



Analytical Resources, LLC
Analytical Chemists and Consultants

14 April 2023

Ali Judkins
Anchor QEA, LLC
1201 3rd Ave, Suite 2600
Seattle, WA 98101

RE: AOC4 UR Phase 3

Please find enclosed sample receipt documentation and analytical results for samples from the project referenced above.

Sample analyses were performed according to ARI's Quality Assurance Plan and any provided project specific Quality Assurance Plan. Each analytical section of this report has been approved and reviewed by an analytical peer, the appropriate Laboratory Supervisor or qualified substitute, and a technical reviewer.

Should you have any questions or problems, please feel free to contact us at your convenience.

Associated Work Order(s)
23C0107

Associated SDG ID(s)
N/A

I certify that this data package is in compliance with the terms and conditions of the contract, both technically and for completeness, for other than the conditions detailed in the enclosed Narrative. ARI, an accredited laboratory, certifies that the report results for which ARI is accredited meets all the requirements of the accrediting body. A list of certified analyses, accreditations, and expiration dates is included in this report.

Release of the data contained in this hardcopy data package has been authorized by the Laboratory Manager or his/her designee, as verified by the following signature.

Analytical Resources, LLC

Susan Dunninghoo, Director, Client Services

The results in this report apply to the samples analyzed in accordance with the chain of custody document. This analytical report must be reproduced in its entirety.



1 of 1

23C0147

CHAIN-OF-CUSTODY/TEST REQUEST FORM

No 3449

Project/Client Name: AOC4 UR Phase 3
 Project Number: 180067.02.04
 Contact Name: Amara Vandevort
 Sampled By: Windward

Ship to: ARL
 Attn: Sue Dunnington
 Shipper: Conner
 Form filled out by: TDJ

Shipping Date: 3/3/23
 Airbill Number: _____
 Turnaround requested: std

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)							Comments / Instructions (Jar tag number(s))	
					WALD AN	PLGS	TOC/TS	Araive					
03.02.23	1625	LDW12-55B26	3	Sediment	X	X	X						Hold pending communication from Windward AW
AW 3/3/23													
Total Number of Containers			3	Purchase Order / Statement of Work # APJ-110222-AOC4-ARL									
1) Released by: Amara Vandevort			1) Rec'd by: <u>Phyllis</u>			2) Released by:			2) Rec'd by:				
Signature: <u>[Signature]</u>			Company: <u>AR</u>			Print name:			Company:				
Company: <u>Windward</u>			Date/Time: <u>3/3/23 1635</u>			Signature:			Date/Time:				
Date/Time: <u>3/3/23 1635</u>			Date/Time: <u>3/3/23 1635</u>			Company:			Date/Time:				

* Distribution: White copies accompany shipment; yellow retained by consignor.



200 1st Ave W, Suite 500
 Seattle, WA 98119
 206.378.1364

To be completed by Laboratory upon sample receipt:

Date of receipt: _____	Laboratory W.O. #: _____
Condition upon receipt: _____	Time of receipt: _____
Cooler temperature: _____	Received by: _____

Archive

1 of 4 23C0107 / 23C0107 CHAIN-OF-CUSTODY/TEST REQUEST FORM

Project/Client Name: Project ACCU
Project Number: 180007-0222
Contact Name: Amos Thompson
Sampled By: Wardward

Ship to: ARI
Attn: Sam Richardson
Shippers: Cowling
Form filled out by: Nickie Edwards
Shipping Date: 7/13/21
Airbill Number: -
Turnaround requested: STB

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)						Comments / Instructions [bar tag number(s)]
					Archive						
7/13/2021	0659	LAW21-IT 608 B	3	Sediment	X						
		LAW21-IT 608 C	3		X						
		LAW21-IT 608 D	3		X						
		LAW21-IT 608 E	3		X						
	0659	LAW21-IT 608 F	3		X						
	0659	LAW21-IT 608 G	3		X						
	0740	LAW21-IT 602 A	4		X						
		LAW21-IT 602 B	4		X						
		LAW21-IT 602 F	4		X						
	0742	LAW21-IT 602 G	4		X						
	0817	LAW21-IT 609 A	4		X						
		LAW21-IT 609 D	4	Sediment	X						

7/13/2021 0817 Total Number of Containers: 42 Purchase Order / Statement of Work # CLF-052021-ARI

1) Released by: Print name: <u>A. Wardward</u> Signature: <u>[Signature]</u> Company: <u>Wardward</u> Date/Time: <u>7/13/21 1724</u>	1) Rec'd by: Print name: <u>[Signature]</u> Signature: <u>[Signature]</u> Company: <u>ARI</u> Date/Time: <u>07/13/21 1724</u>	2) Released by: Print name: Signature: Company: Date/Time:	2) Rec'd by: Print name: Signature: Company: Date/Time:
--	---	--	---

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To be completed by Laboratory upon sample receipt:

Date of receipt:	Laboratory W.O. #:
Condition upon receipt:	Time of receipt:
Cooler temperature:	Received by:

2 of 4 202209 / 23C0107
CHAIN-OF-CUSTODY/TEST REQUEST FORM

Project/Client Name: Dunsmuir ACCY
 Project Number: 18027-0202
 Contact Name: Anna Vandervort
 Sampled by: Windward

Ship to: ACI
 Attn: Sue Dunsmuir Shipping Date: 7/13/21
 Shipper: Lowry Airbill Number: _____
 Form filled out by: Micaela Falkner Turnaround requested: STP

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check tests required)						Comments / Instructions (for tag numbers)
					Asbestos						
7/13/2021	0817	LW021-IT 659 F	4	Sediment	X						
	↓	LW021-IT 659 G	4		X						
	0817	LW021-IT 659 G-651	1		X						
	0901	LW021-IT 658 A	4		X						
	↓	LW021-IT 658 D	4		X						
	↓	LW021-IT 658 F	4		X						
	0901	LW021-IT 658 G	4		X						
	1020	LW021-IT 657 A	4		X						
	↓	LW021-IT 657 D	4		X						
	↓	LW021-IT 657 E	4		X						
	↓	LW021-IT 657 F	4		X						
7/13/2021	1020	LW021-IT 657-651	1	Sediment	X						
Total Number of Containers			42	Purchase Order / Statement of Work # CLF-052021-ARI							

1) Released by: <u>A. Vandervort</u>	1) Rec'd by: <u>Micaela Falkner</u>	2) Released by:	2) Rec'd by:
Print name: <u>A. Vandervort</u>	Print name: <u>Micaela Falkner</u>	Print name:	Print name:
Signature: <u>[Signature]</u>	Signature: <u>[Signature]</u>	Signature:	Signature:
Company: <u>Windward</u>	Company: <u>ACI</u>	Company:	Company:
Date/Time: <u>7/13/21 1424</u>	Date/Time: <u>07/13/21 1324</u>	Date/Time:	Date/Time:

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To be completed by Laboratory upon sample receipt:

Date of receipt:	Laboratory W.O. #:
Condition upon receipt:	Time of receipt:
Cooler temperature:	Received by:

3 of 4 230229/23cd147
CHAIN-OF-CUSTODY/TEST REQUEST FORM

ARL 3708

Project/Client Name: DuPont AOC4
 Project Number: 170017-0200
 Contact Name: Amanda Woodworth
 Sampled By: Ward

Ship to: ARI
 Attn: So. Dunbar
 Shipper: Quint
 Form filled out by: Ward
 Shipping Date: 7/13/21
 Airbill Number: ---
 Turnaround requested: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Tests Requested (check tests required)						Comments / Instructions (for tag number(s))
					Asph						
7/13/2021	1020	LDW21-IT657-GS2	1	Sediment	X						
	1136	LDW21-IT648A	4		X						
		LDW21-IT648D	4		X						
		LDW21-IT648E	4		X						
		LDW21-IT648F	4		X						
		LDW21-IT648-GS1	1		X						
	1138	LDW21-IT648-GS2	1		X						
	1216	LDW21-IT630B	TR4 3		X						
		LDW21-IT630C	TR4 3		X						
		LDW21-IT630D	TR4 3		X						
		LDW21-IT630E	TR4 3		X						
7/13/2021	1216	LDW21-IT630F	TR4 3	Sediment	X						
Total Number of Containers			<u>24</u>	Purchase Order / Statement of Work # <u>CLF-052021-ARI</u>							
1: Released by:			1: Rec'd by:			2: Released by:			2: Rec'd by:		
Print name: <u>A Woodworth</u>			Print name: <u>Ward</u>			Print name:			Print name:		
Signature: <u>A Woodworth</u>			Signature: <u>Ward</u>			Signature:			Signature:		
Company: <u>Ward</u>			Company: <u>ARI</u>			Company:			Company:		
Date/Time: <u>7/13/21 1724</u>			Date/Time: <u>07/13/21 1724</u>			Date/Time:			Date/Time:		

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To be completed by Laboratory upon sample receipt:

Date of receipt:	Laboratory W.O. #:
Condition upon receipt:	Time of receipt:
Cooler temperature:	Received by:



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4 of 4 23379/23cd1d7

CHAIN-OF-CUSTODY/TEST REQUEST FORM

Analysis 12 3850

Project/Client Name: Demingville AOC9
 Project Number: 180007-0202
 Contact Name: Amara Vanderjart
 Sampled By: Ward

Ship to: AKI
 Attn: See Demingville
 Shipper: Enviro
 Form filled out by: Amara Vanderjart
 Shipping Date: 7/13/21
 Airbill Number: -
 Turnaround requested: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)						Comments / Instructions (jar tag numbers)
					Analyte						
7/13/2021	1307	LDWZ1-SC596A	3	Sediment	X						
		LDWZ1-SC596B	3		X						
		LDWZ1-SC596C	3		X						
		LDWZ1-SC596D	3		X						
		LDWZ1-SC596E	3		X						
	1307	LDWZ1-SC596F	3		X						
	1359	LDWZ1-SC562A	3		X						
		LDWZ1-SC562B	3		X						
		LDWZ1-SC562C	3		X						
	1359	LDWZ1-SC562D	3		X						

Total Number of Containers: 30 Purchase Order / Statement of Work #

1) Released by: <u>Amara Vanderjart</u>	1) Rec'd by: <u>Amara Vanderjart</u>	2) Released by:	2) Rec'd by:
Print name: <u>Amara Vanderjart</u>	Print name: <u>Amara Vanderjart</u>	Print name:	Print name:
Signature: <u>[Signature]</u>	Signature: <u>[Signature]</u>	Signature:	Signature:
Company: <u>Ward Environmental LLC</u>	Company: <u>AKI</u>	Company:	Company:
Date/Time: <u>7/13/21 1724</u>	Date/Time: <u>7/13/21 1724</u>	Date/Time:	Date/Time:

* Distribution: White copies accompany shipment; yellow retained by consignee.

To be completed by Laboratory upon sample receipt:

Date of receipt:	Laboratory W.O. #:
Condition upon receipt:	Time of receipt:
Cooler temperature:	Received by:



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1 of 4 2160229 **CHAIN-OF-CUSTODY/TEST REQUEST FORM**

Archive

Project/Client Name: Duwanish AOC4
 Project Number: 180067-02.02
 Contact Name: Amara Vandervoort
 Sampled By: Windward

Ship to: ARI
 Attn: Sue Dunningham Shipping Date: 7/13/21
 Shipper: Courier Airbill Number: _____
 Form filled out by: Nicolas Eckhardt Turnaround requested: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)						Comments / Instructions [Jar tag number(s)]
					Archive						
7/13/2021	0659	LDW21-IT 608 B	3	Sediment	X						
		LDW21-IT 608 C	3		X						
		LDW21-IT 608 D	3		X						
		LDW21-IT 608 E	3		X						
	0659	LDW21-IT 608 F	3		X						
	0659	LDW21-IT 608 G	3		X						
	0742	LDW21-IT 662 A	4		X						
		LDW21-IT 662 D	4		X						
		LDW21-IT 662 F	4		X						
	0742	LDW21-IT 662 G	4		X						
	0817	LDW21-IT 659 A	4		X						
		LDW21-IT 659 D	4	Sediment	X						

7/13/2021 0817 **Total Number of Containers** 42 Purchase Order / Statement of Work # CLF-052021-ARI

1) Released by: <u>Amara Vandervoort</u> Print name: <u>Amara Vandervoort</u> Signature: <u>[Signature]</u> Company: <u>Windward</u> Date/Time: <u>7/13/21 1724</u>	1) Rec'd by: <u>Nicolas Eckhardt</u> Print name: <u>Nicolas Eckhardt</u> Signature: <u>[Signature]</u> Company: <u>ARI</u> Date/Time: <u>07/13/2021 1724</u>	2) Released by: _____ Print name: _____ Signature: _____ Company: _____ Date/Time: _____	2) Rec'd by: _____ Print name: _____ Signature: _____ Company: _____ Date/Time: _____
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To be completed by Laboratory upon sample receipt:

Date of receipt: _____	Laboratory W.O. #: _____
Condition upon receipt: _____	Time of receipt: _____
Cooler temperature: _____	Received by: _____

2 of 4 2160009 **CHAIN-OF-CUSTODY/TEST REQUEST FORM**

Archive

No 3204

Project/Client Name: Duwamish AOCY
 Project Number: 180067-02.02
 Contact Name: Amara Vandervort
 Sampled By: Windward

Ship to: ARI
 Attn: Sue Dunnihoo Shipping Date: 7/13/21
 Shipper: Courier Airbill Number: —
 Form filled out by: Nicolas Eckhardt Turnaround requested: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)						Comments / Instructions [Jar tag number(s)]
					Archive						
7/13/2021	0817	LDW21-IT 659 F	4	Sediment	X						
	↓	LDW21-IT 659 G	4		X						
	0817	LDW21-IT 659 G-6S1	1		X						
	0901	LDW21-IT 658 A	4		X						
	↓	LDW21-IT 658 D	4		X						
	↓	LDW21-IT 658 F	4		X						
	0901	LDW21-IT 658 G	4		X						
	1020	LDW21-IT 657 A	4		X						
	↓	LDW21-IT 657 D	4		X						
	↓	LDW21-IT 657 E	4		X						
	↓	LDW21-IT 657 F	4		X						
7/13/2021	1020	LDW21-IT 657-6S1	1	Sediment	X						
Total Number of Containers			<u>42</u>	Purchase Order / Statement of Work # <u>CLF-052021-ARI</u>							
1) Released by: <u>A. Vandervort</u> Print name: <u>A. Vandervort</u> Signature: <u>[Signature]</u> Company: <u>Windward</u> Date/Time: <u>7/13/21 1424</u>			1) Rec'd by: <u>[Signature]</u> <u>Jacob Walter</u> Company: <u>ARI</u> Date/Time: <u>07/13/2021 1724</u>			2) Released by: _____ Print name: _____ Signature: _____ Company: _____ Date/Time: _____			2) Rec'd by: _____ Print name: _____ Signature: _____ Company: _____ Date/Time: _____		

* Distribution: White copies accompany shipment; yellow retained by consignor.



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To be completed by Laboratory upon sample receipt:

Date of receipt: _____	Laboratory W.O. #: _____
Condition upon receipt: _____	Time of receipt: _____
Cooler temperature: _____	Received by: _____

3 of 4 2160229

CHAIN-OF-CUSTODY/TEST REQUEST FORM

Archive

No 3206

Project/Client Name: Duwamish AOCY
 Project Number: 180067-02.02
 Contact Name: Amara Vanderwort
 Sampled By: Windward

Ship to: ARI
 Attn: Sue Punniho Shipping Date: 7/13/2021
 Shipper: Courier Airbill Number: ---
 Form filled out by: Nicolas Eckhardt Turnaround requested: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)						Comments / Instructions [Jar tag number(s)]
					Archive						
7/13/2021	1020	LDW21-IT657-GS2	1	Sediment	X						
	1132	LDW21-IT648A	4		X						
		LDW21-IT648D	4		X						
		LDW21-IT648E	4		X						
		LDW21-IT648F	4		X						
		LDW21-IT648-GS1	1		X						
	1132	LDW21-IT648-GS2	1		X						
	1216	LDW21-IT630B	4 3		X						
		LDW21-IT630C	4 3		X						
		LDW21-IT630D	4 3		X						
		LDW21-IT630E	4 3		X						
7/13/2021	1216	LDW21-IT630F	4 3	Sediment	X						
Total Number of Containers			24	Purchase Order / Statement of Work # <u>CLF-052021-ARI</u>							
1) Released by:		1) Rec'd by:		2) Released by:		2) Rec'd by:					
Print name: <u>Amara Vanderwort</u>		Print name: <u>Jacob Walter</u>		Print name:		Print name:					
Signature: <u>[Signature]</u>		Signature: <u>[Signature]</u>		Signature:		Signature:					
Company: <u>Windward</u>		Company: <u>ARI</u>		Company:		Company:					
Date/Time: <u>7/13/21 1724</u>		Date/Time: <u>07/13/2021 1724</u>		Date/Time:		Date/Time:					

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To be completed by Laboratory upon sample receipt:

Date of receipt::	Laboratory W.O. #:
Condition upon receipt:	Time of receipt:
Cooler temperature:	Received by:

4 of 4 260229

CHAIN-OF-CUSTODY/TEST REQUEST FORM

Archive

No 3960

Project/Client Name: Duwamish AOCY
 Project Number: 180067-02.02
 Contact Name: Amara Vanderwort
 Sampled By: Windward

Ship to: ARI
 Attn: Sue Dunning Shipping Date: 7/13/21
 Shipper: Courier Airbill Number: —
 Form filled out by: Nicolas Eckhardt Turnaround requested: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Test(s) Requested (check test(s) required)						Comments / Instructions [Jar tag number(s)]
					Archive						
7/13/2021	1307	LDW21-SC596A	3	Sediment	X						
	↓	LDW21-SC596B	3		X						
		LDW21-SC596C	3		X						
		LDW21-SC596D	3		X						
	↓	LDW21-SC596E	3		X						
	1307	LDW21-SC596F	3		X						
	1359	LDW21-SC562A	3		X						
	↓	LDW21-SC562B	3		X						
		LDW21-SC562C	3		X						
	1359	LDW21-SC562D	3		X						
Total Number of Containers			30	Purchase Order / Statement of Work #							
1) Released by: <u>Amara Vanderwort</u>			1) Rec'd by: <u>Jacob Lehto</u>			2) Released by:			2) Rec'd by:		
Print name: <u>Amara Vanderwort</u>			Company: <u>AW</u>			Print name:			Company:		
Signature: <u>[Signature]</u>			Date/Time: <u>07/13/2021 1724</u>			Signature:			Date/Time:		
Company: <u>Windward</u>						Company:			Date/Time:		
Date/Time: <u>7/13/21 1724</u>						Date/Time:			Date/Time:		

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To be completed by Laboratory upon sample receipt:

Date of receipt:	Laboratory W.O. #:
Condition upon receipt:	Time of receipt:
Cooler temperature:	Received by:

AOC4 samples received on 7/15/21

Brandi Quinlisk <brandiq@windwardenv.com>

Sun 7/18/2021 12:56 PM

2160009

To: Sue Dunning <lmsadm@arilabs.com>

Cc: Amara Vandervort <amarav@windwardenv.com>; Cindy Fields <cfields@anchorqea.com>

Hi Sue,

The three samples shown below were sent to you on July 15th for analysis. Can you please put all of the samples noted below on hold and **do not analyze** at this time.

- LDW21-SC621B
- LDW21-SC621C
- LDW21-SC621E

1 of 2

CHAIN-OF-CUSTODY/TEST REQUEST FORM

3157

Project/Client Name: Downtown AOC4 Ship to: ARI

Project Number: 180067-02.02 ARI: Sue Dunning Shipping Date: 7/15/21

Contact Name: Amara Vandervort Shipper: Curator Airbill Number: _____

Sampled By: Windward Form filled out by: Nicola's Eckhardt Turnaround request: STD

Sample Collection Date (m/d/y)	Time	Sample Identification	Volume of Sample / # of Containers	Matrix	Tests Requested (check test(s) required)						Comments / Instructions (for tag numbers)
					PCBs	PAHs / PCBs / SCOPs	Aroclor	Mercury	PCBs / PBB	Lead / Cu	
7/17/2021											
0942		LDW21-SC621 B	3		X	X	X				
↓		LDW21-SC621 C	3		X	X	X				
0942		LDW21-SC621 E	3		X	X	X				

Also, can you please revise these two sample IDs as follows:

- Received 7/12: LDW21-IT652F-GS1, please revise to LDW21-IT652-GS1
- Received 7/13: LDW21-IT659G-GS1, please revise to LDW21-IT659-GS1

Thank you so much,
Brandi

Brandi Quinlisk
Database Manager

200 1st Avenue W., Suite 500 | Seattle, WA 98119
Direct: 206-812-5408
brandiq@windwardenv.com | www.windwardenv.com



RE: Six receipts to review

Amara Vandervort <amarav@windwardenv.com>

Mon 3/6/2023 8:53 PM

To: Sue Dunnihoo <lmsadm@arilabs.com>; Labdata <labdata@anchorqea.com>

Cc: Ali Judkins <ajudkins@anchorqea.com>; Anastasia Barr <anastasiab@windwardenv.com>

Hi Sue,

Per our conversation on Friday, please add the following two samples to the AOC4 WO 23C0107

Sample ID	Date Collected	ARI Login WO ID	PCBs	TOC/Percent Solids
LDW21-IT608C	7/13/2021	21G0229-02	x	x
LDW21-IT608D	7/13/2021	21G0229-03	x	x

Thanks.

Amara Vandervort

Associate

Direct line: 206-812-5415

E-mail: amarav@windwardenv.com | www.windwardenv.com

200 First Avenue West, Suite 500 | Seattle, WA 98119



From: Sue Dunnihoo <lmsadm@arilabs.com>

Sent: Monday, March 6, 2023 10:09 AM

To: Labdata <labdata@anchorqea.com>

Cc: Ali Judkins <ajudkins@anchorqea.com>; Amara Vandervort <amarav@windwardenv.com>; Anastasia Barr <anastasiab@windwardenv.com>

Subject: Six receipts to review

CAUTION: This email originated from outside of the organization. Do not click links or open attachments unless you recognize the sender and know the content is safe.

Please find attached.

Regards, Sue

I will be out of the office March 8th and working part-time and very remotely March 9-24.

**** NOTE - TATs are running 3 to 6 weeks depending on analysis ****

Courier requests cannot be guaranteed for same day pick-ups or deliveries. Please schedule in advance.

****Please email you bottle requests to your project manager several days before needed to give staff time to assemble the kit - we may not be able to accommodate your request as a walk-in****

Susan D. Dunnihoo

She/her/hers

Client Services Director

Analytical Resources, LLC

Analytical Chemists and Consultants

4611 South 134th Place, Suite 100

Tukwila, WA 98168

(206) 695-6207 office

sue.dunnihoo@arilabs.com

www.arilabs.com

As some of our staff are still working remotely we have changed sample receiving hours to 8 to 5 until further notice. Thank you for understanding.

Versions have been updated for SW-846 8260 and 8270 analyses, see our [post](#) explaining the details.

"Don't wish it were easier. Wish you were better." - Jim Rohn

*****Before printing, think about ENVIRONMENTAL responsibility*****

How was your customer experience?

Please take our 5 minute [Online Customer Survey](#)



Cooler Receipt Form

ARI Client: Anchar Kundnand
 COC No(s): 3449 NA
 Assigned ARI Job No: 23C0107 JS
23/07/23

Project Name: AOC4 UR Phase 3
 Delivered by: Fed-Ex UPS Courier Hand Delivered Other: _____
 Tracking No: _____ NA

Preliminary Examination Phase:

Were intact, properly signed and dated custody seals attached to the outside of the cooler? YES NO
 Were custody papers included with the cooler? YES NO
 Were custody papers properly filled out (ink, signed, etc.) YES NO
 Temperature of Cooler(s) (°C) (recommended 2.0-6.0 °C for chemistry)

Time 1650 1.8 0.4 _____
 If cooler temperature is out of compliance fill out form 00070F Temp Gun ID#: Jca 9718
 Cooler Accepted by: JS for PI Date: 03/07/23 Time: 1635

Complete custody forms and attach all shipping documents

Log-In Phase:

Was a temperature blank included in the cooler? YES NO
 What kind of packing material was used? ... Bubble Wrap Wet Ice Gel Packs Baggies Foam Block Paper Other: _____
 Was sufficient ice used (if appropriate)? NA YES NO
 How were bottles sealed in plastic bags? Individually Grouped Not
 Did all bottles arrive in good condition (unbroken)? YES NO
 Were all bottle labels complete and legible? YES NO
 Did the number of containers listed on COC match with the number of containers received? YES NO
 Did all bottle labels and tags agree with custody papers? YES NO
 Were all bottles used correct for the requested analyses? YES NO
 Do any of the analyses (bottles) require preservation? (attach preservation sheet, excluding VOCs) ... NA YES NO
 Were all VOC vials free of air bubbles? NA YES NO
 Was sufficient amount of sample sent in each bottle? YES NO
 Date VOC Trip Blank was made at ARI NA _____
 Were the sample(s) split by ARI? NA YES Date/Time: _____ Equipment: _____ Split by: _____

Samples Logged by: JS Date: 03/07/23 Time: _____ Labels checked by: JS

**** Notify Project Manager of discrepancies or concerns ****

Sample ID on Bottle	Sample ID on COC	Sample ID on Bottle	Sample ID on COC

Additional Notes, Discrepancies, & Resolutions:

By: _____ Date: _____



Cooler Receipt Form

ARI Client: Whitford / Archer
 COC No(s): 3203, 3204, 3206, 3268 NA
 Assigned ARI Job No: 216-0229

Project Name: Lower Devonian Sh AOC4
 Delivered by: Fed-Ex UPS Courier Hand Delivered Other: _____
 Tracking No: _____ NA

Preliminary Examination Phase:

Were intact, properly signed and dated custody seals attached to the outside of the cooler? YES NO
 Were custody papers included with the cooler? YES NO
 Were custody papers properly filled out (ink, signed, etc.) YES NO
 Temperature of Cooler(s) (°C) (recommended 2.0-6.0 °C for chemistry)

Time 1757 5:11 1:3 4:1 0:9
 If cooler temperature is out of compliance fill out form 00070F Temp Gun ID#: DOO-5206

Cooler Accepted by: TS Date: 07/13/2001 Time: 1725

Complete custody forms and attach all shipping documents

Log-In Phase:

Was a temperature blank included in the cooler? YES NO
 What kind of packing material was used? ... Bubble Wrap Wet Ice Gel Packs Baggies Foam Block Paper Other: _____
 Was sufficient ice used (if appropriate)? NA YES NO
 How were bottles sealed in plastic bags? Individually Grouped Not
 Did all bottles arrive in good condition (unbroken)? YES NO
 Were all bottle labels complete and legible? YES NO
 Did the number of containers listed on COC match with the number of containers received? YES NO
 Did all bottle labels and tags agree with custody papers? YES NO
 Were all bottles used correct for the requested analyses? YES NO
 Do any of the analyses (bottles) require preservation? (attach preservation sheet, excluding VOCs) ... NA YES NO
 Were all VOC vials free of air bubbles? NA YES NO
 Was sufficient amount of sample sent in each bottle? YES NO
 Date VOC Trip Blank was made at ARI: _____ NA
 Were the sample(s) split by ARI? NA YES Date/Time: _____ Equipment: _____ Split by: _____

Samples Logged by: TS/SC Date: 07/20/2001 Time: 1607 Labels checked by: TS/SC

**** Notify Project Manager of discrepancies or concerns ****

Sample ID on Bottle	Sample ID on COC	Sample ID on Bottle	Sample ID on COC

Additional Notes, Discrepancies, & Resolutions:

By: _____ Date: _____



Anchor QEA, LLC

1201 3rd Ave, Suite 2600

Seattle, WA 98101

Project: AOC4 UR Phase 3

Project Number: 180067-02.04

Project Manager: Ali Judkins

Reported:

04/14/2023 15:02

ANALYTICAL REPORT FOR SAMPLES

Laboratory ID	Sample ID	Matrix	Date Sampled	Date Received
23C0107-01	LDW22-SS826	Solid	03/02/23 16:25	03/03/23 16:35
23C0107-02	LDW21-IT608C	Solid	07/13/21 06:54	03/03/23 16:35
23C0107-03	LDW21-IT608D	Solid	07/13/21 06:54	03/03/23 16:35



Anchor QEA, LLC
1201 3rd Ave, Suite 2600
Seattle WA, 98101

Project: AOC4 UR Phase 3
Project Number: 180067-02.04
Project Manager: Ali Judkins

Reported:
14-Apr-2023 15:02

Case Narrative

Client: Anchor QEA, LLC
Project: AOC4 UR Phase 3
Work Order: 23C0107

Sample receipt

One sample as listed on the preceding page was received 03-Mar-2023 16:35 under ARI work order 23C0107. For details regarding sample receipt, please refer to the Cooler Receipt Form.

Two additional samples were pulled from frozen archive, originally logged under ARI work order 21G0229, and logged under ARI work order 23C0107. For details regarding sample receipt, please refer to the original Cooler Receipt Forms.

PCB Aroclors - EPA Method SW8082A

The 2023 sample(s) were prepared and analyzed within the recommended holding times. The 2021 samples were prepared outside recommended holding times and have been "H"-flagged noting the excursion.

Initial and continuing calibrations were within method requirements, with accepted excursions outside the 20% window. Associated positive results have been "Q"-flagged.

Aroclor 1260 failed high in SLC0215-CCV4 and SLC0215-CCV6 on the ZB5 column. Decachlorobiphenyl (DCBP) failed in SLC0215-ICV1 and SLC0215-ICV2 on the ZB5 column. Results have been reported from the ZB35 column as primary.

Hexabromobiphenyl (HBB) failed on the ZB5 column for several extracts, attributed to continued issues with the sample matrix from the project. Results have been reported from the ZB35 column as primary.

The surrogate percent recoveries were within control limits.

The method blank(s) were clean at the reporting limits.

The blank spike and blank spike duplicate (BS/LCS and BSD/LCSD) spike recoveries and relative percent difference (RPD) were within control limits.

The matrix spike/matrix spike duplicate (MS/MSD) percent recoveries and relative percent difference (RPD) were within advisory control limits.

Wet Chemistry

The 2023 sample(s) were prepared and analyzed within the recommended holding times. The 2021 samples were prepared outside recommended holding times and have been "H"-flagged noting the excursion.

Initial and continuing calibrations were within method requirements.

The method blank(s) were clean at the reporting limits.

The blank spike (BS/LCS) percent recoveries were within control limits.

The reference material (SRM) percent recoveries were within control limits.

The batch BLD0117 matrix spike (MS) percent recoveries and the duplicate (DUP) relative percent difference (RPD) were within advisory control limits, reported under work order 23C0071.



QUALIFIERS AND NOTES

<u>Qualifier</u>	<u>Definition</u>
U	This analyte is not detected above the reporting limit (RL) or if noted, not detected above the limit of detection (LOD).
Q	Indicates a detected analyte with an initial or continuing calibration that does not meet established acceptance criteria (<20% RSD, <20% drift or minimum RRF)
HC	The natural concentration of the spiked analyte is so much greater than the concentration spiked that an accurate determination of spike recovery is not possible
H	Hold time violation - Hold time was exceeded.
*	Flagged value is not within established control limits.
DET	Analyte DETECTED
ND	Analyte NOT DETECTED at or above the reporting limit
NR	Not Reported
dry	Sample results reported on a dry weight basis
RPD	Relative Percent Difference



Dual Column

ORGANIC ANALYSIS DATA SHEET
EPA 8082A

Laboratory: <u>Analytical Resources, LLC</u>		SDG: <u>23C0107</u>
Client: <u>Anchor QEA, LLC</u>		
Project: <u>AOC4 UR Phase 3</u>		
Matrix: <u>Solid</u>	Laboratory ID: <u>23C0107-01A</u>	File ID: <u>03152319ECD7.D</u>
Sampled: <u>03/02/23 16:25</u>	Prepared: <u>03/10/23 11:58</u>	Analyzed: <u>03/15/23 19:14</u>
% Solids: <u>41.90</u>	Preparation: <u>EPA 3546 (Microwave)</u>	Initial/Final: <u>29.89 g Wet / 2.5 mL</u>
Batch: <u>BLC0219</u>	Sequence: <u>SLC0215</u>	Calibration: <u>GB00069</u>
Instrument: <u>ECD7</u>	Column 1: <u>ZB5</u>	Column 2: <u>ZB35</u>

CAS NO.	COMPOUND	Col #	DILUTION	CONC. (ug/kg dry)	MDL	MRL	Q
12674-11-2	Aroclor 1016	1	1	4.0	1.6	4.0	U
11104-28-2	Aroclor 1221	1	1	4.0	1.6	4.0	U
11141-16-5	Aroclor 1232	1	1	4.0	1.6	4.0	U
53469-21-9	Aroclor 1242	1	1	4.0	1.6	4.0	U
12672-29-6	Aroclor 1248	1	1	17.1	1.6	4.0	
11097-69-1	Aroclor 1254	2	1	31.4	1.6	4.0	
11096-82-5	Aroclor 1260	2	1	31.5	0.6	4.0	

SURROGATES	Col #	ADDED (ug/kg dry)	CONC (ug/kg dry)	% REC	QC LIMITS	Q
<i>Decachlorobiphenyl</i>	1	7.9847	5.86	73.3	40 - 126	Q
<i>Tetrachlorometaxylene</i>	1	7.9847	4.81	60.2	44 - 120	
<i>Decachlorobiphenyl</i>	2	7.9847	5.85	73.3	40 - 126	
<i>Tetrachlorometaxylene</i>	2	7.9847	5.39	67.5	44 - 120	

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152319ECD7.D
Data file 2: /230315.b/230315.b/03152319ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 23C0107-01
Client ID:
Injection Date: 15-MAR-2023 19:14
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.807	-0.004	363990	5.687	-0.006	147256	24.1	27.0	11.3	Tetrachloro-m-xylene
13.887	-0.010	297931	14.112	-0.009	183388	29.3	29.3	0.0	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1011772	50.2
Hexabromobiphenyl	1429847	1031369	-27.9

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	371913	18.0
Hexabromobiphenyl	513946	410686	-20.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	---			0.0	1	---			0.0	
Aroclor-1016	2	---			0.0	2	---			0.0	
Aroclor-1016	3	---			0.0	3	---			0.0	
Aroclor-1016	4	---			0.0	4	---			0.0	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1221	1	---			0.0	1	---			0.0	
Aroclor-1221	2	---			0.0	2	---			0.0	
Aroclor-1221	3	---			0.0	3	---			0.0	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1232	1	---			0.0	1	---			0.0	
Aroclor-1232	2	---			0.0	2	---			0.0	
Aroclor-1232	3	---			0.0	3	---			0.0	
Aroclor-1232	4	---			0.0	4	---			0.0	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1242	1	---			0.0	1	---			0.0	
Aroclor-1242	2	---			0.0	2	---			0.0	
Aroclor-1242	3	---			0.0	3	---			0.0	
Aroclor-1242	4	---			0.0	4	---			0.0	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1248	1	8.400	-0.011	32388	65.6	1	8.302	-0.013	12602	71.0	
Aroclor-1248	2	8.567	-0.022	27821	44.3	2	8.707	-0.014	11422	62.2	
Aroclor-1248	3	8.986	-0.012	82096	69.3	3	9.142	-0.042	18358	86.9	
Aroclor-1248	4	9.287	-0.014	98724	163.8	4	9.533	-0.079	20728	84.7	
Total CollAve (4 peaks):				85.8	Total Col2Ave (4 peaks):				75.4	RPD = 13	
Corrected Ave (3 peaks):				59.8	Corrected Ave (3 peaks):				71.6	RPD = 18	
73.37											
Aroclor-1254	1	9.287	-0.011	98724	97.2	1	9.440	-0.017	41259	146.0	
Aroclor-1254	2	9.363	-0.014	39406	86.2	2	9.958	-0.021	22099	97.2	
Aroclor-1254	3	9.663	-0.005	98766	151.2	3	10.108	-0.026	70341	143.0	
Aroclor-1254	4	9.789	-0.019	162550	128.0	4	10.355	-0.027	84120	175.4	
Aroclor-1254	5	10.123	-0.053	86005	108.0	5	10.556	-0.022	65414	224.0	
Total CollAve (5 peaks):				114.1	Total Col2Ave (5 peaks):				157.1	RPD = 32	
Corrected Ave (4 peaks):				104.9	Corrected Ave (4 peaks):				140.4	RPD = 29	
115.65											
Aroclor-1260	1	11.033	-0.011	61405	165.5	1	11.643	-0.015	40820	169.0	
Aroclor-1260	2	11.348	-0.013	52080	134.3	2	11.905	-0.020	68071	110.5	
Aroclor-1260	3	11.719	-0.015	133954	130.3	3	12.418	-0.022	38452	235.1	
Aroclor-1260	4	12.119	-0.020	77458	149.6	4	12.488	-0.020	48155	115.9	
Aroclor-1260	5	12.234	-0.010	34500	154.8	NS	---			---	
Total CollAve (5 peaks):				146.9	Total Col2Ave (4 peaks):				157.6	RPD = 7	
Corrected Ave (4 peaks):				142.3	Corrected Ave (3 peaks):				131.8	RPD = 8	
Aroclor-1262	1	---			0.0	1	---			0.0	
Aroclor-1262	2	---			0.0	2	---			0.0	
Aroclor-1262	3	---			0.0	3	---			0.0	
Aroclor-1262	4	---			0.0	4	---			0.0	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1268	1	---			0.0	1	---			0.0	
Aroclor-1268	2	---			0.0	2	---			0.0	
Aroclor-1268	3	---			0.0	3	---			0.0	
Aroclor-1268	4	---			0.0	4	---			0.0	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						

Total PCB Area Col1 (5.911 - 13.797) = 3077079 Col1 Total PCB = 0.3 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 1278223 Col2 Total PCB = 0.3 ppm*

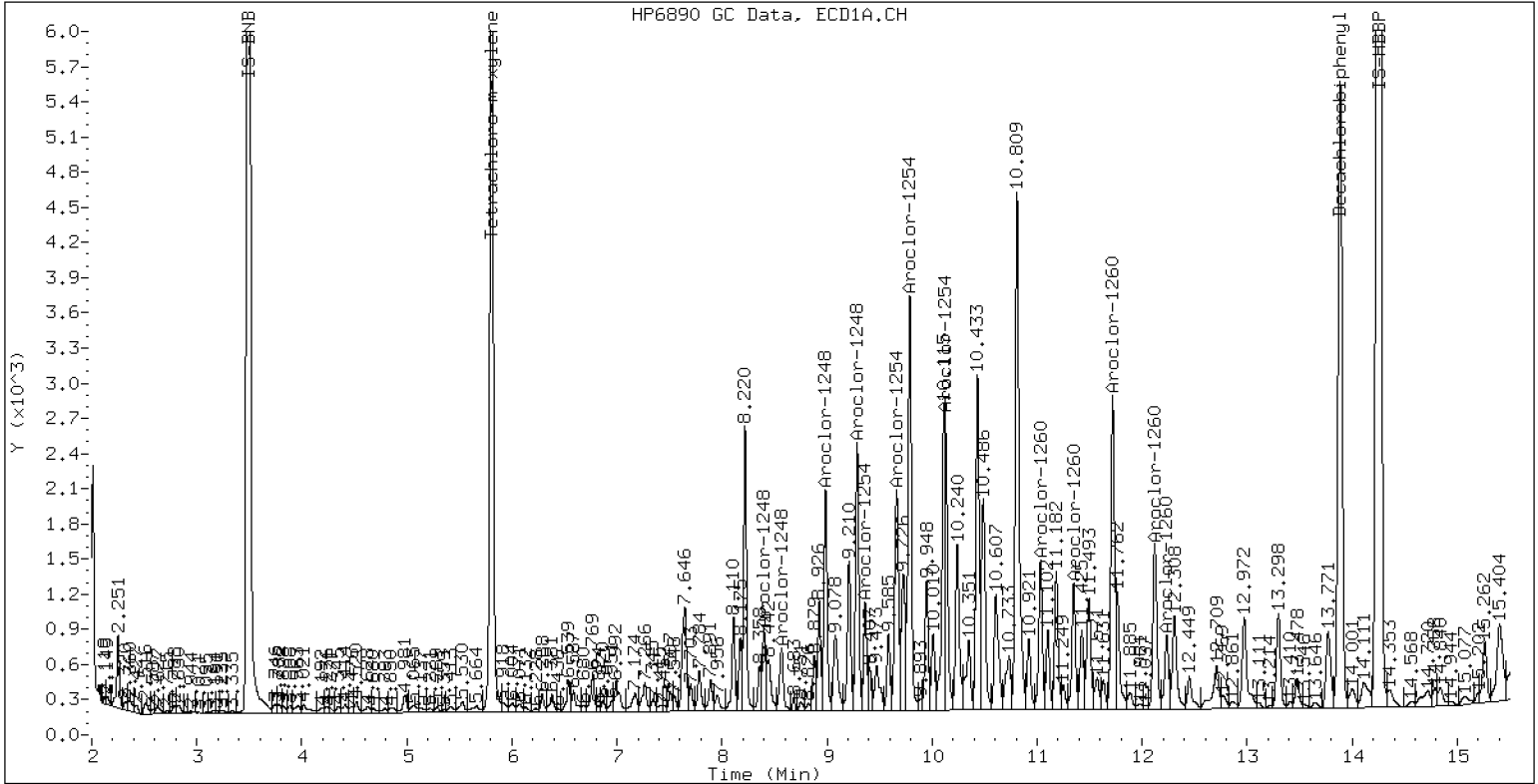
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 23C0107-01

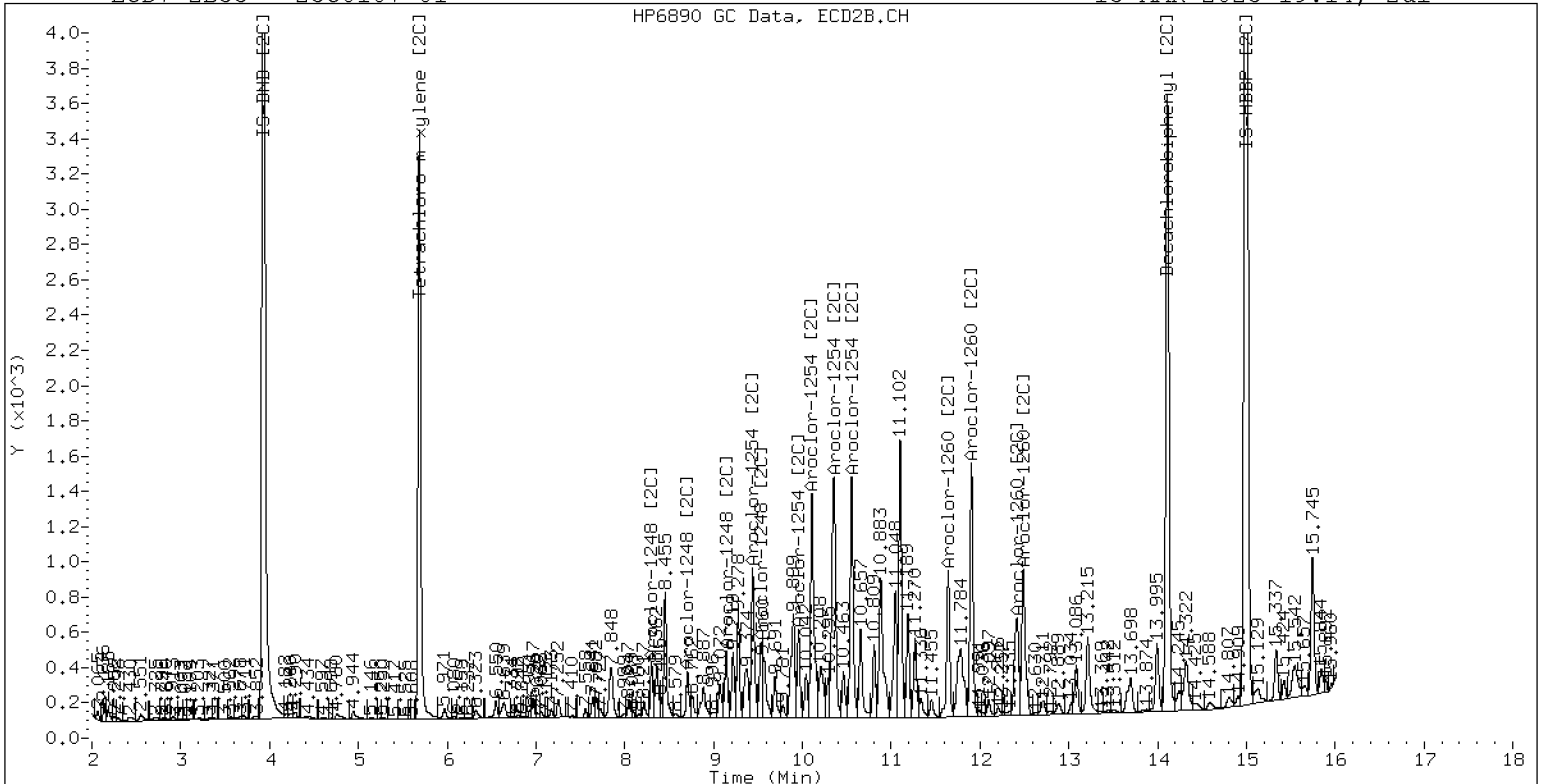
15-MAR-2023 19:14, 2u1



ZB-5 Manual Integration: YES

ECD7-ZB35 23C0107-01

15-MAR-2023 19:14, 2u1



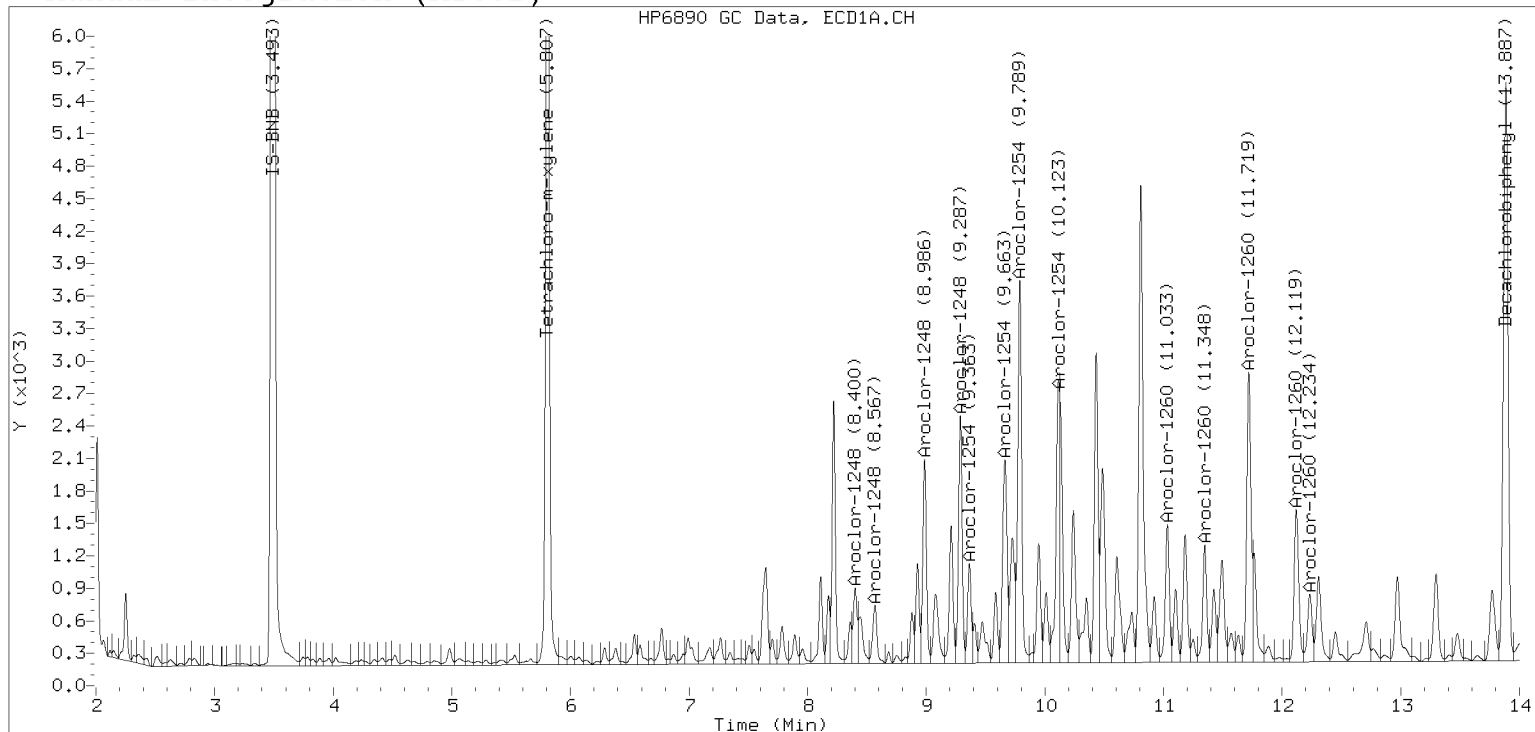
ZB-35 Manual Integration: YES

Manual Peak Adjustment, ZB-5

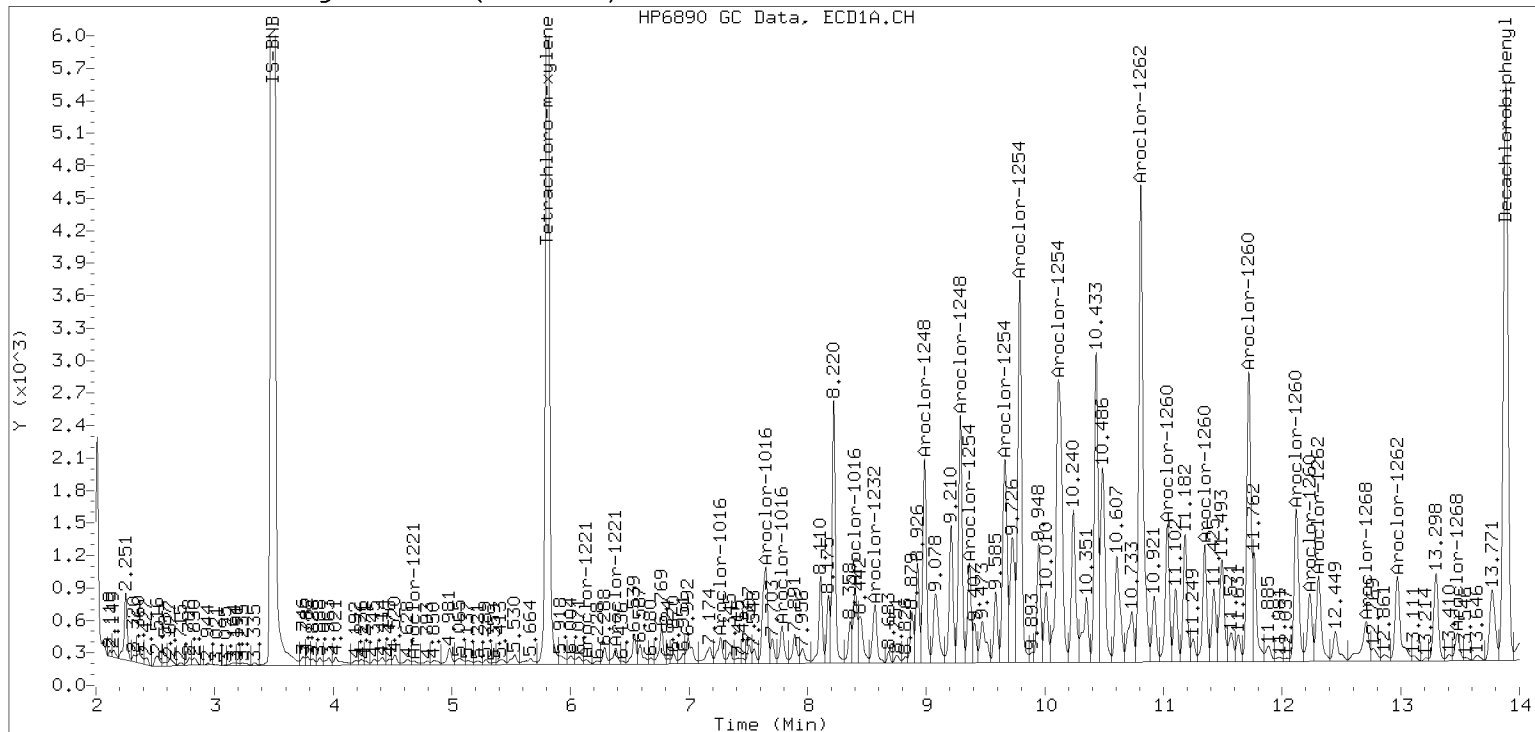
Datafile: ecd7.i/230315.b/03152319ECD7.D

Injection Date: 15-MAR-2023 19:14

Manual Integration (After)



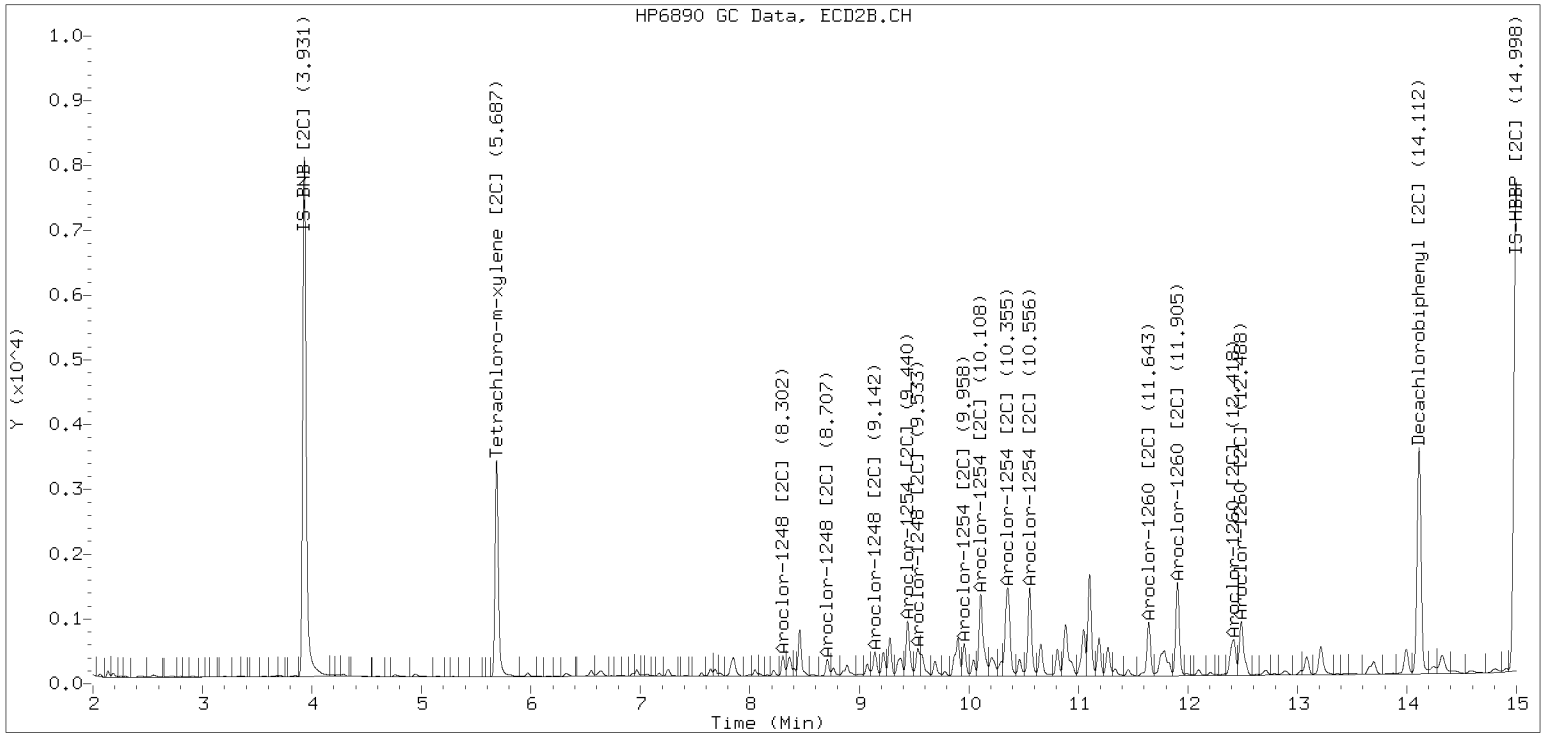
Processed Integration (Before)



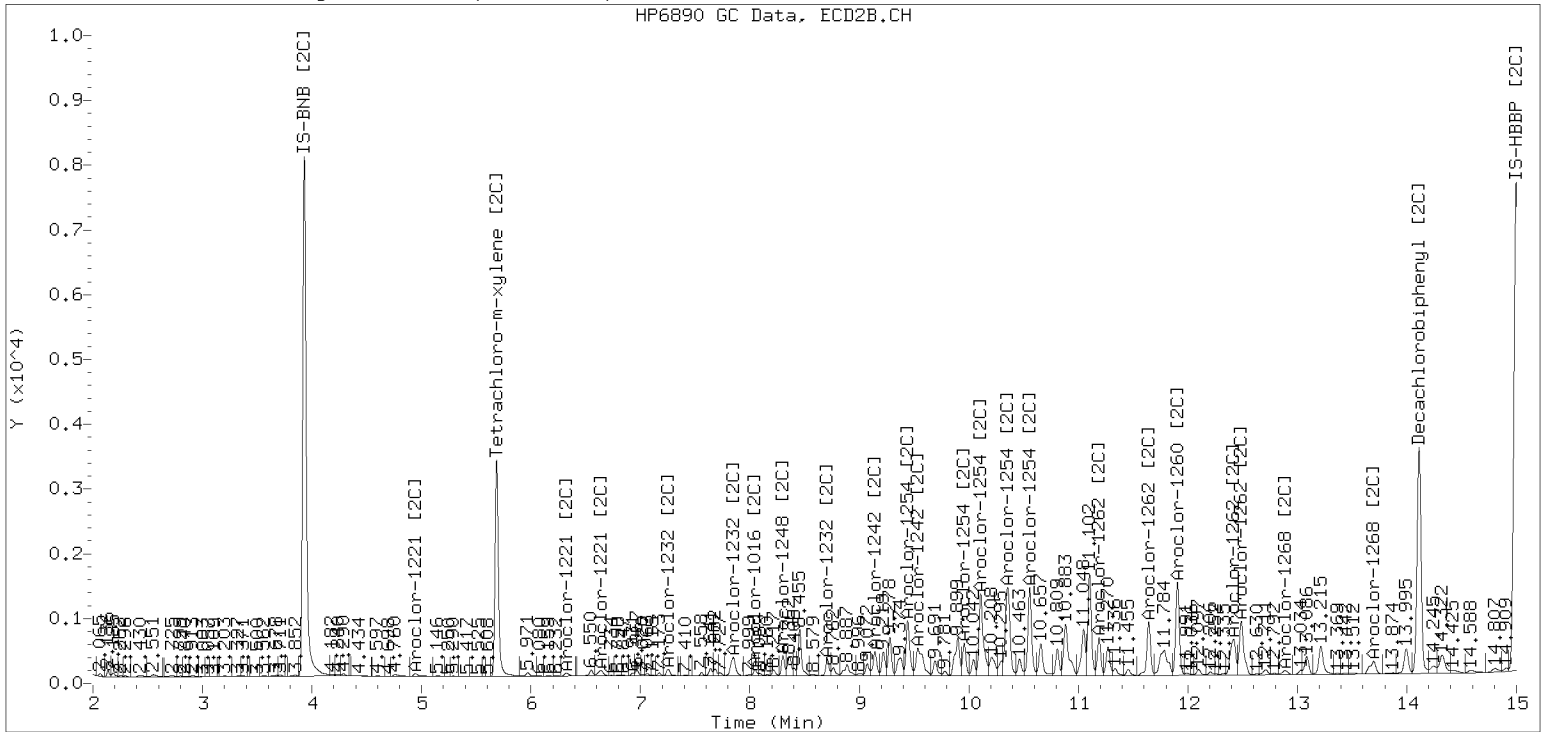
Manual Peak Adjustment, ZB-35

Datafile: ecd7.i/230315.b/230315.b/03152319ECD7.D Injection Date: 15-MAR-2023

Manual Integration (After)



Processed Integration (Before)





Dual Column

ORGANIC ANALYSIS DATA SHEET
EPA 8082A

Laboratory: Analytical Resources, LLC SDG: 23C0107
 Client: Anchor QEA, LLC
 Project: AOC4 UR Phase 3
 Matrix: Solid Laboratory ID: 23C0107-02 A File ID: 03152320ECD7.D
 Sampled: 07/13/21 06:54 Prepared: 03/10/23 11:58 Analyzed: 03/15/23 19:35
 % Solids: 67.19 Preparation: EPA 3546 (Microwave) Initial/Final: 18.61 g Wet / 2.5 mL
 Batch: BLC0219 Sequence: SLC0215 Calibration: GB00069
 Instrument: ECD7 Column 1: ZB5 Column 2: ZB35

CAS NO.	COMPOUND	Col #	DILUTION	CONC. (ug/kg dry)	MDL	MRL	Q
12674-11-2	Aroclor 1016	1	1	4.0	1.6	4.0	H, U
11104-28-2	Aroclor 1221	1	1	4.0	1.6	4.0	H, U
11141-16-5	Aroclor 1232	1	1	4.0	1.6	4.0	H, U
53469-21-9	Aroclor 1242	1	1	4.0	1.6	4.0	H, U
12672-29-6	Aroclor 1248	1	1	4.0	1.6	4.0	H, U
11097-69-1	Aroclor 1254	1	1	4.0	1.6	4.0	H, U
11096-82-5	Aroclor 1260	1	1	4.0	0.6	4.0	H, U

SURROGATES	Col #	ADDED (ug/kg dry)	CONC (ug/kg dry)	% REC	QC LIMITS	Q
<i>Decachlorobiphenyl</i>	1	7.9974	6.83	85.4	40 - 126	Q
<i>Tetrachlorometaxylene</i>	1	7.9974	6.11	76.4	44 - 120	

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152320ECD7.D
Data file 2: /230315.b/230315.b/03152320ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 23C0107-02
Client ID:
Injection Date: 15-MAR-2023 19:35
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	-0.002	463032	5.690	-0.003	170954	30.5	30.1	1.6	Tetrachloro-m-xylene
13.892	-0.005	513008	14.118	-0.003	264110	34.2	35.2	2.9	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1015111	50.7
Hexabromobiphenyl	1429847	1524021	6.6

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	387409	22.9
Hexabromobiphenyl	513946	492712	-4.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	---			0.0	1	---			0.0
Aroclor-1016	2	---			0.0	2	---			0.0
Aroclor-1016	3	---			0.0	3	---			0.0
Aroclor-1016	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	---			0.0	2	---			0.0
Aroclor-1221	3	---			0.0	3	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	---			0.0	2	---			0.0
Aroclor-1232	3	---			0.0	3	---			0.0
Aroclor-1232	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1242	1	---			0.0	1	---			0.0
Aroclor-1242	2	---			0.0	2	---			0.0
Aroclor-1242	3	---			0.0	3	---			0.0
Aroclor-1242	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1248	1	---			0.0	1	---			0.0
Aroclor-1248	2	---			0.0	2	---			0.0
Aroclor-1248	3	---			0.0	3	---			0.0
Aroclor-1248	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1254	1	---			0.0	1	---			0.0
Aroclor-1254	2	---			0.0	2	---			0.0
Aroclor-1254	3	---			0.0	3	---			0.0
Aroclor-1254	4	---			0.0	4	---			0.0
Aroclor-1254	5	---			0.0	5	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1260	1	---			0.0	1	---			0.0
Aroclor-1260	2	---			0.0	2	---			0.0
Aroclor-1260	3	---			0.0	3	---			0.0
Aroclor-1260	4	---			0.0	4	---			0.0
Aroclor-1260	5	---			0.0	NS	---			----
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1262	1	---			0.0	1	---			0.0
Aroclor-1262	2	---			0.0	2	---			0.0
Aroclor-1262	3	---			0.0	3	---			0.0
Aroclor-1262	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1268	1	---			0.0	1	---			0.0
Aroclor-1268	2	---			0.0	2	---			0.0
Aroclor-1268	3	---			0.0	3	---			0.0
Aroclor-1268	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				

Total PCB Area Coll (5.911 - 13.797) = 596193

Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 199492 Col2 Total PCB = 0.0 ppm*

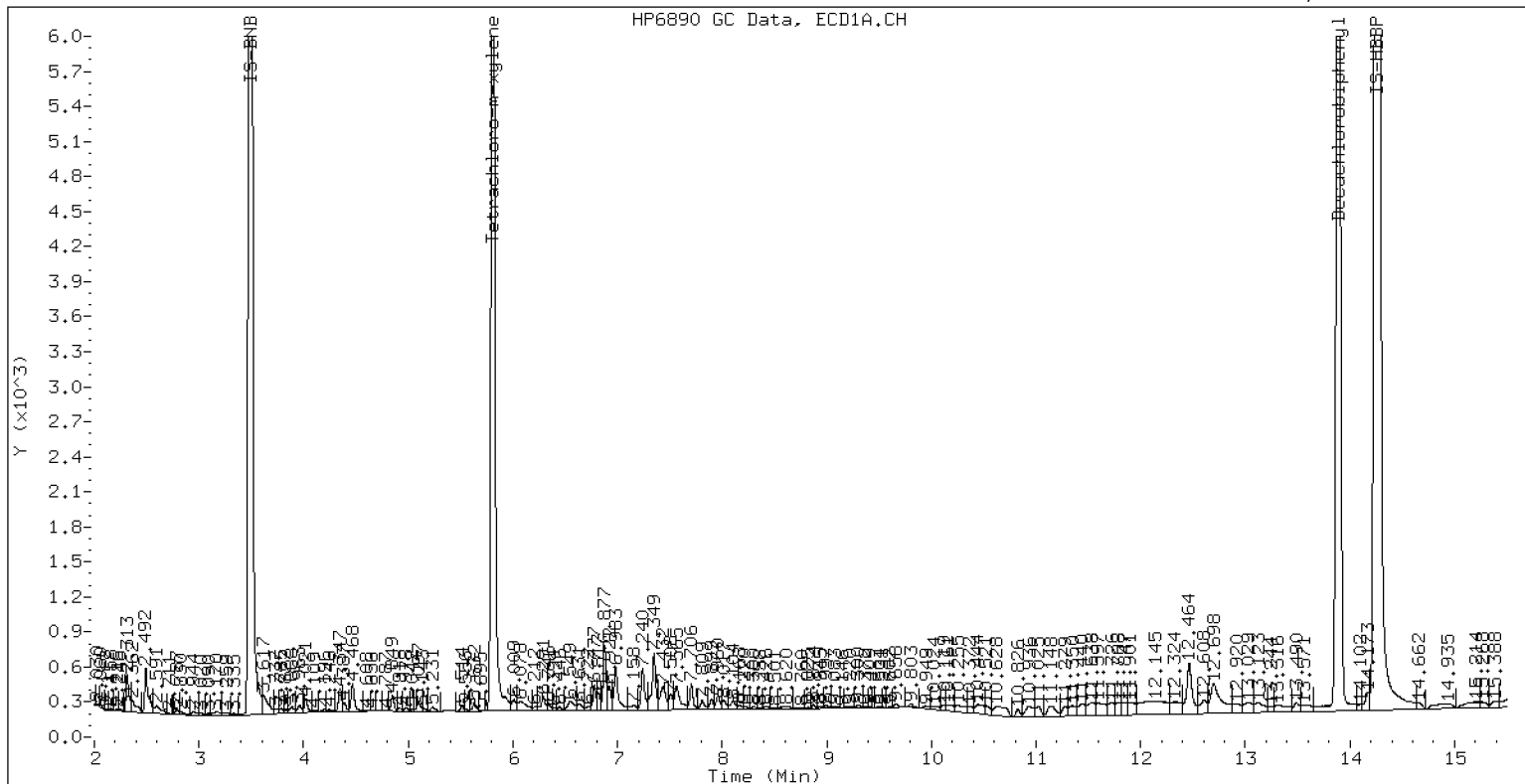
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 23C0107-02

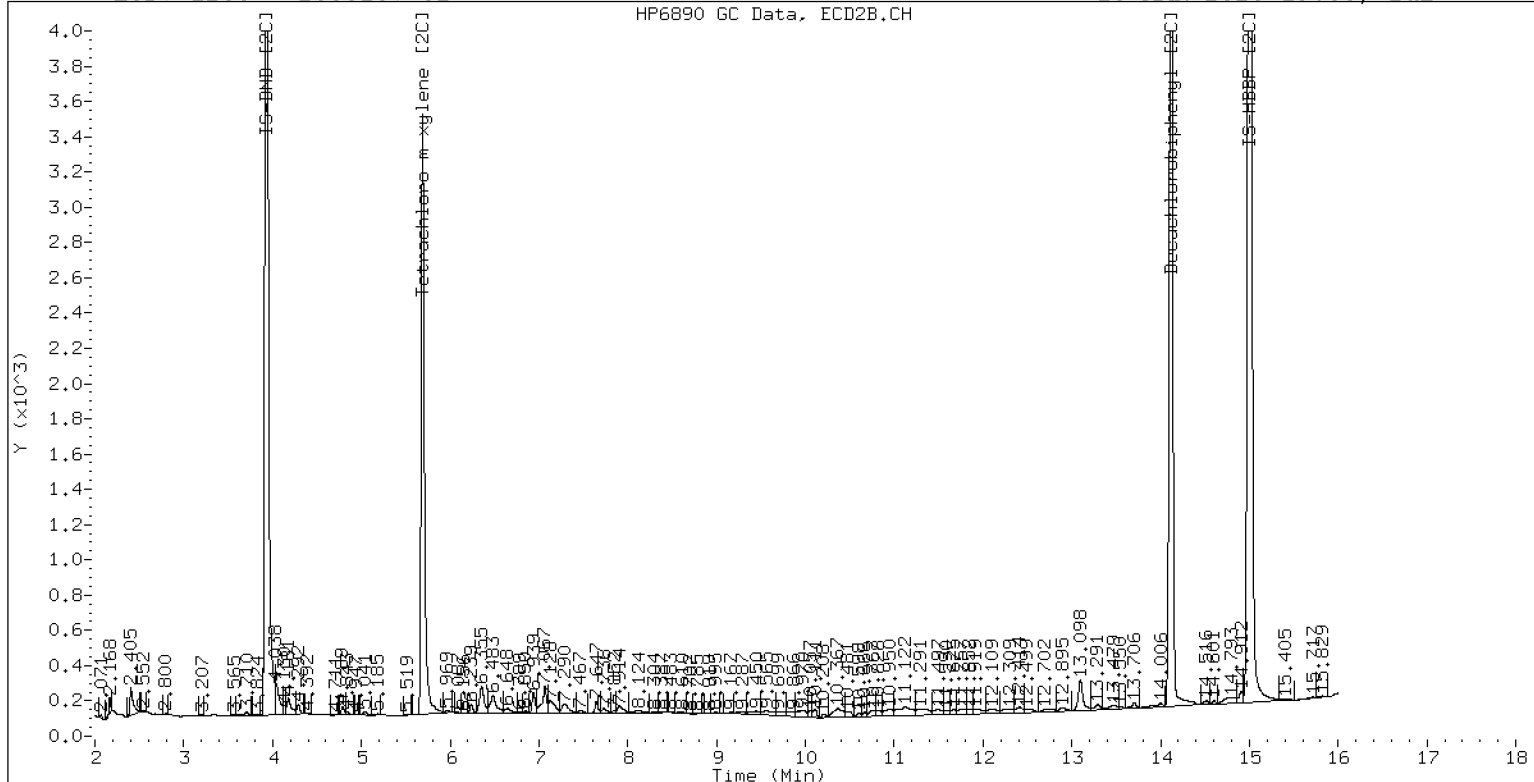
15-MAR-2023 19:35, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 23C0107-02

15-MAR-2023 19:35, 2ul



ZB-35 Manual Integration: NO



ORGANIC ANALYSIS DATA SHEET
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Laboratory ID: 23C0107-03 A

File ID: 03152321ECD7.D

Sampled: 07/13/21 06:54

Prepared: 03/10/23 11:58

Analyzed: 03/15/23 19:56

% Solids: 72.77

Preparation: EPA 3546 (Microwave)

Initial/Final: 17.2 g Wet / 2.5 mL

Batch: BLC0219

Sequence: SLC0215

Calibration: GB00069

Instrument: ECD7

Column 1: ZB5

Column 2: ZB35

CAS NO.	COMPOUND	Col #	DILUTION	CONC. (ug/kg dry)	MDL	MRL	Q
12674-11-2	Aroclor 1016	1	1	4.0	1.6	4.0	H, U
11104-28-2	Aroclor 1221	1	1	4.0	1.6	4.0	H, U
11141-16-5	Aroclor 1232	1	1	4.0	1.6	4.0	H, U
53469-21-9	Aroclor 1242	1	1	4.0	1.6	4.0	H, U
12672-29-6	Aroclor 1248	1	1	4.0	1.6	4.0	H, U
11097-69-1	Aroclor 1254	1	1	4.0	1.6	4.0	H, U
11096-82-5	Aroclor 1260	1	1	4.0	0.6	4.0	H, U

SURROGATES	Col #	ADDED (ug/kg dry)	CONC (ug/kg dry)	% REC	QC LIMITS	Q
<i>Decachlorobiphenyl</i>	1	7.9895	5.65	70.7	40 - 126	Q
<i>Tetrachlorometaxylene</i>	1	7.9895	5.79	72.4	44 - 120	

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152321ECD7.D
Data file 2: /230315.b/230315.b/03152321ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 23C0107-03
Client ID:
Injection Date: 15-MAR-2023 19:56
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.809	-0.002	477566	5.690	-0.002	169193	29.0	28.4	1.9	Tetrachloro-m-xylene
13.893	-0.004	616100	14.119	-0.002	293042	28.3	34.6	20.1	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1104252	63.9
Hexabromobiphenyl	1429847	2211995	54.7

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	405699	28.7
Hexabromobiphenyl	513946	556397	8.3

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	---			0.0	1	---			0.0
Aroclor-1016	2	---			0.0	2	---			0.0
Aroclor-1016	3	---			0.0	3	---			0.0
Aroclor-1016	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	---			0.0	2	---			0.0
Aroclor-1221	3	---			0.0	3	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	---			0.0	2	---			0.0
Aroclor-1232	3	---			0.0	3	---			0.0
Aroclor-1232	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1242	1	---			0.0	1	---			0.0
Aroclor-1242	2	---			0.0	2	---			0.0
Aroclor-1242	3	---			0.0	3	---			0.0
Aroclor-1242	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1248	1	---			0.0	1	---			0.0
Aroclor-1248	2	---			0.0	2	---			0.0
Aroclor-1248	3	---			0.0	3	---			0.0
Aroclor-1248	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1254	1	---			0.0	1	---			0.0
Aroclor-1254	2	---			0.0	2	---			0.0
Aroclor-1254	3	---			0.0	3	---			0.0
Aroclor-1254	4	---			0.0	4	---			0.0
Aroclor-1254	5	---			0.0	5	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1260	1	---			0.0	1	---			0.0
Aroclor-1260	2	---			0.0	2	---			0.0
Aroclor-1260	3	---			0.0	3	---			0.0
Aroclor-1260	4	---			0.0	4	---			0.0
Aroclor-1260	5	---			0.0	NS	---			----
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1262	1	---			0.0	1	---			0.0
Aroclor-1262	2	---			0.0	2	---			0.0
Aroclor-1262	3	---			0.0	3	---			0.0
Aroclor-1262	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1268	1	---			0.0	1	---			0.0
Aroclor-1268	2	---			0.0	2	---			0.0
Aroclor-1268	3	---			0.0	3	---			0.0
Aroclor-1268	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				

Total PCB Area Coll (5.911 - 13.797) = 381367

Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 191885 Col2 Total PCB = 0.0 ppm*

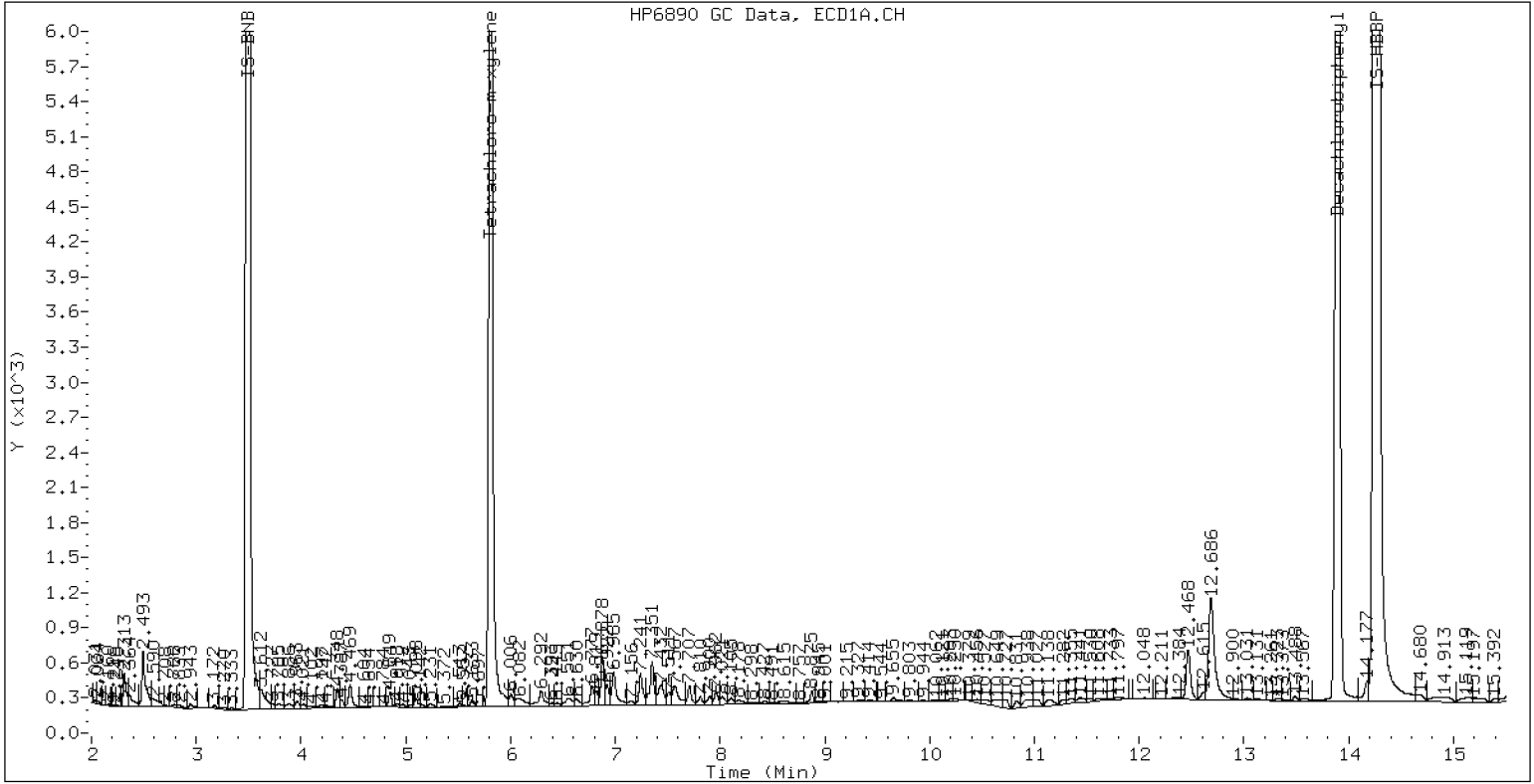
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 23C0107-03

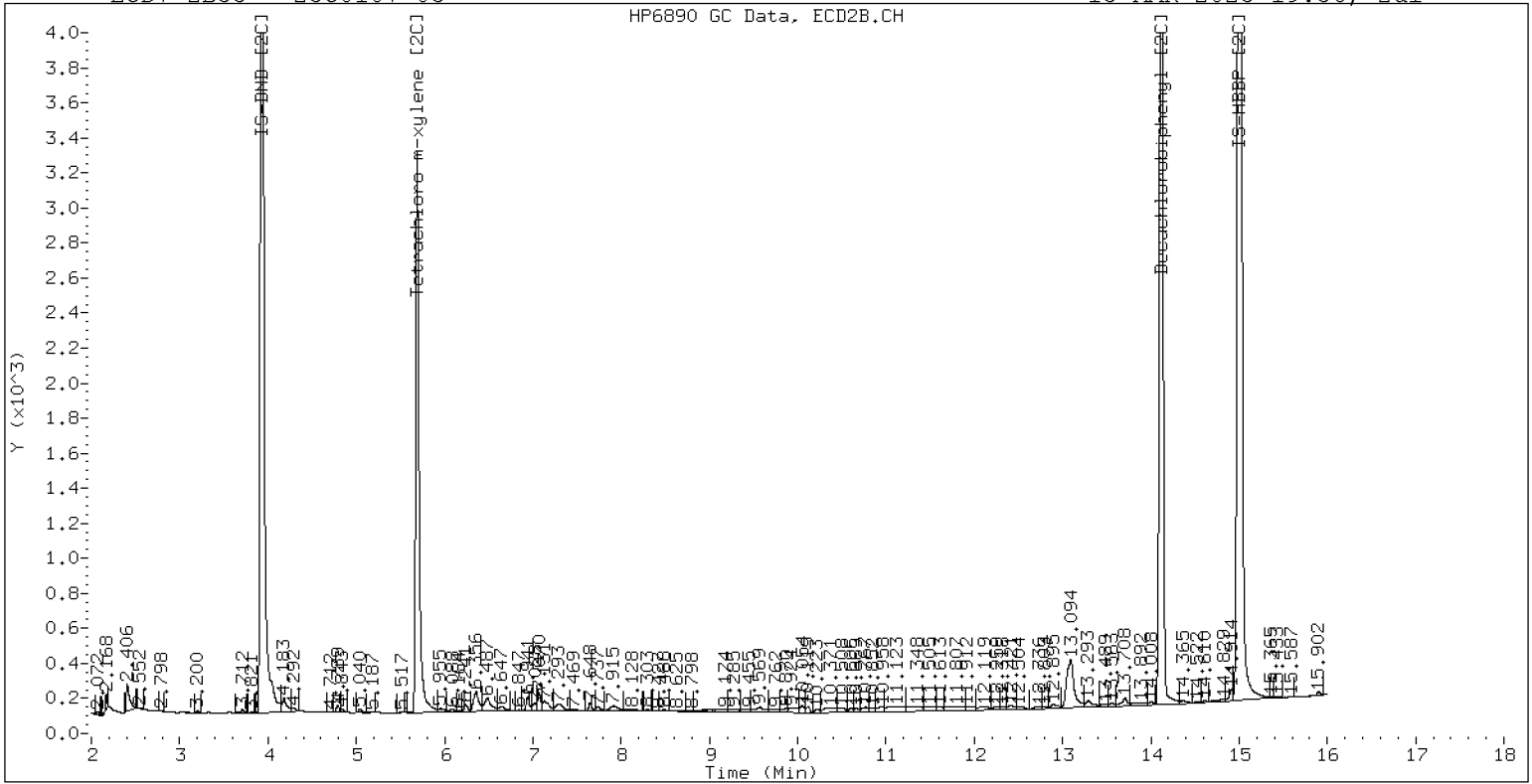
15-MAR-2023 19:56, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 23C0107-03

15-MAR-2023 19:56, 2ul



ZB-35 Manual Integration: NO



Batch: BLC0219

Prepared using: EPA 3546 (Microwave)

8082A PCB Solid 4 in Solid (Version:7 Aroclors)

Matrix: Solid

Date Prepared: 03/14/23

Balance ID: 3146462614

Set Up By: CFO 3/9/23

WO Comments

23C0107: <C>BPR SRM, MS, DUP <C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36,K011477-79, MS/MSD <E>
<H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)
23C0108: <C>BPR SRM, MS, DUP <C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36,K011477-79, MS/MSD <E>
<H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)
23C0109: <C>BPR SRM, MS, DUP <C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36,K011477-79, MS/MSD <E>
<H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)

The following standards may be missing from this batch!

Designator	Description
QLS 5	QLS Spike

Analysis: 8082A PCB Solid 4

Lab Number & Container	% Solids	Initial (g)		(REQ) Acid C/U (5mL)	(REQ) Sulfur C/U (5mL)	(REQ) Silica Gel C/U (2:5)	Final Effective Vol (mL)	Vol (mL) to Lab	Extraction Comments
		Target Dry: 12.5 (Wet)	Actual						
23C0107-01 A	41.9	(29.83)	29.89	5mL	5mL	2mL	2.5	1.0	
23C0107-02 A	67.2	(18.60)	18.61	5mL	5mL	2mL	2.5	1.0	
23C0107-03 A	72.8	(17.18)	17.24	5mL	5mL	2mL	2.5	1.0	
23C0108-01 A	53.6	(23.34)	23.37	5mL	5mL	2mL	2.5	1.0	
23C0108-02 A	52.7	(23.70)	23.70	5mL	5mL	2mL	2.5	1.0	
23C0108-03 A	48.8	(25.63)	25.63	5mL	5mL	2mL	2.5	1.0	
23C0108-04 A	47.4	(26.37)	26.39	5mL	5mL	2mL	2.5	1.0	
23C0108-05 A	48.6	(25.74)	25.79	5mL	5mL	2mL	2.5	1.0	
23C0108-06 A	43.3	(28.85)	28.88	5mL	5mL	2mL	2.5	1.0	
23C0108-07 A	40.2	(31.10)	31.13	5mL	5mL	2mL	2.5	1.0	
23C0108-08 A	46.4	(26.95)	26.97	5mL	5mL	2mL	2.5	1.0	
23C0108-09 A	42.6	(29.33)	29.33	5mL	5mL	2mL	2.5	1.0	
23C0108-10 A	47.3	(26.43)	26.43	5mL	5mL	2mL	2.5	1.0	
23C0109-01 A	43.3	(28.84)	28.87	5mL	5mL	2mL	2.5	1.0	
23C0109-02 A	35.9	(34.82)	34.89	5mL	5mL	2mL	2.5	1.0	
23C0109-03 A	36.1	(34.60)	34.60	5mL	5mL	2mL	2.5	1.0	

Batch QC

Lab Number	% Solids	Initial (g)		(REQ) Acid C/U (5mL)	(REQ) Sulfur C/U (5mL)	(REQ) Silica Gel C/U (2:5)	Final Effective Vol (mL)	Vol (mL) to Lab	Extraction Comments
		Target Dry: 12.5 (Wet)	Actual						
BLC0219-BLK1	100.0	(12.50)	12.54	5mL	5mL	2mL	2.5	1.0	(10g Actual Wt.)
BLC0219-BS1	100.0	(12.50)	12.54	5mL	5mL	2mL	2.5	1.0	(10g Actual Wt.)
BLC0219-BSD1	100.0	(12.50)	12.54	5mL	5mL	2mL	2.5	1.0	(10g Actual Wt.)
BLC0219-MS1	72.8	(17.18)	17.18	5mL	5mL	2mL	2.5	1.0	Use 23C0107-03
BLC0219-MSD1	72.8	(17.18)	17.18	5mL	5mL	2mL	2.5	1.0	Use 23C0107-03
BLC0219-SRM1	100.0	(12.50)	12.54	5mL	5mL	2mL	2.5	1.0	Use K003527

+1g DI WATER



Batch: BLC0219

Prepared using: EPA 3546 (Microwave)

8082A PCB Solid 4 in Solid (Version:7 Aroclors)

WO Comments

23C0107: <C>BPR SRM, MS, DUP </C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36, K011477-79, MS/MSD </E>

<H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)

23C0108: <C>BPR SRM, MS, DUP </C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36, K011477-79, MS/MSD </E>

<H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)

23C0109: <C>BPR SRM, MS, DUP </C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36, K011477-79, MS/MSD </E>

<H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)

<p>Client ID Verified By <u>aj 03/14/23</u></p>	<p>Date</p>	<p>Preparation Reviewed By <u>LJ 3/15/23</u></p>	<p>Date</p>	<p>Extraction Date and Time <u>03/14/23 11:58</u></p>
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Batch: BLC0219

Prepared using: EPA 3546 (Microwave)
8082A PCB Solid 4 in Solid (Version:7 Aroclors)

WO Comments
 23C0107: <C>BPR SRM, MS, DUP <C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36, K011477-79, MS/MSD <E>
 <H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)
 23C0108: <C>BPR SRM, MS, DUP <C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36, K011477-79, MS/MSD <E>
 <H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)
 23C0109: <C>BPR SRM, MS, DUP <C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36, K011477-79, MS/MSD <E>
 <H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)

Prep Steps	Reagents Used	Standard ID	Surrogates & Spike Standards Used				
Microwave ① 2 3 CT 3/11/23 Analyst/Date	Station/Reagent Microwave Analyst: CT/196 Date: 3/11/23	Standard ID Neutral Glass Wool L000497	Type Surrogate 2µg/mL Spike 20µg/mL	Vial ID / Standard ID N L000773 Exp Date: 7/21/23 1 L001587 Exp Date: 8/13/23	Vol uL 50µL 63µL	Analyst CT G	Witness ✓ ✓
	1:1 Hexane/Acetone Hexane Anhydrous Sodium Sulfate	L0002245 L001957 L002484					
KD 100°C Hexane Exchange (2 X 20 mL) ① 2 3 4 ⑤ 6 LD 3-14 Analyst/Date	KD Analyst: LD Date: 3-14-23	Anhydrous Sodium Sulfate Hexane	MANUALLY ENTER EXPIRATION DATES! (V) indicates a virtual standard combining two or more physical standards. In these cases the Standard ID refers to the virtual standard, not the parent standards. If a Standard ID is missing, but should be present, check the standard definition in Element LIMS to be sure Standard Info 6 has the correct letter or number designator matching the vial designator in the Standard ID column. If it is correct, check the batch and bench sheet in Element LIMS to be sure the correct standards are selected for surrogate(s) and spike(s).				
	Vialing Analyst: LJ Date: 3/15/23	Hexane Concentrated Sulfuric Acid					
TurboVap Pre Cleanups 1 2 3 ④ 5 LJ 3/14/23 Analyst/Date	Hexane Concentrated Sulfuric Acid	L000881 L001033					
	TurboVap Post Cleanups 1 2 ③ 4 5 LJ 3/15/23 Analyst/Date	Silica Gel (SPE) Darts Sodium Sulfite Tetrabutylammonium hydrogensulfate (TBAS)					L002256 L002437 L001601
Vialing LJ 3/15/23 Analyst/Date							



Batch: BLC0219

Prepared using: EPA 3546 (Microwave)
8082A PCB Solid 4 in Solid (Version:7 Aroclors)

WO Comments
 23C0107: <C>BPR SRM, MS, DUP </C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36,K011477-79, MS/MSD </E>
 <H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)
 23C0108: <C>BPR SRM, MS, DUP </C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36,K011477-79, MS/MSD </E>
 <H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)
 23C0109: <C>BPR SRM, MS, DUP </C> <M>BPR PS, MS/MSD </M> <E>BPR 8270E RM K000591, SIM PAH RM I009127 PCB RM J006840-43, 7935-36,K011477-79, MS/MSD </E>
 <H>BPR J006840-43, 7935-36, K011477-79, Dup </H> Store in freezer (except GS)

Prep Instructions	
<p>SPECIAL INSTRUCTIONS:</p> <ol style="list-style-type: none"> 1. Weigh soil/sed into beakers-lightly dry with sodium sulfate. 2. Transfer to microwave vessel(s). Note: (do not fill vessels more than 2/3rd full. Some samples may require two vessels). 3. Add 1:1 Hexane/Acetone until the solvent layer is 3 inches above the soil layer after homogenization. 4. Add surr/spike. 5. Microwave on appropriate power setting determined by # of samples. 6. After microwave-Re-homogenize while hot then cool vessels in R-05 15 minutes. Re-homogenize while cool. 7. Decant 1:1 Hex/Ace into Erlenmeyer flask with sodium sulfate in bottom and funnel with neutral glasswool plug. 8. Re-homogenize and rinse with 1:1 Hexane/Acetone. 9. Let cool and decant solvent then empty the soil into the funnel and rinse with Hexane. 10. KD on 100° bath. 11. Exchange (2 X with 20mL) Hexane. 12. TurboVap. 13. Clean-ups. 14. TurboVap. 15. Vial with Hexane. <p>A. Need Total Solids Y <input type="checkbox"/> N <input checked="" type="checkbox"/></p> <p>B. Archive/Freeze <input checked="" type="checkbox"/> Y <input type="checkbox"/> N</p>	



Extraction Parameter: PCB Extraction Batch BL0219

Total Solids Batch: BLG BL0101 Work Order(s): 2302107, 108, 109

Screens: Soil/Sediment/Solid/Other:	Analyst/Date
<input checked="" type="checkbox"/> No Anomalies (standard soil/wet sediment/sand/gravel)= 108, 109	CR 3/7/23
<input checked="" type="checkbox"/> Standing Water Decanted (Not shared)= 107:01	CR 3/7/23
<input type="checkbox"/> Standing Water Homogenized (Shared samples)=	
<input type="checkbox"/> Clay/Clumps (Difficult to homogenize)=	
<input type="checkbox"/> Rocks (%+size)?	
<input type="checkbox"/> Organics (Leaves/sticks/grass)=	
<input checked="" type="checkbox"/> Oily, obvious fuel/sulfur odors= 107:01	CR 3/7/23
<input type="checkbox"/> Received in 32oz jar(s)=Homogenized in Pyrex dish=	
<input checked="" type="checkbox"/> Previously Frozen = 107, 108, 109	CR 3/7/23
<input type="checkbox"/> Other (Details)=	
Aqueous:	
<input checked="" type="checkbox"/> No Anomalies	
<input type="checkbox"/> Turbid/Color=	
<input type="checkbox"/> Particulates(%)=(Note: >5%=Notify Supervisor/Lead)	
<input type="checkbox"/> Emulsions (%)=	
<input type="checkbox"/> Oily, obvious fuel/sulfur odors=	
<input type="checkbox"/> Other (Details)=	
<input type="checkbox"/> Received in 1.0L Bottle(s)=No Bottle Rinse=	
<input type="checkbox"/> Other Notes/Comments= (Note problems, concerns, corrective actions).	
<u>Dropped Sci. filtration vial & spilled ~20% of BSI dunks sulfur cleanup</u>	<u>LJ 3/14/23</u>
<input checked="" type="checkbox"/> Share Samples Y/N	CR 3/7/23
<input checked="" type="checkbox"/> Multiple Jars Y/N	CR 3/7/23
<input type="checkbox"/> Sample Pre-Screens indicate analyte activity=	
<input type="checkbox"/> Sample weights/volumes reduced based on Pre-Screen=	



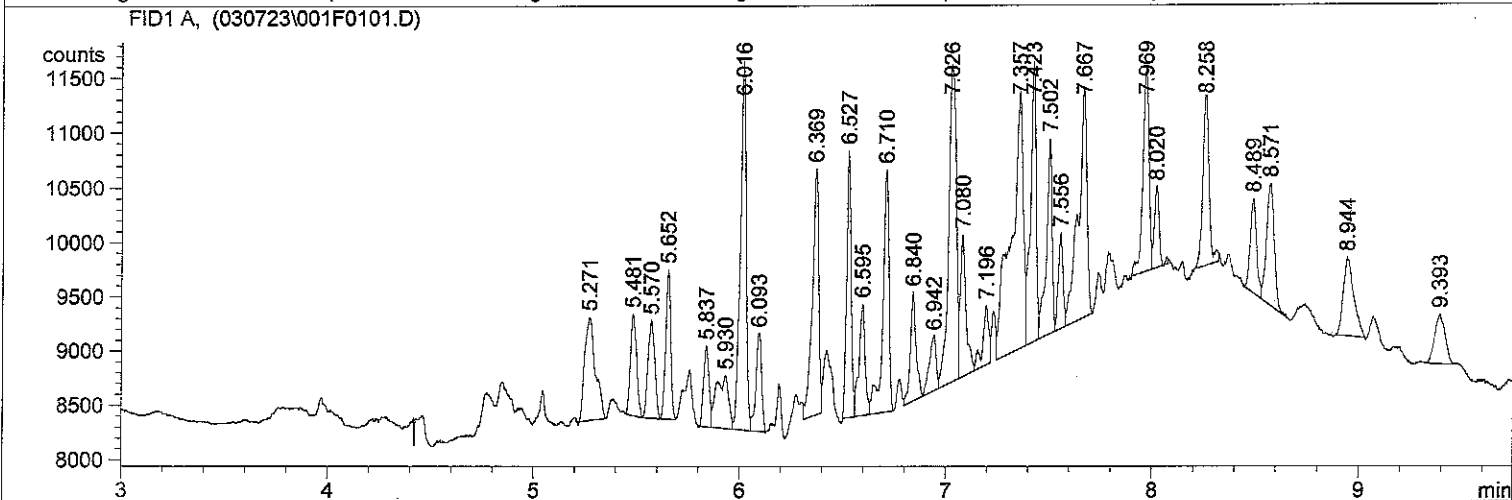
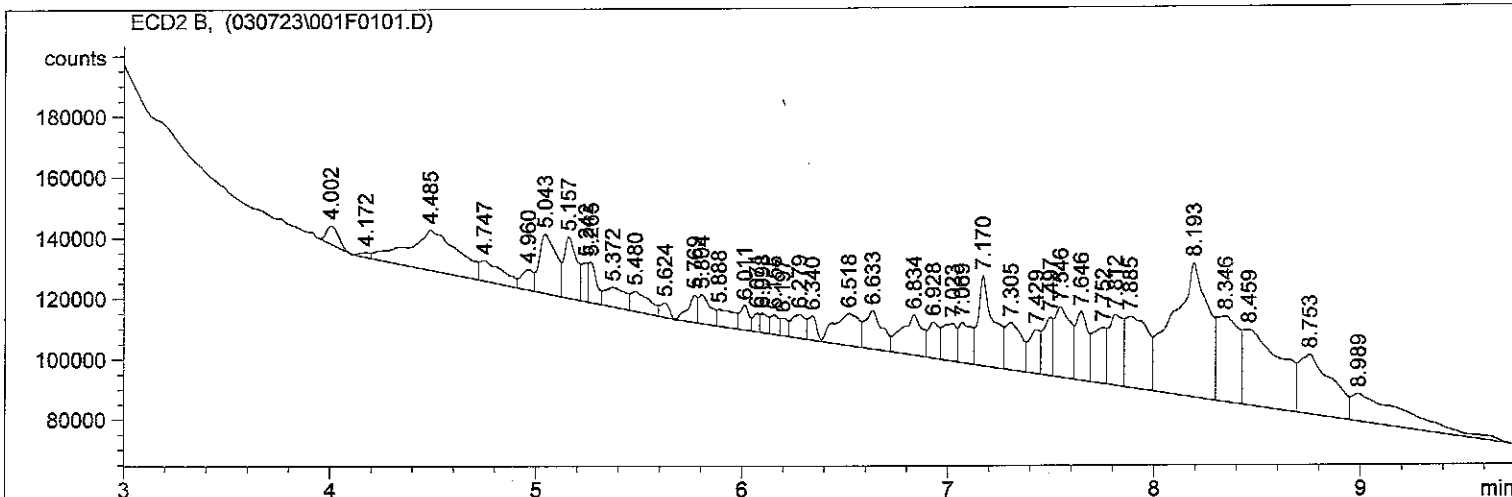
Extraction Parameter: PUB Extraction Batch BL0019

Total Solids Batch: BL0150 Work Order(s): 230809

Screens: Soil/Sediment/Solid/Other:	Analyst/Date
<input checked="" type="checkbox"/> No Anomalies (standard soil/wet sediment/sand/gravel)= <u>φ2, φ3</u>	<u>φ3/φ8/23</u>
<input type="checkbox"/> Standing Water Decanted (Not shared)=	
<input type="checkbox"/> Standing Water Homogenized (Shared samples)=	
<input type="checkbox"/> Clay/Clumps (Difficult to homogenize)=	
<input type="checkbox"/> Rocks (%+size)?	
<input type="checkbox"/> Organics (Leaves/sticks/grass)=	
<input type="checkbox"/> Oily, obvious fuel/sulfur odors=	
<input type="checkbox"/> Received in 32oz jar(s)=Homogenized in Pyrex dish=	
<input type="checkbox"/> Previously Frozen =	
<input type="checkbox"/> Other (Details)=	
Aqueous:	
<input checked="" type="checkbox"/> No Anomalies	
<input type="checkbox"/> Turbid/Color=	
<input type="checkbox"/> Particulates(%)=(Note: >5%=Notify Supervisor/Lead)	
<input type="checkbox"/> Emulsions (%)=	
<input type="checkbox"/> Oily, obvious fuel/sulfur odors=	
<input type="checkbox"/> Other (Details)=	
<input type="checkbox"/> Received in 1.0L Bottle(s)=No Bottle Rinse=	
<input type="checkbox"/> Other Notes/Comments= (Note problems, concerns, corrective actions).	
<input checked="" type="checkbox"/> Share Samples Y / <u>N</u>	<u>φ3/φ8/23</u>
<input checked="" type="checkbox"/> Multiple Jars Y / <u>N</u>	<u>φ3/φ8/23</u>
<input type="checkbox"/> Sample Pre-Screens indicate analyte activity=	
<input type="checkbox"/> Sample weights/volumes reduced based on Pre-Screen=	

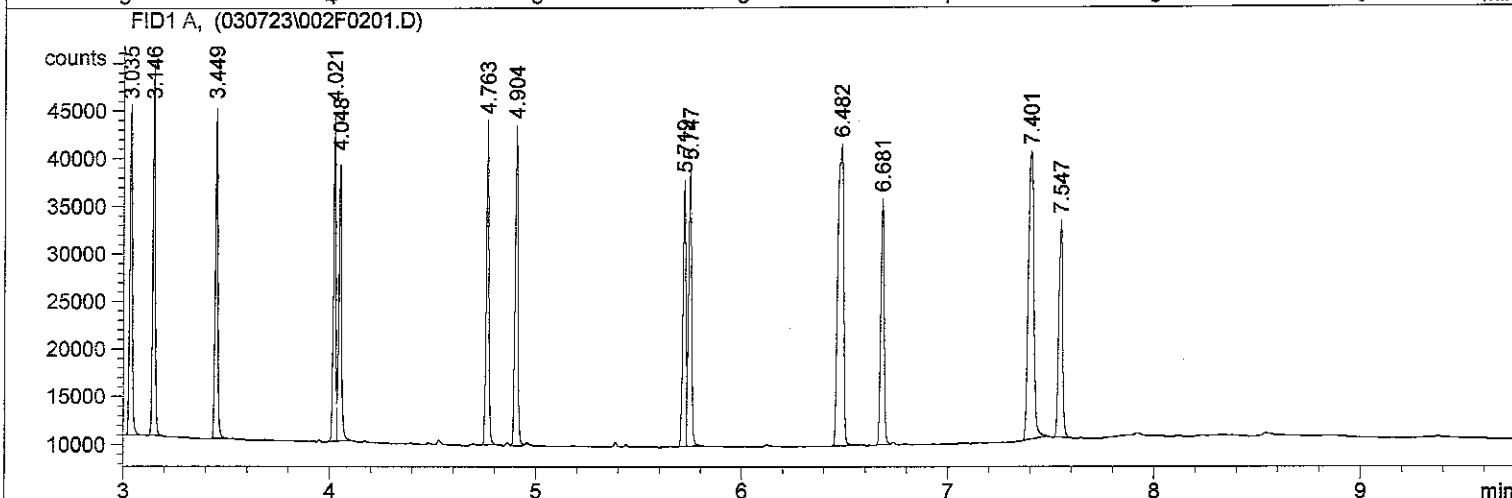
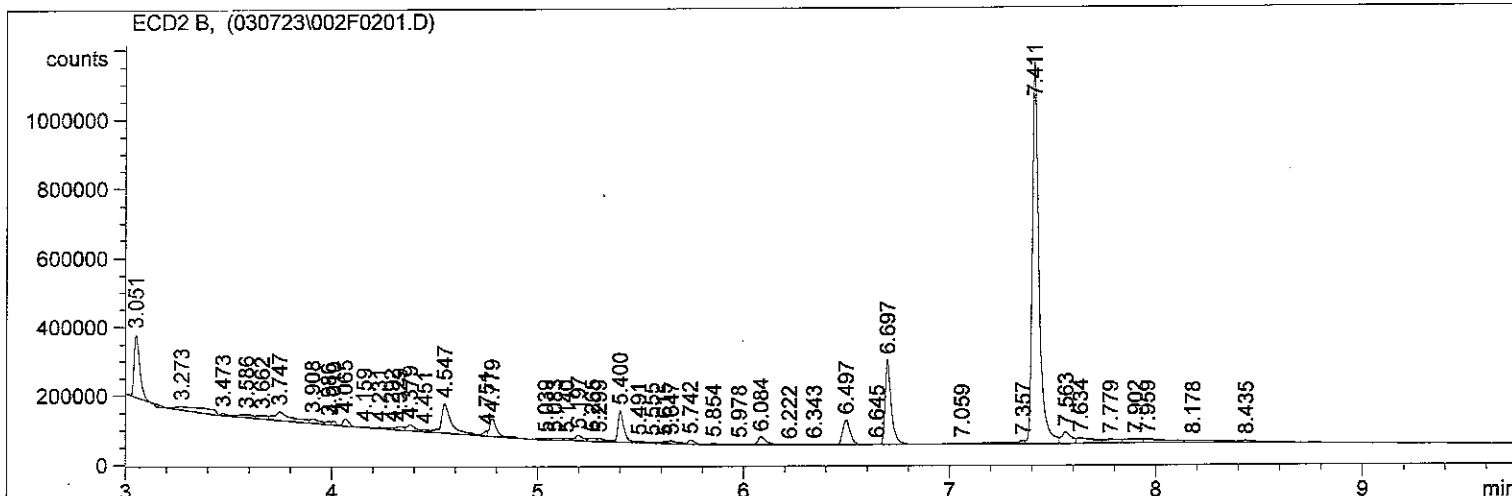
Injection Date : 3/7/2023 3:57:14 PM Seq. Line : 1
Sample Name : DCM RINSE Location : Vial 1
Acq. Operator : CR Inj : 1
Inj Volume : 1 µl

Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***

Injection Date : 3/7/2023 4:11:12 PM Seq. Line : 2
Sample Name : PNA STD 10PPM Location : Vial 2
Acq. Operator : CR Inj : 1
 Inj Volume : 1 µl
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD

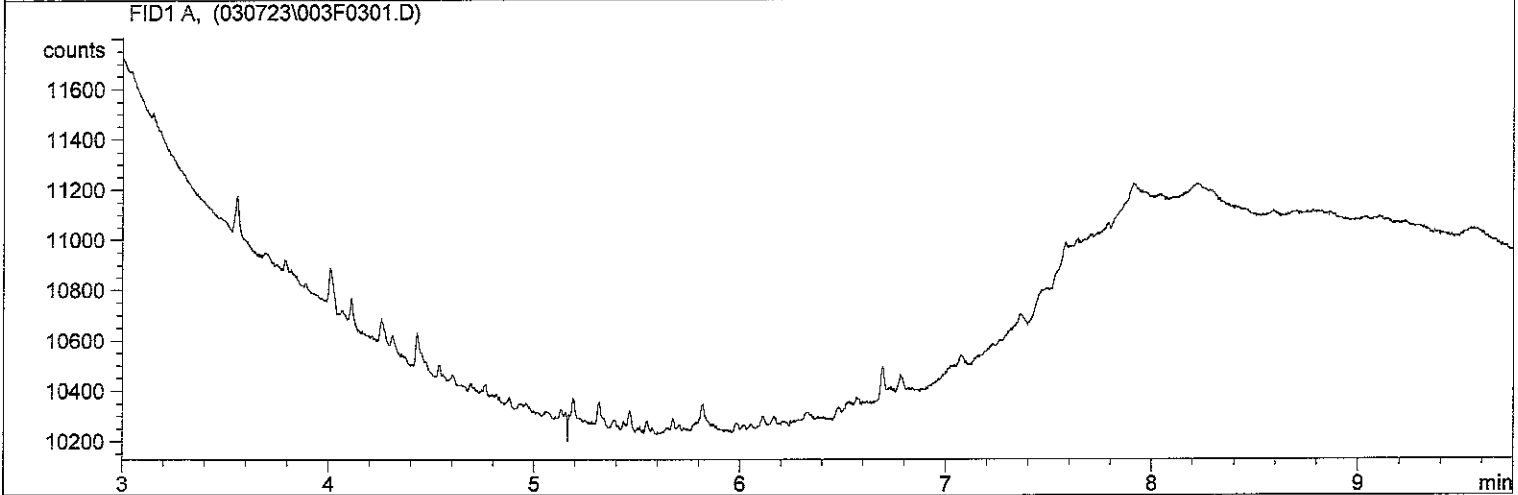
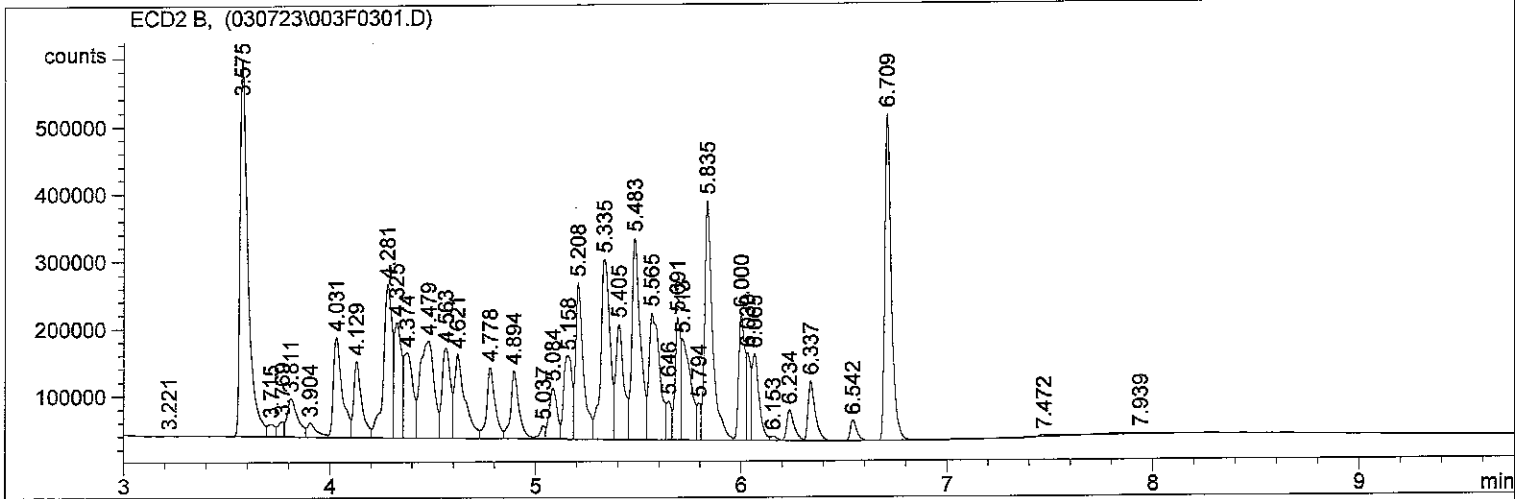


*** End of Report ***

```

=====
Injection Date  : 3/7/2023 4:25:27 PM          Seq. Line :    3
Sample Name    : AR1660 1PPM                  Location  : Vial 3
Acq. Operator  : CR                            Inj       :    1
                                           Inj Volume: 1 µl
Sequence File  : C:\HPCHEM\1\SEQUENCE\030723.S
Method         : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed   : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====

```



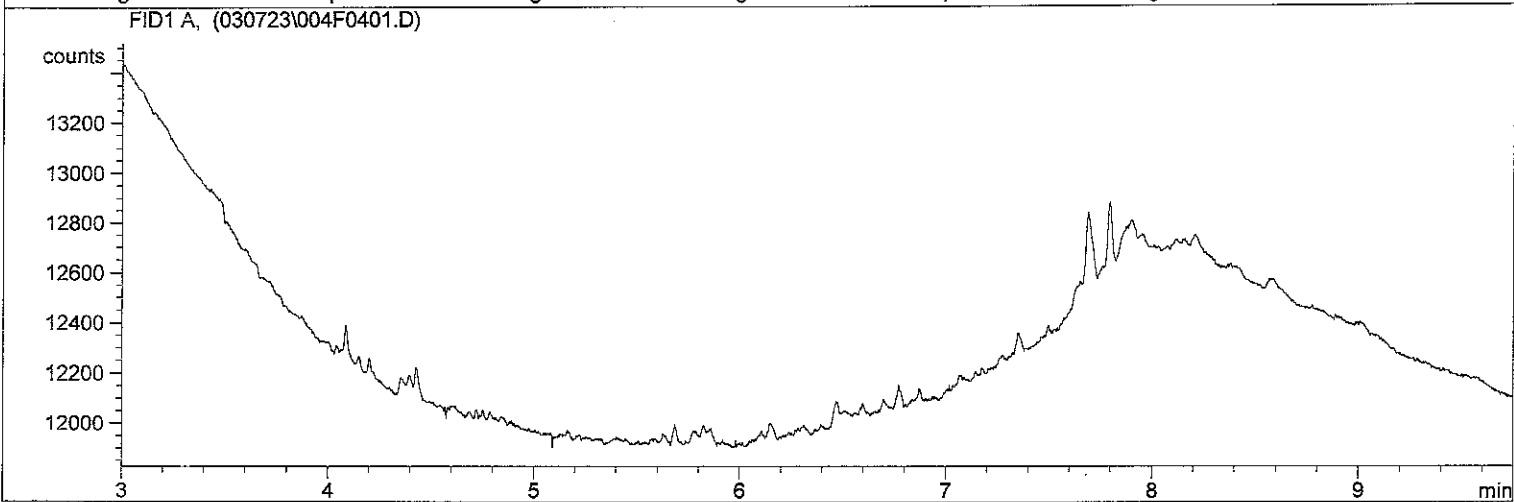
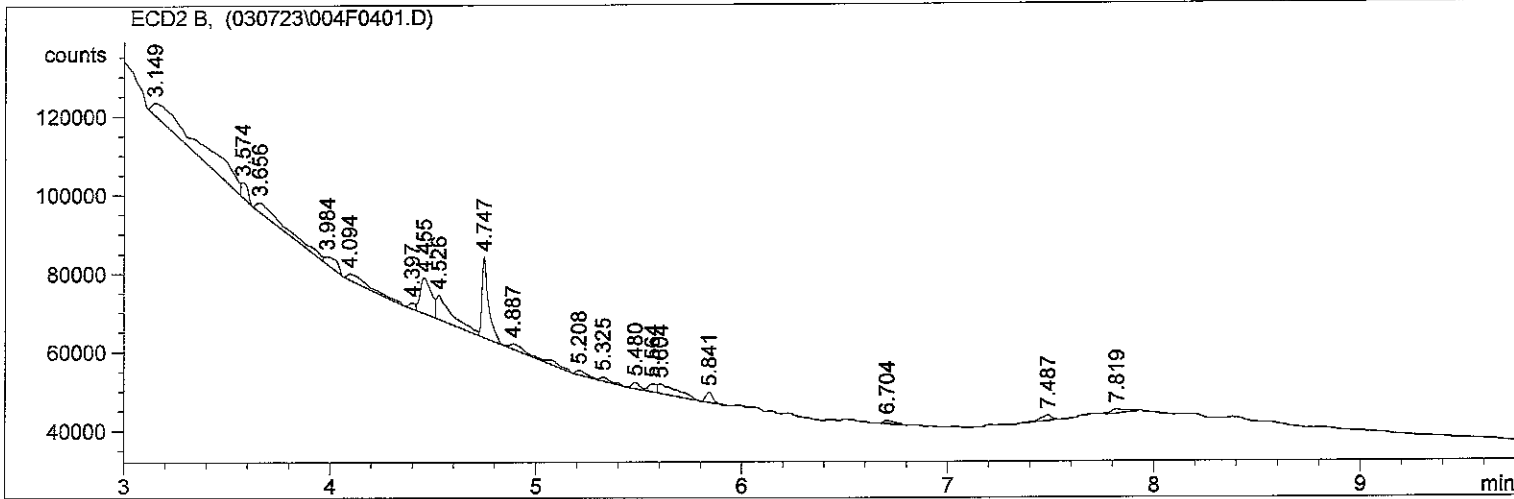
*** End of Report ***

```

Injection Date : 3/7/2023 4:39:28 PM      Seq. Line : 4
Sample Name    : 23C0107 01                Location  : Vial 4
Acq. Operator  : CR                        Inj      : 1
                                           Inj Volume: 1 µl

Sequence File  : C:\HPCHEM\1\SEQUENCE\030723.S
Method         : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed   : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD

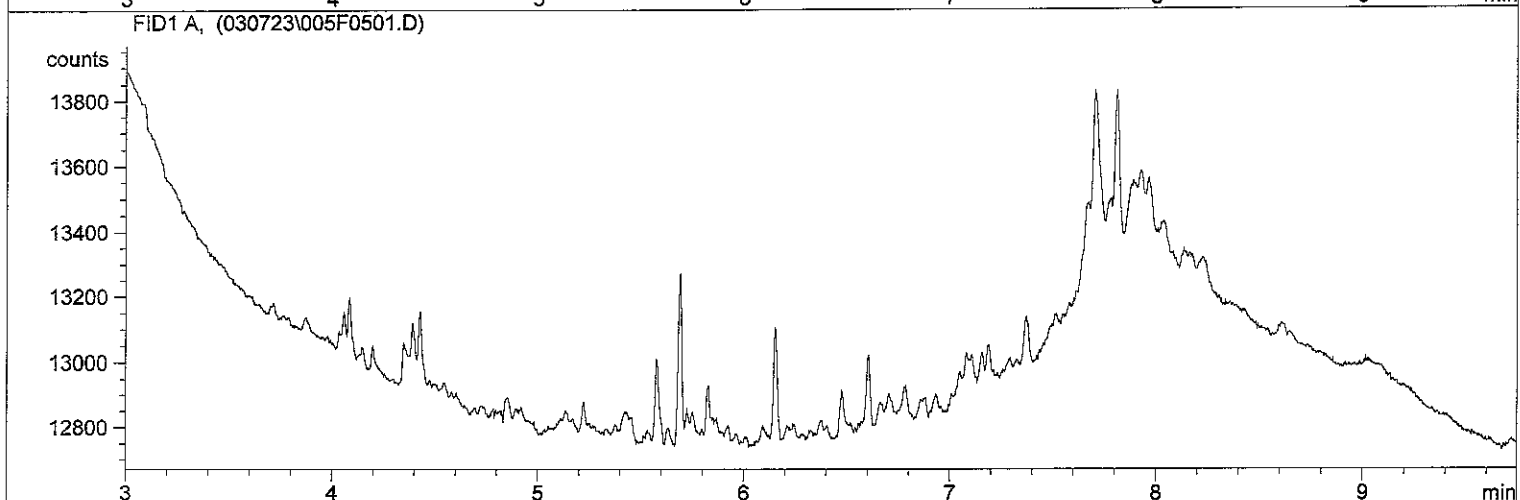
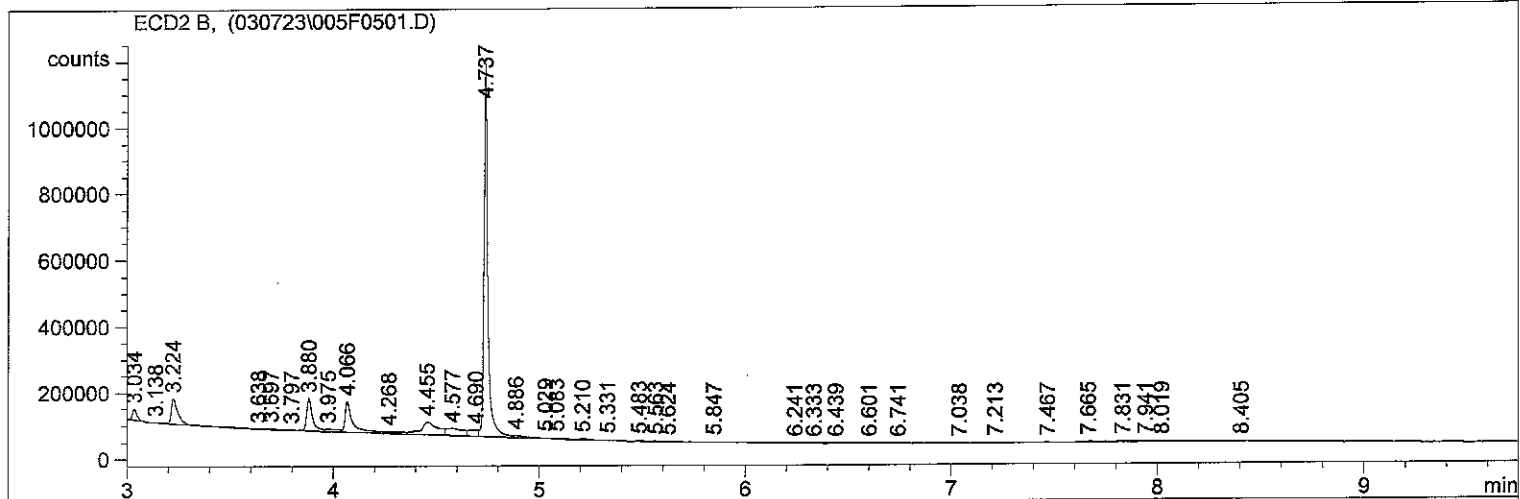
```



*** End of Report ***

```
=====
Injection Date   : 3/7/2023 4:53:45 PM           Seq. Line   :    5
Sample Name     : 23C0108 01                     Location    : Vial 5
Acq. Operator   : CR                             Inj         :    1
                                                    Inj Volume  : 1 µl

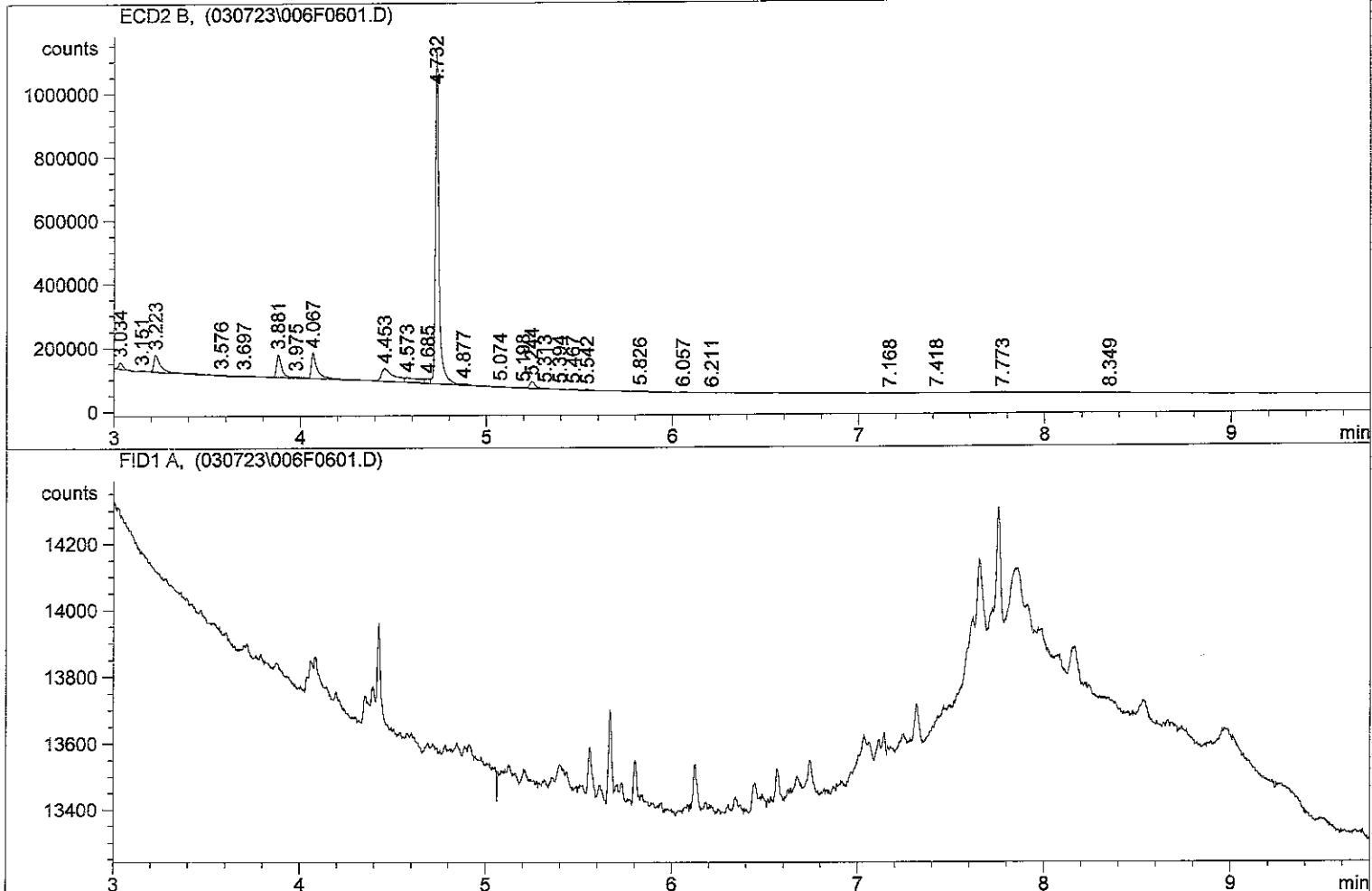
Sequence File   : C:\HPCHEM\1\SEQUENCE\030723.S
Method          : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed    : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====
```



*** End of Report ***

Injection Date : 3/7/2023 5:07:45 PM Seq. Line : 6
Sample Name : 23C0108 02 Location : Vial 6
Acq. Operator : CR Inj : 1
 Inj Volume : 1 µl

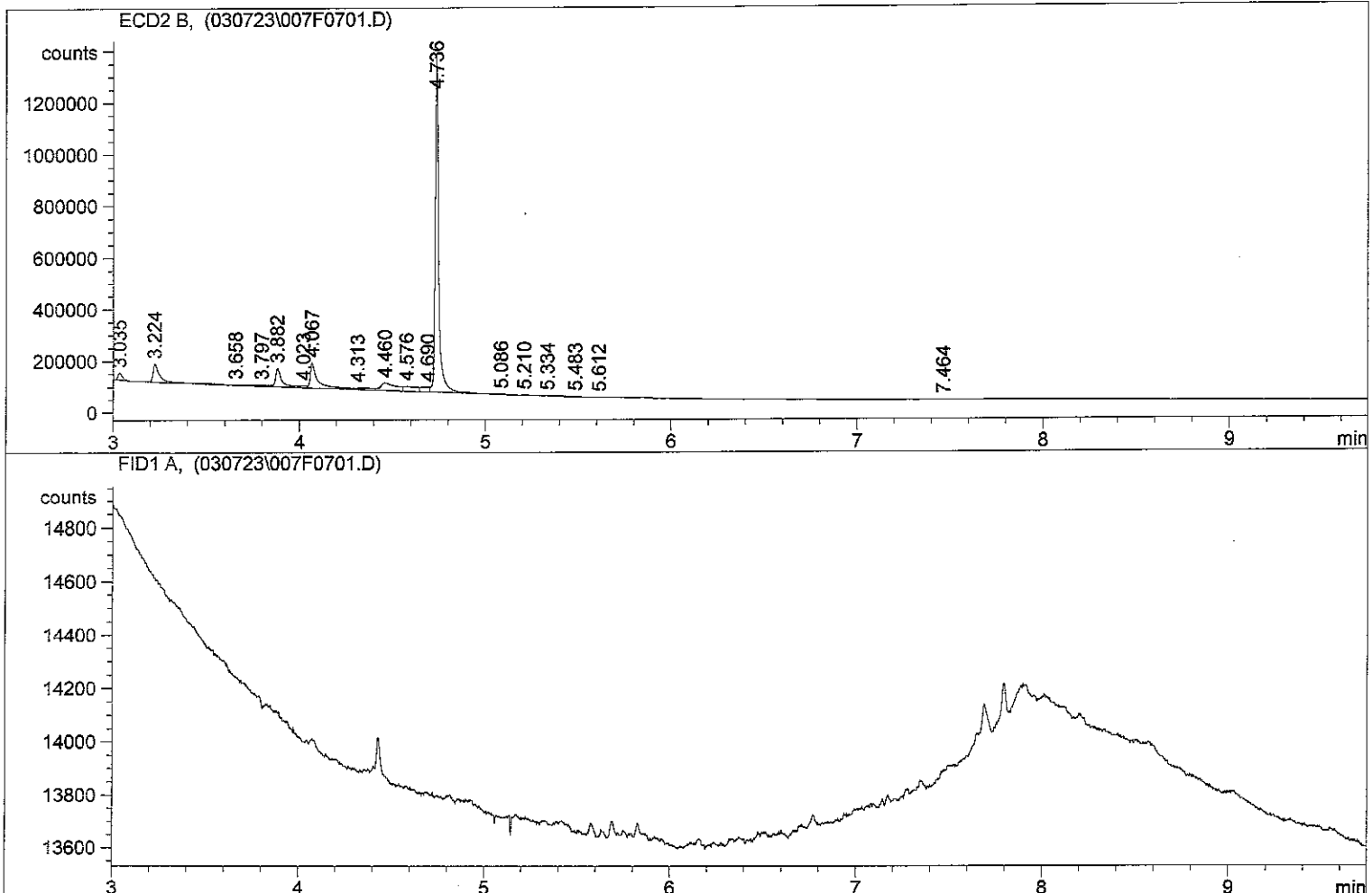
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



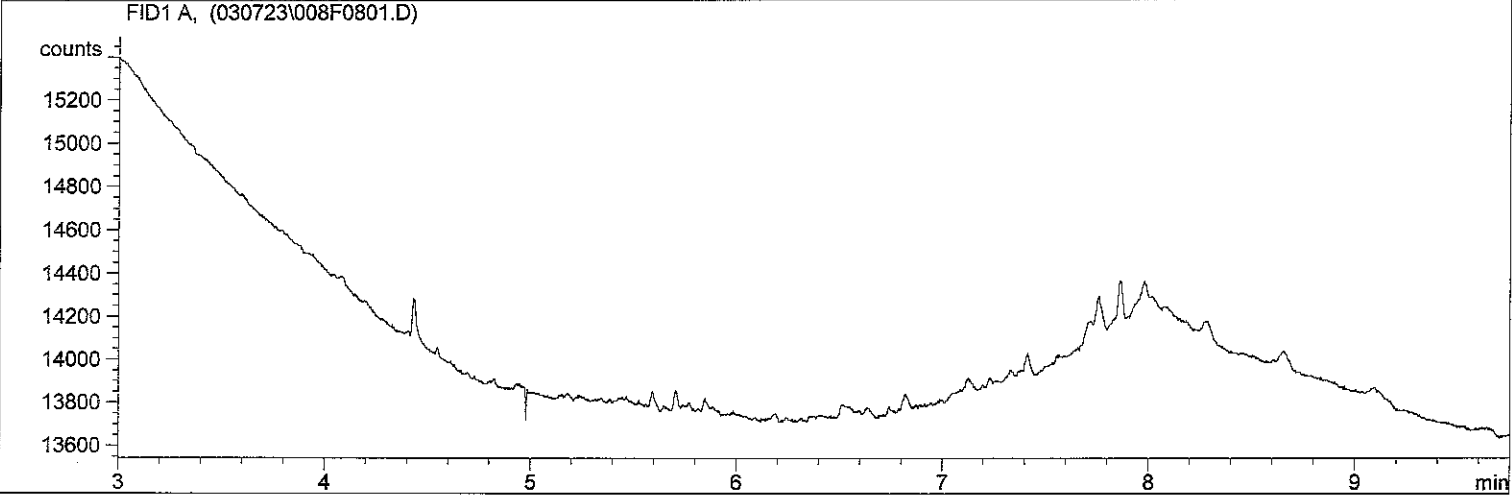
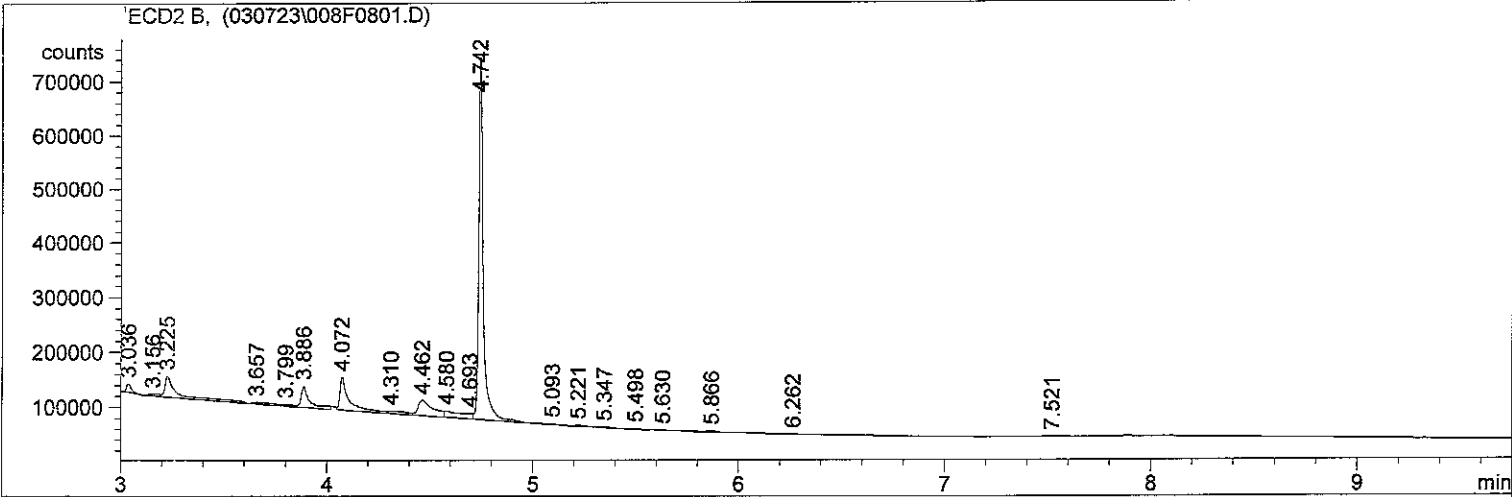
*** End of Report ***

Injection Date : 3/7/2023 5:22:02 PM Seq. Line : 7
Sample Name : 23C0108 03 Location : Vial 7
Acq. Operator : CR Inj : 1
Inj Volume : 1 µl

Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



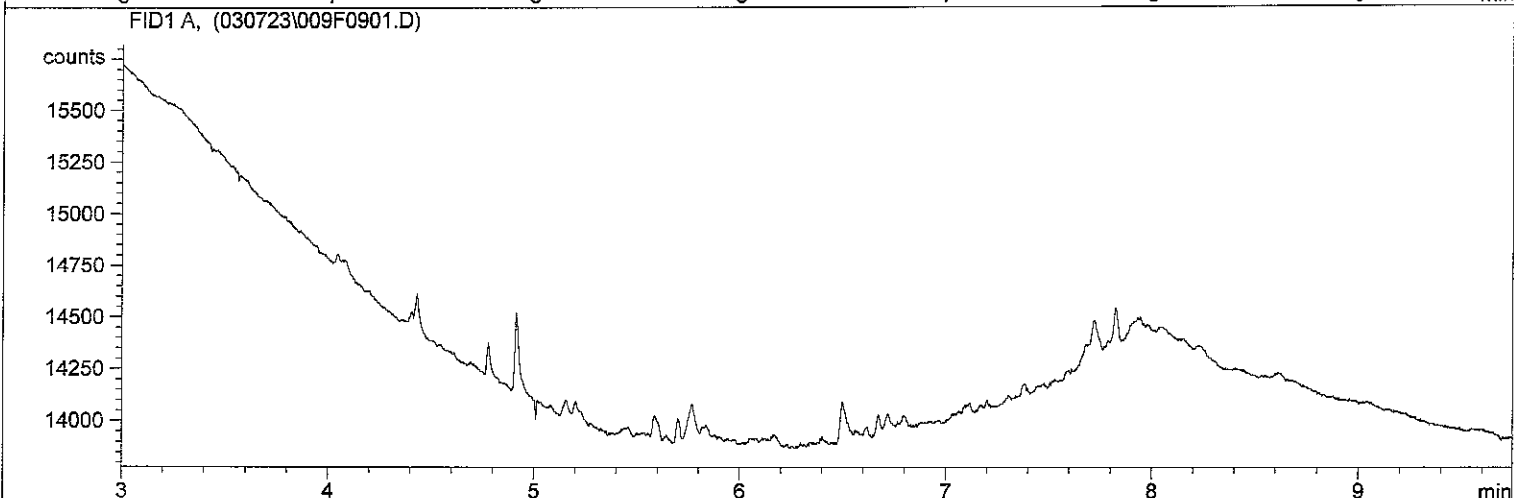
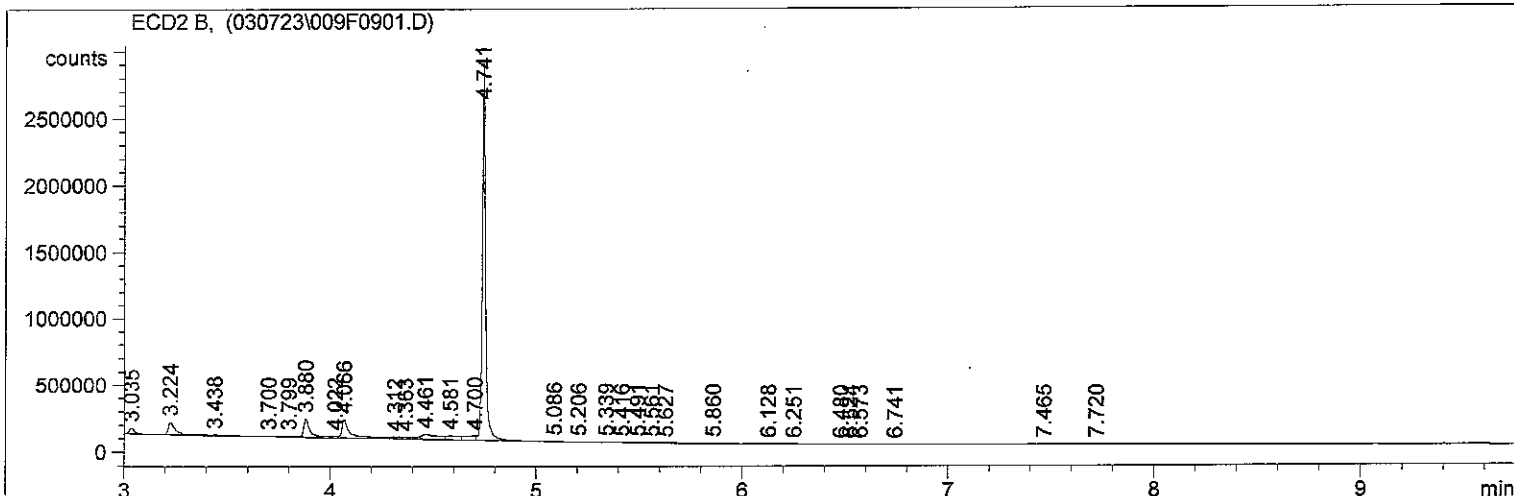
Injection Date : 3/7/2023 5:37:05 PM Seq. Line : 8
Sample Name : 23C0108 04 Location : Vial 8
Acq. Operator : CR Inj : 1
Inj Volume : 1 µl
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***

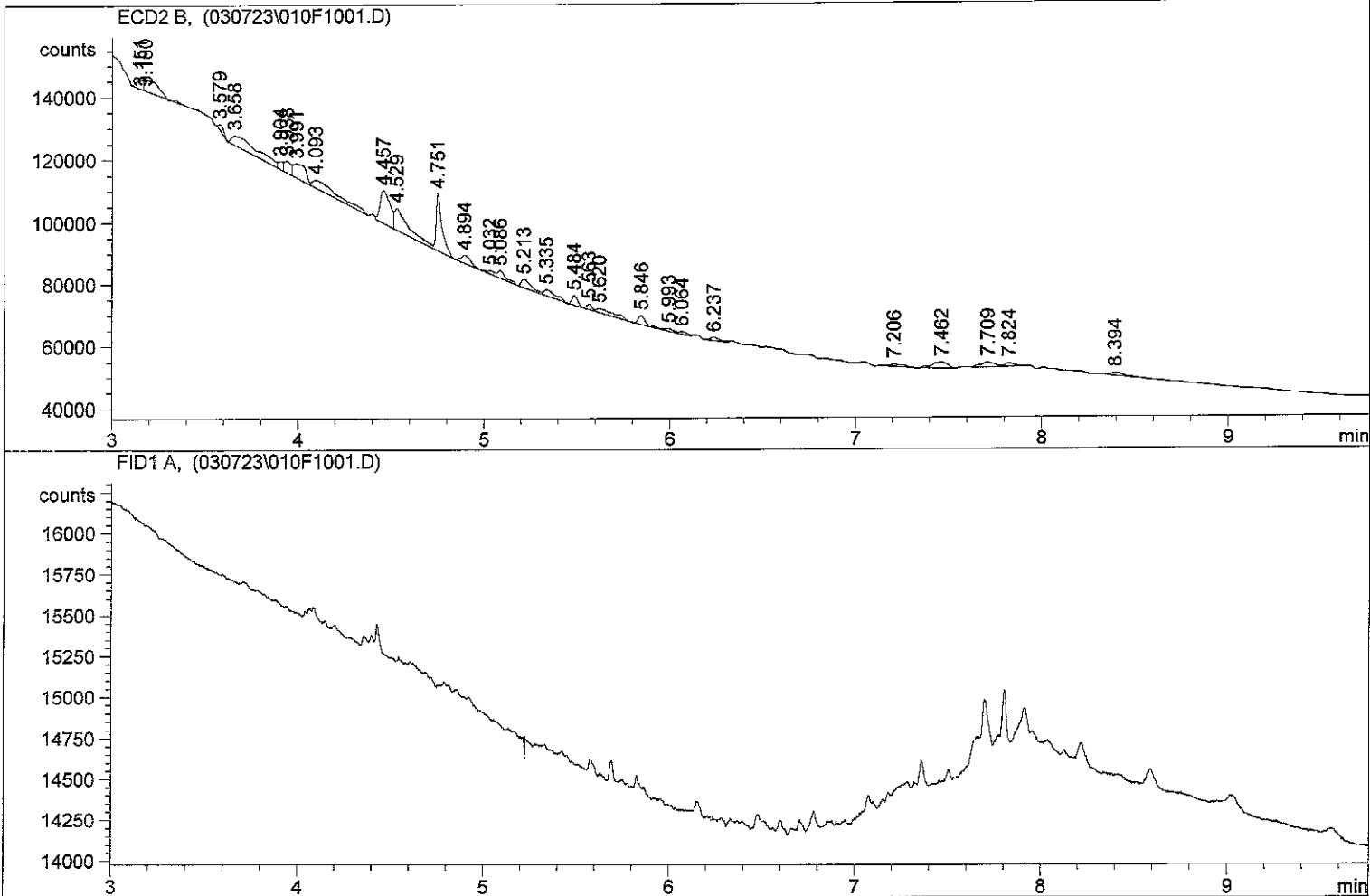
```
=====
Injection Date : 3/7/2023 5:51:06 PM      Seq. Line :    9
Sample Name    : 23C0108 05                Location  : Vial 9
Acq. Operator  : CR                       Inj       :    1
                                           Inj Volume: 1 µl

Sequence File  : C:\HPCHEM\1\SEQUENCE\030723.S
Method         : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed   : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====
```



*** End of Report ***

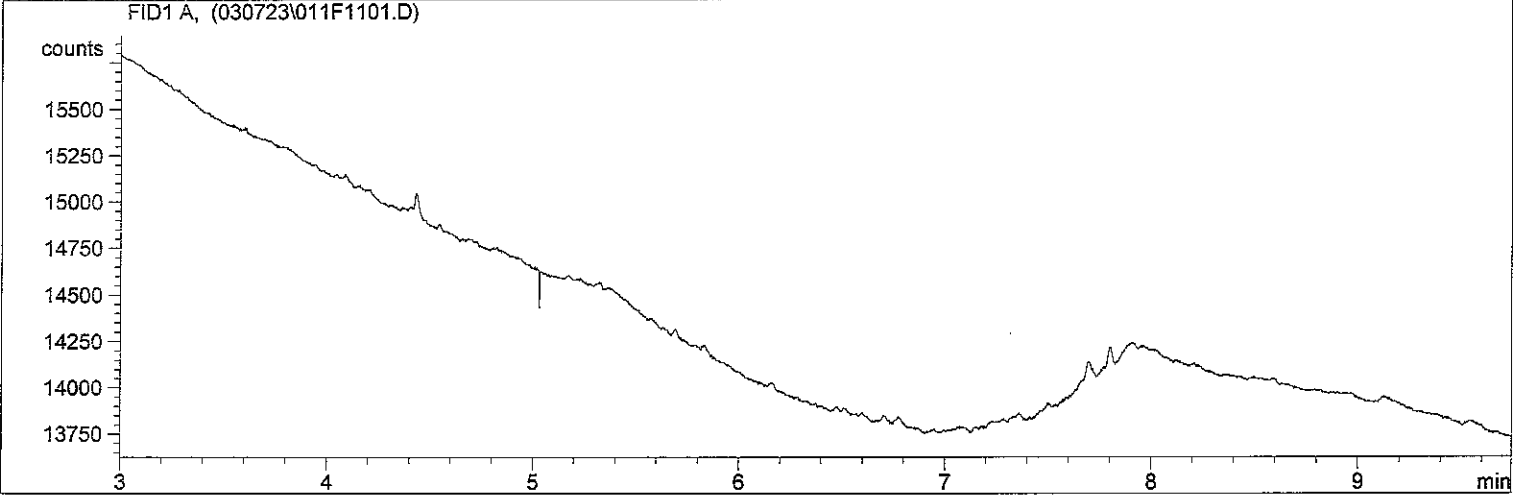
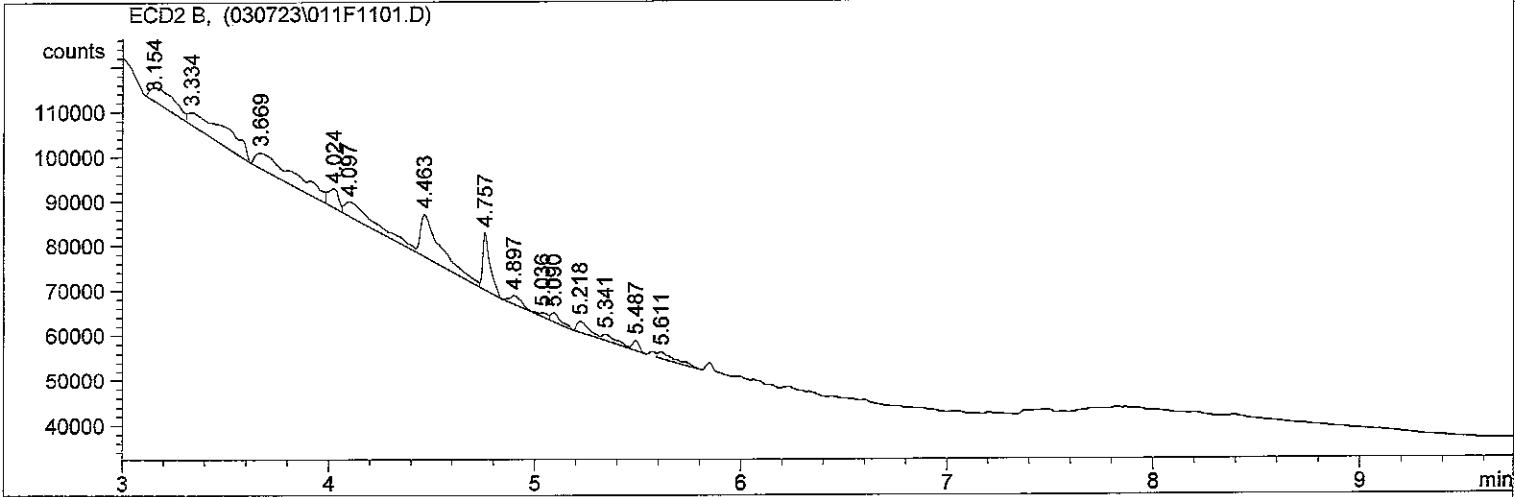
Injection Date : 3/7/2023 6:05:25 PM Seq. Line : 10
Sample Name : 23C0108 06 Location : Vial 10
Acq. Operator : CR Inj : 1
Inj Volume : 1 µl
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***

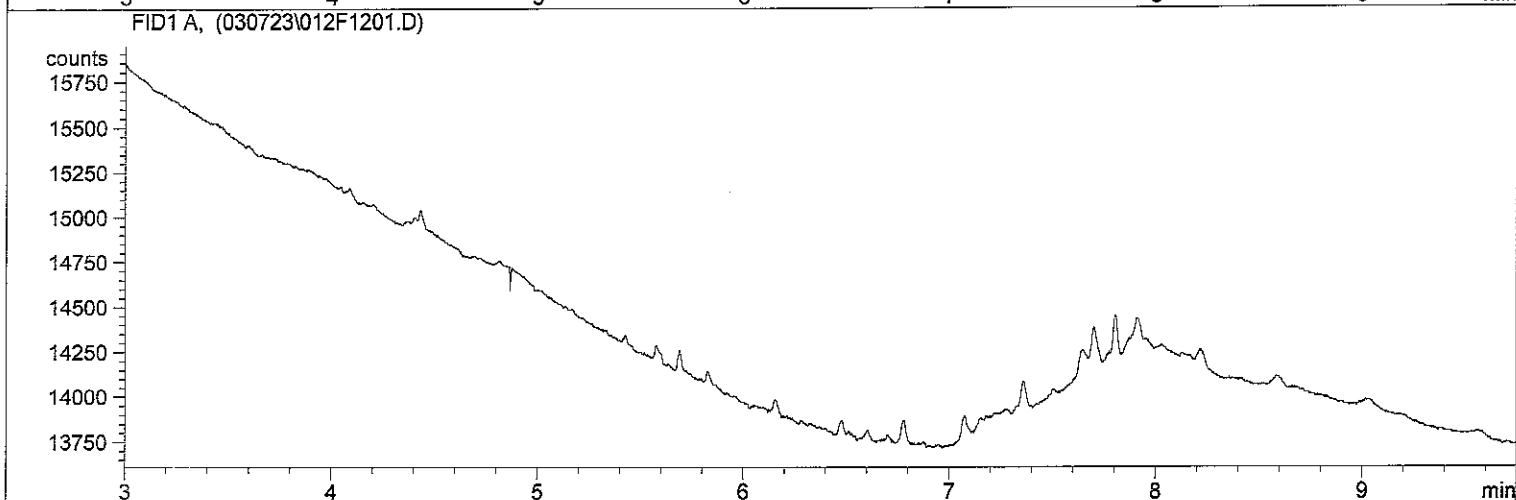
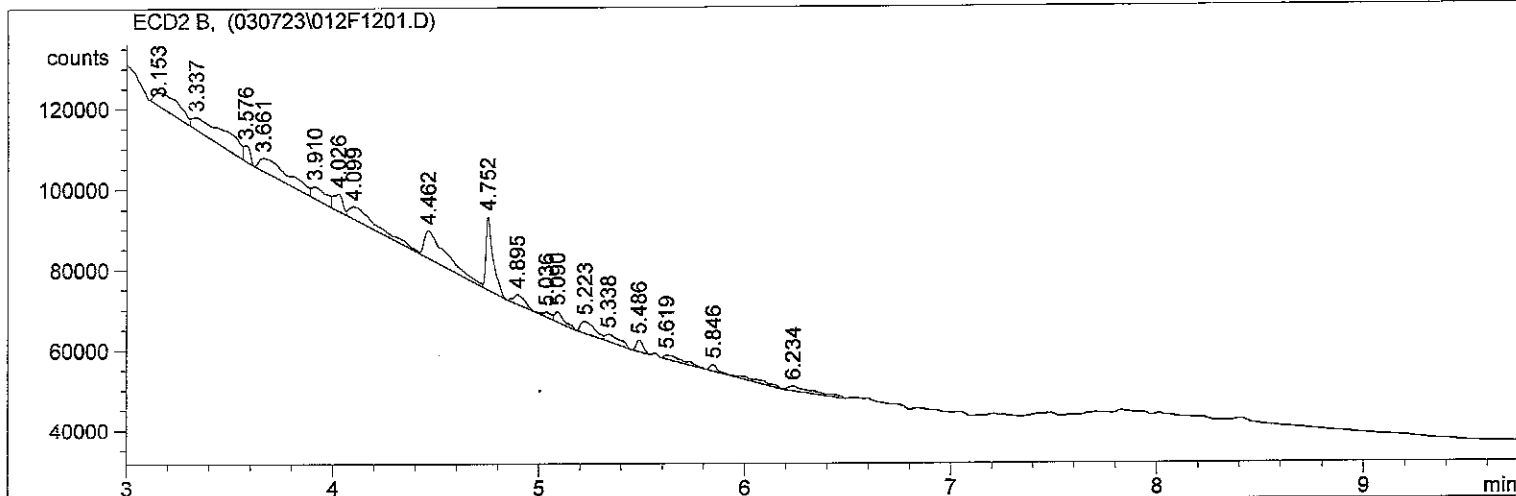
Injection Date : 3/7/2023 6:20:27 PM Seq. Line : 11
Sample Name : 23C0108 07 Location : Vial 11
Acq. Operator : CR Inj : 1
 Inj Volume : 1 µl

Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***

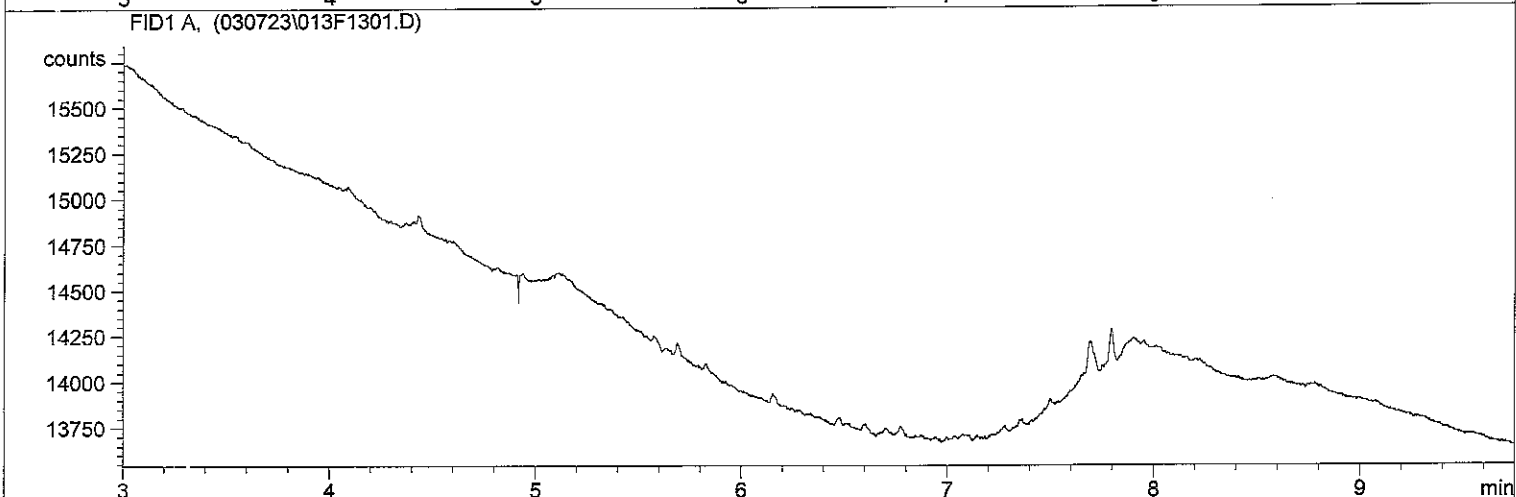
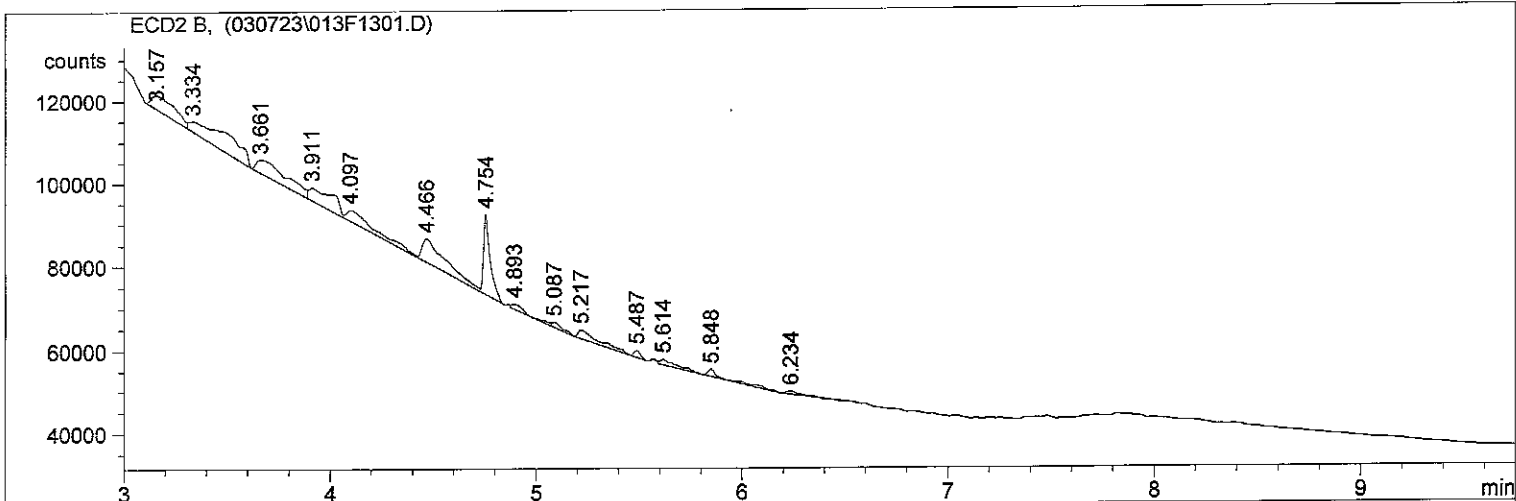
Injection Date : 3/7/2023 6:35:24 PM Seq. Line : 12
Sample Name : 23C0108 08 Location : Vial 12
Acq. Operator : CR Inj : 1
 Inj Volume : 1 µl
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***

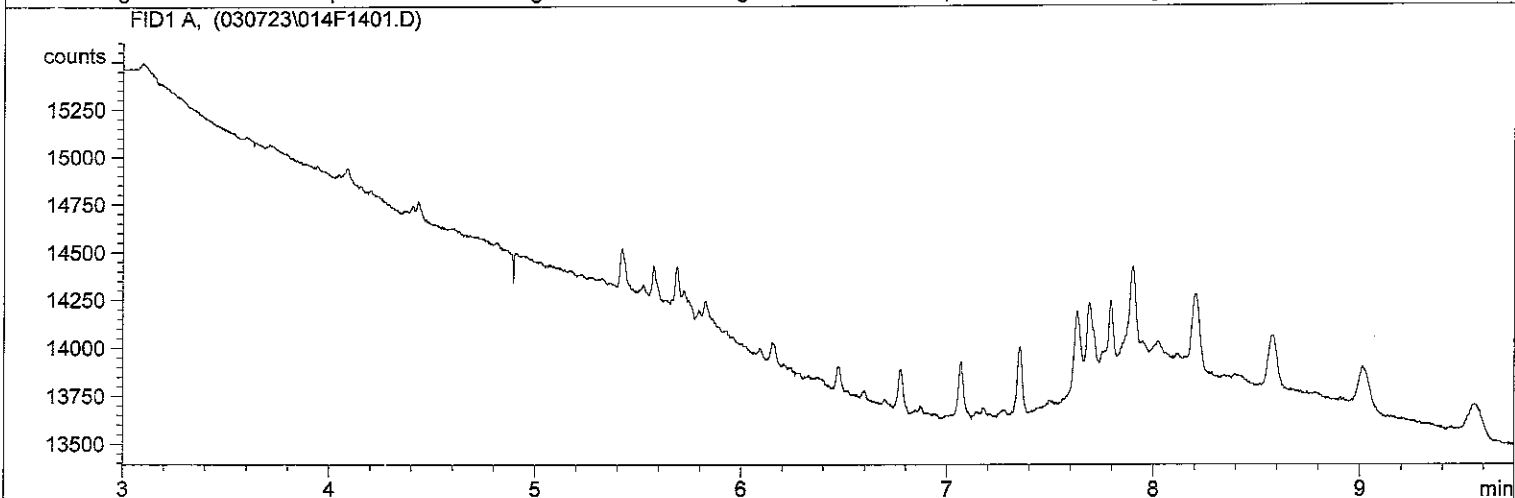
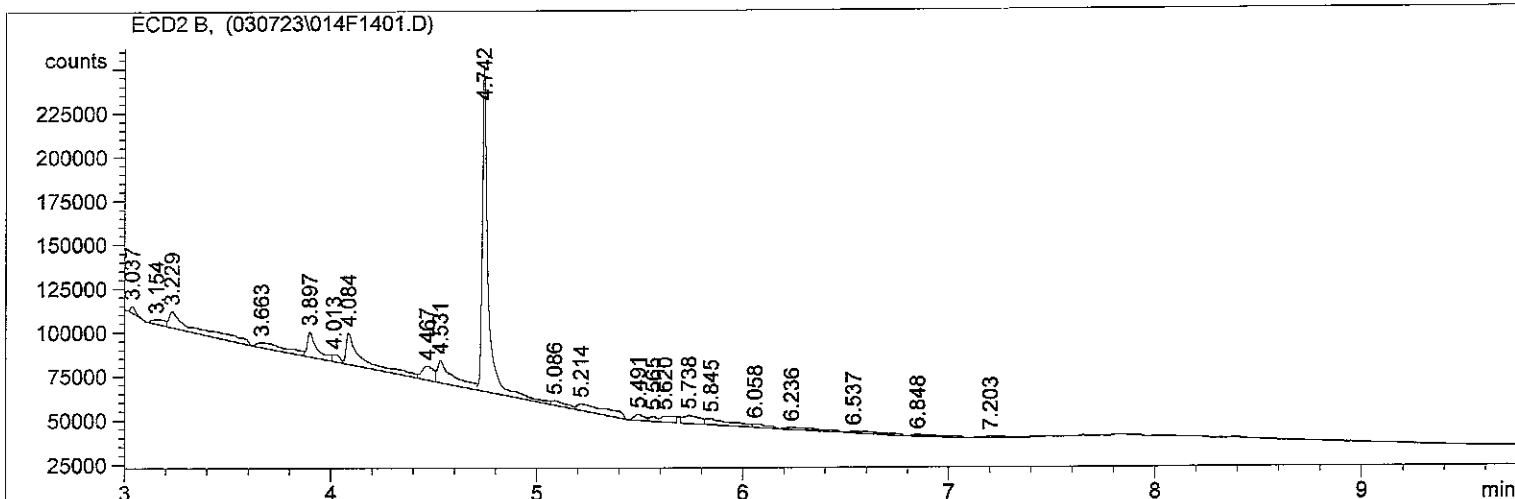
Injection Date : 3/7/2023 6:49:03 PM Seq. Line : 13
Sample Name : 23C0108 09 Location : Vial 13
Acq. Operator : CR Inj : 1
 Inj Volume : 1 µl

Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***

Injection Date : 3/7/2023 7:05:21 PM Seq. Line : 14
Sample Name : 23C0108 10 Location : Vial 14
Acq. Operator : CR Inj : 1
 Inj Volume : 1 µl
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***


```

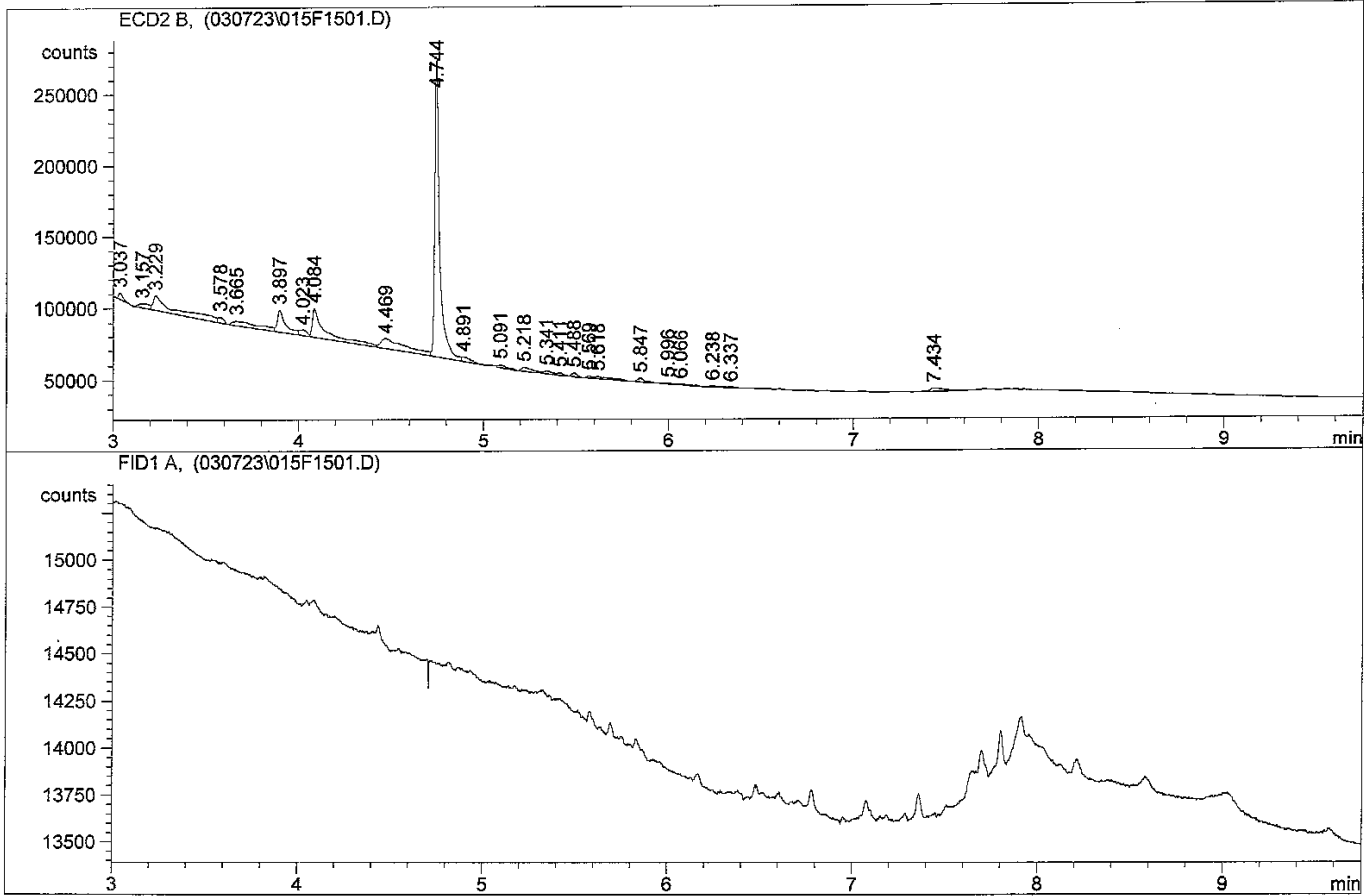
=====
Injection Date   : 3/7/2023 7:35:19 PM      Seq. Line : 16
Sample Name     : 23C0109 02                Location  : Vial 16
Acq. Operator   : CR                       Inj      : 1
                                           Inj Volume: 1 µl
Sequence File   : C:\HPCHEM\1\SEQUENCE\030723.S
=====

```

```

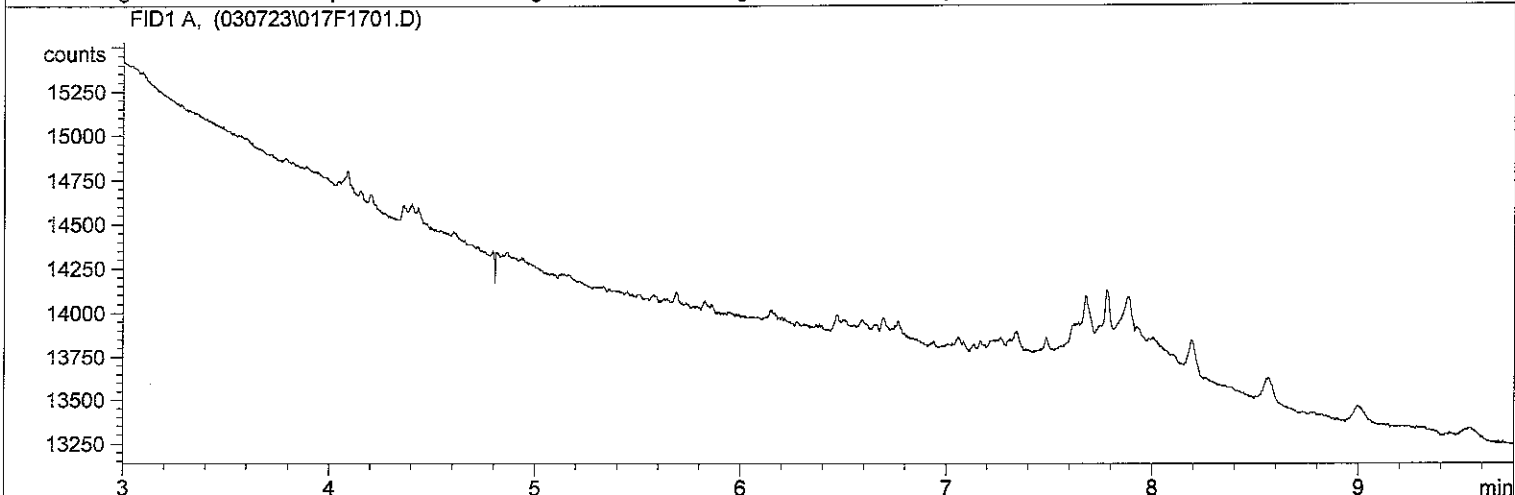
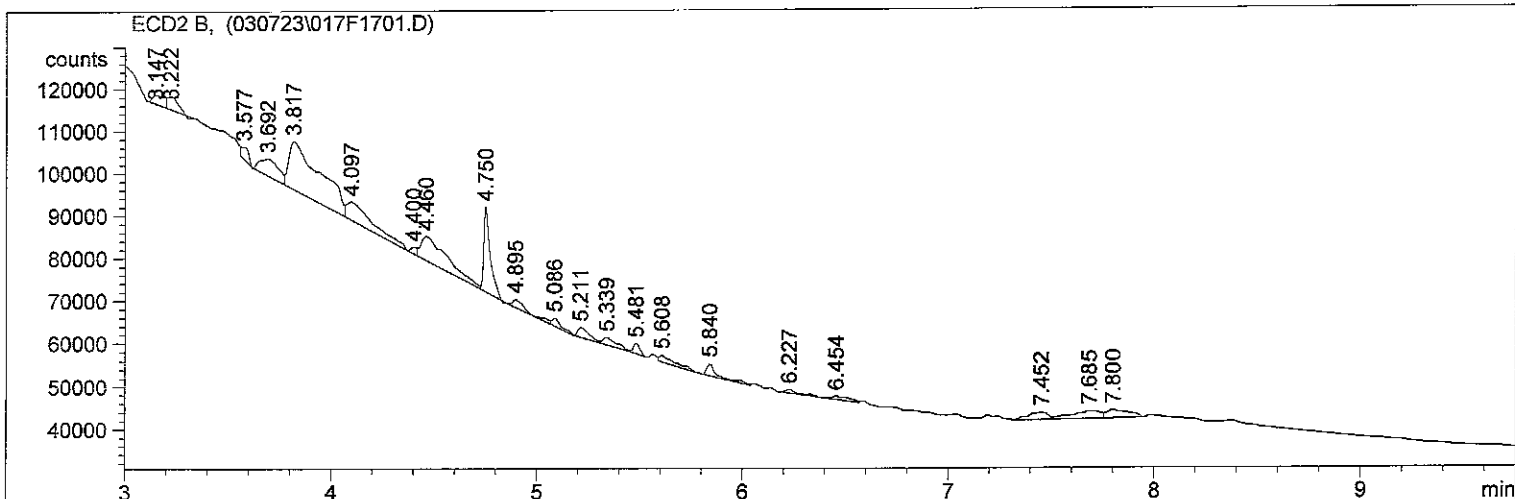
=====
Injection Date   : 3/7/2023 7:20:19 PM      Seq. Line : 15
Sample Name     : 23C0109 01                Location  : Vial 15
Acq. Operator   : CR                       Inj      : 1
                                           Inj Volume: 1 µl
Sequence File   : C:\HPCHEM\1\SEQUENCE\030723.S
Method         : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed    : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====

```



*** End of Report ***

Injection Date : 3/7/2023 7:50:16 PM Seq. Line : 17
Sample Name : 23C0109 03 Location : Vial 17
Acq. Operator : CR Inj : 1
Inj Volume : 1 µl
Sequence File : C:\HPCHEM\1\SEQUENCE\030723.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



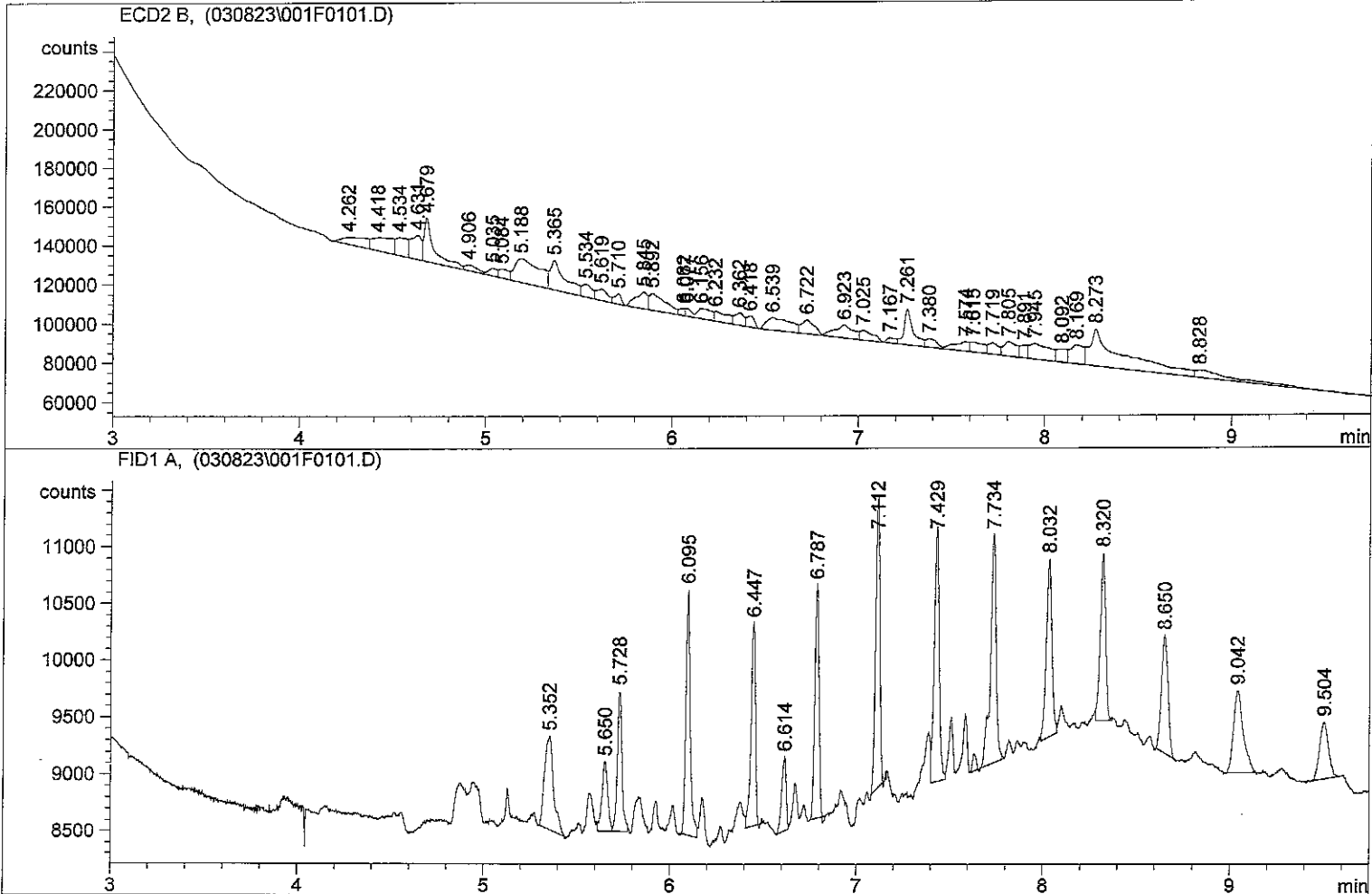
*** End of Report ***

```

Injection Date   : 3/8/2023 1:58:41 PM      Seq. Line   :    1
Sample Name     : DCM RINSE                  Location    : Vial 1
Acq. Operator   : YL                        Inj         :    1
                                           Inj Volume  : 1 µl

Sequence File   : C:\HPCHEM\1\SEQUENCE\030823.S
Method          : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed    : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD

```

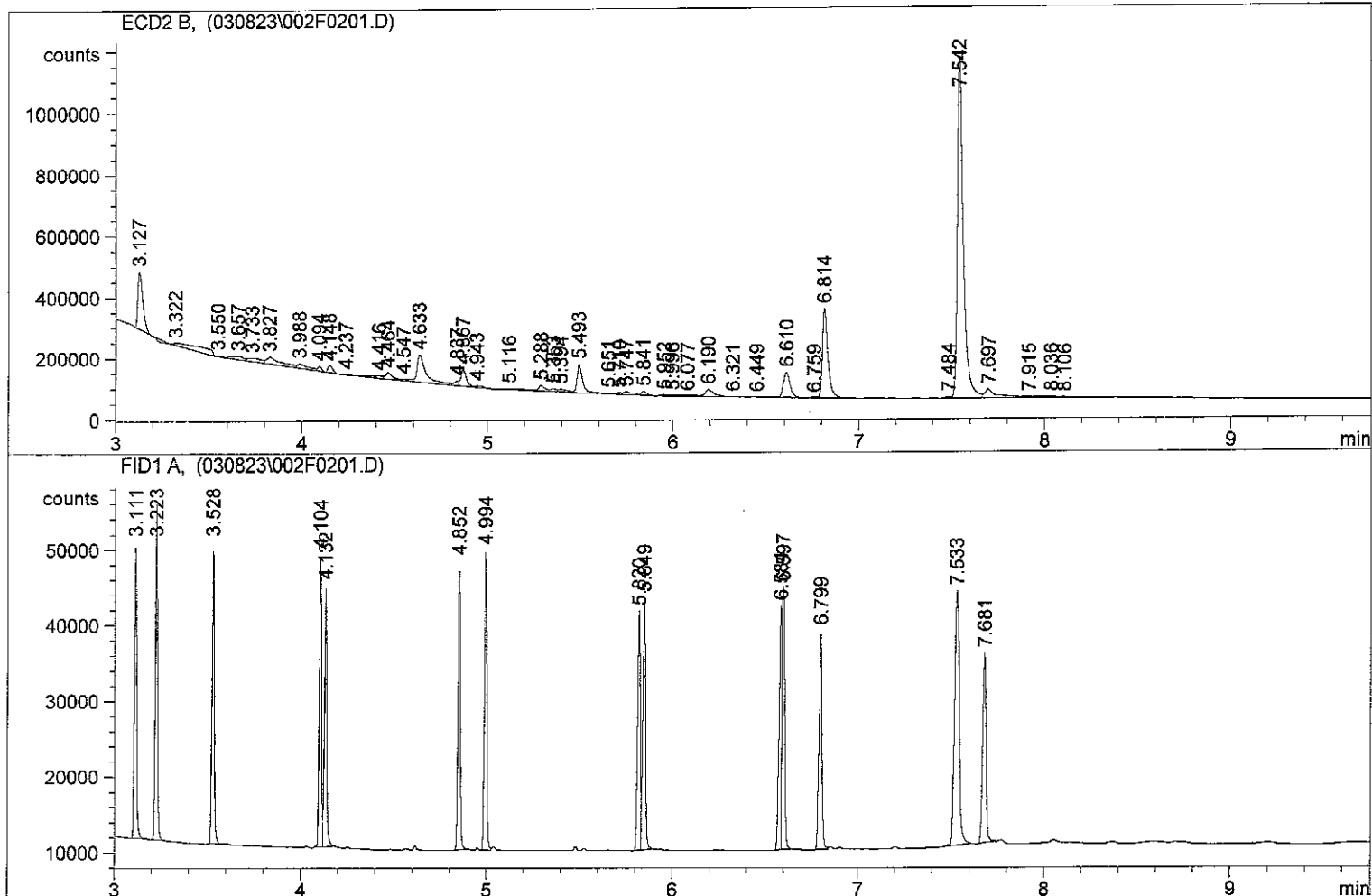


*** End of Report ***

```

=====
Injection Date   : 3/8/2023 2:10:01 PM      Seq. Line   :    2
Sample Name     : PNA STD 10PPM            Location    : Vial 2
Acq. Operator  : YL                        Inj         :    1
                                           Inj Volume  : 1 µl

Sequence File   : C:\HPCHEM\1\SEQUENCE\030823.S
Method          : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed   : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====
    
```



*** End of Report ***

```

=====
Injection Date   : 3/8/2023 2:42:15 PM      Seq. Line :    4
Sample Name     : 23C0107 02                Location  : Vial 4
Acq. Operator   : YL                        Inj       :    1
                                           Inj Volume: 1 µl

Sequence File   : C:\HPCHEM\1\SEQUENCE\030823.S
Method          : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed    : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====

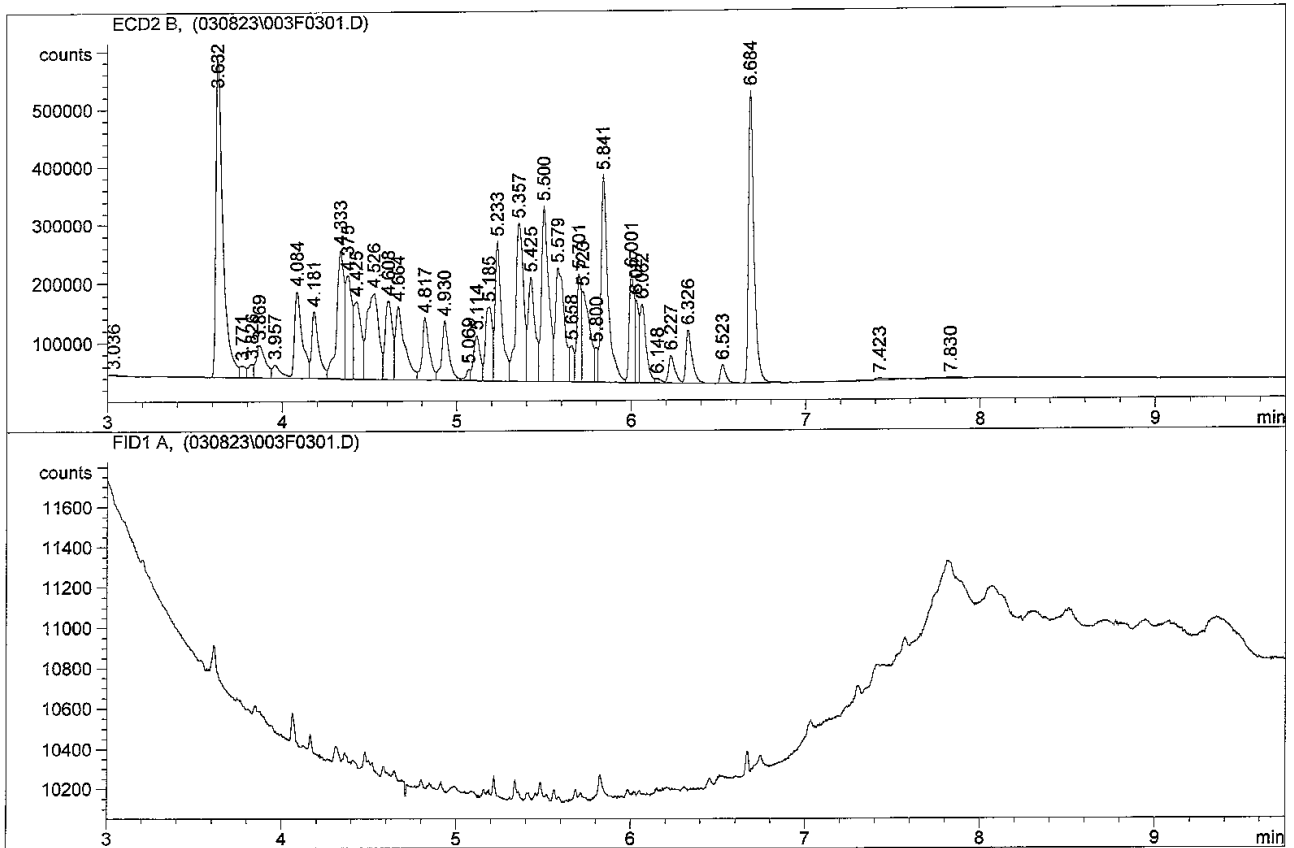
```

```

=====
Injection Date   : 3/8/2023 2:27:57 PM      Seq. Line :    3
Sample Name     : AR1660 1PPM                Location  : Vial 3
Acq. Operator   : YL                        Inj       :    1
                                           Inj Volume: 1 µl

Sequence File   : C:\HPCHEM\1\SEQUENCE\030823.S
Method          : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed    : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD
=====

```

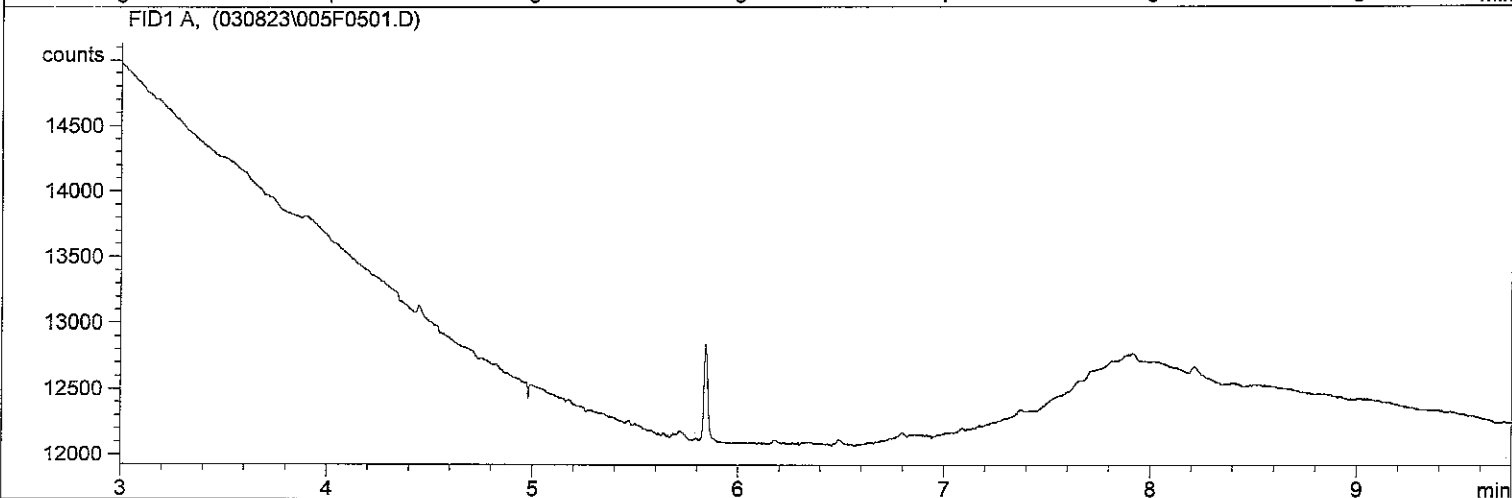
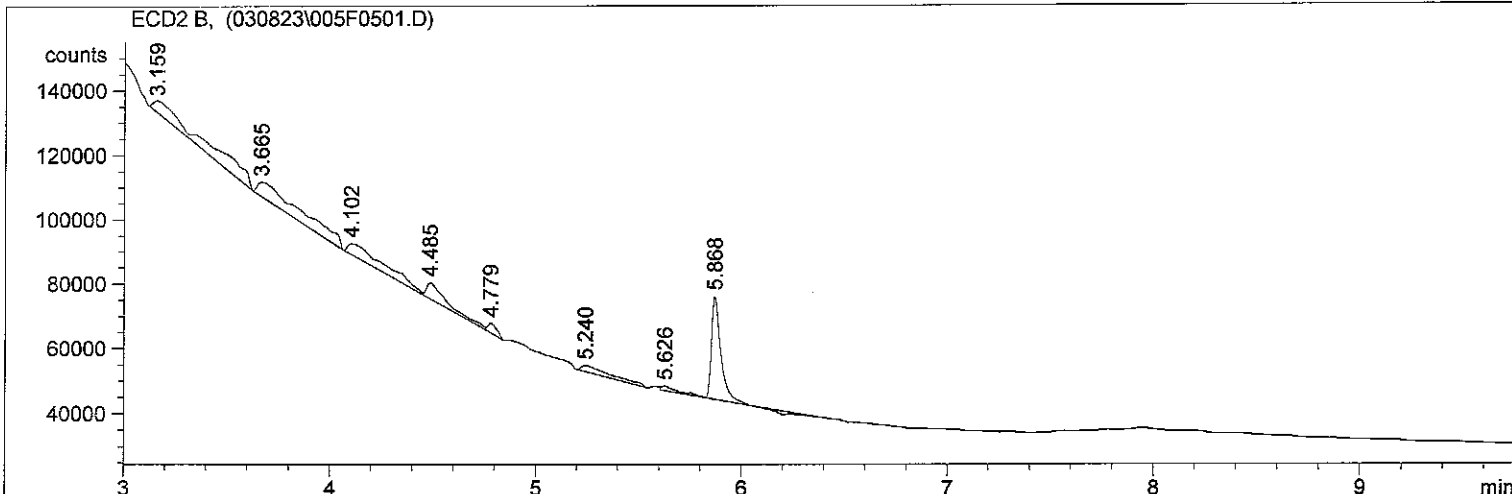


*** End of Report ***

Injection Date : 3/8/2023 2:57:30 PM
Sample Name : 23C0107 03
Acq. Operator : YL

Seq. Line : 5
Location : Vial 5
Inj : 1
Inj Volume : 1 µl

Sequence File : C:\HPCHEM\1\SEQUENCE\030823.S
Method : C:\HPCHEM\1\METHODS\SCREEN.M
Last changed : 7/9/2021 3:37:33 AM by TW
SCREEN METHOD



*** End of Report ***



CLEANUP BATCH SUMMARY

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Cleanup Batch: CLC0117

Cleanup Type: Silica Gel

Cleanup Method: EPA 3630C Silica Gel Cleanup - uL

Analysis: EPA 8082A

SAMPLE NAME	LAB SAMPLE ID	LAB FILE ID	DATE PREPARED	OBSERVATIONS
Matrix Spike	BLC0219-MS1	03152322ECD7.D	03/15/2023	
Matrix Spike Dup	BLC0219-MSD1	03152323ECD7.D	03/15/2023	
LDW21-IT608C	23C0107-02	03152320ECD7.D	03/15/2023	
LCS Dup	BLC0219-BSD1	03152317ECD7.D	03/15/2023	
LCS	BLC0219-BS1	03152316ECD7.D	03/15/2023	
Blank	BLC0219-BLK1	03152315ECD7.D	03/15/2023	
LDW22-SS826	23C0107-01	03152319ECD7.D	03/15/2023	
LDW21-IT608D	23C0107-03	03152321ECD7.D	03/15/2023	
Reference	BLC0219-SRM1	03152318ECD7.D	03/15/2023	



CLEANUP BENCH SHEET

CLC0117

Matrix: Solid

Cleanup using: Organics - EPA 3630C Silica Gel Cleanup - uL

Printed: 3/15/2023 11:26:42AM

Lab Number	Sample Container	Sample Name	Extract Container	Initial (uL)	Final (uL)	Analysis	Clean Up Date	Cleaned By	Cleanup Comments
23C0107-01	A	LDW22-SS826	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0107-02	A	LDW21-IT608C	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0107-03	A	LDW21-IT608D	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-01	A	LDW23-SC1037	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-02	A	LDW23-SC1044	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-03	A	LDW23-SC1107	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-04	A	LDW23-SC1106	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-05	A	LDW23-SC1118	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-06	A	LDW23-SS1106	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-07	A	LDW23-SS1107	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-08	A	LDW23-SS1111	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-09	A	LDW23-SS1118	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-10	A	LDW23-SC1111	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-01	A	LDW23-SC1104	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-02	A	LDW23-SS1104	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-03	A	LDW23-SS1105	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
BLC0219-BLK1	-	Blank	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-BS1	-	LCS	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-BSD1	-	LCS Dup	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-MS1	-	Matrix Spike	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-MSD1	-	Matrix Spike Dup	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-SRM1	-	Reference	-	2.5	2.5	-	3/15/2023	LMJ	



CLEANUP BENCH SHEET

CLC0117

Matrix: Solid

Cleanup using: Organics - EPA 3630C Silica Gel Cleanup - uL

Printed: 3/15/2023 11:26:42AM

Lab Number	Sample Container	Sample Name	Extract Container	Initial (uL)	Final (uL)	Analysis	Clean Up Date	Cleaned By	Cleanup Comments
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CLEANUP BATCH SUMMARY

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Cleanup Batch: CLC0118

Cleanup Type: Sulfur

Cleanup Method: EPA 3660B Sulfur Cleanup - uL

Analysis: EPA 8082A

SAMPLE NAME	LAB SAMPLE ID	LAB FILE ID	DATE PREPARED	OBSERVATIONS
LDW22-SS826	23C0107-01	03152319ECD7.D	03/15/2023	
LDW21-IT608D	23C0107-03	03152321ECD7.D	03/15/2023	
LDW21-IT608C	23C0107-02	03152320ECD7.D	03/15/2023	
Matrix Spike	BLC0219-MS1	03152322ECD7.D	03/15/2023	
Reference	BLC0219-SRM1	03152318ECD7.D	03/15/2023	
Matrix Spike Dup	BLC0219-MSD1	03152323ECD7.D	03/15/2023	
Blank	BLC0219-BLK1	03152315ECD7.D	03/15/2023	
LCS	BLC0219-BS1	03152316ECD7.D	03/15/2023	
LCS Dup	BLC0219-BSD1	03152317ECD7.D	03/15/2023	



CLEANUP BENCH SHEET

CLC0118

Matrix: Solid

Cleanup using: Organics - EPA 3660B Sulfur Cleanup - uL

Printed: 3/15/2023 11:27:26AM

Lab Number	Sample Container	Sample Name	Extract Container	Initial (uL)	Final (uL)	Analysis	Clean Up Date	Cleaned By	Cleanup Comments
23C0107-01	A	LDW22-SS826	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0107-02	A	LDW21-IT608C	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0107-03	A	LDW21-IT608D	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-01	A	LDW23-SC1037	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-02	A	LDW23-SC1044	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-03	A	LDW23-SC1107	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-04	A	LDW23-SC1106	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-05	A	LDW23-SC1118	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-06	A	LDW23-SS1106	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-07	A	LDW23-SS1107	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-08	A	LDW23-SS1111	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-09	A	LDW23-SS1118	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-10	A	LDW23-SC1111	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-01	A	LDW23-SC1104	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-02	A	LDW23-SS1104	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-03	A	LDW23-SS1105	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
BLC0219-BLK1	-	Blank	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-BS1	-	LCS	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-BSD1	-	LCS Dup	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-MS1	-	Matrix Spike	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-MSD1	-	Matrix Spike Dup	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-SRM1	-	Reference	-	2.5	2.5	-	3/15/2023	LMJ	



CLEANUP BENCH SHEET

CLC0118

Matrix: Solid

Cleanup using: Organics - EPA 3660B Sulfur Cleanup - uL

Printed: 3/15/2023 11:27:26AM

Lab Number	Sample Container	Sample Name	Extract Container	Initial (uL)	Final (uL)	Analysis	Clean Up Date	Cleaned By	Cleanup Comments
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CLEANUP BATCH SUMMARY

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Cleanup Batch: CLC0119

Cleanup Type: Sulfuric Acid

Cleanup Method: EPA 3665 Sulfuric Acid Cleanup - uL

Analysis: EPA 8082A

SAMPLE NAME	LAB SAMPLE ID	LAB FILE ID	DATE PREPARED	OBSERVATIONS
LDW22-SS826	23C0107-01	03152319ECD7.D	03/15/2023	
Reference	BLC0219-SRM1	03152318ECD7.D	03/15/2023	
Blank	BLC0219-BLK1	03152315ECD7.D	03/15/2023	
LCS Dup	BLC0219-BSD1	03152317ECD7.D	03/15/2023	
LDW21-IT608D	23C0107-03	03152321ECD7.D	03/15/2023	
LDW21-IT608C	23C0107-02	03152320ECD7.D	03/15/2023	
Matrix Spike	BLC0219-MS1	03152322ECD7.D	03/15/2023	
Matrix Spike Dup	BLC0219-MSD1	03152323ECD7.D	03/15/2023	
LCS	BLC0219-BS1	03152316ECD7.D	03/15/2023	



CLEANUP BENCH SHEET

CLC0119

Matrix: Solid

Cleanup using: Organics - EPA 3665 Sulfuric Acid Cleanup - uL

Printed: 3/15/2023 11:28:07AM

Lab Number	Sample Container	Sample Name	Extract Container	Initial (uL)	Final (uL)	Analysis	Clean Up Date	Cleaned By	Cleanup Comments
23C0107-01	A	LDW22-SS826	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0107-02	A	LDW21-IT608C	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0107-03	A	LDW21-IT608D	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-01	A	LDW23-SC1037	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-02	A	LDW23-SC1044	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-03	A	LDW23-SC1107	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-04	A	LDW23-SC1106	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-05	A	LDW23-SC1118	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-06	A	LDW23-SS1106	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-07	A	LDW23-SS1107	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-08	A	LDW23-SS1111	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-09	A	LDW23-SS1118	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0108-10	A	LDW23-SC1111	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-01	A	LDW23-SC1104	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-02	A	LDW23-SS1104	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
23C0109-03	A	LDW23-SS1105	A 05	2.5	2.5	8082A PCB Solid 4	3/15/2023	LMJ	
BLC0219-BLK1	-	Blank	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-BS1	-	LCS	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-BSD1	-	LCS Dup	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-MS1	-	Matrix Spike	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-MSD1	-	Matrix Spike Dup	-	2.5	2.5	-	3/15/2023	LMJ	
BLC0219-SRM1	-	Reference	-	2.5	2.5	-	3/15/2023	LMJ	



CLEANUP BENCH SHEET

CLC0119

Matrix: Solid

Cleanup using: Organics - EPA 3665 Sulfuric Acid Cleanup - uL

Printed: 3/15/2023 11:28:07AM

Lab Number	Sample Container	Sample Name	Extract Container	Initial (uL)	Final (uL)	Analysis	Clean Up Date	Cleaned By	Cleanup Comments
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Form I
METHOD BLANK DATA SHEET
EPA 8082A

Blank

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Matrix:	<u>Solid</u>	Laboratory ID:	<u>BLC0219-BLK1</u>
Sampled:	<u>N/A</u>	Prepared:	<u>03/10/23 11:58</u>
Solids:		Preparation:	<u>EPA 3546 (Microwave)</u>
Batch:	<u>BLC0219</u>	Sequence:	<u>SLC0215</u>
Instrument:	<u>ECD7</u>	Column:	<u>ZB5</u>
		File ID:	<u>03152315ECD7.D</u>
		Analyzed:	<u>03/15/23 17:51</u>
		Initial/Final:	<u>12.5 g / 2.5 mL</u>
		Calibration:	<u>GB00069</u>
		Cleanups:	<u>Silica Gel, Sulfur, Sulfuric Acid</u>

CAS NO.	COMPOUND	DILUTION	CONC: (ug/kg wet)	Q	DL	RL
12674-11-2	Aroclor 1016	1	4.0	U	1.6	4.0
11104-28-2	Aroclor 1221	1	4.0	U	1.6	4.0
11141-16-5	Aroclor 1232	1	4.0	U	1.6	4.0
53469-21-9	Aroclor 1242	1	4.0	U	1.6	4.0
12672-29-6	Aroclor 1248	1	4.0	U	1.6	4.0
11097-69-1	Aroclor 1254	1	4.0	U	1.6	4.0
11096-82-5	Aroclor 1260	1	4.0	U	0.6	4.0

SURROGATES	ADDED: (ug/kg wet)	FOUND: (ug/kg wet)	% REC	QC LIMITS	Q
Decachlorobiphenyl	8.0000	5.40	67.5	40 - 126	Q
Tetrachlorometaxylene	8.0000	6.11	76.4	44 - 120	
Decachlorobiphenyl [2C]	8.0000	7.57	94.6	40 - 126	
Tetrachlorometaxylene [2C]	8.0000	6.09	76.1	44 - 120	

[2C] indicates second-column analyte, present if quantification on any batch samples used second column data.

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152315ECD7.D
Data file 2: /230315.b/230315.b/03152315ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: BLC0219-BLK1
Client ID:
Injection Date: 15-MAR-2023 17:51
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	-0.000	522873	5.693	0.000	178764	30.6	30.5	0.3	Tetrachloro-m-xylene
13.895	-0.002	913292	14.120	-0.001	364285	27.0	37.8	33.4	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1146280	70.1
Hexabromobiphenyl	1429847	3433528	140.1 <-

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	400044	26.9
Hexabromobiphenyl	513946	632392	23.0

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	---			0.0	1	---			0.0
Aroclor-1016	2	---			0.0	2	---			0.0
Aroclor-1016	3	---			0.0	3	---			0.0
Aroclor-1016	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	---			0.0	2	---			0.0
Aroclor-1221	3	---			0.0	3	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	---			0.0	2	---			0.0
Aroclor-1232	3	---			0.0	3	---			0.0
Aroclor-1232	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1242	1	---			0.0	1	---			0.0
Aroclor-1242	2	---			0.0	2	---			0.0
Aroclor-1242	3	---			0.0	3	---			0.0
Aroclor-1242	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1248	1	---			0.0	1	---			0.0
Aroclor-1248	2	---			0.0	2	---			0.0
Aroclor-1248	3	---			0.0	3	---			0.0
Aroclor-1248	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1254	1	---			0.0	1	---			0.0
Aroclor-1254	2	---			0.0	2	---			0.0
Aroclor-1254	3	---			0.0	3	---			0.0
Aroclor-1254	4	---			0.0	4	---			0.0
Aroclor-1254	5	---			0.0	5	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1260	1	---			0.0	1	---			0.0
Aroclor-1260	2	---			0.0	2	---			0.0
Aroclor-1260	3	---			0.0	3	---			0.0
Aroclor-1260	4	---			0.0	4	---			0.0
Aroclor-1260	5	---			0.0	NS	---			----
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1262	1	---			0.0	1	---			0.0
Aroclor-1262	2	---			0.0	2	---			0.0
Aroclor-1262	3	---			0.0	3	---			0.0
Aroclor-1262	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1268	1	---			0.0	1	---			0.0
Aroclor-1268	2	---			0.0	2	---			0.0
Aroclor-1268	3	---			0.0	3	---			0.0
Aroclor-1268	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				

Total PCB Area Coll (5.911 - 13.797) = 460481

Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 42186 Col2 Total PCB = 0.0 ppm*

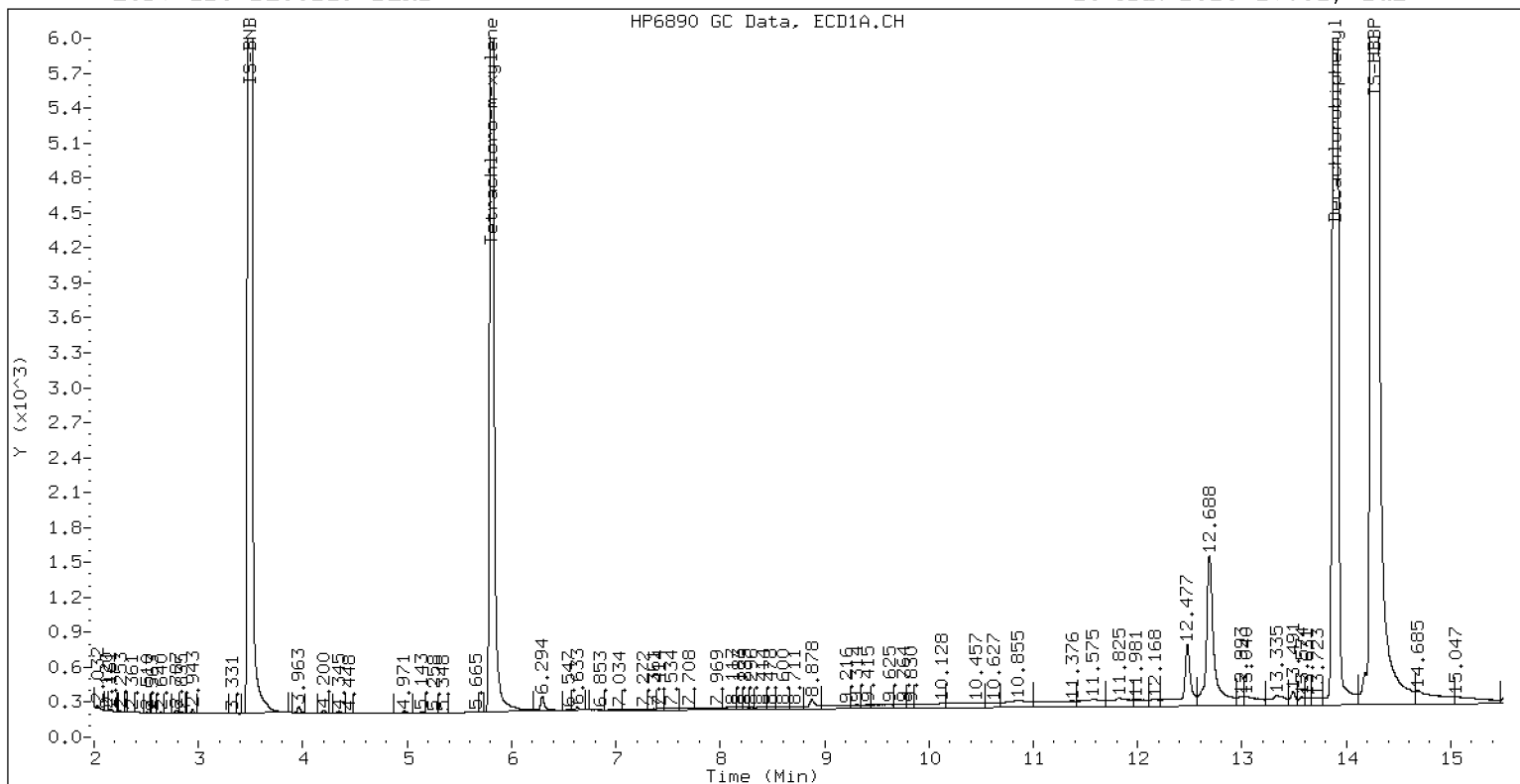
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 BLC0219-BLK1

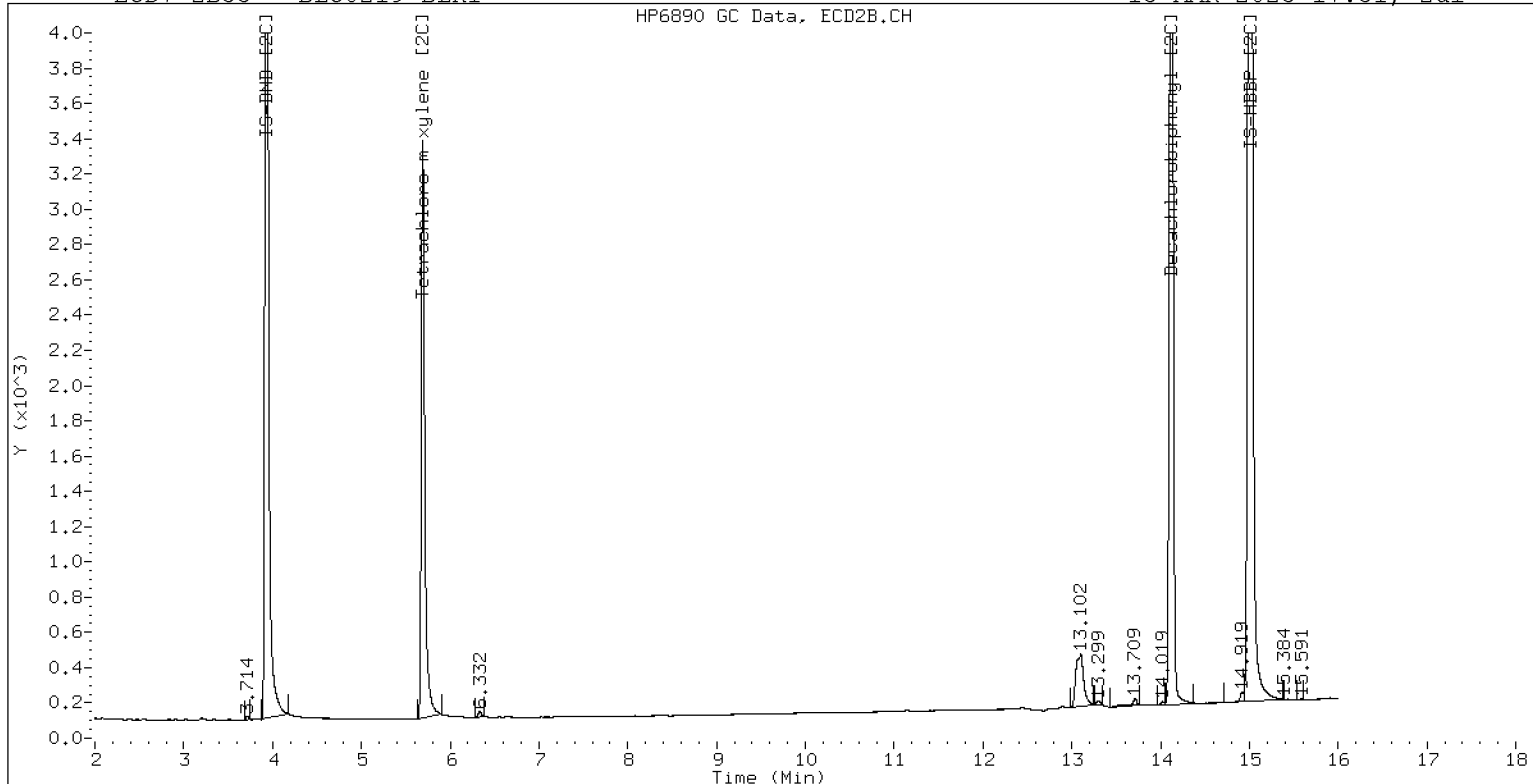
15-MAR-2023 17:51, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 BLC0219-BLK1

15-MAR-2023 17:51, 2u1



ZB-35 Manual Integration: NO



LCS / LCS DUPLICATE RECOVERY
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Analyzed: 03/15/23 18:12

Batch: BLC0219

Laboratory ID: BLC0219-BS1

Preparation: EPA 3546 (Microwave)

Sequence Name: LCS

Initial/Final: 12.5 g / 2.5 mL

COMPOUND	SPIKE ADDED (ug/kg wet)	LCS CONCENTRATION (ug/kg wet)	Q	LCS % REC. #	QC LIMITS REC.
Aroclor 1016	101	77.5		76.9	56 - 120
Aroclor 1260 [2C]	101	84.1		83.4	58 - 120

* Indicates values outside of QC limits

COMPOUND	SPIKE ADDED (ug/kg wet)	LCSD CONCENTRATION (ug/kg wet)	Q	LCSD % REC. #	% RPD #	QC LIMITS	
						RPD	REC.
Aroclor 1016	101	75.1		74.5	3.16	30	56 - 120
Aroclor 1260 [2C]	101	80.2		79.5	4.80	30	58 - 120

* Indicates values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152316ECD7.D
Data file 2: /230315.b/230315.b/03152316ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: BLC0219-BS1
Client ID:
Injection Date: 15-MAR-2023 18:12
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	ZB5 Col Response	RT	ZB35 Col Shift	ZB35 Col Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.810	-0.001	505228	5.692	0.000	169498	28.8	28.4	1.4	Tetrachloro-m-xylene
13.895	-0.002	820992	14.121	-0.000	345231	25.4	35.4	33.0	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1175579	74.5
Hexabromobiphenyl	1429847	3282396	129.6 <-

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	406961	29.1
Hexabromobiphenyl	513946	639731	24.5

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.272	0.001	165452	370.6	1	7.258	-0.001	84544	354.9
Aroclor-1016	2	7.659	0.004	542962	398.9	2	7.865	-0.007	193216	399.9
Aroclor-1016	3	7.795	0.005	256696	386.3	3	8.062	-0.007	82957	380.2
Aroclor-1016	4	8.408	0.003	169765	395.2	4	8.310	-0.003	65082	380.2
Total CollAve (4 peaks):				387.7		Total Col2Ave (4 peaks):				378.8 RPD = 2
Corrected Ave (3 peaks):				384.0		Corrected Ave (3 peaks):				371.8 RPD = 3
Aroclor-1221	1	4.732	0.001	1056	10.0	1	---			0.0
Aroclor-1221	2	6.133	0.001	20438	108.5	2	6.303	0.006	7640	104.8
Aroclor-1221	3	6.385	0.003	101771	232.8	3	6.627	0.005	37823	318.7
Total CollAve (3 peaks):				117.1		Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	4.732	0.002	1056	16.8	1	---			0.0
Aroclor-1232	2	6.133	0.002	20438	163.6	2	7.258	0.004	84544	817.1
Aroclor-1232	3	7.659	0.003	542962	960.3	3	7.865	0.005	193216	933.9
Aroclor-1232	4	8.583	0.002	221624	922.2	4	8.717	0.002	60876	1022.4
Total CollAve (4 peaks):				515.7		Total Col2Ave (3 peaks):				924.5 RPD = 57*
Corrected Ave (3 peaks):				367.6		Corrected Ave: < 3 Peaks				
Aroclor-1242	1	7.272	0.001	165452	454.1	1	7.258	-0.000	84544	447.1
Aroclor-1242	2	7.659	0.003	542962	490.8	2	7.865	-0.007	193216	486.1
Aroclor-1242	3	8.408	0.003	169765	493.2	3	9.172	-0.012	12712	102.8
Aroclor-1242	4	8.583	0.004	221624	435.5	4	9.596	-0.019	4847	32.2
Total CollAve (4 peaks):				468.4		Total Col2Ave (4 peaks):				267.1 RPD = 55*
Corrected Ave (3 peaks):				460.1		Corrected Ave (3 peaks):				194.0 RPD = 81*
Aroclor-1248	1	8.408	-0.003	169765	296.0	1	8.310	-0.004	65082	334.9
Aroclor-1248	2	8.583	-0.006	221624	304.0	2	8.717	-0.004	60876	303.0
Aroclor-1248	3	8.996	-0.003	249587	181.5	3	9.172	-0.012	12712	55.0
Aroclor-1248	4	9.304	0.002	204400	291.9	4	9.596	-0.016	4847	17.5
Total CollAve (4 peaks):				268.3		Total Col2Ave (4 peaks):				177.6 RPD = 41*
Corrected Ave (3 peaks):				256.4		Corrected Ave (3 peaks):				125.2 RPD = 69*
Aroclor-1254	1	9.304	0.005	204400	173.2	1	9.455	-0.003	57853	187.0
Aroclor-1254	2	---			0.0	2	9.977	-0.001	12977	52.2
Aroclor-1254	3	9.672	0.004	36986	48.7	3	10.150	0.016	121936	226.5
Aroclor-1254	4	9.811	0.003	110263	74.7	4	10.374	-0.008	158759	302.5
Aroclor-1254	5	10.124	-0.053	532040	575.2	5	10.572	-0.006	214656	671.7
Total CollAve (4 peaks):				218.0		Total Col2Ave (5 peaks):				288.0 RPD = 28
Corrected Ave (3 peaks):				98.9		Corrected Ave (4 peaks):				192.0 RPD = 64*
Aroclor-1260	1	11.046	0.002	445965	377.7	1	11.656	-0.002	163908	435.7
Aroclor-1260	2	11.366	0.005	472295	382.8	2	11.922	-0.003	419560	437.0
Aroclor-1260	3	11.740	0.006	1247475	381.2	3	12.438	-0.002	104052	408.4
Aroclor-1260	4	12.145	0.005	630448	382.6	4	12.506	-0.002	259564	401.1
Aroclor-1260	5	12.248	0.003	262882	370.6	NS	---			----
Total CollAve (5 peaks):				379.0		Total Col2Ave (4 peaks):				420.6 RPD = 10
Corrected Ave (4 peaks):				378.0		Corrected Ave (3 peaks):				415.1 RPD = 9
Aroclor-1262	1	10.832	0.003	874128	868.1	1	11.202	0.002	155807	285.2
Aroclor-1262	2	12.248	0.004	262882	160.4	2	11.656	0.005	163908	352.3
Aroclor-1262	3	12.322	0.004	317745	180.4	3	12.438	0.005	104052	197.1
Aroclor-1262	4	12.991	0.003	282892	175.7	4	12.506	0.004	259564	313.9
Total CollAve (4 peaks):				346.2		Total Col2Ave (4 peaks):				287.1 RPD = 19
Corrected Ave (3 peaks):				172.2		Corrected Ave (3 peaks):				265.4 RPD = 43*
Aroclor-1268	1	12.248	0.001	262882	62.5	1	12.438	0.006	104052	80.8
Aroclor-1268	2	12.322	0.005	317745	76.3	2	12.506	0.006	259564	187.4
Aroclor-1268	3	12.724	0.024	295205	82.8	3	12.894	0.002	8197	6.9
Aroclor-1268	4	13.489	-0.001	58733	5.0	4	13.710	0.001	32458	8.6
Total CollAve (4 peaks):				56.7		Total Col2Ave (4 peaks):				70.9 RPD = 22
Corrected Ave (3 peaks):				47.9		Corrected Ave (3 peaks):				32.1 RPD = 40

Total PCB Area Col1 (5.911 - 13.797) = 12127048 Col1 Total PCB = 0.9 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 3866444 Col2 Total PCB = 0.8 ppm*

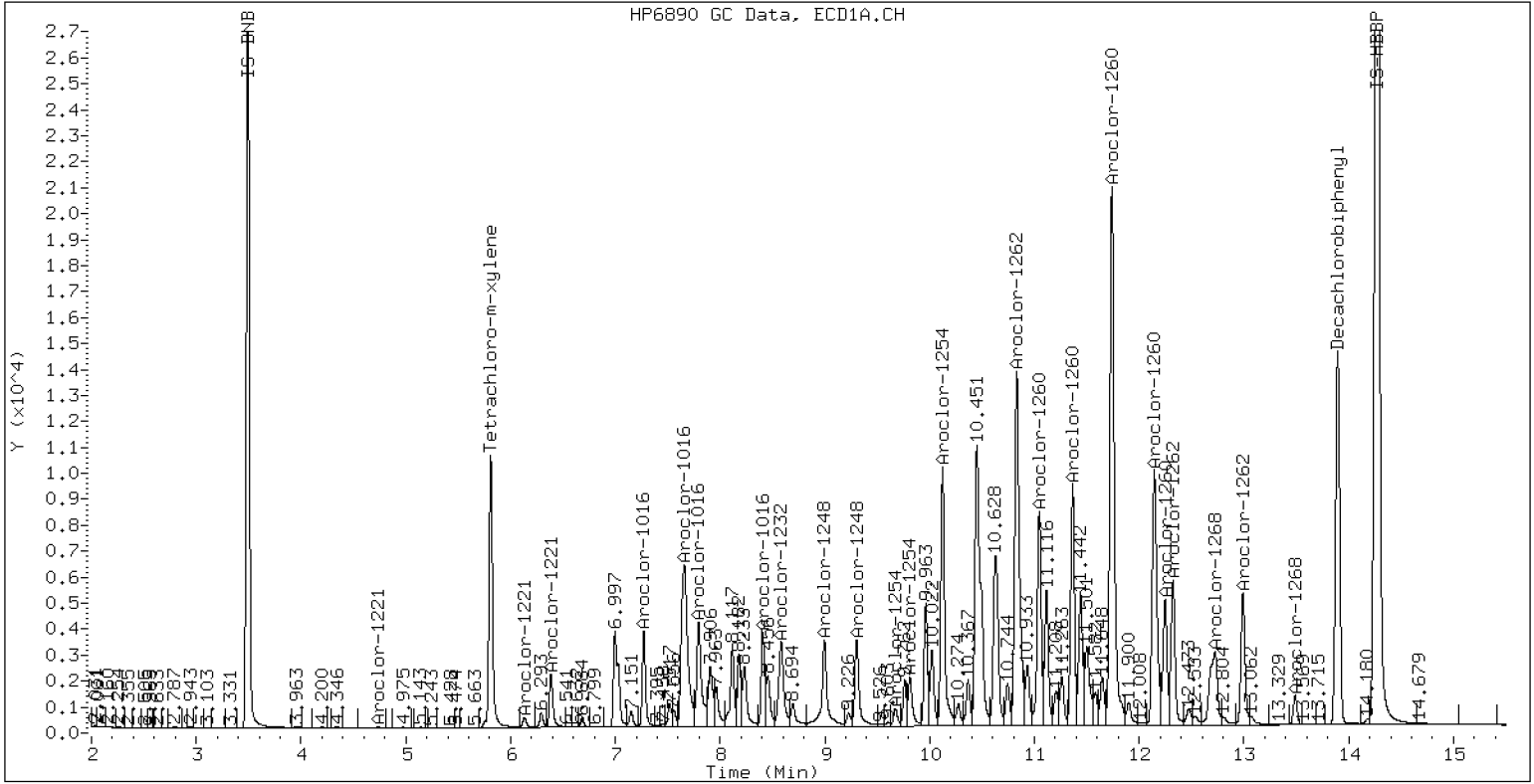
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 BLC0219-BS1

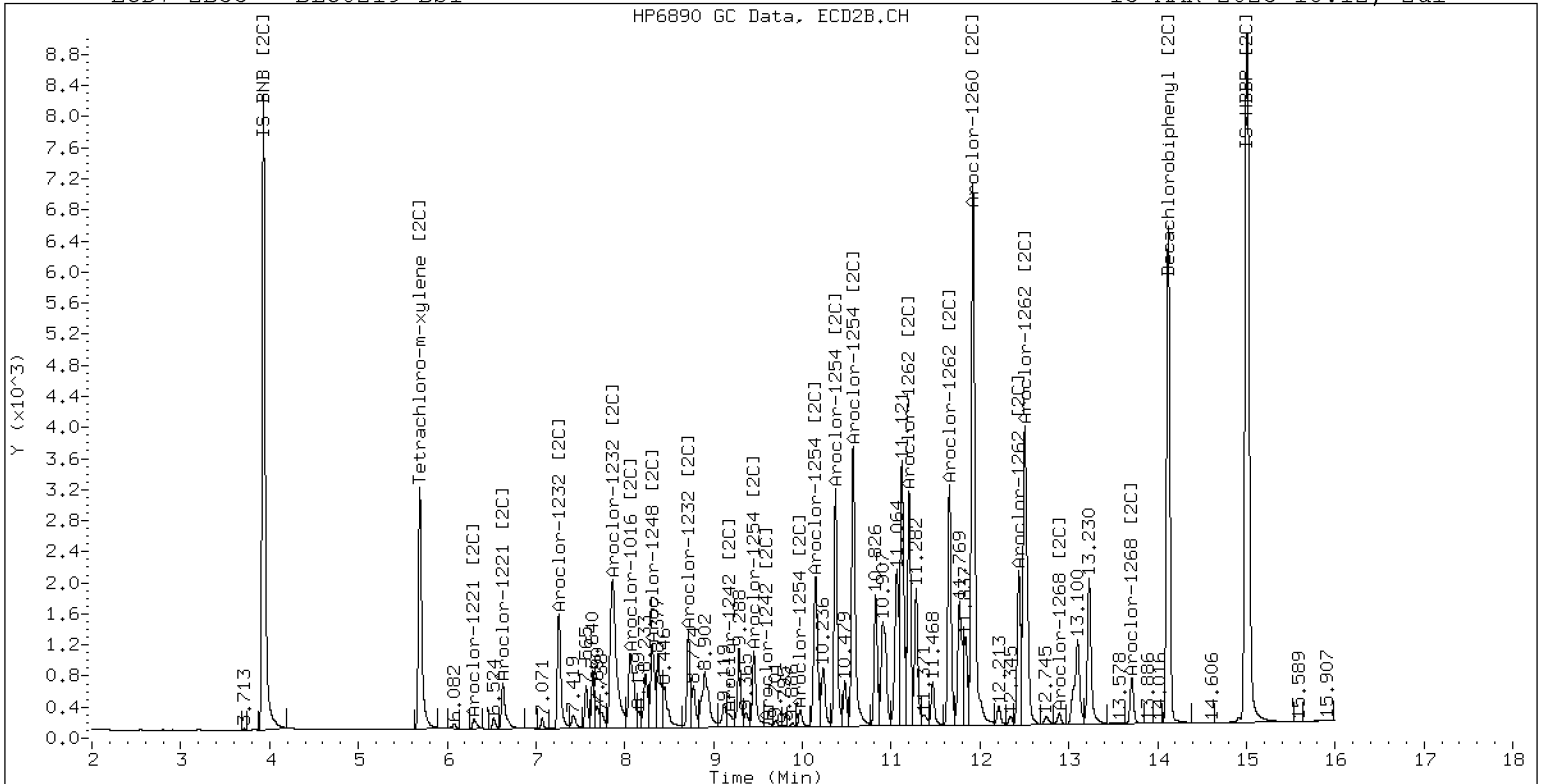
15-MAR-2023 18:12, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 BLC0219-BS1

15-MAR-2023 18:12, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152317ECD7.D
Data file 2: /230315.b/230315.b/03152317ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: BLC0219-BSD1
Client ID:
Injection Date: 15-MAR-2023 18:32
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.811	-0.000	509497	5.691	-0.001	168621	29.7	29.0	2.6	Tetrachloro-m-xylene
13.895	-0.002	856490	14.121	0.000	356293	25.9	36.8	34.8	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1147217	70.3
Hexabromobiphenyl	1429847	3363429	135.2 <-

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	396521	25.8
Hexabromobiphenyl	513946	636625	23.9

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.272	0.001	156687	359.6	1	7.258	-0.001	79128	340.9
Aroclor-1016	2	7.659	0.005	511641	385.2	2	7.864	-0.009	179841	382.1
Aroclor-1016	3	7.795	0.004	240839	371.4	3	8.061	-0.008	76688	360.7
Aroclor-1016	4	8.407	0.002	162007	386.5	4	8.310	-0.003	60260	361.3
Total CollAve (4 peaks):				375.7		Total Col2Ave (4 peaks):				361.2 RPD = 4
Corrected Ave (3 peaks):				372.1		Corrected Ave (3 peaks):				354.3 RPD = 5
Aroclor-1221	1	4.732	0.001	800	7.8	1	---			0.0
Aroclor-1221	2	6.133	0.001	19310	105.1	2	6.302	0.005	7181	101.1
Aroclor-1221	3	6.386	0.003	95967	225.0	3	6.627	0.005	34588	299.1
Total CollAve (3 peaks):				112.6		Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	4.732	0.001	800	13.0	1	---			0.0
Aroclor-1232	2	6.133	0.002	19310	158.4	2	7.258	0.004	79128	784.9
Aroclor-1232	3	7.659	0.003	511641	927.3	3	7.864	0.003	179841	892.1
Aroclor-1232	4	8.581	0.001	208312	888.3	4	8.717	0.002	56684	977.0
Total CollAve (4 peaks):				496.8		Total Col2Ave (3 peaks):				884.7 RPD = 56*
Corrected Ave (3 peaks):				353.2		Corrected Ave: < 3 Peaks				
Aroclor-1242	1	7.272	0.001	156687	440.7	1	7.258	0.000	79128	429.5
Aroclor-1242	2	7.659	0.003	511641	473.9	2	7.864	-0.009	179841	464.4
Aroclor-1242	3	8.407	0.002	162007	482.3	3	9.169	-0.015	11884	98.6
Aroclor-1242	4	8.581	0.002	208312	419.5	4	9.595	-0.020	4463	30.4
Total CollAve (4 peaks):				454.1		Total Col2Ave (4 peaks):				255.7 RPD = 56*
Corrected Ave (3 peaks):				444.7		Corrected Ave (3 peaks):				186.2 RPD = 82*
Aroclor-1248	1	8.407	-0.004	162007	289.4	1	8.310	-0.004	60260	318.3
Aroclor-1248	2	8.581	-0.008	208312	292.8	2	8.717	-0.004	56684	289.6
Aroclor-1248	3	8.996	-0.003	236013	175.8	3	9.169	-0.015	11884	52.8
Aroclor-1248	4	9.303	0.001	193824	283.6	4	9.595	-0.017	4463	16.5
Total CollAve (4 peaks):				260.4		Total Col2Ave (4 peaks):				169.3 RPD = 42*
Corrected Ave (3 peaks):				249.6		Corrected Ave (3 peaks):				119.6 RPD = 70*
Aroclor-1254	1	9.303	0.004	193824	168.3	1	9.454	-0.003	54042	179.3
Aroclor-1254	2	---			0.0	2	9.975	-0.003	11939	49.2
Aroclor-1254	3	9.672	0.004	35440	47.9	3	10.149	0.015	115278	219.8
Aroclor-1254	4	9.810	0.002	105459	73.2	4	10.374	-0.008	149660	292.7
Aroclor-1254	5	10.123	-0.054	507832	562.6	5	10.571	-0.006	202596	650.7
Total CollAve (4 peaks):				213.0		Total Col2Ave (5 peaks):				278.3 RPD = 27
Corrected Ave (3 peaks):				96.4		Corrected Ave (4 peaks):				185.2 RPD = 63*
Aroclor-1260	1	11.046	0.002	427442	353.3	1	11.656	-0.002	155028	414.1
Aroclor-1260	2	11.364	0.003	447379	353.9	2	11.921	-0.004	396758	415.3
Aroclor-1260	3	11.740	0.006	1197748	357.2	3	12.439	-0.001	98681	389.2
Aroclor-1260	4	12.145	0.005	603082	357.2	4	12.506	-0.002	247781	384.8
Aroclor-1260	5	12.247	0.003	251936	346.6	NS	---			----
Total CollAve (5 peaks):				353.6		Total Col2Ave (4 peaks):				400.9 RPD = 13
Corrected Ave (4 peaks):				352.7		Corrected Ave (3 peaks):				396.0 RPD = 12
Aroclor-1262	1	10.831	0.002	840450	814.6	1	11.202	0.001	146805	270.1
Aroclor-1262	2	12.247	0.003	251936	150.1	2	11.656	0.005	155028	334.9
Aroclor-1262	3	12.322	0.003	304569	168.7	3	12.439	0.005	98681	187.9
Aroclor-1262	4	12.992	0.005	270844	164.2	4	12.506	0.004	247781	301.1
Total CollAve (4 peaks):				324.4		Total Col2Ave (4 peaks):				273.5 RPD = 17
Corrected Ave (3 peaks):				161.0		Corrected Ave (3 peaks):				253.0 RPD = 44*
Aroclor-1268	1	12.247	0.000	251936	58.5	1	12.439	0.007	98681	77.0
Aroclor-1268	2	12.322	0.005	304569	71.4	2	12.506	0.006	247781	179.8
Aroclor-1268	3	12.725	0.026	228129	62.5	3	12.895	0.003	7373	6.3
Aroclor-1268	4	13.489	-0.001	59947	5.0	4	13.711	0.002	30712	8.2
Total CollAve (4 peaks):				49.3		Total Col2Ave (4 peaks):				67.8 RPD = 32
Corrected Ave (3 peaks):				42.0		Corrected Ave (3 peaks):				30.5 RPD = 32

Total PCB Area Col1 (5.911 - 13.797) = 11521339 Col1 Total PCB = 0.8 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 3622606 Col2 Total PCB = 0.8 ppm*

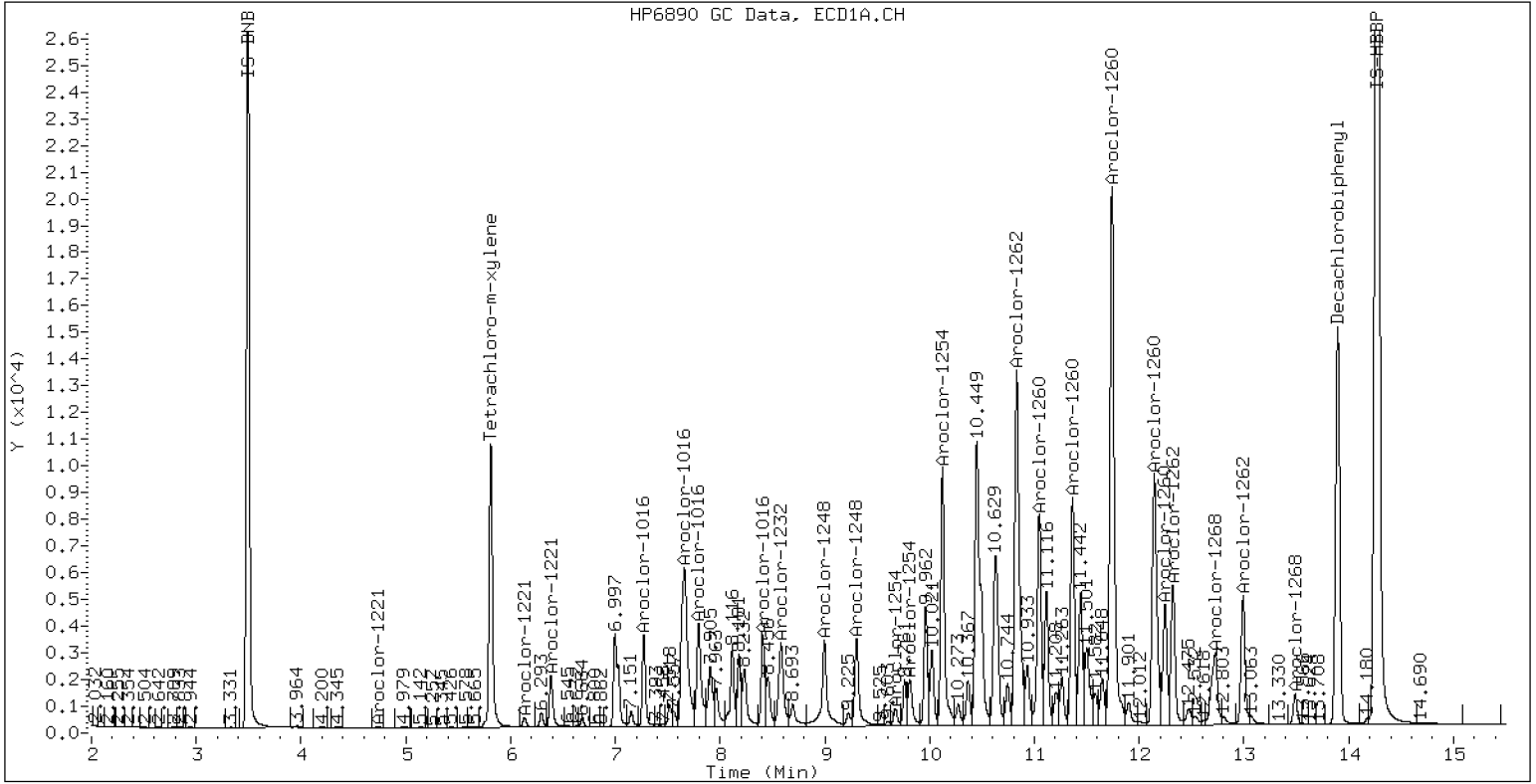
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 BLC0219-BSD1

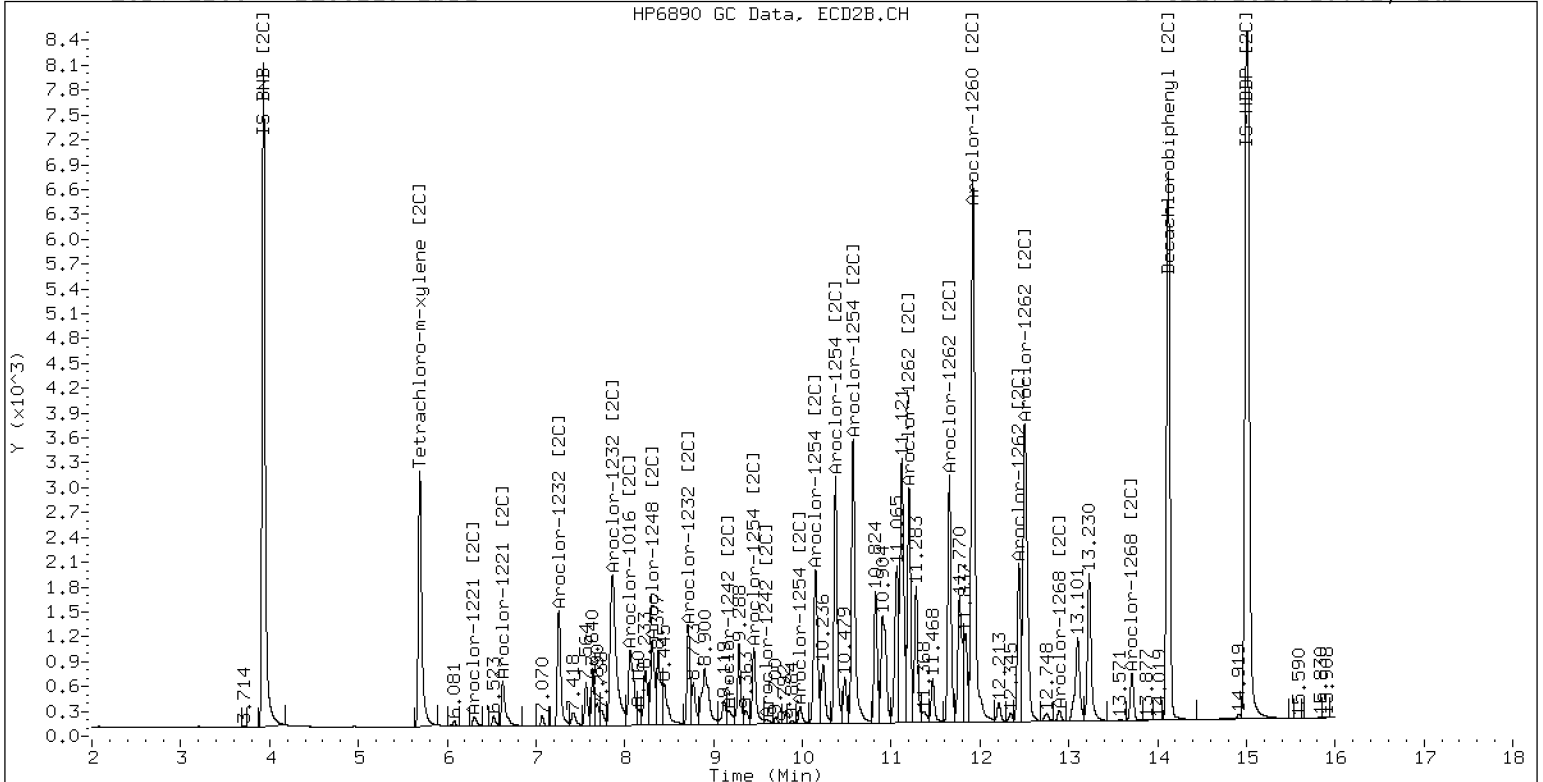
15-MAR-2023 18:32, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 BLC0219-BSD1

15-MAR-2023 18:32, 2u1



ZB-35 Manual Integration: NO



MS / MS DUPLICATE RECOVERY
EPA 8082A

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor OEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Matrix:	<u>Solid</u>	Analyzed:	<u>03/15/23 20:16</u>
Batch:	<u>BLC0219</u>	Laboratory ID:	<u>BLC0219-MS1</u>
Preparation:	<u>EPA 3546 (Microwave)</u>	Sequence Name:	<u>Matrix Spike</u>
Initial/Final:	<u>17.18 g / 2.5 mL</u>	Source Sample:	<u>LDW21-IT608D</u>

COMPOUND	SPIKE ADDED (ug/kg dry)	SAMPLE CONCENTRATION (ug/kg dry)	Q	MS CONCENTRATION (ug/kg dry)	Q	MS % REC. #	QC LIMITS REC.
Aroclor 1016	101	ND	H, U	73.7		73.0	56 - 120
Aroclor 1260	101	ND	H, U	80.3		79.5	58 - 120

* Values outside of QC limits

[2C] indicates second-column analyte, present if quantification on any batch samples used second column data.



MS / MS DUPLICATE RECOVERY
EPA 8082A

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Matrix:	<u>Solid</u>	Analyzed:	<u>03/15/23 20:37</u>
Batch:	<u>BLC0219</u>	Laboratory ID:	<u>BLC0219-MSD1</u>
Preparation:	<u>EPA 3546 (Microwave)</u>	Sequence Name:	<u>Matrix Spike Dup</u>
Initial/Final:	<u>17.18 g / 2.5 mL</u>	Source Sample:	<u>LDW21-IT608D</u>

COMPOUND	SPIKE ADDED (ug/kg dry)	MSD CONCENTRATION (ug/kg dry)	Q	MSD % REC. #	% RPD #	QC LIMITS	
						RPD	REC.
Aroclor 1016	101	73.9		73.2	0.259	30	56 - 120
Aroclor 1260	101	75.9		75.1	5.61	30	58 - 120

* Values outside of QC limits

[2C] indicates second-column analyte, present if quantification on any batch samples used second column data.

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152322ECD7.D
Data file 2: /230315.b/230315.b/03152322ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: BLC0219-MS1
Client ID:
Injection Date: 15-MAR-2023 20:16
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.809	-0.002	496321	5.691	-0.002	171887	30.1	29.3	2.8	Tetrachloro-m-xylene
13.894	-0.003	650336	14.118	-0.003	305974	28.5	35.1	20.9	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1102984	63.7
Hexabromobiphenyl	1429847	2318945	62.2

Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	399893	26.8
Hexabromobiphenyl	513946	571920	11.3

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.271	0.001	159476	380.7	1	7.256	-0.003	82251	351.4
Aroclor-1016	2	7.654	-0.000	498148	390.1	2	7.860	-0.013	177454	373.8
Aroclor-1016	3	7.792	0.001	212078	340.2	3	8.058	-0.012	74804	348.9
Aroclor-1016	4	8.405	0.000	146421	363.3	4	8.308	-0.005	56766	337.5
Total CollAve (4 peaks):				368.6		Total Col2Ave (4 peaks):				352.9 RPD = 4
Corrected Ave (3 peaks):				361.4		Corrected Ave (3 peaks):				345.9 RPD = 4
Aroclor-1221	1	4.734	0.003	821	8.3	1	4.964	0.007	164	4.3
Aroclor-1221	2	6.131	-0.001	25738	145.7	2	6.301	0.004	6319	88.2
Aroclor-1221	3	6.384	0.001	95281	232.3	3	6.626	0.004	38978	334.2
Total CollAve (3 peaks):				128.8		Total Col2Ave (3 peaks):				142.2 RPD = 10
Corrected Ave: < 3 Peaks						Corrected Ave: < 3 Peaks				
Aroclor-1232	1	4.734	0.003	821	13.9	1	4.964	0.007	164	8.0
Aroclor-1232	2	6.131	0.000	25738	219.6	2	7.256	0.002	82251	808.9
Aroclor-1232	3	7.654	-0.002	498148	939.0	3	7.860	-0.001	177454	872.9
Aroclor-1232	4	8.579	-0.002	195277	866.1	4	8.715	-0.000	50548	863.9
Total CollAve (4 peaks):				509.7		Total Col2Ave (4 peaks):				638.4 RPD = 22
Corrected Ave (3 peaks):				366.5		Corrected Ave (3 peaks):				560.3 RPD = 42*
Aroclor-1242	1	7.271	0.000	159476	466.6	1	7.256	-0.002	82251	442.7
Aroclor-1242	2	7.654	-0.002	498148	479.9	2	7.860	-0.013	177454	454.4
Aroclor-1242	3	8.405	0.000	146421	453.3	3	9.162	-0.022	10827	89.1
Aroclor-1242	4	8.579	-0.001	195277	409.0	4	9.576	-0.039	7385	49.9
Total CollAve (4 peaks):				452.2		Total Col2Ave (4 peaks):				259.0 RPD = 54*
Corrected Ave (3 peaks):				443.0		Corrected Ave (3 peaks):				193.9 RPD = 78*
Aroclor-1248	1	8.405	-0.006	146421	272.1	1	8.308	-0.006	56766	297.3
Aroclor-1248	2	8.579	-0.011	195277	285.5	2	8.715	-0.007	50548	256.0
Aroclor-1248	3	8.994	-0.004	196362	152.2	3	9.162	-0.023	10827	47.7
Aroclor-1248	4	9.300	-0.001	169223	257.6	4	9.576	-0.036	7385	27.1
Total CollAve (4 peaks):				241.8		Total Col2Ave (4 peaks):				157.0 RPD = 43*
Corrected Ave (3 peaks):				227.3		Corrected Ave (3 peaks):				110.3 RPD = 69*
Aroclor-1254	1	9.300	0.002	169223	152.8	1	9.450	-0.007	48536	159.7
Aroclor-1254	2	---			0.0	2	9.969	-0.009	11244	46.0
Aroclor-1254	3	9.666	-0.002	36398	51.1	3	10.139	0.005	85322	161.3
Aroclor-1254	4	9.803	-0.004	95817	69.2	4	10.370	-0.012	137338	266.3
Aroclor-1254	5	10.120	-0.057	433245	499.2	5	10.567	-0.011	152598	486.0
Total CollAve (4 peaks):				193.1		Total Col2Ave (5 peaks):				223.8 RPD = 15
Corrected Ave (3 peaks):				91.0		Corrected Ave (4 peaks):				158.3 RPD = 54*
Aroclor-1260	1	11.043	-0.002	318609	381.9	1	11.653	-0.005	127168	378.1
Aroclor-1260	2	11.361	-0.001	351842	403.7	2	11.917	-0.009	324586	378.2
Aroclor-1260	3	11.734	-0.000	937191	405.4	3	12.434	-0.006	89512	393.0
Aroclor-1260	4	12.136	-0.003	489535	420.5	4	12.501	-0.007	209067	361.4
Aroclor-1260	5	12.244	-0.000	198524	396.2	NS	---			----
Total CollAve (5 peaks):				401.5		Total Col2Ave (4 peaks):				377.7 RPD = 6
Corrected Ave (4 peaks):				396.8		Corrected Ave (3 peaks):				372.6 RPD = 6
Aroclor-1262	1	10.827	-0.002	521379	732.9	1	11.199	-0.001	126559	259.2
Aroclor-1262	2	12.244	-0.000	198524	171.5	2	11.653	0.001	127168	305.8
Aroclor-1262	3	12.318	-0.000	241617	194.2	3	12.434	0.001	89512	189.7
Aroclor-1262	4	12.987	-0.001	194892	171.4	4	12.501	-0.001	209067	282.8
Total CollAve (4 peaks):				317.5		Total Col2Ave (4 peaks):				259.4 RPD = 20
Corrected Ave (3 peaks):				179.0		Corrected Ave (3 peaks):				243.9 RPD = 31
Aroclor-1268	1	12.244	-0.003	198524	66.8	1	12.434	0.002	89512	77.7
Aroclor-1268	2	12.318	0.001	241617	82.1	2	12.501	0.001	209067	168.9
Aroclor-1268	3	12.721	0.022	157818	62.7	3	12.891	-0.001	6354	6.0
Aroclor-1268	4	13.487	-0.003	57678	7.0	4	13.708	-0.001	28070	8.3
Total CollAve (4 peaks):				54.6		Total Col2Ave (4 peaks):				65.2 RPD = 18

Corrected Ave (3 peaks): 45.5 Corrected Ave (3 peaks): 30.7 RPD = 39

Total PCB Area Col1 (5.911 - 13.797) = 9595905 Col1 Total PCB = 0.7 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 3256318 Col2 Total PCB = 0.7 ppm*

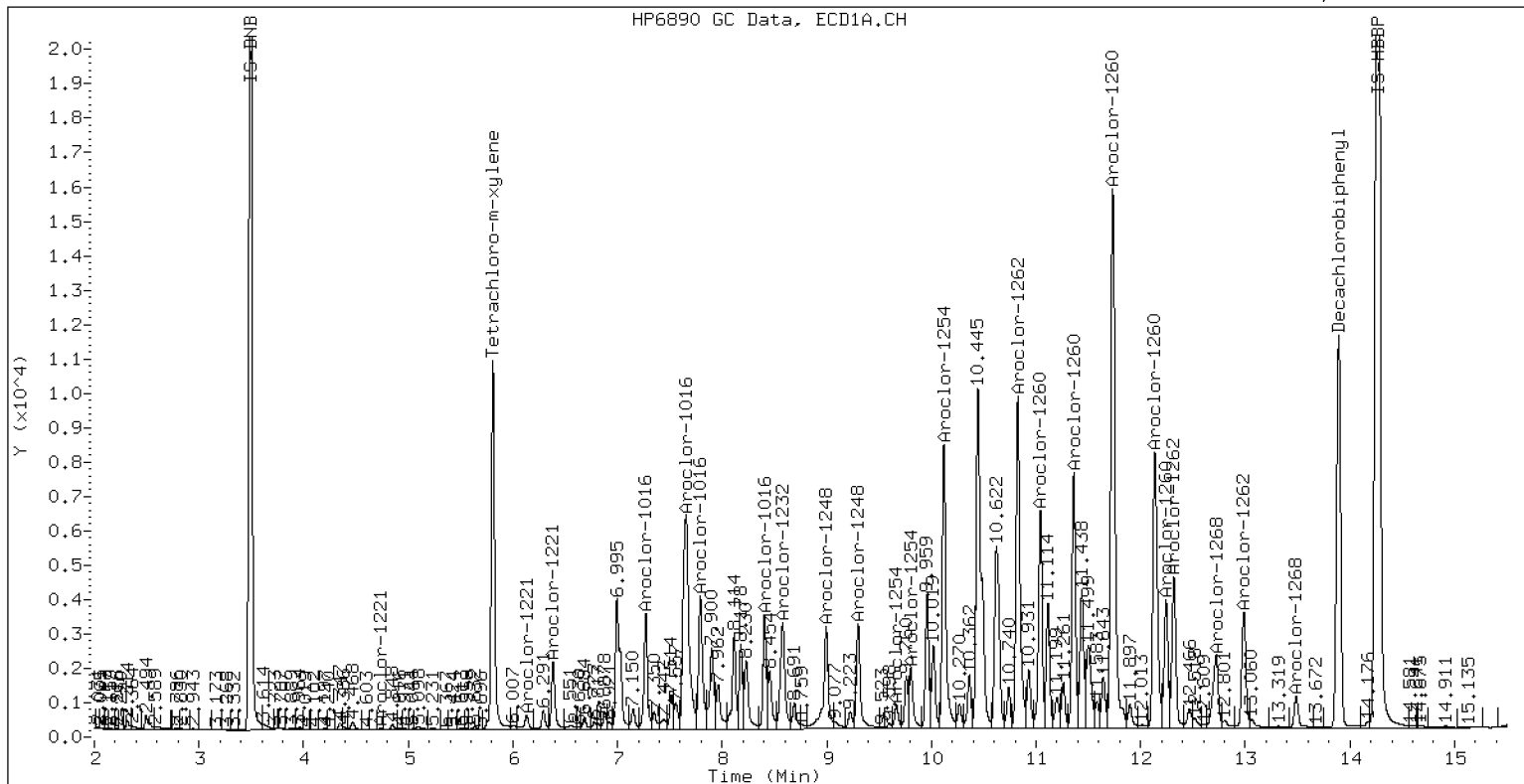
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 BLC0219-MS1

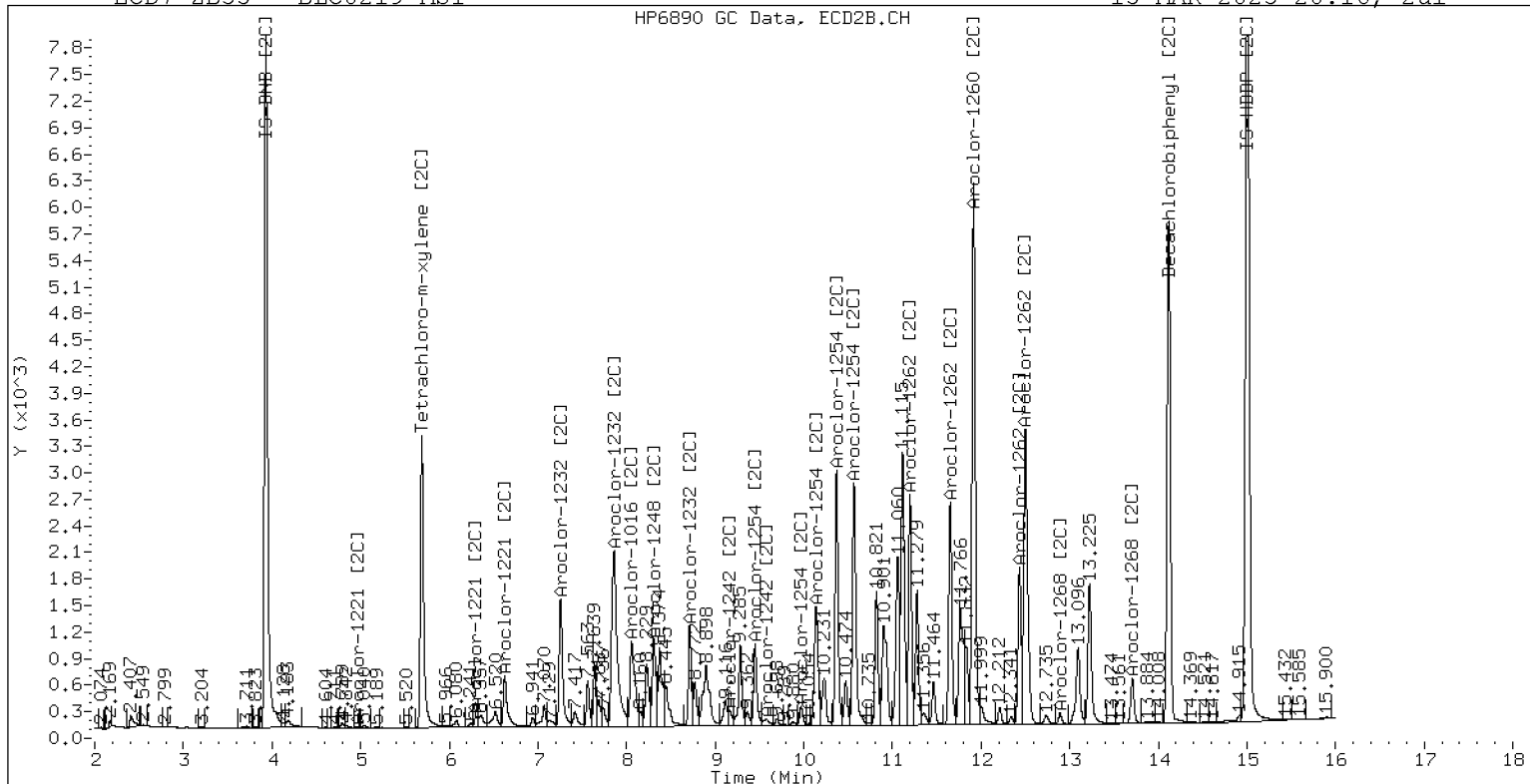
15-MAR-2023 20:16, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 BLC0219-MS1

15-MAR-2023 20:16, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152323ECD7.D
Data file 2: /230315.b/230315.b/03152323ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: BLC0219-MSD1
Client ID:
Injection Date: 15-MAR-2023 20:37
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.809	-0.002	490258	5.691	-0.002	164381	30.0	28.9	3.8	Tetrachloro-m-xylene
13.894	-0.003	634311	14.119	-0.002	303574	27.6	35.5	25.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1094891	62.5
Hexabromobiphenyl	1429847	2334158	63.2

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	388222	23.1
Hexabromobiphenyl	513946	560920	9.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.271	0.000	160694	386.4	1	7.256	-0.003	80873	355.9
Aroclor-1016	2	7.656	0.002	496134	391.3	2	7.858	-0.014	175030	379.8
Aroclor-1016	3	7.792	0.001	209200	338.0	3	8.057	-0.013	73280	352.1
Aroclor-1016	4	8.405	0.000	144914	362.2	4	8.308	-0.005	55082	337.3
Total CollAve (4 peaks):				369.5		Total Col2Ave (4 peaks):				356.3 RPD = 4
Corrected Ave (3 peaks):				362.2		Corrected Ave (3 peaks):				348.4 RPD = 4
Aroclor-1221	1	4.732	0.001	3106	31.7	1	4.965	0.009	121	3.3
Aroclor-1221	2	6.131	-0.001	30180	172.1	2	6.300	0.004	5352	76.9
Aroclor-1221	3	6.384	0.002	95316	234.1	3	6.625	0.003	36699	324.1
Total CollAve (3 peaks):				146.0		Total Col2Ave (3 peaks):				134.8 RPD = 8
Corrected Ave: < 3 Peaks						Corrected Ave: < 3 Peaks				
Aroclor-1232	1	4.732	0.002	3106	53.0	1	4.965	0.009	121	6.1
Aroclor-1232	2	6.131	0.000	30180	259.4	2	7.256	0.002	80873	819.3
Aroclor-1232	3	7.656	-0.000	496134	942.2	3	7.858	-0.002	175030	886.8
Aroclor-1232	4	8.579	-0.002	196530	878.1	4	8.714	-0.001	50243	884.5
Total CollAve (4 peaks):				533.2		Total Col2Ave (4 peaks):				649.2 RPD = 20
Corrected Ave (3 peaks):				396.8		Corrected Ave (3 peaks):				570.0 RPD = 36
Aroclor-1242	1	7.271	-0.000	160694	473.6	1	7.256	-0.002	80873	448.4
Aroclor-1242	2	7.656	-0.000	496134	481.5	2	7.858	-0.014	175030	461.6
Aroclor-1242	3	8.405	-0.000	144914	452.0	3	9.159	-0.025	9775	82.9
Aroclor-1242	4	8.579	-0.001	196530	414.7	4	9.580	-0.035	5554	38.6
Total CollAve (4 peaks):				455.4		Total Col2Ave (4 peaks):				257.9 RPD = 55*
Corrected Ave (3 peaks):				446.8		Corrected Ave (3 peaks):				190.0 RPD = 81*
Aroclor-1248	1	8.405	-0.006	144914	271.3	1	8.308	-0.006	55082	297.2
Aroclor-1248	2	8.579	-0.011	196530	289.4	2	8.714	-0.007	50243	262.2
Aroclor-1248	3	8.994	-0.004	177597	138.6	3	9.159	-0.025	9775	44.3
Aroclor-1248	4	9.299	-0.002	163353	250.5	4	9.580	-0.032	5554	21.0
Total CollAve (4 peaks):				237.5		Total Col2Ave (4 peaks):				156.2 RPD = 41*
Corrected Ave (3 peaks):				220.1		Corrected Ave (3 peaks):				109.1 RPD = 67*
Aroclor-1254	1	9.299	0.001	163353	148.6	1	9.450	-0.008	46993	159.3
Aroclor-1254	2	---	---	---	0.0	2	9.970	-0.008	10393	43.8
Aroclor-1254	3	9.666	-0.002	33025	46.7	3	10.139	0.005	74133	144.3
Aroclor-1254	4	9.803	-0.004	91104	66.3	4	10.370	-0.012	133276	266.2
Aroclor-1254	5	10.119	-0.057	417225	484.3	5	10.567	-0.010	134535	441.3
Total CollAve (4 peaks):				186.5		Total Col2Ave (5 peaks):				211.0 RPD = 12
Corrected Ave (3 peaks):				87.2		Corrected Ave (4 peaks):				153.4 RPD = 55*
Aroclor-1260	1	11.041	-0.003	289753	345.1	1	11.653	-0.005	119148	361.2
Aroclor-1260	2	11.359	-0.002	324578	370.0	2	11.917	-0.008	310025	368.3
Aroclor-1260	3	11.733	-0.001	903769	388.4	3	12.434	-0.006	85142	381.2
Aroclor-1260	4	12.136	-0.003	482414	411.7	4	12.500	-0.007	196579	346.5
Aroclor-1260	5	12.244	0.000	193221	383.1	NS	---	---	---	---
Total CollAve (5 peaks):				379.6		Total Col2Ave (4 peaks):				364.3 RPD = 4
Corrected Ave (4 peaks):				371.6		Corrected Ave (3 peaks):				358.7 RPD = 4
Aroclor-1262	1	10.826	-0.003	441137	616.1	1	11.199	-0.001	123253	257.3
Aroclor-1262	2	12.244	0.000	193221	165.8	2	11.653	0.002	119148	292.1
Aroclor-1262	3	12.318	-0.001	233834	186.7	3	12.434	0.001	85142	184.0
Aroclor-1262	4	12.986	-0.001	182325	159.3	4	12.500	-0.002	196579	271.1
Total CollAve (4 peaks):				282.0		Total Col2Ave (4 peaks):				251.1 RPD = 12
Corrected Ave (3 peaks):				170.6		Corrected Ave (3 peaks):				237.5 RPD = 33
Aroclor-1268	1	12.244	-0.002	193221	64.6	1	12.434	0.002	85142	75.4
Aroclor-1268	2	12.318	0.001	233834	79.0	2	12.500	0.000	196579	161.9
Aroclor-1268	3	12.718	0.018	224993	88.8	3	12.891	-0.000	5921	5.7
Aroclor-1268	4	13.487	-0.003	41034	4.9	4	13.708	-0.001	25956	7.8
Total CollAve (4 peaks):				59.3		Total Col2Ave (4 peaks):				62.7 RPD = 6

Corrected Ave (3 peaks): 49.5 Corrected Ave (3 peaks): 29.6 RPD = 50*

Total PCB Area Col1 (5.911 - 13.797) = 9288063 Col1 Total PCB = 0.7 ppm*
Total PCB Area Col2 (5.792 - 14.021) = 3129184 Col2 Total PCB = 0.7 ppm*

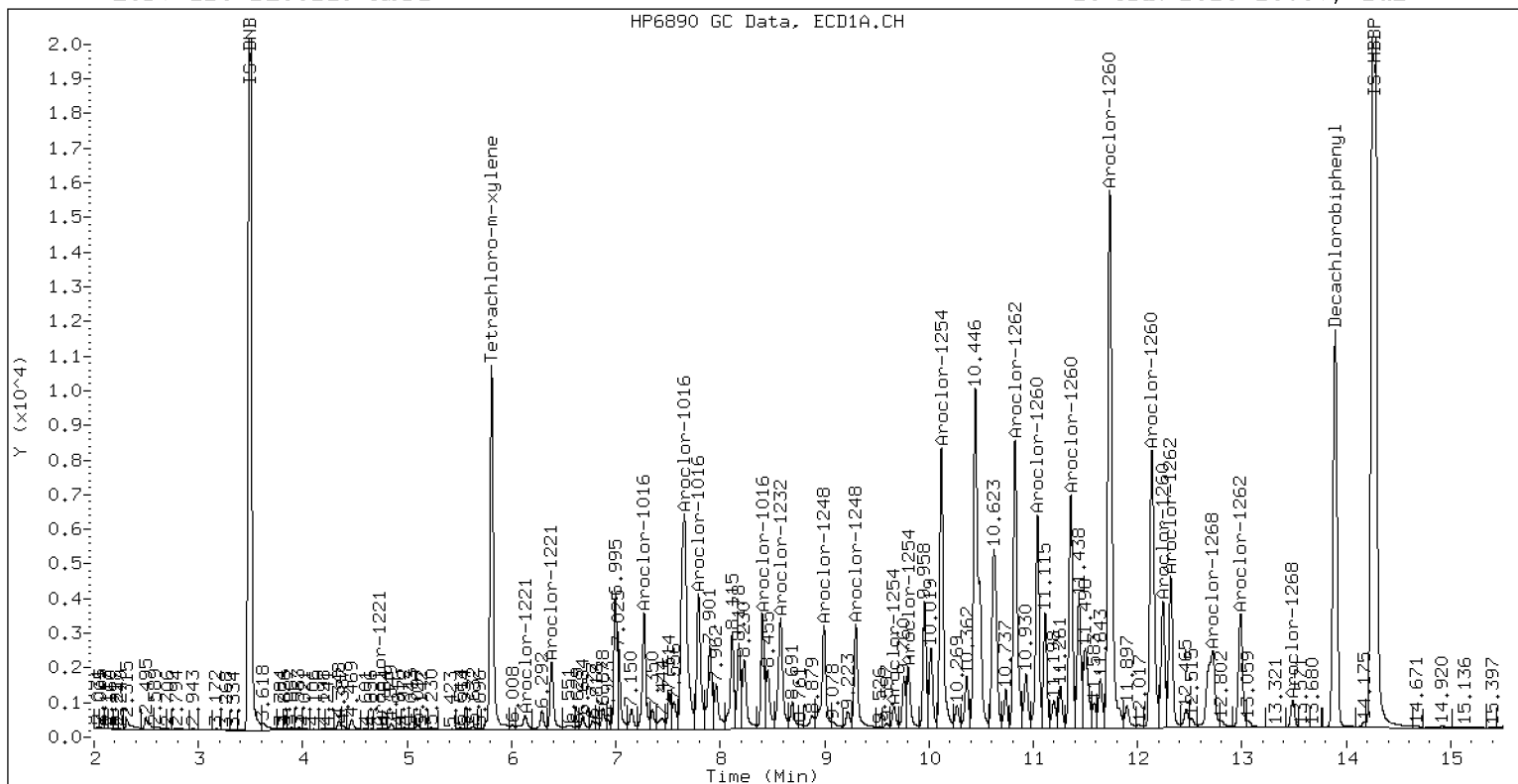
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 BLC0219-MSD1

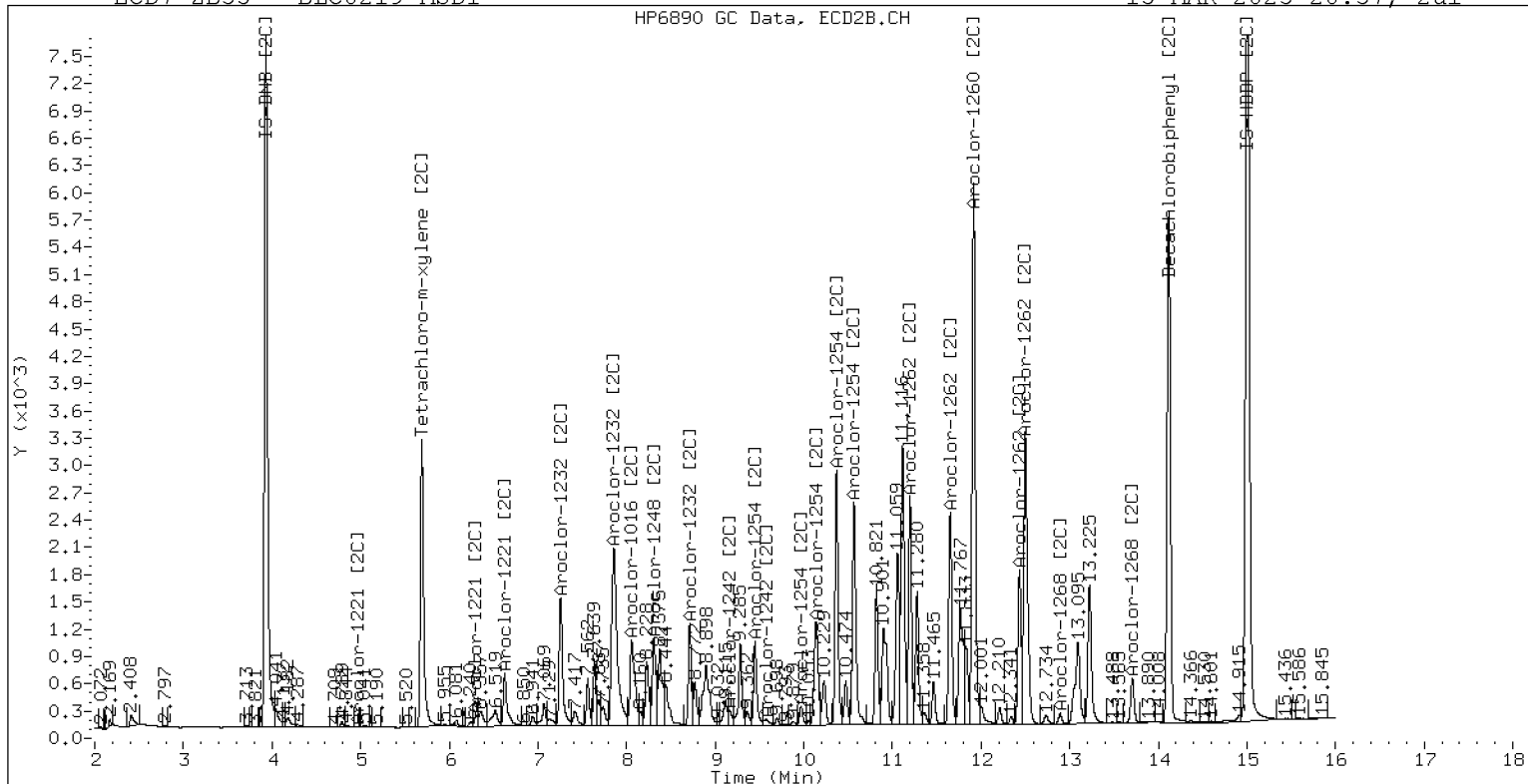
15-MAR-2023 20:37, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 BLC0219-MSD1

15-MAR-2023 20:37, 2u1



ZB-35 Manual Integration: NO



STANDARD REFERENCE MATERIAL RECOVERY

EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Laboratory ID: BLC0219-SRM1

Batch: BLC0219

Initial/Final: 2.5 g / 2.5 mL

Preparation: EPA 3546 (Microwave)

Analyzed: 03/15/2023 18:53

Standard ID: K003527

Expires: 04/12/2023

Standard Lot#: PSRM0150

Description: Puget Sound reference-SRM

ANALYTE	TRUE (ug/kg wet)	FOUND (ug/kg wet)	MDL	MRL	Q	SRM % REC.	QC LIMITS REC.
Aroclor 1260	108.00	94.4	2.9	20.0		87.5	38 - 167
Aroclor 1260 [2C]	108.00	104	2.9	20.0		96.2	38 - 167

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152318ECD7.D
Data file 2: /230315.b/230315.b/03152318ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: BLC0219-SRM1
Client ID:
Injection Date: 15-MAR-2023 18:53
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.809	-0.002	470266	5.689	-0.003	166609	28.5	29.8	4.3	Tetrachloro-m-xylene
13.890	-0.007	535631	14.116	-0.005	262363	27.9	31.9	13.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1103257	63.7
Hexabromobiphenyl	1429847	1949653	36.4

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	380915	20.8
Hexabromobiphenyl	513946	540684	5.2

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.240	-0.030	25069	59.8	1	7.266	0.007	6634	29.8
Aroclor-1016	2	7.653	-0.001	15881	12.4	2	7.858	-0.014	9864	21.8
Aroclor-1016	3	7.798	0.008	9834	15.8	3	8.054	-0.015	1215	5.9
Aroclor-1016	4	8.404	-0.001	13537	33.6	4	8.306	-0.008	6601	41.2
Total CollAve (4 peaks):				30.4		Total Col2Ave (4 peaks):				24.7 RPD = 21
Corrected Ave (3 peaks):				20.6		Corrected Ave (3 peaks):				19.2 RPD = 7
Aroclor-1221	1	4.788	0.057	287	2.9	1	4.945	-0.011	817	22.7
Aroclor-1221	2	6.077	-0.055	10520	59.5	2	6.352	0.056	11300	165.6
Aroclor-1221	3	6.393	0.011	3516	8.6	3	6.644	0.022	3922	35.3
Total CollAve (3 peaks):				23.7		Total Col2Ave (3 peaks):				74.5 RPD = 104*
Corrected Ave: < 3 Peaks						Corrected Ave: < 3 Peaks				
Aroclor-1232	1	4.788	0.058	287	4.9	1	4.945	-0.011	817	41.9
Aroclor-1232	2	6.077	-0.054	10520	89.7	2	7.266	0.012	6634	68.5
Aroclor-1232	3	7.653	-0.003	15881	29.9	3	7.858	-0.003	9864	50.9
Aroclor-1232	4	8.574	-0.006	13130	58.2	4	8.712	-0.003	4576	82.1
Total CollAve (4 peaks):				45.7		Total Col2Ave (4 peaks):				60.9 RPD = 28
Corrected Ave (3 peaks):				31.0		Corrected Ave (3 peaks):				53.8 RPD = 54*
Aroclor-1242	1	7.240	-0.031	25069	73.3	1	7.266	0.008	6634	37.5
Aroclor-1242	2	7.653	-0.003	15881	15.3	2	7.858	-0.014	9864	26.5
Aroclor-1242	3	8.404	-0.002	13537	41.9	3	9.155	-0.029	6244	53.9
Aroclor-1242	4	8.574	-0.005	13130	27.5	4	9.575	-0.040	5456	38.7
Total CollAve (4 peaks):				39.5		Total Col2Ave (4 peaks):				39.2 RPD = 1
Corrected Ave (3 peaks):				28.2		Corrected Ave (3 peaks):				34.2 RPD = 19
Aroclor-1248	1	8.404	-0.008	13537	25.1	1	8.306	-0.009	6601	36.3
Aroclor-1248	2	8.574	-0.015	13130	19.2	2	8.712	-0.009	4576	24.3
Aroclor-1248	3	8.991	-0.007	41836	32.4	3	9.155	-0.030	6244	28.9
Aroclor-1248	4	9.293	-0.008	60727	92.4	4	9.575	-0.037	5456	21.0
Total CollAve (4 peaks):				42.3		Total Col2Ave (4 peaks):				27.6 RPD = 42*
Corrected Ave (3 peaks):				25.6		Corrected Ave (3 peaks):				24.7 RPD = 3
Aroclor-1254	1	9.293	-0.005	60727	54.8	1	9.446	-0.011	19435	67.1
Aroclor-1254	2	9.369	-0.008	22786	45.7	2	9.965	-0.013	9644	41.4
Aroclor-1254	3	9.666	-0.002	34617	48.6	3	10.122	-0.012	39308	78.0
Aroclor-1254	4	9.796	-0.011	89159	64.4	4	10.365	-0.017	49057	99.9
Aroclor-1254	5	10.116	-0.060	129692	149.4	5	10.562	-0.016	49337	164.9
Total CollAve (5 peaks):				72.6		Total Col2Ave (5 peaks):				90.3 RPD = 22
Corrected Ave (4 peaks):				53.4		Corrected Ave (4 peaks):				71.6 RPD = 29
Aroclor-1260	1	11.039	-0.006	75477	107.6	1	11.649	-0.009	35430	111.4
Aroclor-1260	2	11.351	-0.011	59611	81.3	2	11.911	-0.014	77993	96.1
Aroclor-1260	3	11.724	-0.009	186631	96.0	3	12.429	-0.011	25444	118.2
Aroclor-1260	4	12.127	-0.012	98294	100.4	4	12.495	-0.013	49190	89.9
Aroclor-1260	5	12.240	-0.004	36578	86.8	NS	---			----
Total CollAve (5 peaks):				94.4		Total Col2Ave (4 peaks):				103.9 RPD = 10
Corrected Ave (4 peaks):				91.2		Corrected Ave (3 peaks):				99.2 RPD = 8
Aroclor-1262	1	10.817	-0.012	171663	287.0	1	11.195	-0.005	31216	67.6
Aroclor-1262	2	12.240	-0.004	36578	37.6	2	11.649	-0.002	35430	90.1
Aroclor-1262	3	12.313	-0.006	46004	44.0	3	12.429	-0.005	25444	57.0
Aroclor-1262	4	12.979	-0.008	48725	51.0	4	12.495	-0.007	49190	70.4
Total CollAve (4 peaks):				104.9		Total Col2Ave (4 peaks):				71.3 RPD = 38
Corrected Ave (3 peaks):				44.2		Corrected Ave (3 peaks):				65.0 RPD = 38
Aroclor-1268	1	12.240	-0.007	36578	14.6	1	12.429	-0.003	25444	23.4
Aroclor-1268	2	12.313	-0.004	46004	18.6	2	12.495	-0.005	49190	42.0
Aroclor-1268	3	12.668	-0.031	128279	60.6	3	12.889	-0.002	2516	2.5
Aroclor-1268	4	13.481	-0.009	8398	1.2	4	13.704	-0.005	8852	2.8
Total CollAve (4 peaks):				23.8		Total Col2Ave (4 peaks):				17.7 RPD = 29

Corrected Ave (3 peaks): 11.5 Corrected Ave (3 peaks): 9.6 RPD = 18

Total PCB Area Col1 (5.911 - 13.797) = 2322673 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 904909 Col2 Total PCB = 0.2 ppm*

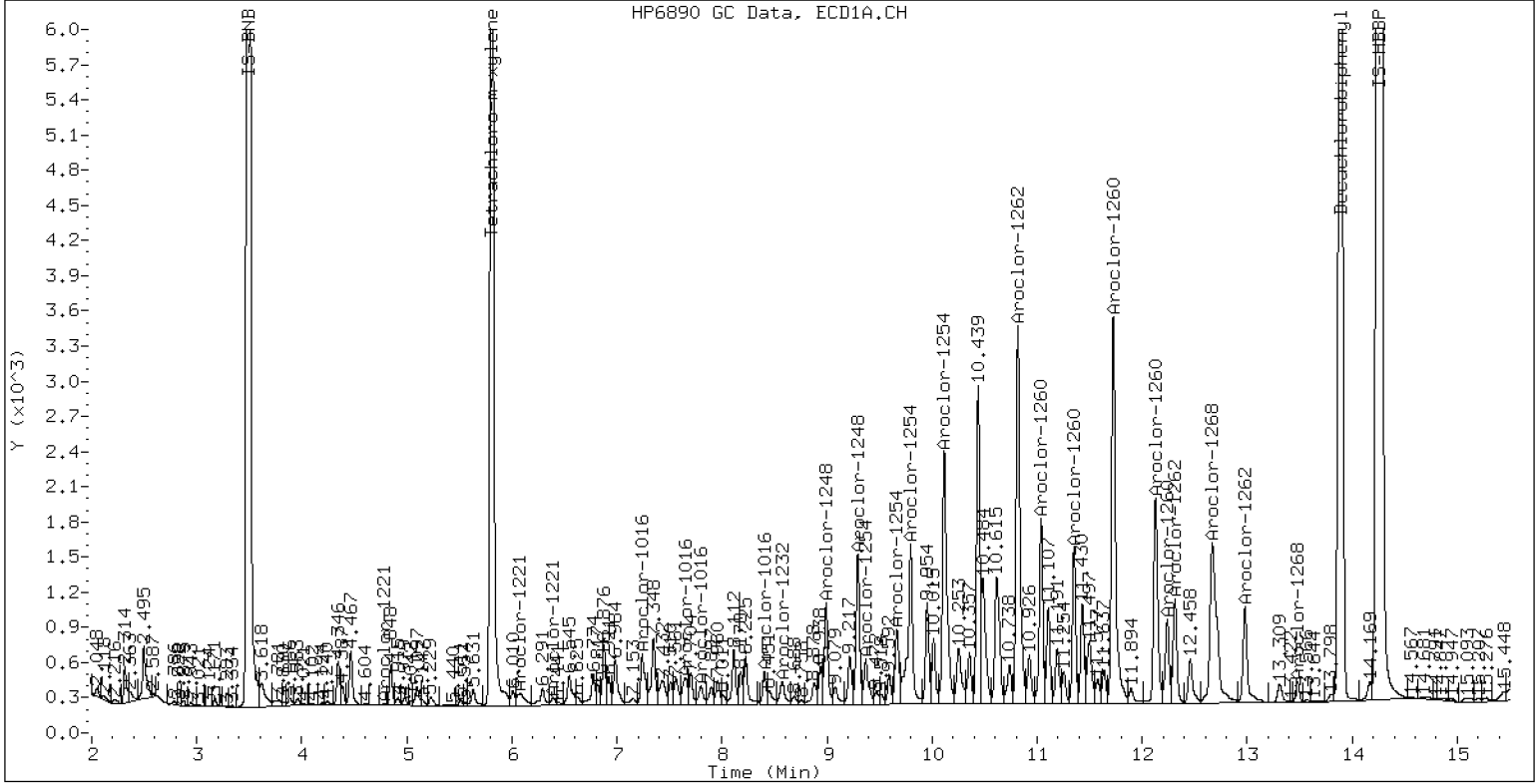
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 BLC0219-SRM1

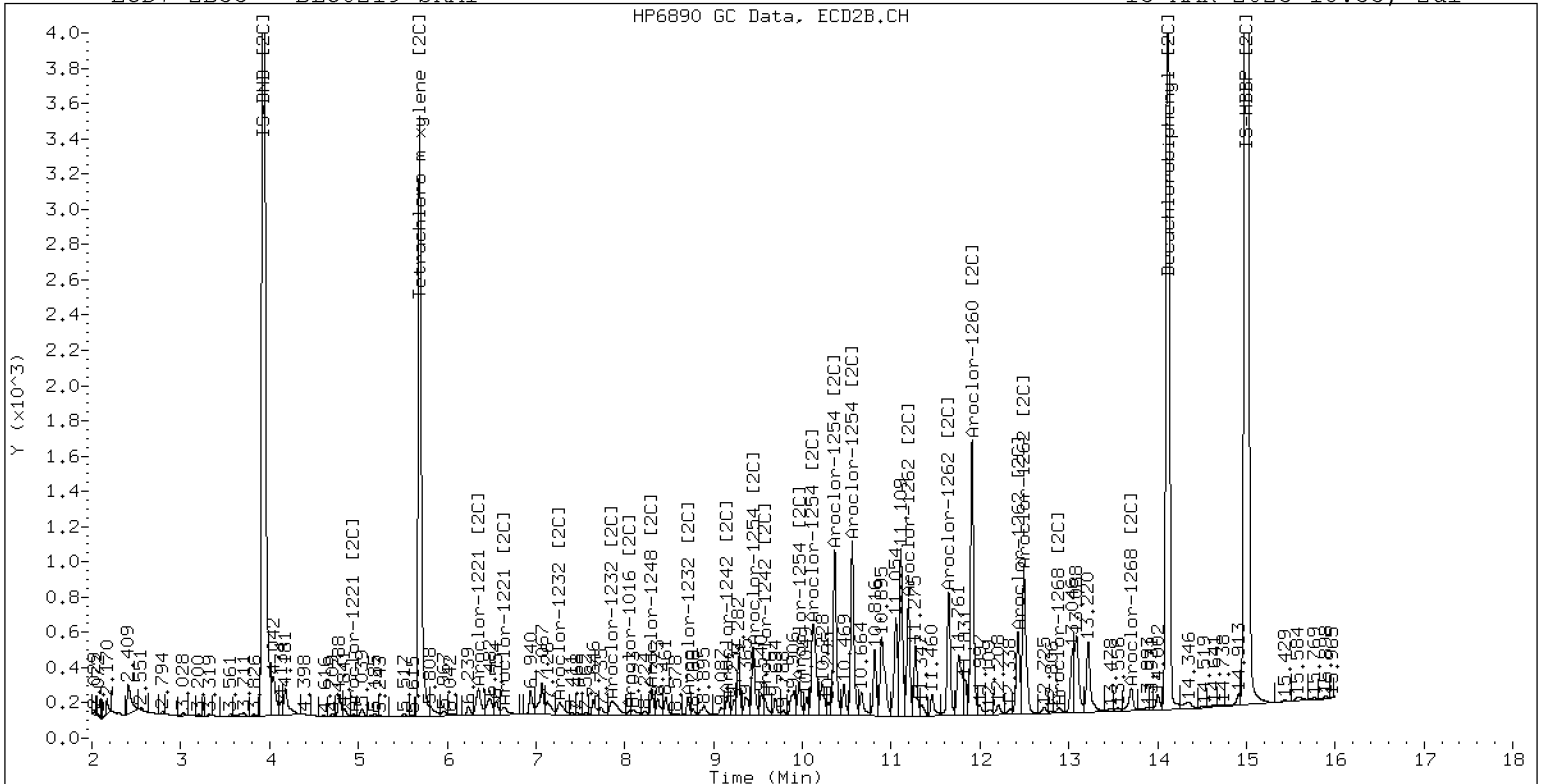
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ZB-5 Manual Integration: NO

ECD7-ZB35 BLC0219-SRM1

15-MAR-2023 18:53, 2u1



ZB-35 Manual Integration: NO



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (1):	ZB5

Compound	Level 01		Level 02		Level 03		Level 04		Level 05		Level 06	
	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
Aroclor 1016	250	0.0511017	20	0.0514466	50	5.107478E-02	1000	4.502727E-02	100	5.036259E-02	500	0.0471841
Aroclor-1016 (1)	250	0.031405	20	3.172321E-02	50	3.253176E-02	1000	2.667138E-02	100	3.141686E-02	500	2.856283E-02
Aroclor-1016 (2)	250	9.848704E-02	20	9.239415E-02	50	9.245774E-02	1000	8.848657E-02	100	9.221759E-02	500	0.0917416
Aroclor-1016 (3)	250	4.393471E-02	20	5.165382E-02	50	5.037363E-02	1000	3.720718E-02	100	4.822959E-02	500	3.990906E-02
Aroclor-1016 (4)	250	3.058004E-02	20	3.001523E-02	50	2.893599E-02	1000	2.774395E-02	100	2.958631E-02	500	2.852291E-02
Aroclor 1260	250	4.264611E-02	20	3.933745E-02	50	3.914748E-02	1000	0.0377098	100	3.888069E-02	500	3.753326E-02
Aroclor-1260 (1)	250	3.096387E-02	20	2.926415E-02	50	2.920486E-02	1000	2.746159E-02	100	2.841034E-02	500	2.736642E-02
Aroclor-1260 (2)	250	3.291004E-02	20	2.966791E-02	50	3.006192E-02	1000	2.856573E-02	100	3.010757E-02	500	2.910054E-02
Aroclor-1260 (3)	250	8.575373E-02	20	8.087657E-02	50	8.045158E-02	1000	7.674417E-02	100	7.953737E-02	500	7.514663E-02
Aroclor-1260 (4)	250	4.484933E-02	20	3.904963E-02	50	3.886754E-02	1000	3.922291E-02	100	3.955346E-02	500	3.941669E-02
Aroclor-1260 (5)	250	1.875356E-02	20	1.782901E-02	50	1.715148E-02	1000	1.655457E-02	100	1.679471E-02	500	1.663603E-02
Decachlorobiphenyl	40	0.7880759	3.2	0.8290115	8	0.8055828	160	0.797423	16	0.7758675	80	0.7312517
Tetrachlorometaxylene	40	1.205085	3.2	1.168271	8	1.244015	160	1.241136	16	1.185465	80	1.122954



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (1):	ZB5

Compound	Level 07		Level 08		Level 09		Level 10		Level 11		Level 12	
	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
Aroclor 1221							250	1.657582E-02				
Aroclor-1221 (1)							250	7.164712E-03				
Aroclor-1221 (2)							250	0.0128135				
Aroclor-1221 (3)							250	2.974924E-02				
Aroclor 1232									250	1.690391E-02		
Aroclor-1232 (1)									250	4.285984E-03		
Aroclor-1232 (2)									250	8.499602E-03		
Aroclor-1232 (3)									250	3.847671E-02		
Aroclor-1232 (4)									250	1.635336E-02		
Aroclor 1242	250	3.953397E-02										
Aroclor-1242 (1)	250	2.479209E-02										
Aroclor-1242 (2)	250	7.528986E-02										
Aroclor-1242 (3)	250	2.342574E-02										
Aroclor-1242 (4)	250	3.462819E-02										
Aroclor 1248			250	5.747549E-02								
Aroclor-1248 (1)			250	3.903293E-02								
Aroclor-1248 (2)			250	0.0496149								
Aroclor-1248 (3)			250	9.360202E-02								
Aroclor-1248 (4)			250	4.765213E-02								
Aroclor 1254					250	6.629494E-02						
Aroclor-1254 (1)					250	8.033306E-02						
Aroclor-1254 (2)					250	0.0361302						
Aroclor-1254 (3)					250	5.164705E-02						
Aroclor-1254 (4)					250	0.100423						
Aroclor-1254 (5)					250	6.294139E-02						
Aroclor 1262							250	3.665955E-02				
Aroclor-1262 (1)							250	2.454122E-02				
Aroclor-1262 (2)							250	3.993338E-02				



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (1):	ZB5

Compound	Level 07		Level 08		Level 09		Level 10		Level 11		Level 12	
	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
Aroclor-1262 (3)							250	4.292945E-02				
Aroclor-1262 (4)							250	3.923413E-02				
Aroclor 1268									250	0.1442124		
Aroclor-1268 (1)									250	0.102504		
Aroclor-1268 (2)									250	0.1015072		
Aroclor-1268 (3)									250	8.685666E-02		
Aroclor-1268 (4)									250	0.2859818		



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (1):	ZB5

COMPOUND	Mean RRF	RRF RSD	Linear COD	Quad COD	Limit Type & Limit	Q
Aroclor 1016	4.936617E-02	5.3			RSD (20)	
Aroclor-1016 (1)	3.038517E-02	7.4			RSD (20)	
Aroclor-1016 (2)	9.263078E-02	3.5			RSD (20)	
Aroclor-1016 (3)	0.045218	12.9			RSD (20)	
Aroclor-1016 (4)	2.923074E-02	3.5			RSD (20)	
Aroclor 1221		0.0			RSD (20)	
Aroclor-1221 (1)		0.0			RSD (20)	
Aroclor-1221 (2)		0.0			RSD (20)	
Aroclor-1221 (3)		0.0			RSD (20)	
Aroclor 1232		0.0			RSD (20)	
Aroclor-1232 (1)		0.0			RSD (20)	
Aroclor-1232 (2)		0.0			RSD (20)	
Aroclor-1232 (3)		0.0			RSD (20)	
Aroclor-1232 (4)		0.0			RSD (20)	
Aroclor 1242		0.0			RSD (20)	
Aroclor-1242 (1)		0.0			RSD (20)	
Aroclor-1242 (2)		0.0			RSD (20)	
Aroclor-1242 (3)		0.0			RSD (20)	
Aroclor-1242 (4)		0.0			RSD (20)	
Aroclor 1248		0.0			RSD (20)	
Aroclor-1248 (1)		0.0			RSD (20)	
Aroclor-1248 (2)		0.0			RSD (20)	
Aroclor-1248 (3)		0.0			RSD (20)	
Aroclor-1248 (4)		0.0			RSD (20)	
Aroclor 1254		0.0			RSD (20)	
Aroclor-1254 (1)		0.0			RSD (20)	
Aroclor-1254 (2)		0.0			RSD (20)	
Aroclor-1254 (3)		0.0			RSD (20)	
Aroclor-1254 (4)		0.0			RSD (20)	
Aroclor-1254 (5)		0.0			RSD (20)	
Aroclor 1260	3.920913E-02	4.7			RSD (20)	



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (1):	ZB5

COMPOUND	Mean RRF	RRF RSD	Linear COD	Quad COD	Limit Type & Limit	Q
Aroclor-1260 (1)	2.877854E-02	4.7			RSD (20)	
Aroclor-1260 (2)	3.006895E-02	5.0			RSD (20)	
Aroclor-1260 (3)	7.975167E-02	4.6			RSD (20)	
Aroclor-1260 (4)	4.015993E-02	5.8			RSD (20)	
Aroclor-1260 (5)	1.728656E-02	5.0			RSD (20)	
Aroclor 1262		0.0			RSD (20)	
Aroclor-1262 (1)		0.0			RSD (20)	
Aroclor-1262 (2)		0.0			RSD (20)	
Aroclor-1262 (3)		0.0			RSD (20)	
Aroclor-1262 (4)		0.0			RSD (20)	
Aroclor 1268		0.0			RSD (20)	
Aroclor-1268 (1)		0.0			RSD (20)	
Aroclor-1268 (2)		0.0			RSD (20)	
Aroclor-1268 (3)		0.0			RSD (20)	
Aroclor-1268 (4)		0.0			RSD (20)	
Decachlorobiphenyl	0.7878687	4.2			RSD (20)	
Tetrachlorometaxylene	1.194488	3.9			RSD (20)	



INITIAL CALIBRATION DATA

EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (2):	ZB35

Compound	Level 01		Level 02		Level 03		Level 04		Level 05		Level 06	
	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
Aroclor 1016 [2C]	250	0.0571297	20	5.099991E-02	50	0.0573721	1000	5.075893E-02	100	5.836783E-02	500	5.288542E-02
Aroclor-1016 (1) [2C]	250	4.732763E-02	20	5.070692E-02	50	5.021603E-02	1000	4.080107E-02	100	4.868029E-02	500	4.325569E-02
Aroclor-1016 (2) [2C]	250	0.1025919	20	8.142537E-02	50	9.407053E-02	1000	9.361548E-02	100	0.1015897	500	9.651233E-02
Aroclor-1016 (3) [2C]	250	4.410181E-02	20	4.005508E-02	50	4.718351E-02	1000	3.925581E-02	100	4.613223E-02	500	4.062487E-02
Aroclor-1016 (4) [2C]	250	3.449742E-02	20	3.181228E-02	50	3.801833E-02	1000	0.0293633	100	0.0370691	500	3.114879E-02
Aroclor 1260 [2C]	250	7.266587E-02	20	0.0760446	50	7.181489E-02	1000	0.0636872	100	6.942709E-02	500	6.617305E-02
Aroclor-1260 (1) [2C]	250	4.801376E-02	20	5.286013E-02	50	4.911343E-02	1000	4.201242E-02	100	4.695569E-02	500	4.328842E-02
Aroclor-1260 (2) [2C]	250	0.1266443	20	0.1297611	50	0.1243096	1000	0.1054494	100	0.1209452	500	0.1132043
Aroclor-1260 (3) [2C]	250	3.207621E-02	20	3.524009E-02	50	3.146502E-02	1000	0.0319805	100	2.936945E-02	500	3.102287E-02
Aroclor-1260 (4) [2C]	250	8.392913E-02	20	8.631709E-02	50	8.237154E-02	1000	7.530648E-02	100	0.080438	500	7.717658E-02
Decachlorobiphenyl [2C]	40	1.310398	3.2	1.170661	8	1.20406	160	1.207975	16	1.205489	80	1.211045
Tetrachlorometaxylene [2C]	40	1.219073	3.2	1.21526	8	1.195453	160	1.111394	16	1.175548	80	1.125598



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (2):	ZB35

Compound	Level 07		Level 08		Level 09		Level 10		Level 11		Level 12	
	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
Aroclor 1221 [2C]							250	1.507982E-02				
Aroclor-1221 (1) [2C]							250	7.573277E-03				
Aroclor-1221 (2) [2C]							250	0.0143332				
Aroclor-1221 (3) [2C]							250	2.333296E-02				
Aroclor 1232 [2C]									250	1.920227E-02		
Aroclor-1232 (1) [2C]									250	4.09321E-03		
Aroclor-1232 (2) [2C]									250	2.034072E-02		
Aroclor-1232 (3) [2C]									250	4.067023E-02		
Aroclor-1232 (4) [2C]									250	1.170493E-02		
Aroclor 1242 [2C]	250	4.230924E-02										
Aroclor-1242 (1) [2C]	250	3.716893E-02										
Aroclor-1242 (2) [2C]	250	7.813249E-02										
Aroclor-1242 (3) [2C]	250	2.431205E-02										
Aroclor-1242 (4) [2C]	250	0.0296235										
Aroclor 1248 [2C]			250	4.442703E-02								
Aroclor-1248 (1) [2C]			250	3.819713E-02								
Aroclor-1248 (2) [2C]			250	3.949349E-02								
Aroclor-1248 (3) [2C]			250	4.544987E-02								
Aroclor-1248 (4) [2C]			250	5.456762E-02								
Aroclor 1254 [2C]					250	0.0763106						
Aroclor-1254 (1) [2C]					250	6.080523E-02						
Aroclor-1254 (2) [2C]					250	4.891616E-02						
Aroclor-1254 (3) [2C]					250	0.1058376						
Aroclor-1254 (4) [2C]					250	0.103175						
Aroclor-1254 (5) [2C]					250	6.281905E-02						
Aroclor 1262 [2C]							250	7.397596E-02				
Aroclor-1262 (1) [2C]							250	6.830764E-02				
Aroclor-1262 (2) [2C]							250	5.817803E-02				



INITIAL CALIBRATION DATA

EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (2):	ZB35

Compound	Level 07		Level 08		Level 09		Level 10		Level 11		Level 12	
	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF	Conc	RRF
Aroclor-1262 (3) [2C]							250	6.600951E-02				
Aroclor-1262 (4) [2C]							250	0.1034087				
Aroclor 1268 [2C]									250	0.2386862		
Aroclor-1268 (1) [2C]									250	0.1610947		
Aroclor-1268 (2) [2C]									250	0.1731794		
Aroclor-1268 (3) [2C]									250	0.1478672		
Aroclor-1268 (4) [2C]									250	0.4726034		



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (2):	ZB35

COMPOUND	Mean RRF	RRF RSD	Linear COD	Quad COD	Limit Type & Limit	Q
Aroclor 1016 [2C]	5.458565E-02	6.3			RSD (20)	
Aroclor-1016 (1) [2C]	4.683127E-02	8.5			RSD (20)	
Aroclor-1016 (2) [2C]	9.496755E-02	8.0			RSD (20)	
Aroclor-1016 (3) [2C]	4.289222E-02	7.9			RSD (20)	
Aroclor-1016 (4) [2C]	3.365154E-02	10.3			RSD (20)	
Aroclor 1221 [2C]		0.0			RSD (20)	
Aroclor-1221 (1) [2C]		0.0			RSD (20)	
Aroclor-1221 (2) [2C]		0.0			RSD (20)	
Aroclor-1221 (3) [2C]		0.0			RSD (20)	
Aroclor 1232 [2C]		0.0			RSD (20)	
Aroclor-1232 (1) [2C]		0.0			RSD (20)	
Aroclor-1232 (2) [2C]		0.0			RSD (20)	
Aroclor-1232 (3) [2C]		0.0			RSD (20)	
Aroclor-1232 (4) [2C]		0.0			RSD (20)	
Aroclor 1242 [2C]		0.0			RSD (20)	
Aroclor-1242 (1) [2C]		0.0			RSD (20)	
Aroclor-1242 (2) [2C]		0.0			RSD (20)	
Aroclor-1242 (3) [2C]		0.0			RSD (20)	
Aroclor-1242 (4) [2C]		0.0			RSD (20)	
Aroclor 1248 [2C]		0.0			RSD (20)	
Aroclor-1248 (1) [2C]		0.0			RSD (20)	
Aroclor-1248 (2) [2C]		0.0			RSD (20)	
Aroclor-1248 (3) [2C]		0.0			RSD (20)	
Aroclor-1248 (4) [2C]		0.0			RSD (20)	
Aroclor 1254 [2C]		0.0			RSD (20)	
Aroclor-1254 (1) [2C]		0.0			RSD (20)	
Aroclor-1254 (2) [2C]		0.0			RSD (20)	
Aroclor-1254 (3) [2C]		0.0			RSD (20)	
Aroclor-1254 (4) [2C]		0.0			RSD (20)	
Aroclor-1254 (5) [2C]		0.0			RSD (20)	
Aroclor 1260 [2C]	6.996878E-02	6.4			RSD (20)	



INITIAL CALIBRATION DATA
EPA 8082A

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	GB00069	Instrument:	ECD7
Calibration Date:	02/24/2023	Column (2):	ZB35

COMPOUND	Mean RRF	RRF RSD	Linear COD	Quad COD	Limit Type & Limit	Q
Aroclor-1260 (1) [2C]	4.704064E-02	8.4			RSD (20)	
Aroclor-1260 (2) [2C]	0.1200523	7.6			RSD (20)	
Aroclor-1260 (3) [2C]	3.185902E-02	6.0			RSD (20)	
Aroclor-1260 (4) [2C]	8.092314E-02	5.1			RSD (20)	
Aroclor 1262 [2C]		0.0			RSD (20)	
Aroclor-1262 (1) [2C]		0.0			RSD (20)	
Aroclor-1262 (2) [2C]		0.0			RSD (20)	
Aroclor-1262 (3) [2C]		0.0			RSD (20)	
Aroclor-1262 (4) [2C]		0.0			RSD (20)	
Aroclor 1268 [2C]		0.0			RSD (20)	
Aroclor-1268 (1) [2C]		0.0			RSD (20)	
Aroclor-1268 (2) [2C]		0.0			RSD (20)	
Aroclor-1268 (3) [2C]		0.0			RSD (20)	
Aroclor-1268 (4) [2C]		0.0			RSD (20)	
Decachlorobiphenyl [2C]	1.218271	3.9			RSD (20)	
Tetrachlorometaxylene [2C]	1.173721	3.9			RSD (20)	



ANALYSIS SEQUENCE

SLB0342

Instrument: ECD7
Calibration ID: GB00069

Printed: 2/28/2023 9:54:44AM

Lab Number	Analysis	Container	Order	Position	STD ID	ISTD ID	Client	Comments
SLB0342-CAL1	QC		1		L000856	L000844		
SLB0342-CAL2	QC		2		L000859	L000844		
SLB0342-CAL3	QC		3		L000858	L000844		
SLB0342-CAL4	QC		4		L000731	L000844		
SLB0342-CAL5	QC		5		L000857	L000844		
SLB0342-CAL6	QC		6		L000855	L000844		
SLB0342-CAL7	QC		7		L000860	L000844		
SLB0342-CAL8	QC		8		L000861	L000844		
SLB0342-CAL9	QC		9		L000862	L000844		
SLB0342-CALA	QC		10		L000863	L000844		
SLB0342-CALB	QC		11		L000864	L000844		
SLB0342-SCV1	QC		12		L002065	L000844		
SLB0342-SCV2	QC		13		K007656	L000844		
SLB0342-SCV3	QC		14		L002066	L000844		
SLB0342-SCV4	QC		15		L002067	L000844		
SLB0342-SCV5	QC		16		L002068	L000844		
SLB0342-SCV6	QC		17		L002069	L000844		

Samples Loaded By Date

Data Processed By Date

GC LOG SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

	Inject	Date/Time	Filename	DF	LabID	ClientID
1	24-FEB-2023	10:51	02242301ECD7.D	1	IB	
2	24-FEB-2023	11:12	02242302ECD7.D	1	0.25PPMAR1660	
3	24-FEB-2023	11:33	02242303ECD7.D	1	0.02PPMAR1660	
4	24-FEB-2023	11:54	02242304ECD7.D	1	0.05PPMAR1660	
5	24-FEB-2023	12:15	02242305ECD7.D	1	1.0PPMAR1660	
6	24-FEB-2023	12:36	02242306ECD7.D	1	0.1PPMAR1660	
7	24-FEB-2023	12:57	02242307ECD7.D	1	0.5PPMAR1660	
8	24-FEB-2023	13:18	02242308ECD7.D	1	0.25PPMAR1242	
9	24-FEB-2023	13:39	02242309ECD7.D	1	0.25PPMAR1248	
10	24-FEB-2023	14:00	02242310ECD7.D	1	0.25PPMAR1254	
11	24-FEB-2023	14:21	02242311ECD7.D	1	0.25PPMAR2162	
12	24-FEB-2023	14:42	02242312ECD7.D	1	0.25PPMAR3268	
13	24-FEB-2023	15:03	02242313ECD7.D	1	AR1660SCV	
14	24-FEB-2023	15:24	02242314ECD7.D	1	AR1242SCV	
15	24-FEB-2023	15:45	02242315ECD7.D	1	AR1248SCV	
16	24-FEB-2023	16:06	02242316ECD7.D	1	AR1254SCV	
17	24-FEB-2023	16:27	02242317ECD7.D	1	AR2162SCV	
18	24-FEB-2023	16:48	02242318ECD7.D	1	AR3268SCV	
19	24-FEB-2023	17:09	02242319ECD7.D	1	DDTS	
20	24-FEB-2023	17:30	02242320ECD7.D	1	DDT BD	

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

ARI Job No.: IB Method: PCB.m Instrument: ecd7.i Date: 24-FEB-2023

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1051	02242301ECD7.D	IB		1	NO MANUAL INTEGRATION
1112	02242302ECD7.D	0.25PPMAR1660		1	NO MANUAL INTEGRATION
1133	02242303ECD7.D	0.02PPMAR1660		1	NO MANUAL INTEGRATION
1154	02242304ECD7.D	0.05PPMAR1660		1	NO MANUAL INTEGRATION
1215	02242305ECD7.D	1.0PPMAR1660		1	NO MANUAL INTEGRATION
1236	02242306ECD7.D	0.1PPMAR1660		1	NO MANUAL INTEGRATION
1257	02242307ECD7.D	0.5PPMAR1660		1	NO MANUAL INTEGRATION
1318	02242308ECD7.D	0.25PPMAR1242		1	NO MANUAL INTEGRATION
1339	02242309ECD7.D	0.25PPMAR1248		1	NO MANUAL INTEGRATION
1400	02242310ECD7.D	0.25PPMAR1254		1	NO MANUAL INTEGRATION
1421	02242311ECD7.D	0.25PPMAR2162		1	NO MANUAL INTEGRATION
1442	02242312ECD7.D	0.25PPMAR3268		1	NO MANUAL INTEGRATION
1503	02242313ECD7.D	AR1660SCV		1	NO MANUAL INTEGRATION
1524	02242314ECD7.D	AR1242SCV		1	NO MANUAL INTEGRATION
1545	02242315ECD7.D	AR1248SCV		1	NO MANUAL INTEGRATION
1606	02242316ECD7.D	AR1254SCV		1	NO MANUAL INTEGRATION
1627	02242317ECD7.D	AR2162SCV		1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1648	02242318ECD7.D	AR3268SCV		1	NO MANUAL INTEGRATION
1709	02242319ECD7.D	DDTS		1	NO MANUAL INTEGRATION
1730	02242320ECD7.D	DDT BD		1	NO MANUAL INTEGRATION
1751	02242321ECD7.D			1	NO MANUAL INTEGRATION
1812	02242322ECD7.D			1	NO MANUAL INTEGRATION
1833	02242323ECD7.D			1	NO MANUAL INTEGRATION
1854	02242324ECD7.D			1	NO MANUAL INTEGRATION
1915	02242325ECD7.D			1	NO MANUAL INTEGRATION
1936	02242326ECD7.D			1	NO MANUAL INTEGRATION
1957	02242327ECD7.D			1	NO MANUAL INTEGRATION
2018	02242328ECD7.D			1	NO MANUAL INTEGRATION
2039	02242329ECD7.D			1	NO MANUAL INTEGRATION
2059	02242330ECD7.D			1	NO MANUAL INTEGRATION
2120	02242331ECD7.D			1	NO MANUAL INTEGRATION
2141	02242332ECD7.D			1	NO MANUAL INTEGRATION
2202	02242333ECD7.D			1	NO MANUAL INTEGRATION
2223	02242334ECD7.D			1	NO MANUAL INTEGRATION
2244	02242335ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
2305	02242336ECD7.D			1	NO MANUAL INTEGRATION
2326	02242337ECD7.D			1	NO MANUAL INTEGRATION
2347	02242338ECD7.D			1	NO MANUAL INTEGRATION
0008	02242339ECD7.D			1	NO MANUAL INTEGRATION
0029	02242340ECD7.D			1	NO MANUAL INTEGRATION
0050	02242341ECD7.D			1	NO MANUAL INTEGRATION
0111	02242342ECD7.D			1	NO MANUAL INTEGRATION
0132	02242343ECD7.D			1	NO MANUAL INTEGRATION
0153	02242344ECD7.D			1	NO MANUAL INTEGRATION
0214	02242345ECD7.D			1	NO MANUAL INTEGRATION
0235	02242346ECD7.D			1	NO MANUAL INTEGRATION
0256	02242347ECD7.D			1	NO MANUAL INTEGRATION
0317	02242348ECD7.D			1	NO MANUAL INTEGRATION
0338	02242349ECD7.D			1	NO MANUAL INTEGRATION
0359	02242350ECD7.D			1	NO MANUAL INTEGRATION
0420	02242351ECD7.D			1	NO MANUAL INTEGRATION
0441	02242352ECD7.D			1	NO MANUAL INTEGRATION
0502	02242353ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0523	02242354ECD7.D			1	NO MANUAL INTEGRATION
0544	02242355ECD7.D			1	NO MANUAL INTEGRATION
0605	02242356ECD7.D			1	NO MANUAL INTEGRATION
0626	02242357ECD7.D			1	NO MANUAL INTEGRATION
0647	02242358ECD7.D			1	NO MANUAL INTEGRATION
0708	02242359ECD7.D			1	NO MANUAL INTEGRATION
0729	02242360ECD7.D			1	NO MANUAL INTEGRATION
0750	02242361ECD7.D			1	NO MANUAL INTEGRATION
0811	02242362ECD7.D			1	NO MANUAL INTEGRATION
0832	02242363ECD7.D			1	NO MANUAL INTEGRATION
0853	02242364ECD7.D			1	NO MANUAL INTEGRATION
0914	02242365ECD7.D			1	NO MANUAL INTEGRATION
0935	02242366ECD7.D			1	NO MANUAL INTEGRATION
0956	02242367ECD7.D			1	NO MANUAL INTEGRATION
1017	02242368ECD7.D			1	NO MANUAL INTEGRATION
1038	02242369ECD7.D			1	NO MANUAL INTEGRATION
1059	02242370ECD7.D			1	NO MANUAL INTEGRATION
1120	02242371ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1141	02242372ECD7.D			1	NO MANUAL INTEGRATION
1202	02242373ECD7.D			1	NO MANUAL INTEGRATION
1223	02242374ECD7.D			1	NO MANUAL INTEGRATION
1244	02242375ECD7.D			1	NO MANUAL INTEGRATION
1305	02242376ECD7.D			1	NO MANUAL INTEGRATION
1326	02242377ECD7.D			1	NO MANUAL INTEGRATION
1347	02242378ECD7.D			1	NO MANUAL INTEGRATION
1408	02242379ECD7.D			1	NO MANUAL INTEGRATION
1429	02242380ECD7.D			1	NO MANUAL INTEGRATION
1450	02242381ECD7.D			1	NO MANUAL INTEGRATION
1511	02242382ECD7.D			1	NO MANUAL INTEGRATION
1532	02242383ECD7.D			1	NO MANUAL INTEGRATION
1553	02242384ECD7.D			1	NO MANUAL INTEGRATION
1051	02242301ECD7.D IB			1	NO MANUAL INTEGRATION
1112	02242302ECD7.D 0.25PPMAR1660			1	NO MANUAL INTEGRATION
1133	02242303ECD7.D 0.02PPMAR1660			1	Aroclor-1016 [2C],
1154	02242304ECD7.D 0.05PPMAR1660			1	NO MANUAL INTEGRATION
1215	02242305ECD7.D 1.0PPMAR1660			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1236	02242306ECD7.D	0.1PPMAR1660		1	NO MANUAL INTEGRATION
1257	02242307ECD7.D	0.5PPMAR1660		1	NO MANUAL INTEGRATION
1318	02242308ECD7.D	0.25PPMAR1242		1	NO MANUAL INTEGRATION
1339	02242309ECD7.D	0.25PPMAR1248		1	NO MANUAL INTEGRATION
1400	02242310ECD7.D	0.25PPMAR1254		1	NO MANUAL INTEGRATION
1421	02242311ECD7.D	0.25PPMAR2162		1	NO MANUAL INTEGRATION
1442	02242312ECD7.D	0.25PPMAR3268		1	NO MANUAL INTEGRATION
1503	02242313ECD7.D	AR1660SCV		1	NO MANUAL INTEGRATION
1524	02242314ECD7.D	AR1242SCV		1	NO MANUAL INTEGRATION
1545	02242315ECD7.D	AR1248SCV		1	NO MANUAL INTEGRATION
1606	02242316ECD7.D	AR1254SCV		1	NO MANUAL INTEGRATION
1627	02242317ECD7.D	AR2162SCV		1	NO MANUAL INTEGRATION
1648	02242318ECD7.D	AR3268SCV		1	NO MANUAL INTEGRATION
1709	02242319ECD7.D	DDTS		1	NO MANUAL INTEGRATION
1730	02242320ECD7.D	DDT BD		1	NO MANUAL INTEGRATION
1751	02242321ECD7.D			1	NO MANUAL INTEGRATION
1812	02242322ECD7.D			1	NO MANUAL INTEGRATION
1833	02242323ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1854	02242324ECD7.D			1	NO MANUAL INTEGRATION
1915	02242325ECD7.D			1	NO MANUAL INTEGRATION
1936	02242326ECD7.D			1	NO MANUAL INTEGRATION
1957	02242327ECD7.D			1	NO MANUAL INTEGRATION
2018	02242328ECD7.D			1	NO MANUAL INTEGRATION
2038	02242329ECD7.D			1	NO MANUAL INTEGRATION
2059	02242330ECD7.D			1	NO MANUAL INTEGRATION
2120	02242331ECD7.D			1	NO MANUAL INTEGRATION
2141	02242332ECD7.D			1	NO MANUAL INTEGRATION
2202	02242333ECD7.D			1	NO MANUAL INTEGRATION
2223	02242334ECD7.D			1	NO MANUAL INTEGRATION
2244	02242335ECD7.D			1	NO MANUAL INTEGRATION
2305	02242336ECD7.D			1	NO MANUAL INTEGRATION
2326	02242337ECD7.D			1	NO MANUAL INTEGRATION
2347	02242338ECD7.D			1	NO MANUAL INTEGRATION
0008	02242339ECD7.D			1	NO MANUAL INTEGRATION
0029	02242340ECD7.D			1	NO MANUAL INTEGRATION
0050	02242341ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0111	02242342ECD7.D			1	NO MANUAL INTEGRATION
0132	02242343ECD7.D			1	NO MANUAL INTEGRATION
0153	02242344ECD7.D			1	NO MANUAL INTEGRATION
0214	02242345ECD7.D			1	NO MANUAL INTEGRATION
0235	02242346ECD7.D			1	NO MANUAL INTEGRATION
0256	02242347ECD7.D			1	NO MANUAL INTEGRATION
0317	02242348ECD7.D			1	NO MANUAL INTEGRATION
0338	02242349ECD7.D			1	NO MANUAL INTEGRATION
0359	02242350ECD7.D			1	NO MANUAL INTEGRATION
0420	02242351ECD7.D			1	NO MANUAL INTEGRATION
0441	02242352ECD7.D			1	NO MANUAL INTEGRATION
0502	02242353ECD7.D			1	NO MANUAL INTEGRATION
0523	02242354ECD7.D			1	NO MANUAL INTEGRATION
0544	02242355ECD7.D			1	NO MANUAL INTEGRATION
0605	02242356ECD7.D			1	NO MANUAL INTEGRATION
0626	02242357ECD7.D			1	NO MANUAL INTEGRATION
0647	02242358ECD7.D			1	NO MANUAL INTEGRATION
0708	02242359ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0729	02242360ECD7.D			1	NO MANUAL INTEGRATION
0750	02242361ECD7.D			1	NO MANUAL INTEGRATION
0811	02242362ECD7.D			1	NO MANUAL INTEGRATION
0832	02242363ECD7.D			1	NO MANUAL INTEGRATION
0853	02242364ECD7.D			1	NO MANUAL INTEGRATION
0914	02242365ECD7.D			1	NO MANUAL INTEGRATION
0935	02242366ECD7.D			1	NO MANUAL INTEGRATION
0956	02242367ECD7.D			1	NO MANUAL INTEGRATION
1017	02242368ECD7.D			1	NO MANUAL INTEGRATION
1038	02242369ECD7.D			1	NO MANUAL INTEGRATION
1059	02242370ECD7.D			1	NO MANUAL INTEGRATION
1120	02242371ECD7.D			1	NO MANUAL INTEGRATION
1141	02242372ECD7.D			1	NO MANUAL INTEGRATION
1202	02242373ECD7.D			1	NO MANUAL INTEGRATION
1223	02242374ECD7.D			1	NO MANUAL INTEGRATION
1244	02242375ECD7.D			1	NO MANUAL INTEGRATION
1305	02242376ECD7.D			1	NO MANUAL INTEGRATION
1326	02242377ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1347	02242378ECD7.D			1	NO MANUAL INTEGRATION
1408	02242379ECD7.D			1	NO MANUAL INTEGRATION
1429	02242380ECD7.D			1	NO MANUAL INTEGRATION
1450	02242381ECD7.D			1	NO MANUAL INTEGRATION
1511	02242382ECD7.D			1	NO MANUAL INTEGRATION
1532	02242383ECD7.D			1	NO MANUAL INTEGRATION
1553	02242384ECD7.D			1	NO MANUAL INTEGRATION

Security Status Report

Date: 28-Feb-2023 09:27

02242301ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242302ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242303ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242304ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242305ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242306ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242307ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242308ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242309ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242310ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242311ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242312ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242313ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242314ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242315ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242316ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242317ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242318ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242319ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27
02242320ECD7.D	Data Locked	richardl, 28-Feb-2023 09:27

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Calibration File Names:

Level 1: \\target\share\chem4\ecd7.i\230224.b\02242303ECD7.D
 Level 2: \\target\share\chem4\ecd7.i\230224.b\02242304ECD7.D
 Level 3: \\target\share\chem4\ecd7.i\230224.b\02242306ECD7.D
 Level 4: \\target\share\chem4\ecd7.i\230224.b\02242302ECD7.D
 Level 5: \\target\share\chem4\ecd7.i\230224.b\02242307ECD7.D
 Level 6: \\target\share\chem4\ecd7.i\230224.b\02242305ECD7.D
 Level 7: \\target\share\chem4\ecd7.i\230224.b\02242312ECD7.D

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
2 Aroclor-1221 (1)	+++++ 0.00716	+++++	+++++	+++++	+++++	+++++	0.00716	0.000
(2)	+++++ 0.01281	+++++	+++++	+++++	+++++	+++++	0.01281	0.000
(3)	+++++ 0.02975	+++++	+++++	+++++	+++++	+++++	0.02975	0.000
3 Aroclor-1242 (1)	+++++ 0.02479	+++++	+++++	+++++	+++++	+++++	0.02479	0.000
(2)	+++++ 0.07529	+++++	+++++	+++++	+++++	+++++	0.07529	0.000
(3)	+++++ 0.02343	+++++	+++++	+++++	+++++	+++++	0.02343	0.000
(4)	+++++ 0.03463	+++++	+++++	+++++	+++++	+++++	0.03463	0.000
4 Aroclor-1232 (1)	+++++ 0.00429	+++++	+++++	+++++	+++++	+++++	0.00429	0.000

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(2)	++++ 0.00850	++++	++++	++++	++++	++++	0.00850	0.000
(3)	++++ 0.03848	++++	++++	++++	++++	++++	0.03848	0.000
(4)	++++ 0.01635	++++	++++	++++	++++	++++	0.01635	0.000
7 Aroclor-1016(1)	0.03172 ++++	0.03253	0.03142	0.03141	0.02856	0.02667	0.03039	7.449
(2)	0.09239 ++++	0.09246	0.09222	0.09849	0.09174	0.08849	0.09263	3.499
(3)	0.05165 ++++	0.05037	0.04823	0.04393	0.03991	0.03721	0.04522	12.936
(4)	0.03002 ++++	0.02894	0.02959	0.03058	0.02852	0.02774	0.02923	3.542
6 Aroclor-1248(1)	++++ 0.03903	++++	++++	++++	++++	++++	0.03903	0.000
(2)	++++ 0.04961	++++	++++	++++	++++	++++	0.04961	0.000
(3)	++++ 0.09360	++++	++++	++++	++++	++++	0.09360	0.000
(4)	++++ 0.04765	++++	++++	++++	++++	++++	0.04765	0.000

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
8 Aroclor-1254 (1)	++++ 0.08033	++++	++++	++++	++++	++++	0.08033	0.000
(2)	++++ 0.03613	++++	++++	++++	++++	++++	0.03613	0.000
(3)	++++ 0.05165	++++	++++	++++	++++	++++	0.05165	0.000
(4)	++++ 0.10042	++++	++++	++++	++++	++++	0.10042	0.000
(5)	++++ 0.06294	++++	++++	++++	++++	++++	0.06294	0.000
9 Aroclor-1260 (1)	0.02926 ++++	0.02920	0.02841	0.03096	0.02737	0.02746	0.02878	4.677
(2)	0.02967 ++++	0.03006	0.03011	0.03291	0.02910	0.02857	0.03007	5.029
(3)	0.08088 ++++	0.08045	0.07954	0.08575	0.07515	0.07674	0.07975	4.627
(4)	0.03905 ++++	0.03887	0.03955	0.04485	0.03942	0.03922	0.04016	5.753
(5)	0.01783 ++++	0.01715	0.01679	0.01875	0.01664	0.01655	0.01729	4.953
10 Aroclor-1262 (1)	++++ 0.02454	++++	++++	++++	++++	++++	0.02454	0.000

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INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(2)	++++ 0.03993	++++	++++	++++	++++	++++	0.03993	0.000
(3)	++++ 0.04293	++++	++++	++++	++++	++++	0.04293	0.000
(4)	++++ 0.03923	++++	++++	++++	++++	++++	0.03923	0.000
11 Aroclor-1268(1)	++++ 0.10250	++++	++++	++++	++++	++++	0.10250	0.000
(2)	++++ 0.10151	++++	++++	++++	++++	++++	0.10151	0.000
(3)	++++ 0.08686	++++	++++	++++	++++	++++	0.08686	0.000
(4)	++++ 0.28598	++++	++++	++++	++++	++++	0.28598	0.000
42 2,4-DDE	++++ ++++	++++	++++	++++	++++	++++	++++	++++
43 2,4-DDD	++++ ++++	++++	++++	++++	++++	++++	++++	++++
44 2,4-DDT	++++ ++++	++++	++++	++++	++++	++++	++++	++++
46 4,4-DDE	++++ ++++	++++	++++	++++	++++	++++	++++	++++

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	250.000 Level 7	RRF	% RSD
47 4,4-DDD	++++	++++	++++	++++	++++	++++		++++	++++
48 4,4-DDT	++++	++++	++++	++++	++++	++++		++++	++++
49 Hexachlorobutadiene	++++	++++	++++	++++	++++	++++		++++	++++
50 Hexachlorobenzene	++++	++++	++++	++++	++++	++++		++++	++++
1 Tetrachloro-m-xylene	1.16827	1.24402	1.18546	1.20509	1.12295	1.24114		1.19449	3.860
13 Decachlorobiphenyl	0.82901	0.80558	0.77587	0.78808	0.73125	0.79742		0.78787	4.189

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INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Calibration File Names:

Level 1: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242303ECD7.D
 Level 2: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242304ECD7.D
 Level 3: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242306ECD7.D
 Level 4: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242302ECD7.D
 Level 5: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242307ECD7.D
 Level 6: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242305ECD7.D
 Level 7: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242312ECD7.D

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
1 Aroclor-1221 [2C] (1)	+++++	+++++	+++++	+++++	+++++	+++++	0.00757	0.000
(2)	0.00757						0.00757	0.000
(3)	0.01433						0.01433	0.000
4 Aroclor-1232 [2C] (1)	+++++	+++++	+++++	+++++	+++++	+++++	0.00409	0.000
(2)	0.00409						0.00409	0.000
(3)	0.02034						0.02034	0.000
(4)	0.04067						0.04067	0.000
3 Aroclor-1242 [2C] (1)	+++++	+++++	+++++	+++++	+++++	+++++	0.03717	0.000
(2)	0.03717						0.03717	0.000

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INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(2)	++++ 0.07813	++++	++++	++++	++++	++++	0.07813	0.000
(3)	++++ 0.02431	++++	++++	++++	++++	++++	0.02431	0.000
(4)	++++ 0.02962	++++	++++	++++	++++	++++	0.02962	0.000
6 Aroclor-1248 [2C] (1)	++++ 0.03820	++++	++++	++++	++++	++++	0.03820	0.000
(2)	++++ 0.03949	++++	++++	++++	++++	++++	0.03949	0.000
(3)	++++ 0.04545	++++	++++	++++	++++	++++	0.04545	0.000
(4)	++++ 0.05457	++++	++++	++++	++++	++++	0.05457	0.000
7 Aroclor-1016 [2C] (1)	0.05071 ++++	0.05022	0.04868	0.04733	0.04326	0.04080	0.04683	8.503
(2)	0.08143 ++++	0.09407	0.10159	0.10259	0.09651	0.09362	0.09497	8.025
(3)	0.04006 ++++	0.04718	0.04613	0.04410	0.04062	0.03926	0.04289	7.857
(4)	0.03181 ++++	0.03802	0.03707	0.03450	0.03115	0.02936	0.03365	10.251

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
8 Aroclor-1254 [2C] (1)	++++ 0.06081	++++	++++	++++	++++	++++	0.06081	0.000
(2)	++++ 0.04892	++++	++++	++++	++++	++++	0.04892	0.000
(3)	++++ 0.10584	++++	++++	++++	++++	++++	0.10584	0.000
(4)	++++ 0.10317	++++	++++	++++	++++	++++	0.10317	0.000
(5)	++++ 0.06282	++++	++++	++++	++++	++++	0.06282	0.000
10 Aroclor-1262 [2C] (1)	++++ 0.06831	++++	++++	++++	++++	++++	0.06831	0.000
(2)	++++ 0.05818	++++	++++	++++	++++	++++	0.05818	0.000
(3)	++++ 0.06601	++++	++++	++++	++++	++++	0.06601	0.000
(4)	++++ 0.10341	++++	++++	++++	++++	++++	0.10341	0.000
9 Aroclor-1260 [2C] (1)	0.05286 ++++	0.04911	0.04696	0.04801	0.04329	0.04201	0.04704	8.422
(2)	0.12976 ++++	0.12431	0.12095	0.12664	0.11320	0.10545	0.12005	7.605

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(3)	0.03524 ++++	0.03147	0.02937	0.03208	0.03102	0.03198	0.03186	6.045
(4)	0.08632 ++++	0.08237	0.08044	0.08393	0.07718	0.07531	0.08092	5.126
11 Aroclor-1268 [2C] (1)	++++ 0.16109	++++	++++	++++	++++	++++	0.16109	0.000
(2)	++++ 0.17318	++++	++++	++++	++++	++++	0.17318	0.000
(3)	++++ 0.14787	++++	++++	++++	++++	++++	0.14787	0.000
(4)	++++ 0.47260	++++	++++	++++	++++	++++	0.47260	0.000
41 2,4-DDE [2C]	++++ ++++	++++	++++	++++	++++	++++	++++	++++
42 2,4-DDD [2C]	++++ ++++	++++	++++	++++	++++	++++	++++	++++
44 4,4-DDE [2C]	++++ ++++	++++	++++	++++	++++	++++	++++	++++
45 4,4-DDD/2,4-DDT [2C]	++++ ++++	++++	++++	++++	++++	++++	++++	++++
46 4,4-DDT [2C]	++++ ++++	++++	++++	++++	++++	++++	++++	++++

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
48 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
49 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
\$ 2 Tetrachloro-m-xylene [2C]	1.21526 +++++	1.19545	1.17555	1.21907	1.12560	1.11139	1.17372	3.897
\$ 13 Decachlorobiphenyl [2C]	1.17066 +++++	1.20406	1.20549	1.31040	1.21104	1.20797	1.21827	3.898

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Batch File: \\target\share\chem4\ecd7.i\230224.b
Inst ID: ecd7.i

ID:	RT01	RT02	RT03	RT04	RT05	RT06
FILENAME:	02242302ECD7	02242303ECD7	02242304ECD7	02242305ECD7	02242306ECD7	02242307ECD7
INJ. DATE:	24-FEB-2023	24-FEB-2023	24-FEB-2023	24-FEB-2023	24-FEB-2023	24-FEB-2023
INJ. TIME:	11:12	11:33	11:54	12:15	12:36	12:57

Compound	RT01	RT02	RT03	RT04	RT05	RT06	EXPEC RT	RT WINDOW	AVG RT	STD DEV
* 41 IS-BNB	3.493	3.492	3.492	3.492	3.491	3.491	3.493	3.393-3.593	3.492	0.001
§ 1 Tetrachloro-m-xylene	5.811	5.809	5.809	5.813	5.809	5.810	5.811	5.711-5.911	5.810	0.002
2 Aroclor-1221	+++++	+++++	+++++	+++++	+++++	+++++	4.732	4.632-4.832	+++++	+++++
3 Aroclor-1242	+++++	+++++	+++++	+++++	+++++	+++++	7.269	7.169-7.369	+++++	+++++
4 Aroclor-1232	+++++	+++++	+++++	+++++	+++++	+++++	4.732	4.632-4.832	+++++	+++++
7 Aroclor-1016	7.272	7.272	7.272	7.270	7.271	7.270	7.272	7.172-7.372	7.271	0.001
6 Aroclor-1248	+++++	+++++	+++++	+++++	+++++	+++++	8.403	8.303-8.503	+++++	+++++
8 Aroclor-1254	+++++	+++++	+++++	+++++	+++++	+++++	9.295	9.195-9.395	+++++	+++++
9 Aroclor-1260	11.046	11.047	11.046	11.044	11.045	11.044	11.046	10.946-11.146	11.045	0.001
10 Aroclor-1262	+++++	+++++	+++++	+++++	+++++	+++++	10.824	10.724-10.924	+++++	+++++
11 Aroclor-1268	+++++	+++++	+++++	+++++	+++++	+++++	12.243	12.143-12.343	+++++	+++++
§ 13 Decachlorobiphenyl	13.897	13.893	13.893	13.899	13.892	13.898	13.897	13.797-13.997	13.895	0.003
* 12 IS-HBBP	14.269	14.268	14.268	14.267	14.268	14.268	14.269	14.169-14.369	14.268	0.001
42 2,4-DDE	+++++	+++++	+++++	+++++	+++++	+++++	9.260	9.210-9.310	+++++	+++++
43 2,4-DDD	+++++	+++++	+++++	+++++	+++++	+++++	9.801	9.751-9.851	+++++	+++++
44 2,4-DDT	+++++	+++++	+++++	+++++	+++++	+++++	10.293	10.243-10.343	+++++	+++++
46 4,4-DDE	+++++	+++++	+++++	+++++	+++++	+++++	9.683	9.583-9.783	+++++	+++++

Reviewer 1 _____ Date: _____
Reviewer 2 _____ Date: _____

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Batch File: \\target\share\chem4\ecd7.i\230224.b
 Inst ID: ecd7.i

Compound	RT01	RT02	RT03	RT04	RT05	RT06	EXPEC RT	RT WINDOW	AVG RT	STD DEV
47 4,4-DDD	+++++	+++++	+++++	+++++	+++++	+++++	10.254	10.154-10.354	+++++	+++++
48 4,4-DDT	+++++	+++++	+++++	+++++	+++++	+++++	10.754	10.654-10.854	+++++	+++++
49 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	1.842	1.742-1.942	+++++	+++++
50 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	6.708	6.608-6.808	+++++	+++++

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
Batch File: \\target\share\chem4\ecd7.i\230224.b\230224.b
Inst ID: ecd7.i

ID: RT01 RT02 RT03 RT04 RT05 RT06
FILENAME: 02242302ECD7 02242303ECD7 02242304ECD7 02242305ECD7 02242306ECD7 02242307ECD7
INJ. DATE: 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023
INJ. TIME: 11:12 11:33 11:54 12:15 12:36 12:57

Table with 11 columns: Compound, RT01, RT02, RT03, RT04, RT05, RT06, EXPEC RT, RT WINDOW, AVG RT, STD DEV. Rows include various chemical compounds like IS-BNB, Tetrachloro-m-xylene, Aroclor-1221, etc.

Reviewer 1 _____ Date: _____
Reviewer 2 _____ Date: _____

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Batch File: \\target\share\chem4\ecd7.i\230224.b\230224.b
 Inst ID: ecd7.i

Compound	RT01	RT02	RT03	RT04	RT05	RT06	EXPEC RT	RT WINDOW	AVG RT	STD DEV
46 4,4-DDT [2C]	+++++	+++++	+++++	+++++	+++++	+++++	11.092	10.992-11.192	+++++	+++++
48 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	1.703	1.603-1.803	+++++	+++++
49 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	7.178	7.078-7.278	+++++	+++++

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242301ECD7.D
Data file 2: /230224.b/230224.b/02242301ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: IB
Client ID:
Injection Date: 24-FEB-2023 10:51
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.826	0.019	382217	5.683	-0.002	180378	33.8	36.5	7.7	Tetrachloro-m-xylene
13.904	0.011	534110	14.120	0.001	295605	35.3	37.2	5.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	756896	12.3
Hexabromobiphenyl	1429847	1534275	7.3

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	336543	6.8
Hexabromobiphenyl	513946	521508	1.5

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	---			0.0	1	---			0.0
Aroclor-1016	2	---			0.0	2	---			0.0
Aroclor-1016	3	---			0.0	3	---			0.0
Aroclor-1016	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	---			0.0	2	6.321	0.025	1873	31.1
Aroclor-1221	3	---			0.0	3	6.633	0.012	314	3.2
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	---			0.0	2	---			0.0
Aroclor-1232	3	7.698	0.043	2193	6.0	3	---			0.0
Aroclor-1232	4	8.505	-0.076	11525	74.5	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1242	1	---			0.0	1	---			0.0
Aroclor-1242	2	7.698	0.042	2193	3.1	2	---			0.0
Aroclor-1242	3	---			0.0	3	---			0.0
Aroclor-1242	4	8.505	-0.074	11525	35.2	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1248	1	---			0.0	1	---			0.0
Aroclor-1248	2	---			0.0	2	---			0.0
Aroclor-1248	3	---			0.0	3	---			0.0
Aroclor-1248	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1254	1	---			0.0	1	---			0.0
Aroclor-1254	2	---			0.0	2	---			0.0
Aroclor-1254	3	9.596	-0.072	31424	64.3	3	---			0.0
Aroclor-1254	4	---			0.0	4	---			0.0
Aroclor-1254	5	10.167	-0.010	18361	30.8	5	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1260	1	11.098	0.054	6994	12.7	1	---			0.0
Aroclor-1260	2	---			0.0	2	---			0.0
Aroclor-1260	3	11.706	-0.027	7806	5.1	3	---			0.0
Aroclor-1260	4	---			0.0	4	---			0.0
Aroclor-1260	5	---			0.0	NS	---			----
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1262	1	10.824	-0.005	16873	35.8	1	---			0.0
Aroclor-1262	2	---			0.0	2	---			0.0
Aroclor-1262	3	---			0.0	3	---			0.0
Aroclor-1262	4	13.040	0.053	14031	18.6	4	---			0.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1268	1	---			0.0	1	---			0.0
Aroclor-1268	2	---			0.0	2	---			0.0
Aroclor-1268	3	12.709	0.009	6037	3.6	3	12.891	-0.001	659	0.7
Aroclor-1268	4	13.499	0.010	12396	2.3	4	13.710	0.001	1848	0.6
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				

Total PCB Area Coll (5.906 - 13.793) = 260205

Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 18252 Col2 Total PCB = 0.0 ppm*

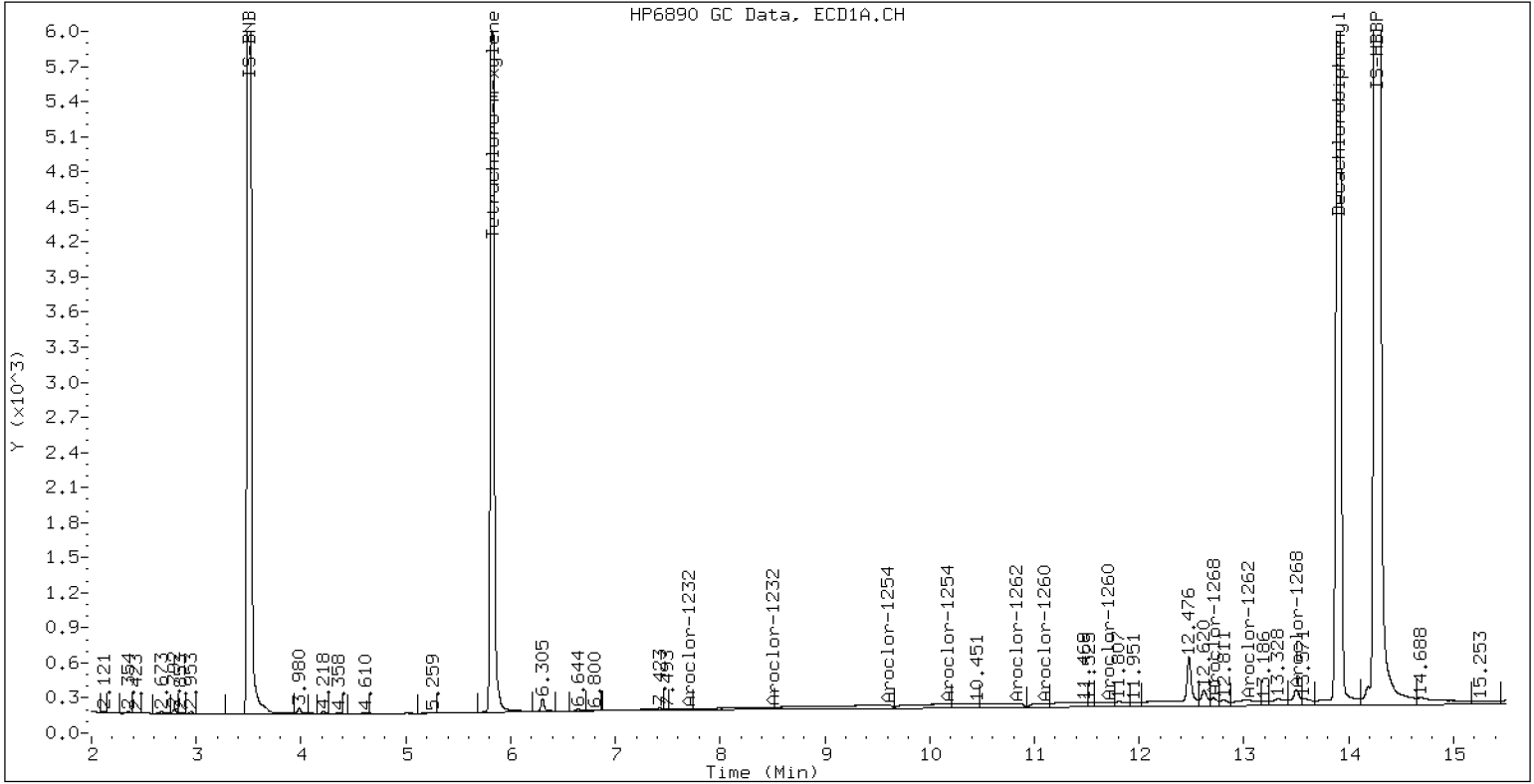
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 IB

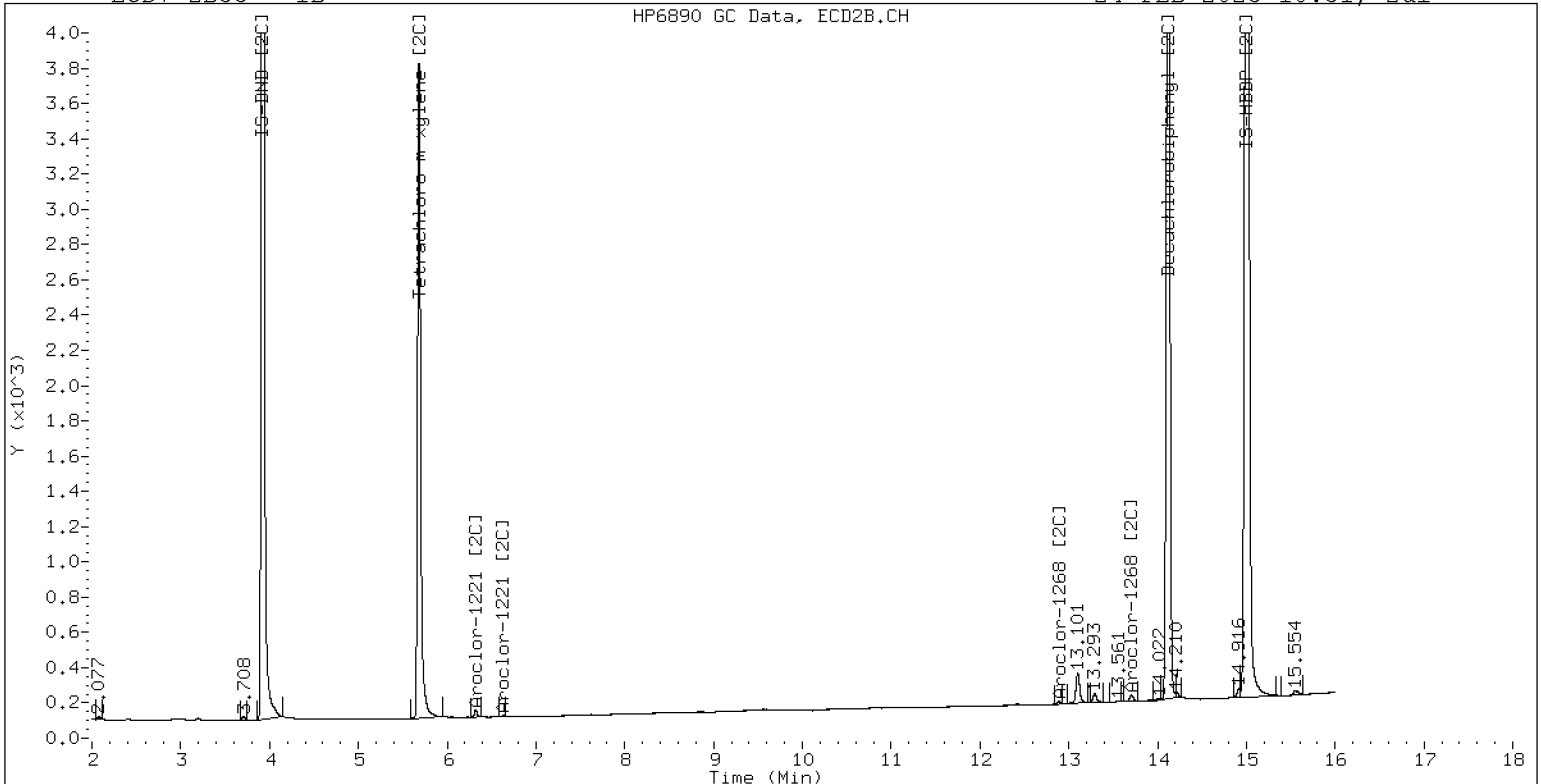
24-FEB-2023 10:51, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 IB

24-FEB-2023 10:51, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242302ECD7.D
Data file 2: /230224.b/230224.b/02242302ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 11:12
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	0.005	405980	5.687	0.002	192160	40.4	41.5	2.9	Tetrachloro-m-xylene
13.897	0.004	563414	14.120	0.001	336737	40.0	43.0	7.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	673778	0.0
Hexabromobiphenyl	1429847	1429847	0.0

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	315256	0.0
Hexabromobiphenyl	513946	513946	0.0

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.272	0.001	66125	258.4	1	7.255	-0.001	46626	252.6
Aroclor-1016	2	7.654	-0.000	207370	265.8	2	7.855	-0.001	101071	270.1
Aroclor-1016	3	7.792	0.002	92507	242.9	3	8.055	0.001	43448	257.1
Aroclor-1016	4	8.406	0.001	64388	261.5	4	8.306	-0.000	33986	256.3
Total CollAve (4 peaks):				257.2		Total Col2Ave (4 peaks):				259.0 RPD = 1
Corrected Ave (3 peaks):				254.3		Corrected Ave (3 peaks):				255.3 RPD = 0

CalAmt %D: 2.9

CalAmt %D: 3.6

Aroclor-1260	1	11.046	0.001	138355	269.0	1	11.653	0.001	77114	255.2
Aroclor-1260	2	11.363	0.002	147051	273.6	2	11.918	0.001	203401	263.7
Aroclor-1260	3	11.736	0.003	383171	268.8	3	12.435	-0.000	51517	251.7
Aroclor-1260	4	12.141	0.002	200399	279.2	4	12.502	0.001	134797	259.3
Aroclor-1260	5	12.247	0.003	83796	271.2	NS	---			----
Total CollAve (5 peaks):				272.4		Total Col2Ave (4 peaks):				257.5 RPD = 6
Corrected Ave (4 peaks):				270.7		Corrected Ave (3 peaks):				255.4 RPD = 6

CalAmt %D: 8.9

CalAmt %D: 3.0

Total PCB Area Coll (5.906 - 13.793) = 4024419 Coll Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1889311 Col2 Total PCB = 0.5 ppm*

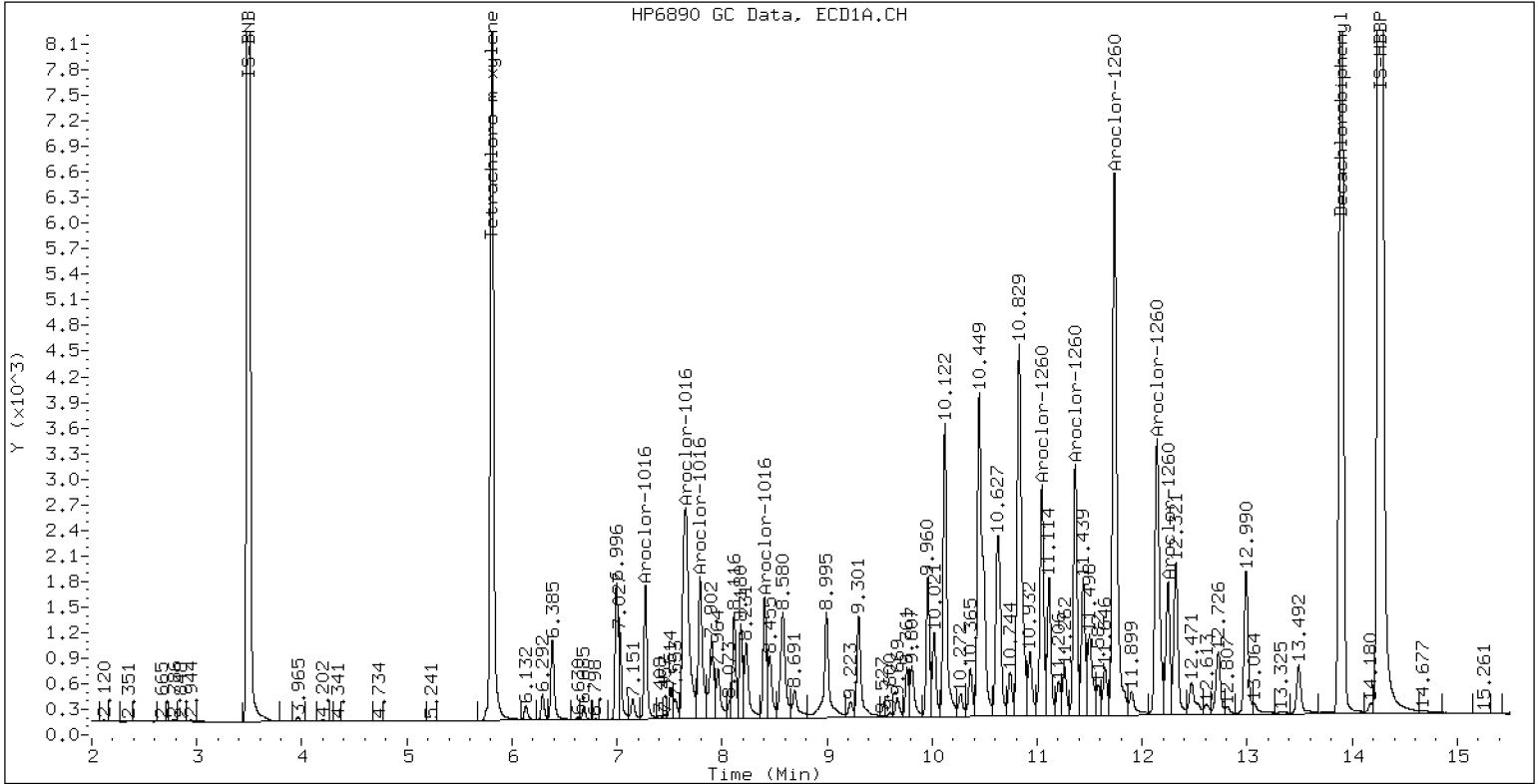
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1660

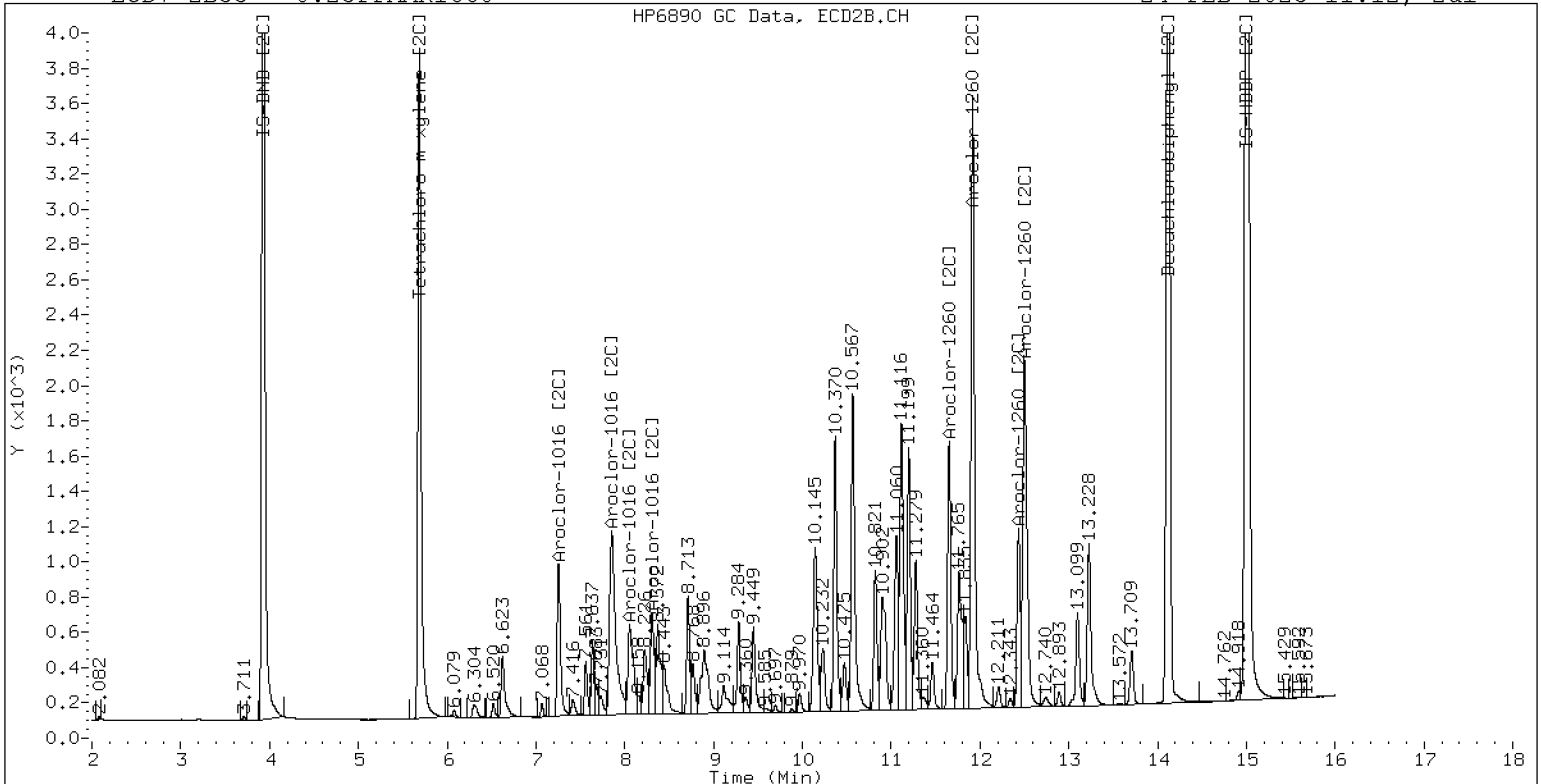
24-FEB-2023 11:12, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1660

24-FEB-2023 11:12, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242303ECD7.D
Data file 2: /230224.b/230224.b/02242303ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.02PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 11:33
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.002	29768	5.688	0.003	14932	3.1	3.3	5.7	Tetrachloro-m-xylene
13.893	0.000	45992	14.120	0.000	23950	3.4	3.1	9.1	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	637010	-5.5
Hexabromobiphenyl	1429847	1386953	-3.0

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	307177	-2.6
Hexabromobiphenyl	513946	511463	-0.5

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.272	0.001	5052	20.9	1	7.256	0.000	3894	21.7	
Aroclor-1016	2	7.659	0.005	14714	19.9	2	7.864	0.008	6253	17.1	
Aroclor-1016	3	7.795	0.005	8226	22.8	3	8.060	0.006	3076	18.7	
Aroclor-1016	4	8.407	0.002	4780	20.5	4	8.309	0.002	2443	18.9	
Total CollAve (4 peaks):				21.1	Total Col2Ave (4 peaks):				19.1	RPD = 10	
Corrected Ave (3 peaks):				20.5	Corrected Ave (3 peaks):				18.2	RPD = 11	
CalAmt %D:				5.3	CalAmt %D:				-4.5		
Aroclor-1260	1	11.047	0.003	10147	20.3	1	11.656	0.003	6759	22.5	
Aroclor-1260	2	11.364	0.003	10287	19.7	2	11.922	0.005	16592	21.6	
Aroclor-1260	3	11.740	0.006	28043	20.3	3	12.438	0.002	4506	22.1	
Aroclor-1260	4	12.145	0.006	13540	19.4	4	12.505	0.004	11037	21.3	
Aroclor-1260	5	12.246	0.002	6182	20.6	NS	---			----	
Total CollAve (5 peaks):				20.1	Total Col2Ave (4 peaks):				21.9	RPD = 9	
Corrected Ave (4 peaks):				19.9	Corrected Ave (3 peaks):				21.7	RPD = 8	
CalAmt %D:				0.4	CalAmt %D:				9.4		

Total PCB Area Coll (5.906 - 13.793) = 324832 Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 157149 Col2 Total PCB = 0.0 ppm*

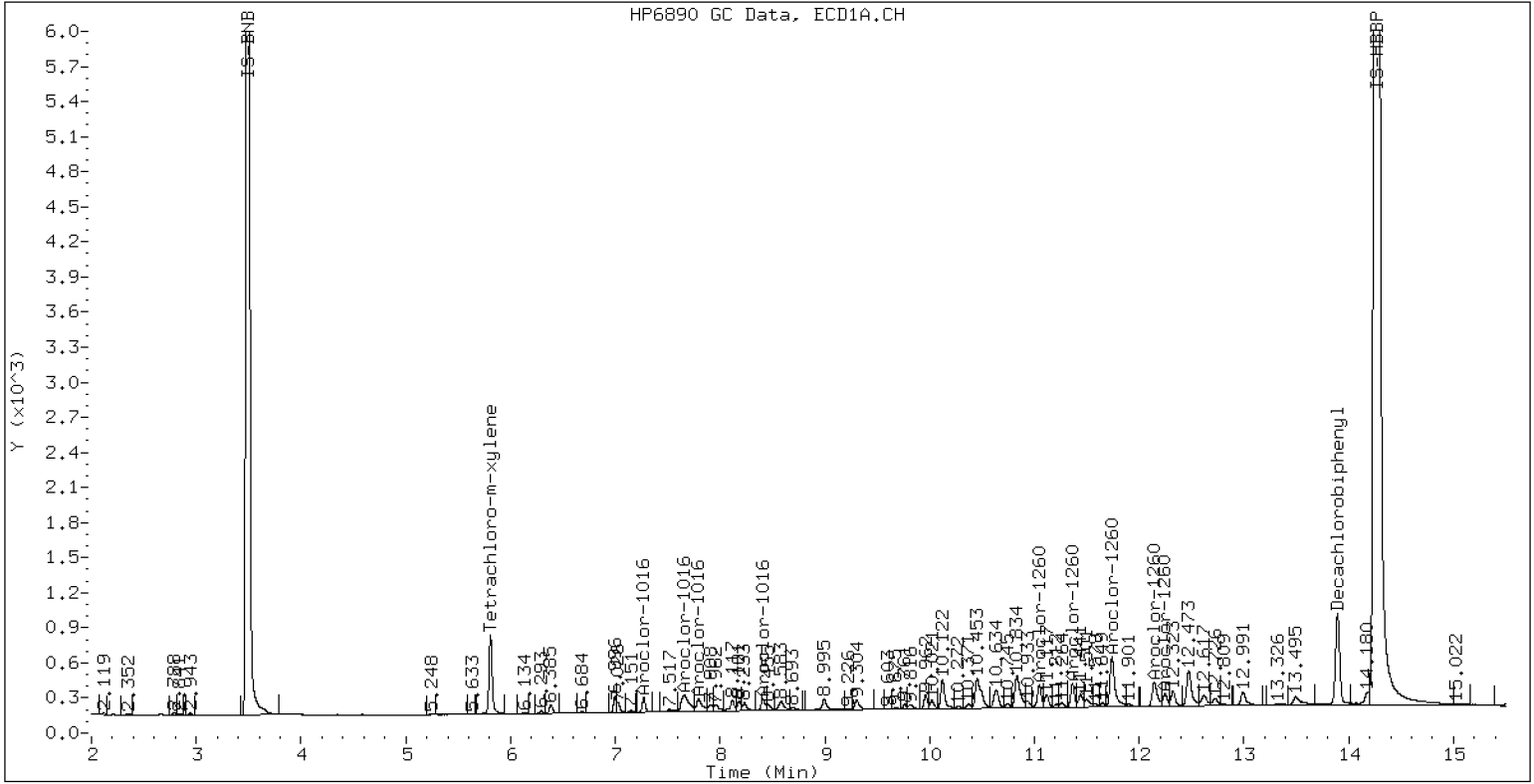
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.02PPMAR1660

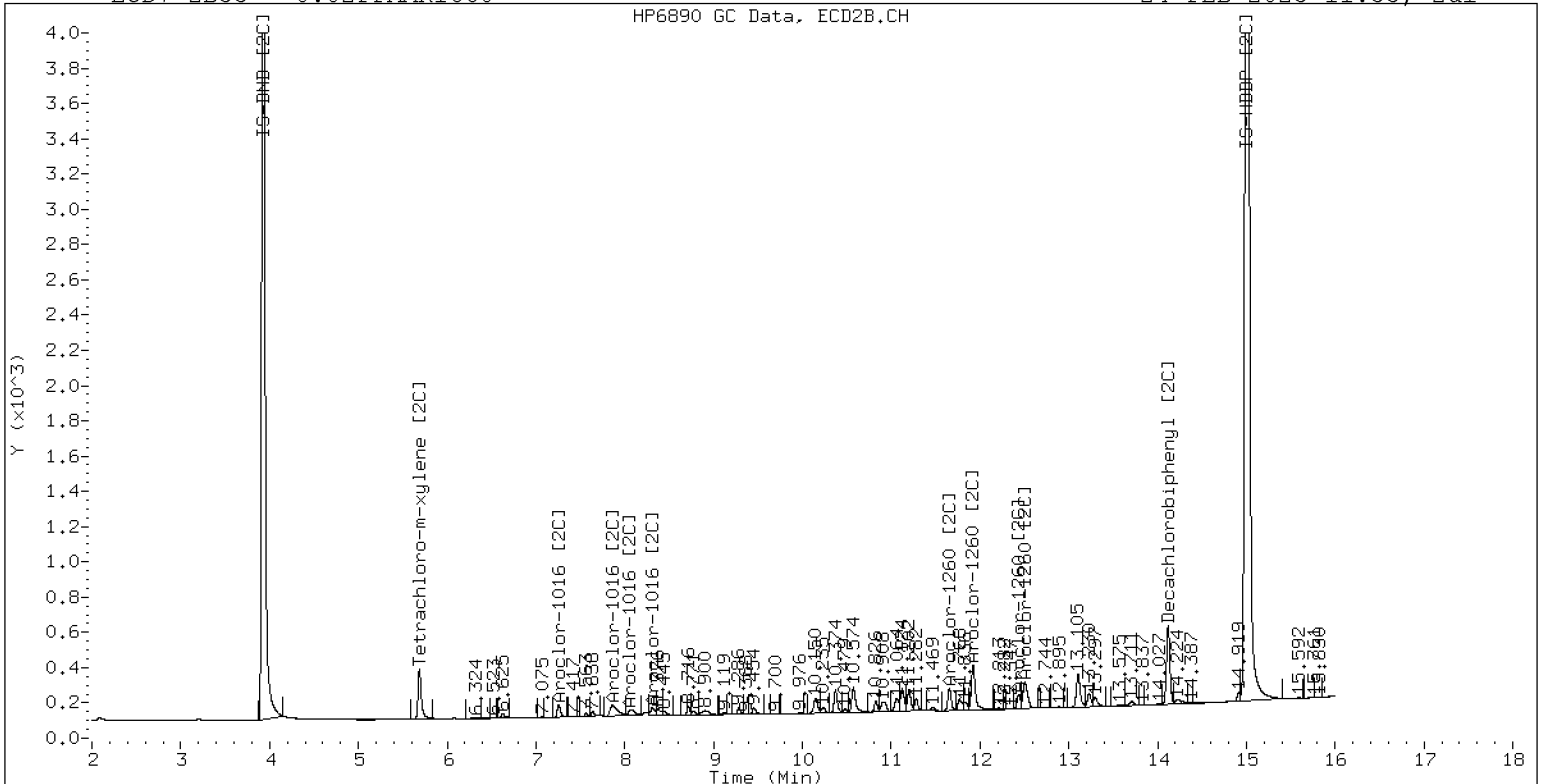
24-FEB-2023 11:33, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.02PPMAR1660

24-FEB-2023 11:33, 2ul

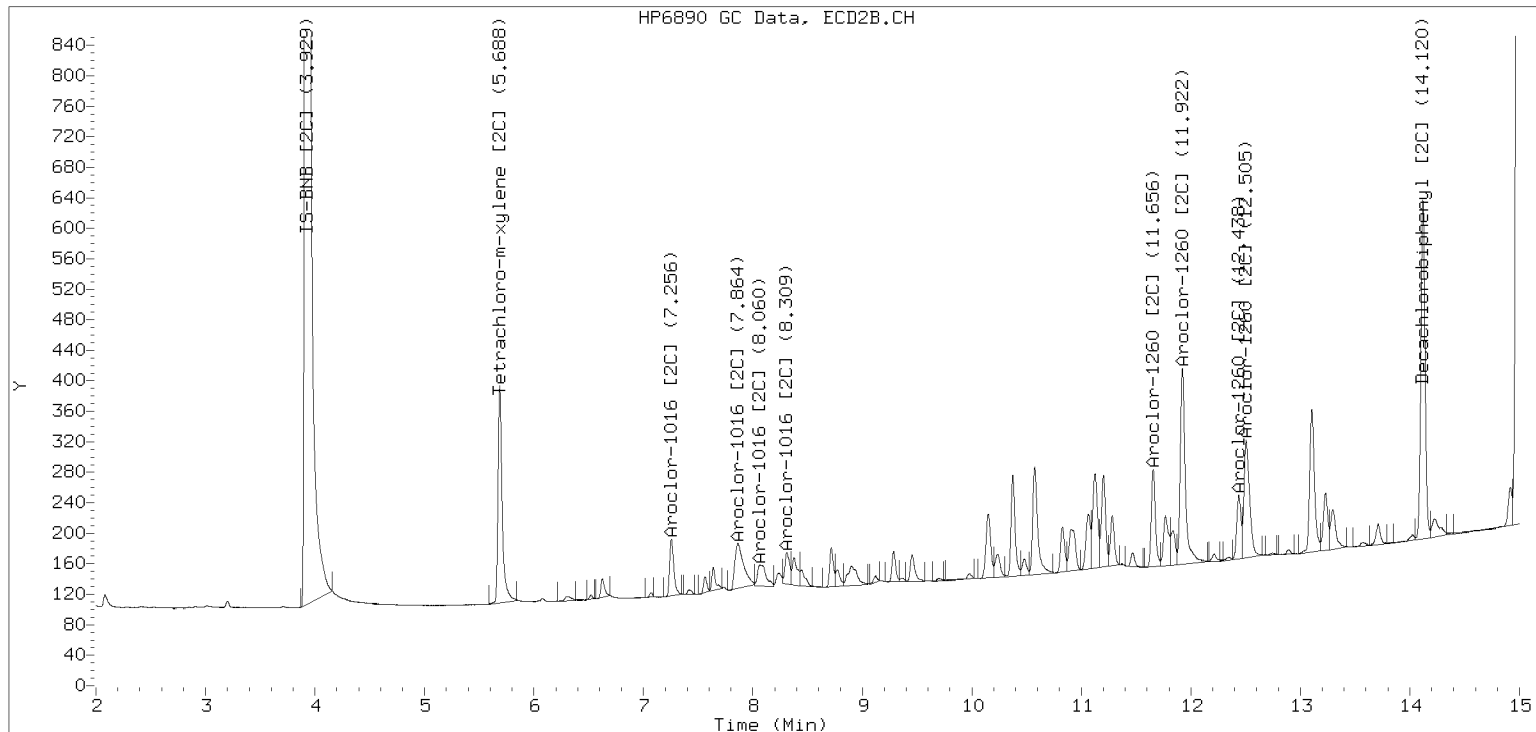


ZB-35 Manual Integration: YES

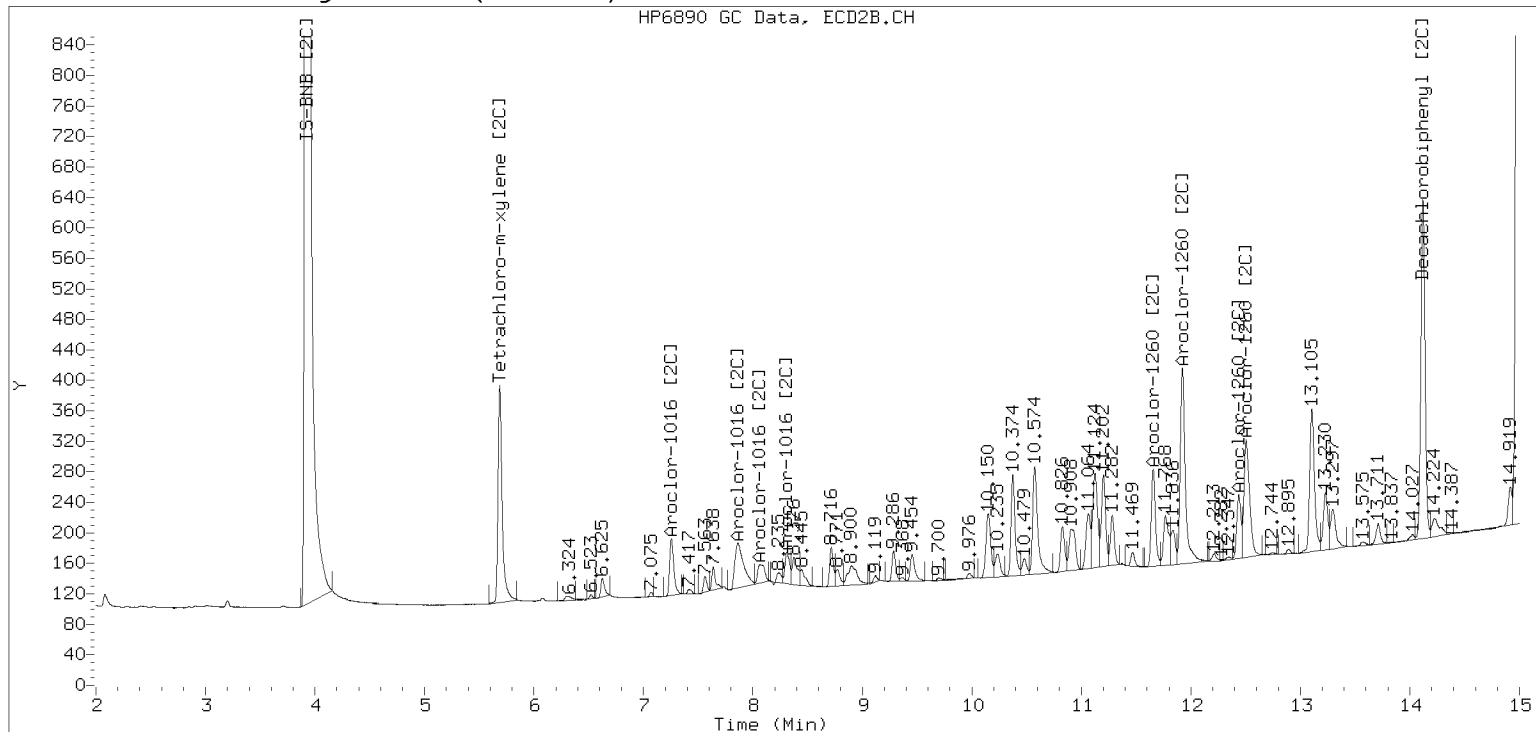
Manual Peak Adjustment, ZB-35

Datafile: ecd7.i/230224.b/230224.b/02242303ECD7.D Injection Date: 24-FEB-2023

Manual Integration (After)



Processed Integration (Before)



Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242304ECD7.D
Data file 2: /230224.b/230224.b/02242304ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.05PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 11:54
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col		ZB5	ZB35	RPD	Compound/Flag		
RT	Shift Response	RT	Shift Response	on col	on col				
5.809	0.003	78493	5.688	0.003	36772	8.3	8.1	2.2	Tetrachloro-m-xylene
13.893	-0.000	113544	14.119	-0.000	62745	8.2	7.9	3.4	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	630965	-6.4
Hexabromobiphenyl	1429847	1409464	-1.4

Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	307599	-2.4
Hexabromobiphenyl	513946	521112	1.4

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.272	0.002	12829	53.5	1	7.256	0.000	9654	53.6	
Aroclor-1016	2	7.660	0.006	36461	49.9	2	7.864	0.008	18085	49.5	
Aroclor-1016	3	7.795	0.005	19865	55.7	3	8.063	0.008	9071	55.0	
Aroclor-1016	4	8.408	0.003	11411	49.5	4	8.310	0.003	7309	56.5	
Total CollAve (4 peaks):				52.2	Total Col2Ave (4 peaks):				53.7	RPD = 3	
Corrected Ave (3 peaks):				51.0	Corrected Ave (3 peaks):				52.7	RPD = 3	
CalAmt %D:				4.3	CalAmt %D:				7.3		
Aroclor-1260	1	11.046	0.002	25727	50.7	1	11.655	0.002	15996	52.2	
Aroclor-1260	2	11.363	0.002	26482	50.0	2	11.922	0.004	40487	51.8	
Aroclor-1260	3	11.739	0.005	70871	50.4	3	12.437	0.002	10248	49.4	
Aroclor-1260	4	12.143	0.004	34239	48.4	4	12.506	0.004	26828	50.9	
Aroclor-1260	5	12.246	0.002	15109	49.6	NS	---			----	
Total CollAve (5 peaks):				49.8	Total Col2Ave (4 peaks):				51.1	RPD = 2	
Corrected Ave (4 peaks):				49.6	Corrected Ave (3 peaks):				50.7	RPD = 2	
CalAmt %D:				-0.3	CalAmt %D:				2.1		

Total PCB Area Coll (5.906 - 13.793) = 758292 Coll Total PCB = 0.1 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 386383 Col2 Total PCB = 0.1 ppm*

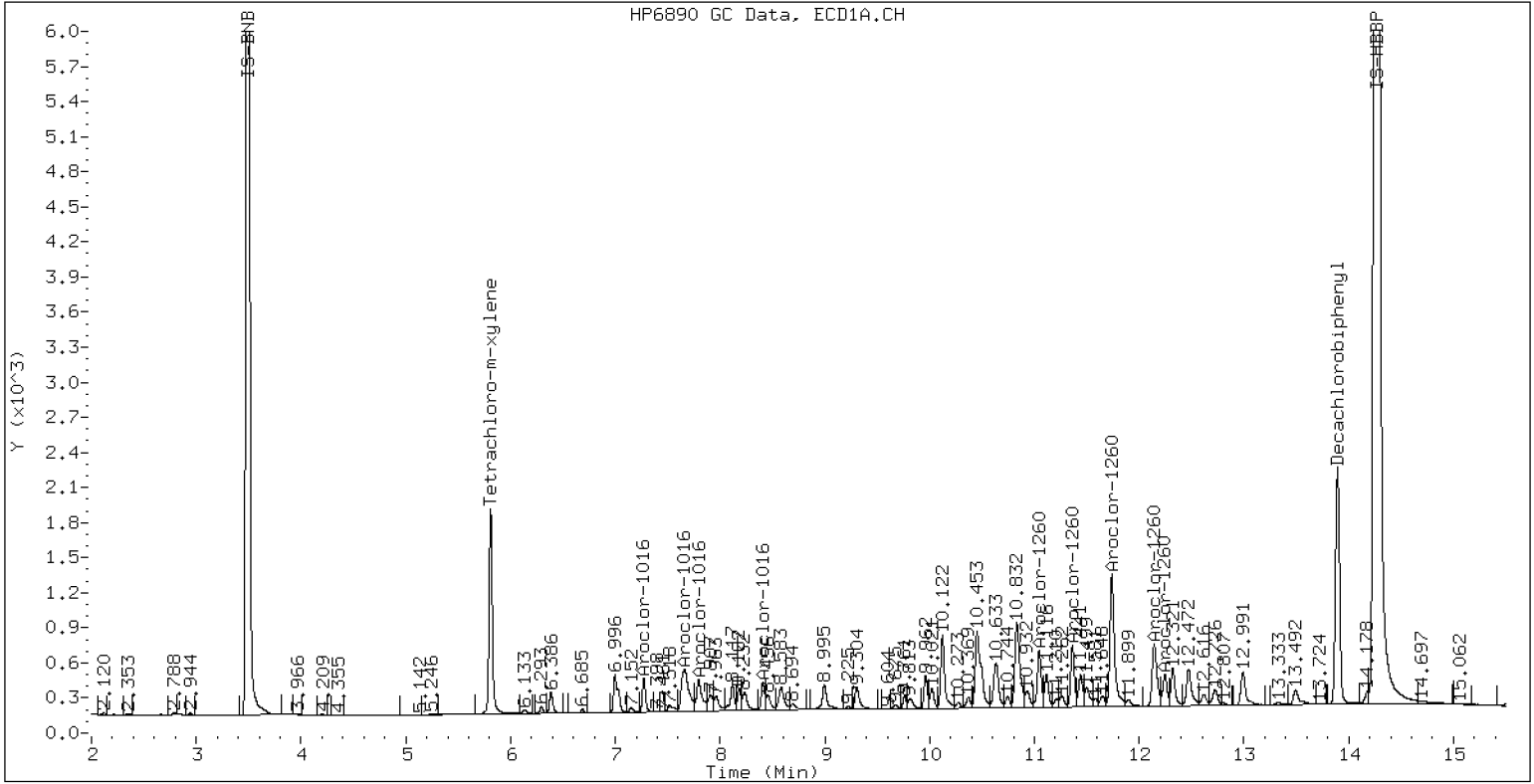
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.05PPMAR1660

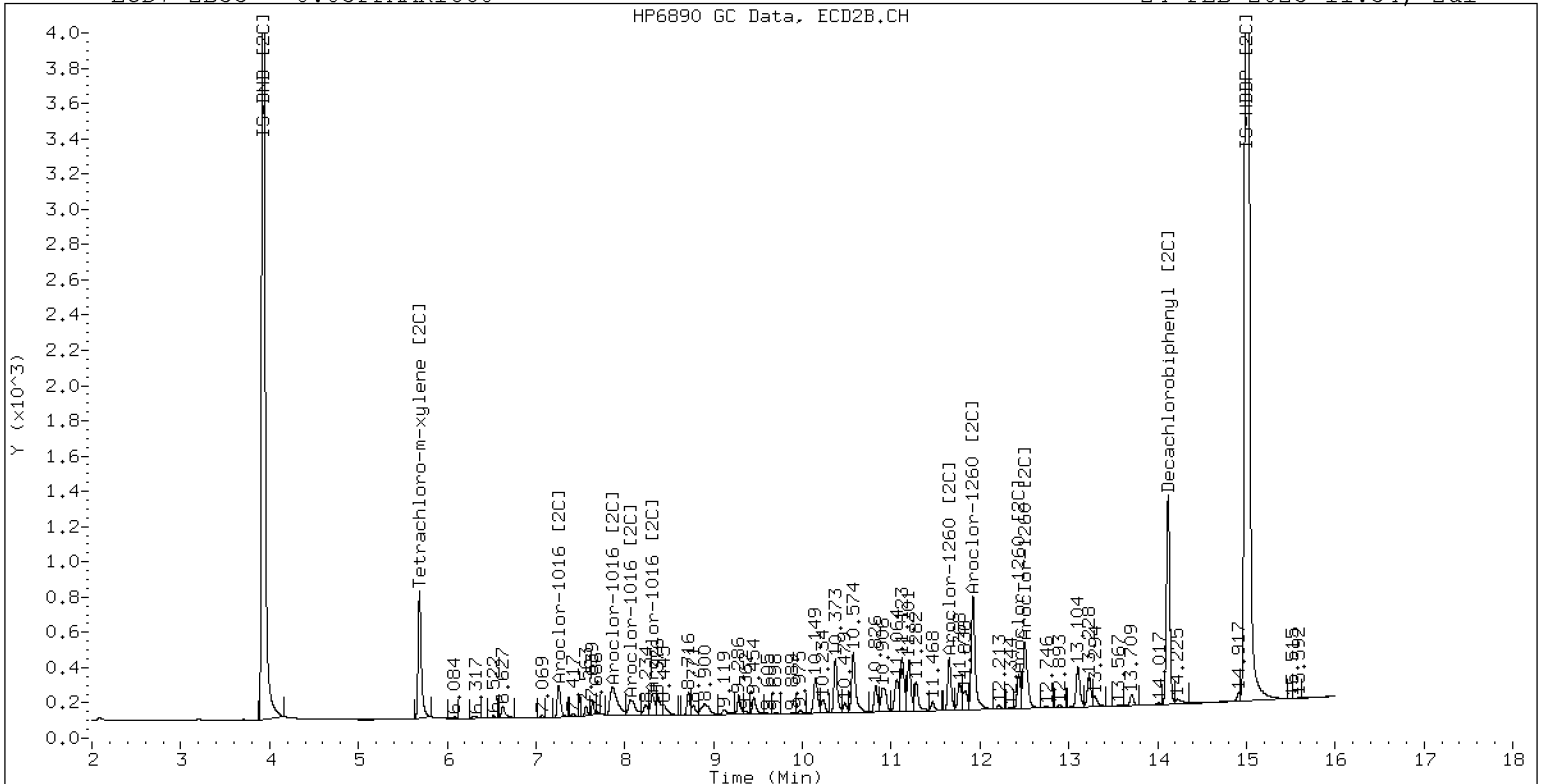
24-FEB-2023 11:54, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.05PPMAR1660

24-FEB-2023 11:54, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242305ECD7.D
Data file 2: /230224.b/230224.b/02242305ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 1.0PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 12:15
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.813	0.006	1641874	5.688	0.003	709674	166.2	151.5	9.3	Tetrachloro-m-xylene
13.899	0.006	2344583	14.122	0.002	1300114	161.9	158.6	2.1	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	661440	-1.8
Hexabromobiphenyl	1429847	1470100	2.8
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	319272	1.3
Hexabromobiphenyl	513946	538138	4.7

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.270	-0.000	220519	877.8	1	7.254	-0.001	162833	871.2
Aroclor-1016	2	7.652	-0.002	731607	955.3	2	7.852	-0.004	373610	985.8
Aroclor-1016	3	7.789	-0.001	307629	822.8	3	8.051	-0.003	156666	915.2
Aroclor-1016	4	8.404	-0.001	229387	949.1	4	8.305	-0.002	117186	872.6
Total CollAve (4 peaks):				901.3		Total Col2Ave (4 peaks):				911.2 RPD = 1
Corrected Ave (3 peaks):				883.3		Corrected Ave (3 peaks):				886.3 RPD = 0

CalAmt %D: -9.9

CalAmt %D: -8.9

Aroclor-1260	1	11.044	-0.000	504641	954.2	1	11.652	-0.000	282606	893.1
Aroclor-1260	2	11.360	-0.001	524931	950.0	2	11.917	-0.000	709329	878.4
Aroclor-1260	3	11.734	-0.000	1410270	962.3	3	12.434	-0.001	215124	1003.8
Aroclor-1260	4	12.137	-0.002	720770	976.7	4	12.501	-0.001	506566	930.6
Aroclor-1260	5	12.243	-0.001	304211	957.7	NS	---			----
Total CollAve (5 peaks):				960.2		Total Col2Ave (4 peaks):				926.5 RPD = 4
Corrected Ave (4 peaks):				956.0		Corrected Ave (3 peaks):				900.7 RPD = 6

CalAmt %D: -4.0

CalAmt %D: -7.4

Total PCB Area Coll (5.906 - 13.793) = 14454279 Coll Total PCB = 1.8 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 7029563 Col2 Total PCB = 1.8 ppm*

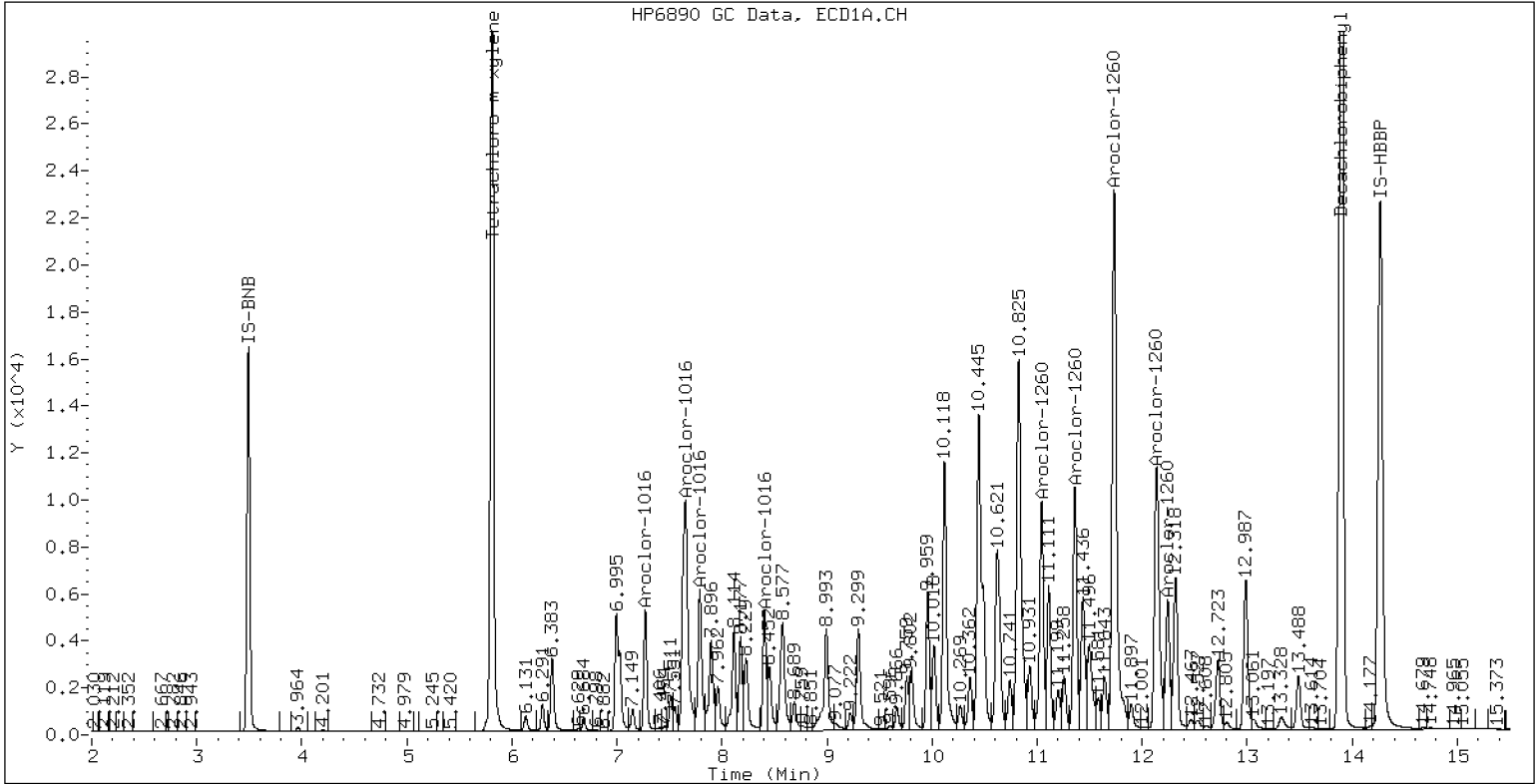
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 1.0PPMAR1660

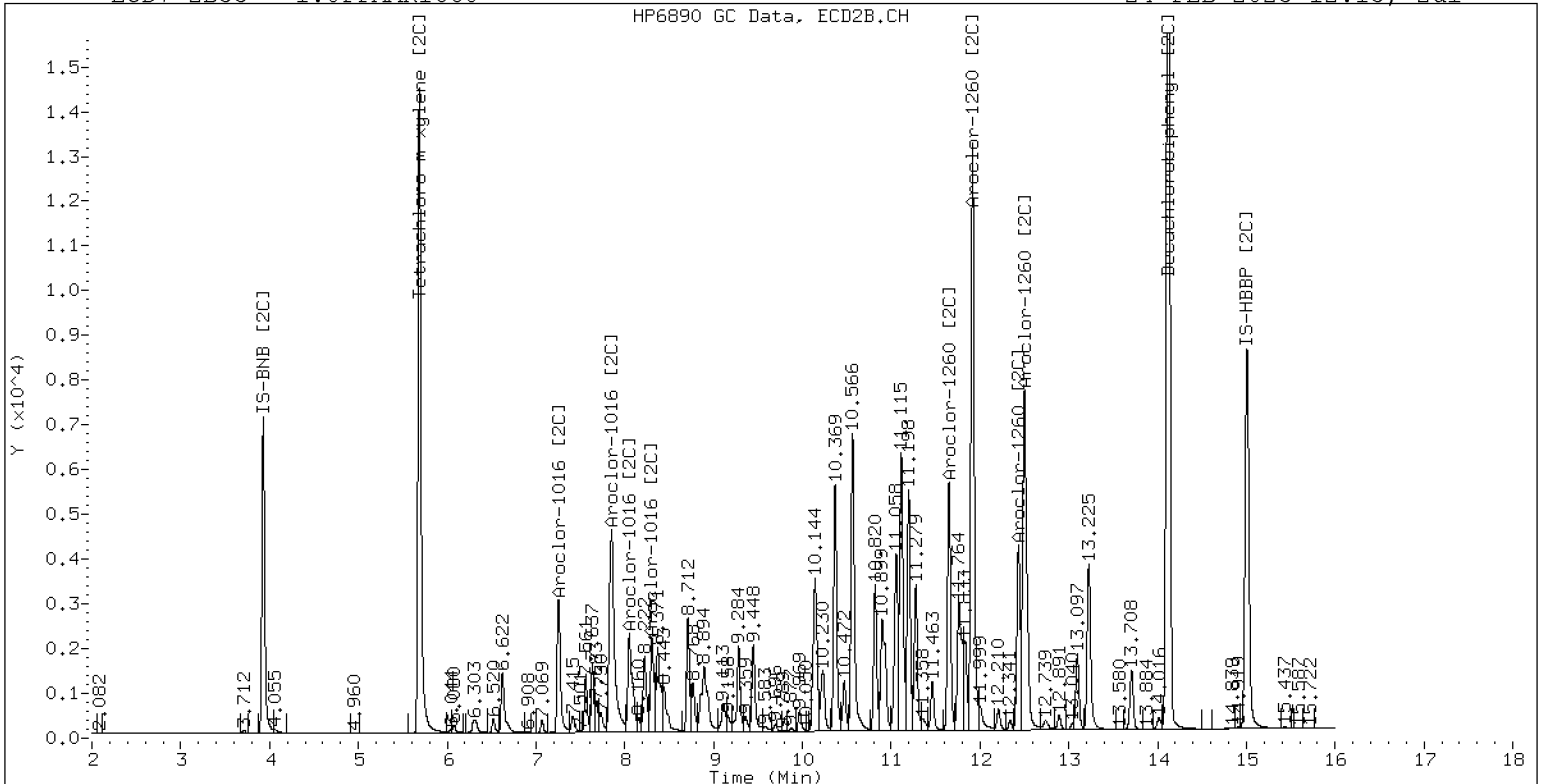
24-FEB-2023 12:15, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 1.0PPMAR1660

24-FEB-2023 12:15, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242306ECD7.D
Data file 2: /230224.b/230224.b/02242306ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.1PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 12:36
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col		ZB5	ZB35	RPD	Compound/Flag		
RT	Shift Response	RT	Shift Response	on col	on col				
5.809	0.002	155528	5.688	0.003	74628	15.9	16.0	0.9	Tetrachloro-m-xylene
13.892	-0.001	227253	14.119	-0.000	128496	15.8	15.8	0.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	655979	-2.6
Hexabromobiphenyl	1429847	1464509	2.4

Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	317418	0.7
Hexabromobiphenyl	513946	532962	3.7

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.271	0.000	25761	103.4	1	7.255	-0.000	19315	103.9
Aroclor-1016	2	7.657	0.003	75616	99.6	2	7.863	0.007	40308	107.0
Aroclor-1016	3	7.794	0.004	39547	106.7	3	8.059	0.005	18304	107.6
Aroclor-1016	4	8.406	0.001	24260	101.2	4	8.309	0.002	14708	110.2
Total CollAve (4 peaks):				102.7		Total Col2Ave (4 peaks):				107.2 RPD = 4
Corrected Ave (3 peaks):				101.4		Corrected Ave (3 peaks):				106.2 RPD = 5
CalAmt %D:				2.7		CalAmt %D:				7.2
Aroclor-1260	1	11.045	0.000	52009	98.7	1	11.655	0.002	31282	99.8
Aroclor-1260	2	11.362	0.001	55116	100.1	2	11.920	0.003	80574	100.7
Aroclor-1260	3	11.738	0.004	145604	99.7	3	12.437	0.002	19566	92.2
Aroclor-1260	4	12.141	0.002	72408	98.5	4	12.503	0.001	53588	99.4
Aroclor-1260	5	12.245	0.001	30745	97.2	NS	---			----
Total CollAve (5 peaks):				98.8		Total Col2Ave (4 peaks):				98.0 RPD = 1
Corrected Ave (4 peaks):				98.5		Corrected Ave (3 peaks):				97.1 RPD = 1
CalAmt %D:				-1.2		CalAmt %D:				-2.0

Total PCB Area Coll (5.906 - 13.793) = 1555762 Coll Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 764924 Col2 Total PCB = 0.2 ppm*

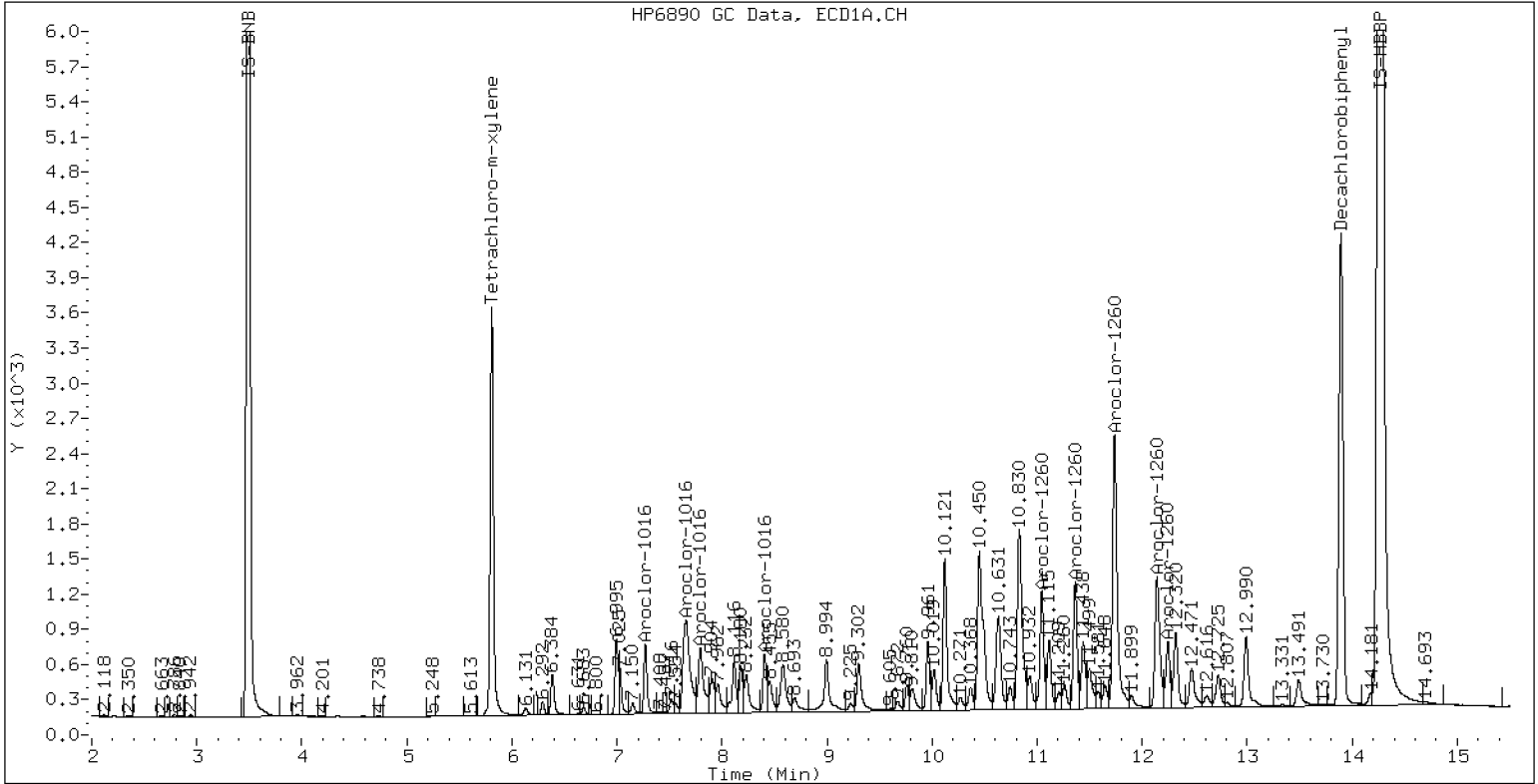
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.1PPMAR1660

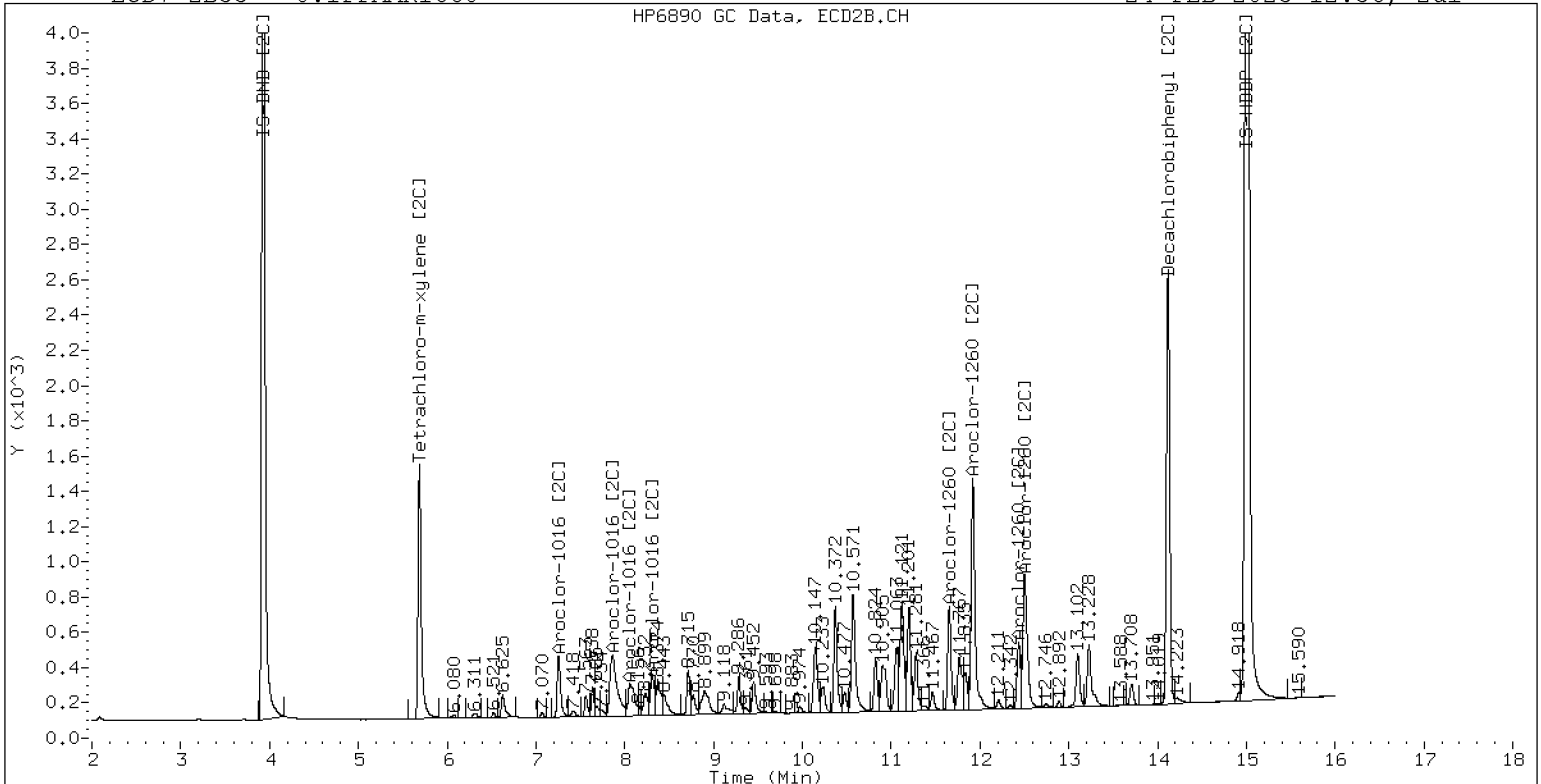
24-FEB-2023 12:36, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 0.1PPMAR1660

24-FEB-2023 12:36, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242307ECD7.D
Data file 2: /230224.b/230224.b/02242307ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.5PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 12:57
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.810	0.004	724614	5.688	0.003	359257	75.2	76.7	2.0	Tetrachloro-m-xylene
13.898	0.005	1056911	14.120	0.000	650153	74.3	79.5	6.9	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	645275	-4.2
Hexabromobiphenyl	1429847	1445345	1.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	319170	1.2
Hexabromobiphenyl	513946	536853	4.5

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	0.000	115193	470.0	1	7.256	0.000	86287	461.8	
Aroclor-1016	2	7.654	0.000	369991	495.2	2	7.856	0.000	192524	508.1	
Aroclor-1016	3	7.790	0.000	160952	441.3	3	8.055	0.000	81039	473.6	
Aroclor-1016	4	8.405	0.000	115032	487.9	4	8.307	0.000	62136	462.8	
Total CollAve (4 peaks):				473.6		Total Col2Ave (4 peaks):				476.6	RPD = 1
Corrected Ave (3 peaks):				466.4		Corrected Ave (3 peaks):				466.1	RPD = 0

CalAmt %D: -5.3

CalAmt %D: -4.7

Aroclor-1260	1	11.044	0.000	247212	475.5	1	11.653	0.000	145247	460.1	
Aroclor-1260	2	11.361	0.000	262877	483.9	2	11.918	0.000	379838	471.5	
Aroclor-1260	3	11.734	0.000	678830	471.1	3	12.436	0.000	104092	486.9	
Aroclor-1260	4	12.139	0.000	356067	490.7	4	12.502	0.000	258953	476.9	
Aroclor-1260	5	12.244	0.000	150280	481.2	NS	---			----	
Total CollAve (5 peaks):				480.5		Total Col2Ave (4 peaks):				473.8	RPD = 1
Corrected Ave (4 peaks):				477.9		Corrected Ave (3 peaks):				469.5	RPD = 2

CalAmt %D: -3.9

CalAmt %D: -5.2

Total PCB Area Coll (5.906 - 13.793) = 7134169 Coll Total PCB = 0.9 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 3589735 Col2 Total PCB = 0.9 ppm*

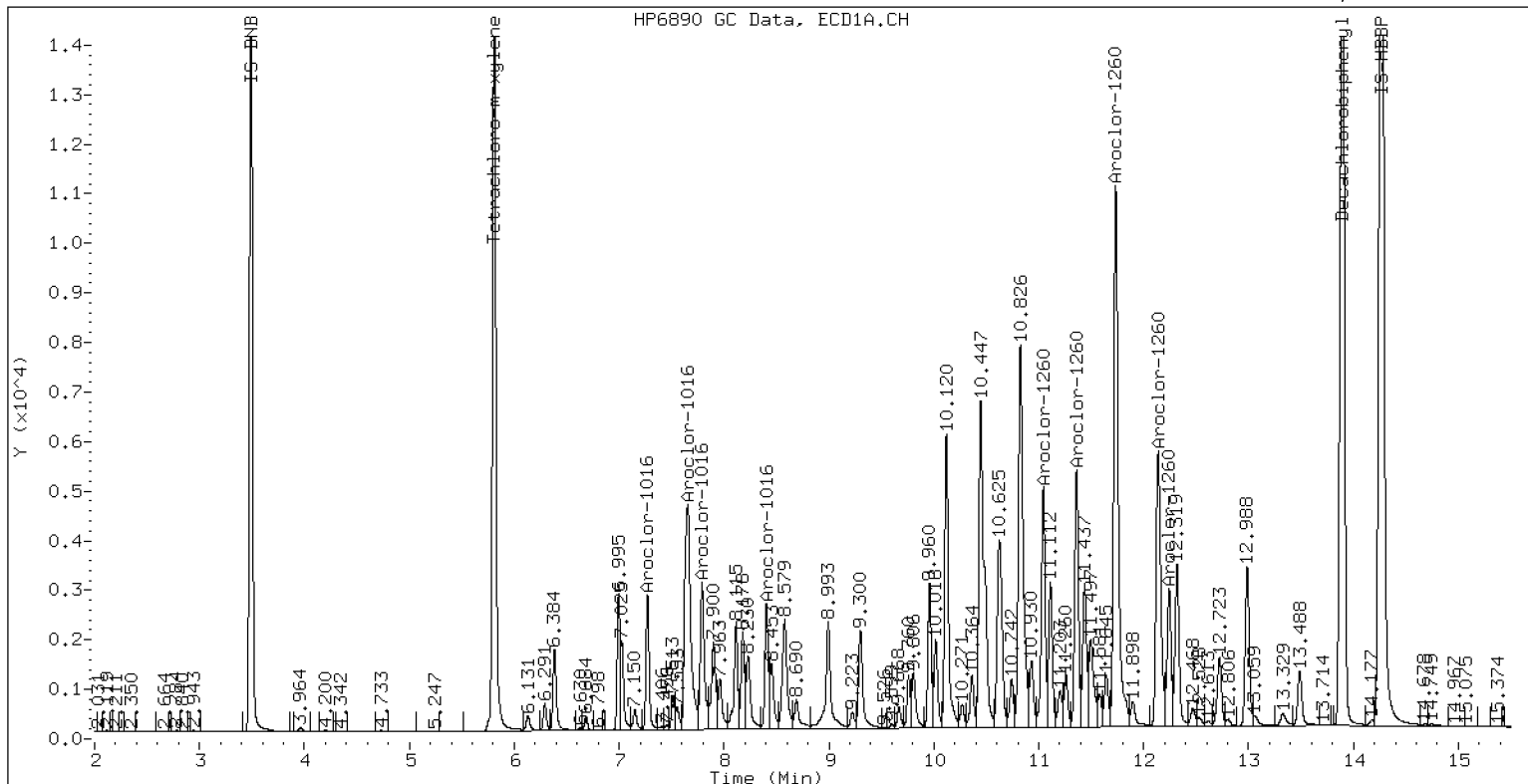
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.5PPMAR1660

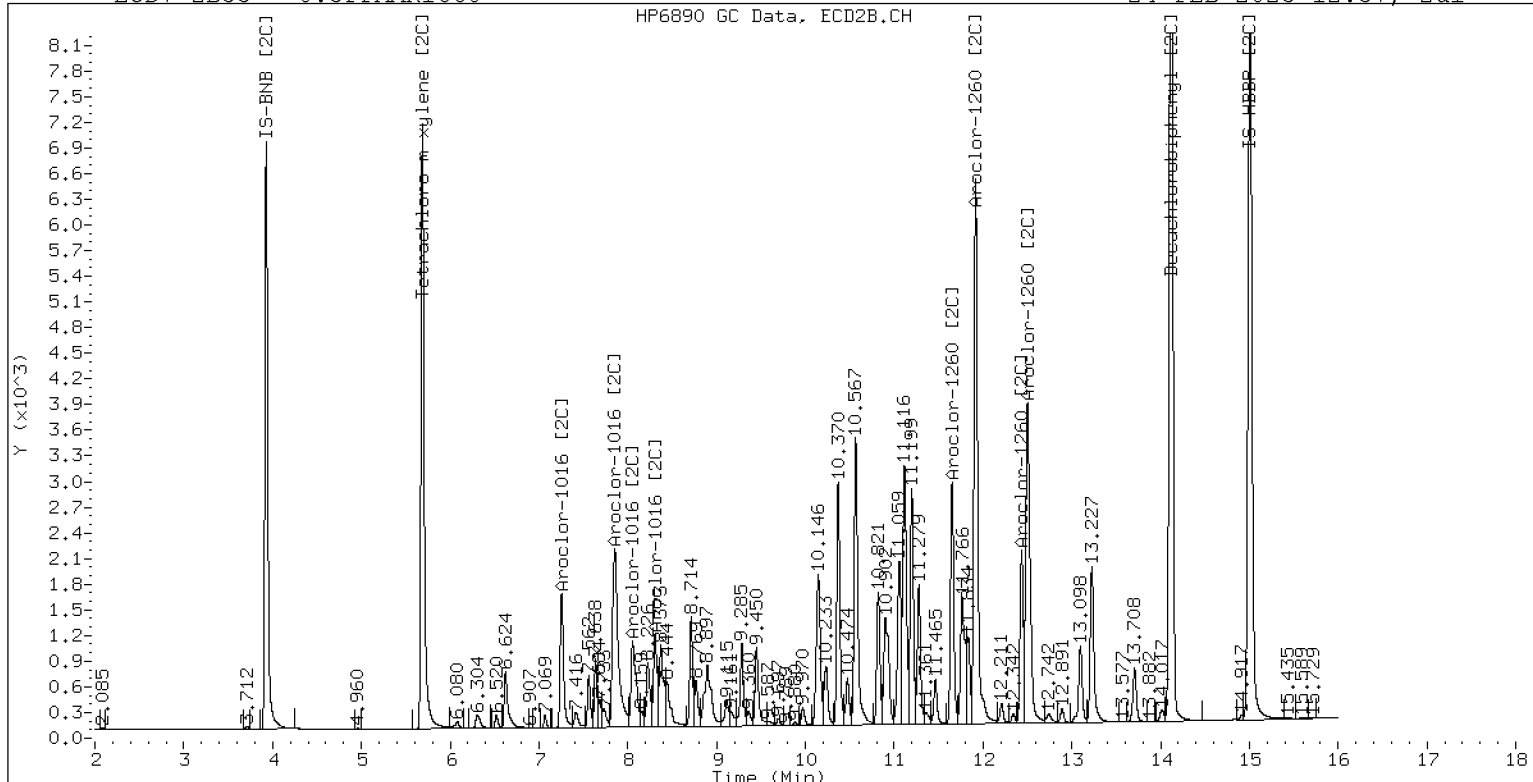
24-FEB-2023 12:57, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 0.5PPMAR1660

24-FEB-2023 12:57, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242308ECD7.D
Data file 2: /230224.b/230224.b/02242308ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1242.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1242
Client ID:
Injection Date: 24-FEB-2023 13:18
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.003	434187	5.688	0.003	214306	46.0	46.5	1.1	Tetrachloro-m-xylene
13.894	0.000	515867	14.119	-0.001	312943	35.6	38.5	7.6	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	632576	-6.1
Hexabromobiphenyl	1429847	1469715	2.8

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	314129	-0.4
Hexabromobiphenyl	513946	534294	4.0

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col ZB35 Col

Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1242	1	7.271	0.000	49009	250.0	1	7.255	0.000	36487	250.0
Aroclor-1242	2	7.656	0.000	148833	250.0	2	7.858	0.000	76699	250.0
Aroclor-1242	3	8.405	0.000	46308	250.0	3	9.167	0.000	23866	250.0
Aroclor-1242	4	8.579	0.000	68453	250.0	4	9.597	0.000	29080	250.0
Total Col1Ave (4 peaks):				250.0	Total Col2Ave (4 peaks):				250.0	RPD = 0
Corrected Ave (3 peaks):				250.0	Corrected Ave (3 peaks):				250.0	RPD = 0

Total PCB Area Col1 (5.906 - 13.793) = 1221467 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 572067 Col2 Total PCB = 0.2 ppm*

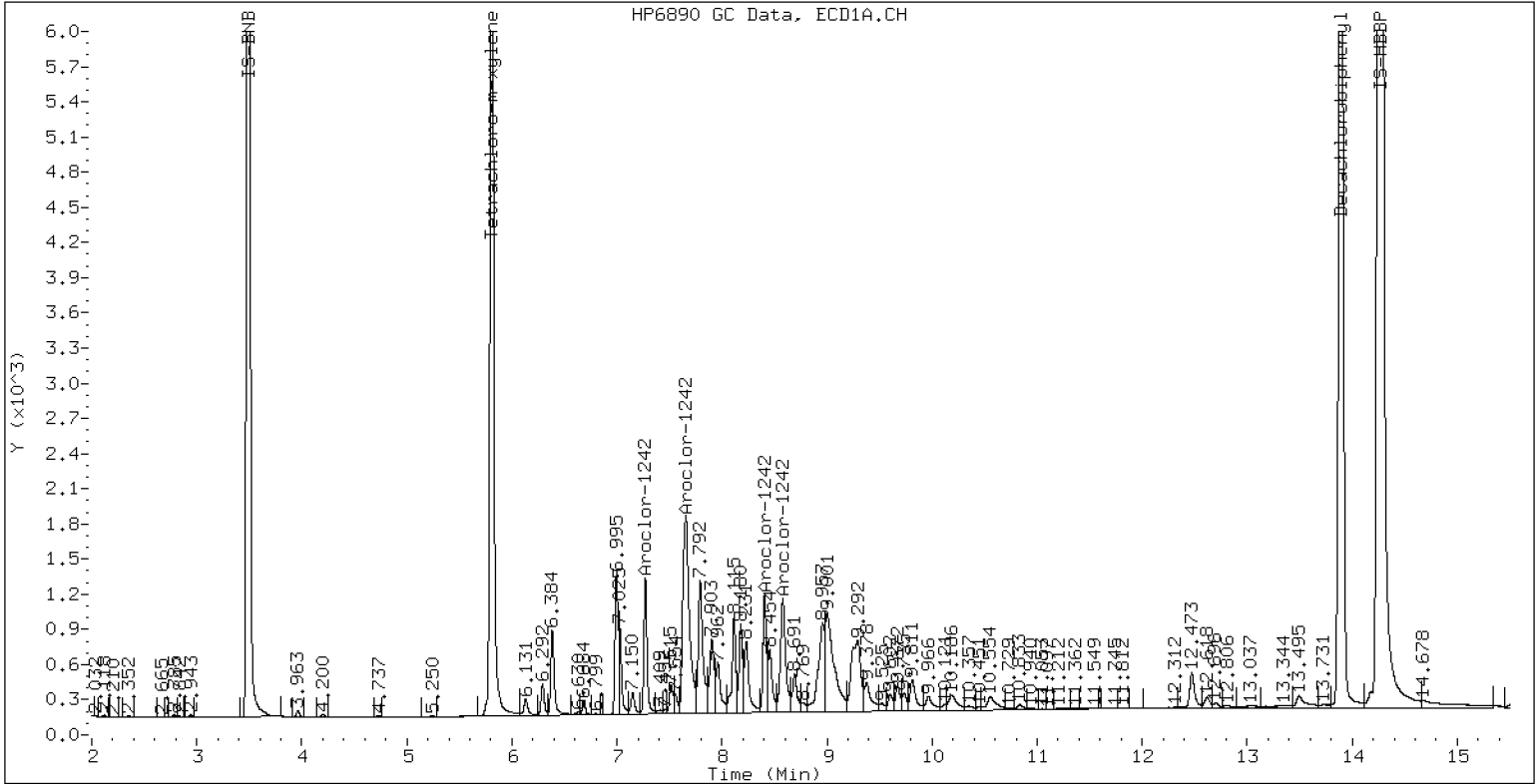
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1242

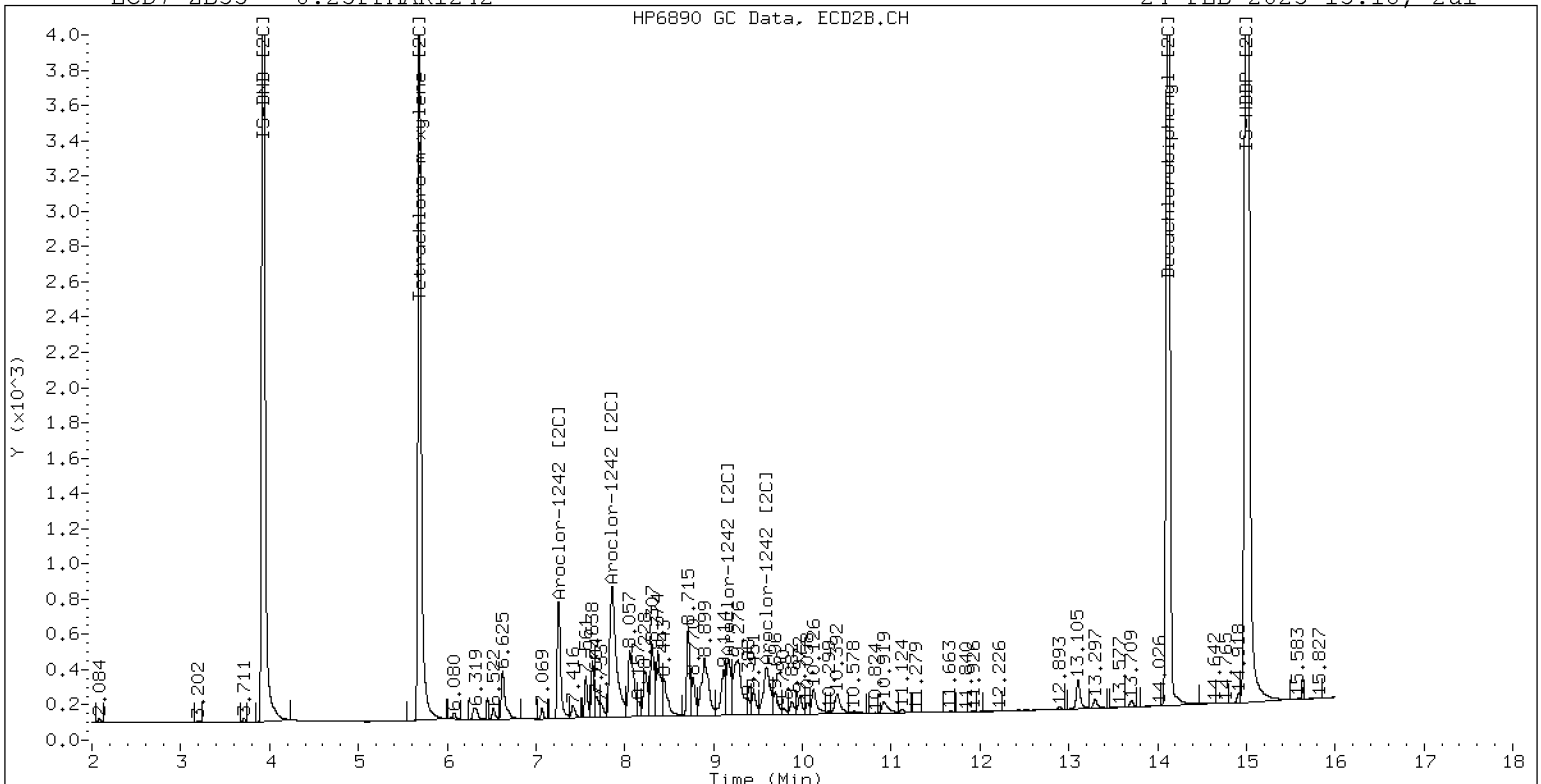
24-FEB-2023 13:18, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1242

24-FEB-2023 13:18, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242309ECD7.D
Data file 2: /230224.b/230224.b/02242309ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1248.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1248
Client ID:
Injection Date: 24-FEB-2023 13:39
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.003	349513	5.688	0.003	176615	36.6	37.9	3.4	Tetrachloro-m-xylene
13.894	0.001	523008	14.121	0.001	322054	36.4	39.3	7.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	639911	-5.0
Hexabromobiphenyl	1429847	1458696	2.0

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	317938	0.9
Hexabromobiphenyl	513946	538760	4.8

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col ZB35 Col

Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1248	1	8.405	0.000	78055	250.0	1	8.308	0.000	37951	250.0
Aroclor-1248	2	8.580	0.000	99216	250.0	2	8.714	0.000	39239	250.0
Aroclor-1248	3	8.999	0.000	187178	250.0	3	9.166	0.000	45157	250.0
Aroclor-1248	4	9.295	0.000	95291	250.0	4	9.590	0.000	54216	250.0
Total Col1Ave (4 peaks):				250.0	Total Col2Ave (4 peaks):				250.0	RPD = 0
Corrected Ave (3 peaks):				250.0	Corrected Ave (3 peaks):				250.0	RPD = 0

Total PCB Area Col1 (5.906 - 13.793) = 1565180 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 754991 Col2 Total PCB = 0.2 ppm*

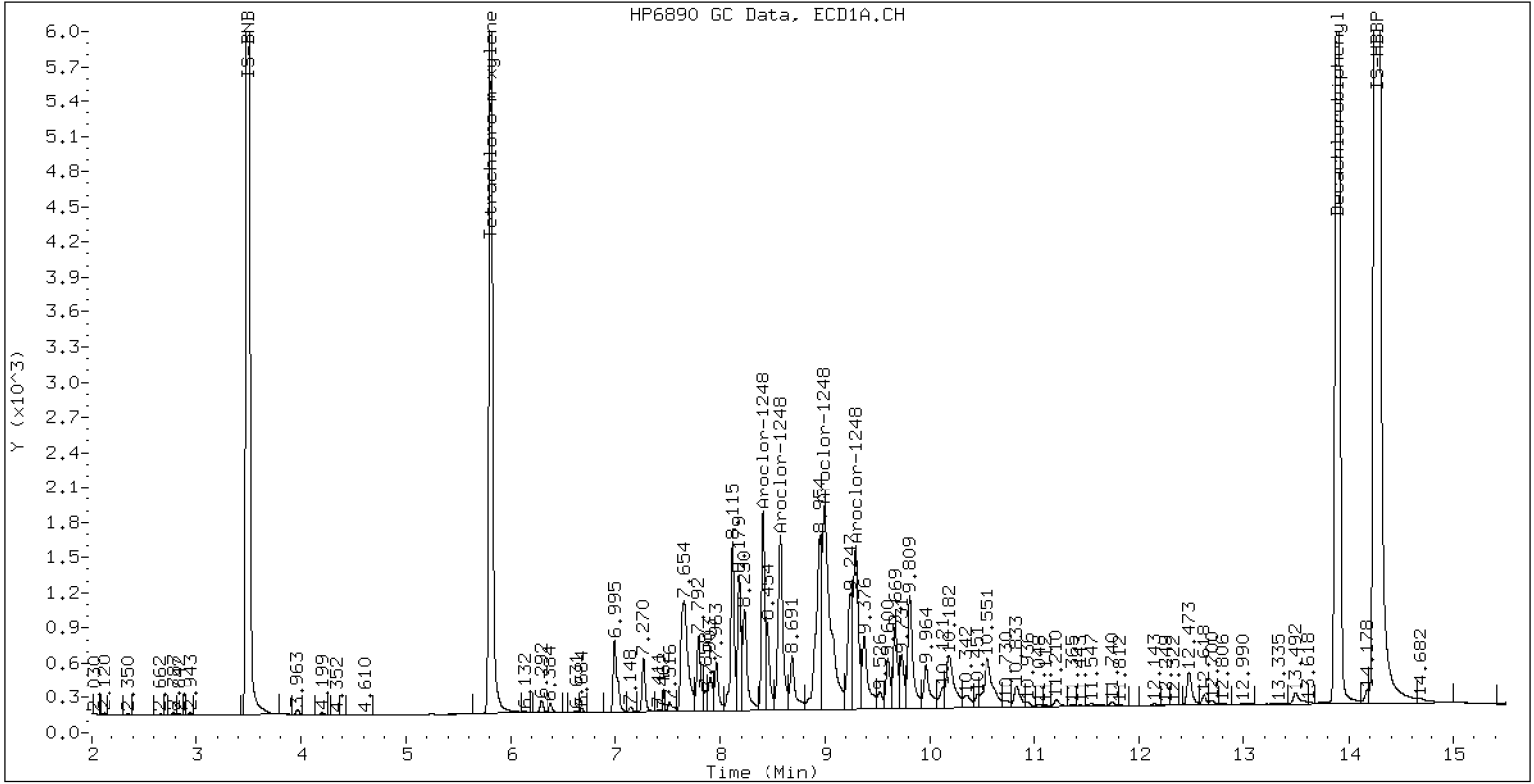
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1248

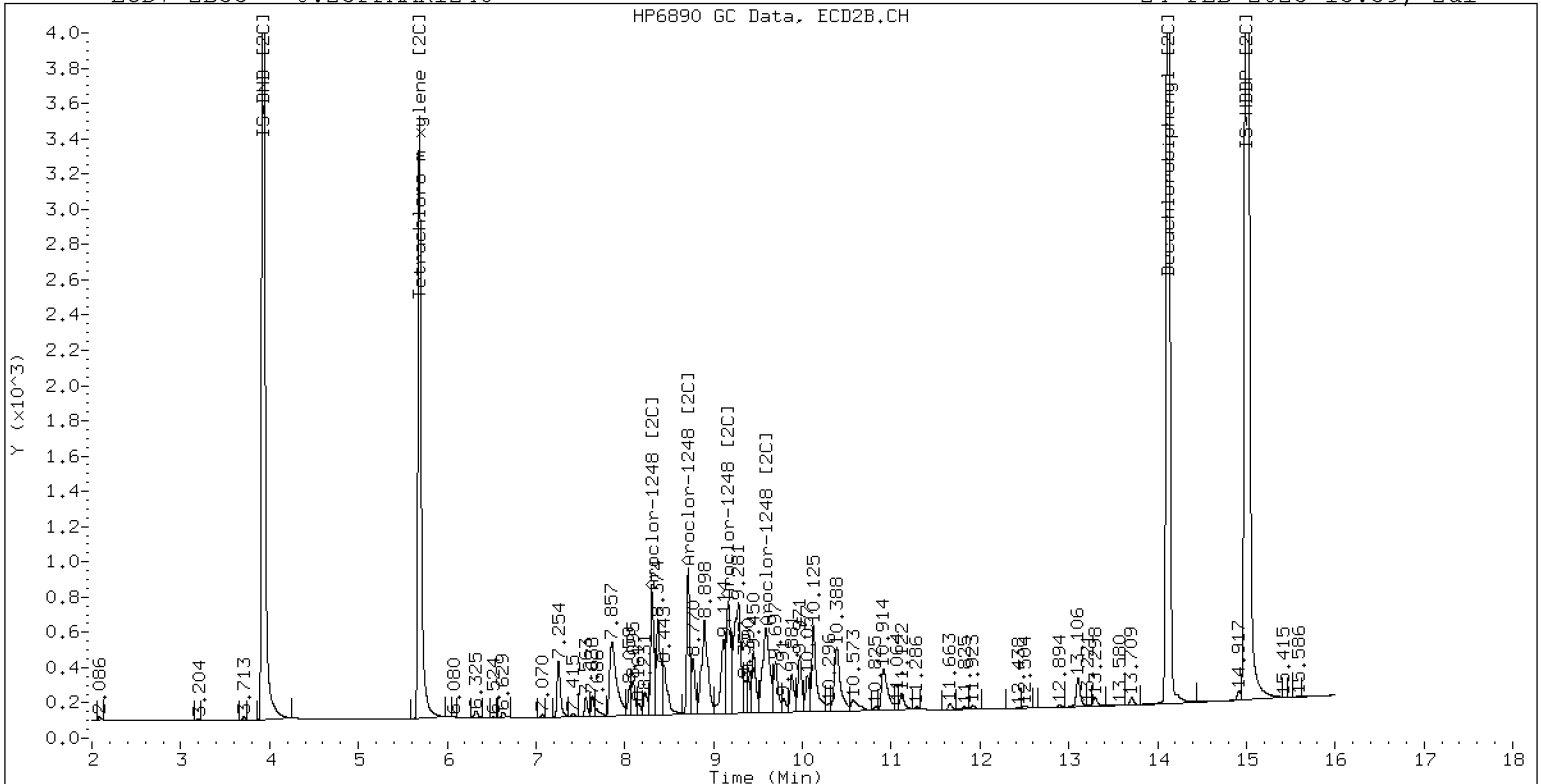
24-FEB-2023 13:39, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1248

24-FEB-2023 13:39, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242310ECD7.D
Data file 2: /230224.b/230224.b/02242310ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1254.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1254
Client ID:
Injection Date: 24-FEB-2023 14:00
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.808	0.002	352587	5.687	0.002	177502	37.3	38.6	3.4	Tetrachloro-m-xylene
13.895	0.002	532500	14.119	0.000	325903	37.0	40.2	8.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	633407	-6.0
Hexabromobiphenyl	1429847	1460265	2.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	313673	-0.5
Hexabromobiphenyl	513946	532442	3.6

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1254	1	9.299	0.000	159011	250.0	1	9.449	0.000	59603	250.0
Aroclor-1254	2	9.377	0.000	71516	250.0	2	9.970	0.000	47949	250.0
Aroclor-1254	3	9.668	0.000	102230	250.0	3	10.124	0.000	103745	250.0
Aroclor-1254	4	9.807	0.000	198777	250.0	4	10.373	0.000	101135	250.0
Aroclor-1254	5	10.176	0.000	124586	250.0	5	10.569	0.000	61577	250.0
Total CollAve (5 peaks):				250.0		Total Col2Ave (5 peaks):				250.0 RPD = 0
Corrected Ave (4 peaks):				250.0		Corrected Ave (4 peaks):				250.0 RPD = 0

Total PCB Area Coll (5.906 - 13.793) = 2179224 Coll Total PCB = 0.3 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1022156 Col2 Total PCB = 0.3 ppm*

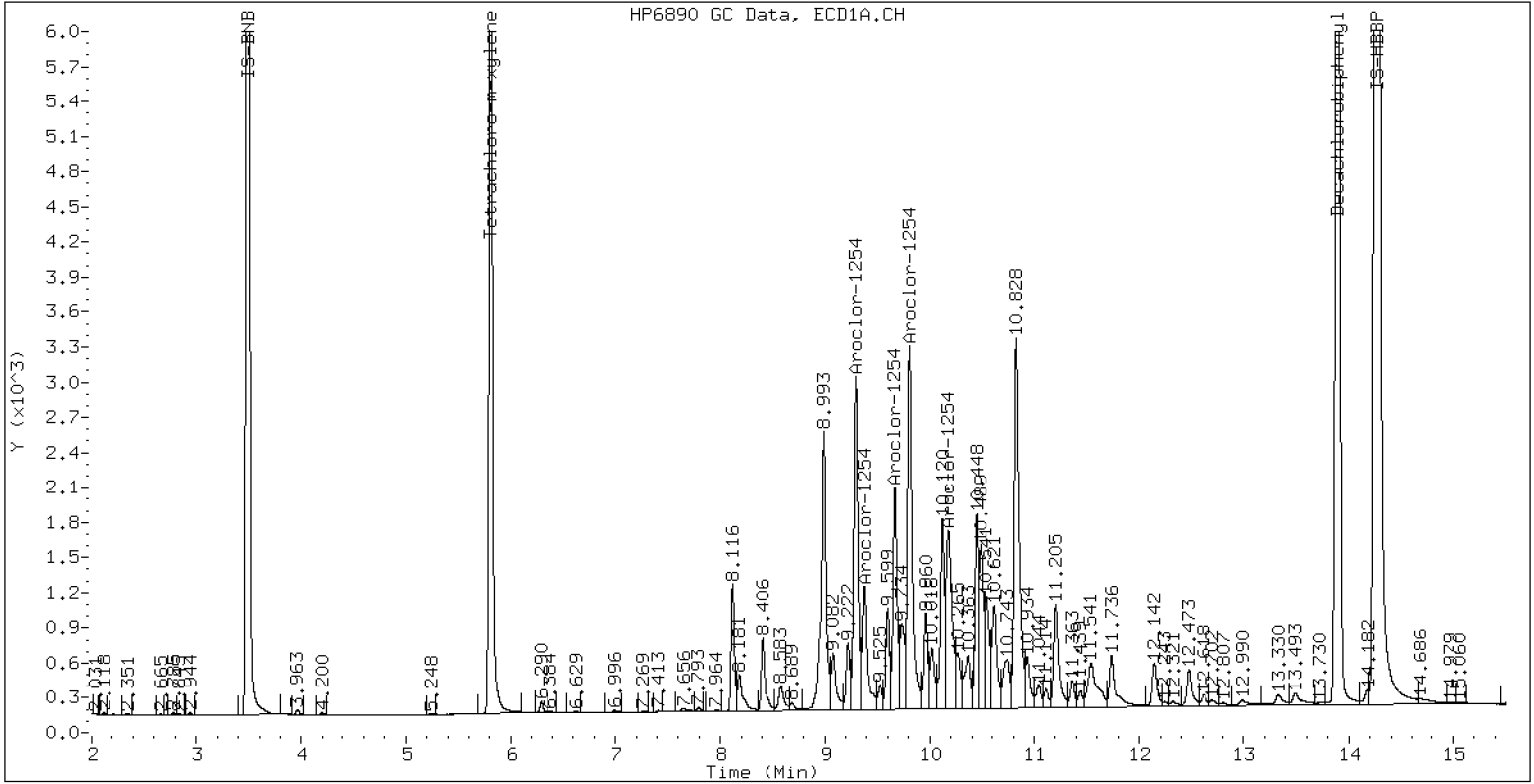
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1254

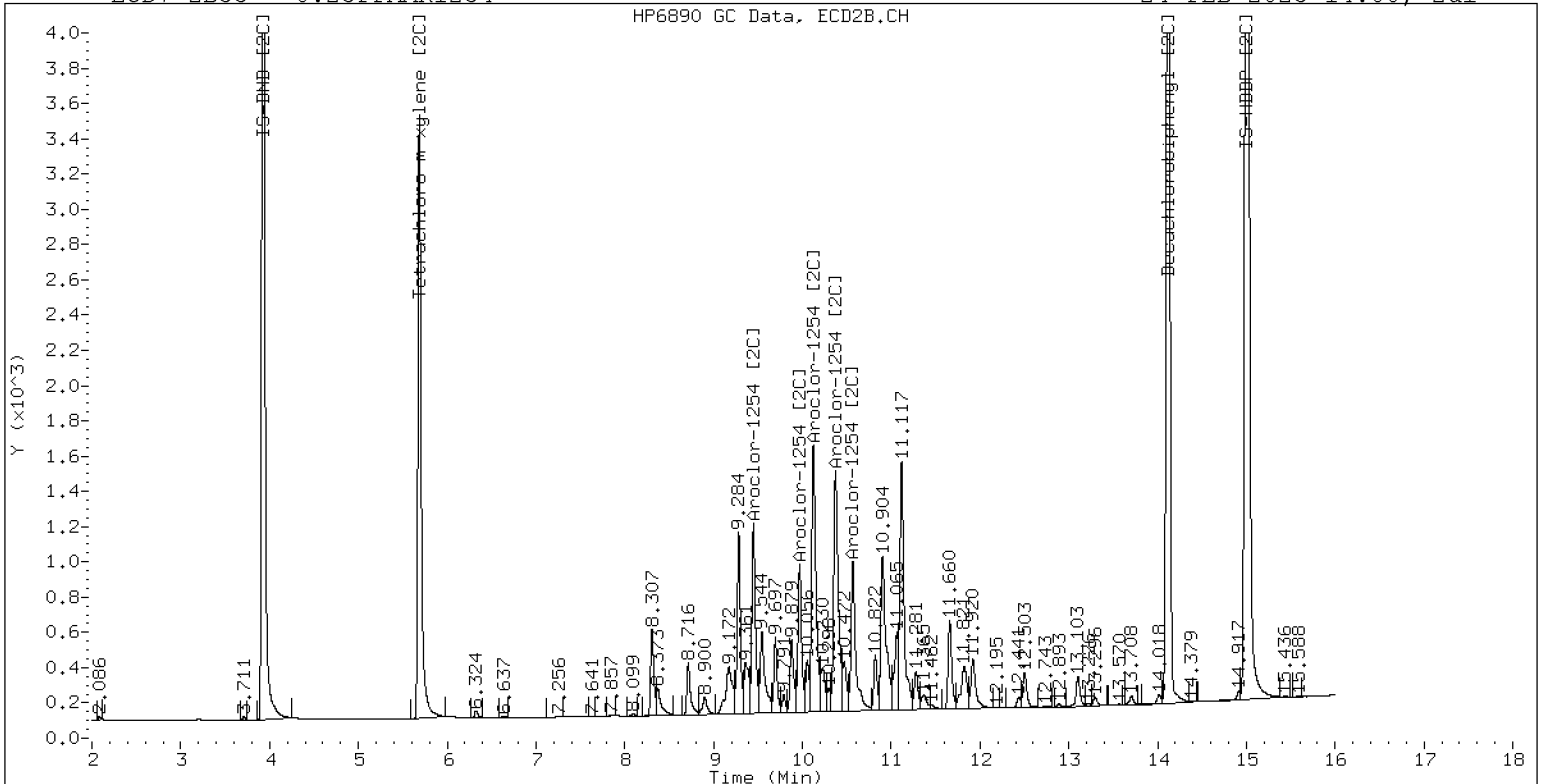
24-FEB-2023 14:00, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1254

24-FEB-2023 14:00, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242311ECD7.D
Data file 2: /230224.b/230224.b/02242311ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR2162.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR2162
Client ID:
Injection Date: 24-FEB-2023 14:21
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.807	0.001	362236	5.686	0.000	177349	38.4	39.2	2.1	Tetrachloro-m-xylene
13.894	0.001	523254	14.119	-0.000	321034	36.0	39.2	8.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	632433	-6.1
Hexabromobiphenyl	1429847	1474039	3.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	308453	-2.2
Hexabromobiphenyl	513946	538177	4.7

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col ZB35 Col

Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1221	1	4.731	0.000	14160	250.0	1	4.956	0.000	7300	250.0	
Aroclor-1221	2	6.132	0.000	25324	250.0	2	6.296	0.000	13816	250.0	
Aroclor-1221	3	6.382	0.000	58795	250.0	3	6.622	0.000	22491	250.0	
Total CollAve (3 peaks):				250.0	Total Col2Ave (3 peaks):				250.0	RPD = 0	
Corrected Ave: < 3 Peaks					Corrected Ave: < 3 Peaks						

Aroclor-1262	1	10.829	0.000	113046	250.0	1	11.200	0.000	114880	250.0	
Aroclor-1262	2	12.244	0.000	183948	250.0	2	11.652	0.000	97844	250.0	
Aroclor-1262	3	12.319	0.000	197749	250.0	3	12.434	0.000	111015	250.0	
Aroclor-1262	4	12.987	0.000	180727	250.0	4	12.502	0.000	173913	250.0	
Total CollAve (4 peaks):				250.0	Total Col2Ave (4 peaks):				250.0	RPD = 0	
Corrected Ave (3 peaks):				250.0	Corrected Ave (3 peaks):				250.0	RPD = 0	

Total PCB Area Coll (5.906 - 13.793) = 3105316 Coll Total PCB = 0.4 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1573107 Col2 Total PCB = 0.4 ppm*

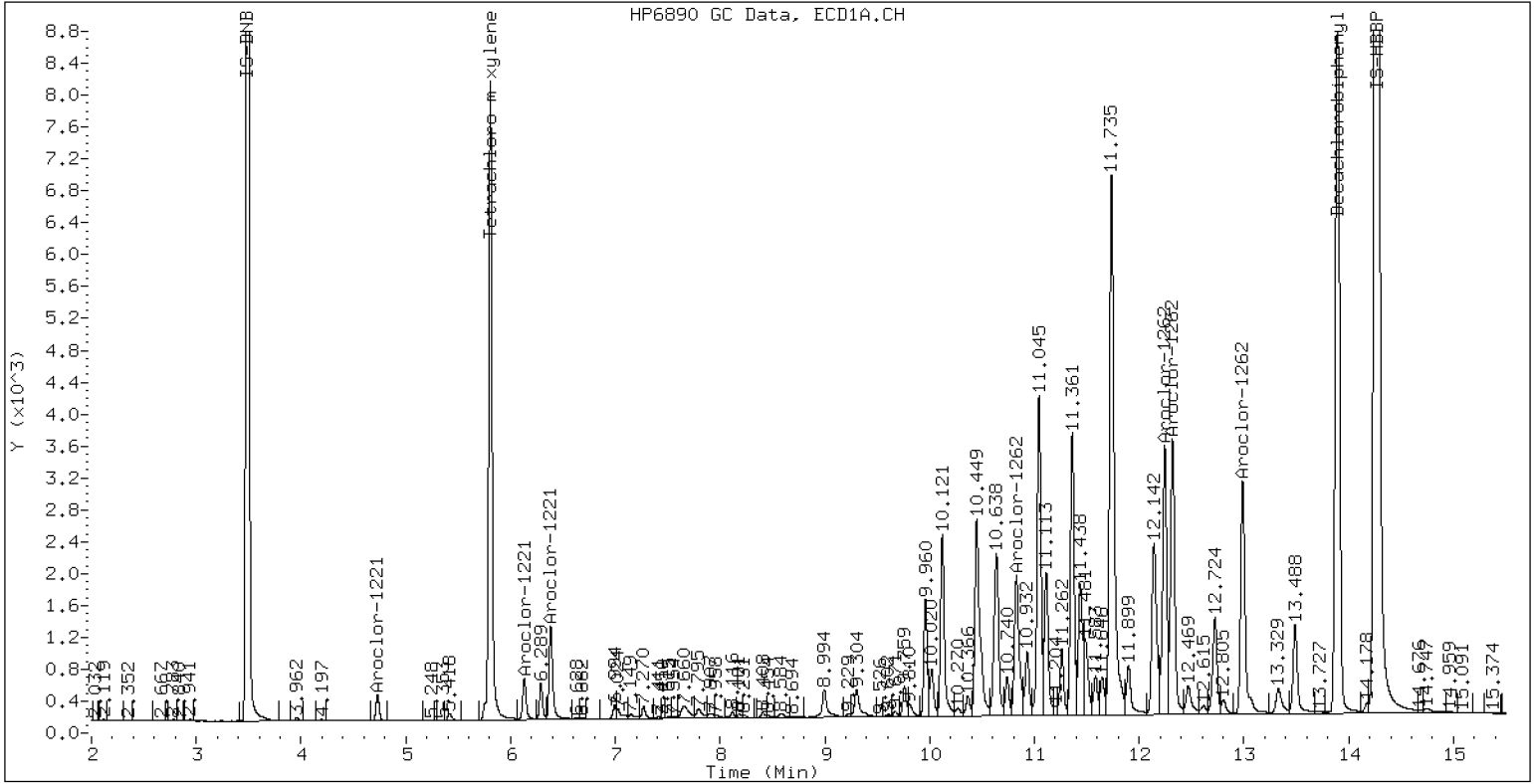
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR2162

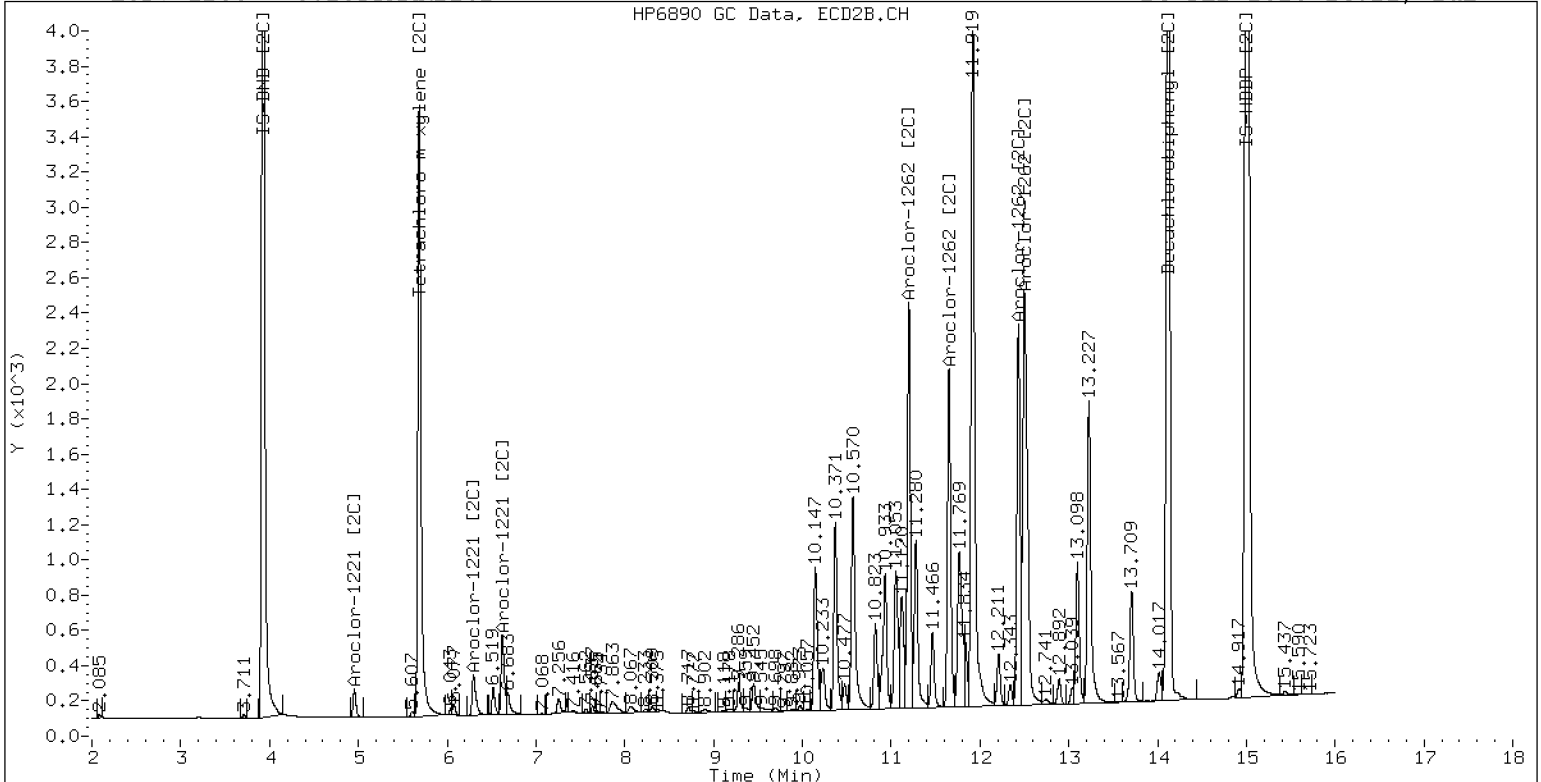
24-FEB-2023 14:21, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR2162

24-FEB-2023 14:21, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242312ECD7.D
Data file 2: /230224.b/230224.b/02242312ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR3268.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR3268
Client ID:
Injection Date: 24-FEB-2023 14:42
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.806	0.000	366416	5.685	0.000	179450	38.0	38.9	2.4	Tetrachloro-m-xylene
13.893	0.000	778191	14.119	0.000	477889	53.0	57.5	8.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	645602	-4.2
Hexabromobiphenyl	1429847	1492154	4.4

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	314042	-0.4
Hexabromobiphenyl	513946	545458	6.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col ZB35 Col

Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1232	1	4.730	0.000	8647	250.0	1	4.956	0.000	4017	250.0
Aroclor-1232	2	6.131	0.000	17148	250.0	2	7.254	0.000	19962	250.0
Aroclor-1232	3	7.656	0.000	77627	250.0	3	7.861	0.000	39913	250.0
Aroclor-1232	4	8.581	0.000	32993	250.0	4	8.715	0.000	11487	250.0
Total CollAve (4 peaks):				250.0		Total Col2Ave (4 peaks):				250.0 RPD = 0
Corrected Ave (3 peaks):				250.0		Corrected Ave (3 peaks):				250.0 RPD = 0

Aroclor-1268	1	12.247	0.000	477974	250.0	1	12.432	0.000	274595	250.0
Aroclor-1268	2	12.317	0.000	473326	250.0	2	12.500	0.000	295194	250.0
Aroclor-1268	3	12.699	0.000	405011	250.0	3	12.892	0.000	252048	250.0
Aroclor-1268	4	13.490	0.000	1333528	250.0	4	13.709	0.000	805579	250.0
Total CollAve (4 peaks):				250.0		Total Col2Ave (4 peaks):				250.0 RPD = 0
Corrected Ave (3 peaks):				250.0		Corrected Ave (3 peaks):				250.0 RPD = 0

Total PCB Area Coll (5.906 - 13.793) = 3998414 Coll Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 2300029 Col2 Total PCB = 0.6 ppm*

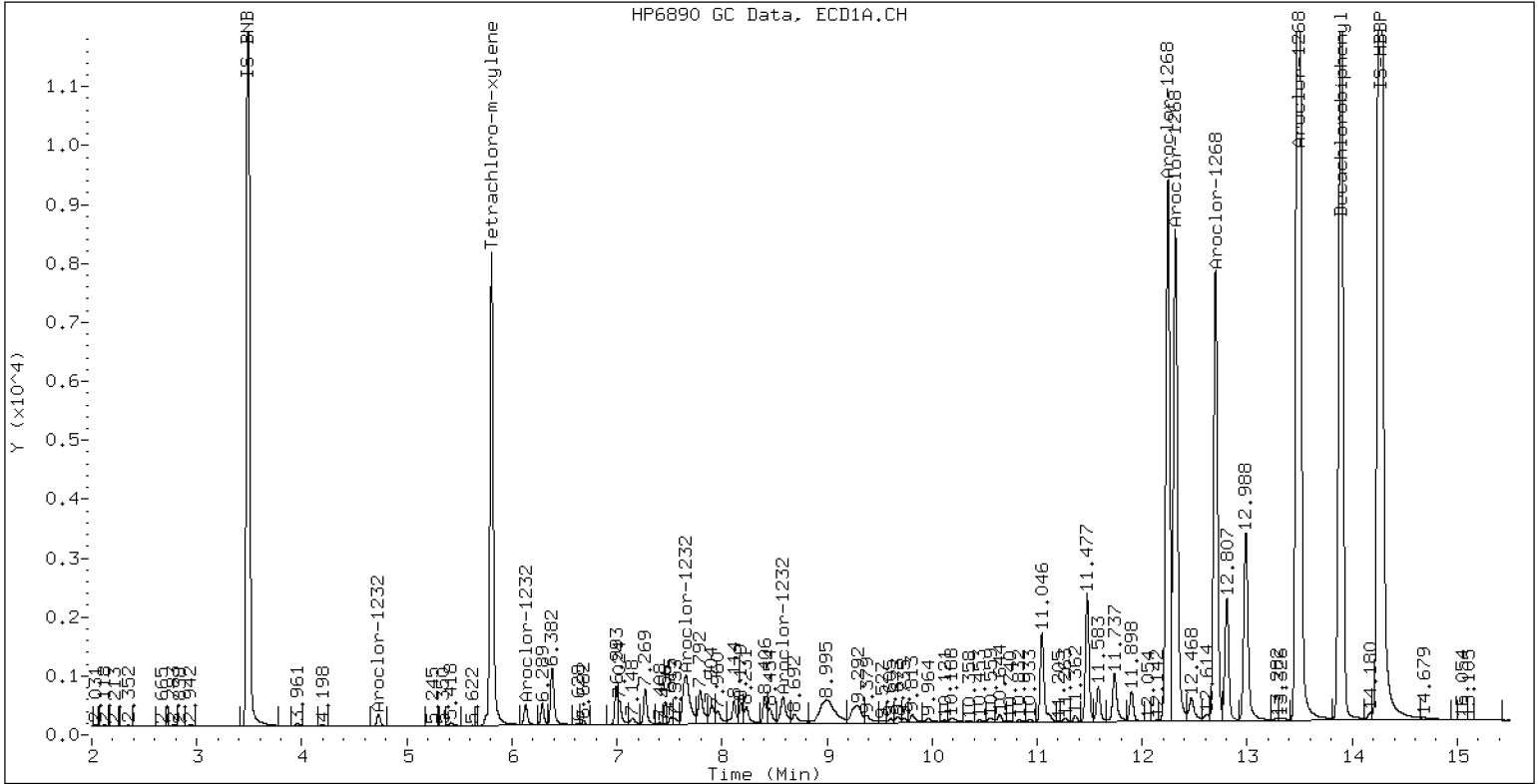
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR3268

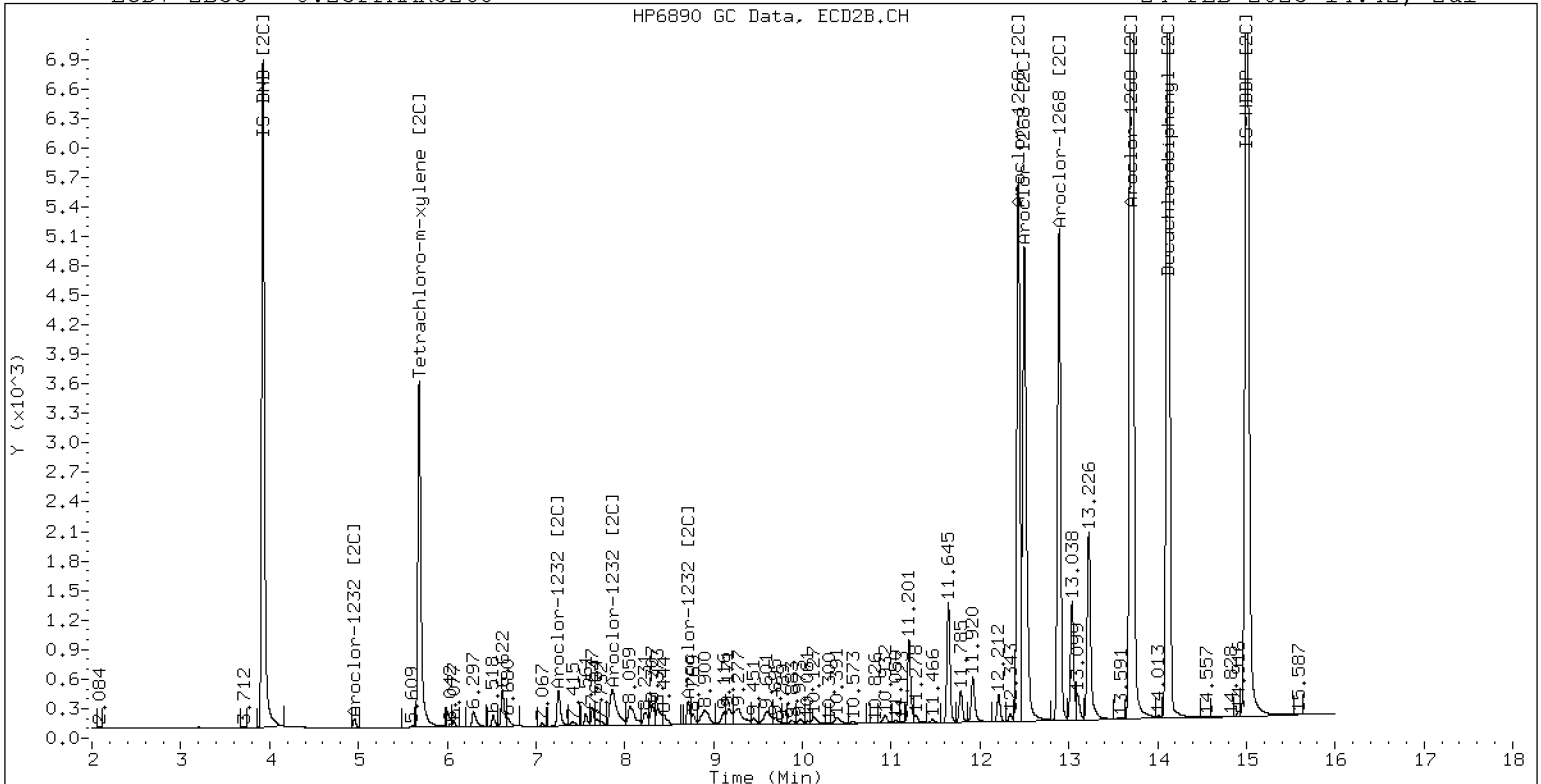
24-FEB-2023 14:42, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR3268

24-FEB-2023 14:42, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242313ECD7.D
Data file 2: /230224.b/230224.b/02242313ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1660SCV
Client ID:
Injection Date: 24-FEB-2023 15:03
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.807	0.001	337070	5.686	0.001	165848	34.9	35.8	2.3	Tetrachloro-m-xylene
13.895	0.002	515407	14.119	-0.000	316730	34.3	37.3	8.4	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	645975	-4.1
Hexabromobiphenyl	1429847	1524245	6.6

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	316115	0.3
Hexabromobiphenyl	513946	556950	8.4

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.269	-0.002	59491	242.5	1	7.254	-0.002	44576	240.9	
Aroclor-1016	2	7.655	0.001	181090	242.1	2	7.857	0.002	95386	254.2	
Aroclor-1016	3	7.790	0.000	88470	242.3	3	8.056	0.002	42160	248.8	
Aroclor-1016	4	8.404	-0.001	57980	245.6	4	8.307	0.000	32197	242.1	
Total CollAve (4 peaks):				243.1	Total Col2Ave (4 peaks):				246.5	RPD = 1	
Corrected Ave (3 peaks):				242.3	Corrected Ave (3 peaks):				243.9	RPD = 1	
Aroclor-1221	1	4.731	0.000	464	8.0	1	---			0.0	
Aroclor-1221	2	6.130	-0.002	9233	89.2	2	6.300	0.004	5379	95.0	
Aroclor-1221	3	6.382	-0.001	42570	177.2	3	6.623	0.001	20952	227.2	
Total CollAve (3 peaks):				91.5	Col2Ave: <3 Quant Peaks						
Aroclor-1232	1	4.731	0.001	464	13.4	1	---			0.0	
Aroclor-1232	2	6.130	-0.001	9233	134.5	2	7.254	-0.000	44576	554.6	
Aroclor-1232	3	7.655	-0.001	181090	582.9	3	7.857	-0.003	95386	593.5	
Aroclor-1232	4	8.580	-0.001	79916	605.2	4	8.713	-0.002	29795	644.2	
Total CollAve (4 peaks):				334.0	Total Col2Ave (3 peaks):				597.4	RPD = 57*	
Corrected Ave (3 peaks):				243.6	Corrected Ave: < 3 Peaks						
Aroclor-1242	1	7.269	-0.002	59491	297.2	1	7.254	-0.002	44576	303.5	
Aroclor-1242	2	7.655	-0.001	181090	297.9	2	7.857	-0.000	95386	309.0	
Aroclor-1242	3	8.404	-0.001	57980	306.5	3	9.115	-0.052	18754	195.2	
Aroclor-1242	4	8.580	0.000	79916	285.8	4	9.697	0.100	1355	11.6	
Total CollAve (4 peaks):				296.8	Total Col2Ave (4 peaks):				204.8	RPD = 37	
Corrected Ave (3 peaks):				293.6	Corrected Ave (3 peaks):				170.1	RPD = 53*	
Aroclor-1248	1	8.404	-0.001	57980	184.0	1	8.307	-0.001	32197	213.3	
Aroclor-1248	2	8.580	-0.001	79916	199.5	2	8.713	-0.001	29795	190.9	
Aroclor-1248	3	8.993	-0.006	71805	95.0	3	9.115	-0.050	18754	104.4	
Aroclor-1248	4	9.300	0.006	47348	123.1	4	---			0.0	
Total CollAve (4 peaks):				150.4	Total Col2Ave (3 peaks):				169.6	RPD = 12	
Corrected Ave (3 peaks):				134.0	Corrected Ave: < 3 Peaks						
Aroclor-1254	1	9.300	0.002	47348	73.0	1	9.451	0.001	22438	93.4	
Aroclor-1254	2	---			0.0	2	9.972	0.001	2694	13.9	
Aroclor-1254	3	9.670	0.002	5461	13.1	3	10.147	0.024	52914	126.5	
Aroclor-1254	4	9.807	-0.000	18944	23.4	4	10.370	-0.003	70430	172.8	
Aroclor-1254	5	10.121	-0.056	154170	303.3	5	10.568	-0.000	98525	396.9	
Total CollAve (4 peaks):				103.2	Total Col2Ave (5 peaks):				160.7	RPD = 44*	
Corrected Ave (3 peaks):				36.5	Corrected Ave (4 peaks):				101.7	RPD = 94*	
Aroclor-1260	1	11.044	0.000	149195	272.1	1	11.653	0.000	82210	251.0	
Aroclor-1260	2	11.361	-0.000	153832	268.5	2	11.919	0.001	222226	265.9	
Aroclor-1260	3	11.736	0.002	396660	261.0	3	12.435	-0.000	59148	266.7	
Aroclor-1260	4	12.140	0.001	190448	248.9	4	12.504	0.002	147180	261.2	
Aroclor-1260	5	12.244	-0.000	91385	277.5	NS	---			----	
Total CollAve (5 peaks):				265.6	Total Col2Ave (4 peaks):				261.2	RPD = 2	
Corrected Ave (4 peaks):				262.6	Corrected Ave (3 peaks):				259.4	RPD = 1	
Aroclor-1262	1	10.827	-0.002	220238	471.0	1	11.199	-0.001	84479	177.6	
Aroclor-1262	2	12.244	0.000	91385	120.1	2	11.653	0.002	82210	203.0	
Aroclor-1262	3	12.320	0.001	113066	138.2	3	12.435	0.002	59148	128.7	
Aroclor-1262	4	12.988	0.001	102156	136.7	4	12.504	0.002	147180	204.4	
Total CollAve (4 peaks):				216.5	Total Col2Ave (4 peaks):				178.4	RPD = 19	
Corrected Ave (3 peaks):				131.7	Corrected Ave (3 peaks):				169.8	RPD = 25	
Aroclor-1268	1	12.244	-0.003	91385	46.8	1	12.435	0.003	59148	52.7	
Aroclor-1268	2	12.320	0.003	113066	58.5	2	12.504	0.004	147180	122.1	
Aroclor-1268	3	12.726	0.027	46633	28.2	3	12.893	0.001	2874	2.8	
Aroclor-1268	4	13.489	-0.000	25567	4.7	4	13.709	-0.000	13041	4.0	
Total CollAve (4 peaks):				34.5	Total Col2Ave (4 peaks):				45.4	RPD = 27	
Corrected Ave (3 peaks):				26.6	Corrected Ave (3 peaks):				19.8	RPD = 29	

Total PCB Area Col1 (5.906 - 13.793) = 3743076 Col1 Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1897008 Col2 Total PCB = 0.5 ppm*

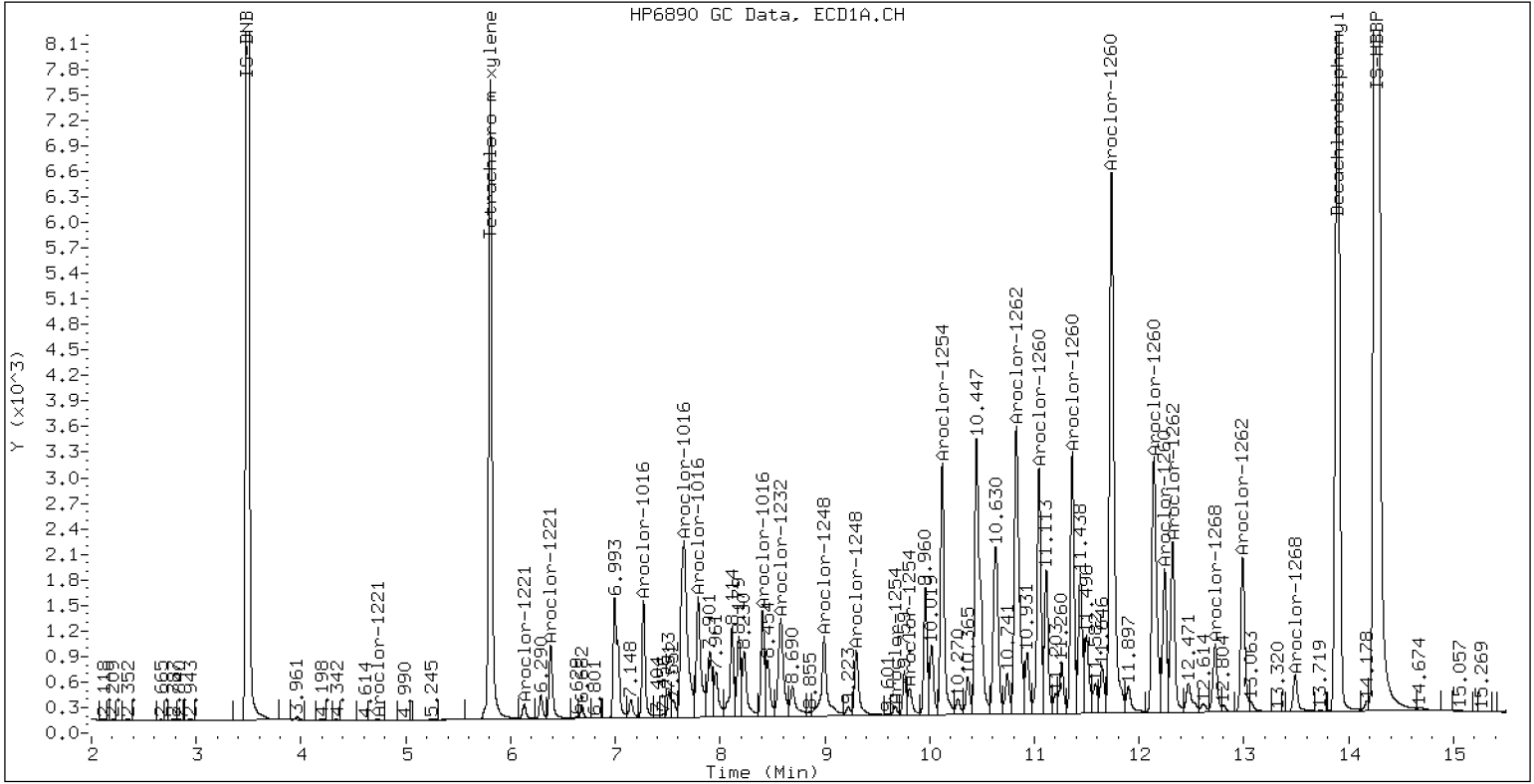
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1660SCV

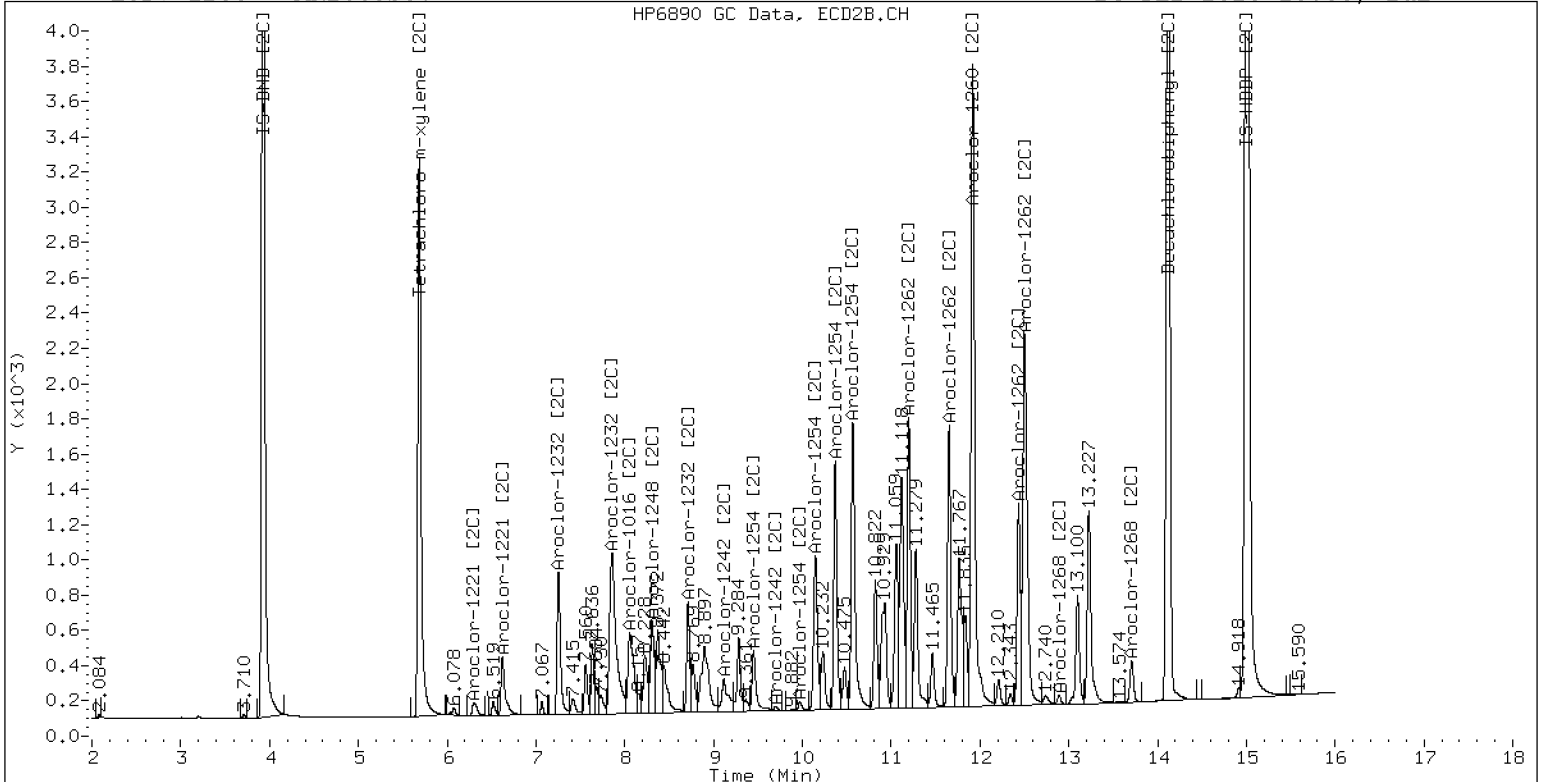
24-FEB-2023 15:03, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1660SCV

24-FEB-2023 15:03, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242314ECD7.D
Data file 2: /230224.b/230224.b/02242314ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1242SCV
Client ID:
Injection Date: 24-FEB-2023 15:24
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.808	0.002	354283	5.686	0.001	172455	33.6	34.5	2.6	Tetrachloro-m-xylene
13.895	0.002	567088	14.120	0.001	347430	37.0	40.3	8.6	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	705650	4.7
Hexabromobiphenyl	1429847	1555683	8.8

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	340433	8.0
Hexabromobiphenyl	513946	565609	10.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.270	-0.000	39927	149.0	1	7.256	0.000	32417	162.7
Aroclor-1016	2	7.653	-0.001	132339	162.0	2	7.856	0.001	69235	171.3
Aroclor-1016	3	7.791	0.001	59310	148.7	3	8.055	0.000	29473	161.5
Aroclor-1016	4	8.405	0.000	42537	165.0	4	8.307	-0.000	22792	159.2
Total CollAve (4 peaks):				156.2		Total Col2Ave (4 peaks):				163.7 RPD = 5
Corrected Ave (3 peaks):				153.2		Corrected Ave (3 peaks):				161.1 RPD = 5
Aroclor-1221	1	4.733	0.002	319	5.0	1	---			0.0
Aroclor-1221	2	6.131	-0.001	6534	57.8	2	6.319	0.022	4365	71.6
Aroclor-1221	3	6.384	0.001	29664	113.0	3	6.624	0.002	14916	150.2
Total CollAve (3 peaks):				58.6		Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	4.733	0.003	319	8.4	1	---			0.0
Aroclor-1232	2	6.131	0.000	6534	87.2	2	7.256	0.002	32417	374.5
Aroclor-1232	3	7.653	-0.003	132339	389.9	3	7.856	-0.004	69235	400.0
Aroclor-1232	4	8.579	-0.002	69445	481.4	4	8.714	-0.001	22167	445.0
Total CollAve (4 peaks):				241.7		Total Col2Ave (3 peaks):				406.5 RPD = 51*
Corrected Ave (3 peaks):				161.8		Corrected Ave: < 3 Peaks				
Aroclor-1242	1	7.270	-0.001	39927	182.6	1	7.256	0.000	32417	205.0
Aroclor-1242	2	7.653	-0.003	132339	199.3	2	7.856	-0.002	69235	208.2
Aroclor-1242	3	8.405	-0.000	42537	205.9	3	9.164	-0.004	23068	223.0
Aroclor-1242	4	8.579	-0.000	69445	227.4	4	9.587	-0.010	31021	246.1
Total CollAve (4 peaks):				203.8		Total Col2Ave (4 peaks):				220.6 RPD = 8
Corrected Ave (3 peaks):				195.9		Corrected Ave (3 peaks):				212.1 RPD = 8
Aroclor-1248	1	8.405	0.000	42537	123.5	1	8.307	-0.001	22792	140.2
Aroclor-1248	2	8.579	-0.001	69445	158.7	2	8.714	-0.000	22167	131.9
Aroclor-1248	3	9.001	0.003	91942	111.4	3	9.164	-0.002	23068	119.3
Aroclor-1248	4	9.294	-0.000	38711	92.1	4	9.587	-0.003	31021	133.6
Total CollAve (4 peaks):				121.4		Total Col2Ave (4 peaks):				131.2 RPD = 8
Corrected Ave (3 peaks):				109.0		Corrected Ave (3 peaks):				128.3 RPD = 16
Aroclor-1254	1	9.294	-0.005	38711	54.6	1	9.450	0.001	13131	50.7
Aroclor-1254	2	9.377	-0.000	17371	54.5	2	9.970	0.000	8340	40.1
Aroclor-1254	3	9.668	-0.000	16373	35.9	3	10.123	-0.000	16364	36.3
Aroclor-1254	4	9.807	-0.001	27490	31.0	4	10.382	0.009	16062	36.6
Aroclor-1254	5	10.175	-0.001	20494	36.9	5	10.572	0.004	4818	18.0
Total CollAve (5 peaks):				42.6		Total Col2Ave (5 peaks):				36.4 RPD = 16
Corrected Ave (4 peaks):				39.6		Corrected Ave (4 peaks):				32.8 RPD = 19
Aroclor-1260	1	11.048	0.003	794	1.4	1	11.665	0.012	1652	5.0
Aroclor-1260	2	11.366	0.005	814	1.4	2	11.926	0.008	842	1.0
Aroclor-1260	3	11.739	0.006	1848	1.2	3	12.438	0.002	483	2.1
Aroclor-1260	4	12.145	0.006	1372	1.8	4	12.506	0.004	790	1.4
Aroclor-1260	5	---			0.0	NS	---			---
Total CollAve (4 peaks):				1.4		Total Col2Ave (4 peaks):				2.4 RPD = 49*
Corrected Ave (3 peaks):				1.3		Corrected Ave (3 peaks):				1.5 RPD = 12
Aroclor-1262	1	10.832	0.003	13157	27.6	1	11.121	-0.079	6113	12.7
Aroclor-1262	2	12.145	-0.098	1372	1.8	2	11.665	0.013	1652	4.0
Aroclor-1262	3	---			0.0	3	12.438	0.004	483	1.0
Aroclor-1262	4	13.038	0.051	842	1.1	4	12.506	0.004	790	1.1
Total CollAve (3 peaks):				10.1		Total Col2Ave (4 peaks):				4.7 RPD = 73*
Corrected Ave: < 3 Peaks						Corrected Ave (3 peaks):				2.0
Aroclor-1268	1	---			0.0	1	12.438	0.006	483	0.4
Aroclor-1268	2	---			0.0	2	12.506	0.006	790	0.6
Aroclor-1268	3	12.617	-0.082	5851	3.5	3	12.899	0.007	491	0.5
Aroclor-1268	4	13.500	0.010	1745	0.3	4	13.714	0.005	379	0.1
CollAve: <3 Quant Peaks						Col2Ave:				0.4

Total PCB Area Col1 (5.906 - 13.793) = 1149784 Col1 Total PCB = 0.1 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 572210 Col2 Total PCB = 0.1 ppm*

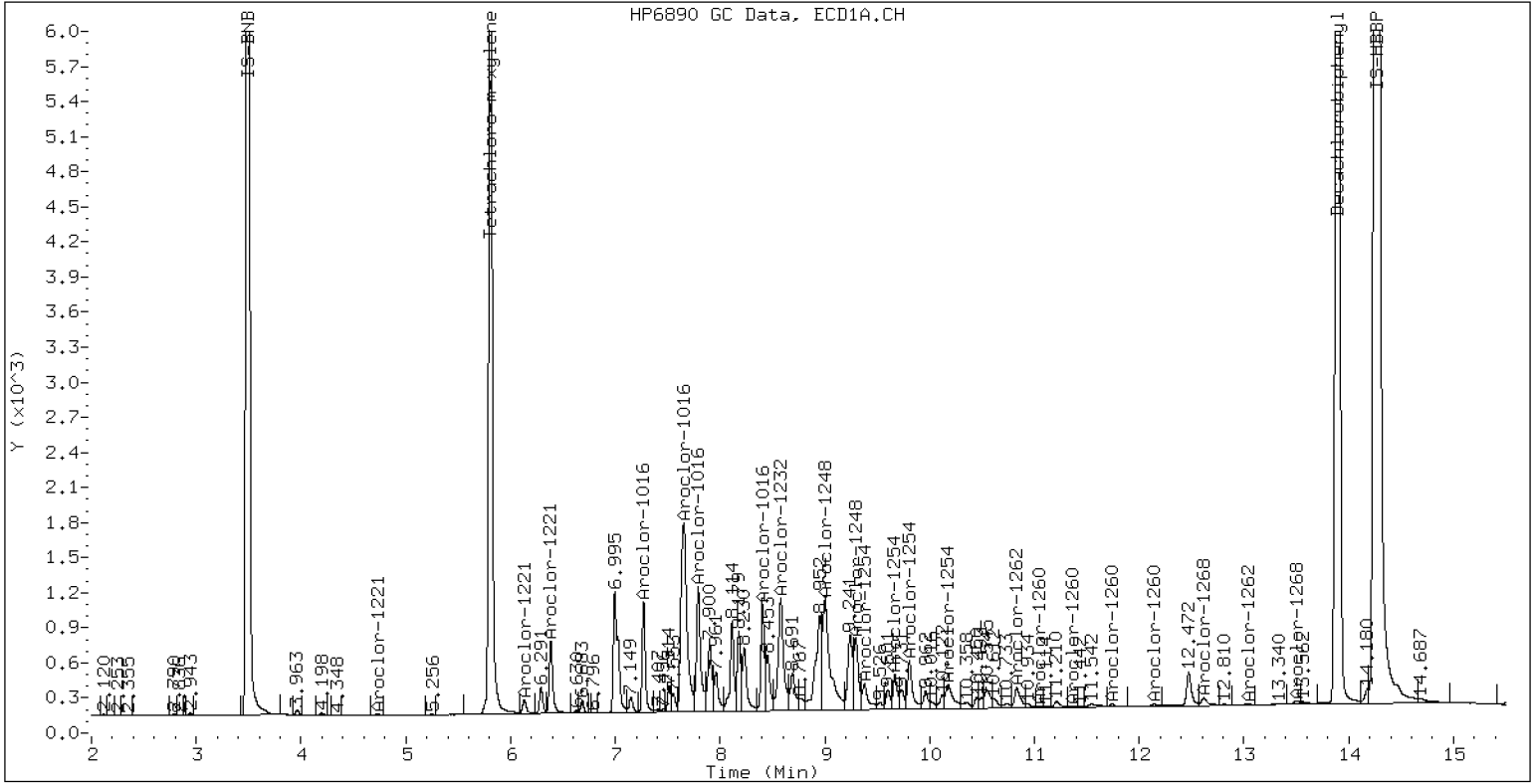
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1242SCV

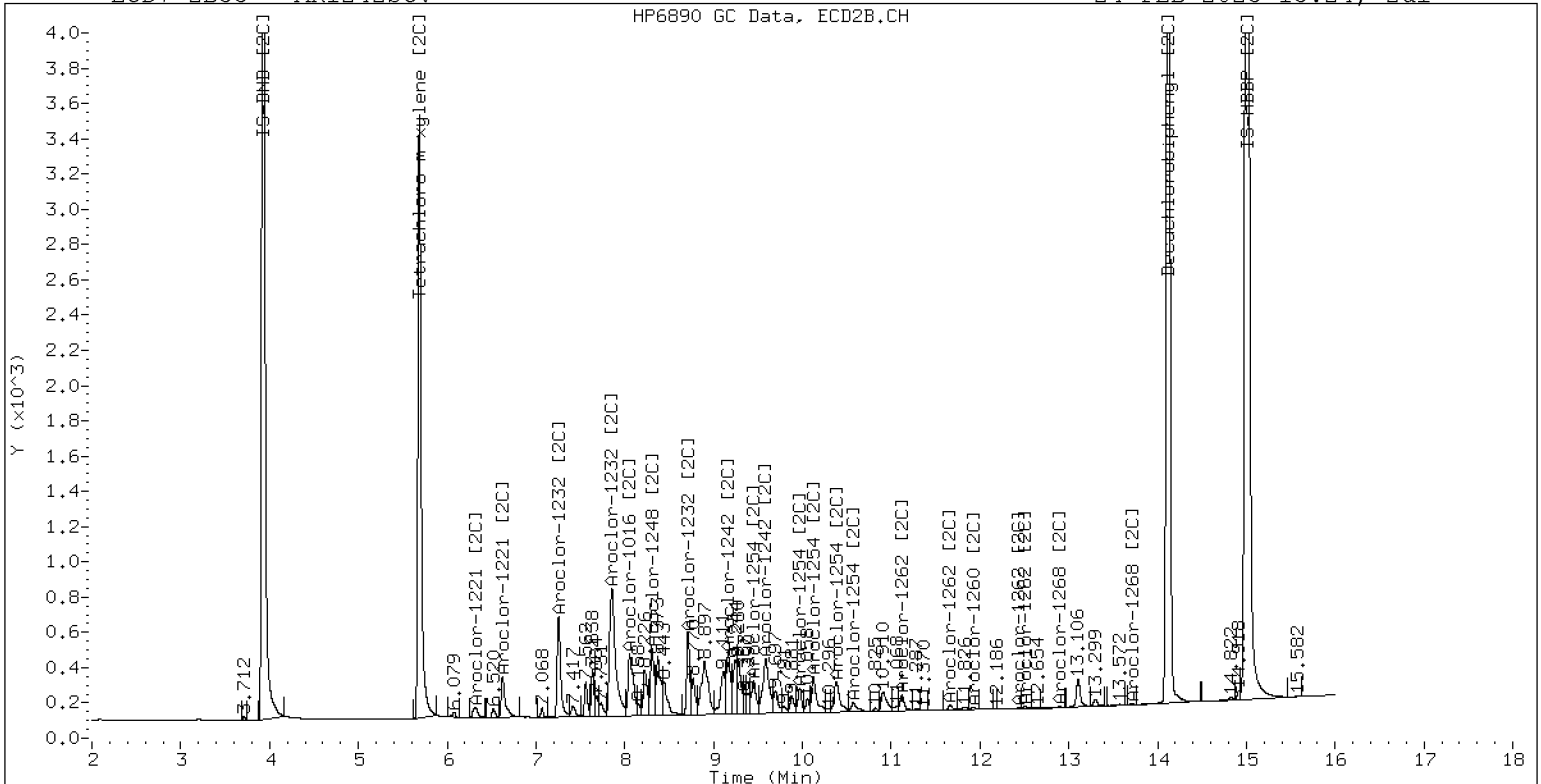
24-FEB-2023 15:24, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1242SCV

24-FEB-2023 15:24, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242315ECD7.D
Data file 2: /230224.b/230224.b/02242315ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1248SCV
Client ID:
Injection Date: 24-FEB-2023 15:45
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.002	336655	5.687	0.002	168719	34.9	36.4	4.2	Tetrachloro-m-xylene
13.894	0.001	499162	14.118	-0.001	308317	33.1	36.3	9.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	646554	-4.0
Hexabromobiphenyl	1429847	1529451	7.0

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	316066	0.3
Hexabromobiphenyl	513946	557213	8.4

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.271	0.000	19773	80.5	1	7.254	-0.002	16926	91.5
Aroclor-1016	2	7.653	-0.001	88099	117.7	2	7.857	0.001	45733	121.9
Aroclor-1016	3	7.794	0.003	35915	98.3	3	8.060	0.005	8078	47.7
Aroclor-1016	4	8.406	0.001	77842	329.5	4	8.307	0.000	37348	280.9
Total CollAve (4 peaks):				156.5		Total Col2Ave (4 peaks):				135.5 RPD = 14
Corrected Ave (3 peaks):				98.8		Corrected Ave (3 peaks):				87.0 RPD = 13
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	6.133	0.001	680	6.6	2	6.326	0.030	1966	34.7
Aroclor-1221	3	6.384	0.002	3390	14.1	3	6.631	0.009	1571	17.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	6.133	0.002	680	9.9	2	7.254	-0.000	16926	210.6
Aroclor-1232	3	7.653	-0.002	88099	283.3	3	7.857	-0.004	45733	284.6
Aroclor-1232	4	8.581	-0.000	99572	753.4	4	8.714	-0.001	38224	826.6
Total CollAve (3 peaks):				348.9		Total Col2Ave (3 peaks):				440.6 RPD = 23
Corrected Ave: < 3 Peaks						Corrected Ave: < 3 Peaks				
Aroclor-1242	1	7.271	-0.000	19773	98.7	1	7.254	-0.002	16926	115.3
Aroclor-1242	2	7.653	-0.003	88099	144.8	2	7.857	-0.001	45733	148.2
Aroclor-1242	3	8.406	0.000	77842	411.2	3	9.165	-0.002	45021	468.7
Aroclor-1242	4	8.581	0.001	99572	355.8	4	9.590	-0.008	53613	458.1
Total CollAve (4 peaks):				252.6		Total Col2Ave (4 peaks):				297.6 RPD = 16
Corrected Ave (3 peaks):				199.8		Corrected Ave (3 peaks):				240.5 RPD = 19
Aroclor-1248	1	8.406	0.000	77842	246.8	1	8.307	-0.001	37348	247.5
Aroclor-1248	2	8.581	0.000	99572	248.3	2	8.714	-0.000	38224	245.0
Aroclor-1248	3	8.998	-0.000	186857	247.0	3	9.165	-0.000	45021	250.7
Aroclor-1248	4	9.294	-0.000	98398	255.5	4	9.590	-0.001	53613	248.7
Total CollAve (4 peaks):				249.4		Total Col2Ave (4 peaks):				248.0 RPD = 1
Corrected Ave (3 peaks):				247.4		Corrected Ave (3 peaks):				247.0 RPD = 0
Aroclor-1254	1	9.294	-0.004	98398	151.6	1	9.450	0.001	21823	90.8
Aroclor-1254	2	9.377	-0.001	49616	169.9	2	9.971	0.001	19450	100.6
Aroclor-1254	3	9.669	0.001	40230	96.4	3	10.124	0.000	36574	87.5
Aroclor-1254	4	9.808	0.001	68500	84.4	4	10.389	0.016	35100	86.1
Aroclor-1254	5	10.183	0.007	47365	93.1	5	10.573	0.004	5676	22.9
Total CollAve (5 peaks):				119.1		Total Col2Ave (5 peaks):				77.6 RPD = 42*
Corrected Ave (4 peaks):				106.4		Corrected Ave (4 peaks):				71.8 RPD = 39
Aroclor-1260	1	11.047	0.003	1670	3.0	1	11.662	0.009	2055	6.3
Aroclor-1260	2	11.362	0.001	1111	1.9	2	11.924	0.007	1466	1.8
Aroclor-1260	3	11.739	0.005	2107	1.4	3	12.434	-0.002	573	2.6
Aroclor-1260	4	12.144	0.005	1379	1.8	4	12.505	0.003	1003	1.8
Aroclor-1260	5	12.251	0.006	698	2.1	NS	---			----
Total CollAve (5 peaks):				2.1		Total Col2Ave (4 peaks):				3.1 RPD = 41*
Corrected Ave (4 peaks):				1.8		Corrected Ave (3 peaks):				2.0 RPD = 12
Aroclor-1262	1	10.833	0.005	15355	32.7	1	11.122	-0.079	7225	15.2
Aroclor-1262	2	12.251	0.007	698	0.9	2	11.662	0.011	2055	5.1
Aroclor-1262	3	12.321	0.002	836	1.0	3	12.434	0.000	573	1.2
Aroclor-1262	4	12.991	0.004	1043	1.4	4	12.505	0.003	1003	1.4
Total CollAve (4 peaks):				9.0		Total Col2Ave (4 peaks):				5.7 RPD = 45*
Corrected Ave (3 peaks):				1.1		Corrected Ave (3 peaks):				2.6 RPD = 80*
Aroclor-1268	1	12.251	0.004	698	0.4	1	12.434	0.002	573	0.5
Aroclor-1268	2	12.321	0.004	836	0.4	2	12.505	0.005	1003	0.8
Aroclor-1268	3	12.700	0.001	2449	1.5	3	12.892	0.001	721	0.7
Aroclor-1268	4	13.493	0.003	7547	1.4	4	13.708	-0.001	2265	0.7
Total CollAve (4 peaks):				0.9		Total Col2Ave (4 peaks):				0.7 RPD = 29
Corrected Ave (3 peaks):				0.7		Corrected Ave (3 peaks):				0.6 RPD = 13

Total PCB Area Col1 (5.906 - 13.793) = 1574335 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 746330 Col2 Total PCB = 0.2 ppm*

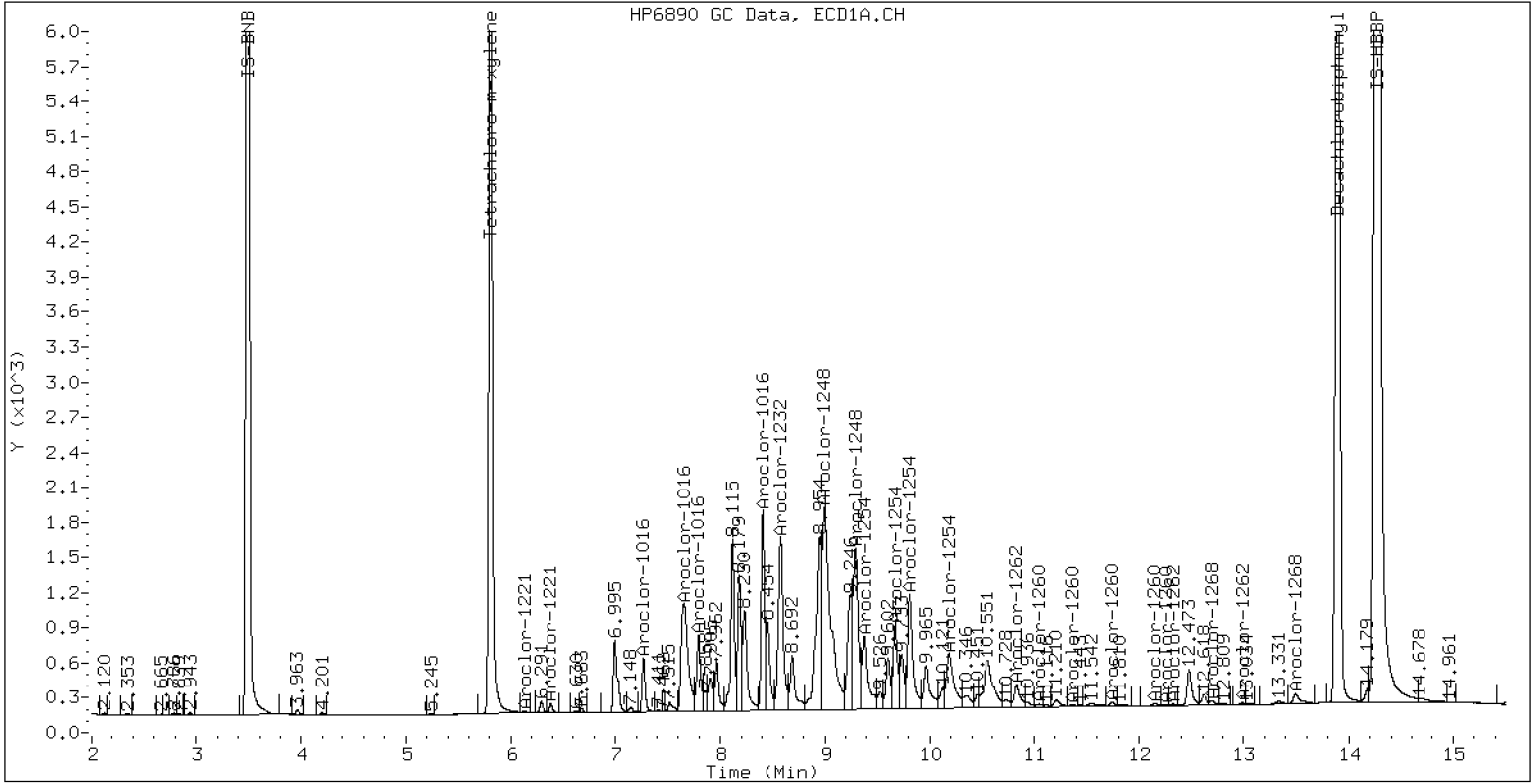
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1248SCV

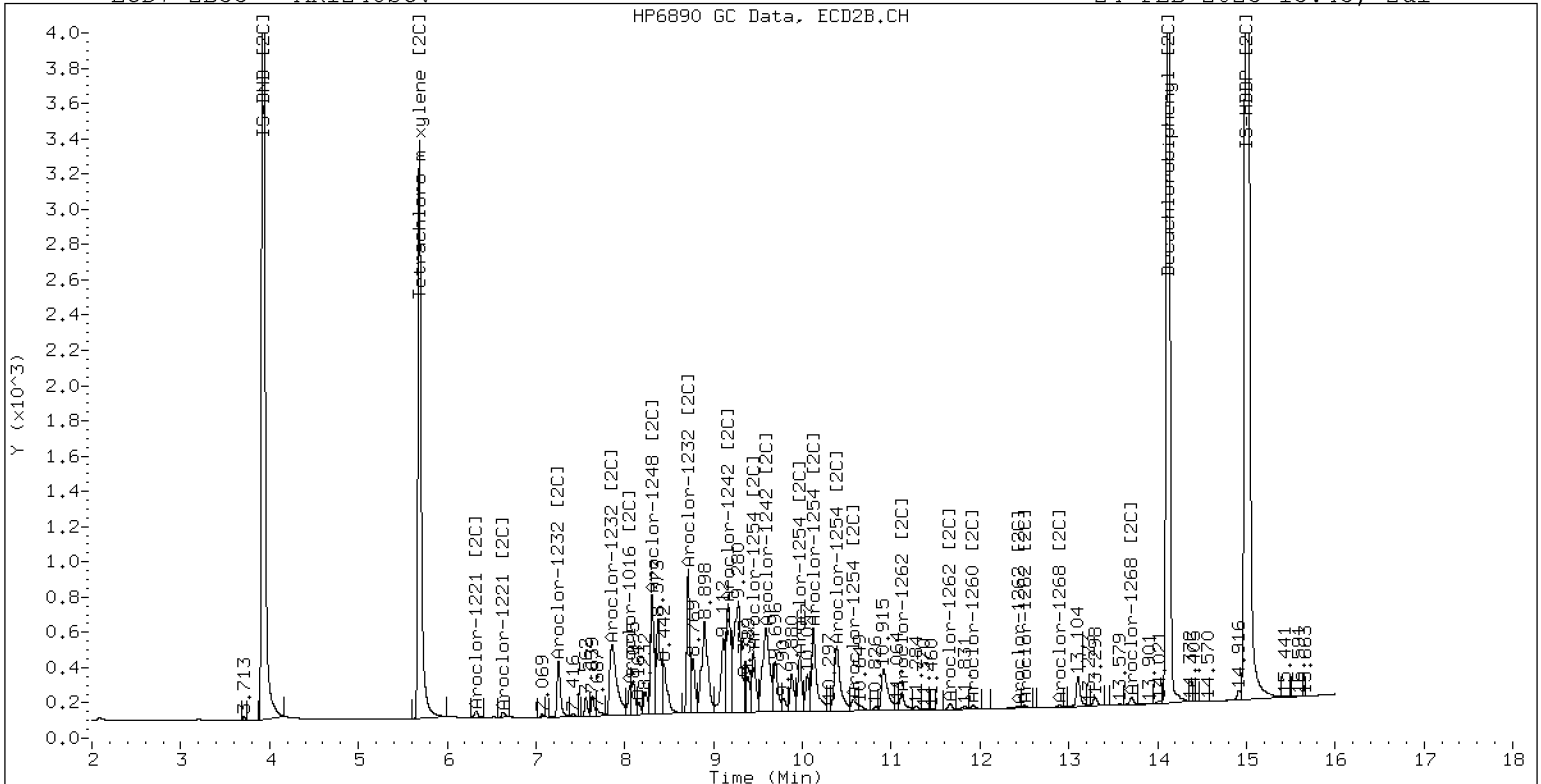
24-FEB-2023 15:45, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1248SCV

24-FEB-2023 15:45, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242316ECD7.D
Data file 2: /230224.b/230224.b/02242316ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1254SCV
Client ID:
Injection Date: 24-FEB-2023 16:06
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.806	-0.000	354312	5.686	0.001	174604	36.1	37.1	2.6	Tetrachloro-m-xylene
13.895	0.002	540961	14.119	-0.000	329134	34.6	37.9	9.0	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	656887	-2.5
Hexabromobiphenyl	1429847	1585505	10.9

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	320936	1.8
Hexabromobiphenyl	513946	570006	10.9

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	0.000	565	2.3	1	7.255	-0.001	387	2.1	
Aroclor-1016	2	7.656	0.002	1875	2.5	2	7.854	-0.002	860	2.3	
Aroclor-1016	3	7.792	0.002	1106	3.0	3	8.098	0.043	578	3.4	
Aroclor-1016	4	8.405	0.000	29924	124.7	4	8.307	0.000	21985	162.9	
Total CollAve (4 peaks):				33.1	Total Col2Ave (4 peaks):				42.6	RPD = 25	
Corrected Ave (3 peaks):				2.6	Corrected Ave (3 peaks):				2.6	RPD = 0	
Aroclor-1221	1	---			0.0	1	---			0.0	
Aroclor-1221	2	---			0.0	2	6.325	0.028	1947	33.9	
Aroclor-1221	3	---			0.0	3	6.637	0.015	368	3.9	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1232	1	---			0.0	1	---			0.0	
Aroclor-1232	2	---			0.0	2	7.255	0.001	387	4.7	
Aroclor-1232	3	7.656	0.000	1875	5.9	3	7.854	-0.007	860	5.3	
Aroclor-1232	4	8.583	0.002	12327	91.8	4	8.715	0.000	15013	319.7	
CollAve: <3 Quant Peaks					Col2Ave: 109.9						
Aroclor-1242	1	7.270	-0.000	565	2.8	1	7.255	-0.001	387	2.6	
Aroclor-1242	2	7.656	0.000	1875	3.0	2	7.854	-0.004	860	2.7	
Aroclor-1242	3	8.405	-0.000	29924	155.6	3	9.169	0.002	21933	224.9	
Aroclor-1242	4	8.583	0.003	12327	43.4	4	9.545	-0.053	34065	286.6	
Total CollAve (4 peaks):				51.2	Total Col2Ave (4 peaks):				129.2	RPD = 87*	
Corrected Ave (3 peaks):				16.4	Corrected Ave (3 peaks):				76.7	RPD = 130*	
Aroclor-1248	1	8.405	0.000	29924	93.4	1	8.307	-0.001	21985	143.5	
Aroclor-1248	2	8.583	0.002	12327	30.3	2	8.715	0.001	15013	94.8	
Aroclor-1248	3	8.992	-0.007	145580	189.4	3	9.169	0.004	21933	120.3	
Aroclor-1248	4	9.298	0.003	155450	397.3	4	9.545	-0.046	34065	155.6	
Total CollAve (4 peaks):				177.6	Total Col2Ave (4 peaks):				128.5	RPD = 32	
Corrected Ave (3 peaks):				104.3	Corrected Ave (3 peaks):				119.5	RPD = 14	
Aroclor-1254	1	9.298	-0.001	155450	235.7	1	9.450	0.001	58639	240.4	
Aroclor-1254	2	9.377	-0.001	69801	235.3	2	9.971	0.000	47008	239.5	
Aroclor-1254	3	9.668	-0.000	100839	237.8	3	10.124	0.000	100062	235.7	
Aroclor-1254	4	9.807	0.000	190544	231.1	4	10.373	0.000	99535	240.5	
Aroclor-1254	5	10.176	-0.000	122321	236.7	5	10.570	0.001	61549	244.2	
Total CollAve (5 peaks):				235.3	Total Col2Ave (5 peaks):				240.1	RPD = 2	
Corrected Ave (4 peaks):				234.7	Corrected Ave (4 peaks):				239.0	RPD = 2	
Aroclor-1260	1	11.043	-0.002	12288	21.5	1	11.661	0.008	29062	86.7	
Aroclor-1260	2	11.361	-0.001	13660	22.9	2	11.921	0.003	22238	26.0	
Aroclor-1260	3	11.736	0.002	37632	23.8	3	12.441	0.005	3555	15.7	
Aroclor-1260	4	12.141	0.002	27105	34.1	4	12.503	0.001	13126	22.8	
Aroclor-1260	5	12.320	0.076	2381	6.9	NS	---			---	
Total CollAve (5 peaks):				21.9	Total Col2Ave (4 peaks):				37.8	RPD = 53*	
Corrected Ave (4 peaks):				18.8	Corrected Ave (3 peaks):				21.5	RPD = 13	
Aroclor-1262	1	10.827	-0.002	220626	453.6	1	11.281	0.081	13562	27.9	
Aroclor-1262	2	12.320	0.076	2381	3.0	2	11.661	0.009	29062	70.1	
Aroclor-1262	3	---			0.0	3	12.441	0.007	3555	7.6	
Aroclor-1262	4	12.989	0.002	3225	4.1	4	12.503	0.001	13126	17.8	
Total CollAve (3 peaks):				153.6	Total Col2Ave (4 peaks):				30.8	RPD = 133*	
Corrected Ave: < 3 Peaks					Corrected Ave (3 peaks):				17.7		
Aroclor-1268	1	12.320	0.074	2381	1.2	1	12.441	0.009	3555	3.1	
Aroclor-1268	2	---			0.0	2	12.503	0.003	13126	10.6	
Aroclor-1268	3	12.701	0.002	2939	1.7	3	12.892	0.000	772	0.7	
Aroclor-1268	4	13.493	0.003	9164	1.6	4	13.707	-0.002	2801	0.8	
Total CollAve (3 peaks):				1.5	Total Col2Ave (4 peaks):				3.8	RPD = 87*	
Corrected Ave: < 3 Peaks					Corrected Ave (3 peaks):				1.6		

Total PCB Area Col1 (5.906 - 13.793) = 2118645 Col1 Total PCB = 0.3 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1007601 Col2 Total PCB = 0.3 ppm*

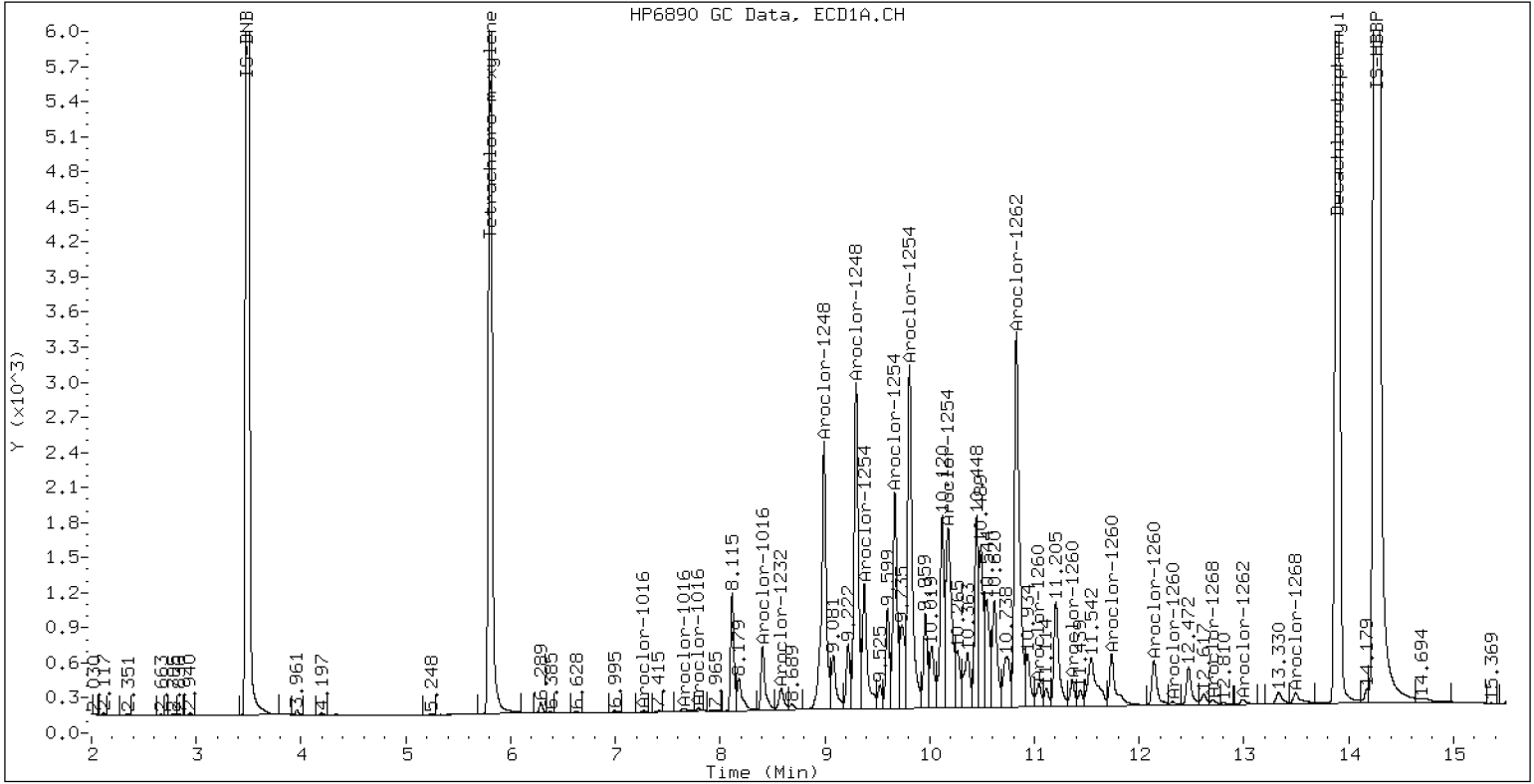
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1254SCV

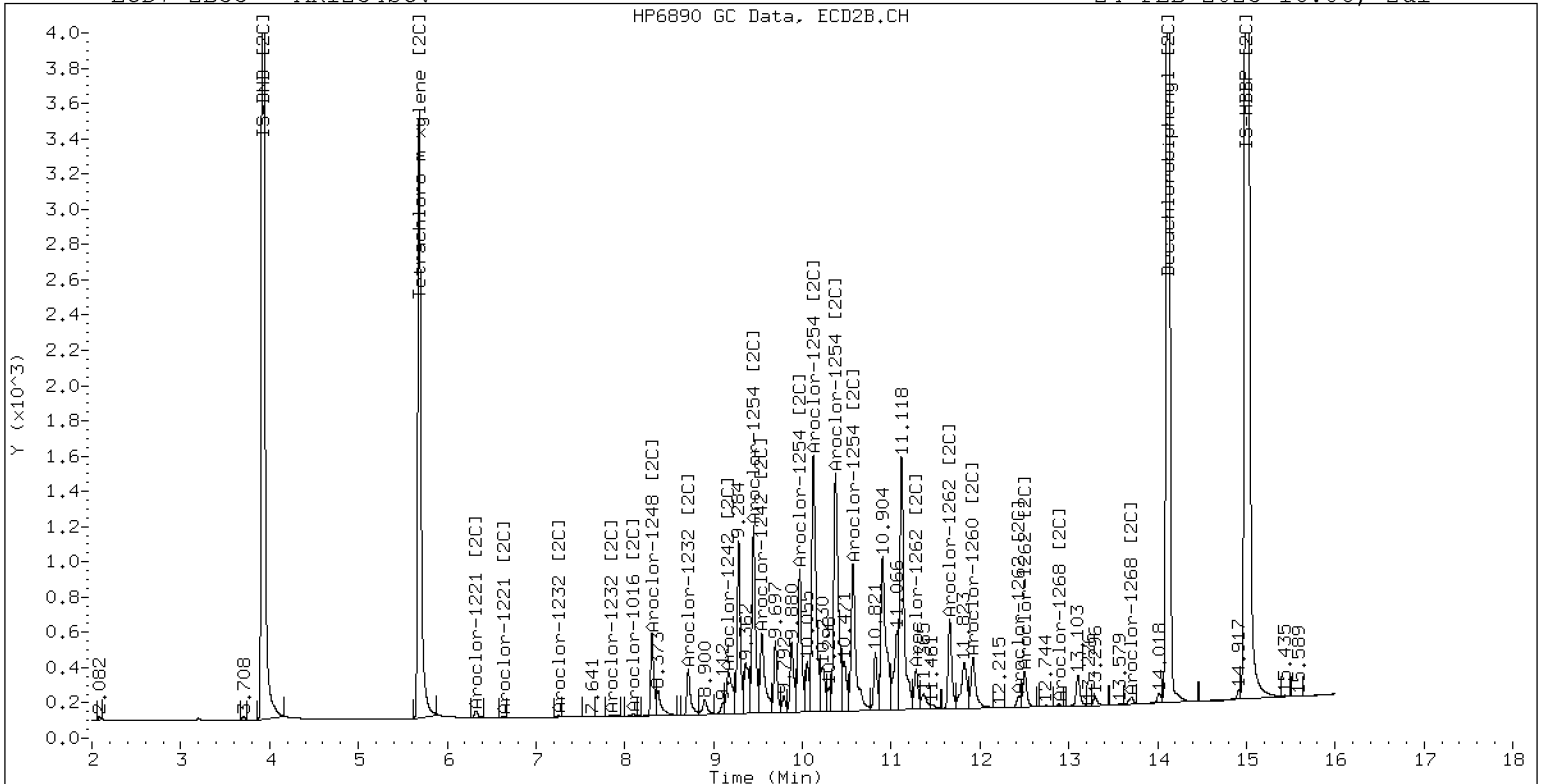
24-FEB-2023 16:06, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1254SCV

24-FEB-2023 16:06, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242317ECD7.D
Data file 2: /230224.b/230224.b/02242317ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR2162SCV
Client ID:
Injection Date: 24-FEB-2023 16:27
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.807	0.000	356001	5.685	0.000	170882	36.0	36.6	1.7	Tetrachloro-m-xylene
13.895	0.002	533971	14.119	0.000	326235	34.4	37.9	9.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	661953	-1.8
Hexabromobiphenyl	1429847	1574993	10.2

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	317807	0.8
Hexabromobiphenyl	513946	565951	10.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.269	-0.001	7175	28.5	1	7.256	0.000	3727	20.0	
Aroclor-1016	2	7.659	0.005	12893	16.8	2	7.863	0.007	5834	15.5	
Aroclor-1016	3	7.794	0.004	6936	18.5	3	8.063	0.009	2963	17.4	
Aroclor-1016	4	8.408	0.003	3610	14.9	4	8.308	0.002	2045	15.3	
Total CollAve (4 peaks):				19.7	Total Col2Ave (4 peaks):				17.0	RPD = 14	
Corrected Ave (3 peaks):				16.8	Corrected Ave (3 peaks):				16.1	RPD = 4	
Aroclor-1221	1	4.730	-0.000	15803	266.6	1	4.955	-0.001	7909	262.9	
Aroclor-1221	2	6.131	-0.001	26946	254.1	2	6.296	-0.000	14303	251.2	
Aroclor-1221	3	6.382	-0.000	62477	253.8	3	6.622	0.000	23612	254.7	
Total CollAve (3 peaks):				258.2	Total Col2Ave (3 peaks):				256.3	RPD = 1	
Corrected Ave: < 3 Peaks					Corrected Ave: < 3 Peaks						
Aroclor-1232	1	4.730	0.000	15803	445.6	1	4.955	-0.001	7909	486.4	
Aroclor-1232	2	6.131	0.000	26946	383.1	2	7.256	0.002	3727	46.1	
Aroclor-1232	3	7.659	0.003	12893	40.5	3	7.863	0.002	5834	36.1	
Aroclor-1232	4	8.583	0.003	2684	19.8	4	8.716	0.002	1189	25.6	
Total CollAve (4 peaks):				222.3	Total Col2Ave (4 peaks):				148.5	RPD = 40	
Corrected Ave (3 peaks):				147.8	Corrected Ave (3 peaks):				35.9	RPD = 122*	
Aroclor-1242	1	7.269	-0.001	7175	35.0	1	7.256	0.000	3727	25.2	
Aroclor-1242	2	7.659	0.003	12893	20.7	2	7.863	0.005	5834	18.8	
Aroclor-1242	3	8.408	0.002	3610	18.6	3	9.175	0.008	1082	11.2	
Aroclor-1242	4	8.583	0.004	2684	9.4	4	9.543	-0.054	1390	11.8	
Total CollAve (4 peaks):				20.9	Total Col2Ave (4 peaks):				16.8	RPD = 22	
Corrected Ave (3 peaks):				16.2	Corrected Ave (3 peaks):				13.9	RPD = 15	
Aroclor-1248	1	8.408	0.002	3610	11.2	1	8.308	0.001	2045	13.5	
Aroclor-1248	2	8.583	0.003	2684	6.5	2	8.716	0.002	1189	7.6	
Aroclor-1248	3	8.994	-0.005	24440	31.6	3	9.175	0.009	1082	6.0	
Aroclor-1248	4	9.302	0.008	26328	66.8	4	9.543	-0.048	1390	6.4	
Total CollAve (4 peaks):				29.0	Total Col2Ave (4 peaks):				8.4	RPD = 110*	
Corrected Ave (3 peaks):				16.4	Corrected Ave (3 peaks):				6.7	RPD = 85*	
Aroclor-1254	1	9.302	0.004	26328	39.6	1	9.452	0.003	9571	39.6	
Aroclor-1254	2	---			0.0	2	9.972	0.002	1733	8.9	
Aroclor-1254	3	9.670	0.002	3721	8.7	3	10.147	0.023	49218	117.1	
Aroclor-1254	4	9.808	0.000	9653	11.6	4	10.370	-0.002	59603	145.4	
Aroclor-1254	5	10.120	-0.056	131179	251.9	5	10.569	0.001	79533	318.7	
Total CollAve (4 peaks):				78.0	Total Col2Ave (5 peaks):				125.9	RPD = 47*	
Corrected Ave (3 peaks):				20.0	Corrected Ave (4 peaks):				77.8	RPD = 118*	
Aroclor-1260	1	11.044	-0.000	223208	394.0	1	11.652	-0.001	104071	312.7	
Aroclor-1260	2	11.361	-0.001	190166	321.2	2	11.919	0.002	251579	296.2	
Aroclor-1260	3	11.737	0.003	458281	291.9	3	12.435	-0.001	113645	504.2	
Aroclor-1260	4	12.141	0.002	149720	189.4	4	12.501	-0.001	182951	319.6	
Aroclor-1260	5	12.244	0.000	196033	576.0	NS	---			----	
Total CollAve (5 peaks):				354.5	Total Col2Ave (4 peaks):				358.2	RPD = 1	
Corrected Ave (4 peaks):				299.1	Corrected Ave (3 peaks):				309.5	RPD = 3	
Aroclor-1262	1	10.828	-0.001	121431	251.3	1	11.201	0.000	121335	251.1	
Aroclor-1262	2	12.244	0.000	196033	249.3	2	11.652	0.000	104071	252.9	
Aroclor-1262	3	12.319	0.001	211092	249.8	3	12.435	0.001	113645	243.4	
Aroclor-1262	4	12.988	0.001	183455	237.5	4	12.501	-0.001	182951	250.1	
Total CollAve (4 peaks):				247.0	Total Col2Ave (4 peaks):				249.3	RPD = 1	
Corrected Ave (3 peaks):				245.5	Corrected Ave (3 peaks):				248.2	RPD = 1	
Aroclor-1268	1	12.244	-0.002	196033	97.1	1	12.435	0.003	113645	99.7	
Aroclor-1268	2	12.319	0.002	211092	105.6	2	12.501	0.001	182951	149.3	
Aroclor-1268	3	12.723	0.024	77240	45.2	3	12.891	-0.000	7755	7.4	
Aroclor-1268	4	13.488	-0.002	65479	11.6	4	13.709	0.000	35146	10.5	
Total CollAve (4 peaks):				64.9	Total Col2Ave (4 peaks):				66.7	RPD = 3	

Corrected Ave (3 peaks): 51.3 Corrected Ave (3 peaks): 39.2 RPD = 27

Total PCB Area Col1 (5.906 - 13.793) = 3239932 Col1 Total PCB = 0.4 ppm*
Total PCB Area Col2 (5.785 - 14.019) = 1655522 Col2 Total PCB = 0.4 ppm*

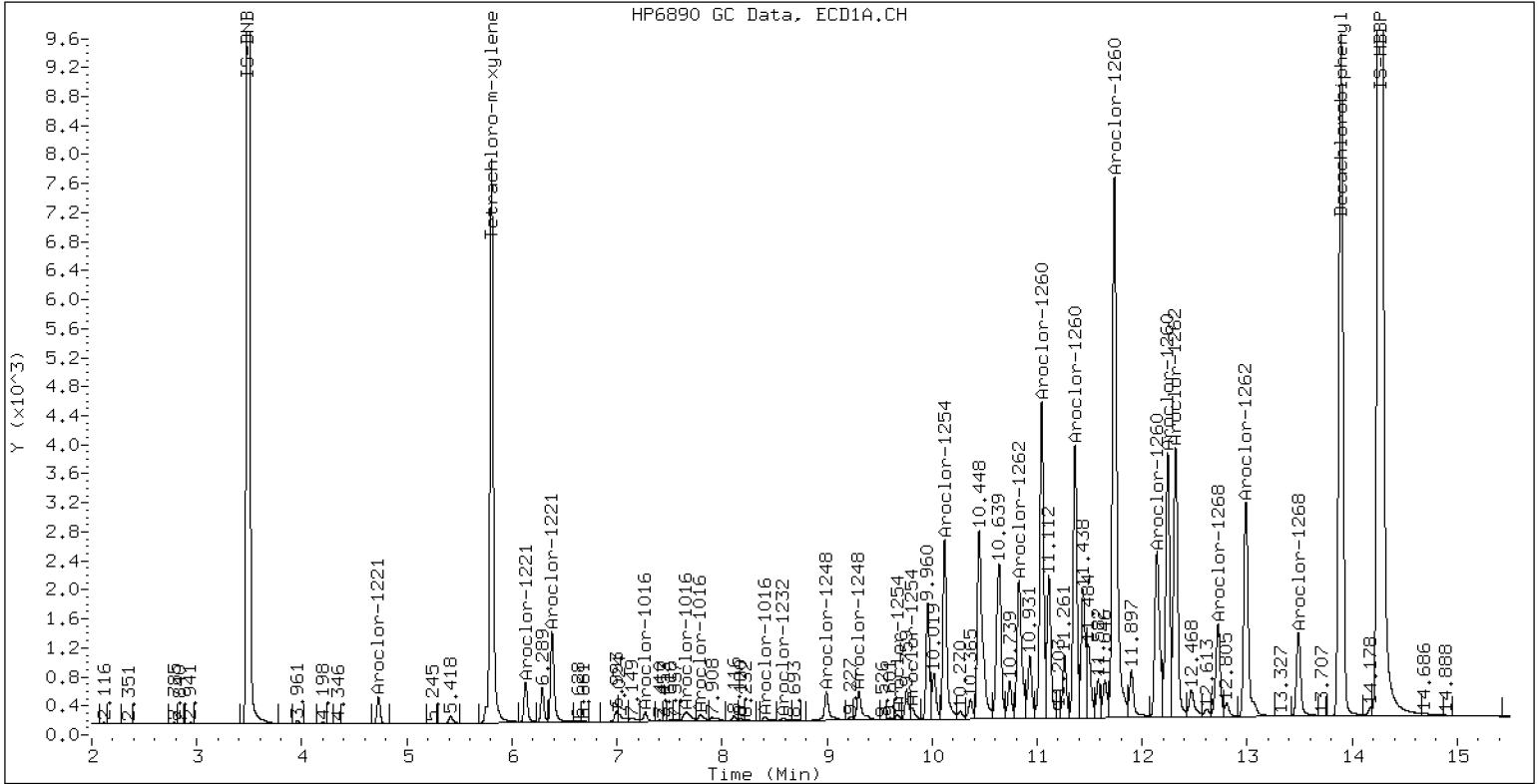
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR2162SCV

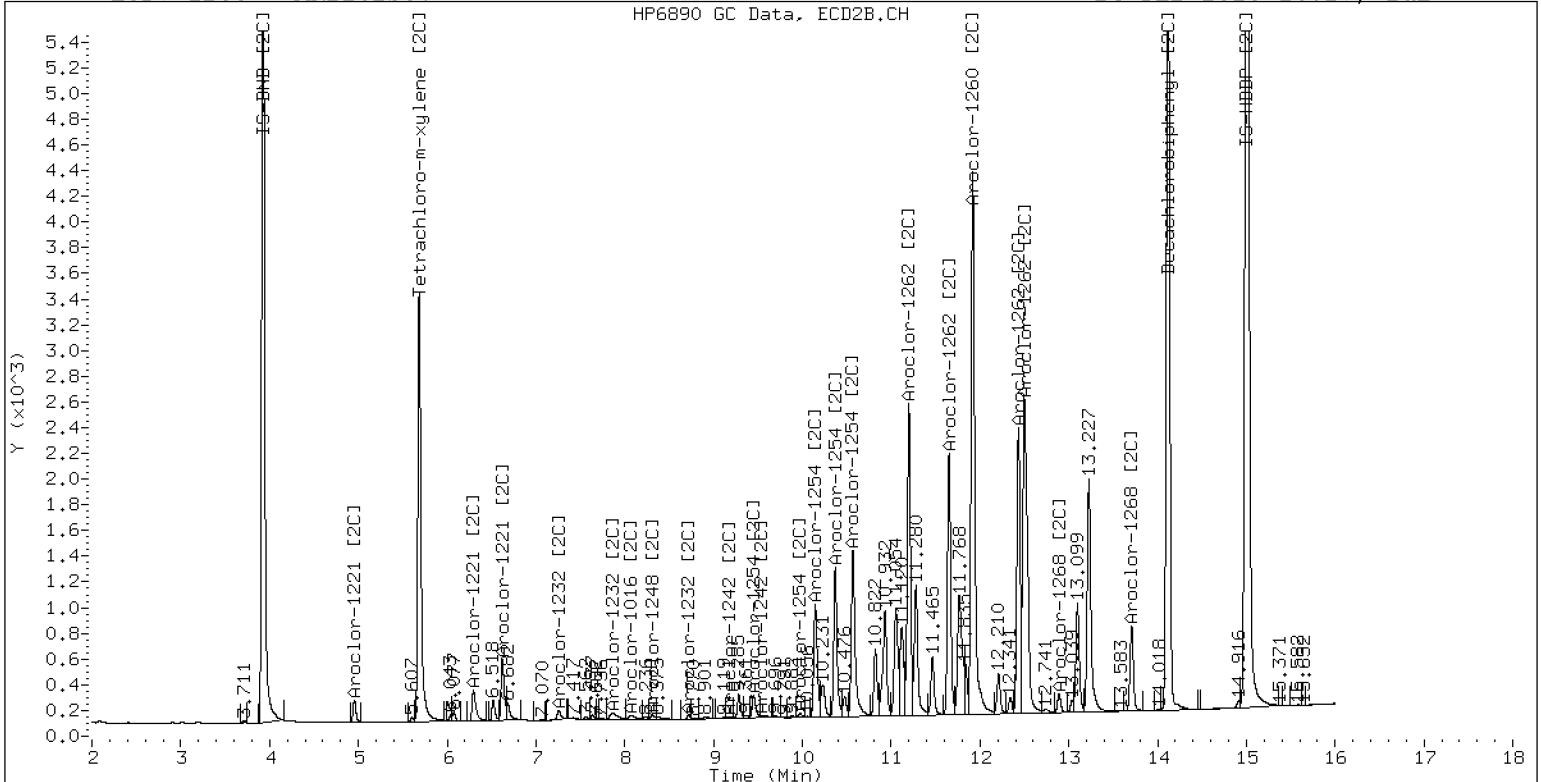
24-FEB-2023 16:27, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR2162SCV

24-FEB-2023 16:27, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242318ECD7.D
Data file 2: /230224.b/230224.b/02242318ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR3268SCV
Client ID:
Injection Date: 24-FEB-2023 16:48
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.806	0.000	363331	5.685	0.000	176204	37.1	38.2	2.9	Tetrachloro-m-xylene
13.894	0.001	800845	14.118	-0.001	488290	51.3	56.4	9.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	656592	-2.6
Hexabromobiphenyl	1429847	1584453	10.8

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	314741	-0.2
Hexabromobiphenyl	513946	568346	10.6

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	-0.001	28327	113.6	1	7.254	-0.001	20651	112.1	
Aroclor-1016	2	7.657	0.003	80668	106.1	2	7.861	0.005	41326	110.6	
Aroclor-1016	3	7.793	0.003	40661	109.6	3	8.060	0.005	20446	121.2	
Aroclor-1016	4	8.407	0.002	24680	102.9	4	8.308	0.001	13576	102.5	
Total CollAve (4 peaks):				108.0	Total Col2Ave (4 peaks):				111.6	RPD = 3	
Corrected Ave (3 peaks):				106.2	Corrected Ave (3 peaks):				108.4	RPD = 2	
Aroclor-1221	1	4.729	-0.001	8535	145.1	1	4.956	-0.000	3965	133.1	
Aroclor-1221	2	6.132	-0.000	15523	147.6	2	6.297	0.001	8689	154.1	
Aroclor-1221	3	6.382	-0.000	45872	187.9	3	6.622	0.001	22272	242.6	
Total CollAve (3 peaks):				160.2	Total Col2Ave (3 peaks):				176.6	RPD = 10	
Corrected Ave: < 3 Peaks					Corrected Ave: < 3 Peaks						
Aroclor-1232	1	4.729	-0.001	8535	242.6	1	4.956	0.000	3965	246.2	
Aroclor-1232	2	6.132	0.001	15523	222.5	2	7.254	0.000	20651	258.1	
Aroclor-1232	3	7.657	0.001	80668	255.4	3	7.861	0.001	41326	258.3	
Aroclor-1232	4	8.582	0.001	34784	259.2	4	8.714	-0.001	12504	271.5	
Total CollAve (4 peaks):				244.9	Total Col2Ave (4 peaks):				258.5	RPD = 5	
Corrected Ave (3 peaks):				240.2	Corrected Ave (3 peaks):				254.2	RPD = 6	
Aroclor-1242	1	7.270	-0.001	28327	139.2	1	7.254	-0.001	20651	141.2	
Aroclor-1242	2	7.657	0.001	80668	130.5	2	7.861	0.003	41326	134.4	
Aroclor-1242	3	8.407	0.001	24680	128.4	3	9.170	0.003	12830	134.1	
Aroclor-1242	4	8.582	0.003	34784	122.4	4	9.600	0.003	14836	127.3	
Total CollAve (4 peaks):				130.1	Total Col2Ave (4 peaks):				134.3	RPD = 3	
Corrected Ave (3 peaks):				127.1	Corrected Ave (3 peaks):				132.0	RPD = 4	
Aroclor-1248	1	8.407	0.001	24680	77.0	1	8.308	0.000	13576	90.3	
Aroclor-1248	2	8.582	0.001	34784	85.4	2	8.714	-0.000	12504	80.5	
Aroclor-1248	3	8.996	-0.003	83592	108.8	3	9.170	0.004	12830	71.8	
Aroclor-1248	4	9.292	-0.003	39603	101.3	4	9.600	0.010	14836	69.1	
Total CollAve (4 peaks):				93.1	Total Col2Ave (4 peaks):				77.9	RPD = 18	
Corrected Ave (3 peaks):				87.9	Corrected Ave (3 peaks):				73.8	RPD = 17	
Aroclor-1254	1	9.292	-0.007	39603	60.1	1	9.452	0.003	4590	19.2	
Aroclor-1254	2	9.377	-0.000	11450	38.6	2	9.973	0.003	2892	15.0	
Aroclor-1254	3	9.674	0.005	6387	15.1	3	10.131	0.007	6052	14.5	
Aroclor-1254	4	9.813	0.006	10162	12.3	4	10.390	0.017	5324	13.1	
Aroclor-1254	5	10.189	0.012	6862	13.3	5	10.572	0.004	1891	7.7	
Total CollAve (5 peaks):				27.9	Total Col2Ave (5 peaks):				13.9	RPD = 67*	
Corrected Ave (4 peaks):				19.8	Corrected Ave (4 peaks):				12.6	RPD = 45*	
Aroclor-1260	1	11.046	0.002	87033	152.7	1	11.645	-0.008	62543	187.1	
Aroclor-1260	2	11.362	0.001	6300	10.6	2	11.920	0.003	28552	33.5	
Aroclor-1260	3	11.738	0.004	54524	34.5	3	12.432	-0.004	285450	1261.2	
Aroclor-1260	4	12.144	0.005	1727	2.2	4	12.499	-0.002	306992	534.0	
Aroclor-1260	5	12.246	0.002	502931	1469.0	NS	---			----	
Total CollAve (5 peaks):				333.8	Total Col2Ave (4 peaks):				503.9	RPD = 41*	
Corrected Ave (4 peaks):				50.0	Corrected Ave (3 peaks):				251.5	RPD = 134*	
Aroclor-1262	1	10.832	0.004	3395	7.0	1	11.201	0.001	44255	91.2	
Aroclor-1262	2	12.246	0.002	502931	635.9	2	11.645	-0.007	62543	151.3	
Aroclor-1262	3	12.318	-0.000	497006	584.5	3	12.432	-0.002	285450	608.7	
Aroclor-1262	4	12.987	-0.000	202197	260.2	4	12.499	-0.003	306992	417.9	
Total CollAve (4 peaks):				371.9	Total Col2Ave (4 peaks):				317.3	RPD = 16	
Corrected Ave (3 peaks):				283.9	Corrected Ave (3 peaks):				220.1	RPD = 25	
Aroclor-1268	1	12.246	-0.001	502931	247.7	1	12.432	-0.000	285450	249.4	
Aroclor-1268	2	12.318	0.002	497006	247.2	2	12.499	-0.001	306992	249.5	
Aroclor-1268	3	12.699	-0.000	422793	245.8	3	12.892	0.000	260893	248.4	
Aroclor-1268	4	13.490	0.000	1386953	244.9	4	13.709	-0.000	829733	247.1	
Total CollAve (4 peaks):				246.4	Total Col2Ave (4 peaks):				248.6	RPD = 1	

Corrected Ave (3 peaks): 246.0 Corrected Ave (3 peaks): 248.3 RPD = 1

Total PCB Area Col1 (5.906 - 13.793) = 4180607 Col1 Total PCB = 0.5 ppm*
Total PCB Area Col2 (5.785 - 14.019) = 2376912 Col2 Total PCB = 0.6 ppm*

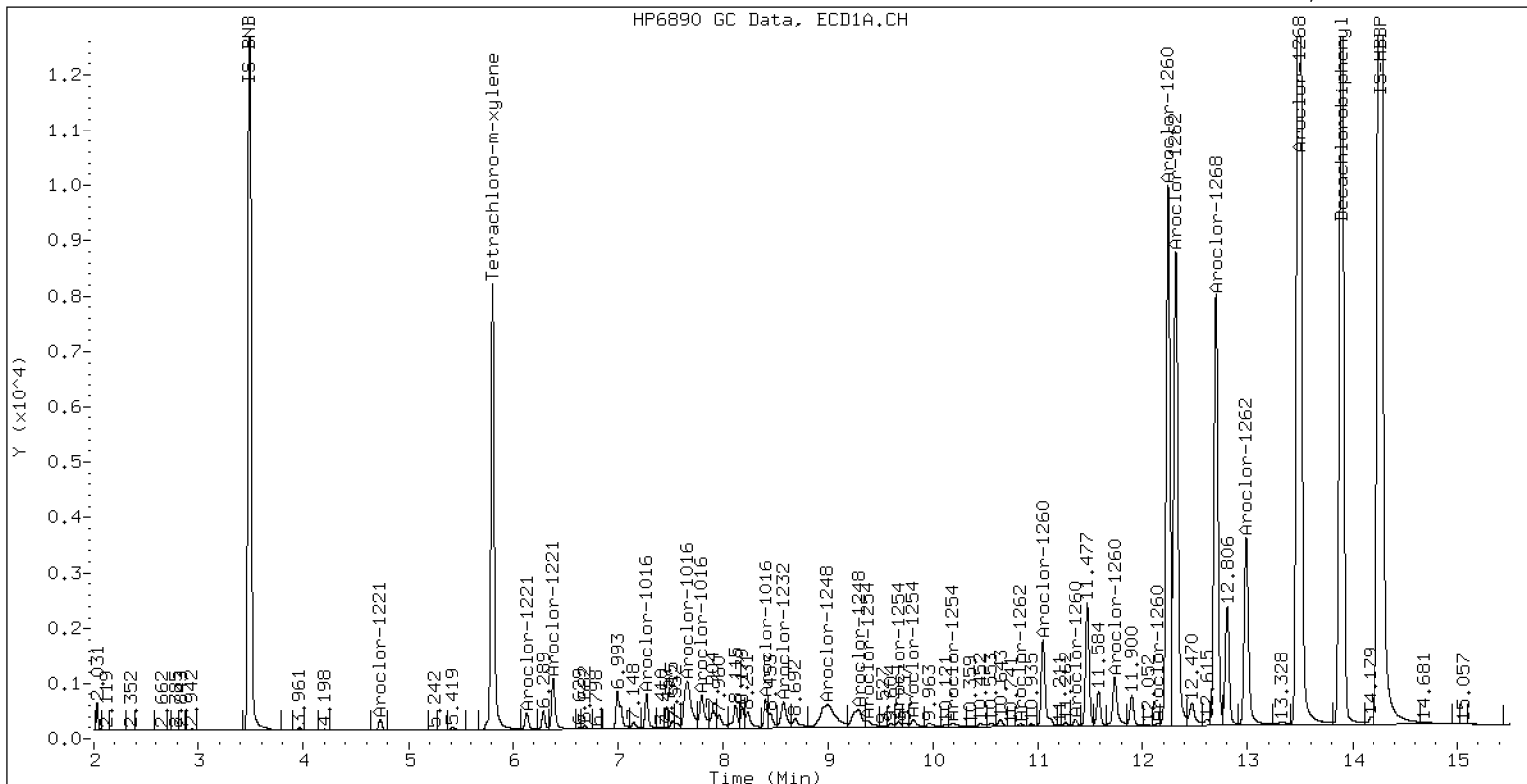
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR3268SCV

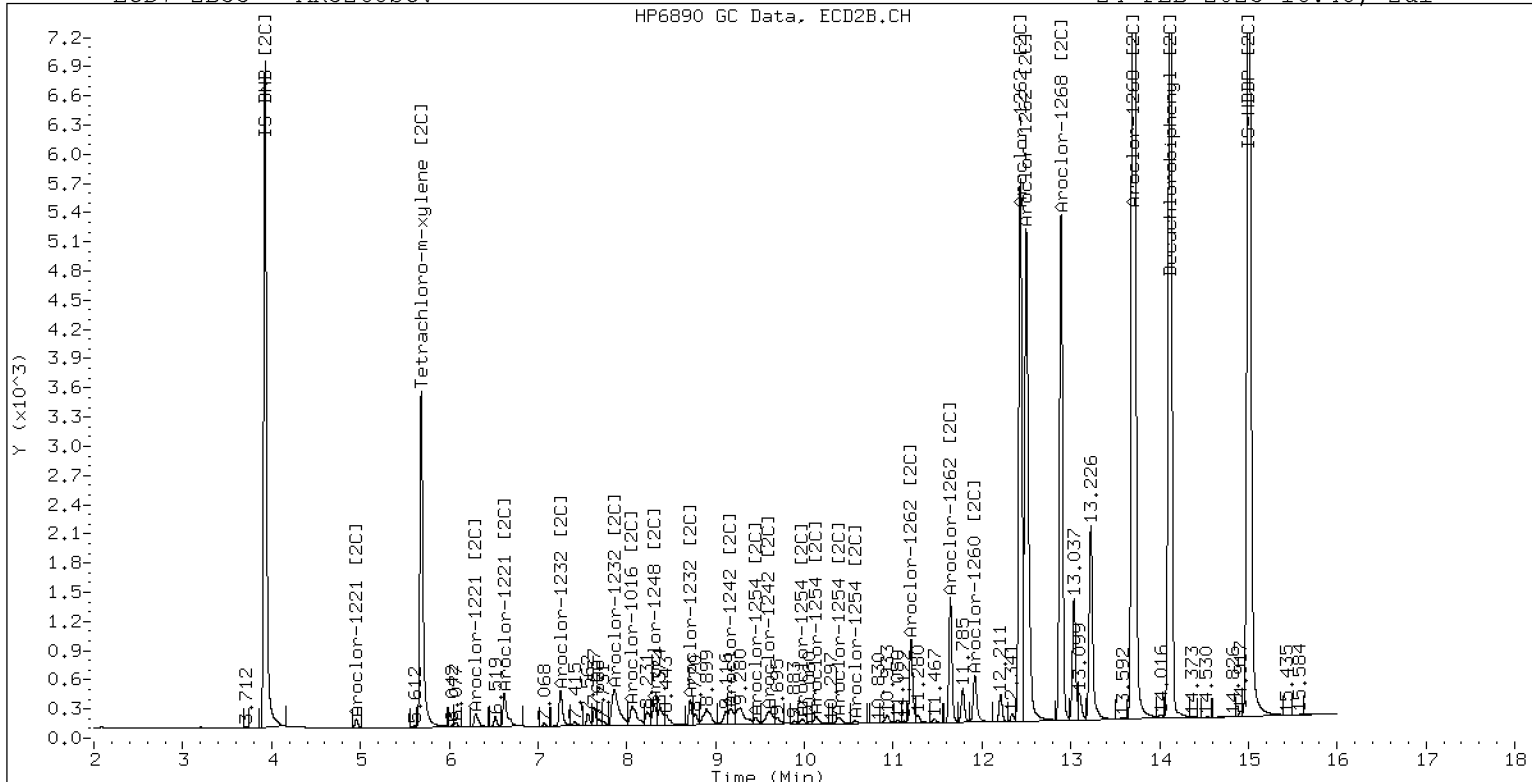
24-FEB-2023 16:48, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 AR3268SCV

24-FEB-2023 16:48, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
8082 DDT SCREEN REPORT

Data file 1: /230224.b/02242319ECD7.D

ARI ID: DDTS

RT	ZB5 Col Shift Response		RT	ZB35 Col Shift Response		ZB5 on col	ZB35 on col	RPD	Compound/Flag
9.261	0.000	694353	9.912	0.000	580269	0.100	0.100	0.0	2,4-DDE
0.000	-10.293	0	10.672	0.000	673479	0.000	0.200#	----	2,4-DDT
9.686	0.000	1191406	10.212	0.000	433373	0.100	0.100	0.0	4,4-DDE
10.259	0.000	1721760	10.672	0.000	673479	0.100	0.200#	66.7*	4,4-DDD

Indicates value is from co-eluting peaks

* Indicates RPD > 40%

Analytical Resources Inc.
8082 DDT SCREEN REPORT

Data file 1: /230224.b/02242320ECD7.D

ARI ID: DDT BD

RT	ZB5 Col Shift Response		RT	ZB35 Col Shift Response		ZB5 on col	ZB35 on col	RPD	Compound/Flag
9.285	0.023	4923	9.921	0.009	9972	0.001	0.002	84.3*	2,4-DDE
0.000	-10.293	0	10.677	0.004	249094	0.000	0.074#	----	2,4-DDT
9.692	0.006	12128	10.221	0.009	528	0.001	0.000	156.7*	4,4-DDE
10.265	0.006	410017	10.677	0.004	249094	0.023	0.074#	103.6*	4,4-DDD

Indicates value is from co-eluting peaks

* Indicates RPD > 40%



ANALYSIS SEQUENCE

SLB0342

Instrument: ECD7
Calibration ID: GB00069

Printed: 2/28/2023 9:54:44AM

Lab Number	Analysis	Container	Order	Position	STD ID	ISTD ID	Client	Comments
SLB0342-CAL1	QC		1		L000856	L000844		
SLB0342-CAL2	QC		2		L000859	L000844		
SLB0342-CAL3	QC		3		L000858	L000844		
SLB0342-CAL4	QC		4		L000731	L000844		
SLB0342-CAL5	QC		5		L000857	L000844		
SLB0342-CAL6	QC		6		L000855	L000844		
SLB0342-CAL7	QC		7		L000860	L000844		
SLB0342-CAL8	QC		8		L000861	L000844		
SLB0342-CAL9	QC		9		L000862	L000844		
SLB0342-CALA	QC		10		L000863	L000844		
SLB0342-CALB	QC		11		L000864	L000844		
SLB0342-SCV1	QC		12		L002065	L000844		
SLB0342-SCV2	QC		13		K007656	L000844		
SLB0342-SCV3	QC		14		L002066	L000844		
SLB0342-SCV4	QC		15		L002067	L000844		
SLB0342-SCV5	QC		16		L002068	L000844		
SLB0342-SCV6	QC		17		L002069	L000844		

Samples Loaded By Date

Data Processed By Date

GC LOG SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

	Inject	Date/Time	Filename	DF	LabID	ClientID
1	24-FEB-2023	10:51	02242301ECD7.D	1	IB	
2	24-FEB-2023	11:12	02242302ECD7.D	1	0.25PPMAR1660	
3	24-FEB-2023	11:33	02242303ECD7.D	1	0.02PPMAR1660	
4	24-FEB-2023	11:54	02242304ECD7.D	1	0.05PPMAR1660	
5	24-FEB-2023	12:15	02242305ECD7.D	1	1.0PPMAR1660	
6	24-FEB-2023	12:36	02242306ECD7.D	1	0.1PPMAR1660	
7	24-FEB-2023	12:57	02242307ECD7.D	1	0.5PPMAR1660	
8	24-FEB-2023	13:18	02242308ECD7.D	1	0.25PPMAR1242	
9	24-FEB-2023	13:39	02242309ECD7.D	1	0.25PPMAR1248	
10	24-FEB-2023	14:00	02242310ECD7.D	1	0.25PPMAR1254	
11	24-FEB-2023	14:21	02242311ECD7.D	1	0.25PPMAR2162	
12	24-FEB-2023	14:42	02242312ECD7.D	1	0.25PPMAR3268	
13	24-FEB-2023	15:03	02242313ECD7.D	1	AR1660SCV	
14	24-FEB-2023	15:24	02242314ECD7.D	1	AR1242SCV	
15	24-FEB-2023	15:45	02242315ECD7.D	1	AR1248SCV	
16	24-FEB-2023	16:06	02242316ECD7.D	1	AR1254SCV	
17	24-FEB-2023	16:27	02242317ECD7.D	1	AR2162SCV	
18	24-FEB-2023	16:48	02242318ECD7.D	1	AR3268SCV	
19	24-FEB-2023	17:09	02242319ECD7.D	1	DDTS	
20	24-FEB-2023	17:30	02242320ECD7.D	1	DDT BD	

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

ARI Job No.: IB Method: PCB.m Instrument: ecd7.i Date: 24-FEB-2023

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1051	02242301ECD7.D	IB		1	NO MANUAL INTEGRATION
1112	02242302ECD7.D	0.25PPMAR1660		1	NO MANUAL INTEGRATION
1133	02242303ECD7.D	0.02PPMAR1660		1	NO MANUAL INTEGRATION
1154	02242304ECD7.D	0.05PPMAR1660		1	NO MANUAL INTEGRATION
1215	02242305ECD7.D	1.0PPMAR1660		1	NO MANUAL INTEGRATION
1236	02242306ECD7.D	0.1PPMAR1660		1	NO MANUAL INTEGRATION
1257	02242307ECD7.D	0.5PPMAR1660		1	NO MANUAL INTEGRATION
1318	02242308ECD7.D	0.25PPMAR1242		1	NO MANUAL INTEGRATION
1339	02242309ECD7.D	0.25PPMAR1248		1	NO MANUAL INTEGRATION
1400	02242310ECD7.D	0.25PPMAR1254		1	NO MANUAL INTEGRATION
1421	02242311ECD7.D	0.25PPMAR2162		1	NO MANUAL INTEGRATION
1442	02242312ECD7.D	0.25PPMAR3268		1	NO MANUAL INTEGRATION
1503	02242313ECD7.D	AR1660SCV		1	NO MANUAL INTEGRATION
1524	02242314ECD7.D	AR1242SCV		1	NO MANUAL INTEGRATION
1545	02242315ECD7.D	AR1248SCV		1	NO MANUAL INTEGRATION
1606	02242316ECD7.D	AR1254SCV		1	NO MANUAL INTEGRATION
1627	02242317ECD7.D	AR2162SCV		1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1648	02242318ECD7.D	AR3268SCV		1	NO MANUAL INTEGRATION
1709	02242319ECD7.D	DDTS		1	NO MANUAL INTEGRATION
1730	02242320ECD7.D	DDT BD		1	NO MANUAL INTEGRATION
1751	02242321ECD7.D			1	NO MANUAL INTEGRATION
1812	02242322ECD7.D			1	NO MANUAL INTEGRATION
1833	02242323ECD7.D			1	NO MANUAL INTEGRATION
1854	02242324ECD7.D			1	NO MANUAL INTEGRATION
1915	02242325ECD7.D			1	NO MANUAL INTEGRATION
1936	02242326ECD7.D			1	NO MANUAL INTEGRATION
1957	02242327ECD7.D			1	NO MANUAL INTEGRATION
2018	02242328ECD7.D			1	NO MANUAL INTEGRATION
2039	02242329ECD7.D			1	NO MANUAL INTEGRATION
2059	02242330ECD7.D			1	NO MANUAL INTEGRATION
2120	02242331ECD7.D			1	NO MANUAL INTEGRATION
2141	02242332ECD7.D			1	NO MANUAL INTEGRATION
2202	02242333ECD7.D			1	NO MANUAL INTEGRATION
2223	02242334ECD7.D			1	NO MANUAL INTEGRATION
2244	02242335ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
2305	02242336ECD7.D			1	NO MANUAL INTEGRATION
2326	02242337ECD7.D			1	NO MANUAL INTEGRATION
2347	02242338ECD7.D			1	NO MANUAL INTEGRATION
0008	02242339ECD7.D			1	NO MANUAL INTEGRATION
0029	02242340ECD7.D			1	NO MANUAL INTEGRATION
0050	02242341ECD7.D			1	NO MANUAL INTEGRATION
0111	02242342ECD7.D			1	NO MANUAL INTEGRATION
0132	02242343ECD7.D			1	NO MANUAL INTEGRATION
0153	02242344ECD7.D			1	NO MANUAL INTEGRATION
0214	02242345ECD7.D			1	NO MANUAL INTEGRATION
0235	02242346ECD7.D			1	NO MANUAL INTEGRATION
0256	02242347ECD7.D			1	NO MANUAL INTEGRATION
0317	02242348ECD7.D			1	NO MANUAL INTEGRATION
0338	02242349ECD7.D			1	NO MANUAL INTEGRATION
0359	02242350ECD7.D			1	NO MANUAL INTEGRATION
0420	02242351ECD7.D			1	NO MANUAL INTEGRATION
0441	02242352ECD7.D			1	NO MANUAL INTEGRATION
0502	02242353ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0523	02242354ECD7.D			1	NO MANUAL INTEGRATION
0544	02242355ECD7.D			1	NO MANUAL INTEGRATION
0605	02242356ECD7.D			1	NO MANUAL INTEGRATION
0626	02242357ECD7.D			1	NO MANUAL INTEGRATION
0647	02242358ECD7.D			1	NO MANUAL INTEGRATION
0708	02242359ECD7.D			1	NO MANUAL INTEGRATION
0729	02242360ECD7.D			1	NO MANUAL INTEGRATION
0750	02242361ECD7.D			1	NO MANUAL INTEGRATION
0811	02242362ECD7.D			1	NO MANUAL INTEGRATION
0832	02242363ECD7.D			1	NO MANUAL INTEGRATION
0853	02242364ECD7.D			1	NO MANUAL INTEGRATION
0914	02242365ECD7.D			1	NO MANUAL INTEGRATION
0935	02242366ECD7.D			1	NO MANUAL INTEGRATION
0956	02242367ECD7.D			1	NO MANUAL INTEGRATION
1017	02242368ECD7.D			1	NO MANUAL INTEGRATION
1038	02242369ECD7.D			1	NO MANUAL INTEGRATION
1059	02242370ECD7.D			1	NO MANUAL INTEGRATION
1120	02242371ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1141	02242372ECD7.D			1	NO MANUAL INTEGRATION
1202	02242373ECD7.D			1	NO MANUAL INTEGRATION
1223	02242374ECD7.D			1	NO MANUAL INTEGRATION
1244	02242375ECD7.D			1	NO MANUAL INTEGRATION
1305	02242376ECD7.D			1	NO MANUAL INTEGRATION
1326	02242377ECD7.D			1	NO MANUAL INTEGRATION
1347	02242378ECD7.D			1	NO MANUAL INTEGRATION
1408	02242379ECD7.D			1	NO MANUAL INTEGRATION
1429	02242380ECD7.D			1	NO MANUAL INTEGRATION
1450	02242381ECD7.D			1	NO MANUAL INTEGRATION
1511	02242382ECD7.D			1	NO MANUAL INTEGRATION
1532	02242383ECD7.D			1	NO MANUAL INTEGRATION
1553	02242384ECD7.D			1	NO MANUAL INTEGRATION
1051	02242301ECD7.D IB			1	NO MANUAL INTEGRATION
1112	02242302ECD7.D 0.25PPMAR1660			1	NO MANUAL INTEGRATION
1133	02242303ECD7.D 0.02PPMAR1660			1	Aroclor-1016 [2C],
1154	02242304ECD7.D 0.05PPMAR1660			1	NO MANUAL INTEGRATION
1215	02242305ECD7.D 1.0PPMAR1660			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1236	02242306ECD7.D	0.1PPMAR1660		1	NO MANUAL INTEGRATION
1257	02242307ECD7.D	0.5PPMAR1660		1	NO MANUAL INTEGRATION
1318	02242308ECD7.D	0.25PPMAR1242		1	NO MANUAL INTEGRATION
1339	02242309ECD7.D	0.25PPMAR1248		1	NO MANUAL INTEGRATION
1400	02242310ECD7.D	0.25PPMAR1254		1	NO MANUAL INTEGRATION
1421	02242311ECD7.D	0.25PPMAR2162		1	NO MANUAL INTEGRATION
1442	02242312ECD7.D	0.25PPMAR3268		1	NO MANUAL INTEGRATION
1503	02242313ECD7.D	AR1660SCV		1	NO MANUAL INTEGRATION
1524	02242314ECD7.D	AR1242SCV		1	NO MANUAL INTEGRATION
1545	02242315ECD7.D	AR1248SCV		1	NO MANUAL INTEGRATION
1606	02242316ECD7.D	AR1254SCV		1	NO MANUAL INTEGRATION
1627	02242317ECD7.D	AR2162SCV		1	NO MANUAL INTEGRATION
1648	02242318ECD7.D	AR3268SCV		1	NO MANUAL INTEGRATION
1709	02242319ECD7.D	DDTS		1	NO MANUAL INTEGRATION
1730	02242320ECD7.D	DDT BD		1	NO MANUAL INTEGRATION
1751	02242321ECD7.D			1	NO MANUAL INTEGRATION
1812	02242322ECD7.D			1	NO MANUAL INTEGRATION
1833	02242323ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1854	02242324ECD7.D			1	NO MANUAL INTEGRATION
1915	02242325ECD7.D			1	NO MANUAL INTEGRATION
1936	02242326ECD7.D			1	NO MANUAL INTEGRATION
1957	02242327ECD7.D			1	NO MANUAL INTEGRATION
2018	02242328ECD7.D			1	NO MANUAL INTEGRATION
2038	02242329ECD7.D			1	NO MANUAL INTEGRATION
2059	02242330ECD7.D			1	NO MANUAL INTEGRATION
2120	02242331ECD7.D			1	NO MANUAL INTEGRATION
2141	02242332ECD7.D			1	NO MANUAL INTEGRATION
2202	02242333ECD7.D			1	NO MANUAL INTEGRATION
2223	02242334ECD7.D			1	NO MANUAL INTEGRATION
2244	02242335ECD7.D			1	NO MANUAL INTEGRATION
2305	02242336ECD7.D			1	NO MANUAL INTEGRATION
2326	02242337ECD7.D			1	NO MANUAL INTEGRATION
2347	02242338ECD7.D			1	NO MANUAL INTEGRATION
0008	02242339ECD7.D			1	NO MANUAL INTEGRATION
0029	02242340ECD7.D			1	NO MANUAL INTEGRATION
0050	02242341ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0111	02242342ECD7.D			1	NO MANUAL INTEGRATION
0132	02242343ECD7.D			1	NO MANUAL INTEGRATION
0153	02242344ECD7.D			1	NO MANUAL INTEGRATION
0214	02242345ECD7.D			1	NO MANUAL INTEGRATION
0235	02242346ECD7.D			1	NO MANUAL INTEGRATION
0256	02242347ECD7.D			1	NO MANUAL INTEGRATION
0317	02242348ECD7.D			1	NO MANUAL INTEGRATION
0338	02242349ECD7.D			1	NO MANUAL INTEGRATION
0359	02242350ECD7.D			1	NO MANUAL INTEGRATION
0420	02242351ECD7.D			1	NO MANUAL INTEGRATION
0441	02242352ECD7.D			1	NO MANUAL INTEGRATION
0502	02242353ECD7.D			1	NO MANUAL INTEGRATION
0523	02242354ECD7.D			1	NO MANUAL INTEGRATION
0544	02242355ECD7.D			1	NO MANUAL INTEGRATION
0605	02242356ECD7.D			1	NO MANUAL INTEGRATION
0626	02242357ECD7.D			1	NO MANUAL INTEGRATION
0647	02242358ECD7.D			1	NO MANUAL INTEGRATION
0708	02242359ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0729	02242360ECD7.D			1	NO MANUAL INTEGRATION
0750	02242361ECD7.D			1	NO MANUAL INTEGRATION
0811	02242362ECD7.D			1	NO MANUAL INTEGRATION
0832	02242363ECD7.D			1	NO MANUAL INTEGRATION
0853	02242364ECD7.D			1	NO MANUAL INTEGRATION
0914	02242365ECD7.D			1	NO MANUAL INTEGRATION
0935	02242366ECD7.D			1	NO MANUAL INTEGRATION
0956	02242367ECD7.D			1	NO MANUAL INTEGRATION
1017	02242368ECD7.D			1	NO MANUAL INTEGRATION
1038	02242369ECD7.D			1	NO MANUAL INTEGRATION
1059	02242370ECD7.D			1	NO MANUAL INTEGRATION
1120	02242371ECD7.D			1	NO MANUAL INTEGRATION
1141	02242372ECD7.D			1	NO MANUAL INTEGRATION
1202	02242373ECD7.D			1	NO MANUAL INTEGRATION
1223	02242374ECD7.D			1	NO MANUAL INTEGRATION
1244	02242375ECD7.D			1	NO MANUAL INTEGRATION
1305	02242376ECD7.D			1	NO MANUAL INTEGRATION
1326	02242377ECD7.D			1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230224.b\230224.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1347	02242378ECD7.D			1	NO MANUAL INTEGRATION
1408	02242379ECD7.D			1	NO MANUAL INTEGRATION
1429	02242380ECD7.D			1	NO MANUAL INTEGRATION
1450	02242381ECD7.D			1	NO MANUAL INTEGRATION
1511	02242382ECD7.D			1	NO MANUAL INTEGRATION
1532	02242383ECD7.D			1	NO MANUAL INTEGRATION
1553	02242384ECD7.D			1	NO MANUAL INTEGRATION

Security Status Report

Date: 28-Feb-2023 10:53

02242301ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242302ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242303ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242304ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242305ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242306ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242307ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242308ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242309ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242310ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242311ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242312ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242313ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242314ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242315ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242316ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242317ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242318ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242319ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53
02242320ECD7.D	Data Locked	richardl, 28-Feb-2023 10:53

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Calibration File Names:

Level 1: \\target\share\chem4\ecd7.i\230224.b\02242303ECD7.D
 Level 2: \\target\share\chem4\ecd7.i\230224.b\02242304ECD7.D
 Level 3: \\target\share\chem4\ecd7.i\230224.b\02242306ECD7.D
 Level 4: \\target\share\chem4\ecd7.i\230224.b\02242302ECD7.D
 Level 5: \\target\share\chem4\ecd7.i\230224.b\02242307ECD7.D
 Level 6: \\target\share\chem4\ecd7.i\230224.b\02242305ECD7.D
 Level 7: \\target\share\chem4\ecd7.i\230224.b\02242312ECD7.D

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
2 Aroclor-1221 (1)	+++++ 0.00716	+++++	+++++	+++++	+++++	+++++	0.00716	0.000
(2)	+++++ 0.01281	+++++	+++++	+++++	+++++	+++++	0.01281	0.000
(3)	+++++ 0.02975	+++++	+++++	+++++	+++++	+++++	0.02975	0.000
3 Aroclor-1242 (1)	+++++ 0.02479	+++++	+++++	+++++	+++++	+++++	0.02479	0.000
(2)	+++++ 0.07529	+++++	+++++	+++++	+++++	+++++	0.07529	0.000
(3)	+++++ 0.02343	+++++	+++++	+++++	+++++	+++++	0.02343	0.000
(4)	+++++ 0.03463	+++++	+++++	+++++	+++++	+++++	0.03463	0.000
4 Aroclor-1232 (1)	+++++ 0.00429	+++++	+++++	+++++	+++++	+++++	0.00429	0.000

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(2)	++++ 0.00850	++++	++++	++++	++++	++++	0.00850	0.000
(3)	++++ 0.03848	++++	++++	++++	++++	++++	0.03848	0.000
(4)	++++ 0.01635	++++	++++	++++	++++	++++	0.01635	0.000
7 Aroclor-1016(1)	0.03172 ++++	0.03253	0.03142	0.03141	0.02856	0.02667	0.03039	7.449
(2)	0.09239 ++++	0.09246	0.09222	0.09849	0.09174	0.08849	0.09263	3.499
(3)	0.05165 ++++	0.05037	0.04823	0.04393	0.03991	0.03721	0.04522	12.936
(4)	0.03002 ++++	0.02894	0.02959	0.03058	0.02852	0.02774	0.02923	3.542
6 Aroclor-1248(1)	++++ 0.03903	++++	++++	++++	++++	++++	0.03903	0.000
(2)	++++ 0.04961	++++	++++	++++	++++	++++	0.04961	0.000
(3)	++++ 0.09360	++++	++++	++++	++++	++++	0.09360	0.000
(4)	++++ 0.04765	++++	++++	++++	++++	++++	0.04765	0.000

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
8 Aroclor-1254 (1)	++++ 0.08033	++++	++++	++++	++++	++++	0.08033	0.000
(2)	++++ 0.03613	++++	++++	++++	++++	++++	0.03613	0.000
(3)	++++ 0.05165	++++	++++	++++	++++	++++	0.05165	0.000
(4)	++++ 0.10042	++++	++++	++++	++++	++++	0.10042	0.000
(5)	++++ 0.06294	++++	++++	++++	++++	++++	0.06294	0.000
9 Aroclor-1260 (1)	0.02926 ++++	0.02920	0.02841	0.03096	0.02737	0.02746	0.02878	4.677
(2)	0.02967 ++++	0.03006	0.03011	0.03291	0.02910	0.02857	0.03007	5.029
(3)	0.08088 ++++	0.08045	0.07954	0.08575	0.07515	0.07674	0.07975	4.627
(4)	0.03905 ++++	0.03887	0.03955	0.04485	0.03942	0.03922	0.04016	5.753
(5)	0.01783 ++++	0.01715	0.01679	0.01875	0.01664	0.01655	0.01729	4.953
10 Aroclor-1262 (1)	++++ 0.02454	++++	++++	++++	++++	++++	0.02454	0.000

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(2)	++++ 0.03993	++++	++++	++++	++++	++++	0.03993	0.000
(3)	++++ 0.04293	++++	++++	++++	++++	++++	0.04293	0.000
(4)	++++ 0.03923	++++	++++	++++	++++	++++	0.03923	0.000
11 Aroclor-1268(1)	++++ 0.10250	++++	++++	++++	++++	++++	0.10250	0.000
(2)	++++ 0.10151	++++	++++	++++	++++	++++	0.10151	0.000
(3)	++++ 0.08686	++++	++++	++++	++++	++++	0.08686	0.000
(4)	++++ 0.28598	++++	++++	++++	++++	++++	0.28598	0.000
42 2,4-DDE	++++ ++++	++++	++++	++++	++++	++++	++++	++++
43 2,4-DDD	++++ ++++	++++	++++	++++	++++	++++	++++	++++
44 2,4-DDT	++++ ++++	++++	++++	++++	++++	++++	++++	++++
46 4,4-DDE	++++ ++++	++++	++++	++++	++++	++++	++++	++++

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Last Edit : 24-Feb-2023 15:31 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
47 4,4-DDD	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
48 4,4-DDT	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
49 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
50 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
1 Tetrachloro-m-xylene	1.16827	1.24402	1.18546	1.20509	1.12295	1.24114	1.19449	3.860
13 Decachlorobiphenyl	0.82901	0.80558	0.77587	0.78808	0.73125	0.79742	0.78787	4.189

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Calibration File Names:

Level 1: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242303ECD7.D
 Level 2: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242304ECD7.D
 Level 3: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242306ECD7.D
 Level 4: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242302ECD7.D
 Level 5: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242307ECD7.D
 Level 6: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242305ECD7.D
 Level 7: \\target\share\chem4\ecd7.i\230224.b\230224.b\02242312ECD7.D

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
1 Aroclor-1221 [2C] (1)	+++++ 0.00757	+++++	+++++	+++++	+++++	+++++	0.00757	0.000
(2)	+++++ 0.01433	+++++	+++++	+++++	+++++	+++++	0.01433	0.000
(3)	+++++ 0.02333	+++++	+++++	+++++	+++++	+++++	0.02333	0.000
4 Aroclor-1232 [2C] (1)	+++++ 0.00409	+++++	+++++	+++++	+++++	+++++	0.00409	0.000
(2)	+++++ 0.02034	+++++	+++++	+++++	+++++	+++++	0.02034	0.000
(3)	+++++ 0.04067	+++++	+++++	+++++	+++++	+++++	0.04067	0.000
(4)	+++++ 0.01170	+++++	+++++	+++++	+++++	+++++	0.01170	0.000
3 Aroclor-1242 [2C] (1)	+++++ 0.03717	+++++	+++++	+++++	+++++	+++++	0.03717	0.000

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
(2)	+++++	+++++	+++++	+++++	+++++	+++++	0.07813	0.000
(3)	+++++	+++++	+++++	+++++	+++++	+++++	0.02431	0.000
(4)	+++++	+++++	+++++	+++++	+++++	+++++	0.02962	0.000
6 Aroclor-1248 [2C] (1)	+++++	+++++	+++++	+++++	+++++	+++++	0.03820	0.000
(2)	+++++	+++++	+++++	+++++	+++++	+++++	0.03949	0.000
(3)	+++++	+++++	+++++	+++++	+++++	+++++	0.04545	0.000
(4)	+++++	+++++	+++++	+++++	+++++	+++++	0.05457	0.000
7 Aroclor-1016 [2C] (1)	0.05071	0.05022	0.04868	0.04733	0.04326	0.04080	0.04683	8.503
(2)	0.08143	0.09407	0.10159	0.10259	0.09651	0.09362	0.09497	8.025
(3)	0.04006	0.04718	0.04613	0.04410	0.04062	0.03926	0.04289	7.857
(4)	0.03181	0.03802	0.03707	0.03450	0.03115	0.02936	0.03365	10.251

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
8 Aroclor-1254 [2C] (1)	+++++ 0.06081	+++++	+++++	+++++	+++++	+++++	0.06081	0.000
(2)	+++++ 0.04892	+++++	+++++	+++++	+++++	+++++	0.04892	0.000
(3)	+++++ 0.10584	+++++	+++++	+++++	+++++	+++++	0.10584	0.000
(4)	+++++ 0.10317	+++++	+++++	+++++	+++++	+++++	0.10317	0.000
(5)	+++++ 0.06282	+++++	+++++	+++++	+++++	+++++	0.06282	0.000
10 Aroclor-1262 [2C] (1)	+++++ 0.06831	+++++	+++++	+++++	+++++	+++++	0.06831	0.000
(2)	+++++ 0.05818	+++++	+++++	+++++	+++++	+++++	0.05818	0.000
(3)	+++++ 0.06601	+++++	+++++	+++++	+++++	+++++	0.06601	0.000
(4)	+++++ 0.10341	+++++	+++++	+++++	+++++	+++++	0.10341	0.000
9 Aroclor-1260 [2C] (1)	0.05286 +++++	0.04911	0.04696	0.04801	0.04329	0.04201	0.04704	8.422
(2)	0.12976 +++++	0.12431	0.12095	0.12664	0.11320	0.10545	0.12005	7.605

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	2500.000 Level 7	RRF	% RSD
(3)	0.03524	0.03147	0.02937	0.03208	0.03102	0.03198		0.03186	6.045
(4)	0.08632	0.08237	0.08044	0.08393	0.07718	0.07531		0.08092	5.126
11 Aroclor-1268 [2C] (1)	0.16109							0.16109	0.000
(2)	0.17318							0.17318	0.000
(3)	0.14787							0.14787	0.000
(4)	0.47260							0.47260	0.000
41 2,4-DDE [2C]									
42 2,4-DDD [2C]									
44 4,4-DDE [2C]									
45 4,4-DDD/2,4-DDT [2C]									
46 4,4-DDT [2C]									

ARI Labs, Inc.

INITIAL CALIBRATION DATA

Start Cal Date : 24-FEB-2023 11:12
 End Cal Date : 24-FEB-2023 14:42
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.14
 Integrator : HP Genie
 Method file : \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Last Edit : 24-Feb-2023 15:29 richardl
 Curve Type : Average

Compound	20.000 Level 1	50.000 Level 2	100.000 Level 3	250.000 Level 4	500.000 Level 5	1000.000 Level 6	RRF	% RSD
48 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
49 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++
\$ 2 Tetrachloro-m-xylene [2C]	1.21526 +++++	1.19545	1.17555	1.21907	1.12560	1.11139	1.17372	3.897
\$ 13 Decachlorobiphenyl [2C]	1.17066 +++++	1.20406	1.20549	1.31040	1.21104	1.20797	1.21827	3.898

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Batch File: \\target\share\chem4\ecd7.i\230224.b
Inst ID: ecd7.i

ID: RT01 RT02 RT03 RT04 RT05 RT06
FILENAME: 02242302ECD7 02242303ECD7 02242304ECD7 02242305ECD7 02242306ECD7 02242307ECD7
INJ. DATE: 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023
INJ. TIME: 11:12 11:33 11:54 12:15 12:36 12:57

Table with 11 columns: Compound, RT01, RT02, RT03, RT04, RT05, RT06, EXPEC RT, RT WINDOW, AVG RT, STD DEV. Rows include various chemical compounds like IS-BNB, Tetrachloro-m-xylene, Aroclor-1221, etc.

Reviewer 1 _____ Date: _____
Reviewer 2 _____ Date: _____

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Batch File: \\target\share\chem4\ecd7.i\230224.b
 Inst ID: ecd7.i

Compound	RT01	RT02	RT03	RT04	RT05	RT06	EXPEC RT	RT WINDOW	AVG RT	STD DEV
47 4,4-DDD	+++++	+++++	+++++	+++++	+++++	+++++	10.254	10.154-10.354	+++++	+++++
48 4,4-DDT	+++++	+++++	+++++	+++++	+++++	+++++	10.754	10.654-10.854	+++++	+++++
49 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	1.842	1.742-1.942	+++++	+++++
50 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	6.708	6.608-6.808	+++++	+++++

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
Batch File: \\target\share\chem4\ecd7.i\230224.b\230224.b
Inst ID: ecd7.i

ID: RT01 RT02 RT03 RT04 RT05 RT06
FILENAME: 02242302ECD7 02242303ECD7 02242304ECD7 02242305ECD7 02242306ECD7 02242307ECD7
INJ. DATE: 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023 24-FEB-2023
INJ. TIME: 11:12 11:33 11:54 12:15 12:36 12:57

Table with 11 columns: Compound, RT01, RT02, RT03, RT04, RT05, RT06, EXPEC RT, RT WINDOW, AVG RT, STD DEV. Rows include various chemical compounds like Aroclor-1221, Aroclor-1232, etc., with their respective retention times and standard deviations.

Reviewer 1 _____ Date: _____
Reviewer 2 _____ Date: _____

ARI Labs, Inc.
RETENTION TIME SUMMARY REPORT

Method File: \\target\share\chem4\ecd7.i\230224.b\PCB.m\PCB2.m
 Batch File: \\target\share\chem4\ecd7.i\230224.b\230224.b
 Inst ID: ecd7.i

Compound	RT01	RT02	RT03	RT04	RT05	RT06	EXPEC RT	RT WINDOW	AVG RT	STD DEV
46 4,4-DDT [2C]	+++++	+++++	+++++	+++++	+++++	+++++	11.092	10.992-11.192	+++++	+++++
48 Hexachlorobutadiene	+++++	+++++	+++++	+++++	+++++	+++++	1.703	1.603-1.803	+++++	+++++
49 Hexachlorobenzene	+++++	+++++	+++++	+++++	+++++	+++++	7.178	7.078-7.278	+++++	+++++

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242301ECD7.D
Data file 2: /230224.b/230224.b/02242301ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: IB
Client ID:
Injection Date: 24-FEB-2023 10:51
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.826	0.019	382217	5.683	-0.002	180378	33.8	36.5	7.7	Tetrachloro-m-xylene
13.904	0.011	534110	14.120	0.001	295605	35.3	37.2	5.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	756896	12.3
Hexabromobiphenyl	1429847	1534275	7.3

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	336543	6.8
Hexabromobiphenyl	513946	521508	1.5

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	---			0.0	1	---			0.0
Aroclor-1016	2	---			0.0	2	---			0.0
Aroclor-1016	3	---			0.0	3	---			0.0
Aroclor-1016	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	---			0.0	2	6.321	0.025	1873	31.1
Aroclor-1221	3	---			0.0	3	6.633	0.012	314	3.2
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	---			0.0	2	---			0.0
Aroclor-1232	3	7.698	0.043	2193	6.0	3	---			0.0
Aroclor-1232	4	8.505	-0.076	11525	74.5	4	---			0.0
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1242	1	---			0.0	1	---			0.0
Aroclor-1242	2	7.698	0.042	2193	3.1	2	---			0.0
Aroclor-1242	3	---			0.0	3	---			0.0
Aroclor-1242	4	8.505	-0.074	11525	35.2	4	---			0.0
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1248	1	---			0.0	1	---			0.0
Aroclor-1248	2	---			0.0	2	---			0.0
Aroclor-1248	3	---			0.0	3	---			0.0
Aroclor-1248	4	---			0.0	4	---			0.0
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1254	1	---			0.0	1	---			0.0
Aroclor-1254	2	---			0.0	2	---			0.0
Aroclor-1254	3	9.596	-0.072	31424	64.3	3	---			0.0
Aroclor-1254	4	---			0.0	4	---			0.0
Aroclor-1254	5	10.167	-0.010	18361	30.8	5	---			0.0
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1260	1	11.098	0.054	6994	12.7	1	---			0.0
Aroclor-1260	2	---			0.0	2	---			0.0
Aroclor-1260	3	11.706	-0.027	7806	5.1	3	---			0.0
Aroclor-1260	4	---			0.0	4	---			0.0
Aroclor-1260	5	---			0.0	NS	---			----
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1262	1	10.824	-0.005	16873	35.8	1	---			0.0
Aroclor-1262	2	---			0.0	2	---			0.0
Aroclor-1262	3	---			0.0	3	---			0.0
Aroclor-1262	4	13.040	0.053	14031	18.6	4	---			0.0
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					
Aroclor-1268	1	---			0.0	1	---			0.0
Aroclor-1268	2	---			0.0	2	---			0.0
Aroclor-1268	3	12.709	0.009	6037	3.6	3	12.891	-0.001	659	0.7
Aroclor-1268	4	13.499	0.010	12396	2.3	4	13.710	0.001	1848	0.6
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks					

Total PCB Area Coll (5.906 - 13.793) = 260205

Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 18252 Col2 Total PCB = 0.0 ppm*

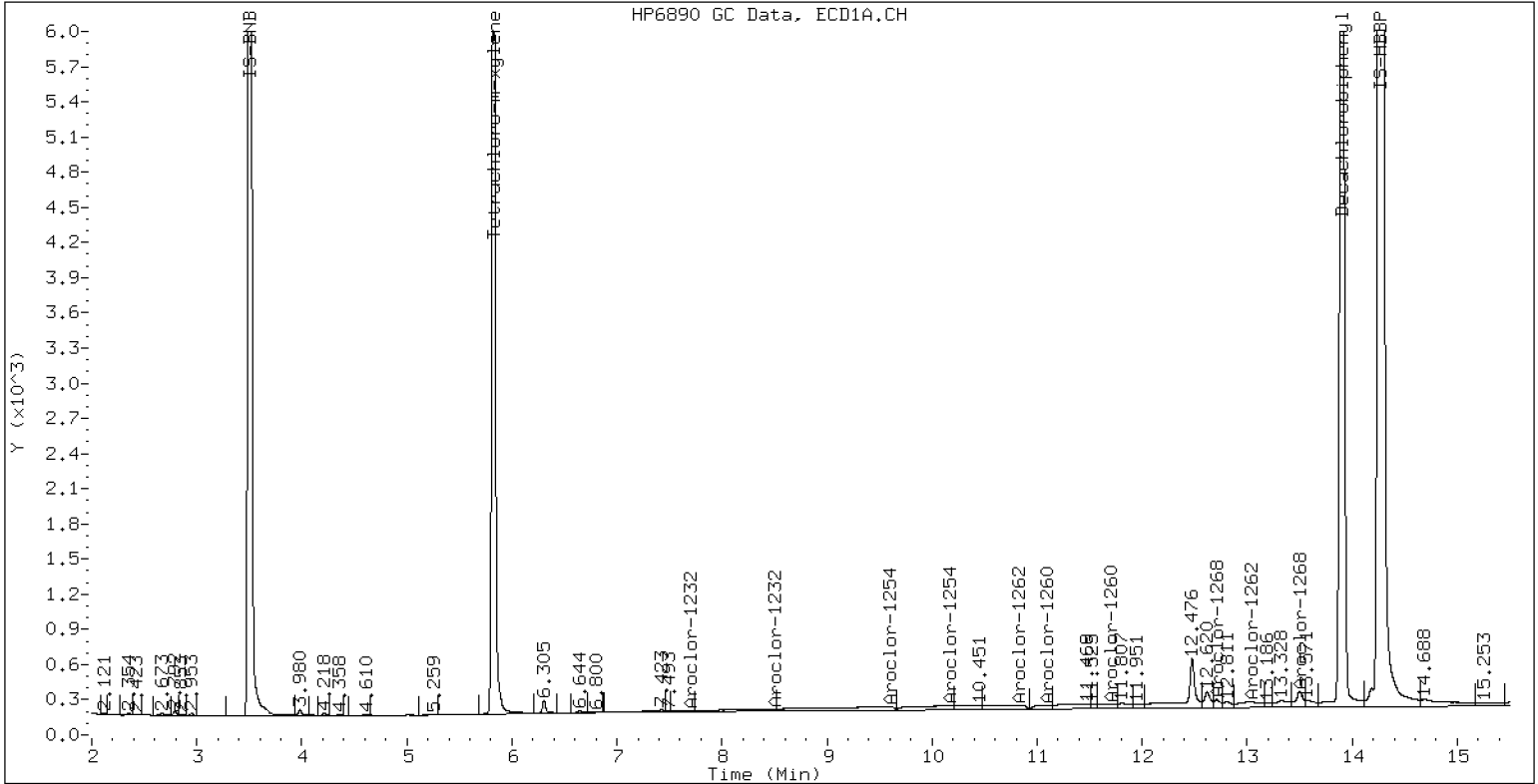
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 IB

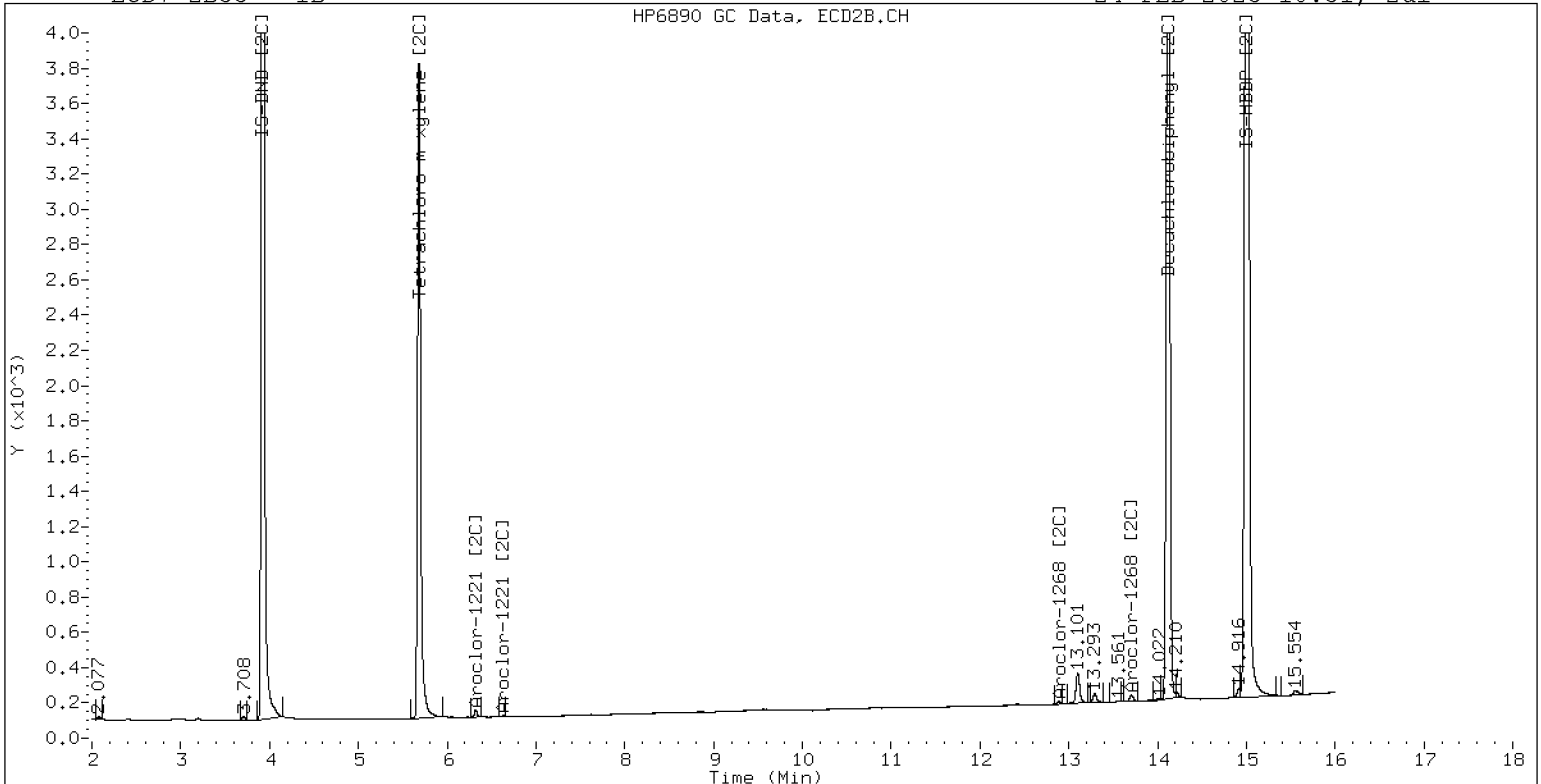
24-FEB-2023 10:51, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 IB

24-FEB-2023 10:51, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242302ECD7.D
 Data file 2: /230224.b/230224.b/02242302ECD7.D
 Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
 Compound Sublist: AR1660.sub
 Instrument, Inj. Vol.: ecd7.i, 2ul
 Quant Method: Internal Std

ARI ID: 0.25PPMAR1660
 Client ID:
 Injection Date: 24-FEB-2023 11:12
 Report Date: 02/28/2023 09:50
 Matrix: NONE
 Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	0.005	405980	5.687	0.002	192160	40.4	41.5	2.9	Tetrachloro-m-xylene
13.897	0.004	563414	14.120	0.001	336737	40.0	43.0	7.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	673778	0.0
Hexabromobiphenyl	1429847	1429847	0.0
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	315256	0.0
Hexabromobiphenyl	513946	513946	0.0

* Standard Areas taken from Initial Cal Level 3
 Initial Calibration Date: 24-FEB-2023
 <- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.272	0.001	66125	258.4	1	7.255	-0.001	46626	252.6
Aroclor-1016	2	7.654	-0.000	207370	265.8	2	7.855	-0.001	101071	270.1
Aroclor-1016	3	7.792	0.002	92507	242.9	3	8.055	0.001	43448	257.1
Aroclor-1016	4	8.406	0.001	64388	261.5	4	8.306	-0.000	33986	256.3
Total CollAve (4 peaks):				257.2		Total Col2Ave (4 peaks):				259.0 RPD = 1
Corrected Ave (3 peaks):				254.3		Corrected Ave (3 peaks):				255.3 RPD = 0
CalAmt %D:				2.9		CalAmt %D:				3.6
Aroclor-1260	1	11.046	0.001	138355	269.0	1	11.653	0.001	77114	255.2
Aroclor-1260	2	11.363	0.002	147051	273.6	2	11.918	0.001	203401	263.7
Aroclor-1260	3	11.736	0.003	383171	268.8	3	12.435	-0.000	51517	251.7
Aroclor-1260	4	12.141	0.002	200399	279.2	4	12.502	0.001	134797	259.3
Aroclor-1260	5	12.247	0.003	83796	271.2	NS	---			----
Total CollAve (5 peaks):				272.4		Total Col2Ave (4 peaks):				257.5 RPD = 6
Corrected Ave (4 peaks):				270.7		Corrected Ave (3 peaks):				255.4 RPD = 6
CalAmt %D:				8.9		CalAmt %D:				3.0

Total PCB Area Coll (5.906 - 13.793) = 4024419 Coll Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1889311 Col2 Total PCB = 0.5 ppm*

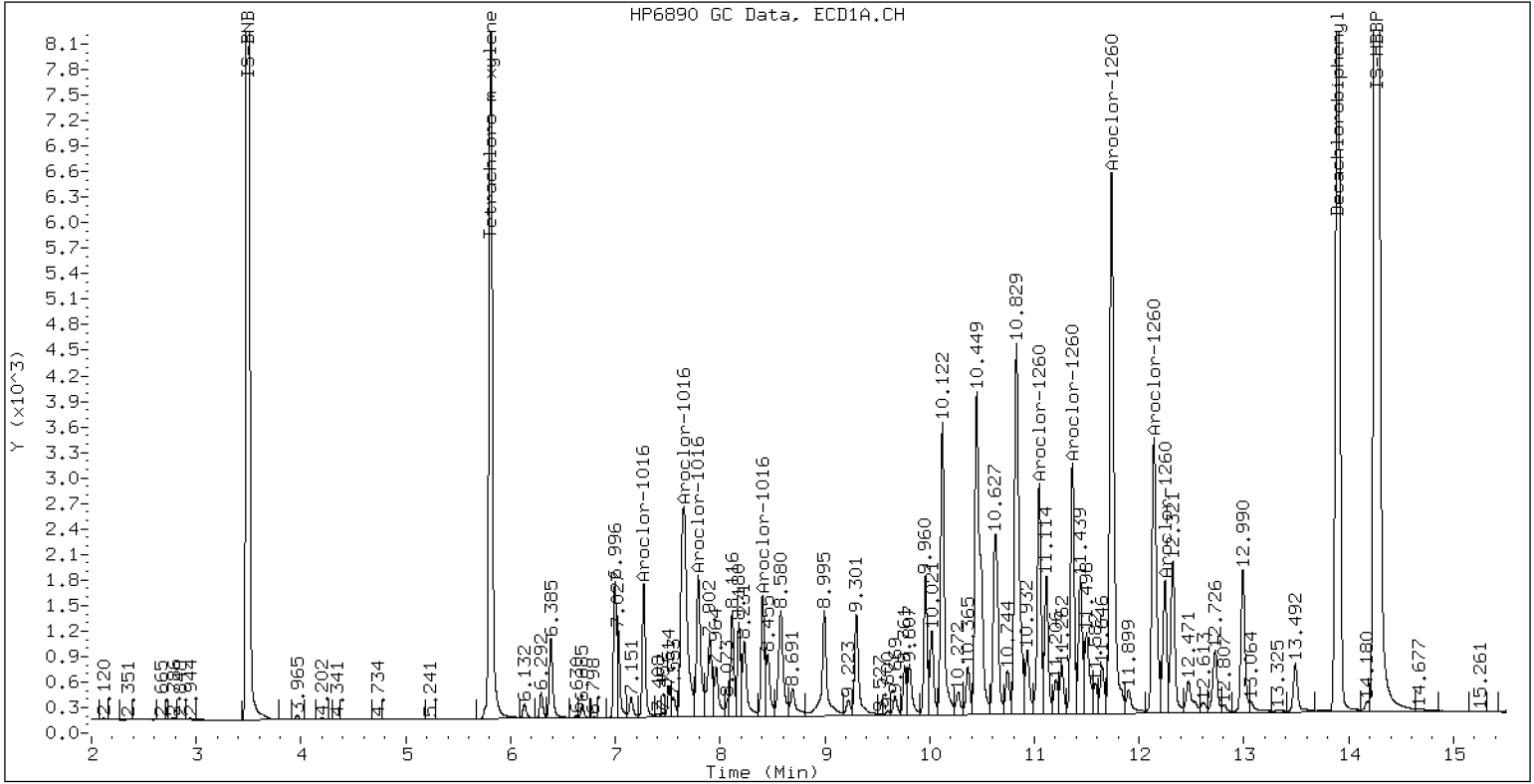
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1660

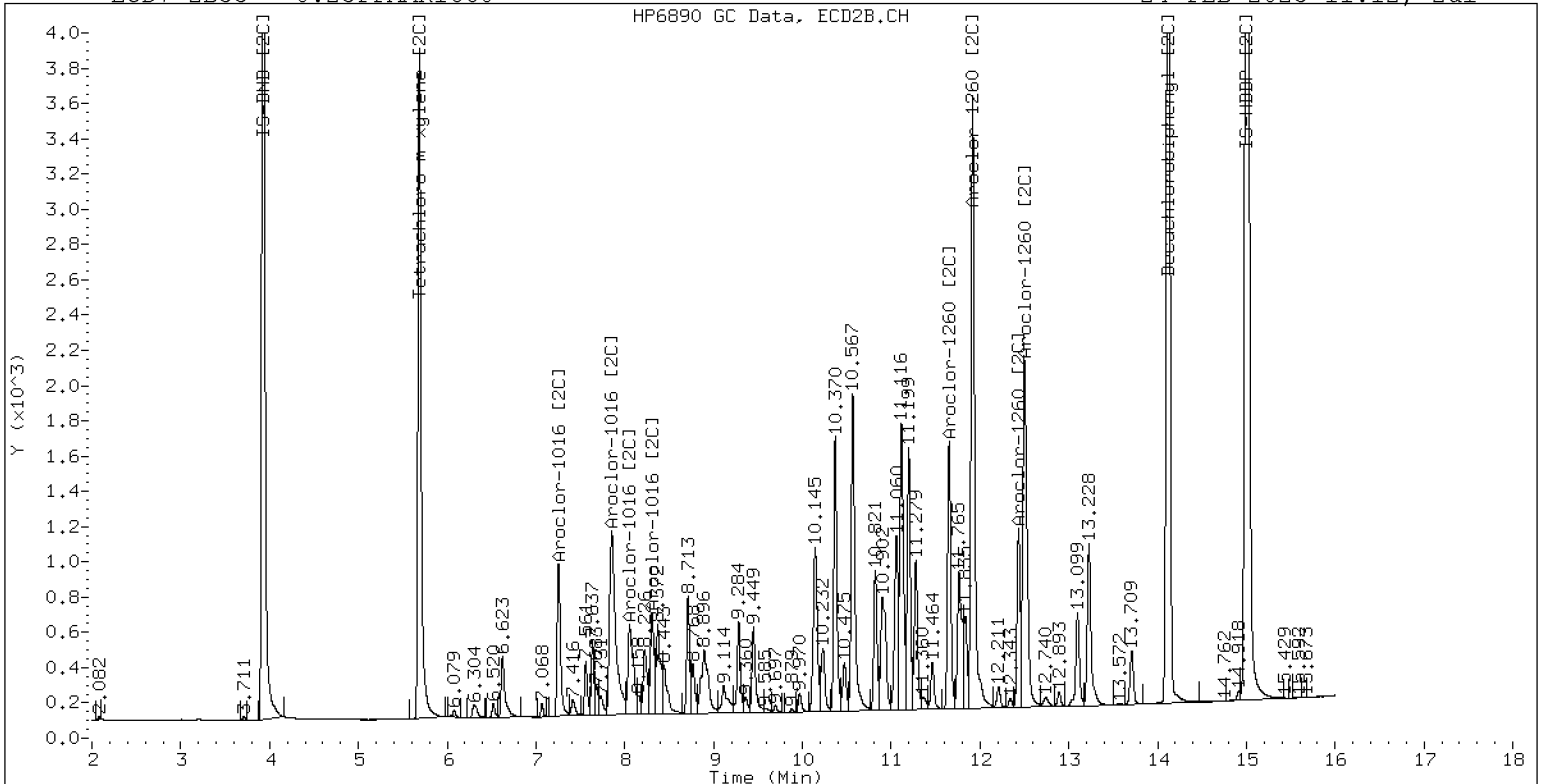
24-FEB-2023 11:12, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1660

24-FEB-2023 11:12, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242303ECD7.D
Data file 2: /230224.b/230224.b/02242303ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.02PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 11:33
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.002	29768	5.688	0.003	14932	3.1	3.3	5.7	Tetrachloro-m-xylene
13.893	0.000	45992	14.120	0.000	23950	3.4	3.1	9.1	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	637010	-5.5
Hexabromobiphenyl	1429847	1386953	-3.0

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	307177	-2.6
Hexabromobiphenyl	513946	511463	-0.5

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.272	0.001	5052	20.9	1	7.256	0.000	3894	21.7	
Aroclor-1016	2	7.659	0.005	14714	19.9	2	7.864	0.008	6253	17.1	
Aroclor-1016	3	7.795	0.005	8226	22.8	3	8.060	0.006	3076	18.7	
Aroclor-1016	4	8.407	0.002	4780	20.5	4	8.309	0.002	2443	18.9	
Total CollAve (4 peaks):				21.1	Total Col2Ave (4 peaks):				19.1	RPD = 10	
Corrected Ave (3 peaks):				20.5	Corrected Ave (3 peaks):				18.2	RPD = 11	
CalAmt %D:				5.3	CalAmt %D:				-4.5		
Aroclor-1260	1	11.047	0.003	10147	20.3	1	11.656	0.003	6759	22.5	
Aroclor-1260	2	11.364	0.003	10287	19.7	2	11.922	0.005	16592	21.6	
Aroclor-1260	3	11.740	0.006	28043	20.3	3	12.438	0.002	4506	22.1	
Aroclor-1260	4	12.145	0.006	13540	19.4	4	12.505	0.004	11037	21.3	
Aroclor-1260	5	12.246	0.002	6182	20.6	NS	---			----	
Total CollAve (5 peaks):				20.1	Total Col2Ave (4 peaks):				21.9	RPD = 9	
Corrected Ave (4 peaks):				19.9	Corrected Ave (3 peaks):				21.7	RPD = 8	
CalAmt %D:				0.4	CalAmt %D:				9.4		

Total PCB Area Coll (5.906 - 13.793) = 324832 Coll Total PCB = 0.0 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 157149 Col2 Total PCB = 0.0 ppm*

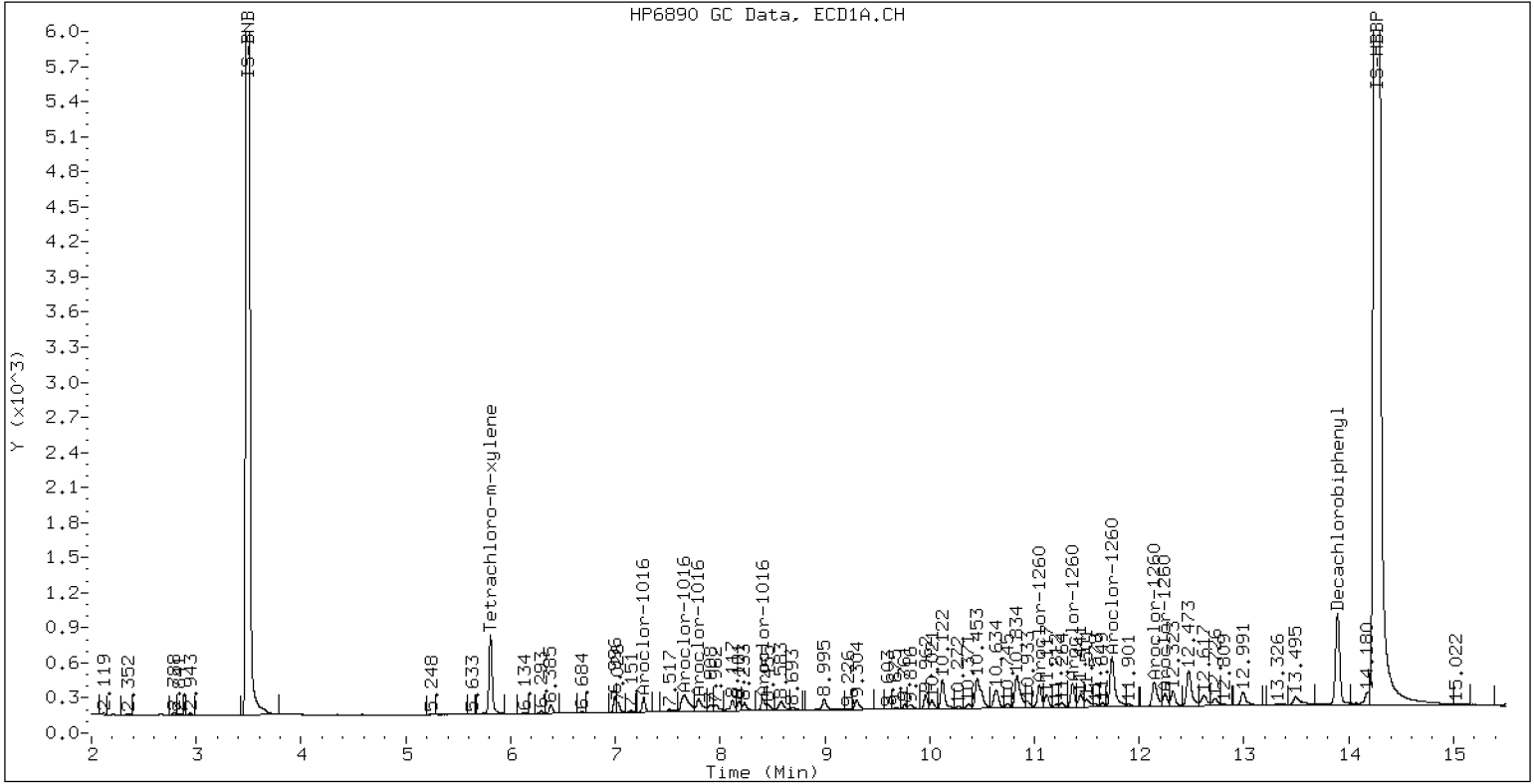
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.02PPMAR1660

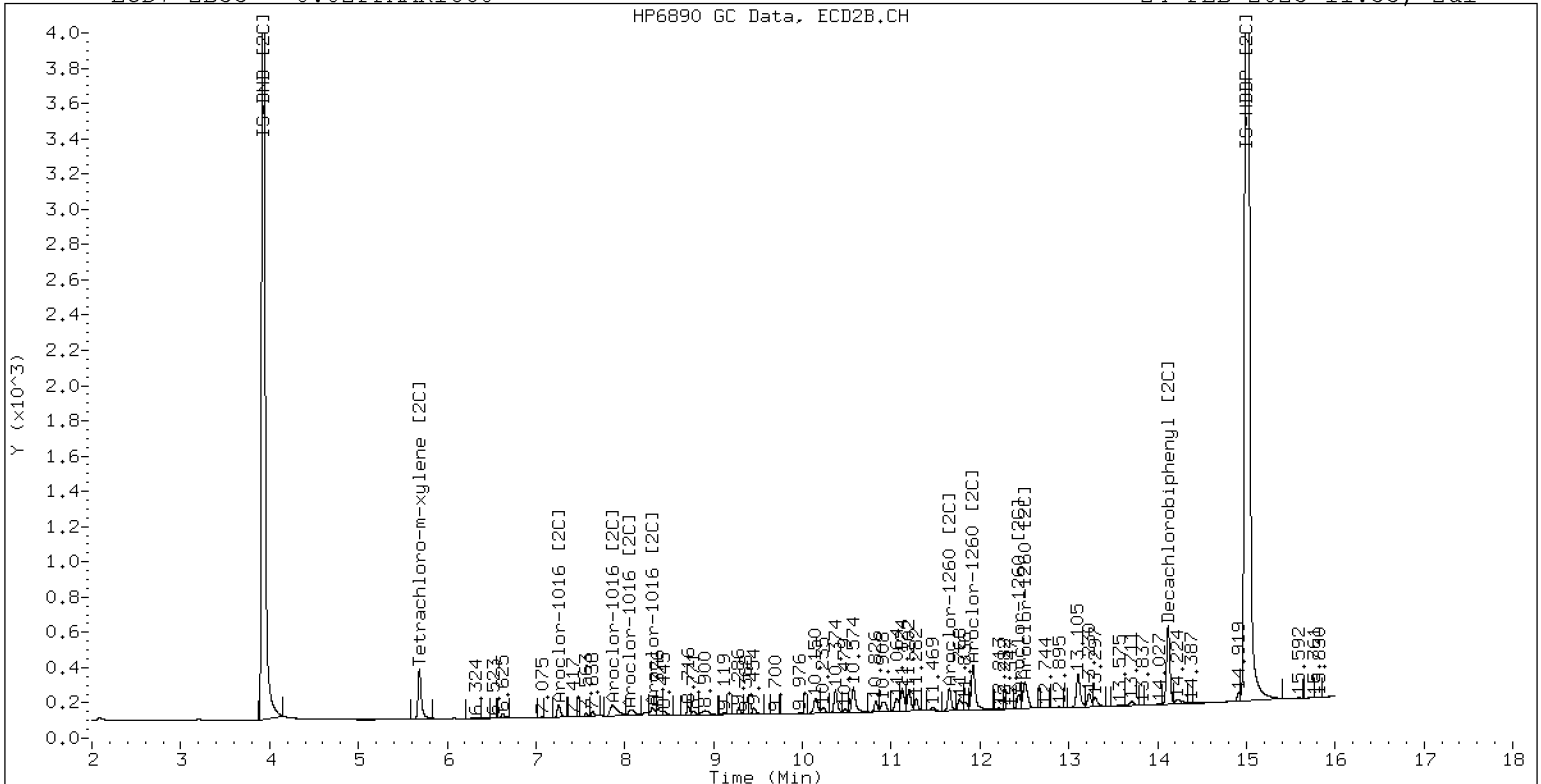
24-FEB-2023 11:33, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.02PPMAR1660

24-FEB-2023 11:33, 2ul

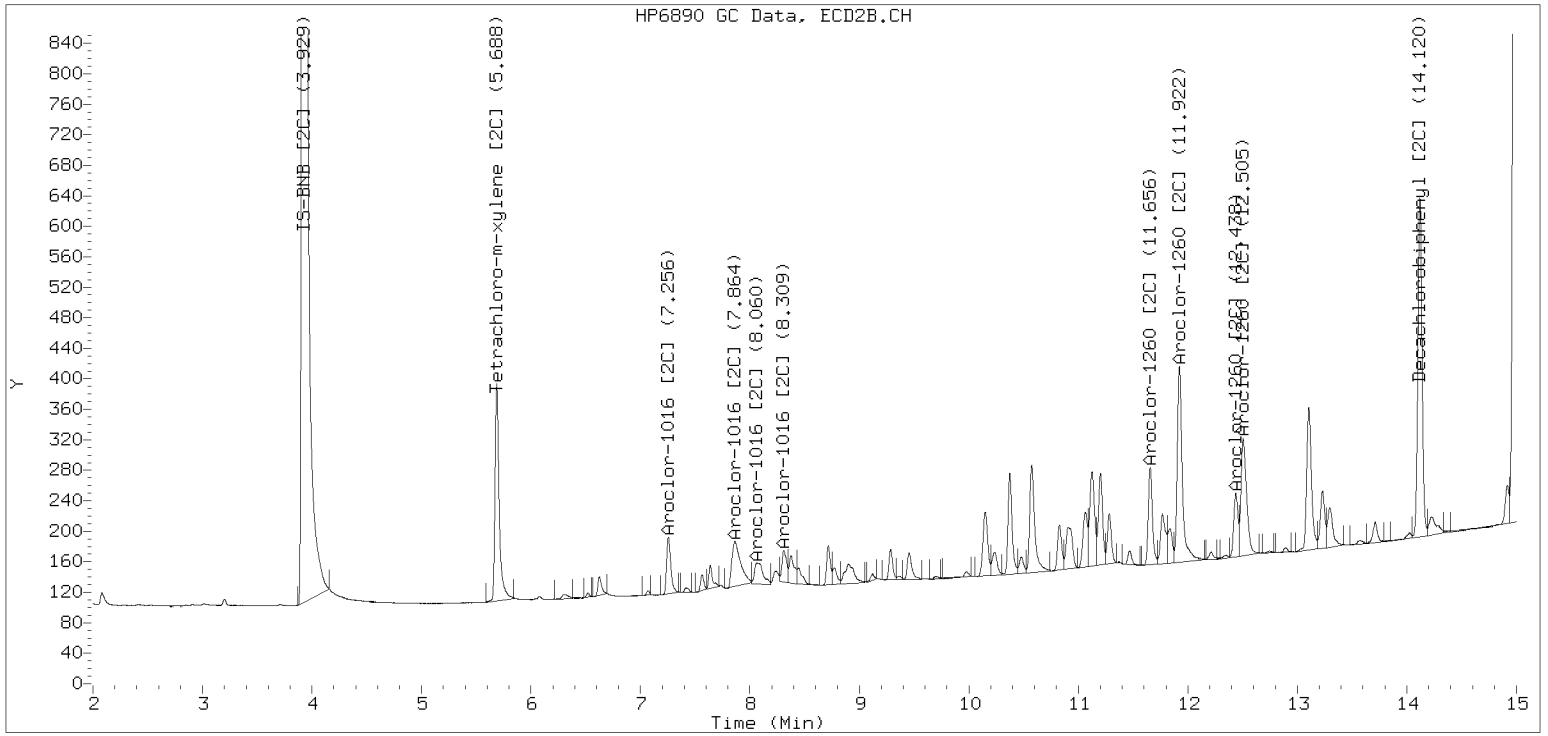


ZB-35 Manual Integration: YES

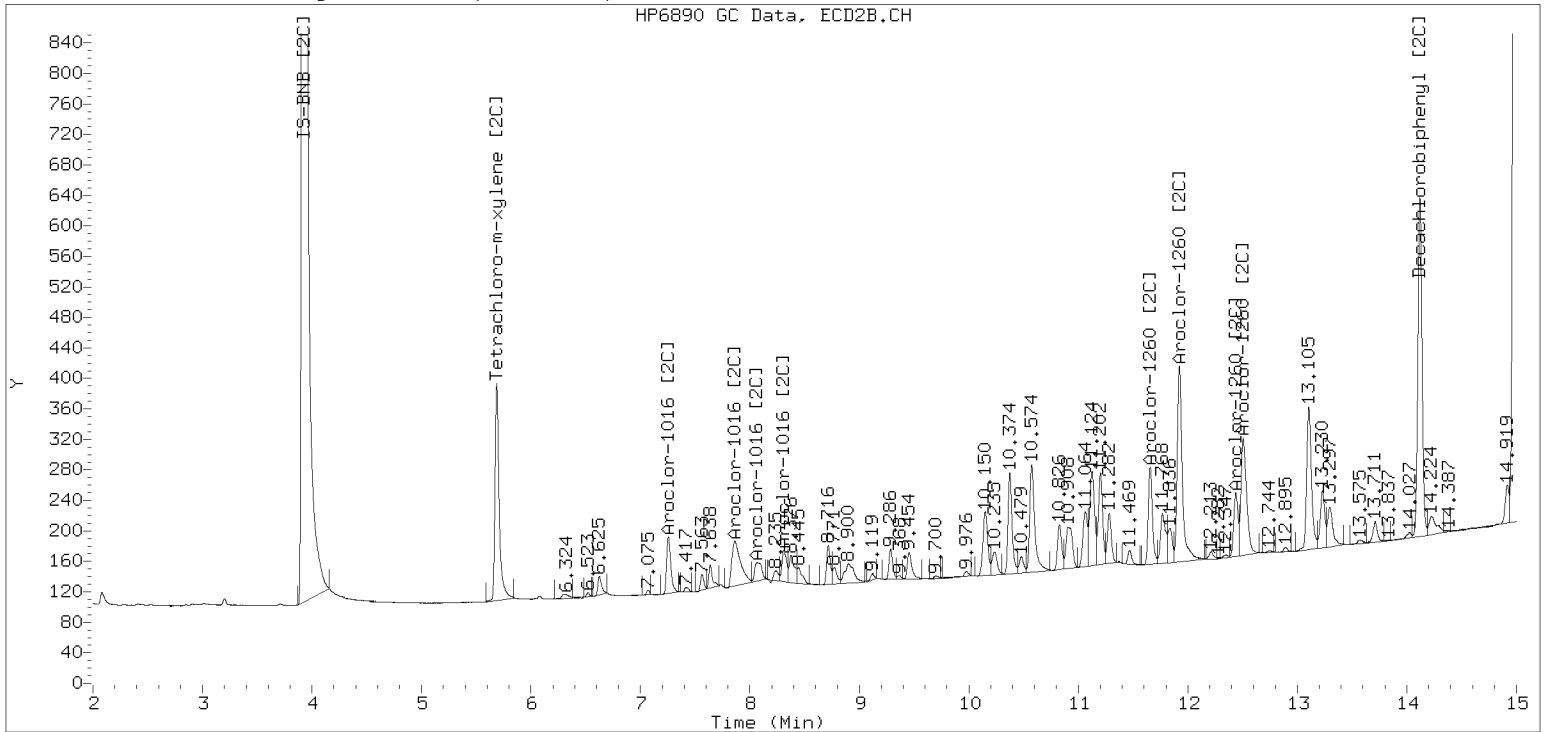
Manual Peak Adjustment, ZB-35

Datafile: ecd7.i/230224.b/230224.b/02242303ECD7.D Injection Date: 24-FEB-2023

Manual Integration (After)



Processed Integration (Before)



Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242304ECD7.D
Data file 2: /230224.b/230224.b/02242304ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.05PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 11:54
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col		ZB5	ZB35	RPD	Compound/Flag		
RT	Shift Response	RT	Shift Response	on col	on col				
5.809	0.003	78493	5.688	0.003	36772	8.3	8.1	2.2	Tetrachloro-m-xylene
13.893	-0.000	113544	14.119	-0.000	62745	8.2	7.9	3.4	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	630965	-6.4
Hexabromobiphenyl	1429847	1409464	-1.4

Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	307599	-2.4
Hexabromobiphenyl	513946	521112	1.4

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.272	0.002	12829	53.5	1	7.256	0.000	9654	53.6	
Aroclor-1016	2	7.660	0.006	36461	49.9	2	7.864	0.008	18085	49.5	
Aroclor-1016	3	7.795	0.005	19865	55.7	3	8.063	0.008	9071	55.0	
Aroclor-1016	4	8.408	0.003	11411	49.5	4	8.310	0.003	7309	56.5	
Total CollAve (4 peaks):				52.2	Total Col2Ave (4 peaks):				53.7	RPD = 3	
Corrected Ave (3 peaks):				51.0	Corrected Ave (3 peaks):				52.7	RPD = 3	
CalAmt %D:				4.3	CalAmt %D:				7.3		
Aroclor-1260	1	11.046	0.002	25727	50.7	1	11.655	0.002	15996	52.2	
Aroclor-1260	2	11.363	0.002	26482	50.0	2	11.922	0.004	40487	51.8	
Aroclor-1260	3	11.739	0.005	70871	50.4	3	12.437	0.002	10248	49.4	
Aroclor-1260	4	12.143	0.004	34239	48.4	4	12.506	0.004	26828	50.9	
Aroclor-1260	5	12.246	0.002	15109	49.6	NS	---			----	
Total CollAve (5 peaks):				49.8	Total Col2Ave (4 peaks):				51.1	RPD = 2	
Corrected Ave (4 peaks):				49.6	Corrected Ave (3 peaks):				50.7	RPD = 2	
CalAmt %D:				-0.3	CalAmt %D:				2.1		

Total PCB Area Coll (5.906 - 13.793) = 758292 Coll Total PCB = 0.1 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 386383 Col2 Total PCB = 0.1 ppm*

* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242305ECD7.D
Data file 2: /230224.b/230224.b/02242305ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 1.0PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 12:15
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.813	0.006	1641874	5.688	0.003	709674	166.2	151.5	9.3	Tetrachloro-m-xylene
13.899	0.006	2344583	14.122	0.002	1300114	161.9	158.6	2.1	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	661440	-1.8
Hexabromobiphenyl	1429847	1470100	2.8
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	319272	1.3
Hexabromobiphenyl	513946	538138	4.7

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	-0.000	220519	877.8	1	7.254	-0.001	162833	871.2	
Aroclor-1016	2	7.652	-0.002	731607	955.3	2	7.852	-0.004	373610	985.8	
Aroclor-1016	3	7.789	-0.001	307629	822.8	3	8.051	-0.003	156666	915.2	
Aroclor-1016	4	8.404	-0.001	229387	949.1	4	8.305	-0.002	117186	872.6	
Total CollAve (4 peaks):				901.3		Total Col2Ave (4 peaks):				911.2	RPD = 1
Corrected Ave (3 peaks):				883.3		Corrected Ave (3 peaks):				886.3	RPD = 0

CalAmt %D: -9.9

CalAmt %D: -8.9

Aroclor-1260	1	11.044	-0.000	504641	954.2	1	11.652	-0.000	282606	893.1	
Aroclor-1260	2	11.360	-0.001	524931	950.0	2	11.917	-0.000	709329	878.4	
Aroclor-1260	3	11.734	-0.000	1410270	962.3	3	12.434	-0.001	215124	1003.8	
Aroclor-1260	4	12.137	-0.002	720770	976.7	4	12.501	-0.001	506566	930.6	
Aroclor-1260	5	12.243	-0.001	304211	957.7	NS	---			----	
Total CollAve (5 peaks):				960.2		Total Col2Ave (4 peaks):				926.5	RPD = 4
Corrected Ave (4 peaks):				956.0		Corrected Ave (3 peaks):				900.7	RPD = 6

CalAmt %D: -4.0

CalAmt %D: -7.4

Total PCB Area Coll (5.906 - 13.793) = 14454279 Coll Total PCB = 1.8 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 7029563 Col2 Total PCB = 1.8 ppm*

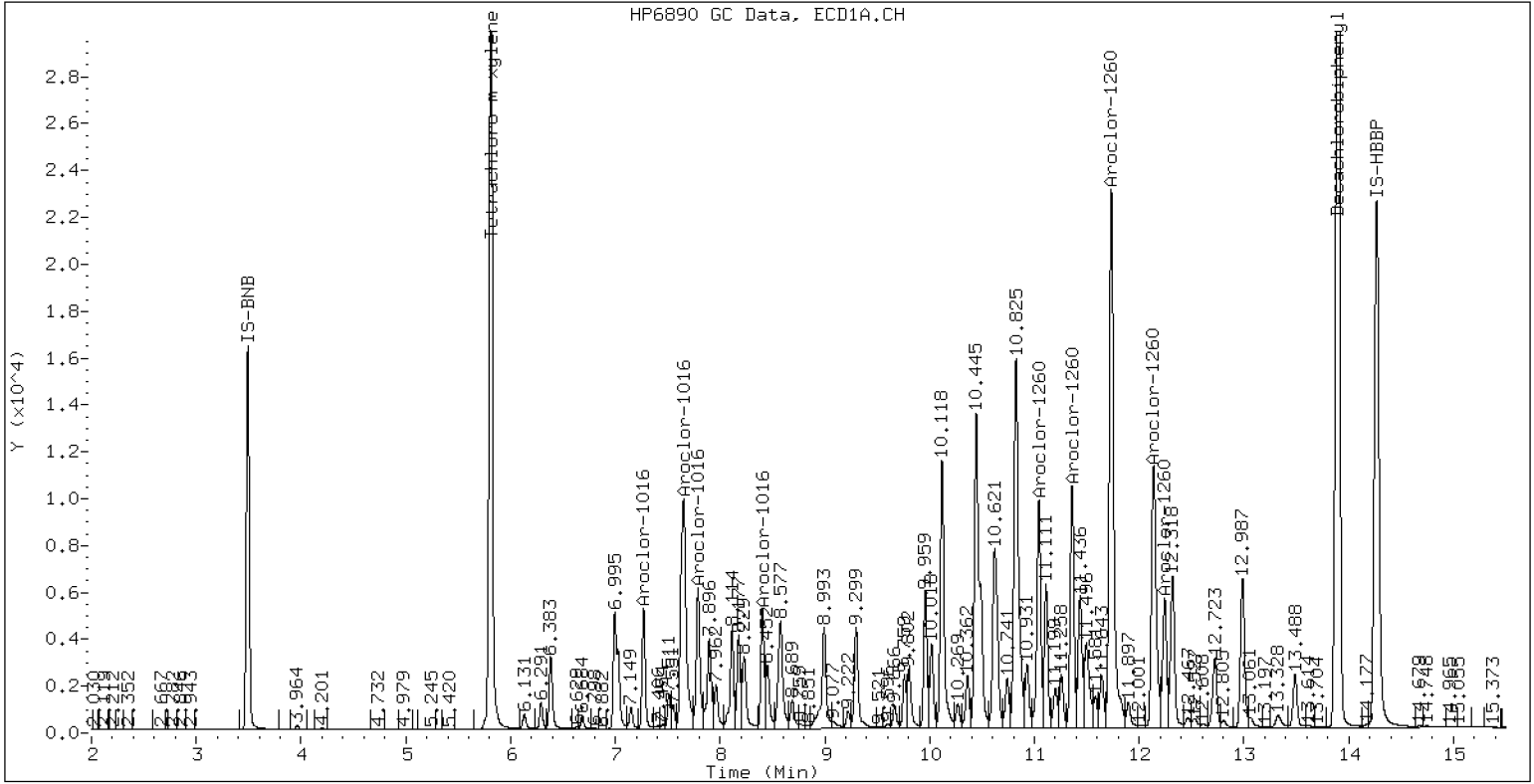
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 1.0PPMAR1660

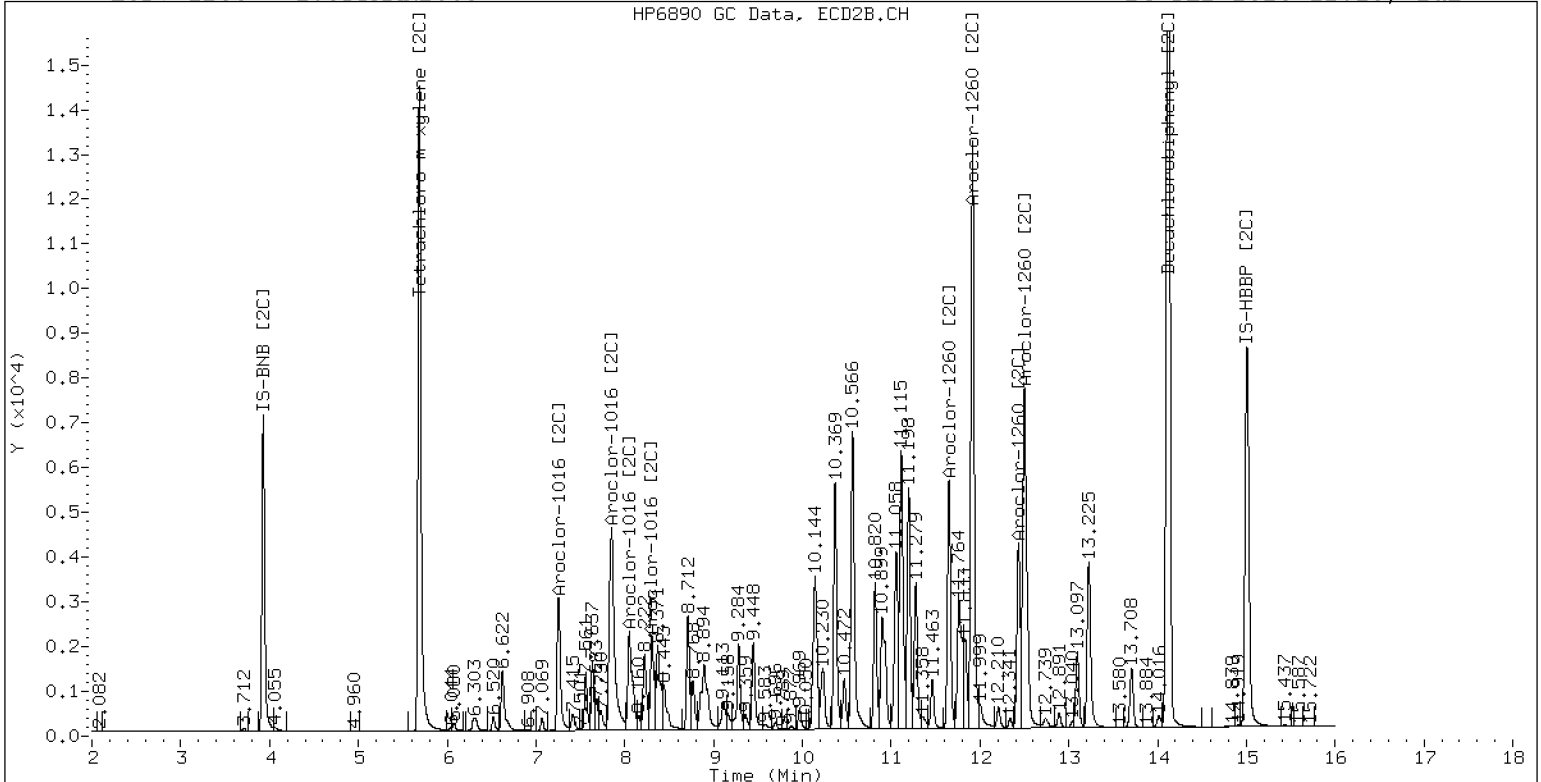
24-FEB-2023 12:15, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 1.0PPMAR1660

24-FEB-2023 12:15, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242306ECD7.D
Data file 2: /230224.b/230224.b/02242306ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.1PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 12:36
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.002	155528	5.688	0.003	74628	15.9	16.0	0.9	Tetrachloro-m-xylene
13.892	-0.001	227253	14.119	-0.000	128496	15.8	15.8	0.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	655979	-2.6
Hexabromobiphenyl	1429847	1464509	2.4

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	317418	0.7
Hexabromobiphenyl	513946	532962	3.7

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.271	0.000	25761	103.4	1	7.255	-0.000	19315	103.9
Aroclor-1016	2	7.657	0.003	75616	99.6	2	7.863	0.007	40308	107.0
Aroclor-1016	3	7.794	0.004	39547	106.7	3	8.059	0.005	18304	107.6
Aroclor-1016	4	8.406	0.001	24260	101.2	4	8.309	0.002	14708	110.2
Total CollAve (4 peaks):				102.7		Total Col2Ave (4 peaks):				107.2 RPD = 4
Corrected Ave (3 peaks):				101.4		Corrected Ave (3 peaks):				106.2 RPD = 5
CalAmt %D:				2.7		CalAmt %D:				7.2
Aroclor-1260	1	11.045	0.000	52009	98.7	1	11.655	0.002	31282	99.8
Aroclor-1260	2	11.362	0.001	55116	100.1	2	11.920	0.003	80574	100.7
Aroclor-1260	3	11.738	0.004	145604	99.7	3	12.437	0.002	19566	92.2
Aroclor-1260	4	12.141	0.002	72408	98.5	4	12.503	0.001	53588	99.4
Aroclor-1260	5	12.245	0.001	30745	97.2	NS	---			----
Total CollAve (5 peaks):				98.8		Total Col2Ave (4 peaks):				98.0 RPD = 1
Corrected Ave (4 peaks):				98.5		Corrected Ave (3 peaks):				97.1 RPD = 1
CalAmt %D:				-1.2		CalAmt %D:				-2.0

Total PCB Area Coll (5.906 - 13.793) = 1555762 Coll Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 764924 Col2 Total PCB = 0.2 ppm*

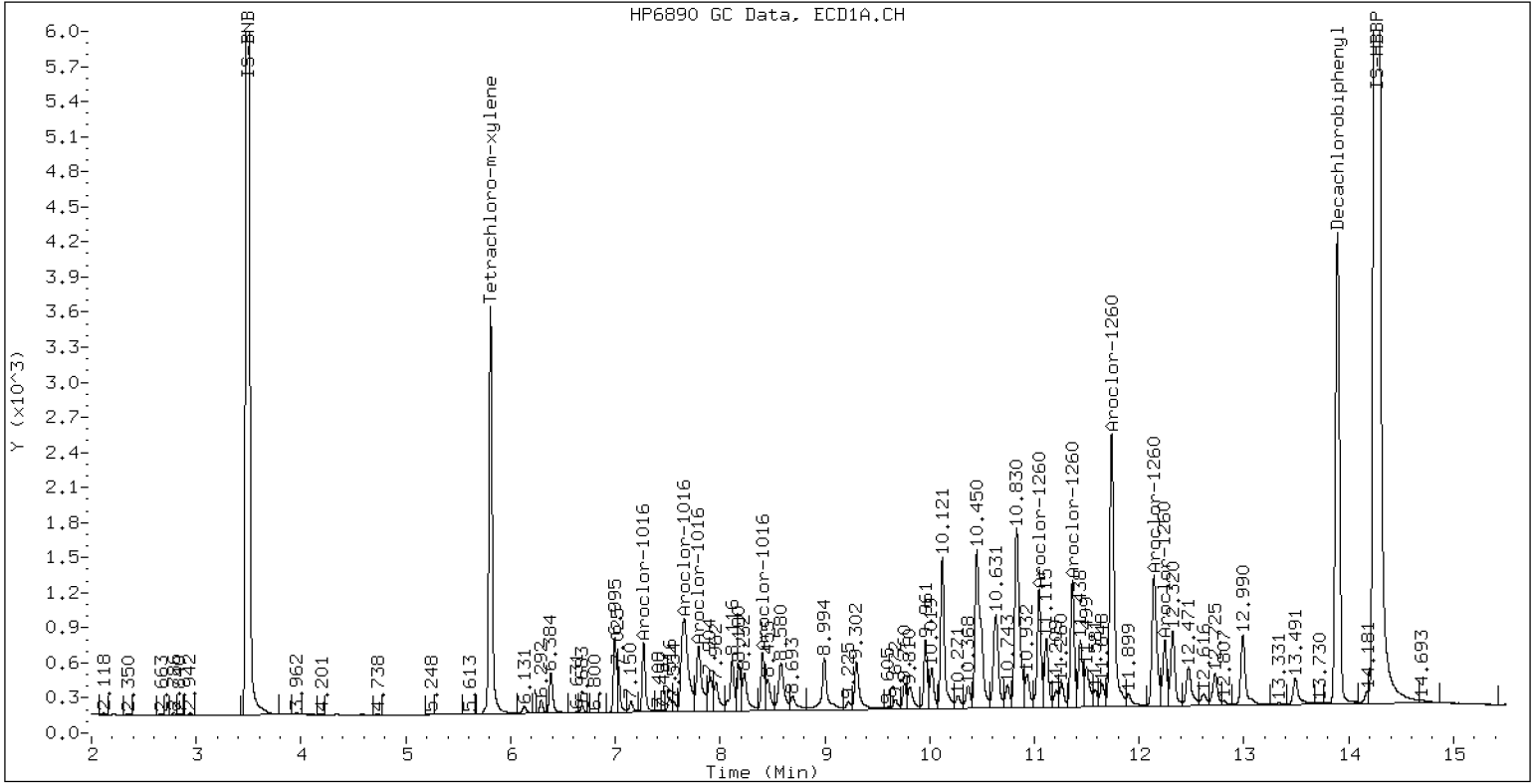
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.1PPMAR1660

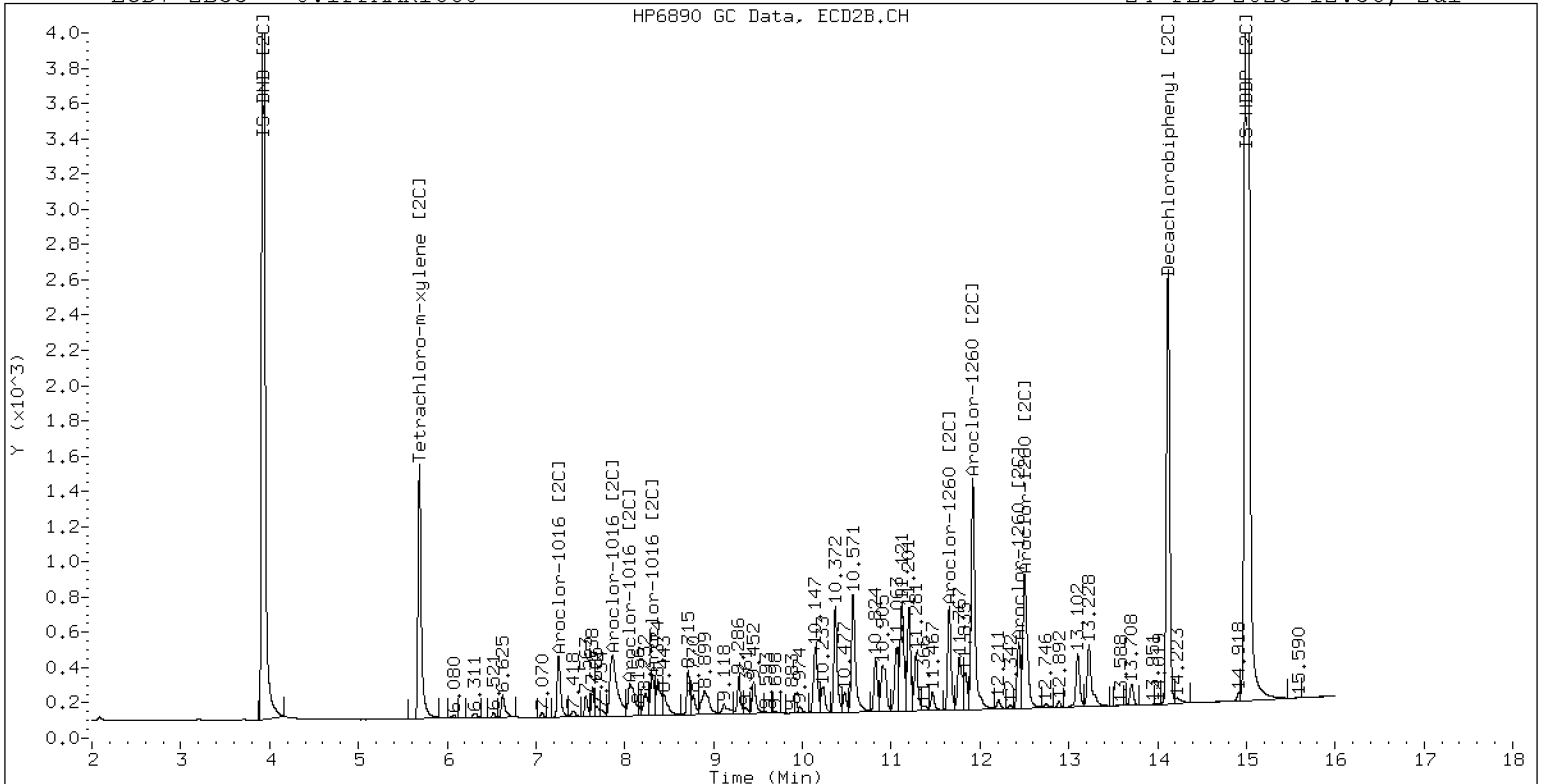
24-FEB-2023 12:36, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 0.1PPMAR1660

24-FEB-2023 12:36, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242307ECD7.D
Data file 2: /230224.b/230224.b/02242307ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.5PPMAR1660
Client ID:
Injection Date: 24-FEB-2023 12:57
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.810	0.004	724614	5.688	0.003	359257	75.2	76.7	2.0	Tetrachloro-m-xylene
13.898	0.005	1056911	14.120	0.000	650153	74.3	79.5	6.9	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	645275	-4.2
Hexabromobiphenyl	1429847	1445345	1.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	319170	1.2
Hexabromobiphenyl	513946	536853	4.5

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	0.000	115193	470.0	1	7.256	0.000	86287	461.8	
Aroclor-1016	2	7.654	0.000	369991	495.2	2	7.856	0.000	192524	508.1	
Aroclor-1016	3	7.790	0.000	160952	441.3	3	8.055	0.000	81039	473.6	
Aroclor-1016	4	8.405	0.000	115032	487.9	4	8.307	0.000	62136	462.8	
Total CollAve (4 peaks):				473.6		Total Col2Ave (4 peaks):				476.6	RPD = 1
Corrected Ave (3 peaks):				466.4		Corrected Ave (3 peaks):				466.1	RPD = 0

CalAmt %D: -5.3

CalAmt %D: -4.7

Aroclor-1260	1	11.044	0.000	247212	475.5	1	11.653	0.000	145247	460.1	
Aroclor-1260	2	11.361	0.000	262877	483.9	2	11.918	0.000	379838	471.5	
Aroclor-1260	3	11.734	0.000	678830	471.1	3	12.436	0.000	104092	486.9	
Aroclor-1260	4	12.139	0.000	356067	490.7	4	12.502	0.000	258953	476.9	
Aroclor-1260	5	12.244	0.000	150280	481.2	NS	---			----	
Total CollAve (5 peaks):				480.5		Total Col2Ave (4 peaks):				473.8	RPD = 1
Corrected Ave (4 peaks):				477.9		Corrected Ave (3 peaks):				469.5	RPD = 2

CalAmt %D: -3.9

CalAmt %D: -5.2

Total PCB Area Coll (5.906 - 13.793) = 7134169 Coll Total PCB = 0.9 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 3589735 Col2 Total PCB = 0.9 ppm*

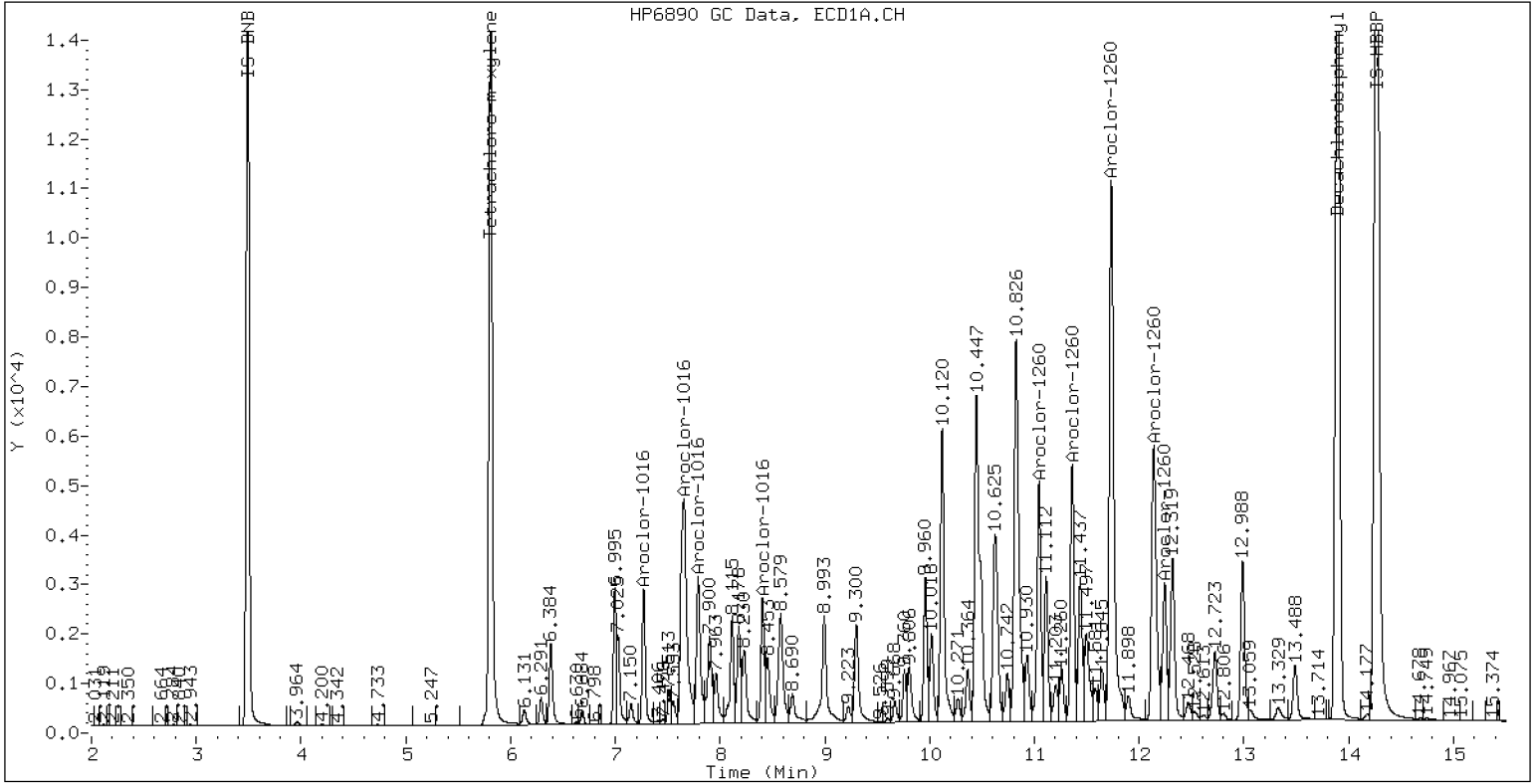
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.5PPMAR1660

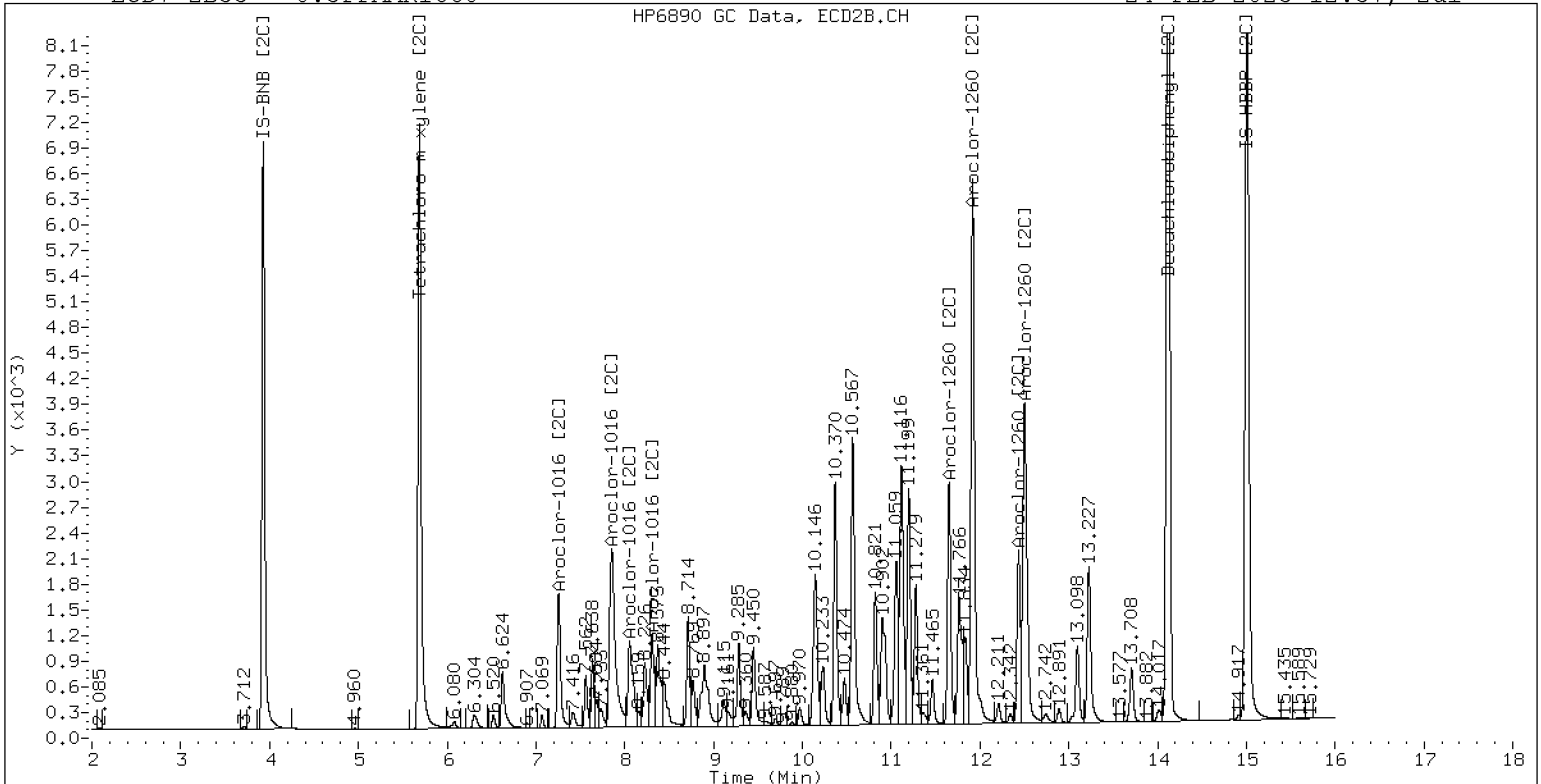
24-FEB-2023 12:57, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 0.5PPMAR1660

24-FEB-2023 12:57, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242308ECD7.D
Data file 2: /230224.b/230224.b/02242308ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1242.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1242
Client ID:
Injection Date: 24-FEB-2023 13:18
Report Date: 02/28/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.003	434187	5.688	0.003	214306	46.0	46.5	1.1	Tetrachloro-m-xylene
13.894	0.000	515867	14.119	-0.001	312943	35.6	38.5	7.6	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	632576	-6.1
Hexabromobiphenyl	1429847	1469715	2.8

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	314129	-0.4
Hexabromobiphenyl	513946	534294	4.0

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1242	1	7.271	0.000	49009	250.0	1	7.255	0.000	36487	250.0
Aroclor-1242	2	7.656	0.000	148833	250.0	2	7.858	0.000	76699	250.0
Aroclor-1242	3	8.405	0.000	46308	250.0	3	9.167	0.000	23866	250.0
Aroclor-1242	4	8.579	0.000	68453	250.0	4	9.597	0.000	29080	250.0
Total Col1Ave (4 peaks):				250.0	Total Col2Ave (4 peaks):				250.0	RPD = 0
Corrected Ave (3 peaks):				250.0	Corrected Ave (3 peaks):				250.0	RPD = 0

Total PCB Area Col1 (5.906 - 13.793) = 1221467 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 572067 Col2 Total PCB = 0.2 ppm*

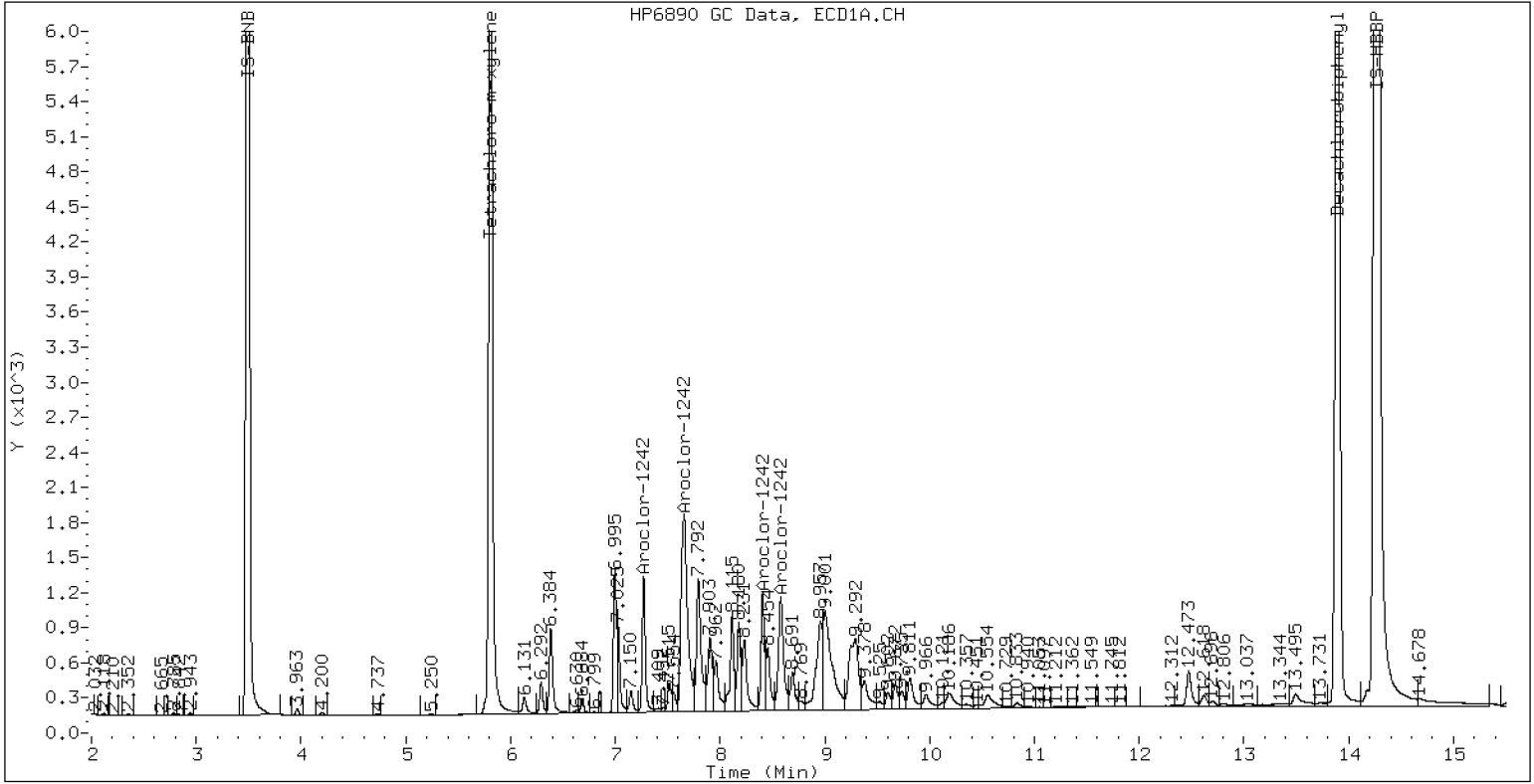
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1242

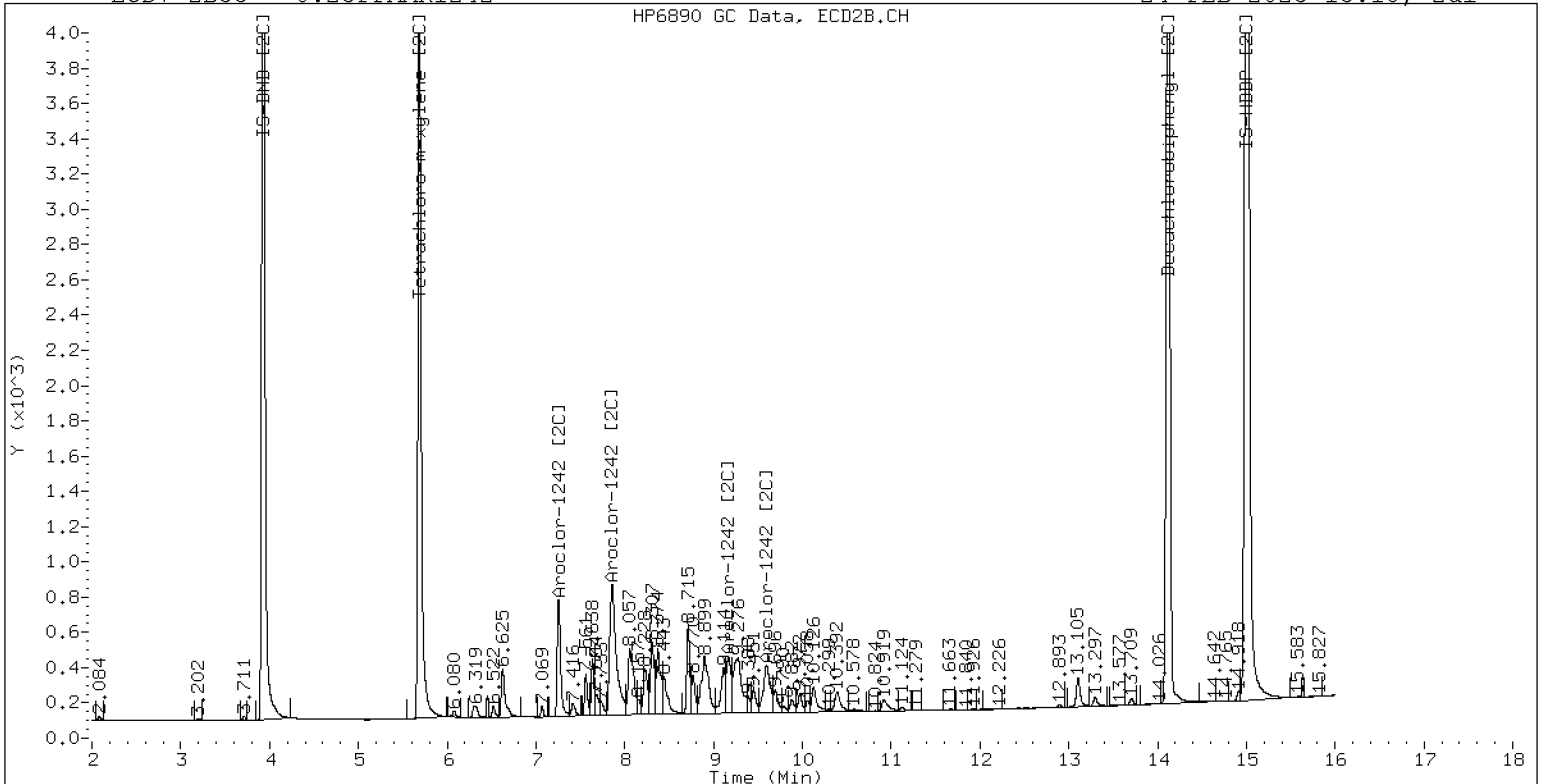
24-FEB-2023 13:18, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1242

24-FEB-2023 13:18, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242309ECD7.D
Data file 2: /230224.b/230224.b/02242309ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1248.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1248
Client ID:
Injection Date: 24-FEB-2023 13:39
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.809	0.003	349513	5.688	0.003	176615	36.6	37.9	3.4	Tetrachloro-m-xylene
13.894	0.001	523008	14.121	0.001	322054	36.4	39.3	7.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	639911	-5.0
Hexabromobiphenyl	1429847	1458696	2.0
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	317938	0.9
Hexabromobiphenyl	513946	538760	4.8

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1248	1	8.405	0.000	78055	250.0	1	8.308	0.000	37951	250.0
Aroclor-1248	2	8.580	0.000	99216	250.0	2	8.714	0.000	39239	250.0
Aroclor-1248	3	8.999	0.000	187178	250.0	3	9.166	0.000	45157	250.0
Aroclor-1248	4	9.295	0.000	95291	250.0	4	9.590	0.000	54216	250.0
Total CollAve (4 peaks):				250.0	Total Col2Ave (4 peaks):				250.0	RPD = 0
Corrected Ave (3 peaks):				250.0	Corrected Ave (3 peaks):				250.0	RPD = 0

Total PCB Area Coll (5.906 - 13.793) = 1565180 Coll Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 754991 Col2 Total PCB = 0.2 ppm*

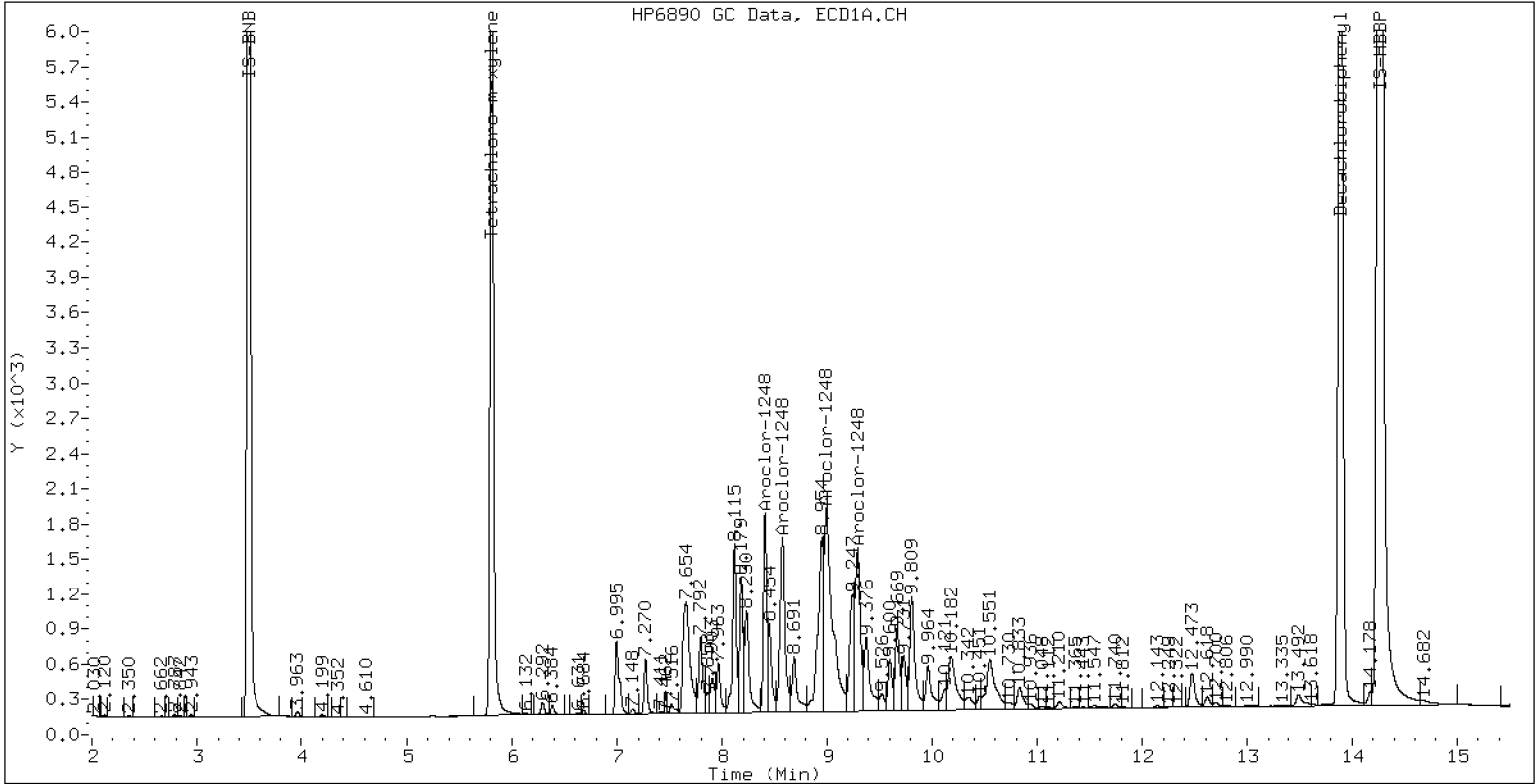
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1248

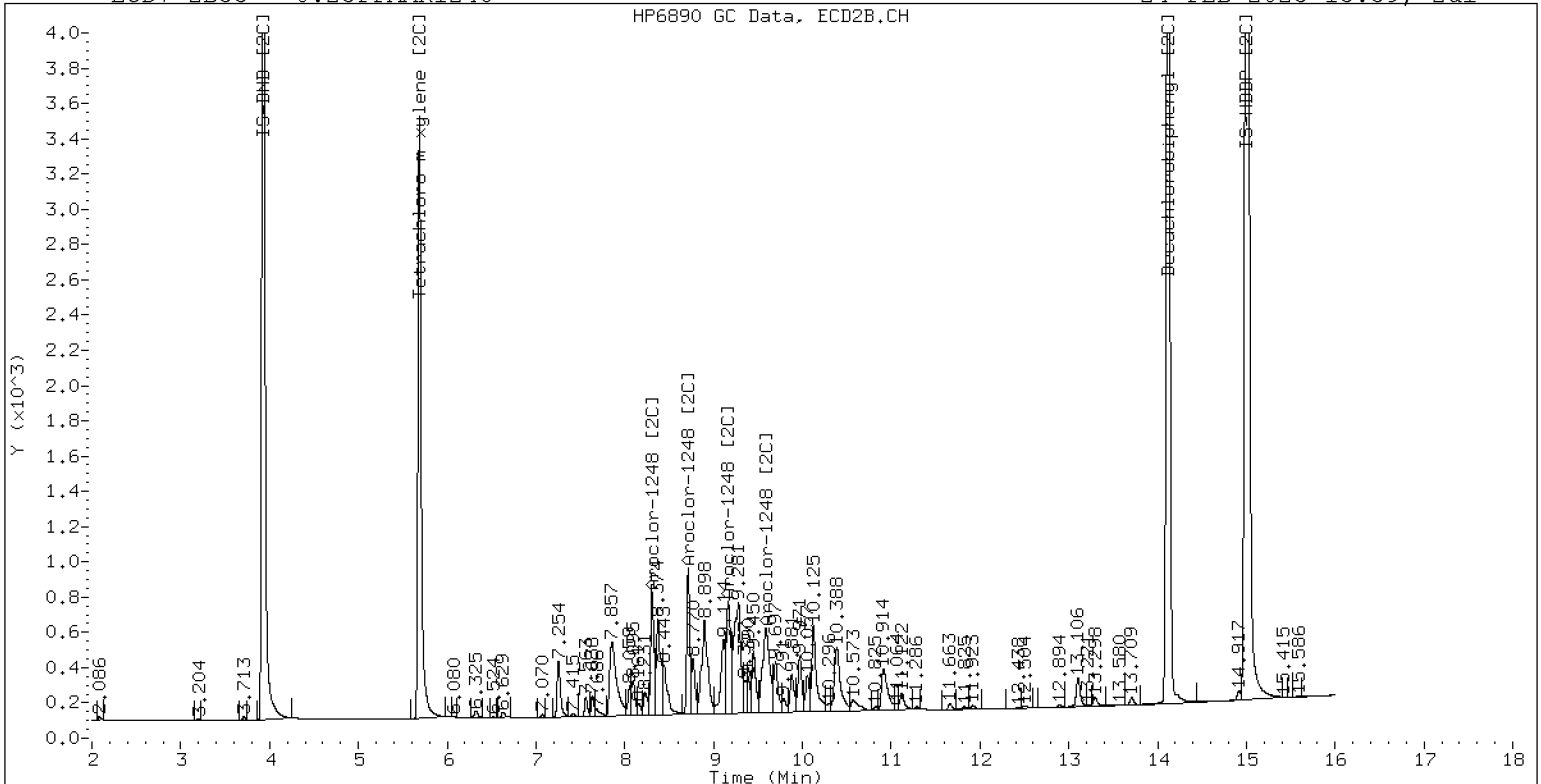
24-FEB-2023 13:39, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1248

24-FEB-2023 13:39, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242310ECD7.D
Data file 2: /230224.b/230224.b/02242310ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR1254.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR1254
Client ID:
Injection Date: 24-FEB-2023 14:00
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.808	0.002	352587	5.687	0.002	177502	37.3	38.6	3.4	Tetrachloro-m-xylene
13.895	0.002	532500	14.119	0.000	325903	37.0	40.2	8.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	633407	-6.0
Hexabromobiphenyl	1429847	1460265	2.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	313673	-0.5
Hexabromobiphenyl	513946	532442	3.6

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col ZB35 Col

Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1254	1	9.299	0.000	159011	250.0	1	9.449	0.000	59603	250.0
Aroclor-1254	2	9.377	0.000	71516	250.0	2	9.970	0.000	47949	250.0
Aroclor-1254	3	9.668	0.000	102230	250.0	3	10.124	0.000	103745	250.0
Aroclor-1254	4	9.807	0.000	198777	250.0	4	10.373	0.000	101135	250.0
Aroclor-1254	5	10.176	0.000	124586	250.0	5	10.569	0.000	61577	250.0
Total CollAve (5 peaks):				250.0		Total Col2Ave (5 peaks):				250.0 RPD = 0
Corrected Ave (4 peaks):				250.0		Corrected Ave (4 peaks):				250.0 RPD = 0

Total PCB Area Coll (5.906 - 13.793) = 2179224 Coll Total PCB = 0.3 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1022156 Col2 Total PCB = 0.3 ppm*

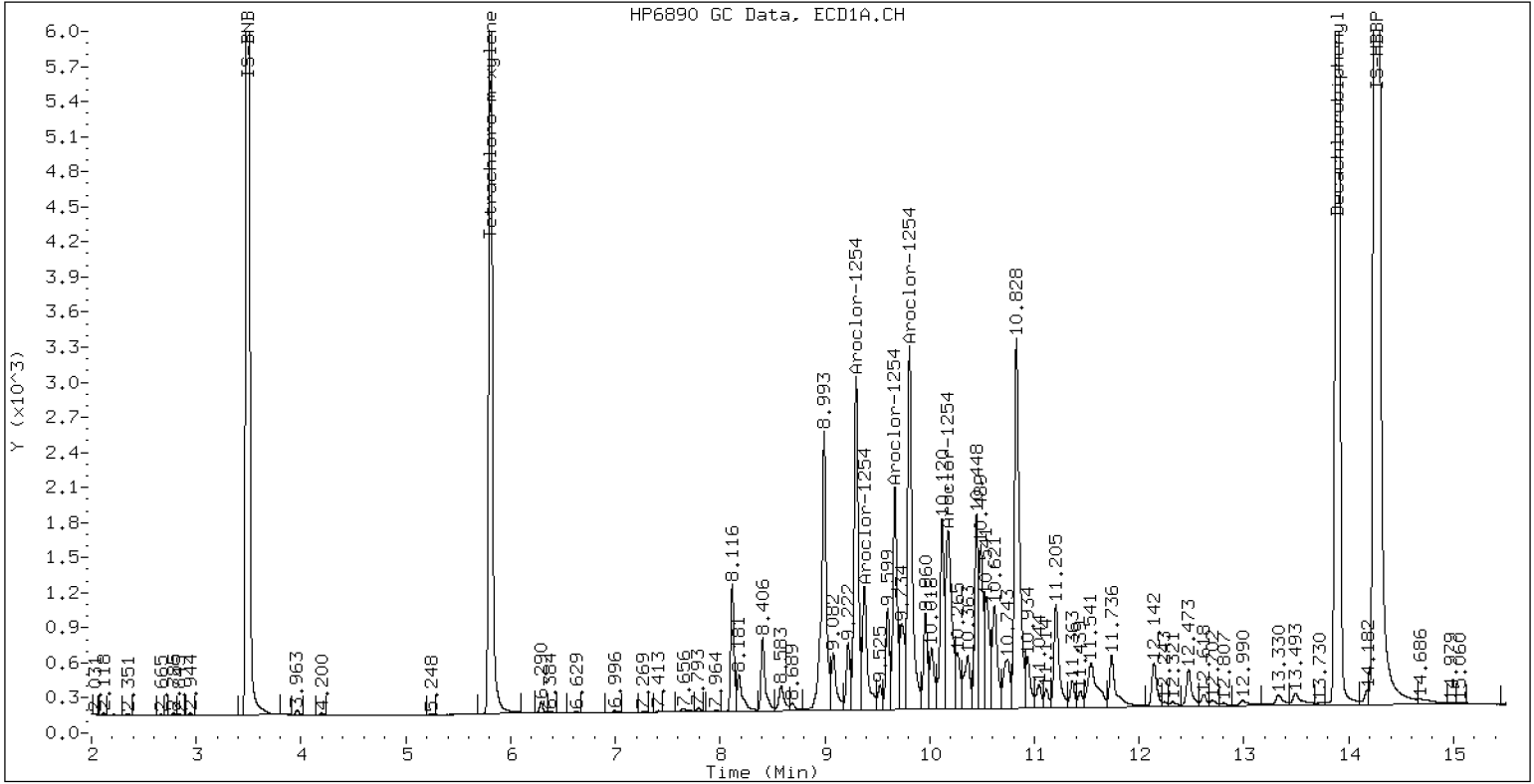
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR1254

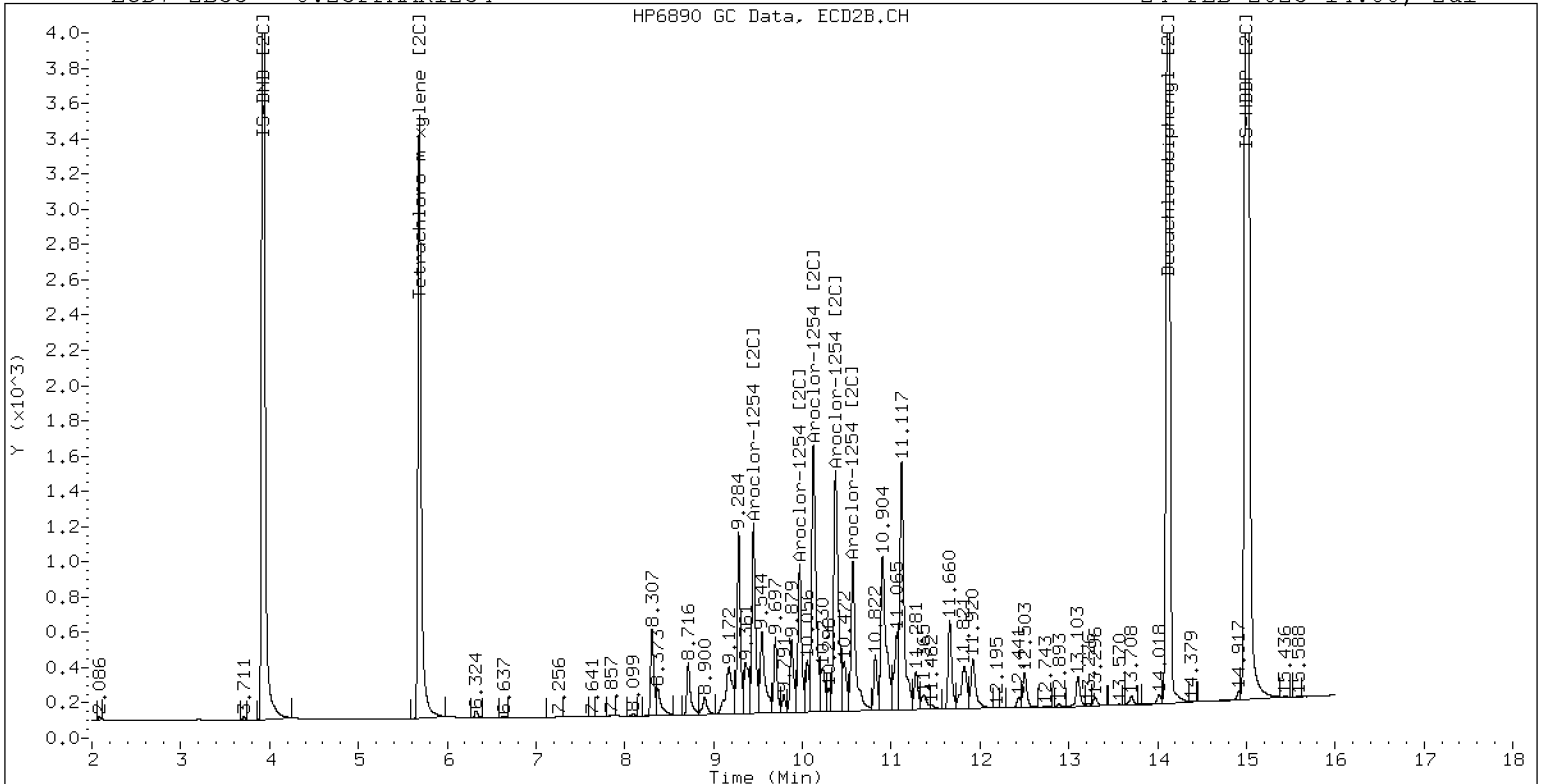
24-FEB-2023 14:00, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR1254

24-FEB-2023 14:00, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242311ECD7.D
Data file 2: /230224.b/230224.b/02242311ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR2162.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR2162
Client ID:
Injection Date: 24-FEB-2023 14:21
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.807	0.001	362236	5.686	0.000	177349	38.4	39.2	2.1	Tetrachloro-m-xylene
13.894	0.001	523254	14.119	-0.000	321034	36.0	39.2	8.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	632433	-6.1
Hexabromobiphenyl	1429847	1474039	3.1
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	308453	-2.2
Hexabromobiphenyl	513946	538177	4.7

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1221	1	4.731	0.000	14160	250.0	1	4.956	0.000	7300	250.0
Aroclor-1221	2	6.132	0.000	25324	250.0	2	6.296	0.000	13816	250.0
Aroclor-1221	3	6.382	0.000	58795	250.0	3	6.622	0.000	22491	250.0
Total CollAve (3 peaks):				250.0		Total Col2Ave (3 peaks):				250.0 RPD = 0
Corrected Ave: < 3 Peaks						Corrected Ave: < 3 Peaks				

Aroclor-1262	1	10.829	0.000	113046	250.0	1	11.200	0.000	114880	250.0
Aroclor-1262	2	12.244	0.000	183948	250.0	2	11.652	0.000	97844	250.0
Aroclor-1262	3	12.319	0.000	197749	250.0	3	12.434	0.000	111015	250.0
Aroclor-1262	4	12.987	0.000	180727	250.0	4	12.502	0.000	173913	250.0
Total CollAve (4 peaks):				250.0		Total Col2Ave (4 peaks):				250.0 RPD = 0
Corrected Ave (3 peaks):				250.0		Corrected Ave (3 peaks):				250.0 RPD = 0

Total PCB Area Coll (5.906 - 13.793) = 3105316 Coll Total PCB = 0.4 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1573107 Col2 Total PCB = 0.4 ppm*

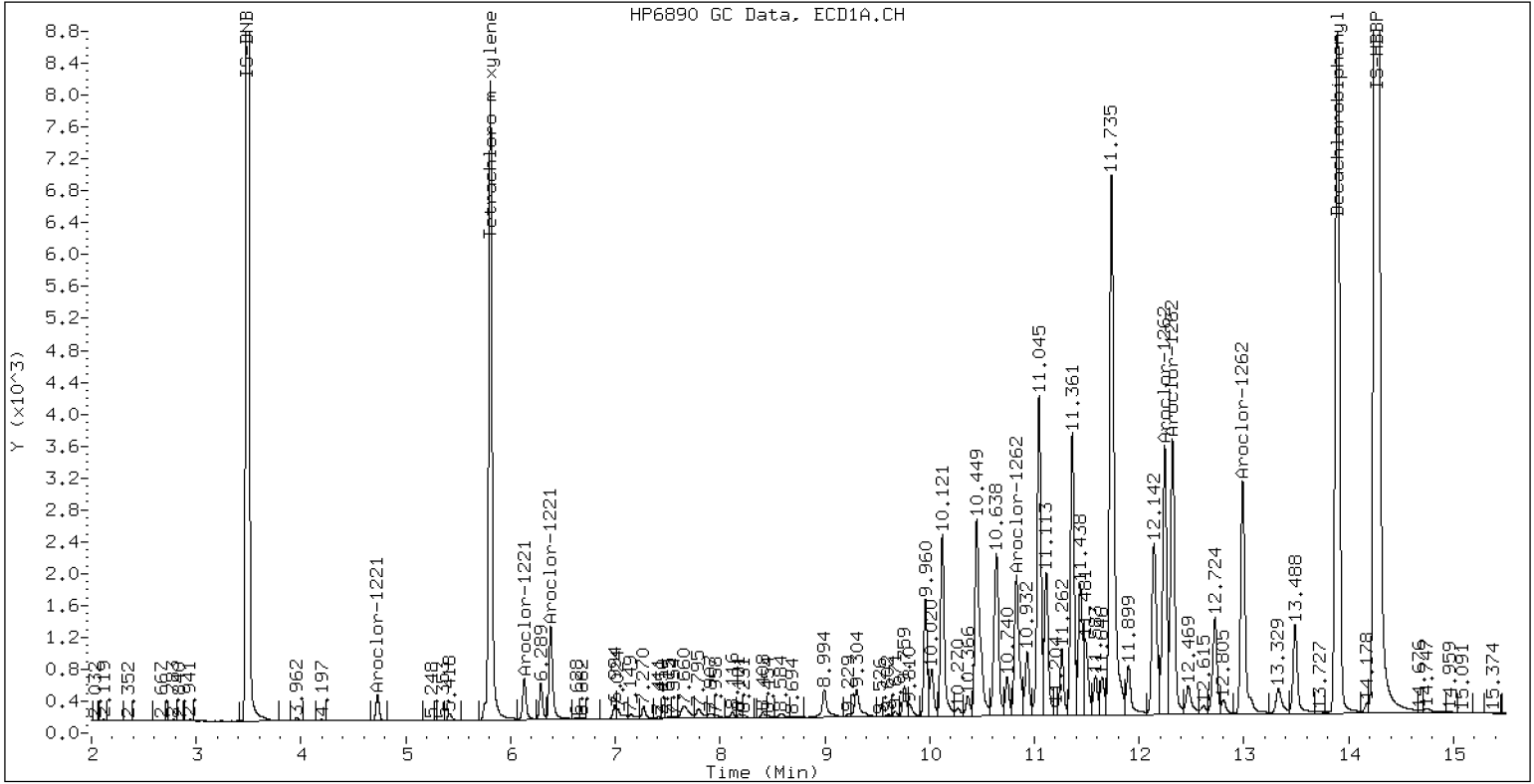
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR2162

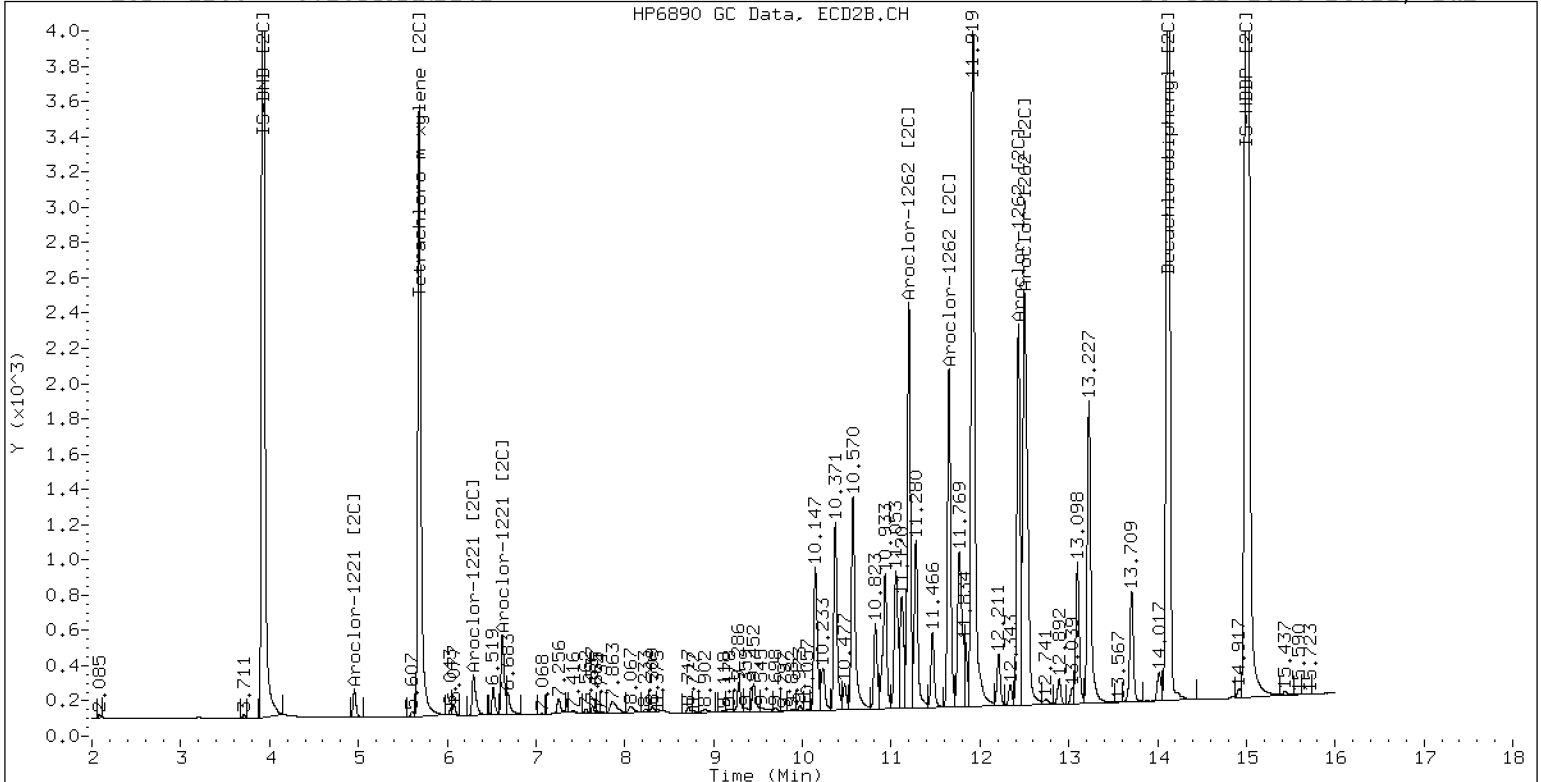
24-FEB-2023 14:21, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR2162

24-FEB-2023 14:21, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242312ECD7.D
Data file 2: /230224.b/230224.b/02242312ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: AR3268.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: 0.25PPMAR3268
Client ID:
Injection Date: 24-FEB-2023 14:42
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.806	0.000	366416	5.685	0.000	179450	38.0	38.9	2.4	Tetrachloro-m-xylene
13.893	0.000	778191	14.119	0.000	477889	53.0	57.5	8.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	645602	-4.2
Hexabromobiphenyl	1429847	1492154	4.4

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	314042	-0.4
Hexabromobiphenyl	513946	545458	6.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col				
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1232	1	4.730	0.000	8647	250.0	1	4.956	0.000	4017	250.0
Aroclor-1232	2	6.131	0.000	17148	250.0	2	7.254	0.000	19962	250.0
Aroclor-1232	3	7.656	0.000	77627	250.0	3	7.861	0.000	39913	250.0
Aroclor-1232	4	8.581	0.000	32993	250.0	4	8.715	0.000	11487	250.0
Total CollAve (4 peaks):				250.0		Total Col2Ave (4 peaks):				250.0 RPD = 0
Corrected Ave (3 peaks):				250.0		Corrected Ave (3 peaks):				250.0 RPD = 0
Aroclor-1268	1	12.247	0.000	477974	250.0	1	12.432	0.000	274595	250.0
Aroclor-1268	2	12.317	0.000	473326	250.0	2	12.500	0.000	295194	250.0
Aroclor-1268	3	12.699	0.000	405011	250.0	3	12.892	0.000	252048	250.0
Aroclor-1268	4	13.490	0.000	1333528	250.0	4	13.709	0.000	805579	250.0
Total CollAve (4 peaks):				250.0		Total Col2Ave (4 peaks):				250.0 RPD = 0
Corrected Ave (3 peaks):				250.0		Corrected Ave (3 peaks):				250.0 RPD = 0

Total PCB Area Coll (5.906 - 13.793) = 3998414 Coll Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 2300029 Col2 Total PCB = 0.6 ppm*

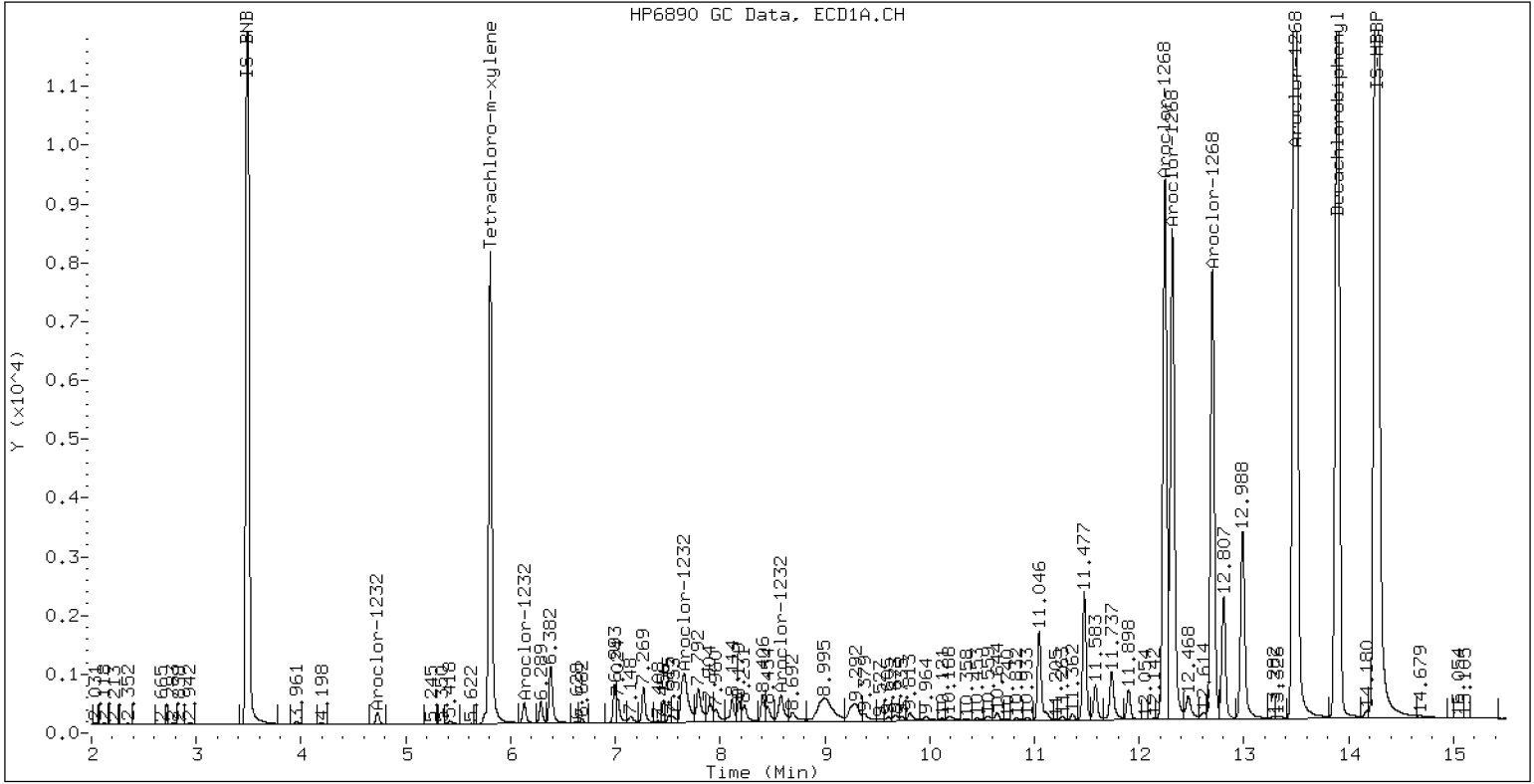
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 0.25PPMAR3268

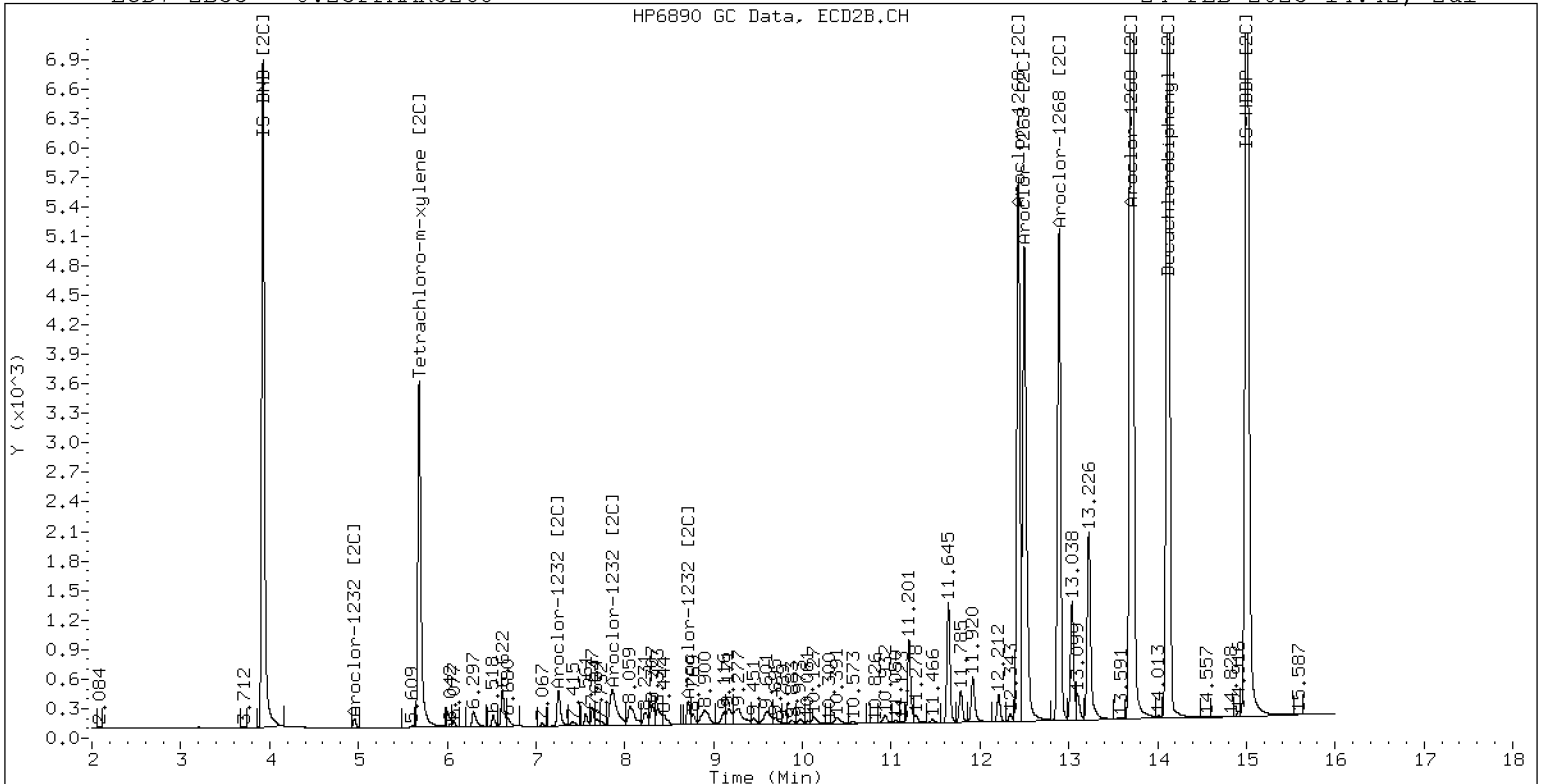
24-FEB-2023 14:42, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 0.25PPMAR3268

24-FEB-2023 14:42, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242313ECD7.D
Data file 2: /230224.b/230224.b/02242313ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1660SCV
Client ID:
Injection Date: 24-FEB-2023 15:03
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.807	0.001	337070	5.686	0.001	165848	34.9	35.8	2.3	Tetrachloro-m-xylene
13.895	0.002	515407	14.119	-0.000	316730	34.3	37.3	8.4	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	645975	-4.1
Hexabromobiphenyl	1429847	1524245	6.6

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	316115	0.3
Hexabromobiphenyl	513946	556950	8.4

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.269	-0.002	59491	242.5	1	7.254	-0.002	44576	240.9	
Aroclor-1016	2	7.655	0.001	181090	242.1	2	7.857	0.002	95386	254.2	
Aroclor-1016	3	7.790	0.000	88470	242.3	3	8.056	0.002	42160	248.8	
Aroclor-1016	4	8.404	-0.001	57980	245.6	4	8.307	0.000	32197	242.1	
Total CollAve (4 peaks):				243.1	Total Col2Ave (4 peaks):				246.5	RPD = 1	
Corrected Ave (3 peaks):				242.3	Corrected Ave (3 peaks):				243.9	RPD = 1	
Aroclor-1221	1	4.731	0.000	464	8.0	1	---			0.0	
Aroclor-1221	2	6.130	-0.002	9233	89.2	2	6.300	0.004	5379	95.0	
Aroclor-1221	3	6.382	-0.001	42570	177.2	3	6.623	0.001	20952	227.2	
Total CollAve (3 peaks):				91.5	Col2Ave: <3 Quant Peaks						
Aroclor-1232	1	4.731	0.001	464	13.4	1	---			0.0	
Aroclor-1232	2	6.130	-0.001	9233	134.5	2	7.254	-0.000	44576	554.6	
Aroclor-1232	3	7.655	-0.001	181090	582.9	3	7.857	-0.003	95386	593.5	
Aroclor-1232	4	8.580	-0.001	79916	605.2	4	8.713	-0.002	29795	644.2	
Total CollAve (4 peaks):				334.0	Total Col2Ave (3 peaks):				597.4	RPD = 57*	
Corrected Ave (3 peaks):				243.6	Corrected Ave: < 3 Peaks						
Aroclor-1242	1	7.269	-0.002	59491	297.2	1	7.254	-0.002	44576	303.5	
Aroclor-1242	2	7.655	-0.001	181090	297.9	2	7.857	-0.000	95386	309.0	
Aroclor-1242	3	8.404	-0.001	57980	306.5	3	9.115	-0.052	18754	195.2	
Aroclor-1242	4	8.580	0.000	79916	285.8	4	9.697	0.100	1355	11.6	
Total CollAve (4 peaks):				296.8	Total Col2Ave (4 peaks):				204.8	RPD = 37	
Corrected Ave (3 peaks):				293.6	Corrected Ave (3 peaks):				170.1	RPD = 53*	
Aroclor-1248	1	8.404	-0.001	57980	184.0	1	8.307	-0.001	32197	213.3	
Aroclor-1248	2	8.580	-0.001	79916	199.5	2	8.713	-0.001	29795	190.9	
Aroclor-1248	3	8.993	-0.006	71805	95.0	3	9.115	-0.050	18754	104.4	
Aroclor-1248	4	9.300	0.006	47348	123.1	4	---			0.0	
Total CollAve (4 peaks):				150.4	Total Col2Ave (3 peaks):				169.6	RPD = 12	
Corrected Ave (3 peaks):				134.0	Corrected Ave: < 3 Peaks						
Aroclor-1254	1	9.300	0.002	47348	73.0	1	9.451	0.001	22438	93.4	
Aroclor-1254	2	---			0.0	2	9.972	0.001	2694	13.9	
Aroclor-1254	3	9.670	0.002	5461	13.1	3	10.147	0.024	52914	126.5	
Aroclor-1254	4	9.807	-0.000	18944	23.4	4	10.370	-0.003	70430	172.8	
Aroclor-1254	5	10.121	-0.056	154170	303.3	5	10.568	-0.000	98525	396.9	
Total CollAve (4 peaks):				103.2	Total Col2Ave (5 peaks):				160.7	RPD = 44*	
Corrected Ave (3 peaks):				36.5	Corrected Ave (4 peaks):				101.7	RPD = 94*	
Aroclor-1260	1	11.044	0.000	149195	272.1	1	11.653	0.000	82210	251.0	
Aroclor-1260	2	11.361	-0.000	153832	268.5	2	11.919	0.001	222226	265.9	
Aroclor-1260	3	11.736	0.002	396660	261.0	3	12.435	-0.000	59148	266.7	
Aroclor-1260	4	12.140	0.001	190448	248.9	4	12.504	0.002	147180	261.2	
Aroclor-1260	5	12.244	-0.000	91385	277.5	NS	---			----	
Total CollAve (5 peaks):				265.6	Total Col2Ave (4 peaks):				261.2	RPD = 2	
Corrected Ave (4 peaks):				262.6	Corrected Ave (3 peaks):				259.4	RPD = 1	
Aroclor-1262	1	10.827	-0.002	220238	471.0	1	11.199	-0.001	84479	177.6	
Aroclor-1262	2	12.244	0.000	91385	120.1	2	11.653	0.002	82210	203.0	
Aroclor-1262	3	12.320	0.001	113066	138.2	3	12.435	0.002	59148	128.7	
Aroclor-1262	4	12.988	0.001	102156	136.7	4	12.504	0.002	147180	204.4	
Total CollAve (4 peaks):				216.5	Total Col2Ave (4 peaks):				178.4	RPD = 19	
Corrected Ave (3 peaks):				131.7	Corrected Ave (3 peaks):				169.8	RPD = 25	
Aroclor-1268	1	12.244	-0.003	91385	46.8	1	12.435	0.003	59148	52.7	
Aroclor-1268	2	12.320	0.003	113066	58.5	2	12.504	0.004	147180	122.1	
Aroclor-1268	3	12.726	0.027	46633	28.2	3	12.893	0.001	2874	2.8	
Aroclor-1268	4	13.489	-0.000	25567	4.7	4	13.709	-0.000	13041	4.0	
Total CollAve (4 peaks):				34.5	Total Col2Ave (4 peaks):				45.4	RPD = 27	
Corrected Ave (3 peaks):				26.6	Corrected Ave (3 peaks):				19.8	RPD = 29	

Total PCB Area Col1 (5.906 - 13.793) = 3743076 Col1 Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1897008 Col2 Total PCB = 0.5 ppm*

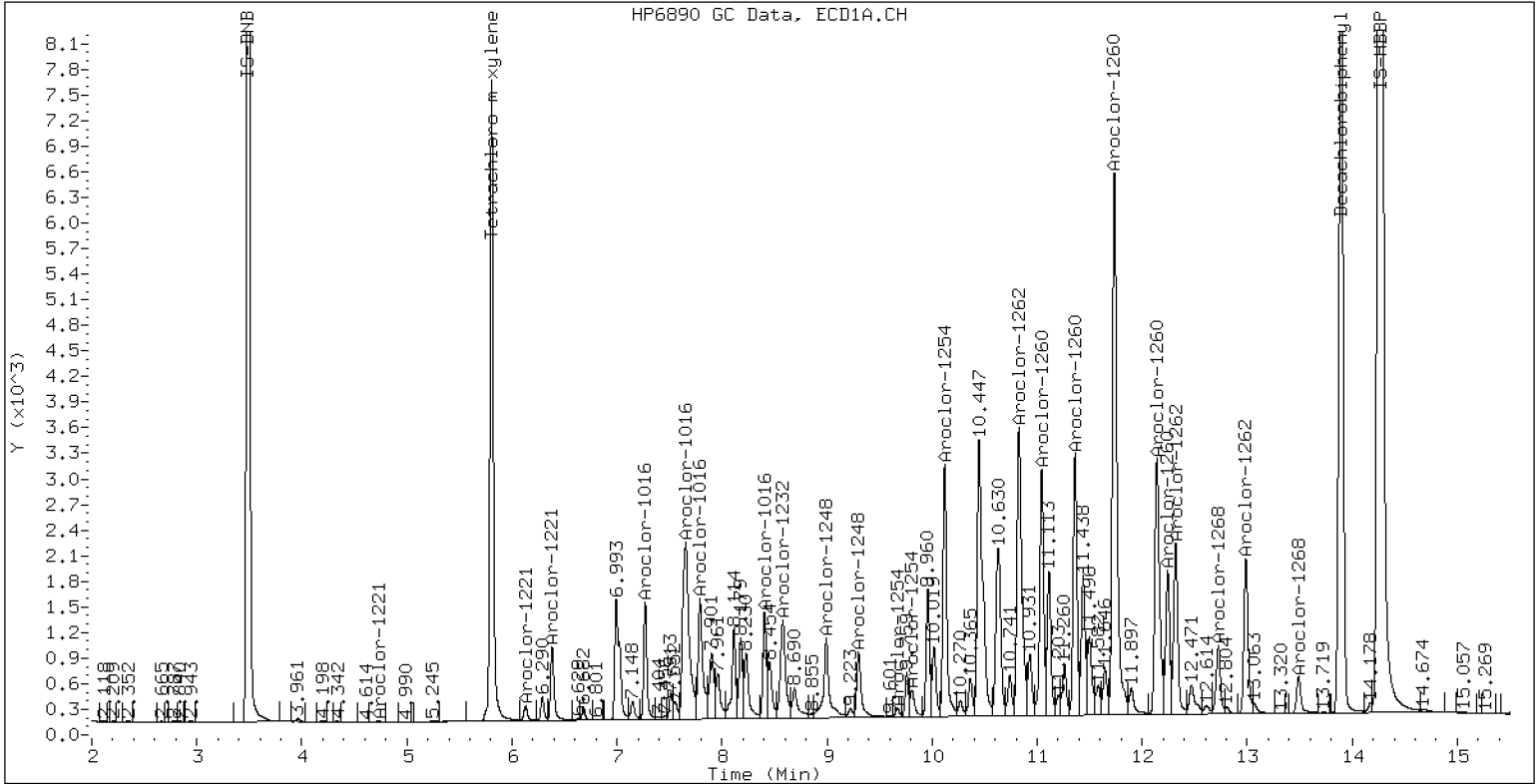
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1660SCV

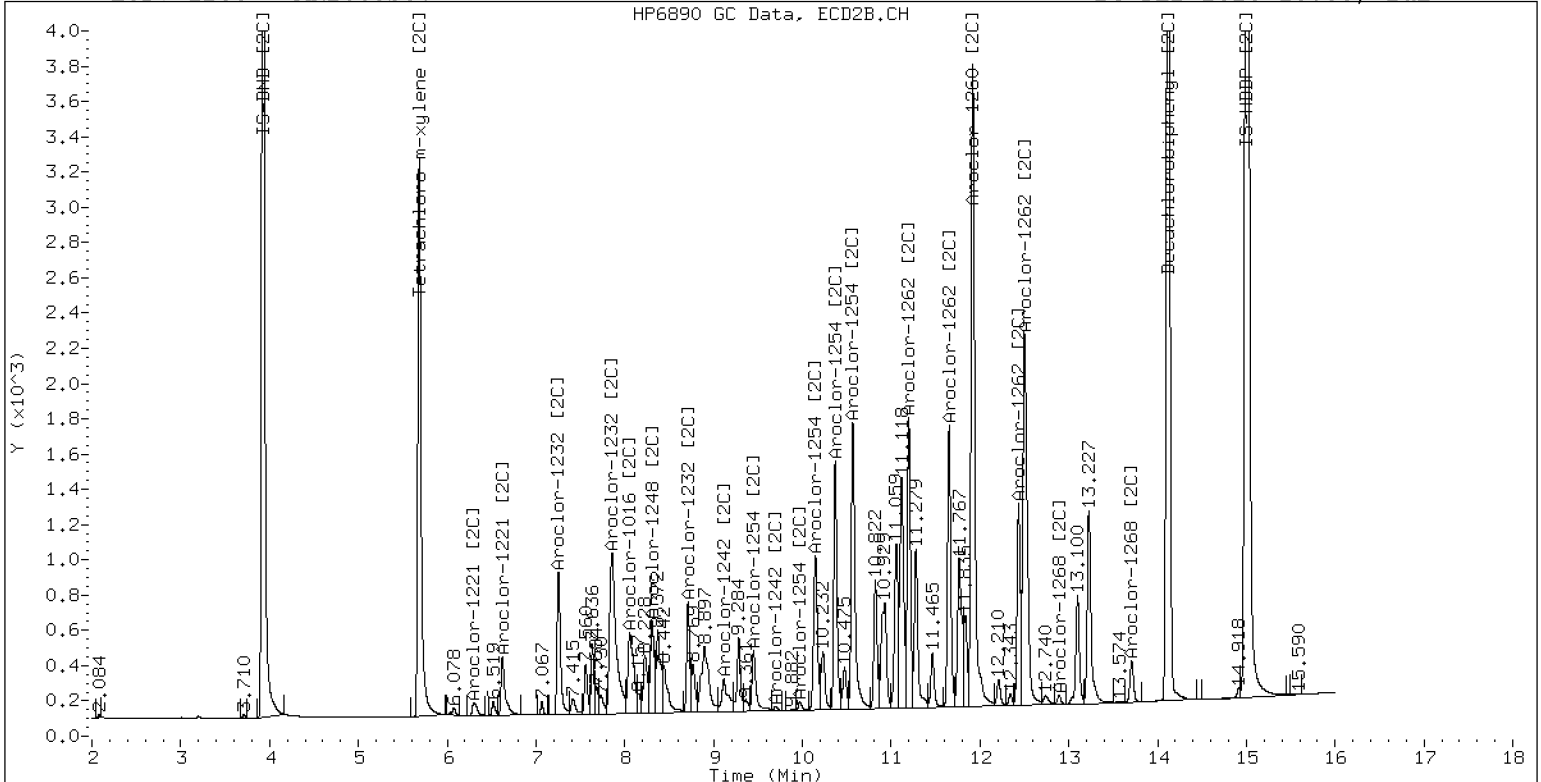
24-FEB-2023 15:03, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1660SCV

24-FEB-2023 15:03, 2u1



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242314ECD7.D
Data file 2: /230224.b/230224.b/02242314ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1242SCV
Client ID:
Injection Date: 24-FEB-2023 15:24
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.808	0.002	354283	5.686	0.001	172455	33.6	34.5	2.6	Tetrachloro-m-xylene
13.895	0.002	567088	14.120	0.001	347430	37.0	40.3	8.6	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	705650	4.7
Hexabromobiphenyl	1429847	1555683	8.8
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	340433	8.0
Hexabromobiphenyl	513946	565609	10.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.270	-0.000	39927	149.0	1	7.256	0.000	32417	162.7
Aroclor-1016	2	7.653	-0.001	132339	162.0	2	7.856	0.001	69235	171.3
Aroclor-1016	3	7.791	0.001	59310	148.7	3	8.055	0.000	29473	161.5
Aroclor-1016	4	8.405	0.000	42537	165.0	4	8.307	-0.000	22792	159.2
Total CollAve (4 peaks):				156.2		Total Col2Ave (4 peaks):				163.7 RPD = 5
Corrected Ave (3 peaks):				153.2		Corrected Ave (3 peaks):				161.1 RPD = 5
Aroclor-1221	1	4.733	0.002	319	5.0	1	---			0.0
Aroclor-1221	2	6.131	-0.001	6534	57.8	2	6.319	0.022	4365	71.6
Aroclor-1221	3	6.384	0.001	29664	113.0	3	6.624	0.002	14916	150.2
Total CollAve (3 peaks):				58.6		Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	4.733	0.003	319	8.4	1	---			0.0
Aroclor-1232	2	6.131	0.000	6534	87.2	2	7.256	0.002	32417	374.5
Aroclor-1232	3	7.653	-0.003	132339	389.9	3	7.856	-0.004	69235	400.0
Aroclor-1232	4	8.579	-0.002	69445	481.4	4	8.714	-0.001	22167	445.0
Total CollAve (4 peaks):				241.7		Total Col2Ave (3 peaks):				406.5 RPD = 51*
Corrected Ave (3 peaks):				161.8		Corrected Ave: < 3 Peaks				
Aroclor-1242	1	7.270	-0.001	39927	182.6	1	7.256	0.000	32417	205.0
Aroclor-1242	2	7.653	-0.003	132339	199.3	2	7.856	-0.002	69235	208.2
Aroclor-1242	3	8.405	-0.000	42537	205.9	3	9.164	-0.004	23068	223.0
Aroclor-1242	4	8.579	-0.000	69445	227.4	4	9.587	-0.010	31021	246.1
Total CollAve (4 peaks):				203.8		Total Col2Ave (4 peaks):				220.6 RPD = 8
Corrected Ave (3 peaks):				195.9		Corrected Ave (3 peaks):				212.1 RPD = 8
Aroclor-1248	1	8.405	0.000	42537	123.5	1	8.307	-0.001	22792	140.2
Aroclor-1248	2	8.579	-0.001	69445	158.7	2	8.714	-0.000	22167	131.9
Aroclor-1248	3	9.001	0.003	91942	111.4	3	9.164	-0.002	23068	119.3
Aroclor-1248	4	9.294	-0.000	38711	92.1	4	9.587	-0.003	31021	133.6
Total CollAve (4 peaks):				121.4		Total Col2Ave (4 peaks):				131.2 RPD = 8
Corrected Ave (3 peaks):				109.0		Corrected Ave (3 peaks):				128.3 RPD = 16
Aroclor-1254	1	9.294	-0.005	38711	54.6	1	9.450	0.001	13131	50.7
Aroclor-1254	2	9.377	-0.000	17371	54.5	2	9.970	0.000	8340	40.1
Aroclor-1254	3	9.668	-0.000	16373	35.9	3	10.123	-0.000	16364	36.3
Aroclor-1254	4	9.807	-0.001	27490	31.0	4	10.382	0.009	16062	36.6
Aroclor-1254	5	10.175	-0.001	20494	36.9	5	10.572	0.004	4818	18.0
Total CollAve (5 peaks):				42.6		Total Col2Ave (5 peaks):				36.4 RPD = 16
Corrected Ave (4 peaks):				39.6		Corrected Ave (4 peaks):				32.8 RPD = 19
Aroclor-1260	1	11.048	0.003	794	1.4	1	11.665	0.012	1652	5.0
Aroclor-1260	2	11.366	0.005	814	1.4	2	11.926	0.008	842	1.0
Aroclor-1260	3	11.739	0.006	1848	1.2	3	12.438	0.002	483	2.1
Aroclor-1260	4	12.145	0.006	1372	1.8	4	12.506	0.004	790	1.4
Aroclor-1260	5	---			0.0	NS	---			---
Total CollAve (4 peaks):				1.4		Total Col2Ave (4 peaks):				2.4 RPD = 49*
Corrected Ave (3 peaks):				1.3		Corrected Ave (3 peaks):				1.5 RPD = 12
Aroclor-1262	1	10.832	0.003	13157	27.6	1	11.121	-0.079	6113	12.7
Aroclor-1262	2	12.145	-0.098	1372	1.8	2	11.665	0.013	1652	4.0
Aroclor-1262	3	---			0.0	3	12.438	0.004	483	1.0
Aroclor-1262	4	13.038	0.051	842	1.1	4	12.506	0.004	790	1.1
Total CollAve (3 peaks):				10.1		Total Col2Ave (4 peaks):				4.7 RPD = 73*
Corrected Ave: < 3 Peaks						Corrected Ave (3 peaks):				2.0
Aroclor-1268	1	---			0.0	1	12.438	0.006	483	0.4
Aroclor-1268	2	---			0.0	2	12.506	0.006	790	0.6
Aroclor-1268	3	12.617	-0.082	5851	3.5	3	12.899	0.007	491	0.5
Aroclor-1268	4	13.500	0.010	1745	0.3	4	13.714	0.005	379	0.1
CollAve: <3 Quant Peaks						Col2Ave:				0.4

Total PCB Area Col1 (5.906 - 13.793) = 1149784 Col1 Total PCB = 0.1 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 572210 Col2 Total PCB = 0.1 ppm*

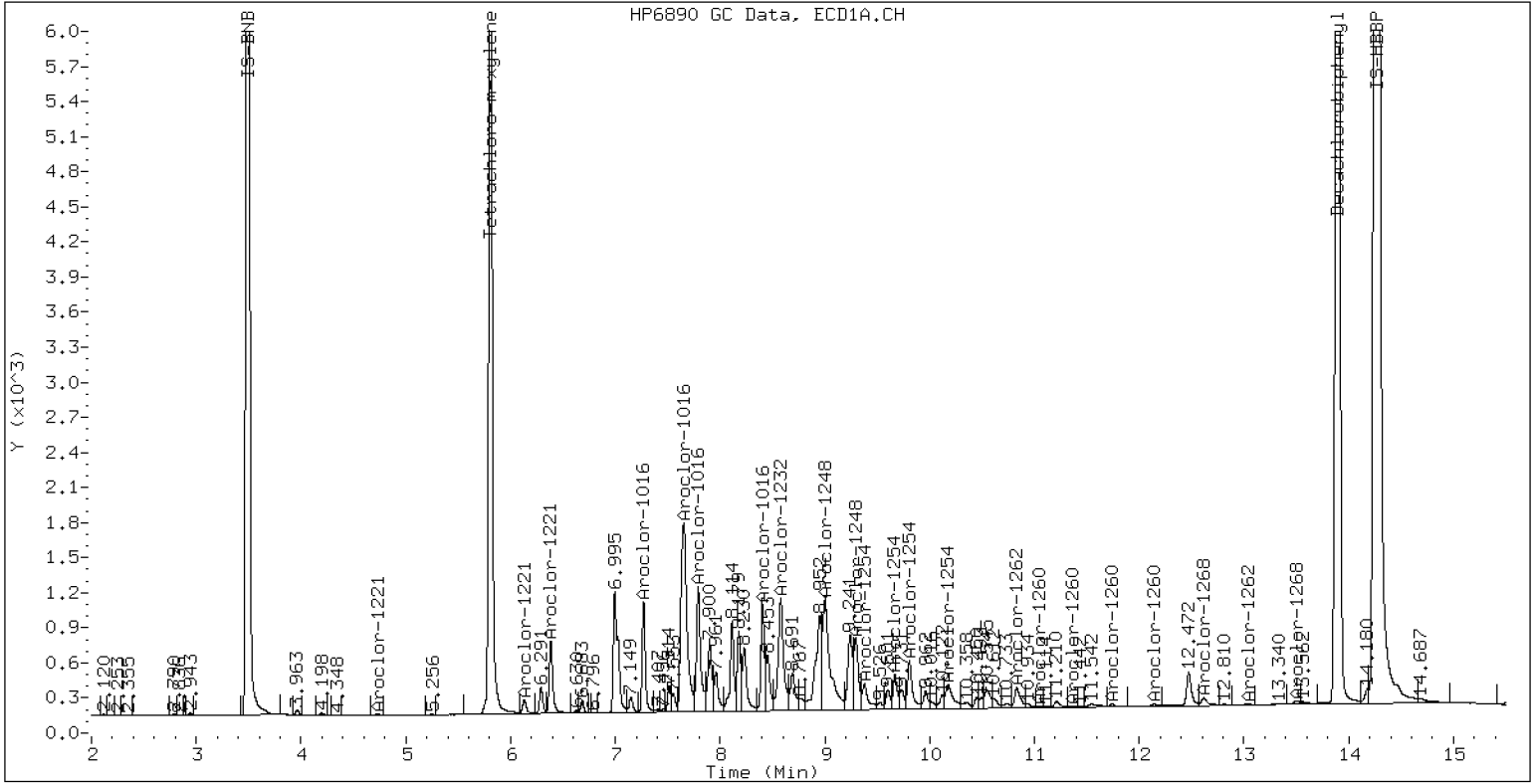
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1242SCV

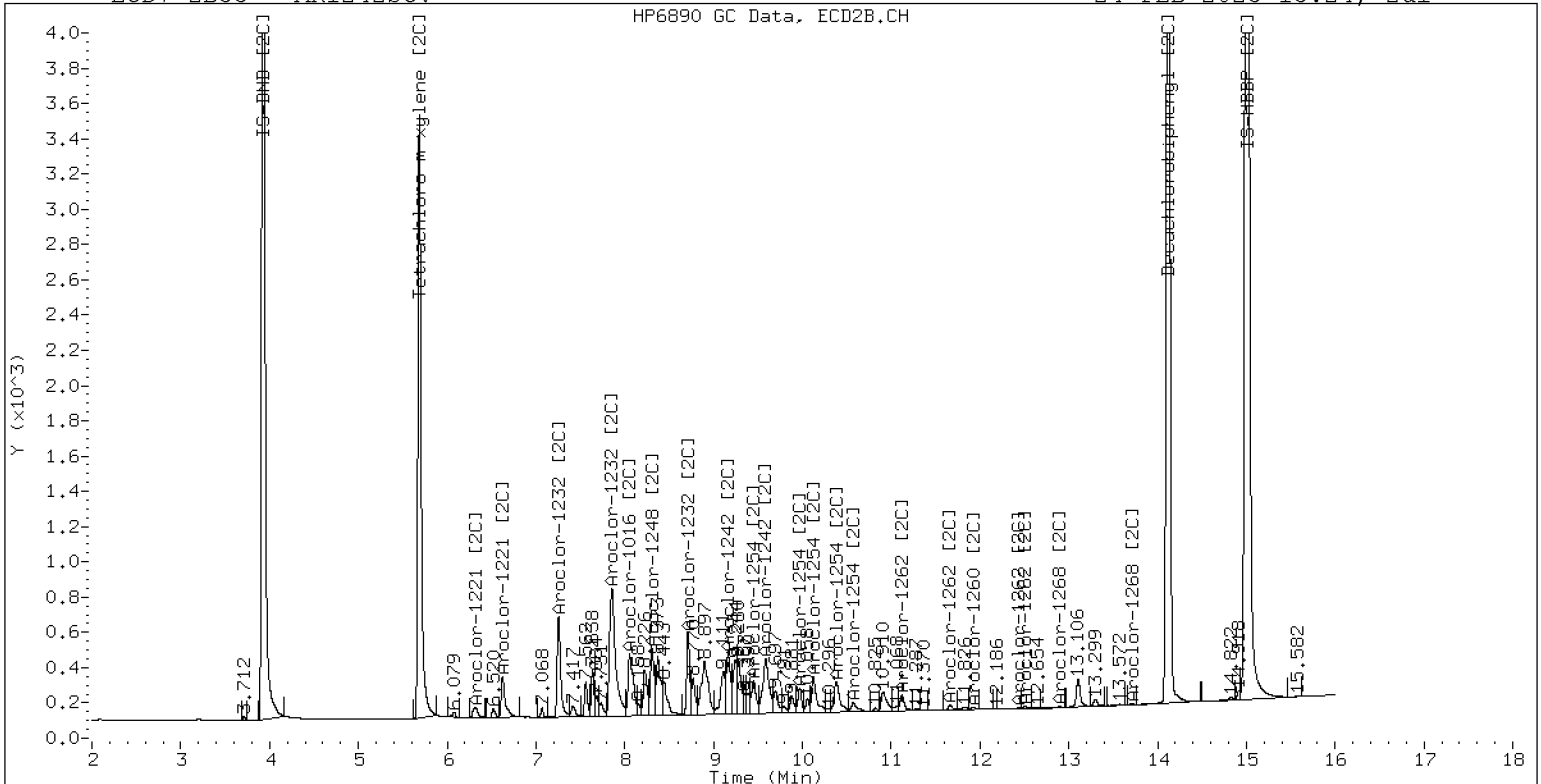
24-FEB-2023 15:24, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1242SCV

24-FEB-2023 15:24, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242315ECD7.D
Data file 2: /230224.b/230224.b/02242315ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1248SCV
Client ID:
Injection Date: 24-FEB-2023 15:45
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col		ZB5	ZB35	RPD	Compound/Flag		
RT	Shift	Response	RT	Shift	Response				
5.809	0.002	336655	5.687	0.002	168719	34.9	36.4	4.2	Tetrachloro-m-xylene
13.894	0.001	499162	14.118	-0.001	308317	33.1	36.3	9.2	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	646554	-4.0
Hexabromobiphenyl	1429847	1529451	7.0
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	316066	0.3
Hexabromobiphenyl	513946	557213	8.4

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.271	0.000	19773	80.5	1	7.254	-0.002	16926	91.5
Aroclor-1016	2	7.653	-0.001	88099	117.7	2	7.857	0.001	45733	121.9
Aroclor-1016	3	7.794	0.003	35915	98.3	3	8.060	0.005	8078	47.7
Aroclor-1016	4	8.406	0.001	77842	329.5	4	8.307	0.000	37348	280.9
Total CollAve (4 peaks):				156.5		Total Col2Ave (4 peaks):				135.5 RPD = 14
Corrected Ave (3 peaks):				98.8		Corrected Ave (3 peaks):				87.0 RPD = 13
Aroclor-1221	1	---			0.0	1	---			0.0
Aroclor-1221	2	6.133	0.001	680	6.6	2	6.326	0.030	1966	34.7
Aroclor-1221	3	6.384	0.002	3390	14.1	3	6.631	0.009	1571	17.0
CollAve: <3 Quant Peaks						Col2Ave: <3 Quant Peaks				
Aroclor-1232	1	---			0.0	1	---			0.0
Aroclor-1232	2	6.133	0.002	680	9.9	2	7.254	-0.000	16926	210.6
Aroclor-1232	3	7.653	-0.002	88099	283.3	3	7.857	-0.004	45733	284.6
Aroclor-1232	4	8.581	-0.000	99572	753.4	4	8.714	-0.001	38224	826.6
Total CollAve (3 peaks):				348.9		Total Col2Ave (3 peaks):				440.6 RPD = 23
Corrected Ave: < 3 Peaks						Corrected Ave: < 3 Peaks				
Aroclor-1242	1	7.271	-0.000	19773	98.7	1	7.254	-0.002	16926	115.3
Aroclor-1242	2	7.653	-0.003	88099	144.8	2	7.857	-0.001	45733	148.2
Aroclor-1242	3	8.406	0.000	77842	411.2	3	9.165	-0.002	45021	468.7
Aroclor-1242	4	8.581	0.001	99572	355.8	4	9.590	-0.008	53613	458.1
Total CollAve (4 peaks):				252.6		Total Col2Ave (4 peaks):				297.6 RPD = 16
Corrected Ave (3 peaks):				199.8		Corrected Ave (3 peaks):				240.5 RPD = 19
Aroclor-1248	1	8.406	0.000	77842	246.8	1	8.307	-0.001	37348	247.5
Aroclor-1248	2	8.581	0.000	99572	248.3	2	8.714	-0.000	38224	245.0
Aroclor-1248	3	8.998	-0.000	186857	247.0	3	9.165	-0.000	45021	250.7
Aroclor-1248	4	9.294	-0.000	98398	255.5	4	9.590	-0.001	53613	248.7
Total CollAve (4 peaks):				249.4		Total Col2Ave (4 peaks):				248.0 RPD = 1
Corrected Ave (3 peaks):				247.4		Corrected Ave (3 peaks):				247.0 RPD = 0
Aroclor-1254	1	9.294	-0.004	98398	151.6	1	9.450	0.001	21823	90.8
Aroclor-1254	2	9.377	-0.001	49616	169.9	2	9.971	0.001	19450	100.6
Aroclor-1254	3	9.669	0.001	40230	96.4	3	10.124	0.000	36574	87.5
Aroclor-1254	4	9.808	0.001	68500	84.4	4	10.389	0.016	35100	86.1
Aroclor-1254	5	10.183	0.007	47365	93.1	5	10.573	0.004	5676	22.9
Total CollAve (5 peaks):				119.1		Total Col2Ave (5 peaks):				77.6 RPD = 42*
Corrected Ave (4 peaks):				106.4		Corrected Ave (4 peaks):				71.8 RPD = 39
Aroclor-1260	1	11.047	0.003	1670	3.0	1	11.662	0.009	2055	6.3
Aroclor-1260	2	11.362	0.001	1111	1.9	2	11.924	0.007	1466	1.8
Aroclor-1260	3	11.739	0.005	2107	1.4	3	12.434	-0.002	573	2.6
Aroclor-1260	4	12.144	0.005	1379	1.8	4	12.505	0.003	1003	1.8
Aroclor-1260	5	12.251	0.006	698	2.1	NS	---			----
Total CollAve (5 peaks):				2.1		Total Col2Ave (4 peaks):				3.1 RPD = 41*
Corrected Ave (4 peaks):				1.8		Corrected Ave (3 peaks):				2.0 RPD = 12
Aroclor-1262	1	10.833	0.005	15355	32.7	1	11.122	-0.079	7225	15.2
Aroclor-1262	2	12.251	0.007	698	0.9	2	11.662	0.011	2055	5.1
Aroclor-1262	3	12.321	0.002	836	1.0	3	12.434	0.000	573	1.2
Aroclor-1262	4	12.991	0.004	1043	1.4	4	12.505	0.003	1003	1.4
Total CollAve (4 peaks):				9.0		Total Col2Ave (4 peaks):				5.7 RPD = 45*
Corrected Ave (3 peaks):				1.1		Corrected Ave (3 peaks):				2.6 RPD = 80*
Aroclor-1268	1	12.251	0.004	698	0.4	1	12.434	0.002	573	0.5
Aroclor-1268	2	12.321	0.004	836	0.4	2	12.505	0.005	1003	0.8
Aroclor-1268	3	12.700	0.001	2449	1.5	3	12.892	0.001	721	0.7
Aroclor-1268	4	13.493	0.003	7547	1.4	4	13.708	-0.001	2265	0.7
Total CollAve (4 peaks):				0.9		Total Col2Ave (4 peaks):				0.7 RPD = 29
Corrected Ave (3 peaks):				0.7		Corrected Ave (3 peaks):				0.6 RPD = 13

Total PCB Area Col1 (5.906 - 13.793) = 1574335 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 746330 Col2 Total PCB = 0.2 ppm*

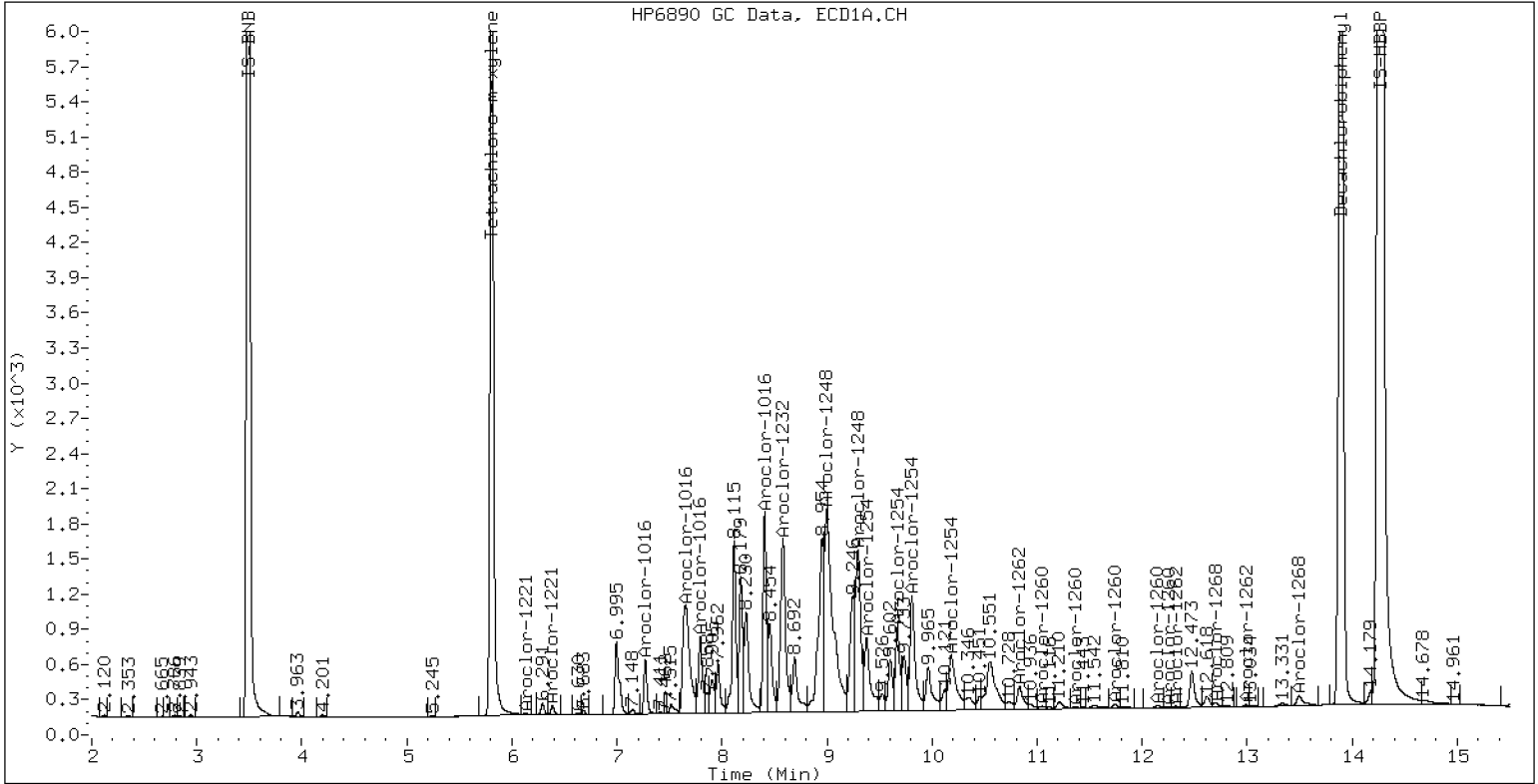
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1248SCV

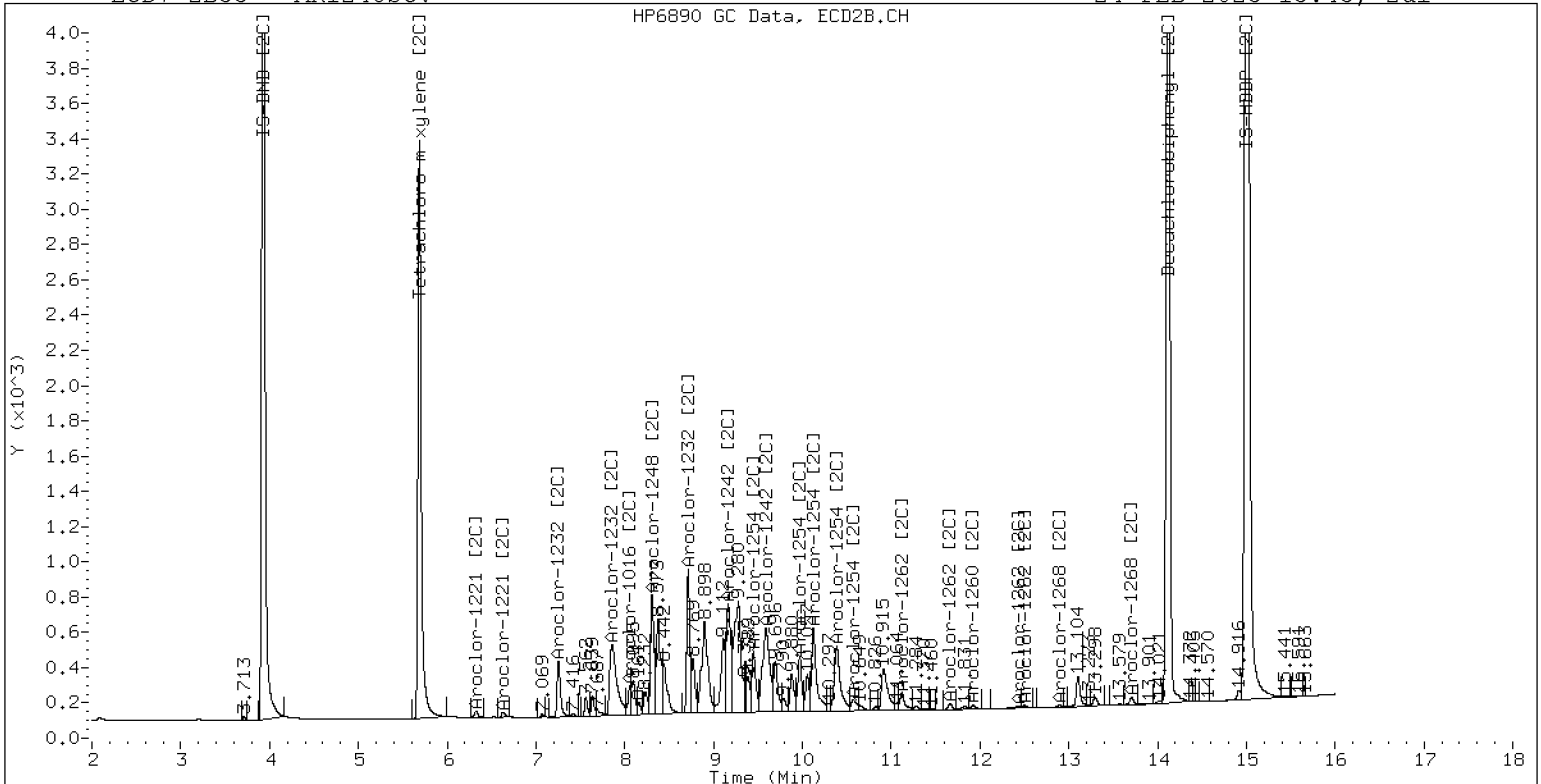
24-FEB-2023 15:45, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1248SCV

24-FEB-2023 15:45, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242316ECD7.D
Data file 2: /230224.b/230224.b/02242316ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1254SCV
Client ID:
Injection Date: 24-FEB-2023 16:06
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.806	-0.000	354312	5.686	0.001	174604	36.1	37.1	2.6	Tetrachloro-m-xylene
13.895	0.002	540961	14.119	-0.000	329134	34.6	37.9	9.0	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	656887	-2.5
Hexabromobiphenyl	1429847	1585505	10.9

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	320936	1.8
Hexabromobiphenyl	513946	570006	10.9

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	0.000	565	2.3	1	7.255	-0.001	387	2.1	
Aroclor-1016	2	7.656	0.002	1875	2.5	2	7.854	-0.002	860	2.3	
Aroclor-1016	3	7.792	0.002	1106	3.0	3	8.098	0.043	578	3.4	
Aroclor-1016	4	8.405	0.000	29924	124.7	4	8.307	0.000	21985	162.9	
Total CollAve (4 peaks):				33.1	Total Col2Ave (4 peaks):				42.6	RPD = 25	
Corrected Ave (3 peaks):				2.6	Corrected Ave (3 peaks):				2.6	RPD = 0	
Aroclor-1221	1	---			0.0	1	---			0.0	
Aroclor-1221	2	---			0.0	2	6.325	0.028	1947	33.9	
Aroclor-1221	3	---			0.0	3	6.637	0.015	368	3.9	
CollAve: <3 Quant Peaks					Col2Ave: <3 Quant Peaks						
Aroclor-1232	1	---			0.0	1	---			0.0	
Aroclor-1232	2	---			0.0	2	7.255	0.001	387	4.7	
Aroclor-1232	3	7.656	0.000	1875	5.9	3	7.854	-0.007	860	5.3	
Aroclor-1232	4	8.583	0.002	12327	91.8	4	8.715	0.000	15013	319.7	
CollAve: <3 Quant Peaks					Col2Ave: 109.9						
Aroclor-1242	1	7.270	-0.000	565	2.8	1	7.255	-0.001	387	2.6	
Aroclor-1242	2	7.656	0.000	1875	3.0	2	7.854	-0.004	860	2.7	
Aroclor-1242	3	8.405	-0.000	29924	155.6	3	9.169	0.002	21933	224.9	
Aroclor-1242	4	8.583	0.003	12327	43.4	4	9.545	-0.053	34065	286.6	
Total CollAve (4 peaks):				51.2	Total Col2Ave (4 peaks):				129.2	RPD = 87*	
Corrected Ave (3 peaks):				16.4	Corrected Ave (3 peaks):				76.7	RPD = 130*	
Aroclor-1248	1	8.405	0.000	29924	93.4	1	8.307	-0.001	21985	143.5	
Aroclor-1248	2	8.583	0.002	12327	30.3	2	8.715	0.001	15013	94.8	
Aroclor-1248	3	8.992	-0.007	145580	189.4	3	9.169	0.004	21933	120.3	
Aroclor-1248	4	9.298	0.003	155450	397.3	4	9.545	-0.046	34065	155.6	
Total CollAve (4 peaks):				177.6	Total Col2Ave (4 peaks):				128.5	RPD = 32	
Corrected Ave (3 peaks):				104.3	Corrected Ave (3 peaks):				119.5	RPD = 14	
Aroclor-1254	1	9.298	-0.001	155450	235.7	1	9.450	0.001	58639	240.4	
Aroclor-1254	2	9.377	-0.001	69801	235.3	2	9.971	0.000	47008	239.5	
Aroclor-1254	3	9.668	-0.000	100839	237.8	3	10.124	0.000	100062	235.7	
Aroclor-1254	4	9.807	0.000	190544	231.1	4	10.373	0.000	99535	240.5	
Aroclor-1254	5	10.176	-0.000	122321	236.7	5	10.570	0.001	61549	244.2	
Total CollAve (5 peaks):				235.3	Total Col2Ave (5 peaks):				240.1	RPD = 2	
Corrected Ave (4 peaks):				234.7	Corrected Ave (4 peaks):				239.0	RPD = 2	
Aroclor-1260	1	11.043	-0.002	12288	21.5	1	11.661	0.008	29062	86.7	
Aroclor-1260	2	11.361	-0.001	13660	22.9	2	11.921	0.003	22238	26.0	
Aroclor-1260	3	11.736	0.002	37632	23.8	3	12.441	0.005	3555	15.7	
Aroclor-1260	4	12.141	0.002	27105	34.1	4	12.503	0.001	13126	22.8	
Aroclor-1260	5	12.320	0.076	2381	6.9	NS	---			---	
Total CollAve (5 peaks):				21.9	Total Col2Ave (4 peaks):				37.8	RPD = 53*	
Corrected Ave (4 peaks):				18.8	Corrected Ave (3 peaks):				21.5	RPD = 13	
Aroclor-1262	1	10.827	-0.002	220626	453.6	1	11.281	0.081	13562	27.9	
Aroclor-1262	2	12.320	0.076	2381	3.0	2	11.661	0.009	29062	70.1	
Aroclor-1262	3	---			0.0	3	12.441	0.007	3555	7.6	
Aroclor-1262	4	12.989	0.002	3225	4.1	4	12.503	0.001	13126	17.8	
Total CollAve (3 peaks):				153.6	Total Col2Ave (4 peaks):				30.8	RPD = 133*	
Corrected Ave: < 3 Peaks					Corrected Ave (3 peaks):				17.7		
Aroclor-1268	1	12.320	0.074	2381	1.2	1	12.441	0.009	3555	3.1	
Aroclor-1268	2	---			0.0	2	12.503	0.003	13126	10.6	
Aroclor-1268	3	12.701	0.002	2939	1.7	3	12.892	0.000	772	0.7	
Aroclor-1268	4	13.493	0.003	9164	1.6	4	13.707	-0.002	2801	0.8	
Total CollAve (3 peaks):				1.5	Total Col2Ave (4 peaks):				3.8	RPD = 87*	
Corrected Ave: < 3 Peaks					Corrected Ave (3 peaks):				1.6		

Total PCB Area Col1 (5.906 - 13.793) = 2118645 Col1 Total PCB = 0.3 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 1007601 Col2 Total PCB = 0.3 ppm*

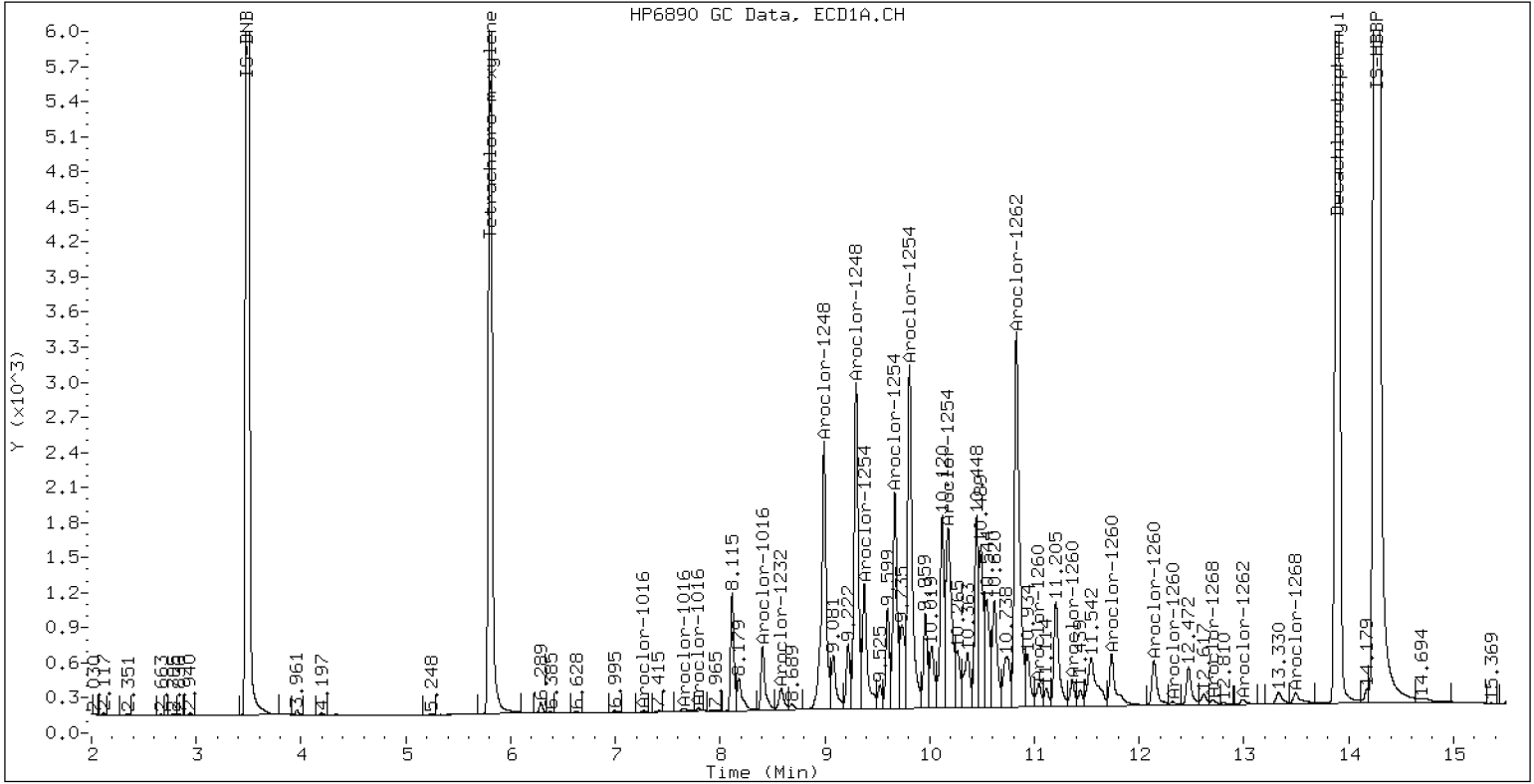
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1254SCV

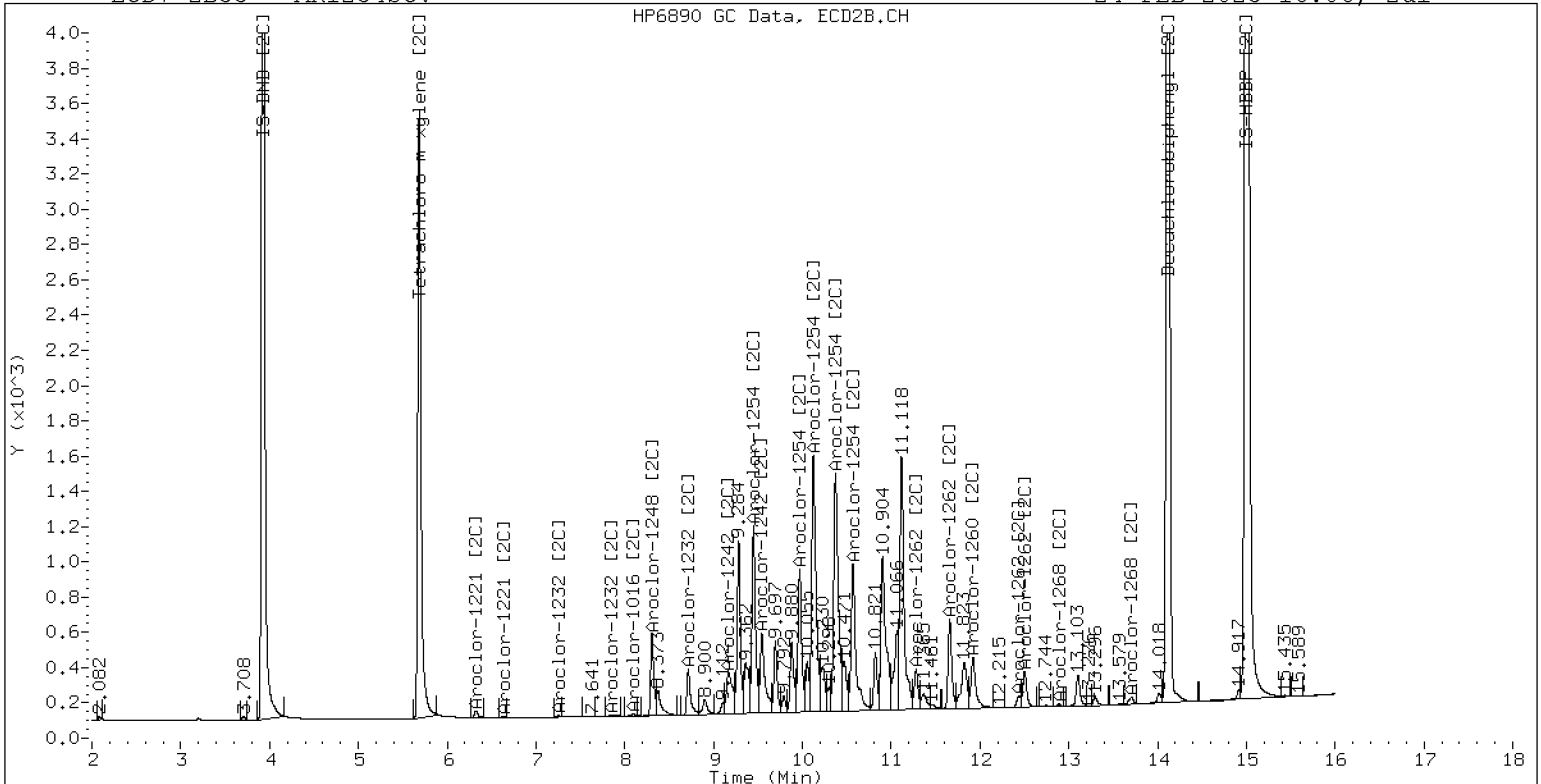
24-FEB-2023 16:06, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1254SCV

24-FEB-2023 16:06, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242317ECD7.D
Data file 2: /230224.b/230224.b/02242317ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR2162SCV
Client ID:
Injection Date: 24-FEB-2023 16:27
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.807	0.000	356001	5.685	0.000	170882	36.0	36.6	1.7	Tetrachloro-m-xylene
13.895	0.002	533971	14.119	0.000	326235	34.4	37.9	9.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	661953	-1.8
Hexabromobiphenyl	1429847	1574993	10.2

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	317807	0.8
Hexabromobiphenyl	513946	565951	10.1

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.269	-0.001	7175	28.5	1	7.256	0.000	3727	20.0	
Aroclor-1016	2	7.659	0.005	12893	16.8	2	7.863	0.007	5834	15.5	
Aroclor-1016	3	7.794	0.004	6936	18.5	3	8.063	0.009	2963	17.4	
Aroclor-1016	4	8.408	0.003	3610	14.9	4	8.308	0.002	2045	15.3	
Total CollAve (4 peaks):				19.7	Total Col2Ave (4 peaks):				17.0	RPD = 14	
Corrected Ave (3 peaks):				16.8	Corrected Ave (3 peaks):				16.1	RPD = 4	
Aroclor-1221	1	4.730	-0.000	15803	266.6	1	4.955	-0.001	7909	262.9	
Aroclor-1221	2	6.131	-0.001	26946	254.1	2	6.296	-0.000	14303	251.2	
Aroclor-1221	3	6.382	-0.000	62477	253.8	3	6.622	0.000	23612	254.7	
Total CollAve (3 peaks):				258.2	Total Col2Ave (3 peaks):				256.3	RPD = 1	
Corrected Ave: < 3 Peaks					Corrected Ave: < 3 Peaks						
Aroclor-1232	1	4.730	0.000	15803	445.6	1	4.955	-0.001	7909	486.4	
Aroclor-1232	2	6.131	0.000	26946	383.1	2	7.256	0.002	3727	46.1	
Aroclor-1232	3	7.659	0.003	12893	40.5	3	7.863	0.002	5834	36.1	
Aroclor-1232	4	8.583	0.003	2684	19.8	4	8.716	0.002	1189	25.6	
Total CollAve (4 peaks):				222.3	Total Col2Ave (4 peaks):				148.5	RPD = 40	
Corrected Ave (3 peaks):				147.8	Corrected Ave (3 peaks):				35.9	RPD = 122*	
Aroclor-1242	1	7.269	-0.001	7175	35.0	1	7.256	0.000	3727	25.2	
Aroclor-1242	2	7.659	0.003	12893	20.7	2	7.863	0.005	5834	18.8	
Aroclor-1242	3	8.408	0.002	3610	18.6	3	9.175	0.008	1082	11.2	
Aroclor-1242	4	8.583	0.004	2684	9.4	4	9.543	-0.054	1390	11.8	
Total CollAve (4 peaks):				20.9	Total Col2Ave (4 peaks):				16.8	RPD = 22	
Corrected Ave (3 peaks):				16.2	Corrected Ave (3 peaks):				13.9	RPD = 15	
Aroclor-1248	1	8.408	0.002	3610	11.2	1	8.308	0.001	2045	13.5	
Aroclor-1248	2	8.583	0.003	2684	6.5	2	8.716	0.002	1189	7.6	
Aroclor-1248	3	8.994	-0.005	24440	31.6	3	9.175	0.009	1082	6.0	
Aroclor-1248	4	9.302	0.008	26328	66.8	4	9.543	-0.048	1390	6.4	
Total CollAve (4 peaks):				29.0	Total Col2Ave (4 peaks):				8.4	RPD = 110*	
Corrected Ave (3 peaks):				16.4	Corrected Ave (3 peaks):				6.7	RPD = 85*	
Aroclor-1254	1	9.302	0.004	26328	39.6	1	9.452	0.003	9571	39.6	
Aroclor-1254	2	---	---	---	0.0	2	9.972	0.002	1733	8.9	
Aroclor-1254	3	9.670	0.002	3721	8.7	3	10.147	0.023	49218	117.1	
Aroclor-1254	4	9.808	0.000	9653	11.6	4	10.370	-0.002	59603	145.4	
Aroclor-1254	5	10.120	-0.056	131179	251.9	5	10.569	0.001	79533	318.7	
Total CollAve (4 peaks):				78.0	Total Col2Ave (5 peaks):				125.9	RPD = 47*	
Corrected Ave (3 peaks):				20.0	Corrected Ave (4 peaks):				77.8	RPD = 118*	
Aroclor-1260	1	11.044	-0.000	223208	394.0	1	11.652	-0.001	104071	312.7	
Aroclor-1260	2	11.361	-0.001	190166	321.2	2	11.919	0.002	251579	296.2	
Aroclor-1260	3	11.737	0.003	458281	291.9	3	12.435	-0.001	113645	504.2	
Aroclor-1260	4	12.141	0.002	149720	189.4	4	12.501	-0.001	182951	319.6	
Aroclor-1260	5	12.244	0.000	196033	576.0	NS	---	---	---	---	
Total CollAve (5 peaks):				354.5	Total Col2Ave (4 peaks):				358.2	RPD = 1	
Corrected Ave (4 peaks):				299.1	Corrected Ave (3 peaks):				309.5	RPD = 3	
Aroclor-1262	1	10.828	-0.001	121431	251.3	1	11.201	0.000	121335	251.1	
Aroclor-1262	2	12.244	0.000	196033	249.3	2	11.652	0.000	104071	252.9	
Aroclor-1262	3	12.319	0.001	211092	249.8	3	12.435	0.001	113645	243.4	
Aroclor-1262	4	12.988	0.001	183455	237.5	4	12.501	-0.001	182951	250.1	
Total CollAve (4 peaks):				247.0	Total Col2Ave (4 peaks):				249.3	RPD = 1	
Corrected Ave (3 peaks):				245.5	Corrected Ave (3 peaks):				248.2	RPD = 1	
Aroclor-1268	1	12.244	-0.002	196033	97.1	1	12.435	0.003	113645	99.7	
Aroclor-1268	2	12.319	0.002	211092	105.6	2	12.501	0.001	182951	149.3	
Aroclor-1268	3	12.723	0.024	77240	45.2	3	12.891	-0.000	7755	7.4	
Aroclor-1268	4	13.488	-0.002	65479	11.6	4	13.709	0.000	35146	10.5	
Total CollAve (4 peaks):				64.9	Total Col2Ave (4 peaks):				66.7	RPD = 3	

Corrected Ave (3 peaks): 51.3 Corrected Ave (3 peaks): 39.2 RPD = 27

Total PCB Area Col1 (5.906 - 13.793) = 3239932 Col1 Total PCB = 0.4 ppm*
Total PCB Area Col2 (5.785 - 14.019) = 1655522 Col2 Total PCB = 0.4 ppm*

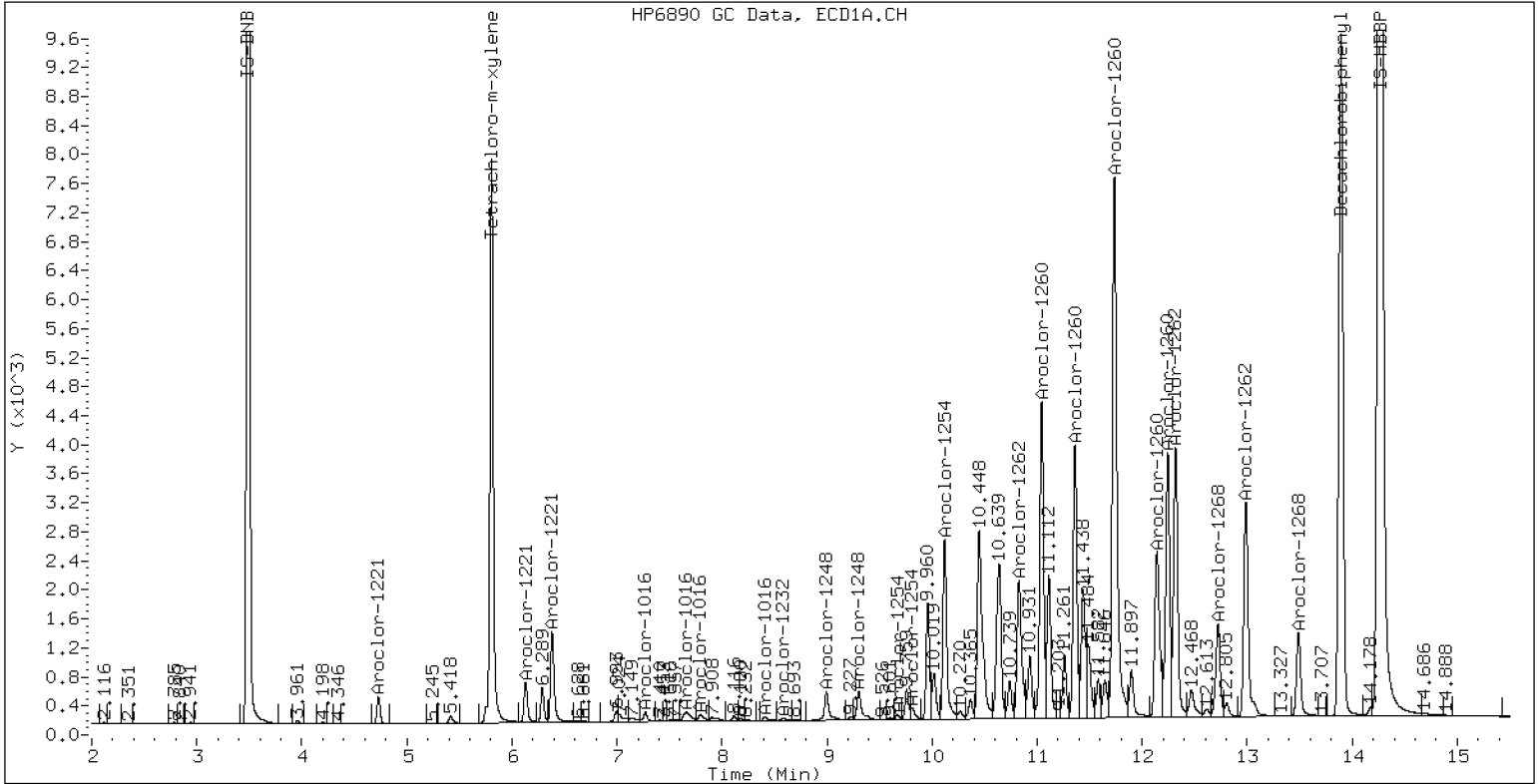
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR2162SCV

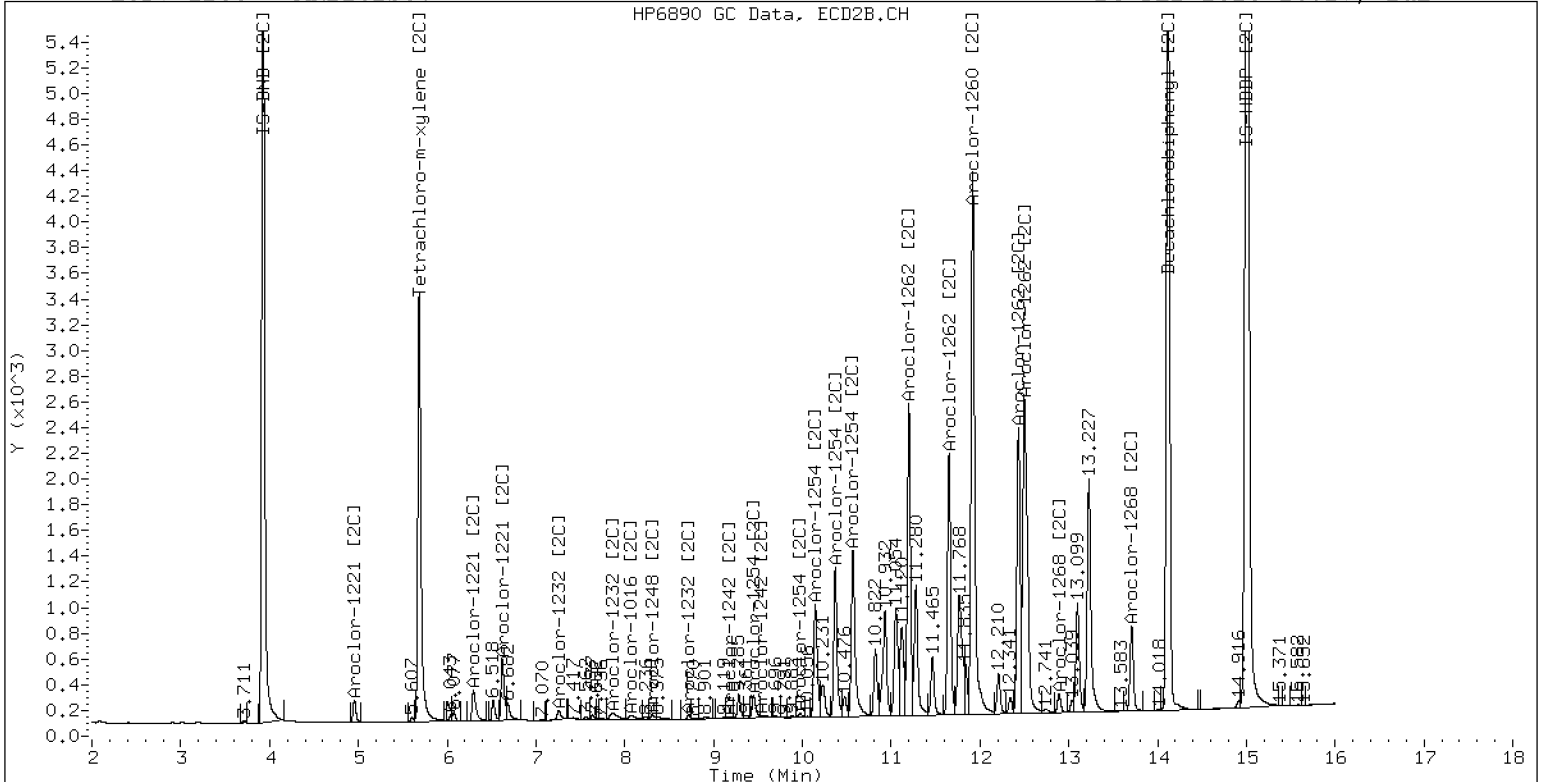
24-FEB-2023 16:27, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR2162SCV

24-FEB-2023 16:27, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230224.b/02242318ECD7.D
Data file 2: /230224.b/230224.b/02242318ECD7.D
Method: \\target\share\chem4\ecd7.i\230224.b\PCB.m
Compound Sublist: PCB.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR3268SCV
Client ID:
Injection Date: 24-FEB-2023 16:48
Report Date: 02/28/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.806	0.000	363331	5.685	0.000	176204	37.1	38.2	2.9	Tetrachloro-m-xylene
13.894	0.001	800845	14.118	-0.001	488290	51.3	56.4	9.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	656592	-2.6
Hexabromobiphenyl	1429847	1584453	10.8

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	314741	-0.2
Hexabromobiphenyl	513946	568346	10.6

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.270	-0.001	28327	113.6	1	7.254	-0.001	20651	112.1	
Aroclor-1016	2	7.657	0.003	80668	106.1	2	7.861	0.005	41326	110.6	
Aroclor-1016	3	7.793	0.003	40661	109.6	3	8.060	0.005	20446	121.2	
Aroclor-1016	4	8.407	0.002	24680	102.9	4	8.308	0.001	13576	102.5	
Total CollAve (4 peaks):				108.0	Total Col2Ave (4 peaks):				111.6	RPD = 3	
Corrected Ave (3 peaks):				106.2	Corrected Ave (3 peaks):				108.4	RPD = 2	
Aroclor-1221	1	4.729	-0.001	8535	145.1	1	4.956	-0.000	3965	133.1	
Aroclor-1221	2	6.132	-0.000	15523	147.6	2	6.297	0.001	8689	154.1	
Aroclor-1221	3	6.382	-0.000	45872	187.9	3	6.622	0.001	22272	242.6	
Total CollAve (3 peaks):				160.2	Total Col2Ave (3 peaks):				176.6	RPD = 10	
Corrected Ave: < 3 Peaks					Corrected Ave: < 3 Peaks						
Aroclor-1232	1	4.729	-0.001	8535	242.6	1	4.956	0.000	3965	246.2	
Aroclor-1232	2	6.132	0.001	15523	222.5	2	7.254	0.000	20651	258.1	
Aroclor-1232	3	7.657	0.001	80668	255.4	3	7.861	0.001	41326	258.3	
Aroclor-1232	4	8.582	0.001	34784	259.2	4	8.714	-0.001	12504	271.5	
Total CollAve (4 peaks):				244.9	Total Col2Ave (4 peaks):				258.5	RPD = 5	
Corrected Ave (3 peaks):				240.2	Corrected Ave (3 peaks):				254.2	RPD = 6	
Aroclor-1242	1	7.270	-0.001	28327	139.2	1	7.254	-0.001	20651	141.2	
Aroclor-1242	2	7.657	0.001	80668	130.5	2	7.861	0.003	41326	134.4	
Aroclor-1242	3	8.407	0.001	24680	128.4	3	9.170	0.003	12830	134.1	
Aroclor-1242	4	8.582	0.003	34784	122.4	4	9.600	0.003	14836	127.3	
Total CollAve (4 peaks):				130.1	Total Col2Ave (4 peaks):				134.3	RPD = 3	
Corrected Ave (3 peaks):				127.1	Corrected Ave (3 peaks):				132.0	RPD = 4	
Aroclor-1248	1	8.407	0.001	24680	77.0	1	8.308	0.000	13576	90.3	
Aroclor-1248	2	8.582	0.001	34784	85.4	2	8.714	-0.000	12504	80.5	
Aroclor-1248	3	8.996	-0.003	83592	108.8	3	9.170	0.004	12830	71.8	
Aroclor-1248	4	9.292	-0.003	39603	101.3	4	9.600	0.010	14836	69.1	
Total CollAve (4 peaks):				93.1	Total Col2Ave (4 peaks):				77.9	RPD = 18	
Corrected Ave (3 peaks):				87.9	Corrected Ave (3 peaks):				73.8	RPD = 17	
Aroclor-1254	1	9.292	-0.007	39603	60.1	1	9.452	0.003	4590	19.2	
Aroclor-1254	2	9.377	-0.000	11450	38.6	2	9.973	0.003	2892	15.0	
Aroclor-1254	3	9.674	0.005	6387	15.1	3	10.131	0.007	6052	14.5	
Aroclor-1254	4	9.813	0.006	10162	12.3	4	10.390	0.017	5324	13.1	
Aroclor-1254	5	10.189	0.012	6862	13.3	5	10.572	0.004	1891	7.7	
Total CollAve (5 peaks):				27.9	Total Col2Ave (5 peaks):				13.9	RPD = 67*	
Corrected Ave (4 peaks):				19.8	Corrected Ave (4 peaks):				12.6	RPD = 45*	
Aroclor-1260	1	11.046	0.002	87033	152.7	1	11.645	-0.008	62543	187.1	
Aroclor-1260	2	11.362	0.001	6300	10.6	2	11.920	0.003	28552	33.5	
Aroclor-1260	3	11.738	0.004	54524	34.5	3	12.432	-0.004	285450	1261.2	
Aroclor-1260	4	12.144	0.005	1727	2.2	4	12.499	-0.002	306992	534.0	
Aroclor-1260	5	12.246	0.002	502931	1469.0	NS	---			----	
Total CollAve (5 peaks):				333.8	Total Col2Ave (4 peaks):				503.9	RPD = 41*	
Corrected Ave (4 peaks):				50.0	Corrected Ave (3 peaks):				251.5	RPD = 134*	
Aroclor-1262	1	10.832	0.004	3395	7.0	1	11.201	0.001	44255	91.2	
Aroclor-1262	2	12.246	0.002	502931	635.9	2	11.645	-0.007	62543	151.3	
Aroclor-1262	3	12.318	-0.000	497006	584.5	3	12.432	-0.002	285450	608.7	
Aroclor-1262	4	12.987	-0.000	202197	260.2	4	12.499	-0.003	306992	417.9	
Total CollAve (4 peaks):				371.9	Total Col2Ave (4 peaks):				317.3	RPD = 16	
Corrected Ave (3 peaks):				283.9	Corrected Ave (3 peaks):				220.1	RPD = 25	
Aroclor-1268	1	12.246	-0.001	502931	247.7	1	12.432	-0.000	285450	249.4	
Aroclor-1268	2	12.318	0.002	497006	247.2	2	12.499	-0.001	306992	249.5	
Aroclor-1268	3	12.699	-0.000	422793	245.8	3	12.892	0.000	260893	248.4	
Aroclor-1268	4	13.490	0.000	1386953	244.9	4	13.709	-0.000	829733	247.1	
Total CollAve (4 peaks):				246.4	Total Col2Ave (4 peaks):				248.6	RPD = 1	

Corrected Ave (3 peaks): 246.0 Corrected Ave (3 peaks): 248.3 RPD = 1

Total PCB Area Col1 (5.906 - 13.793) = 4180607 Col1 Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.785 - 14.019) = 2376912 Col2 Total PCB = 0.6 ppm*

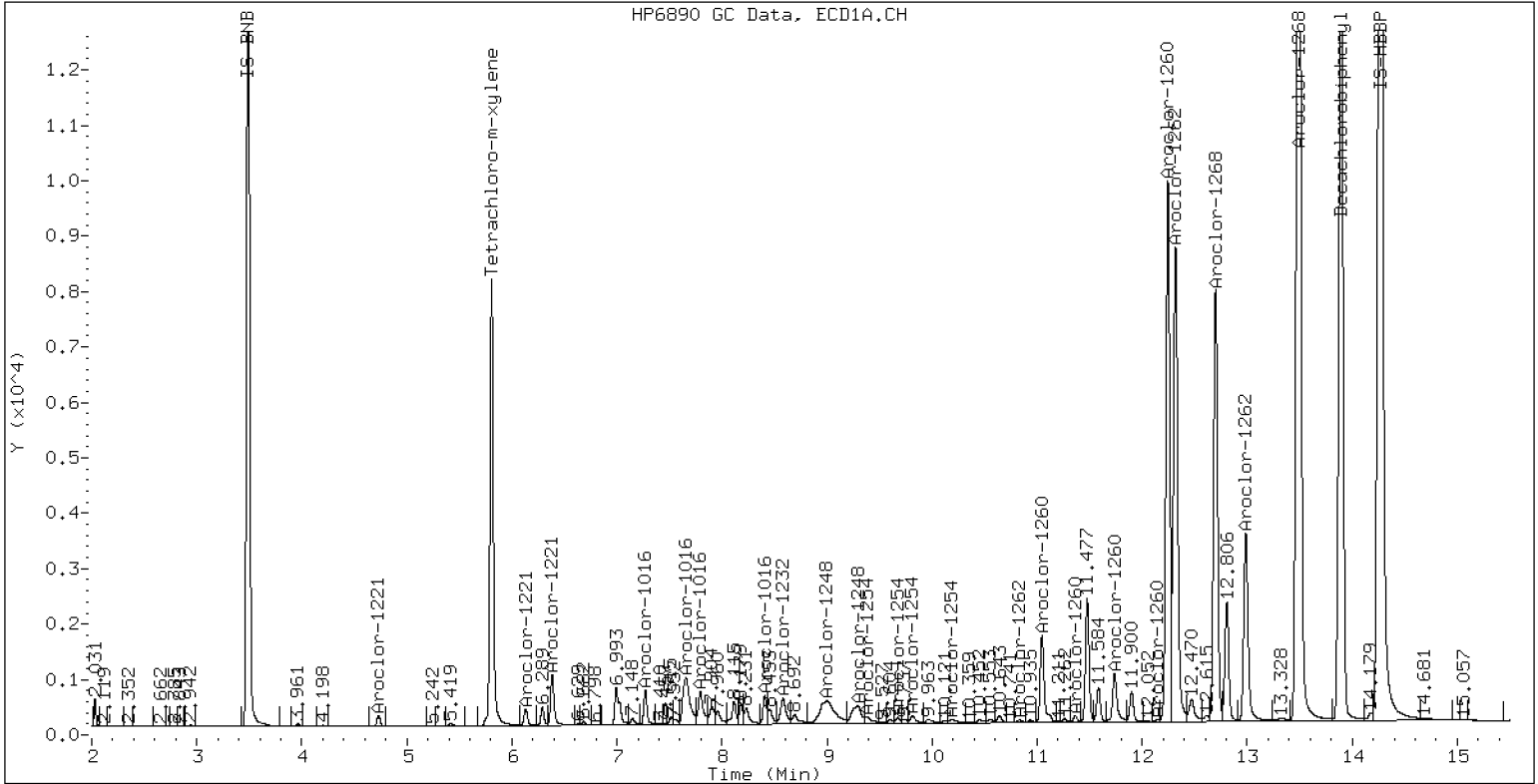
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR3268SCV

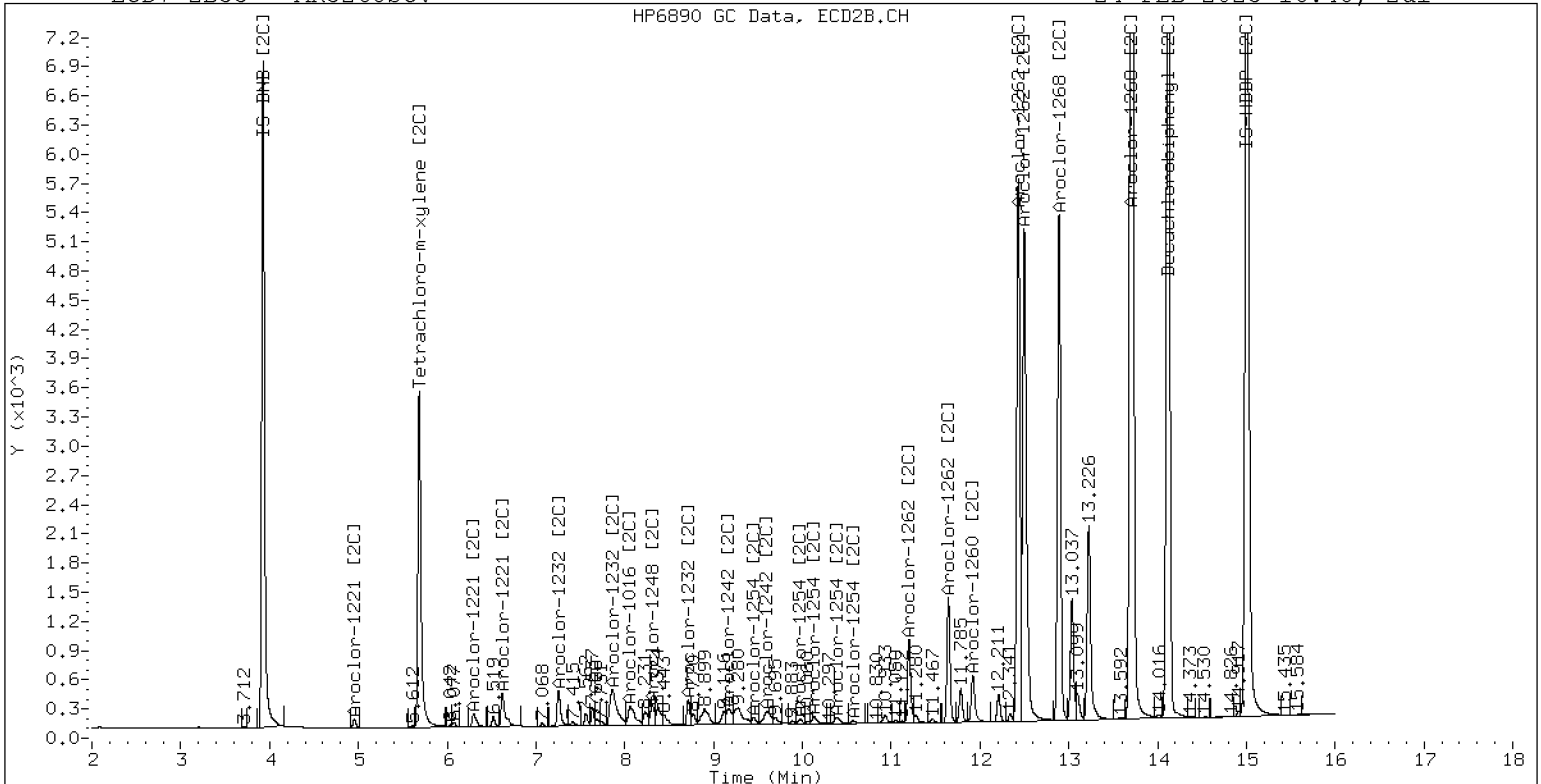
24-FEB-2023 16:48, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR3268SCV

24-FEB-2023 16:48, 2ul



ZB-35 Manual Integration: NO

Analytical Resources Inc.
8082 DDT SCREEN REPORT

Data file 1: /230224.b/02242319ECD7.D

ARI ID: DDTS

RT	ZB5 Col Shift Response		ZB35 Col Shift Response		ZB5 on col	ZB35 on col	RPD	Compound/Flag	
9.261	0.000	694353	9.912	0.000	580269	0.100	0.100	0.0	2,4-DDE
0.000	-10.293	0	10.672	0.000	673479	0.000	0.200#	----	2,4-DDT
9.686	0.000	1191406	10.212	0.000	433373	0.100	0.100	0.0	4,4-DDE
10.259	0.000	1721760	10.672	0.000	673479	0.100	0.200#	66.7*	4,4-DDD

Indicates value is from co-eluting peaks

* Indicates RPD > 40%

Analytical Resources Inc.
8082 DDT SCREEN REPORT

Data file 1: /230224.b/02242320ECD7.D

ARI ID: DDT BD

RT	ZB5 Col Shift Response	ZB35 Col Shift Response	RT	ZB5 on col	ZB35 on col	RPD	Compound/Flag		
9.285	0.023	4923	9.921	0.009	9972	0.001	0.002	84.3*	2,4-DDE
0.000	-10.293	0	10.677	0.004	249094	0.000	0.074#	----	2,4-DDT
9.692	0.006	12128	10.221	0.009	528	0.001	0.000	156.7*	4,4-DDE
10.265	0.006	410017	10.677	0.004	249094	0.023	0.074#	103.6*	4,4-DDD

Indicates value is from co-eluting peaks

* Indicates RPD > 40%



SECOND-SOURCE CALIBRATION VERIFICATION
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: GB00069

Laboratory ID: SLB0342-SCV1

Sequence: SLB0342

Sequence Name: AR1660SCV1

Standard ID: L002065

ANALYTE	EXPECTED (ug/L)	FOUND (ug/L)	% DRIFT	QC LIMIT
Aroclor 1016	250.00	243	-2.7	20.00
Aroclor 1016 [2C]	250.00	246	-1.4	20.00
Aroclor 1260	250.00	266	6.2	20.00
Aroclor 1260 [2C]	250.00	261	4.5	20.00
Decachlorobiphenyl	40.000	34.3	-14.2	20.00
Tetrachlorometaxylene	40.000	34.9	-12.6	20.00
Decachlorobiphenyl [2C]	40.000	37.3	-6.6	20.00
Tetrachlorometaxylene [2C]	40.000	35.8	-10.6	20.00

* Indicates values outside of QC limits
[2C] indicates second-column analyte.



SECOND-SOURCE CALIBRATION VERIFICATION
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: GB00069

Laboratory ID: SLB0342-SCV2

Sequence: SLB0342

Sequence Name: AR1242SCV2

Standard ID: K007656

ANALYTE	EXPECTED (ug/L)	FOUND (ug/L)	% DRIFT	QC LIMIT
Aroclor 1242	250.00	204	-18.5	20.00
Aroclor 1242 [2C]	250.00	221	-11.8	20.00
Decachlorobiphenyl	40.000	37.0	-7.5	20.00
Tetrachlorometaxylene	40.000	33.6	-15.9	20.00
Decachlorobiphenyl [2C]	40.000	40.3	0.8	20.00
Tetrachlorometaxylene [2C]	40.000	34.5	-13.7	20.00

* Indicates values outside of QC limits
[2C] indicates second-column analyte.



SECOND-SOURCE CALIBRATION VERIFICATION
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: GB00069

Laboratory ID: SLB0342-SCV3

Sequence: SLB0342

Sequence Name: AR1248SCV3

Standard ID: L002066

ANALYTE	EXPECTED (ug/L)	FOUND (ug/L)	% DRIFT	QC LIMIT
Aroclor 1248	250.00	249	-0.2	20.00
Aroclor 1248 [2C]	250.00	248	-0.8	20.00
Decachlorobiphenyl	40.000	33.1	-17.2	20.00
Tetrachlorometaxylene	40.000	34.9	-12.8	20.00
Decachlorobiphenyl [2C]	40.000	36.3	-9.2	20.00
Tetrachlorometaxylene [2C]	40.000	36.4	-9.0	20.00

* Indicates values outside of QC limits

[2C] indicates second-column analyte.



SECOND-SOURCE CALIBRATION VERIFICATION
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: GB00069

Laboratory ID: SLB0342-SCV4

Sequence: SLB0342

Sequence Name: AR1254SCV4

Standard ID: L002067

ANALYTE	EXPECTED (ug/L)	FOUND (ug/L)	% DRIFT	QC LIMIT
Aroclor 1254	250.00	235	-5.9	20.00
Aroclor 1254 [2C]	250.00	240	-4.0	20.00
Decachlorobiphenyl	40.000	34.6	-13.4	20.00
Tetrachlorometaxylene	40.000	36.1	-9.7	20.00
Decachlorobiphenyl [2C]	40.000	37.9	-5.2	20.00
Tetrachlorometaxylene [2C]	40.000	37.1	-7.3	20.00

* Indicates values outside of QC limits

[2C] indicates second-column analyte.



SECOND-SOURCE CALIBRATION VERIFICATION
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: GB00069

Laboratory ID: SLB0342-SCV5

Sequence: SLB0342

Sequence Name: AR2162SCV5

Standard ID: L002068

ANALYTE	EXPECTED (ug/L)	FOUND (ug/L)	% DRIFT	QC LIMIT
Aroclor 1221	250.00	258	3.3	20.00
Aroclor 1221 [2C]	250.00	256	2.5	20.00
Aroclor 1262	250.00	247	-1.2	20.00
Aroclor 1262 [2C]	250.00	249	-0.3	20.00
Decachlorobiphenyl	40.000	34.4	-13.9	20.00
Tetrachlorometaxylene	40.000	36.0	-10.0	20.00
Decachlorobiphenyl [2C]	40.000	37.9	-5.4	20.00
Tetrachlorometaxylene [2C]	40.000	36.6	-8.4	20.00

* Indicates values outside of QC limits
[2C] indicates second-column analyte.



SECOND-SOURCE CALIBRATION VERIFICATION
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: GB00069

Laboratory ID: SLB0342-SCV6

Sequence: SLB0342

Sequence Name: AR3268SCV6

Standard ID: L002069

ANALYTE	EXPECTED (ug/L)	FOUND (ug/L)	% DRIFT	QC LIMIT
Aroclor 1232	250.00	245	-2.0	20.00
Aroclor 1232 [2C]	250.00	259	3.4	20.00
Aroclor 1268	250.00	246	-1.4	20.00
Aroclor 1268 [2C]	250.00	249	-0.6	20.00
Decachlorobiphenyl	40.000	51.3	28.3	20.00
Tetrachlorometaxylene	40.000	37.1	-7.3	20.00
Decachlorobiphenyl [2C]	40.000	56.4	41.0	20.00
Tetrachlorometaxylene [2C]	40.000	38.2	-4.6	20.00

* Indicates values outside of QC limits
[2C] indicates second-column analyte.



INITIAL CALIBRATION CHECK
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor OEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: ECD7

Calibration: GB00069

Lab File ID: 03152306ECD7.D

Calibration Date: 02/24/2023

Sequence: SLC0215

Injection Date: 03/15/23

Lab Sample ID: SLC0215-ICV1

Injection Time: 14:43

Sequence Name: AR1254ICV1

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR			% DRIFT/DIFF	
		STD	ICV	ICAL	ICV	MIN	ICV	LIMIT
Aroclor 1254	A	250.00	278	0.0662949	0.0735796		11.1	
Aroclor-1254 (1)	A	250.00	264	0.0803331	0.0846742			
Aroclor-1254 (2)	A	250.00	267	0.0361302	0.0386075			
Aroclor-1254 (3)	A	250.00	264	0.0516471	0.0545564			
Aroclor-1254 (4)	A	250.00	271	0.1004230	0.1088147			
Aroclor-1254 (5)	A	250.00	323	0.0629414	0.0812452			
Aroclor 1254 [2C]	A	250.00	242	0.0763106	0.0739290		-3.1	
Aroclor-1254 (1) [2C]	A	250.00	241	0.0608052	0.0587271			
Aroclor-1254 (2) [2C]	A	250.00	241	0.0489162	0.0472448			
Aroclor-1254 (3) [2C]	A	250.00	242	0.1058376	0.1023944			
Aroclor-1254 (4) [2C]	A	250.00	242	0.1031750	0.0997841			
Aroclor-1254 (5) [2C]	A	250.00	245	0.0628191	0.0614945			
Decachlorobiphenyl	A	40.000	30.8	0.7878687	0.6070139		-23.0	
Tetrachlorometaxylene	A	40.000	35.5	1.1944880	1.0593570		-11.3	
Decachlorobiphenyl [2C]	A	40.000	37.5	1.2182710	1.1409820		-6.3	
Tetrachlorometaxylene [2C]	A	40.000	35.7	1.1737210	1.0466760		-10.8	

* Values outside of QC limits

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152306ECD7.D
Data file 2: /230315.b/230315.b/03152306ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1254.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1254ICV1
Client ID:
Injection Date: 15-MAR-2023 14:43
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.810	-0.001	523127	5.692	-0.001	188744	35.5	35.7	0.5	Tetrachloro-m-xylene
13.898	0.001	757146	14.120	-0.001	305511	30.8	37.5	19.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		%D
	Standard Area*	Sample Area	
Bromo-Nitrobenzene	673778	987631	46.6
Hexabromobiphenyl	1429847	2494658	74.5

Standard Cpnd	Column 2		%D
	Standard Area*	Sample Area	
Bromo-Nitrobenzene	315256	360654	14.4
Hexabromobiphenyl	513946	535523	4.2

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1254	1	9.304	0.005	261334	263.5	1	9.457	-0.001	66188	241.5	
Aroclor-1254	2	9.385	0.007	119156	267.1	2	9.978	0.000	53247	241.5	
Aroclor-1254	3	9.676	0.008	168380	264.1	3	10.134	0.000	115403	241.9	
Aroclor-1254	4	9.817	0.009	335840	270.9	4	10.382	-0.000	112461	241.8	
Aroclor-1254	5	10.189	0.013	250751	322.7	5	10.577	-0.001	69307	244.7	
Total CollAve (5 peaks):				277.7	Total Col2Ave (5 peaks):				242.3	RPD = 14	
Corrected Ave (4 peaks):				266.4	Corrected Ave (4 peaks):				241.6	RPD = 10	
CalAmt %D:				11.1	CalAmt %D:				-3.1		

Total PCB Area Col1 (5.911 - 13.797) = 3476528 Col1 Total PCB = 0.3 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 1119243 Col2 Total PCB = 0.3 ppm*

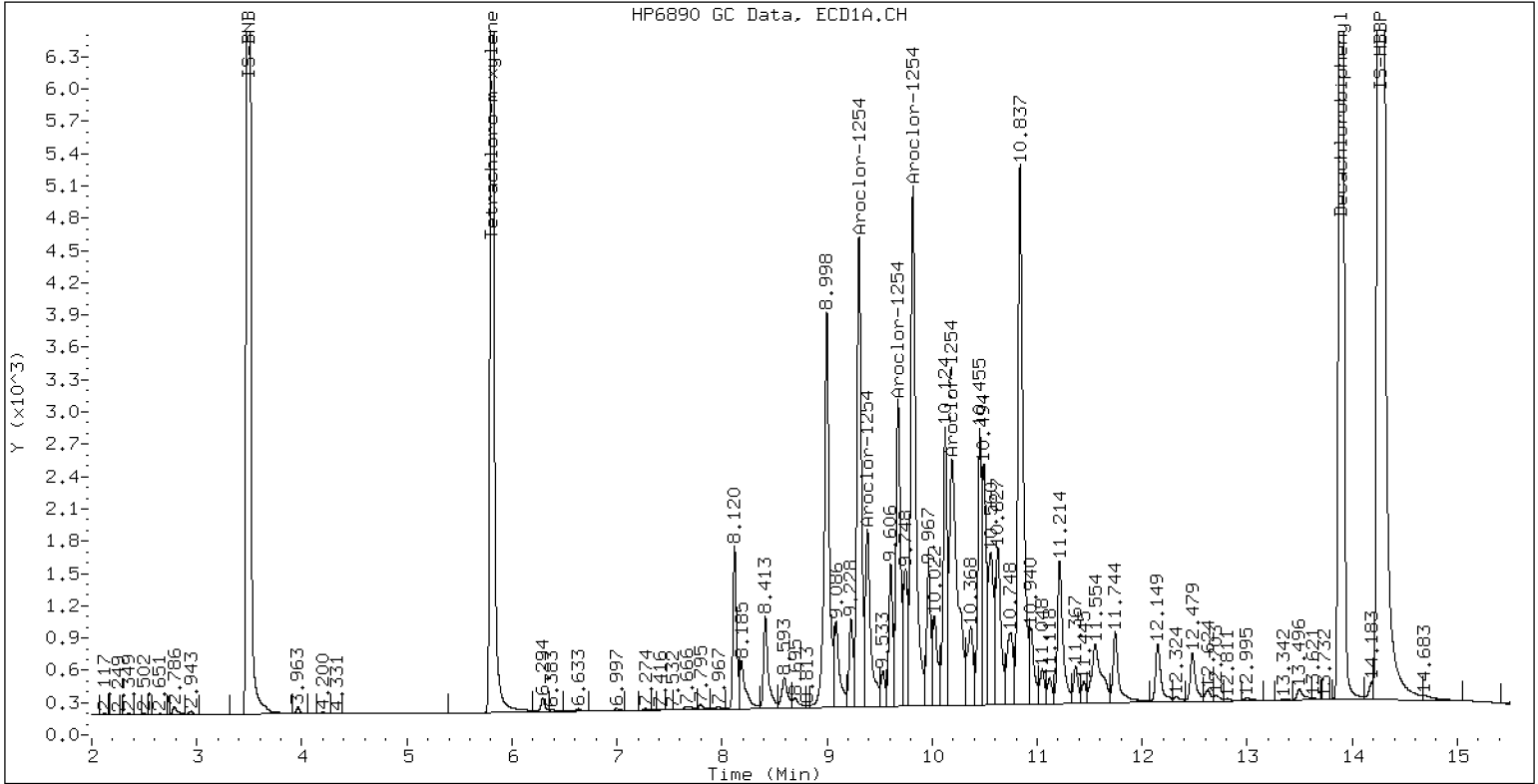
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1254ICV1

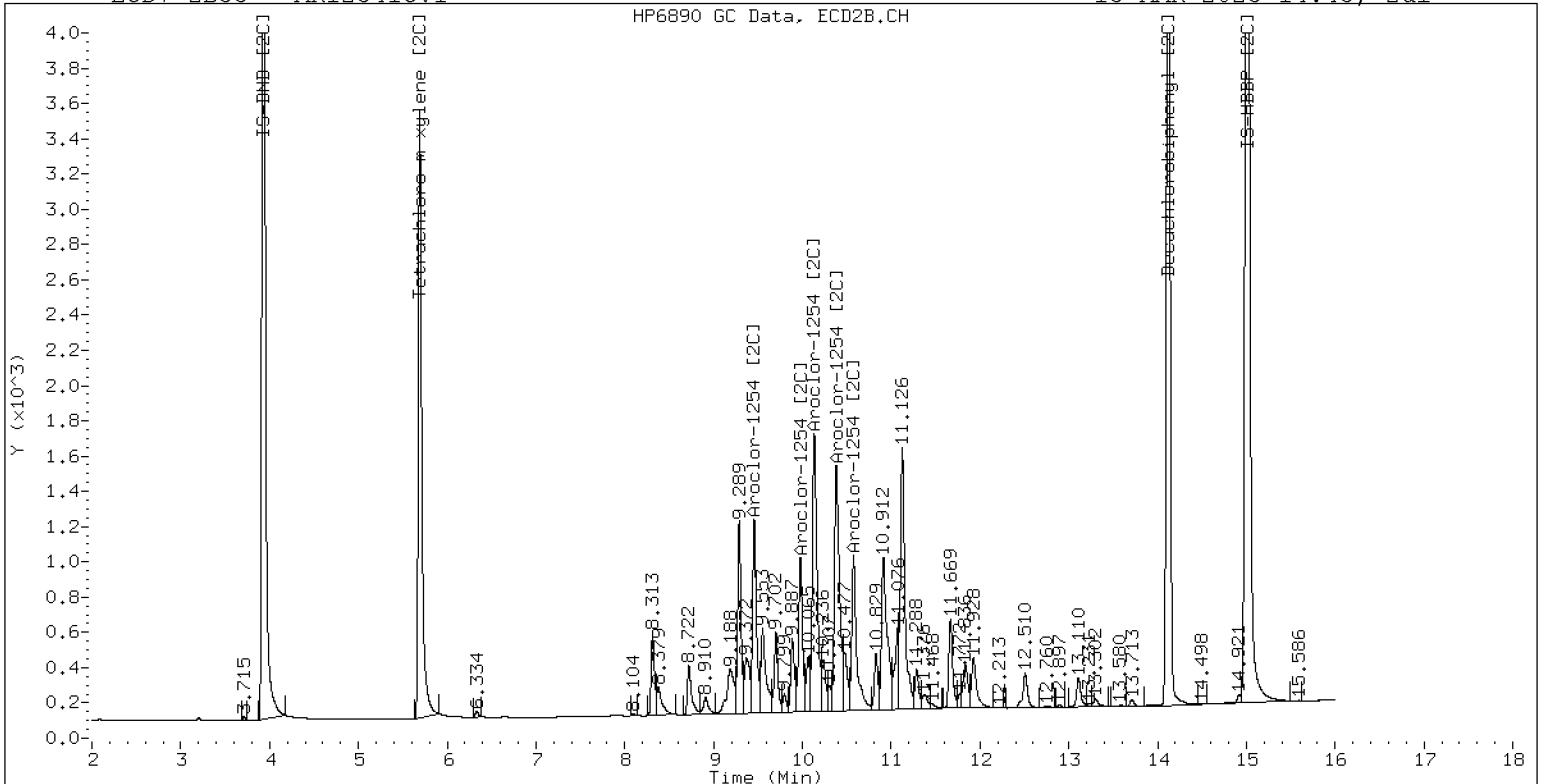
15-MAR-2023 14:43, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1254ICV1

15-MAR-2023 14:43, 2ul



ZB-35 Manual Integration: NO



INITIAL CALIBRATION CHECK
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: ECD7

Calibration: GB00069

Lab File ID: 03152307ECD7.D

Calibration Date: 02/24/2023

Sequence: SLC0215

Injection Date: 03/15/23

Lab Sample ID: SLC0215-ICV2

Injection Time: 15:04

Sequence Name: AR1660ICV2

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR			% DRIFT/DIFF	
		STD	ICV	ICAL	ICV	MIN	ICV	LIMIT
Aroclor 1016	A	250.00	246	0.0493662	0.0484376		-1.6	
Aroclor-1016 (1)	A	250.00	240	0.0303852	0.0291161		-4.0	
Aroclor-1016 (2)	A	250.00	240	0.0926308	0.0889754		-4.0	
Aroclor-1016 (3)	A	250.00	262	0.0452180	0.0473193		4.8	
Aroclor-1016 (4)	A	250.00	242	0.0292307	0.0283397		-3.2	
Aroclor 1016 [2C]	A	250.00	248	0.0545857	0.0537049		-0.9	
Aroclor-1016 (1) [2C]	A	250.00	228	0.0468313	0.0427876		-8.8	
Aroclor-1016 (2) [2C]	A	250.00	241	0.0949676	0.0916476		-3.6	
Aroclor-1016 (3) [2C]	A	250.00	274	0.0428922	0.0470415		9.6	
Aroclor-1016 (4) [2C]	A	250.00	248	0.0336515	0.0333430		-0.8	
Aroclor 1260	A	250.00	224	0.0392091	0.0349117		-10.4	
Aroclor-1260 (1)	A	250.00	234	0.0287785	0.0269957		-6.4	
Aroclor-1260 (2)	A	250.00	227	0.0300690	0.0273171		-9.2	
Aroclor-1260 (3)	A	250.00	219	0.0797517	0.0697586		-12.4	
Aroclor-1260 (4)	A	250.00	219	0.0401599	0.0351835		-12.4	
Aroclor-1260 (5)	A	250.00	221	0.0172866	0.0153035		-11.6	
Aroclor 1260 [2C]	A	250.00	251	0.0699688	0.0712829		0.3	
Aroclor-1260 (1) [2C]	A	250.00	257	0.0470406	0.0483227		2.8	
Aroclor-1260 (2) [2C]	A	250.00	260	0.1200523	0.1247405		4.0	
Aroclor-1260 (3) [2C]	A	250.00	230	0.0318590	0.0292543		-8.0	
Aroclor-1260 (4) [2C]	A	250.00	256	0.0809231	0.0828142		2.4	
Decachlorobiphenyl	A	40.000	31.5	0.7878687	0.6207508		-21.3	
Tetrachlorometaxylene	A	40.000	36.6	1.1944880	1.0920760		-8.5	
Decachlorobiphenyl [2C]	A	40.000	38.0	1.2182710	1.1582140		-5.0	
Tetrachlorometaxylene [2C]	A	40.000	36.6	1.1737210	1.0750530		-8.5	

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152307ECD7.D
Data file 2: /230315.b/230315.b/03152307ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1660ICV2
Client ID:
Injection Date: 15-MAR-2023 15:04
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.810	-0.001	547342	5.692	0.000	196074	36.6	36.6	0.2	Tetrachloro-m-xylene
13.896	-0.001	815056	14.122	0.001	318744	31.5	38.0	18.7	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1002388	48.8
Hexabromobiphenyl	1429847	2626033	83.7

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	364771	15.7
Hexabromobiphenyl	513946	550406	7.1

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.273	0.003	91205	239.6	1	7.259	0.001	48774	228.4	
Aroclor-1016	2	7.665	0.011	278712	240.1	2	7.871	-0.001	104470	241.3	
Aroclor-1016	3	7.798	0.008	148226	261.6	3	8.069	-0.000	53623	274.2	
Aroclor-1016	4	8.412	0.007	88773	242.4	4	8.314	0.001	38008	247.7	
Total CollAve (4 peaks):				245.9		Total Col2Ave (4 peaks):				247.9	RPD = 1
Corrected Ave (3 peaks):				240.7		Corrected Ave (3 peaks):				239.1	RPD = 1

CalAmt %D: -1.6

CalAmt %D: -0.8

Aroclor-1260	1	11.050	0.005	221536	234.5	1	11.658	0.000	83116	256.8	
Aroclor-1260	2	11.367	0.006	224174	227.1	2	11.925	0.000	214556	259.8	
Aroclor-1260	3	11.743	0.009	572464	218.7	3	12.441	0.000	50318	229.6	
Aroclor-1260	4	12.148	0.009	288728	219.0	4	12.509	0.001	142442	255.8	
Aroclor-1260	5	12.249	0.005	125586	221.3	NS	---			----	
Total CollAve (5 peaks):				224.1		Total Col2Ave (4 peaks):				250.5	RPD = 11
Corrected Ave (4 peaks):				221.5		Corrected Ave (3 peaks):				247.4	RPD = 11

CalAmt %D: -10.3

CalAmt %D: 0.2

Total PCB Area Col1 (5.911 - 13.797) = 6074384 Col1 Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 2011382 Col2 Total PCB = 0.5 ppm*

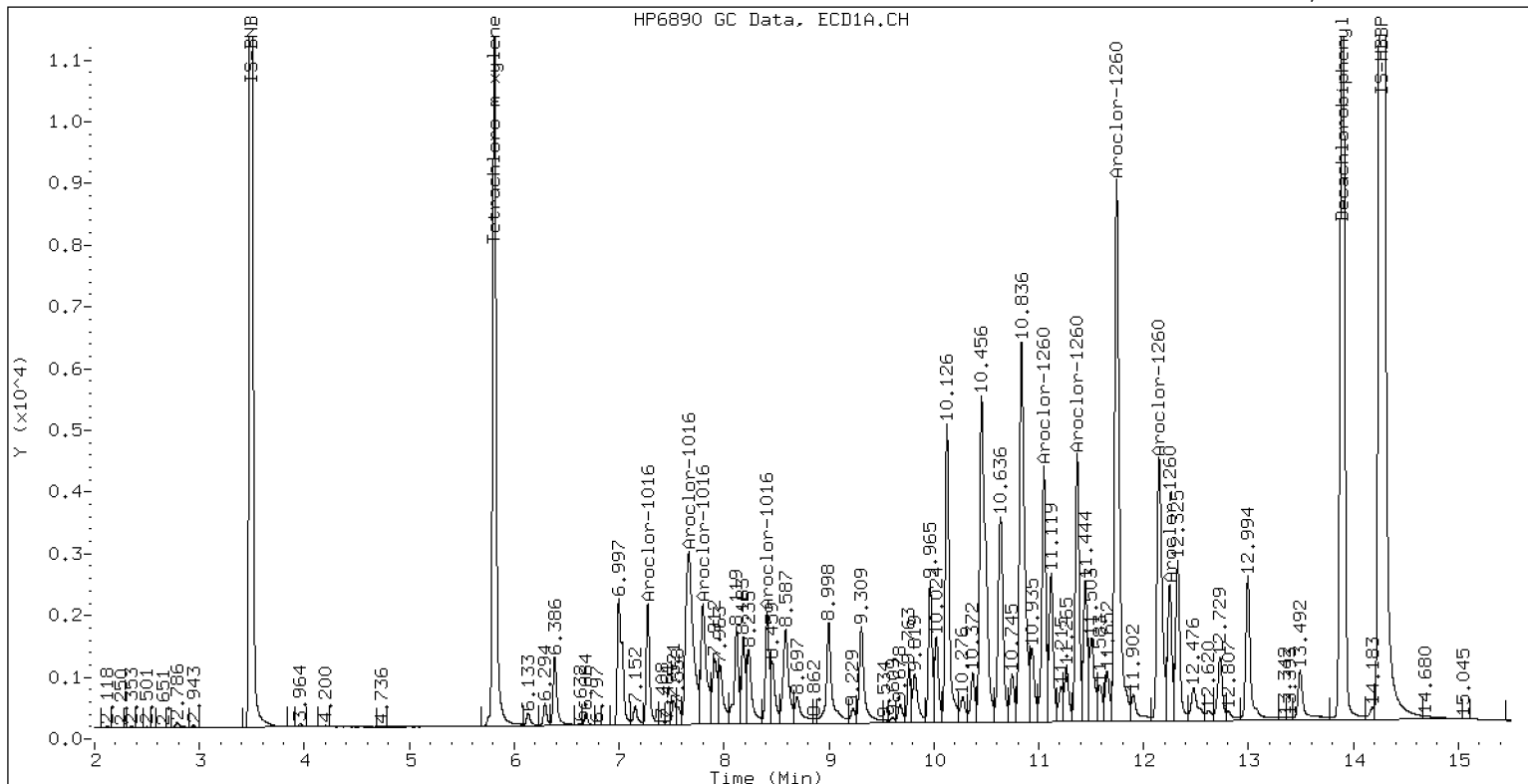
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1660ICV2

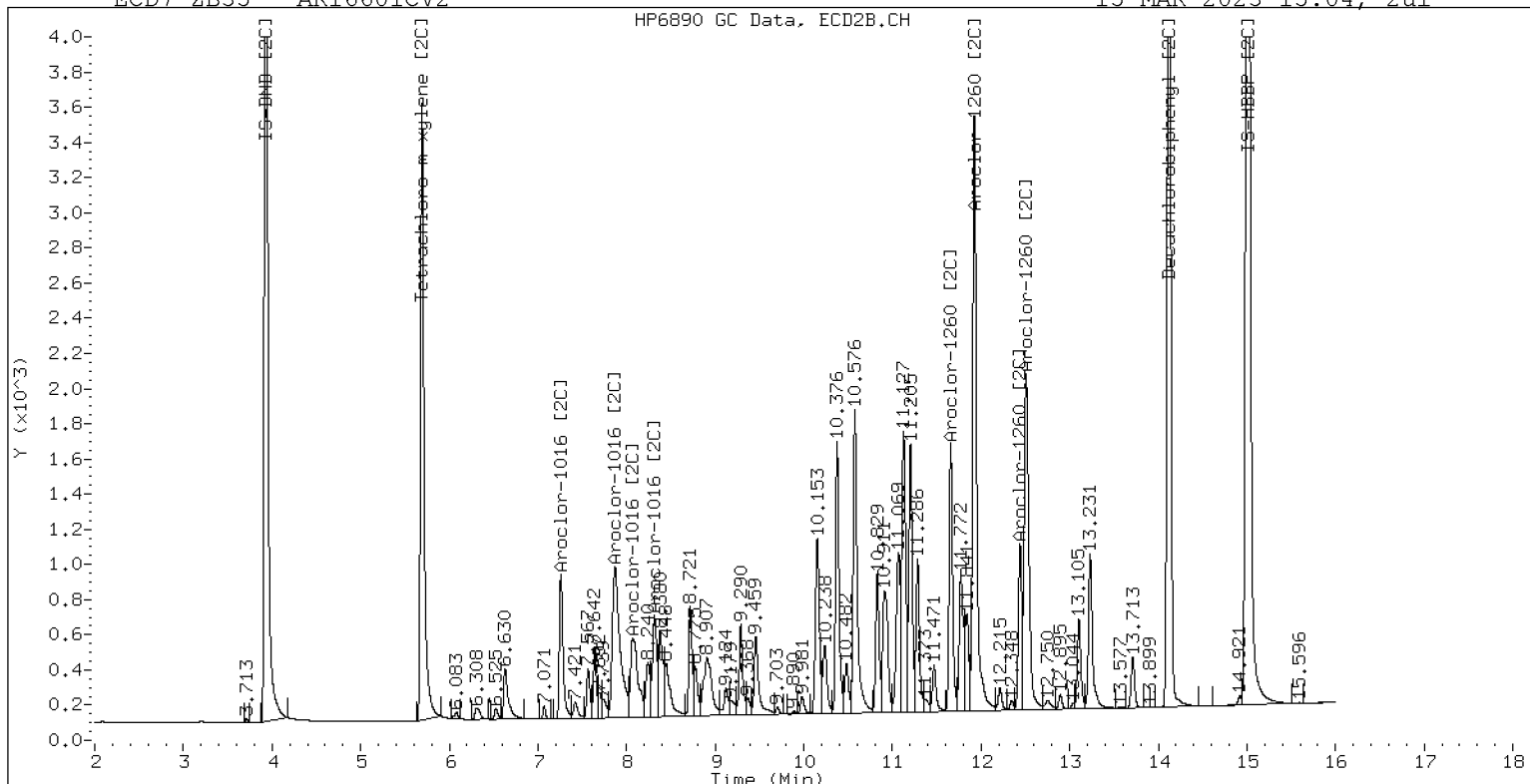
15-MAR-2023 15:04, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1660ICV2

15-MAR-2023 15:04, 2ul



ZB-35 Manual Integration: NO



**SECOND-SOURCE
CONTINUING CALIBRATION CHECK
EPA 8082A**

Laboratory: <u>Analytical Resources, LLC</u>	SDG: <u>23C0107</u>
Client: <u>Anchor QEA, LLC</u>	Project: <u>AOC4 UR Phase 3</u>
Instrument ID: <u>ECD7</u>	Calibration: <u>GB00069</u>
Lab File ID: <u>02242313ECD7.D</u>	Calibration Date: <u>02/24/2023</u>
Sequence: <u>SLB0342</u>	Injection Date: <u>02/24/23</u>
Lab Sample ID: <u>SLB0342-SCV1</u>	Injection Time: <u>15:03</u>
Sequence Name: <u>AR1660SCV1</u>	

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1016	A	250.00	243	0.0493662	0.0479314		-2.7	+/-20
Aroclor 1016 [2C]	A	250.00	246	0.0545857	0.0542382		-1.4	+/-20
Aroclor 1260	A	250.00	266	0.0392091	0.0412121		6.2	+/-20
Aroclor 1260 [2C]	A	250.00	261	0.0699688	0.0733659		4.5	+/-20
Decachlorobiphenyl	A	40.000	34.3	0.7878687	0.6762784		-14.2	+/-20
Tetrachlorometaxylene	A	40.000	34.9	1.1944880	1.0436010		-12.6	+/-20
Decachlorobiphenyl [2C]	A	40.000	37.3	1.2182710	1.1373730		-6.6	+/-20
Tetrachlorometaxylene [2C]	A	40.000	35.8	1.1737210	1.0492890		-10.6	+/-20

* Values outside of QC limits

* Values outside of QC limits



**SECOND-SOURCE
CONTINUING CALIBRATION CHECK
EPA 8082A**

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>02242314ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLB0342</u>	Injection Date:	<u>02/24/23</u>
Lab Sample ID:	<u>SLB0342-SCV2</u>	Injection Time:	<u>15:24</u>
Sequence Name:	<u>AR1242SCV2</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1242	A	250.00	204	0.0395340	0.0322254		-18.5	+/-20
Aroclor 1242 [2C]	A	250.00	221	0.0423092	0.0365983		-11.8	+/-20
Decachlorobiphenyl	A	40.000	37.0	0.7878687	0.7290534		-7.5	+/-20
Tetrachlorometaxylene	A	40.000	33.6	1.1944880	1.0041320		-15.9	+/-20
Decachlorobiphenyl [2C]	A	40.000	40.3	1.2182710	1.2285170		0.8	+/-20
Tetrachlorometaxylene [2C]	A	40.000	34.5	1.1737210	1.0131510		-13.7	+/-20

* Values outside of QC limits

* Values outside of QC limits



**SECOND-SOURCE
CONTINUING CALIBRATION CHECK
EPA 8082A**

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>02242315ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLB0342</u>	Injection Date:	<u>02/24/23</u>
Lab Sample ID:	<u>SLB0342-SCV3</u>	Injection Time:	<u>15:45</u>
Sequence Name:	<u>AR1248SCV3</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1248	A	250.00	249	0.0574755	0.0572474		-0.2	+/-20
Aroclor 1248 [2C]	A	250.00	248	0.0444270	0.0440936		-0.8	+/-20
Decachlorobiphenyl	A	40.000	33.1	0.7878687	0.6527336		-17.2	+/-20
Tetrachlorometaxylene	A	40.000	34.9	1.1944880	1.0413820		-12.8	+/-20
Decachlorobiphenyl [2C]	A	40.000	36.3	1.2182710	1.1066400		-9.2	+/-20
Tetrachlorometaxylene [2C]	A	40.000	36.4	1.1737210	1.0676190		-9.0	+/-20

* Values outside of QC limits



**SECOND-SOURCE
CONTINUING CALIBRATION CHECK
EPA 8082A**

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>02242316ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLB0342</u>	Injection Date:	<u>02/24/23</u>
Lab Sample ID:	<u>SLB0342-SCV4</u>	Injection Time:	<u>16:06</u>
Sequence Name:	<u>AR1254SCV4</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1254	A	250.00	235	0.0662949	0.0622529		-5.9	+/-20
Aroclor 1254 [2C]	A	250.00	240	0.0763106	0.0731447		-4.0	+/-20
Decachlorobiphenyl	A	40.000	34.6	0.7878687	0.6823832		-13.4	+/-20
Tetrachlorometaxylene	A	40.000	36.1	1.1944880	1.0787610		-9.7	+/-20
Decachlorobiphenyl [2C]	A	40.000	37.9	1.2182710	1.1548440		-5.2	+/-20
Tetrachlorometaxylene [2C]	A	40.000	37.1	1.1737210	1.0880920		-7.3	+/-20

* Values outside of QC limits

* Values outside of QC limits



**SECOND-SOURCE
CONTINUING CALIBRATION CHECK
EPA 8082A**

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>02242317ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLB0342</u>	Injection Date:	<u>02/24/23</u>
Lab Sample ID:	<u>SLB0342-SCV5</u>	Injection Time:	<u>16:27</u>
Sequence Name:	<u>AR2162SCV5</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1221	A	250.00	258	0.0165758	0.0169561		3.3	+/-20
Aroclor 1221 [2C]	A	250.00	256	0.0150798	0.0153801		2.5	+/-20
Aroclor 1262	A	250.00	247	0.0366596	0.0361658		-1.2	+/-20
Aroclor 1262 [2C]	A	250.00	249	0.0739760	0.0737876		-0.3	+/-20
Decachlorobiphenyl	A	40.000	34.4	0.7878687	0.6780614		-13.9	+/-20
Tetrachlorometaxylene	A	40.000	36.0	1.1944880	1.0756080		-10.0	+/-20
Decachlorobiphenyl [2C]	A	40.000	37.9	1.2182710	1.1528740		-5.4	+/-20
Tetrachlorometaxylene [2C]	A	40.000	36.6	1.1737210	1.0753820		-8.4	+/-20

* Values outside of QC limits

* Values outside of QC limits



**SECOND-SOURCE
CONTINUING CALIBRATION CHECK
EPA 8082A**

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>02242318ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLB0342</u>	Injection Date:	<u>02/24/23</u>
Lab Sample ID:	<u>SLB0342-SCV6</u>	Injection Time:	<u>16:48</u>
Sequence Name:	<u>AR3268SCV6</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1232	A	250.00	245	0.0169039	0.0169981		-2.0	+/-20
Aroclor 1232 [2C]	A	250.00	259	0.0192023	0.0199392		3.4	+/-20
Aroclor 1268	A	250.00	246	0.1442124	0.1418626		-1.4	+/-20
Aroclor 1268 [2C]	A	250.00	249	0.2386862	0.2369075		-0.6	+/-20
Decachlorobiphenyl	A	40.000	51.3	0.7878687	1.0108790		28.3	+/-20
Tetrachlorometaxylene	A	40.000	37.1	1.1944880	1.1067180		-7.3	+/-20
Decachlorobiphenyl [2C]	A	40.000	56.4	1.2182710	1.7182840		41.0	+/-20
Tetrachlorometaxylene [2C]	A	40.000	38.2	1.1737210	1.1196760		-4.6	+/-20

* Values outside of QC limits



CONTINUING CALIBRATION CHECK
EPA 8082A

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>03152313ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLC0215</u>	Injection Date:	<u>03/15/23</u>
Lab Sample ID:	<u>SLC0215-CCV1</u>	Injection Time:	<u>17:09</u>
Sequence Name:	<u>AR1248CCV1</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1248	A	250.00	268	0.0574755	0.0623427		7.2	
Aroclor-1248 (1)	A	250.00	249		0.0388222			
Aroclor-1248 (2)	A	250.00	253		0.0501795			
Aroclor-1248 (3)	A	250.00	280		0.1050406			
Aroclor-1248 (4)	A	250.00	290		0.0553285			
Aroclor 1248 [2C]	A	250.00	241	0.0444270	0.0427369		-3.7	
Aroclor-1248 (1) [2C]	A	250.00	243		0.0371097			
Aroclor-1248 (2) [2C]	A	250.00	237		0.0374862			
Aroclor-1248 (3) [2C]	A	250.00	248		0.0451214			
Aroclor-1248 (4) [2C]	A	250.00	235		0.0512302			
Decachlorobiphenyl	A	40.000	28.7	0.7878687	0.5650591		-28.3	
Tetrachlorometaxylene	A	40.000	35.1	1.1944880	1.0475430		-12.3	
Decachlorobiphenyl [2C]	A	40.000	37.9	1.2182710	1.1537860		-5.3	
Tetrachlorometaxylene [2C]	A	40.000	36.0	1.1737210	1.0557330		-10.0	

* Values outside of QC limits

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152313ECD7.D
Data file 2: /230315.b/230315.b/03152313ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1248.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1248CCV1
Client ID:
Injection Date: 15-MAR-2023 17:09
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	0.000	521016	5.693	0.001	185706	35.1	36.0	2.5	Tetrachloro-m-xylene
13.897	0.000	836924	14.123	0.002	327643	28.7	37.9	27.6	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	994739	47.6
Hexabromobiphenyl	1429847	2962253	107.2 <-

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	351805	11.6
Hexabromobiphenyl	513946	567944	10.5

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1248	1	8.411	0.000	120681	248.7	1	8.315	0.000	40798	242.9	
Aroclor-1248	2	8.589	0.000	155986	252.8	2	8.722	0.000	41212	237.3	
Aroclor-1248	3	8.998	0.000	326525	280.6	3	9.185	0.000	49606	248.2	
Aroclor-1248	4	9.302	0.000	171992	290.3	4	9.612	0.000	56322	234.7	
Total CollAve (4 peaks):				268.1		Total Col2Ave (4 peaks):				240.8	RPD = 11
Corrected Ave (3 peaks):				260.7		Corrected Ave (3 peaks):				238.3	RPD = 9
CalAmt %D:				7.2		CalAmt %D:				-3.7	

Total PCB Area Col1 (5.911 - 13.797) = 2501360 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 818626 Col2 Total PCB = 0.2 ppm*

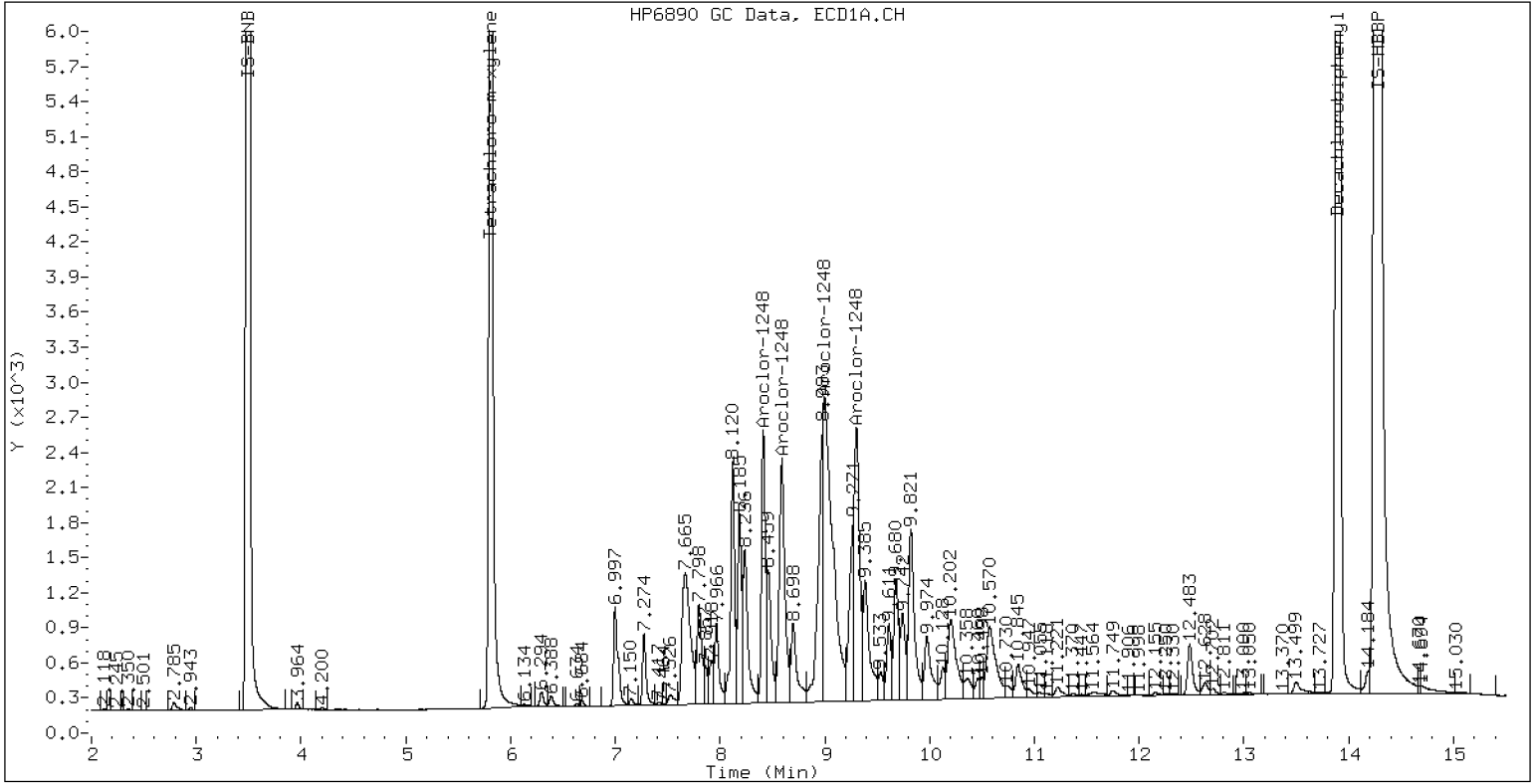
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

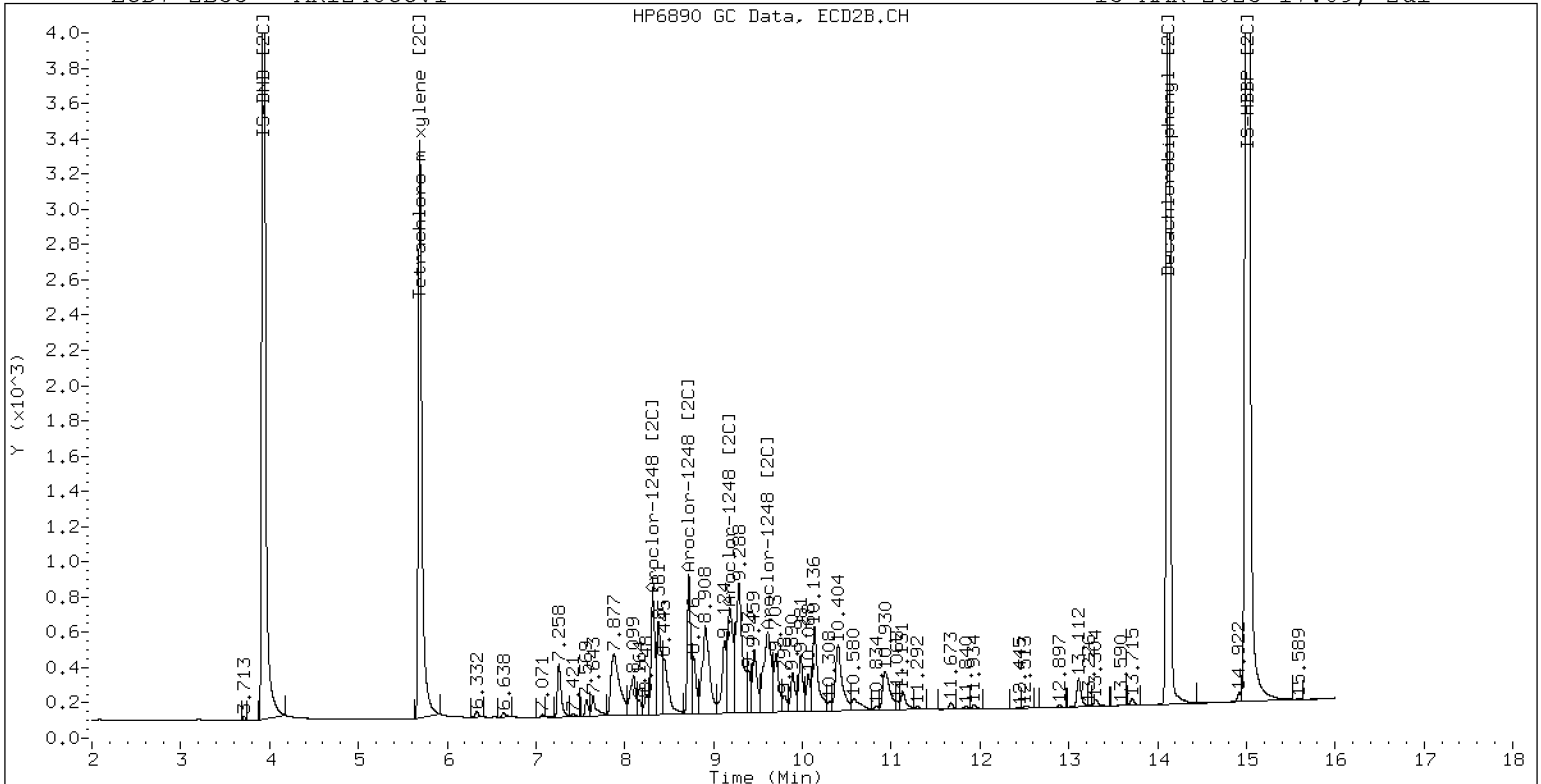
ECD7-ZB5 AR1248CCV1

15-MAR-2023 17:09, 2ul



ECD7-ZB35 AR1248CCV1

15-MAR-2023 17:09, 2ul

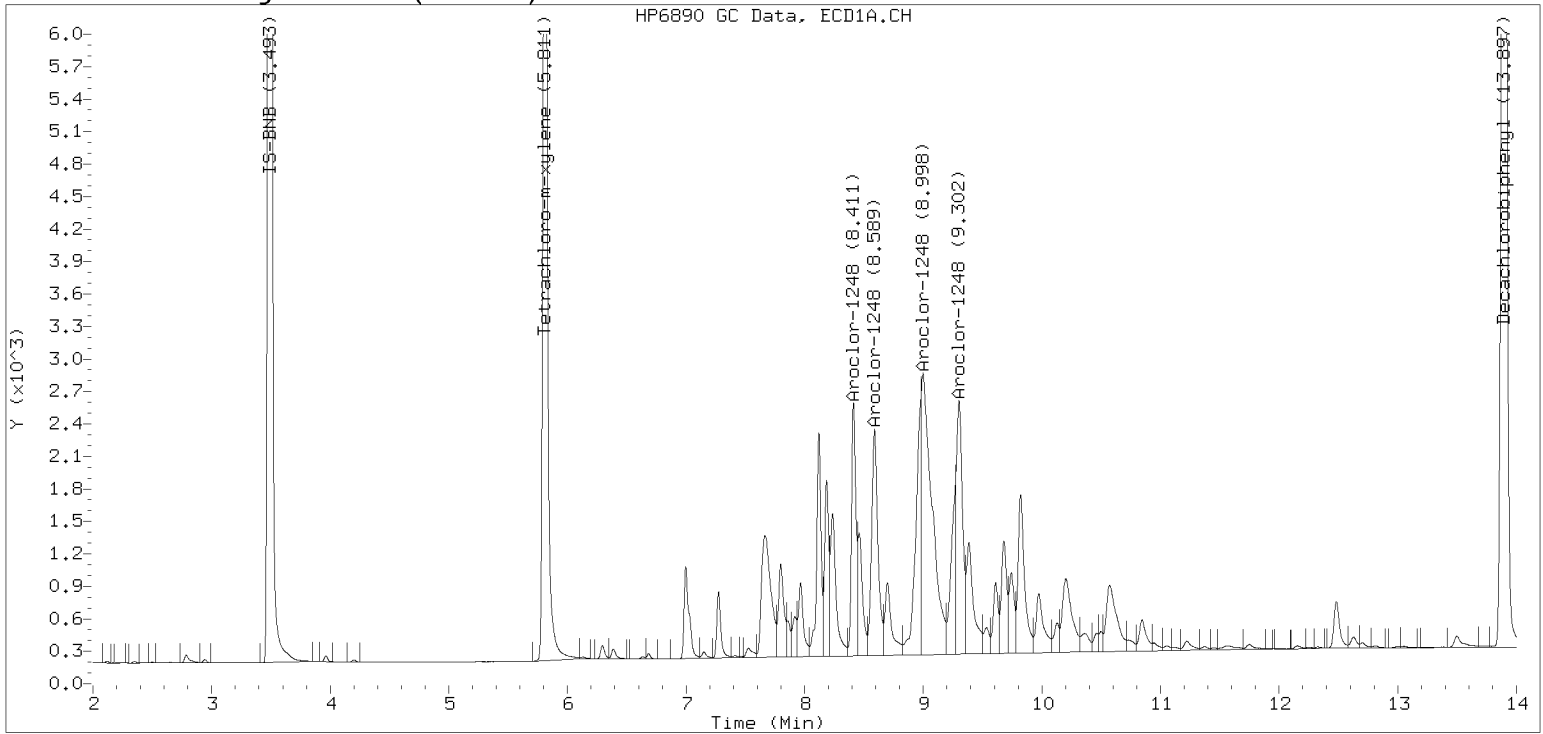


Manual Peak Adjustment, ZB-5

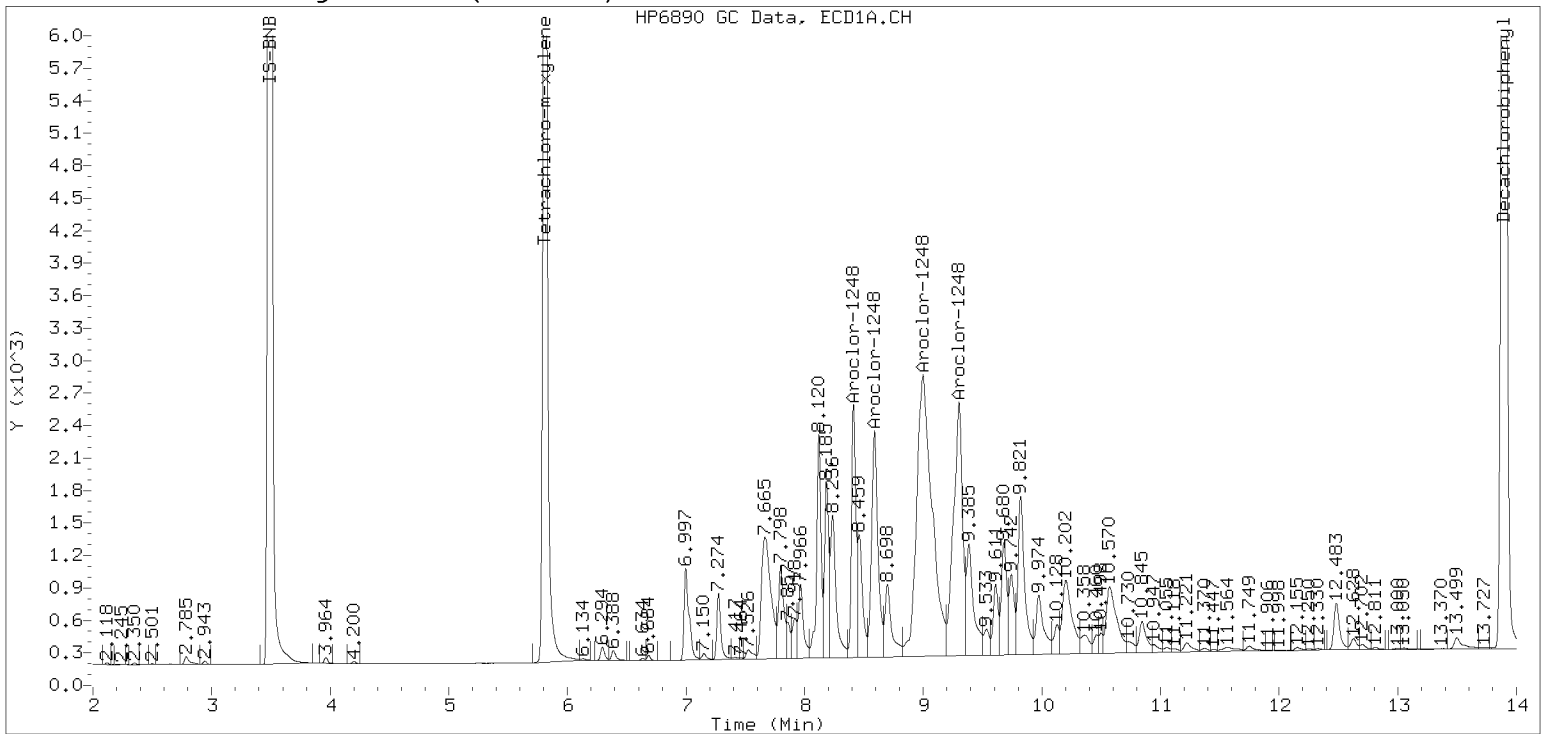
Datafile: ecd7.i/230315.b/03152313ECD7.D

Injection Date: 15-MAR-2023 17:09

Manual Integration (After)



Processed Integration (Before)





CONTINUING CALIBRATION CHECK
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: ECD7

Calibration: GB00069

Lab File ID: 03152314ECD7.D

Calibration Date: 02/24/2023

Sequence: SLC0215

Injection Date: 03/15/23

Lab Sample ID: SLC0215-CCV2

Injection Time: 17:30

Sequence Name: AR1660CCV2

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1016	A	250.00	241	0.0493662	0.0472976		-3.7	
Aroclor-1016 (1)	A	250.00	235	0.0303852	0.0285751		-6.0	
Aroclor-1016 (2)	A	250.00	233	0.0926308	0.0862735		-6.8	
Aroclor-1016 (3)	A	250.00	258	0.0452180	0.0466187		3.2	
Aroclor-1016 (4)	A	250.00	237	0.0292307	0.0277233		-5.2	
Aroclor 1016 [2C]	A	250.00	246	0.0545857	0.0531343		-1.7	
Aroclor-1016 (1) [2C]	A	250.00	225	0.0468313	0.0422156		-10.0	
Aroclor-1016 (2) [2C]	A	250.00	238	0.0949676	0.0902447		-4.8	
Aroclor-1016 (3) [2C]	A	250.00	274	0.0428922	0.0469445		9.6	
Aroclor-1016 (4) [2C]	A	250.00	246	0.0336515	0.0331325		-1.6	
Aroclor 1260	A	250.00	203	0.0392091	0.0317575		-19.0	
Aroclor-1260 (1)	A	250.00	209	0.0287785	0.0240518		-16.4	
Aroclor-1260 (2)	A	250.00	205	0.0300690	0.0246626		-18.0	
Aroclor-1260 (3)	A	250.00	202	0.0797517	0.0645661		-19.2	
Aroclor-1260 (4)	A	250.00	197	0.0401599	0.0316730		-21.2	
Aroclor-1260 (5)	A	250.00	200	0.0172866	0.0138340		-20.0	
Aroclor 1260 [2C]	A	250.00	238	0.0699688	0.0676874		-5.0	
Aroclor-1260 (1) [2C]	A	250.00	243	0.0470406	0.0456724		-2.8	
Aroclor-1260 (2) [2C]	A	250.00	246	0.1200523	0.1184016		-1.6	
Aroclor-1260 (3) [2C]	A	250.00	217	0.0318590	0.0276458		-13.2	
Aroclor-1260 (4) [2C]	A	250.00	244	0.0809231	0.0790299		-2.4	
Decachlorobiphenyl	A	40.000	28.6	0.7878687	0.5631173		-28.5	
Tetrachlorometaxylene	A	40.000	35.6	1.1944880	1.0640570		-11.0	
Decachlorobiphenyl [2C]	A	40.000	38.5	1.2182710	1.1730990		-3.8	
Tetrachlorometaxylene [2C]	A	40.000	36.1	1.1737210	1.0597080		-9.8	

* Values outside of QC limits

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152314ECD7.D
Data file 2: /230315.b/230315.b/03152314ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1660CCV2
Client ID:
Injection Date: 15-MAR-2023 17:30
Report Date: 03/16/2023 09:50
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	-0.000	557813	5.692	0.000	195087	35.6	36.1	1.3	Tetrachloro-m-xylene
13.897	-0.000	878626	14.123	0.002	346391	28.6	38.5	29.6	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1048465	55.6
Hexabromobiphenyl	1429847	3120579	118.2 <-

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	368190	16.8
Hexabromobiphenyl	513946	590557	14.9

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.274	0.003	93625	235.1	1	7.259	0.000	48573	225.4	
Aroclor-1016	2	7.667	0.013	282671	232.8	2	7.873	0.001	103835	237.6	
Aroclor-1016	3	7.799	0.009	152744	257.7	3	8.074	0.005	54014	273.6	
Aroclor-1016	4	8.411	0.006	90834	237.1	4	8.316	0.002	38122	246.1	
Total CollAve (4 peaks):				240.7		Total Col2Ave (4 peaks):				245.7	RPD = 2
Corrected Ave (3 peaks):				235.0		Corrected Ave (3 peaks):				236.4	RPD = 1

CalAmt %D: -3.7

CalAmt %D: -1.7

Aroclor-1260	1	11.050	0.006	234549	208.9	1	11.659	0.001	84288	242.7	
Aroclor-1260	2	11.367	0.006	240505	205.1	2	11.926	0.000	218509	246.6	
Aroclor-1260	3	11.745	0.011	629636	202.4	3	12.441	0.000	51020	216.9	
Aroclor-1260	4	12.150	0.011	308869	197.2	4	12.510	0.002	145849	244.2	
Aroclor-1260	5	12.250	0.006	134906	200.1	NS	---			----	
Total CollAve (5 peaks):				202.7		Total Col2Ave (4 peaks):				237.6	RPD = 16
Corrected Ave (4 peaks):				201.2		Corrected Ave (3 peaks):				234.6	RPD = 15

CalAmt %D: -18.9

CalAmt %D: -5.0

Total PCB Area Coll (5.911 - 13.797) = 6319137 Coll Total PCB = 0.5 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 2033687 Col2 Total PCB = 0.5 ppm*

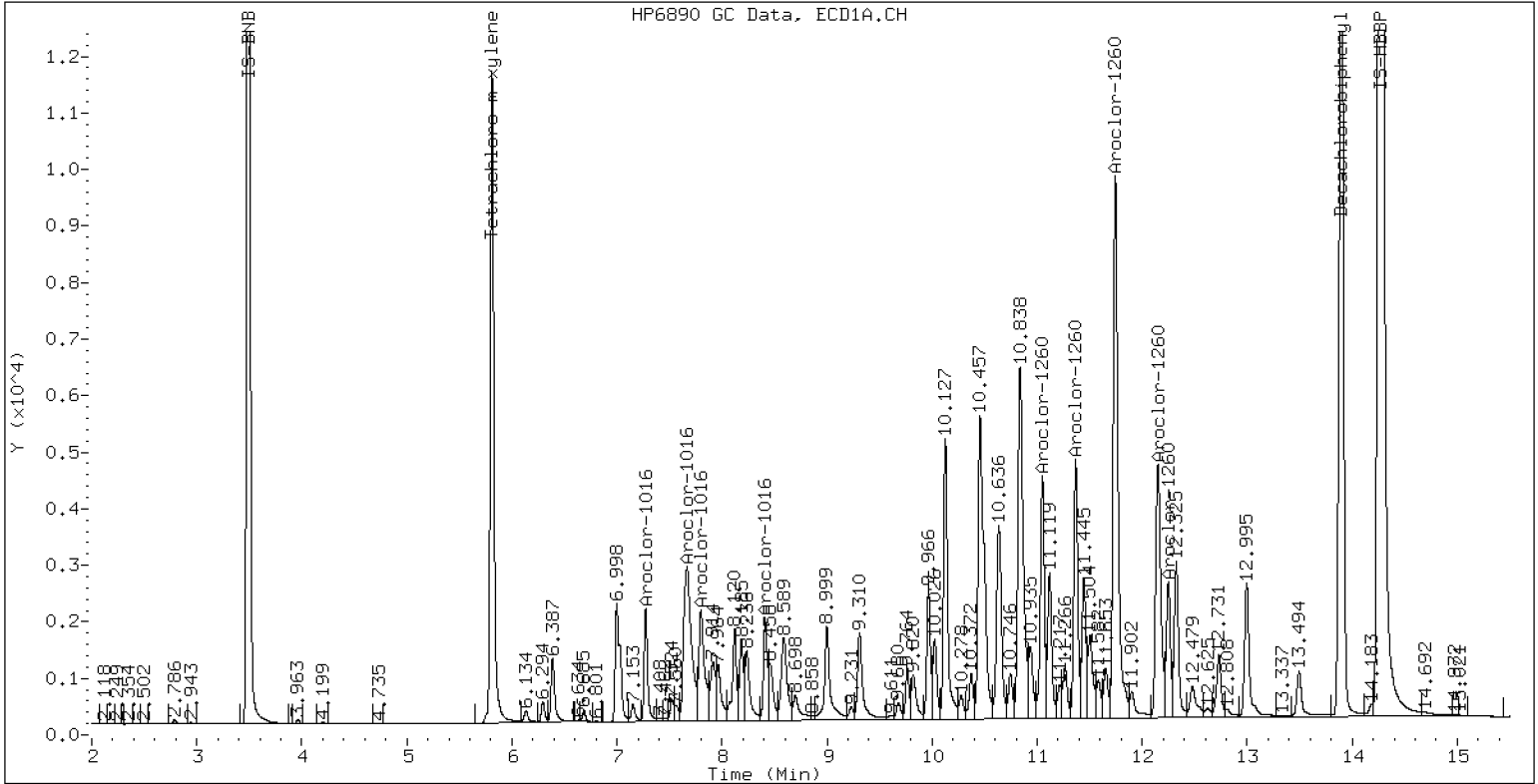
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1660CCV2

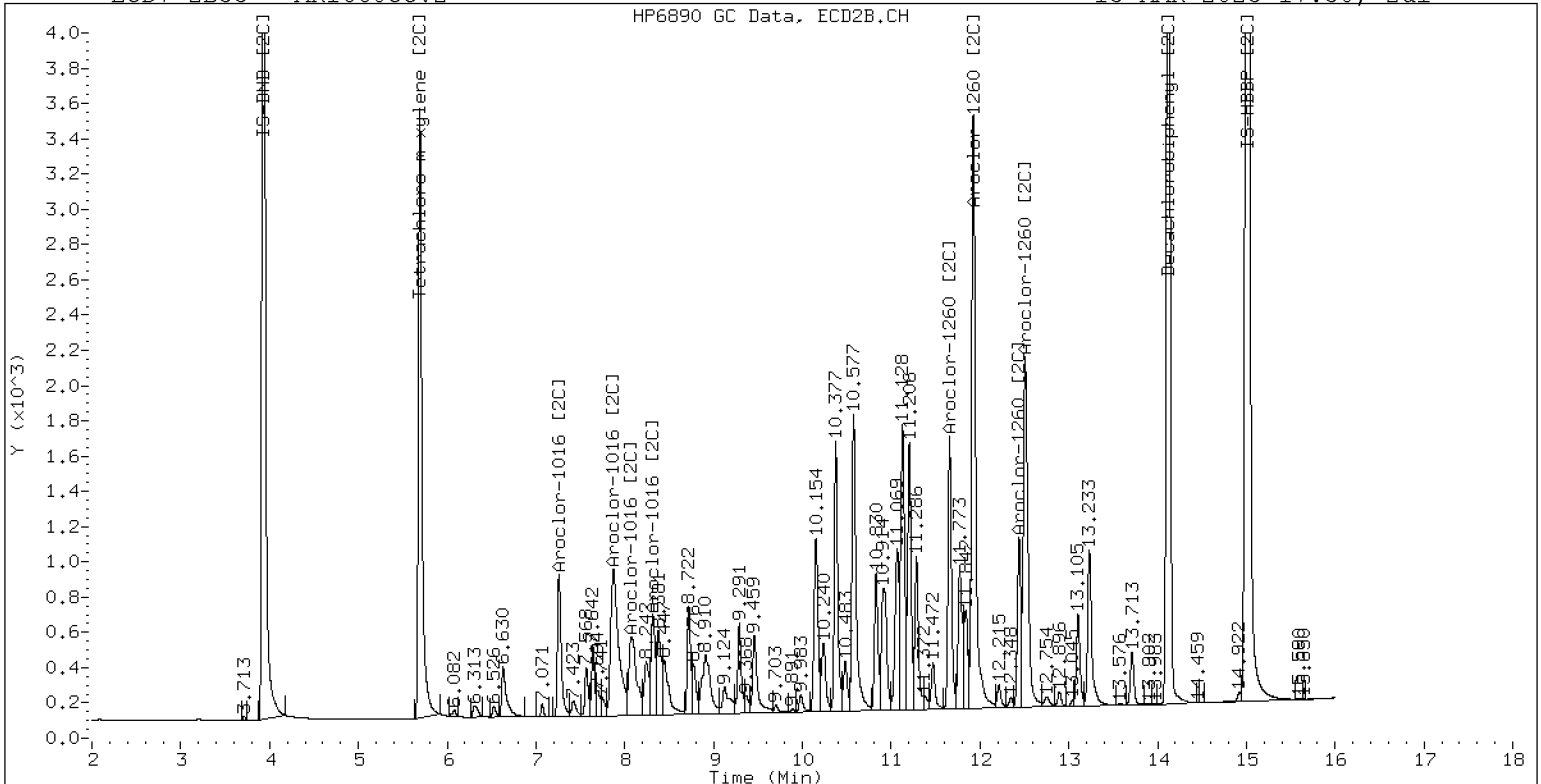
15-MAR-2023 17:30, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1660CCV2

15-MAR-2023 17:30, 2ul



ZB-35 Manual Integration: NO



CONTINUING CALIBRATION CHECK
EPA 8082A

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Instrument ID:	<u>ECD7</u>	Calibration:	<u>GB00069</u>
Lab File ID:	<u>03152331ECD7.D</u>	Calibration Date:	<u>02/24/2023</u>
Sequence:	<u>SLC0215</u>	Injection Date:	<u>03/15/23</u>
Lab Sample ID:	<u>SLC0215-CCV3</u>	Injection Time:	<u>23:24</u>
Sequence Name:	<u>AR1242CCV3</u>		

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1242	A	250.00	219	0.0395340	0.0348168		-12.3	
Aroclor-1242 (1)	A	250.00	227		0.0225326			
Aroclor-1242 (2)	A	250.00	223		0.0672613			
Aroclor-1242 (3)	A	250.00	215		0.0201247			
Aroclor-1242 (4)	A	250.00	212		0.0293487			
Aroclor 1242 [2C]	A	250.00	219	0.0423092	0.0373942		-12.3	
Aroclor-1242 (1) [2C]	A	250.00	225		0.0335097			
Aroclor-1242 (2) [2C]	A	250.00	224		0.0701311			
Aroclor-1242 (3) [2C]	A	250.00	222		0.0215855			
Aroclor-1242 (4) [2C]	A	250.00	206		0.0243505			
Decachlorobiphenyl	A	40.000	33.1	0.7878687	0.6515041		-17.3	
Tetrachlorometaxylene	A	40.000	41.5	1.1944880	1.2387200		3.8	
Decachlorobiphenyl [2C]	A	40.000	35.9	1.2182710	1.0941240		-10.3	
Tetrachlorometaxylene [2C]	A	40.000	42.1	1.1737210	1.2344640		5.3	

* Values outside of QC limits

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152331ECD7.D
Data file 2: /230315.b/230315.b/03152331ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1242.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1242CCV3
Client ID:
Injection Date: 15-MAR-2023 23:24
Report Date: 03/16/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

ZB5 Col		ZB35 Col			ZB5	ZB35	RPD	Compound/Flag	
RT	Shift	Response	RT	Shift	Response	on col			on col
5.811	-0.001	673609	5.691	-0.001	232510	41.5	42.1	1.4	Tetrachloro-m-xylene
13.896	-0.001	396151	14.122	0.001	228272	33.1	35.9	8.3	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1087589	61.4
Hexabromobiphenyl	1429847	1216112	-14.9

Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	376698	19.5
Hexabromobiphenyl	513946	417269	-18.8

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1242	1	7.273	0.002	76582	227.2	1	7.258	0.000	39447	225.4	
Aroclor-1242	2	7.666	0.010	228602	223.3	2	7.872	0.000	82557	224.4	
Aroclor-1242	3	8.412	0.006	68398	214.8	3	9.184	0.000	25410	222.0	
Aroclor-1242	4	8.588	0.009	99748	211.9	4	9.615	0.000	28665	205.5	
Total CollAve (4 peaks):				219.3	Total Col2Ave (4 peaks):				219.3	RPD = 0	
Corrected Ave (3 peaks):				216.7	Corrected Ave (3 peaks):				217.3	RPD = 0	
CalAmt %D:				-12.3	CalAmt %D:				-12.3		

Total PCB Area Col1 (5.911 - 13.797) = 1787261 Col1 Total PCB = 0.1 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 603488 Col2 Total PCB = 0.1 ppm*

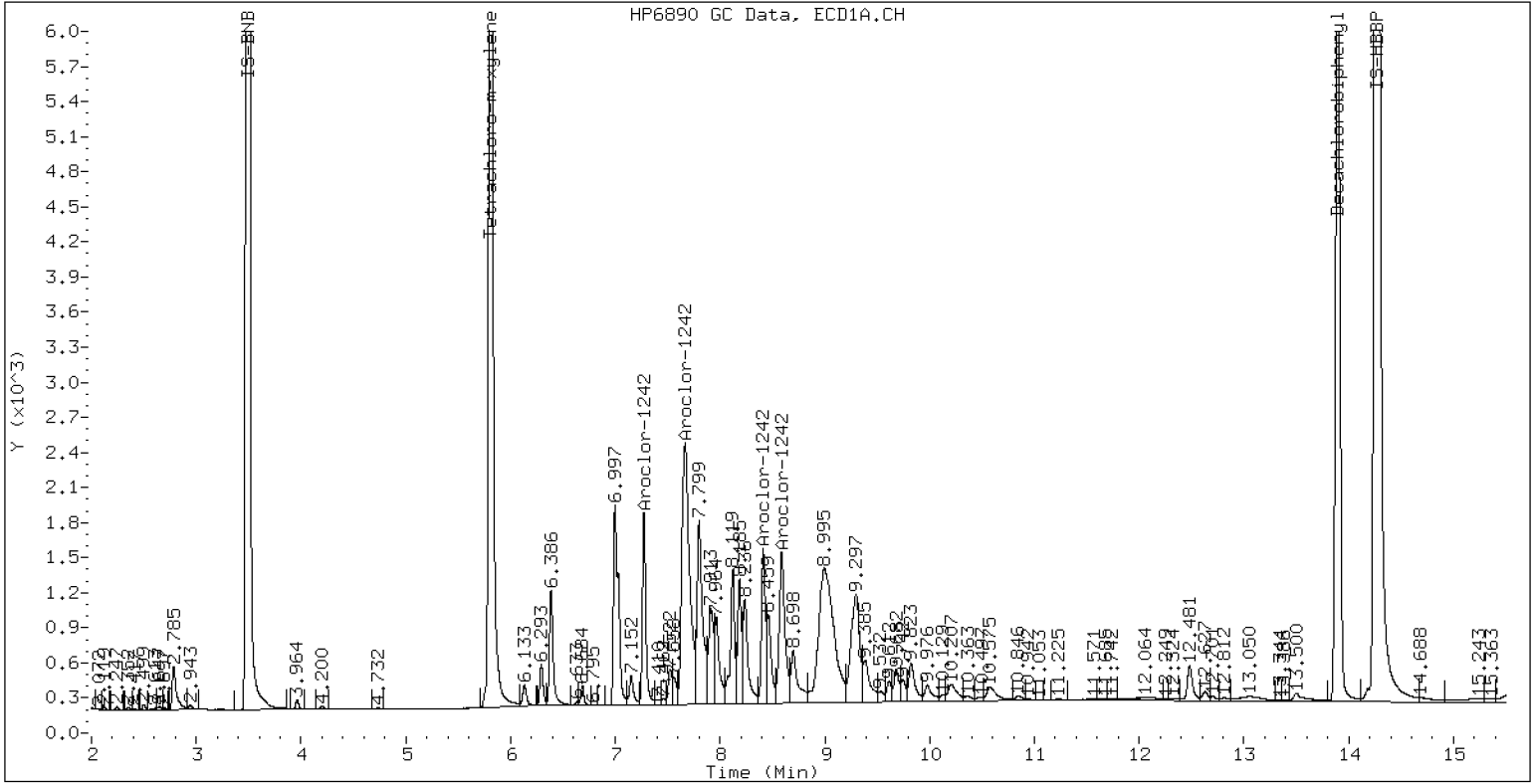
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1242CCV3

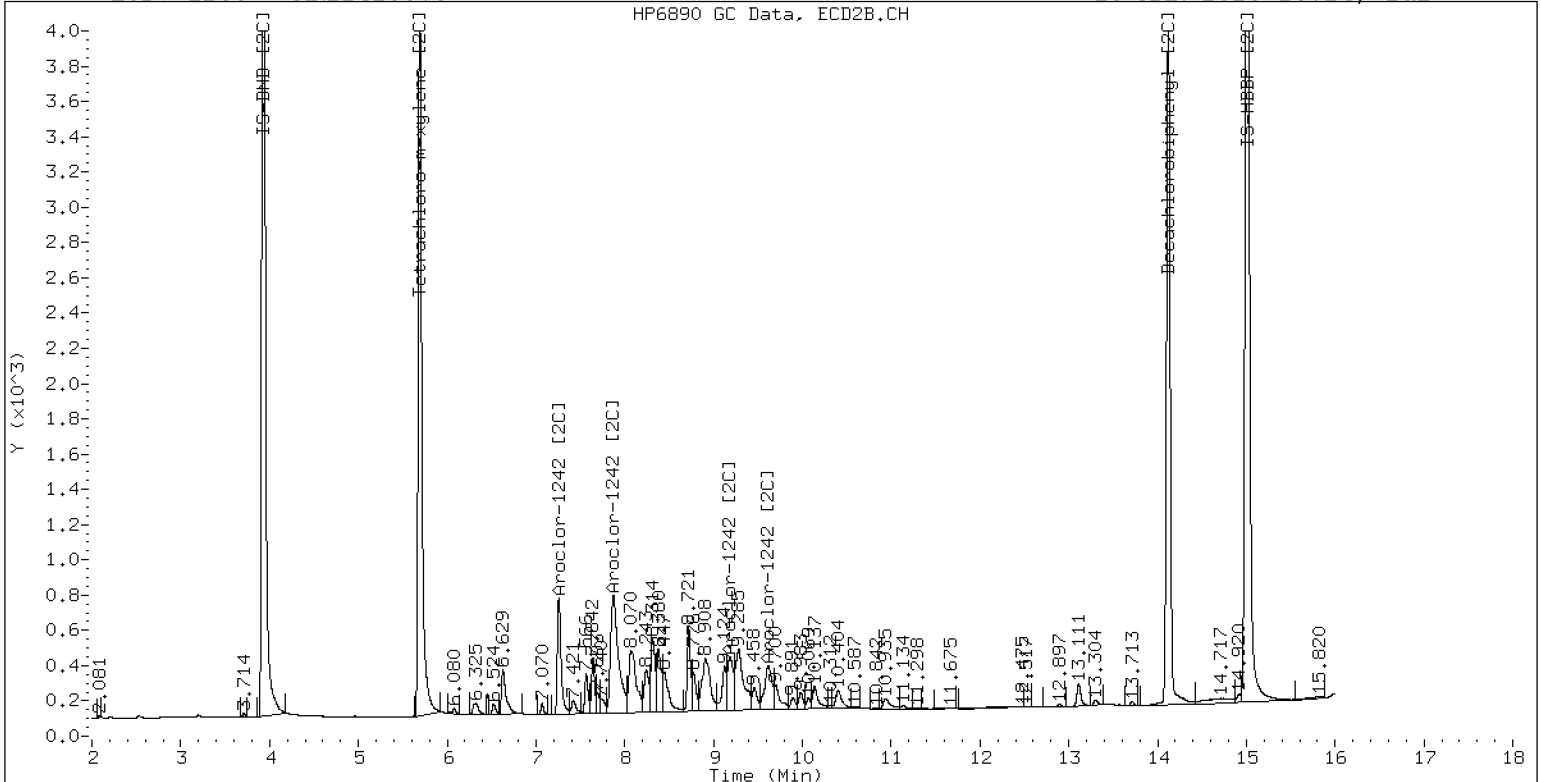
15-MAR-2023 23:24, 2u1



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1242CCV3

15-MAR-2023 23:24, 2u1



ZB-35 Manual Integration: NO



CONTINUING CALIBRATION CHECK
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: ECD7

Calibration: GB00069

Lab File ID: 03152332ECD7.D

Calibration Date: 02/24/2023

Sequence: SLC0215

Injection Date: 03/15/23

Lab Sample ID: SLC0215-CCV4

Injection Time: 23:44

Sequence Name: AR1660CCV4

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1016	A	250.00	243	0.0493662	0.0477128		-3.0	
Aroclor-1016 (1)	A	250.00	236	0.0303852	0.0286670		-5.6	
Aroclor-1016 (2)	A	250.00	237	0.0926308	0.0876814		-5.2	
Aroclor-1016 (3)	A	250.00	258	0.0452180	0.0465798		3.2	
Aroclor-1016 (4)	A	250.00	239	0.0292307	0.0279229		-4.4	
Aroclor 1016 [2C]	A	250.00	248	0.0545857	0.0536477		-1.0	
Aroclor-1016 (1) [2C]	A	250.00	227	0.0468313	0.0425063		-9.2	
Aroclor-1016 (2) [2C]	A	250.00	242	0.0949676	0.0918102		-3.2	
Aroclor-1016 (3) [2C]	A	250.00	274	0.0428922	0.0470015		9.6	
Aroclor-1016 (4) [2C]	A	250.00	247	0.0336515	0.0332726		-1.2	
Aroclor 1260	A	250.00	338	0.0392091	0.0529336		35.1	
Aroclor-1260 (1)	A	250.00	366	0.0287785	0.0421283		46.4	
Aroclor-1260 (2)	A	250.00	354	0.0300690	0.0425492		41.6	
Aroclor-1260 (3)	A	250.00	336	0.0797517	0.1071622		34.4	
Aroclor-1260 (4)	A	250.00	318	0.0401599	0.0510658		27.2	
Aroclor-1260 (5)	A	250.00	315	0.0172866	0.0217625		26.0	
Aroclor 1260 [2C]	A	250.00	269	0.0699688	0.0762886		7.4	
Aroclor-1260 (1) [2C]	A	250.00	278	0.0470406	0.0523470		11.2	
Aroclor-1260 (2) [2C]	A	250.00	279	0.1200523	0.1338828		11.6	
Aroclor-1260 (3) [2C]	A	250.00	247	0.0318590	0.0314995		-1.2	
Aroclor-1260 (4) [2C]	A	250.00	270	0.0809231	0.0874253		8.0	
Decachlorobiphenyl	A	40.000	36.4	0.7878687	0.7167904		-9.0	
Tetrachlorometaxylene	A	40.000	36.7	1.1944880	1.0969820		-8.3	
Decachlorobiphenyl [2C]	A	40.000	38.8	1.2182710	1.1826760		-3.0	
Tetrachlorometaxylene [2C]	A	40.000	36.5	1.1737210	1.0709110		-8.8	

* Values outside of QC limits

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152332ECD7.D
Data file 2: /230315.b/230315.b/03152332ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1660CCV4
Client ID:
Injection Date: 15-MAR-2023 23:44
Report Date: 03/16/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	0.000	608747	5.691	-0.001	202061	36.7	36.5	0.7	Tetrachloro-m-xylene
13.897	0.000	569266	14.121	-0.000	280717	36.4	38.8	6.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1109858	64.7
Hexabromobiphenyl	1429847	1588375	11.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	377363	19.7
Hexabromobiphenyl	513946	474715	-7.6

* Standard Areas taken from Initial Cal Level 3

Initial Calibration Date: 24-FEB-2023

<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount
Aroclor-1016	1	7.273	0.003	99426	235.9	1	7.259	0.000	50126	226.9
Aroclor-1016	2	7.667	0.013	304106	236.6	2	7.871	-0.001	108268	241.7
Aroclor-1016	3	7.799	0.009	161553	257.5	3	8.070	0.000	55427	274.0
Aroclor-1016	4	8.412	0.007	96845	238.8	4	8.314	0.000	39237	247.2
Total CollAve (4 peaks):				242.2		Total Col2Ave (4 peaks):				247.4 RPD = 2
Corrected Ave (3 peaks):				237.1		Corrected Ave (3 peaks):				238.6 RPD = 1

CalAmt %D: -3.1

CalAmt %D: -1.0

Aroclor-1260	1	11.050	0.005	209111	366.0	1	11.658	-0.000	77656	278.2
Aroclor-1260	2	11.368	0.006	211200	353.8	2	11.926	0.000	198613	278.8
Aroclor-1260	3	11.743	0.009	531918	335.9	3	12.441	0.001	46729	247.2
Aroclor-1260	4	12.148	0.009	253474	317.9	4	12.509	0.001	129694	270.1
Aroclor-1260	5	12.249	0.005	108022	314.7	NS	---			----
Total CollAve (5 peaks):				337.7		Total Col2Ave (4 peaks):				268.6 RPD = 23
Corrected Ave (4 peaks):				330.6		Corrected Ave (3 peaks):				265.2 RPD = 22

CalAmt %D: 35.1

CalAmt %D: 7.4

Total PCB Area Col1 (5.911 - 13.797) = 5861640 Col1 Total PCB = 0.4 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 1948467 Col2 Total PCB = 0.4 ppm*

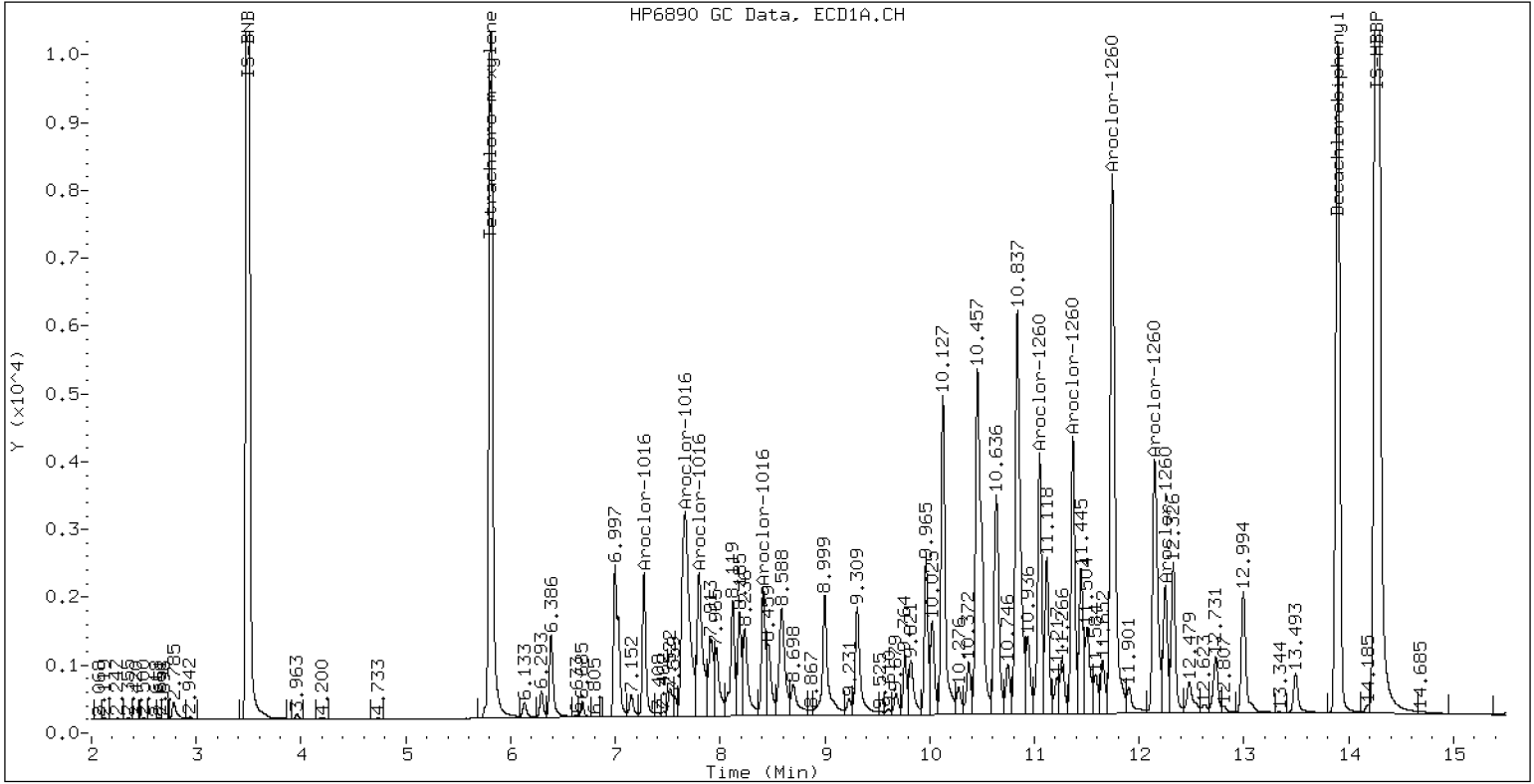
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1660CCV4

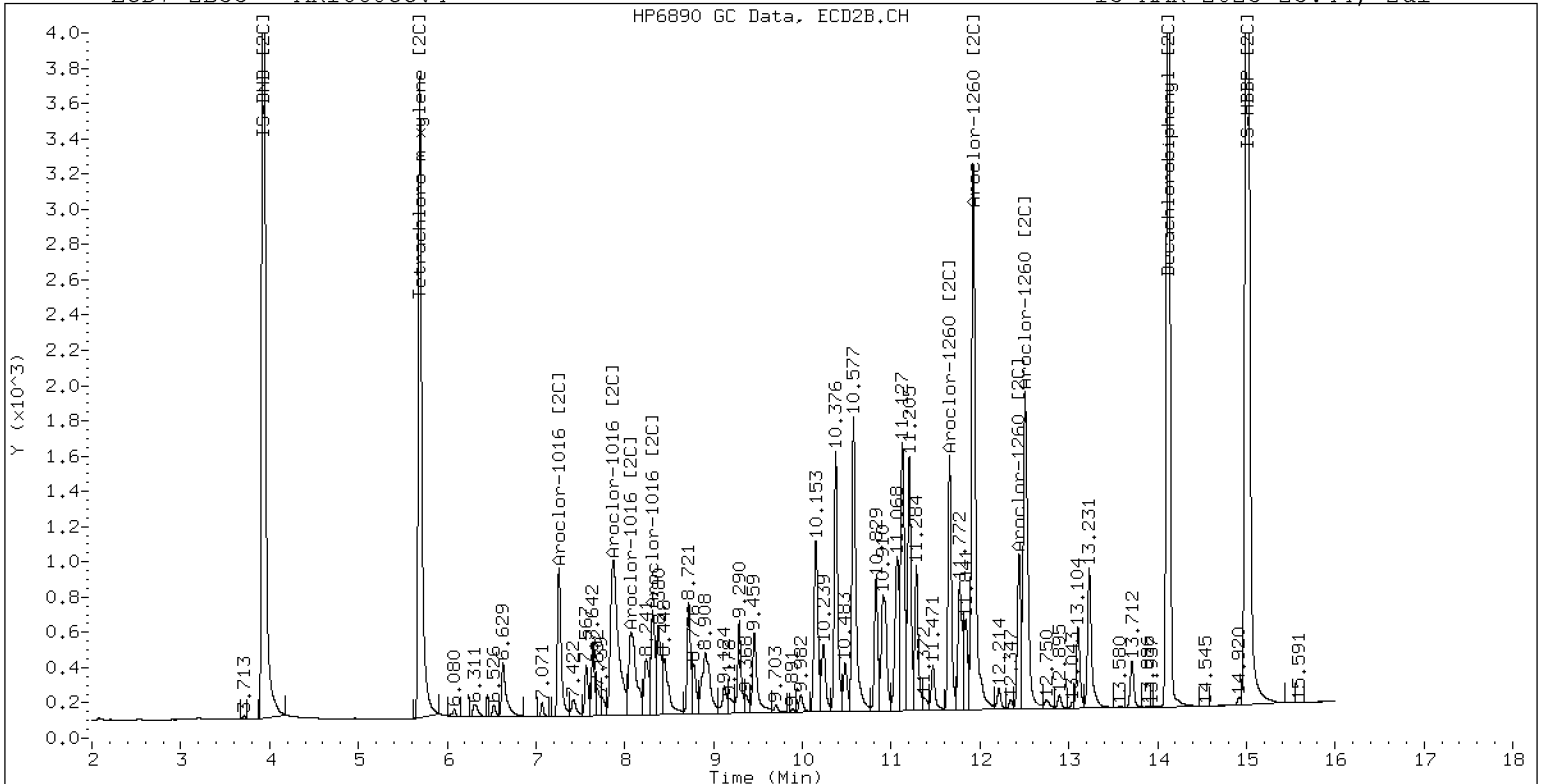
15-MAR-2023 23:44, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1660CCV4

15-MAR-2023 23:44, 2ul



ZB-35 Manual Integration: NO



CONTINUING CALIBRATION CHECK
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: ECD7

Calibration: GB00069

Lab File ID: 03152339ECD7.D

Calibration Date: 02/24/2023

Sequence: SLC0215

Injection Date: 03/16/23

Lab Sample ID: SLC0215-CCV5

Injection Time: 02:10

Sequence Name: AR1254CCV5

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1254	A	250.00	209	0.0662949	0.0552045		-16.3	
Aroclor-1254 (1)	A	250.00	210		0.0675453			
Aroclor-1254 (2)	A	250.00	210		0.0303713			
Aroclor-1254 (3)	A	250.00	200		0.0412775			
Aroclor-1254 (4)	A	250.00	198		0.0795102			
Aroclor-1254 (5)	A	250.00	228		0.0573183			
Aroclor 1254 [2C]	A	250.00	205	0.0763106	0.0624824		-17.8	
Aroclor-1254 (1) [2C]	A	250.00	214		0.0519430			
Aroclor-1254 (2) [2C]	A	250.00	208		0.0406154			
Aroclor-1254 (3) [2C]	A	250.00	204		0.0863452			
Aroclor-1254 (4) [2C]	A	250.00	202		0.0835949			
Aroclor-1254 (5) [2C]	A	250.00	199		0.0499137			
Decachlorobiphenyl	A	40.000	35.9	0.7878687	0.7079897		-10.3	
Tetrachlorometaxylene	A	40.000	34.0	1.1944880	1.0166700		-15.0	
Decachlorobiphenyl [2C]	A	40.000	37.4	1.2182710	1.1389010		-6.5	
Tetrachlorometaxylene [2C]	A	40.000	35.6	1.1737210	1.0449950		-11.0	

* Values outside of QC limits

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152339ECD7.D
Data file 2: /230315.b/230315.b/03152339ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1254.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1254CCV5
Client ID:
Injection Date: 16-MAR-2023 02:10
Report Date: 03/16/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.811	0.000	542525	5.692	-0.000	196179	34.0	35.6	4.5	Tetrachloro-m-xylene
13.895	-0.002	368789	14.121	-0.000	222966	35.9	37.4	4.0	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Standard Cpnd	Column 1		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1067259	58.4
Hexabromobiphenyl	1429847	1041792	-27.1

Standard Cpnd	Column 2		
	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	375464	19.1
Hexabromobiphenyl	513946	391546	-23.8

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col						ZB35 Col					
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1254	1	9.306	0.007	225276	210.2	1	9.458	0.000	60946	213.6	
Aroclor-1254	2	9.385	0.008	101294	210.2	2	9.978	0.000	47655	207.6	
Aroclor-1254	3	9.678	0.010	137668	199.8	3	10.134	0.000	101311	204.0	
Aroclor-1254	4	9.817	0.010	265181	197.9	4	10.382	0.000	98084	202.6	
Aroclor-1254	5	10.189	0.013	191167	227.7	5	10.577	0.000	58565	198.6	
Total CollAve (5 peaks):				209.2		Total Col2Ave (5 peaks):				205.3	RPD = 2
Corrected Ave (4 peaks):				204.5		Corrected Ave (4 peaks):				203.2	RPD = 1
CalAmt %D:				-16.3		CalAmt %D:				-17.9	

Total PCB Area Col1 (5.911 - 13.797) = 2697055 Col1 Total PCB = 0.2 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 974875 Col2 Total PCB = 0.2 ppm*

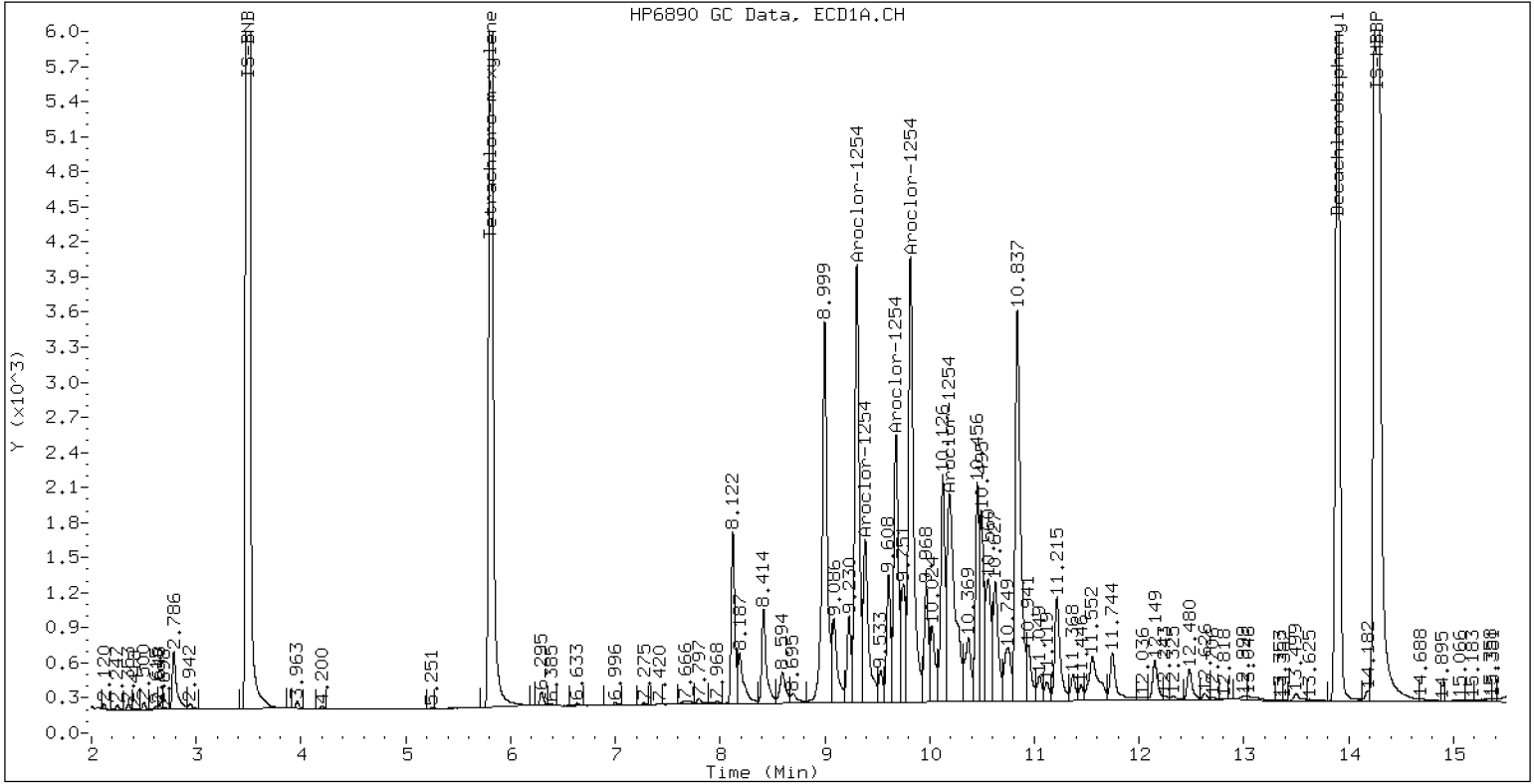
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1254CCV5

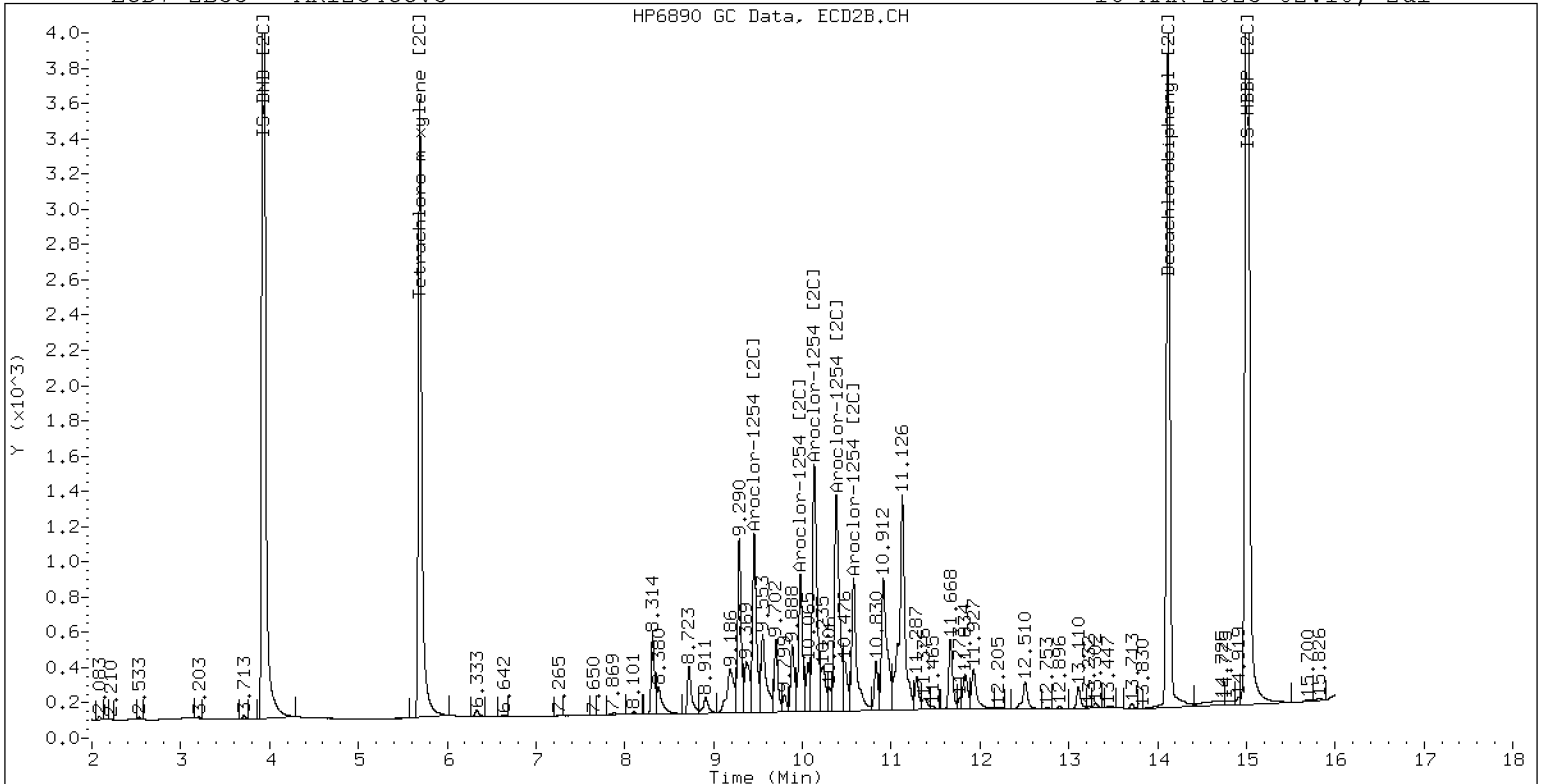
16-MAR-2023 02:10, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1254CCV5

16-MAR-2023 02:10, 2ul



ZB-35 Manual Integration: NO



CONTINUING CALIBRATION CHECK
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: ECD7

Calibration: GB00069

Lab File ID: 03152340ECD7.D

Calibration Date: 02/24/2023

Sequence: SLC0215

Injection Date: 03/16/23

Lab Sample ID: SLC0215-CCV6

Injection Time: 02:31

Sequence Name: AR1660CCV6

COMPOUND	TYPE	CONC. (ug/L)		RESPONSE FACTOR (RRF)			% DRIFT/DIFF	
		STD	CCV	ICAL	CCV	MIN	CCV	LIMIT
Aroclor 1016	A	250.00	233	0.0493662	0.0459448		-6.8	
Aroclor-1016 (1)	A	250.00	230	0.0303852	0.0279355		-8.0	
Aroclor-1016 (2)	A	250.00	229	0.0926308	0.0847361		-8.4	
Aroclor-1016 (3)	A	250.00	246	0.0452180	0.0445537		-1.6	
Aroclor-1016 (4)	A	250.00	227	0.0292307	0.0265541		-9.2	
Aroclor 1016 [2C]	A	250.00	239	0.0545857	0.0519617		-4.3	
Aroclor-1016 (1) [2C]	A	250.00	222	0.0468313	0.0416049		-11.2	
Aroclor-1016 (2) [2C]	A	250.00	235	0.0949676	0.0892226		-6.0	
Aroclor-1016 (3) [2C]	A	250.00	263	0.0428922	0.0451126		5.2	
Aroclor-1016 (4) [2C]	A	250.00	237	0.0336515	0.0319067		-5.2	
Aroclor 1260	A	250.00	329	0.0392091	0.0513805		31.8	
Aroclor-1260 (1)	A	250.00	367	0.0287785	0.0422885		46.8	
Aroclor-1260 (2)	A	250.00	349	0.0300690	0.0420063		39.6	
Aroclor-1260 (3)	A	250.00	321	0.0797517	0.1024213		28.4	
Aroclor-1260 (4)	A	250.00	305	0.0401599	0.0490746		22.0	
Aroclor-1260 (5)	A	250.00	305	0.0172866	0.0211116		22.0	
Aroclor 1260 [2C]	A	250.00	269	0.0699688	0.0763094		7.6	
Aroclor-1260 (1) [2C]	A	250.00	278	0.0470406	0.0523738		11.2	
Aroclor-1260 (2) [2C]	A	250.00	278	0.1200523	0.1336908		11.2	
Aroclor-1260 (3) [2C]	A	250.00	250	0.0318590	0.0318730		0.0	
Aroclor-1260 (4) [2C]	A	250.00	270	0.0809231	0.0873000		8.0	
Decachlorobiphenyl	A	40.000	35.9	0.7878687	0.7073895		-10.3	
Tetrachlorometaxylene	A	40.000	35.5	1.1944880	1.0598660		-11.3	
Decachlorobiphenyl [2C]	A	40.000	37.9	1.2182710	1.1557950		-5.3	
Tetrachlorometaxylene [2C]	A	40.000	36.5	1.1737210	1.0714630		-8.8	

* Values outside of QC limits

Analytical Resources Inc.
Dual Column 608/8082 PCB Quantitation Report

Data file 1: /230315.b/03152340ECD7.D
Data file 2: /230315.b/230315.b/03152340ECD7.D
Method: \\target\share\chem4\ecd7.i\230315.b\PCB.m
Compound Sublist: AR1660.sub
Instrument, Inj. Vol.: ecd7.i, 2ul
Quant Method: Internal Std

ARI ID: AR1660CCV6
Client ID:
Injection Date: 16-MAR-2023 02:31
Report Date: 03/16/2023 09:51
Matrix: NONE
Dilution Factor: 1.0

SURROGATES

RT	ZB5 Col Shift	Response	RT	ZB35 Col Shift	Response	ZB5 on col	ZB35 on col	RPD	Compound/Flag
5.812	0.001	574773	5.692	0.000	200579	35.5	36.5	2.8	Tetrachloro-m-xylene
13.895	-0.002	506065	14.121	0.000	257841	35.9	37.9	5.5	Decachlorobiphenyl

* Indicates RPD > 40%

INTERNAL STANDARD SUMMARY

Column 1			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	673778	1084614	61.0
Hexabromobiphenyl	1429847	1430796	0.1
Column 2			
Standard Cpnd	Standard Area*	Sample Area	%D
Bromo-Nitrobenzene	315256	374402	18.8
Hexabromobiphenyl	513946	446171	-13.2

* Standard Areas taken from Initial Cal Level 3
Initial Calibration Date: 24-FEB-2023
<- Indicates standard response outside Limits (-50 to +100%)

ZB5 Col					ZB35 Col						
Aroclor	Peak#	RT	Shift	Area	Amount	Peak#	RT	Shift	Area	Amount	
Aroclor-1016	1	7.272	0.002	94685	229.8	1	7.259	0.000	48678	222.1	
Aroclor-1016	2	7.664	0.010	287206	228.7	2	7.872	0.000	104391	234.9	
Aroclor-1016	3	7.798	0.008	151011	246.3	3	8.069	0.000	52782	262.9	
Aroclor-1016	4	8.411	0.006	90003	227.1	4	8.313	0.000	37331	237.0	
Total CollAve (4 peaks):				233.0		Total Col2Ave (4 peaks):				239.2	RPD = 3
Corrected Ave (3 peaks):				228.5		Corrected Ave (3 peaks):				231.3	RPD = 1

CalAmt %D: -6.8

CalAmt %D: -4.3

Aroclor-1260	1	11.048	0.004	189082	367.4	1	11.658	0.000	73024	278.3	
Aroclor-1260	2	11.367	0.005	187820	349.2	2	11.925	0.000	186403	278.4	
Aroclor-1260	3	11.742	0.008	457950	321.1	3	12.440	0.000	44440	250.1	
Aroclor-1260	4	12.147	0.008	219424	305.5	4	12.508	0.000	121721	269.7	
Aroclor-1260	5	12.248	0.004	94395	305.3	NS	---			----	
Total CollAve (5 peaks):				329.7		Total Col2Ave (4 peaks):				269.1	RPD = 20
Corrected Ave (4 peaks):				320.3		Corrected Ave (3 peaks):				266.1	RPD = 18

CalAmt %D: 31.9

CalAmt %D: 7.7

Total PCB Area Coll (5.911 - 13.797) = 5391813 Coll Total PCB = 0.4 ppm*

Total PCB Area Col2 (5.792 - 14.021) = 1843359 Col2 Total PCB = 0.4 ppm*

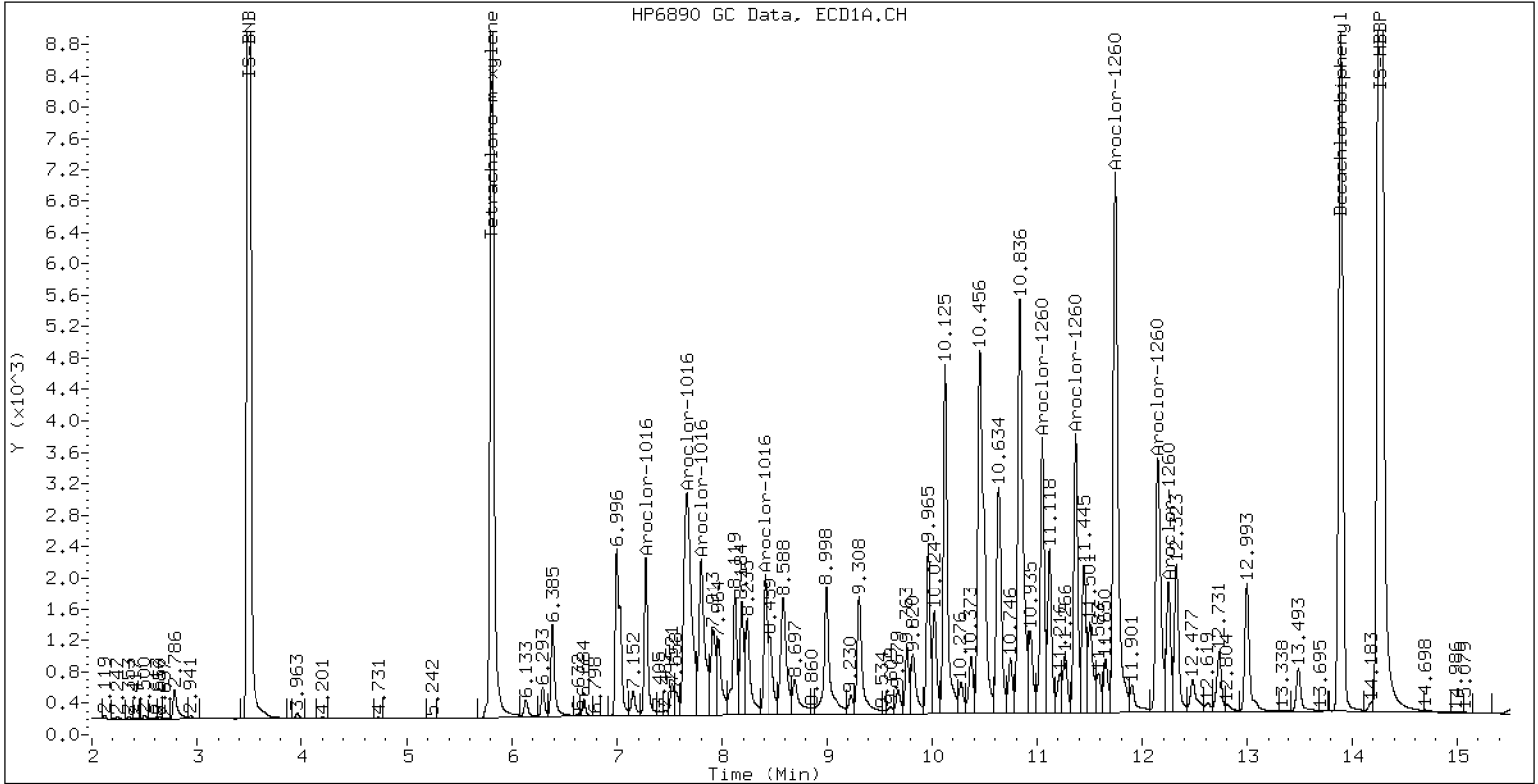
* Quantitated against AR1660 0.25ppm in Ical

PCB-Form 10 Mod.

PCB Dual Column Chromatograms

ECD7-ZB5 AR1660CCV6

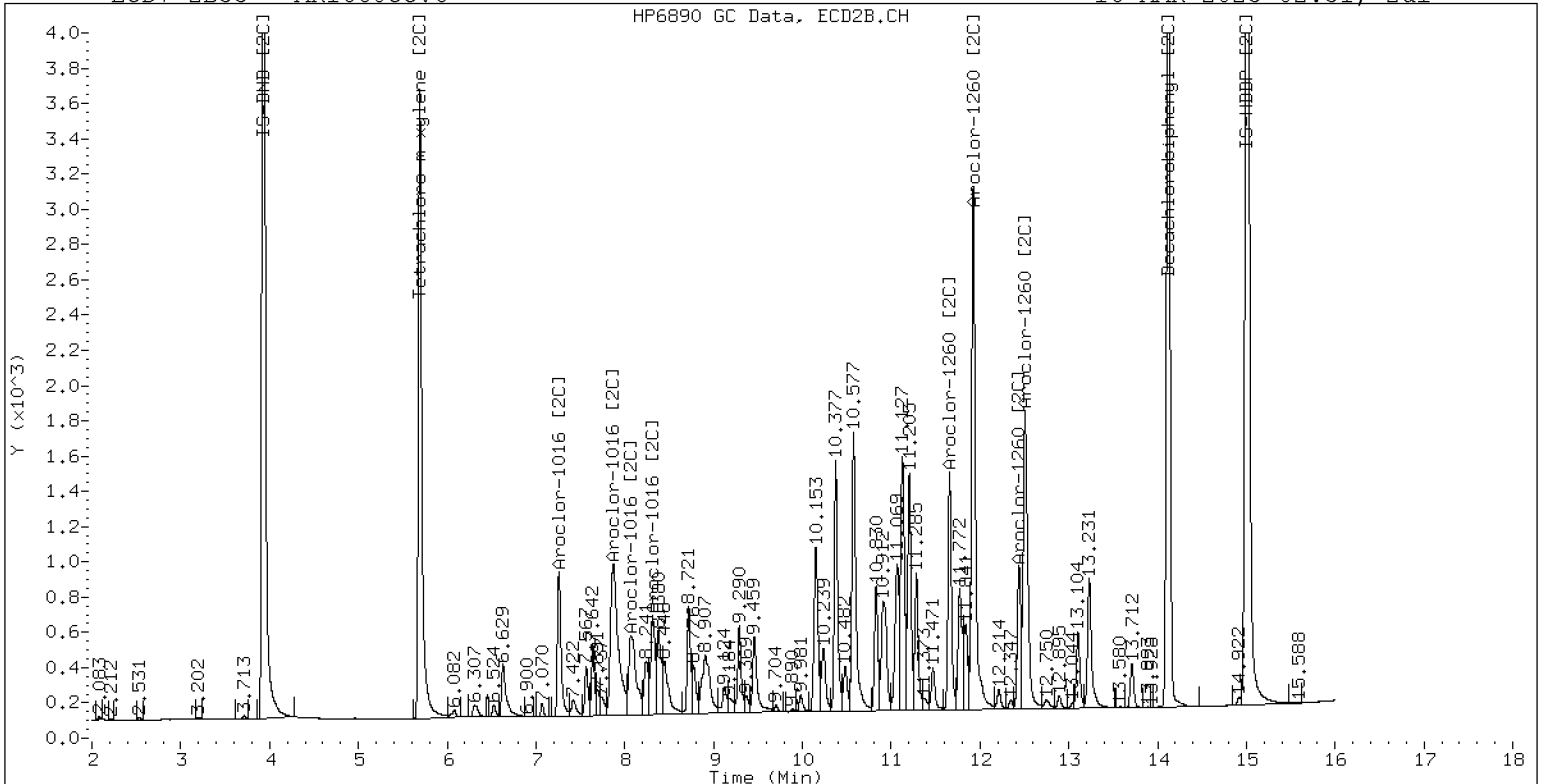
16-MAR-2023 02:31, 2ul



ZB-5 Manual Integration: NO

ECD7-ZB35 AR1660CCV6

16-MAR-2023 02:31, 2ul



ZB-35 Manual Integration: NO



Dual Column
ANALYSIS BATCH (SEQUENCE) SUMMARY
EPA 8082A

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor OEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Sequence:	<u>SLB0342</u>	Instrument:	<u>ECD7</u>
		Calibration:	<u>GB00069</u>

Sample Name	Lab Sample ID	Column 1 File ID	Column 2 File ID	Matrix	Analysis Date/Time
Cal Standard	SLB0342-CAL1	02242302ECD7.D	02242302ECD7.D	NA	02/24/23 11:12
Cal Standard	SLB0342-CAL2	02242303ECD7.D	02242303ECD7.D	NA	02/24/23 11:33
Cal Standard	SLB0342-CAL3	02242304ECD7.D	02242304ECD7.D	NA	02/24/23 11:54
Cal Standard	SLB0342-CAL4	02242305ECD7.D	02242305ECD7.D	NA	02/24/23 12:15
Cal Standard	SLB0342-CAL5	02242306ECD7.D	02242306ECD7.D	NA	02/24/23 12:36
Cal Standard	SLB0342-CAL6	02242307ECD7.D	02242307ECD7.D	NA	02/24/23 12:57
Cal Standard	SLB0342-CAL7	02242308ECD7.D	02242308ECD7.D	NA	02/24/23 13:18
Cal Standard	SLB0342-CAL8	02242309ECD7.D	02242309ECD7.D	NA	02/24/23 13:39
Cal Standard	SLB0342-CAL9	02242310ECD7.D	02242310ECD7.D	NA	02/24/23 14:00
Cal Standard	SLB0342-CALA	02242311ECD7.D	02242311ECD7.D	NA	02/24/23 14:21
Cal Standard	SLB0342-CALB	02242312ECD7.D	02242312ECD7.D	NA	02/24/23 14:42
Secondary Cal Check	SLB0342-SCV1	02242313ECD7.D	02242313ECD7.D	NA	02/24/23 15:03
Secondary Cal Check	SLB0342-SCV2	02242314ECD7.D	02242314ECD7.D	NA	02/24/23 15:24
Secondary Cal Check	SLB0342-SCV3	02242315ECD7.D	02242315ECD7.D	NA	02/24/23 15:45
Secondary Cal Check	SLB0342-SCV4	02242316ECD7.D	02242316ECD7.D	NA	02/24/23 16:06
Secondary Cal Check	SLB0342-SCV5	02242317ECD7.D	02242317ECD7.D	NA	02/24/23 16:27
Secondary Cal Check	SLB0342-SCV6	02242318ECD7.D	02242318ECD7.D	NA	02/24/23 16:48



Dual Column
ANALYSIS BATCH (SEQUENCE) SUMMARY
EPA 8082A

Laboratory: Analytical Resources, LLC SDG: 23C0107
Client: Anchor OEA, LLC Project: AOC4 UR Phase 3
Sequence: SLC0215 Instrument: ECD7
Calibration: GB00069

Sample Name	Lab Sample ID	Column 1 File ID	Column 2 File ID	Matrix	Analysis Date/Time
Initial Cal Check	SLC0215-ICV1	03152306ECD7.D	03152306ECD7.D	NA	03/15/23 14:43
Initial Cal Check	SLC0215-ICV2	03152307ECD7.D	03152307ECD7.D	NA	03/15/23 15:04
Calibration Check	SLC0215-CCV1	03152313ECD7.D	03152313ECD7.D	NA	03/15/23 17:09
Calibration Check	SLC0215-CCV2	03152314ECD7.D	03152314ECD7.D	NA	03/15/23 17:30
Blank	BLC0219-BLK1	03152315ECD7.D	03152315ECD7.D	Solid	03/15/23 17:51
LCS	BLC0219-BS1	03152316ECD7.D	03152316ECD7.D	Solid	03/15/23 18:12
LCS Dup	BLC0219-BSD1	03152317ECD7.D	03152317ECD7.D	Solid	03/15/23 18:32
Reference	BLC0219-SRM1	03152318ECD7.D	03152318ECD7.D	Solid	03/15/23 18:53
LDW22-SS826	23C0107-01	03152319ECD7.D	03152319ECD7.D	Solid	03/15/23 19:14
LDW21-IT608C	23C0107-02	03152320ECD7.D	03152320ECD7.D	Solid	03/15/23 19:35
LDW21-IT608D	23C0107-03	03152321ECD7.D	03152321ECD7.D	Solid	03/15/23 19:56
LDW21-IT608D	BLC0219-MS1	03152322ECD7.D	03152322ECD7.D	Solid	03/15/23 20:16
LDW21-IT608D	BLC0219-MSD1	03152323ECD7.D	03152323ECD7.D	Solid	03/15/23 20:37
Calibration Check	SLC0215-CCV3	03152331ECD7.D	03152331ECD7.D	NA	03/15/23 23:24
Calibration Check	SLC0215-CCV4	03152332ECD7.D	03152332ECD7.D	NA	03/15/23 23:44
Calibration Check	SLC0215-CCV5	03152339ECD7.D	03152339ECD7.D	NA	03/16/23 02:10
Calibration Check	SLC0215-CCV6	03152340ECD7.D	03152340ECD7.D	NA	03/16/23 02:31



ANALYSIS SEQUENCE

SLC0215

Instrument: ECD7
Calibration ID: GB00069

Printed: 3/16/2023 10:26:54AM

Lab Number	Analysis	Container	Order	Position	STD ID	ISTD ID	Client	Comments
SLC0215-ICV1	QC		1		L000862	L000844		
SLC0215-ICV2	QC		2		L000856	L000844		
BLC0266-BLK1	QC		3			L000844		
BLC0266-BS1	QC		4			L000844		
BLC0266-BSD1	QC		5			L000844		
BLC0266-MRL1	QC		6			L000844		
23C0221-01	8082A PCB Water 0.01	I 01	7			L000844	DH Environmental Inc	
SLC0215-CCV1	QC		8		L000861	L000844		
SLC0215-CCV2	QC		9		L000856	L000844		
BLC0219-BLK1	QC		10			L000844		
BLC0219-BS1	QC		11			L000844		
BLC0219-BSD1	QC		12			L000844		
BLC0219-SRM1	QC		13			L000844		
23C0107-01	8082A PCB Solid 4	A 05	14			L000844	Anchor QEA, LLC	
23C0107-02	8082A PCB Solid 4	A 05	15			L000844	Anchor QEA, LLC	
23C0107-03	8082A PCB Solid 4	A 05	16			L000844	Anchor QEA, LLC	
BLC0219-MS1	QC		17			L000844		
BLC0219-MSD1	QC		18			L000844		
23C0108-01	8082A PCB Solid 4	A 05	19			L000844	Anchor QEA, LLC	
23C0108-02	8082A PCB Solid 4	A 05	20			L000844	Anchor QEA, LLC	
23C0108-03	8082A PCB Solid 4	A 05	21			L000844	Anchor QEA, LLC	

Samples Loaded By _____ Date _____

Data Processed By _____ Date _____



ANALYSIS SEQUENCE

SLC0215

Instrument: ECD7
Calibration ID: GB00069

Printed: 3/16/2023 10:26:54AM

Lab Number	Analysis	Container	Order	Position	STD ID	ISTD ID	Client	Comments
23C0108-04	8082A PCB Solid 4	A 05	22			L000844	Anchor QEA, LLC	
23C0108-05	8082A PCB Solid 4	A 05	23			L000844	Anchor QEA, LLC	
23C0108-06	8082A PCB Solid 4	A 05	24			L000844	Anchor QEA, LLC	
23C0108-07	8082A PCB Solid 4	A 05	25			L000844	Anchor QEA, LLC	
SLC0215-CCV3	QC		26		L000860	L000844		
SLC0215-CCV4	QC		27		L000856	L000844		
23C0108-08	8082A PCB Solid 4	A 05	28			L000844	Anchor QEA, LLC	
23C0108-09	8082A PCB Solid 4	A 05	29			L000844	Anchor QEA, LLC	
23C0108-10	8082A PCB Solid 4	A 05	30			L000844	Anchor QEA, LLC	
23C0109-01	8082A PCB Solid 4	A 05	31			L000844	Anchor QEA, LLC	
23C0109-02	8082A PCB Solid 4	A 05	32			L000844	Anchor QEA, LLC	
23C0109-03	8082A PCB Solid 4	A 05	33			L000844	Anchor QEA, LLC	
SLC0215-CCV5	QC		34		L000862	L000844		
SLC0215-CCV6	QC		35		L000856	L000844		

Samples Loaded By _____ Date _____

Data Processed By _____ Date _____

GC LOG SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230315.b

	Inject	Date/Time	Filename	DF	LabID	ClientID
1	15-MAR-2023	12:59	03152301ECD7.D	1	AR1254	
2	15-MAR-2023	13:20	03152302ECD7.D	1	AR1660	
3	15-MAR-2023	13:41	03152303ECD7.D	1	AR1248	
4	15-MAR-2023	14:01	03152304ECD7.D	1	AR1242	
5	15-MAR-2023	14:22	03152305ECD7.D	1	DDTS	
6	15-MAR-2023	14:43	03152306ECD7.D	1	AR1254ICV1	
7	15-MAR-2023	15:04	03152307ECD7.D	1	AR1660ICV2	
8	15-MAR-2023	15:25	03152308ECD7.D	1	BLC0266-BLK1	
9	15-MAR-2023	15:46	03152309ECD7.D	1	BLC0266-BS1	
10	15-MAR-2023	16:06	03152310ECD7.D	1	BLC0266-BSD1	
11	15-MAR-2023	16:27	03152311ECD7.D	1	BLC0266-MRL1	
12	15-MAR-2023	16:48	03152312ECD7.D	1	23C0221-01	
13	15-MAR-2023	17:09	03152313ECD7.D	1	AR1248CCV1	
14	15-MAR-2023	17:30	03152314ECD7.D	1	AR1660CCV2	
15	15-MAR-2023	17:51	03152315ECD7.D	1	BLC0219-BLK1	
16	15-MAR-2023	18:12	03152316ECD7.D	1	BLC0219-BS1	
17	15-MAR-2023	18:32	03152317ECD7.D	1	BLC0219-BSD1	
18	15-MAR-2023	18:53	03152318ECD7.D	1	BLC0219-SRM1	
19	15-MAR-2023	19:14	03152319ECD7.D	1	23C0107-01	
20	15-MAR-2023	19:35	03152320ECD7.D	1	23C0107-02	
21	15-MAR-2023	19:56	03152321ECD7.D	1	23C0107-03	
22	15-MAR-2023	20:16	03152322ECD7.D	1	BLC0219-MS1	
23	15-MAR-2023	20:37	03152323ECD7.D	1	BLC0219-MSD1	
24	15-MAR-2023	20:58	03152324ECD7.D	1	23C0108-01	
25	15-MAR-2023	21:19	03152325ECD7.D	1	23C0108-02	
26	15-MAR-2023	21:40	03152326ECD7.D	1	23C0108-03	
27	15-MAR-2023	22:00	03152327ECD7.D	1	23C0108-04	
28	15-MAR-2023	22:21	03152328ECD7.D	1	23C0108-05	
29	15-MAR-2023	22:42	03152329ECD7.D	1	23C0108-06	
30	15-MAR-2023	23:03	03152330ECD7.D	1	23C0108-07	
31	15-MAR-2023	23:24	03152331ECD7.D	1	AR1242CCV3	
32	15-MAR-2023	23:44	03152332ECD7.D	1	AR1660CCV4	
33	16-MAR-2023	00:05	03152333ECD7.D	1	23C0108-08	
34	16-MAR-2023	00:26	03152334ECD7.D	1	23C0108-09	
35	16-MAR-2023	00:47	03152335ECD7.D	1	23C0108-10	
36	16-MAR-2023	01:07	03152336ECD7.D	1	23C0109-01	
37	16-MAR-2023	01:28	03152337ECD7.D	1	23C0109-02	
38	16-MAR-2023	01:49	03152338ECD7.D	1	23C0109-03	
39	16-MAR-2023	02:10	03152339ECD7.D	1	AR1254CCV5	
40	16-MAR-2023	02:31	03152340ECD7.D	1	AR1660CCV6	

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230315.b

ARI Job No.: AR12 Method: PCB.m Instrument: ecd7.i Date: 15-MAR-2023

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1259	03152301ECD7.D	AR1254		1	NO MANUAL INTEGRATION
1320	03152302ECD7.D	AR1660		1	NO MANUAL INTEGRATION
1341	03152303ECD7.D	AR1248		1	Aroclor-1248,
1401	03152304ECD7.D	AR1242		1	NO MANUAL INTEGRATION
1422	03152305ECD7.D	DDTS		1	NO MANUAL INTEGRATION
1443	03152306ECD7.D	AR1254ICV1		1	NO MANUAL INTEGRATION
1504	03152307ECD7.D	AR1660ICV2		1	NO MANUAL INTEGRATION
1525	03152308ECD7.D	BLC0266-BLK1		1	NO MANUAL INTEGRATION
1546	03152309ECD7.D	BLC0266-BS1		1	NO MANUAL INTEGRATION
1606	03152310ECD7.D	BLC0266-BSD1		1	NO MANUAL INTEGRATION
1627	03152311ECD7.D	BLC0266-MRL1		1	NO MANUAL INTEGRATION
1648	03152312ECD7.D	23C0221-01		1	NO MANUAL INTEGRATION
1709	03152313ECD7.D	AR1248CCV1		1	Aroclor-1248,
1730	03152314ECD7.D	AR1660CCV2		1	NO MANUAL INTEGRATION
1751	03152315ECD7.D	BLC0219-BLK1		1	NO MANUAL INTEGRATION
1812	03152316ECD7.D	BLC0219-BS1		1	NO MANUAL INTEGRATION
1832	03152317ECD7.D	BLC0219-BSD1		1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230315.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1853	03152318ECD7.D	BLC0219-SRM1		1	NO MANUAL INTEGRATION
1914	03152319ECD7.D	23C0107-01		1	Aroclor-1254,
1935	03152320ECD7.D	23C0107-02		1	NO MANUAL INTEGRATION
1956	03152321ECD7.D	23C0107-03		1	NO MANUAL INTEGRATION
2016	03152322ECD7.D	BLC0219-MS1		1	NO MANUAL INTEGRATION
2037	03152323ECD7.D	BLC0219-MSD1		1	NO MANUAL INTEGRATION
2058	03152324ECD7.D	23C0108-01		1	NO MANUAL INTEGRATION
2119	03152325ECD7.D	23C0108-02		1	Aroclor-1254,
2140	03152326ECD7.D	23C0108-03		1	Aroclor-1254,
2200	03152327ECD7.D	23C0108-04		1	Aroclor-1254,
2221	03152328ECD7.D	23C0108-05		1	Aroclor-1254,
2242	03152329ECD7.D	23C0108-06		1	Aroclor-1254,
2303	03152330ECD7.D	23C0108-07		1	NO MANUAL INTEGRATION
2324	03152331ECD7.D	AR1242CCV3		1	NO MANUAL INTEGRATION
2344	03152332ECD7.D	AR1660CCV4		1	NO MANUAL INTEGRATION
0005	03152333ECD7.D	23C0108-08		1	Aroclor-1254,
0026	03152334ECD7.D	23C0108-09		1	Aroclor-1254,
0047	03152335ECD7.D	23C0108-10		1	Aroclor-1254,

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230315.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
0107	03152336ECD7.D	23C0109-01		1	Aroclor-1254,
0128	03152337ECD7.D	23C0109-02		1	Aroclor-1254,
0149	03152338ECD7.D	23C0109-03		1	Aroclor-1254,
0210	03152339ECD7.D	AR1254CCV5		1	NO MANUAL INTEGRATION
0231	03152340ECD7.D	AR1660CCV6		1	NO MANUAL INTEGRATION
1259	03152301ECD7.D	AR1254		1	NO MANUAL INTEGRATION
1320	03152302ECD7.D	AR1660		1	NO MANUAL INTEGRATION
1341	03152303ECD7.D	AR1248		1	NO MANUAL INTEGRATION
1401	03152304ECD7.D	AR1242		1	NO MANUAL INTEGRATION
1422	03152305ECD7.D	DDTS		1	NO MANUAL INTEGRATION
1443	03152306ECD7.D	AR1254ICV1		1	NO MANUAL INTEGRATION
1504	03152307ECD7.D	AR1660ICV2		1	NO MANUAL INTEGRATION
1525	03152308ECD7.D	BLC0266-BLK1		1	NO MANUAL INTEGRATION
1546	03152309ECD7.D	BLC0266-BS1		1	NO MANUAL INTEGRATION
1606	03152310ECD7.D	BLC0266-BSD1		1	NO MANUAL INTEGRATION
1627	03152311ECD7.D	BLC0266-MRL1		1	NO MANUAL INTEGRATION
1648	03152312ECD7.D	23C0221-01		1	NO MANUAL INTEGRATION
1709	03152313ECD7.D	AR1248CCV1		1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230315.b\230315.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
1730	03152314ECD7.D	AR1660CCV2		1	NO MANUAL INTEGRATION
1751	03152315ECD7.D	BLC0219-BLK1		1	NO MANUAL INTEGRATION
1812	03152316ECD7.D	BLC0219-BS1		1	NO MANUAL INTEGRATION
1832	03152317ECD7.D	BLC0219-BSD1		1	NO MANUAL INTEGRATION
1853	03152318ECD7.D	BLC0219-SRM1		1	NO MANUAL INTEGRATION
1914	03152319ECD7.D	23C0107-01		1	Aroclor-1248 [2C],
1935	03152320ECD7.D	23C0107-02		1	NO MANUAL INTEGRATION
1956	03152321ECD7.D	23C0107-03		1	NO MANUAL INTEGRATION
2016	03152322ECD7.D	BLC0219-MS1		1	NO MANUAL INTEGRATION
2037	03152323ECD7.D	BLC0219-MSD1		1	NO MANUAL INTEGRATION
2058	03152324ECD7.D	23C0108-01		1	Aroclor-1248 [2C],
2119	03152325ECD7.D	23C0108-02		1	Aroclor-1248 [2C],
2140	03152326ECD7.D	23C0108-03		1	Aroclor-1248 [2C],
2200	03152327ECD7.D	23C0108-04		1	Aroclor-1248 [2C],
2221	03152328ECD7.D	23C0108-05		1	Aroclor-1248 [2C],
2242	03152329ECD7.D	23C0108-06		1	Aroclor-1248 [2C],
2303	03152330ECD7.D	23C0108-07		1	Aroclor-1248 [2C],
2324	03152331ECD7.D	AR1242CCV3		1	NO MANUAL INTEGRATION

MANUAL INTEGRATION SUMMARY FOR DATABATCH - \\target\share\chem4\ecd7.i\230315.b\230315.b

Time	Filename	LabID	ClientId	DF	Manually Integrated Compounds
2344	03152332ECD7.D	AR1660CCV4		1	NO MANUAL INTEGRATION
0005	03152333ECD7.D	23C0108-08		1	Aroclor-1248 [2C],
0026	03152334ECD7.D	23C0108-09		1	Aroclor-1248 [2C],
0047	03152335ECD7.D	23C0108-10		1	Aroclor-1248 [2C],
0107	03152336ECD7.D	23C0109-01		1	Aroclor-1248 [2C],
0128	03152337ECD7.D	23C0109-02		1	Aroclor-1248 [2C],
0149	03152338ECD7.D	23C0109-03		1	Aroclor-1248 [2C],
0210	03152339ECD7.D	AR1254CCV5		1	NO MANUAL INTEGRATION
0231	03152340ECD7.D	AR1660CCV6		1	NO MANUAL INTEGRATION

Security Status Report

Date: 16-Mar-2023 10:23

03152301ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152302ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152303ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152304ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152305ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152306ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152307ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152308ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152309ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152310ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152311ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152312ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152313ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152314ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152315ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152316ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152317ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152318ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152319ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152320ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152321ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152322ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152323ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152324ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152325ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152326ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152327ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152328ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152329ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152330ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152331ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152332ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152333ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152334ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152335ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152336ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152337ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152338ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152339ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23
03152340ECD7.D	Data Locked	richardl, 16-Mar-2023 10:23



SURROGATE RECOVERY AND RT SUMMARY
EPA 8082A

Laboratory:	<u>Analytical Resources, LLC</u>	SDG/WO:	<u>23C0107</u>
Client:	<u>Anchor OEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Sequence:	<u>SLB0342</u>	Instrument:	<u>ECD7</u>
Calibration:	<u>GB00069</u>	Calibration Date:	<u>02/24/2023</u>

Surrogate Compound	Spike Level ug/L	% Recovery	Recovery Limits	RT	Calibration Mean RT	RT Diff	RT Diff Limit	Q
SLB0342-SCV1 (Water) Lab File ID: 02242313ECD7.D Analyzed: 02/24/23 15:03								
Decachlorobiphenyl	40.000	85.8	80 - 120	13.894	13.89483	-0.0008	N/A	
Tetrachlorometaxylene	40.000	87.4	80 - 120	5.807	5.8095	-0.0025	N/A	
Decachlorobiphenyl [2C]	40.000	93.4	80 - 120	14.119	14.11917	-0.0002	N/A	
Tetrachlorometaxylene [2C]	40.000	89.4	80 - 120	5.685	5.687167	-0.0022	N/A	
SLB0342-SCV2 (Water) Lab File ID: 02242314ECD7.D Analyzed: 02/24/23 15:24								
Decachlorobiphenyl	40.000	92.5	80 - 120	13.895	13.89483	0.0002	N/A	
Tetrachlorometaxylene	40.000	84.1	80 - 120	5.808	5.8095	-0.0015	N/A	
Decachlorobiphenyl [2C]	40.000	101	80 - 120	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	40.000	86.3	80 - 120	5.686	5.687167	-0.0012	N/A	
SLB0342-SCV3 (Water) Lab File ID: 02242315ECD7.D Analyzed: 02/24/23 15:45								
Decachlorobiphenyl	40.000	82.8	80 - 120	13.893	13.89483	-0.0018	N/A	
Tetrachlorometaxylene	40.000	87.2	80 - 120	5.808	5.8095	-0.0015	N/A	
Decachlorobiphenyl [2C]	40.000	90.8	80 - 120	14.118	14.11917	-0.0012	N/A	
Tetrachlorometaxylene [2C]	40.000	91.0	80 - 120	5.687	5.687167	-0.0002	N/A	
SLB0342-SCV4 (Water) Lab File ID: 02242316ECD7.D Analyzed: 02/24/23 16:06								
Decachlorobiphenyl	40.000	86.6	80 - 120	13.894	13.89483	-0.0008	N/A	
Tetrachlorometaxylene	40.000	90.3	80 - 120	5.806	5.8095	-0.0035	N/A	
Decachlorobiphenyl [2C]	40.000	94.8	80 - 120	14.119	14.11917	-0.0002	N/A	
Tetrachlorometaxylene [2C]	40.000	92.7	80 - 120	5.685	5.687167	-0.0022	N/A	
SLB0342-SCV5 (Water) Lab File ID: 02242317ECD7.D Analyzed: 02/24/23 16:27								
Decachlorobiphenyl	40.000	86.1	80 - 120	13.894	13.89483	-0.0008	N/A	
Tetrachlorometaxylene	40.000	90.0	80 - 120	5.806	5.8095	-0.0035	N/A	
Decachlorobiphenyl [2C]	40.000	94.6	80 - 120	14.119	14.11917	-0.0002	N/A	
Tetrachlorometaxylene [2C]	40.000	91.6	80 - 120	5.685	5.687167	-0.0022	N/A	
SLB0342-SCV6 (Water) Lab File ID: 02242318ECD7.D Analyzed: 02/24/23 16:48								
Decachlorobiphenyl	40.000	128	80 - 120	13.893	13.89483	-0.0018	N/A	
Tetrachlorometaxylene	40.000	92.7	80 - 120	5.806	5.8095	-0.0035	N/A	
Decachlorobiphenyl [2C]	40.000	141	80 - 120	14.118	14.11917	-0.0012	N/A	
Tetrachlorometaxylene [2C]	40.000	95.4	80 - 120	5.685	5.687167	-0.0022	N/A	



SURROGATE RECOVERY AND RT SUMMARY EPA 8082A

Laboratory: Analytical Resources, LLC SDG/WO: 23C0107
Client: Anchor QEA, LLC Project: AOC4 UR Phase 3
Sequence: SLC0215 Instrument: ECD7
Calibration: GB00069 Calibration Date: 02/24/2023

Surrogate Compound	Spike Level ug/L	% Recovery	Recovery Limits	RT	Calibration Mean RT	RT Diff	RT Diff Limit	Q
SLC0215-ICV1 (Water)		Lab File ID: 03152306ECD7.D			Analyzed: 03/15/23 14:43			
Decachlorobiphenyl	40.000	77.0	0 - 200	13.898	13.89483	0.0032	N/A	
Tetrachlorometaxylene	40.000	88.8	0 - 200	5.81	5.8095	0.0005	N/A	
Decachlorobiphenyl [2C]	40.000	93.8	0 - 200	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	40.000	89.3	0 - 200	5.691	5.687167	0.0038	N/A	
SLC0215-ICV2 (Water)		Lab File ID: 03152307ECD7.D			Analyzed: 03/15/23 15:04			
Decachlorobiphenyl	40.000	78.8	0 - 200	13.896	13.89483	0.0012	N/A	
Tetrachlorometaxylene	40.000	91.5	0 - 200	5.81	5.8095	0.0005	N/A	
Decachlorobiphenyl [2C]	40.000	95.0	0 - 200	14.121	14.11917	0.0018	N/A	
Tetrachlorometaxylene [2C]	40.000	91.5	0 - 200	5.692	5.687167	0.0048	N/A	
SLC0215-CCV1 (Water)		Lab File ID: 03152313ECD7.D			Analyzed: 03/15/23 17:09			
Decachlorobiphenyl	40.000	71.8	0 - 200	13.896	13.89483	0.0012	N/A	
Tetrachlorometaxylene	40.000	87.8	0 - 200	5.811	5.8095	0.0015	N/A	
Decachlorobiphenyl [2C]	40.000	94.8	0 - 200	14.123	14.11917	0.0038	N/A	
Tetrachlorometaxylene [2C]	40.000	90.0	0 - 200	5.692	5.687167	0.0048	N/A	
SLC0215-CCV2 (Water)		Lab File ID: 03152314ECD7.D			Analyzed: 03/15/23 17:30			
Decachlorobiphenyl	40.000	71.5	0 - 200	13.896	13.89483	0.0012	N/A	
Tetrachlorometaxylene	40.000	89.0	0 - 200	5.81	5.8095	0.0005	N/A	
Decachlorobiphenyl [2C]	40.000	96.3	0 - 200	14.122	14.11917	0.0028	N/A	
Tetrachlorometaxylene [2C]	40.000	90.3	0 - 200	5.692	5.687167	0.0048	N/A	
BLC0219-BLK1 (Solid)		Lab File ID: 03152315ECD7.D			Analyzed: 03/15/23 17:51			
Decachlorobiphenyl	8.0000	67.5	40 - 126	13.895	13.89483	0.0002	N/A	
Tetrachlorometaxylene	8.0000	76.4	44 - 120	5.81	5.8095	0.0005	N/A	
Decachlorobiphenyl [2C]	8.0000	94.6	40 - 126	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	8.0000	76.1	44 - 120	5.692	5.687167	0.0048	N/A	
BLC0219-BS1 (Solid)		Lab File ID: 03152316ECD7.D			Analyzed: 03/15/23 18:12			
Decachlorobiphenyl	8.0000	63.5	40 - 126	13.894	13.89483	-0.0008	N/A	
Tetrachlorometaxylene	8.0000	72.0	44 - 120	5.81	5.8095	0.0005	N/A	
Decachlorobiphenyl [2C]	8.0000	88.6	40 - 126	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	8.0000	71.0	44 - 120	5.692	5.687167	0.0048	N/A	



SURROGATE RECOVERY AND RT SUMMARY
EPA 8082A

Laboratory: Analytical Resources, LLC
Client: Anchor QEA, LLC
Sequence: SLC0215
Calibration: GB00069

SDG/WO: 23C0107
Project: AOC4 UR Phase 3
Instrument: ECD7
Calibration Date: 02/24/2023

Surrogate Compound	Spike Level ug/kg wet	% Recovery	Recovery Limits	RT	Calibration Mean RT	RT Diff	RT Diff Limit	Q
BLC0219-BSD1 (Solid)		Lab File ID: 03152317ECD7.D			Analyzed: 03/15/23 18:32			
Decachlorobiphenyl	8.0000	64.6	40 - 126	13.895	13.89483	0.0002	N/A	
Tetrachlorometaxylene	8.0000	74.4	44 - 120	5.811	5.8095	0.0015	N/A	
Decachlorobiphenyl [2C]	8.0000	91.9	40 - 126	14.121	14.11917	0.0018	N/A	
Tetrachlorometaxylene [2C]	8.0000	72.5	44 - 120	5.691	5.687167	0.0038	N/A	
BLC0219-SRM1 (Solid)		Lab File ID: 03152318ECD7.D			Analyzed: 03/15/23 18:53			
Decachlorobiphenyl	40.000	69.7	40 - 126	13.889	13.89483	-0.0058	N/A	
Tetrachlorometaxylene	40.000	71.4	44 - 120	5.808	5.8095	-0.0015	N/A	
Decachlorobiphenyl [2C]	40.000	79.7	40 - 126	14.116	14.11917	-0.0032	N/A	
Tetrachlorometaxylene [2C]	40.000	74.5	44 - 120	5.689	5.687167	0.0018	N/A	
23C0107-01 (Solid)		Lab File ID: 03152319ECD7.D			Analyzed: 03/15/23 19:14			
Decachlorobiphenyl	7.9847	73.3	40 - 126	13.886	13.89483	-0.0088	N/A	
Tetrachlorometaxylene	7.9847	60.2	44 - 120	5.806	5.8095	-0.0035	N/A	
Decachlorobiphenyl [2C]	7.9847	73.3	40 - 126	14.111	14.11917	-0.0082	N/A	
Tetrachlorometaxylene [2C]	7.9847	67.5	44 - 120	5.686	5.687167	-0.0012	N/A	
23C0107-02 (Solid)		Lab File ID: 03152320ECD7.D			Analyzed: 03/15/23 19:35			
Decachlorobiphenyl	7.9974	85.4	40 - 126	13.891	13.89483	-0.0038	N/A	
Tetrachlorometaxylene	7.9974	76.4	44 - 120	5.809	5.8095	-0.0005	N/A	
Decachlorobiphenyl [2C]	7.9974	88.0	40 - 126	14.117	14.11917	-0.0022	N/A	
Tetrachlorometaxylene [2C]	7.9974	75.2	44 - 120	5.689	5.687167	0.0018	N/A	
23C0107-03 (Solid)		Lab File ID: 03152321ECD7.D			Analyzed: 03/15/23 19:56			
Decachlorobiphenyl	7.9895	70.7	40 - 126	13.892	13.89483	-0.0028	N/A	
Tetrachlorometaxylene	7.9895	72.4	44 - 120	5.809	5.8095	-0.0005	N/A	
Decachlorobiphenyl [2C]	7.9895	86.5	40 - 126	14.118	14.11917	-0.0012	N/A	
Tetrachlorometaxylene [2C]	7.9895	71.1	44 - 120	5.69	5.687167	0.0028	N/A	
BLC0219-MS1 (Solid)		Lab File ID: 03152322ECD7.D			Analyzed: 03/15/23 20:16			
Decachlorobiphenyl	7.9988	71.2	40 - 126	13.893	13.89483	-0.0018	N/A	
Tetrachlorometaxylene	7.9988	75.3	44 - 120	5.809	5.8095	-0.0005	N/A	
Decachlorobiphenyl [2C]	7.9988	87.8	40 - 126	14.117	14.11917	-0.0022	N/A	
Tetrachlorometaxylene [2C]	7.9988	73.2	44 - 120	5.69	5.687167	0.0028	N/A	



SURROGATE RECOVERY AND RT SUMMARY
EPA 8082A

Laboratory: Analytical Resources, LLC
Client: Anchor QEA, LLC
Sequence: SLC0215
Calibration: GB00069

SDG/WO: 23C0107
Project: AOC4 UR Phase 3
Instrument: ECD7
Calibration Date: 02/24/2023

Surrogate Compound	Spike Level ug/kg dry	% Recovery	Recovery Limits	RT	Calibration Mean RT	RT Diff	RT Diff Limit	Q
BLC0219-MSD1 (Solid)			Lab File ID: 03152323ECD7.D			Analyzed: 03/15/23 20:37		
Decachlorobiphenyl	7.9988	69.0	40 - 126	13.893	13.89483	-0.0018	N/A	
Tetrachlorometaxylene	7.9988	75.0	44 - 120	5.809	5.8095	-0.0005	N/A	
Decachlorobiphenyl [2C]	7.9988	88.8	40 - 126	14.118	14.11917	-0.0012	N/A	
Tetrachlorometaxylene [2C]	7.9988	72.2	44 - 120	5.69	5.687167	0.0028	N/A	
SLC0215-CCV3 (Water)			Lab File ID: 03152331ECD7.D			Analyzed: 03/15/23 23:24		
Decachlorobiphenyl	40.000	82.8	0 - 200	13.895	13.89483	0.0002	N/A	
Tetrachlorometaxylene	40.000	104	0 - 200	5.81	5.8095	0.0005	N/A	
Decachlorobiphenyl [2C]	40.000	89.8	0 - 200	14.121	14.11917	0.0018	N/A	
Tetrachlorometaxylene [2C]	40.000	105	0 - 200	5.69	5.687167	0.0028	N/A	
SLC0215-CCV4 (Water)			Lab File ID: 03152332ECD7.D			Analyzed: 03/15/23 23:44		
Decachlorobiphenyl	40.000	91.0	0 - 200	13.897	13.89483	0.0022	N/A	
Tetrachlorometaxylene	40.000	91.8	0 - 200	5.811	5.8095	0.0015	N/A	
Decachlorobiphenyl [2C]	40.000	97.0	0 - 200	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	40.000	91.3	0 - 200	5.691	5.687167	0.0038	N/A	
SLC0215-CCV5 (Water)			Lab File ID: 03152339ECD7.D			Analyzed: 03/16/23 02:10		
Decachlorobiphenyl	40.000	89.8	0 - 200	13.894	13.89483	-0.0008	N/A	
Tetrachlorometaxylene	40.000	85.0	0 - 200	5.811	5.8095	0.0015	N/A	
Decachlorobiphenyl [2C]	40.000	93.5	0 - 200	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	40.000	89.0	0 - 200	5.692	5.687167	0.0048	N/A	
SLC0215-CCV6 (Water)			Lab File ID: 03152340ECD7.D			Analyzed: 03/16/23 02:31		
Decachlorobiphenyl	40.000	89.8	0 - 200	13.895	13.89483	0.0002	N/A	
Tetrachlorometaxylene	40.000	88.8	0 - 200	5.812	5.8095	0.0025	N/A	
Decachlorobiphenyl [2C]	40.000	94.8	0 - 200	14.12	14.11917	0.0008	N/A	
Tetrachlorometaxylene [2C]	40.000	91.3	0 - 200	5.692	5.687167	0.0048	N/A	



INTERNAL STANDARD AREA AND RT SUMMARY
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Sequence: SLB0342

Instrument: ECD7

Calibration: GB00069

Internal Standard	Response	RT	Reference Response	Reference RT	Area %	Area % Limits	RT Diff	RT Diff Limit	Q
Secondary Cal Check (SLB0342-SCV1)		(Water)	Lab File ID: 02242313ECD7.D			Analyzed: 02/24/23 15:03			
1-Bromo-2-Nitrobenzene	645975	3.489	673778	3.493	96	50 - 200	-0.004	+/-0.50	
Hexabromobiphenyl	1524245	14.268	1429847	14.268	107	50 - 200	0.000	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	316115	3.927	315256	3.928	100	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	556950	15.007	513946	15.008	108	50 - 200	-0.001	+/-0.50	
Secondary Cal Check (SLB0342-SCV2)		(Water)	Lab File ID: 02242314ECD7.D			Analyzed: 02/24/23 15:24			
1-Bromo-2-Nitrobenzene	705650	3.493	673778	3.493	105	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl	1555683	14.267	1429847	14.268	109	50 - 200	-0.001	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	340433	3.929	315256	3.928	108	50 - 200	0.001	+/-0.50	
Hexabromobiphenyl [2C]	565609	15.008	513946	15.008	110	50 - 200	0.000	+/-0.50	
Secondary Cal Check (SLB0342-SCV3)		(Water)	Lab File ID: 02242315ECD7.D			Analyzed: 02/24/23 15:45			
1-Bromo-2-Nitrobenzene	646554	3.49	673778	3.493	96	50 - 200	-0.003	+/-0.50	
Hexabromobiphenyl	1529451	14.268	1429847	14.268	107	50 - 200	0.000	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	316066	3.928	315256	3.928	100	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl [2C]	557213	15.008	513946	15.008	108	50 - 200	0.000	+/-0.50	
Secondary Cal Check (SLB0342-SCV4)		(Water)	Lab File ID: 02242316ECD7.D			Analyzed: 02/24/23 16:06			
1-Bromo-2-Nitrobenzene	656887	3.488	673778	3.493	97	50 - 200	-0.005	+/-0.50	
Hexabromobiphenyl	1585505	14.267	1429847	14.268	111	50 - 200	-0.001	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	320936	3.925	315256	3.928	102	50 - 200	-0.003	+/-0.50	
Hexabromobiphenyl [2C]	570006	15.007	513946	15.008	111	50 - 200	-0.001	+/-0.50	
Secondary Cal Check (SLB0342-SCV5)		(Water)	Lab File ID: 02242317ECD7.D			Analyzed: 02/24/23 16:27			
1-Bromo-2-Nitrobenzene	661953	3.489	673778	3.493	98	50 - 200	-0.004	+/-0.50	
Hexabromobiphenyl	1574993	14.268	1429847	14.268	110	50 - 200	0.000	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	317807	3.926	315256	3.928	101	50 - 200	-0.002	+/-0.50	
Hexabromobiphenyl [2C]	565951	15.007	513946	15.008	110	50 - 200	-0.001	+/-0.50	
Secondary Cal Check (SLB0342-SCV6)		(Water)	Lab File ID: 02242318ECD7.D			Analyzed: 02/24/23 16:48			
1-Bromo-2-Nitrobenzene	656592	3.489	673778	3.493	97	50 - 200	-0.004	+/-0.50	
Hexabromobiphenyl	1584453	14.268	1429847	14.268	111	50 - 200	0.000	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	314741	3.926	315256	3.928	100	50 - 200	-0.002	+/-0.50	
Hexabromobiphenyl [2C]	568346	15.007	513946	15.008	111	50 - 200	-0.001	+/-0.50	



INTERNAL STANDARD AREA AND RT SUMMARY
EPA 8082A

Laboratory: Analytical Resources, LLC
Client: Anchor QEA, LLC
Sequence: SLC0215

SDG: 23C0107
Project: AOC4 UR Phase 3
Instrument: ECD7
Calibration: GB00069

Internal Standard	Response	RT	Reference Response	Reference RT	Area %	Area % Limits	RT Diff	RT Diff Limit	Q
Initial Cal Check (SLC0215-ICV1)		(Water)	Lab File ID: 03152306ECD7.D			Analyzed: 03/15/23 14:43			
1-Bromo-2-Nitrobenzene	987631	3.492	987631	3.492	100	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl	2494658	14.275	2494658	14.275	100	50 - 200	0.000	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	360654	3.931	360654	3.931	100	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl [2C]	535523	15.01	535523	15.01	100	50 - 200	0.000	+/-0.50	
Initial Cal Check (SLC0215-ICV2)		(Water)	Lab File ID: 03152307ECD7.D			Analyzed: 03/15/23 15:04			
1-Bromo-2-Nitrobenzene	1002388	3.492	1002388	3.492	100	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl	2626033	14.275	2626033	14.275	100	50 - 200	0.000	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	364771	3.931	364771	3.931	100	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl [2C]	550406	15.011	550406	15.011	100	50 - 200	0.000	+/-0.50	
Blank (BLC0219-BLK1)		(Solid)	Lab File ID: 03152315ECD7.D			Analyzed: 03/15/23 17:51			
1-Bromo-2-Nitrobenzene	1146280	3.494	1002388	3.492	114	50 - 200	0.002	+/-0.50	
Hexabromobiphenyl	3433528	14.274	2626033	14.275	131	50 - 200	-0.001	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	400044	3.931	364771	3.931	110	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl [2C]	632392	15.01	550406	15.011	115	50 - 200	-0.001	+/-0.50	
LCS (BLC0219-BS1)		(Solid)	Lab File ID: 03152316ECD7.D			Analyzed: 03/15/23 18:12			
1-Bromo-2-Nitrobenzene	1175579	3.494	1002388	3.492	117	50 - 200	0.002	+/-0.50	
Hexabromobiphenyl	3282396	14.273	2626033	14.275	125	50 - 200	-0.002	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	406961	3.931	364771	3.931	112	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl [2C]	639731	15.008	550406	15.011	116	50 - 200	-0.003	+/-0.50	
LCS Dup (BLC0219-BSD1)		(Solid)	Lab File ID: 03152317ECD7.D			Analyzed: 03/15/23 18:32			
1-Bromo-2-Nitrobenzene	1147217	3.494	1002388	3.492	114	50 - 200	0.002	+/-0.50	
Hexabromobiphenyl	3363429	14.274	2626033	14.275	128	50 - 200	-0.001	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	396521	3.931	364771	3.931	109	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl [2C]	636625	15.008	550406	15.011	116	50 - 200	-0.003	+/-0.50	
Reference (BLC0219-SRM1)		(Solid)	Lab File ID: 03152318ECD7.D			Analyzed: 03/15/23 18:53			
1-Bromo-2-Nitrobenzene	1103257	3.493	1002388	3.492	110	50 - 200	0.001	+/-0.50	
Hexabromobiphenyl	1949653	14.262	2626033	14.275	74	50 - 200	-0.013	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	380915	3.93	364771	3.931	104	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	540684	15.002	550406	15.011	98	50 - 200	-0.009	+/-0.50	



INTERNAL STANDARD AREA AND RT SUMMARY
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor OEA, LLC

Project: AOC4 UR Phase 3

Sequence: SLC0215

Instrument: ECD7

Calibration: GB00069

Internal Standard	Response	RT	Reference Response	Reference RT	Area %	Area % Limits	RT Diff	RT Diff Limit	Q
LDW22-SS826 (23C0107-01)		(Solid)	Lab File ID: 03152319ECD7.D			Analyzed: 03/15/23 19:14			
1-Bromo-2-Nitrobenzene	1011772	3.492	1002388	3.492	101	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl	1031369	14.254	2626033	14.275	39	50 - 200	-0.021	+/-0.50	*
1-Bromo-2-Nitrobenzene [2C]	371913	3.93	364771	3.931	102	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	410686	14.997	550406	15.011	75	50 - 200	-0.014	+/-0.50	
LDW21-IT608C (23C0107-02)		(Solid)	Lab File ID: 03152320ECD7.D			Analyzed: 03/15/23 19:35			
1-Bromo-2-Nitrobenzene	1015111	3.492	1002388	3.492	101	50 - 200	0.000	+/-0.50	
Hexabromobiphenyl	1524021	14.263	2626033	14.275	58	50 - 200	-0.012	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	387409	3.93	364771	3.931	106	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	492712	15.004	550406	15.011	90	50 - 200	-0.007	+/-0.50	
LDW21-IT608D (23C0107-03)		(Solid)	Lab File ID: 03152321ECD7.D			Analyzed: 03/15/23 19:56			
1-Bromo-2-Nitrobenzene	1104252	3.493	1002388	3.492	110	50 - 200	0.001	+/-0.50	
Hexabromobiphenyl	2211995	14.268	2626033	14.275	84	50 - 200	-0.007	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	405699	3.93	364771	3.931	111	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	556397	15.006	550406	15.011	101	50 - 200	-0.005	+/-0.50	
Matrix Spike (BLC0219-MS1)		(Solid)	Lab File ID: 03152322ECD7.D			Analyzed: 03/15/23 20:16			
1-Bromo-2-Nitrobenzene	1102984	3.493	1002388	3.492	110	50 - 200	0.001	+/-0.50	
Hexabromobiphenyl	2318945	14.268	2626033	14.275	88	50 - 200	-0.007	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	399893	3.93	364771	3.931	110	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	571920	15.007	550406	15.011	104	50 - 200	-0.004	+/-0.50	
Matrix Spike Dup (BLC0219-MSD1)		(Solid)	Lab File ID: 03152323ECD7.D			Analyzed: 03/15/23 20:37			
1-Bromo-2-Nitrobenzene	1094891	3.493	1002388	3.492	109	50 - 200	0.001	+/-0.50	
Hexabromobiphenyl	2334158	14.267	2626033	14.275	89	50 - 200	-0.008	+/-0.50	
1-Bromo-2-Nitrobenzene [2C]	388222	3.93	364771	3.931	106	50 - 200	-0.001	+/-0.50	
Hexabromobiphenyl [2C]	560920	15.007	550406	15.011	102	50 - 200	-0.004	+/-0.50	



DUAL COLUMN CONFIRMATION SUMMARY

Laboratory: Analytical Resources, LLC SDG: 23C0107
Client: Anchor OEA, LLC Project: AOC4 UR Phase 3
Matrix: Sediment Laboratory ID: 23C0107-01 File ID: 03152319ECD7.D
Sampled: 03/02/23 16:25 Prepared: 03/10/23 11:58 Analyzed: 03/15/23 19:14
Solids: 41.90 Preparation: EPA 3546 (Microwave) Instrument: ECD7
Batch: BLC0219 Sequence: SLC0215
GC Column(1): ZB5 GC Column(2): ZB35

COMPOUND	COL	RT	EXP RT	RT DIFF	AREA	CONC	RPD
Aroclor 1248	* 1	8.4	8.405	0.005	60257.25	17.1	15.8
	2	8.301	8.307	0.006	15777.5	14.6	
Aroclor 1254	1	9.287	9.298	0.011	97090.2	23.1	30.5
	* 2	9.44	9.449	0.009	56646.6	31.4	
Aroclor 1260	1	11.032	11.04467	0.0127	71879.4	29.3	7.2
	* 2	11.643	11.6535	0.0105	48874.5	31.5	

* Column used for quantitation



HOLDING TIME SUMMARY

Analysis: EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Sample Name	Date Collected	Date Received	Date Prepared	Days to Prep	Max Days to Prep	Date Analyzed	Days to Analysis	Max Days to Analysis	Q
LDW22-SS826 23C0107-01	03/02/23 16:25	03/03/23 16:35	03/10/23 11:58	7	365	03/15/23 19:14	5	40	
LDW21-IT608C 23C0107-02	07/13/21 06:54	03/03/23 16:35	03/10/23 11:58	605	365	03/15/23 19:35	5	40	*
LDW21-IT608D 23C0107-03	07/13/21 06:54	03/03/23 16:35	03/10/23 11:58	605	365	03/15/23 19:56	5	40	*
Matrix Spike BLC0219-MS1	07/13/21 06:54	03/03/23 16:35	03/10/23 11:58	605	365	03/15/23 20:16	5	40	*
Matrix Spike Dup BLC0219-MSD1	07/13/21 06:54	03/03/23 16:35	03/10/23 11:58	605	365	03/15/23 20:37	5	40	*

* Indicates hold time exceedance.



**METHOD DETECTION
AND REPORTING LIMITS**
EPA 8082A

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Instrument: ECD7

Analyte	MDL	RL	Units
Aroclor 1016	1.6	4.0	ug/kg
Aroclor 1016 [2C]	1.6	4.0	ug/kg
Aroclor 1221	1.6	4.0	ug/kg
Aroclor 1221 [2C]	1.6	4.0	ug/kg
Aroclor 1232	1.6	4.0	ug/kg
Aroclor 1232 [2C]	1.6	4.0	ug/kg
Aroclor 1242	1.6	4.0	ug/kg
Aroclor 1242 [2C]	1.6	4.0	ug/kg
Aroclor 1248	1.6	4.0	ug/kg
Aroclor 1248 [2C]	1.6	4.0	ug/kg
Aroclor 1254	1.6	4.0	ug/kg
Aroclor 1254 [2C]	1.6	4.0	ug/kg
Aroclor 1260	0.6	4.0	ug/kg
Aroclor 1260 [2C]	0.6	4.0	ug/kg

CERTIFICATE OF ANALYSIS

Catalog No: S-279N
Description: Tetrachloro-m-xylene
Lot: 0052481B-1
Solvent: N/A
Hazards: Refer to SDS for complete safety information

Date Certified: Jul 28, 2005
Expiration: Jul 28, 2015
Sample Size: 100 mg
Components: 1
Storage Condition: Ambient (>5 °C)



Signal Word: Warning

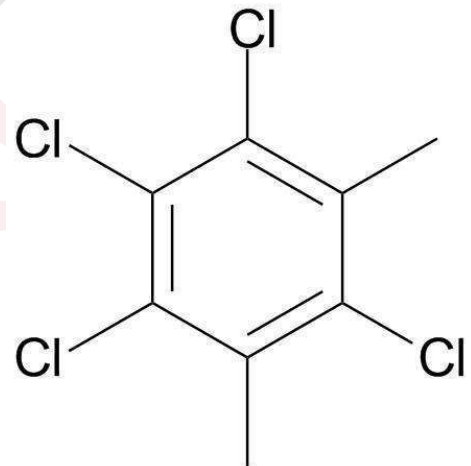
Certified Reference Material



Component	CAS #	Purity % (GC/FID)	Prepared Concentration	Certified Analyte Concentration ¹
Tetrachloro-meta-xylene	877-09-8	96.0	N/A	N/A

Identification:

Molecular formula: C₈H₆Cl₄
Molecular weight: 243.94



C000147

tetrachlorometaxylene

Expires 1/15/2020

Prepared By Joshua Rains 1/15/2014

This Certified Reference Material was verified in accordance with ISO/IEC 17025

A product with a suffix (-1A, -2B, etc. or -01, -02, etc.) on its lot number has had its expiration date extended and is identical to the same lot number without the suffix.

¹ The Uncertainty calculated for this product is ±2.4%. These values are the expanded uncertainty and represent an estimated standard deviation equal to the positive square root of the total variation of the uncertainty of components. A normal distribution is assumed and a coverage factor of K=2 is chosen using approximately a 95% confidence level.

Labels and certificates follow U.S. Conventions in reporting numerical values: A comma (,) is used to separate units of one-thousand or greater. A period (.) is used as a decimal place marker.

Metrological traceability is established through in-house validated methods.

Purity, if stated, is equal to 100% minus found impurity components. Impurity components have not been identified.

The information on this certificate may not be reproduced without the express permission of the manufacturer. See reverse side for additional information

Hazard Information: Please refer to the SDS for information regarding the hazards associated with using this material.

This product was prepared according to in-house procedures and is guaranteed to be homogeneous.

Certified By:

Larry Decker, Organic QC Manager



AccuStandard

125 Market Street
New Haven, CT 06513
(203) 786-5290

CERTIFICATE OF PRODUCT DATA

PRODUCT: C-209N

EXPIRATION: Jul 28, 2015

DESCRIPTION: 2,2',3,3',4,4',5,5',6,6'-Decachlorobiphenyl

LOT #: 990521LB-AC

SOLVENT: N/A

This product is guaranteed accurate to $\pm 0.5\%$ of the Certified Analyte concentration through the Expiration Date on the Label.

Component	CAS #	Purity % (GC/MS)	Prepared Concentration ¹	Certified Analyte Concentration ²
2,2',3,3',4,4',5,5',6,6'-Decachlorobiphenyl	2051-24-3	100	N/A	N/A

2;

C000148

decachlorobiphenyl
Expires 1/15/2020

Prepared By Joshua Rains 1/15/2014

** I 1768 A*

Certified by: *R. Cooper*

Please note: AccuStandard follows the U.S. conventions in reporting numerical values, on both certificates and labels.

A comma (,) is used to separate units of one-thousand or greater.
A period (.) is used as a decimal place marker.

1. All weights are traceable through National Institute of Standards & Technology, Test No. 822/254480
 2. Certified Analyte Concentration = Purity x Prepared Concentration. The Uncertainty calculated for this product is $\pm 0.5\%$ which is the Combined Uncertainty $U_c(y)$. It represents an estimated standard deviation equal to the positive square root of the total variance of the uncertainty of components. The Expanded Uncertainty is U which is $U_c(y) * K$ where K is the coverage factor at the 95% confidence level ($K=2$).
 3. A product with a suffix (-1A, -2B, etc.) on its lot# has had its expiration date extended and is identical to the same lot# without the suffix.

This product was manufactured in accordance to quality system requirements of ISO 9001:2000 and ISO 17025

** Recertified ~ 4-6-09 (S)*



Analytical Standard Record
Standard ID: C000148

Printed: 4/23/2015 11:54:44AM

Description:	decachlorobiphenyl	Expires:	15-Jan-2020
Standard Type:	Other	Prepared:	15-Jan-2014
Solvent:	na/a	Prepared By:	Joshua Rains
Final Volume (mls):	1	Department:	Organics
Vials:	1	Last Edit:	27-Feb-2015 13:03 by JGR
Vendor:	Accustandard	Lot #:	9905211b-ac
Vendor Catalog #:			

Comments

see i1768a
SOM calibrations added 06/12/14 sdrd

Analyte	CAS Number	Concentration	Units
Decachlorobiphenyl [2C]	2051-24-3	1000000	ug/mL
Decachlorobiphenyl	2051-24-3	1000000	ug/mL
DCB 1660 [2C]	2051-24-3	1000000	ug/mL
DCB 1660	2051-24-3	1000000	ug/mL
DCB [2C]	2051-24-3	1000000	ug/mL
DCB (A) [2C]	2051-24-3	1000000	ug/mL
DCB (A)	2051-24-3	1000000	ug/mL
DCB	2051-24-3	1000000	ug/mL

Reviewed By

Date

Certificate of Analysis



Phenova Certified Reference Materials are sold by Phenomenex.

411 Madrid Ave., Torrance, CA 90501 USA ■ Tel: 310-212-0555 ■ Fax: 310-328-7768 ■ info@phenomenex.com

Access your MSDS and digital C of A at www.phenomenex.com/mysupport. Re-order at www.phenomenex.com/standards

Certified Reference Material

This product is included in Phenova's ISO/IEC 17025 and ISO Guide 34 Scopes of Accreditation

Catalog No.: AL0-101461

Lot Number: CL13053

Description: Aroclor 1254

Certification Date: November 29, 2018

Storage: 4 °C

Expiration Date: November 30, 2026

Provided As: 1 mL in 2 mL Ampoule in Hexane

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1254	11097-69-1	1000	± 0.246%

I 09808
Recd.
02/24/20



Reference Material Producer
Certificate No. 2427.02



Manufactured by Phenova, Inc.

Phenova's testing and calibration results are internationally recognized through the ILAC-MRA. Phenova is an accredited ISO Guide 34 Reference Material Provider and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03

Certificate of Analysis

Produced by Phenova

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Access your Safety Data Sheets and digital Certificates at www.phenova.com/documents.

Certified Reference Material

This product is certified in accordance with Phenova's ISO 17034 accreditation and supported by Phenova's ISO/IEC 17025 chemical testing accreditation

Catalog No.: AL0-101468

Lot Number: CL14017

Description: Aroclor 1221

Certification Date: August 20, 2019

Storage: 4 °C

Expiration Date: August 31, 2027

Provided As: 1 mL in 2 mL Ampoule in Isooctane

Andrea Gill

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1221	11104-28-2	1000	± 0.553%

J006466
Recd of
06/18/21



Reference Material Producer
Certificate No. 2427.02



Phenova is an accredited ISO/IEC 17034 Reference Material
Producer and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03

Certificate of Analysis



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1. **Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
2. **Quality Standards:** Phenova is accredited by A2LA to ISO 17034³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
3. **Intended Use:** The product is manufactured for use in calibration, calibration verification, quantification, identification and other appropriate analytical control applications. The product is intended for routine laboratory analysis and research purposes only. Only trained personnel should handle this product.
4. **Handling and Usage Notes:** Store according to recommended conditions listed and avoid prolonged exposure to light. Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all analytes in the mixture. Considerations should be made related to repeated use of the opened product. Once opened, exposure to light, air, heat, objects, and additional transfer vessels may cause evaporation, degradation or contamination resulting in changes in concentration, uncertainty and stability duration. Store opened standards in a clean, tightly capped vessel under the recommended temperature. Appropriate controls, such as the use of additional verification standards should be used to confirm the opened product is fit for purpose under repeated use conditions.
5. **Hazardous Situation:** The product is intended for use by experienced professional personnel. A Safety Data Sheet (SDS) is available at www.phenova.com/documents.
6. **Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
7. **Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
8. **Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
9. **Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).

$$u_{CRM} = k \sqrt{u_M^2 + u_H^2 + u_{LTS}^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.

10. **Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO 17034. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
11. **Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO 17034.
12. **Period of Validity:** The Certified Values, Uncertainties and Expiration Date are based on the unopened product being stored according to the recommended storage condition listed and are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

¹ ISO Guide 31 – Reference Materials – Contents of Certificates and Labels.

² ISO Guide 35 – Reference Material – General and Statistical Principles for Certification.

³ ISO 17034 – General Requirements for the Competence of Reference Material Producers.

⁴ ISO/IEC 17025 – General Requirements for the Competence of Testing and Calibration Laboratories.

⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



Reference Material Producer
Certificate No. 2427.02



Phenova is an accredited ISO/IEC 17034 Reference Material
Producer and ISO/IEC 17025 accredited Chemical Testing Laboratory.

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Company



Chemical Testing Laboratory
Certificate No. 2427.03

Certificate of Analysis

Produced by Phenova

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Access your Safety Data Sheets and digital Certificates at www.phenova.com/documents.

Certified Reference Material

This product is certified in accordance with Phenova's ISO 17034 accreditation and supported by Phenova's ISO/IEC 17025 chemical testing accreditation

Catalog No.: AL0-101469

Lot Number: CL14914

Description: Aroclor 1232

Certification Date: January 31, 2020

Storage: 4 °C

Expiration Date: January 31, 2028

Provided As: 1 mL in 2 mL Ampoule in Isooctane



Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1232	11141-16-5	1000	± 0.738%

J 006467
read
06/18/21



Reference Material Producer
Certificate No. 2427.02



Phenova is an accredited ISO/IEC 17034 Reference Material Producer and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03

Certificate of Analysis



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1. **Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
2. **Quality Standards:** Phenova is accredited by A2LA to ISO 17034³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
3. **Intended Use:** The product is manufactured for use in calibration, calibration verification, quantification, identification and other appropriate analytical control applications. The product is intended for routine laboratory analysis and research purposes only. Only trained personnel should handle this product.
4. **Handling and Usage Notes:** Store according to recommended conditions listed and avoid prolonged exposure to light. Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all analytes in the mixture. Considerations should be made related to repeated use of the opened product. Once opened, exposure to light, air, heat, objects, and additional transfer vessels may cause evaporation, degradation or contamination resulting in changes in concentration, uncertainty and stability duration. Store opened standards in a clean, tightly capped vessel under the recommended temperature. Appropriate controls, such as the use of additional verification standards should be used to confirm the opened product is fit for purpose under repeated use conditions.
5. **Hazardous Situation:** The product is intended for use by experienced professional personnel. A Safety Data Sheet (SDS) is available at www.phenova.com/documents.
6. **Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
7. **Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
8. **Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
9. **Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).

$$uCRM = k\sqrt{uM^2 + uH^2 + uLTS^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.

10. **Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO 17034. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
11. **Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO 17034.
12. **Period of Validity:** The Certified Values, Uncertainties and Expiration Date are based on the unopened product being stored according to the recommended storage condition listed and are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

- ¹ ISO Guide 31 – Reference Materials – Contents of Certificates and Labels.
- ² ISO Guide 35 – Reference Material – General and Statistical Principles for Certification.
- ³ ISO 17034 – General Requirements for the Competence of Reference Material Producers.
- ⁴ ISO/IEC 17025 – General Requirements for the Competence of Testing and Calibration Laboratories.
- ⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



Reference Material Producer
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Phenova is an accredited ISO/IEC 17034 Reference Material
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Certified Reference Material

This product is certified in accordance with Phenova's ISO 17034 accreditation and supported by Phenova's ISO/IEC 17025 chemical testing accreditation

Catalog No.: AL0-101470

Lot Number: CL14018

Description: Aroclor 1242

Certification Date: August 20, 2019

Storage: 4 °C

Expiration Date: August 31, 2027

Provided As: 1 mL in 2 mL Ampoule in Isooctane

Andrea Gill

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1242	53469-21-9	1000	± 0.553%

J006468
feed JR
06/18/21



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1. **Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
2. **Quality Standards:** Phenova is accredited by A2LA to ISO 17034³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
3. **Intended Use:** The product is manufactured for use in calibration, calibration verification, quantification, identification and other appropriate analytical control applications. The product is intended for routine laboratory analysis and research purposes only. Only trained personnel should handle this product.
4. **Handling and Usage Notes:** Store according to recommended conditions listed and avoid prolonged exposure to light. Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all analytes in the mixture. Considerations should be made related to repeated use of the opened product. Once opened, exposure to light, air, heat, objects, and additional transfer vessels may cause evaporation, degradation or contamination resulting in changes in concentration, uncertainty and stability duration. Store opened standards in a clean, tightly capped vessel under the recommended temperature. Appropriate controls, such as the use of additional verification standards should be used to confirm the opened product is fit for purpose under repeated use conditions.
5. **Hazardous Situation:** The product is intended for use by experienced professional personnel. A Safety Data Sheet (SDS) is available at www.phenova.com/documents.
6. **Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
7. **Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
8. **Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
9. **Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).

$$u_{CRM} = k\sqrt{u_M^2 + u_H^2 + u_{LTS}^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.

10. **Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO 17034. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
11. **Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO 17034.
12. **Period of Validity:** The Certified Values, Uncertainties and Expiration Date are based on the unopened product being stored according to the recommended storage condition listed and are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

¹ ISO Guide 31 – Reference Materials – Contents of Certificates and Labels.

² ISO Guide 35 – Reference Material – General and Statistical Principles for Certification.

³ ISO 17034 – General Requirements for the Competence of Reference Material Producers.

⁴ ISO/IEC 17025 – General Requirements for the Competence of Testing and Calibration Laboratories.

⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



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Certified Reference Material

This product is certified in accordance with Phenova's ISO 17034 accreditation and supported by Phenova's ISO/IEC 17025 chemical testing accreditation

Catalog No.: AL0-101471

Lot Number: CL15384

Description: Aroclor 1248

Certification Date: June 19, 2020

Storage: 4 °C

Expiration Date: June 30, 2028

Provided As: 1 mL in 2 mL Ampoule in Isooctane

Andrea L Gill

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1248	12672-29-6	1000	± 0.520%

*# J006469
Reed, JR
06/18/21*



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- 1. Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
- 2. Quality Standards:** Phenova is accredited by A2LA to ISO 17034³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
- 3. Intended Use:** The product is manufactured for use in calibration, calibration verification, quantification, identification and other appropriate analytical control applications. The product is intended for routine laboratory analysis and research purposes only. Only trained personnel should handle this product.
- 4. Handling and Usage Notes:** Store according to recommended conditions listed and avoid prolonged exposure to light. Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all analytes in the mixture. Considerations should be made related to repeated use of the opened product. Once opened, exposure to light, air, heat, objects, and additional transfer vessels may cause evaporation, degradation or contamination resulting in changes in concentration, uncertainty and stability duration. Store opened standards in a clean, tightly capped vessel under the recommended temperature. Appropriate controls, such as the use of additional verification standards should be used to confirm the opened product is fit for purpose under repeated use conditions.
- 5. Hazardous Situation:** The product is intended for use by experienced professional personnel. A Safety Data Sheet (SDS) is available at www.phenova.com/documents.
- 6. Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
- 7. Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
- 8. Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
- 9. Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).
$$u_{CRM} = k\sqrt{u_M^2 + u_H^2 + u_{LTS}^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.
- 10. Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO 17034. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
- 11. Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO 17034.
- 12. Period of Validity:** The Certified Values, Uncertainties and Expiration Date are based on the unopened product being stored according to the recommended storage condition listed and are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

- ¹ ISO Guide 31 – Reference Materials – Contents of Certificates and Labels.
- ² ISO Guide 35 – Reference Material – General and Statistical Principles for Certification.
- ³ ISO 17034 – General Requirements for the Competence of Reference Material Producers.
- ⁴ ISO/IEC 17025 – General Requirements for the Competence of Testing and Calibration Laboratories.
- ⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



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Certified Reference Material

This product is included in Phenova's ISO/IEC 17025 and ISO Guide 34 Scopes of Accreditation

Catalog No.: AL0-101474

Lot Number: CL11330

Description: Aroclor 1262

Certification Date: May 15, 2015

Storage: 4 °C

Expiration Date: April 30, 2023

Provided As: 1 mL in 2 mL Ampoule in Isooctane

Revision Date: April 2, 2018



Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1262	37324-23-5	1000	± 0.516%

J 00647H
Reed JK
06/18/21



Reference Material Producer
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1. **Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
 2. **Quality Standards:** Phenova is accredited by A2LA to ISO Guide 34³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
 3. **Intended Use:** The product is manufactured for use in the calibration and calibration verification of chromatographic instrumentation performed in routine laboratory analysis.
 4. **Instruction:** Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all certified analytes in the mixture.
 5. **Hazardous Situation:** The product is intended for use by experienced professional personnel. A Material Safety Data Sheet (MSDS) is available at www.phenomenex.com/mysupport.
 6. **Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
 7. **Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
 8. **Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
 9. **Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).
$$uCRM = k\sqrt{uM^2 + uH^2 + uLTS^2}$$
- Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.
10. **Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO Guide 34. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
 11. **Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO Guide 34.
 12. **Period of Validity:** The Certified Values and their uncertainties are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

- ¹ ISO Guide 31:2000(E) – Reference Materials – Contents of Certificates and Labels.
- ² ISO Guide 35:2006(E) – Reference Material – General and Statistical Principles for Certification.
- ³ ISO Guide 34:2009(E) – General Requirements for the Competence of Reference Material Producers.
- ⁴ ISO/IEC 17025:2005(E) – General Requirements for the Competence of Testing and Calibration Laboratories.
- ⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



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Certified Reference Material

This product is included in Phenova's ISO/IEC 17025 and ISO Guide 34 Scopes of Accreditation

Catalog No.: AL0-101475

Lot Number: CL11331

Description: Aroclor 1268

Certification Date: May 15, 2015

Storage: 4 °C

Expiration Date: April 30, 2023

Provided As: 1 mL in 2 mL Ampoule in Isooctane

Revision Date: April 2, 2018

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1268	11100-14-4	1000	± 0.516%

J006472
Rec'd. JK
06/18/21



Reference Material Producer
Certificate No. 2427.02



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2. **Quality Standards:** Phenova is accredited by A2LA to ISO Guide 34³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
3. **Intended Use:** The product is manufactured for use in the calibration and calibration verification of chromatographic instrumentation performed in routine laboratory analysis.
4. **Instruction:** Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all certified analytes in the mixture.
5. **Hazardous Situation:** The product is intended for use by experienced professional personnel. A Material Safety Data Sheet (MSDS) is available at www.phenomenex.com/mysupport.
6. **Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
7. **Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
8. **Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
9. **Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).

$$uCRM = k \cdot \sqrt{uM^2 + uH^2 + uLTS^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.

10. **Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO Guide 34. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
11. **Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO Guide 34.
12. **Period of Validity:** The Certified Values and their uncertainties are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

¹ ISO Guide 31:2000(E) – Reference Materials – Contents of Certificates and Labels.

² ISO Guide 35:2006(E) – Reference Material – General and Statistical Principles for Certification.

³ ISO Guide 34:2009(E) – General Requirements for the Competence of Reference Material Producers.

⁴ ISO/IEC 17025:2005(E) – General Requirements for the Competence of Testing and Calibration Laboratories.

⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



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Certificate of Analysis

Aroclor 1016 Solution

Product Number: PP-282

Page: 1 of 1

Lot Number: CR-0761

Lot Issue Date: 28-Feb-2017

Expiration Date: 31-Mar-2025

This ISO Guide 34 Reference Material (RM) was manufactured and verified in accordance with ULTRA's ISO 9001 registered quality system, and the analyte concentrations were verified by our ISO 17025 accredited laboratory. The true value and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	True Value
Aroclor 1016	012674-11-2	NT01016	100.2 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

Storage: Store at Room Temperature (15° to 30°C).

*K1254
Rec'd JP
02/05/17*

ULTRA uses balances calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1 and ISO 9001, and calibrated Class A glassware in the manufacturing of these standards.



ISO 9001
Registered
TUV USA, Inc.

John Russo
President

Monica Bourgeois
Director of QA/RA



Certificate of Analysis

Product Name: Aroclor 1260 Standard

Product Number: PP-362-1

Lot Issue Date: 20-Jan-2021

Lot Number: 0006582048

Expiration Date: 28-Feb-2025

Description:

This analytical reference material (RM) was manufactured and verified in accordance with an ISO 9001 registered quality system and analyte concentrations were verified by an ISO 17025 accredited laboratory. The concentration and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	Concentration ± Uncertainty
Aroclor 1260	011096-82-5	NT01023	100.4 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

K 1255

Storage Conditions: Store at Room Temperature (15° to 30°C).

Traceability:

The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCCL Z540.3, ISO 9001, ISO 17025, and ISO 17034. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 1088.

Homogeneity:

This RM was unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods, and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening the container and should be processed without delay for the certified values to be valid within the stated uncertainties.

Hazards:

Refer to the Safety Data Sheet on www.agilent.com for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.

Maintenance of Certification:

If substantive changes are noted that affect the certification before the expiration of this certificate, Agilent will notify the purchaser.

Sample lot approver:

Monica Bourgeois

QMS Representative



ISO 17034 Cert
No. AR-1936

RM was produced in accordance with TUV USA Inc registered ISO 9001 Quality Management System. Cert # 56 100 18560026

Page: 1 of 1

www.agilent.com/quality/
CSD-QA-015.1



ISO 17025 Cert
No. AT-1937



Certificate of Analysis ISO Guide 34

Aroclor 1242 Solution

Product Number: PP-312

Page: 1 of 1

Lot Number: CS-6293

Lot Issue Date: 04-Jan-2019

Expiration Date: 31-Jan-2023

This ISO Guide 34 Reference Material (RM) was manufactured and verified in accordance with Agilent's ISO 9001 registered quality system, and the analyte concentrations were verified by our ISO 17025 accredited laboratory. The true value and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	True Value
Aroclor 1242	053469-21-9	NT01020	100.4 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

Storage: Store at Room Temperature (15° to 30°C).

K1256

Agilent uses balances calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1 and ISO 9001, and calibrated Class A glassware in the manufacturing of these standards.


Monica Bourgeois
QMS Representative



ISO Guide 34 Cert No.
AR-1936

Produced in accordance with TUV USA Inc 56 100 18560026
registered ISO 9001 Quality Management System



ISO17025 Cert No.
AT-1937

ISO 17034



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Reference Material Certificate

Product Name: Aroclor 1248 Standard **Lot Number:** 0006626997
Product Number: PP-342-1 **Lot Issue Date:** 17-Aug-2021
Storage Conditions: Store at Room Temperature (15° to 30°C). **Expiration Date:** 30-Sep-2025

Component Name	CERTIFIED VALUES			CAS#	Analyte Lot
	Concentration	Expanded Uncertainty			
Aroclor 1248	100.3	± 0.5 µg/mL		012672-29-6	NT01582

Matrix: isooctane (2,2,4-trimethylpentane)

K1257

Description:

This document is prepared in accordance with ISO 17034 and Guide 31. This analytical reference material standard was manufactured and verified in accordance with an ISO 9001 registered quality system and analyte concentrations were verified by an ISO 17025 accredited laboratory. The concentration and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed above.

Traceability:

The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z540.3, ISO 9001, ISO 17025, and ISO 17034. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 1088.

Homogeneity:

This analytical reference standard was unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening the container and should be processed without delay for the certified values to be valid within the stated uncertainties.

Safety:

Refer to the Safety Data Sheet on www.agilent.com for information regarding this analytical reference material.

Intended Use:

This analytical reference standard is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods, and continuing calibration verification.

Expiration of Certification:

The certification of this analytical reference standard is valid until the expiration date specified above, provided the material is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the material is damaged, contaminated, or otherwise modified.



Certificate of Analysis

Aroclor 1254 Solution

Product Number: PP-352

Page: 1 of 1

Lot Number: CS-2321

Lot Issue Date: 04-May-2018

Expiration Date: 31-May-2026

This ISO Guide 34 Reference Material (RM) was manufactured and verified in accordance with ULTRA's ISO 9001 registered quality system, and the analyte concentrations were verified by our ISO 17025 accredited laboratory. The true value and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	True Value
Aroclor 1254	011097-69-1	RM00922	100.4 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

Storage: Store at Room Temperature (15° to 30°C).

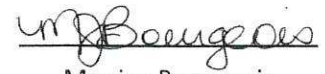
K-1250

ULTRA uses balances calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1 and ISO 9001, and calibrated Class A glassware in the manufacturing of these standards.



ISO 9001
Registered
TUV USA, Inc.


John Russo
President


Monica Bourgeois
Director of QA/RA



Certificate of Analysis

Product Name: Aroclor 1221 Standard

Product Number: PP-292-1

Lot Issue Date: 28-Apr-2020

Lot Number: 0006535333

Expiration Date: 31-May-2024

Description:

This analytical reference material (RM) was manufactured and verified in accordance with an ISO 9001 registered quality system, and analyte concentrations were verified by an ISO 17025 accredited laboratory. The concentration and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	Concentration ± Uncertainty
Aroclor 1221	011104-28-2	RM04278	100.2 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

Storage Conditions: Store at Room Temperature (15° to 30°C).

K1259

Traceability:

The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCCL Z540.3, ISO 9001, ISO 17025, and ISO 17034. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 1088.

Homogeneity:

This RM was unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods, and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening the container and should be processed without delay for the certified values to be valid within the stated uncertainties.

Hazards:

Refer to the Safety Data Sheet on www.agilent.com for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.

Maintenance of Certification:

If substantive changes are noted that affect the certification before the expiration of this certificate, Agilent will notify the purchaser.

Sample lot approver:

Monica Bourgeois
QMS Representative



ISO 17034 Cert No.
AR-1936

RM was produced in accordance with TUV USA Inc registered ISO 9001 Quality Management System. Cert # 56 100 18560026

Page: 1 of 1

www.agilent.com/quality/
CSD-QA-015.1



ISO 17025 Cert
No. AT-1937



Certificate of Analysis ISO 17034

Aroclor 1262 Standard

Product Number: PP-372-1

Page: 1 of 1

Lot Number: 0006499800

Lot Issue Date: 04-Nov-2019

Expiration Date: 30-Nov-2023

This ISO 17034 Reference Material (RM) was manufactured and verified in accordance with Agilent Technologies ISO 9001 registered quality system. A review of the gravimetric preparation data by our ISO 17025 accredited laboratory serves to verify the concentration of each analyte. The true value and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	True Value
Aroclor 1262	037324-23-5	RM14263	100.0 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

Storage: Store at Room Temperature (15° to 30°C).

K1260

Agilent uses balances calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1 and ISO 9001, and calibrated Class A glassware in the manufacturing of these standards.


Monica Bourgeois
QMS Representative



ISO 17034 Cert No.
AR-1936

Produced in accordance with TUV USA Inc 56 100 18560026
registered ISO 9001 Quality Management System



ISO 17025 Cert No.
AT-1937



Certificate of Analysis ISO 17034

Aroclor 1232 Standard

Product Number: PP-302-1

Page: 1 of 1

Lot Number: CF-2197A

Lot Issue Date: 05-Jul-2016

Expiration Date: 31-Aug-2023

This ISO 17034 Reference Material (RM) was manufactured and verified in accordance with Agilent's ISO 9001 registered quality system, and the analyte concentrations were verified by our ISO 17025 accredited laboratory. The true value and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	True Value
Aroclor 1232	011141-16-5	NT01717	100.4 ± 0.5 µg/mL

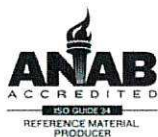
Matrix: isooctane (2,2,4-trimethylpentane)

Storage: Store at Room Temperature (15° to 30°C).

K1261

Agilent uses balances calibrated with weights traceable to NIST in compliance with ANSI/NCCL Z-540-1 and ISO 9001, and calibrated Class A glassware in the manufacturing of these standards.


Monica Bourgeois
QMS Representative



ISO 17034 Cert No.
AR-1936

Produced in accordance with TUV USA Inc 56 100 18560026
registered ISO 9001 Quality Management System



ISO17025 Cert No.
AT-1937



Certificate of Analysis

Product Name: Aroclor 1268 Standard

Product Number: PP-382-1

Lot Issue Date: 09-Feb-2021

Lot Number: 0006587800

Expiration Date: 31-Mar-2029

Description:

This analytical reference material (RM) was manufactured and verified in accordance with an ISO 9001 registered quality system and analyte concentrations were verified by an ISO 17025 accredited laboratory. The concentration and uncertainty value at the 95% confidence level for each analyte, determined gravimetrically, is listed below.

Analyte	CAS#	Analyte Lot	Concentration ± Uncertainty
Aroclor 1268	011100-14-4	RM00937	100.0 ± 0.5 µg/mL

Matrix: isooctane (2,2,4-trimethylpentane)

Storage Conditions: Store at Room Temperature (15° to 30°C).

K1262

Traceability:

The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCCL Z540.3, ISO 9001, ISO 17025, and ISO 17034. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 1088.

Homogeneity:

This RM was unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods, and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening the container and should be processed without delay for the certified values to be valid within the stated uncertainties.

Hazards:

Refer to the Safety Data Sheet on www.agilent.com for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.

Maintenance of Certification:

If substantive changes are noted that affect the certification before the expiration of this certificate, Agilent will notify the purchaser.

Sample lot approver:


Monica Bourgeois
QMS Representative



ISO 17034 Cert
No. AR-1936

RM was produced in accordance with TUV USA Inc registered ISO 9001 Quality Management System. Cert # 56 100 18560026

Page: 1 of 1

www.agilent.com/quality/
CSD-QA-015.1



ISO 17025 Cert
No. AT-1937

Recipient Copy

CHAIN-OF-CUSTODY RECORD

COC No. 15350

Order Number: CB014765

Date Shipped: 4/11/2022

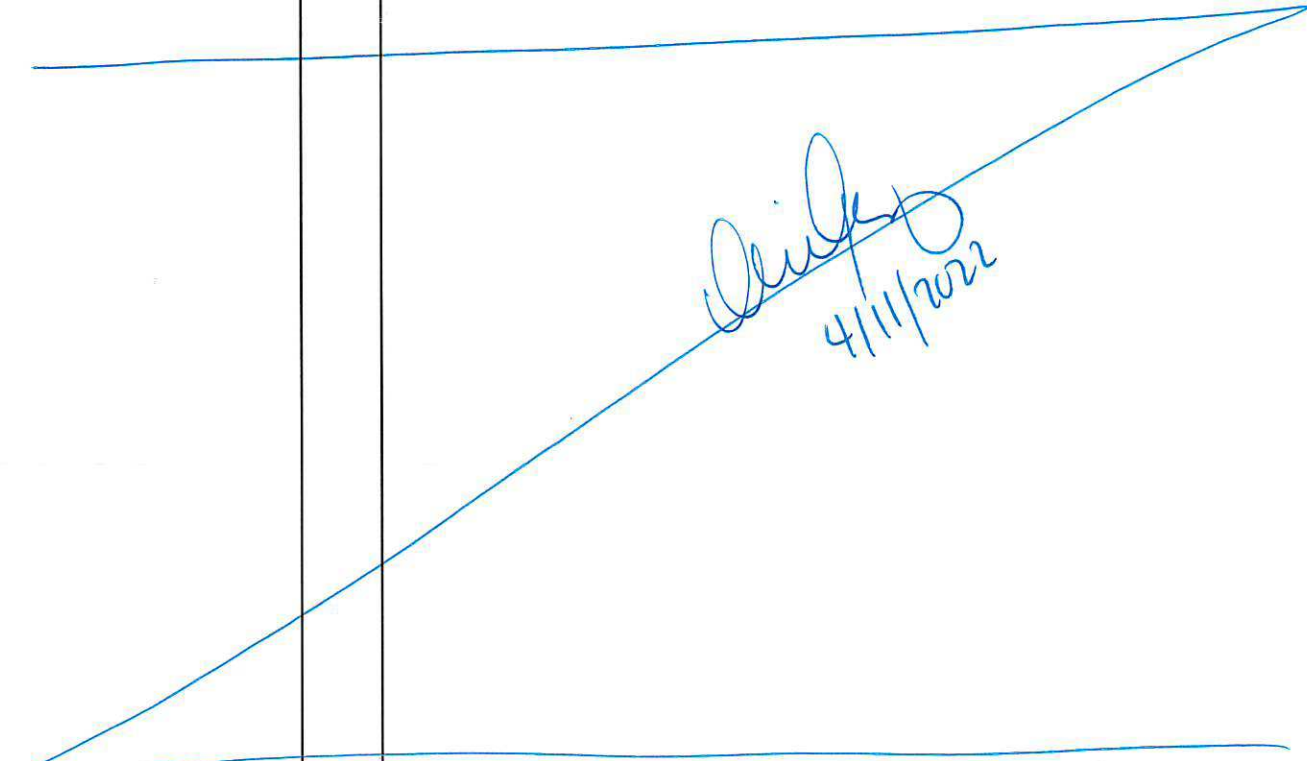
AirBill No(s):

From: QATS LABORATORY
 2700 CHANDLER AVENUE, BLDG. B
 LAS VEGAS, NV 89120
 PHONE: 1-702-895-8712

To: Kelly Bottem
 Analytical Resources, Inc.
 4611 S. 134th Place SUITE 100
 Tukwila WA 98168
 206-695-6211

519204140444

K003525 7
K003528

Sample ID	Qty	Description/Remarks	→ Catalogue Number
PSRM0148	1	PUGET SOUND SEDIMENT RM	PS-SRM
PSRM0149	1	PUGET SOUND SEDIMENT RM	PS-SRM
PSRM0150	1	PUGET SOUND SEDIMENT RM	PS-SRM
PSRM0151	1	PUGET SOUND SEDIMENT RM	PS-SRM
			
		BOEING PLANT 2	

Signature
 4/11/2022

Please use the enclosed Sample Preparation Instructions. If catalogue number(s) are listed at the top of the Sample Preparation Instructions use the Sample Preparation Instructions with catalogue number(s) matching the catalogue number(s) of each of the samples listed above.

Relinquished by: (Signature) <i>[Signature]</i>	Date/Time <i>1400</i> <i>4/11/2022</i>	Received by: (Signature) <i>[Signature]</i>	Date/Time <i>0955</i> <i>04/12/22</i>
Custody Seal(s): Present/Absent <i>PRESENT</i>	Remarks:		
Relinquished by: (Signature)	Date/Time	Received by: (Signature)	Date/Time



**PUGET SOUND SEDIMENT REFERENCE MATERIAL
QATS LABORATORY INSTRUCTIONS FOR
HRGC/HRMS CDD/CDF/CB CONGENER AND GC/ECD AROCLOR ANALYSIS**

NOTE: These instructions are for advisory purposes only. If any apparent conflict exists between these instructions and the analytical protocols or your contract, disregard these instructions.

APPLICATION: For the analysis of CDD/CDF and CB Congener analytes using project-specified HRGC/HRMS methods, and Aroclors using project-specified GC/ECD methods.

CAUTION: Read instructions carefully before opening bottles and proceeding with the analyses.

Contains CDD/CDF, CB Congener, and/or Aroclors
HAZARDOUS MATERIAL
Safety Data Sheets
Available Upon Request

(A) SAMPLE DESCRIPTION

Enclosed is a Puget Sound (Washington State) Sediment Reference Material (SRM) set for chlorinated dibenzo-p-dioxins/chlorinated dibenzofurans (CDD/CDF), and/or chlorinated biphenyl (CB) congener analysis using project-specified high resolution gas chromatography/ high resolution mass spectrometry (HRGC/HRMS) methods. This SRM is also suitable for Aroclors analysis using project-specified gas chromatography/electron capture detection (GC/ECD) methods. This set consists of one (1) or more bottles, each with approximately 30 grams of Puget Sound SRM containing CDD/CDF, CB Congener, and/or Aroclor analytes. Check the chain-of-custody record to determine the number of bottles provided for CDD/CDF, CB Congener, and/or Aroclor analysis. None of the bottles are to be opened until SRM preparation/analysis is to occur.

CAUTION: The SRM could contain compounds that are light sensitive and should be protected from light during storage. Store the SRM at $\leq 6^{\circ}\text{C}$, preferably at $< 0^{\circ}\text{C}$, until SRM preparation and analysis is to occur. Allow the bottle(s) to reach ambient temperature before opening.

(B) BREAKAGE OR MISSING ITEMS

Check the contents of the shipment carefully for any broken, leaking, or missing items. Refer to the enclosed chain-of-custody record. Report any problems to Mr. Keith Strout, APTIM Federal Services, LLC, at (702) 895-8722. If requested, return the chain-of-custody record with appropriate annotations and signatures to the address provided below.

QUALITY ASSURANCE TECHNICAL SUPPORT LABORATORY
APTIM Federal Services, LLC
2700 Chandler Avenue - Building C
Las Vegas, NV 89120



(C) ANALYSIS REQUIREMENTS

The SRM is to be analyzed as described in the project-specified methods employed for the analysis of CDD/CDF and/or CB Congener analytes using HRGC/HRMS instrumentation and/or Aroclors using GC/ECD instrumentation. These instructions are for advisory purposes only. If any apparent conflict exists between these instructions and the project-specified methods, or your contract, disregard these instructions.

(D) SAMPLE ANALYSIS

General Instructions

The SRM contains CDD/CDF, CB Congener, and Aroclor analytes which are known or suspected to have severe health effects. Employing appropriate safety precautions, this SRM is to be handled, prepared, and analyzed exactly as you would process samples received from a known or suspected hazardous waste site. The SRM should be handled only by trained and experienced analysts in facilities expressly designed to handle such materials. When calculating the concentrations of analytes, use 0% as the soil moisture content.

Allow the bottle(s) to reach ambient temperature before opening and removing gravimetric amounts for sample preparation. To begin the extraction and analysis procedure, break the seal and open the bottle carefully. Weigh out the appropriate aliquot for extraction and analysis as prescribed in the project-specified methods (typically 10 grams for HRGC/HRMS methods and 30 grams for GC/ECD methods), or in accordance with your contract.

Proceed immediately with the extraction and analysis as described in the project-specified methods or your contract.

(E) REPORTING

Report the results for the prepared SRM as received.

Report the analytical results for the SRM to EPA or other appropriate Agency, using the format and other instructions for submission of data packages as specified in your contract.

Certificate of Analysis



Phenova Certified Reference Materials are sold by Phenomenex.

411 Madrid Ave., Torrance, CA 90501 USA ■ Tel: 310-212-0555 ■ Fax: 310-328-7768 ■ info@phenomenex.com

Access your MSDS and digital C of A at www.phenomenex.com/mysupport. Re-order at www.phenomenex.com/standards

Certified Reference Material

This product is included in Phenova's ISO/IEC 17025 and ISO Guide 34 Scopes of Accreditation

Catalog No.: AL0-101467

Lot Number: CL12975

Description: Aroclor 1016

Certification Date: November 19, 2018

Storage: 4 °C

Expiration Date: October 31, 2026

Provided As: 1 mL in 2 mL Ampoule in Isooctane

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1016	12674-11-2	1000	± 0.553%

125829



Reference Material Producer
Certificate No. 2427.02



Manufactured by Phenova, Inc.

Phenova's testing and calibration results are internationally recognized through the ILAC-MRA. Phenova is an accredited ISO Guide 34 Reference Material Provider and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03

Certificate of Analysis



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Access your MSDS and digital C of A at www.phenomenex.com/mysupport. Re-order at www.phenomenex.com/standards

1. **Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
2. **Quality Standards:** Phenova is accredited by A2LA to ISO Guide 34³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
3. **Intended Use:** The product is manufactured for use in the calibration and calibration verification of chromatographic instrumentation performed in routine laboratory analysis.
4. **Instruction:** Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all certified analytes in the mixture.
5. **Hazardous Situation:** The product is intended for use by experienced professional personnel. A Material Safety Data Sheet (MSDS) is available at www.phenomenex.com/mysupport.
6. **Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
7. **Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
8. **Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
9. **Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).

$$u_{CRM} = k \sqrt{uM^2 + uH^2 + uLTS^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.

10. **Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO Guide 34. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
11. **Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO Guide 34.
12. **Period of Validity:** The Certified Values and their uncertainties are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

¹ ISO Guide 31:2000(E) – Reference Materials – Contents of Certificates and Labels.

² ISO Guide 35:2006(E) – Reference Material – General and Statistical Principles for Certification.

³ ISO Guide 34:2009(E) – General Requirements for the Competence of Reference Material Producers.

⁴ ISO/IEC 17025:2005(E) – General Requirements for the Competence of Testing and Calibration Laboratories.

⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



Reference Material Producer
Certificate No. 2427.02



Manufactured by Phenova, Inc.

Phenova's testing and calibration results are internationally recognized through the ILAC MRA. Phenova is an accredited ISO Guide 34 Reference Material Provider and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03

IL111063_US

Certificate of Analysis

Produced by Phenova

6390 Joyce Drive STE 100, Golden, CO 80403 USA ■ Tel: 303-940-0033 ■ Fax: 303-940-0043 ■ info@phenova.com
Access your Safety Data Sheets and digital Certificates at www.phenova.com/documents.

Certified Reference Material

This product is certified in accordance with Phenova's ISO 17034 accreditation and supported by Phenova's ISO/IEC 17025 chemical testing accreditation

Catalog No.: AL0-101462

Lot Number: CL18021

Description: Aroclor 1260

Certification Date: February 14, 2022

Storage: 4 °C

Expiration Date: February 28, 2030

Provided As: 1 mL in 2 mL Ampoule in Hexane

Andrea L Gill

Andrea Gill, Certified Reference Materials Manager

Component	CAS #	Certified Value µg/mL	Expanded Uncertainty
Aroclor 1260	11096-82-5	1000	± 0.553%

K005830



Reference Material Producer
Certificate No. 2427.02



Phenova is an accredited ISO/IEC 17034 Reference Material Producer and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03

Certificate of Analysis



Page 2 of 2

Produced by Phenova

6390 Joyce Drive STE 100, Golden, CO 80403 USA ■ Tel: 303-940-0033 ■ Fax: 303-940-0043 ■ info@phenova.com
Access your Safety Data Sheets and digital Certificates at www.phenova.com/documents.

- Quality Document:** This Certificate of Analysis has been created in accordance with ISO Guide 31¹ and ISO Guide 35.²
- Quality Standards:** Phenova is accredited by A2LA to ISO 17034³ and ISO/IEC 17025⁴ as a producer of Certified Reference Materials and Reference Materials. This ensures that our manufacturing processes have been accredited to and meet strict international standards.
- Intended Use:** The product is manufactured for use in calibration, calibration verification, quantification, identification and other appropriate analytical control applications. The product is intended for routine laboratory analysis and research purposes only. Only trained personnel should handle this product.
- Handling and Usage Notes:** Store according to recommended conditions listed and avoid prolonged exposure to light. Visually inspect the solution inside the ampoule for any un-dissolved material. If particulate is visible, sonicate the unopened ampoule until material is fully dissolved. Dilute as required, use only class A glassware and diluents compatible with all analytes in the mixture. Considerations should be made related to repeated use of the opened product. Once opened, exposure to light, air, heat, objects, and additional transfer vessels may cause evaporation, degradation or contamination resulting in changes in concentration, uncertainty and stability duration. Store opened standards in a clean, tightly capped vessel under the recommended temperature. Appropriate controls, such as the use of additional verification standards should be used to confirm the opened product is fit for purpose under repeated use conditions.
- Hazardous Situation:** The product is intended for use by experienced professional personnel. A Safety Data Sheet (SDS) is available at www.phenova.com/documents.
- Level of Homogeneity:** The product has been certified to guarantee the certified values and their uncertainties at a volume of 2 µL.
- Certified Value:** Certified Value is based upon gravimetric and volumetric preparation using calibrated balances and Class A glassware.
- Raw Materials and Purity:** Phenova reference standard products are prepared from the highest quality starting materials. The purity of this material was verified using an ISO/IEC 17025 methodology.
- Expanded Uncertainty:** The expanded uncertainty (uCRM) as stated is determined in accordance with ISO/IEC Guide 98⁵ and ISO Guide 35 incorporating Type A standard uncertainty at a 95% confidence level. The uncertainty contains elements of manufacturing (uM), homogeneity analysis (uH) and long-term stability testing (uLTS). The uncertainty is calculated based on the root-sum-of-squares equation times a coverage factor (k=2).
$$u_{CRM} = k\sqrt{u_M^2 + u_H^2 + u_{LTS}^2}$$

Transport conditions (short-term stability) have been tested such that there is no contribution to the uncertainty reported. The expanded uncertainty applies to the product as received.
- Metrological Traceability:** The property value (certified value and its uncertainty) are traceable through an unbroken chain of calibration to the SI base unit kg through a NIST traceable weight in accordance with ISO 17034. This is achieved through calibration of balances, verification of weights, use of national methodology for glassware calibration and product homogeneity and stability testing utilizing an ISO/IEC 17025 methodology.
- Values Obtained During Product Testing:** This product is subjected to verification, homogeneity and stability testing using an ISO/IEC 17025 chromatographic methodology. All values obtained during testing meet criteria in accordance with ISO 17034.
- Period of Validity:** The Certified Values, Uncertainties and Expiration Date are based on the unopened product being stored according to the recommended storage condition listed and are guaranteed until the expiration date. This product will be monitored during the period of validity and customers notified of any significant changes in stability.

References:

- ¹ ISO Guide 31 – Reference Materials – Contents of Certificates and Labels.
- ² ISO Guide 35 – Reference Material – General and Statistical Principles for Certification.
- ³ ISO 17034 – General Requirements for the Competence of Reference Material Producers.
- ⁴ ISO/IEC 17025 – General Requirements for the Competence of Testing and Calibration Laboratories.
- ⁵ ISO/IEC Guide 98-3:2008(E) – Uncertainty of Measurement – Part 3: Guide to Expression of Uncertainty in Measurement (GUM: 1995)



Reference Material Producer
Certificate No. 2427.02



Phenova is an accredited ISO/IEC 17034 Reference Material
Producer and ISO/IEC 17025 accredited Chemical Testing Laboratory.



Chemical Testing Laboratory
Certificate No. 2427.03



Form I
INORGANIC ANALYSIS DATA SHEET
EPA 9060A m

LDW22-SS826

Laboratory: Analytical Resources, LLC
 Client: Anchor QEA, LLC
 Project: AOC4 UR Phase 3
 Matrix: Sediment Laboratory ID: 23C0107-01 C SDG: 23C0107
 Sampled: 03/02/23 16:25 Prepared: 04/05/23 15:19 File ID: CubeData_04072023@1412-030
 % Solids: 43.35 Preparation: PSEP 1986 (modified) Analyzed: 04/07/23 01:37
 Batch: BLD0117 Sequence: SLD0078 Initial/Final: 0.6154 g Wet / 0.6154 mL
 Instrument: TOC Cube Calibration: FD00070

CAS NO.	Analyte	Concentration (% dry)	Dilution Factor	MDL	MRL	Q
	Total Organic Carbon	2.37	1	0.02	0.02	



Form I
INORGANIC ANALYSIS DATA SHEET
EPA 9060A m

LDW21-IT608C

Laboratory: Analytical Resources, LLC
 Client: Anchor QEA, LLC
 Project: AOC4 UR Phase 3
 Matrix: Sediment Laboratory ID: 23C0107-02 B SDG: 23C0107
 Sampled: 07/13/21 06:54 Prepared: 04/05/23 15:19 File ID: CubeData_04072023@1412-031
 % Solids: 67.49 Preparation: PSEP 1986 (modified) Analyzed: 04/07/23 02:07
 Batch: BLD0117 Sequence: SLD0078 Initial/Final: 0.5175 g Wet / 0.5175 mL
 Instrument: TOC Cube Calibration: FD00070

CAS NO.	Analyte	Concentration (% dry)	Dilution Factor	MDL	MRL	Q
	Total Organic Carbon	0.86	1	0.02	0.02	H



Form I
INORGANIC ANALYSIS DATA SHEET
EPA 9060A m

LDW21-IT608D

Laboratory: Analytical Resources, LLC
 Client: Anchor QEA, LLC
 Project: AOC4 UR Phase 3
 Matrix: Sediment Laboratory ID: 23C0107-03 B SDG: 23C0107
 Sampled: 07/13/21 06:54 Prepared: 04/05/23 15:19 File ID: CubeData_04072023@1412-032
 % Solids: 73.57 Preparation: PSEP 1986 (modified) Analyzed: 04/07/23 02:38
 Batch: BLD0117 Sequence: SLD0078 Initial/Final: 0.5157 g Wet / 0.5157 mL
 Instrument: TOC Cube Calibration: FD00070

CAS NO.	Analyte	Concentration (% dry)	Dilution Factor	MDL	MRL	Q
	Total Organic Carbon	0.37	1	0.02	0.02	H



PREPARATION BATCH SUMMARY

EPA 9060A m

Laboratory: Analytical Resources, LLC SDG: 23C0107
Client: Anchor QEA, LLC Project: AOC4 UR Phase 3
Batch: BLD0117 Batch Matrix: Solid Preparation: PSEP 1986 (modified)

SAMPLE NAME	LAB SAMPLE ID	LAB FILE ID	DATE PREPARED	OBSERVATIONS
LDW22-SS826	23C0107-01	eData_04072023@1412	04/05/23 15:19	
LDW21-IT608C	23C0107-02	eData_04072023@1412	04/05/23 15:19	
LDW21-IT608D	23C0107-03	eData_04072023@1412	04/05/23 15:19	
Blank	BLD0117-BLK1	eData_04072023@1412	04/05/23 15:19	
LCS	BLD0117-BS1	eData_04072023@1412	04/05/23 15:19	
MRL Check	BLD0117-MRL1	eData_04072023@1412	04/05/23 15:19	
Reference	BLD0117-SRM1	eData_04072023@1412	04/05/23 15:19	



Form I
METHOD BLANK DATA SHEET
EPA 9060A m
TotalAnalytes

Blank

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Batch: BLD0117

Laboratory ID: BLD0117-BLK1

Prepared: 04/05/23 15:19

Matrix: Solid

Preparation: PSEP 1986 (modified)

Analyzed: 04/06/23 13:27

Sequence: SLD0078

Calibration: FD00070

Instrument: TOC Cube

CAS NO.	Analyte	Concentration (% wet)	Dilution Factor	MDL	MRL	Q
	Total Organic Carbon	ND	1	0.02	0.02	U



LCS / LCS DUPLICATE RECOVERY
EPA 9060A m

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Matrix:	<u>Solid</u>	Analyzed:	<u>04/06/23 13:58</u>
Batch:	<u>BLD0117</u>	Laboratory ID:	<u>BLD0117-BS1</u>
Preparation:	<u>PSEP 1986 (modified)</u>	Sequence Name:	<u>LCS</u>
Initial/Final:	<u>0.0255 g / 0.0255 mL</u>		

COMPOUND	SPIKE ADDED (% wet)	LCS CONCENTRATION (% wet)	Q	LCS % REC. #	QC LIMITS REC.
Total Organic Carbon	44.4	45.3		102	80 - 120

* Indicates values outside of QC limits



ANALYSIS BATCH (SEQUENCE) SUMMARY

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Sequence: SKD0371

Instrument: TOC Cube

Calibration: FD00070

Sample Name	Lab Sample ID	Lab File ID	Matrix	Analysis Date/Time
Cal Standard	SKD0371-CAL1	CubeData_04272022@1136-001	NA	04/26/22 12:30
Cal Standard	SKD0371-CAL2	CubeData_04272022@1136-002	NA	04/26/22 13:00
Cal Standard	SKD0371-CAL3	CubeData_04272022@1136-003	NA	04/26/22 13:30
Cal Standard	SKD0371-CAL4	CubeData_04272022@1136-004	NA	04/26/22 14:00
Cal Standard	SKD0371-CAL5	CubeData_04272022@1136-005	NA	04/26/22 14:30
Cal Standard	SKD0371-CAL6	CubeData_04272022@1136-006	NA	04/26/22 15:00
Cal Standard	SKD0371-CAL7	CubeData_04272022@1136-007	NA	04/26/22 15:30
Cal Standard	SKD0371-CAL8	CubeData_04272022@1136-008	NA	04/26/22 16:00
Cal Standard	SKD0371-CAL9	CubeData_04272022@1136-009	NA	04/26/22 16:30
Cal Standard	SKD0371-CALA	CubeData_04272022@1136-010	NA	04/26/22 17:00
Cal Standard	SKD0371-CALB	CubeData_04272022@1136-011	NA	04/26/22 17:30
Cal Standard	SKD0371-CALC	CubeData_04272022@1136-012	NA	04/26/22 18:00
Cal Standard	SKD0371-CALD	CubeData_04272022@1136-013	NA	04/26/22 18:30
Cal Standard	SKD0371-CALE	CubeData_04272022@1136-014	NA	04/26/22 19:00
Cal Standard	SKD0371-CALF	CubeData_04272022@1136-015	NA	04/26/22 19:31
Cal Standard	SKD0371-CALG	CubeData_04272022@1136-016	NA	04/26/22 20:01
Cal Standard	SKD0371-CALH	CubeData_04272022@1136-017	NA	04/26/22 20:31
Cal Standard	SKD0371-CALI	CubeData_04272022@1136-018	NA	04/26/22 21:01
Cal Standard	SKD0371-CALJ	CubeData_04272022@1136-019	NA	04/26/22 21:31
Cal Standard	SKD0371-CALK	CubeData_04272022@1136-020	NA	04/26/22 22:01
Initial Cal Check	SKD0371-ICV1	CubeData_04272022@1136-027	NA	04/27/22 02:03
Initial Cal Blank	SKD0371-ICB1	CubeData_04272022@1136-028	NA	04/27/22 02:33
Cal Standard	SKD0371-CALL	CubeData_04272022@1136-021	NA	04/27/22 11:08
Cal Standard	SKD0371-CALM	CubeData_04272022@1136-022	NA	04/27/22 11:08
Cal Standard	SKD0371-CALN	CubeData_04272022@1136-023	NA	04/27/22 11:09
Cal Standard	SKD0371-CALO	CubeData_04272022@1136-024	NA	04/27/22 11:09



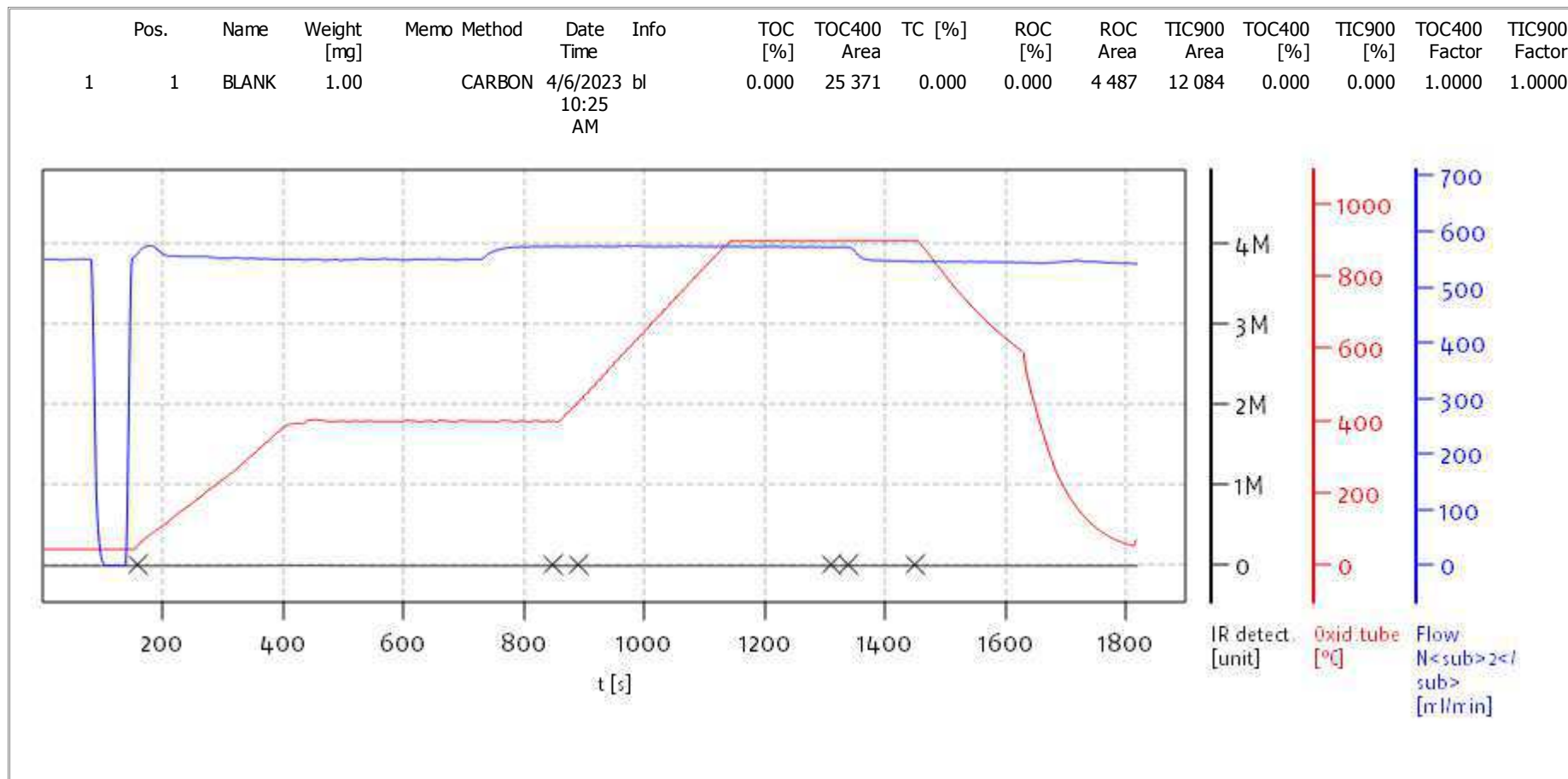
ANALYSIS BATCH (SEQUENCE) SUMMARY

EPA 9060A m

Laboratory:	<u>Analytical Resources, LLC</u>	SDG:	<u>23C0107</u>
Client:	<u>Anchor QEA, LLC</u>	Project:	<u>AOC4 UR Phase 3</u>
Sequence:	<u>SLD0078</u>	Instrument:	<u>TOC Cube</u>
		Calibration:	<u>FD00070</u>

Sample Name	Lab Sample ID	Lab File ID	Matrix	Analysis Date/Time
Initial Cal Check	SLD0078-ICV1	CubeData_04072023@1412-003	NA	04/06/23 11:57
Initial Cal Blank	SLD0078-ICB1	CubeData_04072023@1412-004	NA	04/06/23 12:27
MRL Check	BLD0117-MRL1	CubeData_04072023@1412-005	Solid	04/06/23 12:57
Blank	BLD0117-BLK1	CubeData_04072023@1412-006	Solid	04/06/23 13:27
LCS	BLD0117-BS1	CubeData_04072023@1412-007	Solid	04/06/23 13:58
Reference	BLD0117-SRM1	CubeData_04072023@1412-008	Solid	04/06/23 14:28
Calibration Check	SLD0078-CCV1	CubeData_04072023@1412-015	NA	04/06/23 18:00
Calibration Blank	SLD0078-CCB1	CubeData_04072023@1412-016	NA	04/06/23 18:30
Calibration Check	SLD0078-CCV2	CubeData_04072023@1412-027	NA	04/07/23 00:06
Calibration Blank	SLD0078-CCB2	CubeData_04072023@1412-028	NA	04/07/23 00:36
LDW22-SS826	23C0107-01	CubeData_04072023@1412-030	Solid	04/07/23 01:37
LDW21-IT608C	23C0107-02	CubeData_04072023@1412-031	Solid	04/07/23 02:07
LDW21-IT608D	23C0107-03	CubeData_04072023@1412-032	Solid	04/07/23 02:38
Calibration Check	SLD0078-CCV3	CubeData_04072023@1412-039	NA	04/07/23 06:11
Calibration Blank	SLD0078-CCB3	CubeData_04072023@1412-040	NA	04/07/23 06:42
Calibration Check	SLD0078-CCV4	CubeData_04072023@1412-050	NA	04/07/23 12:17
Calibration Blank	SLD0078-CCB4	CubeData_04072023@1412-051	NA	04/07/23 12:47

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

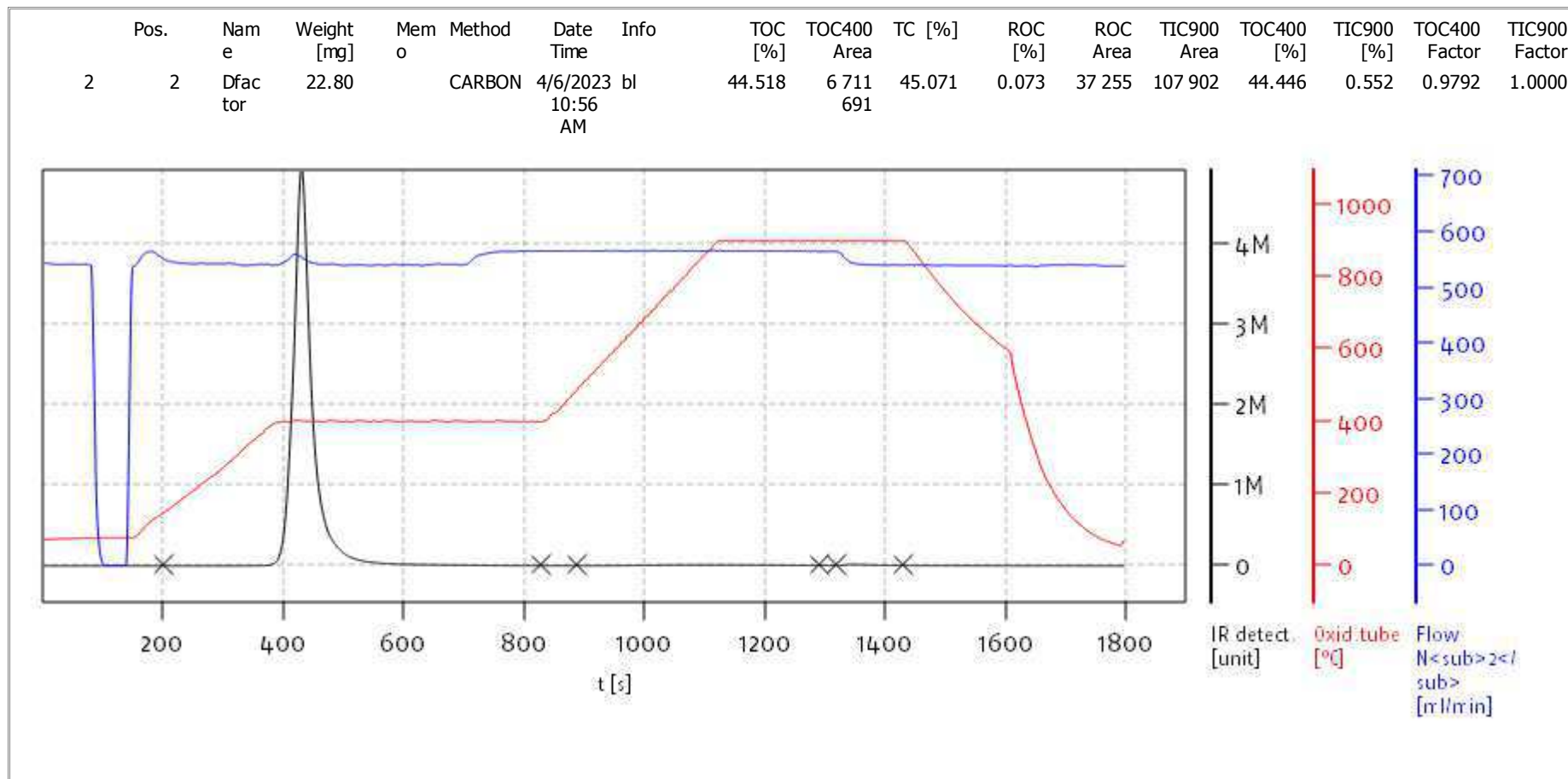
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Date: Fri Apr 7 14:09:03 2023



soliTOC V2.0.2 (31015f9) 2018-11-19
 Serial No: 0300.181017
 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

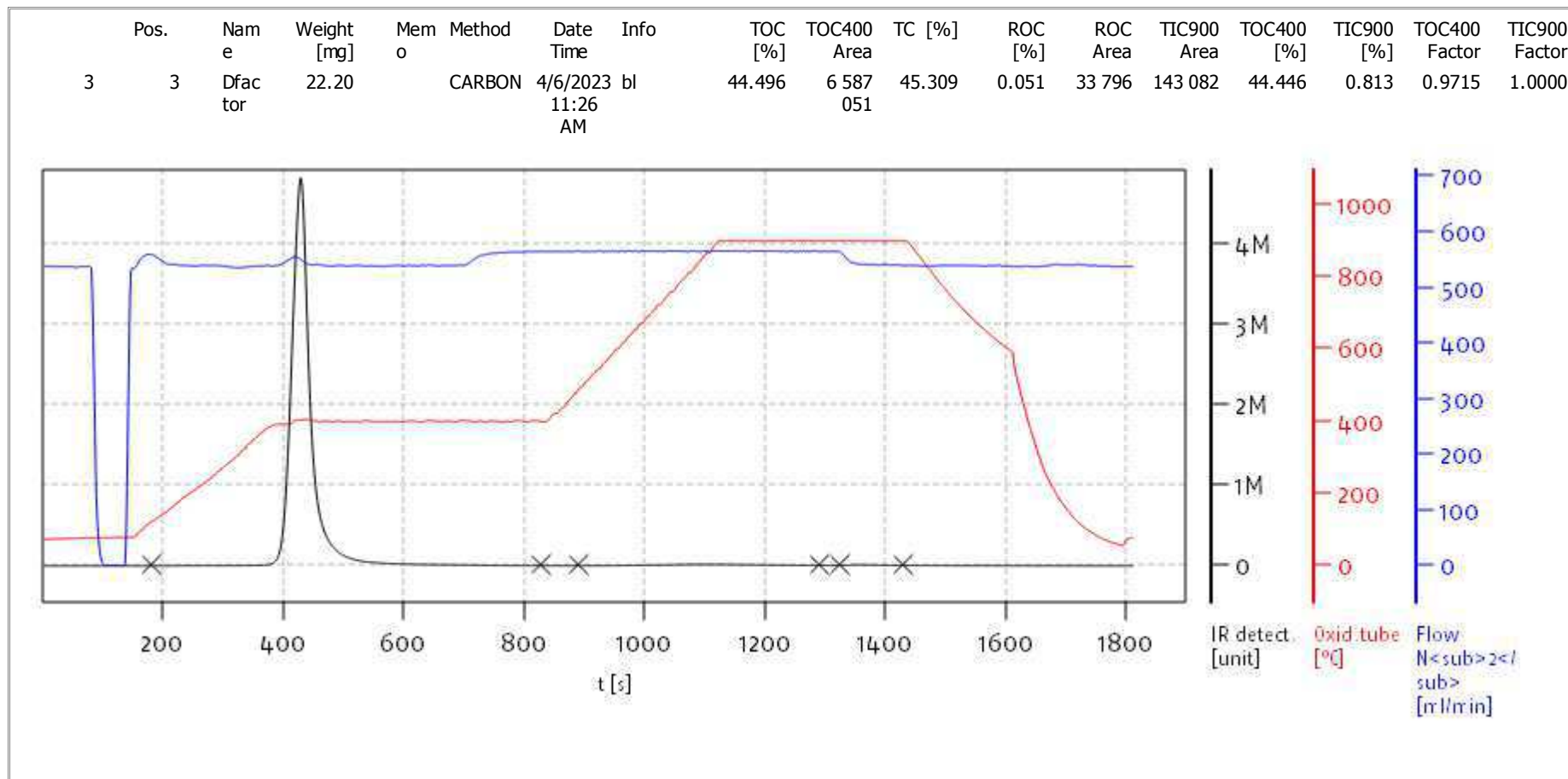
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 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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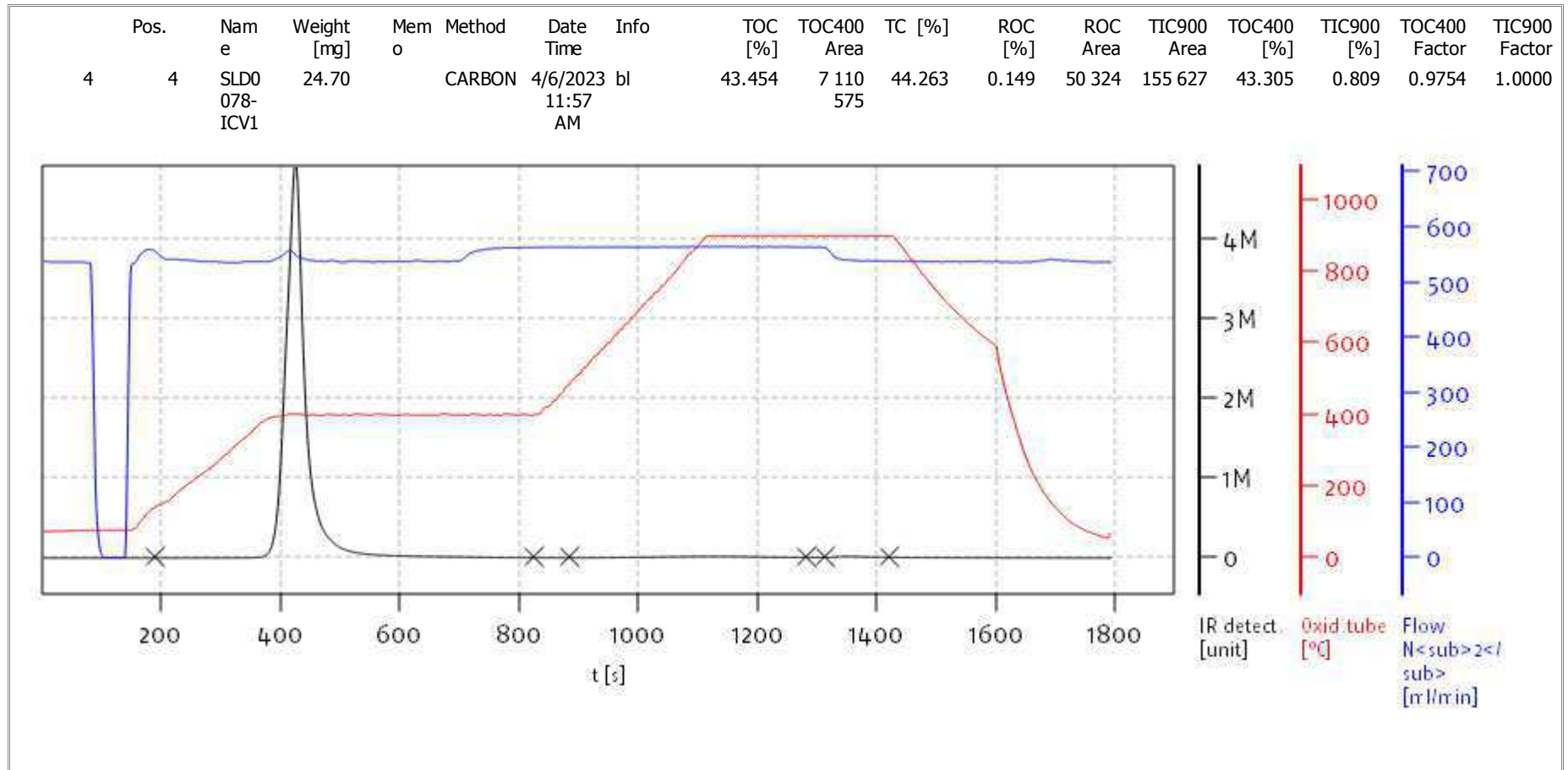
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 Mode CCC

Soli TOC Cube, Carbon
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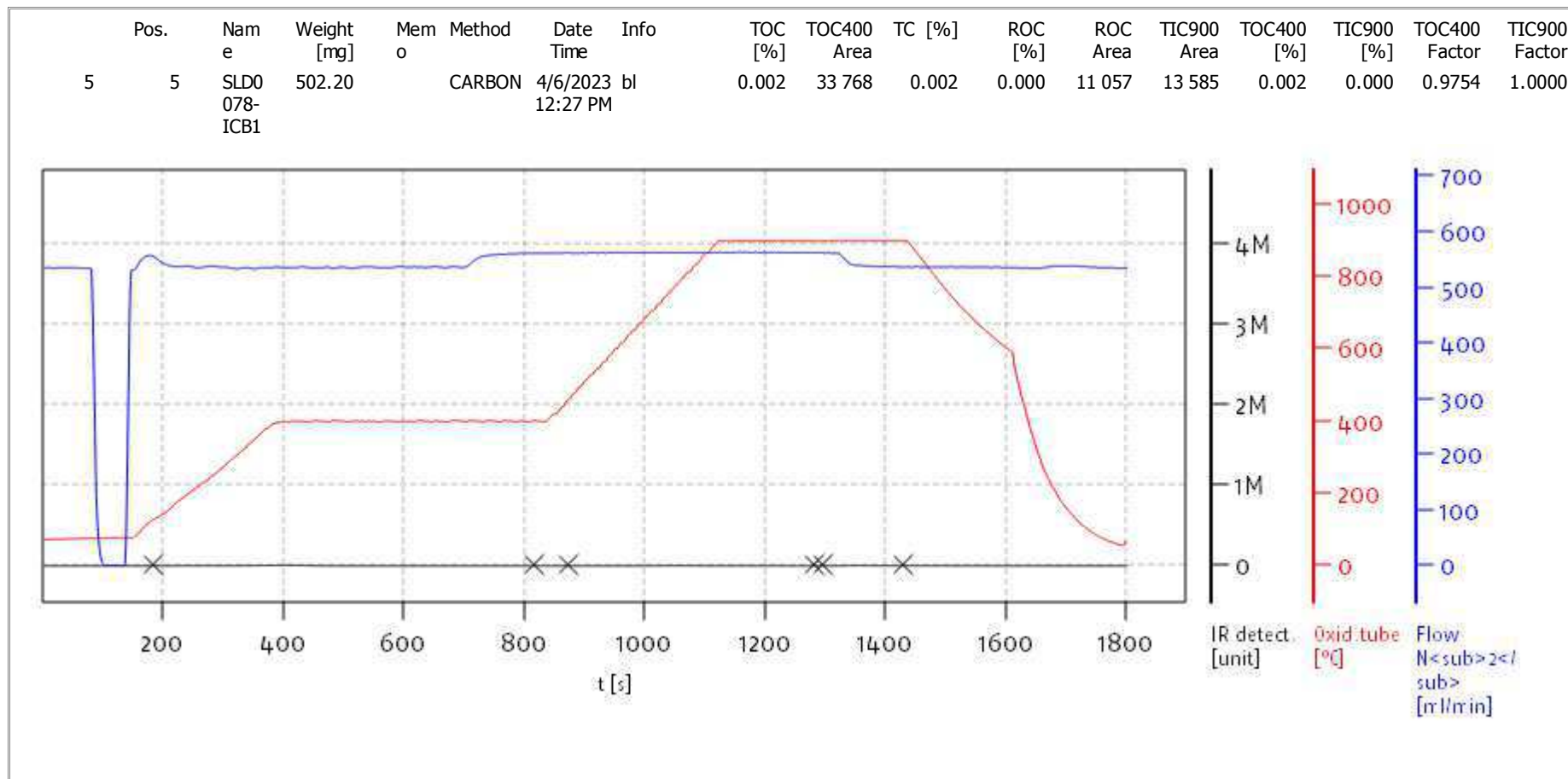
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Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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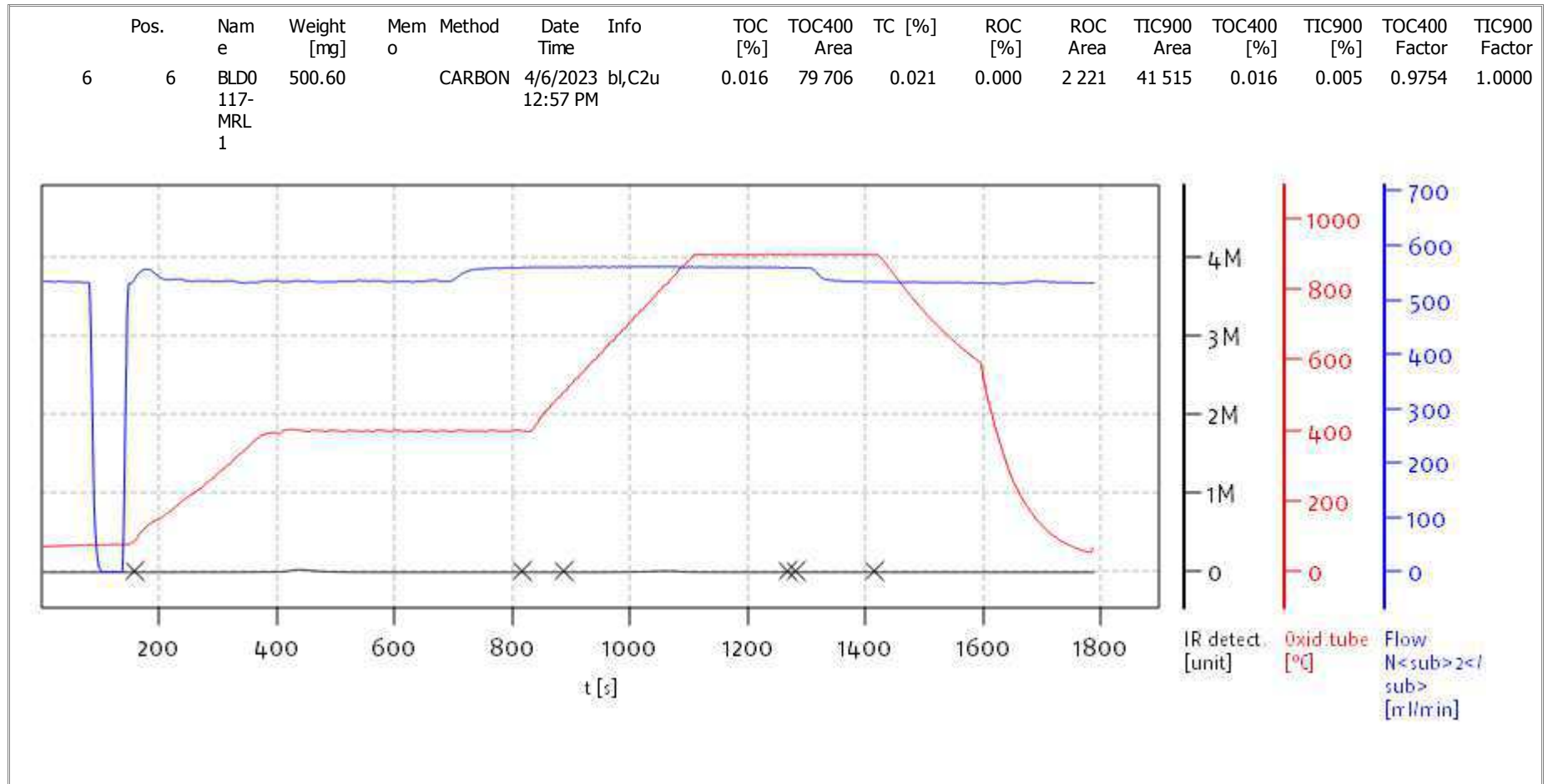
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 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

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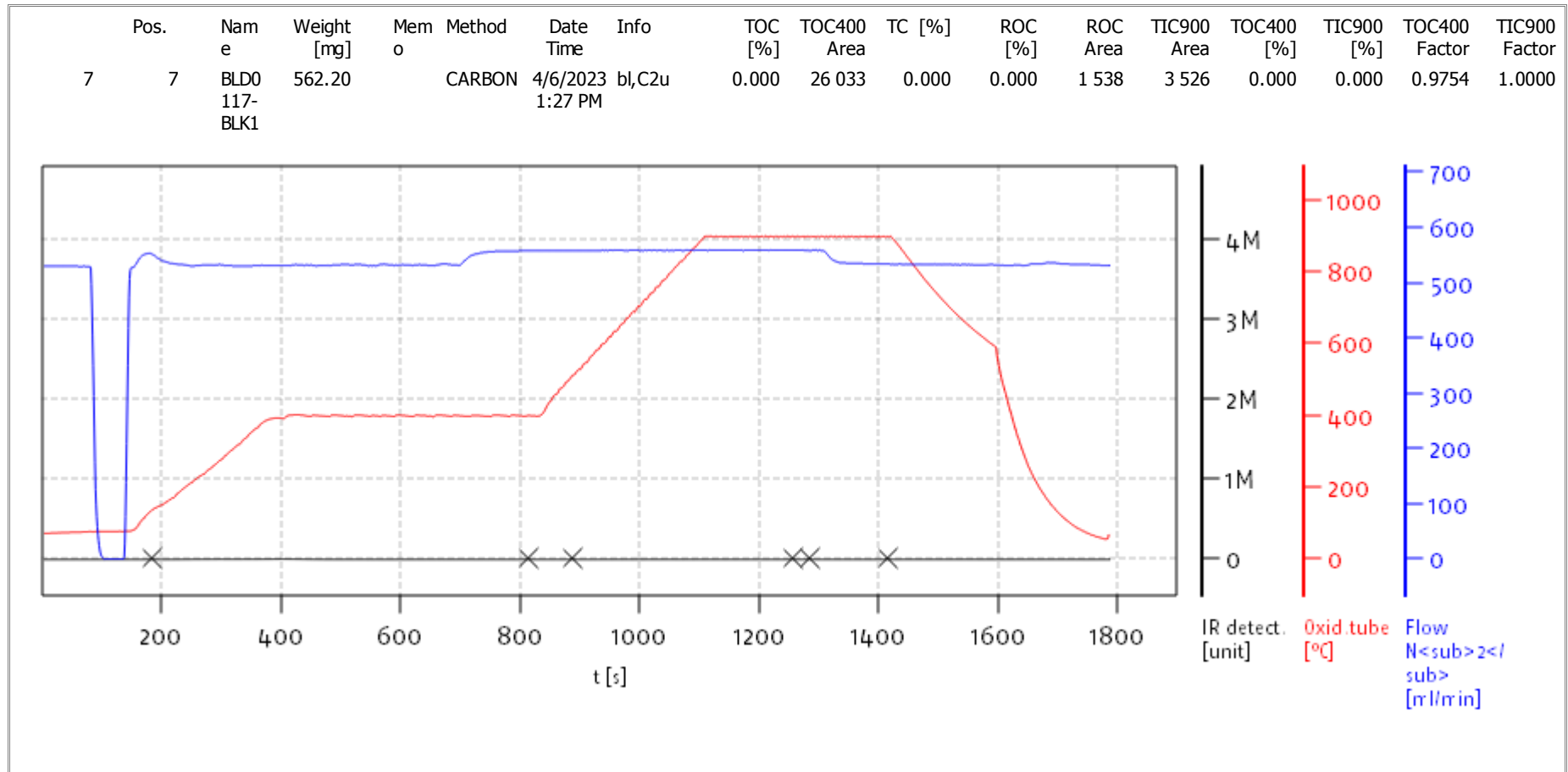
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 Serial No: 0300.18107
 Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: CDE



Name:

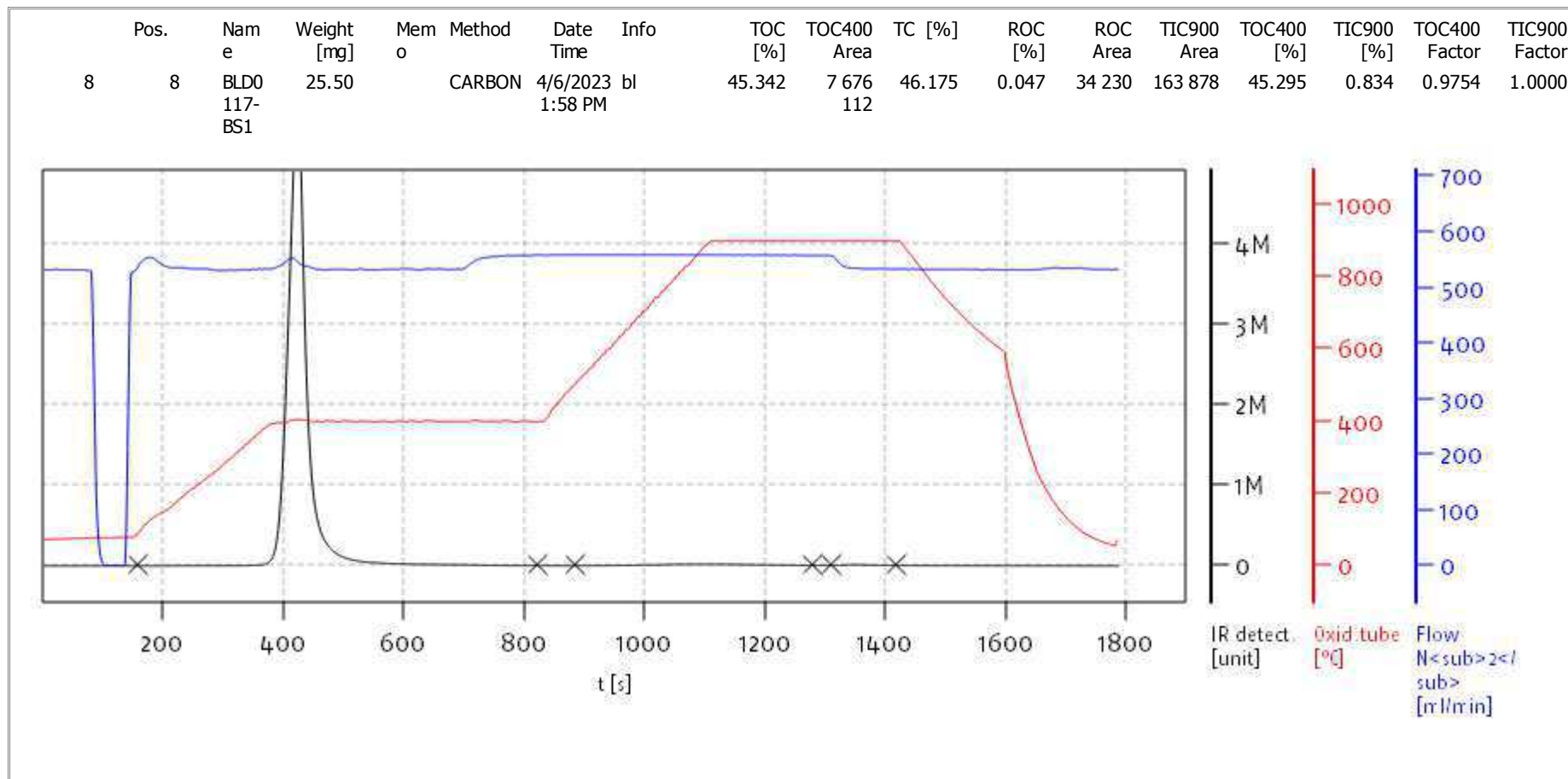
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Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

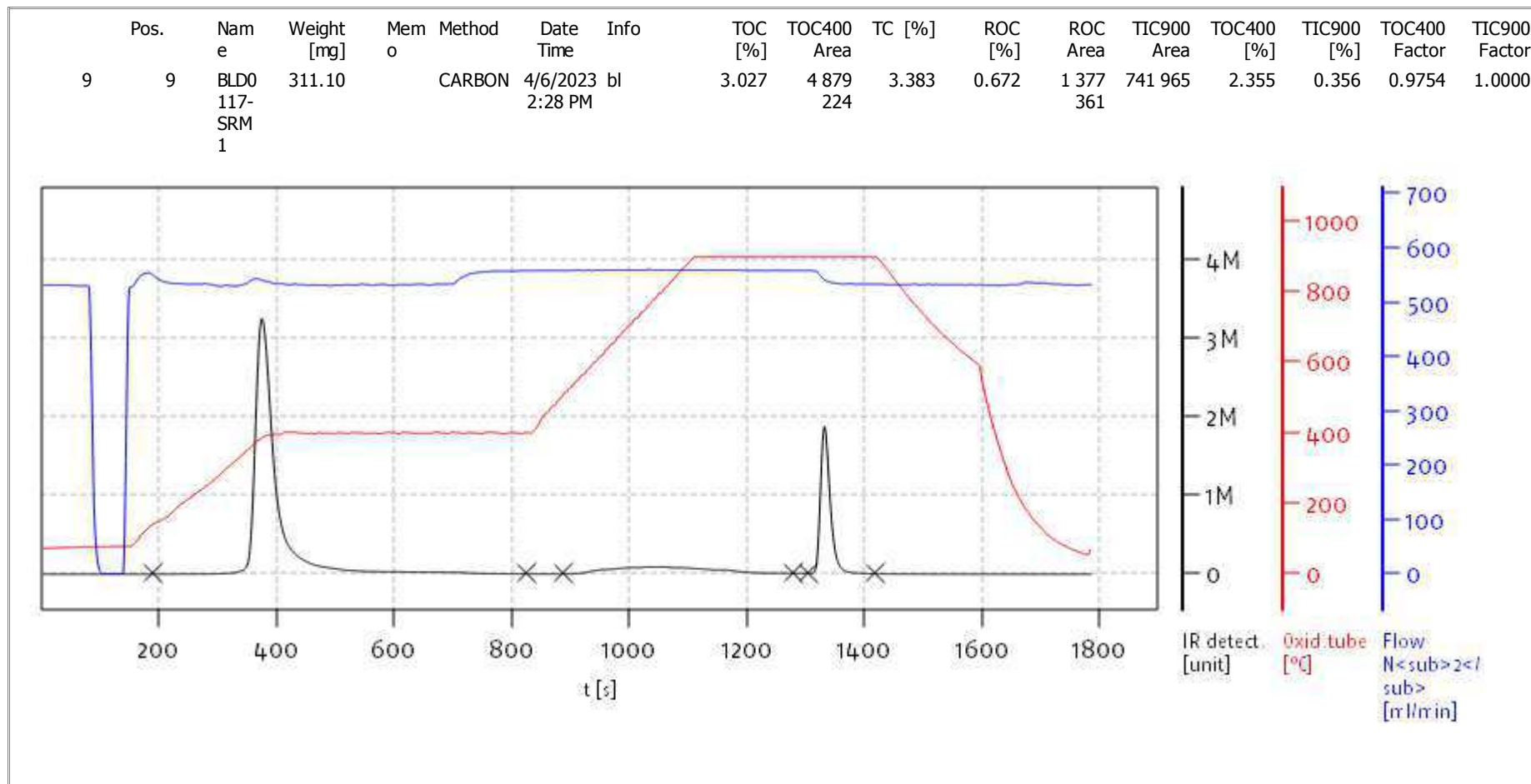
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Soli TOC Cube, Carbon
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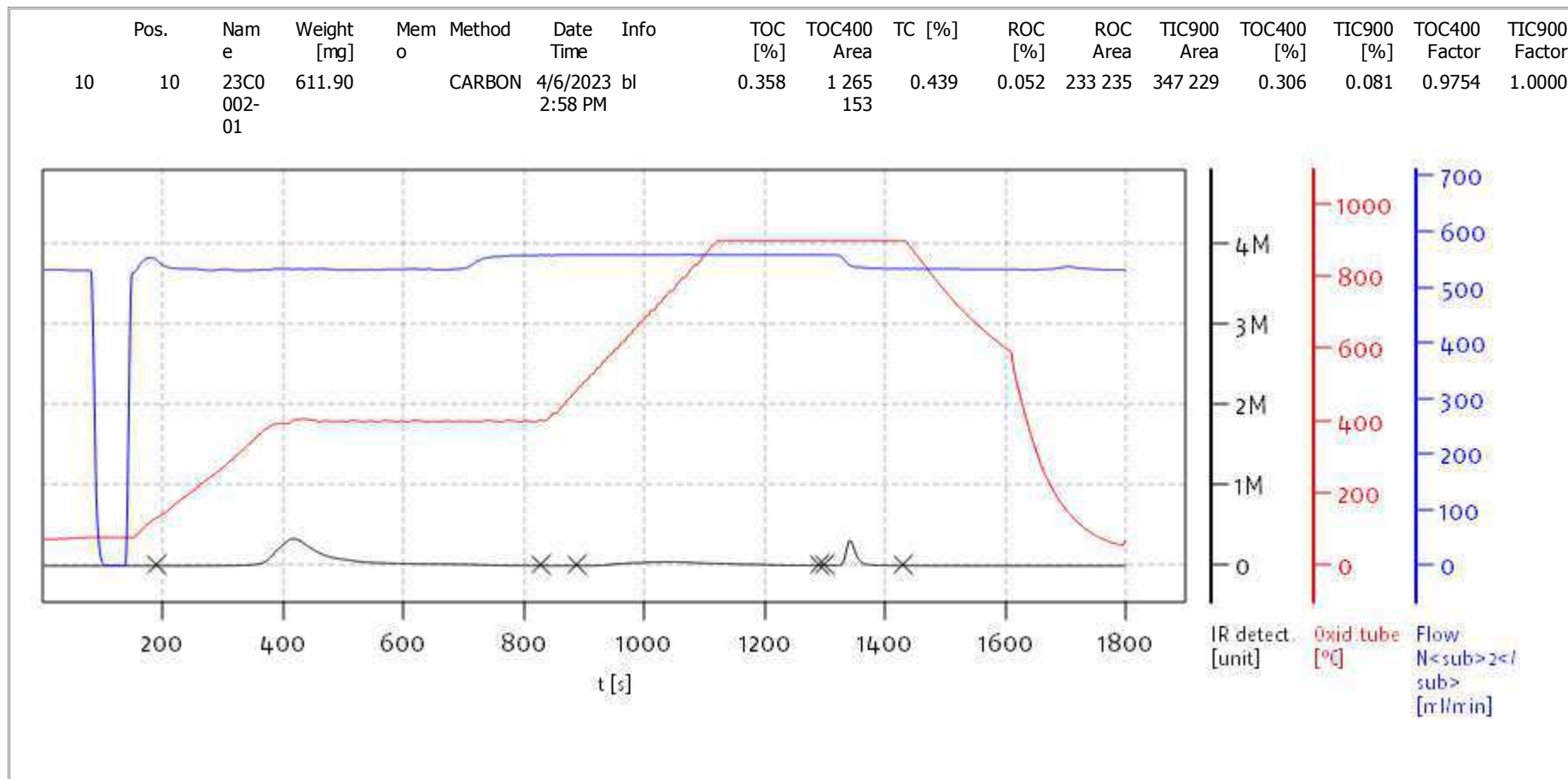
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Soli TOC Cube, Carbon
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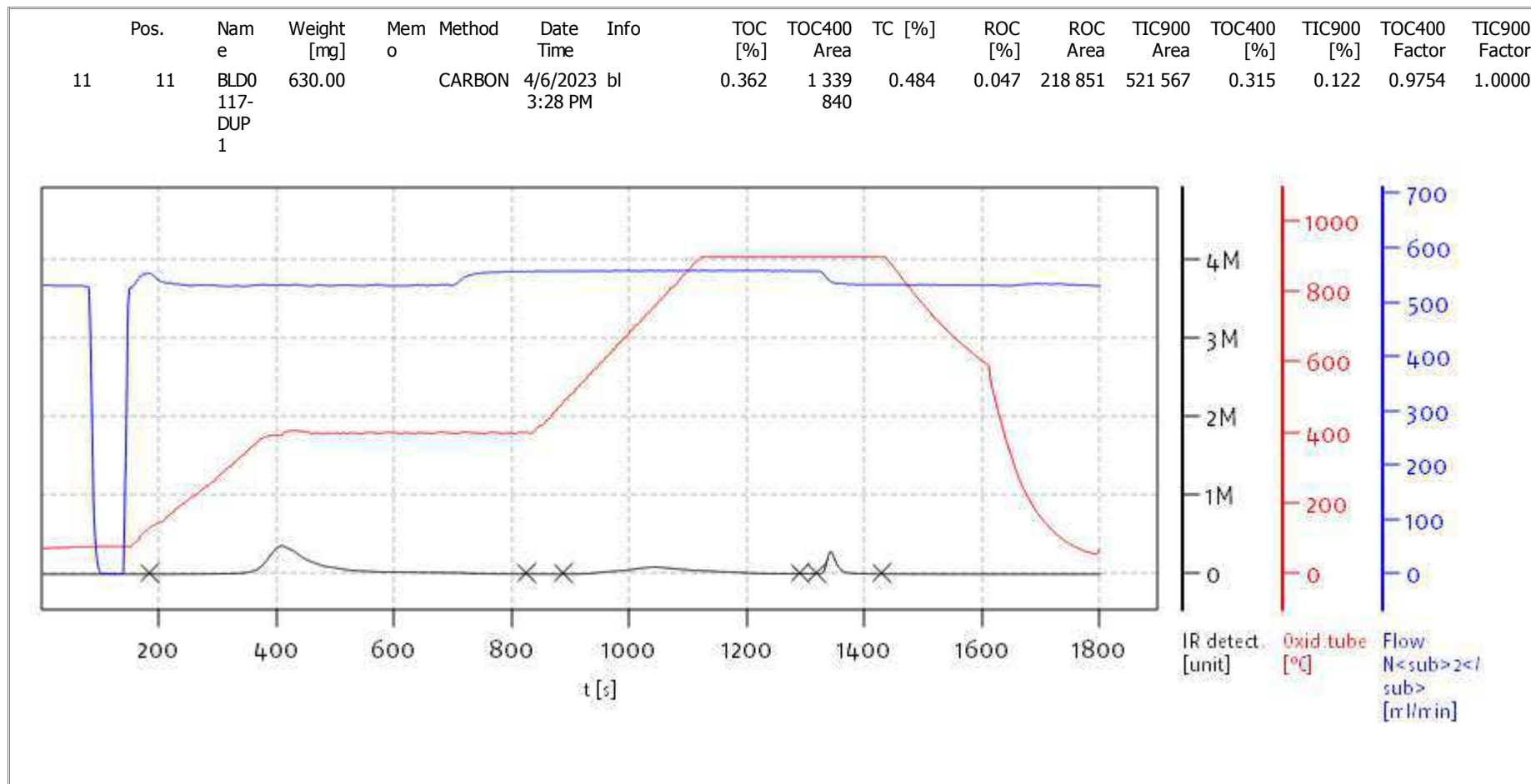
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Soli TOC Cube, Carbon
 Balance: BAL3
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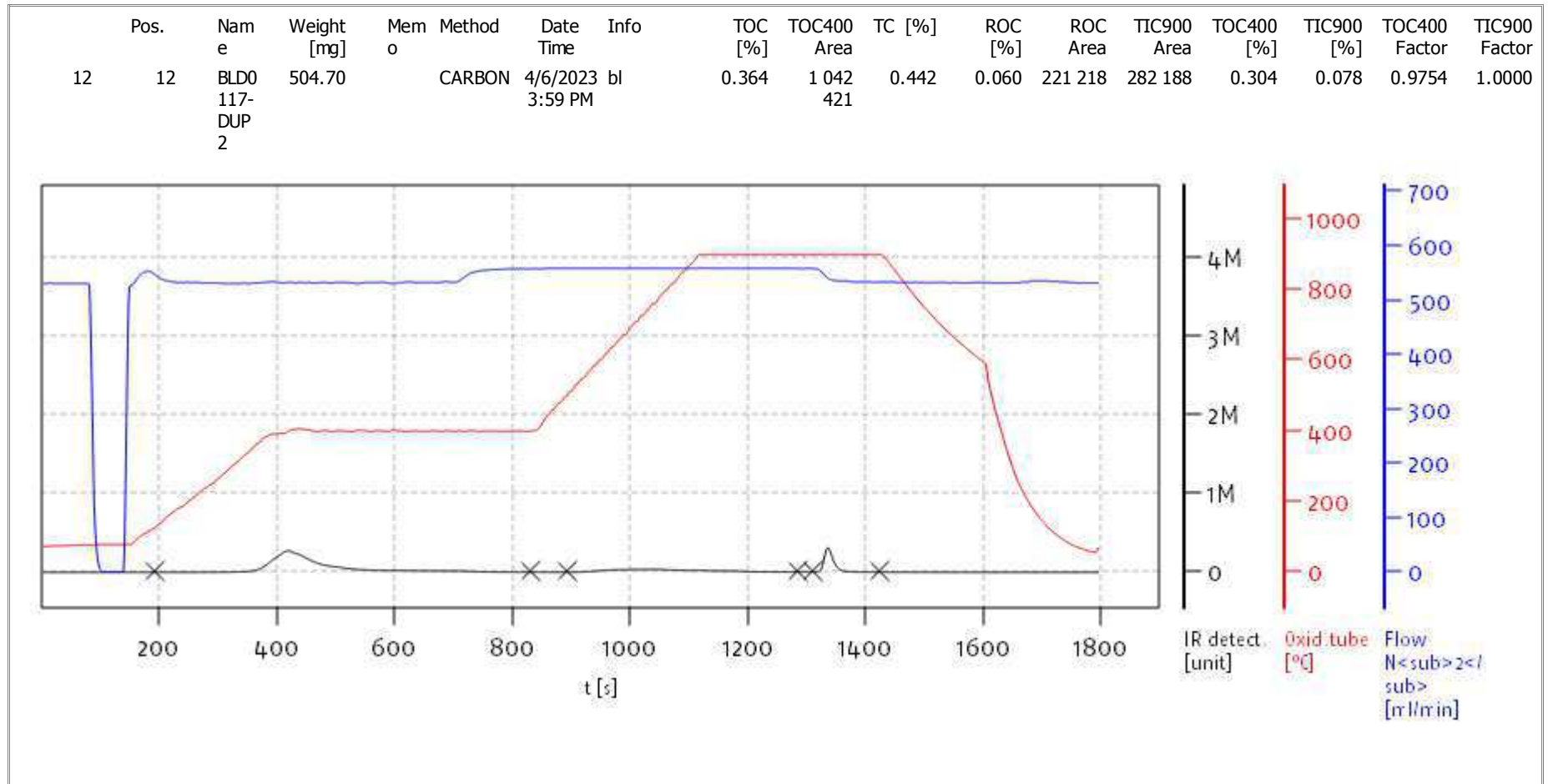
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Soli TOC Cube, Carbon
Balance: BAL3
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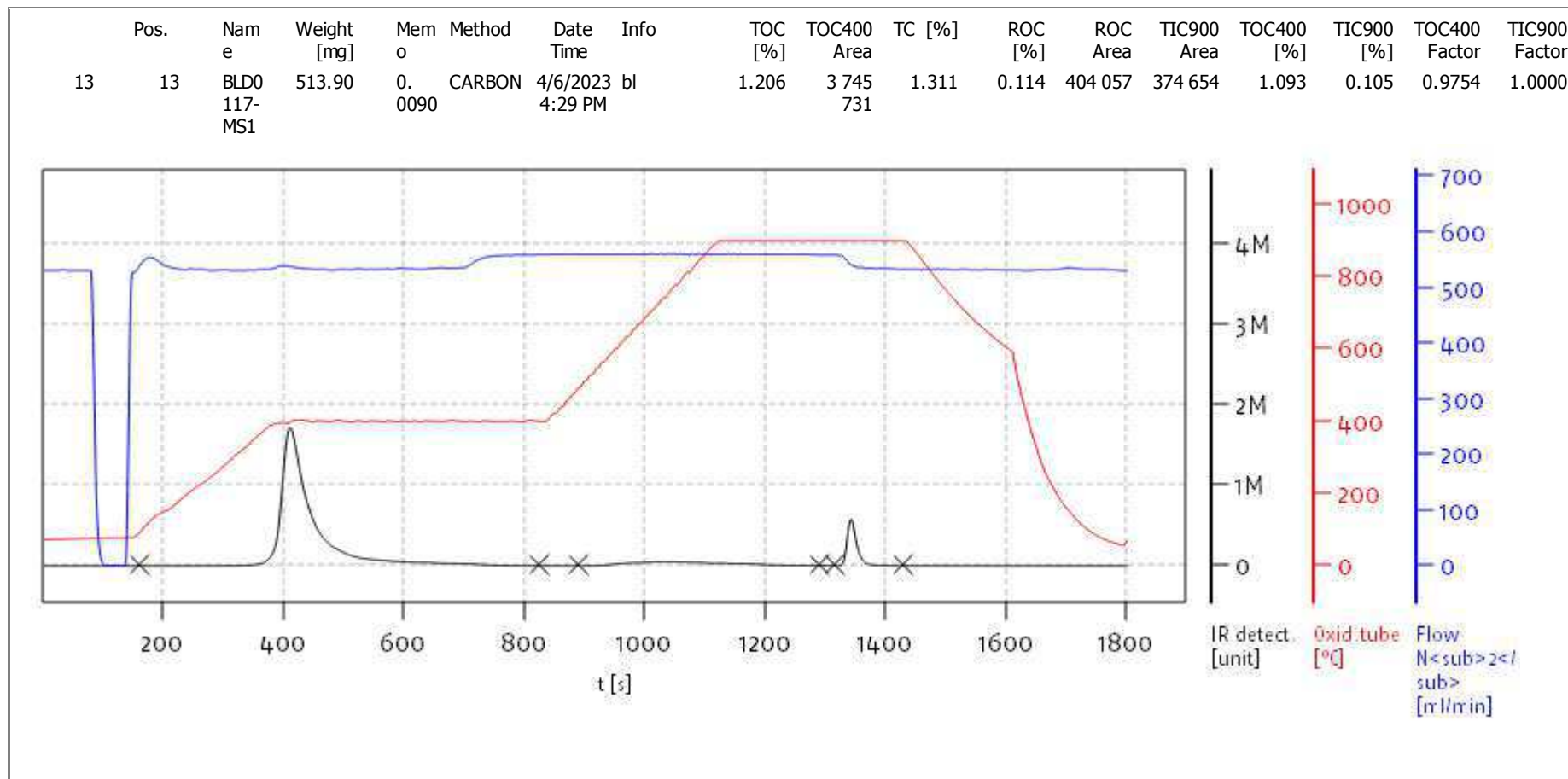
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Soli TOC Cube, Carbon
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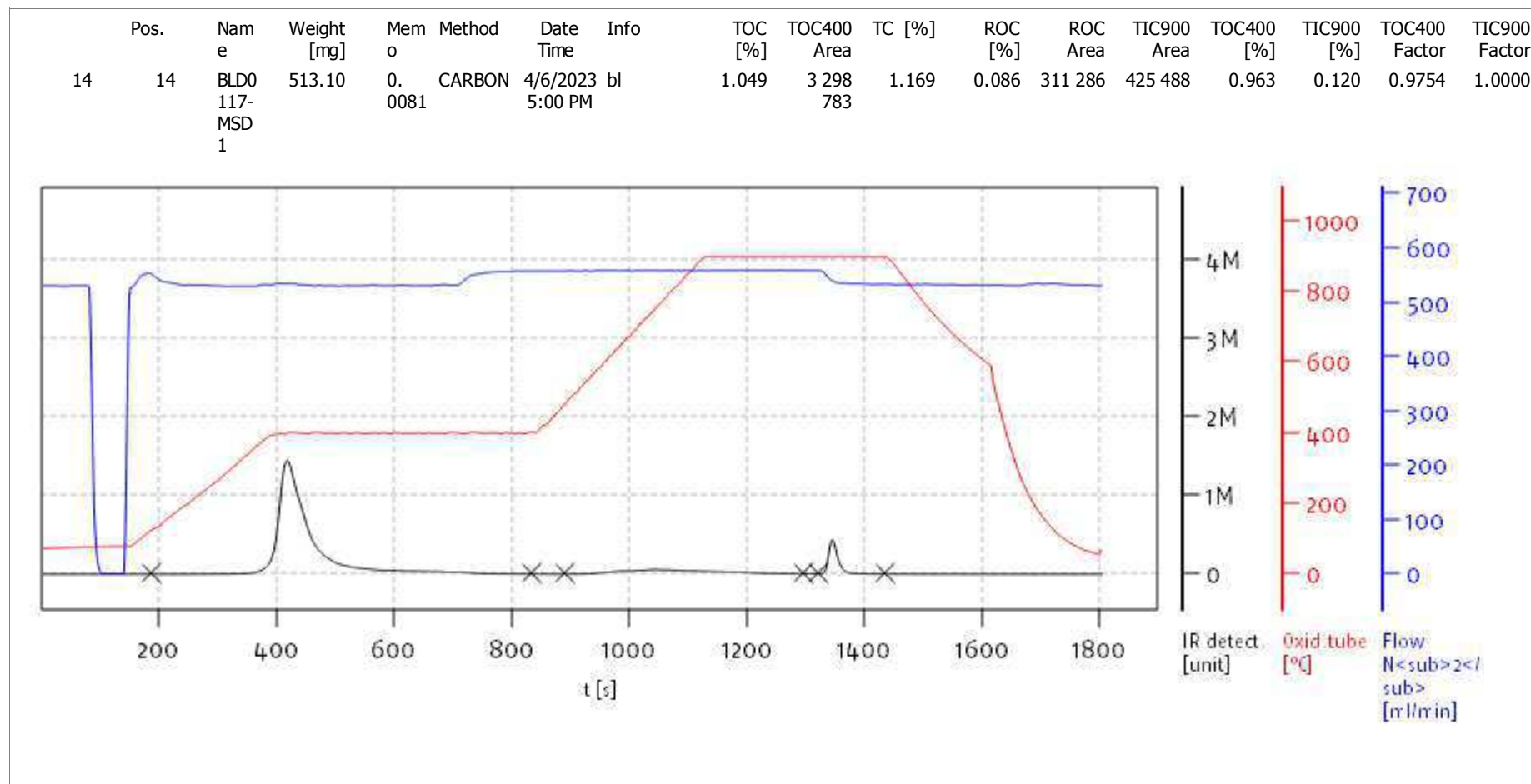
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 Balance: BAL3
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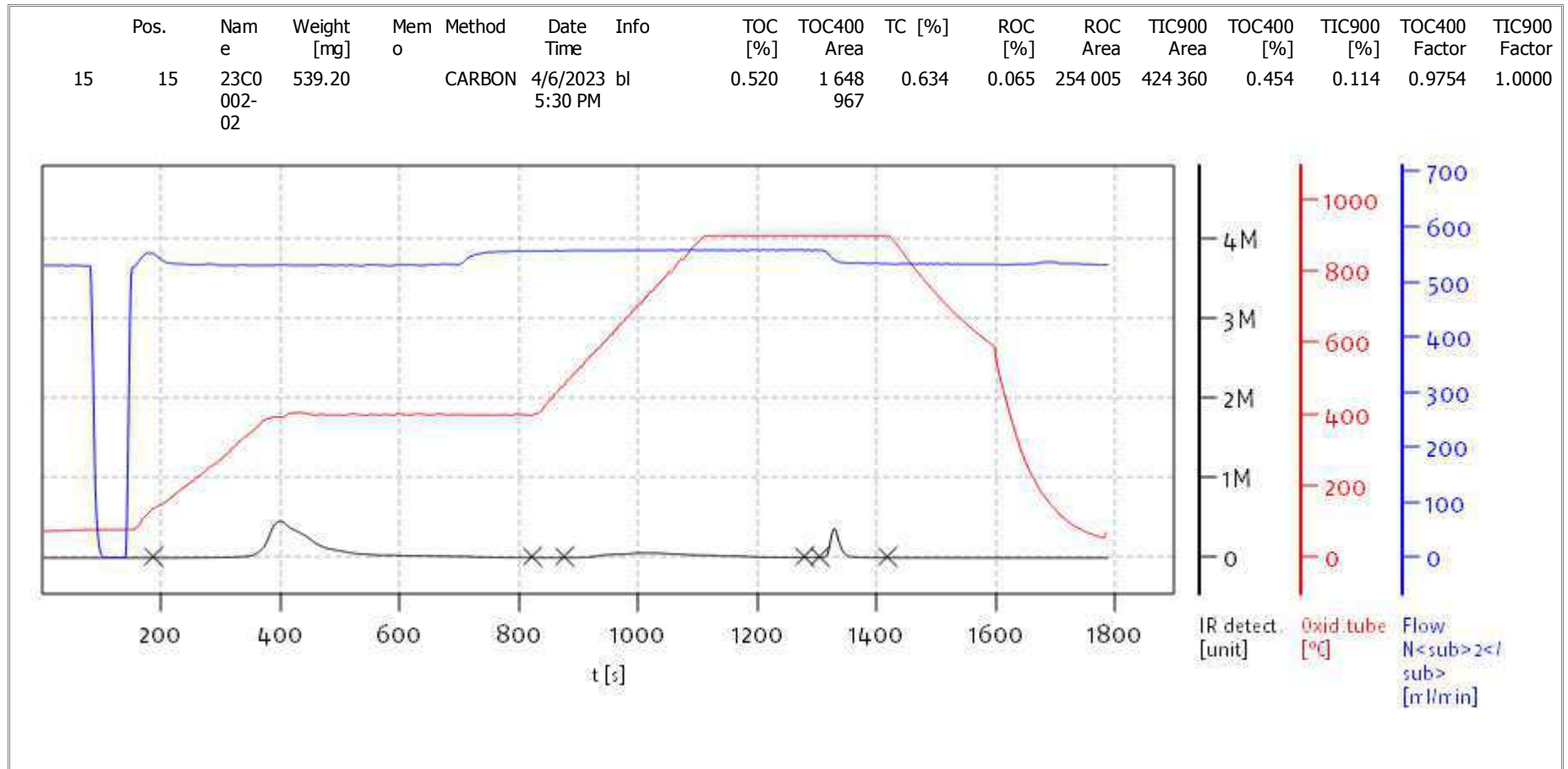
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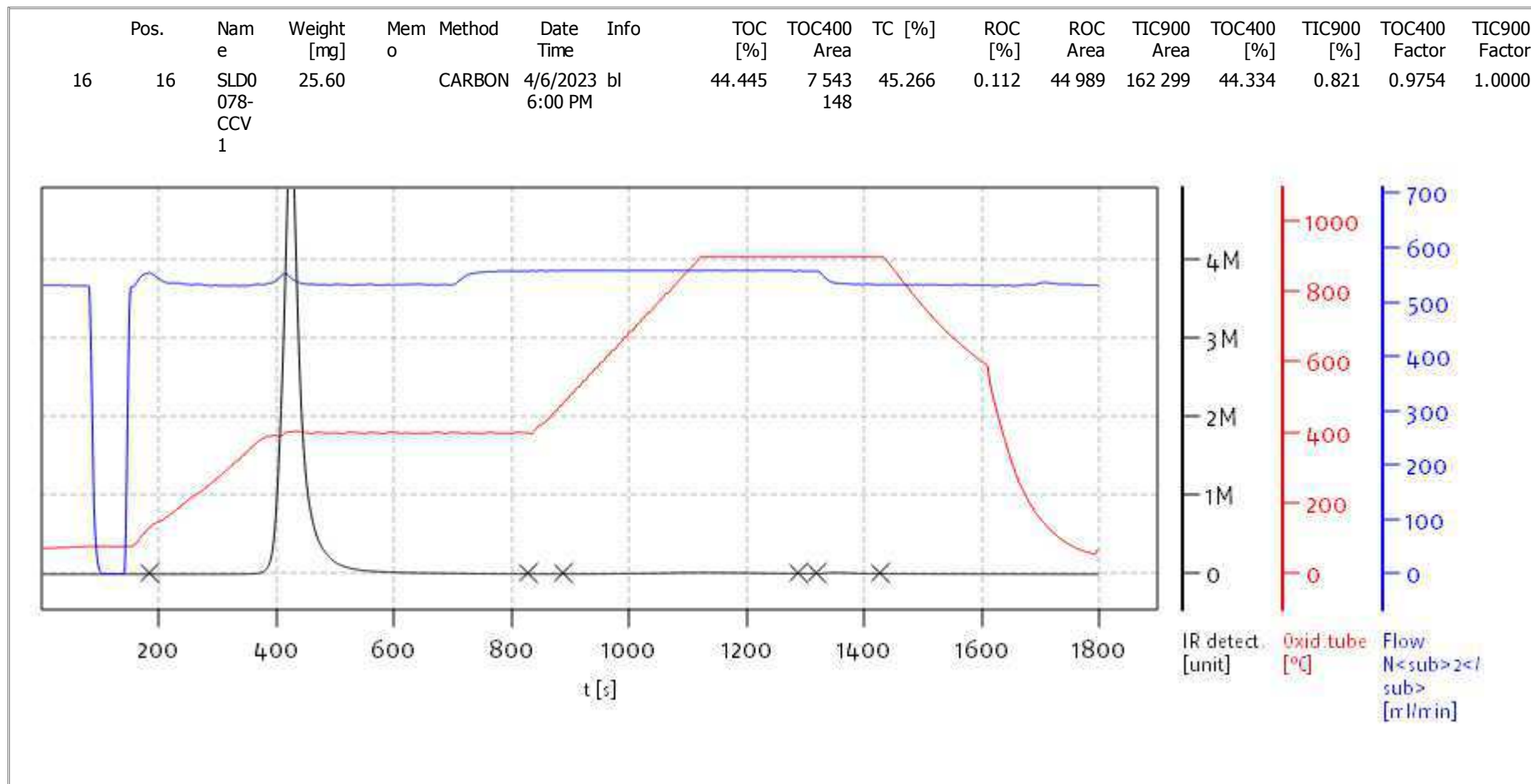
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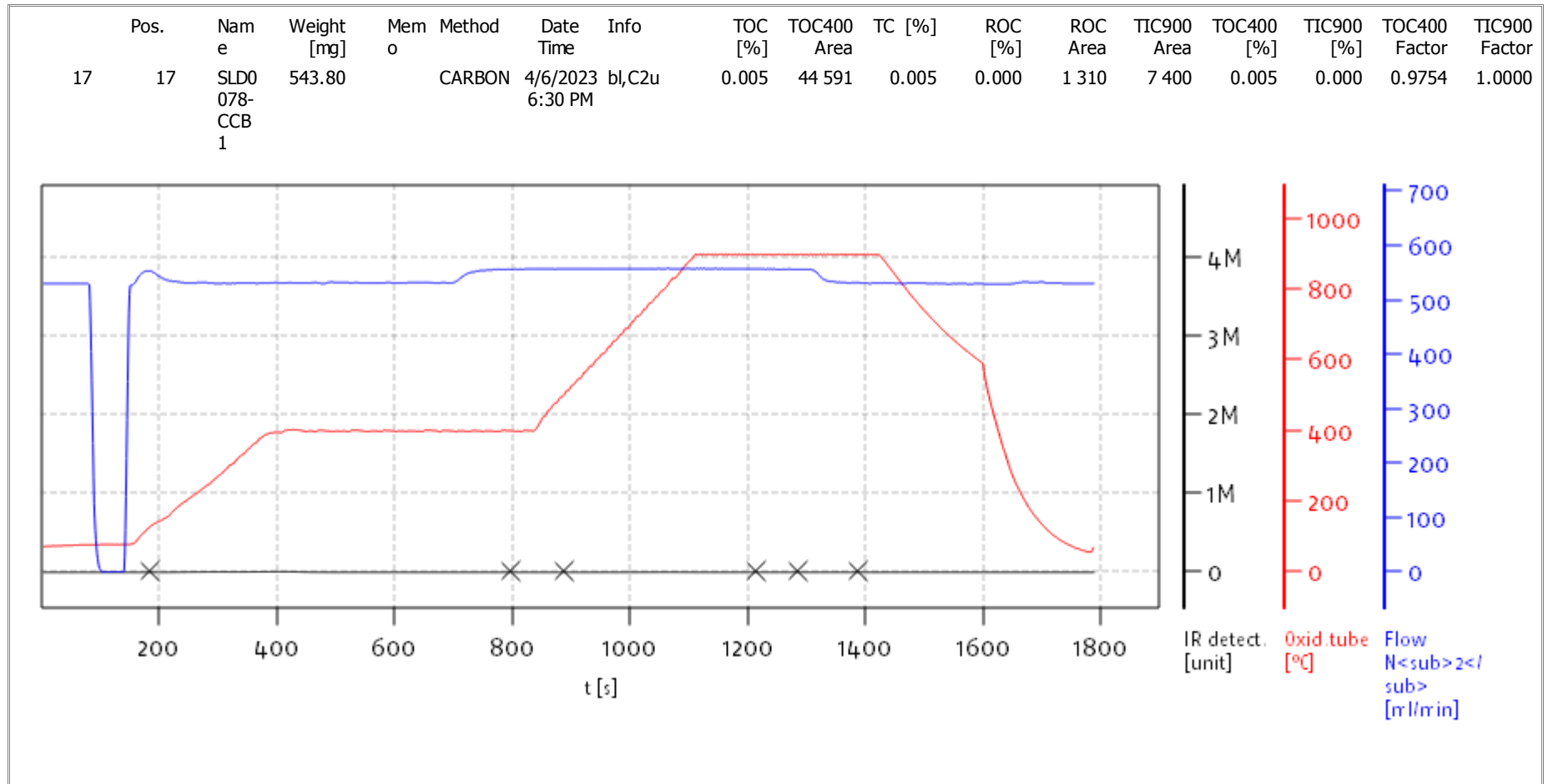
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Balance: BAL3
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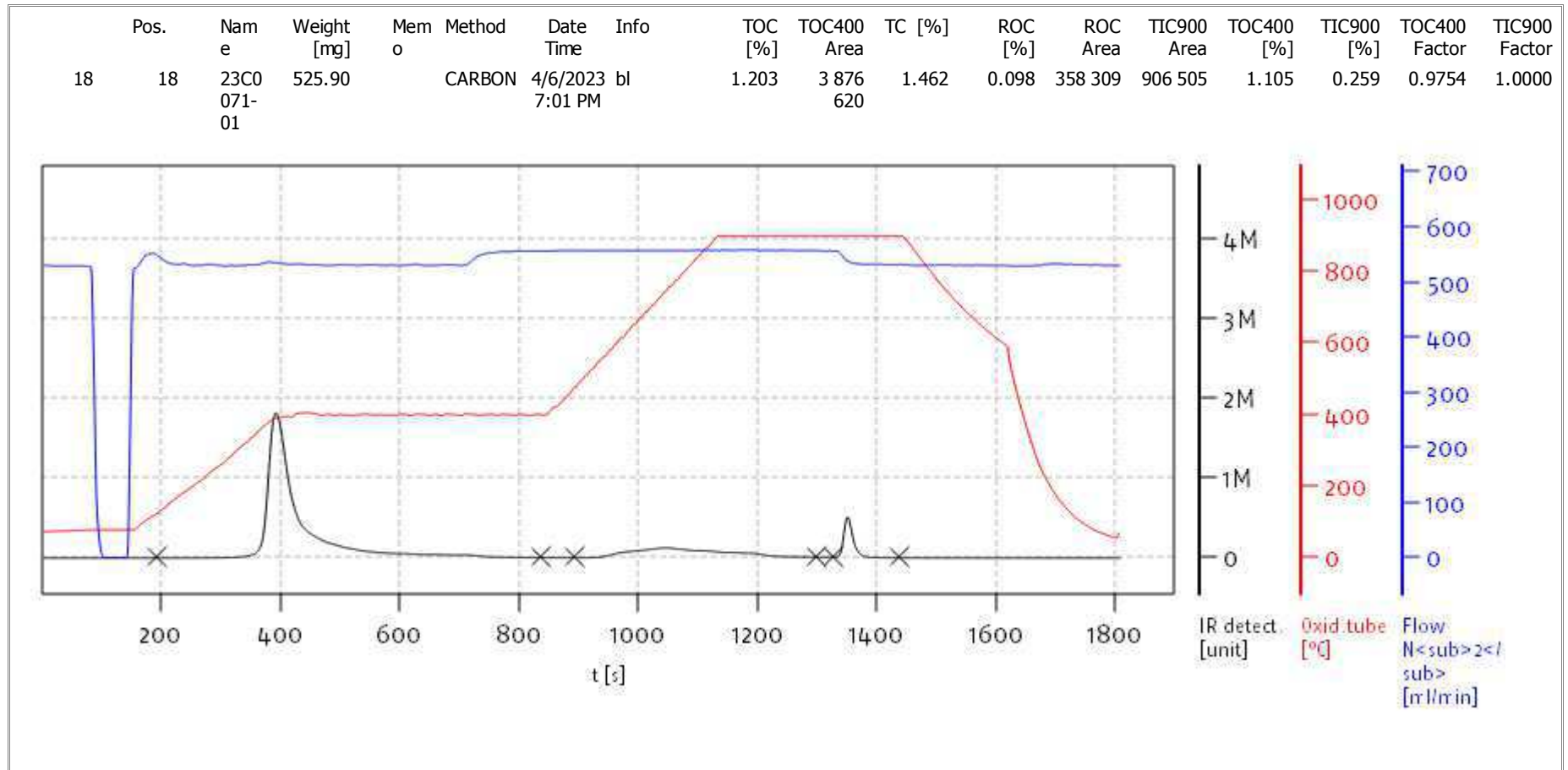
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 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
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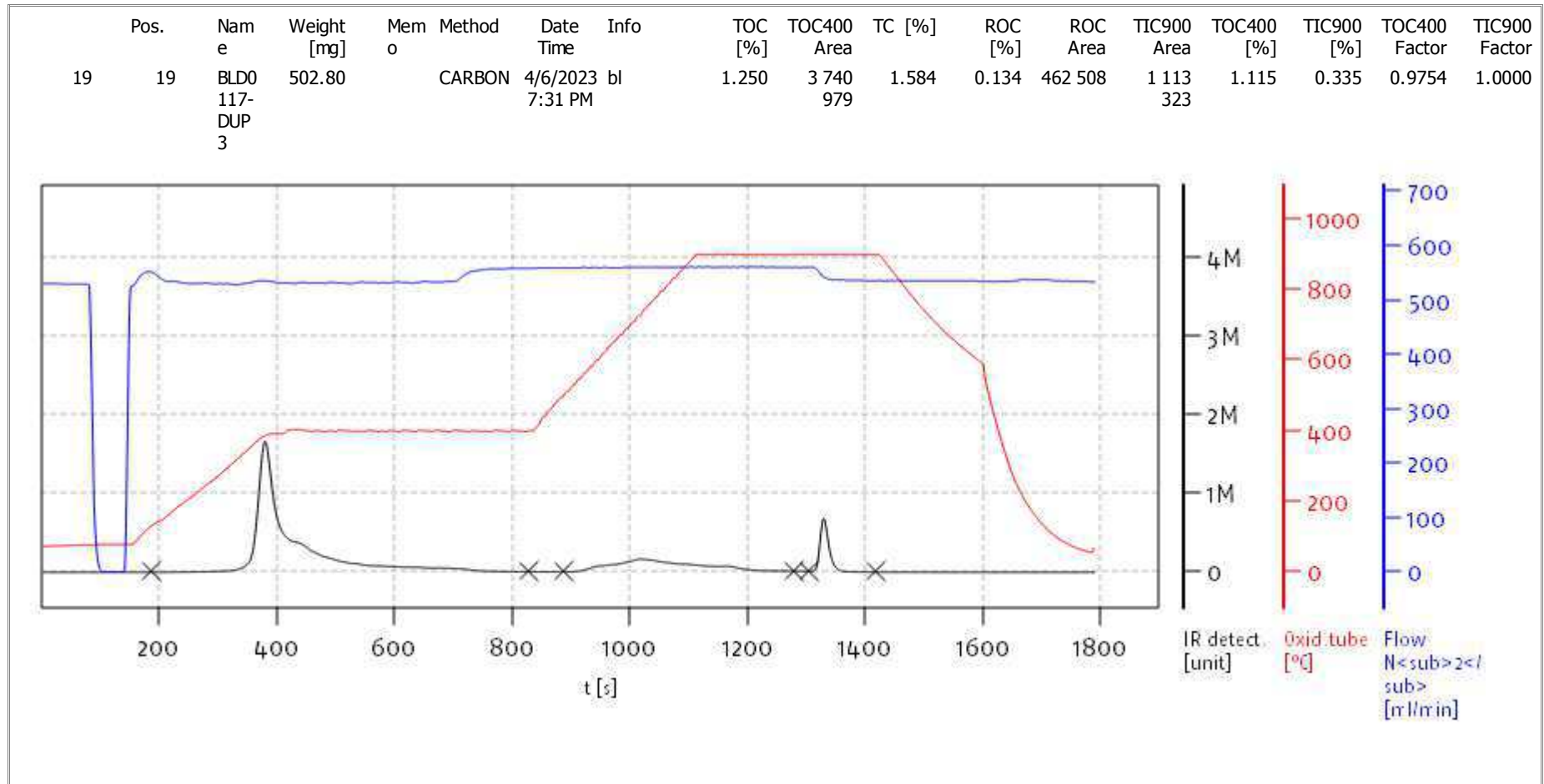
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 Serial No: 0300.181017
 Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: CDE



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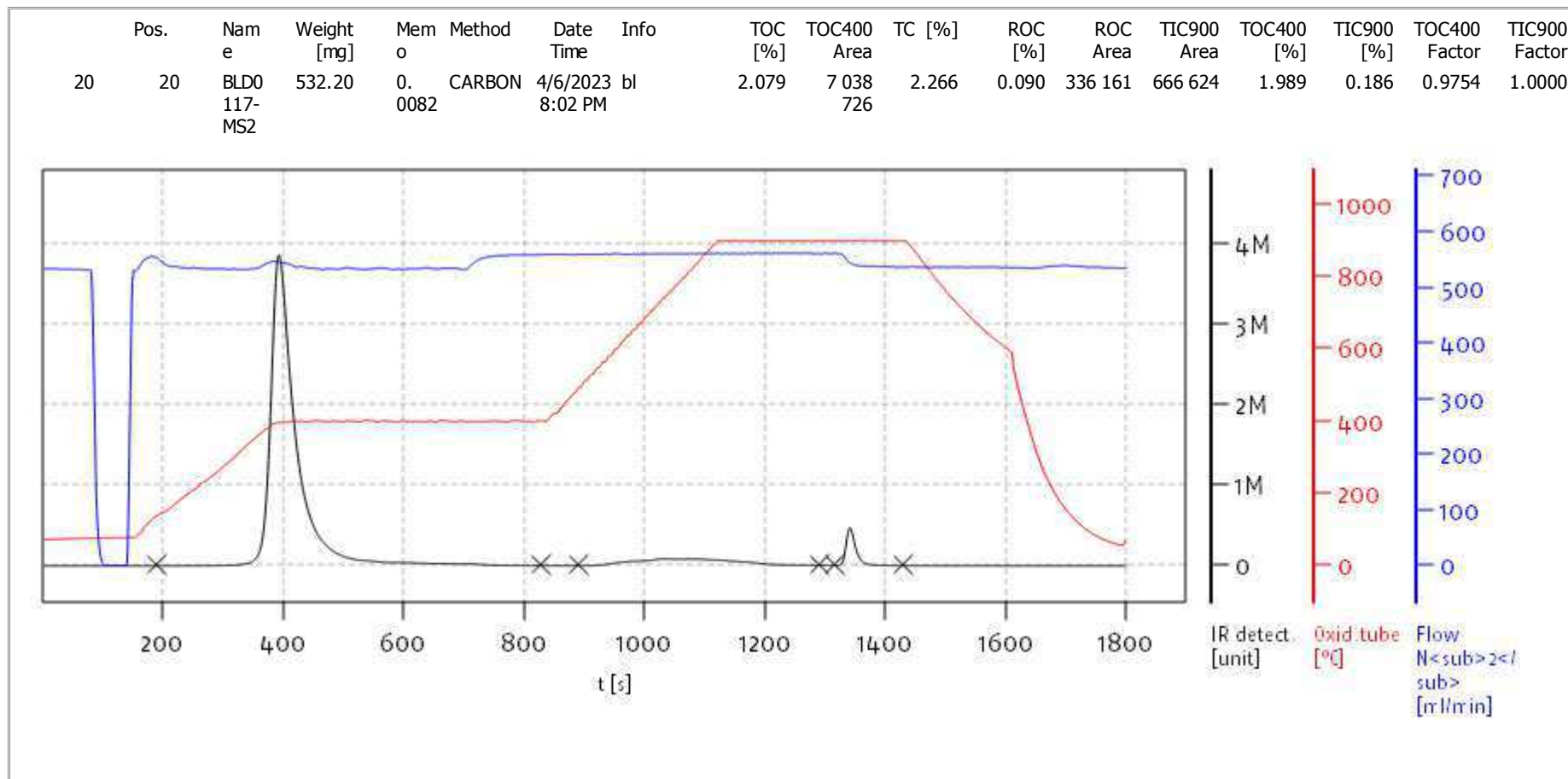
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Soli TOC Cube, Carbon
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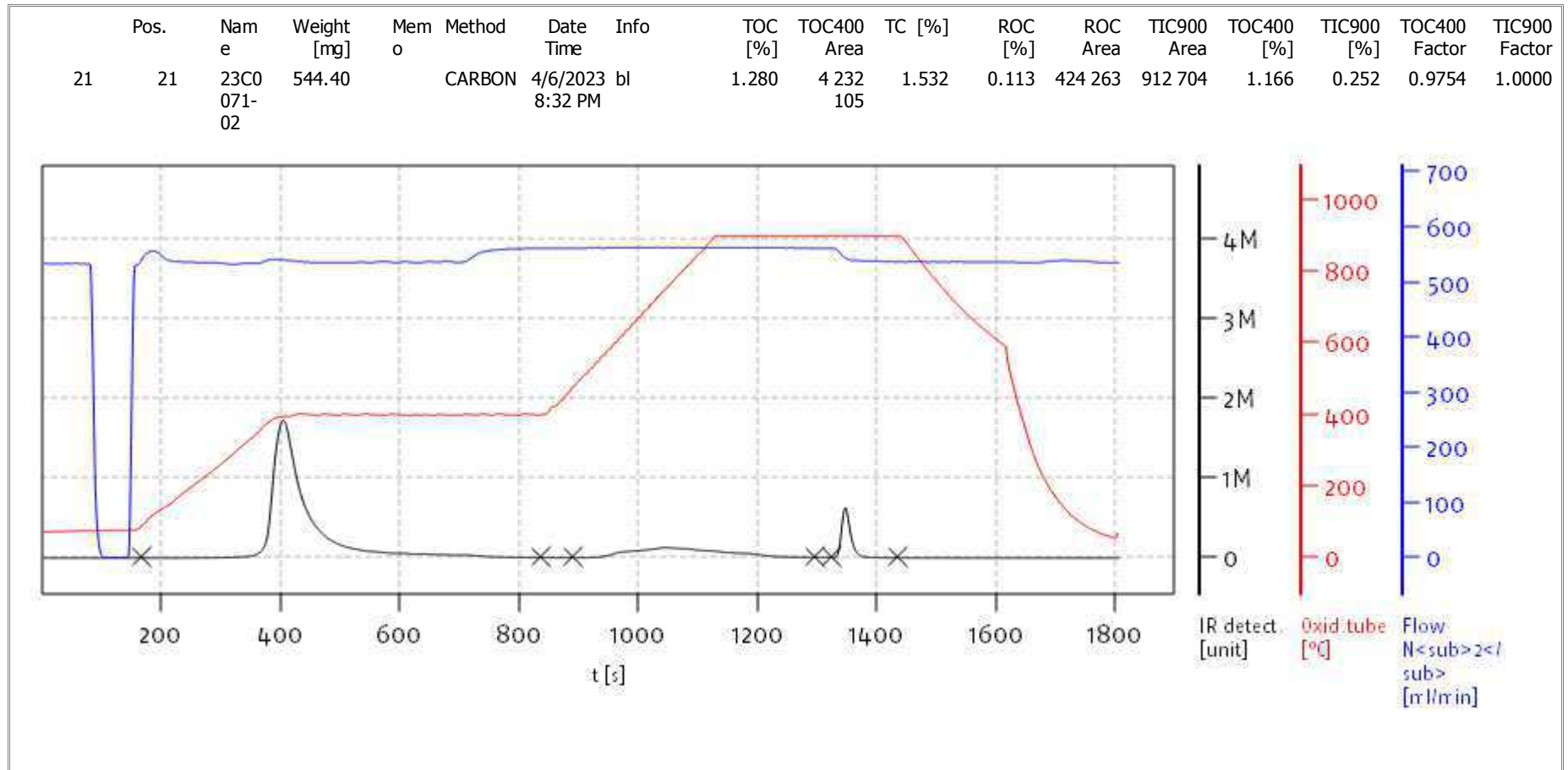
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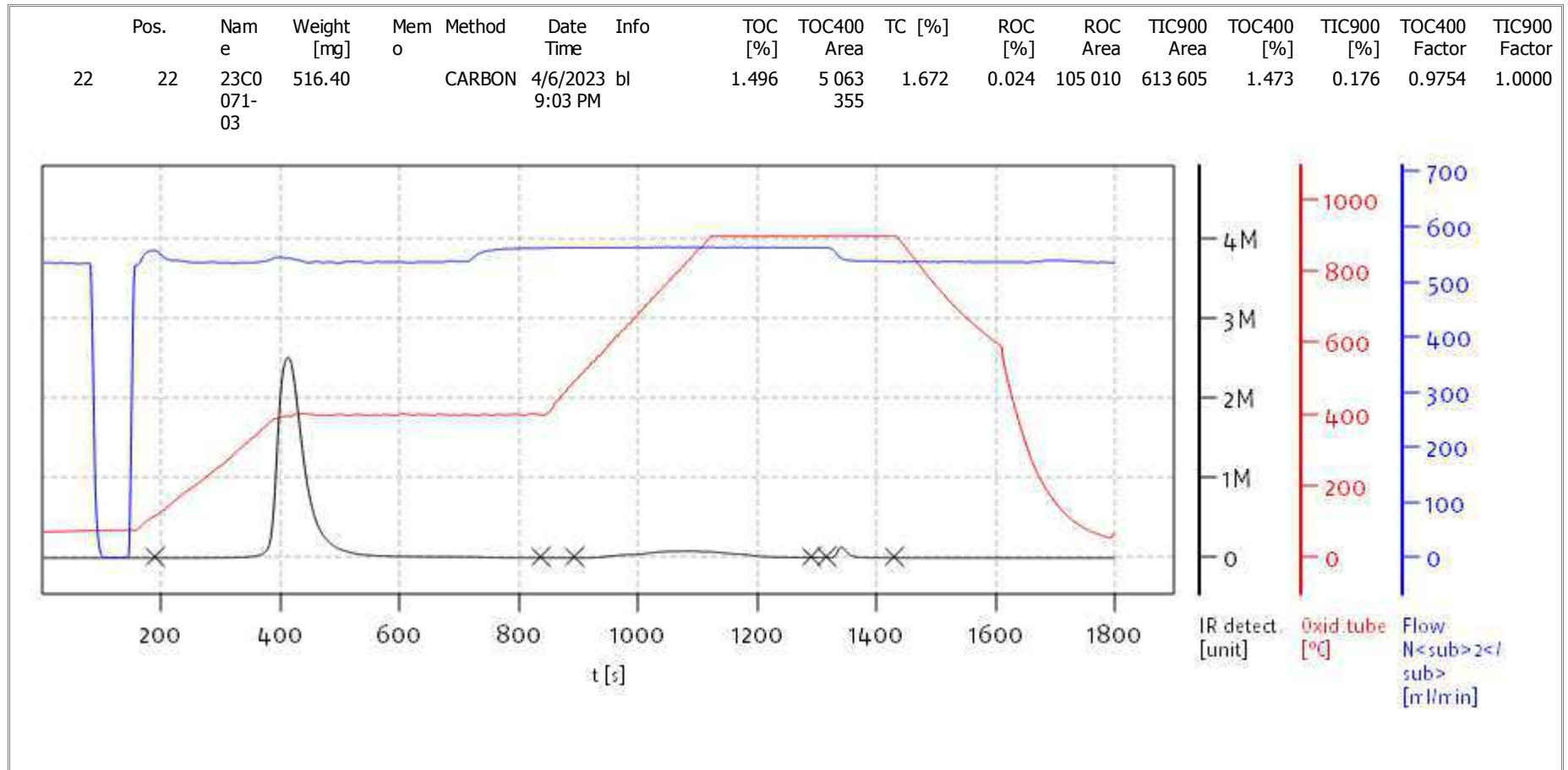
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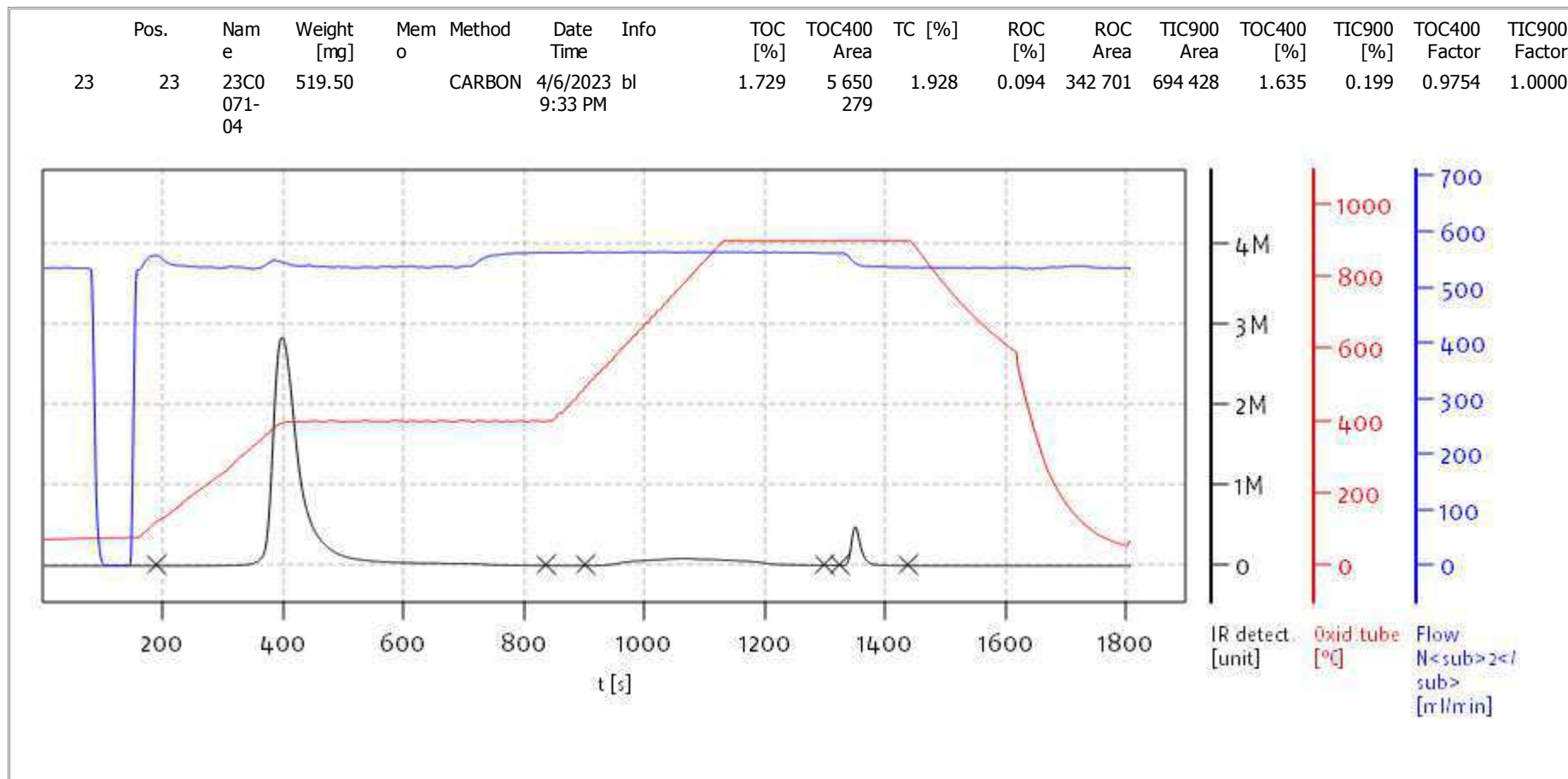
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Soli TOC Cube, Carbon
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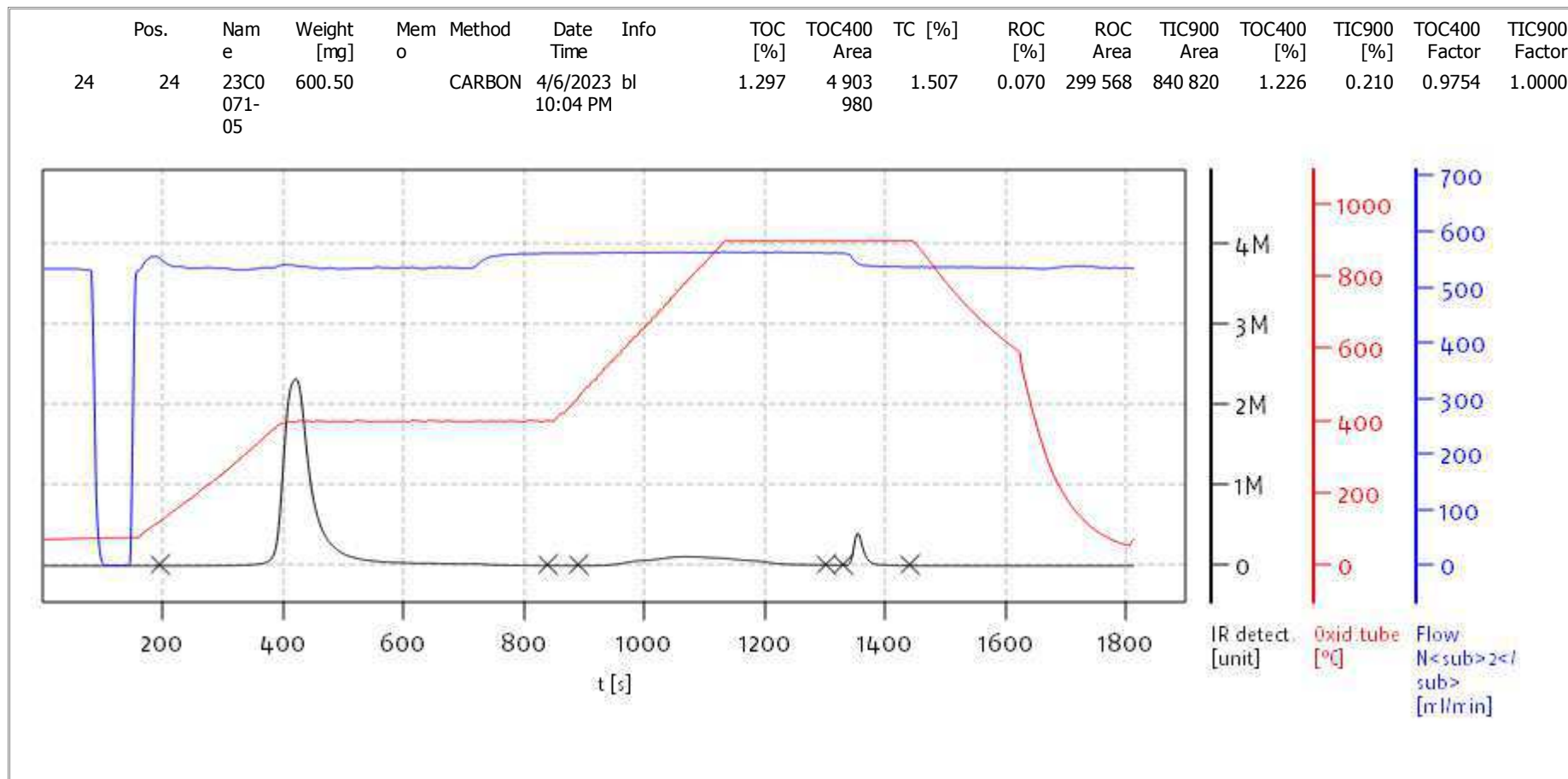
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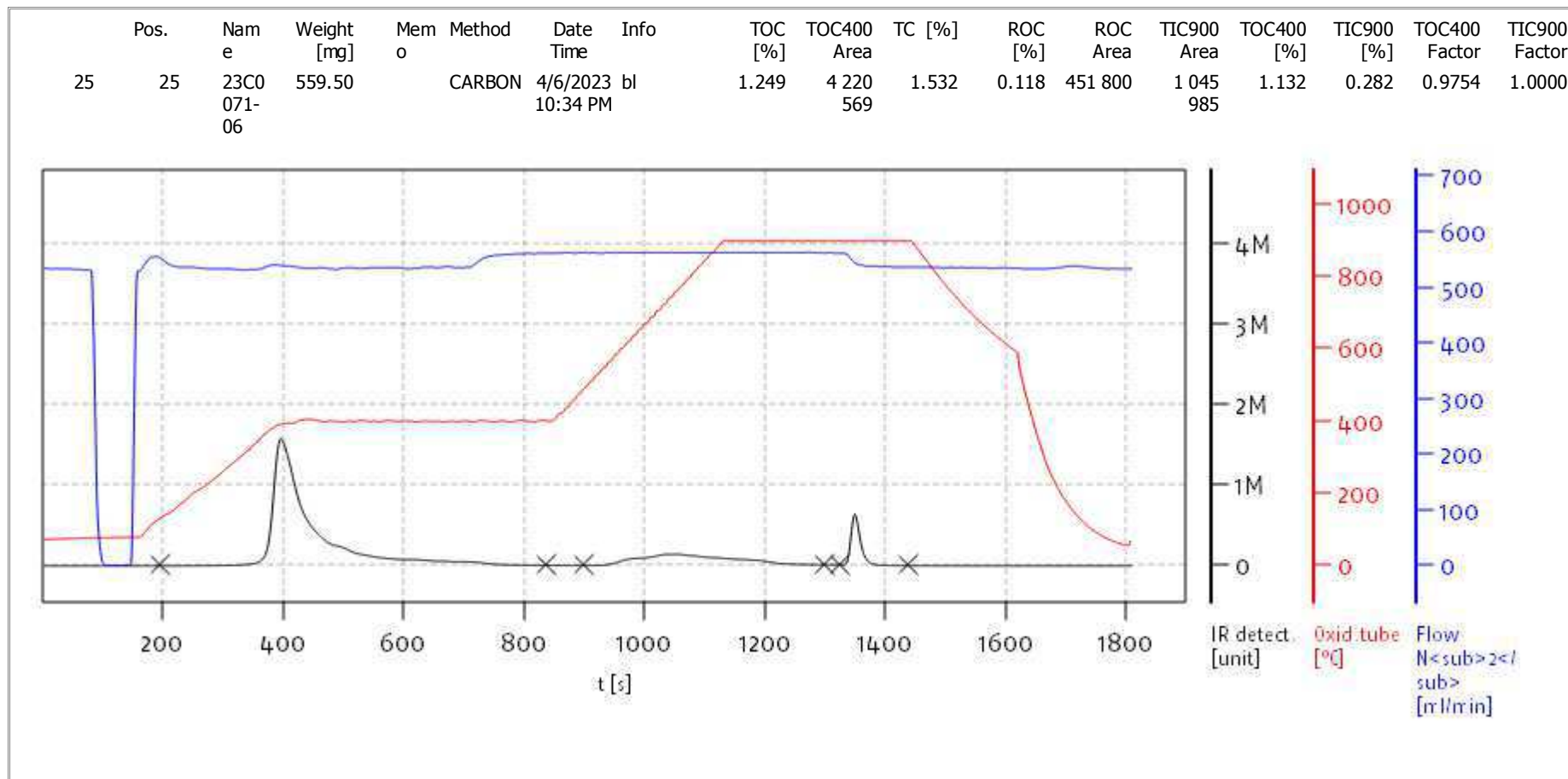
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Name:

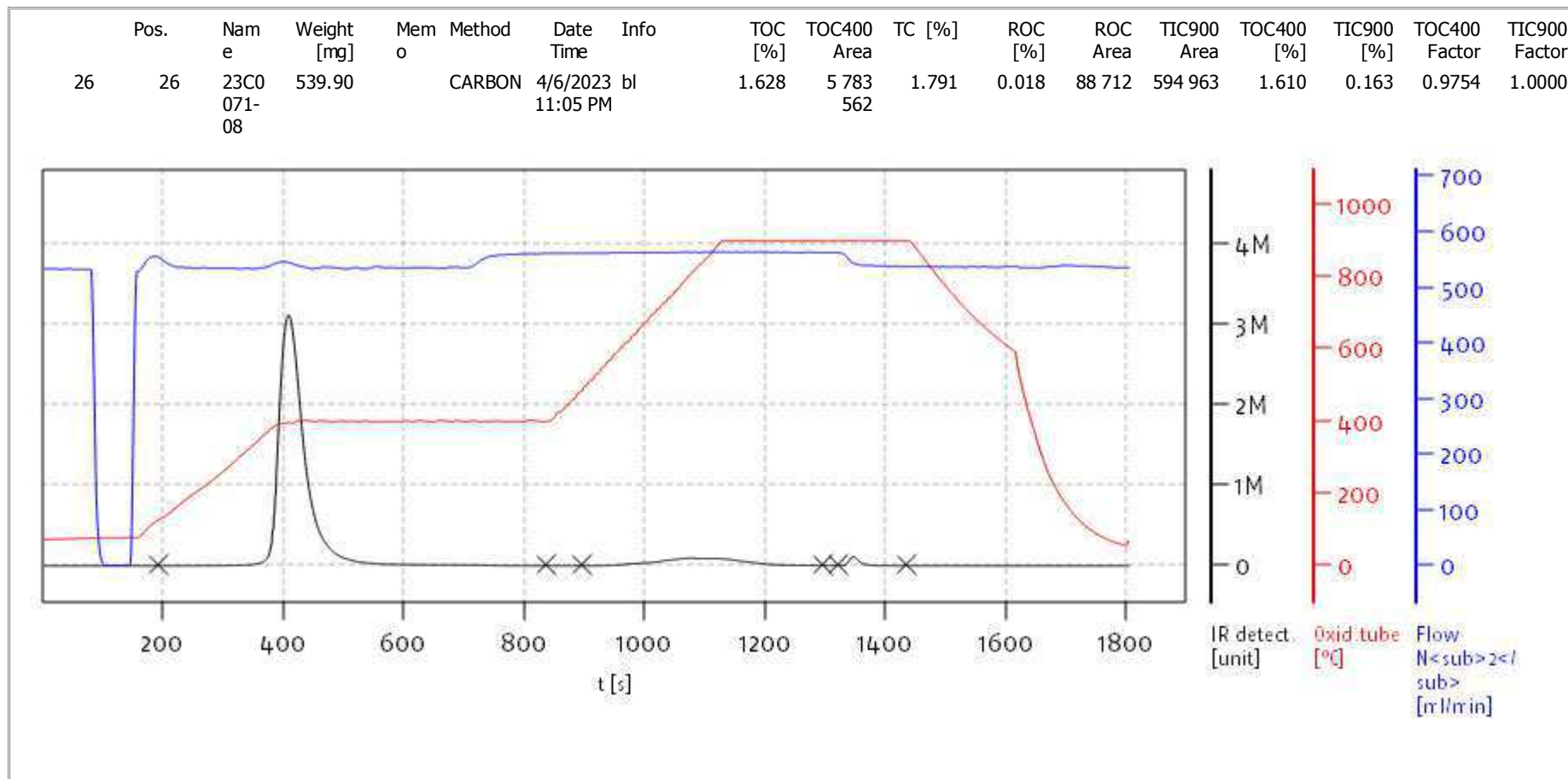
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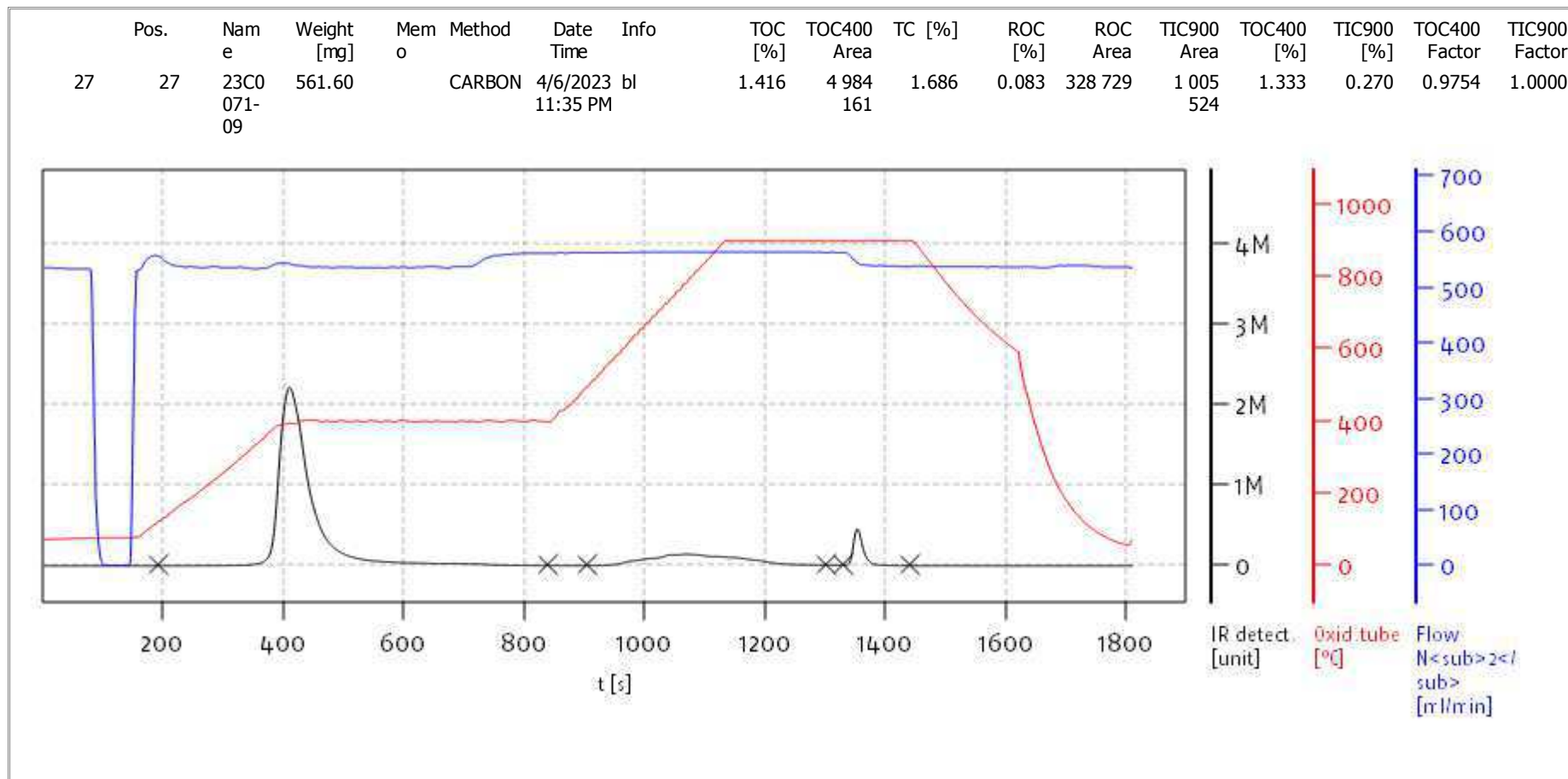
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Soli TOC Cube, Carbon
 Balance: BAL3
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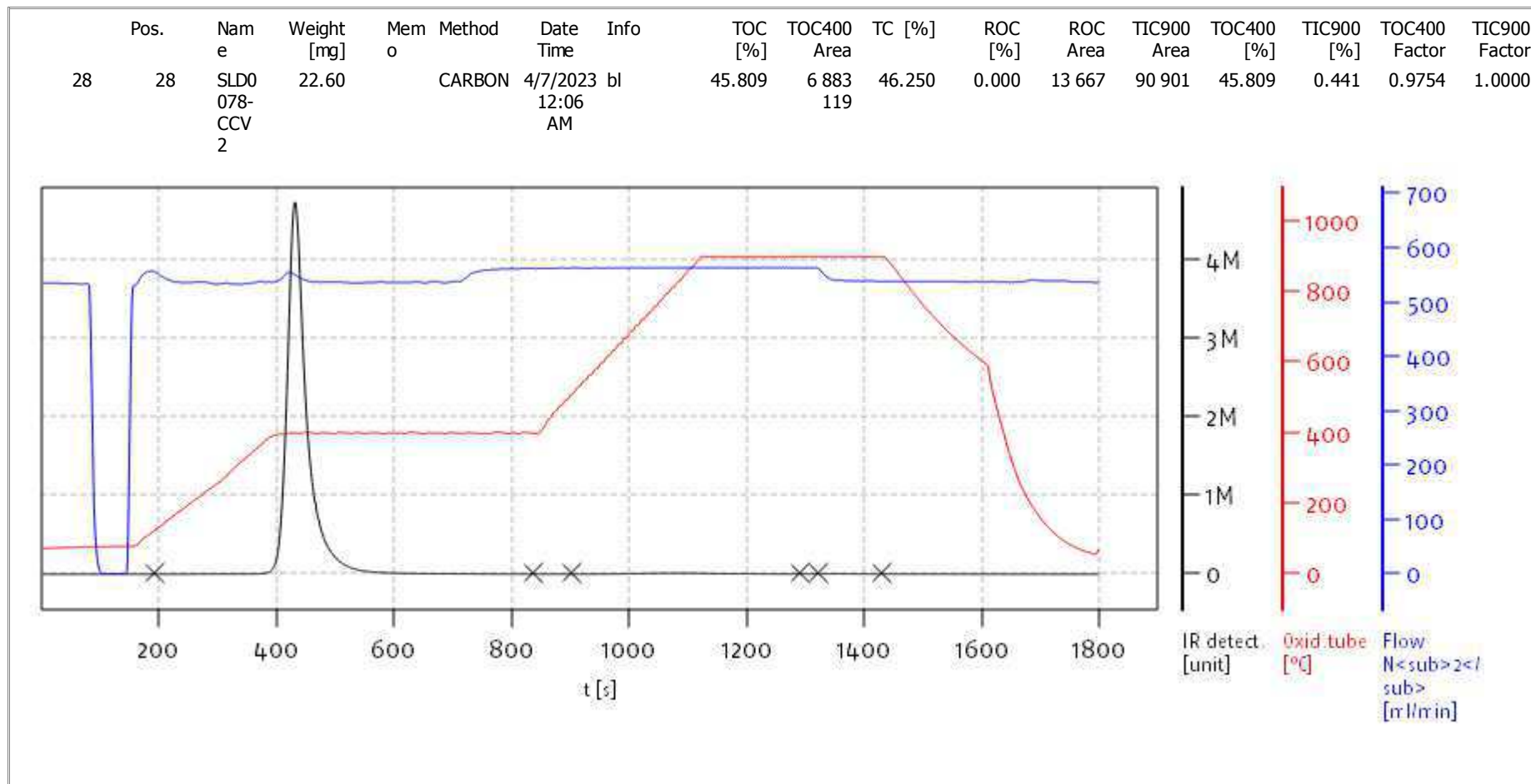
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Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

Access: solITOC superuser

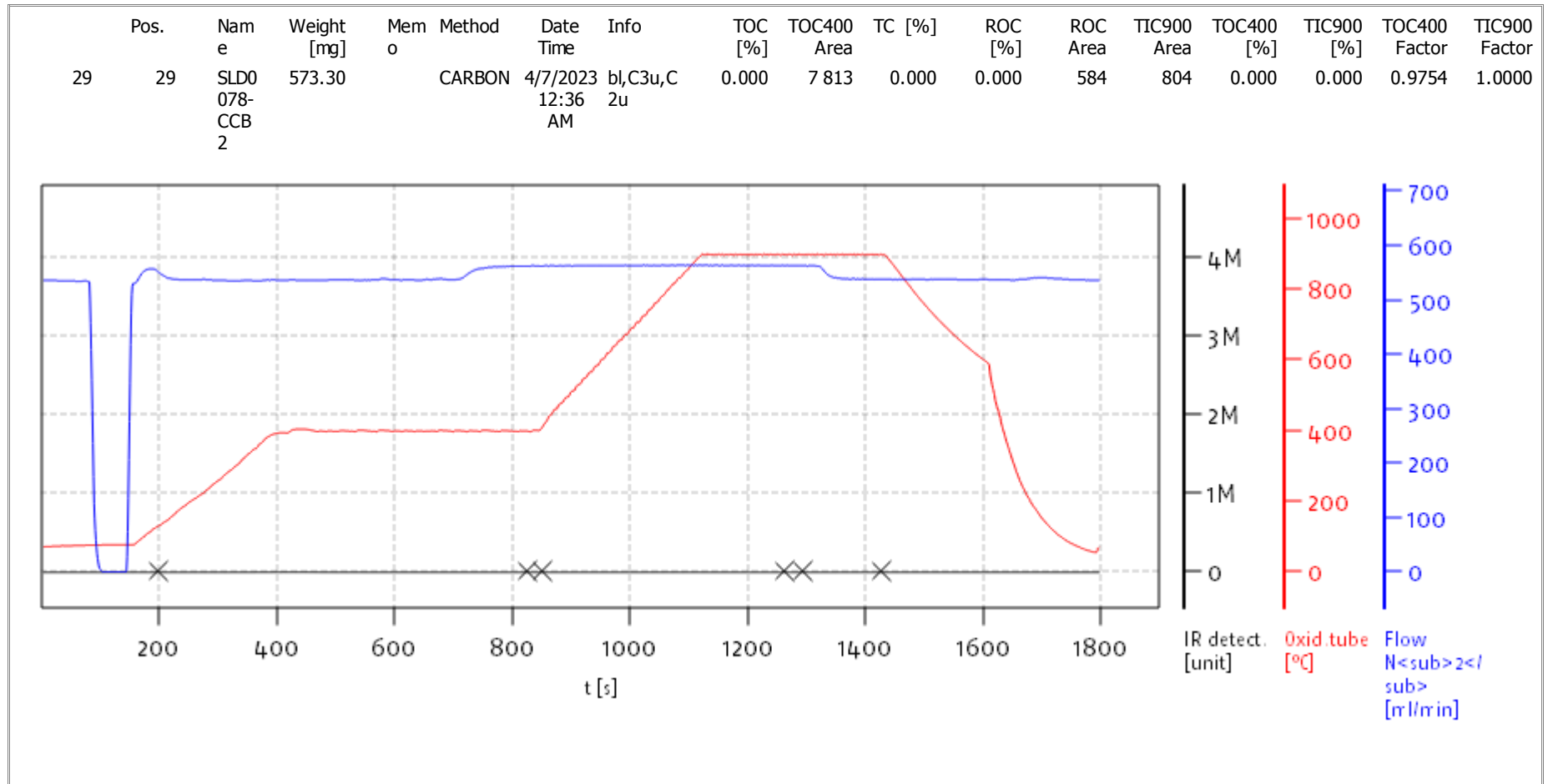
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Soli TOC Cube, Carbon
Balance: BAL3
Analyst: CDE



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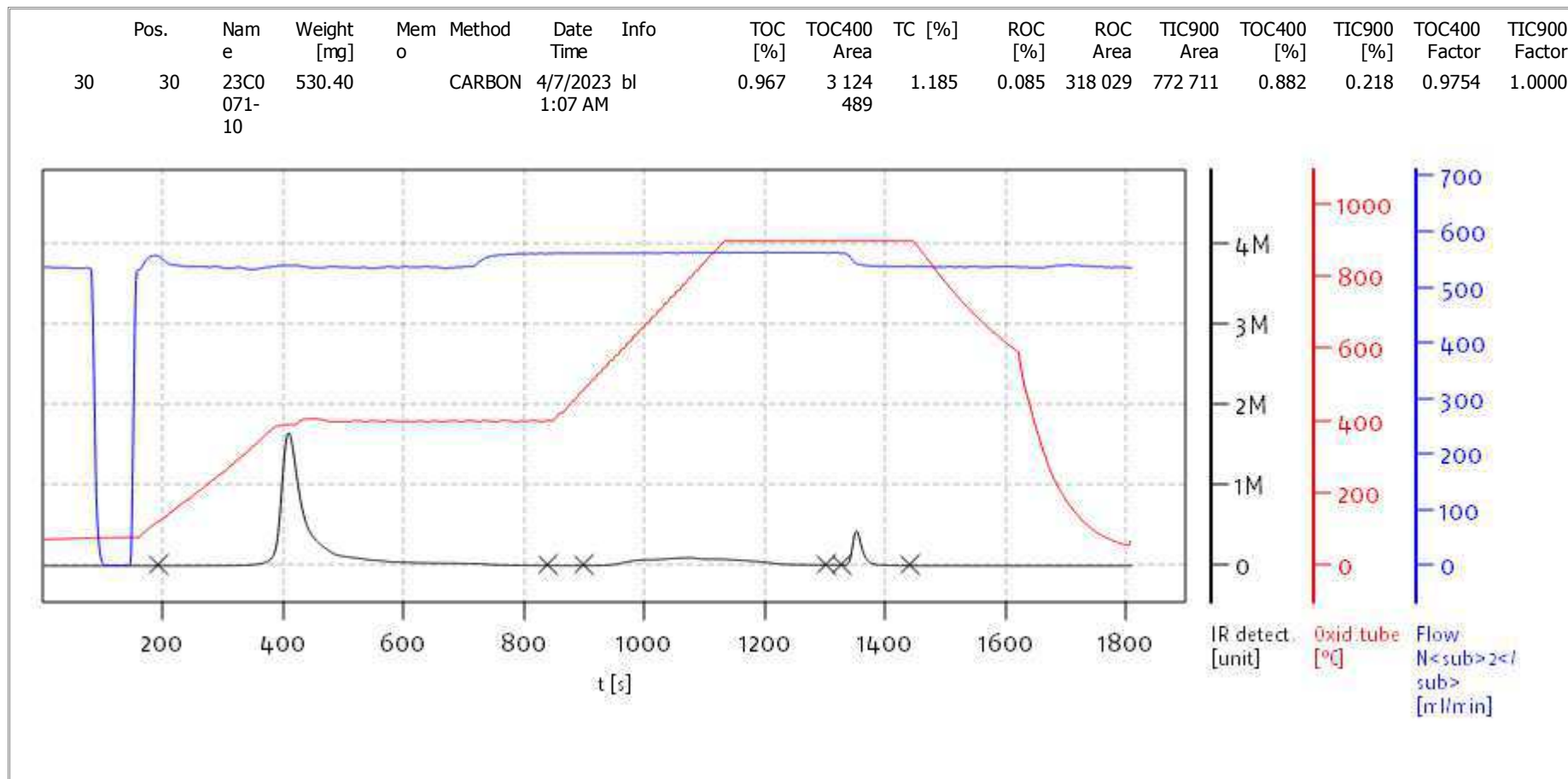
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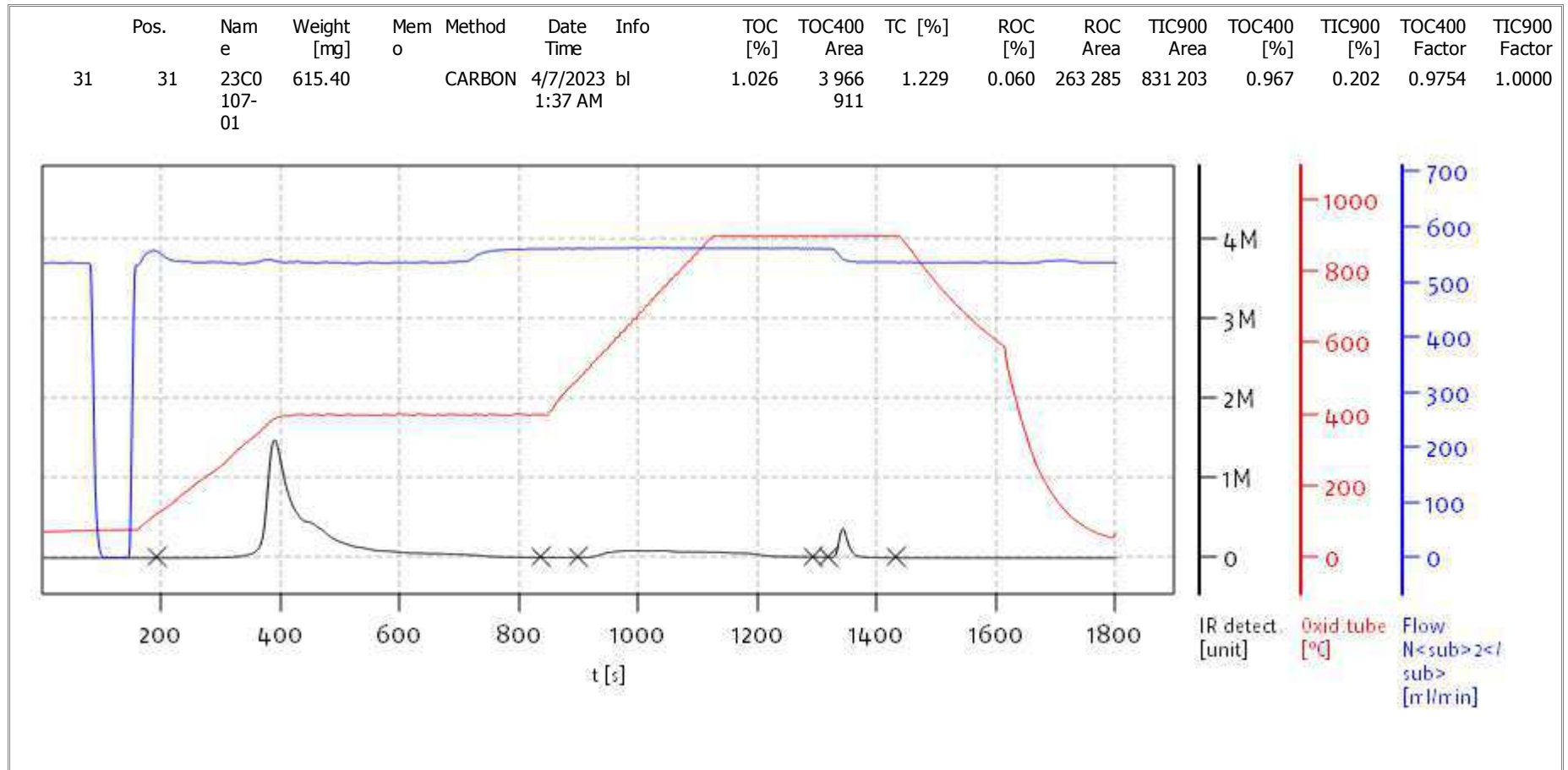
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Soli TOC Cube, Carbon
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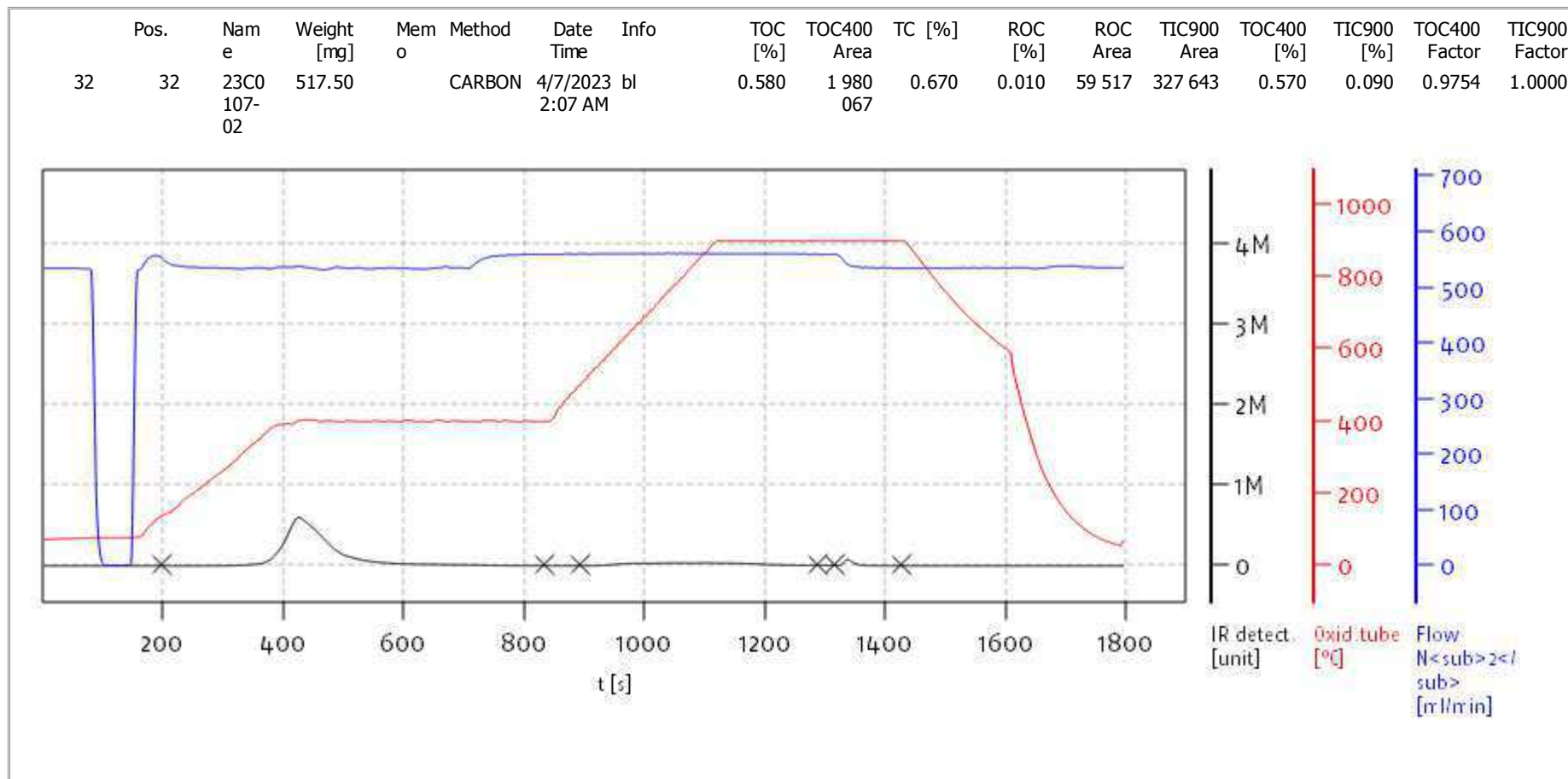
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Soli TOC Cube, Carbon
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Name:

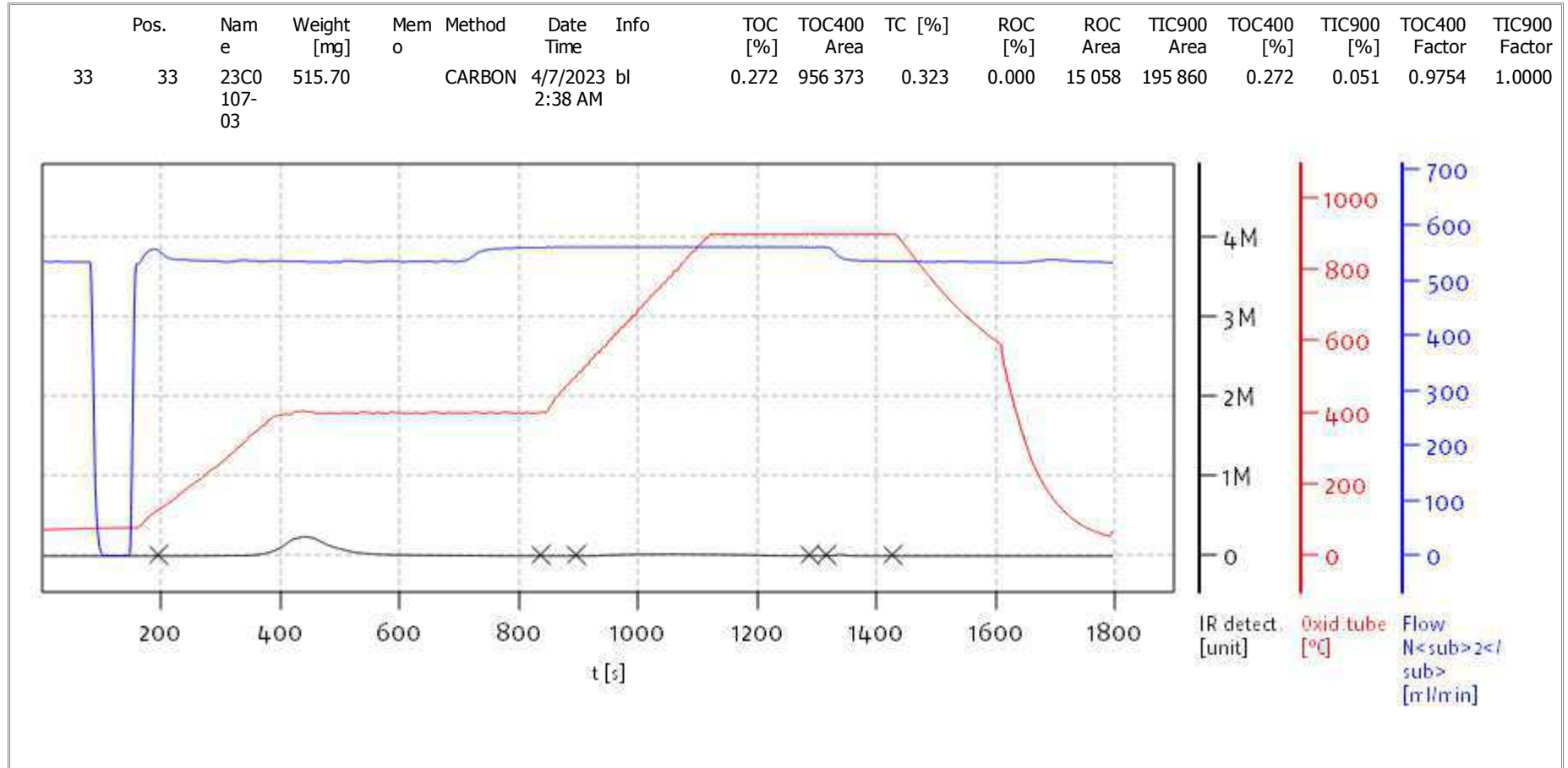
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Soli TOC Cube, Carbon
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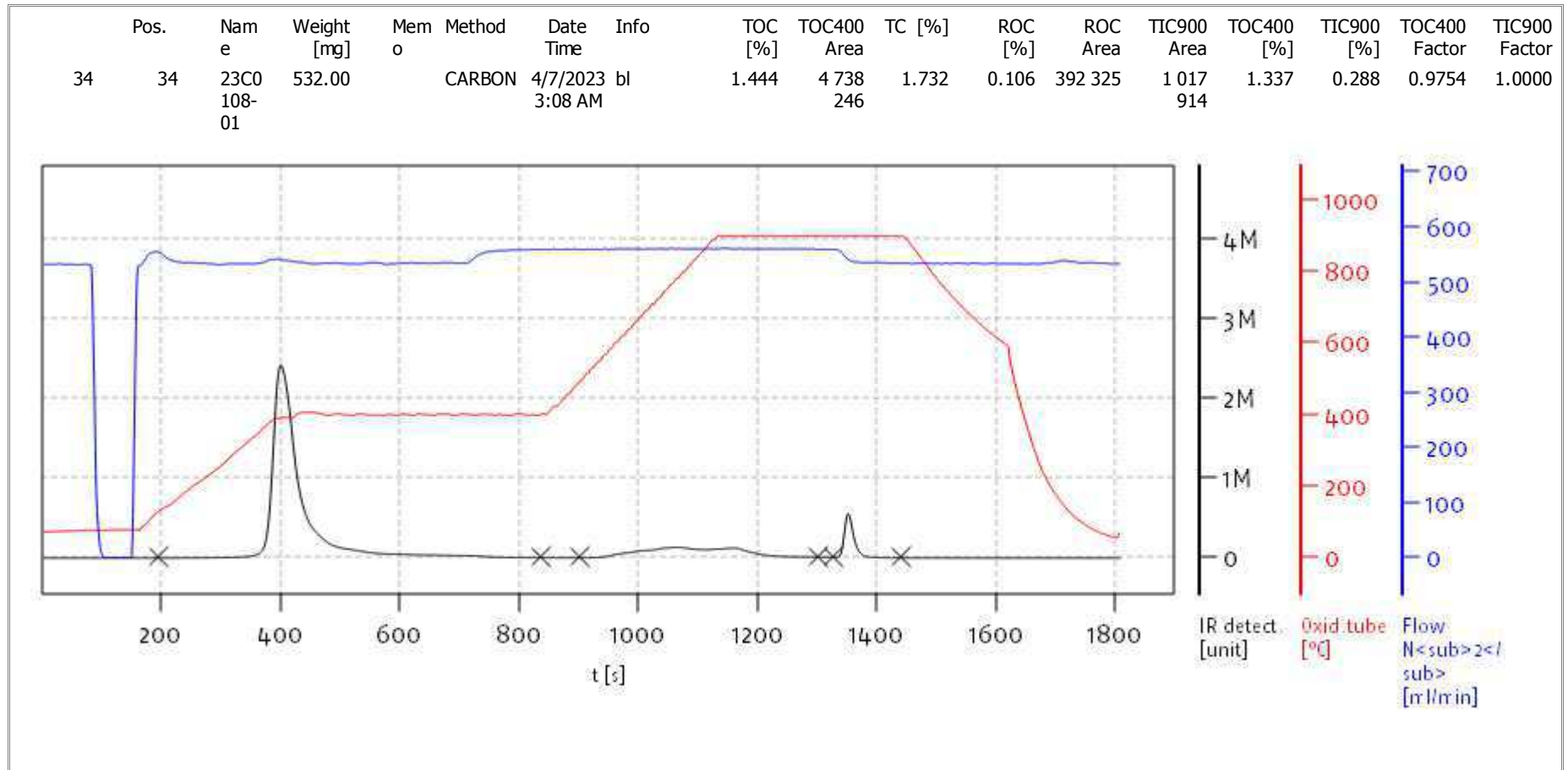
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Soli TOC Cube, Carbon
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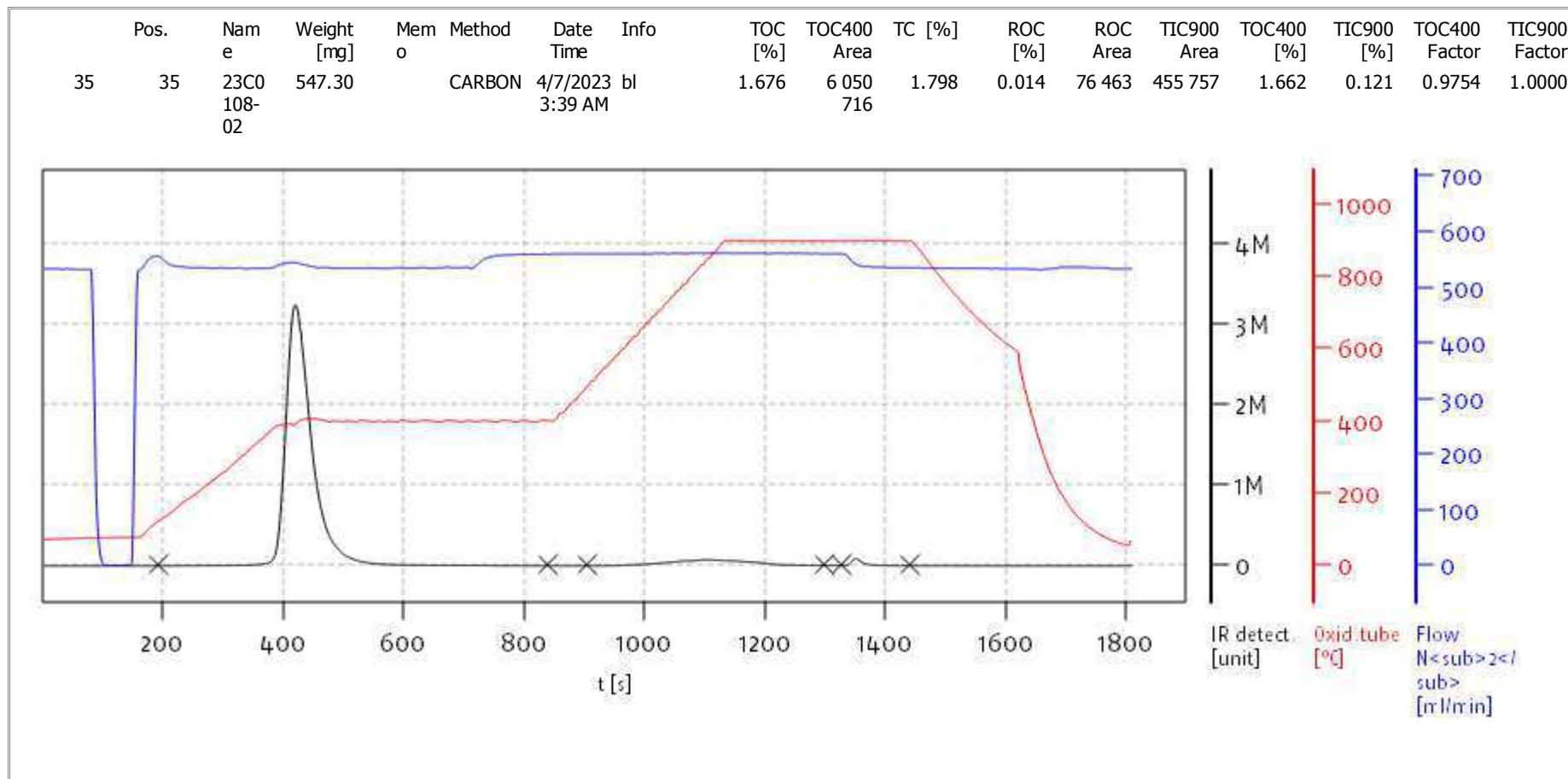
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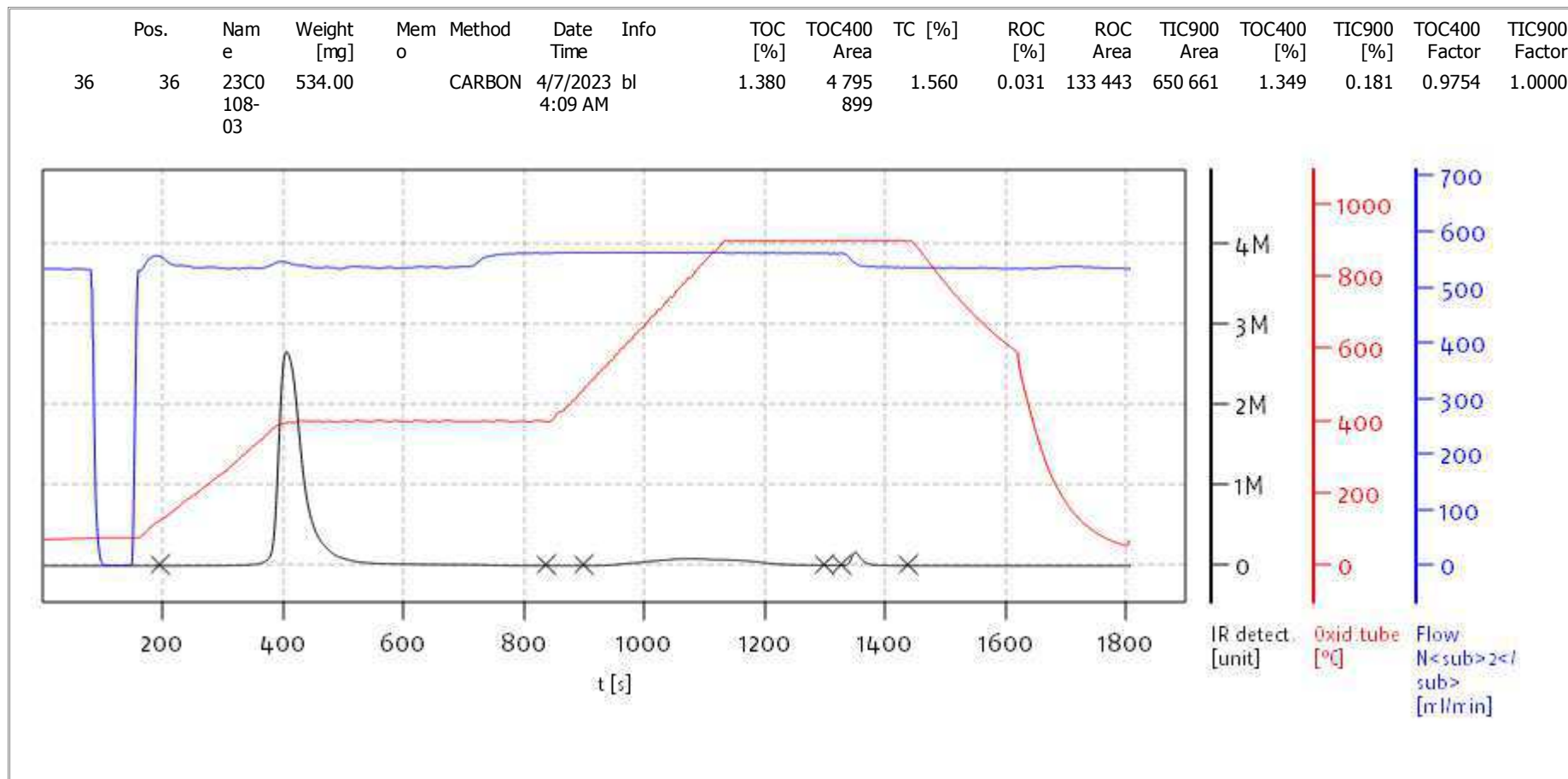
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 Analyst: CDE



Name:

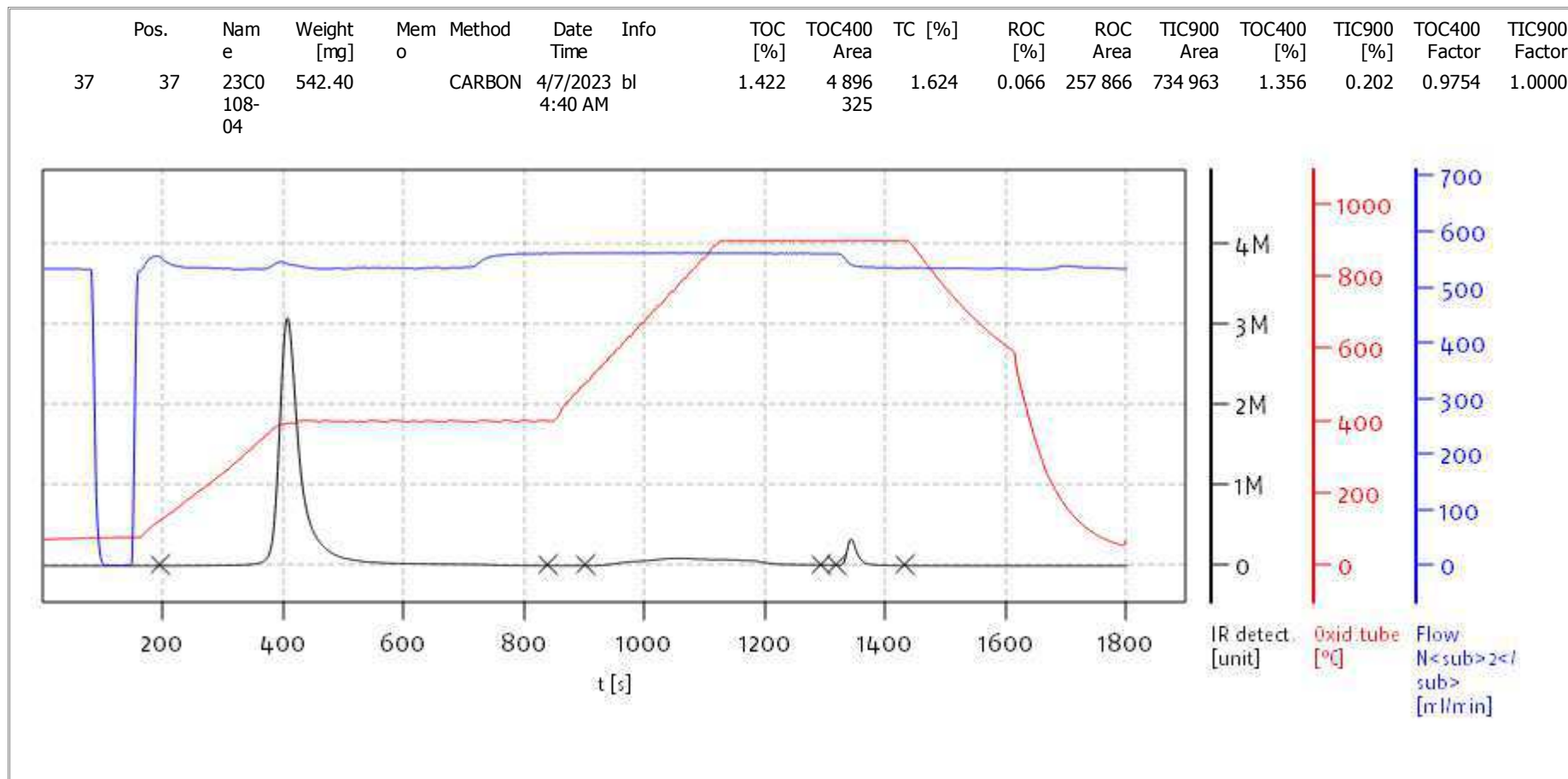
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Date: Fri Apr 7 14:09:03 2023



solITOC V2.0.2 (31015f9) 2018-11-19
 Serial No: 0300.181017
 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

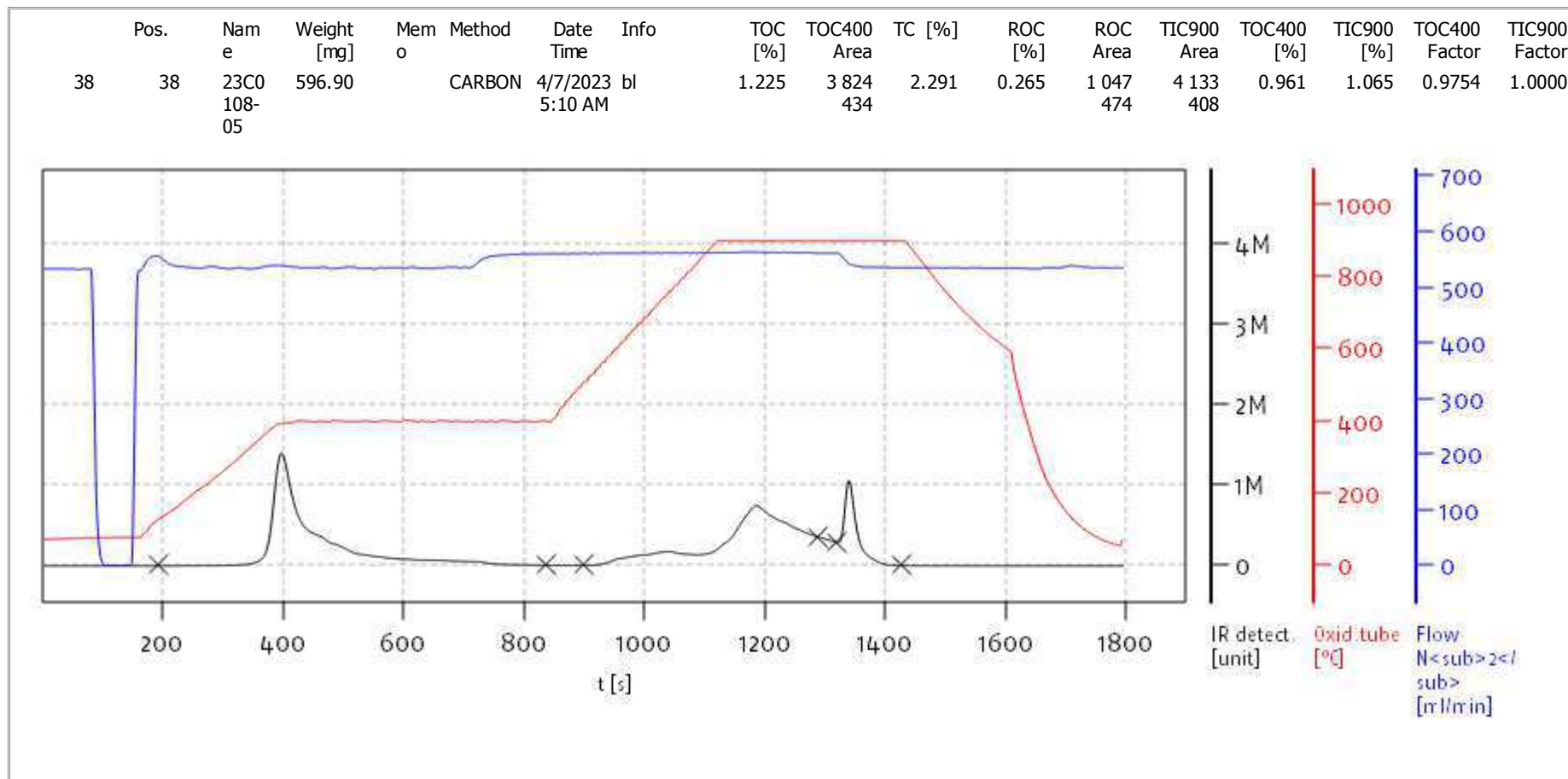
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 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



Name:

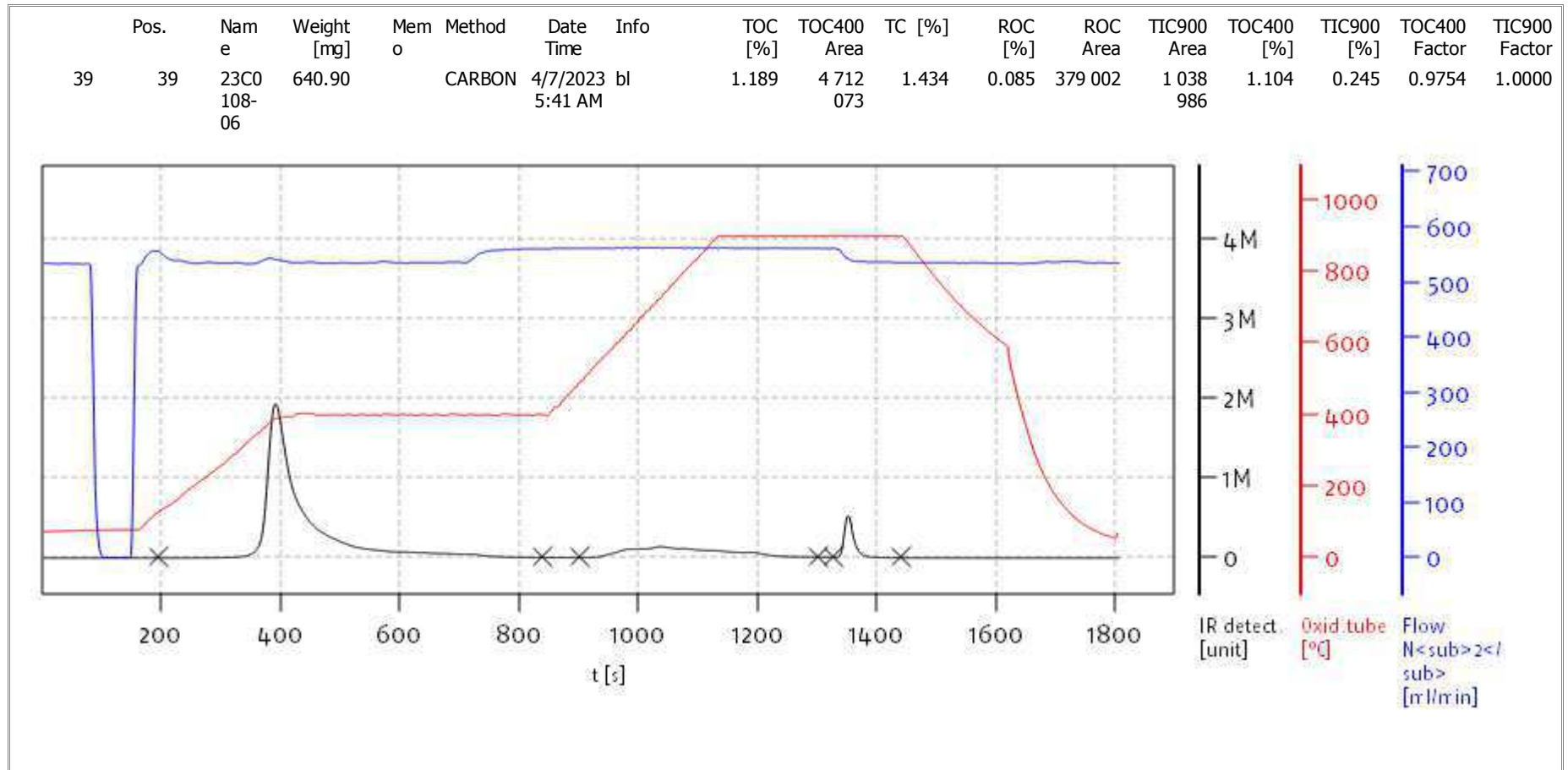
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solITOC V2.0.2 (31015f9) 2018-11-19
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 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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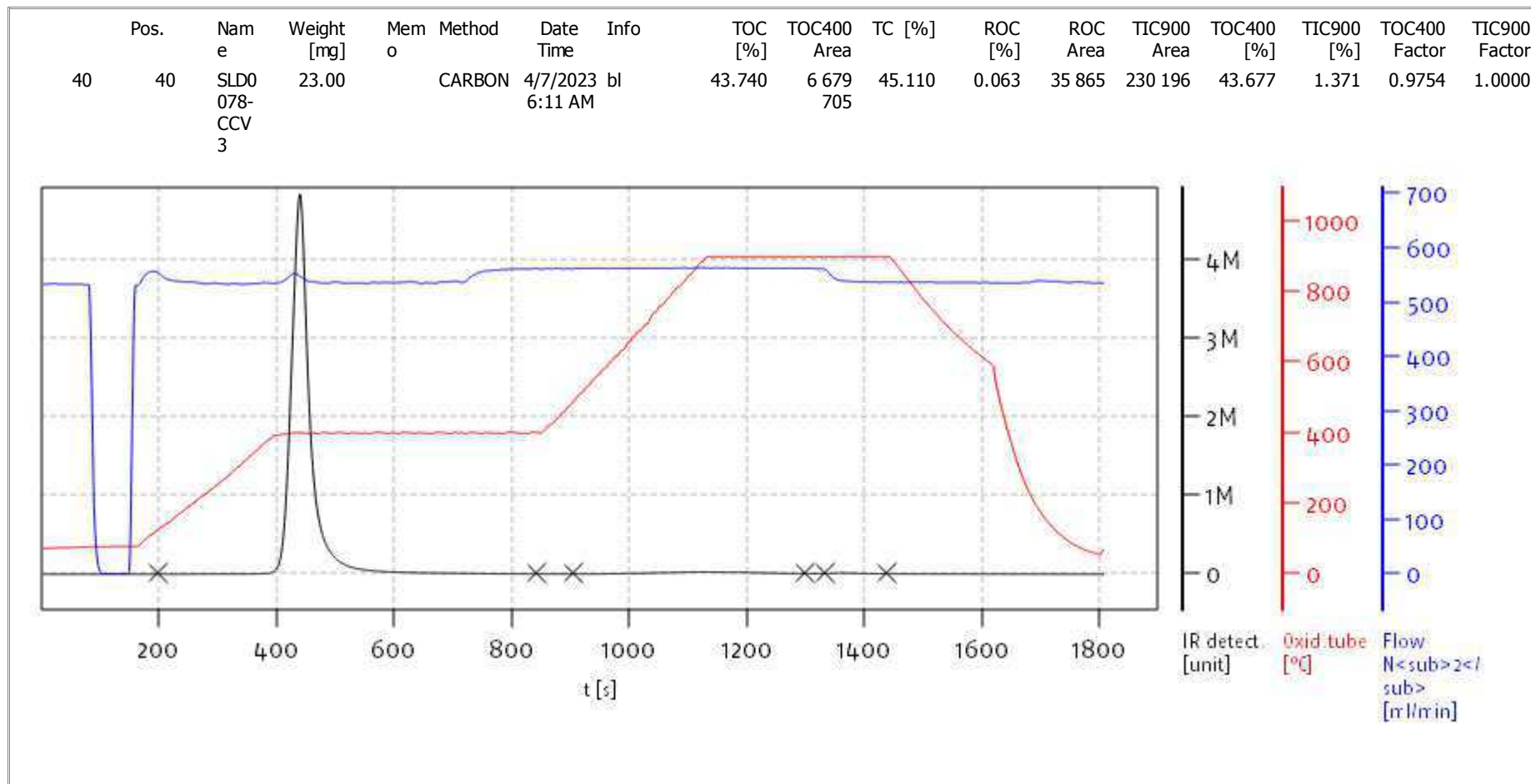
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solITOC V2.0.2 (31015f9) 2018-11-19
 Serial No: 0300.181017
 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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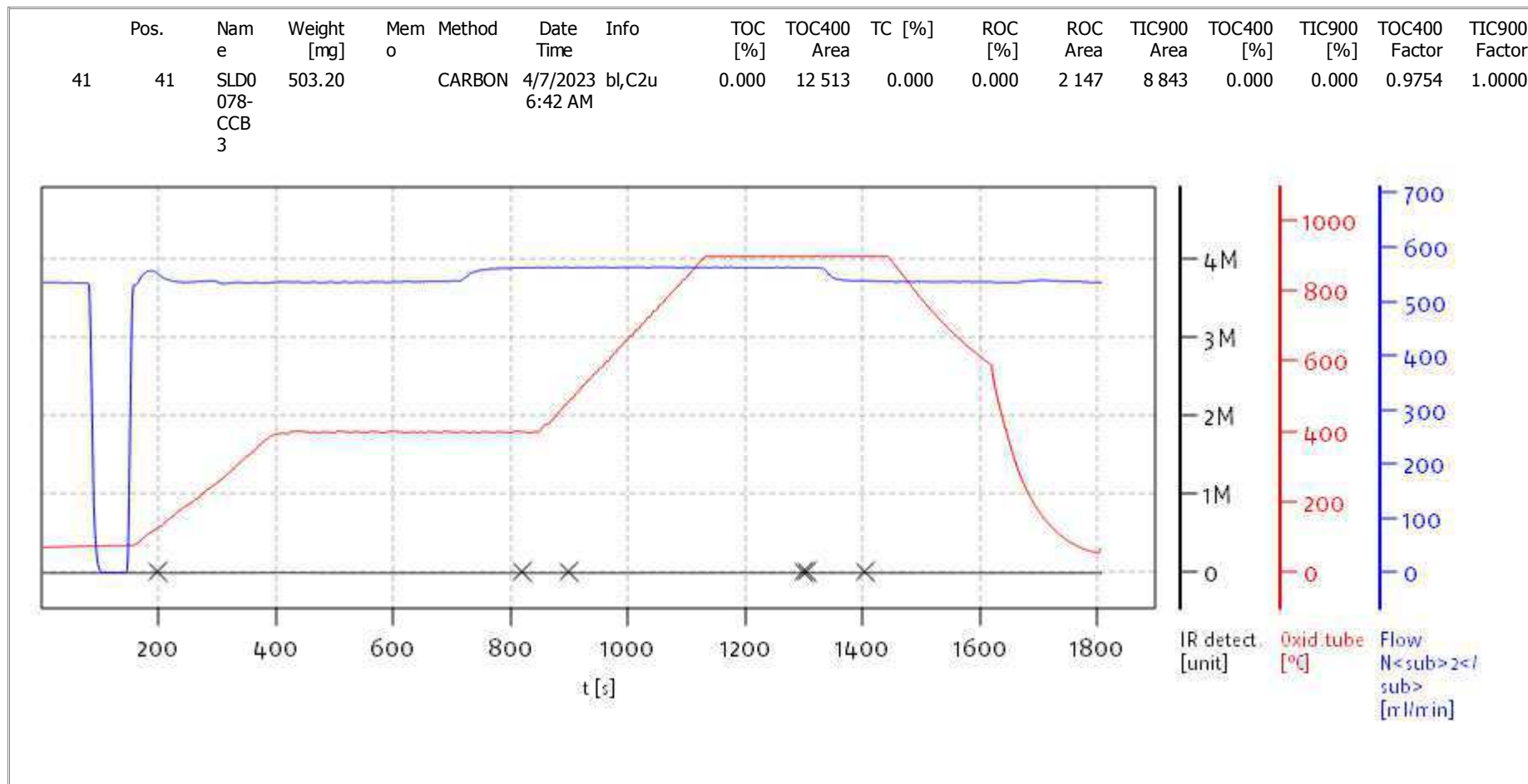
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Date: Fri Apr 7 14:09:03 2023



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 Serial No: 0300.181017
 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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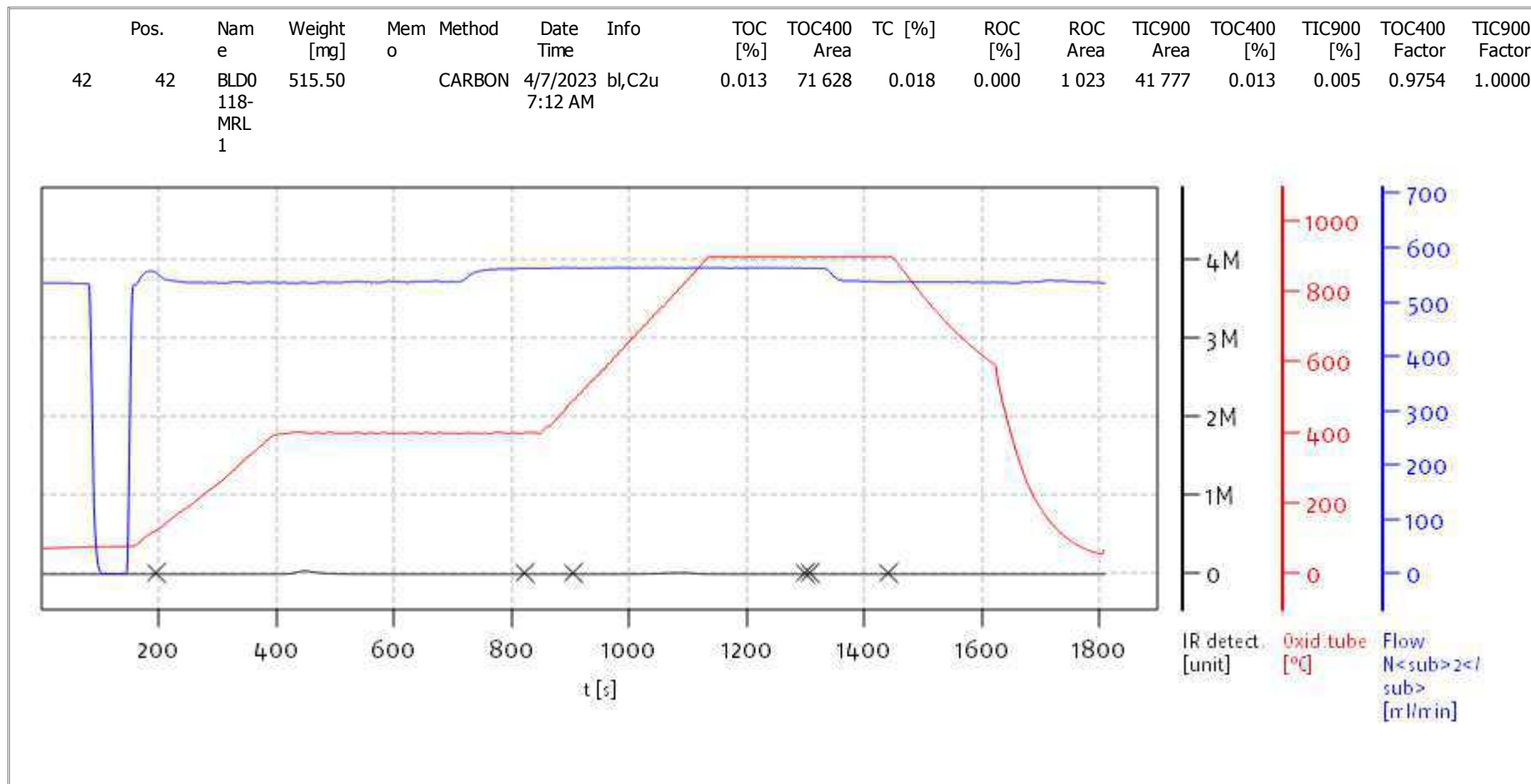
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 Mode CCC

Soli TOC Cube, Carbon
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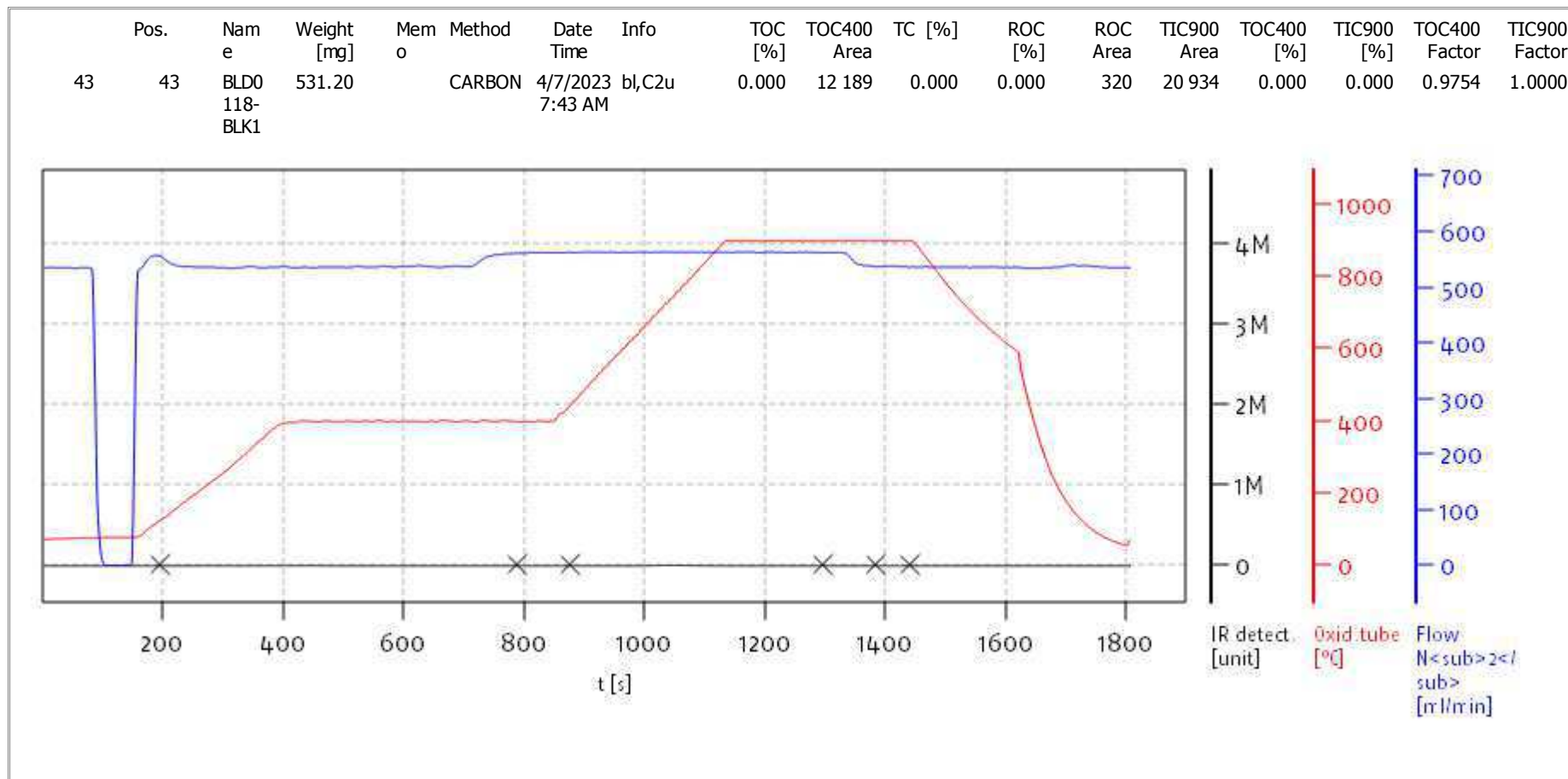
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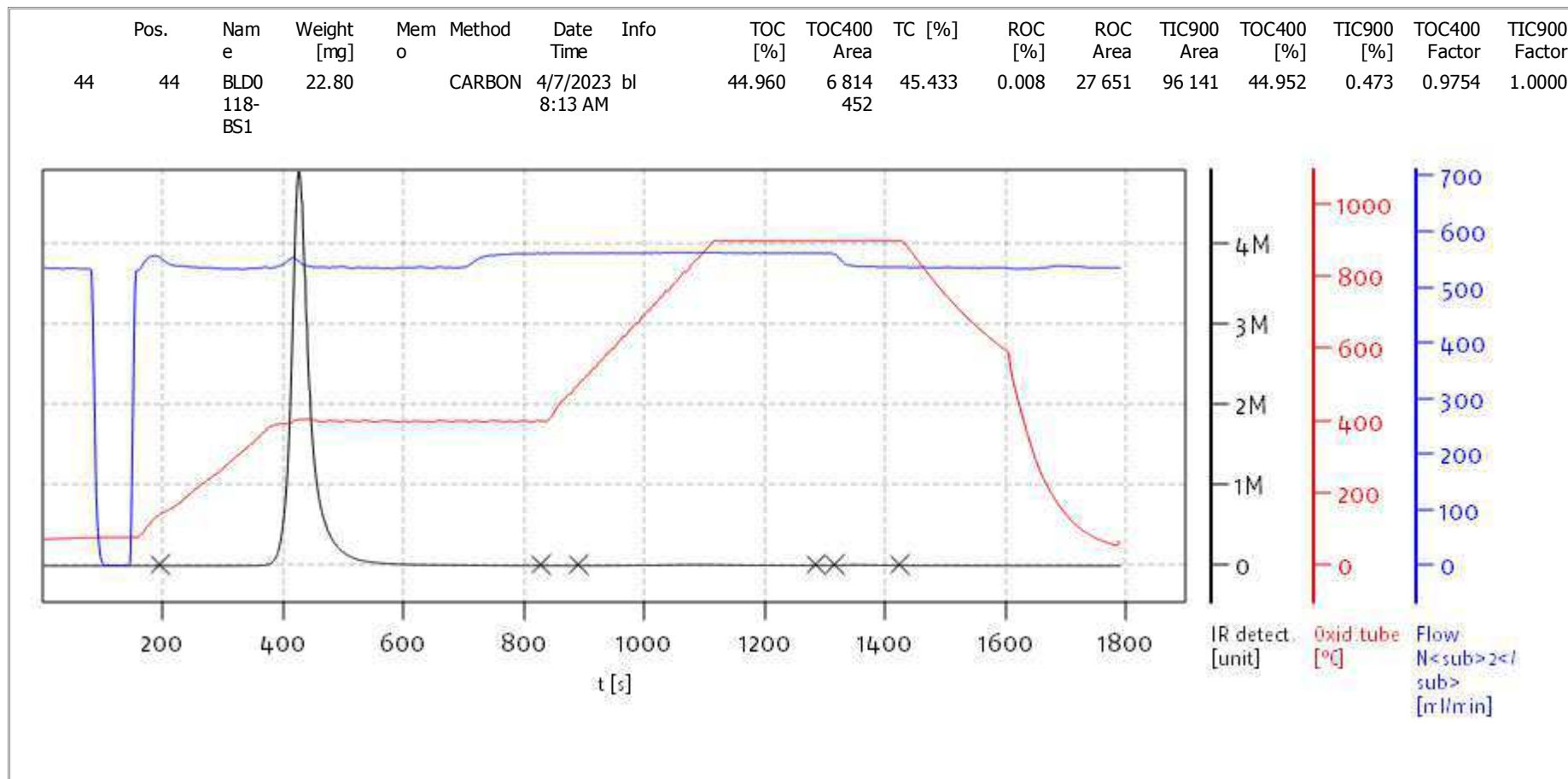
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solITOC V2.0.2 (31015f9) 2018-11-19
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Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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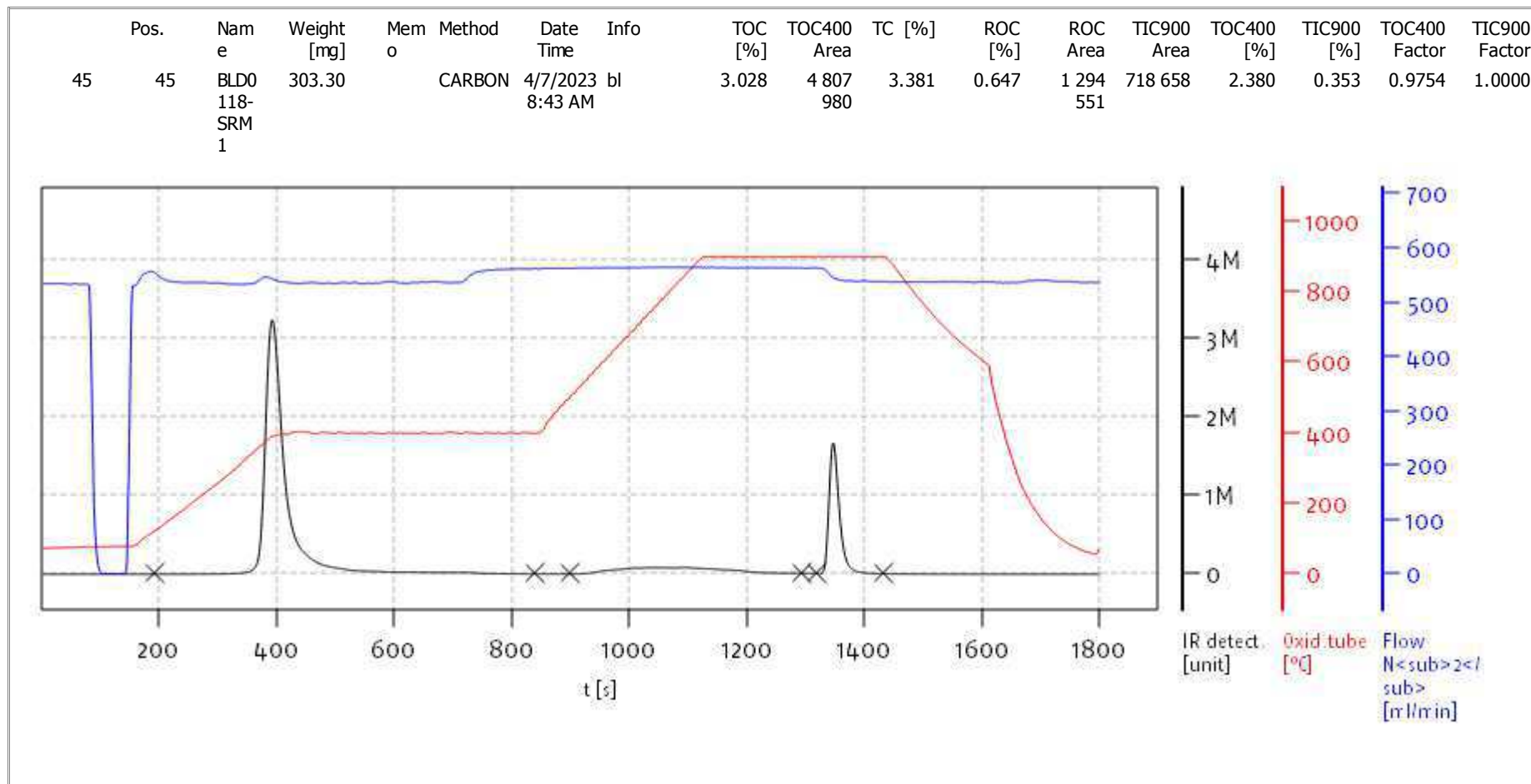
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 Mode CCC

Soli TOC Cube, Carbon
 Balance: BAL3
 Analyst: CDE



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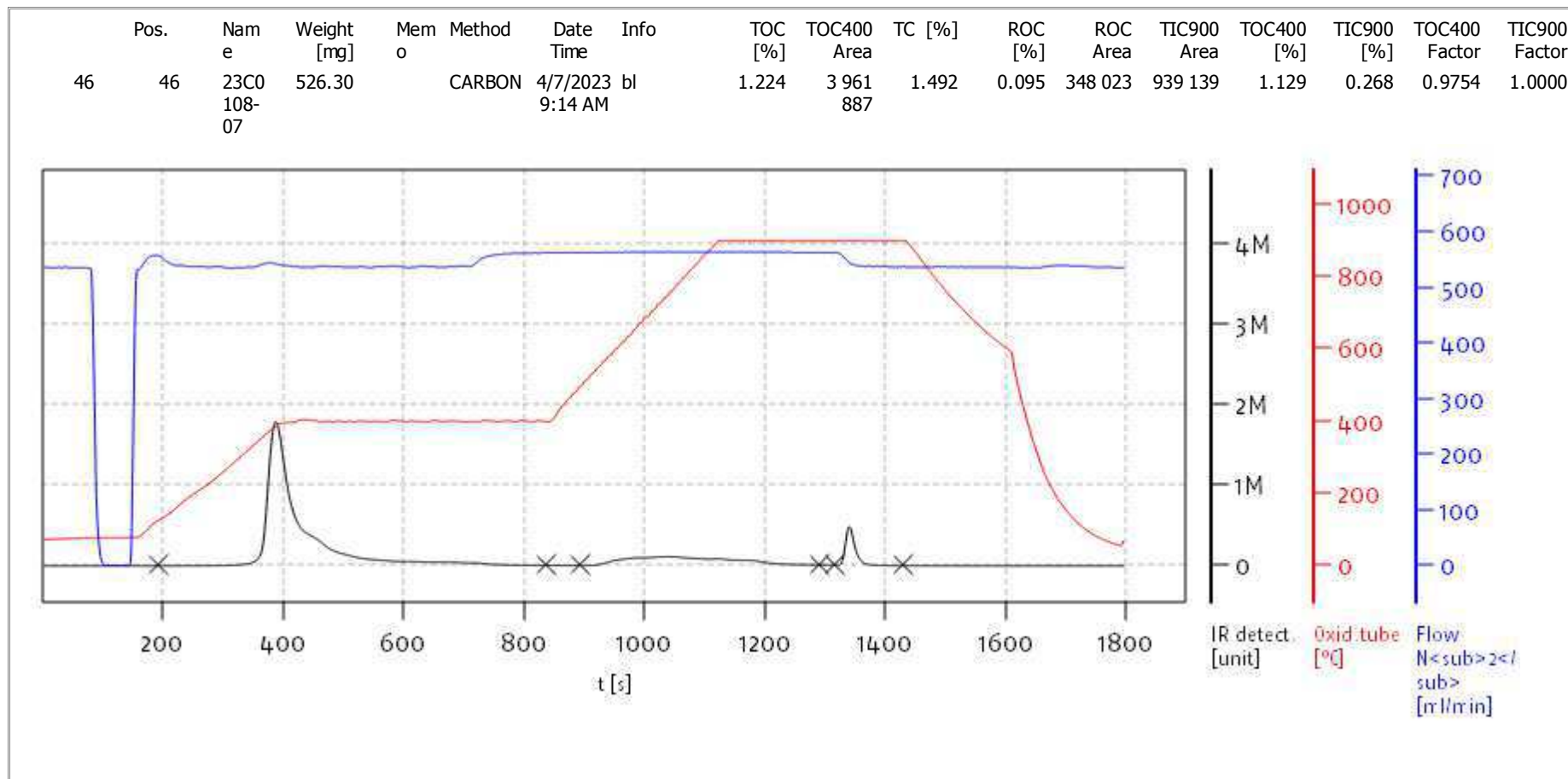
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 Balance: BAL3
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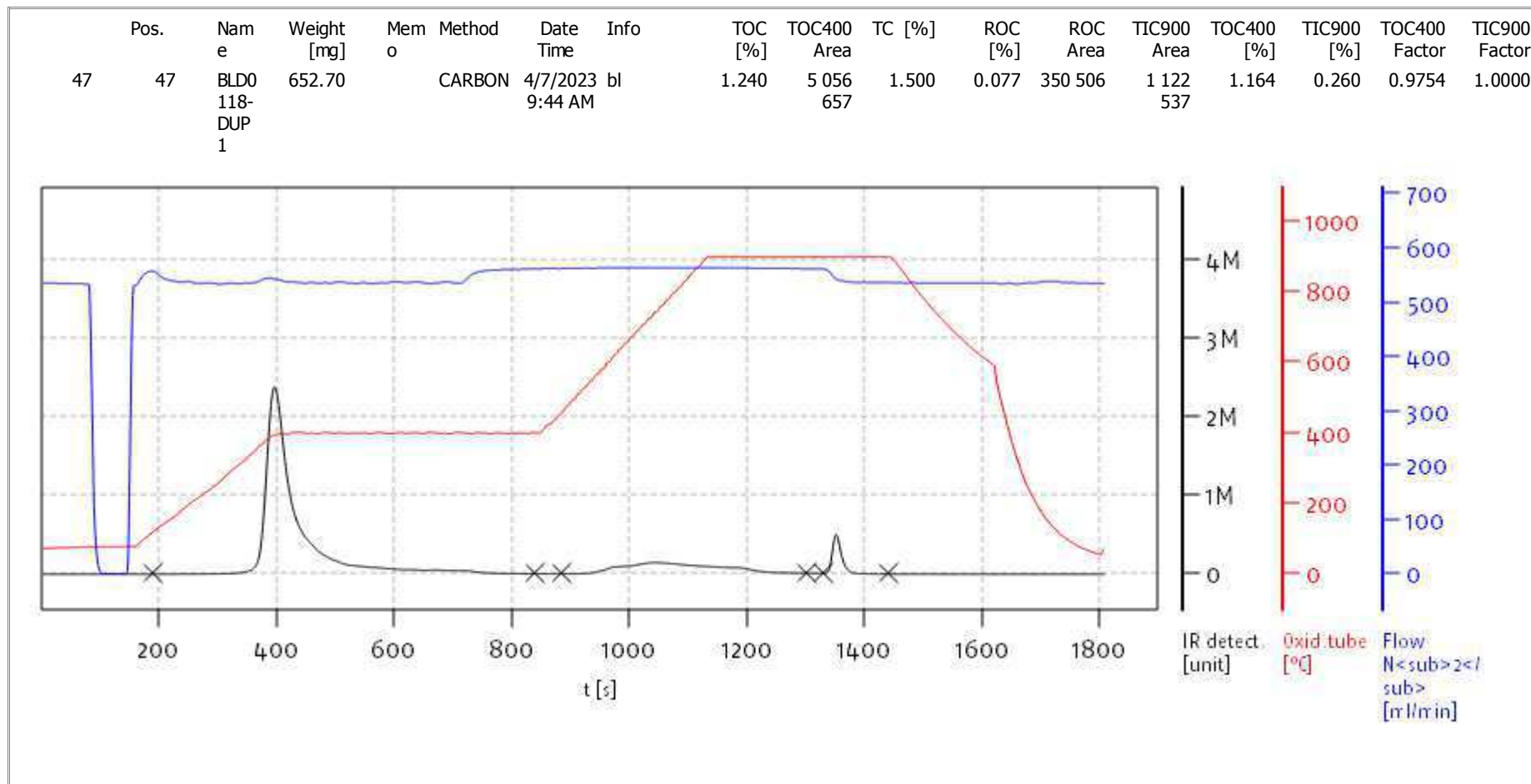
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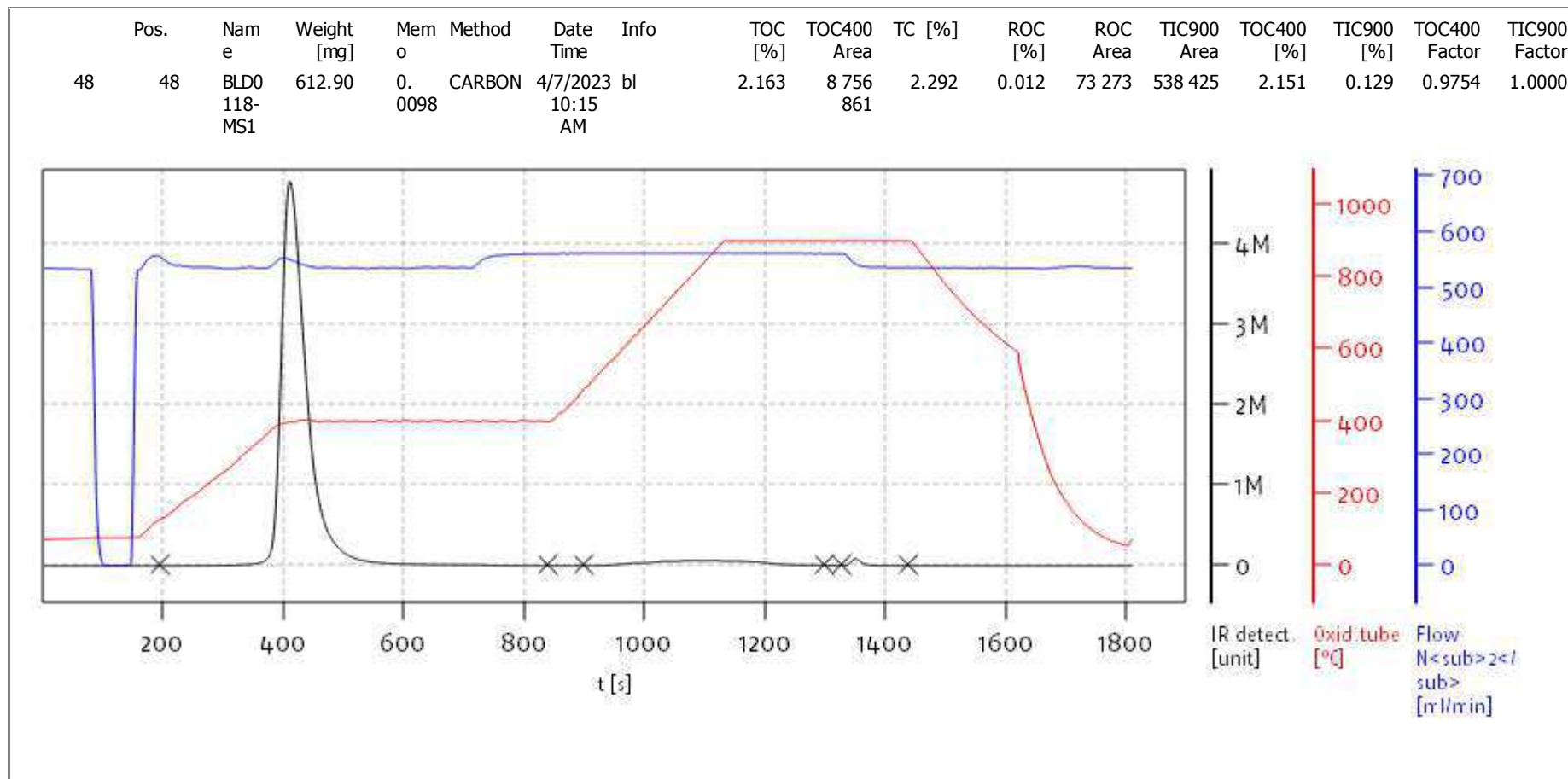
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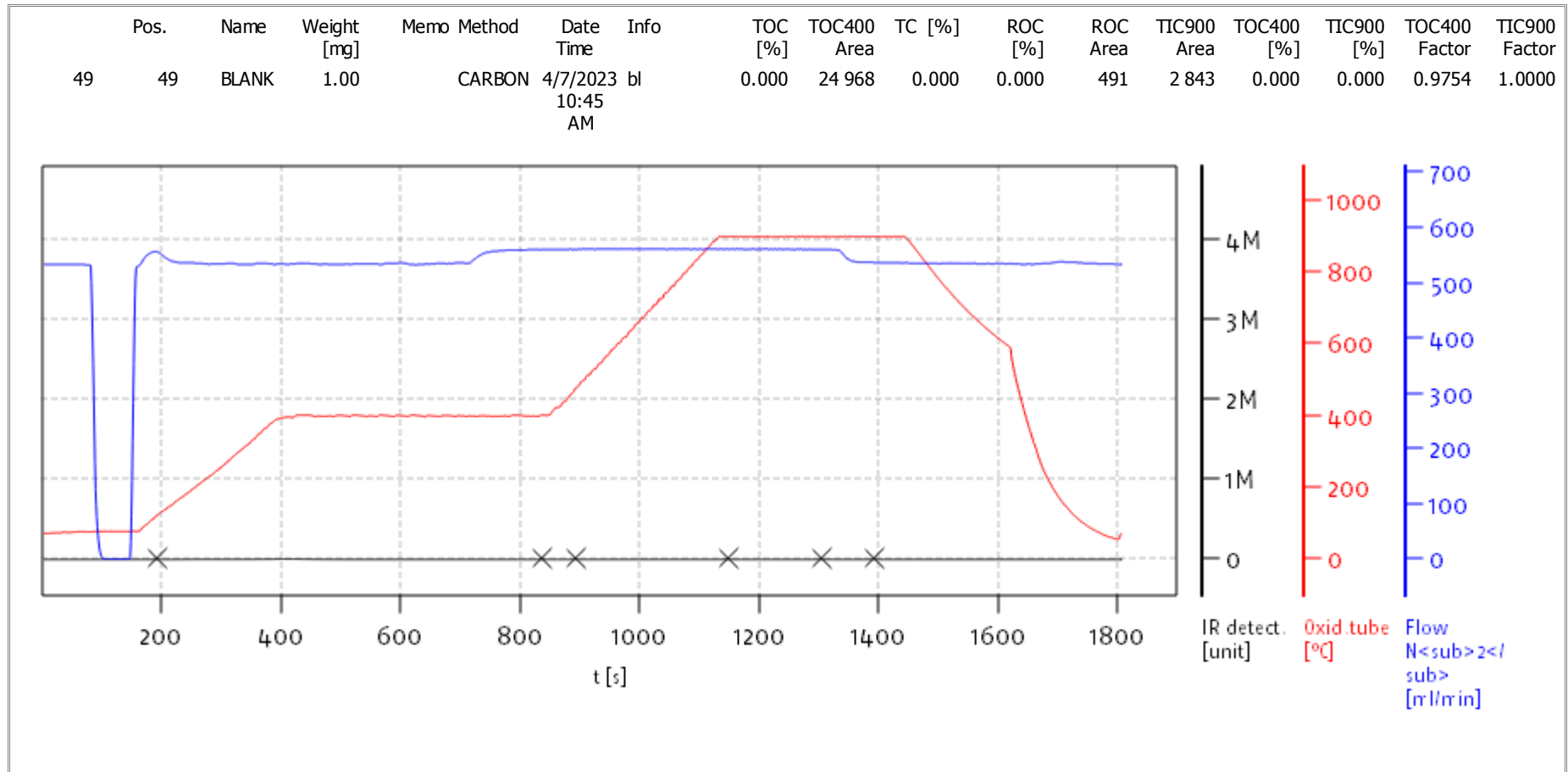
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Balance: BAL3
Analyst: CDE



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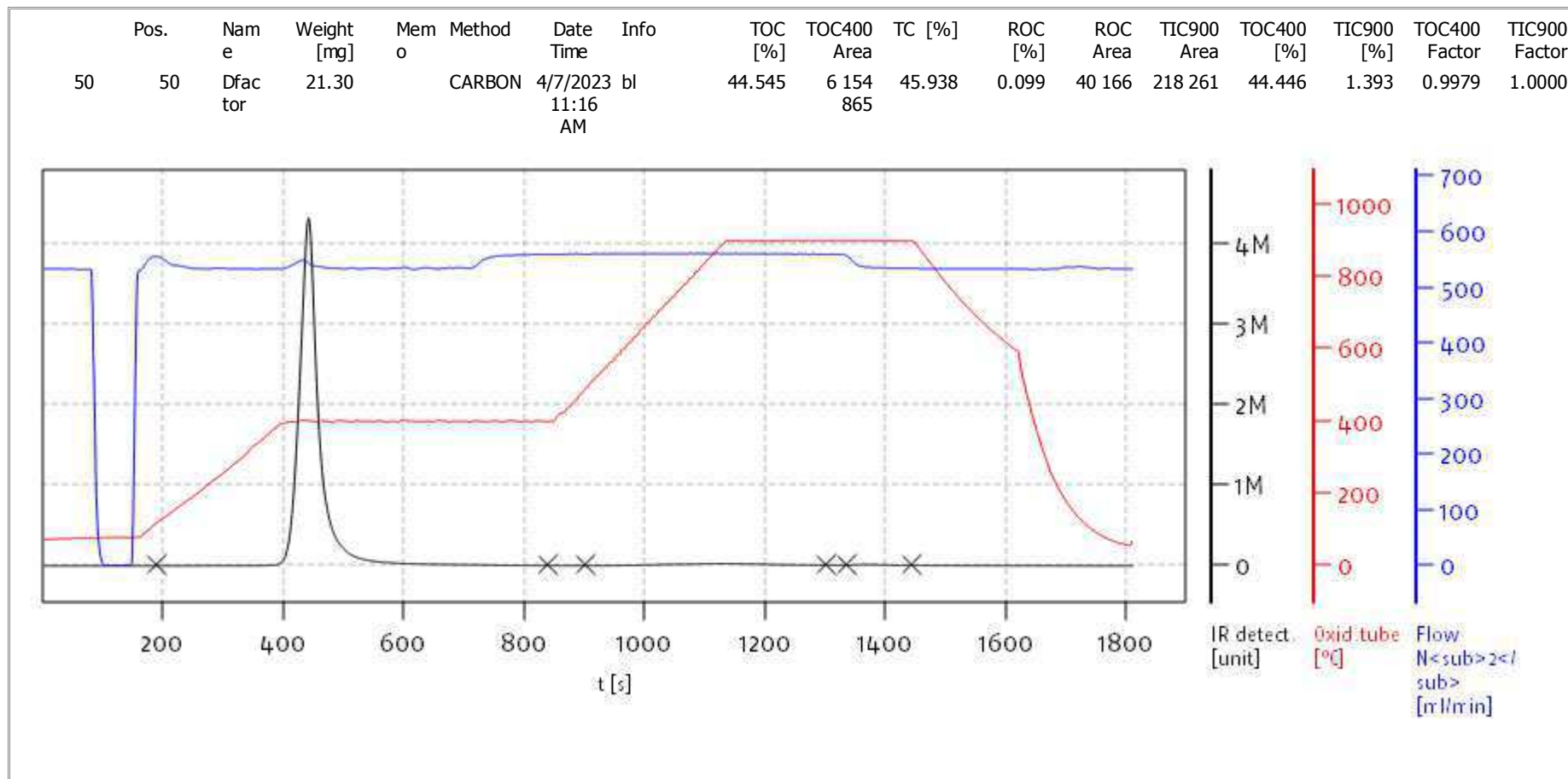
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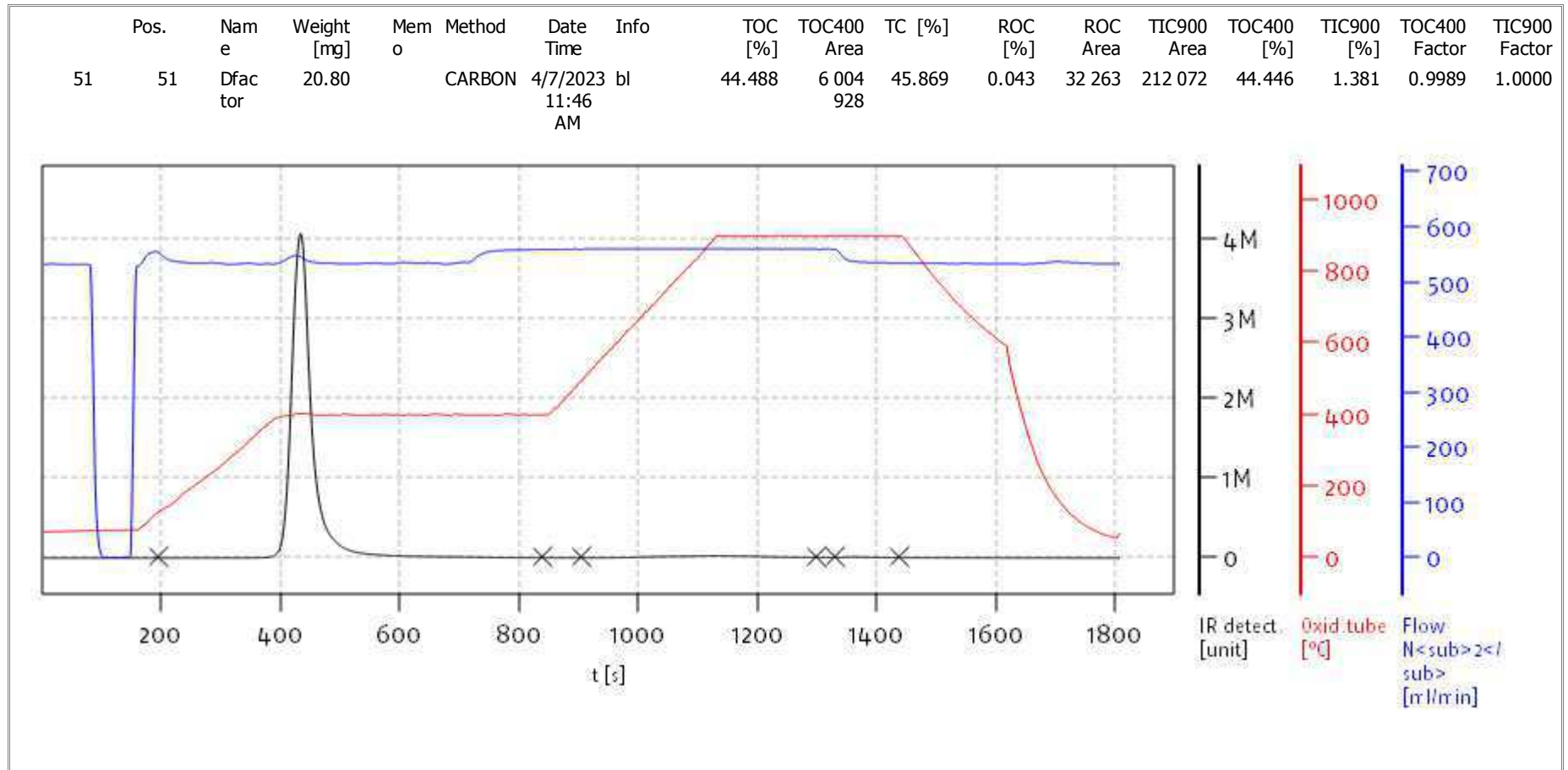
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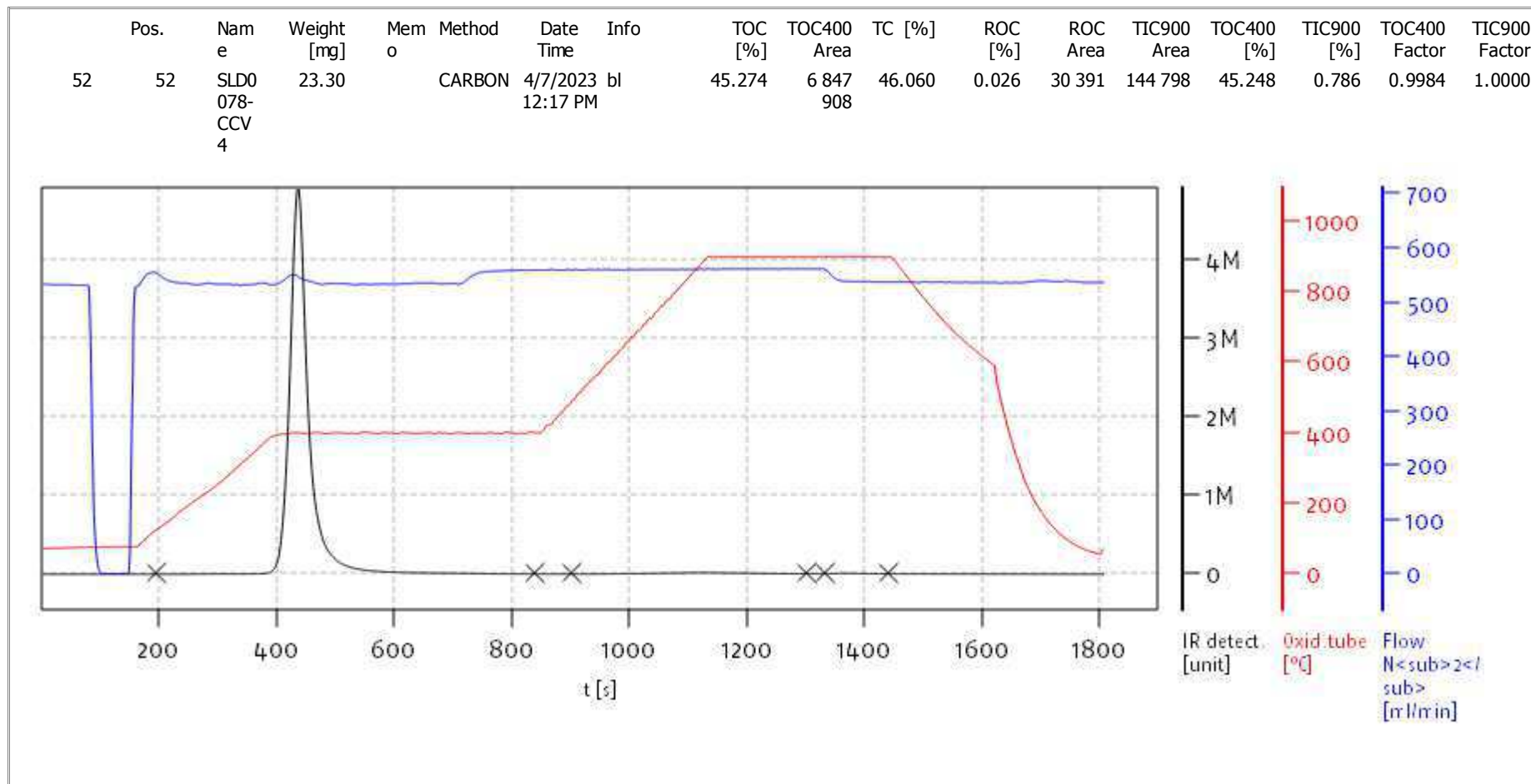
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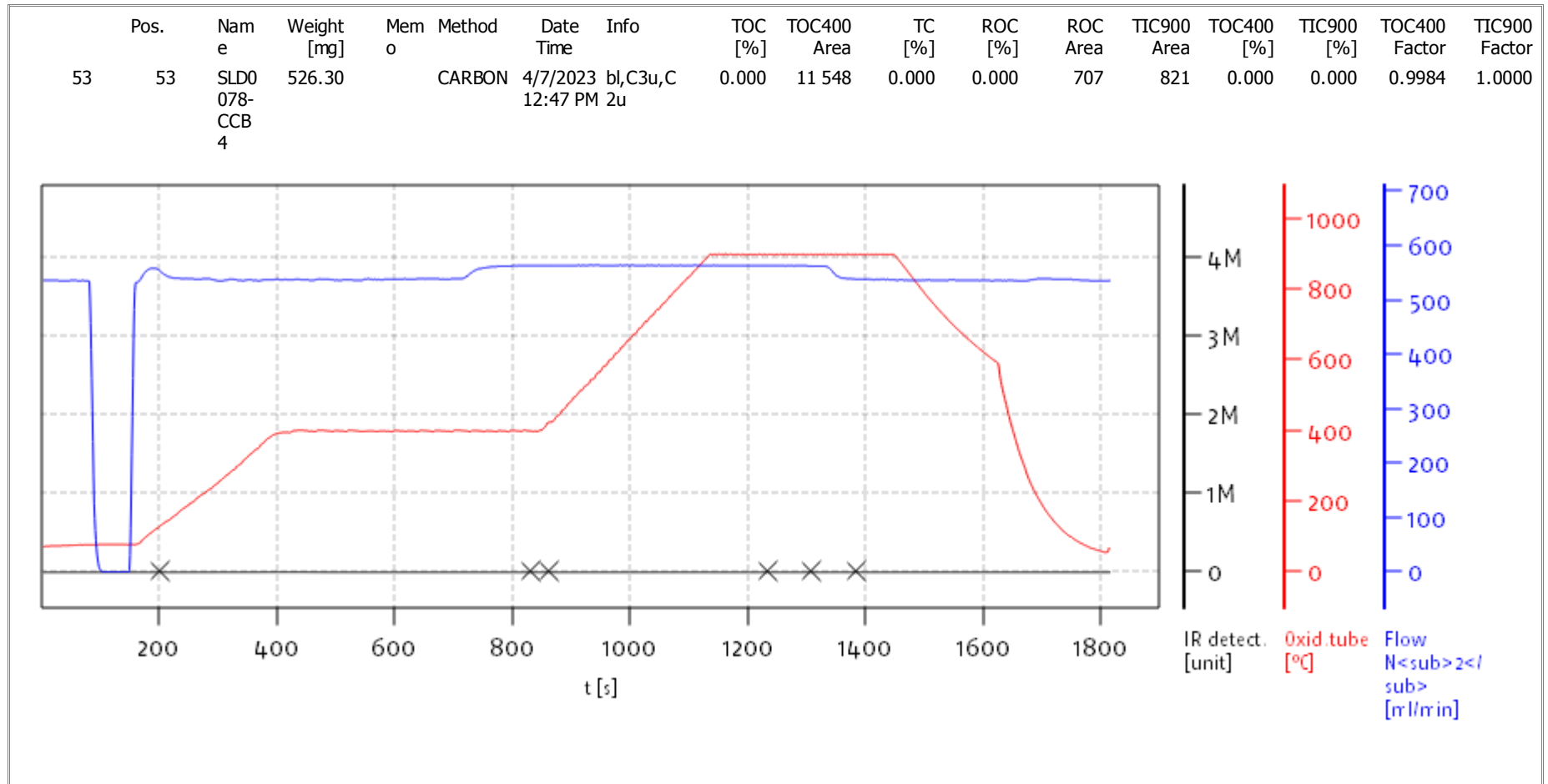
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Soli TOC Cube, Carbon
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Date: Fri Apr 7 14:09:03 2023



solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



INITIAL CALIBRATION DATA

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: FD00070

Instrument: TOC Cube

Calibration Date: 04/26/2022 11:29

Compound	Level 01		Level 02		Level 03		Level 04		Level 05		Level 06	
	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF
Total Organic Carbon	0.0080973	1449743	0.014695	1300238	0.021293	1292913	0.02939	1293535	0.044385	2094063	0.05878	1400085
Total Carbon	0.0080973	1449743	0.014695	1300238	0.021293	1292913	0.02939	1293535	0.044385	2094063	0.05878	1400085
Total Inorganic Carbon	0.0080973	1449743	0.014695	1300238	0.021293	1292913	0.02939	1293535	0.044385	2094063	0.05878	1400085
% Soot	0.0080973	1449743	0.014695	1300238	0.021293	1292913	0.02939	1293535	0.044385	2094063	0.05878	1400085



INITIAL CALIBRATION DATA

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: FD00070

Instrument: TOC Cube

Calibration Date: 04/26/2022 11:29

Compound	Level 07		Level 08		Level 09		Level 10		Level 11		Level 12	
	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF
Total Organic Carbon	0.074075	1370638	0.08937	1351930	0.12056	2158544	0.14995	1559046	0.24	1346463	0.288	1430135
Total Carbon	0.074075	1370638	0.08937	1351930	0.12056	2158544	0.14995	1559046	0.24	1346463	0.288	1430135
Total Inorganic Carbon	0.074075	1370638	0.08937	1351930	0.12056	2158544	0.14995	1559046	0.24	1346463	0.288	1430135
% Soot	0.074075	1370638	0.08937	1351930	0.12056	2158544	0.14995	1559046	0.24	1346463	0.288	1430135



INITIAL CALIBRATION DATA

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: FD00070

Instrument: TOC Cube

Calibration Date: 04/26/2022 11:29

Compound	Level 13		Level 14		Level 15		Level 16		Level 17		Level 18	
	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF
Total Organic Carbon	0.414	1337053	0.606	1385937	0.894	1382774	1.188	1379790	1.5	1375927	1.818	1372882
Total Carbon	0.414	1337053	0.606	1385937	0.894	1382774	1.188	1379790	1.5	1375927	1.818	1372882
Total Inorganic Carbon	0.414	1337053	0.606	1385937	0.894	1382774	1.188	1379790	1.5	1375927	1.818	1372882
% Soot	0.414	1337053	0.606	1385937	0.894	1382774	1.188	1379790	1.5	1375927	1.818	1372882



INITIAL CALIBRATION DATA

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Calibration: FD00070

Instrument: TOC Cube

Calibration Date: 04/26/2022 11:29

Compound	Level 19		Level 20		Level 21		Level 22		Level 23		Level 24	
	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF	Conc	RF
Total Organic Carbon	2.49	1398606	2.982	1376871	4.188	1256057	4.818	1279542	5.406	1283358	7.2	1301408
Total Carbon	2.49	1398606	2.982	1376871	4.188	1256057	4.818	1279542	5.406	1283358	7.2	1301408
Total Inorganic Carbon	2.49	1398606	2.982	1376871	4.188	1256057	4.818	1279542	5.406	1283358	7.2	1301408
% Soot	2.49	1398606	2.982	1376871	4.188	1256057	4.818	1279542	5.406	1283358	7.2	1301408



INITIAL CALIBRATION DATA

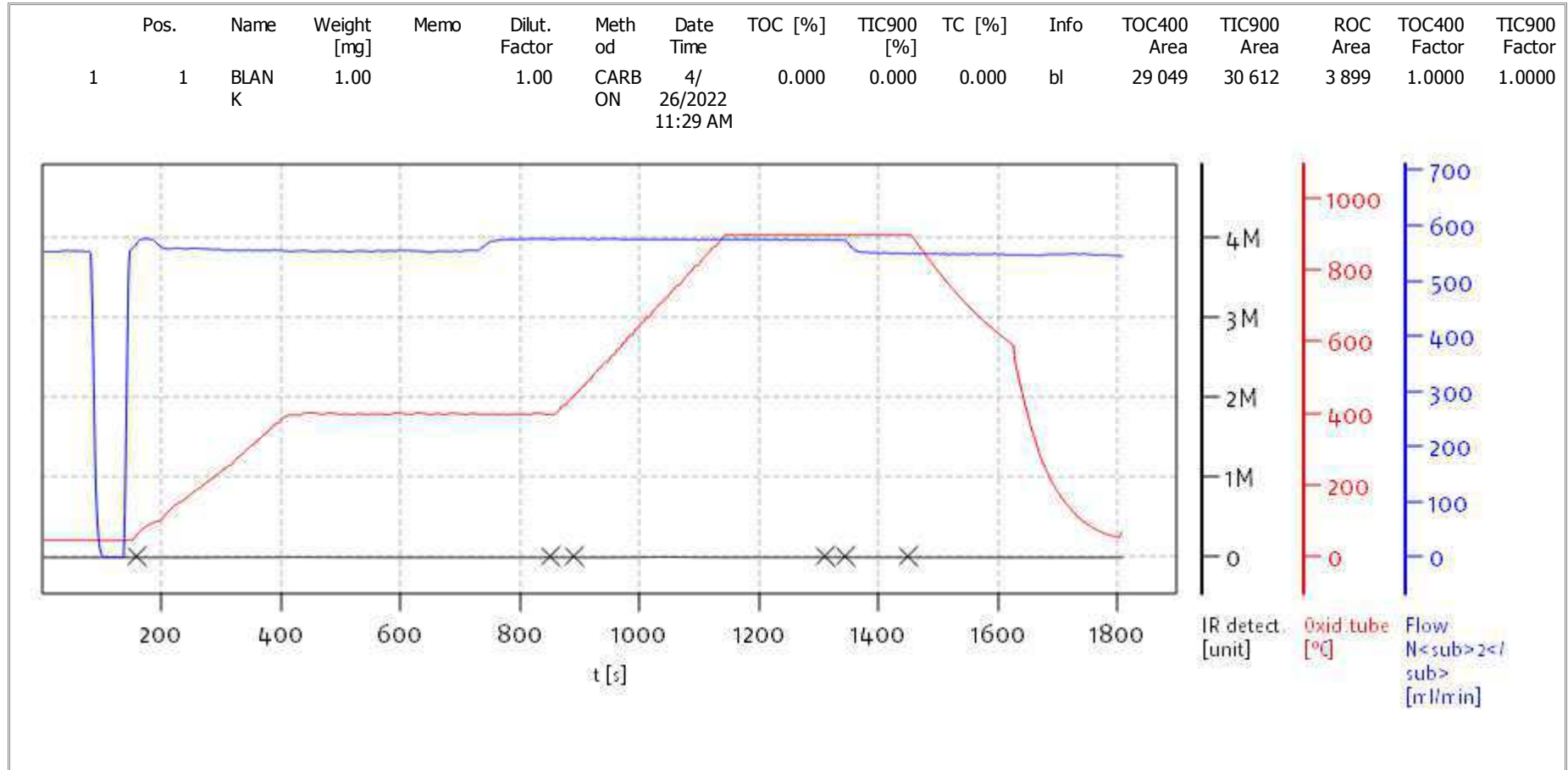
EPA 9060A m

Laboratory:	Analytical Resources, LLC	SDG:	23C0107
Client:	Anchor QEA, LLC	Project:	AOC4 UR Phase 3
Calibration:	FD00070	Instrument:	TOC Cube
Calibration Date:	04/26/2022 11:29		

COMPOUND	Mean RF	RF RSD	Linear COD	Quad COD	COD Limit	Q
Total Organic Carbon	1424064	15.9	0.9988			
Total Carbon	1424064	15.9	0.9988			
Total Inorganic Carbon	1424064	15.9	0.9988			
% Soot	1424064	15.9	0.9988			



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

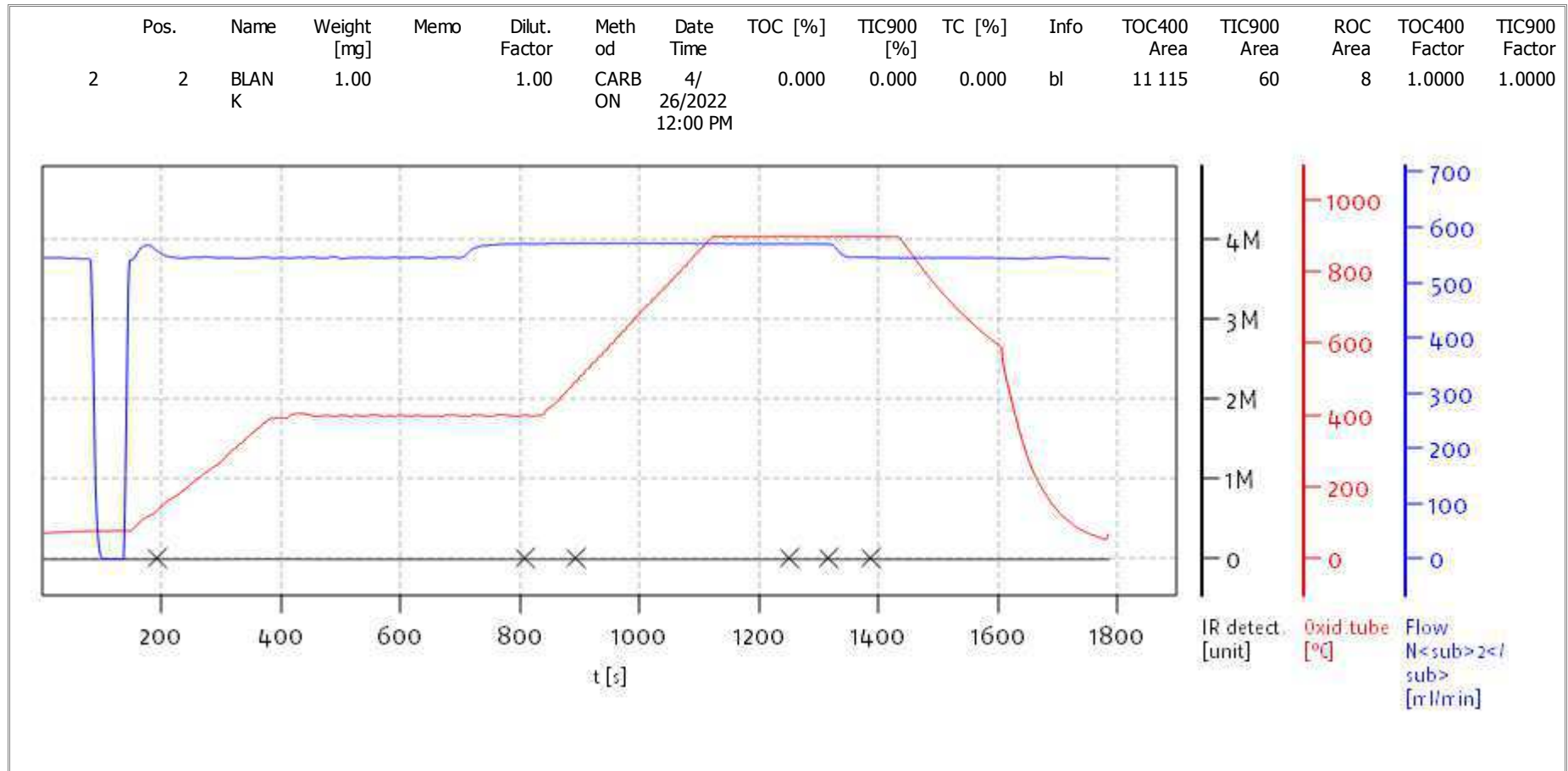
Date: Wed Apr 27 11:07:12 2022



solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

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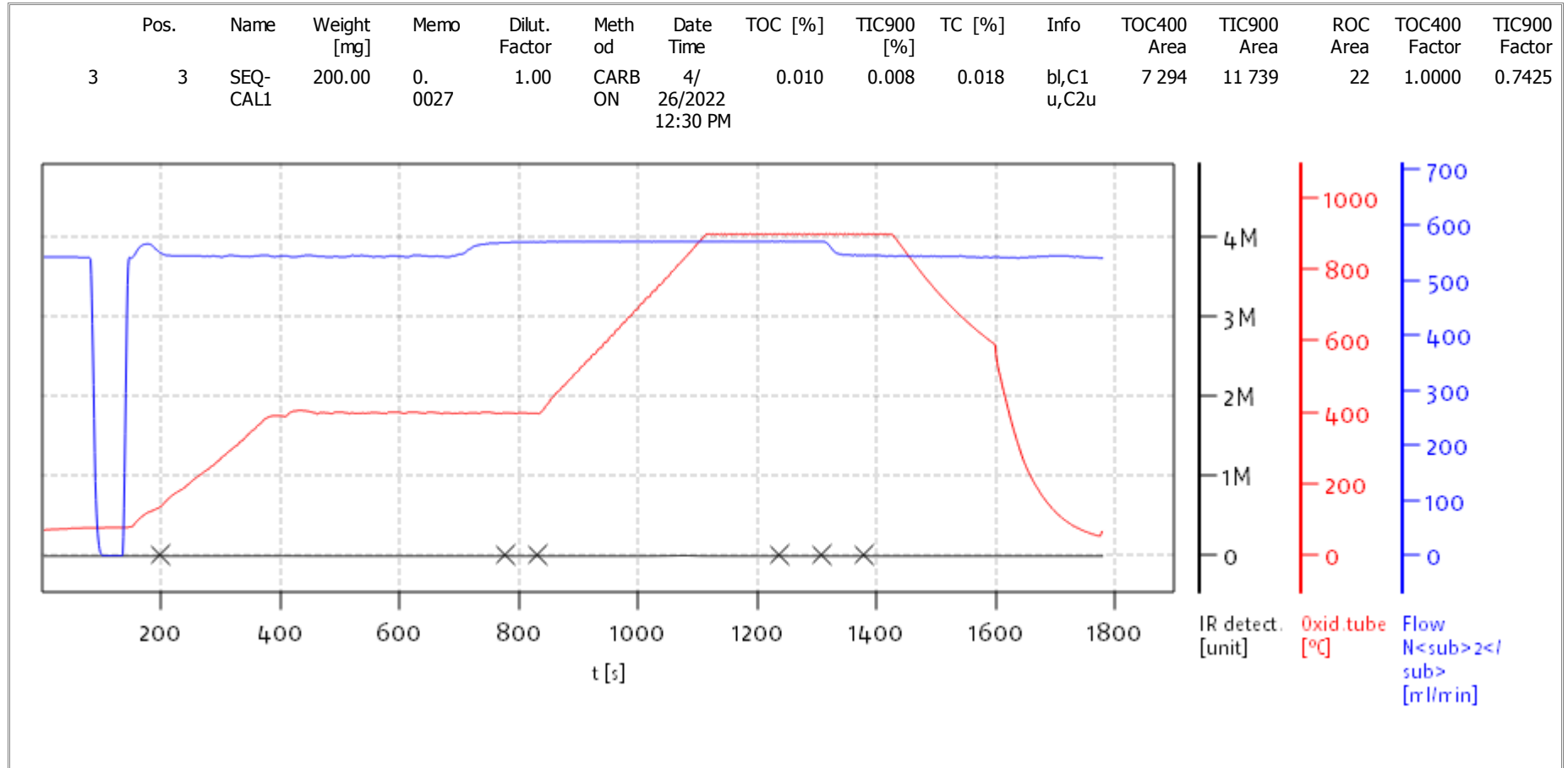
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Soli TOC Cube, Carbon
Balance: BAL3
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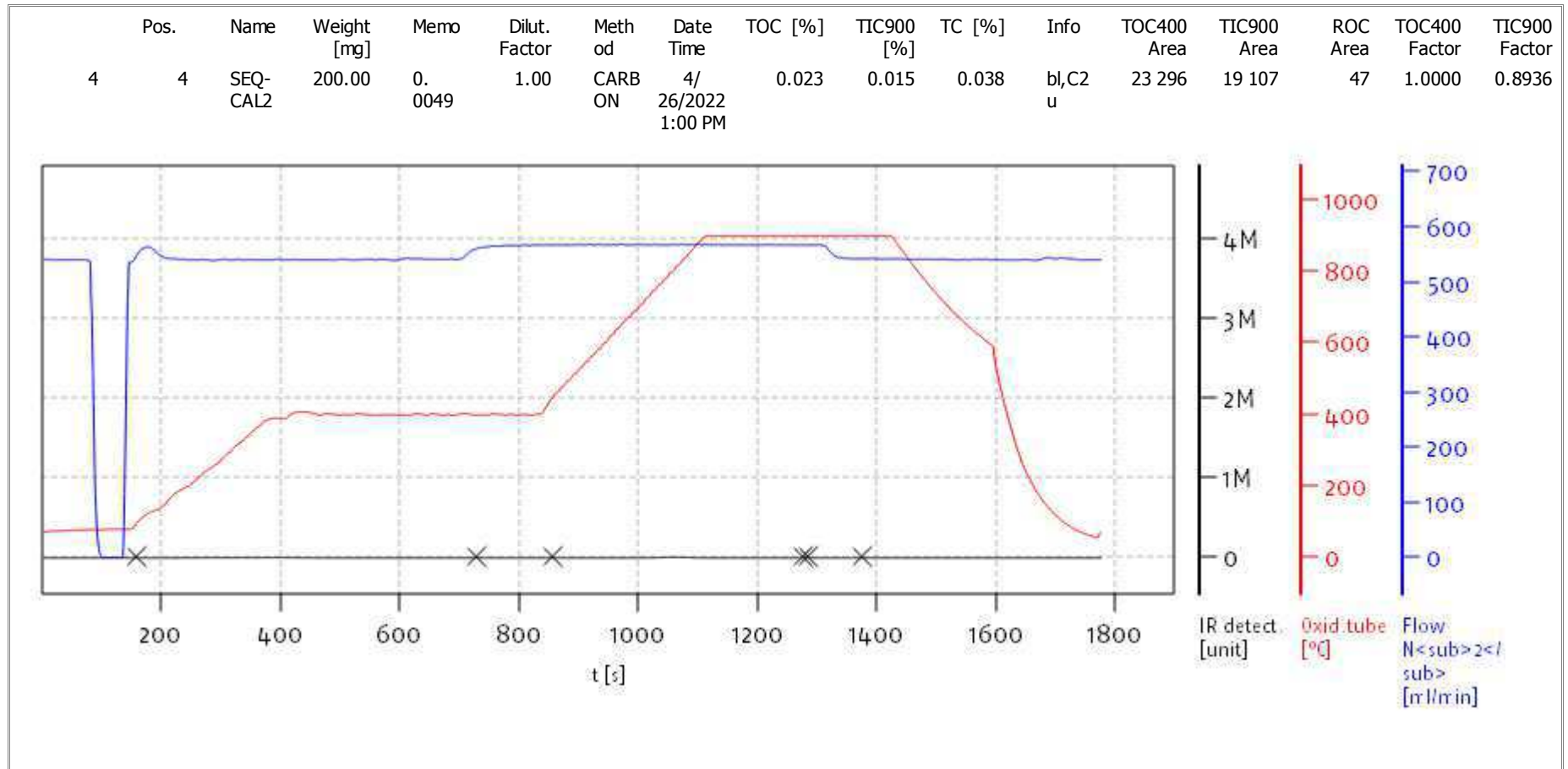
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Soli TOC Cube, Carbon
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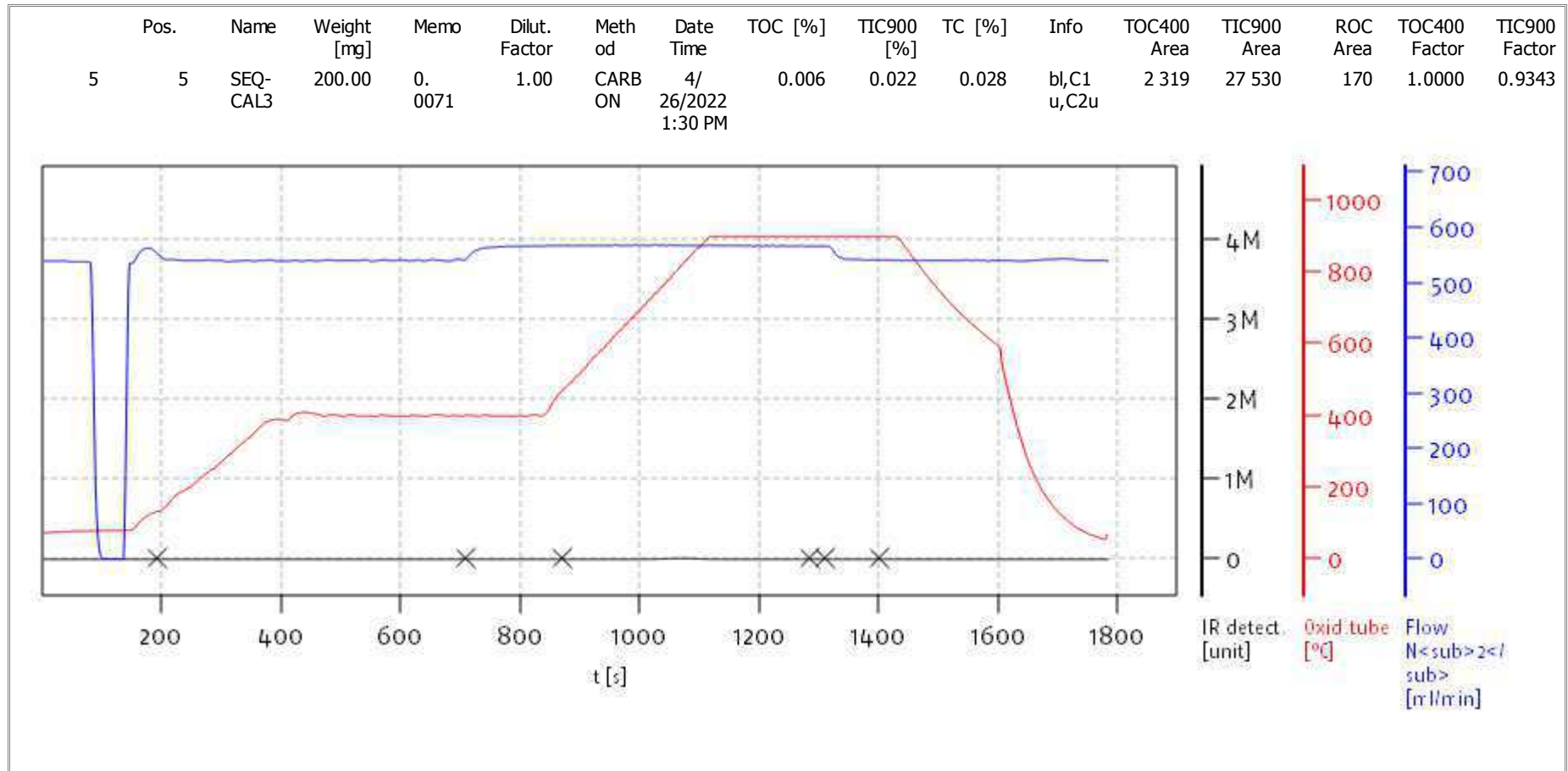
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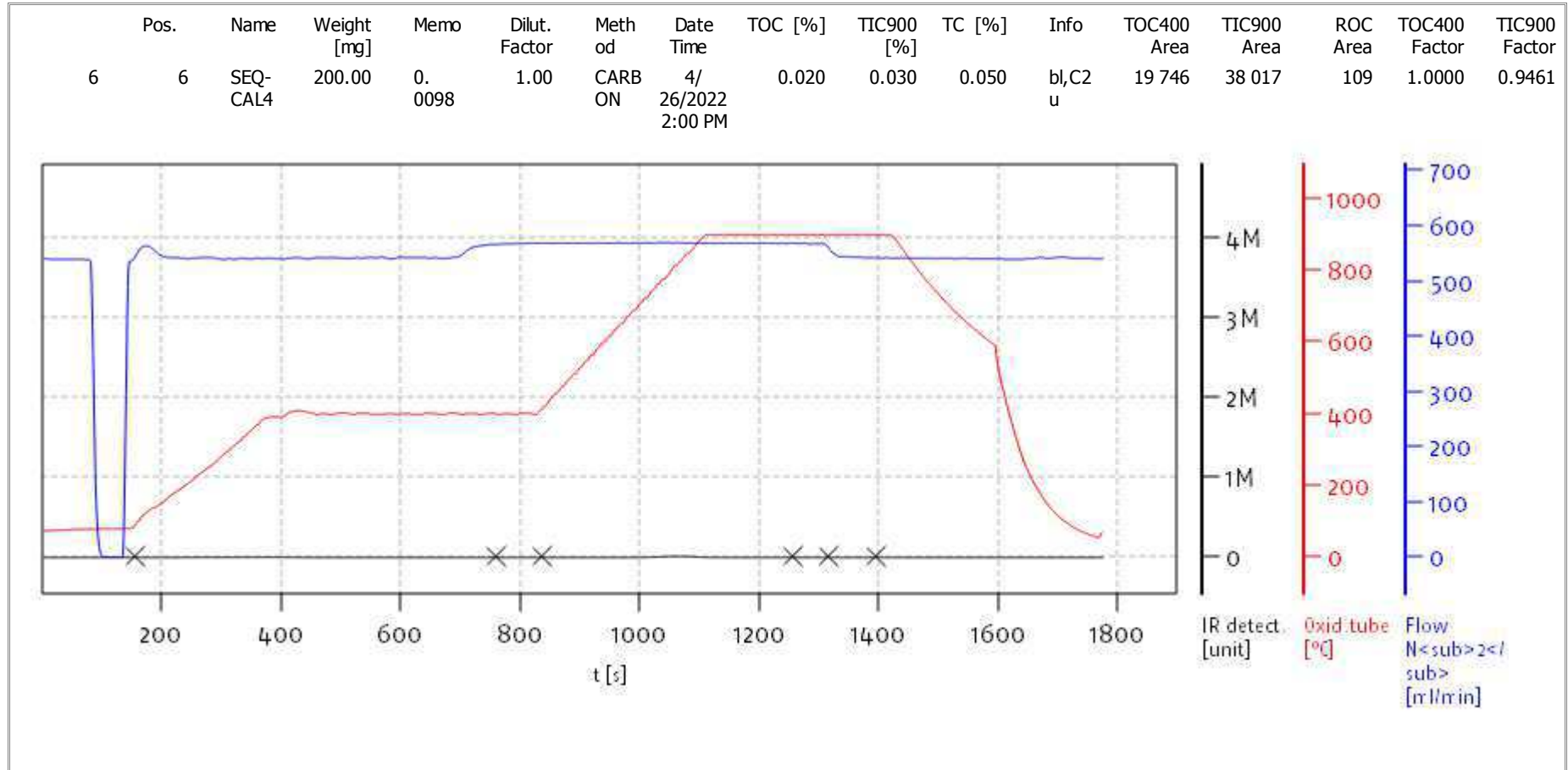
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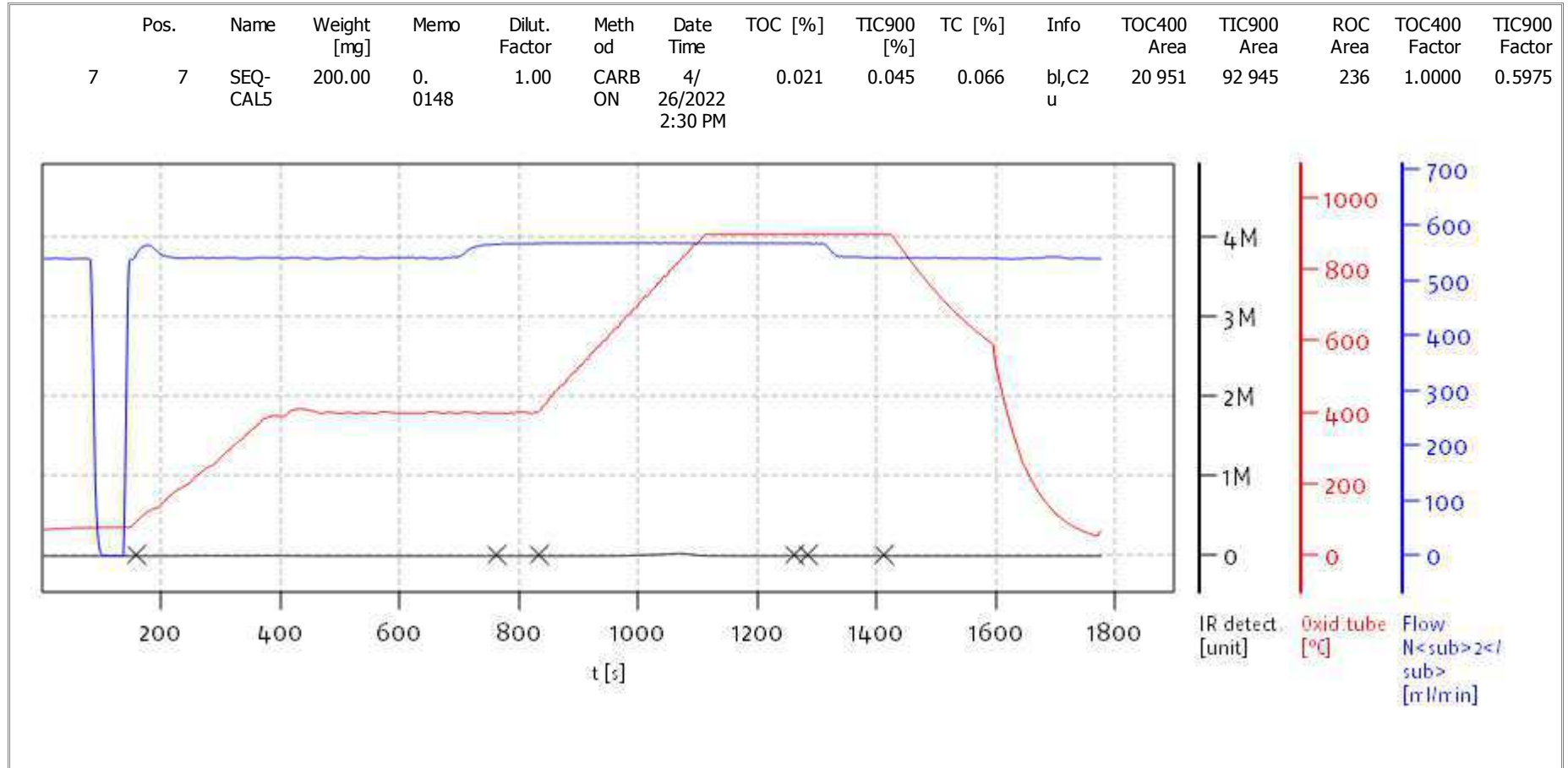
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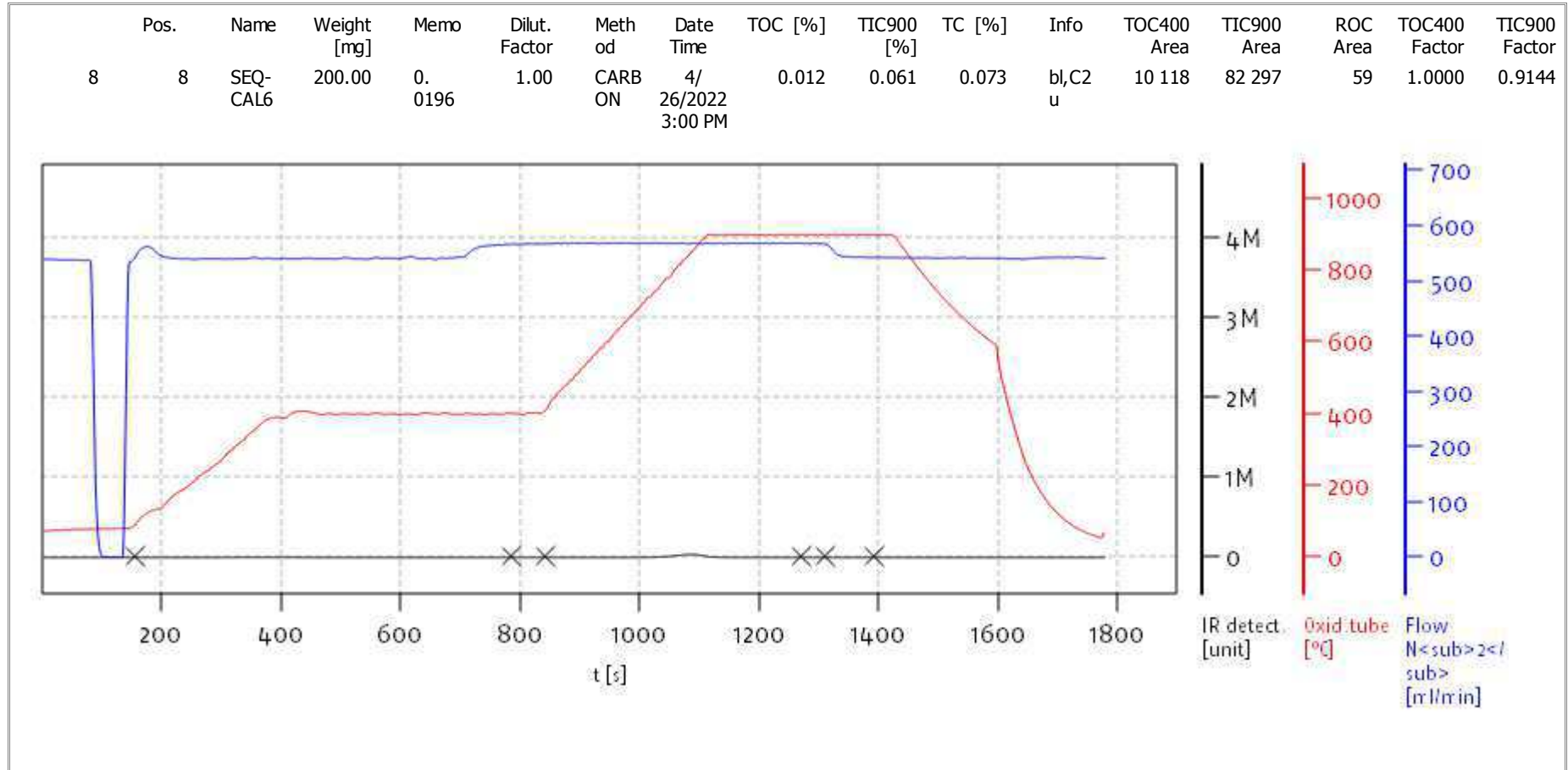
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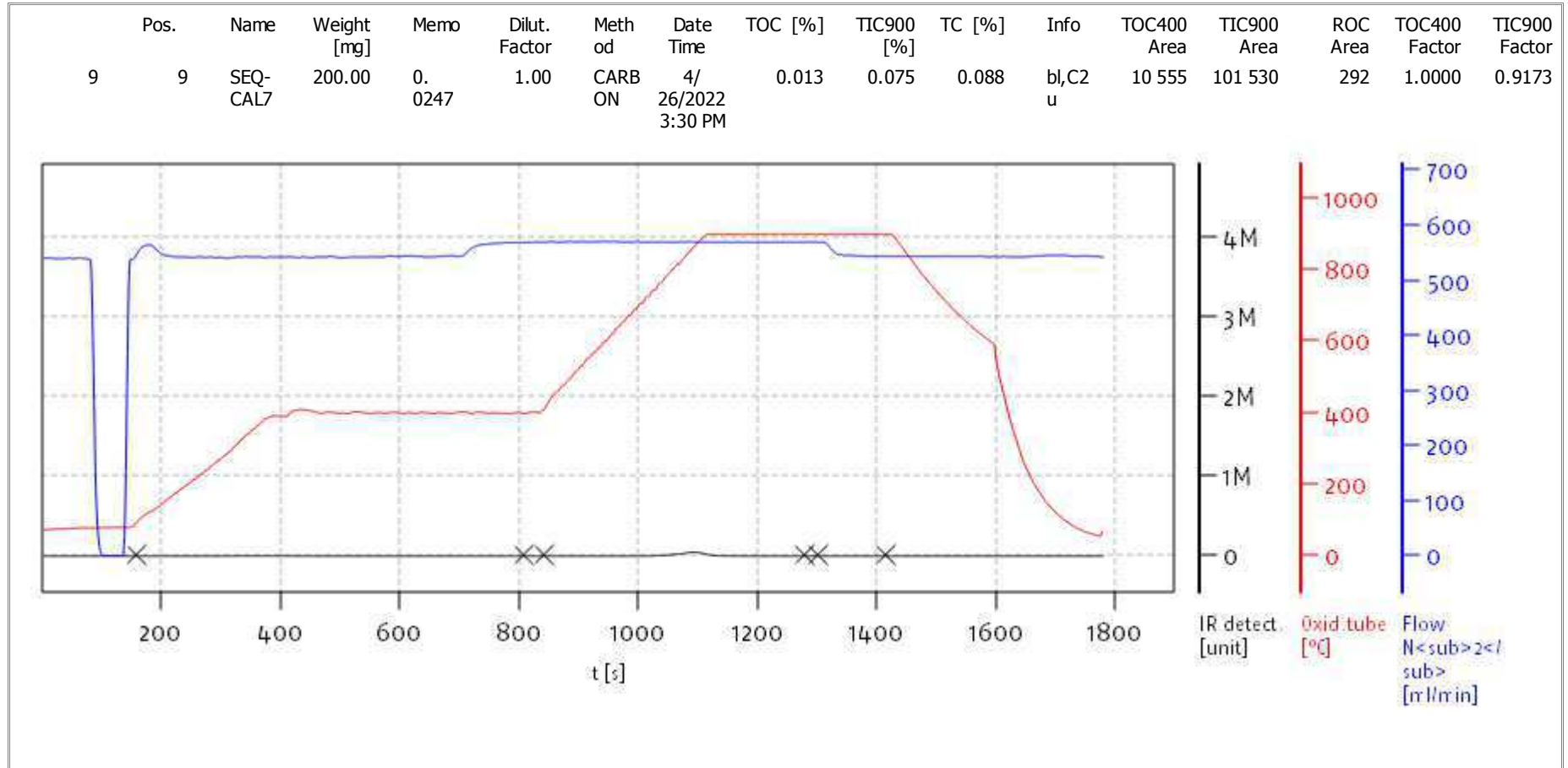
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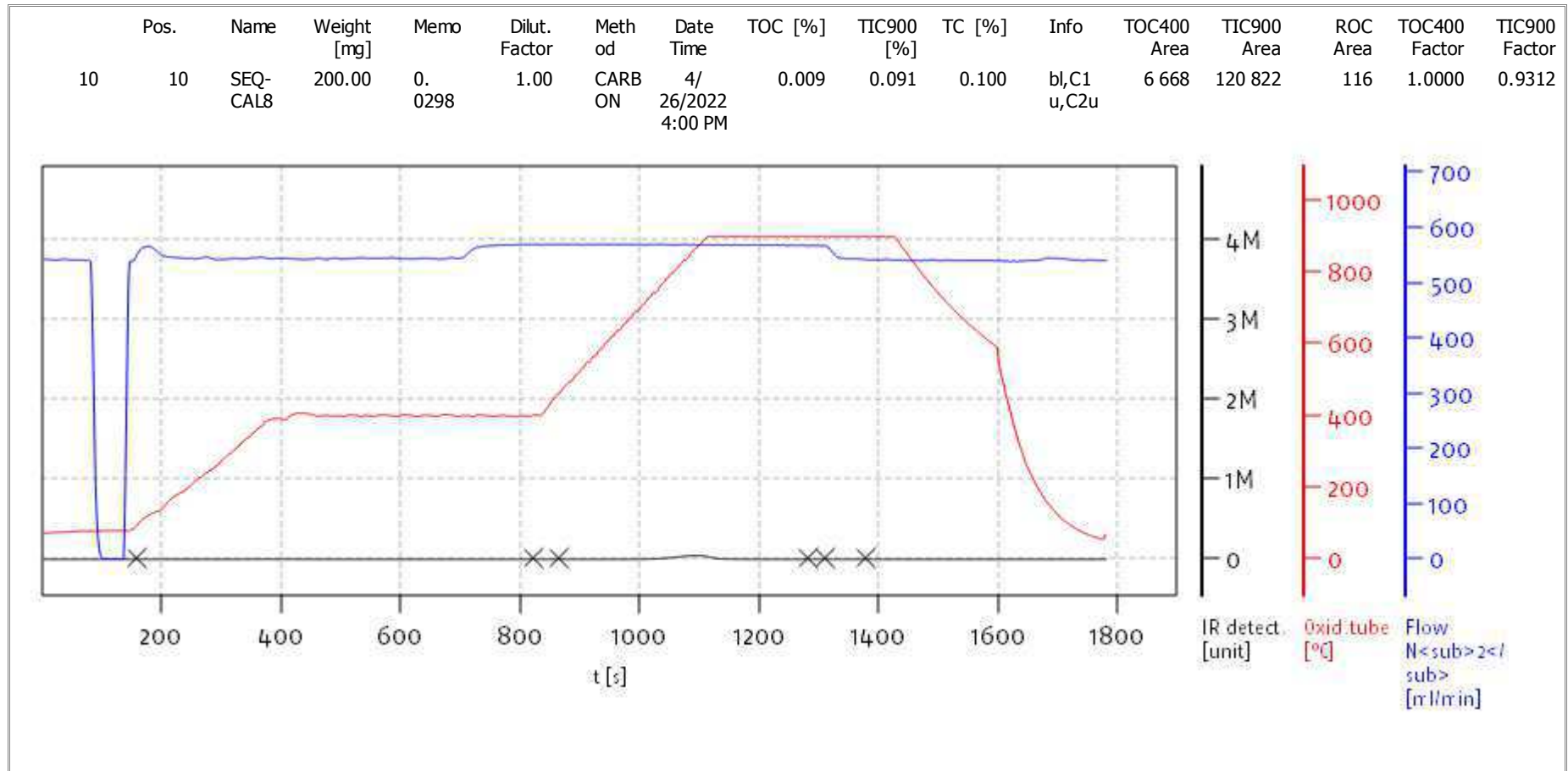
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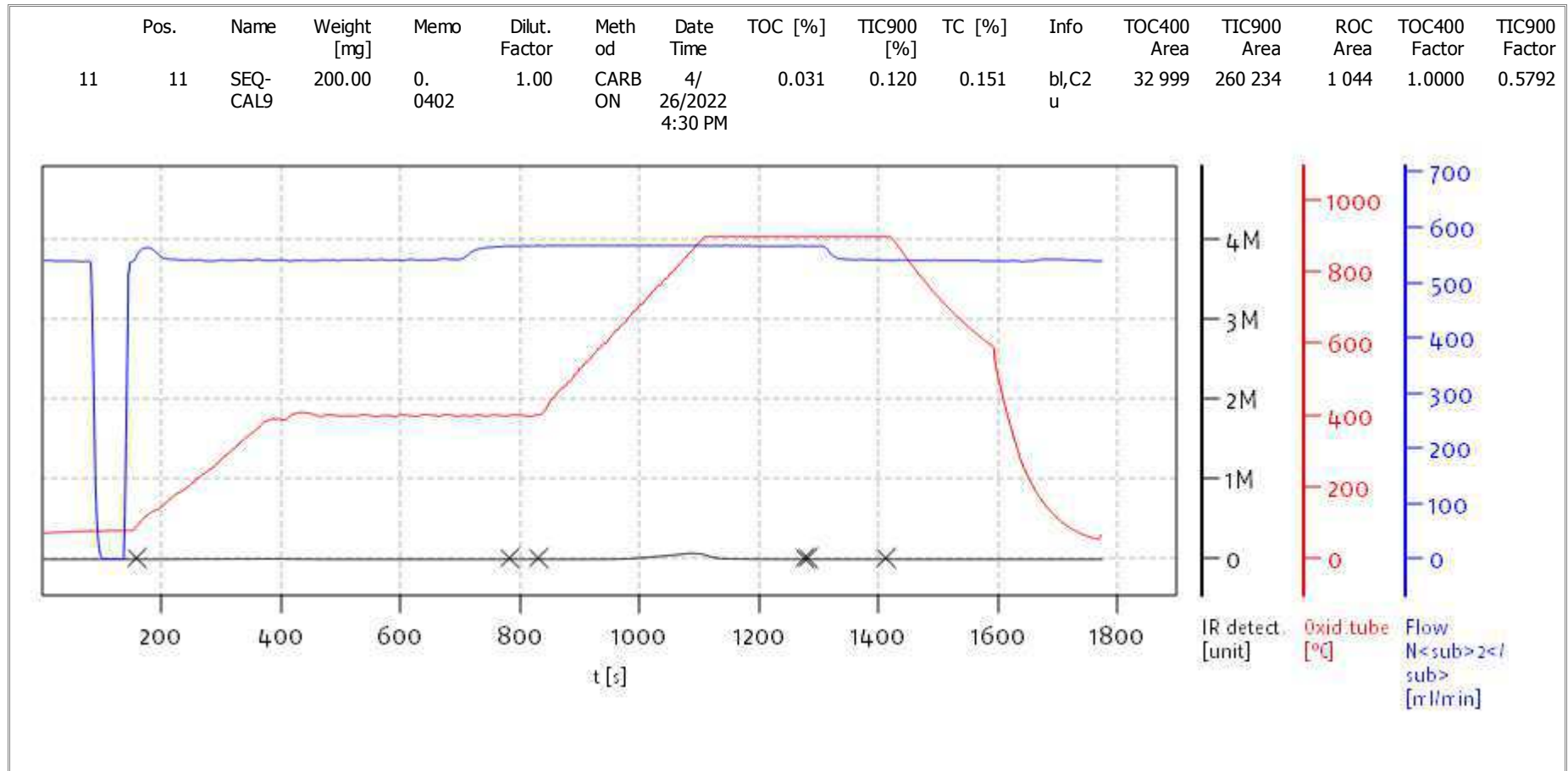
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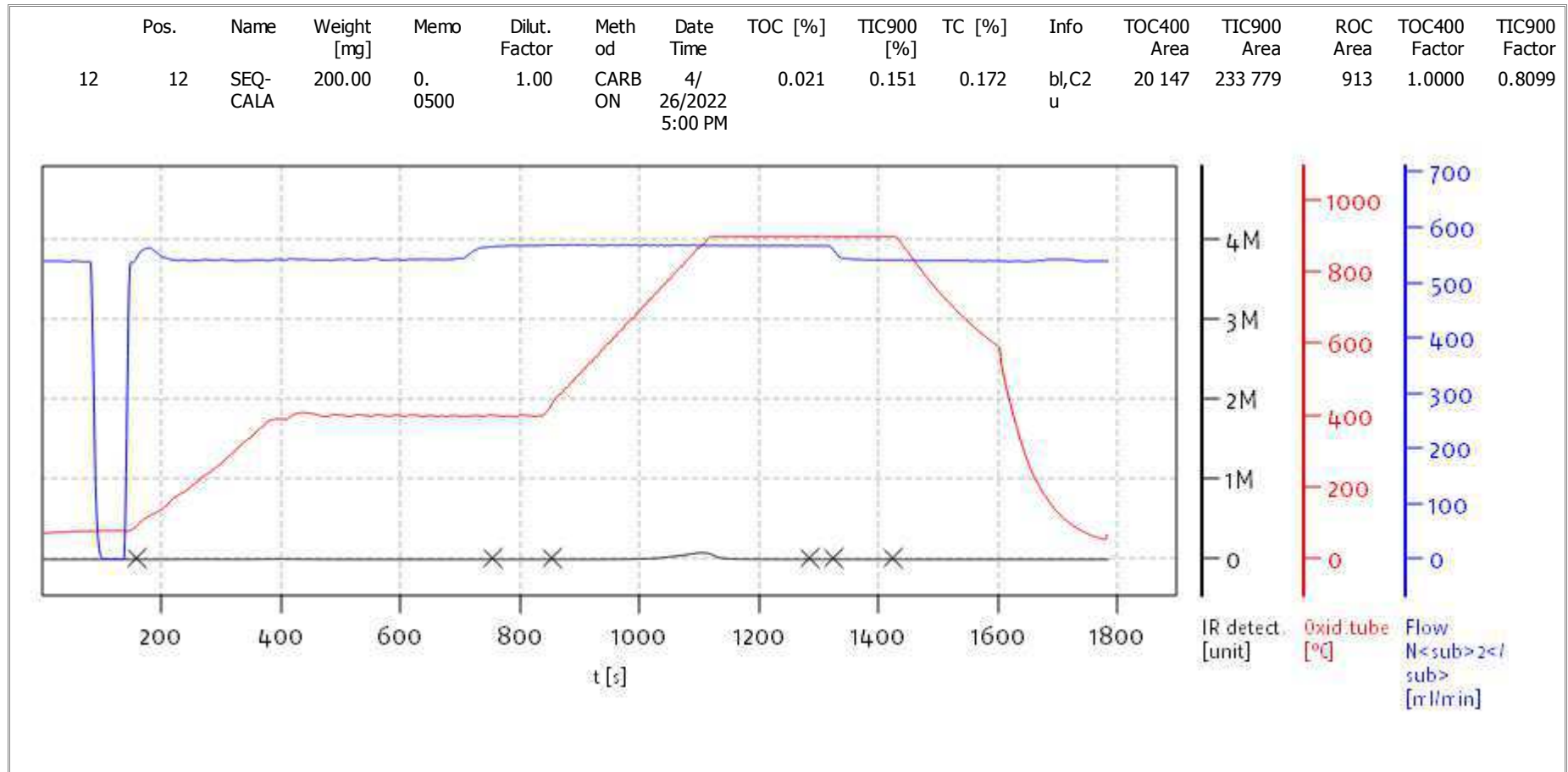
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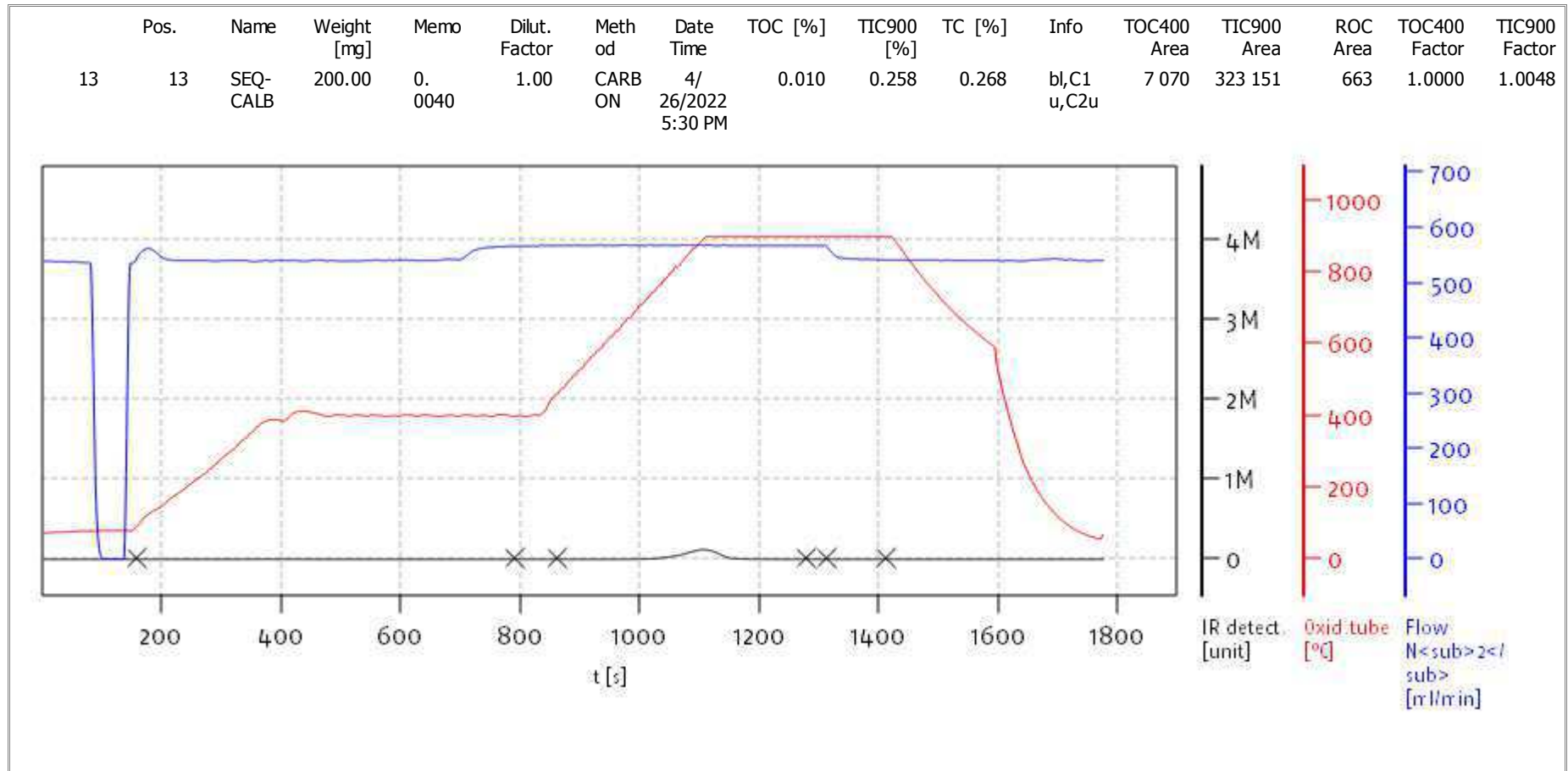
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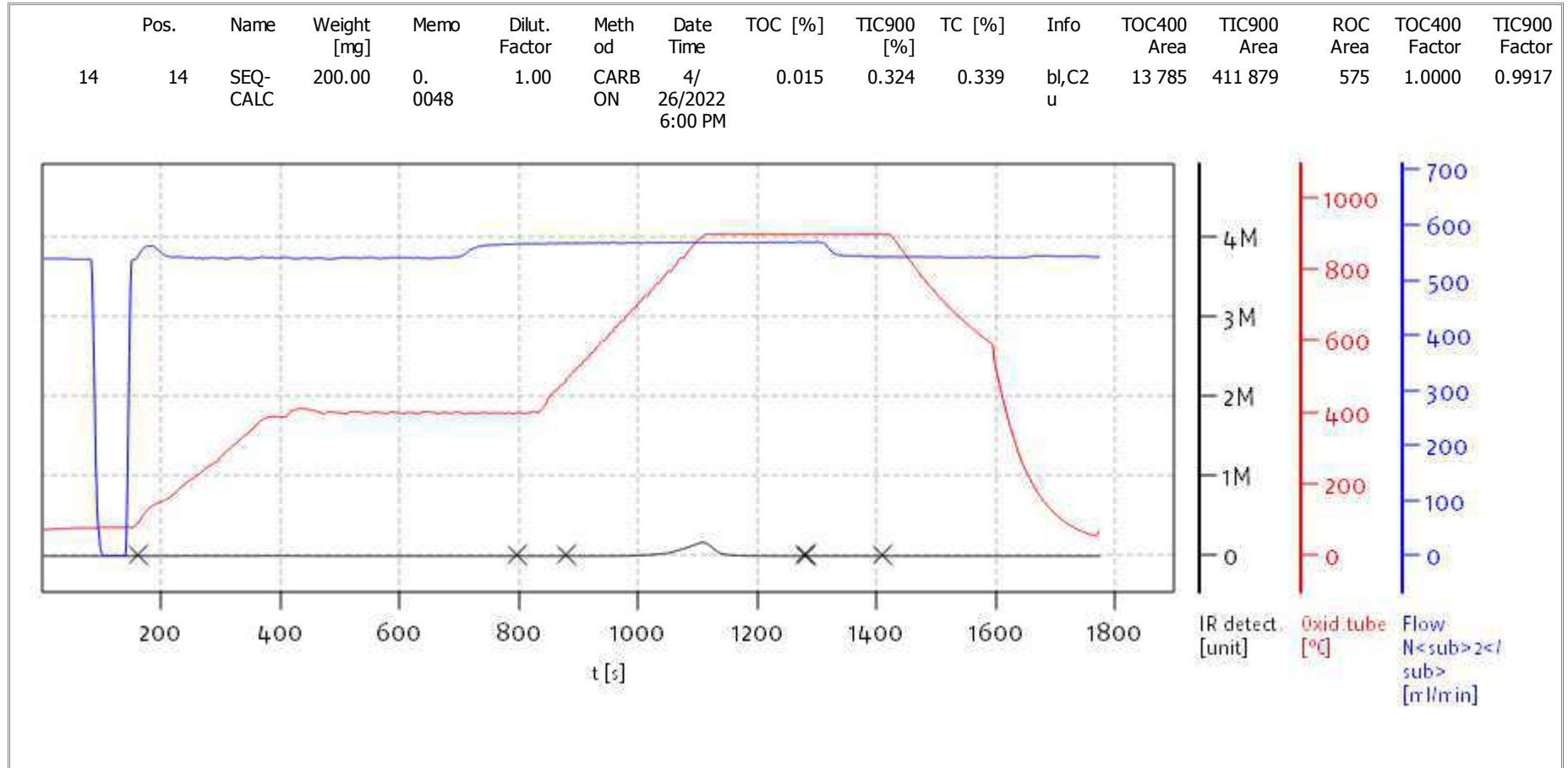
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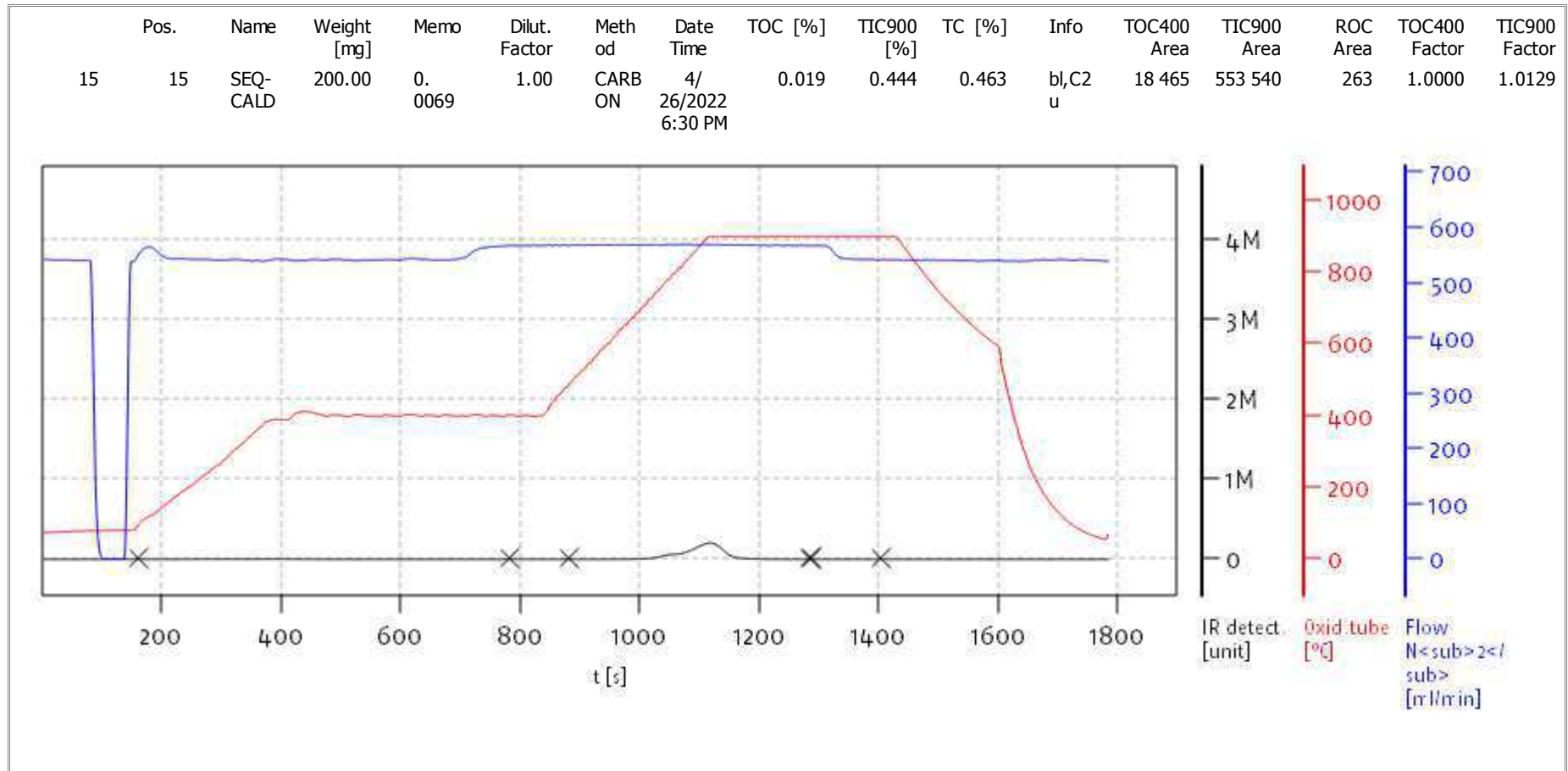
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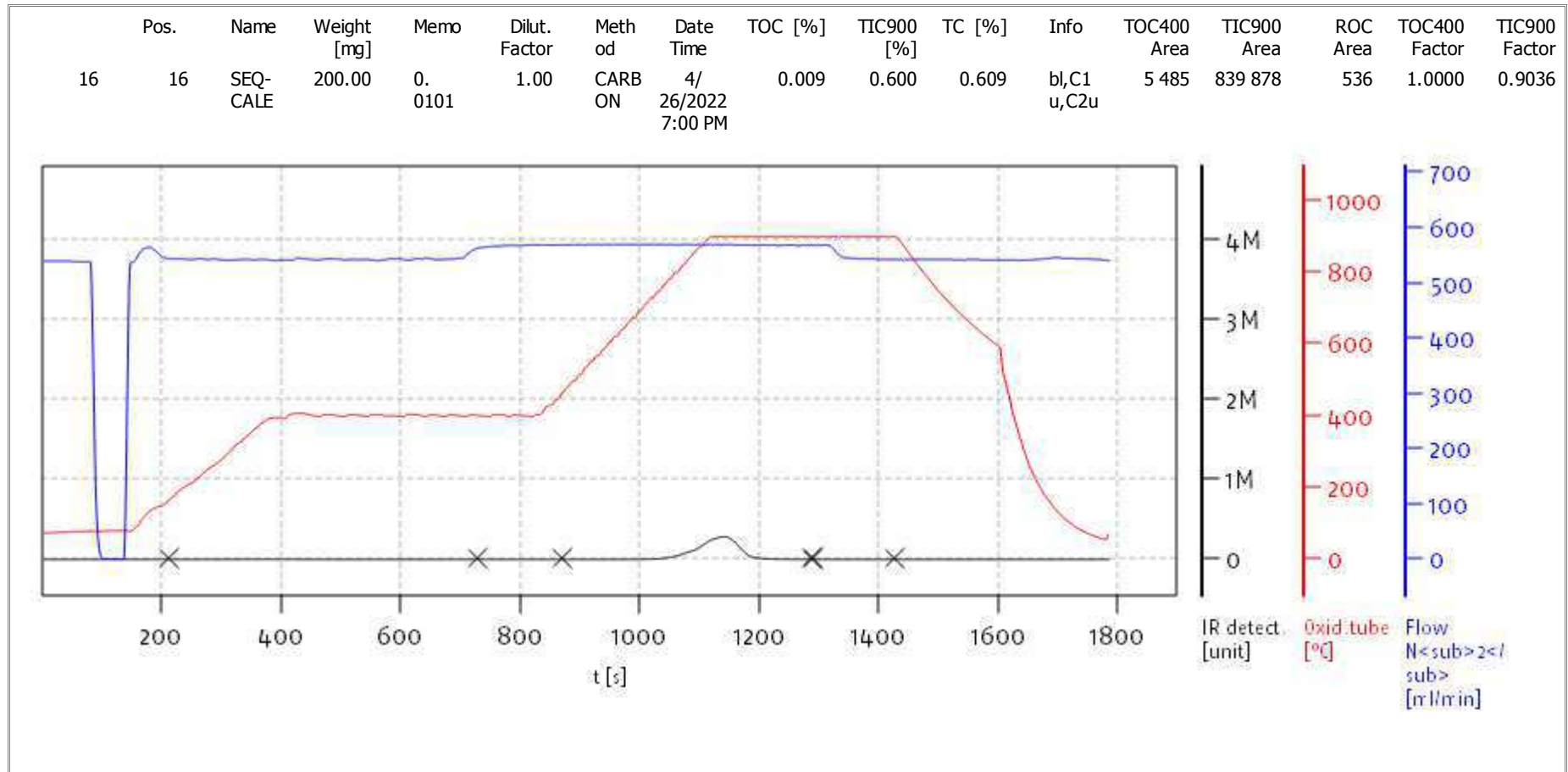
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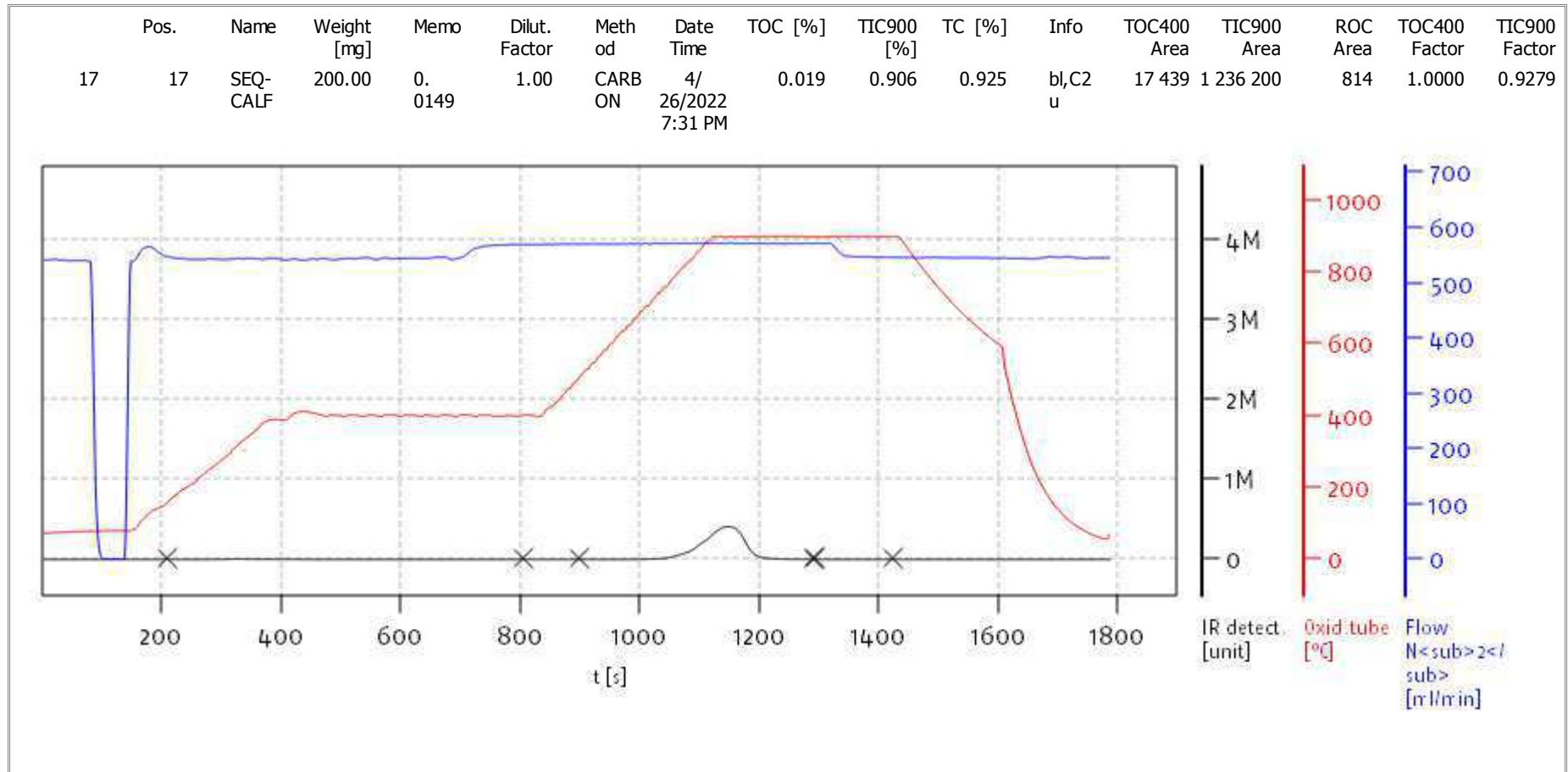
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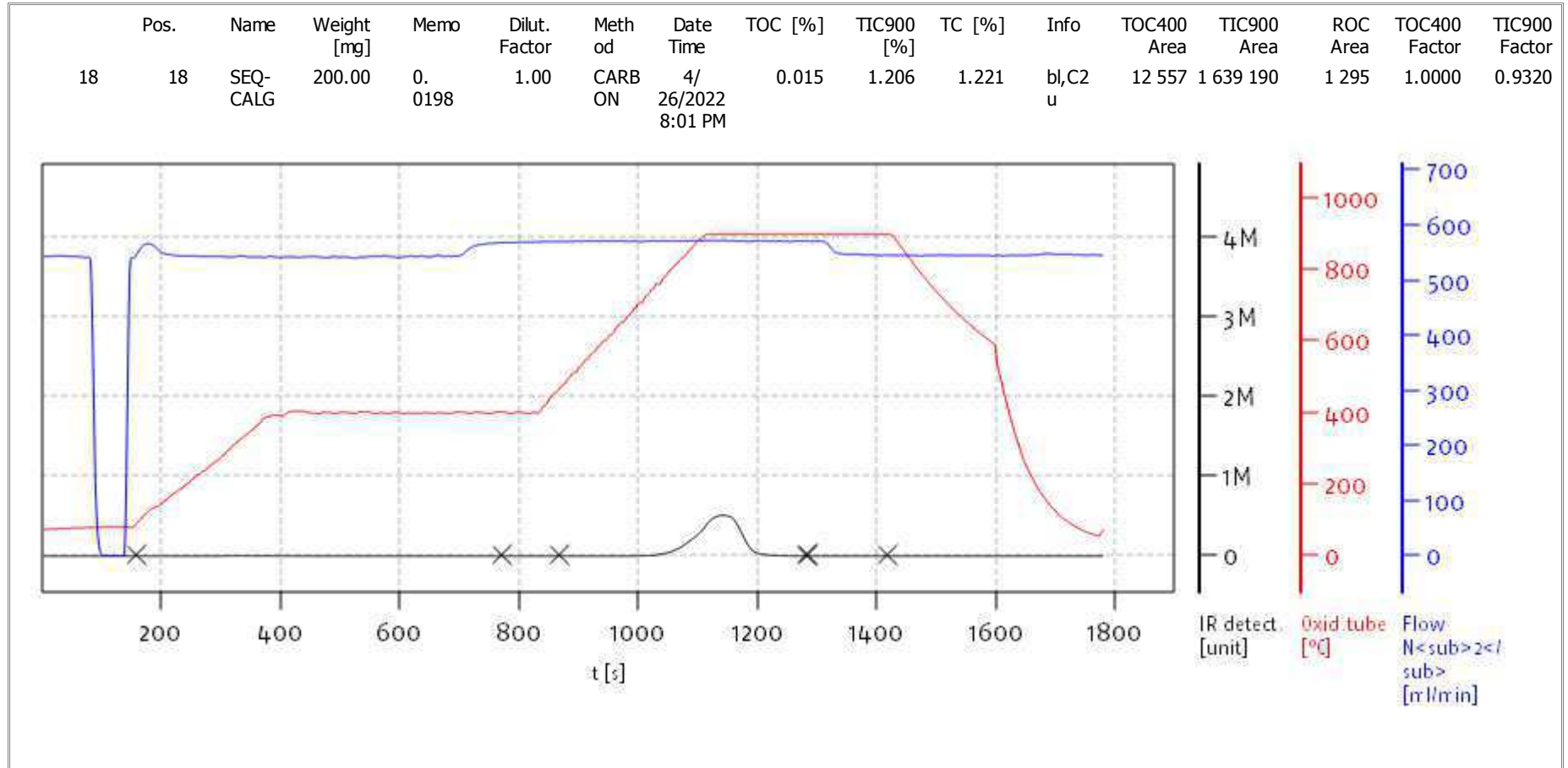
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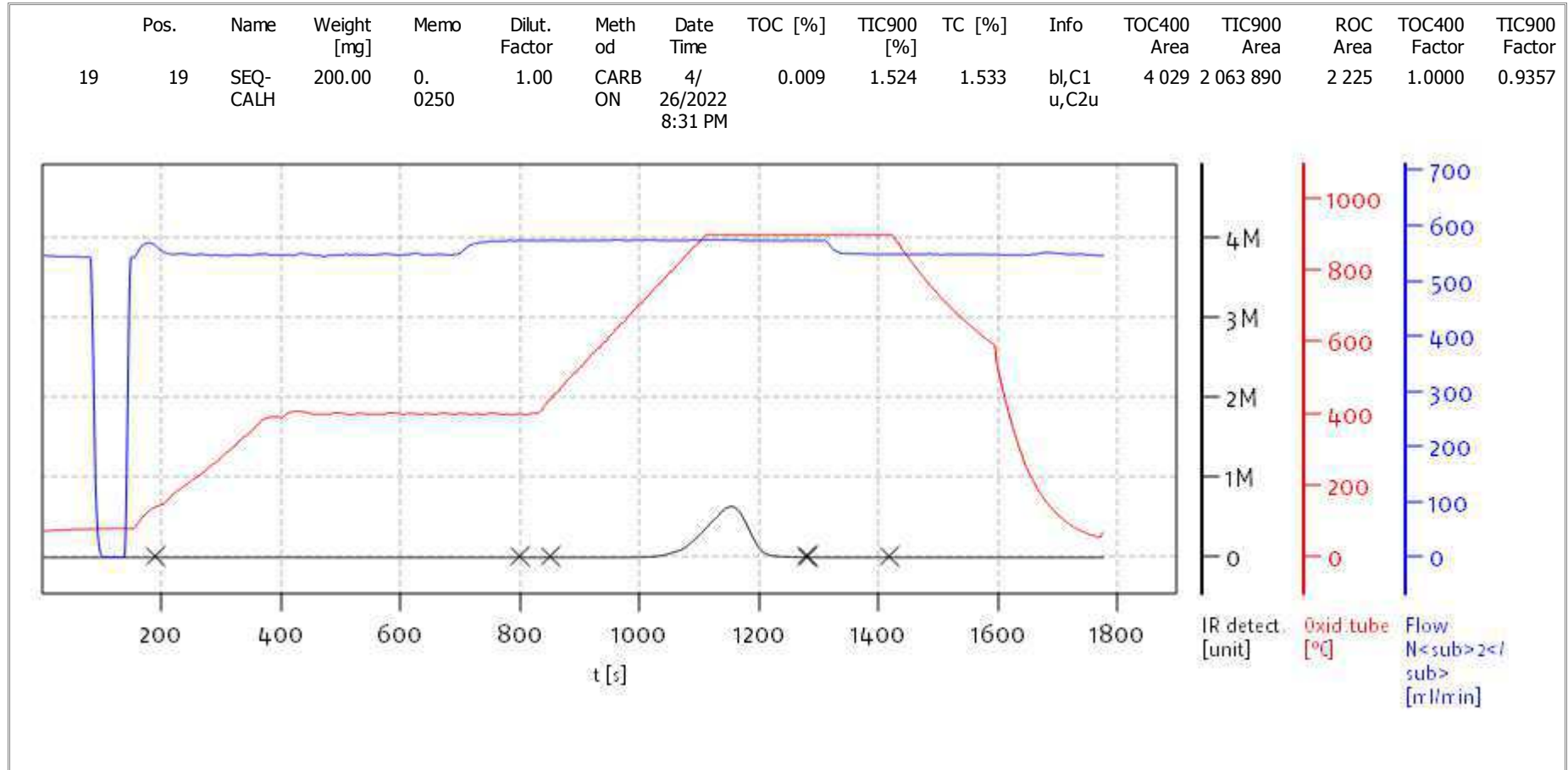
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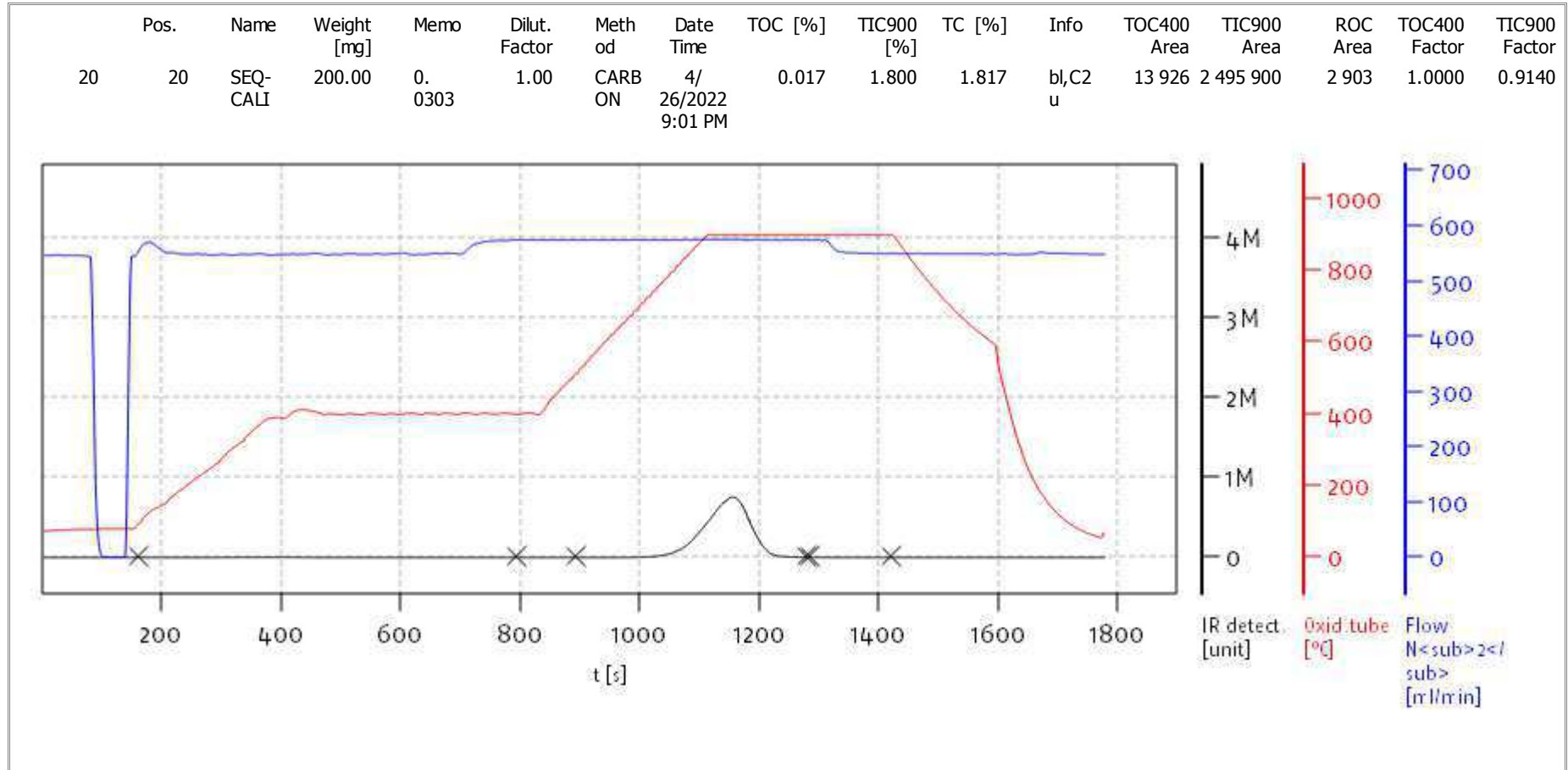
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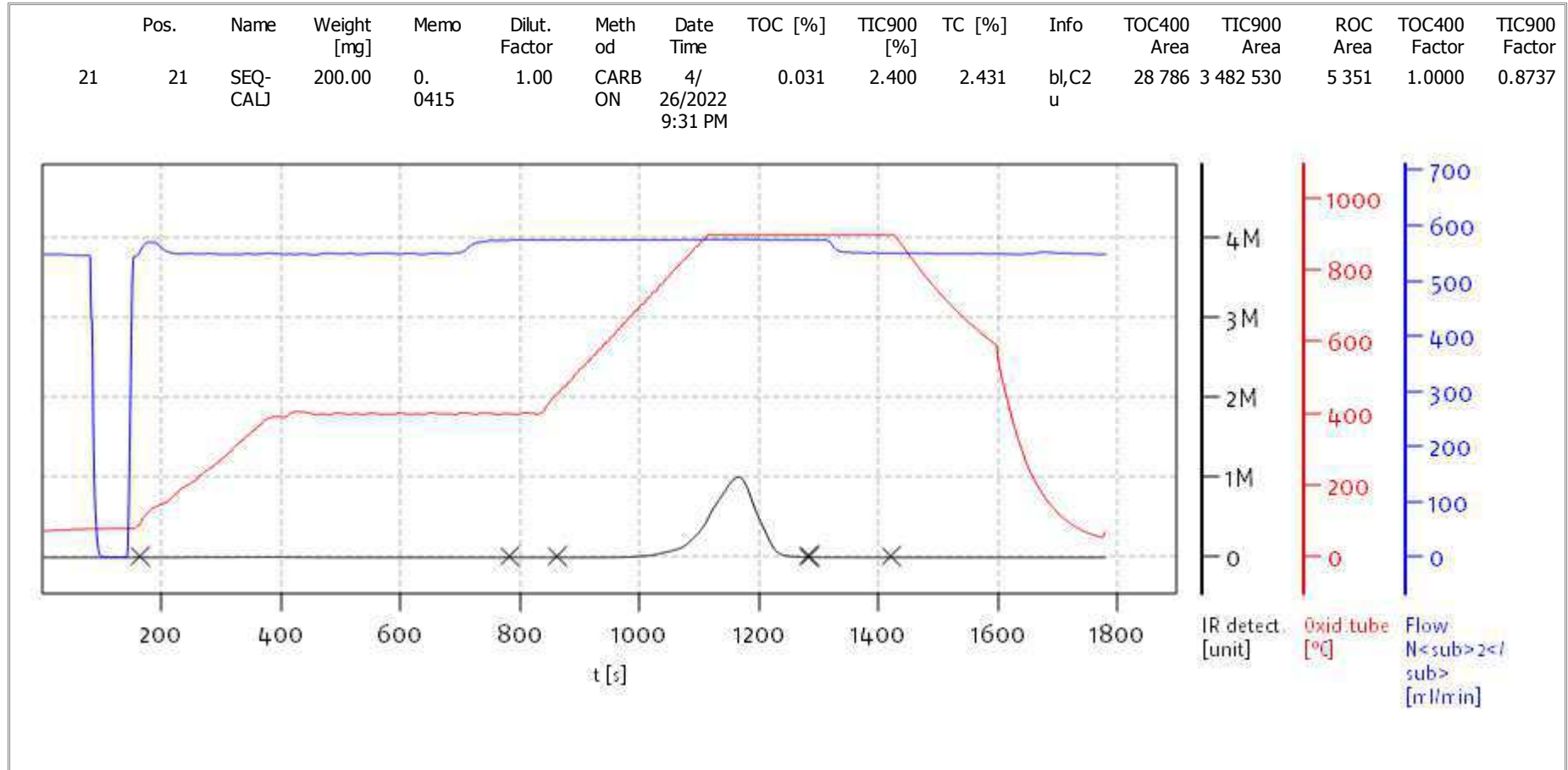
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solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

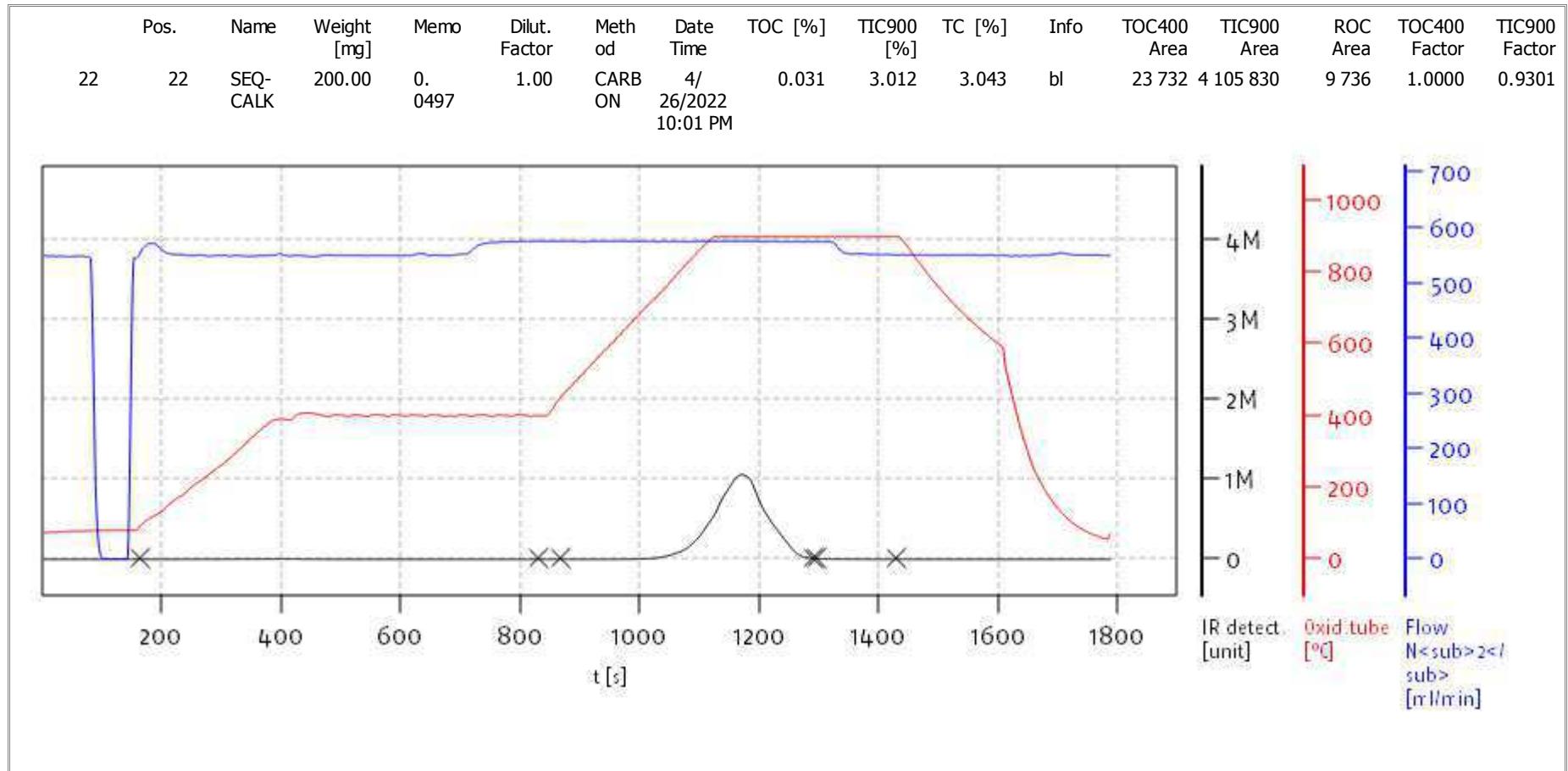
Date: Wed Apr 27 11:07:12 2022



solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

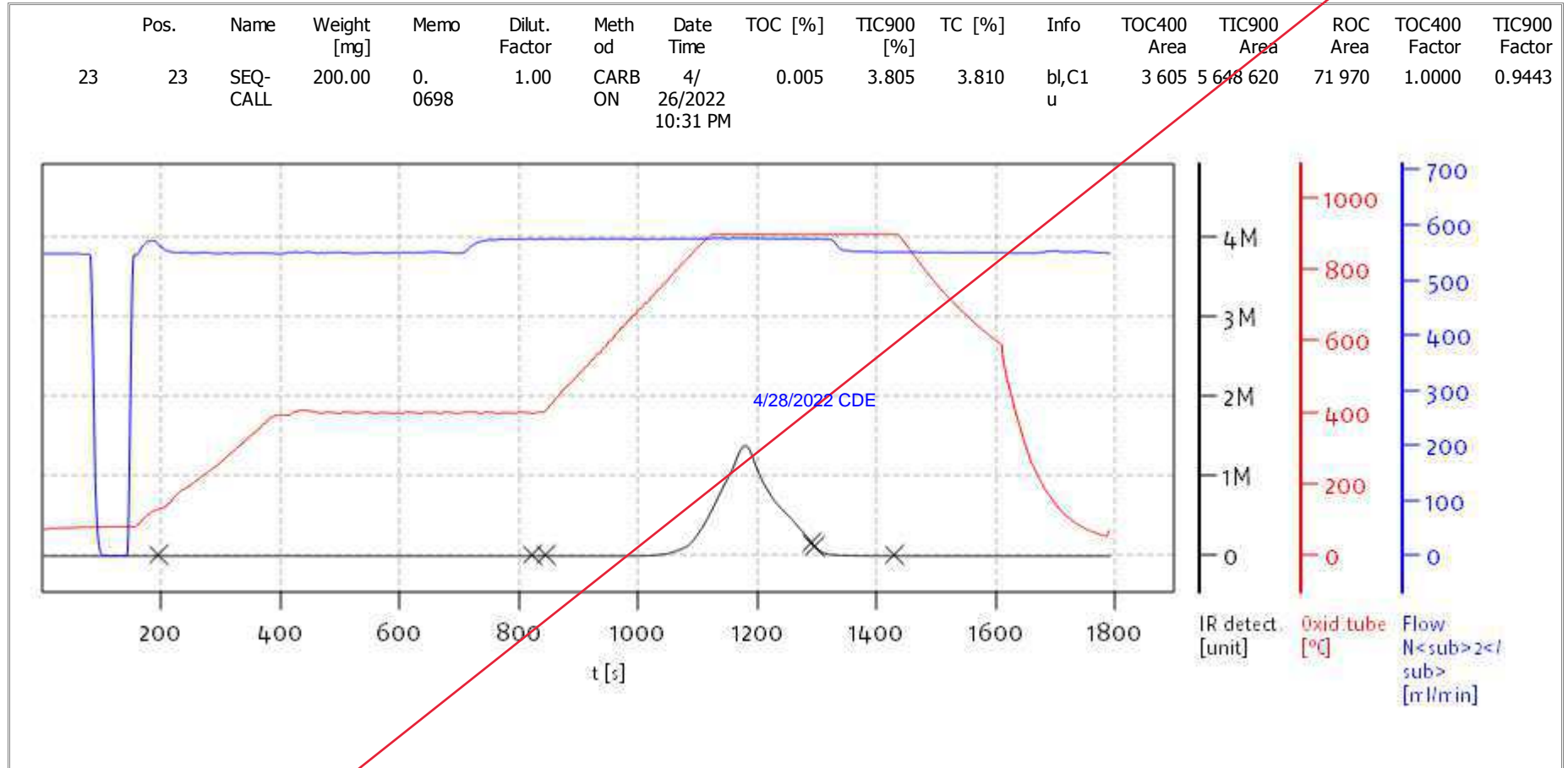
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solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

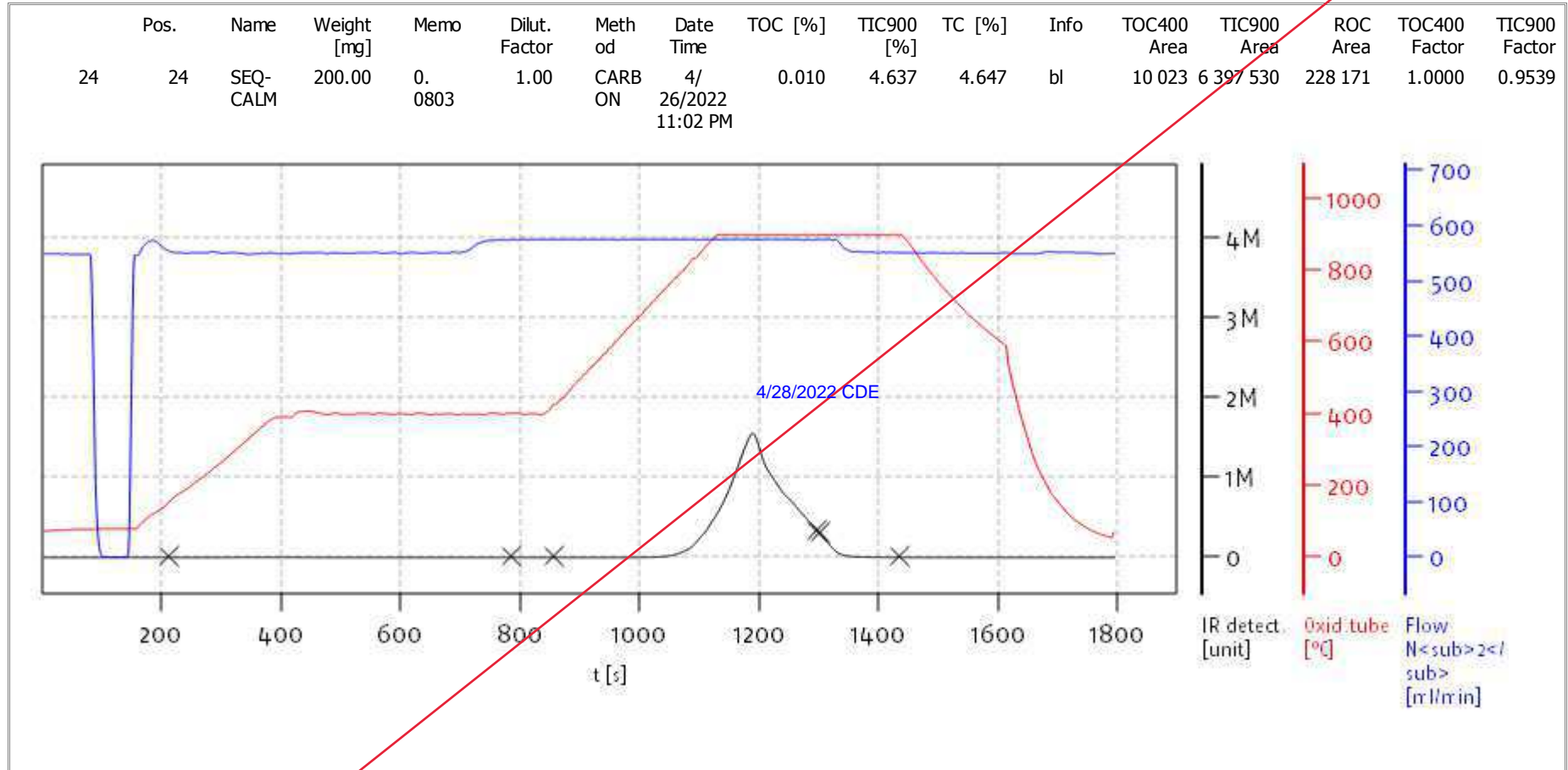
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solITOC V2.0.2 (31015f9) 2018-11-19
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Soli TOC Cube, Carbon
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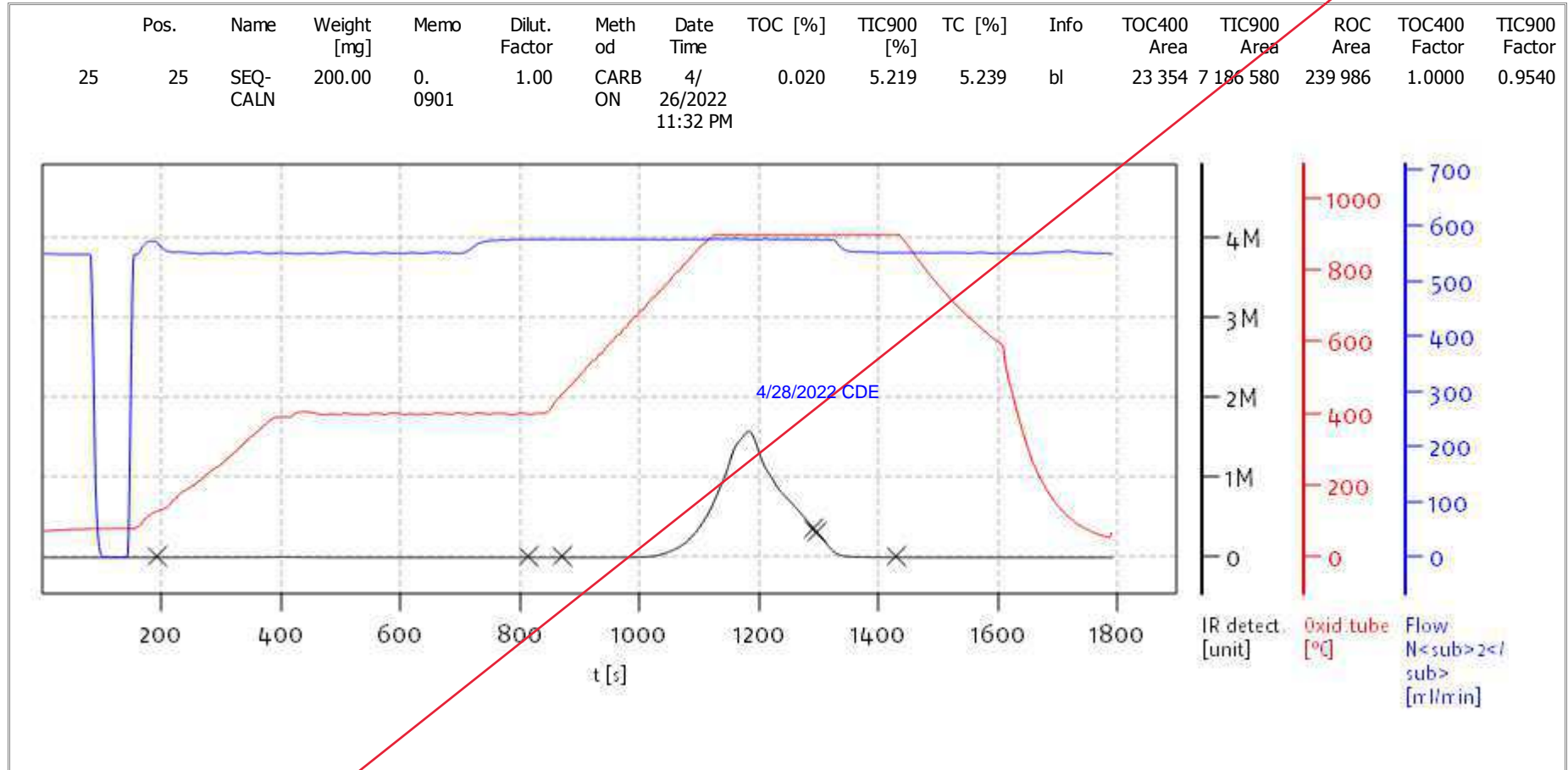
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solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

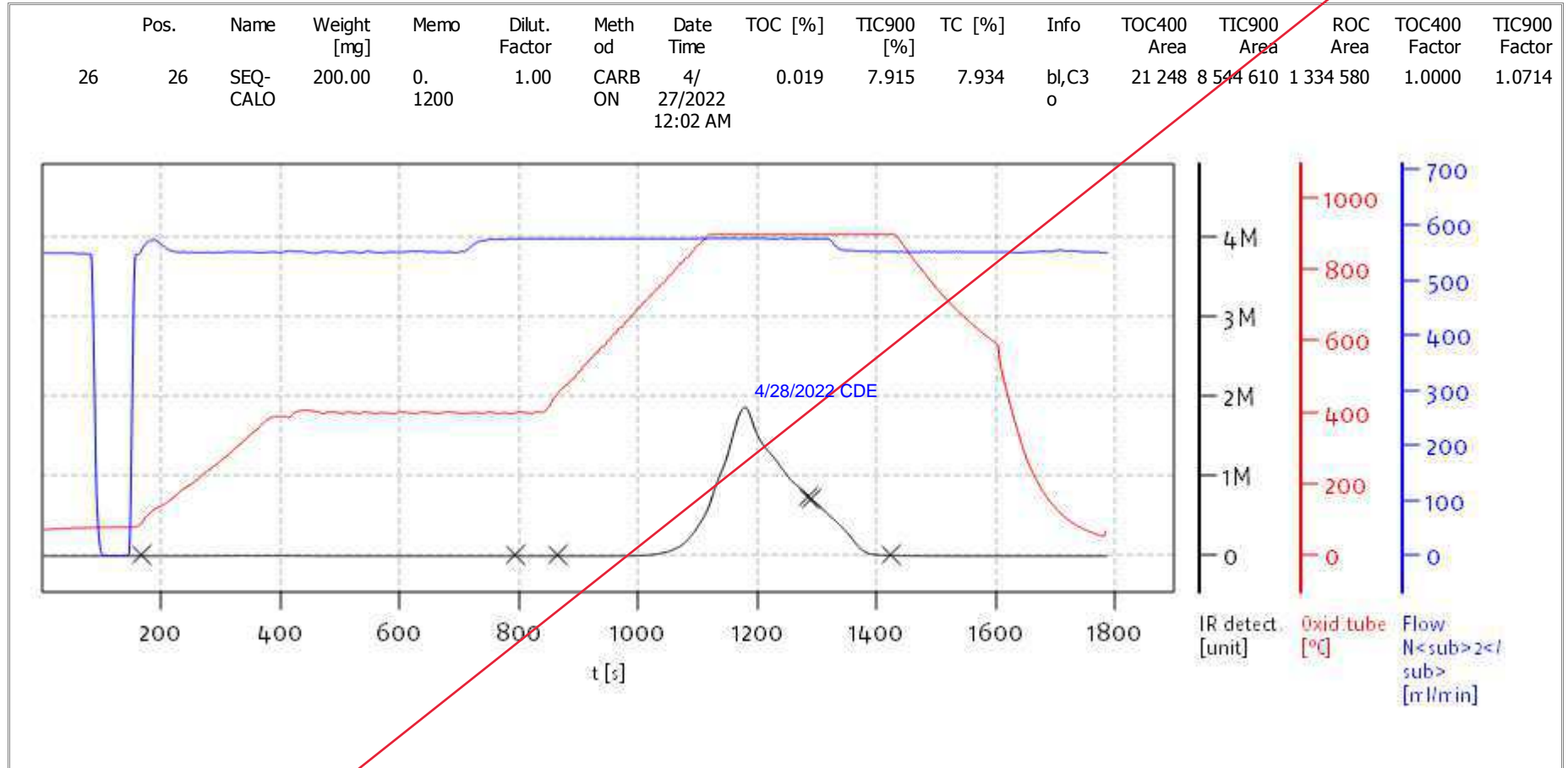
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solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
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Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

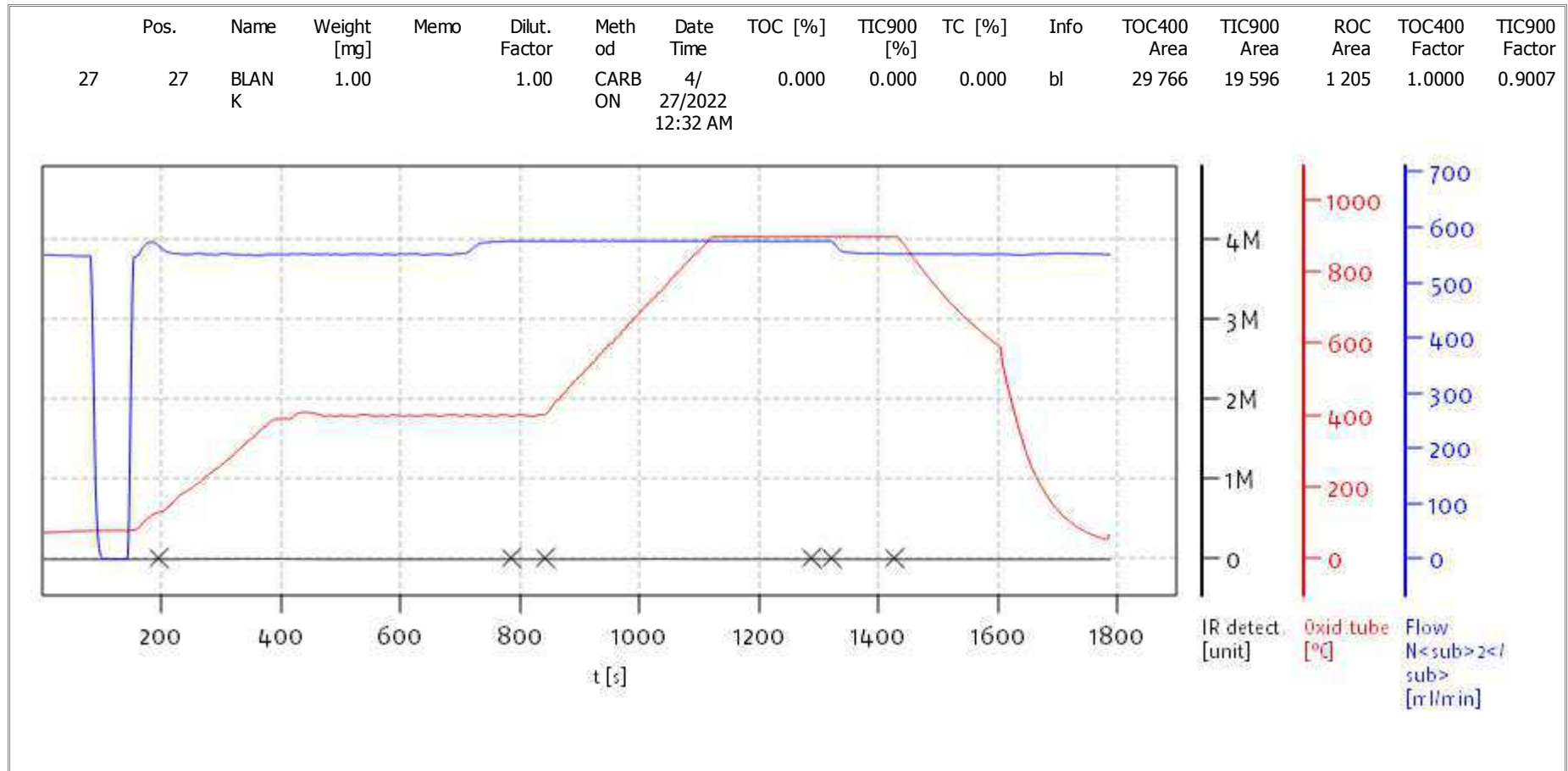
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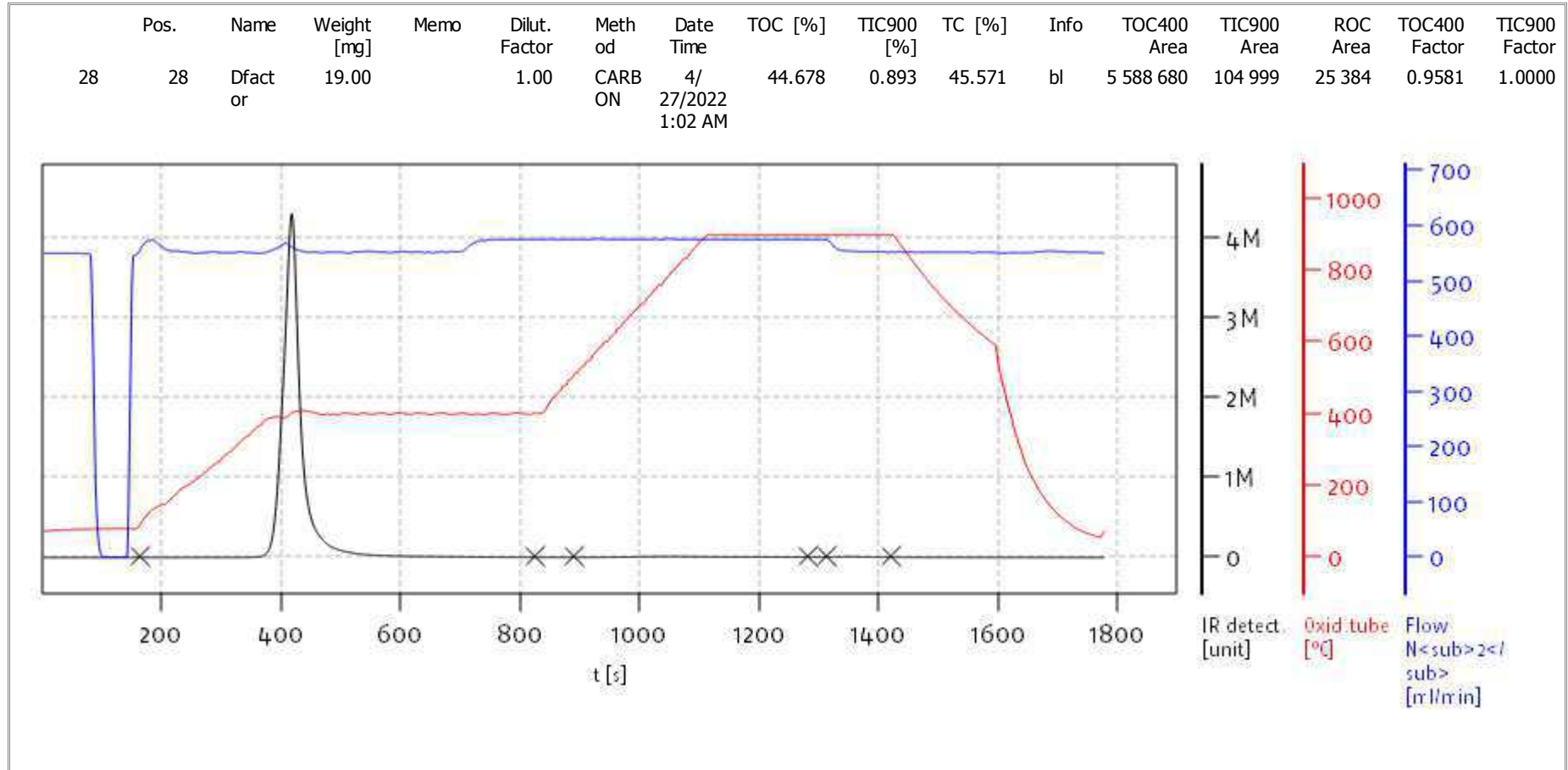
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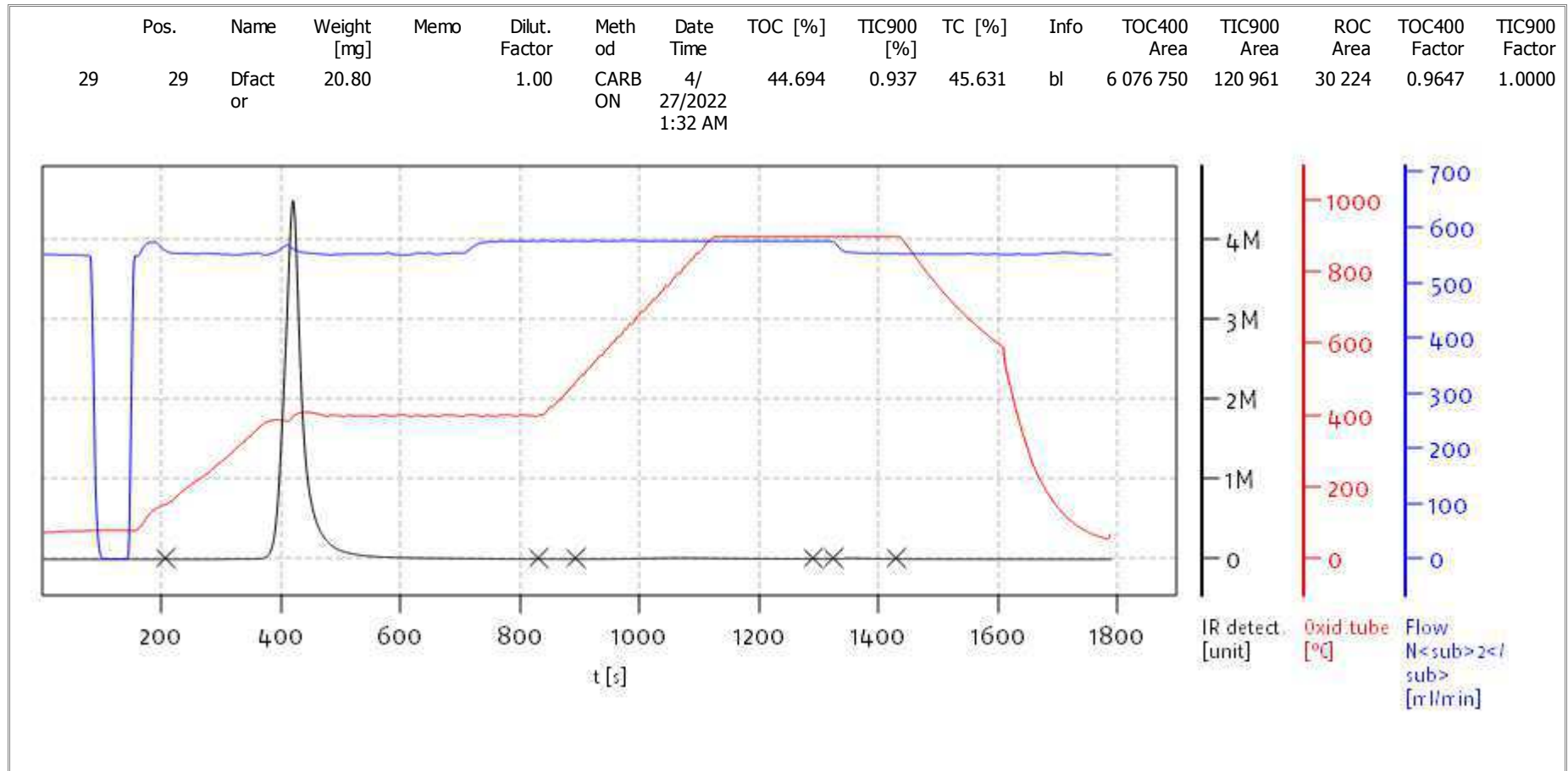
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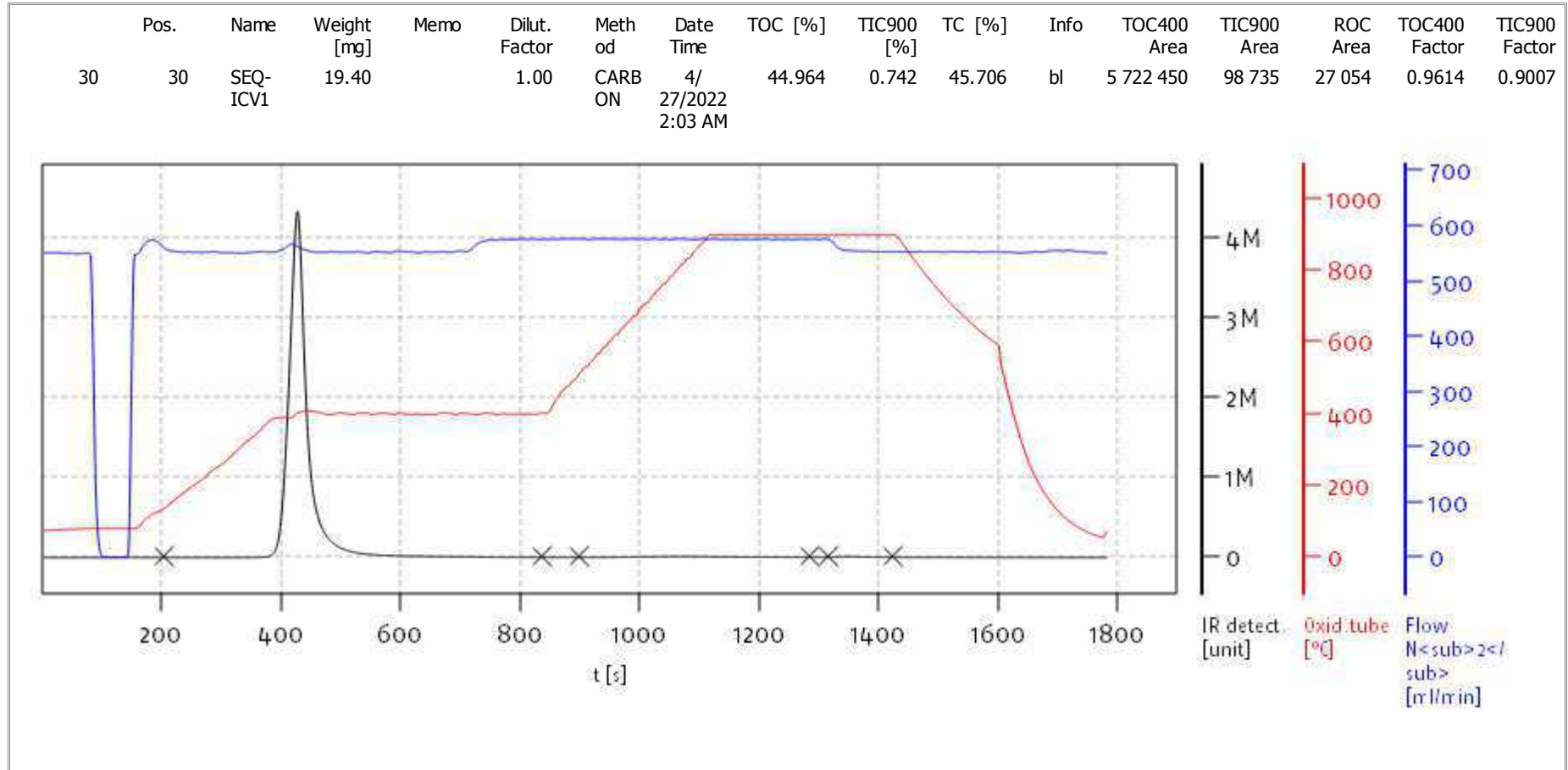
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Name:

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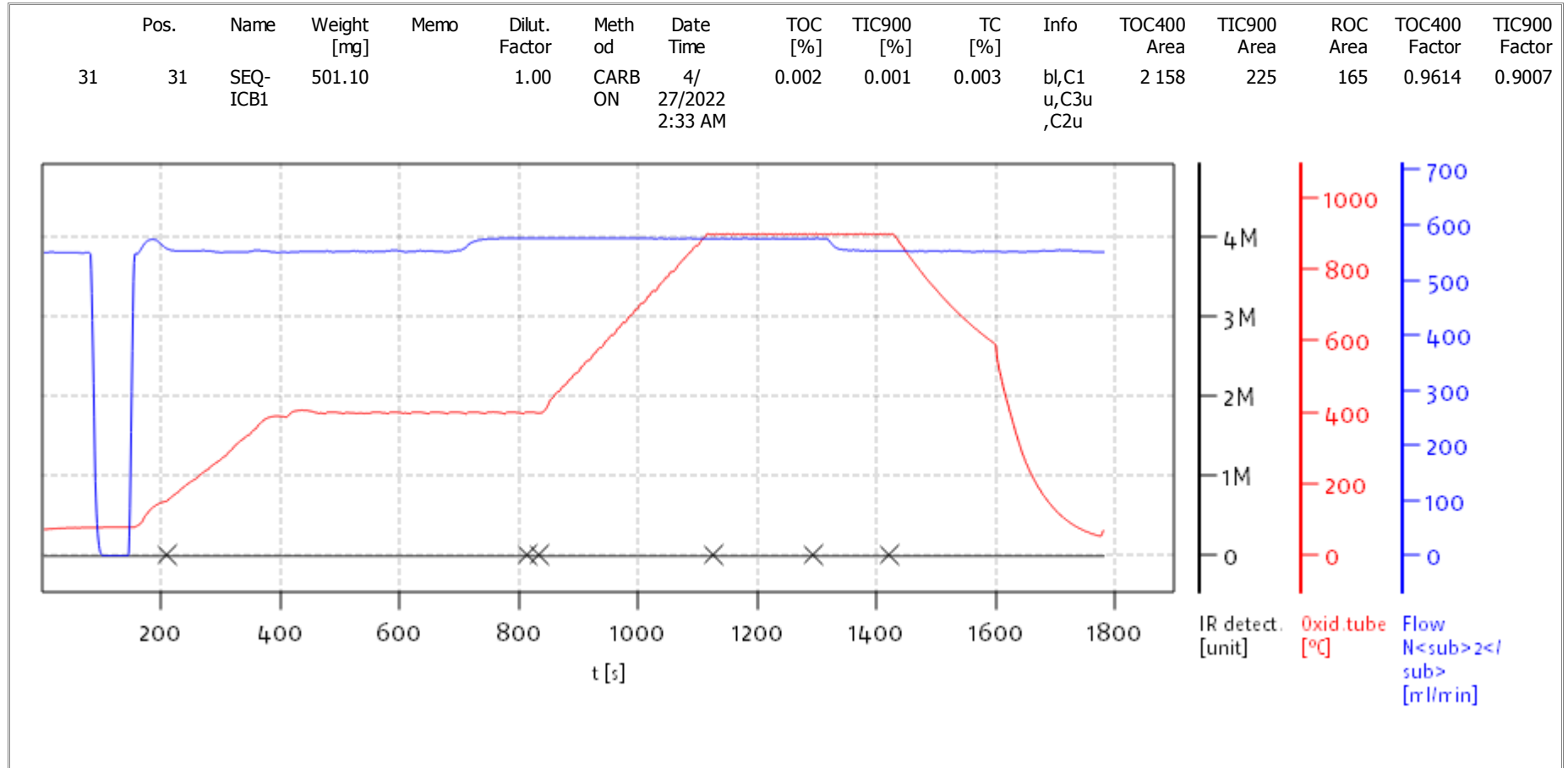
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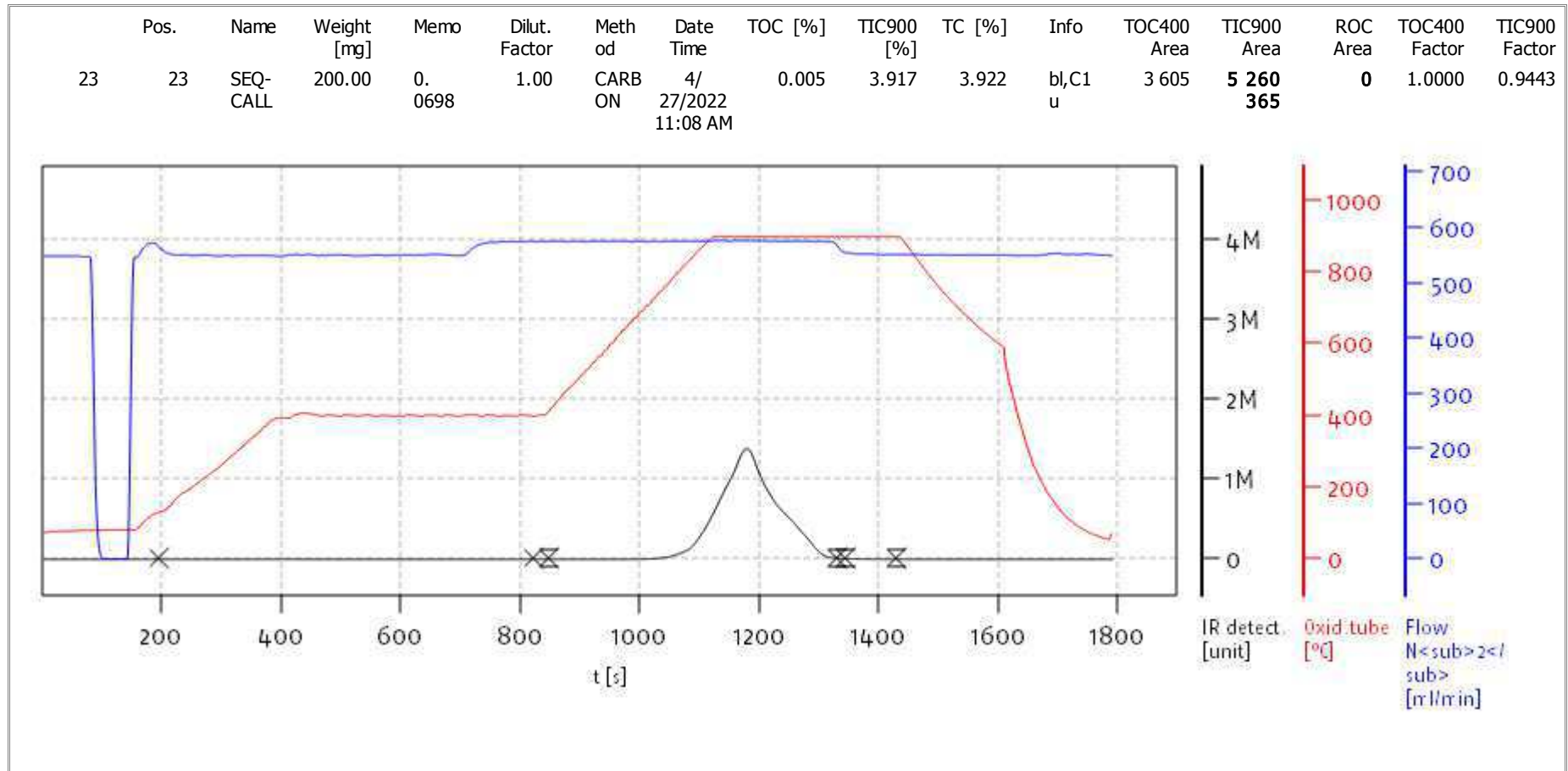
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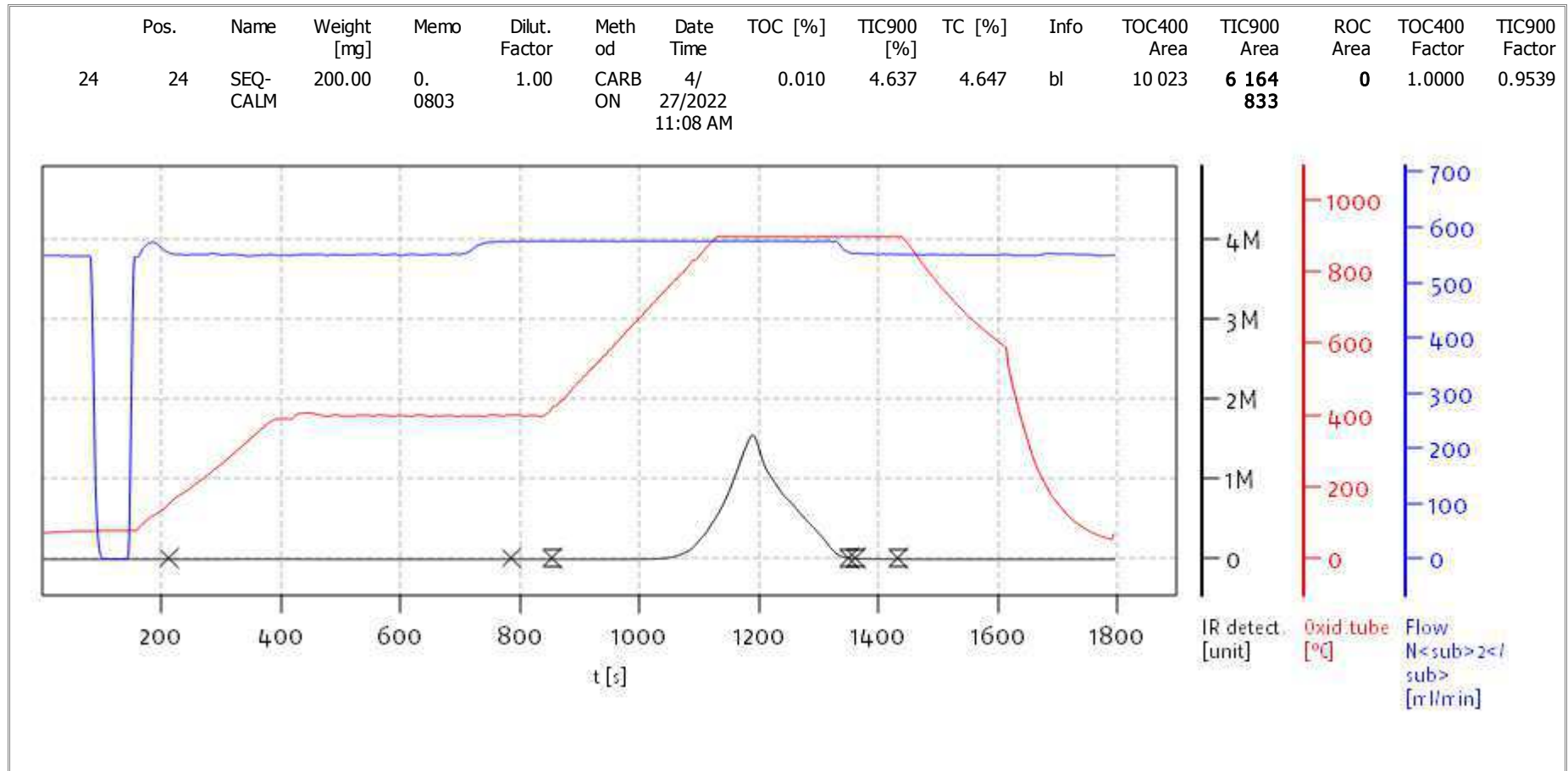
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solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
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Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



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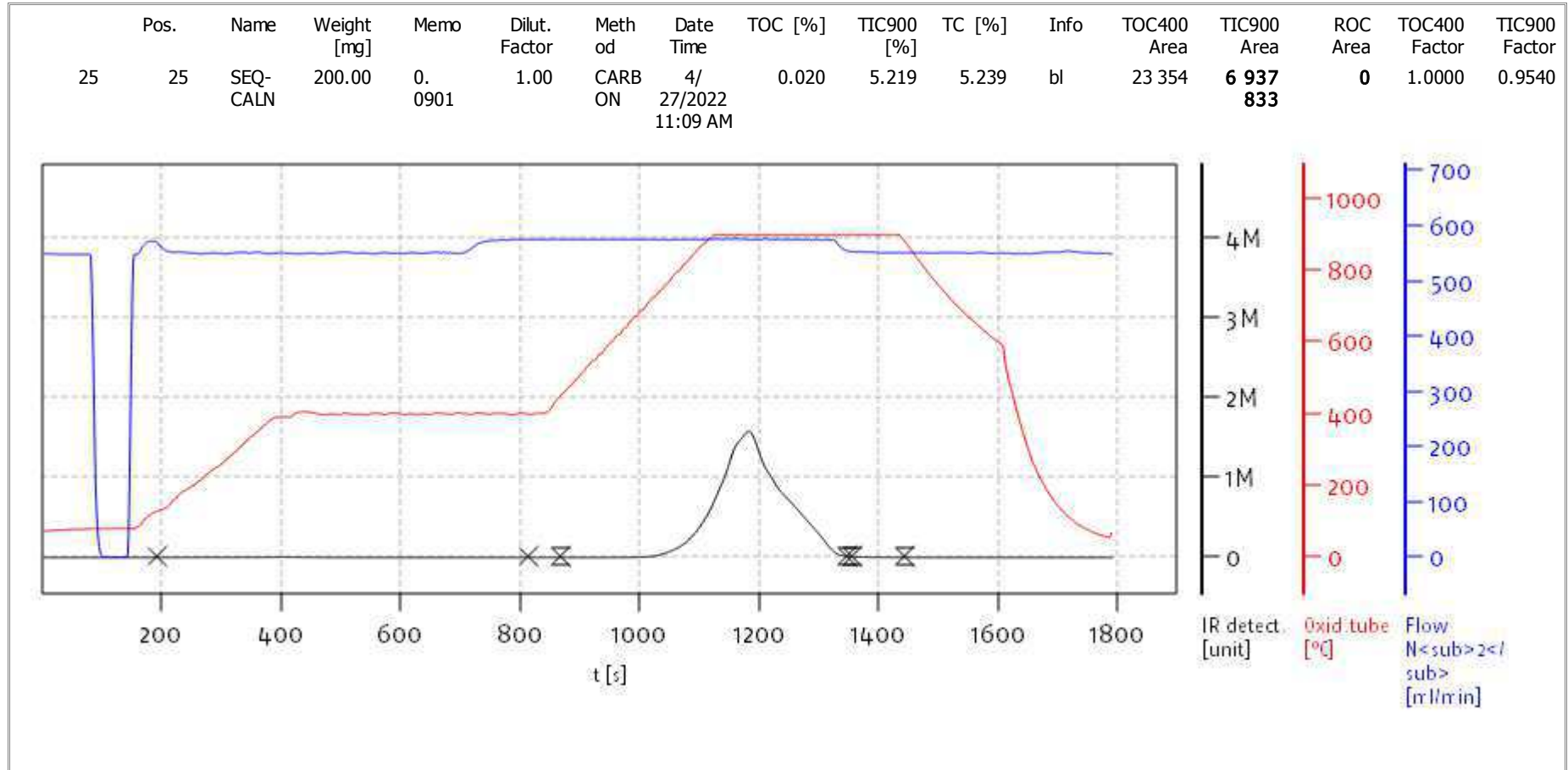
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solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

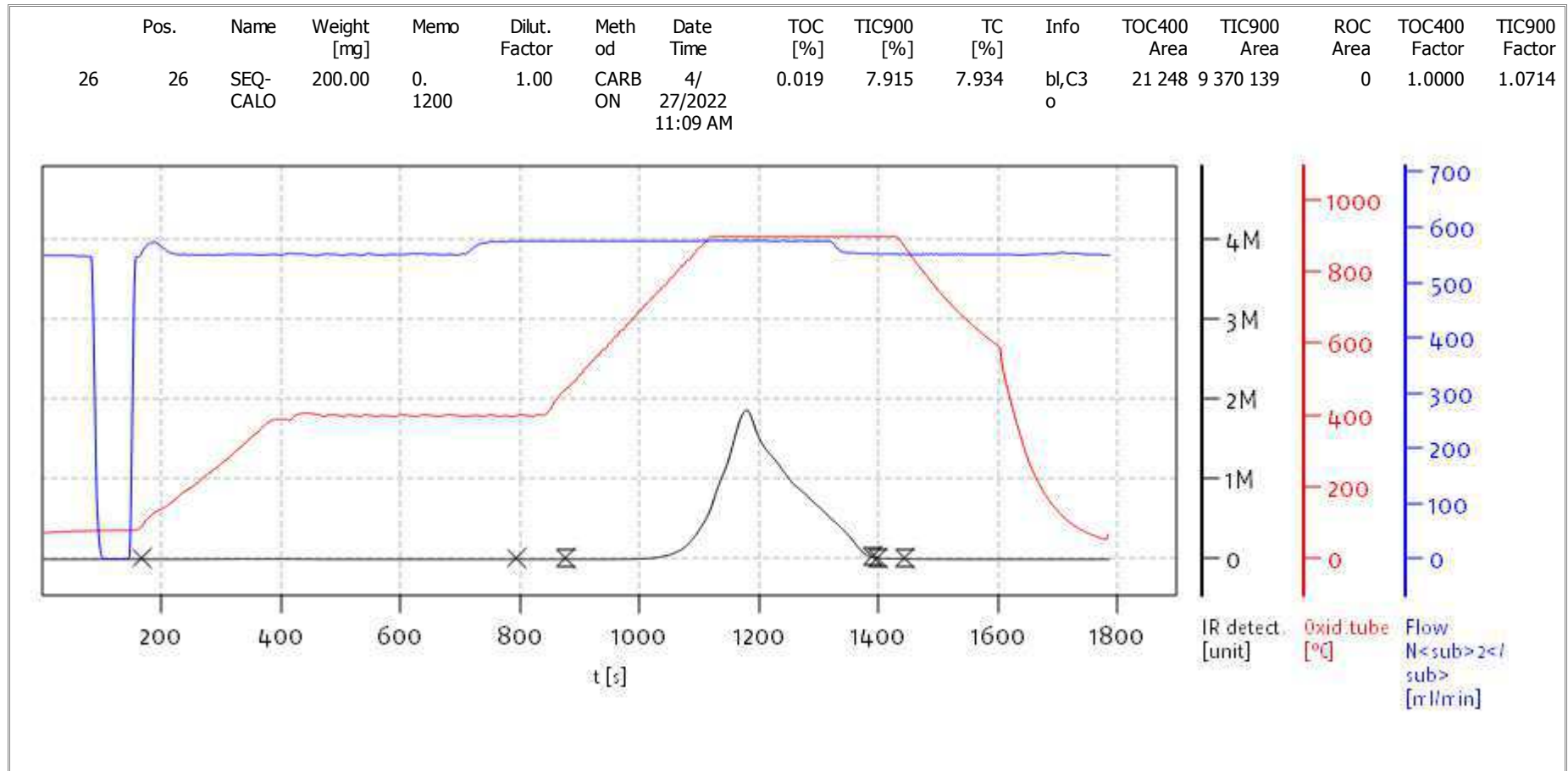
Date: Wed Apr 27 11:10:16 2022



solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC



Soli TOC Cube, Carbon
Balance: BAL3
Analyst: DOE



Name:

Access: solITOC superuser

Date: Wed Apr 27 11:10:16 2022

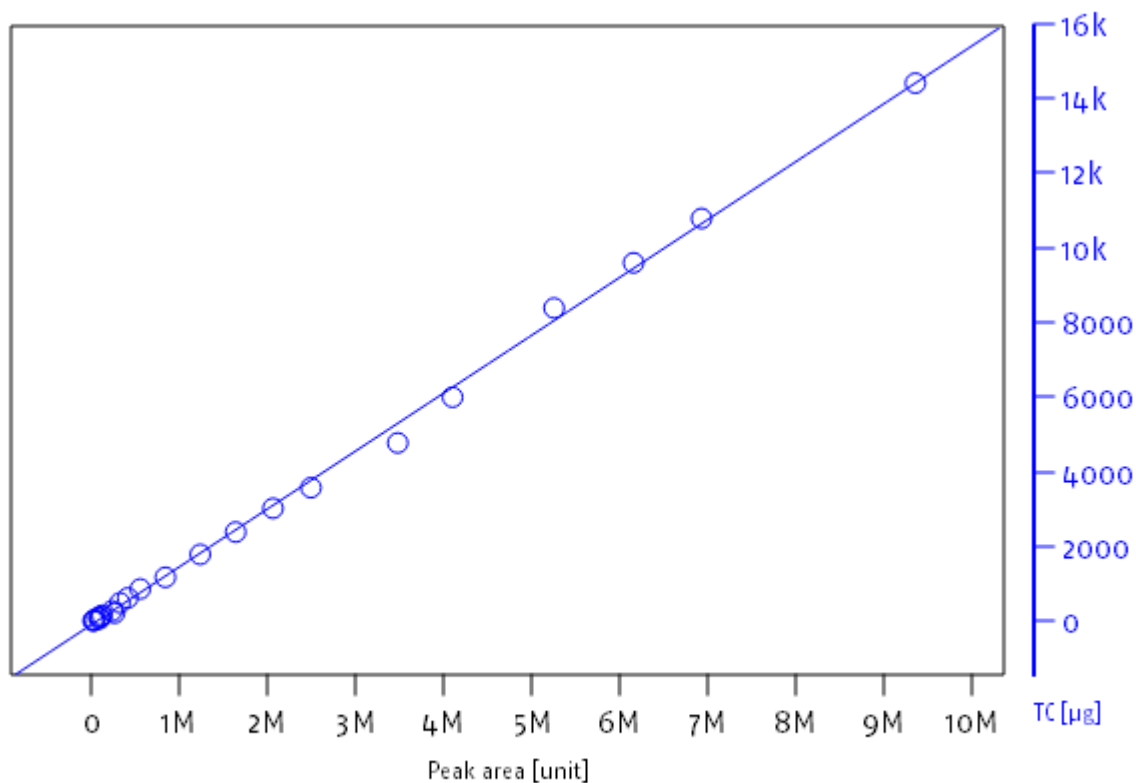


solITOC V2.0.2 (31015f9) 2018-11-19
Serial No: 0300.181017
Mode CCC

Calibration parameters TC, Whole range

a	-4.107546e-02
b	+1.548032e-06
c	+0.000000e+00
d	+0.000000e+00
e	+0.000000e+00
r	0.998372
r_old	0.998372
Proc.-SD	166.070255 µg

Calibration graph TC, Whole range



Name:

Access: solITOC superuser

Date: Wed Apr 27 11:19:56 2022



solITOC V2.0.2 (31015f9) 2018-11-19

Serial No: 0300.181017

Mode CCC



INSTRUMENT BLANKS
EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: TOC Cube

Calibration: FD00070

Sequence: SKD0371

Date Analyzed: 04/27/22 02:33

Lab Sample ID	Analyte	Found	MDL	MRL	Units	C
SKD0371-ICB1	Total Organic Carbon	0.00	0.02	0.02	%	



INSTRUMENT BLANKS
EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: TOC Cube

Calibration: FD00070

Sequence: SLD0078

Date Analyzed: 04/06/23 12:27

Lab Sample ID	Analyte	Found	MDL	MRL	Units	C
SLD0078-ICB1	Total Organic Carbon	0.002	0.02	0.02	%	
SLD0078-CCB1	Total Organic Carbon	0.005	0.02	0.02	%	
SLD0078-CCB2	Total Organic Carbon	0.00	0.02	0.02	%	
SLD0078-CCB3	Total Organic Carbon	0.00	0.02	0.02	%	
SLD0078-CCB4	Total Organic Carbon	0.00	0.02	0.02	%	



**INITIAL AND CONTINUING
CALIBRATION CHECK
EPA 9060A m**

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: TOC Cube

Calibration: FD00070

Control Limit: +/- 10.00%

Sequence: SKD0371

Lab Sample ID	Analyte	True	Found	%R	Units	Method
SKD0371-ICV1	Total Organic Carbon	44.446	43.7	98.3	%	EPA 9060A m
	Total Carbon	44.446	44.1	99.2	%	EPA 9060A m
	Total Inorganic Carbon	0.0000	0.40		%	EPA 9060A m
	% Soot	0.0000	0.004		%	EPA 9060A m

* Values outside of QC limits



**INITIAL AND CONTINUING
CALIBRATION CHECK**
EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Instrument ID: TOC Cube

Calibration: FD00070

Control Limit: +/- 10.00%

Sequence: SLD0078

Lab Sample ID	Analyte	True	Found	%R	Units	Method
SLD0078-ICV1	Total Organic Carbon	44.446	43.5	97.8	%	EPA 9060A m
SLD0078-CCV1	Total Organic Carbon	44.446	44.4	100	%	EPA 9060A m
SLD0078-CCV2	Total Organic Carbon	44.446	45.8	103	%	EPA 9060A m
SLD0078-CCV3	Total Organic Carbon	44.446	43.7	98.4	%	EPA 9060A m
SLD0078-CCV4	Total Organic Carbon	44.446	45.3	102	%	EPA 9060A m

* Values outside of QC limits



STANDARD REFERENCE MATERIAL RECOVERY

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Laboratory ID: BLD0117-SRM1

Batch: BLD0117

Initial/Final: 0.3111 g / 0.3111 mL

Preparation: PSEP 1986 (modified)

Analyzed: 04/06/2023 14:28

Standard ID: L000790

Expires: 02/07/2023

Standard Lot#: NA

Description: 1941B - Organics in Marine Sediment (Conv

ANALYTE	TRUE (% wet)	FOUND (% wet)	MDL	MRL	Q	SRM % REC.	QC LIMITS REC.
Total Organic Carbon	2.9900	3.03	0.02	0.02		101	80 - 120

* Values outside of QC limits



HOLDING TIME SUMMARY

Analysis: EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Sample Name	Date Collected	Date Received	Date Prepared	Days to Prep	Max Days to Prep	Date Analyzed	Days to Analysis	Max Days to Analysis	Q
LDW22-SS826 23C0107-01	03/02/23 16:25	03/03/23 16:35	04/05/23 15:19	33	180	04/07/23 01:37			
LDW21-IT608C 23C0107-02	07/13/21 06:54	03/03/23 16:35	04/05/23 15:19	631	180	04/07/23 02:07			*
LDW21-IT608D 23C0107-03	07/13/21 06:54	03/03/23 16:35	04/05/23 15:19	631	180	04/07/23 02:38			*

* Indicates hold time exceedance.



Analytical Resources, LLC
Analytical Chemists and Consultants

METHOD DETECTION AND REPORTING LIMITS

EPA 9060A m

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Instrument: TOC Cube

Analyte	MDL	RL	Units
Total Organic Carbon	0.02	0.02	%



National Institute of Standards & Technology

Certificate of Analysis

Standard Reference Material® 1941b

Organics in Marine Sediment

This Standard Reference Material (SRM) is marine sediment collected at the mouth of the Baltimore (MD) Harbor. SRM 1941b is intended for use in evaluating analytical methods for the determination of selected polycyclic aromatic hydrocarbons (PAHs), polychlorinated biphenyl (PCB) congeners, and chlorinated pesticides in marine sediment and similar matrices. Information values are also provided for total organic carbon (TOC), total carbon, hydrogen, and nitrogen. All of the constituents for which certified, reference, and information values are provided in SRM 1941b were naturally present in the sediment before processing. A unit of SRM 1941b consists of a bottle containing 50 g of radiation-sterilized, freeze-dried sediment.

Certified Mass Fraction Values: Certified mass fraction values for PAHs, PCB congeners, and chlorinated pesticides are provided in Table 1 through Table 3. The certified values for the PAHs, PCB congeners, and chlorinated pesticides are based on the agreement of results obtained at NIST from two or more chemically independent analytical techniques along with results from an interlaboratory comparison study [1]. A NIST certified value is a value for which NIST has the highest confidence in its accuracy in that all known or suspected sources of bias have been investigated or taken into account [1].

Reference Mass Fraction Values: Reference mass fraction values for additional PAHs (some in combination), additional PCB congeners, and additional chlorinated pesticides are provided in Table 4 through Table 7. Reference values for alkylated PAH groups are provided in Table 8 and for selected hopanes and steranes in Table 9. A reference value for total organic carbon is provided in Table 10. Reference values are noncertified values that are the best estimate of the true value; however, the values do not meet the NIST criteria for certification and are provided with associated uncertainties that may reflect only measurement precision, may not include all sources of uncertainty, or may reflect a lack of sufficient statistical agreement among multiple analytical methods [1].

Information Mass Fraction Values: Information mass fraction values are provided in Table 11 for carbon, hydrogen, and nitrogen. An information value is considered to be a value that will be of use to the SRM user, but insufficient information is available to assess the uncertainty associated with the value [1]. Information values cannot be used to establish metrological traceability.

Expiration of Certification: The certification of **SRM 1941b** is valid, within the measurement uncertainty specified, until **01 October 2020**, provided the SRM is handled and stored in accordance with the instructions given in this certificate (see "Instructions for Handling, Storage, and Use"). This certification is nullified if the SRM is damaged, contaminated, or otherwise modified.

Maintenance of SRM Certification: NIST will monitor this SRM over the period of its certification. If substantive technical changes occur that affect the certification before the expiration of this certificate, NIST will notify the purchaser. Registration (see attached sheet or register online) will facilitate notification.

Coordination of the technical measurements leading to the certification of this material was under the leadership of M.M. Schantz and S.A. Wise of the NIST Chemical Sciences Division.

Analytical measurements for the certification of SRM 1941b were performed at NIST by J.R. Kucklick, B.J. Porter, D.L. Poster, M.M. Schantz, P. Schubert, S. Tutschku, and L.L. Yu of the NIST Chemical Sciences Division.

Carlos A. Gonzalez, Chief
Chemical Sciences Division

Measurements for TOC were provided by a commercial laboratory and T.L. Wade of the Geochemical and Environmental Research Group, Texas A&M University (College Station, TX). The carbon, hydrogen, and nitrogen data were provided by a commercial laboratory. Results for the PAHs, PCBs, and chlorinated pesticides from 38 laboratories (see Appendix A) that participated in an interlaboratory comparison exercise coordinated by NIST were used. Results for the alkylated PAH groups, hopanes, and steranes from 33 laboratories (see Appendix B) that participated in another interlaboratory comparison exercise coordinated by NIST were also used.

Collection and preparation of SRM 1941b were performed by M.P. Cronise and C.N. Fales of the NIST Office of Reference Materials and B.J. Porter and M.M. Schantz of the NIST Chemical Sciences Division. The sediment material was collected with the assistance of G.G. Lauenstein, J. Collier, and J. Lewis (National Oceanic and Atmospheric Administration, Silver Spring, MD).

Consultation on the statistical design of the experimental work and evaluation of the data were provided by S.D. Leigh and J.H. Yen of the NIST Statistical Engineering Division.

Support aspects involved in the issuance of this SRM were coordinated through the NIST Office of Reference Materials.

INSTRUCTIONS FOR HANDLING, STORAGE, AND USE

Handling: This material is naturally occurring marine sediment from an urban area and may contain constituents of unknown toxicities; therefore, caution and care should be exercised during its handling and use.

Storage: SRM 1941b must be stored in its original bottle at temperatures less than 30 °C and away from direct sunlight.

Use: Prior to removal of subsamples for analysis, the contents of the bottle should be mixed. The mass fractions of constituents in SRM 1941b are reported on a dry-mass basis. The SRM, as received, contains a mass fraction of approximately 2.4 % moisture (see "Conversion to Dry-Mass Basis"). The sediment sample should be dried to a constant mass before weighing for analysis; or a separate subsample of the sediment should be removed from the bottle at the time of analysis and dried to determine the mass fraction on a dry-mass basis. If the constituents of interest are volatile, then the moisture must be determined with a separate subsample.

PREPARATION AND ANALYSIS⁽¹⁾

Sample Collection and Preparation: The sediment used to prepare this SRM was collected from the Chesapeake Bay at the mouth of the Baltimore (MD) Harbor near the Francis Scott Key Bridge (39°12.3'N and 76°31.4'W). This location is very near the site where SRM 1941 and SRM 1941a were collected. The sediment was collected using a Kynar-coated modified Van Veen-type grab sampler. A total of approximately 3300 kg of wet sediment was collected from the site. The sediment was freeze-dried, sieved at 150 µm (100 % passing), homogenized in a cone blender, radiation sterilized (⁶⁰Co), and then packaged in screw-capped amber glass bottles each containing approximately 50 g.

Conversion to Dry-Mass Basis: The results for the constituents in SRM 1941b are reported on a dry-mass basis; however, the material "as received" contains residual moisture. The amount of moisture in SRM 1941b was determined by measuring the mass loss after freeze-drying subsamples of 1.1 g to 1.3 g for four days at 1 Pa with a -10 °C shelf temperature and a -50 °C condenser temperature. The moisture content in SRM 1941b at the time of the certification analyses was 2.39 % ± 0.08 % (95 % confidence level). Analytical results for the organic constituents were determined on an as-received basis and then converted to a dry-mass basis by dividing by the conversion factor of 0.9761 (gram dry mass per gram as-received mass).

Polycyclic Aromatic Hydrocarbons: The general approach used for the value assignment of the PAHs in SRM 1941b was similar to that reported in detail elsewhere [2]. The approach consisted of combining results from analyses using various combinations of different extraction techniques and solvents, clean-up/isolation procedures, and chromatographic separation and detection techniques: Soxhlet extraction and pressurized-fluid extraction (PFE) using dichloromethane (DCM) or a hexane/acetone mixture, cleanup of the extracts using solid-phase extraction (SPE) or normal-phase liquid chromatography (LC), followed by analysis using the following techniques: (1) reversed-phase liquid chromatography with fluorescence detection (LC-FL) analysis of the total PAH fraction, (2) reversed-phase

⁽¹⁾ Certain commercial equipment, instruments or materials are identified in this certificate to adequately specify the experimental procedure. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology.

LC-FL analysis of isomeric PAH fractions isolated by normal-phase LC (i.e., multidimensional LC), (3) gas chromatography/mass spectrometry (GC/MS) analysis of the PAH fraction on three stationary phases of different selectivity, i.e., a 5 % (all column compositions are given as mole fractions in %) phenyl-substituted methylpolysiloxane phase, a 50 % phenyl-substituted methylpolysiloxane phase, and a relatively non-polar proprietary phase.

Three sets of GC/MS results, designated as GC/MS (I), GC/MS (II), and GC/MS (III), were obtained using three columns with different selectivities for the separation of PAHs. For GC/MS (I) analyses, duplicate subsamples of approximately 1 g from ten bottles of SRM 1941b were extracted using PFE with DCM. Copper powder was added to the extract to remove elemental sulfur. The concentrated extract was passed through an aminopropyl SPE cartridge and eluted with 2 % DCM in hexane (all solvent concentrations are given as volume fractions in %). The processed extract was then analyzed by GC/MS using a 0.25 mm i.d. × 60 m fused silica capillary column with a 5 % phenyl-substituted methylpolysiloxane phase (0.25 μm film thickness; DB-5 MS, J&W Scientific, Folsom, CA). The GC/MS (II) analyses were performed using 5 g subsamples from six bottles of SRM 1941b. These samples were extracted using PFE with DCM. The high molecular mass compounds were removed from the extracts using size exclusion chromatography (SEC) with a preparative-scale divinylbenzene-polystyrene column (10 μm particle size with 10 nm diameter pores), and the sulfur was removed from the extracts by adding copper powder. The concentrated extract was passed through an aminopropyl SPE cartridge and eluted with 10 % DCM in hexane. The analysis was by GC/MS using a 0.25 mm i.d. × 60 m fused silica capillary column with a 50 % phenyl-substituted methylpolysiloxane phase (0.25 μm film thickness; DB-17 MS, J&W Scientific). For the GC/MS (III), 9 g subsamples from six bottles of SRM 1941b were Soxhlet-extracted for 18 h with 250 mL of a mixture of 50 % hexane/50 % acetone. Copper powder was added to the extract to remove elemental sulfur, and the concentrated extract was passed through a silica SPE cartridge and eluted with 10 % DCM in hexane. The processed extract was then analyzed by GC/MS using a 0.25 mm i.d. × 60 m fused silica capillary column with a relatively non-polar proprietary phase (0.25 μm film thickness; DB-XLB, J&W Scientific).

Two sets of LC-FL results, designated as LC-FL (total) and LC-FL (isomer), were used in the certification process. For the LC-FL (total), subsamples of approximately 1 g from six bottles of SRM 1941b were extracted using PFE with a mixture of 50 % hexane/50 % acetone. The extracts were concentrated and then processed through an aminopropylsilane SPE cartridge using 2 % DCM in hexane to obtain the total PAH fraction. For the LC-FL (isomer), a 5 g subsample from the six bottles was extracted using PFE with DCM and processed through an aminopropylsilane SPE cartridge using 10 % DCM in hexane; the PAH fraction was then fractionated further on a semi-preparative aminopropylsilane column (μBondapak NH₂, 9 mm i.d. × 30 cm, Waters Associates, Milford, MA) to isolate isomeric PAH fractions as described previously [3–6]. The total PAH fraction and the isomeric PAH fractions were analyzed using a 5 μm particle-size polymeric octadecylsilane (C₁₈) column (4.6 mm i.d. × 25 cm, Hypersil-PAH, Keystone Scientific, Inc., Bellefonte, PA) with wavelength-programmed fluorescence detection [4,5].

For the GC/MS and LC-FL measurements described above, selected perdeuterated PAHs were added to the sediment prior to solvent extraction for use as internal standards for quantification purposes.

In addition to the analyses performed at NIST, SRM 1941b was used in an interlaboratory comparison exercise in 1999 as part of the NIST Intercomparison Exercise Program for Organic Contaminants in the Marine Environment [7]. Results from 38 laboratories that participated in this exercise were used as the sixth data set in the determination of the certified values for PAHs in SRM 1941b. The laboratories participating in this exercise used the analytical procedures routinely used in their laboratories to measure the analytes of interest.

Homogeneity Assessment for PAHs: The homogeneity of SRM 1941b was assessed by analyzing duplicate samples of approximately 1 g from ten bottles selected by stratified random sampling. Samples were extracted, processed, and analyzed as described above for GC/MS (I). No statistically significant differences among bottles were observed for the PAHs at this sample size.

PAH Isomers of Molecular Mass 300 and 302: For the determination of the molecular mass 300 and 302 isomers, three subsamples of approximately 5 g each were extracted using PFE with DCM. The extracts were then concentrated with a solvent change to hexane and passed through an aminopropyl SPE cartridge and eluted with 10 % DCM in hexane. The processed extract was then analyzed by GC/MS using a 0.25 mm i.d. × 60 m fused silica capillary column with a 50 % phenyl-substituted methylpolysiloxane phase (0.25 μm film thickness; DB-17MS, J&W Scientific). Perdeuterated dibenzo[*a,i*]pyrene was added to the sediment prior to extraction for use as an internal standard [8].

PCBs and Chlorinated Pesticides: The general approach used for the determination of PCBs and chlorinated pesticides in SRM 1941b consisted of combining results from analyses using various combinations of different extraction techniques and solvents, cleanup/isolation procedures, and chromatographic separation and detection techniques. Techniques and solvents included Soxhlet extraction and PFE using DCM or a hexane/acetone mixture.

clean-up/isolation using SPE or LC, followed by analysis using GC/MS and gas chromatography with electron capture detection (GC-ECD) on two columns with different selectivity for the separation of PCBs and chlorinated pesticides. The analytical methods are described in detail elsewhere [2].

Six sets of results were obtained and designated as GC-ECD (I) A and B, GC/MS (I) A and B, GC/MS (II), and Interlaboratory Comparison Exercise. For the GC-ECD (I) analyses, approximately 10 g subsamples from six bottles of SRM 1941b were extracted using PFE with DCM. Copper powder was added to the extract to remove elemental sulfur, and SEC, as described above, was used to remove the high molecular mass compounds. The concentrated extract was then fractionated on a semi-preparative aminopropylsilane column to isolate two fractions containing: (1) the PCBs and lower-polarity pesticides and (2) the more polar pesticides. GC-ECD analyses of the two fractions were performed on two columns of different selectivities for PCB separations: 0.25 mm × 60 m fused silica capillary column with a 5 % phenyl-substituted methylpolysiloxane phase (0.25 μm film thickness; DB-5, J&W Scientific), and a 0.25 mm × 60 m fused silica capillary column with a non-polar proprietary phase (0.25 μm film thickness; DB-XLB, J&W Scientific). The results from the 5 % phenyl phase are designated as GC-ECD (IA) and the results from the proprietary phase are designated as GC-ECD (IB). For the GC-ECD analyses, two PCB congeners that are not significantly present in the sediment extract (PCB 103 and PCB 198 [9,10]) and endosulfan I-*d*₄, 4,4'-DDE-*d*₈, 4,4'-DD-*d*₈, and 4,4'-DDT-*d*₈ were added to the sediment prior to extraction for use as internal standards for quantification purposes.

Two sets of results were obtained by GC/MS. For GC/MS (I), approximately 9 g subsamples from six bottles were Soxhlet- extracted with a mixture of 50 % hexane/50 % acetone for approximately 18 h. Copper powder was added to the extract to remove elemental sulfur, and the concentrated extract was passed through a silica SPE cartridge and eluted with 10 % DCM in hexane. The processed extract was then analyzed by GC/MS with two ionization modes, electron impact (EI) and negative ion chemical ionization (NICI). The GC/MS EI method, GC/MS (IA), used a 0.25 mm i.d. × 60 m fused silica capillary column with a relatively non-polar proprietary phase (0.25 μm film thickness; DB-XLB, J&W Scientific). The GC/MS NICI method, GC/MS (IB), used a 0.25 mm i.d. × 60 m fused silica capillary column with a 5 % phenyl-substituted methylpolysiloxane phase (0.25 μm film thickness; DB-5MS, J&W Scientific). The GC/MS (II) results were obtained in the same manner as the GC/MS (IA) analyses except that three subsamples were Soxhlet-extracted with DCM for approximately 18 h. For the GC/MS analyses, selected carbon-13 labeled PCB congeners and chlorinated pesticides were added to the sediment prior to extraction for use as internal standards for quantification purposes.

In addition to the analyses performed at NIST, SRM 1941b was used in an interlaboratory comparison exercise in 1999 as part of the NIST Intercomparison Exercise Program for Organic Contaminants in the Marine Environment [7]. Results from 38 laboratories that participated in this exercise were used as the sixth data set in the determination of the certified values for PCB congeners and chlorinated pesticides in SRM 1941b. The laboratories participating in this exercise used the analytical procedures routinely used in their laboratories to measure the analytes of interest.

The reference value for PCB 77 was determined from a separate fraction. The samples were extracted and processed as for GC-ECD (I) above. The first (PCB and lower-polarity pesticide) fraction from the semi-preparative aminopropylsilane column was further fractionated using a Cosmosil PYE (pyrenylethyl group bonded) column (5 μm particle size, 4.6 mm i.d. × 25 cm; Phenomenex, Torrance, CA) [11]. Three fractions were collected: the first fraction contained the pesticides and multi-*ortho* PCBs, the second fraction contained the polychlorinated naphthalenes, non-*ortho* PCB congeners, and some mono-*ortho* PCB congeners, and the third fraction removed the residual planar compounds from the column. The second fraction was analyzed by GC/MS NICI using the same column as GC/MS (IB) above. Carbon-13 labeled PCB 77 was used as an internal standard for quantification purposes.

Alkylated PAH Groups, Hopanes, and Steranes: SRM 1941b was used in an interlaboratory comparison exercise in 2011 [12]. Results from 33 laboratories that participated in this exercise were used in the determination of the reference values for alkylated PAH groups, hopanes, and steranes in SRM 1941b. Note that not all laboratories returned data for each analyte. The laboratories participating in this exercise used the analytical procedures routinely used in their laboratories to measure the analytes of interest. For the alkylated PAHs, the majority of the laboratories (>90 %) used the parent PAH for determination of the response factor for the corresponding alkylated group.

Total Organic Carbon (TOC): Two laboratories provided results for TOC using similar procedures. Briefly, subsamples of approximately 200 mg were reacted with 6 mol/L hydrochloric acid and rinsed with deionized water prior to combustion in a gas fusion furnace. The carbon monoxide and carbon dioxide produced were measured and compared to a blank for calculation of the percent TOC. Each laboratory analyzed subsamples from three bottles of SRM 1941b. One of the laboratories also analyzed three subsamples from three bottles of SRM 1941b for carbon, hydrogen, and nitrogen.

Table 1. Certified Mass Fraction Values for PAHs in SRM 1941b

PAHs	Mass Fractions ^(a) ($\mu\text{g}/\text{kg}$)
Naphthalene ^(b,c,d,e,f,g)	848 \pm 95 ^(h)
Fluorene ^(b,c,d,e,f,g)	85 \pm 15 ^(h)
Phenanthrene ^(b,c,d,e,f,g)	406 \pm 44 ^(h)
Anthracene ^(b,c,d,e,f,g)	184 \pm 18 ^(h)
3-Methylphenanthrene ^(b,c,d)	105 \pm 13 ^(h)
2-Methylphenanthrene ^(b,c,d)	128 \pm 14 ^(h)
1-Methylphenanthrene ^(b,c,d,g)	73.2 \pm 5.9 ^(h)
Fluoranthene ^(b,c,d,e,f,g)	651 \pm 50 ^(h)
Pyrene ^(b,c,d,e,f,g)	581 \pm 39 ^(h)
Benz[<i>a</i>]anthracene ^(b,c,d,e,f,g)	335 \pm 25 ^(h)
Chrysene ^(d,f)	291 \pm 31 ^(h)
Triphenylene ^(d,f)	108 \pm 5 ⁽ⁱ⁾
Benzo[<i>b</i>]fluoranthene ^(c,e)	453 \pm 21 ^(h)
Benzo[<i>k</i>]fluoranthene ^(b,c,d,e)	225 \pm 18 ^(h)
Benzo[<i>e</i>]pyrene ^(b,c,d,g)	325 \pm 25 ^(h)
Benzo[<i>a</i>]pyrene ^(b,c,d,f,g)	358 \pm 17 ^(h)
Perylene ^(b,c,d,f,g)	397 \pm 45 ^(h)
Benzo[<i>ghi</i>]perylene ^(b,c,d,f,g)	307 \pm 45 ^(h)
Indeno[1,2,3- <i>cd</i>]pyrene ^(b,c,d,f,g)	341 \pm 57 ^(h)
Dibenz[<i>a,j</i>]anthracene ^(b,c,d,f)	48.9 \pm 4.6 ^(h)
Dibenz[<i>a,c</i>]anthracene ^(c,f)	36.7 \pm 5.2 ^(h)
Dibenz[<i>a,h</i>]anthracene ^(c,f)	53 \pm 10 ^(h)
Benzo[<i>b</i>]chrysene ^(b,c,d,f)	53 \pm 12 ^(h)
Picene ^(b,c,d)	46.6 \pm 4.7 ^(h)

^(a) Mass fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(b) GC/MS (I) on 5 % phenyl-substituted methylpolysiloxane phase after PFE with DCM.

^(c) GC/MS (II) on 50 % phenyl-substituted methylpolysiloxane phase after PFE with DCM.

^(d) GC/MS (III) on a relatively non-polar proprietary phase after Soxhlet extraction with 50 % hexane/50 % acetone mixture.

^(e) LC-FL (total) of total PAH fraction after PFE with DCM.

^(f) LC-FL (isomer) of isomeric PAH fractions after PFE with DCM.

^(g) 1999 Interlaboratory Comparison Study [7] with 21 to 29 laboratories submitting data for each PAH.

^(h) Certified values are weighted means of the results from two to six analytical methods [13]. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2 (approximately 95 % confidence), calculated by combining a between-method variance incorporating inter-method bias with a pooled within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the constituent listed and the values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

⁽ⁱ⁾ The certified value is an unweighted mean of the results from two analytical methods. The uncertainty listed with the value is an expanded uncertainty about the mean, with coverage factor 2, calculated by combining a between-method variance [16] with a pooled, within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the constituent listed and the value is metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

Table 2. Certified Mass Fraction Values for PCB Congeners^(a) in SRM 1941b

PCB Congeners		Mass Fractions ^(b) ($\mu\text{g}/\text{kg}$)
PCB	8 (2,4'-Dichlorobiphenyl) ^(c,d,e,f,g)	1.65 \pm 0.19 ^(h)
PCB	18 (2,2',5-Trichlorobiphenyl) ^(c,d,e,f,g)	2.39 \pm 0.29 ^(h)
PCB	28 (2,4,4'-Trichlorobiphenyl) ^(c,d,e,f,g)	4.52 \pm 0.57 ^(h)
PCB	31 (2,4',5-Trichlorobiphenyl) ^(c,e,f)	3.18 \pm 0.41 ^(h)
PCB	44 (2,2',3,5'-Tetrachlorobiphenyl) ^(c,d,e,f,g)	3.85 \pm 0.20 ⁽ⁱ⁾
PCB	49 (2,2',4,5'-Tetrachlorobiphenyl) ^(c,d,e,f)	4.34 \pm 0.28 ⁽ⁱ⁾
PCB	52 (2,2',5,5'-Tetrachlorobiphenyl) ^(c,d,e,f,g)	5.24 \pm 0.28 ⁽ⁱ⁾
PCB	66 (2,3',4,4'-Tetrachlorobiphenyl) ^(c,e,f,g,j)	4.96 \pm 0.53 ⁽ⁱ⁾
PCB	87 (2,2',3,4,5'-Pentachlorobiphenyl) ^(c,d,f,j)	1.14 \pm 0.16 ^(h)
PCB	95 (2,2',3,5',6-Pentachlorobiphenyl) ^(c,e,f,g)	3.93 \pm 0.62 ⁽ⁱ⁾
PCB	99 (2,2',4,4',5-Pentachlorobiphenyl) ^(c,d,e,f,g)	2.90 \pm 0.36 ⁽ⁱ⁾
PCB	101 (2,2',4,5,5'-Pentachlorobiphenyl) ^(c,e,f,g,j)	5.11 \pm 0.34 ⁽ⁱ⁾
PCB	105 (2,3,3',4,4'-Pentachlorobiphenyl) ^(c,d,e,f,g,j)	1.43 \pm 0.10 ⁽ⁱ⁾
PCB	110 (2,3,3',4',6-Pentachlorobiphenyl) ^(c,e,f,j)	4.62 \pm 0.36 ⁽ⁱ⁾
PCB	118 (2,3',4,4',5-Pentachlorobiphenyl) ^(c,d,e,f,g,j)	4.23 \pm 0.19 ⁽ⁱ⁾
PCB	128 (2,2',3,3',4,4'-Hexachlorobiphenyl) ^(c,d,e,f,g,j)	0.696 \pm 0.044 ⁽ⁱ⁾
PCB	138 (2,2',3,4,4',5'-Hexachlorobiphenyl) ^(c,e,f,j)	3.60 \pm 0.28 ⁽ⁱ⁾
PCB	149 (2,2',3,4',5,6-Hexachlorobiphenyl) ^(c,d,e,j)	4.35 \pm 0.26 ^(h)
PCB	153 (2,2',4,4',5,5'-Hexachlorobiphenyl) ^(c,d,e,f,g,j)	5.47 \pm 0.32 ⁽ⁱ⁾
PCB	156 (2,3,3',4,4',5-Hexachlorobiphenyl) ^(c,d,e,f,j)	0.507 \pm 0.090 ^(h)
PCB	170 (2,2',3,3',4,4',5-Heptachlorobiphenyl) ^(c,d,e,f,g,j)	1.35 \pm 0.09 ⁽ⁱ⁾
PCB	180 (2,2',3,4,4',5,5'-Heptachlorobiphenyl) ^(c,d,e,f,g,j)	3.24 \pm 0.51 ⁽ⁱ⁾
PCB	183 (2,2',3,4,4',5,6-Heptachlorobiphenyl) ^(c,d,e,j)	0.979 \pm 0.087 ^(h)
PCB	187 (2,2',3,4',5,5',6-Heptachlorobiphenyl) ^(c,d,e,f,g,j)	2.17 \pm 0.22 ⁽ⁱ⁾
PCB	194 (2,2',3,3',4,4',5,5'-Octachlorobiphenyl) ^(c,d,e,j)	1.04 \pm 0.06 ^(h)
PCB	195 (2,2',3,3',4,4',5,6-Octachlorobiphenyl) ^(c,e,g,j)	0.645 \pm 0.060 ⁽ⁱ⁾
PCB	201 (2,2',3,3',4,5',6'-Octachlorobiphenyl) ^(c,e,j)	0.777 \pm 0.034 ^(h)
PCB	206 (2,2',3,3',4,4',5,5',6-Nonachlorobiphenyl) ^(c,e,f,g,j)	2.42 \pm 0.19 ⁽ⁱ⁾
PCB	209 Decachlorobiphenyl ^(c,d,e,f,g,j)	4.86 \pm 0.45 ⁽ⁱ⁾

^(a) PCB congeners are numbered according to the scheme proposed by Ballschmiter and Zell [9] and later revised by Schulte and Malisch [10] to conform to IUPAC rules, except PCB 201. Under the Ballschmiter and Zell numbering system, the IUPAC PCB 201 is listed as PCB 200.

^(b) Mass fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(c) GC/MS (IA) on a relatively non-polar proprietary phase after Soxhlet extraction with 50 % hexane/50 % acetone mixture.

^(d) GC-ECD (IA) on 5 % phenyl-substituted methylpolysiloxane phase after PFE extraction with DCM.

^(e) GC-ECD (IB) on a relatively non-polar proprietary phase; same extracts analyzed as in GC-ECD (IA).

^(f) GC/MS (II) on a relatively non-polar proprietary phase after Soxhlet extraction with DCM.

^(g) 1999 Interlaboratory Comparison Study [7] with 13 to 31 laboratories submitting data for each PCB congener.

^(h) Certified values are unweighted means of the results from three to five analytical methods. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2, calculated by combining a between-method variance [16] with a pooled, within method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the constituent listed and the values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

⁽ⁱ⁾ Certified values are weighted means of the results from three to six analytical methods [13]. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2 (approximately 95 % confidence), calculated by combining a between-method variance incorporating inter-method bias with a pooled within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the constituent listed and the values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

^(j) GC/MS (IB) on 5 % phenyl-substituted methylpolysiloxane phase; same extracts analyzed as in GC/MS (IA).

Table 3. Certified Mass Fraction Values for Chlorinated Pesticides in SRM 1941b

Chlorinated Pesticides	Mass Fractions ^(a) ($\mu\text{g}/\text{kg}$)
Hexachlorobenzene ^(b,c,d,e)	5.83 \pm 0.38 ^(f)
<i>cis</i> -Chlordane ^(b,c,d,e,g)	0.85 \pm 0.11 ^(h)
<i>trans</i> -Chlordane ^(b,c,e)	0.566 \pm 0.093 ^(f)
<i>cis</i> -Nonachlor ^(b,e,g)	0.378 \pm 0.053 ^(h)
<i>trans</i> -Nonachlor ^(b,c,d,e,g)	0.438 \pm 0.073 ^(f)
4,4'-DDE ^(b,d,e,g)	3.22 \pm 0.28 ^(h)
4,4'-DDD ^(b,d,e,g)	4.66 \pm 0.46 ^(h)

^(a) Mass fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(b) GC/MS (IA) on a relatively non-polar proprietary phase after Soxhlet extraction with 50 % hexane/50 % acetone mixture.

^(c) GC/MS (IB) on 5 % phenyl-substituted methylpolysiloxane phase; same extracts analyzed as in GC/MS (IA).

^(d) GC/MS (II) on a relatively non-polar proprietary phase after Soxhlet extraction with DCM.

^(e) 1999 Interlaboratory Comparison Study [7] with 13 to 31 laboratories submitting data for each pesticide.

^(f) Certified values are unweighted means of the results from three to five analytical methods. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2, calculated by combining a between-method variance [16] with a pooled, within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the constituent listed and the values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

^(g) GC-ECD (IA) on 5 % phenyl-substituted methylpolysiloxane phase after PFE extraction with DCM.

^(h) Certified values are weighted means of the results from three to five analytical methods [13]. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2 (approximately 95 % confidence), calculated by combining a between-method variance incorporating inter-method bias with a pooled within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the chlorinated pesticides listed and the values listed are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

Table 4. Reference Mass Fraction Values for PAHs in SRM 1941b

PAHs	Mass Fractions ^(a)		
	(μg/kg)		
1-Methylnaphthalene ^(b,c,d,e)	127	±	14 ^(f)
2-Methylnaphthalene ^(b,c,d,e)	276	±	53 ^(f)
2,6-Dimethylnaphthalene ^(b,c,d,e)	75.9	±	4.5 ^(f)
2,3,5-Trimethylnaphthalene ^(b,c,d,e)	25.5	±	5.1 ^(f)
Biphenyl ^(b,c,d,e)	74.0	±	8.0 ^(f)
Acenaphthylene ^(b,c,d,e)	53.3	±	6.4 ^(f)
Acenaphthene ^(b,c,d,e)	38.4	±	5.2 ^(f)
9-Methylphenanthrene ^(c)	63.5	±	2.5 ^(g)
4-Methylphenanthrene and 9-Methylphenanthrene ^(b,d)	80.1	±	4.8 ^(f)
2-Methylanthracene ^(c,d)	36	±	15 ^(f)
8-Methylfluoranthene ^(b)	49.5	±	2.7 ^(g)
7-Methylfluoranthene ^(b)	45.4	±	1.5 ^(g)
1-Methylfluoranthene ^(b)	42.4	±	2.1 ^(g)
3-Methylfluoranthene ^(b)	28.8	±	1.3 ^(g)
2-Methylpyrene ^(b)	78.7	±	4.0 ^(g)
4-Methylpyrene ^(b)	66.4	±	2.6 ^(g)
1-Methylpyrene ^(b)	52.5	±	2.3 ^(g)
Acephenanthrene ^(d)	30.5	±	1.9 ^(g)
Benzo[<i>c</i>]phenanthrene ^(b,c,d)	58	±	15 ^(f)
Benzo[<i>a</i>]fluoranthene ^(b,c,d)	73	±	18 ^(f)
Benzo[<i>j</i>]fluoranthene ^(c)	217	±	5 ^(g)
Indeno[1,2,3- <i>cd</i>]fluoranthene ^(d)	9.63	±	0.34 ^(g)
Pentaphene ^(d)	25.3	±	1.0 ^(g)

^(a) Mass fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(b) GC/MS (I) on 5 % phenyl-substituted methylpolysiloxane phase after PFE with DCM.

^(c) GC/MS (II) on 50 % phenyl-substituted methylpolysiloxane phase after PFE with DCM.

^(d) GC/MS (III) on a relatively non-polar proprietary phase after Soxhlet extraction with 50 % hexane/50 % acetone mixture.

^(e) 1999 Interlaboratory Comparison Study [7] with 14 to 26 laboratories submitting data for each PAH.

^(f) Reference values are weighted means of the results from two to four analytical methods [13]. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2 (approximately 95 % confidence), calculated by combining a between-method variance incorporating inter-method bias with a pooled within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of PAHs listed as determined by the methods indicated. The values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

^(g) Reference values are the means of results obtained by NIST using one analytical technique. The expanded uncertainty, U , is calculated as $U = k u_c$, where u_c is one standard deviation of the analyte mean, and the coverage factor, k , is determined from the Student's t -distribution for the associated degrees of freedom (19 for footnote b and 5 for footnotes c and d) and 95 % confidence level for each analyte. The measurand is the total mass fraction of the PAHs listed as determined by the method indicated. The values listed are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

Table 5. Reference Mass Fraction Values for PAHs of Molecular Mass 300 and 302 in SRM 1941b

PAHs of Molecular Mass 300 and 302	Mass Fractions ^(a,b,c) ($\mu\text{g}/\text{kg}$)
Coronene	72.6 \pm 4.7
Dibenzo[<i>b,e</i>]fluoranthene	10.3 \pm 0.3
Naphtho[1,2- <i>b</i>]fluoranthene	91.0 \pm 3.1
Naphtho[1,2- <i>k</i>]fluoranthene and Naphtho[2,3- <i>j</i>]fluoranthene	79.8 \pm 2.5
Naphtho[2,3- <i>b</i>]fluoranthene	23.5 \pm 0.3
Dibenzo[<i>b,k</i>]fluoranthene	95.6 \pm 3.1
Dibenzo[<i>a,k</i>]fluoranthene	26.6 \pm 0.4
Dibenzo[<i>j,l</i>]fluoranthene	63.8 \pm 1.8
Dibenzo[<i>a,l</i>]pyrene	11.1 \pm 1.0
Naphtho[2,3- <i>k</i>]fluoranthene	10.7 \pm 0.6
Naphtho[1,2- <i>a</i>]pyrene	16.7 \pm 1.4
Naphtho[2,3- <i>e</i>]pyrene	33.2 \pm 2.3
Dibenzo[<i>a,e</i>]pyrene	76.1 \pm 3.6
Naphtho[2,1- <i>a</i>]pyrene	59.2 \pm 1.8
Dibenzo[<i>e,i</i>]pyrene	35.0 \pm 2.4
Naphtho[2,3- <i>a</i>]pyrene	16.5 \pm 0.6
Benzo[<i>b</i>]perylene	38.2 \pm 1.2
Dibenzo[<i>a,i</i>]pyrene	25.5 \pm 1.0
Dibenzo[<i>a,h</i>]pyrene	6.94 \pm 0.29

^(a) Mass fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(b) Reference values are the means of results obtained by NIST using one analytical technique. The expanded uncertainty, U , is calculated as $U = ku_c$, where u_c is one standard deviation of the analyte mean, and the coverage factor, k , is determined from the Student's t -distribution for two degrees of freedom and 95 % confidence level for each analyte. The measurand is the total mass fraction of the constituent listed as determined by the method indicated. The values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

^(c) GC/MS on 50 % phenyl-substituted methylpolysiloxane phase after PFE with DCM [8].

Table 6. Reference Mass Fraction Values for PCB Congeners^(a) in SRM 1941b

PCB Congeners			Mass Fractions ^(b,c) ($\mu\text{g}/\text{kg}$)		
PCB	45	(2,2',3,6-Tetrachlorobiphenyl) ^(d,e)	0.73	±	0.12
PCB	56	(2,3,3',4'-Tetrachlorobiphenyl) ^(d,f,g)	1.21	±	0.11
PCB	63	(2,3,4',5-Tetrachlorobiphenyl) ^(e,f,g)	0.213	±	0.040
PCB	70	(2,3',4',5-Tetrachlorobiphenyl) ^(e,f,g)	4.99	±	0.29
PCB	74	(2,4,4',5-Tetrachlorobiphenyl) ^(e,f,g)	2.04	±	0.15
PCB	77	(3,3',4,4'-Tetrachlorobiphenyl) ^(h)	0.31	±	0.03
PCB	107	(2,3,3',4',5-Pentachlorobiphenyl) ^(d,e,f,g)	0.628	±	0.028
PCB	132	(2,2',3,3',4,6'-Hexachlorobiphenyl) ^(d,f,g)	1.28	±	0.27
PCB	146	(2,2',3,4',5,5'-Hexachlorobiphenyl) ^(e,f,g)	1.22	±	0.12
PCB	158	(2,3,3',4,4',6-Hexachlorobiphenyl) ^(d,e,f,g)	0.65	±	0.15
PCB	163	(2,3,3',4',5,6-Hexachlorobiphenyl) ^(e,f,g)	1.28	±	0.06
PCB	174	(2,2',3,3',4,5,6'-Heptachlorobiphenyl) ^(d,e,f,g)	1.51	±	0.39
PCB	193	(2,3,3',4',5,5',6-Heptachlorobiphenyl) ^(d,e,f,g)	0.292	±	0.075

^(a) PCB congeners are numbered according to the scheme proposed by Ballschmiter and Zell [9] and later revised by Schulte and Malisch [10] to conform with IUPAC rules, except PCB 107. Under the Ballschmiter and Zell numbering system, the IUPAC PCB 107 is listed as PCB 108.

^(b) Mass fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(c) For these PCB congeners except PCB 77, the reference values are unweighted means of the results from two to four analytical methods. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2, calculated by combining a between-method variance [16] with a pooled within-method variance following the ISO/JCGM Guide [14,15]. For PCB 77, the reference value is the mean of results obtained by NIST using one analytical technique. The expanded uncertainty, U , is calculated as $U = k u_c$, where u_c is one standard deviation of the analyte mean, and the coverage factor, k , is determined from the Student's t -distribution corresponding to two degrees of freedom and 95 % confidence level for PCB 77. The measurand is the total mass fraction of the PCB Congeners listed as determined by the method or methods indicated. The values listed are metrologically traceable to the SI unit of mass, expressed as microgram per kilogram on a dry-mass basis.

^(d) GC-ECD (IA) on 5 % phenyl-substituted methylpolysiloxane phase after PFE extraction with DCM.

^(e) GC-ECD (IB) on a relatively non-polar proprietary phase; same extracts analyzed as in GC-ECD (IA).

^(f) GC/MS (IA) on a relatively non-polar proprietary phase after Soxhlet extraction with 50 % hexane/50 % acetone mixture.

^(g) GC/MS (IB) on 5 % phenyl-substituted methylpolysiloxane phase; same extracts analyzed as in GC/MS (IA).

^(h) GC/MS NICI on a 5 % phenyl-substituted methylpolysiloxane phase; same extracts analyzed as in GC-ECD (I) fractionated using a PYE column.

Table 7. Reference Mass Fraction Values for Selected Chlorinated Pesticides in SRM 1941b

Chlorinated Pesticides	Mass Fractions ^(a,b) ($\mu\text{g}/\text{kg}$)
2,4'-DDE ^(c,d)	0.38 \pm 0.12
4,4'-DDT ^(e,f)	1.12 \pm 0.42

^(a) Mass Fractions reported on dry-mass basis; material as received contains approximately 2.4 % moisture.

^(b) The reference values are unweighted means of the results from two analytical methods. The uncertainty listed with each value is an expanded uncertainty about the mean, with coverage factor 2, calculated by combining a between-method variance [16] with a pooled, within-method variance following the ISO/JCGM Guide [14,15]. The measurand is the total mass fraction of the chlorinated pesticides listed as determined by the methods indicated. The values listed are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.

^(c) GC/MS (IB) on 5 % phenyl-substituted methylpolysiloxane phase; same extracts analyzed as in GC/MS (IA).

^(d) GC-ECD (IB) on a relatively non-polar proprietary phase; same extracts analyzed as in GC-ECD (IA).

^(e) GC/MS (II) on a relatively non-polar proprietary phase after Soxhlet extraction with DCM.

^(f) 1999 Interlaboratory Comparison Study [7] with 10 laboratories submitting data for 4,4'-DDT.

Table 8. Reference Mass Fraction Values for Alkylated PAH Groups in SRM 1941b

Alkylated PAH Group	Mass Fraction ^(a,b) ($\mu\text{g}/\text{kg}$)
C2-decalins	18 \pm 5
C4-decalins	41 \pm 4
C2-naphthalenes	187 \pm 53
C3-naphthalenes	158 \pm 42
C1-benzothiophenes	25 \pm 14
C2-benzothiophenes	20 \pm 11
C3-benzothiophenes	22 \pm 13
C4-benzothiophenes	18 \pm 5
C1-fluorenes	57 \pm 18
C2-fluorenes	122 \pm 43
C3-fluorenes	128 \pm 31
C1-phenanthrenes/anthracenes	313 \pm 99
C2-phenanthrenes/anthracenes	247 \pm 62
C3-phenanthrenes/anthracenes	165 \pm 46
C4-phenanthrenes/anthracenes	87 \pm 36
C1-dibenzothiophenes	54 \pm 13
C2-dibenzothiophenes	91 \pm 18
C3-dibenzothiophenes	84 \pm 15
C4-dibenzothiophenes	57 \pm 13
C1-fluoranthenes/pyrenes	252 \pm 48
C2-fluoranthenes/pyrenes	205 \pm 38
C3-fluoranthenes/pyrenes	102 \pm 22
C4-fluoranthenes/pyrenes	121 \pm 59
C1-benzanthracenes/chrysenes/triphenylenes	208 \pm 43
C2-benzanthracenes/chrysenes/triphenylenes	120 \pm 24
C3-benzanthracenes/chrysenes/triphenylenes	73 \pm 31
C4-benzanthracenes/chrysenes/triphenylenes	41 \pm 11

^(a) The reference mass fraction value reported on a dry-mass basis is the median of results using one analytical technique. The expanded uncertainty, U , is calculated as $U = ku_c$, where u_c is one standard deviation of the median, and the coverage factor, $k = 2$. The measurand is the total mass fraction of the alkylated PAH groups listed as determined by the interlaboratory study methods. The values listed are metrologically traceable to the SI unit of mass fraction, expressed as micrograms per kilogram on a dry-mass basis.

^(b) Data from the interlaboratory study [12].

Table 9. Reference Mass Fraction Values for Hopanes and Steranes in SRM 1941b

Hopane or Sterane	Mass Fraction ^(a,b) (µg/kg)
17α(H)-22,29,30-Trisnorhopane	54 ± 18
17α(H)-21β(H)-30-Norhopane	137 ± 21
17α(H)-21β(H)-30-Hopane	215 ± 44
17α(H)-21β(H)-22R-Homohopane	44 ± 10
17α(H)-21β(H)-22S-Homohopane	48 ± 13
5α(H)-14α(H),17α(H)-Cholestane 20R	41 ± 11
5α(H)-14β(H),17β(H)-Cholestane 20R	27 ± 6
5α(H)-14β(H),17β(H)-24-Methylcholestane 20R	21 ± 8
5α(H)-14α(H),17α(H)-24-Ethylcholestane 20R	19 ± 5
5α(H)-14β(H),17β(H)-24-Ethylcholestane 20R	41 ± 9

- ^(a) The reference mass fraction value reported on a dry-mass basis is the median of results using one analytical technique. The expanded uncertainty, U , is calculated as $U = k u_c$, where u_c is one standard deviation of the median, and the coverage factor, $k = 2$. The measurand is the total mass fraction of the constituent listed as determined by the methods used during the interlaboratory study. The values are metrologically traceable to the SI unit of mass, expressed as micrograms per kilogram on a dry-mass basis.
- ^(b) Data from the interlaboratory study [12].

Table 10. Reference Mass Fraction Value for Total Organic Carbon in SRM 1941b

Total Organic Carbon (TOC)	2.99 % ± 0.24 % ^(a,b)
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- ^(a) Mass fraction is reported on a dry-mass basis; material as received contains approximately 2.4 % moisture.
- ^(b) The reference value for total organic carbon is a weighted mean value from routine measurements made by two laboratories [21]. The uncertainty listed is an expanded uncertainty about the mean, with coverage factor 2 (approximately 95 % confidence), calculated by combining a between-method variance incorporating inter-method bias with a pooled within-method variance. The reporting follows the ISO/JCGM Guides [2]. The measurand is the total mass fraction of TOC listed as determined by the methods indicated. The values listed are metrologically traceable to the SI unit of mass, expressed as a percent on a dry-mass basis.

Table 11. Information Mass Fraction Values for Carbon, Hydrogen, and Nitrogen in SRM 1941b

Elements	Mass Fractions ^(a) (%)
Carbon	3.3
Hydrogen	1.2
Nitrogen	<0.5

- ^(a) Mass fraction is reported on a dry-mass basis; material as received contains approximately 2.4 % moisture.

REFERENCES

- [1] May, W.E.; Parris, R.M.; Beck II, C.M.; Fassett, J.D.; Greenberg, R.R.; Guenther, F.R.; Kramer, G.W.; Wise, S.A.; Gills, T.E.; Colbert, J.C.; Gettings, R.J.; MacDonald, B.R.; *Definition of Terms and Modes Used at NIST for Value-Assignment of Reference Materials for Chemical Measurements*; NIST Special Publication 260-136 (2000); available at <http://www.nist.gov/srm/publications.cfm> (accessed Jan 2015).
- [2] Wise, S.A.; Poster, D.L.; Schantz, M.M.; Kucklick, J.R.; Sander, L.C.; Lopez de Alda, M.; Schubert, P.; Parris, R.M.; Porter, B.J.; *Two New Marine Sediment Standard Reference Materials (SRMs) for the Determination of Organic Contaminants*; Anal. Bioanal. Chem., Vol. 378, pp. 1251–1264 (2004).
- [3] Wise, S.A.; Chesler, S.N.; Hertz, H.S.; Hilpert, L.R.; May, W.E.; *Chemically-Bonded Aminosilane Stationary Phase for the High Performance Liquid Chromatographic Separation of Polynuclear Aromatic Hydrocarbons*; Anal. Chem., Vol. 49, pp. 2306–2310 (1977).
- [4] May, W.E.; Wise, S.A.; *Liquid Chromatographic Determination of Polycyclic Aromatic Hydrocarbons in Air Particulate Extracts*; Anal. Chem., Vol. 56, pp. 225–232 (1984).
- [5] Wise, S.A.; Benner, B.A.; Byrd, G.D.; Chesler, S.N.; Rebbert, R.E.; Schantz, M.M.; *Determination of Polycyclic Aromatic Hydrocarbons in a Coal Tar Standard Reference Material*; Anal. Chem., Vol. 60, pp. 887–894 (1988).
- [6] Wise, S.A.; Deissler, A.; Sander, L.C.; *Liquid Chromatographic Determination of Polycyclic Aromatic Hydrocarbon Isomers of Molecular Weight 278 and 302 in Environmental Standard Reference Materials*; Polycyclic Aromat. Compd., Vol. 3, pp. 169–184 (1993).
- [7] Schantz, M.M.; Parris, R.M.; Wise, S.A.; *NIST/NOAA Intercomparison Exercise Program for Organic Contaminants in the Marine Environment: Description and Results of 1999 Organic Intercomparison Exercises*; NOAA Technical Memorandum NOS NCCOS CCMA 146, Silver Spring, MD (2000).
- [8] Schubert, P.; Schantz, M.M.; Sander, L.C.; Wise, S.A.; *Determination of Polycyclic Aromatic Hydrocarbons with Molecular Mass 300 and 302 in Environmental-Matrix Standard Reference Materials by Gas Chromatography-Mass Spectrometry*; Anal. Chem., Vol. 75, pp. 234–246 (2003).
- [9] Ballschmiter, K.; Zell, M.; *Analysis of Polychlorinated Biphenyls (PCB) by Glass Capillary Gas Chromatography - Composition of Technical Aroclor- and Clophen-PCB Mixtures*; Fresenius' Z. Anal. Chem., Vol. 302, pp. 20–31 (1980).
- [10] Schulte, E.; Malisch, R.; *Calculation of the Real PCB Content in Environmental Samples. I. Investigation of the Composition of Two Technical PCB Mixtures*; Fresenius' Z. Anal. Chem., Vol. 314, pp. 545–551 (1983).
- [11] Brubaker, W.W., Jr.; Schantz, M.M.; Wise, S.A.; *Determination of Non-ortho Polychlorinated Biphenyls in Environmental Standard Reference Materials*; Fresenius' J. Anal. Chem., Vol. 367, pp. 401–406 (2000).
- [12] Schantz, M.M.; Kucklick, J.R.; *NIST Interlaboratory Analytical Comparison Study to Support Deepwater Horizon Natural Resource Damage Assessment: Description and Results for Crude Oil QA10OIL01*; NISTIR 7793 (2011).
- [13] Ruhkin, A.L.; Vangel, M.G.; *Estimation of a Common Mean and Weighted Means Statistics*; J. Am. Statist. Assoc., Vol. 93, pp. 303–308 (1998).
- [14] JCGM 100:2008; *Evaluation of Measurement Data — Guide to the Expression of Uncertainty in Measurement (GUM 1995 with Minor Corrections)*; Joint Committee for Guides in Metrology (2008); available at http://www.bipm.org/utis/common/documents/jcgm/JCGM_100_2008_E.pdf (accessed Jan 2015); see also Taylor, B.N.; Kuyatt, C.E.; *Guidelines for Evaluating and Expressing the Uncertainty of NIST Measurement Results*; NIST Technical Note 1297; U.S. Government Printing Office: Washington, DC (1994); available at <http://www.nist.gov/pml/pubs/tn1297/index.cfm> (accessed Jan 2015).
- [15] JCGM 101:2008; *Evaluation of measurement data – Supplement 1 to the “Guide to the expression of uncertainty in measurement” - Propagation of distributions using a Monte Carlo method*; JCGM (2008); available at http://www.bipm.org/utis/common/documents/jcgm/JCGM_101_2008_E.pdf (accessed Jan 2015).
- [16] Levenson, M.S.; Banks, D.L.; Eberhardt, K.R.; Gill, L.M.; Guthrie, W.F.; Liu, H.-K.; Vangel, M.G.; Yen, J.H.; Zhang, N.F.; *An Approach to Combining Results from Multiple Methods Motivated by the ISO GUM*; J. Res. Natl. Inst. Stand. Technol., Vol. 105, pp. 571–579 (2000).

Certificate Revision History: 16 January 2015 (Corrected IUPAC name for PCB-56 and PCB-107; editorial changes); 10 June 2014 (Units corrected from mg/kg to µg/kg in Tables 8 and 9; editorial changes); 10 April 2012 (Reference value added for alkylated PAH groups, hopanes, and steranes; extension of certification period; editorial changes); 16 August 2004 (Reference values for the butyl tins removed; editorial changes); 15 July 2002 (Original certificate date).

Users of this SRM should ensure that the Certificate of Analysis in their possession is current. This can be accomplished by contacting the SRM Program: telephone (301) 975-2200; fax (301) 948-3730; e-mail srminfo@nist.gov; or via the Internet at <http://www.nist.gov/srm>.

APPENDIX A

The laboratories listed below performed measurements that contributed to the certification of PAHs, PCBs, and chlorinated pesticides in SRM 1941b Organics in Marine Sediment.

Arthur D. Little, Inc; Cambridge, MA
Axys Analytical Services; Sidney, BC, Canada
B & B Laboratories; College Station, TX
Battelle Ocean Sciences; Duxbury, MA
Bedford Institute of Oceanography; Dartmouth, NS, Canada
California Department of Fish and Game; Rancho Cordova, CA
Central Contra Costa Sanitary District; Martinez, CA
Chesapeake Biological Laboratory; Solomons, MD
Centro de Investigaciones Energeticas Medioambientales y Tecnologicas; Madrid, Spain
City of Los Angeles Environmental Monitoring Division; Playa del Rey, CA
City of San Jose Environmental Services Department; San Jose, CA
Columbia Analytical Services; Kelso, WA
East Bay Municipal Utility District; Oakland, CA
Florida Department of Environmental Protection; Tallahassee, FL
Manchester Environmental Laboratory; Port Orchard, WA
Murray State University; Murray, KY
Massachusetts Water Resources Authority Central Lab; Winthrop, MA
National Research Council of Canada; Ottawa, Ontario, Canada
National Oceanic and Atmospheric Association (NOAA), National Marine Fisheries Service (NMFS), Auke Bay Laboratory; Juneau, AK
NOAA, National Ocean Service/Center for Coastal Environmental Health and Biomolecular Research; Charleston, SC
NOAA, NMFS, Sandy Hook Marine Laboratory; Highlands, NJ
NOAA, NMFS, Northwest Fisheries Science Center; Seattle, WA
Orange County Sanitation District; Fountain Valley, CA
Philip Analytical Services; Burlington, Ontario, Canada
Serv de Hidrografia Naval; Buenos Aires, Argentina
Skidaway Institute of Technology; Savannah, GA
Southwest Laboratory of Oklahoma; Broken Arrow, OK
Severn Trent Knoxville Laboratory; Knoxville, TN
Texas A&M University, Geochemical and Environmental Research Group; College Station, TX
Texas Parks and Wildlife Department; San Marcos, TX
University of California at Los Angeles, Institute of Geophysics and Planetary Physics; Los Angeles, CA
University of Connecticut, Environmental Research Institute; Storrs, CT
University of Rhode Island, Graduate School of Oceanography; Narragansett, RI
US Department of Agriculture, Environmental Chemistry Laboratory; Beltsville, MD
US Environmental Protection Agency, Atlantic Ecology Division; Narragansett, RI
US Geological Survey, National Water Quality Laboratory; Denver, CO
Woods Hole Group Environmental Lab; Raynham, MA
Wright State University; Dayton, OH

APPENDIX B

The laboratories listed below performed measurements that contributed to the certification of alkylated PAH groups, hopanes, and steranes in SRM 1941b Organics in Marine Sediment.

Alpha Analytical, Inc.; Mansfield, MA
Analytical Resources, Inc.; Tukwila, WA
Axy's Analytical Services; Sydney, BC, Canada
Battelle Analytical & Environmental Chemistry Laboratory; Duxbury, MA
Center for Laboratory Sciences; Pasco, WA
Columbia Analytical Services; Jacksonville, FL
Columbia Analytical Services; Rochester, NY
Columbia Analytical Services, Kelso, WA
Florida Department of Environmental Protection; Tallahassee, FL
Florida International University; North Miami, FL
Michigan Department of Natural Resources and Environment; Lansing, MI
Mississippi State Chemical Laboratory; Mississippi State, MS
NIST; Charleston, SC
NIST; Gaithersburg, MD
NOAA/NCCOS/NOS; Charleston, SC
NOAA/NMFS/Alaska Fisheries Science Center; Juneau, AK
NY State Department of Health; Albany, NY
Pace Analytical Services, Inc. Minneapolis; Minneapolis, MN
RJ Lee Group, Inc; Monroeville, PA
TDI/B&B Laboratories, Inc.; College Station, TX
TestAmerica Laboratories; Mobile, AL
TestAmerica Laboratories; West Sacramento, CA
TestAmerica Laboratories; University Park, IL
TestAmerica Laboratories; Schriever, LA
TestAmerica Laboratories; Edison, NJ
TestAmerica Laboratories; Knoxville, TN
TestAmerica Laboratories; Pittsburgh, PA
TestAmerica Laboratories; South Burlington, VT
TestAmerica Laboratories; Tacoma, WA
US Army Engineer Research and Development Center; Vicksburg, MS
USGS Columbia Environmental Research Center; Columbia, MO
University of Iowa, State Hygienic Laboratory; Iowa City, IO
Washington State Public Health Laboratories; Shoreline, WA



Date of Issue:
31 March 2014

SAFETY DATA SHEET

1. SUBSTANCE AND SOURCE IDENTIFICATION

Product Identifier

SRM Number: 1941b
SRM Name: Organics in Marine Sediment
Other Means of Identification: Not applicable.

Recommended Use of This Material and Restrictions of Use

This Standard Reference Material (SRM) is marine sediment collected at the mouth of the Baltimore (MD) Harbor. SRM 1941b is intended for use in evaluating analytical methods for the determination of selected polycyclic aromatic hydrocarbons (PAHs), polychlorinated biphenyl (PCB) congeners, and chlorinated pesticides in marine sediment and similar matrices. All of the constituents for which certified, reference, and information values are provided in SRM 1941b were naturally present in the sediment before processing. A unit of SRM 1941b consists of a bottle containing 50 g of radiation-sterilized, freeze-dried sediment.

Company Information

National Institute of Standards and Technology
Standard Reference Materials Program
100 Bureau Drive, Stop 2300
Gaithersburg, Maryland 20899-2300

Telephone: 301-975-2200
FAX: 301-948-3730
E-mail: SRMMSDS@nist.gov
Website: <http://www.nist.gov/srm>

Emergency Telephone ChemTrec:
1-800-424-9300 (North America)
+1-703-527-3887 (International)

2. HAZARDS IDENTIFICATION

Classification

Physical Hazard: Not classified.
Health Hazard: Not classified.

Label Elements

Symbol
No Symbol/Pictogram

Signal Word
Not applicable.

Hazard Statement(s): Not applicable.

Precautionary Statement(s): Not applicable.

Hazards Not Otherwise Classified: Not applicable.

Ingredients(s) with Unknown Acute Toxicity: Not applicable.

3. COMPOSITION AND INFORMATION ON HAZARDOUS INGREDIENTS

Substance: Marine sediment

Other Designations: Sediment.

This material is naturally occurring marine sediment from an urban area. The material contains trace amounts of polycyclic aromatic hydrocarbons (PAHs), polychlorinated biphenyl (PCB) congeners, and should be handled with care. Components are listed in compliance with OSHA's 29 CFR 1910.1200; for the actual values see the Certificate of Analysis.

Hazardous Component(s)	CAS Number	EC Number (EINECS)	Nominal Mass Concentration (%)
Marine Sediment	Not available	Not available	23C0107 CLPLIKE (Rev0) - Page 515 of 540 100

4. FIRST AID MEASURES

Description of First Aid Measures:

Inhalation: If adverse effects occur, remove to uncontaminated area. If not breathing, give artificial respiration or oxygen by qualified personnel. Seek immediate medical attention.

Skin Contact: Wash skin with soap and water.

Eye Contact: Flush eyes with water for at least 15 minutes. If necessary, seek medical attention.

Ingestion: If adverse effects occur after ingestion, seek medical treatment.

Most Important Symptoms/Effects, Acute and Delayed: May cause irritation.

Indication of any immediate medical attention and special treatment needed, if necessary: If any of the above symptoms are present, seek medical attention if needed.

5. FIRE FIGHTING MEASURES

Fire and Explosion Hazards: Negligible fire hazard. Avoid generating dust. See Section 9, "Physical and Chemical Properties" for flammability properties.

Extinguishing Media:

Suitable: Use extinguishing media appropriate for surrounding fire.

Unsuitable: None listed.

Specific Hazards Arising from the Chemical: None listed.

Special Protective Equipment and Precautions for Fire-Fighters: Avoid inhalation of material or combustion byproducts. Wear full protective clothing and NIOSH approved self-contained breathing apparatus (SCBA).

NFPA Ratings (0 = Minimal; 1 = Slight; 2 = Moderate; 3 = Serious; 4 = Severe)

Health = 1

Fire = 0

Reactivity = 0

6. ACCIDENTAL RELEASE MEASURES

Personal Precautions, Protective Equipment and Emergency Procedures: Any accumulated material on surfaces should be removed and properly disposed of. Use suitable protective equipment; see Section 8, "Exposure Controls and Personal Protection".

Methods and Materials for Containment and Clean up: Collect spilled material in appropriate container for disposal. Keep out of water supplies and sewers. Keep unnecessary people away, isolate hazard area and deny entry.

7. HANDLING AND STORAGE

Safe Handling Precautions: Minimize dust generation and accumulation on surfaces. Routine housekeeping should be instituted to ensure that dusts do not accumulate on surfaces. See Section 8, "Exposure Controls and Personal Protection".

Storage: Store and handling in accordance with all current regulations and standards.

8. EXPOSURE CONTROLS AND PERSONAL PROTECTION

Exposure Limits: No occupational exposure limits have been established for marine sediment. This material is a particulate matter and adequate inhalation/respiratory protection should be used to minimize exposure. The exposure limits for Particulates Not Otherwise Regulated (PNOR) are applicable.

OSHA (PEL): 15 mg/m³ (TWA, total particulates not otherwise regulated)

OSHA (PEL) 5 mg/m³ (TWA, respirable particulates not otherwise regulated)

NIOSH (REL): 10 mg/m³ (TWA, total particulates not otherwise regulated, 8 h)

NIOSH (REL): 5 mg/m³ (TWA, respirable particulates not otherwise regulated)

Engineering Controls: Provide local exhaust or process enclosure ventilation system. Ensure compliance with applicable exposure limits.

Personal Protection: In accordance with OSHA 29 CFR 1910.132, subpart I, wear appropriate Personal Protective Equipment (PPE) to minimize exposure to this material.

Respiratory Protection: If workplace conditions warrant a respirator, a respiratory protection program that meets OSHA 29CFR 1910.134 must be followed. Refer to NIOSH 42 CFR 84 for applicable certified respirators.

Eye/Face Protection: Wear splash resistant safety goggles with a face shield. An eye wash station should be readily available near areas of use.

Skin and Body Protection: Personal protective equipment for the body should be selected based on the task being performed and the risks involved and should be approved by a specialist before handling this product. Chemical-resistant gloves should be worn at all times when handling chemicals.

9. PHYSICAL AND CHEMICAL PROPERTIES

Descriptive Properties:

Appearance (physical state, color, etc.):	amorphous powder
Molecular Formula:	not applicable
Molar Mass (g/mol):	not applicable
Odor:	not available
Odor threshold:	not available
pH:	not available
Evaporation rate:	not applicable
Melting point/freezing point (°C):	not available
Specific Gravity (water=1)	not available
Vapor Pressure (mmHg):	not applicable
Vapor Density (air = 1):	not applicable
Viscosity (cP):	not applicable
Solubility(ies):	not available
Partition coefficient (n-octanol/water):	not available
Particle Size:	<150 µm

Thermal Stability Properties:

Autoignition Temperature (°C):	not available
Thermal Decomposition (°C):	not available
Initial boiling point and boiling range (°C):	not available
Explosive Limits, LEL (Volume %):	not available
Explosive Limits, UEL (Volume %):	not available
Flash Point (°C):	not available
Flammability (solid, gas):	not available

10. STABILITY AND REACTIVITY

Reactivity: Stable at normal temperatures and pressure.

Stability: X Stable Unstable

Possible Hazardous Reactions: None listed.

Conditions to Avoid: Avoid generating dust.

Incompatible Materials: None listed.

Fire/Explosion Information: See Section 5, "Fire Fighting Measures".

Hazardous Decomposition: Thermal decomposition will produce oxides of carbon.

Hazardous Polymerization: Will Occur X Will Not Occur

11. TOXICOLOGICAL INFORMATION

Route of Exposure: Inhalation Skin Ingestion

Symptoms Related to the Physical, Chemical and Toxicological Characteristics: Generated dust may cause irritation if inhaled.

Potential Health Effects (Acute, Chronic and Delayed):

Inhalation: Generated dust may cause irritation.

Skin Contact: May cause mechanical irritation.

Eye Contact: May cause mechanical irritation.

Ingestion: No data available.

Numerical Measures of Toxicity:

Acute Toxicity: Not classified; no data available.

Skin Corrosion/Irritation: Not classified; no data available.

Serious Eye damage/ Eye irritation: Not classified; no data available.

Respiratory Sensitization: Not classified; no data available.

Skin Sensitization: Not classified; no data available.

Germ Cell Mutagenicity: Not classified; no data available.

Carcinogenicity: Not classified.

Listed as a Carcinogen/Potential Carcinogen Yes No
Marine sediment is not listed by NTP, IARC or OSHA as a carcinogen.

Reproductive Toxicity: Not classified; no data available.

Specific Target Organ Toxicity, Single Exposure: Not classified; no data available.

Specific Target Organ Toxicity, Repeated Exposure: Not classified; no data available.

Aspiration Hazard: Not classified; no data available.

12. ECOLOGICAL INFORMATION

Ecotoxicity Data: No data available.

Persistence and Degradability: No data available.

Bioaccumulative Potential: No data available.

Mobility in Soil: No data available.

Other Adverse effects: No data available.

13. DISPOSAL CONSIDERATIONS

Waste Disposal: Dispose of waste in accordance with all applicable federal, state, and local regulations.

14. TRANSPORTATION INFORMATION

U.S. DOT and IATA: Not regulated by DOT or IATA.

15. REGULATORY INFORMATION

U.S. Regulations:

CERCLA Sections 102a/103 (40 CFR 302.4): Not regulated.

SARA Title III Section 302 (40 CFR 355.30): Not regulated.

SARA Title III Section 304 (40 CFR 355.40): Not regulated.

SARA Title III Sections 311/312 Hazardous Categories (40 CFR 370.21):

ACUTE HEALTH: No.
CHRONIC HEALTH: No.
FIRE: No.
REACTIVE: No.
PRESSURE: No.

State Regulations:

California Proposition 65: Not listed.

U.S. TSCA Inventory: Not listed.

TSCA 12(b), Export Notification: Not listed.

Canadian Regulations:

WHMIS Information: Not provided for this material.

16. OTHER INFORMATION

Issue Date: 31 March 2014

Sources: 29 CFR Occupational Health and Safety Office (OSHA) 1910.1000, *Limits for Air Contaminants*, Table Z-1; available at http://www.osha.gov/pls/oshaweb/owadisp.show_document?p_table=STANDARDS&p_id=9992 (accessed Mar 2014).

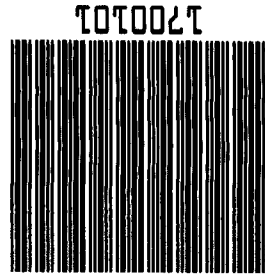
Center for Disease Control (CDC) NIOSH Pocket Guide to Chemical Hazards, *Particulates not otherwise regulated*; available at <http://www.cdc.gov/niosh/npg/npgd0480.html> (accessed Mar 2014).

Key of Acronyms:

ACGIH	American Conference of Governmental Industrial Hygienists	NRC	Nuclear Regulatory Commission
ALI	Annual Limit on Intake	NTP	National Toxicology Program
CAS	Chemical Abstracts Service	OSHA	Occupational Safety and Health Administration
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act	PEL	Permissible Exposure Limit
CFR	Code of Federal Regulations	RCRA	Resource Conservation and Recovery Act
DOT	Department of Transportation	REL	Recommended Exposure Limit
EC50	Effective Concentration, 50 %	RM	Reference Material
EINECS	European Inventory of Existing Commercial Chemical Substances	RQ	Reportable Quantity
EPCRA	Emergency Planning and Community Right-to-Know Act	RTECS	Registry of Toxic Effects of Chemical Substances
IARC	International Agency for Research on Cancer	SARA	Superfund Amendments and Reauthorization Act
IATA	International Air Transportation Agency	SCBA	Self-Contained Breathing Apparatus
IDLH	Immediately Dangerous to Life and Health	SRM	Standard Reference Material
LC50	Lethal Concentration, 50 %	STEL	Short Term Exposure Limit
LD50	Lethal Dose, 50 %	TLV	Threshold Limit Value
LEL	Lower Explosive Limit	TPQ	Threshold Planning Quantity
MSDS	Material Safety Data Sheet	TSCA	Toxic Substances Control Act
NFPA	National Fire Protection Association	TWA	Time Weighted Average
NIOSH	National Institute for Occupational Safety and Health	UEL	Upper Explosive Limit
NIST	National Institute of Standards and Technology	WHMIS	Workplace Hazardous Materials Information System

Disclaimer: Physical and chemical data contained in this SDS are provided only for use in assessing the hazardous nature of the material. The SDS was prepared carefully, using current references; however, NIST does not certify the data in the SDS. The certified values for this material are given in the NIST Certificate of Analysis.

Users of this SRM should ensure that the SDS in their possession is current. This can be accomplished by contacting the SRM Program: telephone (301) 975-2200; fax (301) 948-3730; 2300a1b7r0undsk@nist.gov; Page 518 of 540
Internet at <http://www.nist.gov/srm/>



Picked by
9/21/16 04:04 PM

Weight	
# of pieces	
Packed by	
Picked by	

Order	UOM	Ship	UOM	B/O	UOM	Item	Description
1	EACH	1	EACH	0	EACH	1941B	Organics in Marine Sediment
						Total qty:	1 / EACH
NOT FOR HUMAN CONSUMPTION, LABORATORY USE ONLY.							

Order discrepancies (other than back ordered items) must be reported to our Customer Relations Department at 301-975-6776 within 5 days of receipt of shipment or this order will be considered complete. NIST SRMs/RMs are generally not returnable - with the exception of defective goods or shipments made in error by NIST. To return a SRM/RM, please call for instructions and a Return SRM/RM Authorization Number before shipment. Returns WILL NOT BE ACCEPTED without prior authorization.

Ship via	UFS Ground	Description	
Salesmen	MCMIDM2	Instructions	
Contact	DAVE MITCHELL	Prof	
		Truck#	
		Blanket	
		Ship from	

Ship to:
68456
DAVE MITCHELL
ANALYTICAL RESOURCES INC
4611 S 134TH PLACE
SUITE 100
TUKWILA, WA 98168-3240
1 (206) 695-6205

Bill to:
68456
DAVE MITCHELL
ANALYTICAL RESOURCES INC
4611 S 134TH PLACE
SUITE 100
TUKWILA, WA 98168-3240
1 (206) 695-6205



MP Biomedicals, LLC

29525 Fountain Parkway
Solon, Ohio 44139

Telephone: 440/337-1200
Toll Free: 800/854-0530

Fax: 440/337-1180
web: www.mpbio.com

Certificate of Analysis

Product Description: Microcrystalline Cellulose Powder_
Catalog Number: 191499_
Lot: Q9483_


Formula: (C ₆ H ₁₀ O ₅) _n CAS #: 9004-34-6 Physical Description: White Powder	Formula Weight: N/A Storage: 15 - 30°C
---	---

Test	Specification	Result
Identity Test	Passes	Passes
Purity	97.0 - 102.0%	97.0 - 102.0%
Moisture	<5.0%	3.4%
Particle Size/Mesh	Wt %	
+60 mesh	<8%	<1%
+200 mesh	>45%	55%
pH	5 - 7	6.73
Residue on Ignition	<0.05%	<0.05%
Water Soluble Substances	<12.0 mg/5 g	4.5 mg/5 g
Heavy Metals	<10 ppm	<10 ppm

H001822
Microcrystalline Cellulose Powder (TOC)
Expires 11/30/2022
Prepared By Casey English 2/22/2019

Identification A & B: Passes
Bulk Density: 0.29 g/ml
Bulk Density (graduated cylinder): 0.31 g/ml
Conductivity: 18 µS/cm
Starch: Negative
Ether Soluble Substances: 0.01%
Total Aerobic microbial Count: 100 cfu/g
Total Mold and Yeast Count: 20 cfu/g
Staphylococcus aureus: Absent/1 g
Pseudomonas aeruginosa: Absent/1 g
E. coli: Absent/1 g
Salmonella: Absent/10 g
Particle size:

- 450 mesh: 77%
- d10: 37 um
- d50: 139 um
- d90: 271 um
TUP: <9/600 cm²
Degree of brightness: >88%
Powder flow-angle of repose: <42°
Recommended Retest Date: 11/30/2022



07/26/2018 - John Huang, PhD
MP Biomedicals, LLC.
Quality Control Manager

This is an electronically generated document
<mailto:biotech@mpbio.com>
<http://www.mpbio.com>

Online Ordering, MSDSs, certificates of analysis and data sheets now available on our web site
Technical Service: 1-800-279-5490 (440-337-1200) Customer Service: 1-800-854-0530 (440-337-1200)



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Certificate of Analysis

Product Description: Microcrystalline Cellulose Powder_
Catalog Number: 191499_
Lot: Q9483_

Formula: (C₆H₁₀O₅)_n
CAS #: 9004-34-6
Physical Description: White Powder

Formula Weight: N/A
Storage: 15 - 30°C


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Identity Test	Passes	Passes
Purity	97.0 - 102.0%	97.0 - 102.0%
Moisture	<5.0%	3.4%
Particle Size/Mesh	Wt %	
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+200 mesh	>45%	55%
pH	5 - 7	6.73
Residue on Ignition	<0.05%	<0.05%
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H001822

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Expires 11/30/2022
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- d50: 139 um
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Powder flow-angle of repose: <42°
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Lot: Q9483_

Formula: (C₆H₁₀O₅)_n
CAS #: 9004-34-6
Physical Description: White Powder

Formula Weight: N/A
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
Test	Specification	Result
Identity Test	Passes	Passes
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Moisture	<5.0%	3.4%
Particle Size/Mesh	Wt %	
+60 mesh	<8%	<1%
+200 mesh	>45%	55%
pH	5 - 7	6.73
Residue on Ignition	<0.05%	<0.05%
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07/26/2018 - John Huang, PhD
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Form I
INORGANIC ANALYSIS DATA SHEET
SM 2540 G-97

LDW22-SS826

Laboratory: Analytical Resources, LLC

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Sediment Laboratory ID: 23C0107-01 A SDG: 23C0107

Sampled: 03/02/23 16:25 Prepared: 03/09/23 09:37 File ID:

% Solids: 43.35 Preparation: No Prep Wet Chem Analyzed: 03/09/23 09:41

Batch: BLC0220 Sequence:

Instrument: BAL2 Calibration: 5 g Wet / 5 g

CAS NO.	Analyte	Concentration (%)	Dilution Factor	MDL	MRL	Q
	Total Solids	43.35	1	0.04	0.04	



Form I
INORGANIC ANALYSIS DATA SHEET
SM 2540 G-97

LDW21-IT608C

Laboratory: Analytical Resources, LLC

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Sediment Laboratory ID: 23C0107-02 A SDG: 23C0107

Sampled: 07/13/21 06:54 Prepared: 03/09/23 09:37 File ID:

% Solids: 67.49 Preparation: No Prep Wet Chem Analyzed: 03/09/23 09:41

Batch: BLC0220 Sequence:

Instrument: BAL2 Calibration: 5 g Wet / 5 g

CAS NO.	Analyte	Concentration (%)	Dilution Factor	MDL	MRL	Q
	Total Solids	67.49	1	0.04	0.04	*



Form I
INORGANIC ANALYSIS DATA SHEET
SM 2540 G-97

LDW21-IT608D

Laboratory: Analytical Resources, LLC

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Sediment Laboratory ID: 23C0107-03 A SDG: 23C0107

Sampled: 07/13/21 06:54 Prepared: 03/09/23 09:37 File ID:

% Solids: 73.57 Preparation: No Prep Wet Chem Analyzed: 03/09/23 09:41

Batch: BLC0220 Sequence:

Instrument: BAL2 Calibration: 5 g Wet / 5 g

CAS NO.	Analyte	Concentration (%)	Dilution Factor	MDL	MRL	Q
	Total Solids	73.57	1	0.04	0.04	*



PREPARATION BATCH SUMMARY
SM 2540 G-97

Laboratory: Analytical Resources, LLC SDG: 23C0107
Client: Anchor QEA, LLC Project: AOC4 UR Phase 3
Batch: BLC0220 Batch Matrix: Solid Preparation: No Prep Wet Chem

SAMPLE NAME	LAB SAMPLE ID	LAB FILE ID	DATE PREPARED	OBSERVATIONS
LDW22-SS826	23C0107-01		03/09/23 09:37	
LDW21-IT608C	23C0107-02		03/09/23 09:37	
LDW21-IT608D	23C0107-03		03/09/23 09:37	
Blank	BLC0220-BLK1		03/09/23 09:37	
LDW22-SS826	BLC0220-DUP1		03/09/23 09:37	
LDW22-SS826	BLC0220-DUP2		03/09/23 09:37	

TOTAL SOLIDS/VOLATILE SOLIDS (TS / TVS) BENCHSHEET for Solid samples													Batch:		BLC0220		
Method: PSEP 1986, SM2540, EPA 160.1													Date:		3/9/2023 9:41		
(dry at 104 (12-24 hr) then combust at 550 (30 min))													Analyst:		UW		
Instrumentation			Drying Ovens:			12			Analytical Balance:			BAL2					
			Muffle Furnace:			2											
Batch drying time						Oven Temps, °C						TVS (mg/kg dry wt) calculated as: Final ash wt (g) = (min ash wt - tare wt) TVS (mg/kg) = [(Dry wt-Ash wt)/ (dry weight)] *1,000,000 if ash wt > dry wt, "Chk for Err" if dry wt-ash wt < 0.001 g, "< (1/dry wt)*1,000,000					
record times as mm/dd/yy hh:mm			TS (%) calculated as:			Start Temp			106								
date/time in oven:			3/9/2023 10:15			Final dry wt (g) = (Dry Wt - Tare Wt)			Dry Cycle 1						105		
date/time out:			3/10/2023 13:35			TS = (Final Dry Wt)/(grams Sample-Tare)			Dry Cycle 2								
elapsed hrs =			27.3			> 24 hr			Dry Cycle 3								
Balance Calibration Check																	
Record weights to 4 places																	
Cal Weight ID:		CV-02		CV-02		CV-02		CV-02		CV-02		CV-02		CV-02			
Date & Time:		3/9/23 9:45		3/9/23 9:50		3/10/23 14:00											
Cal Wt (g):		10.0000		10.0000		9.9999											
		Cal OK!		Cal OK!		Cal OK!											
Sample ID	Dish #	Tare Wt. (g)	Dish & Sample (g)	Dry Wt 104C (grams)			dry Wt (g)	TS (%)	Notes	ASH WT 550C (grams)			Ash Wt (g)	TVS		Notes	
				1	2	3				1	2	3		(mg/kg)	(%)		
BLC0220-BLK1	1	0.8137	0.0000	0.8134			-0.0003	0.04%									
23C0107-01	2	0.8011	7.2723	3.6063			2.8052	43.35%									
BLC0220-DUP1	3	0.8104	7.5570	3.7489			2.9385	43.56%	RPD=0.5								
BLC0220-DUP2	4	0.8088	7.3915	3.6702			2.8614	43.47%	RSD=0.2								
23C0107-02	5	0.7989	6.9080	4.9222			4.1233	67.49%									
23C0107-03	6	0.8116	7.5421	5.7632			4.9516	73.57%									
23C0108-01	7	0.7831	8.2550	4.9336			4.1505	55.55%									
23C0108-02	8	0.8087	9.4611	5.3583			4.5496	52.58%									
23C0108-03	9	0.8120	8.2007	4.4442			3.6322	49.16%									
23C0108-04	10	0.7802	9.1662	4.9118			4.1316	49.27%									
23C0108-05	11	0.7787	9.3389	5.3973			4.6186	53.95%									
23C0108-06	12	0.7903	7.4789	3.6011			2.8108	42.02%									
23C0108-07	13	0.7757	8.5513	3.9932			3.2175	41.38%									
23C0108-08	14	0.8050	7.1523	3.8375			3.0325	47.78%									
23C0108-09	15	0.8102	7.0271	3.5741			2.7639	44.46%									
23C0108-10	16	0.8081	7.7303	4.0760			3.2679	47.21%									
23C0109-01	17	0.8407	9.9869	4.8972			4.0565	44.35%									
23C0109-02	18	0.8211	5.6131	2.6322			1.8111	37.79%									
23C0109-03	19	0.8338	7.0443	3.0898			2.2560	36.33%									



Form I
METHOD BLANK DATA SHEET
SM 2540 G-97
TotalAnalytes

Blank

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Batch: BLC0220

Laboratory ID: BLC0220-BLK1

Prepared: 03/09/23 09:37

Matrix: Solid

Preparation: No Prep Wet Chem

Analyzed: 03/09/23 09:41

Sequence:

Calibration:

Instrument: BAL2

CAS NO.	Analyte	Concentration (%)	Dilution Factor	MDL	MRL	Q
	Total Solids	ND	1	0.04	0.04	U



DUPLICATES
SM 2540 G-97

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Laboratory ID: BLC0220-DUP1

Batch: BLC0220

Lab Source ID: 23C0107-01

Preparation: No Prep Wet Chem

Initial/Final: 5 g / 5 g

Source Sample Name: LDW22-SS826

% Solids: 43.35

ANALYTE	CONTROL LIMIT	SAMPLE CONCENTRATION	DUPLICATE CONCENTRATION	RPD %	Q
Total Solids	20	43.35	43.56	0.475	

*: Values outside of QC limits

L: Analyte concentration is <=5 times the reporting limit and the replicate control limit defaults to Dup = +/-RL instead of 20% RPD



DUPLICATES
SM 2540 G-97

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Laboratory ID: BLC0220-DUP2

Batch: BLC0220

Lab Source ID: 23C0107-01

Preparation: No Prep Wet Chem

Initial/Final: 5 g / 5 g

Source Sample Name: LDW22-SS826

% Solids: 43.35

ANALYTE	CONTROL LIMIT	SAMPLE CONCENTRATION	DUPLICATE CONCENTRATION	RPD %	Q
Total Solids	20	43.35	43.47	0.275	

*: Values outside of QC limits

L: Analyte concentration is <=5 times the reporting limit and the replicate control limit defaults to Dup = +/-RL instead of 20% RPD



HOLDING TIME SUMMARY

Analysis: SM 2540 G-97

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Sample Name	Date Collected	Date Received	Date Prepared	Days to Prep	Max Days to Prep	Date Analyzed	Days to Analysis	Max Days to Analysis	Q
LDW22-SS826 23C0107-01	03/02/23 16:25	03/03/23 16:35	03/09/23 09:37	6	180	03/09/23 09:41	7	180	
LDW21-IT608C 23C0107-02	07/13/21 06:54	03/03/23 16:35	03/09/23 09:37	604	180	03/09/23 09:41	604	180	*
LDW21-IT608D 23C0107-03	07/13/21 06:54	03/03/23 16:35	03/09/23 09:37	604	180	03/09/23 09:41	604	180	*
Duplicate BLC0220-DUP1	03/02/23 16:25	03/03/23 16:35	03/09/23 09:37	6	180	03/09/23 09:41	7	180	
Duplicate BLC0220-DUP2	03/02/23 16:25	03/03/23 16:35	03/09/23 09:37	6	180	03/09/23 09:41	7	180	

* Indicates hold time exceedance.



Analytical Resources, LLC
Analytical Chemists and Consultants

METHOD DETECTION AND REPORTING LIMITS

SM 2540 G-97

Laboratory: Analytical Resources, LLC

SDG: 23C0107

Client: Anchor QEA, LLC

Project: AOC4 UR Phase 3

Matrix: Solid

Instrument:

Analyte	MDL	RL	Units
Total Solids	0.04	0.04	%

TOTAL SOLIDS BENCHSHEET						Batch:	BLC0101	
Method: PSEP 1986 (dry at 103-105 C)						Date:	3/6/2023 8:53	
						Analyst:	cr	
Instrumentation:						Drying Oven:	15	
						Analytical Balance:	B139298002	
Batch drying time			Oven Temp, C				TS (%) calculated as:	
Record times as mm/dd/yy hh:mm			Final dry wt (g) = (Dry Wt - Tare Wt)				Oven Temps, °C	
Date/time in oven:	3/7/2023 13:15		TS = (Final Dry Wt X 100)/(sample & dish -dish tare)				Start Temp:	108
Date/time out:	3/8/2023 9:10						End Temp:	104
Elapsed hrs:	19.9							
SAMPLE ID	Dish Tare Wt (g)	Dish with Sample (g)	Dry Wt (g)	Solids Wt (g)	TS (%)	Sample Decanted		
23C0107-01	0.8000	12.4700	5.6900	4.89	41.90%	Yes		
23C0108-01	0.7800	12.4300	7.0200	6.24	53.56%	No		
23C0108-02	0.8000	11.0200	6.1900	5.39	52.74%	No		
23C0108-03	0.8000	12.2600	6.3900	5.59	48.78%	No		
23C0108-04	0.8000	11.4100	5.8300	5.03	47.41%	No		
23C0108-05	0.8000	12.7000	6.5800	5.78	48.57%	No		
23C0108-06	0.8000	12.2000	5.7400	4.94	43.33%	No		
23C0108-07	0.8100	11.8300	5.2400	4.43	40.20%	No		
23C0108-08	0.8300	12.5600	6.2700	5.44	46.38%	No		
23C0108-09	0.8000	11.4300	5.3300	4.53	42.62%	No		
23C0108-10	0.7700	12.3800	6.2600	5.49	47.29%	No		
23C0109-01	0.8400	12.7700	6.0100	5.17	43.34%	No		
23C0109-02	0.8000	11.4400	4.6200	3.82	35.90%	No		
23C0109-03	0.8000	12.2600	4.9400	4.14	36.13%	No		

TOTAL SOLIDS BENCHSHEET			Batch:	BLC0101
Method: PSEP 1986 (dry at 103-105 C)			Date:	3/6/2023 8:53
Instrumentation			Analyst:	CR
			Drying Oven:	015
			Analytical Balance:	B139298002
Batch drying time				
Record times as mm/dd/yy hh:mm		Oven Temp, C	TS (%) calculated as:	
Date/time in oven:	3/7/23 13:15	108	Final dry wt (g) = (Dry Wt - Tare Wt)	
Date/time out:	3/7/23 09:14	104	TS = (Final Dry Wt X 100)/(sample & dish -dish tare)	
Elapsed hrs:	0.0			
		Oven Temps, °C		
		Start Temp:		108
		End Temp:		104

SAMPLE ID	Dish Tare Wt (g)	Dish with Sample (g)	Dry Wt (g)	Solids Wt (g)	TS (%)	Sample Decanted
23C0107-01 A	0.80	12.47	5.69			No Yes
23C0108-01	0.78	12.43	7.02			No
23C0108-02	0.80	11.02	6.19			No
23C0108-03	0.80	12.26	6.39			No
23C0108-04	0.80	11.41	5.83			No
23C0108-05	0.80	12.70	6.58			No
23C0108-06	0.80	12.20	5.74			No
23C0108-07	0.81	11.83	5.24			No
23C0108-08	0.83	12.56	6.27			No
23C0108-09	0.80	11.43	5.33			No
23C0108-10	0.77	12.38	6.26			No
23C0109-01	0.84	12.77	6.01			No
23C0109-02	0.80	11.44	4.62			No
23C0109-03 A	0.80	12.26	4.94			No

T/S + Screens

2 copies

TOTAL SOLIDS BENCHSHEET						Batch:	BLC0150
Method: PSEP 1986						Date:	3/7/2023 12:33
(dry at 103-105 C)						Analyst:	YL
Instrumentation						Drying Oven:	15
						Analytical Balance:	B146462614
Batch drying time			Oven Temp, C		TS (%) calculated as:		Oven Temps, °C
Record times as mm/dd/yy hh:mm					Final dry wt (g) = (Dry Wt - Tare Wt)		Start Temp: 105
Date/time in oven:	3/8/2023 10:30				TS = (Final Dry Wt X 100)/ (sample & dish -dish tare)		End Temp: 104
Date/time out:	3/9/2023 8:42						
Elapsed hrs:	22.2						
SAMPLE ID	Dish Tare Wt (g)	Dish with Sample (g)	Dry Wt (g)	Solids Wt (g)	TS (%)	Sample Decanted	
23C0107-02	0.8000	12.1700	8.4400	7.64	67.19%	No	
23C0107-03	0.8000	12.1100	9.0300	8.23	72.77%	No	

TOTAL SOLIDS BENCHSHEET				Batch:	BLC0150	
Method: PSEP 1986				Date:	3/7/2023 12:33	
(dry at 103-105 C)				Analyst:	JL	
Instrumentation				Drying Oven:	Φ15	
				Analytical Balance:	3146462614	
Batch drying time			Oven Temp, C	TS (%) calculated as:		
Record times as mm/dd/yy hh:mm				Final dry wt (g) = (Dry Wt - Tare Wt)		
Date/time in oven:	03/08/23	10:32	105	TS = (Final Dry Wt X 100) / (sample & dish - dish tare)		
Date/time out:	3/9/23	08:42	104			
Elapsed hrs:	0.0					
				Oven Temps, °C		
				Start Temp:	105	
				End Temp:	104	
SAMPLE ID	Dish Tare Wt (g)	Dish with Sample (g)	Dry Wt (g)	Solids Wt (g)	TS (%)	Sample Decanted
23C0107-02 A	0.80	12.17	8.44			No
23C0107-03 A	0.80	12.11	9.03			No

3/9/23

T/S + Surung
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