

User: Galos, Marie  
Westchester Dept. of Labs & Research  
Date: 12/11/2001 Time: 16:54:26

Sample: AD26985  
Analysis: \$GCMS GCMS Scan For Unknowns  
Units: ug  
Started: 12/11/01 16:53 Ended: 12/11/01 16:53 Analyst: MG  
Page: 1

|   | Component Name          | Viol | Result      | MDL |
|---|-------------------------|------|-------------|-----|
| 1 | Other unknown compounds |      | see comment |     |
| 2 | Surr_2,4-DDD            |      |             | 5.0 |

**Comments for Sample AD26985 - Analysis \$GCMS**

Westchester Dept. of Labs & Research

User: Galos, Marie

Date: 12-11-2001 Time: 16:54:30

A scan of the extract prepared for the 508 analysis, from 35-550 amu using a mass spectrometer, does not reveal the presence of non-target compounds.

## Quantitation Report

(QT Reviewed)

Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Acq On : 7 Dec 2001 2:24 pm  
 Sample : gc/ms unknown  
 Misc :  
 MS Integration Params: RTEINT.P  
 Quant Time: Dec 11 11:44 2001

Vial: 31  
 Operator: GALOS  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Results File: NP112701.RES

Quant Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
 Title : 208 625/TCLP calibration table  
 Last Update : Wed Nov 28 15:33:44 2001  
 Response via : Initial Calibration  
 DataAcq Meth : NP112701

| Internal Standards        | R.T.  | QIon | Response | Conc  | Units | Dev(Min) |
|---------------------------|-------|------|----------|-------|-------|----------|
| 1) 1,4-Dichlorobenzene-d4 | 11.89 | 152  | 1112743  | 20.00 | ng/uL | -0.05    |
| 19) Naphthalene-d8        | 15.50 | 136  | 4187522  | 20.00 | ng/uL | -0.05    |
| 31) Acenaphthene-d10      | 20.77 | 164  | 1913233  | 20.00 | ng/uL | -0.05    |
| 49) Phenanthrene-d10      | 25.18 | 188  | 2793469  | 20.00 | ng/uL | -0.06    |
| 59) Chrysene-d12          | 33.19 | 240  | 1961446  | 20.00 | ng/uL | -0.06    |
| 68) Perylene-d12          | 37.28 | 264  | 1398003  | 20.00 | ng/uL | -0.06    |

## System Monitoring Compounds

|                           |               |     |            |        |       |       |
|---------------------------|---------------|-----|------------|--------|-------|-------|
| 4) 2-Fluorophenol         | 0.00          | 112 | 0          | 0.00   | ng/uL |       |
| Spiked Amount 75.000      | Range 18 - 42 |     | Recovery = | 0.00%# |       |       |
| 5) Phenol-d5              | 0.00          | 99  | 0          | 0.00   | ng/uL |       |
| Spiked Amount 75.000      | Range 19 - 32 |     | Recovery = | 0.00%# |       |       |
| 6) 2-Chlorophenol-d4      | 11.89         | 132 | 763        | 0.01   | ng/uL | 0.46  |
| Spiked Amount 75.000      | Range 34 - 84 |     | Recovery = | 0.01%# |       |       |
| 7) 1,2-Dichlorobenzene-d4 | 12.24         | 152 | 321        | 0.01   | ng/uL | -0.21 |
| Spiked Amount 50.000      | Range 26 - 73 |     | Recovery = | 0.02%# |       |       |
| 20) Nitrobenzene-d5       | 0.00          | 82  | 0          | 0.00   | ng/uL |       |
| Spiked Amount 50.000      | Range 55 - 98 |     | Recovery = | 0.00%# |       |       |
| 32) 2-Fluorobiphenyl      | 18.92         | 172 | 2164       | 0.02   | ng/uL | 0.09  |
| Spiked Amount 50.000      | Range 42 - 78 |     | Recovery = | 0.04%# |       |       |
| 33) 2,4,6-Tribromophenol  | 0.00          | 330 | 0          | 0.00   | ng/uL |       |
| Spiked Amount 75.000      | Range 45 - 96 |     | Recovery = | 0.00%# |       |       |
| 62) Terphenyl-d14         | 0.00          | 244 | 0          | 0.00   | ng/uL |       |
| Spiked Amount 50.000      | Range 58 - 98 |     | Recovery = | 0.00%# |       |       |

## Target Compounds

|                                |      |     |   | Qvalue |
|--------------------------------|------|-----|---|--------|
| 2) Pyridine                    | 0.00 | 79  | 0 | N.D.   |
| 3) N-nitrosodimethylamine      | 0.00 | 74  | 0 | N.D.   |
| 8) Phenol                      | 0.00 | 94  | 0 | N.D.   |
| 9) bis(2-Chloroethyl) ether    | 0.00 | 93  | 0 | N.D.   |
| 10) 2-Chlorophenol             | 0.00 | 128 | 0 | N.D.   |
| 11) 1,3-Dichlorobenzene        | 0.00 | 146 | 0 | N.D.   |
| 12) 1,4-Dichlorobenzene        | 0.00 | 146 | 0 | N.D.   |
| 13) 1,2-Dichlorobenzene        | 0.00 | 146 | 0 | N.D.   |
| 14) 2-Methylphenol             | 0.00 | 108 | 0 | N.D.   |
| 15) 2,2'-oxybis(1-Chloropropan | 0.00 | 45  | 0 | N.D.   |
| 16) 3&4-Methylphenol           | 0.00 | 108 | 0 | N.D.   |

(#) = qualifier out of range (m) = manual integration

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## Quantitation Report

(QT Reviewed)

Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Acq On : 7 Dec 2001 2:24 pm  
 Sample : gc/ms unknown  
 Misc :  
 MS Integration Params: RTEINT.P  
 Quant Time: Dec 11 11:44 2001

Vial: 31  
 Operator: GALOS  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Results File: NP112701.RES

Quant Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
 Title : 208 625/TCLP calibration table  
 Last Update : Wed Nov 28 15:33:44 2001  
 Response via : Initial Calibration  
 DataAcq Meth : NP112701

| Compound                        | R.T.  | QIon | Response | Conc | Unit | Qvalue |
|---------------------------------|-------|------|----------|------|------|--------|
| 17) N-Nitrosodi-n-propylamine   | 0.00  | 70   | 0        | N.D. |      |        |
| 18) Hexachloroethane            | 0.00  | 117  | 0        | N.D. |      |        |
| 21) Nitrobenzene                | 0.00  | 77   | 0        | N.D. |      |        |
| 22) Isophorone                  | 0.00  | 82   | 0        | N.D. |      |        |
| 23) 2-Nitrophenol               | 0.00  | 139  | 0        | N.D. |      |        |
| 24) 2,4-Dimethylphenol          | 0.00  | 107  | 0        | N.D. |      |        |
| 25) bis(2-Chloroethoxy)methane  | 0.00  | 93   | 0        | N.D. |      |        |
| 26) 2,4-Dichlorophenol          | 0.00  | 162  | 0        | N.D. |      |        |
| 27) 1,2,4-Trichlorobenzene      | 0.00  | 180  | 0        | N.D. |      |        |
| 28) Naphthalene                 | 0.00  | 128  | 0        | N.D. |      |        |
| 29) Hexachlorobutadiene         | 0.00  | 225  | 0        | N.D. |      |        |
| 30) 4-Chloro-3-methylphenol     | 0.00  | 107  | 0        | N.D. |      |        |
| 34) Hexachlorocyclopentadiene   | 0.00  | 237  | 0        | N.D. |      |        |
| 35) 2,4,6-Trichlorophenol       | 0.00  | 196  | 0        | N.D. |      |        |
| 36) 2,4,5-Trichlorophenol       | 0.00  | 196  | 0        | N.D. |      |        |
| 37) 2-Chloronaphthalene         | 0.00  | 162  | 0        | N.D. |      |        |
| 38) Dimethylphthalate           | 0.00  | 163  | 0        | N.D. |      |        |
| 39) 2,6-Dinitrotoluene          | 0.00  | 165  | 0        | N.D. | d    |        |
| 40) Acenaphthylene              | 0.00  | 152  | 0        | N.D. |      |        |
| 41) Acenaphthene                | 0.00  | 153  | 0        | N.D. |      |        |
| 42) 2,4-Dinitrophenol           | 0.00  | 184  | 0        | N.D. |      |        |
| 43) 4-Nitrophenol               | 0.00  | 65   | 0        | N.D. |      |        |
| 44) 2,4-Dinitrotoluene          | 0.00  | 165  | 0        | N.D. |      |        |
| 45) Diethylphthalate            | 0.00  | 149  | 0        | N.D. |      |        |
| 46) 4-Chlorophenylphenylether   | 0.00  | 204  | 0        | N.D. |      |        |
| 47) Fluorene                    | 0.00  | 166  | 0        | N.D. |      |        |
| 48) Azobenzene/ 1,2-Diphenylhyd | 0.00  | 77   | 0        | N.D. |      |        |
| 50) 4,6-Dinitro-2-methylphenol  | 0.00  | 198  | 0        | N.D. |      |        |
| 51) N-Nitrosodiphenylamine      | 0.00  | 169  | 0        | N.D. |      |        |
| 52) 4-Bromophenyl-phenylether   | 0.00  | 248  | 0        | N.D. |      |        |
| 53) Hexachlorobenzene           | 0.00  | 284  | 0        | N.D. |      |        |
| 54) Pentachlorophenol           | 0.00  | 266  | 0        | N.D. |      |        |
| 55) Phenanthrene                | 0.00  | 178  | 0        | N.D. |      |        |
| 56) Anthracene                  | 0.00  | 178  | 0        | N.D. |      |        |
| 57) Di-n-butylphthalate         | 27.17 | 149  | 4130     | N.D. |      |        |
| 58) Fluoranthene                | 0.00  | 202  | 0        | N.D. |      |        |
| 60) 3,3'-Dichlorobenzidine      | 0.00  | 252  | 0        | N.D. |      |        |

(#) = qualifier out of range (m) = manual integration

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Quantitation Report

(QT Reviewed)

Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Acq On : 7 Dec 2001 2:24 pm  
 Sample : gc/ms unknown  
 Misc :  
 MS Integration Params: RTEINT.P  
 Quant Time: Dec 11 11:44 2001

Vial: 31  
 Operator: GALOS  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Results File: NP112701.RES

Quant Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
 Title : 208 625/TCLP calibration table  
 Last Update : Wed Nov 28 15:33:44 2001  
 Response via : Initial Calibration  
 DataAcq Meth : NP112701

| Compound                       | R.T.  | QIon | Response | Conc | Unit | Qvalue |
|--------------------------------|-------|------|----------|------|------|--------|
| 61) Benzidine                  | 0.00  | 184  | 0        | N.D. |      |        |
| 63) Pyrene                     | 0.00  | 202  | 0        | N.D. |      |        |
| 64) Butylbenzylphthalate       | 0.00  | 149  | 0        | N.D. |      |        |
| 65) Benzo(a)anthracene         | 33.19 | 228  | 4100     | N.D. |      |        |
| 66) Bis(2-ethylhexyl)phthalate | 33.46 | 149  | 3025     | N.D. |      |        |
| 67) Chrysene                   | 33.19 | 228  | 4100     | N.D. |      |        |
| 69) Di-n-Octylphthalate        | 0.00  | 149  | 0        | N.D. |      |        |
| 70) Benzo(b)fluoranthene       | 0.00  | 252  | 0        | N.D. |      |        |
| 71) Benzo(k)fluoranthene       | 0.00  | 252  | 0        | N.D. |      |        |
| 72) Benzo(a)pyrene             | 37.28 | 252  | 4194     | N.D. |      |        |
| 73) Indeno(1,2,3-cd)pyrene     | 0.00  | 276  | 0        | N.D. |      |        |
| 74) Dibenz(a,h)anthracene      | 0.00  | 278  | 0        | N.D. |      |        |
| 75) Benzo(g,h,i)perylene       | 0.00  | 276  | 0        | N.D. |      |        |

(#) = qualifier out of range (m) = manual integration

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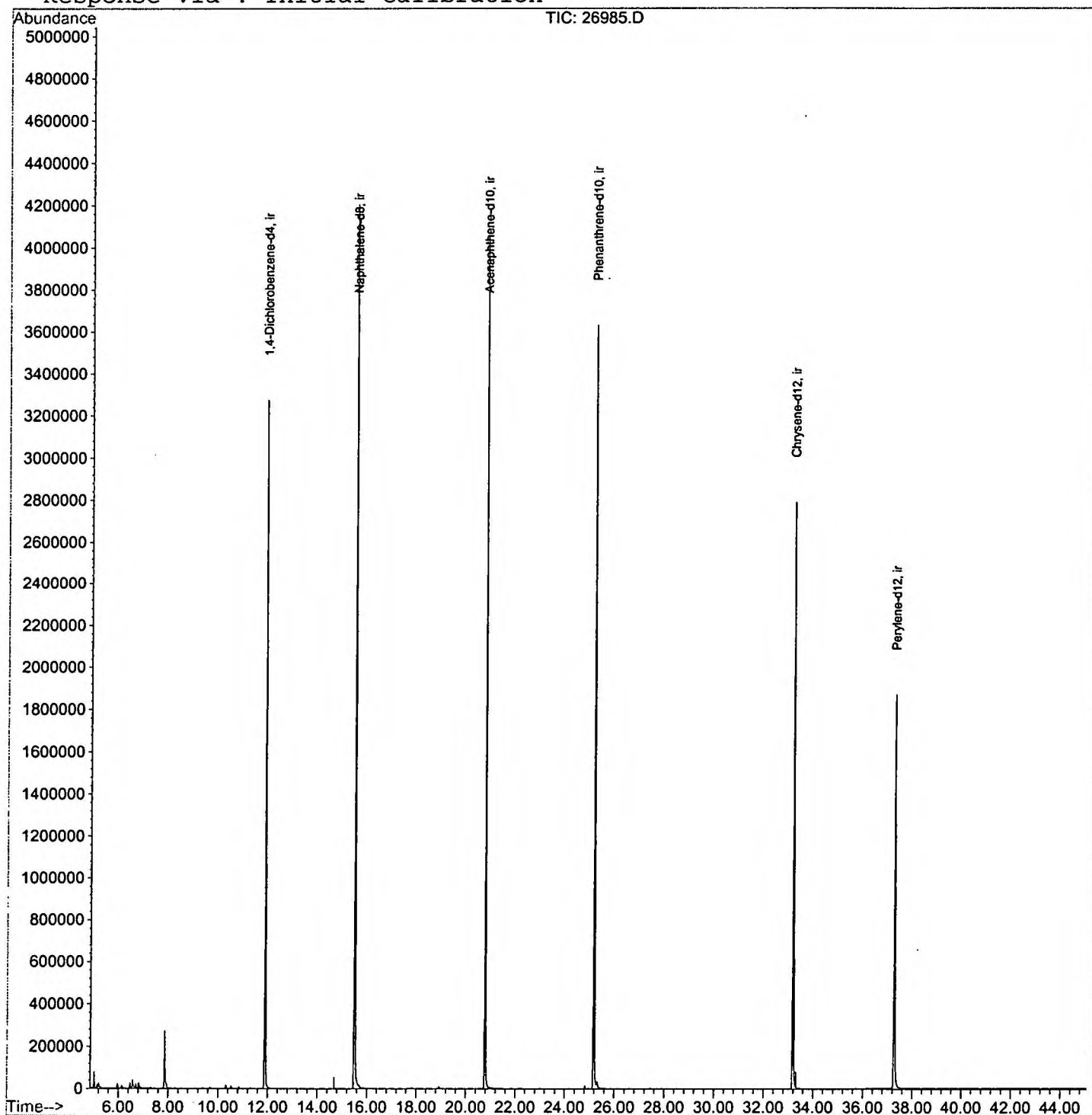
## Quantitation Report

Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
Acq On : 7 Dec 2001 2:24 pm  
Sample : gc/ms unknown  
Misc :  
MS Integration Params: RTEINT.P  
Quant Time: Dec 11 11:44 2001

Vial: 31  
Operator: GALOS  
Inst : GC/MS Ins  
Multiplr: 1.00

Quant Results File: NP112701.RES

Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
Title : 208 625/TCLP calibration table  
Last Update : Wed Nov 28 15:33:44 2001  
Response via : Initial Calibration



## LSC Area Percent Report

Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D

Acq On : 7 Dec 2001 2:24 pm

Sample : gc/ms unknown

Misc :

MS Integration Params: LSCINT.P

Vial: 31

Operator: GALOS

Inst : GC/MS Ins

Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)

Title : 208 625/TCLP calibration table

Smoothing : OFF

Filtering: 5

Sampling : 1

Min Area: 1 % of largest Peak

Start Thrs: 0.2

Max Peaks: 100

Stop Thrs : 0

Peak Location: TOP

If leading or trailing edge &lt; 100 prefer &lt; Baseline drop else tangent &gt;

Peak separation: 5

Signal : TIC

| peak<br># | R.T.<br>min | first<br>scan | max<br>scan | last<br>scan | PK<br>TY | peak<br>height | peak<br>area | peak<br>% max. | % of<br>total |
|-----------|-------------|---------------|-------------|--------------|----------|----------------|--------------|----------------|---------------|
| 1         | 5.028       | 15            | 20          | 25           | rBV      | 77975          | 151907       | 1.78%          | 0.346%        |
| 2         | 6.578       | 185           | 189         | 200          | rVV2     | 42475          | 106266       | 1.25%          | 0.242%        |
| 3         | 7.852       | 324           | 328         | 344          | rVB      | 272432         | 609434       | 7.16%          | 1.387%        |
| 4         | 11.887      | 763           | 768         | 787          | rBV      | 3275354        | 6869095      | 80.65%         | 15.637%       |
| 5         | 15.500      | 1156          | 1162        | 1180         | rBV      | 4203638        | 8516789      | 100.00%        | 19.388%       |
| 6         | 20.773      | 1731          | 1737        | 1752         | rBV      | 4047032        | 8420495      | 98.87%         | 19.168%       |
| 7         | 25.184      | 2211          | 2218        | 2229         | rBV2     | 3630462        | 7849589      | 92.17%         | 17.869%       |
| 8         | 25.321      | 2230          | 2233        | 2243         | rVB      | 32276          | 85302        | 1.00%          | 0.194%        |
| 9         | 33.189      | 3084          | 3091        | 3103         | rBV2     | 2788835        | 6217537      | 73.00%         | 14.154%       |
| 10        | 37.279      | 3528          | 3537        | 3550         | rBV      | 1865627        | 5102799      | 59.91%         | 11.616%       |

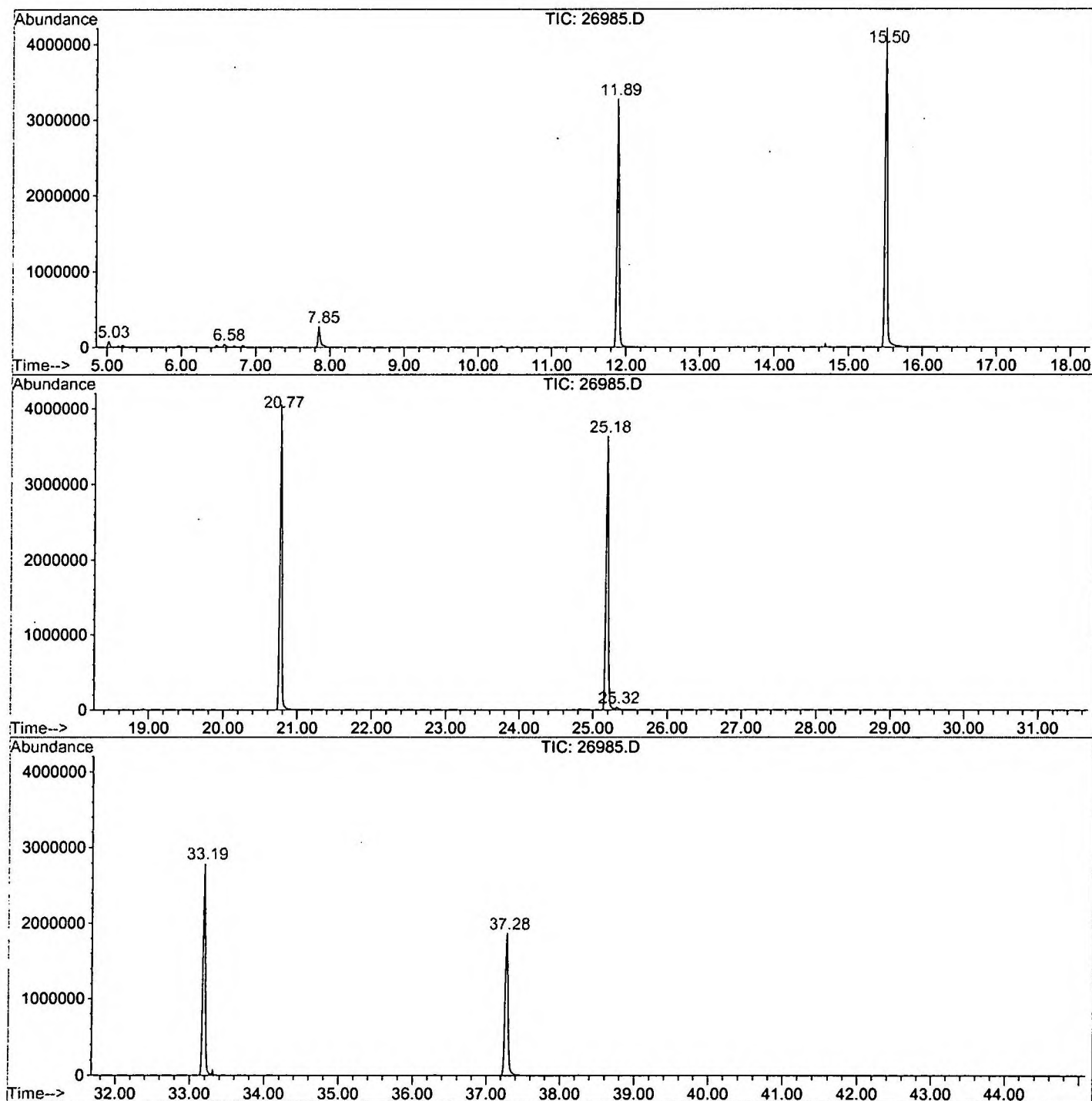
Sum of corrected areas: 43929213

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Tue Dec 11 14:50:18 2001

# LSC Report - Integrated Chromatogram

File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Operator : GALOS  
 Acquired : 7 Dec 2001 2:24 pm using AcqMethod NP112701  
 Instrument : GC/MS Ins  
 Sample Name: gc/ms unknown  
 Misc Info :  
 Vial Number: 31  
 Quant File :NP112701.RES (RTE Integrator)



26985.D NP112701.M

Tue Dec 11 14:50:21 2001

# Library Search Compound Report

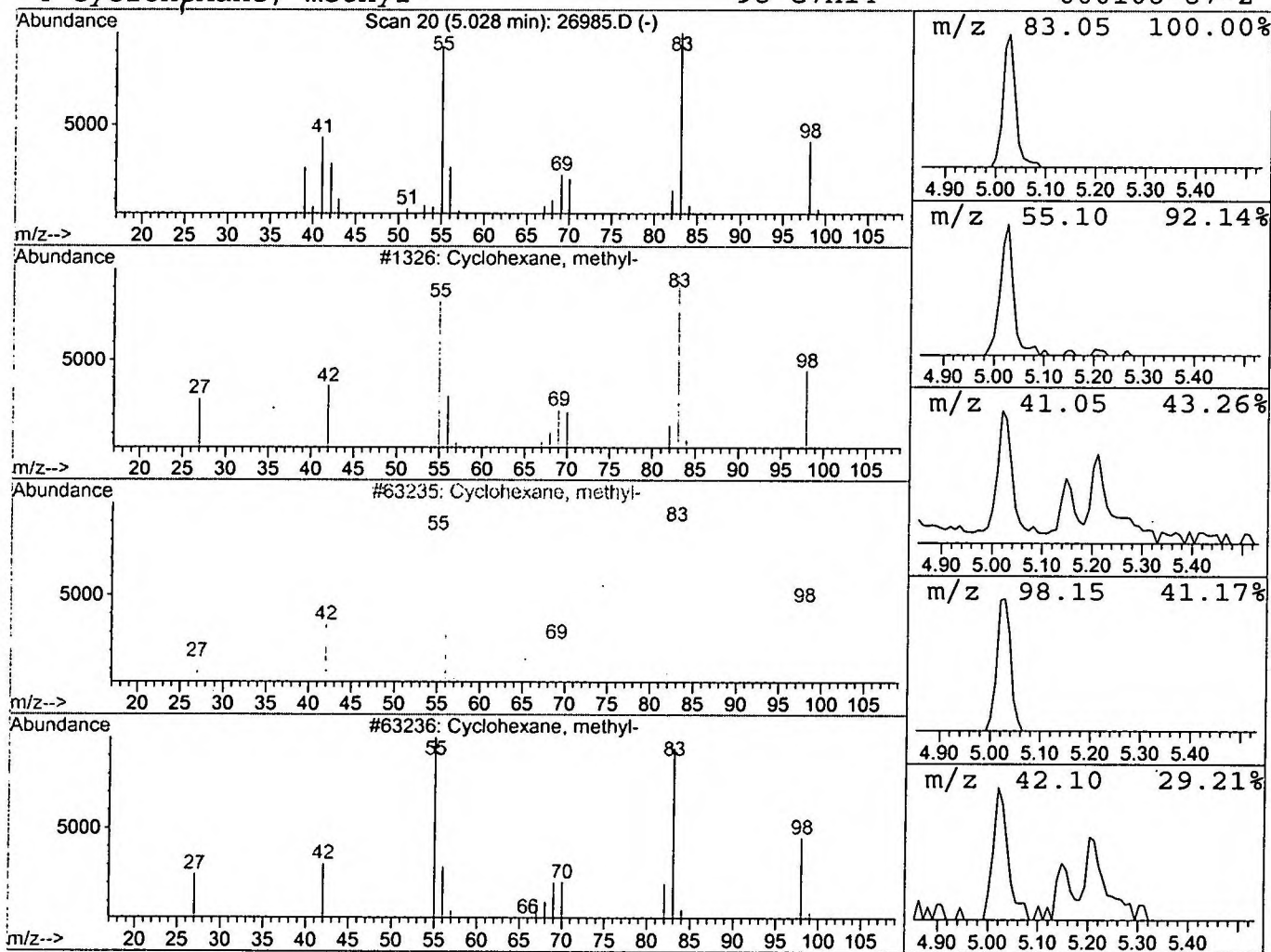
Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Acq On : 7 Dec 2001 2:24 pm  
 Sample : gc/ms unknown  
 Misc :  
 MS Integration Params: LSCINT.P

Vial: 31  
 Operator: GALOS  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
 Title : 208 625/TCLP calibration table  
 Library : C:\DATABASE\NBS75K.L

\*\*\*\*\*  
 Peak Number 1 Cyclohexane, methyl- Concentration Rank 2

| R.T.      | EstConc              | Area   | Relative to ISTD       | R.T.        |      |
|-----------|----------------------|--------|------------------------|-------------|------|
| 5.03      | 0.44 ng/uL           | 151907 | 1,4-Dichlorobenzene-d4 | 11.89       |      |
| Hit# of 5 | Tentative ID         | MW     | MolForm                | CAS#        | Qual |
| 1         | Cyclohexane, methyl- | 98     | C7H14                  | 000108-87-2 | 97   |
| 2         | Cyclohexane, methyl- | 98     | C7H14                  | 000108-87-2 | 97   |
| 3         | Cyclohexane, methyl- | 98     | C7H14                  | 000108-87-2 | 95   |
| 4         | Cyclohexane, methyl- | 98     | C7H14                  | 000108-87-2 | 94   |



# Library Search Compound Report

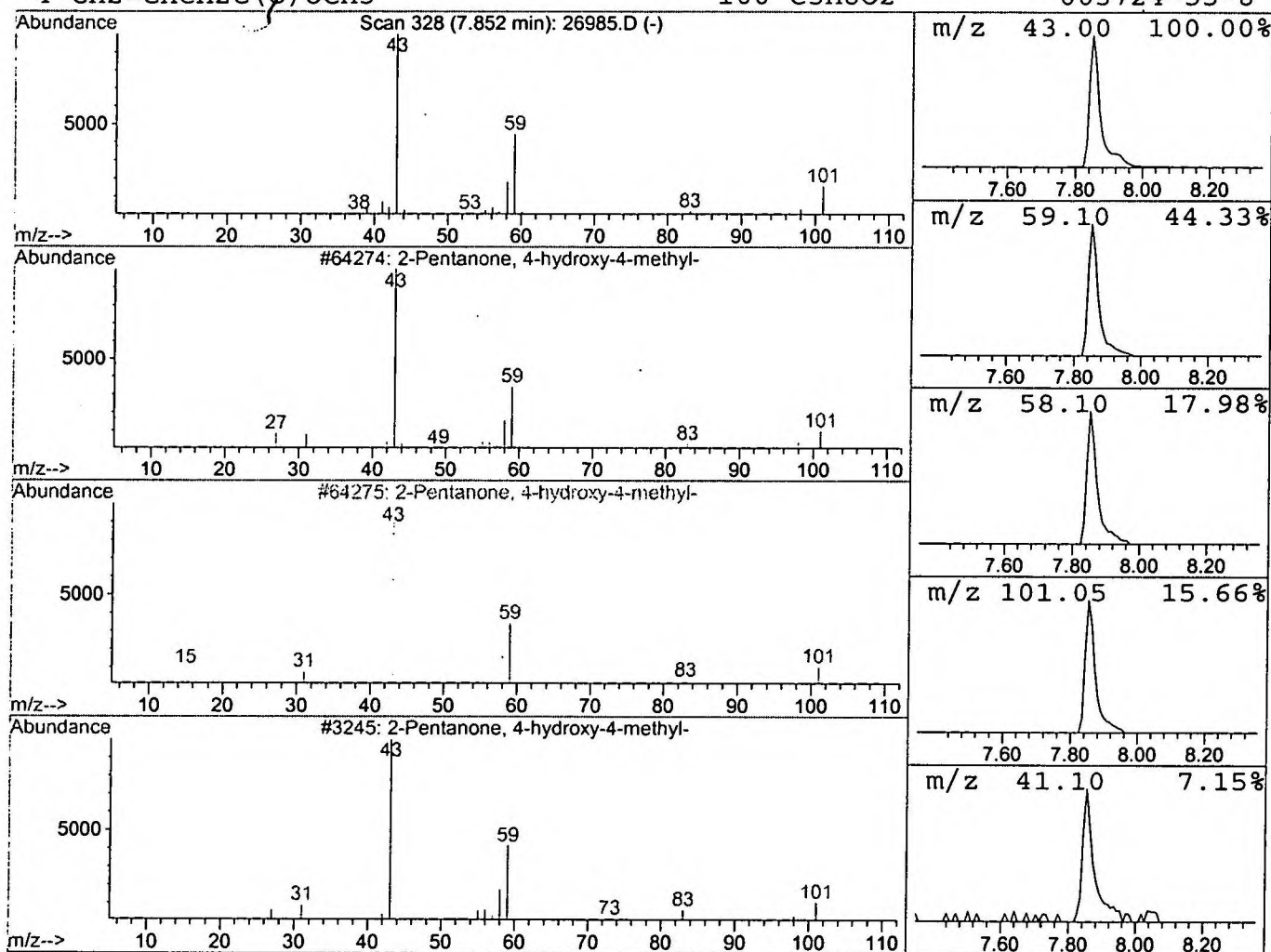
Data File : C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Acq On : 7 Dec 2001 2:24 pm  
 Sample : gc/ms unknown  
 Misc :  
 MS Integration Params: LSCINT.P

Vial: 31  
 Operator: GALOS  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Method : C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
 Title : 208 625/TCLP calibration table  
 Library : C:\DATABASE\NBS75K.L

\*\*\*\*\*  
 Peak Number 2 2-Pentanone, 4-hydroxy-4-methy Concentration Rank 1

| R.T.      | EstConc                          | Area   | Relative to ISTD       | R.T.        |      |
|-----------|----------------------------------|--------|------------------------|-------------|------|
| 7.85      | 1.77 ng/uL                       | 609434 | 1,4-Dichlorobenzene-d4 | 11.89       |      |
| Hit# of 5 | Tentative ID                     | MW     | MolForm                | CAS#        | Qual |
| 1         | 2-Pentanone, 4-hydroxy-4-methyl- | 116    | C6H12O2                | 000123-42-2 | 59   |
| 2         | 2-Pentanone, 4-hydroxy-4-methyl- | 116    | C6H12O2                | 000123-42-2 | 53   |
| 3         | 2-Pentanone, 4-hydroxy-4-methyl- | 116    | C6H12O2                | 000123-42-2 | 43   |
| 4         | CH2=CHCH2C(O)OCH3                | 100    | C5H8O2                 | 003724-55-8 | 9    |



# Tentatively Identified Compound (LSC) summary

Operator ID: GALOS      Date Acquired: 7 Dec 2001      2:24 pm  
 Data File: C:\HPCHEM\1\DATA\DEC0601\26985.D  
 Name: gc/ms unknown  
 Misc:  
 Method: C:\HPCHEM\1\METHODS\NP112701.M (RTE Integrator)  
 Title: 208 625/TCLP calibration table  
 Library Searched: C:\DATABASE\NBS75K.L

| TIC Top Hit name     | RT   | EstConc | Units | Area   | IntStd | ISRT  | ISArea  | ISConc |
|----------------------|------|---------|-------|--------|--------|-------|---------|--------|
| -----                |      |         |       |        |        |       |         |        |
| Cyclohexane, methyl- | 5.03 | 0.4     | ng/uL | 151907 | ISTD01 | 11.89 | 6869100 | 20.0   |
| 2-Pentanone, 4-hydro | 7.85 | 1.8     | ng/uL | 609434 | ISTD01 | 11.89 | 6869100 | 20.0   |

26985.D    NP112701.M      Tue Dec 11 14:50:27 2001