

User: Huhn, William  
 Westchester Dept. of Labs & Research  
 Date: 11/14/2001 Time: 14:54:09

Sample: AD27118  
 Analysis: \$F\_524 Volatile Organic Compounds-Field Blank  
 Units: ug  
 Started: 11/14/01 00:04 Ended: 11/14/01 00:04 Analyst: BH  
 Result source: m:\instruments\206\rrfiles\27118f.rr  
 Page: 1

|    | Component Name                 | Viol | Result | Qualifier | MDL |
|----|--------------------------------|------|--------|-----------|-----|
| 1  | Surr_1,2-DICHLOROETHANE-       |      | 6.4    |           | .5  |
| 2  | Surr_4-BROMOFLUOROBENZE        |      | 4.6    |           | .5  |
| 3  | Dichlorodifluoromethane        |      | < MDL  |           | .5  |
| 4  | Chloromethane                  |      | < MDL  |           | .5  |
| 5  | Vinyl chloride                 |      | < MDL  |           | .5  |
| 6  | Bromomethane                   |      | < MDL  |           | .5  |
| 7  | Chloroethane                   |      | < MDL  |           | .5  |
| 8  | Trichlorofluoromethane         |      | < MDL  |           | .5  |
| 9  | 1,1-Dichloroethene             |      | < MDL  |           | .5  |
| 10 | Methylene Chloride             |      | < MDL  |           | .5  |
| 11 | Methyl tert-butyl ether (MTBE) |      | < MDL  |           | 1.0 |
| 12 | trans-1,2-dichloroethene       |      | < MDL  |           | .5  |
| 13 | 1,1-dichloroethane             |      | < MDL  |           | .5  |
| 14 | 2-butanone (MEK)               |      | < MDL  |           | 2.0 |
| 15 | 2,2-dichloropropane            |      | < MDL  |           | .5  |
| 16 | cis-1,2-dichloroethene         |      | < MDL  |           | .5  |
| 17 | Chloroform                     |      | < MDL  |           | .5  |
| 18 | Bromochloromethane             |      | < MDL  |           | .5  |
| 19 | 1,2-dichloroethane             |      | < MDL  |           | .5  |
| 20 | Surr_TOLUENE-D8                |      | 5.0    |           | .5  |
| 21 | 1,1,1- trichloroethane         |      | < MDL  |           | .5  |
| 22 | 1,1-dichloropropene            |      | < MDL  |           | .5  |
| 23 | Carbon tetrachloride           |      | < MDL  |           | .5  |
| 24 | Benzene                        |      | < MDL  |           | .5  |
| 25 | Trichloroethene                |      | < MDL  |           | .5  |
| 26 | 1,2-dichloropropane            |      | < MDL  |           | .5  |
| 27 | Dibromomethane                 |      | < MDL  |           | .5  |
| 28 | Bromodichloromethane           |      | < MDL  |           | .5  |
| 29 | Methy iso-butyl ketone (MIBK)  |      | < MDL  |           | 1.0 |
| 30 | cis-1,3-dichloropropene        |      | < MDL  |           | .5  |
| 31 | Toluene                        |      | < MDL  |           | .5  |
| 32 | trans-1,3-dichloropropene      |      | < MDL  |           | .5  |
| 33 | 1,1,2-trichloroethane          |      | < MDL  |           | .5  |
| 34 | Tetrachloroethene              |      | < MDL  |           | .5  |
| 35 | 1,3-dichloropropane            |      | < MDL  |           | .5  |
| 36 | Dibromochloromethane           |      | < MDL  |           | 2.0 |
| 37 | Chlorobenzene                  |      | < MDL  |           | .5  |
| 38 | 1,1,1,2-tetrachloroethane      |      | < MDL  |           | .5  |
| 39 | Ethylbenzene                   |      | < MDL  |           | .5  |
| 40 | P & M-xylene                   |      | < MDL  |           | .5  |
| 41 | O-xylene                       |      | < MDL  |           | .5  |
| 42 | Styrene                        |      | < MDL  |           | .5  |
| 43 | 1,1,2,2-tetrachloroethane      |      | < MDL  |           | .5  |
| 44 | Bromoform                      |      | < MDL  |           | 2.0 |
| 45 | Isopropylbenzene               |      | < MDL  |           | .5  |
| 46 | 1,2,3-trichloropropane         |      | < MDL  |           | .5  |
| 47 | N-propylbenzene                |      | < MDL  |           | .5  |
| 48 | Bromobenzene                   |      | < MDL  |           | .5  |

*Acetone - 61*

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Units: ug

Started: 11/14/01 00:04 Ended: 11/14/01 00:04 Analyst: BH

Result source: m:\instruments\206\rrfiles\27118f.rr

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|    | Component Name         | Viol | Result | Qualifier | MDL |
|----|------------------------|------|--------|-----------|-----|
| 49 | 1,3,5-trimethylbenzene |      | < MDL  |           | .5  |
| 50 | 2-chlorotoluene        |      | < MDL  |           | .5  |
| 51 | 4-chlorotoluene        |      | < MDL  |           | .5  |
| 52 | TERT-butylbenzene      |      | < MDL  |           | .5  |
| 53 | 1,2,4-trimethylbenzene |      | < MDL  |           | .5  |
| 54 | SEC-butylbenzene       |      | < MDL  |           | .5  |
| 55 | P-isopropyltoluene     |      | < MDL  |           | .5  |
| 56 | 1,3-dichlorobenzene    |      | < MDL  |           | .5  |
| 57 | 1,4-dichlorobenzene    |      | < MDL  |           | .5  |
| 58 | N-butylbenzene         |      | < MDL  |           | .5  |
| 59 | 1,2-dichlorobenzene    |      | < MDL  |           | .5  |
| 60 | 1,2,4-trichlorobenzene |      | < MDL  |           | .5  |
| 61 | Hexachlobutadiene      |      | < MDL  |           | .5  |
| 62 | Naphthalene            |      | < MDL  |           | .5  |
| 63 | 1,2,3-trichlorobenzene |      | < MDL  |           | .5  |

Quantitation Report

(Not Reviewed)

Data File : C:\HPCHEM\1\DATA\01nov13\27118F.D  
 Acq On : 14 Nov 2001 4:41 am  
 Sample : fb  
 Misc :  
 MS Integration Params: LSCINT.P  
 Quant Time: Nov 14 5:19 2001

Vial: 15  
 Operator:  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Results File: HP110101.RES

Quant Method : C:\HPCHEM\1\METHODS\HP110101.M (RTE Integrator)  
 Title : VOCs BY EPA METHOD 524  
 Last Update : Fri Nov 02 15:17:46 2001  
 Response via : Initial Calibration  
 DataAcq Meth : HP110101

| Internal Standards          | R.T.  | QIon | Response | Conc  | Units   | Dev(Min)     |
|-----------------------------|-------|------|----------|-------|---------|--------------|
| 1) 1,4-DIFLUOROBENZENE      | 14.37 | 114  | 477180   | 5.00  | ug/L    | -0.07        |
| 22) CHLOROBENZENE-D5        | 20.96 | 117  | 426104   | 5.00  | ug/L    | -0.07        |
| 50) 1,4-DICHLOROBENZENE-D4  | 26.13 | 152  | 168802   | 5.00  | ug/L    | -0.07        |
| System Monitoring Compounds |       |      |          |       |         |              |
| 2) 1,2-DICHLOROETHANE-D4    | 13.59 | 65   | 180653   | 6.42  | ug/L    | -0.07        |
| Spiked Amount 5.000         |       |      | Recovery | =     | 128.40% |              |
| 3) 4-BROMOFLUOROBENZENE     | 23.54 | 174  | 113813   | 4.59  | ug/L    | -0.07        |
| Spiked Amount 5.000         |       |      | Recovery | =     | 91.80%  |              |
| 23) TOLUENE-D8              | 17.69 | 98   | 638041   | 4.96  | ug/L    | -0.05        |
| Spiked Amount 5.000         |       |      | Recovery | =     | 99.20%  |              |
| Target Compounds            |       |      |          |       |         |              |
| 10) ACETONE                 | 7.63  | 43   | 158636   | 61.36 | ug/L    | Qvalue<br>94 |

(#) = qualifier out of range (m) = manual integration  
 27118F.D HP110101.M Wed Nov 14 05:20:05 2001

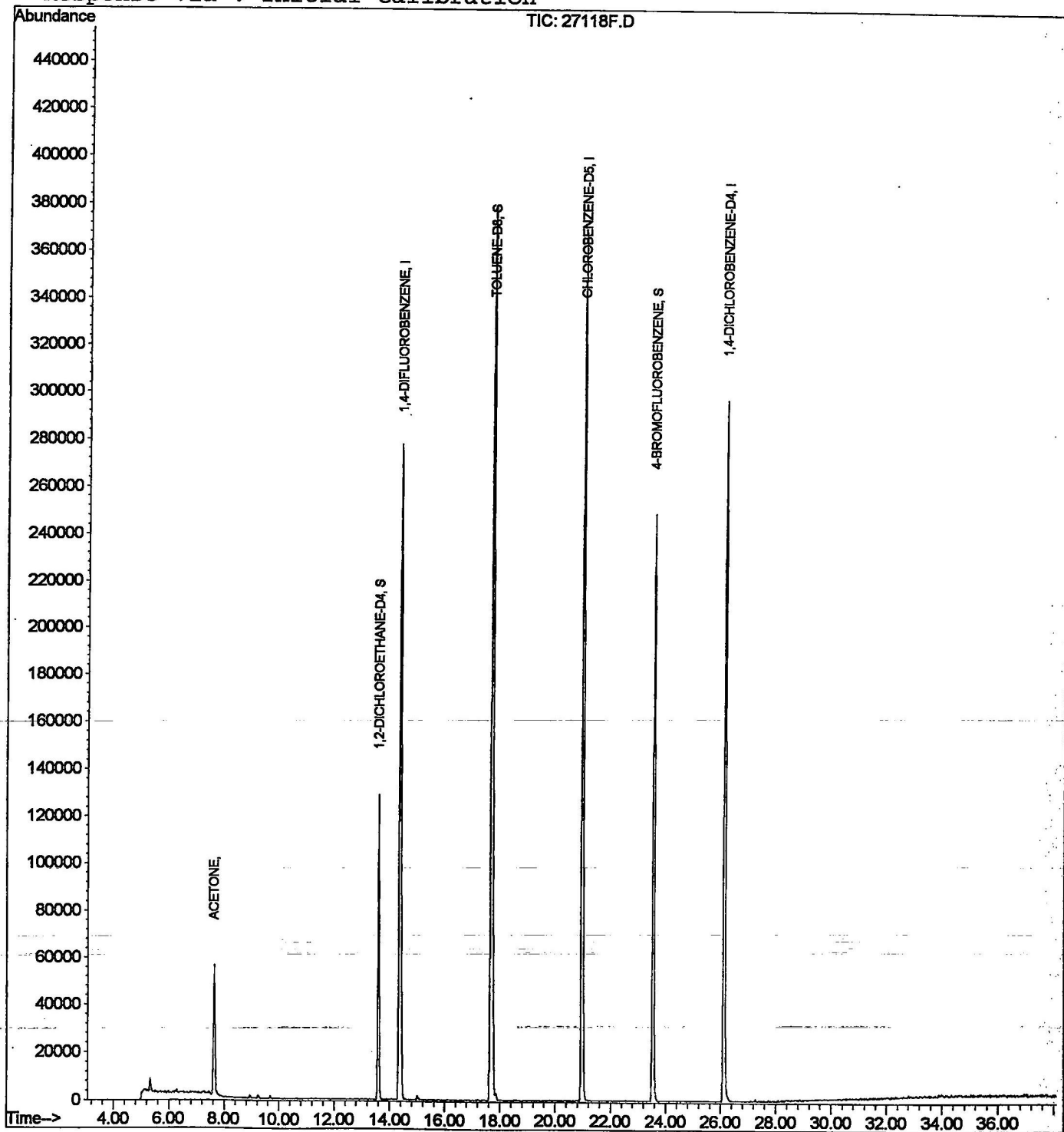
## Quantitation Report

Data File : C:\HPCHEM\1\DATA\01nov13\27118F.D  
Acq On : 14 Nov 2001 4:41 am  
Sample : fb  
Misc :  
MS Integration Params: LSCINT.P  
Quant Time: Nov 14 5:19 2001

Vial: 15  
Operator:  
Inst : GC/MS Ins  
Multiplr: 1.00

Quant Results File: HP110101.RES

Method : C:\HPCHEM\1\METHODS\HP110101.M (RTE Integrator)  
Title : VOCs BY EPA METHOD 524  
Last Update : Fri Nov 02 15:17:46 2001  
Response via : Initial Calibration



# Library Search Compound Report

Data File : C:\HPCHEM\1\DATA\01nov13\27118F.D  
 Acq On : 14 Nov 2001 4:41 am  
 Sample : fb  
 Misc :  
 MS Integration Params: LSCINT.P

Vial: 15  
 Operator:  
 Inst : GC/MS Ins  
 Multiplr: 1.00

Quant Method : C:\HPCHEM\1\METHODS\HP110101.M (RTE Integrator)  
 Title : VOCs BY EPA METHOD 524  
 Library : C:\DATABASE\NBS75K.L

\*\*\*\*\*  
 Peak Number 1 1-(Propenylthio)prop-1-ene Concentration Rank 1

| R.T.  | EstConc   | Area | Relative to ISTD    | R.T.  |
|-------|-----------|------|---------------------|-------|
| 15.00 | 0.02 ug/L | 6052 | 1,4-DIFLUOROBENZENE | 14.37 |

| Hit# | of | Tentative ID                        | MW  | MolForm  | CAS#        | Qual |
|------|----|-------------------------------------|-----|----------|-------------|------|
| 1    | 5  | 1-(Propenylthio)prop-1-ene          | 114 | C6H10S   | 000000-00-0 | 3    |
| 2    |    | 2,4-Imidazolidinedione, 3-methyl-   | 114 | C4H6N2O2 | 006843-45-4 | 3    |
| 3    |    | 3-Cyclobutene-1,2-dione, 3,4-dihydr | 114 | C4H2O4   | 002892-51-5 | 2    |
| 4    |    | 2,4,5-Trioxoimidazolidine           | 114 | C3H2N2O3 | 000120-89-8 | 2    |

