

## Atmospheric Analysis & Consulting, Inc.

Client : SWAPE  
Client Project Name : Bridgeton Sanitary Landfill Air Quality Assessment  
Client Project No. : NA  
AAC Project No. : 131099  
Reporting Date : 08/22/2013

On August 15, 2013, Atmospheric Analysis & Consulting, Inc. received four (4) DNPH impregnated silica gel cartridges for Carbonyls analysis by EPA Method TO-11A. Upon receipt the samples were assigned unique Laboratory ID numbers as follows:

Client Sample ID	AAC Sample ID
U-1 K DNPH	131099-65668
U-2 W2 DNPH	131099-65669
D-1 W6E DNPH	131099-65670
D-2 W6W DNPH	131099-65671

TO-11A - HPLC/UV analysis - A 10 $\mu$ L aliquot of the extract was analyzed by HPLC/UV following the analytical protocols of EPA Method TO-11A as specified in the SOW. Holding times for preparation and analysis were complied with.

No problems were encountered during the receipt, preparation, and/or analysis of these samples. The test results included in this report meet all requirements of the NELAC Standards and/or AAC SOP# TO.11.09.

I certify that this data is technically accurate, complete and in compliance with the terms and conditions of the contract. The Laboratory Director or his designee, as verified by the following signature, has authorized release of the data contained in this hardcopy data package.

If you have any questions or require further explanation of data results, please contact the undersigned.

  
Marcus Hueppe  
Laboratory Director

This report consists of 55 pages.



4424 131099

CHAIN OF CUSTODY RECORD / ANALYTICAL REQUEST FORM

Bridgeton Sanitary Landfill Air Quality Assessment

Client Name: SOIL / WATER AIR PROTECTION ENTERPRISE Telephone No. / Fax No.: (310) 434-0110 / (310) 434-0011 Date: August 14th Page 1 of 1

Project Manager: PAUL ROSENFELD, PH.D. Address: 1640 FIFTH STREET, SUITE 204, SANTA MONICA, CA 90401

Project Name and Location: BRIDGETON SANITARY LANDFILL AIR QUALITY ASSESSMENT

Sampled By: John Blank Sampler Signature: *John Blank*

LAB ID	SAMPLE ID NUMBER	Type	Date	Time	VOCS - EPA TO-15	Reduced Sulfur Compounds - ASTM D5504	Carbonyls - EPA TO-11A	Carboxylic Acids - Tube GC-MS	HCL - NIOSH 7903	Ammonia - OSHA ID-188	SO2 - OSHA ID-200	HCN - NIOSH 6010	Amines - NIOSH 2010M	Fixed Gases - EPA 3C	PAHs / Dioxins EPA TO-13A / 9A	Mercury - NIOSH 6009	Odor Evaluation	Special Instructions / Conditions of Receipt
<del>65586</del>	<del>U-1 K</del>	Tube	August 14th	240 min			X											Tube # 4440600652 240.7 L
65569	U-2 W2	Tube	August 14th	245 min			X											Tube # 4440600649 238.8 L
65570	D-1 W6E	Tube	August 14th	240 min			X											Tube # 4440600651 269.6 L
65571	D-1 W6W	Tube	August 14th	240 min			X											Tube # 4440600648 245.3 L

Requested Turnaround Time: Standard turn-around for all analyses. If possible deliver report within 2 weeks. OC Requirements: Provide Level IV OC Package for all Analyses.

Relinquished By: John Blank Date: August 14th Time: 12 Noon Received By: Date: Time:

Relinquished By: *John Blank* Date: August 14th Time: Received By: *WJF* Date: 8/15/13 Time: 1200

Samples received @ 5:00 PM

# AIR SAMPLING PUMP CALIBRATION LOG

Bridgeton Sanitary Landfill Air Quality Assessment

COMPLETED BY: John Blank



DATE: August 14th, 2013

PAGE: 1 OF 1

CALIBRATION INSTRUMENT : Biocore Defender510

INITIAL PUMP SETUP (PRE-SAMPLING FLOW CHECK)

Sample ID e.g. acetaldehyde	Analyte	SKC Tube ID	Air Pump Serial No.	START		END		Average Flow Rate L/Min	Total Sampled Minutes	Total Volume Liters
				Flow Rate (L/min)	Start Time (24 Hour)	Flow Rate (L/min)	Stop Time (24 Hour)			
U-1 K	Aldehydes	4440600652	67992	1.010	16:54:00	0.9958	20:54:00	1.003	240	240.696
U-2 W2	Aldehydes	4440600649	67385	1.028	17:55:00	0.962	21:55:00	0.995	240	238.800
D-1 W6E	Aldehydes	4440600651	71526	1.129	17:20:00	1.072	21:25:00	1.101	245	269.623
D-1 W6W	Aldehydes	4440600648	59912	1.082	17:30:00	0.962	21:30:00	1.022	240	245.280

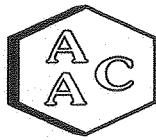
NOTES / LOCATION REFERENCES

TUBES:

ANALYTE	SKC TUBE ID
Aldehydes	226-120
Amines	226-10
Ammonia	226-29
Carboxylic Acids	226-55

SKC TUBE ID
226-10-03
226-28
226-17-1A
226-80

SOIL / WATER / AIR PROTECTION ENTERPRISE



### SAMPLE RECEIPT / LOG-IN REPORT

AAC Project 131099

Received By: W. Horn

<u>Sample Receipt Date</u>	<u>Project Desc</u>	<u>Clients ID</u>	<u>Matrix</u>	<u>Sampling Date/Time</u>	<u>Sampled By</u>	<u>Sample #</u>	<u>Analysis Requested</u>
8/15/2013 1100	Soil Water Air Protection Enterprise (SWAPE) Bridgeton Sanitary Landfill Air Quality Assessment	U-1 K DNPH	Tube	8/14/2013	Client	65568	TO-11A
8/15/2013 1100	Soil Water Air Protection Enterprise (SWAPE) Bridgeton Sanitary Landfill Air Quality Assessment	U-2 W2 DNPH	Tube	8/14/2013	Client	65569	TO-11A
8/15/2013 1100	Soil Water Air Protection Enterprise (SWAPE) Bridgeton Sanitary Landfill Air Quality Assessment	D-1 W6E DNPH	Tube	8/14/2013	Client	65570	TO-11A
8/15/2013 1100	Soil Water Air Protection Enterprise (SWAPE) Bridgeton Sanitary Landfill Air Quality Assessment	D-1 W6W DNPH	Tube	8/14/2013	Client	65571	TO-11A

**TURN AROUND TIME:** Normal (10days)

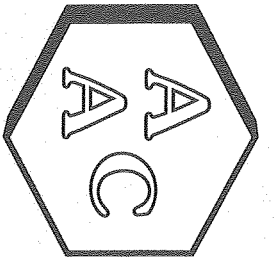
Lab Due Date: 8/22/2013

Total Samples: 4

**REMARKS:**

Samples received at 5.0°C. "Standard turn-around for all analyses. If possible deliver report within 2 weeks. Provide Level IV QC Package for all Analyses".

# Results



# Atmospheric Analysis & Consulting, Inc.


## LABORATORY ANALYSIS REPORT Analysis of Carbonyls by EPA Method TO-11A

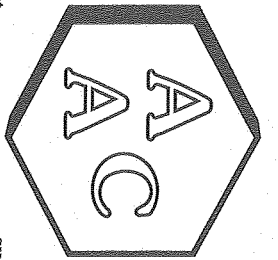
Client : SWAPE  
 Client Project Name : Bridgetown Sanitary Landfill Air Quality Assessment  
 AAC Project No. : 131099  
 Analyst : EG/HP  
 Units : ppbv

Sampling Date (s) : 08/14/2013  
 Receiving Date : 08/15/2013  
 Analysis Date : 08/22/2013  
 Reporting Date : 08/22/2013

Client ID	AAC Sample ID	Formaldehyde	Acetaldehyde	Acrolein	Acetone	Propionaldehyde	Crotonaldehyde	Methacrolein	MEK & Butyraldehyde	Benzaldehyde	Valeraldehyde	m-Tolaldehyde	Hexaldehyde
U-1 K DNPH	131099-65568	<SRL	0.204	<SRL	1.63	<SRL	<SRL	<SRL	0.220	<SRL	<SRL	<SRL	<SRL
		SRL	0.173	0.136	0.131	0.131	0.109	0.109	0.106	0.072	0.088	0.063	0.076
U-2 W2 DNPH	131099-65569	<SRL	0.222	<SRL	1.60	<SRL	<SRL	<SRL	0.229	<SRL	<SRL	<SRL	<SRL
		SRL	0.174	0.137	0.132	0.132	0.110	0.110	0.106	0.072	0.089	0.064	0.077
D-1 W6E DNPH	131099-65570	<SRL	0.159	<SRL	1.47	<SRL	<SRL	<SRL	0.175	<SRL	<SRL	<SRL	<SRL
		SRL	0.154	0.121	0.117	0.117	0.097	0.097	0.094	0.064	0.079	0.057	0.068
D-2 W6W DNPH	131099-65571	0.702	0.326	<SRL	0.938	<SRL	0.492	<SRL	0.130	<SRL	<SRL	<SRL	<SRL
		SRL	0.170	0.133	0.129	<SRL	0.107	0.107	0.104	0.070	0.087	0.062	0.075

<SRL - compound was analyzed for but not detected at or above the SRL (Sample Reporting Limit)

  
 Marcus Hueppe  
 Laboratory Director



# Atmospheric Analysis & Consulting, Inc.


## LABORATORY ANALYSIS REPORT Analysis of Carbonyls by EPA Method TO-11A

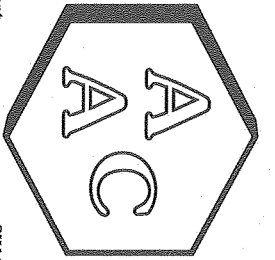
Client : SWAPE  
 Client Project Name : Bridgetown Sanitary Landfill Air Quality Assessment  
 AAC Project No. : 131099  
 Analyst : EG/HP  
 Units : ug/m<sup>3</sup>

Sampling Date (s) : 08/14/2013  
 Receiving Date : 08/15/2013  
 Analysis Date : 08/22/2013  
 Reporting Date : 08/22/2013

Client ID	AAC Sample ID	Formaldehyde	Acetaldehyde	Acrolein	Acetone	Propionaldehyde	Crotonaldehyde	Methacrolein	MEK & Butyraldehyde	Benzaldehyde	Valeraldehyde	m-Tolaldehyde	Hexaldehyde
U-1 K DNPH	131099-65568	<SRL	0.368	<SRL	3.87	<SRL	<SRL	<SRL	0.648	<SRL	<SRL	<SRL	<SRL
		0.312	0.312	0.312	0.312	0.312	0.312	0.312	0.312	0.312	0.312	0.312	0.312
U-2 W2 DNPH	131099-65569	<SRL	0.399	<SRL	3.80	<SRL	<SRL	<SRL	0.675	<SRL	<SRL	<SRL	<SRL
		0.314	0.314	0.314	0.314	0.314	0.314	0.314	0.314	0.314	0.314	0.314	0.314
D-1 W6E DNPH	131099-65570	<SRL	0.287	<SRL	3.48	<SRL	<SRL	<SRL	0.515	<SRL	<SRL	<SRL	<SRL
		0.278	0.278	0.278	0.278	0.278	0.278	0.278	0.278	0.278	0.278	0.278	0.278
D-2 W6W DNPH	131099-65571	0.862	0.587	<SRL	2.23	<SRL	<SRL	<SRL	0.383	<SRL	<SRL	<SRL	<SRL
		0.306	0.306	0.306	0.306	0.306	0.306	0.306	0.306	0.306	0.306	0.306	0.306

<SRL-compound was analyzed for but not detected at or above the SRL (Sample Reporting Limit)

  
 Marcus Hueppe  
 Laboratory Director



# Atmospheric Analysis & Consulting, Inc.

## LABORATORY ANALYSIS REPORT Analysis of Carbonyls by EPA Method TO-11A

Client : SWAPE  
 Client Project Name : Bridgetown Sanitary Landfill Air Quality Assessment  
 AAC Project No. : 131099  
 Analyst : EG/HP  
 Units : ug/sample

Sampling Date (s) : 08/14/2013  
 Receiving Date : 08/15/2013  
 Analysis Date : 08/22/2013  
 Reporting Date : 08/22/2013

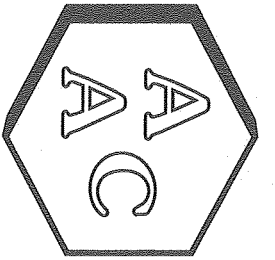
Client ID	AAC Sample ID	Formaldehyde	Acetaldehyde	Acrolein	Acetone	Propionaldehyde	Crotonaldehyde	Methacrolein	MEK & Butyraldehyde	Benzaldehyde	Valeraldehyde	m-Tolualdehyde	Hexaldehyde
U-1 K DNPH	131099-65568	<SRL	0.089	<SRL	0.931	<SRL	<SRL	<SRL	0.156	<SRL	<SRL	<SRL	<SRL
		SRL	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075
U-2 W2 DNPH	131099-65569	<SRL	0.095	<SRL	0.908	<SRL	<SRL	<SRL	0.161	<SRL	<SRL	<SRL	<SRL
		SRL	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075
D-1 W6E DNPH	131099-65570	<SRL	0.077	<SRL	0.939	<SRL	<SRL	<SRL	0.139	<SRL	<SRL	<SRL	<SRL
		SRL	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075
D-2 W6W DNPH	131099-65571	<SRL	0.144	<SRL	0.547	<SRL	<SRL	<SRL	0.094	<SRL	<SRL	<SRL	<SRL
		SRL	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075	0.075

<SRL-compound was analyzed for but not detected at or above the SRL (Sample Reporting Limit)

  
 Marcus Huepfe  
 Laboratory Director



# QA/QC Summary



# Atmospheric Analysis & Consulting, Inc.

## Quality Control/Quality Assurance Report

TO-11A

HPLC Calibration Verification of the 06/11/2013 Calibration

Analysis Date : 08/22/2013  
Analyst : HPEEG

Instrument ID : HPLC 01

Opening CCV

Standard Concentration (ug/ml)	Formaldehyde (ug/ml)	Acetaldehyde (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyde (ug/ml)	Crtonaldehyde (ug/ml)	Methacrolein (ug/ml)	MEK & Butyraldehyde (ug/ml)	Benzaldehyde (ug/ml)	Valeraldehyde (ug/ml)	m-Tolualdehyde (ug/ml)	Hexaldehyde (ug/ml)
2.50	2.30	2.33	2.31	2.33	2.35	2.31	2.32	4.65	2.35	2.32	2.30	2.34
Accuracy (%)*	92.0	93.2	92.4	93.2	94.0	92.4	92.8	93.0	94.0	92.8	92.0	93.6

Continuing CCV

Standard Concentration (ug/ml)	Formaldehyde (ug/ml)	Acetaldehyde (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyde (ug/ml)	Crtonaldehyde (ug/ml)	Methacrolein (ug/ml)	MEK & Butyraldehyde (ug/ml)	Benzaldehyde (ug/ml)	Valeraldehyde (ug/ml)	m-Tolualdehyde (ug/ml)	Hexaldehyde (ug/ml)
2.50	2.28	2.29	2.28	2.29	2.32	2.29	2.30	4.60	2.32	2.29	2.28	2.32
Accuracy (%)*	91.2	91.6	91.2	91.6	92.8	91.6	92.0	92.0	92.8	91.6	91.2	92.8

Continuing CCV

Standard Concentration (ug/ml)	Formaldehyde (ug/ml)	Acetaldehyde (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyde (ug/ml)	Crtonaldehyde (ug/ml)	Methacrolein (ug/ml)	MEK & Butyraldehyde (ug/ml)	Benzaldehyde (ug/ml)	Valeraldehyde (ug/ml)	m-Tolualdehyde (ug/ml)	Hexaldehyde (ug/ml)
2.50	2.34	2.36	2.35	2.37	2.38	2.35	2.38	4.72	2.39	2.35	2.35	2.39
Accuracy (%)*	93.6	94.4	94.0	94.8	95.2	94.0	95.2	94.4	95.6	94.0	94.0	95.6

Closing CCV

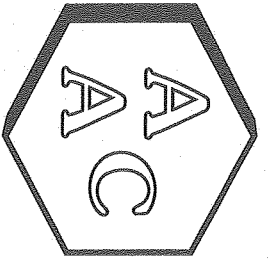
Standard Concentration (ug/ml)	Formaldehyde (ug/ml)	Acetaldehyde (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyde (ug/ml)	Crtonaldehyde (ug/ml)	Methacrolein (ug/ml)	MEK & Butyraldehyde (ug/ml)	Benzaldehyde (ug/ml)	Valeraldehyde (ug/ml)	m-Tolualdehyde (ug/ml)	Hexaldehyde (ug/ml)
2.50	2.30	2.34	2.33	2.34	2.35	2.33	2.34	4.69	2.36	2.32	2.32	2.36
Accuracy (%)*	92.0	93.6	93.2	93.6	94.0	93.2	93.6	93.8	94.4	92.8	92.8	94.4

Second Source

Standard Concentration (ug/ml)	Formaldehyde (ug/ml)	Acetaldehyde (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyde (ug/ml)	Crtonaldehyde (ug/ml)	Methacrolein (ug/ml)	MEK & Butyraldehyde (ug/ml)	Benzaldehyde (ug/ml)	Valeraldehyde (ug/ml)	m-Tolualdehyde (ug/ml)	Hexaldehyde (ug/ml)
2.50	2.29	2.30	2.29	2.31	2.34	2.29	2.30	4.63	2.33	2.30	2.30	2.33
Accuracy (%)*	91.6	92.0	91.6	92.4	93.6	91.6	92.0	92.6	93.2	92.0	92.0	93.2

\*Must be 100 ± 10%

*Marcus Hueppe*  
 Marcus Hueppe  
 Laboratory Director



# Atmospheric Analysis & Consulting, Inc.

## Quality Control/Quality Assurance Report TO-11A Laboratory Control Spike Analysis

Analysis Date : 08/22/2013

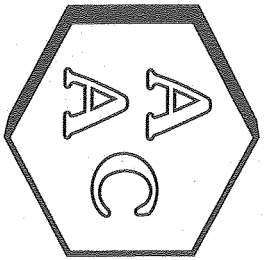
Analyst : HPEG

Instrument ID : HPLC 01

Airbytes	Formaldehyde (ug/ml)	Acetaldehyde (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyde (ug/ml)	Crotonaldehyde (ug/ml)	Methacrolein (ug/ml)	MEK & Butyraldehyde (ug/ml)	Benzaldehyde (ug/ml)	Valeraldehyde (ug/ml)	m-Tolualdehyde (ug/ml)	Hexaldehyde (ug/ml)
Laboratory Control Spike 1												
Sample Concentration (ug/ml)	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
Spike Concentration (ug/ml)	0.379	0.379	0.379	0.379	0.379	0.379	0.379	0.758	0.379	0.379	0.379	0.379
Spiked Sample Concentration (ug/ml)	0.378	0.366	0.383	0.365	0.393	0.371	0.418	0.718	0.368	0.371	0.383	0.377
Spike Recovery (%)*	100	96.7	101	96.4	104	97.8	110	94.7	97.1	97.9	101	100

\*Must be 100 ± 15%

  
Marcus Hueppe  
Laboratory Director



# Atmospheric Analysis & Consulting, Inc.

## Quality Control/Quality Assurance Report TO-11A Matrix Spike Analysis

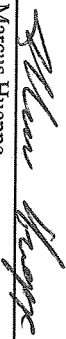
Analysis Date : 08/22/2013

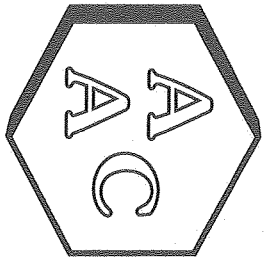
Analyst : HP/EG

Instrument ID : HPLC 01

Analytes	Formaldehyde (ug/mL)	Acetaldehyde (ug/mL)	Acrolein (ug/mL)	Acetone (ug/mL)	Propionaldehyde (ug/mL)	Crotonaldehyde (ug/mL)	Methacrolein (ug/mL)	MEK & Butyraldehyde (ug/mL)	Benzaldehyde (ug/mL)	Valeraldehyde (ug/mL)	m-Tolualdehyde (ug/mL)	Hexaldehyde (ug/mL)
Sample ID 131114-65647												
Sample Concentration (ug/mL)	0.000	0.007	0.000	0.063	0.004	0.007	0.000	0.008	0.006	0.009	0.015	0.005
Spike Concentration (ug/mL)	1.25	1.25	1.25	1.25	1.25	1.25	1.25	2.50	1.25	1.25	1.25	1.25
Spiked Sample Concentration (ug/mL)	1.15	1.15	1.18	1.25	1.19	1.20	1.23	2.37	1.20	1.19	1.21	1.17
Duplicate Spiked Sample Concentration (ug/mL)	1.15	1.15	1.18	1.25	1.19	1.19	1.23	2.35	1.20	1.19	1.20	1.17
Spike Recovery (%)*	92.0	91.4	94.4	95.0	94.9	95.5	98.4	94.5	95.6	94.5	95.6	93.2
Duplicate Spike Recovery (%)*	92.0	91.4	94.4	95.0	94.9	94.7	98.4	93.7	95.6	94.5	94.8	93.2
RPD**	0.0	0.0	0.0	0.0	0.0	0.8	0.0	0.8	0.0	0.0	0.8	0.0

\*Must be 100± 25%  
\*\* Must be ≤ 25%

  
 Marcus Hueppe  
 Laboratory Director



# Atmospheric Analysis & Consulting, Inc.

## Quality Control/Quality Assurance Report TO-11A Duplicate Analysis

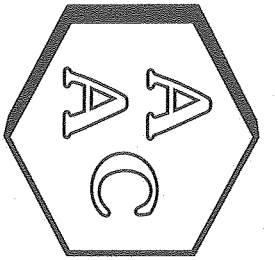
Analysis Date : 08/22/2013  
Analyst : HP/EG

Instrument ID : HPLC 01

Analytic	Formaldehyde (ug/mL)	Acetaldehyde (ug/mL)	Acrolein (ug/mL)	Acetone (ug/mL)	Propionaldehyde (ug/mL)	Crystalline (ug/mL)	Methacrolein (ug/mL)	MIBK & Butyraldehyde (ug/mL)	Benzaldehyde (ug/mL)	Valeraldehyde (ug/mL)	m- Toluylaldehyde (ug/mL)	Heptaldehyde (ug/mL)
Sample ID	131114-65647											
Sample Concentration (ug/mL)	ND	<RL	ND	0.126	<RL	<RL	ND	<RL	<RL	<RL	0.030	<RL
Duplicate Sample Concentration (ug/mL)	ND	<RL	ND	0.128	<RL	<RL	ND	<RL	<RL	<RL	0.032	<RL
RPD**	NA	NA	NA	1.5	NA	NA	NA	NA	NA	NA	5.5	NA
Sample ID	131114-65640											
Sample Concentration (ug/mL)	0.136	0.113	<RL	0.476	0.034	0.039	<RL	0.044	<RL	0.042	0.034	<RL
Duplicate Sample Concentration (ug/mL)	0.136	0.114	<RL	0.477	0.035	0.038	<RL	0.046	<RL	0.041	0.037	<RL
RPD**	0.1	0.7	NA	0.2	2.3	4.4	NA	4.5	NA	3.4	7.1	NA
Sample ID	131099-65568											
Sample Concentration (ug/mL)	<RL	0.030	ND	0.310	<RL	<RL	<RL	0.052	ND	<RL	ND	<RL
Duplicate Sample Concentration (ug/mL)	<RL	0.029	ND	0.308	<RL	<RL	<RL	0.055	ND	<RL	ND	<RL
RPD**	NA	2.1	NA	0.6	NA	NA	NA	4.9	NA	NA	NA	NA

\*\* Must be ≤ 20%  
<RL = less than the Reporting Limit  
ND = Not Detected  
NA = Not Applicable

  
 Marcus Hueppe  
 Laboratory Director



# Atmospheric Analysis & Consulting, Inc.

## Quality Control/Quality Assurance Report

TO-11A

System and Method Blank Analysis

Analysis Date : 08/22/2013  
Analyst : HP/EG

Instrument ID : HPLC 01

Analyte	Formaldehyd de (ug/ml)	Acetaldehyd de (ug/ml)	Acrolein (ug/ml)	Acetone (ug/ml)	Propionaldehyd de (ug/ml)	Crumaldehyd de (ug/ml)	Methacrol an (ug/ml)	MEK & Butyraldehyd de (ug/ml)	Benzaldehyd de (ug/ml)	Valeroldehyd e (ug/ml)	m- Toluoldehyd e (ug/ml)	Hexaldehyd e (ug/ml)
Opening Acetonitrile Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL
Method Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL
Continuing Acetonitrile Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL
Continuing Acetonitrile Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL
Closing Acetonitrile Blank	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL	<RL
Reporting Limit	0.025	0.025	0.025	0.025	0.025	0.025	0.025	0.025	0.025	0.025	0.025	0.025

RL = Reporting Limit

<RL = less than the Reporting Limit

Marcus Hueppe  
Laboratory Director

# Calibration Summary

File Name: C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL  
Version: 6

Creator: EG/HP  
Description: EPA TO-11

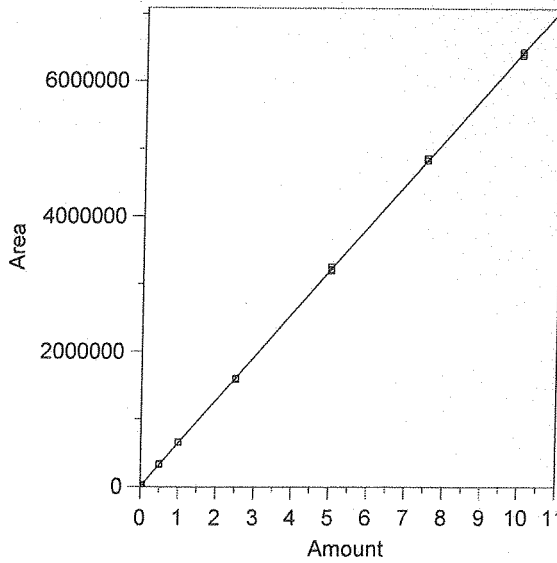
External standard calibration  
No injection volume correction  
No sample weight correction  
Area reject threshold: 1000  
Reference peak area reject threshold: 1000  
Amount units: ug/ml  
No default component

Method of calculating data point averages: Equal weight for all updates  
No calibration update report

All levels are normal data points.



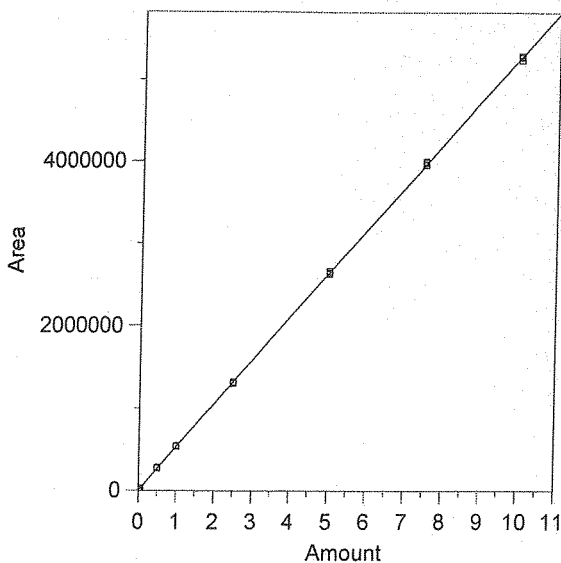
1 Formaldehyde



Expected retention time: 2.691 minutes  
 Search window: 0.1 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 $Y = 643471.9 X + 0$   
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999344  
 Average error: 1.932%  
 Average CF: 653816.2  
 RSD: 2.187%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	3408.358	681671.6	5.937	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	3325.732	665146.4	3.368	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	3366.37	673274.1	4.631	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	16194.19	647767.6	0.668	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	16179.21	647168.4	0.574	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	16232.13	649285.2	0.903	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	33956.52	679130.4	5.542	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	33691.46	673829.2	4.718	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	33082.47	661649.4	2.825	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	342894.1	685788.2	6.576	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011	6/13/2013 9:25:13 AM
11	0.5	326442.5	652885	1.463	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	328901	657802	2.227	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	659323.3	659323.3	2.463	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	652994.7	652994.7	1.480	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	653589.6	653589.6	1.572	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	1593441	637376.4	-0.947	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	1584858	633943.2	-1.481	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	1606654	642661.6	-0.126	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	3196329	639265.8	-0.654	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	3251038	650207.6	1.047	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	3213737	642747.4	-0.113	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	4829078	643877.1	0.063	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	4862793	648372.4	0.762	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	4865866	648782.1	0.825	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	6411879	641187.9	-0.355	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	6443770	644377	0.141	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	6389328	638932.8	-0.705	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

2 Acetaldehyde



Expected retention time: 3.293 minutes  
 Search window: 0.3 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0

Single peak quantification by area

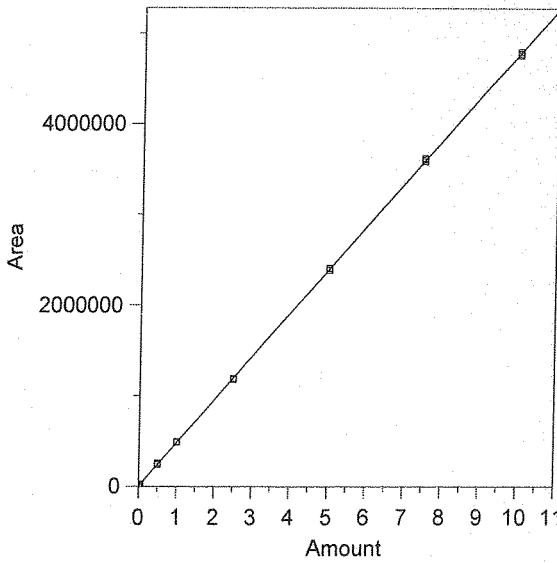
$Y = 528411.2 X + 0$

Linear fit with equal weighting, forced to origin

Coefficient of determination: 0.9999362  
 Average error: 1.513%  
 Average CF: 534461.4  
 RSD: 1.856%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	2781.029	556205.8	5.260	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	2639.74	527948	-0.088	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	2713.861	542772.3	2.718	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	13324.46	532978.4	0.864	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	13196.94	527877.6	-0.101	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	13319.8	532792	0.829	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	27360.74	547214.8	3.559	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	27153.9	543078	2.776	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	26813.58	536271.6	1.488	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	281357.4	562714.8	6.492	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	269068.5	538137	1.841	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	269886.8	539773.6	2.150	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	541849.3	541849.3	2.543	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	535548.3	535548.3	1.351	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	536037.3	536037.3	1.443	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	1304421	521768.4	-1.257	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	1298271	519308.4	-1.723	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	1316882	526752.8	-0.314	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	2629002	525800.4	-0.494	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	2663350	532670	0.806	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	2644688	528937.6	0.100	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	3962025	528270	-0.027	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	3985977	531463.6	0.578	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	4000077	533343.6	0.933	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	5275343	527534.3	-0.166	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	5292054	529205.4	0.150	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	5242042	524204.2	-0.796	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

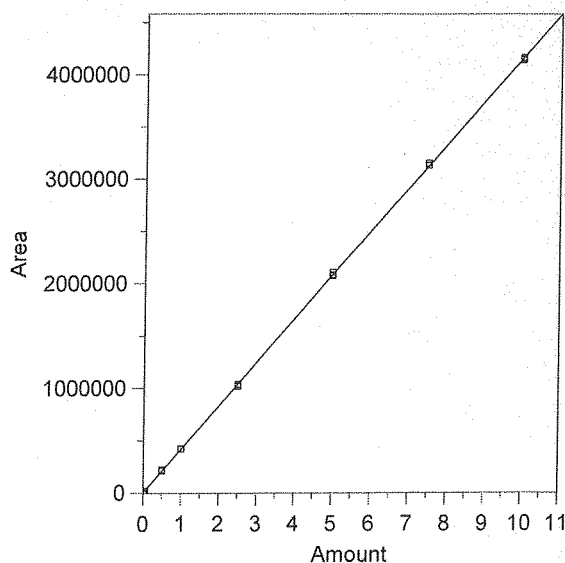
3 Acrolein



Expected retention time: 3.986 minutes  
 Search window: 0.2 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 479129.3 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999534  
 Average error: 1.520%  
 Average CF: 483677.6  
 RSD: 1.887%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	2333.751	466750.2	-2.584	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	2468.499	493699.8	3.041	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	2507.324	501464.8	4.662	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	11981.09	479243.6	0.024	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	12224.04	488961.6	2.052	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	11986.11	479444.4	0.066	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	24721.62	494432.4	3.194	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	24405.32	488106.4	1.874	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	24448.33	488966.6	2.053	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	254413	508826	6.198	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	242138.8	484277.6	1.075	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	244413.3	488826.6	2.024	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	490244.2	490244.2	2.320	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	484969.3	484969.3	1.219	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	484927.5	484927.5	1.210	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	1183948	473579.2	-1.158	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	1176976	470790.4	-1.740	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	1188060	475224	-0.815	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	2384550	476910	-0.463	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	2408812	481762.4	0.550	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	2403943	480788.6	0.346	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	3586781	478237.5	-0.186	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	3607897	481052.9	0.401	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	3621213	482828.4	0.772	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	4781989	478198.9	-0.194	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	4803733	480373.3	0.260	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	4764090	476409	-0.568	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

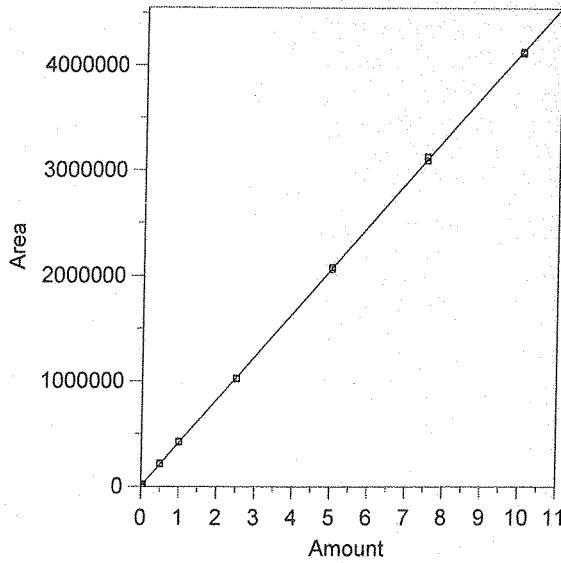
4 Acetone



Expected retention time: 4.179 minutes  
 Search window: 0.4 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 $Y = 416415.2 X + 0$   
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999405  
 Average error: 1.960%  
 Average CF: 422955.5  
 RSD: 2.274%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	2238.368	447673.6	7.507	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	2171.32	434264	4.286	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	2130.784	426156.8	2.339	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	10615.52	424620.8	1.971	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	10418.99	416759.6	0.083	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	10743.64	429745.6	3.201	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	21509.85	430197	3.310	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	21494.17	429883.4	3.234	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	21202.1	424042	1.832	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	223642.5	447285	7.413	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	212278.1	424556.2	1.955	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	214246.7	428493.4	2.901	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	426914.8	426914.8	2.521	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	422587.8	422587.8	1.482	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	420868.8	420868.8	1.070	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	1025289	410115.6	-1.513	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	1019650	407860	-2.054	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	1038209	415283.6	-0.272	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	2070625	414125	-0.550	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	2103557	420711.4	1.032	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	2080934	416186.8	-0.055	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	3130817	417442.3	0.247	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	3151942	420258.9	0.923	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	3133901	417853.5	0.345	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	4164627	416462.7	0.011	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	4152960	415296	-0.269	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	4141528	414152.8	-0.543	C:\Chromep\perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

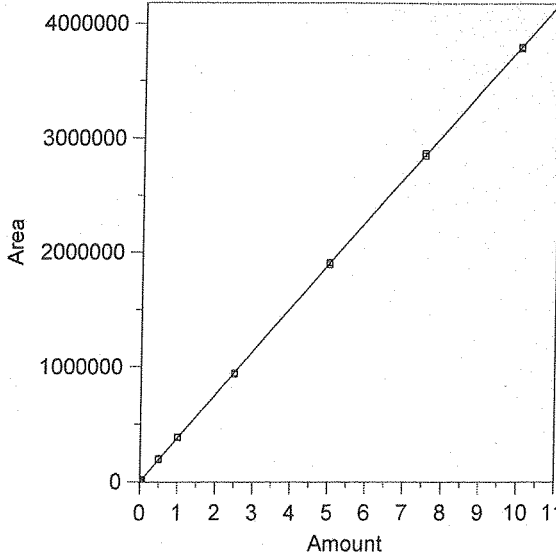
5 Propionaldehyde



Expected retention time: 4.45 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 413491.1 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999471  
 Average error: 1.596%  
 Average CF: 416900.7  
 RSD: 1.885%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	2017.78	403556	-2.403	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	2120.531	424106.2	2.567	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	2110.294	422058.8	2.072	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	10270.25	410810	-0.648	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	10463.54	418541.6	1.221	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	10129.01	405160.4	-2.015	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	21474.82	429496.4	3.871	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	21155.03	423100.6	2.324	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	21206.9	424138	2.575	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	218583.6	437167.2	5.726	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	211193.6	422387.2	2.151	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	211752.1	423504.2	2.422	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	425236.3	425236.3	2.841	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	416753.3	416753.3	0.789	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	420042.3	420042.3	1.584	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	1017464	406985.6	-1.573	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	1016426	406570.4	-1.674	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	1025583	410233.2	-0.788	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	2060419	412083.8	-0.340	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	2076059	415211.8	0.416	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	2079987	415997.4	0.606	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	3092180	412290.7	-0.290	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	3130391	417385.5	0.942	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	3118990	415865.3	0.574	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	4134994	413499.4	0.002	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	4125538	412553.8	-0.227	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	4115842	411584.2	-0.461	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

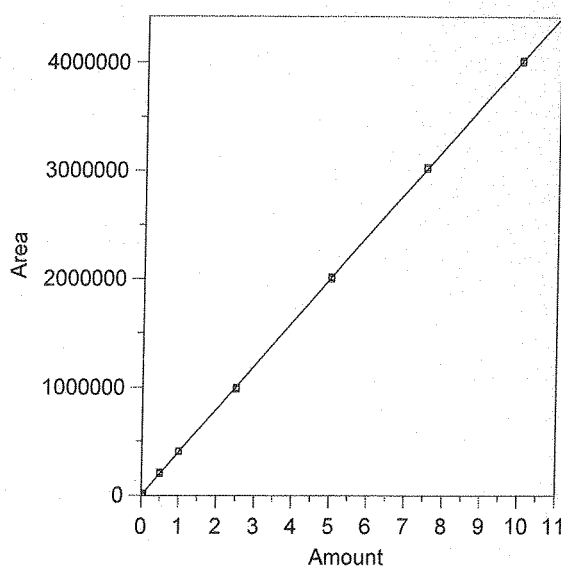
6 Crotonaldehyde



Expected retention time: 5.222 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 380159.3 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999585  
 Average error: 1.483%  
 Average CF: 383492.5  
 RSD: 2.008%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	1954.227	390845.4	2.811	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	1853.591	370718.2	-2.483	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	2006.267	401253.4	5.549	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	9500.876	380035	-0.033	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	9677.464	387098.6	1.825	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	9491.502	379660.1	-0.131	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	19612.18	392243.6	3.179	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	19377.38	387547.6	1.943	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	19062.43	381248.6	0.287	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	202454.9	404909.8	6.511	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	192705	385410	1.381	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	193772.2	387544.4	1.943	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	389857.3	389857.3	2.551	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	384117.9	384117.9	1.041	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	383544.6	383544.6	0.890	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	931914.6	372765.8	-1.945	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	933845.9	373538.3	-1.742	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	943212	377284.8	-0.756	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	1892096	378419.2	-0.458	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	1909513	381902.6	0.459	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	1905140	381028	0.229	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	2846489	379531.9	-0.165	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	2869201	382560.1	0.632	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	2868674	382489.9	0.613	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	3796762	379676.2	-0.127	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	3802364	380236.4	0.020	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	3788300	378830	-0.350	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

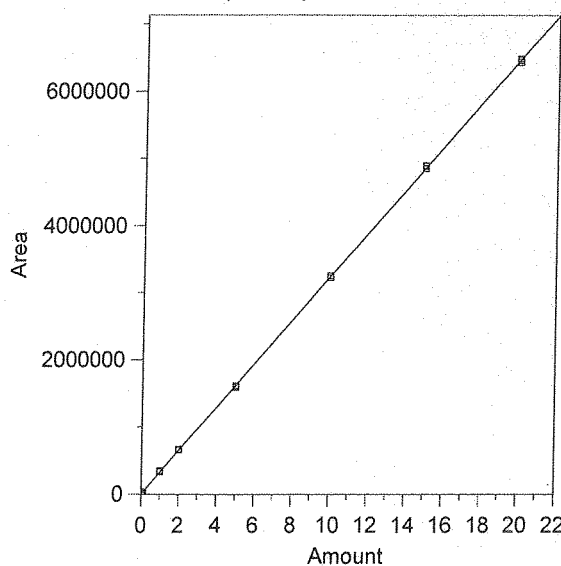
7 Methacrolein



Expected retention time: 5.582 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 402103.5 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999555  
 Average error: 1.593%  
 Average CF: 405803.3  
 RSD: 2.245%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	2046.606	409321.2	1.795	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	1941.76	388352	-3.420	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	2152.434	430486.8	7.059	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	10133.4	405336	0.804	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	10129.37	405174.8	0.764	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	10215.62	408624.8	1.622	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	20786.57	415731.4	3.389	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	20543.71	410874.2	2.181	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	20057.09	401141.8	-0.239	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	215228.8	430457.6	7.051	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	202696.3	405392.6	0.818	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	205546.4	411092.8	2.236	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	411542.4	411542.4	2.347	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	406241.6	406241.6	1.029	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	405954.8	405954.8	0.958	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	988074.2	395229.7	-1.709	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	985936.9	394374.8	-1.922	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	1000575	400230	-0.466	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	1996175	399235	-0.713	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	2017778	403555.6	0.361	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	2011258	402251.6	0.037	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	3018119	402415.9	0.078	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	3035201	404693.5	0.644	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	3035088	404678.4	0.640	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	4025485	402548.5	0.111	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	4010411	401041.1	-0.264	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	4007106	400710.6	-0.346	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

8 MEK & Butyraldehyde

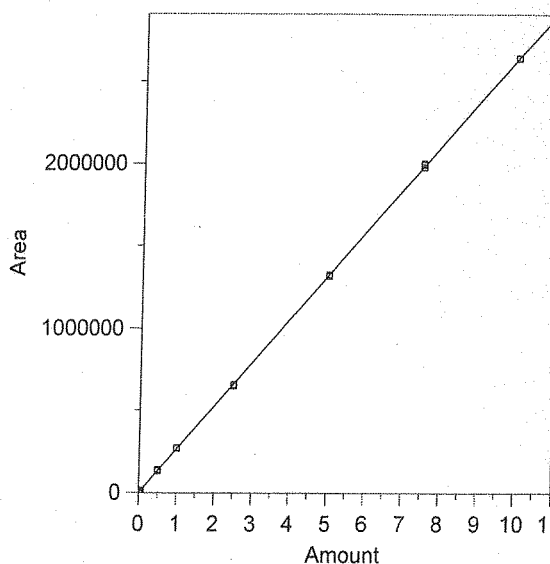


Expected retention time: 5.966 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 323725.3 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999455  
 Average error: 1.583%  
 Average CF: 327616.8  
 RSD: 1.902%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.01	3295.589	329558.9	1.802	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.01	3245.783	324578.3	0.264	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.01	3411.926	341192.6	5.396	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.05	16421.55	328431	1.454	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.05	16468.17	329363.4	1.742	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.05	16425.41	328508.2	1.477	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.1	33572.15	335721.5	3.706	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.1	33048.58	330485.8	2.088	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.1	32802.24	328022.4	1.327	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	1	346679.9	346679.9	7.091	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	1	328309.7	328309.7	1.416	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	1	331244	331244	2.323	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	2	664752.6	332376.3	2.672	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	2	656210.7	328105.3	1.353	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	2	654874.3	327437.2	1.147	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	5	1594650	318930	-1.481	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	5	1588729	317745.8	-1.847	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	5	1611625	322325	-0.433	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	10	3222726	322272.6	-0.449	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	10	3250978	325097.8	0.424	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	10	3247845	324784.5	0.327	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	15	4851557	323437.1	-0.089	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	15	4886541	325769.4	0.631	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	15	4893723	326248.2	0.779	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	20	6462091	323104.6	-0.192	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	20	6486439	324321.9	0.184	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	20	6432050	321602.5	-0.656	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM



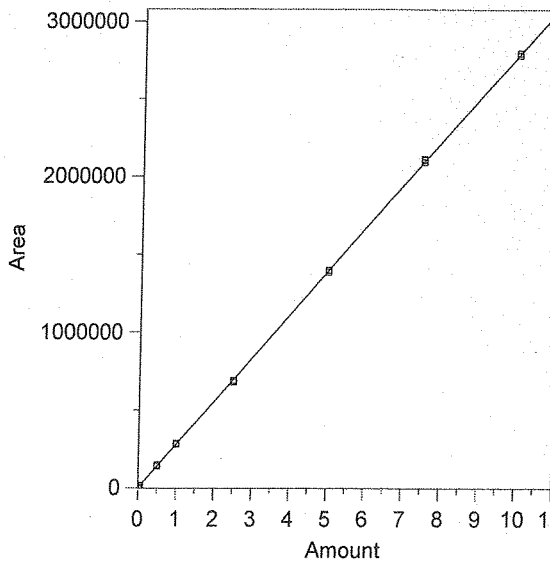
9 Benzaldehyde



Expected retention time: 6.375 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 $Y = 264532.8 X + 0$   
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999563  
 Average error: 1.328%  
 Average CF: 264682.8  
 RSD: 1.937%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	1271.628	254325.6	-3.859	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	1273.601	254720.2	-3.709	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	1382.946	276589.2	4.558	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	6506.057	260242.3	-1.622	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	6640.884	265635.3	0.417	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	6582.594	263303.8	-0.465	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	13464.09	269281.8	1.795	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	13243.82	264876.4	0.130	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	13138.29	262765.8	-0.668	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	138265.6	276531.2	4.536	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	131281	262562	-0.745	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	134626.8	269253.6	1.785	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	270957.1	270957.1	2.429	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	266032	266032	0.567	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	266541.5	266541.5	0.759	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	647530.3	259012.1	-2.087	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	648400.6	259360.3	-1.955	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	655172.9	262069.2	-0.931	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	1315721	263144.2	-0.525	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	1324105	264821	0.109	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	1325810	265162	0.238	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	1978244	263765.9	-0.290	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	2001675	266890	0.891	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	1993485	265798	0.478	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	2645002	264500.2	-0.012	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	2641466	264146.6	-0.146	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	2641496	264149.6	-0.145	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

10 Valeraldehyde



Expected retention time: 8.329 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0

High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0

Single peak quantification by area

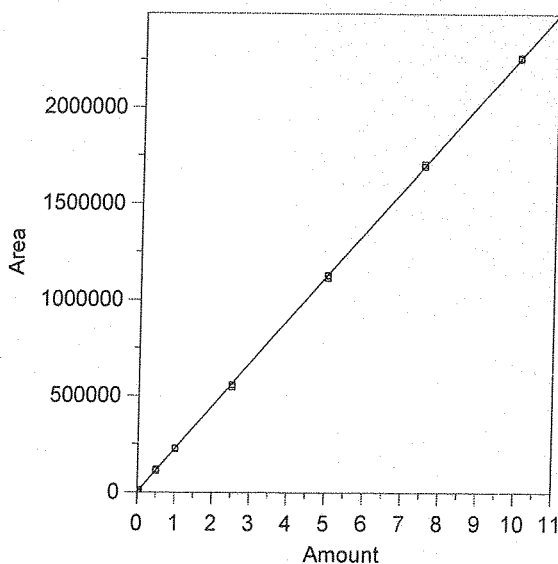
$Y = 279178.6 X + 0$

Linear fit with equal weighting, forced to origin

Coefficient of determination: 0.9999412  
 Average error: 1.679%  
 Average CF: 282346.4  
 RSD: 2.394%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	1484.718	296943.6	6.363	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	1440.975	288195	3.230	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	1478.778	295755.6	5.938	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	6975.488	279019.5	-0.057	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	6979.106	279164.2	-0.005	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	7027.2	281088	0.684	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	14633.92	292678.4	4.836	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	13928.58	278571.6	-0.217	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	14125.8	282516	1.195	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	149315.6	298631.2	6.968	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	140804	281608	0.870	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	142885.6	285771.2	2.361	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	286175.2	286175.2	2.506	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	280760.4	280760.4	0.567	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	280828.8	280828.8	0.591	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	681125.6	272450.3	-2.410	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	681786.7	272714.7	-2.315	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	691632.1	276652.8	-0.905	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	1386088	277217.6	-0.702	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	1398275	279655	0.171	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	1395245	279049	-0.046	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	2091460	278861.3	-0.114	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	2112053	281607.1	0.870	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	2105825	280776.7	0.572	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	2799263	279926.3	0.268	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	2783838	278383.8	-0.285	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	2783513	278351.3	-0.296	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

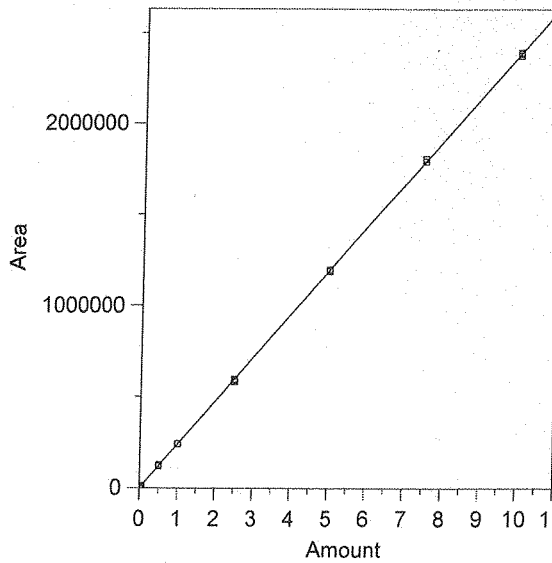
11 m-Tolualdehyde



Expected retention time: 8.864 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 225671.2 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.9999189  
 Average error: 1.845%  
 Average CF: 225055.9  
 RSD: 2.753%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	1205.651	241130.2	6.850	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	1133.419	226683.8	0.449	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	1115.328	223065.6	-1.155	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	5301.758	212070.3	-6.027	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	5574.778	222991.1	-1.188	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	5291.765	211670.6	-6.204	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	11500.01	230000.2	1.918	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	10974.64	219492.8	-2.738	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	11585.74	231714.8	2.678	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	118273.4	236546.8	4.819	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	112401.6	224803.2	-0.385	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	114850.5	229701	1.786	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	228083.9	228083.9	1.069	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	224659.6	224659.6	-0.448	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	224437.7	224437.7	-0.547	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	545930.8	218372.3	-3.234	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	545333.1	218133.3	-3.340	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	555975.6	222390.3	-1.454	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	1112742	222548.4	-1.384	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	1127437	225487.4	-0.081	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	1124909	224981.8	-0.305	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	1693020	225736	0.029	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	1703770	227169.3	0.664	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	1705210	227361.3	0.749	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	2261498	226149.8	0.212	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	2256941	225694.1	0.010	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	2254343	225434.3	-0.105	C:\Chromep perfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

12 Hexaldehyde



Expected retention time: 11.919 minutes  
 Search window: 0.5 minutes  
 No retention time reference component  
 Group number: 0  
 High alarm limit: 0  
 Low alarm limit: 0  
 Component constant: 0  
 Single peak quantification by area  
 Y = 238926 X + 0  
 Linear fit with equal weighting, forced to origin  
 Coefficient of determination: 0.999934  
 Average error: 1.592%  
 Average CF: 241305.9  
 RSD: 2.114%

Level	Amount	Response	Cal Factor	Error, %	Source	Date and time
1	0.005	1202.478	240495.6	0.657	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0002	6/13/2013 9:05:08 AM
2	0.005	1239.538	247907.6	3.759	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0003	6/13/2013 9:05:23 AM
3	0.005	1271.401	254280.2	6.426	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0004	6/13/2013 7:21:34 AM
4	0.025	5947.313	237892.5	-0.433	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0033	6/13/2013 7:22:04 AM
5	0.025	6068.35	242734	1.594	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0006	6/13/2013 7:22:28 AM
6	0.025	5933.474	237339	-0.664	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0007	6/13/2013 7:22:52 AM
7	0.05	12531.54	250630.8	4.899	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0008	6/13/2013 7:23:11 AM
8	0.05	12251.32	245026.4	2.553	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0009	6/13/2013 7:23:32 AM
9	0.05	12152.7	243054	1.728	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0010	6/13/2013 7:23:49 AM
10	0.5	126196	252392	5.636	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0011.BND	6/13/2013 9:25:13 AM
11	0.5	120634.2	241268.4	0.980	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0012	6/13/2013 7:24:58 AM
12	0.5	121148.8	242297.6	1.411	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0013	6/13/2013 7:25:12 AM
13	1	242571.7	242571.7	1.526	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0014	6/13/2013 7:25:25 AM
14	1	240813.4	240813.4	0.790	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0015	6/13/2013 7:25:42 AM
15	1	241116.1	241116.1	0.917	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0016	6/13/2013 7:25:59 AM
16	2.5	587005.4	234802.2	-1.726	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0017	6/13/2013 7:26:15 AM
17	2.5	579802.9	231921.2	-2.932	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0018	6/13/2013 7:26:31 AM
18	2.5	594270.9	237708.3	-0.510	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0019	6/13/2013 7:26:46 AM
19	5	1188236	237647.2	-0.535	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0020	6/13/2013 7:27:07 AM
20	5	1187571	237514.2	-0.591	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0021	6/13/2013 7:27:30 AM
21	5	1196602	239320.4	0.165	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0022	6/13/2013 7:27:52 AM
22	7.5	1792229	238963.9	0.016	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0023	6/13/2013 7:28:15 AM
23	7.5	1807473	240996.4	0.867	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0024	6/13/2013 7:28:35 AM
24	7.5	1806138	240818.4	0.792	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0025	6/13/2013 7:28:58 AM
25	10	2394549	239454.9	0.221	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0026	6/13/2013 7:29:18 AM
26	10	2387175	238717.5	-0.087	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0027	6/13/2013 7:29:40 AM
27	10	2375749	237574.9	-0.565	C:\Chromperfect 2\Data\HPLC #1\2013\061113(cal)\061113.0028	6/13/2013 7:29:57 AM

Chrom Perfect Sequence File

File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113(cal).SEQ

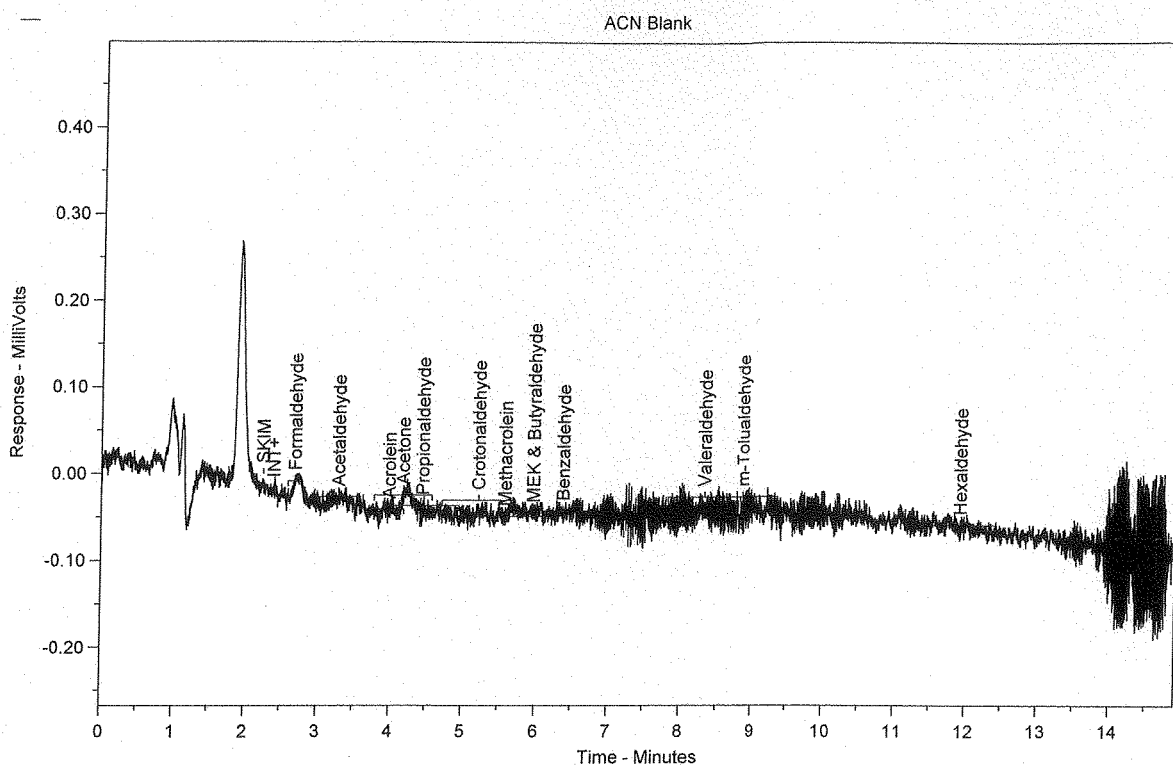
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Analyst=

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2	061113.0002.raw	061113 TO-11A.MET	STD 1-1 (.005ug/ml [PS061113-01]x2000)	2	1
3	061113.0003.raw	061113 TO-11A.MET	STD 1-2 (.005ug/ml [PS061113-01]x2000)	2	1
4	061113.0004.raw	061113 TO-11A.MET	STD 1-3 (.005ug/ml [PS061113-01]x2000)	2	1
5	061113.0005.raw	061113 TO-11A.MET	STD 2-1 (.025ug/ml [PS061113-01]x400)	3	1
6	061113.0006.raw	061113 TO-11A.MET	STD 2-2 (.025ug/ml [PS061113-01]x400)	3	1
7	061113.0007.raw	061113 TO-11A.MET	STD 2-3 (.025ug/ml [PS061113-01]x400)	3	1
8	061113.0008.raw	061113 TO-11A.MET	STD 3-1 (.050ug/ml [PS061113-01]x200)	4	1
9	061113.0009.raw	061113 TO-11A.MET	STD 3-2 (.050ug/ml [PS061113-01]x200)	4	1
10	061113.0010.raw	061113 TO-11A.MET	STD 3-3 (.050ug/ml [PS061113-01]x200)	4	1
11	061113.0011.raw	061113 TO-11A.MET	STD 4-1 (.5ug/ml [PS061113-01]x20)	5	1
12	061113.0012.raw	061113 TO-11A.MET	STD 4-2 (.5ug/ml [PS061113-01]x20)	5	1
13	061113.0013.raw	061113 TO-11A.MET	STD 4-3 (.5ug/ml [PS061113-01]x20)	5	1
14	061113.0014.raw	061113 TO-11A.MET	STD 5-1 (1.0ug/ml [PS061113-01]x10)	6	1
15	061113.0015.raw	061113 TO-11A.MET	STD 5-2 (1.0ug/ml [PS061113-01]x10)	6	1
16	061113.0016.raw	061113 TO-11A.MET	STD 5-3 (1.0ug/ml [PS061113-01]x10)	6	1
17	061113.0017.raw	061113 TO-11A.MET	STD 6-1 (2.5ug/ml [PS061113-01]x4)	7	1
18	061113.0018.raw	061113 TO-11A.MET	STD 6-2 (2.5ug/ml [PS061113-01]x4)	7	1
19	061113.0019.raw	061113 TO-11A.MET	STD 6-3 (2.5ug/ml [PS061113-01]x4)	7	1
20	061113.0020.raw	061113 TO-11A.MET	STD 7-1 (5.0ug/ml [PS061113-01]x2)	8	1
21	061113.0021.raw	061113 TO-11A.MET	STD 7-2 (5.0ug/ml [PS061113-01]x2)	8	1
22	061113.0022.raw	061113 TO-11A.MET	STD 7-3 (5.0ug/ml [PS061113-01]x2)	8	1
23	061113.0023.raw	061113 TO-11A.MET	STD 8-1 (7.5ug/ml [PS061113-01]x1.3)	9	1
24	061113.0024.raw	061113 TO-11A.MET	STD 8-2 (7.5ug/ml [PS061113-01]x1.3)	9	1
25	061113.0025.raw	061113 TO-11A.MET	STD 8-3 (7.5ug/ml [PS061113-01]x1.3)	9	1
26	061113.0026.raw	061113 TO-11A.MET	STD 9-1 (10.0ug/ml [PS061113-01]x1)	10	1
27	061113.0027.raw	061113 TO-11A.MET	STD 9-2 (10.0ug/ml [PS061113-01]x1)	10	1
28	061113.0028.raw	061113 TO-11A.MET	STD 9-3 (10.0ug/ml [PS061113-01]x1)	10	1
29	061113.0029.raw	061113 TO-11A.MET	ACN Blank	11	1
30	061113.0030.raw	061113 TO-11A.MET	CCV (2.5ug/ml [PS061113-01]x4)	12	1
31	061113.0031.raw	061113 TO-11A.MET	CCV (2.5ug/ml [PS061113-01]x4)	13	1
32	061113.0032.raw	061113 TO-11A.MET	ACN Blank	14	1
33	061113.0033.raw	061113 TO-11A.MET	STD 2-1 (.025ug/ml [PS061113-01]x400)	15	1

# Raw Data

Chrom Perfect Chromatogram Report



Sample Name = ACN Blank

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0001.RAW

Date Taken (end) = 8/22/2013 6:01:22 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 1

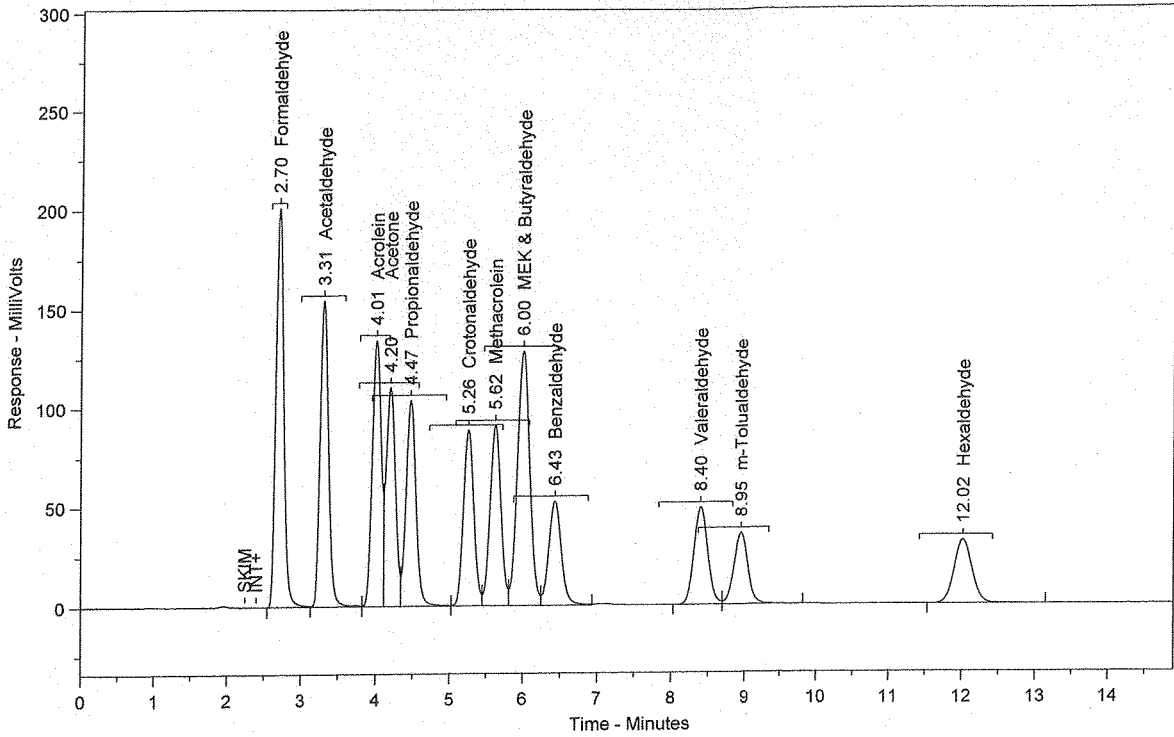
Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
Total Area = 0			Total Height = 0		Total Amount = 0			

Chrom Perfect Chromatogram Report

CCV 2.5 ug/mL (PS061113-01)



Sample Name = CCV 2.5 ug/mL (PS061113-01)

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0002.RAW

Date Taken (end) = 8/22/2013 6:18:01 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 2

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.70	Formaldehyde	2.3004	7.615	1480263	12.957	SBB	0.11
2	3.31	Acetaldehyde	2.3330	7.723	1232801	10.791	TBV	0.12
3	4.01	Acrolein	2.3062	7.634	1104955	9.672	TVV	0.14
4	4.20	Acetone	2.3295	7.711	970037	8.491	TVV	0.14
5	4.47	Propionaldehyde	2.3517	7.785	972425	8.512	TVV	0.14
6	5.26	Crotonaldehyde	2.3119	7.653	878893	7.693	TVV	0.15
7	5.62	Methacrolein	2.3249	7.696	934866	8.183	TVV	0.15
8	6.00	MEK & Butyraldehyde	4.6486	15.388	1504857	13.172	TVV	0.18
9	6.43	Benzaldehyde	2.3478	7.772	621062	5.436	TVB	0.18
10	8.40	Valeraldehyde	2.3166	7.669	646738	5.661	BV	0.20
11	8.95	m-Tolualdehyde	2.3022	7.621	519535	4.548	VB	0.22
12	12.02	Hexaldehyde	2.3356	7.732	558044	4.885	BB	0.27

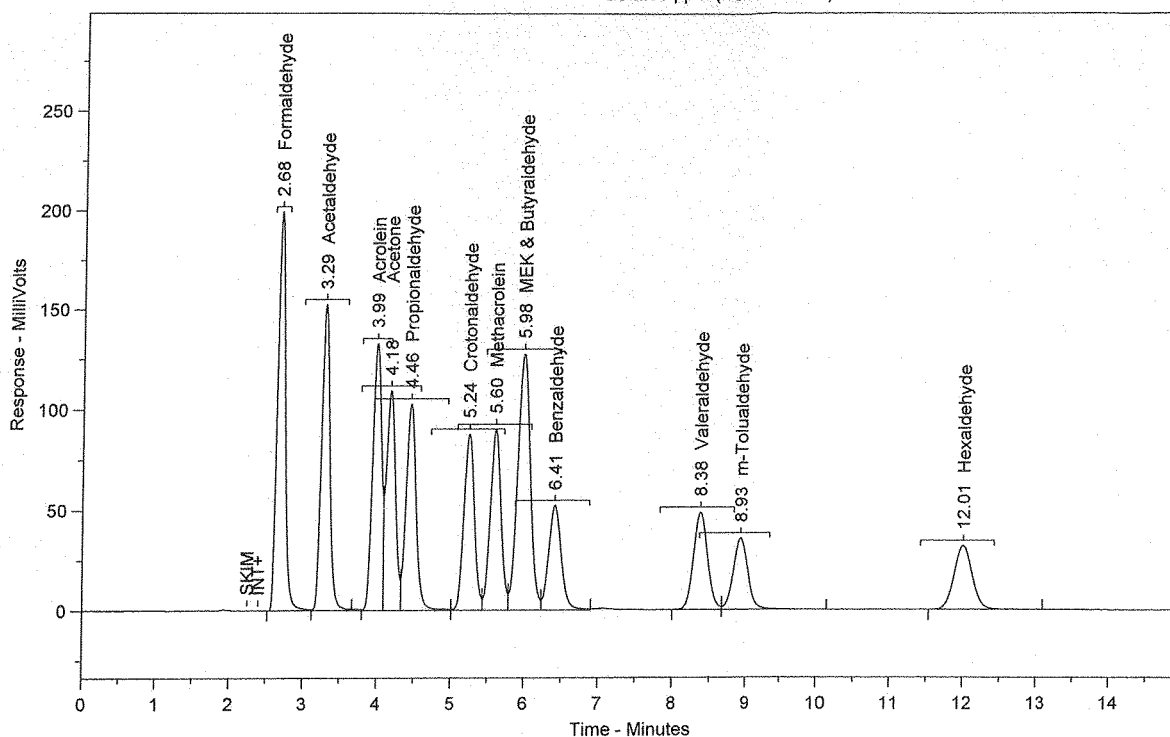
Total Area = 1.142447E+07

Total Height = 1181462

Total Amount = 30.20844



SS 2.50 ppm (PS011613-01)



Sample Name = SS 2.50 ppm (PS011613-01)

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0003.RAW

Date Taken (end) = 8/22/2013 6:34:40 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 3

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.68	Formaldehyde	2.2884	7.623	1472508	12.977	SBB	0.11
2	3.29	Acetaldehyde	2.3009	7.665	1215813	10.715	TBV	0.12
3	3.99	Acrolein	2.2936	7.640	1098935	9.685	TVV	0.14
4	4.18	Acetone	2.3054	7.680	960011	8.461	TVV	0.14
5	4.46	Propionaldehyde	2.3392	7.792	967227	8.524	TVV	0.14
6	5.24	Crotonaldehyde	2.2930	7.638	871716	7.682	TVV	0.15
7	5.60	Methacrolein	2.3029	7.671	926013	8.161	TVV	0.15
8	5.98	MEK & Butyraldehyde	4.6303	15.424	1498945	13.210	TVV	0.18
9	6.41	Benzaldehyde	2.3339	7.775	617392	5.441	TVB	0.18
10	8.38	Valeraldehyde	2.2998	7.661	642068	5.659	BV	0.20
11	8.93	m-Tolualdehyde	2.3048	7.678	520133	4.584	VB	0.22
12	12.01	Hexaldehyde	2.3274	7.753	556080	4.901	BB	0.27

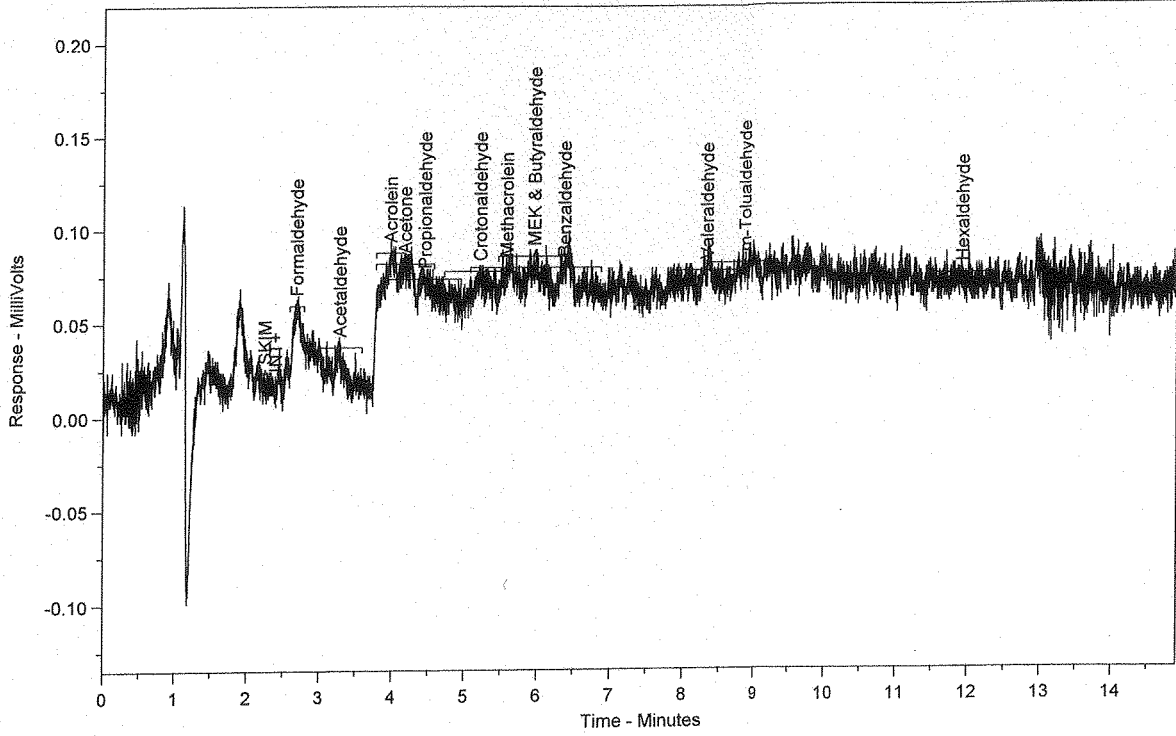
Total Area = 1.134684E+07

Total Height = 1170301

Total Amount = 30.0197

Chrom Perfect Chromatogram Report

TO-11 Method Blank



Sample Name = TO-11 Method Blank

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\0822\13TO-11\082213.0004.RAW

Date Taken (end) = 8/22/2013 6:51:20 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

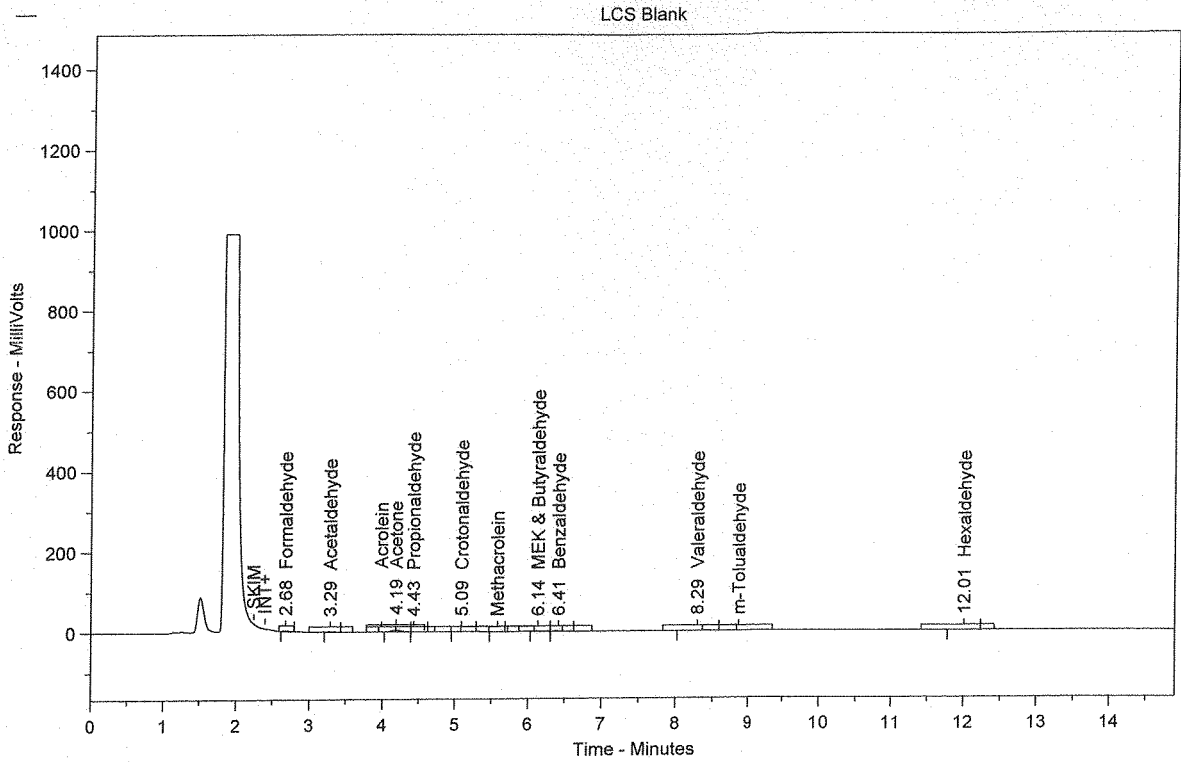
Vial Number = 4

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
Total Area = 0			Total Height = 0		Total Amount = 0			

Chrom Perfect Chromatogram Report



Sample Name = LCS Blank

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0005.RAW

Date Taken (end) = 8/22/2013 7:07:59 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 5

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.68	Formaldehyde	0.0095	4.615	6124	7.858	BB	0.13
2	3.29	Acetaldehyde	0.0107	5.204	5671	7.277	BB	0.15
3	4.19	Acetone	0.0881	42.731	36692	47.083	BV	0.13
4	4.43	Propionaldehyde	0.0040	1.953	1665	2.137	VB	0.14
5	5.09	Crotonaldehyde	0.0083	4.029	3159	4.053	BB	0.24
7	6.14	MEK & Butyraldehyde	0.0295	14.319	9558	12.265	BV	0.19
8	6.41	Benzaldehyde	0.0173	8.400	4582	5.879	VB	0.20
9	8.29	Valeraldehyde	0.0309	14.977	8622	11.064	BB	0.22
10	12.01	Hexaldehyde	0.0078	3.771	1858	2.384	BB	0.28

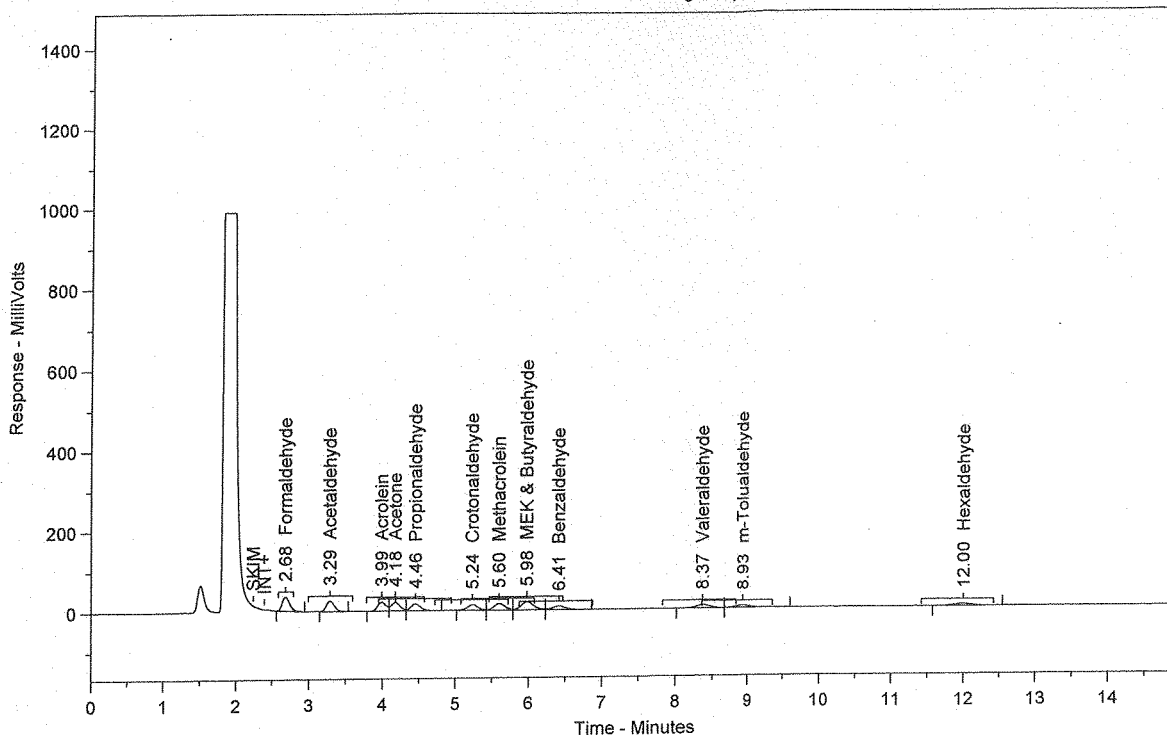
Total Area = 77930.99

Total Height = 8448.571

Total Amount = 0.2062064

Chrom Perfect Chromatogram Report

LCS .379ug/mL (PS011013-01)



Sample Name = LCS .379ug/mL (PS011013-01)

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0006.RAW

Date Taken (end) = 8/22/2013 7:24:37 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 6

Injection Volume = 10

Dilution Factor = 1

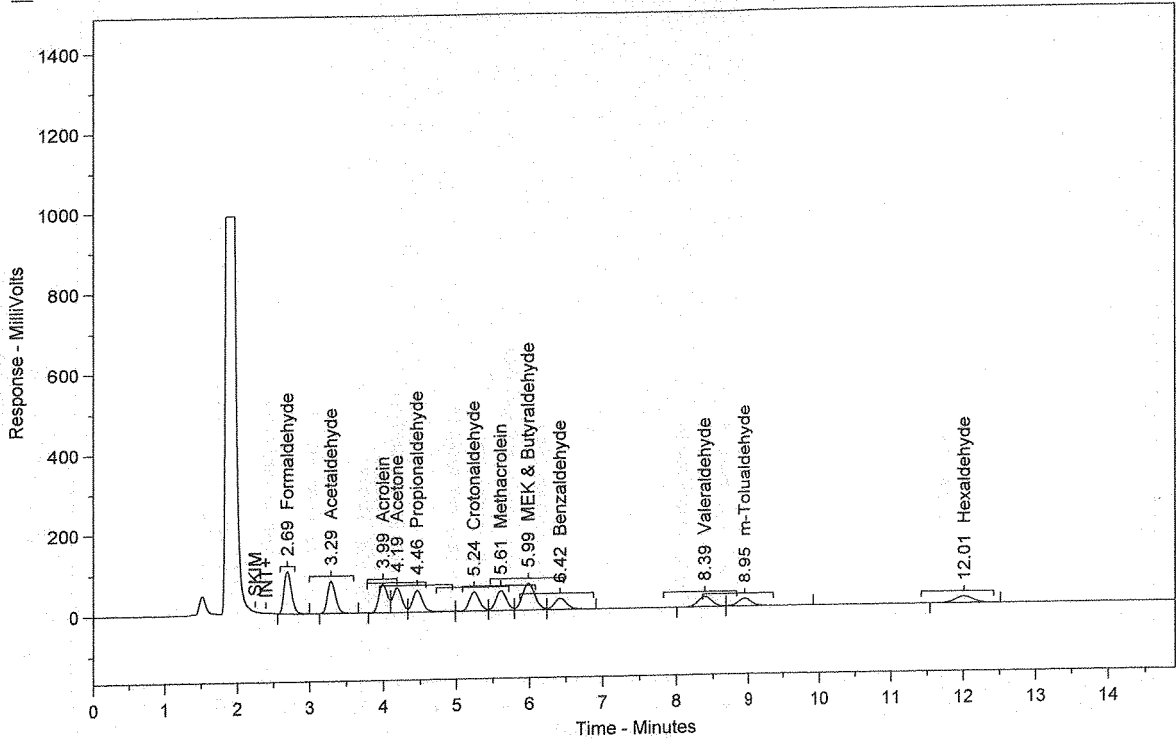
Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.68	Formaldehyde	0.3876	7.605	249418	12.914	BB	0.11
2	3.29	Acetaldehyde	0.3771	7.399	199273	10.317	BB	0.12
3	3.99	Acrolein	0.3829	7.512	183450	9.498	BV	0.14
4	4.18	Acetone	0.4533	8.894	188757	9.773	VV	0.13
5	4.46	Propionaldehyde	0.3966	7.782	163990	8.491	VB	0.14
6	5.24	Crotonaldehyde	0.3788	7.432	143996	7.455	BV	0.15
7	5.60	Methacrolein	0.4184	8.209	168231	8.710	VV	0.15
8	5.98	MEK & Butyraldehyde	0.7472	14.660	241883	12.524	VV	0.18
9	6.41	Benzaldehyde	0.3852	7.558	101898	5.276	VB	0.18
10	8.37	Valeraldehyde	0.4017	7.881	112139	5.806	BV	0.21
11	8.93	m-Tolualdehyde	0.3832	7.519	86481	4.478	VB	0.22
12	12.00	Hexaldehyde	0.3847	7.548	91910	4.759	BB	0.27

Total Area = 1931426

Total Height = 201370.8

Total Amount = 5.096613

MS 131114-65647 1.25 ppm [(PS061113-01x2)]



Sample Name = MS 131114-65647 1.25 ppm [(PS061113-01x2)]

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\0822\13TO-11\082213.0007.RAW

Date Taken (end) = 8/22/2013 7:41:16 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889  
Injection Volume = 10

Vial Number = 7  
Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.69	Formaldehyde	1.1506	7.426	740375	12.655	BB	0.11
2	3.29	Acetaldehyde	1.1549	7.453	610267	10.431	BB	0.12
3	3.99	Acrolein	1.1845	7.644	567529	9.701	BV	0.14
4	4.19	Acetone	1.2534	8.089	521928	8.921	VV	0.13
5	4.46	Propionaldehyde	1.1926	7.696	493118	8.429	VV	0.14
6	5.24	Crotonaldehyde	1.1958	7.717	454592	7.770	VV	0.15
7	5.61	Methacrolein	1.2320	7.951	495411	8.468	VV	0.15
8	5.99	MEK & Butyraldehyde	2.3683	15.284	766684	13.105	VV	0.18
9	6.42	Benzaldehyde	1.1951	7.713	316152	5.404	VB	0.18
10	8.39	Valeraldehyde	1.1889	7.673	331916	5.673	BV	0.21
11	8.95	m-Tolualdehyde	1.2075	7.792	272488	4.658	VB	0.22
12	12.01	Hexaldehyde	1.1716	7.561	279916	4.785	BB	0.27

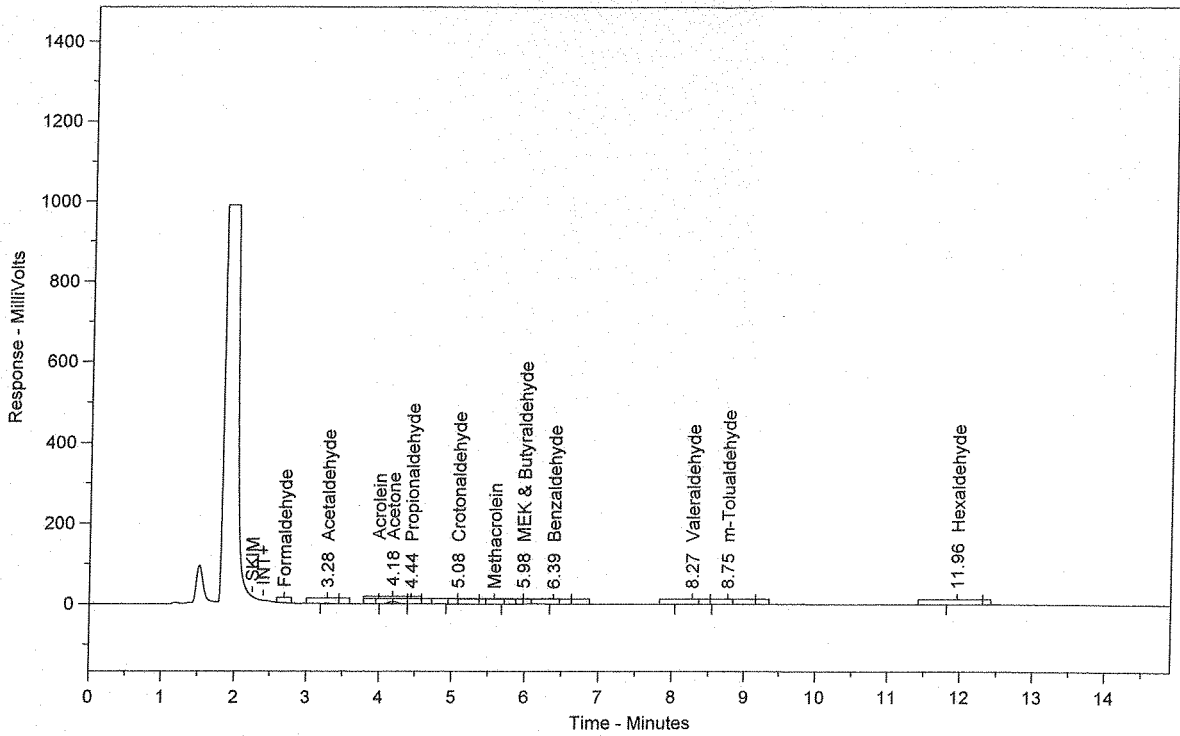
Total Area = 5850375

Total Height = 609053.1

Total Amount = 15.49517

Chrom Perfect Chromatogram Report

131114-65647 dup



Sample Name = 131114-65647 dup

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0010.RAW

Date Taken (end) = 8/22/2013 8:31:17 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 10

Injection Volume = 10

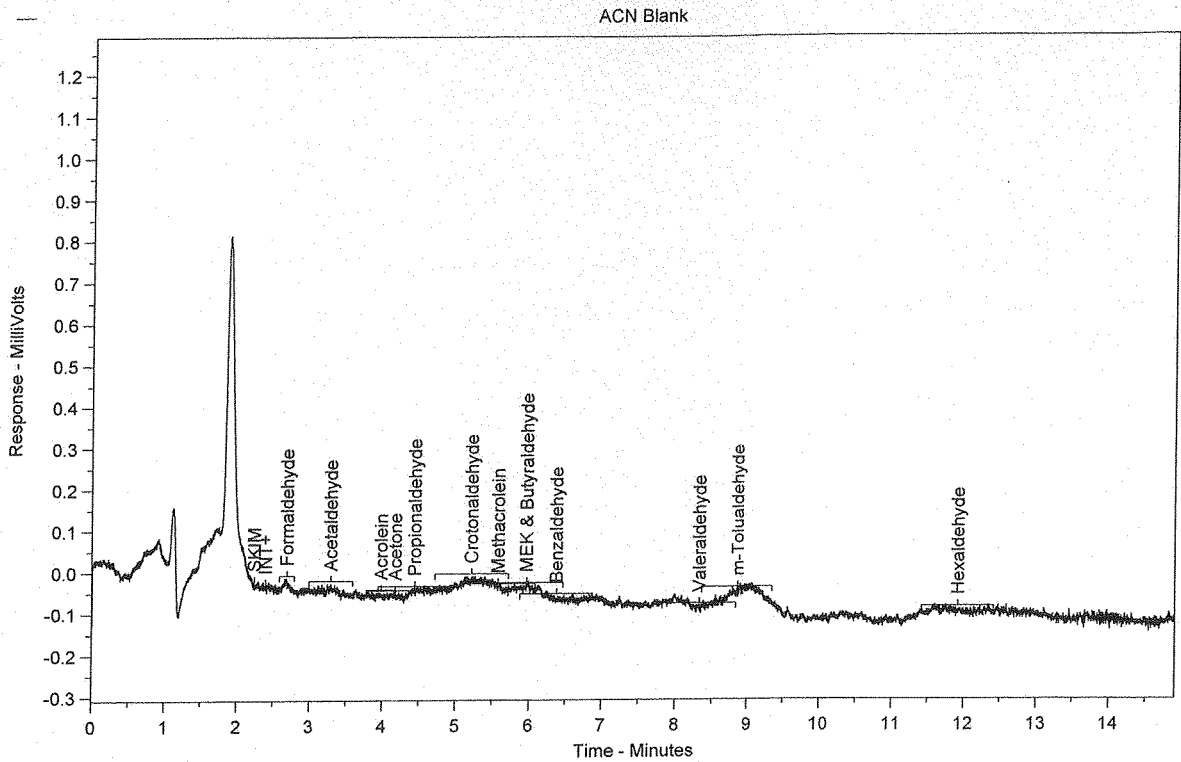
Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	3.28	Acetaldehyde	0.0137	5.433	7259	7.817	BB	0.16
2	4.18	Acetone	0.1281	50.673	53357	57.457	BV	0.13
3	4.44	Propionaldehyde	0.0089	3.505	3665	3.947	VB	0.12
4	5.08	Crotonaldehyde	0.0135	5.322	5116	5.509	BB	0.28
5	5.98	MEK & Butyraldehyde	0.0197	7.797	6382	6.872	BB	0.17
6	6.39	Benzaldehyde	0.0106	4.197	2807	3.023	BB	0.16
7	8.27	Valeraldehyde	0.0189	7.462	5268	5.673	BV	0.23
8	8.75	m-Tolualdehyde	0.0317	12.548	7160	7.711	VB	0.25
9	11.96	Hexaldehyde	0.0077	3.064	1851	1.993	BB	0.29

Total Area = 92865.34

Total Height = 9758.865

Total Amount = 0.2528643



Sample Name = ACN Blank

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213\TO-11\082213.0012.RAW

Date Taken (end) = 8/22/2013 9:04:35 AM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 12

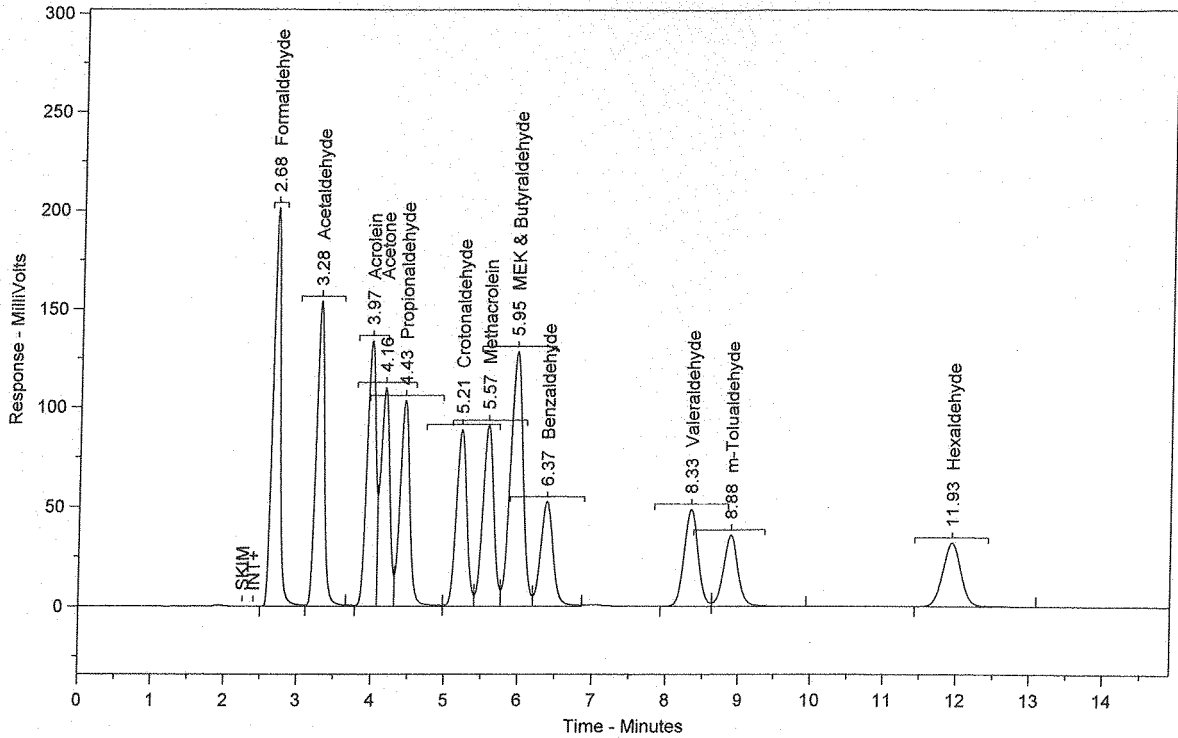
Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
Total Area = 0			Total Height = 0		Total Amount = 0			

Chrom Perfect Chromatogram Report

CCV 2.5 ug/mL (PS011613-01)



Sample Name = **CCV 2.5 ug/mL (PS011613-01)**

Instrument = HPLC #1

Raw File Name = C:\Chromep perfect 2\Data\HPLC #1\2013\082213TO-11\082213.0013.RAW

Date Taken (end) = 8/22/2013 9:21:14 AM

Method File Name = C:\Chromep perfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromep perfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 13

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.68	Formaldehyde	2.2762	7.623	1464689	12.977	SBB	0.11
2	3.28	Acetaldehyde	2.2873	7.660	1208632	10.708	TBV	0.12
3	3.97	Acrolein	2.2809	7.639	1092832	9.682	TVV	0.14
4	4.16	Acetone	2.2915	7.675	954235	8.454	TVV	0.13
5	4.43	Propionaldehyde	2.3170	7.760	958077	8.489	TVV	0.14
6	5.21	Crotonaldehyde	2.2904	7.671	870730	7.715	TVV	0.15
7	5.57	Methacrolein	2.3036	7.715	926279	8.207	TVV	0.15
8	5.95	MEK & Butyraldehyde	4.5994	15.404	1488945	13.192	TVV	0.17
9	6.37	Benzaldehyde	2.3232	7.781	614552	5.445	TVB	0.18
10	8.33	Valeraldehyde	2.2862	7.657	638269	5.655	BV	0.20
11	8.88	m-Tolualdehyde	2.2828	7.645	515155	4.564	VB	0.22
12	11.93	Hexaldehyde	2.3201	7.770	554331	4.911	BB	0.27

Total Area = 1.128673E+07

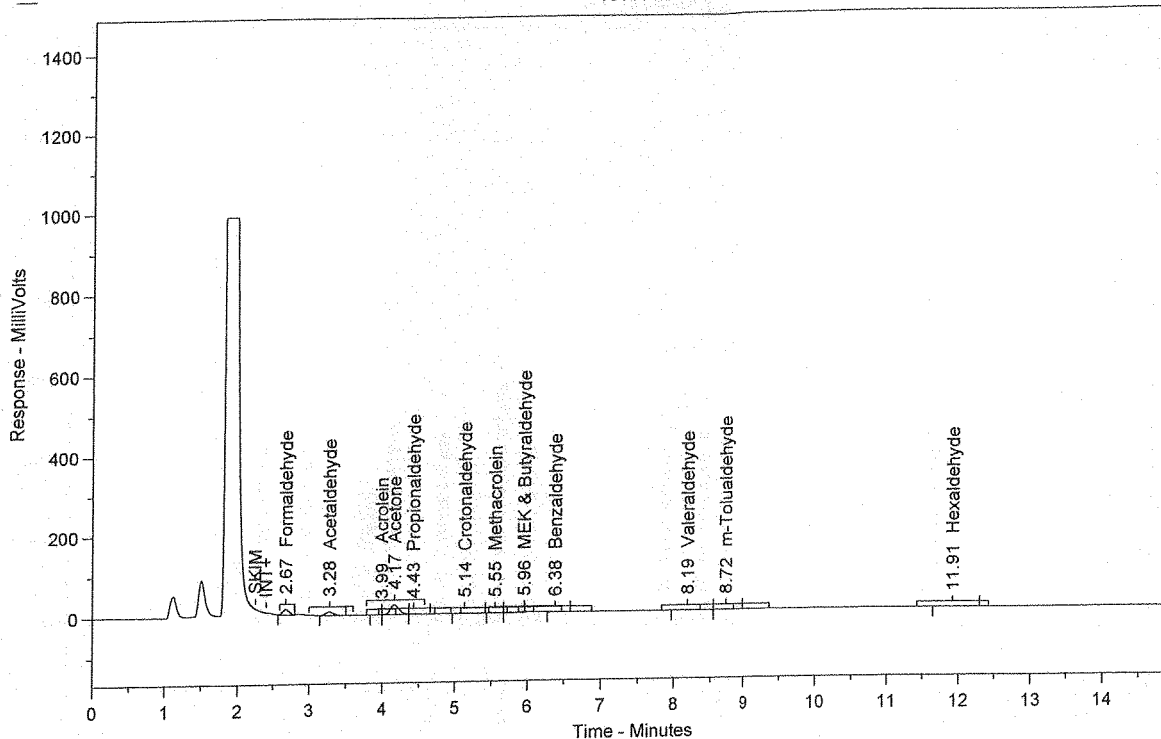
Total Height = 1179057

Total Amount = 29.85867



Chrom Perfect Chromatogram Report

131114-65640



Sample Name = 131114-65640

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0014.RAW  
 Date Taken (end) = 8/22/2013 9:37:53 AM

Method File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0014.BND  
 Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0014.BND  
 Concentration Units = ug/ml

Run Time = 14.89889  
 Injection Volume = 10

Vial Number = 14  
 Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.67	Formaldehyde	0.1364	14.063	87791	20.750	BB	0.10
2	3.28	Acetaldehyde	0.1132	11.670	59826	14.140	BB	0.12
3	3.99	Acrolein	0.0042	0.432	2010	0.475	BV	0.08
4	4.17	Acetone	0.4757	49.037	198105	46.822	VV	0.13
5	4.43	Propionaldehyde	0.0342	3.530	14159	3.347	VB	0.14
6	5.14	Crotonaldehyde	0.0394	4.063	14985	3.542	BB	0.20
7	5.55	Methacrolein	0.0058	0.597	2330	0.551	BV	0.14
8	5.96	MEK & Butyraldehyde	0.0438	4.517	14186	3.353	VB	0.11
9	6.38	Benzaldehyde	0.0167	1.725	4428	1.047	BB	0.19
10	8.19	Valeraldehyde	0.0424	4.370	11837	2.798	BV	0.27
11	8.72	m-Tolualdehyde	0.0342	3.527	7722	1.825	VB	0.29
12	11.91	Hexaldehyde	0.0239	2.468	5720	1.352	BB	0.30

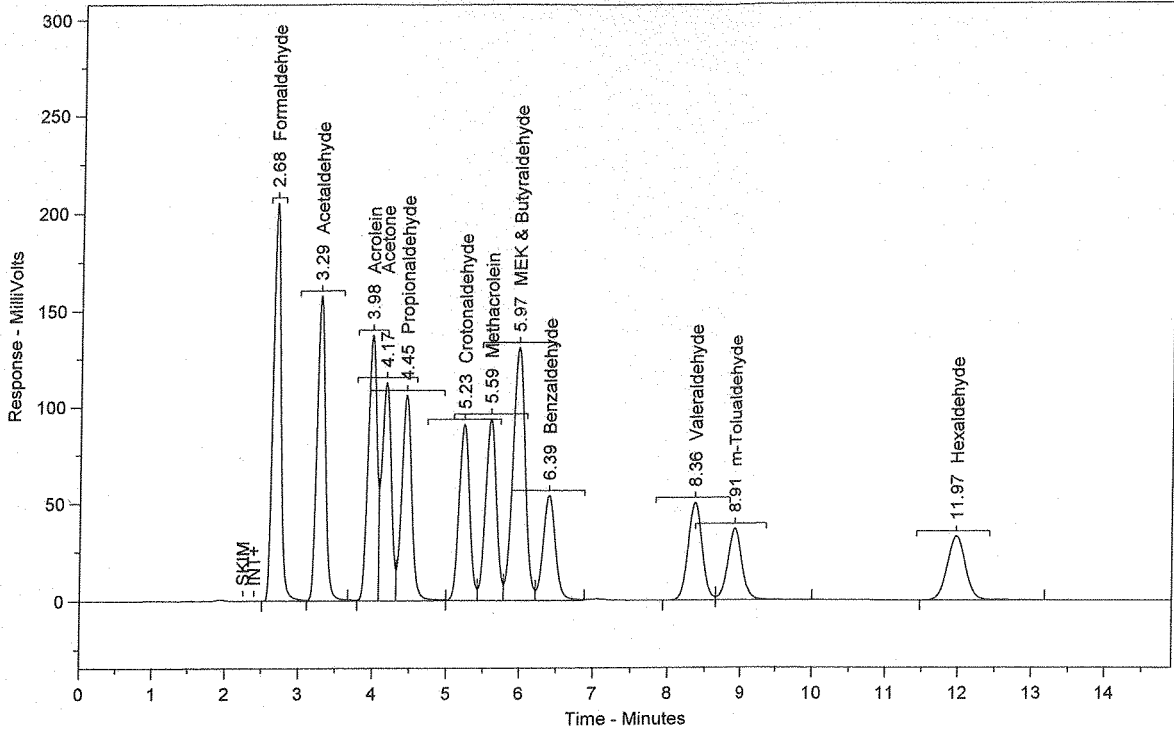
Total Area = 423099.3

Total Height = 50735.92

Total Amount = 0.9701592

Chrom Perfect Chromatogram Report

CCV 2.5 ug/mL (PS011613-01)



Sample Name = CCV 2.5 ug/mL (PS011613-01)

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0023.RAW

Date Taken (end) = 8/22/2013 12:07:46 PM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 23

Injection Volume = 10

Dilution Factor = 1

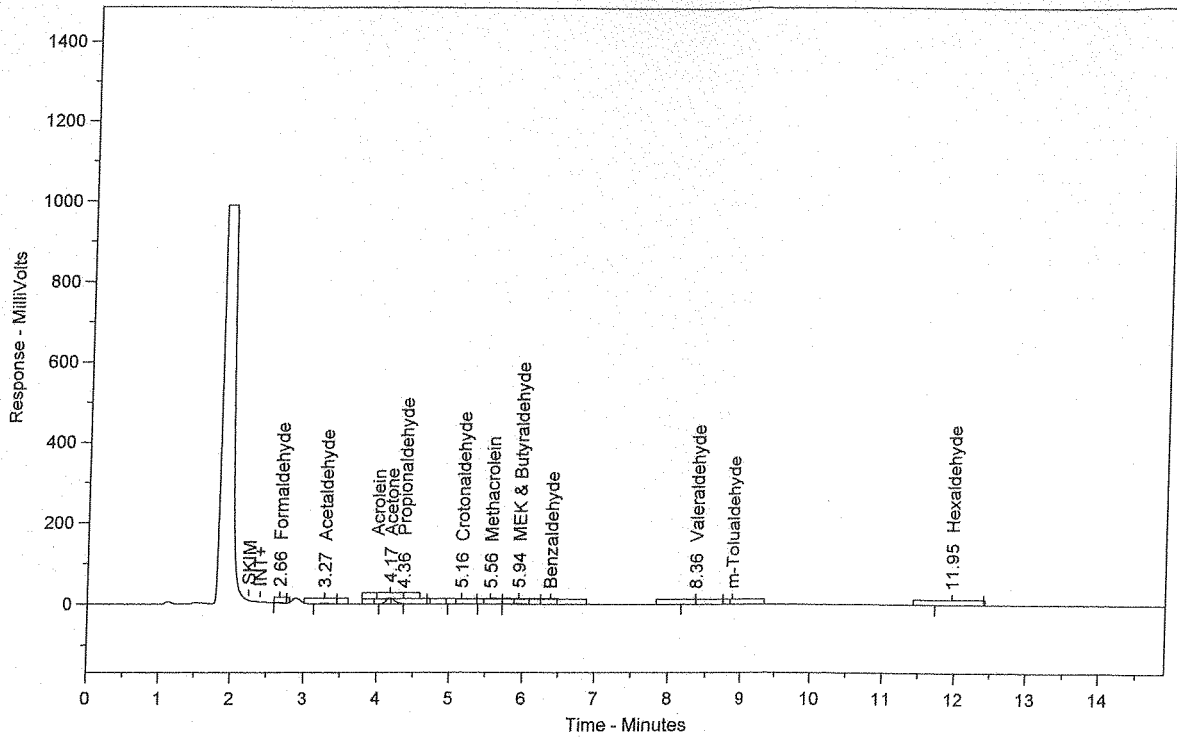
Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.68	Formaldehyde	2.3380	7.606	1504441	12.946	SBB	0.11
2	3.29	Acetaldehyde	2.3581	7.671	1246023	10.723	TBV	0.12
3	3.98	Acrolein	2.3486	7.640	1125268	9.684	TVV	0.14
4	4.17	Acetone	2.3713	7.714	987444	8.497	TVV	0.14
5	4.45	Propionaldehyde	2.3844	7.757	985946	8.485	TVV	0.14
6	5.23	Crotonaldehyde	2.3547	7.660	895143	7.703	TVV	0.15
7	5.59	Methacrolein	2.3841	7.756	958649	8.250	TVV	0.15
8	5.97	MEK & Butyraldehyde	4.7196	15.354	1527840	13.148	TVV	0.18
9	6.39	Benzaldehyde	2.3881	7.769	631732	5.436	TVB	0.18
10	8.36	Valeraldehyde	2.3506	7.647	656248	5.647	BV	0.20
11	8.91	m-Tolualdehyde	2.3542	7.659	531276	4.572	VB	0.22
12	11.97	Hexaldehyde	2.3875	7.767	570447	4.909	BB	0.27

Total Area = 1.162046E+07

Total Height = 1206828

Total Amount = 30.73916

131099-65568



Sample Name = 131099-65568

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0024.RAW

Date Taken (end) = 8/22/2013 12:24:25 PM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 24

Injection Volume = 10

Dilution Factor = 1

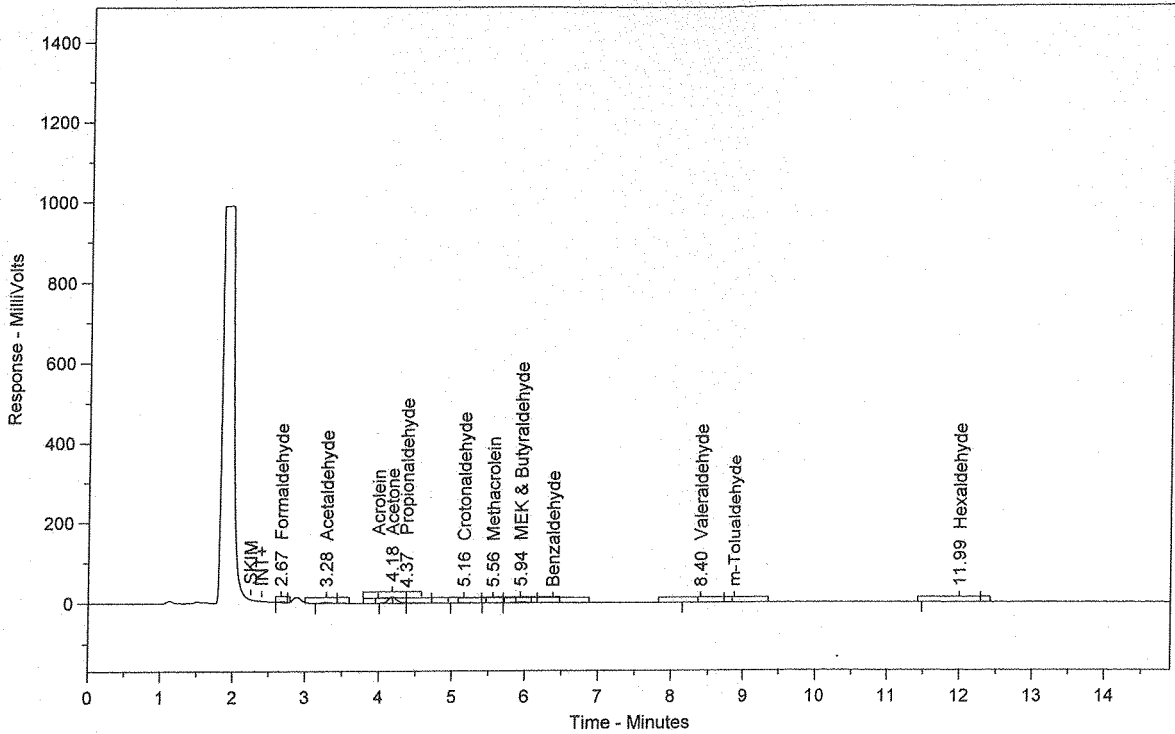
Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.66	Formaldehyde	0.0122	2.584	7868	4.054	BB	0.13
2	3.27	Acetaldehyde	0.0295	6.242	15606	8.041	BB	0.12
3	4.17	Acetone	0.3102	65.570	129185	66.560	SBB	0.13
4	4.36	Propionaldehyde	0.0198	4.177	8172	4.211	TBB	0.12
5	5.16	Crotonaldehyde	0.0204	4.307	7746	3.991	BB	0.17
6	5.56	Methacrolein	0.0081	1.722	3276	1.688	BV	0.14
7	5.94	MEK & Butyraldehyde	0.0520	10.997	16843	8.678	VB	0.17
8	8.36	Valeraldehyde	0.0103	2.182	2882	1.485	BB	0.40
9	11.95	Hexaldehyde	0.0105	2.218	2508	1.292	BB	0.29

Total Area = 194087

Total Height = 22630.36

Total Amount = 0.473129

131099-65568 dup



Sample Name = 131099-65568 dup

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0025.RAW

Date Taken (end) = 8/22/2013 12:41:03 PM

Method File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0025.BND

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0025.BND

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 25

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.67	Formaldehyde	0.0120	2.549	7725	4.006	BB	0.12
2	3.28	Acetaldehyde	0.0289	6.141	15282	7.925	BB	0.12
3	4.18	Acetone	0.3084	65.491	128429	66.601	BV	0.13
4	4.37	Propionaldehyde	0.0190	4.031	7848	4.070	VV	0.12
5	5.16	Crotonaldehyde	0.0199	4.232	7576	3.929	VB	0.17
6	5.56	Methacrolein	0.0072	1.525	2888	1.498	BV	0.14
7	5.94	MEK & Butyraldehyde	0.0546	11.597	17679	9.168	VB	0.17
8	8.40	Valeraldehyde	0.0104	2.206	2900	1.504	BB	0.34
9	11.99	Hexaldehyde	0.0105	2.229	2508	1.300	BB	0.28

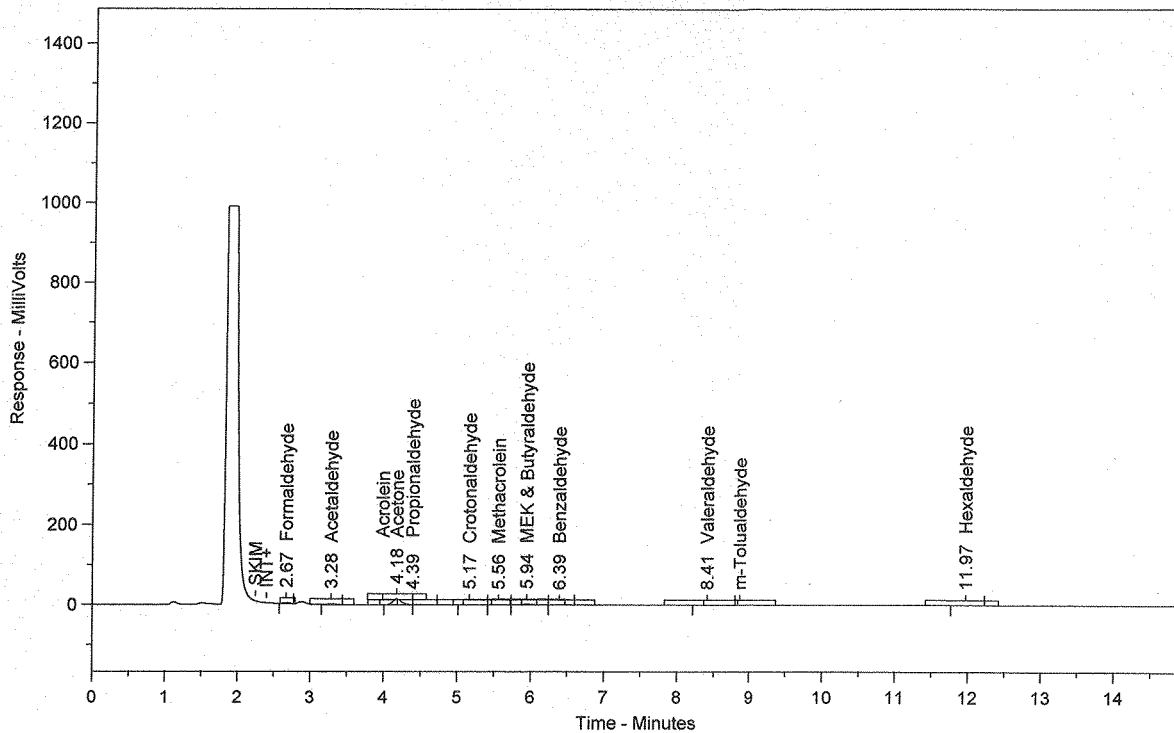
Total Area = 192834.5

Total Height = 22619.08

Total Amount = 0.4709261

Chrom Perfect Chromatogram Report

131099-65569



Sample Name = 131099-65569

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0026.RAW

Date Taken (end) = 8/22/2013 1:01:31 PM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 26

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.67	Formaldehyde	0.0199	4.219	12801	6.553	BB	0.11
2	3.28	Acetaldehyde	0.0318	6.750	16819	8.609	BB	0.12
3	4.18	Acetone	0.3025	64.140	125949	64.469	BV	0.13
4	4.39	Propionaldehyde	0.0163	3.452	6730	3.445	VV	0.11
5	5.17	Crotonaldehyde	0.0150	3.173	5689	2.912	VV	0.18
6	5.56	Methacrolein	0.0106	2.253	4271	2.186	VV	0.15
7	5.94	MEK & Butyraldehyde	0.0537	11.392	17391	8.902	VV	0.17
8	6.39	Benzaldehyde	0.0038	0.814	1016	0.520	VB	0.25
9	8.41	Valeraldehyde	0.0101	2.145	2823	1.445	BB	0.39
10	11.97	Hexaldehyde	0.0078	1.663	1874	0.959	BB	0.24

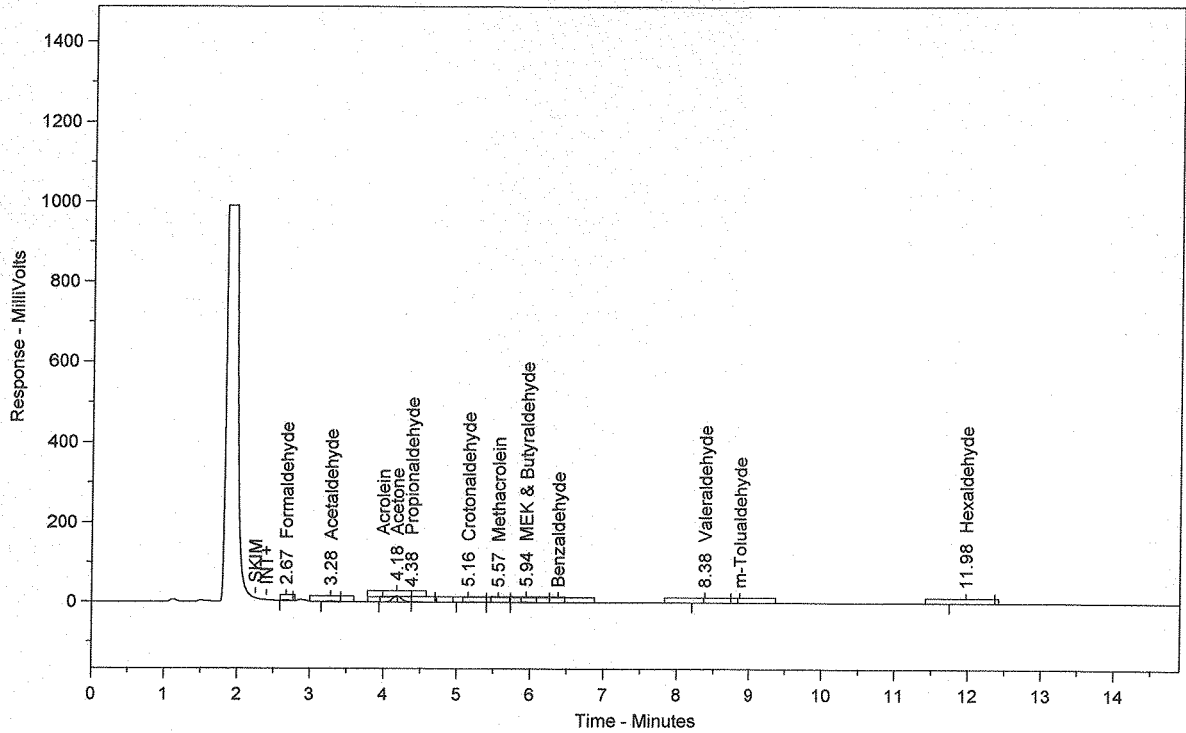
Total Area = 195362.9

Total Height = 22716.19

Total Amount = 0.4715627

Chrom Perfect Chromatogram Report

131099-65570



Sample Name = 131099-65570

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0027.RAW

Date Taken (end) = 8/22/2013 1:23:21 PM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 27

Injection Volume = 10

Dilution Factor = 1

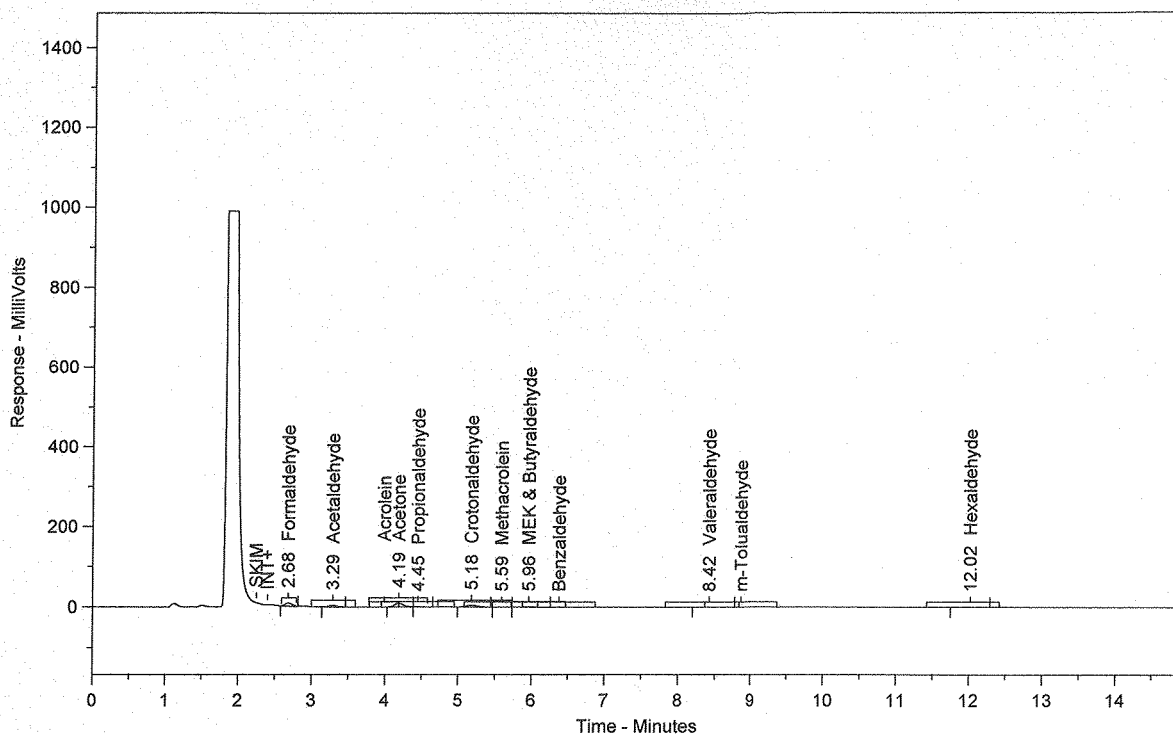
Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.67	Formaldehyde	0.0175	3.773	11256	5.890	BB	0.11
2	3.28	Acetaldehyde	0.0258	5.564	13631	7.133	BB	0.12
3	4.18	Acetone	0.3130	67.509	130323	68.193	BV	0.13
4	4.38	Propionaldehyde	0.0151	3.254	6237	3.264	VV	0.12
5	5.16	Crotonaldehyde	0.0152	3.280	5781	3.025	VB	0.18
6	5.57	Methacrolein	0.0071	1.527	2847	1.490	BV	0.14
7	5.94	MEK & Butyraldehyde	0.0463	9.977	14973	7.835	VB	0.17
8	8.38	Valeraldehyde	0.0098	2.114	2736	1.432	BB	0.37
9	11.98	Hexaldehyde	0.0139	3.002	3325	1.740	BB	0.29

Total Area = 191108.9

Total Height = 22523

Total Amount = 0.4635923

131099-65571



Sample Name = 131099-65571

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0028.RAW

Date Taken (end) = 8/22/2013 1:40:00 PM

Method File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0028.BND

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0028.BND

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 28

Injection Volume = 10

Dilution Factor = 1

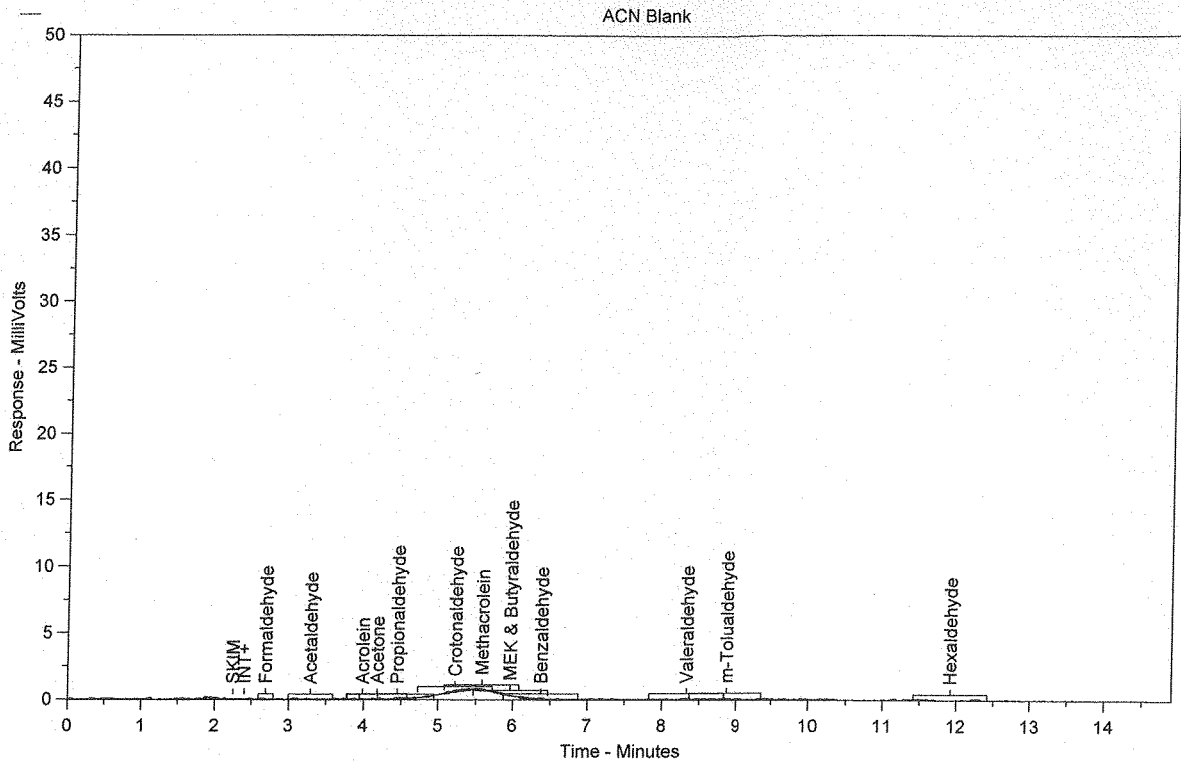
Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.68	Formaldehyde	0.0705	14.168	45355	20.953	BB	0.11
2	3.29	Acetaldehyde	0.0480	9.658	25389	11.729	BB	0.12
3	4.19	Acetone	0.1822	36.625	75873	35.052	BV	0.13
4	4.45	Propionaldehyde	0.0126	2.523	5190	2.398	VB	0.14
5	5.18	Crotonaldehyde	0.1153	23.173	43825	20.246	BB	0.17
6	5.59	Methacrolein	0.0064	1.290	2581	1.193	BV	0.14
7	5.96	MEK & Butyraldehyde	0.0313	6.285	10123	4.676	VB	0.19
8	8.42	Valeraldehyde	0.0165	3.313	4601	2.126	BB	0.33
9	12.02	Hexaldehyde	0.0147	2.965	3524	1.628	BB	0.27

Total Area = 216461.4

Total Height = 25989.21

Total Amount = 0.4974892

Chrom Perfect Chromatogram Report



Sample Name = ACN Blank

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0034.RAW

Date Taken (end) = 8/22/2013 3:19:55 PM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 34

Injection Volume = 10

Dilution Factor = 1

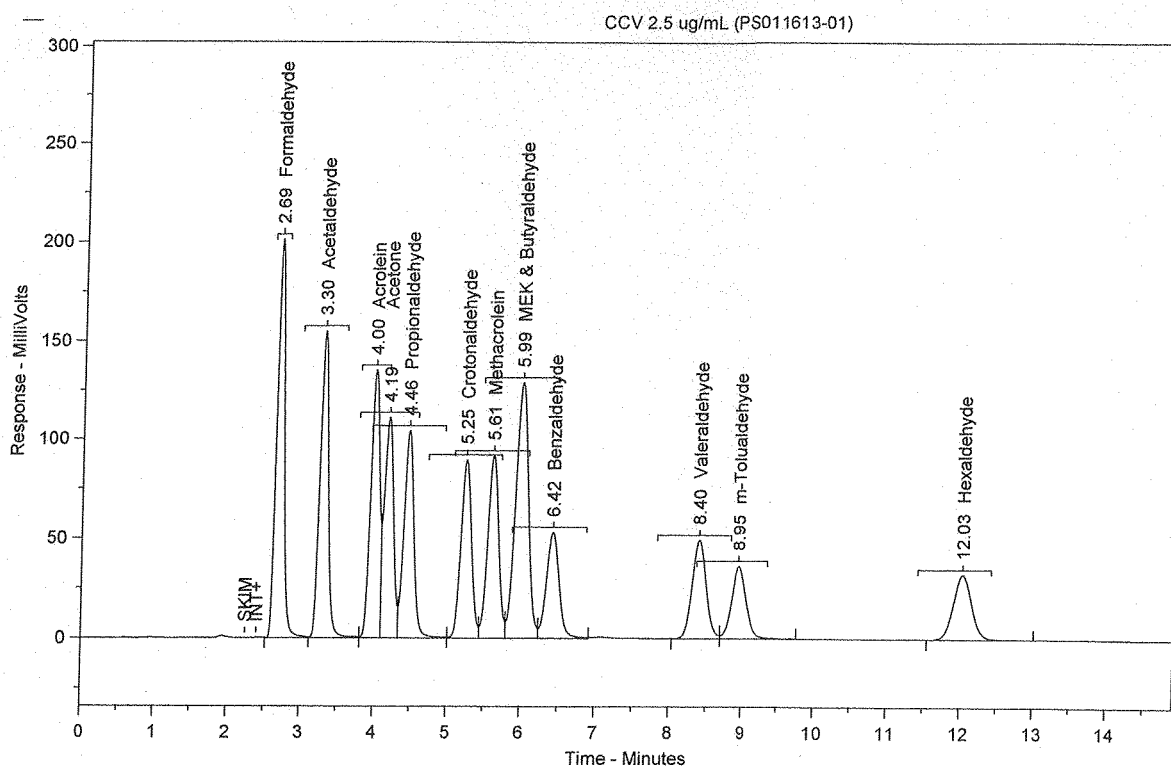
Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
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Total Area = 0

Total Height = 0

Total Amount = 0





Sample Name = CCV 2.5 ug/mL (PS011613-01)

Instrument = HPLC #1

Raw File Name = C:\Chromperfect 2\Data\HPLC #1\2013\082213TO-11\082213.0035.RAW

Date Taken (end) = 8/22/2013 3:36:34 PM

Method File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.MET

Method Description=TO-11A Carbonyls

Calibration File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\061113 TO-11A.CAL

Concentration Units = ug/ml

Run Time = 14.89889

Vial Number = 35

Injection Volume = 10

Dilution Factor = 1

Peak #	Ret. Time	Name	Amount	Amt %	Area	Area %	Type	Width
1	2.69	Formaldehyde	2.2992	7.568	1479446	12.883	SBB	0.11
2	3.30	Acetaldehyde	2.3403	7.703	1236663	10.768	TBV	0.12
3	4.00	Acrolein	2.3307	7.671	1116729	9.724	TWV	0.14
4	4.19	Acetone	2.3363	7.690	972880	8.472	TWV	0.14
5	4.46	Propionaldehyde	2.3536	7.747	973196	8.474	TWV	0.14
6	5.25	Crotonaldehyde	2.3330	7.679	886904	7.723	TWV	0.15
7	5.61	Methacrolein	2.3377	7.694	939985	8.185	TWV	0.15
8	5.99	MEK & Butyraldehyde	4.6927	15.446	1519158	13.228	TWV	0.18
9	6.42	Benzaldehyde	2.3628	7.777	625028	5.443	TVB	0.18
10	8.40	Valeraldehyde	2.3197	7.635	647611	5.639	BV	0.20
11	8.95	m-Tolualdehyde	2.3156	7.622	522568	4.550	VB	0.22
12	12.03	Hexaldehyde	2.3603	7.769	563946	4.911	BB	0.27

Total Area = 1.148411E+07

Total Height = 1187265

Total Amount = 30.38198

## Chrom Perfect Sequence File

File Name = C:\Chromperfect 2\Methods and Sequences\#1 HPLC #1\2013\082213 (TO-11).SEQ

File Date = 8/22/2013 12:42:57 PM

Analyst=

Entry #	Raw File Name	Method File Name	Sample Name	Vial #	DF
1	082213.0001.raw	061113 TO-11A.MET	ACN Blank	1	1
2	082213.0002.raw	061113 TO-11A.MET	CCV 2.5 ug/mL (PS061113-01)	2	1
3	082213.0003.raw	061113 TO-11A.MET	SS 2.50 ppm (PS011613-01)	3	1
4	082213.0004.raw	061113 TO-11A.MET	TO-11 Method Blank	4	1
5	082213.0005.raw	061113 TO-11A.MET	LCS Blank	5	1
6	082213.0006.raw	061113 TO-11A.MET	LCS .379ug/mL (PS011013-01)	6	1
7	082213.0007.raw	061113 TO-11A.MET	MS 131114-65647 1.25 ppm [(PS061113-01x2]	7	1
8	082213.0008.raw	061113 TO-11A.MET	MSD 131114-65647 1.25 ppm [(PS061113-01x2]	8	1
9	082213.0009.raw	061113 TO-11A.MET	131114-65647	9	1
10	082213.0010.raw	061113 TO-11A.MET	131114-65647 dup	10	1
11	082213.0011.raw	061113 TO-11A.MET	131114-65639	11	1
12	082213.0012.raw	061113 TO-11A.MET	ACN Blank	12	1
13	082213.0013.raw	061113 TO-11A.MET	CCV 2.5 ug/mL (PS011613-01)	13	1
14	082213.0014.raw	061113 TO-11A.MET	131114-65640	14	1
15	082213.0015.raw	061113 TO-11A.MET	131114-65640 dup	15	1
16	082213.0016.raw	061113 TO-11A.MET	131114-65641	16	1
17	082213.0017.raw	061113 TO-11A.MET	131114-65642	17	1
18	082213.0018.raw	061113 TO-11A.MET	131114-65643	18	1
19	082213.0019.raw	061113 TO-11A.MET	131114-65644	19	1
20	082213.0020.raw	061113 TO-11A.MET	131114-65645	20	1
21	082213.0021.raw	061113 TO-11A.MET	131114-65646	21	1
22	082213.0022.raw	061113 TO-11A.MET	ACN Blank	22	1
23	082213.0023.raw	061113 TO-11A.MET	CCV 2.5 ug/mL (PS011613-01)	23	1
24	082213.0024.raw	061113 TO-11A.MET	131099-65568	24	1
25	082213.0025.raw	061113 TO-11A.MET	131099-65568 dup	25	1
26	082213.0026.raw	061113 TO-11A.MET	131099-65569	26	1
27	082213.0027.raw	061113 TO-11A.MET	131099-65570	27	1
28	082213.0028.raw	061113 TO-11A.MET	131099-65571	28	1
29	082213.0029.raw	061113 TO-11A.MET	131xxx-D1	29	1
30	082213.0030.raw	061113 TO-11A.MET	131xxx-D2	30	1
31	082213.0031.raw	061113 TO-11A.MET	131xxx-U1	31	1
32	082213.0032.raw	061113 TO-11A.MET	131xxx-U2	32	1
33	082213.0033.raw	061113 TO-11A.MET	ACN Blank	33	1
34	082213.0034.raw	061113 TO-11A.MET	ACN Blank	34	1
35	082213.0035.raw	061113 TO-11A.MET	CCV 2.5 ug/mL (PS011613-01)	35	1

