

## **DATA PACKAGE**

SEMI-VOLATILE ORGANICS

**PROJECT NAME : MONROE, NJ**

**G ENVIRONMENTAL**

**8 Carriage Ln**

**Succasunna, NJ - 07876**

**Phone No: 973-294-1771**

**ORDER ID : Q3273**

**ATTENTION : Gary Landis**



**Laboratory Certification ID # 20012**



|                                       |    |
|---------------------------------------|----|
| 1) Signature Page                     | 3  |
| 2) Case Narrative                     | 5  |
| 2.1) SVOC-TCL BNA -20- Case Narrative | 5  |
| 3) Qualifier Page                     | 7  |
| 4) QA Checklist                       | 8  |
| 5) SVOC-TCL BNA -20 Data              | 9  |
| 6) Shipping Document                  | 94 |
| 6.1) CHAIN OF CUSTODY                 | 95 |
| 6.2) Lab Certificate                  | 96 |

|   |
|---|
| 1 |
| 2 |
| 3 |
| 4 |
| 5 |
| 6 |

## DATA OF KNOWN QUALITY CONFORMANCE/NON-CONFORMANCE SUMMARY QUESTIONNAIRE

Laboratory Name : Alliance Technical Group LLC Client : G Environmental  
 Project Location : NJ Project Number : - Monroe, NJ  
 Laboratory Sample ID(s) : Q3273 Sampling Date(s) : 10/01/2025  
 List DKQP Methods Used (e.g., 8260,8270, et Cetra) **8270E,SOP**

|    |                                                                                                                                                                                                                                                                                                     |                                                                                                                                                                             |
|----|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 1  | For each analytical method referenced in this laboratory report package, were all specified QA/QC performance criteria followed, including the requirement to explain any criteria falling outside of acceptable guidelines, as specified in the NJDEP Data of Known Quality performance standards? | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No                                                                                                         |
| 1A | Were the method specified handling, preservation, and holding time requirements met?                                                                                                                                                                                                                | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No                                                                                                         |
| 1B | EPH Method: Was the EPH method conducted without significant modifications (see Section 11.3 of respective DKQ methods)                                                                                                                                                                             | <input type="checkbox"/> Yes <input type="checkbox"/> No <input checked="" type="checkbox"/> N/A                                                                            |
| 2  | Were all samples received by the laboratory in a condition consistent with that described on the associated chain-of-custody document(s)?                                                                                                                                                           | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No                                                                                                         |
| 3  | Were samples received at an appropriate temperature (4±2° C)?                                                                                                                                                                                                                                       | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A                                                                            |
| 4  | Were all QA/QC performance criteria specified in the NJDEP DKQP standards achieved?                                                                                                                                                                                                                 | <input type="checkbox"/> Yes <input checked="" type="checkbox"/> No                                                                                                         |
| 5  | a)Were reporting limits specified or referenced on the chain-of-custody or communicated to the laboratory prior to sample receipt?<br><br>b)Were these reporting limits met?                                                                                                                        | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No<br><br><input checked="" type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A |
| 6  | For each analytical method referenced in this laboratory report package, were results reported for all constituents identified in the method-specific analyte lists presented in the DKQP documents and/or site-specific QAPP?                                                                      | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No                                                                                                         |
| 7  | Are project-specific matrix spikes and/or laboratory duplicates included in this data set?                                                                                                                                                                                                          | <input checked="" type="checkbox"/> Yes <input type="checkbox"/> No                                                                                                         |

Notes: For all questions to which the response was "No" (with the exception of question #7), additional information should be provided in an attached narrative. If the answer to question #1, #1A, or #1B is "No", the data package does not meet the requirements for "Data of Known Quality."

## Cover Page

**Order ID :** Q3273

**Project ID :** Monroe, NJ

**Client :** G Environmental

**Lab Sample Number**

Q3273-01

**Client Sample Number**

MW1

I certify that the data package is in compliance with the terms and conditions of the contract, both technically and for completeness, for other than the conditions detailed above. Release of the data contained in this hard copy data package has been authorized by the laboratory manager or his designee, as verified by the following signature.

Signature :

**APPROVED**

*By Nimisha Pandya, QA/QC Supervisor at 3:36 pm, Oct 10, 2025*

Date: 10/10/2025

NYDOH CERTIFICATION NO - 11376

NJDEP CERTIFICATION NO - 20012

## **CASE NARRATIVE**

### **G Environmental**

**Project Name: Monroe, NJ**

**Project # N/A**

**Order ID # Q3273**

**Test Name: SVOC-TCL BNA -20**

### **A. Number of Samples and Date of Receipt:**

1 Water sample was received on 10/02/2025.

### **B. Parameters**

According to the Chain of Custody document, the following analyses were requested: SVOC-TCL BNA -20. This data package contains results for SVOC-TCL BNA -20.

### **C. Analytical Techniques:**

The samples were analyzed on instrument BNA\_G using GC Column ZB-SemiVolatiles Guardian which is 30 meters, 0.25 mm ID, 0.5 um df, Catalog # 7HG-G027-17-GGA. The analysis of SVOC-TCL BNA -20 was based on method 8270E and extraction was done based on method 3510.

### **D. QA/ QC Samples:**

The Holding Times were met for all analysis.

The Surrogate recoveries were met for all analysis.

The Internal Standards Areas were met for all analysis.

The Retention Times were met for all analysis.

The RPD for {PB169973BSD} with File ID: BG064490.D met criteria except for 4-Chloroaniline[21%], due to difference in results of BS and BSD.

The Blank Spike for {PB169973BS} with File ID: BG064489.D met requirements for all compounds except for 3,3-Dichlorobenzidine[61%], 3-Nitroaniline[60%] and 4-Chloroaniline[38%], these compounds did not meet the NJDKQP criteria but met the in-house criteria.

The Blank Spike Duplicate for {PB169973BSD} with File ID: BG064490.D met requirements for all compounds except for 3,3-Dichlorobenzidine[55%], 3-Nitroaniline[52%] and 4-Chloroaniline[30%], these compounds did not meet the NJDKQP criteria but met the in-house criteria.

The Blank analysis did not indicate the presence of lab contamination.

The Initial Calibration met the requirements.

The Continuous Calibration met the requirements.

The Tuning criteria met requirements.



284 Sheffield Street, Mountainside, NJ 07092  
Phone: 908 789 8900 Fax: 908 789 8922

**E. Additional Comments:**

The Form 6 is not included in the data package because the Initial Calibration was performed using 7 points.

**F. Manual Integration Comments:**

Please refer to the Manual integration Report included with the Run Logs for information on the manual integrations performed.

---

I certify that the data package is in compliance with the terms and conditions of the contract, both technically and for completeness, for other than the conditions detailed above. The laboratory manager or his designee, as verified by the following signature has authorized release of the data contained in this hard copy data package.

**APPROVED**

Signature\_\_\_\_\_

*By Nimisha Pandya, QA/QC Supervisor at 3:36 pm, Oct 10, 2025*

## DATA REPORTING QUALIFIERS- ORGANIC

For reporting results, the following “ Results Qualifiers” are used:

|           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
|-----------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Value     | If the result is a value greater than or equal to the detection limit, report the value                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
| <b>U</b>  | Indicates the compound was analyzed for but was not detected. Report the minimum detection limit for the sample with the U, i.e. “10 U”. This is not necessarily the instrument detection limit attainable for this particular sample based on any concentration or dilution that may have been required.                                                                                                                                                                                                                                                                                      |
| <b>ND</b> | Indicates the analyte was analyzed for, but not detected                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| <b>J</b>  | Indicates an estimated value. This flag is used: <ol style="list-style-type: none"> <li>(1) When estimating a concentration for a tentatively identified compound (library search hits, where a 1:1 response is assumed.)</li> <li>(2) When the mass spectral data indicated the identification, however the result was less than the specified detection limit greater than zero. If the detection limit was 10ug/L and a concentration of 3 ug/L was calculated report as 3 J. This flag is used when similar situation arise on any organic parameter i.e. Pest, PCB and others.</li> </ol> |
| <b>B</b>  | Indicates the analyte was found in the blank as well as the sample report as “12 B”.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| <b>E</b>  | Indicates the analyte ‘s concentration exceeds the calibrated range of the instrument for that specific analysis.                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| <b>D</b>  | This flag identifies all compounds identified in an analysis at a secondary dilution factor.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| <b>P</b>  | This flag is used for Pesticide/PCB target analyte when there is >25% difference for detected concentrations between the two GC columns. The lower of the two values is reported on Form 1 and flagged with a “P”.                                                                                                                                                                                                                                                                                                                                                                             |
| <b>N</b>  | This flag indicates presumptive evidence of a compound. This is only used for tentatively identified compounds (TICs), where the identification is based on a mass spectral library search. It applies to all TIC results. For generic characterization of a TIC, such as chlorinated hydrocarbon, the flag is not used.                                                                                                                                                                                                                                                                       |
| <b>A</b>  | This flag indicates that a Tentatively Identified Compound is a suspected aldol-condensation product.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| <b>Q</b>  | Indicates the LCS did not meet the control limits requirements                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |

## APPENDIX A

### QA REVIEW GENERAL DOCUMENTATION

Project #: Q3273

Completed

For thorough review, the report must have the following:

#### GENERAL:

Are all original paperwork present (chain of custody, record of communication,airbill, sample management lab chronicle, login page)

✓

Check chain-of-custody for proper relinquish/return of samples

✓

Is the chain of custody signed and complete

✓

Check internal chain-of-custody for proper relinquish/return of samples /sample extracts

✓

Collect information for each project id from server. Were all requirements followed

✓

#### COVER PAGE:

Do numbers of samples correspond to the number of samples in the Chain of Custody on login page

✓

Do lab numbers and client Ids on cover page agree with the Chain of Custody

✓

#### CHAIN OF CUSTODY:

Do requested analyses on Chain of Custody agree with form I results

✓

Do requested analyses on Chain of Custody agree with the log-in page

✓

Were the correct method log-in for analysis according to the Analytical Request and Chain of Castody

✓

Were the samples received within hold time

✓

Were any problems found with the samples at arrival recorded in the Sample Management Laboratory Chronicle

✓

#### ANALYTICAL:

Was method requirement followed?

✓

Was client requirement followed?

✓

Does the case narrative summarize all QC failure?

✓

All runlogs and manual integration are reviewed for requirements

✓

All manual calculations and /or hand notations verified

✓

QA Review Signature: SOHIL JODHANI

Date: 10/10/2025



### Hit Summary Sheet SW-846

**SDG No.:** Q3273  
**Client:** G Environmental

| Sample ID              | Client ID | Parameter                                  | Concentration | C | MDL | RDL | Units |
|------------------------|-----------|--------------------------------------------|---------------|---|-----|-----|-------|
| <b>Client ID : MW1</b> |           |                                            |               |   |     |     |       |
| Q3273-01               | MW1       | WATER Pyrene                               | 4.700         | J | 0.5 | 5   | ug/L  |
|                        |           | <b>Total Svoc :</b>                        | <b>4.70</b>   |   |     |     |       |
| Q3273-01               | MW1       | WATER 1H-Indene, 2,3-dihydro-4,7-dimet *   | 10.600        | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER 5-Eicosene, (E)- *                   | 7.300         | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Decalin, syn-1-methyl-, cis- *       | 9.300         | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Heptadecane, 2,6,10,15-tetramethy *  | 21.300        | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Hexadecane, 2,6,10,14-tetramethy *   | 20.000        | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Naphthalene, 1,2,3,4-tetrahydro-1. * | 15.200        | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Naphthalene, 1,2,3,4-tetrahydro-5. * | 14.100        | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Naphthalene, 2,3,6-trimethyl- *      | 11.100        | J | 0   | 0   | ug/L  |
| Q3273-01               | MW1       | WATER Octadecane, 2,6-dimethyl- *          | 23.300        | J | 0   | 0   | ug/L  |
|                        |           | <b>Total Tics :</b>                        | <b>132.20</b> |   |     |     |       |
|                        |           | <b>Total Concentration:</b>                | <b>136.90</b> |   |     |     |       |



# SAMPLE DATA

A

B

C

D

E

F

G

H

I

J

K

### Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: | 10/01/25         |
| Project:           | Monroe, NJ       | Date Received:  | 10/02/25         |
| Client Sample ID:  | MW1              | SDG No.:        | Q3273            |
| Lab Sample ID:     | Q3273-01         | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064493.D        | 1         | 10/03/25 08:35 | 10/03/25 18:59 | PB169973      |

| CAS Number     | Parameter                   | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|----------------|-----------------------------|-------|-----------|------|------------|-------|
| <b>TARGETS</b> |                             |       |           |      |            |       |
| 100-52-7       | Benzaldehyde                | 3.90  | U         | 3.90 | 10.0       | ug/L  |
| 108-95-2       | Phenol                      | 0.91  | U         | 0.91 | 5.00       | ug/L  |
| 111-44-4       | bis(2-Chloroethyl)ether     | 0.81  | U         | 0.81 | 5.00       | ug/L  |
| 95-57-8        | 2-Chlorophenol              | 0.58  | U         | 0.58 | 5.00       | ug/L  |
| 95-48-7        | 2-Methylphenol              | 1.10  | U         | 1.10 | 5.00       | ug/L  |
| 108-60-1       | 2,2-oxybis(1-Chloropropane) | 1.30  | U         | 1.30 | 5.00       | ug/L  |
| 98-86-2        | Acetophenone                | 0.74  | U         | 0.74 | 5.00       | ug/L  |
| 65794-96-9     | 3+4-Methylphenols           | 1.10  | U         | 1.10 | 10.0       | ug/L  |
| 621-64-7       | n-Nitroso-di-n-propylamine  | 1.40  | U         | 1.40 | 2.50       | ug/L  |
| 67-72-1        | Hexachloroethane            | 0.65  | U         | 0.65 | 5.00       | ug/L  |
| 98-95-3        | Nitrobenzene                | 0.76  | U         | 0.76 | 5.00       | ug/L  |
| 78-59-1        | Isophorone                  | 0.75  | U         | 0.75 | 5.00       | ug/L  |
| 88-75-5        | 2-Nitrophenol               | 1.80  | U         | 1.80 | 5.00       | ug/L  |
| 105-67-9       | 2,4-Dimethylphenol          | 1.90  | U         | 1.90 | 5.00       | ug/L  |
| 111-91-1       | bis(2-Chloroethoxy)methane  | 0.68  | U         | 0.68 | 5.00       | ug/L  |
| 120-83-2       | 2,4-Dichlorophenol          | 0.52  | U         | 0.52 | 5.00       | ug/L  |
| 91-20-3        | Naphthalene                 | 0.50  | U         | 0.50 | 5.00       | ug/L  |
| 106-47-8       | 4-Chloroaniline             | 0.84  | UQ        | 0.84 | 5.00       | ug/L  |
| 87-68-3        | Hexachlorobutadiene         | 0.54  | U         | 0.54 | 5.00       | ug/L  |
| 105-60-2       | Caprolactam                 | 1.10  | U         | 1.10 | 10.0       | ug/L  |
| 59-50-7        | 4-Chloro-3-methylphenol     | 0.59  | U         | 0.59 | 5.00       | ug/L  |
| 91-57-6        | 2-Methylnaphthalene         | 0.56  | U         | 0.56 | 5.00       | ug/L  |
| 77-47-4        | Hexachlorocyclopentadiene   | 3.60  | U         | 3.60 | 10.0       | ug/L  |
| 88-06-2        | 2,4,6-Trichlorophenol       | 0.51  | U         | 0.51 | 5.00       | ug/L  |
| 95-95-4        | 2,4,5-Trichlorophenol       | 0.62  | U         | 0.62 | 5.00       | ug/L  |
| 92-52-4        | 1,1-Biphenyl                | 0.53  | U         | 0.53 | 5.00       | ug/L  |
| 91-58-7        | 2-Chloronaphthalene         | 0.61  | U         | 0.61 | 5.00       | ug/L  |
| 88-74-4        | 2-Nitroaniline              | 1.30  | U         | 1.30 | 5.00       | ug/L  |
| 131-11-3       | Dimethylphthalate           | 0.61  | U         | 0.61 | 5.00       | ug/L  |

## Report of Analysis

|                    |                        |                 |                  |
|--------------------|------------------------|-----------------|------------------|
| Client:            | G Environmental        | Date Collected: | 10/01/25         |
| Project:           | Monroe, NJ             | Date Received:  | 10/02/25         |
| Client Sample ID:  | MW1                    | SDG No.:        | Q3273            |
| Lab Sample ID:     | Q3273-01               | Matrix:         | Water            |
| Analytical Method: | 8270E                  | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000      Units:    mL | Final Vol:      | 1000      uL     |
| Soil Aliquot Vol:  | uL                     | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted :    N        | Level :         | LOW              |
| Injection Volume : | GPC Factor :    1.0    | GPC Cleanup :   | N      PH :      |
| Prep Method :      |                        |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064493.D        | 1         | 10/03/25 08:35 | 10/03/25 18:59 | PB169973      |

| CAS Number | Parameter                  | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|------------|----------------------------|-------|-----------|------|------------|-------|
| 208-96-8   | Acenaphthylene             | 0.75  | U         | 0.75 | 5.00       | ug/L  |
| 606-20-2   | 2,6-Dinitrotoluene         | 0.92  | U         | 0.92 | 5.00       | ug/L  |
| 99-09-2    | 3-Nitroaniline             | 1.10  | UQ        | 1.10 | 5.00       | ug/L  |
| 83-32-9    | Acenaphthene               | 0.55  | U         | 0.55 | 5.00       | ug/L  |
| 51-28-5    | 2,4-Dinitrophenol          | 6.00  | U         | 6.00 | 10.0       | ug/L  |
| 100-02-7   | 4-Nitrophenol              | 2.40  | U         | 2.40 | 10.0       | ug/L  |
| 132-64-9   | Dibenzofuran               | 0.61  | U         | 0.61 | 5.00       | ug/L  |
| 121-14-2   | 2,4-Dinitrotoluene         | 1.20  | U         | 1.20 | 5.00       | ug/L  |
| 84-66-2    | Diethylphthalate           | 0.69  | U         | 0.69 | 5.00       | ug/L  |
| 7005-72-3  | 4-Chlorophenyl-phenylether | 0.68  | U         | 0.68 | 5.00       | ug/L  |
| 86-73-7    | Fluorene                   | 0.63  | U         | 0.63 | 5.00       | ug/L  |
| 100-01-6   | 4-Nitroaniline             | 1.50  | U         | 1.50 | 5.00       | ug/L  |
| 534-52-1   | 4,6-Dinitro-2-methylphenol | 2.90  | U         | 2.90 | 10.0       | ug/L  |
| 86-30-6    | n-Nitrosodiphenylamine     | 0.58  | U         | 0.58 | 5.00       | ug/L  |
| 101-55-3   | 4-Bromophenyl-phenylether  | 0.40  | U         | 0.40 | 5.00       | ug/L  |
| 118-74-1   | Hexachlorobenzene          | 0.52  | U         | 0.52 | 5.00       | ug/L  |
| 1912-24-9  | Atrazine                   | 1.00  | U         | 1.00 | 5.00       | ug/L  |
| 87-86-5    | Pentachlorophenol          | 1.60  | U         | 1.60 | 10.0       | ug/L  |
| 85-01-8    | Phenanthrene               | 0.50  | U         | 0.50 | 5.00       | ug/L  |
| 120-12-7   | Anthracene                 | 0.61  | U         | 0.61 | 5.00       | ug/L  |
| 86-74-8    | Carbazole                  | 0.72  | U         | 0.72 | 5.00       | ug/L  |
| 84-74-2    | Di-n-butylphthalate        | 1.20  | U         | 1.20 | 5.00       | ug/L  |
| 206-44-0   | Fluoranthene               | 0.82  | U         | 0.82 | 5.00       | ug/L  |
| 129-00-0   | Pyrene                     | 4.70  | J         | 0.50 | 5.00       | ug/L  |
| 85-68-7    | Butylbenzylphthalate       | 1.90  | U         | 1.90 | 5.00       | ug/L  |
| 91-94-1    | 3,3-Dichlorobenzidine      | 0.93  | UQ        | 0.93 | 10.0       | ug/L  |
| 56-55-3    | Benzo(a)anthracene         | 0.45  | U         | 0.45 | 5.00       | ug/L  |
| 218-01-9   | Chrysene                   | 0.44  | U         | 0.44 | 5.00       | ug/L  |
| 117-81-7   | Bis(2-ethylhexyl)phthalate | 1.60  | U         | 1.60 | 5.00       | ug/L  |
| 117-84-0   | Di-n-octyl phthalate       | 2.30  | U         | 2.30 | 10.0       | ug/L  |
| 205-99-2   | Benzo(b)fluoranthene       | 0.49  | U         | 0.49 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: | 10/01/25         |
| Project:           | Monroe, NJ       | Date Received:  | 10/02/25         |
| Client Sample ID:  | MW1              | SDG No.:        | Q3273            |
| Lab Sample ID:     | Q3273-01         | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064493.D        | 1         | 10/03/25 08:35 | 10/03/25 18:59 | PB169973      |

| CAS Number                            | Parameter                          | Conc.  | Qualifier | MDL                 | LOQ / CRQL | Units    |
|---------------------------------------|------------------------------------|--------|-----------|---------------------|------------|----------|
| 207-08-9                              | Benzo(k)fluoranthene               | 0.48   | U         | 0.48                | 5.00       | ug/L     |
| 50-32-8                               | Benzo(a)pyrene                     | 0.55   | U         | 0.55                | 5.00       | ug/L     |
| 193-39-5                              | Indeno(1,2,3-cd)pyrene             | 0.59   | U         | 0.59                | 5.00       | ug/L     |
| 53-70-3                               | Dibenzo(a,h)anthracene             | 0.67   | U         | 0.67                | 5.00       | ug/L     |
| 191-24-2                              | Benzo(g,h,i)perylene               | 0.69   | U         | 0.69                | 5.00       | ug/L     |
| 95-94-3                               | 1,2,4,5-Tetrachlorobenzene         | 0.52   | U         | 0.52                | 5.00       | ug/L     |
| 123-91-1                              | 1,4-Dioxane                        | 1.00   | U         | 1.00                | 5.00       | ug/L     |
| 58-90-2                               | 2,3,4,6-Tetrachlorophenol          | 0.72   | U         | 0.72                | 5.00       | ug/L     |
| <b>SURROGATES</b>                     |                                    |        |           |                     |            |          |
| 367-12-4                              | 2-Fluorophenol                     | 68.2   |           | 15 (23) - 110 (138) | 45%        | SPK: 150 |
| 13127-88-3                            | Phenol-d6                          | 40.6   |           | 15 (10) - 110 (134) | 27%        | SPK: 150 |
| 4165-60-0                             | Nitrobenzene-d5                    | 90.5   |           | 30 (67) - 130 (132) | 91%        | SPK: 100 |
| 321-60-8                              | 2-Fluorobiphenyl                   | 91.7   |           | 30 (52) - 130 (132) | 92%        | SPK: 100 |
| 118-79-6                              | 2,4,6-Tribromophenol               | 134    |           | 15 (44) - 110 (137) | 89%        | SPK: 150 |
| 1718-51-0                             | Terphenyl-d14                      | 86.3   |           | 30 (42) - 130 (152) | 86%        | SPK: 100 |
| <b>INTERNAL STANDARDS</b>             |                                    |        |           |                     |            |          |
| 3855-82-1                             | 1,4-Dichlorobenzene-d4             | 20100  | 7.82      |                     |            |          |
| 1146-65-2                             | Naphthalene-d8                     | 82400  | 10.611    |                     |            |          |
| 15067-26-2                            | Acenaphthene-d10                   | 60800  | 14.459    |                     |            |          |
| 1517-22-2                             | Phenanthrene-d10                   | 130000 | 17.197    |                     |            |          |
| 1719-03-5                             | Chrysene-d12                       | 137000 | 21.416    |                     |            |          |
| 1520-96-3                             | Perylene-d12                       | 150000 | 24.389    |                     |            |          |
| <b>TENTATIVE IDENTIFIED COMPOUNDS</b> |                                    |        |           |                     |            |          |
| 1000158-89-1                          | Decalin, syn-1-methyl-, cis-       | 9.30   | J         |                     | 9.81       | ug/L     |
| 006682-71-9                           | 1H-Indene, 2,3-dihydro-4,7-dimethy | 10.6   | J         |                     | 11.5       | ug/L     |
| 004175-54-6                           | Naphthalene, 1,2,3,4-tetrahydro-1, | 15.2   | J         |                     | 12.5       | ug/L     |
| 020027-77-4                           | Naphthalene, 1,2,3,4-tetrahydro-5, | 14.1   | J         |                     | 13.4       | ug/L     |
| 054833-48-6                           | Heptadecane, 2,6,10,15-tetramethyl | 21.3   | J         |                     | 14.0       | ug/L     |
| 000829-26-5                           | Naphthalene, 2,3,6-trimethyl-      | 11.1   | J         |                     | 15.1       | ug/L     |
| 075163-97-2                           | Octadecane, 2,6-dimethyl-          | 23.3   | J         |                     | 16.2       | ug/L     |

## Report of Analysis

|                    |                        |                 |                              |
|--------------------|------------------------|-----------------|------------------------------|
| Client:            | G Environmental        | Date Collected: | 10/01/25                     |
| Project:           | Monroe, NJ             | Date Received:  | 10/02/25                     |
| Client Sample ID:  | MW1                    | SDG No.:        | Q3273                        |
| Lab Sample ID:     | Q3273-01               | Matrix:         | Water                        |
| Analytical Method: | 8270E                  | % Solid:        | 0                            |
| Sample Wt/Vol:     | 1000      Units:    mL | Final Vol:      | 1000                      uL |
| Soil Aliquot Vol:  | uL                     | Test:           | SVOC-TCL BNA -20             |
| Extraction Type :  | Decanted :      N      | Level :         | LOW                          |
| Injection Volume : | GPC Factor :    1.0    | GPC Cleanup :   | N                      PH :  |
| Prep Method :      |                        |                 |                              |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064493.D        | 1         | 10/03/25 08:35 | 10/03/25 18:59 | PB169973      |

| CAS Number  | Parameter                          | Conc. | Qualifier | MDL | LOQ / CRQL | Units |
|-------------|------------------------------------|-------|-----------|-----|------------|-------|
| 000638-36-8 | Hexadecane, 2,6,10,14-tetramethyl- | 20.0  | J         |     | 17.0       | ug/L  |
| 074685-30-6 | 5-Eicosene, (E)-                   | 7.30  | J         |     | 21.1       | ug/L  |

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

\* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products



# QC SUMMARY

## Surrogate Summary

SW-846

SDG No.: Q3273

Client: G Environmental

Analytical Method: 8270E

| Lab Sample ID | Client ID   | Parameter            | Spike (PPM) | Result (PPM) | Recovery (%) | Qual | Limits (%) |           |
|---------------|-------------|----------------------|-------------|--------------|--------------|------|------------|-----------|
|               |             |                      |             |              |              |      | Low        | High      |
| PB169973BL    | PB169973BL  | 2-Fluorophenol       | 150         | 162          | 108          |      | 15 (23)    | 110 (138) |
|               |             | Phenol-d6            | 150         | 150          | 100          |      | 15 (10)    | 110 (134) |
|               |             | Nitrobenzene-d5      | 100         | 94.0         | 94           |      | 30 (67)    | 130 (132) |
|               |             | 2-Fluorobiphenyl     | 100         | 90.7         | 91           |      | 30 (52)    | 130 (132) |
|               |             | 2,4,6-Tribromophenol | 150         | 138          | 92           |      | 15 (44)    | 110 (137) |
|               |             | Terphenyl-d14        | 100         | 98.8         | 99           |      | 30 (42)    | 130 (152) |
| PB169973BS    | PB169973BS  | 2-Fluorophenol       | 150         | 157          | 105          |      | 15 (23)    | 110 (138) |
|               |             | Phenol-d6            | 150         | 153          | 102          |      | 15 (10)    | 110 (134) |
|               |             | Nitrobenzene-d5      | 100         | 94.3         | 94           |      | 30 (67)    | 130 (132) |
|               |             | 2-Fluorobiphenyl     | 100         | 91.8         | 92           |      | 30 (52)    | 130 (132) |
|               |             | 2,4,6-Tribromophenol | 150         | 149          | 99           |      | 15 (44)    | 110 (137) |
|               |             | Terphenyl-d14        | 100         | 97.3         | 97           |      | 30 (42)    | 130 (152) |
| PB169973BSD   | PB169973BSD | 2-Fluorophenol       | 150         | 163          | 108          |      | 15 (23)    | 110 (138) |
|               |             | Phenol-d6            | 150         | 155          | 103          |      | 15 (10)    | 110 (134) |
|               |             | Nitrobenzene-d5      | 100         | 91.5         | 91           |      | 30 (67)    | 130 (132) |
|               |             | 2-Fluorobiphenyl     | 100         | 90.3         | 90           |      | 30 (52)    | 130 (132) |
|               |             | 2,4,6-Tribromophenol | 150         | 144          | 96           |      | 15 (44)    | 110 (137) |
|               |             | Terphenyl-d14        | 100         | 96.5         | 97           |      | 30 (42)    | 130 (152) |
| Q3273-01      | MW1         | 2-Fluorophenol       | 150         | 68.2         | 45           |      | 15 (23)    | 110 (138) |
|               |             | Phenol-d6            | 150         | 40.6         | 27           |      | 15 (10)    | 110 (134) |
|               |             | Nitrobenzene-d5      | 100         | 90.5         | 91           |      | 30 (67)    | 130 (132) |
|               |             | 2-Fluorobiphenyl     | 100         | 91.7         | 92           |      | 30 (52)    | 130 (132) |
|               |             | 2,4,6-Tribromophenol | 150         | 134          | 89           |      | 15 (44)    | 110 (137) |
|               |             | Terphenyl-d14        | 100         | 86.3         | 86           |      | 30 (42)    | 130 (152) |

() = LABORATORY INHOUSE LIMIT



# Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

|          |                        |                    |                   |
|----------|------------------------|--------------------|-------------------|
| SDG No.: | <u>Q3273</u>           | Analytical Method: | <u>8270E</u>      |
| Client:  | <u>G Environmental</u> | DataFile:          | <u>BG064489.D</u> |

| Lab Sample ID | Parameter                   | Spike | Result | Unit | Rec | RPD | Qual | RPD  | Limits  |           | RPD |
|---------------|-----------------------------|-------|--------|------|-----|-----|------|------|---------|-----------|-----|
|               |                             |       |        |      |     |     |      | Qual | Low     | High      |     |
| PB169973BS    | Benzaldehyde                | 50    | 26.5   | ug/L | 53  |     |      |      | 20 (10) | 160 (162) |     |
|               | Phenol                      | 50    | 53.3   | ug/L | 107 |     |      |      | 20 (66) | 160 (118) |     |
|               | bis(2-Chloroethyl)ether     | 50    | 49.4   | ug/L | 99  |     |      |      | 70 (62) | 130 (103) |     |
|               | 2-Chlorophenol              | 50    | 51.6   | ug/L | 103 |     |      |      | 70 (70) | 130 (117) |     |
|               | 2-Methylphenol              | 50    | 52.4   | ug/L | 105 |     |      |      | 70 (69) | 130 (109) |     |
|               | 2,2-oxybis(1-Chloropropane) | 50    | 49.5   | ug/L | 99  |     |      |      | 70 (65) | 130 (100) |     |
|               | Acetophenone                | 50    | 57.0   | ug/L | 114 |     |      |      | 70 (60) | 130 (104) |     |
|               | 3+4-Methylphenols           | 50    | 53.3   | ug/L | 107 |     |      |      | 20 (67) | 160 (106) |     |
|               | N-Nitroso-di-n-propylamine  | 50    | 51.0   | ug/L | 102 |     |      |      | 70 (57) | 130 (107) |     |
|               | Hexachloroethane            | 50    | 49.2   | ug/L | 98  |     |      |      | 20 (76) | 160 (118) |     |
|               | Nitrobenzene                | 50    | 50.7   | ug/L | 101 |     |      |      | 70 (58) | 130 (106) |     |
|               | Isophorone                  | 50    | 48.4   | ug/L | 97  |     |      |      | 70 (61) | 130 (102) |     |
|               | 2-Nitrophenol               | 50    | 49.1   | ug/L | 98  |     |      |      | 70 (70) | 130 (115) |     |
|               | 2,4-Dimethylphenol          | 50    | 58.0   | ug/L | 116 |     |      |      | 70 (42) | 130 (142) |     |
|               | bis(2-Chloroethoxy)methane  | 50    | 50.5   | ug/L | 101 |     |      |      | 70 (58) | 130 (109) |     |
|               | 2,4-Dichlorophenol          | 50    | 55.8   | ug/L | 112 |     |      |      | 70 (66) | 130 (115) |     |
|               | Naphthalene                 | 50    | 50.7   | ug/L | 101 |     |      |      | 70 (64) | 130 (107) |     |
|               | 4-Chloroaniline             | 50    | 18.8   | ug/L | 38  |     | *    |      | 70 (10) | 130 (85)  |     |
|               | Hexachlorobutadiene         | 50    | 49.1   | ug/L | 98  |     |      |      | 70 (69) | 130 (101) |     |
|               | Caprolactam                 | 50    | 52.0   | ug/L | 104 |     |      |      | 20 (58) | 160 (128) |     |
|               | 4-Chloro-3-methylphenol     | 50    | 53.9   | ug/L | 108 |     |      |      | 70 (65) | 130 (114) |     |
|               | 2-Methylnaphthalene         | 50    | 46.2   | ug/L | 92  |     |      |      | 70 (64) | 130 (107) |     |
|               | Hexachlorocyclopentadiene   | 100   | 150    | ug/L | 150 |     |      |      | 20 (36) | 160 (160) |     |
|               | 2,4,6-Trichlorophenol       | 50    | 53.5   | ug/L | 107 |     |      |      | 70 (61) | 130 (110) |     |
|               | 2,4,5-Trichlorophenol       | 50    | 54.4   | ug/L | 109 |     |      |      | 70 (70) | 130 (106) |     |
|               | 1,1-Biphenyl                | 50    | 56.4   | ug/L | 113 |     |      |      | 70 (72) | 130 (98)  |     |
|               | 2-Chloronaphthalene         | 50    | 50.1   | ug/L | 100 |     |      |      | 70 (59) | 130 (106) |     |
|               | 2-Nitroaniline              | 50    | 53.0   | ug/L | 106 |     |      |      | 70 (73) | 130 (114) |     |
|               | Dimethylphthalate           | 50    | 52.1   | ug/L | 104 |     |      |      | 70 (64) | 130 (103) |     |
|               | Acenaphthylene              | 50    | 57.9   | ug/L | 116 |     |      |      | 70 (79) | 130 (103) |     |
|               | 2,6-Dinitrotoluene          | 50    | 53.0   | ug/L | 106 |     |      |      | 70 (64) | 130 (110) |     |
|               | 3-Nitroaniline              | 50    | 30.0   | ug/L | 60  |     | *    |      | 70 (28) | 130 (100) |     |
|               | Acenaphthene                | 50    | 49.8   | ug/L | 100 |     |      |      | 70 (59) | 130 (113) |     |
|               | 2,4-Dinitrophenol           | 100   | 95.5   | ug/L | 96  |     |      |      | 20 (36) | 160 (166) |     |
|               | 4-Nitrophenol               | 100   | 110    | ug/L | 110 |     |      |      | 20 (45) | 160 (147) |     |
|               | Dibenzofuran                | 50    | 51.4   | ug/L | 103 |     |      |      | 70 (65) | 130 (106) |     |
|               | 2,4-Dinitrotoluene          | 50    | 52.2   | ug/L | 104 |     |      |      | 70 (60) | 130 (115) |     |
|               | Diethylphthalate            | 50    | 51.8   | ug/L | 104 |     |      |      | 70 (63) | 130 (105) |     |
|               | 4-Chlorophenyl-phenylether  | 50    | 50.4   | ug/L | 101 |     |      |      | 70 (61) | 130 (104) |     |
|               | Fluorene                    | 50    | 52.1   | ug/L | 104 |     |      |      | 70 (64) | 130 (107) |     |
|               | 4-Nitroaniline              | 50    | 53.5   | ug/L | 107 |     |      |      | 70 (55) | 130 (125) |     |
|               | 4,6-Dinitro-2-methylphenol  | 50    | 51.5   | ug/L | 103 |     |      |      | 70 (62) | 130 (132) |     |
|               | N-Nitrosodiphenylamine      | 50    | 52.3   | ug/L | 105 |     |      |      | 70 (61) | 130 (109) |     |
|               | 4-Bromophenyl-phenylether   | 50    | 51.0   | ug/L | 102 |     |      |      | 70 (73) | 130 (103) |     |

() = LABORATORY INHOUSE LIMIT

## Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

|                                       |                                        |
|---------------------------------------|----------------------------------------|
| <b>SDG No.:</b> <u>Q3273</u>          | <b>Analytical Method:</b> <u>8270E</u> |
| <b>Client:</b> <u>G Environmental</u> | <b>DataFile:</b> <u>BG064489.D</u>     |

| Lab Sample ID | Parameter                  | Spike | Result | Unit | Rec | RPD | Qual | RPD  | Limits  |           | RPD |
|---------------|----------------------------|-------|--------|------|-----|-----|------|------|---------|-----------|-----|
|               |                            |       |        |      |     |     |      | Qual | Low     | High      |     |
| PB169973BS    | Hexachlorobenzene          | 50    | 50.8   | ug/L | 102 |     |      |      | 70 (73) | 130 (106) |     |
|               | Atrazine                   | 50    | 55.0   | ug/L | 110 |     |      |      | 70 (76) | 130 (120) |     |
|               | Pentachlorophenol          | 100   | 110    | ug/L | 110 |     |      |      | 20 (47) | 160 (114) |     |
|               | Phenanthrene               | 50    | 51.6   | ug/L | 103 |     |      |      | 70 (62) | 130 (109) |     |
|               | Anthracene                 | 50    | 53.0   | ug/L | 106 |     |      |      | 70 (65) | 130 (110) |     |
|               | Carbazole                  | 50    | 53.2   | ug/L | 106 |     |      |      | 70 (62) | 130 (106) |     |
|               | Di-n-butylphthalate        | 50    | 52.6   | ug/L | 105 |     |      |      | 70 (64) | 130 (106) |     |
|               | Fluoranthene               | 50    | 52.1   | ug/L | 104 |     |      |      | 70 (64) | 130 (110) |     |
|               | Pyrene                     | 50    | 51.7   | ug/L | 103 |     |      |      | 70 (71) | 130 (103) |     |
|               | Butylbenzylphthalate       | 50    | 52.8   | ug/L | 106 |     |      |      | 70 (61) | 130 (105) |     |
|               | 3,3-Dichlorobenzidine      | 50    | 30.6   | ug/L | 61  |     | *    |      | 70 (43) | 130 (108) |     |
|               | Benzo(a)anthracene         | 50    | 51.5   | ug/L | 103 |     |      |      | 70 (62) | 130 (107) |     |
|               | Chrysene                   | 50    | 51.3   | ug/L | 103 |     |      |      | 70 (61) | 130 (108) |     |
|               | bis(2-Ethylhexyl)phthalate | 50    | 52.7   | ug/L | 105 |     |      |      | 70 (59) | 130 (110) |     |
|               | Di-n-octyl phthalate       | 50    | 52.0   | ug/L | 104 |     |      |      | 70 (52) | 130 (139) |     |
|               | Benzo(b)fluoranthene       | 50    | 54.2   | ug/L | 108 |     |      |      | 70 (77) | 130 (113) |     |
|               | Benzo(k)fluoranthene       | 50    | 51.8   | ug/L | 104 |     |      |      | 70 (77) | 130 (105) |     |
|               | Benzo(a)pyrene             | 50    | 54.9   | ug/L | 110 |     |      |      | 70 (72) | 130 (131) |     |
|               | Indeno(1,2,3-cd)pyrene     | 50    | 53.9   | ug/L | 108 |     |      |      | 70 (72) | 130 (105) |     |
|               | Dibenz(a,h)anthracene      | 50    | 55.2   | ug/L | 110 |     |      |      | 70 (78) | 130 (115) |     |
|               | Benzo(g,h,i)perylene       | 50    | 54.7   | ug/L | 109 |     |      |      | 70 (75) | 130 (118) |     |
|               | 1,2,4,5-Tetrachlorobenzene | 50    | 54.2   | ug/L | 108 |     |      |      | 70 (72) | 130 (101) |     |
|               | 1,4-Dioxane                | 50    | 37.9   | ug/L | 76  |     |      |      | 20 (38) | 160 (125) |     |
|               | 2,3,4,6-Tetrachlorophenol  | 50    | 54.7   | ug/L | 109 |     |      |      | 70 (63) | 130 (116) |     |

() = LABORATORY INHOUSE LIMIT

# Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q3273

Analytical Method:

8270E

Client: G Environmental

DataFile:

BG064490.D

| Lab Sample ID | Parameter                   | Spike | Result | Unit | Rec | RPD | Qual | RPD  |  | Limits  |           | RPD     |
|---------------|-----------------------------|-------|--------|------|-----|-----|------|------|--|---------|-----------|---------|
|               |                             |       |        |      |     |     |      | Qual |  | Low     | High      |         |
| PB169973BSD   | Benzaldehyde                | 50    | 26.1   | ug/L | 52  | 2   |      |      |  | 20 (10) | 160 (162) | 20 (20) |
|               | Phenol                      | 50    | 55.3   | ug/L | 111 | 4   |      |      |  | 20 (66) | 160 (118) | 20 (20) |
|               | bis(2-Chloroethyl)ether     | 50    | 49.7   | ug/L | 99  | 1   |      |      |  | 70 (62) | 130 (103) | 20 (20) |
|               | 2-Chlorophenol              | 50    | 55.2   | ug/L | 110 | 7   |      |      |  | 70 (70) | 130 (117) | 20 (20) |
|               | 2-Methylphenol              | 50    | 55.0   | ug/L | 110 | 5   |      |      |  | 70 (69) | 130 (109) | 20 (20) |
|               | 2,2-oxybis(1-Chloropropane) | 50    | 48.3   | ug/L | 97  | 2   |      |      |  | 70 (65) | 130 (100) | 20 (20) |
|               | Acetophenone                | 50    | 54.9   | ug/L | 110 | 4   |      |      |  | 70 (60) | 130 (104) | 20 (20) |
|               | 3+4-Methylphenols           | 50    | 55.3   | ug/L | 111 | 4   |      |      |  | 20 (67) | 160 (106) | 20 (20) |
|               | N-Nitroso-di-n-propylamine  | 50    | 52.4   | ug/L | 105 | 3   |      |      |  | 70 (57) | 130 (107) | 20 (20) |
|               | Hexachloroethane            | 50    | 49.3   | ug/L | 99  | 0   |      |      |  | 20 (76) | 160 (118) | 20 (20) |
|               | Nitrobenzene                | 50    | 47.6   | ug/L | 95  | 6   |      |      |  | 70 (58) | 130 (106) | 20 (20) |
|               | Isophorone                  | 50    | 47.7   | ug/L | 95  | 1   |      |      |  | 70 (61) | 130 (102) | 20 (20) |
|               | 2-Nitrophenol               | 50    | 48.4   | ug/L | 97  | 1   |      |      |  | 70 (70) | 130 (115) | 20 (20) |
|               | 2,4-Dimethylphenol          | 50    | 57.0   | ug/L | 114 | 2   |      |      |  | 70 (42) | 130 (142) | 20 (20) |
|               | bis(2-Chloroethoxy)methane  | 50    | 49.3   | ug/L | 99  | 2   |      |      |  | 70 (58) | 130 (109) | 20 (20) |
|               | 2,4-Dichlorophenol          | 50    | 52.6   | ug/L | 105 | 6   |      |      |  | 70 (66) | 130 (115) | 20 (20) |
|               | Naphthalene                 | 50    | 48.5   | ug/L | 97  | 4   |      |      |  | 70 (64) | 130 (107) | 20 (20) |
|               | 4-Chloroaniline             | 50    | 15.2   | ug/L | 30  | 21  | *    | *    |  | 70 (10) | 130 (85)  | 20 (20) |
|               | Hexachlorobutadiene         | 50    | 47.0   | ug/L | 94  | 4   |      |      |  | 70 (69) | 130 (101) | 20 (20) |
|               | Caprolactam                 | 50    | 50.1   | ug/L | 100 | 4   |      |      |  | 20 (58) | 160 (128) | 20 (20) |
|               | 4-Chloro-3-methylphenol     | 50    | 51.6   | ug/L | 103 | 4   |      |      |  | 70 (65) | 130 (114) | 20 (20) |
|               | 2-Methylnaphthalene         | 50    | 44.5   | ug/L | 89  | 4   |      |      |  | 70 (64) | 130 (107) | 20 (20) |
|               | Hexachlorocyclopentadiene   | 100   | 140    | ug/L | 140 | 7   |      |      |  | 20 (36) | 160 (160) | 20 (20) |
|               | 2,4,6-Trichlorophenol       | 50    | 51.9   | ug/L | 104 | 3   |      |      |  | 70 (61) | 130 (110) | 20 (20) |
|               | 2,4,5-Trichlorophenol       | 50    | 49.0   | ug/L | 98  | 10  |      |      |  | 70 (70) | 130 (106) | 20 (20) |
|               | 1,1-Biphenyl                | 50    | 53.2   | ug/L | 106 | 6   |      |      |  | 70 (72) | 130 (98)  | 20 (20) |
|               | 2-Chloronaphthalene         | 50    | 47.4   | ug/L | 95  | 6   |      |      |  | 70 (59) | 130 (106) | 20 (20) |
|               | 2-Nitroaniline              | 50    | 50.6   | ug/L | 101 | 5   |      |      |  | 70 (73) | 130 (114) | 20 (20) |
|               | Dimethylphthalate           | 50    | 48.6   | ug/L | 97  | 7   |      |      |  | 70 (64) | 130 (103) | 20 (20) |
|               | Acenaphthylene              | 50    | 55.5   | ug/L | 111 | 4   |      |      |  | 70 (79) | 130 (103) | 20 (20) |
|               | 2,6-Dinitrotoluene          | 50    | 50.2   | ug/L | 100 | 5   |      |      |  | 70 (64) | 130 (110) | 20 (20) |
|               | 3-Nitroaniline              | 50    | 26.1   | ug/L | 52  | 14  | *    |      |  | 70 (28) | 130 (100) | 20 (20) |
|               | Acenaphthene                | 50    | 47.3   | ug/L | 95  | 5   |      |      |  | 70 (59) | 130 (113) | 20 (20) |
|               | 2,4-Dinitrophenol           | 100   | 96.1   | ug/L | 96  | 1   |      |      |  | 20 (36) | 160 (166) | 20 (20) |
|               | 4-Nitrophenol               | 100   | 110    | ug/L | 110 | 0   |      |      |  | 20 (45) | 160 (147) | 20 (20) |
|               | Dibenzofuran                | 50    | 49.0   | ug/L | 98  | 5   |      |      |  | 70 (65) | 130 (106) | 20 (20) |
|               | 2,4-Dinitrotoluene          | 50    | 53.2   | ug/L | 106 | 2   |      |      |  | 70 (60) | 130 (115) | 20 (20) |
|               | Diethylphthalate            | 50    | 49.5   | ug/L | 99  | 5   |      |      |  | 70 (63) | 130 (105) | 20 (20) |
|               | 4-Chlorophenyl-phenylether  | 50    | 48.0   | ug/L | 96  | 5   |      |      |  | 70 (61) | 130 (104) | 20 (20) |
|               | Fluorene                    | 50    | 50.3   | ug/L | 101 | 4   |      |      |  | 70 (64) | 130 (107) | 20 (20) |
|               | 4-Nitroaniline              | 50    | 52.3   | ug/L | 105 | 2   |      |      |  | 70 (55) | 130 (125) | 20 (20) |
|               | 4,6-Dinitro-2-methylphenol  | 50    | 47.9   | ug/L | 96  | 7   |      |      |  | 70 (62) | 130 (132) | 20 (20) |
|               | N-Nitrosodiphenylamine      | 50    | 49.3   | ug/L | 99  | 6   |      |      |  | 70 (61) | 130 (109) | 20 (20) |
|               | 4-Bromophenyl-phenylether   | 50    | 48.1   | ug/L | 96  | 6   |      |      |  | 70 (73) | 130 (103) | 20 (20) |

() = LABORATORY INHOUSE LIMIT

## Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

|                                       |                                        |
|---------------------------------------|----------------------------------------|
| <b>SDG No.:</b> <u>Q3273</u>          | <b>Analytical Method:</b> <u>8270E</u> |
| <b>Client:</b> <u>G Environmental</u> | <b>DataFile:</b> <u>BG064490.D</u>     |

| Lab Sample ID | Parameter                  | Spike | Result | Unit | Rec | RPD | Qual | RPD  |  | Limits  |           | RPD     |
|---------------|----------------------------|-------|--------|------|-----|-----|------|------|--|---------|-----------|---------|
|               |                            |       |        |      |     |     |      | Qual |  | Low     | High      |         |
| PB169973BSD   | Hexachlorobenzene          | 50    | 48.9   | ug/L | 98  | 4   |      |      |  | 70 (73) | 130 (106) | 20 (20) |
|               | Atrazine                   | 50    | 52.8   | ug/L | 106 | 4   |      |      |  | 70 (76) | 130 (120) | 20 (20) |
|               | Pentachlorophenol          | 100   | 100    | ug/L | 100 | 10  |      |      |  | 20 (47) | 160 (114) | 20 (20) |
|               | Phenanthrene               | 50    | 48.6   | ug/L | 97  | 6   |      |      |  | 70 (62) | 130 (109) | 20 (20) |
|               | Anthracene                 | 50    | 49.9   | ug/L | 100 | 6   |      |      |  | 70 (65) | 130 (110) | 20 (20) |
|               | Carbazole                  | 50    | 52.1   | ug/L | 104 | 2   |      |      |  | 70 (62) | 130 (106) | 20 (20) |
|               | Di-n-butylphthalate        | 50    | 51.7   | ug/L | 103 | 2   |      |      |  | 70 (64) | 130 (106) | 20 (20) |
|               | Fluoranthene               | 50    | 51.0   | ug/L | 102 | 2   |      |      |  | 70 (64) | 130 (110) | 20 (20) |
|               | Pyrene                     | 50    | 49.5   | ug/L | 99  | 4   |      |      |  | 70 (71) | 130 (103) | 20 (20) |
|               | Butylbenzylphthalate       | 50    | 50.7   | ug/L | 101 | 4   |      |      |  | 70 (61) | 130 (105) | 20 (20) |
|               | 3,3-Dichlorobenzidine      | 50    | 27.3   | ug/L | 55  | 11  | *    |      |  | 70 (43) | 130 (108) | 20 (20) |
|               | Benzo(a)anthracene         | 50    | 50.0   | ug/L | 100 | 3   |      |      |  | 70 (62) | 130 (107) | 20 (20) |
|               | Chrysene                   | 50    | 49.6   | ug/L | 99  | 3   |      |      |  | 70 (61) | 130 (108) | 20 (20) |
|               | bis(2-Ethylhexyl)phthalate | 50    | 51.0   | ug/L | 102 | 3   |      |      |  | 70 (59) | 130 (110) | 20 (20) |
|               | Di-n-octyl phthalate       | 50    | 50.3   | ug/L | 101 | 3   |      |      |  | 70 (52) | 130 (139) | 20 (20) |
|               | Benzo(b)fluoranthene       | 50    | 50.4   | ug/L | 101 | 7   |      |      |  | 70 (77) | 130 (113) | 20 (20) |
|               | Benzo(k)fluoranthene       | 50    | 51.7   | ug/L | 103 | 0   |      |      |  | 70 (77) | 130 (105) | 20 (20) |
|               | Benzo(a)pyrene             | 50    | 53.3   | ug/L | 107 | 3   |      |      |  | 70 (72) | 130 (131) | 20 (20) |
|               | Indeno(1,2,3-cd)pyrene     | 50    | 51.2   | ug/L | 102 | 5   |      |      |  | 70 (72) | 130 (105) | 20 (20) |
|               | Dibenz(a,h)anthracene      | 50    | 52.4   | ug/L | 105 | 5   |      |      |  | 70 (78) | 130 (115) | 20 (20) |
|               | Benzo(g,h,i)perylene       | 50    | 52.7   | ug/L | 105 | 4   |      |      |  | 70 (75) | 130 (118) | 20 (20) |
|               | 1,2,4,5-Tetrachlorobenzene | 50    | 53.7   | ug/L | 107 | 1   |      |      |  | 70 (72) | 130 (101) | 20 (20) |
|               | 1,4-Dioxane                | 50    | 40.6   | ug/L | 81  | 7   |      |      |  | 20 (38) | 160 (125) | 20 (20) |
|               | 2,3,4,6-Tetrachlorophenol  | 50    | 54.3   | ug/L | 109 | 1   |      |      |  | 70 (63) | 130 (116) | 20 (20) |

() = LABORATORY INHOUSE LIMIT

4B

SEMIVOLATILE METHOD BLANK SUMMARY

Client ID

PB169973BL

Lab Name: Alliance

Contract: GENV01

Lab Code: ACE

SDG NO.: Q3273

Lab File ID: BG064488.D

Lab Sample ID: PB169973BL

Instrument ID: BNA\_G

Date Extracted: 10/03/2025

Matrix: (soil/water) Water

Date Analyzed: 10/03/2025

Level: (low/med) LOW

Time Analyzed: 15:35

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, MS AND MSD:

| EPA<br>SAMPLE NO. | LAB<br>SAMPLE ID | LAB<br>FILE ID | DATE<br>ANALYZED |
|-------------------|------------------|----------------|------------------|
| PB169973BS        | PB169973BS       | BG064489.D     | 10/03/2025       |
| PB169973BSD       | PB169973BSD      | BG064490.D     | 10/03/2025       |
| MW1               | Q3273-01         | BG064493.D     | 10/03/2025       |

COMMENTS: \_\_\_\_\_

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK  
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: Alliance  
Lab Code: ACE  
Lab File ID: BG064472.D  
Instrument ID: BNA\_G

Contract: GENV01  
SDG NO.: Q3273  
DFTPP Injection Date: 10/02/2025  
DFTPP Injection Time: 08:21

| m/e | ION ABUNDANCE CRITERIA             | % RELATIVE ABUNDANCE |
|-----|------------------------------------|----------------------|
| 68  | Less than 2.0% of mass 69          | 0.3 ( 1.1 ) 1        |
| 69  | Mass 69 relative abundance         | 29.9                 |
| 70  | Less than 2.0% of mass 69          | 0.1 ( 0.4 ) 1        |
| 197 | Less than 2.0% of mass 198         | 0.0                  |
| 198 | Base Peak, 100% relative abundance | 100                  |
| 199 | 5.0 to 9.0% of mass 198            | 6.4                  |
| 365 | Greater than 1% of mass 198        | 6.7                  |
| 441 | Present, but less than mass 443    | 16.8                 |
| 442 | Greater than 50% of mass 198       | 100                  |
| 443 | 15.0 - 24.0% of mass 442           | 19.5 ( 19.5 ) 2      |

1-Value is % mass 69

2-Value is % mass 442

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

| EPA<br>SAMPLE NO. | LAB<br>SAMPLE ID | LAB<br>FILE ID | DATE<br>ANALYZED | TIME<br>ANALYZED |
|-------------------|------------------|----------------|------------------|------------------|
| SSTDICC2.5        | SSTDICC2.5       | BG064473.D     | 10/02/2025       | 09:01            |
| SSTDICC005        | SSTDICC005       | BG064474.D     | 10/02/2025       | 09:42            |
| SSTDICC020        | SSTDICC020       | BG064476.D     | 10/02/2025       | 11:33            |
| SSTDICCC040       | SSTDICCC040      | BG064477.D     | 10/02/2025       | 12:14            |
| SSTDICC050        | SSTDICC050       | BG064478.D     | 10/02/2025       | 12:55            |
| SSTDICC060        | SSTDICC060       | BG064479.D     | 10/02/2025       | 13:35            |
| SSTDICC080        | SSTDICC080       | BG064480.D     | 10/02/2025       | 14:16            |
| SSTDICC010        | SSTDICC010       | BG064481.D     | 10/02/2025       | 15:49            |

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK  
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: Alliance  
Lab Code: ACE  
Lab File ID: BG064484.D  
Instrument ID: BNA\_G

Contract: GENV01  
SDG NO.: Q3273  
DFTPP Injection Date: 10/03/2025  
DFTPP Injection Time: 12:52

| m/e | ION ABUNDANCE CRITERIA             | % RELATIVE ABUNDANCE |
|-----|------------------------------------|----------------------|
| 68  | Less than 2.0% of mass 69          | 0.4 ( 1.4 ) 1        |
| 69  | Mass 69 relative abundance         | 28.7                 |
| 70  | Less than 2.0% of mass 69          | 0.1 ( 0.3 ) 1        |
| 197 | Less than 2.0% of mass 198         | 0.3                  |
| 198 | Base Peak, 100% relative abundance | 100                  |
| 199 | 5.0 to 9.0% of mass 198            | 6.1                  |
| 365 | Greater than 1% of mass 198        | 6.1                  |
| 441 | Present, but less than mass 443    | 11.1                 |
| 442 | Greater than 50% of mass 198       | 100                  |
| 443 | 15.0 - 24.0% of mass 442           | 19.9 ( 19.9 ) 2      |

1-Value is % mass 69

2-Value is % mass 442

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

| EPA<br>SAMPLE NO. | LAB<br>SAMPLE ID | LAB<br>FILE ID | DATE<br>ANALYZED | TIME<br>ANALYZED |
|-------------------|------------------|----------------|------------------|------------------|
| SSTDCCC040        | SSTDCCC040       | BG064485.D     | 10/03/2025       | 13:33            |
| PB169973BL        | PB169973BL       | BG064488.D     | 10/03/2025       | 15:35            |
| PB169973BS        | PB169973BS       | BG064489.D     | 10/03/2025       | 16:16            |
| PB169973BSD       | PB169973BSD      | BG064490.D     | 10/03/2025       | 16:57            |
| MW1               | Q3273-01         | BG064493.D     | 10/03/2025       | 18:59            |

8B

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: Alliance

Lab Code: ACE

SDG NO.: Q3273

Client ID : SSTDCCC040

Date Analyzed: 10/03/2025

Lab File ID: BG064485.D

Time Analyzed: 13:33

Instrument ID: BNA\_G

GC Column: ZB-GR ID: 0.25 (mm)

|                | IS1 (DCB)<br>AREA # | RT #  | IS2 (NPT)<br>AREA # | RT #   | IS3 (ANT)<br>AREA # | RT #   |
|----------------|---------------------|-------|---------------------|--------|---------------------|--------|
| 12 HOUR STD    | 17500               | 7.821 | 78873               | 10.61  | 59706               | 14.45  |
| UPPER LIMIT    | 35000               | 8.321 | 157746              | 11.112 | 119412              | 14.948 |
| LOWER LIMIT    | 8750                | 7.321 | 39436.5             | 10.112 | 29853               | 13.948 |
| EPA SAMPLE NO. |                     |       |                     |        |                     |        |
| 01 PB169973BL  | 16965               | 7.82  | 71465               | 10.61  | 57081               | 14.45  |
| 02 PB169973BS  | 16264               | 7.82  | 69209               | 10.61  | 53762               | 14.45  |
| 03 PB169973BSD | 15312               | 7.83  | 68886               | 10.61  | 53563               | 14.45  |
| 04 MW1         | 20058               | 7.82  | 82373               | 10.61  | 60802               | 14.46  |

IS1 (DCB) = 1,4-Dichlorobenzene-d4

IS2 (NPT) = Naphthalene-d8

IS3 (ANT) = Acenaphthene-d10

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = -50% of internal standard area

RT UPPER LIMIT = +0.50 minutes of internal standard RT

RT UPPER LIMIT = -0.50 minutes of internal standard RT

# Column used to flag values outside QC limits with an asterisk.

\* Values outside of QC limits.



8C

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: Alliance

Lab Code: ACE

SDG NO.: Q3273

Client ID: SSTDCCC040

Date Analyzed: 10/03/2025

Lab File ID: BG064485.D

Time Analyzed: 13:33

Instrument ID: BNA\_G

GC Column: ZB-GR ID: 0.25 (mm)

|                | IS4 (PHN)<br>AREA # | RT #   | IS5 (CRY)<br>AREA # | RT #   | IS6 (PRY)<br>AREA # | RT #  |
|----------------|---------------------|--------|---------------------|--------|---------------------|-------|
| 12 HOUR STD    | 149943              | 17.186 | 155011              | 21.417 | 163411              | 24.39 |
| UPPER LIMIT    | 299886              | 17.686 | 310022              | 21.917 | 326822              | 24.89 |
| LOWER LIMIT    | 74971.5             | 16.686 | 77505.5             | 20.917 | 81705.5             | 23.89 |
| EPA SAMPLE NO. |                     |        |                     |        |                     |       |
| 01 PB169973BL  | 145353              | 17.19  | 149348              | 21.41  | 157878              | 24.39 |
| 02 PB169973BS  | 136086              | 17.19  | 136322              | 21.41  | 146226              | 24.39 |
| 03 PB169973BSD | 138632              | 17.19  | 141515              | 21.42  | 151583              | 24.39 |
| 04 MW1         | 129947              | 17.20  | 136555              | 21.42  | 149983              | 24.39 |

IS4 (PHN) = Phenanthrene-d10

IS5 (CRY) = Chrysene-d12

IS6 (PRY) = Perylene-d12

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = -50% of internal standard area

RT UPPER LIMIT = +0.50 minutes of internal standard RT

RT UPPER LIMIT = -0.50 minutes of internal standard RT

# Column used to flag values outside QC limits with an asterisk.

\* Values outside of QC limits.



# QC SAMPLE DATA

## Report of Analysis

|                    |                        |                 |                  |
|--------------------|------------------------|-----------------|------------------|
| Client:            | G Environmental        | Date Collected: |                  |
| Project:           | Monroe, NJ             | Date Received:  |                  |
| Client Sample ID:  | PB169973BL             | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BL             | Matrix:         | Water            |
| Analytical Method: | 8270E                  | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000      Units:    mL | Final Vol:      | 1000      uL     |
| Soil Aliquot Vol:  | uL                     | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted :    N        | Level :         | LOW              |
| Injection Volume : | GPC Factor :    1.0    | GPC Cleanup :   | N      PH :      |
| Prep Method :      |                        |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064488.D        | 1         | 10/03/25 08:35 | 10/03/25 15:35 | PB169973      |

| CAS Number     | Parameter                   | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|----------------|-----------------------------|-------|-----------|------|------------|-------|
| <b>TARGETS</b> |                             |       |           |      |            |       |
| 100-52-7       | Benzaldehyde                | 3.90  | U         | 3.90 | 10.0       | ug/L  |
| 108-95-2       | Phenol                      | 0.91  | U         | 0.91 | 5.00       | ug/L  |
| 111-44-4       | bis(2-Chloroethyl)ether     | 0.81  | U         | 0.81 | 5.00       | ug/L  |
| 95-57-8        | 2-Chlorophenol              | 0.58  | U         | 0.58 | 5.00       | ug/L  |
| 95-48-7        | 2-Methylphenol              | 1.10  | U         | 1.10 | 5.00       | ug/L  |
| 108-60-1       | 2,2-oxybis(1-Chloropropane) | 1.30  | U         | 1.30 | 5.00       | ug/L  |
| 98-86-2        | Acetophenone                | 0.74  | U         | 0.74 | 5.00       | ug/L  |
| 65794-96-9     | 3+4-Methylphenols           | 1.10  | U         | 1.10 | 10.0       | ug/L  |
| 621-64-7       | n-Nitroso-di-n-propylamine  | 1.40  | U         | 1.40 | 2.50       | ug/L  |
| 67-72-1        | Hexachloroethane            | 0.65  | U         | 0.65 | 5.00       | ug/L  |
| 98-95-3        | Nitrobenzene                | 0.76  | U         | 0.76 | 5.00       | ug/L  |
| 78-59-1        | Isophorone                  | 0.75  | U         | 0.75 | 5.00       | ug/L  |
| 88-75-5        | 2-Nitrophenol               | 1.80  | U         | 1.80 | 5.00       | ug/L  |
| 105-67-9       | 2,4-Dimethylphenol          | 1.90  | U         | 1.90 | 5.00       | ug/L  |
| 111-91-1       | bis(2-Chloroethoxy)methane  | 0.68  | U         | 0.68 | 5.00       | ug/L  |
| 120-83-2       | 2,4-Dichlorophenol          | 0.52  | U         | 0.52 | 5.00       | ug/L  |
| 91-20-3        | Naphthalene                 | 0.50  | U         | 0.50 | 5.00       | ug/L  |
| 106-47-8       | 4-Chloroaniline             | 0.84  | U         | 0.84 | 5.00       | ug/L  |
| 87-68-3        | Hexachlorobutadiene         | 0.54  | U         | 0.54 | 5.00       | ug/L  |
| 105-60-2       | Caprolactam                 | 1.10  | U         | 1.10 | 10.0       | ug/L  |
| 59-50-7        | 4-Chloro-3-methylphenol     | 0.59  | U         | 0.59 | 5.00       | ug/L  |
| 91-57-6        | 2-Methylnaphthalene         | 0.56  | U         | 0.56 | 5.00       | ug/L  |
| 77-47-4        | Hexachlorocyclopentadiene   | 3.60  | U         | 3.60 | 10.0       | ug/L  |
| 88-06-2        | 2,4,6-Trichlorophenol       | 0.51  | U         | 0.51 | 5.00       | ug/L  |
| 95-95-4        | 2,4,5-Trichlorophenol       | 0.62  | U         | 0.62 | 5.00       | ug/L  |
| 92-52-4        | 1,1-Biphenyl                | 0.53  | U         | 0.53 | 5.00       | ug/L  |
| 91-58-7        | 2-Chloronaphthalene         | 0.61  | U         | 0.61 | 5.00       | ug/L  |
| 88-74-4        | 2-Nitroaniline              | 1.30  | U         | 1.30 | 5.00       | ug/L  |
| 131-11-3       | Dimethylphthalate           | 0.61  | U         | 0.61 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BL       | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BL       | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064488.D        | 1         | 10/03/25 08:35 | 10/03/25 15:35 | PB169973      |

| CAS Number | Parameter                  | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|------------|----------------------------|-------|-----------|------|------------|-------|
| 208-96-8   | Acenaphthylene             | 0.75  | U         | 0.75 | 5.00       | ug/L  |
| 606-20-2   | 2,6-Dinitrotoluene         | 0.92  | U         | 0.92 | 5.00       | ug/L  |
| 99-09-2    | 3-Nitroaniline             | 1.10  | U         | 1.10 | 5.00       | ug/L  |
| 83-32-9    | Acenaphthene               | 0.55  | U         | 0.55 | 5.00       | ug/L  |
| 51-28-5    | 2,4-Dinitrophenol          | 6.00  | U         | 6.00 | 10.0       | ug/L  |
| 100-02-7   | 4-Nitrophenol              | 2.40  | U         | 2.40 | 10.0       | ug/L  |
| 132-64-9   | Dibenzofuran               | 0.61  | U         | 0.61 | 5.00       | ug/L  |
| 121-14-2   | 2,4-Dinitrotoluene         | 1.20  | U         | 1.20 | 5.00       | ug/L  |
| 84-66-2    | Diethylphthalate           | 0.69  | U         | 0.69 | 5.00       | ug/L  |
| 7005-72-3  | 4-Chlorophenyl-phenylether | 0.68  | U         | 0.68 | 5.00       | ug/L  |
| 86-73-7    | Fluorene                   | 0.63  | U         | 0.63 | 5.00       | ug/L  |
| 100-01-6   | 4-Nitroaniline             | 1.50  | U         | 1.50 | 5.00       | ug/L  |
| 534-52-1   | 4,6-Dinitro-2-methylphenol | 2.90  | U         | 2.90 | 10.0       | ug/L  |
| 86-30-6    | n-Nitrosodiphenylamine     | 0.58  | U         | 0.58 | 5.00       | ug/L  |
| 101-55-3   | 4-Bromophenyl-phenylether  | 0.40  | U         | 0.40 | 5.00       | ug/L  |
| 118-74-1   | Hexachlorobenzene          | 0.52  | U         | 0.52 | 5.00       | ug/L  |
| 1912-24-9  | Atrazine                   | 1.00  | U         | 1.00 | 5.00       | ug/L  |
| 87-86-5    | Pentachlorophenol          | 1.60  | U         | 1.60 | 10.0       | ug/L  |
| 85-01-8    | Phenanthrene               | 0.50  | U         | 0.50 | 5.00       | ug/L  |
| 120-12-7   | Anthracene                 | 0.61  | U         | 0.61 | 5.00       | ug/L  |
| 86-74-8    | Carbazole                  | 0.72  | U         | 0.72 | 5.00       | ug/L  |
| 84-74-2    | Di-n-butylphthalate        | 1.20  | U         | 1.20 | 5.00       | ug/L  |
| 206-44-0   | Fluoranthene               | 0.82  | U         | 0.82 | 5.00       | ug/L  |
| 129-00-0   | Pyrene                     | 0.50  | U         | 0.50 | 5.00       | ug/L  |
| 85-68-7    | Butylbenzylphthalate       | 1.90  | U         | 1.90 | 5.00       | ug/L  |
| 91-94-1    | 3,3-Dichlorobenzidine      | 0.93  | U         | 0.93 | 10.0       | ug/L  |
| 56-55-3    | Benzo(a)anthracene         | 0.45  | U         | 0.45 | 5.00       | ug/L  |
| 218-01-9   | Chrysene                   | 0.44  | U         | 0.44 | 5.00       | ug/L  |
| 117-81-7   | Bis(2-ethylhexyl)phthalate | 1.60  | U         | 1.60 | 5.00       | ug/L  |
| 117-84-0   | Di-n-octyl phthalate       | 2.30  | U         | 2.30 | 10.0       | ug/L  |
| 205-99-2   | Benzo(b)fluoranthene       | 0.49  | U         | 0.49 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BL       | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BL       | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064488.D        | 1         | 10/03/25 08:35 | 10/03/25 15:35 | PB169973      |

| CAS Number                | Parameter                  | Conc.  | Qualifier | MDL                 | LOQ / CRQL | Units    |
|---------------------------|----------------------------|--------|-----------|---------------------|------------|----------|
| 207-08-9                  | Benzo(k)fluoranthene       | 0.48   | U         | 0.48                | 5.00       | ug/L     |
| 50-32-8                   | Benzo(a)pyrene             | 0.55   | U         | 0.55                | 5.00       | ug/L     |
| 193-39-5                  | Indeno(1,2,3-cd)pyrene     | 0.59   | U         | 0.59                | 5.00       | ug/L     |
| 53-70-3                   | Dibenzo(a,h)anthracene     | 0.67   | U         | 0.67                | 5.00       | ug/L     |
| 191-24-2                  | Benzo(g,h,i)perylene       | 0.69   | U         | 0.69                | 5.00       | ug/L     |
| 95-94-3                   | 1,2,4,5-Tetrachlorobenzene | 0.52   | U         | 0.52                | 5.00       | ug/L     |
| 123-91-1                  | 1,4-Dioxane                | 1.00   | U         | 1.00                | 5.00       | ug/L     |
| 58-90-2                   | 2,3,4,6-Tetrachlorophenol  | 0.72   | U         | 0.72                | 5.00       | ug/L     |
| <b>SURROGATES</b>         |                            |        |           |                     |            |          |
| 367-12-4                  | 2-Fluorophenol             | 162    |           | 15 (23) - 110 (138) | 108%       | SPK: 150 |
| 13127-88-3                | Phenol-d6                  | 150    |           | 15 (10) - 110 (134) | 100%       | SPK: 150 |
| 4165-60-0                 | Nitrobenzene-d5            | 94.0   |           | 30 (67) - 130 (132) | 94%        | SPK: 100 |
| 321-60-8                  | 2-Fluorobiphenyl           | 90.7   |           | 30 (52) - 130 (132) | 91%        | SPK: 100 |
| 118-79-6                  | 2,4,6-Tribromophenol       | 138    |           | 15 (44) - 110 (137) | 92%        | SPK: 150 |
| 1718-51-0                 | Terphenyl-d14              | 98.8   |           | 30 (42) - 130 (152) | 99%        | SPK: 100 |
| <b>INTERNAL STANDARDS</b> |                            |        |           |                     |            |          |
| 3855-82-1                 | 1,4-Dichlorobenzene-d4     | 17000  | 7.824     |                     |            |          |
| 1146-65-2                 | Naphthalene-d8             | 71500  | 10.609    |                     |            |          |
| 15067-26-2                | Acenaphthene-d10           | 57100  | 14.446    |                     |            |          |
| 1517-22-2                 | Phenanthrene-d10           | 145000 | 17.189    |                     |            |          |
| 1719-03-5                 | Chrysene-d12               | 149000 | 21.408    |                     |            |          |
| 1520-96-3                 | Perylene-d12               | 158000 | 24.387    |                     |            |          |

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

\* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BS       | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BS       | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064489.D        | 1         | 10/03/25 08:35 | 10/03/25 16:16 | PB169973      |

| CAS Number     | Parameter                   | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|----------------|-----------------------------|-------|-----------|------|------------|-------|
| <b>TARGETS</b> |                             |       |           |      |            |       |
| 100-52-7       | Benzaldehyde                | 26.5  |           | 3.90 | 10.0       | ug/L  |
| 108-95-2       | Phenol                      | 53.3  |           | 0.91 | 5.00       | ug/L  |
| 111-44-4       | bis(2-Chloroethyl)ether     | 49.4  |           | 0.81 | 5.00       | ug/L  |
| 95-57-8        | 2-Chlorophenol              | 51.6  |           | 0.58 | 5.00       | ug/L  |
| 95-48-7        | 2-Methylphenol              | 52.4  |           | 1.10 | 5.00       | ug/L  |
| 108-60-1       | 2,2-oxybis(1-Chloropropane) | 49.5  |           | 1.30 | 5.00       | ug/L  |
| 98-86-2        | Acetophenone                | 57.0  |           | 0.74 | 5.00       | ug/L  |
| 65794-96-9     | 3+4-Methylphenols           | 53.3  |           | 1.10 | 10.0       | ug/L  |
| 621-64-7       | n-Nitroso-di-n-propylamine  | 51.0  |           | 1.40 | 2.50       | ug/L  |
| 67-72-1        | Hexachloroethane            | 49.2  |           | 0.65 | 5.00       | ug/L  |
| 98-95-3        | Nitrobenzene                | 50.7  |           | 0.76 | 5.00       | ug/L  |
| 78-59-1        | Isophorone                  | 48.4  |           | 0.75 | 5.00       | ug/L  |
| 88-75-5        | 2-Nitrophenol               | 49.1  |           | 1.80 | 5.00       | ug/L  |
| 105-67-9       | 2,4-Dimethylphenol          | 58.0  |           | 1.90 | 5.00       | ug/L  |
| 111-91-1       | bis(2-Chloroethoxy)methane  | 50.5  |           | 0.68 | 5.00       | ug/L  |
| 120-83-2       | 2,4-Dichlorophenol          | 55.8  |           | 0.52 | 5.00       | ug/L  |
| 91-20-3        | Naphthalene                 | 50.7  |           | 0.50 | 5.00       | ug/L  |
| 106-47-8       | 4-Chloroaniline             | 18.8  |           | 0.84 | 5.00       | ug/L  |
| 87-68-3        | Hexachlorobutadiene         | 49.1  |           | 0.54 | 5.00       | ug/L  |
| 105-60-2       | Caprolactam                 | 52.0  |           | 1.10 | 10.0       | ug/L  |
| 59-50-7        | 4-Chloro-3-methylphenol     | 53.9  |           | 0.59 | 5.00       | ug/L  |
| 91-57-6        | 2-Methylnaphthalene         | 46.2  |           | 0.56 | 5.00       | ug/L  |
| 77-47-4        | Hexachlorocyclopentadiene   | 150   | E         | 3.60 | 10.0       | ug/L  |
| 88-06-2        | 2,4,6-Trichlorophenol       | 53.5  |           | 0.51 | 5.00       | ug/L  |
| 95-95-4        | 2,4,5-Trichlorophenol       | 54.4  |           | 0.62 | 5.00       | ug/L  |
| 92-52-4        | 1,1-Biphenyl                | 56.4  |           | 0.53 | 5.00       | ug/L  |
| 91-58-7        | 2-Chloronaphthalene         | 50.1  |           | 0.61 | 5.00       | ug/L  |
| 88-74-4        | 2-Nitroaniline              | 53.0  |           | 1.30 | 5.00       | ug/L  |
| 131-11-3       | Dimethylphthalate           | 52.1  |           | 0.61 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BS       | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BS       | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064489.D        | 1         | 10/03/25 08:35 | 10/03/25 16:16 | PB169973      |

| CAS Number | Parameter                  | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|------------|----------------------------|-------|-----------|------|------------|-------|
| 208-96-8   | Acenaphthylene             | 57.9  |           | 0.75 | 5.00       | ug/L  |
| 606-20-2   | 2,6-Dinitrotoluene         | 53.0  |           | 0.92 | 5.00       | ug/L  |
| 99-09-2    | 3-Nitroaniline             | 30.0  |           | 1.10 | 5.00       | ug/L  |
| 83-32-9    | Acenaphthene               | 49.8  |           | 0.55 | 5.00       | ug/L  |
| 51-28-5    | 2,4-Dinitrophenol          | 95.5  | E         | 6.00 | 10.0       | ug/L  |
| 100-02-7   | 4-Nitrophenol              | 110   | E         | 2.40 | 10.0       | ug/L  |
| 132-64-9   | Dibenzofuran               | 51.4  |           | 0.61 | 5.00       | ug/L  |
| 121-14-2   | 2,4-Dinitrotoluene         | 52.2  |           | 1.20 | 5.00       | ug/L  |
| 84-66-2    | Diethylphthalate           | 51.8  |           | 0.69 | 5.00       | ug/L  |
| 7005-72-3  | 4-Chlorophenyl-phenylether | 50.4  |           | 0.68 | 5.00       | ug/L  |
| 86-73-7    | Fluorene                   | 52.1  |           | 0.63 | 5.00       | ug/L  |
| 100-01-6   | 4-Nitroaniline             | 53.5  |           | 1.50 | 5.00       | ug/L  |
| 534-52-1   | 4,6-Dinitro-2-methylphenol | 51.5  |           | 2.90 | 10.0       | ug/L  |
| 86-30-6    | n-Nitrosodiphenylamine     | 52.3  |           | 0.58 | 5.00       | ug/L  |
| 101-55-3   | 4-Bromophenyl-phenylether  | 51.0  |           | 0.40 | 5.00       | ug/L  |
| 118-74-1   | Hexachlorobenzene          | 50.8  |           | 0.52 | 5.00       | ug/L  |
| 1912-24-9  | Atrazine                   | 55.0  |           | 1.00 | 5.00       | ug/L  |
| 87-86-5    | Pentachlorophenol          | 110   | E         | 1.60 | 10.0       | ug/L  |
| 85-01-8    | Phenanthrene               | 51.6  |           | 0.50 | 5.00       | ug/L  |
| 120-12-7   | Anthracene                 | 53.0  |           | 0.61 | 5.00       | ug/L  |
| 86-74-8    | Carbazole                  | 53.2  |           | 0.72 | 5.00       | ug/L  |
| 84-74-2    | Di-n-butylphthalate        | 52.6  |           | 1.20 | 5.00       | ug/L  |
| 206-44-0   | Fluoranthene               | 52.1  |           | 0.82 | 5.00       | ug/L  |
| 129-00-0   | Pyrene                     | 51.7  |           | 0.50 | 5.00       | ug/L  |
| 85-68-7    | Butylbenzylphthalate       | 52.8  |           | 1.90 | 5.00       | ug/L  |
| 91-94-1    | 3,3-Dichlorobenzidine      | 30.6  |           | 0.93 | 10.0       | ug/L  |
| 56-55-3    | Benzo(a)anthracene         | 51.5  |           | 0.45 | 5.00       | ug/L  |
| 218-01-9   | Chrysene                   | 51.3  |           | 0.44 | 5.00       | ug/L  |
| 117-81-7   | Bis(2-ethylhexyl)phthalate | 52.7  |           | 1.60 | 5.00       | ug/L  |
| 117-84-0   | Di-n-octyl phthalate       | 52.0  |           | 2.30 | 10.0       | ug/L  |
| 205-99-2   | Benzo(b)fluoranthene       | 54.2  |           | 0.49 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BS       | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BS       | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064489.D        | 1         | 10/03/25 08:35 | 10/03/25 16:16 | PB169973      |

| CAS Number                | Parameter                  | Conc.  | Qualifier | MDL                 | LOQ / CRQL | Units    |
|---------------------------|----------------------------|--------|-----------|---------------------|------------|----------|
| 207-08-9                  | Benzo(k)fluoranthene       | 51.8   |           | 0.48                | 5.00       | ug/L     |
| 50-32-8                   | Benzo(a)pyrene             | 54.9   |           | 0.55                | 5.00       | ug/L     |
| 193-39-5                  | Indeno(1,2,3-cd)pyrene     | 53.9   |           | 0.59                | 5.00       | ug/L     |
| 53-70-3                   | Dibenzo(a,h)anthracene     | 55.2   |           | 0.67                | 5.00       | ug/L     |
| 191-24-2                  | Benzo(g,h,i)perylene       | 54.7   |           | 0.69                | 5.00       | ug/L     |
| 95-94-3                   | 1,2,4,5-Tetrachlorobenzene | 54.2   |           | 0.52                | 5.00       | ug/L     |
| 123-91-1                  | 1,4-Dioxane                | 37.9   |           | 1.00                | 5.00       | ug/L     |
| 58-90-2                   | 2,3,4,6-Tetrachlorophenol  | 54.7   |           | 0.72                | 5.00       | ug/L     |
| <b>SURROGATES</b>         |                            |        |           |                     |            |          |
| 367-12-4                  | 2-Fluorophenol             | 157    |           | 15 (23) - 110 (138) | 105%       | SPK: 150 |
| 13127-88-3                | Phenol-d6                  | 153    |           | 15 (10) - 110 (134) | 102%       | SPK: 150 |
| 4165-60-0                 | Nitrobenzene-d5            | 94.3   |           | 30 (67) - 130 (132) | 94%        | SPK: 100 |
| 321-60-8                  | 2-Fluorobiphenyl           | 91.8   |           | 30 (52) - 130 (132) | 92%        | SPK: 100 |
| 118-79-6                  | 2,4,6-Tribromophenol       | 149    |           | 15 (44) - 110 (137) | 99%        | SPK: 150 |
| 1718-51-0                 | Terphenyl-d14              | 97.3   |           | 30 (42) - 130 (152) | 97%        | SPK: 100 |
| <b>INTERNAL STANDARDS</b> |                            |        |           |                     |            |          |
| 3855-82-1                 | 1,4-Dichlorobenzene-d4     | 16300  | 7.823     |                     |            |          |
| 1146-65-2                 | Naphthalene-d8             | 69200  | 10.608    |                     |            |          |
| 15067-26-2                | Acenaphthene-d10           | 53800  | 14.45     |                     |            |          |
| 1517-22-2                 | Phenanthrene-d10           | 136000 | 17.188    |                     |            |          |
| 1719-03-5                 | Chrysene-d12               | 136000 | 21.413    |                     |            |          |
| 1520-96-3                 | Perylene-d12               | 146000 | 24.391    |                     |            |          |

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

\* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products



## Report of Analysis

|                    |                        |                 |                   |
|--------------------|------------------------|-----------------|-------------------|
| Client:            | G Environmental        | Date Collected: |                   |
| Project:           | Monroe, NJ             | Date Received:  |                   |
| Client Sample ID:  | PB169973BSD            | SDG No.:        | Q3273             |
| Lab Sample ID:     | PB169973BSD            | Matrix:         | Water             |
| Analytical Method: | 8270E                  | % Solid:        | 0                 |
| Sample Wt/Vol:     | 1000      Units:    mL | Final Vol:      | 1000      uL      |
| Soil Aliquot Vol:  | uL                     | Test:           | SVOC-TCL BNA -20  |
| Extraction Type :  | Decanted :    N        | Level :         | LOW               |
| Injection Volume : | GPC Factor :    1.0    | GPC Cleanup :   | N            PH : |
| Prep Method :      |                        |                 |                   |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064490.D        | 1         | 10/03/25 08:35 | 10/03/25 16:57 | PB169973      |

| CAS Number     | Parameter                   | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|----------------|-----------------------------|-------|-----------|------|------------|-------|
| <b>TARGETS</b> |                             |       |           |      |            |       |
| 100-52-7       | Benzaldehyde                | 26.1  |           | 3.90 | 10.0       | ug/L  |
| 108-95-2       | Phenol                      | 55.3  |           | 0.91 | 5.00       | ug/L  |
| 111-44-4       | bis(2-Chloroethyl)ether     | 49.7  |           | 0.81 | 5.00       | ug/L  |
| 95-57-8        | 2-Chlorophenol              | 55.2  |           | 0.58 | 5.00       | ug/L  |
| 95-48-7        | 2-Methylphenol              | 55.0  |           | 1.10 | 5.00       | ug/L  |
| 108-60-1       | 2,2-oxybis(1-Chloropropane) | 48.3  |           | 1.30 | 5.00       | ug/L  |
| 98-86-2        | Acetophenone                | 54.9  |           | 0.74 | 5.00       | ug/L  |
| 65794-96-9     | 3+4-Methylphenols           | 55.3  |           | 1.10 | 10.0       | ug/L  |
| 621-64-7       | n-Nitroso-di-n-propylamine  | 52.4  |           | 1.40 | 2.50       | ug/L  |
| 67-72-1        | Hexachloroethane            | 49.3  |           | 0.65 | 5.00       | ug/L  |
| 98-95-3        | Nitrobenzene                | 47.6  |           | 0.76 | 5.00       | ug/L  |
| 78-59-1        | Isophorone                  | 47.7  |           | 0.75 | 5.00       | ug/L  |
| 88-75-5        | 2-Nitrophenol               | 48.4  |           | 1.80 | 5.00       | ug/L  |
| 105-67-9       | 2,4-Dimethylphenol          | 57.0  |           | 1.90 | 5.00       | ug/L  |
| 111-91-1       | bis(2-Chloroethoxy)methane  | 49.3  |           | 0.68 | 5.00       | ug/L  |
| 120-83-2       | 2,4-Dichlorophenol          | 52.6  |           | 0.52 | 5.00       | ug/L  |
| 91-20-3        | Naphthalene                 | 48.5  |           | 0.50 | 5.00       | ug/L  |
| 106-47-8       | 4-Chloroaniline             | 15.2  |           | 0.84 | 5.00       | ug/L  |
| 87-68-3        | Hexachlorobutadiene         | 47.0  |           | 0.54 | 5.00       | ug/L  |
| 105-60-2       | Caprolactam                 | 50.1  |           | 1.10 | 10.0       | ug/L  |
| 59-50-7        | 4-Chloro-3-methylphenol     | 51.6  |           | 0.59 | 5.00       | ug/L  |
| 91-57-6        | 2-Methylnaphthalene         | 44.5  |           | 0.56 | 5.00       | ug/L  |
| 77-47-4        | Hexachlorocyclopentadiene   | 140   | E         | 3.60 | 10.0       | ug/L  |
| 88-06-2        | 2,4,6-Trichlorophenol       | 51.9  |           | 0.51 | 5.00       | ug/L  |
| 95-95-4        | 2,4,5-Trichlorophenol       | 49.0  |           | 0.62 | 5.00       | ug/L  |
| 92-52-4        | 1,1-Biphenyl                | 53.2  |           | 0.53 | 5.00       | ug/L  |
| 91-58-7        | 2-Chloronaphthalene         | 47.4  |           | 0.61 | 5.00       | ug/L  |
| 88-74-4        | 2-Nitroaniline              | 50.6  |           | 1.30 | 5.00       | ug/L  |
| 131-11-3       | Dimethylphthalate           | 48.6  |           | 0.61 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BSD      | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BSD      | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064490.D        | 1         | 10/03/25 08:35 | 10/03/25 16:57 | PB169973      |

| CAS Number | Parameter                  | Conc. | Qualifier | MDL  | LOQ / CRQL | Units |
|------------|----------------------------|-------|-----------|------|------------|-------|
| 208-96-8   | Acenaphthylene             | 55.5  |           | 0.75 | 5.00       | ug/L  |
| 606-20-2   | 2,6-Dinitrotoluene         | 50.2  |           | 0.92 | 5.00       | ug/L  |
| 99-09-2    | 3-Nitroaniline             | 26.1  |           | 1.10 | 5.00       | ug/L  |
| 83-32-9    | Acenaphthene               | 47.3  |           | 0.55 | 5.00       | ug/L  |
| 51-28-5    | 2,4-Dinitrophenol          | 96.1  | E         | 6.00 | 10.0       | ug/L  |
| 100-02-7   | 4-Nitrophenol              | 110   | E         | 2.40 | 10.0       | ug/L  |
| 132-64-9   | Dibenzofuran               | 49.0  |           | 0.61 | 5.00       | ug/L  |
| 121-14-2   | 2,4-Dinitrotoluene         | 53.2  |           | 1.20 | 5.00       | ug/L  |
| 84-66-2    | Diethylphthalate           | 49.5  |           | 0.69 | 5.00       | ug/L  |
| 7005-72-3  | 4-Chlorophenyl-phenylether | 48.0  |           | 0.68 | 5.00       | ug/L  |
| 86-73-7    | Fluorene                   | 50.3  |           | 0.63 | 5.00       | ug/L  |
| 100-01-6   | 4-Nitroaniline             | 52.3  |           | 1.50 | 5.00       | ug/L  |
| 534-52-1   | 4,6-Dinitro-2-methylphenol | 47.9  |           | 2.90 | 10.0       | ug/L  |
| 86-30-6    | n-Nitrosodiphenylamine     | 49.3  |           | 0.58 | 5.00       | ug/L  |
| 101-55-3   | 4-Bromophenyl-phenylether  | 48.1  |           | 0.40 | 5.00       | ug/L  |
| 118-74-1   | Hexachlorobenzene          | 48.9  |           | 0.52 | 5.00       | ug/L  |
| 1912-24-9  | Atrazine                   | 52.8  |           | 1.00 | 5.00       | ug/L  |
| 87-86-5    | Pentachlorophenol          | 100   | E         | 1.60 | 10.0       | ug/L  |
| 85-01-8    | Phenanthrene               | 48.6  |           | 0.50 | 5.00       | ug/L  |
| 120-12-7   | Anthracene                 | 49.9  |           | 0.61 | 5.00       | ug/L  |
| 86-74-8    | Carbazole                  | 52.1  |           | 0.72 | 5.00       | ug/L  |
| 84-74-2    | Di-n-butylphthalate        | 51.7  |           | 1.20 | 5.00       | ug/L  |
| 206-44-0   | Fluoranthene               | 51.0  |           | 0.82 | 5.00       | ug/L  |
| 129-00-0   | Pyrene                     | 49.5  |           | 0.50 | 5.00       | ug/L  |
| 85-68-7    | Butylbenzylphthalate       | 50.7  |           | 1.90 | 5.00       | ug/L  |
| 91-94-1    | 3,3-Dichlorobenzidine      | 27.3  |           | 0.93 | 10.0       | ug/L  |
| 56-55-3    | Benzo(a)anthracene         | 50.0  |           | 0.45 | 5.00       | ug/L  |
| 218-01-9   | Chrysene                   | 49.6  |           | 0.44 | 5.00       | ug/L  |
| 117-81-7   | Bis(2-ethylhexyl)phthalate | 51.0  |           | 1.60 | 5.00       | ug/L  |
| 117-84-0   | Di-n-octyl phthalate       | 50.3  |           | 2.30 | 10.0       | ug/L  |
| 205-99-2   | Benzo(b)fluoranthene       | 50.4  |           | 0.49 | 5.00       | ug/L  |

## Report of Analysis

|                    |                  |                 |                  |
|--------------------|------------------|-----------------|------------------|
| Client:            | G Environmental  | Date Collected: |                  |
| Project:           | Monroe, NJ       | Date Received:  |                  |
| Client Sample ID:  | PB169973BSD      | SDG No.:        | Q3273            |
| Lab Sample ID:     | PB169973BSD      | Matrix:         | Water            |
| Analytical Method: | 8270E            | % Solid:        | 0                |
| Sample Wt/Vol:     | 1000 Units: mL   | Final Vol:      | 1000 uL          |
| Soil Aliquot Vol:  | uL               | Test:           | SVOC-TCL BNA -20 |
| Extraction Type :  | Decanted : N     | Level :         | LOW              |
| Injection Volume : | GPC Factor : 1.0 | GPC Cleanup :   | N PH :           |
| Prep Method :      |                  |                 |                  |

|                   |           |                |                |               |
|-------------------|-----------|----------------|----------------|---------------|
| File ID/Qc Batch: | Dilution: | Prep Date      | Date Analyzed  | Prep Batch ID |
| BG064490.D        | 1         | 10/03/25 08:35 | 10/03/25 16:57 | PB169973      |

| CAS Number                | Parameter                  | Conc.  | Qualifier | MDL                 | LOQ / CRQL | Units    |
|---------------------------|----------------------------|--------|-----------|---------------------|------------|----------|
| 207-08-9                  | Benzo(k)fluoranthene       | 51.7   |           | 0.48                | 5.00       | ug/L     |
| 50-32-8                   | Benzo(a)pyrene             | 53.3   |           | 0.55                | 5.00       | ug/L     |
| 193-39-5                  | Indeno(1,2,3-cd)pyrene     | 51.2   |           | 0.59                | 5.00       | ug/L     |
| 53-70-3                   | Dibenzo(a,h)anthracene     | 52.4   |           | 0.67                | 5.00       | ug/L     |
| 191-24-2                  | Benzo(g,h,i)perylene       | 52.7   |           | 0.69                | 5.00       | ug/L     |
| 95-94-3                   | 1,2,4,5-Tetrachlorobenzene | 53.7   |           | 0.52                | 5.00       | ug/L     |
| 123-91-1                  | 1,4-Dioxane                | 40.6   |           | 1.00                | 5.00       | ug/L     |
| 58-90-2                   | 2,3,4,6-Tetrachlorophenol  | 54.3   |           | 0.72                | 5.00       | ug/L     |
| <b>SURROGATES</b>         |                            |        |           |                     |            |          |
| 367-12-4                  | 2-Fluorophenol             | 163    |           | 15 (23) - 110 (138) | 108%       | SPK: 150 |
| 13127-88-3                | Phenol-d6                  | 155    |           | 15 (10) - 110 (134) | 103%       | SPK: 150 |
| 4165-60-0                 | Nitrobenzene-d5            | 91.5   |           | 30 (67) - 130 (132) | 91%        | SPK: 100 |
| 321-60-8                  | 2-Fluorobiphenyl           | 90.3   |           | 30 (52) - 130 (132) | 90%        | SPK: 100 |
| 118-79-6                  | 2,4,6-Tribromophenol       | 144    |           | 15 (44) - 110 (137) | 96%        | SPK: 150 |
| 1718-51-0                 | Terphenyl-d14              | 96.5   |           | 30 (42) - 130 (152) | 97%        | SPK: 100 |
| <b>INTERNAL STANDARDS</b> |                            |        |           |                     |            |          |
| 3855-82-1                 | 1,4-Dichlorobenzene-d4     | 15300  | 7.825     |                     |            |          |
| 1146-65-2                 | Naphthalene-d8             | 68900  | 10.61     |                     |            |          |
| 15067-26-2                | Acenaphthene-d10           | 53600  | 14.453    |                     |            |          |
| 1517-22-2                 | Phenanthrene-d10           | 139000 | 17.191    |                     |            |          |
| 1719-03-5                 | Chrysene-d12               | 142000 | 21.415    |                     |            |          |
| 1520-96-3                 | Perylene-d12               | 152000 | 24.388    |                     |            |          |

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

\* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products



# CALIBRATION SUMMARY

Method Path : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\  
 Method File : 8270-BG100225.M  
 Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
 Last Update : Thu Oct 02 16:40:42 2025  
 Response Via : Initial Calibration

## Calibration Files

2.5 =BG064473.D 5 =BG064474.D 10 =BG064481.D 20 =BG064476.D 40 =BG064477.D 50 =BG064478.D 60 =BG064479.D 80 =BG064480.D

| Compound                   | 2.5            | 5     | 10    | 20    | 40    | 50    | 60    | 80    | Avg   | %RSD  |
|----------------------------|----------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| -----                      |                |       |       |       |       |       |       |       |       |       |
| 1) I 1,4-Dichlorobenzen... | -----ISTD----- |       |       |       |       |       |       |       |       |       |
| 2) 1,4-Dioxane             |                | 0.530 | 0.412 | 0.447 | 0.438 | 0.398 | 0.386 | 0.381 | 0.427 | 12.10 |
| 3) Pyridine                |                | 1.225 | 1.151 | 1.147 | 1.236 | 1.102 | 1.095 | 1.086 | 1.149 | 5.33  |
| 4) n-Nitrosodimet...       |                |       | 0.854 | 0.790 | 0.863 | 0.833 | 0.843 | 0.814 | 0.833 | 3.24  |
| 5) S 2-Fluorophenol        |                | 1.066 | 1.018 | 1.053 | 1.195 | 1.071 | 1.108 | 1.053 | 1.081 | 5.26  |
| 6) Aniline                 |                | 1.913 | 1.738 | 1.827 | 2.020 | 1.772 | 1.823 | 1.729 | 1.832 | 5.68  |
| 7) S Phenol-d6             |                | 1.395 | 1.373 | 1.446 | 1.631 | 1.434 | 1.496 | 1.391 | 1.452 | 6.12  |
| 8) 2-Chlorophenol          |                | 1.387 | 1.264 | 1.312 | 1.455 | 1.264 | 1.336 | 1.271 | 1.327 | 5.45  |
| 9) Benzaldehyde            |                |       | 1.437 | 1.039 | 1.527 | 1.146 | 1.548 | 1.037 | 1.289 | 18.76 |
| 10) C Phenol               |                | 1.518 | 1.397 | 1.530 | 1.683 | 1.517 | 1.583 | 1.496 | 1.532 | 5.68  |
| 11) bis(2-Chloroet...      |                | 1.117 | 0.953 | 1.002 | 1.134 | 1.000 | 1.064 | 0.992 | 1.038 | 6.61  |
| 12) 1,3-Dichlorobe...      |                | 1.558 | 1.459 | 1.396 | 1.582 | 1.385 | 1.418 | 1.363 | 1.452 | 5.97  |
| 13) C 1,4-Dichlorobe...    |                | 1.506 | 1.435 | 1.452 | 1.581 | 1.421 | 1.421 | 1.383 | 1.457 | 4.57  |
| 14) 1,2-Dichlorobe...      |                | 1.583 | 1.389 | 1.434 | 1.533 | 1.377 | 1.399 | 1.335 | 1.436 | 6.24  |
| 15) Benzyl Alcohol         |                |       | 1.224 | 1.300 | 1.447 | 1.342 | 1.399 | 1.312 | 1.337 | 5.87  |
| 16) 2,2'-oxybis(1-...      |                | 1.720 | 1.610 | 1.657 | 1.804 | 1.650 | 1.708 | 1.558 | 1.672 | 4.80  |
| 17) 2-Methylphenol         |                | 1.105 | 1.063 | 1.087 | 1.215 | 1.076 | 1.143 | 1.077 | 1.109 | 4.81  |
| 18) Hexachloroethane       |                | 0.586 | 0.630 | 0.534 | 0.604 | 0.562 | 0.566 | 0.544 | 0.575 | 5.87  |
| 19) P n-Nitroso-di-n...    | 0.813          | 1.008 | 0.913 | 1.006 | 1.096 | 0.959 | 1.048 | 0.926 | 0.971 | 9.10  |
| 20) 3+4-Methylphenols      |                |       | 1.472 | 1.510 | 1.700 | 1.492 | 1.622 | 1.494 | 1.548 | 5.91  |
| -----                      |                |       |       |       |       |       |       |       |       |       |
| 21) I Naphthalene-d8       | -----ISTD----- |       |       |       |       |       |       |       |       |       |
| 22) Acetophenone           |                | 0.476 | 0.450 | 0.477 | 0.516 | 0.491 | 0.495 | 0.486 | 0.484 | 4.21  |
| 23) S Nitrobenzene-d5      |                | 0.360 | 0.352 | 0.378 | 0.396 | 0.369 | 0.381 | 0.369 | 0.372 | 3.84  |
| 24) Nitrobenzene           |                | 0.403 | 0.348 | 0.352 | 0.395 | 0.364 | 0.377 | 0.358 | 0.371 | 5.73  |
| 25) Isophorone             |                | 0.672 | 0.654 | 0.674 | 0.721 | 0.660 | 0.686 | 0.650 | 0.674 | 3.59  |
| 26) C 2-Nitrophenol        |                | 0.197 | 0.169 | 0.181 | 0.192 | 0.185 | 0.188 | 0.181 | 0.185 | 4.93  |
| 27) 2,4-Dimethylph...      |                | 0.277 | 0.257 | 0.279 | 0.299 | 0.285 | 0.298 | 0.281 | 0.282 | 4.97  |
| 28) bis(2-Chloroet...      |                | 0.320 | 0.320 | 0.361 | 0.375 | 0.336 | 0.352 | 0.341 | 0.343 | 5.92  |
| 29) C 2,4-Dichloroph...    |                | 0.323 | 0.305 | 0.321 | 0.343 | 0.322 | 0.336 | 0.332 | 0.326 | 3.78  |
| 30) 1,2,4-Trichlor...      |                | 0.383 | 0.333 | 0.360 | 0.377 | 0.339 | 0.349 | 0.347 | 0.356 | 5.27  |
| 31) Naphthalene            |                | 1.062 | 0.980 | 1.013 | 1.107 | 0.995 | 1.041 | 1.011 | 1.030 | 4.26  |
| 32) Benzoic acid           |                |       | 0.149 | 0.198 | 0.230 | 0.240 | 0.239 | 0.243 | 0.216 | 17.02 |
| 33) 4-Chloroaniline        |                | 0.392 | 0.398 | 0.373 | 0.417 | 0.388 | 0.406 | 0.392 | 0.395 | 3.53  |
| 34) C Hexachlorobuta...    |                | 0.331 | 0.289 | 0.308 | 0.316 | 0.296 | 0.300 | 0.291 | 0.304 | 4.94  |
| 35) Caprolactam            |                |       | 0.108 | 0.120 | 0.115 | 0.111 | 0.110 | 0.107 | 0.112 | 4.36  |
| 36) C 4-Chloro-3-met...    |                | 0.395 | 0.381 | 0.392 | 0.418 | 0.390 | 0.405 | 0.395 | 0.397 | 3.01  |
| 37) 2-Methylnaphth...      |                | 0.783 | 0.771 | 0.790 | 0.843 | 0.778 | 0.799 | 0.770 | 0.791 | 3.18  |
| 38) 1-Methylnaphth...      |                | 0.837 | 0.782 | 0.782 | 0.825 | 0.750 | 0.784 | 0.758 | 0.788 | 4.08  |

Method Path : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\  
 Method File : 8270-BG100225.M

|       |                   |                |       |       |       |       |       |       |       |       |  |
|-------|-------------------|----------------|-------|-------|-------|-------|-------|-------|-------|-------|--|
| 39) I | Acenaphthene-d10  | -----ISTD----- |       |       |       |       |       |       |       |       |  |
| 40)   | 1,2,4,5-Tetrac... | 0.622          | 0.604 | 0.622 | 0.708 | 0.640 | 0.678 | 0.646 | 0.646 | 5.60  |  |
| 41) P | Hexachlorocycl... | 0.279          | 0.296 | 0.363 | 0.356 | 0.378 | 0.356 | 0.338 |       | 11.95 |  |
| 42) S | 2,4,6-Tribromo... | 0.364          | 0.352 | 0.367 | 0.381 | 0.364 | 0.361 | 0.349 | 0.363 | 2.89  |  |
| 43) C | 2,4,6-Trichlor... | 0.391          | 0.390 | 0.396 | 0.445 | 0.416 | 0.436 | 0.414 | 0.413 | 5.30  |  |
| 44)   | 2,4,5-Trichlor... | 0.451          | 0.452 | 0.430 | 0.486 | 0.465 | 0.473 | 0.470 | 0.461 | 3.93  |  |
| 45) S | 2-Fluorobiphenyl  | 1.355          | 1.338 | 1.346 | 1.434 | 1.325 | 1.389 | 1.310 | 1.357 | 3.11  |  |
| 46)   | 1,1'-Biphenyl     | 1.439          | 1.388 | 1.417 | 1.521 | 1.434 | 1.484 | 1.387 | 1.438 | 3.42  |  |
| 47)   | 2-Chloronaphth... | 1.006          | 1.104 | 1.046 | 1.153 | 1.056 | 1.106 | 1.046 | 1.074 | 4.61  |  |
| 48)   | 2-Nitroaniline    | 0.301          | 0.362 | 0.371 | 0.397 | 0.386 | 0.385 | 0.365 | 0.367 | 8.60  |  |
| 49)   | Acenaphthylene    | 1.591          | 1.534 | 1.573 | 1.704 | 1.574 | 1.638 | 1.546 | 1.594 | 3.68  |  |
| 50)   | Dimethylphthalate | 1.564          | 1.584 | 1.582 | 1.624 | 1.531 | 1.586 | 1.493 | 1.566 | 2.72  |  |
| 51)   | 2,6-Dinitrotol... | 0.292          | 0.326 | 0.335 | 0.337 | 0.333 | 0.345 | 0.316 | 0.326 | 5.43  |  |
| 52) C | Acenaphthene      | 1.108          | 1.099 | 1.086 | 1.159 | 1.106 | 1.138 | 1.058 | 1.108 | 2.98  |  |
| 53)   | 3-Nitroaniline    | 0.279          | 0.294 | 0.300 | 0.319 | 0.315 | 0.303 | 0.296 | 0.301 | 4.50  |  |
| 54) P | 2,4-Dinitrophenol | 0.265          | 0.270 | 0.207 | 0.228 | 0.229 | 0.220 | 0.237 |       | 10.76 |  |
| 55)   | Dibenzofuran      | 1.837          | 1.810 | 1.800 | 1.909 | 1.821 | 1.872 | 1.753 | 1.829 | 2.77  |  |
| 56) P | 4-Nitrophenol     | 0.192          | 0.227 | 0.260 | 0.256 | 0.253 | 0.232 | 0.237 |       | 10.85 |  |
| 57)   | 2,4-Dinitrotol... | 0.481          | 0.499 | 0.488 | 0.527 | 0.513 | 0.506 | 0.477 | 0.499 | 3.60  |  |
| 58)   | Fluorene          | 1.518          | 1.511 | 1.535 | 1.601 | 1.488 | 1.532 | 1.458 | 1.520 | 2.91  |  |
| 59)   | 2,3,4,6-Tetrac... | 0.438          | 0.447 | 0.475 | 0.505 | 0.475 | 0.494 | 0.457 | 0.470 | 5.16  |  |
| 60)   | Diethylphthalate  | 1.747          | 1.692 | 1.679 | 1.730 | 1.675 | 1.663 | 1.578 | 1.680 | 3.25  |  |
| 61)   | 4-Chlorophenyl... | 0.866          | 0.809 | 0.839 | 0.868 | 0.817 | 0.819 | 0.794 | 0.830 | 3.42  |  |
| 62)   | 4-Nitroaniline    | 0.229          | 0.296 | 0.331 | 0.334 | 0.345 | 0.340 | 0.320 | 0.314 | 12.96 |  |
| 63)   | Azobenzene        | 1.339          | 1.299 | 1.325 | 1.346 | 1.295 | 1.289 | 1.216 | 1.301 | 3.37  |  |
| 64) I | Phenanthrene-d10  | -----ISTD----- |       |       |       |       |       |       |       |       |  |
| 65)   | 4,6-Dinitro-2-... | 0.125          | 0.139 | 0.145 | 0.138 | 0.145 | 0.144 | 0.139 |       | 5.58  |  |
| 66) c | n-Nitrosodiphe... | 0.540          | 0.519 | 0.534 | 0.569 | 0.508 | 0.541 | 0.529 | 0.534 | 3.62  |  |
| 67)   | 4-Bromophenyl-... | 0.235          | 0.229 | 0.227 | 0.253 | 0.224 | 0.237 | 0.241 | 0.235 | 4.21  |  |
| 68)   | Hexachlorobenzene | 0.256          | 0.261 | 0.272 | 0.288 | 0.260 | 0.280 | 0.270 | 0.269 | 4.36  |  |
| 69)   | Atrazine          | 0.237          | 0.240 | 0.247 | 0.250 | 0.232 | 0.239 | 0.219 | 0.238 | 4.26  |  |
| 70) C | Pentachlorophenol | 0.118          | 0.146 | 0.169 | 0.166 | 0.170 | 0.175 | 0.157 |       | 13.66 |  |
| 71)   | Phenanthrene      | 1.072          | 1.022 | 1.055 | 1.111 | 1.002 | 1.050 | 1.010 | 1.046 | 3.65  |  |
| 72)   | Anthracene        | 1.045          | 1.028 | 1.044 | 1.130 | 1.001 | 1.068 | 1.029 | 1.049 | 3.93  |  |
| 73)   | Carbazole         | 0.883          | 0.895 | 0.904 | 0.987 | 0.877 | 0.921 | 0.880 | 0.907 | 4.24  |  |
| 74)   | Di-n-butylphth... | 1.096          | 1.079 | 1.105 | 1.189 | 1.076 | 1.131 | 1.076 | 1.107 | 3.71  |  |
| 75) C | Fluoranthene      | 1.269          | 1.226 | 1.272 | 1.330 | 1.192 | 1.253 | 1.204 | 1.250 | 3.77  |  |
| 76) I | Chrysene-d12      | -----ISTD----- |       |       |       |       |       |       |       |       |  |
| 77)   | Benzidine         | 0.393          | 0.424 | 0.649 | 0.540 | 0.621 | 0.359 | 0.498 |       | 24.67 |  |
| 78)   | Pyrene            | 1.295          | 1.329 | 1.316 | 1.392 | 1.242 | 1.281 | 1.230 | 1.298 | 4.24  |  |
| 79) S | Terphenyl-d14     | 1.067          | 1.061 | 1.069 | 1.123 | 1.030 | 1.044 | 1.003 | 1.057 | 3.55  |  |
| 80)   | Butylbenzylpht... | 0.437          | 0.438 | 0.464 | 0.497 | 0.447 | 0.479 | 0.458 | 0.460 | 4.78  |  |
| 81)   | Benzo(a)anthra... | 1.280          | 1.239 | 1.285 | 1.353 | 1.238 | 1.291 | 1.253 | 1.277 | 3.15  |  |
| 82)   | 3,3'-Dichlorob... | 0.418          | 0.431 | 0.458 | 0.423 | 0.443 | 0.420 | 0.432 |       | 3.58  |  |
| 83)   | Chrysene          | 1.226          | 1.200 | 1.195 | 1.297 | 1.173 | 1.210 | 1.178 | 1.211 | 3.48  |  |
| 84)   | Bis(2-ethylhex... | 0.605          | 0.637 | 0.640 | 0.692 | 0.641 | 0.670 | 0.658 | 0.649 | 4.26  |  |
| 85) c | Di-n-octyl pht... | 1.055          | 1.050 | 1.153 | 1.073 | 1.125 | 1.089 | 1.091 |       | 3.74  |  |

Method Path : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\  
Method File : 8270-BG100225.M

|       |                   |                |       |       |       |       |       |       |       |      |
|-------|-------------------|----------------|-------|-------|-------|-------|-------|-------|-------|------|
| 86) I | Perylene-d12      | -----ISTD----- |       |       |       |       |       |       |       |      |
| 87)   | Indeno(1,2,3-c... | 1.368          | 1.320 | 1.378 | 1.493 | 1.387 | 1.431 | 1.368 | 1.392 | 3.98 |
| 88)   | Benzo(b)fluora... | 1.077          | 1.084 | 1.171 | 1.212 | 1.142 | 1.169 | 1.109 | 1.138 | 4.40 |
| 89)   | Benzo(k)fluora... | 1.139          | 1.082 | 1.145 | 1.233 | 1.123 | 1.156 | 1.142 | 1.146 | 3.97 |
| 90) C | Benzo(a)pyrene    | 1.017          | 0.999 | 1.059 | 1.122 | 1.031 | 1.075 | 1.037 | 1.049 | 3.91 |
| 91)   | Dibenzo(a,h)an... | 1.047          | 1.053 | 1.121 | 1.190 | 1.115 | 1.151 | 1.116 | 1.113 | 4.55 |
| 92)   | Benzo(g,h,i)pe... | 1.059          | 1.053 | 1.088 | 1.153 | 1.069 | 1.104 | 1.048 | 1.082 | 3.41 |

-----  
(#) = Out of Range

7C

SEMIVOLATILE CONTINUING CALIBRATION CHECK

|                 |                   |                        |                              |
|-----------------|-------------------|------------------------|------------------------------|
| Lab Name:       | <u>Alliance</u>   | Contract:              | <u>GENV01</u>                |
| Lab Code:       | <u>ACE</u>        | SDG No.:               | <u>Q3273</u>                 |
| Instrument ID:  | <u>BNA_G</u>      | Calibration Date/Time: | <u>10/03/2025 13:33</u>      |
| Lab File ID:    | <u>BG064485.D</u> | Init. Calib. Date(s):  | <u>10/02/2025 10/02/2025</u> |
| EPA Sample No.: | <u>SSTDCCC040</u> | Init. Calib. Time(s):  | <u>09:01 15:49</u>           |
| GC Column:      | <u>ZB-GR</u>      | ID:                    | <u>0.25</u> (mm)             |

| COMPOUND                    | RRF   | RRF040 | MIN RRF | %D   | MAX%D |
|-----------------------------|-------|--------|---------|------|-------|
| 2-Fluorophenol              | 1.081 | 1.209  |         | 11.8 |       |
| Benzaldehyde                | 1.290 | 1.512  |         | 17.2 |       |
| Phenol-d6                   | 1.452 | 1.602  |         | 10.3 |       |
| Phenol                      | 1.532 | 1.694  |         | 10.6 | 20.0  |
| bis(2-Chloroethyl)ether     | 1.038 | 1.112  |         | 7.1  |       |
| 2-Chlorophenol              | 1.327 | 1.425  |         | 7.4  |       |
| 2-Methylphenol              | 1.109 | 1.211  |         | 9.2  |       |
| 2,2-oxybis(1-Chloropropane) | 1.672 | 1.832  |         | 9.6  |       |
| Acetophenone                | 0.484 | 0.523  |         | 8.1  |       |
| 3+4-Methylphenols           | 1.548 | 1.704  |         | 10.1 |       |
| n-Nitroso-di-n-propylamine  | 0.971 | 1.074  | 0.050   | 10.6 |       |
| Nitrobenzene-d5             | 0.372 | 0.400  |         | 7.5  |       |
| Hexachloroethane            | 0.575 | 0.621  |         | 8.0  |       |
| Nitrobenzene                | 0.371 | 0.391  |         | 5.4  |       |
| Isophorone                  | 0.674 | 0.706  |         | 4.7  |       |
| 2-Nitrophenol               | 0.185 | 0.195  |         | 5.4  | 20.0  |
| 2,4-Dimethylphenol          | 0.282 | 0.301  |         | 6.7  |       |
| bis(2-Chloroethoxy)methane  | 0.343 | 0.360  |         | 5.0  |       |
| 2,4-Dichlorophenol          | 0.326 | 0.348  |         | 6.7  | 20.0  |
| Naphthalene                 | 1.030 | 1.093  |         | 6.1  |       |
| 4-Chloroaniline             | 0.395 | 0.420  |         | 6.3  |       |
| Hexachlorobutadiene         | 0.304 | 0.315  |         | 3.6  | 20.0  |
| Caprolactam                 | 0.112 | 0.122  |         | 8.9  |       |
| 4-Chloro-3-methylphenol     | 0.397 | 0.428  |         | 7.8  | 20.0  |
| 2-Methylnaphthalene         | 0.791 | 0.823  |         | 4.0  |       |
| Hexachlorocyclopentadiene   | 0.338 | 0.368  | 0.050   | 8.9  |       |
| 2,4,6-Trichlorophenol       | 0.413 | 0.437  |         | 5.8  | 20.0  |
| 2-Fluorobiphenyl            | 1.357 | 1.436  |         | 5.8  |       |
| 2,4,5-Trichlorophenol       | 0.461 | 0.511  |         | 10.8 |       |
| 1,1-Biphenyl                | 1.438 | 1.499  |         | 4.2  |       |
| 2-Chloronaphthalene         | 1.074 | 1.152  |         | 7.3  |       |
| 2-Nitroaniline              | 0.367 | 0.393  |         | 7.1  |       |
| Dimethylphthalate           | 1.566 | 1.632  |         | 4.2  |       |
| Acenaphthylene              | 1.594 | 1.697  |         | 6.5  |       |
| 2,6-Dinitrotoluene          | 0.326 | 0.352  |         | 8.0  |       |
| 3-Nitroaniline              | 0.301 | 0.332  |         | 10.3 |       |
| Acenaphthene                | 1.108 | 1.174  |         | 6.0  | 20.0  |
| 2,4-Dinitrophenol           | 0.237 | 0.222  | 0.050   | -6.3 |       |
| 4-Nitrophenol               | 0.237 | 0.271  | 0.050   | 14.3 |       |



7C

SEMIVOLATILE CONTINUING CALIBRATION CHECK

|                 |                   |                        |                              |
|-----------------|-------------------|------------------------|------------------------------|
| Lab Name:       | <u>Alliance</u>   | Contract:              | <u>GENV01</u>                |
| Lab Code:       | <u>ACE</u>        | SDG No.:               | <u>Q3273</u>                 |
| Instrument ID:  | <u>BNA_G</u>      | Calibration Date/Time: | <u>10/03/2025 13:33</u>      |
| Lab File ID:    | <u>BG064485.D</u> | Init. Calib. Date(s):  | <u>10/02/2025 10/02/2025</u> |
| EPA Sample No.: | <u>SSTDCCC040</u> | Init. Calib. Time(s):  | <u>09:01 15:49</u>           |
| GC Column:      | <u>ZB-GR</u>      | ID:                    | <u>0.25</u> (mm)             |

| COMPOUND                   | RRF   | RRF040 | MIN<br>RRF | %D   | MAX%D |
|----------------------------|-------|--------|------------|------|-------|
| Dibenzofuran               | 1.829 | 1.939  |            | 6.0  |       |
| 2,4-Dinitrotoluene         | 0.499 | 0.537  |            | 7.6  |       |
| Diethylphthalate           | 1.680 | 1.758  |            | 4.6  |       |
| 4-Chlorophenyl-phenylether | 0.830 | 0.863  |            | 4.0  |       |
| Fluorene                   | 1.520 | 1.622  |            | 6.7  |       |
| 4-Nitroaniline             | 0.314 | 0.357  |            | 13.7 |       |
| 4,6-Dinitro-2-methylphenol | 0.139 | 0.147  |            | 5.8  |       |
| n-Nitrosodiphenylamine     | 0.534 | 0.559  |            | 4.7  | 20.0  |
| 2,4,6-Tribromophenol       | 0.363 | 0.380  |            | 4.7  |       |
| 4-Bromophenyl-phenylether  | 0.235 | 0.250  |            | 6.4  |       |
| Hexachlorobenzene          | 0.269 | 0.281  |            | 4.5  |       |
| Atrazine                   | 0.238 | 0.251  |            | 5.5  |       |
| Pentachlorophenol          | 0.157 | 0.168  |            | 7.0  | 20.0  |
| Phenanthrene               | 1.046 | 1.113  |            | 6.4  |       |
| Anthracene                 | 1.049 | 1.103  |            | 5.1  |       |
| Carbazole                  | 0.907 | 0.978  |            | 7.8  |       |
| Di-n-butylphthalate        | 1.107 | 1.201  |            | 8.5  |       |
| Fluoranthene               | 1.250 | 1.372  |            | 9.8  | 20.0  |
| Pyrene                     | 1.298 | 1.371  |            | 5.6  |       |
| Terphenyl-d14              | 1.057 | 1.121  |            | 6.1  |       |
| Butylbenzylphthalate       | 0.460 | 0.487  |            | 5.9  |       |
| 3,3-Dichlorobenzidine      | 0.432 | 0.460  |            | 6.5  |       |
| Benzo (a) anthracene       | 1.277 | 1.353  |            | 6.0  |       |
| Chrysene                   | 1.211 | 1.274  |            | 5.2  |       |
| Bis(2-ethylhexyl)phthalate | 0.649 | 0.681  |            | 4.9  |       |
| Di-n-octyl phthalate       | 1.091 | 1.132  |            | 3.8  | 20.0  |
| Benzo (b) fluoranthene     | 1.138 | 1.230  |            | 8.1  |       |
| Benzo (k) fluoranthene     | 1.146 | 1.207  |            | 5.3  |       |
| Benzo (a) pyrene           | 1.049 | 1.132  |            | 7.9  | 20.0  |
| Indeno (1,2,3-cd) pyrene   | 1.392 | 1.487  |            | 6.8  |       |
| Dibenzo (a,h) anthracene   | 1.113 | 1.209  |            | 8.6  |       |
| Benzo (g,h,i) perylene     | 1.082 | 1.155  |            | 6.7  |       |
| 1,2,4,5-Tetrachlorobenzene | 0.646 | 0.677  |            | 4.8  |       |
| 1,4-Dioxane                | 0.427 | 0.434  |            | 1.6  | 20.0  |
| 2,3,4,6-Tetrachlorophenol  | 0.470 | 0.497  |            | 5.7  |       |

All other compounds must meet a minimum RRF of 0.010.



# SAMPLE RAW DATA

A

B

C

D

E

F

G

H

I

J

K

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Time: Oct 04 00:00:10 2025  
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
QLast Update : Thu Oct 02 16:40:42 2025  
Response via : Initial Calibration

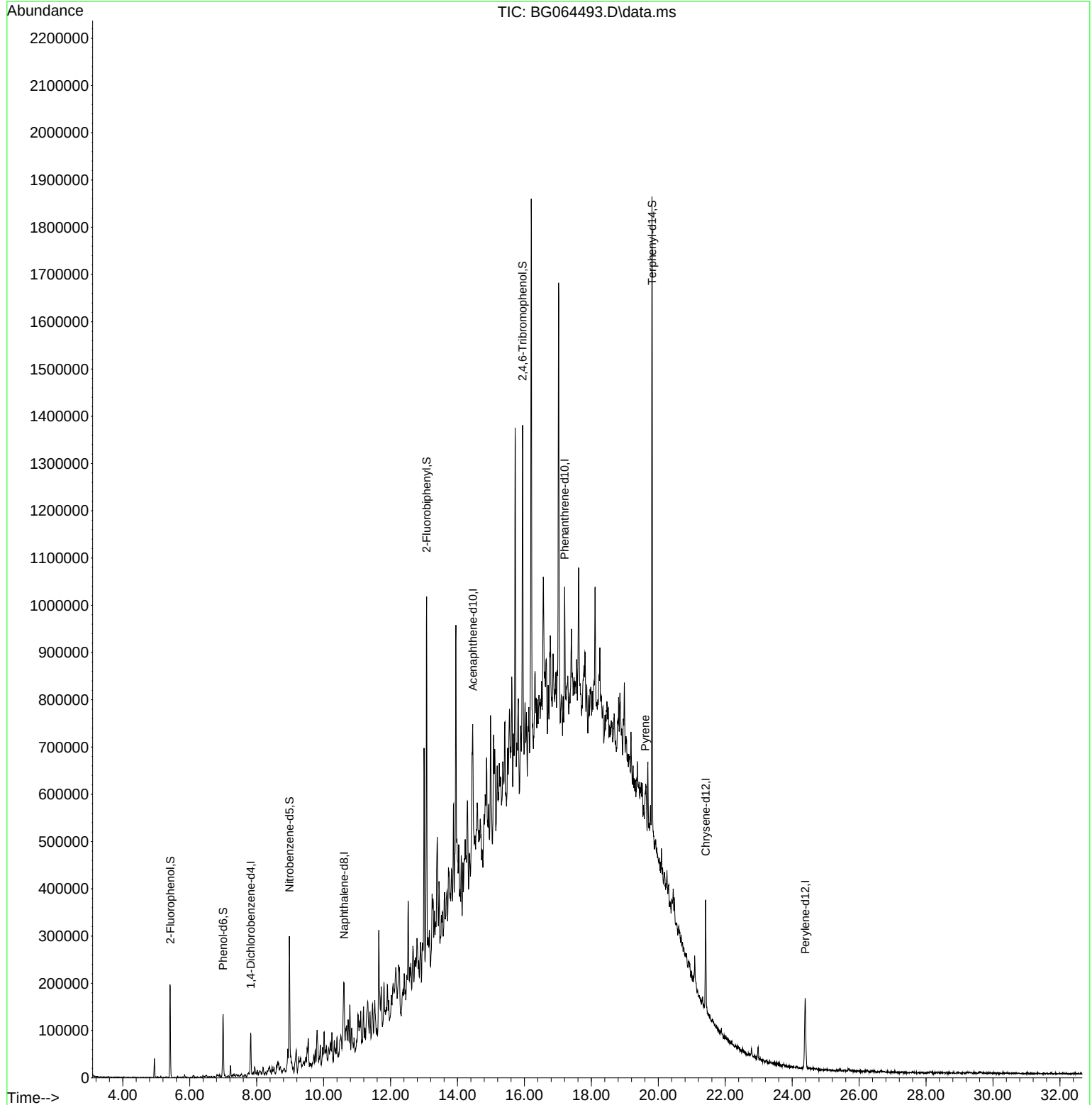
| Compound                    | R.T.   | QIon | Response | Conc    | Units | Dev(Min) |
|-----------------------------|--------|------|----------|---------|-------|----------|
| -----                       |        |      |          |         |       |          |
| Internal Standards          |        |      |          |         |       |          |
| 1) 1,4-Dichlorobenzene-d4   | 7.820  | 152  | 20058    | 20.000  | ng    | 0.00     |
| 21) Naphthalene-d8          | 10.611 | 136  | 82373    | 20.000  | ng    | # 0.00   |
| 39) Acenaphthene-d10        | 14.459 | 164  | 60802    | 20.000  | ng    | 0.01     |
| 64) Phenanthrene-d10        | 17.197 | 188  | 129947   | 20.000  | ng    | # 0.01   |
| 76) Chrysene-d12            | 21.416 | 240  | 136555   | 20.000  | ng    | # 0.00   |
| 86) Perylene-d12            | 24.389 | 264  | 149983   | 20.000  | ng    | # 0.00   |
|                             |        |      |          |         |       |          |
| System Monitoring Compounds |        |      |          |         |       |          |
| 5) 2-Fluorophenol           | 5.417  | 112  | 73920    | 68.203  | ng    | 0.01     |
| 7) Phenol-d6                | 6.997  | 99   | 59110    | 40.584  | ng    | 0.01     |
| 23) Nitrobenzene-d5         | 8.977  | 82   | 138717   | 90.549  | ng    | 0.01     |
| 42) 2,4,6-Tribromophenol    | 15.946 | 330  | 147739   | 134.050 | ng    | 0.01     |
| 45) 2-Fluorobiphenyl        | 13.079 | 172  | 378313   | 91.713  | ng    | 0.01     |
| 79) Terphenyl-d14           | 19.812 | 244  | 622795   | 86.312  | ng    | 0.00     |
|                             |        |      |          |         |       |          |
| Target Compounds            |        |      |          |         |       | Qvalue   |
| 78) Pyrene                  | 19.606 | 202  | 41244    | 4.654   | ng    | # 91     |
| -----                       |        |      |          |         |       |          |

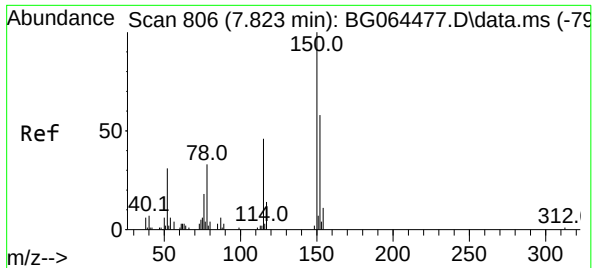
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Time: Oct 04 00:00:10 2025  
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
QLast Update : Thu Oct 02 16:40:42 2025  
Response via : Initial Calibration

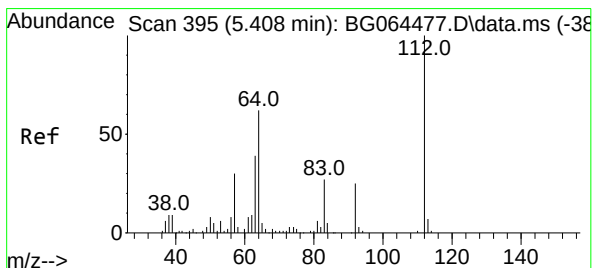
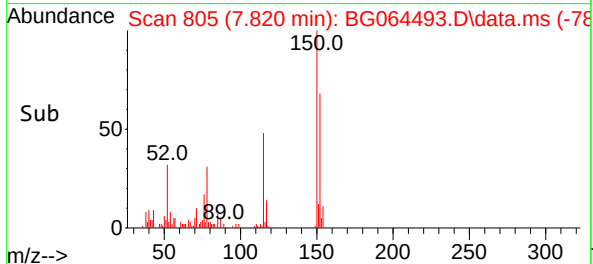
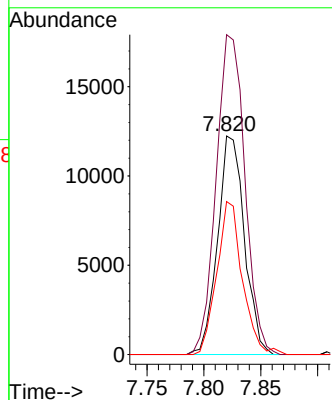
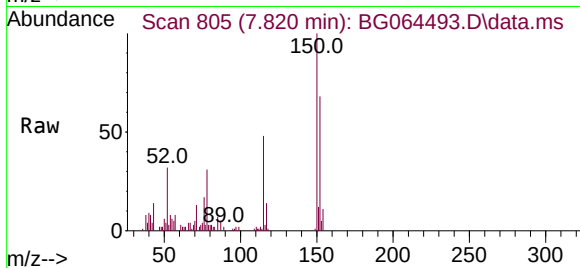




#1  
1,4-Dichlorobenzene-d4  
Concen: 20.000 ng  
RT: 7.820 min Scan# 806  
Delta R.T. -0.000 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

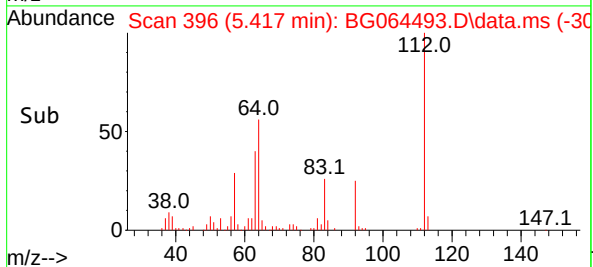
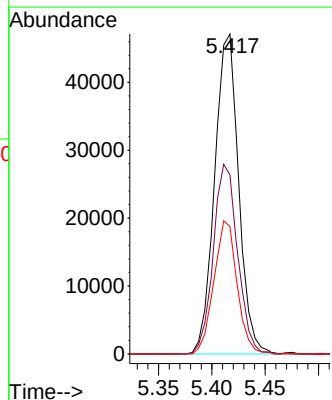
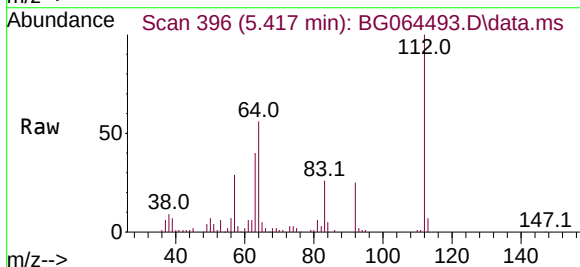
Instrument :  
BNA\_G  
ClientSampleId :  
MW1

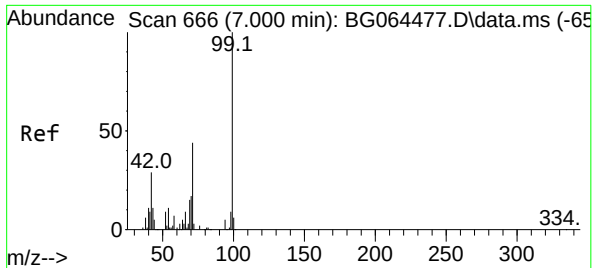
| Tgt Ion   | 152   | Resp  | 20058 |
|-----------|-------|-------|-------|
| Ion Ratio | Lower | Upper |       |
| 152       | 100   |       |       |
| 150       | 146.3 | 138.6 | 207.8 |
| 115       | 70.0  | 63.4  | 95.2  |



#5  
2-Fluorophenol  
Concen: 68.203 ng  
RT: 5.417 min Scan# 396  
Delta R.T. 0.011 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

| Tgt Ion   | 112   | Resp  | 73920 |
|-----------|-------|-------|-------|
| Ion Ratio | Lower | Upper |       |
| 112       | 100   |       |       |
| 64        | 55.9  | 49.8  | 74.8  |
| 63        | 39.8  | 31.3  | 46.9  |

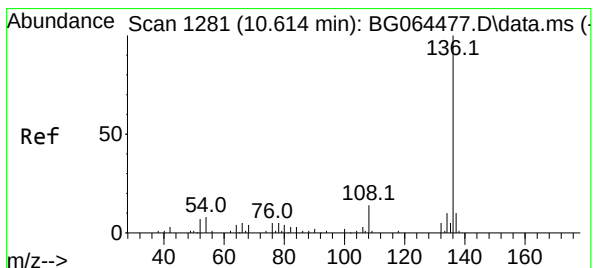
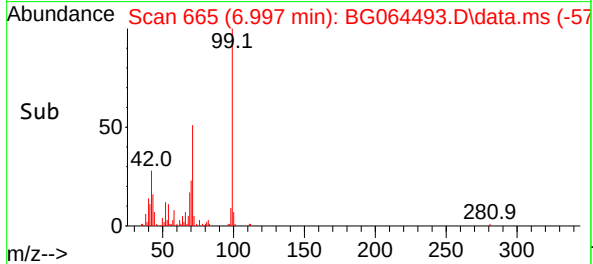
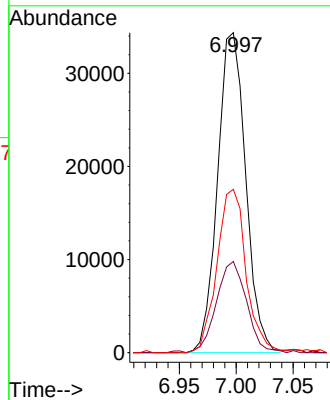
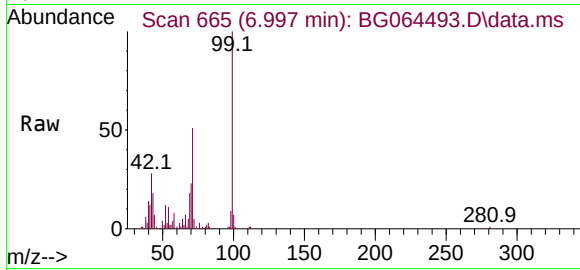




#7  
Phenol-d6  
Concen: 40.584 ng  
RT: 6.997 min Scan# 60  
Delta R.T. 0.011 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

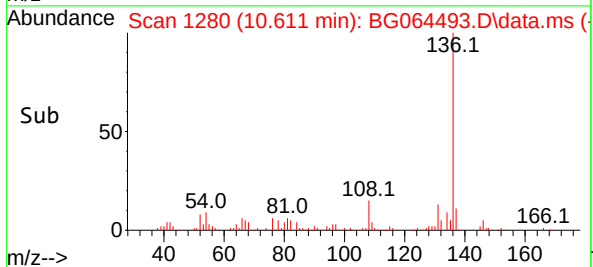
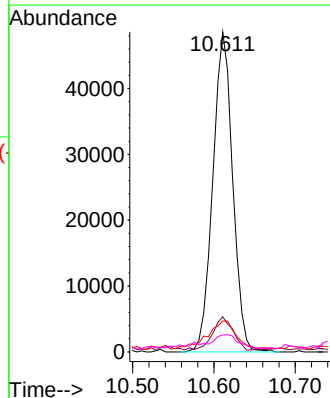
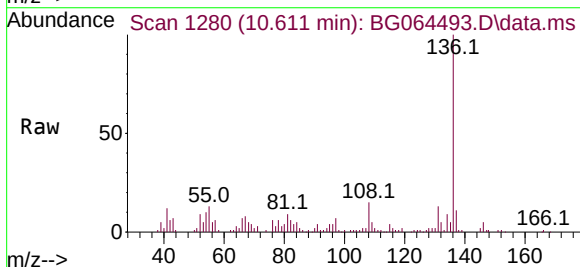
Instrument :  
BNA\_G  
ClientSampleId :  
MW1

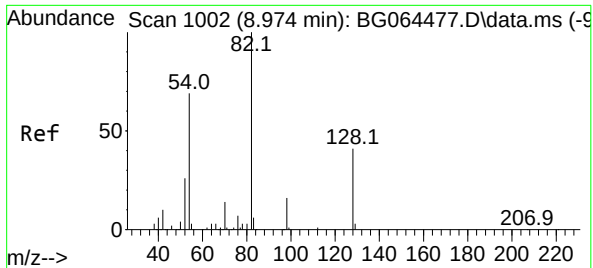
| Tgt Ion | 99   | Ratio | 100  | Lower | Upper |
|---------|------|-------|------|-------|-------|
| 99      | 100  |       |      |       |       |
| 42      | 28.5 |       | 22.9 | 34.3  |       |
| 71      | 51.0 |       | 35.4 | 53.2  |       |



#21  
Naphthalene-d8  
Concen: 20.000 ng  
RT: 10.611 min Scan# 1280  
Delta R.T. 0.005 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

| Tgt Ion | 136  | Ratio | 100 | Lower | Upper |
|---------|------|-------|-----|-------|-------|
| 136     | 100  |       |     |       |       |
| 137     | 11.0 |       | 8.3 | 12.5  |       |
| 54      | 9.6  |       | 6.2 | 9.4#  |       |
| 68      | 5.2  |       | 3.0 | 4.4#  |       |





#23

Nitrobenzene-d5

Concen: 90.549 ng

RT: 8.977 min Scan# 10

Delta R.T. 0.011 min

Lab File: BG064493.D

Acq: 3 Oct 2025 18:59

Instrument :

BNA\_G

ClientSampleId :

MW1

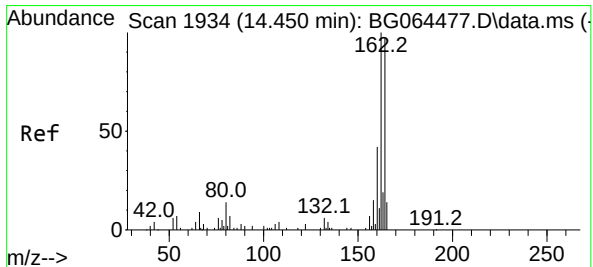
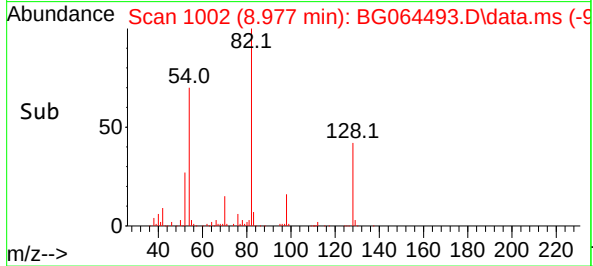
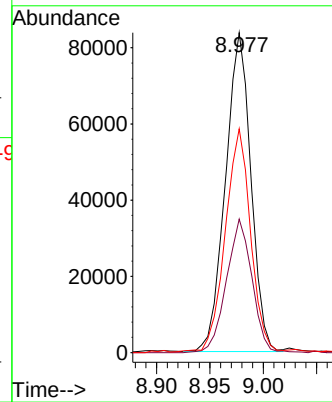
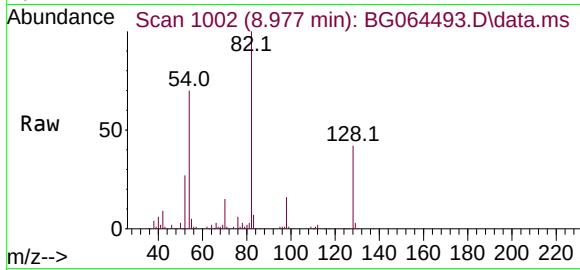
Tgt Ion: 82 Resp: 138717

Ion Ratio Lower Upper

82 100

128 41.7 33.1 49.7

54 69.9 55.4 83.2



#39

Acenaphthene-d10

Concen: 20.000 ng

RT: 14.459 min Scan# 1935

Delta R.T. 0.011 min

Lab File: BG064493.D

Acq: 3 Oct 2025 18:59

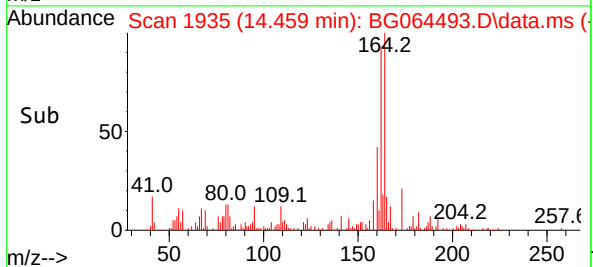
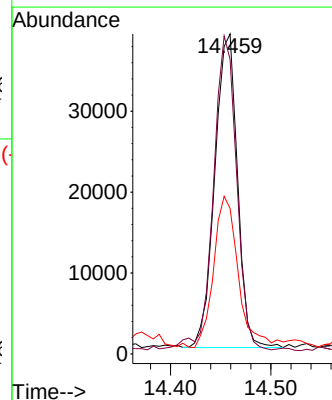
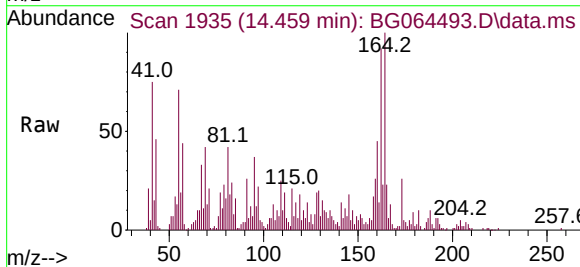
Tgt Ion:164 Resp: 60802

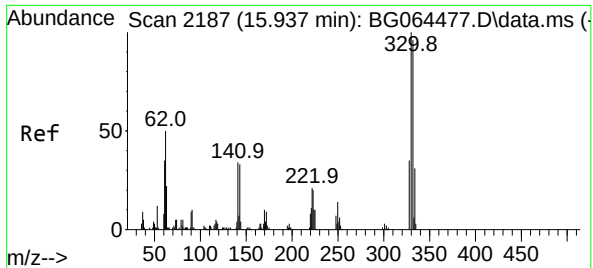
Ion Ratio Lower Upper

164 100

162 92.0 82.2 123.2

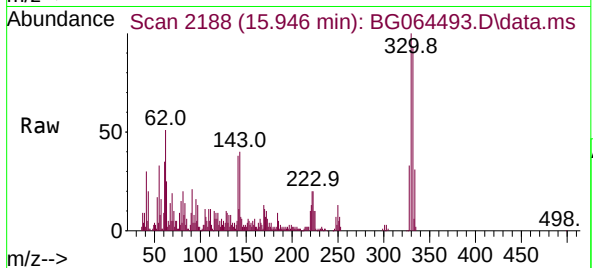
160 45.5 34.3 51.5



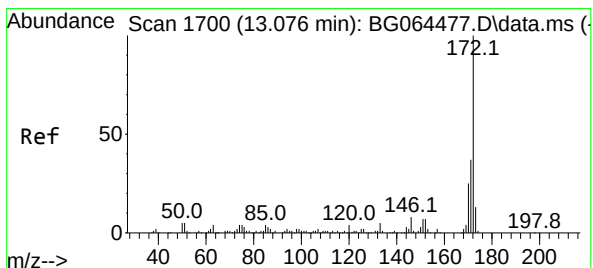
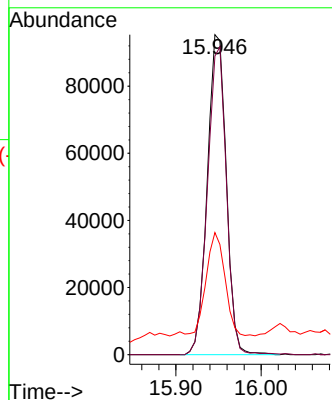
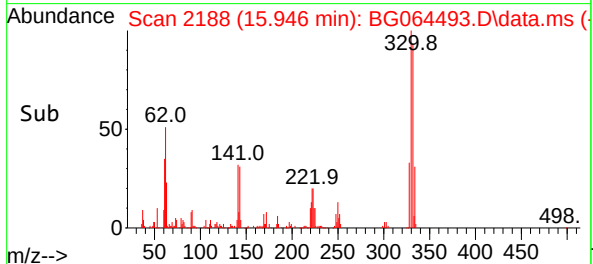


#42  
2,4,6-Tribromophenol  
Concen: 134.050 ng  
RT: 15.946 min Scan# 2187  
Delta R.T. 0.011 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

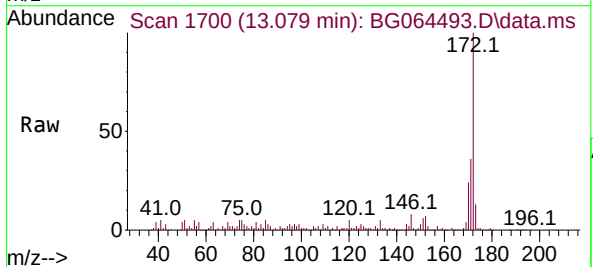
Instrument :  
BNA\_G  
ClientSampleId :  
MW1



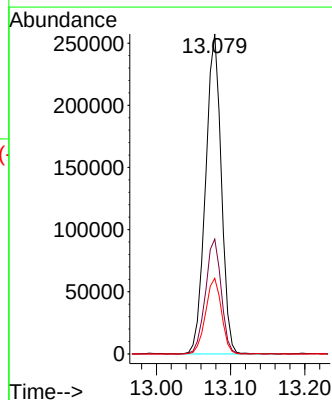
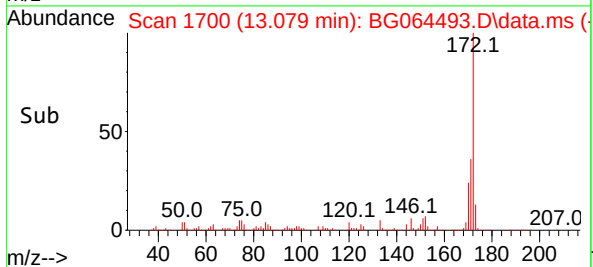
Tgt Ion:330 Resp: 147739  
Ion Ratio Lower Upper  
330 100  
332 95.8 77.4 116.0  
141 34.0 25.8 38.6



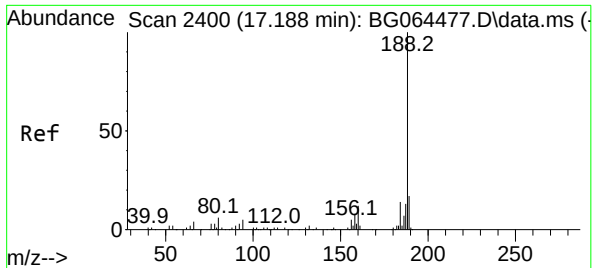
#45  
2-Fluorobiphenyl  
Concen: 91.713 ng  
RT: 13.079 min Scan# 1700  
Delta R.T. 0.011 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59



Tgt Ion:172 Resp: 378313  
Ion Ratio Lower Upper  
172 100  
171 35.9 29.6 44.4  
170 23.6 20.2 30.2



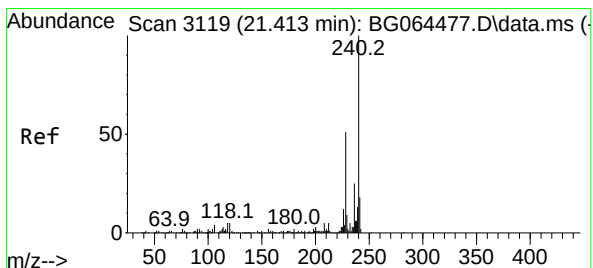
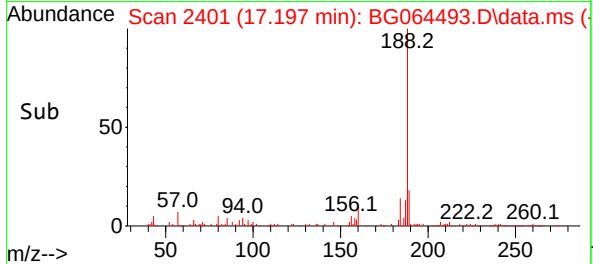
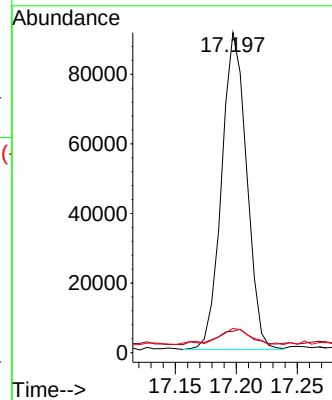
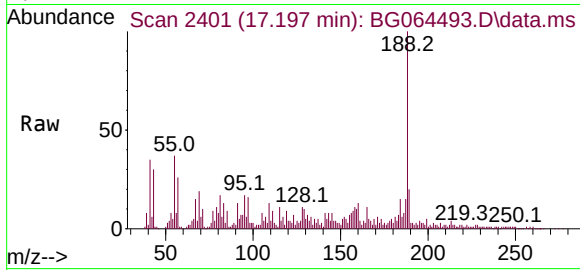




#64  
Phenanthrene-d10  
Concen: 20.000 ng  
RT: 17.197 min Scan# 2401  
Delta R.T. 0.011 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

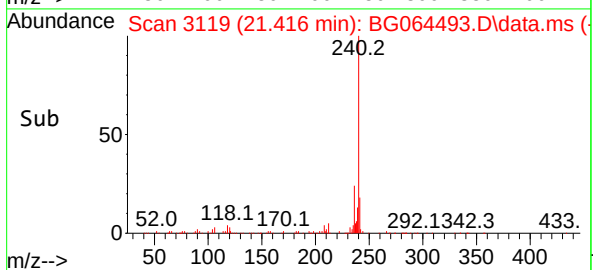
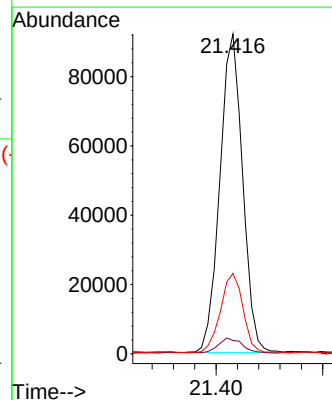
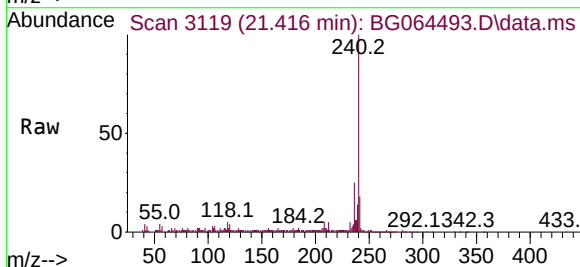
Instrument :  
BNA\_G  
ClientSampleId :  
MW1

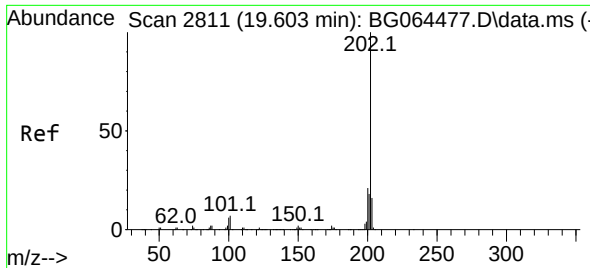
Tgt Ion:188 Resp: 129947  
Ion Ratio Lower Upper  
188 100  
94 6.7 3.7 5.5#  
80 7.6 4.5 6.7#



#76  
Chrysene-d12  
Concen: 20.000 ng  
RT: 21.416 min Scan# 3119  
Delta R.T. 0.006 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

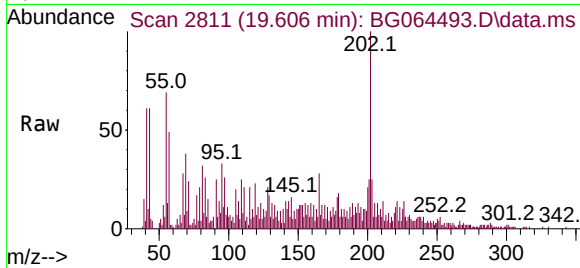
Tgt Ion:240 Resp: 136555  
Ion Ratio Lower Upper  
240 100  
120 4.2 4.3 6.5#  
236 25.2 20.4 30.6



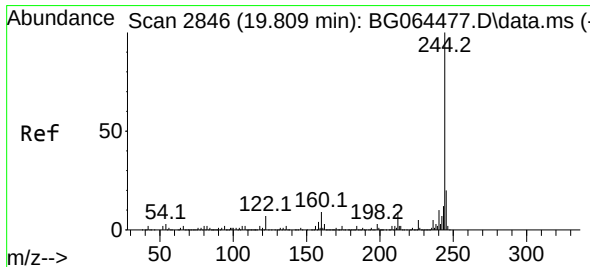
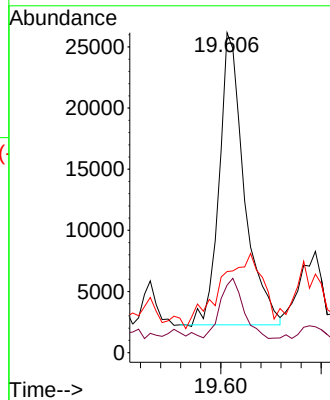
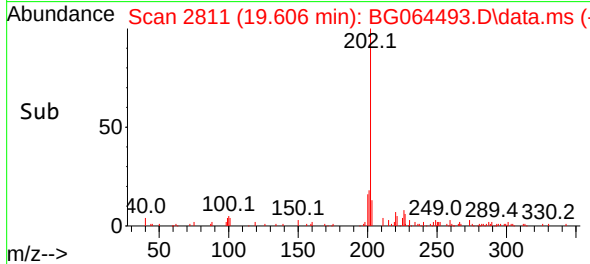


#78  
Pyrene  
Concen: 4.654 ng  
RT: 19.606 min Scan# 2811  
Delta R.T. 0.005 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

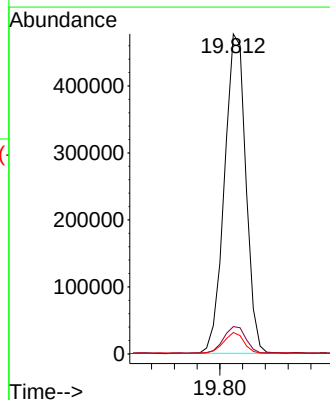
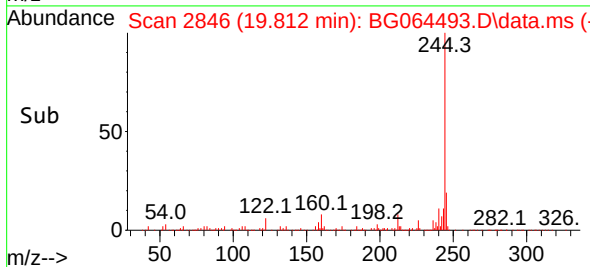
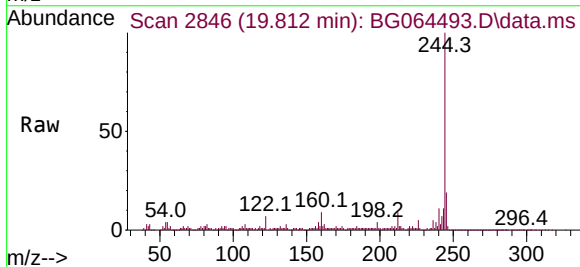


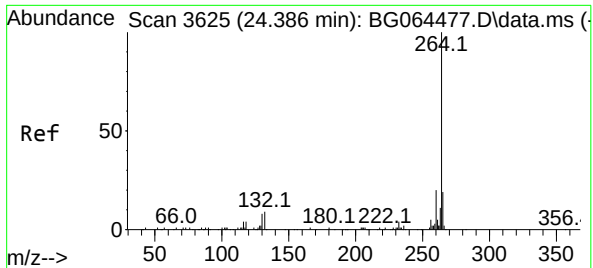
Tgt Ion:202 Resp: 41244  
Ion Ratio Lower Upper  
202 100  
200 21.1 17.0 25.4  
203 25.3 13.0 19.4#



#79  
Terphenyl-d14  
Concen: 86.312 ng  
RT: 19.812 min Scan# 2846  
Delta R.T. 0.005 min  
Lab File: BG064493.D  
Acq: 3 Oct 2025 18:59

Tgt Ion:244 Resp: 622795  
Ion Ratio Lower Upper  
244 100  
212 8.5 7.0 10.6  
122 6.7 5.6 8.4





#86

Perylene-d12

Concen: 20.000 ng

RT: 24.389 min Scan# 30

Delta R.T. -0.000 min

Lab File: BG064493.D

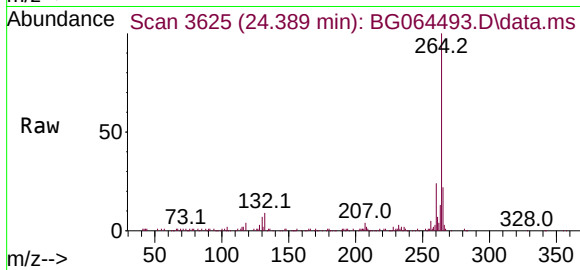
Acq: 3 Oct 2025 18:59

Instrument :

BNA\_G

ClientSampleId :

MW1



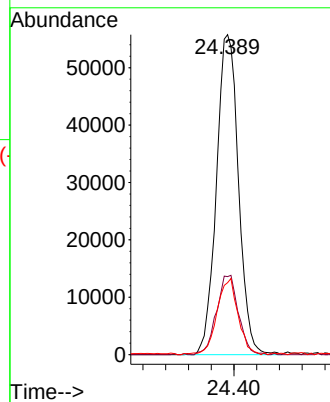
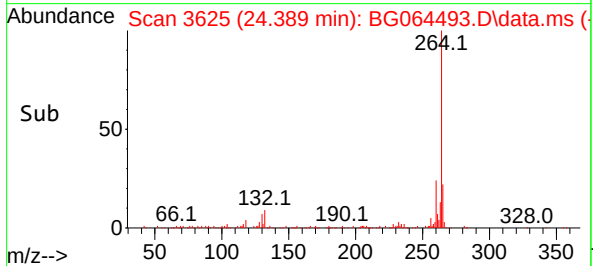
Tgt Ion:264 Resp: 149983

Ion Ratio Lower Upper

264 100

260 24.4 16.2 24.4#

265 22.5 15.0 22.4#



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Integration Parameters: rteint.p

Integrator: RTE

Smoothing : ON

Sampling : 1

Start Thrs: 0.2

Stop Thrs : 0

Filtering: 5

Min Area: 3 % of largest Peak

Max Peaks: 100

Peak Location: TOP

If leading or trailing edge < 100 prefer < Baseline drop else tangent >

Peak separation: 5

Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M

Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

Signal : TIC: BG064493.D\data.ms

| peak<br># | R.T.<br>min | first<br>scan | max<br>scan | last<br>scan | PK<br>TY | peak<br>height | corr.<br>area | corr.<br>% max. | % of<br>total |
|-----------|-------------|---------------|-------------|--------------|----------|----------------|---------------|-----------------|---------------|
| 1         | 4.941       | 310           | 315         | 320          | rBV      | 40001          | 55911         | 2.94%           | 0.289%        |
| 2         | 5.411       | 389           | 395         | 403          | rBV      | 197160         | 310378        | 16.34%          | 1.603%        |
| 3         | 6.997       | 656           | 665         | 673          | rBV      | 132736         | 243525        | 12.82%          | 1.258%        |
| 4         | 7.215       | 697           | 702         | 707          | rBV      | 23646          | 38555         | 2.03%           | 0.199%        |
| 5         | 7.820       | 798           | 805         | 811          | rVB2     | 87062          | 149972        | 7.90%           | 0.774%        |
| 6         | 7.937       | 821           | 825         | 834          | rBV3     | 15596          | 28453         | 1.50%           | 0.147%        |
| 7         | 8.090       | 846           | 851         | 861          | rVB9     | 7598           | 23374         | 1.23%           | 0.121%        |
| 8         | 8.190       | 862           | 868         | 873          | rBV5     | 14847          | 27445         | 1.45%           | 0.142%        |
| 9         | 8.460       | 910           | 914         | 917          | rBV2     | 15811          | 21230         | 1.12%           | 0.110%        |
| 10        | 8.513       | 920           | 923         | 930          | rVB9     | 14241          | 24408         | 1.29%           | 0.126%        |
| 11        | 8.596       | 932           | 937         | 939          | rBV4     | 18375          | 29502         | 1.55%           | 0.152%        |
| 12        | 8.930       | 985           | 994         | 997          | rBV3     | 47247          | 94433         | 4.97%           | 0.488%        |
| 13        | 8.977       | 997           | 1002        | 1008         | rVV      | 260683         | 433060        | 22.80%          | 2.236%        |
| 14        | 9.183       | 1025          | 1037        | 1042         | rBV9     | 48929          | 147964        | 7.79%           | 0.764%        |
| 15        | 9.259       | 1042          | 1050        | 1052         | rBV3     | 34323          | 66740         | 3.51%           | 0.345%        |
| 16        | 9.312       | 1057          | 1059        | 1064         | rVB5     | 19556          | 29449         | 1.55%           | 0.152%        |
| 17        | 9.471       | 1079          | 1086        | 1088         | rBV7     | 17906          | 37995         | 2.00%           | 0.196%        |
| 18        | 9.506       | 1088          | 1092        | 1094         | rVV2     | 37975          | 58066         | 3.06%           | 0.300%        |
| 19        | 9.542       | 1094          | 1098        | 1103         | rVB5     | 59378          | 102568        | 5.40%           | 0.530%        |
| 20        | 9.706       | 1123          | 1126        | 1130         | rVB5     | 16696          | 23023         | 1.21%           | 0.119%        |
| 21        | 9.753       | 1130          | 1134        | 1137         | rBV3     | 27934          | 38358         | 2.02%           | 0.198%        |
| 22        | 9.806       | 1137          | 1143        | 1151         | rVB5     | 68479          | 157941        | 8.32%           | 0.816%        |
| 23        | 9.906       | 1155          | 1160        | 1164         | rVB6     | 42281          | 68126         | 3.59%           | 0.352%        |
| 24        | 9.970       | 1164          | 1171        | 1174         | rBV6     | 37691          | 73029         | 3.85%           | 0.377%        |
| 25        | 10.017      | 1174          | 1179        | 1183         | rVV4     | 48421          | 73879         | 3.89%           | 0.382%        |
| 26        | 10.082      | 1188          | 1190        | 1196         | rVB4     | 31509          | 42583         | 2.24%           | 0.220%        |
| 27        | 10.153      | 1198          | 1202        | 1206         | rBV5     | 24415          | 45870         | 2.42%           | 0.237%        |
| 28        | 10.205      | 1207          | 1211        | 1215         | rVV5     | 23990          | 41352         | 2.18%           | 0.214%        |
| 29        | 10.247      | 1215          | 1218        | 1224         | rVB3     | 61706          | 92561         | 4.87%           | 0.478%        |
| 30        | 10.323      | 1226          | 1231        | 1234         | rBV3     | 41364          | 73354         | 3.86%           | 0.379%        |
| 31        | 10.399      | 1240          | 1244        | 1249         | rVB5     | 42376          | 72150         | 3.80%           | 0.373%        |
| 32        | 10.487      | 1255          | 1259        | 1260         | rBV4     | 31589          | 40173         | 2.12%           | 0.207%        |
| 33        | 10.605      | 1271          | 1279        | 1285         | rBV2     | 132387         | 340070        | 17.91%          | 1.756%        |
| 34        | 10.664      | 1285          | 1289        | 1290         | rBV2     | 24392          | 28925         | 1.52%           | 0.149%        |
| 35        | 10.734      | 1297          | 1301        | 1306         | rVV4     | 46933          | 81078         | 4.27%           | 0.419%        |
| 36        | 10.781      | 1306          | 1309        | 1314         | rVB      | 95062          | 142024        | 7.48%           | 0.733%        |

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

A

B

C

D

E

F

G

H

I

J

K

Integration Parameters: rteint.p

Integrator: RTE

Smoothing : ON

Filtering: 5

Sampling : 1

Min Area: 3 % of largest Peak

Start Thrs: 0.2

Max Peaks: 100

Stop Thrs : 0

Peak Location: TOP

If leading or trailing edge < 100 prefer < Baseline drop else tangent >

Peak separation: 5

Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M

Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

|    |        |      |      |      |      |         |         |         |        |
|----|--------|------|------|------|------|---------|---------|---------|--------|
| 37 | 10.840 | 1314 | 1319 | 1327 | rBV7 | 46359   | 110414  | 5.81%   | 0.570% |
| 38 | 10.910 | 1329 | 1331 | 1335 | rVB4 | 31151   | 35404   | 1.86%   | 0.183% |
| 39 | 11.028 | 1346 | 1351 | 1356 | rBV7 | 59293   | 125142  | 6.59%   | 0.646% |
| 40 | 11.110 | 1362 | 1365 | 1369 | rVB4 | 67533   | 89948   | 4.74%   | 0.464% |
| 41 | 11.193 | 1374 | 1379 | 1383 | rVV4 | 73911   | 122671  | 6.46%   | 0.633% |
| 42 | 11.234 | 1383 | 1386 | 1389 | rVV5 | 26390   | 36194   | 1.91%   | 0.187% |
| 43 | 11.316 | 1390 | 1400 | 1408 | rVV5 | 80482   | 294385  | 15.50%  | 1.520% |
| 44 | 11.386 | 1408 | 1412 | 1417 | rVB4 | 54886   | 98153   | 5.17%   | 0.507% |
| 45 | 11.463 | 1420 | 1425 | 1430 | rVV7 | 67843   | 134139  | 7.06%   | 0.693% |
| 46 | 11.527 | 1430 | 1436 | 1444 | rVB5 | 73278   | 179981  | 9.48%   | 0.929% |
| 47 | 11.651 | 1451 | 1457 | 1465 | rBV2 | 222993  | 520961  | 27.43%  | 2.690% |
| 48 | 11.804 | 1479 | 1483 | 1487 | rBV3 | 81723   | 124171  | 6.54%   | 0.641% |
| 49 | 11.903 | 1497 | 1500 | 1504 | rVB4 | 54243   | 73969   | 3.89%   | 0.382% |
| 50 | 12.409 | 1583 | 1586 | 1590 | rBV4 | 45893   | 59591   | 3.14%   | 0.308% |
| 51 | 12.526 | 1602 | 1606 | 1612 | rVB3 | 168838  | 258535  | 13.61%  | 1.335% |
| 52 | 12.955 | 1674 | 1679 | 1682 | rBV6 | 82116   | 178853  | 9.42%   | 0.924% |
| 53 | 13.002 | 1683 | 1687 | 1692 | rVB  | 464058  | 649169  | 34.18%  | 3.352% |
| 54 | 13.079 | 1694 | 1700 | 1705 | rVB  | 742657  | 1110122 | 58.45%  | 5.733% |
| 55 | 13.243 | 1725 | 1728 | 1730 | rBV3 | 103230  | 132972  | 7.00%   | 0.687% |
| 56 | 13.396 | 1749 | 1754 | 1760 | rVV4 | 193183  | 423809  | 22.32%  | 2.189% |
| 57 | 13.449 | 1761 | 1763 | 1768 | rVB4 | 131725  | 160728  | 8.46%   | 0.830% |
| 58 | 13.731 | 1808 | 1811 | 1813 | rBV4 | 98185   | 134639  | 7.09%   | 0.695% |
| 59 | 13.883 | 1833 | 1837 | 1840 | rVB  | 186984  | 243396  | 12.82%  | 1.257% |
| 60 | 13.954 | 1845 | 1849 | 1853 | rVV2 | 482293  | 641767  | 33.79%  | 3.314% |
| 61 | 14.160 | 1881 | 1884 | 1887 | rBV4 | 109863  | 163549  | 8.61%   | 0.845% |
| 62 | 14.453 | 1928 | 1934 | 1942 | rVB4 | 253459  | 602951  | 31.75%  | 3.114% |
| 63 | 14.988 | 2022 | 2025 | 2034 | rVB6 | 271974  | 474482  | 24.98%  | 2.450% |
| 64 | 15.082 | 2037 | 2041 | 2044 | rBV4 | 187138  | 333489  | 17.56%  | 1.722% |
| 65 | 15.722 | 2146 | 2150 | 2156 | rVB  | 717501  | 1094931 | 57.65%  | 5.654% |
| 66 | 15.946 | 2183 | 2188 | 2195 | rVB3 | 696936  | 1158897 | 61.02%  | 5.985% |
| 67 | 16.204 | 2228 | 2232 | 2243 | rVB  | 1163831 | 1899152 | 100.00% | 9.807% |
| 68 | 17.021 | 2367 | 2371 | 2379 | rVB  | 930979  | 1628576 | 85.75%  | 8.410% |
| 69 | 19.812 | 2842 | 2846 | 2852 | rVB  | 1344246 | 1707441 | 89.91%  | 8.817% |
| 70 | 21.087 | 3061 | 3063 | 3075 | rVB7 | 77829   | 148237  | 7.81%   | 0.766% |
| 71 | 21.416 | 3114 | 3119 | 3127 | rVB  | 241586  | 405107  | 21.33%  | 2.092% |
| 72 | 22.785 | 3350 | 3352 | 3358 | rVB7 | 17330   | 24511   | 1.29%   | 0.127% |
| 73 | 22.985 | 3382 | 3386 | 3389 | rVB5 | 23965   | 33776   | 1.78%   | 0.174% |
| 74 | 24.389 | 3615 | 3625 | 3635 | rVB  | 148355  | 392807  | 20.68%  | 2.028% |

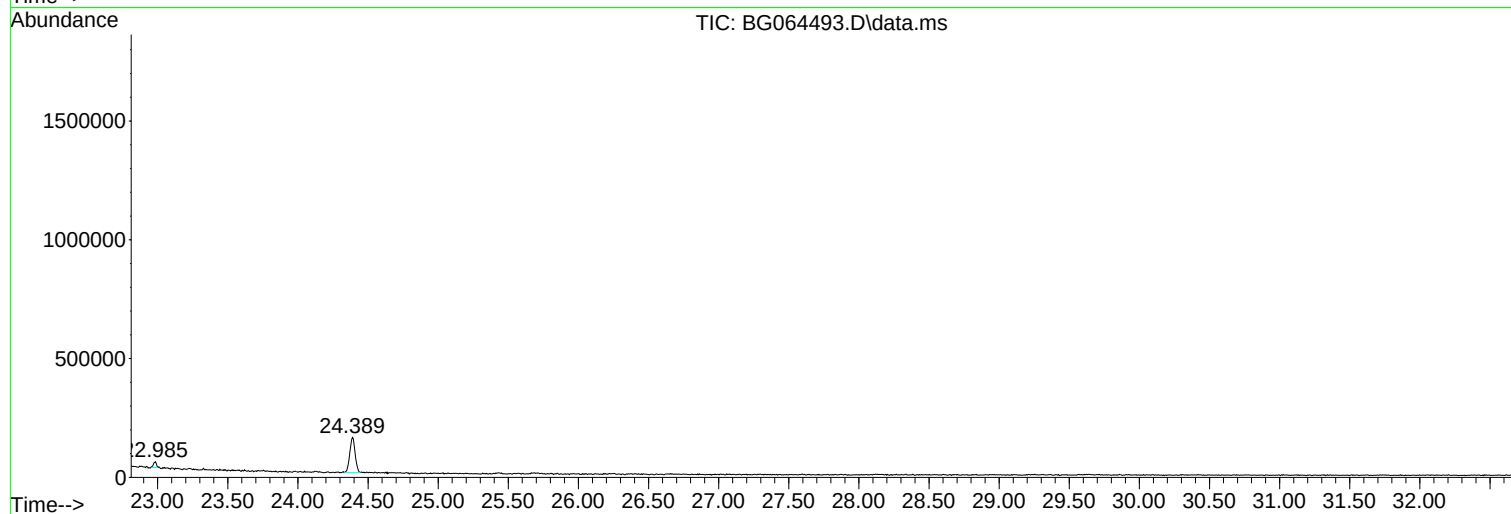
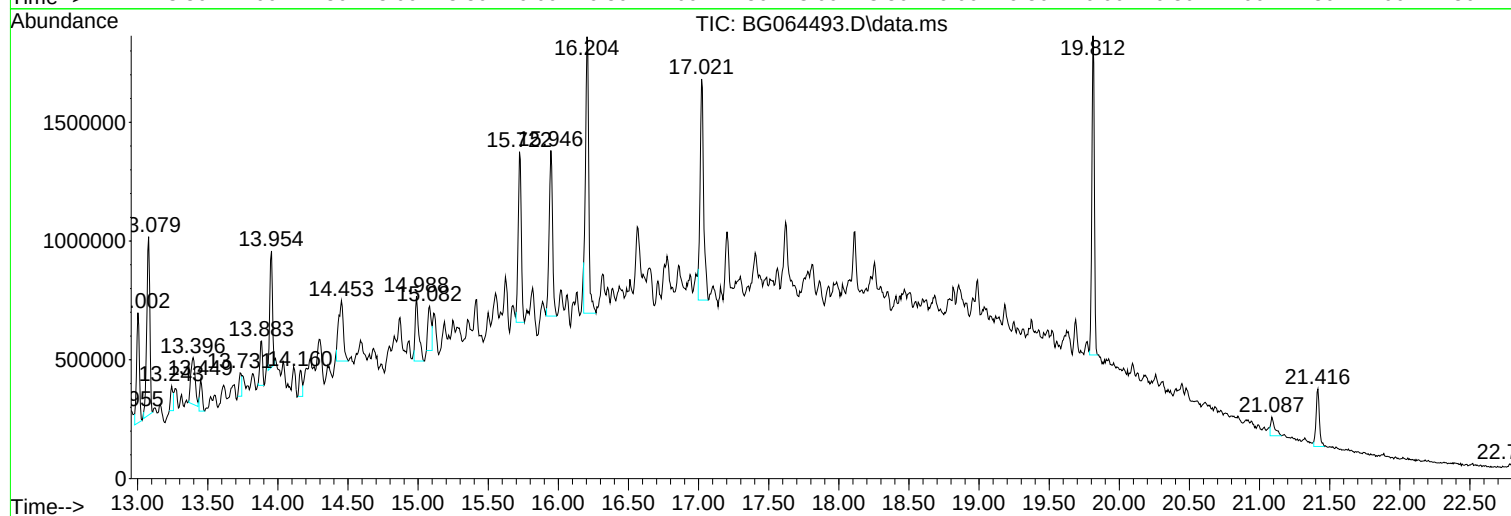
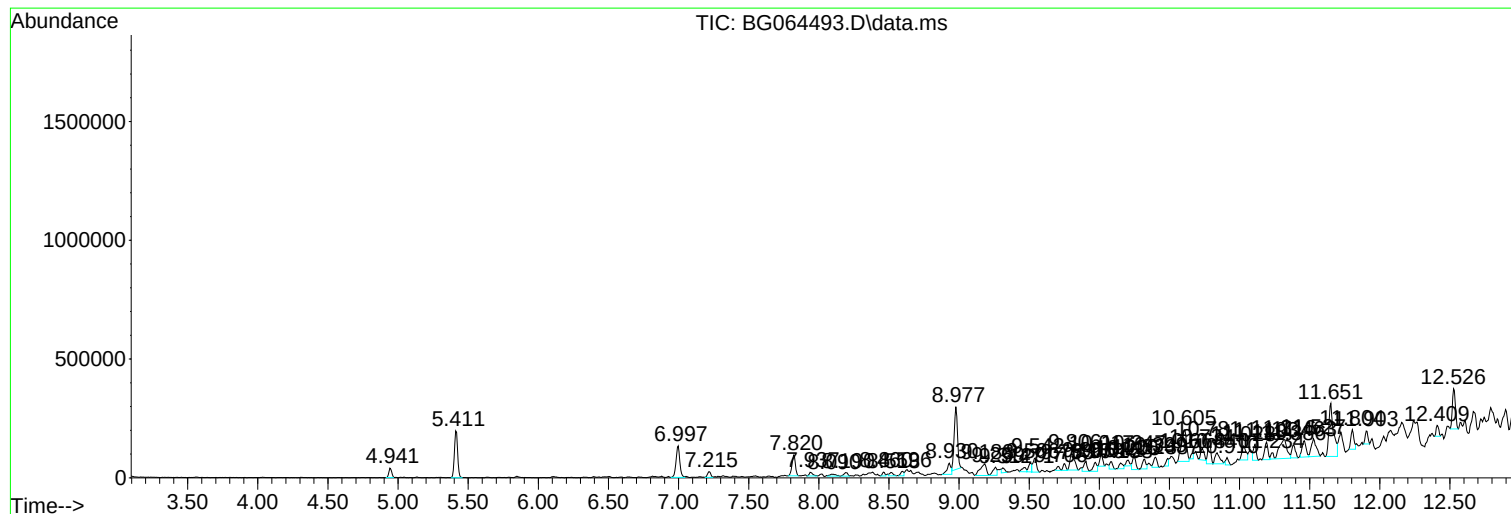
Sum of corrected areas: 19364543

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

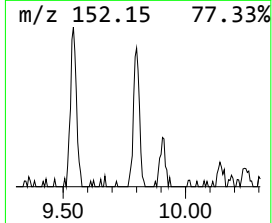
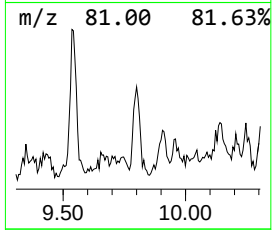
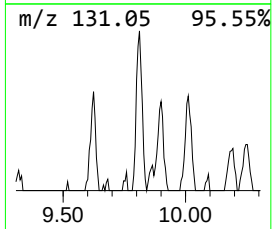
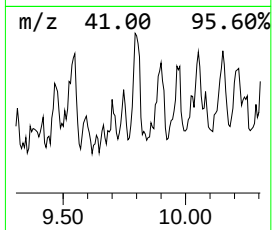
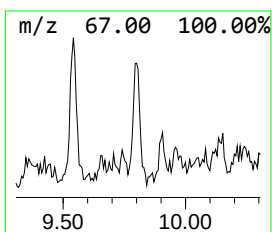
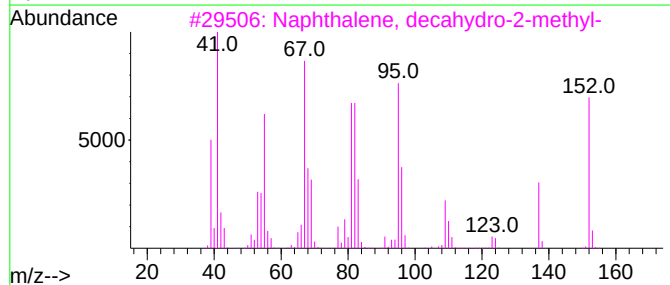
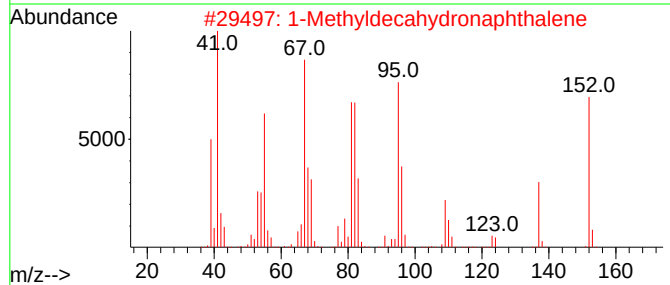
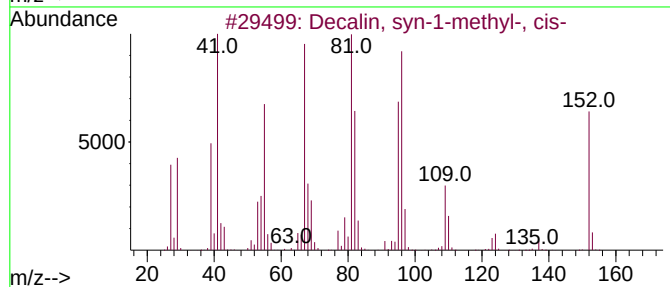
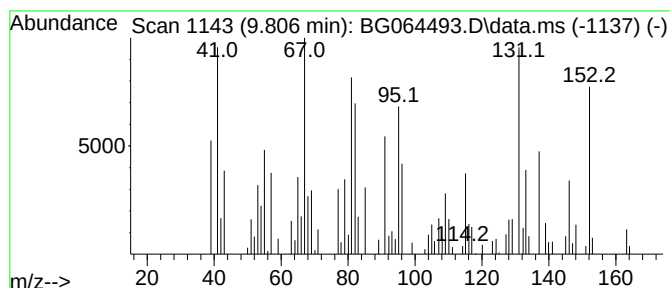
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*

Peak Number 3 Decalin, syn-1-methyl-, cis- Concentration Rank 14

| R.T.  | EstConc | Area                                | Relative to ISTD | R.T.         |      |
|-------|---------|-------------------------------------|------------------|--------------|------|
| 9.806 | 9.29 ng | 157941                              | Naphthalene-d8   | 10.611       |      |
| Hit#  | of 5    | Tentative ID                        | MW MolForm       | CAS#         | Qual |
| 1     |         | Decalin, syn-1-methyl-, cis-        | 152 C11H20       | 1000158-89-1 | 87   |
| 2     |         | 1-Methyldecahydronaphthalene        | 152 C11H20       | 002958-75-0  | 86   |
| 3     |         | Naphthalene, decahydro-2-methyl-    | 152 C11H20       | 002958-76-1  | 86   |
| 4     |         | trans-4a-Methyl-decahydronaphtha... | 152 C11H20       | 002547-27-5  | 55   |
| 5     |         | 1,3,3-Trimethylcyclohex-1-ene-4-... | 152 C10H16O      | 127128-60-3  | 50   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

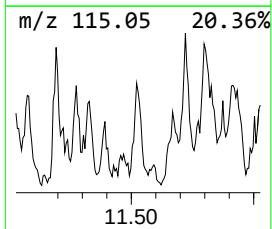
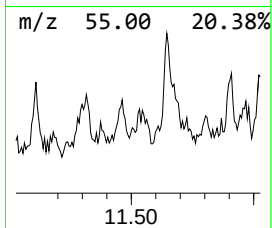
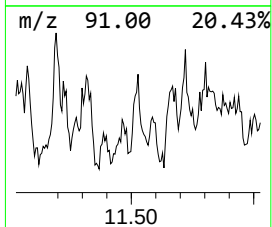
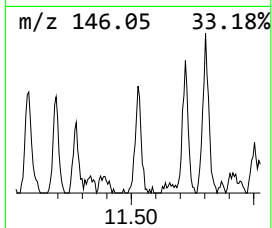
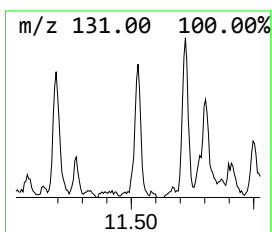
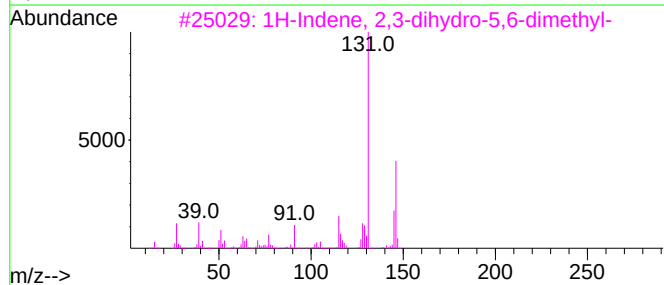
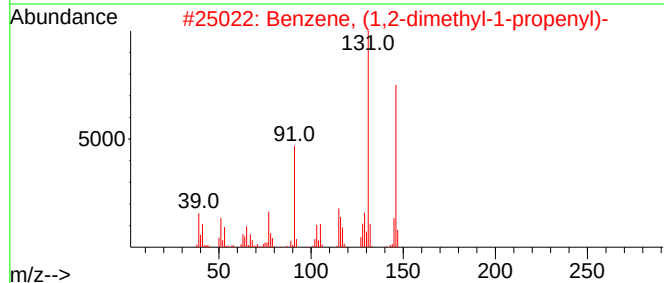
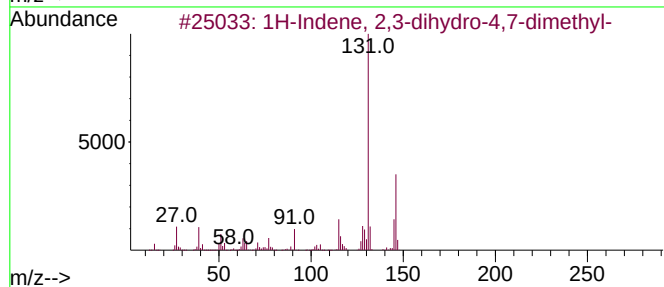
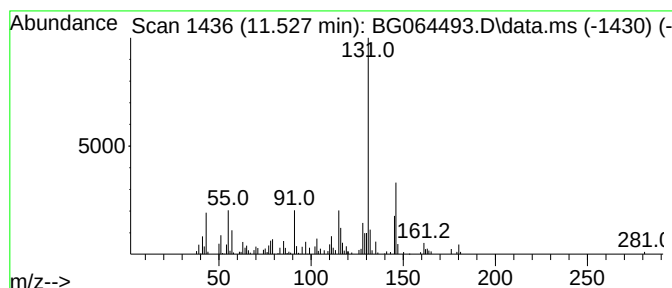
\*\*\*\*\*

Peak Number 8 1H-Indene, 2,3-dihydro-4,7-... Concentration Rank 13

| R.T.   | EstConc  | Area   | Relative to ISTD | R.T.   |
|--------|----------|--------|------------------|--------|
| 11.527 | 10.58 ng | 179981 | Naphthalene-d8   | 10.611 |

| Hit# | of | 5 | Tentative ID                        | MW  | MolForm | CAS#        | Qual |
|------|----|---|-------------------------------------|-----|---------|-------------|------|
| 1    |    |   | 1H-Indene, 2,3-dihydro-4,7-dimet... | 146 | C11H14  | 006682-71-9 | 90   |
| 2    |    |   | Benzene, (1,2-dimethyl-1-propenyl)- | 146 | C11H14  | 000769-57-3 | 90   |
| 3    |    |   | 1H-Indene, 2,3-dihydro-5,6-dimet... | 146 | C11H14  | 001075-22-5 | 90   |
| 4    |    |   | Naphthalene, 1,2,3,4-tetrahydro-... | 146 | C11H14  | 001559-81-5 | 90   |
| 5    |    |   | Benzene, (3-methyl-2-butenyl)-      | 146 | C11H14  | 004489-84-3 | 87   |





Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

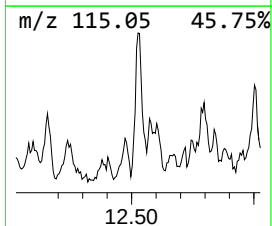
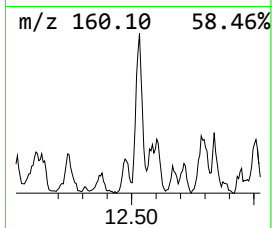
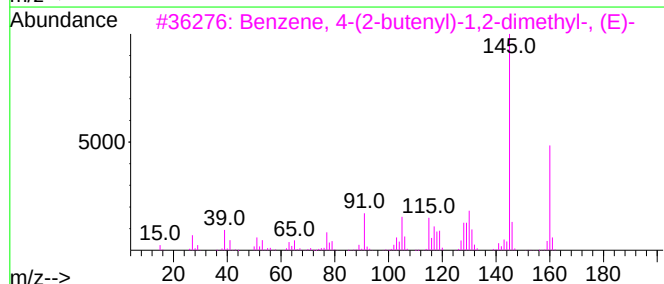
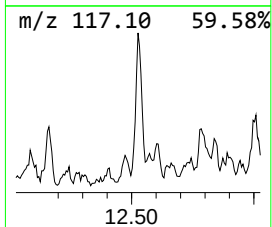
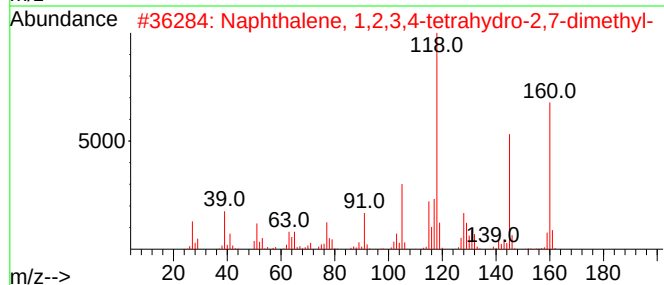
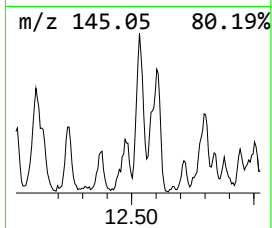
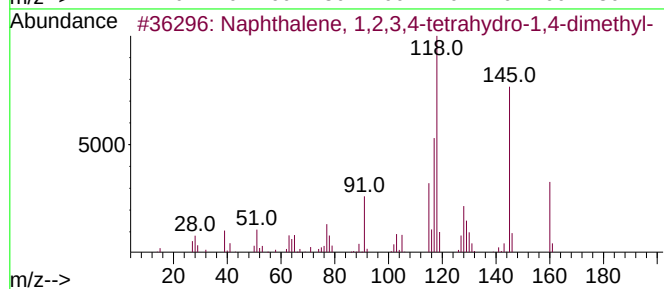
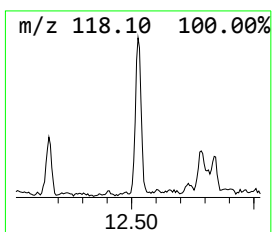
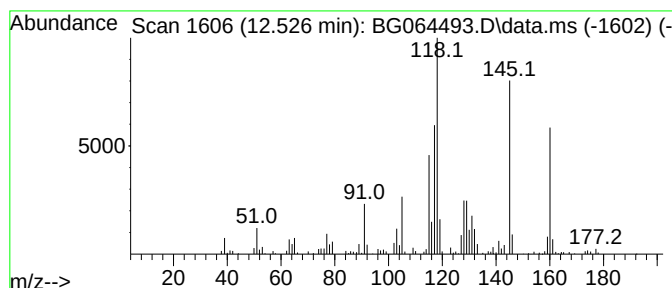
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*

Peak Number 10 Naphthalene, 1,2,3,4-tetrahydro-... Concentration Rank 10

| R.T.   | EstConc  | Area                                | Relative to ISTD | R.T.        |      |
|--------|----------|-------------------------------------|------------------|-------------|------|
| 12.526 | 15.20 ng | 258535                              | Naphthalene-d8   | 10.611      |      |
| Hit#   | of 5     | Tentative ID                        | MW MolForm       | CAS#        | Qual |
| 1      |          | Naphthalene, 1,2,3,4-tetrahydro-... | 160 C12H16       | 004175-54-6 | 95   |
| 2      |          | Naphthalene, 1,2,3,4-tetrahydro-... | 160 C12H16       | 013065-07-1 | 81   |
| 3      |          | Benzene, 4-(2-butenyl)-1,2-dimet... | 160 C12H16       | 054340-86-2 | 78   |
| 4      |          | Naphthalene, 1,2,3,4-tetrahydro-... | 160 C12H16       | 007524-63-2 | 76   |
| 5      |          | Benzene, 1-(2-butenyl)-2,3-dimet... | 160 C12H16       | 054340-85-1 | 55   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

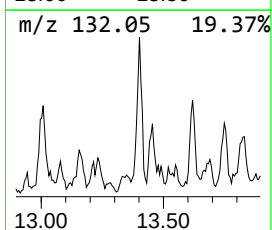
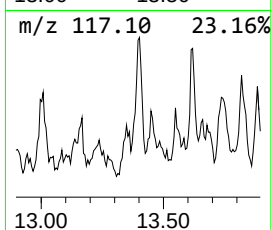
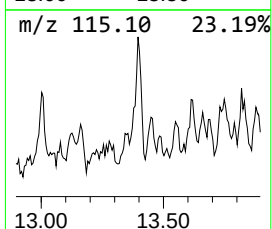
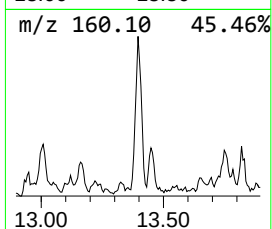
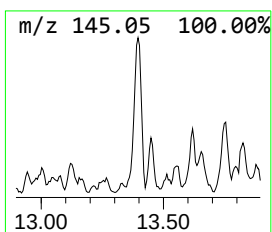
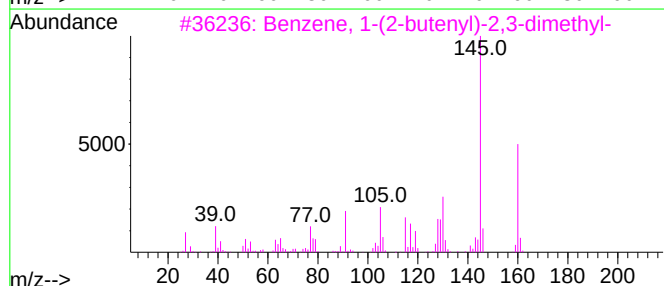
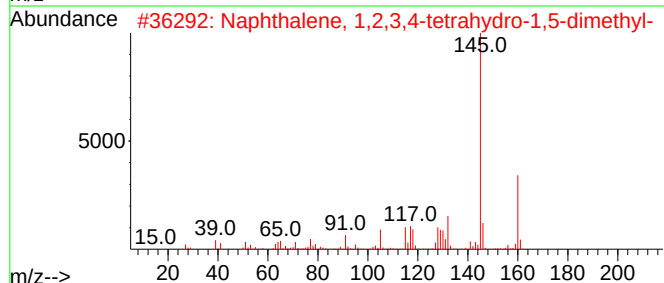
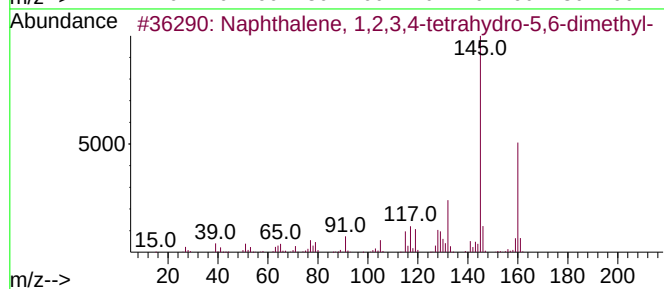
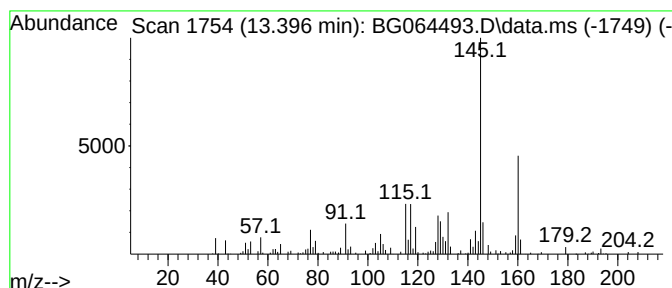
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*

Peak Number 12 Naphthalene, 1,2,3,4-tetrahydro-... Concentration Rank 11

| R.T.      | EstConc                             | Area   | Relative to ISTD | R.T.        |      |
|-----------|-------------------------------------|--------|------------------|-------------|------|
| 13.396    | 14.06 ng                            | 423809 | Acenaphthene-d10 | 14.459      |      |
| Hit# of 5 | Tentative ID                        | MW     | MolForm          | CAS#        | Qual |
| 1         | Naphthalene, 1,2,3,4-tetrahydro-... | 160    | C12H16           | 020027-77-4 | 93   |
| 2         | Naphthalene, 1,2,3,4-tetrahydro-... | 160    | C12H16           | 021564-91-0 | 93   |
| 3         | Benzene, 1-(2-butenyl)-2,3-dimet... | 160    | C12H16           | 054340-85-1 | 89   |
| 4         | Benzene, 1,3,5-trimethyl-2-(1-me... | 160    | C12H16           | 014679-13-1 | 86   |
| 5         | 1H-Indene, 2,3-dihydro-1,1,3-tri... | 160    | C12H16           | 002613-76-5 | 81   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

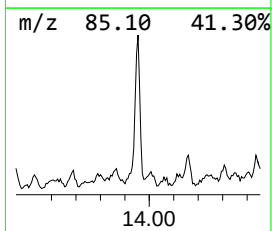
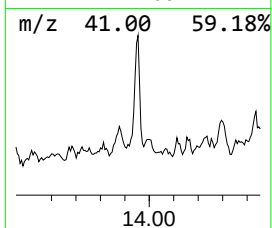
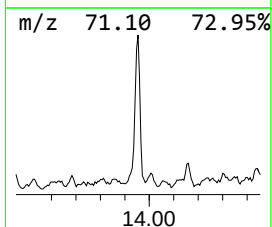
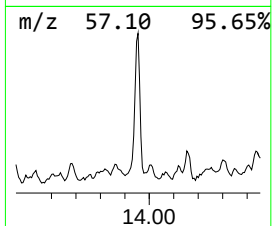
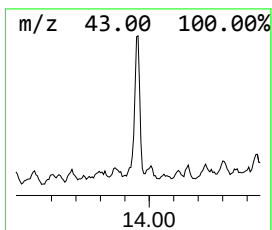
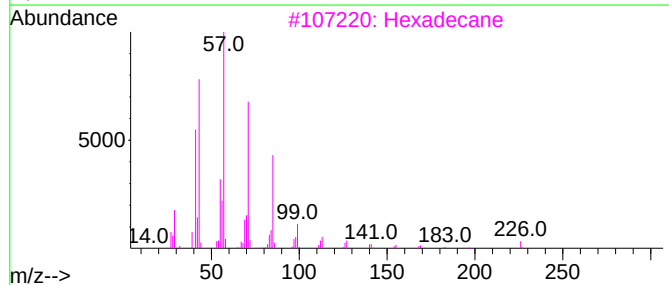
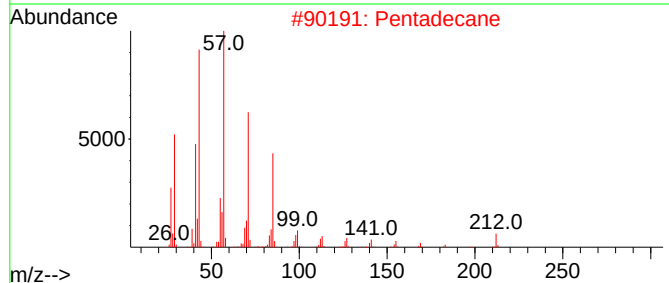
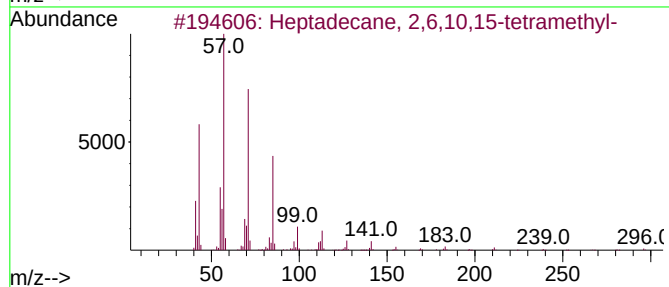
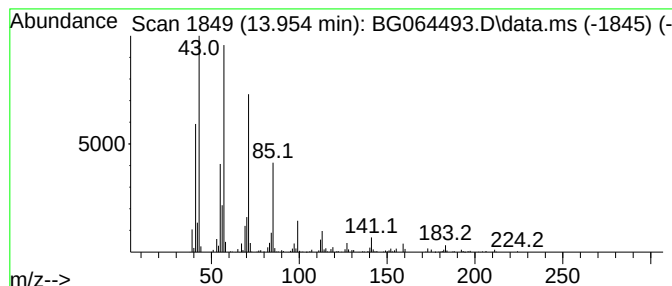
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*

Peak Number 14 Heptadecane, 2,6,10,15-tetr... Concentration Rank 5

| R.T.      | EstConc                             | Area   | Relative to ISTD | R.T.        |      |
|-----------|-------------------------------------|--------|------------------|-------------|------|
| 13.954    | 21.29 ng                            | 641767 | Acenaphthene-d10 | 14.459      |      |
| Hit# of 5 | Tentative ID                        | MW     | MolForm          | CAS#        | Qual |
| 1         | Heptadecane, 2,6,10,15-tetramethyl- | 296    | C21H44           | 054833-48-6 | 91   |
| 2         | Pentadecane                         | 212    | C15H32           | 000629-62-9 | 90   |
| 3         | Hexadecane                          | 226    | C16H34           | 000544-76-3 | 87   |
| 4         | Dodecane, 1-iodo-                   | 296    | C12H25I          | 004292-19-7 | 86   |
| 5         | Dodecane, 2-methyl-6-propyl-        | 226    | C16H34           | 055045-08-4 | 86   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

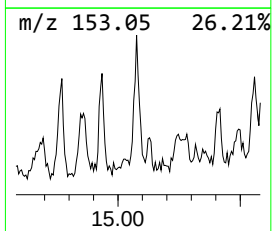
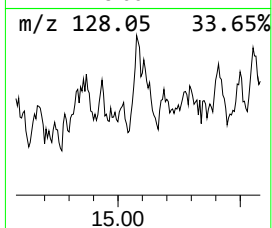
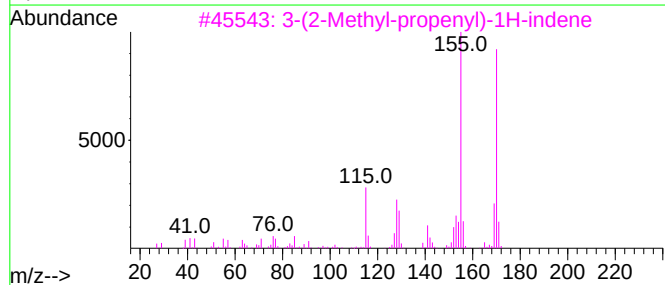
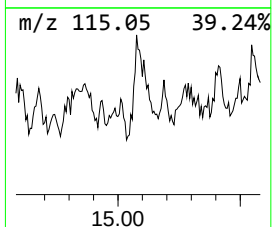
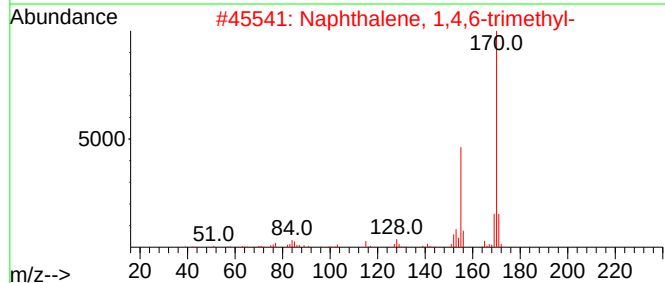
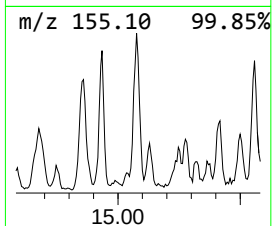
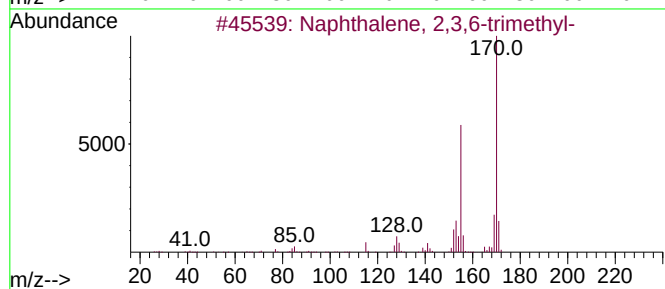
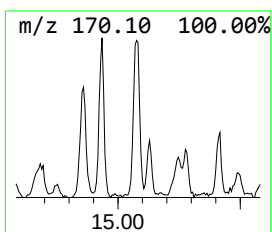
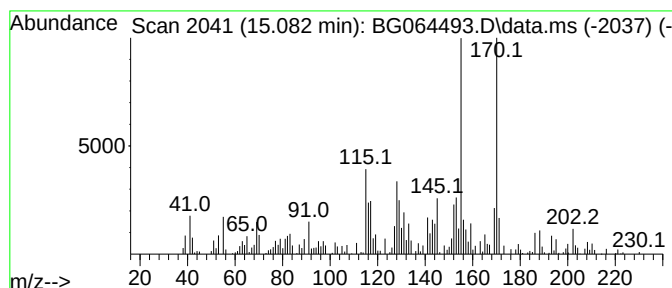
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*

Peak Number 16 Naphthalene, 2,3,6-trimethyl- Concentration Rank 12

| R.T.    | EstConc  | Area                            | Relative to ISTD |         | R.T.         |      |
|---------|----------|---------------------------------|------------------|---------|--------------|------|
| 15.082  | 11.06 ng | 333489                          | Acenaphthene-d10 |         | 14.459       |      |
| Hit# of | 5        | Tentative ID                    | MW               | MolForm | CAS#         | Qual |
| 1       |          | Naphthalene, 2,3,6-trimethyl-   | 170              | C13H14  | 000829-26-5  | 95   |
| 2       |          | Naphthalene, 1,4,6-trimethyl-   | 170              | C13H14  | 002131-42-2  | 93   |
| 3       |          | 3-(2-Methyl-propenyl)-1H-indene | 170              | C13H14  | 1000187-78-5 | 93   |
| 4       |          | Naphthalene, 1,6,7-trimethyl-   | 170              | C13H14  | 002245-38-7  | 93   |
| 5       |          | 4,6,8-Trimethylazulene          | 170              | C13H14  | 000941-81-1  | 86   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

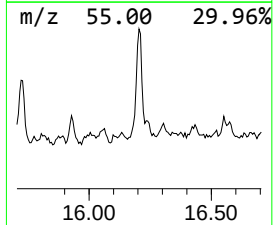
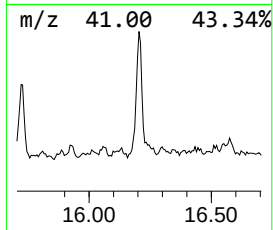
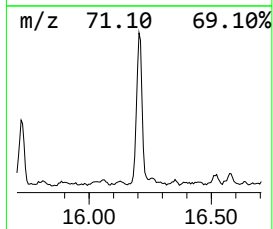
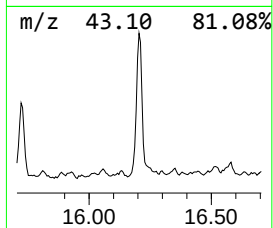
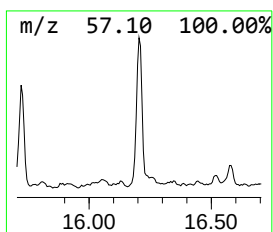
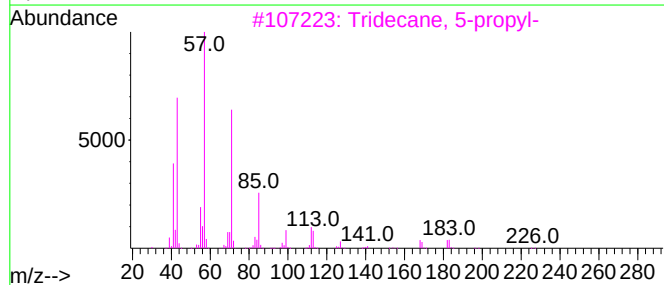
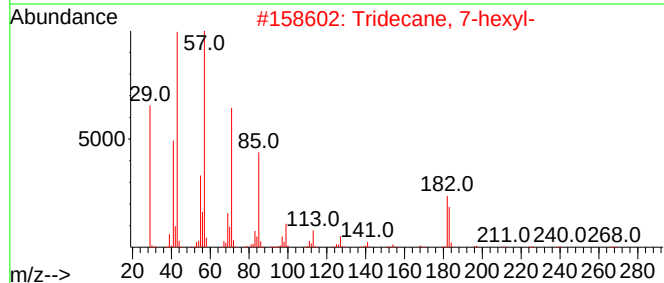
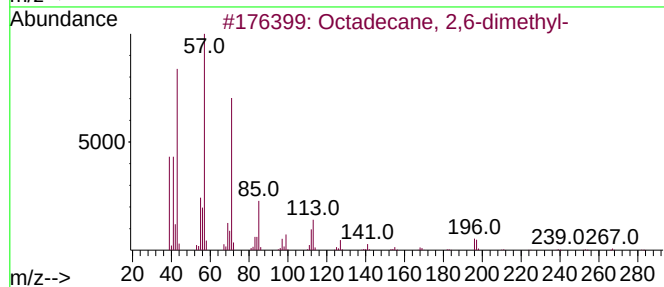
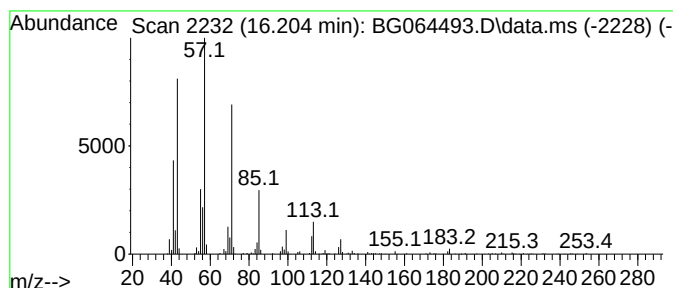
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*

Peak Number 18 Octadecane, 2,6-dimethyl- Concentration Rank 3

| R.T.   | EstConc  | Area                                | Relative to ISTD | R.T.    |             |      |
|--------|----------|-------------------------------------|------------------|---------|-------------|------|
| 16.204 | 23.32 ng | 1899150                             | Phenanthrene-d10 | 17.197  |             |      |
| Hit#   | of 5     | Tentative ID                        | MW               | MolForm | CAS#        | Qual |
| 1      |          | Octadecane, 2,6-dimethyl-           | 282              | C20H42  | 075163-97-2 | 90   |
| 2      |          | Tridecane, 7-hexyl-                 | 268              | C19H40  | 007225-66-3 | 81   |
| 3      |          | Tridecane, 5-propyl-                | 226              | C16H34  | 055045-11-9 | 80   |
| 4      |          | Hexadecane, 2,6,11,15-tetramethyl-  | 282              | C20H42  | 000504-44-9 | 80   |
| 5      |          | Pentadecane, 2,6,10,14-tetramethyl- | 268              | C19H40  | 001921-70-6 | 80   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

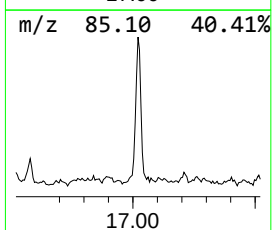
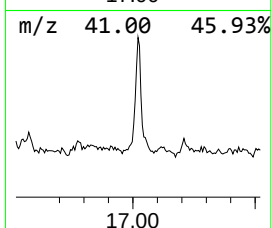
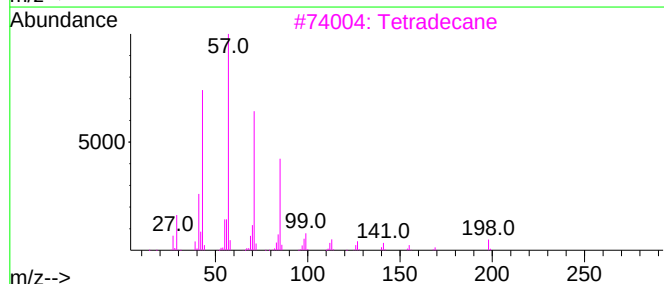
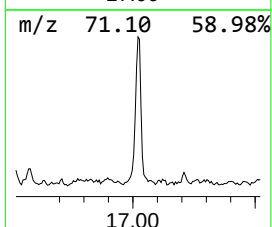
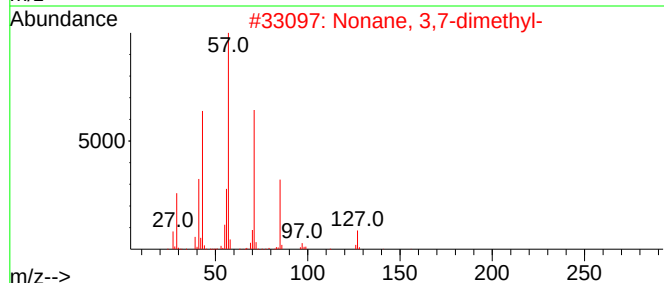
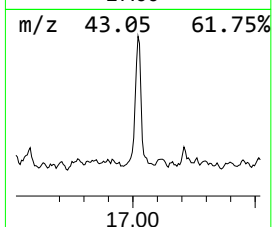
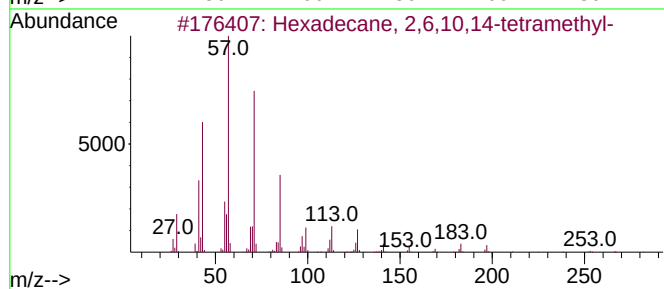
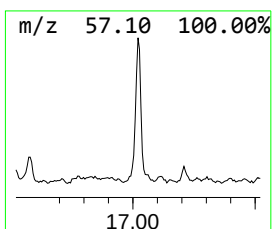
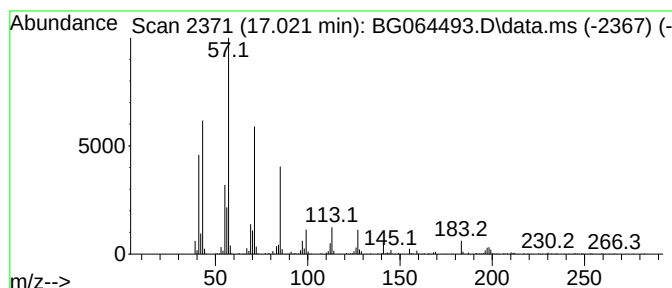
\*\*\*\*\*

Peak Number 19 Hexadecane, 2,6,10,14-tetra... Concentration Rank 6

| R.T.   | EstConc  | Area    | Relative to ISTD | R.T.   |
|--------|----------|---------|------------------|--------|
| 17.021 | 20.00 ng | 1628580 | Phenanthrene-d10 | 17.197 |

| Hit# | of 5 | Tentative ID                        | MW  | MolForm | CAS#        | Qual |
|------|------|-------------------------------------|-----|---------|-------------|------|
| 1    |      | Hexadecane, 2,6,10,14-tetramethyl-  | 282 | C20H42  | 000638-36-8 | 96   |
| 2    |      | Nonane, 3,7-dimethyl-               | 156 | C11H24  | 017302-32-8 | 87   |
| 3    |      | Tetradecane                         | 198 | C14H30  | 000629-59-4 | 87   |
| 4    |      | Heptadecane, 2,6,10,15-tetramethyl- | 296 | C21H44  | 054833-48-6 | 86   |
| 5    |      | Pentadecane, 2,6,10-trimethyl-      | 254 | C18H38  | 003892-00-0 | 81   |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

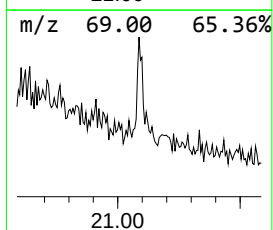
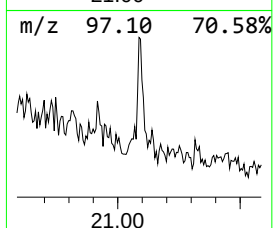
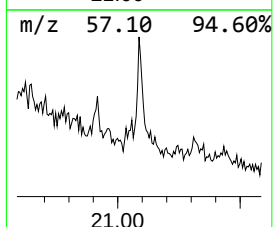
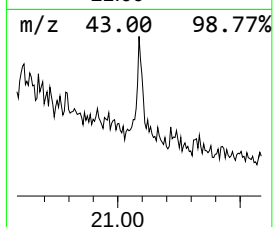
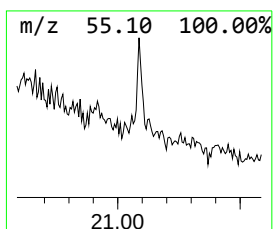
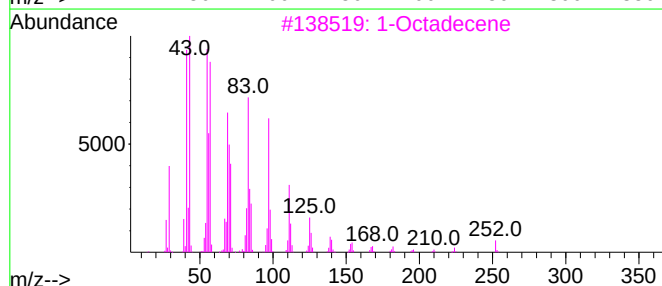
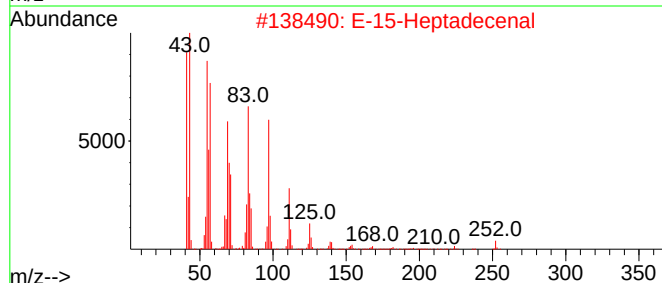
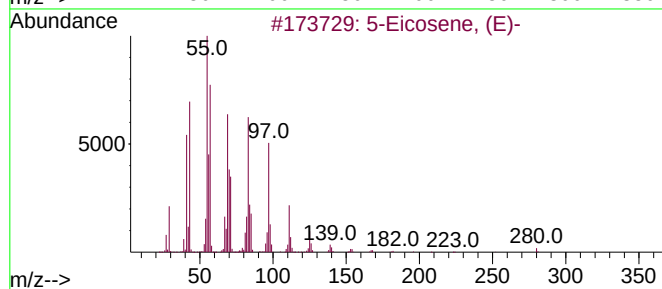
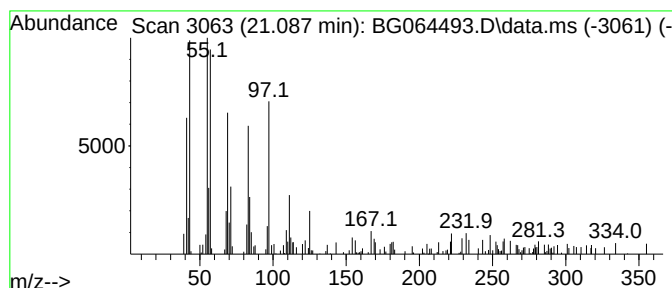
Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

\*\*\*\*\*  
Peak Number 20 5-Eicosene, (E)- Concentration Rank 20

| R.T.      | EstConc                        | Area   | Relative to ISTD |            |              | R.T.   |
|-----------|--------------------------------|--------|------------------|------------|--------------|--------|
| 21.087    | 7.32 ng                        | 148237 | Chrysene-d12     |            |              | 21.416 |
| Hit# of 5 | Tentative ID                   |        | MW               | MolForm    | CAS#         | Qual   |
| 1         | 5-Eicosene, (E)-               |        | 280              | C20H40     | 074685-30-6  | 86     |
| 2         | E-15-Heptadecenal              |        | 252              | C17H32O    | 1000130-97-9 | 76     |
| 3         | 1-Octadecene                   |        | 252              | C18H36     | 000112-88-9  | 76     |
| 4         | Heptadecyl heptafluorobutyrate |        | 452              | C21H35F7O2 | 959085-66-6  | 74     |
| 5         | 1-Nonadecene                   |        | 266              | C19H38     | 018435-45-5  | 72     |



5

A

B

C

D

E

F

G

H

I

J

K

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064493.D  
Acq On : 3 Oct 2025 18:59  
Operator : RC/JU  
Sample : Q3273-01  
Misc :  
ALS Vial : 10 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
MW1

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

| TIC Top Hit name   | RT     | EstConc | Units | Response | --Internal Standard-- |        |         |      |
|--------------------|--------|---------|-------|----------|-----------------------|--------|---------|------|
|                    |        |         |       |          | #                     | RT     | Resp    | Conc |
| Decalin, syn-1-... | 9.806  | 9.3     | ng    | 157941   | 2                     | 10.611 | 340070  | 20.0 |
| 1H-Indene, 2,3-... | 11.527 | 10.6    | ng    | 179981   | 2                     | 10.611 | 340070  | 20.0 |
| Naphthalene, 1,... | 12.526 | 15.2    | ng    | 258535   | 2                     | 10.611 | 340070  | 20.0 |
| Naphthalene, 1,... | 13.396 | 14.1    | ng    | 423809   | 3                     | 14.459 | 602951  | 20.0 |
| Heptadecane, 2,... | 13.954 | 21.3    | ng    | 641767   | 3                     | 14.459 | 602951  | 20.0 |
| Naphthalene, 2,... | 15.082 | 11.1    | ng    | 333489   | 3                     | 14.459 | 602951  | 20.0 |
| Octadecane, 2,6... | 16.204 | 23.3    | ng    | 1899150  | 4                     | 17.197 | 1628580 | 20.0 |
| Hexadecane, 2,6... | 17.021 | 20.0    | ng    | 1628580  | 4                     | 17.197 | 1628580 | 20.0 |
| 5-Eicosene, (E)-   | 21.087 | 7.3     | ng    | 148237   | 5                     | 21.416 | 405107  | 20.0 |



5

A

B

C

D

E

F

G

H

I

J

K

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064488.D  
Acq On : 3 Oct 2025 15:35  
Operator : RC/JU  
Sample : PB169973BL  
Misc :  
ALS Vial : 5 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

Quant Time: Oct 03 16:20:38 2025  
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
QLast Update : Thu Oct 02 16:40:42 2025  
Response via : Initial Calibration

| Compound                    | R.T.   | QIon | Response | Conc    | Units | Dev(Min) |
|-----------------------------|--------|------|----------|---------|-------|----------|
| -----                       |        |      |          |         |       |          |
| Internal Standards          |        |      |          |         |       |          |
| 1) 1,4-Dichlorobenzene-d4   | 7.824  | 152  | 16965    | 20.000  | ng    | 0.00     |
| 21) Naphthalene-d8          | 10.609 | 136  | 71465    | 20.000  | ng    | # 0.00   |
| 39) Acenaphthene-d10        | 14.446 | 164  | 57081    | 20.000  | ng    | 0.00     |
| 64) Phenanthrene-d10        | 17.189 | 188  | 145353   | 20.000  | ng    | # 0.00   |
| 76) Chrysene-d12            | 21.408 | 240  | 149348   | 20.000  | ng    | 0.00     |
| 86) Perylene-d12            | 24.387 | 264  | 157878   | 20.000  | ng    | # 0.00   |
| System Monitoring Compounds |        |      |          |         |       |          |
| 5) 2-Fluorophenol           | 5.415  | 112  | 148699   | 162.213 | ng    | 0.00     |
| 7) Phenol-d6                | 7.001  | 99   | 185155   | 150.303 | ng    | 0.02     |
| 23) Nitrobenzene-d5         | 8.976  | 82   | 124977   | 94.032  | ng    | 0.00     |
| 42) 2,4,6-Tribromophenol    | 15.938 | 330  | 142586   | 137.808 | ng    | 0.00     |
| 45) 2-Fluorobiphenyl        | 13.071 | 172  | 351343   | 90.727  | ng    | 0.00     |
| 79) Terphenyl-d14           | 19.810 | 244  | 779476   | 98.773  | ng    | 0.00     |

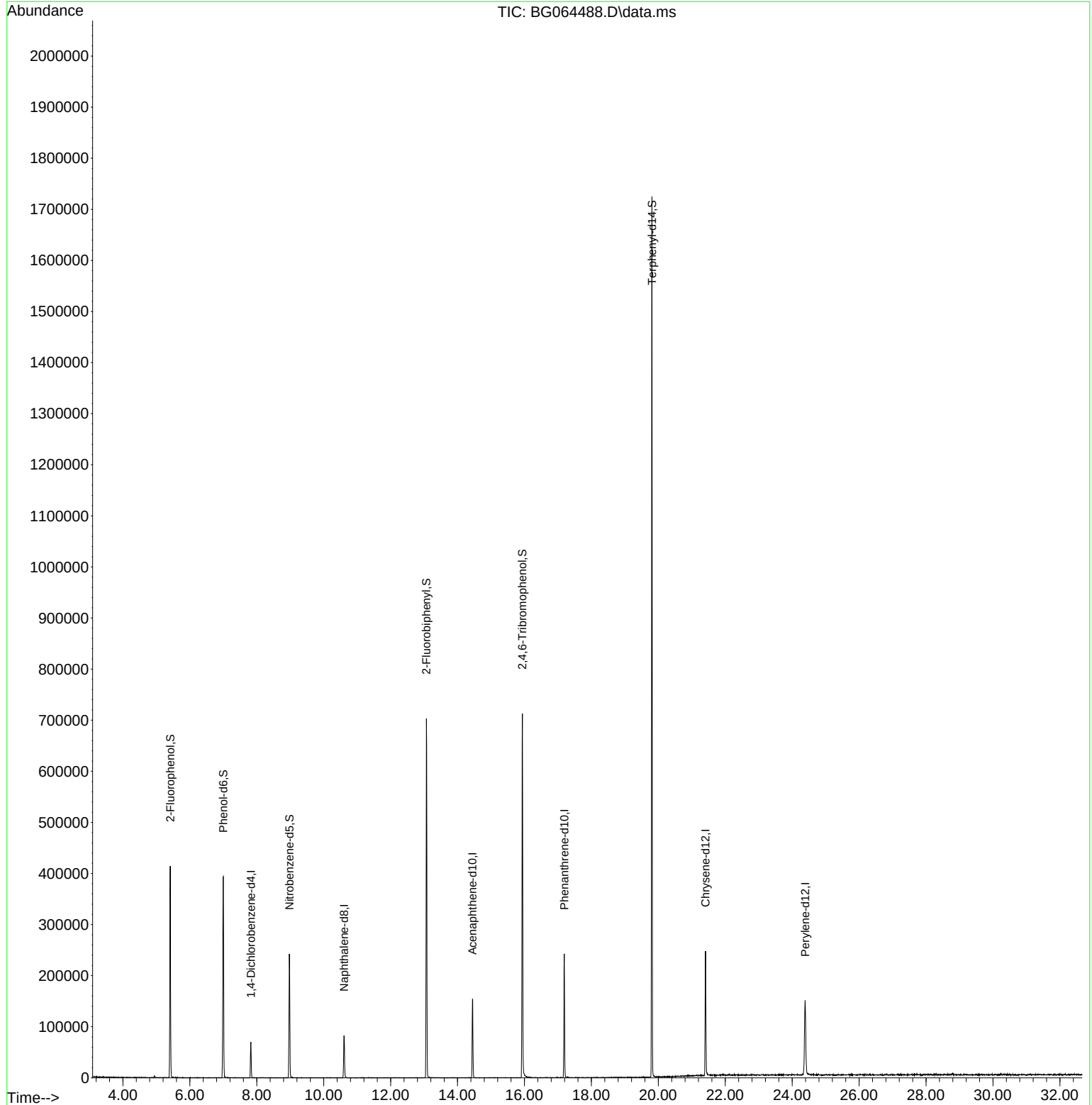
| Target Compounds | Qvalue |
|------------------|--------|
| -----            |        |

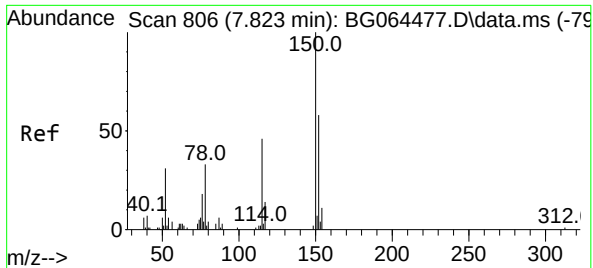
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064488.D  
Acq On : 3 Oct 2025 15:35  
Operator : RC/JU  
Sample : PB169973BL  
Misc :  
ALS Vial : 5 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

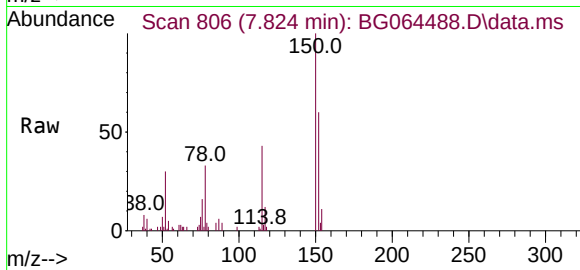
Quant Time: Oct 03 16:20:38 2025  
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
QLast Update : Thu Oct 02 16:40:42 2025  
Response via : Initial Calibration



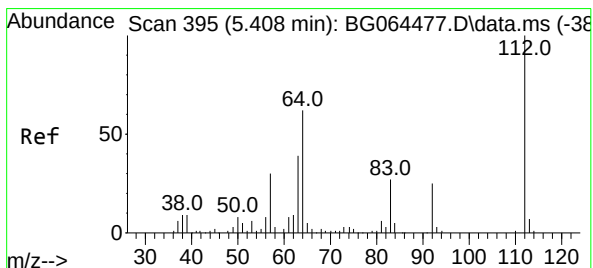
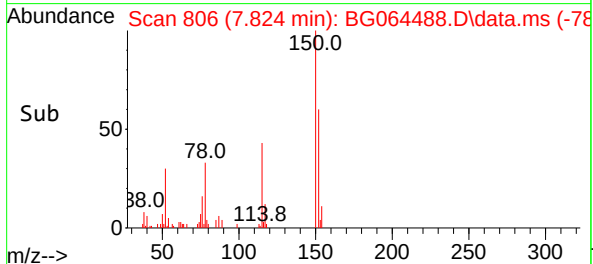
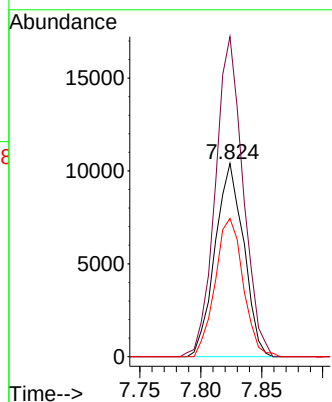


#1  
1,4-Dichlorobenzene-d4  
Concen: 20.000 ng  
RT: 7.824 min Scan# 806  
Delta R.T. 0.004 min  
Lab File: BG064488.D  
Acq: 3 Oct 2025 15:35

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

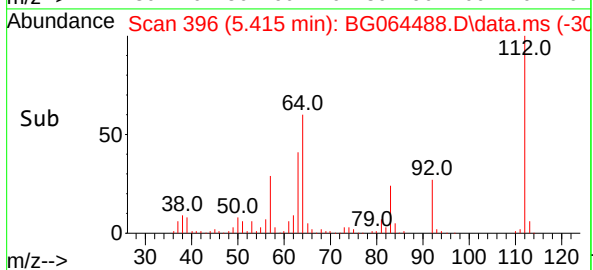
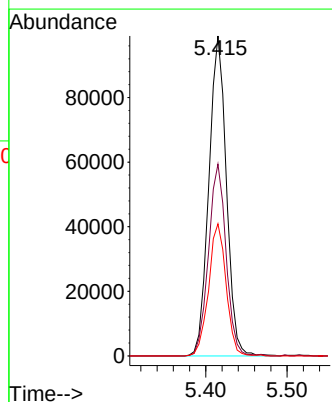
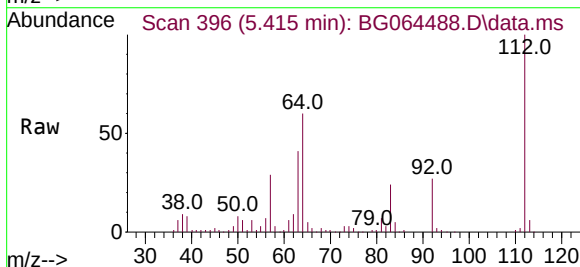


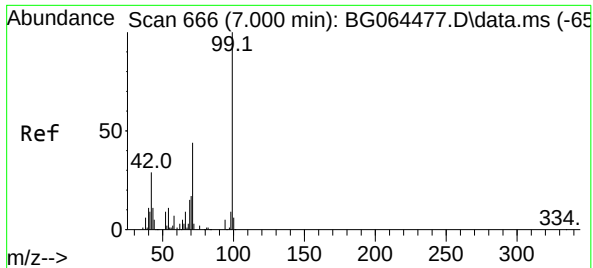
Tgt Ion:152 Resp: 16965  
Ion Ratio Lower Upper  
152 100  
150 165.7 138.6 207.8  
115 71.5 63.4 95.2



#5  
2-Fluorophenol  
Concen: 162.213 ng  
RT: 5.415 min Scan# 396  
Delta R.T. 0.009 min  
Lab File: BG064488.D  
Acq: 3 Oct 2025 15:35

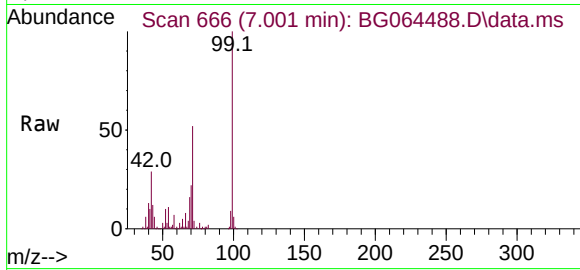
Tgt Ion:112 Resp: 148699  
Ion Ratio Lower Upper  
112 100  
64 59.9 49.8 74.8  
63 41.3 31.3 46.9



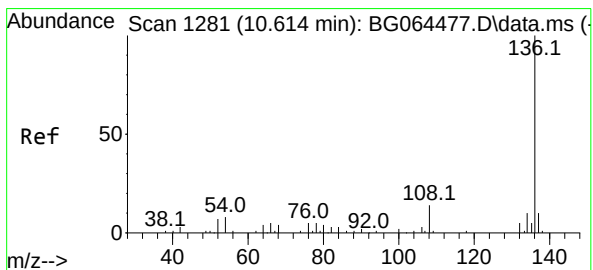
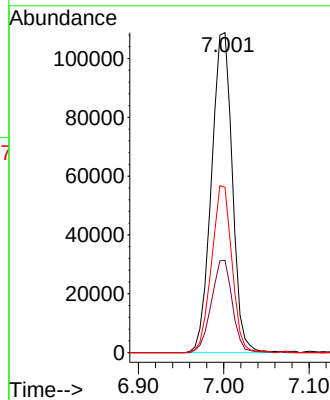
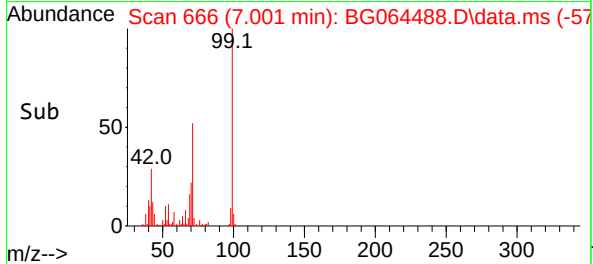


#7  
Phenol-d6  
Concen: 150.303 ng  
RT: 7.001 min Scan# 60  
Delta R.T. 0.015 min  
Lab File: BG064488.D  
Acq: 3 Oct 2025 15:35

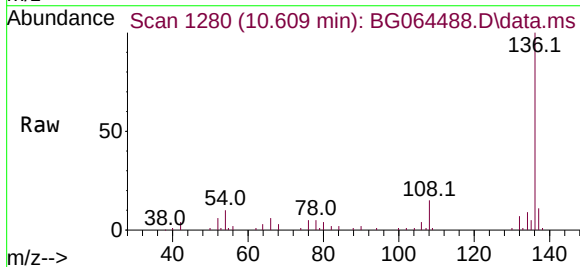
Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL



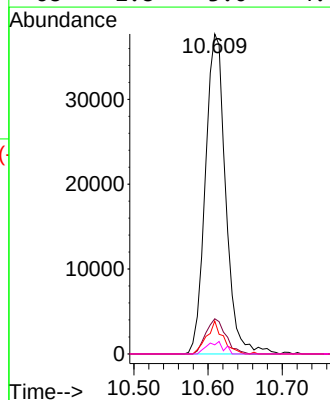
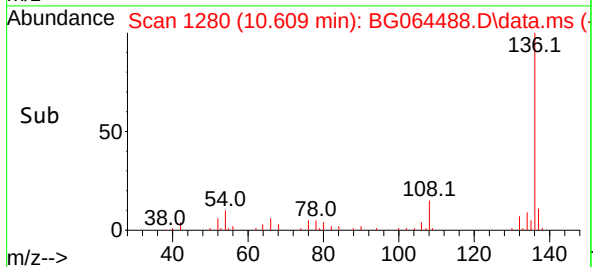
Tgt Ion: 99 Resp: 185155  
Ion Ratio Lower Upper  
99 100  
42 28.8 22.9 34.3  
71 51.7 35.4 53.2

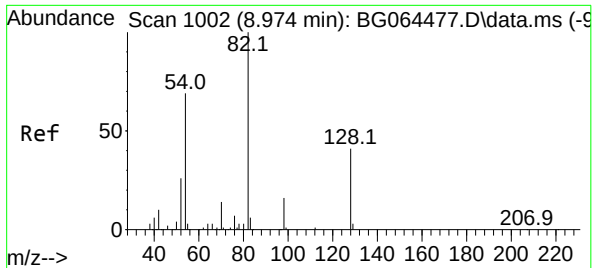


#21  
Naphthalene-d8  
Concen: 20.000 ng  
RT: 10.609 min Scan# 1280  
Delta R.T. 0.003 min  
Lab File: BG064488.D  
Acq: 3 Oct 2025 15:35



Tgt Ion: 136 Resp: 71465  
Ion Ratio Lower Upper  
136 100  
137 11.0 8.3 12.5  
54 10.3 6.2 9.4#  
68 2.8 3.0 4.4#





#23

Nitrobenzene-d5

Concen: 94.032 ng

RT: 8.976 min Scan# 10

Delta R.T. 0.009 min

Lab File: BG064488.D

Acq: 3 Oct 2025 15:35

Instrument :

BNA\_G

ClientSampleId :

PB169973BL

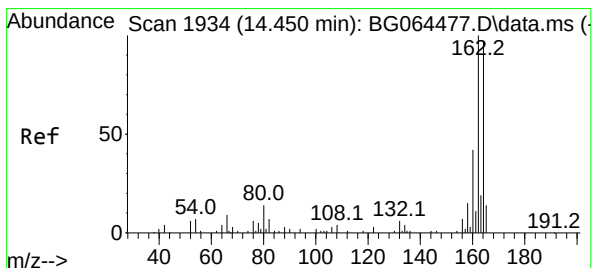
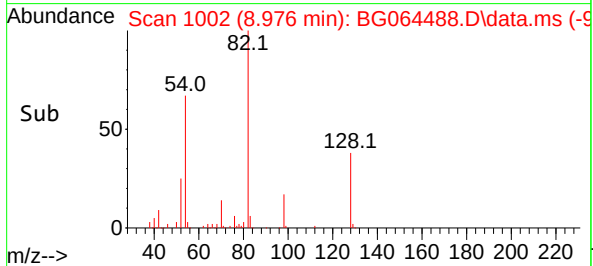
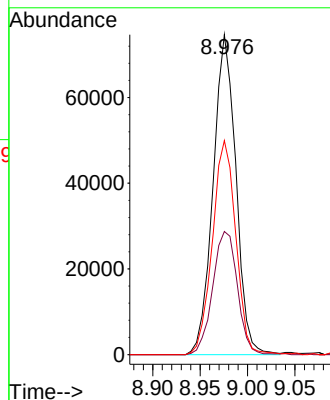
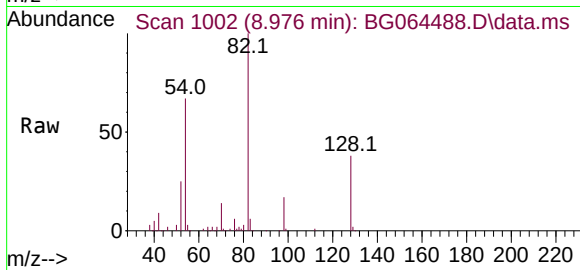
Tgt Ion: 82 Resp: 124977

Ion Ratio Lower Upper

82 100

128 38.5 33.1 49.7

54 66.9 55.4 83.2



#39

Acenaphthene-d10

Concen: 20.000 ng

RT: 14.446 min Scan# 1933

Delta R.T. -0.002 min

Lab File: BG064488.D

Acq: 3 Oct 2025 15:35

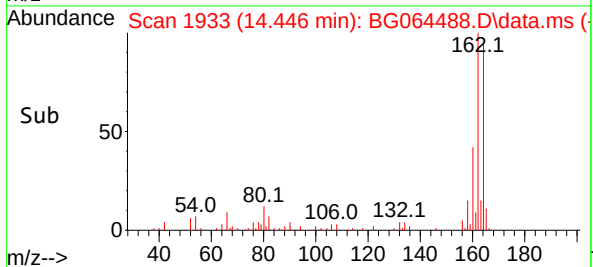
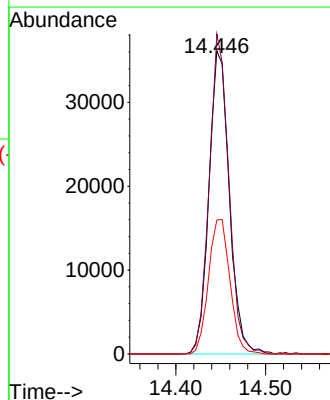
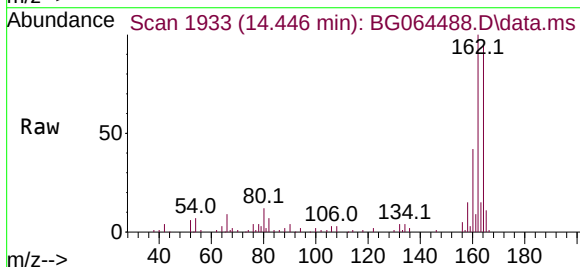
Tgt Ion:164 Resp: 57081

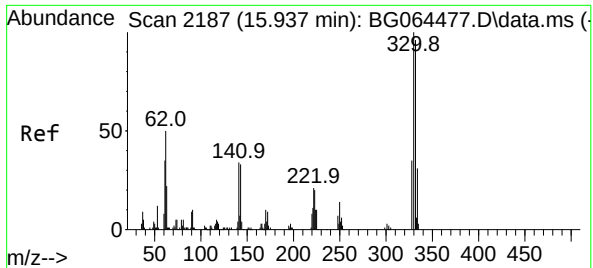
Ion Ratio Lower Upper

164 100

162 105.6 82.2 123.2

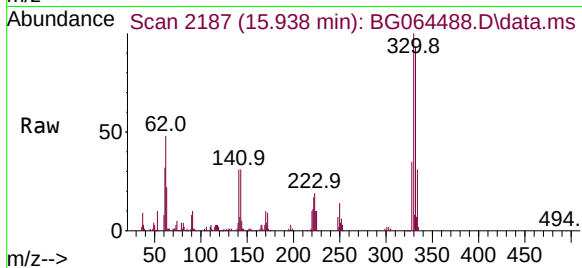
160 44.2 34.3 51.5



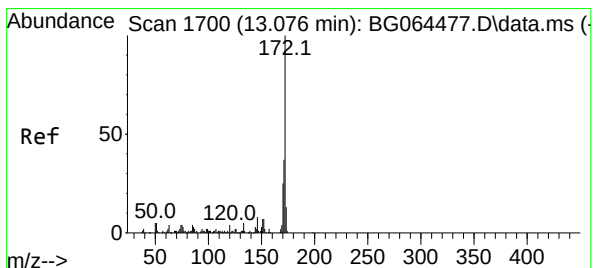
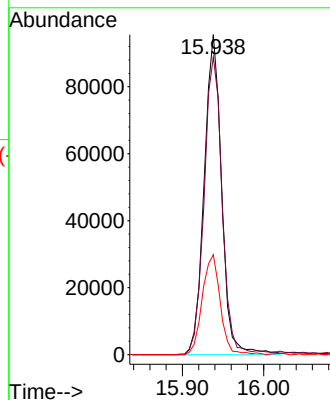
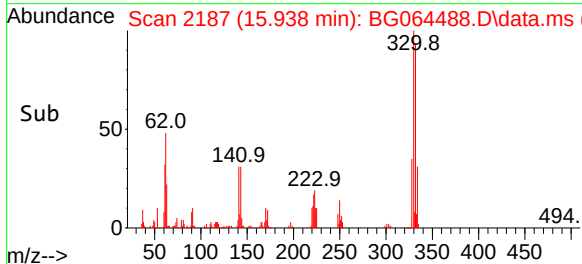


#42  
2,4,6-Tribromophenol  
Concen: 137.808 ng  
RT: 15.938 min Scan# 2187  
Delta R.T. 0.003 min  
Lab File: BG064488.D  
Acq: 3 Oct 2025 15:35

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

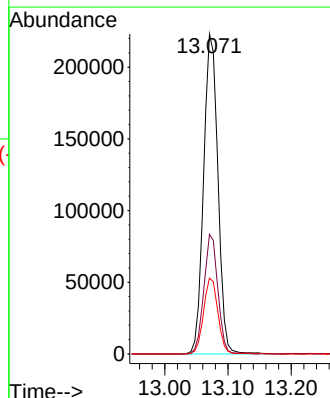
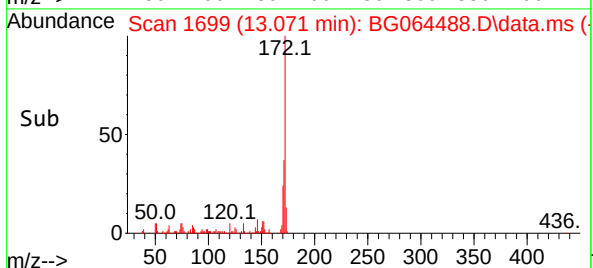
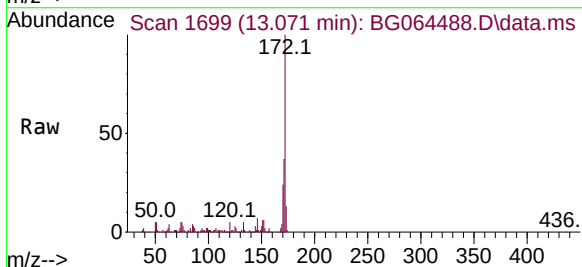


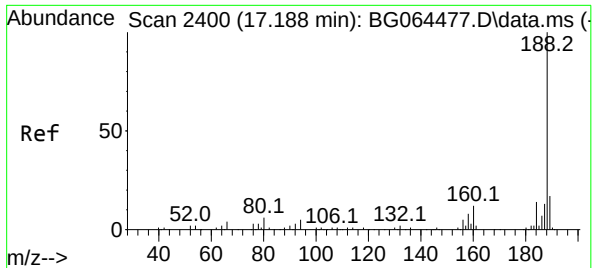
Tgt Ion:330 Resp: 142586  
Ion Ratio Lower Upper  
330 100  
332 96.3 77.4 116.0  
141 32.0 25.8 38.6



#45  
2-Fluorobiphenyl  
Concen: 90.727 ng  
RT: 13.071 min Scan# 1699  
Delta R.T. 0.003 min  
Lab File: BG064488.D  
Acq: 3 Oct 2025 15:35

Tgt Ion:172 Resp: 351343  
Ion Ratio Lower Upper  
172 100  
171 37.4 29.6 44.4  
170 23.7 20.2 30.2





#64

Phenanthrene-d10

Concen: 20.000 ng

RT: 17.189 min Scan# 2400

Delta R.T. 0.003 min

Lab File: BG064488.D

Acq: 3 Oct 2025 15:35

Instrument :

BNA\_G

ClientSampleId :

PB169973BL

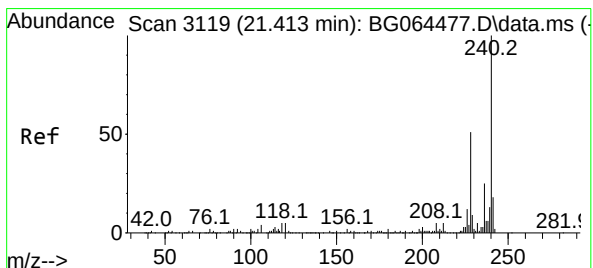
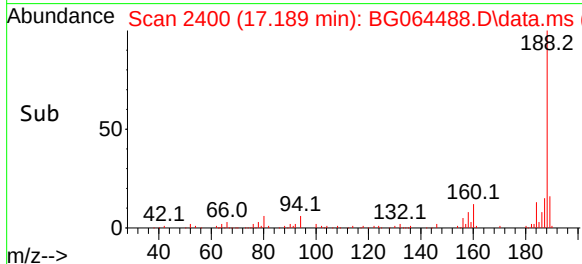
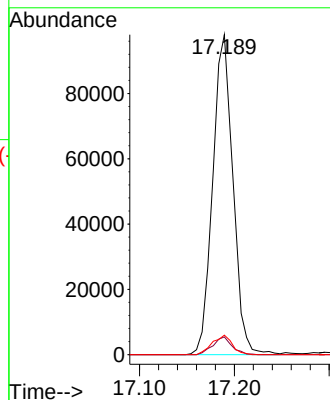
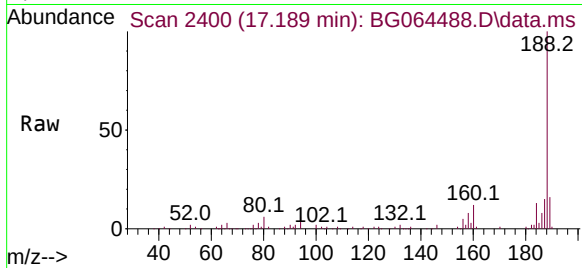
Tgt Ion:188 Resp: 145353

Ion Ratio Lower Upper

188 100

94 5.5 3.7 5.5#

80 6.1 4.5 6.7



#76

Chrysene-d12

Concen: 20.000 ng

RT: 21.408 min Scan# 3118

Delta R.T. -0.002 min

Lab File: BG064488.D

Acq: 3 Oct 2025 15:35

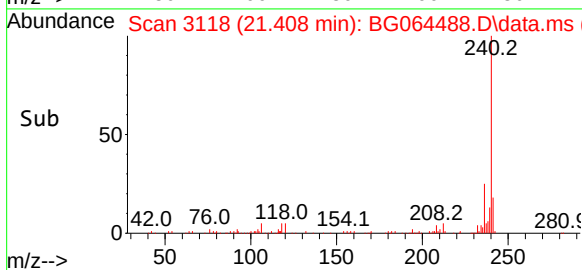
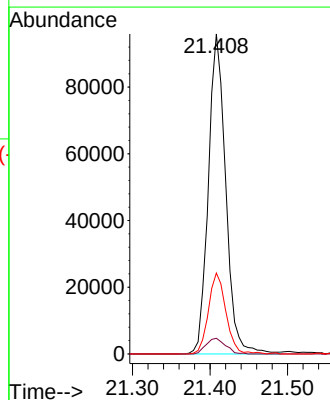
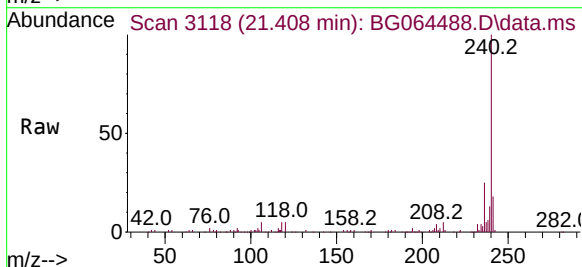
Tgt Ion:240 Resp: 149348

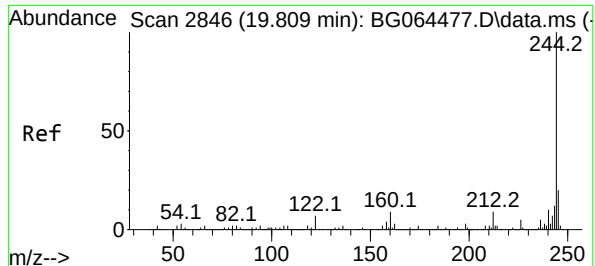
Ion Ratio Lower Upper

240 100

120 4.9 4.3 6.5

236 25.3 20.4 30.6





#79

Terphenyl-d14

Concen: 98.773 ng

RT: 19.810 min Scan# 2846

Delta R.T. 0.003 min

Lab File: BG064488.D

Acq: 3 Oct 2025 15:35

Instrument :

BNA\_G

ClientSampleId :

PB169973BL

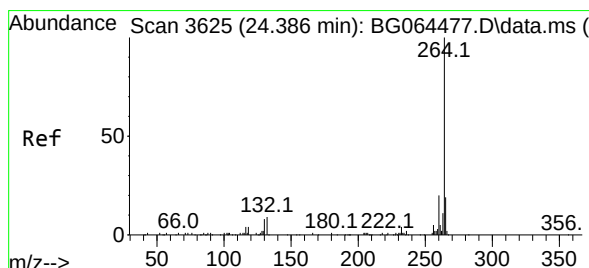
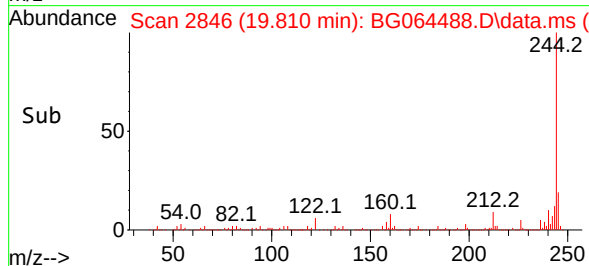
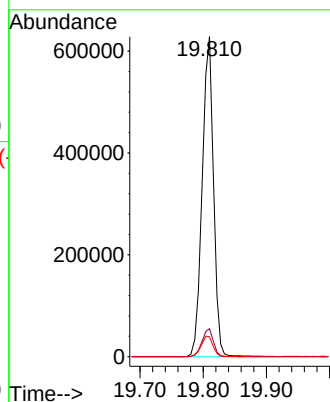
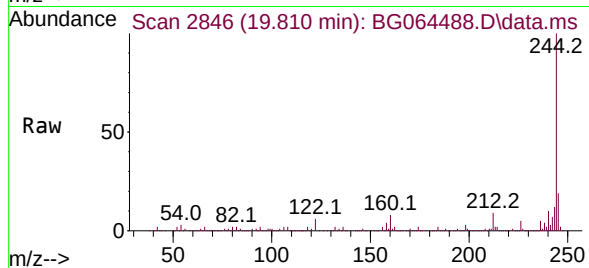
Tgt Ion:244 Resp: 779476

Ion Ratio Lower Upper

244 100

212 8.7 7.0 10.6

122 6.2 5.6 8.4



#86

Perylene-d12

Concen: 20.000 ng

RT: 24.387 min Scan# 3625

Delta R.T. -0.002 min

Lab File: BG064488.D

Acq: 3 Oct 2025 15:35

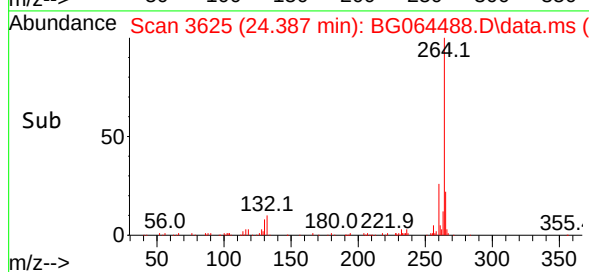
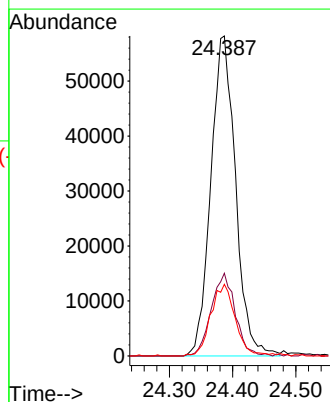
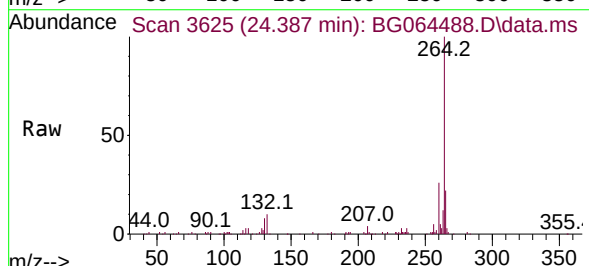
Tgt Ion:264 Resp: 157878

Ion Ratio Lower Upper

264 100

260 25.8 16.2 24.4#

265 22.4 15.0 22.4





5

A

B

C

D

E

F

G

H

I

J

K

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064488.D  
Acq On : 3 Oct 2025 15:35  
Operator : RC/JU  
Sample : PB169973BL  
Misc :  
ALS Vial : 5 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

Integration Parameters: rteint.p

Integrator: RTE

Smoothing : ON

Filtering: 5

Sampling : 1

Min Area: 3 % of largest Peak

Start Thrs: 0.2

Max Peaks: 100

Stop Thrs : 0

Peak Location: TOP

If leading or trailing edge < 100 prefer < Baseline drop else tangent >

Peak separation: 5

Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M

Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

Signal : TIC: BG064488.D\data.ms

| peak<br># | R.T.<br>min | first<br>scan | max<br>scan | last<br>scan | PK<br>TY | peak<br>height | corr.<br>area | corr.<br>% max. | % of<br>total |
|-----------|-------------|---------------|-------------|--------------|----------|----------------|---------------|-----------------|---------------|
| 1         | 5.415       | 389           | 396         | 404          | rBV      | 414112         | 622304        | 28.61%          | 8.093%        |
| 2         | 7.001       | 658           | 666         | 673          | rBV      | 393647         | 667108        | 30.67%          | 8.676%        |
| 3         | 7.824       | 799           | 806         | 813          | rBB      | 69950          | 113447        | 5.22%           | 1.475%        |
| 4         | 8.976       | 993           | 1002        | 1010         | rBV      | 242448         | 422134        | 19.41%          | 5.490%        |
| 5         | 10.609      | 1273          | 1280        | 1295         | rVB      | 82628          | 154573        | 7.11%           | 2.010%        |
| 6         | 13.071      | 1693          | 1699        | 1712         | rBV      | 702849         | 1080017       | 49.65%          | 14.046%       |
| 7         | 14.446      | 1927          | 1933        | 1944         | rBV      | 154134         | 244585        | 11.24%          | 3.181%        |
| 8         | 15.938      | 2180          | 2187        | 2201         | rBV      | 712709         | 1087253       | 49.99%          | 14.140%       |
| 9         | 17.189      | 2391          | 2400        | 2407         | rBV      | 242464         | 354840        | 16.31%          | 4.615%        |
| 10        | 19.810      | 2839          | 2846        | 2854         | rBV      | 1723358        | 2175065       | 100.00%         | 28.287%       |
| 11        | 21.408      | 3112          | 3118        | 3127         | rBV      | 243946         | 383387        | 17.63%          | 4.986%        |
| 12        | 24.387      | 3615          | 3625        | 3635         | rBV2     | 145265         | 384523        | 17.68%          | 5.001%        |

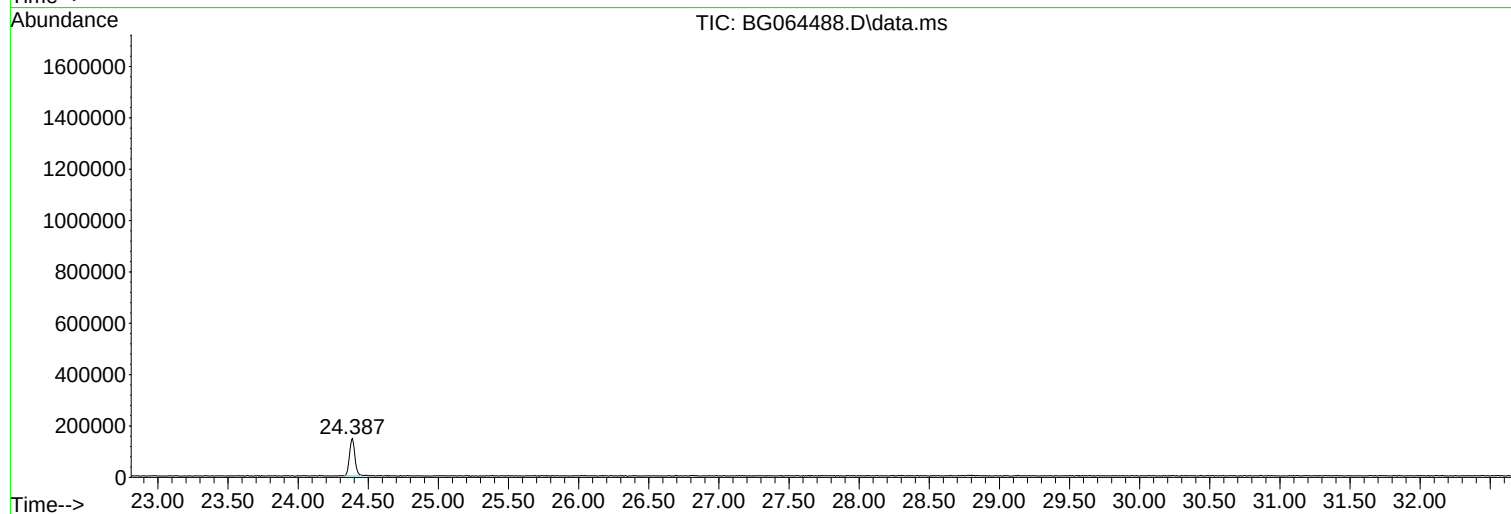
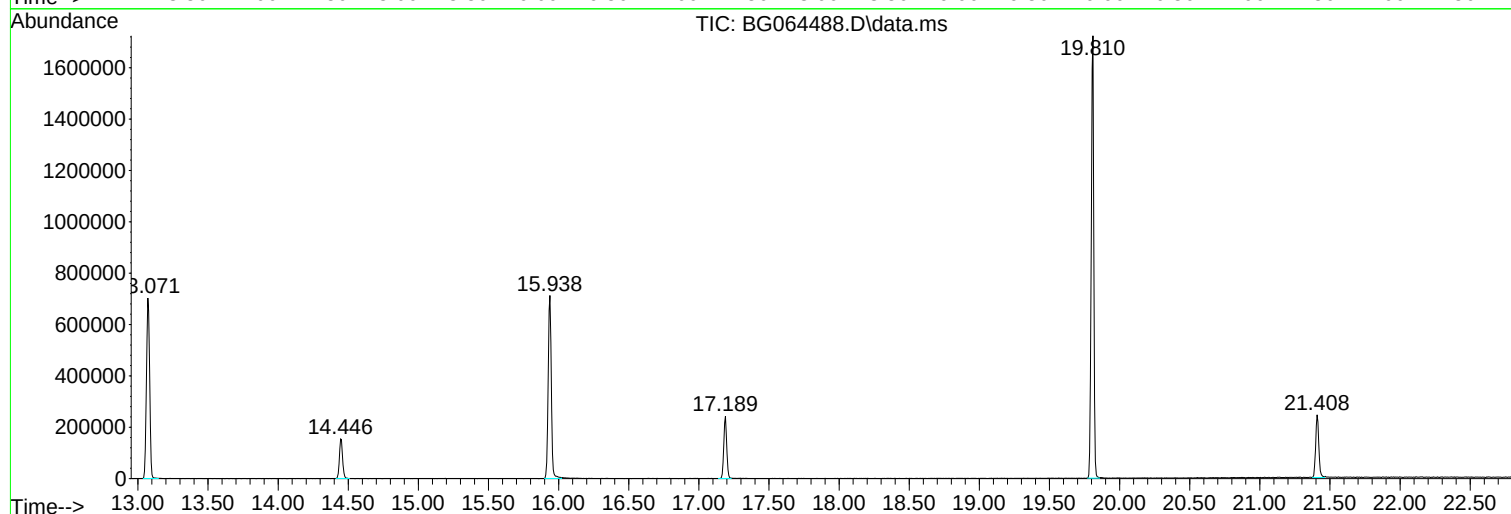
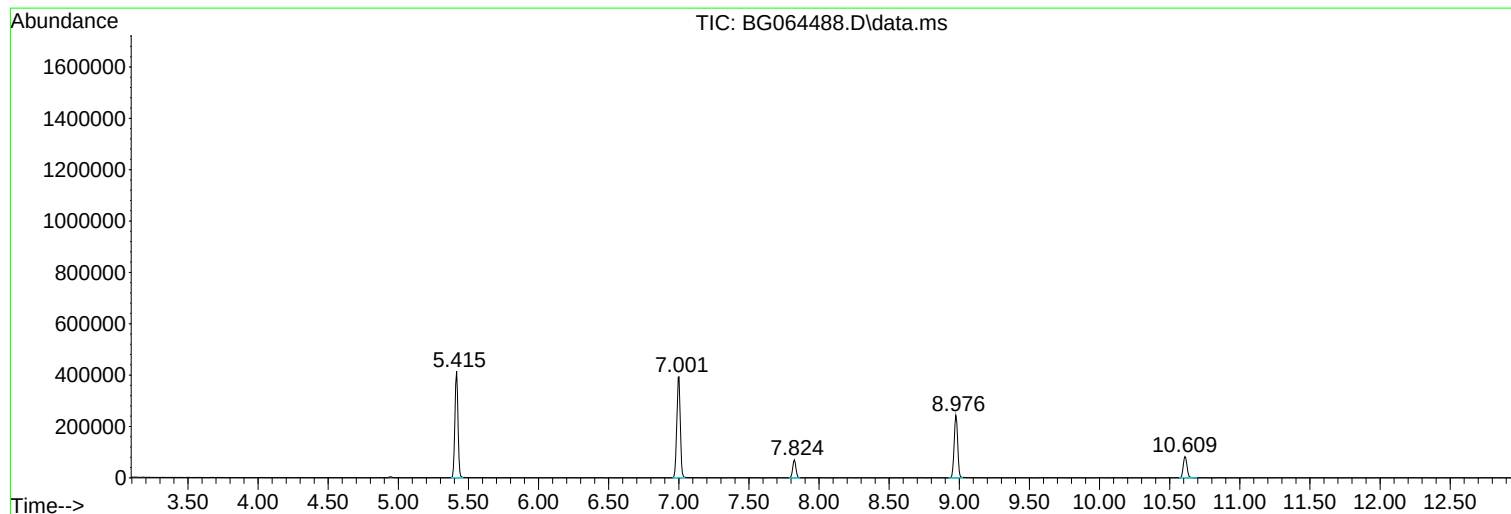
Sum of corrected areas: 7689236

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064488.D  
Acq On : 3 Oct 2025 15:35  
Operator : RC/JU  
Sample : PB169973BL  
Misc :  
ALS Vial : 5 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064488.D  
Acq On : 3 Oct 2025 15:35  
Operator : RC/JU  
Sample : PB169973BL  
Misc :  
ALS Vial : 5 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

No Library Search Compounds Detected

\*\*\*\*\*

5

A

B

C

D

E

F

G

H

I

J

K

5

A

B

C

D

E

F

G

H

I

J

K

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064488.D  
Acq On : 3 Oct 2025 15:35  
Operator : RC/JU  
Sample : PB169973BL  
Misc :  
ALS Vial : 5 Sample Multiplier: 1

Instrument :  
BNA\_G  
ClientSampleId :  
PB169973BL

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

TIC Library : C:\Database\NIST20.L  
TIC Integration Parameters: LSCINT.P

| TIC Top Hit name | RT | EstConc | Units | Response | # | RT | Resp | Conc |
|------------------|----|---------|-------|----------|---|----|------|------|
|------------------|----|---------|-------|----------|---|----|------|------|

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
 Data File : BG064489.D  
 Acq On : 3 Oct 2025 16:16  
 Operator : RC/JU  
 Sample : PB169973BS  
 Misc :  
 ALS Vial : 6 Sample Multiplier: 1

## Instrument :

BNA\_G

## ClientSampleId :

PB169973BS

## Manual Integrations

## APPROVED

Reviewed By :Rahul Chavli 10/06/2025

Supervised By :Jagrut Upadhyay 10/06/2025

Quant Time: Oct 03 17:29:58 2025

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M

Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

QLast Update : Thu Oct 02 16:40:42 2025

Response via : Initial Calibration

| Compound                      | R.T.   | QIon | Response | Conc    | Units | Dev(Min)  |
|-------------------------------|--------|------|----------|---------|-------|-----------|
| Internal Standards            |        |      |          |         |       |           |
| 1) 1,4-Dichlorobenzene-d4     | 7.823  | 152  | 16264    | 20.000  | ng    | # 0.00    |
| 21) Naphthalene-d8            | 10.608 | 136  | 69209    | 20.000  | ng    | # 0.00    |
| 39) Acenaphthene-d10          | 14.450 | 164  | 53762    | 20.000  | ng    | 0.00      |
| 64) Phenanthrene-d10          | 17.188 | 188  | 136086   | 20.000  | ng    | 0.00      |
| 76) Chrysene-d12              | 21.413 | 240  | 136322   | 20.000  | ng    | 0.00      |
| 86) Perylene-d12              | 24.391 | 264  | 146226   | 20.000  | ng    | 0.00      |
| System Monitoring Compounds   |        |      |          |         |       |           |
| 5) 2-Fluorophenol             | 5.414  | 112  | 138141   | 157.191 | ng    | 0.00      |
| 7) Phenol-d6                  | 7.000  | 99   | 180394   | 152.750 | ng    | 0.01      |
| 23) Nitrobenzene-d5           | 8.980  | 82   | 121407   | 94.324  | ng    | 0.01      |
| 42) 2,4,6-Tribromophenol      | 15.943 | 330  | 145062   | 148.856 | ng    | 0.00      |
| 45) 2-Fluorobiphenyl          | 13.075 | 172  | 334812   | 91.796  | ng    | 0.00      |
| 79) Terphenyl-d14             | 19.809 | 244  | 701104   | 97.331  | ng    | 0.00      |
| Target Compounds              |        |      |          |         |       |           |
| 2) 1,4-Dioxane                | 3.316  | 88   | 13173    | 37.896  | ng    | Qvalue 93 |
| 3) Pyridine                   | 3.710  | 79   | 35965    | 38.498  | ng    | 96        |
| 4) n-Nitrosodimethylamine     | 3.628  | 42   | 35975    | 53.120  | ng    | # 94      |
| 6) Aniline                    | 7.153  | 93   | 54677    | 36.704  | ng    | 95        |
| 8) 2-Chlorophenol             | 7.394  | 128  | 55732    | 51.645  | ng    | 95        |
| 9) Benzaldehyde               | 6.959  | 77   | 27733    | 26.454  | ng    | 97        |
| 10) Phenol                    | 7.024  | 94   | 66374    | 53.277  | ng    | 94        |
| 11) bis(2-Chloroethyl)ether   | 7.247  | 93   | 41708    | 49.435  | ng    | 93        |
| 12) 1,3-Dichlorobenzene       | 7.711  | 146  | 59993    | 50.820  | ng    | 93        |
| 13) 1,4-Dichlorobenzene       | 7.858  | 146  | 60392    | 50.973  | ng    | 94        |
| 14) 1,2-Dichlorobenzene       | 8.175  | 146  | 60785    | 52.056  | ng    | 99        |
| 15) Benzyl Alcohol            | 8.064  | 79   | 58102    | 53.422  | ng    | 93        |
| 16) 2,2'-oxybis(1-Chloropr... | 8.346  | 45   | 67381    | 49.546  | ng    | 97        |
| 17) 2-Methylphenol            | 8.269  | 107  | 47294    | 52.425  | ng    | 98        |
| 18) Hexachloroethane          | 8.904  | 117  | 23005    | 49.193  | ng    | 93        |
| 19) n-Nitroso-di-n-propyla... | 8.628  | 70   | 40266    | 50.981  | ng    | # 93      |
| 20) 3+4-Methylphenols         | 8.592  | 107  | 67040    | 53.253  | ng    | 93        |
| 22) Acetophenone              | 8.645  | 105  | 95591    | 57.044  | ng    | # 98      |
| 24) Nitrobenzene              | 9.021  | 77   | 65082    | 50.705  | ng    | 99        |
| 25) Isophorone                | 9.544  | 82   | 112788   | 48.371  | ng    | 99        |
| 26) 2-Nitrophenol             | 9.726  | 139  | 31396    | 49.147  | ng    | 93        |
| 27) 2,4-Dimethylphenol        | 9.791  | 122  | 56577    | 57.985  | ng    | 89        |
| 28) bis(2-Chloroethoxy)met... | 10.026 | 93   | 59992    | 50.474  | ng    | 95        |
| 29) 2,4-Dichlorophenol        | 10.261 | 162  | 63008    | 55.844  | ng    | 91        |
| 30) 1,2,4-Trichlorobenzene    | 10.478 | 180  | 65847    | 53.523  | ng    | # 94      |
| 31) Naphthalene               | 10.661 | 128  | 180768   | 50.722  | ng    | 98        |
| 32) Benzoic acid              | 9.967  | 122  | 46416m   | 61.961  | ng    |           |
| 33) 4-Chloroaniline           | 10.772 | 127  | 25770    | 18.843  | ng    | # 91      |
| 34) Hexachlorobutadiene       | 10.954 | 225  | 51772    | 49.138  | ng    | 97        |
| 35) Caprolactam               | 11.542 | 113  | 20070m   | 51.960  | ng    |           |
| 36) 4-Chloro-3-methylphenol   | 11.900 | 107  | 73890    | 53.851  | ng    | 100       |
| 37) 2-Methylnaphthalene       | 12.270 | 142  | 126325   | 46.176  | ng    | 98        |
| 38) 1-Methylnaphthalene       | 12.494 | 142  | 131440   | 48.184  | ng    | 95        |
| 40) 1,2,4,5-Tetrachloroben... | 12.641 | 216  | 94170    | 54.249  | ng    | # 96      |
| 41) Hexachlorocyclopentadiene | 12.623 | 237  | 132704   | 146.146 | ng    | 94        |
| 43) 2,4,6-Trichlorophenol     | 12.881 | 196  | 59397    | 53.526  | ng    | 96        |

5

A

B

C

D

E

F

G

H

I

J

K

Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
 Data File : BG064489.D  
 Acq On : 3 Oct 2025 16:16  
 Operator : RC/JU  
 Sample : PB169973BS  
 Misc :  
 ALS Vial : 6 Sample Multiplier: 1

## Instrument :

BNA\_G

## ClientSampleId :

PB169973BS

## Manual Integrations

## APPROVED

Reviewed By :Rahul Chavli 10/06/2025

Supervised By :Jagrut Upadhyay 10/06/2025

Quant Time: Oct 03 17:29:58 2025

Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M

Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION

QLast Update : Thu Oct 02 16:40:42 2025

Response via : Initial Calibration

| Compound                      | R.T.   | QIon | Response | Conc    | Units | Dev(Min) |
|-------------------------------|--------|------|----------|---------|-------|----------|
| 44) 2,4,5-Trichlorophenol     | 12.952 | 196  | 67385    | 54.393  | ng    | 96       |
| 46) 1,1'-Biphenyl             | 13.287 | 154  | 218091   | 56.402  | ng    | 98       |
| 47) 2-Chloronaphthalene       | 13.322 | 162  | 144673   | 50.112  | ng    | 98       |
| 48) 2-Nitroaniline            | 13.528 | 65   | 52229    | 53.007  | ng    | 91       |
| 49) Acenaphthylene            | 14.174 | 152  | 248338   | 57.947  | ng    | 99       |
| 50) Dimethylphthalate         | 13.910 | 163  | 219320   | 52.087  | ng    | 98       |
| 51) 2,6-Dinitrotoluene        | 14.027 | 165  | 46526    | 53.043  | ng    | 93       |
| 52) Acenaphthene              | 14.515 | 154  | 148370   | 49.831  | ng    | 98       |
| 53) 3-Nitroaniline            | 14.356 | 138  | 24285    | 30.021  | ng    | # 90     |
| 54) 2,4-Dinitrophenol         | 14.568 | 184  | 60734    | 95.491  | ng    | # 84     |
| 55) Dibenzofuran              | 14.850 | 168  | 252708   | 51.403  | ng    | 99       |
| 56) 4-Nitrophenol             | 14.679 | 139  | 70603    | 110.857 | ng    | 86       |
| 57) 2,4-Dinitrotoluene        | 14.814 | 165  | 70046    | 52.240  | ng    | # 91     |
| 58) Fluorene                  | 15.496 | 166  | 212852   | 52.080  | ng    | 98       |
| 59) 2,3,4,6-Tetrachlorophenol | 15.079 | 232  | 69146m   | 54.690  | ng    |          |
| 60) Diethylphthalate          | 15.279 | 149  | 233804   | 51.761  | ng    | 99       |
| 61) 4-Chlorophenyl-phenyle... | 15.496 | 204  | 112399   | 50.374  | ng    | 98       |
| 62) 4-Nitroaniline            | 15.525 | 138  | 45131    | 53.529  | ng    | 92       |
| 63) Azobenzene                | 15.784 | 77   | 183158   | 52.357  | ng    | 98       |
| 65) 4,6-Dinitro-2-methylph... | 15.578 | 198  | 48834    | 51.538  | ng    | 96       |
| 66) n-Nitrosodiphenylamine    | 15.708 | 169  | 190210   | 52.341  | ng    | 96       |
| 67) 4-Bromophenyl-phenylether | 16.383 | 248  | 81751    | 51.049  | ng    | 92       |
| 68) Hexachlorobenzene         | 16.501 | 284  | 93070    | 50.789  | ng    | 98       |
| 69) Atrazine                  | 16.659 | 200  | 88975    | 55.031  | ng    | 99       |
| 70) Pentachlorophenol         | 16.847 | 266  | 114041   | 106.438 | ng    | 98       |
| 71) Phenanthrene              | 17.235 | 178  | 367467   | 51.635  | ng    | 99       |
| 72) Anthracene                | 17.323 | 178  | 378287   | 52.985  | ng    | 99       |
| 73) Carbazole                 | 17.594 | 167  | 328133   | 53.172  | ng    | 98       |
| 74) Di-n-butylphthalate       | 18.158 | 149  | 396056   | 52.573  | ng    | 99       |
| 75) Fluoranthene              | 19.245 | 202  | 442925   | 52.097  | ng    | 98       |
| 77) Benzidine                 | 19.427 | 184  | 112790   | 33.250  | ng    | 97       |
| 78) Pyrene                    | 19.603 | 202  | 457350   | 51.696  | ng    | 98       |
| 80) Butylbenzylphthalate      | 20.502 | 149  | 165515   | 52.777  | ng    | 97       |
| 81) Benzo(a)anthracene        | 21.395 | 228  | 448603   | 51.538  | ng    | 98       |
| 82) 3,3'-Dichlorobenzidine    | 21.319 | 252  | 90148    | 30.600  | ng    | 98       |
| 83) Chrysene                  | 21.460 | 228  | 423453   | 51.298  | ng    | 97       |
| 84) Bis(2-ethylhexyl)phtha... | 21.319 | 149  | 233098   | 52.679  | ng    | 99       |
| 85) Di-n-octyl phthalate      | 22.441 | 149  | 386957   | 52.033  | ng    | 99       |
| 87) Indeno(1,2,3-cd)pyrene    | 27.746 | 276  | 548210   | 53.861  | ng    | 98       |
| 88) Benzo(b)fluoranthene      | 23.451 | 252  | 450951   | 54.209  | ng    | 98       |
| 89) Benzo(k)fluoranthene      | 23.516 | 252  | 433743   | 51.769  | ng    | 97       |
| 90) Benzo(a)pyrene            | 24.256 | 252  | 420603   | 54.860  | ng    | 99       |
| 91) Dibenzo(a,h)anthracene    | 27.799 | 278  | 449110   | 55.175  | ng    | 97       |
| 92) Benzo(g,h,i)perylene      | 28.792 | 276  | 433026   | 54.744  | ng    | 97       |

(#) = qualifier out of range (m) = manual integration (+) = signals summed

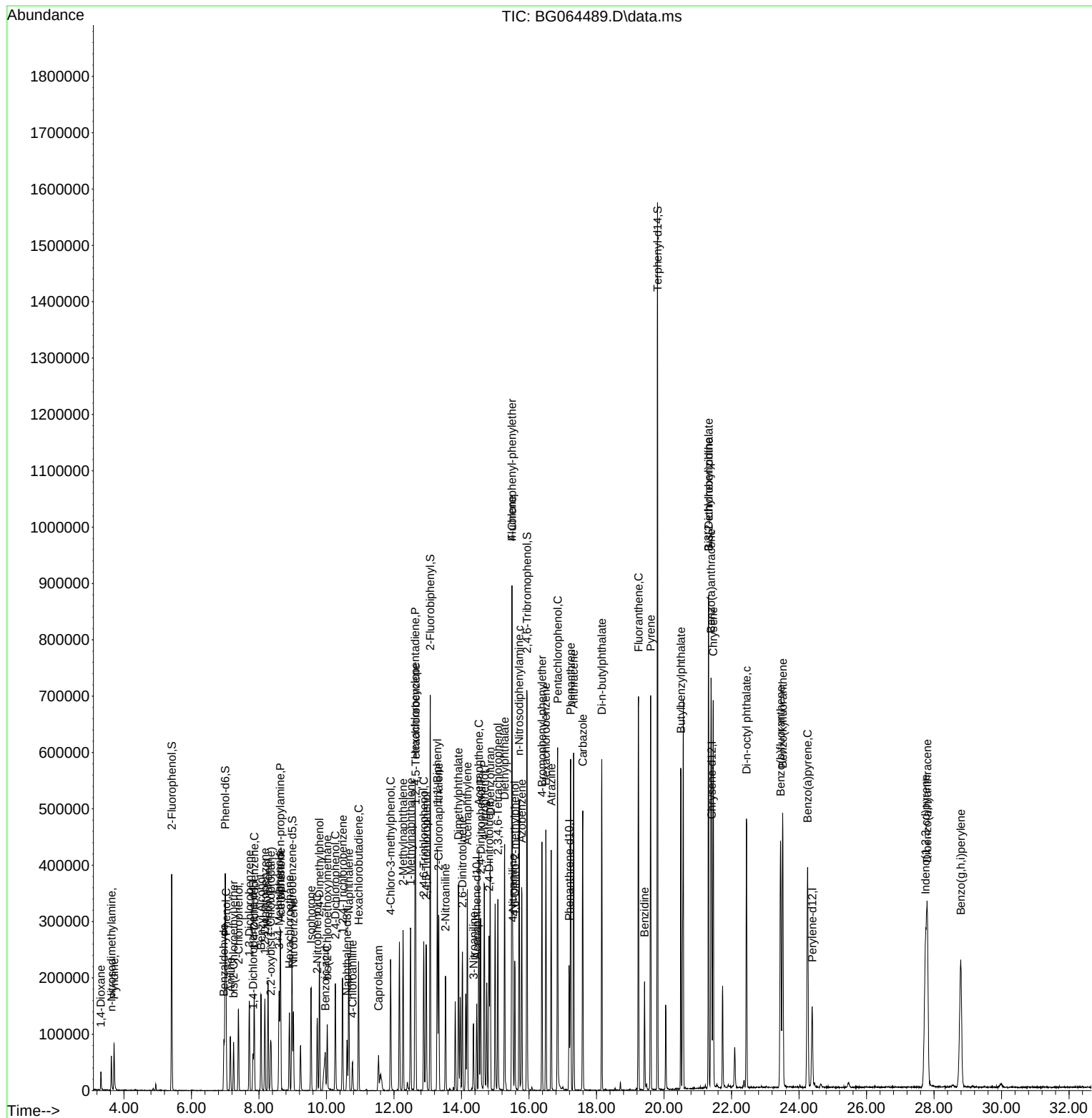
Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
Data File : BG064489.D  
Acq On : 3 Oct 2025 16:16  
Operator : RC/JU  
Sample : PB169973BS  
Misc :  
ALS Vial : 6 Sample Multiplier: 1

**Instrument :**  
BNA\_G  
**ClientSampleId :**  
PB169973BS

## Manual Integrations APPROVED

Reviewed By :Rahul Chavli 10/06/2025  
Supervised By :Jagrut Upadhyay 10/06/2025

Quant Time: Oct 03 17:29:58 2025  
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
QLast Update : Thu Oct 02 16:40:42 2025  
Response via : Initial Calibration



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
 Data File : BG064490.D  
 Acq On : 3 Oct 2025 16:57  
 Operator : RC/JU  
 Sample : PB169973BSD  
 Misc :  
 ALS Vial : 7 Sample Multiplier: 1

Instrument :  
 BNA\_G  
 ClientSampleId :  
 PB169973BSD

Manual Integrations  
 APPROVED

Reviewed By :Rahul Chavli 10/06/2025  
 Supervised By :Jagrut Upadhyay 10/06/2025

Quant Time: Oct 03 17:39:14 2025  
 Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
 Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
 QLast Update : Thu Oct 02 16:40:42 2025  
 Response via : Initial Calibration

| Compound                      | R.T.   | QIon | Response | Conc    | Units | Dev(Min) |
|-------------------------------|--------|------|----------|---------|-------|----------|
| -----                         |        |      |          |         |       |          |
| Internal Standards            |        |      |          |         |       |          |
| 1) 1,4-Dichlorobenzene-d4     | 7.825  | 152  | 15312    | 20.000  | ng    | 0.00     |
| 21) Naphthalene-d8            | 10.610 | 136  | 68886    | 20.000  | ng    | # 0.00   |
| 39) Acenaphthene-d10          | 14.453 | 164  | 53563    | 20.000  | ng    | 0.00     |
| 64) Phenanthrene-d10          | 17.191 | 188  | 138632   | 20.000  | ng    | # 0.00   |
| 76) Chrysene-d12              | 21.415 | 240  | 141515   | 20.000  | ng    | 0.00     |
| 86) Perylene-d12              | 24.388 | 264  | 151583   | 20.000  | ng    | # 0.00   |
|                               |        |      |          |         |       |          |
| System Monitoring Compounds   |        |      |          |         |       |          |
| 5) 2-Fluorophenol             | 5.416  | 112  | 134466   | 162.522 | ng    | 0.01     |
| 7) Phenol-d6                  | 7.003  | 99   | 172569   | 155.209 | ng    | 0.02     |
| 23) Nitrobenzene-d5           | 8.977  | 82   | 117218   | 91.496  | ng    | 0.01     |
| 42) 2,4,6-Tribromophenol      | 15.939 | 330  | 140210   | 144.412 | ng    | 0.00     |
| 45) 2-Fluorobiphenyl          | 13.072 | 172  | 328094   | 90.288  | ng    | 0.00     |
| 79) Terphenyl-d14             | 19.805 | 244  | 721842   | 96.533  | ng    | 0.00     |
|                               |        |      |          |         |       |          |
| Target Compounds              |        |      |          |         |       | Qvalue   |
| 2) 1,4-Dioxane                | 3.319  | 88   | 13298    | 40.634  | ng    | # 92     |
| 3) Pyridine                   | 3.706  | 79   | 34946    | 39.733  | ng    | 94       |
| 4) n-Nitrosodimethylamine     | 3.624  | 42   | 31964    | 50.132  | ng    | 87       |
| 6) Aniline                    | 7.149  | 93   | 48523    | 34.598  | ng    | 97       |
| 8) 2-Chlorophenol             | 7.390  | 128  | 56075    | 55.194  | ng    | 98       |
| 9) Benzaldehyde               | 6.961  | 77   | 25788    | 26.128  | ng    | 91       |
| 10) Phenol                    | 7.026  | 94   | 64826    | 55.269  | ng    | 98       |
| 11) bis(2-Chloroethyl)ether   | 7.249  | 93   | 39465    | 49.685  | ng    | 95       |
| 12) 1,3-Dichlorobenzene       | 7.713  | 146  | 55884    | 50.283  | ng    | 98       |
| 13) 1,4-Dichlorobenzene       | 7.860  | 146  | 59496    | 53.339  | ng    | 91       |
| 14) 1,2-Dichlorobenzene       | 8.172  | 146  | 55839    | 50.794  | ng    | 96       |
| 15) Benzyl Alcohol            | 8.060  | 79   | 55796    | 54.492  | ng    | 90       |
| 16) 2,2'-oxybis(1-Chloropr... | 8.348  | 45   | 61797    | 48.266  | ng    | 98       |
| 17) 2-Methylphenol            | 8.272  | 107  | 46672    | 54.952  | ng    | 98       |
| 18) Hexachloroethane          | 8.900  | 117  | 21715    | 49.321  | ng    | 92       |
| 19) n-Nitroso-di-n-propyla... | 8.624  | 70   | 38963    | 52.398  | ng    | 97       |
| 20) 3+4-Methylphenols         | 8.595  | 107  | 65493    | 55.259  | ng    | 93       |
| 22) Acetophenone              | 8.642  | 105  | 91544    | 54.885  | ng    | # 96     |
| 24) Nitrobenzene              | 9.018  | 77   | 60821    | 47.607  | ng    | 96       |
| 25) Isophorone                | 9.541  | 82   | 110698   | 47.697  | ng    | 98       |
| 26) 2-Nitrophenol             | 9.729  | 139  | 30795    | 48.432  | ng    | 97       |
| 27) 2,4-Dimethylphenol        | 9.793  | 122  | 55310    | 56.953  | ng    | 95       |
| 28) bis(2-Chloroethoxy)met... | 10.023 | 93   | 58363    | 49.333  | ng    | 97       |
| 29) 2,4-Dichlorophenol        | 10.263 | 162  | 59120    | 52.643  | ng    | 96       |
| 30) 1,2,4-Trichlorobenzene    | 10.475 | 180  | 61294    | 50.056  | ng    | 98       |
| 31) Naphthalene               | 10.663 | 128  | 171886   | 48.456  | ng    | 99       |
| 32) Benzoic acid              | 9.958  | 122  | 44961m   | 60.300  | ng    |          |
| 33) 4-Chloroaniline           | 10.769 | 127  | 20695    | 15.203  | ng    | # 97     |
| 34) Hexachlorobutadiene       | 10.951 | 225  | 49245    | 46.959  | ng    | 99       |
| 35) Caprolactam               | 11.538 | 113  | 19269m   | 50.120  | ng    |          |
| 36) 4-Chloro-3-methylphenol   | 11.903 | 107  | 70424    | 51.566  | ng    | 97       |
| 37) 2-Methylnaphthalene       | 12.273 | 142  | 121185   | 44.505  | ng    | 98       |
| 38) 1-Methylnaphthalene       | 12.490 | 142  | 126687   | 46.659  | ng    | 100      |
| 40) 1,2,4,5-Tetrachloroben... | 12.643 | 216  | 92935    | 53.737  | ng    | 98       |
| 41) Hexachlorocyclopentadiene | 12.625 | 237  | 123269   | 136.260 | ng    | 94       |
| 43) 2,4,6-Trichlorophenol     | 12.884 | 196  | 57388    | 51.908  | ng    | 99       |



Data Path : Z:\svoasrv\HPCHEM1\BNA\_G\Data\BG100325\  
 Data File : BG064490.D  
 Acq On : 3 Oct 2025 16:57  
 Operator : RC/JU  
 Sample : PB169973BSD  
 Misc :  
 ALS Vial : 7 Sample Multiplier: 1

Instrument :  
 BNA\_G  
 ClientSampleId :  
 PB169973BSD

### Manual Integrations APPROVED

Reviewed By :Rahul Chavli 10/06/2025  
 Supervised By :Jagrut Upadhyay 10/06/2025

Quant Time: Oct 03 17:39:14 2025  
 Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
 Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
 QLast Update : Thu Oct 02 16:40:42 2025  
 Response via : Initial Calibration

| Compound                      | R.T.   | QIon | Response | Conc    | Units | Dev(Min) |
|-------------------------------|--------|------|----------|---------|-------|----------|
| 44) 2,4,5-Trichlorophenol     | 12.954 | 196  | 60489    | 49.008  | ng    | 95       |
| 46) 1,1'-Biphenyl             | 13.283 | 154  | 204884   | 53.183  | ng    | 99       |
| 47) 2-Chloronaphthalene       | 13.325 | 162  | 136452   | 47.440  | ng    | 96       |
| 48) 2-Nitroaniline            | 13.530 | 65   | 49637    | 50.563  | ng    | 96       |
| 49) Acenaphthylene            | 14.171 | 152  | 236815   | 55.463  | ng    | 99       |
| 50) Dimethylphthalate         | 13.912 | 163  | 203802   | 48.581  | ng    | 97       |
| 51) 2,6-Dinitrotoluene        | 14.024 | 165  | 43905    | 50.241  | ng    | 95       |
| 52) Acenaphthene              | 14.517 | 154  | 140204   | 47.264  | ng    | 95       |
| 53) 3-Nitroaniline            | 14.353 | 138  | 21048    | 26.116  | ng    | 93       |
| 54) 2,4-Dinitrophenol         | 14.564 | 184  | 60905    | 96.115  | ng    | 86       |
| 55) Dibenzofuran              | 14.852 | 168  | 239756   | 48.950  | ng    | 99       |
| 56) 4-Nitrophenol             | 14.676 | 139  | 70541    | 111.171 | ng    | 97       |
| 57) 2,4-Dinitrotoluene        | 14.817 | 165  | 71035    | 53.174  | ng    | 99       |
| 58) Fluorene                  | 15.498 | 166  | 204659   | 50.261  | ng    | 98       |
| 59) 2,3,4,6-Tetrachlorophenol | 15.081 | 232  | 68433    | 54.327  | ng    | 98       |
| 60) Diethylphthalate          | 15.275 | 149  | 222549   | 49.452  | ng    | # 97     |
| 61) 4-Chlorophenyl-phenyle... | 15.493 | 204  | 106708   | 48.001  | ng    | 96       |
| 62) 4-Nitroaniline            | 15.522 | 138  | 43915    | 52.280  | ng    | 89       |
| 63) Azobenzene                | 15.786 | 77   | 173819   | 49.872  | ng    | 96       |
| 65) 4,6-Dinitro-2-methylph... | 15.581 | 198  | 46256    | 47.921  | ng    | 97       |
| 66) n-Nitrosodiphenylamine    | 15.710 | 169  | 182340   | 49.254  | ng    | 93       |
| 67) 4-Bromophenyl-phenylether | 16.386 | 248  | 78451    | 48.089  | ng    | 90       |
| 68) Hexachlorobenzene         | 16.503 | 284  | 91352    | 48.936  | ng    | 98       |
| 69) Atrazine                  | 16.656 | 200  | 87001    | 52.822  | ng    | 95       |
| 70) Pentachlorophenol         | 16.850 | 266  | 109925   | 100.712 | ng    | 100      |
| 71) Phenanthrene              | 17.232 | 178  | 352360   | 48.603  | ng    | 97       |
| 72) Anthracene                | 17.326 | 178  | 362889   | 49.895  | ng    | 98       |
| 73) Carbazole                 | 17.590 | 167  | 327423   | 52.082  | ng    | 99       |
| 74) Di-n-butylphthalate       | 18.154 | 149  | 396998   | 51.730  | ng    | 100      |
| 75) Fluoranthene              | 19.241 | 202  | 441298   | 50.952  | ng    | 99       |
| 77) Benzidine                 | 19.423 | 184  | 100027   | 28.405  | ng    | 96       |
| 78) Pyrene                    | 19.605 | 202  | 454934   | 49.536  | ng    | 98       |
| 80) Butylbenzylphthalate      | 20.498 | 149  | 165196   | 50.742  | ng    | 99       |
| 81) Benzo(a)anthracene        | 21.397 | 228  | 451659   | 49.985  | ng    | 99       |
| 82) 3,3'-Dichlorobenzidine    | 21.321 | 252  | 83367    | 27.260  | ng    | # 96     |
| 83) Chrysene                  | 21.456 | 228  | 424681   | 49.559  | ng    | 96       |
| 84) Bis(2-ethylhexyl)phtha... | 21.321 | 149  | 234171   | 50.979  | ng    | 100      |
| 85) Di-n-octyl phthalate      | 22.443 | 149  | 388583   | 50.334  | ng    | 99       |
| 87) Indeno(1,2,3-cd)pyrene    | 27.743 | 276  | 540606   | 51.237  | ng    | # 97     |
| 88) Benzo(b)fluoranthene      | 23.454 | 252  | 434847   | 50.425  | ng    | 99       |
| 89) Benzo(k)fluoranthene      | 23.513 | 252  | 449174   | 51.716  | ng    | 100      |
| 90) Benzo(a)pyrene            | 24.253 | 252  | 423237   | 53.252  | ng    | 97       |
| 91) Dibenzo(a,h)anthracene    | 27.796 | 278  | 442427   | 52.434  | ng    | 98       |
| 92) Benzo(g,h,i)perylene      | 28.789 | 276  | 432535   | 52.749  | ng    | 98       |

(#) = qualifier out of range (m) = manual integration (+) = signals summed

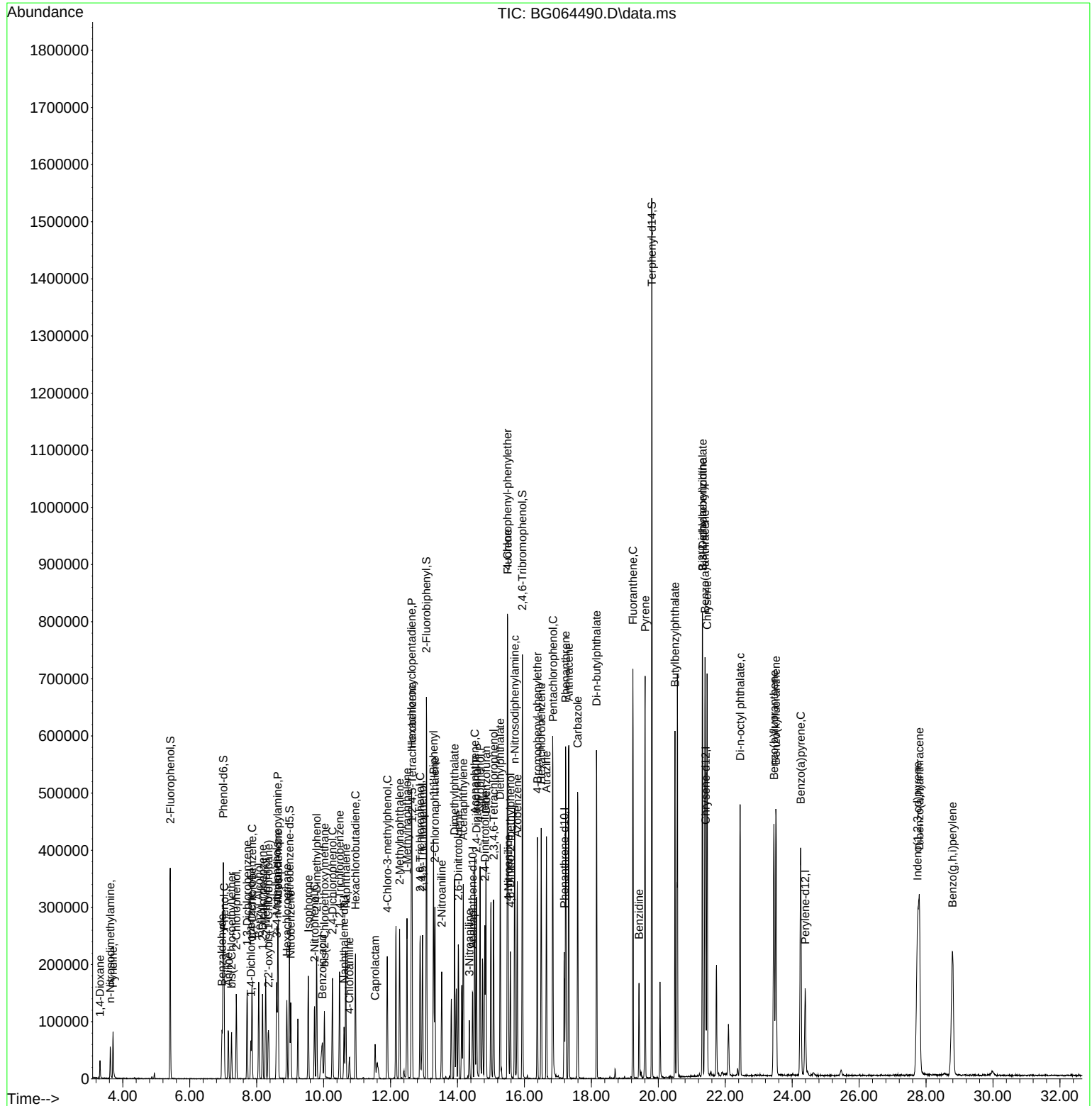
```
Data Path : Z:\svoasrv\HPCHEM1\BNA_G\Data\BG100325\  
Data File : BG064490.D  
Acq On    : 3 Oct 2025 16:57  
Operator  : RC/JU  
Sample    : PB169973BSD  
Misc      :  
ALS Vial  : 7 Sample Multiplier: 1
```

**Instrument :**  
BNA\_G  
**ClientSampleId :**  
PB169973BSD

## Manual Integrations

Reviewed By :Rahul Chavli 10/06/2025  
Supervised By :Jagrut Upadhyay 10/06/2025

Quant Time: Oct 03 17:39:14 2025  
Quant Method : Z:\svoasrv\HPCHEM1\BNA\_G\Methods\8270-BG100225.M  
Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION  
QLast Update : Thu Oct 02 16:40:42 2025  
Response via : Initial Calibration



## Manual Integration Report

|           |          |            |       |
|-----------|----------|------------|-------|
| Sequence: | BG100225 | Instrument | BNA_g |
|-----------|----------|------------|-------|

| Sample ID   | File ID    | Parameter            | Review By | Review On            | Supervised By | Supervised On        | Reason                      |
|-------------|------------|----------------------|-----------|----------------------|---------------|----------------------|-----------------------------|
| SSTDICC005  | BG064474.D | Anthracene           | Rahul     | 10/3/2025 8:37:57 AM | Jagrut        | 10/3/2025 3:25:00 PM | Peak Integrated by Software |
| SSTDICC005  | BG064474.D | Benzo(b)fluoranthene | Rahul     | 10/3/2025 8:37:57 AM | Jagrut        | 10/3/2025 3:25:00 PM | Peak Integrated by Software |
| SSTDICC020  | BG064476.D | 1-Methylnaphthalene  | Rahul     | 10/3/2025 8:38:04 AM | Jagrut        | 10/3/2025 3:25:05 PM | Peak Integrated by Software |
| SSTDICC020  | BG064476.D | Benzaldehyde         | Rahul     | 10/3/2025 8:38:04 AM | Jagrut        | 10/3/2025 3:25:05 PM | Peak Integrated by Software |
| SSTDICC020  | BG064476.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:04 AM | Jagrut        | 10/3/2025 3:25:05 PM | Peak Integrated by Software |
| SSTDICCC040 | BG064477.D | Benzaldehyde         | Rahul     | 10/3/2025 8:38:08 AM | Jagrut        | 10/3/2025 3:25:08 PM | Peak Integrated by Software |
| SSTDICCC040 | BG064477.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:08 AM | Jagrut        | 10/3/2025 3:25:08 PM | Peak Integrated by Software |
| SSTDICC050  | BG064478.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:10 AM | Jagrut        | 10/3/2025 3:25:09 PM | Peak Integrated by Software |
| SSTDICC060  | BG064479.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:13 AM | Jagrut        | 10/3/2025 3:25:12 PM | Peak Integrated by Software |
| SSTDICC080  | BG064480.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:15 AM | Jagrut        | 10/3/2025 3:25:14 PM | Peak Integrated by Software |
| SSTDICC010  | BG064481.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:18 AM | Jagrut        | 10/3/2025 3:25:20 PM | Peak Integrated by Software |
| SSTDICV040  | BG064482.D | Benzoic acid         | Rahul     | 10/3/2025 8:38:21 AM | Jagrut        | 10/3/2025 3:25:22 PM | Peak Integrated by Software |



Manual Integration Report

|           |          |            |       |
|-----------|----------|------------|-------|
| Sequence: | BG100225 | Instrument | BNA_g |
|-----------|----------|------------|-------|

| Sample ID | File ID | Parameter | Review By | Review On | Supervised By | Supervised On | Reason |
|-----------|---------|-----------|-----------|-----------|---------------|---------------|--------|
|-----------|---------|-----------|-----------|-----------|---------------|---------------|--------|

## Manual Integration Report

|           |          |            |       |
|-----------|----------|------------|-------|
| Sequence: | BG100325 | Instrument | BNA_g |
|-----------|----------|------------|-------|

| Sample ID   | File ID    | Parameter                     | Review By | Review On            | Supervised By | Supervised On        | Reason                      |
|-------------|------------|-------------------------------|-----------|----------------------|---------------|----------------------|-----------------------------|
| SSTDCCC040  | BG064485.D | 1-Methylnaphthalene           | Rahul     | 10/6/2025 4:14:13 PM | Jagrut        | 10/6/2025 4:23:08 PM | Peak Integrated by Software |
| SSTDCCC040  | BG064485.D | Benzaldehyde                  | Rahul     | 10/6/2025 4:14:13 PM | Jagrut        | 10/6/2025 4:23:08 PM | Peak Integrated by Software |
| SSTDCCC040  | BG064485.D | Benzoic acid                  | Rahul     | 10/6/2025 4:14:13 PM | Jagrut        | 10/6/2025 4:23:08 PM | Peak Integrated by Software |
| PB169973BS  | BG064489.D | 2,3,4,6-Tetrachlorophen<br>ol | Rahul     | 10/6/2025 4:14:18 PM | Jagrut        | 10/6/2025 4:23:13 PM | Peak Integrated by Software |
| PB169973BS  | BG064489.D | Benzoic acid                  | Rahul     | 10/6/2025 4:14:18 PM | Jagrut        | 10/6/2025 4:23:13 PM | Peak Integrated by Software |
| PB169973BS  | BG064489.D | Caprolactam                   | Rahul     | 10/6/2025 4:14:18 PM | Jagrut        | 10/6/2025 4:23:13 PM | Peak Integrated by Software |
| PB169973BSD | BG064490.D | Benzoic acid                  | Rahul     | 10/6/2025 4:14:20 PM | Jagrut        | 10/6/2025 4:23:34 PM | Peak Integrated by Software |
| PB169973BSD | BG064490.D | Caprolactam                   | Rahul     | 10/6/2025 4:14:20 PM | Jagrut        | 10/6/2025 4:23:34 PM | Peak Integrated by Software |

Instrument ID: BNA\_G

Daily Analysis Runlog For Sequence/QC Batch ID # BG100225

| Review By                | Rahul    | Review On                                               | 10/3/2025 8:38:44 AM |                      |          |
|--------------------------|----------|---------------------------------------------------------|----------------------|----------------------|----------|
| Supervise By             | Jagrut   | Supervise On                                            | 10/3/2025 3:25:38 PM |                      |          |
| SubDirectory             | BG100225 | HP Acquire Method                                       | BNA_G                | HP Processing Method | bg100225 |
| STD. NAME                |          | STD REF.#                                               |                      |                      |          |
| Tune/Reschk              |          | SP6875                                                  |                      |                      |          |
| Initial Calibration Stds |          | SP6857,SP6858,SP6859,SP6860,SP6861,SP6862,SP6863,SP6864 |                      |                      |          |
| CCC                      |          | SP6860                                                  |                      |                      |          |
| Internal Standard/PEM    |          | S13176,10ul/1000ul sample                               |                      |                      |          |
| ICV/I.BLK                |          | SP6872                                                  |                      |                      |          |
| Surrogate Standard       |          |                                                         |                      |                      |          |
| MS/MSD Standard          |          |                                                         |                      |                      |          |
| LCS Standard             |          |                                                         |                      |                      |          |

| Sr# | SampleId    | Data File Name | Date-Time        | Operator | Status |
|-----|-------------|----------------|------------------|----------|--------|
| 1   | DFTPP       | BG064472.D     | 2 Oct 2025 8:21  | RC/JU    | Ok     |
| 2   | SSTDICC2.5  | BG064473.D     | 2 Oct 2025 9:01  | RC/JU    | Ok     |
| 3   | SSTDICC005  | BG064474.D     | 2 Oct 2025 9:42  | RC/JU    | Ok,M   |
| 4   | SSTDICC010  | BG064475.D     | 2 Oct 2025 10:53 | RC/JU    | Not Ok |
| 5   | SSTDICC020  | BG064476.D     | 2 Oct 2025 11:33 | RC/JU    | Ok,M   |
| 6   | SSTDICCC040 | BG064477.D     | 2 Oct 2025 12:14 | RC/JU    | Ok,M   |
| 7   | SSTDICC050  | BG064478.D     | 2 Oct 2025 12:55 | RC/JU    | Ok,M   |
| 8   | SSTDICC060  | BG064479.D     | 2 Oct 2025 13:35 | RC/JU    | Ok,M   |
| 9   | SSTDICC080  | BG064480.D     | 2 Oct 2025 14:16 | RC/JU    | Ok,M   |
| 10  | SSTDICC010  | BG064481.D     | 2 Oct 2025 15:49 | RC/JU    | Ok,M   |
| 11  | SSTDICV040  | BG064482.D     | 2 Oct 2025 18:09 | RC/JU    | Ok,M   |
| 12  | PB169936BL  | BG064483.D     | 2 Oct 2025 18:50 | RC/JU    | Not Ok |

M : Manual Integration

Instrument ID: BNA\_G

Daily Analysis Runlog For Sequence/QC Batch ID # BG100325

|                          |          |                                                         |                      |                      |          |
|--------------------------|----------|---------------------------------------------------------|----------------------|----------------------|----------|
| Review By                | Rahul    | Review On                                               | 10/6/2025 4:14:36 PM |                      |          |
| Supervise By             | Jagrut   | Supervise On                                            | 10/6/2025 4:23:45 PM |                      |          |
| SubDirectory             | BG100325 | HP Acquire Method                                       | BNA_G                | HP Processing Method | bg100225 |
| STD. NAME                |          | STD REF.#                                               |                      |                      |          |
| Tune/Reschk              |          | SP6875                                                  |                      |                      |          |
| Initial Calibration Stds |          | SP6857,SP6858,SP6859,SP6860,SP6861,SP6862,SP6863,SP6864 |                      |                      |          |
| CCC                      |          | SP6860                                                  |                      |                      |          |
| Internal Standard/PEM    |          | S13177,10ul/1000ul sample                               |                      |                      |          |
| ICV/I.BLK                |          | SP6872                                                  |                      |                      |          |
| Surrogate Standard       |          |                                                         |                      |                      |          |
| MS/MSD Standard          |          |                                                         |                      |                      |          |
| LCS Standard             |          |                                                         |                      |                      |          |

| Sr# | SampleId    | Data File Name | Date-Time        | Operator | Status |
|-----|-------------|----------------|------------------|----------|--------|
| 1   | DFTPP       | BG064484.D     | 3 Oct 2025 12:52 | RC/JU    | Ok     |
| 2   | SSTDCCC040  | BG064485.D     | 3 Oct 2025 13:33 | RC/JU    | Ok,M   |
| 3   | PB169969BL  | BG064486.D     | 3 Oct 2025 14:13 | RC/JU    | Ok     |
| 4   | PB169969BS  | BG064487.D     | 3 Oct 2025 14:54 | RC/JU    | Ok,M   |
| 5   | PB169973BL  | BG064488.D     | 3 Oct 2025 15:35 | RC/JU    | Ok     |
| 6   | PB169973BS  | BG064489.D     | 3 Oct 2025 16:16 | RC/JU    | Ok,M   |
| 7   | PB169973BSD | BG064490.D     | 3 Oct 2025 16:57 | RC/JU    | Ok,M   |
| 8   | Q3263-01    | BG064491.D     | 3 Oct 2025 17:38 | RC/JU    | Ok     |
| 9   | Q3263-02    | BG064492.D     | 3 Oct 2025 18:19 | RC/JU    | Ok     |
| 10  | Q3273-01    | BG064493.D     | 3 Oct 2025 18:59 | RC/JU    | Ok     |
| 11  | Q3274-01    | BG064494.D     | 3 Oct 2025 19:40 | RC/JU    | Ok     |
| 12  | Q3272-06    | BG064495.D     | 3 Oct 2025 20:21 | RC/JU    | Ok     |

M : Manual Integration

Instrument ID: BNA\_G

**Daily Analysis Runlog For Sequence/QC Batch ID # BG100225**

|              |          |                      |                      |
|--------------|----------|----------------------|----------------------|
| Review By    | Rahul    | Review On            | 10/3/2025 8:38:44 AM |
| Supervise By | Jagrut   | Supervise On         | 10/3/2025 3:25:38 PM |
| SubDirectory | BG100225 | HP Acquire Method    | BNA_G                |
|              |          | HP Processing Method | bg100225             |

| STD. NAME                | STD REF.#                                               |
|--------------------------|---------------------------------------------------------|
| Tune/Reschk              | SP6875                                                  |
| Initial Calibration Stds | SP6857,SP6858,SP6859,SP6860,SP6861,SP6862,SP6863,SP6864 |
| CCC                      | SP6860                                                  |
| Internal Standard/PEM    | S13176,10ul/1000ul sample                               |
| ICV/I.BLK                | SP6872                                                  |
| Surrogate Standard       |                                                         |
| MS/MSD Standard          |                                                         |
| LCS Standard             |                                                         |

| Sr# | SampleId    | ClientID    | Data File Name | Date-Time        | Comment                                 | Operator | Status |
|-----|-------------|-------------|----------------|------------------|-----------------------------------------|----------|--------|
| 1   | DFTPP       | DFTPP       | BG064472.D     | 2 Oct 2025 8:21  |                                         | RC/JU    | Ok     |
| 2   | SSTDICC2.5  | SSTDICC2.5  | BG064473.D     | 2 Oct 2025 9:01  |                                         | RC/JU    | Ok     |
| 3   | SSTDICC005  | SSTDICC005  | BG064474.D     | 2 Oct 2025 9:42  |                                         | RC/JU    | Ok,M   |
| 4   | SSTDICC010  | SSTDICC010  | BG064475.D     | 2 Oct 2025 10:53 | Not Used                                | RC/JU    | Not Ok |
| 5   | SSTDICC020  | SSTDICC020  | BG064476.D     | 2 Oct 2025 11:33 |                                         | RC/JU    | Ok,M   |
| 6   | SSTDICCC040 | SSTDICCC040 | BG064477.D     | 2 Oct 2025 12:14 | Compound failed in the calibration. #77 | RC/JU    | Ok,M   |
| 7   | SSTDICC050  | SSTDICC050  | BG064478.D     | 2 Oct 2025 12:55 |                                         | RC/JU    | Ok,M   |
| 8   | SSTDICC060  | SSTDICC060  | BG064479.D     | 2 Oct 2025 13:35 |                                         | RC/JU    | Ok,M   |
| 9   | SSTDICC080  | SSTDICC080  | BG064480.D     | 2 Oct 2025 14:16 |                                         | RC/JU    | Ok,M   |
| 10  | SSTDICC010  | SSTDICC010  | BG064481.D     | 2 Oct 2025 15:49 |                                         | RC/JU    | Ok,M   |
| 11  | SSTDICV040  | ICVBG100225 | BG064482.D     | 2 Oct 2025 18:09 |                                         | RC/JU    | Ok,M   |
| 12  | PB169936BL  | PB169936BL  | BG064483.D     | 2 Oct 2025 18:50 | Surrogate Fail                          | RC/JU    | Not Ok |

M : Manual Integration



Instrument ID: BNA\_G

**Daily Analysis Runlog For Sequence/QC Batch ID # BG100325**

|              |          |                      |                      |
|--------------|----------|----------------------|----------------------|
| Review By    | Rahul    | Review On            | 10/6/2025 4:14:36 PM |
| Supervise By | Jagrut   | Supervise On         | 10/6/2025 4:23:45 PM |
| SubDirectory | BG100325 | HP Acquire Method    | BNA_G                |
|              |          | HP Processing Method | bg100225             |

| STD. NAME                | STD REF.#                                               |
|--------------------------|---------------------------------------------------------|
| Tune/Reschk              | SP6875                                                  |
| Initial Calibration Stds | SP6857,SP6858,SP6859,SP6860,SP6861,SP6862,SP6863,SP6864 |
| CCC                      | SP6860                                                  |
| Internal Standard/PEM    | S13177,10ul/1000ul sample                               |
| ICV/I.BLK                | SP6872                                                  |
| Surrogate Standard       |                                                         |
| MS/MSD Standard          |                                                         |
| LCS Standard             |                                                         |

| Sr# | SampleID    | ClientID        | Data File Name | Date-Time        | Comment | Operator | Status |
|-----|-------------|-----------------|----------------|------------------|---------|----------|--------|
| 1   | DFTPP       | DFTPP           | BG064484.D     | 3 Oct 2025 12:52 |         | RC/JU    | Ok     |
| 2   | SSTDCCC040  | SSTDCCC040      | BG064485.D     | 3 Oct 2025 13:33 |         | RC/JU    | Ok,M   |
| 3   | PB169969BL  | PB169969BL      | BG064486.D     | 3 Oct 2025 14:13 |         | RC/JU    | Ok     |
| 4   | PB169969BS  | PB169969BS      | BG064487.D     | 3 Oct 2025 14:54 |         | RC/JU    | Ok,M   |
| 5   | PB169973BL  | PB169973BL      | BG064488.D     | 3 Oct 2025 15:35 |         | RC/JU    | Ok     |
| 6   | PB169973BS  | PB169973BS      | BG064489.D     | 3 Oct 2025 16:16 |         | RC/JU    | Ok,M   |
| 7   | PB169973BSD | PB169973BSD     | BG064490.D     | 3 Oct 2025 16:57 |         | RC/JU    | Ok,M   |
| 8   | Q3263-01    | 251818          | BG064491.D     | 3 Oct 2025 17:38 |         | RC/JU    | Ok     |
| 9   | Q3263-02    | 266380          | BG064492.D     | 3 Oct 2025 18:19 |         | RC/JU    | Ok     |
| 10  | Q3273-01    | MW1             | BG064493.D     | 3 Oct 2025 18:59 |         | RC/JU    | Ok     |
| 11  | Q3274-01    | MW4             | BG064494.D     | 3 Oct 2025 19:40 |         | RC/JU    | Ok     |
| 12  | Q3272-06    | RT5417-CONCRETE | BG064495.D     | 3 Oct 2025 20:21 |         | RC/JU    | Ok     |

M : Manual Integration

**SOP ID:** M3510C,3580A-Extraction SVOC-21

**Clean Up SOP #:** N/A

**Matrix :** Water

**Weigh By:** N/A

**Balance check:** N/A

**Balance ID:** N/A

**pH Strip Lot#:** E3953

**Extraction By:** RJ

**Filter By:** RJ

**pH Meter ID:** N/A

**Hood ID:** 4,6,7

**Extraction Start Date :** 10/03/2025

**Extraction Start Time :** 08:35

**Extraction End Date :** 10/03/2025

**Extraction End Time :** 13:45

**Concentration By:** EH

**Supervisor By :** ritesh

**Extraction Method:** ☒ Separatory Funnel ☐ Continuous Liquid/Liquid ☐ Sonication ☐ Waste Dilution ☐ Soxhlet

| Standard Name | MLS USED | Concentration ug/mL | STD REF. # FROM LOG |
|---------------|----------|---------------------|---------------------|
| Spike Sol 1   | 1.0ML    | 50/100 PPM          | SP6865              |
| Surrogate     | 1.0ML    | 100/150 PPM         | SP6869              |
| N/A           | N/A      | N/A                 | N/A                 |
| N/A           | N/A      | N/A                 | N/A                 |
| N/A           | N/A      | N/A                 | N/A                 |

| Chemical Used      | ML/SAMPLE USED | Lot Number |
|--------------------|----------------|------------|
| Methylene Chloride | N/A            | E3973      |
| Baked Na2SO4       | N/A            | EP2646     |
| H2SO4 1:1          | N/A            | EP2610     |
| 10N NaOH           | N/A            | EP2609     |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |
| N/A                | N/A            | N/A        |

**Extraction Conformance/Non-Conformance Comments:**

pH Adjusted to < 2 with 1:1 H2SO4 & >12 with 10N NaOH, 1.5ML Vial Lot # 2210443.

**KD Bath ID:** WATER BATH-1,2

**KD Bath Temperature:** 60 °C

**Envap ID:** NE VAP-02

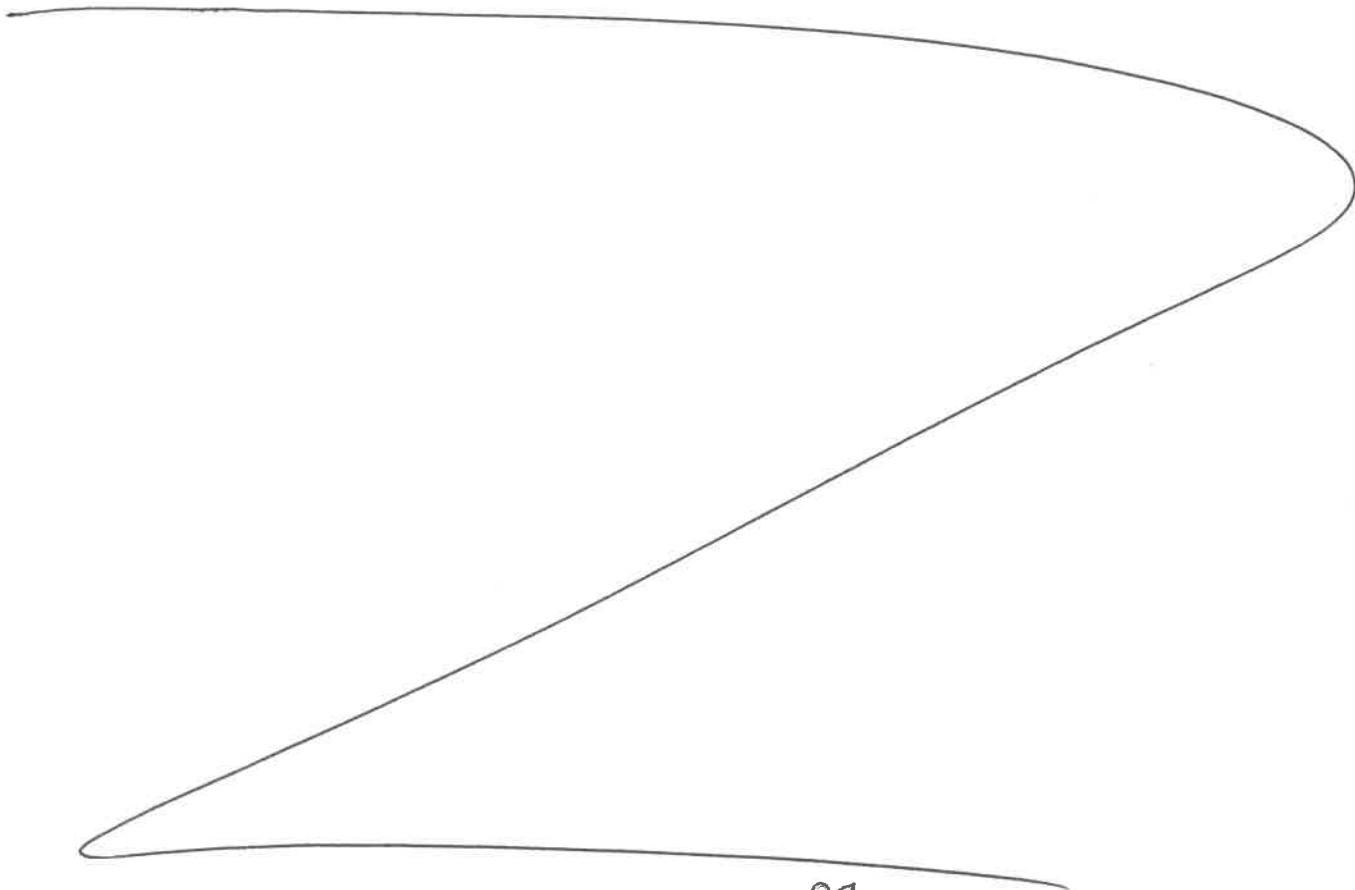
**Envap Temperature:** 40 °C

| Date / Time | Prepped Sample Relinquished By/Location | Received By/Location |
|-------------|-----------------------------------------|----------------------|
| 10/03/25    | RJ CEXT-1067                            | Rclsvoc              |
| 13:50       | Preparation Group                       | Analysis Group       |

Analytical Method: M3510C,3580A-Extraction SVOC-21

Concentration Date: 10/03/2025

| Sample ID       | Client Sample ID | Test                | g / mL | PH | Surr/Spike By: |            | Final Vol.<br>(mL) | JarID | Comments | Prep<br>Pos |
|-----------------|------------------|---------------------|--------|----|----------------|------------|--------------------|-------|----------|-------------|
|                 |                  |                     |        |    | AddedBy        | VerifiedBy |                    |       |          |             |
| PB169973BL      | SBLK973          | SVOC-TCL BNA<br>-20 | 1000   | 6  | ritesh         | Evelyn     | 1                  |       |          | SEP-1       |
| PB169973BS      | SLCS973          | SVOC-TCL BNA<br>-20 | 1000   | 6  | ritesh         | Evelyn     | 1                  |       |          | 2           |
| PB169973BS<br>D | SLCSD973         | SVOC-TCL BNA<br>-20 | 1000   | 6  | ritesh         | Evelyn     | 1                  |       |          | 3           |
| Q3263-01        | 251818           | SVOC-TCL BNA<br>-20 | 1000   | 6  | ritesh         | Evelyn     | 1                  | N     |          | 4           |
| Q3263-02        | 266380           | SVOC-TCL BNA<br>-20 | 1000   | 6  | ritesh         | Evelyn     | 1                  | O     |          | 5           |
| Q3273-01        | MW1              | SVOC-TCL BNA<br>-20 | 1000   | 6  | ritesh         | Evelyn     | 1                  | B     |          | 6           |
| Q3274-01        | MW4              | SVOCMS<br>Group1    | 1000   | 6  | ritesh         | Evelyn     | 1                  | C     |          | 7           |



RJ  
10/03

\* Extracts relinquished on the same date as received.

8:135  
Q3273

WORKLIST(Hardcopy Internal Chain)

WorkList Name : Q3273      WorkList ID : 192268      Department : Extraction      Date : 10-03-2025 08:30:38

| Sample   | Customer Sample | Matrix | Test             | Preservative | Customer | Raw Sample Storage Location | Collect Date | Method |
|----------|-----------------|--------|------------------|--------------|----------|-----------------------------|--------------|--------|
| Q3263-01 | 251818          | Water  | SVOC-TCL BNA -20 | Cool 4 deg C | PSEG03   | D31                         | 10/01/2025   | 8270E  |
| Q3263-02 | 266380          | Water  | SVOC-TCL BNA -20 | Cool 4 deg C | PSEG03   | D31                         | 10/01/2025   | 8270E  |
| Q3273-01 | MW1             | Water  | SVOC-TCL BNA -20 | Cool 4 deg C | GENV01   | D31                         | 10/01/2025   | 8270E  |
| Q3274-01 | MW4             | Water  | SVOCMS Group1    | Cool 4 deg C | GENV01   | D41                         | 10/01/2025   | 8270E  |

Date/Time 10/03/25 8:30  
Raw Sample Received by: R3 (LTA-106)  
Raw Sample Relinquished by: QP Sn

Date/Time 10/03/25 9:00  
Raw Sample Received by: QP Sn  
Raw Sample Relinquished by: R3 (LTA-106)

LAB CHRONICLE

|          |                 |            |                       |
|----------|-----------------|------------|-----------------------|
| OrderID: | Q3273           | OrderDate: | 10/2/2025 3:36:00 PM  |
| Client:  | G Environmental | Project:   | Monroe, NJ            |
| Contact: | Gary Landis     | Location:  | D31,VOA Ref. #3 Water |

| LabID    | ClientID | Matrix | Test             | Method | Sample Date | Prep Date | Anal Date | Received |
|----------|----------|--------|------------------|--------|-------------|-----------|-----------|----------|
| Q3273-01 | MW1      | Water  | SVOC-TCL BNA -20 | 8270E  | 10/01/25    | 10/03/25  | 10/03/25  | 10/02/25 |



# SHIPPING DOCUMENTS

CLIENT INFORMATION

CLIENT PROJECT INFORMATION

CLIENT BILLING INFORMATION

COMPANY: G ENVIRONMENTAL  
ADDRESS: 8 CARRIAGE  
CITY: Succasunna STATE: NJ ZIP: 07876  
ATTENTION:  
PHONE: FAX:

PROJECT NAME: Monroe  
PROJECT NO.: LOCATION: 6 NJ  
PROJECT MANAGER: GL  
e-mail:  
PHONE: FAX:

BILL TO: G ENVIRONMENTAL  
ADDRESS: 8 CARRIAGE  
CITY: Succasunna STATE: NJ ZIP: 07876  
ATTENTION: PHONE:  
ANALYSIS

DATA TURNAROUND INFORMATION

DATA DELIVERABLE INFORMATION

FAX (RUSH) STANDARD DAYS\*  
HARDCOPY (DATA PACKAGE): STANDARD DAYS\*  
EDD: STANDARD DAYS\*  
\*TO BE APPROVED BY CHEMTECH  
STANDARD HARDCOPY TURNAROUND TIME IS 10 BUSINESS

☐ Level 1 (Results Only) ☐ Level 4 (QC + Full Raw Data)  
☐ Level 2 (Results + QC) ☒ NJ Reduced ☐ US EPA CLP  
☐ Level 3 (Results + QC) ☐ NYS ASP A ☐ NYS ASP B  
+ Raw Data ☐ Other  
☒ EDD FORMAT excel, pdf

1. 2. 3. 4. 5. 6. 7. 8. 9. BN TICS

| ALLIANCE<br>SAMPLE<br>ID | PROJECT<br>SAMPLE IDENTIFICATION | SAMPLE<br>MATRIX | SAMPLE<br>TYPE |          | SAMPLE<br>COLLECTION |          | # OF BOTTLES | PRESERVATIVES |   |   |   |   |   |   |   |   | COMMENTS                                                                   |
|--------------------------|----------------------------------|------------------|----------------|----------|----------------------|----------|--------------|---------------|---|---|---|---|---|---|---|---|----------------------------------------------------------------------------|
|                          |                                  |                  | COMP           | GRAB     | DATE                 | TIME     |              | 1             | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 |                                                                            |
| 1.                       | <u>MW1</u>                       | <u>GW</u>        |                | <u>X</u> | <u>10/25/24</u>      | <u>2</u> | <u>2</u>     | <u>X</u>      |   |   |   |   |   |   |   |   | ← Specify Preservatives<br>A-HCl D-NaOH<br>B-HNO3 E-ICE<br>C-H2SO4 F-OTHER |
| 2.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 3.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 4.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 5.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 6.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 7.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 8.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 9.                       |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |
| 10.                      |                                  |                  |                |          |                      |          |              |               |   |   |   |   |   |   |   |   |                                                                            |

SAMPLE CUSTODY MUST BE DOCUMENTED BELOW EACH TIME SAMPLES CHANGE POSSESSION INCLUDING COURIER DELIVERY

|                                                   |                         |                        |                                                                                                                                                                           |
|---------------------------------------------------|-------------------------|------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| RELINQUISHED BY SAMPLER:<br>1. <u>[Signature]</u> | DATE/TIME: <u>10/25</u> | RECEIVED BY: <u>CP</u> | Conditions of bottles or coolers at receipt: <input type="checkbox"/> COMPLIANT <input type="checkbox"/> NON COMPLIANT <input type="checkbox"/> COOLER TEMP <u>2-1</u> °C |
| RELINQUISHED BY SAMPLER:<br>2. <u>[Signature]</u> | DATE/TIME:              | RECEIVED BY:           | Comments: <u>BN Tics</u>                                                                                                                                                  |
| RELINQUISHED BY SAMPLER:<br>3. <u>[Signature]</u> | DATE/TIME:              | RECEIVED BY:           | Comments: <u>2-1</u>                                                                                                                                                      |

Page \_\_\_\_ of CLIENT: ☐ Hand Delivered ☐ Other Shipment Complete  
☐ YES ☐ NO

### Laboratory Certification

| Certified By    | License No.      |
|-----------------|------------------|
|                 |                  |
| Connecticut     | PH-0830          |
|                 |                  |
| DOD ELAP (ANAB) | L2219            |
|                 |                  |
| Maine           | 2024021          |
|                 |                  |
| Maryland        | 296              |
|                 |                  |
| New Hampshire   | 255425           |
|                 |                  |
| New Jersey      | 20012            |
|                 |                  |
| New York        | 11376            |
|                 |                  |
| Pennsylvania    | 68-00548         |
|                 |                  |
| Soil Permit     | 525-24-234-08441 |
|                 |                  |
| Texas           | TX-C25-00189     |
|                 |                  |
| Virginia        | 460312           |