

**REDACTED**

### Data Validation Checklist Semivolatile Organic Analyses

Project: 35<sup>TH</sup> Avenue Superfund Site  
 Laboratory: TestAmerica – Tampa, FL  
 Method: SW-846 8270C Low-Level (PAH)  
 Matrix: Soil and water  
 Reviewer: Jane Lindsey  
 Concurrence<sup>1</sup>: Carol Lovett, Sarah Choyke

Project No: 15268508.20000  
 Job ID.: 680-88766-2  
 Associated Samples: Refer to Attachment A (Sample Summary)  
 Date(s) Collected: 03/25/2013 and 03/26/2013  
 Date: 04/10/2013  
 Date: 04/19/2013

Review Questions	Yes	No	N/A	Samples (Analytes) Affected/Comments	Flag
1. Were sample storage and preservation requirements met? If temperature >6°C, then J/UJ-flag results.	✓				
2. Were all COC records signed and integrity seals intact, indicating that COC was maintained for all samples?	✓				
3. Were there any problems noted in laboratory data package concerning condition of samples upon receipt?		✓			
4. Do any soil samples contain more than 50% water? If yes, then results are to be reported on a wet-weight basis.		✓			
5. Were holding times met (≤7 and 14 days from collection to extraction for aqueous and solid samples, respectively; ≤40 days from extraction to analysis)? If not, then J/UJ-flag sample results. If grossly (2x) exceeded, then flag J/R.	✓				
6. Were results for all project-specified target analytes reported?	✓				
7. Were project-specified Reporting Limits achieved for undiluted sample analyses?	✓				
8. Were samples with analyte concentrations exceeding the calibration range of the instrument re-analyzed at a higher dilution? If not, then J-flag sample result.			✓		
9. Was a method blank extracted with each batch (i.e., one per 20 samples, per batch, per matrix and per level)?	✓				
10. Were target analytes detected in the method blank?		✓			
11. Were target analytes detected in equipment/rinsate blanks?		✓		PAH were not detected during the analysis of rinsate blank 032613-RB-Shovel (680-88766-23).	
12. Are equipment/rinsate blanks associated with every sample? If	✓			According to the QAPP, a rinsate blank is to be collected after each decontamination event, which	

<sup>1</sup> Independent technical reviewer  
 URS Group, Inc.  
 Page 1 of 5

Data Validation Checklist (Continued)

Review Questions	Yes	No	N/A	Samples (Analytes) Affected/Comments	Flag
no, note in DV report.				occurs once per week per the client. A rinsate blank (032613-RB-Shovel) was collected during the week of 03/25/2013. The rinsate blank was analyzed for PAHs under this Test America Job ID.	
13. Were analytes detected in samples below the blank contamination action level? If yes, U-flag positive sample results <5x associated blank concentration (10x for common blank contaminants – phthalates)			✓	Blank contamination does not exist.	
14. Is a field duplicate associated with this Job?		✓			
15. Was precision deemed acceptable as defined by the project plans?			✓		
16. Were DFTPP ion abundance criteria (i.e., Table 3 of SW-846 8270C) met? If no, professional judgment may be applied to determine to what extent the data may be utilized.	✓			Alternate tuning criteria were used by the laboratory (i.e., EPA Method 525.2). All ion abundance criteria were met per EPA Method 525.2.	
17. Were samples analyzed within 12 hours of the DFTPP tune? If no, professional judgment may be applied to determine to what extent the data may be utilized.	✓				
18. Were initial and continuing calibration standards analyzed at the proper frequency for each instrument? <ul style="list-style-type: none"> <li>• Ensure that a minimum of five standards are used for the initial calibration. If no, use professional judgment to determine the effect on the data and note in the reviewer narrative.</li> <li>• An initial calibration is to be associated with each sample analysis.</li> <li>• A continuing calibration standard is to be analyzed for every 12 hours of sample analysis per instrument.</li> </ul>	✓			<ul style="list-style-type: none"> <li>• Initial Calibration: 04/02/2013, instrument BSMC5973</li> <li>• ICV: 04/02/2013 @ 15:34</li> <li>• CCV: 04/03/2013 @ 11:45</li> <li>• CCV: 04/05/2013 @ 12:15</li> </ul>	
19. Were calibration results within laboratory/project specifications? <ul style="list-style-type: none"> <li>• ICAL (Criteria: <math>\leq 15</math> mean %RSD with individual CCC %RSD <math>\leq 30</math> (<math>\leq 50\%</math> for poor performers), OR <math>r \geq 0.995</math>, OR <math>r^2 \geq 0.99</math>, and RRF <math>\geq 0.050</math> (<math>\geq 0.010</math> for poor performers)): <ul style="list-style-type: none"> <li>○ If %RSD <math>&gt; 15</math> (<math>&gt; 50\%</math> for poor performers), or <math>r &lt; 0.995</math>, or <math>r^2 &lt; 0.995</math>, then J-flag positive results and UJ-flag non-detects</li> <li>○ If mean RRF <math>&lt; 0.050</math> (<math>&lt; 0.010</math> for poor performers), then J-flag positive results and R-flag non-detects</li> </ul> </li> <li>• ICV and CCV (Criteria: <math>\leq 20\%D</math> (<math>\leq 50\%</math> for poor performers) and RF <math>\geq 0.050</math> (<math>\geq 0.010</math> for poor performers)):</li> </ul>		✓		ICV of 04/02/2013 @ 15:34, instrument BSMC5973: <ul style="list-style-type: none"> <li>• Pyrene @ -21.4%D (Lab: <math>\leq 35</math>, Project: <math>\leq 20</math>), 78.5%R</li> <li>• Chrysene @ -23.5%D (Lab: <math>\leq 35</math>, Project: <math>\leq 20</math>), 76.5%R</li> <li>• Benzo(b)fluoranthene @ -21.1%D (Lab: <math>\leq 35</math>, Project: <math>\leq 20</math>), 79%R</li> <li>• Benzo(a)pyrene @ -24.3%D (Lab: <math>\leq 35</math>, Project: <math>\leq 20</math>), 75.5%R</li> </ul> A negative bias is indicated by the ICV percent difference, therefore, J and UJ flag results in associated	J, UJ

Data Validation Checklist (Continued)

Review Questions	Yes	No	N/A	Samples (Analytes) Affected/Comments	Flag
<ul style="list-style-type: none"> <li>○ If %D&gt;20 (&gt;50% for poor performers), then J-flag positive results and UJ-flag non-detects</li> <li>○ If RF &lt;0.050 (&lt;0.010 for poor performers), then UJ-flag non-detected semivolatile target compounds</li> </ul>				samples <sup>2</sup> .	
20. Was a LCS prepared for each batch and matrix?	✓				
21. Were LCS recoveries within lab control limits? If no, J-flag positive results when %R >Upper Control Limit (UCL) and J/R-flag results when %R <Lower Control Limit (LCL).	✓				
22. Were LCS/LCSD RPD within lab specifications? If no, J-flag positive results and UJ-flag non-detects	✓			LCSD associated with water prep batch 136013.	
23. Was a MS/MSD pair extracted at the proper frequency (one per 20 samples per batch)?	✓			<ul style="list-style-type: none"> <li>• Water Prep Batch 136013: 680-88766-23 (032613-RB-Shovel), MS only due to limited sample volume. A LCSD analysis was conducted in lieu of the MSD.</li> <li>• Soil Prep Batch 136063: 680-88766-21 (CV0014AB-GS), MS/MSD</li> </ul>	
24. Is the MS/MSD parent sample a project-specific sample?	✓			See above.	
25. Were MS/MSD recoveries within laboratory/project specifications? <i>Only QC results for project samples that are reported under this Job ID are evaluated.</i> <ul style="list-style-type: none"> <li>• If the native sample concentration &gt; 4x spiking level, then an evaluation of interference is not possible.</li> <li>• If either MS or MSD recovery meets control limits, qualification of data is not warranted.</li> <li>• MS and MSD %R&lt;10: J and R Flag positive and ND results, respectively</li> <li>• MS and MSD %R &gt;10 and &lt;LCL: J-Flag positive and UJ-flag non-detect results</li> <li>• MS and MSD R% &gt;UCL (or 140): J-Flag positive results</li> </ul>		✓		032613-RB-Shovel (680-88766-23): Refer to Attachment B, MS recoveries. UJ Flag results due to low MS recoveries.	UJ
26. Were laboratory criteria met for precision during the MS/MSD analysis? <i>Only QC results for project samples that are reported under this Job ID are evaluated.</i> <ul style="list-style-type: none"> <li>• If the native sample concentration &gt; 4x spiking level, then an evaluation of interference is not possible.</li> <li>• If %RPD &gt; UCL, J-flag positive result and UJ-flag non-detect result.</li> </ul>	✓				

Data Validation Checklist (Continued)

Review Questions	Yes	No	N/A	Samples (Analytes) Affected/Comments	Flag
27. Were surrogate recoveries within lab/project specifications? <ul style="list-style-type: none"> <li>• If %R &lt;10, then J-flag positive and R-flag non-detect associated sample results</li> <li>• If %R &gt;UCL, then J-flag positive results</li> <li>• %R ≥10%, but &lt;LCL, then J-flag positive results and UJ-flag non-detect results</li> <li>• If 1 %R &gt;UCL and 1 %R ≥10%, but &lt;LCL, then J-flag positive results and UJ-flag non-detect results</li> </ul>	✓				
28. Were internal standard (IS) results within lab/project specifications? <ul style="list-style-type: none"> <li>• If IS area counts are less than 50% of the midpoint calibration standard, then J-flag positive and UJ-flag non-detect associated sample results</li> <li>• If IS area counts are greater than 100% of the midpoint calibration standard, then J-flag positive results</li> <li>• If extremely low area counts are reported or performance exhibits a major abrupt drop-off, then a severe loss of sensitivity is indicated, J-flag positive and R-flag non-detect results</li> <li>• If retention time of sample's internal standard is not within 30 seconds of the associated calibration standard, R-flag associated data.</li> <li>• The chromatographic profile for that sample must be examined to determine if any false positives or negatives exists. For shifts of large magnitude, the reviewer may consider partial or total rejection of the data for that sample fraction. Positive results need not be qualified as R, if mass spectral criteria are met.</li> </ul>	✓				
29. Were lab comments included in report?	✓			Refer to <b>Attachment C</b> (Case Narrative)	
<p><b>Comments:</b> The data validation was conducted in accordance with the <i>Non-Industrial Use Property Sampling Event QAPP for the 35th Avenue Removal Site, Birmingham, Alabama, Revision 1</i> (OTIE, October 2012). The data review process was modeled after the <i>USEPA Contract Laboratory Program (CLP) National Functional Guidelines (NFG) for Organic Methods Data Review</i> (EPA, October 1999) and <i>USEPA CLP NFG for Low Concentration Organic Methods Data Review</i> (EPA, June 2001). Sample results have been qualified based on the results of the data review process (<b>Attachment D</b>). Criteria for acceptability of data were based upon available site information, analytical method requirements, guidance documents, and professional judgment.</p>					

## Data Validation Checklist (Continued)

**DV Flag Definitions:**

- J The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
- R The sample results are unusable. The analyte may or may not be present in the sample.
- U The analyte was analyzed for, but was not detected above the associated level; blank contamination may exist.
- UJ The analyte was not detected above the limit, and the limit is approximate and may be inaccurate or imprecise.

**ATTACHMENT A**  
**SAMPLE SUMMARY**

# Sample Summary

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

Lab Sample ID	Client Sample ID	Matrix	Collected	Received
680-88766-21	CV0014AB-GS	Solid	03/25/13 15:18	03/28/13 09:37
680-88766-23	032613-RB-shovel	Water	03/26/13 13:00	03/28/13 09:37

1

2

3

4

5

6

7

8

9

10

11

12

**ATTACHMENT B**

**MS RESULTS**



FORM III  
GC/MS SEMI VOA MATRIX SPIKE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Water Level: Low Lab File ID: 1CD05007.D  
 Lab ID: 680-88766-23 MS Client ID: 032613-RB-shovel MS

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENTRATION (ug/L)	MS CONCENTRATION (ug/L)	MS % REC	QC LIMITS REC	#
Acenaphthene	10.4	2.5 U	4.74	45	55-132	F
Acenaphthylene	10.4	1.3 U	5.18	50	39-130	
Anthracene	10.4	0.25 U	4.97	48	39-130	
Benzo[a]anthracene	10.4	0.25 U	4.72	45	54-135	F
Benzo[a]pyrene	10.4	0.25 U	2.13	20	21-130	F
Benzo[b]fluoranthene	10.4	0.25 U	2.68	26	37-130	F
Benzo[g,h,i]perylene	10.4	0.63 U	1.33	13	26-130	F
Benzo[k]fluoranthene	10.4	0.25 U	2.38	23	38-130	F
Chrysene	10.4	0.25 U	3.55	34	56-130	F
Dibenz(a,h)anthracene	10.4	0.25 U	1.12	11	13-130	F
Fluoranthene	10.4	0.63 U	5.46	52	60-130	F
Fluorene	10.4	2.5 U	4.83	46	55-140	F
Indeno[1,2,3-cd]pyrene	10.4	0.25 U	1.28	12	21-130	F
1-Methylnaphthalene	10.4	2.5 U	5.64	54	49-130	
2-Methylnaphthalene	10.4	2.5 U	4.68	45	48-130	F
Naphthalene	10.4	2.5 U	5.39	52	54-133	F
Phenanthrene	10.4	0.63 U	5.42	52	60-136	F
Pyrene	10.4	0.63 U	5.24	50	60-138	F

# Column to be used to flag recovery and RPD values

**ATTACHMENT C**  
**CASE NARRATIVE**

## Case Narrative

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

**Job ID: 680-88766-2**

**Laboratory: TestAmerica Savannah**

Narrative

### CASE NARRATIVE

**Client: Oneida Total Integrated Enterprises LLC**

**Project: 35th Avenue Superfund Site**

**Report Number: 680-88766-2**

With the exceptions noted as flags or footnotes, standard analytical protocols were followed in the analysis of the samples and no problems were encountered or anomalies observed. In addition all laboratory quality control samples were within established control limits, with any exceptions noted below. Each sample was analyzed to achieve the lowest possible reporting limit within the constraints of the method. In some cases, due to interference or analytes present at high concentrations, samples were diluted. For diluted samples, the reporting limits are adjusted relative to the dilution required.

Calculations are performed before rounding to avoid round-off errors in calculated results.

All holding times were met and proper preservation noted for the methods performed on these samples, unless otherwise detailed in the individual sections below.

#### RECEIPT

The samples were received on 03/28/2013; the samples arrived in good condition, properly preserved and on ice. The temperature of the coolers at receipt was 1.4 C.

#### SEMIVOLATILE ORGANIC COMPOUNDS BY GCMS - LOW LEVEL

Sample CV0014AB-GS (680-88766-21) was analyzed for Semivolatile Organic Compounds by GCMS - Low Level in accordance with EPA SW-846 Method 8270C. The samples were prepared on 04/02/2013 and analyzed on 04/03/2013.

Sample CV0014AB-GS (680-88766-21)[4X] required dilution prior to analysis. The reporting limits have been adjusted accordingly.

No difficulties were encountered during the SVOAs analysis.

All quality control parameters were within the acceptance limits.

#### SEMIVOLATILE ORGANIC COMPOUNDS (GC-MS)-WATER

Sample 032613-RB-shovel (680-88766-23) was analyzed for semivolatile organic compounds (GC-MS) in accordance with EPA SW-846 Method 8270C. The samples were prepared on 04/02/2013 and analyzed on 04/05/2013.

Several analytes recovered outside the recovery criteria low for the MS of sample 032613-RB-shovel (680-88766-23) in batch 660-136171.

No other difficulties were encountered during the semivolatiles analysis.

All other quality control parameters were within the acceptance limits.

**ATTACHMENT D**  
**QUALIFIED SAMPLE RESULTS**

# Client Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

**Client Sample ID: CV0014AB-GS**

**Lab Sample ID: 680-88766-21**

Date Collected: 03/25/13 15:18

Matrix: Solid

Date Received: 03/28/13 09:37

Percent Solids: 73.9

Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels									
Analyte	Result	Qualifier	RL	MDL	Unit	D	Prepared	Analyzed	Dil Fac
Acenaphthene	530	U	530	110	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Acenaphthylene	34	J	210	27	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Anthracene	45	U J	45	22	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Benzo[a]anthracene	200	J	43	21	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Benzo[a]pyrene	100	J	55	28	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Benzo[b]fluoranthene	150	J	65	32	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Benzo[g,h,i]perylene	130	J	110	23	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Benzo[k]fluoranthene	79	J	43	19	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Chrysene	210	J	48	24	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Dibenz(a,h)anthracene	62	J	110	22	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Fluoranthene	210	J	110	21	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Fluorene	39	J	110	22	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Indeno[1,2,3-cd]pyrene	76	J	110	38	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
1-Methylnaphthalene	120	J	210	23	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
2-Methylnaphthalene	130	J	210	38	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Naphthalene	100	J	210	23	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Phenanthrene	200	J	43	21	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Pyrene	210	J	110	20	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Surrogate</b>	<b>%Recovery</b>	<b>Qualifier</b>	<b>Limits</b>				<b>Prepared</b>	<b>Analyzed</b>	<b>Dil Fac</b>
<i>o</i> -Terphenyl	95		30 - 130				04/02/13 11:33	04/03/13 16:10	4

**Client Sample ID: 032613-RB-shovel**

**Lab Sample ID: 680-88766-23**

Date Collected: 03/26/13 13:00

Matrix: Water

Date Received: 03/28/13 09:37

Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels									
Analyte	Result	Qualifier	RL	MDL	Unit	D	Prepared	Analyzed	Dil Fac
Acenaphthene	2.5	U F J	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
Acenaphthylene	1.3	U	1.3	0.31	ug/L		04/02/13 08:08	04/05/13 12:54	1
Anthracene	0.25	U J	0.25	0.095	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[a]anthracene	0.25	U F J	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[a]pyrene	0.25	U F J	0.25	0.071	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[b]fluoranthene	0.25	U F J	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[g,h,i]perylene	0.63	U F J	0.63	0.13	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[k]fluoranthene	0.25	U F J	0.25	0.071	ug/L		04/02/13 08:08	04/05/13 12:54	1
Chrysene	0.25	U F J	0.25	0.086	ug/L		04/02/13 08:08	04/05/13 12:54	1
Dibenz(a,h)anthracene	0.25	U F J	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
Fluoranthene	0.63	U F J	0.63	0.068	ug/L		04/02/13 08:08	04/05/13 12:54	1
Fluorene	2.5	U F J	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
Indeno[1,2,3-cd]pyrene	0.25	U F J	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
1-Methylnaphthalene	2.5	U	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
2-Methylnaphthalene	2.5	U F J	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
Naphthalene	2.5	U F J	2.5	0.31	ug/L		04/02/13 08:08	04/05/13 12:54	1
Phenanthrene	0.63	U F J	0.63	0.25	ug/L		04/02/13 08:08	04/05/13 12:54	1
Pyrene	0.63	U F J	0.63	0.11	ug/L		04/02/13 08:08	04/05/13 12:54	1
<b>Surrogate</b>	<b>%Recovery</b>	<b>Qualifier</b>	<b>Limits</b>				<b>Prepared</b>	<b>Analyzed</b>	<b>Dil Fac</b>
<i>o</i> -Terphenyl	67		30 - 130				04/02/13 08:08	04/05/13 12:54	1

TestAmerica Savannah

Sample results have been qualified by URS in accordance with the Non-Industrial Use Property Sampling Event QAPP for the 35th Avenue Removal Site, Birmingham, Alabama, Revision 1 (OTIE, October 2012)

## ANALYTICAL REPORT

Job Number: 680-88766-2

SDG Number: 68088766-2

Job Description: 35th Avenue Superfund Site

For:

Oneida Total Integrated Enterprises LLC

1220 Kennestone Circle

Suite 106

Marietta, GA 30060

Attention: Ms. Limari F Krebs



Approved for release.  
Bernard Kirkland  
Project Manager I  
4/8/2013 9:11 AM

---

Designee for

Lisa Harvey

Project Manager II

[lisa.harvey@testamericainc.com](mailto:lisa.harvey@testamericainc.com)

04/08/2013

The test results in this report meet NELAP requirements for parameters for which accreditation is required or available. Any exceptions to the NELAP requirements are noted. Results pertain only to samples listed in this report. This report may not be reproduced, except in full, without the written approval of the laboratory. Questions should be directed to the person who signed this report.

Savannah Certifications and ID #s: A2LA: 0399.01; AL: 41450; ARDEQ: 88-0692; ARDOH; AZ: AZ0741; CA: 03217CA; CO; CT: PH0161; DE; FL: E87052; GA: 803; Guam; HI; IL: 200022; IN: C-GA-02; IA: 353; KS: E-10322; KY EPPC: 90084; KY UST; LA DEQ: 30690; LA DHH: LA080008; ME: 2008022; MD: 250; MA: M-GA006; MI: 9925; MS; NFESC: 249; NV: GA00006; NJ: GA769; NM; NY: 10842; NC DWQ: 269; NC DHHS: 13701; PA: 68-00474; PR: GA00006; RI: LAO00244; SC: 98001001; TN: TN0296; TX: T104704185; USEPA: GA00006; VT: VT-87052; VA: 00302; WA; WV DEP: 094; WV DHHR: 9950 C; WI DNR: 999819810; WY/EPAR8: 8TMS-Q

**TestAmerica Laboratories, Inc.**

TestAmerica Savannah 5102 LaRoche Avenue, Savannah, GA 31404

Tel (912) 354-7858 Fax (912) 352-0165 [www.testamericainc.com](http://www.testamericainc.com)



# Table of Contents

Cover Title Page . . . . .	1
Data Summaries . . . . .	4
Report Narrative . . . . .	4
Sample Summary . . . . .	5
Method Summary . . . . .	6
Method / Analyst Summary . . . . .	7
Data Qualifiers . . . . .	8
QC Association Summary . . . . .	9
Manual Integration Summary . . . . .	10
Organic Sample Data . . . . .	17
GC/MS Semi VOA . . . . .	17
Method 8270C Low Level . . . . .	17
Method 8270C Low Level QC Summary . . . . .	18
Method 8270C Low Level Sample Data . . . . .	40
Standards Data . . . . .	66
Method 8270C Low Level ICAL Data . . . . .	66
Method 8270C Low Level CCAL Data . . . . .	118
Raw QC Data . . . . .	137
Method 8270C Low Level Tune Data . . . . .	137
Method 8270C Low Level Blank Data . . . . .	157
Method 8270C Low Level LCS/LCSD Data . . . . .	163
Method 8270C Low Level MS/MSD Data . . . . .	178
Method 8270C Low Level Run Logs . . . . .	195
Method 8270C Low Level Prep Data . . . . .	199
Inorganic Sample Data . . . . .	203
General Chemistry Data . . . . .	203

# Table of Contents

Gen Chem Cover Page .....	204
Gen Chem MDL .....	205
Gen Chem Analysis Run Log .....	207
Gen Chem Prep Data .....	209
Shipping and Receiving Documents .....	210
Client Chain of Custody .....	211
Sample Receipt Checklist .....	212



## CASE NARRATIVE

**Client: Oneida Total Integrated Enterprises LLC**

**Project: 35th Avenue Superfund Site**

**Report Number: 680-88766-2**

With the exceptions noted as flags or footnotes, standard analytical protocols were followed in the analysis of the samples and no problems were encountered or anomalies observed. In addition all laboratory quality control samples were within established control limits, with any exceptions noted below. Each sample was analyzed to achieve the lowest possible reporting limit within the constraints of the method. In some cases, due to interference or analytes present at high concentrations, samples were diluted. For diluted samples, the reporting limits are adjusted relative to the dilution required.

Calculations are performed before rounding to avoid round-off errors in calculated results.

All holding times were met and proper preservation noted for the methods performed on these samples, unless otherwise detailed in the individual sections below.

### **RECEIPT**

The samples were received on 03/28/2013; the samples arrived in good condition, properly preserved and on ice. The temperature of the coolers at receipt was 1.4 C.

### **SEMIVOLATILE ORGANIC COMPOUNDS BY GCMS - LOW LEVEL**

Sample CV0014AB-GS (680-88766-21) was analyzed for Semivolatile Organic Compounds by GCMS - Low Level in accordance with EPA SW-846 Method 8270C. The samples were prepared on 04/02/2013 and analyzed on 04/03/2013.

Sample CV0014AB-GS (680-88766-21)[4X] required dilution prior to analysis. The reporting limits have been adjusted accordingly.

No difficulties were encountered during the SVOAs analysis.

All quality control parameters were within the acceptance limits.

### **SEMIVOLATILE ORGANIC COMPOUNDS (GC-MS)-WATER**

Sample 032613-RB-shovel (680-88766-23) was analyzed for semivolatile organic compounds (GC-MS) in accordance with EPA SW-846 Method 8270C. The samples were prepared on 04/02/2013 and analyzed on 04/05/2013.

Several analytes recovered outside the recovery criteria low for the MS of sample 032613-RB-shovel (680-88766-23) in batch 660-136171.

No other difficulties were encountered during the semivolatiles analysis.

All other quality control parameters were within the acceptance limits.

## SAMPLE SUMMARY

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2  
Sdg Number: 68088766-2

<b>Lab Sample ID</b>	<b>Client Sample ID</b>	<b>Client Matrix</b>	<b>Date/Time Sampled</b>	<b>Date/Time Received</b>
680-88766-21	CV0014AB-GS	Solid	03/25/2013 1518	03/28/2013 0937
680-88766-21MS	CV0014AB-GS	Solid	03/25/2013 1518	03/28/2013 0937
680-88766-21MSD	CV0014AB-GS	Solid	03/25/2013 1518	03/28/2013 0937
680-88766-23	032613-RB-shovel	Water	03/26/2013 1300	03/28/2013 0937

## METHOD SUMMARY

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

Sdg Number: 68088766-2

Description	Lab Location	Method	Preparation Method
<b>Matrix: Solid</b>			
Semivolatile Organic Compounds by GCMS - Low Levels	TAL TAM	SW846 8270C LL	
Microwave Extraction	TAL TAM		SW846 3546
Percent Moisture	TAL TAM	EPA Moisture	
<b>Matrix: Water</b>			
Semivolatile Organic Compounds by GCMS - Low Levels	TAL TAM	SW846 8270C LL	
Liquid-Liquid Extraction (Continuous)	TAL TAM		SW846 3520C

### Lab References:

TAL TAM = TestAmerica Tampa

### Method References:

EPA = US Environmental Protection Agency

SW846 = "Test Methods For Evaluating Solid Waste, Physical/Chemical Methods", Third Edition, November 1986 And Its Updates.

## METHOD / ANALYST SUMMARY

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

Sdg Number: 68088766-2

<b>Method</b>	<b>Analyst</b>	<b>Analyst ID</b>
SW846 8270C LL	Cantin, Stephen C	SCC
EPA Moisture	Galio, Andrew	AG

## DATA REPORTING QUALIFIERS

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

Sdg Number: 68088766-2

<b>Lab Section</b>	<b>Qualifier</b>	<b>Description</b>
GC/MS Semi VOA	U	Indicates the analyte was analyzed for but not detected.
	F	MS or MSD exceeds the control limits
	J	Result is less than the RL but greater than or equal to the MDL and the concentration is an approximate value.

## Quality Control Results

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

Sdg Number: 68088766-2

### QC Association Summary

Lab Sample ID	Client Sample ID	Report Basis	Client Matrix	Method	Prep Batch
<b>GC/MS Semi VOA</b>					
<b>Prep Batch: 660-136013</b>					
LCS 660-136013/2-A	Lab Control Sample	T	Water	3520C	
LCSD 660-136013/3-A	Lab Control Sample Duplicate	T	Water	3520C	
MB 660-136013/1-A	Method Blank	T	Water	3520C	
680-88766-23	032613-RB-shovel	T	Water	3520C	
680-88766-23MS	Matrix Spike	T	Water	3520C	
<b>Prep Batch: 660-136063</b>					
LCS 660-136063/2-A	Lab Control Sample	T	Solid	3546	
MB 660-136063/1-A	Method Blank	T	Solid	3546	
680-88766-21	CV0014AB-GS	T	Solid	3546	
680-88766-21MS	Matrix Spike	T	Solid	3546	
680-88766-21MSD	Matrix Spike Duplicate	T	Solid	3546	
<b>Analysis Batch:660-136081</b>					
LCS 660-136063/2-A	Lab Control Sample	T	Solid	8270C LL	660-136063
MB 660-136063/1-A	Method Blank	T	Solid	8270C LL	660-136063
680-88766-21	CV0014AB-GS	T	Solid	8270C LL	660-136063
680-88766-21MS	Matrix Spike	T	Solid	8270C LL	660-136063
680-88766-21MSD	Matrix Spike Duplicate	T	Solid	8270C LL	660-136063
<b>Analysis Batch:660-136164</b>					
LCS 660-136013/2-A	Lab Control Sample	T	Water	8270C LL	660-136013
LCSD 660-136013/3-A	Lab Control Sample Duplicate	T	Water	8270C LL	660-136013
MB 660-136013/1-A	Method Blank	T	Water	8270C LL	660-136013
<b>Analysis Batch:660-136171</b>					
680-88766-23	032613-RB-shovel	T	Water	8270C LL	660-136013
680-88766-23MS	Matrix Spike	T	Water	8270C LL	660-136013

**Report Basis**

T = Total

**General Chemistry**

<b>Analysis Batch:660-135922</b>					
680-88766-A-6 MS	Matrix Spike	T	Solid	Moisture	
680-88766-A-6 MSD	Matrix Spike Duplicate	T	Solid	Moisture	
680-88766-21	CV0014AB-GS	T	Solid	Moisture	

**Report Basis**

T = Total

## GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMC5973 Analysis Batch Number: 136048Lab Sample ID: IC 660-136048/5 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/02/13 13:26 Lab File ID: 1CD02005.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Dibenz(a,h)anthracene	10.09	Baseline Event	cantins	04/02/13 15:44

Lab Sample ID: IC 660-136048/6 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/02/13 13:44 Lab File ID: 1CD02006.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.01	Split Peak	cantins	04/02/13 15:45

Lab Sample ID: IC 660-136048/7 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/02/13 14:02 Lab File ID: 1CD02007.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.00	Split Peak	cantins	04/02/13 15:48

Lab Sample ID: IC 660-136048/8 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/02/13 14:20 Lab File ID: 1CD02008.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.00	Split Peak	cantins	04/02/13 15:49

Lab Sample ID: ICIS 660-136048/9 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/02/13 14:39 Lab File ID: 1CD02009.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.01	Split Peak	cantins	04/02/13 15:39

GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Instrument ID: BSMC5973 Analysis Batch Number: 136048

Lab Sample ID: IC 660-136048/10 Client Sample ID: \_\_\_\_\_

Date Analyzed: 04/02/13 14:57 Lab File ID: 1CD02010.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.01	Split Peak	cantins	04/02/13 15:50

Lab Sample ID: IC 660-136048/11 Client Sample ID: \_\_\_\_\_

Date Analyzed: 04/02/13 15:15 Lab File ID: 1CD02011.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.02	Split Peak	cantins	04/02/13 15:51

Lab Sample ID: ICV 660-136048/12 Client Sample ID: \_\_\_\_\_

Date Analyzed: 04/02/13 15:34 Lab File ID: 1CD02012.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.01	Split Peak	cantins	04/02/13 15:57



## GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMC5973 Analysis Batch Number: 136081Lab Sample ID: CCVIS 660-136081/3 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/03/13 11:45 Lab File ID: 1CD03003.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	9.99	Split Peak	cantins	04/03/13 11:59

Lab Sample ID: LCS 660-136063/2-A Client Sample ID: \_\_\_\_\_Date Analyzed: 04/03/13 15:52 Lab File ID: 1CD03016.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	9.99	Split Peak	cantins	04/04/13 15:32

Lab Sample ID: 680-88766-21 Client Sample ID: CV0014AB-GSDate Analyzed: 04/03/13 16:10 Lab File ID: 1CD03017.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	10.00	Split Peak	cantins	04/04/13 15:33
Dibenz(a,h)anthracene	10.02	Baseline Event	cantins	04/04/13 15:33
Benzo[g,h,i]perylene	10.35	Baseline Event	cantins	04/04/13 15:33

Lab Sample ID: 680-88766-21 MS Client Sample ID: CV0014AB-GS MSDate Analyzed: 04/03/13 16:29 Lab File ID: 1CD03018.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	9.99	Split Peak	cantins	04/04/13 15:34
Dibenz(a,h)anthracene	10.01	Baseline Event	cantins	04/04/13 15:34
Benzo[g,h,i]perylene	10.34	Baseline Event	cantins	04/04/13 15:34

GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Instrument ID: BSMC5973 Analysis Batch Number: 136081

Lab Sample ID: 680-88766-21 MSD Client Sample ID: CV0014AB-GS MSD

Date Analyzed: 04/03/13 16:47 Lab File ID: 1CD03019.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	9.99	Baseline Event	cantins	04/04/13 15:35

GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Instrument ID: BSMC5973 Analysis Batch Number: 136171

Lab Sample ID: CCVIS 660-136171/4 Client Sample ID: \_\_\_\_\_

Date Analyzed: 04/05/13 12:15 Lab File ID: 1CD05004.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	9.96	Split Peak	cantins	04/05/13 12:33

Lab Sample ID: 680-88766-23 MS Client Sample ID: 032613-RB-shovel MS

Date Analyzed: 04/05/13 13:12 Lab File ID: 1CD05007.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	9.96	Split Peak	cantins	04/05/13 13:29

## GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMD5973 Analysis Batch Number: 136164Lab Sample ID: IC 660-136164/15 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 13:49 Lab File ID: 1DD04007.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.73	Split Peak	cantins	04/05/13 12:28
Dibenz(a,h)anthracene	14.76	Baseline Event	cantins	04/05/13 12:28

Lab Sample ID: IC 660-136164/16 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 14:11 Lab File ID: 1DD04008.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.73	Split Peak	cantins	04/05/13 12:29
Dibenz(a,h)anthracene	14.76	Baseline Event	cantins	04/05/13 12:28

Lab Sample ID: IC 660-136164/17 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 14:34 Lab File ID: 1DD04009.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.74	Split Peak	cantins	04/05/13 12:29

Lab Sample ID: IC 660-136164/18 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 14:57 Lab File ID: 1DD04010.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.75	Split Peak	cantins	04/05/13 12:30

Lab Sample ID: ICIS 660-136164/19 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 15:19 Lab File ID: 1DD04011.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.76	Split Peak	cantins	04/05/13 12:26

## GC/MS SEMI VOA MANUAL INTEGRATION SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMD5973 Analysis Batch Number: 136164Lab Sample ID: IC 660-136164/20 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 15:42 Lab File ID: 1DD04012.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.77	Split Peak	cantins	04/05/13 12:30

Lab Sample ID: IC 660-136164/21 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 16:04 Lab File ID: 1DD04013.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.79	Split Peak	cantins	04/05/13 12:30

Lab Sample ID: ICV 660-136164/22 Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 16:27 Lab File ID: 1DD04014.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Carbazole	9.23	Baseline Event	cantins	04/05/13 13:08
Indeno[1,2,3-cd]pyrene	14.76	Split Peak	cantins	04/05/13 13:09

Lab Sample ID: LCS 660-136013/2-A Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 20:13 Lab File ID: 1DD04023.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.74	Split Peak	cantins	04/05/13 13:13

Lab Sample ID: LCSD 660-136013/3-A Client Sample ID: \_\_\_\_\_Date Analyzed: 04/04/13 20:36 Lab File ID: 1DD04024.D GC Column: DB-5MS ID: 250 (um)

COMPOUND NAME	RETENTION TIME	MANUAL INTEGRATION		
		REASON	ANALYST	DATE
Indeno[1,2,3-cd]pyrene	14.73	Split Peak	cantins	04/05/13 13:14

# Method 8270C Low Level

---

Semivolatile Organic Compounds  
(GC/MS) Low Level by Method 8270C

FORM II  
GC/MS SEMI VOA SURROGATE RECOVERY

Lab Name: TestAmerica Tampa

Job No.: 680-88766-2

SDG No.: 68088766-2

Matrix: Solid

Level: Low

GC Column (1): DB-5MS ID: 250 (um)

Client Sample ID	Lab Sample ID	OTPH #
CV0014AB-GS	680-88766-21	95
	MB 660-136063/1-A	80
	LCS 660-136063/2-A	72
CV0014AB-GS MS	680-88766-21 MS	92
CV0014AB-GS MSD	680-88766-21 MSD	97

OTPH = o-Terphenyl

QC LIMITS  
30-130

# Column to be used to flag recovery values

FORM II 8270C LL

FORM II  
GC/MS SEMI VOA SURROGATE RECOVERY

Lab Name: TestAmerica Tampa

Job No.: 680-88766-2

SDG No.: 68088766-2

Matrix: Water

Level: Low

GC Column (1): DB-5MS ID: 250 (um)

Client Sample ID	Lab Sample ID	OTPH #
032613-RB-shovel	680-88766-23	67
	MB 660-136013/1-A	78
	LCS 660-136013/2-A	72
	LCSD 660-136013/3-A	72
032613-RB-shovel MS	680-88766-23 MS	49

OTPH = o-Terphenyl

QC LIMITS  
30-130

# Column to be used to flag recovery values

FORM II 8270C LL



FORM III  
GC/MS SEMI VOA LAB CONTROL SAMPLE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Water Level: Low Lab File ID: 1DD04023.D  
 Lab ID: LCS 660-136013/2-A Client ID: \_\_\_\_\_

COMPOUND	SPIKE ADDED (ug/L)	LCS CONCENTRATION (ug/L)	LCS % REC	QC LIMITS REC	#
Acenaphthene	10.0	7.38	74	55-132	
Acenaphthylene	10.0	7.55	76	39-130	
Anthracene	10.0	7.12	71	39-130	
Benzo[a]anthracene	10.0	7.21	72	54-135	
Benzo[a]pyrene	10.0	4.96	50	21-130	
Benzo[b]fluoranthene	10.0	5.86	59	37-130	
Benzo[g,h,i]perylene	10.0	3.51	35	26-130	
Benzo[k]fluoranthene	10.0	5.75	57	38-130	
Chrysene	10.0	7.28	73	56-130	
Dibenz(a,h)anthracene	10.0	3.33	33	13-130	
Fluoranthene	10.0	7.83	78	60-130	
Fluorene	10.0	8.04	80	55-140	
Indeno[1,2,3-cd]pyrene	10.0	3.17	32	21-130	
1-Methylnaphthalene	10.0	7.86	79	49-130	
2-Methylnaphthalene	10.0	7.57	76	48-130	
Naphthalene	10.0	7.44	74	54-133	
Phenanthrene	10.0	7.52	75	60-136	
Pyrene	10.0	7.50	75	60-138	

# Column to be used to flag recovery and RPD values

FORM III  
GC/MS SEMI VOA LAB CONTROL SAMPLE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Solid Level: Low Lab File ID: 1CD03016.D  
 Lab ID: LCS 660-136063/2-A Client ID: \_\_\_\_\_

COMPOUND	SPIKE ADDED (ug/Kg)	LCS CONCENTRATION (ug/Kg)	LCS % REC	QC LIMITS REC	#
Acenaphthene	668	458	69	39-130	
Acenaphthylene	668	488	73	38-130	
Anthracene	668	469	70	37-130	
Benzo[a]anthracene	668	484	72	40-130	
Benzo[a]pyrene	668	444	66	49-130	
Benzo[b]fluoranthene	668	516	77	37-130	
Benzo[g,h,i]perylene	668	418	63	32-130	
Benzo[k]fluoranthene	668	468	70	32-130	
Chrysene	668	462	69	41-130	
Dibenz(a,h)anthracene	668	471	71	27-130	
Fluoranthene	668	487	73	40-130	
Fluorene	668	444	66	40-130	
Indeno[1,2,3-cd]pyrene	668	399	60	30-130	
1-Methylnaphthalene	668	522	78	31-130	
2-Methylnaphthalene	668	458	69	33-130	
Naphthalene	668	484	72	36-130	
Phenanthrene	668	499	75	42-130	
Pyrene	668	516	77	44-130	

# Column to be used to flag recovery and RPD values

FORM III  
GC/MS SEMI VOA LAB CONTROL SAMPLE DUPLICATE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Water Level: Low Lab File ID: 1DD04024.D  
 Lab ID: LCSD 660-136013/3-A Client ID: \_\_\_\_\_

COMPOUND	SPIKE ADDED (ug/L)	LCSD CONCENTRATION (ug/L)	LCSD % REC	% RPD	QC LIMITS		#
					RPD	REC	
Acenaphthene	10.0	7.35	73	0	35	55-132	
Acenaphthylene	10.0	7.42	74	2	35	39-130	
Anthracene	10.0	7.14	71	0	35	39-130	
Benzo[a]anthracene	10.0	7.19	72	0	35	54-135	
Benzo[a]pyrene	10.0	5.01	50	1	35	21-130	
Benzo[b]fluoranthene	10.0	5.92	59	1	35	37-130	
Benzo[g,h,i]perylene	10.0	3.91	39	11	35	26-130	
Benzo[k]fluoranthene	10.0	5.65	57	2	35	38-130	
Chrysene	10.0	6.91	69	5	35	56-130	
Dibenz(a,h)anthracene	10.0	3.54	35	6	35	13-130	
Fluoranthene	10.0	7.94	79	1	35	60-130	
Fluorene	10.0	7.90	79	2	35	55-140	
Indeno[1,2,3-cd]pyrene	10.0	3.59	36	12	35	21-130	
1-Methylnaphthalene	10.0	7.77	78	1	35	49-130	
2-Methylnaphthalene	10.0	7.56	76	0	35	48-130	
Naphthalene	10.0	7.36	74	1	35	54-133	
Phenanthrene	10.0	7.41	74	1	35	60-136	
Pyrene	10.0	7.46	75	0	35	60-138	

# Column to be used to flag recovery and RPD values

FORM III  
GC/MS SEMI VOA MATRIX SPIKE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Solid Level: Low Lab File ID: 1CD03018.D  
 Lab ID: 680-88766-21 MS Client ID: CV0014AB-GS MS

COMPOUND	SPIKE ADDED (ug/Kg)	SAMPLE CONCENTRATION (ug/Kg)	MS CONCENTRATION (ug/Kg)	MS % REC	QC LIMITS REC	#
Acenaphthene	883	530 U	656	74	39-130	
Acenaphthylene	883	34 J	626	67	38-130	
Anthracene	883	45 U	666	75	37-130	
Benzo[a]anthracene	883	200	833	72	40-130	
Benzo[a]pyrene	883	100	650	62	49-130	
Benzo[b]fluoranthene	883	150	738	66	37-130	
Benzo[g,h,i]perylene	883	130	623	56	32-130	
Benzo[k]fluoranthene	883	79	667	67	32-130	
Chrysene	883	210	780	65	41-130	
Dibenz(a,h)anthracene	883	62 J	622	63	27-130	
Fluoranthene	883	210	697	55	40-130	
Fluorene	883	39 J	639	68	40-130	
Indeno[1,2,3-cd]pyrene	883	76 J	532	52	30-130	
1-Methylnaphthalene	883	120 J	653	60	31-130	
2-Methylnaphthalene	883	130 J	653	59	33-130	
Naphthalene	883	100 J	681	66	36-130	
Phenanthrene	883	200	690	55	42-130	
Pyrene	883	210	750	62	44-130	

# Column to be used to flag recovery and RPD values

FORM III  
GC/MS SEMI VOA MATRIX SPIKE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Water Level: Low Lab File ID: 1CD05007.D  
 Lab ID: 680-88766-23 MS Client ID: 032613-RB-shovel MS

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENTRATION (ug/L)	MS CONCENTRATION (ug/L)	MS % REC	QC LIMITS REC	#
Acenaphthene	10.4	2.5 U	4.74	45	55-132	F
Acenaphthylene	10.4	1.3 U	5.18	50	39-130	
Anthracene	10.4	0.25 U	4.97	48	39-130	
Benzo[a]anthracene	10.4	0.25 U	4.72	45	54-135	F
Benzo[a]pyrene	10.4	0.25 U	2.13	20	21-130	F
Benzo[b]fluoranthene	10.4	0.25 U	2.68	26	37-130	F
Benzo[g,h,i]perylene	10.4	0.63 U	1.33	13	26-130	F
Benzo[k]fluoranthene	10.4	0.25 U	2.38	23	38-130	F
Chrysene	10.4	0.25 U	3.55	34	56-130	F
Dibenz(a,h)anthracene	10.4	0.25 U	1.12	11	13-130	F
Fluoranthene	10.4	0.63 U	5.46	52	60-130	F
Fluorene	10.4	2.5 U	4.83	46	55-140	F
Indeno[1,2,3-cd]pyrene	10.4	0.25 U	1.28	12	21-130	F
1-Methylnaphthalene	10.4	2.5 U	5.64	54	49-130	
2-Methylnaphthalene	10.4	2.5 U	4.68	45	48-130	F
Naphthalene	10.4	2.5 U	5.39	52	54-133	F
Phenanthrene	10.4	0.63 U	5.42	52	60-136	F
Pyrene	10.4	0.63 U	5.24	50	60-138	F

# Column to be used to flag recovery and RPD values

FORM III  
GC/MS SEMI VOA MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Matrix: Solid Level: Low Lab File ID: 1CD03019.D  
 Lab ID: 680-88766-21 MSD Client ID: CV0014AB-GS MSD

COMPOUND	SPIKE ADDED (ug/Kg)	MSD CONCENTRATION (ug/Kg)	MSD % REC	% RPD	QC LIMITS		#
					RPD	REC	
Acenaphthene	927	669	72	2	40	39-130	
Acenaphthylene	927	686	70	9	40	38-130	
Anthracene	927	674	73	1	40	37-130	
Benzo[a]anthracene	927	856	71	3	40	40-130	
Benzo[a]pyrene	927	651	59	0	40	49-130	
Benzo[b]fluoranthene	927	710	60	4	40	37-130	
Benzo[g,h,i]perylene	927	618	53	1	40	32-130	
Benzo[k]fluoranthene	927	822	80	21	40	32-130	
Chrysene	927	739	57	5	40	41-130	
Dibenz(a,h)anthracene	927	713	70	14	40	27-130	
Fluoranthene	927	780	62	11	40	40-130	
Fluorene	927	657	67	3	40	40-130	
Indeno[1,2,3-cd]pyrene	927	659	63	21	40	30-130	
1-Methylnaphthalene	927	709	64	8	40	31-130	
2-Methylnaphthalene	927	671	58	3	40	33-130	
Naphthalene	927	648	59	5	40	36-130	
Phenanthrene	927	758	60	9	40	42-130	
Pyrene	927	878	72	16	40	44-130	

# Column to be used to flag recovery and RPD values

FORM IV  
GC/MS SEMI VOA METHOD BLANK SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
SDG No.: 68088766-2  
Lab File ID: 1DD04022.D Lab Sample ID: MB 660-136013/1-A  
Matrix: Water Date Extracted: 04/02/2013 08:08  
Instrument ID: BSMD5973 Date Analyzed: 04/04/2013 19:51  
Level: (Low/Med) Low

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES:

CLIENT SAMPLE ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
	LCS 660-136013/2-A	1DD04023.D	04/04/2013 20:13
	LCSD 660-136013/3-A	1DD04024.D	04/04/2013 20:36
032613-RB-shovel	680-88766-23	1CD05006.D	04/05/2013 12:54
032613-RB-shovel MS	680-88766-23 MS	1CD05007.D	04/05/2013 13:12

FORM IV  
GC/MS SEMI VOA METHOD BLANK SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
SDG No.: 68088766-2  
Lab File ID: 1CD03015.D Lab Sample ID: MB 660-136063/1-A  
Matrix: Solid Date Extracted: 04/02/2013 11:33  
Instrument ID: BSMC5973 Date Analyzed: 04/03/2013 15:34  
Level: (Low/Med) Low

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES:

CLIENT SAMPLE ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
	LCS 660-136063/2-A	1CD03016.D	04/03/2013 15:52
CV0014AB-GS	680-88766-21	1CD03017.D	04/03/2013 16:10
CV0014AB-GS MS	680-88766-21 MS	1CD03018.D	04/03/2013 16:29
CV0014AB-GS MSD	680-88766-21 MSD	1CD03019.D	04/03/2013 16:47



FORM V  
GC/MS SEMI VOA INSTRUMENT PERFORMANCE CHECK  
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab File ID: 1CD02002.D DFTPP Injection Date: 04/02/2013  
 Instrument ID: BSMC5973 DFTPP Injection Time: 11:31  
 Analysis Batch No.: 136048

M/E	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0 % of mass 198	34.9
68	Less than 2.0 % of mass 69	0.8 (1.6)1
69	Mass 69 relative abundance	49.9
70	Less than 2.0 % of mass 69	0.4 (0.9)1
127	10.0 - 80.0 % of mass 198	42.2
197	Less than 2.0 % of mass 198	0.4
198	Base Peak, 100% relative abundance	100.0
199	5.0 - 9.0 % of mass 198	7.6
275	10.0 - 60.0 % of mass 198	21.5
365	Greater than 1.0 % of mass 198	3.4
441	Present but less than mass 443	10.2
442	Greater than 50.0 % of mass 198	56.7
443	15.0 - 24.0 % of mass 442	11.0 (19.4)2

1-Value is % mass 69

2-Value is % mass 442

THIS CHECK APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS AND STANDARDS:

CLIENT SAMPLE ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
	IC 660-136048/5	1CD02005.D	04/02/2013	13:26
	IC 660-136048/6	1CD02006.D	04/02/2013	13:44
	IC 660-136048/7	1CD02007.D	04/02/2013	14:02
	IC 660-136048/8	1CD02008.D	04/02/2013	14:20
	ICIS 660-136048/9	1CD02009.D	04/02/2013	14:39
	IC 660-136048/10	1CD02010.D	04/02/2013	14:57
	IC 660-136048/11	1CD02011.D	04/02/2013	15:15
	ICV 660-136048/12	1CD02012.D	04/02/2013	15:34

FORM V  
GC/MS SEMI VOA INSTRUMENT PERFORMANCE CHECK  
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab File ID: 1CD03002.D DFTPP Injection Date: 04/03/2013  
 Instrument ID: BSMC5973 DFTPP Injection Time: 11:28  
 Analysis Batch No.: 136081

M/E	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0 % of mass 198	42.7
68	Less than 2.0 % of mass 69	0.6 (1.1)1
69	Mass 69 relative abundance	49.7
70	Less than 2.0 % of mass 69	0.0 (0.0)1
127	10.0 - 80.0 % of mass 198	48.1
197	Less than 2.0 % of mass 198	0.0
198	Base Peak, 100% relative abundance	100.0
199	5.0 - 9.0 % of mass 198	6.2
275	10.0 - 60.0 % of mass 198	19.7
365	Greater than 1.0 % of mass 198	2.4
441	Present but less than mass 443	6.9
442	Greater than 50.0 % of mass 198	61.0
443	15.0 - 24.0 % of mass 442	10.2 (16.7)2

1-Value is % mass 69

2-Value is % mass 442

THIS CHECK APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS AND STANDARDS:

CLIENT SAMPLE ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
	CCVIS 660-136081/3	1CD03003.D	04/03/2013	11:45
	MB 660-136063/1-A	1CD03015.D	04/03/2013	15:34
	LCS 660-136063/2-A	1CD03016.D	04/03/2013	15:52
CV0014AB-GS	680-88766-21	1CD03017.D	04/03/2013	16:10
CV0014AB-GS MS	680-88766-21 MS	1CD03018.D	04/03/2013	16:29
CV0014AB-GS MSD	680-88766-21 MSD	1CD03019.D	04/03/2013	16:47

FORM V  
GC/MS SEMI VOA INSTRUMENT PERFORMANCE CHECK  
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab File ID: 1CD05003.D DFTPP Injection Date: 04/05/2013  
 Instrument ID: BSMC5973 DFTPP Injection Time: 11:57  
 Analysis Batch No.: 136171

M/E	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0 % of mass 198	41.6
68	Less than 2.0 % of mass 69	0.8 (1.4) 1
69	Mass 69 relative abundance	55.3
70	Less than 2.0 % of mass 69	0.3 (0.6) 1
127	10.0 - 80.0 % of mass 198	49.0
197	Less than 2.0 % of mass 198	0.6
198	Base Peak, 100% relative abundance	100.0
199	5.0 - 9.0 % of mass 198	6.7
275	10.0 - 60.0 % of mass 198	19.3
365	Greater than 1.0 % of mass 198	3.0
441	Present but less than mass 443	7.6
442	Greater than 50.0 % of mass 198	55.6
443	15.0 - 24.0 % of mass 442	11.7 (21.0) 2

1-Value is % mass 69

2-Value is % mass 442

THIS CHECK APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS AND STANDARDS:

CLIENT SAMPLE ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
	CCVIS 660-136171/4	1CD05004.D	04/05/2013	12:15
032613-RB-shovel	680-88766-23	1CD05006.D	04/05/2013	12:54
032613-RB-shovel MS	680-88766-23 MS	1CD05007.D	04/05/2013	13:12

FORM V  
GC/MS SEMI VOA INSTRUMENT PERFORMANCE CHECK  
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab File ID: 1DD04003.D DFTPP Injection Date: 04/04/2013  
 Instrument ID: BSMD5973 DFTPP Injection Time: 12:15  
 Analysis Batch No.: 136164

M/E	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0 % of mass 198	44.9
68	Less than 2.0 % of mass 69	0.0 (0.0) 1
69	Mass 69 relative abundance	45.4
70	Less than 2.0 % of mass 69	0.2 (0.3) 1
127	10.0 - 80.0 % of mass 198	50.5
197	Less than 2.0 % of mass 198	0.0
198	Base Peak, 100% relative abundance	100.0
199	5.0 - 9.0 % of mass 198	6.9
275	10.0 - 60.0 % of mass 198	26.7
365	Greater than 1.0 % of mass 198	3.1
441	Present but less than mass 443	3.3
442	Greater than 50.0 % of mass 198	67.1
443	15.0 - 24.0 % of mass 442	13.9 (20.6) 2

1-Value is % mass 69

2-Value is % mass 442

THIS CHECK APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS AND STANDARDS:

CLIENT SAMPLE ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
	IC 660-136164/15	1DD04007.D	04/04/2013	13:49
	IC 660-136164/16	1DD04008.D	04/04/2013	14:11
	IC 660-136164/17	1DD04009.D	04/04/2013	14:34
	IC 660-136164/18	1DD04010.D	04/04/2013	14:57
	ICIS 660-136164/19	1DD04011.D	04/04/2013	15:19
	IC 660-136164/20	1DD04012.D	04/04/2013	15:42
	IC 660-136164/21	1DD04013.D	04/04/2013	16:04
	ICV 660-136164/22	1DD04014.D	04/04/2013	16:27
	MB 660-136013/1-A	1DD04022.D	04/04/2013	19:51
	LCS 660-136013/2-A	1DD04023.D	04/04/2013	20:13
	LCSD 660-136013/3-A	1DD04024.D	04/04/2013	20:36

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: ICIS 660-136048/9 Date Analyzed: 04/02/2013 14:39  
 Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1CD02009.D Heated Purge: (Y/N) N  
 Calibration ID: 2859

	NPT		ANT		PHN		
	AREA #	RT #	AREA #	RT #	AREA #	RT #	
INITIAL CALIBRATION MID-POINT	501011	3.71	361349	4.80	702974	5.75	
UPPER LIMIT	1002022	4.21	722698	5.30	1405948	6.25	
LOWER LIMIT	250506	3.21	180675	4.30	351487	5.25	
LAB SAMPLE ID	CLIENT SAMPLE ID						
ICV 660-136048/12		649122	3.71	500935	4.80	955391	5.75

NPT = Naphthalene-d8  
 ANT = Acenaphthene-d10  
 PHN = Phenanthrene-d10

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: ICIS 660-136048/9 Date Analyzed: 04/02/2013 14:39  
 Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1CD02009.D Heated Purge: (Y/N) N  
 Calibration ID: 2859

	CRY		PRY		AREA #	RT #
	AREA #	RT #	AREA #	RT #		
INITIAL CALIBRATION MID-POINT	875378	7.69	942955	8.86		
UPPER LIMIT	1750756	8.19	1885910	9.36		
LOWER LIMIT	437689	7.19	471478	8.36		
LAB SAMPLE ID	CLIENT SAMPLE ID					
ICV 660-136048/12	1249690	7.69	1306409	8.86		

CRY = Chrysene-d12  
 PRY = Perylene-d12

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: CCVIS 660-136081/3 Date Analyzed: 04/03/2013 11:45  
 Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1CD03003.D Heated Purge: (Y/N) N  
 Calibration ID: 2859

	NPT		ANT		PHN			
	AREA #	RT #	AREA #	RT #	AREA #	RT #		
12/24 HOUR STD	500765	3.70	364027	4.79	687020	5.74		
UPPER LIMIT	1001530	4.20	728054	5.29	1374040	6.24		
LOWER LIMIT	250383	3.20	182014	4.29	343510	5.24		
LAB SAMPLE ID	CLIENT SAMPLE ID							
MB 660-136063/1-A			646271	3.71	483289	4.80	879169	5.75
LCS 660-136063/2-A			631468	3.71	487717	4.79	877192	5.74
680-88766-21	CV0014AB-GS		582428	3.71	437660	4.79	782573	5.74
680-88766-21 MS	CV0014AB-GS MS		604618	3.71	447317	4.79	819020	5.74
680-88766-21 MSD	CV0014AB-GS MSD		599290	3.71	437990	4.79	801261	5.74

NPT = Naphthalene-d8  
 ANT = Acenaphthene-d10  
 PHN = Phenanthrene-d10

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: CCVIS 660-136081/3 Date Analyzed: 04/03/2013 11:45  
 Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1CD03003.D Heated Purge: (Y/N) N  
 Calibration ID: 2859

	CRY		PRY		AREA #	RT #
	AREA #	RT #	AREA #	RT #		
12/24 HOUR STD	857573	7.68	866012	8.85		
UPPER LIMIT	1715146	8.18	1732024	9.35		
LOWER LIMIT	428787	7.18	433006	8.35		
LAB SAMPLE ID	CLIENT SAMPLE ID					
MB 660-136063/1-A		990236	7.68	958431	8.86	
LCS 660-136063/2-A		995572	7.68	964341	8.85	
680-88766-21	CV0014AB-GS	873422	7.68	855093	8.86	
680-88766-21 MS	CV0014AB-GS MS	882360	7.68	885344	8.85	
680-88766-21 MSD	CV0014AB-GS MSD	885431	7.68	855561	8.85	

CRY = Chrysene-d12  
 PRY = Perylene-d12

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits



FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: CCVIS 660-136171/4 Date Analyzed: 04/05/2013 12:15  
 Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1CD05004.D Heated Purge: (Y/N) N  
 Calibration ID: 2859

	NPT		ANT		PHN			
	AREA #	RT #	AREA #	RT #	AREA #	RT #		
12/24 HOUR STD	392528	3.69	289150	4.78	539578	5.72		
UPPER LIMIT	785056	4.19	578300	5.28	1079156	6.22		
LOWER LIMIT	196264	3.19	144575	4.28	269789	5.22		
LAB SAMPLE ID	CLIENT SAMPLE ID							
680-88766-23	032613-RB-shovel		397216	3.69	288931	4.77	552053	5.72
680-88766-23 MS	032613-RB-shovel MS		411549	3.69	313598	4.77	608819	5.72

NPT = Naphthalene-d8  
 ANT = Acenaphthene-d10  
 PHN = Phenanthrene-d10

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: CCVIS 660-136171/4 Date Analyzed: 04/05/2013 12:15  
 Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1CD05004.D Heated Purge: (Y/N) N  
 Calibration ID: 2859

	CRY		PRY		AREA #	RT #
	AREA #	RT #	AREA #	RT #		
12/24 HOUR STD	739705	7.66	746693	8.83		
UPPER LIMIT	1479410	8.16	1493386	9.33		
LOWER LIMIT	369853	7.16	373347	8.33		
LAB SAMPLE ID	CLIENT SAMPLE ID					
680-88766-23	032613-RB-shovel		730677	7.66	752862	8.82
680-88766-23 MS	032613-RB-shovel MS		752169	7.66	802662	8.82

CRY = Chrysene-d12  
 PRY = Perylene-d12

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: ICIS 660-136164/19 Date Analyzed: 04/04/2013 15:19  
 Instrument ID: BSMD5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1DD04011.D Heated Purge: (Y/N) N  
 Calibration ID: 2874

	NPT		ANT		PHN	
	AREA #	RT #	AREA #	RT #	AREA #	RT #
INITIAL CALIBRATION MID-POINT	2475113	6.09	1466924	7.77	2428512	9.03
UPPER LIMIT	4950226	6.59	2933848	8.27	4857024	9.53
LOWER LIMIT	1237557	5.59	733462	7.27	1214256	8.53
LAB SAMPLE ID	CLIENT SAMPLE ID					
ICV 660-136164/22	3619899	6.10	2333423	7.77	3845474	9.03
MB 660-136013/1-A	2711590	6.09	1656380	7.77	2762536	9.03
LCS 660-136013/2-A	2684140	6.09	1602441	7.77	2731968	9.03
LCSD 660-136013/3-A	2700850	6.09	1619728	7.77	2719610	9.03

NPT = Naphthalene-d8  
 ANT = Acenaphthene-d10  
 PHN = Phenanthrene-d10

Area Limit = 50%-200% of internal standard area  
 RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM VIII  
GC/MS SEMI VOA INTERNAL STANDARD AREA AND RETENTION TIME SUMMARY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Sample No.: ICIS 660-136164/19 Date Analyzed: 04/04/2013 15:19  
 Instrument ID: BSMD5973 GC Column: DB-5MS ID: 250 (um)  
 Lab File ID (Standard): 1DD04011.D Heated Purge: (Y/N) N  
 Calibration ID: 2874

	CRY		PRY		AREA #	RT #
	AREA #	RT #	AREA #	RT #		
INITIAL CALIBRATION MID-POINT	2464730	11.34	2515643	13.17		
UPPER LIMIT	4929460	11.84	5031286	13.67		
LOWER LIMIT	1232365	10.84	1257822	12.67		
LAB SAMPLE ID	CLIENT SAMPLE ID					
ICV 660-136164/22		3963674	11.35	3958481	13.18	
MB 660-136013/1-A		2767376	11.34	2802105	13.17	
LCS 660-136013/2-A		2730920	11.34	2751843	13.17	
LCSD 660-136013/3-A		2765140	11.34	2782071	13.17	

CRY = Chrysene-d12

PRY = Perylene-d12

Area Limit = 50%-200% of internal standard area

RT Limit = ± 0.5 minutes of internal standard RT

# Column used to flag values outside QC limits

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: CV0014AB-GS Lab Sample ID: 680-88766-21  
 Matrix: Solid Lab File ID: 1CD03017.D  
 Analysis Method: 8270C LL Date Collected: 03/25/2013 15:18  
 Extract. Method: 3546 Date Extracted: 04/02/2013 11:33  
 Sample wt/vol: 15.25(g) Date Analyzed: 04/03/2013 16:10  
 Con. Extract Vol.: 1(mL) Dilution Factor: 4  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: 26.1 GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136081 Units: ug/Kg

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	530	U	530	110
208-96-8	Acenaphthylene	34	J	210	27
120-12-7	Anthracene	45	U	45	22
56-55-3	Benzo[a]anthracene	200		43	21
50-32-8	Benzo[a]pyrene	100		55	28
205-99-2	Benzo[b]fluoranthene	150		65	32
191-24-2	Benzo[g,h,i]perylene	130		110	23
207-08-9	Benzo[k]fluoranthene	79		43	19
218-01-9	Chrysene	210		48	24
53-70-3	Dibenz(a,h)anthracene	62	J	110	22
206-44-0	Fluoranthene	210		110	21
86-73-7	Fluorene	39	J	110	22
193-39-5	Indeno[1,2,3-cd]pyrene	76	J	110	38
90-12-0	1-Methylnaphthalene	120	J	210	23
91-57-6	2-Methylnaphthalene	130	J	210	38
91-20-3	Naphthalene	100	J	210	23
85-01-8	Phenanthrene	200		43	21
129-00-0	Pyrene	210		110	20

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	95		30-130

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03017.D  
 Lab Smp Id: 680-88766-A-21-D Client Smp ID: CV0014AB-GS  
 Inj Date : 03-APR-2013 16:10  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : 680-88766-a-21-d  
 Misc Info : 680-88766-A-21-D  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\a-bFASTPAHi-m.m  
 Meth Date : 03-Apr-2013 11:59 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 17  
 Dil Factor: 4.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula:

$$\text{Amt} * \text{DF} * 1/\text{Vi} * \text{Vt}/\text{Ws} * 100/(100 - \text{M}) * \text{A} * \text{B} * \text{C} * \text{D} * \text{GPC} * \text{CpndVariable}$$

Name	Value	Description
DF	4.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Ws	15.250	Weight Extracted
M	26.115	% Moisture
A	1000.000	uL to mL conversion
B	1000.000	g to kg conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1 if no conv)
GPC	1.000	GPC FACTOR
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
							ON-COLUMN	FINAL
	MASS						(ug/ml)	(ug/Kg)
* 1 Naphthalene-d8	136		3.710	3.704	(1.000)	582428	40.0000	
* 6 Acenaphthene-d10	164		4.792	4.792	(1.000)	437660	40.0000	
* 10 Phenanthrene-d10	188		5.739	5.739	(1.000)	782573	40.0000	
\$ 14 o-Terphenyl	230		5.992	5.992	(1.044)	21143	2.38061	845.1223
* 18 Chrysene-d12	240		7.680	7.680	(1.000)	873422	40.0000	
* 23 Perylene-d12	264		8.856	8.851	(1.000)	855093	40.0000	
2 Naphthalene	128		3.721	3.722	(1.003)	4269	0.28537	101.3070(Q)
3 2-Methylnaphthalene	142		4.145	4.145	(1.117)	3722	0.36550	129.7549
4 1-Methylnaphthalene	142		4.204	4.210	(1.133)	3101	0.33843	120.1436
5 Acenaphthylene	152		4.710	4.704	(0.983)	1749	0.09656	34.2779(Q)
9 Fluorene	166		5.133	5.133	(1.071)	1631	0.10905	38.7139
11 Phenanthrene	178		5.757	5.757	(1.003)	13094	0.57450	203.9476
13 Carbazole	167		5.904	5.898	(1.029)	1637	0.08270	29.3583(Q)
15 Fluoranthene	202		6.586	6.592	(1.148)	14763	0.58651	208.2114

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/ml)	FINAL (ug/Kg)
16 Pyrene	202	6.757	6.757 (0.880)		14060	0.58112	206.3009
17 Benzo(a)anthracene	228	7.674	7.668 (0.999)		10878	0.56546	200.7412
19 Chrysene	228	7.698	7.698 (1.002)		14637	0.58810	208.7764
20 Benzo(b)fluoranthene	252	8.515	8.509 (0.961)		10463	0.43282	153.6511
21 Benzo(k)fluoranthene	252	8.539	8.533 (0.964)		5203	0.22253	78.9998(Q)
22 Benzo(a)pyrene	252	8.798	8.798 (0.993)		6701	0.29443	104.5224(Q)
24 Indeno(1,2,3-cd)pyrene	276	9.998	9.992 (1.129)		4605	0.21303	75.6245(M)
25 Dibenzo(a,h)anthracene	278	10.015	10.009 (1.131)		3484	0.17447	61.9370(M)
26 Benzo(g,h,i)perylene	276	10.345	10.339 (1.168)		7927	0.35929	127.5493(M)

QC Flag Legend

- Q - Qualifier signal failed the ratio test.
- M - Compound response manually integrated.

Data File: 1CD03017.D

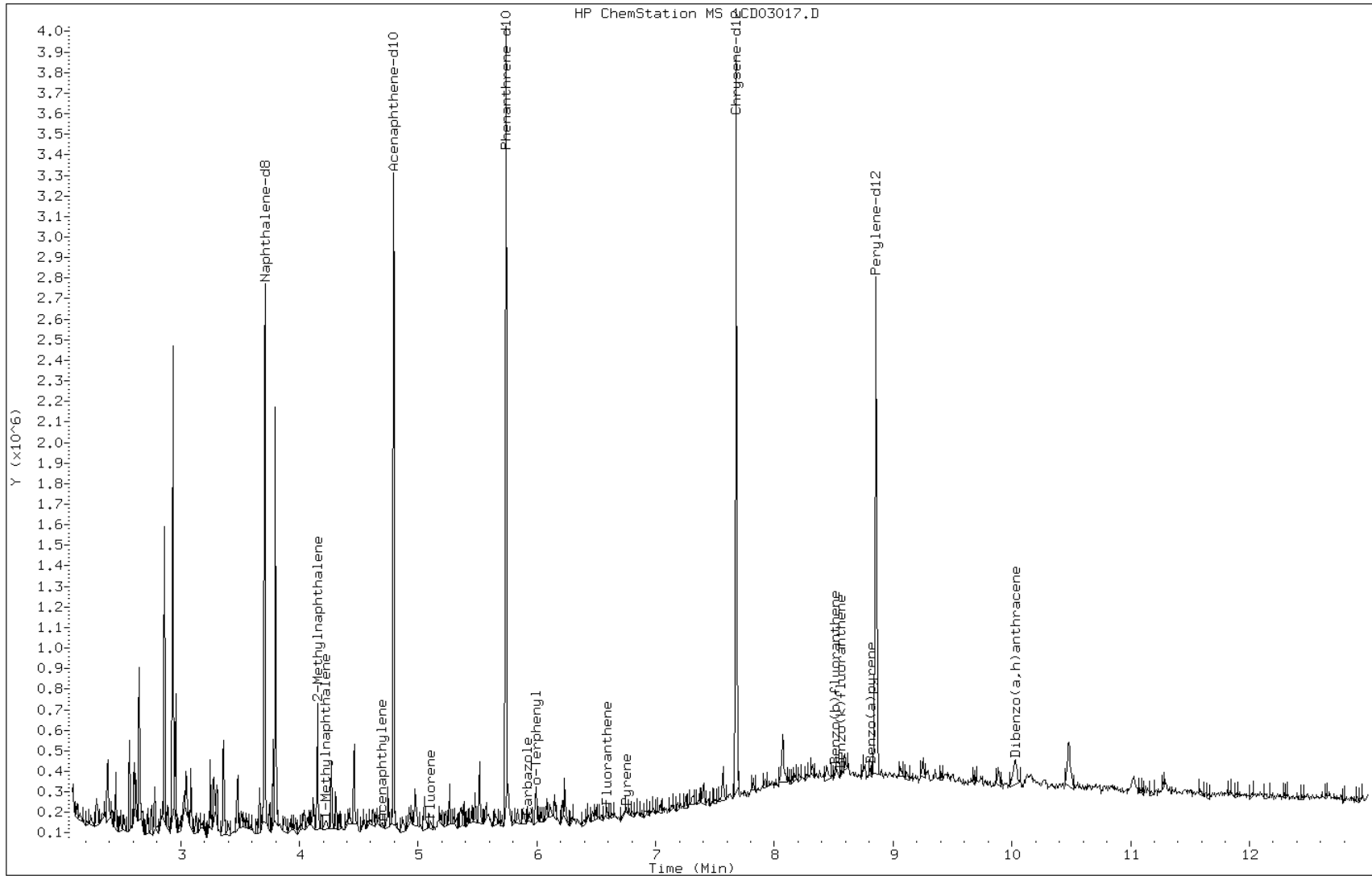
Date: 03-APR-2013 16:10

Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC





Data File: 1CD03017.D

Date: 03-APR-2013 16:10

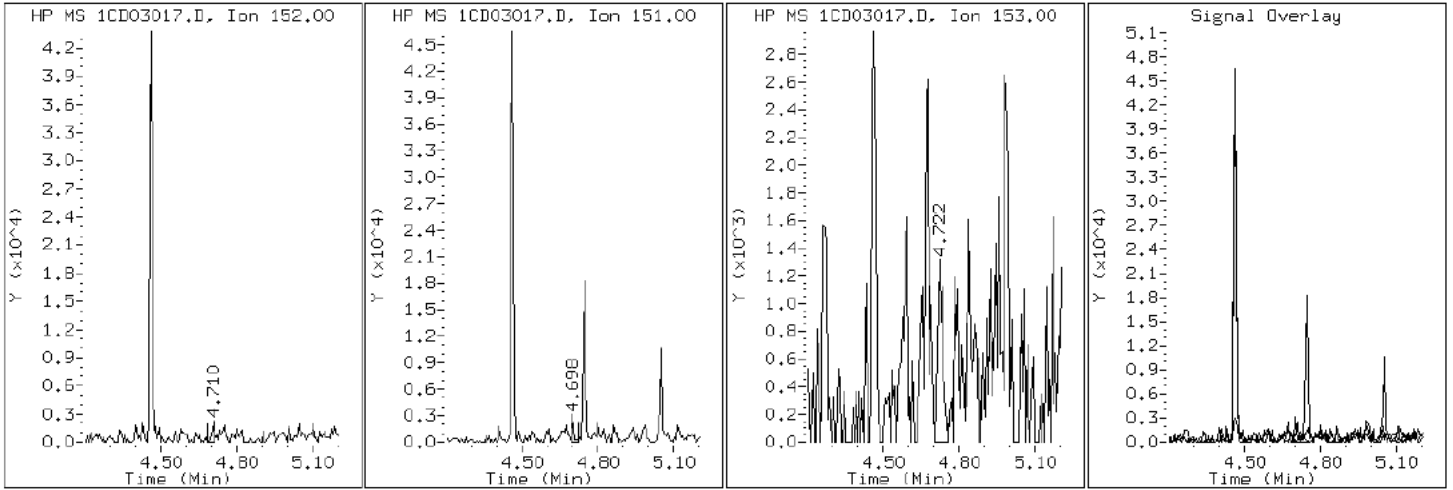
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

5 Acenaphthylene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

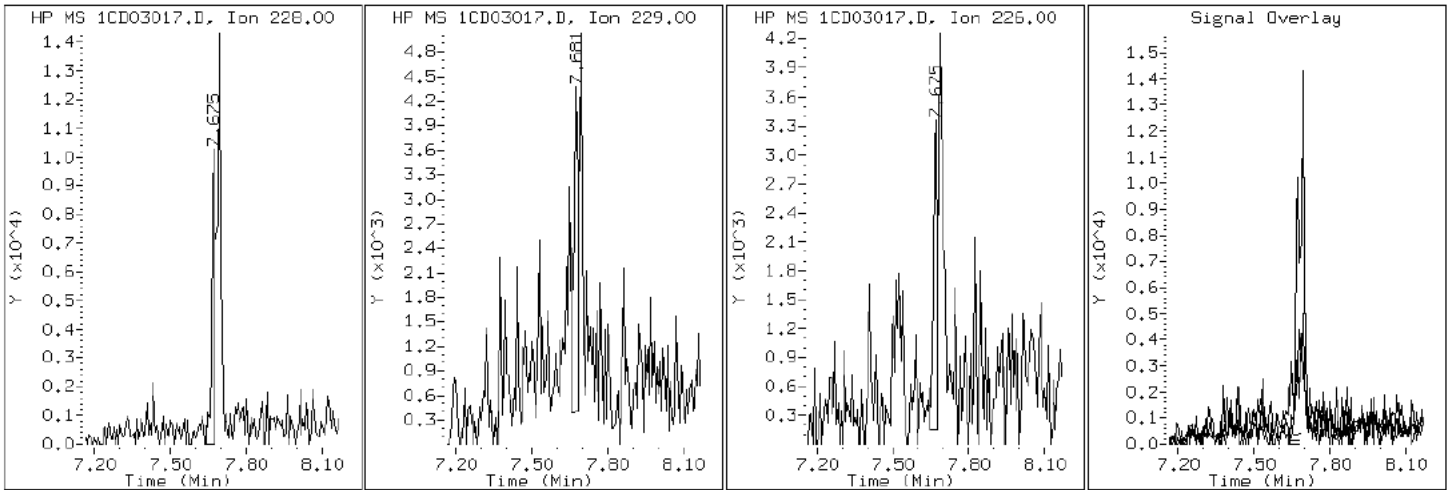
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

17 Benzo(a)anthracene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

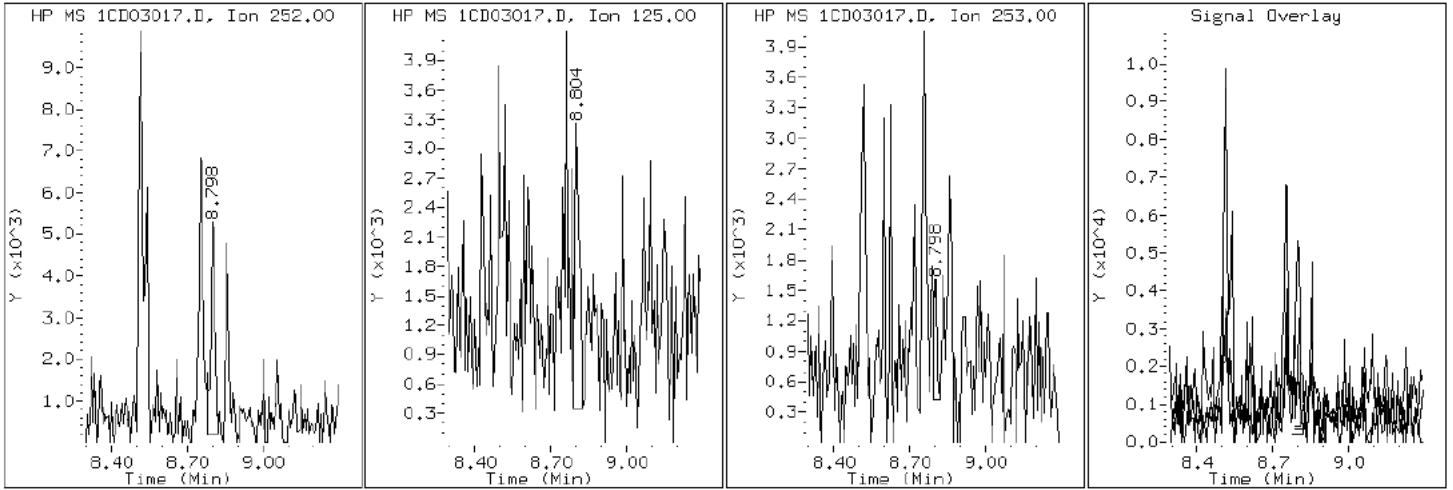
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

22 Benzo(a)pyrene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

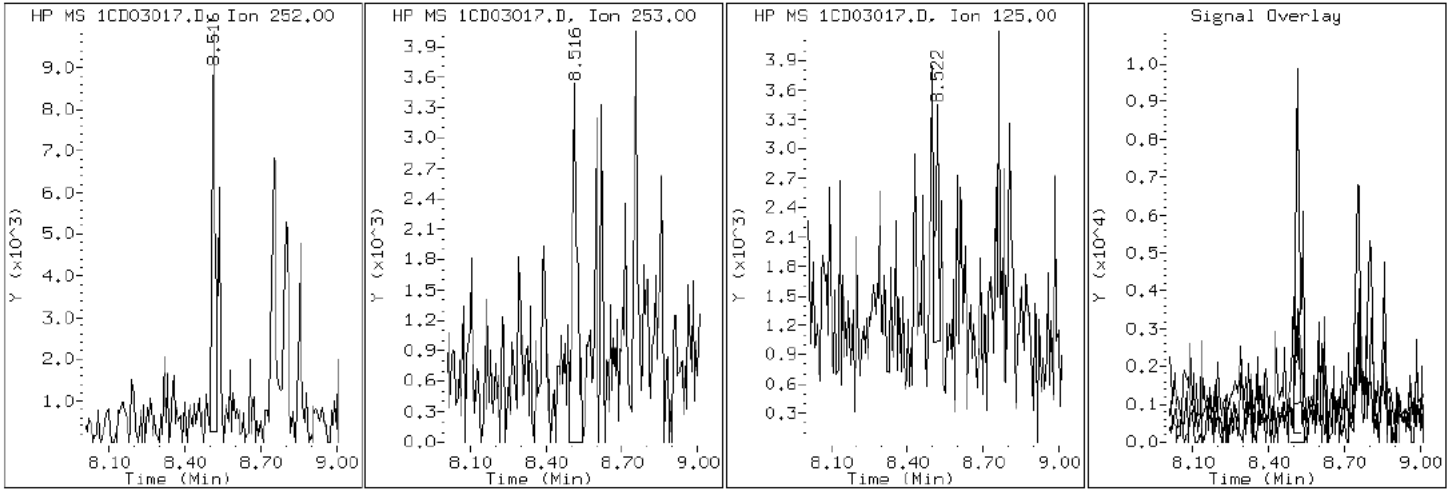
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

20 Benzo (b) fluoranthene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

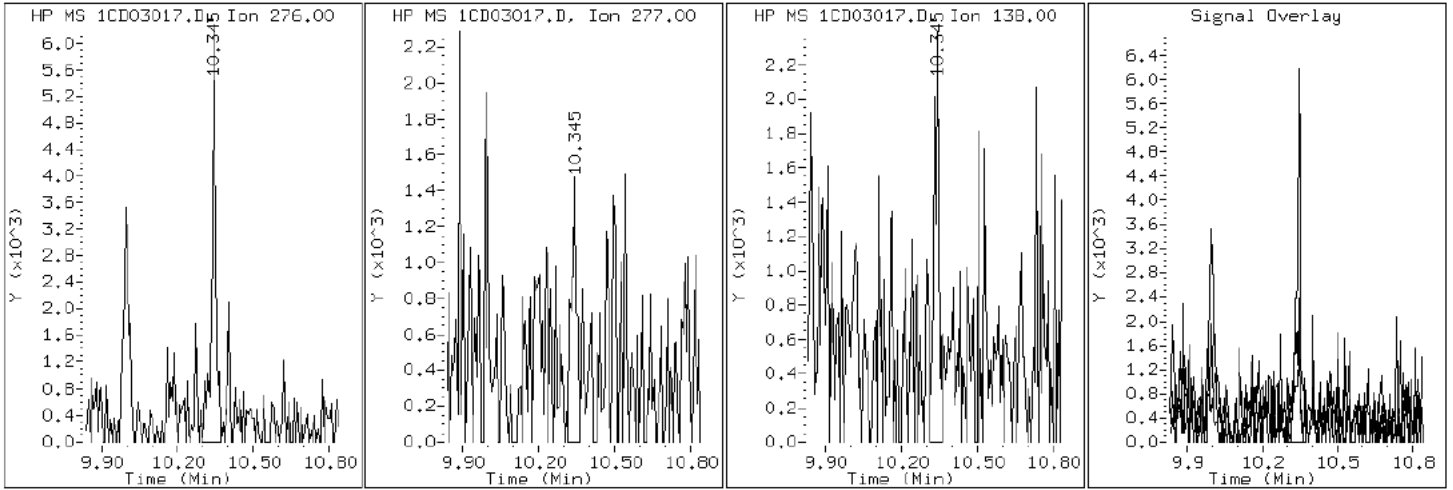
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

26 Benzo(g,h,i)perylene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

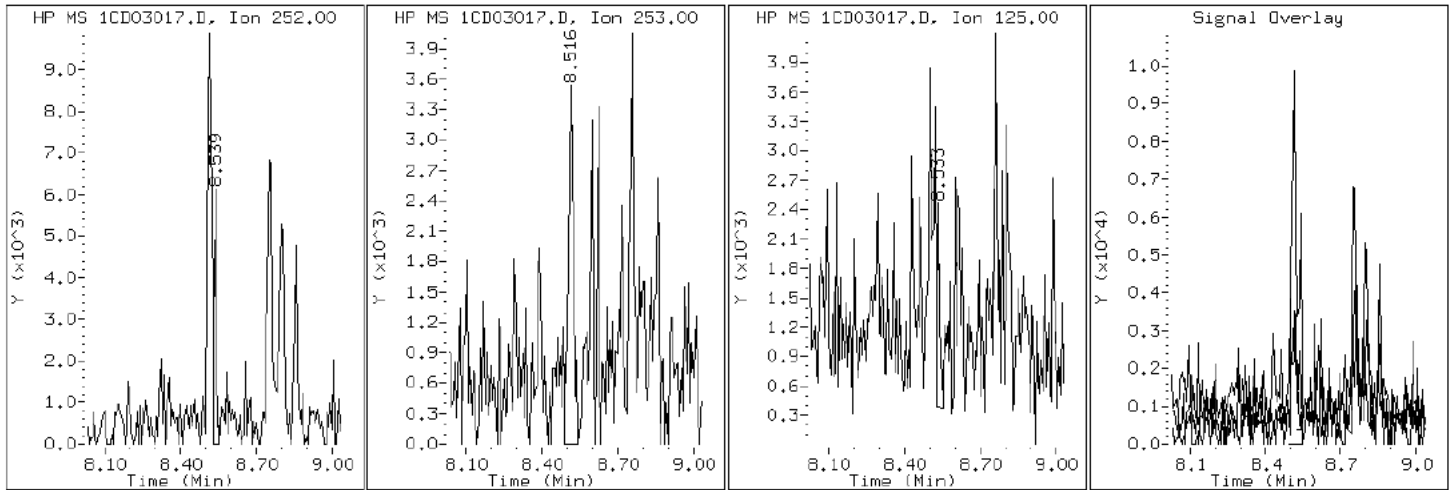
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

21 Benzo(k)fluoranthene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

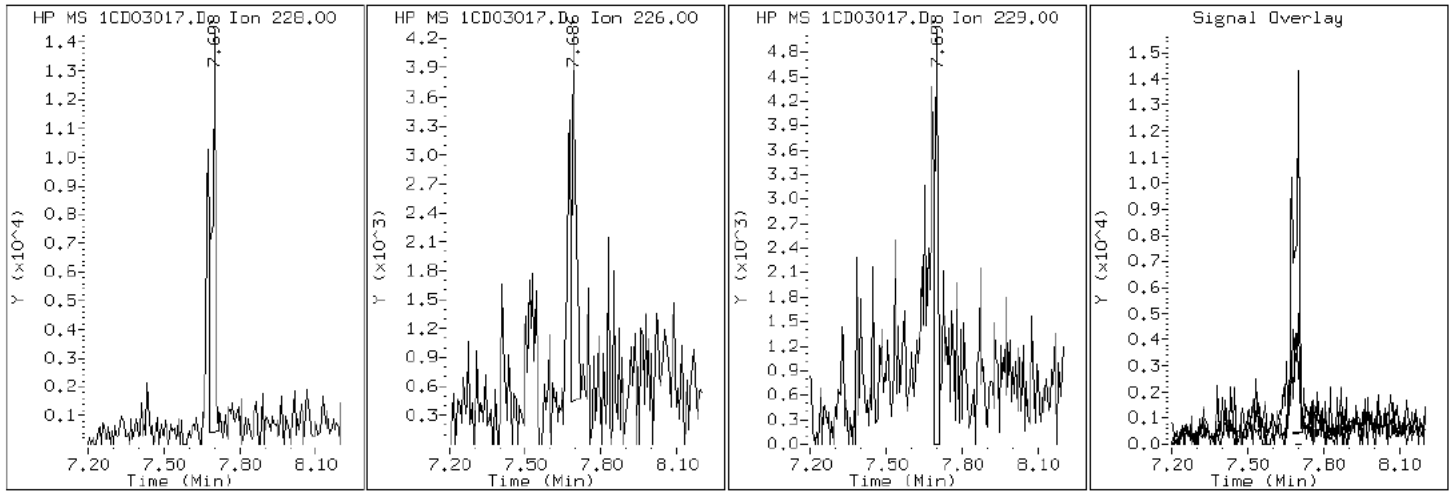
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

19 Chrysene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

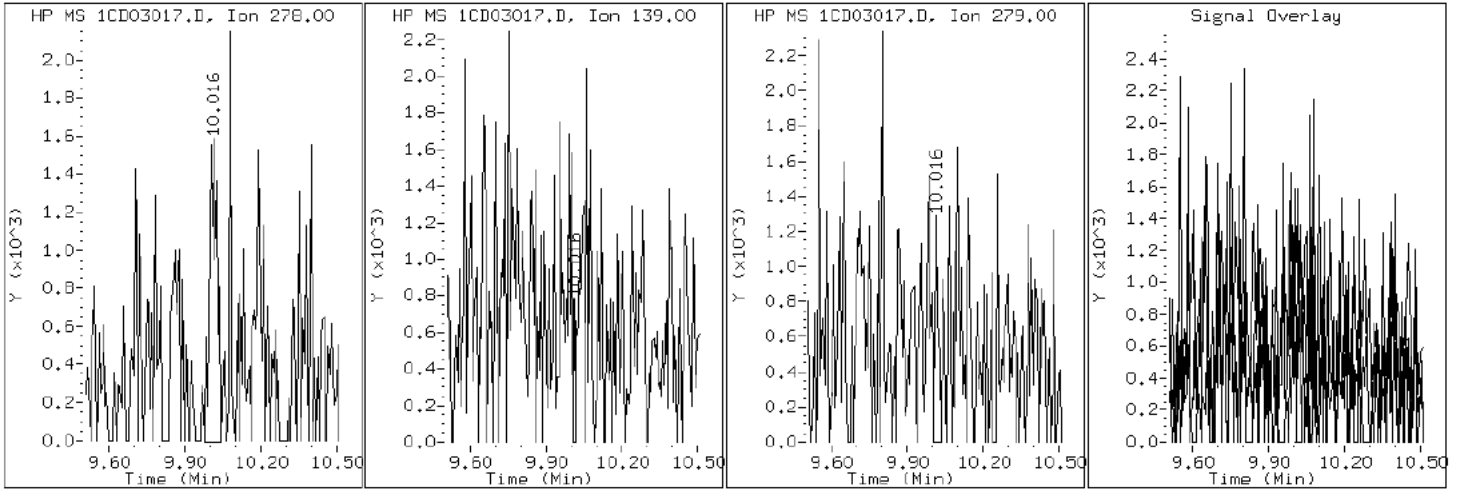
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

25 Dibenzo (a,h) anthracene





Data File: 1CD03017.D

Date: 03-APR-2013 16:10

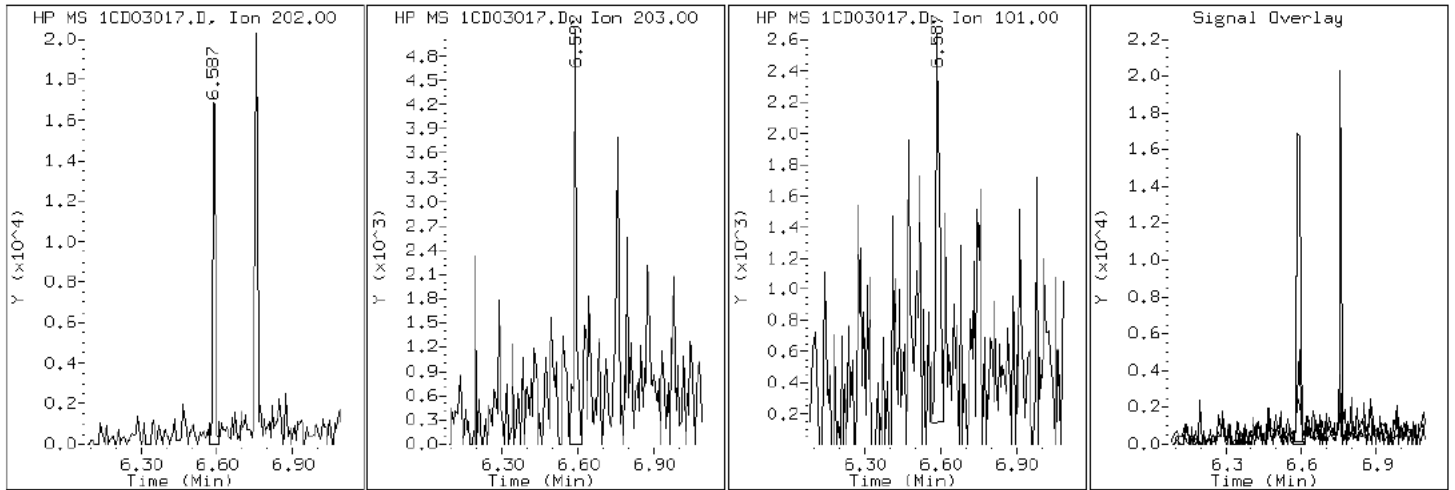
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

15 Fluoranthene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

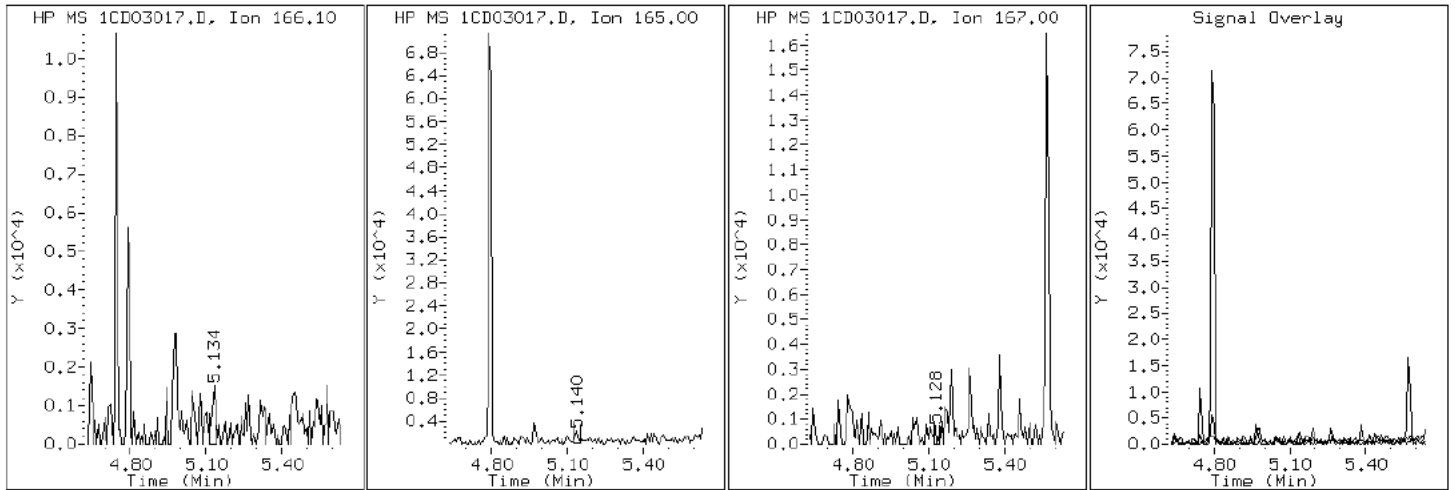
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

9 Fluorene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

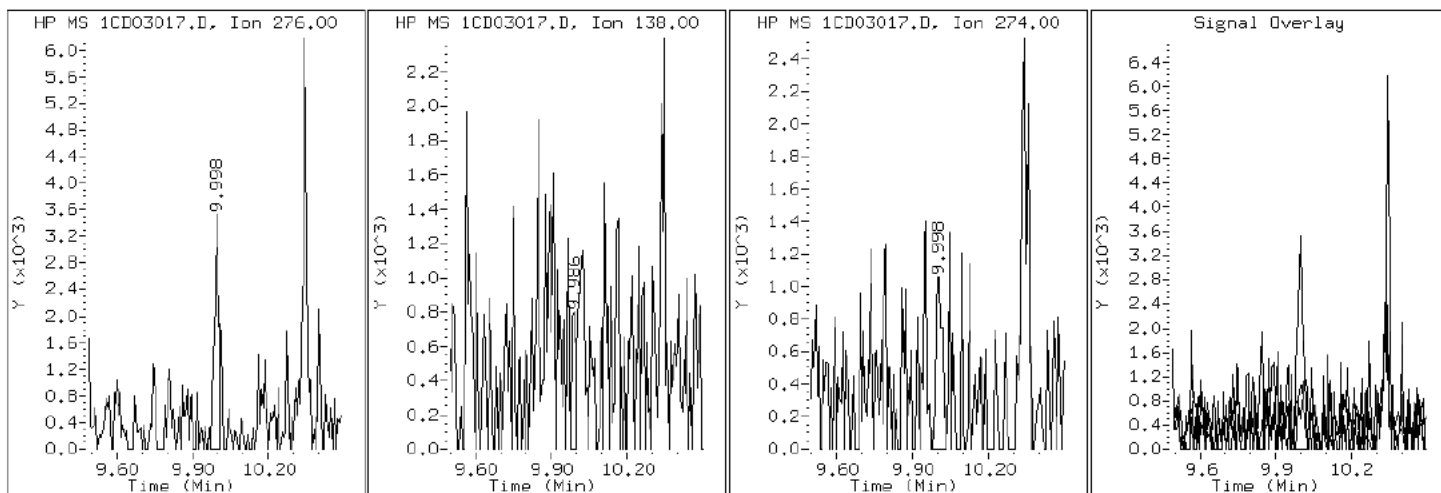
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

24 Indeno(1,2,3-cd)pyrene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

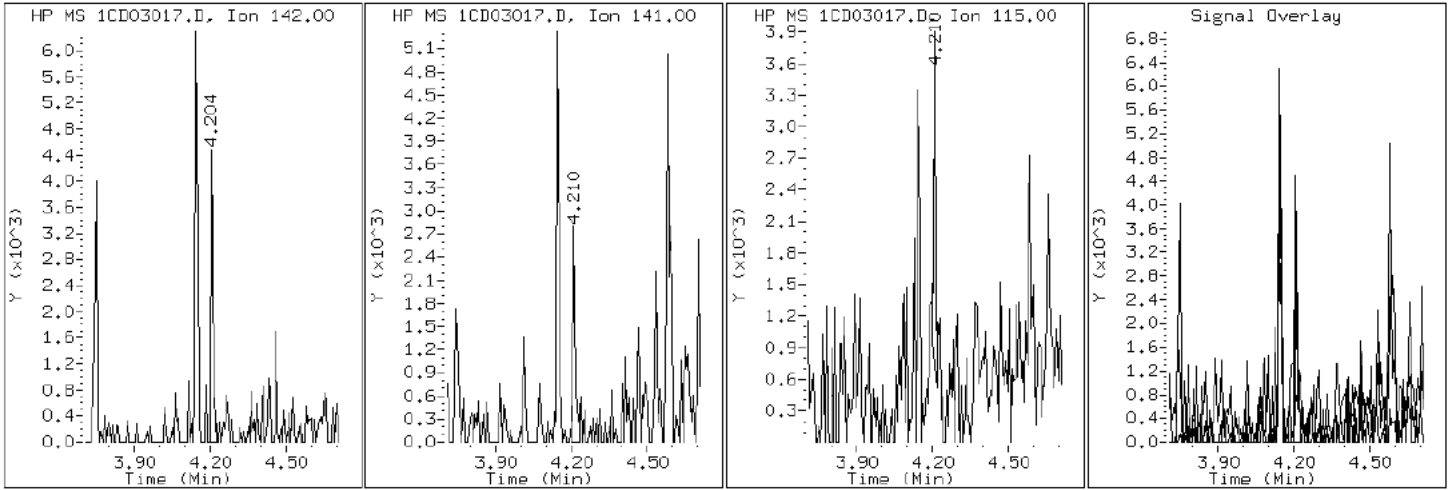
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

4 1-Methylnaphthalene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

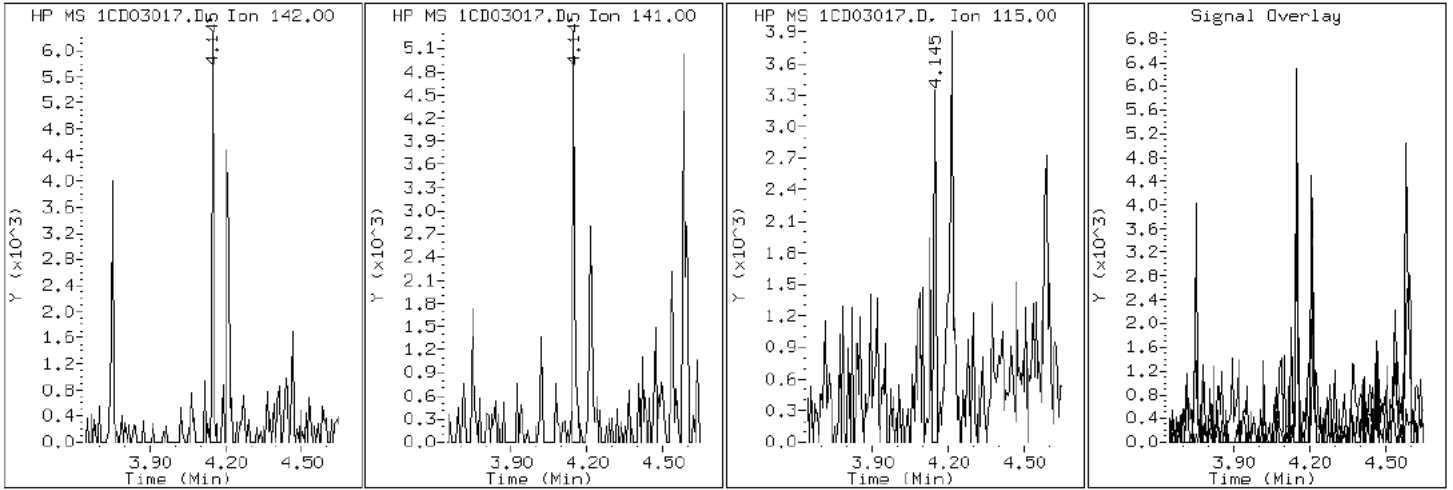
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

3 2-Methylnaphthalene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

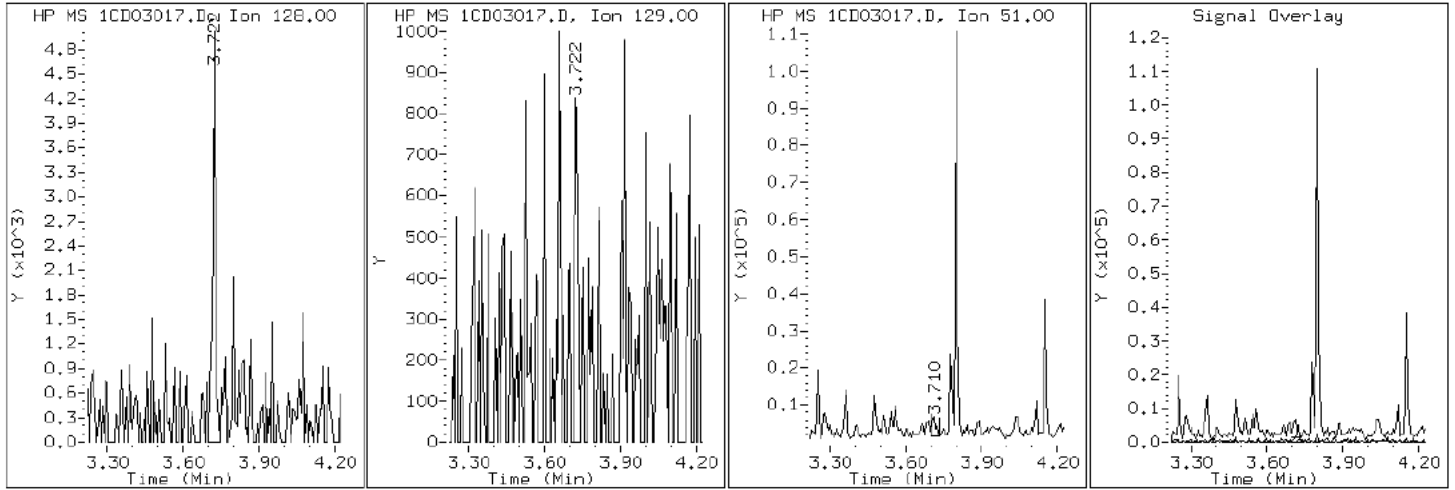
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

2 Naphthalene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

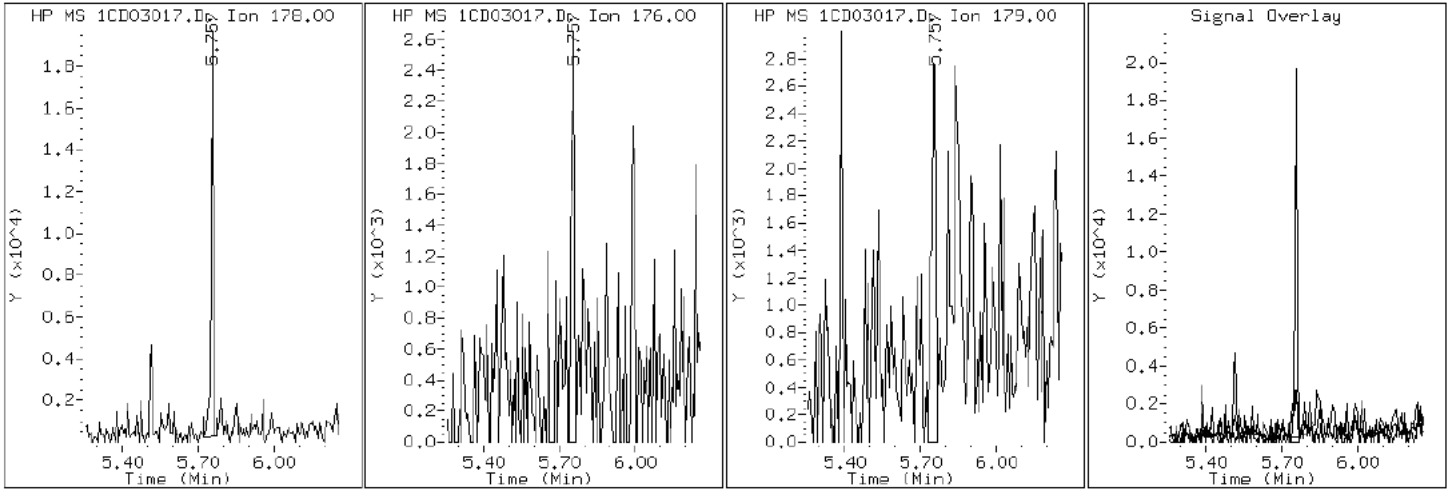
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

11 Phenanthrene



Data File: 1CD03017.D

Date: 03-APR-2013 16:10

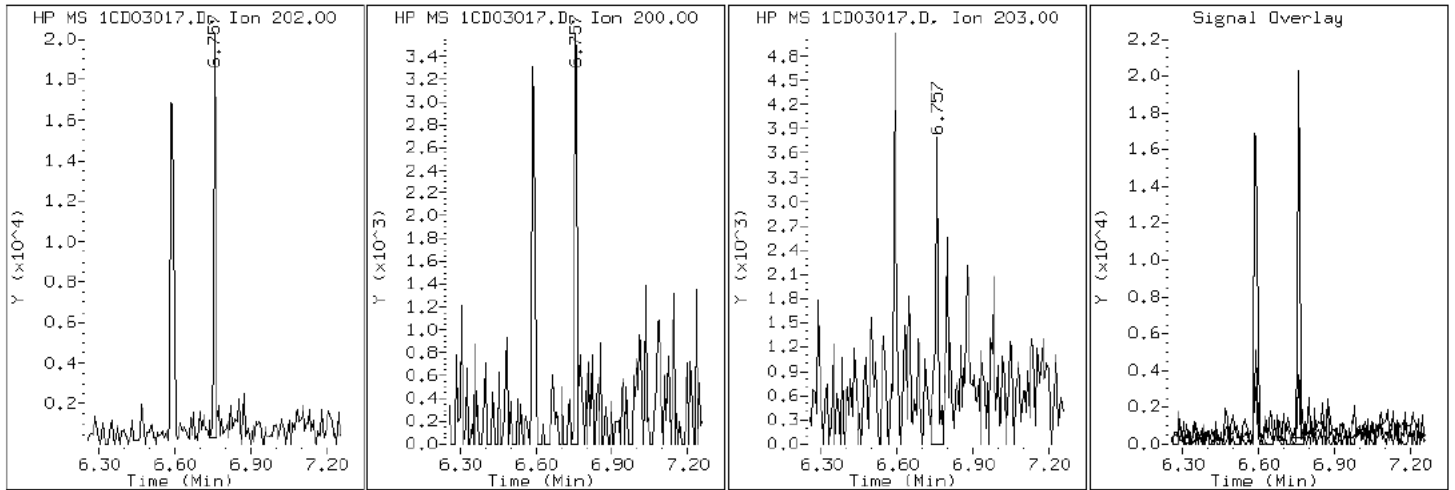
Client ID: CV0014AB-GS

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-d

Operator: SCC

16 Pyrene



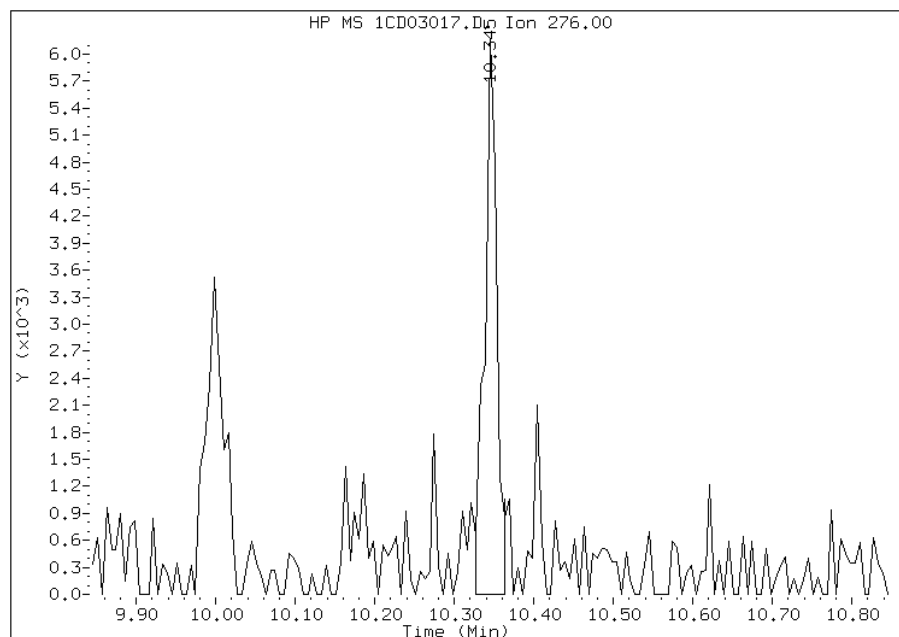


# Manual Integration Report

Data File: 1CD03017.D  
Inj. Date and Time: 03-APR-2013 16:10  
Instrument ID: BSMC5973.i  
Client ID: CV0014AB-GS  
Compound: 26 Benzo(g,h,i)perylene  
CAS #: 191-24-2  
Report Date: 04/04/2013

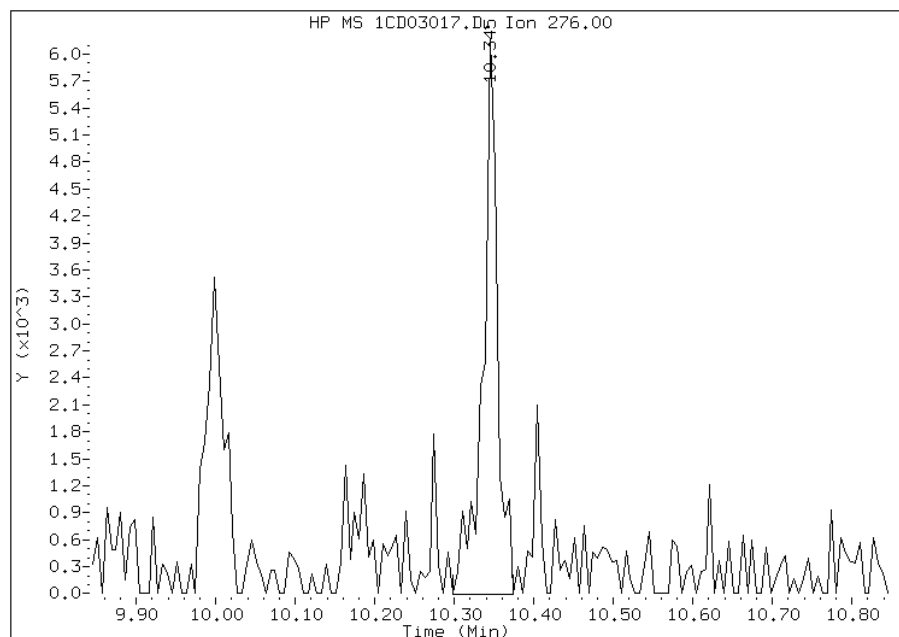
## Processing Integration Results

RT: 10.35  
Response: 6565  
Amount: 0  
Conc: 106



## Manual Integration Results

RT: 10.35  
Response: 7927  
Amount: 0  
Conc: 128



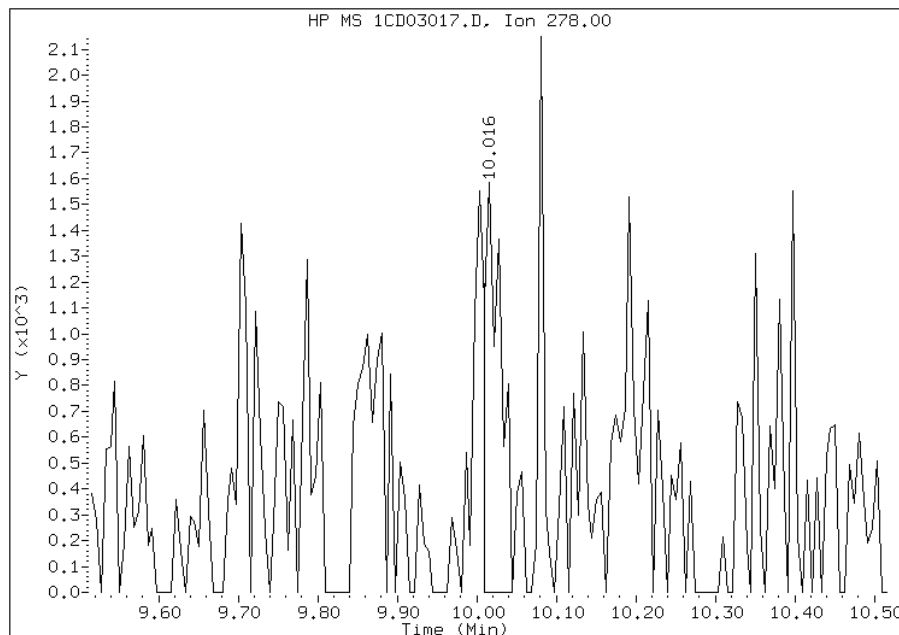
Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:33  
Manual Integration Reason: Baseline Event

# Manual Integration Report

Data File: 1CD03017.D  
Inj. Date and Time: 03-APR-2013 16:10  
Instrument ID: BSMC5973.i  
Client ID: CV0014AB-GS  
Compound: 25 Dibenzo(a,h)anthracene  
CAS #: 53-70-3  
Report Date: 04/04/2013

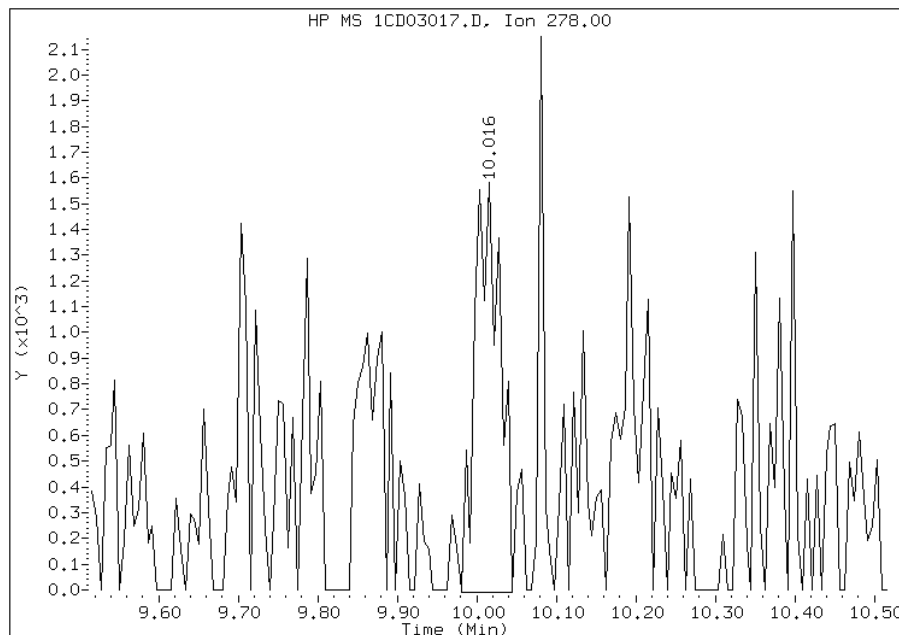
## Processing Integration Results

RT: 10.02  
Response: 2258  
Amount: 0  
Conc: 40



## Manual Integration Results

RT: 10.02  
Response: 3484  
Amount: 0  
Conc: 62



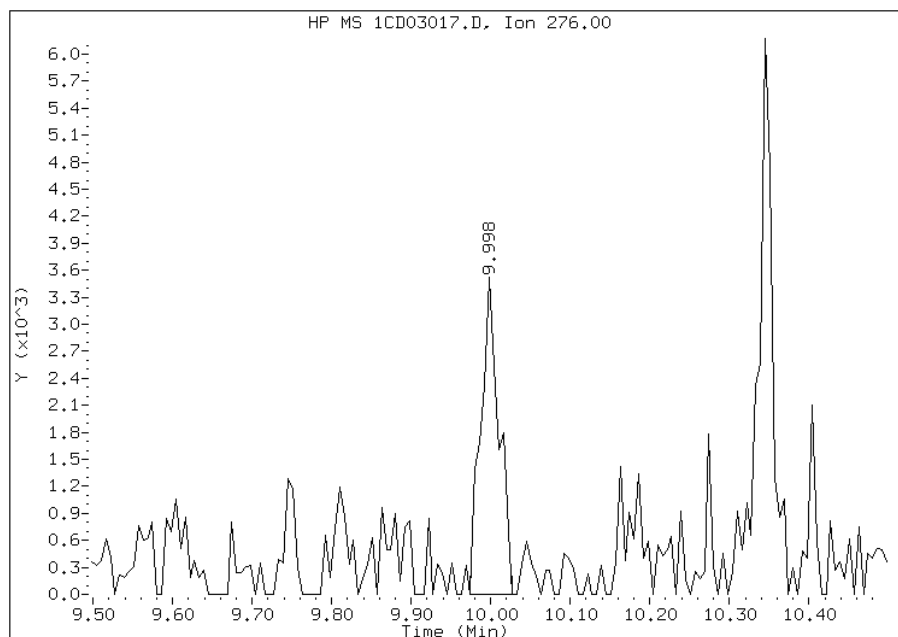
Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:33  
Manual Integration Reason: Baseline Event

# Manual Integration Report

Data File: 1CD03017.D  
Inj. Date and Time: 03-APR-2013 16:10  
Instrument ID: BSMC5973.i  
Client ID: CV0014AB-GS  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/04/2013

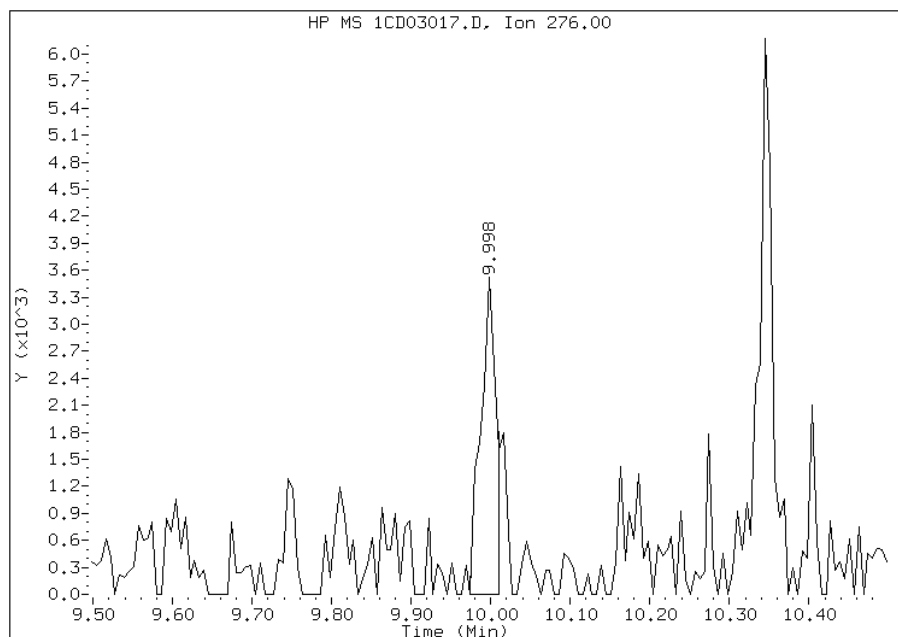
## Processing Integration Results

RT: 10.00  
Response: 5487  
Amount: 0  
Conc: 90



## Manual Integration Results

RT: 10.00  
Response: 4605  
Amount: 0  
Conc: 76



Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:33  
Manual Integration Reason: Split Peak

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: 032613-RB-shovel Lab Sample ID: 680-88766-23  
 Matrix: Water Lab File ID: 1CD05006.D  
 Analysis Method: 8270C LL Date Collected: 03/26/2013 13:00  
 Extract. Method: 3520C Date Extracted: 04/02/2013 08:08  
 Sample wt/vol: 800(mL) Date Analyzed: 04/05/2013 12:54  
 Con. Extract Vol.: 1(mL) Dilution Factor: 1  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136171 Units: ug/L

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	2.5	U F	2.5	0.63
208-96-8	Acenaphthylene	1.3	U	1.3	0.31
120-12-7	Anthracene	0.25	U	0.25	0.095
56-55-3	Benzo[a]anthracene	0.25	U F	0.25	0.063
50-32-8	Benzo[a]pyrene	0.25	U F	0.25	0.071
205-99-2	Benzo[b]fluoranthene	0.25	U F	0.25	0.063
191-24-2	Benzo[g,h,i]perylene	0.63	U F	0.63	0.13
207-08-9	Benzo[k]fluoranthene	0.25	U F	0.25	0.071
218-01-9	Chrysene	0.25	U F	0.25	0.086
53-70-3	Dibenz(a,h)anthracene	0.25	U F	0.25	0.063
206-44-0	Fluoranthene	0.63	U F	0.63	0.068
86-73-7	Fluorene	2.5	U F	2.5	0.63
193-39-5	Indeno[1,2,3-cd]pyrene	0.25	U F	0.25	0.063
90-12-0	1-Methylnaphthalene	2.5	U	2.5	0.63
91-57-6	2-Methylnaphthalene	2.5	U F	2.5	0.63
91-20-3	Naphthalene	2.5	U F	2.5	0.31
85-01-8	Phenanthrene	0.63	U F	0.63	0.25
129-00-0	Pyrene	0.63	U F	0.63	0.11

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	67		30-130

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\1CD05006.D  
 Lab Smp Id: 680-88766-A-23-A Client Smp ID: 032613-RB-shovel  
 Inj Date : 05-APR-2013 12:54  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : 680-88766-A-23-A  
 Misc Info : 680-88766-A-23-A  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\a-bFASTPAHi-m.m  
 Meth Date : 05-Apr-2013 12:31 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 5  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	800.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
							ON-COLUMN (ug/ml)	FINAL ( ug/l)
* 1 Naphthalene-d8	=====	136	3.692	3.692	(1.000)	397216	40.0000	
* 6 Acenaphthene-d10	=====	164	4.774	4.780	(1.000)	288931	40.0000	
* 10 Phenanthrene-d10	=====	188	5.721	5.721	(1.000)	552053	40.0000	
\$ 14 o-Terphenyl	=====	230	5.974	5.974	(1.044)	54191	6.73936	8.4242
* 18 Chrysene-d12	=====	240	7.657	7.662	(1.000)	730677	40.0000	
* 23 Perylene-d12	=====	264	8.821	8.827	(1.000)	752862	40.0000	

Data File: 1CD05006.D

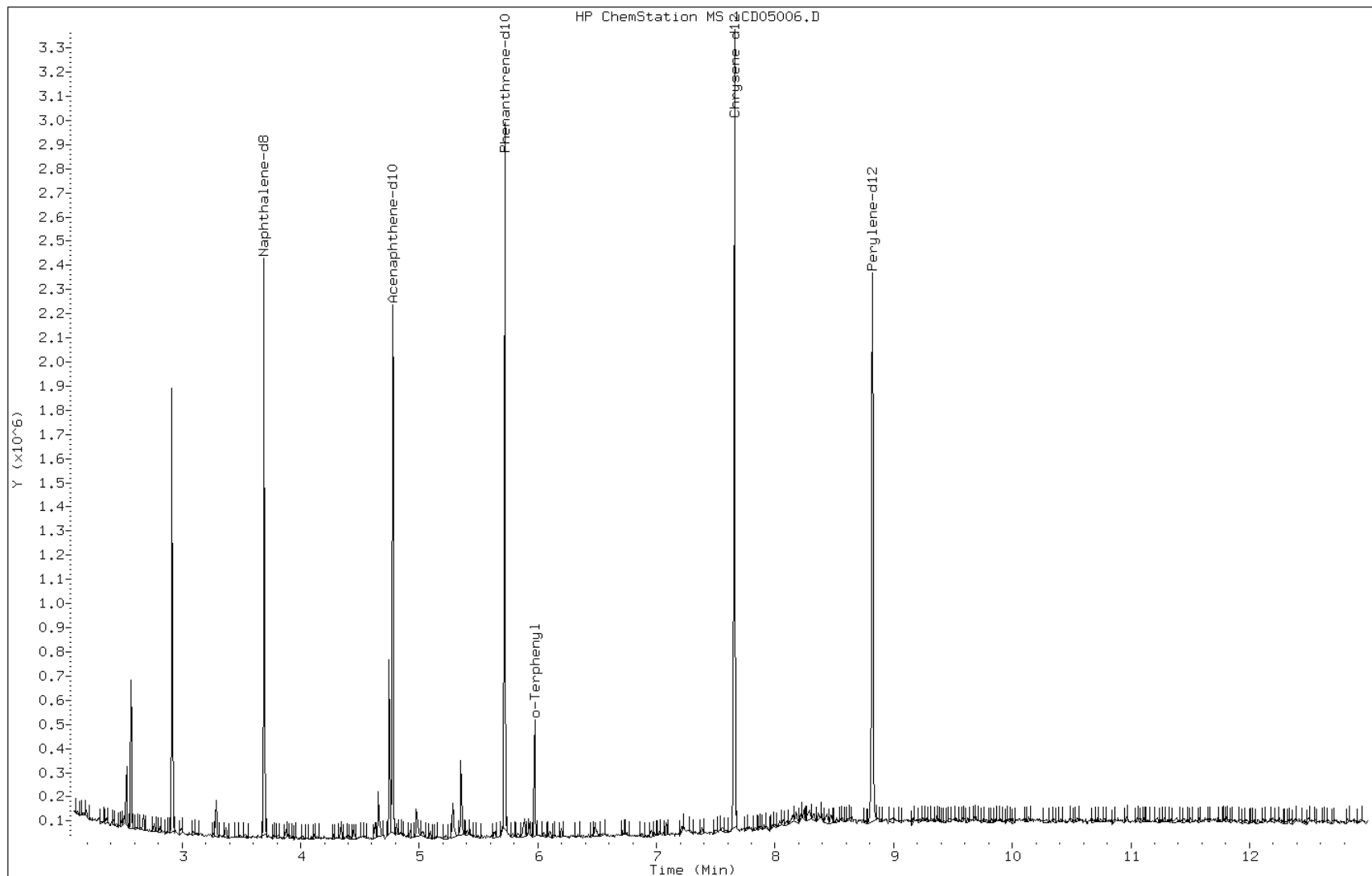
Date: 05-APR-2013 12:54

Client ID: 032613-RB-shovel

Instrument: BSMC5973.i

Sample Info: 680-88766-A-23-A

Operator: SCC



FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD CURVE EVALUATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136048

SDG No.: 68088766-2

Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um) Heated Purge: (Y/N) N

Calibration Start Date: 04/02/2013 13:26 Calibration End Date: 04/02/2013 15:15 Calibration ID: 2859

Calibration Files:

LEVEL:	LAB SAMPLE ID:	LAB FILE ID:
Level 1	IC 660-136048/5	1CD02005.D
Level 2	IC 660-136048/6	1CD02006.D
Level 3	IC 660-136048/7	1CD02007.D
Level 4	IC 660-136048/8	1CD02008.D
Level 5	ICIS 660-136048/9	1CD02009.D
Level 6	IC 660-136048/10	1CD02010.D
Level 7	IC 660-136048/11	1CD02011.D

ANALYTE	RRF					CURVE TYPE	COEFFICIENT			#	MIN RRF	%RSD	#	MAX %RSD	R^2 OR COD	#	MIN R^2 OR COD
	LVL 1	LVL 2	LVL 3	LVL 4	LVL 5		B	M1	M2								
	LVL 6	LVL 7															
Naphthalene	0.9951 1.0462	0.9249 1.0491	1.1511	1.0146	1.0107	Ave		1.0274			0.0000	6.7	15.0				
2-Methylnaphthalene	0.7586 0.6820	0.6817 0.7025	0.6887	0.7485	0.6335	Ave		0.6994			0.0000	6.1	15.0				
1-Methylnaphthalene	0.7248 0.6605	0.4518 0.6576	0.6481	0.6089	0.6533	Ave		0.6293			0.0000	13.6	15.0				
Acenaphthylene	1.4345 1.7430	1.5801 1.7453	1.7015	1.6743	1.7098	Ave		1.6555			0.0000	6.8	15.0				
Acenaphthene	0.8041 1.0063	1.3709 1.0300	0.9518	0.9544	1.0574	Lin		1.0254			0.0000			0.9993		0.9900	
Fluorene	1.2800 1.3623	1.5080 1.3691	1.4076	1.2955	1.3459	Ave		1.3669			0.0000	5.6	15.0				
Phenanthrene	1.2753 1.1465	1.1377 1.2101	1.1311	1.1382	1.1160	Ave		1.1650			0.0000	4.9	15.0				
Anthracene	1.2299 1.2077	1.1082 1.2343	1.1512	1.1740	1.1613	Ave		1.1810			0.0000	3.9	15.0				
Carbazole	0.9389 1.0577	0.8968 1.0652	1.0685	0.9845	1.0709	Ave		1.0118			0.0000	7.1	15.0				
Fluoranthene	1.0844 1.3160	1.1991 1.4023	1.3527	1.3181	1.3335	Ave		1.2866			0.0000	8.4	15.0				
Pyrene	1.0454 1.1504	1.0946 1.1474	1.1166	1.0638	1.1380	Ave		1.1080			0.0000	3.8	15.0				
Benzo[a]anthracene	1.9586 1.1436	1.3015 1.1642	1.1246	1.1267	1.1237	Lin	0.0034	1.1590			0.0000			0.9997		0.9900	
Chrysene	1.0137 1.1434	1.2130 1.1619	1.2029	1.1145	1.1295	Ave		1.1398			0.0000	5.8	15.0				
Benzo[b]fluoranthene	1.4007 1.0698	0.9300 1.1884	1.1544	1.1244	1.0480	Ave		1.1308			0.0000	12.9	15.0				

Note: The m1 coefficient is the same as Ave RRF for an Ave curve type.

FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD CURVE EVALUATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136048

SDG No.: 68088766-2

Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um) Heated Purge: (Y/N) N

Calibration Start Date: 04/02/2013 13:26 Calibration End Date: 04/02/2013 15:15 Calibration ID: 2859

ANALYTE	RRF					CURVE TYPE	COEFFICIENT			#	MIN RRF	%RSD	#	MAX %RSD	R^2 OR COD	#	MIN R^2 OR COD
	LVL 1	LVL 2	LVL 3	LVL 4	LVL 5		B	M1	M2								
	LVL 6	LVL 7															
Benzo[k]fluoranthene	0.9952 1.1459	1.0465 1.1495	1.1058	1.1151	1.0979	Ave		1.0937			0.0000	5.1	15.0				
Benzo[a]pyrene	1.2128 1.0446	0.9589 1.1556	1.0227	1.0341	1.0238	Ave		1.0647			0.0000	8.2	15.0				
Indeno[1,2,3-cd]pyrene	1.2338 1.0436	0.9049 1.0226	1.0384	0.9595	0.8756	Ave		1.0112			0.0000	11.7	15.0				
Dibenz(a,h)anthracene	0.9208 0.9567	0.9397 0.9834	0.8833	0.9304	0.9246	Ave		0.9341			0.0000	3.3	15.0				
Benzo[g,h,i]perylene	1.0683 1.0751	0.9692 1.0455	1.0646	1.0048	0.9970	Ave		1.0321			0.0000	4.0	15.0				
o-Terphenyl	0.8162 0.5958	0.5068 0.6604	0.5759	0.6060	0.6022	Lin	0.0181	0.6529			0.0000			0.9966		0.9900	

Note: The m1 coefficient is the same as Ave RRF for an Ave curve type.



FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD RESPONSE AND CONCENTRATION

Lab Name: TestAmerica Tampa

Job No.: 680-88766-2

Analy Batch No.: 136048

SDG No.: 68088766-2

Instrument ID: BSMC5973

GC Column: DB-5MS

ID: 250 (um)

Heated Purge: (Y/N) N

Calibration Start Date: 04/02/2013 13:26

Calibration End Date: 04/02/2013 15:15

Calibration ID: 2859

Calibration Files:

LEVEL:	LAB SAMPLE ID:	LAB FILE ID:
Level 1	IC 660-136048/5	1CD02005.D
Level 2	IC 660-136048/6	1CD02006.D
Level 3	IC 660-136048/7	1CD02007.D
Level 4	IC 660-136048/8	1CD02008.D
Level 5	ICIS 660-136048/9	1CD02009.D
Level 6	IC 660-136048/10	1CD02010.D
Level 7	IC 660-136048/11	1CD02011.D

ANALYTE	IS REF	CURVE TYPE	RESPONSE					CONCENTRATION (UG/ML)				
			LVL 1 LVL 6	LVL 2 LVL 7	LVL 3	LVL 4	LVL 5	LVL 1 LVL 6	LVL 2 LVL 7	LVL 3	LVL 4	LVL 5
Naphthalene	NPT	Ave	2264 350333	10440 668649	65815	121970	253190	0.200 30.0	1.00 50.0	5.00	10.0	20.0
2-Methylnaphthalene	NPT	Ave	1726 228375	7695 447751	39376	89978	158694	0.200 30.0	1.00 50.0	5.00	10.0	20.0
1-Methylnaphthalene	NPT	Ave	1649 221182	5100 419135	37056	73198	163647	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Acenaphthylene	ANT	Ave	2387 423924	12563 814053	70473	148174	308909	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Acenaphthene	ANT	Lin	1338 244735	10900 480392	39421	84460	191043	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Fluorene	ANT	Ave	2130 331328	11990 638557	58298	114648	243174	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Phenanthrene	PHN	Ave	3900 529536	16838 1077014	88442	194036	392252	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Anthracene	PHN	Ave	3761 557837	16401 1098599	90016	200131	408192	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Carbazole	PHN	Ave	2871 488550	13272 948101	83549	167822	376402	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Fluoranthene	PHN	Ave	3316 607836	17746 1248081	105772	224705	468708	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Pyrene	CRY	Ave	4087 663294	20532 1360548	109963	236267	498076	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[a]anthracene	CRY	Lin	7657 659379	24413 1380443	110756	250220	491852	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Chrysene	CRY	Ave	3963 659226	22752 1377767	118460	247512	494376	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[b]fluoranthene	PRY	Ave	5890 671785	19731 1443812	127315	261073	494109	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[k]fluoranthene	PRY	Ave	4185 719552	22203 1396501	121957	258924	517620	0.200 30.0	1.00 50.0	5.00	10.0	20.0

FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD RESPONSE AND CONCENTRATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136048

SDG No.: 68088766-2

Instrument ID: BSMC5973 GC Column: DB-5MS ID: 250 (um) Heated Purge: (Y/N) N

Calibration Start Date: 04/02/2013 13:26 Calibration End Date: 04/02/2013 15:15 Calibration ID: 2859

ANALYTE	IS REF	CURVE TYPE	RESPONSE					CONCENTRATION (UG/ML)				
			LVL 1	LVL 2	LVL 3	LVL 4	LVL 5	LVL 1	LVL 2	LVL 3	LVL 4	LVL 5
			LVL 6	LVL 7				LVL 6	LVL 7			
Benzo[a]pyrene	PRY	Ave	5100	20343	112782	240110	482722	0.200	1.00	5.00	10.0	20.0
			655944	1403971				30.0	50.0			
Indeno[1,2,3-cd]pyrene	PRY	Ave	5188	19198	114519	222795	412839	0.200	1.00	5.00	10.0	20.0
			655344	1242391				30.0	50.0			
Dibenz(a,h)anthracene	PRY	Ave	3872	19937	97409	216036	435940	0.200	1.00	5.00	10.0	20.0
			600720	1194691				30.0	50.0			
Benzo[g,h,i]perylene	PRY	Ave	4492	20561	117403	233308	470085	0.200	1.00	5.00	10.0	20.0
			675124	1270187				30.0	50.0			
o-Terphenyl	PHN	Lin	2496	7501	45027	103309	211673	0.200	1.00	5.00	10.0	20.0
			275212	587824				30.0	50.0			

Curve Type Legend:

Ave = Average ISTD
Lin = Linear ISTD

TestAmerica Laboratories

Semivolatile 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02005.D  
 Lab Smp Id: IC1  
 Inj Date : 02-APR-2013 13:26  
 Operator : SCC  
 Smp Info : IC1  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 5 Calibration Sample, Level: 1  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)
* 1 Naphthalene-d8	136	3.710	3.710	(1.000)	455021	40.0000	
* 6 Acenaphthene-d10	164	4.804	4.804	(1.000)	332800	40.0000	
* 10 Phenanthrene-d10	188	5.757	5.757	(1.000)	611597	40.0000	
\$ 14 o-Terphenyl	230	6.004	6.004	(1.043)	2496	0.20000	0.2618
* 18 Chrysene-d12	240	7.704	7.704	(1.000)	781900	40.0000	
* 23 Perylene-d12	264	8.909	8.909	(1.000)	841000	40.0000	(H)
2 Naphthalene	128	3.727	3.727	(1.005)	2264	0.20000	0.1937
3 2-Methylnaphthalene	142	4.157	4.157	(1.120)	1726	0.20000	0.2169
4 1-Methylnaphthalene	142	4.216	4.216	(1.136)	1649	0.20000	0.2303
5 Acenaphthylene	152	4.716	4.716	(0.982)	2387	0.20000	0.1733
7 Acenaphthene	154	4.821	4.821	(1.004)	1338	0.20000	0.1568(Q)
9 Fluorene	166	5.145	5.145	(1.071)	2130	0.20000	0.1872
11 Phenanthrene	178	5.768	5.768	(1.002)	3900	0.20000	0.2189
12 Anthracene	178	5.804	5.804	(1.008)	3761	0.20000	0.2082
13 Carbazole	167	5.915	5.915	(1.028)	2871	0.20000	0.1855
15 Fluoranthene	202	6.604	6.604	(1.147)	3316	0.20000	0.1685
16 Pyrene	202	6.774	6.774	(0.879)	4087	0.20000	0.1886
17 Benzo(a)anthracene	228	7.698	7.698	(0.999)	7657	0.20000	0.3066
19 Chrysene	228	7.727	7.727	(1.003)	3963	0.20000	0.1778
20 Benzo(b)fluoranthene	252	8.562	8.562	(0.961)	5890	0.20000	0.2477(H)
21 Benzo(k)fluoranthene	252	8.586	8.586	(0.964)	4185	0.20000	0.1819(H)
22 Benzo(a)pyrene	252	8.851	8.851	(0.993)	5100	0.20000	0.2278(H)
24 Indeno(1,2,3-cd)pyrene	276	10.062	10.062	(1.129)	5188	0.20000	0.2440
25 Dibenzo(a,h)anthracene	278	10.086	10.086	(1.132)	3872	0.20000	0.1971(MH)
26 Benzo(g,h,i)perylene	276	10.415	10.415	(1.169)	4492	0.20000	0.2070(H)

QC Flag Legend

- Q - Qualifier signal failed the ratio test.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1CD02005.D

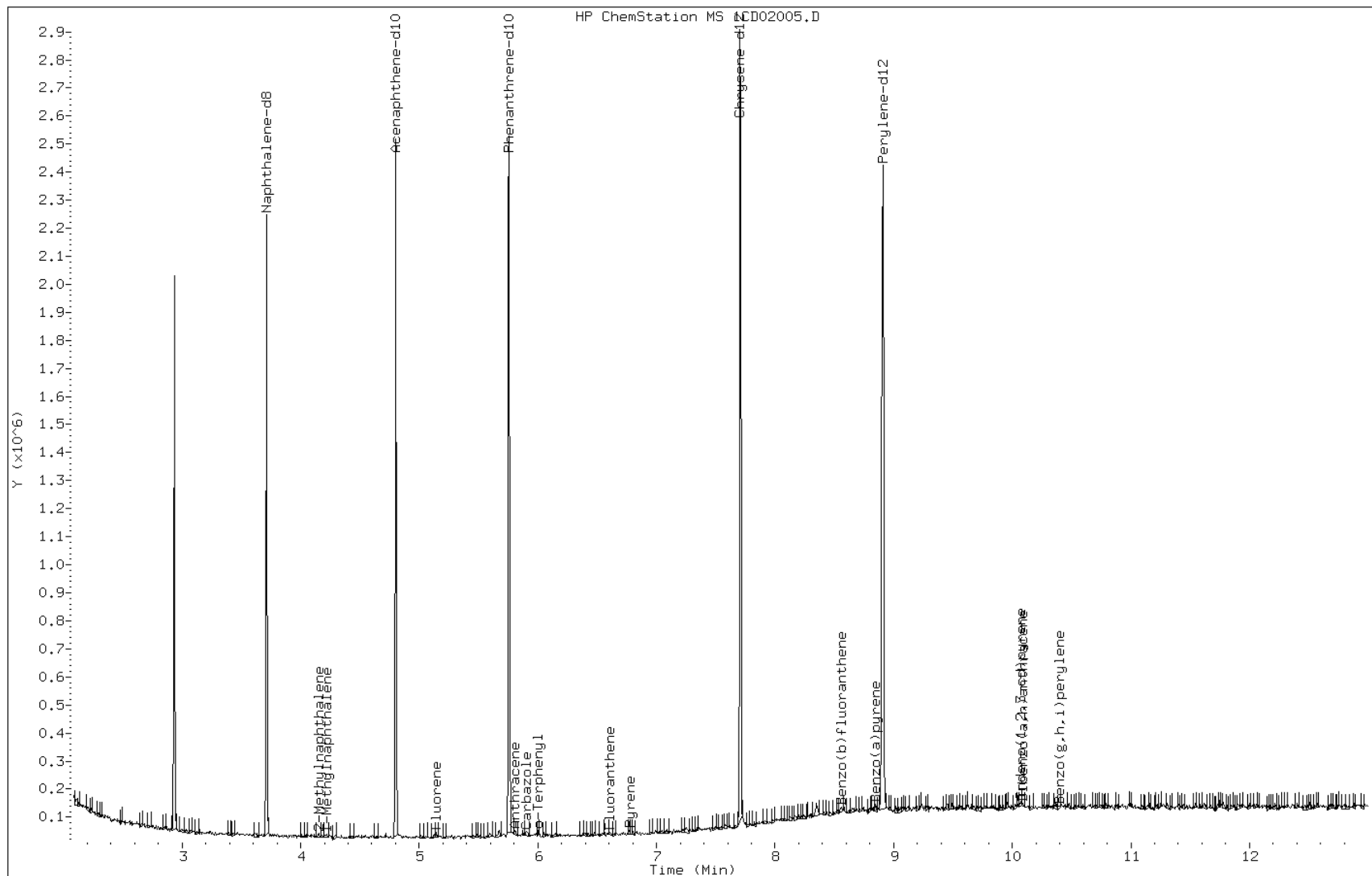
Date: 02-APR-2013 13:26

Client ID:

Instrument: BSMC5973.i

Sample Info: IC1

Operator: SCC

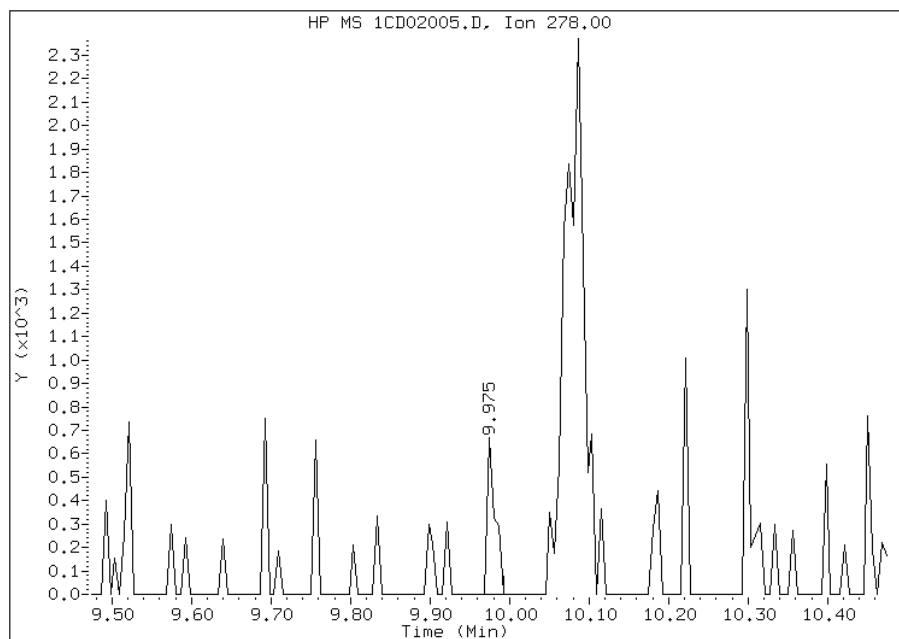


# Manual Integration Report

Data File: 1CD02005.D  
Inj. Date and Time: 02-APR-2013 13:26  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 25 Dibenzo(a,h)anthracene  
CAS #: 53-70-3  
Report Date: 04/02/2013

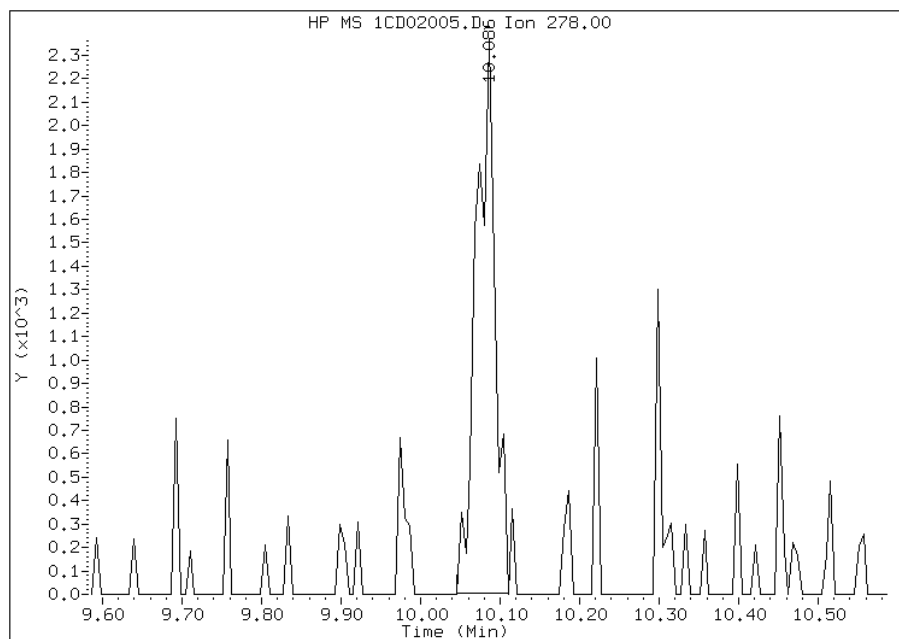
## Processing Integration Results

RT: 9.97  
Response: 454  
Amount: 0  
Conc: 0



## Manual Integration Results

RT: 10.09  
Response: 3872  
Amount: 0  
Conc: 0



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:44  
Manual Integration Reason: Baseline Event

TestAmerica Laboratories

Semivolatile 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02006.D  
 Lab Smp Id: IC2  
 Inj Date : 02-APR-2013 13:44  
 Operator : SCC  
 Smp Info : IC2  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 13:26 Cal File: 1CD02005.D  
 Als bottle: 6 Calibration Sample, Level: 2  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT	SIG	AMOUNTS					
			CAL-AMT	ON-COL	MASS	RT	EXP RT	REL RT
* 1 Naphthalene-d8	136		40.0000		3.710	3.710	(1.000)	451517
* 6 Acenaphthene-d10	164		40.0000		4.798	4.798	(1.000)	318036
* 10 Phenanthrene-d10	188		40.0000		5.745	5.745	(1.000)	591987
\$ 14 o-Terphenyl	230		1.00000	0.8130	5.998	5.998	(1.044)	7501
* 18 Chrysene-d12	240		40.0000	(H)	7.686	7.686	(1.000)	750291
* 23 Perylene-d12	264		40.0000	(H)	8.862	8.862	(1.000)	848618
2 Naphthalene	128		1.00000	0.9002	3.727	3.727	(1.005)	10440
3 2-Methylnaphthalene	142		1.00000	0.9747	4.151	4.151	(1.119)	7695
4 1-Methylnaphthalene	142		1.00000	0.7179(Q)	4.216	4.216	(1.136)	5100
5 Acenaphthylene	152		1.00000	0.9544	4.710	4.710	(0.982)	12563
7 Acenaphthene	154		1.00000	1.3375(Q)	4.821	4.821	(1.005)	10900
9 Fluorene	166		1.00000	1.1032	5.139	5.139	(1.071)	11990
11 Phenanthrene	178		1.00000	0.9766	5.762	5.762	(1.003)	16838
12 Anthracene	178		1.00000	0.9383	5.798	5.798	(1.009)	16401
13 Carbazole	167		1.00000	0.8863	5.904	5.904	(1.028)	13272
15 Fluoranthene	202		1.00000	0.9319	6.598	6.598	(1.148)	17746
16 Pyrene	202		1.00000	0.9878(H)	6.762	6.762	(0.880)	20532
17 Benzo(a)anthracene	228		1.00000	1.0187(H)	7.680	7.680	(0.999)	24413
19 Chrysene	228		1.00000	1.0641	7.704	7.704	(1.002)	22752
20 Benzo(b)fluoranthene	252		1.00000	0.8224(H)	8.521	8.521	(0.962)	19731
21 Benzo(k)fluoranthene	252		1.00000	0.9568(H)	8.539	8.539	(0.963)	22203
22 Benzo(a)pyrene	252		1.00000	0.9006(H)	8.809	8.809	(0.994)	20343
24 Indeno(1,2,3-cd)pyrene	276		1.00000	0.8948(MH)	10.009	10.009	(1.129)	19198
25 Dibenzo(a,h)anthracene	278		1.00000	1.0060(H)	10.027	10.027	(1.131)	19937
26 Benzo(g,h,i)perylene	276		1.00000	0.9390(H)	10.356	10.356	(1.169)	20561

QC Flag Legend

- Q - Qualifier signal failed the ratio test.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1CD02006.D

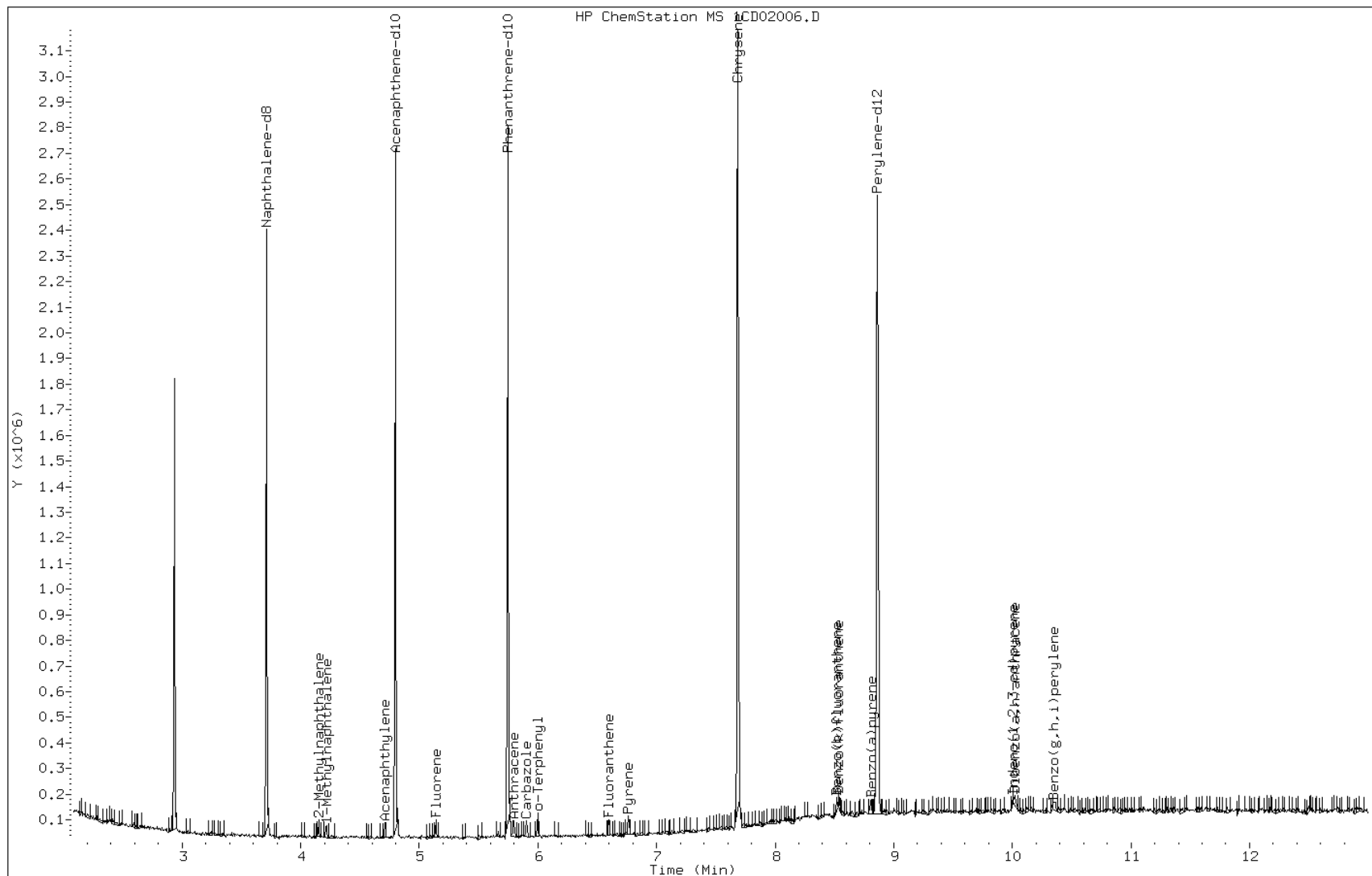
Date: 02-APR-2013 13:44

Client ID:

Instrument: BSMC5973.i

Sample Info: IC2

Operator: SCC

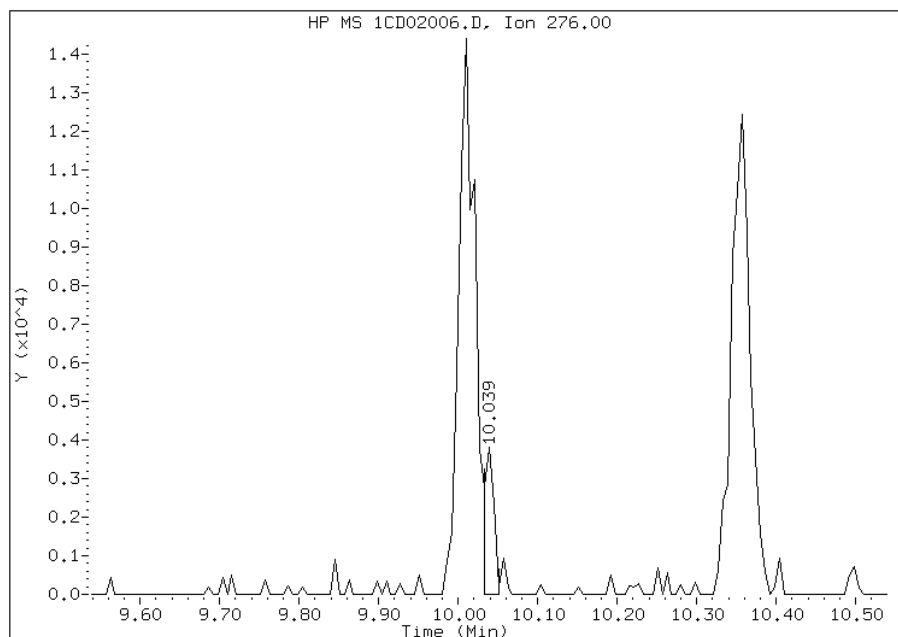


# Manual Integration Report

Data File: 1CD02006.D  
Inj. Date and Time: 02-APR-2013 13:44  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

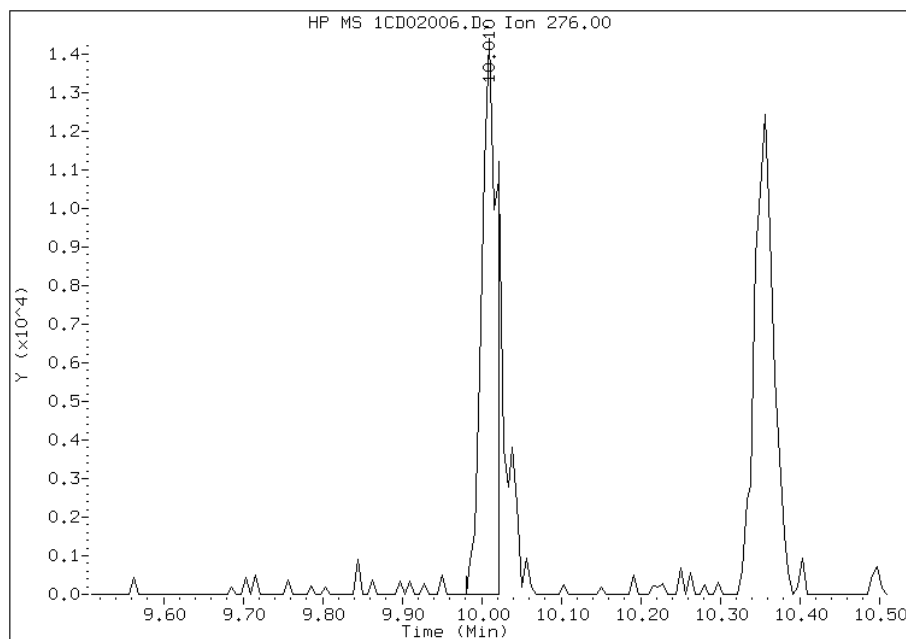
## Processing Integration Results

RT: 10.04  
Response: 3225  
Amount: 0  
Conc: 0



## Manual Integration Results

RT: 10.01  
Response: 19198  
Amount: 1  
Conc: 1



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:45  
Manual Integration Reason: Split Peak



TestAmerica Laboratories

Semivolatile 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02007.D  
 Lab Smp Id: IC3  
 Inj Date : 02-APR-2013 14:02  
 Operator : SCC  
 Smp Info : IC3  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 13:44 Cal File: 1CD02006.D  
 Als bottle: 7 Calibration Sample, Level: 3  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT	SIG	AMOUNTS					
			MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)
* 1 Naphthalene-d8	136		3.710	3.710	(1.000)	457408	40.0000	
* 6 Acenaphthene-d10	164		4.798	4.798	(1.000)	331342	40.0000	
* 10 Phenanthrene-d10	188		5.745	5.745	(1.000)	625535	40.0000	
\$ 14 o-Terphenyl	230		5.998	5.998	(1.044)	45027	5.00000	4.6190
* 18 Chrysene-d12	240		7.686	7.686	(1.000)	787858	40.0000	
* 23 Perylene-d12	264		8.856	8.856	(1.000)	882270	40.0000	(H)
2 Naphthalene	128		3.727	3.727	(1.005)	65815	5.00000	5.6020
3 2-Methylnaphthalene	142		4.151	4.151	(1.119)	39376	5.00000	4.9236
4 1-Methylnaphthalene	142		4.216	4.216	(1.136)	37056	5.00000	5.1494(Q)
5 Acenaphthylene	152		4.710	4.710	(0.982)	70473	5.00000	5.1389
7 Acenaphthene	154		4.821	4.821	(1.005)	39421	5.00000	4.6430
9 Fluorene	166		5.139	5.139	(1.071)	58298	5.00000	5.1486
11 Phenanthrene	178		5.763	5.763	(1.003)	88442	5.00000	4.8545
12 Anthracene	178		5.792	5.792	(1.008)	90016	5.00000	4.8741
13 Carbazole	167		5.904	5.904	(1.028)	83549	5.00000	5.2803
15 Fluoranthene	202		6.598	6.598	(1.148)	105772	5.00000	5.2570
16 Pyrene	202		6.762	6.762	(0.880)	109963	5.00000	5.0385
17 Benzo(a)anthracene	228		7.674	7.674	(0.998)	110756	5.00000	4.4014
19 Chrysene	228		7.704	7.704	(1.002)	118460	5.00000	5.2764(H)
20 Benzo(b)fluoranthene	252		8.515	8.515	(0.961)	127315	5.00000	5.1043
21 Benzo(k)fluoranthene	252		8.539	8.539	(0.964)	121957	5.00000	5.0554(H)
22 Benzo(a)pyrene	252		8.804	8.804	(0.994)	112782	5.00000	4.8027(H)
24 Indeno(1,2,3-cd)pyrene	276		10.003	10.003	(1.129)	114519	5.00000	5.1344(MH)
25 Dibenzo(a,h)anthracene	278		10.021	10.021	(1.131)	97409	5.00000	4.7277(H)
26 Benzo(g,h,i)perylene	276		10.345	10.345	(1.168)	117403	5.00000	5.1573(H)

QC Flag Legend

- Q - Qualifier signal failed the ratio test.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1CD02007.D

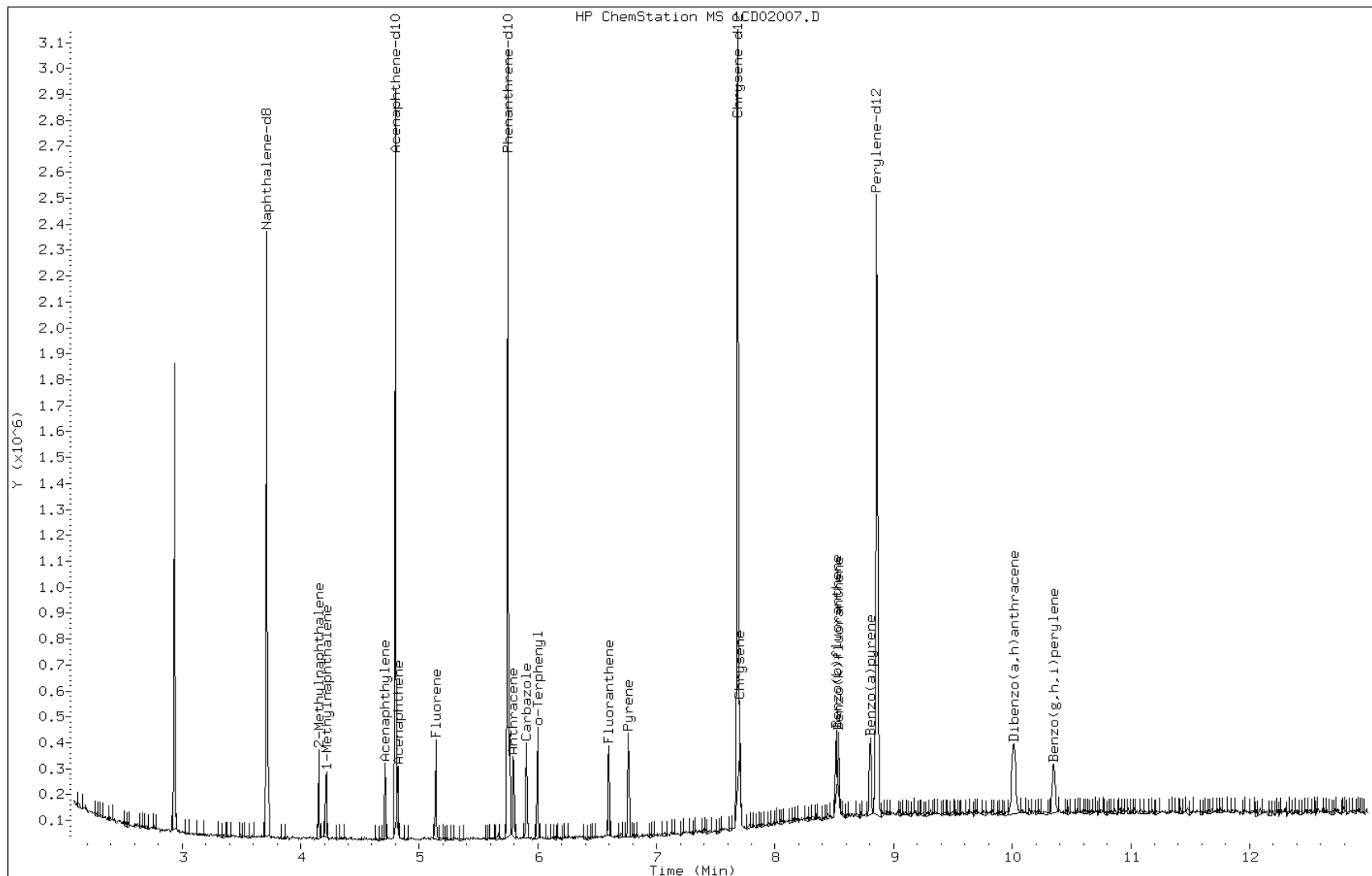
Date: 02-APR-2013 14:02

Client ID:

Instrument: BSMC5973.i

Sample Info: IC3

Operator: SCC

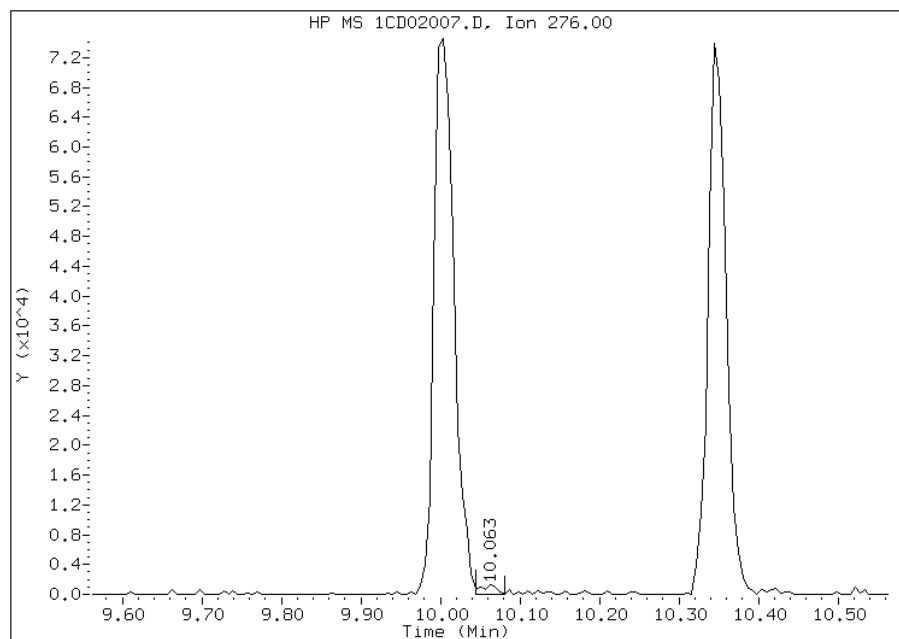


# Manual Integration Report

Data File: 1CD02007.D  
Inj. Date and Time: 02-APR-2013 14:02  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

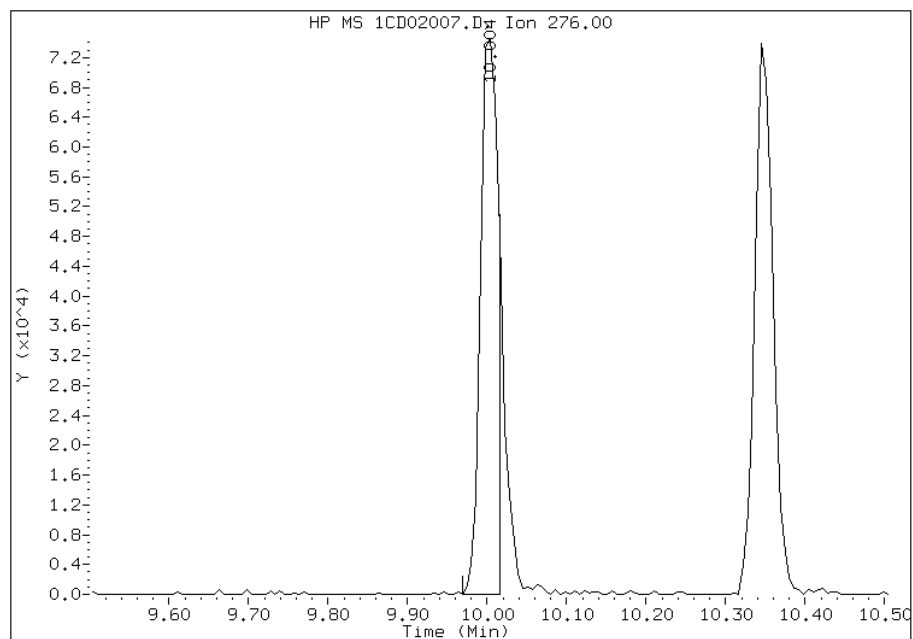
## Processing Integration Results

RT: 10.06  
Response: 1809  
Amount: 0  
Conc: 0



## Manual Integration Results

RT: 10.00  
Response: 114519  
Amount: 5  
Conc: 5



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:48  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02008.D  
 Lab Smp Id: IC4  
 Inj Date : 02-APR-2013 14:20  
 Operator : SCC  
 Smp Info : IC4  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 14:02 Cal File: 1CD02007.D  
 Als bottle: 8 Calibration Sample, Level: 4  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)
* 1 Naphthalene-d8	136	3.710	3.710	(1.000)	480844	40.0000	
* 6 Acenaphthene-d10	164	4.798	4.798	(1.000)	353988	40.0000	
* 10 Phenanthrene-d10	188	5.745	5.745	(1.000)	681887	40.0000	
\$ 14 o-Terphenyl	230	5.998	5.998	(1.044)	103309	10.0000	9.7219
* 18 Chrysene-d12	240	7.686	7.686	(1.000)	888354	40.0000	
* 23 Perylene-d12	264	8.856	8.856	(1.000)	928754	40.0000	
2 Naphthalene	128	3.727	3.727	(1.005)	121970	10.0000	9.8758
3 2-Methylnaphthalene	142	4.151	4.151	(1.119)	89978	10.0000	10.7026
4 1-Methylnaphthalene	142	4.215	4.215	(1.136)	73198	10.0000	9.6761
5 Acenaphthylene	152	4.710	4.710	(0.982)	148174	10.0000	10.1137
7 Acenaphthene	154	4.821	4.821	(1.005)	84460	10.0000	9.3113
9 Fluorene	166	5.139	5.139	(1.071)	114648	10.0000	9.4775
11 Phenanthrene	178	5.762	5.762	(1.003)	194036	10.0000	9.7703
12 Anthracene	178	5.792	5.792	(1.008)	200131	10.0000	9.9409
13 Carbazole	167	5.904	5.904	(1.028)	167822	10.0000	9.7299
15 Fluoranthene	202	6.598	6.598	(1.148)	224705	10.0000	10.2452
16 Pyrene	202	6.762	6.762	(0.880)	236267	10.0000	9.6011
17 Benzo(a)anthracene	228	7.674	7.674	(0.998)	250220	10.0000	8.8188
19 Chrysene	228	7.703	7.703	(1.002)	247512	10.0000	9.7775(H)
20 Benzo(b)fluoranthene	252	8.515	8.515	(0.961)	261073	10.0000	9.9431(H)
21 Benzo(k)fluoranthene	252	8.539	8.539	(0.964)	258924	10.0000	10.1958(H)
22 Benzo(a)pyrene	252	8.803	8.803	(0.994)	240110	10.0000	9.7131
24 Indeno(1,2,3-cd)pyrene	276	10.003	10.003	(1.129)	222795	10.0000	9.4889(MH)
25 Dibenzo(a,h)anthracene	278	10.021	10.021	(1.131)	216036	10.0000	9.9604
26 Benzo(g,h,i)perylene	276	10.350	10.350	(1.169)	233308	10.0000	9.7359(H)

QC Flag Legend

M - Compound response manually integrated.  
 H - Operator selected an alternate compound hit.

Data File: 1CD02008.D

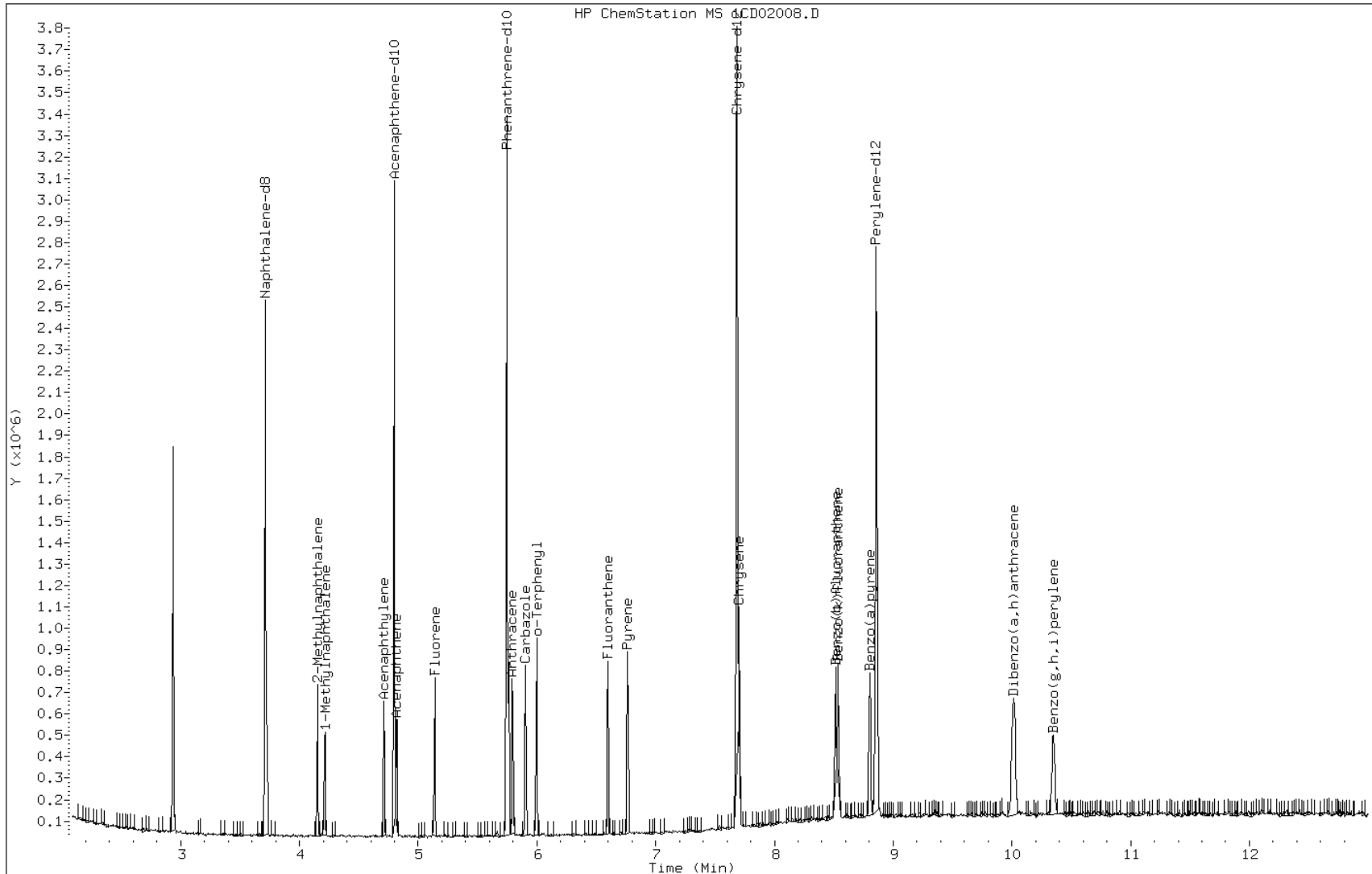
Date: 02-APR-2013 14:20

Client ID:

Instrument: BSMC5973.i

Sample Info: IC4

Operator: SCC

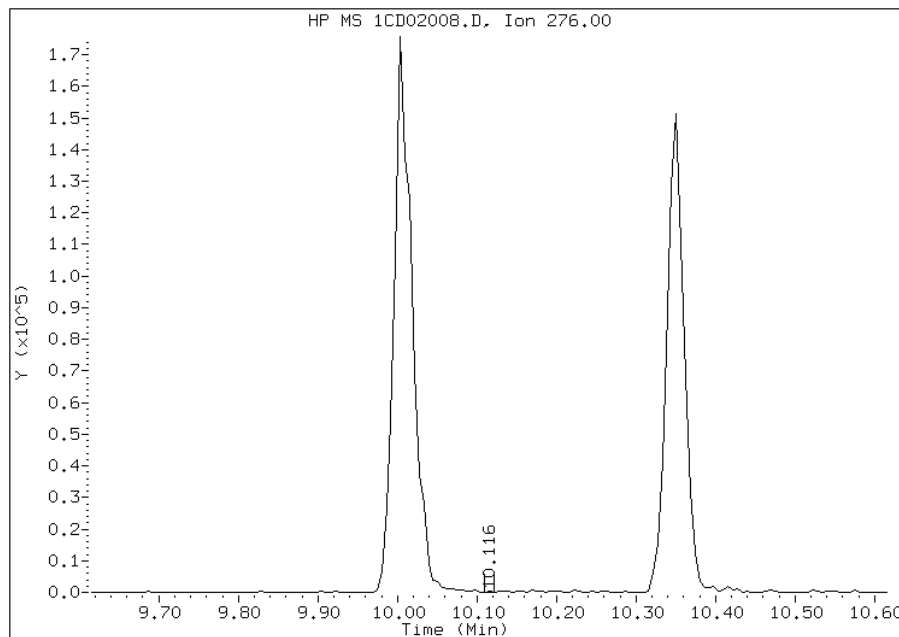


# Manual Integration Report

Data File: 1CD02008.D  
Inj. Date and Time: 02-APR-2013 14:20  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

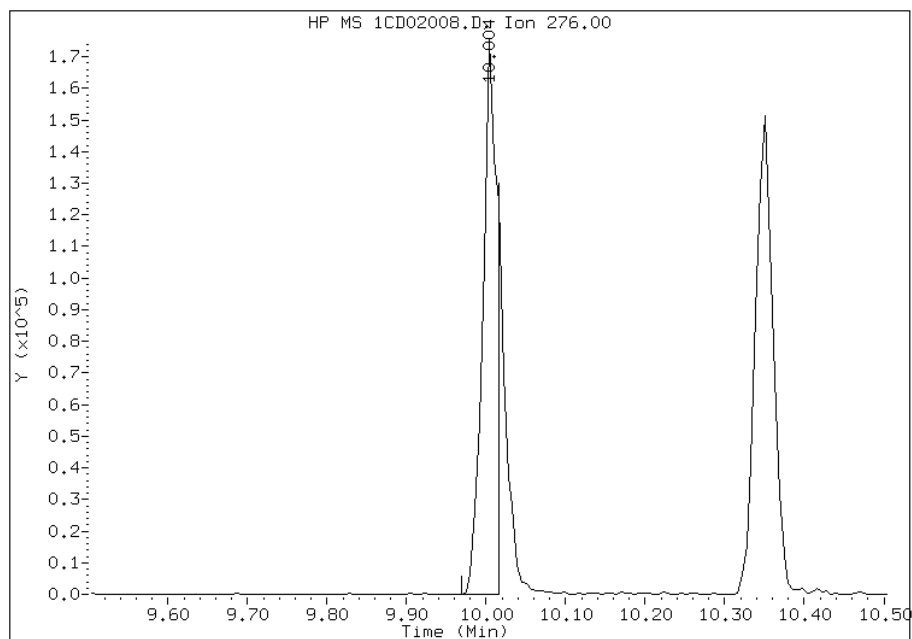
## Processing Integration Results

RT: 10.12  
Response: 142  
Amount: 0  
Conc: 0



## Manual Integration Results

RT: 10.00  
Response: 222795  
Amount: 9  
Conc: 9



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:49  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02009.D  
 Lab Smp Id: IC5  
 Inj Date : 02-APR-2013 14:39  
 Operator : SCC  
 Smp Info : IC5  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 14:20 Cal File: 1CD02008.D  
 Als bottle: 9 Calibration Sample, Level: 5  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG						AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)	
* 1 Naphthalene-d8	136	3.710	3.710	(1.000)	501011	40.0000		
* 6 Acenaphthene-d10	164	4.798	4.798	(1.000)	361349	40.0000		
* 10 Phenanthrene-d10	188	5.745	5.745	(1.000)	702974	40.0000		
\$ 14 o-Terphenyl	230	5.998	5.998	(1.044)	211673	20.0000	19.3221	
* 18 Chrysene-d12	240	7.686	7.686	(1.000)	875378	40.0000		
* 23 Perylene-d12	264	8.862	8.862	(1.000)	942955	40.0000		
2 Naphthalene	128	3.721	3.721	(1.003)	253190	20.0000	19.6753	
3 2-Methylnaphthalene	142	4.151	4.151	(1.119)	158694	20.0000	18.1163	
4 1-Methylnaphthalene	142	4.216	4.216	(1.136)	163647	20.0000	20.7620	
5 Acenaphthylene	152	4.710	4.710	(0.982)	308909	20.0000	20.6554	
7 Acenaphthene	154	4.821	4.821	(1.005)	191043	20.0000	20.6326	
9 Fluorene	166	5.139	5.139	(1.071)	243174	20.0000	19.6928	
11 Phenanthrene	178	5.762	5.762	(1.003)	392252	20.0000	19.1586	
12 Anthracene	178	5.798	5.798	(1.009)	408192	20.0000	19.6676	
13 Carbazole	167	5.904	5.904	(1.028)	376402	20.0000	21.1684	
15 Fluoranthene	202	6.598	6.598	(1.148)	468708	20.0000	20.7293	
16 Pyrene	202	6.762	6.762	(0.880)	498076	20.0000	20.5403	
17 Benzo(a)anthracene	228	7.674	7.674	(0.998)	491852	20.0000	17.5920	
19 Chrysene	228	7.704	7.704	(1.002)	494376	20.0000	19.8190	
20 Benzo(b)fluoranthene	252	8.515	8.515	(0.961)	494109	20.0000	18.5350	
21 Benzo(k)fluoranthene	252	8.539	8.539	(0.963)	517620	20.0000	20.0758	
22 Benzo(a)pyrene	252	8.803	8.803	(0.993)	482722	20.0000	19.2334	
24 Indeno(1,2,3-cd)pyrene	276	10.009	10.009	(1.129)	412839	20.0000	17.3182(M)	
25 Dibenzo(a,h)anthracene	278	10.021	10.021	(1.131)	435940	20.0000	19.7965	
26 Benzo(g,h,i)perylene	276	10.356	10.356	(1.169)	470085	20.0000	19.3212	

QC Flag Legend

M - Compound response manually integrated.

Data File: 1CD02009.D

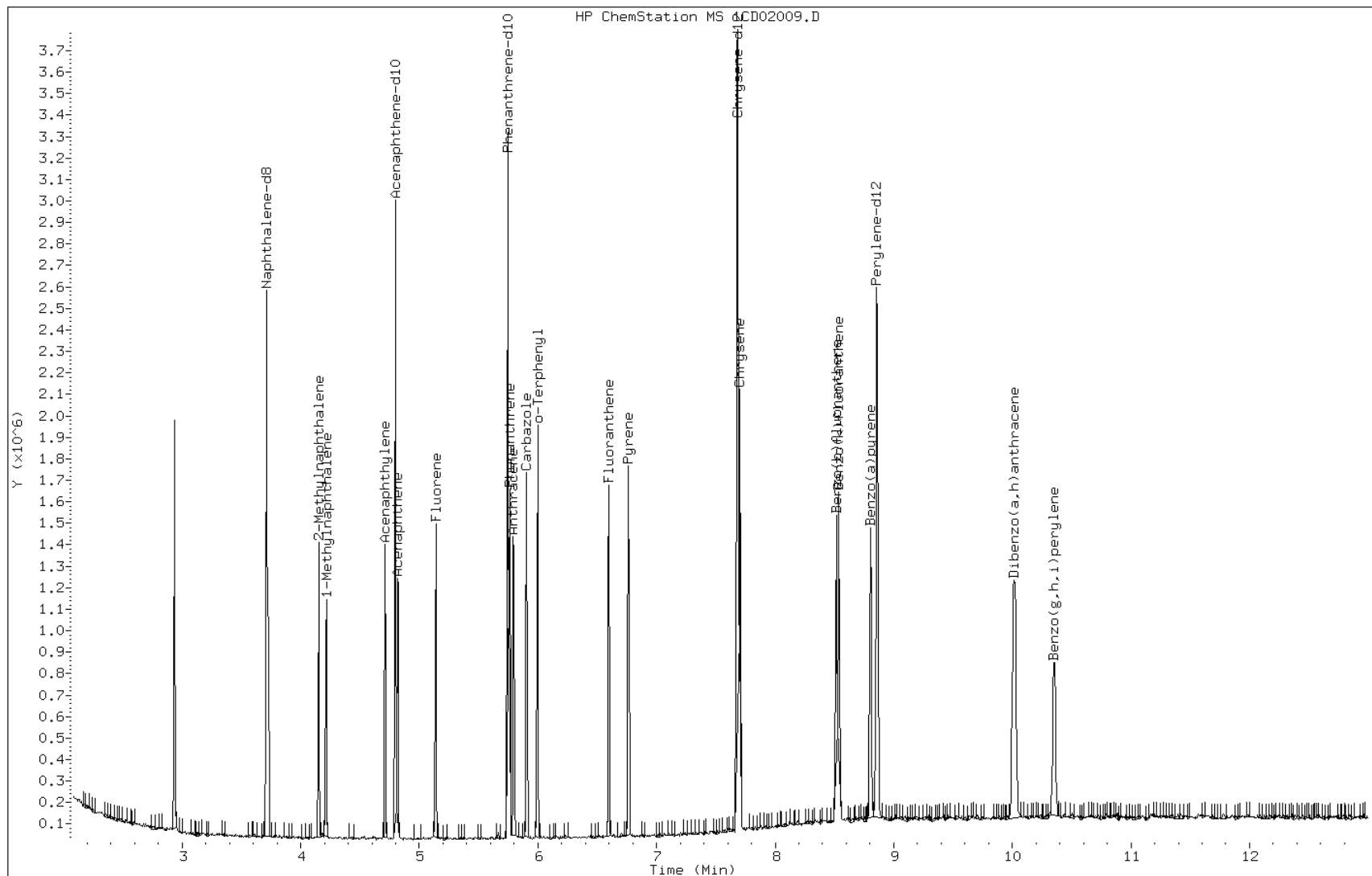
Date: 02-APR-2013 14:39

Client ID:

Instrument: BSMC5973.i

Sample Info: IC5

Operator: SCC



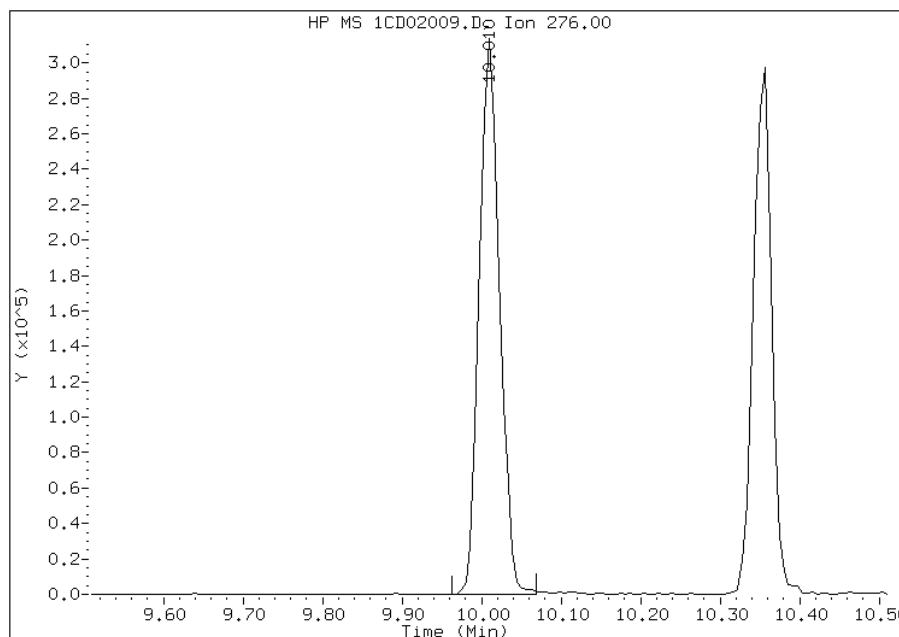


# Manual Integration Report

Data File: 1CD02009.D  
Inj. Date and Time: 02-APR-2013 14:39  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

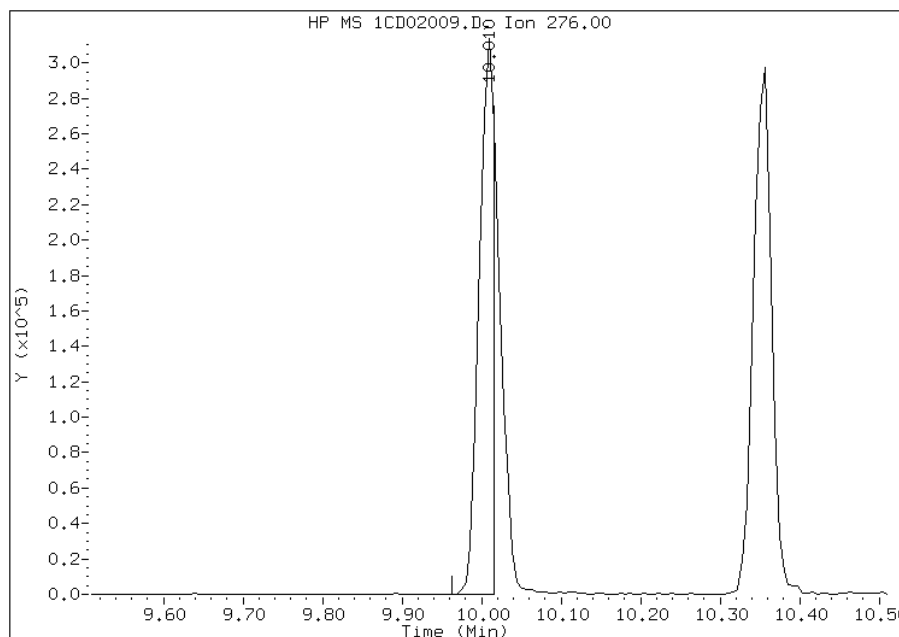
## Processing Integration Results

RT: 10.01  
Response: 550558  
Amount: 32  
Conc: 32



## Manual Integration Results

RT: 10.01  
Response: 412839  
Amount: 17  
Conc: 17



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:39  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02010.D  
 Lab Smp Id: IC6  
 Inj Date : 02-APR-2013 14:57  
 Operator : SCC  
 Smp Info : IC6  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 14:39 Cal File: 1CD02009.D  
 Als bottle: 10 Calibration Sample, Level: 6  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)
* 1 Naphthalene-d8	136	3.710	3.710	(1.000)	446499	40.0000	
* 6 Acenaphthene-d10	164	4.798	4.798	(1.000)	324284	40.0000	
* 10 Phenanthrene-d10	188	5.745	5.745	(1.000)	615852	40.0000	
\$ 14 o-Terphenyl	230	5.998	5.998	(1.044)	275212	30.0000	28.6761
* 18 Chrysene-d12	240	7.686	7.686	(1.000)	768745	40.0000	
* 23 Perylene-d12	264	8.857	8.857	(1.000)	837251	40.0000	
2 Naphthalene	128	3.722	3.722	(1.003)	350333	30.0000	30.5481
3 2-Methylnaphthalene	142	4.151	4.151	(1.119)	228375	30.0000	29.2540
4 1-Methylnaphthalene	142	4.216	4.216	(1.136)	221182	30.0000	31.4875
5 Acenaphthylene	152	4.710	4.710	(0.982)	423924	30.0000	31.5858
7 Acenaphthene	154	4.822	4.822	(1.005)	244735	30.0000	29.4523
9 Fluorene	166	5.139	5.139	(1.071)	331328	30.0000	29.8986
11 Phenanthrene	178	5.763	5.763	(1.003)	529536	30.0000	29.5228
12 Anthracene	178	5.792	5.792	(1.008)	557837	30.0000	30.6801
13 Carbazole	167	5.904	5.904	(1.028)	488550	30.0000	31.3623
15 Fluoranthene	202	6.598	6.598	(1.148)	607836	30.0000	30.6854
16 Pyrene	202	6.763	6.763	(0.880)	663294	30.0000	31.1481
17 Benzo(a)anthracene	228	7.674	7.674	(0.998)	659379	30.0000	26.8553
19 Chrysene	228	7.704	7.704	(1.002)	659226	30.0000	30.0935(H)
20 Benzo(b)fluoranthene	252	8.515	8.515	(0.961)	671785	30.0000	28.3815(H)
21 Benzo(k)fluoranthene	252	8.539	8.539	(0.964)	719552	30.0000	31.4311(H)
22 Benzo(a)pyrene	252	8.804	8.804	(0.994)	655944	30.0000	29.4349
24 Indeno(1,2,3-cd)pyrene	276	10.009	10.009	(1.130)	655344	30.0000	30.9619(MH)
25 Dibenzo(a,h)anthracene	278	10.027	10.027	(1.132)	600720	30.0000	30.7234
26 Benzo(g,h,i)perylene	276	10.356	10.356	(1.169)	675124	30.0000	31.2520(H)

QC Flag Legend

M - Compound response manually integrated.  
 H - Operator selected an alternate compound hit.

Data File: 1CD02010.D

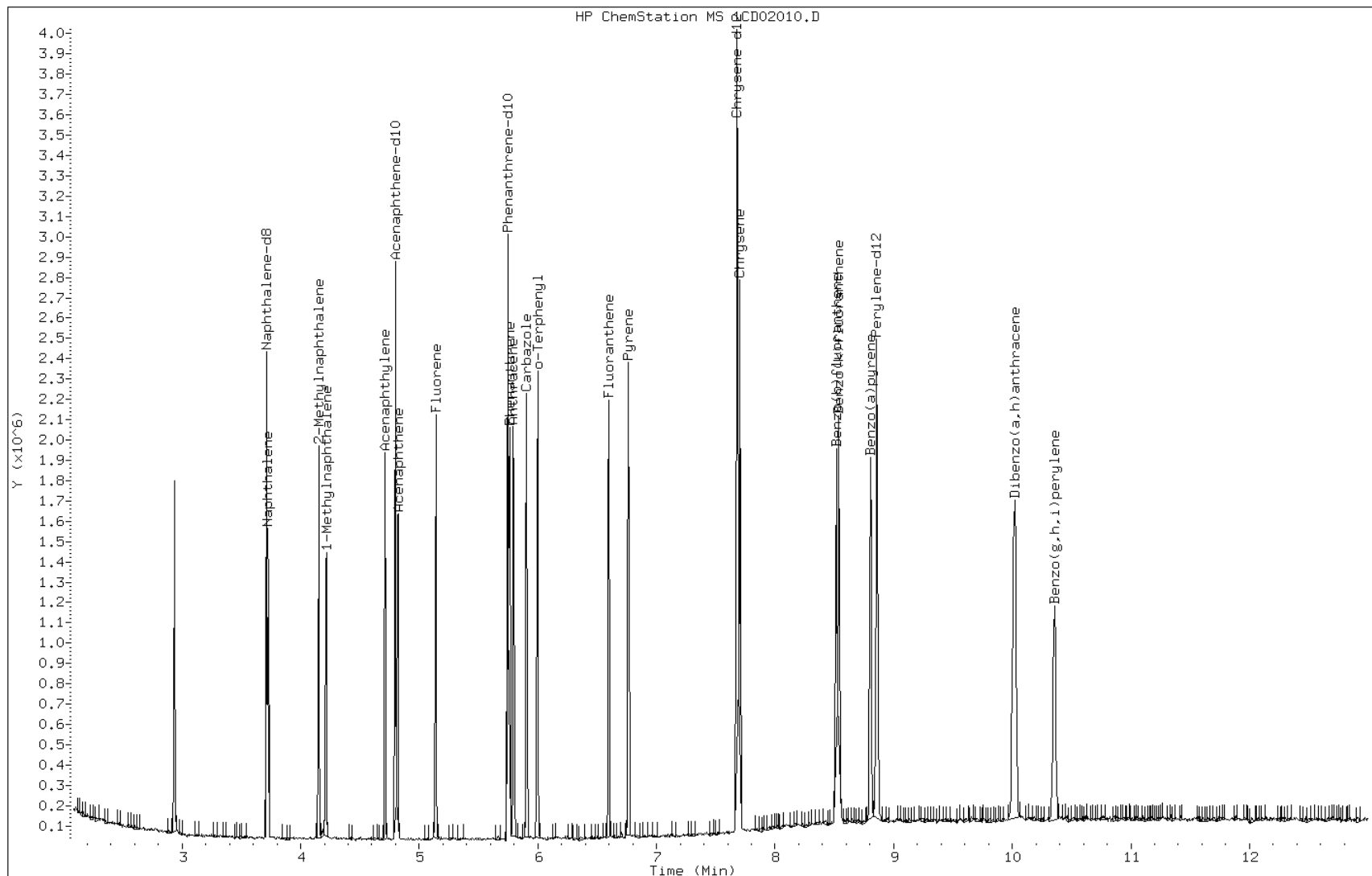
Date: 02-APR-2013 14:57

Client ID:

Instrument: BSMC5973.i

Sample Info: IC6

Operator: SCC

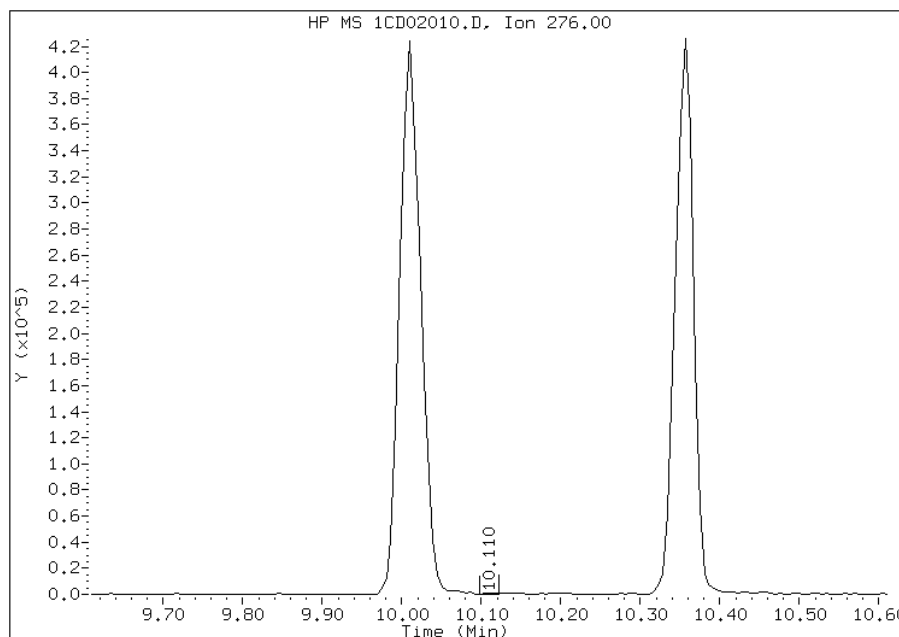


# Manual Integration Report

Data File: 1CD02010.D  
Inj. Date and Time: 02-APR-2013 14:57  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

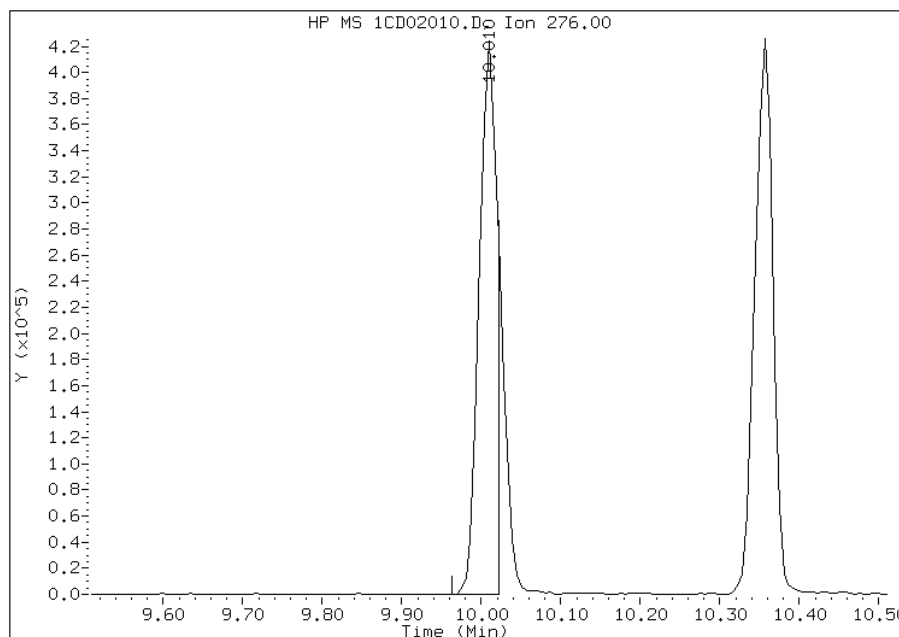
## Processing Integration Results

RT: 10.11  
Response: 1008  
Amount: 0  
Conc: 0



## Manual Integration Results

RT: 10.01  
Response: 655344  
Amount: 31  
Conc: 31



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:50  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02011.D  
 Lab Smp Id: IC7  
 Inj Date : 02-APR-2013 15:15  
 Operator : SCC  
 Smp Info : IC7  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:51 BSMC5973.i Quant Type: ISTD  
 Cal Date : 02-APR-2013 14:57 Cal File: 1CD02010.D  
 Als bottle: 11 Calibration Sample, Level: 7  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)
* 1 Naphthalene-d8	136	3.710	3.710	(1.000)	509868	40.0000	
* 6 Acenaphthene-d10	164	4.798	4.798	(1.000)	373136	40.0000	
* 10 Phenanthrene-d10	188	5.745	5.745	(1.000)	712035	40.0000	
\$ 14 o-Terphenyl	230	5.998	5.998	(1.044)	587824	50.0000	52.9755(A)
* 18 Chrysene-d12	240	7.686	7.686	(1.000)	948633	40.0000	
* 23 Perylene-d12	264	8.862	8.862	(1.000)	971909	40.0000	
2 Naphthalene	128	3.727	3.727	(1.005)	668649	50.0000	51.0580(A)
3 2-Methylnaphthalene	142	4.151	4.151	(1.119)	447751	50.0000	50.2269(A)
4 1-Methylnaphthalene	142	4.215	4.215	(1.136)	419135	50.0000	52.2523(A)
5 Acenaphthylene	152	4.710	4.710	(0.982)	814053	50.0000	52.7127(A)
7 Acenaphthene	154	4.821	4.821	(1.005)	480392	50.0000	50.2433(A)
9 Fluorene	166	5.139	5.139	(1.071)	638557	50.0000	50.0785(A)
11 Phenanthrene	178	5.762	5.762	(1.003)	1077014	50.0000	51.9349(A)
12 Anthracene	178	5.798	5.798	(1.009)	1098599	50.0000	52.2594(A)
13 Carbazole	167	5.904	5.904	(1.028)	948101	50.0000	52.6415(A)
15 Fluoranthene	202	6.598	6.598	(1.148)	1248081	50.0000	54.4959(A)
16 Pyrene	202	6.762	6.762	(0.880)	1360548	50.0000	51.7754(A)
17 Benzo(a)anthracene	228	7.680	7.680	(0.999)	1380443	50.0000	45.5615
19 Chrysene	228	7.709	7.709	(1.003)	1377767	50.0000	50.9681(AH)
20 Benzo(b)fluoranthene	252	8.521	8.521	(0.962)	1443812	50.0000	52.5467(AH)
21 Benzo(k)fluoranthene	252	8.545	8.545	(0.964)	1396501	50.0000	52.5496(AH)
22 Benzo(a)pyrene	252	8.809	8.809	(0.994)	1403971	50.0000	54.2730(A)
24 Indeno(1,2,3-cd)pyrene	276	10.015	10.015	(1.130)	1242391	50.0000	50.5646(AMH)
25 Dibenzo(a,h)anthracene	278	10.033	10.033	(1.132)	1194691	50.0000	52.6360(A)
26 Benzo(g,h,i)perylene	276	10.362	10.362	(1.169)	1270187	50.0000	50.6515(AH)

QC Flag Legend

- A - Target compound detected but, quantitated amount exceeded maximum amount.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1CD02011.D

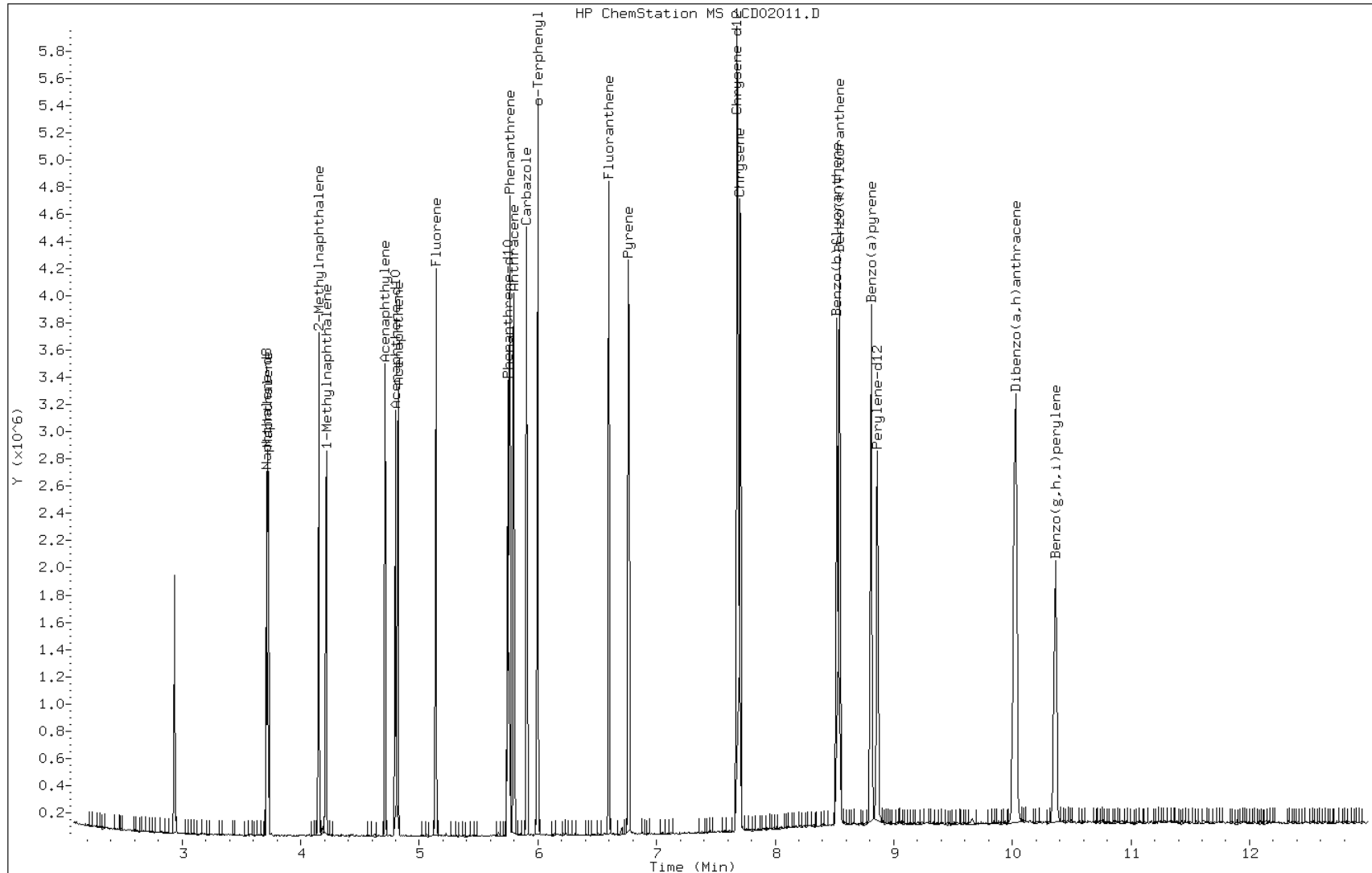
Date: 02-APR-2013 15:15

Client ID:

Instrument: BSMC5973.i

Sample Info: IC7

Operator: SCC

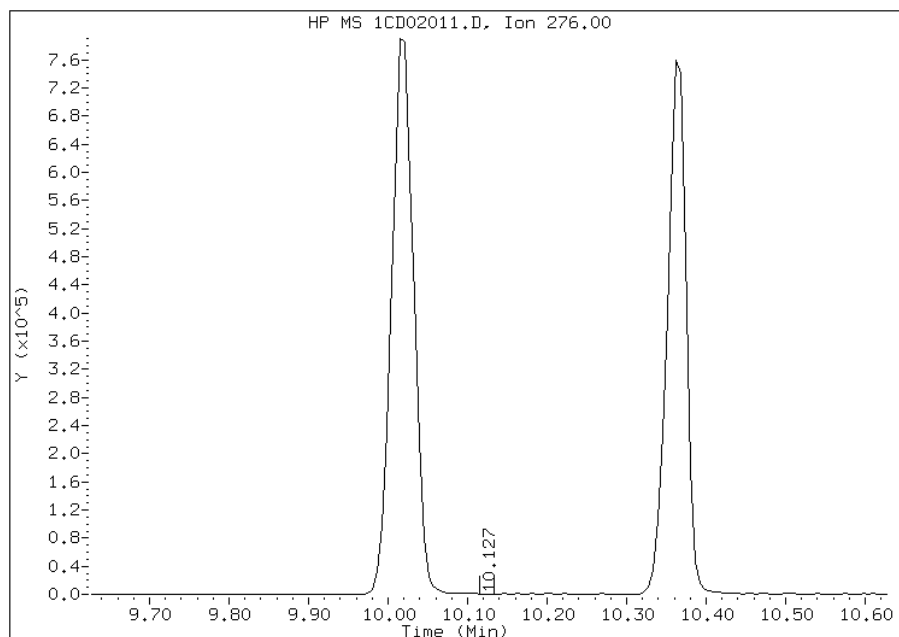


Manual Integration Report

Data File: 1CD02011.D  
Inj. Date and Time: 02-APR-2013 15:15  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

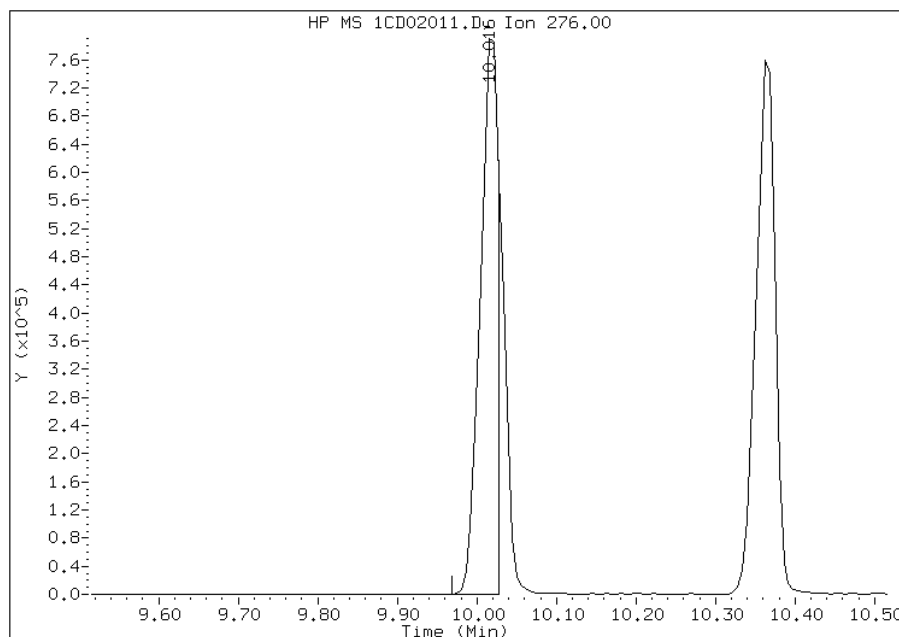
Processing Integration Results

RT: 10.13  
Response: 653  
Amount: 0  
Conc: 0



Manual Integration Results

RT: 10.02  
Response: 1242391  
Amount: 51  
Conc: 51



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:51  
Manual Integration Reason: Split Peak

FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD CURVE EVALUATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136164

SDG No.: 68088766-2

Instrument ID: BSMD5973 GC Column: DB-5MS ID: 250(um) Heated Purge: (Y/N) N

Calibration Start Date: 04/04/2013 13:49 Calibration End Date: 04/04/2013 16:04 Calibration ID: 2874

Calibration Files:

LEVEL:	LAB SAMPLE ID:	LAB FILE ID:
Level 1	IC 660-136164/15	1DD04007.D
Level 2	IC 660-136164/16	1DD04008.D
Level 3	IC 660-136164/17	1DD04009.D
Level 4	IC 660-136164/18	1DD04010.D
Level 5	ICIS 660-136164/19	1DD04011.D
Level 6	IC 660-136164/20	1DD04012.D
Level 7	IC 660-136164/21	1DD04013.D

ANALYTE	RRF					CURVE TYPE	COEFFICIENT			#	MIN RRF	%RSD	#	MAX %RSD	R <sup>2</sup> OR COD	#	MIN R <sup>2</sup> OR COD
	LVL 1	LVL 2	LVL 3	LVL 4	LVL 5		B	M1	M2								
	LVL 6	LVL 7															
Naphthalene	0.9331 1.0230	0.9606 1.0509	1.0286	0.9649	0.9984	Ave	0.9942			0.0000	4.3		15.0				
2-Methylnaphthalene	0.5806 0.6693	0.6114 0.6984	0.6517	0.6297	0.6515	Ave	0.6418			0.0000	6.0		15.0				
1-Methylnaphthalene	0.5558 0.6314	0.5782 0.6544	0.6189	0.5919	0.6119	Ave	0.6061			0.0000	5.5		15.0				
Acenaphthylene	1.4312 1.8297	1.5518 1.8878	1.7317	1.6795	1.7392	Ave	1.6930			0.0000	9.3		15.0				
Acenaphthene	1.0016 1.0873	0.9902 1.1219	1.0649	1.0164	1.0329	Ave	1.0450			0.0000	4.6		15.0				
Fluorene	1.1332 1.3072	1.1795 1.3301	1.2333	1.2265	1.2526	Ave	1.2375			0.0000	5.5		15.0				
Phenanthrene	1.0628 1.1227	1.0409 1.1914	1.1226	1.0753	1.0969	Ave	1.1018			0.0000	4.5		15.0				
Anthracene	0.9667 1.1508	1.0104 1.2102	1.1116	1.0846	1.1206	Ave	1.0936			0.0000	7.6		15.0				
Carbazole	0.8539 0.9974	0.9170 1.0575	0.9788	0.9568	0.9906	Ave	0.9646			0.0000	6.7		15.0				
Fluoranthene	1.0349 1.1765	1.0636 1.2407	1.1552	1.1188	1.1468	Ave	1.1338			0.0000	6.1		15.0				
Pyrene	1.1042 1.2400	1.1445 1.2796	1.2302	1.1952	1.2147	Ave	1.2012			0.0000	5.0		15.0				
Benzo[a]anthracene	1.5223 1.0884	1.1349 1.0935	1.1146	1.0605	1.0812	Ave	1.1565			0.0000	14.1		15.0				
Chrysene	1.1462 1.0803	1.0503 1.1335	1.0831	1.0383	1.0590	Ave	1.0844			0.0000	3.8		15.0				
Benzo[b]fluoranthene	0.9638 1.0305	0.9264 1.0697	1.0233	0.9705	1.0102	Ave	0.9992			0.0000	4.8		15.0				

Note: The m1 coefficient is the same as Ave RRF for an Ave curve type.



FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD CURVE EVALUATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136164  
 SDG No.: 68088766-2  
 Instrument ID: BSMD5973 GC Column: DB-5MS ID: 250 (um) Heated Purge: (Y/N) N  
 Calibration Start Date: 04/04/2013 13:49 Calibration End Date: 04/04/2013 16:04 Calibration ID: 2874

ANALYTE	RRF					CURVE TYPE	COEFFICIENT			#	MIN RRF	%RSD	#	MAX %RSD	R^2 OR COD	#	MIN R^2 OR COD
	LVL 1	LVL 2	LVL 3	LVL 4	LVL 5		B	M1	M2								
	LVL 6	LVL 7															
Benzo[k]fluoranthene	0.9941 1.0870	1.0278 1.1123	1.0413	1.0574	1.0488	Ave		1.0527			0.0000	3.7		15.0			
Benzo[a]pyrene	0.9363 1.0554	0.9330 1.0817	1.0086	0.9978	1.0150	Ave		1.0040			0.0000	5.5		15.0			
Indeno[1,2,3-cd]pyrene	0.9719 1.1444	1.0047 1.2203	1.0673	1.0253	1.0598	Ave		1.0705			0.0000	8.0		15.0			
Dibenz(a,h)anthracene	1.0008 1.0474	0.9200 1.0891	1.0022	0.9846	1.0127	Ave		1.0081			0.0000	5.2		15.0			
Benzo[g,h,i]perylene	0.9959 1.0588	1.0032 1.0675	1.0494	1.0184	1.0221	Ave		1.0308			0.0000	2.7		15.0			
o-Terphenyl	0.5239 0.6240	0.5611 0.6847	0.6139	0.5898	0.6214	Ave		0.6027			0.0000	8.5		15.0			

Note: The m1 coefficient is the same as Ave RRF for an Ave curve type.

FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD RESPONSE AND CONCENTRATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136164

SDG No.: 68088766-2

Instrument ID: BSMD5973 GC Column: DB-5MS ID: 250 (um) Heated Purge: (Y/N) N

Calibration Start Date: 04/04/2013 13:49 Calibration End Date: 04/04/2013 16:04 Calibration ID: 2874

Calibration Files:

LEVEL:	LAB SAMPLE ID:	LAB FILE ID:
Level 1	IC 660-136164/15	1DD04007.D
Level 2	IC 660-136164/16	1DD04008.D
Level 3	IC 660-136164/17	1DD04009.D
Level 4	IC 660-136164/18	1DD04010.D
Level 5	ICIS 660-136164/19	1DD04011.D
Level 6	IC 660-136164/20	1DD04012.D
Level 7	IC 660-136164/21	1DD04013.D

ANALYTE	IS REF	CURVE TYPE	RESPONSE					CONCENTRATION (UG/ML)				
			LVL 1 LVL 6	LVL 2 LVL 7	LVL 3	LVL 4	LVL 5	LVL 1 LVL 6	LVL 2 LVL 7	LVL 3	LVL 4	LVL 5
Naphthalene	NPT	Ave	11503 1777021	59216 3211548	316194	614716	1235557	0.200 30.0	1.00 50.0	5.00	10.0	20.0
2-Methylnaphthalene	NPT	Ave	7158 1162560	37688 2134320	200332	401151	806286	0.200 30.0	1.00 50.0	5.00	10.0	20.0
1-Methylnaphthalene	NPT	Ave	6852 1096847	35645 1999874	190230	377068	757317	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Acenaphthylene	ANT	Ave	10298 1852399	56340 3396591	314191	620756	1275622	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Acenaphthene	ANT	Ave	7207 1100779	35951 2018481	193205	375673	757590	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Fluorene	ANT	Ave	8154 1323451	42826 2393163	223769	453336	918747	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Phenanthrene	PHN	Ave	12866 1932978	63070 3534794	338739	657435	1331875	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Anthracene	PHN	Ave	11703 1981347	61222 3590722	335430	663091	1360668	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Carbazole	PHN	Ave	10338 1717245	55563 3137679	295345	584967	1202897	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Fluoranthene	PHN	Ave	12529 2025512	64445 3681257	348578	684049	1392506	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Pyrene	CRY	Ave	13274 2181708	69252 3965627	374480	738839	1496990	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[a]anthracene	CRY	Ave	18301 1914899	68675 3388838	339292	655565	1332372	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Chrysene	CRY	Ave	13779 1900592	63553 3512644	329706	641842	1305118	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[b]fluoranthene	PRY	Ave	12005 1811151	57946 3290902	323060	612455	1270704	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[k]fluoranthene	PRY	Ave	12382 1910468	64288 3421834	328752	667284	1319239	0.200 30.0	1.00 50.0	5.00	10.0	20.0

FORM VI  
GC/MS SEMI VOA INITIAL CALIBRATION DATA  
INTERNAL STANDARD RESPONSE AND CONCENTRATION

Lab Name: TestAmerica Tampa Job No.: 680-88766-2 Analy Batch No.: 136164

SDG No.: 68088766-2

Instrument ID: BSMD5973 GC Column: DB-5MS ID: 250 (um) Heated Purge: (Y/N) N

Calibration Start Date: 04/04/2013 13:49 Calibration End Date: 04/04/2013 16:04 Calibration ID: 2874

ANALYTE	IS REF	CURVE TYPE	RESPONSE					CONCENTRATION (UG/ML)				
			LVL 1	LVL 2	LVL 3	LVL 4	LVL 5	LVL 1	LVL 2	LVL 3	LVL 4	LVL 5
			LVL 6	LVL 7				LVL 6	LVL 7			
Benzo[a]pyrene	PRY	Ave	11662 1854979	58354 3327888	318431	629684	1276688	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Indeno[1,2,3-cd]pyrene	PRY	Ave	12106 2011375	62840 3754268	336963	647015	1333044	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Dibenz(a,h)anthracene	PRY	Ave	12466 1840819	57541 3350541	316396	621340	1273836	0.200 30.0	1.00 50.0	5.00	10.0	20.0
Benzo[g,h,i]perylene	PRY	Ave	12405 1860821	62750 3284166	331324	642692	1285637	0.200 30.0	1.00 50.0	5.00	10.0	20.0
o-Terphenyl	PHN	Ave	6343 1074388	33997 2031596	185249	360585	754512	0.200 30.0	1.00 50.0	5.00	10.0	20.0

Curve Type Legend:

Ave = Average ISTD

TestAmerica Laboratories

Semivolatile 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04007.D  
 Lab Smp Id: IC-1531396  
 Inj Date : 04-APR-2013 13:49  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : IC-1531396  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 16:04 Cal File: 1DD04013.D  
 Als bottle: 5 Calibration Sample, Level: 1  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	AMOUNTS	
								CAL-AMT ( ug/l)	ON-COL ( ug/l)
* 1 Naphthalene-d8	136			6.091	6.091	(1.000)	2465524	40.0000	
* 6 Acenaphthene-d10	164			7.766	7.766	(1.000)	1439075	40.0000	
* 9 Phenanthrene-d10	188			9.023	9.023	(1.000)	2421253	40.0000	
\$ 13 o-Terphenyl	230			9.329	9.329	(1.034)	6343	0.20000	0.17
* 17 Chrysene-d12	240			11.338	11.338	(1.000)	2404329	40.0000	
* 22 Perylene-d12	264			13.165	13.165	(1.000)	2491199	40.0000	
2 Naphthalene	128			6.109	6.109	(1.003)	11503	0.20000	0.19
3 2-Methylnaphthalene	142			6.814	6.814	(1.119)	7158	0.20000	0.18
4 1-Methylnaphthalene	142			6.908	6.908	(1.134)	6852	0.20000	0.18
5 Acenaphthylene	152			7.637	7.637	(0.983)	10298	0.20000	0.17
7 Acenaphthene	154			7.789	7.789	(1.003)	7207	0.20000	0.19
8 Fluorene	166			8.236	8.236	(1.061)	8154	0.20000	0.18
10 Phenanthrene	178			9.041	9.041	(1.002)	12866	0.20000	0.19
11 Anthracene	178			9.082	9.082	(1.007)	11703	0.20000	0.18
12 Carbazole	167			9.223	9.223	(1.022)	10338	0.20000	0.18
14 Fluoranthene	202			10.022	10.022	(1.111)	12529	0.20000	0.18
15 Pyrene	202			10.210	10.210	(0.901)	13274	0.20000	0.18
16 Benzo(a)anthracene	228			11.321	11.321	(0.998)	18301	0.20000	0.28
18 Chrysene	228			11.356	11.356	(1.002)	13779	0.20000	0.21
19 Benzo(b)fluoranthene	252			12.613	12.613	(0.958)	12005	0.20000	0.19
20 Benzo(k)fluoranthene	252			12.648	12.648	(0.961)	12382	0.20000	0.19
21 Benzo(a)pyrene	252			13.060	13.060	(0.992)	11662	0.20000	0.19
23 Indeno(1,2,3-cd)pyrene	276			14.734	14.734	(1.119)	12106	0.20000	0.18(M)
24 Dibenzo(a,h)anthracene	278			14.758	14.758	(1.121)	12466	0.20000	0.20(M)
25 Benzo(g,h,i)perylene	276			15.175	15.175	(1.153)	12405	0.20000	0.19

QC Flag Legend

M - Compound response manually integrated.

Data File: 1DD04007.D

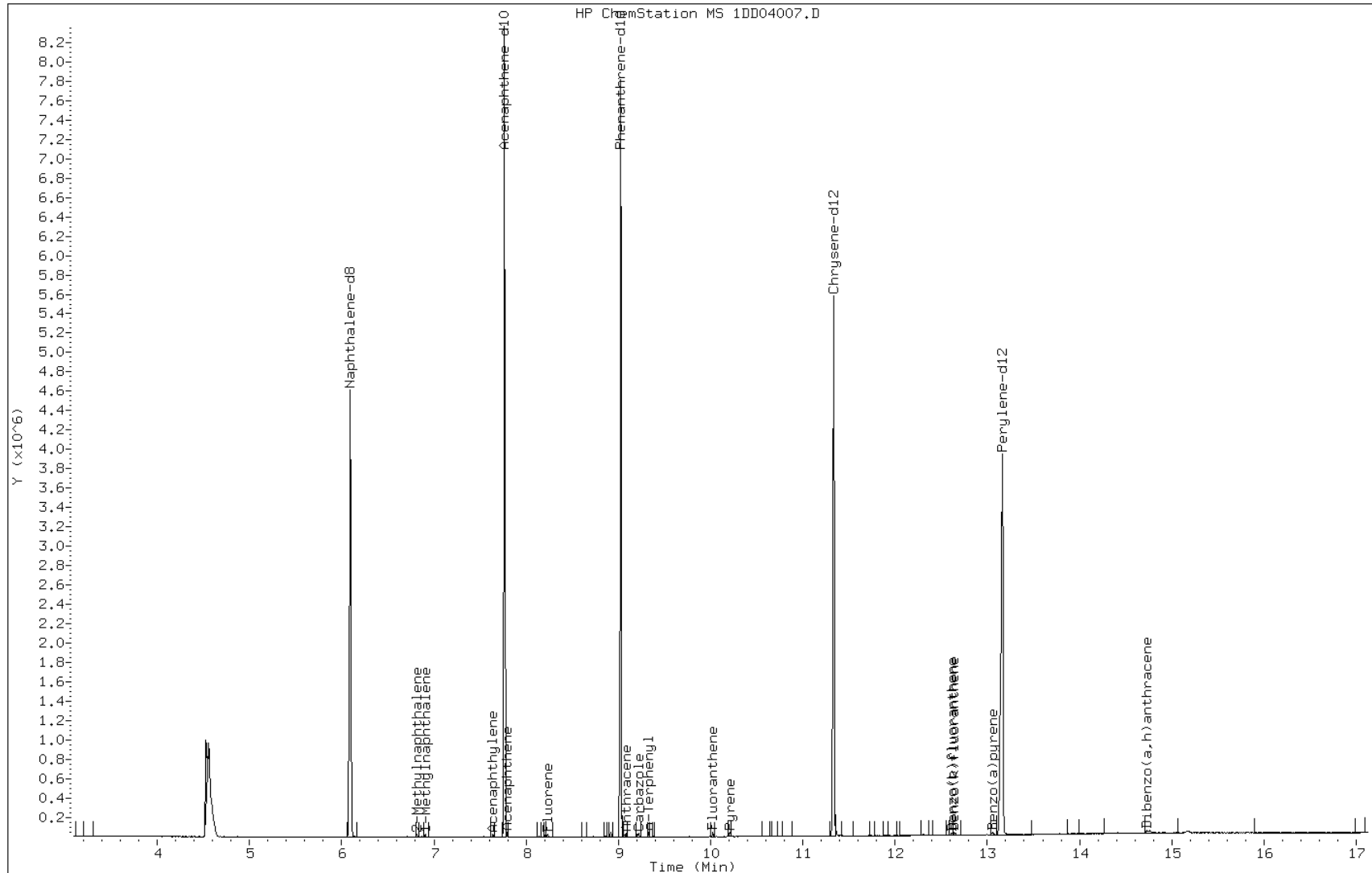
Date: 04-APR-2013 13:49

Client ID:

Instrument: BSMSD.i

Sample Info: IC-1531396

Operator: SCC

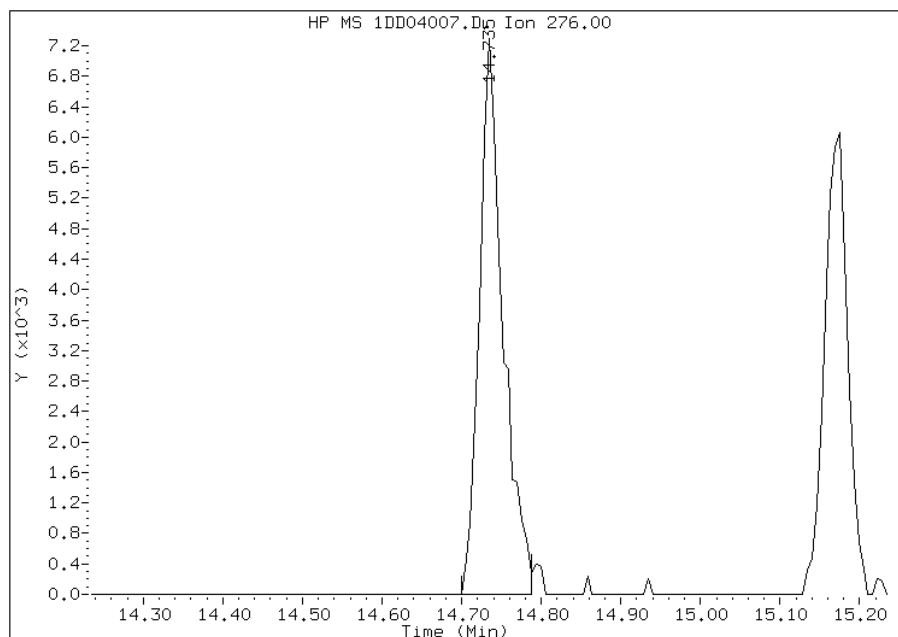


# Manual Integration Report

Data File: 1DD04007.D  
Inj. Date and Time: 04-APR-2013 13:49  
Instrument ID: BSMSD.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

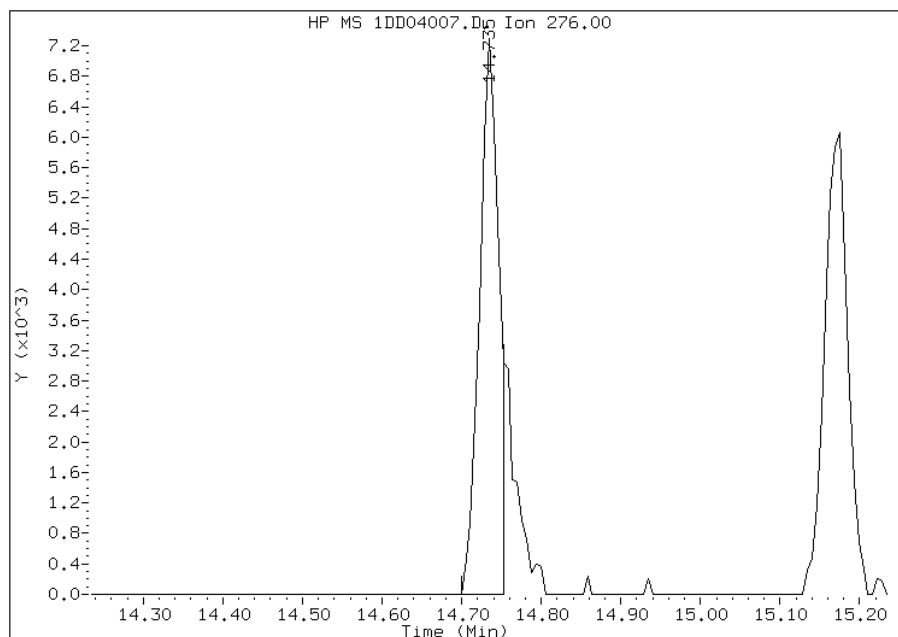
## Processing Integration Results

RT: 14.73  
Response: 14910  
Amount: 0  
Conc: 0



## Manual Integration Results

RT: 14.73  
Response: 12106  
Amount: 0  
Conc: 0



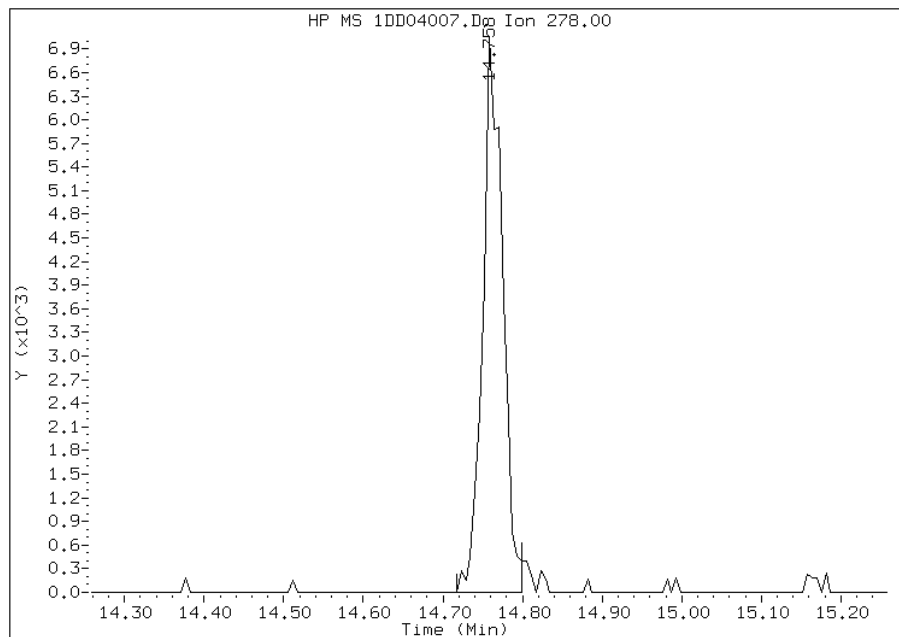
Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:28  
Manual Integration Reason: Split Peak

Manual Integration Report

Data File: 1DD04007.D  
Inj. Date and Time: 04-APR-2013 13:49  
Instrument ID: BSMSD.i  
Client ID:  
Compound: 24 Dibenzo(a,h)anthracene  
CAS #: 53-70-3  
Report Date: 04/05/2013

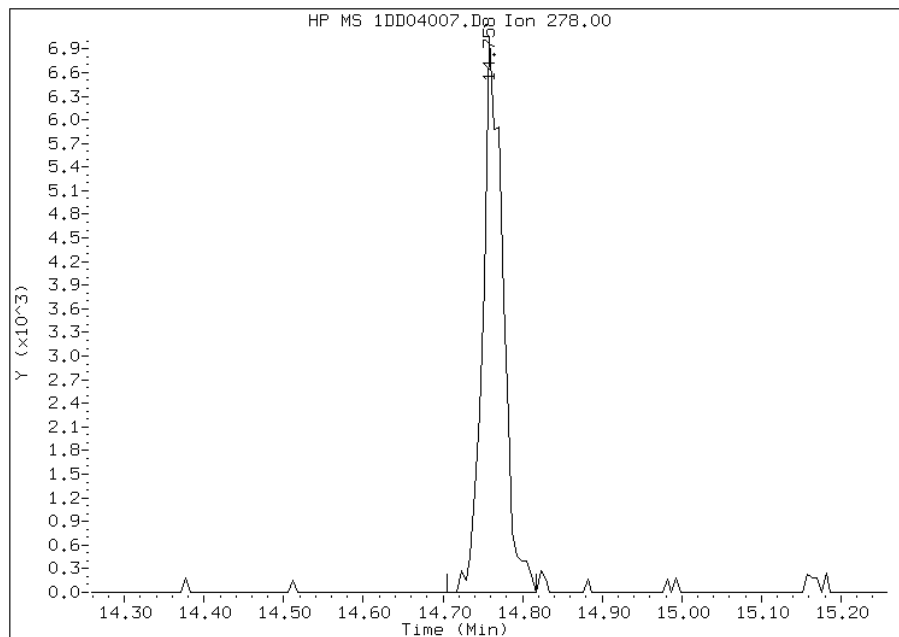
Processing Integration Results

RT: 14.76  
Response: 12250  
Amount: 0  
Conc: 0



Manual Integration Results

RT: 14.76  
Response: 12466  
Amount: 0  
Conc: 0



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:28  
Manual Integration Reason: Baseline Event

TestAmerica Laboratories

Semivolatile 8270 low level PAH

Data file : \\tam-chemsrv\chem\SM\BSMSD.i\1D040413.b\1DD04008.D  
 Lab Smp Id: IC-1531398  
 Inj Date : 04-APR-2013 14:11  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : IC-1531398  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsrv\chem\SM\BSMSD.i\1D040413.b\dFASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 13:49 Cal File: 1DD04007.D  
 Als bottle: 6 Calibration Sample, Level: 2  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG		AMOUNTS				
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT ( ug/l)	ON-COL ( ug/l)
* 1 Naphthalene-d8	136	6.089	6.089	(1.000)	2465772	40.0000	
* 6 Acenaphthene-d10	164	7.769	7.769	(1.000)	1452284	40.0000	
* 9 Phenanthrene-d10	188	9.027	9.027	(1.000)	2423707	40.0000	
\$ 13 o-Terphenyl	230	9.332	9.332	(1.034)	33997	1.00000	0.93
* 17 Chrysene-d12	240	11.336	11.336	(1.000)	2420423	40.0000	
* 22 Perylene-d12	264	13.163	13.163	(1.000)	2501899	40.0000	
2 Naphthalene	128	6.112	6.112	(1.004)	59216	1.00000	0.97
3 2-Methylnaphthalene	142	6.817	6.817	(1.120)	37688	1.00000	0.95
4 1-Methylnaphthalene	142	6.911	6.911	(1.135)	35645	1.00000	0.95
5 Acenaphthylene	152	7.640	7.640	(0.983)	56340	1.00000	0.92
7 Acenaphthene	154	7.793	7.793	(1.003)	35951	1.00000	0.95
8 Fluorene	166	8.233	8.233	(1.060)	42826	1.00000	0.95
10 Phenanthrene	178	9.038	9.038	(1.001)	63070	1.00000	0.94
11 Anthracene	178	9.080	9.080	(1.006)	61222	1.00000	0.92
12 Carbazole	167	9.221	9.221	(1.021)	55563	1.00000	0.95
14 Fluoranthene	202	10.020	10.020	(1.110)	64445	1.00000	0.94
15 Pyrene	202	10.208	10.208	(0.900)	69252	1.00000	0.95
16 Benzo(a)anthracene	228	11.318	11.318	(0.998)	68675	1.00000	1.0
18 Chrysene	228	11.359	11.359	(1.002)	63553	1.00000	0.97
19 Benzo(b)fluoranthene	252	12.611	12.611	(0.958)	57946	1.00000	0.93
20 Benzo(k)fluoranthene	252	12.646	12.646	(0.961)	64288	1.00000	0.98
21 Benzo(a)pyrene	252	13.057	13.057	(0.992)	58354	1.00000	0.93
23 Indeno(1,2,3-cd)pyrene	276	14.732	14.732	(1.119)	62840	1.00000	0.94(M)
24 Dibenzo(a,h)anthracene	278	14.761	14.761	(1.121)	57541	1.00000	0.91(M)
25 Benzo(g,h,i)perylene	276	15.167	15.167	(1.152)	62750	1.00000	0.97

QC Flag Legend

M - Compound response manually integrated.



Data File: 1DD04008.D

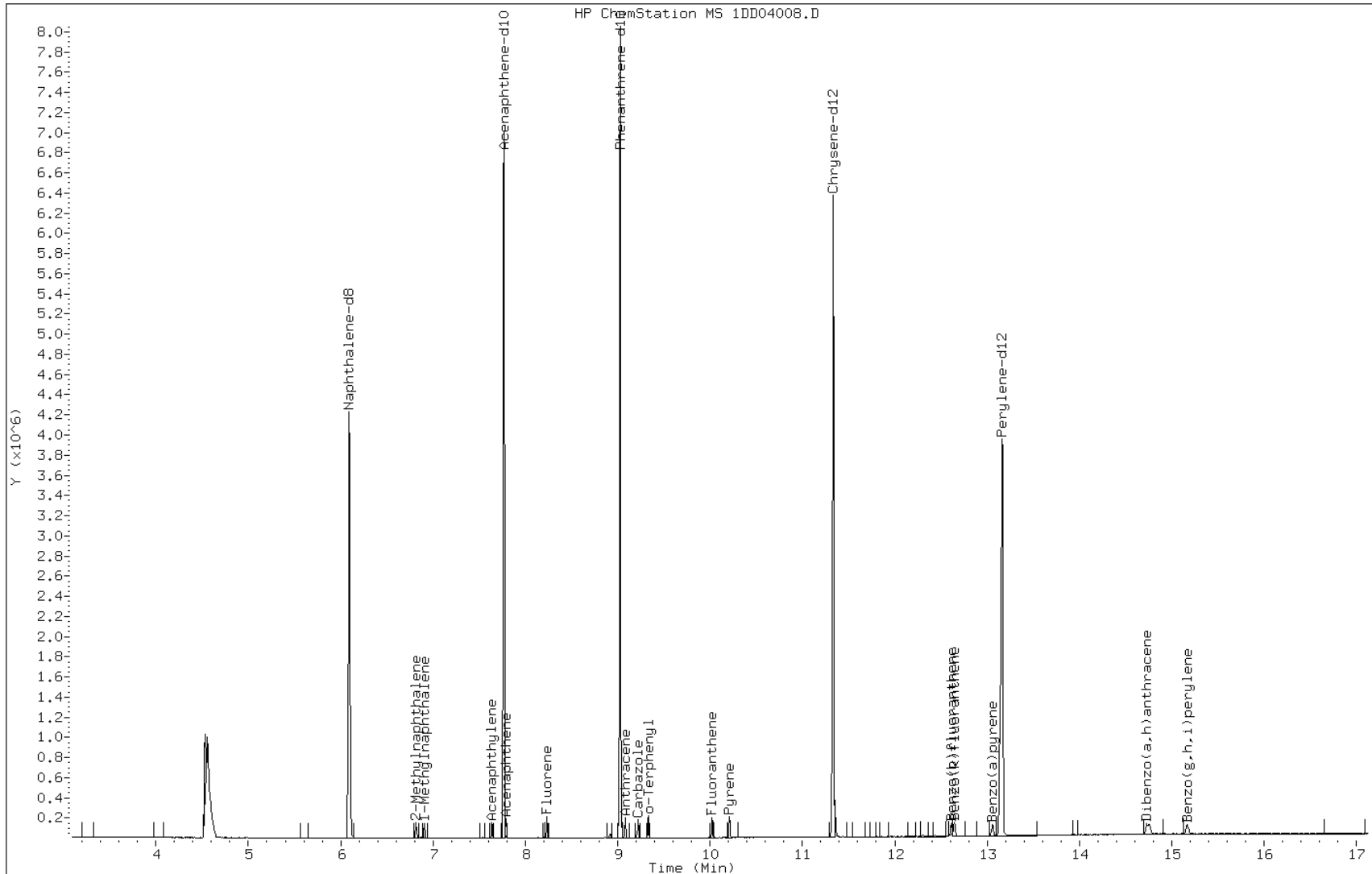
Date: 04-APR-2013 14:11

Client ID:

Instrument: BSMSD.i

Sample Info: IC-1531398

Operator: SCC

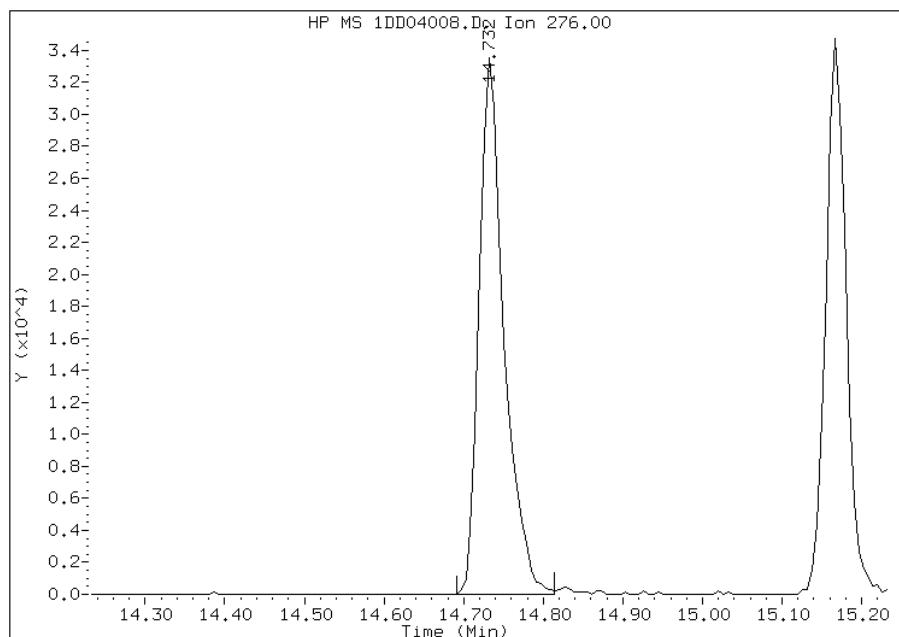


Manual Integration Report

Data File: 1DD04008.D  
Inj. Date and Time: 04-APR-2013 14:11  
Instrument ID: BSMSD.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

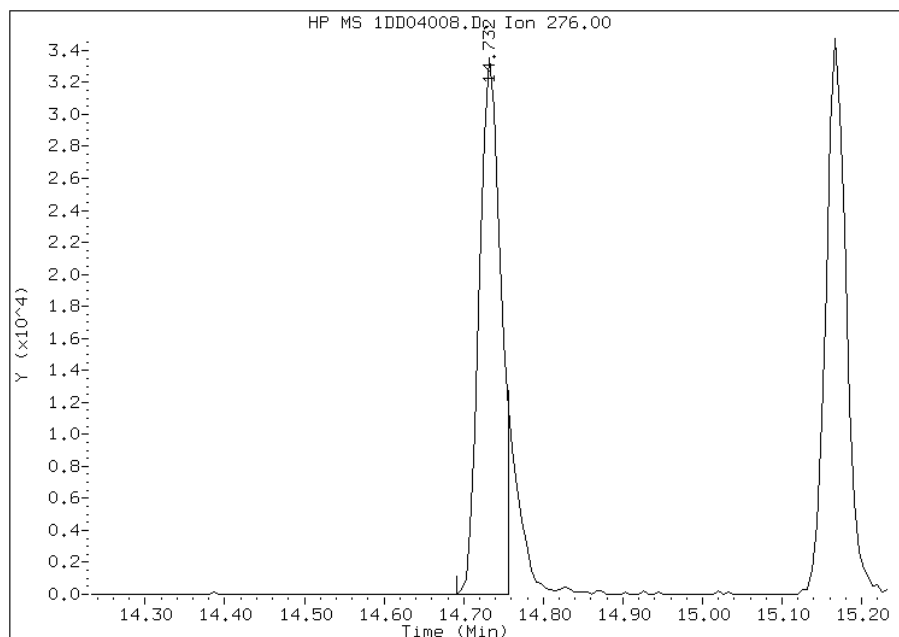
Processing Integration Results

RT: 14.73  
Response: 72512  
Amount: 1  
Conc: 1



Manual Integration Results

RT: 14.73  
Response: 62840  
Amount: 1  
Conc: 1



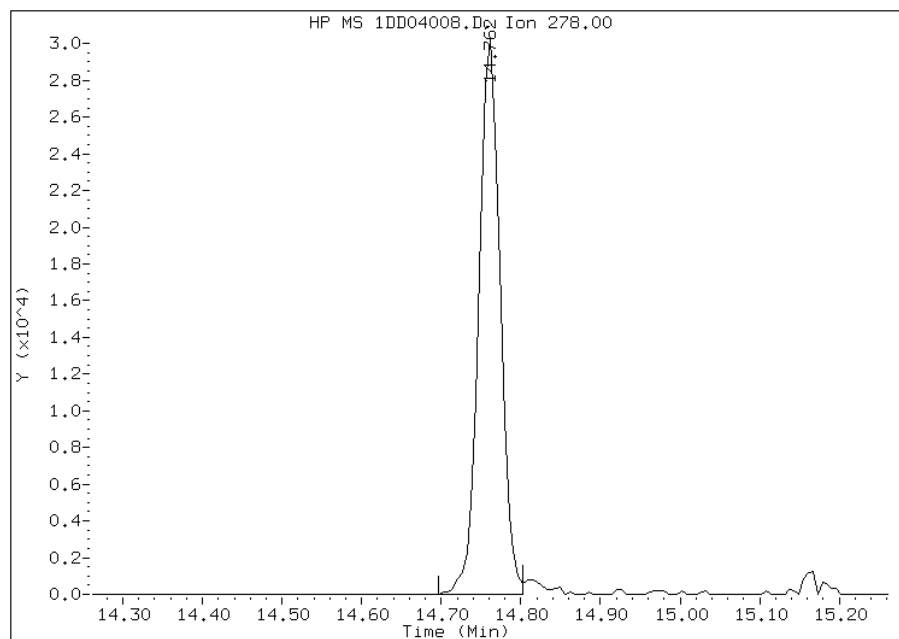
Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:29  
Manual Integration Reason: Split Peak

# Manual Integration Report

Data File: 1DD04008.D  
Inj. Date and Time: 04-APR-2013 14:11  
Instrument ID: BSMSD.i  
Client ID:  
Compound: 24 Dibenzo(a,h)anthracene  
CAS #: 53-70-3  
Report Date: 04/05/2013

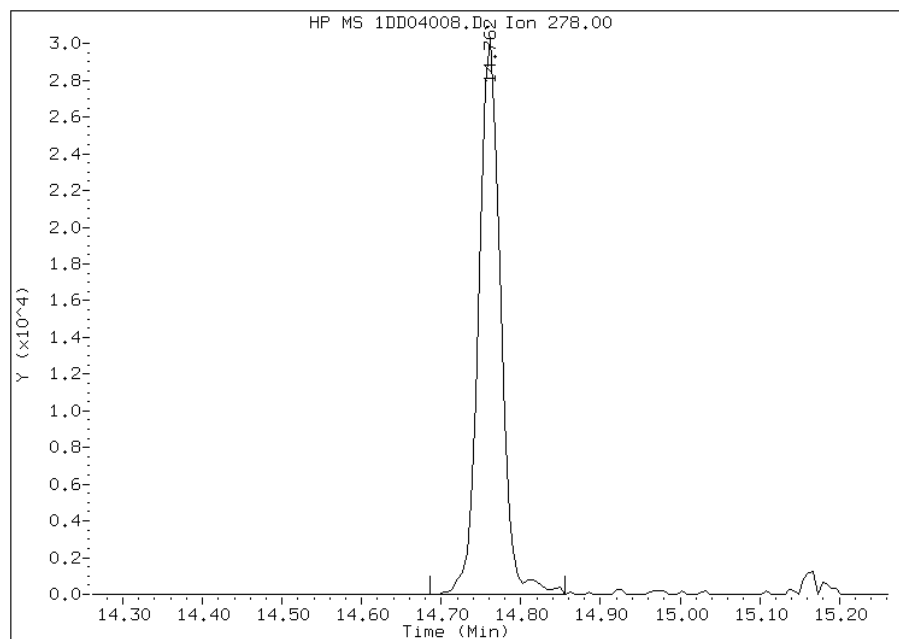
## Processing Integration Results

RT: 14.76  
Response: 56125  
Amount: 1  
Conc: 1



## Manual Integration Results

RT: 14.76  
Response: 57541  
Amount: 1  
Conc: 1



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:28  
Manual Integration Reason: Baseline Event

TestAmerica Laboratories

Semivolatile 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04009.D  
 Lab Smp Id: IC-1531399  
 Inj Date : 04-APR-2013 14:34  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : IC-1531399  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dFASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 14:11 Cal File: 1DD04008.D  
 Als bottle: 7 Calibration Sample, Level: 3  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG		AMOUNTS				
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT ( ug/l)	ON-COL ( ug/l)
* 1 Naphthalene-d8	136	6.093	6.093	(1.000)	2459101	40.0000	
* 6 Acenaphthene-d10	164	7.768	7.768	(1.000)	1451469	40.0000	
* 9 Phenanthrene-d10	188	9.025	9.025	(1.000)	2413975	40.0000	
\$ 13 o-Terphenyl	230	9.331	9.331	(1.034)	185249	5.00000	5.1
* 17 Chrysene-d12	240	11.340	11.340	(1.000)	2435324	40.0000	
* 22 Perylene-d12	264	13.167	13.167	(1.000)	2525708	40.0000	
2 Naphthalene	128	6.111	6.111	(1.003)	316194	5.00000	5.2
3 2-Methylnaphthalene	142	6.816	6.816	(1.119)	200332	5.00000	5.1
4 1-Methylnaphthalene	142	6.910	6.910	(1.134)	190230	5.00000	5.1
5 Acenaphthylene	152	7.639	7.639	(0.983)	314191	5.00000	5.1
7 Acenaphthene	154	7.791	7.791	(1.003)	193205	5.00000	5.1
8 Fluorene	166	8.232	8.232	(1.060)	223769	5.00000	5.0
10 Phenanthrene	178	9.043	9.043	(1.002)	338739	5.00000	5.1
11 Anthracene	178	9.084	9.084	(1.007)	335430	5.00000	5.1
12 Carbazole	167	9.219	9.219	(1.021)	295345	5.00000	5.1
14 Fluoranthene	202	10.024	10.024	(1.111)	348578	5.00000	5.1
15 Pyrene	202	10.212	10.212	(0.901)	374480	5.00000	5.1
16 Benzo(a)anthracene	228	11.323	11.323	(0.998)	339292	5.00000	5.1
18 Chrysene	228	11.358	11.358	(1.002)	329706	5.00000	5.0
19 Benzo(b)fluoranthene	252	12.615	12.615	(0.958)	323060	5.00000	5.1
20 Benzo(k)fluoranthene	252	12.650	12.650	(0.961)	328752	5.00000	4.9
21 Benzo(a)pyrene	252	13.062	13.062	(0.992)	318431	5.00000	5.0
23 Indeno(1,2,3-cd)pyrene	276	14.742	14.742	(1.120)	336963	5.00000	5.0(M)
24 Dibenzo(a,h)anthracene	278	14.766	14.766	(1.121)	316396	5.00000	5.0
25 Benzo(g,h,i)perylene	276	15.177	15.177	(1.153)	331324	5.00000	5.1

QC Flag Legend

M - Compound response manually integrated.

Data File: 1DD04009.D

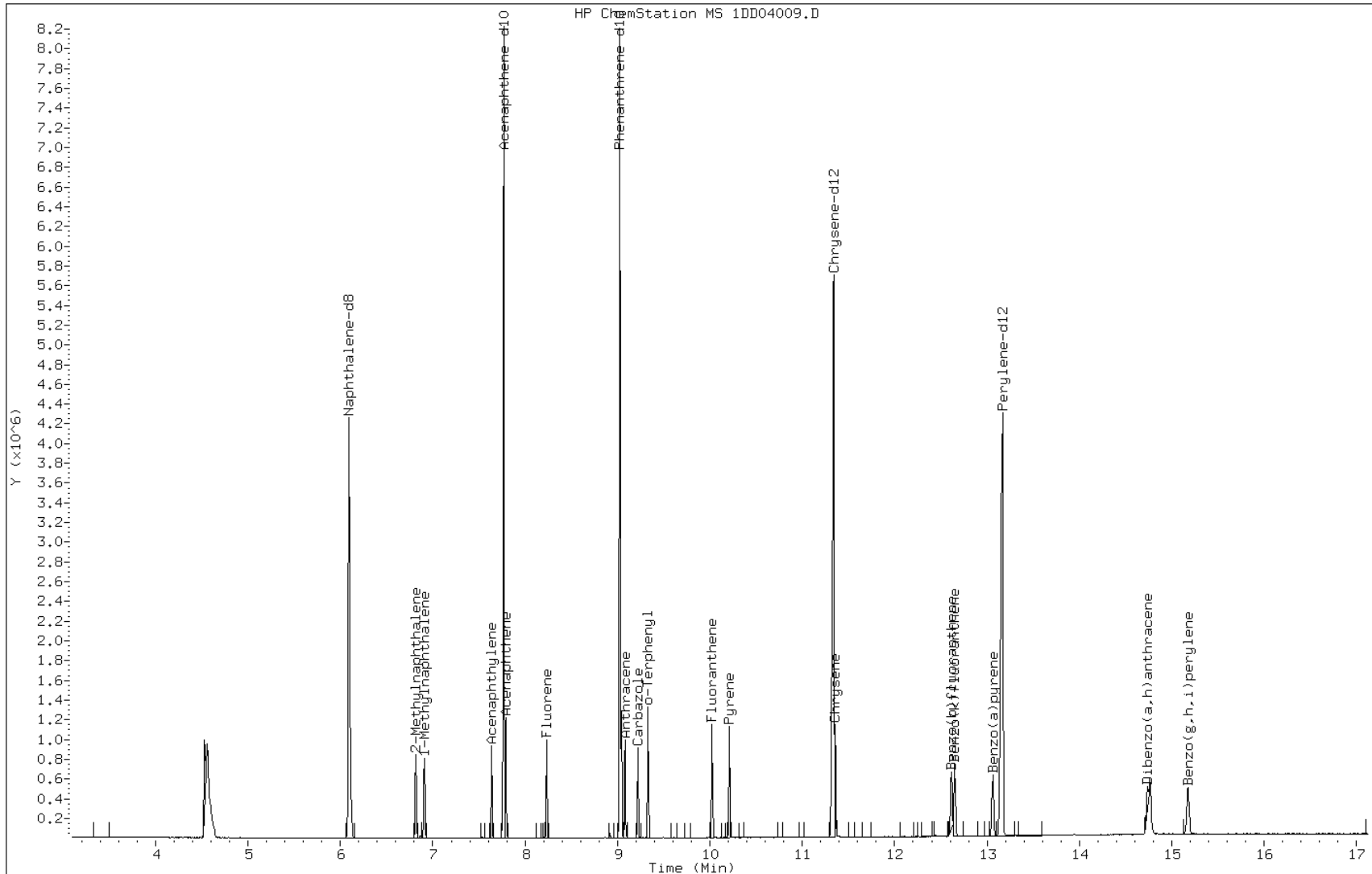
Date: 04-APR-2013 14:34

Client ID:

Instrument: BSMSD.i

Sample Info: IC-1531399

Operator: SCC

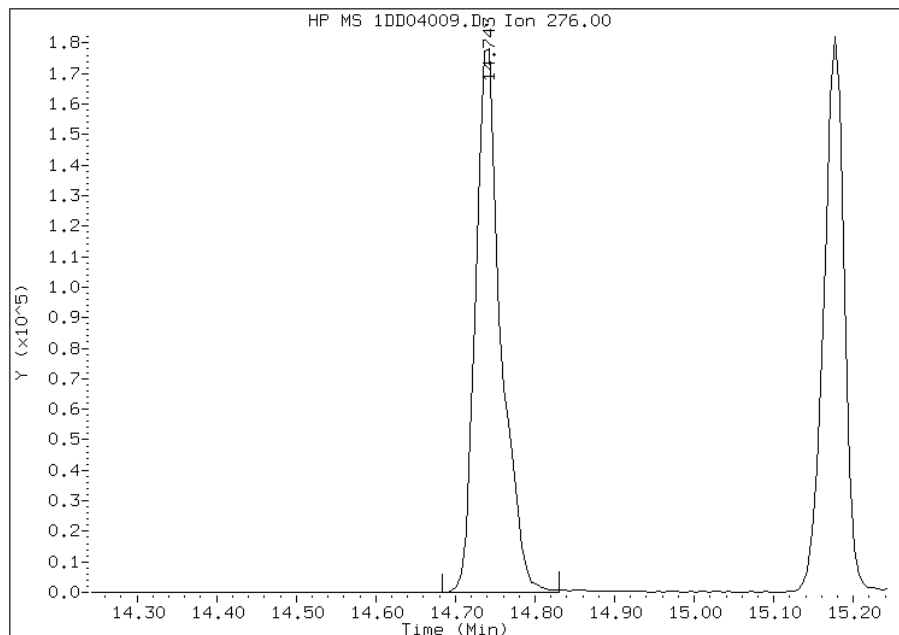


# Manual Integration Report

Data File: 1DD04009.D  
Inj. Date and Time: 04-APR-2013 14:34  
Instrument ID: BSM5D.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

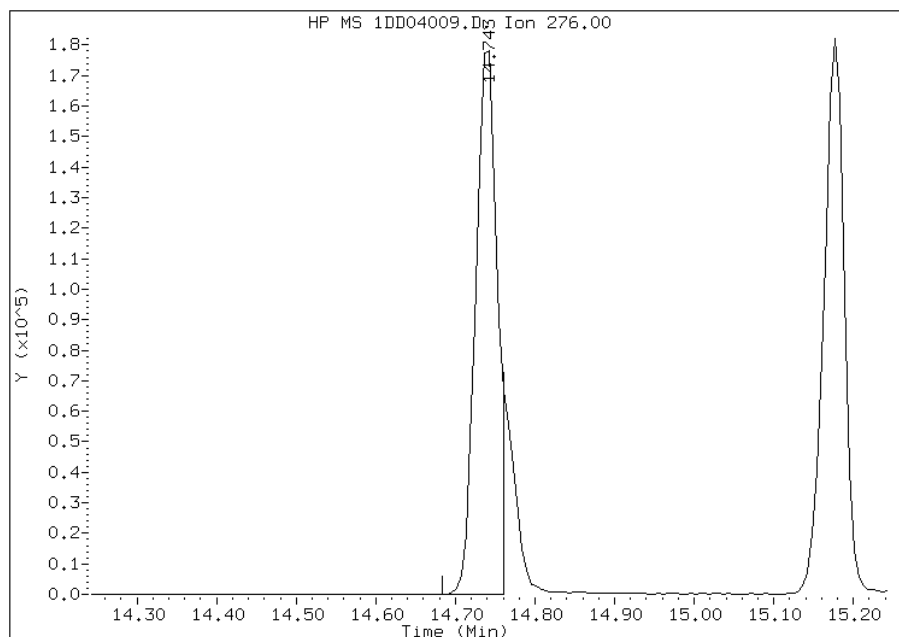
## Processing Integration Results

RT: 14.74  
Response: 395308  
Amount: 5  
Conc: 5



## Manual Integration Results

RT: 14.74  
Response: 336963  
Amount: 5  
Conc: 5



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:29  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatiles 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04010.D  
 Lab Smp Id: IC-1531400  
 Inj Date : 04-APR-2013 14:57  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : IC-1531400  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 14:34 Cal File: 1DD04009.D  
 Als bottle: 8 Calibration Sample, Level: 4  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG		AMOUNTS				
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT ( ug/l)	ON-COL ( ug/l)
* 1 Naphthalene-d8	136	6.093	6.093	(1.000)	2548377	40.0000	
* 6 Acenaphthene-d10	164	7.767	7.767	(1.000)	1478460	40.0000	
* 9 Phenanthrene-d10	188	9.025	9.025	(1.000)	2445573	40.0000	
\$ 13 o-Terphenyl	230	9.330	9.330	(1.034)	360585	10.0000	9.8
* 17 Chrysene-d12	240	11.340	11.340	(1.000)	2472736	40.0000	
* 22 Perylene-d12	264	13.167	13.167	(1.000)	2524268	40.0000	
2 Naphthalene	128	6.110	6.110	(1.003)	614716	10.0000	9.7
3 2-Methylnaphthalene	142	6.816	6.816	(1.119)	401151	10.0000	9.8
4 1-Methylnaphthalene	142	6.910	6.910	(1.134)	377068	10.0000	9.8
5 Acenaphthylene	152	7.638	7.638	(0.983)	620756	10.0000	9.9
7 Acenaphthene	154	7.791	7.791	(1.003)	375673	10.0000	9.7
8 Fluorene	166	8.237	8.237	(1.061)	453336	10.0000	9.9
10 Phenanthrene	178	9.042	9.042	(1.002)	657435	10.0000	9.8
11 Anthracene	178	9.083	9.083	(1.007)	663091	10.0000	9.9
12 Carbazole	167	9.224	9.224	(1.022)	584967	10.0000	9.9
14 Fluoranthene	202	10.024	10.024	(1.111)	684049	10.0000	9.9
15 Pyrene	202	10.212	10.212	(0.901)	738839	10.0000	9.9
16 Benzo(a)anthracene	228	11.322	11.322	(0.998)	655565	10.0000	9.7
18 Chrysene	228	11.363	11.363	(1.002)	641842	10.0000	9.6
19 Benzo(b)fluoranthene	252	12.621	12.621	(0.959)	612455	10.0000	9.7
20 Benzo(k)fluoranthene	252	12.656	12.656	(0.961)	667284	10.0000	10
21 Benzo(a)pyrene	252	13.067	13.067	(0.992)	629684	10.0000	9.9
23 Indeno(1,2,3-cd)pyrene	276	14.747	14.747	(1.120)	647015	10.0000	9.6(M)
24 Dibenzo(a,h)anthracene	278	14.777	14.777	(1.122)	621340	10.0000	9.8
25 Benzo(g,h,i)perylene	276	15.188	15.188	(1.153)	642692	10.0000	9.9

QC Flag Legend

M - Compound response manually integrated.

Data File: 1DD04010.D

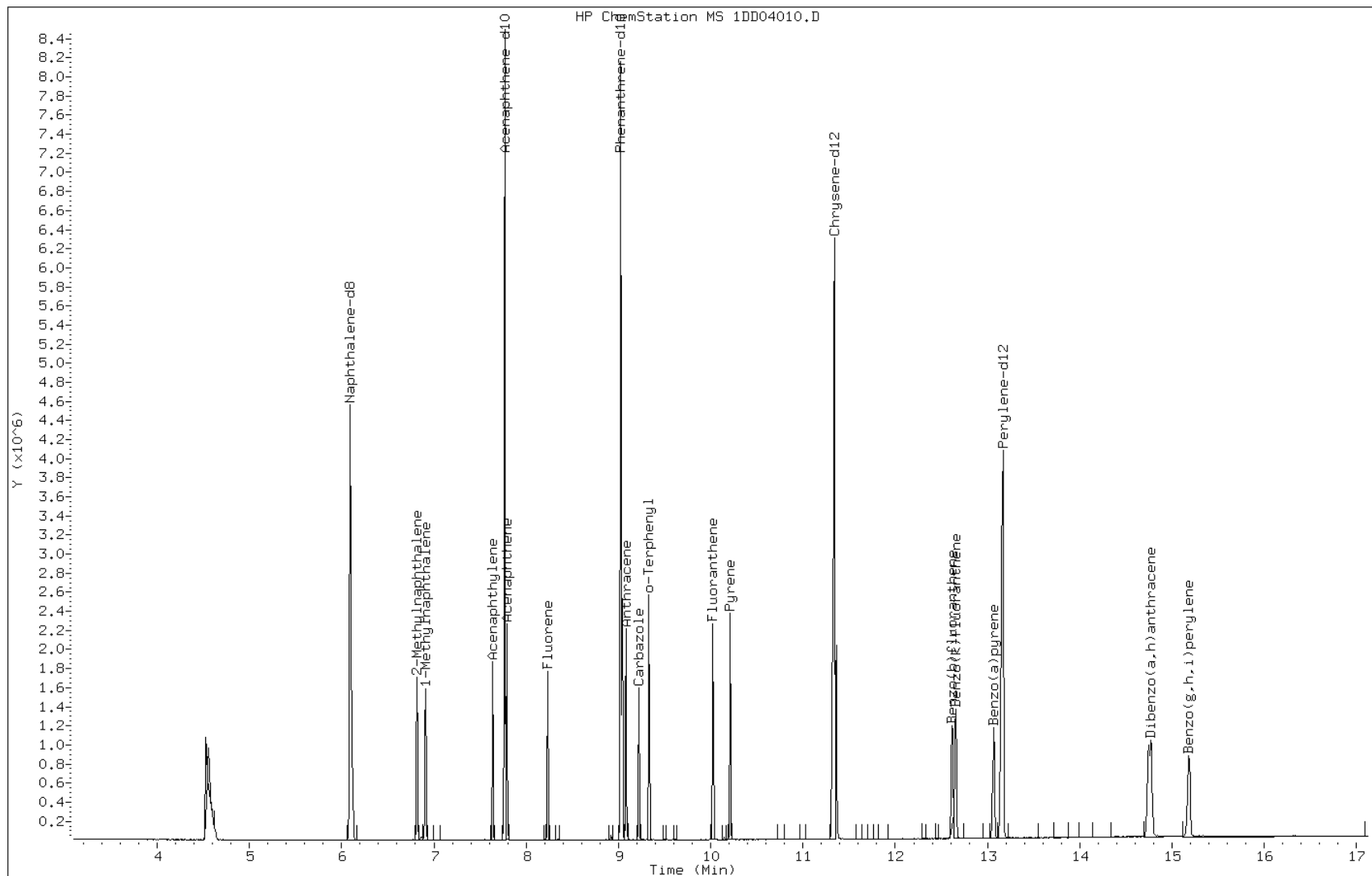
Date: 04-APR-2013 14:57

Client ID:

Instrument: BSMSD.i

Sample Info: IC-1531400

Operator: SCC



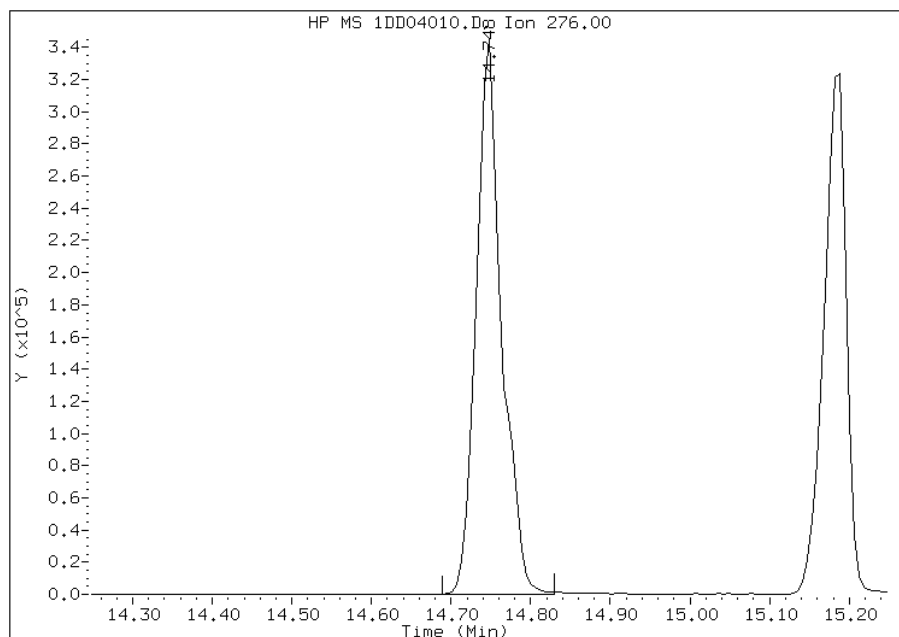


Manual Integration Report

Data File: 1DD04010.D  
Inj. Date and Time: 04-APR-2013 14:57  
Instrument ID: BSMSSD.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

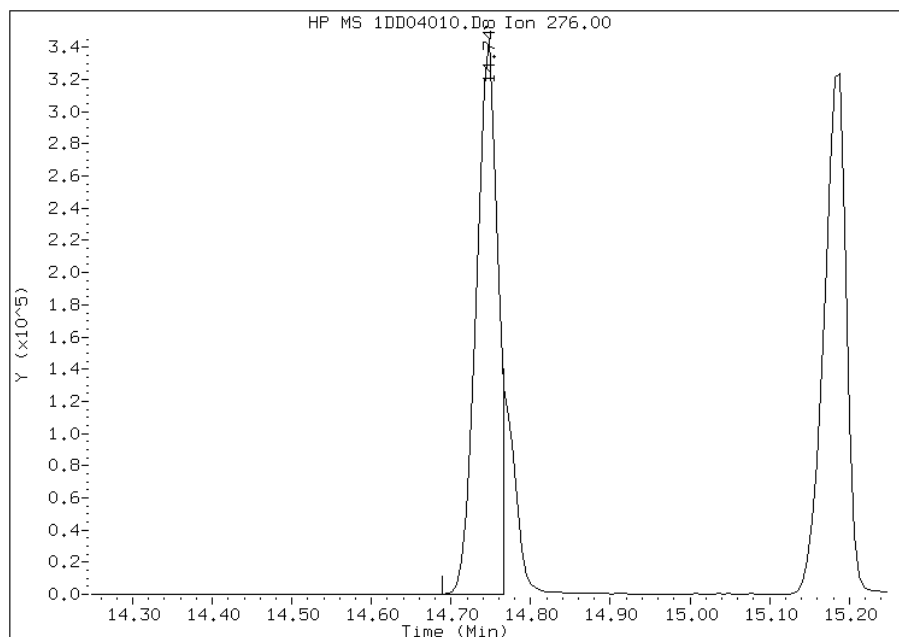
Processing Integration Results

RT: 14.75  
Response: 759012  
Amount: 10  
Conc: 10



Manual Integration Results

RT: 14.75  
Response: 647015  
Amount: 10  
Conc: 10



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:30  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatile 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04011.D  
 Lab Smp Id: ICIS-1531401  
 Inj Date : 04-APR-2013 15:19  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : ICIS-1531401  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 14:57 Cal File: 1DD04010.D  
 Als bottle: 9 Calibration Sample, Level: 5  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT	SIG	AMOUNTS					ON-COL
			MASS	RT	EXP RT	REL RT	RESPONSE	
* 1 Naphthalene-d8	136		6.089	6.089	(1.000)	2475113	40.0000	
* 6 Acenaphthene-d10	164		7.769	7.769	(1.000)	1466924	40.0000	
* 9 Phenanthrene-d10	188		9.027	9.027	(1.000)	2428512	40.0000	
\$ 13 o-Terphenyl	230		9.332	9.332	(1.034)	754512	20.0000	21
* 17 Chrysene-d12	240		11.342	11.342	(1.000)	2464730	40.0000	
* 22 Perylene-d12	264		13.169	13.169	(1.000)	2515643	40.0000	
2 Naphthalene	128		6.113	6.113	(1.004)	1235557	20.0000	20
3 2-Methylnaphthalene	142		6.818	6.818	(1.120)	806286	20.0000	20
4 1-Methylnaphthalene	142		6.912	6.912	(1.135)	757317	20.0000	20
5 Acenaphthylene	152		7.640	7.640	(0.983)	1275622	20.0000	20
7 Acenaphthene	154		7.793	7.793	(1.003)	757590	20.0000	20
8 Fluorene	166		8.234	8.234	(1.060)	918747	20.0000	20
10 Phenanthrene	178		9.044	9.044	(1.002)	1331875	20.0000	20
11 Anthracene	178		9.086	9.086	(1.007)	1360668	20.0000	20
12 Carbazole	167		9.227	9.227	(1.022)	1202897	20.0000	20
14 Fluoranthene	202		10.026	10.026	(1.111)	1392506	20.0000	20
15 Pyrene	202		10.214	10.214	(0.901)	1496990	20.0000	20
16 Benzo(a)anthracene	228		11.324	11.324	(0.998)	1332372	20.0000	20
18 Chrysene	228		11.365	11.365	(1.002)	1305118	20.0000	20
19 Benzo(b)fluoranthene	252		12.623	12.623	(0.959)	1270704	20.0000	20
20 Benzo(k)fluoranthene	252		12.664	12.664	(0.962)	1319239	20.0000	20
21 Benzo(a)pyrene	252		13.075	13.075	(0.993)	1276688	20.0000	20
23 Indeno(1,2,3-cd)pyrene	276		14.761	14.761	(1.121)	1333044	20.0000	20(M)
24 Dibenzo(a,h)anthracene	278		14.785	14.785	(1.123)	1273836	20.0000	20
25 Benzo(g,h,i)perylene	276		15.202	15.202	(1.154)	1285637	20.0000	20

QC Flag Legend

M - Compound response manually integrated.

Data File: 1DD04011.D

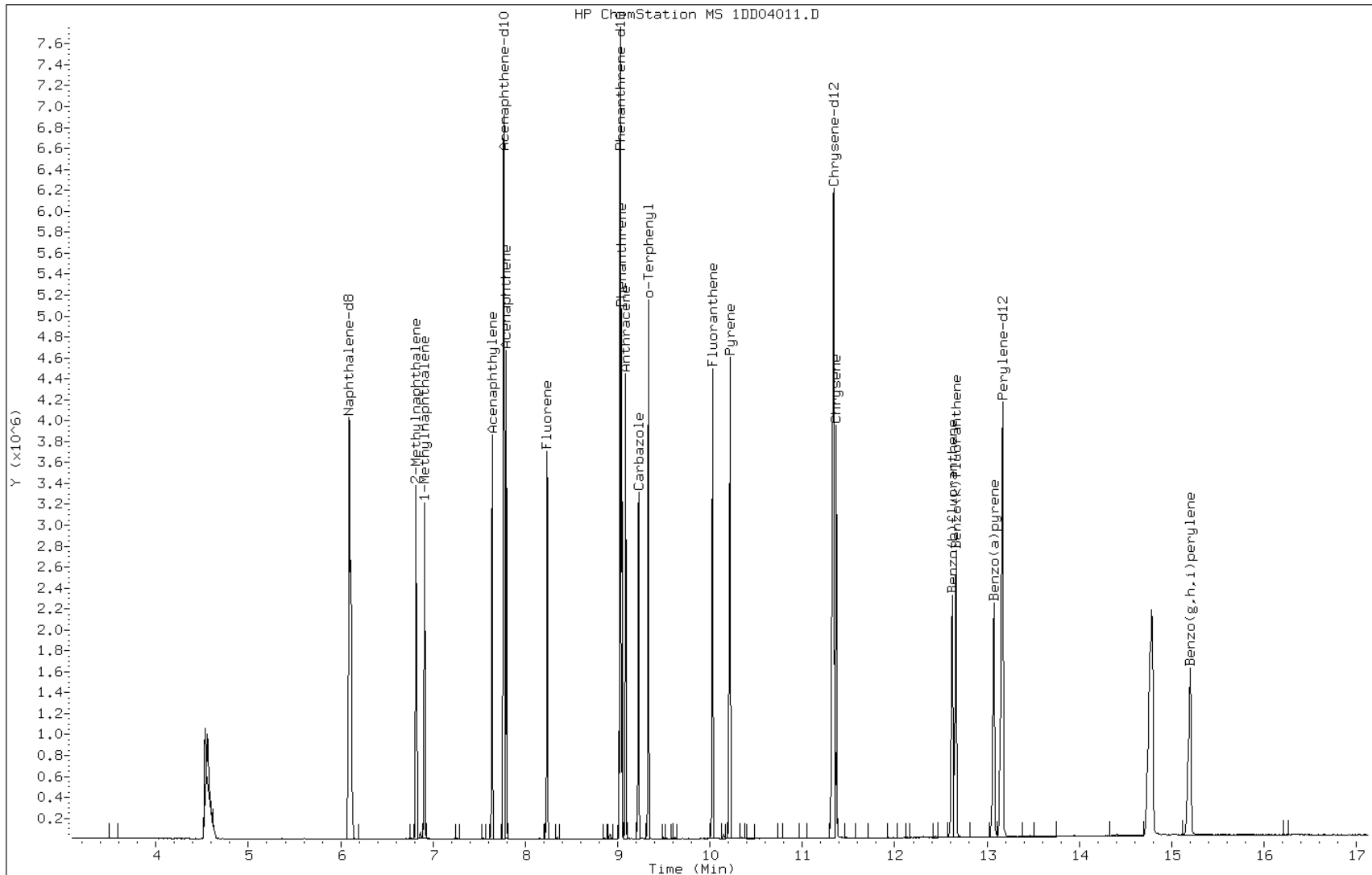
Date: 04-APR-2013 15:19

Client ID:

Instrument: BSMSD.i

Sample Info: ICIS-1531401

Operator: SCC

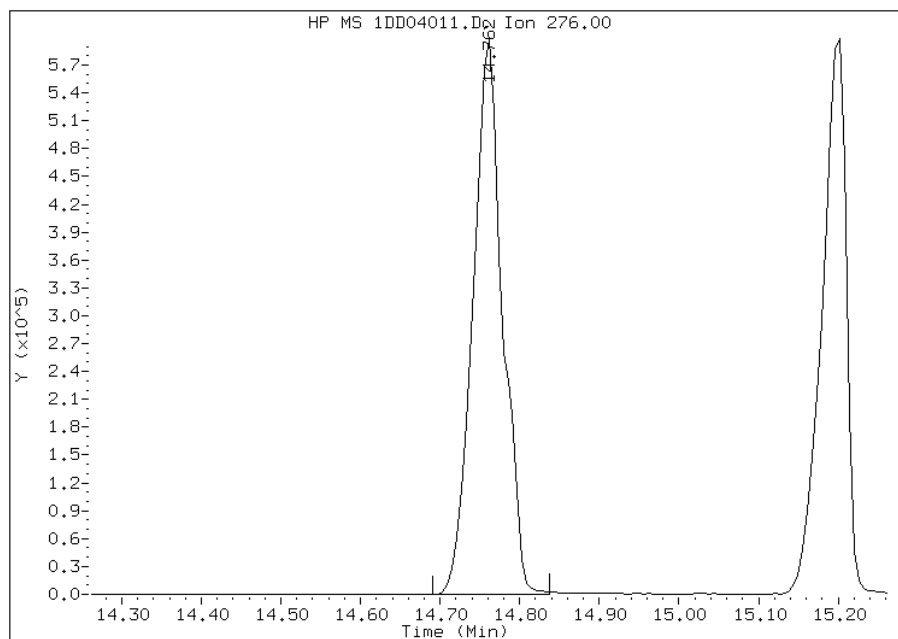


# Manual Integration Report

Data File: 1DD04011.D  
Inj. Date and Time: 04-APR-2013 15:19  
Instrument ID: BSMSD.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

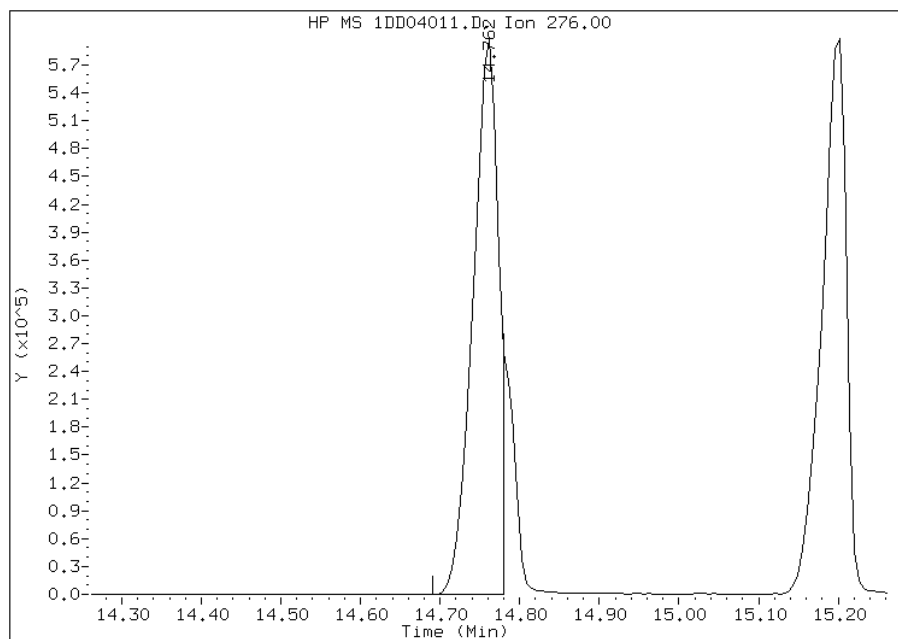
## Processing Integration Results

RT: 14.76  
Response: 1546230  
Amount: 22  
Conc: 22



## Manual Integration Results

RT: 14.76  
Response: 1333044  
Amount: 20  
Conc: 20



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:26  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatiles 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04012.D  
 Lab Smp Id: IC-1531402  
 Inj Date : 04-APR-2013 15:42  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : IC-1531402  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 15:19 Cal File: 1DD04011.D  
 Als bottle: 10 Calibration Sample, Level: 6  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG		AMOUNTS				
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT ( ug/l)	ON-COL ( ug/l)
* 1 Naphthalene-d8	136	6.090	6.090	(1.000)	2316091	40.0000	
* 6 Acenaphthene-d10	164	7.765	7.765	(1.000)	1349878	40.0000	
* 9 Phenanthrene-d10	188	9.028	9.028	(1.000)	2295562	40.0000	
\$ 13 o-Terphenyl	230	9.334	9.334	(1.034)	1074388	30.0000	31
* 17 Chrysene-d12	240	11.343	11.343	(1.000)	2345845	40.0000	
* 22 Perylene-d12	264	13.170	13.170	(1.000)	2343379	40.0000	
2 Naphthalene	128	6.114	6.114	(1.004)	1777021	30.0000	31
3 2-Methylnaphthalene	142	6.819	6.819	(1.120)	1162560	30.0000	31
4 1-Methylnaphthalene	142	6.913	6.913	(1.135)	1096847	30.0000	31
5 Acenaphthylene	152	7.642	7.642	(0.984)	1852399	30.0000	32
7 Acenaphthene	154	7.794	7.794	(1.004)	1100779	30.0000	31
8 Fluorene	166	8.235	8.235	(1.061)	1323451	30.0000	32
10 Phenanthrene	178	9.046	9.046	(1.002)	1932978	30.0000	30
11 Anthracene	178	9.087	9.087	(1.007)	1981347	30.0000	32
12 Carbazole	167	9.228	9.228	(1.022)	1717245	30.0000	31
14 Fluoranthene	202	10.027	10.027	(1.111)	2025512	30.0000	31
15 Pyrene	202	10.215	10.215	(0.901)	2181708	30.0000	31
16 Benzo(a)anthracene	228	11.326	11.326	(0.998)	1914899	30.0000	30
18 Chrysene	228	11.367	11.367	(1.002)	1900592	30.0000	30
19 Benzo(b)fluoranthene	252	12.630	12.630	(0.959)	1811151	30.0000	31
20 Benzo(k)fluoranthene	252	12.671	12.671	(0.962)	1910468	30.0000	31
21 Benzo(a)pyrene	252	13.082	13.082	(0.993)	1854979	30.0000	32
23 Indeno(1,2,3-cd)pyrene	276	14.769	14.769	(1.121)	2011375	30.0000	32(M)
24 Dibenzo(a,h)anthracene	278	14.798	14.798	(1.124)	1840819	30.0000	31
25 Benzo(g,h,i)perylene	276	15.209	15.209	(1.155)	1860821	30.0000	31

QC Flag Legend

M - Compound response manually integrated.

Data File: 1DD04012.D

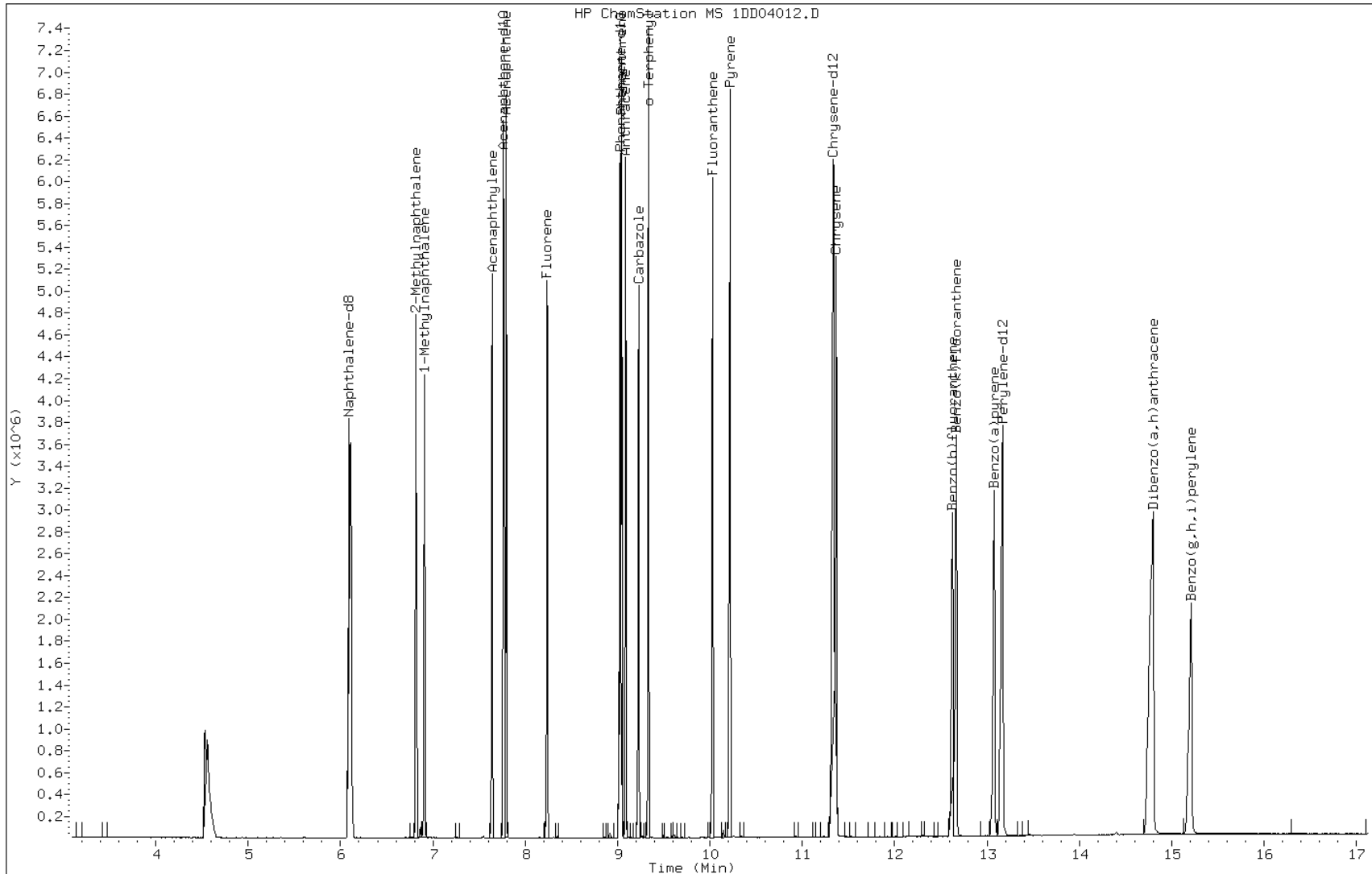
Date: 04-APR-2013 15:42

Client ID:

Instrument: BSMSD.i

Sample Info: IC-1531402

Operator: SCC

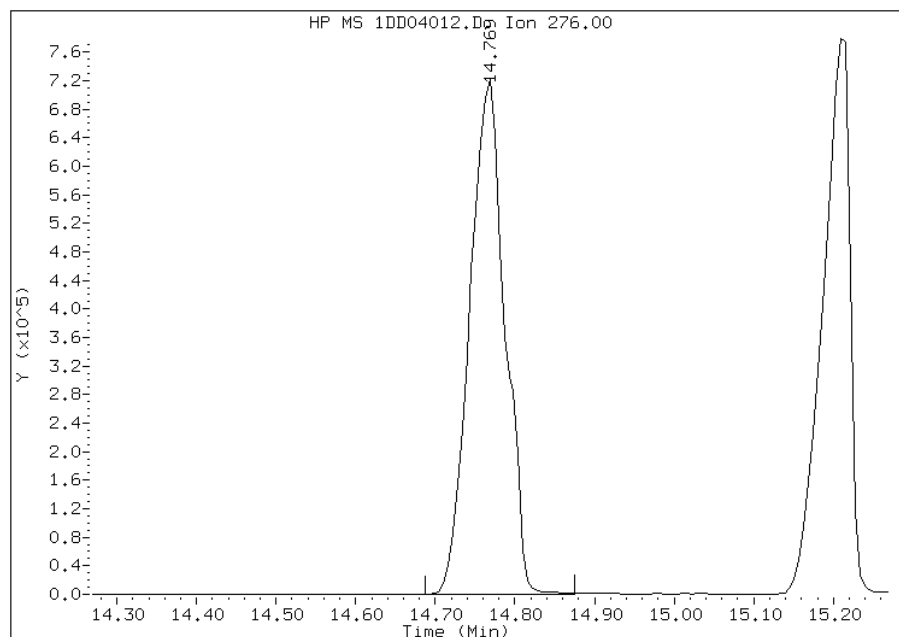


# Manual Integration Report

Data File: 1DD04012.D  
Inj. Date and Time: 04-APR-2013 15:42  
Instrument ID: BSMMSD.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

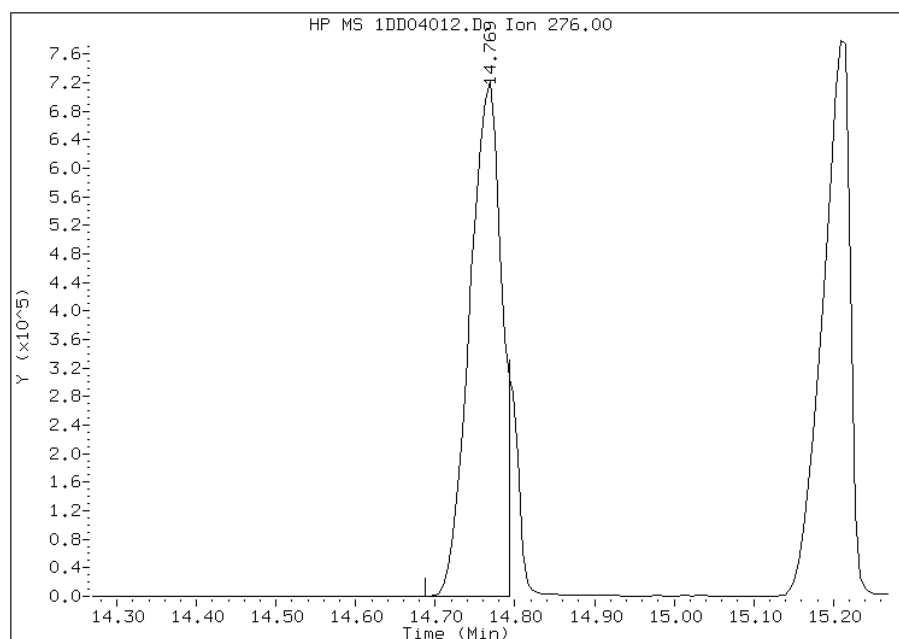
## Processing Integration Results

RT: 14.77  
Response: 2221522  
Amount: 32  
Conc: 32



## Manual Integration Results

RT: 14.77  
Response: 2011375  
Amount: 32  
Conc: 32



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:30  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Semivolatile 8270 low level PAH

Data file : \\tam-chemsrv\chem\SM\BSMSD.i\1D040413.b\1DD04013.D  
 Lab Smp Id: IC-1531403  
 Inj Date : 04-APR-2013 16:04  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : IC-1531403  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsrv\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 12:31 BSMSD.i Quant Type: ISTD  
 Cal Date : 04-APR-2013 15:42 Cal File: 1DD04012.D  
 Als bottle: 11 Calibration Sample, Level: 7  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT ( ug/l)	ON-COL ( ug/l)
* 1 Naphthalene-d8	136	6.090	6.090	(1.000)	2444753	40.0000	
* 6 Acenaphthene-d10	164	7.770	7.770	(1.000)	1439391	40.0000	
* 9 Phenanthrene-d10	188	9.027	9.027	(1.000)	2373597	40.0000	
\$ 13 o-Terphenyl	230	9.339	9.339	(1.034)	2031596	50.0000	57(A)
* 17 Chrysene-d12	240	11.348	11.348	(1.000)	2479223	40.0000	
* 22 Perylene-d12	264	13.175	13.175	(1.000)	2461140	40.0000	
2 Naphthalene	128	6.113	6.113	(1.004)	3211548	50.0000	53(A)
3 2-Methylnaphthalene	142	6.818	6.818	(1.120)	2134320	50.0000	54(A)
4 1-Methylnaphthalene	142	6.912	6.912	(1.135)	1999874	50.0000	54(A)
5 Acenaphthylene	152	7.641	7.641	(0.983)	3396591	50.0000	56(A)
7 Acenaphthene	154	7.799	7.799	(1.004)	2018481	50.0000	54(A)
8 Fluorene	166	8.240	8.240	(1.060)	2393163	50.0000	54(A)
10 Phenanthrene	178	9.051	9.051	(1.003)	3534794	50.0000	54(A)
11 Anthracene	178	9.092	9.092	(1.007)	3590722	50.0000	55(A)
12 Carbazole	167	9.233	9.233	(1.023)	3137679	50.0000	55(A)
14 Fluoranthene	202	10.032	10.032	(1.111)	3681257	50.0000	55(A)
15 Pyrene	202	10.220	10.220	(0.901)	3965627	50.0000	53(A)
16 Benzo(a)anthracene	228	11.325	11.325	(0.998)	3388838	50.0000	50(A)
18 Chrysene	228	11.377	11.377	(1.003)	3512644	50.0000	52(A)
19 Benzo(b)fluoranthene	252	12.635	12.635	(0.959)	3290902	50.0000	54(A)
20 Benzo(k)fluoranthene	252	12.682	12.682	(0.963)	3421834	50.0000	53(A)
21 Benzo(a)pyrene	252	13.093	13.093	(0.994)	3327888	50.0000	54(A)
23 Indeno(1,2,3-cd)pyrene	276	14.785	14.785	(1.122)	3754268	50.0000	57(AM)
24 Dibenzo(a,h)anthracene	278	14.826	14.826	(1.125)	3350541	50.0000	54(A)
25 Benzo(g,h,i)perylene	276	15.238	15.238	(1.157)	3284166	50.0000	52(A)

QC Flag Legend

- A - Target compound detected but, quantitated amount exceeded maximum amount.
- M - Compound response manually integrated.



Data File: 1DD04013.D

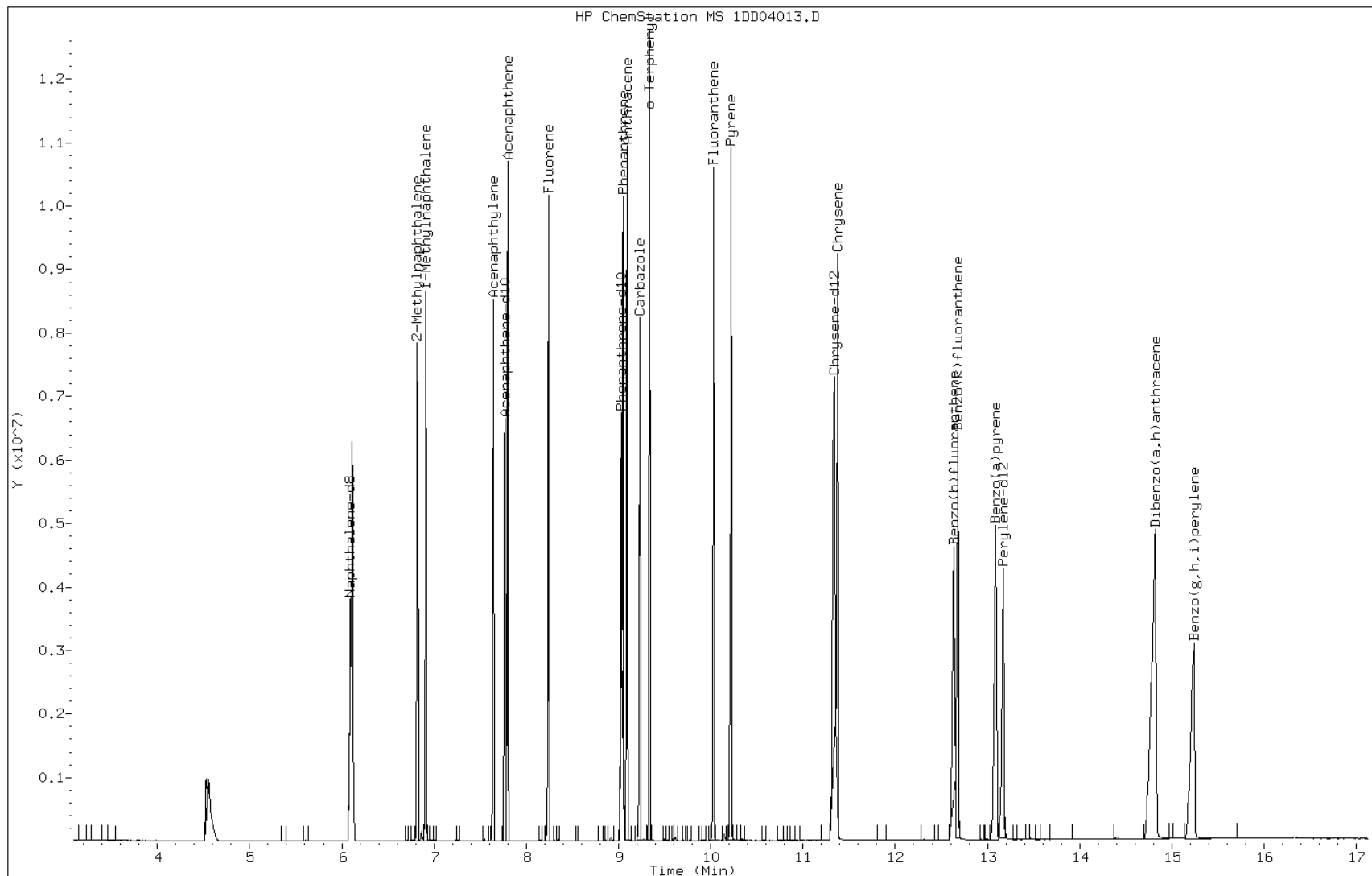
Date: 04-APR-2013 16:04

Client ID:

Instrument: BSMSD.i

Sample Info: IC-1531403

Operator: SCC

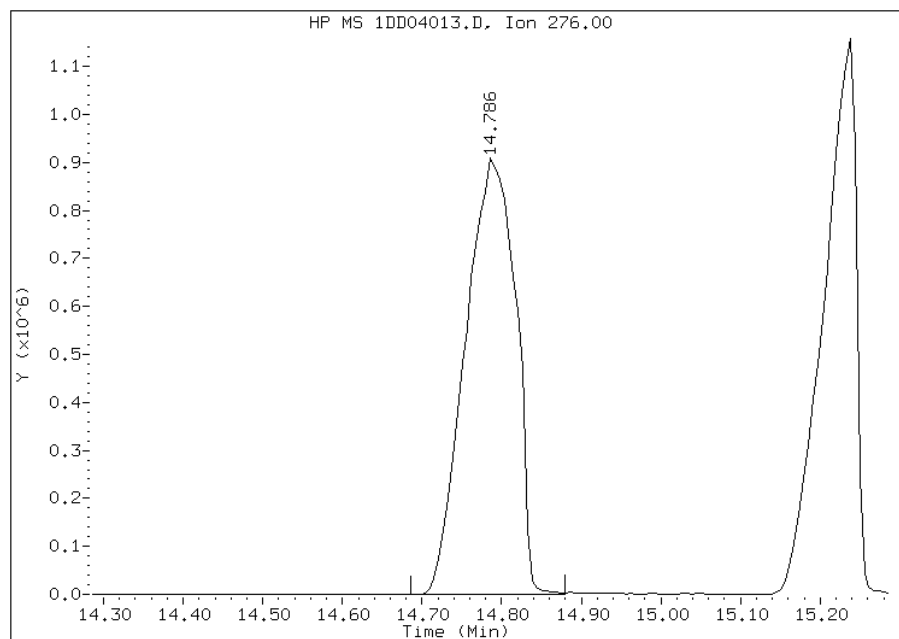


# Manual Integration Report

Data File: 1DD04013.D  
Inj. Date and Time: 04-APR-2013 16:04  
Instrument ID: BSMDS.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

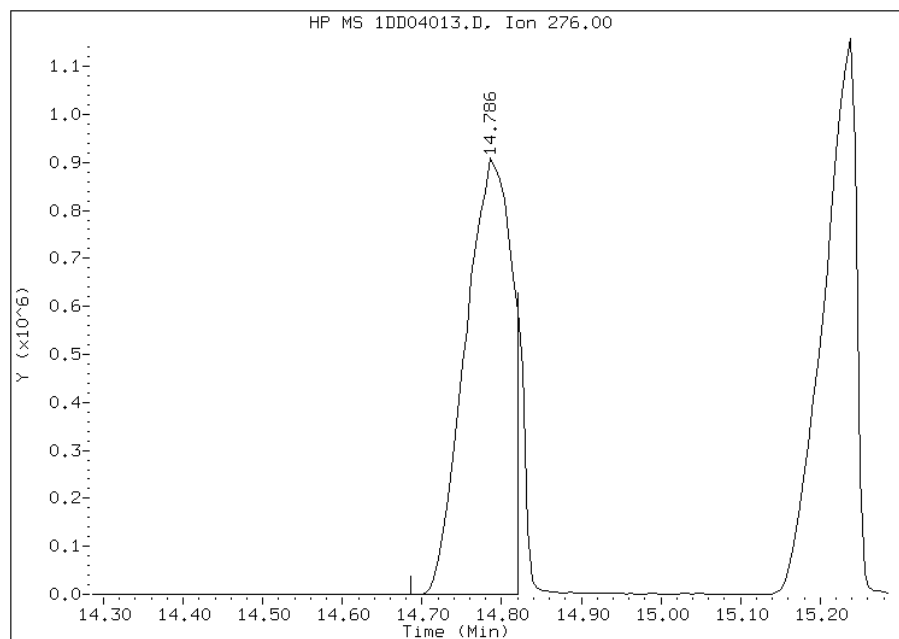
## Processing Integration Results

RT: 14.79  
Response: 3993028  
Amount: 54  
Conc: 54



## Manual Integration Results

RT: 14.79  
Response: 3754268  
Amount: 57  
Conc: 57



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:30  
Manual Integration Reason: Split Peak

FORM VII  
GC/MS SEMI VOA CONTINUING CALIBRATION DATA

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab Sample ID: ICV 660-136048/12 Calibration Date: 04/02/2013 15:34  
 Instrument ID: BSMC5973 Calib Start Date: 04/02/2013 13:26  
 GC Column: DB-5MS ID: 250.00 (um) Calib End Date: 04/02/2013 15:15  
 Lab File ID: 1CD02012.D Conc. Units: ug/Kg

ANALYTE	CURVE TYPE	AVE RRF	RRF	MIN RRF	CALC AMOUNT	SPIKE AMOUNT	%D	MAX %D
Naphthalene	Ave	1.027	0.9549	0.0000	18600	20000	-7.1	35.0
2-Methylnaphthalene	Ave	0.6994	0.5884	0.0000	16800	20000	-15.9	35.0
1-Methylnaphthalene	Ave	0.6293	0.5998	0.0000	19100	20000	-4.7	35.0
Acenaphthylene	Ave	1.656	1.493	0.0000	18000	20000	-9.8	35.0
Acenaphthene	Lin	1.025	0.8508	0.0000	16600	20000	-17.0	35.0
Fluorene	Ave	1.367	1.209	0.0000	17700	20000	-11.5	35.0
Phenanthrene	Ave	1.165	0.9563	0.0000	16400	20000	-17.9	35.0
Anthracene	Ave	1.181	0.9425	0.0000	16000	20000	-20.2	35.0
Carbazole	Ave	1.012	0.8775	0.0000	17300	20000	-13.3	35.0
Fluoranthene	Ave	1.287	1.100	0.0000	17100	20000	-14.5	35.0
Pyrene	Ave	1.108	0.8708	0.0000	15700	20000	-21.4	35.0
Benzo[a]anthracene	Lin	1.278	0.9658	0.0000	16800	20000	-16.0	35.0
Chrysene	Ave	1.140	0.8716	0.0000	15300	20000	-23.5	35.0
Benzo[b]fluoranthene	Ave	1.131	0.8920	0.0000	15800	20000	-21.1	35.0
Benzo[k]fluoranthene	Ave	1.094	0.8978	0.0000	16400	20000	-17.9	35.0
Benzo[a]pyrene	Ave	1.065	0.8060	0.0000	15100	20000	-24.3	35.0
Indeno[1,2,3-cd]pyrene	Ave	1.011	0.8744	0.0000	17300	20000	-13.5	35.0
Dibenz(a,h)anthracene	Ave	0.9341	0.8626	0.0000	18500	20000	-7.7	35.0
Benzo[g,h,i]perylene	Ave	1.032	0.8592	0.0000	16600	20000	-16.8	35.0
o-Terphenyl	Lin	0.6233	0.5049	0.0000	16200	20000	-19.0	35.0

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02012.D  
 Lab Smp Id: ICV-1448440  
 Inj Date : 02-APR-2013 15:34  
 Operator : SCC  
 Smp Info : ICV-1448440  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\a-bFASTPAHi-m.m  
 Meth Date : 02-Apr-2013 15:55 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 12 QC Sample: LCS  
 Dil Factor: 1.00000  
 Integrator: HP RTE  
 Target Version: 4.14  
 Inst ID: BSMC5973.i  
 Compound Sublist: pah.sub

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	1000.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
							ON-COLUMN (ug/ml)	FINAL ( ug/l)
* 1 Naphthalene-d8	136	3.710	3.710	(1.000)	649122	40.0000		
* 6 Acenaphthene-d10	164	4.798	4.798	(1.000)	500935	40.0000		
* 10 Phenanthrene-d10	188	5.745	5.745	(1.000)	955391	40.0000		
\$ 14 o-Terphenyl	230	5.998	5.998	(1.044)	241169	16.1906	16.1906	
* 18 Chrysene-d12	240	7.686	7.686	(1.000)	1249690	40.0000		
* 23 Perylene-d12	264	8.856	8.863	(1.000)	1306409	40.0000		
2 Naphthalene	128	3.727	3.728	(1.005)	309919	18.5886	18.5885	
3 2-Methylnaphthalene	142	4.151	4.151	(1.119)	190970	16.8266	16.8266	
4 1-Methylnaphthalene	142	4.216	4.216	(1.136)	194664	19.0620	19.0620	
5 Acenaphthylene	152	4.710	4.710	(0.982)	373939	18.0364	18.0363	
7 Acenaphthene	154	4.821	4.822	(1.005)	213089	16.5944	16.5943	
9 Fluorene	166	5.139	5.139	(1.071)	302875	17.6930	17.6929	
11 Phenanthrene	178	5.763	5.763	(1.003)	456841	16.4181	16.4181	
12 Anthracene	178	5.798	5.798	(1.009)	450208	15.9610	15.9609	
13 Carbazole	167	5.904	5.904	(1.028)	419186	17.3461	17.3460	
15 Fluoranthene	202	6.598	6.598	(1.148)	525545	17.1022	17.1021	
16 Pyrene	202	6.763	6.763	(0.880)	544110	15.7178	15.7178	
17 Benzo(a)anthracene	228	7.680	7.680	(0.999)	603470	16.8016	16.8016	

Compounds	QUANT SIG		CONCENTRATIONS				
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/ml)	FINAL ( ug/l)
-----	----	----	-----	-----	-----	-----	-----
19 Chrysene	228	7.704	7.710	(1.002)	544600	15.2932	15.2931
20 Benzo(b)fluoranthene	252	8.515	8.522	(0.961)	582649	15.7757	15.7757
21 Benzo(k)fluoranthene	252	8.539	8.545	(0.964)	586474	16.4181	16.4181
22 Benzo(a)pyrene	252	8.804	8.810	(0.994)	526495	15.1414	15.1414
24 Indeno(1,2,3-cd)pyrene	276	10.009	10.016	(1.130)	571166	17.2941	17.2940(M)
25 Dibenzo(a,h)anthracene	278	10.021	10.033	(1.131)	563427	18.4677	18.4676
26 Benzo(g,h,i)perylene	276	10.351	10.363	(1.169)	561199	16.6490	16.6490

QC Flag Legend

M - Compound response manually integrated.

Data File: 1CD02012.D

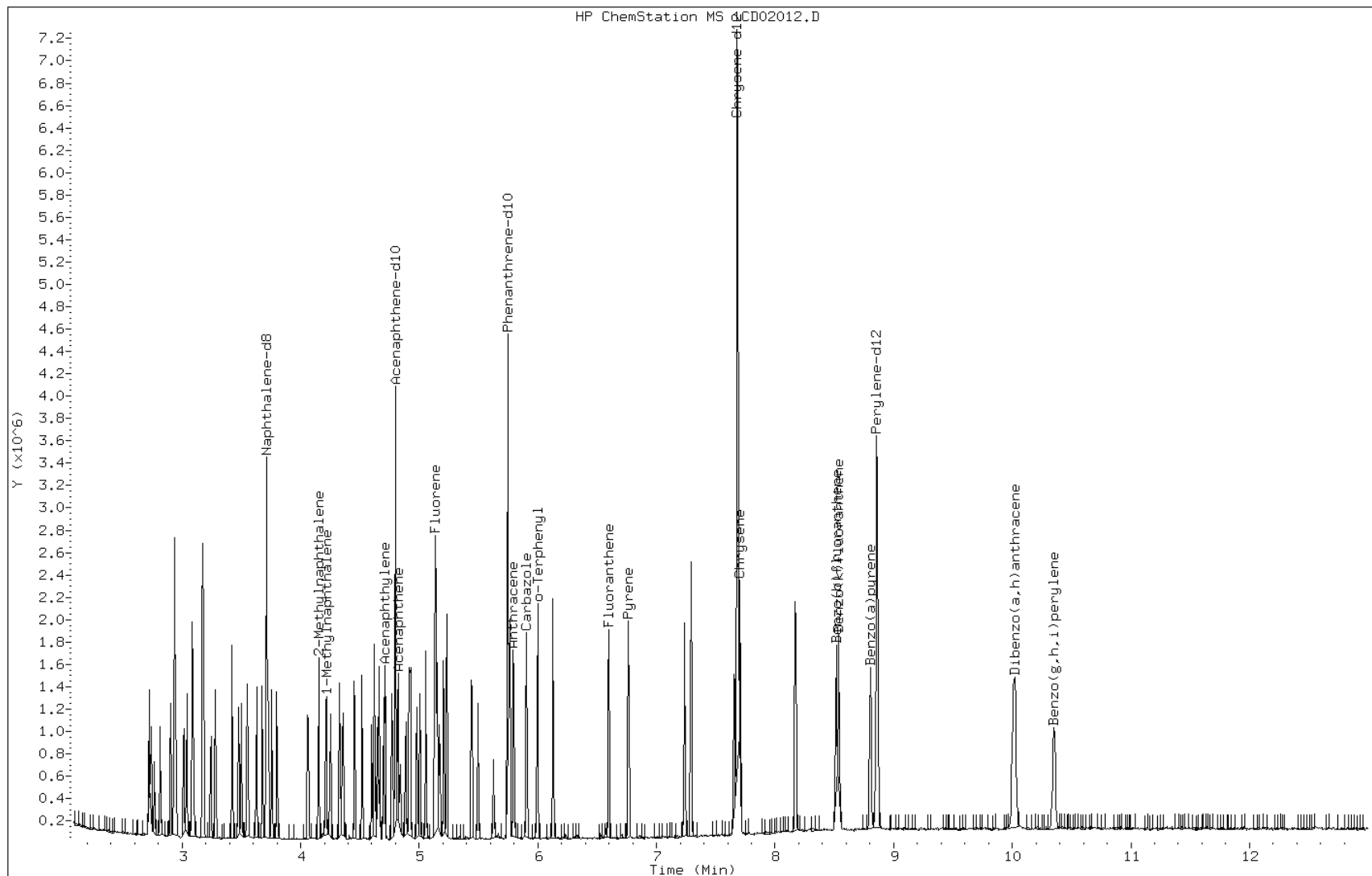
Date: 02-APR-2013 15:34

Client ID:

Instrument: BSMC5973.i

Sample Info: ICV-1448440

Operator: SCC

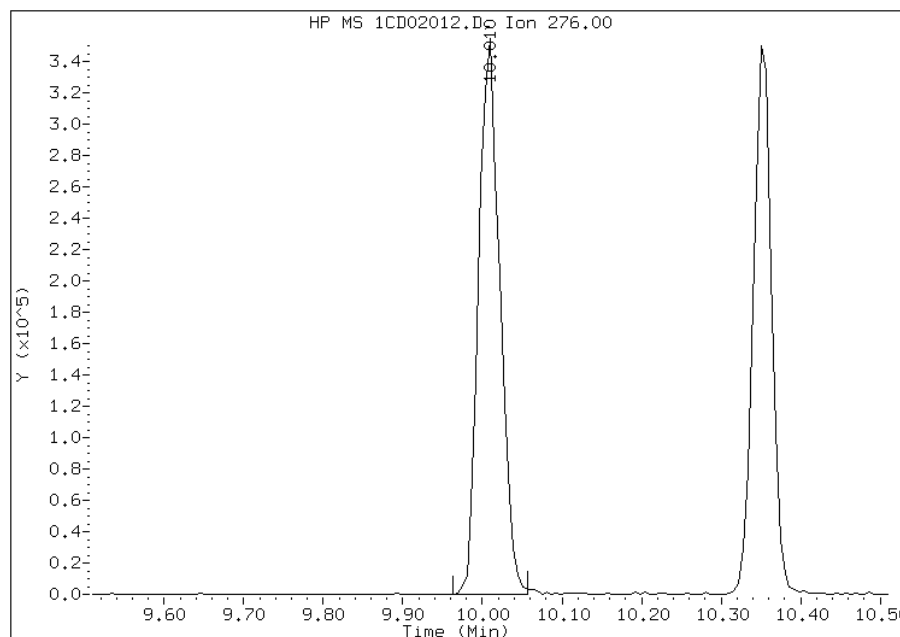


# Manual Integration Report

Data File: 1CD02012.D  
Inj. Date and Time: 02-APR-2013 15:34  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/02/2013

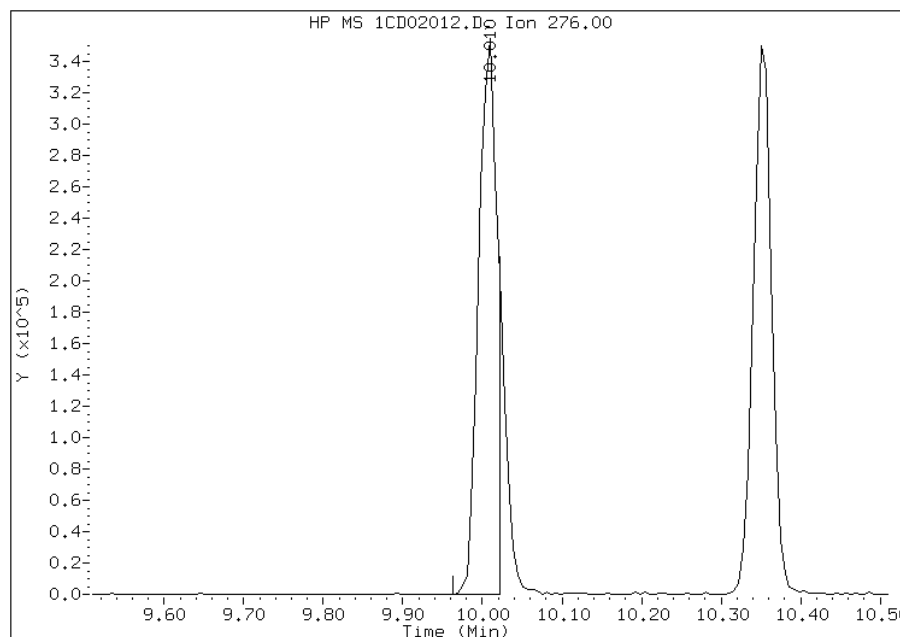
## Processing Integration Results

RT: 10.01  
Response: 653584  
Amount: 20  
Conc: 20



## Manual Integration Results

RT: 10.01  
Response: 571166  
Amount: 17  
Conc: 17



Manually Integrated By: cantins  
Modification Date: 02-Apr-2013 15:57  
Manual Integration Reason: Split Peak

FORM VII  
GC/MS SEMI VOA CONTINUING CALIBRATION DATA

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab Sample ID: CCVIS 660-136081/3 Calibration Date: 04/03/2013 11:45  
 Instrument ID: BSMC5973 Calib Start Date: 04/02/2013 13:26  
 GC Column: DB-5MS ID: 250.00 (um) Calib End Date: 04/02/2013 15:15  
 Lab File ID: 1CD03003.D Conc. Units: ug/Kg

ANALYTE	CURVE TYPE	AVE RRF	RRF	MIN RRF	CALC AMOUNT	SPIKE AMOUNT	%D	MAX %D
Naphthalene	Ave	1.027	0.999	0.0000	19400	20000	-2.8	20.0
2-Methylnaphthalene	Ave	0.6994	0.6776	0.0000	19400	20000	-3.1	20.0
1-Methylnaphthalene	Ave	0.6293	0.6430	0.0000	20400	20000	2.2	20.0
Acenaphthylene	Ave	1.656	1.706	0.0000	20600	20000	3.1	20.0
Acenaphthene	Lin	1.025	0.9817	0.0000	19100	20000	-4.3	20.0
Fluorene	Ave	1.367	1.328	0.0000	19400	20000	-2.9	20.0
Phenanthrene	Ave	1.165	1.152	0.0000	19800	20000	-1.1	20.0
Anthracene	Ave	1.181	1.140	0.0000	19300	20000	-3.5	20.0
Carbazole	Ave	1.012	1.032	0.0000	20400	20000	2.0	20.0
Fluoranthene	Ave	1.287	1.311	0.0000	20400	20000	1.9	20.0
Pyrene	Ave	1.108	1.146	0.0000	20700	20000	3.5	20.0
Benzo[a]anthracene	Lin	1.278	1.115	0.0000	19400	20000	-3.1	20.0
Chrysene	Ave	1.140	1.082	0.0000	19000	20000	-5.1	20.0
Benzo[b]fluoranthene	Ave	1.131	1.174	0.0000	20800	20000	3.8	20.0
Benzo[k]fluoranthene	Ave	1.094	1.068	0.0000	19500	20000	-2.4	20.0
Benzo[a]pyrene	Ave	1.065	1.055	0.0000	19800	20000	-0.9	20.0
Indeno[1,2,3-cd]pyrene	Ave	1.011	0.9599	0.0000	19000	20000	-5.1	20.0
Dibenz(a,h)anthracene	Ave	0.9341	0.9411	0.0000	20200	20000	0.8	20.0
Benzo[g,h,i]perylene	Ave	1.032	0.995	0.0000	19300	20000	-3.6	20.0
o-Terphenyl	Lin	0.6233	0.6314	0.0000	20100	20000	0.3	20.0



TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsrv\chem\SM\BSMC5973.i\1C040313.b\1CD03003.D  
 Lab Smp Id: CCVIS-1531401  
 Inj Date : 03-APR-2013 11:45  
 Operator : SCC  
 Smp Info : CCVIS-1531401  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsrv\chem\SM\BSMC5973.i\1C040313.b\a-bFASTPAHi-m.m  
 Meth Date : 03-Apr-2013 11:59 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 3 Continuing Calibration Sample  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG						AMOUNTS	
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)
* 1 Naphthalene-d8	136	3.704	3.704	(1.000)	500765	40.0000		
* 6 Acenaphthene-d10	164	4.792	4.792	(1.000)	364027	40.0000		
* 10 Phenanthrene-d10	188	5.739	5.739	(1.000)	687020	40.0000		
\$ 14 o-Terphenyl	230	5.992	5.992	(1.044)	216882	20.0000	20.0659	
* 18 Chrysene-d12	240	7.680	7.680	(1.000)	857573	40.0000		
* 23 Perylene-d12	264	8.851	8.851	(1.000)	866012	40.0000		
2 Naphthalene	128	3.722	3.722	(1.005)	250131	20.0000	19.4472	
3 2-Methylnaphthalene	142	4.145	4.145	(1.119)	169659	20.0000	19.3776	
4 1-Methylnaphthalene	142	4.210	4.210	(1.137)	161005	20.0000	20.4368	
5 Acenaphthylene	152	4.704	4.704	(0.982)	310521	20.0000	20.6104	
7 Acenaphthene	154	4.816	4.816	(1.005)	178686	20.0000	19.1486	
9 Fluorene	166	5.133	5.133	(1.071)	241638	20.0000	19.4245	
11 Phenanthrene	178	5.757	5.757	(1.003)	395730	20.0000	19.7773	
12 Anthracene	178	5.792	5.792	(1.009)	391504	20.0000	19.3016	
13 Carbazole	167	5.898	5.898	(1.028)	354598	20.0000	20.4052	
15 Fluoranthene	202	6.592	6.592	(1.149)	450430	20.0000	20.3836	
16 Pyrene	202	6.757	6.757	(0.880)	491523	20.0000	20.6909	
17 Benzo(a)anthracene	228	7.668	7.668	(0.998)	478063	20.0000	19.3750	
19 Chrysene	228	7.698	7.698	(1.002)	463959	20.0000	18.9858	
20 Benzo(b)fluoranthene	252	8.509	8.509	(0.961)	508197	20.0000	20.7572	
21 Benzo(k)fluoranthene	252	8.533	8.533	(0.964)	462286	20.0000	19.5227	
22 Benzo(a)pyrene	252	8.798	8.798	(0.994)	456933	20.0000	19.8234	
24 Indeno(1,2,3-cd)pyrene	276	9.992	9.992	(1.129)	415659	20.0000	18.9857(M)	
25 Dibenzo(a,h)anthracene	278	10.009	10.009	(1.131)	407517	20.0000	20.1500	
26 Benzo(g,h,i)perylene	276	10.339	10.339	(1.168)	431010	20.0000	19.2892	

QC Flag Legend

M - Compound response manually integrated.

Data File: 1CD03003.D

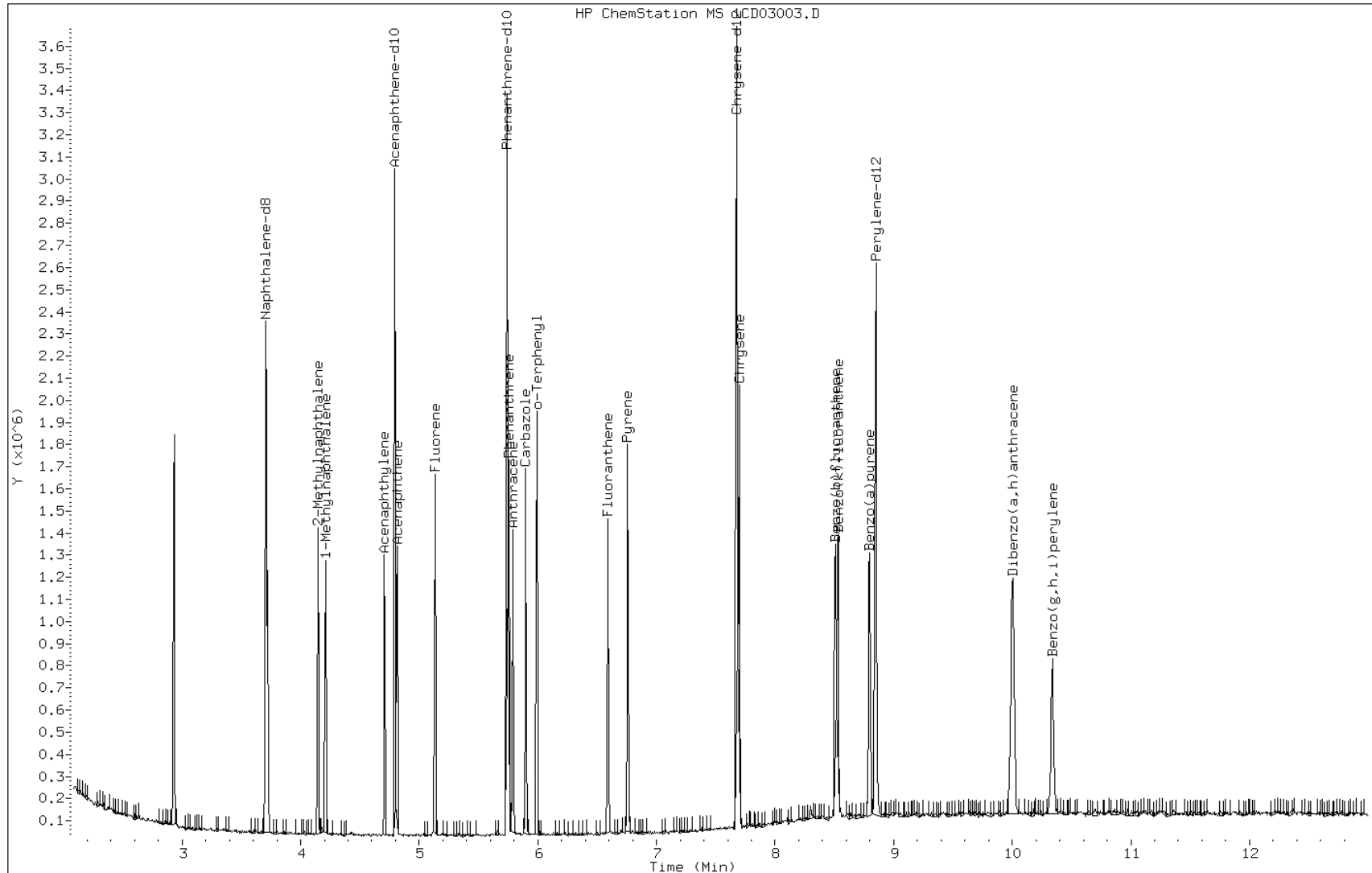
Date: 03-APR-2013 11:45

Client ID:

Instrument: BSMC5973.i

Sample Info: CCVIS-1531401

Operator: SCC

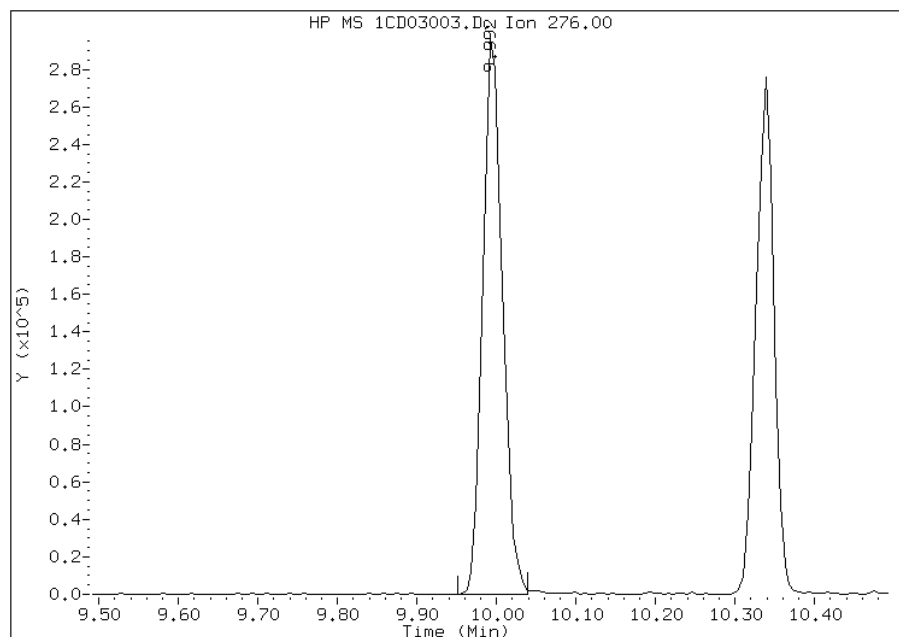


# Manual Integration Report

Data File: 1CD03003.D  
Inj. Date and Time: 03-APR-2013 11:45  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/03/2013

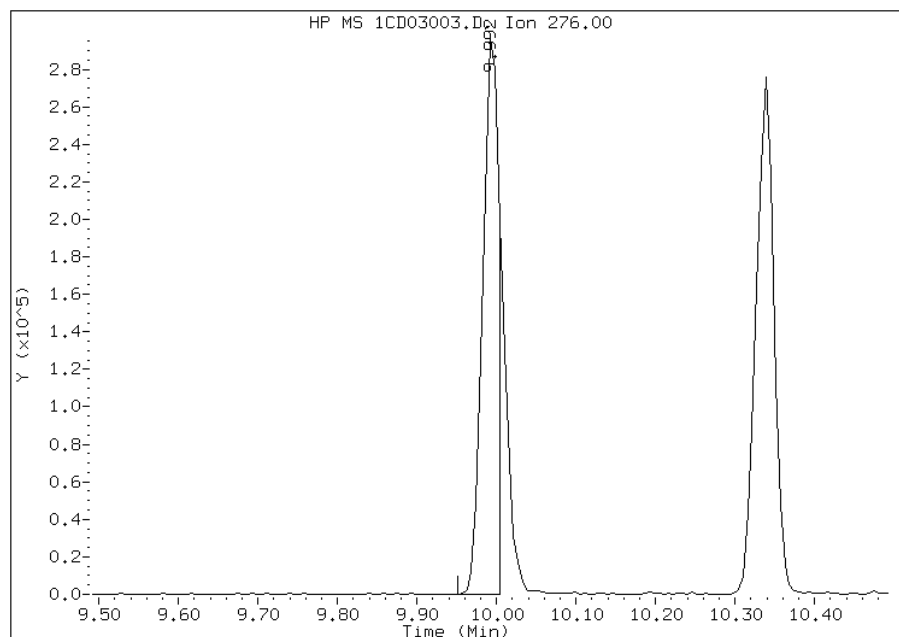
## Processing Integration Results

RT: 9.99  
Response: 506304  
Amount: 23  
Conc: 23



## Manual Integration Results

RT: 9.99  
Response: 415659  
Amount: 19  
Conc: 19



Manually Integrated By: cantins  
Modification Date: 03-Apr-2013 11:59  
Manual Integration Reason: Split Peak

FORM VII  
GC/MS SEMI VOA CONTINUING CALIBRATION DATA

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab Sample ID: CCVIS 660-136171/4 Calibration Date: 04/05/2013 12:15  
 Instrument ID: BSMC5973 Calib Start Date: 04/02/2013 13:26  
 GC Column: DB-5MS ID: 250.00 (um) Calib End Date: 04/02/2013 15:15  
 Lab File ID: 1CD05004.D Conc. Units: ug/mL

ANALYTE	CURVE TYPE	AVE RRF	RRF	MIN RRF	CALC AMOUNT	SPIKE AMOUNT	%D	MAX %D
Naphthalene	Ave	1.027	1.032	0.0000	2000	20.0	0.5	20.0
2-Methylnaphthalene	Ave	0.6994	0.7098	0.0000	2000	20.0	1.5	20.0
1-Methylnaphthalene	Ave	0.6293	0.6324	0.0000	2000	20.0	0.5	20.0
Acenaphthylene	Ave	1.656	1.686	0.0000	1000	20.0	1.8	20.0
Acenaphthene	Lin	1.025	0.9558	0.0000	2000	20.0	-6.8	20.0
Fluorene	Ave	1.367	1.254	0.0000	2000	20.0	-8.2	20.0
Phenanthrene	Ave	1.165	1.127	0.0000	500	20.0	-3.2	20.0
Anthracene	Ave	1.181	1.206	0.0000	200	20.0	2.1	20.0
Carbazole	Ave	1.012	1.040	0.0000	1000	20.0	2.8	20.0
Fluoranthene	Ave	1.287	1.346	0.0000	500	20.0	4.6	20.0
Pyrene	Ave	1.108	1.077	0.0000	500	20.0	-2.8	20.0
Benzo[a]anthracene	Lin	1.278	1.101	0.0000	200	20.0	-4.3	20.0
Chrysene	Ave	1.140	1.074	0.0000	200	20.0	-5.8	20.0
Benzo[b]fluoranthene	Ave	1.131	1.071	0.0000	200	20.0	-5.3	20.0
Benzo[k]fluoranthene	Ave	1.094	1.162	0.0000	200	20.0	6.3	20.0
Benzo[a]pyrene	Ave	1.065	1.057	0.0000	200	20.0	-0.7	20.0
Indeno[1,2,3-cd]pyrene	Ave	1.011	0.9896	0.0000	200	20.0	-2.1	20.0
Dibenz(a,h)anthracene	Ave	0.9341	0.9614	0.0000	200	20.0	2.9	20.0
Benzo[g,h,i]perylene	Ave	1.032	0.9820	0.0000	500	20.0	-4.9	20.0
o-Terphenyl	Lin	0.6233	0.6473	0.0000	20.6	20.0	2.8	20.0

TestAmerica Laboratories

Semivolatile 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\1CD05004.D  
 Lab Smp Id: CCVIS-1531401  
 Inj Date : 05-APR-2013 12:15  
 Operator : SCC  
 Smp Info : CCVIS-1531401  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\a-bFASTPAHi-m.m  
 Meth Date : 05-Apr-2013 12:31 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 3 Continuing Calibration Sample  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/ml)	ON-COL (ug/ml)
* 1 Naphthalene-d8	136	3.692	3.692	(1.000)	392528	40.0000	
* 6 Acenaphthene-d10	164	4.780	4.780	(1.000)	289150	40.0000	
* 10 Phenanthrene-d10	188	5.721	5.721	(1.000)	539578	40.0000	(H)
\$ 14 o-Terphenyl	230	5.974	5.974	(1.044)	174628	20.0000	20.5532
* 18 Chrysene-d12	240	7.662	7.662	(1.000)	739705	40.0000	
* 23 Perylene-d12	264	8.827	8.827	(1.000)	746693	40.0000	(H)
2 Naphthalene	128	3.704	3.704	(1.003)	202593	20.0000	20.0945
3 2-Methylnaphthalene	142	4.133	4.133	(1.119)	139304	20.0000	20.2978
4 1-Methylnaphthalene	142	4.192	4.192	(1.135)	124123	20.0000	20.0997
5 Acenaphthylene	152	4.692	4.692	(0.982)	243681	20.0000	20.3623
7 Acenaphthene	154	4.798	4.798	(1.004)	138184	20.0000	18.6430
9 Fluorene	166	5.116	5.116	(1.070)	181351	20.0000	18.3533
11 Phenanthrene	178	5.739	5.739	(1.003)	304115	20.0000	19.3518(H)
12 Anthracene	178	5.774	5.774	(1.009)	325239	20.0000	20.4162(H)
13 Carbazole	167	5.880	5.880	(1.028)	280645	20.0000	20.5626(H)
15 Fluoranthene	202	6.574	6.574	(1.149)	363056	20.0000	20.9190(H)
16 Pyrene	202	6.739	6.739	(0.879)	398242	20.0000	19.4355
17 Benzo(a)anthracene	228	7.651	7.651	(0.998)	407283	20.0000	19.1383
19 Chrysene	228	7.680	7.680	(1.002)	397270	20.0000	18.8472
20 Benzo(b)fluoranthene	252	8.486	8.486	(0.961)	399687	20.0000	18.9338(H)
21 Benzo(k)fluoranthene	252	8.509	8.509	(0.964)	433951	20.0000	21.2545(H)
22 Benzo(a)pyrene	252	8.774	8.774	(0.994)	394530	20.0000	19.8513(H)
24 Indeno(1,2,3-cd)pyrene	276	9.962	9.962	(1.129)	369463	20.0000	19.5723(MH)
25 Dibenzo(a,h)anthracene	278	9.980	9.980	(1.131)	358939	20.0000	20.5841(H)
26 Benzo(g,h,i)perylene	276	10.303	10.303	(1.167)	366622	20.0000	19.0294(H)

QC Flag Legend

M - Compound response manually integrated.  
 H - Operator selected an alternate compound hit.

Data File: 1CD05004.D

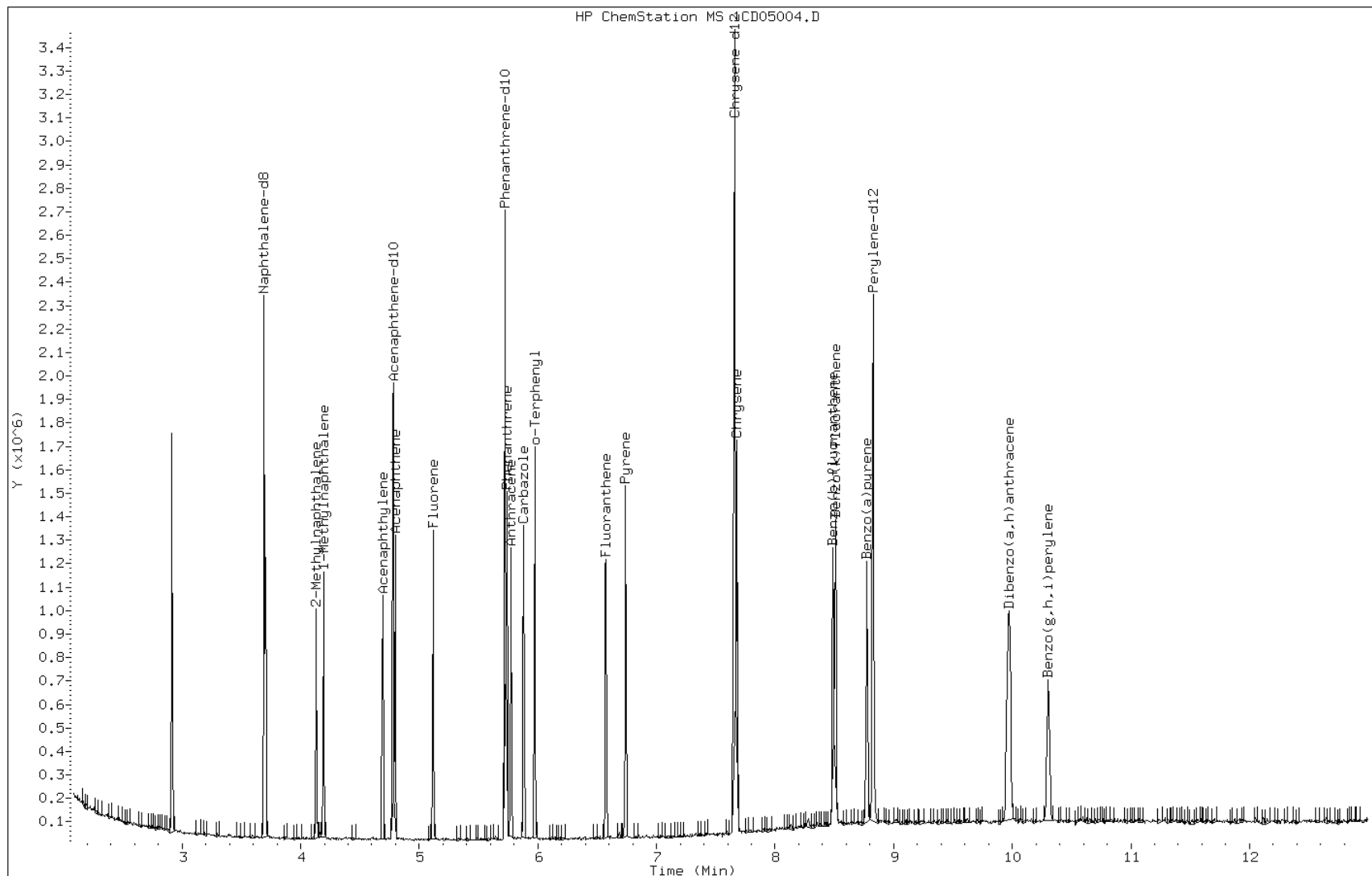
Date: 05-APR-2013 12:15

Client ID:

Instrument: BSMC5973.i

Sample Info: CCVIS-1531401

Operator: SCC

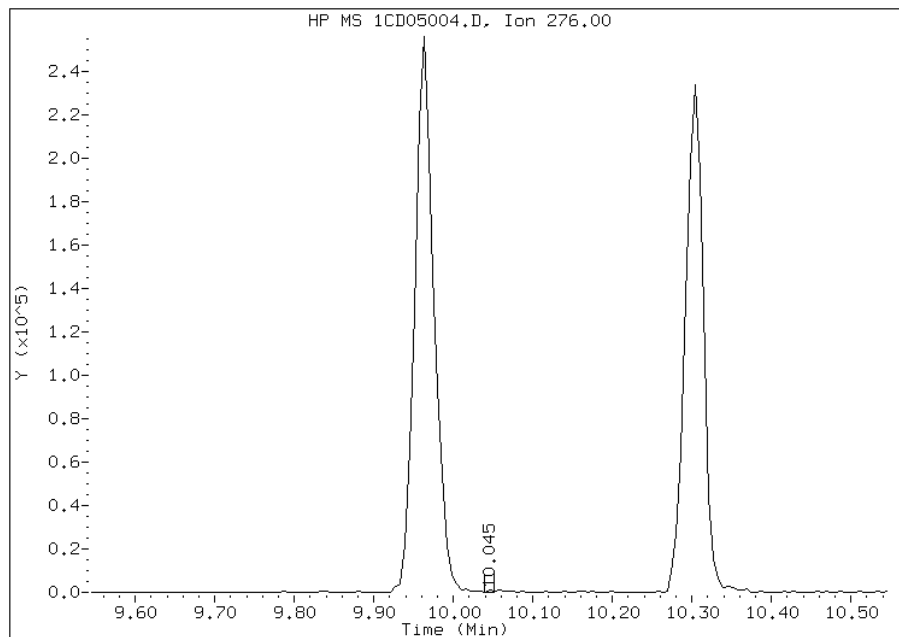


Manual Integration Report

Data File: 1CD05004.D  
Inj. Date and Time: 05-APR-2013 12:15  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

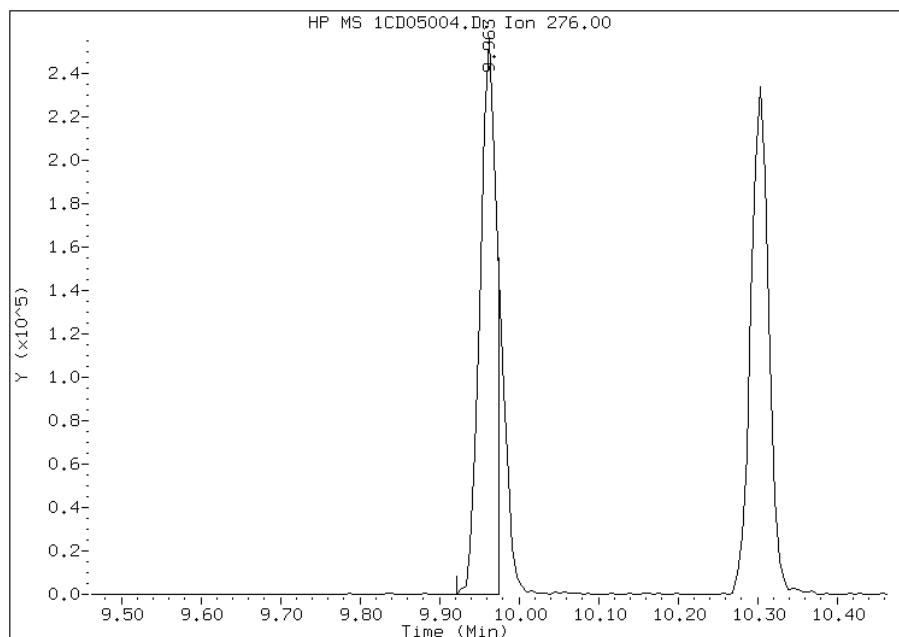
Processing Integration Results

RT: 10.05  
Response: 614  
Amount: 0  
Conc: 0



Manual Integration Results

RT: 9.96  
Response: 369463  
Amount: 20  
Conc: 20



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 12:33  
Manual Integration Reason: Split Peak

FORM VII  
GC/MS SEMI VOA CONTINUING CALIBRATION DATA

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Lab Sample ID: ICV 660-136164/22 Calibration Date: 04/04/2013 16:27  
 Instrument ID: BSMD5973 Calib Start Date: 04/04/2013 13:49  
 GC Column: DB-5MS ID: 250.00 (um) Calib End Date: 04/04/2013 16:04  
 Lab File ID: 1DD04014.D Conc. Units: ug/mL

ANALYTE	CURVE TYPE	AVE RRF	RRF	MIN RRF	CALC AMOUNT	SPIKE AMOUNT	%D	MAX %D
Naphthalene	Ave	0.9942	0.9009	0.0000	2000	20.0	-9.4	35.0
2-Methylnaphthalene	Ave	0.6418	0.5957	0.0000	2000	20.0	-7.2	35.0
1-Methylnaphthalene	Ave	0.6061	0.5697	0.0000	2000	20.0	-6.0	35.0
Acenaphthylene	Ave	1.693	1.431	0.0000	1000	20.0	-15.5	35.0
Acenaphthene	Ave	1.045	0.8522	0.0000	2000	20.0	-18.5	35.0
Fluorene	Ave	1.238	1.099	0.0000	2000	20.0	-11.2	35.0
Phenanthrene	Ave	1.102	0.8997	0.0000	500	20.0	-18.3	35.0
Anthracene	Ave	1.094	0.9197	0.0000	200	20.0	-15.9	35.0
Carbazole	Ave	0.9646	0.6860	0.0000	1000	20.0	-28.9	35.0
Fluoranthene	Ave	1.134	0.9937	0.0000	500	20.0	-12.4	35.0
Pyrene	Ave	1.201	0.9577	0.0000	500	20.0	-20.3	35.0
Benzo[a]anthracene	Ave	1.156	0.9847	0.0000	200	20.0	-14.9	35.0
Chrysene	Ave	1.084	0.8727	0.0000	200	20.0	-19.5	35.0
Benzo[b]fluoranthene	Ave	0.999	0.8893	0.0000	200	20.0	-11.0	35.0
Benzo[k]fluoranthene	Ave	1.053	0.8752	0.0000	200	20.0	-16.9	35.0
Benzo[a]pyrene	Ave	1.004	0.7657	0.0000	200	20.0	-23.7	35.0
Indeno[1,2,3-cd]pyrene	Ave	1.071	0.8560	0.0000	200	20.0	-20.0	35.0
Dibenz(a,h)anthracene	Ave	1.008	0.9464	0.0000	200	20.0	-6.1	35.0
Benzo[g,h,i]perylene	Ave	1.031	0.8761	0.0000	500	20.0	-15.0	35.0
o-Terphenyl	Ave	0.6027	0.4989	0.0000	16.6	20.0	-17.2	35.0



TestAmerica Laboratories

Semivolatile 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04014.D  
 Lab Smp Id: ICV-1448440  
 Inj Date : 04-APR-2013 16:27  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : ICV-1448440  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dFASTPAHi.m  
 Meth Date : 05-Apr-2013 13:07 cantins Quant Type: ISTD  
 Cal Date : 04-APR-2013 16:04 Cal File: 1DD04013.D  
 Als bottle: 12 QC Sample: LCS  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	1000.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
								ON-COLUMN ( ug/l)	FINAL ( ug/l)
* 1 Naphthalene-d8			136	6.096	6.090	(1.000)	3619899	40.0000	
* 6 Acenaphthene-d10			164	7.771	7.770	(1.000)	2333423	40.0000	
* 9 Phenanthrene-d10			188	9.028	9.028	(1.000)	3845474	40.0000	
\$ 13 o-Terphenyl			230	9.334	9.339	(1.034)	959307	16.5566	16
* 17 Chrysene-d12			240	11.349	11.349	(1.000)	3963674	40.0000	
* 22 Perylene-d12			264	13.182	13.176	(1.000)	3958481	40.0000	
2 Naphthalene			128	6.114	6.114	(1.003)	1630598	18.1229	18
3 2-Methylnaphthalene			142	6.819	6.819	(1.119)	1078163	18.5630	18
4 1-Methylnaphthalene			142	6.913	6.913	(1.134)	1031118	18.7992	19
5 Acenaphthylene			152	7.642	7.641	(0.983)	1669244	16.9019	17
7 Acenaphthene			154	7.800	7.800	(1.004)	994282	16.3100	16
8 Fluorene			166	8.241	8.240	(1.060)	1281905	17.7572	18
10 Phenanthrene			178	9.046	9.051	(1.002)	1729949	16.3322	16
11 Anthracene			178	9.087	9.092	(1.007)	1768381	16.8207	17
12 Carbazole			167	9.228	9.233	(1.022)	1319041	14.2242	14(M)
14 Fluoranthene			202	10.027	10.032	(1.111)	1910613	17.5287	18
15 Pyrene			202	10.215	10.220	(0.900)	1898084	15.9464	16

Compounds	QUANT SIG		CONCENTRATIONS				
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN ( ug/l)	FINAL ( ug/l)
16 Benzo(a)anthracene	228	11.325	11.325	(0.998)	1951469	17.0289	17
18 Chrysene	228	11.372	11.378	(1.002)	1729613	16.0966	16
19 Benzo(b)fluoranthene	252	12.630	12.635	(0.958)	1760131	17.8000	18
20 Benzo(k)fluoranthene	252	12.671	12.682	(0.961)	1732123	16.6271	17
21 Benzo(a)pyrene	252	13.076	13.094	(0.992)	1515587	15.2542	15
23 Indeno(1,2,3-cd)pyrene	276	14.763	14.786	(1.120)	1694283	15.9925	16(M)
24 Dibenzo(a,h)anthracene	278	14.798	14.827	(1.123)	1873209	18.7764	19
25 Benzo(g,h,i)perylene	276	15.215	15.238	(1.154)	1734029	16.9990	17(H)

QC Flag Legend

- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1DD04014.D

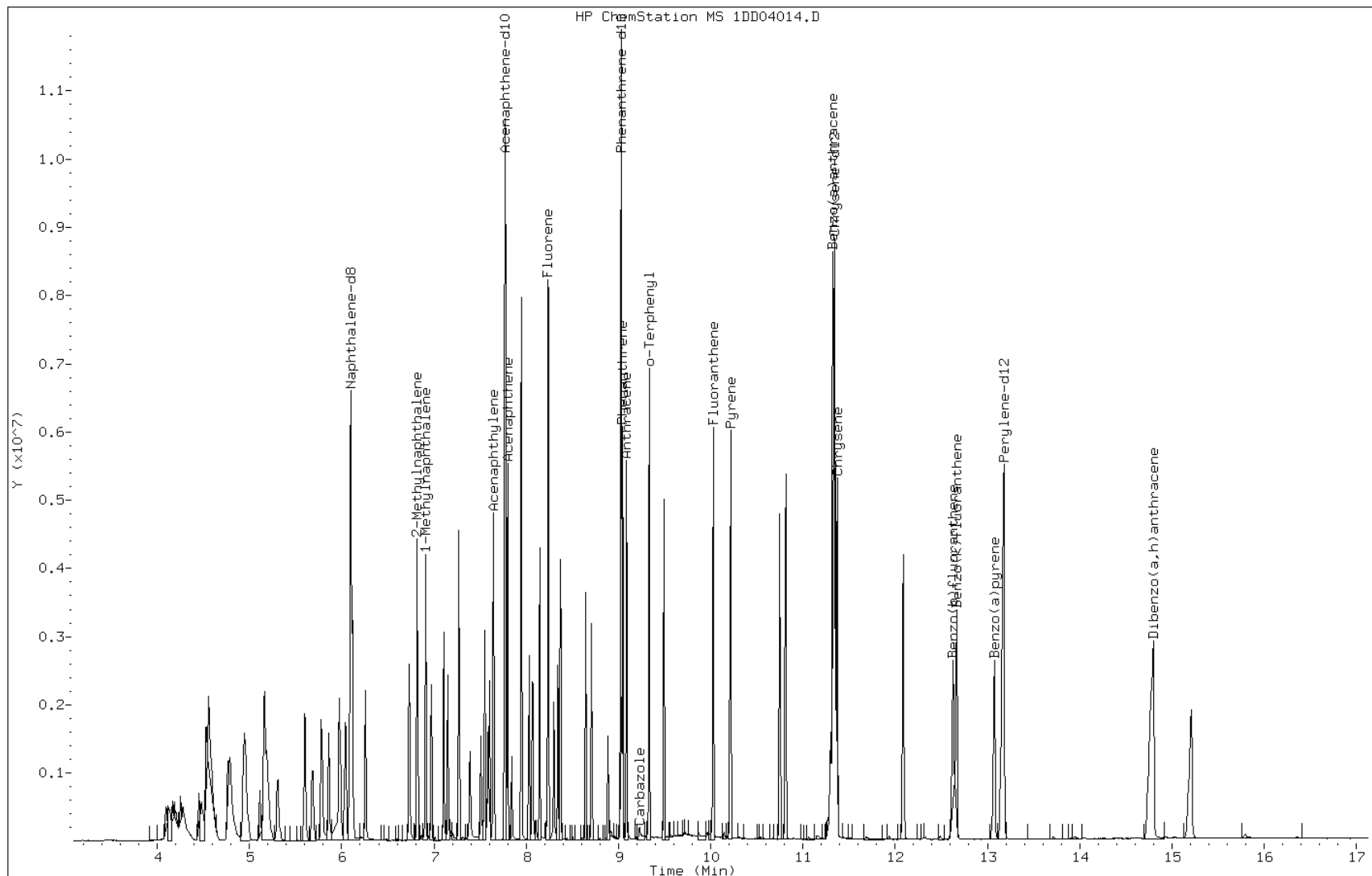
Date: 04-APR-2013 16:27

Client ID:

Instrument: BSMSD.i

Sample Info: ICV-1448440

Operator: SCC

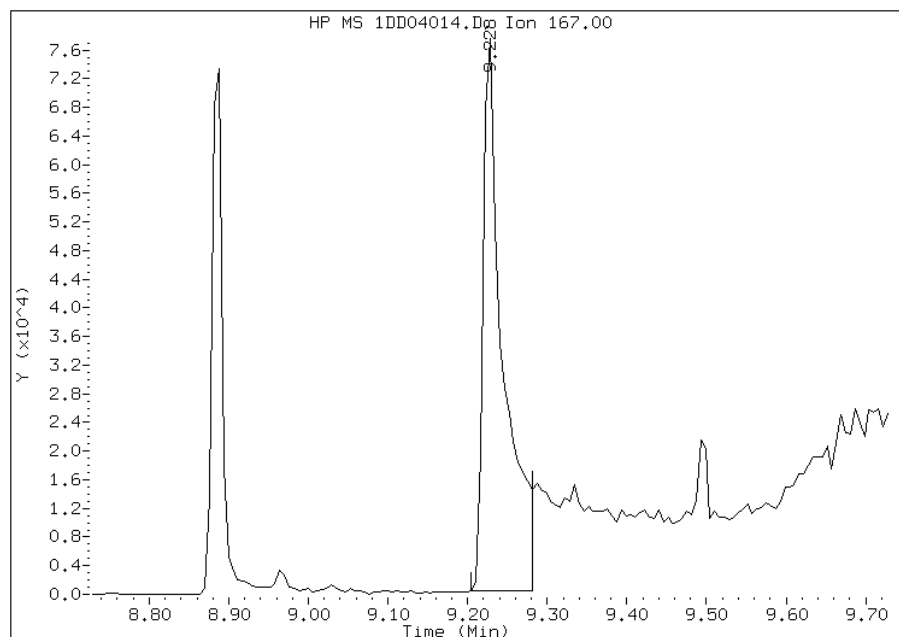


# Manual Integration Report

Data File: 1DD04014.D  
Inj. Date and Time: 04-APR-2013 16:27  
Instrument ID: BSMDS.i  
Client ID:  
Compound: 12 Carbazole  
CAS #: 86-74-8  
Report Date: 04/05/2013

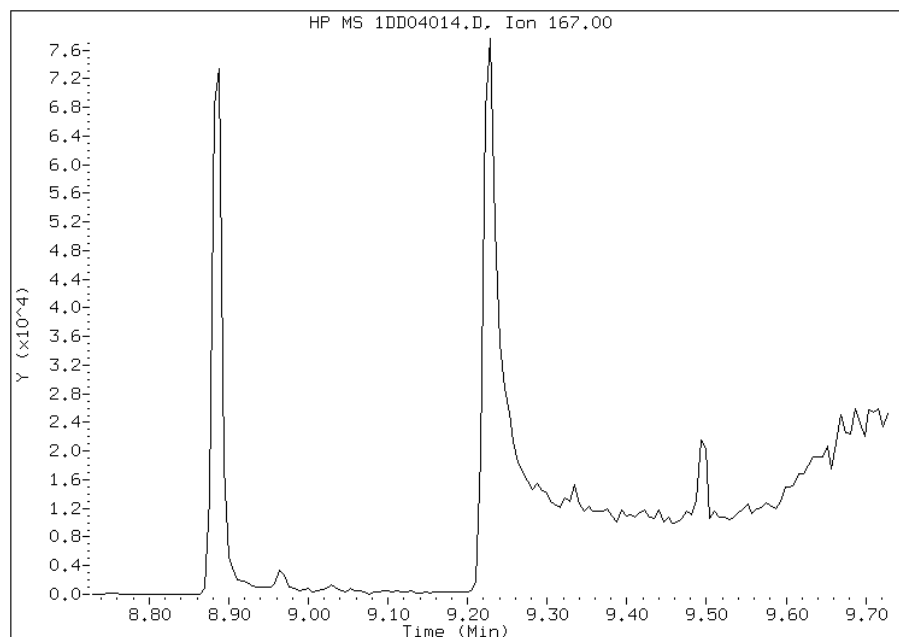
## Processing Integration Results

RT: 9.23  
Response: 136620  
Amount: 1  
Conc: 1



## Manual Integration Results

RT: 9.23  
Response: 1319041  
Amount: 14  
Conc: 14



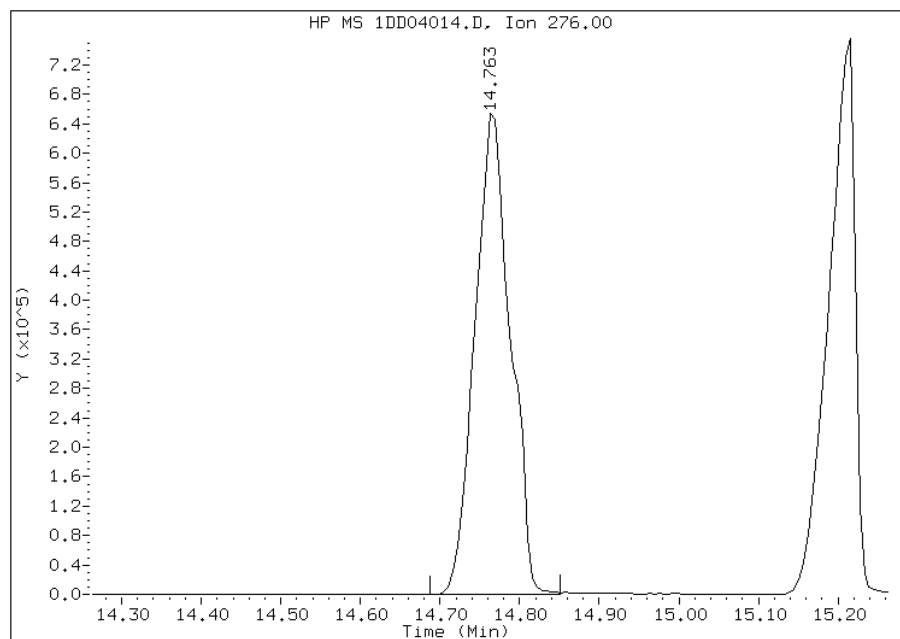
Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 13:08  
Manual Integration Reason: Baseline Event

Manual Integration Report

Data File: 1DD04014.D  
Inj. Date and Time: 04-APR-2013 16:27  
Instrument ID: BSMDS.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

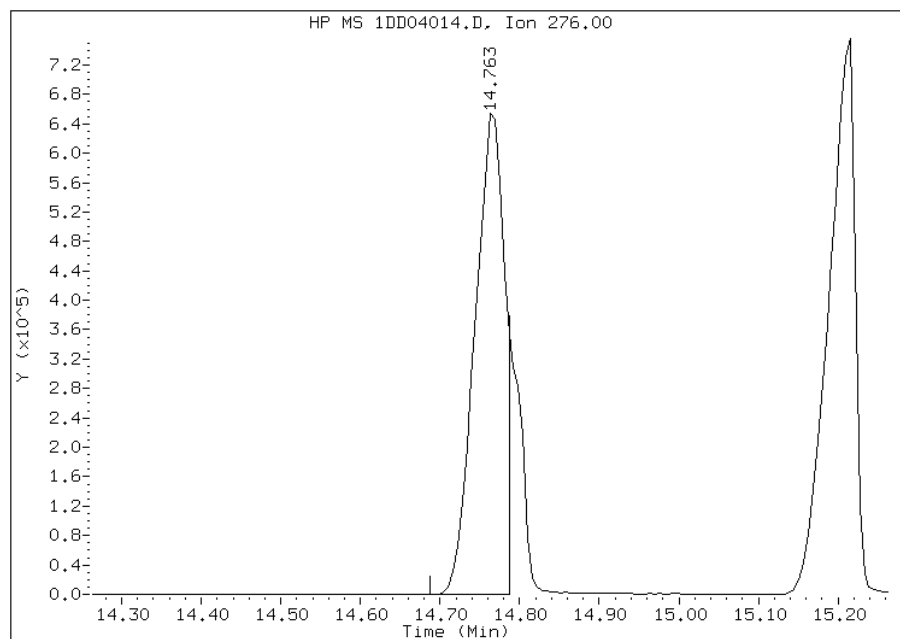
Processing Integration Results

RT: 14.76  
Response: 2024721  
Amount: 19  
Conc: 19



Manual Integration Results

RT: 14.76  
Response: 1694283  
Amount: 16  
Conc: 16



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 13:09  
Manual Integration Reason: Split Peak

TestAmerica Laboratories

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\1CD02002.D  
 Lab Smp Id: DFTPP Client Smp ID: DFTPP  
 Inj Date : 02-APR-2013 11:31  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : DFTPP-1525850  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040213.b\c-dftpp198.m  
 Meth Date : 04-Feb-2013 16:33 cantins Quant Type: ESTD  
 Cal Date : Cal File:  
 Als bottle: 2 QC Sample: DFTPP  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: all.sub  
 Target Version: 4.14 Sample Matrix: None  
 Processing Host: TAM1000

CONCENTRATIONS									
ON-COL FINAL									
RT	EXP RT	DLT RT	MASS	RESPONSE	( ug/L)	( ug/L)	TARGET	RANGE	RATIO
====	=====	=====	====	=====	=====	=====	=====	=====	=====
1 dftpp					CAS #: 5074-71-5				
7.310	7.469	-0.159	198	70432			50.00-	0.00	100.00
7.310	7.469	-0.159	51	24576			10.00-	80.00	34.89
7.310	7.469	-0.159	68	571			0.00-	2.00	1.62
7.310	7.469	-0.159	69	35176			0.00-	0.00	49.94
7.310	7.469	-0.159	70	308			0.00-	2.00	0.88
7.310	7.469	-0.159	127	29688			10.00-	80.00	42.15
7.310	7.469	-0.159	197	310			0.00-	2.00	0.44
7.310	7.469	-0.159	442	39944			50.00-	0.00	56.71
7.310	7.469	-0.159	199	5383			5.00-	9.00	7.64
7.310	7.469	-0.159	275	15117			10.00-	60.00	21.46
7.310	7.469	-0.159	365	2390			1.00-	0.00	3.39
7.310	7.469	-0.159	441	7169			0.01-	99.99	92.67
7.310	7.469	-0.159	443	7736			15.00-	24.00	19.37

Data File: 1CD02002.D

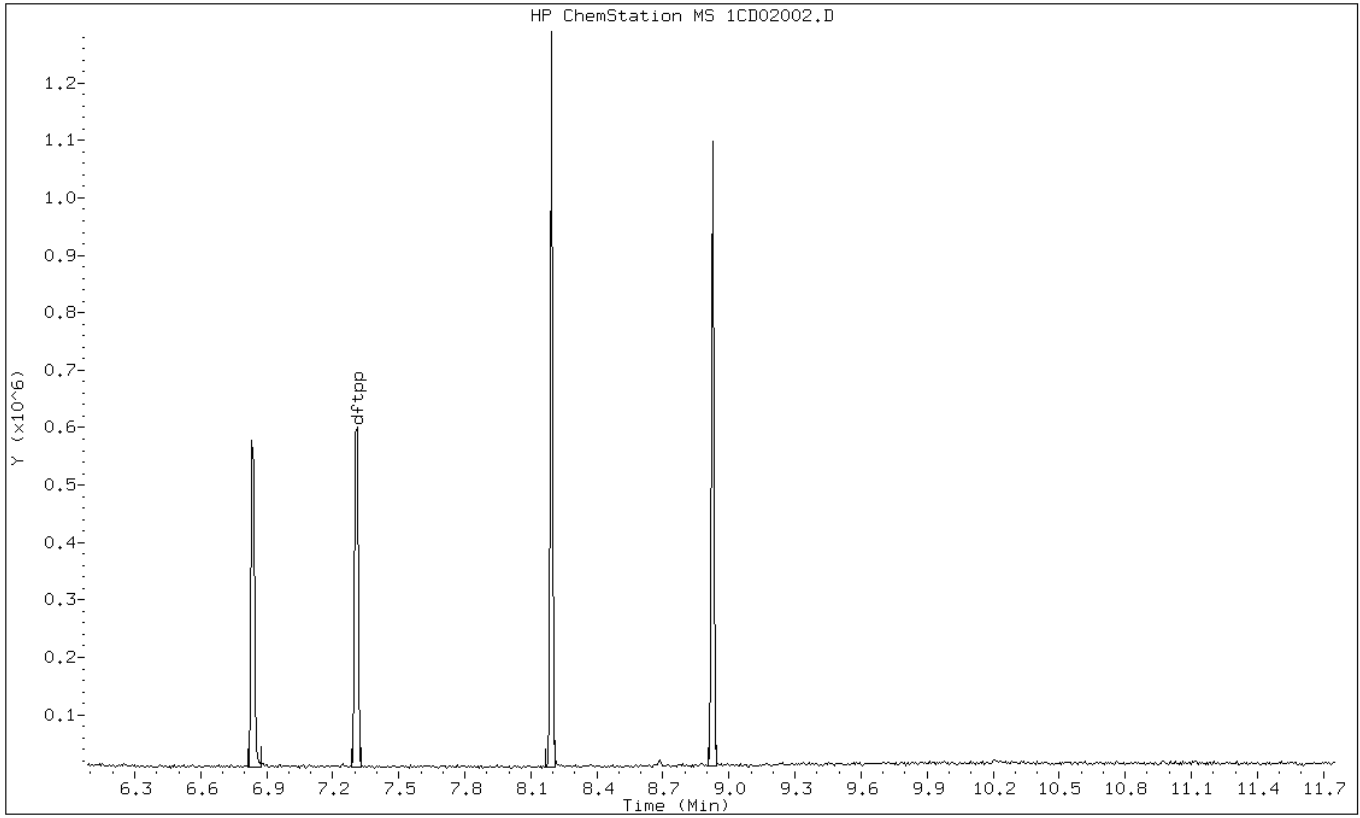
Date: 02-APR-2013 11:31

Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC



Data File: 1CD02002.D

Date: 02-APR-2013 11:31

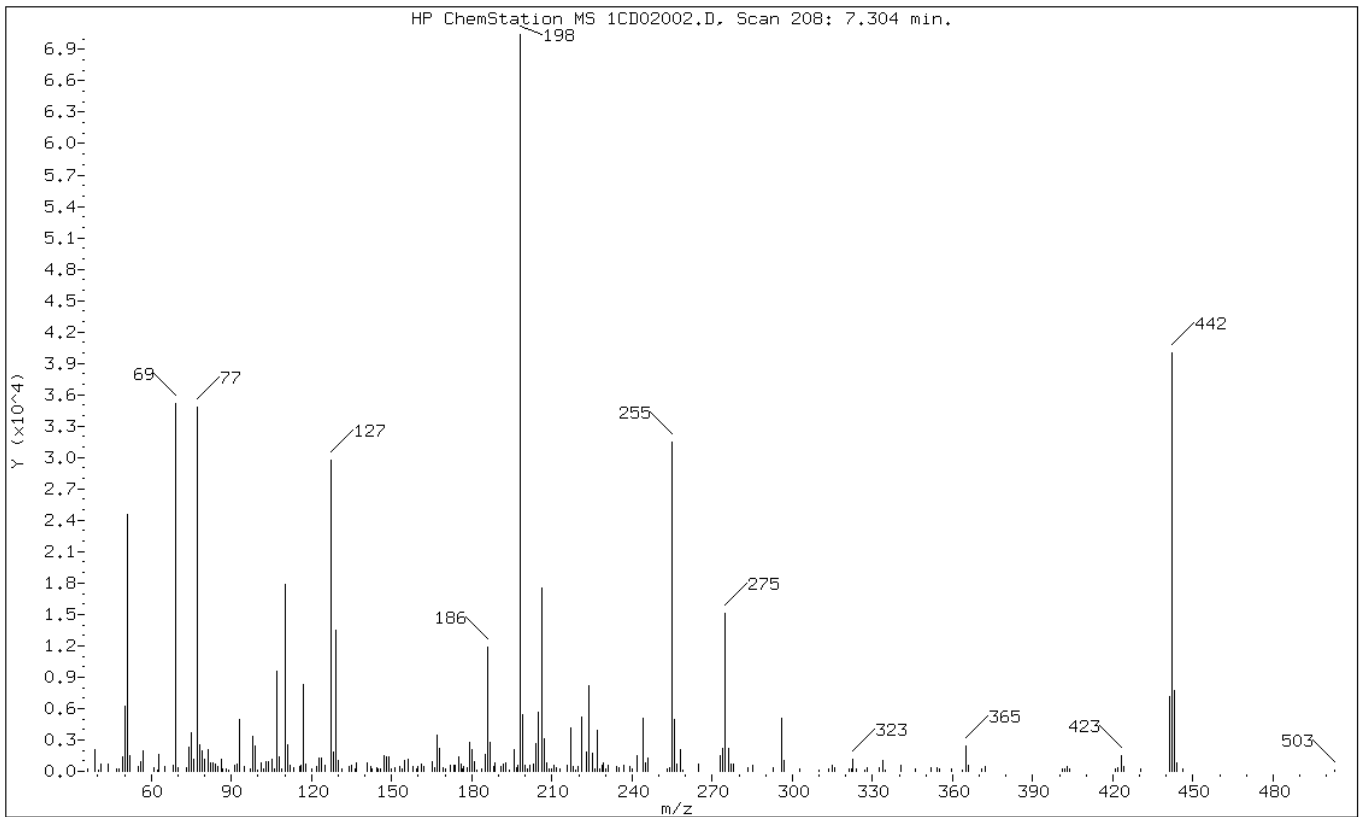
Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC

1 dftpp



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
198	Base Peak, 100% relative abundance	100.00
51	10.00 - 80.00% of mass 198	34.89
68	Less than 2.00% of mass 69	0.81 ( 1.62)
69	Mass 69 relative abundance	49.94
70	Less than 2.00% of mass 69	0.44 ( 0.88)
127	10.00 - 80.00% of mass 198	42.15
197	Less than 2.00% of mass 198	0.44
442	Greater than 50.00% of mass 198	56.71
199	5.00 - 9.00% of mass 198	7.64
275	10.00 - 60.00% of mass 198	21.46
365	Greater than 1.00% of mass 198	3.39
441	Present, but less than mass 442	10.18
443	15.00 - 24.00% of mass 442	10.98 ( 19.37)



Data File: 1CD02002.D

Date: 02-APR-2013 11:31

Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC

Data File: \\tam-chemsrv\chem\SM\BSMC5973.i\1C040213\_PAHIC.b\1CD02002.D

Spectrum: HP ChemStation MS 1CD02002.D, Scan 208: 7.304 min.

Location of Maximum: 198.00

Number of points: 229

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.20	191	113.10	351	185.10	1649	258.00	2060
39.00	2089	115.80	410	186.00	11880	259.00	166
40.10	156	116.20	563	187.00	2755	265.00	700
41.20	672	117.00	8338	188.30	505	273.00	1556
44.00	691	118.00	714	188.80	850	274.00	2191
46.90	264	120.20	251	190.90	451	275.00	15117
48.00	207	122.00	433	192.00	717	276.10	2178
49.10	1329	122.90	1302	192.90	774	276.90	747
50.10	6281	123.80	1270	193.90	161	278.10	714
51.10	24576	125.10	560	195.90	2063	283.20	367
52.10	1487	127.10	29688	196.70	310	285.10	604
55.00	486	128.00	1837	197.10	545	293.00	386
56.10	964	129.10	13517	198.00	70432	296.00	5053
57.00	1965	130.00	1041	199.00	5383	297.00	1014
60.80	304	131.20	273	200.10	567	302.80	285
62.30	156	134.00	480	200.60	270	310.10	151
63.00	1637	134.90	620	201.50	554	313.70	217
65.00	481	136.20	200	203.00	654	315.00	561
68.10	571	137.00	811	204.10	2706	316.00	397
69.00	35176	140.90	765	205.10	5687	321.20	252
69.90	308	142.10	410	206.10	17552	322.00	188
73.00	304	142.70	282	207.10	3108	322.80	1174
74.10	2331	144.30	362	208.00	798	324.00	267
75.00	3676	145.00	189	208.90	282	327.10	153
76.00	1155	145.90	247	210.00	219	328.20	395
77.10	34856	147.10	1448	210.90	584	332.70	292
78.10	2489	148.00	1427	211.50	320	333.90	1034
79.10	1952	149.00	1344	213.00	214	334.60	151
80.10	1105	150.00	235	215.70	551	340.80	534
81.10	2019	151.00	357	217.00	4128	346.10	272
82.00	853	153.00	443	217.90	509	352.10	376
83.00	779	153.90	266	218.80	152	354.20	383
83.80	657	155.00	984	219.60	431	354.90	200
84.90	486	156.00	1110	221.00	5183	359.50	267
86.10	1181	157.80	502	223.10	1793	363.80	168
86.90	260	159.30	205	224.00	8192	365.00	2390
88.00	245	159.90	477	225.20	1759	365.90	597
89.10	155	161.10	679	226.10	240	370.80	193
91.10	583	162.00	441	227.00	3893	372.00	411
92.10	667	165.10	934	227.90	218	401.00	218

93.00	5005	166.00	385	228.70	623	402.10	194
95.00	495	167.00	3405	229.10	783	402.90	407
96.90	195	168.00	2215	230.00	287	403.80	197
98.00	3343	169.20	374	231.10	622	420.70	267
99.00	2408	170.30	186	234.00	423	421.10	211
100.00	162	172.10	634	234.90	390	422.00	318
101.00	782	173.10	602	236.90	598	423.00	1535
102.10	189	173.70	532	239.10	486	424.00	439
103.10	884	175.10	1337	240.10	221	430.30	186
104.00	939	176.00	727	242.00	1442	441.00	7169
105.00	1194	176.60	217	244.10	5072	442.00	39944
106.00	180	177.10	501	245.20	829	443.00	7736
107.00	9612	178.10	387	246.00	1322	444.00	786
108.00	1350	179.00	2811	253.10	269	446.00	182
109.00	183	180.10	2065	254.10	289	503.00	171
110.00	17856	181.00	967	255.00	31424		
111.00	2511	181.80	164	256.00	4972		
112.10	622	183.90	209	256.90	650		

TestAmerica Laboratories

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03002.D  
 Lab Smp Id: DFTPP Client Smp ID: DFTPP  
 Inj Date : 03-APR-2013 11:28  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : DFTPP-1525850  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\c-dftpp198.m  
 Meth Date : 04-Feb-2013 16:33 cantins Quant Type: ESTD  
 Cal Date : Cal File:  
 Als bottle: 2 QC Sample: DFTPP  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: all.sub  
 Target Version: 4.14 Sample Matrix: None  
 Processing Host: TAM1000

CONCENTRATIONS									
ON-COL FINAL									
RT	EXP RT	DLT RT	MASS	RESPONSE ( ug/L)	( ug/L)	TARGET RANGE	RATIO		
====	=====	=====	====	=====	=====	=====	=====		
1 dftpp					CAS #: 5074-71-5				
7.304	7.469	-0.165	198	75560		50.00- 0.00	100.00		
7.304	7.469	-0.165	51	32256		10.00- 80.00	42.69		
7.304	7.469	-0.165	68	431		0.00- 2.00	1.15		
7.304	7.469	-0.165	69	37536		0.00- 0.00	49.68		
7.304	7.469	-0.165	70	0	0.0	0.0	0.00- 2.00	0.00	
7.304	7.469	-0.165	127	36336		10.00- 80.00	48.09		
7.304	7.469	-0.165	197	0	0.0	0.0	0.00- 2.00	0.00	
7.304	7.469	-0.165	442	46072		50.00- 0.00	60.97		
7.304	7.469	-0.165	199	4654		5.00- 9.00	6.16		
7.304	7.469	-0.165	275	14882		10.00- 60.00	19.70		
7.304	7.469	-0.165	365	1786		1.00- 0.00	2.36		
7.304	7.469	-0.165	441	5248		0.01- 99.99	68.32		
7.304	7.469	-0.165	443	7681		15.00- 24.00	16.67		

Data File: 1CD03002.D

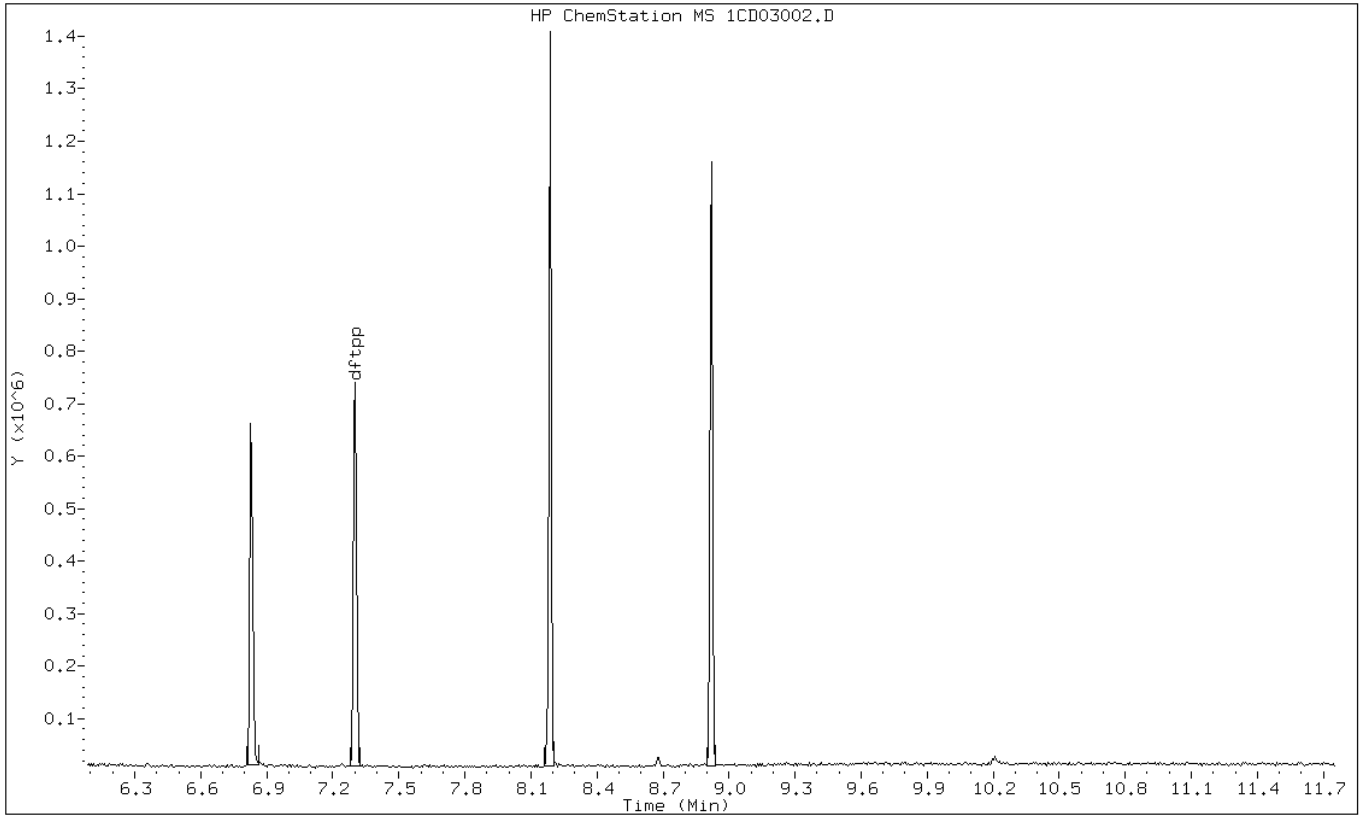
Date: 03-APR-2013 11:28

Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC



Data File: 1CD03002.D

Date: 03-APR-2013 11:28

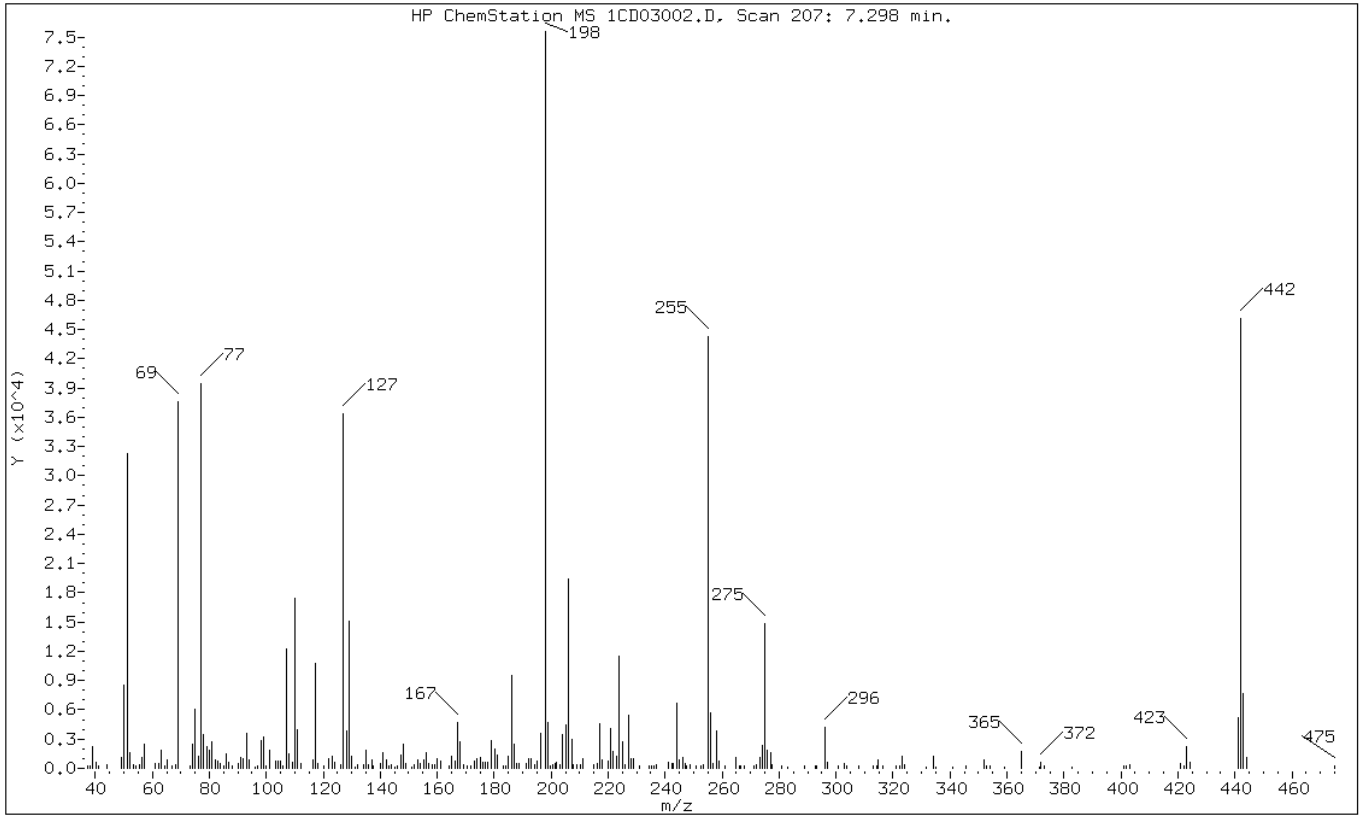
Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC

1 dftpp



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
198	Base Peak, 100% relative abundance	100.00
51	10.00 - 80.00% of mass 198	42.69
68	Less than 2.00% of mass 69	0.57 ( 1.15)
69	Mass 69 relative abundance	49.68
70	Less than 2.00% of mass 69	0.00 ( 0.00)
127	10.00 - 80.00% of mass 198	48.09
197	Less than 2.00% of mass 198	0.00
442	Greater than 50.00% of mass 198	60.97
199	5.00 - 9.00% of mass 198	6.16
275	10.00 - 60.00% of mass 198	19.70
365	Greater than 1.00% of mass 198	2.36
441	Present, but less than mass 443	6.95
443	15.00 - 24.00% of mass 442	10.17 ( 16.67)

Data File: 1CD03002.D

Date: 03-APR-2013 11:28

Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC

Data File: \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03002.D

Spectrum: HP ChemStation MS 1CD03002.D, Scan 207: 7.298 min.

Location of Maximum: 198.00

Number of points: 240

m/z	Y	m/z	Y	m/z	Y	m/z	Y
37.30	192	117.10	10764	188.00	493	266.00	218
38.00	283	118.00	527	188.90	513	266.70	201
39.10	2239	120.10	245	191.10	499	267.90	235
40.10	585	122.00	935	192.00	1043	271.10	199
40.90	297	123.10	1247	192.90	1024	272.10	327
44.10	330	123.90	604	194.10	332	273.10	1066
49.00	1129	126.00	335	195.20	727	274.00	2324
50.10	8491	127.10	36336	196.10	3641	275.00	14882
51.10	32256	128.10	3781	198.00	75560	276.00	1830
52.10	1581	129.00	15116	199.00	4654	277.00	1659
53.20	336	130.00	1263	199.90	276	277.70	402
54.10	256	131.20	159	200.60	335	281.00	220
55.30	348	132.00	334	201.20	451	282.80	180
56.20	1156	134.10	331	202.00	580	289.00	293
57.10	2512	135.10	1813	203.20	332	292.70	189
60.90	464	135.90	409	204.10	3436	293.20	284
62.10	453	137.00	852	205.00	4509	296.00	4228
63.00	1830	137.70	212	206.10	19472	297.00	626
64.20	214	140.10	437	207.10	2922	300.80	203
65.00	826	141.00	1569	207.90	382	303.00	545
66.80	196	142.10	863	209.10	415	303.80	201
68.00	431	143.00	262	210.20	312	308.00	262
69.10	37536	144.00	331	211.00	973	313.00	225
73.10	264	145.00	171	214.90	410	314.10	209
74.10	2431	146.00	305	216.00	506	314.90	815
75.10	6014	147.10	1301	217.00	4568	316.30	293
76.10	1227	148.10	2447	218.00	865	321.00	250
77.10	39480	149.00	456	220.20	718	322.20	285
78.00	3407	151.10	166	221.00	4124	323.00	1264
79.10	2246	152.10	368	221.70	1716	324.00	356
80.00	1895	153.00	900	223.10	1261	331.80	174
81.00	2668	154.10	515	224.00	11463	334.00	1232
82.10	879	155.10	863	225.10	2700	334.90	167
83.10	764	156.10	1663	226.00	418	340.90	178
84.00	475	157.00	538	227.00	5402	345.50	200
85.00	238	158.10	315	227.90	949	345.70	201
86.00	1545	159.10	361	229.00	1026	352.00	846
86.80	591	160.10	1033	231.00	287	352.90	228
88.00	244	161.10	710	234.20	305	354.00	283
90.00	555	164.00	222	235.10	301	359.10	153

90.90	1058	165.10	1243	236.10	289	364.90	1786
92.00	991	166.10	702	237.00	410	371.30	185
93.00	3605	167.00	4706	241.10	620	371.90	659
94.10	821	168.10	2729	242.20	454	373.00	302
96.10	182	169.20	408	242.90	481	382.70	163
97.10	261	170.70	259	244.00	6719	400.90	212
98.00	2850	171.80	303	245.00	898	402.00	306
99.10	3237	173.10	799	246.10	1119	403.00	361
100.10	191	174.10	994	247.10	479	420.80	515
101.10	1802	175.10	1080	247.60	278	422.10	230
103.10	791	176.10	603	248.80	422	422.90	2199
104.10	795	177.00	595	250.70	208	424.10	620
105.10	697	177.60	617	252.60	245	441.00	5248
105.70	251	179.10	2842	253.60	315	442.00	46072
107.00	12235	180.10	1998	255.00	44296	443.00	7681
108.10	1536	180.90	1327	256.10	5661	444.00	1055
109.10	670	183.40	224	256.80	459	475.00	248
110.10	17424	184.10	220	258.00	3829		
111.00	3940	185.00	1287	259.10	709		
112.20	440	186.10	9570	260.90	205		
116.20	866	187.10	2436	265.00	1167		

TestAmerica Laboratories

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\1CD05003.D  
 Lab Smp Id: DFTPP Client Smp ID: DFTPP  
 Inj Date : 05-APR-2013 11:57  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : DFTPP-1525850  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\c-dftpp198.m  
 Meth Date : 04-Feb-2013 16:33 cantins Quant Type: ESTD  
 Cal Date : Cal File:  
 Als bottle: 2 QC Sample: DFTPP  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: all.sub  
 Target Version: 4.14 Sample Matrix: None  
 Processing Host: TAM1000

CONCENTRATIONS									
ON-COL FINAL									
RT	EXP RT	DLT RT	MASS	RESPONSE	( ug/L)	( ug/L)	TARGET	RANGE	RATIO
====	=====	=====	====	=====	=====	=====	=====	=====	=====
1 dftpp					CAS #: 5074-71-5				
7.286	7.469	-0.183	198	70588			50.00-	0.00	100.00
7.286	7.469	-0.183	51	29336			10.00-	80.00	41.56
7.286	7.469	-0.183	68	565			0.00-	2.00	1.45
7.286	7.469	-0.183	69	39020			0.00-	0.00	55.28
7.286	7.469	-0.183	70	218			0.00-	2.00	0.56
7.286	7.469	-0.183	127	34576			10.00-	80.00	48.98
7.286	7.469	-0.183	197	438			0.00-	2.00	0.62
7.286	7.469	-0.183	442	39248			50.00-	0.00	55.60
7.286	7.469	-0.183	199	4704			5.00-	9.00	6.66
7.286	7.469	-0.183	275	13612			10.00-	60.00	19.28
7.286	7.469	-0.183	365	2087			1.00-	0.00	2.96
7.286	7.469	-0.183	441	5332			0.01-	99.99	64.58
7.286	7.469	-0.183	443	8257			15.00-	24.00	21.04



Data File: 1CD05003.D

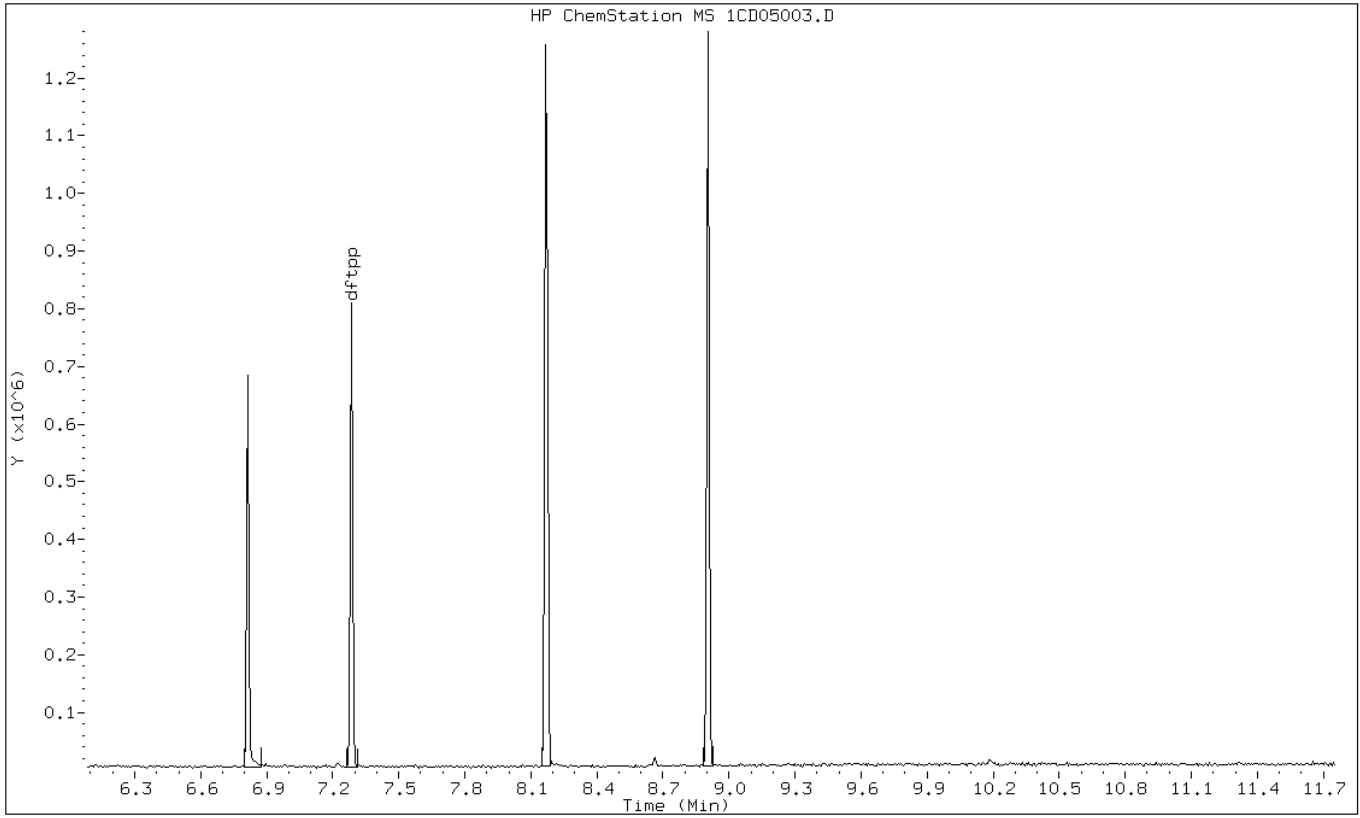
Date: 05-APR-2013 11:57

Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC



Data File: 1CD05003.D

Date: 05-APR-2013 11:57

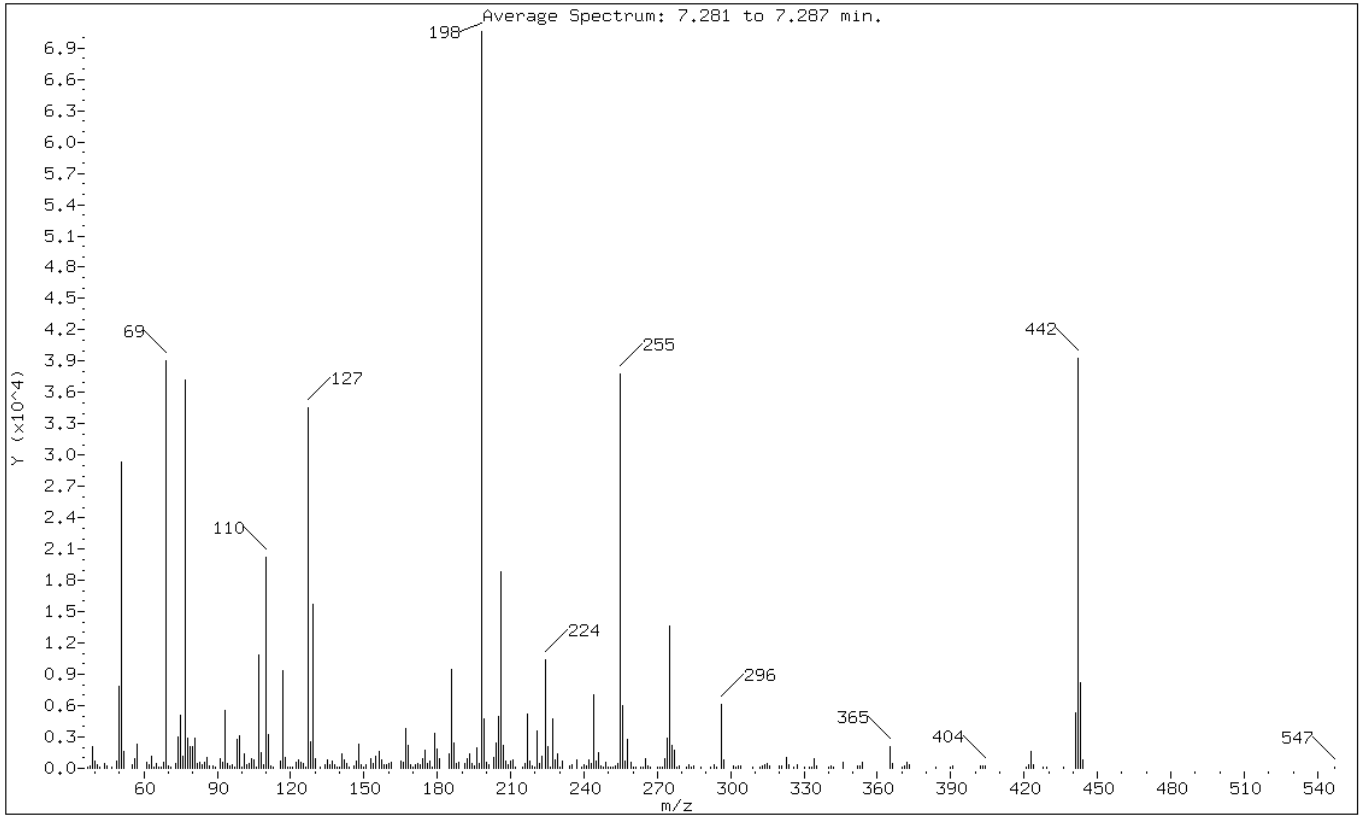
Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC

1 dftpp



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
198	Base Peak, 100% relative abundance	100.00
51	10.00 - 80.00% of mass 198	41.56
68	Less than 2.00% of mass 69	0.80 ( 1.45)
69	Mass 69 relative abundance	55.28
70	Less than 2.00% of mass 69	0.31 ( 0.56)
127	10.00 - 80.00% of mass 198	48.98
197	Less than 2.00% of mass 198	0.62
442	Greater than 50.00% of mass 198	55.60
199	5.00 - 9.00% of mass 198	6.66
275	10.00 - 60.00% of mass 198	19.28
365	Greater than 1.00% of mass 198	2.96
441	Present, but less than mass 443	7.55
443	15.00 - 24.00% of mass 442	11.70 ( 21.04)

Data File: 1CD05003.D

Date: 05-APR-2013 11:57

Client ID: DFTPP

Instrument: BSMC5973.i

Sample Info: DFTPP-1525850

Operator: SCC

Data File: \\tam-chemsrv\chem\SM\BSMC5973.i\1C040513.b\1CD05003.D

Spectrum: Average Spectrum: 7.281 to 7.287 min.

Location of Maximum: 198.00

Number of points: 272

m/z	Y	m/z	Y	m/z	Y	m/z	Y
37.00	87	118.00	1005	198.00	70584	277.00	1778
38.00	215	119.00	77	199.00	4704	278.00	123
39.00	2084	120.00	109	200.00	535	279.00	249
40.00	685	121.00	95	201.00	377	282.00	110
41.00	373	122.00	557	203.00	1061	283.00	366
42.00	99	123.00	824	204.00	2377	284.00	78
44.00	466	124.00	525	205.00	4989	285.00	223
45.00	227	125.00	459	206.00	18816	288.00	110
47.00	162	126.00	156	207.00	2185	292.00	134
49.00	724	127.00	34576	208.00	693	293.00	289
50.00	7845	128.00	2580	209.00	314	294.00	163
51.00	29336	129.00	15715	210.00	666	296.00	6099
52.00	1655	130.00	920	211.00	776	297.00	790
55.00	364	132.00	129	212.00	124	301.00	224
56.00	896	134.00	387	215.00	121	302.00	162
57.00	2300	135.00	780	216.00	407	303.00	275
61.00	566	136.00	313	217.00	5252	304.00	252
62.00	316	137.00	703	218.00	694	309.00	166
63.00	1159	138.00	398	219.00	258	312.00	119
64.00	78	139.00	81	220.00	136	313.00	204
65.00	468	140.00	81	221.00	3530	314.00	309
66.00	82	141.00	1344	222.00	504	315.00	458
67.00	138	142.00	755	223.00	1120	316.00	213
68.00	565	143.00	438	224.00	10390	320.00	205
69.00	39016	144.00	75	225.00	2068	321.00	246
70.00	218	146.00	214	226.00	144	323.00	1081
71.00	79	147.00	650	227.00	4743	324.00	400
73.00	429	148.00	2309	228.00	766	326.00	99
74.00	3044	149.00	380	229.00	1355	327.00	399
75.00	5071	150.00	82	230.00	78	330.00	118
76.00	1173	151.00	300	231.00	697	332.00	99
77.00	37208	153.00	927	234.00	233	333.00	94
78.00	2848	154.00	467	235.00	309	334.00	919
79.00	2133	155.00	1200	237.00	759	335.00	218
80.00	2030	156.00	1561	239.00	78	340.00	87
81.00	2919	157.00	859	240.00	290	341.00	188
82.00	460	158.00	326	241.00	276	342.00	147
83.00	534	159.00	358	242.00	757	346.00	613
84.00	344	160.00	508	243.00	471	352.00	275
85.00	549	161.00	601	244.00	7050	353.00	186

86.00	1030	165.00	731	245.00	670	354.00	548
87.00	176	166.00	623	246.00	1507	365.00	2087
88.00	257	167.00	3864	247.00	284	366.00	490
89.00	87	168.00	2200	248.00	109	370.00	147
91.00	945	169.00	331	249.00	616	371.00	183
92.00	633	170.00	165	250.00	141	372.00	601
93.00	5573	171.00	295	251.00	99	373.00	335
94.00	433	172.00	412	252.00	82	384.00	140
95.00	219	173.00	329	253.00	183	390.00	75
96.00	357	174.00	969	254.00	496	391.00	206
97.00	94	175.00	1726	255.00	37768	402.00	252
98.00	2760	176.00	453	256.00	6014	403.00	188
99.00	3086	177.00	636	257.00	656	404.00	274
100.00	108	178.00	167	258.00	2749	421.00	79
101.00	1360	179.00	3315	259.00	560	422.00	289
102.00	383	180.00	1844	260.00	94	423.00	1582
103.00	417	181.00	957	261.00	110	424.00	356
104.00	900	185.00	1392	263.00	75	428.00	103
105.00	865	186.00	9523	264.00	82	429.00	102
106.00	162	187.00	2465	265.00	976	436.00	121
107.00	10874	188.00	440	266.00	191	441.00	5332
108.00	1494	189.00	611	267.00	142	442.00	39248
109.00	397	191.00	453	270.00	87	443.00	8257
110.00	20224	192.00	872	271.00	101	444.00	752
111.00	3238	193.00	1334	272.00	155	547.00	127
112.00	219	194.00	408	273.00	972		
113.00	166	195.00	217	274.00	2900		
116.00	660	196.00	1965	275.00	13612		
117.00	9344	197.00	438	276.00	2248		

TestAmerica Laboratories

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04003.D  
 Lab Smp Id: DFTPP Client Smp ID: DFTPP  
 Inj Date : 04-APR-2013 12:15  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : DFTPP-1525850  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\d-dftpp198.m  
 Meth Date : 08-Jan-2013 12:23 cantins Quant Type: ESTD  
 Cal Date : Cal File:  
 Als bottle: 2 QC Sample: DFTPP  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: all.sub  
 Target Version: 4.14 Sample Matrix: None  
 Processing Host: TAM1000

CONCENTRATIONS									
ON-COL FINAL									
RT	EXP RT	DLT RT	MASS	RESPONSE	( ug/L)	( ug/L)	TARGET	RANGE	RATIO
====	=====	=====	====	=====	=====	=====	=====	=====	=====
1 dftpp					CAS #: 5074-71-5				
8.382	8.532	-0.150	198	72572			50.00-	0.00	100.00
8.382	8.532	-0.150	51	32556			10.00-	80.00	44.86
8.382	8.532	-0.150	68	0	0.0	0.0	0.00-	2.00	0.00
8.382	8.532	-0.150	69	32936			0.00-	0.00	45.38
8.382	8.532	-0.150	70	114			0.00-	2.00	0.35
8.382	8.532	-0.150	127	36680			10.00-	80.00	50.54
8.382	8.532	-0.150	197	0	0.0	0.0	0.00-	2.00	0.00
8.382	8.532	-0.150	442	48716			50.00-	0.00	67.13
8.382	8.532	-0.150	199	4977			5.00-	9.00	6.86
8.382	8.532	-0.150	275	19350			10.00-	60.00	26.66
8.382	8.532	-0.150	365	2279			1.00-	0.00	3.14
8.382	8.532	-0.150	441	2370			0.01-	99.99	23.58
8.382	8.532	-0.150	443	10052			15.00-	24.00	20.63

Data File: 1DD04003.D

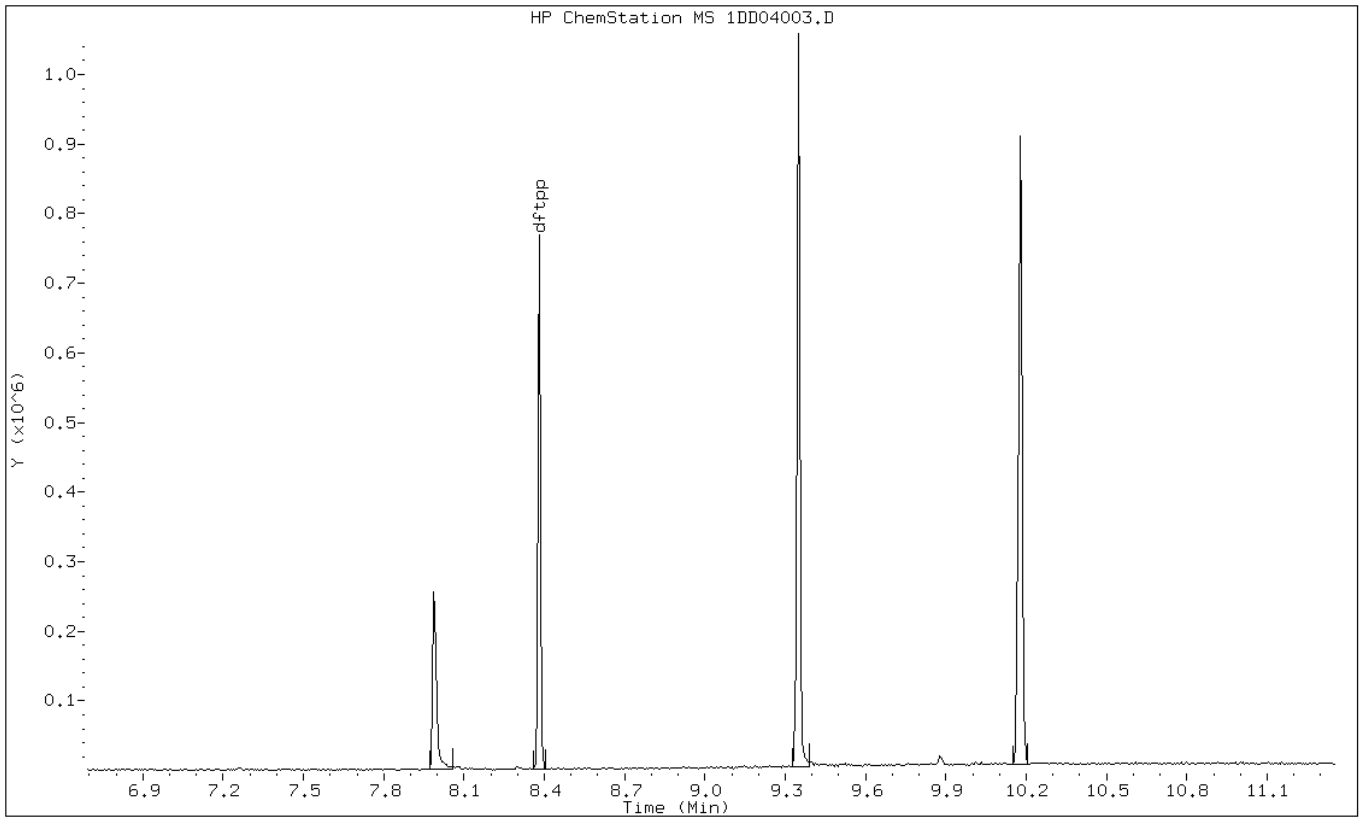
Date: 04-APR-2013 12:15

Client ID: DFTPP

Instrument: BSMSD.i

Sample Info: DFTPP-1525850

Operator: SCC



Data File: 1DD04003.D

Date: 04-APR-2013 12:15

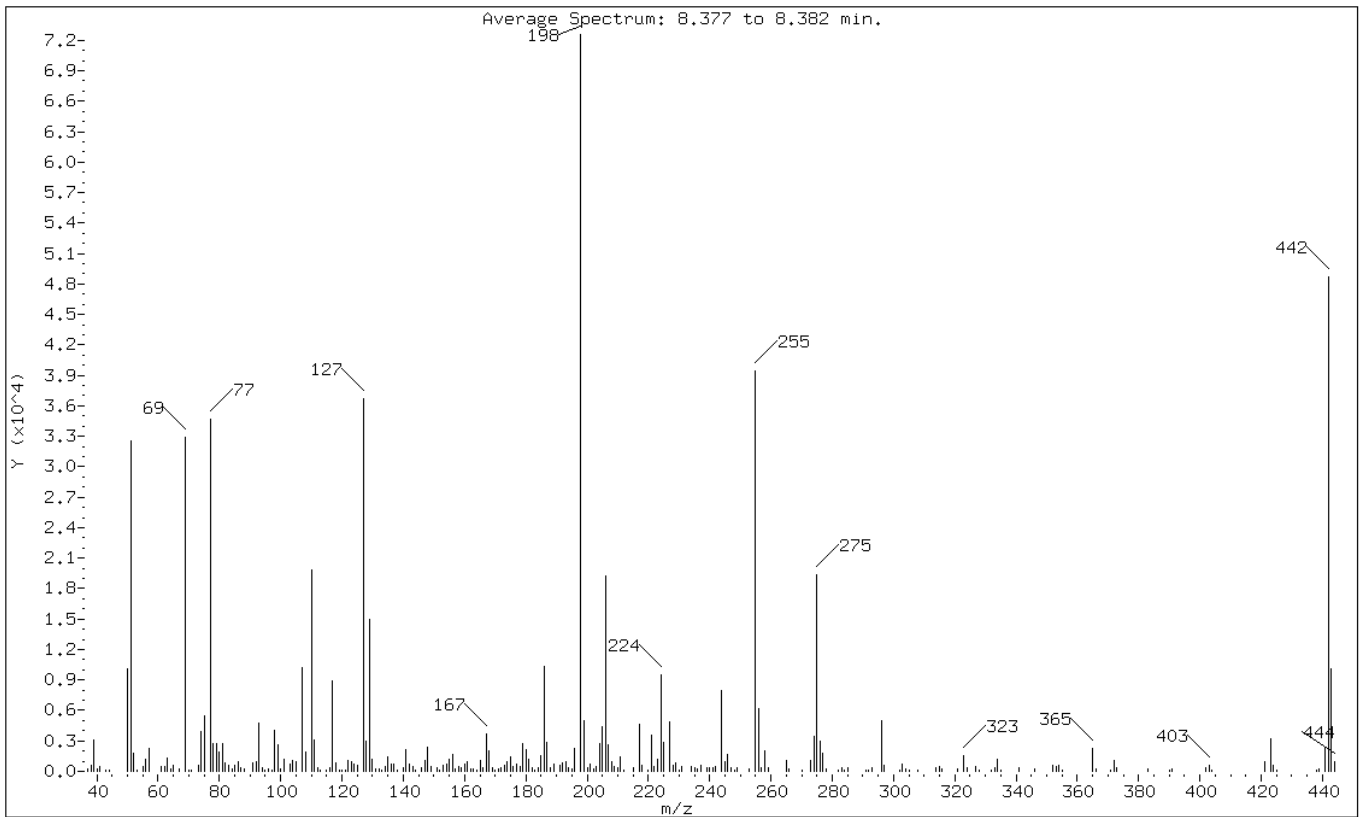
Client ID: DFTPP

Instrument: BSMSD.i

Sample Info: DFTPP-1525850

Operator: SCC

1 dftpp



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
198	Base Peak, 100% relative abundance	100.00
51	10.00 - 80.00% of mass 198	44.86
68	Less than 2.00% of mass 69	0.00 ( 0.00)
69	Mass 69 relative abundance	45.38
70	Less than 2.00% of mass 69	0.16 ( 0.35)
127	10.00 - 80.00% of mass 198	50.54
197	Less than 2.00% of mass 198	0.00
442	Greater than 50.00% of mass 198	67.13
199	5.00 - 9.00% of mass 198	6.86
275	10.00 - 60.00% of mass 198	26.66
365	Greater than 1.00% of mass 198	3.14
441	Present, but less than mass 443	3.27
443	15.00 - 24.00% of mass 442	13.85 ( 20.63)

Data File: 1DD04003.D

Date: 04-APR-2013 12:15

Client ID: DFTPP

Instrument: BSMSD.i

Sample Info: DFTPP-1525850

Operator: SCC

Data File: \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04003.D

Spectrum: Average Spectrum: 8.377 to 8.382 min.

Location of Maximum: 198.00

Number of points: 246

m/z	Y	m/z	Y	m/z	Y	m/z	Y
37.00	274	119.00	120	185.00	1517	270.00	78
38.00	589	120.00	118	186.00	10284	273.00	1081
39.00	3038	121.00	77	187.00	2888	274.00	3485
40.00	277	122.00	1015	188.00	332	275.00	19344
41.00	463	123.00	946	189.00	735	276.00	2999
43.00	124	124.00	666	191.00	579	277.00	1839
44.00	117	125.00	567	192.00	873	278.00	226
50.00	10128	127.00	36680	193.00	975	282.00	81
51.00	32552	128.00	2957	194.00	335	283.00	314
52.00	1767	129.00	14951	195.00	275	284.00	90
53.00	85	130.00	1205	196.00	2233	285.00	356
55.00	420	131.00	194	198.00	72568	291.00	83
56.00	1176	132.00	206	199.00	4977	292.00	80
57.00	2213	133.00	92	200.00	323	293.00	412
61.00	490	134.00	523	201.00	663	296.00	5046
62.00	459	135.00	1404	202.00	210	297.00	576
63.00	1290	136.00	674	203.00	519	302.00	157
64.00	230	137.00	709	204.00	2685	303.00	675
65.00	539	138.00	79	205.00	4398	304.00	185
67.00	251	140.00	333	206.00	19200	305.00	82
69.00	32936	141.00	2082	207.00	2631	308.00	174
70.00	114	142.00	713	208.00	974	314.00	314
71.00	81	143.00	523	209.00	499	315.00	487
73.00	647	144.00	93	210.00	329	316.00	223
74.00	3962	146.00	312	211.00	1393	321.00	206
75.00	5478	147.00	1032	212.00	165	323.00	1494
77.00	34688	148.00	2326	215.00	308	324.00	410
78.00	2711	149.00	488	217.00	4596	327.00	476
79.00	2695	151.00	320	218.00	606	328.00	99
80.00	1923	152.00	103	220.00	76	332.00	111
81.00	2677	153.00	558	221.00	3596	333.00	396
82.00	777	154.00	665	222.00	431	334.00	1163
83.00	630	155.00	1227	223.00	1208	335.00	119
84.00	185	156.00	1628	224.00	9447	341.00	297
85.00	566	157.00	240	225.00	2804	346.00	197
86.00	895	158.00	430	227.00	4861	352.00	557
87.00	384	159.00	320	228.00	637	353.00	477
88.00	184	160.00	765	229.00	843	354.00	558
91.00	856	161.00	1005	230.00	115	355.00	81
92.00	893	162.00	279	231.00	446	365.00	2279



93.00	4736	163.00	190	234.00	485	366.00	181
94.00	298	164.00	105	235.00	402	371.00	117
95.00	167	165.00	1019	236.00	243	372.00	1076
96.00	240	166.00	344	237.00	537	373.00	335
97.00	178	167.00	3671	239.00	320	383.00	219
+-----+							
98.00	4066	168.00	1997	240.00	333	390.00	136
99.00	2655	169.00	349	241.00	361	391.00	180
100.00	295	170.00	112	242.00	472	402.00	362
101.00	1142	171.00	208	244.00	7939	403.00	564
103.00	719	172.00	342	245.00	988	404.00	144
+-----+							
104.00	1122	173.00	643	246.00	1619	421.00	961
105.00	909	174.00	893	247.00	381	423.00	3222
107.00	10195	175.00	1368	248.00	80	424.00	628
108.00	1940	176.00	519	249.00	382	425.00	87
110.00	19784	177.00	713	253.00	265	438.00	129
+-----+							
111.00	3136	178.00	422	255.00	39432	439.00	214
112.00	374	179.00	2728	256.00	6151	441.00	2370
113.00	128	180.00	2151	257.00	340	442.00	48712
115.00	153	181.00	1200	258.00	2068	443.00	10052
116.00	393	182.00	314	259.00	399	444.00	994
+-----+							
117.00	8897	183.00	98	265.00	1086		
118.00	800	184.00	382	266.00	282		
+-----+							

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: \_\_\_\_\_ Lab Sample ID: MB 660-136013/1-A  
 Matrix: Water Lab File ID: 1DD04022.D  
 Analysis Method: 8270C LL Date Collected: \_\_\_\_\_  
 Extract. Method: 3520C Date Extracted: 04/02/2013 08:08  
 Sample wt/vol: 1000(mL) Date Analyzed: 04/04/2013 19:51  
 Con. Extract Vol.: 1(mL) Dilution Factor: 1  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136164 Units: ug/L

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	2.0	U	2.0	0.50
208-96-8	Acenaphthylene	1.0	U	1.0	0.25
120-12-7	Anthracene	0.20	U	0.20	0.076
56-55-3	Benzo[a]anthracene	0.20	U	0.20	0.050
50-32-8	Benzo[a]pyrene	0.20	U	0.20	0.057
205-99-2	Benzo[b]fluoranthene	0.20	U	0.20	0.050
191-24-2	Benzo[g,h,i]perylene	0.50	U	0.50	0.10
207-08-9	Benzo[k]fluoranthene	0.20	U	0.20	0.057
218-01-9	Chrysene	0.20	U	0.20	0.069
53-70-3	Dibenz(a,h)anthracene	0.20	U	0.20	0.050
206-44-0	Fluoranthene	0.50	U	0.50	0.054
86-73-7	Fluorene	2.0	U	2.0	0.50
193-39-5	Indeno[1,2,3-cd]pyrene	0.20	U	0.20	0.050
90-12-0	1-Methylnaphthalene	2.0	U	2.0	0.50
91-57-6	2-Methylnaphthalene	2.0	U	2.0	0.50
91-20-3	Naphthalene	2.0	U	2.0	0.25
85-01-8	Phenanthrene	0.50	U	0.50	0.20
129-00-0	Pyrene	0.50	U	0.50	0.089

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	78		30-130

TestAmerica Laboratories

Semivolatiles 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04022.D  
 Lab Smp Id: MB 660-136013/1-A  
 Inj Date : 04-APR-2013 19:51  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : MB 660-136013/1-A  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dFASTPAHi.m  
 Meth Date : 05-Apr-2013 13:07 cantins Quant Type: ISTD  
 Cal Date : 04-APR-2013 16:04 Cal File: 1DD04013.D  
 Als bottle: 20 QC Sample: BLANK  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	1000.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
							ON-COLUMN ( ug/l)	FINAL ( ug/l)
* 1 Naphthalene-d8		136	6.089	6.090	(1.000)	2711590	40.0000	
* 6 Acenaphthene-d10		164	7.769	7.770	(1.000)	1656380	40.0000	
* 9 Phenanthrene-d10		188	9.027	9.028	(1.000)	2762536	40.0000	
\$ 13 o-Terphenyl		230	9.332	9.339	(1.034)	325411	7.81784	7.8
* 17 Chrysene-d12		240	11.336	11.349	(1.000)	2767376	40.0000	
* 22 Perylene-d12		264	13.169	13.176	(1.000)	2802105	40.0000	

Data File: 1DD04022.D

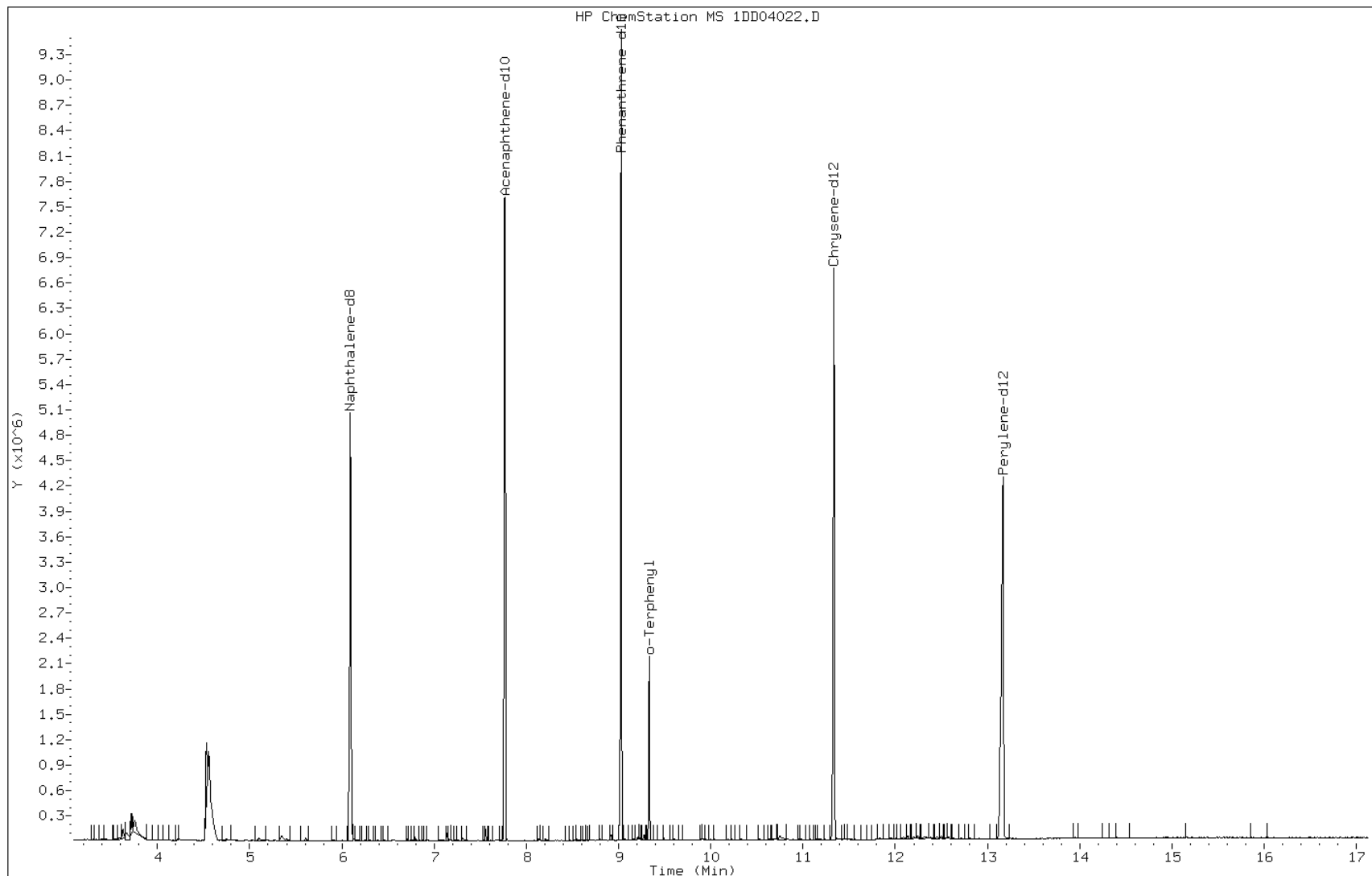
Date: 04-APR-2013 19:51

Client ID:

Instrument: BSMSD.i

Sample Info: MB 660-136013/1-A

Operator: SCC



FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: \_\_\_\_\_ Lab Sample ID: MB 660-136063/1-A  
 Matrix: Solid Lab File ID: 1CD03015.D  
 Analysis Method: 8270C LL Date Collected: \_\_\_\_\_  
 Extract. Method: 3546 Date Extracted: 04/02/2013 11:33  
 Sample wt/vol: 14.99(g) Date Analyzed: 04/03/2013 15:34  
 Con. Extract Vol.: 1(mL) Dilution Factor: 1  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136081 Units: ug/Kg

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	100	U	100	20
208-96-8	Acenaphthylene	40	U	40	5.0
120-12-7	Anthracene	8.4	U	8.4	4.2
56-55-3	Benzo[a]anthracene	8.0	U	8.0	3.9
50-32-8	Benzo[a]pyrene	10	U	10	5.2
205-99-2	Benzo[b]fluoranthene	12	U	12	6.1
191-24-2	Benzo[g,h,i]perylene	20	U	20	4.4
207-08-9	Benzo[k]fluoranthene	8.0	U	8.0	3.6
218-01-9	Chrysene	9.0	U	9.0	4.5
53-70-3	Dibenz(a,h)anthracene	20	U	20	4.1
206-44-0	Fluoranthene	20	U	20	4.0
86-73-7	Fluorene	20	U	20	4.1
193-39-5	Indeno[1,2,3-cd]pyrene	20	U	20	7.1
90-12-0	1-Methylnaphthalene	40	U	40	4.4
91-57-6	2-Methylnaphthalene	40	U	40	7.1
91-20-3	Naphthalene	40	U	40	4.4
85-01-8	Phenanthrene	8.0	U	8.0	3.9
129-00-0	Pyrene	20	U	20	3.7

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	80		30-130

TestAmerica Laboratories

Semivolatile 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03015.D  
 Lab Smp Id: mb 660-136063/1-a  
 Inj Date : 03-APR-2013 15:34  
 Operator : SCC  
 Smp Info : mb 660-136063/1-a  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\a-bFASTPAHi-m.m  
 Meth Date : 03-Apr-2013 11:59 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 15 QC Sample: BLANK  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula:

$$\text{Amt} * \text{DF} * 1/\text{Vi} * \text{Vt}/\text{Ws} * 100/(100 - \text{M}) * \text{A} * \text{B} * \text{C} * \text{D} * \text{GPC} * \text{CpndVariable}$$

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Ws	14.990	Weight Extracted
M	0.00000	% Moisture
A	1000.000	uL to mL conversion
B	1000.000	g to kg conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1 if no conv)
GPC	1.000	GPC FACTOR
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
							ON-COLUMN	FINAL
	MASS						(ug/ml)	(ug/Kg)
* 1 Naphthalene-d8	136		3.710	3.704	(1.000)	646271	40.0000	
* 6 Acenaphthene-d10	164		4.798	4.792	(1.000)	483289	40.0000	
* 10 Phenanthrene-d10	188		5.745	5.739	(1.000)	879169	40.0000	
\$ 14 o-Terphenyl	230		5.992	5.992	(1.043)	104839	8.03116	535.7676
* 18 Chrysene-d12	240		7.680	7.680	(1.000)	990236	40.0000	
* 23 Perylene-d12	264		8.856	8.851	(1.000)	958431	40.0000	

Data File: 1CD03015.D

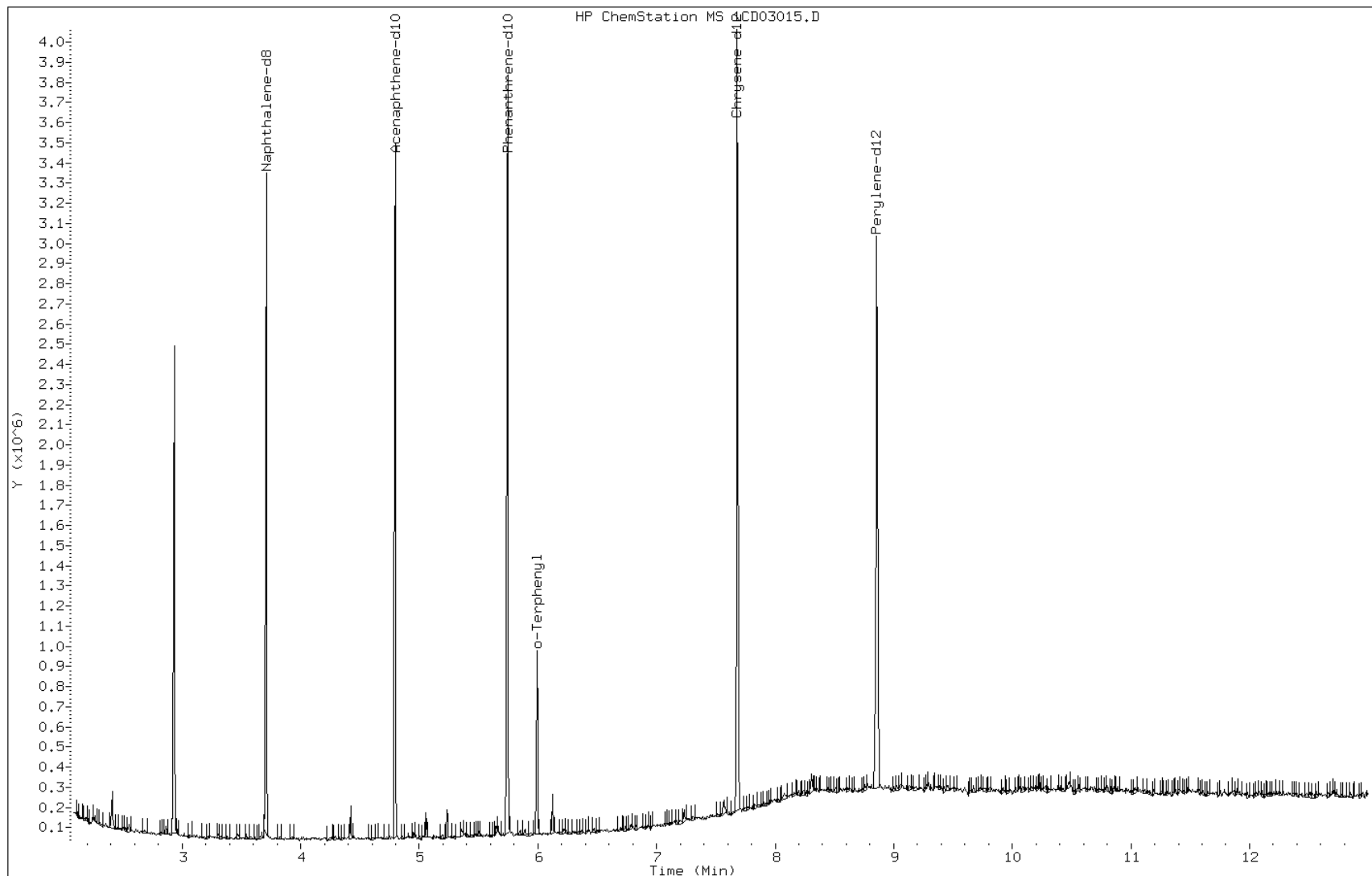
Date: 03-APR-2013 15:34

Client ID:

Instrument: BSMC5973.i

Sample Info: mb 660-136063/1-a

Operator: SCC



FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: \_\_\_\_\_ Lab Sample ID: LCS 660-136013/2-A  
 Matrix: Water Lab File ID: 1DD04023.D  
 Analysis Method: 8270C LL Date Collected: \_\_\_\_\_  
 Extract. Method: 3520C Date Extracted: 04/02/2013 08:08  
 Sample wt/vol: 1000(mL) Date Analyzed: 04/04/2013 20:13  
 Con. Extract Vol.: 1(mL) Dilution Factor: 1  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136164 Units: ug/L

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	7.38		2.0	0.50
208-96-8	Acenaphthylene	7.55		1.0	0.25
120-12-7	Anthracene	7.12		0.20	0.076
56-55-3	Benzo[a]anthracene	7.21		0.20	0.050
50-32-8	Benzo[a]pyrene	4.96		0.20	0.057
205-99-2	Benzo[b]fluoranthene	5.86		0.20	0.050
191-24-2	Benzo[g,h,i]perylene	3.51		0.50	0.10
207-08-9	Benzo[k]fluoranthene	5.75		0.20	0.057
218-01-9	Chrysene	7.28		0.20	0.069
53-70-3	Dibenz(a,h)anthracene	3.33		0.20	0.050
206-44-0	Fluoranthene	7.83		0.50	0.054
86-73-7	Fluorene	8.04		2.0	0.50
193-39-5	Indeno[1,2,3-cd]pyrene	3.17		0.20	0.050
90-12-0	1-Methylnaphthalene	7.86		2.0	0.50
91-57-6	2-Methylnaphthalene	7.57		2.0	0.50
91-20-3	Naphthalene	7.44		2.0	0.25
85-01-8	Phenanthrene	7.52		0.50	0.20
129-00-0	Pyrene	7.50		0.50	0.089

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	72		30-130



TestAmerica Laboratories

Semivolatiles 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04023.D  
 Lab Smp Id: LCS 660-136013/2-A  
 Inj Date : 04-APR-2013 20:13  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : LCS 660-136013/2-A  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 13:07 cantins Quant Type: ISTD  
 Cal Date : 04-APR-2013 16:04 Cal File: 1DD04013.D  
 Als bottle: 21 QC Sample: LCS  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	1000.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
								ON-COLUMN ( ug/l)	FINAL ( ug/l)
* 1 Naphthalene-d8	136		6.091	6.090	(1.000)	2684140	40.0000		
* 6 Acenaphthene-d10	164		7.771	7.770	(1.000)	1602441	40.0000		
* 9 Phenanthrene-d10	188		9.028	9.028	(1.000)	2731968	40.0000		
\$ 13 o-Terphenyl	230		9.328	9.339	(1.033)	295609	7.18133	7.2	
* 17 Chrysene-d12	240		11.337	11.349	(1.000)	2730920	40.0000		
* 22 Perylene-d12	264		13.171	13.176	(1.000)	2751843	40.0000		
2 Naphthalene	128		6.114	6.114	(1.004)	496565	7.44300	7.4	
3 2-Methylnaphthalene	142		6.813	6.819	(1.119)	325909	7.56748	7.6	
4 1-Methylnaphthalene	142		6.907	6.913	(1.134)	319826	7.86387	7.9	
5 Acenaphthylene	152		7.636	7.641	(0.983)	512152	7.55139	7.6	
7 Acenaphthene	154		7.795	7.800	(1.003)	309132	7.38412	7.4	
8 Fluorene	166		8.235	8.240	(1.060)	398793	8.04409	8.0	
10 Phenanthrene	178		9.040	9.051	(1.001)	565842	7.51938	7.5	
11 Anthracene	178		9.081	9.092	(1.006)	531932	7.12196	7.1	
12 Carbazole	167		9.222	9.233	(1.021)	508129	7.71291	7.7	
14 Fluoranthene	202		10.021	10.032	(1.110)	606655	7.83416	7.8	
15 Pyrene	202		10.209	10.220	(0.901)	614790	7.49659	7.5	

Compounds	QUANT SIG		CONCENTRATIONS				
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN ( ug/l)	FINAL ( ug/l)
16 Benzo(a)anthracene	228	11.320	11.325	(0.998)	569571	7.21375	7.2
18 Chrysene	228	11.361	11.378	(1.002)	539084	7.28167	7.3
19 Benzo(b)fluoranthene	252	12.618	12.635	(0.958)	402734	5.85866	5.8
20 Benzo(k)fluoranthene	252	12.648	12.682	(0.960)	416183	5.74683	5.7(H)
21 Benzo(a)pyrene	252	13.065	13.094	(0.992)	342407	4.95744	5.0
23 Indeno(1,2,3-cd)pyrene	276	14.739	14.786	(1.119)	233654	3.17256	3.2(MH)
24 Dibenzo(a,h)anthracene	278	14.763	14.827	(1.121)	230609	3.32512	3.3(H)
25 Benzo(g,h,i)perylene	276	15.168	15.238	(1.152)	249229	3.51456	3.5(H)

QC Flag Legend

- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1DD04023.D

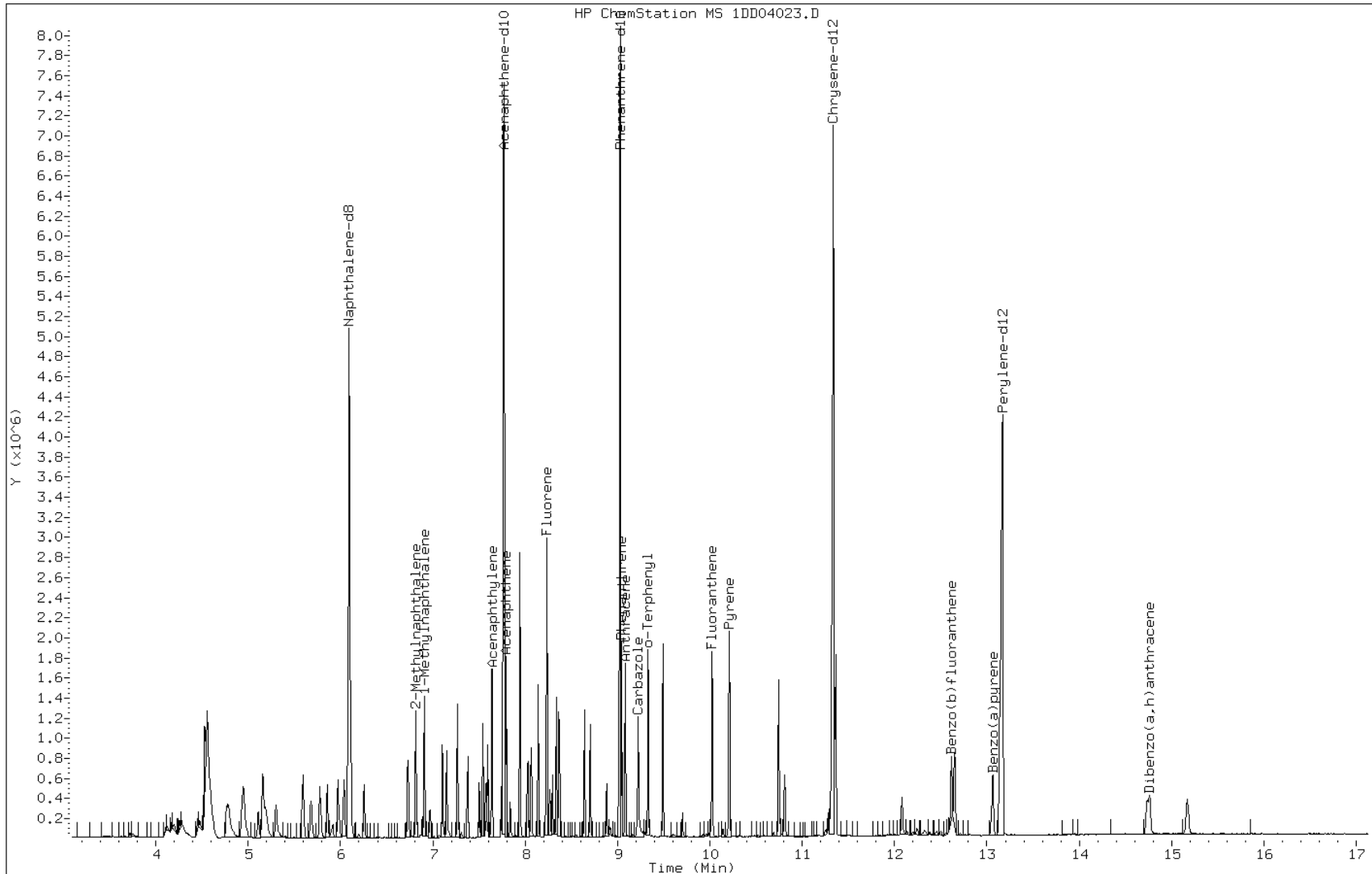
Date: 04-APR-2013 20:13

Client ID:

Instrument: BSMSD.i

Sample Info: LCS 660-136013/2-A

Operator: SCC

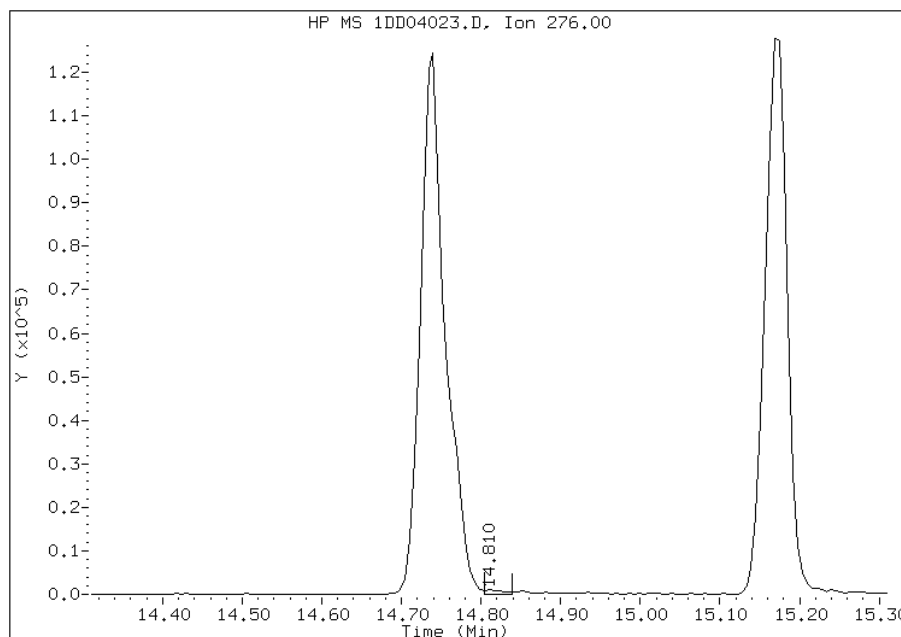


Manual Integration Report

Data File: 1DD04023.D  
Inj. Date and Time: 04-APR-2013 20:13  
Instrument ID: BSMDS.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

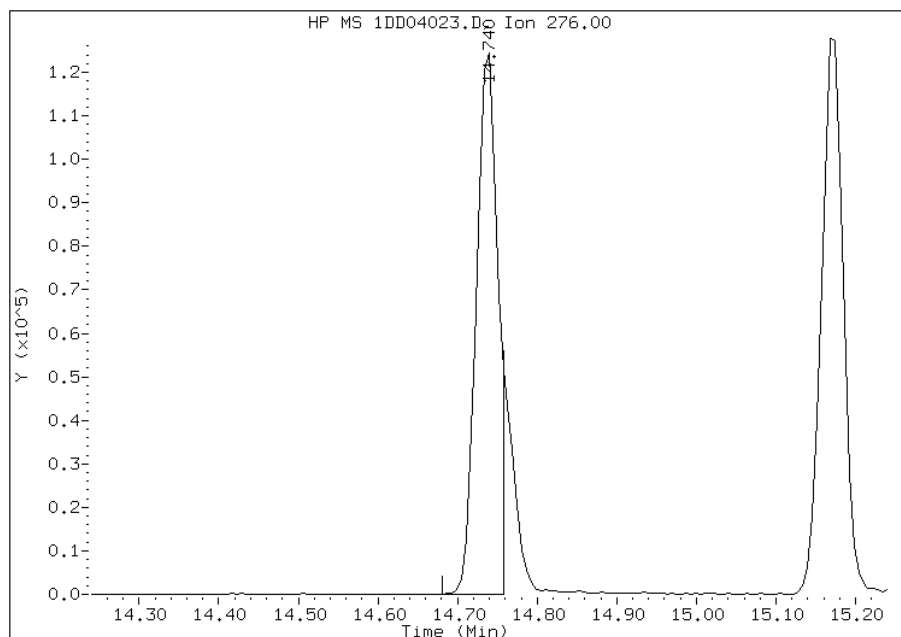
Processing Integration Results

RT: 14.81  
Response: 1999  
Amount: 0  
Conc: 0



Manual Integration Results

RT: 14.74  
Response: 233654  
Amount: 3  
Conc: 3



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 13:13  
Manual Integration Reason: Split Peak

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: \_\_\_\_\_ Lab Sample ID: LCS 660-136063/2-A  
 Matrix: Solid Lab File ID: 1CD03016.D  
 Analysis Method: 8270C LL Date Collected: \_\_\_\_\_  
 Extract. Method: 3546 Date Extracted: 04/02/2013 11:33  
 Sample wt/vol: 14.97(g) Date Analyzed: 04/03/2013 15:52  
 Con. Extract Vol.: 1(mL) Dilution Factor: 1  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136081 Units: ug/Kg

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	458		100	20
208-96-8	Acenaphthylene	488		40	5.0
120-12-7	Anthracene	469		8.4	4.2
56-55-3	Benzo[a]anthracene	484		8.0	3.9
50-32-8	Benzo[a]pyrene	444		10	5.2
205-99-2	Benzo[b]fluoranthene	516		12	6.1
191-24-2	Benzo[g,h,i]perylene	418		20	4.4
207-08-9	Benzo[k]fluoranthene	468		8.0	3.6
218-01-9	Chrysene	462		9.0	4.5
53-70-3	Dibenz(a,h)anthracene	471		20	4.1
206-44-0	Fluoranthene	487		20	4.0
86-73-7	Fluorene	444		20	4.1
193-39-5	Indeno[1,2,3-cd]pyrene	399		20	7.1
90-12-0	1-Methylnaphthalene	522		40	4.4
91-57-6	2-Methylnaphthalene	458		40	7.1
91-20-3	Naphthalene	484		40	4.4
85-01-8	Phenanthrene	499		8.0	3.9
129-00-0	Pyrene	516		20	3.7

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	72		30-130

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03016.D  
 Lab Smp Id: lcs 660-136063/2-a  
 Inj Date : 03-APR-2013 15:52  
 Operator : SCC  
 Smp Info : lcs 660-136063/2-a  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\a-bFASTPAHi-m.m  
 Meth Date : 03-Apr-2013 11:59 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 16 QC Sample: LCS  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula:

$$\text{Amt} * \text{DF} * 1/\text{Vi} * \text{Vt}/\text{Ws} * 100/(100 - \text{M}) * \text{A} * \text{B} * \text{C} * \text{D} * \text{GPC} * \text{CpndVariable}$$

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Ws	14.970	Weight Extracted
M	0.00000	% Moisture
A	1000.000	uL to mL conversion
B	1000.000	g to kg conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1 if no conv)
GPC	1.000	GPC FACTOR
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
							ON-COLUMN	FINAL
	MASS						(ug/ml)	(ug/Kg)
* 1 Naphthalene-d8	136		3.710	3.704	(1.000)	631468	40.0000	
* 6 Acenaphthene-d10	164		4.792	4.792	(1.000)	487717	40.0000	
* 10 Phenanthrene-d10	188		5.739	5.739	(1.000)	877192	40.0000	
\$ 14 o-Terphenyl	230		5.992	5.992	(1.044)	92529	7.18786	480.1508
* 18 Chrysene-d12	240		7.680	7.680	(1.000)	995572	40.0000	
* 23 Perylene-d12	264		8.851	8.851	(1.000)	964341	40.0000	
2 Naphthalene	128		3.722	3.722	(1.003)	117544	7.24725	484.1181
3 2-Methylnaphthalene	142		4.145	4.145	(1.117)	75697	6.85623	457.9979
4 1-Methylnaphthalene	142		4.210	4.210	(1.135)	77643	7.81557	522.0820
5 Acenaphthylene	152		4.710	4.704	(0.983)	147518	7.30815	488.1862
7 Acenaphthene	154		4.816	4.816	(1.005)	85784	6.86151	458.3506
9 Fluorene	166		5.133	5.133	(1.071)	110785	6.64710	444.0278
11 Phenanthrene	178		5.757	5.757	(1.003)	190897	7.47212	499.1395
12 Anthracene	178		5.792	5.792	(1.009)	181998	7.02747	469.4369

Compounds	QUANT SIG		CONCENTRATIONS				
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/ml)	FINAL (ug/Kg)
-----	----	----	-----	-----	-----	-----	-----
13 Carbazole	167	5.898	5.898	(1.028)	177248	7.98844	533.6301
15 Fluoranthene	202	6.592	6.592	(1.149)	205854	7.29605	487.3779
16 Pyrene	202	6.757	6.757	(0.880)	212934	7.72112	515.7731
17 Benzo(a)anthracene	228	7.668	7.668	(0.998)	204915	7.23923	483.5827
19 Chrysene	228	7.698	7.698	(1.002)	196071	6.91134	461.6795
20 Benzo(b)fluoranthene	252	8.509	8.509	(0.961)	210472	7.72013	515.7070
21 Benzo(k)fluoranthene	252	8.527	8.533	(0.963)	184760	7.00698	468.0684
22 Benzo(a)pyrene	252	8.798	8.798	(0.994)	170578	6.64575	443.9378
24 Indeno(1,2,3-cd)pyrene	276	9.992	9.992	(1.129)	145625	5.97338	399.0232(M)
25 Dibenzo(a,h)anthracene	278	10.009	10.009	(1.131)	158866	7.05430	471.2290
26 Benzo(g,h,i)perylene	276	10.339	10.339	(1.168)	155744	6.25938	418.1284

QC Flag Legend

M - Compound response manually integrated.

Data File: 1CD03016.D

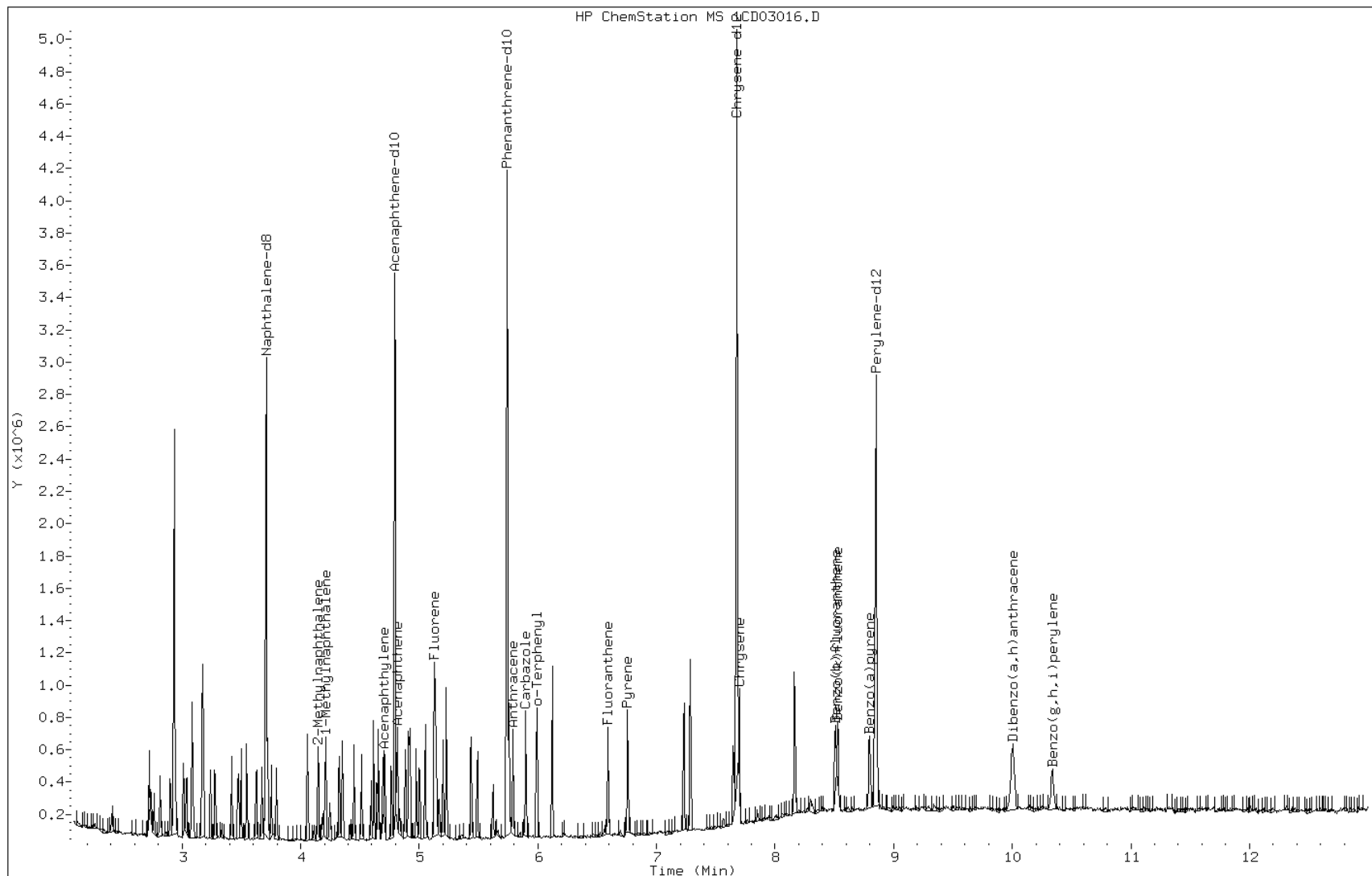
Date: 03-APR-2013 15:52

Client ID:

Instrument: BSMC5973.i

Sample Info: lcs 660-136063/2-a

Operator: SCC



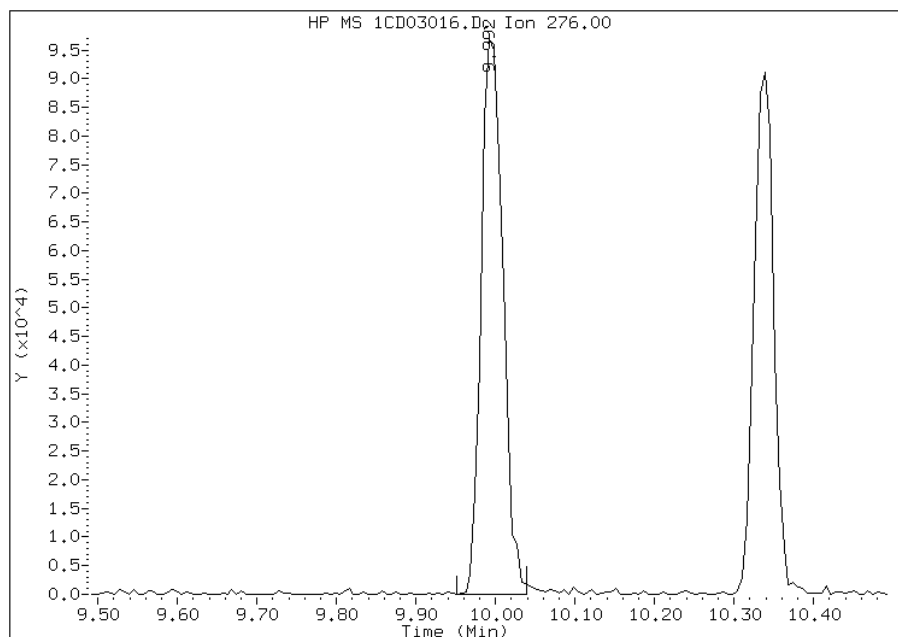


# Manual Integration Report

Data File: 1CD03016.D  
Inj. Date and Time: 03-APR-2013 15:52  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/04/2013

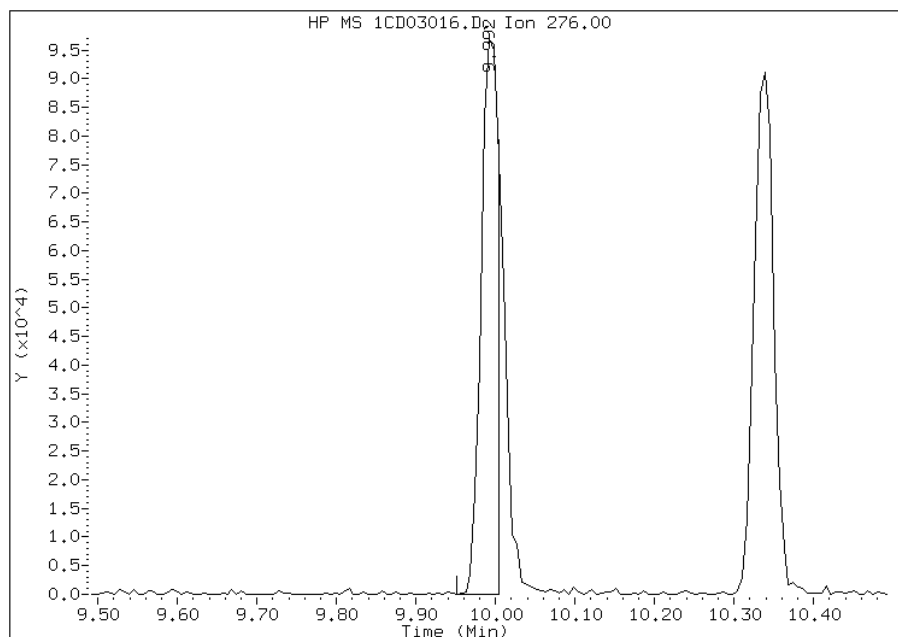
## Processing Integration Results

RT: 9.99  
Response: 182593  
Amount: 7  
Conc: 500



## Manual Integration Results

RT: 9.99  
Response: 145625  
Amount: 6  
Conc: 399



Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:32  
Manual Integration Reason: Split Peak

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: \_\_\_\_\_ Lab Sample ID: LCSD 660-136013/3-A  
 Matrix: Water Lab File ID: 1DD04024.D  
 Analysis Method: 8270C LL Date Collected: \_\_\_\_\_  
 Extract. Method: 3520C Date Extracted: 04/02/2013 08:08  
 Sample wt/vol: 1000(mL) Date Analyzed: 04/04/2013 20:36  
 Con. Extract Vol.: 1(mL) Dilution Factor: 1  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136164 Units: ug/L

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	7.35		2.0	0.50
208-96-8	Acenaphthylene	7.42		1.0	0.25
120-12-7	Anthracene	7.14		0.20	0.076
56-55-3	Benzo[a]anthracene	7.19		0.20	0.050
50-32-8	Benzo[a]pyrene	5.01		0.20	0.057
205-99-2	Benzo[b]fluoranthene	5.92		0.20	0.050
191-24-2	Benzo[g,h,i]perylene	3.91		0.50	0.10
207-08-9	Benzo[k]fluoranthene	5.65		0.20	0.057
218-01-9	Chrysene	6.91		0.20	0.069
53-70-3	Dibenz(a,h)anthracene	3.54		0.20	0.050
206-44-0	Fluoranthene	7.94		0.50	0.054
86-73-7	Fluorene	7.90		2.0	0.50
193-39-5	Indeno[1,2,3-cd]pyrene	3.59		0.20	0.050
90-12-0	1-Methylnaphthalene	7.77		2.0	0.50
91-57-6	2-Methylnaphthalene	7.56		2.0	0.50
91-20-3	Naphthalene	7.36		2.0	0.25
85-01-8	Phenanthrene	7.41		0.50	0.20
129-00-0	Pyrene	7.46		0.50	0.089

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	72		30-130

TestAmerica Laboratories

Semivolatiles 8270 low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\1DD04024.D  
 Lab Smp Id: LCSD 660-136013/3-A  
 Inj Date : 04-APR-2013 20:36  
 Operator : SCC Inst ID: BSMSD.i  
 Smp Info : LCSD 660-136013/3-A  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMSD.i\1D040413.b\dfASTPAHi.m  
 Meth Date : 05-Apr-2013 13:07 cantins Quant Type: ISTD  
 Cal Date : 04-APR-2013 16:04 Cal File: 1DD04013.D  
 Als bottle: 22 QC Sample: LCSD  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	1000.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
								ON-COLUMN ( ug/l)	FINAL ( ug/l)
* 1 Naphthalene-d8			136	6.090	6.090	(1.000)	2700850	40.0000	
* 6 Acenaphthene-d10			164	7.770	7.770	(1.000)	1619728	40.0000	
* 9 Phenanthrene-d10			188	9.028	9.028	(1.000)	2719610	40.0000	
\$ 13 o-Terphenyl			230	9.327	9.339	(1.033)	295542	7.21232	7.2
* 17 Chrysene-d12			240	11.337	11.349	(1.000)	2765140	40.0000	
* 22 Perylene-d12			264	13.170	13.176	(1.000)	2782071	40.0000	
2 Naphthalene			128	6.113	6.114	(1.004)	494316	7.36345	7.4
3 2-Methylnaphthalene			142	6.819	6.819	(1.120)	327714	7.56231	7.6
4 1-Methylnaphthalene			142	6.907	6.913	(1.134)	317804	7.76581	7.8
5 Acenaphthylene			152	7.641	7.641	(0.983)	508668	7.41997	7.4
7 Acenaphthene			154	7.794	7.800	(1.003)	310956	7.34842	7.3
8 Fluorene			166	8.235	8.240	(1.060)	396073	7.90396	7.9
10 Phenanthrene			178	9.039	9.051	(1.001)	555014	7.40900	7.4
11 Anthracene			178	9.081	9.092	(1.006)	530832	7.13953	7.1
12 Carbazole			167	9.222	9.233	(1.021)	507939	7.74506	7.7
14 Fluoranthene			202	10.021	10.032	(1.110)	611727	7.93556	7.9
15 Pyrene			202	10.209	10.220	(0.900)	619595	7.46168	7.5

Compounds	QUANT SIG		CONCENTRATIONS				
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN ( ug/l)	FINAL ( ug/l)
16 Benzo(a)anthracene	228	11.319	11.325	(0.998)	574573	7.18704	7.2
18 Chrysene	228	11.360	11.378	(1.002)	518298	6.91426	6.9
19 Benzo(b)fluoranthene	252	12.612	12.635	(0.958)	411516	5.92137	5.9
20 Benzo(k)fluoranthene	252	12.647	12.682	(0.960)	413704	5.65053	5.6
21 Benzo(a)pyrene	252	13.064	13.094	(0.992)	349884	5.01065	5.0
23 Indeno(1,2,3-cd)pyrene	276	14.733	14.786	(1.119)	267607	3.59410	3.6(MH)
24 Dibenzo(a,h)anthracene	278	14.762	14.827	(1.121)	248551	3.54488	3.5(H)
25 Benzo(g,h,i)perylene	276	15.173	15.238	(1.152)	280575	3.91361	3.9(H)

QC Flag Legend

- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.

Data File: 1DD04024.D

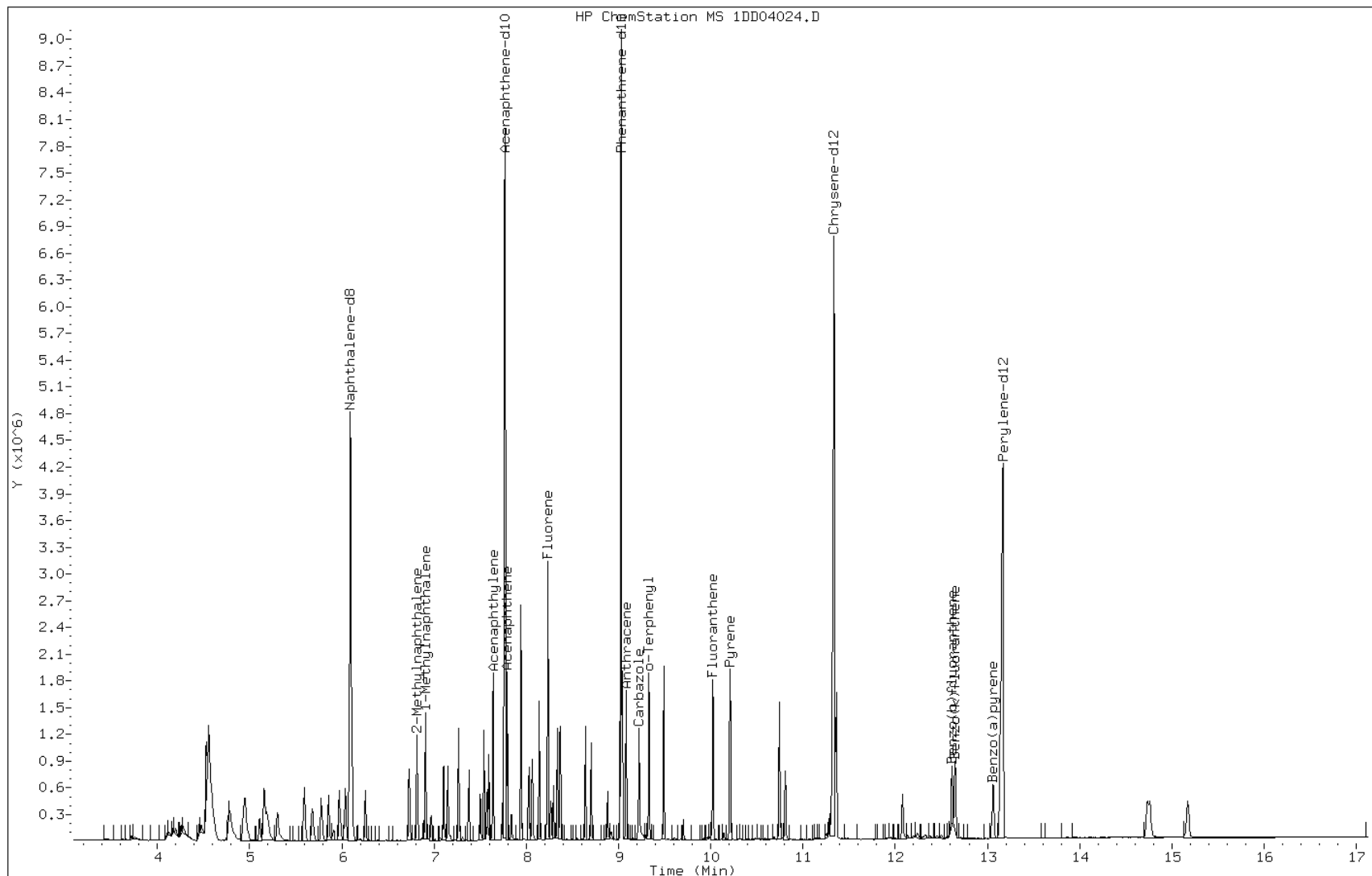
Date: 04-APR-2013 20:36

Client ID:

Instrument: BSMSD.i

Sample Info: LCSD 660-136013/3-A

Operator: SCC

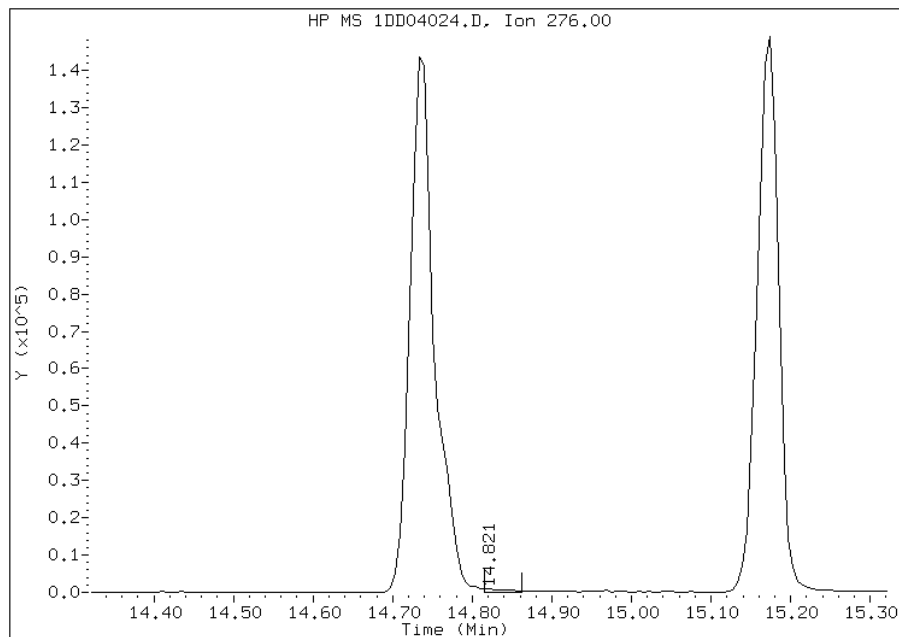


Manual Integration Report

Data File: 1DD04024.D  
Inj. Date and Time: 04-APR-2013 20:36  
Instrument ID: BSMDS.i  
Client ID:  
Compound: 23 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

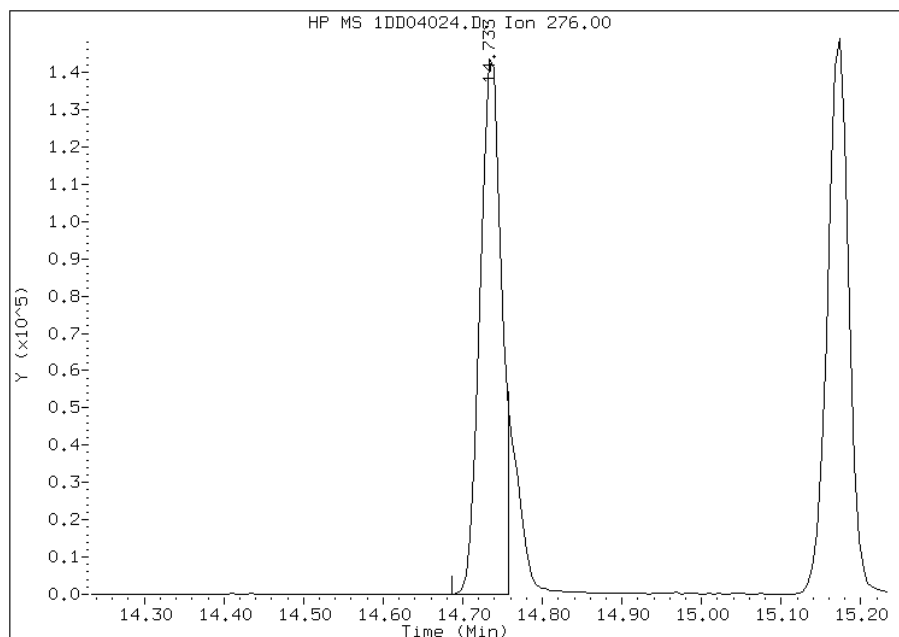
Processing Integration Results

RT: 14.82  
Response: 1981  
Amount: 0  
Conc: 0



Manual Integration Results

RT: 14.73  
Response: 267607  
Amount: 4  
Conc: 4



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 13:14  
Manual Integration Reason: Split Peak

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: CV0014AB-GS MS Lab Sample ID: 680-88766-21 MS  
 Matrix: Solid Lab File ID: 1CD03018.D  
 Analysis Method: 8270C LL Date Collected: 03/25/2013 15:18  
 Extract. Method: 3546 Date Extracted: 04/02/2013 11:33  
 Sample wt/vol: 15.32 (g) Date Analyzed: 04/03/2013 16:29  
 Con. Extract Vol.: 1 (mL) Dilution Factor: 4  
 Injection Volume: 1 (uL) Level: (low/med) Low  
 % Moisture: 26.1 GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136081 Units: ug/Kg

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	656		530	110
208-96-8	Acenaphthylene	626		210	27
120-12-7	Anthracene	666		45	22
56-55-3	Benzo[a]anthracene	833		42	21
50-32-8	Benzo[a]pyrene	650		55	28
205-99-2	Benzo[b]fluoranthene	738		65	32
191-24-2	Benzo[g,h,i]perylene	623		110	23
207-08-9	Benzo[k]fluoranthene	667		42	19
218-01-9	Chrysene	780		48	24
53-70-3	Dibenz(a,h)anthracene	622		110	22
206-44-0	Fluoranthene	697		110	21
86-73-7	Fluorene	639		110	22
193-39-5	Indeno[1,2,3-cd]pyrene	532		110	38
90-12-0	1-Methylnaphthalene	653		210	23
91-57-6	2-Methylnaphthalene	653		210	38
91-20-3	Naphthalene	681		210	23
85-01-8	Phenanthrene	690		42	21
129-00-0	Pyrene	750		110	20

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	92		30-130

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03018.D  
 Lab Smp Id: 680-88766-a-21-e ms  
 Inj Date : 03-APR-2013 16:29  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : 680-88766-a-21-e ms  
 Misc Info : 4.0  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\a-bFASTPAHi-m.m  
 Meth Date : 03-Apr-2013 11:59 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 18 QC Sample: MS  
 Dil Factor: 4.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula:

$$\text{Amt} * \text{DF} * 1/\text{Vi} * \text{Vt}/\text{Ws} * 100/(100 - \text{M}) * \text{A} * \text{B} * \text{C} * \text{D} * \text{GPC} * \text{CpndVariable}$$

Name	Value	Description
DF	4.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Ws	15.320	Weight Extracted
M	0.00000	% Moisture
A	1000.000	uL to mL conversion
B	1000.000	g to kg conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1 if no conv)
GPC	1.000	GPC FACTOR
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
								ON-COLUMN (ug/ml)	FINAL (ug/Kg)
* 1 Naphthalene-d8	136		3.710	3.704	(1.000)	604618	40.0000		
* 6 Acenaphthene-d10	164		4.792	4.792	(1.000)	447317	40.0000		
* 10 Phenanthrene-d10	188		5.739	5.739	(1.000)	819020	40.0000		
\$ 14 o-Terphenyl	230		5.992	5.992	(1.044)	20991	2.29558	599.3677	
* 18 Chrysene-d12	240		7.680	7.680	(1.000)	882360	40.0000		
* 23 Perylene-d12	264		8.851	8.851	(1.000)	885344	40.0000		
2 Naphthalene	128		3.721	3.722	(1.003)	29924	1.92692	503.1110	
3 2-Methylnaphthalene	142		4.145	4.145	(1.117)	19529	1.84738	482.3454	
4 1-Methylnaphthalene	142		4.210	4.210	(1.135)	17586	1.84882	482.7217	
5 Acenaphthylene	152		4.704	4.704	(0.982)	32783	1.77078	462.3434	
7 Acenaphthene	154		4.815	4.816	(1.005)	21291	1.85679	484.8005	
9 Fluorene	166		5.133	5.133	(1.071)	27627	1.80733	471.8875	
11 Phenanthrene	178		5.757	5.757	(1.003)	46547	1.95136	509.4924	
12 Anthracene	178		5.792	5.792	(1.009)	45559	1.88411	491.9353	



Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/ml)	FINAL (ug/Kg)
13 Carbazole	167	5.898	5.898	(1.028)	41473	2.00192	522.6941
15 Fluoranthene	202	6.592	6.592	(1.149)	51944	1.97180	514.8315
16 Pyrene	202	6.757	6.757	(0.880)	51906	2.12364	554.4739
17 Benzo(a)anthracene	228	7.668	7.668	(0.998)	56786	2.35675	615.3405
19 Chrysene	228	7.698	7.698	(1.002)	55519	2.20809	576.5258
20 Benzo(b)fluoranthene	252	8.509	8.509	(0.961)	52279	2.08870	545.3529
21 Benzo(k)fluoranthene	252	8.527	8.533	(0.963)	45683	1.88711	492.7170
22 Benzo(a)pyrene	252	8.792	8.798	(0.993)	43357	1.83992	480.3966
24 Indeno(1,2,3-cd)pyrene	276	9.992	9.992	(1.129)	33696	1.50550	393.0811(M)
25 Dibenzo(a,h)anthracene	278	10.009	10.009	(1.131)	36413	1.76116	459.8320(M)
26 Benzo(g,h,i)perylene	276	10.339	10.339	(1.168)	40267	1.76274	460.2452(M)

QC Flag Legend

M - Compound response manually integrated.

Data File: 1CD03018.D

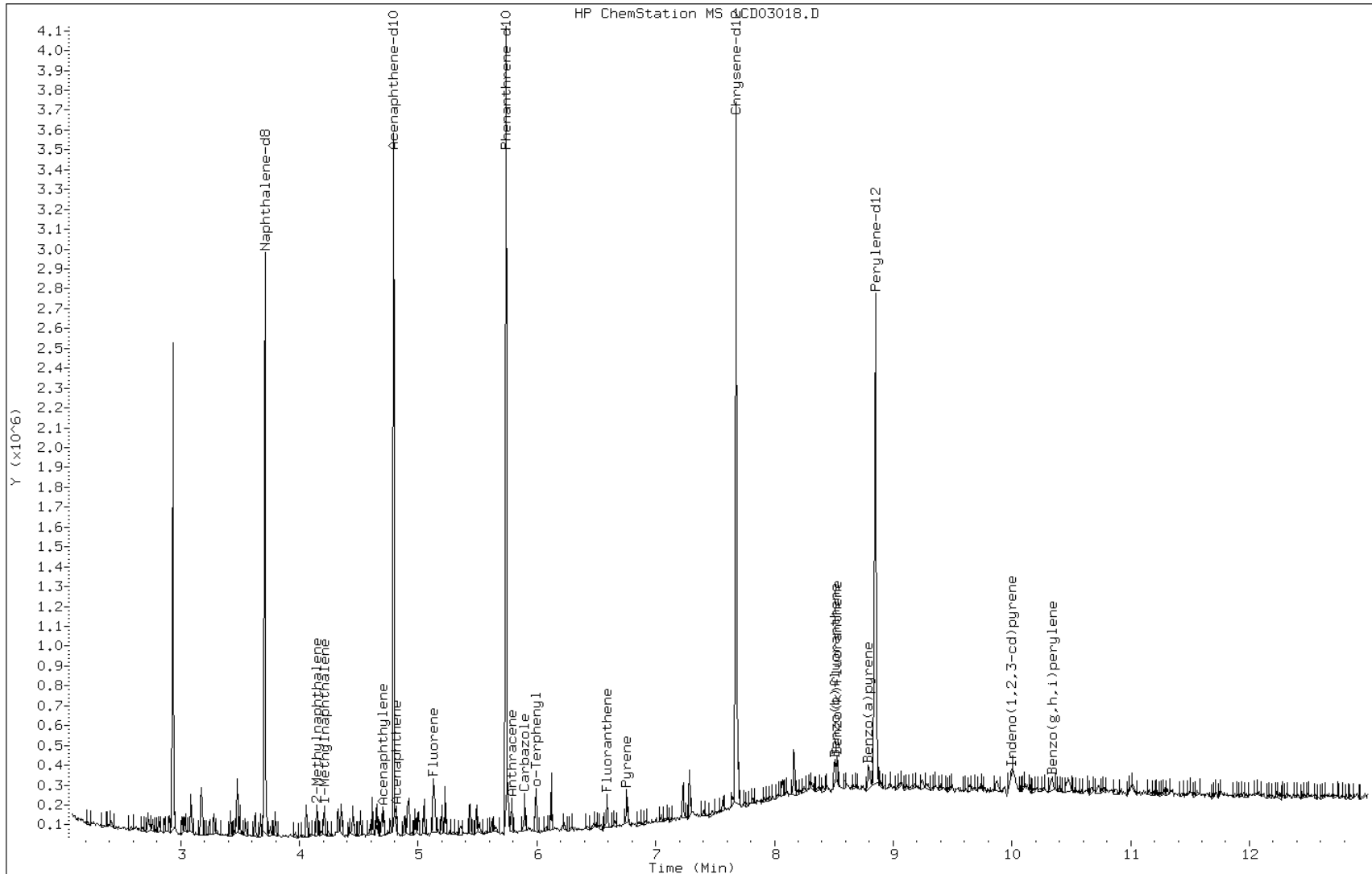
Date: 03-APR-2013 16:29

Client ID:

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-e ms

Operator: SCC

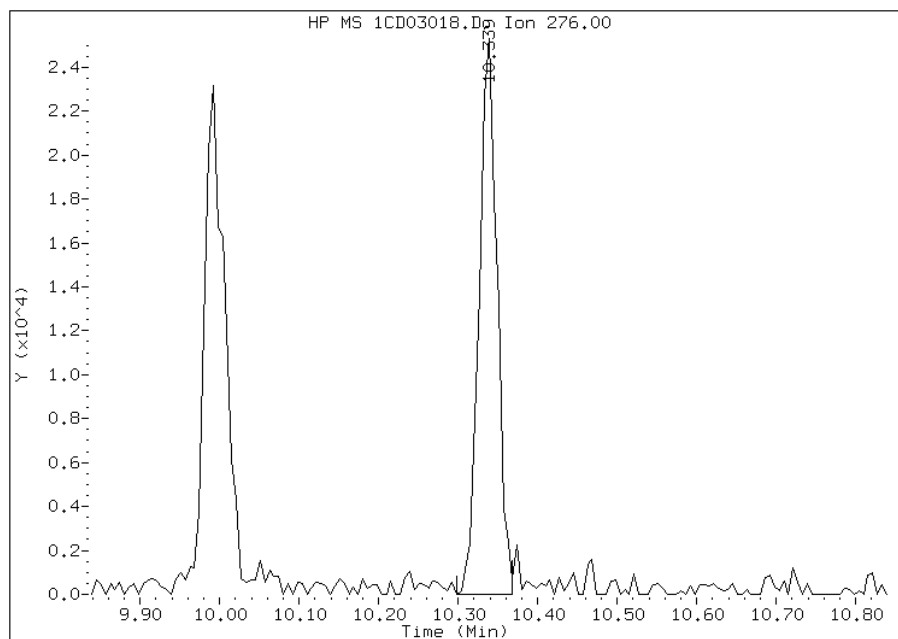


# Manual Integration Report

Data File: 1CD03018.D  
Inj. Date and Time: 03-APR-2013 16:29  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 26 Benzo(g,h,i)perylene  
CAS #: 191-24-2  
Report Date: 04/04/2013

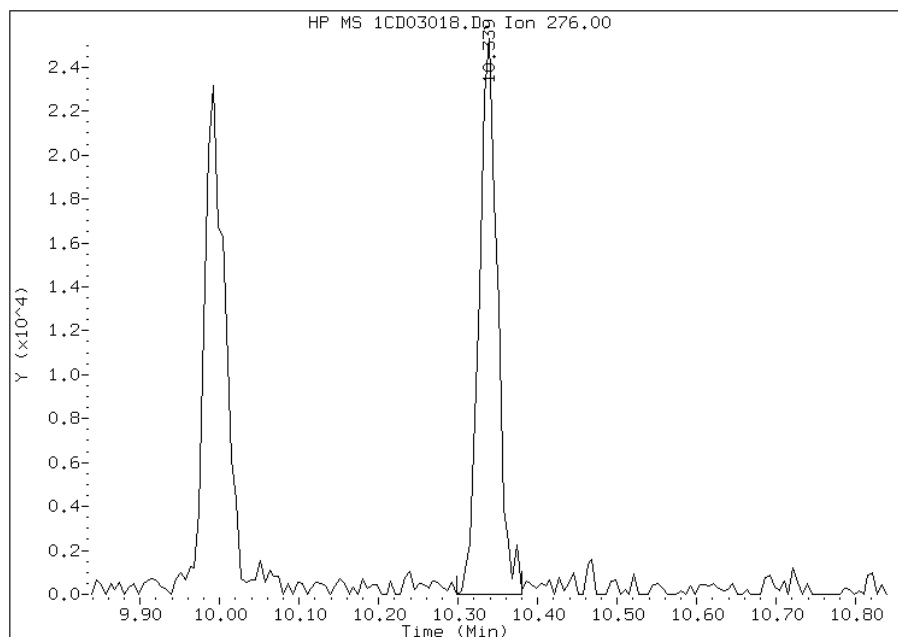
## Processing Integration Results

RT: 10.34  
Response: 39429  
Amount: 2  
Conc: 451



## Manual Integration Results

RT: 10.34  
Response: 40267  
Amount: 2  
Conc: 460



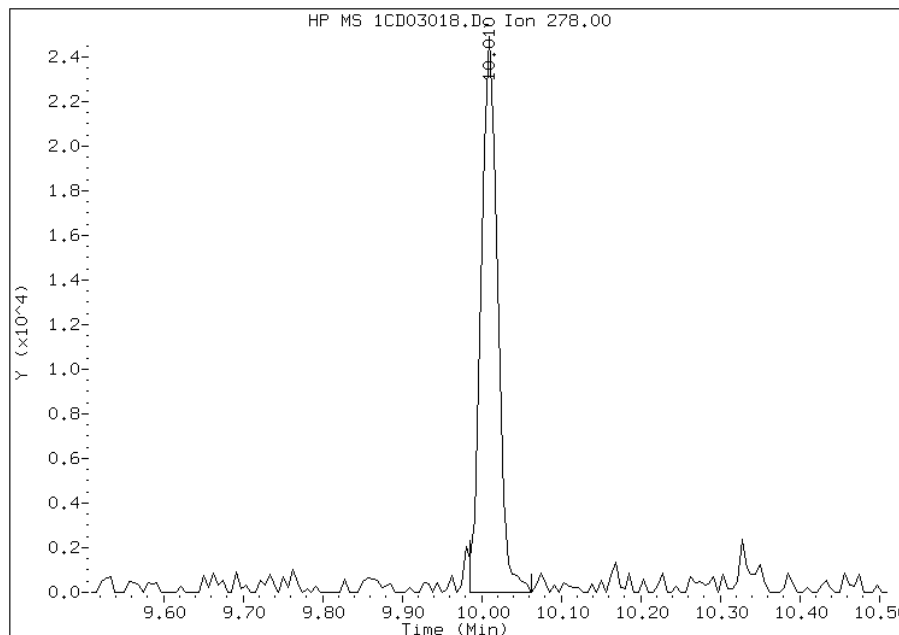
Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:34  
Manual Integration Reason: Baseline Event

# Manual Integration Report

Data File: 1CD03018.D  
Inj. Date and Time: 03-APR-2013 16:29  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 25 Dibenzo(a,h)anthracene  
CAS #: 53-70-3  
Report Date: 04/04/2013

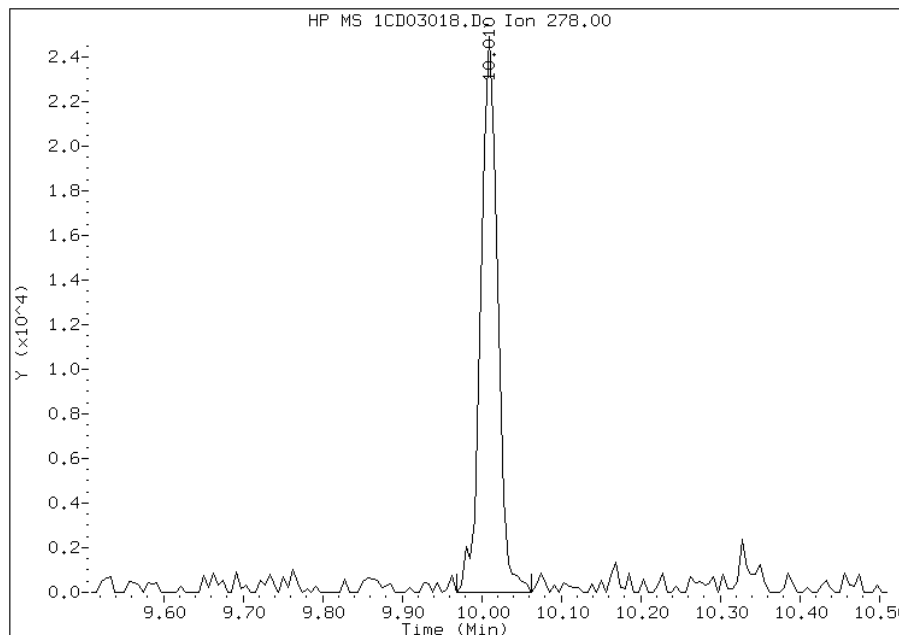
## Processing Integration Results

RT: 10.01  
Response: 35621  
Amount: 2  
Conc: 450



## Manual Integration Results

RT: 10.01  
Response: 36413  
Amount: 2  
Conc: 460



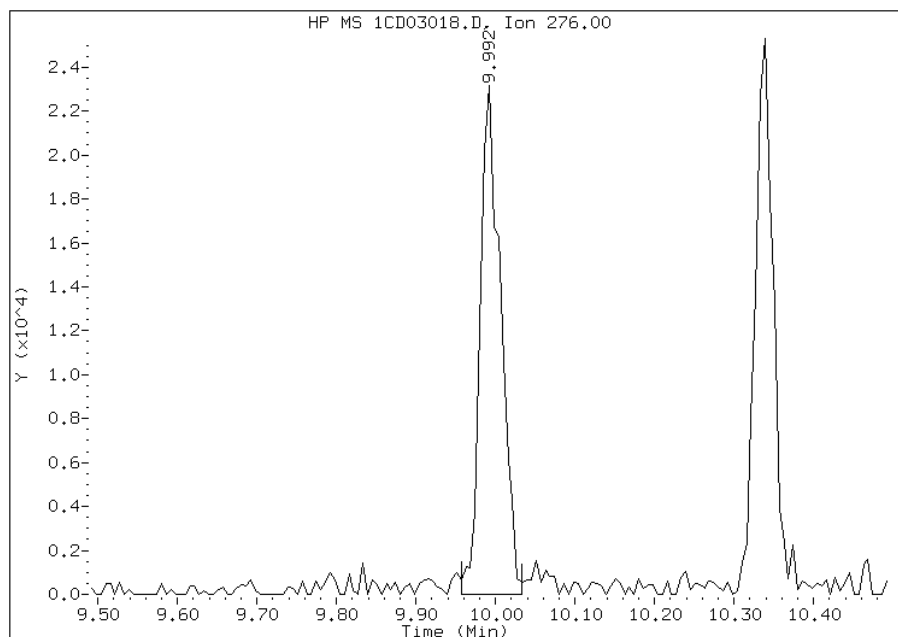
Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:34  
Manual Integration Reason: Baseline Event

# Manual Integration Report

Data File: 1CD03018.D  
Inj. Date and Time: 03-APR-2013 16:29  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/04/2013

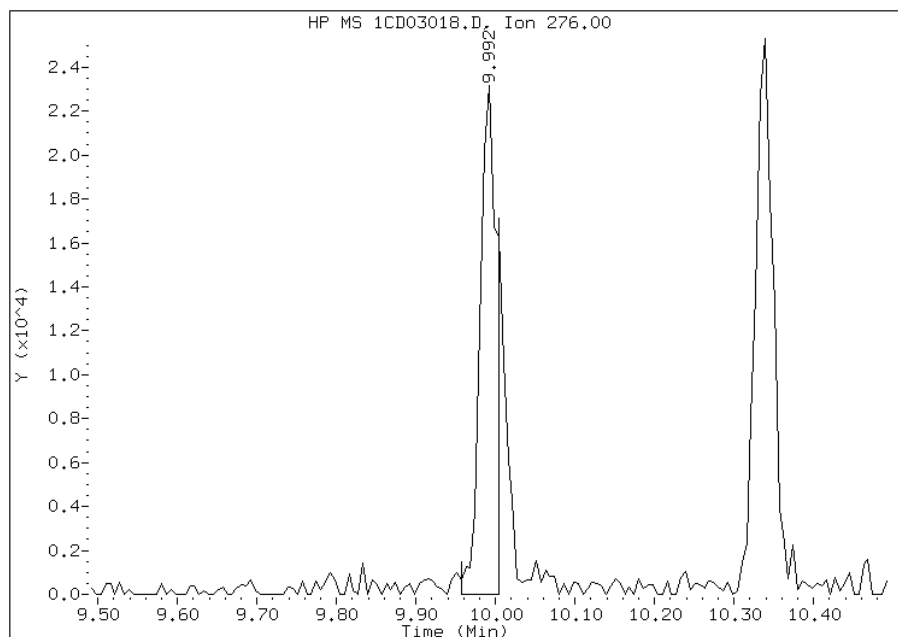
## Processing Integration Results

RT: 9.99  
Response: 41575  
Amount: 2  
Conc: 485



## Manual Integration Results

RT: 9.99  
Response: 33696  
Amount: 2  
Conc: 393



Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:34  
Manual Integration Reason: Split Peak

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: 032613-RB-shovel MS Lab Sample ID: 680-88766-23 MS  
 Matrix: Water Lab File ID: 1CD05007.D  
 Analysis Method: 8270C LL Date Collected: 03/26/2013 13:00  
 Extract. Method: 3520C Date Extracted: 04/02/2013 08:08  
 Sample wt/vol: 960 (mL) Date Analyzed: 04/05/2013 13:12  
 Con. Extract Vol.: 1 (mL) Dilution Factor: 1  
 Injection Volume: 1 (uL) Level: (low/med) Low  
 % Moisture: \_\_\_\_\_ GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136171 Units: ug/L

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	4.74		2.1	0.52
208-96-8	Acenaphthylene	5.18		1.0	0.26
120-12-7	Anthracene	4.97		0.21	0.079
56-55-3	Benzo[a]anthracene	4.72		0.21	0.052
50-32-8	Benzo[a]pyrene	2.13		0.21	0.059
205-99-2	Benzo[b]fluoranthene	2.68		0.21	0.052
191-24-2	Benzo[g,h,i]perylene	1.33		0.52	0.10
207-08-9	Benzo[k]fluoranthene	2.38		0.21	0.059
218-01-9	Chrysene	3.55		0.21	0.072
53-70-3	Dibenz(a,h)anthracene	1.12		0.21	0.052
206-44-0	Fluoranthene	5.46		0.52	0.056
86-73-7	Fluorene	4.83		2.1	0.52
193-39-5	Indeno[1,2,3-cd]pyrene	1.28		0.21	0.052
90-12-0	1-Methylnaphthalene	5.64		2.1	0.52
91-57-6	2-Methylnaphthalene	4.68		2.1	0.52
91-20-3	Naphthalene	5.39		2.1	0.26
85-01-8	Phenanthrene	5.42		0.52	0.21
129-00-0	Pyrene	5.24		0.52	0.093

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	49		30-130

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\1CD05007.D  
 Lab Smp Id: 680-88766-B-23-B MS  
 Inj Date : 05-APR-2013 13:12  
 Operator : SCC  
 Smp Info : 680-88766-B-23-B MS  
 Misc Info :  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040513.b\1CD05007.D  
 Meth Date : 05-Apr-2013 12:31 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 6 QC Sample: MS  
 Dil Factor: 1.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula: Amt \* DF \* 1/Vi \* Vt/Vo \* A \* B \* C \* D \* CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Vo	960.000	Sample Volume
A	1000.000	uL to mL conversion
B	1000.000	mL to L conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1= if no con
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
								ON-COLUMN (ug/ml)	FINAL (ug/l)
* 1 Naphthalene-d8	136		3.692	3.692	(1.000)	411549	40.0000		
* 6 Acenaphthene-d10	164		4.774	4.780	(1.000)	313598	40.0000		
* 10 Phenanthrene-d10	188		5.721	5.721	(1.000)	608819	40.0000		
\$ 14 o-Terphenyl	230		5.974	5.974	(1.044)	41438	4.89529	5.0992	
* 18 Chrysene-d12	240		7.657	7.662	(1.000)	752169	40.0000		
* 23 Perylene-d12	264		8.821	8.827	(1.000)	802662	40.0000		
2 Naphthalene	128		3.704	3.704	(1.003)	54689	5.17371	5.3892(R)	
3 2-Methylnaphthalene	142		4.133	4.133	(1.119)	32325	4.49236	4.6795(R)	
4 1-Methylnaphthalene	142		4.192	4.192	(1.135)	35049	5.41332	5.6388	
5 Acenaphthylene	152		4.692	4.692	(0.983)	64565	4.97455	5.1818	
7 Acenaphthene	154		4.798	4.798	(1.005)	36555	4.54731	4.7367(R)	
9 Fluorene	166		5.116	5.116	(1.071)	49688	4.63657	4.8297(R)	
11 Phenanthrene	178		5.739	5.739	(1.003)	92320	5.20651	5.4234(R)	
12 Anthracene	178		5.768	5.774	(1.008)	85715	4.76865	4.9673	
13 Carbazole	167		5.880	5.880	(1.028)	81580	5.29750	5.5182	
15 Fluoranthene	202		6.568	6.574	(1.148)	102690	5.24400	5.4625(R)	
16 Pyrene	202		6.739	6.739	(0.880)	104769	5.02835	5.2378(R)	

Compounds	QUANT SIG		CONCENTRATIONS					
	MASS		RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/ml)	FINAL ( ug/l)
=====	=====		=====	=====	=====	=====	=====	=====
17 Benzo(a)anthracene	228		7.651	7.651	(0.999)	95789	4.53082	4.7196(R)
19 Chrysene	228		7.674	7.680	(1.002)	73051	3.40826	3.5502(R)
20 Benzo(b)fluoranthene	252		8.480	8.486	(0.961)	58382	2.57281	2.6800(R)
21 Benzo(k)fluoranthene	252		8.504	8.509	(0.964)	50112	2.28330	2.3784(R)
22 Benzo(a)pyrene	252		8.768	8.774	(0.994)	43747	2.04770	2.1330(R)
24 Indeno(1,2,3-cd)pyrene	276		9.956	9.962	(1.129)	24986	1.23134	1.2826(RM)
25 Dibenzo(a,h)anthracene	278		9.968	9.980	(1.130)	20244	1.07998	1.1249(R)
26 Benzo(g,h,i)perylene	276		10.286	10.303	(1.166)	26358	1.27271	1.3257(R)

QC Flag Legend

R - Spike/Surrogate failed recovery limits.  
M - Compound response manually integrated.



Data File: 1CD05007.D

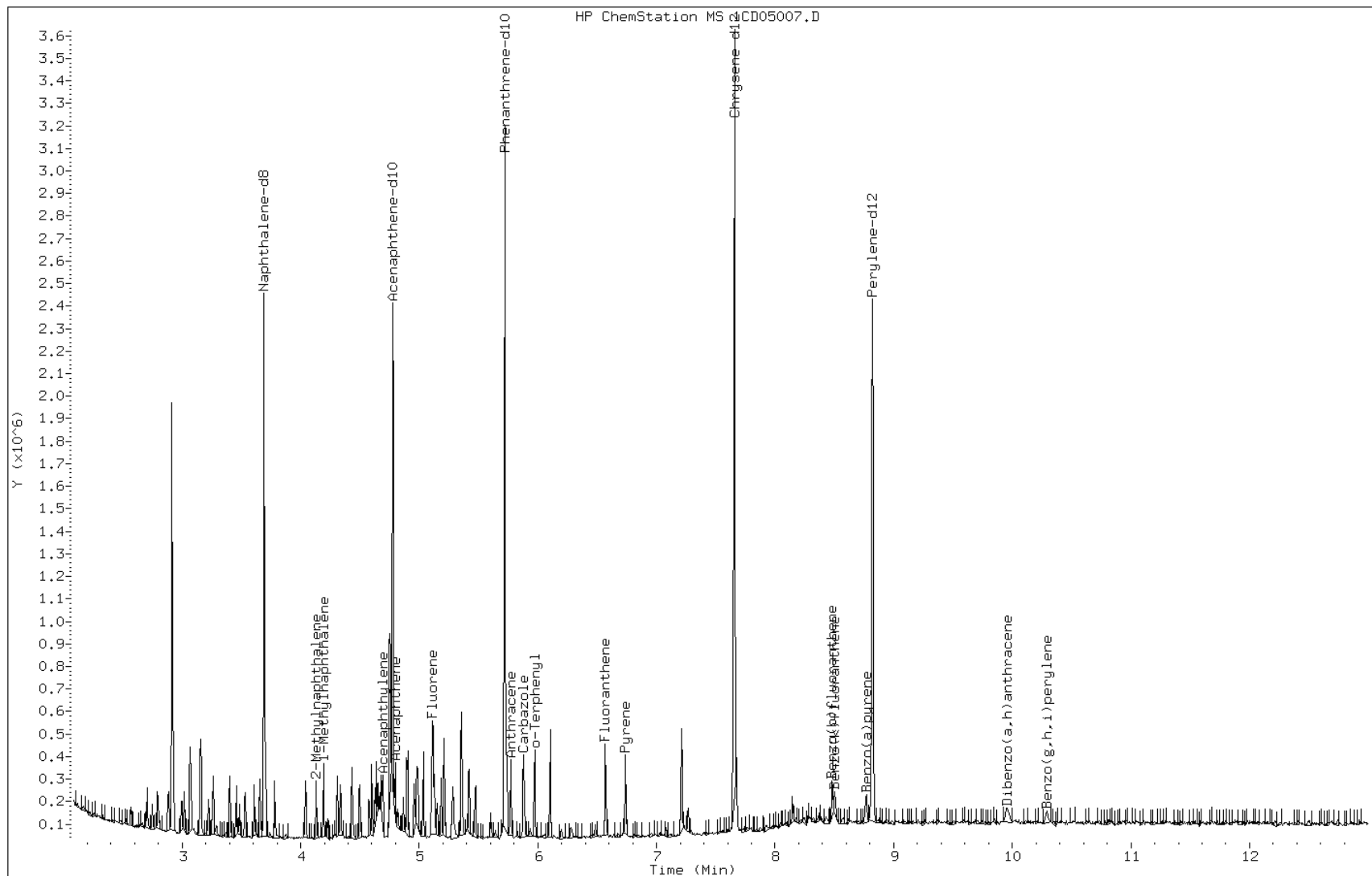
Date: 05-APR-2013 13:12

Client ID:

Instrument: BSMC5973.i

Sample Info: 680-88766-B-23-B MS

Operator: SCC

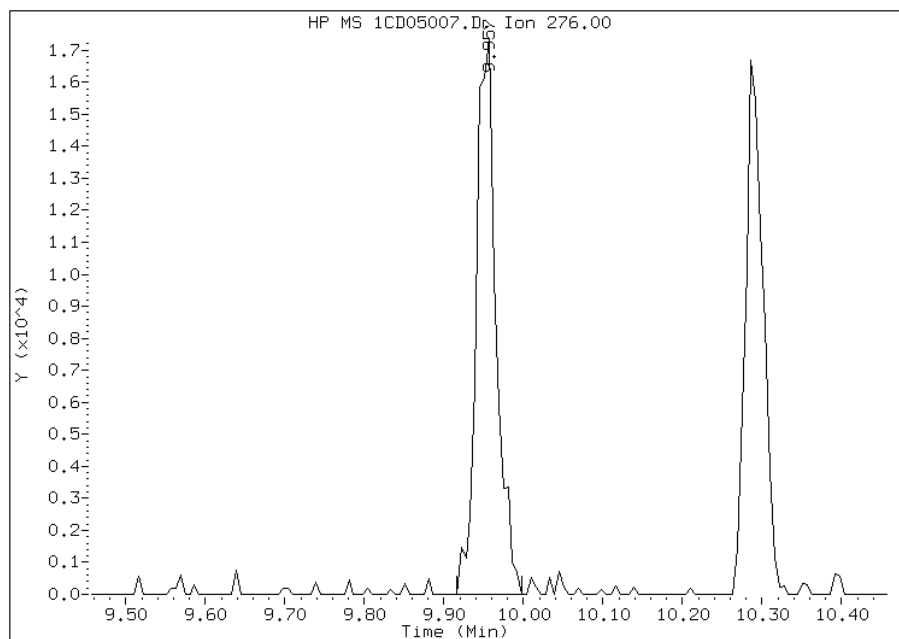


Manual Integration Report

Data File: 1CD05007.D  
Inj. Date and Time: 05-APR-2013 13:12  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/05/2013

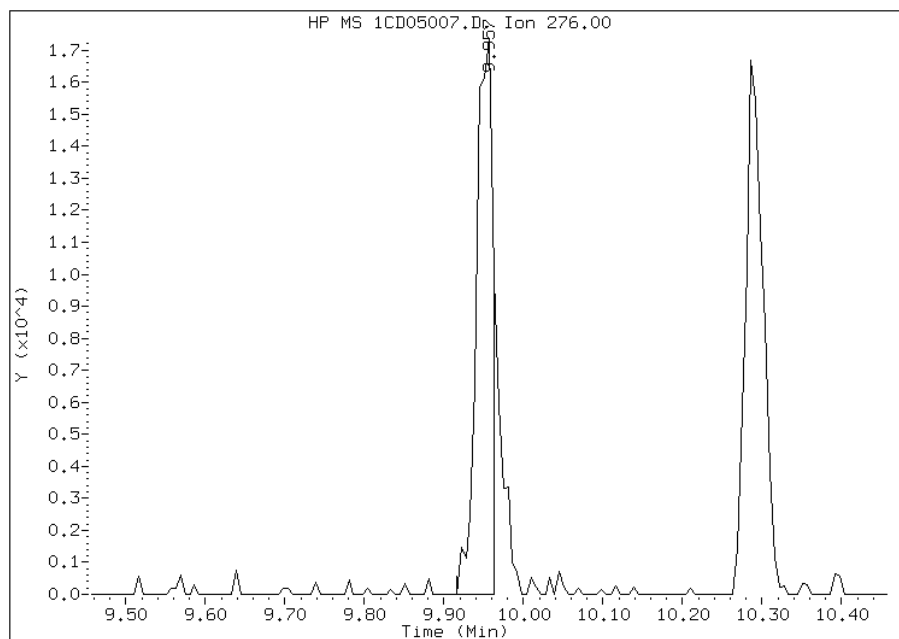
Processing Integration Results

RT: 9.96  
Response: 30035  
Amount: 1  
Conc: 2



Manual Integration Results

RT: 9.96  
Response: 24986  
Amount: 1  
Conc: 1



Manually Integrated By: cantins  
Modification Date: 05-Apr-2013 13:29  
Manual Integration Reason: Split Peak

FORM I  
GC/MS SEMI VOA ORGANICS ANALYSIS DATA SHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2  
 SDG No.: 68088766-2  
 Client Sample ID: CV0014AB-GS MSD Lab Sample ID: 680-88766-21 MSD  
 Matrix: Solid Lab File ID: 1CD03019.D  
 Analysis Method: 8270C LL Date Collected: 03/25/2013 15:18  
 Extract. Method: 3546 Date Extracted: 04/02/2013 11:33  
 Sample wt/vol: 14.60(g) Date Analyzed: 04/03/2013 16:47  
 Con. Extract Vol.: 1(mL) Dilution Factor: 4  
 Injection Volume: 1(uL) Level: (low/med) Low  
 % Moisture: 26.1 GPC Cleanup: (Y/N) N  
 Analysis Batch No.: 136081 Units: ug/Kg

CAS NO.	COMPOUND NAME	RESULT	Q	RL	MDL
83-32-9	Acenaphthene	669		560	110
208-96-8	Acenaphthylene	686		220	28
120-12-7	Anthracene	674		47	23
56-55-3	Benzo[a]anthracene	856		44	22
50-32-8	Benzo[a]pyrene	651		58	29
205-99-2	Benzo[b]fluoranthene	710		68	34
191-24-2	Benzo[g,h,i]perylene	618		110	24
207-08-9	Benzo[k]fluoranthene	822		44	20
218-01-9	Chrysene	739		50	25
53-70-3	Dibenz(a,h)anthracene	713		110	23
206-44-0	Fluoranthene	780		110	22
86-73-7	Fluorene	657		110	23
193-39-5	Indeno[1,2,3-cd]pyrene	659		110	39
90-12-0	1-Methylnaphthalene	709		220	24
91-57-6	2-Methylnaphthalene	671		220	39
91-20-3	Naphthalene	648		220	24
85-01-8	Phenanthrene	758		44	22
129-00-0	Pyrene	878		110	21

CAS NO.	SURROGATE	%REC	Q	LIMITS
84-15-1	o-Terphenyl	97		30-130

TestAmerica Laboratories

Semivolatiles 8270C low level PAH

Data file : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\1CD03019.D  
 Lab Smp Id: 680-88766-a-21-f ms  
 Inj Date : 03-APR-2013 16:47  
 Operator : SCC Inst ID: BSMC5973.i  
 Smp Info : 680-88766-a-21-f msd  
 Misc Info : 4.0  
 Comment :  
 Method : \\tam-chemsvr\chem\SM\BSMC5973.i\1C040313.b\a-bFASTPAHi-m.m  
 Meth Date : 03-Apr-2013 11:59 cantins Quant Type: ISTD  
 Cal Date : 02-APR-2013 15:15 Cal File: 1CD02011.D  
 Als bottle: 19 QC Sample: MSD  
 Dil Factor: 4.00000  
 Integrator: HP RTE Compound Sublist: pah.sub  
 Target Version: 4.14  
 Processing Host: TAM1000

Concentration Formula:

$$\text{Amt} * \text{DF} * 1/\text{Vi} * \text{Vt}/\text{Ws} * 100/(100 - \text{M}) * \text{A} * \text{B} * \text{C} * \text{D} * \text{GPC} * \text{CpndVariable}$$

Name	Value	Description
DF	4.000	Dilution Factor
Vi	1.000	Injection Volume
Vt	1.000	Final Volume
Ws	14.600	Weight Extracted
M	0.00000	% Moisture
A	1000.000	uL to mL conversion
B	1000.000	g to kg conversion
C	0.00100	ng to ug conversion
D	1.000	ug to mg conversion(value = 1 if no conv)
GPC	1.000	GPC FACTOR
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
								ON-COLUMN (ug/ml)	FINAL (ug/Kg)
* 1 Naphthalene-d8	136		3.710	3.704	(1.000)	599290	40.0000		
* 6 Acenaphthene-d10	164		4.792	4.792	(1.000)	437990	40.0000		
* 10 Phenanthrene-d10	188		5.739	5.739	(1.000)	801261	40.0000		
\$ 14 o-Terphenyl	230		5.992	5.992	(1.044)	22282	2.42909	665.5045	
* 18 Chrysene-d12	240		7.680	7.680	(1.000)	885431	40.0000		
* 23 Perylene-d12	264		8.850	8.851	(1.000)	855561	40.0000		
2 Naphthalene	128		3.721	3.722	(1.003)	26909	1.74817	478.9516	
3 2-Methylnaphthalene	142		4.145	4.145	(1.117)	18972	1.81065	496.0680	
4 1-Methylnaphthalene	142		4.210	4.210	(1.135)	18036	1.91299	524.1069	
5 Acenaphthylene	152		4.704	4.704	(0.982)	33560	1.85135	507.2185	
7 Acenaphthene	154		4.815	4.816	(1.005)	20252	1.80379	494.1878	
9 Fluorene	166		5.133	5.133	(1.071)	26513	1.77139	485.3117	
11 Phenanthrene	178		5.757	5.757	(1.003)	47684	2.04333	559.8158	
12 Anthracene	178		5.792	5.792	(1.009)	43006	1.81795	498.0687	

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/ml)	FINAL (ug/Kg)
-----	----	----	-----	-----	-----	-----	-----
13 Carbazole	167	5.898	5.898	(1.028)	40592	2.00282	548.7177
15 Fluoranthene	202	6.592	6.592	(1.149)	54214	2.10359	576.3252
16 Pyrene	202	6.756	6.757	(0.880)	58049	2.36673	648.4183
17 Benzo(a)anthracene	228	7.668	7.668	(0.998)	55755	2.30886	632.5654
19 Chrysene	228	7.698	7.698	(1.002)	50258	1.99192	545.7319
20 Benzo(b)fluoranthene	252	8.509	8.509	(0.961)	46327	1.91533	524.7488
21 Benzo(k)fluoranthene	252	8.527	8.533	(0.963)	51872	2.21736	607.4953
22 Benzo(a)pyrene	252	8.792	8.798	(0.993)	39958	1.75471	480.7413
24 Indeno(1,2,3-cd)pyrene	276	9.992	9.992	(1.129)	38467	1.77849	487.2581(M)
25 Dibenzo(a,h)anthracene	278	10.009	10.009	(1.131)	38424	1.92311	526.8807
26 Benzo(g,h,i)perylene	276	10.333	10.339	(1.167)	36793	1.66673	456.6381

QC Flag Legend

M - Compound response manually integrated.

Data File: 1CD03019.D

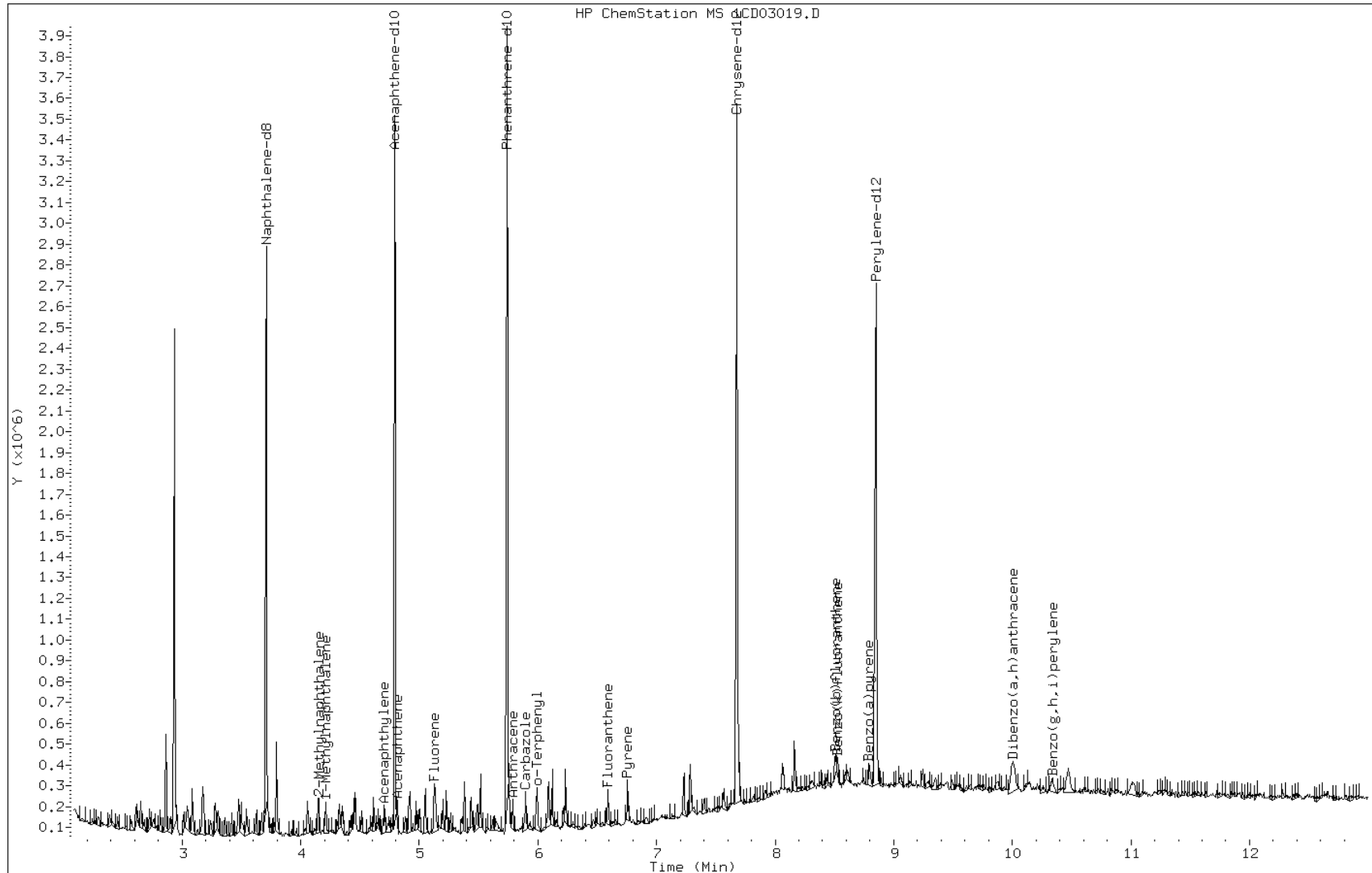
Date: 03-APR-2013 16:47

Client ID:

Instrument: BSMC5973.i

Sample Info: 680-88766-a-21-f msd

Operator: SCC

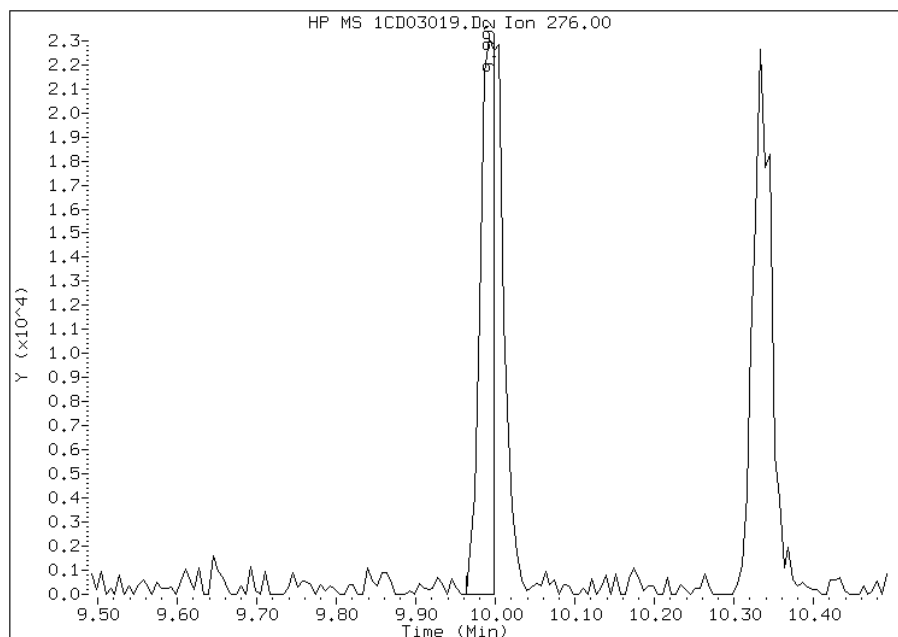


# Manual Integration Report

Data File: 1CD03019.D  
Inj. Date and Time: 03-APR-2013 16:47  
Instrument ID: BSMC5973.i  
Client ID:  
Compound: 24 Indeno(1,2,3-cd)pyrene  
CAS #: 193-39-5  
Report Date: 04/04/2013

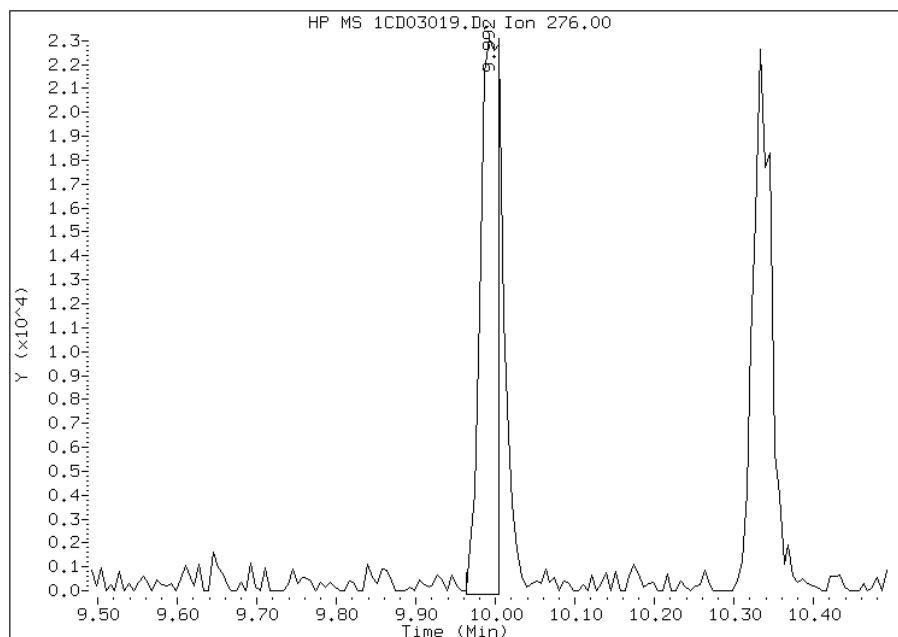
## Processing Integration Results

RT: 9.99  
Response: 30001  
Amount: 1  
Conc: 380



## Manual Integration Results

RT: 9.99  
Response: 38467  
Amount: 2  
Conc: 487



Manually Integrated By: cantins  
Modification Date: 04-Apr-2013 15:35  
Manual Integration Reason: Baseline Event

## GC/MS SEMI VOA ANALYSIS RUN LOG

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMC5973 Start Date: 04/02/2013 10:54Analysis Batch Number: 136048 End Date: 04/02/2013 15:34

LAB SAMPLE ID	CLIENT SAMPLE ID	DATE ANALYZED	DILUTION FACTOR	LAB FILE ID	COLUMN ID
ZZZZZ		04/02/2013 10:54	1		DB-5MS 250 (um)
ZZZZZ		04/02/2013 11:13	1		DB-5MS 250 (um)
DFTPP 660-136048/2		04/02/2013 11:31	1	1CD02002.D	DB-5MS 250 (um)
CCVIS 660-136048/3		04/02/2013 11:49	1		DB-5MS 250 (um)
CCVIS 660-136048/4		04/02/2013 12:09	1		DB-5MS 250 (um)
IC 660-136048/5		04/02/2013 13:26	1	1CD02005.D	DB-5MS 250 (um)
IC 660-136048/6		04/02/2013 13:44	1	1CD02006.D	DB-5MS 250 (um)
IC 660-136048/7		04/02/2013 14:02	1	1CD02007.D	DB-5MS 250 (um)
IC 660-136048/8		04/02/2013 14:20	1	1CD02008.D	DB-5MS 250 (um)
ICIS 660-136048/9		04/02/2013 14:39	1	1CD02009.D	DB-5MS 250 (um)
IC 660-136048/10		04/02/2013 14:57	1	1CD02010.D	DB-5MS 250 (um)
IC 660-136048/11		04/02/2013 15:15	1	1CD02011.D	DB-5MS 250 (um)
ICV 660-136048/12		04/02/2013 15:34	1	1CD02012.D	DB-5MS 250 (um)



## GC/MS SEMI VOA ANALYSIS RUN LOG

Lab Name: TestAmerica TampaJob No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMC5973Start Date: 04/03/2013 10:52Analysis Batch Number: 136081End Date: 04/03/2013 22:35

LAB SAMPLE ID	CLIENT SAMPLE ID	DATE ANALYZED	DILUTION FACTOR	LAB FILE ID	COLUMN ID
ZZZZZ		04/03/2013 10:52	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 11:10	1		DB-5MS 250 (um)
DFTPP 660-136081/2		04/03/2013 11:28	1	1CD03002.D	DB-5MS 250 (um)
CCVIS 660-136081/3		04/03/2013 11:45	1	1CD03003.D	DB-5MS 250 (um)
ZZZZZ		04/03/2013 12:04	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 12:22	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 12:40	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 12:59	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 13:17	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 13:35	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 13:54	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 14:12	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 14:30	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 14:49	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 15:07	4		DB-5MS 250 (um)
MB 660-136063/1-A		04/03/2013 15:34	1	1CD03015.D	DB-5MS 250 (um)
LCS 660-136063/2-A		04/03/2013 15:52	1	1CD03016.D	DB-5MS 250 (um)
680-88766-21	CV0014AB-GS	04/03/2013 16:10	4	1CD03017.D	DB-5MS 250 (um)
680-88766-21 MS	CV0014AB-GS MS	04/03/2013 16:29	4	1CD03018.D	DB-5MS 250 (um)
680-88766-21 MSD	CV0014AB-GS MSD	04/03/2013 16:47	4	1CD03019.D	DB-5MS 250 (um)
ZZZZZ		04/03/2013 17:05	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 17:24	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 17:42	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 18:00	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 18:19	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 18:37	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 18:55	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 19:13	4		DB-5MS 250 (um)
ZZZZZ		04/03/2013 19:32	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 19:50	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 20:08	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 20:27	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 20:45	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 21:03	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 21:21	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 21:40	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 21:58	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 22:17	1		DB-5MS 250 (um)
ZZZZZ		04/03/2013 22:35	1		DB-5MS 250 (um)

## GC/MS SEMI VOA ANALYSIS RUN LOG

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMC5973 Start Date: 04/05/2013 10:58Analysis Batch Number: 136171 End Date: 04/05/2013 13:12

LAB SAMPLE ID	CLIENT SAMPLE ID	DATE ANALYZED	DILUTION FACTOR	LAB FILE ID	COLUMN ID
ZZZZZ		04/05/2013 10:58	1		DB-5MS 250 (um)
ZZZZZ		04/05/2013 11:18	1		DB-5MS 250 (um)
DFTPP 660-136171/2		04/05/2013 11:37	1		DB-5MS 250 (um)
DFTPP 660-136171/3		04/05/2013 11:57	1	1CD05003.D	DB-5MS 250 (um)
CCVIS 660-136171/4		04/05/2013 12:15	1	1CD05004.D	DB-5MS 250 (um)
ZZZZZ		04/05/2013 12:35	1		DB-5MS 250 (um)
680-88766-23	032613-RB-shovel	04/05/2013 12:54	1	1CD05006.D	DB-5MS 250 (um)
680-88766-23 MS	032613-RB-shovel MS	04/05/2013 13:12	1	1CD05007.D	DB-5MS 250 (um)

## GC/MS SEMI VOA ANALYSIS RUN LOG

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Instrument ID: BSMD5973 Start Date: 04/04/2013 11:04Analysis Batch Number: 136164 End Date: 04/04/2013 20:36

LAB SAMPLE ID	CLIENT SAMPLE ID	DATE ANALYZED	DILUTION FACTOR	LAB FILE ID	COLUMN ID
ZZZZZ		04/04/2013 11:04	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 11:30	1		DB-5MS 250 (um)
DFTPP 660-136164/2		04/04/2013 11:55	1		DB-5MS 250 (um)
DFTPP 660-136164/3		04/04/2013 12:15	1	1DD04003.D	DB-5MS 250 (um)
CCVIS 660-136164/4		04/04/2013 12:34	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 13:02	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 13:26	1		DB-5MS 250 (um)
IC 660-136164/15		04/04/2013 13:49	1	1DD04007.D	DB-5MS 250 (um)
IC 660-136164/16		04/04/2013 14:11	1	1DD04008.D	DB-5MS 250 (um)
IC 660-136164/17		04/04/2013 14:34	1	1DD04009.D	DB-5MS 250 (um)
IC 660-136164/18		04/04/2013 14:57	1	1DD04010.D	DB-5MS 250 (um)
ICIS 660-136164/19		04/04/2013 15:19	1	1DD04011.D	DB-5MS 250 (um)
IC 660-136164/20		04/04/2013 15:42	1	1DD04012.D	DB-5MS 250 (um)
IC 660-136164/21		04/04/2013 16:04	1	1DD04013.D	DB-5MS 250 (um)
ICV 660-136164/22		04/04/2013 16:27	1	1DD04014.D	DB-5MS 250 (um)
ZZZZZ		04/04/2013 16:52	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 17:18	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 17:44	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 18:09	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 18:35	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 19:01	1		DB-5MS 250 (um)
ZZZZZ		04/04/2013 19:27	1		DB-5MS 250 (um)
MB 660-136013/1-A		04/04/2013 19:51	1	1DD04022.D	DB-5MS 250 (um)
LCS 660-136013/2-A		04/04/2013 20:13	1	1DD04023.D	DB-5MS 250 (um)
LCSD 660-136013/3-A		04/04/2013 20:36	1	1DD04024.D	DB-5MS 250 (um)

## GC/MS SEMI VOA BATCH WORKSHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Batch Number: 136013 Batch Start Date: 04/02/13 08:08 Batch Analyst: George, AbrahamBatch Method: 3520C Batch End Date: 04/03/13 10:55

Lab Sample ID	Client Sample ID	Method Chain	Basis	InitialAmount	FinalAmount	ReceivedpH	FirstAdjustpH	EX-625LVI SPK 00020	EXLLSURINT 00178
MB 660-136013/1		3520C, 8270C LL		1000 mL	1 mL	8	<2		1 mL
LCS 660-136013/2		3520C, 8270C LL		1000 mL	1 mL	8	<2	1 mL	1 mL
LCSD 660-136013/3		3520C, 8270C LL		1000 mL	1 mL	8	<2	1 mL	1 mL
680-88766-A-23	032613-RB-shovel	3520C, 8270C LL	T	800 mL	1 mL	6	<2		1 mL
680-88766-B-23 MS	032613-RB-shovel	3520C, 8270C LL	T	960 mL	1 mL	6	<2	1 mL	1 mL

Batch Notes	
Acid used for pH adjustment	SULFURIC ACID
Acid used for pH adjust Lot #	EX 10H2S04 _6
Batch Comment	RUSH
Concentration End Time	6.30/4/3/13
Concentration Start Time	5.35/4/3/13
Person's name who did the concentration	AG
Exchange Solvent Lot #	NA/
Time the first extraction ended 24hr	12.45/4/2/13
Time the first extraction started 24 hr	8.45/4/2/13
pH Paper Lot Number	HC 273036
Prep Solvent Lot #	EX MC CYCL _55
Prep Solvent Name	DCM
Prep Solvent Volume Used	210 mL
Person's name who did the prep	AG
Person's name who witnessed reagent drop	RYAN
Sufficient volume for MS/MSD?	MS ONLY
Water Bath ID	TURBOVAP2#1
Water Bath Temperature	40 C Celsius

The pound sign (#) in the amount added field denotes that the reagent was used undiluted. All calculations are performed using the stated concentration for this reagent.

8270C LL

Page 1 of 2

GC/MS SEMI VOA BATCH WORKSHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Batch Number: 136013 Batch Start Date: 04/02/13 08:08 Batch Analyst: George, Abraham

Batch Method: 3520C Batch End Date: 04/03/13 10:55

Basis	Basis Description
T	Total/NA

The pound sign (#) in the amount added field denotes that the reagent was used undiluted. All calculations are performed using the stated concentration for this reagent.

## GC/MS SEMI VOA BATCH WORKSHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2SDG No.: 68088766-2Batch Number: 136063 Batch Start Date: 04/02/13 11:33 Batch Analyst: Cerome, SaurelBatch Method: 3546 Batch End Date: 04/03/13 08:20

Lab Sample ID	Client Sample ID	Method Chain	Basis	InitialAmount	FinalAmount	EX-625LVI SPK 00020	EXLLSURINT 00178		
MB 660-136063/1		3546, 8270C LL		14.99 g	1 mL		1 mL		
LCS 660-136063/2		3546, 8270C LL		14.97 g	1 mL	1 mL	1 mL		
680-88766-A-21	CV0014AB-GS	3546, 8270C LL	T	15.25 g	1 mL		1 mL		
680-88766-A-21 MS	CV0014AB-GS	3546, 8270C LL	T	15.32 g	1 mL	1 mL	1 mL		
680-88766-A-21 MSD	CV0014AB-GS	3546, 8270C LL	T	14.60 g	1 mL	1 mL	1 mL		

Batch Notes	
Acetone Lot #	EX-ACETON BOT 51
Balance ID	B001
Batch Comment	RUSH
Person's name who did the concentration	RYAN
Exchange Solvent Lot #	EX-MC CYCL 55
Exchange Solvent Name	DCM
Final Concentrator Volume	1 mL
MeCl2 Lot #	EX-MC CYCL 55
MeCl2/Acetone Lot #	DCM/ACETON 58
Microwave Start Time	12:30 4/2/13
Microwave Stop Time	13:05 4/2/13
Na2SO4 Lot Number	EX-NA2S04A 65
Ottawa Sand Lot #	EX-OTTOWA SAND 14
Person's name who did the prep	RYAN
SOP Number	TP-EX014
Person who witnessed spiking	SC
Surrogate Lot Number	EXLLSURINT 178
Water Bath ID	TURBOVAP2 #1-4
Water Bath Temperature	40

The pound sign (#) in the amount added field denotes that the reagent was used undiluted. All calculations are performed using the stated concentration for this reagent.

GC/MS SEMI VOA BATCH WORKSHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Batch Number: 136063 Batch Start Date: 04/02/13 11:33 Batch Analyst: Cerome, Saurel

Batch Method: 3546 Batch End Date: 04/03/13 08:20

Basis	Basis Description
T	Total/NA

The pound sign (#) in the amount added field denotes that the reagent was used undiluted. All calculations are performed using the stated concentration for this reagent.

# **GENERAL CHEMISTRY**



COVER PAGE  
GENERAL CHEMISTRY

Lab Name: TestAmerica Tampa Job Number: 680-88766-2

SDG No.: 68088766-2

Project: 35th Avenue Superfund Site

Client Sample ID  
CV0014AB-GS

Lab Sample ID  
680-88766-21

Comments:

9-IN  
DETECTION LIMITS  
GENERAL CHEMISTRY

Lab Name: TestAmerica Tampa Job Number: 680-88766-2  
SDG Number: 68088766-2  
Matrix: Solid Instrument ID: NOEQUIP  
Method: Moisture RL Date: 01/01/2004 18:10

Analyte	Wavelength/ Mass	RL (%)	
Percent Moisture		0.1	

9-IN  
CALIBRATION BLANK DETECTION LIMITS  
GENERAL CHEMISTRY

Lab Name: TestAmerica Tampa Job Number: 680-88766-2  
SDG Number: 68088766-2  
Matrix: Solid Instrument ID: NOEQUIP  
Method: Moisture XRL Date: 04/12/2010 08:14

Analyte	Wavelength/ Mass	XRL (%)	
Percent Moisture		0.1	



13-IN  
ANALYSIS RUN LOG  
GENERAL CHEMISTRY

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Instrument ID: NOEQUIP Method: Moisture

Start Date: 03/29/2013 10:07 End Date: 03/29/2013 10:07

Lab Sample ID	D / F	Type	Time	Analytes															
				M	o	i	s	t											
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
680-88766-A-6 MS	1	T	10:07	X															
680-88766-A-6 MSD	1	T	10:07	X															
680-88766-21	1	T	10:07	X															
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																
ZZZZZZ			10:07																

Prep Types  
T = Total/NA

GENERAL CHEMISTRY BATCH WORKSHEET

Lab Name: TestAmerica Tampa Job No.: 680-88766-2

SDG No.: 68088766-2

Batch Number: 135922 Batch Start Date: 03/29/13 10:07 Batch Analyst: Galio, Andrew

Batch Method: Moisture Batch End Date: \_\_\_\_\_

Lab Sample ID	Client Sample ID	Method Chain	Basis	DISH#	DishWeight	SampleMassWet	SampleMassDry		
680-88766-A-6 MS		Moisture	T	51	0 g	4.48 g	3.80 g		
680-88766-A-6 MSD		Moisture	T	51	0 g	4.48 g	3.80 g		
680-88766-A-21	CV0014AB-GS	Moisture	T	52	0 g	4.71 g	3.48 g		

Batch Notes	
Balance ID	2 No Unit
Date samples were placed in the oven	3.29.13
Date samples were removed from oven	3/30/13
Time Samples were removed from oven	8:30

Basis	Basis Description
T	Total/NA

The pound sign (#) in the amount added field denotes that the reagent was used undiluted. All calculations are performed using the stated concentration for this reagent.

Moisture

# Shipping and Receiving Documents

ANALYSIS REQUEST AND CHAIN OF CUSTODY RECORD

# TestAmerica

THE LEADER IN ENVIRONMENTAL TESTING

TestAmerica Savannah  
5102 LaRoche Avenue  
Savannah, GA 31404

Website: www.testamericainc.com  
Phone: (912) 354-7858  
Fax: (912) 352-0165

Alternate Laboratory Name/Location

Phone:  
Fax:

PROJECT REFERENCE <b>35th Ave Removal</b>	PROJECT NO. <b>2005148-1356</b>	PROJECT LOCATION (STATE) <b>AL</b>	MATRIX TYPE	REQUIRED ANALYSIS	PAGE <b>2</b>	OF <b>2</b>
--	------------------------------------	---------------------------------------	-------------	-------------------	---------------	-------------

TAL (LAB) PROJECT MANAGER <i>[Signature]</i>	P.O. NUMBER	CONTRACT NO.	ENT FAX	COMPOSITE (C) OR GRAB (G) INDICATE	AGUEOUS (WATER)	SOLID OR SEMISOLID	AIR	NONAQUEOUS LIQUID (OIL, SOLVENT, ...)	STANDARD REPORT DELIVERY <input type="radio"/>	DATE DUE _____
<p><b>LLPAH</b> <b>RELA 8 Metals</b></p>								EXPEDITED REPORT DELIVERY (SURCHARGE) <input type="radio"/>	DATE DUE _____	

(b) (6)  
(b) (6)  
(b) (6)

COMPANY CONT. *[Signature]*

**PRESERVATIVE**

NUMBER OF COOLERS SUBMITTED PER SHIPMENT: \_\_\_\_\_

SAMPLE		SAMPLE IDENTIFICATION	COMPOSITE (C) OR GRAB (G)	INDICATE	AGUEOUS (WATER)	SOLID OR SEMISOLID	AIR	NONAQUEOUS LIQUID (OIL, SOLVENT, ...)	NUMBER OF CONTAINERS SUBMITTED										REMARKS		
DATE	TIME								1	2	3	4	5	6	7	8	9	10		11	12
3-25-13	1428	CVO613K-CSD	C	X				X	X												
	1332	CVO613AB-GS	G	X				X													
	1334	CVO613AC-GS	G	X				X													
	1440	CVO610A-CS	C	X				X													
	1442	CVO610B-CS	C	X				X													
	1439	CVO610AB-GS	G	X				X													
	1506	CVO506A-CS	C	X				X													
	1515	CVO506B-CS	C	X				X													
	1518	CVO014AB-GS	G	X				X													
	1343	CVO613E-CS (sieve)	C	X				X	X												
3-26-13	1300	032613-RB-Shovel		X				X	X												

RELINQUISHED BY: (SIGNATURE) <i>[Signature]</i>	DATE 3-26-13	TIME 1330	RELINQUISHED BY: (SIGNATURE)	DATE	TIME	RELINQUISHED BY: (SIGNATURE)	DATE	TIME
RECEIVED BY: (SIGNATURE)	DATE	TIME	RECEIVED BY: (SIGNATURE)	DATE	TIME	RECEIVED BY: (SIGNATURE)	DATE	TIME

LABORATORY USE ONLY

RECEIVED FOR LABORATORY BY: (SIGNATURE) <i>[Signature]</i>	DATE 03/28/13	TIME 0937	CUSTODY INTACT YES <input type="radio"/> NO <input type="radio"/>	CUSTODY SEAL NO.	SAVANNAH LOG NO. 680-88766	LABORATORY REMARKS 1-y <sup>c</sup>
---	------------------	--------------	---	------------------	-------------------------------	--



## Login Sample Receipt Checklist

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

SDG Number: 68088766-2

**Login Number: 88766**

**List Source: TestAmerica Savannah**

**List Number: 1**

**Creator: Barnett, Eddie T**

Question	Answer	Comment
Radioactivity wasn't checked or is <= background as measured by a survey meter.	N/A	
The cooler's custody seal, if present, is intact.	True	
Sample custody seals, if present, are intact.	True	
The cooler or samples do not appear to have been compromised or tampered with.	True	
Samples were received on ice.	True	
Cooler Temperature is acceptable.	True	
Cooler Temperature is recorded.	True	
COC is present.	True	
COC is filled out in ink and legible.	True	
COC is filled out with all pertinent information.	True	
Is the Field Sampler's name present on COC?	N/A	
There are no discrepancies between the containers received and the COC.	True	
Samples are received within Holding Time.	True	
Sample containers have legible labels.	True	
Containers are not broken or leaking.	True	
Sample collection date/times are provided.	True	
Appropriate sample containers are used.	True	
Sample bottles are completely filled.	True	
Sample Preservation Verified.	True	
There is sufficient vol. for all requested analyses, incl. any requested MS/MSDs	True	
Containers requiring zero headspace have no headspace or bubble is <6mm (1/4").	N/A	
Multiphasic samples are not present.	True	
Samples do not require splitting or compositing.	True	
Residual Chlorine Checked.	N/A	

## Login Sample Receipt Checklist

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

SDG Number: 68088766-2

Login Number: 88766

List Source: TestAmerica Tampa

List Number: 1

List Creation: 03/29/13 09:17 AM

Creator: McNulty, Carol

Question	Answer	Comment
Radioactivity wasn't checked or is $\leq$ background as measured by a survey meter.	True	
The cooler's custody seal, if present, is intact.	True	
Sample custody seals, if present, are intact.	True	
The cooler or samples do not appear to have been compromised or tampered with.	True	
Samples were received on ice.	True	
Cooler Temperature is acceptable.	True	
Cooler Temperature is recorded.	True	
COC is present.	True	
COC is filled out in ink and legible.	True	
COC is filled out with all pertinent information.	True	
Is the Field Sampler's name present on COC?	True	
There are no discrepancies between the containers received and the COC.	True	
Samples are received within Holding Time.	True	
Sample containers have legible labels.	True	
Containers are not broken or leaking.	True	
Sample collection date/times are provided.	True	
Appropriate sample containers are used.	True	
Sample bottles are completely filled.	True	
Sample Preservation Verified.	True	
There is sufficient vol. for all requested analyses, incl. any requested MS/MSDs	True	
Containers requiring zero headspace have no headspace or bubble is $<6\text{mm}$ (1/4").	True	
Multiphasic samples are not present.	True	
Samples do not require splitting or compositing.	True	
Residual Chlorine Checked.	True	

# TestAmerica

THE LEADER IN ENVIRONMENTAL TESTING

## ANALYTICAL REPORT

TestAmerica Laboratories, Inc.

TestAmerica Savannah

5102 LaRoche Avenue

Savannah, GA 31404

Tel: (912)354-7858

TestAmerica Job ID: 680-88766-2

TestAmerica Sample Delivery Group: 68088766-2

Client Project/Site: 35th Avenue Superfund Site

For:

Oneida Total Integrated Enterprises LLC

1220 Kennestone Circle

Suite 106

Marietta, Georgia 30060

Attn: Ms. Limari F Krebs



Authorized for release by:

4/8/2013 9:02:06 AM

Bernard Kirkland

Project Manager I

[bernard.kirkland@testamericainc.com](mailto:bernard.kirkland@testamericainc.com)

Designee for

Lisa Harvey

Project Manager II

[lisa.harvey@testamericainc.com](mailto:lisa.harvey@testamericainc.com)

*The test results in this report meet all 2003 NELAC and 2009 TNI requirements for accredited parameters, exceptions are noted in this report. This report may not be reproduced except in full, and with written approval from the laboratory. For questions please contact the Project Manager at the e-mail address or telephone number listed on this page.*

*This report has been electronically signed and authorized by the signatory. Electronic signature is intended to be the legally binding equivalent of a traditionally handwritten signature.*

*Results relate only to the items tested and the sample(s) as received by the laboratory.*



### LINKS

Review your project  
results through

**TotalAccess**

Have a Question?



Visit us at:

[www.testamericainc.com](http://www.testamericainc.com)

1

2

3

4

5

6

7

8

9

10

11

12

# Case Narrative

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

**Job ID: 680-88766-2**

**Laboratory: TestAmerica Savannah**

Narrative

## CASE NARRATIVE

**Client: Oneida Total Integrated Enterprises LLC**

**Project: 35th Avenue Superfund Site**

**Report Number: 680-88766-2**

With the exceptions noted as flags or footnotes, standard analytical protocols were followed in the analysis of the samples and no problems were encountered or anomalies observed. In addition all laboratory quality control samples were within established control limits, with any exceptions noted below. Each sample was analyzed to achieve the lowest possible reporting limit within the constraints of the method. In some cases, due to interference or analytes present at high concentrations, samples were diluted. For diluted samples, the reporting limits are adjusted relative to the dilution required.

Calculations are performed before rounding to avoid round-off errors in calculated results.

All holding times were met and proper preservation noted for the methods performed on these samples, unless otherwise detailed in the individual sections below.

### RECEIPT

The samples were received on 03/28/2013; the samples arrived in good condition, properly preserved and on ice. The temperature of the coolers at receipt was 1.4 C.

### SEMIVOLATILE ORGANIC COMPOUNDS BY GCMS - LOW LEVEL

Sample CV0014AB-GS (680-88766-21) was analyzed for Semivolatile Organic Compounds by GCMS - Low Level in accordance with EPA SW-846 Method 8270C. The samples were prepared on 04/02/2013 and analyzed on 04/03/2013.

Sample CV0014AB-GS (680-88766-21)[4X] required dilution prior to analysis. The reporting limits have been adjusted accordingly.

No difficulties were encountered during the SVOAs analysis.

All quality control parameters were within the acceptance limits.

### SEMIVOLATILE ORGANIC COMPOUNDS (GC-MS)-WATER

Sample 032613-RB-shovel (680-88766-23) was analyzed for semivolatile organic compounds (GC-MS) in accordance with EPA SW-846 Method 8270C. The samples were prepared on 04/02/2013 and analyzed on 04/05/2013.

Several analytes recovered outside the recovery criteria low for the MS of sample 032613-RB-shovel (680-88766-23) in batch 660-136171.

No other difficulties were encountered during the semivolatiles analysis.

All other quality control parameters were within the acceptance limits.

# Sample Summary

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

Lab Sample ID	Client Sample ID	Matrix	Collected	Received
680-88766-21	CV0014AB-GS	Solid	03/25/13 15:18	03/28/13 09:37
680-88766-23	032613-RB-shovel	Water	03/26/13 13:00	03/28/13 09:37

1

2

3

4

5

6

7

8

9

10

11

12

# Method Summary

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

Method	Method Description	Protocol	Laboratory
8270C LL	Semivolatile Organic Compounds by GCMS - Low Levels	SW846	TAL TAM
Moisture	Percent Moisture	EPA	TAL TAM

**Protocol References:**

EPA = US Environmental Protection Agency

SW846 = "Test Methods For Evaluating Solid Waste, Physical/Chemical Methods", Third Edition, November 1986 And Its Updates.

**Laboratory References:**

TAL TAM = TestAmerica Tampa, 6712 Benjamin Road, Suite 100, Tampa, FL 33634, TEL (813)885-7427

1

2

3

4

5

6

7

8

9

10

11

12

## Definitions/Glossary

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

### Qualifiers

#### GC/MS Semi VOA

Qualifier	Qualifier Description
J	Result is less than the RL but greater than or equal to the MDL and the concentration is an approximate value.
U	Indicates the analyte was analyzed for but not detected.
F	MS or MSD exceeds the control limits

### Glossary

Abbreviation	These commonly used abbreviations may or may not be present in this report.
α	Listed under the "D" column to designate that the result is reported on a dry weight basis
%R	Percent Recovery
CNF	Contains no Free Liquid
DER	Duplicate error ratio (normalized absolute difference)
DL, RA, RE, IN	Indicates a Dilution, Re-analysis, Re-extraction, or additional Initial metals/anion analysis of the sample
DLC	Decision level concentration
MDA	Minimum detectable activity
EDL	Estimated Detection Limit
MDC	Minimum detectable concentration
MDL	Method Detection Limit
ML	Minimum Level (Dioxin)
ND	Not detected at the reporting limit (or MDL or EDL if shown)
PQL	Practical Quantitation Limit
QC	Quality Control
RER	Relative error ratio
RL	Reporting Limit or Requested Limit (Radiochemistry)
RPD	Relative Percent Difference, a measure of the relative difference between two points
TEF	Toxicity Equivalent Factor (Dioxin)
TEQ	Toxicity Equivalent Quotient (Dioxin)

# Client Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

**Client Sample ID: CV0014AB-GS**

**Lab Sample ID: 680-88766-21**

Date Collected: 03/25/13 15:18

Matrix: Solid

Date Received: 03/28/13 09:37

Percent Solids: 73.9

**Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels**

Analyte	Result	Qualifier	RL	MDL	Unit	D	Prepared	Analyzed	Dil Fac
Acenaphthene	530	U	530	110	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Acenaphthylene</b>	<b>34</b>	<b>J</b>	210	27	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
Anthracene	45	U	45	22	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Benzo[a]anthracene</b>	<b>200</b>		43	21	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Benzo[a]pyrene</b>	<b>100</b>		55	28	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Benzo[b]fluoranthene</b>	<b>150</b>		65	32	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Benzo[g,h,i]perylene</b>	<b>130</b>		110	23	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Benzo[k]fluoranthene</b>	<b>79</b>		43	19	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Chrysene</b>	<b>210</b>		48	24	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Dibenz(a,h)anthracene</b>	<b>62</b>	<b>J</b>	110	22	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Fluoranthene</b>	<b>210</b>		110	21	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Fluorene</b>	<b>39</b>	<b>J</b>	110	22	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Indeno[1,2,3-cd]pyrene</b>	<b>76</b>	<b>J</b>	110	38	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>1-Methylnaphthalene</b>	<b>120</b>	<b>J</b>	210	23	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>2-Methylnaphthalene</b>	<b>130</b>	<b>J</b>	210	38	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Naphthalene</b>	<b>100</b>	<b>J</b>	210	23	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Phenanthrene</b>	<b>200</b>		43	21	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4
<b>Pyrene</b>	<b>210</b>		110	20	ug/Kg	☼	04/02/13 11:33	04/03/13 16:10	4

Surrogate	%Recovery	Qualifier	Limits	Prepared	Analyzed	Dil Fac
<i>o</i> -Terphenyl	95		30 - 130	04/02/13 11:33	04/03/13 16:10	4

**Client Sample ID: 032613-RB-shovel**

**Lab Sample ID: 680-88766-23**

Date Collected: 03/26/13 13:00

Matrix: Water

Date Received: 03/28/13 09:37

**Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels**

Analyte	Result	Qualifier	RL	MDL	Unit	D	Prepared	Analyzed	Dil Fac
Acenaphthene	2.5	U F	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
Acenaphthylene	1.3	U	1.3	0.31	ug/L		04/02/13 08:08	04/05/13 12:54	1
Anthracene	0.25	U	0.25	0.095	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[a]anthracene	0.25	U F	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[a]pyrene	0.25	U F	0.25	0.071	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[b]fluoranthene	0.25	U F	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[g,h,i]perylene	0.63	U F	0.63	0.13	ug/L		04/02/13 08:08	04/05/13 12:54	1
Benzo[k]fluoranthene	0.25	U F	0.25	0.071	ug/L		04/02/13 08:08	04/05/13 12:54	1
Chrysene	0.25	U F	0.25	0.086	ug/L		04/02/13 08:08	04/05/13 12:54	1
Dibenz(a,h)anthracene	0.25	U F	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
Fluoranthene	0.63	U F	0.63	0.068	ug/L		04/02/13 08:08	04/05/13 12:54	1
Fluorene	2.5	U F	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
Indeno[1,2,3-cd]pyrene	0.25	U F	0.25	0.063	ug/L		04/02/13 08:08	04/05/13 12:54	1
1-Methylnaphthalene	2.5	U	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
2-Methylnaphthalene	2.5	U F	2.5	0.63	ug/L		04/02/13 08:08	04/05/13 12:54	1
Naphthalene	2.5	U F	2.5	0.31	ug/L		04/02/13 08:08	04/05/13 12:54	1
Phenanthrene	0.63	U F	0.63	0.25	ug/L		04/02/13 08:08	04/05/13 12:54	1
Pyrene	0.63	U F	0.63	0.11	ug/L		04/02/13 08:08	04/05/13 12:54	1

Surrogate	%Recovery	Qualifier	Limits	Prepared	Analyzed	Dil Fac
<i>o</i> -Terphenyl	67		30 - 130	04/02/13 08:08	04/05/13 12:54	1

TestAmerica Savannah



# QC Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

## Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels

**Lab Sample ID: MB 660-136013/1-A**

**Matrix: Water**

**Analysis Batch: 136164**

**Client Sample ID: Method Blank**

**Prep Type: Total/NA**

**Prep Batch: 136013**

Analyte	MB Result	MB Qualifier	RL	MDL	Unit	D	Prepared	Analyzed	Dil Fac
Acenaphthene	2.0	U	2.0	0.50	ug/L		04/02/13 08:08	04/04/13 19:51	1
Acenaphthylene	1.0	U	1.0	0.25	ug/L		04/02/13 08:08	04/04/13 19:51	1
Anthracene	0.20	U	0.20	0.076	ug/L		04/02/13 08:08	04/04/13 19:51	1
Benzo[a]anthracene	0.20	U	0.20	0.050	ug/L		04/02/13 08:08	04/04/13 19:51	1
Benzo[a]pyrene	0.20	U	0.20	0.057	ug/L		04/02/13 08:08	04/04/13 19:51	1
Benzo[b]fluoranthene	0.20	U	0.20	0.050	ug/L		04/02/13 08:08	04/04/13 19:51	1
Benzo[g,h,i]perylene	0.50	U	0.50	0.10	ug/L		04/02/13 08:08	04/04/13 19:51	1
Benzo[k]fluoranthene	0.20	U	0.20	0.057	ug/L		04/02/13 08:08	04/04/13 19:51	1
Chrysene	0.20	U	0.20	0.069	ug/L		04/02/13 08:08	04/04/13 19:51	1
Dibenz(a,h)anthracene	0.20	U	0.20	0.050	ug/L		04/02/13 08:08	04/04/13 19:51	1
Fluoranthene	0.50	U	0.50	0.054	ug/L		04/02/13 08:08	04/04/13 19:51	1
Fluorene	2.0	U	2.0	0.50	ug/L		04/02/13 08:08	04/04/13 19:51	1
Indeno[1,2,3-cd]pyrene	0.20	U	0.20	0.050	ug/L		04/02/13 08:08	04/04/13 19:51	1
1-Methylnaphthalene	2.0	U	2.0	0.50	ug/L		04/02/13 08:08	04/04/13 19:51	1
2-Methylnaphthalene	2.0	U	2.0	0.50	ug/L		04/02/13 08:08	04/04/13 19:51	1
Naphthalene	2.0	U	2.0	0.25	ug/L		04/02/13 08:08	04/04/13 19:51	1
Phenanthrene	0.50	U	0.50	0.20	ug/L		04/02/13 08:08	04/04/13 19:51	1
Pyrene	0.50	U	0.50	0.089	ug/L		04/02/13 08:08	04/04/13 19:51	1

Surrogate	MB %Recovery	MB Qualifier	Limits	Prepared	Analyzed	Dil Fac
<i>o</i> -Terphenyl	78		30 - 130	04/02/13 08:08	04/04/13 19:51	1

**Lab Sample ID: LCS 660-136013/2-A**

**Matrix: Water**

**Analysis Batch: 136164**

**Client Sample ID: Lab Control Sample**

**Prep Type: Total/NA**

**Prep Batch: 136013**

Analyte	Spike Added	LCS Result	LCS Qualifier	Unit	D	%Rec	%Rec. Limits
Acenaphthene	10.0	7.38		ug/L		74	55 - 132
Acenaphthylene	10.0	7.55		ug/L		76	39 - 130
Anthracene	10.0	7.12		ug/L		71	39 - 130
Benzo[a]anthracene	10.0	7.21		ug/L		72	54 - 135
Benzo[a]pyrene	10.0	4.96		ug/L		50	21 - 130
Benzo[b]fluoranthene	10.0	5.86		ug/L		59	37 - 130
Benzo[g,h,i]perylene	10.0	3.51		ug/L		35	26 - 130
Benzo[k]fluoranthene	10.0	5.75		ug/L		57	38 - 130
Chrysene	10.0	7.28		ug/L		73	56 - 130
Dibenz(a,h)anthracene	10.0	3.33		ug/L		33	13 - 130
Fluoranthene	10.0	7.83		ug/L		78	60 - 130
Fluorene	10.0	8.04		ug/L		80	55 - 140
Indeno[1,2,3-cd]pyrene	10.0	3.17		ug/L		32	21 - 130
1-Methylnaphthalene	10.0	7.86		ug/L		79	49 - 130
2-Methylnaphthalene	10.0	7.57		ug/L		76	48 - 130
Naphthalene	10.0	7.44		ug/L		74	54 - 133
Phenanthrene	10.0	7.52		ug/L		75	60 - 136
Pyrene	10.0	7.50		ug/L		75	60 - 138

TestAmerica Savannah

# QC Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

## Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels (Continued)

**Lab Sample ID: LCS 660-136013/2-A**  
**Matrix: Water**  
**Analysis Batch: 136164**

**Client Sample ID: Lab Control Sample**  
**Prep Type: Total/NA**  
**Prep Batch: 136013**

Surrogate	LCS		Limits
	%Recovery	Qualifier	
<i>o</i> -Terphenyl	72		30 - 130

**Lab Sample ID: LCSD 660-136013/3-A**  
**Matrix: Water**  
**Analysis Batch: 136164**

**Client Sample ID: Lab Control Sample Dup**  
**Prep Type: Total/NA**  
**Prep Batch: 136013**

Analyte	Spike Added	LCSD		Unit	D	%Rec	%Rec.		RPD	Limit
		Result	Qualifier				Limits	RPD		
Acenaphthene	10.0	7.35		ug/L		73	55 - 132	0	35	
Acenaphthylene	10.0	7.42		ug/L		74	39 - 130	2	35	
Anthracene	10.0	7.14		ug/L		71	39 - 130	0	35	
Benzo[a]anthracene	10.0	7.19		ug/L		72	54 - 135	0	35	
Benzo[a]pyrene	10.0	5.01		ug/L		50	21 - 130	1	35	
Benzo[b]fluoranthene	10.0	5.92		ug/L		59	37 - 130	1	35	
Benzo[g,h,i]perylene	10.0	3.91		ug/L		39	26 - 130	11	35	
Benzo[k]fluoranthene	10.0	5.65		ug/L		57	38 - 130	2	35	
Chrysene	10.0	6.91		ug/L		69	56 - 130	5	35	
Dibenz(a,h)anthracene	10.0	3.54		ug/L		35	13 - 130	6	35	
Fluoranthene	10.0	7.94		ug/L		79	60 - 130	1	35	
Fluorene	10.0	7.90		ug/L		79	55 - 140	2	35	
Indeno[1,2,3-cd]pyrene	10.0	3.59		ug/L		36	21 - 130	12	35	
1-Methylnaphthalene	10.0	7.77		ug/L		78	49 - 130	1	35	
2-Methylnaphthalene	10.0	7.56		ug/L		76	48 - 130	0	35	
Naphthalene	10.0	7.36		ug/L		74	54 - 133	1	35	
Phenanthrene	10.0	7.41		ug/L		74	60 - 136	1	35	
Pyrene	10.0	7.46		ug/L		75	60 - 138	0	35	

Surrogate	LCSD		Limits
	%Recovery	Qualifier	
<i>o</i> -Terphenyl	72		30 - 130

**Lab Sample ID: 680-88766-23 MS**  
**Matrix: Water**  
**Analysis Batch: 136171**

**Client Sample ID: 032613-RB-shovel**  
**Prep Type: Total/NA**  
**Prep Batch: 136013**

Analyte	Sample Result	Sample Qualifier	Spike Added	MS		Unit	D	%Rec	%Rec.	
				Result	Qualifier				Limits	RPD
Acenaphthene	2.5	U F	10.4	4.74	F	ug/L		45	55 - 132	
Acenaphthylene	1.3	U	10.4	5.18		ug/L		50	39 - 130	
Anthracene	0.25	U	10.4	4.97		ug/L		48	39 - 130	
Benzo[a]anthracene	0.25	U F	10.4	4.72	F	ug/L		45	54 - 135	
Benzo[a]pyrene	0.25	U F	10.4	2.13	F	ug/L		20	21 - 130	
Benzo[b]fluoranthene	0.25	U F	10.4	2.68	F	ug/L		26	37 - 130	
Benzo[g,h,i]perylene	0.63	U F	10.4	1.33	F	ug/L		13	26 - 130	
Benzo[k]fluoranthene	0.25	U F	10.4	2.38	F	ug/L		23	38 - 130	
Chrysene	0.25	U F	10.4	3.55	F	ug/L		34	56 - 130	
Dibenz(a,h)anthracene	0.25	U F	10.4	1.12	F	ug/L		11	13 - 130	
Fluoranthene	0.63	U F	10.4	5.46	F	ug/L		52	60 - 130	
Fluorene	2.5	U F	10.4	4.83	F	ug/L		46	55 - 140	
Indeno[1,2,3-cd]pyrene	0.25	U F	10.4	1.28	F	ug/L		12	21 - 130	
1-Methylnaphthalene	2.5	U	10.4	5.64		ug/L		54	49 - 130	

TestAmerica Savannah

# QC Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

## Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels (Continued)

**Lab Sample ID: 680-88766-23 MS**

**Matrix: Water**

**Analysis Batch: 136171**

**Client Sample ID: 032613-RB-shovel**

**Prep Type: Total/NA**

**Prep Batch: 136013**

Analyte	Sample	Sample	Spike	MS	MS	Unit	D	%Rec	%Rec.
	Result	Qualifier	Added	Result	Qualifier				
2-Methylnaphthalene	2.5	U F	10.4	4.68	F	ug/L		45	48 - 130
Naphthalene	2.5	U F	10.4	5.39	F	ug/L		52	54 - 133
Phenanthrene	0.63	U F	10.4	5.42	F	ug/L		52	60 - 136
Pyrene	0.63	U F	10.4	5.24	F	ug/L		50	60 - 138
<b>MS MS</b>									
Surrogate	%Recovery	Qualifier	Limits						
<i>o</i> -Terphenyl	49		30 - 130						

**Lab Sample ID: MB 660-136063/1-A**

**Matrix: Solid**

**Analysis Batch: 136081**

**Client Sample ID: Method Blank**

**Prep Type: Total/NA**

**Prep Batch: 136063**

Analyte	MB	MB	RL	MDL	Unit	D	Prepared	Analyzed	Dil Fac
	Result	Qualifier							
Acenaphthene	100	U	100	20	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Acenaphthylene	40	U	40	5.0	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Anthracene	8.4	U	8.4	4.2	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Benzo[a]anthracene	8.0	U	8.0	3.9	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Benzo[a]pyrene	10	U	10	5.2	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Benzo[b]fluoranthene	12	U	12	6.1	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Benzo[g,h,i]perylene	20	U	20	4.4	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Benzo[k]fluoranthene	8.0	U	8.0	3.6	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Chrysene	9.0	U	9.0	4.5	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Dibenz(a,h)anthracene	20	U	20	4.1	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Fluoranthene	20	U	20	4.0	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Fluorene	20	U	20	4.1	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Indeno[1,2,3-cd]pyrene	20	U	20	7.1	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
1-Methylnaphthalene	40	U	40	4.4	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
2-Methylnaphthalene	40	U	40	7.1	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Naphthalene	40	U	40	4.4	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Phenanthrene	8.0	U	8.0	3.9	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
Pyrene	20	U	20	3.7	ug/Kg		04/02/13 11:33	04/03/13 15:34	1
<b>MB MB</b>									
Surrogate	%Recovery	Qualifier	Limits	Prepared	Analyzed	Dil Fac			
<i>o</i> -Terphenyl	80		30 - 130	04/02/13 11:33	04/03/13 15:34	1			

**Lab Sample ID: LCS 660-136063/2-A**

**Matrix: Solid**

**Analysis Batch: 136081**

**Client Sample ID: Lab Control Sample**

**Prep Type: Total/NA**

**Prep Batch: 136063**

Analyte	Spike	LCS	LCS	Unit	D	%Rec	%Rec.
Acenaphthene	668	458		ug/Kg		69	39 - 130
Acenaphthylene	668	488		ug/Kg		73	38 - 130
Anthracene	668	469		ug/Kg		70	37 - 130
Benzo[a]anthracene	668	484		ug/Kg		72	40 - 130
Benzo[a]pyrene	668	444		ug/Kg		66	49 - 130
Benzo[b]fluoranthene	668	516		ug/Kg		77	37 - 130
Benzo[g,h,i]perylene	668	418		ug/Kg		63	32 - 130
Benzo[k]fluoranthene	668	468		ug/Kg		70	32 - 130

TestAmerica Savannah

## QC Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

### Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels (Continued)

**Lab Sample ID: LCS 660-136063/2-A**

**Matrix: Solid**

**Analysis Batch: 136081**

**Client Sample ID: Lab Control Sample**

**Prep Type: Total/NA**

**Prep Batch: 136063**

Analyte	Spike Added	LCS Result	LCS Qualifier	Unit	D	%Rec	%Rec.	
							Limits	
Chrysene	668	462		ug/Kg		69	41 - 130	
Dibenz(a,h)an hracene	668	471		ug/Kg		71	27 - 130	
Fluoranthene	668	487		ug/Kg		73	40 - 130	
Fluorene	668	444		ug/Kg		66	40 - 130	
Indeno[1,2,3-cd]pyrene	668	399		ug/Kg		60	30 - 130	
1-Methylnaphthalene	668	522		ug/Kg		78	31 - 130	
2-Methylnaphthalene	668	458		ug/Kg		69	33 - 130	
Naphthalene	668	484		ug/Kg		72	36 - 130	
Phenanthrene	668	499		ug/Kg		75	42 - 130	
Pyrene	668	516		ug/Kg		77	44 - 130	

Surrogate	LCS LCS		Limits
	%Recovery	Qualifier	
<i>o</i> -Terphenyl	72		30 - 130

**Lab Sample ID: 680-88766-21 MS**

**Matrix: Solid**

**Analysis Batch: 136081**

**Client Sample ID: CV0014AB-GS**

**Prep Type: Total/NA**

**Prep Batch: 136063**

Analyte	Sample Result	Sample Qualifier	Spike Added	MS Result	MS Qualifier	Unit	D	%Rec	%Rec.	
									Limits	
Acenaphthene	530	U	883	656		ug/Kg	☼	74	39 - 130	
Acenaphthylene	34	J	883	626		ug/Kg	☼	67	38 - 130	
Anthracene	45	U	883	666		ug/Kg	☼	75	37 - 130	
Benzo[a]anthracene	200		883	833		ug/Kg	☼	72	40 - 130	
Benzo[a]pyrene	100		883	650		ug/Kg	☼	62	49 - 130	
Benzo[b]fluoranthene	150		883	738		ug/Kg	☼	66	37 - 130	
Benzo[g,h,i]perylene	130		883	623		ug/Kg	☼	56	32 - 130	
Benzo[k]fluoranthene	79		883	667		ug/Kg	☼	67	32 - 130	
Chrysene	210		883	780		ug/Kg	☼	65	41 - 130	
Dibenz(a,h)an hracene	62	J	883	622		ug/Kg	☼	63	27 - 130	
Fluoranthene	210		883	697		ug/Kg	☼	55	40 - 130	
Fluorene	39	J	883	639		ug/Kg	☼	68	40 - 130	
Indeno[1,2,3-cd]pyrene	76	J	883	532		ug/Kg	☼	52	30 - 130	
1-Methylnaphthalene	120	J	883	653		ug/Kg	☼	60	31 - 130	
2-Methylnaphthalene	130	J	883	653		ug/Kg	☼	59	33 - 130	
Naphthalene	100	J	883	681		ug/Kg	☼	66	36 - 130	
Phenanthrene	200		883	690		ug/Kg	☼	55	42 - 130	
Pyrene	210		883	750		ug/Kg	☼	62	44 - 130	

Surrogate	MS MS		Limits
	%Recovery	Qualifier	
<i>o</i> -Terphenyl	92		30 - 130

**Lab Sample ID: 680-88766-21 MSD**

**Matrix: Solid**

**Analysis Batch: 136081**

**Client Sample ID: CV0014AB-GS**

**Prep Type: Total/NA**

**Prep Batch: 136063**

Analyte	Sample Result	Sample Qualifier	Spike Added	MSD Result	MSD Qualifier	Unit	D	%Rec	%Rec.		RPD	
									Limits		RPD	Limit
Acenaphthene	530	U	927	669		ug/Kg	☼	72	39 - 130		2	40
Acenaphthylene	34	J	927	686		ug/Kg	☼	70	38 - 130		9	40

TestAmerica Savannah

# QC Sample Results

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

## Method: 8270C LL - Semivolatile Organic Compounds by GCMS - Low Levels (Continued)

Lab Sample ID: 680-88766-21 MSD

Matrix: Solid

Analysis Batch: 136081

Client Sample ID: CV0014AB-GS

Prep Type: Total/NA

Prep Batch: 136063

Analyte	Sample	Sample	Spike	MSD	MSD	Unit	D	%Rec	%Rec.	RPD	Limit
	Result	Qualifier	Added	Result	Qualifier				Limits		
Anthracene	45	U	927	674		ug/Kg	☼	73	37 - 130	1	40
Benzo[a]anthracene	200		927	856		ug/Kg	☼	71	40 - 130	3	40
Benzo[a]pyrene	100		927	651		ug/Kg	☼	59	49 - 130	0	40
Benzo[b]fluoranthene	150		927	710		ug/Kg	☼	60	37 - 130	4	40
Benzo[g,h,i]perylene	130		927	618		ug/Kg	☼	53	32 - 130	1	40
Benzo[k]fluoranthene	79		927	822		ug/Kg	☼	80	32 - 130	21	40
Chrysene	210		927	739		ug/Kg	☼	57	41 - 130	5	40
Dibenz(a,h)anthracene	62	J	927	713		ug/Kg	☼	70	27 - 130	14	40
Fluoranthene	210		927	780		ug/Kg	☼	62	40 - 130	11	40
Fluorene	39	J	927	657		ug/Kg	☼	67	40 - 130	3	40
Indeno[1,2,3-cd]pyrene	76	J	927	659		ug/Kg	☼	63	30 - 130	21	40
1-Methylnaphthalene	120	J	927	709		ug/Kg	☼	64	31 - 130	8	40
2-Methylnaphthalene	130	J	927	671		ug/Kg	☼	58	33 - 130	3	40
Naphthalene	100	J	927	648		ug/Kg	☼	59	36 - 130	5	40
Phenanthrene	200		927	758		ug/Kg	☼	60	42 - 130	9	40
Pyrene	210		927	878		ug/Kg	☼	72	44 - 130	16	40
<b>MSD MSD</b>											
<b>Surrogate</b>	<b>%Recovery</b>	<b>Qualifier</b>	<b>Limits</b>								
<i>o</i> -Terphenyl	97		30 - 130								

# QC Association Summary

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

## GC/MS Semi VOA

### Prep Batch: 136013

Lab Sample ID	Client Sample ID	Prep Type	Matrix	Method	Prep Batch
680-88766-23	032613-RB-shovel	Total/NA	Water	3520C	
680-88766-23 MS	032613-RB-shovel	Total/NA	Water	3520C	
LCS 660-136013/2-A	Lab Control Sample	Total/NA	Water	3520C	
LCSD 660-136013/3-A	Lab Control Sample Dup	Total/NA	Water	3520C	
MB 660-136013/1-A	Method Blank	Total/NA	Water	3520C	

### Prep Batch: 136063

Lab Sample ID	Client Sample ID	Prep Type	Matrix	Method	Prep Batch
680-88766-21	CV0014AB-GS	Total/NA	Solid	3546	
680-88766-21 MS	CV0014AB-GS	Total/NA	Solid	3546	
680-88766-21 MSD	CV0014AB-GS	Total/NA	Solid	3546	
LCS 660-136063/2-A	Lab Control Sample	Total/NA	Solid	3546	
MB 660-136063/1-A	Method Blank	Total/NA	Solid	3546	

### Analysis Batch: 136081

Lab Sample ID	Client Sample ID	Prep Type	Matrix	Method	Prep Batch
680-88766-21	CV0014AB-GS	Total/NA	Solid	8270C LL	136063
680-88766-21 MS	CV0014AB-GS	Total/NA	Solid	8270C LL	136063
680-88766-21 MSD	CV0014AB-GS	Total/NA	Solid	8270C LL	136063
LCS 660-136063/2-A	Lab Control Sample	Total/NA	Solid	8270C LL	136063
MB 660-136063/1-A	Method Blank	Total/NA	Solid	8270C LL	136063

### Analysis Batch: 136164

Lab Sample ID	Client Sample ID	Prep Type	Matrix	Method	Prep Batch
LCS 660-136013/2-A	Lab Control Sample	Total/NA	Water	8270C LL	136013
LCSD 660-136013/3-A	Lab Control Sample Dup	Total/NA	Water	8270C LL	136013
MB 660-136013/1-A	Method Blank	Total/NA	Water	8270C LL	136013

### Analysis Batch: 136171

Lab Sample ID	Client Sample ID	Prep Type	Matrix	Method	Prep Batch
680-88766-23	032613-RB-shovel	Total/NA	Water	8270C LL	136013
680-88766-23 MS	032613-RB-shovel	Total/NA	Water	8270C LL	136013

## General Chemistry

### Analysis Batch: 135922

Lab Sample ID	Client Sample ID	Prep Type	Matrix	Method	Prep Batch
680-88766-21	CV0014AB-GS	Total/NA	Solid	Moisture	

# Lab Chronicle

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

**Client Sample ID: CV0014AB-GS**

**Lab Sample ID: 680-88766-21**

**Date Collected: 03/25/13 15:18**

**Matrix: Solid**

**Date Received: 03/28/13 09:37**

**Percent Solids: 73.9**

Prep Type	Batch Type	Batch Method	Run	Dilution Factor	Batch Number	Prepared or Analyzed	Analyst	Lab
Total/NA	Prep	3546			136063	04/02/13 11:33	SC	TAL TAM
Total/NA	Analysis	8270C LL		4	136081	04/03/13 16:10	SCC	TAL TAM
Total/NA	Analysis	Moisture		1	135922	03/29/13 10:07	AG	TAL TAM

**Client Sample ID: 032613-RB-shovel**

**Lab Sample ID: 680-88766-23**

**Date Collected: 03/26/13 13:00**

**Matrix: Water**

**Date Received: 03/28/13 09:37**

Prep Type	Batch Type	Batch Method	Run	Dilution Factor	Batch Number	Prepared or Analyzed	Analyst	Lab
Total/NA	Prep	3520C			136013	04/02/13 08:08	AG	TAL TAM
Total/NA	Analysis	8270C LL		1	136171	04/05/13 12:54	SCC	TAL TAM

**Laboratory References:**

TAL TAM = TestAmerica Tampa, 6712 Benjamin Road, Suite 100, Tampa, FL 33634, TEL (813)885-7427



ANALYSIS REQUEST AND CHAIN OF CUSTODY RECORD

# TestAmerica

THE LEADER IN ENVIRONMENTAL TESTING

TestAmerica Savannah  
5102 LaRoche Avenue  
Savannah, GA 31404

Website: www.testamericainc.com  
Phone: (912) 354-7858  
Fax: (912) 352-0165

Alternate Laboratory Name/Location

Phone:  
Fax:

PROJECT REFERENCE <b>35th Ave Removal</b>	PROJECT NO. <b>2005148-1356</b>	PROJECT LOCATION (STATE) <b>AL</b>	MATRIX TYPE	REQUIRED ANALYSIS					PAGE <b>2</b> OF <b>2</b>
TAL (LAB) PROJECT MANAGER <i>Lee Harmon</i>	P.O. NUMBER	CONTRACT NO.	ENT FAX	STANDARD REPORT DELIVERY <input type="radio"/> DATE DUE _____ EXPEDITED REPORT DELIVERY (SURCHARGE) <input type="radio"/> DATE DUE _____ NUMBER OF COOLERS SUBMITTED PER SHIPMENT:					
(b) (6)									
COMPANY CONT.			COMPOSITE (C) OR GRAB (G) INDICATE	NONAQUEOUS LIQUID (OIL, SOLVENT, ...)	PRESERVATIVE				

DATE	TIME	SAMPLE IDENTIFICATION	COMPOSITE (C) OR GRAB (G)	WATER	SOLID OR SEMISOLID	AIR	NONAQUEOUS LIQUID (OIL, SOLVENT, ...)	NUMBER OF CONTAINERS SUBMITTED					REMARKS
								1	2	3	4	5	
3-25-13	1428	CN0613K-CSD	C	X			X	X					
	1332	CN0613AB-GS	G	X			X						
	1334	CN0613AC-GS	G	X			X						
	1440	CN0610A-CS	C	X			X						
	1442	CN0610B-CS	C	X			X						
	1439	CN0610AB-GS	G	X			X						
	1506	CN0506A-CS	C	X			X						
	1515	CN0506B-CS	C	X			X						
	1518	CN0014AB-GS	G	X			X						
	1343	CN0613E-CS (sieve)	C	X			X	X					
3-26-13	1300	032613-RB-Shovel		X			X	X					

RELINQUISHED BY: (SIGNATURE) <i>[Signature]</i>	DATE 3-26-13	TIME 1330	RELINQUISHED BY: (SIGNATURE)	DATE	TIME	RELINQUISHED BY: (SIGNATURE)	DATE	TIME
RECEIVED BY: (SIGNATURE)	DATE	TIME	RECEIVED BY: (SIGNATURE)	DATE	TIME	RECEIVED BY: (SIGNATURE)	DATE	TIME

LABORATORY USE ONLY								
RECEIVED FOR LABORATORY BY: (SIGNATURE) <i>[Signature]</i>	DATE 03/28/13	TIME 0937	CUSTODY INTACT YES <input type="radio"/> NO <input type="radio"/>	CUSTODY SEAL NO.	SAVANNAH LOG NO. <b>680-88766</b>	LABORATORY REMARKS <b>1-y<sup>c</sup></b>		

Page 14 of 18

4/8/2013





## Login Sample Receipt Checklist

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

SDG Number: 68088766-2

**Login Number: 88766**

**List Number: 1**

**Creator: Barnett, Eddie T**

**List Source: TestAmerica Savannah**

Question	Answer	Comment
Radioactivity wasn't checked or is <=/ background as measured by a survey meter.	N/A	
The cooler's custody seal, if present, is intact.	True	
Sample custody seals, if present, are intact.	True	
The cooler or samples do not appear to have been compromised or tampered with.	True	
Samples were received on ice.	True	
Cooler Temperature is acceptable.	True	
Cooler Temperature is recorded.	True	
COC is present.	True	
COC is filled out in ink and legible.	True	
COC is filled out with all pertinent information.	True	
Is the Field Sampler's name present on COC?	N/A	
There are no discrepancies between the containers received and the COC.	True	
Samples are received within Holding Time.	True	
Sample containers have leg ble labels.	True	
Containers are not broken or leaking.	True	
Sample collection date/times are provided.	True	
Appropriate sample containers are used.	True	
Sample bottles are completely filled.	True	
Sample Preservation Verified.	True	
There is sufficient vol. for all requested analyses, incl. any requested MS/MSDs	True	
Containers requiring zero headspace have no headspace or bubble is <6mm (1/4").	N/A	
Multiphasic samples are not present.	True	
Samples do not require splitting or compositing.	True	
Residual Chlorine Checked.	N/A	



## Login Sample Receipt Checklist

Client: Oneida Total Integrated Enterprises LLC

Job Number: 680-88766-2

SDG Number: 68088766-2

**Login Number: 88766**

**List Number: 1**

**Creator: McNulty, Carol**

**List Source: TestAmerica Tampa**

**List Creation: 03/29/13 09:17 AM**

Question	Answer	Comment
Radioactivity wasn't checked or is <=/ background as measured by a survey meter.	True	
The cooler's custody seal, if present, is intact.	True	
Sample custody seals, if present, are intact.	True	
The cooler or samples do not appear to have been compromised or tampered with.	True	
Samples were received on ice.	True	
Cooler Temperature is acceptable.	True	
Cooler Temperature is recorded.	True	
COC is present.	True	
COC is filled out in ink and legible.	True	
COC is filled out with all pertinent information.	True	
Is the Field Sampler's name present on COC?	True	
There are no discrepancies between the containers received and the COC.	True	
Samples are received within Holding Time.	True	
Sample containers have leg ble labels.	True	
Containers are not broken or leaking.	True	
Sample collection date/times are provided.	True	
Appropriate sample containers are used.	True	
Sample bottles are completely filled.	True	
Sample Preservation Verified.	True	
There is sufficient vol. for all requested analyses, incl. any requested MS/MSDs	True	
Containers requiring zero headspace have no headspace or bubble is <6mm (1/4").	True	
Multiphasic samples are not present.	True	
Samples do not require splitting or compositing.	True	
Residual Chlorine Checked.	True	



## Certification Summary

Client: Oneida Total Integrated Enterprises LLC  
 Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
 SDG: 68088766-2

### Laboratory: TestAmerica Savannah

All certifications held by this laboratory are listed. Not all certifications are applicable to this report.

Authority	Program	EPA Region	Certification ID	Expiration Date
A2LA	DoD ELAP		0399-01	05-31-13
Alabama	State Program	4	41450	06-30-13
Alaska (UST)	State Program	10	UST-104	06-19-13
California	NELAP	9	3217CA	07-31-13
Colorado	State Program	8	N/A	12-31-13
Florida	NELAP	4	E87052	06-30-13
GA Dept. of Agriculture	State Program	4	N/A	12-31-13
Georgia	State Program	4	N/A	06-30-13
Georgia	State Program	4	803	06-30-13
Guam	State Program	9	09-005r	04-17-13
Hawaii	State Program	9	N/A	06-30-13
Illinois	NELAP	5	200022	11-30-13
Indiana	State Program	5	N/A	06-30-13
Iowa	State Program	7	353	07-01-13
Kentucky	State Program	4	90084	12-31-12 *
Kentucky (UST)	State Program	4	18	03-31-13 *
Louisiana	NELAP	6	30690	06-30-13
Louisiana	NELAP	6	LA100015	12-31-13
Maine	State Program	1	GA00006	08-16-14
Maryland	State Program	3	250	12-31-13
Massachusetts	State Program	1	M-GA006	06-30-13
Michigan	State Program	5	9925	06-30-13
Mississippi	State Program	4	N/A	06-30-13
Montana	State Program	8	CERT0081	01-01-14
Nebraska	State Program	7	TestAmerica-Savannah	06-30-13
New Jersey	NELAP	2	GA769	06-30-13
New Mexico	State Program	6	N/A	06-30-13
North Carolina DENR	State Program	4	269	12-31-13
North Carolina DHHS	State Program	4	13701	07-31-13
Oklahoma	State Program	6	9984	08-31-13
Pennsylvania	NELAP	3	68-00474	06-30-13
Puerto Rico	State Program	2	GA00006	01-01-14
South Carolina	State Program	4	98001	06-30-13
Tennessee	State Program	4	TN02961	06-30-13
Texas	NELAP	6	T104704185-08-TX	11-30-13
USDA	Federal		SAV 3-04	04-07-14
Virginia	NELAP	3	460161	06-14-13
Washington	State Program	10	C1794	06-10-13
West Virginia	State Program	3	9950C	12-31-13
West Virginia DEP	State Program	3	94	06-30-13
Wisconsin	State Program	5	999819810	08-31-13
Wyoming	State Program	8	8TMS-Q	06-30-13

### Laboratory: TestAmerica Tampa

All certifications held by this laboratory are listed. Not all certifications are applicable to this report.

Authority	Program	EPA Region	Certification ID	Expiration Date
Alabama	State Program	4	40610	06-30-13
Florida	NELAP	4	E84282	06-30-13
Georgia	State Program	4	905	06-30-13

\* Expired certification is currently pending renewal and is considered valid.

TestAmerica Savannah

# Certification Summary

Client: Oneida Total Integrated Enterprises LLC  
Project/Site: 35th Avenue Superfund Site

TestAmerica Job ID: 680-88766-2  
SDG: 68088766-2

## Laboratory: TestAmerica Tampa (Continued)

All certifications held by this laboratory are listed. Not all certifications are applicable to this report.

Authority	Program	EPA Region	Certification ID	Expiration Date
USDA	Federal		P330-11-00177	04-20-14

- 1
- 2
- 3
- 4
- 5
- 6
- 7
- 8
- 9
- 10
- 11
- 12