

User: Huhn, William
 Westchester Dept. of Labs & Research
 Date: 11/13/2001 Time: 16:31:07

Sample: AD27008

Analysis: \$F_524 Volatile Organic Compounds-Field Blank

Units: ug

Started: 11/10/01 00:04 Ended: 11/10/01 00:04 Analyst: BH

Result source: m:\instruments\206\rrfiles\27008f.rr

Page: 1

	Component Name	Vol	Result	Qualifier	MDL
1	Surr_1,2-DICHLOROETHANE-		6.6		.5
2	Surr_4-BROMOFLUOROBENZE		5.1		.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		5.1		.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 - 21

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	Component Name	Viol	Result	Quantifier	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\01nov9\27008F.D
 Acq On : 10 Nov 2001 4:27 am
 Sample : fb
 Misc :
 MS Integration Params: LSCINT.P
 Quant Time: Nov 10 5:06 2001

Vial: 7
 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	14.39	114	429946	5.00	ug/L	-0.05
22) CHLOROBENZENE-D5	21.00	117	388901	5.00	ug/L	-0.03
50) 1,4-DICHLOROBENZENE-D4	26.16	152	156810	5.00	ug/L	-0.03
System Monitoring Compounds						
2) 1,2-DICHLOROETHANE-D4	13.62	65	168293	6.63	ug/L	-0.03
Spiked Amount 5.000			Recovery	=	132.60%	
3) 4-BROMOFLUOROBENZENE	23.57	174	113618	5.08	ug/L	-0.03
Spiked Amount 5.000			Recovery	=	101.60%	
23) TOLUENE-D8	17.71	98	594650	5.06	ug/L	-0.03
Spiked Amount 5.000			Recovery	=	101.20%	
Target Compounds						
10) ACETONE	7.66	43	49224	21.13	ug/L	Qvalue 96
16) 2-BUTANONE (MEK)	11.36	43	1271	0.27	ug/L	57
72) 1,2,3-TRICHLOROBENZENE	31.87	180	3806	<u>0.38</u>	ug/L	86

C/S

(#) = qualifier out of range (m) = manual integration
 27008F.D HP110101.M Sat Nov 10 05:06:40 2001

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Quantitation Report

Data File : C:\HPCHEM\1\DATA\01nov9\27008F.D

Acq On : 10 Nov 2001 4:27 am

Sample : fb

Misc :

MS Integration Params: LSCINT.P

Quant Time: Nov 10 5:06 2001

Vial: 7

Operator:

Inst : GC/MS Ins

Multiplr: 1.00

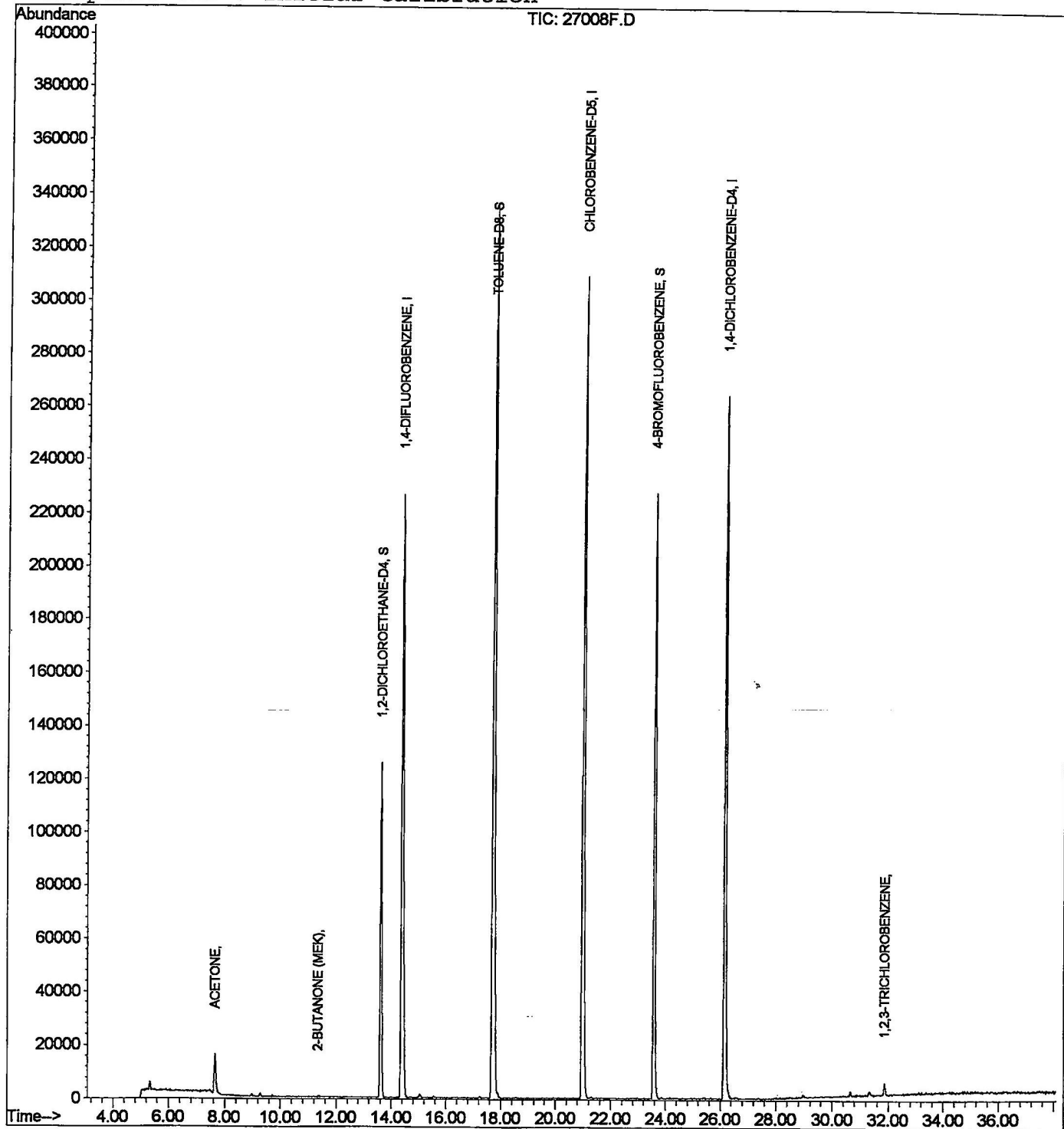
Quant Results File: HP110101.RES

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\01nov9\27008F.D
 Acq On : 10 Nov 2001 4:27 am
 Sample : fb
 Misc :
 MS Integration Params: LSCINT.P

Vial: 7
 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.	
15.05	0.03 ug/L	6252	1,4-DIFLUOROBENZENE		14.39	
Hit# of	5	Tentative ID	MW	MolForm	CAS#	Qual
1		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

