



***Air
Toxics LTD.***
Laboratory Services Since 1989

Electronic Comprehensive Validation Package (eCVP)



AN ENVIRONMENTAL ANALYTICAL LABORATORY

COMPREHENSIVE VALIDATION PACKAGE

Modified TO-15

INVENTORY SHEET

Work Order #: 0706440

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Comments:

Completed by:

Judy Lee

Judy Lee / Document Control

7/10/07

(Signature)

(Print Name & Title)

(Date)



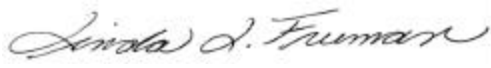
AN ENVIRONMENTAL ANALYTICAL LABORATORY

WORK ORDER #: 0706440

Work Order Summary

CLIENT:	Ms. Sarah Aldridge GEI Consultants, Inc. 455 Winding Brook Dr. Suite 201 Glastonbury, CT 06033	BILL TO:	Ms. Sarah Aldridge GEI Consultants, Inc. 455 Winding Brook Dr. Suite 201 Glastonbury, CT 06033
PHONE:	860-368-5300	P.O. #	NR
FAX:	860-368-5307	PROJECT #	061140-8-1703 Bay Shore OUI
DATE RECEIVED:	06/21/2007	CONTACT:	Bryanna Langley
DATE COMPLETED:	07/05/2007		

<u>FRACTION #</u>	<u>NAME</u>	<u>TEST</u>	<u>RECEIPT VAC./PRES.</u>
01A	AMS 5 DW (4148)	Modified TO-15	7.0 "Hg
02A	AMS 5 DW (3746)	Modified TO-15	7.5 "Hg
02AA	AMS 5 DW (3746) Lab Duplicate	Modified TO-15	7.5 "Hg
03A	AMS 2 UW	Modified TO-15	7.5 "Hg
04A	TRIP BLANK	Modified TO-15	4.4 psi
05A	Lab Blank	Modified TO-15	NA
06A	CCV	Modified TO-15	NA
07A	LCS	Modified TO-15	NA

CERTIFIED BY:  DATE: 07/05/07

Laboratory Director

Certification numbers: CA NELAP - 02110CA, LA NELAP/LELAP- AI 30763, NJ NELAP - CA004
NY NELAP - 11291, UT NELAP - 9166389892

Name of Accrediting Agency: NELAP/Florida Department of Health, Scope of Application: Clean Air Act,
Accreditation number: E87680, Effective date: 07/01/07, Expiration date: 06/30/08
Air Toxics Ltd. certifies that the test results contained in this report meet all requirements of the NELAC standards

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LABORATORY NARRATIVE
Modified TO-15
GEI Consultants, Inc.
Workorder# 0706440

Three 6 Liter Summa Canister and one 6 Liter Summa Canister (100% Certified) samples were received on June 21, 2007. The laboratory performed analysis via modified EPA Method TO-15 using GC/MS in the full scan mode. The method involves concentrating up to 0.2 liters of air. The concentrated aliquot is then flash vaporized and swept through a water management system to remove water vapor. Following dehumidification, the sample passes directly into the GC/MS for analysis.

This workorder was independently validated prior to submittal using 'USEPA National Functional Guidelines' as generally applied to the analysis of volatile organic compounds in air. A rules-based, logic driven, independent validation engine was employed to assess completeness, evaluate pass/fail of relevant project quality control requirements and verification of all quantified amounts.

Method modifications taken to run these samples are summarized in the below table. Specific project requirements may over-ride the ATL modifications.

<i>Requirement</i>	<i>TO-15</i>	<i>ATL Modifications</i>
Daily CCV	+/- 30% Difference	<= 30% Difference with two allowed out up to <=40%.; flag and narrate outliers
Sample collection media	Summa canister	ATL recommends use of summa canisters to insure data defensibility, but will report results from Tedlar bags at client request
Method Detection Limit	Follow 40CFR Pt.136 App. B	The MDL met all relevant requirements in Method TO-15 (statistical MDL less than the LOQ). The concentration of the spiked replicate may have exceeded 10X the calculated MDL in some cases

Receiving Notes

Sample identifications on the Chain of Custody (COC) were not unique for samples AMS 5 DW (4148) and AMS 5 DW (3746). The canister numbers were added to each of the sample identifications to ensure uniqueness.

Analytical Notes

There were no analytical discrepancies.

Definition of Data Qualifying Flags

Eight qualifiers may have been used on the data analysis sheets and indicates as follows:

- B - Compound present in laboratory blank greater than reporting limit (background subtraction no performed).
- J - Estimated value.
- E - Exceeds instrument calibration range.

- S - Saturated peak.
- Q - Exceeds quality control limits.
- U - Compound analyzed for but not detected above the reporting limit.
- UJ- Non-detected compound associated with low bias in the CCV
- N - The identification is based on presumptive evidence.

File extensions may have been used on the data analysis sheets and indicates as follows:

- a-File was requantified
- b-File was quantified by a second column and detector
- r1-File was requantified for the purpose of reissue

Table 1

Client Sample ID	Lab Sample ID	Date Collected	Date Received	Date Extracted	Sample	Sample Extract		Sample Condition
					Holding Time (Days)	Date Analyzed	Holding Time (Days)	
AMS 5 DW (4148)	0706440-01A	6/20/2007	6/21/2007	NA	13	7/ 3/2007	NA	Good
AMS 5 DW (3746)	0706440-02A	6/20/2007	6/21/2007	NA	13	7/ 3/2007	NA	Good
AMS 5 DW (3746) Lab D	0706440-02AA	6/20/2007	6/21/2007	NA	13	7/ 3/2007	NA	Good
AMS 2 UW	0706440-03A	6/20/2007	6/21/2007	NA	13	7/ 3/2007	NA	Good
TRIP BLANK	0706440-04A	6/20/2007	6/21/2007	NA	13	7/ 3/2007	NA	Good
Lab Blank	0706440-05A	NA	NA	NA	NA	7/ 3/2007	NA	Good
CCV	0706440-06A	NA	NA	NA	NA	7/ 3/2007	NA	Good
LCS	0706440-07A	NA	NA	NA	NA	7/ 3/2007	NA	Good

Sample Results and Raw Data



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Summary of Detected Compounds MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

Client Sample ID: AMS 5 DW (4148)

Lab ID#: 0706440-01A

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Methylene Chloride	0.88	1.0	3.0	3.5
Acetone	3.5	4.5	8.3	11



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 5 DW (4148)

Lab ID#: 0706440-01A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070316	Date of Collection:	6/20/07
Dil. Factor:	1.75	Date of Analysis:	7/3/07 06:42 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Freon 12	0.88	Not Detected	4.3	Not Detected
Freon 114	0.88	Not Detected	6.1	Not Detected
Vinyl Chloride	0.88	Not Detected	2.2	Not Detected
Bromomethane	0.88	Not Detected	3.4	Not Detected
Chloroethane	0.88	Not Detected	2.3	Not Detected
Freon 11	0.88	Not Detected	4.9	Not Detected
1,1-Dichloroethene	0.88	Not Detected	3.5	Not Detected
Freon 113	0.88	Not Detected	6.7	Not Detected
Methylene Chloride	0.88	1.0	3.0	3.5
1,1-Dichloroethane	0.88	Not Detected	3.5	Not Detected
cis-1,2-Dichloroethene	0.88	Not Detected	3.5	Not Detected
Chloroform	0.88	Not Detected	4.3	Not Detected
1,1,1-Trichloroethane	0.88	Not Detected	4.8	Not Detected
Carbon Tetrachloride	0.88	Not Detected	5.5	Not Detected
Benzene	0.88	Not Detected	2.8	Not Detected
1,2-Dichloroethane	0.88	Not Detected	3.5	Not Detected
Trichloroethene	0.88	Not Detected	4.7	Not Detected
1,2-Dichloropropane	0.88	Not Detected	4.0	Not Detected
cis-1,3-Dichloropropene	0.88	Not Detected	4.0	Not Detected
Toluene	0.88	Not Detected	3.3	Not Detected
trans-1,3-Dichloropropene	0.88	Not Detected	4.0	Not Detected
1,1,2-Trichloroethane	0.88	Not Detected	4.8	Not Detected
Tetrachloroethene	0.88	Not Detected	5.9	Not Detected
1,2-Dibromoethane (EDB)	0.88	Not Detected	6.7	Not Detected
Chlorobenzene	0.88	Not Detected	4.0	Not Detected
Ethyl Benzene	0.88	Not Detected	3.8	Not Detected
m,p-Xylene	0.88	Not Detected	3.8	Not Detected
o-Xylene	0.88	Not Detected	3.8	Not Detected
Styrene	0.88	Not Detected	3.7	Not Detected
1,1,2,2-Tetrachloroethane	0.88	Not Detected	6.0	Not Detected
1,3,5-Trimethylbenzene	0.88	Not Detected	4.3	Not Detected
1,2,4-Trimethylbenzene	0.88	Not Detected	4.3	Not Detected
1,3-Dichlorobenzene	0.88	Not Detected	5.3	Not Detected
1,4-Dichlorobenzene	0.88	Not Detected	5.3	Not Detected
alpha-Chlorotoluene	0.88	Not Detected	4.5	Not Detected
1,2-Dichlorobenzene	0.88	Not Detected	5.3	Not Detected
1,3-Butadiene	0.88	Not Detected	1.9	Not Detected
Hexane	0.88	Not Detected	3.1	Not Detected
Cyclohexane	0.88	Not Detected	3.0	Not Detected



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 5 DW (4148)

Lab ID#: 0706440-01A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070316	Date of Collection:	6/20/07
Dil. Factor:	1.75	Date of Analysis:	7/3/07 06:42 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Heptane	0.88	Not Detected	3.6	Not Detected
Bromodichloromethane	0.88	Not Detected	5.9	Not Detected
Dibromochloromethane	0.88	Not Detected	7.4	Not Detected
Cumene	0.88	Not Detected	4.3	Not Detected
Propylbenzene	0.88	Not Detected	4.3	Not Detected
Chloromethane	3.5	Not Detected	7.2	Not Detected
1,2,4-Trichlorobenzene	3.5	Not Detected	26	Not Detected
Hexachlorobutadiene	3.5	Not Detected	37	Not Detected
Acetone	3.5	4.5	8.3	11
Carbon Disulfide	0.88	Not Detected	2.7	Not Detected
2-Propanol	3.5	Not Detected	8.6	Not Detected
trans-1,2-Dichloroethene	0.88	Not Detected	3.5	Not Detected
2-Butanone (Methyl Ethyl Ketone)	0.88	Not Detected	2.6	Not Detected
Tetrahydrofuran	0.88	Not Detected	2.6	Not Detected
1,4-Dioxane	3.5	Not Detected	13	Not Detected
4-Methyl-2-pentanone	0.88	Not Detected	3.6	Not Detected
2-Hexanone	3.5	Not Detected	14	Not Detected
Bromoform	0.88	Not Detected	9.0	Not Detected
4-Ethyltoluene	0.88	Not Detected	4.3	Not Detected
Ethanol	3.5	Not Detected	6.6	Not Detected
Methyl tert-butyl ether	0.88	Not Detected	3.2	Not Detected
3-Chloropropene	3.5	Not Detected	11	Not Detected
2,2,4-Trimethylpentane	0.88	Not Detected	4.1	Not Detected
Naphthalene	3.5	Not Detected	18	Not Detected

Container Type: 6 Liter Summa Canister

Surrogates	%Recovery	Method Limits
Toluene-d8	98	70-130
1,2-Dichloroethane-d4	107	70-130
4-Bromofluorobenzene	104	70-130

Report Date: 05-Jul-2007 09:31

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070316.d
 Lab Smp Id: 0706440-01A
 Inj Date : 03-JUL-2007 18:42
 Operator : jdj Inst ID: msd8.i
 Smp Info : 200ml #4148
 Misc Info : 7.0"Hg-5.0psi
 Comment :
 Method : /var/chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 14:39 dpage Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1
 Dil Factor: 1.75000
 Integrator: HP RTE Compound Sublist: AT04.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS								
RT	EXP RT	(REL RT)	MASS	RESPONSE		TARGET RANGE	RATIO	
				(PPBV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====

* 68	Bromochloromethane					CAS #: 74-97-5		
7.387	7.387	(1.000)	130	247531	25.0000	70.00- 130.00	100.00	
7.387	7.387	(1.000)	128	193034		47.25- 107.25	77.98	
7.387	7.387	(1.000)	49	377150		115.07- 175.07	152.36	

* 88	1,4-Difluorobenzene					CAS #: 540-36-3		
9.267	9.267	(1.000)	114	1055614	25.0000	70.00- 130.00	100.00	
9.267	9.267	(1.000)	88	162248		0.00- 44.97	15.37	

* 125	Chlorobenzene-d5					CAS #: 3114-55-4		
14.576	14.576	(1.000)	117	791246	25.0000	70.00- 130.00	100.00	
14.576	14.576	(1.000)	82	443362		0.00- 30.00	56.03	

\$ 82	1,2-Dichloroethane-d4					CAS #: 17060-07-0		
8.465	8.465	(1.146)	65	345682	26.6832	26.683 70.00- 130.00	100.00	
8.465	8.465	(1.146)	67	181350		0.00- 30.00	52.46	

\$ 104	Toluene-d8					CAS #: 2037-26-5		
12.115	12.115	(1.307)	98	899495	24.6133	24.613 70.00- 130.00	100.00	
12.115	12.115	(1.307)	70	90477		0.00- 30.00	10.06	

CONCENTRATIONS

ON-COL FINAL

RT EXP RT (REL RT) MASS RESPONSE (PPEV) (PPBV) TARGET RANGE RATIO
== =====

\$ 104 Toluene-d8 (continued)

12.115 12.115 (1.307) 100 590740 0.00- 30.00 65.67

\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207 16.207 (1.112) 174 488026 25.9228 25.923 70.00- 130.00 100.00

16.207 16.207 (1.112) 95 590024 86.34- 146.34 120.90

16.207 16.207 (1.112) 176 468200 66.11- 126.11 95.94

30 Acetone

CAS #: 67-64-1

4.124 4.124 (0.558) 58 21706 2.55889 4.478 70.00- 130.00 100.00

4.152 4.124 (0.562) 43 66061 0.00- 30.00 304.34

40 Methylene Chloride

CAS #: 75-09-2

4.816 4.815 (0.652) 49 10981 0.57912 1.013 70.00- 130.00 100.00

4.816 4.815 (0.652) 84 6706 37.67- 97.67 61.07

4.843 4.815 (0.656) 51 4714 0.00- 30.00 42.93

Report Date: 05-Jul-2007 09:31

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARYInstrument ID: msd8.i
Lab File ID: 8070316.d
Lab Smp Id: 0706440-01ACalibration Date: 03-JUL-2007
Calibration Time: 09:41

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdg

Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 7.0"Hg-5.0psi

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	247531	-28.45
88 1,4-Difluorobenze	1512218	907331	2117105	1055614	-30.19
125 Chlorobenzene-d5	1120246	672148	1568344	791246	-29.37

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
Sample Matrix: GAS Fraction: VOA
Lab Smp Id: 0706440-01A
Level: LOW Operator: jdg
Data Type: MS DATA SampleType: SAMPLE
SpikeList File: AT04+NA-2.spk Quant Type: ISTD
Sublist File: AT04.sub
Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m
Misc Info: 7.0"Hg-5.0psi

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	26.683	106.73	70-130
\$ 104 Toluene-d8	25.000	24.613	98.45	70-130
\$ 140 Bromofluorobenzene	25.000	25.923	103.69	70-130

Data File: /chem/msd8.1/8-03jul.b/8070316.d

Date: 03-JUL-2007 18:42

Client ID:

Sample Info: 200ml #4148

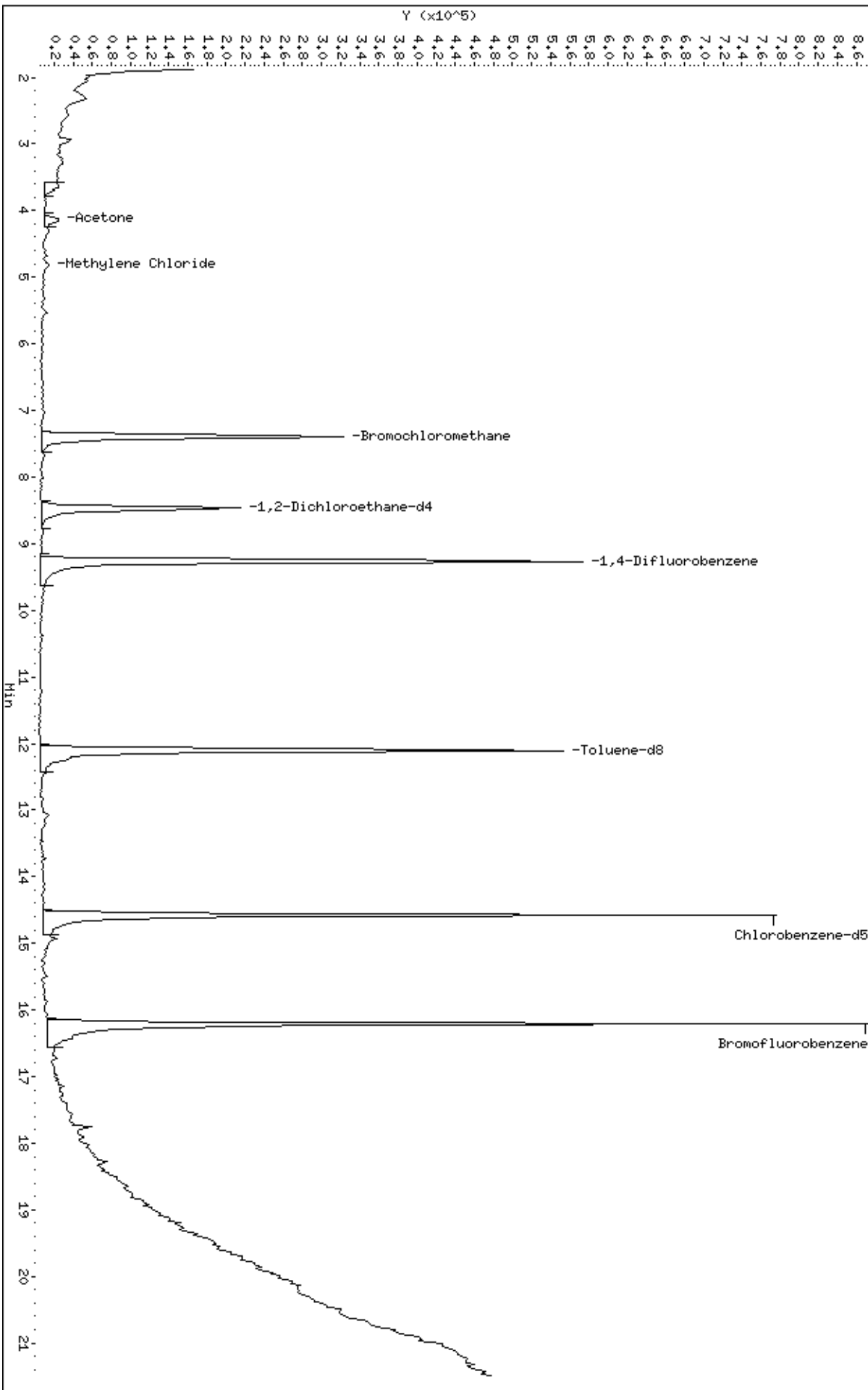
Column phase: RTX-624

Instrument: msd8.1

Operator: jdg

Column diameter: 0.53

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Date : 03-JUL-2007 18:42

Client ID:

Instrument: msd8.i

Sample Info: 200ml #4148

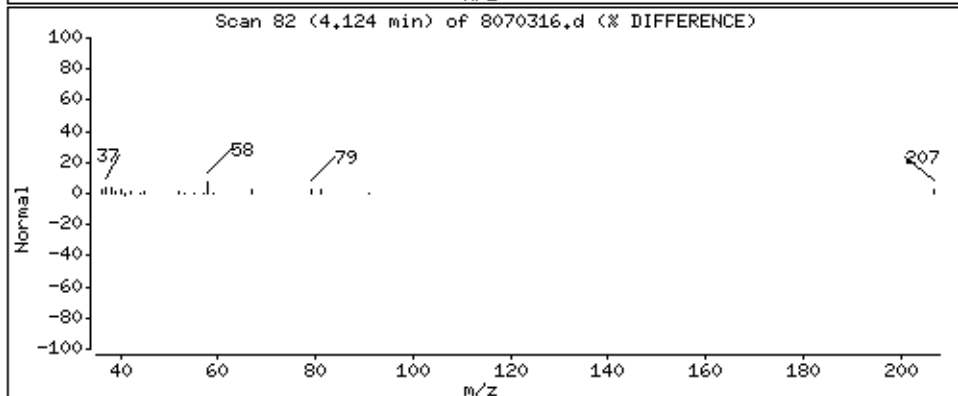
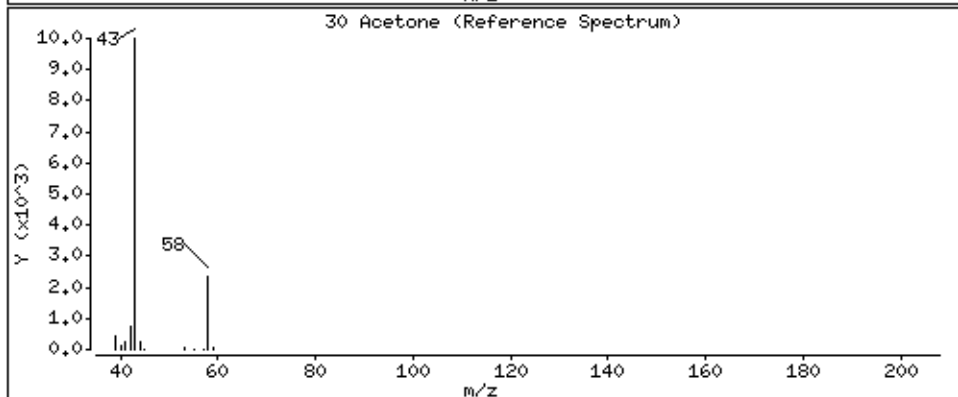
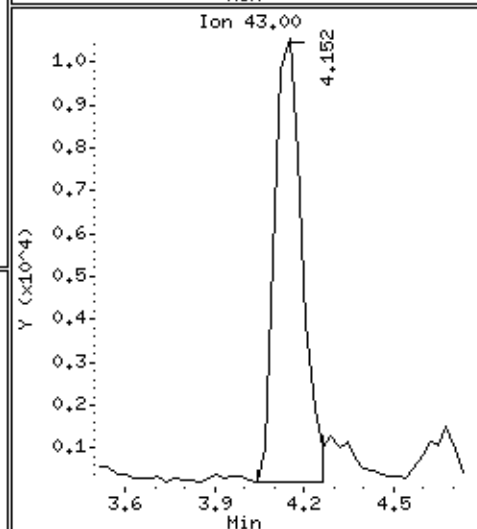
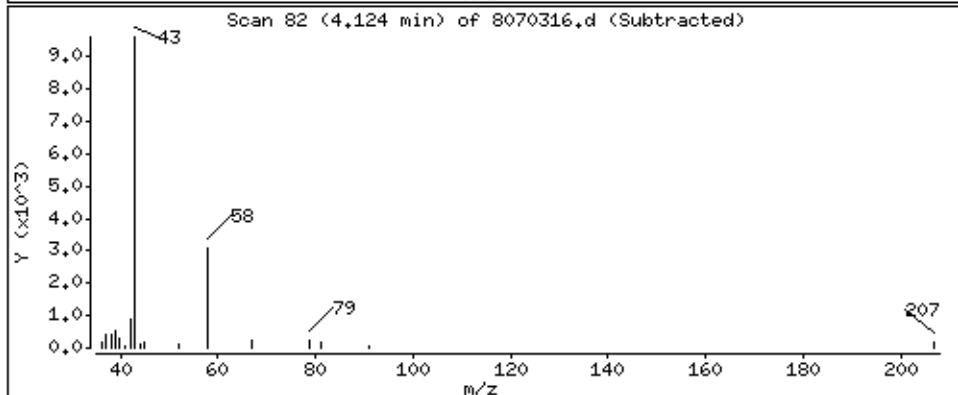
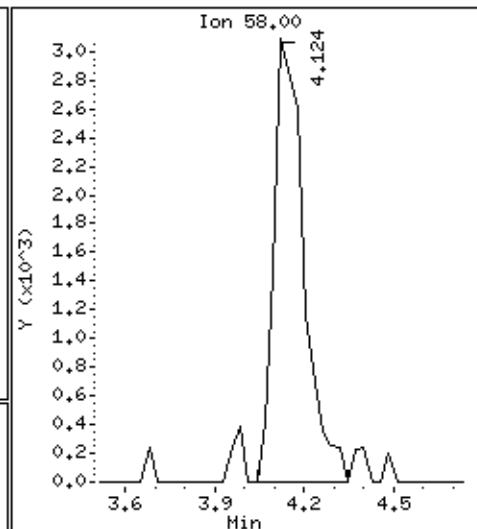
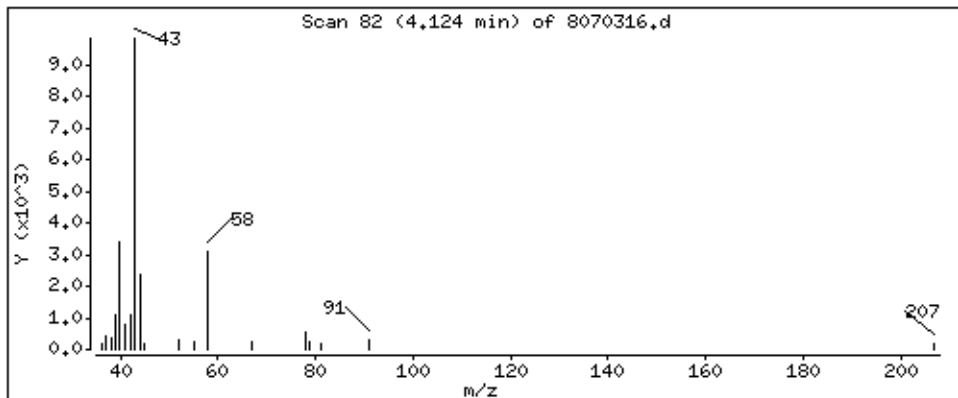
Operator: jdg

Column phase: RTX-624

Column diameter: 0.53

30 Acetone

Concentration: 4.478 PPBV



Date : 03-JUL-2007 18:42

Client ID:

Instrument: msd8,i

Sample Info: 200ml #4148

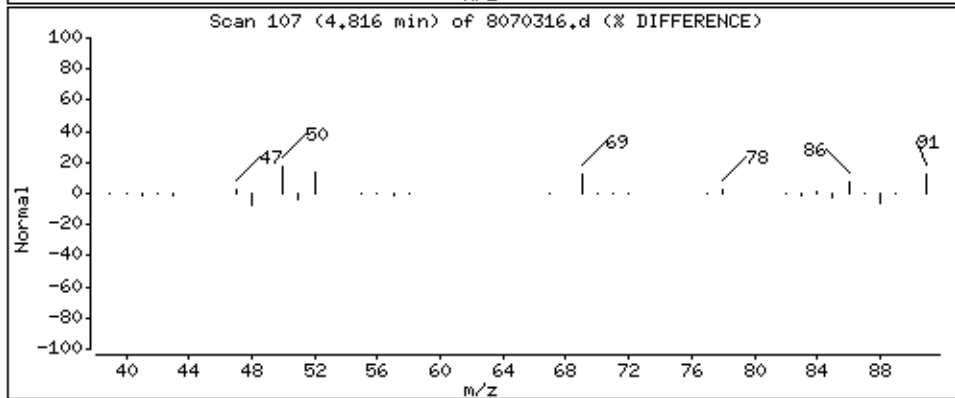
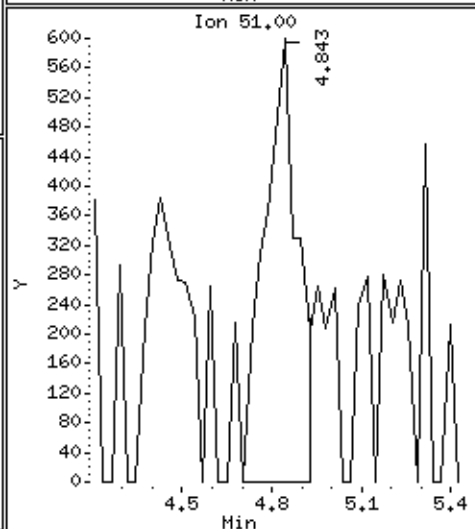
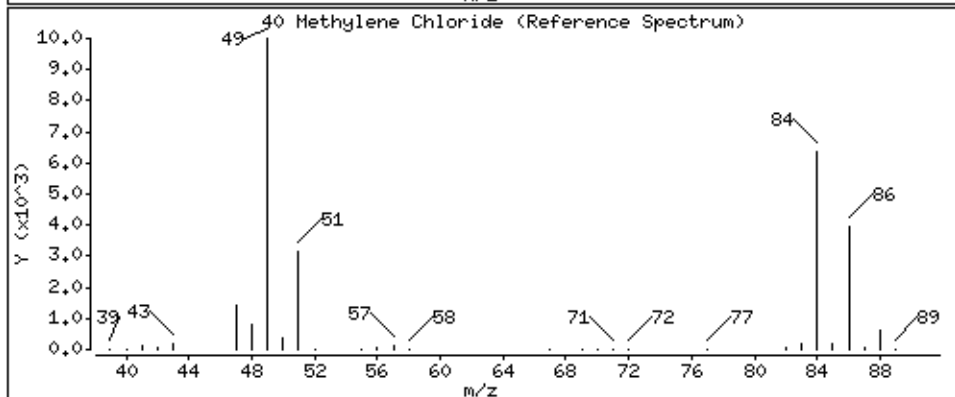
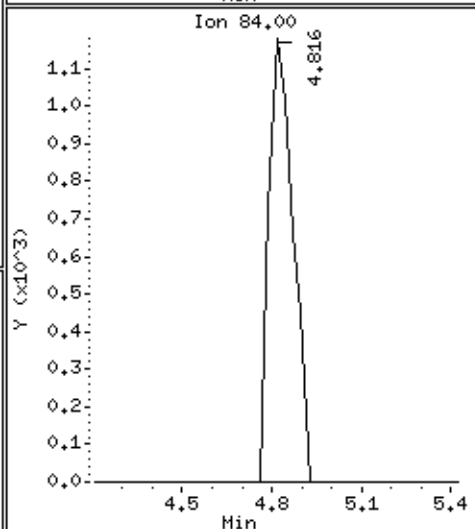
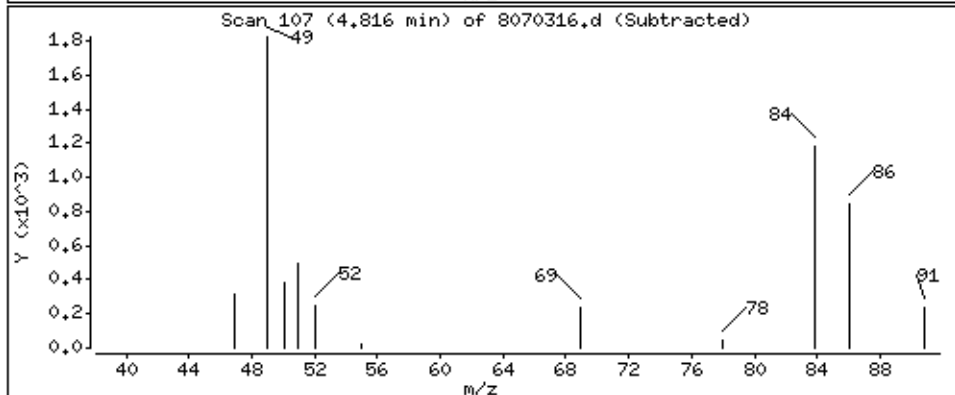
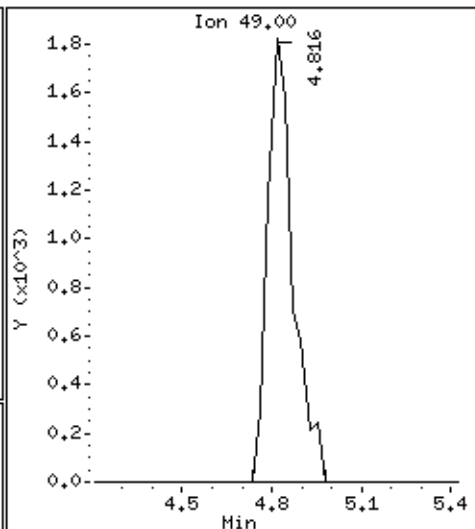
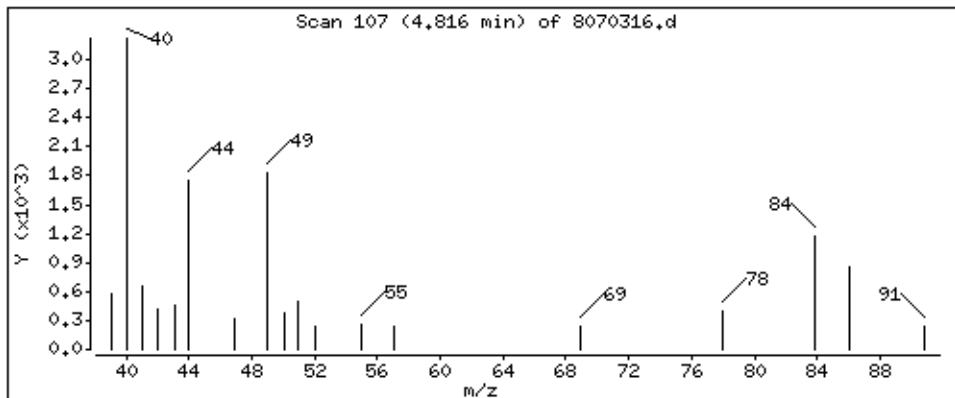
Operator: jdg

Column phase: RTX-624

Column diameter: 0.53

40 Methylene Chloride

Concentration: 1,013 PPBV





AN ENVIRONMENTAL ANALYTICAL LABORATORY

Summary of Detected Compounds MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

Client Sample ID: AMS 5 DW (3746)

Lab ID#: 0706440-02A

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Acetone	3.6	24	8.5	58
2-Butanone (Methyl Ethyl Ketone)	0.90	2.9	2.6	8.6



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 5 DW (3746)

Lab ID#: 0706440-02A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070317	Date of Collection:	6/20/07
Dil. Factor:	1.79	Date of Analysis:	7/3/07 07:24 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Freon 12	0.90	Not Detected	4.4	Not Detected
Freon 114	0.90	Not Detected	6.2	Not Detected
Vinyl Chloride	0.90	Not Detected	2.3	Not Detected
Bromomethane	0.90	Not Detected	3.5	Not Detected
Chloroethane	0.90	Not Detected	2.4	Not Detected
Freon 11	0.90	Not Detected	5.0	Not Detected
1,1-Dichloroethene	0.90	Not Detected	3.5	Not Detected
Freon 113	0.90	Not Detected	6.8	Not Detected
Methylene Chloride	0.90	Not Detected	3.1	Not Detected
1,1-Dichloroethane	0.90	Not Detected	3.6	Not Detected
cis-1,2-Dichloroethene	0.90	Not Detected	3.5	Not Detected
Chloroform	0.90	Not Detected	4.4	Not Detected
1,1,1-Trichloroethane	0.90	Not Detected	4.9	Not Detected
Carbon Tetrachloride	0.90	Not Detected	5.6	Not Detected
Benzene	0.90	Not Detected	2.8	Not Detected
1,2-Dichloroethane	0.90	Not Detected	3.6	Not Detected
Trichloroethene	0.90	Not Detected	4.8	Not Detected
1,2-Dichloropropane	0.90	Not Detected	4.1	Not Detected
cis-1,3-Dichloropropene	0.90	Not Detected	4.1	Not Detected
Toluene	0.90	Not Detected	3.4	Not Detected
trans-1,3-Dichloropropene	0.90	Not Detected	4.1	Not Detected
1,1,2-Trichloroethane	0.90	Not Detected	4.9	Not Detected
Tetrachloroethene	0.90	Not Detected	6.1	Not Detected
1,2-Dibromoethane (EDB)	0.90	Not Detected	6.9	Not Detected
Chlorobenzene	0.90	Not Detected	4.1	Not Detected
Ethyl Benzene	0.90	Not Detected	3.9	Not Detected
m,p-Xylene	0.90	Not Detected	3.9	Not Detected
o-Xylene	0.90	Not Detected	3.9	Not Detected
Styrene	0.90	Not Detected	3.8	Not Detected
1,1,1,2-Tetrachloroethane	0.90	Not Detected	6.1	Not Detected
1,3,5-Trimethylbenzene	0.90	Not Detected	4.4	Not Detected
1,2,4-Trimethylbenzene	0.90	Not Detected	4.4	Not Detected
1,3-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
1,4-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
alpha-Chlorotoluene	0.90	Not Detected	4.6	Not Detected
1,2-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
1,3-Butadiene	0.90	Not Detected	2.0	Not Detected
Hexane	0.90	Not Detected	3.2	Not Detected
Cyclohexane	0.90	Not Detected	3.1	Not Detected



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Client Sample ID: AMS 5 DW (3746)

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MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070317	Date of Collection:	6/20/07
Dil. Factor:	1.79	Date of Analysis:	7/3/07 07:24 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Heptane	0.90	Not Detected	3.7	Not Detected
Bromodichloromethane	0.90	Not Detected	6.0	Not Detected
Dibromochloromethane	0.90	Not Detected	7.6	Not Detected
Cumene	0.90	Not Detected	4.4	Not Detected
Propylbenzene	0.90	Not Detected	4.4	Not Detected
Chloromethane	3.6	Not Detected	7.4	Not Detected
1,2,4-Trichlorobenzene	3.6	Not Detected	26	Not Detected
Hexachlorobutadiene	3.6	Not Detected	38	Not Detected
Acetone	3.6	24	8.5	58
Carbon Disulfide	0.90	Not Detected	2.8	Not Detected
2-Propanol	3.6	Not Detected	8.8	Not Detected
trans-1,2-Dichloroethene	0.90	Not Detected	3.5	Not Detected
2-Butanone (Methyl Ethyl Ketone)	0.90	2.9	2.6	8.6
Tetrahydrofuran	0.90	Not Detected	2.6	Not Detected
1,4-Dioxane	3.6	Not Detected	13	Not Detected
4-Methyl-2-pentanone	0.90	Not Detected	3.7	Not Detected
2-Hexanone	3.6	Not Detected	15	Not Detected
Bromoform	0.90	Not Detected	9.2	Not Detected
4-Ethyltoluene	0.90	Not Detected	4.4	Not Detected
Ethanol	3.6	Not Detected	6.7	Not Detected
Methyl tert-butyl ether	0.90	Not Detected	3.2	Not Detected
3-Chloropropene	3.6	Not Detected	11	Not Detected
2,2,4-Trimethylpentane	0.90	Not Detected	4.2	Not Detected
Naphthalene	3.6	Not Detected	19	Not Detected

Container Type: 6 Liter Summa Canister

Surrogates	%Recovery	Method Limits
Toluene-d8	101	70-130
1,2-Dichloroethane-d4	105	70-130
4-Bromofluorobenzene	104	70-130

Report Date: 05-Jul-2007 09:31

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070317.d
 Lab Smp Id: 0706440-02A
 Inj Date : 03-JUL-2007 19:24
 Operator : jdj Inst ID: msd8.i
 Smp Info : 200ml #3746
 Misc Info : 7.5"Hg-5.0psi
 Comment :
 Method : /var/chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 14:39 dpage Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1
 Dil Factor: 1.79000
 Integrator: HP RTE Compound Sublist: AT04.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS									
ON-COL FINAL									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	(PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	251316	25.0000		70.00- 130.00	100.00	
7.387	7.387	(1.000)	128	198247			47.25- 107.25	78.88	
7.387	7.387	(1.000)	49	378024			115.07- 175.07	150.42	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	1070881	25.0000		70.00- 130.00	100.00	
9.267	9.267	(1.000)	88	160861			0.00- 44.97	15.02	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	801011	25.0000		70.00- 130.00	100.00	
14.576	14.576	(1.000)	82	451272			0.00- 30.00	56.34	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465	(1.146)	65	346721	26.3603	26.360	70.00- 130.00	100.00	
8.465	8.465	(1.146)	67	175077			0.00- 30.00	50.50	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115	(1.307)	98	934771	25.2140	25.214	70.00- 130.00	100.00	
12.115	12.115	(1.307)	70	89814			0.00- 30.00	9.61	

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPEV)	(PPBV)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====

\$ 104 Toluene-d8 (continued)

12.115	12.115	(1.307)	100	602228			0.00- 30.00	64.43
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\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207	16.207	(1.112)	174	493228	25.8798	25.880	70.00- 130.00	100.00
16.207	16.207	(1.112)	95	618390			86.34- 146.34	125.38
16.207	16.207	(1.112)	176	469820			66.11- 126.11	95.25

30 Acetone

CAS #: 67-64-1

4.124	4.124	(0.558)	58	116737	13.5547	24.263	70.00- 130.00	100.00
4.124	4.124	(0.558)	43	342071			0.00- 30.00	293.03

65 2-Butanone

CAS #: 78-93-3

7.027	7.027	(0.951)	72	13608	1.63389	2.925	70.00- 130.00	100.00
7.027	7.027	(0.951)	43	69391			420.80- 480.80	509.93
7.027	7.027	(0.951)	57	5448			0.00- 30.00	40.04

Report Date: 05-Jul-2007 09:31

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARYInstrument ID: msd8.i
Lab File ID: 8070317.d
Lab Smp Id: 0706440-02ACalibration Date: 03-JUL-2007
Calibration Time: 09:41

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdg

Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 7.5"Hg-5.0psi

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	251316	-27.35
88 1,4-Difluorobenze	1512218	907331	2117105	1070881	-29.18
125 Chlorobenzene-d5	1120246	672148	1568344	801011	-28.50

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
Sample Matrix: GAS Fraction: VOA
Lab Smp Id: 0706440-02A
Level: LOW Operator: jdg
Data Type: MS DATA SampleType: SAMPLE
SpikeList File: AT04+NA-2.spk Quant Type: ISTD
Sublist File: AT04.sub
Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m
Misc Info: 7.5"Hg-5.0psi

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	26.360	105.44	70-130
\$ 104 Toluene-d8	25.000	25.214	100.86	70-130
\$ 140 Bromofluorobenzene	25.000	25.880	103.52	70-130

Data File: /chem/msd8.1/8-03jul.b/8070317.d

Date: 03-JUL-2007 19:24

Client ID:

Sample Info: 200ml #3746

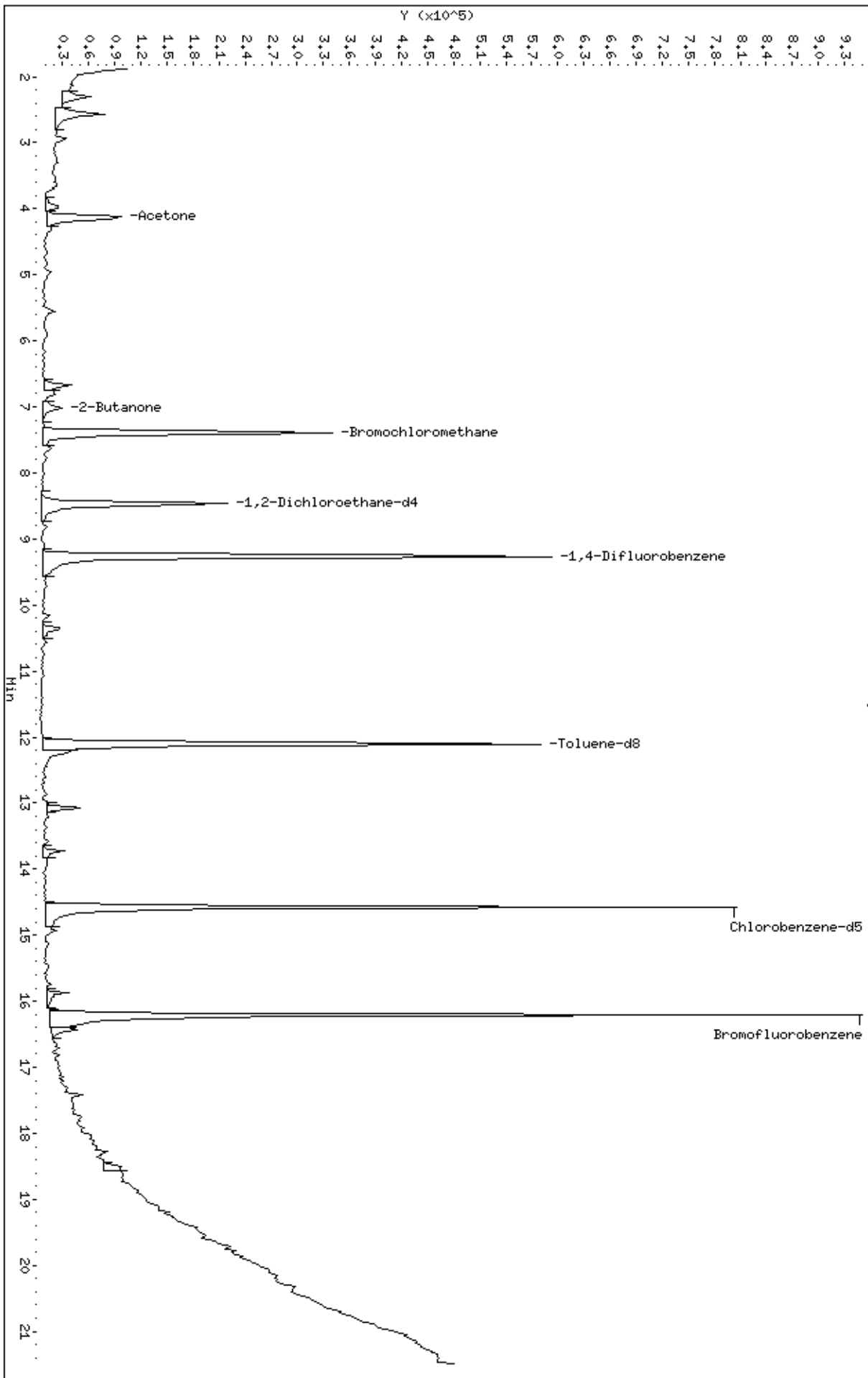
Column phase: RTX-624

Instrument: msd8.1

Operator: jdg

Column diameter: 0.53

/chem/msd8.1/8-03jul.b/8070317.d



Date : 03-JUL-2007 19:24

Client ID:

Instrument: msd8.i

Sample Info: 200ml #3746

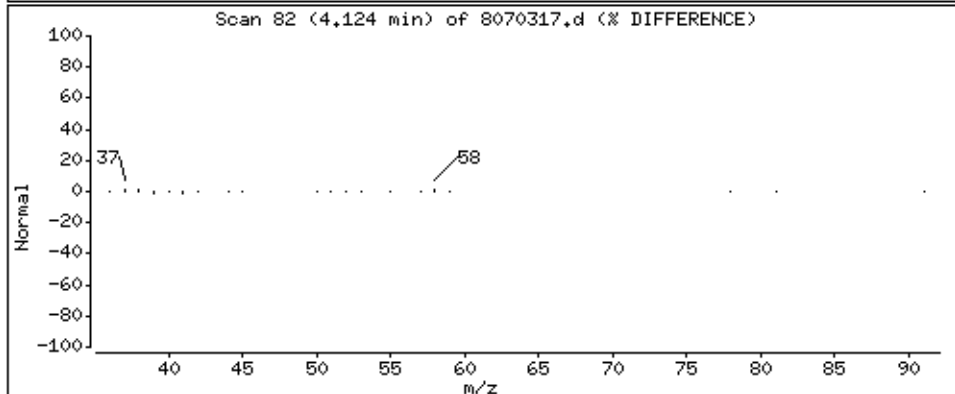
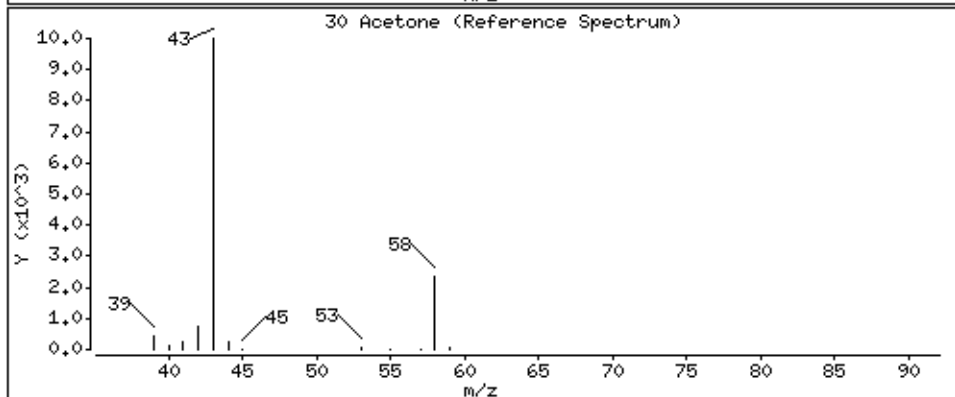
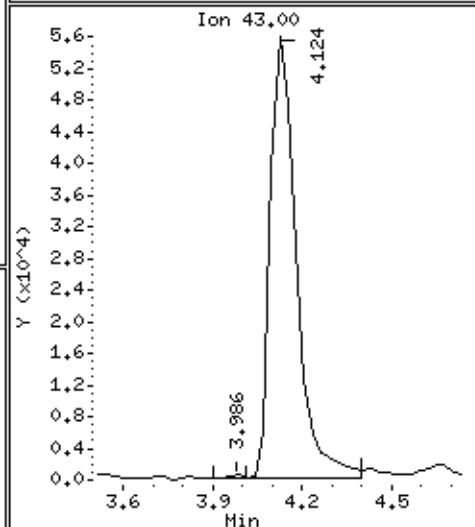
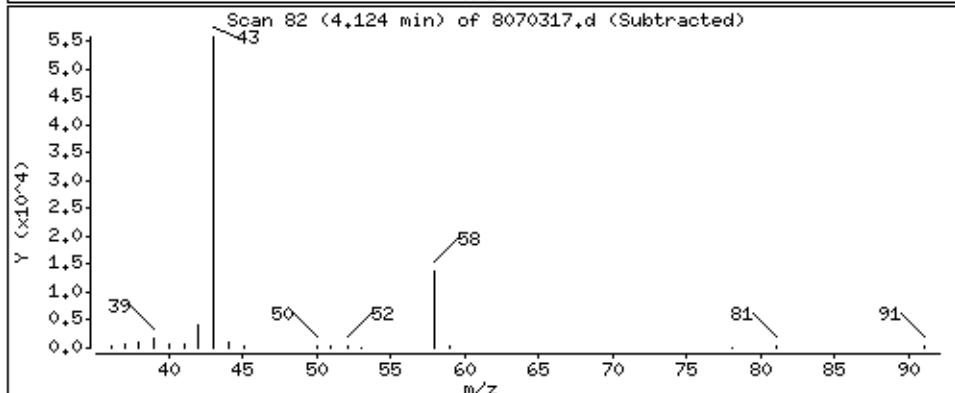
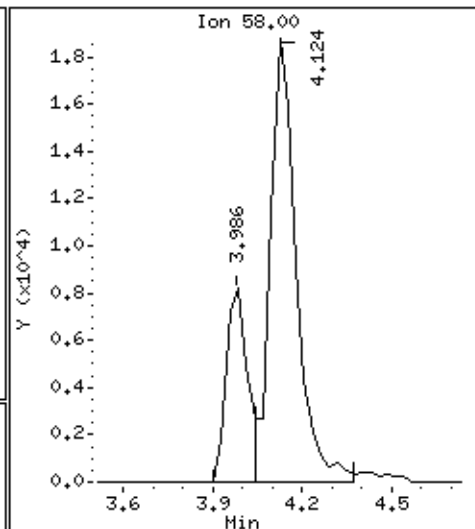
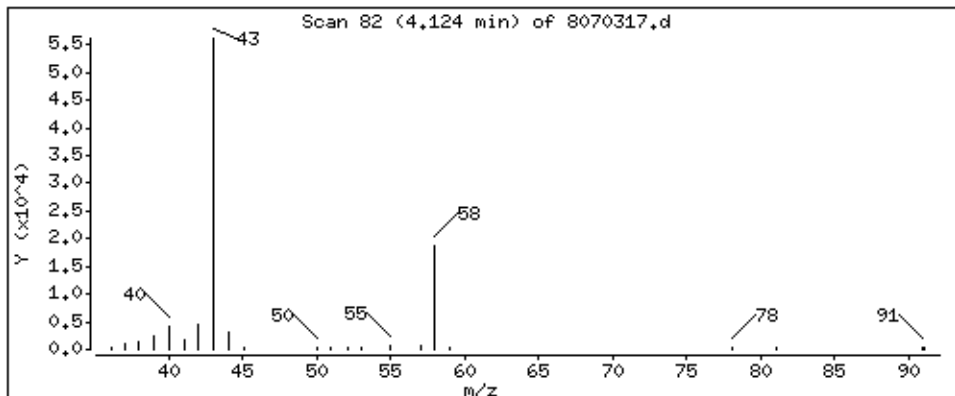
Operator: jdg

Column phase: RTX-624

Column diameter: 0.53

30 Acetone

Concentration: 24,263 PPBV



Date : 03-JUL-2007 19:24

Client ID:

Instrument: msd8.i

Sample Info: 200ml #3746

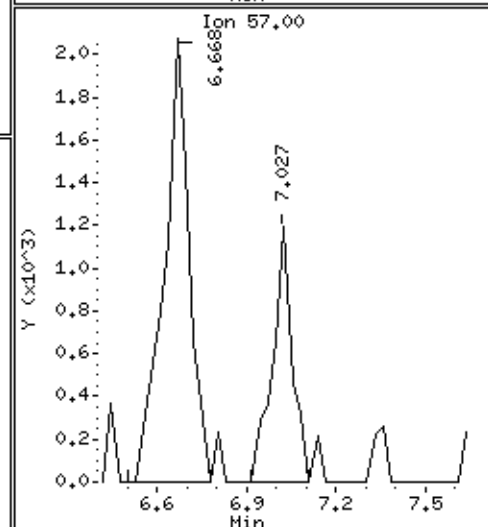
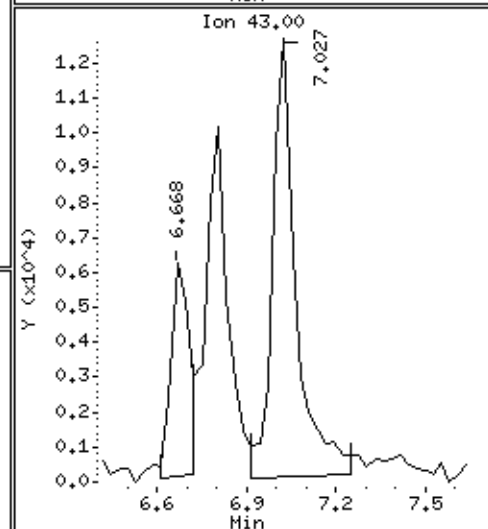
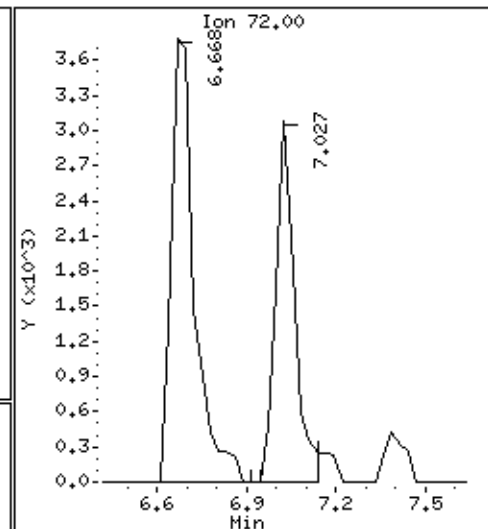
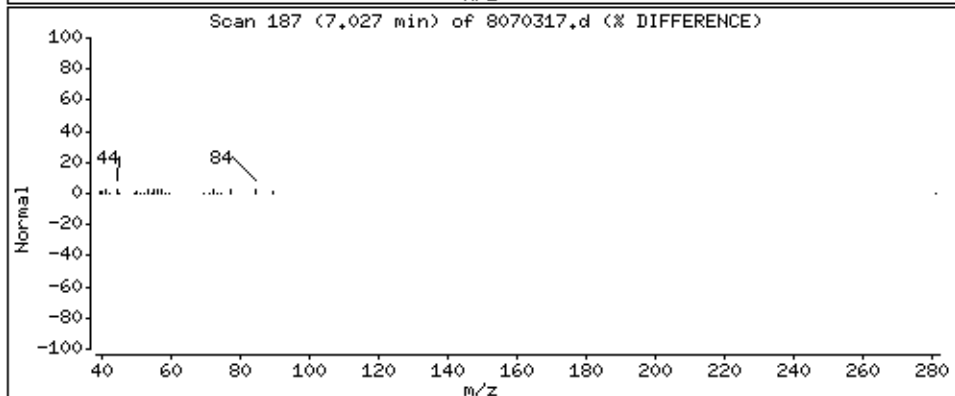
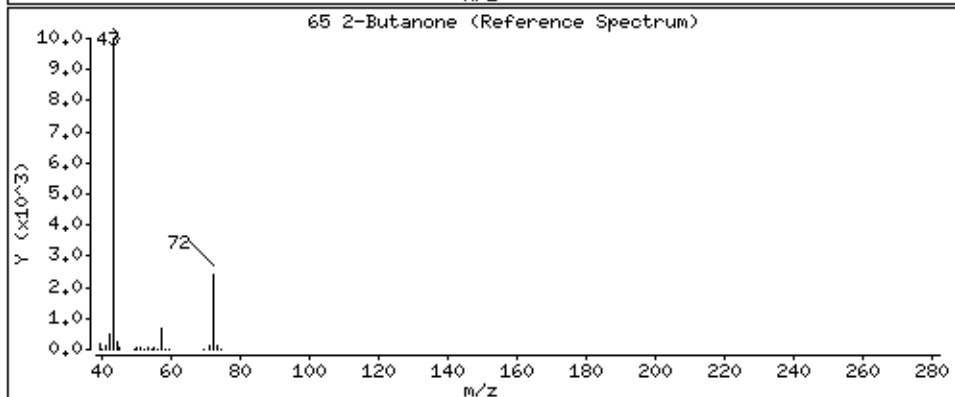
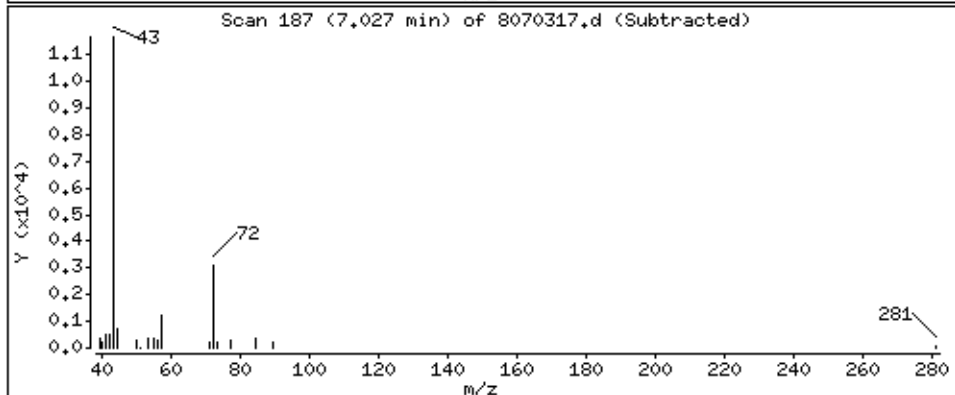
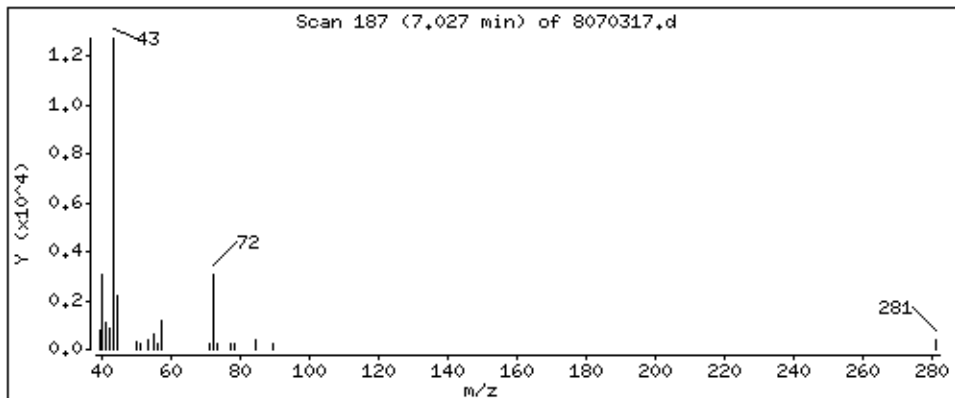
Operator: jdg

Column phase: RTx-624

Column diameter: 0.53

65 2-Butanone

Concentration: 2,925 PPBV





AN ENVIRONMENTAL ANALYTICAL LABORATORY

Summary of Detected Compounds MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

Client Sample ID: AMS 5 DW (3746) Lab Duplicate

Lab ID#: 0706440-02AA

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Acetone	3.6	23	8.5	54
2-Butanone (Methyl Ethyl Ketone)	0.90	2.7	2.6	8.0



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 5 DW (3746) Lab Duplicate

Lab ID#: 0706440-02AA

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070318	Date of Collection:	6/20/07
Dil. Factor:	1.79	Date of Analysis:	7/3/07 08:06 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Freon 12	0.90	Not Detected	4.4	Not Detected
Freon 114	0.90	Not Detected	6.2	Not Detected
Vinyl Chloride	0.90	Not Detected	2.3	Not Detected
Bromomethane	0.90	Not Detected	3.5	Not Detected
Chloroethane	0.90	Not Detected	2.4	Not Detected
Freon 11	0.90	Not Detected	5.0	Not Detected
1,1-Dichloroethene	0.90	Not Detected	3.5	Not Detected
Freon 113	0.90	Not Detected	6.8	Not Detected
Methylene Chloride	0.90	Not Detected	3.1	Not Detected
1,1-Dichloroethane	0.90	Not Detected	3.6	Not Detected
cis-1,2-Dichloroethene	0.90	Not Detected	3.5	Not Detected
Chloroform	0.90	Not Detected	4.4	Not Detected
1,1,1-Trichloroethane	0.90	Not Detected	4.9	Not Detected
Carbon Tetrachloride	0.90	Not Detected	5.6	Not Detected
Benzene	0.90	Not Detected	2.8	Not Detected
1,2-Dichloroethane	0.90	Not Detected	3.6	Not Detected
Trichloroethene	0.90	Not Detected	4.8	Not Detected
1,2-Dichloropropane	0.90	Not Detected	4.1	Not Detected
cis-1,3-Dichloropropene	0.90	Not Detected	4.1	Not Detected
Toluene	0.90	Not Detected	3.4	Not Detected
trans-1,3-Dichloropropene	0.90	Not Detected	4.1	Not Detected
1,1,2-Trichloroethane	0.90	Not Detected	4.9	Not Detected
Tetrachloroethene	0.90	Not Detected	6.1	Not Detected
1,2-Dibromoethane (EDB)	0.90	Not Detected	6.9	Not Detected
Chlorobenzene	0.90	Not Detected	4.1	Not Detected
Ethyl Benzene	0.90	Not Detected	3.9	Not Detected
m,p-Xylene	0.90	Not Detected	3.9	Not Detected
o-Xylene	0.90	Not Detected	3.9	Not Detected
Styrene	0.90	Not Detected	3.8	Not Detected
1,1,2,2-Tetrachloroethane	0.90	Not Detected	6.1	Not Detected
1,3,5-Trimethylbenzene	0.90	Not Detected	4.4	Not Detected
1,2,4-Trimethylbenzene	0.90	Not Detected	4.4	Not Detected
1,3-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
1,4-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
alpha-Chlorotoluene	0.90	Not Detected	4.6	Not Detected
1,2-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
1,3-Butadiene	0.90	Not Detected	2.0	Not Detected
Hexane	0.90	Not Detected	3.2	Not Detected
Cyclohexane	0.90	Not Detected	3.1	Not Detected



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 5 DW (3746) Lab Duplicate

Lab ID#: 0706440-02AA

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070318	Date of Collection: 6/20/07
Dil. Factor:	1.79	Date of Analysis: 7/3/07 08:06 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Heptane	0.90	Not Detected	3.7	Not Detected
Bromodichloromethane	0.90	Not Detected	6.0	Not Detected
Dibromochloromethane	0.90	Not Detected	7.6	Not Detected
Cumene	0.90	Not Detected	4.4	Not Detected
Propylbenzene	0.90	Not Detected	4.4	Not Detected
Chloromethane	3.6	Not Detected	7.4	Not Detected
1,2,4-Trichlorobenzene	3.6	Not Detected	26	Not Detected
Hexachlorobutadiene	3.6	Not Detected	38	Not Detected
Acetone	3.6	23	8.5	54
Carbon Disulfide	0.90	Not Detected	2.8	Not Detected
2-Propanol	3.6	Not Detected	8.8	Not Detected
trans-1,2-Dichloroethene	0.90	Not Detected	3.5	Not Detected
2-Butanone (Methyl Ethyl Ketone)	0.90	2.7	2.6	8.0
Tetrahydrofuran	0.90	Not Detected	2.6	Not Detected
1,4-Dioxane	3.6	Not Detected	13	Not Detected
4-Methyl-2-pentanone	0.90	Not Detected	3.7	Not Detected
2-Hexanone	3.6	Not Detected	15	Not Detected
Bromoform	0.90	Not Detected	9.2	Not Detected
4-Ethyltoluene	0.90	Not Detected	4.4	Not Detected
Ethanol	3.6	Not Detected	6.7	Not Detected
Methyl tert-butyl ether	0.90	Not Detected	3.2	Not Detected
3-Chloropropene	3.6	Not Detected	11	Not Detected
2,2,4-Trimethylpentane	0.90	Not Detected	4.2	Not Detected
Naphthalene	3.6	Not Detected	19	Not Detected

Container Type: 6 Liter Summa Canister

Surrogates	%Recovery	Method Limits
Toluene-d8	98	70-130
1,2-Dichloroethane-d4	106	70-130
4-Bromofluorobenzene	102	70-130

Report Date: 05-Jul-2007 09:31

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070318.d
 Lab Smp Id: 0706440-02AA
 Inj Date : 03-JUL-2007 20:06
 Operator : jdg Inst ID: msd8.i
 Smp Info : 200ml #3746
 Misc Info : 7.5"Hg-5.0psi
 Comment :
 Method : /var/chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 14:39 dpage Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1
 Dil Factor: 1.79000
 Integrator: HP RTE Compound Sublist: AT04.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS								
		ON-COL		FINAL		TARGET RANGE		RATIO
RT	EXP RT (REL RT)	MASS	RESPONSE (PPBV)	(PPBV)				
==	=====	=====	=====	=====	=====	=====	=====	=====

* 68	Bromochloromethane					CAS #: 74-97-5		
7.387	7.387 (1.000)	130	254121	25.0000		70.00- 130.00	100.00	
7.387	7.387 (1.000)	128	198009			47.25- 107.25	77.92	
7.387	7.387 (1.000)	49	386483			115.07- 175.07	152.09	

* 88	1,4-Difluorobenzene					CAS #: 540-36-3		
9.267	9.267 (1.000)	114	1069480	25.0000		70.00- 130.00	100.00	
9.267	9.267 (1.000)	88	162764			0.00- 44.97	15.22	

* 125	Chlorobenzene-d5					CAS #: 3114-55-4		
14.576	14.576 (1.000)	117	804503	25.0000		70.00- 130.00	100.00	
14.576	14.576 (1.000)	82	450306			0.00- 30.00	55.97	

\$ 82	1,2-Dichloroethane-d4					CAS #: 17060-07-0		
8.465	8.465 (1.146)	65	350782	26.3747	26.375	70.00- 130.00	100.00	
8.465	8.465 (1.146)	67	179976			0.00- 30.00	51.31	

\$ 104	Toluene-d8					CAS #: 2037-26-5		
12.115	12.115 (1.307)	98	904394	24.4265	24.426	70.00- 130.00	100.00	
12.115	12.115 (1.307)	70	92862			0.00- 30.00	10.27	

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPEV)	(PPBV)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====

\$ 104 Toluene-d8 (continued)

12.115	12.115	(1.307)	100	588074			0.00- 30.00	65.02
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\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207	16.207	(1.112)	174	486422	25.4119	25.412	70.00- 130.00	100.00
16.207	16.207	(1.112)	95	615255			86.34- 146.34	126.49
16.207	16.207	(1.112)	176	479866			66.11- 126.11	98.65

30 Acetone

CAS #: 67-64-1

4.124	4.124	(0.558)	58	111458	12.7989	22.910	70.00- 130.00	100.00
4.124	4.124	(0.558)	43	346958			0.00- 30.00	311.29

65 2-Butanone

CAS #: 78-93-3

7.027	7.027	(0.951)	72	12858	1.52680	2.733	70.00- 130.00	100.00
7.027	7.027	(0.951)	43	60618			420.80- 480.80	471.44
7.027	7.027	(0.951)	57	6521			0.00- 30.00	50.72

Report Date: 05-Jul-2007 09:31

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 03-JUL-2007

Lab File ID: 8070318.d

Calibration Time: 09:41

Lab Smp Id: 0706440-02AA

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdg

Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 7.5"Hg-5.0psi

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	254121	-26.54
88 1,4-Difluorobenze	1512218	907331	2117105	1069480	-29.28
125 Chlorobenzene-d5	1120246	672148	1568344	804503	-28.19

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
Sample Matrix: GAS Fraction: VOA
Lab Smp Id: 0706440-02AA
Level: LOW Operator: jdg
Data Type: MS DATA SampleType: SAMPLE
SpikeList File: AT04+NA-2.spk Quant Type: ISTD
Sublist File: AT04.sub
Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m
Misc Info: 7.5"Hg-5.0psi

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	26.375	105.50	70-130
\$ 104 Toluene-d8	25.000	24.426	97.71	70-130
\$ 140 Bromofluorobenzene	25.000	25.412	101.65	70-130

Data File: /chem/msd8.1/8-03jul.b/8070318.d

Date: 03-JUL-2007 20:06

Client ID:

Sample Info: 200ml #3746

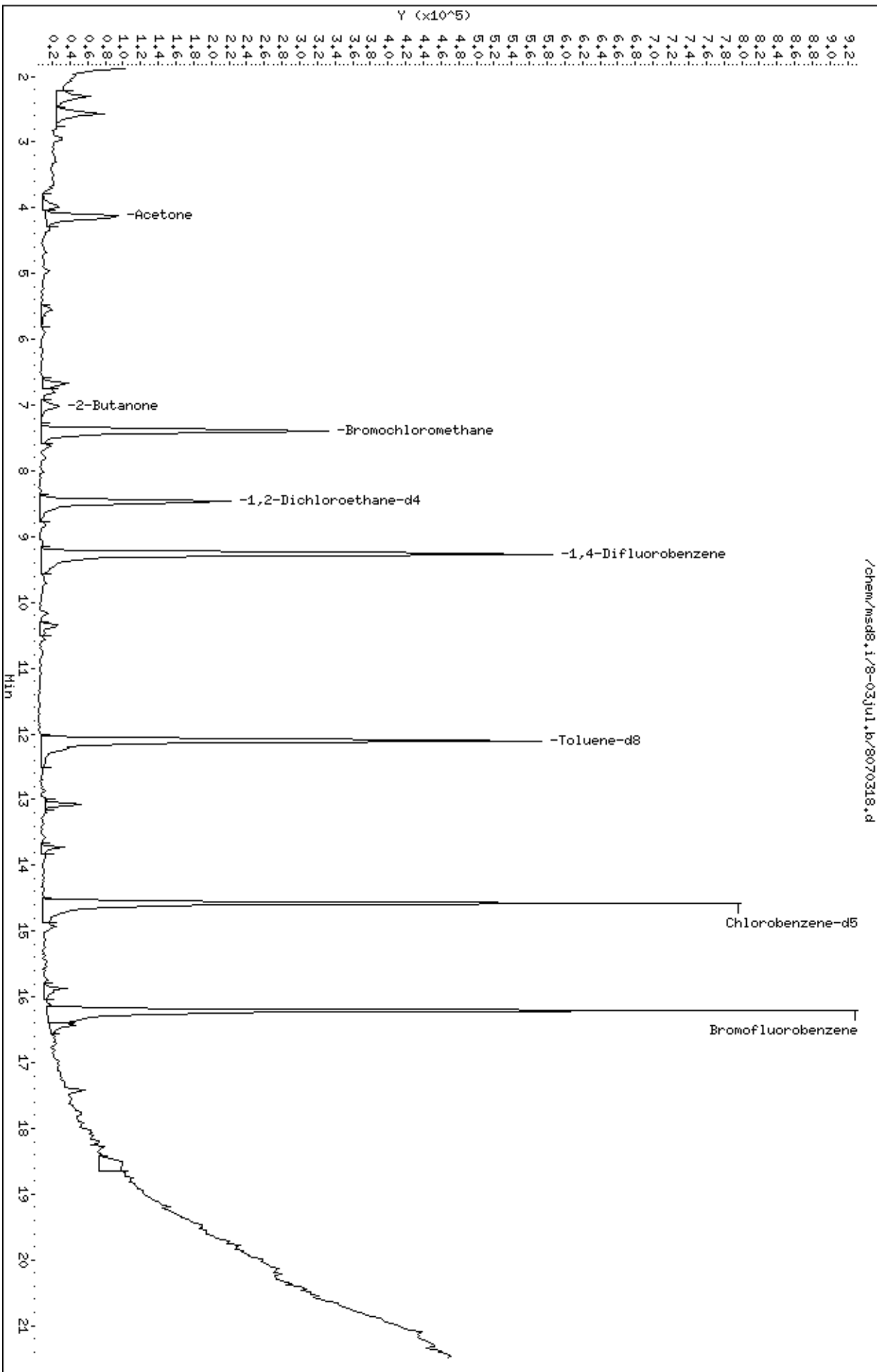
Column phase: RTX-624

Instrument: msd8.1

Operator: jdg

Column diameter: 0.53

/chem/msd8.1/8-03jul.b/8070318.d



Date : 03-JUL-2007 20:06

Client ID:

Instrument: msd8.i

Sample Info: 200ml #3746

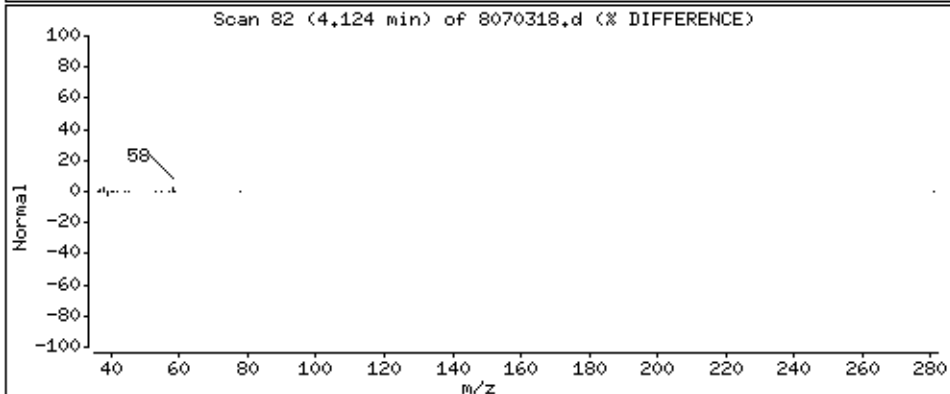
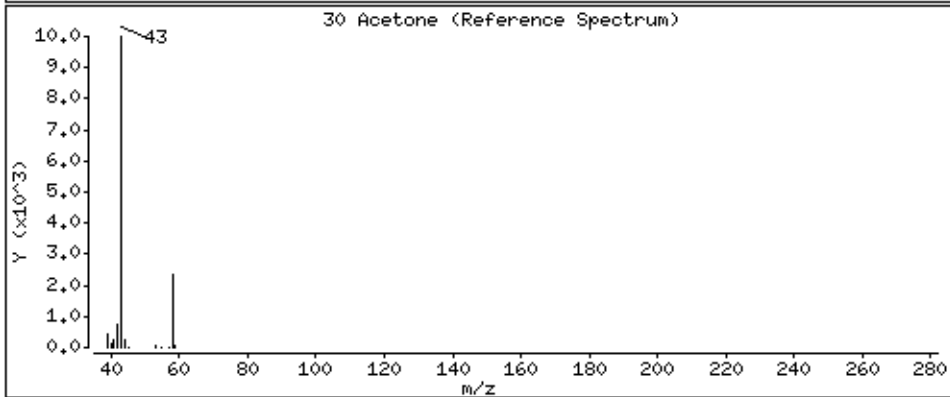
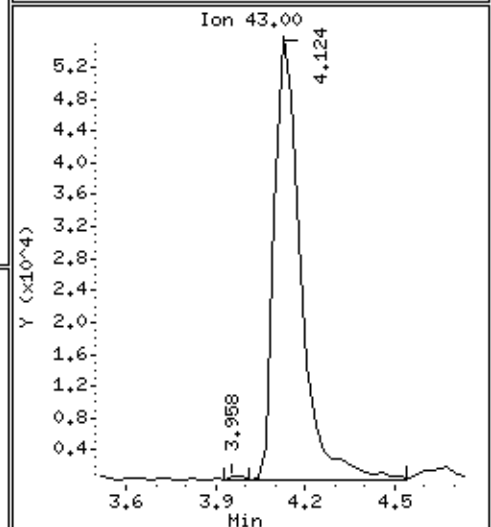
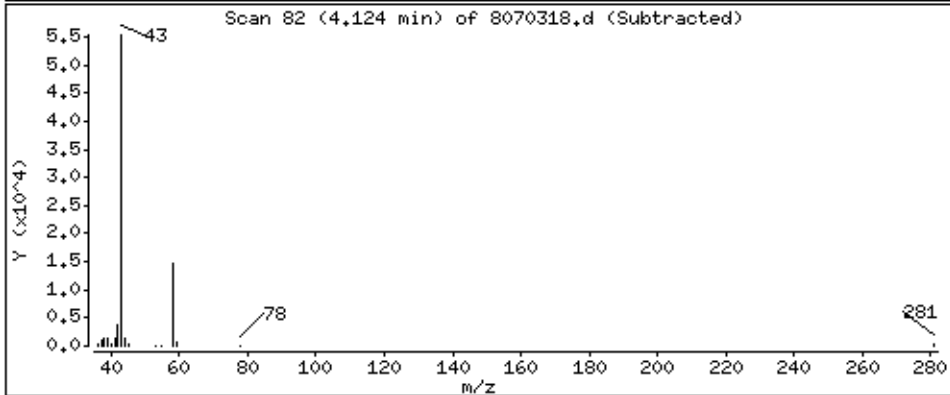
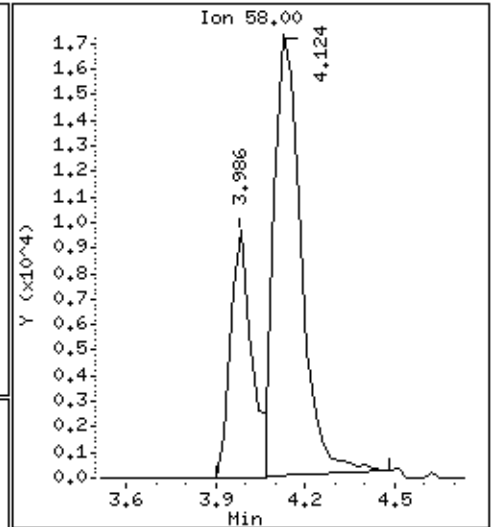
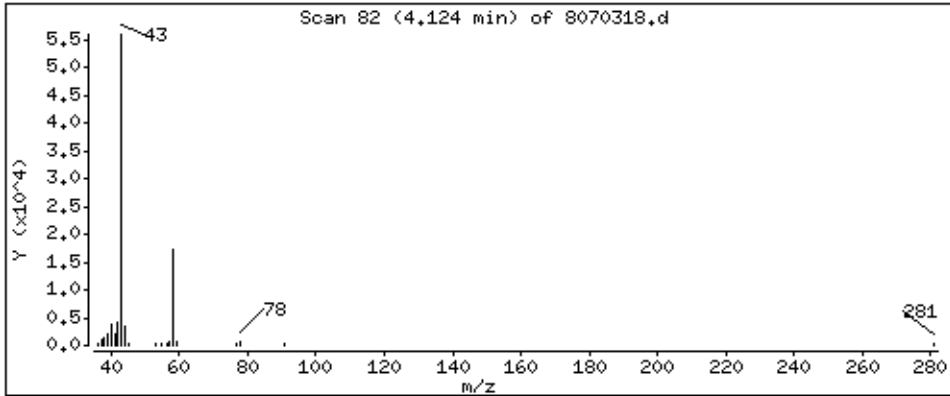
Operator: jdg

Column phase: RTX-624

Column diameter: 0.53

30 Acetone

Concentration: 22,910 PPBV



Date : 03-JUL-2007 20:06

Client ID:

Instrument: msd8.i

Sample Info: 200ml #3746

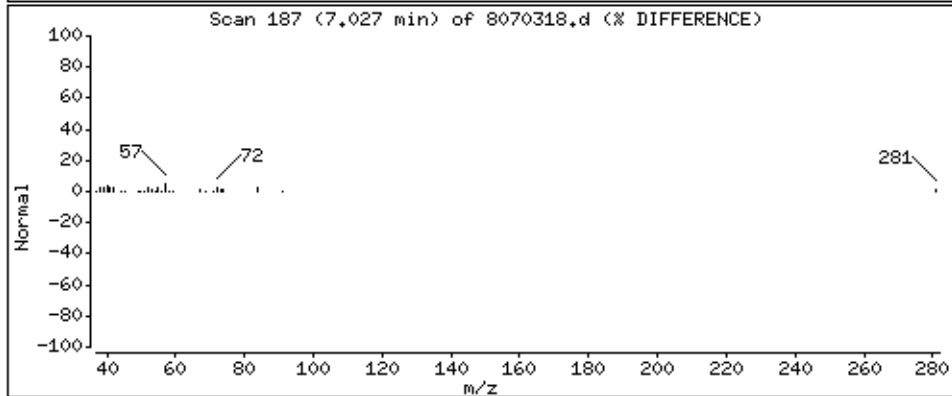
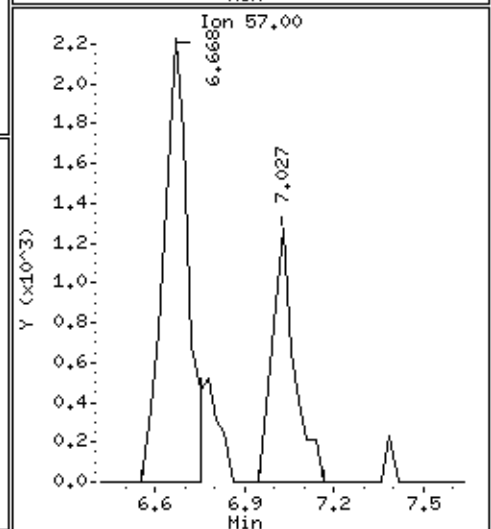
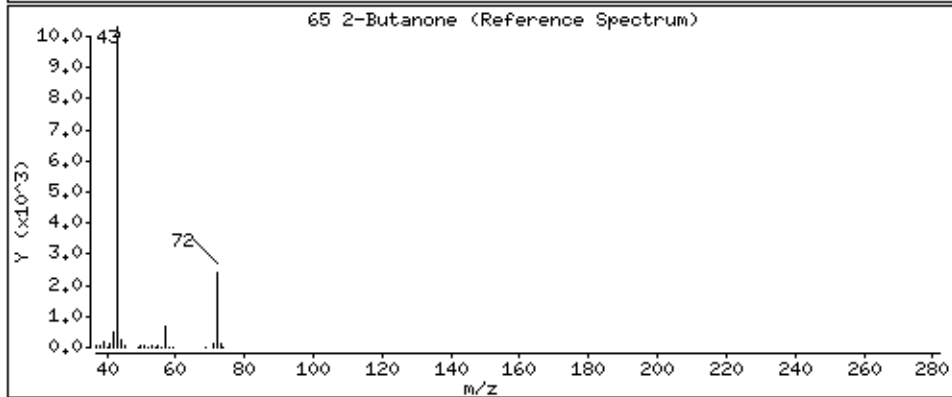
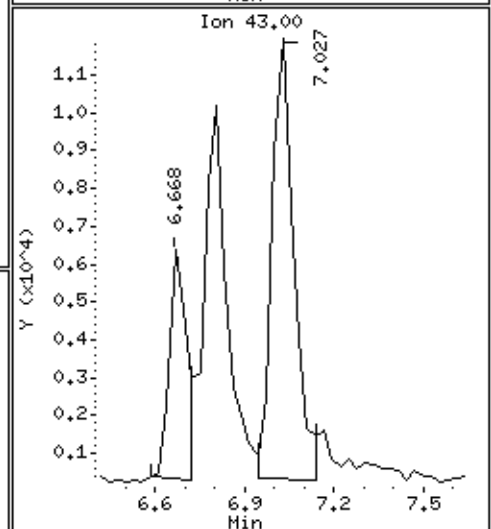
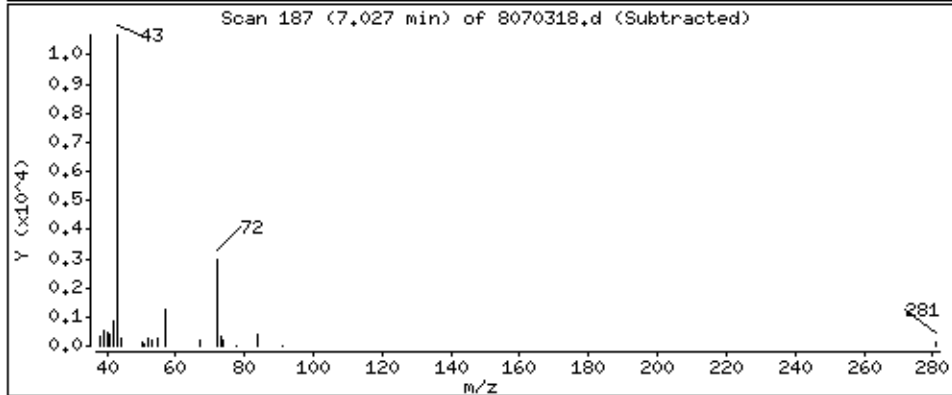
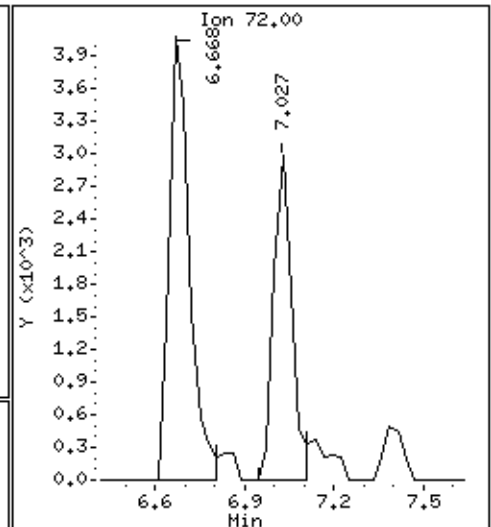
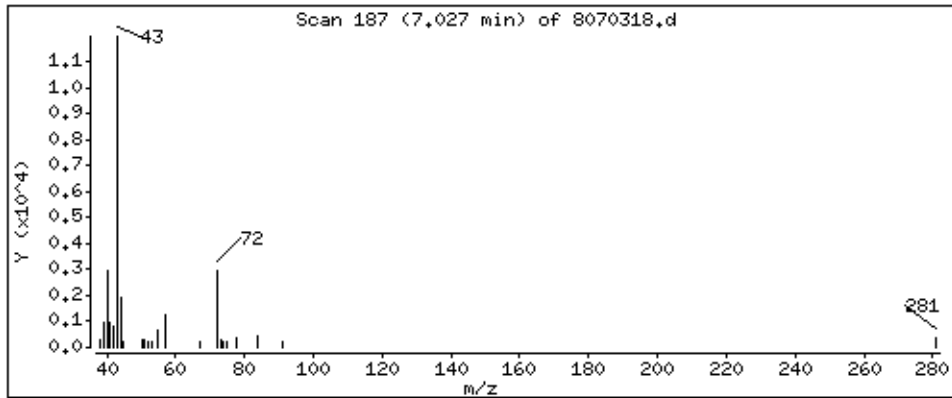
Operator: jdg

Column phase: RTx-624

Column diameter: 0.53

65 2-Butanone

Concentration: 2,733 PPBV





AN ENVIRONMENTAL ANALYTICAL LABORATORY

Summary of Detected Compounds
MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

Client Sample ID: AMS 2 UW

Lab ID#: 0706440-03A

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Toluene	0.90	0.89 J	3.4	3.4
Acetone	3.6	10	8.5	24



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 2 UW

Lab ID#: 0706440-03A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070319	Date of Collection:	6/20/07
Dil. Factor:	1.79	Date of Analysis:	7/3/07 08:49 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Freon 12	0.90	Not Detected	4.4	Not Detected
Freon 114	0.90	Not Detected	6.2	Not Detected
Vinyl Chloride	0.90	Not Detected	2.3	Not Detected
Bromomethane	0.90	Not Detected	3.5	Not Detected
Chloroethane	0.90	Not Detected	2.4	Not Detected
Freon 11	0.90	Not Detected	5.0	Not Detected
1,1-Dichloroethene	0.90	Not Detected	3.5	Not Detected
Freon 113	0.90	Not Detected	6.8	Not Detected
Methylene Chloride	0.90	Not Detected	3.1	Not Detected
1,1-Dichloroethane	0.90	Not Detected	3.6	Not Detected
cis-1,2-Dichloroethene	0.90	Not Detected	3.5	Not Detected
Chloroform	0.90	Not Detected	4.4	Not Detected
1,1,1-Trichloroethane	0.90	Not Detected	4.9	Not Detected
Carbon Tetrachloride	0.90	Not Detected	5.6	Not Detected
Benzene	0.90	Not Detected	2.8	Not Detected
1,2-Dichloroethane	0.90	Not Detected	3.6	Not Detected
Trichloroethene	0.90	Not Detected	4.8	Not Detected
1,2-Dichloropropane	0.90	Not Detected	4.1	Not Detected
cis-1,3-Dichloropropene	0.90	Not Detected	4.1	Not Detected
Toluene	0.90	0.89 J	3.4	3.4
trans-1,3-Dichloropropene	0.90	Not Detected	4.1	Not Detected
1,1,2-Trichloroethane	0.90	Not Detected	4.9	Not Detected
Tetrachloroethene	0.90	Not Detected	6.1	Not Detected
1,2-Dibromoethane (EDB)	0.90	Not Detected	6.9	Not Detected
Chlorobenzene	0.90	Not Detected	4.1	Not Detected
Ethyl Benzene	0.90	Not Detected	3.9	Not Detected
m,p-Xylene	0.90	Not Detected	3.9	Not Detected
o-Xylene	0.90	Not Detected	3.9	Not Detected
Styrene	0.90	Not Detected	3.8	Not Detected
1,1,2,2-Tetrachloroethane	0.90	Not Detected	6.1	Not Detected
1,3,5-Trimethylbenzene	0.90	Not Detected	4.4	Not Detected
1,2,4-Trimethylbenzene	0.90	Not Detected	4.4	Not Detected
1,3-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
1,4-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
alpha-Chlorotoluene	0.90	Not Detected	4.6	Not Detected
1,2-Dichlorobenzene	0.90	Not Detected	5.4	Not Detected
1,3-Butadiene	0.90	Not Detected	2.0	Not Detected
Hexane	0.90	Not Detected	3.2	Not Detected
Cyclohexane	0.90	Not Detected	3.1	Not Detected



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: AMS 2 UW

Lab ID#: 0706440-03A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070319	Date of Collection:	6/20/07
Dil. Factor:	1.79	Date of Analysis:	7/3/07 08:49 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Heptane	0.90	Not Detected	3.7	Not Detected
Bromodichloromethane	0.90	Not Detected	6.0	Not Detected
Dibromochloromethane	0.90	Not Detected	7.6	Not Detected
Cumene	0.90	Not Detected	4.4	Not Detected
Propylbenzene	0.90	Not Detected	4.4	Not Detected
Chloromethane	3.6	Not Detected	7.4	Not Detected
1,2,4-Trichlorobenzene	3.6	Not Detected	26	Not Detected
Hexachlorobutadiene	3.6	Not Detected	38	Not Detected
Acetone	3.6	10	8.5	24
Carbon Disulfide	0.90	Not Detected	2.8	Not Detected
2-Propanol	3.6	Not Detected	8.8	Not Detected
trans-1,2-Dichloroethene	0.90	Not Detected	3.5	Not Detected
2-Butanone (Methyl Ethyl Ketone)	0.90	Not Detected	2.6	Not Detected
Tetrahydrofuran	0.90	Not Detected	2.6	Not Detected
1,4-Dioxane	3.6	Not Detected	13	Not Detected
4-Methyl-2-pentanone	0.90	Not Detected	3.7	Not Detected
2-Hexanone	3.6	Not Detected	15	Not Detected
Bromoform	0.90	Not Detected	9.2	Not Detected
4-Ethyltoluene	0.90	Not Detected	4.4	Not Detected
Ethanol	3.6	Not Detected	6.7	Not Detected
Methyl tert-butyl ether	0.90	Not Detected	3.2	Not Detected
3-Chloropropene	3.6	Not Detected	11	Not Detected
2,2,4-Trimethylpentane	0.90	Not Detected	4.2	Not Detected
Naphthalene	3.6	Not Detected	19	Not Detected

J = Estimated value.

Container Type: 6 Liter Summa Canister

Surrogates	%Recovery	Method Limits
Toluene-d8	98	70-130
1,2-Dichloroethane-d4	102	70-130
4-Bromofluorobenzene	99	70-130

Report Date: 05-Jul-2007 09:32

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070319.d
 Lab Smp Id: 0706440-03A
 Inj Date : 03-JUL-2007 20:49
 Operator : jdg Inst ID: msd8.i
 Smp Info : 200ml #34717
 Misc Info : 7.5"Hg-5.0psi
 Comment :
 Method : /var/chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 14:39 dpage Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1
 Dil Factor: 1.79000
 Integrator: HP RTE Compound Sublist: AT04.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS									
		ON-COL		FINAL		TARGET RANGE		RATIO	
RT	EXP RT (REL RT)	MASS	RESPONSE (PPBV)	(PPBV)					
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387 (1.000)	130	260882	25.0000		70.00-	130.00	100.00	
7.387	7.387 (1.000)	128	199512			47.25-	107.25	76.48	
7.387	7.387 (1.000)	49	389048			115.07-	175.07	149.13	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267 (1.000)	114	1079703	25.0000		70.00-	130.00	100.00	
9.267	9.267 (1.000)	88	159317			0.00-	44.97	14.76	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576 (1.000)	117	796366	25.0000		70.00-	130.00	100.00	
14.576	14.576 (1.000)	82	452940			0.00-	30.00	56.88	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465 (1.146)	65	347272	25.4341	25.434	70.00-	130.00	100.00	
8.465	8.465 (1.146)	67	181716			0.00-	30.00	52.33	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115 (1.307)	98	911136	24.3756	24.376	70.00-	130.00	100.00	
12.115	12.115 (1.307)	70	95643			0.00-	30.00	10.50	

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPEV)	(PPBV)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====

\$ 104 Toluene-d8 (continued)

12.115	12.115	(1.307)	100	579900			0.00- 30.00	63.65
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\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207	16.207	(1.112)	174	467761	24.6867	24.687	70.00- 130.00	100.00
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16.207	16.207	(1.112)	95	610407			86.34- 146.34	130.50
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16.207	16.207	(1.112)	176	471424			66.11- 126.11	100.78
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30 Acetone

CAS #: 67-64-1

4.124	4.124	(0.558)	58	50725	5.67388	10.156	70.00- 130.00	100.00
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4.124	4.124	(0.558)	43	132253			0.00- 30.00	260.73
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105 Toluene

CAS #: 108-88-3

12.225	12.225	(1.319)	91	24762	0.49951	0.8941	70.00- 130.00	100.00(a)
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12.253	12.225	(1.322)	92	12923			30.07- 90.07	52.19
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QC Flag Legend

a - Target compound detected but, quantitated amount
Below Limit Of Quantitation(BLOQ).

Report Date: 05-Jul-2007 09:32

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARYInstrument ID: msd8.i
Lab File ID: 8070319.d
Lab Smp Id: 0706440-03ACalibration Date: 03-JUL-2007
Calibration Time: 09:41

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdg

Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 7.5"Hg-5.0psi

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	260882	-24.59
88 1,4-Difluorobenze	1512218	907331	2117105	1079703	-28.60
125 Chlorobenzene-d5	1120246	672148	1568344	796366	-28.91

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
Sample Matrix: GAS Fraction: VOA
Lab Smp Id: 0706440-03A
Level: LOW Operator: jdg
Data Type: MS DATA SampleType: SAMPLE
SpikeList File: AT04+NA-2.spk Quant Type: ISTD
Sublist File: AT04.sub
Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m
Misc Info: 7.5"Hg-5.0psi

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	25.434	101.74	70-130
\$ 104 Toluene-d8	25.000	24.376	97.50	70-130
\$ 140 Bromofluorobenzene	25.000	24.687	98.75	70-130

Data File: /chem/msd8.1/8-03jul.b/8070319.d

Date: 03-JUL-2007 20:49

Client ID:

Sample Info: 200ml #34717

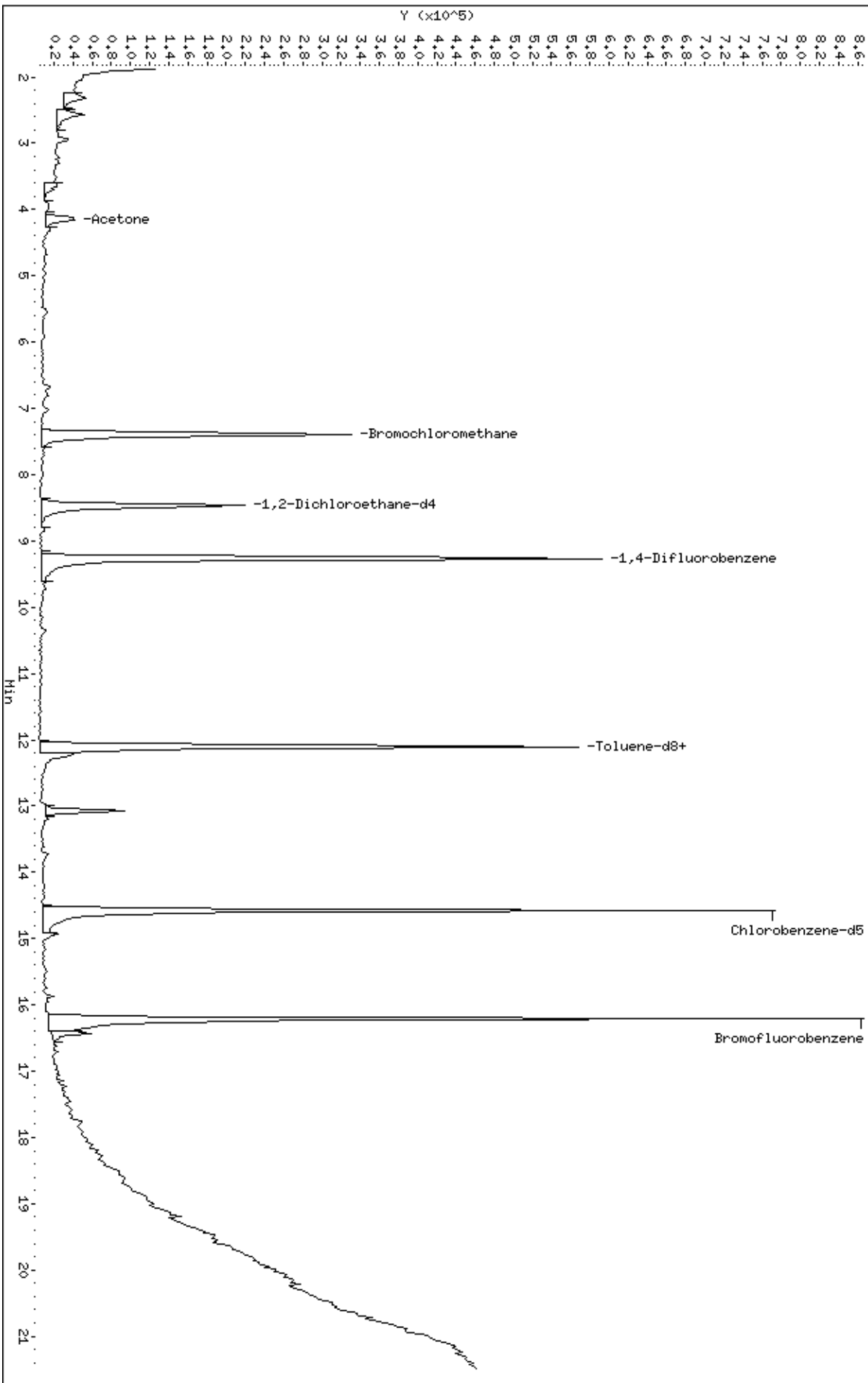
Column phase: RTX-624

Instrument: msd8.1

Operator: jdg

Column diameter: 0.53

/chem/msd8.1/8-03jul.b/8070319.d



Date : 03-JUL-2007 20:49

Client ID:

Instrument: msd8.i

Sample Info: 200ml #34717

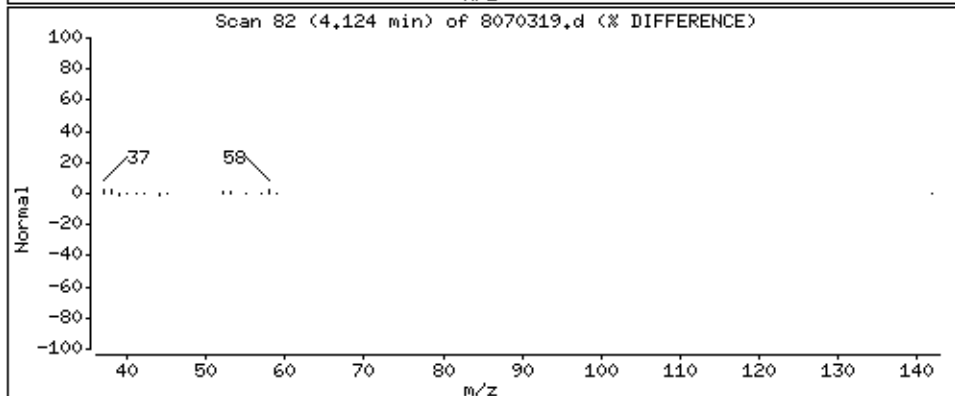
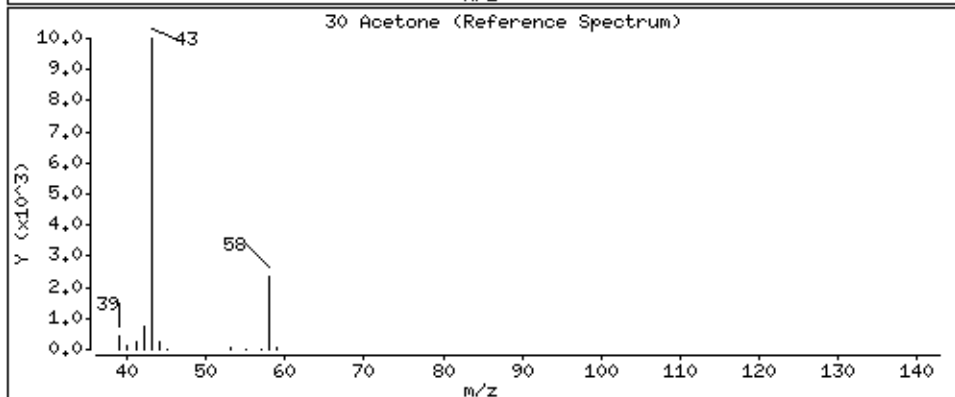
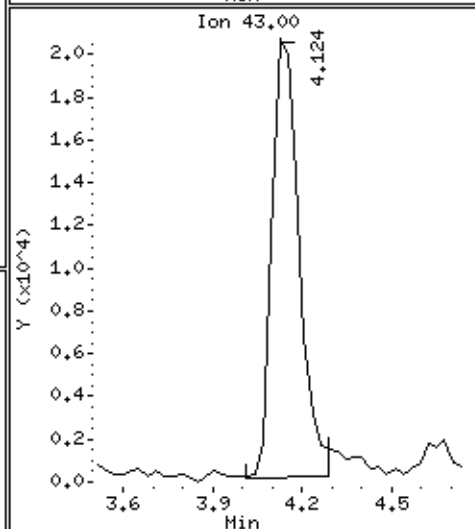
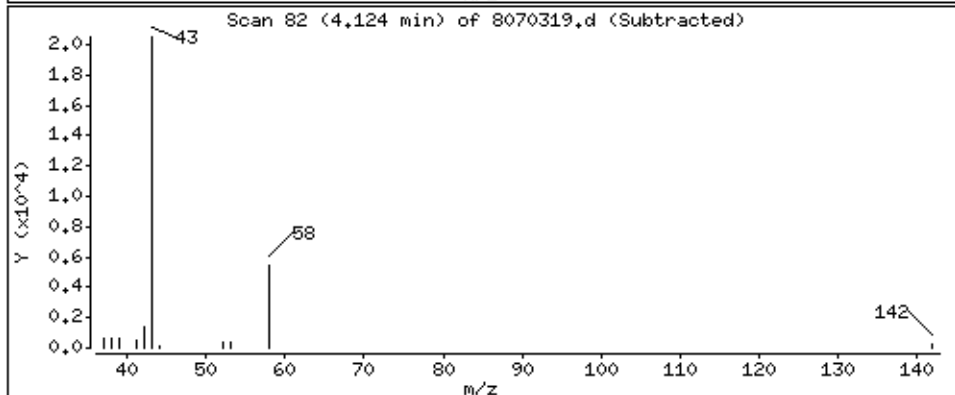
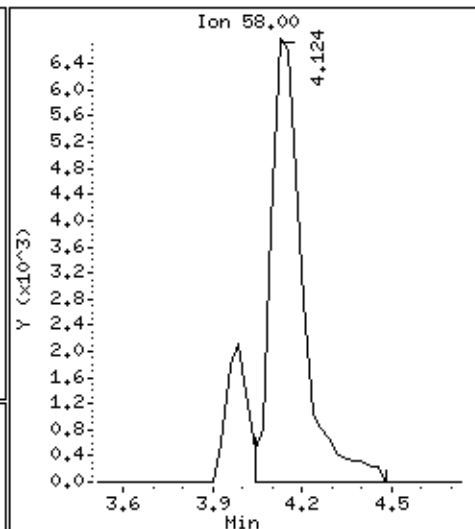
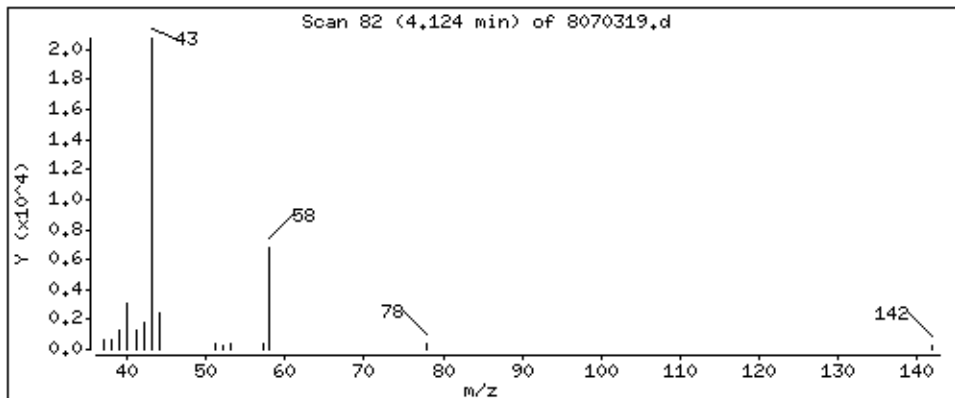
Operator: jdg

Column phase: RTX-624

Column diameter: 0.53

30 Acetone

Concentration: 10,156 PPBV



Date : 03-JUL-2007 20:49

Client ID:

Instrument: msd8.i

Sample Info: 200ml #34717

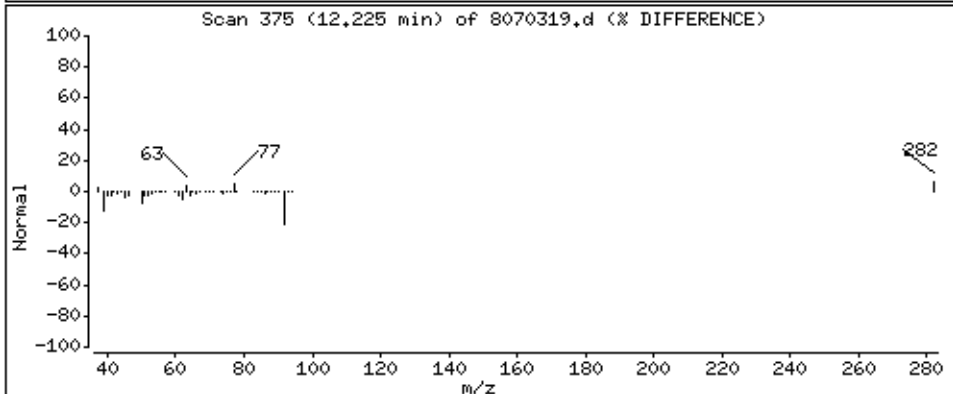
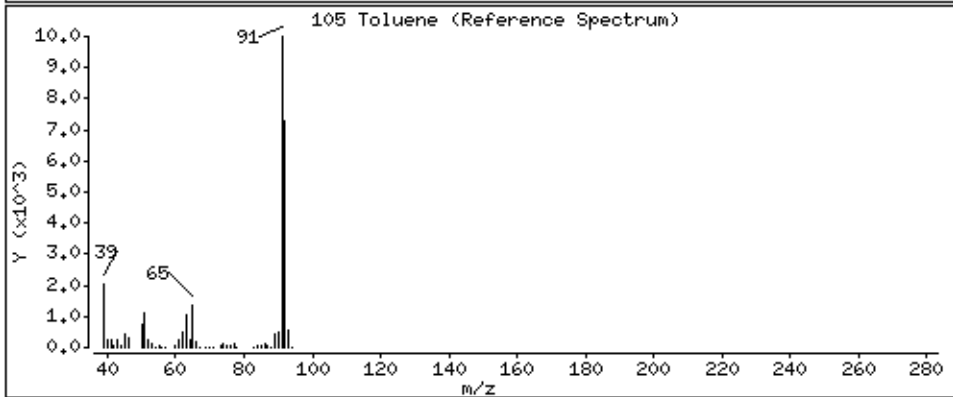
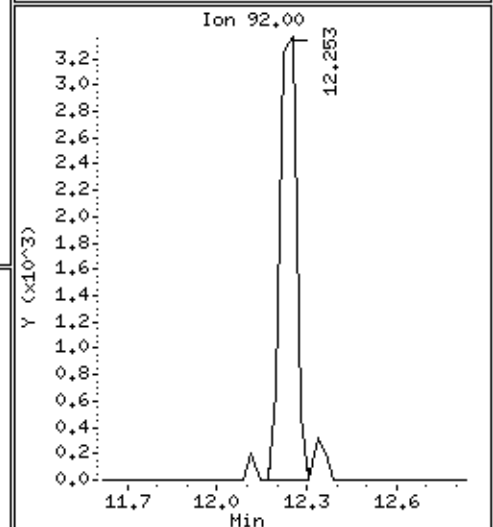
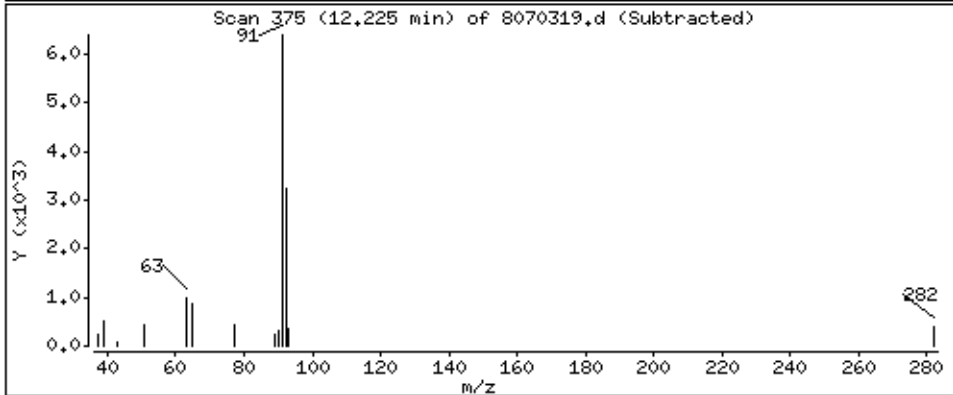
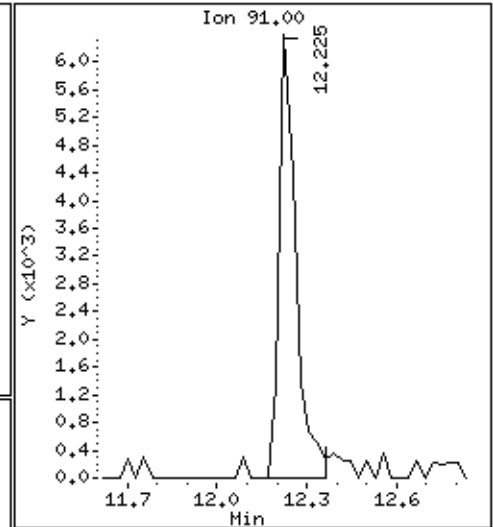
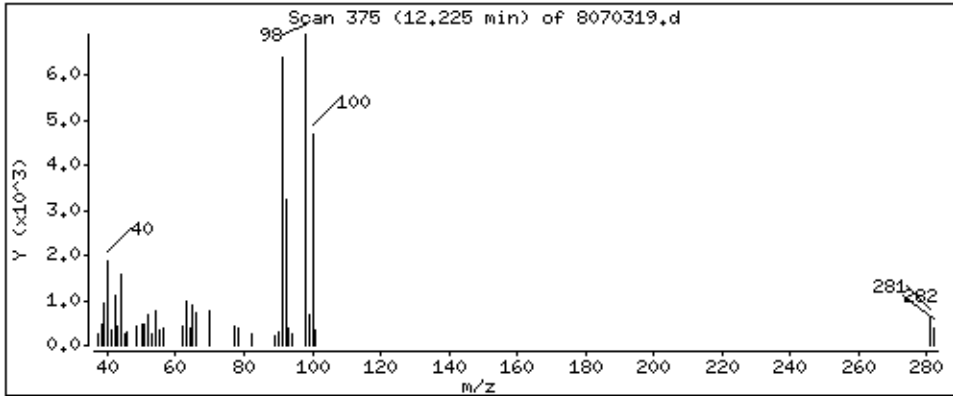
Operator: jdj

Column phase: RTx-624

Column diameter: 0.53

105 Toluene

Concentration: 0.8941 PPBV





AN ENVIRONMENTAL ANALYTICAL LABORATORY

Summary of Detected Compounds
MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

Client Sample ID: TRIP BLANK

Lab ID#: 0706440-04A

No Detections Were Found.



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: TRIP BLANK

Lab ID#: 0706440-04A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070320	Date of Collection: 6/20/07
Dil. Factor:	1.00	Date of Analysis: 7/3/07 09:31 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Freon 12	0.50	Not Detected	2.5	Not Detected
Freon 114	0.50	Not Detected	3.5	Not Detected
Vinyl Chloride	0.50	Not Detected	1.3	Not Detected
Bromomethane	0.50	Not Detected	1.9	Not Detected
Chloroethane	0.50	Not Detected	1.3	Not Detected
Freon 11	0.50	Not Detected	2.8	Not Detected
1,1-Dichloroethene	0.50	Not Detected	2.0	Not Detected
Freon 113	0.50	Not Detected	3.8	Not Detected
Methylene Chloride	0.50	Not Detected	1.7	Not Detected
1,1-Dichloroethane	0.50	Not Detected	2.0	Not Detected
cis-1,2-Dichloroethene	0.50	Not Detected	2.0	Not Detected
Chloroform	0.50	Not Detected	2.4	Not Detected
1,1,1-Trichloroethane	0.50	Not Detected	2.7	Not Detected
Carbon Tetrachloride	0.50	Not Detected	3.1	Not Detected
Benzene	0.50	Not Detected	1.6	Not Detected
1,2-Dichloroethane	0.50	Not Detected	2.0	Not Detected
Trichloroethene	0.50	Not Detected	2.7	Not Detected
1,2-Dichloropropane	0.50	Not Detected	2.3	Not Detected
cis-1,3-Dichloropropene	0.50	Not Detected	2.3	Not Detected
Toluene	0.50	Not Detected	1.9	Not Detected
trans-1,3-Dichloropropene	0.50	Not Detected	2.3	Not Detected
1,1,2-Trichloroethane	0.50	Not Detected	2.7	Not Detected
Tetrachloroethene	0.50	Not Detected	3.4	Not Detected
1,2-Dibromoethane (EDB)	0.50	Not Detected	3.8	Not Detected
Chlorobenzene	0.50	Not Detected	2.3	Not Detected
Ethyl Benzene	0.50	Not Detected	2.2	Not Detected
m,p-Xylene	0.50	Not Detected	2.2	Not Detected
o-Xylene	0.50	Not Detected	2.2	Not Detected
Styrene	0.50	Not Detected	2.1	Not Detected
1,1,2,2-Tetrachloroethane	0.50	Not Detected	3.4	Not Detected
1,3,5-Trimethylbenzene	0.50	Not Detected	2.4	Not Detected
1,2,4-Trimethylbenzene	0.50	Not Detected	2.4	Not Detected
1,3-Dichlorobenzene	0.50	Not Detected	3.0	Not Detected
1,4-Dichlorobenzene	0.50	Not Detected	3.0	Not Detected
alpha-Chlorotoluene	0.50	Not Detected	2.6	Not Detected
1,2-Dichlorobenzene	0.50	Not Detected	3.0	Not Detected
1,3-Butadiene	0.50	Not Detected	1.1	Not Detected
Hexane	0.50	Not Detected	1.8	Not Detected
Cyclohexane	0.50	Not Detected	1.7	Not Detected



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: TRIP BLANK

Lab ID#: 0706440-04A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070320	Date of Collection:	6/20/07
Dil. Factor:	1.00	Date of Analysis:	7/3/07 09:31 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Heptane	0.50	Not Detected	2.0	Not Detected
Bromodichloromethane	0.50	Not Detected	3.4	Not Detected
Dibromochloromethane	0.50	Not Detected	4.2	Not Detected
Cumene	0.50	Not Detected	2.4	Not Detected
Propylbenzene	0.50	Not Detected	2.4	Not Detected
Chloromethane	2.0	Not Detected	4.1	Not Detected
1,2,4-Trichlorobenzene	2.0	Not Detected	15	Not Detected
Hexachlorobutadiene	2.0	Not Detected	21	Not Detected
Acetone	2.0	Not Detected	4.8	Not Detected
Carbon Disulfide	0.50	Not Detected	1.6	Not Detected
2-Propanol	2.0	Not Detected	4.9	Not Detected
trans-1,2-Dichloroethene	0.50	Not Detected	2.0	Not Detected
2-Butanone (Methyl Ethyl Ketone)	0.50	Not Detected	1.5	Not Detected
Tetrahydrofuran	0.50	Not Detected	1.5	Not Detected
1,4-Dioxane	2.0	Not Detected	7.2	Not Detected
4-Methyl-2-pentanone	0.50	Not Detected	2.0	Not Detected
2-Hexanone	2.0	Not Detected	8.2	Not Detected
Bromoform	0.50	Not Detected	5.2	Not Detected
4-Ethyltoluene	0.50	Not Detected	2.4	Not Detected
Ethanol	2.0	Not Detected	3.8	Not Detected
Methyl tert-butyl ether	0.50	Not Detected	1.8	Not Detected
3-Chloropropene	2.0	Not Detected	6.3	Not Detected
2,2,4-Trimethylpentane	0.50	Not Detected	2.3	Not Detected
Naphthalene	2.0	Not Detected	10	Not Detected

Container Type: 6 Liter Summa Canister (100% Certified)

Surrogates	%Recovery	Method Limits
Toluene-d8	100	70-130
1,2-Dichloroethane-d4	104	70-130
4-Bromofluorobenzene	102	70-130

Report Date: 05-Jul-2007 09:32

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070320.d
 Lab Smp Id: 0706440-04A
 Inj Date : 03-JUL-2007 21:31
 Operator : jdg Inst ID: msd8.i
 Smp Info : 200ml #35155
 Misc Info : 4.4psi-4.4psi
 Comment :
 Method : /var/chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 14:39 dpage Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS								
RT	EXP RT	(REL RT)	MASS	RESPONSE		TARGET RANGE	RATIO	
				(PPBV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====

* 68	Bromochloromethane					CAS #: 74-97-5		
7.387	7.387	(1.000)	130	262769	25.0000	70.00- 130.00	100.00	
7.387	7.387	(1.000)	128	200095		47.25- 107.25	76.15	
7.387	7.387	(1.000)	49	386645		115.07- 175.07	147.14	

* 88	1,4-Difluorobenzene					CAS #: 540-36-3		
9.267	9.267	(1.000)	114	1075454	25.0000	70.00- 130.00	100.00	
9.267	9.267	(1.000)	88	158390		0.00- 44.97	14.73	

* 125	Chlorobenzene-d5					CAS #: 3114-55-4		
14.576	14.576	(1.000)	117	811260	25.0000	70.00- 130.00	100.00	
14.576	14.576	(1.000)	82	453330		0.00- 30.00	55.88	

\$ 82	1,2-Dichloroethane-d4					CAS #: 17060-07-0		
8.465	8.465	(1.146)	65	358368	26.0583	26.058 70.00- 130.00	100.00	
8.465	8.465	(1.146)	67	179788		0.00- 30.00	50.17	

\$ 104	Toluene-d8					CAS #: 2037-26-5		
12.115	12.115	(1.307)	98	929799	24.9732	24.973 70.00- 130.00	100.00	
12.115	12.115	(1.307)	70	94093		0.00- 30.00	10.12	

CONCENTRATIONS

ON-COL FINAL

RT EXP RT (REL RT) MASS RESPONSE (PPEV) (PPBV) TARGET RANGE RATIO
== =====

\$ 104 Toluene-d8 (continued)

12.115 12.115 (1.307) 100 590007 0.00- 30.00 63.46

\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207 16.207 (1.112) 174 492601 25.5203 25.520 70.00- 130.00 100.00

16.207 16.207 (1.112) 95 611022 86.34- 146.34 124.04

16.207 16.207 (1.112) 176 476129 66.11- 126.11 96.66

Report Date: 05-Jul-2007 09:32

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARYInstrument ID: msd8.i
Lab File ID: 8070320.d
Lab Smp Id: 0706440-04ACalibration Date: 03-JUL-2007
Calibration Time: 09:41

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdg

Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 4.4psi-4.4psi

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	262769	-24.04
88 1,4-Difluorobenze	1512218	907331	2117105	1075454	-28.88
125 Chlorobenzene-d5	1120246	672148	1568344	811260	-27.58

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
Sample Matrix: GAS Fraction: VOA
Lab Smp Id: 0706440-04A
Level: LOW Operator: jdg
Data Type: MS DATA SampleType: SAMPLE
SpikeList File: AT04+NA-2.spk Quant Type: ISTD
Sublist File: AT04.sub
Method File: /var/chem/msd8.i/8-03jul.b/t14q530b.m
Misc Info: 4.4psi-4.4psi

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	26.058	104.23	70-130
\$ 104 Toluene-d8	25.000	24.973	99.89	70-130
\$ 140 Bromofluorobenzene	25.000	25.520	102.08	70-130

Data File: /chem/msd8.1/8-03jul.b/8070320.d

Date : 03-JUL-2007 21:31

Client ID:

Sample Info: 200ml #35155

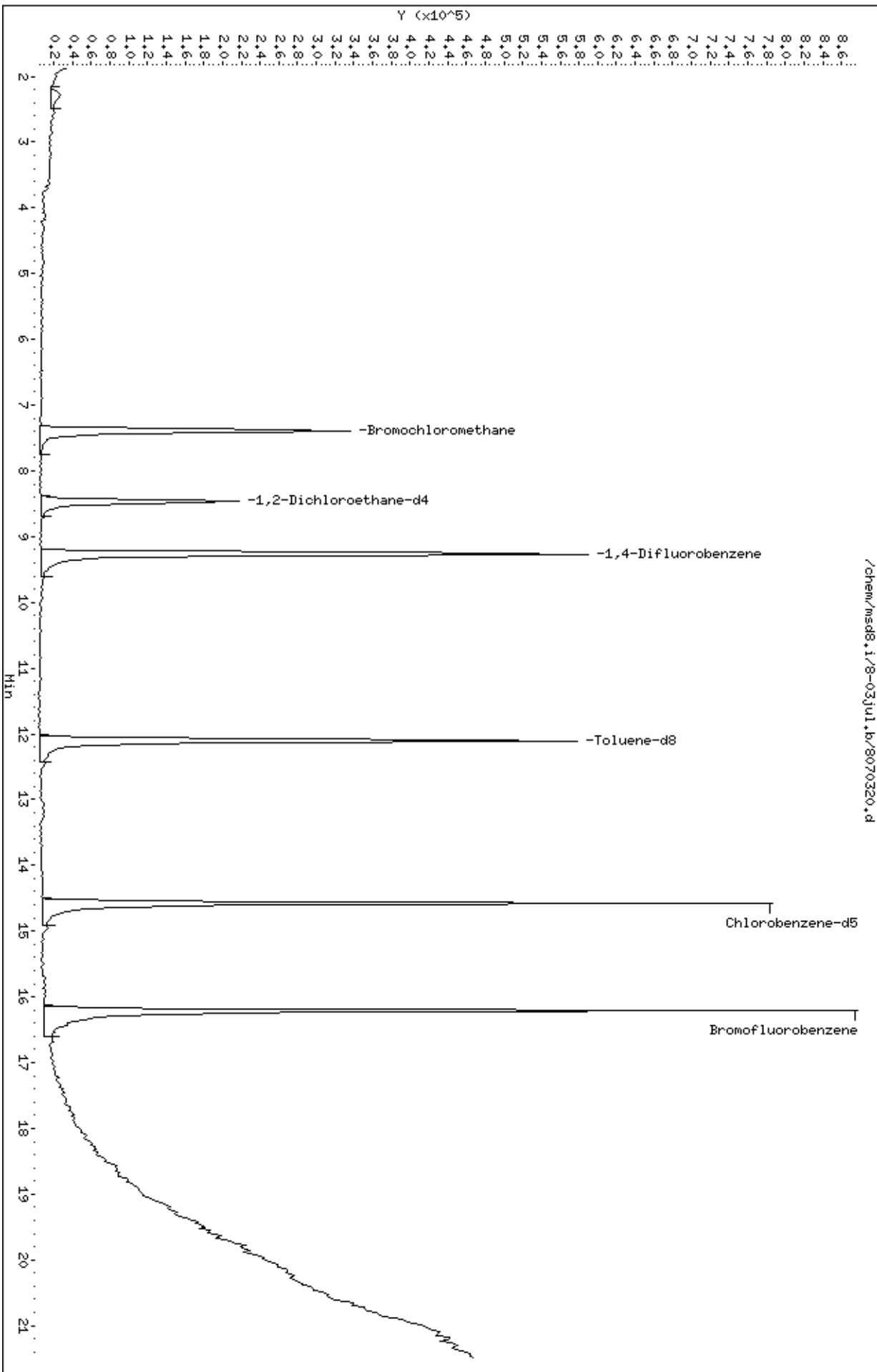
Column phase: RTX-624

Instrument: msd8.1

Operator: jdg

Column diameter: 0.53

/chem/msd8.1/8-03jul.b/8070320.d



QC Results and Raw Data



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: Lab Blank

Lab ID#: 0706440-05A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070307	Date of Collection: NA
Dil. Factor:	1.00	Date of Analysis: 7/3/07 12:17 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Freon 12	0.50	Not Detected	2.5	Not Detected
Freon 114	0.50	Not Detected	3.5	Not Detected
Vinyl Chloride	0.50	Not Detected	1.3	Not Detected
Bromomethane	0.50	Not Detected	1.9	Not Detected
Chloroethane	0.50	Not Detected	1.3	Not Detected
Freon 11	0.50	Not Detected	2.8	Not Detected
1,1-Dichloroethene	0.50	Not Detected	2.0	Not Detected
Freon 113	0.50	Not Detected	3.8	Not Detected
Methylene Chloride	0.50	Not Detected	1.7	Not Detected
1,1-Dichloroethane	0.50	Not Detected	2.0	Not Detected
cis-1,2-Dichloroethene	0.50	Not Detected	2.0	Not Detected
Chloroform	0.50	Not Detected	2.4	Not Detected
1,1,1-Trichloroethane	0.50	Not Detected	2.7	Not Detected
Carbon Tetrachloride	0.50	Not Detected	3.1	Not Detected
Benzene	0.50	Not Detected	1.6	Not Detected
1,2-Dichloroethane	0.50	Not Detected	2.0	Not Detected
Trichloroethene	0.50	Not Detected	2.7	Not Detected
1,2-Dichloropropane	0.50	Not Detected	2.3	Not Detected
cis-1,3-Dichloropropene	0.50	Not Detected	2.3	Not Detected
Toluene	0.50	Not Detected	1.9	Not Detected
trans-1,3-Dichloropropene	0.50	Not Detected	2.3	Not Detected
1,1,2-Trichloroethane	0.50	Not Detected	2.7	Not Detected
Tetrachloroethene	0.50	Not Detected	3.4	Not Detected
1,2-Dibromoethane (EDB)	0.50	Not Detected	3.8	Not Detected
Chlorobenzene	0.50	Not Detected	2.3	Not Detected
Ethyl Benzene	0.50	Not Detected	2.2	Not Detected
m,p-Xylene	0.50	Not Detected	2.2	Not Detected
o-Xylene	0.50	Not Detected	2.2	Not Detected
Styrene	0.50	Not Detected	2.1	Not Detected
1,1,2,2-Tetrachloroethane	0.50	Not Detected	3.4	Not Detected
1,3,5-Trimethylbenzene	0.50	Not Detected	2.4	Not Detected
1,2,4-Trimethylbenzene	0.50	Not Detected	2.4	Not Detected
1,3-Dichlorobenzene	0.50	Not Detected	3.0	Not Detected
1,4-Dichlorobenzene	0.50	Not Detected	3.0	Not Detected
alpha-Chlorotoluene	0.50	Not Detected	2.6	Not Detected
1,2-Dichlorobenzene	0.50	Not Detected	3.0	Not Detected
1,3-Butadiene	0.50	Not Detected	1.1	Not Detected
Hexane	0.50	Not Detected	1.8	Not Detected
Cyclohexane	0.50	Not Detected	1.7	Not Detected



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: Lab Blank

Lab ID#: 0706440-05A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070307	Date of Collection: NA
Dil. Factor:	1.00	Date of Analysis: 7/3/07 12:17 PM

Compound	Rpt. Limit (ppbv)	Amount (ppbv)	Rpt. Limit (uG/m3)	Amount (uG/m3)
Heptane	0.50	Not Detected	2.0	Not Detected
Bromodichloromethane	0.50	Not Detected	3.4	Not Detected
Dibromochloromethane	0.50	Not Detected	4.2	Not Detected
Cumene	0.50	Not Detected	2.4	Not Detected
Propylbenzene	0.50	Not Detected	2.4	Not Detected
Chloromethane	2.0	Not Detected	4.1	Not Detected
1,2,4-Trichlorobenzene	2.0	Not Detected	15	Not Detected
Hexachlorobutadiene	2.0	Not Detected	21	Not Detected
Acetone	2.0	Not Detected	4.8	Not Detected
Carbon Disulfide	0.50	Not Detected	1.6	Not Detected
2-Propanol	2.0	Not Detected	4.9	Not Detected
trans-1,2-Dichloroethene	0.50	Not Detected	2.0	Not Detected
2-Butanone (Methyl Ethyl Ketone)	0.50	Not Detected	1.5	Not Detected
Tetrahydrofuran	0.50	Not Detected	1.5	Not Detected
1,4-Dioxane	2.0	Not Detected	7.2	Not Detected
4-Methyl-2-pentanone	0.50	Not Detected	2.0	Not Detected
2-Hexanone	2.0	Not Detected	8.2	Not Detected
Bromoform	0.50	Not Detected	5.2	Not Detected
4-Ethyltoluene	0.50	Not Detected	2.4	Not Detected
Ethanol	2.0	Not Detected	3.8	Not Detected
Methyl tert-butyl ether	0.50	Not Detected	1.8	Not Detected
3-Chloropropene	2.0	Not Detected	6.3	Not Detected
2,2,4-Trimethylpentane	0.50	Not Detected	2.3	Not Detected
Naphthalene	2.0	Not Detected	10	Not Detected

Container Type: NA - Not Applicable

Surrogates	%Recovery	Method Limits
Toluene-d8	98	70-130
1,2-Dichloroethane-d4	104	70-130
4-Bromofluorobenzene	101	70-130

Report Date: 03-Jul-2007 12:30

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070307.d
 Lab Smp Id: Lab Blank Client Smp ID: Lab Blank
 Inj Date : 03-JUL-2007 12:17
 Operator : jdg Inst ID: msd8.i
 Smp Info : 200mL #13673
 Misc Info : Humid
 Comment :
 Method : /chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 12:18 jgray Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS									
ON-COL FINAL									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	(PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	263558	25.0000		80.00- 120.00	100.00	
7.387	7.387	(1.000)	128	207036			47.25- 107.25	78.55	
7.387	7.387	(1.000)	49	406728			115.07- 175.07	154.32	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	1147745	25.0000		80.00- 120.00	100.00	
9.267	9.267	(1.000)	88	170039			0.00- 44.97	14.82	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	859637	25.0000		80.00- 120.00	100.00	
14.576	14.576	(1.000)	82	480326			0.00- 30.00	55.88	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465	(1.146)	65	359749	26.0804	26.080	80.00- 120.00	100.00	
8.465	8.465	(1.146)	67	182238			0.00- 30.00	50.66	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115	(1.307)	98	975843	24.5590	24.559	80.00- 120.00	100.00	
12.115	12.115	(1.307)	70	98849			0.00- 30.00	10.13	

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPEV)	(PPBV)	TARGET RANGE	RATIO
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\$ 104 Toluene-d8 (continued)

12.115	12.115	(1.307)	100	655421			0.00- 30.00	67.16
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\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207	16.207	(1.112)	174	517793	25.3158	25.316	80.00- 120.00	100.00
16.207	16.207	(1.112)	95	650555			86.34- 146.34	125.64
16.207	16.207	(1.112)	176	497306			66.11- 126.11	96.04

Report Date: 03-Jul-2007 12:30

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 03-JUL-2007

Lab File ID: 8070307.d

Calibration Time: 09:41

Lab Smp Id: Lab Blank

Client Smp ID: Lab Blank

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdg

Method File: /chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: Humid

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	263558	-23.81
88 1,4-Difluorobenze	1512218	907331	2117105	1147745	-24.10
125 Chlorobenzene-d5	1120246	672148	1568344	859637	-23.26

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
Sample Matrix: GAS Fraction: VOA
Lab Smp Id: Lab Blank Client Smp ID: Lab Blank
Level: LOW Operator: jdg
Data Type: MS DATA SampleType: SAMPLE
SpikeList File: AT04+NA-2.spk Quant Type: ISTD
Sublist File: AT04+ENSR.sub
Method File: /chem/msd8.i/8-03jul.b/t14q530b.m
Misc Info: Humid

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	26.080	104.32	70-130
\$ 104 Toluene-d8	25.000	24.559	98.24	70-130
\$ 140 Bromofluorobenzene	25.000	25.316	101.26	70-130

Data File: /chem/msd8.1/8-03jul.b/8070307.d

Date : 03-JUL-2007 12:17

Client ID: Lab Blank

Sample Info: 200mL #13673

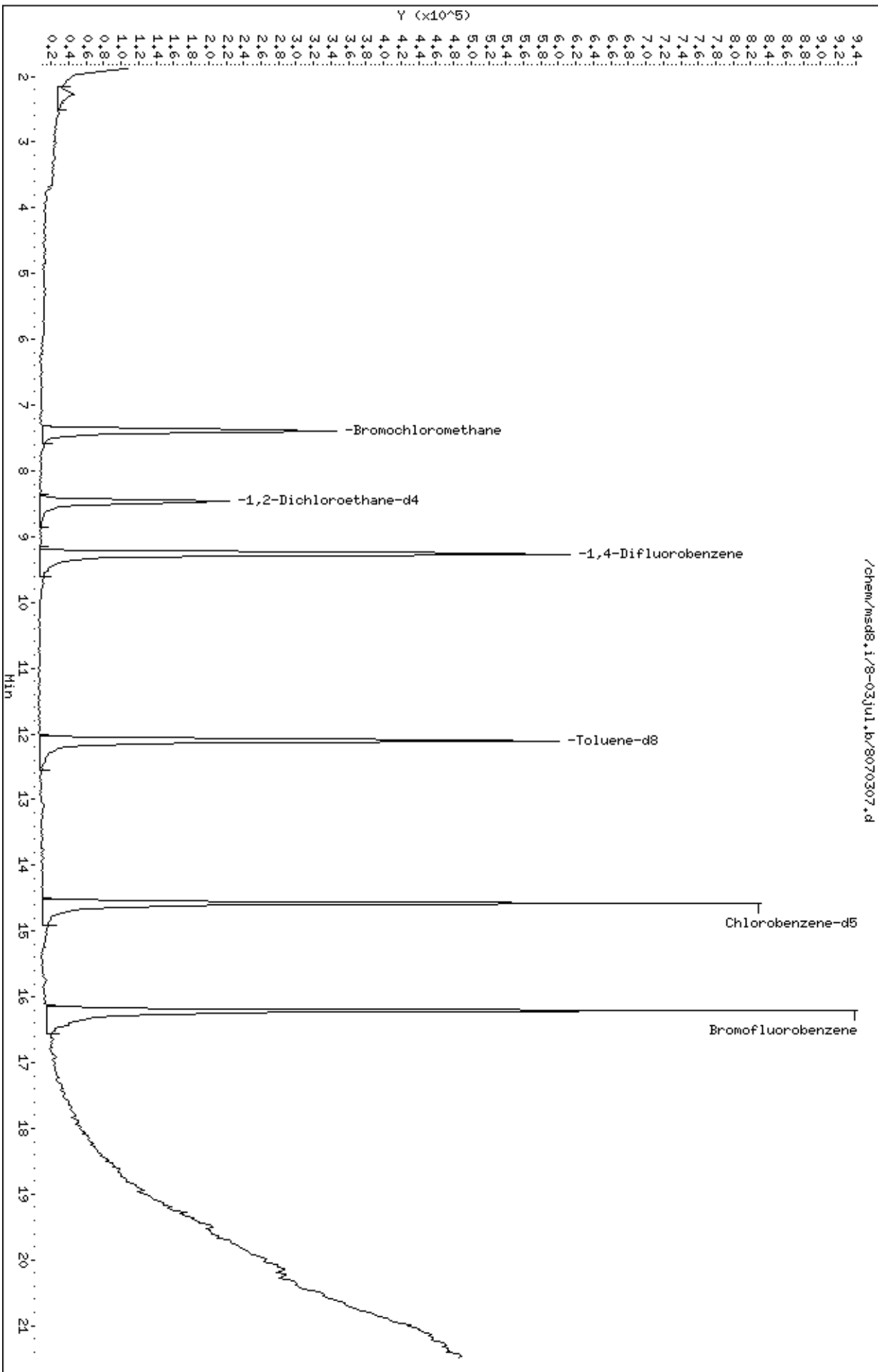
Column phase: RTX-624

Instrument: msd8.1

Operator: jdg

Column diameter: 0.53

/chem/msd8.1/8-03jul.b/8070307.d



LEVEL-IV VALIDATABLE

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

SURROGATE RECOVERY FORM

Lab Name: AIR TOXICS LIMITED.

SDG No.: 0706440

	CLIENT SAMPLE NO.	SURROGATE % RECOVERY						TOTAL OUT
		1,2-Dichloroethane-d 4	#	Toluene-d8	#	4-Bromofluorobenze ne	#	
01	AMS 5 DW (4148)	107		98		104		0
02	AMS 5 DW (3746)	105		101		104		0
03	AMS 5 DW (3746) Lab Duplicate	106		98		102		0
04	AMS 2 UW	102		98		99		0
05	TRIP BLANK	104		100		102		0
06	Lab Blank	104		98		101		0
07	CCV	103		100		106		0
08	LCS	102		103		112		0
09								0
10								0
11								0
12								0
13								0
14								0
15								0
16								0
17								0
18								0
19								0
20								0
21								0
22								0
23								0
24								0

Surrogate Recovery Limits

1,2-Dichloroethane-d4 70 - 130

Toluene-d8 70 - 130

4-Bromofluorobenzene 70 - 130

* Designates values outside of QC limits

LEVEL-IV VALIDATABLE

Modified EPA Method TO-15 GC/MS Full Scan

INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: AIR TOXICS, LTD
 Lab File ID: 8070302.d
 Instrument ID: msd8.i

SDG No: 0706440
 Date Analyzed: 07/03/2007
 Time Analyzed: 09:41 AM

	Chlorobenzene-d5		RT		1,4-Difluorobenzene		RT		Bromochloromethane		RT	
	Area	#		#	Area	#		#	Area	#		#
	24-HOUR STD	1120246		14.58	1512218		9.27		345936		7.39	
	UPPER LIMIT	1568344		14.91	2117105		09.60		484310		07.72	
	LOWER LIMIT	672148		14.25	907331		08.94		207562		07.06	
	CLIENT SAMPLE NO											
01	AMS 5 DW (4148)	791246		14.58	1055614		9.27		247531		7.39	
02	AMS 5 DW (3746)	801011		14.58	1070881		9.27		251316		7.39	
03	AMS 5 DW (3746) Lab Duplicate	804503		14.58	1069480		9.27		254121		7.39	
04	AMS 2 UW	796366		14.58	1079703		9.27		260882		7.39	
05	TRIP BLANK	811260		14.58	1075454		9.27		262769		7.39	
06	Lab Blank	859637		14.58	1147745		9.27		263558		7.39	
07	CCV	1120246		14.58	1512218		9.27		345936		7.39	
08	LCS	882830		14.58	1169163		9.27		278074		7.39	
09												
10												
11												
12												
13												
14												
15												
16												
17												
18												
19												
20												
21												
22												

'Area Upper Limit=+40% of internal standard area'
 'Area Lower Limit=-40% of internal standard area'

RT Upper Limit=+0.33 minutes of internal standard RT
 RT Lower Limit=-0.33 minutes of internal standard RT

* Designates values outside of QC limits

SAMPLE RESULTS/SAMPLE RESULTS DUPLICATE

Lab Name: Air Toxics Ltd.
 Lab Sample ID: 02A & 02AA
 Client Sample ID: &

Lab File ID: 8070318.d & 8070317.d
 Dilution: 1.79 & 1.79
 Date Analyzed: 7/3/07 & 7/3/07

CAS Number	Compound	Original		Duplicate		RPD
		Amount	Flags	Amount	Flags	
71-55-6	1,1,1-Trichloroethane	ND	U	ND	U	0
79-34-5	1,1,2,2-Tetrachloroethane	ND	U	ND	U	0
79-00-5	1,1,2-Trichloroethane	ND	U	ND	U	0
75-34-3	1,1-Dichloroethane	ND	U	ND	U	0
75-35-4	1,1-Dichloroethene	ND	U	ND	U	0
120-82-1	1,2,4-Trichlorobenzene	ND	U	ND	U	0
95-63-6	1,2,4-Trimethylbenzene	ND	U	ND	U	0
106-93-4	1,2-Dibromoethane (EDB)	ND	U	ND	U	0
95-50-1	1,2-Dichlorobenzene	ND	U	ND	U	0
107-06-2	1,2-Dichloroethane	ND	U	ND	U	0
78-87-5	1,2-Dichloropropane	ND	U	ND	U	0
108-67-8	1,3,5-Trimethylbenzene	ND	U	ND	U	0
106-99-0	1,3-Butadiene	ND	U	ND	U	0
541-73-1	1,3-Dichlorobenzene	ND	U	ND	U	0
106-46-7	1,4-Dichlorobenzene	ND	U	ND	U	0
123-91-1	1,4-Dioxane	ND	U	ND	U	0
540-84-1	2,2,4-Trimethylpentane	ND	U	ND	U	0
78-93-3	2-Butanone (Methyl Ethyl Ketone)	2.925		2.733		6.8
591-78-6	2-Hexanone	ND	U	ND	U	0
67-63-0	2-Propanol	ND	U	ND	U	0
107-05-1	3-Chloropropene	ND	U	ND	U	0
622-96-8	4-Ethyltoluene	ND	U	ND	U	0
108-10-1	4-Methyl-2-pentanone	ND	U	ND	U	0
67-64-1	Acetone	24.263		22.91		5.7
100-44-7	alpha-Chlorotoluene	ND	U	ND	U	0
71-43-2	Benzene	ND	U	ND	U	0
75-27-4	Bromodichloromethane	ND	U	ND	U	0
75-25-2	Bromoforr	ND	U	ND	U	0
74-83-9	Bromomethane	ND	U	ND	U	0
75-15-0	Carbon Disulfide	ND	U	ND	U	0
56-23-5	Carbon Tetrachloride	ND	U	ND	U	0
108-90-7	Chlorobenzene	ND	U	ND	U	0
75-00-3	Chloroethane	ND	U	ND	U	0
67-66-3	Chloroforr	ND	U	ND	U	0
74-87-3	Chloromethane	ND	U	ND	U	0
156-59-2	cis-1,2-Dichloroethene	ND	U	ND	U	0
10061-01-5	cis-1,3-Dichloropropene	ND	U	ND	U	0
98-82-8	Cumene	ND	U	ND	U	0
110-82-7	Cyclohexane	ND	U	ND	U	0
124-48-1	Dibromochloromethane	ND	U	ND	U	0
64-17-5	Ethanol	ND	U	ND	U	0
100-41-4	Ethyl Benzene	ND	U	ND	U	0
75-69-4	Freon 11	ND	U	ND	U	0
76-13-1	Freon 113	ND	U	ND	U	0
76-14-2	Freon 114	ND	U	ND	U	0
75-71-8	Freon 12	ND	U	ND	U	0

Note: The results appearing in the Amount columns are the raw, unrounded numbers acquired from the instrument.

SAMPLE RESULTS/SAMPLE RESULTS DUPLICATE

Lab Name: Air Toxics Ltd.
 Lab Sample ID: 02A & 02AA
 Client Sample ID: &

Lab File ID: 8070318.d & 8070317.d
 Dilution: 1.79 & 1.79
 Date Analyzed: 7/3/07 & 7/3/07

CAS Number	Compound	Original		Duplicate		RPD
		Amount	Flags	Amount	Flags	
142-82-5	Heptane	ND	U	ND	U	0
87-68-3	Hexachlorobutadiene	ND	U	ND	U	0
110-54-3	Hexane	ND	U	ND	U	0
108-38-3	m,p-Xylene	ND	U	ND	U	0
1634-04-4	Methyl tert-butyl ether	ND	U	ND	U	0
75-09-2	Methylene Chloride	ND	U	ND	U	0
91-20-3	Naphthalene	ND	U	ND	U	0
95-47-6	o-Xylene	ND	U	ND	U	0
103-65-1	Propylbenzene	ND	U	ND	U	0
100-42-5	Styrene	ND	U	ND	U	0
127-18-4	Tetrachloroethene	ND	U	ND	U	0
109-99-9	Tetrahydrofuran	ND	U	ND	U	0
108-88-3	Toluene	ND	U	ND	U	0
156-60-5	trans-1,2-Dichloroethene	ND	U	ND	U	0
10061-02-6	trans-1,3-Dichloropropene	ND	U	ND	U	0
79-01-6	Trichloroethene	ND	U	ND	U	0
75-01-4	Vinyl Chloride	ND	U	ND	U	0

Note: The results appearing in the Amount columns are the raw, unrounded numbers acquired from the instrument.

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Calibration File Names:

- Level 1: /chem/msd8.i/8-30may.b/8053003.d
- Level 2: /chem/msd8.i/8-30may.b/8053004.d
- Level 3: /chem/msd8.i/8-07jun.b/8060704.d
- Level 4: /chem/msd8.i/8-30may.b/8053006.d
- Level 5: /chem/msd8.i/8-07jun.b/8060705.d
- Level 6: /chem/msd8.i/8-30may.b/8053008.d
- Level 7: /chem/msd8.i/8-07jun.b/8060706.d

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	___	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	
	200.000							
	Level 7							
1 Freon 152a	+++++	+++++	0.98763	+++++	0.78978	+++++		
	0.82069						0.86603	12.290
2 Freon 22	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
3 Propylene	+++++	+++++	1.92982	1.31817	1.20253	1.15722		
	1.15784						1.35311	24.315
4 Dichlorodifluoromethane/Fr12	+++++	4.22735	4.02658	3.34137	3.02222	2.98122		
	2.95137						3.42502	16.496
5 Freon134a	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
6 Freon 114	+++++	3.61995	3.07946	3.11187	3.01070	2.87973		
	2.86380						3.09425	8.944
7 Isobutane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000 Level 1	0.50000 Level 2	2.000 Level 3	25.000 Level 4	50.000 Level 5	100.000 Level 6	Level 7	RRF	% RSD
8 Chloromethane	200.000 1.41963	+++++	1.76358	1.64915	1.55747	1.49684		1.57733	8.482
9 Butane	0.37508	+++++	0.49593	0.43121	0.39247	0.36498		0.41193	12.942
10 1,3-Butadiene	1.37941	2.12647	1.72912	1.45627	1.41158	1.30372		1.56776	19.767
11 Vinyl Chloride	1.64028	2.34753	1.85221	1.78079	1.74379	1.63245		1.83284	14.500
12 Methanol	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
13 Bromomethane	1.20683	1.25815	1.19877	1.26616	1.22534	1.19166		1.22449	2.562
14 Vinyl Bromide	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
15 Isopentane	2.24216	+++++	2.38563	2.37944	2.25961	2.17613		2.28859	3.988
16 Chloroethane	0.87782	1.10636	1.03490	0.98458	0.94664	0.86976		0.97001	9.461
17 Dichlorofluoromethane/Fr21	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
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 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000 Level 1	0.50000 Level 2	2.000 Level 3	25.000 Level 4	50.000 Level 5	100.000 Level 6	Level 7	RRF	% RSD
18 Trichlorofluoromethane/Fr11	200.000 3.70548	4.06616	4.21652	3.95963	3.78415	3.62588		3.89297	5.829
19 Pentane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
20 Freon123a	1.29778	+++++	1.38653	+++++	1.32725	+++++		1.33719	3.380
21 Freon123	0.82221	+++++	0.86166	+++++	0.84444	+++++		0.84277	2.347
22 Dimethyl Ether	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
23 Ethanol	0.57722	+++++	0.67753	0.66346	0.60794	0.58208		0.62165	7.458
24 Freon 13	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
25 Acrolein	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
26 Isobutylene	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
27 Freon142b	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
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 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	—	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	
	200.000							
	Level 7							
28 Freon 113	+++++	2.88068	2.79775	2.60107	2.43015	2.35327		
	2.39146						2.57573	8.639
29 1,1-Dichloroethene	+++++	2.86789	2.72869	2.56996	2.43351	2.34903		
	2.39867						2.55796	8.009
30 Acetone	+++++	+++++	0.98245	0.85434	0.84281	0.79818		
	0.80581						0.85672	8.662
31 Acetaldehyde	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
32 Freon143a	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
33 Carbon Disulfide	+++++	5.58896	4.86914	4.79052	4.54598	4.41760		
	4.55167						4.79398	8.848
34 2-Propanol	+++++	+++++	3.63428	3.09423	2.96019	2.88102		
	2.91819						3.09759	10.029
35 Acetonitrile	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
36 Cyclopentene	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
37 3-Chloropropene	+++++	+++++	0.72576	0.81455	0.78451	0.74233		
	0.75685						0.76480	4.602

Air Toxics Ltd.

INITIAL CALIBRATION DATA

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 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	---	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	
	200.000							
	Level 7							
48 Ethanol-high	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
	+++++						+++++	+++++
49 Isopropyl ether	+++++	+++++	4.44068	+++++	4.38024	+++++		
	4.25213						4.35768	2.209
50 Propanal	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
51 Chloroprene	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
52 1-Propanol	+++++	+++++	0.33412	+++++	0.29773	+++++		
	0.31039						0.31408	5.881
53 Bromoethane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
54 1,1-Dichloroethane	+++++	2.90490	3.23579	2.99995	2.89542	2.78581		
	2.88219						2.95068	5.266
55 Vinyl Acetate	+++++	+++++	0.40978	0.42041	0.40294	0.41753		
	0.43205						0.41654	2.650
56 Iodomethane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
57 2,3-Dimethylbutane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	—	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	
	200.000							
	Level 7							
58 Ethyl-tert-butyl Ether	+++++	+++++	2.61890	+++++	3.32619	+++++		
	2.78316						2.90942	12.723
59 Methyl Acetate	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
60 2,2-Dichloropropane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
61 Ethyl Acetate	+++++	+++++	0.39734	+++++	0.37435	+++++		
	0.36735						0.37968	4.133
62 1-Hexene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
63 Methyl Acrylate	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
64 cis-1,2-Dichloroethene	+++++	3.03723	2.36908	2.27992	2.14250	2.06765		
	2.11545						2.33530	15.487
65 2-Butanone	+++++	1.00709	0.85028	0.78676	0.77311	0.76198		
	0.79178						0.82850	11.187
66 2,4-Dimethylpentane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
67 Tetrahydrofuran	+++++	2.62724	2.20335	2.11722	2.03416	1.93041		
	2.00841						2.15347	11.624

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000 Level 1	0.50000 Level 2	2.000 Level 3	25.000 Level 4	50.000 Level 5	100.000 Level 6	200.000 Level 7	RRF	% RSD
69 Butanal	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
70 Chloroform	4.52339 2.82903	3.38269	2.97755	2.93420	2.84544	2.73693		3.17561	19.826
71 2-Butanol	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
72 1,1-Dichloropropene	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
73 Cyclohexane	+++++ 2.27273	3.13955	2.37507	2.38858	2.29135	2.22343		2.44845	14.063
74 3-Methyl-1-Hexene	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
75 1,1,1-Trichloroethane	+++++ 2.95670	3.69058	3.04827	3.03489	2.94955	2.87166		3.09194	9.709
76 2,3-Dimethylpentane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
77 Carbon Tetrachloride	+++++ 2.87341	2.95018	2.47054	2.94416	2.84417	2.79886		2.81355	6.321
78 Isobutanol	+++++ 0.29396	+++++	0.27113	+++++	0.29573	+++++		0.28694	4.783

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	RRF	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6		
	200.000							
	Level 7							
79 tert-amyl-Methyl Ether	+++++	+++++	2.60292	+++++	2.98159	+++++		
	2.39754						2.66068	11.136
80 2,2,4-Trimethylpentane	+++++	7.83756	7.57621	7.24629	6.93326	6.86073		
	7.40163						7.30928	5.139
81 Benzene	1.46711	1.32521	1.13921	1.09388	1.07898	1.04564		
	1.09326						1.17761	13.352
83 1,2-Dichloroethane	+++++	0.56444	0.46007	0.44951	0.43040	0.41345		
	0.43142						0.45822	11.897
84 Thiopene	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
85 Heptane	+++++	0.19081	0.11421	0.11021	0.10683	0.10369		
	0.10755						0.12222	27.645
86 1-Methoxy-2-Propanol	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
87 2,3,4-Trimethylpentane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
89 1-Butanol	+++++	+++++	0.22243	+++++	0.24711	+++++		
	0.26464						0.24472	8.665
90 Methyl Methacrylate	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000		
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	% RSD
	200.000							
	Level 7							
91 2-Pentanone	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
92 Pentanal	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
93 Ethyl Acrylate	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
94 Trichloroethene	+++++	0.53286	0.44939	0.44327	0.42705	0.41017	0.44757	9.860
95 Methyl Cyclohexane	+++++	3.26599	3.09517	3.05667	2.96150	2.88311	3.05031	4.263
96 Dibromomethane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
97 1,2-Dichloropropane	+++++	0.42737	0.43013	0.38160	0.37306	0.36109	0.39140	7.588
98 1,4-Dioxane	+++++	+++++	0.24770	0.24197	0.23695	0.23009	0.23873	2.747
99 Octane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
100 Bromodichloromethane	+++++	0.73233	0.64459	0.64531	0.63931	0.61664	0.65370	6.123

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000 Level 1	0.50000 Level 2	2.000 Level 3	25.000 Level 4	50.000 Level 5	100.000 Level 6	200.000 Level 7	RRF	% RSD
101 1-Nitropropane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
102 cis-1,3-Dichloropropene	+++++	0.61851	0.52709	0.53649	0.53068	0.51575		0.54392	6.854
103 4-Methyl-2-pentanone	+++++	0.28397	0.30418	0.29328	0.29702	0.29012		0.29426	2.342
105 Toluene	+++++	1.25780	1.14795	1.11923	1.10994	1.09416		1.14782	5.129
106 1-Methoxy-2-propyl acetate	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
107 Epichlorohydrin	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
108 trans-1,3-Dichloropropene	+++++	0.75878	0.70145	0.71553	0.72836	0.71539		0.72478	2.694
109 1,3-Dichloropropane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
110 1,1,2-Trichloroethane	+++++	0.66231	0.52902	0.51019	0.50076	0.48912		0.53155	12.321
111 alpha-Pinene	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	—	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	
	200.000							
	Level 7							
122 Dicyclopentadiene	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
123 1,1,1,2-Tetrachloroethane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
124 D-Limonene	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
126 Chlorobenzene	+++++	1.22227	1.44203	1.24119	1.25172	1.24989		
	1.29500						1.28368	6.323
127 Bis(2-chloroethyl) ether	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
128 Nonane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
129 Ethyl Benzene	+++++	0.80931	0.66703	0.64128	0.64323	0.65061		
	0.66455						0.67934	9.504
130 m,p-Xylene	+++++	0.84552	0.85011	0.79451	0.81241	0.81316		
	0.85120						0.82782	2.919
131 Undecane	+++++	+++++	+++++	+++++	+++++	+++++		
	+++++						+++++	+++++
132 o-Xylene	+++++	0.91319	0.84572	0.80595	0.81315	0.80154		
	0.83252						0.83535	4.986

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000 Level 1	0.50000 Level 2	2.000 Level 3	25.000 Level 4	50.000 Level 5	100.000 Level 6	Level 7	RRF	% RSD
133 2-Heptanone	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
134 Styrene	1.11092 1.30443	1.18281	1.11362	1.12817	1.19071	1.22490		1.17936	5.944
135 Bromoform	+++++	0.63208	0.65434	0.73553	0.76857	0.78441		0.73275	10.236
136 Cyclohexanone	+++++	+++++	0.55672	+++++	0.57050	+++++		0.57218	2.859
137 Cumene	2.79880 2.13898	2.40146	2.35517	2.21000	2.28171	2.31208		2.35689	9.067
138 1-chloro-2-Bromopropane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
139 Bromobenzene	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
141 1,2,3-Trichloropropane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
142 Dodecane	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++
143 2-Chlorotoluene	+++++	+++++	+++++	+++++	+++++	+++++		+++++	+++++

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	RRF	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6		
	200.000							
	Level 7							
144 1,1,2,2-Tetrachloroethane	+++++	1.33097	1.23407	1.15272	1.19774	1.19102		
	1.22319						1.22162	4.962
145 Propylbenzene	+++++	2.82083	2.74232	2.62546	2.75408	2.80075		
	1.94674						2.61503	12.788
146 4-Chlorotoluene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
147 4-Ethyltoluene	+++++	2.24895	2.26379	2.34932	2.40819	2.44980		
	1.73174						2.24196	11.686
148 1,3,5-Trimethylbenzene	+++++	2.38847	2.33379	2.17383	2.24009	2.30283		
	1.49096						2.15500	15.486
149 2,6-Dimethyl-1-propanol	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
150 tert-Butylbenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
151 Pentachloroethane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
152 sec-Butylbenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
153 1,2,4-Trimethylbenzene	+++++	2.40359	2.29481	2.25241	2.34010	2.38632		
	1.33793						2.16919	18.952

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000	0.50000	2.000	25.000	50.000	100.000	RRF	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6		
	200.000							
	Level 7							
164 Aniline	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
165 1,2-Dibromo-3-Chloropropane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
166 Isooctyl Alcohol	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
167 1,2,4-Trichlorobenzene	+++++ 1.22253	+++++	1.48368	1.26108	1.43814	1.47814	1.37671	9.090
168 Hexachlorobutadiene	+++++ 0.73297	+++++	0.95260	0.75132	0.77092	0.74328	0.79022	11.621
169 Naphthalene	+++++ 1.31458	+++++	3.90045	3.19414	3.61060	2.49731	2.90342	35.577 <-
170 1,2,3-Trichlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
171 1,3,5-Trichlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
172 Isooctyl Acrylate	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
\$ 82 1,2-Dichloroethane-d4	1.29108 1.39339	1.28164	1.33015	1.25825	1.26386	1.34062	1.30843	3.724

Air Toxics Ltd.

INITIAL CALIBRATION DATA

Start Cal Date : 30-MAY-2007 14:12
 End Cal Date : 07-JUN-2007 12:08
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 3.50
 Integrator : HP RTE
 Method file : /chem/msd8.i/8-07jun.b/t14q530b.m
 Cal Date : 07-Jun-2007 14:14 jgray
 Curve Type : Average

Compound	0.20000 Level 1	0.50000 Level 2	2.000 Level 3	25.000 Level 4	50.000 Level 5	100.000 Level 6	200.000 Level 7	RRF	% RSD
\$ 104 Toluene-d8	0.89493	0.88890	0.87548	0.83261	0.85871	0.85180		0.86549	2.548
\$ 140 Bromofluorobenzene	0.57145	0.56913	0.58528	0.59982	0.61223	0.61820		0.59482	3.318

Calibration History

Method : /chem/msd8.i/8-07jun.b/t14q530b.m
Start Cal Date: 30-MAY-2007 14:12
End Cal Date : 07-JUN-2007 12:08

Initial Calibration

Injection Date	Sublist	Calibration File
Cal Level: 1 , Cal Amount: 0.20000		
30-MAY-2007 14:12	AFCEElow	/chem/msd8.i/8-30may.b/8053003.d
Cal Level: 2 , Cal Amount: 0.50000		
30-MAY-2007 14:39	AT04Low+ENSR	/chem/msd8.i/8-30may.b/8053004.d
Cal Level: 3 , Cal Amount: 2.00000		
07-JUN-2007 11:09	sp16b	/chem/msd8.i/8-07jun.b/8060704.d
30-MAY-2007 15:07	AT04mdl+ENSR	/chem/msd8.i/8-30may.b/8053005.d
Cal Level: 4 , Cal Amount: 25.00000		
30-MAY-2007 15:35	AT04mdl+ENSR	/chem/msd8.i/8-30may.b/8053006.d
Cal Level: 5 , Cal Amount: 50.00000		
07-JUN-2007 11:37	sp16b	/chem/msd8.i/8-07jun.b/8060705.d
30-MAY-2007 16:03	AT04mdl+ENSR	/chem/msd8.i/8-30may.b/8053007.d
Cal Level: 6 , Cal Amount: 100.00000		
30-MAY-2007 16:31	AT04mdl+ENSR	/chem/msd8.i/8-30may.b/8053008.d
Cal Level: 7 , Cal Amount: 200.00000		
07-JUN-2007 12:08	sp16b	/chem/msd8.i/8-07jun.b/8060706.d
30-MAY-2007 17:02	AT04mdl+ENSR	/chem/msd8.i/8-30may.b/8053009.d

Continuing Calibration
Ccal Level Mode: GLOBAL LEVEL 5

+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	Ccal Level: 5 , Ccal Amount: 50.000	
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	07-JUN-2007 10:01 AT04+ENSR	/chem/msd8.i/8-07jun.b/8060702.d
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	Ccal Level: 5 , Ccal Amount: 50.000	
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	07-JUN-2007 11:37 sp16bCCV	/chem/msd8.i/8-07jun.b/8060705a.d
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	Ccal Level: 5 , Ccal Amount: 50.000	
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	07-JUN-2007 11:37 sp16b	/chem/msd8.i/8-07jun.b/8060705.d
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	Ccal Level: 5 , Ccal Amount: 50.000	
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		
	07-JUN-2007 11:37 sp16bCCV	/chem/msd8.i/8-07jun.b/8060705a.d
+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+		

@ Air Toxics Ltd.

MSD-8

% REL. ABUNDANCE

m/z	ION ABUNDANCE CRITERIA	% REL. ABUNDANCE
50	15.0 - 40.0% of mass 95	17.60
75	30.0 - 60.0% of mass 95	44.95
95	Base peak, 100.00% relative abundance	100.00
96	5.0 - 9.0% of mass 95	6.34
173	Less than 2.0% of mass 174	(0.00) ¹
174	Greater than 50.0% of mass 95	82.17
175	5.0 - 9.0% of mass 174	(7.22) ¹
176	Greater than 95.0% but less than 101.0% of mass 174	(97.95) ¹
177	5.0 - 9.0% of mass 176	(6.12) ²

¹ - value in parenthesis is % mass 174

² - value in parenthesis is % mass 176

Verify 176/174 m/z Ratio: ~~164.6880/168.0596~~ * 100 = 97.93

DB 5/31/07

NOAH Cart #: _____

File #: _____

BFB Injection Date: 5/30/07 Logbook #: 1478
 BFB Injection Time: 1320
 BFB File ID: 8053001
 Tekmar Purge Flow: 16.3 ml/min
 Vacuum: 8.1 x 10⁻⁶
 IS/Std #: 1443-64 Exp. Date: 7/30/07
 BCM 441133
 1,4-DFB 1992312
 CB-d5 1475337
 Verified CCVIS vs ICAL mid-point (-40%D) 279
 initials

Calculation Check:

ppbv of compound = $\frac{\text{Area}_{\text{sample}}}{\text{Area}_{\text{std}}} \times \text{Conc.}_{\text{std}} \times \text{RRF}$ = $\left(\frac{\text{DB } 5/31/07}{\text{DB } 5/31/07} \right) \times \left(\frac{\text{DB } 5/31/07}{\text{DB } 5/31/07} \right)$

Reported Result _____

File ID: _____
 Compound: _____
 Initials: _____

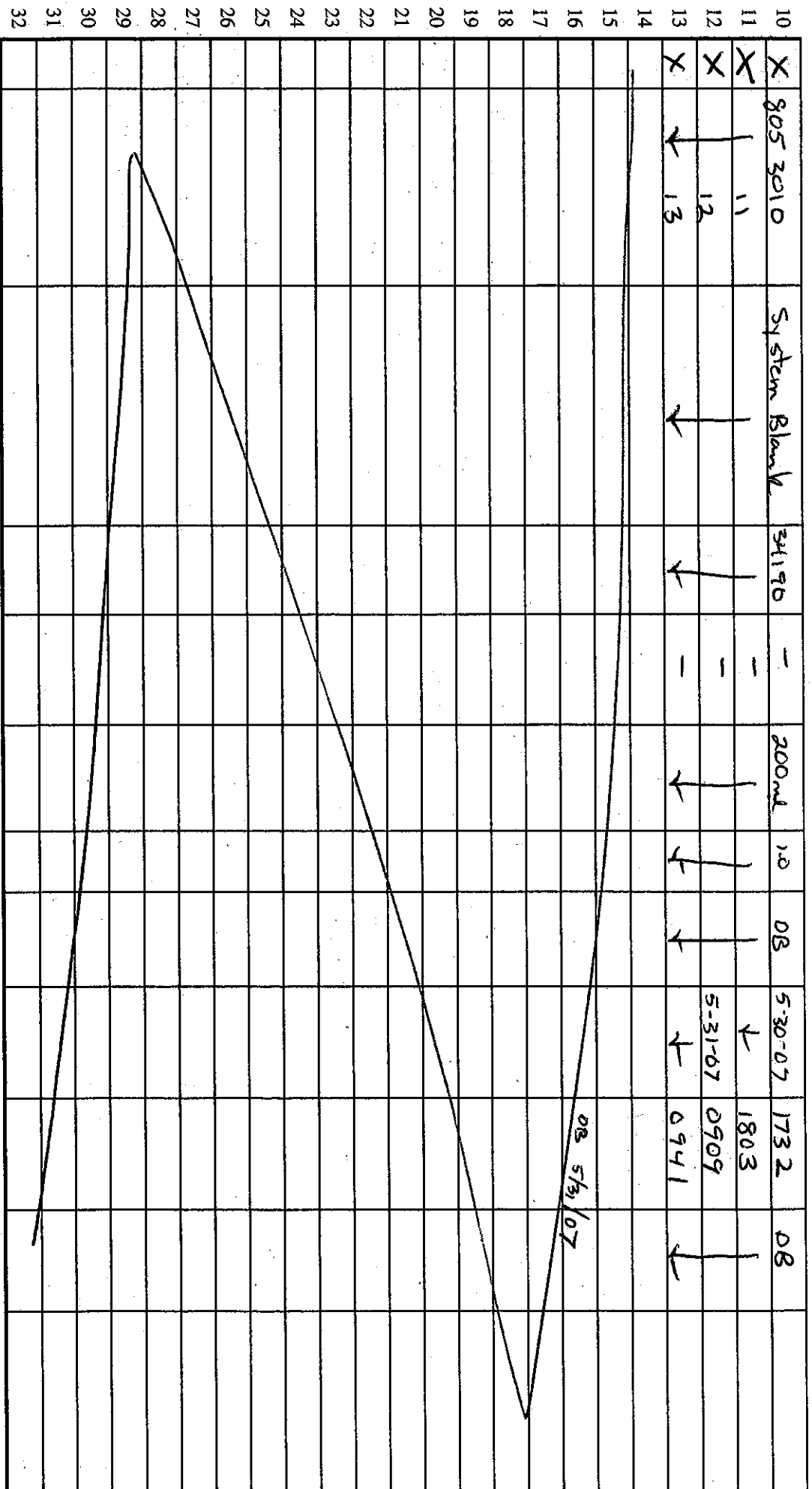
File #	Sample / Client Name	Can #	Pressure	Ampl Loaded	DF	Loader Init.	Date Analyzed	Time Analyzed	Review Init.	Comments
1	8053001	843-2981	50mg	200ul	10	DB	5-30-07	1320	DB	
2	X 8053002	34190	-	200ul				1344		
3	ICAR Level 1	1187289	0.2ppbv	0.2ul				1412		
4	04		0.5ppbv	0.5ul				1439		
5	05		20ppbv	2.0ul				1507		
6	06		25ppbv	2.5ul				1535		
7	07		50ppbv	5.0ul				1603		
8	08		100ppbv	10.0ul				1631		
9	09		200ppbv	20.0ul				1702		

Signature

Steve Butson

Date

5/31/07



Comments: 1st Flow Meter: OSE 2760 exp 8/19/07

220 ml/min → 25 ml/min

Flow meter: AA 9506172

Signature Laura Boston

Date 5/31/07

Revision 05/2005
Page 300

ION ABUNDANCE CRITERIA % REL. ABUNDANCE

m/z	REL. ABUNDANCE
50	15.0 - 40.0% of mass 95
75	30.0 - 60.0% of mass 95
95	Base peak, 100.00% relative abundance
96	5.0 - 9.0% of mass 95
173	Less than 2.0% of mass 174
174	Greater than 50.0% of mass 95
175	5.0 - 9.0% of mass 174
176	Greater than 95.0% but less than 101.0% of mass 174
177	5.0 - 9.0% of mass 176

BFB Injection Date: 5/31/07

BFB Injection Time: 1648

BFB File ID: 8053101

Tekmar Purge Flow: [Signature]

Vacuum: 0.5/31/07

IS/S Std #: 1443-64 Exp. Date: 7/30/07

BCM: 510024

1,4-DFB: 2335946

CB-d5: 1717992

Verified CCV IS vs ICAL mid-point (-40% D) [Signature]

NOAH Cart #: 8/14 File #: F052304/5052205

Verify 176/174 m/z Ratio: 126/176 / 1305/100 = 109 = 0.2725

Calculation Check:

$$\text{ppbv of compound} = \frac{\text{Area}_{\text{sample}}}{\text{Area}_{\text{std}}} \times \frac{\text{Conc.}_{\text{std}}}{\text{Conc.}_{\text{sample}}} \times \text{RRF}$$

$$= \left(\frac{1947437}{2335966} \right) \times \left(\frac{25}{0.80549} \right) \times 2235966$$

Reported Result: 24.081

File ID: 8053102
 Compound: Tbl-d8
 Initials: [Signature]

File #	Sample/Client Name	Conc. is	RRF	Area Sample	Area Std	ppbv of compound
8053101	BFB Time Check	7961				
02	CEV 4 1487-289	200ppb	SD ppb	5D vial	1.00	OR
03	LES * 1487-225	13693	Handsd	200 vial	1.00	OR
04	Lab Blank					
05	Lab Blank					
06	0705480-02A	1883	25" H ₂ O - 15psi		2.20	OR
07	07	2105	10" H ₂ O - 15psi		2.02	OR
08	08	5025			2.02	OR
09	09	-04AA	30psi	5D vial	8.36	OR

Signature: [Signature]

Date: 5/31/07

Revision 05/2005
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MSD-8

Logbook #: 1478

ION ABUNDANCE CRITERIA

m/z	REL. ABUNDANCE
50	15.0 - 40.0% of mass 95
75	30.0 - 60.0% of mass 95
95	Base peak, 100.00% relative abundance
96	5.0 - 9.0% of mass 95
173	Less than 2.0% of mass 174
174	Greater than 50.0% of mass 95
175	5.0 - 9.0% of mass 174
176	Greater than 95.0% but less than 101.0% of mass 174
177	5.0 - 9.0% of mass 176

BFB Injection Date: 1/21/02

BFB Injection Time: 0944

BFB File ID: 80e0701

Tekmar Purge Flow: 2 at 61362

Vacuum: 2 at 61362

IS/S Std #:	1493-64	Exp. Date:	2/25/02
BCM	432026		
1,4-DFB	1441557		
CB-d5	1439615		

Verified CCV IS vs ICAL mid-point (-40% D) DL

Verify 176/174 m/z Ratio: 123.806 / 126.9760 = 100 = 97.50%

NOAH Cart #:

File #:

Calculation Check:

ppbv of compound = $\frac{\text{Area}_{\text{sample}}}{\text{Areas}} \times \text{Conc.}_{\text{is}} \times \text{RRF}$

$= \frac{(1460776)}{(1441557)} \times (25) = 24.208$

Reported Result 24.728

File ID:	80e0702
Compound:	T61-28
Initials:	DL

File #	Sample / Client Name	Cart #	Pressure	Ampl. Loaded	DF	Loader Int.	Date Analyzed	Time Analyzed	Review Int.	Comments
1	BFB Tone Check	2481	50mg	2µl	100	DL	1/21/02	0944	DL	
2	DL	200	50µl	50µl				1001	DL	
3	DL	200	50µl	50µl				1029	DL	
4	DL	1493-64	2µl	2µl				1109	DL	splide
5	DL	5	50µl	50µl				1137	DL	
6	DL	7	200µl	200µl				1208	DL	
7	DL	131023	Humid	200µl	DL			1326	DL	
8	DL	DL	Lab Blank	1				1356	DL	
9	DL	DL	DL	DL						

Signature

DL

Date

1/21/02

Initial Calibration Narrative

A seven-point initial calibration was analyzed on MSD-8 on May 30, 2007.

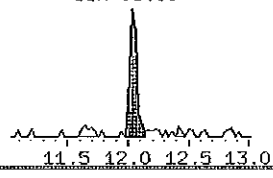
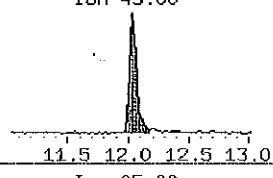
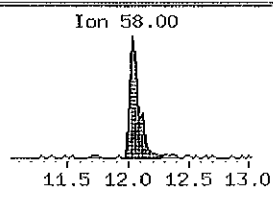
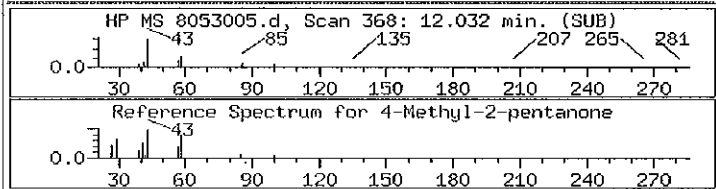
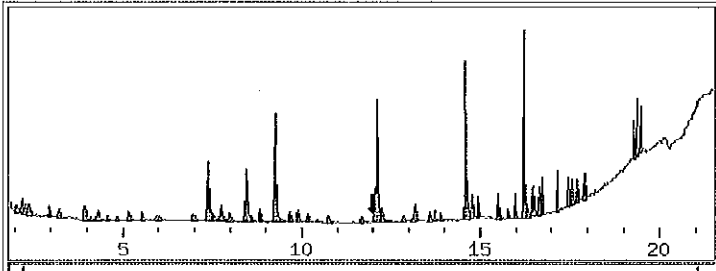
The following compounds used 0.2 as the lowest calibration concentration:
Benzene, Chloroform, Styrene and Cumene.

before

File Security Edit Display Process Spectra Help

Sample: ICAL Type: CALIB_3 Inj.Date: 30-MAY-2007 15:07

- + 67 Tetrahydrofural
- + 70 Chloroform
- + 73 Cyclohexane
- + 75 1,1,1-Trichloro
- + 77 Carbon Tetrach.
- + 81 Benzene
- + 80 2,2,4-Trimethy.
- + 83 1,2-Dichloroeth.
- + 85 Heptane
- + 94 Trichloroethen.
- + 95 Methyl Cyclohe.
- + 97 1,2-Dichloropr.
- + 98 1,4-Dioxane
- + 100 Bromodichlorom.
- + 102 cis-1,3-Dichlo
- + 103 4-Methyl-2-pentanone**
- + 105 Toluene
- + 108 trans-1,3-Dich.
- + 110 1,1,2-Trichloro
- + 112 Tetrachloroeth.
- + 114 2-Hexanone
- + 116 Dibromochlorom.
- + 117 1,2-Dibromoeth.
- + 126 Chlorobenzene
- + 129 Ethyl Benzene




HP MS 8053005.d, Scan 368: 12.032 min. (SUB)

Reference Spectrum for 4-Methyl-2-pentanone

Hit#	RT(min)	Response	Amount	Conc	Ratio	Flags	Report:
1	11.645	2880			234		
2	12.032	63447	2.526	2.526	100		
	12.032	123002			194		
	12.032	21458			34		
3	12.364	2625	0.1045	0.1045	100	T*	

8053005.d

Jun 1 2007



Sample / Initial	6/1107 DL
Poor Integration	
Split Peak	✓
Peak Tailing	
Background Subtraction	
Zoom In	
Missed Peak	

after
Ne 6/1107

File Security Edit Display Process Spectra Help

Sample: ICAL Type: CALIB_3 Inj.Date: 30-MAY-2007 15:07

- + 67 Tetrahydrofuran
- + 70 Chloroform
- + 73 Cyclohexane
- + 75 1,1,1-Trichloroethane
- + 77 Carbon Tetrachloride
- + 81 Benzene
- + 80 2,2,4-Trimethylpentane
- + 83 1,2-Dichloroethane
- + 85 Heptane
- + 94 Trichloroethylene
- + 95 Methyl Cyclohexane
- + 97 1,2-Dichloropropane
- + 98 1,4-Dioxane
- + 100 Bromodichloromethane
- + 102 cis-1,3-Dichloropropane
- + 103 4-Methyl-2-pentanone**
- + 105 Toluene
- + 108 trans-1,3-Dichloropropane
- + 110 1,1,2-Trichloroethane
- + 112 Tetrachloroethane
- + 114 2-Hexanone
- + 116 Dibromochloromethane
- + 117 1,2-Dibromoethane
- + 126 Chlorobenzene
- + 129 Ethyl Benzene

Ion 58.00

Ion 43.00

Ion 85.00

HP MS 8053005.d, Scan 368: 12.032 min. (SUB)

Reference Spectrum for 4-Methyl-2-pentanone

Hit#	RT(min)	Response	Amount	Conc	Ratio	Flags	Report:
1	12.032	49522	2.067	2.067	100	M	
	12.032	123001			248		
	12.032	21457			43		

- Mark 4-Methyl-2-pentanone Undetected.

8053005.d

Air Toxics Ltd.
 Modified EPA Methods TO-14A/TO-15
 Internal Standard and Associated Target Compounds and Surrogates

Bromochloromethane
Target Compounds:
Freon 12
Freon 114
Chloromethane
Vinyl Chloride
1,3-Butadiene
Bromomethane
Chloroethane
Freon 11
Ethanol
Freon 113
1,1-Dichloroethene
Acetone
2-Propanol
Carbon Disulfide
3-Chloropropene
Methylene Chloride
Methyl tert-butyl ether
trans-1,2-Dichloroethene
Hexane
1,1-Dichloroethane
2-Butanone (Methyl Ethyl Ketone)
cis-1,2-Dichloroethene
Tetrahydrofuran
Chloroform
1,1,1-Trichloroethane
Cyclohexane
Carbon Tetrachloride
2,2,4-Trimethylpentane
Surrogates:
1,2-Dichloroethane-d4

1,4-Difluorobenzene
Target Compounds:
Benzene
1,2-Dichloroethane
Heptane
Trichloroethene
1,2-Dichloropropane
1,4-Dioxane
Bromodichloromethane
cis-1,3-Dichloropropene
4-Methyl-2-pentanone
Toluene
Surrogates:
Toluene-d8

Chlorobenzene-d5
Target Compounds:
trans-1,3-Dichloropropene
1,1,2-Trichloroethane
Tetrachloroethene
2-Hexanone
Dibromochloromethane
1,2-Dibromoethane (EDB)
Chlorobenzene
Ethyl Benzene
m,p-Xylene
o-Xylene
Styrene
Bromoform
Cumene
1,1,2,2-Tetrachloroethane
Propylbenzene
4-Ethyltoluene
1,3,5-Trimethylbenzene
1,2,4-Trimethylbenzene
1,3-Dichlorobenzene
1,4-Dichlorobenzene
alpha-Chlorotoluene
1,2-Dichlorobenzene
1,2,4-Trichlorobenzene
Hexachlorobutadiene
Surrogates:
Bromofluorobenzene

Report Date: 01-Jun-2007 10:48

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-31may.b/8053103.d
 Lab Smp Id: LCS-1 Client Smp ID: LCS-1
 Inj Date : 31-MAY-2007 11:48
 Operator : JG Inst ID: msd8.i
 Smp Info : 50ml #1487-275
 Misc Info : 200ppbv-50ppbv
 Comment :
 Method : /chem/msd8.i/8-31may.b/t14q530a.m
 Meth Date : 01-Jun-2007 10:48 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 17:02 Cal File: 8053009.d
 Als bottle: 1 QC Sample: LCS
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS								
RT	EXP RT	(REL RT)	MASS	RESPONSE		TARGET RANGE	RATIO	
				(PPBV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====

* 68	Bromochloromethane					CAS #: 74-97-5		
7.387	7.415	(1.000)	130	440756	25.0000	80.00- 120.00	100.00	
7.387	7.415	(1.000)	128	346888		48.60- 108.60	78.70	
7.387	7.387	(1.000)	49	625716		114.98- 174.98	141.96	

* 88	1,4-Difluorobenzene					CAS #: 540-36-3		
9.267	9.267	(1.000)	114	2010798	25.0000	80.00- 120.00	100.00	
9.267	9.267	(1.000)	88	311662		0.00- 45.16	15.50	

* 125	Chlorobenzene-d5					CAS #: 3114-55-4		
14.576	14.576	(1.000)	117	1506758	25.0000	80.00- 120.00	100.00	
14.576	14.576	(1.000)	82	856520		0.00- 30.00	56.85	

\$ 82	1,2-Dichloroethane-d4					CAS #: 17060-07-0		
8.465	8.465	(1.146)	65	572200	24.8051	24.805 80.00- 120.00	100.00	
8.465	8.465	(1.146)	67	351230		0.00- 30.00	61.38	

\$ 104	Toluene-d8					CAS #: 2037-26-5		
12.115	12.115	(1.307)	98	1737699	24.9622	24.962 80.00- 120.00	100.00	
12.115	12.115	(1.307)	70	186801		0.00- 30.00	10.75	

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT (REL RT)	MASS	RESPONSE	(PPEV)	(PPBV)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====

\$ 104 Toluene-d8 (continued)

12.115	12.115 (1.307)	100	1549990			0.00- 30.00	89.20
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\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207	16.207 (1.112)	174	897285	25.0287	25.029	80.00- 120.00	100.00
16.207	16.207 (1.112)	95	1158459			105.65- 165.65	129.11
16.207	16.207 (1.112)	176	894688			66.34- 126.34	99.71

4 Dichlorodifluoromethane/Fr12

CAS #: 75-71-8

2.050	2.078 (0.278)	85	2778941	46.0212	46.021	80.00- 120.00	100.00
2.050	2.078 (0.278)	87	891336			0.00- 30.00	32.07

6 Freon 114

CAS #: 76-14-2

2.161	2.189 (0.293)	135	2562967	46.9817	46.982	80.00- 120.00	100.00
2.161	2.189 (0.293)	137	809100			0.71- 60.71	31.57

8 Chloromethane

CAS #: 74-87-3

2.272	2.299 (0.308)	50	1366974	49.1562	49.156	80.00- 120.00	100.00
2.272	2.299 (0.308)	52	419326			0.00- 30.00	30.68

11 Vinyl Chloride

CAS #: 75-01-4

2.410	2.438 (0.326)	62	1506942	46.6352	46.635	80.00- 120.00	100.00
2.410	2.438 (0.326)	64	471709			0.00- 30.00	31.30

10 1,3-Butadiene

CAS #: 106-99-0

2.382	2.410 (0.322)	54	1138555	41.1922	41.192	80.00- 120.00	100.00
2.382	2.410 (0.322)	39	1152974			0.00- 30.00	101.27

13 Bromomethane

CAS #: 74-83-9

2.852	2.880 (0.386)	94	1047024	48.5003	48.500	80.00- 120.00	100.00
2.852	2.880 (0.386)	96	1003279			63.65- 123.65	95.82

16 Chloroethane

CAS #: 75-00-3

2.963	2.991 (0.401)	64	798701	46.7036	46.704	80.00- 120.00	100.00
2.935	2.963 (0.397)	49	208062			0.00- 30.00	26.05
2.963	2.991 (0.401)	66	241751			0.00- 30.00	30.27

18 Trichlorofluoromethane/Fr11

CAS #: 75-69-4

3.212	3.239 (0.435)	101	3306964	48.1826	48.183	80.00- 120.00	100.00
3.212	3.239 (0.435)	103	2152698			34.50- 94.50	65.10

23 Ethanol

CAS #: 64-17-5

3.516	3.544 (0.476)	45	588478	53.6942	53.694	80.00- 120.00	100.00
3.516	3.544 (0.476)	43	117595			0.00- 30.00	19.98
3.516	3.544 (0.476)	46	241557			0.00- 30.00	41.05

CONCENTRATIONS

ON-COL FINAL

RT EXP RT (REL RT) MASS RESPONSE (PPBV) (PPBV) TARGET RANGE RATIO
 == == ===== == ===== ===== =====

28 Freon 113 CAS #: 76-13-1
 3.931 3.958 (0.532) 151 2312964 50.9343 50.934 80.00- 120.00 100.00
 3.931 3.958 (0.532) 153 1528932 34.32- 94.32 66.10
 3.931 3.958 (0.532) 101 2826353 85.50- 145.50 122.20

29 1,1-Dichloroethene CAS #: 75-35-4
 3.958 3.986 (0.536) 61 2385056 52.8867 52.887 80.00- 120.00 100.00
 3.958 3.986 (0.536) 96 1431121 29.81- 89.81 60.00
 3.958 3.986 (0.536) 98 913818 9.00- 69.00 38.31

30 Acetone CAS #: 67-64-1
 4.124 4.124 (0.558) 58 750144 49.6648 49.665 80.00- 120.00 100.00
 4.124 4.124 (0.558) 43 2233113 0.00- 30.00 297.69

34 2-Propanol CAS #: 67-63-0
 4.318 4.318 (0.584) 45 2687206 49.2062 49.206 80.00- 120.00 100.00
 4.318 4.318 (0.584) 43 509221 0.00- 30.00 18.95
 4.318 4.318 (0.584) 59 100187 0.00- 30.00 3.73

33 Carbon Disulfide CAS #: 75-15-0
 4.290 4.290 (0.581) 76 4015505 47.5101 47.510 80.00- 120.00 100.00

37 3-Chloropropene CAS #: 107-05-1
 4.566 4.594 (0.618) 76 692936 51.3911 51.391 80.00- 120.00 100.00
 4.566 4.594 (0.618) 41 2090707 0.00- 30.00 301.72

40 Methylene Chloride CAS #: 75-09-2
 4.815 4.815 (0.652) 49 1736664 51.4364 51.436 80.00- 120.00 100.00
 4.815 4.843 (0.652) 84 1236611 40.47- 100.47 71.21
 4.815 4.815 (0.652) 51 518270 0.00- 30.00 29.84

43 MTBE CAS #: 1634-04-4
 5.147 5.175 (0.697) 73 2334781 48.6230 48.623 80.00- 120.00 100.00
 5.147 5.175 (0.697) 57 526586 0.00- 52.63 22.55
 5.147 5.175 (0.697) 41 546280 0.00- 30.00 23.40

45 trans-1,2-Dichloroethene CAS #: 156-60-5
 5.175 5.203 (0.701) 96 1480166 46.3832 46.383 80.00- 120.00 100.00
 5.175 5.203 (0.701) 61 2223377 119.47- 179.47 150.21
 5.175 5.203 (0.701) 98 951148 0.00- 30.00 64.26

46 Hexane CAS #: 110-54-3
 5.534 5.534 (0.749) 57 2449048 50.6003 50.600 80.00- 120.00 100.00
 5.534 5.534 (0.749) 43 1575746 0.00- 30.00 64.34
 5.534 5.534 (0.749) 86 364966 0.00- 30.00 14.90

CONCENTRATIONS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	ON-COL (PPEV)	FINAL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====	=====	

54	1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	2639497	50.7389	50.739	80.00-	120.00	100.00	
5.949	5.949	(0.805)	65	834971			1.86-	61.86	31.63	

65	2-Butanone					CAS #: 78-93-3				
7.027	7.027	(0.951)	72	692091	47.3821	47.382	80.00-	120.00	100.00	
7.027	7.027	(0.951)	43	2937806			409.65-	469.65	424.48	
7.027	7.027	(0.951)	57	229143			0.00-	30.00	33.11	

64	cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.972	6.972	(0.944)	61	1927506	46.8160	46.816	80.00-	120.00	100.00	
6.972	6.972	(0.944)	96	1448922			43.20-	103.20	75.17	
6.972	6.972	(0.944)	98	903977			17.85-	77.85	46.90	

67	Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.415	(1.000)	42	1762470	46.4222	46.422	80.00-	120.00	100.00	
7.387	7.415	(1.000)	71	616972			5.48-	65.48	35.01	
7.387	7.415	(1.000)	72	683813			0.00-	30.00	38.80	

70	Chloroform					CAS #: 67-66-3				
7.525	7.553	(1.019)	83	2568490	45.8768	45.877	80.00-	120.00	100.00	
7.525	7.553	(1.019)	85	1587019			32.67-	92.67	61.79	

75	1,1,1-Trichloroethane					CAS #: 71-55-6				
7.774	7.774	(1.052)	97	2636611	48.3678	48.368	80.00-	120.00	100.00	
7.774	7.774	(1.052)	99	1692005			34.55-	94.55	64.17	

73	Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	2011710	46.6032	46.603	80.00-	120.00	100.00	
7.746	7.746	(1.049)	56	2573375			96.25-	156.25	127.92	
7.746	7.746	(1.049)	41	1297745			32.28-	92.28	64.51	

77	Carbon Tetrachloride					CAS #: 56-23-5				
7.995	8.023	(1.082)	119	2542777	51.2619	51.262	80.00-	120.00	100.00	
7.995	8.023	(1.082)	117	2618298			73.30-	133.30	102.97	

80	2,2,4-Trimethylpentane					CAS #: 540-84-1				
8.465	8.465	(1.146)	57	6011320	46.6484	46.648	80.00-	120.00	100.00	
8.465	8.465	(1.146)	56	1918609			0.00-	30.00	31.92	
8.465	8.465	(1.146)	41	1389836			0.00-	30.00	23.12	

81	Benzene					CAS #: 71-43-2				
8.437	8.438	(0.910)	78	4287698	45.2681	45.268	80.00-	120.00	100.00	
8.437	8.438	(0.910)	77	959130			0.00-	30.00	22.37	

83	1,2-Dichloroethane					CAS #: 107-06-2				
8.603	8.603	(0.928)	62	1763231	47.8421	47.842	80.00-	120.00	100.00	

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	ON-COL (PPEV)	FINAL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
83 1,2-Dichloroethane (continued)									
8.603	8.603	(0.928)	64	570080			0.00- 30.00	32.33	

85 Heptane CAS #: 142-82-5									
8.852	8.852	(0.955)	100	422399	42.9697	42.970	80.00- 120.00	100.00	
8.852	8.852	(0.955)	43	2420809			0.00- 30.00	573.11	
8.852	8.852	(0.955)	71	1375374			0.00- 30.00	325.61	

94 Trichloroethene CAS #: 79-01-6									
9.682	9.682	(1.045)	95	1722090	47.8377	47.838	80.00- 120.00	100.00	
9.682	9.682	(1.045)	130	1803299			78.57- 138.57	104.72	
9.682	9.682	(1.045)	97	1080675			34.08- 94.08	62.75	

97 1,2-Dichloropropane CAS #: 78-87-5									
10.179	10.179	(1.098)	63	1484383	47.1518	47.152	80.00- 120.00	100.00	
10.179	10.179	(1.098)	62	1055399			42.26- 102.26	71.10	
10.179	10.179	(1.098)	41	846166			26.77- 86.77	57.00	

98 1,4-Dioxane CAS #: 123-91-1									
10.428	10.428	(1.125)	88	953301	49.6479	49.648	80.00- 120.00	100.00	
10.428	10.428	(1.125)	58	672923			41.45- 101.45	70.59	
10.428	10.428	(1.125)	57	202353			0.00- 30.00	21.23	

100 Bromodichloromethane CAS #: 75-27-4									
10.732	10.732	(1.158)	83	2523493	47.9946	47.995	80.00- 120.00	100.00	
10.732	10.732	(1.158)	85	1562917			31.75- 91.75	61.93	

102 cis-1,3-Dichloropropene CAS #: 10061-01-5									
11.672	11.673	(1.260)	75	2069212	47.2977	47.298	80.00- 120.00	100.00	
11.672	11.673	(1.260)	77	655311			1.36- 61.36	31.67	
11.672	11.673	(1.260)	39	1004071			17.70- 77.70	48.52	

103 4-Methyl-2-pentanone CAS #: 108-10-1									
12.032	12.032	(1.298)	58	1179076	49.8183	49.818	80.00- 120.00	100.00	
12.032	12.032	(1.298)	43	2851980			0.00- 30.00	241.88	
12.032	12.032	(1.298)	85	496153			0.00- 30.00	42.08	

105 Toluene CAS #: 108-88-3									
12.253	12.253	(1.322)	91	4606596	49.8972	49.897	80.00- 120.00	100.00	
12.253	12.253	(1.322)	92	2757068			29.69- 89.69	59.85	

108 trans-1,3-Dichloropropene CAS #: 10061-02-6									
12.834	12.861	(0.880)	75	2120788	48.5501	48.550	80.00- 120.00	100.00	
12.834	12.861	(0.880)	77	642673			0.08- 60.08	30.30	
12.834	12.834	(0.880)	39	956206			15.97- 75.97	45.09	

CONCENTRATIONS										
RT	EXP RT	(REL RT)	MASS	ON-COL		FINAL	TARGET RANGE	RATIO		
				RESPONSE	(PPEV)	(PPBV)				
==	=====	=====	=====	=====	=====	=====	=====	=====	=====	

110	1,1,2-Trichloroethane					CAS #:	79-00-5			
13.138	13.138	(0.901)	97	1450072	45.2633	45.263	80.00-	120.00	100.00	
13.138	13.138	(0.901)	99	903189			29.85-	89.85	62.29	
13.138	13.138	(0.901)	83	1268499			54.93-	114.93	87.48	

112	Tetrachloroethene					CAS #:	127-18-4			
13.193	13.193	(0.905)	166	2052640	47.3396	47.340	80.00-	120.00	100.00	
13.193	13.193	(0.905)	129	1598570			45.08-	105.08	77.88	
13.193	13.193	(0.905)	131	1573037			44.22-	104.22	76.63	

114	2-Hexanone					CAS #:	591-78-6			
13.580	13.580	(0.932)	58	1578878	49.7385	49.738	80.00-	120.00	100.00	
13.580	13.580	(0.932)	43	2839029			148.62-	208.62	179.81	
13.580	13.580	(0.932)	100	328947			0.00-	30.00	20.83	

116	Dibromochloromethane					CAS #:	124-48-1			
13.718	13.719	(0.941)	129	2441815	48.6342	48.634	80.00-	120.00	100.00	
13.718	13.719	(0.941)	127	1862028			0.00-	30.00	76.26	

117	1,2-Dibromoethane					CAS #:	106-93-4			
13.884	13.884	(0.953)	107	2264782	46.3953	46.395	80.00-	120.00	100.00	
13.884	13.884	(0.953)	109	2141958			63.81-	123.81	94.58	

126	Chlorobenzene					CAS #:	108-90-7			
14.603	14.631	(1.002)	112	3600449	46.5367	46.537	80.00-	120.00	100.00	
14.603	14.631	(1.002)	114	1156467			1.84-	61.84	32.12	
14.603	14.603	(1.002)	77	2040419			26.78-	86.78	56.67	

129	Ethyl Benzene					CAS #:	100-41-4			
14.769	14.769	(1.013)	106	1875253	45.8007	45.801	80.00-	120.00	100.00	
14.769	14.769	(1.013)	91	5813094			0.00-	30.00	309.99	

130	m,p-Xylene					CAS #:	108-38-3			
14.935	14.935	(1.025)	106	2373682	47.5756	47.576	80.00-	120.00	100.00	
14.935	14.935	(1.025)	91	4516302			0.00-	30.00	190.27	

132	o-Xylene					CAS #:	95-47-6			
15.488	15.488	(1.063)	106	2318082	46.0425	46.042	80.00-	120.00	100.00	
15.488	15.488	(1.063)	91	4803654			178.74-	238.74	207.23	

134	Styrene					CAS #:	100-42-5			
15.516	15.516	(1.064)	104	3405235	47.9066	47.907	80.00-	120.00	100.00	
15.516	15.516	(1.064)	78	1716218			20.27-	80.27	50.40	

135	Bromoform					CAS #:	75-25-2			
15.764	15.765	(1.082)	173	2241846	50.7630	50.763	80.00-	120.00	100.00	

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	ON-COL (PPEV)	FINAL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
135 Bromoform (continued)									
15.764	15.765	(1.082)	171	1157469			20.67- 80.67	51.63	

144 1,1,2,2-Tetrachloroethane CAS #: 79-34-5									
16.456	16.456	(1.129)	83	3338269	45.3401	45.340	80.00- 120.00	100.00	
16.456	16.456	(1.129)	85	2060617			31.59- 91.59	61.73	

147 4-Ethyltoluene CAS #: 622-96-8									
16.649	16.649	(1.142)	105	6752548	49.9730	49.973	80.00- 120.00	100.00	
16.649	16.649	(1.142)	120	2056264			0.00- 59.85	30.45	

148 1,3,5-Trimethylbenzene CAS #: 108-67-8									
16.732	16.732	(1.148)	105	6260050	48.1979	48.198	80.00- 120.00	100.00	
16.732	16.732	(1.148)	120	3095852			0.00- 30.00	49.45	

153 1,2,4-Trimethylbenzene CAS #: 95-63-6									
17.147	17.147	(1.176)	105	6608084	50.5445	50.544	80.00- 120.00	100.00	
17.147	17.147	(1.176)	120	2956336			15.22- 75.22	44.74	

156 1,3-Dichlorobenzene CAS #: 541-73-1									
17.451	17.451	(1.197)	146	4297022	50.4188	50.419	80.00- 120.00	100.00	
17.451	17.451	(1.197)	148	2703733			0.00- 30.00	62.92	
17.451	17.451	(1.197)	111	1752089			0.00- 30.00	40.77	

157 1,4-Dichlorobenzene CAS #: 106-46-7									
17.562	17.562	(1.205)	146	3673574	45.4817	45.482	80.00- 120.00	100.00	
17.562	17.562	(1.205)	148	2397507			0.00- 30.00	65.26	
17.562	17.562	(1.205)	111	1301488			0.00- 30.00	35.43	

158 alpha-Chlorotoluene CAS #: 100-44-7									
17.700	17.700	(1.214)	91	6425001	54.7247	54.725	80.00- 120.00	100.00	
17.700	17.700	(1.214)	126	1322597			0.00- 30.00	20.59	

161 1,2-Dichlorobenzene CAS #: 95-50-1									
17.921	17.921	(1.230)	146	3730128	45.4352	45.435	80.00- 120.00	100.00	
17.921	17.921	(1.230)	148	2410442			33.37- 93.37	64.62	
17.921	17.921	(1.230)	111	1412557			7.65- 67.65	37.87	

167 1,2,4-Trichlorobenzene CAS #: 120-82-1									
19.276	19.276	(1.322)	180	3988477	48.0685	48.068	80.00- 120.00	100.00	
19.276	19.276	(1.322)	182	3802153			64.68- 124.68	95.33	

168 Hexachlorobutadiene CAS #: 87-68-3									
19.359	19.359	(1.328)	225	2159382	45.3397	45.340	80.00- 120.00	100.00	
19.359	19.359	(1.328)	223	1379410			33.22- 93.22	63.88	

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPEV)	FINAL	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

145 Propylbenzene						CAS #:	103-65-1		
16.483	16.484	(1.131)	91	8066062	51.1778	51.178	80.00-	120.00	100.00
16.483	16.484	(1.131)	120	1852431			0.00-	30.00	22.97
16.483	16.484	(1.131)	105	285850			0.00-	30.00	3.54

137 Cumene						CAS #:	98-82-8		
15.958	15.958	(1.095)	105	6813817	47.9676	47.968	80.00-	120.00	100.00
15.958	15.986	(1.095)	120	1785636			0.00-	30.00	26.21
15.958	15.958	(1.095)	51	636171			0.00-	30.00	9.34

169 Naphthalene						CAS #:	91-20-3		
19.470	19.470	(1.336)	128	9185790	52.4933	52.493	80.00-	120.00	100.00
19.470	19.470	(1.336)	127	1071085			0.00-	30.00	11.66

9 Butane						CAS #:	106-97-8		
2.327	2.355	(0.315)	58	350243	48.2264	48.226	70.00-	130.00	100.00
2.327	2.355	(0.315)	43	2575697			0.00-	30.00	735.40

15 Isopentane						CAS #:	78-78-4		
2.963	2.963	(0.401)	43	1966389	48.7352	48.735	70.00-	130.00	100.00
2.963	2.963	(0.401)	57	1333134			0.00-	30.00	67.80
2.963	2.963	(0.401)	72	143213			0.00-	30.00	7.28

95 Methyl Cyclohexane						CAS #:	108-87-2		
9.903	9.903	(1.341)	83	2603625	48.4146	48.414	70.00-	130.00	100.00
9.903	9.903	(1.341)	98	1166714			0.00-	30.00	44.81
9.903	9.903	(1.341)	55	2051809			0.00-	30.00	78.81

Report Date: 01-Jun-2007 10:48

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 31-MAY-2007

Lab File ID: 8053103.d

Calibration Time: 11:20

Lab Smp Id: LCS-1

Client Smp ID: LCS-1

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: JG

Method File: /chem/msd8.i/8-31may.b/t14q530a.m

Misc Info: 200ppbv-50ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	510079	306047	714111	440756	-13.59
88 1,4-Difluorobenze	2335966	1401580	3270352	2010798	-13.92
125 Chlorobenzene-d5	1717992	1030795	2405189	1506758	-12.30

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.41	7.08	7.74	7.39	-0.37
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Air Toxics Ltd.

RECOVERY REPORT

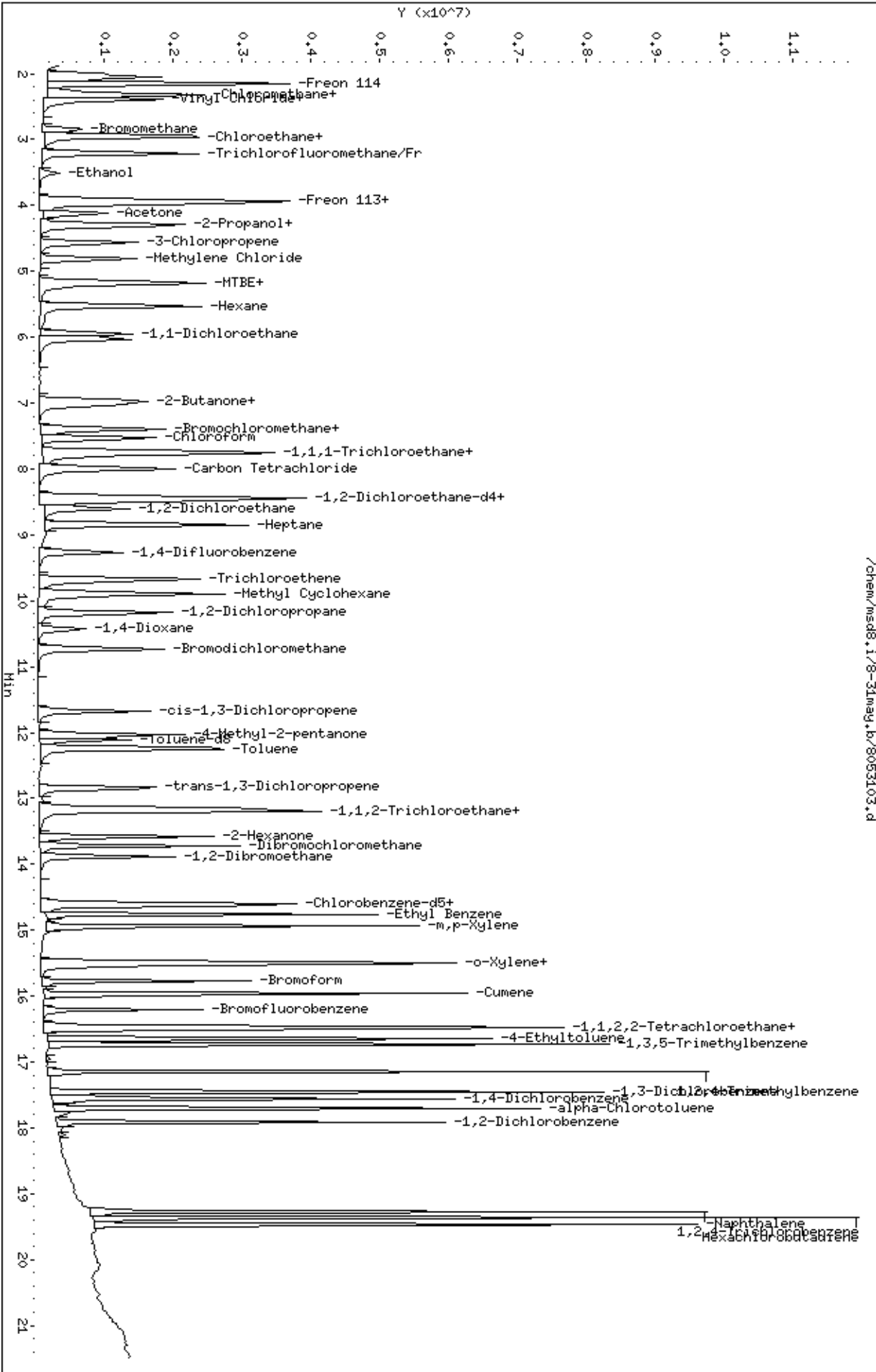
Client Name: Client SDG: 8-31may
 Sample Matrix: GAS Fraction: VOA
 Lab Smp Id: LCS-1 Client Smp ID: LCS-1
 Level: LOW Operator: JG
 Data Type: MS DATA SampleType: LCS
 SpikeList File: Spectra.spk Quant Type: ISTD
 Sublist File: AT04+ENSR.sub
 Method File: /chem/msd8.i/8-31may.b/t14q530a.m
 Misc Info: 200ppbv-50ppbv

SPIKE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
134 Styrene	50.000	47.907	95.81	70-130
108 trans-1,3-Dichloro	50.000	48.550	97.10	70-130
4 Dichlorodifluorome	50.000	46.021	92.04	70-130
6 Freon 114	50.000	46.982	93.96	70-130
8 Chloromethane	50.000	49.156	98.31	70-130
11 Vinyl Chloride	50.000	46.635	93.27	70-130
10 1,3-Butadiene	50.000	41.192	82.38	60-140
13 Bromomethane	50.000	48.500	97.00	70-130
16 Chloroethane	50.000	46.704	93.41	70-130
18 Trichlorofluoromet	50.000	48.183	96.37	70-130
23 Ethanol	50.000	53.694	107.39	60-140
28 Freon 113	50.000	50.934	101.87	70-130
29 1,1-Dichloroethene	50.000	52.887	105.77	70-130
30 Acetone	50.000	49.665	99.33	60-140
33 Carbon Disulfide	50.000	47.510	95.02	60-140
34 2-Propanol	50.000	49.206	98.41	60-140
40 Methylene Chloride	50.000	51.436	102.87	70-130
43 MTBE	50.000	48.623	97.25	60-140
45 trans-1,2-Dichloro	50.000	46.383	92.77	60-140
46 Hexane	50.000	50.600	101.20	60-140
54 1,1-Dichloroethane	50.000	50.739	101.48	70-130
64 cis-1,2-Dichloroet	50.000	46.816	93.63	70-130
65 2-Butanone	50.000	47.382	94.76	60-140
67 Tetrahydrofuran	50.000	46.422	92.84	60-140
70 Chloroform	50.000	45.877	91.75	70-130
73 Cyclohexane	50.000	46.603	93.21	60-140
75 1,1,1-Trichloroeth	50.000	48.368	96.74	70-130
77 Carbon Tetrachlori	50.000	51.262	102.52	70-130
81 Benzene	50.000	45.268	90.54	70-130
83 1,2-Dichloroethane	50.000	47.842	95.68	70-130
85 Heptane	50.000	42.970	85.94	60-140
94 Trichloroethene	50.000	47.838	95.68	70-130
97 1,2-Dichloropropan	50.000	47.152	94.30	70-130

Report Date: 01-Jun-2007 10:48

SPIKE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
98 1,4-Dioxane	50.000	49.648	99.30	60-140
100 Bromodichlorometha	50.000	47.995	95.99	60-140
102 cis-1,3-Dichloropr	50.000	47.298	94.60	70-130
103 4-Methyl-2-pentano	50.000	49.818	99.64	60-140
105 Toluene	50.000	49.897	99.79	70-130
110 1,1,2-Trichloroeth	50.000	45.263	90.53	70-130
112 Tetrachloroethene	50.000	47.340	94.68	70-130
114 2-Hexanone	50.000	49.738	99.48	60-140
116 Dibromochlorometha	50.000	48.634	97.27	60-140
117 1,2-Dibromoethane	50.000	46.395	92.79	70-130
126 Chlorobenzene	50.000	46.537	93.07	70-130
129 Ethyl Benzene	50.000	45.801	91.60	70-130
130 m,p-Xylene	50.000	47.576	95.15	70-130
132 o-Xylene	50.000	46.042	92.08	70-130
135 Bromoform	50.000	50.763	101.53	60-140
144 1,1,2,2-Tetrachlor	50.000	45.340	90.68	70-130
147 4-Ethyltoluene	50.000	49.973	99.95	60-140
148 1,3,5-Trimethylben	50.000	48.198	96.40	70-130
153 1,2,4-Trimethylben	50.000	50.544	101.09	70-130
156 1,3-Dichlorobenzen	50.000	50.419	100.84	70-130
157 1,4-Dichlorobenzen	50.000	45.482	90.96	70-130
158 alpha-Chlorotoluen	50.000	54.725	109.45	70-130
161 1,2-Dichlorobenzen	50.000	45.435	90.87	70-130
167 1,2,4-Trichloroben	50.000	48.068	96.14	70-130
168 Hexachlorobutadien	50.000	45.340	90.68	70-130
137 Cumene	50.000	47.968	95.94	60-140
145 Propylbenzene	50.000	51.178	102.36	60-140
37 3-Chloropropene	50.000	51.391	102.78	60-140
80 2,2,4-Trimethylpen	50.000	46.648	93.30	60-140
169 Naphthalene	50.000	52.493	104.99	60-140
9 Butane	50.000	48.226	96.45	70-130
15 Isopentane	50.000	48.735	97.47	70-130
95 Methyl Cyclohexane	50.000	48.414	96.83	70-130

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	24.805	99.22	70-130
\$ 104 Toluene-d8	25.000	24.962	99.85	70-130
\$ 140 Bromofluorobenzene	25.000	25.029	100.11	70-130



Report Date: 31-May-2007 14:52

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053003.d
 Lab Smp Id: ICAL Client Smp ID: Level 1
 Inj Date : 30-MAY-2007 14:12
 Operator : db Inst ID: msd8.i
 Smp Info : 0.2ml #1487-289
 Misc Info : 200ppbv-0.2ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 31-May-2007 14:52 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 14:12 Cal File: 8053003.d
 Als bottle: 1 Calibration Sample, Level: 1
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AFCEElow.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS								
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	ON-COL	TARGET RANGE	RATIO
==	=====	=====	====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5								
7.387	7.387	(1.000)	130	449663	25.0000		70.00- 130.00	100.00
7.387	7.387	(1.000)	128	347373			47.57- 107.57	77.25
7.387	7.387	(1.000)	49	653261			113.47- 173.47	145.28

* 88 1,4-Difluorobenzene CAS #: 540-36-3								
9.267	9.267	(1.000)	114	2011611	25.0000		70.00- 130.00	100.00
9.267	9.267	(1.000)	88	307684			0.00- 45.68	15.30

* 125 Chlorobenzene-d5 CAS #: 3114-55-4								
14.576	14.576	(1.000)	117	1511139	25.0000		70.00- 130.00	100.00
14.576	14.576	(1.000)	82	870029			0.00- 30.00	57.57

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0								
8.465	8.465	(1.146)	65	580550	25.0000	24.668	70.00- 130.00	100.00
8.465	8.465	(1.146)	67	304406			0.00- 30.00	52.43

\$ 104 Toluene-d8 CAS #: 2037-26-5								
12.115	12.115	(1.307)	98	1800242	25.0000	25.850	70.00- 130.00	100.00
12.115	12.115	(1.307)	70	162095			0.00- 30.00	9.00

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
\$ 104 Toluene-d8 (continued)										
12.115	12.115	(1.307)	100	1155287			0.00- 30.00	64.17		

\$ 140 Bromofluorobenzene										
						CAS #: 460-00-4				
16.207	16.207	(1.112)	174	863540	25.0000	24.018	70.00- 130.00	100.00		
16.207	16.207	(1.112)	95	1149176			102.16- 162.16	133.08		
16.207	16.207	(1.112)	176	834188			64.31- 124.31	96.60		

70 Chloroform										
						CAS #: 67-66-3				
7.525	7.525	(1.019)	83	16272	0.20000	0.3066	70.00- 130.00	100.00(a)		
7.525	7.525	(1.019)	85	9678			31.98- 91.98	59.48		

81 Benzene										
						CAS #: 71-43-2				
8.437	8.437	(0.910)	78	23610	0.20000	0.2598	70.00- 130.00	100.00(a)		
8.437	8.437	(0.910)	77	8272			0.00- 30.00	35.04		

134 Styrene										
						CAS #: 100-42-5				
15.516	15.516	(1.064)	104	13430	0.20000	0.1866	70.00- 130.00	100.00(a)		
15.516	15.516	(1.064)	78	8927			20.25- 80.25	66.47		

137 Cumene										
						CAS #: 98-82-8				
15.958	15.958	(1.095)	105	33835	0.20000	0.2452	70.00- 130.00	100.00(a)		
15.958	15.958	(1.095)	120	10282			0.00- 30.00	30.39		
15.958	15.958	(1.095)	51	4623			0.00- 30.00	13.66		

QC Flag Legend

a - Target compound detected but, quantitated amount
 Below Limit Of Quantitation(BLOQ).

Report Date: 31-May-2007 14:52

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053003.d

Calibration Time: 16:03

Lab Smp Id: ICAL

Client Smp ID: Level 1

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-0.2ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	441133	264680	617586	449663	1.93
88 1,4-Difluorobenze	1992312	1195387	2789237	2011611	0.97
125 Chlorobenzene-d5	1475337	885202	2065472	1511139	2.43

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-30may.b/8053003.d

Date: 30-May-2007 14:12

Client ID: Level 1

Sample Info: 0.2ml #1487-289

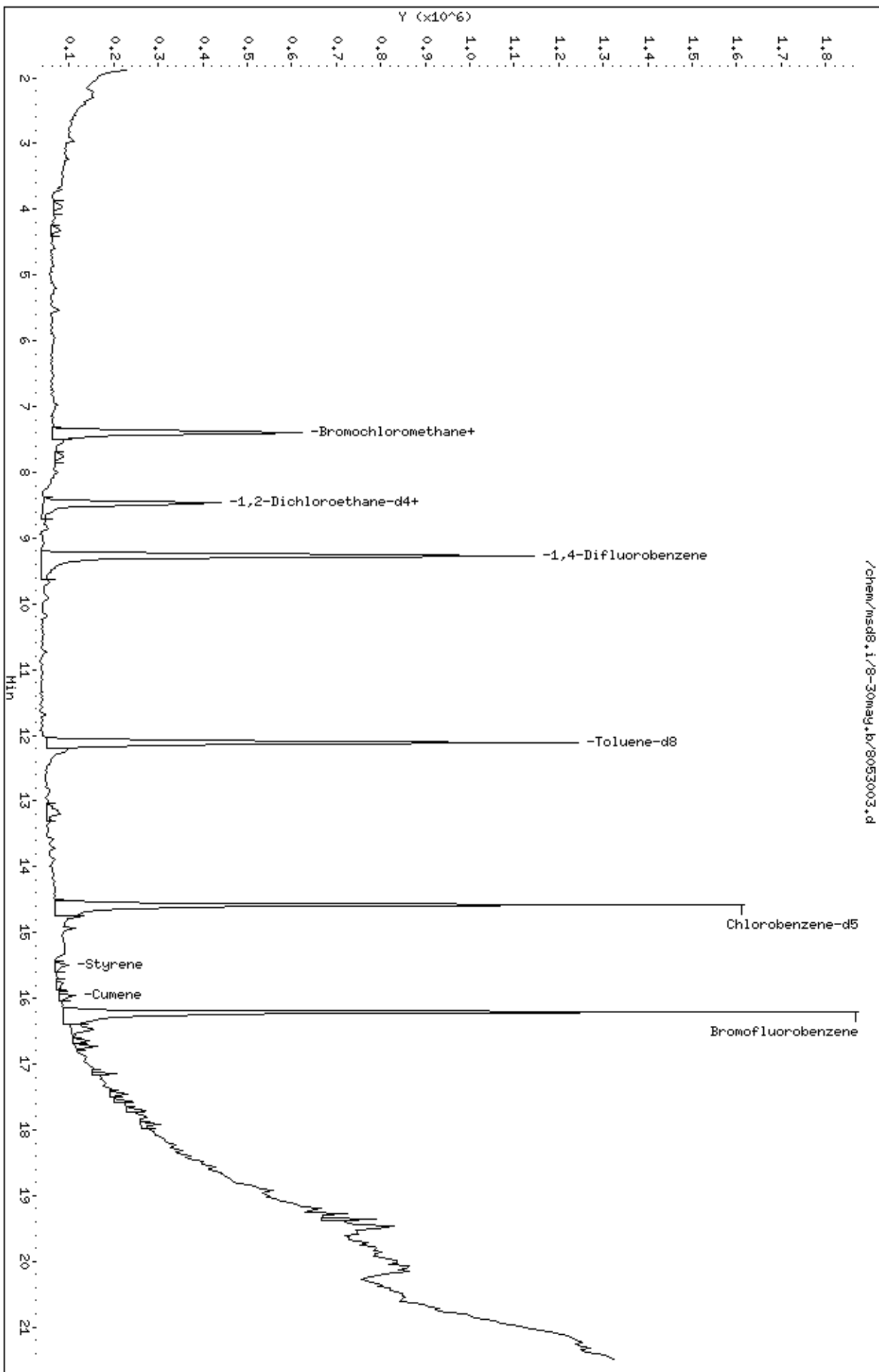
Column phase: RTX-624

Instrument: msd8.1

Operator: db

Column diameter: 0.53

/chem/msd8.1/8-30may.b/8053003.d



Report Date: 01-Jun-2007 13:36

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053004.d
 Lab Smp Id: ICAL Client Smp ID: Level 2
 Inj Date : 30-MAY-2007 14:39
 Operator : db Inst ID: msd8.i
 Smp Info : 0.5ml #1487-289
 Misc Info : 200ppbv-0.5ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 01-Jun-2007 10:56 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 14:39 Cal File: 8053004.d
 Als bottle: 1 Calibration Sample, Level: 2
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04Low+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS								
RT	EXP RT	(REL RT)	MASS	CAL-AMT	ON-COL	RESPONSE (PPBV)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5								
7.387	7.415	(1.000)	130	441719	25.0000		80.00- 120.00	100.00
7.387	7.415	(1.000)	128	338788			48.60- 108.60	76.70
7.387	7.415	(1.000)	49	651629			114.98- 174.98	147.52

* 88 1,4-Difluorobenzene CAS #: 540-36-3								
9.267	9.267	(1.000)	114	1974518	25.0000		80.00- 120.00	100.00
9.267	9.267	(1.000)	88	311817			0.00- 45.16	15.79

* 125 Chlorobenzene-d5 CAS #: 3114-55-4								
14.576	14.576	(1.000)	117	1505104	25.0000		80.00- 120.00	100.00
14.576	14.576	(1.000)	82	859236			0.00- 30.00	57.09

§ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0								
8.465	8.465	(1.146)	65	566125	25.0000	24.488	80.00- 120.00	100.00
8.465	8.465	(1.146)	67	303961			0.00- 30.00	53.69

§ 104 Toluene-d8 CAS #: 2037-26-5								
12.115	12.115	(1.307)	98	1755155	25.0000	25.676	80.00- 120.00	100.00
12.115	12.115	(1.307)	70	183881			0.00- 30.00	10.48

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
\$ 104 Toluene-d8 (continued)										
12.115	12.115	(1.307)	100	1118700			0.00- 30.00	63.74		

\$ 140 Bromofluorobenzene										
						CAS #: 460-00-4				
16.207	16.207	(1.112)	174	856607	25.0000	23.920	80.00- 120.00	100.00		
16.207	16.207	(1.112)	95	1122836			105.65- 165.65	131.08		
16.207	16.207	(1.112)	176	830280			66.34- 126.34	96.93		

4 Dichlorodifluoromethane/Fr12						CAS #: 75-71-8				
2.050	2.078	(0.278)	85	37346	0.50000	0.6171	80.00- 120.00	100.00		
2.050	2.078	(0.278)	87	12534			0.00- 30.00	33.56		

6 Freon 114						CAS #: 76-14-2				
2.216	2.189	(0.300)	135	31980	0.50000	0.5849	80.00- 120.00	100.00(H)		
2.216	2.189	(0.300)	137	10844			0.71- 60.71	33.91		

11 Vinyl Chloride						CAS #: 75-01-4				
2.438	2.438	(0.330)	62	20739	0.50000	0.6404	80.00- 120.00	100.00		
2.438	2.438	(0.330)	64	16204			0.00- 30.00	78.13		

10 1,3-Butadiene						CAS #: 106-99-0				
2.438	2.410	(0.330)	54	18786	0.50000	0.6782	80.00- 120.00	100.00		
2.410	2.410	(0.326)	39	32703			0.00- 30.00	174.08		

13 Bromomethane						CAS #: 74-83-9				
2.852	2.880	(0.386)	94	11115	0.50000	0.5137	80.00- 120.00	100.00		
2.852	2.880	(0.386)	96	7716			63.65- 123.65	69.42		

16 Chloroethane						CAS #: 75-00-3				
2.991	2.991	(0.405)	64	9774	0.50000	0.5703	80.00- 120.00	100.00		
2.991	2.991	(0.405)	49	2239			0.00- 30.00	22.91		
2.963	2.991	(0.401)	66	2848			0.00- 30.00	29.14		

18 Trichlorofluoromethane/Fr11						CAS #: 75-69-4				
3.239	3.239	(0.439)	101	35922	0.50000	0.5222	80.00- 120.00	100.00		
3.239	3.239	(0.439)	103	24332			34.50- 94.50	67.74		

28 Freon 113						CAS #: 76-13-1				
3.931	3.958	(0.532)	151	25449	0.50000	0.5592	80.00- 120.00	100.00		
3.931	3.958	(0.532)	153	17314			34.32- 94.32	68.03		
3.931	3.958	(0.532)	101	27551			85.50- 145.50	108.26		

29 1,1-Dichloroethene						CAS #: 75-35-4				
3.986	3.986	(0.540)	61	25336	0.50000	0.5606	80.00- 120.00	100.00		
3.986	3.986	(0.540)	96	14575			29.81- 89.81	57.53		
3.986	3.986	(0.540)	98	11440			9.00- 69.00	45.15		

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	====	=====	=====	=====	=====	=====		

33	Carbon Disulfide					CAS #:	75-15-0			
4.318	4.290	(0.585)	76	49375	0.50000	0.5829	80.00-	120.00	100.00	

40	Methylene Chloride					CAS #:	75-09-2			
4.843	4.815	(0.656)	49	19995	0.50000	0.5909	80.00-	120.00	100.00	
4.871	4.815	(0.659)	84	18154			40.47-	100.47	90.79	
4.843	4.815	(0.656)	51	6239			0.00-	30.00	31.20	

43	MTBE					CAS #:	1634-04-4			
5.175	5.175	(0.701)	73	22887	0.50000	0.4756	80.00-	120.00	100.00(a)	
5.147	5.175	(0.697)	57	8958			0.00-	52.63	39.14	
5.175	5.175	(0.701)	41	6914			0.00-	30.00	30.21	

45	trans-1,2-Dichloroethene					CAS #:	156-60-5			
5.203	5.203	(0.704)	96	19355	0.50000	0.6052	80.00-	120.00	100.00	
5.203	5.203	(0.704)	61	29045			119.47-	179.47	150.06	
5.203	5.203	(0.704)	98	12410			0.00-	30.00	64.12	

46	Hexane					CAS #:	110-54-3			
5.534	5.534	(0.749)	57	24684	0.50000	0.5089	80.00-	120.00	100.00	
5.534	5.534	(0.749)	43	18247			0.00-	30.00	73.92	
5.534	5.534	(0.749)	86	4409			0.00-	30.00	17.86	

54	1,1-Dichloroethane					CAS #:	75-34-3			
5.949	5.949	(0.805)	63	25663	0.50000	0.4922	80.00-	120.00	100.00(a)	
5.949	5.949	(0.805)	65	13490			1.86-	61.86	52.57	

65	2-Butanone					CAS #:	78-93-3			
7.027	7.027	(0.951)	72	8897	0.50000	0.6078	80.00-	120.00	100.00	
7.027	7.027	(0.951)	43	35422			409.65-	469.65	398.13	
7.055	7.027	(0.955)	57	4439			0.00-	30.00	49.89	

64	cis-1,2-Dichloroethene					CAS #:	156-59-2			
6.972	6.972	(0.944)	61	26832	0.50000	0.6503	80.00-	120.00	100.00	
6.972	6.972	(0.944)	96	19889			43.20-	103.20	74.12	
6.972	6.972	(0.944)	98	13061			17.85-	77.85	48.68	

67	Tetrahydrofuran					CAS #:	109-99-9			
7.415	7.415	(1.004)	42	23210	0.50000	0.6100	80.00-	120.00	100.00	
7.415	7.415	(1.004)	71	9141			5.48-	65.48	39.38	
7.415	7.415	(1.004)	72	10737			0.00-	30.00	46.26	

70	Chloroform					CAS #:	67-66-3			
7.525	7.553	(1.019)	83	29884	0.50000	0.5731	80.00-	120.00	100.00	
7.525	7.553	(1.019)	85	17860			32.67-	92.67	59.76	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

75	1,1,1-Trichloroethane					CAS #:	71-55-6			
7.774	7.774	(1.052)	97	32604	0.50000	0.5968	80.00-	120.00	100.00	
7.774	7.774	(1.052)	99	18613			34.55-	94.55	57.09	

73	Cyclohexane					CAS #:	110-82-7			
7.746	7.746	(1.049)	84	27736	0.50000	0.6411	80.00-	120.00	100.00	
7.746	7.746	(1.049)	56	29634			96.25-	156.25	106.84	
7.746	7.746	(1.049)	41	14639			32.28-	92.28	52.78	

77	Carbon Tetrachloride					CAS #:	56-23-5			
7.995	8.023	(1.082)	119	26063	0.50000	0.5243	80.00-	120.00	100.00	
7.995	8.023	(1.082)	117	24950			73.30-	133.30	95.73	

81	Benzene					CAS #:	71-43-2			
8.438	8.438	(0.910)	78	52333	0.50000	0.5867	80.00-	120.00	100.00	
8.438	8.438	(0.910)	77	11248			0.00-	30.00	21.49	

83	1,2-Dichloroethane					CAS #:	107-06-2			
8.603	8.603	(0.928)	62	22290	0.50000	0.6159	80.00-	120.00	100.00	
8.603	8.603	(0.928)	64	6977			0.00-	30.00	31.30	

85	Heptane					CAS #:	142-82-5			
8.852	8.852	(0.955)	100	7535	0.50000	0.7806	80.00-	120.00	100.00	
8.852	8.852	(0.955)	43	29890			0.00-	30.00	396.68	
8.852	8.852	(0.955)	71	17485			0.00-	30.00	232.05	

94	Trichloroethene					CAS #:	79-01-6			
9.682	9.682	(1.045)	95	21043	0.50000	0.5953	80.00-	120.00	100.00	
9.682	9.682	(1.045)	130	22414			78.57-	138.57	106.52	
9.682	9.682	(1.045)	97	14838			34.08-	94.08	70.51	

97	1,2-Dichloropropane					CAS #:	78-87-5			
10.179	10.179	(1.098)	63	16877	0.50000	0.5460	80.00-	120.00	100.00	
10.179	10.179	(1.098)	62	13213			42.26-	102.26	78.29	
10.179	10.179	(1.098)	41	10990			26.77-	86.77	65.12	

100	Bromodichloromethane					CAS #:	75-27-4			
10.732	10.732	(1.158)	83	28920	0.50000	0.5601	80.00-	120.00	100.00	
10.732	10.732	(1.158)	85	18413			31.75-	91.75	63.67	

102	cis-1,3-Dichloropropene					CAS #:	10061-01-5			
11.672	11.673	(1.260)	75	24425	0.50000	0.5686	80.00-	120.00	100.00	
11.672	11.673	(1.260)	77	6970			1.36-	61.36	28.54	
11.700	11.673	(1.263)	39	8862			17.70-	77.70	36.28	

103	4-Methyl-2-pentanone					CAS #:	108-10-1			
12.032	12.032	(1.298)	58	11214	0.50000	0.4825	80.00-	120.00	100.00(a)	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
103 4-Methyl-2-pentanone (continued)									
12.032	12.032	(1.298)	43	37485			0.00- 30.00	334.27	
12.032	12.032	(1.298)	85	6047			0.00- 30.00	53.92	

105 Toluene CAS #: 108-88-3									
12.253	12.253	(1.322)	91	49671	0.50000	0.5479	80.00- 120.00	100.00	
12.253	12.253	(1.322)	92	32962			29.69- 89.69	66.36	

108 trans-1,3-Dichloropropene CAS #: 10061-02-6									
12.861	12.861	(0.882)	75	22841	0.50000	0.5235	80.00- 120.00	100.00	
12.834	12.861	(0.880)	77	6413			0.08- 60.08	28.08	
12.834	12.861	(0.880)	39	12864			15.97- 75.97	56.32	

110 1,1,2-Trichloroethane CAS #: 79-00-5									
13.138	13.138	(0.901)	97	19937	0.50000	0.6230	80.00- 120.00	100.00	
13.138	13.138	(0.901)	99	10911			29.85- 89.85	54.73	
13.138	13.138	(0.901)	83	15123			54.93- 114.93	75.85	

112 Tetrachloroethene CAS #: 127-18-4									
13.193	13.193	(0.905)	166	23583	0.50000	0.5445	80.00- 120.00	100.00	
13.193	13.193	(0.905)	129	18864			45.08- 105.08	79.99	
13.193	13.193	(0.905)	131	17800			44.22- 104.22	75.48	

114 2-Hexanone CAS #: 591-78-6									
13.580	13.580	(0.932)	58	16251	0.50000	0.5125	80.00- 120.00	100.00	
13.580	13.580	(0.932)	43	29468			148.62- 208.62	181.33	
13.580	13.580	(0.932)	100	5765			0.00- 30.00	35.47	

116 Dibromochloromethane CAS #: 124-48-1									
13.719	13.719	(0.941)	129	25600	0.50000	0.5104	80.00- 120.00	100.00	
13.719	13.719	(0.941)	127	16875			0.00- 30.00	65.92	

117 1,2-Dibromoethane CAS #: 106-93-4									
13.884	13.884	(0.953)	107	26910	0.50000	0.5519	80.00- 120.00	100.00	
13.884	13.884	(0.953)	109	23468			63.81- 123.81	87.21	

126 Chlorobenzene CAS #: 108-90-7									
14.631	14.631	(1.004)	112	36793	0.50000	0.4761	80.00- 120.00	100.00(a)	
14.631	14.631	(1.004)	114	11501			1.84- 61.84	31.26	
14.603	14.631	(1.002)	77	32356			26.78- 86.78	87.94	

129 Ethyl Benzene CAS #: 100-41-4									
14.769	14.769	(1.013)	106	24362	0.50000	0.5957	80.00- 120.00	100.00	
14.769	14.769	(1.013)	91	64080			0.00- 30.00	263.03	

130 m,p-Xylene CAS #: 108-38-3									
14.935	14.935	(1.025)	106	25452	0.50000	0.5107	80.00- 120.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
130 m,p-Xylene (continued)									
14.935	14.935	(1.025)	91	55732			0.00- 30.00	218.97	

132 o-Xylene CAS #: 95-47-6									
15.488	15.488	(1.063)	106	27489	0.50000	0.5466	80.00- 120.00	100.00	
15.488	15.488	(1.063)	91	51508			178.74- 238.74	187.38	

134 Styrene CAS #: 100-42-5									
15.516	15.516	(1.064)	104	35605	0.50000	0.4966	80.00- 120.00	100.00(a)	
15.516	15.516	(1.064)	78	18620			20.27- 80.27	52.30	

135 Bromoform CAS #: 75-25-2									
15.765	15.765	(1.082)	173	19027	0.50000	0.4313	80.00- 120.00	100.00(a)	
15.765	15.765	(1.082)	171	11415			20.67- 80.67	59.99	

144 1,1,2,2-Tetrachloroethane CAS #: 79-34-5									
16.456	16.456	(1.129)	83	40065	0.50000	0.5448	80.00- 120.00	100.00	
16.456	16.456	(1.129)	85	25337			31.59- 91.59	63.24	

147 4-Ethyltoluene CAS #: 622-96-8									
16.649	16.649	(1.142)	105	67698	0.50000	0.5016	80.00- 120.00	100.00	
16.649	16.649	(1.142)	120	22846			0.00- 59.85	33.75	

148 1,3,5-Trimethylbenzene CAS #: 108-67-8									
16.732	16.732	(1.148)	105	71898	0.50000	0.5542	80.00- 120.00	100.00	
16.732	16.732	(1.148)	120	34076			0.00- 30.00	47.39	

153 1,2,4-Trimethylbenzene CAS #: 95-63-6									
17.147	17.147	(1.176)	105	72353	0.50000	0.5540	80.00- 120.00	100.00	
17.147	17.147	(1.176)	120	33530			15.22- 75.22	46.34	

156 1,3-Dichlorobenzene CAS #: 541-73-1									
17.451	17.451	(1.197)	146	40043	0.50000	0.4704	80.00- 120.00	100.00(a)	
17.451	17.451	(1.197)	148	25065			0.00- 30.00	62.60	
17.451	17.451	(1.197)	111	16317			0.00- 30.00	40.75	

157 1,4-Dichlorobenzene CAS #: 106-46-7									
17.562	17.562	(1.205)	146	46404	0.50000	0.5751	80.00- 120.00	100.00	
17.562	17.562	(1.205)	148	26549			0.00- 30.00	57.21	
17.562	17.562	(1.205)	111	15041			0.00- 30.00	32.41	

158 alpha-Chlorotoluene CAS #: 100-44-7									
17.700	17.700	(1.214)	91	56800	0.50000	0.4843	80.00- 120.00	100.00(a)	
17.700	17.700	(1.214)	126	13675			0.00- 30.00	24.08	

161 1,2-Dichlorobenzene CAS #: 95-50-1									
17.921	17.921	(1.230)	146	44406	0.50000	0.5415	80.00- 120.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
161 1,2-Dichlorobenzene (continued)									
17.921	17.921	(1.230)	148	29125			33.37- 93.37	65.59	
17.921	17.921	(1.230)	111	15806			7.65- 67.65	35.59	

137 Cumene CAS #: 98-82-8									
15.958	15.958	(1.095)	105	72289	0.50000	0.5259	80.00- 120.00	100.00	
15.958	15.958	(1.095)	120	21844			0.00- 30.00	30.22	
15.958	15.958	(1.095)	51	7145			0.00- 30.00	9.88	

145 Propylbenzene CAS #: 103-65-1									
16.484	16.484	(1.131)	91	84913	0.50000	0.5394	80.00- 120.00	100.00	
16.484	16.484	(1.131)	120	21938			0.00- 30.00	25.84	
16.484	16.484	(1.131)	105	3685			0.00- 30.00	4.34	

80 2,2,4-Trimethylpentane CAS #: 540-84-1									
8.465	8.465	(1.146)	57	69240	0.50000	0.5361	80.00- 120.00	100.00	
8.465	8.465	(1.146)	56	27170			0.00- 30.00	39.24	
8.465	8.465	(1.146)	41	17762			0.00- 30.00	25.65	

95 Methyl Cyclohexane CAS #: 108-87-2									
9.903	9.903	(1.341)	83	28853	0.50000	0.5354	70.00- 130.00	100.00	
9.903	9.903	(1.341)	98	14551			0.00- 30.00	50.43	
9.903	9.903	(1.341)	55	24257			0.00- 30.00	84.07	

QC Flag Legend

- a - Target compound detected but, quantitated amount
 Below Limit Of Quantitation(BLOQ).
- H - Operator selected an alternate compound hit.

Report Date: 01-Jun-2007 13:36

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053004.d

Calibration Time: 14:39

Lab Smp Id: ICAL

Client Smp ID: Level 2

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-0.5ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	448309	268985	627633	441719	-1.47
88 1,4-Difluorobenze	2033490	1220094	2846886	1974518	-2.90
125 Chlorobenzene-d5	1524596	914758	2134434	1505104	-1.28

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-30may.b/8053004.d

Date: 30-May-2007 14:39

Client ID: Level 2

Sample Info: 0.5ml #1487-289

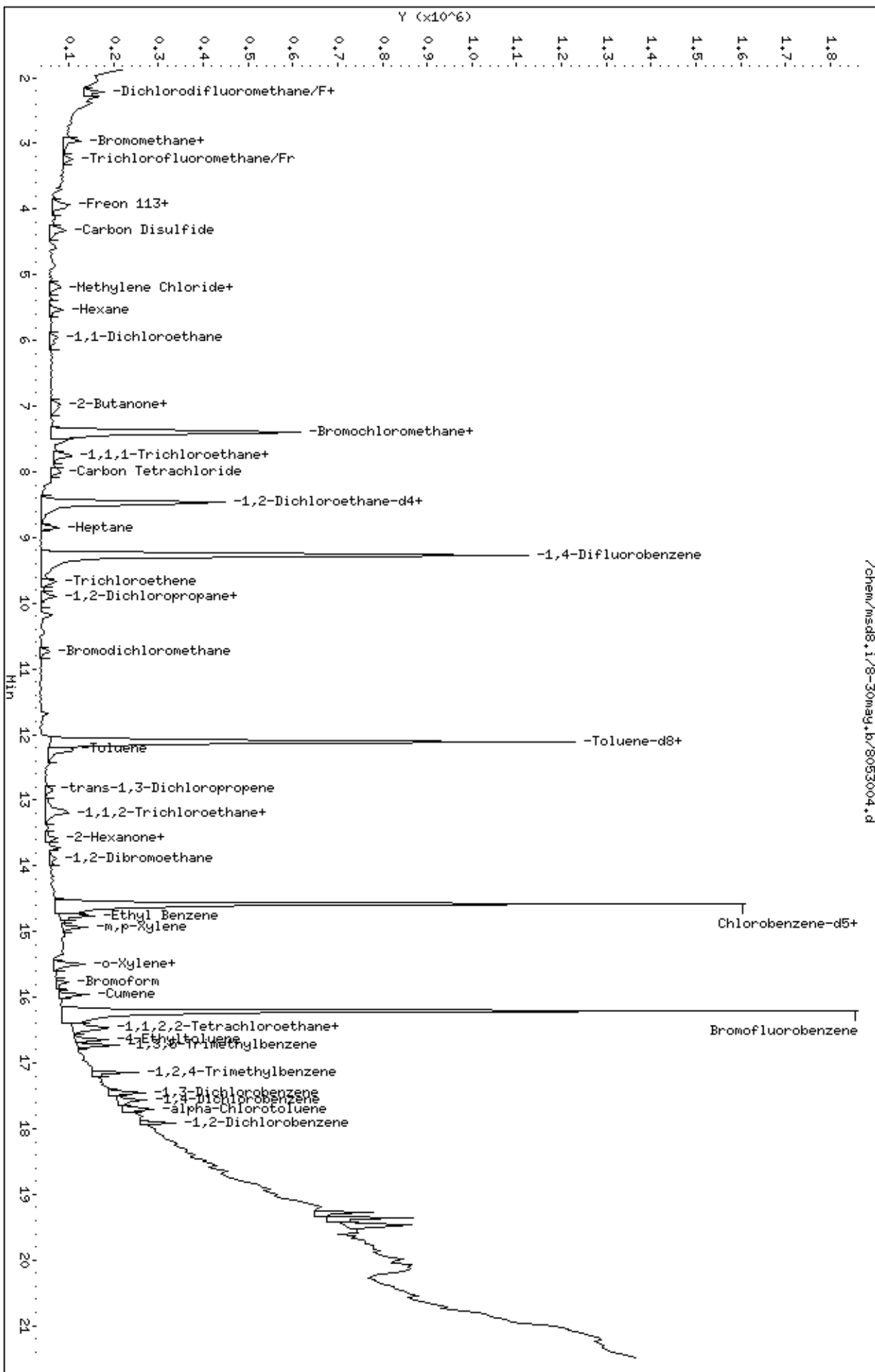
Column phase: RTX-624

Instrument: msd8.1

Operator: db

Column diameter: 0.53

Page 1



/chem/msd8.1/8-30may.b/8053004.d

Report Date: 07-Jun-2007 13:41

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-07jun.b/8060704.d
 Lab Smp Id: ICAL Client Smp ID: Level 3
 Inj Date : 07-JUN-2007 11:09
 Operator : JG Inst ID: msd8.i
 Smp Info : 2mL #1443-96
 Misc Info : 200ppbv-2ppbv
 Comment :
 Method : /chem/msd8.i/8-07jun.b/t14q530b.m
 Meth Date : 07-Jun-2007 13:41 jgray Quant Type: ISTD
 Cal Date : 07-JUN-2007 11:09 Cal File: 8060704.d
 Als bottle: 1 Calibration Sample, Level: 3
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: sp16b.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	CAL-AMT	ON-COL	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	359827	25.0000			70.00- 130.00	100.00
7.387	7.387	(1.000)	128	276445				42.98- 102.98	76.83
7.387	7.387	(1.000)	49	515800				112.53- 172.53	143.35

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	1618571	25.0000			70.00- 130.00	100.00
9.267	9.267	(1.000)	88	239337				0.00- 44.58	14.79

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	1231123	25.0000			70.00- 130.00	100.00
14.576	14.576	(1.000)	82	682772				0.00- 30.00	55.46

1 Freon 152a CAS #: 75-37-6									
2.050	2.050	(0.278)	65	28430	2.00000	2.281		70.00- 130.00	100.00
2.078	2.078	(0.281)	51	113235				0.00- 30.00	398.29

20 Freon123a CAS #: 354-23-4									
3.682	3.682	(0.498)	67	39913	2.00000	2.074		70.00- 130.00	100.00
3.709	3.709	(0.502)	117	33578				0.00- 30.00	84.13

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
21 Freon123						CAS #:	306-83-2			
3.792	3.792	(0.513)	83	24804	2.00000	2.045	70.00- 130.00	100.00		
3.792	3.792	(0.513)	133	7379			0.00- 30.00	29.75		
3.792	3.792	(0.513)	85	18073			0.00- 30.00	72.86		

38 tert-Butyl-Alcohol						CAS #:	75-65-0			
4.954	4.954	(0.671)	59	67488	2.00000	2.691	70.00- 130.00	100.00		
4.954	4.954	(0.671)	41	16170			0.00- 30.00	23.96		
4.954	4.954	(0.671)	57	7979			0.00- 30.00	11.82		

49 Isopropyl ether						CAS #:	108-20-3			
5.949	5.949	(0.805)	45	127830	2.00000	2.038	70.00- 130.00	100.00		
5.977	5.977	(0.809)	87	31904			0.00- 30.00	24.96		
5.977	5.977	(0.809)	59	14776			0.00- 30.00	11.56		

52 1-Propanol						CAS #:	71-23-8			
6.170	6.170	(0.835)	42	9618	2.00000	2.128	70.00- 130.00	100.00		
6.170	6.170	(0.835)	59	11102			0.00- 30.00	115.43		
6.170	6.170	(0.835)	41	10796			0.00- 30.00	112.25		

58 Ethyl-tert-butyl Ether						CAS #:	637-92-3			
6.585	6.585	(0.891)	59	75388	2.00000	1.800	70.00- 130.00	100.00(a)		
6.585	6.585	(0.891)	87	27007			0.00- 30.00	35.82		
6.585	6.585	(0.891)	41	10624			0.00- 30.00	14.09		

61 Ethyl Acetate						CAS #:	141-78-6			
7.083	7.083	(0.959)	70	11438	2.00000	2.093	70.00- 130.00	100.00		
7.083	7.083	(0.959)	43	92650			0.00- 30.00	810.02		
7.083	7.083	(0.959)	61	15040			0.00- 30.00	131.49		

78 Isobutanol						CAS #:	78-83-1			
8.437	8.437	(0.910)	43	35107	2.00000	1.890	70.00- 130.00	100.00		
8.437	8.437	(0.910)	41	23753			0.00- 30.00	67.66		

79 tert-amyl-Methyl Ether						CAS #:	994-05-8			
8.631	8.631	(1.168)	73	74928	2.00000	1.956	70.00- 130.00	100.00(a)		
8.631	8.631	(1.168)	87	18030			0.00- 30.00	24.06		
8.631	8.631	(1.168)	55	22510			0.00- 30.00	30.04		

89 1-Butanol						CAS #:	71-36-3			
9.737	9.737	(1.051)	56	28801	2.00000	1.818	70.00- 130.00	100.00(a)		
9.737	9.737	(1.051)	41	18725			0.00- 30.00	65.02		
9.737	9.737	(1.051)	43	17406			0.00- 30.00	60.44		

136 Cyclohexanone						CAS #:	108-94-1			
16.152	16.152	(1.108)	55	54831	2.00000	1.946	70.00- 130.00	100.00(a)		

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	====	=====	=====	=====	=====	=====	
136 Cyclohexanone (continued)									
16.152	16.152	(1.108)	98	24425			0.00- 30.00	44.55	
16.152	16.152	(1.108)	42	35102			0.00- 30.00	64.02	

QC Flag Legend

a - Target compound detected but, quantitated amount
Below Limit Of Quantitation(BLOQ).

Report Date: 07-Jun-2007 13:41

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 07-JUN-2007

Lab File ID: 8060704.d

Calibration Time: 11:37

Lab Smp Id: ICAL

Client Smp ID: Level 3

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: JG

Method File: /chem/msd8.i/8-07jun.b/t14q530b.m

Misc Info: 200ppbv-2ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	350593	210356	490830	359827	2.63
88 1,4-Difluorobenze	1524282	914569	2133995	1618571	6.19
125 Chlorobenzene-d5	1168126	700876	1635376	1231123	5.39

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-07jun.b/8060704.d

Date: 07-JUN-2007 11:09

Client ID: Level 3

Sample Info: 2mL #1443-96

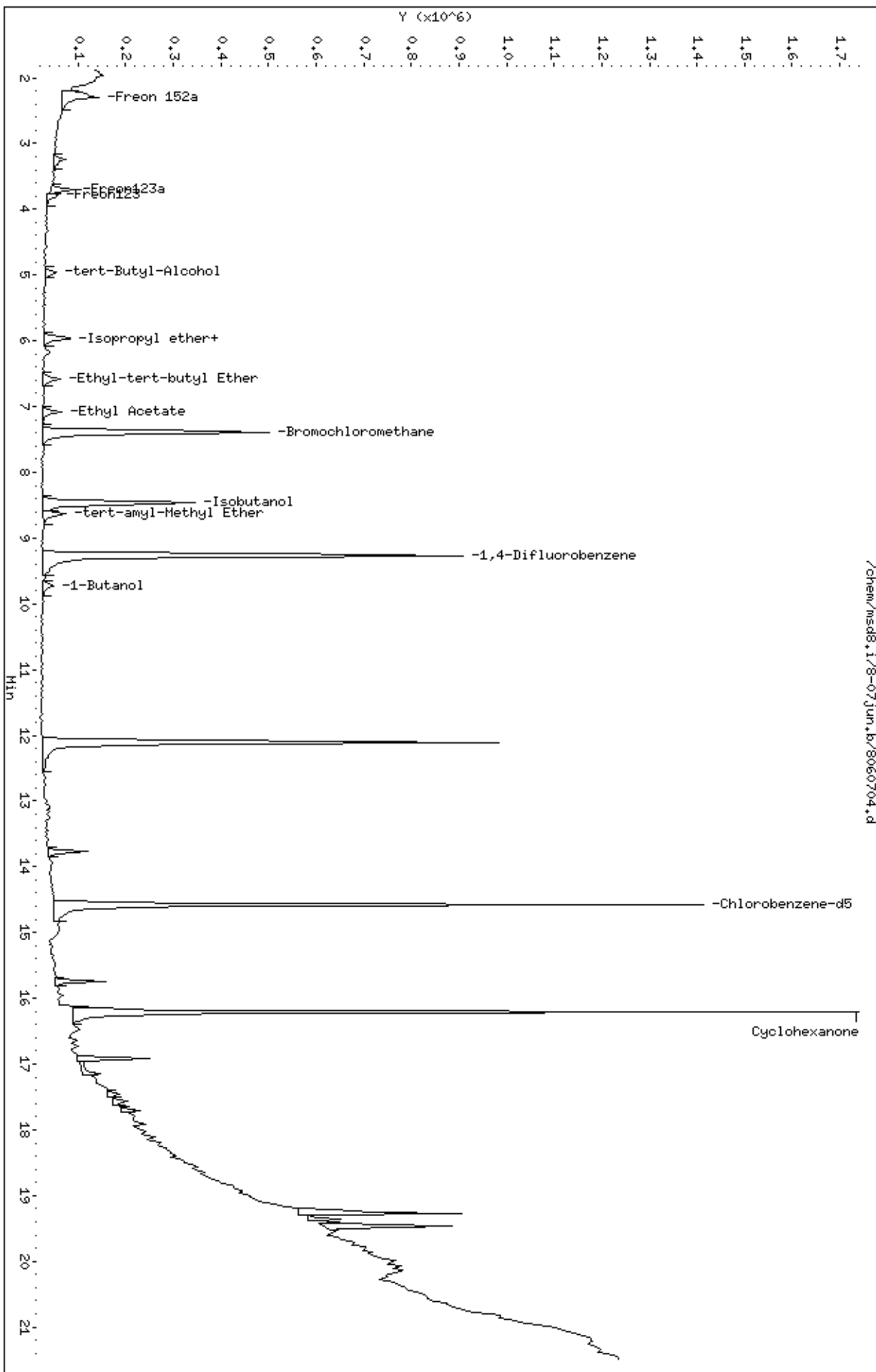
Column phase: RTX-624

Instrument: msd8.1

Operator: JG

Column diameter: 0.53

/chem/msd8.1/8-07jun.b/8060704.d



Report Date: 31-May-2007 14:52

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053005.d
 Lab Smp Id: ICAL Client Smp ID: Level 3
 Inj Date : 30-MAY-2007 15:07
 Operator : db Inst ID: msd8.i
 Smp Info : 2.0ml #1487-289
 Misc Info : 200ppbv-2.0ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 31-May-2007 14:52 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 15:07 Cal File: 8053005.d
 Als bottle: 1 Calibration Sample, Level: 3
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04mdl+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS								
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	ON-COL	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5								
7.387	7.387	(1.000)	130	444018	25.0000		70.00- 130.00	100.00
7.387	7.387	(1.000)	128	343097			47.57- 107.57	77.27
7.387	7.387	(1.000)	49	663820			113.47- 173.47	149.50

* 88 1,4-Difluorobenzene CAS #: 540-36-3								
9.267	9.267	(1.000)	114	2035031	25.0000		70.00- 130.00	100.00
9.267	9.267	(1.000)	88	289330			0.00- 45.68	14.22

* 125 Chlorobenzene-d5 CAS #: 3114-55-4								
14.576	14.576	(1.000)	117	1499190	25.0000		70.00- 130.00	100.00
14.576	14.576	(1.000)	82	846297			0.00- 30.00	56.45

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0								
8.465	8.465	(1.146)	65	590610	25.0000	25.415	70.00- 130.00	100.00
8.465	8.465	(1.146)	67	303638			0.00- 30.00	51.41

\$ 104 Toluene-d8 CAS #: 2037-26-5								
12.115	12.115	(1.307)	98	1781631	25.0000	25.288	70.00- 130.00	100.00
12.115	12.115	(1.307)	70	175095			0.00- 30.00	9.83

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
\$ 104 Toluene-d8 (continued)										
12.115	12.115	(1.307)	100	1161193			0.00- 30.00	65.18		

\$ 140 Bromofluorobenzene										
						CAS #: 460-00-4				
16.207	16.207	(1.112)	174	877440	25.0000	24.599	70.00- 130.00	100.00		
16.207	16.207	(1.112)	95	1139577			102.16- 162.16	129.88		
16.207	16.207	(1.112)	176	835648			64.31- 124.31	95.24		

3 Propylene										
						CAS #: 115-07-1				
2.023	2.023	(0.274)	41	68550	2.00000	2.852	70.00- 130.00	100.00		
2.023	2.023	(0.274)	42	32909			0.00- 30.00	48.01		
2.023	2.023	(0.274)	39	33753			0.00- 30.00	49.24		

4 Dichlorodifluoromethane/Fr12										
						CAS #: 75-71-8				
2.050	2.050	(0.278)	85	143030	2.00000	2.351	70.00- 130.00	100.00		
2.050	2.050	(0.278)	87	45048			0.00- 30.00	31.50		

6 Freon 114										
						CAS #: 76-14-2				
2.216	2.216	(0.300)	135	109387	2.00000	1.990	70.00- 130.00	100.00		
2.216	2.216	(0.300)	137	29718			1.53- 61.53	27.17		

8 Chloromethane										
						CAS #: 74-87-3				
2.299	2.299	(0.311)	50	62645	2.00000	2.236	70.00- 130.00	100.00		
2.299	2.299	(0.311)	52	20724			0.00- 30.00	33.08		

9 Butane										
						CAS #: 106-97-8				
2.382	2.382	(0.322)	58	17616	2.00000	2.408	70.00- 130.00	100.00		
2.382	2.382	(0.322)	43	128408			0.00- 30.00	728.93		

11 Vinyl Chloride										
						CAS #: 75-01-4				
2.410	2.410	(0.326)	62	65793	2.00000	2.021	70.00- 130.00	100.00		
2.437	2.437	(0.330)	64	27327			0.00- 30.00	41.53		

10 1,3-Butadiene										
						CAS #: 106-99-0				
2.410	2.410	(0.326)	54	61421	2.00000	2.206	70.00- 130.00	100.00		
2.410	2.410	(0.326)	39	64322			0.00- 30.00	104.72		

13 Bromomethane										
						CAS #: 74-83-9				
2.852	2.852	(0.386)	94	42582	2.00000	1.958	70.00- 130.00	100.00		
2.852	2.852	(0.386)	96	42273			65.03- 125.03	99.27		

16 Chloroethane										
						CAS #: 75-00-3				
2.990	2.990	(0.405)	64	36761	2.00000	2.134	70.00- 130.00	100.00		
2.990	2.990	(0.405)	49	9519			0.00- 30.00	25.89		
2.990	2.990	(0.405)	66	10406			0.00- 30.00	28.31		

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
15 Isopentane						CAS #:	78-78-4			
2.963	2.963	(0.401)	43	84741	2.00000	2.085	70.00-	130.00	100.00	
2.963	2.963	(0.401)	57	57480			0.00-	30.00	67.83	
2.963	2.963	(0.401)	72	7193			0.00-	30.00	8.49	

18 Trichlorofluoromethane/Fr11						CAS #:	75-69-4			
3.239	3.239	(0.439)	101	149777	2.00000	2.166	70.00-	130.00	100.00	
3.239	3.239	(0.439)	103	91623			34.97-	94.97	61.17	

23 Ethanol						CAS #:	64-17-5			
3.516	3.516	(0.476)	45	24067	2.00000	2.180	70.00-	130.00	100.00	
3.516	3.516	(0.476)	43	13031			0.00-	30.00	54.14	
3.516	3.516	(0.476)	46	11087			0.00-	30.00	46.07	

28 Freon 113						CAS #:	76-13-1			
3.931	3.931	(0.532)	151	99380	2.00000	2.172	70.00-	130.00	100.00	
3.931	3.931	(0.532)	153	59189			33.71-	93.71	59.56	
3.931	3.931	(0.532)	101	109122			86.34-	146.34	109.80	

29 1,1-Dichloroethene						CAS #:	75-35-4			
3.986	3.986	(0.540)	61	96927	2.00000	2.133	70.00-	130.00	100.00	
3.986	3.986	(0.540)	96	59254			30.42-	90.42	61.13	
3.986	3.986	(0.540)	98	44431			8.39-	68.39	45.84	

30 Acetone						CAS #:	67-64-1			
4.124	4.124	(0.558)	58	34898	2.00000	2.294	70.00-	130.00	100.00	
4.124	4.124	(0.558)	43	95643			0.00-	30.00	274.06	

33 Carbon Disulfide						CAS #:	75-15-0			
4.318	4.318	(0.584)	76	172959	2.00000	2.031	70.00-	130.00	100.00	

34 2-Propanol						CAS #:	67-63-0			
4.318	4.318	(0.584)	45	129095	2.00000	2.346	70.00-	130.00	100.00	
4.318	4.318	(0.584)	43	34823			0.00-	30.00	26.97	
4.318	4.318	(0.584)	59	7801			0.00-	30.00	6.04	

37 3-Chloropropene						CAS #:	107-05-1			
4.566	4.566	(0.618)	76	25780	2.00000	1.898	70.00-	130.00	100.00(a)	
4.566	4.566	(0.618)	41	78835			0.00-	30.00	305.80	

40 Methylene Chloride						CAS #:	75-09-2			
4.843	4.843	(0.656)	49	72037	2.00000	2.118	70.00-	130.00	100.00	
4.843	4.843	(0.656)	84	56488			41.01-	101.01	78.42	
4.843	4.843	(0.656)	51	22744			0.00-	30.00	31.57	

43 MTBE						CAS #:	1634-04-4			
5.175	5.175	(0.701)	73	78059	2.00000	1.614	70.00-	130.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
43 MTBE (continued)									
5.147	5.147	(0.697)	57	21684			0.00- 53.82	27.78	
5.147	5.147	(0.697)	41	16336			0.00- 30.00	20.93	

45 trans-1,2-Dichloroethene						CAS #: 156-60-5			
5.202	5.202	(0.704)	96	70934	2.00000	2.206	70.00- 130.00	100.00	
5.202	5.202	(0.704)	61	94977			119.04- 179.04	133.89	
5.202	5.202	(0.704)	98	40611			0.00- 30.00	57.25	

46 Hexane						CAS #: 110-54-3			
5.534	5.534	(0.749)	57	95925	2.00000	1.967	70.00- 130.00	100.00	
5.534	5.534	(0.749)	43	60460			0.00- 30.00	63.03	
5.534	5.534	(0.749)	86	15672			0.00- 30.00	16.34	

54 1,1-Dichloroethane						CAS #: 75-34-3			
5.949	5.949	(0.805)	63	114940	2.00000	2.193	70.00- 130.00	100.00	
5.949	5.949	(0.805)	65	34286			1.22- 61.22	29.83	

55 Vinyl Acetate						CAS #: 108-05-4			
6.032	6.032	(0.817)	86	14556	2.00000	1.968	70.00- 130.00	100.00(a)	
6.032	6.032	(0.817)	43	135869			0.00- 30.00	933.42	
6.032	6.032	(0.817)	42	12791			0.00- 30.00	87.87	

64 cis-1,2-Dichloroethene						CAS #: 156-59-2			
6.972	6.972	(0.944)	61	84153	2.00000	2.029	70.00- 130.00	100.00	
6.972	6.972	(0.944)	96	66205			45.09- 105.09	78.67	
6.972	6.972	(0.944)	98	40871			17.66- 77.66	48.57	

65 2-Butanone						CAS #: 78-93-3			
7.027	7.027	(0.951)	72	30203	2.00000	2.052	70.00- 130.00	100.00	
7.027	7.027	(0.951)	43	124260			398.56- 458.56	411.42	
7.027	7.027	(0.951)	57	9637			0.00- 30.00	31.91	

67 Tetrahydrofuran						CAS #: 109-99-9			
7.414	7.414	(1.004)	42	78266	2.00000	2.046	70.00- 130.00	100.00	
7.414	7.414	(1.004)	71	36816			5.14- 65.14	47.04	
7.414	7.414	(1.004)	72	33830			0.00- 30.00	43.22	

70 Chloroform						CAS #: 67-66-3			
7.525	7.525	(1.019)	83	105767	2.00000	2.018	70.00- 130.00	100.00	
7.525	7.525	(1.019)	85	69135			31.98- 91.98	65.37	

73 Cyclohexane						CAS #: 110-82-7			
7.746	7.746	(1.049)	84	84366	2.00000	1.940	70.00- 130.00	100.00	
7.746	7.746	(1.049)	56	109014			99.30- 159.30	129.22	
7.746	7.746	(1.049)	41	61541			33.84- 93.84	72.95	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

75	1,1,1-Trichloroethane					CAS #:	71-55-6			
7.774	7.774	(1.052)	97	108279	2.00000	1.972	70.00-	130.00	100.00	
7.774	7.774	(1.052)	99	66999			34.73-	94.73	61.88	

77	Carbon Tetrachloride					CAS #:	56-23-5			
7.995	7.995	(1.082)	119	87757	2.00000	1.756	70.00-	130.00	100.00	
7.995	7.995	(1.082)	117	99969			74.04-	134.04	113.92	

81	Benzene					CAS #:	71-43-2			
8.437	8.437	(0.910)	78	185467	2.00000	2.017	70.00-	130.00	100.00	
8.437	8.437	(0.910)	77	44486			0.00-	30.00	23.99	

80	2,2,4-Trimethylpentane					CAS #:	540-84-1			
8.465	8.465	(1.146)	57	269118	2.00000	2.073	70.00-	130.00	100.00	
8.465	8.465	(1.146)	56	84149			0.00-	30.00	31.27	
8.465	8.465	(1.146)	41	62827			0.00-	30.00	23.35	

83	1,2-Dichloroethane					CAS #:	107-06-2			
8.603	8.603	(0.928)	62	74900	2.00000	2.008	70.00-	130.00	100.00	
8.603	8.603	(0.928)	64	24637			0.00-	30.00	32.89	

85	Heptane					CAS #:	142-82-5			
8.852	8.852	(0.955)	100	18594	2.00000	1.869	70.00-	130.00	100.00	
8.852	8.852	(0.955)	43	110807			0.00-	30.00	595.93	
8.852	8.852	(0.955)	71	61004			0.00-	30.00	328.08	

94	Trichloroethene					CAS #:	79-01-6			
9.682	9.682	(1.045)	95	73161	2.00000	2.008	70.00-	130.00	100.00	
9.682	9.682	(1.045)	130	72926			77.84-	137.84	99.68	
9.682	9.682	(1.045)	97	48621			34.22-	94.22	66.46	

95	Methyl Cyclohexane					CAS #:	108-87-2			
9.903	9.903	(1.341)	83	109945	2.00000	2.029	70.00-	130.00	100.00	
9.903	9.903	(1.341)	98	54584			0.00-	30.00	49.65	
9.903	9.903	(1.341)	55	89274			0.00-	30.00	81.20	

97	1,2-Dichloropropane					CAS #:	78-87-5			
10.179	10.179	(1.098)	63	70027	2.00000	2.198	70.00-	130.00	100.00	
10.179	10.179	(1.098)	62	47177			42.15-	102.15	67.37	
10.179	10.179	(1.098)	41	36977			26.47-	86.47	52.80	

98	1,4-Dioxane					CAS #:	123-91-1			
10.428	10.428	(1.125)	88	40326	2.00000	2.075	70.00-	130.00	100.00	
10.428	10.428	(1.125)	58	27930			44.03-	104.03	69.26	
10.428	10.428	(1.125)	57	7847			0.00-	30.00	19.46	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT	ON-COL	TARGET RANGE	RATIO	
					(PPEV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

100	Bromodichloromethane				CAS #:		75-27-4		
10.732	10.732	(1.158)	83	104941	2.00000	1.972	70.00-	130.00	100.00
10.732	10.732	(1.158)	85	62504			31.03-	91.03	59.56

102	cis-1,3-Dichloropropene				CAS #:		10061-01-5		
11.672	11.672	(1.260)	75	85811	2.00000	1.938	70.00-	130.00	100.00
11.672	11.672	(1.260)	77	26235			1.19-	61.19	30.57
11.672	11.672	(1.260)	39	39774			16.93-	76.93	46.35

103	4-Methyl-2-pentanone				CAS #:		108-10-1		
12.032	12.032	(1.298)	58	49522	2.00000	2.067	70.00-	130.00	100.00(M)
12.032	12.032	(1.298)	43	123001			0.00-	30.00	248.38
12.032	12.032	(1.298)	85	21457			0.00-	30.00	43.33

105	Toluene				CAS #:		108-88-3		
12.253	12.253	(1.322)	91	186889	2.00000	2.000	70.00-	130.00	100.00
12.253	12.253	(1.322)	92	110346			30.46-	90.46	59.04

108	trans-1,3-Dichloropropene				CAS #:		10061-02-6		
12.861	12.861	(0.882)	75	84129	2.00000	1.936	70.00-	130.00	100.00
12.861	12.861	(0.882)	77	21596			0.11-	60.11	25.67
12.834	12.834	(0.880)	39	40030			15.75-	75.75	47.58

110	1,1,2-Trichloroethane				CAS #:		79-00-5		
13.138	13.138	(0.901)	97	63448	2.00000	1.990	70.00-	130.00	100.00
13.138	13.138	(0.901)	99	41614			31.47-	91.47	65.59
13.138	13.138	(0.901)	83	50435			58.25-	118.25	79.49

112	Tetrachloroethene				CAS #:		127-18-4		
13.193	13.193	(0.905)	166	89856	2.00000	2.083	70.00-	130.00	100.00
13.193	13.193	(0.905)	129	69545			45.90-	105.90	77.40
13.193	13.193	(0.905)	131	66172			43.88-	103.88	73.64

114	2-Hexanone				CAS #:		591-78-6		
13.580	13.580	(0.932)	58	52105	2.00000	1.650	70.00-	130.00	100.00(a)
13.580	13.580	(0.932)	43	110802			147.03-	207.03	212.65
13.580	13.580	(0.932)	100	12762			0.00-	30.00	24.49

116	Dibromochloromethane				CAS #:		124-48-1		
13.718	13.718	(0.941)	129	95939	2.00000	1.920	70.00-	130.00	100.00
13.718	13.718	(0.941)	127	73156			0.00-	30.00	76.25

117	1,2-Dibromoethane				CAS #:		106-93-4		
13.884	13.884	(0.953)	107	96620	2.00000	1.989	70.00-	130.00	100.00
13.884	13.884	(0.953)	109	96596			65.18-	125.18	99.98

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
126 Chlorobenzene						CAS #:	108-90-7			
14.631	14.631	(1.004)	112	172950	2.00000	2.247	70.00-	130.00	100.00	
14.631	14.631	(1.004)	114	49642			1.91-	61.91	28.70	
14.603	14.603	(1.002)	77	98312			26.50-	86.50	56.84	

129 Ethyl Benzene						CAS #:	100-41-4			
14.769	14.769	(1.013)	106	80000	2.00000	1.964	70.00-	130.00	100.00	
14.769	14.769	(1.013)	91	248489			0.00-	30.00	310.61	

130 m,p-Xylene						CAS #:	108-38-3			
14.935	14.935	(1.025)	106	101958	2.00000	2.054	70.00-	130.00	100.00	
14.935	14.935	(1.025)	91	191884			0.00-	30.00	188.20	

132 o-Xylene						CAS #:	95-47-6			
15.488	15.488	(1.063)	106	101432	2.00000	2.025	70.00-	130.00	100.00	
15.488	15.488	(1.063)	91	208881			177.47-	237.47	205.93	

134 Styrene						CAS #:	100-42-5			
15.516	15.516	(1.064)	104	133562	2.00000	1.870	70.00-	130.00	100.00	
15.516	15.516	(1.064)	78	71778			20.25-	80.25	53.74	

135 Bromoform						CAS #:	75-25-2			
15.764	15.764	(1.082)	173	78478	2.00000	1.786	70.00-	130.00	100.00	
15.764	15.764	(1.082)	171	40496			21.05-	81.05	51.60	

137 Cumene						CAS #:	98-82-8			
15.958	15.958	(1.095)	105	282468	2.00000	2.063	70.00-	130.00	100.00	
15.958	15.958	(1.095)	120	74218			0.00-	30.00	26.27	
15.958	15.958	(1.095)	51	28040			0.00-	30.00	9.93	

144 1,1,2,2-Tetrachloroethane						CAS #:	79-34-5			
16.456	16.456	(1.129)	83	148008	2.00000	2.020	70.00-	130.00	100.00	
16.456	16.456	(1.129)	85	94627			31.99-	91.99	63.93	

145 Propylbenzene						CAS #:	103-65-1			
16.483	16.483	(1.131)	91	328901	2.00000	2.097	70.00-	130.00	100.00	
16.483	16.483	(1.131)	120	75360			0.00-	30.00	22.91	
16.483	16.483	(1.131)	105	9519			0.00-	30.00	2.89	

147 4-Ethyltoluene						CAS #:	622-96-8			
16.649	16.649	(1.142)	105	271508	2.00000	2.019	70.00-	130.00	100.00	
16.649	16.649	(1.142)	120	87836			0.00-	59.60	32.35	

148 1,3,5-Trimethylbenzene						CAS #:	108-67-8			
16.732	16.732	(1.148)	105	279904	2.00000	2.166	70.00-	130.00	100.00	
16.732	16.732	(1.148)	120	137880			0.00-	30.00	49.26	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

153	17.147	17.147	(1.176)	105	275228	2.00000	2.116	70.00-	130.00	100.00
	17.147	17.147	(1.176)	120	129173			15.41-	75.41	46.93
	CAS #: 95-63-6									

156	17.451	17.451	(1.197)	146	167109	2.00000	1.971	70.00-	130.00	100.00
	17.451	17.451	(1.197)	148	106537			0.00-	30.00	63.75
	17.451	17.451	(1.197)	111	73203			0.00-	30.00	43.81
	CAS #: 541-73-1									

157	17.562	17.562	(1.205)	146	138434	2.00000	1.722	70.00-	130.00	100.00
	17.562	17.562	(1.205)	148	127853			0.00-	30.00	92.36
	17.562	17.562	(1.205)	111	69398			0.00-	30.00	50.13
	CAS #: 106-46-7									

158	17.700	17.700	(1.214)	91	230635	2.00000	1.974	70.00-	130.00	100.00
	17.700	17.700	(1.214)	126	47342			0.00-	30.00	20.53
	CAS #: 100-44-7									

161	17.921	17.921	(1.230)	146	165520	2.00000	2.026	70.00-	130.00	100.00
	17.921	17.921	(1.230)	148	106977			32.70-	92.70	64.63
	17.921	17.921	(1.230)	111	68354			7.07-	67.07	41.30
	CAS #: 95-50-1									

167	19.276	19.276	(1.322)	180	177945	2.00000	2.155	70.00-	130.00	100.00
	19.276	19.276	(1.322)	182	166734			65.19-	125.19	93.70
	CAS #: 120-82-1									

168	19.359	19.359	(1.328)	225	114250	2.00000	2.411	70.00-	130.00	100.00
	19.359	19.359	(1.328)	223	74622			33.26-	93.26	65.31
	CAS #: 87-68-3									

169	19.470	19.470	(1.336)	128	467801	2.00000	2.687	70.00-	130.00	100.00
	19.470	19.470	(1.336)	127	67084			0.00-	30.00	14.34
	CAS #: 91-20-3									

QC Flag Legend

- a - Target compound detected but, quantitated amount Below Limit Of Quantitation(BLOQ).
- M - Compound response manually integrated.

Report Date: 31-May-2007 14:52

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053005.d

Calibration Time: 16:03

Lab Smp Id: ICAL

Client Smp ID: Level 3

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-2.0ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	441133	264680	617586	444018	0.65
88 1,4-Difluorobenze	1992312	1195387	2789237	2035031	2.14
125 Chlorobenzene-d5	1475337	885202	2065472	1499190	1.62

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

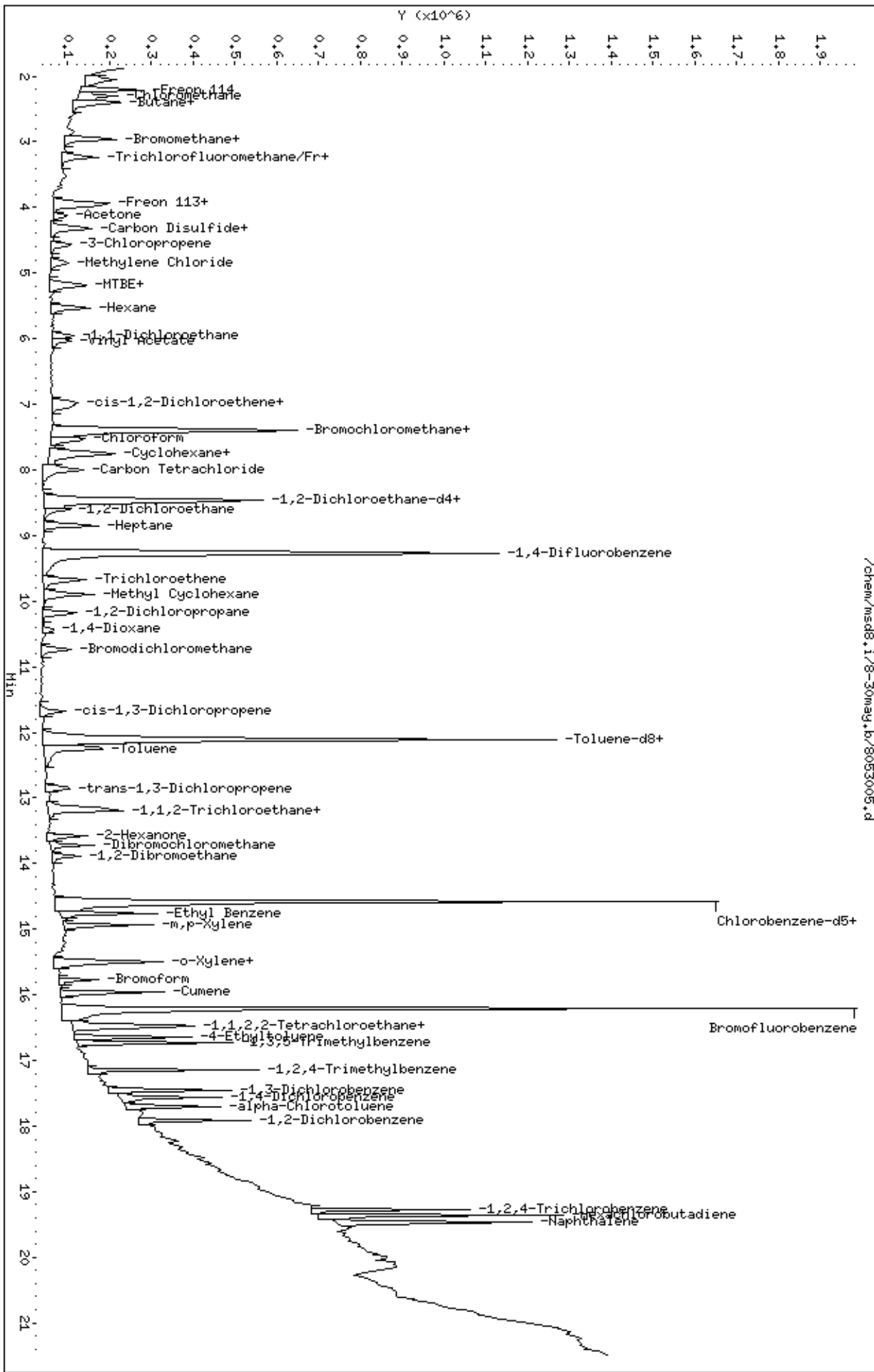
AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-30may.b/8053005.d
 Date: 30-May-2007 15:07
 Client ID: Level 3
 Sample Info: 2.0ml #1487-289
 Column phase: RTX-624

Instrument: msd8.1
 Operator: db
 Column diameter: 0.53



Report Date: 01-Jun-2007 13:51

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053006.d
 Lab Smp Id: ICAL Client Smp ID: Level 4
 Inj Date : 30-MAY-2007 15:35
 Operator : db Inst ID: msd8.i
 Smp Info : 25ml #1487-289
 Misc Info : 200ppbv-25ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 01-Jun-2007 10:56 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 15:35 Cal File: 8053006.d
 Als bottle: 1 Calibration Sample, Level: 4
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04mdl+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS									
		CAL-AMT		ON-COL		TARGET RANGE		RATIO	
RT	EXP RT (REL RT)	MASS	RESPONSE (PPBV)	(PPBV)					
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.415 (1.000)	130	442807 25.0000			80.00- 120.00		100.00	
7.387	7.415 (1.000)	128	348882			48.60- 108.60		78.79	
7.387	7.415 (1.000)	49	642051			114.98- 174.98		145.00	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267 (1.000)	114	2029767 25.0000			80.00- 120.00		100.00	
9.267	9.267 (1.000)	88	305336			0.00- 45.16		15.04	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576 (1.000)	117	1539410 25.0000			80.00- 120.00		100.00	
14.576	14.576 (1.000)	82	880295			0.00- 30.00		57.18	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465 (1.146)	65	557163 25.0000	24.041		80.00- 120.00		100.00	
8.465	8.465 (1.146)	67	328511			0.00- 30.00		58.96	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115 (1.307)	98	1689996 25.0000	24.050		80.00- 120.00		100.00	
12.115	12.115 (1.307)	70	173996			0.00- 30.00		10.30	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
\$ 104 Toluene-d8 (continued)										
12.115	12.115	(1.307)	100	1337051			0.00- 30.00	79.12		

\$ 140 Bromofluorobenzene										
						CAS #: 460-00-4				
16.207	16.207	(1.112)	174	923372	25.0000	25.210	80.00- 120.00	100.00		
16.207	16.207	(1.112)	95	1221505			105.65- 165.65	132.29		
16.207	16.207	(1.112)	176	870157			66.34- 126.34	94.24		

3 Propylene						CAS #: 115-07-1				
1.995	1.995	(0.270)	41	583695	25.0000	24.354	70.00- 130.00	100.00		
1.995	1.995	(0.270)	42	385435			0.00- 30.00	66.03		
1.995	1.995	(0.270)	39	385271			0.00- 30.00	66.01		

4 Dichlorodifluoromethane/Fr12						CAS #: 75-71-8				
2.050	2.078	(0.278)	85	1479581	25.0000	24.389	80.00- 120.00	100.00		
2.050	2.078	(0.278)	87	460934			0.00- 30.00	31.15		

6 Freon 114						CAS #: 76-14-2				
2.161	2.189	(0.293)	135	1377957	25.0000	25.142	80.00- 120.00	100.00		
2.161	2.189	(0.293)	137	436681			0.71- 60.71	31.69		

8 Chloromethane						CAS #: 74-87-3				
2.272	2.299	(0.308)	50	730257	25.0000	26.138	80.00- 120.00	100.00		
2.272	2.299	(0.308)	52	232404			0.00- 30.00	31.82		

9 Butane						CAS #: 106-97-8				
2.327	2.355	(0.315)	58	190942	25.0000	26.170	70.00- 130.00	100.00		
2.327	2.355	(0.315)	43	1372202			0.00- 30.00	718.65		

11 Vinyl Chloride						CAS #: 75-01-4				
2.410	2.438	(0.326)	62	788546	25.0000	24.290	80.00- 120.00	100.00		
2.410	2.438	(0.326)	64	250595			0.00- 30.00	31.78		

10 1,3-Butadiene						CAS #: 106-99-0				
2.382	2.410	(0.322)	54	644846	25.0000	23.222	80.00- 120.00	100.00		
2.382	2.410	(0.322)	39	672011			0.00- 30.00	104.21		

13 Bromomethane						CAS #: 74-83-9				
2.852	2.880	(0.386)	94	560666	25.0000	25.851	80.00- 120.00	100.00		
2.852	2.880	(0.386)	96	514411			63.65- 123.65	91.75		

16 Chloroethane						CAS #: 75-00-3				
2.935	2.991	(0.397)	64	435981	25.0000	25.376	80.00- 120.00	100.00		
2.935	2.991	(0.397)	49	106716			0.00- 30.00	24.48		
2.935	2.991	(0.397)	66	126764			0.00- 30.00	29.08		

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
15 Isopentane						CAS #: 78-78-4			
2.935	2.963	(0.397)	43	1053632	25.0000	25.992	70.00- 130.00	100.00	
2.963	2.963	(0.401)	57	687436			0.00- 30.00	65.24	
2.963	2.963	(0.401)	72	78712			0.00- 30.00	7.47	

18 Trichlorofluoromethane/Fr11						CAS #: 75-69-4			
3.212	3.239	(0.435)	101	1753350	25.0000	25.428	80.00- 120.00	100.00	
3.212	3.239	(0.435)	103	1132129			34.50- 94.50	64.57	

23 Ethanol						CAS #: 64-17-5			
3.516	3.544	(0.476)	45	293783	25.0000	26.681	80.00- 120.00	100.00	
3.516	3.544	(0.476)	43	57563			0.00- 30.00	19.59	
3.516	3.544	(0.476)	46	113176			0.00- 30.00	38.52	

28 Freon 113						CAS #: 76-13-1			
3.930	3.958	(0.532)	151	1151774	25.0000	25.246	80.00- 120.00	100.00	
3.930	3.958	(0.532)	153	703274			34.32- 94.32	61.06	
3.930	3.958	(0.532)	101	1334210			85.50- 145.50	115.84	

29 1,1-Dichloroethene						CAS #: 75-35-4			
3.958	3.986	(0.536)	61	1137998	25.0000	25.117	80.00- 120.00	100.00	
3.958	3.986	(0.536)	96	688816			29.81- 89.81	60.53	
3.958	3.986	(0.536)	98	424092			9.00- 69.00	37.27	

30 Acetone						CAS #: 67-64-1			
4.124	4.124	(0.558)	58	378307	25.0000	24.930	80.00- 120.00	100.00	
4.124	4.124	(0.558)	43	1131658			0.00- 30.00	299.14	

33 Carbon Disulfide						CAS #: 75-15-0			
4.290	4.290	(0.581)	76	2121278	25.0000	24.982	80.00- 120.00	100.00	

34 2-Propanol						CAS #: 67-63-0			
4.318	4.318	(0.584)	45	1370148	25.0000	24.973	80.00- 120.00	100.00	
4.318	4.318	(0.584)	43	262313			0.00- 30.00	19.14	
4.318	4.318	(0.584)	59	54411			0.00- 30.00	3.97	

37 3-Chloropropene						CAS #: 107-05-1			
4.566	4.594	(0.618)	76	360688	25.0000	26.626	80.00- 120.00	100.00	
4.566	4.594	(0.618)	41	1082431			0.00- 30.00	300.10	

40 Methylene Chloride						CAS #: 75-09-2			
4.815	4.815	(0.652)	49	845544	25.0000	24.927	80.00- 120.00	100.00	
4.815	4.815	(0.652)	84	605378			40.47- 100.47	71.60	
4.815	4.815	(0.652)	51	259145			0.00- 30.00	30.65	

43 MTBE						CAS #: 1634-04-4			
5.147	5.175	(0.697)	73	1407624	25.0000	29.179	80.00- 120.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
43 MTBE (continued)									
5.147	5.175	(0.697)	57	327200			0.00- 52.63	23.24	
5.147	5.175	(0.697)	41	310806			0.00- 30.00	22.08	

45 trans-1,2-Dichloroethene					CAS #: 156-60-5				
5.175	5.203	(0.701)	96	762528	25.0000	23.784	80.00- 120.00	100.00	
5.175	5.203	(0.701)	61	1150624			119.47- 179.47	150.90	
5.175	5.203	(0.701)	98	496217			0.00- 30.00	65.08	

46 Hexane					CAS #: 110-54-3				
5.534	5.534	(0.749)	57	1205663	25.0000	24.795	80.00- 120.00	100.00	
5.534	5.534	(0.749)	43	773376			0.00- 30.00	64.15	
5.534	5.534	(0.749)	86	198943			0.00- 30.00	16.50	

54 1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	1328399	25.0000	25.417	80.00- 120.00	100.00	
5.949	5.949	(0.805)	65	419788			1.86- 61.86	31.60	

55 Vinyl Acetate					CAS #: 108-05-4				
6.032	6.032	(0.817)	86	186162	25.0000	25.232	70.00- 130.00	100.00	
6.032	6.032	(0.817)	43	2005122			0.00- 30.00	1077.08	
6.032	6.032	(0.817)	42	151750			0.00- 30.00	81.52	

64 cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.972	6.972	(0.944)	61	1009566	25.0000	24.407	80.00- 120.00	100.00	
6.972	6.972	(0.944)	96	744531			43.20- 103.20	73.75	
6.972	6.972	(0.944)	98	478476			17.85- 77.85	47.39	

65 2-Butanone					CAS #: 78-93-3				
7.027	7.027	(0.951)	72	348381	25.0000	23.740	80.00- 120.00	100.00	
7.027	7.027	(0.951)	43	1511122			409.65- 469.65	433.76	
7.027	7.027	(0.951)	57	115798			0.00- 30.00	33.24	

67 Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.415	(1.000)	42	937522	25.0000	24.579	80.00- 120.00	100.00	
7.387	7.415	(1.000)	71	331869			5.48- 65.48	35.40	
7.414	7.415	(1.004)	72	366437			0.00- 30.00	39.09	

70 Chloroform					CAS #: 67-66-3				
7.525	7.553	(1.019)	83	1299284	25.0000	24.858	80.00- 120.00	100.00	
7.525	7.553	(1.019)	85	810995			32.67- 92.67	62.42	

73 Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	1057678	25.0000	24.389	80.00- 120.00	100.00	
7.746	7.746	(1.049)	56	1367225			96.25- 156.25	129.27	
7.746	7.746	(1.049)	41	682191			32.28- 92.28	64.50	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

75	1,1,1-Trichloroethane					CAS #:	71-55-6			
7.774	7.774	(1.052)	97	1343871	25.0000	24.539	80.00-	120.00	100.00	
7.774	7.774	(1.052)	99	876393			34.55-	94.55	65.21	

77	Carbon Tetrachloride					CAS #:	56-23-5			
7.995	8.023	(1.082)	119	1303695	25.0000	26.160	80.00-	120.00	100.00	
7.995	8.023	(1.082)	117	1329642			73.30-	133.30	101.99	

81	Benzene					CAS #:	71-43-2			
8.437	8.438	(0.910)	78	2220324	25.0000	24.214	80.00-	120.00	100.00	
8.437	8.438	(0.910)	77	477752			0.00-	30.00	21.52	

80	2,2,4-Trimethylpentane					CAS #:	540-84-1			
8.465	8.465	(1.146)	57	3208710	25.0000	24.784	80.00-	120.00	100.00	
8.465	8.465	(1.146)	56	1043397			0.00-	30.00	32.52	
8.465	8.465	(1.146)	41	660582			0.00-	30.00	20.59	

83	1,2-Dichloroethane					CAS #:	107-06-2			
8.603	8.603	(0.928)	62	912402	25.0000	24.525	80.00-	120.00	100.00	
8.603	8.603	(0.928)	64	284806			0.00-	30.00	31.21	

85	Heptane					CAS #:	142-82-5			
8.852	8.852	(0.955)	100	223706	25.0000	22.544	80.00-	120.00	100.00	
8.852	8.852	(0.955)	43	1246808			0.00-	30.00	557.34	
8.852	8.852	(0.955)	71	718610			0.00-	30.00	321.23	

94	Trichloroethene					CAS #:	79-01-6			
9.682	9.682	(1.045)	95	899732	25.0000	24.760	80.00-	120.00	100.00	
9.682	9.682	(1.045)	130	950218			78.57-	138.57	105.61	
9.682	9.682	(1.045)	97	567162			34.08-	94.08	63.04	

95	Methyl Cyclohexane					CAS #:	108-87-2			
9.903	9.903	(1.341)	83	1353516	25.0000	25.052	70.00-	130.00	100.00	
9.903	9.903	(1.341)	98	623110			0.00-	30.00	46.04	
9.903	9.903	(1.341)	55	1057240			0.00-	30.00	78.11	

97	1,2-Dichloropropane					CAS #:	78-87-5			
10.179	10.179	(1.098)	63	774557	25.0000	24.374	80.00-	120.00	100.00	
10.179	10.179	(1.098)	62	537358			42.26-	102.26	69.38	
10.179	10.179	(1.098)	41	444960			26.77-	86.77	57.45	

98	1,4-Dioxane					CAS #:	123-91-1			
10.428	10.428	(1.125)	88	491146	25.0000	25.340	80.00-	120.00	100.00	
10.428	10.428	(1.125)	58	356106			41.45-	101.45	72.51	
10.428	10.428	(1.125)	57	107866			0.00-	30.00	21.96	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

100 Bromodichloromethane						CAS #: 75-27-4			
10.732	10.732	(1.158)	83	1309837	25.0000	24.679	80.00- 120.00	100.00	
10.732	10.732	(1.158)	85	812676			31.75- 91.75	62.04	

102 cis-1,3-Dichloropropene						CAS #: 10061-01-5			
11.672	11.673	(1.260)	75	1088954	25.0000	24.658	80.00- 120.00	100.00	
11.672	11.673	(1.260)	77	332460			1.36- 61.36	30.53	
11.672	11.673	(1.260)	39	522028			17.70- 77.70	47.94	

103 4-Methyl-2-pentanone						CAS #: 108-10-1			
12.032	12.032	(1.298)	58	595289	25.0000	24.917	80.00- 120.00	100.00	
12.032	12.032	(1.298)	43	1462778			0.00- 30.00	245.73	
12.032	12.032	(1.298)	85	250389			0.00- 30.00	42.06	

105 Toluene						CAS #: 108-88-3			
12.253	12.253	(1.322)	91	2271771	25.0000	24.377	80.00- 120.00	100.00	
12.253	12.253	(1.322)	92	1302497			29.69- 89.69	57.33	

108 trans-1,3-Dichloropropene						CAS #: 10061-02-6			
12.834	12.861	(0.880)	75	1101495	25.0000	24.681	80.00- 120.00	100.00	
12.834	12.861	(0.880)	77	341616			0.08- 60.08	31.01	
12.834	12.861	(0.880)	39	503742			15.97- 75.97	45.73	

110 1,1,2-Trichloroethane						CAS #: 79-00-5			
13.138	13.138	(0.901)	97	785392	25.0000	23.996	80.00- 120.00	100.00	
13.138	13.138	(0.901)	99	476630			29.85- 89.85	60.69	
13.138	13.138	(0.901)	83	661660			54.93- 114.93	84.25	

112 Tetrachloroethene						CAS #: 127-18-4			
13.193	13.193	(0.905)	166	1075849	25.0000	24.286	80.00- 120.00	100.00	
13.193	13.193	(0.905)	129	832653			45.08- 105.08	77.39	
13.193	13.193	(0.905)	131	811520			44.22- 104.22	75.43	

114 2-Hexanone						CAS #: 591-78-6			
13.580	13.580	(0.932)	58	818770	25.0000	25.246	80.00- 120.00	100.00	
13.580	13.580	(0.932)	43	1453939			148.62- 208.62	177.58	
13.580	13.580	(0.932)	100	165065			0.00- 30.00	20.16	

116 Dibromochloromethane						CAS #: 124-48-1			
13.718	13.719	(0.941)	129	1251408	25.0000	24.396	80.00- 120.00	100.00	
13.718	13.719	(0.941)	127	988822			0.00- 30.00	79.02	

117 1,2-Dibromoethane						CAS #: 106-93-4			
13.884	13.884	(0.953)	107	1209824	25.0000	24.258	80.00- 120.00	100.00	
13.884	13.884	(0.953)	109	1141289			63.81- 123.81	94.34	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
126 Chlorobenzene						CAS #:	108-90-7			
14.603	14.631	(1.002)	112	1910705	25.0000	24.172	80.00-	120.00	100.00	
14.631	14.631	(1.004)	114	623141			1.84-	61.84	32.61	
14.603	14.631	(1.002)	77	1069076			26.78-	86.78	55.95	

129 Ethyl Benzene						CAS #:	100-41-4			
14.769	14.769	(1.013)	106	987199	25.0000	23.600	80.00-	120.00	100.00	
14.769	14.769	(1.013)	91	3031265			0.00-	30.00	307.06	

130 m,p-Xylene						CAS #:	108-38-3			
14.935	14.935	(1.025)	106	1223079	25.0000	23.994	80.00-	120.00	100.00	
14.935	14.935	(1.025)	91	2365743			0.00-	30.00	193.43	

132 o-Xylene						CAS #:	95-47-6			
15.488	15.488	(1.063)	106	1240684	25.0000	24.120	80.00-	120.00	100.00	
15.488	15.488	(1.063)	91	2526182			178.74-	238.74	203.61	

134 Styrene						CAS #:	100-42-5			
15.516	15.516	(1.064)	104	1736719	25.0000	23.686	80.00-	120.00	100.00	
15.516	15.516	(1.064)	78	866978			20.27-	80.27	49.92	

135 Bromoform						CAS #:	75-25-2			
15.764	15.765	(1.082)	173	1132279	25.0000	25.095	80.00-	120.00	100.00	
15.764	15.765	(1.082)	171	579428			20.67-	80.67	51.17	

137 Cumene						CAS #:	98-82-8			
15.958	15.958	(1.095)	105	3402095	25.0000	24.198	80.00-	120.00	100.00	
15.958	15.958	(1.095)	120	915816			0.00-	30.00	26.92	
15.958	15.958	(1.095)	51	322973			0.00-	30.00	9.49	

144 1,1,2,2-Tetrachloroethane						CAS #:	79-34-5			
16.456	16.456	(1.129)	83	1774511	25.0000	23.590	80.00-	120.00	100.00	
16.456	16.456	(1.129)	85	1091597			31.59-	91.59	61.52	

145 Propylbenzene						CAS #:	103-65-1			
16.483	16.484	(1.131)	91	4041657	25.0000	25.100	80.00-	120.00	100.00	
16.483	16.484	(1.131)	120	954510			0.00-	30.00	23.62	
16.483	16.484	(1.131)	105	145225			0.00-	30.00	3.59	

147 4-Ethyltoluene						CAS #:	622-96-8			
16.649	16.649	(1.142)	105	3616568	25.0000	26.197	80.00-	120.00	100.00	
16.649	16.649	(1.142)	120	1114974			0.00-	59.85	30.83	

148 1,3,5-Trimethylbenzene						CAS #:	108-67-8			
16.732	16.732	(1.148)	105	3346408	25.0000	25.218	80.00-	120.00	100.00	
16.732	16.732	(1.148)	120	1670014			0.00-	30.00	49.90	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

153	17.147	17.147 (1.176)	105	3467382	25.0000	25.959	80.00- 120.00	100.00	
	17.147	17.147 (1.176)	120	1561865			15.22- 75.22	45.04	

156	17.451	17.451 (1.197)	146	2193262	25.0000	25.189	80.00- 120.00	100.00	
	17.451	17.451 (1.197)	148	1371901			0.00- 30.00	62.55	
	17.451	17.451 (1.197)	111	918858			0.00- 30.00	41.89	

157	17.562	17.562 (1.205)	146	1979487	25.0000	23.988	80.00- 120.00	100.00	
	17.562	17.562 (1.205)	148	1208680			0.00- 30.00	61.06	
	17.562	17.562 (1.205)	111	686076			0.00- 30.00	34.66	

158	17.700	17.700 (1.214)	91	3074757	25.0000	25.634	80.00- 120.00	100.00	
	17.700	17.700 (1.214)	126	653675			0.00- 30.00	21.26	

161	17.921	17.921 (1.230)	146	1978729	25.0000	23.591	80.00- 120.00	100.00	
	17.921	17.921 (1.230)	148	1274964			33.37- 93.37	64.43	
	17.921	17.921 (1.230)	111	743013			7.65- 67.65	37.55	

167	19.276	19.276 (1.322)	180	1941314	25.0000	22.900	80.00- 120.00	100.00	
	19.276	19.276 (1.322)	182	1852644			64.68- 124.68	95.43	

168	19.359	19.359 (1.328)	225	1156590	25.0000	23.769	80.00- 120.00	100.00	
	19.359	19.359 (1.328)	223	748967			33.22- 93.22	64.76	

169	19.469	19.470 (1.336)	128	4917090	25.0000	27.503	80.00- 120.00	100.00	
	19.469	19.470 (1.336)	127	601613			0.00- 30.00	12.24	

Report Date: 01-Jun-2007 13:51

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053006.d

Calibration Time: 15:35

Lab Smp Id: ICAL

Client Smp ID: Level 4

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-25ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	448309	268985	627633	442807	-1.23
88 1,4-Difluorobenze	2033490	1220094	2846886	2029767	-0.18
125 Chlorobenzene-d5	1524596	914758	2134434	1539410	0.97

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

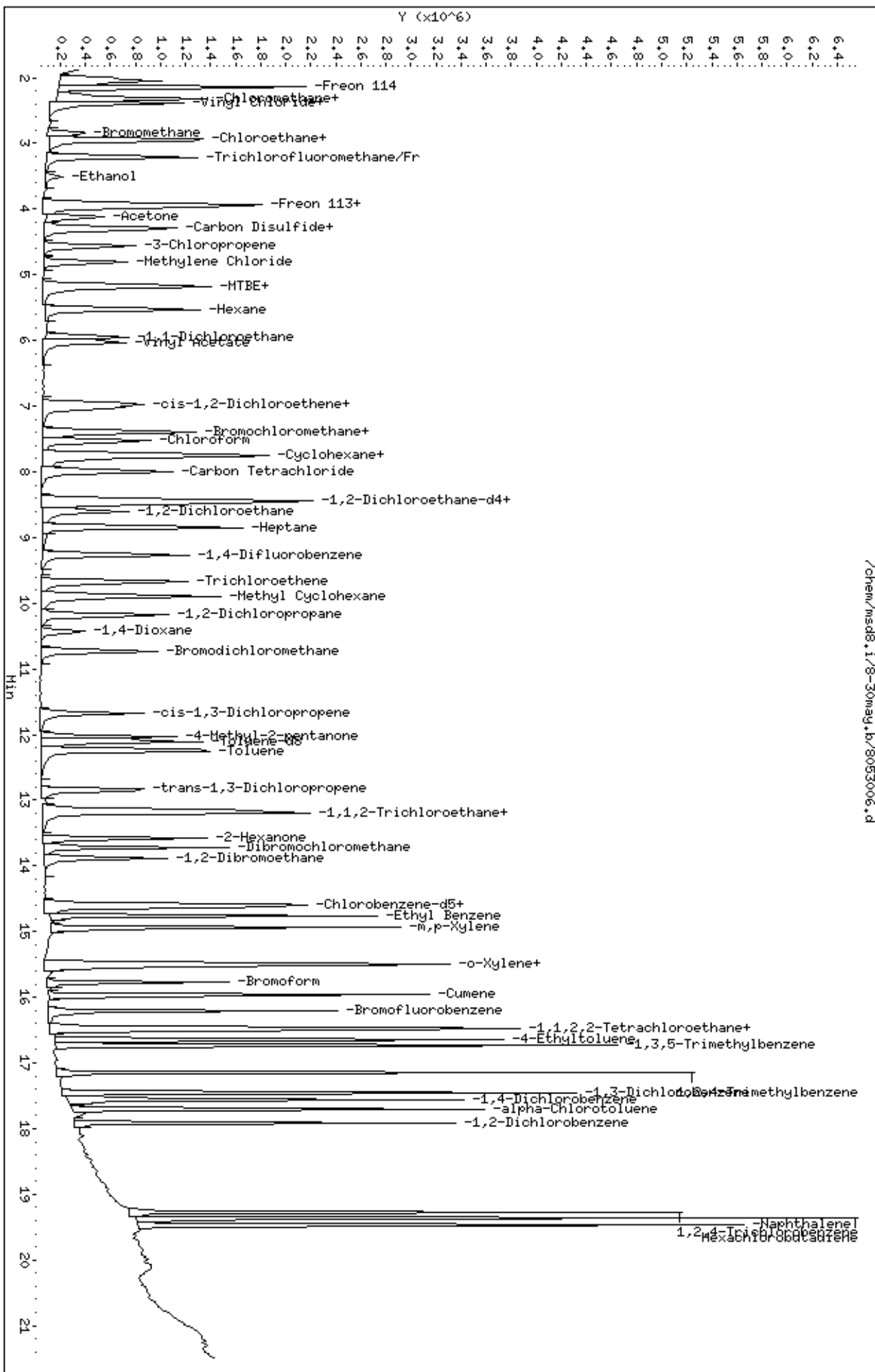
RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-30may.b/8053006.d
Date: 30-May-2007 15:35
Client ID: Level 4
Sample Info: 25ml #1487-289

Column phase: RTX-624

Instrument: msd8.1
Operator: db
Column diameter: 0.53



Report Date: 07-Jun-2007 13:41

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-07jun.b/8060705.d
 Lab Smp Id: ICAL Client Smp ID: Level 5
 Inj Date : 07-JUN-2007 11:37
 Operator : JG Inst ID: msd8.i
 Smp Info : 50ml #1443-96
 Misc Info : 200ppbv-50ppbv
 Comment :
 Method : /chem/msd8.i/8-07jun.b/t14q530b.m
 Meth Date : 07-Jun-2007 13:41 jgray Quant Type: ISTD
 Cal Date : 07-JUN-2007 11:37 Cal File: 8060705.d
 Als bottle: 1 Calibration Sample, Level: 5
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: sp16b.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable

Local Compound Variable

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	CAL-AMT	ON-COL	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	350593	25.0000			80.00- 120.00	100.00
7.387	7.387	(1.000)	128	255876				42.98- 102.98	72.98
7.387	7.387	(1.000)	49	499697				112.53- 172.53	142.53

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	1524282	25.0000			80.00- 120.00	100.00
9.267	9.267	(1.000)	88	222191				0.00- 44.58	14.58

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	1168126	25.0000			80.00- 120.00	100.00
14.576	14.576	(1.000)	82	655064				26.08- 86.08	56.08

1 Freon 152a CAS #: 75-37-6									
1.995	1.995	(0.270)	65	553785	50.0000	45.598		80.00- 120.00	100.00
2.050	2.050	(0.278)	51	2608865				441.10- 501.10	471.10

20 Freon123a CAS #: 354-23-4									
3.682	3.682	(0.498)	67	930646	50.0000	49.628		80.00- 120.00	100.00
3.682	3.682	(0.498)	117	739080				49.42- 109.42	79.42

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
21 Freon123						CAS #: 306-83-2			
3.792	3.792	(0.513)	83	592112	50.0000	50.099	80.00- 120.00	100.00	
3.792	3.792	(0.513)	133	115001			0.00- 49.42	19.42	
3.792	3.792	(0.513)	85	439319			44.20- 104.20	74.20	

38 tert-Butyl-Alcohol						CAS #: 75-65-0			
4.954	4.954	(0.671)	59	1213562	50.0000	49.665	80.00- 120.00	100.00	
4.954	4.954	(0.671)	41	281263			0.00- 53.18	23.18	
4.954	4.954	(0.671)	57	119569			0.00- 39.85	9.85	

49 Isopropyl ether						CAS #: 108-20-3			
5.949	5.949	(0.805)	45	3071363	50.0000	50.259	80.00- 120.00	100.00	
5.949	5.949	(0.805)	87	793048			0.00- 55.82	25.82	
5.949	5.949	(0.805)	59	334819			0.00- 40.90	10.90	

52 1-Propanol						CAS #: 71-23-8			
6.170	6.170	(0.835)	42	208766	50.0000	47.398	80.00- 120.00	100.00	
6.170	6.170	(0.835)	59	279123			103.70- 163.70	133.70	
6.170	6.170	(0.835)	41	166970			49.98- 109.98	79.98	

58 Ethyl-tert-butyl Ether						CAS #: 637-92-3			
6.585	6.585	(0.891)	59	2332280	50.0000	57.162	80.00- 120.00	100.00	
6.585	6.585	(0.891)	87	900443			8.61- 68.61	38.61	
6.585	6.585	(0.891)	41	405190			0.00- 47.37	17.37	

61 Ethyl Acetate						CAS #: 141-78-6			
7.083	7.083	(0.959)	70	262488	50.0000	49.298	80.00- 120.00	100.00	
7.083	7.083	(0.959)	43	2435365			897.80- 957.80	927.80	
7.083	7.083	(0.959)	61	345569			101.65- 161.65	131.65	

78 Isobutanol						CAS #: 78-83-1			
8.437	8.437	(0.910)	43	901547	50.0000	51.531	80.00- 120.00	100.00	
8.437	8.437	(0.910)	41	603783			36.97- 96.97	66.97	

79 tert-amyl-Methyl Ether						CAS #: 994-05-8			
8.631	8.631	(1.168)	73	2090649	50.0000	56.030	80.00- 120.00	100.00	
8.631	8.631	(1.168)	87	519620			0.00- 54.85	24.85	
8.631	8.631	(1.168)	55	584782			0.00- 57.97	27.97	

89 1-Butanol						CAS #: 71-36-3			
9.709	9.709	(1.048)	56	753331	50.0000	50.488	80.00- 120.00	100.00	
9.709	9.709	(1.048)	41	501173			36.53- 96.53	66.53	
9.709	9.709	(1.048)	43	411551			24.63- 84.63	54.63	

136 Cyclohexanone						CAS #: 108-94-1			
16.152	16.152	(1.108)	55	1332835	50.0000	49.854	80.00- 120.00	100.00	

AMOUNTS

RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO
16.152	16.152	(1.108)	98	595161			14.65- 74.65	44.65
16.124	16.124	(1.106)	42	881662			36.15- 96.15	66.15

136 Cyclohexanone (continued)

Report Date: 07-Jun-2007 13:41

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 07-JUN-2007

Lab File ID: 8060705.d

Calibration Time: 11:37

Lab Smp Id: ICAL

Client Smp ID: Level 5

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: JG

Method File: /chem/msd8.i/8-07jun.b/t14q530b.m

Misc Info: 200ppbv-50ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	350593	210356	490830	350593	0.00
88 1,4-Difluorobenze	1524282	914569	2133995	1524282	0.00
125 Chlorobenzene-d5	1168126	700876	1635376	1168126	0.00

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-07jun.b/8060705.d

Date: 07-JUN-2007 11:37

Client ID: Level 5

Sample Info: 50ml #1443-96

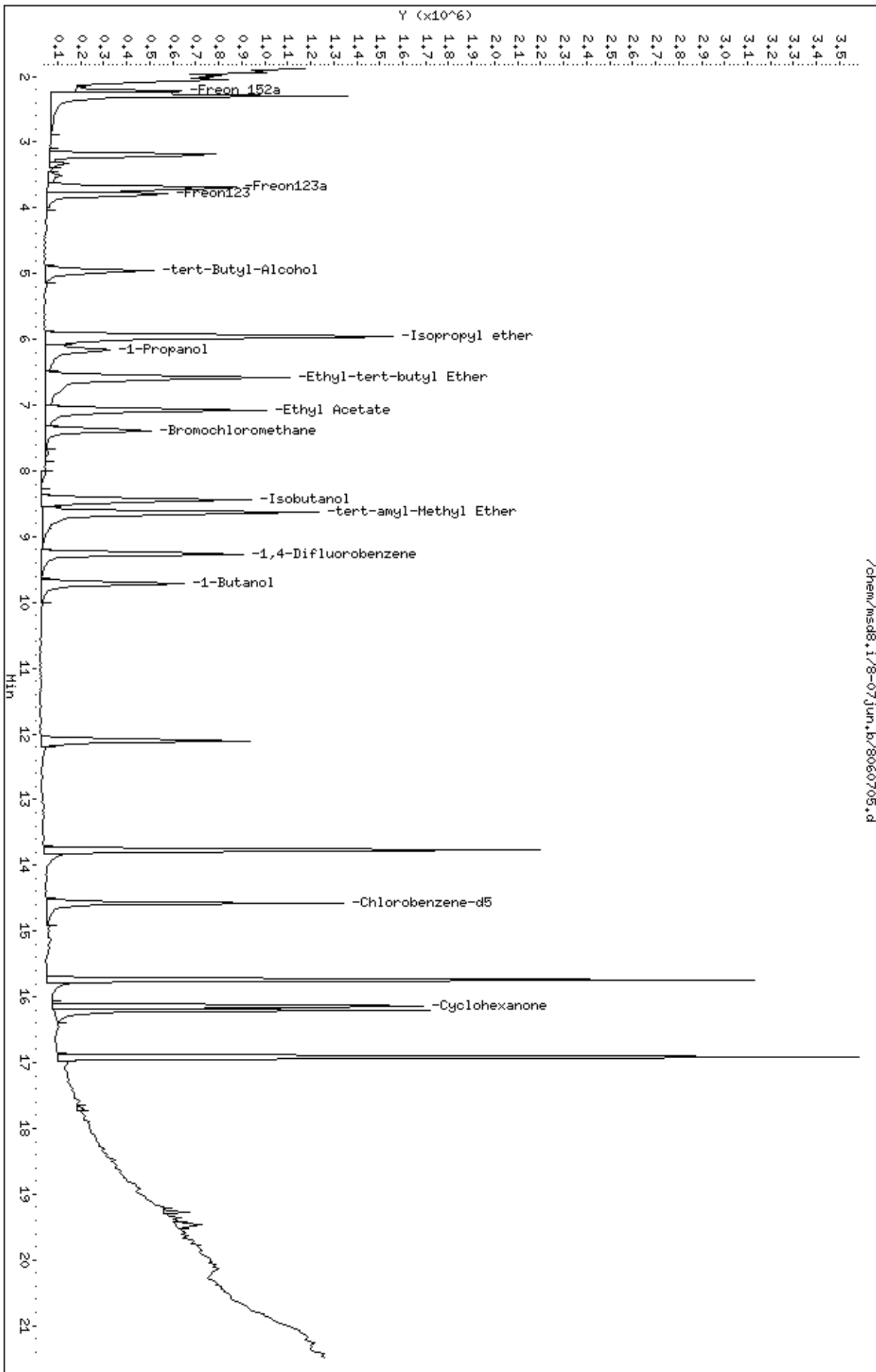
Column phase: RTX-624

Instrument: msd8.1

Operator: JG

Column diameter: 0.53

/chem/msd8.1/8-07jun.b/8060705.d



Report Date: 31-May-2007 14:53

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053007.d
 Lab Smp Id: ICAL Client Smp ID: Level 5
 Inj Date : 30-MAY-2007 16:03
 Operator : db Inst ID: msd8.i
 Smp Info : 50ml #1487-289
 Misc Info : 200ppbv-50ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 31-May-2007 14:53 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 16:03 Cal File: 8053007.d
 Als bottle: 1 Calibration Sample, Level: 5
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04mdl+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	(PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	441133	25.0000		80.00- 120.00	100.00	
7.387	7.387	(1.000)	128	342185			47.57- 107.57	77.57	
7.387	7.387	(1.000)	49	632872			113.47- 173.47	143.47	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	1992312	25.0000		80.00- 120.00	100.00	
9.267	9.267	(1.000)	88	312375			0.00- 45.68	15.68	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	1475337	25.0000		80.00- 120.00	100.00	
14.576	14.576	(1.000)	82	860714			28.34- 88.34	58.34	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465	(1.146)	65	557531	25.0000	24.148	80.00- 120.00	100.00	
8.465	8.465	(1.146)	67	334456			29.99- 89.99	59.99	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115	(1.307)	98	1710823	25.0000	24.804	80.00- 120.00	100.00	
12.115	12.115	(1.307)	70	174905			0.00- 40.22	10.22	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
\$ 104 Toluene-d8 (continued)									
12.115	12.115	(1.307)	100	1606916			63.93- 123.93	93.93	

\$ 140 Bromofluorobenzene									
						CAS #: 460-00-4			
16.207	16.207	(1.112)	174	903252	25.0000	25.732	80.00- 120.00	100.00	
16.207	16.207	(1.112)	95	1193763			102.16- 162.16	132.16	
16.207	16.207	(1.112)	176	851833			64.31- 124.31	94.31	

3 Propylene									
						CAS #: 115-07-1			
1.995	1.995	(0.270)	41	1060951	50.0000	44.436	80.00- 120.00	100.00	
1.995	1.995	(0.270)	42	724676			38.30- 98.30	68.30	
1.995	1.995	(0.270)	39	723770			38.22- 98.22	68.22	

4 Dichlorodifluoromethane/Fr12									
						CAS #: 75-71-8			
2.050	2.050	(0.278)	85	2666402	50.0000	44.120	80.00- 120.00	100.00	
2.050	2.050	(0.278)	87	867034			2.52- 62.52	32.52	

6 Freon 114									
						CAS #: 76-14-2			
2.161	2.161	(0.293)	135	2656241	50.0000	48.650	80.00- 120.00	100.00	
2.161	2.161	(0.293)	137	837548			1.53- 61.53	31.53	

8 Chloromethane									
						CAS #: 74-87-3			
2.272	2.272	(0.308)	50	1374103	50.0000	49.370	80.00- 120.00	100.00	
2.272	2.272	(0.308)	52	422135			0.72- 60.72	30.72	

9 Butane									
						CAS #: 106-97-8			
2.327	2.327	(0.315)	58	346263	50.0000	47.638	80.00- 120.00	100.00	
2.327	2.327	(0.315)	43	2556532			708.32- 768.32	738.32	

11 Vinyl Chloride									
						CAS #: 75-01-4			
2.410	2.410	(0.326)	62	1538485	50.0000	47.571	80.00- 120.00	100.00	
2.410	2.410	(0.326)	64	484291			1.48- 61.48	31.48	

10 1,3-Butadiene									
						CAS #: 106-99-0			
2.382	2.382	(0.322)	54	1245393	50.0000	45.019	80.00- 120.00	100.00	
2.382	2.382	(0.322)	39	1201450			66.47- 126.47	96.47	

13 Bromomethane									
						CAS #: 74-83-9			
2.852	2.852	(0.386)	94	1081074	50.0000	50.035	80.00- 120.00	100.00	
2.852	2.852	(0.386)	96	1027301			65.03- 125.03	95.03	

16 Chloroethane									
						CAS #: 75-00-3			
2.963	2.963	(0.401)	64	835185	50.0000	48.795	80.00- 120.00	100.00	
2.963	2.963	(0.401)	49	206572			0.00- 54.73	24.73	
2.963	2.963	(0.401)	66	259149			1.03- 61.03	31.03	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

15 Isopentane						CAS #: 78-78-4			
2.963	2.963	(0.401)	43	1993575	50.0000	49.367	80.00- 120.00	100.00	
2.963	2.963	(0.401)	57	1377018			39.07- 99.07	69.07	
2.963	2.963	(0.401)	72	149398			0.00- 37.49	7.49	

18 Trichlorofluoromethane/Fr11						CAS #: 75-69-4			
3.212	3.212	(0.435)	101	3338623	50.0000	48.602	80.00- 120.00	100.00	
3.212	3.212	(0.435)	103	2168945			34.97- 94.97	64.97	

23 Ethanol						CAS #: 64-17-5			
3.516	3.516	(0.476)	45	536367	50.0000	48.898	80.00- 120.00	100.00	
3.516	3.516	(0.476)	43	110348			0.00- 50.57	20.57	
3.516	3.516	(0.476)	46	225695			12.08- 72.08	42.08	

28 Freon 113						CAS #: 76-13-1			
3.931	3.931	(0.532)	151	2144041	50.0000	47.174	80.00- 120.00	100.00	
3.931	3.931	(0.532)	153	1365964			33.71- 93.71	63.71	
3.931	3.931	(0.532)	101	2494396			86.34- 146.34	116.34	

29 1,1-Dichloroethene						CAS #: 75-35-4			
3.958	3.958	(0.536)	61	2147006	50.0000	47.567	80.00- 120.00	100.00	
3.958	3.958	(0.536)	96	1297826			30.45- 90.45	60.45	
3.958	3.958	(0.536)	98	824285			8.39- 68.39	38.39	

30 Acetone						CAS #: 67-64-1			
4.124	4.124	(0.558)	58	743587	50.0000	49.188	80.00- 120.00	100.00	
4.124	4.124	(0.558)	43	2174812			262.48- 322.48	292.48	

33 Carbon Disulfide						CAS #: 75-15-0			
4.290	4.290	(0.581)	76	4010765	50.0000	47.413	80.00- 120.00	100.00	

34 2-Propanol						CAS #: 67-63-0			
4.318	4.318	(0.584)	45	2611678	50.0000	47.782	80.00- 120.00	100.00	
4.318	4.318	(0.584)	43	497434			0.00- 49.05	19.05	
4.318	4.318	(0.584)	59	105698			0.00- 34.05	4.05	

37 3-Chloropropene						CAS #: 107-05-1			
4.566	4.566	(0.618)	76	692146	50.0000	51.289	80.00- 120.00	100.00	
4.566	4.566	(0.618)	41	2087482			271.60- 331.60	301.60	

40 Methylene Chloride						CAS #: 75-09-2			
4.815	4.815	(0.652)	49	1613264	50.0000	47.741	80.00- 120.00	100.00	
4.815	4.815	(0.652)	84	1145605			41.01- 101.01	71.01	
4.815	4.815	(0.652)	51	477098			0.00- 59.57	29.57	

43 MTBE						CAS #: 1634-04-4			
5.147	5.147	(0.697)	73	2615636	50.0000	54.425	80.00- 120.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	CAL-AMT		ON-COL	TARGET RANGE	RATIO	
				RESPONSE	(PPEV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
43 MTBE (continued)									
5.147	5.147	(0.697)	57	623120			0.00- 53.82	23.82	
5.147	5.147	(0.697)	41	585377			0.00- 52.38	22.38	

45 trans-1,2-Dichloroethene					CAS #: 156-60-5				
5.175	5.175	(0.701)	96	1487814	50.0000	46.583	80.00- 120.00	100.00	
5.175	5.175	(0.701)	61	2217443			119.04- 179.04	149.04	
5.202	5.202	(0.704)	98	952041			33.99- 93.99	63.99	

46 Hexane					CAS #: 110-54-3				
5.534	5.534	(0.749)	57	2496765	50.0000	51.542	80.00- 120.00	100.00	
5.534	5.534	(0.749)	43	1570026			32.88- 92.88	62.88	
5.534	5.534	(0.749)	86	367327			0.00- 44.71	14.71	

54 1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	2554528	50.0000	49.064	80.00- 120.00	100.00	
5.949	5.949	(0.805)	65	797507			1.22- 61.22	31.22	

55 Vinyl Acetate					CAS #: 108-05-4				
6.032	6.032	(0.817)	86	355502	50.0000	48.367	80.00- 120.00	100.00	
6.032	6.032	(0.817)	43	3900404			1067.15-1127.15	1097.15	
6.032	6.032	(0.817)	42	297238			53.61- 113.61	83.61	

64 cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.972	6.972	(0.944)	61	1890254	50.0000	45.872	80.00- 120.00	100.00	
6.972	6.972	(0.944)	96	1419326			45.09- 105.09	75.09	
6.972	6.972	(0.944)	98	900856			17.66- 77.66	47.66	

65 2-Butanone					CAS #: 78-93-3				
7.027	7.027	(0.951)	72	682086	50.0000	46.657	80.00- 120.00	100.00	
7.027	7.027	(0.951)	43	2923118			398.56- 458.56	428.56	
7.027	7.027	(0.951)	57	230736			3.83- 63.83	33.83	

67 Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.387	(1.000)	42	1794674	50.0000	47.230	80.00- 120.00	100.00	
7.387	7.387	(1.000)	71	630736			5.14- 65.14	35.14	
7.387	7.387	(1.000)	72	683952			8.11- 68.11	38.11	

70 Chloroform					CAS #: 67-66-3				
7.525	7.525	(1.019)	83	2510438	50.0000	48.212	80.00- 120.00	100.00	
7.525	7.525	(1.019)	85	1556022			31.98- 91.98	61.98	

73 Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	2021584	50.0000	46.792	80.00- 120.00	100.00	
7.746	7.746	(1.049)	56	2613838			99.30- 159.30	129.30	
7.746	7.746	(1.049)	41	1290550			33.84- 93.84	63.84	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====	=====	

75	1,1,1-Trichloroethane					CAS #:	71-55-6			
7.774	7.774	(1.052)	97	2602285	50.0000	47.697	80.00-	120.00	100.00	
7.774	7.774	(1.052)	99	1684374			34.73-	94.73	64.73	

77	Carbon Tetrachloride					CAS #:	56-23-5			
7.995	7.995	(1.082)	119	2509316	50.0000	50.544	80.00-	120.00	100.00	
7.995	7.995	(1.082)	117	2610640			74.04-	134.04	104.04	

81	Benzene					CAS #:	71-43-2			
8.437	8.437	(0.910)	78	4299343	50.0000	47.769	80.00-	120.00	100.00	
8.437	8.437	(0.910)	77	953019			0.00-	52.17	22.17	

80	2,2,4-Trimethylpentane					CAS #:	540-84-1			
8.465	8.465	(1.146)	57	6116976	50.0000	47.428	80.00-	120.00	100.00	
8.465	8.465	(1.146)	56	1934694			1.63-	61.63	31.63	
8.465	8.465	(1.146)	41	1435732			0.00-	53.47	23.47	

83	1,2-Dichloroethane					CAS #:	107-06-2			
8.603	8.603	(0.928)	62	1714995	50.0000	46.965	80.00-	120.00	100.00	
8.603	8.603	(0.928)	64	541369			1.57-	61.57	31.57	

85	Heptane					CAS #:	142-82-5			
8.852	8.852	(0.955)	100	425691	50.0000	43.706	80.00-	120.00	100.00	
8.852	8.852	(0.955)	43	2479146			552.38-	612.38	582.38	
8.852	8.852	(0.955)	71	1375573			293.14-	353.14	323.14	

94	Trichloroethene					CAS #:	79-01-6			
9.682	9.682	(1.045)	95	1701628	50.0000	47.708	80.00-	120.00	100.00	
9.682	9.682	(1.045)	130	1834985			77.84-	137.84	107.84	
9.682	9.682	(1.045)	97	1092814			34.22-	94.22	64.22	

95	Methyl Cyclohexane					CAS #:	108-87-2			
9.903	9.903	(1.341)	83	2612830	50.0000	48.544	80.00-	120.00	100.00	
9.903	9.903	(1.341)	98	1224638			16.87-	76.87	46.87	
9.903	9.903	(1.341)	55	2044095			48.23-	108.23	78.23	

97	1,2-Dichloropropane					CAS #:	78-87-5			
10.179	10.179	(1.098)	63	1486513	50.0000	47.658	80.00-	120.00	100.00	
10.179	10.179	(1.098)	62	1072565			42.15-	102.15	72.15	
10.179	10.179	(1.098)	41	839449			26.47-	86.47	56.47	

98	1,4-Dioxane					CAS #:	123-91-1			
10.428	10.428	(1.125)	88	944170	50.0000	49.629	80.00-	120.00	100.00	
10.428	10.428	(1.125)	58	698982			44.03-	104.03	74.03	
10.428	10.428	(1.125)	57	215615			0.00-	52.84	22.84	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

100	Bromodichloromethane					CAS #:	75-27-4			
10.732	10.732	(1.158)	83	2547396	50.0000	48.899	80.00-	120.00	100.00	
10.732	10.732	(1.158)	85	1554700			31.03-	91.03	61.03	

102	cis-1,3-Dichloropropene					CAS #:	10061-01-5			
11.672	11.672	(1.260)	75	2114572	50.0000	48.783	80.00-	120.00	100.00	
11.672	11.672	(1.260)	77	659599			1.19-	61.19	31.19	
11.672	11.672	(1.260)	39	992332			16.93-	76.93	46.93	

103	4-Methyl-2-pentanone					CAS #:	108-10-1			
12.032	12.032	(1.298)	58	1183517	50.0000	50.470	80.00-	120.00	100.00	
12.032	12.032	(1.298)	43	2867386			212.28-	272.28	242.28	
12.032	12.032	(1.298)	85	492895			11.65-	71.65	41.65	

105	Toluene					CAS #:	108-88-3			
12.253	12.253	(1.322)	91	4422713	50.0000	48.350	80.00-	120.00	100.00	
12.253	12.253	(1.322)	92	2673976			30.46-	90.46	60.46	

108	trans-1,3-Dichloropropene					CAS #:	10061-02-6			
12.834	12.834	(0.880)	75	2149165	50.0000	50.248	80.00-	120.00	100.00	
12.834	12.834	(0.880)	77	647075			0.11-	60.11	30.11	
12.834	12.834	(0.880)	39	983279			15.75-	75.75	45.75	

110	1,1,2-Trichloroethane					CAS #:	79-00-5			
13.138	13.138	(0.901)	97	1477573	50.0000	47.104	80.00-	120.00	100.00	
13.138	13.138	(0.901)	99	908334			31.47-	91.47	61.47	
13.138	13.138	(0.901)	83	1303964			58.25-	118.25	88.25	

112	Tetrachloroethene					CAS #:	127-18-4			
13.193	13.193	(0.905)	166	2055564	50.0000	48.417	80.00-	120.00	100.00	
13.193	13.193	(0.905)	129	1560241			45.90-	105.90	75.90	
13.193	13.193	(0.905)	131	1518741			43.88-	103.88	73.88	

114	2-Hexanone					CAS #:	591-78-6			
13.580	13.580	(0.932)	58	1624483	50.0000	52.265	80.00-	120.00	100.00	
13.580	13.580	(0.932)	43	2875801			147.03-	207.03	177.03	
13.580	13.580	(0.932)	100	334216			0.00-	50.57	20.57	

116	Dibromochloromethane					CAS #:	124-48-1			
13.718	13.718	(0.941)	129	2439103	50.0000	49.615	80.00-	120.00	100.00	
13.718	13.718	(0.941)	127	1880746			47.11-	107.11	77.11	

117	1,2-Dibromoethane					CAS #:	106-93-4			
13.884	13.884	(0.953)	107	2349832	50.0000	49.163	80.00-	120.00	100.00	
13.884	13.884	(0.953)	109	2236461			65.18-	125.18	95.18	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
126 Chlorobenzene						CAS #:	108-90-7			
14.631	14.631	(1.004)	112	3693405	50.0000	48.755	80.00-	120.00	100.00	
14.631	14.631	(1.004)	114	1178415			1.91-	61.91	31.91	
14.603	14.603	(1.002)	77	2086900			26.50-	86.50	56.50	

129 Ethyl Benzene						CAS #:	100-41-4			
14.769	14.769	(1.013)	106	1897967	50.0000	47.343	80.00-	120.00	100.00	
14.769	14.769	(1.013)	91	5944387			283.20-	343.20	313.20	

130 m,p-Xylene						CAS #:	108-38-3			
14.935	14.935	(1.025)	106	2397145	50.0000	49.069	80.00-	120.00	100.00	
14.935	14.935	(1.025)	91	4593983			161.64-	221.64	191.64	

132 o-Xylene						CAS #:	95-47-6			
15.488	15.488	(1.063)	106	2399345	50.0000	48.671	80.00-	120.00	100.00	
15.488	15.488	(1.063)	91	4977982			177.47-	237.47	207.47	

134 Styrene						CAS #:	100-42-5			
15.516	15.516	(1.064)	104	3513401	50.0000	49.997	80.00-	120.00	100.00	
15.516	15.516	(1.064)	78	1765544			20.25-	80.25	50.25	

135 Bromoform						CAS #:	75-25-2			
15.764	15.764	(1.082)	173	2267799	50.0000	52.444	80.00-	120.00	100.00	
15.764	15.764	(1.082)	171	1157789			21.05-	81.05	51.05	

137 Cumene						CAS #:	98-82-8			
15.958	15.958	(1.095)	105	6732590	50.0000	49.967	80.00-	120.00	100.00	
15.958	15.958	(1.095)	120	1777182			0.00-	56.40	26.40	
15.958	15.958	(1.095)	51	616754			0.00-	39.16	9.16	

144 1,1,2,2-Tetrachloroethane						CAS #:	79-34-5			
16.456	16.456	(1.129)	83	3534141	50.0000	49.023	80.00-	120.00	100.00	
16.456	16.456	(1.129)	85	2190780			31.99-	91.99	61.99	

145 Propylbenzene						CAS #:	103-65-1			
16.483	16.483	(1.131)	91	8126384	50.0000	52.659	80.00-	120.00	100.00	
16.483	16.483	(1.131)	120	1859596			0.00-	52.88	22.88	
16.483	16.483	(1.131)	105	291527			0.00-	33.59	3.59	

147 4-Ethyltoluene						CAS #:	622-96-8			
16.649	16.649	(1.142)	105	7105769	50.0000	53.707	80.00-	120.00	100.00	
16.649	16.649	(1.142)	120	2103349			0.00-	59.60	29.60	

148 1,3,5-Trimethylbenzene						CAS #:	108-67-8			
16.732	16.732	(1.148)	105	6609789	50.0000	51.974	80.00-	120.00	100.00	
16.732	16.732	(1.148)	120	3263943			19.38-	79.38	49.38	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

153	1,2,4-Trimethylbenzene					CAS #: 95-63-6			
17.147	17.147	(1.176)	105	6904880	50.0000	53.940	80.00- 120.00	100.00	
17.147	17.147	(1.176)	120	3135375			15.41- 75.41	45.41	

156	1,3-Dichlorobenzene					CAS #: 541-73-1			
17.451	17.451	(1.197)	146	4379647	50.0000	52.483	80.00- 120.00	100.00	
17.451	17.451	(1.197)	148	2790179			33.71- 93.71	63.71	
17.451	17.451	(1.197)	111	1812557			11.39- 71.39	41.39	

157	1,4-Dichlorobenzene					CAS #: 106-46-7			
17.562	17.562	(1.205)	146	3953538	50.0000	49.990	80.00- 120.00	100.00	
17.562	17.562	(1.205)	148	2487194			32.91- 92.91	62.91	
17.562	17.562	(1.205)	111	1393312			5.24- 65.24	35.24	

158	alpha-Chlorotoluene					CAS #: 100-44-7			
17.700	17.700	(1.214)	91	6430403	50.0000	55.937	80.00- 120.00	100.00	
17.700	17.700	(1.214)	126	1298111			0.00- 50.19	20.19	

161	1,2-Dichlorobenzene					CAS #: 95-50-1			
17.921	17.921	(1.230)	146	3963669	50.0000	49.308	80.00- 120.00	100.00	
17.921	17.921	(1.230)	148	2485367			32.70- 92.70	62.70	
17.921	17.921	(1.230)	111	1469135			7.07- 67.07	37.07	

167	1,2,4-Trichlorobenzene					CAS #: 120-82-1			
19.276	19.276	(1.322)	180	4243469	50.0000	52.231	80.00- 120.00	100.00	
19.276	19.276	(1.322)	182	4039439			65.19- 125.19	95.19	

168	Hexachlorobutadiene					CAS #: 87-68-3			
19.359	19.359	(1.328)	225	2274747	50.0000	48.779	80.00- 120.00	100.00	
19.359	19.359	(1.328)	223	1438980			33.26- 93.26	63.26	

169	Naphthalene					CAS #: 91-20-3			
19.470	19.470	(1.336)	128	10653699	50.0000	62.178	80.00- 120.00	100.00	
19.470	19.470	(1.336)	127	1203008			0.00- 41.29	11.29	

Report Date: 31-May-2007 14:53

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053007.d

Calibration Time: 16:03

Lab Smp Id: ICAL

Client Smp ID: Level 5

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-50ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	441133	264680	617586	441133	0.00
88 1,4-Difluorobenze	1992312	1195387	2789237	1992312	0.00
125 Chlorobenzene-d5	1475337	885202	2065472	1475337	0.00

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

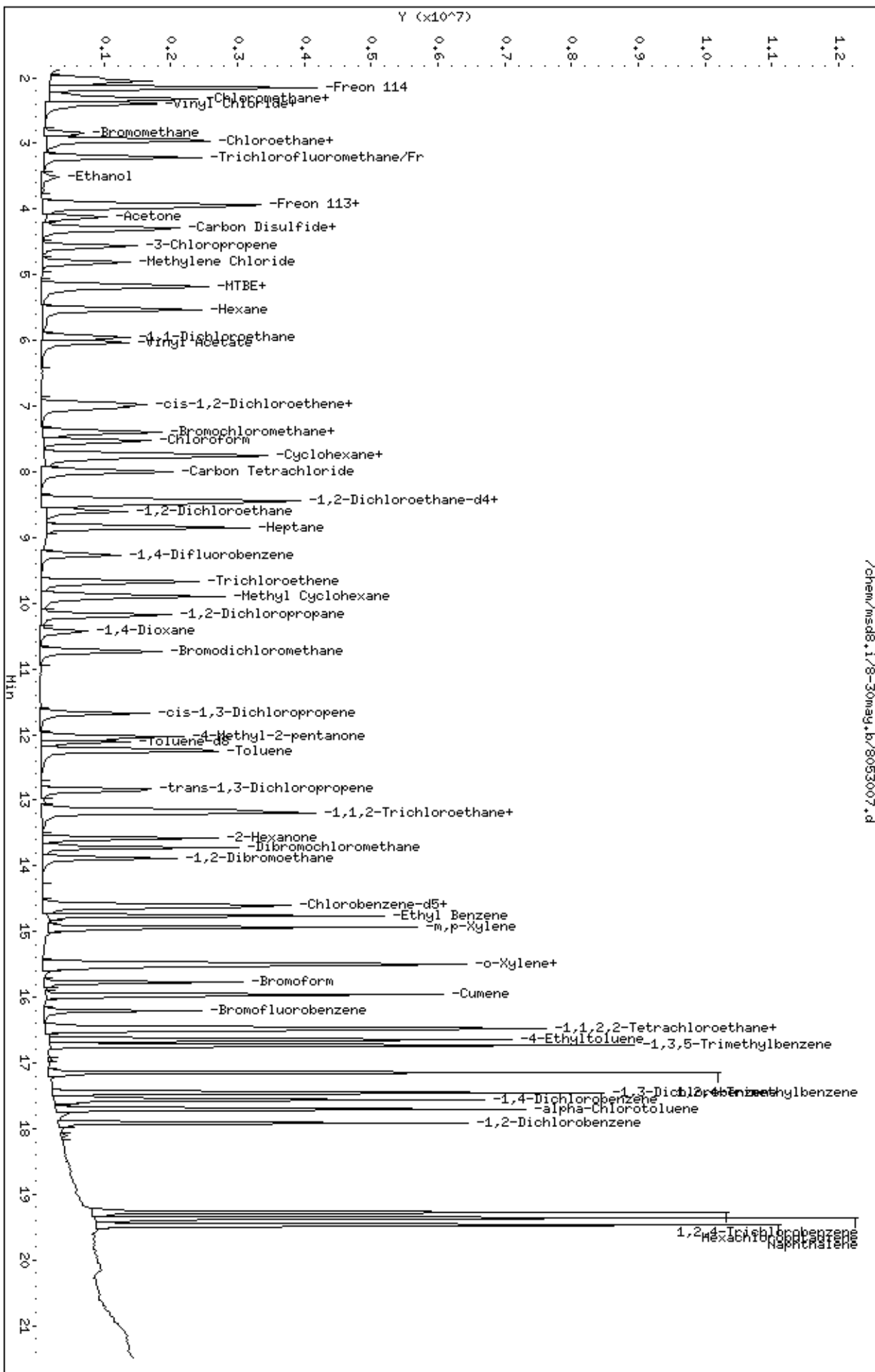
RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-30may.b/8053007.d
Date: 30-May-2007 16:03
Client ID: Level 5
Sample Info: 50ml #1487-289

Column phase: RTX-624

Instrument: msd8.1
Operator: db
Column diameter: 0.53

/chem/msd8.1/8-30may.b/8053007.d



Report Date: 31-May-2007 14:53

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053008.d
 Lab Smp Id: ICAL Client Smp ID: Level 6
 Inj Date : 30-MAY-2007 16:31
 Operator : db Inst ID: msd8.i
 Smp Info : 100ml #1487-289
 Misc Info : 200ppbv-100ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 31-May-2007 14:53 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 16:31 Cal File: 8053008.d
 Als bottle: 1 Calibration Sample, Level: 6
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04mdl+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	ON-COL	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	456242	25.0000		70.00- 130.00	100.00	
7.387	7.387	(1.000)	128	338679			47.57- 107.57	74.23	
7.387	7.387	(1.000)	49	641995			113.47- 173.47	140.71	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	2079239	25.0000		70.00- 130.00	100.00	
9.267	9.267	(1.000)	88	315764			0.00- 45.68	15.19	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	1515229	25.0000		70.00- 130.00	100.00	
14.576	14.576	(1.000)	82	883171			0.00- 30.00	58.29	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465	(1.146)	65	611646	25.0000	25.615	70.00- 130.00	100.00	
8.465	8.465	(1.146)	67	381750			0.00- 30.00	62.41	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115	(1.307)	98	1771106	25.0000	24.605	70.00- 130.00	100.00	
12.115	12.115	(1.307)	70	167250			0.00- 30.00	9.44	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
\$ 104 Toluene-d8 (continued)										
12.115	12.115	(1.307)	100	1418894			0.00- 30.00	80.11		

\$ 140 Bromofluorobenzene										
						CAS #: 460-00-4				
16.207	16.207	(1.112)	174	936709	25.0000	25.982	70.00- 130.00	100.00		
16.207	16.207	(1.112)	95	1233267			102.16- 162.16	131.66		
16.207	16.207	(1.112)	176	887102			64.31- 124.31	94.70		

3 Propylene						CAS #: 115-07-1				
1.995	1.995	(0.270)	41	2111882	100.000	85.522	70.00- 130.00	100.00		
1.995	1.995	(0.270)	42	1405317			0.00- 30.00	66.54		
1.995	1.995	(0.270)	39	1426212			0.00- 30.00	67.53		

4 Dichlorodifluoromethane/Fr12						CAS #: 75-71-8				
2.050	2.050	(0.278)	85	5440637	100.000	87.042	70.00- 130.00	100.00		
2.050	2.050	(0.278)	87	1735219			0.00- 30.00	31.89		

6 Freon 114						CAS #: 76-14-2				
2.161	2.161	(0.293)	135	5255420	100.000	93.067	70.00- 130.00	100.00		
2.161	2.161	(0.293)	137	1643521			1.53- 61.53	31.27		

8 Chloromethane						CAS #: 74-87-3				
2.272	2.272	(0.308)	50	2731679	100.000	94.896	70.00- 130.00	100.00		
2.272	2.272	(0.308)	52	810595			0.00- 30.00	29.67		

9 Butane						CAS #: 106-97-8				
2.327	2.327	(0.315)	58	666073	100.000	88.601	70.00- 130.00	100.00		
2.327	2.327	(0.315)	43	5010913			0.00- 30.00	752.31		

11 Vinyl Chloride						CAS #: 75-01-4				
2.410	2.410	(0.326)	62	2979171	100.000	89.067	70.00- 130.00	100.00		
2.410	2.410	(0.326)	64	906402			0.00- 30.00	30.42		

10 1,3-Butadiene						CAS #: 106-99-0				
2.410	2.410	(0.326)	54	2379256	100.000	83.158	70.00- 130.00	100.00		
2.382	2.382	(0.322)	39	2438472			0.00- 30.00	102.49		

13 Bromomethane						CAS #: 74-83-9				
2.852	2.852	(0.386)	94	2174749	100.000	97.320	70.00- 130.00	100.00		
2.852	2.852	(0.386)	96	2040837			65.03- 125.03	93.84		

16 Chloroethane						CAS #: 75-00-3				
2.963	2.963	(0.401)	64	1587282	100.000	89.665	70.00- 130.00	100.00		
2.963	2.963	(0.401)	49	396910			0.00- 30.00	25.01		
2.963	2.963	(0.401)	66	488935			0.00- 30.00	30.80		

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

15 Isopentane						CAS #: 78-78-4			
2.963	2.963	(0.401)	43	3971363	100.000	95.086	70.00- 130.00	100.00	
2.963	2.963	(0.401)	57	2686310			0.00- 30.00	67.64	
2.963	2.963	(0.401)	72	284025			0.00- 30.00	7.15	

18 Trichlorofluoromethane/Fr11						CAS #: 75-69-4			
3.212	3.212	(0.435)	101	6617115	100.000	93.139	70.00- 130.00	100.00	
3.212	3.212	(0.435)	103	4259322			34.97- 94.97	64.37	

23 Ethanol						CAS #: 64-17-5			
3.543	3.543	(0.480)	45	1062280	100.000	93.635	70.00- 130.00	100.00	
3.516	3.516	(0.476)	43	191015			0.00- 30.00	17.98	
3.516	3.516	(0.476)	46	431356			0.00- 30.00	40.61	

28 Freon 113						CAS #: 76-13-1			
3.930	3.930	(0.532)	151	4294641	100.000	91.363	70.00- 130.00	100.00	
3.930	3.930	(0.532)	153	2715258			33.71- 93.71	63.22	
3.930	3.930	(0.532)	101	5038552			86.34- 146.34	117.32	

29 1,1-Dichloroethene						CAS #: 75-35-4			
3.958	3.958	(0.536)	61	4286913	100.000	91.832	70.00- 130.00	100.00	
3.958	3.958	(0.536)	96	2549826			30.45- 90.45	59.48	
3.958	3.958	(0.536)	98	1633650			8.39- 68.39	38.11	

30 Acetone						CAS #: 67-64-1			
4.124	4.124	(0.558)	58	1456647	100.000	93.167	70.00- 130.00	100.00	
4.124	4.124	(0.558)	43	4343030			0.00- 30.00	298.15	

33 Carbon Disulfide						CAS #: 75-15-0			
4.290	4.290	(0.581)	76	8061972	100.000	92.149	70.00- 130.00	100.00	

34 2-Propanol						CAS #: 67-63-0			
4.318	4.318	(0.584)	45	5257774	100.000	93.009	70.00- 130.00	100.00	
4.318	4.318	(0.584)	43	986622			0.00- 30.00	18.77	
4.318	4.318	(0.584)	59	208508			0.00- 30.00	3.97	

37 3-Chloropropene						CAS #: 107-05-1			
4.566	4.566	(0.618)	76	1354721	100.000	97.062	70.00- 130.00	100.00	
4.566	4.566	(0.618)	41	4132615			0.00- 30.00	305.05	

40 Methylene Chloride						CAS #: 75-09-2			
4.815	4.815	(0.652)	49	3152614	100.000	90.205	70.00- 130.00	100.00	
4.815	4.815	(0.652)	84	2300704			41.01- 101.01	72.98	
4.815	4.815	(0.652)	51	949529			0.00- 30.00	30.12	

43 MTBE						CAS #: 1634-04-4			
5.147	5.147	(0.697)	73	5123497	100.000	103.08	70.00- 130.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
43 MTBE (continued)									
5.147	5.147	(0.697)	57	1222306			0.00- 53.82	23.86	
5.147	5.147	(0.697)	41	1170037			0.00- 30.00	22.84	

45 trans-1,2-Dichloroethene					CAS #: 156-60-5				
5.175	5.175	(0.701)	96	2948613	100.000	89.263	70.00- 130.00	100.00	
5.175	5.175	(0.701)	61	4422228			119.04- 179.04	149.98	
5.175	5.175	(0.701)	98	1894246			0.00- 30.00	64.24	

46 Hexane					CAS #: 110-54-3				
5.534	5.534	(0.749)	57	4711392	100.000	94.039	70.00- 130.00	100.00	
5.534	5.534	(0.749)	43	2957338			0.00- 30.00	62.77	
5.534	5.534	(0.749)	86	725012			0.00- 30.00	15.39	

54 1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	5084008	100.000	94.412	70.00- 130.00	100.00	
5.949	5.949	(0.805)	65	1596380			1.22- 61.22	31.40	

55 Vinyl Acetate					CAS #: 108-05-4				
6.032	6.032	(0.817)	86	761982	100.000	100.24	70.00- 130.00	100.00	
6.032	6.032	(0.817)	43	8054687			0.00- 30.00	1057.07	
6.032	6.032	(0.817)	42	616418			0.00- 30.00	80.90	

64 cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.972	6.972	(0.944)	61	3773397	100.000	88.539	70.00- 130.00	100.00	
6.972	6.972	(0.944)	96	2782950			45.09- 105.09	73.75	
6.972	6.972	(0.944)	98	1774497			17.66- 77.66	47.03	

65 2-Butanone					CAS #: 78-93-3				
7.027	7.027	(0.951)	72	1390585	100.000	91.971	70.00- 130.00	100.00	
7.027	7.027	(0.951)	43	5954215			398.56- 458.56	428.18	
7.027	7.027	(0.951)	57	451831			0.00- 30.00	32.49	

67 Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.387	(1.000)	42	3522933	100.000	89.642	70.00- 130.00	100.00	
7.387	7.387	(1.000)	71	1272954			5.14- 65.14	36.13	
7.387	7.387	(1.000)	72	1374375			0.00- 30.00	39.01	

70 Chloroform					CAS #: 67-66-3				
7.525	7.525	(1.019)	83	4994804	100.000	92.746	70.00- 130.00	100.00	
7.525	7.525	(1.019)	85	3098846			31.98- 91.98	62.04	

73 Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	4057695	100.000	90.810	70.00- 130.00	100.00	
7.746	7.746	(1.049)	56	5201149			99.30- 159.30	128.18	
7.746	7.746	(1.049)	41	2605789			33.84- 93.84	64.22	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

75	1,1,1-Trichloroethane				CAS #:		71-55-6		
7.774	7.774	(1.052)	97	5240694	100.000	92.876	70.00-	130.00	100.00
7.774	7.774	(1.052)	99	3350182			34.73-	94.73	63.93

77	Carbon Tetrachloride				CAS #:		56-23-5		
7.995	7.995	(1.082)	119	5107823	100.000	99.478	70.00-	130.00	100.00
7.995	7.995	(1.082)	117	5255553			74.04-	134.04	102.89

81	Benzene				CAS #:		71-43-2		
8.437	8.437	(0.910)	78	8696555	100.000	92.587	70.00-	130.00	100.00
8.437	8.437	(0.910)	77	1944596			0.00-	30.00	22.36

80	2,2,4-Trimethylpentane				CAS #:		540-84-1		
8.465	8.465	(1.146)	57	12520618	100.000	93.863	70.00-	130.00	100.00
8.465	8.465	(1.146)	56	3975572			0.00-	30.00	31.75
8.465	8.465	(1.146)	41	2931448			0.00-	30.00	23.41

83	1,2-Dichloroethane				CAS #:		107-06-2		
8.603	8.603	(0.928)	62	3438647	100.000	90.230	70.00-	130.00	100.00
8.603	8.603	(0.928)	64	1083383			0.00-	30.00	31.51

85	Heptane				CAS #:		142-82-5		
8.852	8.852	(0.955)	100	862354	100.000	84.838	70.00-	130.00	100.00
8.852	8.852	(0.955)	43	4963883			0.00-	30.00	575.62
8.852	8.852	(0.955)	71	2771790			0.00-	30.00	321.42

94	Trichloroethene				CAS #:		79-01-6		
9.682	9.682	(1.045)	95	3411326	100.000	91.643	70.00-	130.00	100.00
9.682	9.682	(1.045)	130	3669068			77.84-	137.84	107.56
9.682	9.682	(1.045)	97	2155039			34.22-	94.22	63.17

95	Methyl Cyclohexane				CAS #:		108-87-2		
9.903	9.903	(1.341)	83	5261578	100.000	94.518	70.00-	130.00	100.00
9.903	9.903	(1.341)	98	2392828			0.00-	30.00	45.48
9.903	9.903	(1.341)	55	4199922			0.00-	30.00	79.82

97	1,2-Dichloropropane				CAS #:		78-87-5		
10.179	10.179	(1.098)	63	3003131	100.000	92.255	70.00-	130.00	100.00
10.179	10.179	(1.098)	62	2137034			42.15-	102.15	71.16
10.179	10.179	(1.098)	41	1697497			26.47-	86.47	56.52

98	1,4-Dioxane				CAS #:		123-91-1		
10.428	10.428	(1.125)	88	1913662	100.000	96.383	70.00-	130.00	100.00
10.428	10.428	(1.125)	58	1398870			44.03-	104.03	73.10
10.428	10.428	(1.125)	57	419838			0.00-	30.00	21.94

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

100	Bromodichloromethane					CAS #:	75-27-4			
10.732	10.732	(1.158)	83	5128594	100.000	94.331	70.00-	130.00	100.00	
10.732	10.732	(1.158)	85	3168965			31.03-	91.03	61.79	

102	cis-1,3-Dichloropropene					CAS #:	10061-01-5			
11.672	11.672	(1.260)	75	4289445	100.000	94.820	70.00-	130.00	100.00	
11.672	11.672	(1.260)	77	1336780			1.19-	61.19	31.16	
11.672	11.672	(1.260)	39	1995148			16.93-	76.93	46.51	

103	4-Methyl-2-pentanone					CAS #:	108-10-1			
12.032	12.032	(1.298)	58	2412926	100.000	98.595	70.00-	130.00	100.00	
12.032	12.032	(1.298)	43	5915330			0.00-	30.00	245.15	
12.032	12.032	(1.298)	85	996247			0.00-	30.00	41.29	

105	Toluene					CAS #:	108-88-3			
12.253	12.253	(1.322)	91	9100045	100.000	95.324	70.00-	130.00	100.00	
12.253	12.253	(1.322)	92	5435801			30.46-	90.46	59.73	

108	trans-1,3-Dichloropropene					CAS #:	10061-02-6			
12.834	12.834	(0.880)	75	4335904	100.000	98.705	70.00-	130.00	100.00	
12.834	12.834	(0.880)	77	1335466			0.11-	60.11	30.80	
12.834	12.834	(0.880)	39	2001936			15.75-	75.75	46.17	

110	1,1,2-Trichloroethane					CAS #:	79-00-5			
13.138	13.138	(0.901)	97	2964497	100.000	92.018	70.00-	130.00	100.00	
13.138	13.138	(0.901)	99	1840259			31.47-	91.47	62.08	
13.138	13.138	(0.901)	83	2551394			58.25-	118.25	86.06	

112	Tetrachloroethene					CAS #:	127-18-4			
13.193	13.193	(0.905)	166	4186065	100.000	96.002	70.00-	130.00	100.00	
13.193	13.193	(0.905)	129	3258954			45.90-	105.90	77.85	
13.193	13.193	(0.905)	131	3031301			43.88-	103.88	72.41	

114	2-Hexanone					CAS #:	591-78-6			
13.580	13.580	(0.932)	58	3291331	100.000	103.10	70.00-	130.00	100.00	
13.580	13.580	(0.932)	43	6040547			147.03-	207.03	183.53	
13.580	13.580	(0.932)	100	705295			0.00-	30.00	21.43	

116	Dibromochloromethane					CAS #:	124-48-1			
13.718	13.718	(0.941)	129	5095518	100.000	100.92	70.00-	130.00	100.00	
13.718	13.718	(0.941)	127	3903792			0.00-	30.00	76.61	

117	1,2-Dibromoethane					CAS #:	106-93-4			
13.884	13.884	(0.953)	107	4784702	100.000	97.469	70.00-	130.00	100.00	
13.884	13.884	(0.953)	109	4502499			65.18-	125.18	94.10	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
126 Chlorobenzene						CAS #:	108-90-7		
14.631	14.631	(1.004)	112	7575451	100.000	97.367	70.00- 130.00	100.00	
14.631	14.631	(1.004)	114	2379711			1.91- 61.91	31.41	
14.603	14.603	(1.002)	77	4237044			26.50- 86.50	55.93	

129 Ethyl Benzene						CAS #:	100-41-4		
14.769	14.769	(1.013)	106	3943287	100.000	95.771	70.00- 130.00	100.00	
14.769	14.769	(1.013)	91	12558154			0.00- 30.00	318.47	

130 m,p-Xylene						CAS #:	108-38-3		
14.935	14.935	(1.025)	106	4928477	100.000	98.229	70.00- 130.00	100.00	
14.935	14.935	(1.025)	91	9732569			0.00- 30.00	197.48	

132 o-Xylene						CAS #:	95-47-6		
15.488	15.488	(1.063)	106	4858089	100.000	95.953	70.00- 130.00	100.00	
15.488	15.488	(1.063)	91	10291979			177.47- 237.47	211.85	

134 Styrene						CAS #:	100-42-5		
15.516	15.516	(1.064)	104	7423994	100.000	102.86	70.00- 130.00	100.00	
15.516	15.516	(1.064)	78	3649528			20.25- 80.25	49.16	

135 Bromoform						CAS #:	75-25-2		
15.764	15.764	(1.082)	173	4754269	100.000	107.05	70.00- 130.00	100.00	
15.764	15.764	(1.082)	171	2426878			21.05- 81.05	51.05	

137 Cumene						CAS #:	98-82-8		
15.958	15.958	(1.095)	105	14013350	100.000	101.26	70.00- 130.00	100.00	
15.958	15.958	(1.095)	120	3600114			0.00- 30.00	25.69	
15.958	15.958	(1.095)	51	1268255			0.00- 30.00	9.05	

144 1,1,2,2-Tetrachloroethane						CAS #:	79-34-5		
16.456	16.456	(1.129)	83	7218673	100.000	97.495	70.00- 130.00	100.00	
16.456	16.456	(1.129)	85	4420519			31.99- 91.99	61.24	

145 Propylbenzene						CAS #:	103-65-1		
16.483	16.483	(1.131)	91	16975123	100.000	107.10	70.00- 130.00	100.00	
16.483	16.483	(1.131)	120	3752339			0.00- 30.00	22.10	
16.483	16.483	(1.131)	105	582164			0.00- 30.00	3.43	

147 4-Ethyltoluene						CAS #:	622-96-8		
16.649	16.649	(1.142)	105	14848016	100.000	109.27	70.00- 130.00	100.00	
16.649	16.649	(1.142)	120	4383266			0.00- 59.60	29.52	

148 1,3,5-Trimethylbenzene						CAS #:	108-67-8		
16.732	16.732	(1.148)	105	13957272	100.000	106.86	70.00- 130.00	100.00	
16.732	16.732	(1.148)	120	6759537			0.00- 30.00	48.43	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

153	1,2,4-Trimethylbenzene					CAS #: 95-63-6			
17.147	17.147	(1.176)	105	14463263	100.000	110.01	70.00- 130.00	100.00	
17.147	17.147	(1.176)	120	6341397			15.41- 75.41	43.84	

156	1,3-Dichlorobenzene					CAS #: 541-73-1			
17.451	17.451	(1.197)	146	9209832	100.000	107.46	70.00- 130.00	100.00	
17.451	17.451	(1.197)	148	5794867			0.00- 30.00	62.92	
17.451	17.451	(1.197)	111	3806522			0.00- 30.00	41.33	

157	1,4-Dichlorobenzene					CAS #: 106-46-7			
17.562	17.562	(1.205)	146	8545658	100.000	105.21	70.00- 130.00	100.00	
17.562	17.562	(1.205)	148	5364507			0.00- 30.00	62.77	
17.562	17.562	(1.205)	111	2979869			0.00- 30.00	34.87	

158	alpha-Chlorotoluene					CAS #: 100-44-7			
17.700	17.700	(1.214)	91	14388865	100.000	121.87	70.00- 130.00	100.00	
17.700	17.700	(1.214)	126	2812628			0.00- 30.00	19.55	

161	1,2-Dichlorobenzene					CAS #: 95-50-1			
17.921	17.921	(1.230)	146	8345519	100.000	101.08	70.00- 130.00	100.00	
17.921	17.921	(1.230)	148	5277126			32.70- 92.70	63.23	
17.921	17.921	(1.230)	111	3065390			7.07- 67.07	36.73	

167	1,2,4-Trichlorobenzene					CAS #: 120-82-1			
19.276	19.276	(1.322)	180	8958905	100.000	107.37	70.00- 130.00	100.00	
19.276	19.276	(1.322)	182	8611749			65.19- 125.19	96.13	

168	Hexachlorobutadiene					CAS #: 87-68-3			
19.359	19.359	(1.328)	225	4504987	100.000	94.060	70.00- 130.00	100.00	
19.359	19.359	(1.328)	223	2843237			33.26- 93.26	63.11	

169	Naphthalene					CAS #: 91-20-3			
19.469	19.469	(1.336)	128	15136012	100.000	86.013	70.00- 130.00	100.00	
19.469	19.469	(1.336)	127	2518408			0.00- 30.00	16.64	

Report Date: 31-May-2007 14:53

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053008.d

Calibration Time: 16:03

Lab Smp Id: ICAL

Client Smp ID: Level 6

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-100ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	441133	264680	617586	456242	3.43
88 1,4-Difluorobenze	1992312	1195387	2789237	2079239	4.36
125 Chlorobenzene-d5	1475337	885202	2065472	1515229	2.70

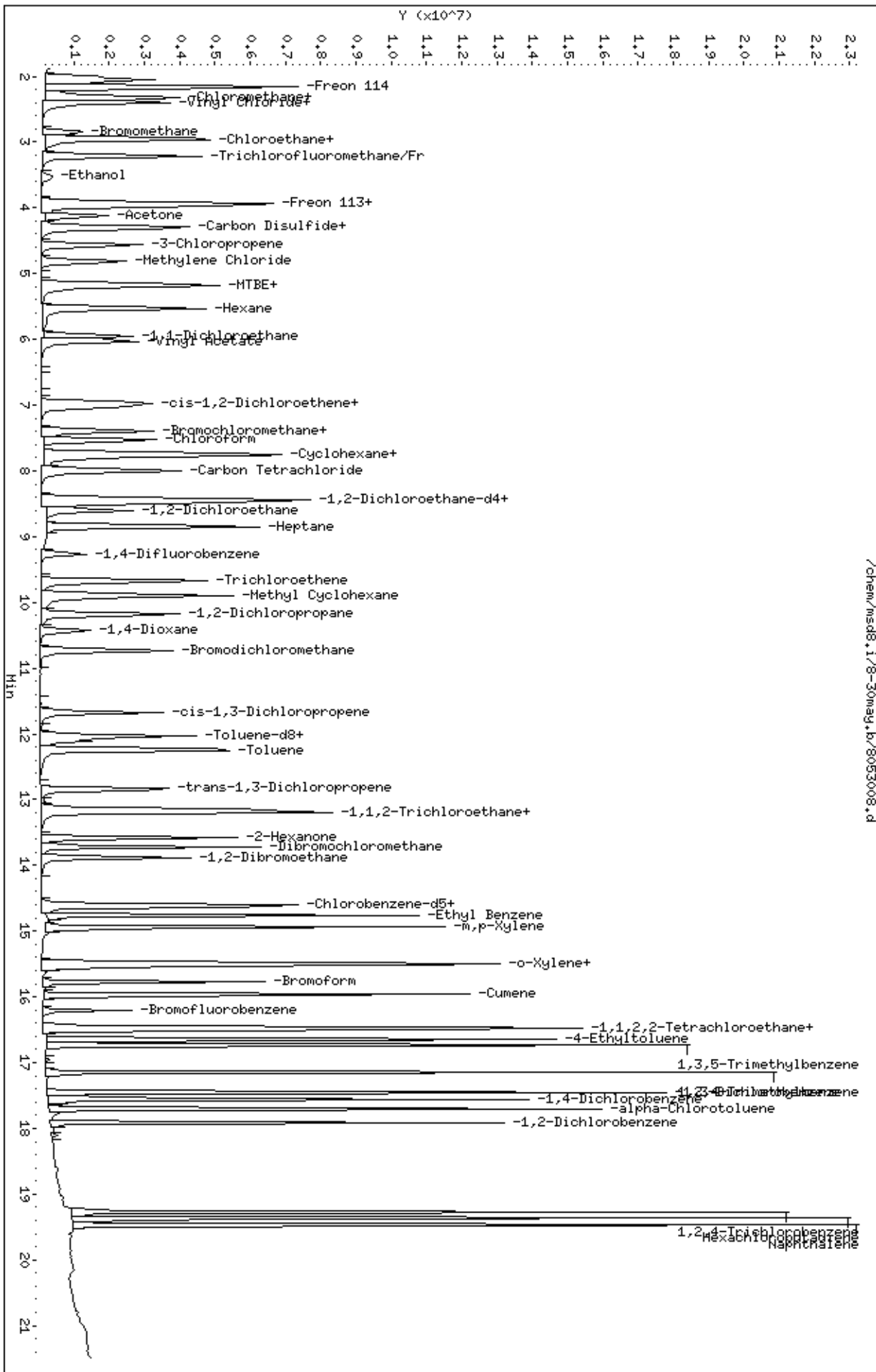
COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.



Report Date: 07-Jun-2007 13:41

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-07jun.b/8060706.d
 Lab Smp Id: ICAL Client Smp ID: Level 7
 Inj Date : 07-JUN-2007 12:08
 Operator : JG Inst ID: msd8.i
 Smp Info : 200ml #1443-96
 Misc Info : 200ppbv-200ppbv
 Comment :
 Method : /chem/msd8.i/8-07jun.b/t14q530b.m
 Meth Date : 07-Jun-2007 13:41 jgray Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1 Calibration Sample, Level: 7
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: sp16b.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS								
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	ON-COL	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====

* 68	Bromochloromethane					CAS #:	74-97-5	
7.387	7.387	(1.000)	130	361211	25.0000		70.00- 130.00	100.00
7.387	7.387	(1.000)	128	275936			42.98- 102.98	76.39
7.387	7.387	(1.000)	49	511706			112.53- 172.53	141.66

* 88	1,4-Difluorobenzene					CAS #:	540-36-3	
9.267	9.267	(1.000)	114	1595379	25.0000		70.00- 130.00	100.00
9.267	9.267	(1.000)	88	244093			0.00- 44.58	15.30

* 125	Chlorobenzene-d5					CAS #:	3114-55-4	
14.576	14.576	(1.000)	117	1171779	25.0000		70.00- 130.00	100.00
14.576	14.576	(1.000)	82	674436			0.00- 30.00	57.56

1	Freon 152a					CAS #:	75-37-6	
2.023	2.023	(0.274)	65	2371527	200.000	189.53	70.00- 130.00	100.00
2.078	2.078	(0.281)	51	10140161			0.00- 30.00	427.58

20	Freon123a					CAS #:	354-23-4	
3.682	3.682	(0.498)	67	3750189	200.000	194.11	70.00- 130.00	100.00
3.710	3.710	(0.502)	117	2671033			0.00- 30.00	71.22

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
21 Freon123						CAS #: 306-83-2			
3.792	3.792	(0.513)	83	2375930	200.000	195.12	70.00- 130.00	100.00	
3.792	3.792	(0.513)	133	448165			0.00- 30.00	18.86	
3.792	3.792	(0.513)	85	1758646			0.00- 30.00	74.02	

38 tert-Butyl-Alcohol						CAS #: 75-65-0			
4.954	4.954	(0.671)	59	3329003	200.000	132.23	70.00- 130.00	100.00	
4.954	4.954	(0.671)	41	767923			0.00- 30.00	23.07	
4.954	4.954	(0.671)	57	334446			0.00- 30.00	10.05	

49 Isopropyl ether						CAS #: 108-20-3			
5.949	5.949	(0.805)	45	12287330	200.000	195.16	70.00- 130.00	100.00	
5.949	5.949	(0.805)	87	3425229			0.00- 30.00	27.88	
5.949	5.949	(0.805)	59	1325426			0.00- 30.00	10.79	

52 1-Propanol						CAS #: 71-23-8			
6.170	6.170	(0.835)	42	896926	200.000	197.65	70.00- 130.00	100.00	
6.170	6.170	(0.835)	59	1152290			0.00- 30.00	128.47	
6.143	6.143	(0.832)	41	735160			0.00- 30.00	81.96	

58 Ethyl-tert-butyl Ether						CAS #: 637-92-3			
6.585	6.585	(0.891)	59	8042475	200.000	191.32	70.00- 130.00	100.00	
6.585	6.585	(0.891)	87	3100400			0.00- 30.00	38.55	
6.585	6.585	(0.891)	41	1355089			0.00- 30.00	16.85	

61 Ethyl Acetate						CAS #: 141-78-6			
7.083	7.083	(0.959)	70	1061516	200.000	193.50	70.00- 130.00	100.00	
7.083	7.083	(0.959)	43	9985386			0.00- 30.00	940.67	
7.083	7.083	(0.959)	61	1428422			0.00- 30.00	134.56	

78 Isobutanol						CAS #: 78-83-1			
8.438	8.438	(0.910)	43	3751872	200.000	204.90	70.00- 130.00	100.00(A)	
8.438	8.438	(0.910)	41	2497173			0.00- 30.00	66.56	

79 tert-amyl-Methyl Ether						CAS #: 994-05-8			
8.631	8.631	(1.168)	73	6928131	200.000	180.22	70.00- 130.00	100.00	
8.631	8.631	(1.168)	87	1658341			0.00- 30.00	23.94	
8.631	8.631	(1.168)	55	1860950			0.00- 30.00	26.86	

89 1-Butanol						CAS #: 71-36-3			
9.709	9.709	(1.048)	56	3377553	200.000	216.27	70.00- 130.00	100.00(A)	
9.709	9.709	(1.048)	41	2212090			0.00- 30.00	65.49	
9.709	9.709	(1.048)	43	1815594			0.00- 30.00	53.75	

136 Cyclohexanone						CAS #: 108-94-1			
16.152	16.152	(1.108)	55	5524328	200.000	205.99	70.00- 130.00	100.00(A)	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
136 Cyclohexanone (continued)									
16.152	16.152	(1.108)	98	2497048			0.00- 30.00	45.20	
16.124	16.124	(1.106)	42	3645343			0.00- 30.00	65.99	

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Report Date: 07-Jun-2007 13:41

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 07-JUN-2007

Lab File ID: 8060706.d

Calibration Time: 11:37

Lab Smp Id: ICAL

Client Smp ID: Level 7

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: JG

Method File: /chem/msd8.i/8-07jun.b/t14q530b.m

Misc Info: 200ppbv-200ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	350593	210356	490830	361211	3.03
88 1,4-Difluorobenze	1524282	914569	2133995	1595379	4.66
125 Chlorobenzene-d5	1168126	700876	1635376	1171779	0.31

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

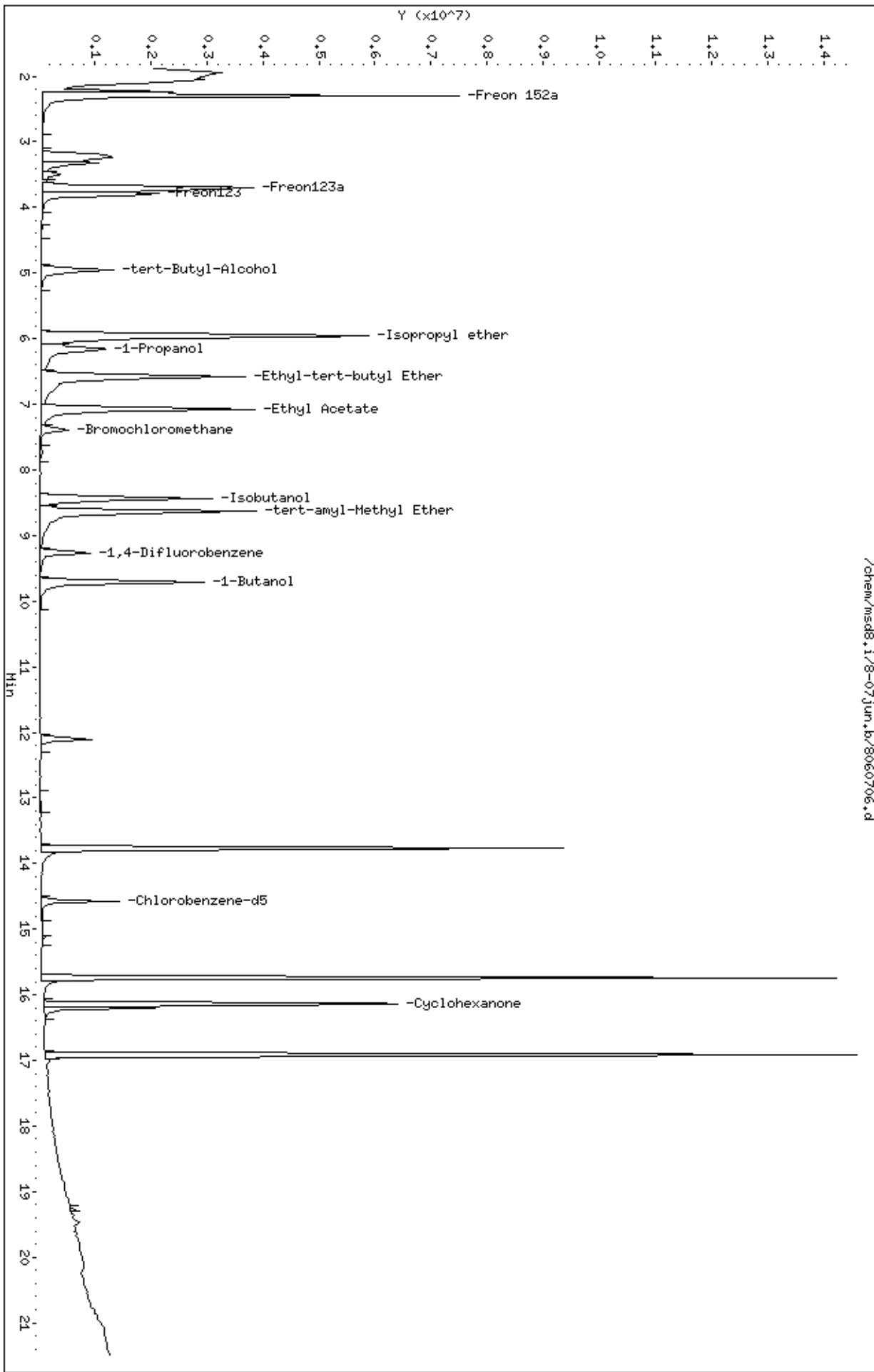
RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-07jun.b/8060706.d
Date : 07-JUN-2007 12:08
Client ID: Level 7
Sample Info: 200ml #1443-96

Column phase: RTX-624

Instrument: msd8.1
Operator: JG
Column diameter: 0.53

/chem/msd8.1/8-07jun.b/8060706.d



Report Date: 31-May-2007 14:53

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-30may.b/8053009.d
 Lab Smp Id: ICAL Client Smp ID: Level 7
 Inj Date : 30-MAY-2007 17:02
 Operator : db Inst ID: msd8.i
 Smp Info : 200ml #1487-289
 Misc Info : 200ppbv-200ppbv
 Comment :
 Method : /chem/msd8.i/8-30may.b/t14q530a.m
 Meth Date : 31-May-2007 14:53 jgray Quant Type: ISTD
 Cal Date : 30-MAY-2007 17:02 Cal File: 8053009.d
 Als bottle: 1 Calibration Sample, Level: 7
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04mdl+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	ON-COL	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	448309	25.0000		70.00- 130.00	100.00	
7.387	7.387	(1.000)	128	332770			47.57- 107.57	74.23	
7.387	7.387	(1.000)	49	652713			113.47- 173.47	145.59	

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	2033490	25.0000		70.00- 130.00	100.00	
9.267	9.267	(1.000)	88	307791			0.00- 45.68	15.14	

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	1524596	25.0000		70.00- 130.00	100.00	
14.576	14.576	(1.000)	82	864175			0.00- 30.00	56.68	

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465	(1.146)	65	624669	25.0000	26.623	70.00- 130.00	100.00	
8.465	8.465	(1.146)	67	466938			0.00- 30.00	74.75	

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115	(1.307)	98	1740690	25.0000	24.726	70.00- 130.00	100.00	
12.115	12.115	(1.307)	70	177697			0.00- 30.00	10.21	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		
\$ 104 Toluene-d8 (continued)										
12.115	12.115	(1.307)	100	1188238			0.00- 30.00	68.26		

\$ 140 Bromofluorobenzene										
						CAS #: 460-00-4				
16.207	16.207	(1.112)	174	926433	25.0000	25.539	70.00- 130.00	100.00		
16.207	16.207	(1.112)	95	1239953			102.16- 162.16	133.84		
16.207	16.207	(1.112)	176	905386			64.31- 124.31	97.73		

3 Propylene						CAS #: 115-07-1				
1.995	1.995	(0.270)	41	4152556	200.000	171.14	70.00- 130.00	100.00		
1.995	1.995	(0.270)	42	2781935			0.00- 30.00	66.99		
1.995	1.995	(0.270)	39	2816950			0.00- 30.00	67.84		

4 Dichlorodifluoromethane/Fr12						CAS #: 75-71-8				
2.050	2.050	(0.278)	85	10585010	200.000	172.34	70.00- 130.00	100.00		
2.050	2.050	(0.278)	87	3334935			0.00- 30.00	31.51		

6 Freon 114						CAS #: 76-14-2				
2.189	2.189	(0.296)	135	10270951	200.000	185.10	70.00- 130.00	100.00		
2.189	2.189	(0.296)	137	3216633			1.53- 61.53	31.32		

8 Chloromethane						CAS #: 74-87-3				
2.299	2.299	(0.311)	50	5091461	200.000	180.00	70.00- 130.00	100.00		
2.299	2.299	(0.311)	52	1478081			0.00- 30.00	29.03		

9 Butane						CAS #: 106-97-8				
2.355	2.355	(0.319)	58	1345216	200.000	182.11	70.00- 130.00	100.00		
2.355	2.355	(0.319)	43	9990258			0.00- 30.00	742.65		

11 Vinyl Chloride						CAS #: 75-01-4				
2.410	2.410	(0.326)	62	5882802	200.000	178.99	70.00- 130.00	100.00		
2.410	2.410	(0.326)	64	1795563			0.00- 30.00	30.52		

10 1,3-Butadiene						CAS #: 106-99-0				
2.410	2.410	(0.326)	54	4947217	200.000	175.97	70.00- 130.00	100.00		
2.410	2.410	(0.326)	39	5464185			0.00- 30.00	110.45		

13 Bromomethane						CAS #: 74-83-9				
2.852	2.852	(0.386)	94	4328268	200.000	197.12	70.00- 130.00	100.00		
2.852	2.852	(0.386)	96	4126213			65.03- 125.03	95.33		

16 Chloroethane						CAS #: 75-00-3				
2.963	2.963	(0.401)	64	3148287	200.000	180.99	70.00- 130.00	100.00		
2.963	2.963	(0.401)	49	753176			0.00- 30.00	23.92		
2.963	2.963	(0.401)	66	936359			0.00- 30.00	29.74		

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

15 Isopentane						CAS #: 78-78-4			
2.963	2.963	(0.401)	43	8041442	200.000	195.94	70.00- 130.00	100.00	
2.963	2.963	(0.401)	57	5344427			0.00- 30.00	66.46	
2.963	2.963	(0.401)	72	583435			0.00- 30.00	7.26	

18 Trichlorofluoromethane/Fr11						CAS #: 75-69-4			
3.212	3.212	(0.435)	101	13289588	200.000	190.37	70.00- 130.00	100.00	
3.212	3.212	(0.435)	103	8582872			34.97- 94.97	64.58	

23 Ethanol						CAS #: 64-17-5			
3.571	3.571	(0.483)	45	2070197	200.000	185.71	70.00- 130.00	100.00	
3.571	3.571	(0.483)	43	359833			0.00- 30.00	17.38	
3.571	3.571	(0.483)	46	831102			0.00- 30.00	40.15	

28 Freon 113						CAS #: 76-13-1			
3.931	3.931	(0.532)	151	8576901	200.000	185.69	70.00- 130.00	100.00	
3.931	3.931	(0.532)	153	5456307			33.71- 93.71	63.62	
3.931	3.931	(0.532)	101	10076601			86.34- 146.34	117.49	

29 1,1-Dichloroethene						CAS #: 75-35-4			
3.958	3.958	(0.536)	61	8602773	200.000	187.54	70.00- 130.00	100.00	
3.958	3.958	(0.536)	96	5115455			30.45- 90.45	59.46	
3.958	3.958	(0.536)	98	3278527			8.39- 68.39	38.11	

30 Acetone						CAS #: 67-64-1			
4.124	4.124	(0.558)	58	2890019	200.000	188.12	70.00- 130.00	100.00	
4.124	4.124	(0.558)	43	8716480			0.00- 30.00	301.61	

33 Carbon Disulfide						CAS #: 75-15-0			
4.290	4.290	(0.581)	76	16324453	200.000	189.89	70.00- 130.00	100.00	

34 2-Propanol						CAS #: 67-63-0			
4.318	4.318	(0.584)	45	10466020	200.000	188.42	70.00- 130.00	100.00	
4.318	4.318	(0.584)	43	1932601			0.00- 30.00	18.47	
4.318	4.318	(0.584)	59	413505			0.00- 30.00	3.95	

37 3-Chloropropene						CAS #: 107-05-1			
4.566	4.566	(0.618)	76	2714411	200.000	197.92	70.00- 130.00	100.00	
4.566	4.566	(0.618)	41	8255613			0.00- 30.00	304.14	

40 Methylene Chloride						CAS #: 75-09-2			
4.815	4.815	(0.652)	49	6217595	200.000	181.05	70.00- 130.00	100.00	
4.815	4.815	(0.652)	84	4498494			41.01- 101.01	72.35	
4.815	4.815	(0.652)	51	1878140			0.00- 30.00	30.21	

43 MTBE						CAS #: 1634-04-4			
5.147	5.147	(0.697)	73	9333899	200.000	191.11	70.00- 130.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
43 MTBE (continued)									
5.147	5.147	(0.697)	57	2192207			0.00- 53.82	23.49	
5.147	5.147	(0.697)	41	2086738			0.00- 30.00	22.36	

45 trans-1,2-Dichloroethene					CAS #: 156-60-5				
5.175	5.175	(0.701)	96	5912048	200.000	182.14	70.00- 130.00	100.00	
5.175	5.175	(0.701)	61	8855088			119.04- 179.04	149.78	
5.175	5.175	(0.701)	98	3761119			0.00- 30.00	63.62	

46 Hexane					CAS #: 110-54-3				
5.534	5.534	(0.749)	57	10195455	200.000	207.10	70.00- 130.00	100.00(A)	
5.534	5.534	(0.749)	43	6386604			0.00- 30.00	62.64	
5.534	5.534	(0.749)	86	1626253			0.00- 30.00	15.95	

54 1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	10336900	200.000	195.36	70.00- 130.00	100.00	
5.949	5.949	(0.805)	65	3213837			1.22- 61.22	31.09	

55 Vinyl Acetate					CAS #: 108-05-4				
6.032	6.032	(0.817)	86	1549543	200.000	207.45	70.00- 130.00	100.00(A)	
6.032	6.032	(0.817)	43	16691748			0.00- 30.00	1077.20	
6.032	6.032	(0.817)	42	1228627			0.00- 30.00	79.29	

64 cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.972	6.972	(0.944)	61	7586985	200.000	181.17	70.00- 130.00	100.00	
6.972	6.972	(0.944)	96	5613120			45.09- 105.09	73.98	
6.972	6.972	(0.944)	98	3566701			17.66- 77.66	47.01	

65 2-Butanone					CAS #: 78-93-3				
7.027	7.027	(0.951)	72	2839689	200.000	191.14	70.00- 130.00	100.00	
7.027	7.027	(0.951)	43	12152318			398.56- 458.56	427.95	
7.027	7.027	(0.951)	57	913237			0.00- 30.00	32.16	

67 Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.387	(1.000)	42	7203120	200.000	186.53	70.00- 130.00	100.00	
7.387	7.387	(1.000)	71	2561625			5.14- 65.14	35.56	
7.387	7.387	(1.000)	72	2765395			0.00- 30.00	38.39	

70 Chloroform					CAS #: 67-66-3				
7.525	7.525	(1.019)	83	10146243	200.000	191.74	70.00- 130.00	100.00	
7.525	7.525	(1.019)	85	6278598			31.98- 91.98	61.88	

73 Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	8151077	200.000	185.65	70.00- 130.00	100.00	
7.746	7.746	(1.049)	56	10487225			99.30- 159.30	128.66	
7.746	7.746	(1.049)	41	5082554			33.84- 93.84	62.35	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

75	1,1,1-Trichloroethane				CAS #:		71-55-6		
7.774	7.774	(1.052)	97	10604133	200.000	191.25	70.00-	130.00	100.00
7.774	7.774	(1.052)	99	6750218			34.73-	94.73	63.66

77	Carbon Tetrachloride				CAS #:		56-23-5		
7.995	7.995	(1.082)	119	10305397	200.000	204.25	70.00-	130.00	100.00(A)
7.995	7.995	(1.082)	117	10656606			74.04-	134.04	103.41

81	Benzene				CAS #:		71-43-2		
8.437	8.437	(0.910)	78	17785145	200.000	193.61	70.00-	130.00	100.00
8.437	8.437	(0.910)	77	3896089			0.00-	30.00	21.91

80	2,2,4-Trimethylpentane				CAS #:		540-84-1		
8.465	8.465	(1.146)	57	26545736	200.000	202.53	70.00-	130.00	100.00(A)
8.465	8.465	(1.146)	56	8248292			0.00-	30.00	31.07
8.465	8.465	(1.146)	41	6074066			0.00-	30.00	22.88

83	1,2-Dichloroethane				CAS #:		107-06-2		
8.603	8.603	(0.928)	62	7018378	200.000	188.31	70.00-	130.00	100.00
8.603	8.603	(0.928)	64	2171145			0.00-	30.00	30.94

85	Heptane				CAS #:		142-82-5		
8.852	8.852	(0.955)	100	1749662	200.000	176.00	70.00-	130.00	100.00
8.852	8.852	(0.955)	43	10101737			0.00-	30.00	577.35
8.852	8.852	(0.955)	71	5661505			0.00-	30.00	323.58

94	Trichloroethene				CAS #:		79-01-6		
9.682	9.682	(1.045)	95	6875905	200.000	188.87	70.00-	130.00	100.00
9.682	9.682	(1.045)	130	7400546			77.84-	137.84	107.63
9.682	9.682	(1.045)	97	4412159			34.22-	94.22	64.17

95	Methyl Cyclohexane				CAS #:		108-87-2		
9.903	9.903	(1.341)	83	10900787	200.000	199.28	70.00-	130.00	100.00
9.903	9.903	(1.341)	98	4807211			0.00-	30.00	44.10
9.903	9.903	(1.341)	55	8581305			0.00-	30.00	78.72

97	1,2-Dichloropropane				CAS #:		78-87-5		
10.179	10.179	(1.098)	63	6102724	200.000	191.69	70.00-	130.00	100.00
10.179	10.179	(1.098)	62	4318168			42.15-	102.15	70.76
10.179	10.179	(1.098)	41	3413820			26.47-	86.47	55.94

98	1,4-Dioxane				CAS #:		123-91-1		
10.428	10.428	(1.125)	88	3854134	200.000	198.48	70.00-	130.00	100.00
10.428	10.428	(1.125)	58	2794898			44.03-	104.03	72.52
10.428	10.428	(1.125)	57	841647			0.00-	30.00	21.84

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
==	=====	=====	=====	=====	=====	=====	=====	=====		

100	Bromodichloromethane					CAS #:	75-27-4			
10.732	10.732	(1.158)	83	10477147	200.000	197.04	70.00- 130.00	100.00		
10.732	10.732	(1.158)	85	6434109			31.03- 91.03	61.41		

102	cis-1,3-Dichloropropene					CAS #:	10061-01-5			
11.672	11.672	(1.260)	75	8703573	200.000	196.72	70.00- 130.00	100.00		
11.672	11.672	(1.260)	77	2719406			1.19- 61.19	31.24		
11.672	11.672	(1.260)	39	4066285			16.93- 76.93	46.72		

103	4-Methyl-2-pentanone					CAS #:	108-10-1			
12.032	12.032	(1.298)	58	4830872	200.000	201.84	70.00- 130.00	100.00(A)		
12.032	12.032	(1.298)	43	12195996			0.00- 30.00	252.46		
12.032	12.032	(1.298)	85	2017678			0.00- 30.00	41.77		

105	Toluene					CAS #:	108-88-3			
12.253	12.253	(1.322)	91	18836110	200.000	201.75	70.00- 130.00	100.00(A)		
12.253	12.253	(1.322)	92	11087412			30.46- 90.46	58.86		

108	trans-1,3-Dichloropropene					CAS #:	10061-02-6			
12.834	12.834	(0.880)	75	8893044	200.000	201.20	70.00- 130.00	100.00(A)		
12.834	12.834	(0.880)	77	2758588			0.11- 60.11	31.02		
12.834	12.834	(0.880)	39	4059278			15.75- 75.75	45.65		

110	1,1,2-Trichloroethane					CAS #:	79-00-5			
13.138	13.138	(0.901)	97	6072445	200.000	187.33	70.00- 130.00	100.00		
13.138	13.138	(0.901)	99	3749841			31.47- 91.47	61.75		
13.138	13.138	(0.901)	83	5216234			58.25- 118.25	85.90		

112	Tetrachloroethene					CAS #:	127-18-4			
13.193	13.193	(0.905)	166	8509996	200.000	193.97	70.00- 130.00	100.00		
13.193	13.193	(0.905)	129	6421962			45.90- 105.90	75.46		
13.193	13.193	(0.905)	131	6172896			43.88- 103.88	72.54		

114	2-Hexanone					CAS #:	591-78-6			
13.580	13.580	(0.932)	58	6995194	200.000	217.79	70.00- 130.00	100.00(A)		
13.580	13.580	(0.932)	43	12710091			147.03- 207.03	181.70		
13.580	13.580	(0.932)	100	1440713			0.00- 30.00	20.60		

116	Dibromochloromethane					CAS #:	124-48-1			
13.718	13.718	(0.941)	129	10582353	200.000	208.30	70.00- 130.00	100.00(A)		
13.718	13.718	(0.941)	127	8100146			0.00- 30.00	76.54		

117	1,2-Dibromoethane					CAS #:	106-93-4			
13.884	13.884	(0.953)	107	9615029	200.000	194.66	70.00- 130.00	100.00		
13.884	13.884	(0.953)	109	9190853			65.18- 125.18	95.59		

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
126 Chlorobenzene						CAS #: 108-90-7			
14.603	14.603	(1.002)	112	15794834	200.000	201.76	70.00- 130.00	100.00(A)	
14.631	14.631	(1.004)	114	4886367			1.91- 61.91	30.94	
14.603	14.603	(1.002)	77	8721903			26.50- 86.50	55.22	

129 Ethyl Benzene						CAS #: 100-41-4			
14.769	14.769	(1.013)	106	8105324	200.000	195.65	70.00- 130.00	100.00	
14.769	14.769	(1.013)	91	22645192			0.00- 30.00	279.39	

130 m,p-Xylene						CAS #: 108-38-3			
14.935	14.935	(1.025)	106	10381915	200.000	205.65	70.00- 130.00	100.00(A)	
14.935	14.935	(1.025)	91	19724408			0.00- 30.00	189.99	

132 o-Xylene						CAS #: 95-47-6			
15.488	15.488	(1.063)	106	10154080	200.000	199.32	70.00- 130.00	100.00	
15.488	15.488	(1.063)	91	18196909			177.47- 237.47	179.21	

134 Styrene						CAS #: 100-42-5			
15.516	15.516	(1.064)	104	15909823	200.000	219.09	70.00- 130.00	100.00(A)	
15.516	15.516	(1.064)	78	7701877			20.25- 80.25	48.41	

135 Bromoform						CAS #: 75-25-2			
15.765	15.765	(1.082)	173	10020398	200.000	224.24	70.00- 130.00	100.00(A)	
15.765	15.765	(1.082)	171	5036697			21.05- 81.05	50.26	

137 Cumene						CAS #: 98-82-8			
15.958	15.958	(1.095)	105	26088631	200.000	187.36	70.00- 130.00	100.00	
15.958	15.958	(1.095)	120	7362914			0.00- 30.00	28.22	
15.958	15.958	(1.095)	51	2543543			0.00- 30.00	9.75	

144 1,1,2,2-Tetrachloroethane						CAS #: 79-34-5			
16.456	16.456	(1.129)	83	14918929	200.000	200.26	70.00- 130.00	100.00(A)	
16.456	16.456	(1.129)	85	9007640			31.99- 91.99	60.38	

145 Propylbenzene						CAS #: 103-65-1			
16.483	16.483	(1.131)	91	23743906	200.000	148.89	70.00- 130.00	100.00	
16.483	16.483	(1.131)	120	7677511			0.00- 30.00	32.33	
16.483	16.483	(1.131)	105	1188043			0.00- 30.00	5.00	

147 4-Ethyltoluene						CAS #: 622-96-8			
16.649	16.649	(1.142)	105	21121664	200.000	154.48	70.00- 130.00	100.00	
16.649	16.649	(1.142)	120	9170339			0.00- 59.60	43.42	

148 1,3,5-Trimethylbenzene						CAS #: 108-67-8			
16.732	16.732	(1.148)	105	18184900	200.000	138.37	70.00- 130.00	100.00	
16.732	16.732	(1.148)	120	14177241			0.00- 30.00	77.96	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	

153	17.147	17.147 (1.176)	105	16318393	200.000	123.36	70.00- 130.00	100.00	
	17.147	17.147 (1.176)	120	12694037			15.41- 75.41	77.79	

156	17.451	17.451 (1.197)	146	16249745	200.000	188.43	70.00- 130.00	100.00	
	17.451	17.451 (1.197)	148	12045123			0.00- 30.00	74.12	
	17.451	17.451 (1.197)	111	7776993			0.00- 30.00	47.86	

157	17.562	17.562 (1.205)	146	15969247	200.000	195.40	70.00- 130.00	100.00	
	17.562	17.562 (1.205)	148	11683869			0.00- 30.00	73.16	
	17.562	17.562 (1.205)	111	6235138			0.00- 30.00	39.04	

158	17.700	17.700 (1.214)	91	16188926	200.000	136.28	70.00- 130.00	100.00	
	17.700	17.700 (1.214)	126	5969592			0.00- 30.00	36.87	

161	17.921	17.921 (1.230)	146	16002972	200.000	192.64	70.00- 130.00	100.00	
	17.921	17.921 (1.230)	148	10877445			32.70- 92.70	67.97	
	17.921	17.921 (1.230)	111	6353120			7.07- 67.07	39.70	

167	19.276	19.276 (1.322)	180	14910912	200.000	177.60	70.00- 130.00	100.00	
	19.276	19.276 (1.322)	182	14865436			65.19- 125.19	99.70	

168	19.359	19.359 (1.328)	225	8939871	200.000	185.51	70.00- 130.00	100.00	
	19.359	19.359 (1.328)	223	5565949			33.26- 93.26	62.26	

169	19.470	19.470 (1.336)	128	16033683	200.000	90.554	70.00- 130.00	100.00	
	19.470	19.470 (1.336)	127	5377959			0.00- 30.00	33.54	

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Report Date: 31-May-2007 14:53

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 30-MAY-2007

Lab File ID: 8053009.d

Calibration Time: 16:03

Lab Smp Id: ICAL

Client Smp ID: Level 7

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: db

Method File: /chem/msd8.i/8-30may.b/t14q530a.m

Misc Info: 200ppbv-200ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	441133	264680	617586	448309	1.63
88 1,4-Difluorobenze	1992312	1195387	2789237	2033490	2.07
125 Chlorobenzene-d5	1475337	885202	2065472	1524596	3.34

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

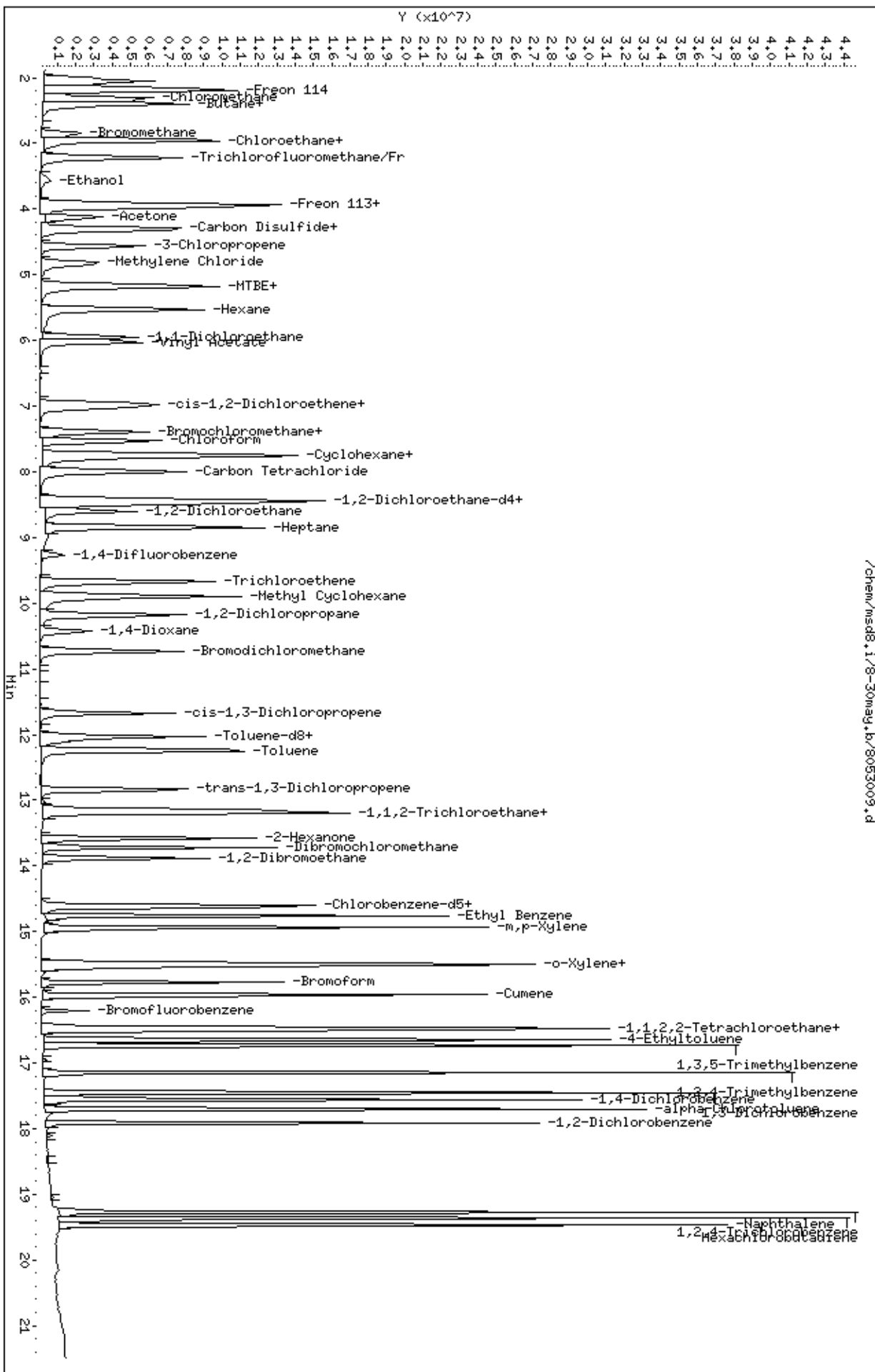
RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-30may.b/8053009.d
 Date: 30-May-2007 17:02
 Client ID: Level 7
 Sample Info: 200ml #1487-289

Column phase: RTX-624

Instrument: msd8.1
 Operator: db
 Column diameter: 0.53





AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: CCV

Lab ID#: 0706440-06A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070302	Date of Collection: NA
Dil. Factor:	1.00	Date of Analysis: 7/3/07 09:41 AM

Compound	%Recovery
Freon 12	71
Freon 114	88
Vinyl Chloride	86
Bromomethane	93
Chloroethane	84
Freon 11	90
1,1-Dichloroethene	86
Freon 113	87
Methylene Chloride	88
1,1-Dichloroethane	86
cis-1,2-Dichloroethene	81
Chloroform	79
1,1,1-Trichloroethane	85
Carbon Tetrachloride	92
Benzene	79
1,2-Dichloroethane	90
Trichloroethene	86
1,2-Dichloropropane	83
cis-1,3-Dichloropropene	84
Toluene	83
trans-1,3-Dichloropropene	87
1,1,2-Trichloroethane	81
Tetrachloroethene	88
1,2-Dibromoethane (EDB)	85
Chlorobenzene	84
Ethyl Benzene	81
m,p-Xylene	85
o-Xylene	80
Styrene	83
1,1,2,2-Tetrachloroethane	78
1,3,5-Trimethylbenzene	81
1,2,4-Trimethylbenzene	87
1,3-Dichlorobenzene	89
1,4-Dichlorobenzene	78
alpha-Chlorotoluene	85
1,2-Dichlorobenzene	75
1,3-Butadiene	83
Hexane	88
Cyclohexane	77



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: CCV

Lab ID#: 0706440-06A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070302	Date of Collection: NA
Dil. Factor:	1.00	Date of Analysis: 7/3/07 09:41 AM

Compound	%Recovery
Heptane	87
Bromodichloromethane	87
Dibromochloromethane	89
Cumene	86
Propylbenzene	93
Chloromethane	96
1,2,4-Trichlorobenzene	77
Hexachlorobutadiene	109
Acetone	88
Carbon Disulfide	84
2-Propanol	85
trans-1,2-Dichloroethene	82
2-Butanone (Methyl Ethyl Ketone)	79
Tetrahydrofuran	84
1,4-Dioxane	84
4-Methyl-2-pentanone	86
2-Hexanone	79
Bromoform	95
4-Ethyltoluene	84
Ethanol	91
Methyl tert-butyl ether	108
3-Chloropropene	93
2,2,4-Trimethylpentane	76
Naphthalene	82

Container Type: NA - Not Applicable

Surrogates	%Recovery	Method Limits
Toluene-d8	100	70-130
1,2-Dichloroethane-d4	103	70-130
4-Bromofluorobenzene	106	70-130

Report Date: 03-Jul-2007 12:18

Air Toxics Ltd.

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: msd8.i Injection Date: 03-JUL-2007 09:41
 Lab File ID: 8070302.d Init. Cal. Date(s): 30-MAY-2007 07-JUN-2007
 Analysis Type: AIR Init. Cal. Times: 14:12 12:08
 Lab Sample ID: CCV-1 Quant Type: ISTD
 Method: /chem/msd8.i/8-03jul.b/t14q530b.m

COMPOUND	RRF / AMOUNT	RF50	MIN	MAX	CURVE TYPE	
			RRF	%D / %DRIFT	%D / %DRIFT	
\$ 82 1,2-Dichloroethane-d4	1.30843	1.34331	0.010	-2.66641	30.00000	Averaged
\$ 104 Toluene-d8	0.86549	0.86923	0.010	-0.43180	30.00000	Averaged
\$ 140 Bromofluorobenzene	0.59482	0.62959	0.010	-5.84412	30.00000	Averaged
3 Propylene	1.35311	1.15293	0.010	14.79416	30.00000	Averaged
4 Dichlorodifluoromethane/Fr1	3.42502	2.43812	0.010	28.81428	30.00000	Averaged
6 Freon 114	3.09425	2.71791	0.010	12.16265	30.00000	Averaged
8 Chloromethane	1.57733	1.52091	0.010	3.57740	30.00000	Averaged
11 Vinyl Chloride	1.83284	1.57095	0.010	14.28904	30.00000	Averaged
10 1,3-Butadiene	1.56776	1.29822	0.010	17.19266	30.00000	Averaged
13 Bromomethane	1.22449	1.14240	0.010	6.70370	30.00000	Averaged
16 Chloroethane	0.97001	0.81765	0.010	15.70720	30.00000	Averaged
18 Trichlorofluoromethane/Fr11	3.89297	3.50226	0.010	10.03618	30.00000	Averaged
23 Ethanol	0.62165	0.56569	0.010	9.00176	30.00000	Averaged
28 Freon 113	2.57573	2.24356	0.010	12.89606	30.00000	Averaged
29 1,1-Dichloroethene	2.55796	2.20735	0.010	13.70667	30.00000	Averaged
30 Acetone	0.85672	0.75285	0.010	12.12375	30.00000	Averaged
34 2-Propanol	3.09759	2.62690	0.010	15.19517	30.00000	Averaged
33 Carbon Disulfide	4.79398	4.01180	0.010	16.31589	30.00000	Averaged
37 3-Chloropropene	0.76480	0.70909	0.010	7.28450	30.00000	Averaged
40 Methylene Chloride	1.91508	1.69528	0.010	11.47740	30.00000	Averaged
43 MTBE	2.72362	2.95657	0.010	-8.55301	30.00000	Averaged
45 trans-1,2-Dichloroethene	1.81006	1.48694	0.010	17.85098	30.00000	Averaged
46 Hexane	2.74528	2.41562	0.010	12.00817	30.00000	Averaged
54 1,1-Dichloroethane	2.95068	2.52911	0.010	14.28723	30.00000	Averaged
55 Vinyl Acetate	0.41654	0.36010	0.010	13.55162	30.00000	Averaged
65 2-Butanone	0.82850	0.65740	0.010	20.65160	30.00000	Averaged
64 cis-1,2-Dichloroethene	2.33530	1.89912	0.010	18.67774	30.00000	Averaged
67 Tetrahydrofuran	2.15347	1.80533	0.010	16.16637	30.00000	Averaged
70 Chloroform	3.17561	2.52424	0.010	20.51162	30.00000	Averaged
75 1,1,1-Trichloroethane	3.09194	2.63278	0.010	14.85015	30.00000	Averaged
73 Cyclohexane	2.44845	1.87897	0.010	23.25905	30.00000	Averaged
77 Carbon Tetrachloride	2.81355	2.59442	0.010	7.78837	30.00000	Averaged
80 2,2,4-Trimethylpentane	7.30928	5.55972	0.010	23.93611	30.00000	Averaged
81 Benzene	1.17761	0.93562	0.010	20.54962	30.00000	Averaged
83 1,2-Dichloroethane	0.45822	0.41177	0.010	10.13694	30.00000	Averaged

Air Toxics Ltd.

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: msd8.i Injection Date: 03-JUL-2007 09:41
 Lab File ID: 8070302.d Init. Cal. Date(s): 30-MAY-2007 07-JUN-2007
 Analysis Type: AIR Init. Cal. Times: 14:12 12:08
 Lab Sample ID: CCV-1 Quant Type: ISTD
 Method: /chem/msd8.i/8-03jul.b/t14q530b.m

COMPOUND	RRF / AMOUNT	RF50	MIN		MAX		CURVE TYPE
			RRF	%D / %DRIFT	%D / %DRIFT	%D / %DRIFT	
85 Heptane	0.12222	0.10593	0.010	13.32307	30.00000	Averaged	
94 Trichloroethene	0.44757	0.38326	0.010	14.36818	30.00000	Averaged	
97 1,2-Dichloropropane	0.39140	0.32442	0.010	17.11363	30.00000	Averaged	
98 1,4-Dioxane	0.23873	0.20172	0.010	15.50009	30.00000	Averaged	
100 Bromodichloromethane	0.65370	0.56880	0.010	12.98827	30.00000	Averaged	
102 cis-1,3-Dichloropropene	0.54392	0.45648	0.010	16.07633	30.00000	Averaged	
103 4-Methyl-2-pentanone	0.29426	0.25197	0.010	14.37072	30.00000	Averaged	
105 Toluene	1.14782	0.94940	0.010	17.28684	30.00000	Averaged	
108 trans-1,3-Dichloropropene	0.72478	0.62891	0.010	13.22739	30.00000	Averaged	
110 1,1,2-Trichloroethane	0.53155	0.43108	0.010	18.89976	30.00000	Averaged	
112 Tetrachloroethene	0.71942	0.63011	0.010	12.41460	30.00000	Averaged	
114 2-Hexanone	0.52669	0.41865	0.010	20.51222	30.00000	Averaged	
116 Dibromochloromethane	0.83304	0.73836	0.010	11.36544	30.00000	Averaged	
117 1,2-Dibromoethane	0.80993	0.69007	0.010	14.79885	30.00000	Averaged	
126 Chlorobenzene	1.28368	1.08070	0.010	15.81290	30.00000	Averaged	
129 Ethyl Benzene	0.67934	0.55344	0.010	18.53146	30.00000	Averaged	
130 m,p-Xylene	0.82782	0.70312	0.010	15.06313	30.00000	Averaged	
132 o-Xylene	0.83535	0.66569	0.010	20.30976	30.00000	Averaged	
134 Styrene	1.17936	0.98231	0.010	16.70846	30.00000	Averaged	
135 Bromoform	0.73275	0.69348	0.010	5.35944	30.00000	Averaged	
144 1,1,2,2-Tetrachloroethane	1.22162	0.94716	0.010	22.46713	30.00000	Averaged	
147 4-Ethyltoluene	2.24196	1.88648	0.010	15.85610	30.00000	Averaged	
148 1,3,5-Trimethylbenzene	2.15500	1.74088	0.010	19.21653	30.00000	Averaged	
153 1,2,4-Trimethylbenzene	2.16919	1.89778	0.010	12.51195	30.00000	Averaged	
156 1,3-Dichlorobenzene	1.41407	1.26403	0.010	10.61044	30.00000	Averaged	
157 1,4-Dichlorobenzene	1.34013	1.04104	0.010	22.31794	30.00000	Averaged	
158 alpha-Chlorotoluene	1.94799	1.65570	0.010	15.00471	30.00000	Averaged	
161 1,2-Dichlorobenzene	1.36216	1.02533	0.010	24.72766	30.00000	Averaged	
167 1,2,4-Trichlorobenzene	1.37671	1.06113	0.010	22.92313	30.00000	Averaged	
168 Hexachlorobutadiene	0.79022	0.86358	0.010	-9.28292	30.00000	Averaged	
145 Propylbenzene	2.61503	2.42455	0.010	7.28392	30.00000	Averaged	
137 Cumene	2.35689	2.01773	0.010	14.38998	30.00000	Averaged	
169 Naphthalene	2.90342	2.36832	0.010	18.43006	30.00000	Averaged	
9 Butane	0.41193	0.35949	0.010	12.73193	30.00000	Averaged	
15 Isopentane	2.28859	2.12593	0.010	7.10775	30.00000	Averaged	

Air Toxics Ltd.

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: msd8.i Injection Date: 03-JUL-2007 09:41
Lab File ID: 8070302.d Init. Cal. Date(s): 30-MAY-2007 07-JUN-2007
Analysis Type: AIR Init. Cal. Times: 14:12 12:08
Lab Sample ID: CCV-1 Quant Type: ISTD
Method: /chem/msd8.i/8-03jul.b/t14q530b.m

COMPOUND	RRF / AMOUNT	RF50	MIN	MAX	CURVE TYPE	
95 Methyl Cyclohexane	3.05031	2.40267	0.010	21.23209	30.00000	Averaged

Report Date: 03-Jul-2007 12:18

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070302.d
 Lab Smp Id: CCV-1 Client Smp ID: CCV-1
 Inj Date : 03-JUL-2007 09:41
 Operator : jdg Inst ID: msd8.i
 Smp Info : 50ml #1443-137
 Misc Info : 200ppbv-50ppbv
 Comment :
 Method : /chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 12:18 jgray Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1 Continuing Calibration Sample
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	(PPBV)	CAL-AMT	ON-COL	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====	=====
* 68 Bromochloromethane CAS #: 74-97-5									
7.387	7.387	(1.000)	130	345936	25.0000			80.00- 120.00	100.00
7.387	7.387	(1.000)	128	267223				47.25- 107.25	77.25
7.387	7.387	(1.000)	49	501857				115.07- 175.07	145.07

* 88 1,4-Difluorobenzene CAS #: 540-36-3									
9.267	9.267	(1.000)	114	1512218	25.0000			80.00- 120.00	100.00
9.267	9.267	(1.000)	88	226425				0.00- 44.97	14.97

* 125 Chlorobenzene-d5 CAS #: 3114-55-4									
14.576	14.576	(1.000)	117	1120246	25.0000			80.00- 120.00	100.00
14.576	14.576	(1.000)	82	598446				0.00- 30.00	53.42

\$ 82 1,2-Dichloroethane-d4 CAS #: 17060-07-0									
8.465	8.465	(1.146)	65	464701	25.0000	25.667		80.00- 120.00	100.00
8.465	8.465	(1.146)	67	261843				0.00- 30.00	56.35

\$ 104 Toluene-d8 CAS #: 2037-26-5									
12.115	12.115	(1.307)	98	1314464	25.0000	25.108		80.00- 120.00	100.00
12.115	12.115	(1.307)	70	126819				0.00- 30.00	9.65

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
\$ 104 Toluene-d8 (continued)									
12.115	12.115	(1.307)	100	874522			0.00- 30.00	66.53	

\$ 140 Bromofluorobenzene									
						CAS #: 460-00-4			
16.207	16.207	(1.112)	174	705292	25.0000	26.461	80.00- 120.00	100.00	
16.207	16.207	(1.112)	95	820514			86.34- 146.34	116.34	
16.207	16.207	(1.112)	176	677873			66.11- 126.11	96.11	

3 Propylene									
						CAS #: 115-07-1			
2.023	2.023	(0.274)	41	797682	50.0000	42.603	80.00- 120.00	100.00	
2.023	2.023	(0.274)	42	535093			0.00- 30.00	67.08	
2.023	2.023	(0.274)	39	543456			0.00- 30.00	68.13	

4 Dichlorodifluoromethane/Fr12									
						CAS #: 75-71-8			
2.078	2.078	(0.281)	85	1686870	50.0000	35.593	80.00- 120.00	100.00	
2.078	2.078	(0.281)	87	547621			0.00- 30.00	32.46	

6 Freon 114									
						CAS #: 76-14-2			
2.161	2.161	(0.293)	135	1880446	50.0000	43.919	80.00- 120.00	100.00	
2.161	2.161	(0.293)	137	586000			1.16- 61.16	31.16	

8 Chloromethane									
						CAS #: 74-87-3			
2.299	2.299	(0.311)	50	1052273	50.0000	48.211	80.00- 120.00	100.00	
2.299	2.299	(0.311)	52	320766			0.00- 30.00	30.48	

11 Vinyl Chloride									
						CAS #: 75-01-4			
2.437	2.437	(0.330)	62	1086893	50.0000	42.855	80.00- 120.00	100.00	
2.437	2.437	(0.330)	64	333808			0.00- 30.00	30.71	

10 1,3-Butadiene									
						CAS #: 106-99-0			
2.410	2.410	(0.326)	54	898204	50.0000	41.404	80.00- 120.00	100.00	
2.410	2.410	(0.326)	39	1077694			0.00- 30.00	119.98	

13 Bromomethane									
						CAS #: 74-83-9			
2.880	2.880	(0.390)	94	790395	50.0000	46.648	80.00- 120.00	100.00	
2.880	2.880	(0.390)	96	741488			63.81- 123.81	93.81	

16 Chloroethane									
						CAS #: 75-00-3			
2.963	2.963	(0.401)	64	565708	50.0000	42.146	80.00- 120.00	100.00	
2.935	2.935	(0.397)	49	153885			0.00- 30.00	27.20	
2.963	2.963	(0.401)	66	176124			0.00- 30.00	31.13	

18 Trichlorofluoromethane/Fr11									
						CAS #: 75-69-4			
3.212	3.212	(0.435)	101	2423118	50.0000	44.982	80.00- 120.00	100.00	
3.239	3.239	(0.439)	103	1576036			35.04- 95.04	65.04	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
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23 Ethanol						CAS #: 64-17-5			
3.516	3.516	(0.476)	45	391384	50.0000	45.499	80.00- 120.00	100.00	
3.516	3.516	(0.476)	43	83465			0.00- 30.00	21.33	
3.516	3.516	(0.476)	46	156156			0.00- 30.00	39.90	

28 Freon 113						CAS #: 76-13-1			
3.958	3.958	(0.536)	151	1552258	50.0000	43.552	80.00- 120.00	100.00	
3.958	3.958	(0.536)	153	1007728			34.92- 94.92	64.92	
3.931	3.931	(0.532)	101	1767499			83.87- 143.87	113.87	

29 1,1-Dichloroethene						CAS #: 75-35-4			
3.986	3.986	(0.540)	61	1527203	50.0000	43.147	80.00- 120.00	100.00	
3.986	3.986	(0.540)	96	894539			28.57- 88.57	58.57	
3.986	3.986	(0.540)	98	580714			8.02- 68.02	38.02	

30 Acetone						CAS #: 67-64-1			
4.124	4.124	(0.558)	58	520877	50.0000	43.938	80.00- 120.00	100.00	
4.124	4.124	(0.558)	43	1608692			0.00- 30.00	308.84	

34 2-Propanol						CAS #: 67-63-0			
4.318	4.318	(0.584)	45	1817480	50.0000	42.402	80.00- 120.00	100.00	
4.318	4.318	(0.584)	43	386785			0.00- 30.00	21.28	
4.318	4.318	(0.584)	59	68606			0.00- 30.00	3.77	

33 Carbon Disulfide						CAS #: 75-15-0			
4.290	4.290	(0.581)	76	2775652	50.0000	41.842	80.00- 120.00	100.00	

37 3-Chloropropene						CAS #: 107-05-1			
4.594	4.594	(0.622)	76	490597	50.0000	46.358	80.00- 120.00	100.00	
4.566	4.566	(0.618)	41	1489477			0.00- 30.00	303.60	

40 Methylene Chloride						CAS #: 75-09-2			
4.815	4.815	(0.652)	49	1172915	50.0000	44.261	80.00- 120.00	100.00	
4.815	4.815	(0.652)	84	793716			37.67- 97.67	67.67	
4.815	4.815	(0.652)	51	350864			0.00- 30.00	29.91	

43 MTBE						CAS #: 1634-04-4			
5.175	5.175	(0.701)	73	2045568	50.0000	54.276	80.00- 120.00	100.00	
5.175	5.175	(0.701)	57	493559			0.00- 54.13	24.13	
5.147	5.147	(0.697)	41	505369			0.00- 30.00	24.71	

45 trans-1,2-Dichloroethene						CAS #: 156-60-5			
5.202	5.202	(0.704)	96	1028774	50.0000	41.074	80.00- 120.00	100.00	
5.202	5.202	(0.704)	61	1541557			119.84- 179.84	149.84	
5.202	5.202	(0.704)	98	666476			0.00- 30.00	64.78	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
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46 Hexane					CAS #: 110-54-3				
5.534	5.534	(0.749)	57	1671300	50.0000	43.996	80.00- 120.00	100.00	
5.534	5.534	(0.749)	43	1142700			0.00- 30.00	68.37	
5.534	5.534	(0.749)	86	260328			0.00- 30.00	15.58	

54 1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	1749818	50.0000	42.856	80.00- 120.00	100.00	
5.949	5.949	(0.805)	65	562732			2.16- 62.16	32.16	

55 Vinyl Acetate					CAS #: 108-05-4				
6.032	6.032	(0.817)	86	249140	50.0000	43.224	80.00- 120.00	100.00	
6.032	6.032	(0.817)	43	2736813			0.00- 30.00	1098.50	
6.032	6.032	(0.817)	42	215402			0.00- 30.00	86.46	

65 2-Butanone					CAS #: 78-93-3				
7.027	7.027	(0.951)	72	454836	50.0000	39.674	80.00- 120.00	100.00	
7.027	7.027	(0.951)	43	2050395			420.80- 480.80	450.80	
7.027	7.027	(0.951)	57	153603			0.00- 30.00	33.77	

64 cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.972	6.972	(0.944)	61	1313949	50.0000	40.661	80.00- 120.00	100.00	
6.972	6.972	(0.944)	96	952972			42.53- 102.53	72.53	
6.972	6.972	(0.944)	98	616245			16.90- 76.90	46.90	

67 Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.387	(1.000)	42	1249056	50.0000	41.917	80.00- 120.00	100.00	
7.387	7.387	(1.000)	71	426402			4.14- 64.14	34.14	
7.387	7.387	(1.000)	72	457937			0.00- 30.00	36.66	

70 Chloroform					CAS #: 67-66-3				
7.525	7.525	(1.019)	83	1746449	50.0000	39.744	80.00- 120.00	100.00	
7.525	7.525	(1.019)	85	1089812			32.40- 92.40	62.40	

75 1,1,1-Trichloroethane					CAS #: 71-55-6				
7.774	7.774	(1.052)	97	1821550	50.0000	42.575	80.00- 120.00	100.00	
7.774	7.774	(1.052)	99	1140569			32.62- 92.62	62.62	

73 Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	1300004	50.0000	38.370	80.00- 120.00	100.00	
7.746	7.746	(1.049)	56	1748924			104.53- 164.53	134.53	
7.746	7.746	(1.049)	41	895089			38.85- 98.85	68.85	

77 Carbon Tetrachloride					CAS #: 56-23-5				
7.995	7.995	(1.082)	119	1795008	50.0000	46.106	80.00- 120.00	100.00	
7.995	7.995	(1.082)	117	1840898			72.56- 132.56	102.56	

AMOUNTS										
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO		
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80	2,2,4-Trimethylpentane					CAS #: 540-84-1				
8.465	8.465	(1.146)	57	3846617	50.0000	38.032	80.00- 120.00	100.00		
8.465	8.465	(1.146)	56	1243228			0.00- 30.00	32.32		
8.465	8.465	(1.146)	41	954069			0.00- 30.00	24.80		

81	Benzene					CAS #: 71-43-2				
8.437	8.437	(0.910)	78	2829721	50.0000	39.725	80.00- 120.00	100.00		
8.437	8.437	(0.910)	77	650913			0.00- 30.00	23.00		

83	1,2-Dichloroethane					CAS #: 107-06-2				
8.603	8.603	(0.928)	62	1245363	50.0000	44.932	80.00- 120.00	100.00		
8.603	8.603	(0.928)	64	388708			0.00- 30.00	31.21		

85	Heptane					CAS #: 142-82-5				
8.852	8.852	(0.955)	100	320391	50.0000	43.338	80.00- 120.00	100.00		
8.852	8.852	(0.955)	43	1841976			0.00- 30.00	574.92		
8.852	8.852	(0.955)	71	979212			0.00- 30.00	305.63		

94	Trichloroethene					CAS #: 79-01-6				
9.682	9.682	(1.045)	95	1159143	50.0000	42.816	80.00- 120.00	100.00		
9.682	9.682	(1.045)	130	1240778			77.04- 137.04	107.04		
9.682	9.682	(1.045)	97	734565			33.37- 93.37	63.37		

97	1,2-Dichloropropane					CAS #: 78-87-5				
10.179	10.179	(1.098)	63	981175	50.0000	41.443	80.00- 120.00	100.00		
10.179	10.179	(1.098)	62	714021			42.77- 102.77	72.77		
10.179	10.179	(1.098)	41	608053			31.97- 91.97	61.97		

98	1,4-Dioxane					CAS #: 123-91-1				
10.428	10.428	(1.125)	88	610100	50.0000	42.250	80.00- 120.00	100.00		
10.428	10.428	(1.125)	58	468421			46.78- 106.78	76.78		
10.428	10.428	(1.125)	57	146481			0.00- 30.00	24.01		

100	Bromodichloromethane					CAS #: 75-27-4				
10.732	10.732	(1.158)	83	1720296	50.0000	43.506	80.00- 120.00	100.00		
10.732	10.732	(1.158)	85	1061670			31.71- 91.71	61.71		

102	cis-1,3-Dichloropropene					CAS #: 10061-01-5				
11.672	11.672	(1.260)	75	1380591	50.0000	41.962	80.00- 120.00	100.00		
11.672	11.672	(1.260)	77	437051			1.66- 61.66	31.66		
11.672	11.672	(1.260)	39	717038			21.94- 81.94	51.94		

103	4-Methyl-2-pentanone					CAS #: 108-10-1				
12.032	12.032	(1.298)	58	762063	50.0000	42.815	80.00- 120.00	100.00		
12.032	12.032	(1.298)	43	1942464			0.00- 30.00	254.90		
12.032	12.032	(1.298)	85	315989			0.00- 30.00	41.46		

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
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105 Toluene						CAS #: 108-88-3			
12.225	12.225	(1.319)	91	2871405	50.0000	41.356	80.00- 120.00	100.00	
12.225	12.225	(1.319)	92	1724784			30.07- 90.07	60.07	

108 trans-1,3-Dichloropropene						CAS #: 10061-02-6			
12.834	12.834	(0.880)	75	1409060	50.0000	43.386	80.00- 120.00	100.00	
12.834	12.834	(0.880)	77	433607			0.77- 60.77	30.77	
12.834	12.834	(0.880)	39	691519			19.08- 79.08	49.08	

110 1,1,2-Trichloroethane						CAS #: 79-00-5			
13.138	13.138	(0.901)	97	965841	50.0000	40.550	80.00- 120.00	100.00	
13.138	13.138	(0.901)	99	598592			31.98- 91.98	61.98	
13.138	13.138	(0.901)	83	838646			56.83- 116.83	86.83	

112 Tetrachloroethene						CAS #: 127-18-4			
13.193	13.193	(0.905)	166	1411757	50.0000	43.793	80.00- 120.00	100.00	
13.193	13.193	(0.905)	129	1069106			45.73- 105.73	75.73	
13.193	13.193	(0.905)	131	1029397			42.92- 102.92	72.92	

114 2-Hexanone						CAS #: 591-78-6			
13.580	13.580	(0.932)	58	937985	50.0000	39.744	80.00- 120.00	100.00	
13.580	13.580	(0.932)	43	1778479			159.61- 219.61	189.61	
13.580	13.580	(0.932)	100	210412			0.00- 30.00	22.43	

116 Dibromochloromethane						CAS #: 124-48-1			
13.718	13.718	(0.941)	129	1654298	50.0000	44.317	80.00- 120.00	100.00	
13.718	13.718	(0.941)	127	1306341			0.00- 30.00	78.97	

117 1,2-Dibromoethane						CAS #: 106-93-4			
13.884	13.884	(0.953)	107	1546100	50.0000	42.600	80.00- 120.00	100.00	
13.884	13.884	(0.953)	109	1468071			64.95- 124.95	94.95	

126 Chlorobenzene						CAS #: 108-90-7			
14.603	14.603	(1.002)	112	2421290	50.0000	42.094	80.00- 120.00	100.00	
14.603	14.603	(1.002)	114	775897			2.04- 62.04	32.04	
14.603	14.603	(1.002)	77	1385709			27.23- 87.23	57.23	

129 Ethyl Benzene						CAS #: 100-41-4			
14.769	14.769	(1.013)	106	1239988	50.0000	40.734	80.00- 120.00	100.00	
14.769	14.769	(1.013)	91	3833499			0.00- 30.00	309.16	

130 m,p-Xylene						CAS #: 108-38-3			
14.935	14.935	(1.025)	106	1575341	50.0000	42.468	80.00- 120.00	100.00	
14.935	14.935	(1.025)	91	3079533			0.00- 30.00	195.48	

132 o-Xylene						CAS #: 95-47-6			
15.488	15.488	(1.063)	106	1491473	50.0000	39.845	80.00- 120.00	100.00	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
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132 o-Xylene (continued)									
15.488	15.488	(1.063)	91	3069708			175.82- 235.82	205.82	

134 Styrene CAS #: 100-42-5									
15.516	15.516	(1.064)	104	2200860	50.0000	41.646	80.00- 120.00	100.00	
15.516	15.516	(1.064)	78	1172145			23.26- 83.26	53.26	

135 Bromoform CAS #: 75-25-2									
15.765	15.765	(1.082)	173	1553731	50.0000	47.320	80.00- 120.00	100.00	
15.765	15.765	(1.082)	171	793970			21.10- 81.10	51.10	

144 1,1,2,2-Tetrachloroethane CAS #: 79-34-5									
16.456	16.456	(1.129)	83	2122094	50.0000	38.766	80.00- 120.00	100.00	
16.456	16.456	(1.129)	85	1312582			31.85- 91.85	61.85	

147 4-Ethyltoluene CAS #: 622-96-8									
16.649	16.649	(1.142)	105	4226634	50.0000	42.072	80.00- 120.00	100.00	
16.649	16.649	(1.142)	120	1297770			0.70- 60.70	30.70	

148 1,3,5-Trimethylbenzene CAS #: 108-67-8									
16.732	16.732	(1.148)	105	3900430	50.0000	40.392	80.00- 120.00	100.00	
16.732	16.732	(1.148)	120	1980510			0.00- 30.00	50.78	

153 1,2,4-Trimethylbenzene CAS #: 95-63-6									
17.147	17.147	(1.176)	105	4251969	50.0000	43.744	80.00- 120.00	100.00	
17.147	17.147	(1.176)	120	1993469			16.88- 76.88	46.88	

156 1,3-Dichlorobenzene CAS #: 541-73-1									
17.451	17.451	(1.197)	146	2832058	50.0000	44.695	80.00- 120.00	100.00	
17.451	17.451	(1.197)	148	1804580			0.00- 30.00	63.72	
17.451	17.451	(1.197)	111	1110308			0.00- 30.00	39.20	

157 1,4-Dichlorobenzene CAS #: 106-46-7									
17.562	17.562	(1.205)	146	2332451	50.0000	38.841	80.00- 120.00	100.00	
17.562	17.562	(1.205)	148	1511896			0.00- 30.00	64.82	
17.562	17.562	(1.205)	111	866983			0.00- 30.00	37.17	

158 alpha-Chlorotoluene CAS #: 100-44-7									
17.700	17.700	(1.214)	91	3709577	50.0000	42.498	80.00- 120.00	100.00	
17.700	17.700	(1.214)	126	829027			0.00- 30.00	22.35	

161 1,2-Dichlorobenzene CAS #: 95-50-1									
17.921	17.921	(1.230)	146	2297242	50.0000	37.636	80.00- 120.00	100.00	
17.921	17.921	(1.230)	148	1477315			34.31- 94.31	64.31	
17.921	17.921	(1.230)	111	924019			10.22- 70.22	40.22	

AMOUNTS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	CAL-AMT (PPEV)	ON-COL (PPBV)	TARGET RANGE	RATIO	
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167	1,2,4-Trichlorobenzene					CAS #: 120-82-1			
19.276	19.276	(1.322)	180	2377446	50.0000	38.538	80.00- 120.00	100.00	
19.276	19.276	(1.322)	182	2310203			67.17- 127.17	97.17	

168	Hexachlorobutadiene					CAS #: 87-68-3			
19.359	19.359	(1.328)	225	1934833	50.0000	54.641	80.00- 120.00	100.00	
19.359	19.359	(1.328)	223	1232092			33.68- 93.68	63.68	

145	Propylbenzene					CAS #: 103-65-1			
16.483	16.483	(1.131)	91	5432193	50.0000	46.358	80.00- 120.00	100.00	
16.483	16.483	(1.131)	120	1253064			0.00- 30.00	23.07	
16.483	16.483	(1.131)	105	193545			0.00- 30.00	3.56	

137	Cumene					CAS #: 98-82-8			
15.958	15.958	(1.095)	105	4520711	50.0000	42.805	80.00- 120.00	100.00	
15.958	15.958	(1.095)	120	1200281			0.00- 30.00	26.55	
15.958	15.958	(1.095)	51	447430			0.00- 30.00	9.90	

169	Naphthalene					CAS #: 91-20-3			
19.470	19.470	(1.336)	128	5306192	50.0000	40.785	80.00- 120.00	100.00	
19.470	19.470	(1.336)	127	643005			0.00- 30.00	12.12	

9	Butane					CAS #: 106-97-8			
2.355	2.355	(0.319)	58	248718	50.0000	43.634	80.00- 120.00	100.00	
2.355	2.355	(0.319)	43	1885509			0.00- 30.00	758.09	

15	Isopentane					CAS #: 78-78-4			
2.963	2.963	(0.401)	43	1470868	50.0000	46.446	80.00- 120.00	100.00	
2.963	2.963	(0.401)	57	967997			0.00- 30.00	65.81	
2.963	2.963	(0.401)	72	99701			0.00- 30.00	6.78	

95	Methyl Cyclohexane					CAS #: 108-87-2			
9.903	9.903	(1.341)	83	1662337	50.0000	39.384	80.00- 120.00	100.00	
9.903	9.903	(1.341)	98	764465			0.00- 30.00	45.99	
9.903	9.903	(1.341)	55	1372915			0.00- 30.00	82.59	

Report Date: 03-Jul-2007 12:18

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 03-JUL-2007

Lab File ID: 8070302.d

Calibration Time: 11:45

Lab Smp Id: CCV-1

Client Smp ID: CCV-1

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdj

Method File: /chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 200ppbv-50ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	273335	164001	382669	345936	26.56
88 1,4-Difluorobenze	1172440	703464	1641416	1512218	28.98
125 Chlorobenzene-d5	875856	525514	1226198	1120246	27.90

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

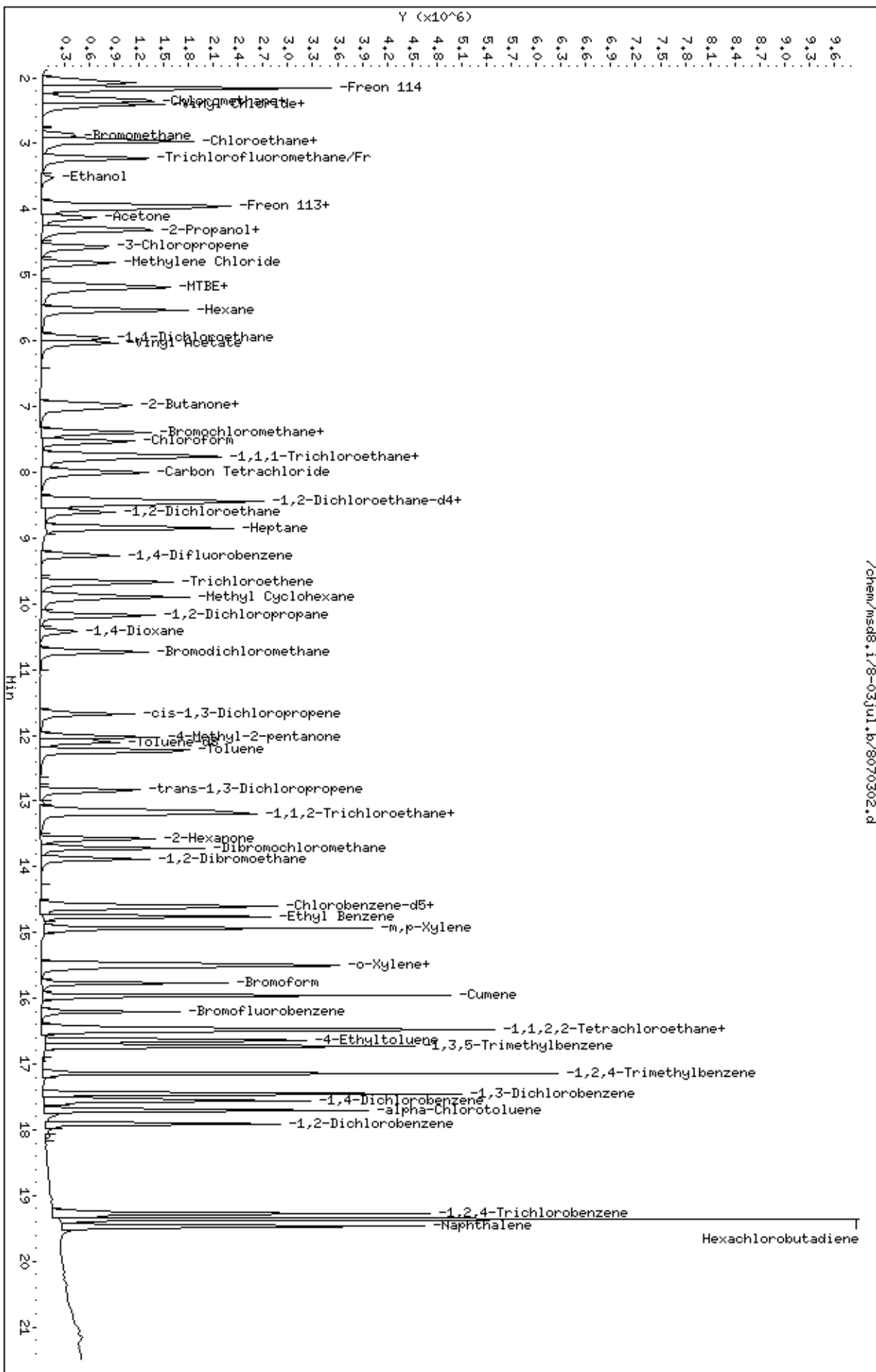
RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-03jul.b/8070302.d
Date: 03-JUL-2007 09:41
Client ID: CCV-1
Sample Info: 50ml #1443-137

Column phase: RTX-624

Instrument: msd8.1
Operator: jdg
Column diameter: 0.53

/chem/msd8.1/8-03jul.b/8070302.d





AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: LCS

Lab ID#: 0706440-07A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070303	Date of Collection: NA
Dil. Factor:	1.00	Date of Analysis: 7/3/07 10:09 AM

Compound	%Recovery
Freon 12	88
Freon 114	86
Vinyl Chloride	84
Bromomethane	92
Chloroethane	85
Freon 11	88
1,1-Dichloroethene	97
Freon 113	98
Methylene Chloride	96
1,1-Dichloroethane	92
cis-1,2-Dichloroethene	83
Chloroform	81
1,1,1-Trichloroethane	88
Carbon Tetrachloride	93
Benzene	85
1,2-Dichloroethane	97
Trichloroethene	92
1,2-Dichloropropane	87
cis-1,3-Dichloropropene	87
Toluene	93
trans-1,3-Dichloropropene	91
1,1,2-Trichloroethane	86
Tetrachloroethene	93
1,2-Dibromoethane (EDB)	88
Chlorobenzene	89
Ethyl Benzene	83
m,p-Xylene	90
o-Xylene	84
Styrene	93
1,1,2,2-Tetrachloroethane	79
1,3,5-Trimethylbenzene	82
1,2,4-Trimethylbenzene	90
1,3-Dichlorobenzene	94
1,4-Dichlorobenzene	80
alpha-Chlorotoluene	90
1,2-Dichlorobenzene	76
1,3-Butadiene	82
Hexane	88
Cyclohexane	79



AN ENVIRONMENTAL ANALYTICAL LABORATORY

Client Sample ID: LCS

Lab ID#: 0706440-07A

MODIFIED EPA METHOD TO-15 GC/MS FULL SCAN

File Name:	8070303	Date of Collection: NA
Dil. Factor:	1.00	Date of Analysis: 7/3/07 10:09 AM

Compound	%Recovery
Heptane	90
Bromodichloromethane	91
Dibromochloromethane	92
Cumene	94
Propylbenzene	100
Chloromethane	97
1,2,4-Trichlorobenzene	80
Hexachlorobutadiene	115
Acetone	88
Carbon Disulfide	85
2-Propanol	86
trans-1,2-Dichloroethene	82
2-Butanone (Methyl Ethyl Ketone)	83
Tetrahydrofuran	85
1,4-Dioxane	90
4-Methyl-2-pentanone	92
2-Hexanone	84
Bromoform	100
4-Ethyltoluene	90
Ethanol	99
Methyl tert-butyl ether	81
3-Chloropropene	92
2,2,4-Trimethylpentane	78
Naphthalene	77

Container Type: NA - Not Applicable

Surrogates	%Recovery	Method Limits
Toluene-d8	103	70-130
1,2-Dichloroethane-d4	102	70-130
4-Bromofluorobenzene	112	70-130

Air Toxics Ltd.

RECOVERY REPORT

Client Name: Client SDG: 8-03jul
 Sample Matrix: GAS Fraction: VOA
 Lab Smp Id: LCS-1 Client Smp ID: LCS-1
 Level: LOW Operator: jdg
 Data Type: MS DATA SampleType: LCS
 SpikeList File: Spectra.spk Quant Type: ISTD
 Sublist File: AT04+ENSR.sub
 Method File: /chem/msd8.i/8-03jul.b/t14q530b.m
 Misc Info: 200ppbv-50ppbv

SPIKE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
134 Styrene	50.000	46.308	92.62	70-130
108 trans-1,3-Dichloro	50.000	45.396	90.79	70-130
3 Propylene	50.000	46.081	92.16	60-140
4 Dichlorodifluorome	50.000	44.268	88.54	70-130
6 Freon 114	50.000	42.994	85.99	70-130
8 Chloromethane	50.000	48.332	96.66	70-130
11 Vinyl Chloride	50.000	42.184	84.37	70-130
10 1,3-Butadiene	50.000	41.126	82.25	60-140
13 Bromomethane	50.000	46.248	92.50	70-130
16 Chloroethane	50.000	42.311	84.62	70-130
18 Trichlorofluoromet	50.000	44.083	88.17	70-130
23 Ethanol	50.000	49.493	98.99	60-140
28 Freon 113	50.000	48.986	97.97	70-130
29 1,1-Dichloroethene	50.000	48.490	96.98	70-130
30 Acetone	50.000	43.937	87.87	60-140
33 Carbon Disulfide	50.000	42.371	84.74	60-140
34 2-Propanol	50.000	43.294	86.59	60-140
40 Methylene Chloride	50.000	48.153	96.31	70-130
43 MTBE	50.000	40.626	81.25	60-140
45 trans-1,2-Dichloro	50.000	41.253	82.51	60-140
46 Hexane	50.000	44.078	88.16	60-140
54 1,1-Dichloroethane	50.000	46.082	92.16	70-130
55 Vinyl Acetate	50.000	42.029	84.06	60-140
64 cis-1,2-Dichloroet	50.000	41.680	83.36	70-130
65 2-Butanone	50.000	41.309	82.62	60-140
67 Tetrahydrofuran	50.000	42.662	85.32	60-140
70 Chloroform	50.000	40.747	81.49	70-130
73 Cyclohexane	50.000	39.488	78.98	60-140
75 1,1,1-Trichloroeth	50.000	43.794	87.59	70-130
77 Carbon Tetrachlori	50.000	46.689	93.38	70-130
81 Benzene	50.000	42.328	84.66	70-130
83 1,2-Dichloroethane	50.000	48.420	96.84	70-130
85 Heptane	50.000	45.070	90.14	60-140

Report Date: 03-Jul-2007 10:39

SPIKE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
94 Trichloroethene	50.000	45.926	91.85	70-130
97 1,2-Dichloropropan	50.000	43.382	86.76	70-130
98 1,4-Dioxane	50.000	45.090	90.18	60-140
100 Bromodichlorometha	50.000	45.700	91.40	60-140
102 cis-1,3-Dichloropr	50.000	43.335	86.67	70-130
103 4-Methyl-2-pentano	50.000	46.060	92.12	60-140
105 Toluene	50.000	46.427	92.85	70-130
110 1,1,2-Trichloroeth	50.000	42.918	85.84	70-130
112 Tetrachloroethene	50.000	46.578	93.16	70-130
114 2-Hexanone	50.000	42.051	84.10	60-140
116 Dibromochlorometha	50.000	45.825	91.65	60-140
117 1,2-Dibromoethane	50.000	43.888	87.78	70-130
126 Chlorobenzene	50.000	44.379	88.76	70-130
129 Ethyl Benzene	50.000	41.546	83.09	70-130
130 m,p-Xylene	50.000	44.959	89.92	70-130
132 o-Xylene	50.000	42.002	84.00	70-130
135 Bromoform	50.000	50.025	100.05	60-140
144 1,1,2,2-Tetrachlor	50.000	39.489	78.98	70-130
147 4-Ethyltoluene	50.000	45.074	90.15	60-140
148 1,3,5-Trimethylben	50.000	41.144	82.29	70-130
153 1,2,4-Trimethylben	50.000	44.852	89.70	70-130
156 1,3-Dichlorobenzen	50.000	46.981	93.96	70-130
157 1,4-Dichlorobenzen	50.000	39.750	79.50	70-130
158 alpha-Chlorotoluen	50.000	45.001	90.00	70-130
161 1,2-Dichlorobenzen	50.000	37.984	75.97	70-130
167 1,2,4-Trichloroben	50.000	40.101	80.20	70-130
168 Hexachlorobutadien	50.000	57.740	115.48	70-130
137 Cumene	50.000	47.133	94.27	60-140
145 Propylbenzene	50.000	50.252	100.50	60-140
37 3-Chloropropene	50.000	46.261	92.52	60-140
80 2,2,4-Trimethylpen	50.000	39.016	78.03	60-140
169 Naphthalene	50.000	38.643	77.29	60-140
9 Butane	50.000	43.039	86.08	70-130
15 Isopentane	50.000	45.686	91.37	70-130
95 Methyl Cyclohexane	50.000	40.292	80.59	70-130

SURROGATE COMPOUND	CONC ADDED PPBV	CONC RECOVERED PPBV	% RECOVERED	LIMITS
\$ 82 1,2-Dichloroethane	25.000	25.497	101.99	70-130
\$ 104 Toluene-d8	25.000	25.693	102.77	70-130
\$ 140 Bromofluorobenzene	25.000	27.917	111.67	70-130

Report Date: 03-Jul-2007 10:39

Air Toxics Ltd.

AMBIENT AIR METHOD TO14A/TO15

Data file : /chem/msd8.i/8-03jul.b/8070303.d
 Lab Smp Id: LCS-1 Client Smp ID: LCS-1
 Inj Date : 03-JUL-2007 10:09
 Operator : jdg Inst ID: msd8.i
 Smp Info : 50ml #1443-146
 Misc Info : 200ppbv-50ppbv
 Comment :
 Method : /chem/msd8.i/8-03jul.b/t14q530b.m
 Meth Date : 03-Jul-2007 09:51 jgray Quant Type: ISTD
 Cal Date : 07-JUN-2007 12:08 Cal File: 8060706.d
 Als bottle: 1 QC Sample: LCS
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: AT04+ENSR.sub
 Target Version: 3.50 Sample Matrix: AIR
 Processing Host: eeyore

Concentration Formula: Amt * DF * CpndVariable

Cpnd Variable Local Compound Variable

CONCENTRATIONS								
RT	EXP RT	(REL RT)	MASS	RESPONSE		TARGET RANGE	RATIO	
				(PPBV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====

* 68	Bromochloromethane					CAS #: 74-97-5		
7.387	7.387	(1.000)	130	278074	25.0000	80.00- 120.00	100.00	
7.387	7.387	(1.000)	128	201465		47.25- 107.25	72.45	
7.387	7.387	(1.000)	49	418940		115.07- 175.07	150.66	

* 88	1,4-Difluorobenzene					CAS #: 540-36-3		
9.267	9.267	(1.000)	114	1169163	25.0000	80.00- 120.00	100.00	
9.267	9.267	(1.000)	88	178860		0.00- 44.97	15.30	

* 125	Chlorobenzene-d5					CAS #: 3114-55-4		
14.576	14.576	(1.000)	117	882830	25.0000	80.00- 120.00	100.00	
14.576	14.576	(1.000)	82	495979		0.00- 30.00	56.18	

\$ 82	1,2-Dichloroethane-d4					CAS #: 17060-07-0		
8.465	8.465	(1.146)	65	371071	25.4969	25.497 80.00- 120.00	100.00	
8.465	8.465	(1.146)	67	222065		0.00- 30.00	59.84	

\$ 104	Toluene-d8					CAS #: 2037-26-5		
12.115	12.115	(1.307)	98	1039944	25.6928	25.693 80.00- 120.00	100.00	
12.087	12.115	(1.304)	70	107685		0.00- 30.00	10.35	

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT (REL RT)	MASS	RESPONSE (PPEV)	(PPBV)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====

\$ 104 Toluene-d8 (continued)

12.115	12.115 (1.307)	100	747694		0.00- 30.00	71.90
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\$ 140 Bromofluorobenzene

CAS #: 460-00-4

16.207	16.207 (1.112)	174	586410	27.9174	27.917	80.00- 120.00	100.00
16.207	16.207 (1.112)	95	675445			86.34- 146.34	115.18
16.207	16.207 (1.112)	176	574515			66.11- 126.11	97.97

3 Propylene

CAS #: 115-07-1

1.995	2.023 (0.270)	41	693553	46.0813	46.081	80.00- 120.00	100.00
1.995	2.023 (0.270)	42	471702			0.00- 30.00	68.01
1.995	2.023 (0.270)	39	458268			0.00- 30.00	66.08

4 Dichlorodifluoromethane/Fr12

CAS #: 75-71-8

2.050	2.078 (0.278)	85	1686445	44.2679	44.268	80.00- 120.00	100.00
2.050	2.078 (0.278)	87	545919			0.00- 30.00	32.37

6 Freon 114

CAS #: 76-14-2

2.133	2.161 (0.289)	135	1479730	42.9938	42.994	80.00- 120.00	100.00
2.133	2.161 (0.289)	137	470484			1.16- 61.16	31.80

8 Chloromethane

CAS #: 74-87-3

2.272	2.299 (0.308)	50	847972	48.3323	48.332	80.00- 120.00	100.00
2.272	2.299 (0.308)	52	251762			0.00- 30.00	29.69

11 Vinyl Chloride

CAS #: 75-01-4

2.410	2.437 (0.326)	62	859992	42.1842	42.184	80.00- 120.00	100.00
2.410	2.437 (0.326)	64	264998			0.00- 30.00	30.81

10 1,3-Butadiene

CAS #: 106-99-0

2.382	2.410 (0.322)	54	717155	41.1256	41.126	80.00- 120.00	100.00
2.382	2.410 (0.322)	39	745460			0.00- 30.00	103.95

13 Bromomethane

CAS #: 74-83-9

2.825	2.880 (0.382)	94	629895	46.2481	46.248	80.00- 120.00	100.00
2.852	2.880 (0.386)	96	589138			63.81- 123.81	93.53

16 Chloroethane

CAS #: 75-00-3

2.935	2.963 (0.397)	64	456507	42.3108	42.311	80.00- 120.00	100.00
2.935	2.935 (0.397)	49	120912			0.00- 30.00	26.49
2.935	2.963 (0.397)	66	136748			0.00- 30.00	29.96

18 Trichlorofluoromethane/Fr11

CAS #: 75-69-4

3.212	3.212 (0.435)	101	1908843	44.0828	44.083	80.00- 120.00	100.00
3.212	3.239 (0.435)	103	1234630			35.04- 95.04	64.68

CONCENTRATIONS

ON-COL FINAL

RT EXP RT (REL RT) MASS RESPONSE (PPEV) (PPBV) TARGET RANGE RATIO
 == == ===== == ===== ===== =====

23 Ethanol CAS #: 64-17-5
 3.516 3.516 (0.476) 45 342222 49.4929 49.493 80.00- 120.00 100.00
 3.516 3.516 (0.476) 43 71293 0.00- 30.00 20.83
 3.516 3.516 (0.476) 46 138791 0.00- 30.00 40.56

28 Freon 113 CAS #: 76-13-1
 3.931 3.958 (0.532) 151 1403442 48.9862 48.986 80.00- 120.00 100.00
 3.931 3.958 (0.532) 153 897198 34.92- 94.92 63.93
 3.931 3.931 (0.532) 101 1581209 83.87- 143.87 112.67

29 1,1-Dichloroethene CAS #: 75-35-4
 3.958 3.986 (0.536) 61 1379628 48.4895 48.490 80.00- 120.00 100.00
 3.958 3.986 (0.536) 96 803469 28.57- 88.57 58.24
 3.958 3.986 (0.536) 98 522498 8.02- 68.02 37.87

30 Acetone CAS #: 67-64-1
 4.124 4.124 (0.558) 58 418690 43.9374 43.937 80.00- 120.00 100.00
 4.096 4.124 (0.555) 43 1300405 0.00- 30.00 310.59

34 2-Propanol CAS #: 67-63-0
 4.290 4.318 (0.581) 45 1491672 43.2942 43.294 80.00- 120.00 100.00
 4.290 4.318 (0.581) 43 308520 0.00- 30.00 20.68
 4.290 4.318 (0.581) 59 59631 0.00- 30.00 4.00

33 Carbon Disulfide CAS #: 75-15-0
 4.290 4.290 (0.581) 76 2259352 42.3709 42.371 80.00- 120.00 100.00

37 3-Chloropropene CAS #: 107-05-1
 4.567 4.594 (0.618) 76 393531 46.2607 46.261 80.00- 120.00 100.00
 4.567 4.566 (0.618) 41 1202229 0.00- 30.00 305.50

40 Methylene Chloride CAS #: 75-09-2
 4.815 4.815 (0.652) 49 1025720 48.1528 48.153 80.00- 120.00 100.00
 4.815 4.815 (0.652) 84 699580 37.67- 97.67 68.20
 4.815 4.815 (0.652) 51 307915 0.00- 30.00 30.02

43 MTBE CAS #: 1634-04-4
 5.147 5.175 (0.697) 73 1230764 40.6264 40.626 80.00- 120.00 100.00
 5.147 5.175 (0.697) 57 291453 0.00- 54.13 23.68
 5.147 5.147 (0.697) 41 293933 0.00- 30.00 23.88

45 trans-1,2-Dichloroethene CAS #: 156-60-5
 5.175 5.202 (0.701) 96 830548 41.2527 41.253 80.00- 120.00 100.00
 5.175 5.202 (0.701) 61 1261947 119.84- 179.84 151.94
 5.175 5.202 (0.701) 98 533520 0.00- 30.00 64.24

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE		ON-COL	FINAL	TARGET RANGE	RATIO
				(PPEV)	(PPEV)				
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

46 Hexane					CAS #: 110-54-3				
5.534	5.534	(0.749)	57	1345951	44.0781	44.078	80.00-	120.00	100.00
5.534	5.534	(0.749)	43	920194			0.00-	30.00	68.37
5.534	5.534	(0.749)	86	208005			0.00-	30.00	15.45

54 1,1-Dichloroethane					CAS #: 75-34-3				
5.949	5.949	(0.805)	63	1512422	46.0820	46.082	80.00-	120.00	100.00
5.949	5.949	(0.805)	65	480396			2.16-	62.16	31.76

55 Vinyl Acetate					CAS #: 108-05-4				
6.032	6.032	(0.817)	86	194728	42.0288	42.029	80.00-	120.00	100.00
6.032	6.032	(0.817)	43	2242918			0.00-	30.00	1151.82
6.032	6.032	(0.817)	42	177879			0.00-	30.00	91.35

65 2-Butanone					CAS #: 78-93-3				
7.000	7.027	(0.948)	72	380677	41.3091	41.309	80.00-	120.00	100.00
7.000	7.027	(0.948)	43	1686845			420.80-	480.80	443.12
7.000	7.027	(0.948)	57	125075			0.00-	30.00	32.86

64 cis-1,2-Dichloroethene					CAS #: 156-59-2				
6.944	6.972	(0.940)	61	1082669	41.6804	41.680	80.00-	120.00	100.00
6.972	6.972	(0.944)	96	790908			42.53-	102.53	73.05
6.972	6.972	(0.944)	98	508829			16.90-	76.90	47.00

67 Tetrahydrofuran					CAS #: 109-99-9				
7.387	7.387	(1.000)	42	1021884	42.6622	42.662	80.00-	120.00	100.00
7.387	7.387	(1.000)	71	341966			4.14-	64.14	33.46
7.387	7.387	(1.000)	72	374351			0.00-	30.00	36.63

70 Chloroform					CAS #: 67-66-3				
7.525	7.525	(1.019)	83	1439278	40.7472	40.747	80.00-	120.00	100.00
7.525	7.525	(1.019)	85	898365			32.40-	92.40	62.42

75 1,1,1-Trichloroethane					CAS #: 71-55-6				
7.774	7.774	(1.052)	97	1506129	43.7936	43.794	80.00-	120.00	100.00
7.774	7.774	(1.052)	99	937825			32.62-	92.62	62.27

73 Cyclohexane					CAS #: 110-82-7				
7.746	7.746	(1.049)	84	1075406	39.4876	39.488	80.00-	120.00	100.00
7.746	7.746	(1.049)	56	1429383			104.53-	164.53	132.92
7.746	7.746	(1.049)	41	742271			38.85-	98.85	69.02

77 Carbon Tetrachloride					CAS #: 56-23-5				
7.995	7.995	(1.082)	119	1461122	46.6886	46.689	80.00-	120.00	100.00
7.995	7.995	(1.082)	117	1501242			72.56-	132.56	102.75

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE		ON-COL	FINAL	TARGET RANGE	RATIO
				(PPEV)	(PPEV)				
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

80	2,2,4-Trimethylpentane					CAS #: 540-84-1			
8.465	8.465	(1.146)	57	3172029	39.0159	39.016	80.00-	120.00	100.00
8.465	8.465	(1.146)	56	1014519			0.00-	30.00	31.98
8.437	8.465	(1.142)	41	787495			0.00-	30.00	24.83

81	Benzene					CAS #: 71-43-2			
8.410	8.437	(0.908)	78	2331131	42.3281	42.328	80.00-	120.00	100.00
8.410	8.437	(0.908)	77	517986			0.00-	30.00	22.22

83	1,2-Dichloroethane					CAS #: 107-06-2			
8.603	8.603	(0.928)	62	1037606	48.4203	48.420	80.00-	120.00	100.00
8.603	8.603	(0.928)	64	329154			0.00-	30.00	31.72

85	Heptane					CAS #: 142-82-5			
8.852	8.852	(0.955)	100	257604	45.0698	45.070	80.00-	120.00	100.00
8.852	8.852	(0.955)	43	1502572			0.00-	30.00	583.29
8.852	8.852	(0.955)	71	802153			0.00-	30.00	311.39

94	Trichloroethene					CAS #: 79-01-6			
9.654	9.682	(1.042)	95	961294	45.9265	45.926	80.00-	120.00	100.00
9.654	9.682	(1.042)	130	1025530			77.04-	137.04	106.68
9.654	9.682	(1.042)	97	602741			33.37-	93.37	62.70

97	1,2-Dichloropropane					CAS #: 78-87-5			
10.179	10.179	(1.098)	63	794079	43.3820	43.382	80.00-	120.00	100.00
10.179	10.179	(1.098)	62	577144			42.77-	102.77	72.68
10.179	10.179	(1.098)	41	492060			31.97-	91.97	61.97

98	1,4-Dioxane					CAS #: 123-91-1			
10.401	10.428	(1.122)	88	503403	45.0900	45.090	80.00-	120.00	100.00
10.401	10.428	(1.122)	58	380284			46.78-	106.78	75.54
10.401	10.428	(1.122)	57	116914			0.00-	30.00	23.22

100	Bromodichloromethane					CAS #: 75-27-4			
10.732	10.732	(1.158)	83	1397122	45.7002	45.700	80.00-	120.00	100.00
10.732	10.732	(1.158)	85	867086			31.71-	91.71	62.06

102	cis-1,3-Dichloropropene					CAS #: 10061-01-5			
11.672	11.672	(1.260)	75	1102335	43.3354	43.335	80.00-	120.00	100.00
11.672	11.672	(1.260)	77	355938			1.66-	61.66	32.29
11.672	11.672	(1.260)	39	572678			21.94-	81.94	51.95

103	4-Methyl-2-pentanone					CAS #: 108-10-1			
12.032	12.032	(1.298)	58	633840	46.0596	46.060	80.00-	120.00	100.00
12.032	12.032	(1.298)	43	1624043			0.00-	30.00	256.22
12.032	12.032	(1.298)	85	254734			0.00-	30.00	40.19

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	ON-COL		FINAL	TARGET RANGE	RATIO	
				RESPONSE	(PPEV)	(PPBV)			
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

105	Toluene			CAS #: 108-88-3					
12.225	12.225	(1.319)	91	2492205	46.4273	46.427	80.00-	120.00	100.00
12.225	12.225	(1.319)	92	1445640			30.07-	90.07	58.01

108	trans-1,3-Dichloropropene			CAS #: 10061-02-6					
12.834	12.834	(0.880)	75	1161878	45.3963	45.396	80.00-	120.00	100.00
12.834	12.834	(0.880)	77	359009			0.77-	60.77	30.90
12.834	12.834	(0.880)	39	566437			19.08-	79.08	48.75

110	1,1,2-Trichloroethane			CAS #: 79-00-5					
13.138	13.138	(0.901)	97	805603	42.9184	42.918	80.00-	120.00	100.00
13.138	13.138	(0.901)	99	490762			31.98-	91.98	60.92
13.138	13.138	(0.901)	83	685452			56.83-	116.83	85.09

112	Tetrachloroethene			CAS #: 127-18-4					
13.193	13.193	(0.905)	166	1183328	46.5783	46.578	80.00-	120.00	100.00
13.165	13.193	(0.903)	129	874196			45.73-	105.73	73.88
13.193	13.193	(0.905)	131	849348			42.92-	102.92	71.78

114	2-Hexanone			CAS #: 591-78-6					
13.580	13.580	(0.932)	58	782102	42.0508	42.051	80.00-	120.00	100.00
13.553	13.580	(0.930)	43	1472714			159.61-	219.61	188.30
13.580	13.580	(0.932)	100	156531			0.00-	30.00	20.01

116	Dibromochloromethane			CAS #: 124-48-1					
13.718	13.718	(0.941)	129	1348045	45.8247	45.825	80.00-	120.00	100.00
13.718	13.718	(0.941)	127	1069227			0.00-	30.00	79.32

117	1,2-Dibromoethane			CAS #: 106-93-4					
13.884	13.884	(0.953)	107	1255262	43.8883	43.888	80.00-	120.00	100.00
13.884	13.884	(0.953)	109	1185514			64.95-	124.95	94.44

126	Chlorobenzene			CAS #: 108-90-7					
14.603	14.603	(1.002)	112	2011743	44.3790	44.379	80.00-	120.00	100.00
14.603	14.603	(1.002)	114	640113			2.04-	62.04	31.82
14.603	14.603	(1.002)	77	1153821			27.23-	87.23	57.35

129	Ethyl Benzene			CAS #: 100-41-4					
14.769	14.769	(1.013)	106	996669	41.5461	41.546	80.00-	120.00	100.00
14.742	14.769	(1.011)	91	3106016			0.00-	30.00	311.64

130	m,p-Xylene			CAS #: 108-38-3					
14.935	14.935	(1.025)	106	1314283	44.9590	44.959	80.00-	120.00	100.00
14.935	14.935	(1.025)	91	2582698			0.00-	30.00	196.51

132	o-Xylene			CAS #: 95-47-6					
15.488	15.488	(1.063)	106	1239020	42.0024	42.002	80.00-	120.00	100.00

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE	ON-COL (PPEV)	FINAL (PPBV)	TARGET RANGE	RATIO	
==	=====	=====	=====	=====	=====	=====	=====	=====	
132 o-Xylene (continued)									
15.488	15.488	(1.063)	91	2515113			175.82- 235.82	202.99	

134 Styrene CAS #: 100-42-5									
15.516	15.516	(1.064)	104	1928614	46.3084	46.308	80.00- 120.00	100.00	
15.516	15.516	(1.064)	78	964771			23.26- 83.26	50.02	

135 Bromoform CAS #: 75-25-2									
15.765	15.765	(1.082)	173	1294443	50.0254	50.025	80.00- 120.00	100.00	
15.765	15.765	(1.082)	171	658261			21.10- 81.10	50.85	

144 1,1,2,2-Tetrachloroethane CAS #: 79-34-5									
16.456	16.456	(1.129)	83	1703512	39.4887	39.489	80.00- 120.00	100.00	
16.456	16.456	(1.129)	85	1055668			31.85- 91.85	61.97	

147 4-Ethyltoluene CAS #: 622-96-8									
16.649	16.649	(1.142)	105	3568588	45.0745	45.074	80.00- 120.00	100.00	
16.649	16.649	(1.142)	120	1097023			0.70- 60.70	30.74	

148 1,3,5-Trimethylbenzene CAS #: 108-67-8									
16.732	16.732	(1.148)	105	3131039	41.1438	41.144	80.00- 120.00	100.00	
16.732	16.732	(1.148)	120	1583281			0.00- 30.00	50.57	

153 1,2,4-Trimethylbenzene CAS #: 95-63-6									
17.147	17.147	(1.176)	105	3435677	44.8515	44.852	80.00- 120.00	100.00	
17.147	17.147	(1.176)	120	1630581			16.88- 76.88	47.46	

156 1,3-Dichlorobenzene CAS #: 541-73-1									
17.451	17.451	(1.197)	146	2346038	46.9814	46.981	80.00- 120.00	100.00	
17.451	17.451	(1.197)	148	1503465			0.00- 30.00	64.09	
17.451	17.451	(1.197)	111	906985			0.00- 30.00	38.66	

157 1,4-Dichlorobenzene CAS #: 106-46-7									
17.562	17.562	(1.205)	146	1881136	39.7498	39.750	80.00- 120.00	100.00	
17.562	17.562	(1.205)	148	1197721			0.00- 30.00	63.67	
17.562	17.562	(1.205)	111	728119			0.00- 30.00	38.71	

158 alpha-Chlorotoluene CAS #: 100-44-7									
17.700	17.700	(1.214)	91	3095601	45.0010	45.001	80.00- 120.00	100.00	
17.700	17.700	(1.214)	126	703472			0.00- 30.00	22.72	

161 1,2-Dichlorobenzene CAS #: 95-50-1									
17.921	17.921	(1.230)	146	1827128	37.9843	37.984	80.00- 120.00	100.00	
17.921	17.921	(1.230)	148	1175305			34.31- 94.31	64.33	
17.894	17.921	(1.228)	111	766950			10.22- 70.22	41.98	

CONCENTRATIONS									
RT	EXP RT	(REL RT)	MASS	RESPONSE		ON-COL	FINAL	TARGET RANGE	RATIO
				(PPEV)	(PPBV)				
==	=====	=====	=====	=====	=====	=====	=====	=====	=====

167	1,2,4-Trichlorobenzene						CAS #: 120-82-1		
19.276	19.276	(1.322)	180	1949568	40.1013	40.101	80.00-	120.00	100.00
19.276	19.276	(1.322)	182	1879622			67.17-	127.17	96.41

168	Hexachlorobutadiene						CAS #: 87-68-3		
19.359	19.359	(1.328)	225	1611250	57.7402	57.740	80.00-	120.00	100.00
19.359	19.359	(1.328)	223	1030031			33.68-	93.68	63.93

145	Propylbenzene						CAS #: 103-65-1		
16.483	16.483	(1.131)	91	4640561	50.2524	50.252	80.00-	120.00	100.00
16.483	16.483	(1.131)	120	1122970			0.00-	30.00	24.20
16.483	16.483	(1.131)	105	171733			0.00-	30.00	3.70

137	Cumene						CAS #: 98-82-8		
15.958	15.958	(1.095)	105	3922826	47.1328	47.133	80.00-	120.00	100.00
15.958	15.958	(1.095)	120	1054441			0.00-	30.00	26.88
15.958	15.958	(1.095)	51	391325			0.00-	30.00	9.98

169	Naphthalene						CAS #: 91-20-3		
19.470	19.470	(1.336)	128	3962023	38.6430	38.643	80.00-	120.00	100.00
19.470	19.470	(1.336)	127	482850			0.00-	30.00	12.19

9	Butane						CAS #: 106-97-8		
2.327	2.355	(0.315)	58	197200	43.0388	43.039	80.00-	120.00	100.00
2.327	2.355	(0.315)	43	1536552			0.00-	30.00	779.18

15	Isopentane						CAS #: 78-78-4		
2.935	2.963	(0.397)	43	1162968	45.6856	45.686	80.00-	120.00	100.00
2.935	2.963	(0.397)	57	748985			0.00-	30.00	64.40
2.935	2.963	(0.397)	72	77284			0.00-	30.00	6.65

95	Methyl Cyclohexane						CAS #: 108-87-2		
9.903	9.903	(1.341)	83	1367065	40.2926	40.292	80.00-	120.00	100.00
9.903	9.903	(1.341)	98	628218			0.00-	30.00	45.95
9.903	9.903	(1.341)	55	1110705			0.00-	30.00	81.25

Report Date: 03-Jul-2007 10:39

Air Toxics Ltd.

INTERNAL STANDARD COMPOUNDS
AREA AND RT SUMMARY

Instrument ID: msd8.i

Calibration Date: 03-JUL-2007

Lab File ID: 8070303.d

Calibration Time: 09:41

Lab Smp Id: LCS-1

Client Smp ID: LCS-1

Analysis Type: VOA

Level: LOW

Quant Type: ISTD

Sample Type: AIR

Operator: jdj

Method File: /chem/msd8.i/8-03jul.b/t14q530b.m

Misc Info: 200ppbv-50ppbv

COMPOUND	STANDARD	AREA LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	345936	207562	484310	278074	-19.62
88 1,4-Difluorobenze	1512218	907331	2117105	1169163	-22.69
125 Chlorobenzene-d5	1120246	672148	1568344	882830	-21.19

COMPOUND	STANDARD	RT LIMIT		SAMPLE	%DIFF
		LOWER	UPPER		
68 Bromochloromethan	7.39	7.06	7.72	7.39	0.00
88 1,4-Difluorobenze	9.27	8.94	9.60	9.27	0.00
125 Chlorobenzene-d5	14.58	14.25	14.91	14.58	0.00

AREA UPPER LIMIT = + 40% of internal standard area.

AREA LOWER LIMIT = - 40% of internal standard area.

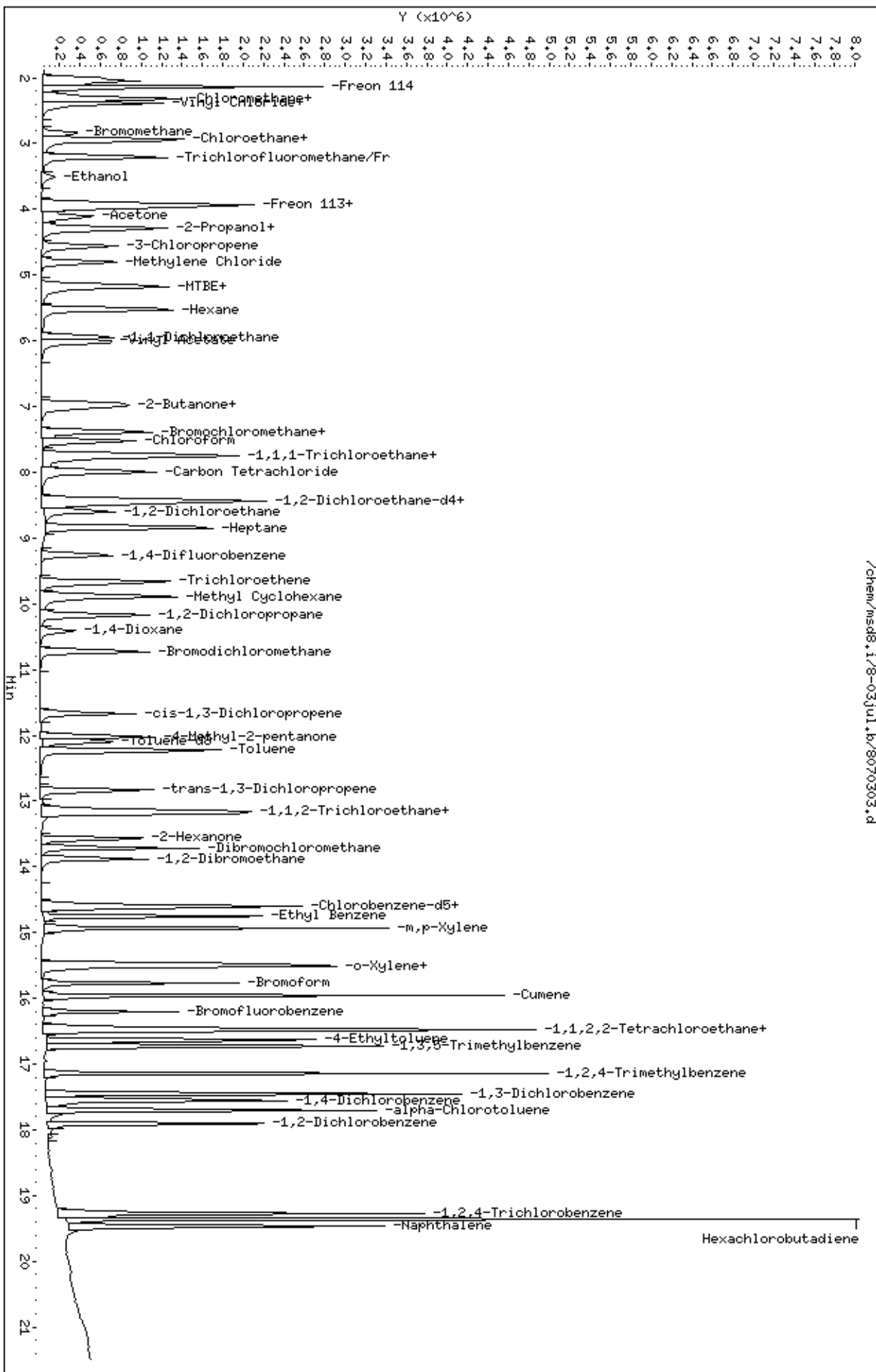
RT UPPER LIMIT = + 0.33 minutes of internal standard RT.

RT LOWER LIMIT = - 0.33 minutes of internal standard RT.

Data File: /chem/msd8.1/8-03jul.b/8070303.d
Date: 03-JUL-2007 10:09
Client ID: LCS-1
Sample Info: 50ml #1443-146

Column phase: RTX-624

Instrument: msd8.1
Operator: jdg
Column diameter: 0.53



@ Air Toxics Ltd.

MSD-8

Logbook #: 1478

m/z	ION ABUNDANCE CRITERIA	% REL. ABUNDANCE
50	15.0 - 40.0% of mass 95	18.46
75	30.0 - 60.0% of mass 95	46.42
95	Base peak, 100.00% relative abundance	100.00
96	5.0 - 9.0% of mass 95	6.78
173	Less than 2.0% of mass 174	(0.21) ¹
174	Greater than 50.0% of mass 95	70.27
175	5.0 - 9.0% of mass 174	(7.58) ¹
176	Greater than 95.0% but less than 101.0% of mass 174	(28.44) ¹
177	5.0 - 9.0% of mass 176	(6.38) ²

BFB Injection Date: 7/3/07
 BFB Injection Time: 0918
 BFB File ID: 8070301
 Tekmar Purge Flow: —
 Vacuum: —
 Verified CCV IS vs ICAL mid-point (-40% D) Q-
 Initials

ISIS Std #:	1487-312	Exp. Date:	4/18/07
BCM	345936		
1,4-DFB	1512218		
CB-d5	1120246		

Verify 176/174 m/z Ratio: $\frac{28.44}{70.27} = 0.4047$

NOAH Cart #: 15/8 File #: 8070205/8070206

Calculation Check:

ppbv of compound = $\frac{\text{Area}_{\text{m/z}}}{\text{Area}_{95}} \times \frac{\text{Conc.}_{95}}{\text{RRF}} = \frac{(1314444)}{(1512218)} \times \frac{(25)}{(0.86549)} = 25.108$

Reported Result 25.108

File ID: 8070302
 Compound: Tol-d8
 Initials: Q-

eg	File #	Sample/Client Name	Can #	Pressure	Amt Loaded	DF	Loader Init.	Date Analyzed	Time Analyzed	Review Init.	Comments
✓	B070301	BER Time Check	043-2461	SD wgt	2 g/L	1.00	Q-	7/3/07	0918	Q-	
✓	02	CV # 1443-137	200ppm	SD ppm	SD w/L	1.00	Q-		0941	Q-	
✓	03	LC8 # 1443-146							1009	Q-	
X	04	CV sp #1487-315							1040	Q-	had load
X	05	System Blank	13673	Horizontal	200 w/L	1.00	Q-		1111	Q-	
X	06	CV sp #1487-315	200ppm	SD ppm	SD w/L	1.00	Q-		1145	Q-	
✓	07	Lab Blank	13673	Horizontal	200 w/L	1.00	Q-		1247	Q-	
✓	08	Cart Car #14, Leg 6							1315	Q-	
✓	09	0706413A-04A	30818	35% 15.5%		2.29	Q-		1424	Q-	

Signature: AG Date: 7/3/07
 Revision 05/2005 Page 357

10	✓	6070310	67020413A-03A	341612	5.5" H ⁺ -15" F ⁻	200ml	2.17	Q ⁻	3/3/07	1506	Q ⁻	
11	X	11	-02A	341116	5.5" H ⁺ -15" F ⁻	2ml	2.47			1541	Q ⁻	W/O 7.5ml
12	X	12	-01A	35635	5.0" H ⁺ -15" F ⁻	1.5ml	3.23			1613	Q ⁻	W/O 4ml
13	X	13	-02A	341116	5.5" H ⁺ -15" F ⁻	7.5ml	6.59			1707	UR	PR @ 10ml
14	X	14	-01A	35235	5.0" H ⁺ -15" F ⁻	4ml	1.21			1737	UR	dil for PT
15	✓	15	-01AA							1807	UR	
16	✓	16	07020413A-01A	4148	7.0" H ⁺ -5" F ⁻	200ml	1.75			1842	UR	
17	✓	17	-02A	3746	7.5" H ⁺ -5" F ⁻		1.39			1924	UR	
18	✓	18	-02AA							2000	UR	
19	✓	19	-03A	34917			1.29			2049	UR	
20	✓	20	-04A	35155	4.4" H ⁺ -4.4" F ⁻		1.00			2131	UR	TB
21	✓	21	07020413A-02A	34116	5.5" H ⁺ -15" F ⁻	10ml	2.44	UR		2216	UR	dil for PT
22	✓	22	07020413A-01A	1855	4.0" H ⁺ -5" F ⁻	200ml	1.55			2307	UR	
23	✓	23	-01AA							2349	Q ⁻	
24	✓	24	-02A	34349	6.5" H ⁺ -5" F ⁻		1.71			2381	Q ⁻	
25	✓	25	-03A	25237	5.0" H ⁺ -15" F ⁻		1.41			2414	Q ⁻	
26	✓	26	-01A	34234	4.5" H ⁺ -5" F ⁻		1.38			2446	Q ⁻	
27	✓	27	07020413A-01A	2-16	6.5" H ⁺ -5" F ⁻		1.71			2538	Q ⁻	
28	✓	28	-02A	34480	6.5" H ⁺ -5" F ⁻		1.46			2621	Q ⁻	
29	✓	29	-04A	12082	4.0" H ⁺ -5" F ⁻		1.55			2703	Q ⁻	
30	✓	30	-03A	33531	1.0" H ⁺ -5" F ⁻		1.39			2746	Q ⁻	
31												
32												

Comments:

[Handwritten signature/initials]

Signature *[Handwritten signature]*

Date *3/4/07*

Report Date: 30-May-2007 13:12

Air Toxics Ltd.

Data file : /var/chem/msd8.i/8-30may.b/8053001.d
 Lab Smp Id: BFB Client Smp ID: BFB
 Inj Date : 30-MAY-2007 13:20
 Operator : db Inst ID: msd8.i
 Smp Info : BFB Tune Check
 Misc Info : 50ng 2uL #843-2981
 Comment :
 Method : /var/chem/msd8.i/8-30may.b/bfb30.m
 Meth Date : 30-May-2007 13:12 Quant Type: ESTD
 Cal Date : Cal File:
 Als bottle: 1 QC Sample: BFB
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: all.sub
 Target Version: 3.50 Sample Matrix: WATER
 Processing Host: eeyore

Concentration Formula: Amt * DF * Uf * Vf * Vi * CpndVariable

Name	Value	Description
DF	1.00000	Dilution Factor
Uf	1.00000	ng unit correction factor
Vf	1.00000	Volumetric correction factor
Vi	2.00000	Injection Volume

Cpnd Variable Local Compound Variable

CONCENTRATIONS

ON-COL FINAL

RT EXP RT DLT RT MASS RESPONSE (ug/L) (ug/L) TARGET RANGE RATIO
 == =====

RT	EXP RT	DLT RT	MASS	RESPONSE	(ug/L)	(ug/L)	TARGET RANGE	RATIO
1	bfb						CAS #: 460-00-4	
3.693	3.748	-0.055	95	2045811			100.00- 100.00	100.00
3.693	3.748	-0.055	50	359998			15.00- 40.00	17.60
3.693	3.748	-0.055	75	919533			30.00- 60.00	44.95
3.693	3.748	-0.055	96	129781			5.00- 9.00	6.34
3.693	3.748	-0.055	173	0			0.00- 2.00	0.00
3.693	3.748	-0.055	174	1680993			50.00- 100.00	82.17
3.693	3.748	-0.055	175	121284			5.00- 9.00	7.22
3.693	3.748	-0.055	176	1646565			95.00- 101.00	97.95
3.693	3.748	-0.055	177	100750			5.00- 9.00	6.12

Date : 30-MAY-2007 13:20

Client ID: BFB

Instrument: msd8.i

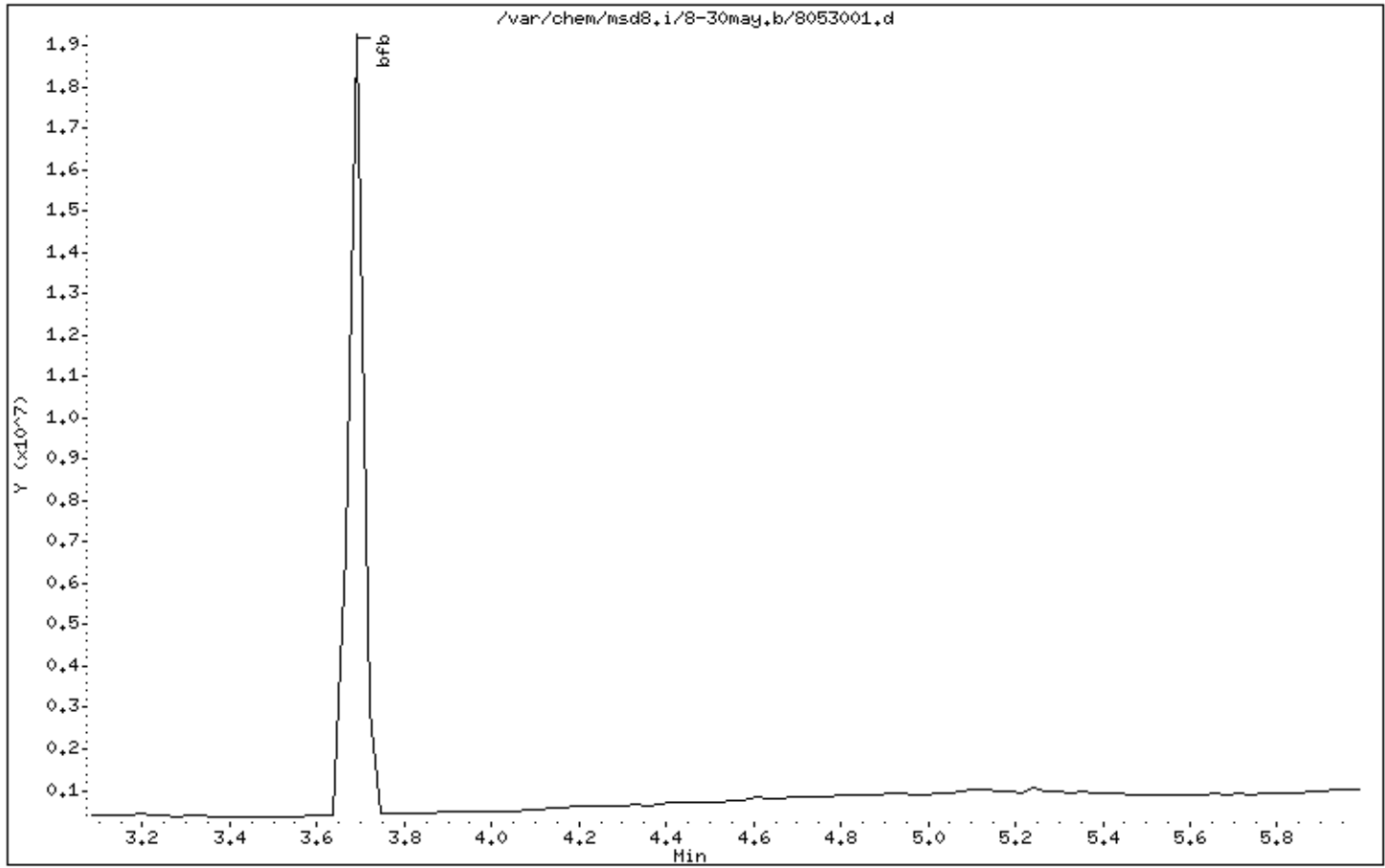
Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: db

Column phase:

Column diameter: 0.53



Date : 30-MAY-2007 13:20

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

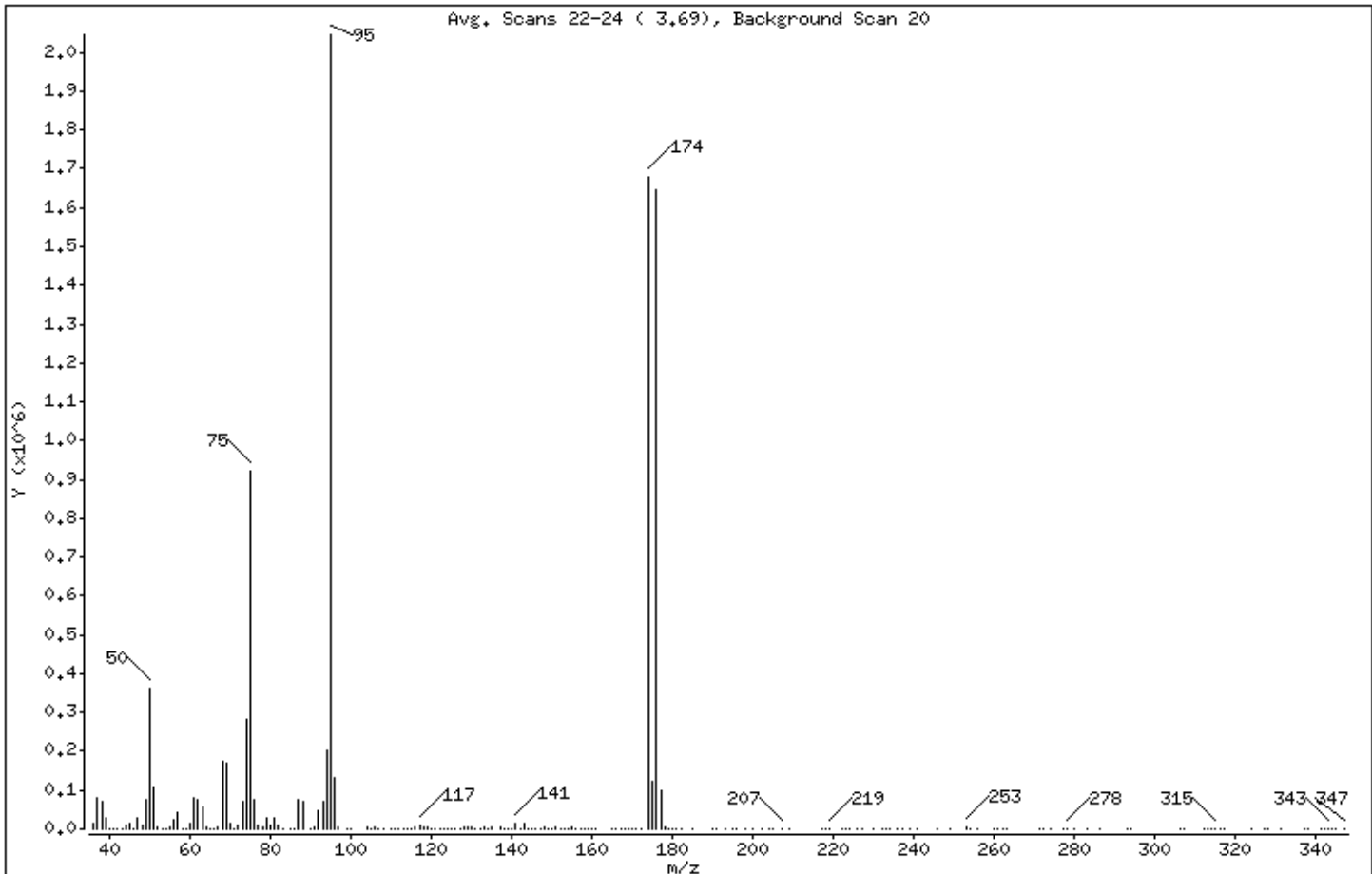
Volume Injected (uL): 2.0

Operator: db

Column phase:

Column diameter: 0.53

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95	Base Peak, 100% relative abundance	100.00
50	15.00 - 40.00% of mass 95	17.60
75	30.00 - 60.00% of mass 95	44.95
96	5.00 - 9.00% of mass 95	6.34
173	Less than 2.00% of mass 174	0.00 (0.00)
174	50.00 - 100.00% of mass 95	82.17
175	5.00 - 9.00% of mass 174	5.93 (7.22)
176	95.00 - 101.00% of mass 174	80.48 (97.95)
177	5.00 - 9.00% of mass 176	4.92 (6.12)

Date : 30-MAY-2007 13:20

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: db

Column phase:

Column diameter: 0.53

Data File: 8053001.d

Spectrum: Avg. Scans 22-24 (3.69), Background Scan 20

Location of Maximum: 95.00

Number of points: 203

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	14576	88.00	69600	146.00	2289	224.00	96
37.00	81912	90.00	71	147.00	735	226.00	106
38.00	72104	91.00	5332	148.00	3537	227.00	79
39.00	27744	92.00	45488	149.00	517	230.00	71
40.00	666	93.00	72336	150.00	1526	232.00	7
41.00	282	94.00	200896	151.00	3455	233.00	69
42.00	266	95.00	2045440	152.00	1099	234.00	53
43.00	1037	96.00	129776	153.00	1348	236.00	162
44.00	8880	97.00	3615	154.00	793	237.00	106
45.00	14430	99.00	151	155.00	3949	239.00	128
46.00	773	100.00	254	156.00	1096	241.00	12
47.00	25872	104.00	5055	157.00	2094	246.00	153
48.00	9871	105.00	596	158.00	402	249.00	187
49.00	74368	106.00	4503	159.00	2163	253.00	2495
50.00	359936	107.00	988	160.00	268	254.00	409
51.00	108176	108.00	555	161.00	1464	256.00	228
52.00	4377	110.00	383	165.00	310	260.00	2065
53.00	38	111.00	999	166.00	280	261.00	288
54.00	179	112.00	318	167.00	31	262.00	141
55.00	3691	113.00	871	168.00	406	263.00	49
56.00	23056	114.00	168	169.00	116	271.00	132
57.00	42176	115.00	1417	170.00	525	272.00	213
58.00	1681	116.00	4417	171.00	529	274.00	165
59.00	870	117.00	7444	172.00	147	277.00	227
60.00	14254	118.00	4536	174.00	1680896	278.00	250
61.00	78008	119.00	5766	175.00	121280	280.00	96
62.00	77280	120.00	141	176.00	1646080	283.00	9
63.00	54928	121.00	37	177.00	100744	286.00	82
64.00	4760	122.00	334	178.00	2548	293.00	62
65.00	299	123.00	262	179.00	303	294.00	193
66.00	139	124.00	1026	180.00	158	306.00	23
67.00	4680	125.00	80	181.00	28	307.00	84
68.00	174208	126.00	853	182.00	414	312.00	77
69.00	170368	127.00	1542	185.00	250	313.00	20
70.00	12581	128.00	6593	190.00	92	314.00	35

Date : 30-MAY-2007 13:20

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: db

Column phase:

Column diameter: 0.53

Data File: 8053001.d
Spectrum: Avg. Scans 22-24 (3.69), Background Scan 20
Location of Maximum: 95.00
Number of points: 203

m/z	Y	m/z	Y	m/z	Y	m/z	Y
71.00	479	129.00	2422	191.00	1655	315.00	123
72.00	8178	130.00	5089	193.00	1111	316.00	69
73.00	70936	131.00	2310	195.00	988	317.00	75
74.00	282304	132.00	264	196.00	87	324.00	162
75.00	919488	133.00	5559	198.00	244	327.00	112
76.00	77456	134.00	717	200.00	181	328.00	238
77.00	9607	135.00	3857	202.00	196	331.00	341
78.00	6315	137.00	2383	204.00	252	337.00	8
79.00	28224	138.00	384	205.00	48	338.00	80
80.00	10980	139.00	23	207.00	1232	341.00	15
81.00	30280	140.00	1176	209.00	15	342.00	184
82.00	8002	141.00	14199	217.00	163	343.00	1020
83.00	870	142.00	1290	218.00	108	344.00	224
85.00	271	143.00	13753	219.00	292	345.00	257
86.00	2288	144.00	540	222.00	218	347.00	112
87.00	76888	145.00	517	223.00	143		

Report Date: 31-May-2007 10:40

Air Toxics Ltd.

Data file : /var/chem/msd8.i/8-31may.b/8053101.d
 Lab Smp Id: BFB Client Smp ID: BFB
 Inj Date : 31-MAY-2007 10:48
 Operator : JG Inst ID: msd8.i
 Smp Info : BFB Tune Check
 Misc Info : 50ng 2uL #843-2981
 Comment :
 Method : /var/chem/msd8.i/8-31may.b/bfb30.m
 Meth Date : 31-May-2007 10:40 Quant Type: ESTD
 Cal Date : Cal File:
 Als bottle: 1 QC Sample: BFB
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: all.sub
 Target Version: 3.50 Sample Matrix: WATER
 Processing Host: eeyore

Concentration Formula: Amt * DF * Uf * Vf * Vi * CpndVariable

Name	Value	Description
DF	1.00000	Dilution Factor
Uf	1.00000	ng unit correction factor
Vf	1.00000	Volumetric correction factor
Vi	2.00000	Injection Volume

Cpnd Variable Local Compound Variable

CONCENTRATIONS

ON-COL FINAL

RT	EXP RT	DLT RT	MASS	RESPONSE	(ug/L)	(ug/L)	TARGET RANGE	RATIO
==	=====	=====	=====	=====	=====	=====	=====	=====

CAS #: 460-00-4

1 bfb								
3.665	3.748	-0.083	95	2098550			100.00- 100.00	100.00
3.665	3.748	-0.083	50	381786			15.00- 40.00	18.19
3.665	3.748	-0.083	75	968533			30.00- 60.00	46.15
3.665	3.748	-0.083	96	132439			5.00- 9.00	6.31
3.665	3.748	-0.083	173	0			0.00- 2.00	0.00
3.665	3.748	-0.083	174	1305650			50.00- 100.00	62.22
3.665	3.748	-0.083	175	94421			5.00- 9.00	7.23
3.665	3.748	-0.083	176	1269961			95.00- 101.00	97.27
3.665	3.748	-0.083	177	79443			5.00- 9.00	6.26

Date : 31-MAY-2007 10:48

Client ID: BFB

Instrument: msd8.i

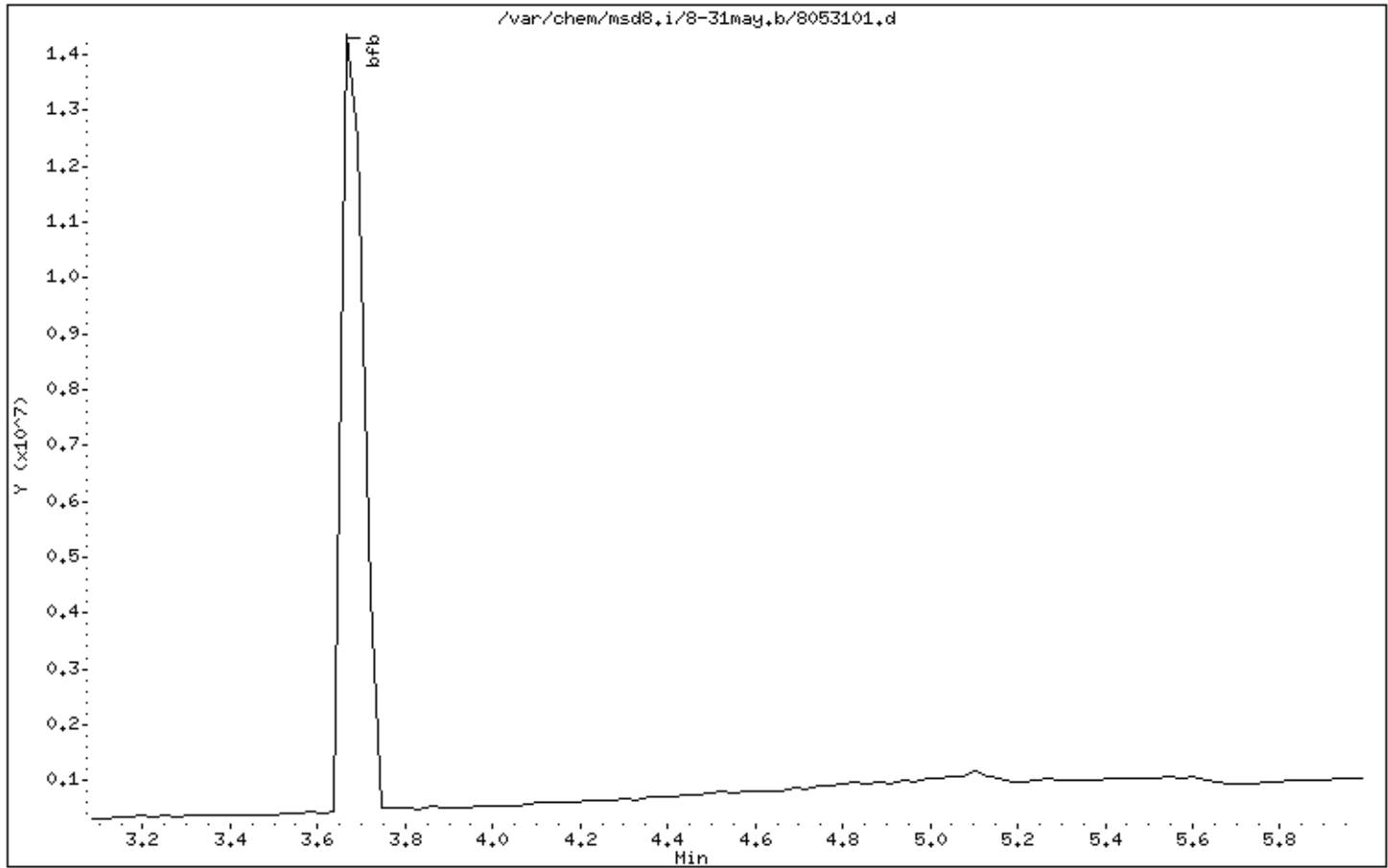
Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53



Date : 31-MAY-2007 10:48

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

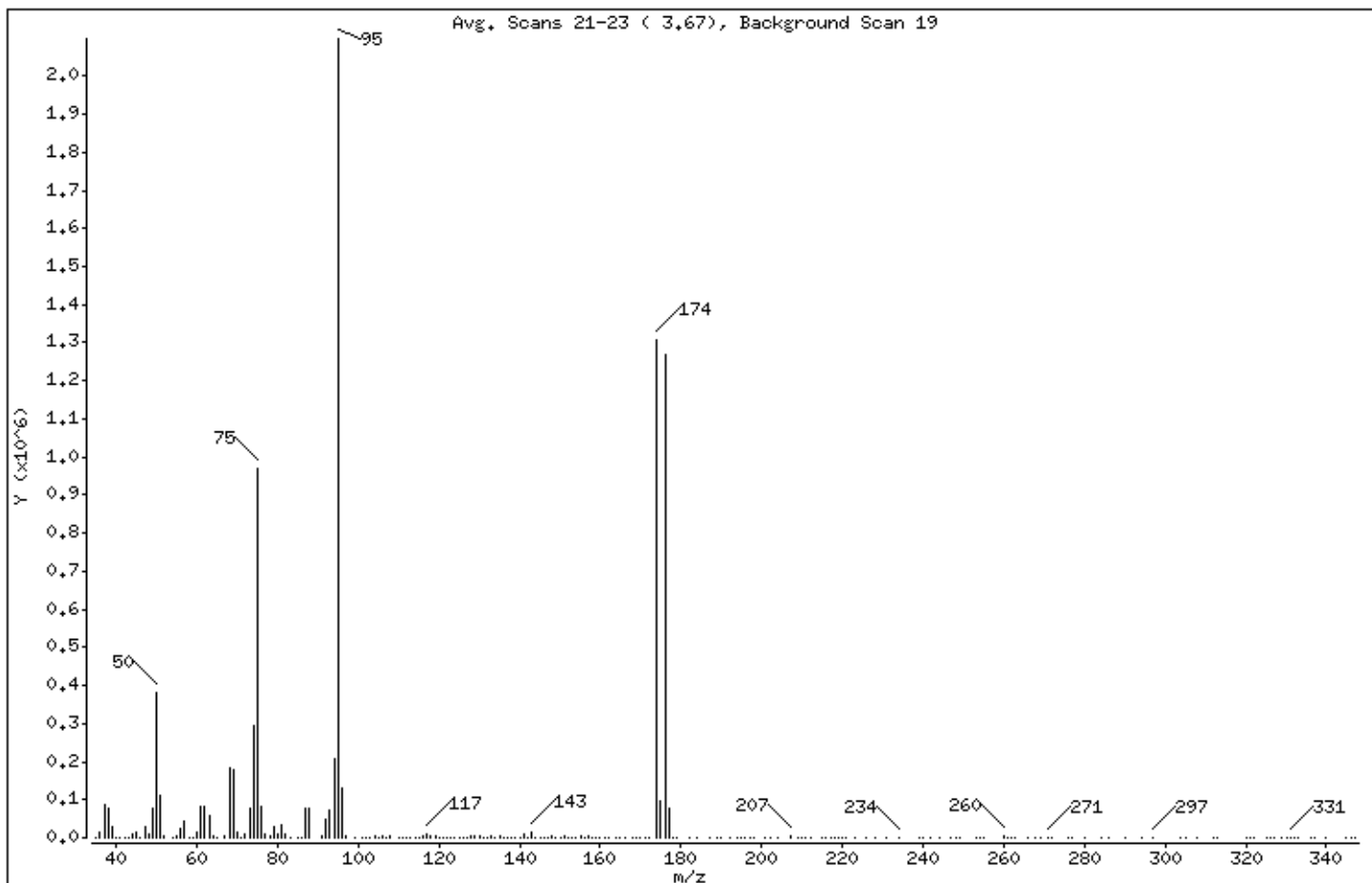
Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95	Base Peak, 100% relative abundance	100.00
50	15.00 - 40.00% of mass 95	18.19
75	30.00 - 60.00% of mass 95	46.15
96	5.00 - 9.00% of mass 95	6.31
173	Less than 2.00% of mass 174	0.00 (0.00)
174	50.00 - 100.00% of mass 95	62.22
175	5.00 - 9.00% of mass 174	4.50 (7.23)
176	95.00 - 101.00% of mass 174	60.52 (97.27)
177	5.00 - 9.00% of mass 176	3.79 (6.26)

Date : 31-MAY-2007 10:48

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

Data File: 8053101.d

Spectrum: Avg. Scans 21-23 (3.67), Background Scan 19

Location of Maximum: 95.00

Number of points: 214

m/z	Y	m/z	Y	m/z	Y	m/z	Y
35.00	137	94.00	205184	151.00	3262	228.00	71
36.00	14156	95.00	2098176	152.00	1052	231.00	253
37.00	84784	96.00	132416	153.00	1057	234.00	294
38.00	75080	97.00	3341	154.00	736	239.00	62
39.00	28376	99.00	48	155.00	3200	240.00	97
40.00	1047	101.00	315	156.00	909	242.00	73
41.00	106	102.00	203	157.00	2816	244.00	149
42.00	55	103.00	657	158.00	559	247.00	572
43.00	211	104.00	5350	159.00	1429	248.00	254
44.00	7388	105.00	2049	160.00	60	249.00	409
45.00	14858	106.00	5376	161.00	1271	253.00	1015
46.00	633	107.00	1173	162.00	125	254.00	514
47.00	26592	108.00	2447	164.00	421	255.00	420
48.00	10731	110.00	632	165.00	504	260.00	2797
49.00	76568	111.00	502	166.00	207	261.00	644
50.00	381760	112.00	639	168.00	29	262.00	211
51.00	113200	113.00	462	169.00	293	263.00	573
52.00	4745	114.00	191	170.00	825	266.00	105
54.00	329	115.00	1370	171.00	202	268.00	539
55.00	4012	116.00	4406	172.00	1168	269.00	2080
56.00	23784	117.00	7635	174.00	1305600	271.00	417
57.00	44856	118.00	4301	175.00	94416	272.00	226
58.00	1605	119.00	5713	176.00	1269760	276.00	73
59.00	94	120.00	206	177.00	79440	277.00	70
60.00	15955	121.00	85	178.00	2019	280.00	67
61.00	80456	122.00	442	179.00	232	284.00	33
62.00	80808	123.00	276	182.00	265	286.00	285
63.00	59272	124.00	626	184.00	73	290.00	69
64.00	4612	125.00	654	187.00	309	294.00	279
65.00	347	126.00	665	189.00	56	297.00	97
67.00	4333	127.00	383	190.00	468	304.00	67
68.00	183360	128.00	6237	192.00	470	305.00	75
69.00	178944	129.00	2514	194.00	219	308.00	6
70.00	12538	130.00	5415	195.00	187	312.00	97
71.00	154	131.00	1580	196.00	51	313.00	93

Date : 31-MAY-2007 10:48

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

Data File: 8053101.d

Spectrum: Avg. Scans 21-23 (3.67), Background Scan 19

Location of Maximum: 95.00

Number of points: 214

m/z	Y	m/z	Y	m/z	Y	m/z	Y
72,00	9221	132,00	642	197,00	155	320,00	79
73,00	77056	133,00	4651	198,00	68	321,00	66
74,00	293888	134,00	94	201,00	293	322,00	31
75,00	968512	135,00	3077	202,00	2	325,00	87
76,00	80984	136,00	556	204,00	196	326,00	336
77,00	10372	137,00	1762	207,00	4437	327,00	374
78,00	6903	138,00	535	209,00	907	329,00	379
79,00	30680	139,00	313	210,00	269	330,00	28
80,00	11058	140,00	959	211,00	98	331,00	523
81,00	32776	141,00	12035	212,00	77	332,00	108
82,00	7583	142,00	1761	215,00	383	333,00	61
83,00	817	143,00	13357	216,00	126	336,00	57
85,00	193	144,00	630	217,00	405	337,00	293
86,00	1722	145,00	1013	218,00	98	340,00	72
87,00	76424	146,00	1173	219,00	148	345,00	27
88,00	75128	147,00	1632	220,00	54	346,00	86
91,00	5521	148,00	3013	221,00	230	347,00	328
92,00	47968	149,00	1108	223,00	29		
93,00	70904	150,00	1355	226,00	7		

Report Date: 07-Jun-2007 09:36

Air Toxics Ltd.

Data file : /var/chem/msd8.i/8-07jun.b/8060701.d
 Lab Smp Id: BFB Client Smp ID: BFB
 Inj Date : 07-JUN-2007 09:44
 Operator : JG Inst ID: msd8.i
 Smp Info : BFB Tune Check
 Misc Info : 50ng 2uL #843-2981
 Comment :
 Method : /var/chem/msd8.i/8-07jun.b/bfb30.m
 Meth Date : 07-Jun-2007 09:36 Quant Type: ESTD
 Cal Date : Cal File:
 Als bottle: 1 QC Sample: BFB
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: all.sub
 Target Version: 3.50 Sample Matrix: WATER
 Processing Host: eeyore

Concentration Formula: Amt * DF * Uf * Vf * Vi * CpndVariable

Name	Value	Description
DF	1.00000	Dilution Factor
Uf	1.00000	ng unit correction factor
Vf	1.00000	Volumetric correction factor
Vi	2.00000	Injection Volume

Cpnd Variable Local Compound Variable

CONCENTRATIONS

ON-COL FINAL

RT EXP RT DLT RT MASS RESPONSE (ug/L) (ug/L) TARGET RANGE RATIO
 == =====

RT	EXP RT	DLT RT	MASS	RESPONSE	(ug/L)	(ug/L)	TARGET RANGE	RATIO
1	bfb						CAS #: 460-00-4	
3.693	3.748	-0.055	95	1547009			100.00- 100.00	100.00
3.693	3.748	-0.055	50	260935			15.00- 40.00	16.87
3.693	3.748	-0.055	75	686126			30.00- 60.00	44.35
3.693	3.748	-0.055	96	100570			5.00- 9.00	6.50
3.693	3.748	-0.055	173	0			0.00- 2.00	0.00
3.693	3.748	-0.055	174	1270075			50.00- 100.00	82.10
3.693	3.748	-0.055	175	91158			5.00- 9.00	7.18
3.693	3.748	-0.055	176	1238372			95.00- 101.00	97.50
3.693	3.748	-0.055	177	79712			5.00- 9.00	6.44

Data File: /var/chem/msd8.i/8-07jun,b/8060701.d

Page 1

Date : 07-JUN-2007 09:44

Client ID: BFB

Instrument: msd8.i

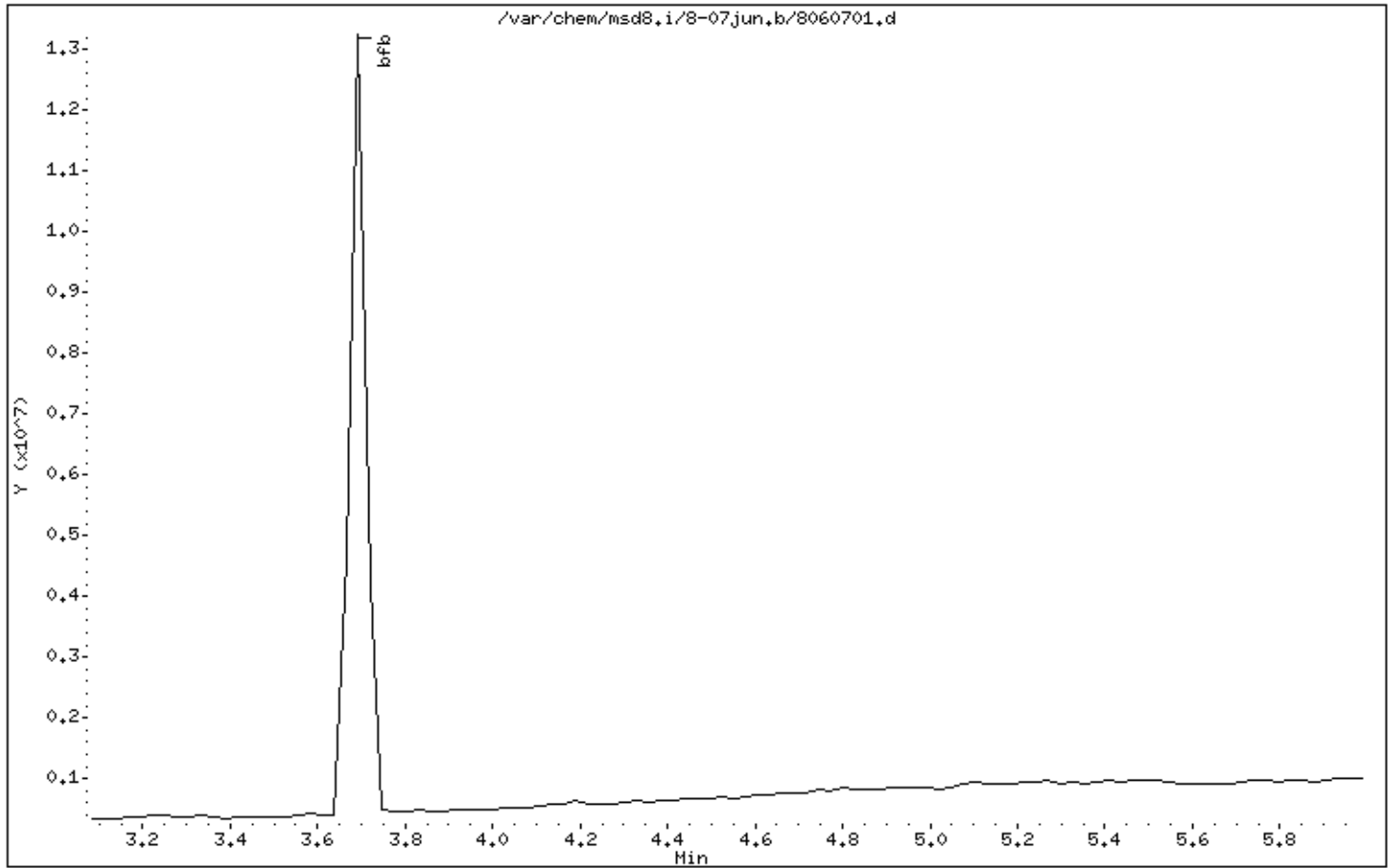
Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53



Date : 07-JUN-2007 09:44

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

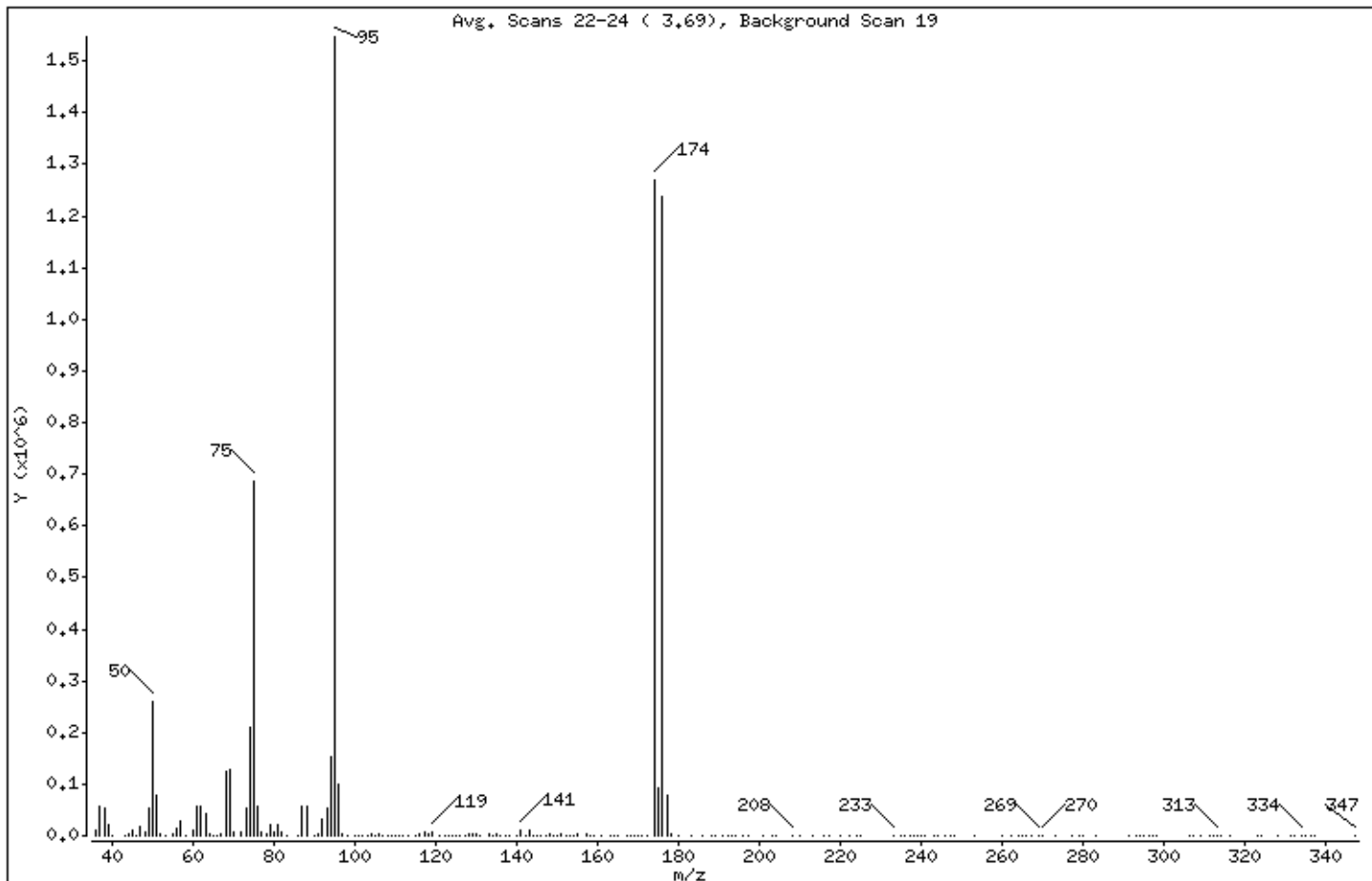
Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95	Base Peak, 100% relative abundance	100.00
50	15.00 - 40.00% of mass 95	16.87
75	30.00 - 60.00% of mass 95	44.35
96	5.00 - 9.00% of mass 95	6.50
173	Less than 2.00% of mass 174	0.00 (0.00)
174	50.00 - 100.00% of mass 95	82.10
175	5.00 - 9.00% of mass 174	5.89 (7.18)
176	95.00 - 101.00% of mass 174	80.05 (97.50)
177	5.00 - 9.00% of mass 176	5.15 (6.44)

Date : 07-JUN-2007 09:44

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

Data File: 8060701.d

Spectrum: Avg. Scans 22-24 (3.69), Background Scan 19

Location of Maximum: 95.00

Number of points: 200

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	9970	95.00	1546752	151.00	1962	239.00	118
37.00	57896	96.00	100568	152.00	814	240.00	95
38.00	52136	97.00	2753	153.00	984	241.00	353
39.00	19648	98.00	161	154.00	1010	243.00	71
40.00	588	100.00	167	155.00	3800	244.00	176
43.00	197	101.00	285	157.00	2246	246.00	120
44.00	4251	102.00	16	158.00	327	247.00	6
45.00	9185	103.00	22	159.00	1499	248.00	11
46.00	342	104.00	4146	161.00	1320	253.00	36
47.00	19488	105.00	117	163.00	412	260.00	741
48.00	7926	106.00	4264	164.00	210	262.00	130
49.00	52552	107.00	1026	165.00	348	264.00	100
50.00	260928	108.00	312	167.00	140	265.00	313
51.00	78664	109.00	104	168.00	485	266.00	130
52.00	3036	110.00	448	169.00	66	267.00	225
53.00	118	111.00	533	170.00	415	269.00	1562
55.00	2142	112.00	413	171.00	196	270.00	1087
56.00	15891	113.00	724	172.00	123	273.00	281
57.00	28648	115.00	1346	174.00	1269760	277.00	76
58.00	1367	116.00	3207	175.00	91152	279.00	262
60.00	10831	117.00	5498	176.00	1238016	280.00	71
61.00	56968	118.00	4049	177.00	79712	283.00	90
62.00	56216	119.00	5853	178.00	2338	291.00	69
63.00	41432	121.00	1635	180.00	88	293.00	255
64.00	3731	122.00	373	183.00	322	294.00	81
65.00	363	123.00	369	186.00	156	295.00	90
66.00	70	124.00	635	188.00	13	296.00	257
67.00	2845	125.00	463	189.00	96	297.00	347
68.00	125112	126.00	226	191.00	373	298.00	75
69.00	129024	127.00	748	192.00	133	306.00	333
70.00	8669	128.00	3748	193.00	163	307.00	133
72.00	6050	129.00	2217	194.00	325	309.00	140
73.00	53624	130.00	4252	196.00	3	311.00	123
74.00	208320	131.00	1446	197.00	94	312.00	69
75.00	686080	133.00	2990	201.00	184	313.00	385

Date : 07-JUN-2007 09:44

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

Data File: 8060701.d

Spectrum: Avg. Scans 22-24 (3.69), Background Scan 19

Location of Maximum: 95.00

Number of points: 200

m/z	Y	m/z	Y	m/z	Y	m/z	Y
76.00	57288	134.00	774	203.00	75	314.00	315
77.00	7929	135.00	3276	204.00	97	316.00	128
78.00	5211	136.00	534	208.00	463	323.00	154
79.00	20992	137.00	1598	210.00	12	324.00	52
80.00	8519	138.00	533	213.00	69	328.00	104
81.00	22688	140.00	855	216.00	79	331.00	184
82.00	6628	141.00	10503	217.00	7	332.00	155
83.00	282	142.00	1289	220.00	40	334.00	738
86.00	1293	143.00	10201	222.00	82	335.00	292
87.00	57576	144.00	550	224.00	74	336.00	256
88.00	55552	145.00	522	225.00	34	337.00	184
90.00	170	146.00	1223	233.00	391	347.00	72
91.00	4158	147.00	355	235.00	251		
92.00	33184	148.00	2820	236.00	228		
93.00	54200	149.00	926	237.00	151		
94.00	153536	150.00	1082	238.00	199		

Report Date: 03-Jul-2007 09:10

Air Toxics Ltd.

Data file : /var/chem/msd8.i/8-03jul.b/8070301.d
 Lab Smp Id: BFB Client Smp ID: BFB
 Inj Date : 03-JUL-2007 09:18
 Operator : JG Inst ID: msd8.i
 Smp Info : BFB Tune Check
 Misc Info : 50ng 2uL #843-2981
 Comment :
 Method : /var/chem/msd8.i/8-03jul.b/bfb30.m
 Meth Date : 03-Jul-2007 09:10 Quant Type: ESTD
 Cal Date : Cal File:
 Als bottle: 1 QC Sample: BFB
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: all.sub
 Target Version: 3.50 Sample Matrix: WATER
 Processing Host: eeyore

Concentration Formula: Amt * DF * Uf * Vf * Vi * CpndVariable

Name	Value	Description
DF	1.00000	Dilution Factor
Uf	1.00000	ng unit correction factor
Vf	1.00000	Volumetric correction factor
Vi	2.00000	Injection Volume

Cpnd Variable Local Compound Variable

CONCENTRATIONS

ON-COL FINAL

RT EXP RT DLT RT MASS RESPONSE (ug/L) (ug/L) TARGET RANGE RATIO
 == =====

RT	EXP RT	DLT RT	MASS	RESPONSE	(ug/L)	(ug/L)	TARGET RANGE	RATIO
1	bfb						CAS #: 460-00-4	
3.693	3.748	-0.055	95	1231465			100.00- 100.00	100.00
3.693	3.748	-0.055	50	227274			15.00- 40.00	18.46
3.693	3.748	-0.055	75	577782			30.00- 60.00	46.92
3.693	3.748	-0.055	96	83517			5.00- 9.00	6.78
3.693	3.748	-0.055	173	2005			0.00- 2.00	0.21
3.693	3.748	-0.055	174	939285			50.00- 100.00	76.27
3.693	3.748	-0.055	175	71213			5.00- 9.00	7.58
3.693	3.748	-0.055	176	924608			95.00- 101.00	98.44
3.693	3.748	-0.055	177	58944			5.00- 9.00	6.38

Data File: /var/chem/msd8.i/8-03jul.b/8070301.d

Page 1

Date : 03-JUL-2007 09:18

Client ID: BFB

Instrument: msd8.i

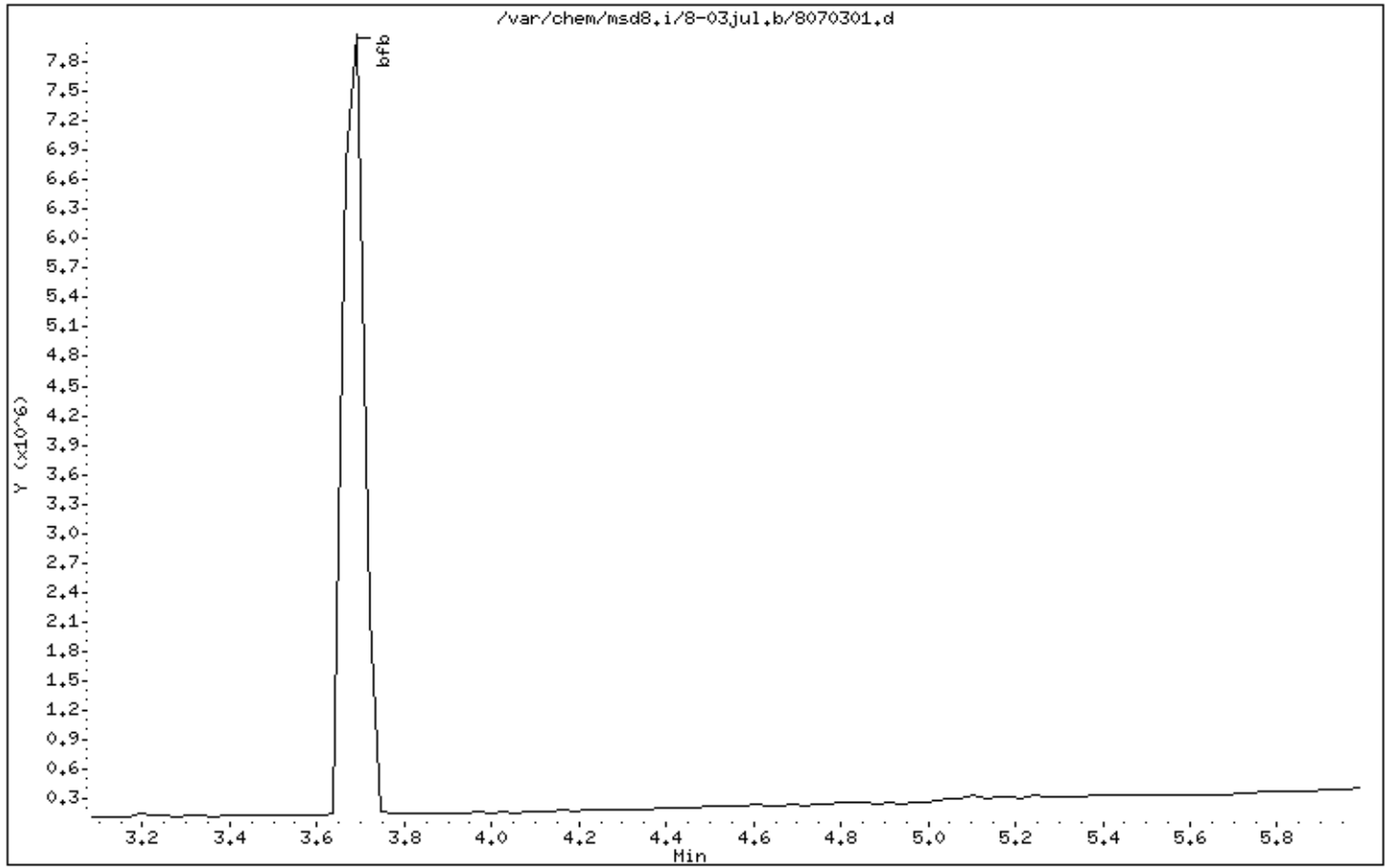
Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53



Date : 03-JUL-2007 09:18

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

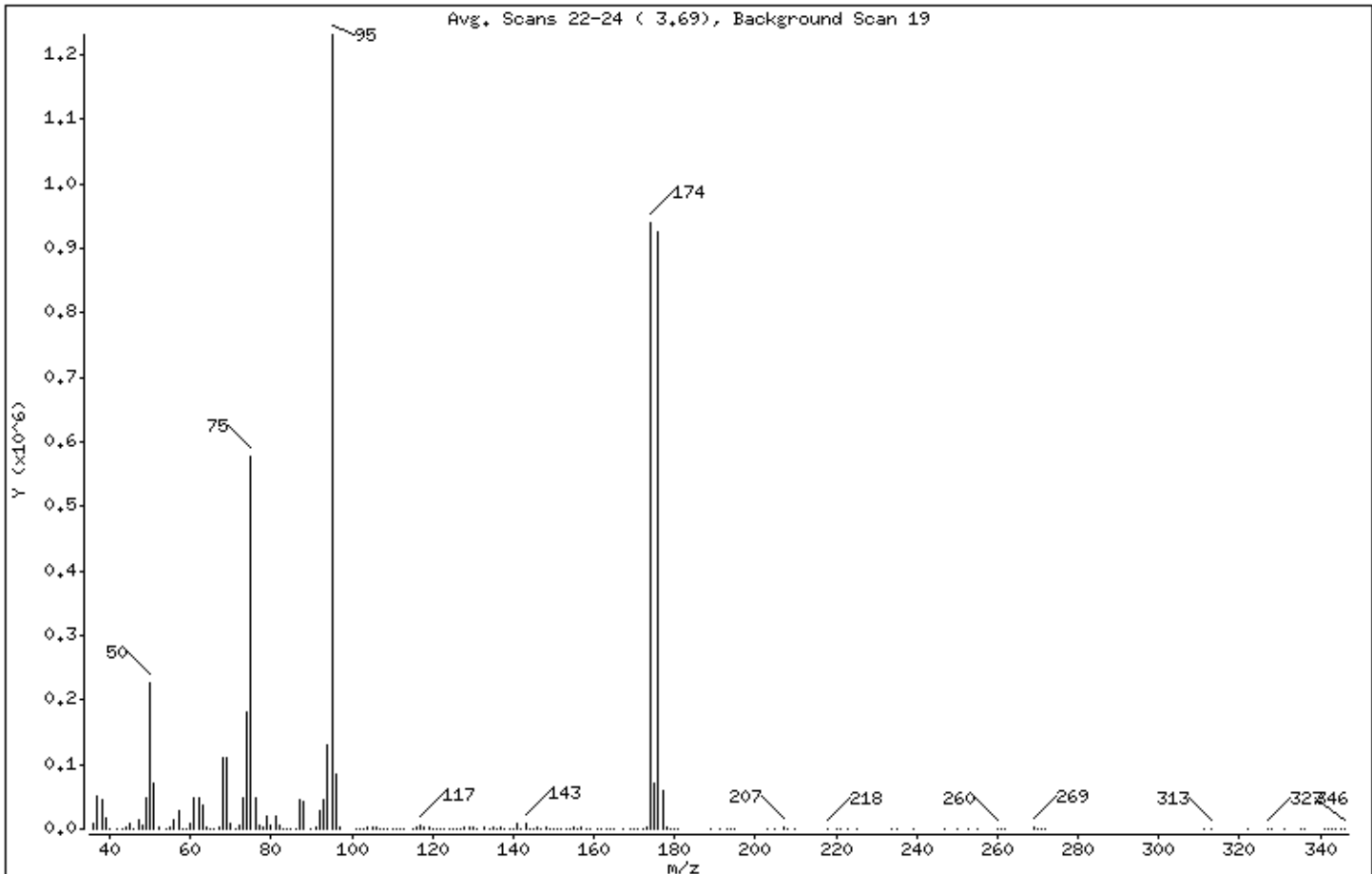
Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95	Base Peak, 100% relative abundance	100.00
50	15.00 - 40.00% of mass 95	18.46
75	30.00 - 60.00% of mass 95	46.92
96	5.00 - 9.00% of mass 95	6.78
173	Less than 2.00% of mass 174	0.16 (0.21)
174	50.00 - 100.00% of mass 95	76.27
175	5.00 - 9.00% of mass 174	5.78 (7.58)
176	95.00 - 101.00% of mass 174	75.08 (98.44)
177	5.00 - 9.00% of mass 176	4.79 (6.38)

Date : 03-JUL-2007 09:18

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

Data File: 8070301.d

Spectrum: Avg. Scans 22-24 (3.69), Background Scan 19

Location of Maximum: 95.00

Number of points: 178

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	8978	83.00	669	134.00	255	189.00	77
37.00	51568	84.00	97	135.00	1976	191.00	24
38.00	45408	85.00	107	136.00	423	193.00	48
39.00	17304	86.00	1104	137.00	1532	194.00	170
40.00	543	87.00	44176	138.00	250	195.00	101
42.00	57	88.00	42120	139.00	449	203.00	23
43.00	753	90.00	258	140.00	756	205.00	37
44.00	4175	91.00	3022	141.00	9194	207.00	1539
45.00	9489	92.00	29384	142.00	1138	208.00	253
46.00	570	93.00	45136	143.00	9618	210.00	289
47.00	15388	94.00	128904	144.00	752	218.00	312
48.00	6631	95.00	1231360	145.00	650	220.00	77
49.00	47312	96.00	83512	146.00	1667	221.00	207
50.00	227264	97.00	2210	147.00	1051	223.00	40
51.00	69808	101.00	240	148.00	2483	225.00	69
52.00	2845	102.00	370	149.00	941	234.00	156
54.00	42	103.00	598	150.00	1090	235.00	73
55.00	2084	104.00	3395	151.00	1058	239.00	71
56.00	14859	105.00	1482	152.00	703	247.00	14
57.00	27488	106.00	3623	153.00	882	250.00	117
58.00	860	107.00	965	154.00	829	253.00	666
59.00	448	108.00	156	155.00	2836	255.00	8
60.00	9850	109.00	376	156.00	412	260.00	805
61.00	48800	110.00	519	157.00	2040	261.00	240
62.00	49192	111.00	761	158.00	509	262.00	542
63.00	36760	112.00	658	159.00	1108	269.00	1542
64.00	3372	113.00	739	161.00	1320	270.00	135
65.00	373	115.00	820	162.00	230	271.00	374
66.00	135	116.00	2822	163.00	42	272.00	254
67.00	2780	117.00	5115	164.00	198	311.00	72
68.00	109816	118.00	3125	165.00	26	313.00	166
69.00	110408	119.00	4204	167.00	161	322.00	303
70.00	7576	120.00	296	169.00	256	327.00	527
71.00	187	121.00	64	170.00	353	328.00	77
72.00	5003	122.00	120	171.00	293	331.00	222

Date : 03-JUL-2007 09:18

Client ID: BFB

Instrument: msd8.i

Sample Info: BFB Tune Check

Volume Injected (uL): 2.0

Operator: JG

Column phase:

Column diameter: 0.53

Data File: 8070301.d

Spectrum: Avg. Scans 22-24 (3.69), Background Scan 19

Location of Maximum: 95.00

Number of points: 178

m/z	Y	m/z	Y	m/z	Y	m/z	Y
73.00	46992	123.00	271	172.00	818	335.00	69
74.00	180736	124.00	461	173.00	2005	336.00	213
75.00	577728	125.00	203	174.00	939264	341.00	204
76.00	47608	126.00	696	175.00	71208	342.00	358
77.00	6102	127.00	338	176.00	924608	343.00	61
78.00	3341	128.00	3517	177.00	58944	344.00	402
79.00	18944	129.00	1951	178.00	1839	345.00	76
80.00	7009	130.00	3812	179.00	283	346.00	254
81.00	20144	131.00	1310	180.00	157		
82.00	5270	133.00	1638	181.00	23		

Shipping/ Receiving Documents



AN ENVIRONMENTAL ANALYTICAL LABORATORY

**180 Blue Ravine Road, Suite B
Folsom, CA 95630**

**Phone (916) 985-1000 FAX (916) 985-1020
Hours 8:00 A.M. to 6:00 P.M. Pacific**

COMPANY: _____ GEI Consultants, Inc.
ATTENTION: _____ Ms. Sarah Aldridge
FAX #: _____ 860-368-5307
FROM: _____ Sample Receiving
Workorder #: _____ 0706440
of pages (Including Cover): _____ 1

7/10/2007

Thank you for selecting Air Toxics Ltd. We have received your samples and have found discrepancies. In order to expedite analysis and reporting, please review the attached information for accuracy. Corrections can be faxed to **Alicia Sullivan at 916-985-1020**. ATL will proceed with the analysis as specified on the Chain of Custody and Sample Login page.

The following discrepancy has been observed:

Sample identifications on the Chain of Custody were not unique for samples AMS 5 DW (4148) and AMS 5 DW (3746). The canister numbers were added to each identification to ensure uniqueness. ATL will proceed with the analysis unless otherwise notified.

Your prompt response is appreciated.



CHAIN-OF-CUSTODY RECORD

Sample Transportation Notice
 Relinquishing signature on this document indicates that sample is being shipped in compliance with all applicable local, State, Federal, National, and International laws, regulations and ordinances of every kind. Air Toxics Limited assumes no liability with respect to the collection, handling or shipping of these samples. Relinquishing signature also indicates agreement to hold harmless, defend, and indemnify Air Toxics Limited against any claim, demand, or action, of any kind, related to the collection, handling, or shipping of samples. D.O.T. Hotline (800) 487-4922

180 BLUE RAVINE ROAD, SUITE B
 FOLSOM, CA 95630-4719
 (916) 985-1000 FAX (916) 985-1020

Project Manager: BRIAN McCarthy

Collected by: (Print and Sign) M. FOSTER

Company: PEI Consultants, INC Email: _____

Address: 455 WINDING BROOK CRY CROSBURY State CT Zip 06033

Phone: _____ Fax: _____

Project Info:
 P.O. # _____
 Project # 061140-8-1703
 Project Name PEI STORE OUI

Turn Around Time:
 Normal
 Rush
 Pressurized by: BF
 Date: 6/26/07
 Pressurization Gas: N₂ He

Lab I.D.	Field Sample I.D. (Location)	Can #	Date of Collection	Time of Collection	Analyses Requested	Canister Pressure/Vacuum		
						Initial	Final	Final %
D1A	AMNS 5 DU	4148	6/26/07		TO-15 + NAPHTHALENE			
D2A	AMNS 5 DU	3746	6/26/07		TO-15 + NAPHTHALENE			
D3A	AMNS 2 UU	34712	6/26/07		TO-15 + NAPHTHALENE			
D7A	TREX BLANK	35155	6/26/07					

Relinquished by: (signature) _____ Date: Time _____
 Received by: (signature) _____ Date: Time _____

Relinquished by: (signature) _____ Date: Time _____
 Received by: (signature) _____ Date: Time _____

Relinquished by: (signature) _____ Date: Time _____
 Received by: (signature) _____ Date: Time _____

Shipper Name: _____ Air Bill # _____ Temp (°C): _____ Condition: _____ Custody Seals Intact? Yes No None Work Order #: _____
 Use Only: Fed Ex 861758707617 NA Good Yes No None 0706440

Notes:
 used flow controllers included
 send data pack to Lisa McDougall
 and EOD to detroy@pei-consultants.com



AN ENVIRONMENTAL ANALYTICAL LABORATORY

SAMPLE RECEIPT SUMMARY

WORKORDER 0706440

Client

Ms. Sarah Aldridge
GEI Consultants, Inc.
455 Winding Brook Dr. Suite 201
Glastonbury, CT 06033

Phone

860-368-5300

Fax

860-368-5307

Date Promised: 07/06/07

Date Completed: 7/5/07

Date Received: 6/21/07

PO#: NR

Project#: 061140-8-1703 Bay Shore OU1

Sales Rep: ANS

Total \$: \$ 1,233.00

Logged By: MW

<u>Fraction</u>	<u>Sample #</u>	<u>Analysis</u>	<u>Collected</u>	<u>Receipt Vac./Pres.</u>	<u>Amount\$</u>
01A	AMS 5 DW (4148)	Modified TO-15	6/20/2007	7.0 "Hg	\$225.00
02A	AMS 5 DW (3746)	Modified TO-15	6/20/2007	7.5 "Hg	\$225.00
02AA	AMS 5 DW (3746) Lab Duplicate	Modified TO-15	6/20/2007	7.5 "Hg	\$0.00
03A	AMS 2 UW	Modified TO-15	6/20/2007	7.5 "Hg	\$225.00
04A	TRIP BLANK	Modified TO-15	6/20/2007	4.4 psi	\$225.00
05A	Lab Blank	Modified TO-15	NA	NA	\$0.00
06A	CCV	Modified TO-15	NA	NA	\$0.00
07A	LCS	Modified TO-15	NA	NA	\$0.00

Misc. Charges 6 Liter Summa Canister (3) @ \$50.00 each.	\$150.00
6 Liter Summa Canister (100% Certified) (1) @ \$65.00 each.	\$65.00
Blue Body Flow Controller (3) @ \$35.00 each.	\$105.00
Fuel Surcharge (4) @ \$2.00 each.	\$8.00
Duplicate Sampling T (100% Certified) (1) @ \$5.00 each.	\$5.00

Note: Samples received after 3 P.M. PST are considered to be received on the following work day.
Atlas Project Name/Profile#: Bay Shore OU1 South Perimeter Air/9699

BILL TO: Ms. Sarah Aldridge
GEI Consultants, Inc.
455 Winding Brook Dr. Suite 201
Glastonbury, CT 06033

Analysis Code: TO-14A

TERMS:

Reporting Method: Modified TO-15 + Naph

180 BLUE RAVINE ROAD, SUITE B FOLSOM, CA - 95630
(916) 985-1000 . (800) 985-5955 . FAX (916) 985-1020

Sample Discrepancy Report

Identification

Initiated By: DNW Date: 6/21

Discrepancy Type:
(circle all that apply)

I. II. III.

Workorder(s) affected: 0706440 Sample(s) affected: OIA 302A

I. Sample Receipt Discrepancies

Document on Cover Page of Sample Receipt Confirmation and in Receiving Notes of Lab Narrative

Narration not required:

- COC was not filled out in ink.
- Sample container (cartridge/tube/VOA vial) was received broken, however sample was intact.
- Flow controller used - canister samples received at ambient or under pressure.
- No brass cap on canister.
- VOA vial for RSK-175 analysis received with headspace bubble <5mm.

Narration Required:

- COC improperly relinquished / received.
- Sample tags / can numbers do not match the COC.
- Samples received at wrong temperature (up to 10°C); ice / blue ice (circle one) was present. A temp. blank was / was not present (circle one).
- Custody Seal on the outside of the container was broken / improperly placed (circle one).
- Other (describe below).

Describe the Discrepancy: OIA 302A non unique IDs - added can #s

II. Sample Receipt/Screening Discrepancies requiring CSR notification

Document on Cover Page of Sample Receipt Confirmation and in Receiving Notes of Lab Narrative

If Section II. is filled out CSR must be notified within 24 hrs of Initiation

- COC was not received with samples.
- Analysis method(s) is not specified / incorrectly specified (circle one) on the COC.
- Number of samples on the COC does not match the number of samples that were received.
- Samples were received expired.
- Sampling date / time (sulfur only) is not documented for some / any samples (circle one).
- Sample received with significant (pooling) volume of H₂O in the Tedlar Bag.
- Sample container (cartridge/tube/VOA vial/DNPH Bottle, etc.) was received broken / leaking (circle one); sample can / cannot be analyzed (circle one).
- VOA vial for RSK-175 analysis received with headspace bubble >5mm.
- Samples for RSK-175 CO₂ analysis received preserved with HCl.
- Tedlar Bag received leaking / flat (circle one). Sample can / cannot (circle one) be analyzed.
- Canister was at ambient pressure at time of pressurization and (check all that apply): canister failed leak check on two manifolds, canister valve was open, brass nut was loose. Sample can / cannot be analyzed (circle one).
- Tedlar bag / canister received emitting a strong odor; sample can / cannot (circle one) be analyzed.
- Canister sample received with a vacuum difference >7.0"Hg between the receipt vac. and the final vac. reported on the COC, indicating loss of vacuum.
- Canister sample received at >15"Hg (not identified as a Trip/Field Blank).
- Trip Blank received at low vacuum (< 25"Hg).
- Tedlar Bag for Sulfur analysis has metal fitting.
- Incorrect sampling media / container for analysis requested.
- Sample was received at ≥ 10°C.
- Other (describe below)

Initials: ANC
(if not the original initiator)

Date: 6/23/07

CSR Notified
(see section below)

Describe the Discrepancy: Trip Blank (04A) rec'd @ 4.4psi.

III. Lab Discrepancies requiring Team Leader/CSR notification

Document in Analytical Notes of Lab Narrative

If Section III. is filled out CSR must be notified within 24 hrs of Initiation

- Tedlar Bag found to be leaking at the time of analysis; sample can / cannot (circle one) be analyzed.
- Tedlar Bag found to be flat at the time of analysis.
- Canister found to be leaking at the time of analysis.
- Tedlar Bag received at low volume; sample cannot be analyzed.
- Sulfur samples received with insufficient time to analyze prior to expiration.
- VOST tube saturated; bag dilution necessary.
- Sample loss due to instrument malfunction / broken glassware.
- Other (describe below).

Initials: _____
(if not the original initiator)

Date: _____

CSR Notified
(see section below)

Team Lead Initials: _____

Date: _____

Describe the Discrepancy: _____

Client Services Use Only

Client Services Notification

CSR notified: BL

Date: 6/23/07

Action:

- It is not necessary to notify the client. ~~Narrate the discrepancy by documenting on cover page of Sample Receipt Confirmation and in Receiving Notes/Analytical Notes of Lab Narrative.~~

CSR Initials: BL Date: 6-25-07

- Client notification required. See attached client contact / email, or comments below:

Client Notification:

Person notified: _____ Date: _____

Comments: TRIP BLANKS ARE PRESSURIZED TO 5 PSI PRIOR TO SHIPPING.
DO NOT NARRATE.

Lab notified Name: _____ Date: _____

Additional Notifications

CSR notified: _____

Date: _____

Action:

- It is not necessary to notify the client. Narrate the discrepancy by documenting on cover page of Sample Receipt Confirmation and in Receiving Notes/Analytical Notes of Lab Narrative.

CSR Initials: _____ Date: _____

- Client notification required. See attached client contact / email, or comments below:

Client Notification:

Person notified: _____ Date: _____

Comments: _____

Lab notified Name: _____ Date: _____

- Additional notifications attached.

Other Records

DILUTION FACTORS

$$\text{Dilution Factor} = \frac{\text{Final Pressure}}{\text{Initial Vacuum}} = \frac{14.7 \text{ psi} + \text{Final Pressure (psi)}}{14.7 \text{ psi} - [(\text{Initial Pressure ("Hg)}) (14.7 \text{ psi} / 30 \text{ "Hg})]}$$

$$\text{Dilution Factor} = \frac{\text{Final Pressure}}{\text{Initial Pressure}} = \frac{14.7 \text{ psi} + \text{Final Pressure (psi)}}{14.7 \text{ psi} + \text{Initial Pressure (psi)}}$$

Initial Vacuum ("Hg)	5 psi Final Press. Dil. Factor	10 psi Final Press. Dil. Factor	15 psi Final Press. Dil. Factor
0.0	1.34	1.68	2.02
0.5	1.36	1.71	2.05
1.0	1.39	1.74	2.09
1.5	1.41	1.77	2.13
2.0	1.44	1.80	2.16
2.5	1.46	1.83	2.20
3.0	1.49	1.87	2.24
3.5	1.52	1.90	2.29
4.0	1.55	1.94	2.33
4.5	1.58	1.98	2.38
5.0	1.61	2.02	2.42
5.5	1.64	2.06	2.47
6.0	1.68	2.10	2.53
6.5	1.71	2.15	2.58
7.0	1.75	2.19	2.64
7.5	1.79	2.24	2.69
8.0	1.83	2.29	2.76
8.5	1.87	2.34	2.82
9.0	1.91	2.40	2.89
9.5	1.96	2.46	2.96
10.0	2.01	2.52	3.03
10.5	2.06	2.59	3.11
11.0	2.12	2.65	3.19
11.5	2.17	2.72	3.28
12.0	2.23	2.80	3.37
12.5	2.30	2.88	3.46
13.0	2.36	2.97	3.57
13.5	2.44	3.06	3.67
14.0	2.51	3.15	3.79
14.5	2.59	3.25	3.91
15.0	2.68	3.36	4.04
15.5	2.77	3.48	4.18
16.0	2.87	3.60	4.33
16.5	2.98	3.73	4.49
17.0	3.09	3.88	4.66
17.5	3.22	4.03	4.85
18.0	3.35	4.20	5.05
18.5	3.50	4.38	5.27
19.0	3.65	4.58	5.51
19.5	3.83	4.80	5.77
20.0	4.02	5.04	6.06
20.5	4.23	5.31	6.38

Initial Vacuum ("Hg)	5 psi Final Press. Dil. Factor	10 psi Final Press. Dil. Factor	15 psi Final Press. Dil. Factor
21.0	4.47	5.60	6.73
21.5	4.73	5.93	7.13
22.0	5.03	6.30	7.58
22.5	5.36	6.72	8.08
23.0	5.74	7.20	8.66
23.5	6.19	7.76	9.32
24.0	6.70	8.40	10.10
24.5	7.31	9.17	11.02
25.0	8.04	10.08	12.12
25.5	8.93	11.20	13.47
26.0	10.05	12.60	15.15
26.5	11.49	14.40	17.32
27.0	13.40	16.80	20.20
27.5	16.08	20.16	24.24
28.0	20.10	25.20	30.31
28.5	26.80	33.61	40.41
29.0	40.20	50.41	60.61

Initial Pressure (psi)	5 psi Final Press. Dil. Factor	10 psi Final Press. Dil. Factor	15 psi Final Press. Dil. Factor
0.0	1.34	1.68	2.02
0.2	1.32	1.66	1.99
0.4	1.30	1.64	1.97
0.6	1.29	1.61	1.94
0.8	1.27	1.59	1.92
1.0	1.25	1.57	1.89
1.2	1.24	1.55	1.87
1.4	1.22	1.53	1.84
1.6	1.21	1.52	1.82
1.8	1.19	1.50	1.80
2.0	1.18	1.48	1.78
2.2	1.17	1.46	1.76
2.4	1.15	1.44	1.74
2.6	1.14	1.43	1.72
2.8	1.13	1.41	1.70
3.0	1.11	1.40	1.68
3.2	1.10	1.38	1.66
3.4	1.09	1.36	1.64
3.6	1.08	1.35	1.62
3.8	1.06	1.34	1.61
4.0	1.05	1.32	1.59

DILUTION FACTORS

$$\text{Dilution Factor} = \frac{\text{Final Pressure}}{\text{Initial Pressure}} = \frac{14.7 \text{ psi} + \text{Final Pressure (psi)}}{14.7 \text{ psi} + \text{Initial Pressure (psi)}}$$

Initial Pressure (psi)	5 psi Final Press. Dil. Factor	10 psi Final Press. Dil. Factor	15 psi Final Press. Dil. Factor
0.0	1.34	1.68	2.02
0.2	1.32	1.66	1.99
0.4	1.30	1.64	1.97
0.6	1.29	1.61	1.94
0.8	1.27	1.59	1.92
1.0	1.25	1.57	1.89
1.2	1.24	1.55	1.87
1.4	1.22	1.53	1.84
1.6	1.21	1.52	1.82
1.8	1.19	1.50	1.80
2.0	1.18	1.48	1.78
2.2	1.17	1.46	1.76
2.4	1.15	1.44	1.74
2.6	1.14	1.43	1.72
2.8	1.13	1.41	1.70
3.0	1.11	1.40	1.68
3.2	1.10	1.38	1.66
3.4	1.09	1.36	1.64
3.6	1.08	1.35	1.62
3.8	1.06	1.34	1.61
4.0	1.05	1.32	1.59
4.2	1.04	1.31	1.57
4.4	1.03	1.29	1.55
4.6	1.02	1.28	1.54
4.8	1.01	1.27	1.52
5.0	1.00	1.25	1.51
5.2	NA	1.24	1.49
5.4	NA	1.23	1.48
5.6	NA	1.22	1.46
5.8	NA	1.20	1.45
6.0	NA	1.19	1.43
6.2	NA	1.18	1.42
6.4	NA	1.17	1.41
6.6	NA	1.16	1.39
6.8	NA	1.15	1.38
7.0	NA	1.14	1.37
7.2	NA	1.13	1.36
7.4	NA	1.12	1.34

Initial Pressure (psi)	5 psi Final Press. Dil. Factor	10 psi Final Press. Dil. Factor	15 psi Final Press. Dil. Factor
7.6	NA	1.11	1.33
7.8	NA	1.10	1.32
8.0	NA	1.09	1.31
8.2	NA	1.08	1.30
8.4	NA	1.07	1.29
8.6	NA	1.06	1.27
8.8	NA	1.05	1.26
9.0	NA	1.04	1.25
9.2	NA	1.03	1.24
9.4	NA	1.02	1.23
9.6	NA	1.02	1.22
9.8	NA	1.01	1.21
10.0	NA	1.00	1.20
10.2	NA	NA	1.19
10.4	NA	NA	1.18
10.6	NA	NA	1.17
10.8	NA	NA	1.16
11.0	NA	NA	1.16
11.2	NA	NA	1.15
11.4	NA	NA	1.14
11.6	NA	NA	1.13
11.8	NA	NA	1.12
12.0	NA	NA	1.11
12.2	NA	NA	1.10
12.4	NA	NA	1.10
12.6	NA	NA	1.09
12.8	NA	NA	1.08
13.0	NA	NA	1.07
13.2	NA	NA	1.06
13.4	NA	NA	1.06
13.6	NA	NA	1.05
13.8	NA	NA	1.04
14.0	NA	NA	1.03
14.2	NA	NA	1.03
14.4	NA	NA	1.02
14.6	NA	NA	1.01
14.8	NA	NA	1.01

Compound Listing

Modified TO-15 + Naph

CAS Number	Compound	Detection Limit	Type
		ppbv	
75-71-8	Freon 12	0.50	
76-14-2	Freon 114	0.50	
108-38-3	m,p-Xylene	0.50	
95-47-6	o-Xylene	0.50	
100-42-5	Styrene	0.50	
79-34-5	1,1,2,2-Tetrachloroethane	0.50	
108-67-8	1,3,5-Trimethylbenzene	0.50	
95-63-6	1,2,4-Trimethylbenzene	0.50	
541-73-1	1,3-Dichlorobenzene	0.50	
106-46-7	1,4-Dichlorobenzene	0.50	
100-44-7	alpha-Chlorotoluene	0.50	
95-50-1	1,2-Dichlorobenzene	0.50	
106-99-0	1,3-Butadiene	0.50	
110-54-3	Hexane	0.50	
110-82-7	Cyclohexane	0.50	
142-82-5	Heptane	0.50	
75-27-4	Bromodichloromethane	0.50	
124-48-1	Dibromochloromethane	0.50	
98-82-8	Cumene	0.50	
103-65-1	Propylbenzene	0.50	
74-87-3	Chloromethane	2.0	
120-82-1	1,2,4-Trichlorobenzene	2.0	
87-68-3	Hexachlorobutadiene	2.0	
67-64-1	Acetone	2.0	
75-15-0	Carbon Disulfide	0.50	
67-63-0	2-Propanol	2.0	
156-60-5	trans-1,2-Dichloroethene	0.50	
78-93-3	2-Butanone (Methyl Ethyl Ketone)	0.50	
109-99-9	Tetrahydrofuran	0.50	
123-91-1	1,4-Dioxane	2.0	
108-10-1	4-Methyl-2-pentanone	0.50	
591-78-6	2-Hexanone	2.0	
75-25-2	Bromoform	0.50	
622-96-8	4-Ethyltoluene	0.50	
64-17-5	Ethanol	2.0	
1634-04-4	Methyl tert-butyl ether	0.50	
91-20-3	Naphthalene	2.0	
107-05-1	3-Chloropropene	2.0	
540-84-1	2,2,4-Trimethylpentane	0.50	
2037-26-5	Toluene-d8		
17060-07-0	1,2-Dichloroethane-d4		
460-00-4	4-Bromofluorobenzene		
75-01-4	Vinyl Chloride	0.50	
74-83-9	Bromomethane	0.50	
75-00-3	Chloroethane	0.50	
75-69-4	Freon 11	0.50	

Compound Listing

Modified TO-15 + Naph

CAS Number	Compound	Detection Limit	Type
		ppbv	
75-35-4	1,1-Dichloroethene	0.50	
76-13-1	Freon 113	0.50	
75-09-2	Methylene Chloride	0.50	
75-34-3	1,1-Dichloroethane	0.50	
156-59-2	cis-1,2-Dichloroethene	0.50	
67-66-3	Chloroform	0.50	
71-55-6	1,1,1-Trichloroethane	0.50	
56-23-5	Carbon Tetrachloride	0.50	
71-43-2	Benzene	0.50	
107-06-2	1,2-Dichloroethane	0.50	
79-01-6	Trichloroethene	0.50	
78-87-5	1,2-Dichloropropane	0.50	
10061-01-5	cis-1,3-Dichloropropene	0.50	
108-88-3	Toluene	0.50	
10061-02-6	trans-1,3-Dichloropropene	0.50	
79-00-5	1,1,2-Trichloroethane	0.50	
127-18-4	Tetrachloroethene	0.50	
106-93-4	1,2-Dibromoethane (EDB)	0.50	
108-90-7	Chlorobenzene	0.50	
100-41-4	Ethyl Benzene	0.50	



www.airtoxics.com
1-800-985-5955

Media Certification Report

Canister Number: F051741;6L#35155:1

Date: 5/18/2007 01:54:14

Peak #	Quantification	CAS	Type	Conc.	Units
	1,1,1,2-Tetrafluoroethane		Not Found	0.000	ppbv
	1,1-Difluoroethane		Not Found	0.000	ppbv
	Freon 12		Not Found	0.000	ppbv
	Freon 114		Not Found	0.000	ppbv
	Chloromethane		Not Found	0.000	ppbv
	Butane		Not Found	0.000	ppbv
	Vinyl Chloride		Not Found	0.000	ppbv
	1,3-Butadiene		Not Found	0.000	ppbv
	Bromomethane		Not Found	0.000	ppbv
	Chloroethane		Not Found	0.000	ppbv
	Isopentane		Not Found	0.000	ppbv
	Vinyl bromide		Not Found	0.000	ppbv
	Freon 11		Not Found	0.000	ppbv
	Ethanol		Not Found	0.000	ppbv
	Freon 113		Not Found	0.000	ppbv
	1,1-Dichloroethene		Not Found	0.000	ppbv
	2-Methylpentane		Not Found	0.000	ppbv
	3-Chloropropene		Not Found	0.000	ppbv
	Methyl Acetate		Not Found	0.000	ppbv
	tert-Butyl alcohol		Not Found	0.000	ppbv
	trans-1,2-Dichloroethene		Not Found	0.000	ppbv
	Methyl tert-butyl ether		Not Found	0.000	ppbv
	Hexane		Not Found	0.000	ppbv
	Isopropyl ether		Not Found	0.000	ppbv
	1,1-Dichloroethane		Not Found	0.000	ppbv
	Vinyl Acetate		Not Found	0.000	ppbv
	Chloroprene		Not Found	0.000	ppbv
	Ethyl-tert-butyl ether		Not Found	0.000	ppbv
	2,2-Dichloropropane		Not Found	0.000	ppbv
	cis-1,2-Dichloroethene		Not Found	0.000	ppbv
	2-Butanone (Methyl Ethyl Ketone)		Not Found	0.000	ppbv
	Chloroform		Not Found	0.000	ppbv
	Cyclohexane		Not Found	0.000	ppbv
	2,3-Dimethylpentane		Not Found	0.000	ppbv
	1,1,1-Trichloroethane		Not Found	0.000	ppbv
	Carbon Tetrachloride		Not Found	0.000	ppbv
	1,1-Dichloropropene		Not Found	0.000	ppbv
	2,2,4-Trimethylpentane		Not Found	0.000	ppbv
	tert-Amyl Methyl ether		Not Found	0.000	ppbv
	Heptane		Not Found	0.000	ppbv
	1,2-Dichloroethane		Not Found	0.000	ppbv
	Thiophene		Not Found	0.000	ppbv
	Trichloroethene		Not Found	0.000	ppbv
	Methylcyclohexane		Not Found	0.000	ppbv
	1,2-Dichloropropane		Not Found	0.000	ppbv
	1,4-Dioxane		Not Found	0.000	ppbv
	Dibromomethane		Not Found	0.000	ppbv
	Bromodichloromethane		Not Found	0.000	ppbv



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1-800-985-5955

Media Certification Report

Canister Number: F051741;6L#35155:1

Date: 5/18/2007 01:54:14

Peak #	Quantification	CAS	Type	Conc.	Units
	cis-1,3-Dichloropropene		Not Found	0.000	ppbv
	4-Methyl-2-pentanone		Not Found	0.000	ppbv
	trans-1,3-Dichloropropene		Not Found	0.000	ppbv
	1,1,2-Trichloroethane		Not Found	0.000	ppbv
	Tetrachloroethene		Not Found	0.000	ppbv
	2-Hexanone		Not Found	0.000	ppbv
	Dibromochloromethane		Not Found	0.000	ppbv
	1,2-Dibromoethane (EDB)		Not Found	0.000	ppbv
	Chlorobenzene		Not Found	0.000	ppbv
	Nonane		Not Found	0.000	ppbv
	Ethyl Benzene		Not Found	0.000	ppbv
	1,1,1,2-Tetrachloroethane		Not Found	0.000	ppbv
	m,p-Xylene		Not Found	0.000	ppbv
	o-Xylene		Not Found	0.000	ppbv
	Styrene		Not Found	0.000	ppbv
	Bromoform		Not Found	0.000	ppbv
	Cumene		Not Found	0.000	ppbv
	1,1,2,2-Tetrachloroethane		Not Found	0.000	ppbv
	Propylbenzene		Not Found	0.000	ppbv
	1,2,3-Trichloropropane		Not Found	0.000	ppbv
	4-Ethyltoluene		Not Found	0.000	ppbv
	1,3,5-Trimethylbenzene		Not Found	0.000	ppbv
	tert-Butylbenzene		Not Found	0.000	ppbv
	1,2,4-Trimethylbenzene		Not Found	0.000	ppbv
	Pentachloroethane		Not Found	0.000	ppbv
	sec-Butylbenzene		Not Found	0.000	ppbv
	p-Cymene		Not Found	0.000	ppbv
	1,3-Dichlorobenzene		Not Found	0.000	ppbv
	1,4-Dichlorobenzene		Not Found	0.000	ppbv
	alpha-Chlorotoluene		Not Found	0.000	ppbv
	Indan		Not Found	0.000	ppbv
	Butylbenzene		Not Found	0.000	ppbv
	1,2-Dichlorobenzene		Not Found	0.000	ppbv
	Indene		Not Found	0.000	ppbv
	Hexachloroethane		Not Found	0.000	ppbv
	1,2-Dibromo-3-chloropropane		Not Found	0.000	ppbv
	1,2,4-Trichlorobenzene		Not Found	0.000	ppbv
	Hexachlorobutadiene		Not Found	0.000	ppbv
	Naphthalene		Not Found	0.000	ppbv
	1,2,3-Trichlorobenzene		Not Found	0.000	ppbv
1	Propylene	55255-50-0	Quantified	0.02137	ppbv
8	Acrolein	55255-50-0	Quantified	0.08169	ppbv
10	Carbon Disulfide	3294-96-0	Quantified	0.02359	ppbv
11	Acetone	1073-91-2	Quantified	0.1756	ppbv
12	2-Propanol	0-00-0	Quantified	0.08855	ppbv
14	Methylene Chloride	75-09-2	Quantified	0.02411	ppbv
15	Acrylonitrile	466-96-6	Quantified	0.02881	ppbv
20	Ethyl Acetate	523-14-8	Quantified	0.000	ppbv



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1-800-985-5955

Media Certification Report

Canister Number: F051741;6L #35155:1

Date: 5/18/2007 01:54:14

Peak #	Quantification	CAS	Type	Conc.	Units
21	Bromochloromethane-IS	74-97-5	Quantified	0.000	ppbv
22	Tetrahydrofuran	150832-18-1	Quantified	0.09349	ppbv
23	Benzene	1111-97-3	Quantified	0.03402	ppbv
24	1,2-Dichloroethane-d4	121505-32-6	Quantified	4.781	ppbv
25	1,4-Difluorobenzene-IS	540-36-3	Quantified	0.000	ppbv
29	Toluene-D8	2037-26-5	Quantified	5.067	ppbv
30	Toluene	571-31-3	Quantified	0.01255	ppbv
33	Chlorobenzene-d5-IS	3114-55-4	Quantified	0.000	ppbv
34	Bromofluorobenzene	460-00-4	Quantified	4.917	ppbv

DATA REVIEW CHECKLIST

Work Order #:

0700440

- | | | | | |
|-------------------------------------|-------------------------------------|--------------------------|--------------------------|--------------------------|
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
- Analysis/Reporting vs. Project Profile/SOP requirements checked (i.e. 100% Dups, J-Flag to MDL, etc)
 - The final report has the correct reporting list, special units, and header info.
 - Lab Narrative is correct (proper method & description/Receiving & Analytical notes correct)
-
- Corrective Action issued: # _____
- Unusual circumstances have been documented in the notes section below

LUMEN validation report present and initiated

CIRCLE (YES / NO)

- | | | | | | |
|-------------------------------------|--------------------------|-------------------------------------|-------------------------------------|--------------------------|---|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Lab Blank, CCV, LCS and DUP met QC criteria |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Hold time is met for all samples |
| <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Appropriate data qualifier flags are applied |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Manual integrations for samples and QC are properly documented |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Samples analyzed within the project or method specific clock |
| <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Retention times have been verified |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Appropriate ICAL(s) included |
| <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | At least one result per sample is verified against the target quant sheets/raw data |

- | | | | | | |
|-------------------------------------|-------------------------------------|--------------------------|--------------------------|--------------------------|---|
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Dilution factor correctly calculated (sample load volume, syringe and bag dilutions, can pressurization(s)) |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Correct amount of sample analyzed (i.e. sample not over-diluted) |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Spectra verified - documentation of spectral defense included (Section 5A of eCVP pkg) |
-
- | | | | | | |
|-------------------------------------|-------------------------------------|--------------------------|-------------------------------------|--------------------------|--|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | TICs resemble reference spectra |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | TICs between duplicate samples are consistent |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Checked samples for trends (i.e. Influent>Effluent, Landfill or Ambient etc) |
-
- | | | | | | |
|-------------------------------------|-------------------------------------|--------------------------|--------------------------|--------------------------|--|
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Special units for all samples in the final report are correctly calculated |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Manually entered results checked (i.e. special CCV compounds) |
-
- | | | | | | |
|-------------------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | TPH/NMOC (verify calculations and correct reference compound used) |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Chain of Custody scanned correctly |
| <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Verify sample id's vs. chain of custody |
-
- | | | | | | |
|-------------------------------------|-------------------------------------|--------------------------|--------------------------|--------------------------|--|
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Samples pressurized w/ appropriate gas (N ₂ or He) <input type="checkbox"/> Tedlar Bag only |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Final pressure consistent with canister size (6L vs. 1L) |
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Verify receipt pressures against logbook and Target |
-
- | | | | | | |
|-------------------------------------|-------------------------------------|-------------------------------------|-------------------------------------|--------------------------|---|
| <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Verify canister ID #'s |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Extra printed copies are provided per client profile |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | Final invoice amount correct (adjusted for TAT, Penalties, Re-issue Charges etc.) |
-
- | | | | | | |
|-------------------------------------|--------------------------|-------------------------------------|--------------------------|--------------------------|--|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | Client LUMEN report reviewed for accuracy and completeness |
|-------------------------------------|--------------------------|-------------------------------------|--------------------------|--------------------------|--|

Notes: (to include: noting samples with QA/QC problems, Blanks with positive hits, narratives, etc.)

A/R: Dup on 02A
φ out in CCV, LCS

M/Q: _____

A (Analytical Review/Date)	R/T (Reporting Review/Date)	M (Management Review/Date)	Q (QA Review/Date)
DA 7/4/07	R: [Signature] 7/5/07	[Signature] 7-5-07	

T: _____

Not Applicable