

2023 Q3 MOBILE MONITORING VAN COMMERCE CITY NORTH DENVER COMMUNITY AIR MONITORING NETWORK COMMERCE CITY, COLORADO

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EXECUTIVE SUMMARY

In response to feedback received by Suncor Energy (U.S.A.) Inc. (Suncor) through community engagement conducted in the fall of 2020, Suncor voluntarily committed to developing a continuous, near real-time air monitoring program to gain insight into air quality for neighborhoods in the vicinity of the Suncor refinery in Commerce City, Colorado. Montrose Environmental Group - Air Quality Services, LLC (Montrose) was contracted by Suncor to deploy, operate and maintain the network in the Commerce City and North Denver (CCND) neighborhoods. Air monitoring was accomplished through three separate technical approaches: (1) continuous, near real-time monitoring for the following analytes¹: carbon monoxide (CO), sulfur dioxide (SO₂), hydrogen sulfide (H₂S), nitric oxide or nitrogen oxide (NO), nitrogen dioxide (NO₂), particulate matter (PM_{2.5}) and total volatile organic compounds (VOCs); (2) periodic collection and laboratory analysis for the presence of specific VOCs from Summa canisters; and (3) periodic real-time air monitoring throughout neighborhoods using a mobile monitoring van to detect the presence of specific VOCs and hydrogen sulfide (H₂S). This report details approach number three, the periodic real-time air monitoring through six neighborhoods with the mobile monitoring van and a screening health risk analysis of the detected chemicals. Continuous real-time air monitoring and Summa canister sampling data are presented in separate reports.

The mobile monitoring van contains the equipment necessary to identify and quantitate individual chemicals present in ambient air at ultra-low concentrations. This equipment measures and reports concentrations of select chemicals at sub-parts per billion (ppb) levels and as quickly as one measurement per second. During the monitoring period, the mobile monitoring van followed a dense route through each of the six CCND residential neighborhoods that fall within a three-mile radius around the refinery. Accessible streets in the monitored neighborhoods were traversed at approximately 10 miles per hour (MPH) while collecting a data point for each chemical every 1 second. During the third quarter 2023 sampling period (July 31-August 4), the mobile monitoring van was in a total of six neighborhoods and collected more than 66,584 data points across five days of monitoring, resulting in approximately 45,422 1-hour rolling average concentrations. Meteorological conditions were also reported in real time.

Health scientists from CTEH, LLC (CTEH®) (a subsidiary company of Montrose) performed a screening-level human health risk assessment based on the data collected by Montrose. This evaluation was consistent with federal and state risk assessment guidelines and was conducted to determine whether the estimated 1-hour maximum measured concentrations of individual or cumulative (combined) VOCs could potentially pose acute (short-term) health hazards. The air monitoring data and health risk assessment indicate:

- Air monitoring data and health risk assessment indicate all measured individual and combined air concentrations were below their respective acute health reference levels in all neighborhoods.
- Results indicate the measured concentrations are likely to be without any appreciable risk of adverse acute health effects, even for sensitive sub-populations.

¹An “analyte” is a material that a measuring device is designed to detect and measure. It may be a chemical gas, an airborne particle, or other type of material.

1.0 INTRODUCTION

In response to feedback received by Suncor Energy (U.S.A.) Inc. (Suncor) through community engagement conducted in the fall of 2020, voluntarily committed to developing a continuous, near real-time air monitoring program to gain insight into air quality for neighborhoods in the vicinity of the Suncor refinery in Commerce City, Colorado. Montrose Environmental Group - Air Quality Services, LLC (Montrose) was contracted by Suncor to deploy, operate and maintain the network in the Commerce City and North Denver (CCND) neighborhoods. Air monitoring was accomplished through three separate technical approaches: (1) continuous, near real-time stationary monitoring for the following analytes: carbon monoxide (CO), sulfur dioxide (SO₂), hydrogen sulfide (H₂S), nitric oxide or nitrogen oxide (NO), nitrogen dioxide (NO₂), particulate matter (PM_{2.5}) and total volatile organic compounds (VOCs); (2) periodic collection and laboratory analysis for the presence of specific VOCs from Summa canisters; and (3) periodic real-time air monitoring throughout neighborhoods using a mobile monitoring van to detect the presence of specific chemicals. An “analyte” is a material that a measuring device is designed to detect and measure. It may be a chemical gas, an airborne particle, or other type of material. This report details approach number three. The continuous real-time community air monitoring and Summa canister sampling data are presented in separate reports. Air monitoring, sampling and analysis from approaches (1) and (2) were conducted in accordance with the Quality Assurance Project Plan (QAPP) that can be found online at ccnd-air.com/documents.

2.0 MOBILE SAMPLING PROGRAM

2.1 Mobile Van Air Sampling Description

The mobile monitoring van is a Mercedes 2500 Sprinter Van outfitted with equipment necessary to identify and quantitate individual chemicals present in ambient air to ultra-low concentrations. The mobile monitoring van is equipped with an Ionicon Model 6000-X2 proton transfer reaction time-of-flight mass spectrometer (PTR-TOF-MS). This instrument provides concentrations of select chemicals at sub-parts per billion (ppb) levels and as quickly as one measurement per second. The mobile monitoring van is outfitted with an external sampling system, which transports ambient air from outside of the van into the PTR-TOF-MS sample inlet for immediate real-time analysis. The entire sampling system is comprised of Teflon or Teflon-coated materials, which ensures the lowest amount of sample loss due to surface absorption of chemical molecules. The mobile monitoring van incorporates a high-precision global positioning system (GPS), a sonic anemometer to measure wind direction and wind velocity and other incorporated meteorological (MET) sensors.

During the mobile monitoring program, groups of structurally similar chemicals (called isomers) that include the list of 65 chemicals in Table 2-1 were measured to determine the instantaneous ambient concentrations. Appendix A provides more detail on the need for isomer grouping. This list of chemicals was compiled based on the typical chemicals that are monitored in urban and industrial areas and the mobile monitoring van analysis capabilities.

The mobile monitoring van followed a driving route through each of the six CCND residential neighborhoods that fall within a three-mile radius around the refinery operations. Accessible streets in the neighborhoods were traversed at approximately 10 MPH while collecting a data point approximately every 1 second. The details of the monitored neighborhoods are listed in Table 2-2 and are shown in Figure 2-1.

TABLE 2-1
MOBILE MONITORING VAN PROGRAM CHEMICALS²

o-Diethylbenzene	2-Methylhexane	Neopentane	Methyl-cyclopentane	o-Ethyltoluene (2-ethyltoluene)
1,3-Butadiene	2-Methylpentane	Ethylbenzene	m-Ethyltoluene	p-Diethylbenzene (1,4-diethylbenzene)
1-Butene	3-Methylheptane	Ethylcyclohexane	m/o/p-Xylenes	p-Ethyltoluene (4-ethyltoluene)
1-Hexene	3-Methylhexane	Ethylene	n-Butane	1,2,4-trimethylbenzene
1-Pentene	3-Methylpentane	Hydrogen Cyanide	n-Decane	Propylene (Propene)
Styrene	Acetylene	Hydrogen Sulfide	n-Dodecane	2,2,4-Trimethylpentane
2,2-Dimethylbutane	Benzene	i-Butane	n-Heptane	Tetrachloroethylene
Toluene	Carbon disulfide	i-Pentane	n-Hexane	2,3,4-Trimethylpentane
2,3-Dimethylbutane	trans-2-Butene	Isopentane	n-Nonane	trans-1,2-Dimethylcyclohexane
2,3-Dimethylpentane	cis-2-Butene	Isoprene	n-Octane	trans-1,3-Dimethylcyclohexane
2,4-Dimethylpentane	cis-2-Pentene	m-Diethylbenzene	n-Pentane	cis-1,3-dimethylcyclohexane
2-Methyl-2-butene	Cumene	Methanol	n-Propylbenzene	trans-2-Pentene
2-Methylheptane	Cyclohexane	Methyl-cyclohexane	n-Undecane	Cyclopentane

² See Appendix A for isomer analysis details

TABLE 2-2
NEIGHBORHOOD MONITORING PROGRAM DETAILS

Neighborhood	Area (square miles)	Sampling Date	Start Time	End Time	Total Data Points Collected	Total Hourly Rolling Averages Calculated*
Adams City	0.41	8/4/23	9:13	12:03	10,146	6,619
Dupont	1.4	8/2/23	16:57	20:06	11,273	7,746
Elyria-Swansea	1.2	8/1/23	18:43	20:51	7,613	4,086
Globeville	0.44	8/1/23	16:26	18:34	7,626	4,099
Pioneer Park	1.7	7/31/23	11:30	15:37	14,772	11,245
Western Hills	1.6	8/3/23	11:48	16:02	15,154	11,627

*Data completeness threshold set at 98%

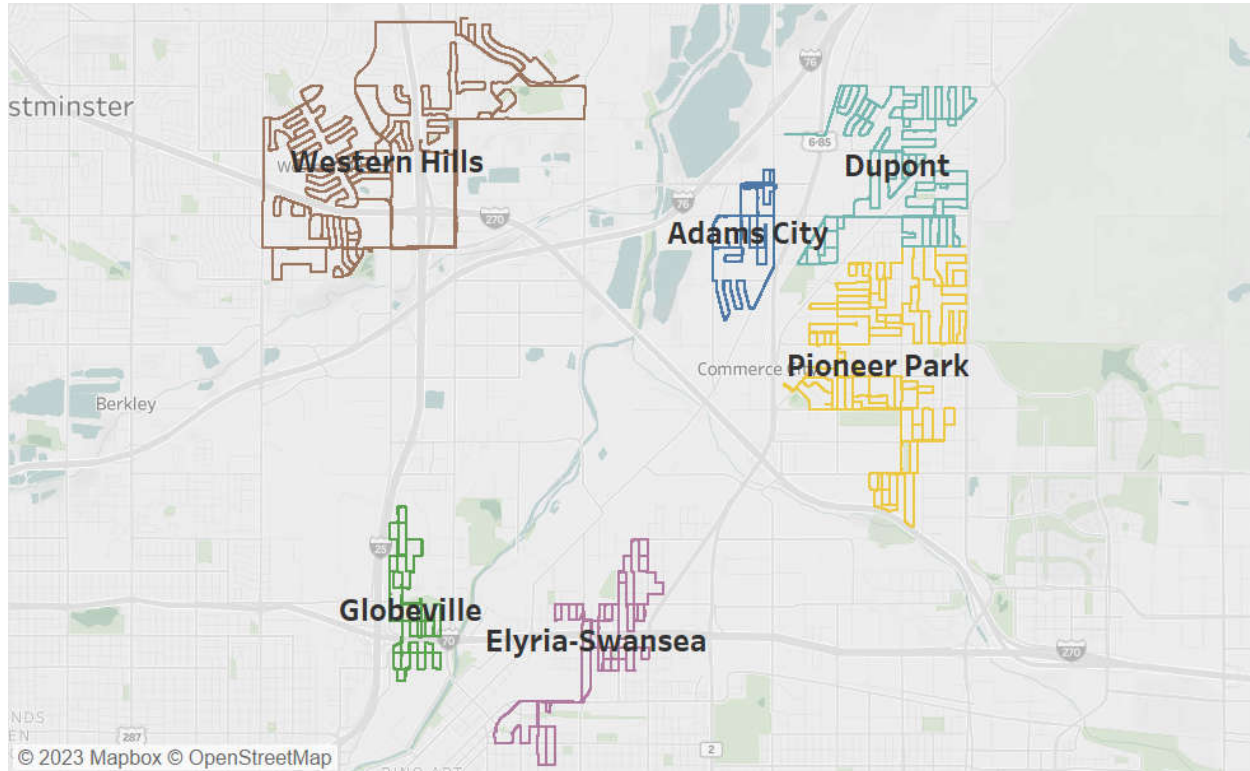
2.2 Mobile Monitoring Van Air Sampling Methods

The PTR-TOF-MS calibration was checked, and the instrument was zeroed each day prior to collection of any ambient air data. The instrument was calibrated using United States Environmental Protection Agency (USEPA) protocol certified calibration gases. The multi-chemical cylinder standards were used to generate multiple point calibration curves for each commercially available chemical present in the standard. Note: Not all chemicals listed in Table 2-1 are available as certified calibration gases. The chemical dilutions were made using an Environics Model 4040 gas dilution system. The gas dilution system was validated using the appropriate USEPA methodology (40 Code of Federal Regulation Part 51 Appendix M, Method 205). Zero-count measurements were obtained to ensure proper baseline measurements were incorporated into the calculation of each chemical's concentration. Zero-count measurements were performed through the entire sampling system using ultra-high purity air. Post-testing calibration checks were performed on the instrument to ensure there was no significant drift during the course of the sampling event. Drift can cause an increase or decrease in the measured chemical concentrations, which can lead to both positive and negative biasing of the obtained results.

The mobile monitoring van collected continuous measurements throughout each neighborhood following the routes shown in Figure 2-1. The measurements were collected from the ambient environment at a height of 15 feet above grade at approximately 8 liters per minute using a Teflon-coated sampling boom and pump. The PTR-TOF-MS sampled a slip stream of this flow at approximately 100 ml/min. The sample was introduced into the reaction tube of the PTR-TOF-MS

and results were collected in 1-second intervals. See the attached Appendix D for specific PTR-TOF-MS instrument operation conditions.

**FIGURE 2-1
MOBILE MONITORING VAN PROGRAM ROUTE THROUGH SIX NEIGHBORHOOD AREAS**



2.3 Screening Health Risk Assessment Methods

CTEH® conducted a screening-level public health risk assessment, consistent with federal risk assessment guidelines, to determine whether exposure to the detected concentrations of individual or cumulative (combined) chemicals in the air could potentially pose acute (short-term) health impacts. A tiered approach to the risk assessment was used. This approach involves one or more iterative steps (or tiers) being performed in which health risks are calculated and evaluated multiple times. In most cases, risk assessors cannot know exactly the level of chemical exposure experienced by individuals or communities. Therefore, the first tier involves use of exposure assumptions that are health-conservative. This means that data reflecting maximum exposure potential are plugged into the risk calculations. These are worst-case scenarios that typically represent exposure conditions higher than would be reasonably expected. Such calculations are very simple and assume a person is constantly exposed to the highest one hour rolling average concentration for each detected chemical. If the resulting risk values indicate the lack of likely acute adverse health effects under these worst-case conditions, then the risk assessment is complete. However, if the risk values suggest a potential for acute adverse health effects, then a second tier of risk calculations are performed, but this time using more detailed assumptions about exposure that are still simple representations of the real world but are more realistic than the first-tier worst-case assumptions. Each successive tier represents a more

complete characterization of exposure variability and/or uncertainty that requires a corresponding increase in calculation complexity and scientific level of effort.

The first tier of this risk assessment process is called a screening-level risk assessment. The conservative assumptions used for this level of risk calculation typically represent exposure conditions higher than would be reasonably expected. As such, an exceedance of an acceptable risk level (defined below) does not necessarily indicate that adverse health effects are likely. The Agency for Toxic Substances and Disease Registry (ATSDR) states, “*when health assessors find exposures higher than the MRLs (ATSDR’s specific health-based reference levels), it means that they may want to look more closely at a site*”³. In other words, screening-level findings of an estimated exposure to a specific or cumulative set of chemical(s) being higher than its reference level (RL) does NOT indicate an actual likelihood of adverse effects but does indicate a need to move to a second tier of analysis and refine the risk assessment process with more realistic detail to determine if an actual risk exists that needs to be mitigated.

The screening-level risk assessment reported here includes calculated acute risks from exposure to individually measured chemicals as well as exposure to all measured chemicals at once (cumulative). For individual chemicals, an acute health risk value was calculated as the exposure concentration (EC) divided by the chemical-specific federal or state established acute RL (Equation 1). The result is referred to as the hazard quotient (HQ). Estimates of EC were derived from 1-hour rolling average concentrations of each chemical for the entire measurement time in an individual CCND neighborhood. The RLs used to calculate the HQs are previously established exposure levels below which no adverse effect in humans is expected. If available, RLs adopted by the Colorado Department of Public Health and Environment (CDPHE) were selected for use within this assessment and include ATSDR acute minimum risk levels (MRL), California EPA Office of Environmental Health Hazard Assessment (OEHHA) acute risk levels and Texas Commission on Environmental Quality (TCEQ) acute exposure guideline levels. If the chemical was not listed by CDPHE, a federal and state recommended hierarchy for selection of RLs was used⁴. For chemical isomer groups which were unable to be differentiated the lowest, most health-protective RL of the isomer group was selected for use in this assessment.

Acute HQs were calculated as follows:

Eq. 1 – Hazard Quotient (HQ) Equation

$$HQ = EC/RL$$

Where:

HQ = Hazard Quotient

EC = Maximum 1-hour rolling average air concentration

RL = Acute Health-based Reference Level (ATSDR, Cal EPA OEHHA and TCEQ)

Health risks from potential cumulative exposures to all detected chemicals were calculated by adding together each individual chemical's HQ calculated for a given neighborhood. The sum of all the individual HQs is called a Hazard Index (HI). Adding together all the HQs is also a very

³[https://www.atsdr.cdc.gov/minimalrisklevels/#:~:text=The%20ATSDR%2C%20in%20response%20to,minimal%20risk%20levels%20\(MRLs\)](https://www.atsdr.cdc.gov/minimalrisklevels/#:~:text=The%20ATSDR%2C%20in%20response%20to,minimal%20risk%20levels%20(MRLs))

⁴ <https://drive.google.com/file/d/1P2KEvu0MFiyzQAOQtjQUclqR-WGh1bEX/view>

health-conservative approach because it assumes that all the measured chemicals exert an adverse effect on the body in a similar manner, which is rarely the case.

An HQ or HI of less than or equal to one is an indication that the estimated exposure is likely to be without an appreciable risk of adverse acute health effects, even for sensitive sub-populations. The potential for adverse health effects increases as HQ or HI increase above one, but it is not known by how much. HQ or HI values of greater than one would prompt a second-tier risk assessment beyond the screening-level assessment.

According to the USEPA and ATSDR, the federal agencies that establish these RLs, these values “*are set below levels that, based on current information, might cause adverse health effects in the people most sensitive.*”⁵ This is because RLs are based on observed toxicity in human or animal studies with an added safety factor to account for uncertainties and variabilities in the toxicity data. For example, ATSDR identified the lowest observed adverse effect level (LOAEL) for acute exposure to benzene as 10,200 parts per billion (ppb), based on a study of mice exposed six hours per day for six days. ATSDR then applied a combined safety factor of 300 to derive the final RL to account for several uncertainties, including differences between mice and humans and for sensitive individuals. Therefore, it is scientifically incorrect to assume that all real-world exposures to a chemical at levels higher than a RL will likely result in an adverse effect.

Using the maximum 1-hour rolling average for the EC conservatively assumes that a hypothetical maximally exposed individual occupies the monitored neighborhood and breathes the maximum 1-hour detected concentration continuously for an hour up to multiple days (an acute exposure). A 1-hour average concentration is more appropriate than a 1-second or 1-minute concentration for use in an acute health risk assessment. This is because 1-second exposures to the chemicals measured in this study do not cause adverse effects unless the levels are extremely high (i.e., tens of thousands to millions of ppb). Guidance values for use in emergency situations with extremely elevated levels of these chemicals are available and are discussed below. Across all neighborhoods, 45,422 1-hour rolling averages of chemical concentrations were calculated to derive the estimated ECs (Table 2-2). The range between the average and maximum rolling 1-hour average values provides a robust estimate of plausible outdoor exposures of persons occupying the monitored neighborhood while the mobile monitoring van was present (Figures 3-1 to 3-6).

The USEPA also has established values for use in emergency situations, termed Acute Exposure Guideline Levels (AEGLs). Unlike RLs that can be thousands of times below exposure levels where adverse effects are observed, AEGL values are levels at which different acute adverse health effects may be anticipated to occur. According to USEPA, “*AEGL-1 represent exposure levels that could produce mild and progressively increasing but transient and non-disabling odor, taste and sensory irritation or certain asymptomatic, non-sensory effects. With increasing airborne concentration above each AEGL, there is a progressive increase in the likelihood of occurrence and the severity of effects described for each corresponding AEGL [i.e., AEGL-2 or AEGL-3].*”⁶ The AEGL-1 60-minute value, if available for the applicable chemical, was also used for comparison purposes because it is more precautionary (than AEGL-2 or AEGL-3) as the AEGL-1 level reflects protecting against acute health effects that are reversible upon cessation of exposure.

⁵

<https://www.atsdr.cdc.gov/mrls/index.html#:~:text=ATSDR%20uses%20the%20no%20observed,to%20such%20substance%2Dinduced%20effects>.

⁶ <https://www.epa.gov/aegl/about-acute-exposure-guideline-levels-aegls>

3.0 SUMMARY AND DISCUSSION OF RESULTS

3.1 Summary of Mobile Monitoring Van Results

A summary of mobile monitoring van results by neighborhood can be found in Table 2-2. Over five days, six neighborhoods were monitored for 65 chemicals, collecting more than 66,584 total data points. Individual neighborhood results are detailed in Figures 3-1 through 3-6. Each figure shows a map of the monitoring locations within each neighborhood, the chemicals that resulted in the five highest calculated acute HQs and time profiles of the measured levels of these chemicals. The time profiles show all the 1-second data (orange) and calculated 1-hour rolling averages (green) of the monitoring data. Each green 1-hour average data point shown in these profiles reflects all the 1-second measurements collected over the previous hour. Thus, 1-hour rolling average values are shown on the time profiles only after one hour of data collection (Figure 3-1 through 3-6).

Wind roses for each sampling day are provided in Appendix B. The data used to derive the wind roses were collected from the CCND community sensor location most local to the neighborhood being monitored on each day because the stationary source of MET data is more reliable than the MET station on the mobile monitoring van when the lab is moving.

3.2 Screening Health Risk Assessment Results

Acute health risks were calculated for each neighborhood. According to USEPA guidelines, an acute HQ or HI less than or equal to one (1) indicates that exposures are likely to be without any acute adverse health effects, even for sensitive sub-populations.

Maximum 1-hour rolling average concentrations for 65 chemicals measured in each neighborhood were compared to acute RLs to derive HQs. Figures 3-1 through 3-6 show concentrations of chemicals over the sampling time and summaries of results for chemicals resulting in the five highest HQs by neighborhood (if available). The estimated HI values (if available) shown in Figures 3-1 through 3-6 were calculated by summing the HQs of all detected chemicals measured in a given neighborhood. If any measured chemical resulted in a HQ greater than 1, then a separate figure would be shown for that chemical alone. Complete results for HQs for all chemicals detected in each neighborhood are available in Appendix C.

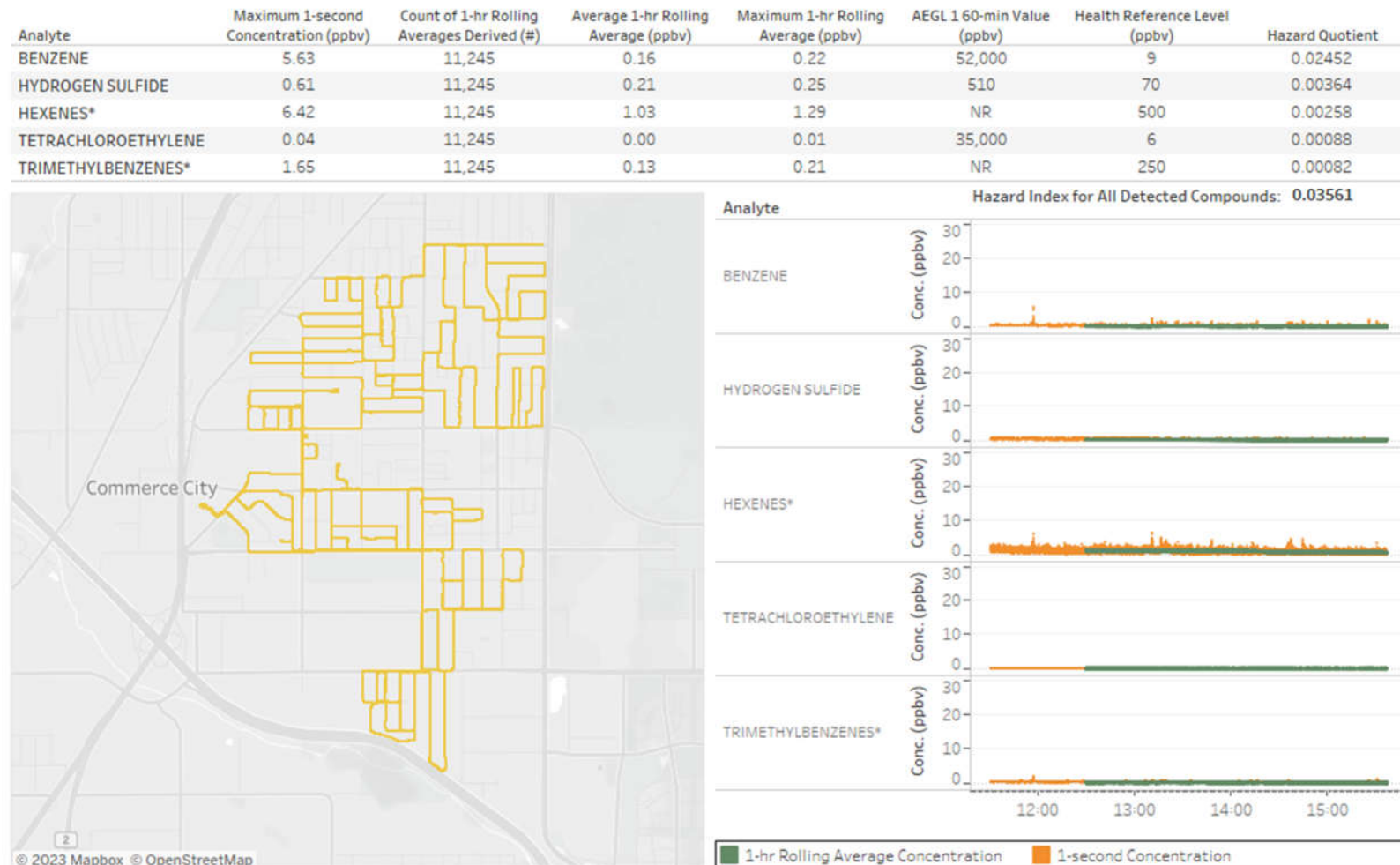
In conclusion, the data collected during this study phase did not indicate a potential for acute adverse health effects from exposure to the measured chemicals, both individually and combined.

- All HQs were less than one for all detected chemicals, indicating that the maximum 1-hour rolling average concentration for each chemical was below its respective acute RL in all six neighborhoods (Figure 3-1 through 3-6).
- In this quarter, benzene, tetrachloroethylene, hydrogen sulfide, hexene group, hydrogen cyanide, xylenes, and trimethylbenzene group were the chemicals or isomer groupings resulting in the highest HQ in each neighborhood, accounting for over 80% of the total calculated HI values. However, all HI values calculated in all six neighborhoods were below one (Figures 3-1 through 3-6).

CCND Mobile Monitoring Van
2023 Q3

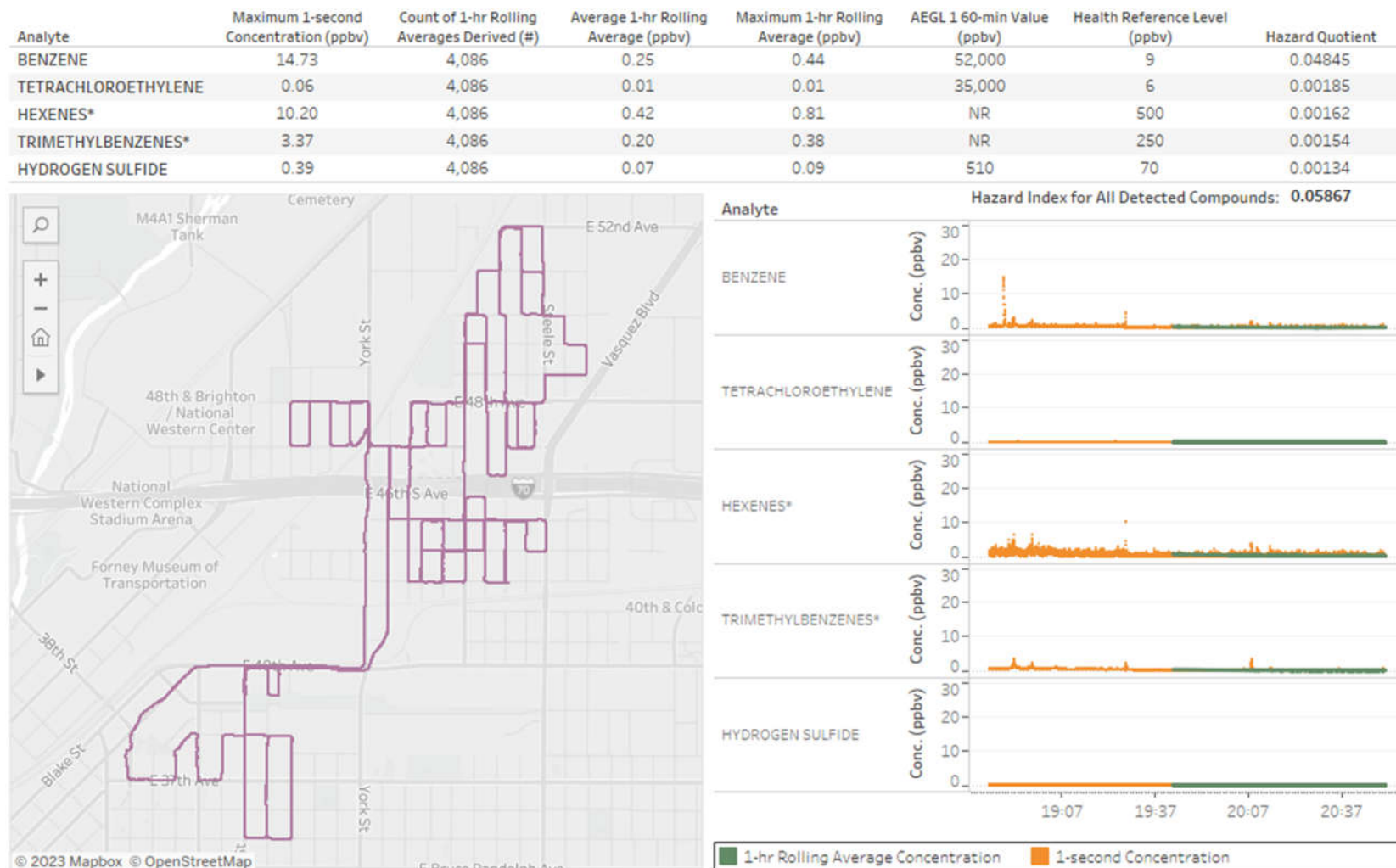
- These results indicate that the measured concentrations of chemicals, both individually and cumulative (combined), are likely to be without an appreciable risk of acute adverse health effects, even for sensitive sub-populations.

FIGURE 3-1
PIONEER PARK NEIGHBORHOOD: JULY 31, 2023



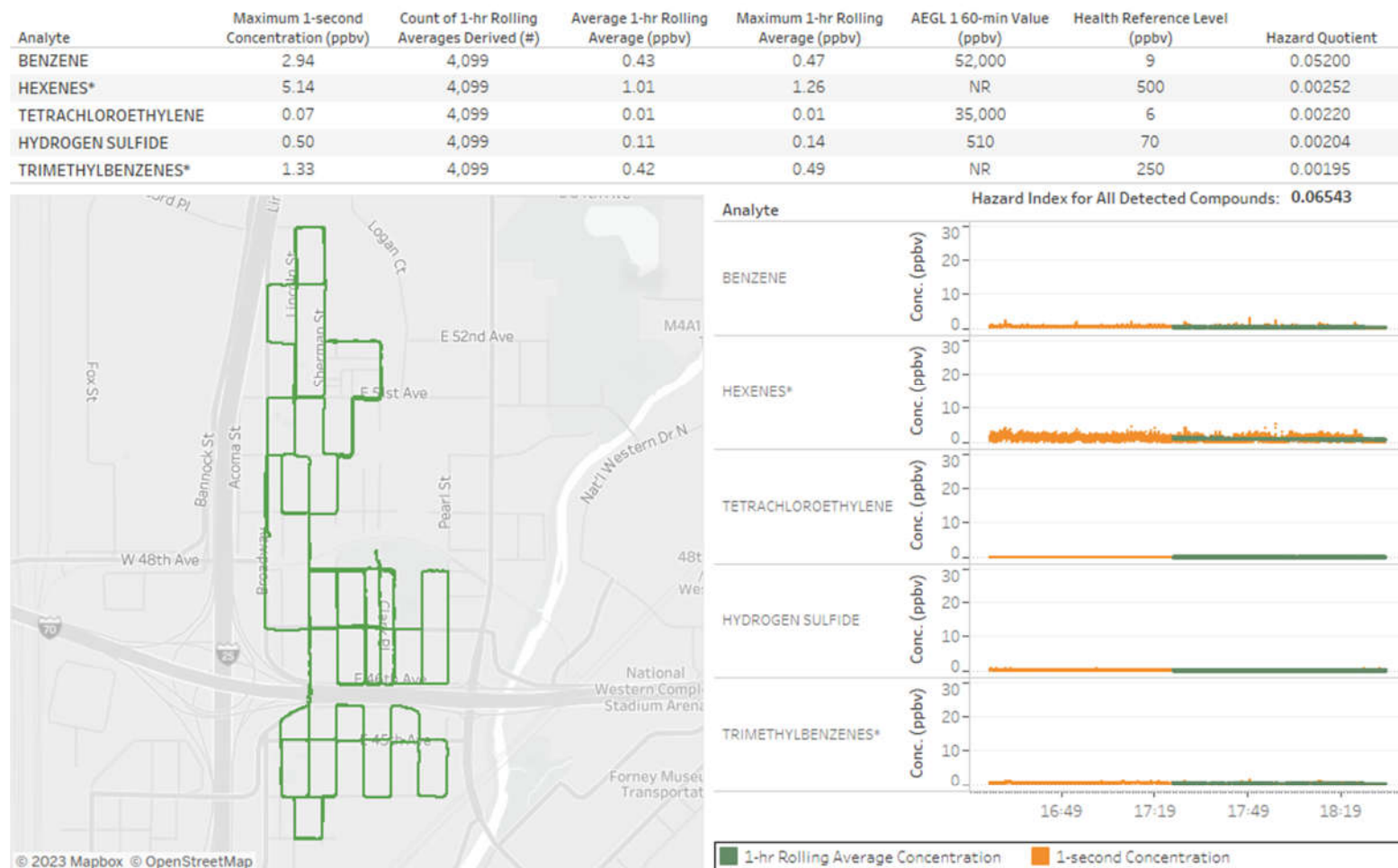
The top 5 hazard quotients are reported in this dashboard. The hazard index represents cumulative risks including all unlisted analytes. The hazard quotient was calculated by comparing the acute health reference level to the maximum 1-hour rolling average. The comparative AEGL value is shown for comparison purposes. NR = According to EPA, the AEGL value is "not recommended due to insufficient data". *For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the group was selected for use in this assessment (Appendix A).

FIGURE 3-2
ELYRIA-SWANSEA NEIGHBORHOOD: AUGUST 1, 2023



The top 5 hazard quotients are reported in this dashboard. The hazard index represents cumulative risks including all unlisted analytes. The hazard quotient was calculated by comparing the acute health reference level to the maximum 1-hour rolling average. The comparative AEGL value is shown for comparison purposes. NR = According to EPA, the AEGL value is "not recommended due to insufficient data". *For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the group was selected for use in this assessment (Appendix A).

FIGURE 3-3
GLOBEVILLE NEIGHBORHOOD: AUGUST 1, 2023



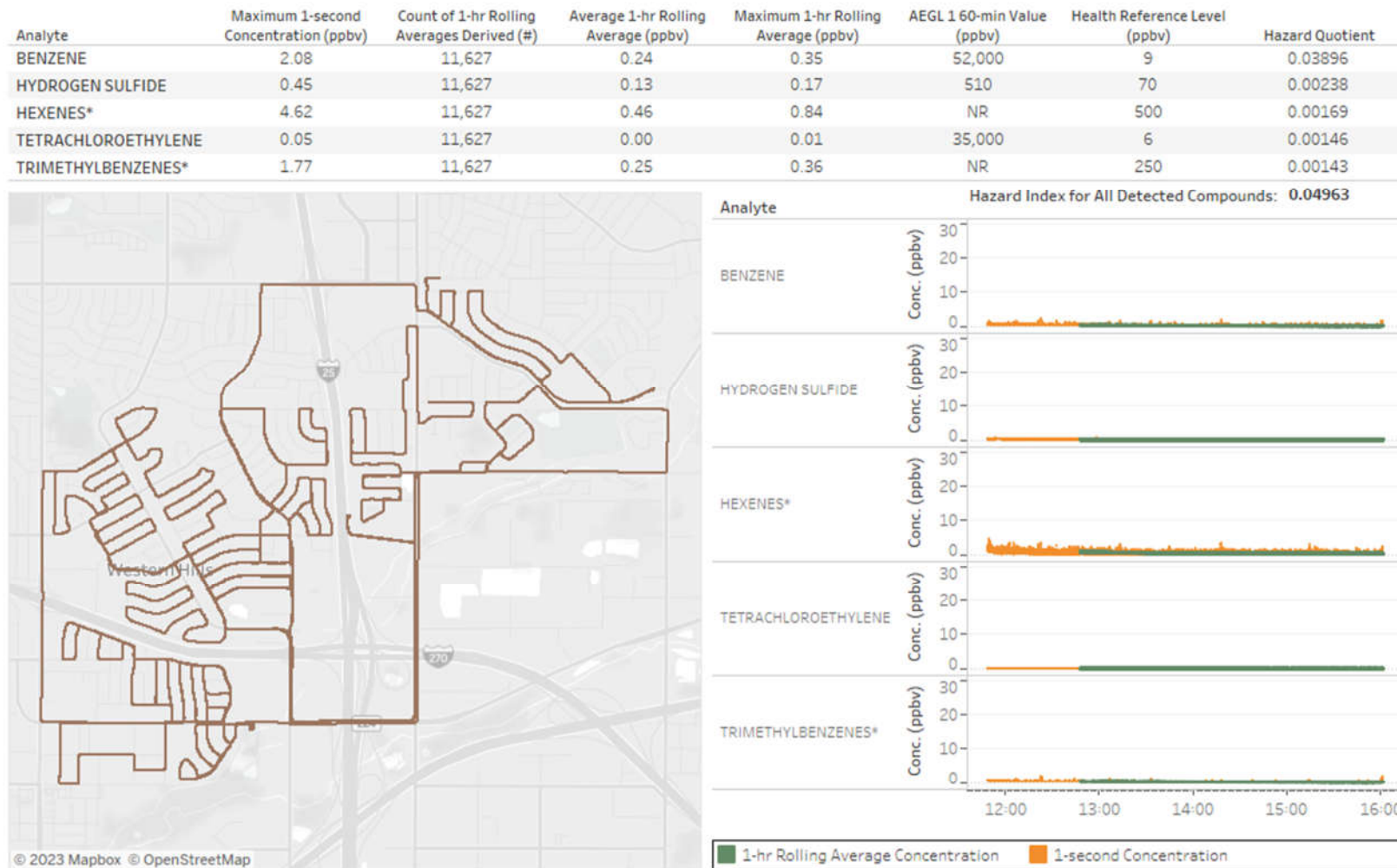
The top 5 hazard quotients are reported in this dashboard. The hazard index represents cumulative risks including all unlisted analytes. The hazard quotient was calculated by comparing the acute health reference level to the maximum 1-hour rolling average. The comparative AEGL value is shown for comparison purposes. NR = According to EPA, the AEGL value is "not recommended due to insufficient data". For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the group was selected for use in this assessment (Appendix A).

FIGURE 3-4
DUPONT NEIGHBORHOOD: AUGUST 2, 2023



The top 5 hazard quotients are reported in this dashboard. The hazard index represents cumulative risks including all unlisted analytes. The hazard quotient was calculated by comparing the acute health reference level to the maximum 1-hour rolling average. The comparative AEGL value is shown for comparison purposes. NR = According to EPA, the AEGL value is "not recommended due to insufficient data". *For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the group was selected for use in this assessment (Appendix A).

FIGURE 3-5
WESTERN HILLS NEIGHBORHOOD: AUGUST 3, 2023



The top 5 hazard quotients are reported in this dashboard. The hazard index represents cumulative risks including all unlisted analytes. The hazard quotient was calculated by comparing the acute health reference level to the maximum 1-hour rolling average. The comparative AEGL value is shown for comparison purposes. NR = According to EPA, the AEGL value is "not recommended due to insufficient data". *For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the group was selected for use in this assessment (Appendix A).

FIGURE 3-6
ADAMS CITY NEIGHBORHOOD: AUGUST 4, 2023



The top 5 hazard quotients are reported in this dashboard. The hazard index represents cumulative risks including all unlisted analytes. The hazard quotient was calculated by comparing the acute health reference level to the maximum 1-hour rolling average. The comparative AEGL value is shown for comparison purposes. NR = According to EPA, the AEGL value is "not recommended due to insufficient data". *For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the group was selected for use in this assessment (Appendix A).

3.3 Uncertainty Evaluation

Scientific uncertainty is inherent in each step of the risk assessment process because all risk assessments incorporate a variety of assumptions and professional judgments. Therefore, the acute hazard estimates presented in this assessment are estimates of risk due to a number of assumptions about exposure and toxicity. This screening-level risk assessment relied on a combination of health-protective exposure scenarios and input values (i.e., high-end exposures and conservative selection of lowest reference value per isomer). Because of these assumptions, the estimates of acute hazards are themselves uncertain but likely to be over-estimates of actual risk.

This risk assessment did not address past or present health outcomes associated with current or past exposures. As such, this risk assessment cannot be used to make realistic predictions of biological effects and/or used to determine whether someone is ill (cancer or other adverse health effects) due to past or current exposures. This risk assessment was limited to inhalation exposures from outdoor exposures to all potential sources.

3.4 Program Changes

No program changes occurred during this reporting period.

Respectfully Submitted:



Steven Yuchs, PhD.
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Michael Lumpkin, PhD, DABT
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CTEH®, LLC

APPENDIX A

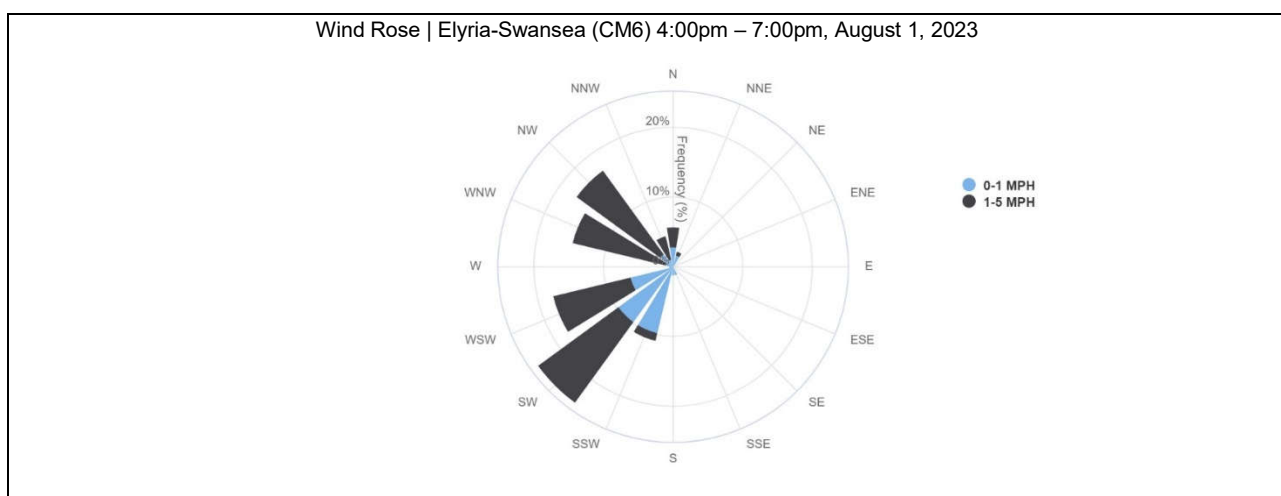
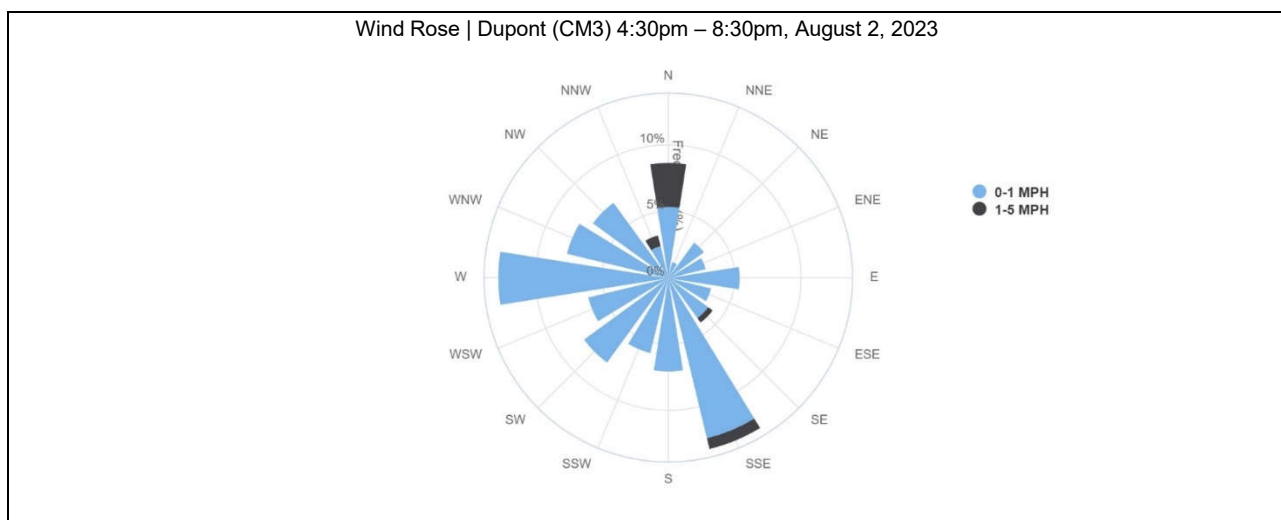
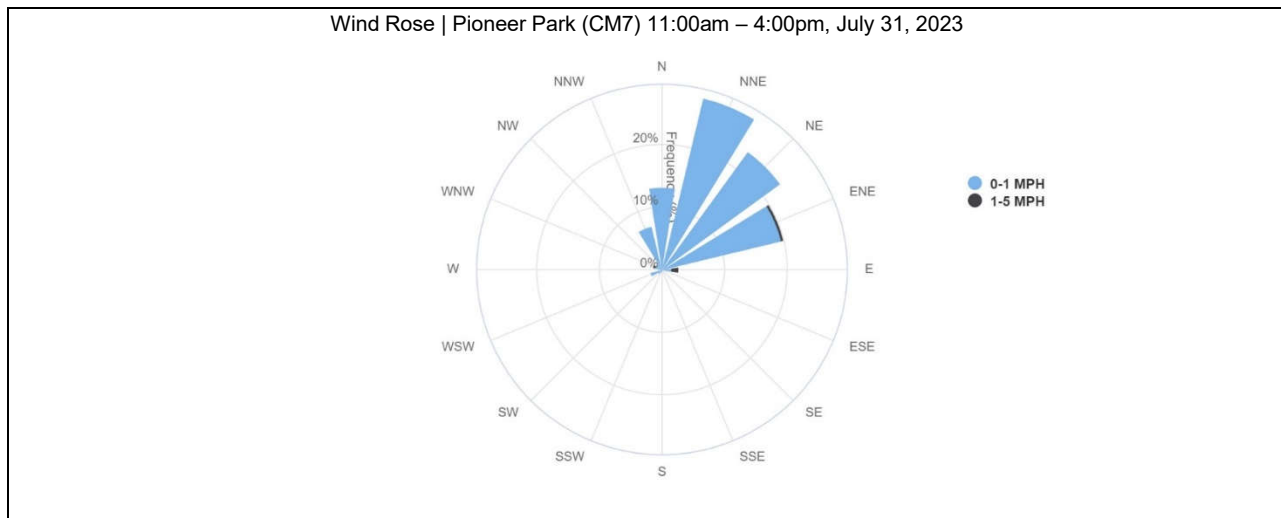
ISOMER CHEMICAL SAMPLING DETAILS

In a real-time PTR-TOF analysis, it is not possible to speciate isomers, or chemical compounds that have the same molecular weight. For example, n-hexane, 2-methyl pentane and 2,2-dimethyl butane all have a molecular mass of 86.178 g/mol. In order to provide the most conservative determination of concentration during this mapping program, each isomer's concentration is reported as the sum of all isomers with the same molecular weight. For the sake of simplicity, the calculations in the report refer to generic names for a group of specific isomers. The following table defines a simplified list of the many isomers that may comprise the generic groups reported.

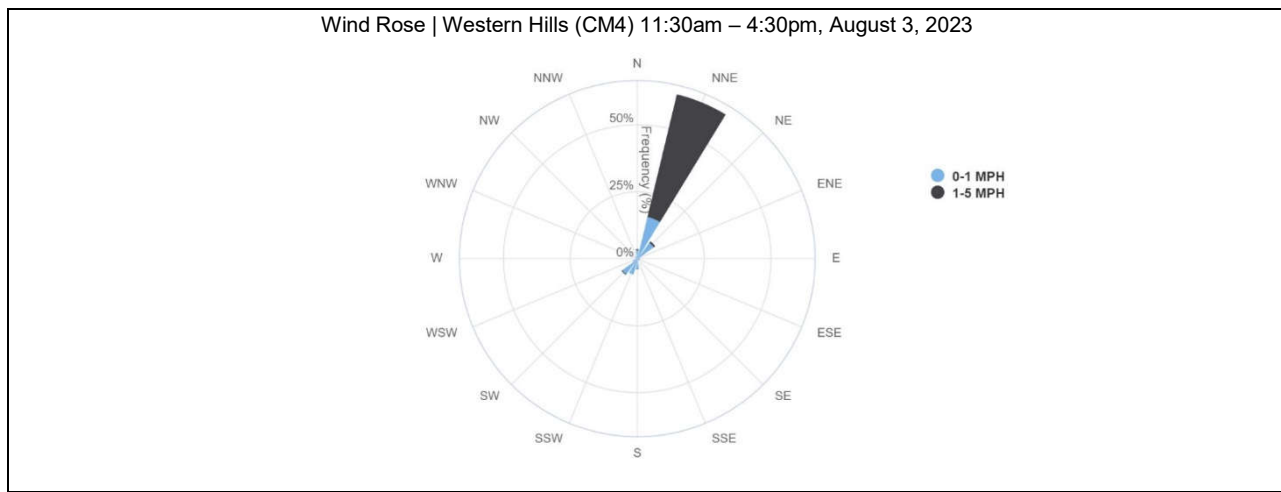
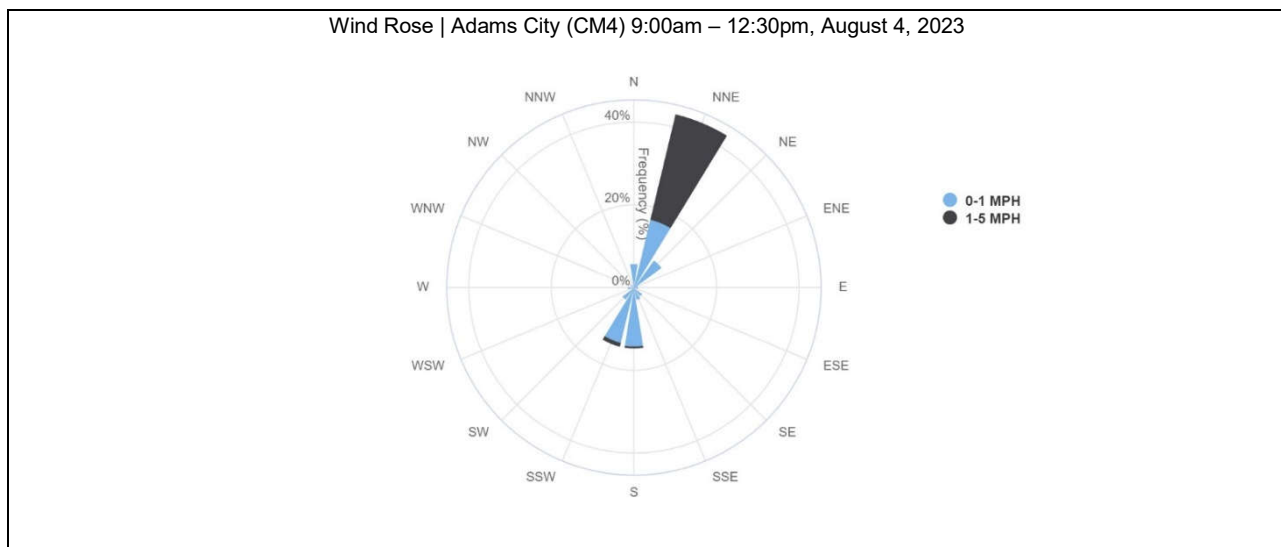
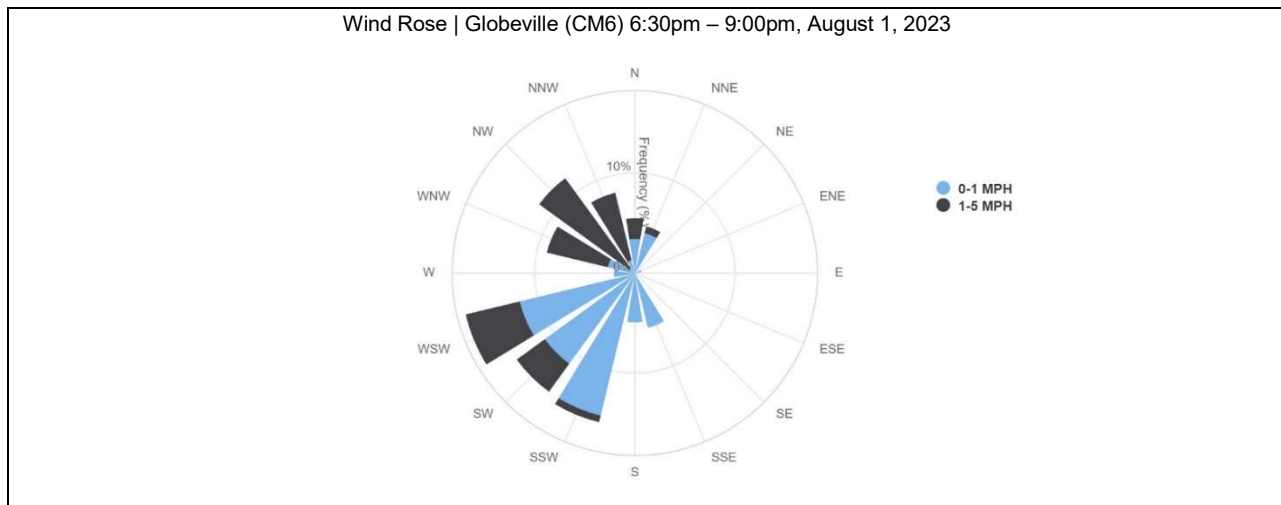
Group Name	Specific Isomers	Group Name	Specific Isomers
Butenes	1-Butene cis-2-Butene trans-2-Butene	Xylenes	Ethyl Benzene o-Xylene m-Xylene p-Xylene
Butanes	iso-Butane n-Butane	Dimethylcyclohexanes	Ethylcyclohexane cis-1,3-Dimethylcyclohexane trans-1,2- Dimethylcyclohexane trans-1,3- Dimethylcyclohexane
Cyclopentanes	Cyclopentane 1-Pentene 2-Methyl-2-butene cis-2-Pentene trans-2-Pentene	Octanes	n-Octane 2-Methylheptane 3-Methylheptane 2,2,4-Trimethylpentane 2,3,4-Trimethylpentane
Pentanes	iso-Pentane n-Pentane neo-Pentane	Trimethylbenzenes	Cumene 1,2,4-Trimethylbenzene o-Ethyltoluene m-Ethyltoluene p-Ethyltoluene n-Propylbenzene 1,3,5-Trimethylbenzene
Hexenes	1-Hexene Cyclohexane Methylcyclopentane	Diethylbenzenes	o-Diethylbenzene m-Diethylbenzene p-Diethylbenzene All other C ₁₀ H ₁₄ Isomers
Hexanes	n-Hexane 2-Methylpentane 3-Methylpentane 2,2-Dimethylbutane 2,3-Dimethylbutane		
Heptanes	n-Heptane 2-Methylhexane 3-Methylhexane 2,3- Dimethylpentane 2,4- Dimethylpentane		

APPENDIX B DAILY WIND ROSES

CCND Mobile Monitoring Van
2023 Q3



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2023 Q3



APPENDIX C
SCREENING RISK ASSESSMENT DETAILS
(ALPHABETICAL ORDER BY NEIGHBORHOOD NAME)

CCND Mobile Monitoring Van

2023 Q3

Mobile Laboratory Sampling Data Summary and Risk Assessment

Adams City Neighborhood | August 4, 2023

Analyte	Cas No	Count of 1-second Concentrations (#)	Maximum 1-second Concentration (ppbv)	Count of 1-hr Rolling Averages Derived (#)	Average 1-hr Rolling Average (ppbv)	Maximum 1-hr Rolling Average (ppbv)	AEGL 1 60-min Value	Health Reference Level (ppbv)	Screening Value Source	Hazard Quotient
1,3 BUTADIENE	106-99-0	10,146	0.12	6,619	0.01	0.01	670,000	298	OEHHA Acute REL	0.00004
ACETYLENE	74-86-2	10,146	0.73	6,619	0.09	0.11	NR	25,000	TCEQ Short-Term AMCV Health	0.00000
BENZENE	71-43-2	10,146	24.83	6,619	0.37	0.42	52,000	9	ATSDR Acute MRL	0.04685
BUTANES*	75-28-5	10,146	14.87	6,619	2.17	2.67	NR	33000	TCEQ Short-Term AMCV Health	0.00008
BUTENES*	590-18-1	10,146	145.92	6,619	1.99	2.75	NR	15000	TCEQ Short-Term AMCV Health	0.00018
CARBON DISULFIDE	75-15-0	10,146	0.05	6,619	0.00	0.00	13,000	1,991	OEHHA Acute REL	0.00000
CYCLOPENTANES*	287-92-3	10,146	67.83	6,619	2.19	2.63	NR	5,900	TCEQ Short-Term AMCV Health	0.00045
DECANES	124-18-5	10,146	0.04	6,619	0.00	0.00	NR	1,000	TCEQ Short-Term AMCV Health	0.00000
DIETHYLBENZENES*	141-93-5	10,146	0.23	6,619	0.08	0.09	NR	450	TCEQ Short-Term AMCV Health	0.00020
DIMETHYLCYCLOHEXANES*	638-04-0	10,146	0.91	6,619	0.04	0.04	NR	4,000	CDPHE	0.00001
DODECANES	112-40-3	10,146	0.01	6,619	0.00	0.00	NR	1720	CDPHE	0.00000
ETHYLENE	74-85-1	10,146	9.83	6,619	5.30	5.31	NR	500,000	TCEQ Short-Term AMCV Health	0.00001
HEPTANES*	142-82-5	10,146	1.21	6,619	0.08	0.10	NR	8,300	TCEQ Short-Term AMCV Health	0.00001
HEXANES*	110-54-3	10,146	0.32	6,619	0.06	0.07	NR	5,400	TCEQ Short-Term AMCV Health	0.00001
HEXENES*	592-41-6	10,146	37.41	6,619	0.62	0.83	NR	500	TCEQ Short-Term AMCV Health	0.00166
HYDROGEN CYANIDE	74-90-8	10,146	0.55	6,619	0.16	0.18	2,000	308	OEHHA Acute REL	0.00058
HYDROGEN SULFIDE	7783-06-4	10,146	0.57	6,619	0.20	0.22	510	70	ATSDR Acute MRL	0.00307
ISOPRENE	78-79-5	10,146	3.93	6,619	0.21	0.24	NR	1,400	TCEQ Short-Term AMCV Health	0.00017
METHANOL	67-56-1	10,146	20.12	6,619	6.83	6.98	530,000	21,366	OEHHA Acute REL	0.00033
METHYLCYCLOHEXANE	108-87-2	10,146	1.60	6,619	0.03	0.04	NR	4,000	TCEQ Short-Term AMCV Health	0.00001
NONANES	111-84-2	10,146	0.09	6,619	0.03	0.03	NR	3,000	TCEQ Short-Term AMCV Health	0.00001
OCTANES*	111-65-9	10,146	0.16	6,619	0.04	0.05	NR	4,100	TCEQ Short-Term AMCV Health	0.00001
PENTANES*	109-66-0	10,146	0.49	6,619	0.04	0.05	NR	68,000	TCEQ Short-Term AMCV Health	0.00000
PROPYLENE	115-07-1	10,146	27.98	6,619	0.52	0.67	NR	NA	NE	
STYRENE	100-42-5	10,146	0.13	6,619	0.00	0.00	20,000	5,000	ATSDR Acute MRL	0.00000
TETRACHLOROETHYLENE	127-18-4	10,146	0.04	6,619	0.00	0.00	35,000	6	ATSDR Acute MRL	0.00083
TOLUENE	108-88-3	10,146	158.26	6,619	0.70	1.04	67,000	2,000	ATSDR Acute MRL	0.00052
TRIMETHYLBENZENES*	622-96-8	10,146	13.30	6,619	0.38	0.46	50,000	250	TCEQ Short-Term AMCV Health	0.00183
UNDECANES	1120-21-4	10,146	0.04	6,619	0.00	0.00	NR	550	TCEQ Short-Term AMCV Health	0.00000
XYLENES*	1330-20-7	10,146	104.29	6,619	0.83	1.15	130,000	2,000	ATSDR Acute MRL	0.00057
								Hazard Index	0.05745	

NR= According to EPA, AEGL is "not recommended due to insufficient data"

NA= Not Available

NC= Not Calculated

For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the isomer group was selected for use in this assessment (See Appendix A)

CCND Mobile Monitoring Van

2023 Q3

Mobile Laboratory Sampling Data Summary and Risk Assessment

Dupont Neighborhood | August 2, 2023

Analyte	Cas No	Count of 1-second Concentrations (#)	Maximum 1-second Concentration (ppbv)	Count of 1-hr Rolling Averages Derived (#)	Average 1-hr Rolling Average (ppbv)	Maximum 1-hr Rolling Average (ppbv)	AEGL 1 60-min Value	Health Reference Level (ppbv)	Screening Value Source	Hazard Quotient
1,3 BUTADIENE	106-99-0	11,273	0.12	7,746	0.01	0.01	670,000	298	OEHHA Acute REL	0.00005
ACETYLENE	74-86-2	11,273	0.85	7,746	0.11	0.17	NR	25,000	TCEQ Short-Term AMCV Health	0.00001
BENZENE	71-43-2	11,273	7.09	7,746	0.17	0.27	52,000	9	ATSDR Acute MRL	0.03037
BUTANES*	75-28-5	11,273	23.29	7,746	3.66	4.42	NR	33000	TCEQ Short-Term AMCV Health	0.00013
BUTENES*	590-18-1	11,273	49.73	7,746	1.70	2.32	NR	15000	TCEQ Short-Term AMCV Health	0.00015
CARBON DISULFIDE	75-15-0	11,273	0.03	7,746	0.00	0.00	13,000	1,991	OEHHA Acute REL	0.00000
CYCLOPENTANES*	287-92-3	11,273	42.30	7,746	2.53	3.57	NR	5,900	TCEQ Short-Term AMCV Health	0.00060
DECANES	124-18-5	11,273	0.06	7,746	0.01	0.02	NR	1,000	TCEQ Short-Term AMCV Health	0.00002
DIETHYLBENZENES*	141-93-5	11,273	0.28	7,746	0.10	0.12	NR	450	TCEQ Short-Term AMCV Health	0.00027
DIMETHYLCYCLOHEXANES*	638-04-0	11,273	0.13	7,746	0.04	0.04	NR	4,000	CDPHE	0.00001
DODECANES	112-40-3	11,273	0.05	7,746	0.00	0.00	NR	1720	CDPHE	0.00000
ETHYLENE	74-85-1	11,273	8.65	7,746	7.24	7.26	NR	500,000	TCEQ Short-Term AMCV Health	0.00001
HEPTANES*	142-82-5	11,273	0.22	7,746	0.11	0.12	NR	8,300	TCEQ Short-Term AMCV Health	0.00001
HEXANES*	110-54-3	11,273	0.24	7,746	0.07	0.08	NR	5,400	TCEQ Short-Term AMCV Health	0.00001
HEXENES*	592-41-6	11,273	8.76	7,746	0.61	0.92	NR	500	TCEQ Short-Term AMCV Health	0.00184
HYDROGEN CYANIDE	74-90-8	11,273	0.88	7,746	0.18	0.24	2,000	308	OEHHA Acute REL	0.00077
HYDROGEN SULFIDE	7783-06-4	11,273	0.69	7,746	0.13	0.18	510	70	ATSDR Acute MRL	0.00256
ISOPRENE	78-79-5	11,273	2.04	7,746	0.16	0.21	NR	1,400	TCEQ Short-Term AMCV Health	0.00015
METHANOL	67-56-1	11,273	246.58	7,746	3.57	4.04	530,000	21,366	OEHHA Acute REL	0.00019
METHYLCYCLOHEXANE	108-87-2	11,273	0.26	7,746	0.04	0.05	NR	4,000	TCEQ Short-Term AMCV Health	0.00001
NONANES	111-84-2	11,273	0.07	7,746	0.01	0.01	NR	3,000	TCEQ Short-Term AMCV Health	0.00000
OCTANES*	111-65-9	11,273	0.10	7,746	0.04	0.05	NR	4,100	TCEQ Short-Term AMCV Health	0.00001
PENTANES*	109-66-0	11,273	0.49	7,746	0.05	0.08	NR	68,000	TCEQ Short-Term AMCV Health	0.00000
PROPYLENE	115-07-1	11,273	21.63	7,746	0.27	0.51	NR	NA	NE	
STYRENE	100-42-5	11,273	0.39	7,746	0.03	0.05	20,000	5,000	ATSDR Acute MRL	0.00001
TETRACHLOROETHYLENE	127-18-4	11,273	0.11	7,746	0.01	0.01	35,000	6	ATSDR Acute MRL	0.00147
TOLUENE	108-88-3	11,273	19.89	7,746	0.57	0.93	67,000	2,000	ATSDR Acute MRL	0.00047
TRIMETHYLBENZENES*	622-96-8	11,273	3.75	7,746	0.10	0.21	50,000	250	TCEQ Short-Term AMCV Health	0.00084
UNDECANES	1120-21-4	11,273	0.05	7,746	0.01	0.01	NR	550	TCEQ Short-Term AMCV Health	0.00001
XYLENES*	1330-20-7	11,273	15.10	7,746	0.48	0.98	130,000	2,000	ATSDR Acute MRL	0.00049
								Hazard Index	0.04051	

NR=According to EPA, AEGL is "not recommended due to insufficient data"

NA= Not Available

NC= Not Calculated

For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the isomer group was selected for use in this assessment (See Appendix A)

CCND Mobile Monitoring Van

2023 Q3

Mobile Laboratory Sampling Data Summary and Risk Assessment

Elyria-Swansea Neighborhood | August 1, 2023

Analyte	Cas No	Count of 1-second Concentrations (#)	Maximum 1-second Concentration (ppbv)	Count of 1-hr Rolling Averages Derived (#)	Average 1-hr Rolling Average (ppbv)	Maximum 1-hr Rolling Average (ppbv)	AEGL 1 60-min Value	Health Reference Level (ppbv)	Screening Value Source	Hazard Quotient
1,3 BUTADIENE	106-99-0	7,613	0.11	4,086	0.00	0.00	670,000	298	OEHHA Acute REL	0.00002
ACETYLENE	74-86-2	7,613	0.92	4,086	0.15	0.22	NR	25,000	TCEQ Short-Term AMCV Health	0.00001
BENZENE	71-43-2	7,613	14.73	4,086	0.25	0.44	52,000	9	ATSDR Acute MRL	0.04845
BUTANES*	75-28-5	7,613	5.46	4,086	1.31	2.27	NR	33000	TCEQ Short-Term AMCV Health	0.00007
BUTENES*	590-18-1	7,613	30.56	4,086	0.96	1.87	NR	15000	TCEQ Short-Term AMCV Health	0.00012
CARBON DISULFIDE	75-15-0	7,613	0.02	4,086	0.00	0.00	13,000	1,991	OEHHA Acute REL	0.00000
CYCLOPENTANES*	287-92-3	7,613	23.15	4,086	0.69	1.72	NR	5,900	TCEQ Short-Term AMCV Health	0.00029
DECANES	124-18-5	7,613	0.17	4,086	0.07	0.09	NR	1,000	TCEQ Short-Term AMCV Health	0.00009
DIETHYLBENZENES*	141-93-5	7,613	0.20	4,086	0.06	0.09	NR	450	TCEQ Short-Term AMCV Health	0.00019
DIMETHYLCYCLOHEXANES*	638-04-0	7,613	0.18	4,086	0.04	0.05	NR	4,000	CDPHE	0.00001
DODECANES	112-40-3	7,613	0.01	4,086	0.00	0.00	NR	1720	CDPHE	0.00000
ETHYLENE	74-85-1	7,613	7.85	4,086	7.43	7.44	NR	500,000	TCEQ Short-Term AMCV Health	0.00001
HEPTANES*	142-82-5	7,613	0.27	4,086	0.12	0.15	NR	8,300	TCEQ Short-Term AMCV Health	0.00002
HEXANES*	110-54-3	7,613	0.34	4,086	0.11	0.15	NR	5,400	TCEQ Short-Term AMCV Health	0.00003
HEXENES*	592-41-6	7,613	10.20	4,086	0.42	0.81	NR	500	TCEQ Short-Term AMCV Health	0.00162
HYDROGEN CYANIDE	74-90-8	7,613	0.76	4,086	0.13	0.14	2,000	308	OEHHA Acute REL	0.00046
HYDROGEN SULFIDE	7783-06-4	7,613	0.39	4,086	0.07	0.09	510	70	ATSDR Acute MRL	0.00134
ISOPRENE	78-79-5	7,613	1.27	4,086	0.13	0.23	NR	1,400	TCEQ Short-Term AMCV Health	0.00016
METHANOL	67-56-1	7,613	16.85	4,086	5.63	6.68	530,000	21,366	OEHHA Acute REL	0.00031
METHYLCYCLOHEXANE	108-87-2	7,613	0.29	4,086	0.07	0.10	NR	4,000	TCEQ Short-Term AMCV Health	0.00002
NONANES	111-84-2	7,613	0.14	4,086	0.04	0.05	NR	3,000	TCEQ Short-Term AMCV Health	0.00002
OCTANES*	111-65-9	7,613	0.21	4,086	0.07	0.09	NR	4,100	TCEQ Short-Term AMCV Health	0.00002
PENTANES*	109-66-0	7,613	0.52	4,086	0.27	0.28	NR	68,000	TCEQ Short-Term AMCV Health	0.00000
PROPYLENE	115-07-1	7,613	11.75	4,086	0.58	0.93	NR	NA	NE	
STYRENE	100-42-5	7,613	0.21	4,086	0.03	0.06	20,000	5,000	ATSDR Acute MRL	0.00001
TETRACHLOROETHYLENE	127-18-4	7,613	0.06	4,086	0.01	0.01	35,000	6	ATSDR Acute MRL	0.00185
TOLUENE	108-88-3	7,613	12.94	4,086	1.37	2.06	67,000	2,000	ATSDR Acute MRL	0.00103
TRIMETHYLBENZENES*	622-96-8	7,613	3.37	4,086	0.20	0.38	50,000	250	TCEQ Short-Term AMCV Health	0.00154
UNDECANES	1120-21-4	7,613	0.13	4,086	0.06	0.07	NR	550	TCEQ Short-Term AMCV Health	0.00013
XYLENES*	1330-20-7	7,613	14.70	4,086	0.98	1.70	130,000	2,000	ATSDR Acute MRL	0.00085
								Hazard Index	0.05867	

NR=According to EPA, AEGL is "not recommended due to insufficient data"

NA= Not Available

NC= Not Calculated

For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the isomer group was selected for use in this assessment (See Appendix A)

CCND Mobile Monitoring Van

2023 Q3

Mobile Laboratory Sampling Data Summary and Risk Assessment Globeville Neighborhood | August 1, 2023

Analyte	Cas No	Count of 1-second Concentrations (#)	Maximum 1-second Concentration (ppbv)	Count of 1-hr Rolling Averages Derived (#)	Average 1-hr Rolling Average (ppbv)	Maximum 1-hr Rolling Average (ppbv)	AEGL 1 60-min Value	Health Reference Level (ppbv)	Screening Value Source	Hazard Quotient
1,3 BUTADIENE	106-99-0	7,626	0.11	4,099	0.01	0.01	670,000	298	OEHHA Acute REL	0.00002
ACETYLENE	74-86-2	7,626	0.93	4,099	0.19	0.21	NR	25,000	TCEQ Short-Term AMCV Health	0.00001
BENZENE	71-43-2	7,626	2.94	4,099	0.43	0.47	52,000	9	ATSDR Acute MRL	0.05200
BUTANES*	75-28-5	7,626	27.94	4,099	3.75	4.58	NR	33000	TCEQ Short-Term AMCV Health	0.00014
BUTENES*	590-18-1	7,626	25.63	4,099	2.20	2.61	NR	15000	TCEQ Short-Term AMCV Health	0.00017
CARBON DISULFIDE	75-15-0	7,626	0.02	4,099	0.00	0.00	13,000	1,991	OEHHA Acute REL	0.00000
CYCLOPENTANES*	287-92-3	7,626	28.56	4,099	2.41	2.82	NR	5,900	TCEQ Short-Term AMCV Health	0.00048
DECANES	124-18-5	7,626	0.19	4,099	0.10	0.10	NR	1,000	TCEQ Short-Term AMCV Health	0.00010
DIETHYLBENZENES*	141-93-5	7,626	0.37	4,099	0.11	0.14	NR	450	TCEQ Short-Term AMCV Health	0.00031
DIMETHYLCYCLOHEXANES*	638-04-0	7,626	0.13	4,099	0.06	0.06	NR	4,000	CDPHE	0.00001
DODECANES	112-40-3	7,626	0.01	4,099	0.00	0.00	NR	1720	CDPHE	0.00000
ETHYLENE	74-85-1	7,626	7.97	4,099	7.45	7.47	NR	500,000	TCEQ Short-Term AMCV Health	0.00001
HEPTANES*	142-82-5	7,626	0.42	4,099	0.22	0.24	NR	8,300	TCEQ Short-Term AMCV Health	0.00003
HEXANES*	110-54-3	7,626	0.45	4,099	0.18	0.19	NR	5,400	TCEQ Short-Term AMCV Health	0.00003
HEXENES*	592-41-6	7,626	5.14	4,099	1.01	1.26	NR	500	TCEQ Short-Term AMCV Health	0.00252
HYDROGEN CYANIDE	74-90-8	7,626	0.55	4,099	0.19	0.25	2,000	308	OEHHA Acute REL	0.00081
HYDROGEN SULFIDE	7783-06-4	7,626	0.50	4,099	0.11	0.14	510	70	ATSDR Acute MRL	0.00204
ISOPRENE	78-79-5	7,626	1.17	4,099	0.30	0.31	NR	1,400	TCEQ Short-Term AMCV Health	0.00022
METHANOL	67-56-1	7,626	51.34	4,099	6.37	7.01	530,000	21,366	OEHHA Acute REL	0.00033
METHYLCYCLOHEXANE	108-87-2	7,626	0.25	4,099	0.12	0.13	NR	4,000	TCEQ Short-Term AMCV Health	0.00003
NONANES	111-84-2	7,626	0.15	4,099	0.06	0.06	NR	3,000	TCEQ Short-Term AMCV Health	0.00002
OCTANES*	111-65-9	7,626	0.35	4,099	0.11	0.12	NR	4,100	TCEQ Short-Term AMCV Health	0.00003
PENTANES*	109-66-0	7,626	1.16	4,099	0.28	0.35	NR	68,000	TCEQ Short-Term AMCV Health	0.00001
PROPYLENE	115-07-1	7,626	6.19	4,099	1.13	1.28	NR	NA	NE	
STYRENE	100-42-5	7,626	0.37	4,099	0.08	0.10	20,000	5,000	ATSDR Acute MRL	0.00002
TETRACHLOROETHYLENE	127-18-4	7,626	0.07	4,099	0.01	0.01	35,000	6	ATSDR Acute MRL	0.00220
TOLUENE	108-88-3	7,626	11.72	4,099	0.79	1.55	67,000	2,000	ATSDR Acute MRL	0.00078
TRIMETHYLBENZENES*	622-96-8	7,626	1.33	4,099	0.42	0.49	50,000	250	TCEQ Short-Term AMCV Health	0.00195
UNDECANES	1120-21-4	7,626	0.13	4,099	0.08	0.08	NR	550	TCEQ Short-Term AMCV Health	0.00014
XYLENES*	1330-20-7	7,626	7.23	4,099	1.83	2.02	130,000	2,000	ATSDR Acute MRL	0.00101
								Hazard Index	0.06543	

NR=According to EPA, AEGL is "not recommended due to insufficient data"

NA= Not Available

NC= Not Calculated

For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the isomer group was selected for use in this assessment (See Appendix A)

CCND Mobile Monitoring Van

2023 Q3

Mobile Laboratory Sampling Data Summary and Risk Assessment

Pioneer Park Neighborhood | July 31, 2023

Analyte	Cas No	Count of 1-second Concentrations (#)	Maximum 1-second Concentration (ppbv)	Count of 1-hr Rolling Averages Derived (#)	Average 1-hr Rolling Average (ppbv)	Maximum 1-hr Rolling Average (ppbv)	AEGL 1 60-min Value	Health Reference Level (ppbv)	Screening Value Source	Hazard Quotient
1,3 BUTADIENE	106-99-0	14,772	0.15	11,245	0.02	0.02	670,000	298	OEHHA Acute REL	0.00006
ACETYLENE	74-86-2	14,772	3.87	11,245	0.19	0.31	NR	25,000	TCEQ Short-Term AMCV Health	0.00001
BENZENE	71-43-2	14,772	5.63	11,245	0.16	0.22	52,000	9	ATSDR Acute MRL	0.02452
BUTANES*	75-28-5	14,772	19.65	11,245	3.62	4.38	NR	33000	TCEQ Short-Term AMCV Health	0.00013
BUTENES*	590-18-1	14,772	29.51	11,245	2.62	3.25	NR	15000	TCEQ Short-Term AMCV Health	0.00022
CARBON DISULFIDE	75-15-0	14,772	0.04	11,245	0.00	0.00	13,000	1,991	OEHHA Acute REL	0.00000
CYCLOPENTANES*	287-92-3	14,772	30.28	11,245	1.61	2.65	NR	5,900	TCEQ Short-Term AMCV Health	0.00045
DECANES	124-18-5	14,772	0.13	11,245	0.06	0.07	NR	1,000	TCEQ Short-Term AMCV Health	0.00007
DIETHYLBENZENES*	141-93-5	14,772	0.29	11,245	0.05	0.13	NR	450	TCEQ Short-Term AMCV Health	0.00028
DIMETHYLCYCLOHEXANES*	638-04-0	14,772	0.12	11,245	0.02	0.03	NR	4,000	CDPHE	0.00001
DODECANES	112-40-3	14,772	0.01	11,245	0.00	0.00	NR	1720	CDPHE	0.00000
ETHYLENE	74-85-1	14,772	28.08	11,245	6.81	7.11	NR	500,000	TCEQ Short-Term AMCV Health	0.00001
HEPTANES*	142-82-5	14,772	0.40	11,245	0.18	0.21	NR	8,300	TCEQ Short-Term AMCV Health	0.00002
HEXANES*	110-54-3	14,772	0.40	11,245	0.13	0.15	NR	5,400	TCEQ Short-Term AMCV Health	0.00003
HEXENES*	592-41-6	14,772	6.42	11,245	1.03	1.29	NR	500	TCEQ Short-Term AMCV Health	0.00258
HYDROGEN CYANIDE	74-90-8	14,772	1.24	11,245	0.12	0.14	2,000	308	OEHHA Acute REL	0.00047
HYDROGEN SULFIDE	7783-06-4	14,772	0.61	11,245	0.21	0.25	510	70	ATSDR Acute MRL	0.00364
ISOPRENE	78-79-5	14,772	1.18	11,245	0.16	0.20	NR	1,400	TCEQ Short-Term AMCV Health	0.00014
METHANOL	67-56-1	14,772	29.07	11,245	2.59	3.39	530,000	21,366	OEHHA Acute REL	0.00016
METHYLCYCLOHEXANE	108-87-2	14,772	0.21	11,245	0.05	0.07	NR	4,000	TCEQ Short-Term AMCV Health	0.00002
NONANES	111-84-2	14,772	0.10	11,245	0.02	0.03	NR	3,000	TCEQ Short-Term AMCV Health	0.00001
OCTANES*	111-65-9	14,772	0.20	11,245	0.07	0.08	NR	4,100	TCEQ Short-Term AMCV Health	0.00002
PENTANES*	109-66-0	14,772	0.56	11,245	0.26	0.27	NR	68,000	TCEQ Short-Term AMCV Health	0.00000
PROPYLENE	115-07-1	14,772	6.85	11,245	0.39	0.58	NR	NA	NE	
STYRENE	100-42-5	14,772	0.27	11,245	0.01	0.04	20,000	5,000	ATSDR Acute MRL	0.00001
TETRACHLOROETHYLENE	127-18-4	14,772	0.04	11,245	0.00	0.01	35,000	6	ATSDR Acute MRL	0.00088
TOLUENE	108-88-3	14,772	18.19	11,245	0.57	0.83	67,000	2,000	ATSDR Acute MRL	0.00042
TRIMETHYLBENZENES*	622-96-8	14,772	1.65	11,245	0.13	0.21	50,000	250	TCEQ Short-Term AMCV Health	0.00082
UNDECANES	1120-21-4	14,772	0.06	11,245	0.01	0.01	NR	550	TCEQ Short-Term AMCV Health	0.00003
XYLENES*	1330-20-7	14,772	18.92	11,245	0.65	1.20	130,000	2,000	ATSDR Acute MRL	0.00060
								Hazard Index	0.03561	

NR=According to EPA, AEGL is "not recommended due to insufficient data"

NA= Not Available

NC= Not Calculated

For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the isomer group was selected for use in this assessment (See Appendix A)

CCND Mobile Monitoring Van

2023 Q3

Mobile Laboratory Sampling Data Summary and Risk Assessment

Western Hills Neighborhood | August 3, 2023

Analyte	Cas No	Count of 1-second Concentrations (#)	Maximum 1-second Concentration (ppbv)	Count of 1-hr Rolling Averages Derived (#)	Average 1-hr Rolling Average (ppbv)	Maximum 1-hr Rolling Average (ppbv)	AEGL 1 60-min Value	Health Reference Level (ppbv)	Screening Value Source	Hazard Quotient
1,3 BUTADIENE	106-99-0	15,154	0.11	11,627	0.01	0.01	670,000	298	OEHHA Acute REL	0.00004
ACETYLENE	74-86-2	15,154	0.76	11,627	0.10	0.16	NR	25,000	TCEQ Short-Term AMCV Health	0.00001
BENZENE	71-43-2	15,154	2.08	11,627	0.24	0.35	52,000	9	ATSDR Acute MRL	0.03896
BUTANES*	75-28-5	15,154	13.43	11,627	2.51	3.23	NR	33000	TCEQ Short-Term AMCV Health	0.00010
BUTENES*	590-18-1	15,154	15.22	11,627	1.35	2.22	NR	15000	TCEQ Short-Term AMCV Health	0.00015
CARBON DISULFIDE	75-15-0	15,154	0.02	11,627	0.00	0.00	13,000	1,991	OEHHA Acute REL	0.00000
CYCLOPENTANES*	287-92-3	15,154	16.90	11,627	1.27	2.96	NR	5,900	TCEQ Short-Term AMCV Health	0.00050
DECANES	124-18-5	15,154	0.13	11,627	0.05	0.06	NR	1,000	TCEQ Short-Term AMCV Health	0.00006
DIETHYLBENZENES*	141-93-5	15,154	0.34	11,627	0.07	0.10	NR	450	TCEQ Short-Term AMCV Health	0.00023
DIMETHYLCYCLOHEXANES*	638-04-0	15,154	0.12	11,627	0.03	0.04	NR	4,000	CDPHE	0.00001
DODECANES	112-40-3	15,154	0.01	11,627	0.00	0.00	NR	1720	CDPHE	0.00000
ETHYLENE	74-85-1	15,154	8.66	11,627	7.21	7.23	NR	500,000	TCEQ Short-Term AMCV Health	0.00001
HEPTANES*	142-82-5	15,154	0.37	11,627	0.09	0.10	NR	8,300	TCEQ Short-Term AMCV Health	0.00001
HEXANES*	110-54-3	15,154	0.38	11,627	0.10	0.13	NR	5,400	TCEQ Short-Term AMCV Health	0.00002
HEXENES*	592-41-6	15,154	4.62	11,627	0.46	0.84	NR	500	TCEQ Short-Term AMCV Health	0.00169
HYDROGEN CYANIDE	74-90-8	15,154	0.49	11,627	0.14	0.19	2,000	308	OEHHA Acute REL	0.00063
HYDROGEN SULFIDE	7783-06-4	15,154	0.45	11,627	0.13	0.17	510	70	ATSDR Acute MRL	0.00238
ISOPRENE	78-79-5	15,154	0.78	11,627	0.20	0.25	NR	1,400	TCEQ Short-Term AMCV Health	0.00018
METHANOL	67-56-1	15,154	17.87	11,627	5.42	6.02	530,000	21,366	OEHHA Acute REL	0.00028
METHYLCYCLOHEXANE	108-87-2	15,154	0.26	11,627	0.08	0.10	NR	4,000	TCEQ Short-Term AMCV Health	0.00002
NONANES	111-84-2	15,154	0.08	11,627	0.01	0.02	NR	3,000	TCEQ Short-Term AMCV Health	0.00001
OCTANES*	111-65-9	15,154	0.20	11,627	0.06	0.08	NR	4,100	TCEQ Short-Term AMCV Health	0.00002
PENTANES*	109-66-0	15,154	0.44	11,627	0.03	0.03	NR	68,000	TCEQ Short-Term AMCV Health	0.00000
PROPYLENE	115-07-1	15,154	3.46	11,627	0.17	0.46	NR	NA	NE	
STYRENE	100-42-5	15,154	1.19	11,627	0.10	0.13	20,000	5,000	ATSDR Acute MRL	0.00003
TETRACHLOROETHYLENE	127-18-4	15,154	0.05	11,627	0.00	0.01	35,000	6	ATSDR Acute MRL	0.00146
TOLUENE	108-88-3	15,154	7.66	11,627	0.74	1.15	67,000	2,000	ATSDR Acute MRL	0.00057
TRIMETHYLBENZENES*	622-96-8	15,154	1.77	11,627	0.25	0.36	50,000	250	TCEQ Short-Term AMCV Health	0.00143
UNDECANES	1120-21-4	15,154	0.10	11,627	0.04	0.05	NR	550	TCEQ Short-Term AMCV Health	0.00009
XYLENES*	1330-20-7	15,154	9.01	11,627	0.91	1.43	130,000	2,000	ATSDR Acute MRL	0.00071
								Hazard Index	0.04963	

NR=According to EPA, AEGL is "not recommended due to insufficient data"

NA= Not Available

NC= Not Calculated

For analyte isomer groups which were unable to be differentiated, the lowest health reference value of the isomer group was selected for use in this assessment (See Appendix A)

APPENDIX D

PTR CALIBRATION AND QA/QC DATA

Notable Sampling Events During Test Program

7-31-23 Pioneer Park Neighborhood

11:57 66th and Newport: Benzene, toluene and xylene spike, unknow source.
13:11 Holly and 64th: Hexenes, Circle K Gas Station.

8-1-23 Globeville and Elyria-Swansea Neighborhoods

17:29 Lincoln and 43rd: Benzene, toluene auto exhaust.
19:28 York and 47th: Benzene, Trimethylbenzenes, train exhaust

8-2-23 Dupont Neighborhood

17:03 Intersection: Benzene, toluene car exhaust
19:15 Ivy and 72nd: Hexenes, trimethylbenzenes, benzene car exhaust

8-3-23 Western Hills Neighborhood

No notable sampling occurrences

8-4-23 Adams City Neighborhood

11:37 Gas Station: Benzene, Toluene, Trimethylbenzenes and hexenes, Van refueling

CCND Mobile Monitoring Van
2023 Q3

Commerce City North Denver Neighborhood Monitoring
Third Quarter 2023
PTR Screen Shots

7-30-23 Initial Calibration Screen Shots.

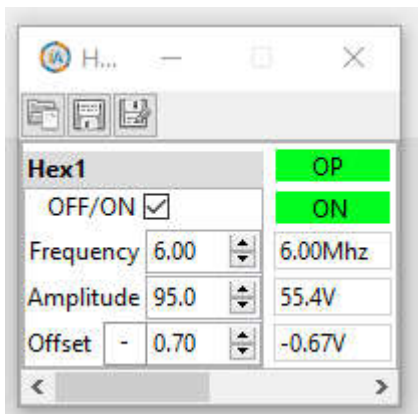
The screenshot shows a software interface for PTR calibration. At the top, there are icons for file operations (New, Open, Save, Print) and a search function. Below this is a table for setting parameters:

Setting	Odor	▼	▼
Primary Ion	H ₃ O ⁺	▼	▼
Transmission	DC	▼	▼

Below the settings are two columns: "Man/Ctrl" and "Ctrl". The "Man/Ctrl" column contains numerical values with up/down arrows for adjustment, and the "Ctrl" column shows the current controlled values. The parameters listed are:

PC	343.9	343.88 mbar		
p Drift	2.30	2.31 mbar		
TofLens		4.79E-5 mbar		
TOF		2.21E-6 mbar		
E/N		119 Td		
Temps	79.90 °C	80.10 °C		
SrcValve	50.0			
H ₂ O	6.0	6.00 sccm		
O ₂	0.0	0.00 sccm		
NO	0.0	0.00 sccm		
Ihc	4	4.0 mA		
	On/Off	On		
FCinlet	60.0	60.01 sccm		
U	FU	°C	D+	D-
Us	150	145.0 V		
Uso	80	78.6 V		
Udrift	525	526.1 V		

Production Settings



Hex Settings

TPS 4-6-23 MCP Tune.iTPS *Changed*				
Lens 1	14.0	14.0 V	All on	<input checked="" type="checkbox"/>
Lens 2	30.0	30.0 V	Lenses	<input checked="" type="checkbox"/>
Lens 3	20.0	20.0 V		
Lens 4	60.0	60.0 V		
Lens 5	70.0	69.0 V		
Lens 6	80.0	80.0 V		
Lens 7	17.0	18.0 V		
Push L	16.5	16.0 V	<input checked="" type="checkbox"/>	3 mA
Push H	790.0	790.0 V	<input checked="" type="checkbox"/>	3 mA
Pull L	80.0	80.0 V	<input checked="" type="checkbox"/>	3 mA
Pull H	680.0	680.0 V	<input checked="" type="checkbox"/>	3 mA
Grid	2400.0	2282.0 V	<input checked="" type="checkbox"/>	1 µA
Cage	5020.0	4766 V	<input checked="" type="checkbox"/>	99 µA
Refl. Grid	667.0	634.0 V	<input checked="" type="checkbox"/>	76 µA
Refl. Back	900.0	855.0 V	<input checked="" type="checkbox"/>	167 µA
MCP F	5400	5134 V	<input checked="" type="checkbox"/>	17 µA
MCP B	2496	2381 V	<input checked="" type="checkbox"/>	214 µA

TOF Lenses

CCND Mobile Monitoring Van
2023 Q3

Defined Peaks

	Mass	Value	Unit
*(H ₂ O)+	18.01000	19.27	ppb
*(H ₃ N)H+	18.03380	336.18	ppb
*(H ₂ O)H+	19.01780	14.45	ppb
✓ *(H ₂ O)H+	21.02210	1.19E+5	ppb
[HCN]+	27.02000	2.33	ppb
*(N ₂)+	28.00600	0.00	ppb
✓ (HCN)H+	28.02100	0.79	ppb
✓ (C ₂ H ₄)+	28.03508	12.30	ppb
*(N ₂)H+	29.01340	107.96	ppb
Ethylene[C ₂ H ₄]	29.04400	1.25	ppb
*(NO)+ [NO+]	29.99740	196.11	ppb

21 of 239 Peaks selected from
"2-5-23 Suncor Working Peak Table.upta"

Instrument

Description	Value	Unit
TPS_Lens1_Act	15.000	V
TPS_Lens2_Act	30.000	V
TPS_Lens3_Act	20.000	V
TPS_Lens4_Act	60.000	V
TPS_Lens5_Act	69.000	V

Calculated

Trace	Value	Unit
NO+	0.4003	%
O ₂ +	4.775	%
H ₃ O+(H ₂ O)	1.090	%
PI	7.130E+7	ncps
H ₃ O+	93.73	%

Corrected H₃O+ Calc Traces.iCT

Peaks and Traces

Acquisition ACQ active

Single Spec Time (ms) 1000
Extraction time (μs) 4.0 372.0 amu
max Flighttime(μs) 32.0 31.25 kHz

Data Save Settings

Spec Trace Raw
Time Duration
02:00:00 Single File Duration
24 Number of Files To Store
C:\lonicon\data

Add File Count Extension
 New ACQ for new file
<year>_<month>_<day>\
Data_<hour>_<minute>_<second>
2023_07_29\Data_19_36_43_part_XXX

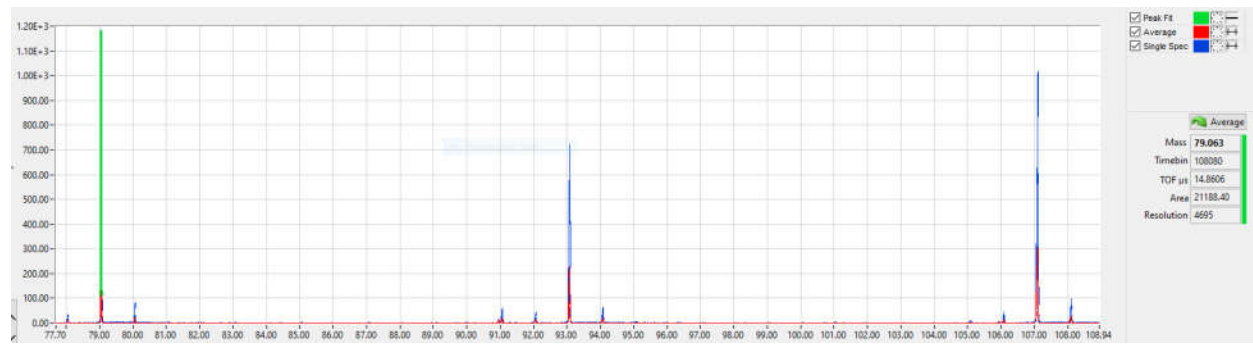
Mass Axis Calibration

Cal 60 sec
Mass TimeBin

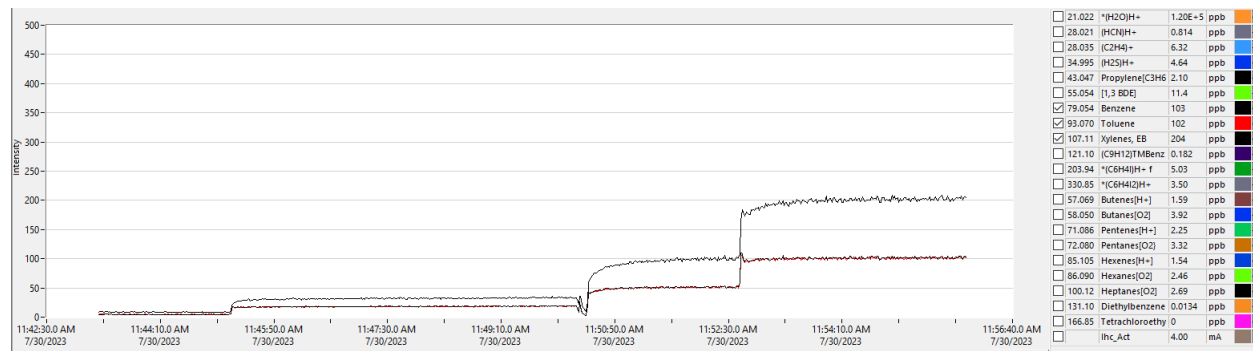
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330.8500	247759	<input type="button" value="Delete"/>	<input type="button" value="Up"/>	-25496.5	
59.0491	89943	<input type="button" value="Delete"/>	<input type="button" value="Down"/>		

Acquisition Settings

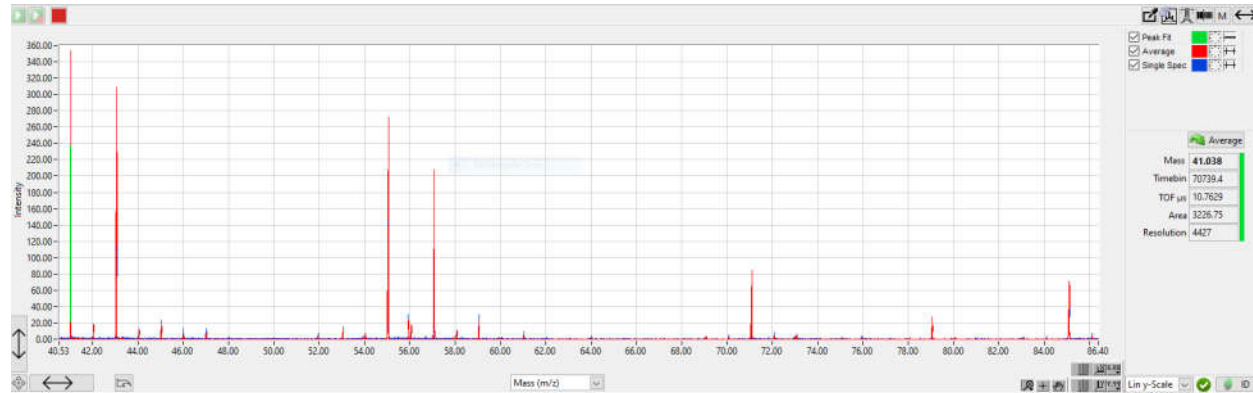
CCND Mobile Monitoring Van
2023 Q3



BTEX Peaks

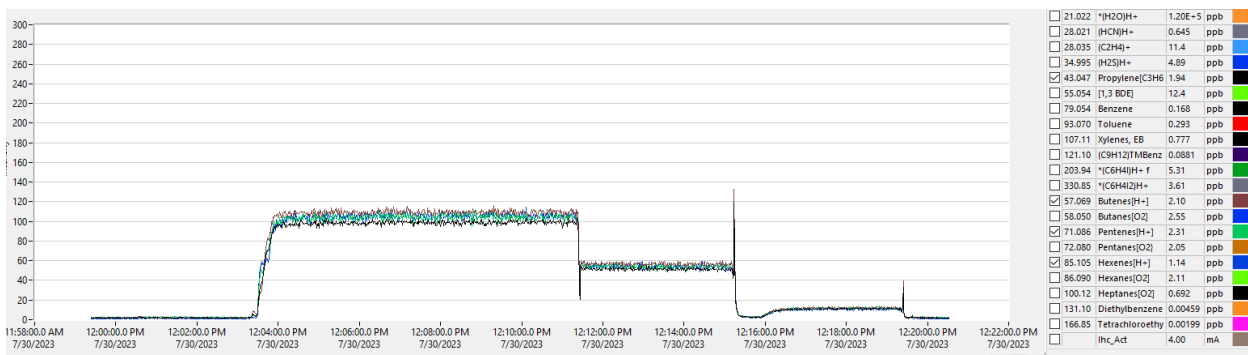


Calibration BTEX

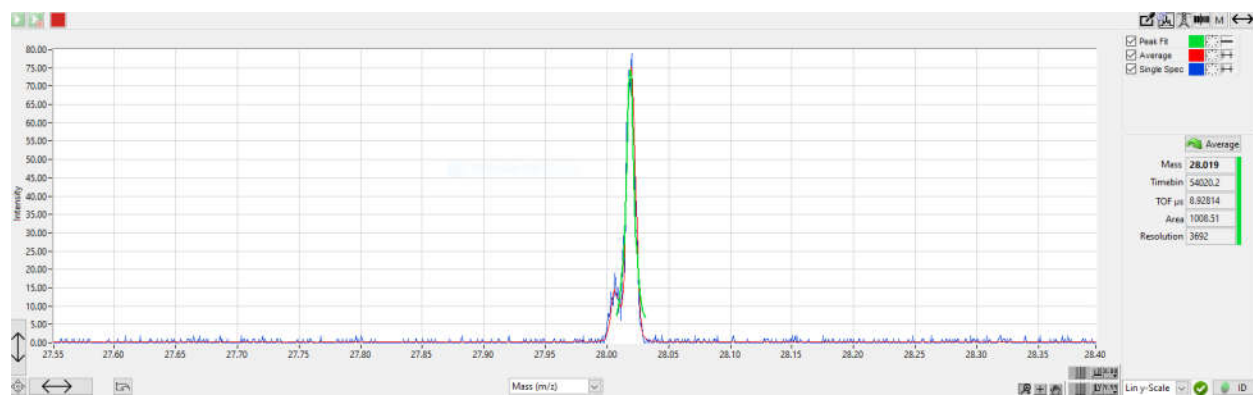


Alkenes Peaks

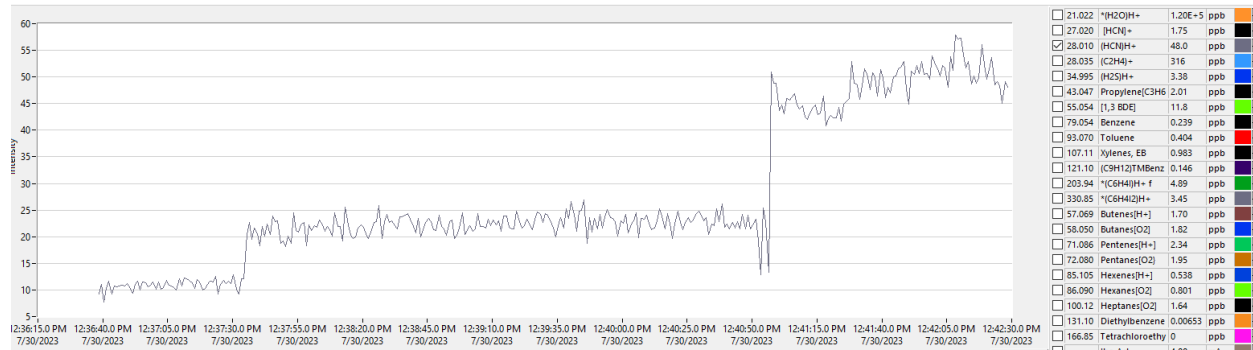
CCND Mobile Monitoring Van 2023 Q3



Calibration Alkenes



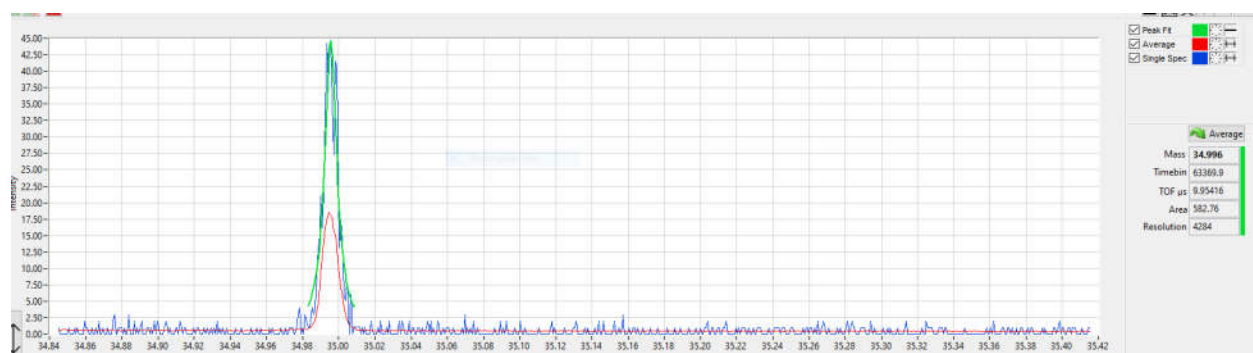
HCN Peak



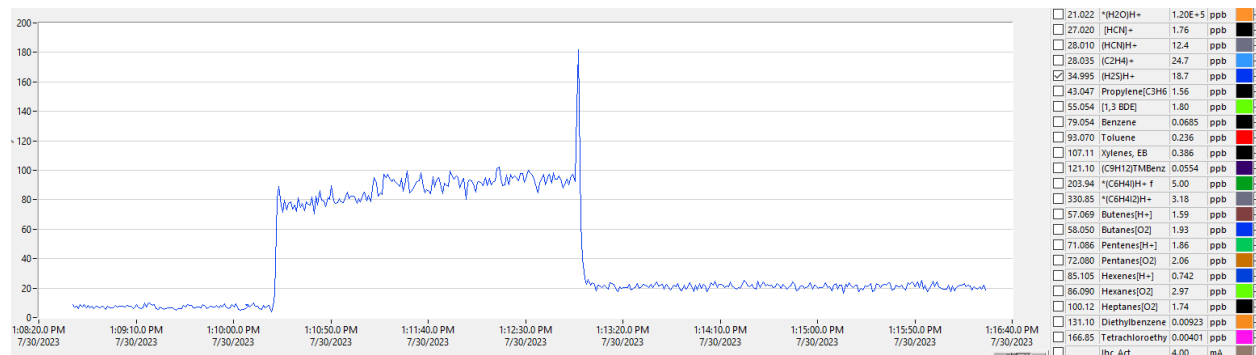
HCN Calibrations

CCND Mobile Monitoring Van

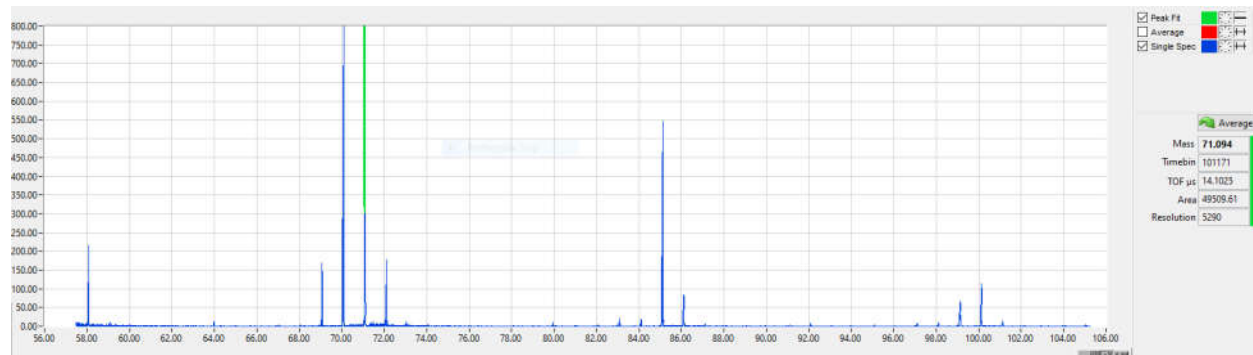
2023 Q3



H₂S Peak

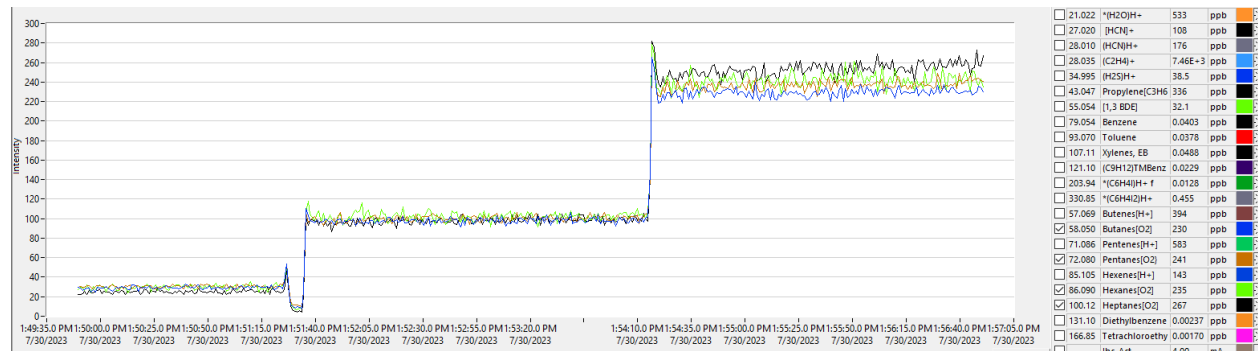


H₂S Calibrations



Alkane Peaks

CCND Mobile Monitoring Van
2023 Q3



Alkane Calibration

CCND Mobile Monitoring Van
2023 Q3

7-31-23 CCND Neighborhood Monitoring
Pioneer Park
PTR Screen Shots

The screenshot shows a software interface for a PTR mass spectrometer. At the top, there are icons for file operations (New, Open, Save, Print) and system controls (Stop, Start). Below this is a parameter configuration section with dropdown menus for 'Setting' (Odor), 'Primary Ion' (H₃O⁺), and 'Transmission' (DC). The main area displays various operational parameters in a grid format:

	Man/Ctrl	Ctrl
PC	344.3	344.44 mbar
p Drift	2.30	2.29 mbar
TofLens		4.64E-5 mbar
TOF		1.32E-6 mbar
E/N		120 Td
Temps	80.00 °C	80.20 °C
SrcValve	50.0	
H ₂ O	6.0	5.99 sccm
O ₂	0.0	0.00 sccm
NO	0.0	0.00 sccm
Ihc	4	4.0 mA
	On/Off	On
FCinlet	60.0	59.97 sccm
U	FU °C	D+ D-
Us	150	145.0 V
Uso	80	78.6 V
Udrift	525	526.1 V

Production settings

TPS 4-6-23 MCP Tune.iTPS *Changed*

Lens 1	14.0	14.0 V	All on <input checked="" type="checkbox"/>
Lens 2	30.0	30.0 V	Lenses <input checked="" type="checkbox"/>
Lens 3	20.0	20.0 V	
Lens 4	60.0	60.0 V	
Lens 5	70.0	69.0 V	
Lens 6	80.0	80.0 V	
Lens 7	17.0	17.0 V	
Push L	16.5	16.0 V	<input checked="" type="checkbox"/> 3 mA
Push H	790.0	790.0 V	<input checked="" type="checkbox"/> 3 mA
Pull L	80.0	80.0 V	<input checked="" type="checkbox"/> 3 mA
Pull H	680.0	680.0 V	<input checked="" type="checkbox"/> 3 mA
Grid	2400.0	2283.0 V	<input checked="" type="checkbox"/> 1 µA
Cage	5020.0	4766 V	<input checked="" type="checkbox"/> 99 µA
Refl. Grid	667.0	634.0 V	<input checked="" type="checkbox"/> 75 µA
Refl. Back	900.0	855.0 V	<input checked="" type="checkbox"/> 167 µA
MCP F	5400	5134 V	<input checked="" type="checkbox"/> 17 µA
MCP B	2496	2379 V	<input checked="" type="checkbox"/> 216 µA

TOF Lenses

CCND Mobile Monitoring Van
2023 Q3

Defined Peaks

	Mass	Value	Unit
✓ Pentanes[O2]	72.08000	4.40	ppb
Pentanes[H+]	73.16000	0.03	ppb
Hexenes[O2]	84.16000	0.01	ppb
✓ Hexenes[H+]	85.10500	3.32	ppb
✓ Hexanes[O2]	86.09000	8.64	ppb
Hexanes[H+]	87.11680	0.21	ppb
✓ Heptanes[O2]	100.12000	5.87	ppb
Heptanes[H+]	100.91130	0.01	ppb
Dimethylcyclohe	112.21000	0.01	ppb
Octanes[O2]	114.23000	0.01	ppb
Octanes[H+]	115.12000	0.12	ppb

21 of 239 Peaks selected from
"2-5-23 Suncor Working Peak Table.ipta"

Instrument

TOFSupply

Description	Value	Unit
TPS_Lens1_Act	14.000	V
TPS_Lens2_Act	30.000	V
TPS_Lens3_Act	20.000	V
TPS_Lens4_Act	60.000	V
TPS_Lens5_Act	69.000	V

Calculated

Trace	Value	Unit
NO+	0.7041	%
O2+	3.530	%
H3O+(H2O)	3.316	%
PI	6.776E+7	ncps
H3O+	92.45	%

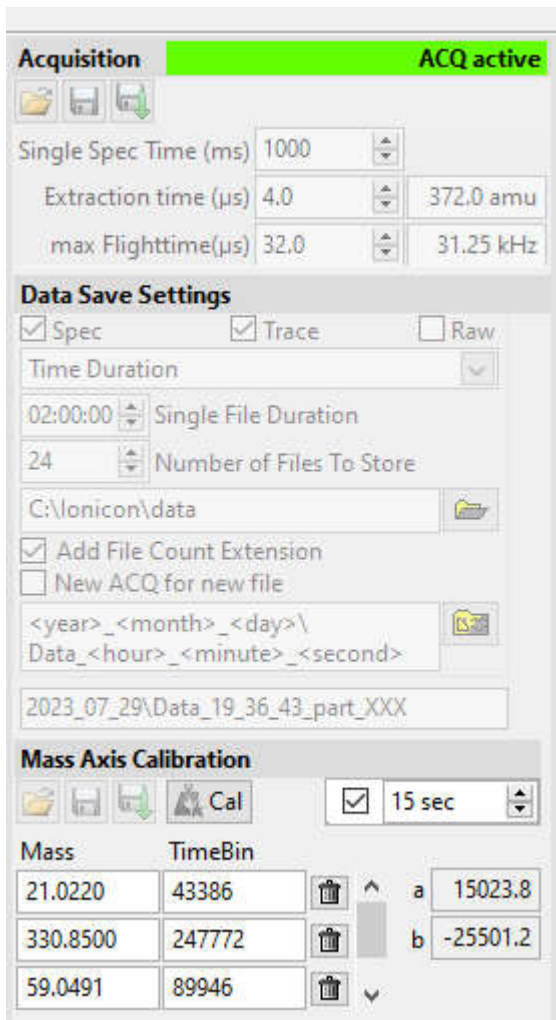
Corrected H3O+ Calc Traces.iCT

Hex1

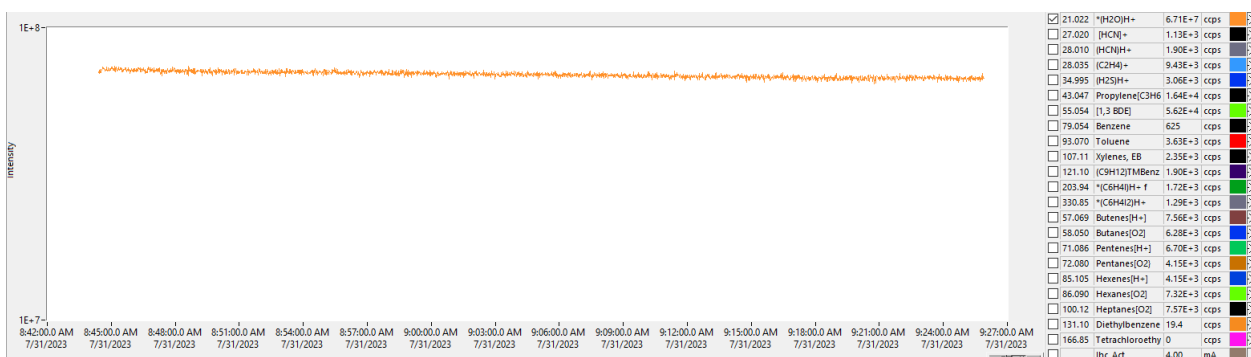
OFF/ON	OP
<input checked="" type="checkbox"/>	ON
Frequency	6.00
Amplitude	95.0
Offset	-0.70

Peaks and Traces, Hexapole Settings

CCND Mobile Monitoring Van
2023 Q3

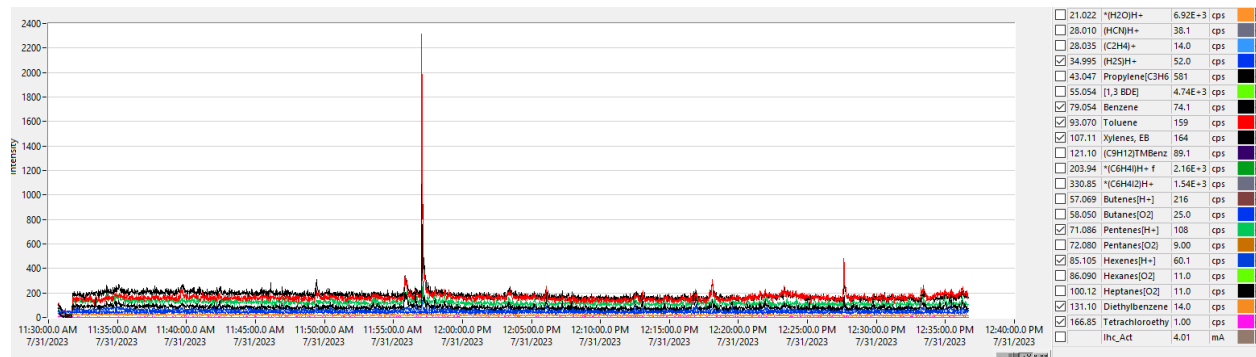


Acquisition Settings



Hydronium Stability

CCND Mobile Monitoring Van 2023 Q3



Pioneer Park Raw Data

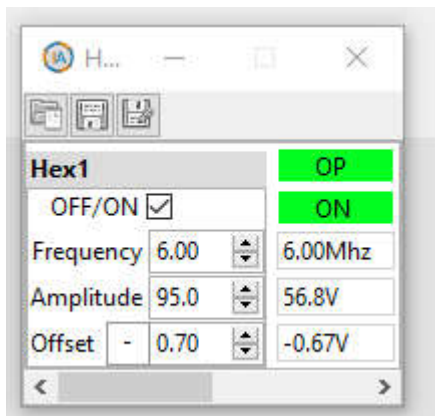
CCND Mobile Monitoring Van
2023 Q3

8/1/23 CCND Afternoon/Evening Monitoring
Globeville and Swansea
PTR Screen Shots

The screenshot shows a software interface for a PTR mass spectrometer. At the top, there are icons for file operations (New, Open, Save, Print) and a search function. Below this is a section for 'Setting' (Odor), 'Primary Ion' (H₃O⁺), and 'Transmission' (DC). The main area displays various parameters with their current values and control buttons:

	Man/Ctrl	Ctrl
PC	343.9	343.89 mbar
p Drift	2.30	2.29 mbar
TofLens		4.72E-5 mbar
TOF		1.12E-6 mbar
E/N		120 Td
Temps	80.00 °C	80.10 °C
SrcValve	50.0	
H ₂ O	6.0	6.00 sccm
O ₂	0.0	0.00 sccm
NO	0.0	0.00 sccm
Ihc	4	4.0 mA
	On/Off	On
FCinlet	60.0	60.03 sccm
U	FU	°C
Us	150	145.0 V
Uso	80	78.6 V
Udrift	525	526.1 V

Production Settings



Hexapole Settings

TPS *Changed*			
Lens 1	14.0	14.0 V	All on <input checked="" type="checkbox"/>
Lens 2	30.0	30.0 V	Lenses <input checked="" type="checkbox"/>
Lens 3	20.0	20.0 V	
Lens 4	60.0	60.0 V	
Lens 5	70.0	69.0 V	
Lens 6	80.0	80.0 V	
Lens 7	17.0	18.0 V	
Push L	16.5	16.0 V	<input checked="" type="checkbox"/> 3 mA
Push H	790.0	790.0 V	<input checked="" type="checkbox"/> 2 mA
Pull L	80.0	80.0 V	<input checked="" type="checkbox"/> 3 mA
Pull H	680.0	680.0 V	<input checked="" type="checkbox"/> 3 mA
Grid	2400.0	2282.0 V	<input checked="" type="checkbox"/> 1 µA
Cage	5020.0	4766 V	<input checked="" type="checkbox"/> 99 µA
Refl. Grid	667.0	634.0 V	<input checked="" type="checkbox"/> 75 µA
Refl. Back	900.0	855.0 V	<input checked="" type="checkbox"/> 167 µA
MCP F	5400	5134 V	<input checked="" type="checkbox"/> 17 µA
MCP B	2496	2378 V	<input checked="" type="checkbox"/> 215 µA

TOF Lenses Voltages

CCND Mobile Monitoring Van
2023 Q3

Defined Peaks

	Mass	Value	Unit
*	(H ₂ O)+	18.01000	1.23E+4 ccps
*	(H ₃ N)H+	18.03380	1.98E+5 ccps
*	(H ₂ O)H+	19.01780	5.19E+3 ccps
✓	(H ₂ O)H+	21.02210	6.53E+7 ccps
	[HCN]H+	27.02000	1.41E+3 ccps
	*(N ₂)H+	28.00600	0.00 ccps
✓	(HCN)H+	28.01000	1.75E+3 ccps
✓	(C ₂ H ₄)H+	28.03508	1.11E+4 ccps
	*(N ₂)H+	29.01340	2.65E+4 ccps
	Ethylene[C ₂ H ₄]	29.04400	570.68 ccps
	*(NO)H+ [NO+]	29.99740	8.94E+4 ccps

21 of 239 Peaks selected from
"2-5-23 Suncor Working Peak Table.upta"

Instrume

Description	Value	Unit
TOFSupply		
TPS_Lens1_Act	14.000	V
TPS_Lens2_Act	30.000	V
TPS_Lens3_Act	20.000	V
TPS_Lens4_Act	60.000	V
TPS_Lens5_Act	70.000	V

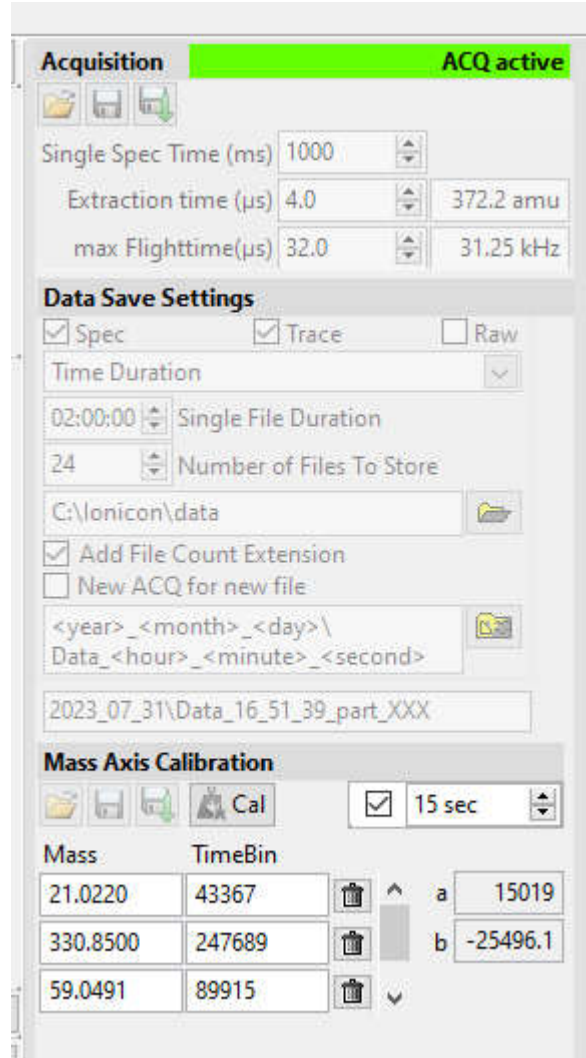
Calculated

Trace	Value	Unit
NO+	2.010	%
O ₂ +	3.567	%
H ₃ O+(H ₂ O)	5.058	%
PI	7.303E+7	ncps
H ₃ O+	89.36	%

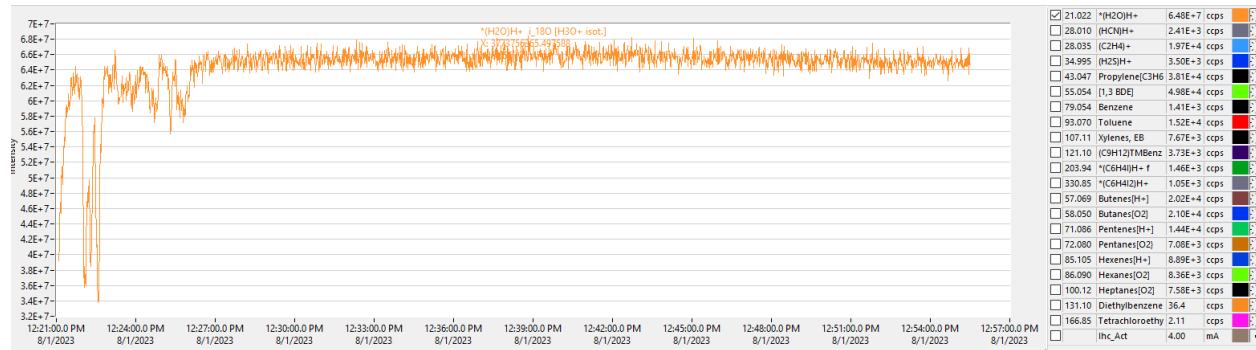
Corrected H₃O+ Calc Traces.iCT

Peak and Traces

CCND Mobile Monitoring Van
2023 Q3

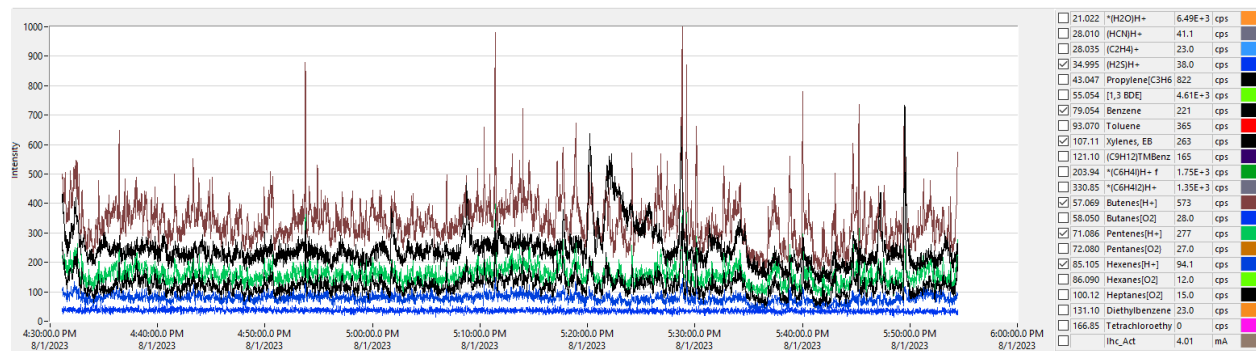


Acquisition Settings



Hydronium Stability

CCND Mobile Monitoring Van 2023 Q3



Globeville Raw Data

CCND Mobile Monitoring Van
2023 Q3

8/2/23 CCND Afternoon/Evening Monitoring

PTR Settings

Dupont

Acquisition Idle

Single Spec Time (ms)
Extraction time (μ s) 372.1 amu
max Flighttime(μ s) 31.25 kHz

Data Save Settings

Spec Trace Raw

Time Duration

02:00:00 Single File Duration

24 Number of Files To Store

C:\lonicon\data

Add File Count Extension
 New ACQ for new file

<year>_<month>_<day>\
Data_<hour>_<minute>_<second>

2023_07_31\Data_16_51_39_part_XXX

Mass Axis Calibration

Cal 60 sec

Mass	TimeBin		
21.0220	43375	<input type="button"/>	a 15020.8
330.8500	247719	<input type="button"/>	b -25497.5
59.0491	89926	<input type="button"/>	

Acquisition Settings

CCND Mobile Monitoring Van
2023 Q3

Defined Peaks

Mass Value Unit

(C4H8S)H+	89.04190	0.04	ppb
(C4H8O2)H+	89.05970	0.03	ppb
(C5H12O)H+	89.09610	0.03	ppb
*(FeH2O2)H+	90.94770	0.34	ppb
(C4H10S)H+	91.05760	0.03	ppb
(C3H10OSi)H+	91.05800	0.03	ppb
(C4H10O2)H+	91.07540	0.03	ppb
(C3H10OSi)H+ i	92.05900	0.01	ppb
<input checked="" type="checkbox"/> Toluene	93.06990	0.28	ppb
(C2H6S2)H+	94.99840	0.02	ppb
(C6H6O)H+	95.04910	0.05	ppb

21 of 239 Peaks selected from
"2-5-23 Suncor Working Peak Table.ipta"

Instrument

TOFSupply

Description Value Unit

TPS_Lens1_Act	15.000	V
TPS_Lens2_Act	31.000	V
TPS_Lens3_Act	21.000	V
TPS_Lens4_Act	61.000	V
TPS_Lens5_Act	70.000	V

Calculated

Trace Value Unit

NO+	0.2520	%
O2+	4.076	%
H3O+(H2O)	0.3490	%
PI	6.102E+7	ncps
H3O+	95.32	%

Corrected H3O+ Calc Traces.iCT

Peaks & Traces

CCND Mobile Monitoring Van
2023 Q3

TPS *Changed*			
Lens 1	14.0	15.0 V	All on <input checked="" type="checkbox"/>
Lens 2	30.0	31.0 V	Lenses <input checked="" type="checkbox"/>
Lens 3	20.0	21.0 V	
Lens 4	60.0	61.0 V	
Lens 5	70.0	70.0 V	
Lens 6	80.0	80.0 V	
Lens 7	17.0	18.0 V	
Push L	16.5	16.0 V	<input checked="" type="checkbox"/> 0 mA
Push H	790.0	790.0 V	<input checked="" type="checkbox"/> 0 mA
Pull L	80.0	80.0 V	<input checked="" type="checkbox"/> 0 mA
Pull H	680.0	680.0 V	<input checked="" type="checkbox"/> 0 mA
Grid	2400.0	2283.0 V	<input checked="" type="checkbox"/> 1 µA
Cage	5020.0	4768 V	<input checked="" type="checkbox"/> 100 µA
Refl. Grid	667.0	634.0 V	<input checked="" type="checkbox"/> 77 µA
Refl. Back	900.0	855.0 V	<input checked="" type="checkbox"/> 169 µA
MCP F	5400	5136 V	<input checked="" type="checkbox"/> 17 µA
MCP B	2496	2379 V	<input checked="" type="checkbox"/> 211 µA

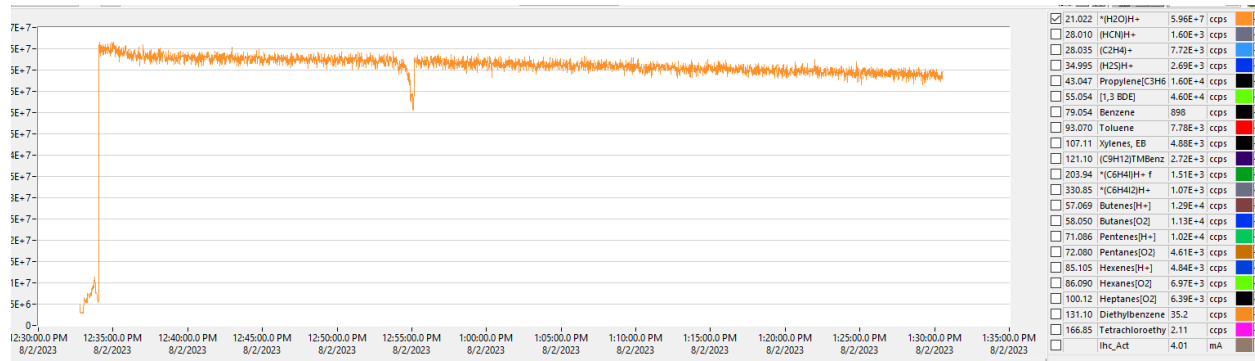
TOF Lenses Voltages

Hex1

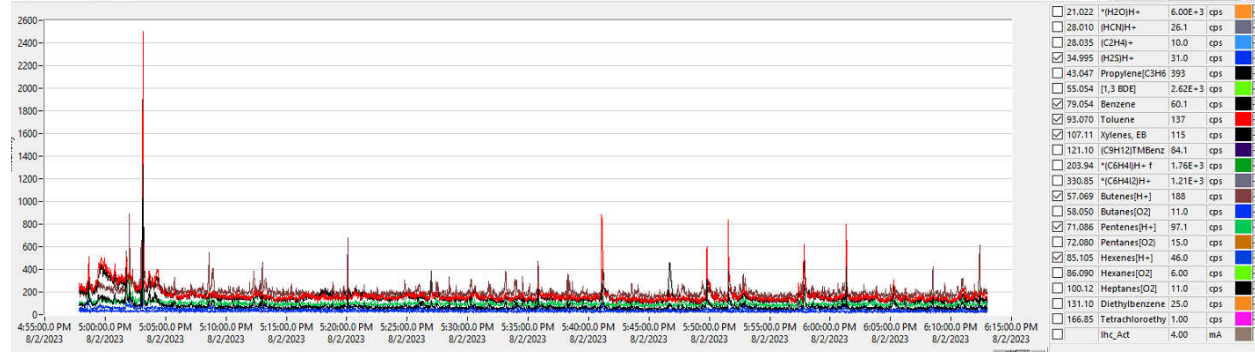
OFF/ON <input checked="" type="checkbox"/>	OP
Frequency 6.00	6.00Mhz
Amplitude 95.0	56.8V
Offset - 0.70	-0.67V

Hexapole Settings

CCND Mobile Monitoring Van
2023 Q3



Hydronium Stability Check



Dupont Raw Data

CCND Mobile Monitoring Van
2023 Q3

8/3/23 CCND Monitoring
Western Hills
PTR Settings

Setting	Odor			
Primary Ion	H ₃ O ⁺			
Transmission	DC			
Man/Ctrl Ctrl				
PC	343.3	343.32 mbar		
p Drift	2.30	2.29 mbar		
TofLens		4.58E-5 mbar		
TOF		9.84E-7 mbar		
E/N		120 Td		
Temps	80.10 °C	80.10 °C		
SrcValve	50.0			
H ₂ O	6.0	6.00 sccm		
O ₂	0.0	0.00 sccm		
NO	0.0	0.00 sccm		
Ihc	4	4.0 mA		
	On/Off	On		
FCinlet	60.0	59.97 sccm		
U	FU	°C	D \rightarrow	D \leftarrow
Us	150	145.0 V		
Uso	80	78.6 V		
Udrift	525	526.1 V		

Production Settings

TPS *Changed*

MCP TOP

Lens 1	14.0	1.0 V	All on	<input type="checkbox"/>
Lens 2	30.0	1.0 V	Lenses	<input type="checkbox"/>
Lens 3	20.0	1.0 V		
Lens 4	60.0	1.0 V		
Lens 5	70.0	1.0 V		
Lens 6	80.0	1.0 V		
Lens 7	17.0	1.0 V		
Push L	16.5	0.0 V	<input type="checkbox"/>	0 mA
Push H	790.0	1.0 V	<input type="checkbox"/>	0 mA
Pull L	80.0	0.0 V	<input type="checkbox"/>	0 mA
Pull H	680.0	3.0 V	<input type="checkbox"/>	0 mA
Grid	2400.0	11.0 V	<input type="checkbox"/>	1 µA
Cage	5020.0	10 V	<input type="checkbox"/>	0 µA
Refl. Grid	667.0	3.0 V	<input type="checkbox"/>	3 µA
Refl. Back	900.0	2.0 V	<input type="checkbox"/>	5 µA
MCP F	5400	26 V	<input type="checkbox"/>	2 µA
MCP B	2496	21 V	<input type="checkbox"/>	3 µA

Lens Settings

CCND Mobile Monitoring Van
2023 Q3

Defined Peaks

Mass Value Unit

(C7H14)H+	99.11680	0.03	ppb
(C5H8O2)H+	101.05970	0.02	ppb
(C6H12O)H+	101.09610	0.02	ppb
(C7H16)H+	101.13240	0.02	ppb
(C6H15N)H+	102.12770	0.01	ppb
(C6H14O)H+	103.11170	0.02	ppb
Styrene	105.06990	0.04	ppb
*(HO3Fe)H+	105.93480	0.06	ppb
(C7H6O)H+	107.04910	0.13	ppb
✓ Xylenes, EB	107.10550	0.56	ppb
(C6H4O2)H+ [p-]	109.02840	0.06	ppb

21 of 239 Peaks selected from
"2-5-23 Suncor Working Peak Table.ipta"

Instrument

TOFSupply

Description Value Unit

TPS_Lens1_Act	1.000	V
TPS_Lens2_Act	1.000	V
TPS_Lens3_Act	1.000	V
TPS_Lens4_Act	1.000	V
TPS_Lens5_Act	1.000	V

Calculated

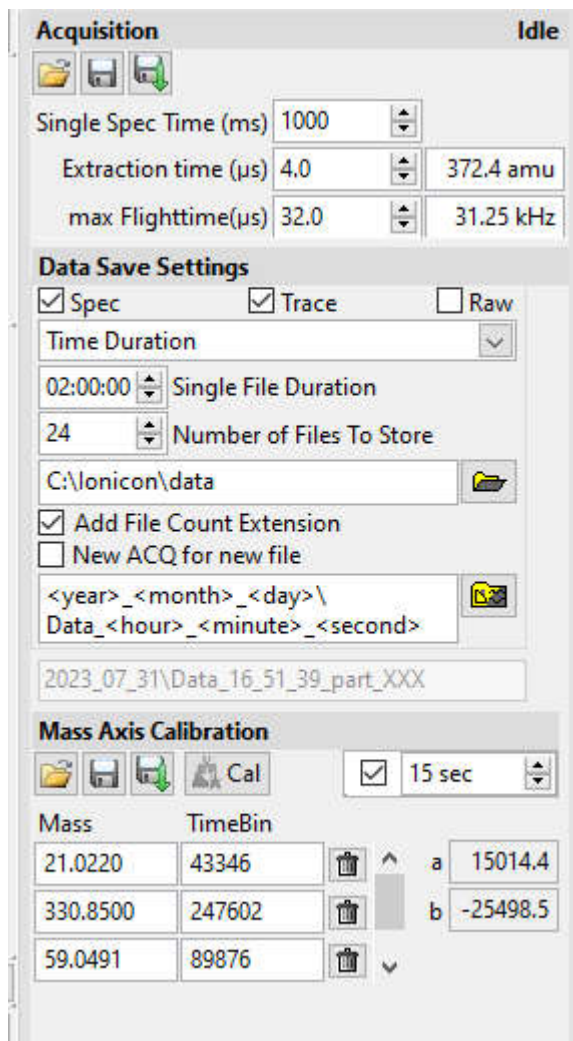
Trace Value Unit

NO+	0.3168	%
O2+	3.734	%
H3O+(H2O)	0.3726	%
PI	5.325E+7	ncps
H3O+	95.58	%

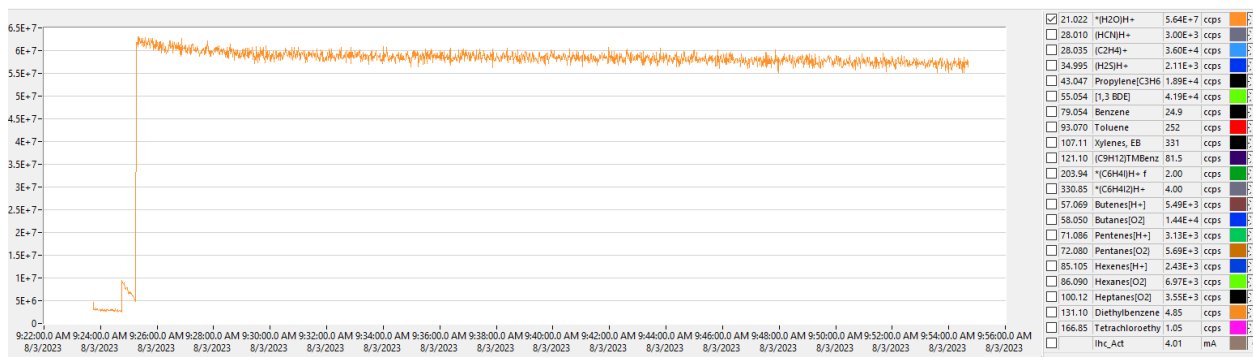
Corrected H3O+ Calc Traces.iCT

Peaks & Traces

CCND Mobile Monitoring Van
2023 Q3

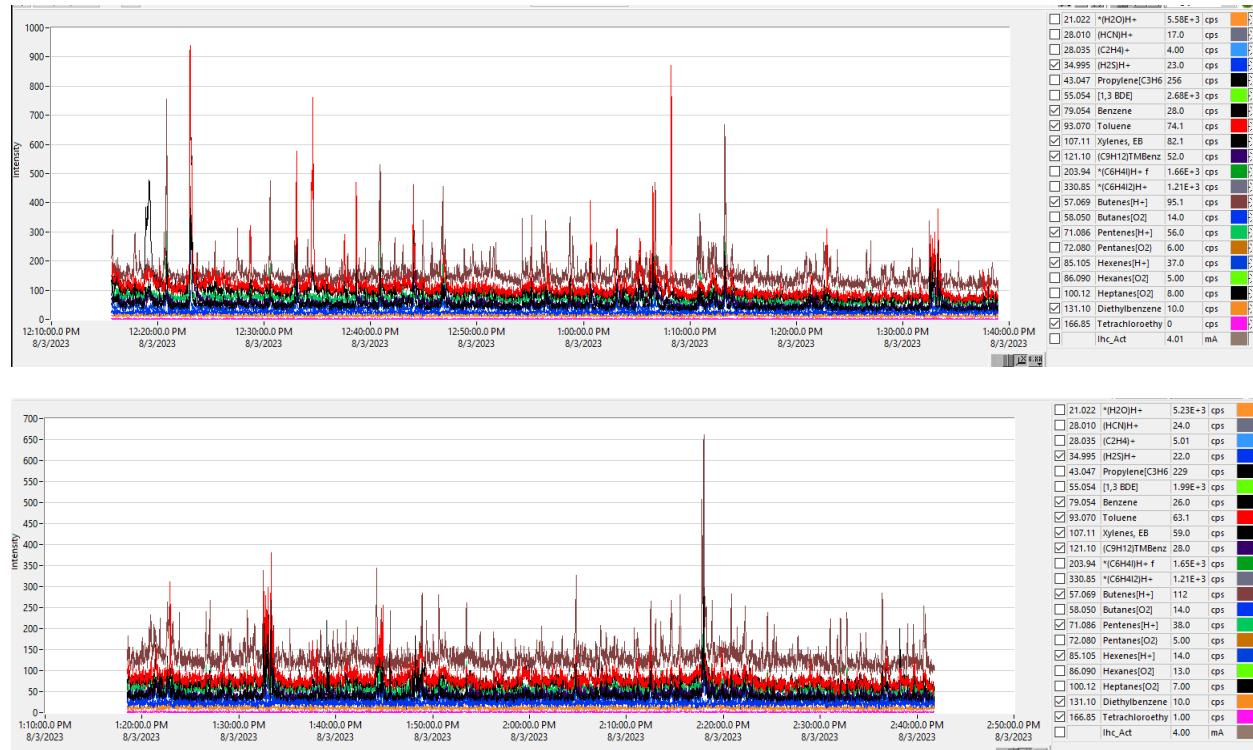


Acquisition Settings



Hydronium Settings

CCND Mobile Monitoring Van 2023 Q3



Western Hills Raw Data

8-4-23 CCND Monitoring
Adams City

CCND Mobile Monitoring Van
2023 Q3

PTR Screenshots

Setting	Odor			
Primary Ion	H ₃ O ⁺			
Transmission	DC			
	Man/Ctrl	Ctrl		
PC	344.6	344.60 mbar		
p Drift	2.30	2.30 mbar		
TofLens		4.72E-5 mbar		
TOF		9.49E-7 mbar		
E/N		120 Td		
Temps	80.10 °C	79.90 °C		
SrcValve	50.0			
H ₂ O	6.0	6.00 sccm		
O ₂	0.0	0.00 sccm		
NO	0.0	0.00 sccm		
Ihc	4	4.0 mA		
	On/Off	On		
FCinlet	60.0	60.00 sccm		
U	FU	°C		
Us	150		145.0 V	
Uso	80		78.6 V	
Udrift	525		526.1 V	

Production Settings

CCND Mobile Monitoring Van
2023 Q3

TPS		*Changed*		
Lens 1	14.0	14.0 V	All on	<input checked="" type="checkbox"/>
Lens 2	30.0	30.0 V	Lenses	<input checked="" type="checkbox"/>
Lens 3	20.0	20.0 V		
Lens 4	60.0	60.0 V		
Lens 5	70.0	70.0 V		
Lens 6	80.0	80.0 V		
Lens 7	17.0	18.0 V		
Push L	16.5	16.0 V	<input checked="" type="checkbox"/>	3 mA
Push H	790.0	790.0 V	<input checked="" type="checkbox"/>	2 mA
Pull L	80.0	80.0 V	<input checked="" type="checkbox"/>	3 mA
Pull H	680.0	680.0 V	<input checked="" type="checkbox"/>	3 mA
Grid	2400.0	2282.0 V	<input checked="" type="checkbox"/>	1 µA
Cage	5020.0	4766 V	<input checked="" type="checkbox"/>	99 µA
Refl. Grid	667.0	634.0 V	<input checked="" type="checkbox"/>	75 µA
Refl. Back	900.0	855.0 V	<input checked="" type="checkbox"/>	167 µA
MCP F	5400	5134 V	<input checked="" type="checkbox"/>	17 µA
MCP B	2496	2379 V	<input checked="" type="checkbox"/>	216 µA

TOF Lenses

CCND Mobile Monitoring Van
2023 Q3

Defined Peaks

	Mass	Value	Unit
(C7H14)H+	99.11680	0.08	ppb
(C5H8O2)H+	101.05970	0.18	ppb
(C6H12O)H+	101.09610	0.17	ppb
(C7H16)H+	101.13240	0.15	ppb
(C6H15N)H+	102.12770	0.02	ppb
(C6H14O)H+	103.11170	0.04	ppb
Styrene	105.06990	0.12	ppb
*(HO3Fe)H+	105.93480	0.02	ppb
(C7H6O)H+	107.04910	0.47	ppb
✓ Xylenes, EB	107.10550	1.92	ppb
(C6H4O2)H+ [p-]	109.02840	0.17	ppb

21 of 239 Peaks selected from
"2-5-23 Suncor Working Peak Table.ipta"

Instrument

Description	Value	Unit
TPS_Lens1_Act	14.000	V
TPS_Lens2_Act	30.000	V
TPS_Lens3_Act	20.000	V
TPS_Lens4_Act	60.000	V
TPS_Lens5_Act	70.000	V

Calculated

Trace	Value	Unit
NO+	0.5549	%
O2+	2.387	%
H3O+(H2O)	2.936	%
PI	5.817E+7	ncps
H3O+	94.12	%

Corrected H3O+ Calc Traces.iCT

Peaks and Traces

CCND Mobile Monitoring Van
2023 Q3

Acquisition ACQ active

Single Spec Time (ms)	1000.	▲ ▼
Extraction time (μs)	4.0.	▲ ▼
max Flighttime(μs)	32.0	▲ ▼
	372.6 amu	
	31.25 kHz	

Data Save Settings

Spec Trace Raw

Time Duration:

02:00:00 Single File Duration

24 Number of Files To Store

C:\Ionicon\data

Add File Count Extension
 New ACQ for new file

<year>_<month>_<day>\
Data_<hour>_<minute>_<second>

2023_07_31\Data_16_51_39_part_XXX

Mass Axis Calibration

15 sec

Mass	TimeBin	<input type="button" value="刪"/>	<input type="button" value="▲"/>	a	15012.2
21.0220	43334	<input type="button" value="刪"/>	<input type="button" value="▼"/>	b	-25499.6
330.8500	247562	<input type="button" value="刪"/>			
59.0491	89858	<input type="button" value="刪"/>	<input type="button" value="▼"/>		

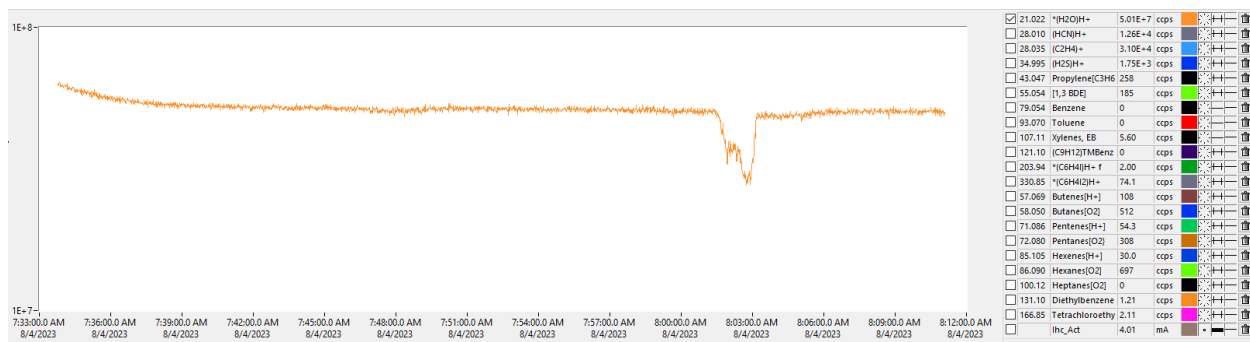
Acquisitions

H...

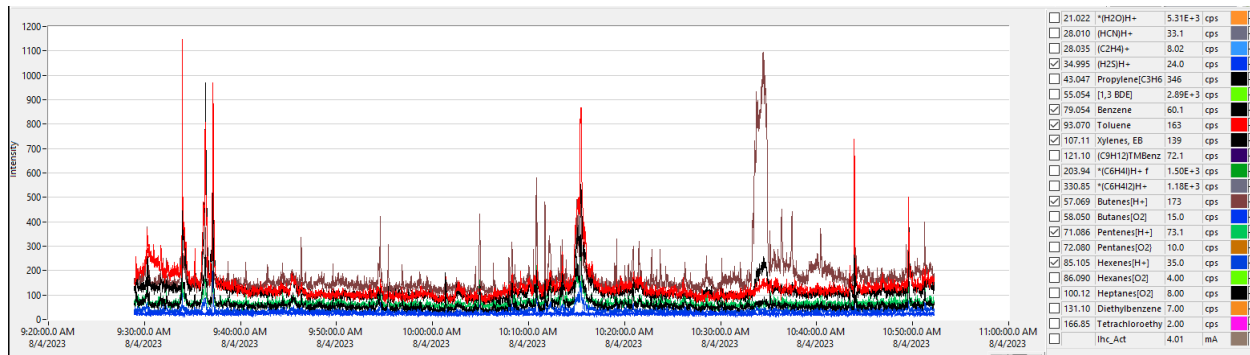
Hex1		OP
OFF/ON	<input checked="" type="checkbox"/>	ON
Frequency	6.00	6.00Mhz
Amplitude	95.0	57.1V
Offset	-0.70	-0.67V

Hexapole Settings

CCND Mobile Monitoring Van 2023 Q3



Hydronium Stability



Adams City Raw Data

CCND Mobile Monitoring Van
2023 Q3

PTR Daily Calibration Checks

Date	Time	Initial Instrument Calibration				
		Calibration Gas Component	Calibration Value (ppb v)	Response (ppb v)	Difference (% of value)	Pass/Fail
7/30/2023	11:55	Benzene	100	99.7	-0.3	Pass
		Toluene	100	99.2	-0.8	Pass
		Xylenes	200	202	1.0	Pass
	11:51	Benzene	50	51.4	2.8	Pass
		Toluene	50	50.1	0.2	Pass
		Xylenes	100	101	1.0	Pass
	11:48	Benzene	20	19.4	-3.0	Pass
		Toluene	20	19.2	-4.0	Pass
		Xylenes	40	34	-15.0	Pass
	11:43	Benzene	5	4.84	-3.2	Pass
		Toluene	5	4.94	-1.2	Pass
		Xylenes	10	9.1	-9.0	Pass
12:10	12:10	Ethylene	100	104	4.0	Pass
		Propylene	100	99.2	-0.8	Pass
		1-Butene	100	110	10.0	Pass
		1-Pentene	100	108	8.0	Pass
		1-Hexene	100	104	4.0	Pass
		1,3-Butadiene	100	113	13.0	Pass
	12:14	Ethylene	50	50.9	1.8	Pass
		Propylene	50	50.6	1.2	Pass
		1-Butene	50	55.8	11.6	Pass
		1-Pentene	50	55.9	11.8	Pass
		1-Hexene	50	55.5	11.0	Pass
		1,3-Butadiene	50	58.1	16.2	Pass
	12:16	Ethylene	10	11.5	15.0	Pass
		Propylene	10	10.9	9.0	Pass
		1-Butene	10	11.3	13.0	Pass
		1-Pentene	10	11.4	14.0	Pass
		1-Hexene	10	11.3	13.0	Pass
		1,3-Butadiene	10	10.3	3	Pass
12:37	12:37	HCN	50	50.4	0.8	Pass
	12:40	HCN	25	24.1	-3.6	Pass
		HCN	10	10.6	6.0	Pass
	13:12	H ₂ S	100	97	-3.0	Pass
	13:15	H ₂ S	20	20.2	1.0	Pass
	13:09	H ₂ S	5	5.8	16.0	Pass
	13:55	Butane	250	232	-7.2	Pass
		Pentane	250	236	-5.6	Pass
		Hexane	250	244	-2.4	Pass
		Heptane	250	248	-0.8	Pass
13:53	13:53	Butane	100	100	0.0	Pass
		Pentane	100	106	6.0	Pass
		Hexane	100	103	3.0	Pass
		Heptane	100	102	2.0	Pass
13:50	13:50	Butane	25	27.1	8.4	Pass
		Pentane	25	28	12.0	Pass
		Hexane	25	25.2	0.8	Pass
		Heptane	25	26.5	6.0	Pass

CCND Mobile Monitoring Van
2023 Q3

Instrument Calibration Check						
Date	Time	Calibration Gas Component	Calibration Value (ppb v)	Response (ppb v)	Difference (% of value)	Pass/Fail
7/31/2023 Pioneer Park	9:53	Ethylene	50	52.3	4.6	Pass
		Propylene	50	51.5	3.0	Pass
		1-Butene	50	54.1	8.2	Pass
		1-Pentene	50	52	4.0	Pass
		1-Hexene	50	53.4	6.8	Pass
		1,3-Butadiene	50	48.6	-2.8	Pass
	9:56	Benzene	100	105	5.0	Pass
		Toluene	100	99.8	-0.2	Pass
		Xylenes	200	197	-1.5	Pass
	10:02	Benzene	20	19.6	-2.0	Pass
		Toluene	20	19.1	-4.5	Pass
		Xylenes	40	36	-10.0	Pass
.						
	10:08	HCN	25	24.9	-0.4	Pass
	10:14	H ₂ S	100	96.9	-3.1	Pass
		H ₂ S	20	19.9	-0.5	Pass
	10:16	Butane	150	151	0.7	Pass
		Pentane	150	146	-2.7	Pass
		Hexane	150	144	-4.0	Pass
		Heptane	150	147	-2.0	Pass
.						
14:19	HCN	25	24.8	-0.8	Pass	
16:25	H ₂ S	20	18.9	-5.5	Pass	
16:14	Butane	150	160	6.7	Pass	
	Pentane	150	145	-3.3	Pass	
	Hexane	150	145	-3.3	Pass	
	Heptane	150	142	-5.3	Pass	
	16:38	Benzene	20	20.2	1.0	Pass
		Toluene	20	19.5	-2.5	Pass
		Xylenes	40	36.7	-8.2	Pass
	16:27	Ethylene	50	51.9	3.8	Pass
		Propylene	50	48.1	-3.8	Pass
		1-Butene	50	53	6.0	Pass
		1-Pentene	50	50.6	1.2	Pass
		1-Hexene	50	49.5	-1.0	Pass
		1,3-Butadiene	50	53.1	6.2	Pass

CCND Mobile Monitoring Van
2023 Q3

Date	Time	Calibration Gas Component	Instrument Calibration Check			
			Calibration Value (ppb v)	Response (ppb v)	Difference (% of value)	Pass/Fail
8/1/2023 Globeville Swansea	14:26	Ethylene	50	51.8	3.6	Pass
		Propylene	50	51.1	2.2	Pass
		1-Butene	50	55.4	10.8	Pass
		1-Pentene	50	51.6	3.2	Pass
		1-Hexene	50	51.4	2.8	Pass
		1,3-Butadiene	50	54.1	8.2	Pass
	14:19	Benzene	100	110	10.0	Pass
		Toluene	100	106	6.0	Pass
		Xylenes	200	221	10.5	Pass
		Benzene	20	21.6	8.0	Pass
14:35		Toluene	20	21.1	5.5	Pass
		Xylenes	40	41.1	2.8	Pass
	14:35	HCN	25	25.5	2.0	Pass
	14:45	H ₂ S	100	102	2.0	Pass
	14:49	H ₂ S	20	23.3	16.5	Pass
	14:39	Butane	150	158	5.3	Pass
		Pentane	150	161	7.3	Pass
		Hexane	150	151	0.7	Pass
		Heptane	150	146	-2.7	Pass
21:32	21:32	HCN	25	26.1	4.4	Pass
	21:38	H ₂ S	20	20.6	3.0	Pass
	21:36	Butane	150	156	4.0	Pass
		Pentane	150	142	-5.3	Pass
		Hexane	150	149	-0.7	Pass
		Heptane	150	144	-4.0	Pass
	21:29	Benzene	20	20	0.0	Pass
		Toluene	20	18.2	-9.0	Pass
		Xylenes	40	36.9	-7.8	Pass
	21:34	Ethylene	50	55	10.0	Pass
		Propylene	50	51.1	2.2	Pass
		1-Butene	50	49.3	-1.4	Pass
		1-Pentene	50	50.2	0.4	Pass
		1-Hexene	50	49.1	-1.8	Pass
		1,3-Butadiene	50	51.6	3.2	Pass

CCND Mobile Monitoring Van
2023 Q3

Instrument Calibration Check						
Date	Time	Calibration Gas Component	Calibration Value (ppb v)	Response (ppb v)	Difference (% of value)	Pass/Fail
8/2/2023 Dupont	14:14	Ethylene	50	53.4	6.8	Pass
		Propylene	50	51.1	2.2	Pass
		1-Butene	50	51.9	3.8	Pass
		1-Pentene	50	51.3	2.6	Pass
		1-Hexene	50	51.8	3.6	Pass
		1,3-Butadiene	50	53.9	7.8	Pass
	14:06	Benzene	100	109	9.0	Pass
		Toluene	100	103	3.0	Pass
		Xylenes	200	232	16.0	Pass
	14:11	Benzene	20	20.4	2.0	Pass
		Toluene	20	19.5	-2.5	Pass
		Xylenes	40	38.7	-3.2	Pass
	14:19	HCN	25	262	948.0	Fail
	14:24	H ₂ S	100	98.6	-1.4	Pass
	14:26		20	22.4	12.0	Pass
	14:16	Butane	150	157	4.7	Pass
		Pentane	150	150	0.0	Pass
		Hexane	150	156	4.0	Pass
		Heptane	150	151	0.7	Pass
	20:41	HCN	25	25.9	3.6	Pass
	20:39	H ₂ S	20	19.5	-2.5	Pass
	20:53	Butane	150	153	2.0	Pass
		Pentane	150	139	-7.3	Pass
		Hexane	150	141	-6.0	Pass
		Heptane	150	144	-4.0	Pass
	20:51	Benzene	20	22.1	10.5	Pass
		Toluene	20	21.6	8.0	Pass
		Xylenes	40	38.9	-2.8	Pass
	20:44	Ethylene	50	55.5	11.0	Pass
		Propylene	50	48.8	-2.4	Pass
		1-Butene	50	52.2	4.4	Pass
		1-Pentene	50	46.6	-6.8	Pass
		1-Hexene	50	48.1	-3.8	Pass
		1,3-Butadiene	50	50.8	1.6	Pass

CCND Mobile Monitoring Van
2023 Q3

Instrument Calibration Check						
Date	Time	Calibration Gas Component	Calibration Value (ppb v)	Response (ppb v)	Difference (% of value)	Pass/Fail
8/3/2023 Western Hills	10:00	Ethylene	50	52.7	5.4	Pass
		Propylene	50	50.1	0.2	Pass
		1-Butene	50	49.4	-1.2	Pass
		1-Pentene	50	54.5	9.0	Pass
		1-Hexene	50	51.4	2.8	Pass
		1,3-Butadiene	50	49.2	-1.6	Pass
	10:03	Benzene	100	107	7.0	Pass
		Toluene	100	104	4.0	Pass
		Xylenes	200	220	10.0	Pass
	10:08	Benzene	20	20.4	2.0	Pass
		Toluene	20	18.7	-6.5	Pass
		Xylenes	40	37.8	-5.5	Pass
	10:11	HCN	25	27.2	8.8	Pass
	10:19	H ₂ S	100	110	10.0	Pass
	10:21		20	21.3	6.5	Pass
	10:13	Butane	150	154	2.7	Pass
		Pentane	150	144	-4.0	Pass
		Hexane	150	146	-2.7	Pass
		Heptane	150	141	-6.0	Pass
	16:44	HCN	25	23.9	-4.4	Pass
	16:49	H ₂ S	20	22.1	10.5	Pass
	16:46	Butane	150	153	2.0	Pass
		Pentane	150	146	-2.7	Pass
		Hexane	150	148	-1.3	Pass
		Heptane	150	142	-5.3	Pass
	16:39	Benzene	20	20.9	4.5	Pass
		Toluene	20	18.9	-5.5	Pass
		Xylenes	40	40.3	0.7	Pass
	16:42	Ethylene	50	50.6	1.2	Pass
		Propylene	50	49.9	-0.2	Pass
		1-Butene	50	52.2	4.4	Pass
		1-Pentene	50	48.6	-2.8	Pass
		1-Hexene	50	52	4.0	Pass
		1,3-Butadiene	50	51.6	3.2	Pass

CCND Mobile Monitoring Van
2023 Q3

Date	Time	Gas Component	Instrument Calibration Check			
			Calibration Value (ppb v)	Response (ppb v)	Difference (% of value)	Pass/Fail
8/4/2023 Adams City	8:23	Ethylene	50	51.4	2.8	Pass
		Propylene	50	56.1	12.2	Pass
		1-Butene	50	51.4	2.8	Pass
		1-Pentene	50	58.1	16.2	Pass
		1-Hexene	50	55.5	11.0	Pass
		1,3-Butadiene	50	56.1	12.2	Pass
	8:32	Benzene	100	104	4.0	Pass
		Toluene	100	100	0.0	Pass
		Xylenes	200	207	3.5	Pass
	8:39	Benzene	20	19.6	-2.0	Pass
		Toluene	20	18.5	-7.5	Pass
		Xylenes	40	36.8	-8.0	Pass
	8:29	HCN	25	23.5	-6.0	Pass
	8:19	H ₂ S	100	104	4.0	Pass
	8:20		20	20.5	2.5	Pass
	8:25	Butane	150	157	4.7	Pass
		Pentane	150	160	6.7	Pass
		Hexane	150	156	4.0	Pass
		Heptane	150	153	2.0	Pass
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12:44 12:47 12:40 12:37 12:41	12:44	HCN	25	23.9	-4.4	Pass
	12:47	H ₂ S	20	18.9	-5.5	Pass
	12:40	Butane	150	154	2.7	Pass
		Pentane	150	166	10.7	Pass
		Hexane	150	160	6.7	Pass
		Heptane	150	161	7.3	Pass
		Benzene	20	19.4	-3.0	Pass
		Toluene	20	18.2	-9.0	Pass
		Xylenes	40	37.9	-5.3	Pass
		Ethylene	50	47	-6.0	Pass
		Propylene	50	55.8	11.6	Pass
		1-Butene	50	54.2	8.4	Pass
		1-Pentene	50	58.1	16.2	Pass
		1-Hexene	50	56.4	12.8	Pass
		1,3-Butadiene	50	56	12.0	Pass

APPENDIX E

CALIBRATION GAS CERTIFICATION SHEETS

CERTIFICATE OF ANALYSIS

Grade of Product: CERTIFIED STANDARD-SPEC

Customer:	MONTROSE AIR QUALITY SERVICES LLC	Reference Number:	126-402278540-1
Part Number:	X02NI99C15W0061	Cylinder Volume:	144.3 CF
Cylinder Number:	CC519990	Cylinder Pressure:	2015 PSIG
Laboratory:	124 - La Porte Mix - TX	Valve Outlet:	330
Analysis Date:	Dec 14, 2021		
Lot Number:	126-402278540-1		
	Expiration Date: Dec 14, 2024		

Product composition verified by direct comparison to calibration standards traceable to N.I.S.T. weights and/or N.I.S.T. Gas Mixture reference materials.

ANALYTICAL RESULTS

Component	Req Conc	Actual Concentration (Mole %)	Analytical Uncertainty
HYDROGEN SULFIDE	1.000 PPM	1.084 PPM	+/-5%
NITROGEN	Balance		

Notes: MONTROSE AIR QUALITY SERVICES LLC
PO3: PO018078



Signature on file

Approved for Release

Page 1 of 1



an Air Liquide company

Airgas Specialty Gases
Airgas USA, LLC
616 Miller Cut Off Road
La Porte, TX 77571
Airgas.com

CERTIFICATE OF ANALYSIS

Grade of Product: CERTIFIED STANDARD-SPEC

Customer: *CRYSTAL LAKE , IL* MONTROSE AIR QUALITY SERVICES
Part X06NI99C15A00A3

Reference Number: 126-402159020-1

Number:
Cylinder CC344804

Cylinder Volume: 144.3 CF

Number:
Laboratory: 124 - La Porte Mix - TX
Analysis Jul 30, 2021

Cylinder Pressure: 2015 PSIG
Valve Outlet: 350

Date:
Lot Number: 126-402159020-1

Expiration Date: Jul 30, 2024

Product composition verified by direct comparison to calibration standards traceable to N.I.S.T. weights and/or N.I.S.T. Gas Mixture reference materials.

ANALYTICAL RESULTS

Component	Req Conc	Actual Concentration (Mole %)	Analytical Uncertainty
HEXANE	1.000 PPM	0.9950 PPM	+/- 5%
N BUTANE	1.000 PPM	1.002 PPM	+/- 5%
N HEPTANE	1.000 PPM	1.000 PPM	+/- 5%
N PENTANE	1.000 PPM	1.000 PPM	+/- 5%
PROPANE	1.000 PPM	1.009 PPM	+/- 5%
NITROGEN	Balance		

Notes:

PO # PO-011307




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Airgas USA, LLC
4646 Linden Rd
Rockford, IL 61109
Airgas.com

CERTIFICATE OF BATCH ANALYSIS

Grade of Product: ZERO

Part Number:	AI Z15A	Reference Number:	152-402047887-1
Cylinder Analyzed:	CC235228	Cylinder Volume:	146.0 CF
Laboratory:	192 - Rockford IL Fill Plant (N513) - IL	Cylinder Pressure:	2000 PSIG
Analysis Date:	Mar 03, 2021	Valve Outlet:	590
Lot Number:	152-402047887-1		

ANALYTICAL RESULTS

Component	Requested Purity	Certified Concentration
AIR		
THC	< 1.0 PPM	0.043 PPM
Percent Oxygen	20-22 %	20.82 %
Moisture	< 3.0 PPM	0.07 PPM

Cylinders in Batch:

CC235228, XC002876B

Impurities verified against analytical standards traceable to NIST by weight and/or analysis.

Signature on file

Approved for Release

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an Air Liquide company

Airgas Specialty Gases

Airgas USA, LLC
616 Miller Cut Off Road
La Porte, TX 77571
Airgas.com

CERTIFICATE OF ANALYSIS

Grade of Product: CERTIFIED STANDARD-SPEC

Customer: MONTROSE AIR QUALITY SERVICES LLC - CRYSTAL

LAKE,

Part X07NI99C15A00A9

Reference Number: 126-402159021-1

Number:

Cylinder CC164840

Cylinder Volume: 144.3 CF

Number:

Laboratory: 124 - La Porte Mix - TX

Cylinder Pressure: 2015 PSIG

Analysis Aug 09, 2021

Valve Outlet: 350

Date:

Lot Number: 126-402159021-1

Expiration Date: Aug 09, 2023

Product composition verified by direct comparison to calibration standards traceable to N.I.S.T. weights and/or N.I.S.T. Gas Mixture reference materials.

ANALYTICAL RESULTS

Component	Req Conc	Actual Concentration (Mole %)	Analytical Uncertainty
1 BUTENE	1.000 PPM	0.9918 PPM	+/- 5%
1 HEXENE	1.000 PPM	1.003 PPM	+/- 5%
1 PENTENE	1.000 PPM	1.005 PPM	+/- 5%
1,3 BUTADIENE	1.000 PPM	1.005 PPM	+/- 5%
ETHYLENE	1.000 PPM	1.087 PPM	+/- 5%
PROPYLENE	1.000 PPM	1.006 PPM	+/- 5%
NITROGEN	Balance		

Notes:

MONTROSE AIR QUALITY SERVICES LLC

PO#: PO-011307

NITROGEN BALANCE : 99.99939022%



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