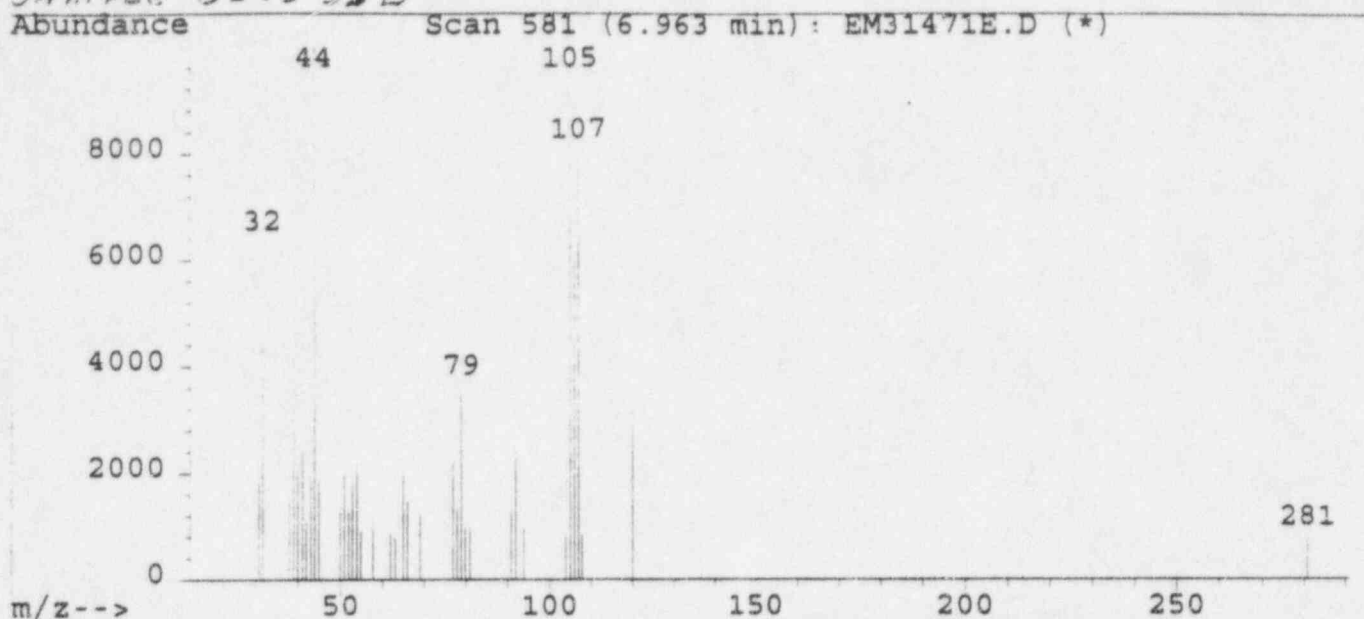


FIGURE 12

Library Searched : C:\DATABASE\NBS75K.L  
Quality : 68  
ID : Pentanedioic acid, diethyl ester

SAMPLE 0395-373

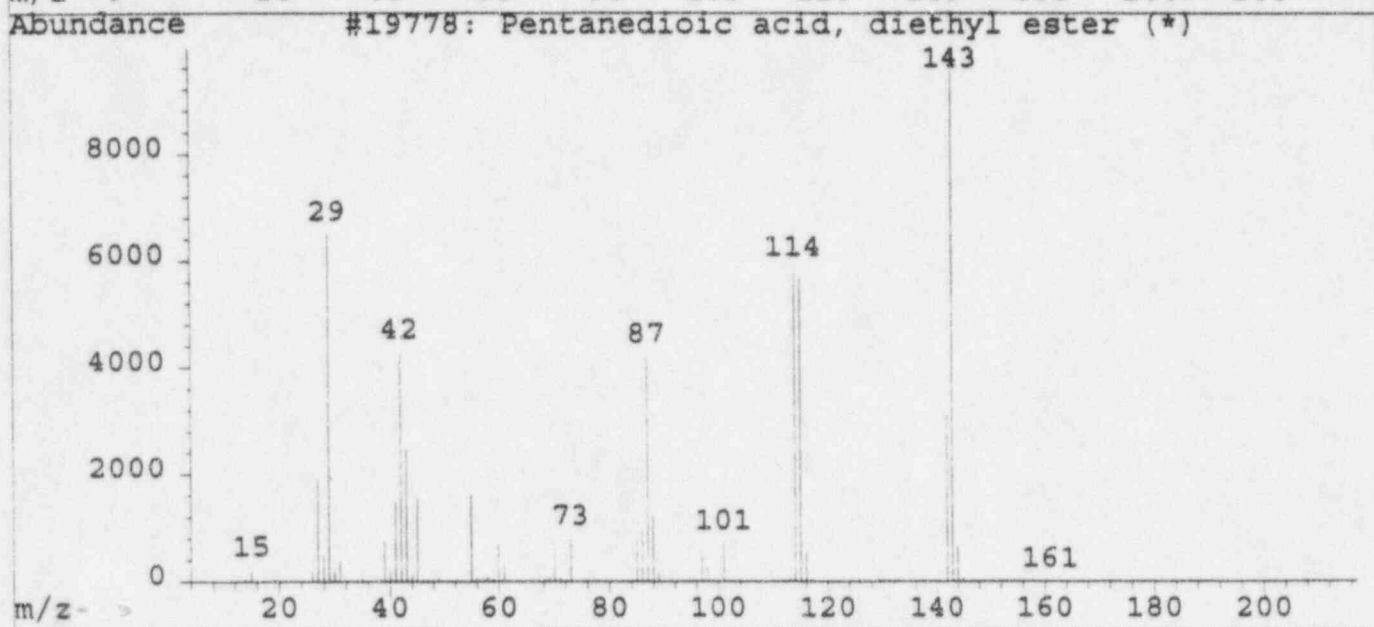
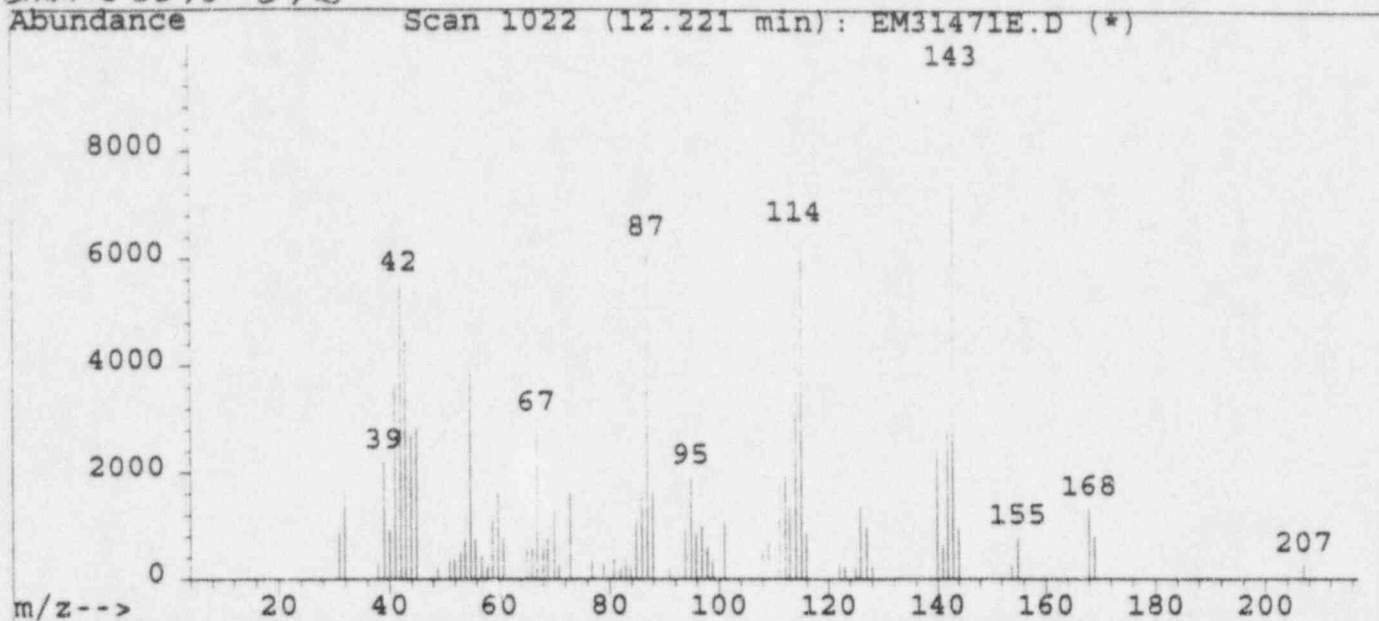


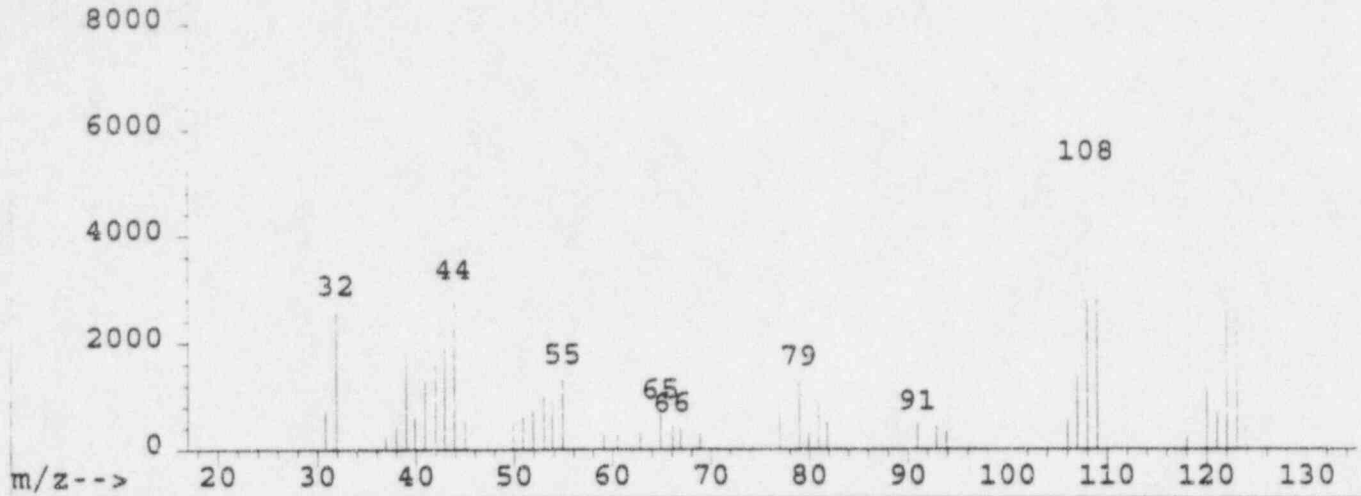
FIGURE 13

Library Searched : C:\DATABASE\NBS75K.L  
Quality : 91  
ID : 1H-Pyrrole, 2,3,4,5-tetramethyl-

SAMPLE 0395-373

Abundance Scan 794 (9.502 min): EM31471E.D (\*)

122



Abundance #4056: 1H-Pyrrole, 2,3,4,5-tetramethyl- (\*)

122

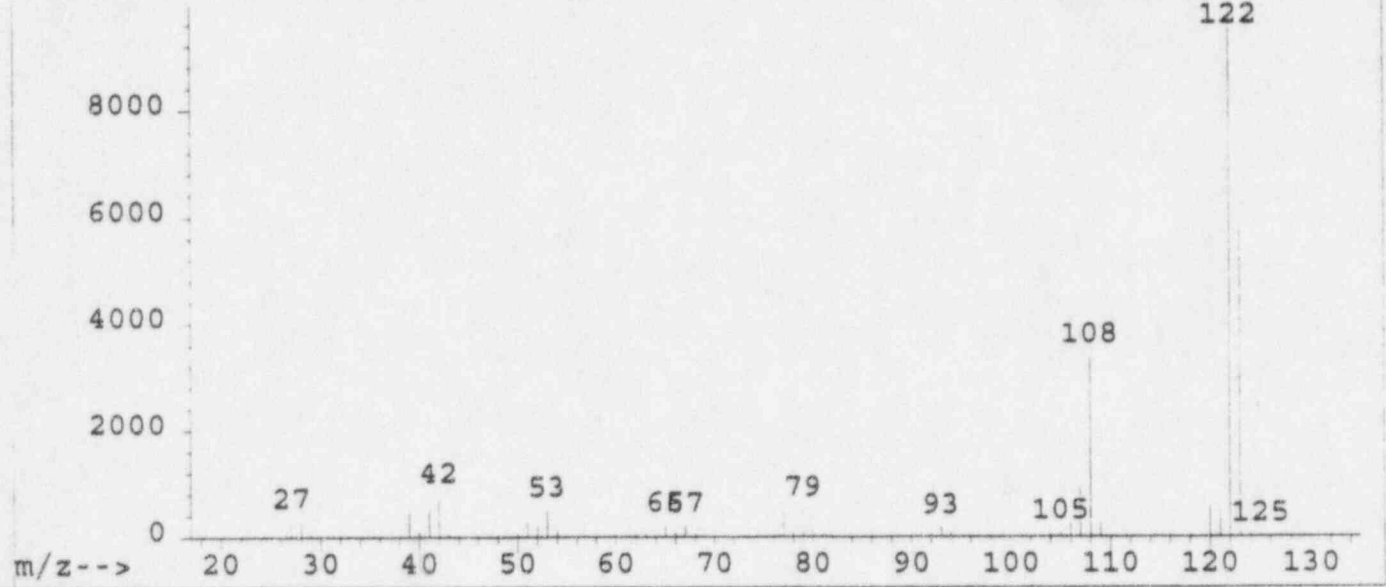


FIGURE 14

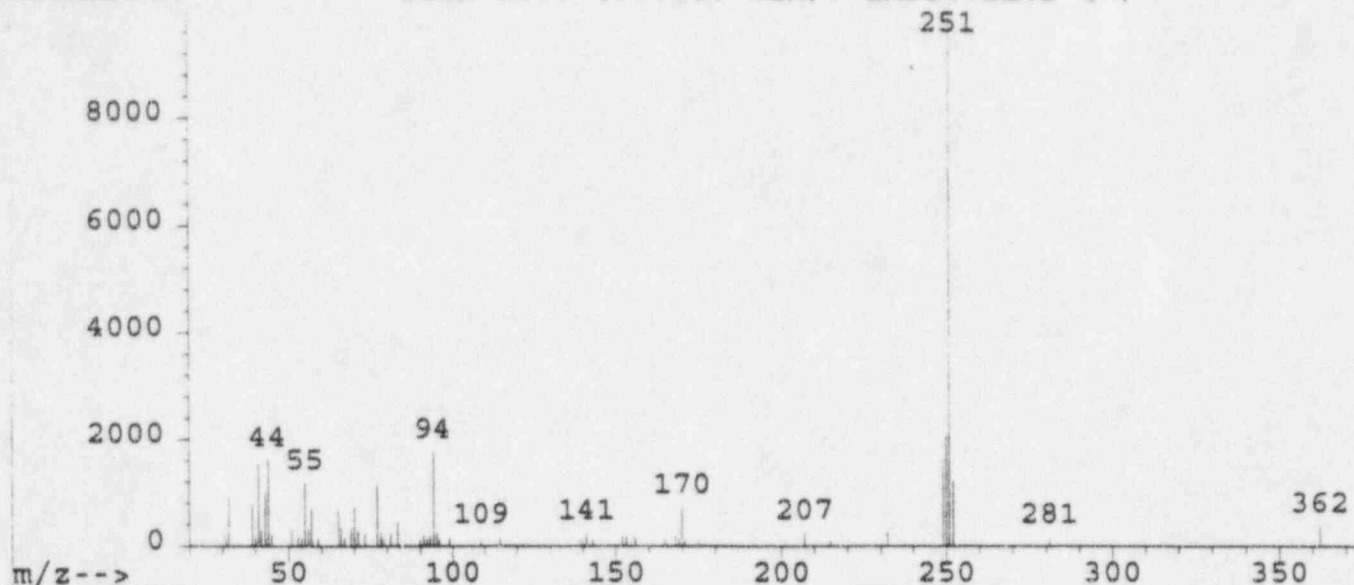
Library Searched : C:\DATABASE\NBS75K.L

Quality : 49

ID : Octicizer

SAMPLE 0395-378

Abundance Scan 2300 (27.460 min): EM31471E.D (\*)



Abundance #50542: Octicizer (\*)

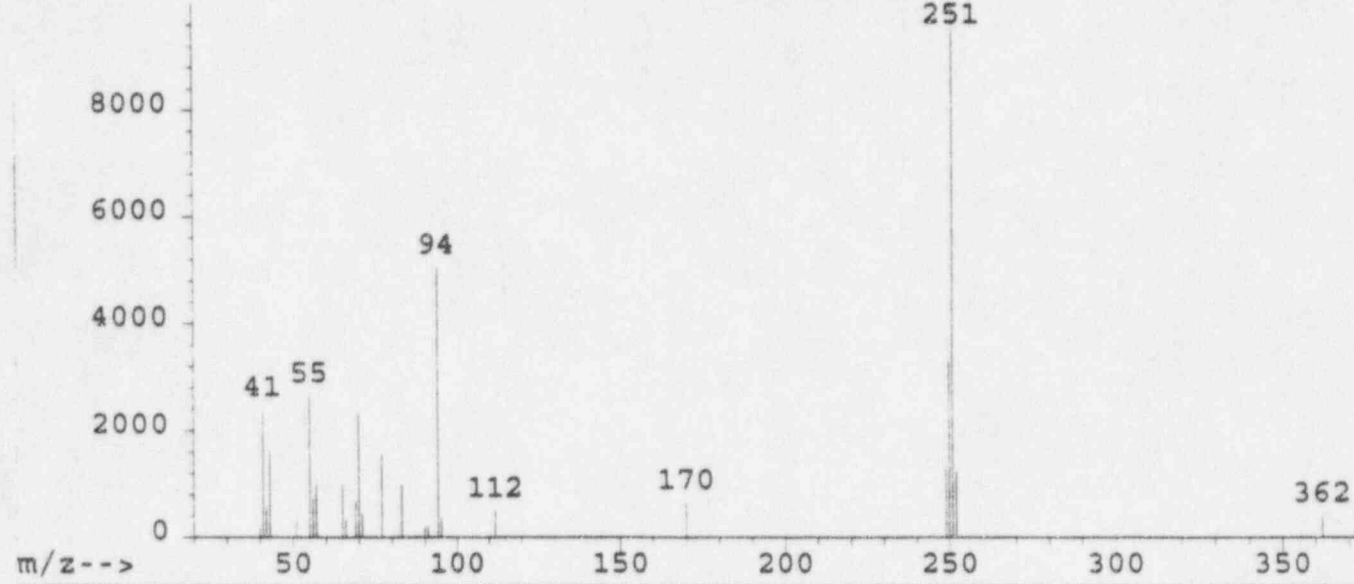


FIGURE 15

**ATTACHMENT No. 3**

**UPDATED**

**GENERIC LETTER 86-10**

**EVALUATION OF**

**WATERFORD 3**

**THERMO-LAG APPLICATIONS**