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TABLE OF CONTENTS

| <u>SEC</u> | TION | <u>N</u> <u>P</u> | AGE | | | | | | |
|-----------------------------|------|--|-----|--|--|--|--|--|--|
| EXE | CUT | IVE SUMMARY | 3 | | | | | | |
| 1.0 | INTF | RODUCTION | 4 | | | | | | |
| 2.0 MOBILE SAMPLING PROGRAM | | | | | | | | | |
| | 2.1 | Mobile Van Air Sampling Description | 4 | | | | | | |
| | 2.2 | Mobile Monitoring Van Air Sampling Methods | 6 | | | | | | |
| | 2.3 | Screening Health Risk Assessment Methods | 7 | | | | | | |
| 3.0 | SUM | IMARY AND DISCUSSION OF RESULTS | 10 | | | | | | |
| | 3.1 | Summary of Mobile Monitoring Van Results | 10 | | | | | | |
| | 3.2 | Screening Health Risk Assessment Results | 10 | | | | | | |
| | 3.3 | Uncertainty Evaluation | 18 | | | | | | |
| | 3.4 | Program Changes | 18 | | | | | | |
| 1 16. | | | | | | | | | |

LIST OF APPENDICES

- A ISOMER CHEMICAL SAMPLING DETAILS
- B DAILY WIND ROSES
- C SCREENING RISK ASSESSMENT DETAILS (ALPHABETICAL ORDER BY NEIGHBORHOOD NAME)
- D CALIBRATION AND QA/QC DATA
- E CALIBRATION GAS CERTIFICATION SHEETS

LIST OF TABLES

| 2-1 | MOBILE MONITORING VAN PROGRAM CHEMICALS | 5 |
|-----|---|---|
| 2-2 | NEIGHBORHOOD MONITORING PROGRAM DETAILS | 6 |

LIST OF FIGURES

| 2-1 | MOBILE MONITORING VAN PROGRAM ROUTE THROUGH SIX NEIGHBORHOOD AREAS | 7 |
|-----|---|-----|
| 3-1 | ADAMS CITY NEIGHBORHOOD: NOVEMBER 18, 2022 | .12 |
| 3-2 | DUPONT NEIGHBORHOOD: NOVEMBER 15, 2022 | .13 |
| 3-3 | ELYRA-SWANSEA NEIGHBORHOOD: NOVEMBER 17, 2022 | .14 |
| 3-4 | GLOBEVILLE NEIGHBORHOOD: NOVEMBER 17, 2022 | .15 |
| 3-5 | PIONEER PARK NEIGHBORHOOD: NOVEMBER 16, 2022 | .16 |
| 3-6 | WESTERN HILLS NEIGHBORHOOD: NOVEMBER 14, 2022 | .17 |



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 fourth quarter 2022 sampling period (November 14-18), the mobile monitoring van was in a total of six neighborhoods and collected more than 74,800 data points across five days of monitoring, resulting in approximately 53,700 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.

¹ 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.



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, the list of 65 chemicals in Table 2-1 were measured to determine the instantaneous ambient concentrations. 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 every 1 second. The details of the monitored neighborhoods are listed in Table 2-2 and are shown in Figure 2-1.



| 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 |

TABLE 2-1 MOBILE MONITORING VAN PROGRAM CHEMICALS²



² See Appendix A for isomer analysis details

| Neighborhood | Area (square miles) | Sampling Date | Start Time | End Time | Total Data Points Collected | Total Hourly Rolling Averages Calculated |
|----------------|---------------------------|---------------|------------|----------|-----------------------------------|--|
| Adams City | 0.41 | 11/18/22 | 10:23 | 13:33 | 11,433 | 7,906 |
| Dupont | 1.4 | 11/15/22 | 12:26 | 16:29 | 14,527 | 11,000 |
| Elyria-Swansea | 1.2 | 11/17/22 | 13:58 | 16:30 | 9,105 | 5,578 |
| Globeville | 0.44 | 11/17/22 | 10:48 | 13:39 | 10,274 | 6,747 |
| Pioneer Park | 1.7 | 11/16/22 | 10:31 | 14:36 | 14,654 | 11,127 |
| Western Hills | 1.6 | 11/14/22 | 10:16 | 14:25 | 14,874 | 11,347 |

TABLE 2-2 NEIGHBORHOOD MONITORING PROGRAM DETAILS

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 multichemical 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. Measurements that were collected from transition periods or from moving between neighborhoods were excluded in this assessment.

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)



³https://www.atsdr.cdc.gov/minimalrisklevels/#:~:text=The%20ATSDR%2C%20in%20response%20to,minimal%20risk%20levels%2 0(MRLs)

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

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 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, 39,7831-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 3-8).

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



⁵

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

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

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.

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 74,800 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 top five 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 and trimethylbenzene group were the chemicals or isomer groupings resulting in the highest HQ in each neighborhood, accounting for over



70% of the total calculated HI values. However, all HI values calculated in all six neighborhoods were below one (Figures 3-1 through 3-6).

• 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 ADAMS CITY NEIGHBORHOOD: NOVEMBER 18, 2022



FIGURE 3-2 DUPONT NEIGHBORHOOD: NOVEMBER 15, 2022







FIGURE 3-3 ELYRA-SWANSEA NEIGHBORHOOD: NOVEMBER 17, 2022



FIGURE 3-4 GLOBEVILLE NEIGHBORHOOD: NOVEMBER 17, 2022





FIGURE 3-5 PIONEER PARK NEIGHBORHOOD: NOVEMBER 16, 2022





FIGURE 3-6 WESTERN HILLS NEIGHBORHOOD: NOVEMBER 14, 2022





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:

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



APPENDIX B DAILY WIND ROSES

















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



Mobile Laboratory Sampling Data Summary and Risk Assessment Adams City Neighborhood | November 18, 2022

| 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 | 11,433 | 0.15 | 7,906 | 0.02 | 0.02 | 670,000 | 298 | OEHHA Acute REL | 0.00008 |
| ACETYLENE | 74-86-2 | 11,433 | 0.71 | 7,906 | 0.15 | 0.19 | NR | 25,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| BENZENE | 71-43-2 | 11,433 | 9.00 | 7,906 | 0.29 | 0.33 | 52,000 | 9 | ATSDR Acute MRL | 0.03613 |
| BUTANES* | 75-28-5 | 11,433 | 10.09 | 7,906 | 2.61 | 2.71 | NR | 33000 | TCEQ Short-Term AMCV Health | 0.00008 |
| BUTENES* | 590-18-1 | 11,433 | 33.92 | 7,906 | 1.61 | 1.76 | NR | 15000 | TCEQ Short-Term AMCV Health | 0.00012 |
| CARBON DISULFIDE | 75-15-0 | 11,433 | 0.03 | 7,906 | 0.00 | 0.00 | 13,000 | 1,991 | OEHHA Acute REL | 0.00000 |
| CYCLOPENTANES* | 287-92-3 | 11,433 | 23.65 | 7,906 | 2.62 | 2.74 | NR | 5,900 | TCEQ Short-Term AMCV Health | 0.00047 |
| DECANES | 124-18-5 | 11,433 | 0.07 | 7,906 | 0.03 | 0.03 | NR | 1,000 | TCEQ Short-Term AMCV Health | 0.00003 |
| DIETHYLBENZENES* | 141-93-5 | 11,433 | 0.28 | 7,906 | 0.12 | 0.13 | NR | 450 | TCEQ Short-Term AMCV Health | 0.00030 |
| DIMETHYLCYCLOHEXANES* | 638-04-0 | 11,433 | 0.17 | 7,906 | 0.11 | 0.11 | NR | 4,000 | CDPHE | 0.00003 |
| DODECANES | 112-40-3 | 11,433 | 0.02 | 7,906 | 0.00 | 0.00 | NR | 1720 | CDPHE | 0.00000 |
| ETHYLENE | 74-85-1 | 11,433 | 22.44 | 7,906 | 5.93 | 6.01 | NR | 500,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEPTANES* | 142-82-5 | 11,433 | 0.15 | 7,906 | 0.08 | 0.08 | NR | 8,300 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEXANES* | 110-54-3 | 11,433 | 0.23 | 7,906 | 0.12 | 0.12 | NR | 5,400 | TCEQ Short-Term AMCV Health | 0.00002 |
| HEXENES* | 592-41-6 | 11,433 | 10.91 | 7,906 | 0.76 | 0.83 | NR | 500 | TCEQ Short-Term AMCV Health | 0.00165 |
| HYDROGEN CYANIDE | 74-90-8 | 11,433 | 1.48 | 7,906 | 0.14 | 0.24 | 2,000 | 308 | OEHHA Acute REL | 0.00078 |
| HYDROGEN SULFIDE | 7783-06-4 | 11,433 | 1.13 | 7,906 | 0.15 | 0.22 | 510 | 70 | ATSDR Acute MRL | 0.00315 |
| ISOPRENE | 78-79-5 | 11,433 | 1.69 | 7,906 | 0.17 | 0.19 | NR | 1,400 | TCEQ Short-Term AMCV Health | 0.00014 |
| METHANOL | 67-56-1 | 11,433 | 34.79 | 7,906 | 5.06 | 5.36 | 530,000 | 21,366 | OEHHA Acute REL | 0.00025 |
| METHYLCYCLOHEXANE | 108-87-2 | 11,433 | 0.30 | 7,906 | 0.05 | 0.05 | NR | 4,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| NONANES | 111-84-2 | 11,433 | 0.12 | 7,906 | 0.07 | 0.07 | NR | 3,000 | TCEQ Short-Term AMCV Health | 0.00002 |
| OCTANES* | 111-65-9 | 11,433 | 0.13 | 7,906 | 0.07 | 0.08 | NR | 4,100 | TCEQ Short-Term AMCV Health | 0.00002 |
| PENTANES* | 109-66-0 | 11,433 | 0.47 | 7,906 | 0.20 | 0.20 | NR | 68,000 | TCEQ Short-Term AMCV Health | 0.00000 |
| PROPYLENE | 115-07-1 | 11,433 | 11.83 | 7,906 | 0.32 | 0.38 | NR | NA | NE | |
| STYRENE | 100-42-5 | 11,433 | 0.39 | 7,906 | 0.06 | 0.07 | 20,000 | 5,000 | ATSDR Acute MRL | 0.00001 |
| TETRACHLOROETHYLENE | 127-18-4 | 11,433 | 0.15 | 7,906 | 0.01 | 0.01 | 35,000 | 6 | ATSDR Acute MRL | 0.00228 |
| TOLUENE | 108-88-3 | 11,433 | 17.98 | 7,906 | 0.61 | 0.71 | 67,000 | 2,000 | ATSDR Acute MRL | 0.00035 |
| TRIMETHYLBENZENES* | 622-96-8 | 11,433 | 2.65 | 7,906 | 0.21 | 0.26 | 50,000 | 250 | TCEQ Short-Term AMCV Health | 0.00102 |
| UNDECANES | 1120-21-4 | 11,433 | 0.05 | 7,906 | 0.02 | 0.02 | NR | 550 | TCEQ Short-Term AMCV Health | 0.00004 |
| XYLENES* | 1330-20-7 | 11,433 | 17.35 | 7,906 | 1.26 | 1.40 | 130,000 | 2,000 | ATSDR Acute MRL | 0.00070 |
| | | | | | | | | | Hazard Index | 0.04772 |

NR= According to EPA, AEGL is "not recommended due to insufficient data" NA= Not Available NC= Not Calculated



Mobile Laboratory Sampling Data Summary and Risk Assessment

Dupont Neighborhood | November 15, 2022

| 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 | 14,527 | 0.15 | 11,000 | 0.01 | 0.01 | 670,000 | 298 | OEHHA Acute REL | 0.00005 |
| ACETYLENE | 74-86-2 | 14,527 | 0.70 | 11,000 | 0.10 | 0.14 | NR | 25,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| BENZENE | 71-43-2 | 14,527 | 6.42 | 11,000 | 0.24 | 0.26 | 52,000 | 9 | ATSDR Acute MRL | 0.02924 |
| BUTANES* | 75-28-5 | 14,527 | 26.78 | 11,000 | 3.69 | 3.87 | NR | 33000 | TCEQ Short-Term AMCV Health | 0.00012 |
| BUTENES* | 590-18-1 | 14,527 | 18.22 | 11,000 | 3.08 | 3.25 | NR | 15000 | TCEQ Short-Term AMCV Health | 0.00022 |
| CARBON DISULFIDE | 75-15-0 | 14,527 | 0.04 | 11,000 | 0.00 | 0.00 | 13,000 | 1,991 | OEHHA Acute REL | 0.00000 |
| CYCLOPENTANES* | 287-92-3 | 14,527 | 11.33 | 11,000 | 1.49 | 1.65 | NR | 5,900 | TCEQ Short-Term AMCV Health | 0.00028 |
| DECANES | 124-18-5 | 14,527 | 0.11 | 11,000 | 0.05 | 0.06 | NR | 1,000 | TCEQ Short-Term AMCV Health | 0.00006 |
| DIETHYLBENZENES* | 141-93-5 | 14,527 | 0.16 | 11,000 | 0.07 | 0.08 | NR | 450 | TCEQ Short-Term AMCV Health | 0.00017 |
| DIMETHYLCYCLOHEXANES* | 638-04-0 | 14,527 | 0.08 | 11,000 | 0.03 | 0.03 | NR | 4,000 | CDPHE | 0.00001 |
| DODECANES | 112-40-3 | 14,527 | 0.01 | 11,000 | 0.00 | 0.00 | NR | 1720 | CDPHE | 0.00000 |
| ETHYLENE | 74-85-1 | 14,527 | 33.55 | 11,000 | 5.30 | 5.41 | NR | 500,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEPTANES* | 142-82-5 | 14,527 | 0.16 | 11,000 | 0.06 | 0.06 | NR | 8,300 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEXANES* | 110-54-3 | 14,527 | 0.22 | 11,000 | 0.05 | 0.06 | NR | 5,400 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEXENES* | 592-41-6 | 14,527 | 7.45 | 11,000 | 0.93 | 1.01 | NR | 500 | TCEQ Short-Term AMCV Health | 0.00202 |
| HYDROGEN CYANIDE | 74-90-8 | 14,527 | 0.80 | 11,000 | 0.17 | 0.24 | 2,000 | 308 | OEHHA Acute REL | 0.00077 |
| HYDROGEN SULFIDE | 7783-06-4 | 14,527 | 1.07 | 11,000 | 0.18 | 0.24 | 510 | 70 | ATSDR Acute MRL | 0.00343 |
| ISOPRENE | 78-79-5 | 14,527 | 0.90 | 11,000 | 0.08 | 0.10 | NR | 1,400 | TCEQ Short-Term AMCV Health | 0.00007 |
| METHANOL | 67-56-1 | 14,527 | 23.54 | 11,000 | 2.67 | 2.86 | 530,000 | 21,366 | OEHHA Acute REL | 0.00013 |
| METHYLCYCLOHEXANE | 108-87-2 | 14,527 | 0.13 | 11,000 | 0.06 | 0.06 | NR | 4,000 | TCEQ Short-Term AMCV Health | 0.00002 |
| NONANES | 111-84-2 | 14,527 | 0.07 | 11,000 | 0.02 | 0.02 | NR | 3,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| OCTANES* | 111-65-9 | 14,527 | 0.41 | 11,000 | 0.03 | 0.04 | NR. | 4,100 | TCEQ Short-Term AMCV Health | 0.00001 |
| PENTANES* | 109-66-0 | 14,527 | 1.43 | 11,000 | 1.13 | 1.13 | NR. | 68,000 | TCEQ Short-Term AMCV Health | 0.00002 |
| PROPYLENE | 115-07-1 | 14,527 | 3.44 | 11,000 | 0.13 | 0.20 | NR. | NA | NE | |
| STYRENE | 100-42-5 | 14,527 | 0.37 | 11,000 | 0.09 | 0.10 | 20,000 | 5,000 | ATSDR Acute MRL | 0.00002 |
| TETRACHLOROETHYLENE | 127-18-4 | 14,527 | 0.09 | 11,000 | 0.01 | 0.01 | 35,000 | 6 | ATSDR Acute MRL | 0.00108 |
| TOLUENE | 108-88-3 | 14,527 | 21.12 | 11,000 | 0.42 | 0.51 | 67,000 | 2,000 | ATSDR Acute MRL | 0.00025 |
| TRIMETHYLBENZENES* | 622-96-8 | 14,527 | 3.28 | 11,000 | 0.19 | 0.23 | 50,000 | 250 | TCEQ Short-Term AMCV Health | 0.00092 |
| UNDECANES | 1120-21-4 | 14,527 | 0.04 | 11,000 | 0.00 | 0.01 | NR. | 550 | TCEQ Short-Term AMCV Health | 0.00001 |
| XYLENES* | 1330-20-7 | 14,527 | 21.73 | 11,000 | 0.62 | 0.75 | 130,000 | 2,000 | ATSDR Acute MRL | 0.00038 |
| | | | | | | | | | Hazard Index | 0.03931 |

NR= According to EPA, AEGL is "not recommended due to insufficient data" NA= Not Available NC= Not Calculated



Mobile Laboratory Sampling Data Summary and Risk Assessment

Elyria-Swansea Neighborhood | November 17, 2022

| 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 | 9,105 | 0.18 | 5,578 | 0.04 | 0.04 | 670,000 | 298 | OEHHA Acute REL | 0.00015 |
| ACETYLENE | 74-86-2 | 9,105 | 0.68 | 5,578 | 0.11 | 0.14 | NR | 25,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| BENZENE | 71-43-2 | 9,105 | 17.92 | 5,578 | 0.29 | 0.35 | 52,000 | 9 | ATSDR Acute MRL | 0.03933 |
| BUTANES* | 75-28-5 | 9,105 | 10.18 | 5,578 | 3.52 | 3.63 | NR | 33000 | TCEQ Short-Term AMCV Health | 0.00011 |
| BUTENES* | 590-18-1 | 9,105 | 67.72 | 5,578 | 1.49 | 1.80 | NR | 15000 | TCEQ Short-Term AMCV Health | 0.00012 |
| CARBON DISULFIDE | 75-15-0 | 9,105 | 0.03 | 5,578 | 0.00 | 0.00 | 13,000 | 1,991 | OEHHA Acute REL | 0.00000 |
| CYCLOPENTANES* | 287-92-3 | 9,105 | 87.88 | 5,578 | 2.87 | 3.19 | NR | 5,900 | TCEQ Short-Term AMCV Health | 0.00054 |
| DECANES | 124-18-5 | 9,105 | 0.07 | 5,578 | 0.03 | 0.03 | NR | 1,000 | TCEQ Short-Term AMCV Health | 0.00003 |
| DIETHYLBENZENES* | 141-93-5 | 9,105 | 0.40 | 5,578 | 0.11 | 0.12 | NR | 450 | TCEQ Short-Term AMCV Health | 0.00028 |
| DIMETHYLCYCLOHEXANES* | 638-04-0 | 9,105 | 0.39 | 5,578 | 0.02 | 0.02 | NR | 4,000 | CDPHE | 0.00001 |
| DODECANES | 112-40-3 | 9,105 | 0.01 | 5,578 | 0.00 | 0.00 | NR | 1720 | CDPHE | 0.00000 |
| ETHYLENE | 74-85-1 | 9,105 | 10.37 | 5,578 | 8.84 | 8.85 | NR | 500,000 | TCEQ Short-Term AMCV Health | 0.00002 |
| HEPTANES* | 142-82-5 | 9,105 | 0.13 | 5,578 | 0.04 | 0.04 | NR | 8,300 | TCEQ Short-Term AMCV Health | 0.00000 |
| HEXANES* | 110-54-3 | 9,105 | 0.49 | 5,578 | 0.26 | 0.26 | NR | 5,400 | TCEQ Short-Term AMCV Health | 0.00005 |
| HEXENES* | 592-41-6 | 9,105 | 57.70 | 5,578 | 1.28 | 1.43 | NR | 500 | TCEQ Short-Term AMCV Health | 0.00285 |
| HYDROGEN CYANIDE | 74-90-8 | 9,105 | 1.43 | 5,578 | 0.17 | 0.21 | 2,000 | 308 | OEHHA Acute REL | 0.00069 |
| HYDROGEN SULFIDE | 7783-06-4 | 9,105 | 4.92 | 5,578 | 0.18 | 0.24 | 510 | 70 | ATSDR Acute MRL | 0.00346 |
| ISOPRENE | 78-79-5 | 9,105 | 3.42 | 5,578 | 0.22 | 0.23 | NR | 1,400 | TCEQ Short-Term AMCV Health | 0.00017 |
| METHANOL | 67-56-1 | 9,105 | 44.19 | 5,578 | 5.83 | 6.77 | 530,000 | 21,366 | OEHHA Acute REL | 0.00032 |
| METHYLCYCLOHEXANE | 108-87-2 | 9,105 | 0.73 | 5,578 | 0.15 | 0.15 | NR | 4,000 | TCEQ Short-Term AMCV Health | 0.00004 |
| NONANES | 111-84-2 | 9,105 | 0.05 | 5,578 | 0.01 | 0.01 | NR | 3,000 | TCEQ Short-Term AMCV Health | 0.00000 |
| OCTANES* | 111-65-9 | 9,105 | 0.13 | 5,578 | 0.03 | 0.03 | NR | 4,100 | TCEQ Short-Term AMCV Health | 0.00001 |
| PENTANES* | 109-66-0 | 9,105 | 0.93 | 5,578 | 0.90 | 0.90 | NR | 68,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| PROPYLENE | 115-07-1 | 9,105 | 11.99 | 5,578 | 0.25 | 0.28 | NR | NA | NE | |
| STYRENE | 100-42-5 | 9,105 | 0.53 | 5,578 | 0.07 | 0.08 | 20,000 | 5,000 | ATSDR Acute MRL | 0.00002 |
| TETRACHLOROETHYLENE | 127-18-4 | 9,105 | 0.05 | 5,578 | 0.01 | 0.01 | 35,000 | 6 | ATSDR Acute MRL | 0.00128 |
| TOLUENE | 108-88-3 | 9,105 | 84.62 | 5,578 | 0.86 | 1.19 | 67,000 | 2,000 | ATSDR Acute MRL | 0.00060 |
| TRIMETHYLBENZENES* | 622-96-8 | 9,105 | 19.66 | 5,578 | 0.35 | 0.45 | 50,000 | 250 | TCEQ Short-Term AMCV Health | 0.00179 |
| UNDECANES | 1120-21-4 | 9,105 | 0.05 | 5,578 | 0.02 | 0.02 | NR | 550 | TCEQ Short-Term AMCV Health | 0.00004 |
| XYLENES* | 1330-20-7 | 9,105 | 95.23 | 5,578 | 1.58 | 1.98 | 130,000 | 2,000 | ATSDR Acute MRL | 0.00099 |
| | | | | | | | | | Hazard Index | 0.05291 |

NR= According to EPA, AEGL is "not recommended due to insufficient data" NA= Not Available NC= Not Calculated



Mobile Laboratory Sampling Data Summary and Risk Assessment Globeville Neighborhood | November 17, 2022

| 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 | 10,274 | 0.16 | 6,747 | 0.01 | 0.02 | 670,000 | 298 | OEHHA Acute REL | 0.00007 |
| ACETYLENE | 74-86-2 | 10,274 | 0.87 | 6,747 | 0.11 | 0.17 | NR | 25,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| BENZENE | 71-43-2 | 10,274 | 2.72 | 6,747 | 0.24 | 0.29 | 52,000 | 9 | ATSDR Acute MRL | 0.03199 |
| BUTANES* | 75-28-5 | 10,274 | 28.39 | 6,747 | 1.75 | 1.86 | NR | 33000 | TCEQ Short-Term AMCV Health | 0.00006 |
| BUTENES* | 590-18-1 | 10,274 | 9.10 | 6,747 | 0.91 | 1.16 | NR | 15000 | TCEQ Short-Term AMCV Health | 0.00008 |
| CARBON DISULFIDE | 75-15-0 | 10,274 | 0.03 | 6,747 | 0.00 | 0.00 | 13,000 | 1,991 | OEHHA Acute REL | 0.00000 |
| CYCLOPENTANES* | 287-92-3 | 10,274 | 39.69 | 6,747 | 1.14 | 1.61 | NR | 5,900 | TCEQ Short-Term AMCV Health | 0.00027 |
| DECANES | 124-18-5 | 10,274 | 0.09 | 6,747 | 0.03 | 0.04 | NR | 1,000 | TCEQ Short-Term AMCV Health | 0.00004 |
| DIETHYLBENZENES* | 141-93-5 | 10,274 | 0.26 | 6,747 | 0.13 | 0.14 | NR | 450 | TCEQ Short-Term AMCV Health | 0.00031 |
| DIMETHYLCYCLOHEXANES* | 638-04-0 | 10,274 | 0.10 | 6,747 | 0.02 | 0.02 | NR | 4,000 | CDPHE | 0.00001 |
| DODECANES | 112-40-3 | 10,274 | 0.02 | 6,747 | 0.00 | 0.00 | NR | 1720 | CDPHE | 0.00000 |
| ETHYLENE | 74-85-1 | 10,274 | 11.92 | 6,747 | 6.79 | 6.84 | NR | 500,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEPTANES* | 142-82-5 | 10,274 | 0.14 | 6,747 | 0.04 | 0.05 | NR | 8,300 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEXANES* | 110-54-3 | 10,274 | 0.24 | 6,747 | 0.07 | 0.07 | NR | 5,400 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEXENES* | 592-41-6 | 10,274 | 4.59 | 6,747 | 0.51 | 0.73 | NR | 500 | TCEQ Short-Term AMCV Health | 0.00146 |
| HYDROGEN CYANIDE | 74-90-8 | 10,274 | 0.82 | 6,747 | 0.12 | 0.25 | 2,000 | 308 | OEHHA Acute REL | 0.00081 |
| HYDROGEN SULFIDE | 7783-06-4 | 10,274 | 0.70 | 6,747 | 0.17 | 0.26 | 510 | 70 | ATSDR Acute MRL | 0.00369 |
| ISOPRENE | 78-79-5 | 10,274 | 0.95 | 6,747 | 0.22 | 0.24 | NR | 1,400 | TCEQ Short-Term AMCV Health | 0.00017 |
| METHANOL | 67-56-1 | 10,274 | 13.41 | 6,747 | 5.16 | 5.28 | 530,000 | 21,366 | OEHHA Acute REL | 0.00025 |
| METHYLCYCLOHEXANE | 108-87-2 | 10,274 | 0.18 | 6,747 | 0.10 | 0.11 | NR | 4,000 | TCEQ Short-Term AMCV Health | 0.00003 |
| NONANES | 111-84-2 | 10,274 | 0.06 | 6,747 | 0.02 | 0.02 | NR | 3,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| OCTANES* | 111-65-9 | 10,274 | 2.75 | 6,747 | 0.03 | 0.08 | NR | 4,100 | TCEQ Short-Term AMCV Health | 0.00002 |
| PENTANES* | 109-66-0 | 10,274 | 0.72 | 6,747 | 0.26 | 0.27 | NR | 68,000 | TCEQ Short-Term AMCV Health | 0.00000 |
| PROPYLENE | 115-07-1 | 10,274 | 4.28 | 6,747 | 0.35 | 0.42 | NR | NA | NE | |
| STYRENE | 100-42-5 | 10,274 | 1.63 | 6,747 | 0.09 | 0.11 | 20,000 | 5,000 | ATSDR Acute MRL | 0.00002 |
| TETRACHLOROETHYLENE | 127-18-4 | 10,274 | 0.15 | 6,747 | 0.00 | 0.00 | 35,000 | 6 | ATSDR Acute MRL | 0.00076 |
| TOLUENE | 108-88-3 | 10,274 | 10.01 | 6,747 | 0.43 | 0.47 | 67,000 | 2,000 | ATSDR Acute MRL | 0.00023 |
| TRIMETHYLBENZENES* | 622-96-8 | 10,274 | 1.57 | 6,747 | 0.10 | 0.12 | 50,000 | 250 | TCEQ Short-Term AMCV Health | 0.00047 |
| UNDECANES | 1120-21-4 | 10,274 | 0.07 | 6,747 | 0.02 | 0.03 | NR | 550 | TCEQ Short-Term AMCV Health | 0.00005 |
| XYLENES* | 1330-20-7 | 10,274 | 9.91 | 6,747 | 0.64 | 0.70 | 130,000 | 2,000 | ATSDR Acute MRL | 0.00035 |
| | | | | | | | | | Hazard Index | 0.04117 |

NR= According to EPA, AEGL is "not recommended due to insufficient data" NA= Not Available NC= Not Calculated



Mobile Laboratory Sampling Data Summary and Risk Assessment

Pioneer Park Neighborhood | November 16, 2022

| 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 | 14,654 | 0.18 | 11,127 | 0.03 | 0.04 | 670,000 | 298 | OEHHA Acute REL | 0.00012 |
| ACETYLENE | 74-86-2 | 14,654 | 0.84 | 11,127 | 0.16 | 0.20 | NR | 25,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| BENZENE | 71-43-2 | 14,654 | 6.57 | 11,127 | 0.24 | 0.27 | 52,000 | 9 | ATSDR Acute MRL | 0.02954 |
| BUTANES* | 75-28-5 | 14,654 | 16.72 | 11,127 | 1.72 | 1.82 | NR | 33000 | TCEQ Short-Term AMCV Health | 0.00006 |
| BUTENES* | 590-18-1 | 14,654 | 23.56 | 11,127 | 1.30 | 1.58 | NR | 15000 | TCEQ Short-Term AMCV Health | 0.00011 |
| CARBON DISULFIDE | 75-15-0 | 14,654 | 0.03 | 11,127 | 0.00 | 0.00 | 13,000 | 1,991 | OEHHA Acute REL | 0.00000 |
| CYCLOPENTANES* | 287-92-3 | 14,654 | 18.02 | 11,127 | 1.29 | 1.57 | NR | 5,900 | TCEQ Short-Term AMCV Health | 0.00027 |
| DECANES | 124-18-5 | 14,654 | 0.08 | 11,127 | 0.04 | 0.04 | NR | 1,000 | TCEQ Short-Term AMCV Health | 0.00004 |
| DIETHYLBENZENES* | 141-93-5 | 14,654 | 0.19 | 11,127 | 0.09 | 0.11 | NR | 450 | TCEQ Short-Term AMCV Health | 0.00024 |
| DIMETHYLCYCLOHEXANES* | 638-04-0 | 14,654 | 0.11 | 11,127 | 0.02 | 0.02 | NR | 4,000 | CDPHE | 0.00001 |
| DODECANES | 112-40-3 | 14,654 | 0.03 | 11,127 | 0.00 | 0.00 | NR | 1720 | CDPHE | 0.00000 |
| ETHYLENE | 74-85-1 | 14,654 | 8.27 | 11,127 | 4.96 | 5.02 | NR | 500,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEPTANES* | 142-82-5 | 14,654 | 0.24 | 11,127 | 0.17 | 0.17 | NR | 8,300 | TCEQ Short-Term AMCV Health | 0.00002 |
| HEXANES* | 110-54-3 | 14,654 | 0.28 | 11,127 | 0.18 | 0.18 | NR | 5,400 | TCEQ Short-Term AMCV Health | 0.00003 |
| HEXENES* | 592-41-6 | 14,654 | 9.48 | 11,127 | 0.69 | 0.84 | NR | 500 | TCEQ Short-Term AMCV Health | 0.00168 |
| HYDROGEN CYANIDE | 74-90-8 | 14,654 | 1.47 | 11,127 | 0.12 | 0.24 | 2,000 | 308 | OEHHA Acute REL | 0.00079 |
| HYDROGEN SULFIDE | 7783-06-4 | 14,654 | 0.65 | 11,127 | 0.24 | 0.29 | 510 | 70 | ATSDR Acute MRL | 0.00410 |
| ISOPRENE | 78-79-5 | 14,654 | 1.06 | 11,127 | 0.22 | 0.24 | NR | 1,400 | TCEQ Short-Term AMCV Health | 0.00017 |
| METHANOL | 67-56-1 | 14,654 | 4.81 | 11,127 | 2.71 | 2.78 | 530,000 | 21,366 | OEHHA Acute REL | 0.00013 |
| METHYLCYCLOHEXANE | 108-87-2 | 14,654 | 0.17 | 11,127 | 0.04 | 0.05 | NR | 4,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| NONANES | 111-84-2 | 14,654 | 0.06 | 11,127 | 0.02 | 0.02 | NR. | 3,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| OCTANES* | 111-65-9 | 14,654 | 0.10 | 11,127 | 0.03 | 0.03 | NR | 4,100 | TCEQ Short-Term AMCV Health | 0.00001 |
| PENTANES* | 109-66-0 | 14,654 | 0.46 | 11,127 | 0.22 | 0.22 | NR | 68,000 | TCEQ Short-Term AMCV Health | 0.00000 |
| PROPYLENE | 115-07-1 | 14,654 | 4.72 | 11,127 | 0.28 | 0.30 | NR | NA | NE | |
| STYRENE | 100-42-5 | 14,654 | 0.31 | 11,127 | 0.08 | 0.09 | 20,000 | 5,000 | ATSDR Acute MRL | 0.00002 |
| TETRACHLOROETHYLENE | 127-18-4 | 14,654 | 0.10 | 11,127 | 0.01 | 0.01 | 35,000 | 6 | ATSDR Acute MRL | 0.00131 |
| TOLUENE | 108-88-3 | 14,654 | 17.67 | 11,127 | 0.38 | 0.49 | 67,000 | 2,000 | ATSDR Acute MRL | 0.00025 |
| TRIMETHYLBENZENES* | 622-96-8 | 14,654 | 4.06 | 11,127 | 0.19 | 0.23 | 50,000 | 250 | TCEQ Short-Term AMCV Health | 0.00093 |
| UNDECANES | 1120-21-4 | 14,654 | 0.06 | 11,127 | 0.03 | 0.03 | NR | 550 | TCEQ Short-Term AMCV Health | 0.00005 |
| XYLENES* | 1330-20-7 | 14,654 | 21.97 | 11,127 | 0.73 | 0.88 | 130,000 | 2,000 | ATSDR Acute MRL | 0.00044 |
| | | | | | | | | | Hazard Index | 0.04036 |

NR= According to EPA, AEGL is "not recommended due to insufficient data" NA= Not Available NC= Not Calculated



Mobile Laboratory Sampling Data Summary and Risk Assessment Western Hills Neighborhood | November 14, 2022

| 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 | 14,874 | 3.04 | 11,347 | 0.02 | 0.03 | 670,000 | 298 | OEHHA Acute REL | 0.00010 |
| ACETYLENE | 74-86-2 | 14,874 | 0.87 | 11,347 | 0.17 | 0.23 | NR | 25,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| BENZENE | 71-43-2 | 14,874 | 6.47 | 11,347 | 0.14 | 0.20 | 52,000 | 9 | ATSDR Acute MRL | 0.02214 |
| BUTANES* | 75-28-5 | 14,874 | 34.48 | 11,347 | 2.60 | 2.88 | NR | 33000 | TCEQ Short-Term AMCV Health | 0.00009 |
| BUTENES* | 590-18-1 | 14,874 | 17.85 | 11,347 | 1.78 | 2.04 | NR | 15000 | TCEQ Short-Term AMCV Health | 0.00014 |
| CARBON DISULFIDE | 75-15-0 | 14,874 | 0.25 | 11,347 | 0.00 | 0.00 | 13,000 | 1,991 | OEHHA Acute REL | 0.00000 |
| CYCLOPENTANES* | 287-92-3 | 14,874 | 22.85 | 11,347 | 2.82 | 3.14 | NR | 5,900 | TCEQ Short-Term AMCV Health | 0.00053 |
| DECANES | 124-18-5 | 14,874 | 0.12 | 11,347 | 0.05 | 0.05 | NR | 1,000 | TCEQ Short-Term AMCV Health | 0.00005 |
| DIETHYLBENZENES* | 141-93-5 | 14,874 | 0.20 | 11,347 | 0.09 | 0.11 | NR | 450 | TCEQ Short-Term AMCV Health | 0.00024 |
| DIMETHYLCYCLOHEXANES* | 638-04-0 | 14,874 | 0.24 | 11,347 | 0.15 | 0.15 | NR | 4,000 | CDPHE | 0.00004 |
| DODECANES | 112-40-3 | 14,874 | 4.44 | 11,347 | 0.01 | 0.02 | NR | 1720 | CDPHE | 0.00001 |
| ETHYLENE | 74-85-1 | 14,874 | 50.42 | 11,347 | 5.80 | 6.03 | NR | 500,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| HEPTANES* | 142-82-5 | 14,874 | 0.13 | 11,347 | 0.02 | 0.02 | NR | 8,300 | TCEQ Short-Term AMCV Health | 0.00000 |
| HEXANES* | 110-54-3 | 14,874 | 0.60 | 11,347 | 0.12 | 0.12 | NR | 5,400 | TCEQ Short-Term AMCV Health | 0.00002 |
| HEXENES* | 592-41-6 | 14,874 | 12.73 | 11,347 | 0.56 | 0.69 | NR | 500 | TCEQ Short-Term AMCV Health | 0.00138 |
| HYDROGEN CYANIDE | 74-90-8 | 14,874 | 0.68 | 11,347 | 0.10 | 0.20 | 2,000 | 308 | OEHHA Acute REL | 0.00065 |
| HYDROGEN SULFIDE | 7783-06-4 | 14,874 | 0.64 | 11,347 | 0.21 | 0.26 | 510 | 70 | ATSDR Acute MRL | 0.00369 |
| ISOPRENE | 78-79-5 | 14,874 | 2.88 | 11,347 | 0.15 | 0.18 | NR | 1,400 | TCEQ Short-Term AMCV Health | 0.00013 |
| METHANOL | 67-56-1 | 14,874 | 5.97 | 11,347 | 4.28 | 4.37 | 530,000 | 21,366 | OEHHA Acute REL | 0.00020 |
| METHYLCYCLOHEXANE | 108-87-2 | 14,874 | 0.31 | 11,347 | 0.05 | 0.05 | NR | 4,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| NONANES | 111-84-2 | 14,874 | 0.19 | 11,347 | 0.02 | 0.02 | NR | 3,000 | TCEQ Short-Term AMCV Health | 0.00001 |
| OCTANES* | 111-65-9 | 14,874 | 0.39 | 11,347 | 0.09 | 0.09 | NR | 4,100 | TCEQ Short-Term AMCV Health | 0.00002 |
| PENTANES* | 109-66-0 | 14,874 | 0.39 | 11,347 | 0.11 | 0.11 | NR | 68,000 | TCEQ Short-Term AMCV Health | 0.00000 |
| PROPYLENE | 115-07-1 | 14,874 | 3.53 | 11,347 | 0.25 | 0.30 | NR | NA | NE | |
| STYRENE | 100-42-5 | 14,874 | 1.38 | 11,347 | 0.08 | 0.09 | 20,000 | 5,000 | ATSDR Acute MRL | 0.00002 |
| TETRACHLOROETHYLENE | 127-18-4 | 14,874 | 0.08 | 11,347 | 0.00 | 0.01 | 35,000 | 6 | ATSDR Acute MRL | 0.00101 |
| TOLUENE | 108-88-3 | 14,874 | 20.91 | 11,347 | 0.75 | 0.94 | 67,000 | 2,000 | ATSDR Acute MRL | 0.00047 |
| TRIMETHYLBENZENES* | 622-96-8 | 14,874 | 4.30 | 11,347 | 0.21 | 0.28 | 50,000 | 250 | TCEQ Short-Term AMCV Health | 0.00114 |
| UNDECANES | 1120-21-4 | 14,874 | 0.07 | 11,347 | 0.03 | 0.04 | NR | 550 | TCEQ Short-Term AMCV Health | 0.00006 |
| XYLENES* | 1330-20-7 | 14,874 | 21.48 | 11,347 | 0.91 | 1.14 | 130,000 | 2,000 | ATSDR Acute MRL | 0.00057 |
| | | | | | | | | | Hazard Index | 0.03275 |

NR= According to EPA, AEGL is "not recommended due to insufficient data" NA= Not Available NC= Not Calculated



APPENDIX D CALIBRATION AND QA/QC DATA



| | | Initial Inst | trument Calibration an | d Verification | | |
|------------|--------|------------------|------------------------|----------------|--------------|-----------|
| | | Calibration | Calibration Value | Response | Difference | |
| Date | Time | Gas Component | (ppb v) | (ppb v) | (% of value) | Pass/Fail |
| | | | | | | |
| 11/13/2022 | 12:53 | Benzene | 250 | 225 | -10.0 | Pass |
| | | Toluene | 250 | 228 | -8.8 | Pass |
| | | Xylenes | 500 | 475 | -5.0 | Pass |
| | 42.50 | | 50 | 50.0 | | D |
| | 12:59 | Benzene | 50 | 50.8 | 1.6 | Pass |
| | • | Toluene | 50 | 49.9 | -0.2 | Pass |
| | | Ayleries | 100 | 110 | 10.0 | PdSS |
| | 1.04 | Benzene | 20 | 18.9 | -5.5 | Pass |
| | 1.04 | Toluene | 20 | 19.8 | -1.0 | Pass |
| | | Xvlenes | 40 | 39.7 | -0.7 | Pass |
| | | | | | | |
| | 13:06 | Benzene | 5 | 5.4 | 8.0 | Pass |
| | | Toluene | 5 | 5.5 | 10.0 | Pass |
| | | Xylenes | 10 | 10.4 | 4.0 | Pass |
| | | | | | | |
| | 13:23 | Ethylene | 100 | 98 | -2.0 | Pass |
| | | Propylene | 100 | 101 | 1.0 | Pass |
| | | 1-Butene | 100 | 99 | -1.0 | Pass |
| | | 1-Pentene | 100 | 103 | 3.0 | Pass |
| | | 1-Hexene | 100 | 98 | -2.0 | Pass |
| | | 1,3-Butadiene | 100 | 98 | -2.0 | Pass |
| | | | | | | |
| | 13:27 | Ethylene | 50 | 54.6 | 9.2 | Pass |
| | | Propylene | 50 | 51.8 | 3.6 | Pass |
| | | 1-Butene | 50 | 52.3 | 2.2 | Pass |
| | | 1-Pentene | 50 | 51.1 | 6.8 | Pass |
| | | 1-Hexene | 50 | 53.4 | 3.2 | Pass |
| | | 1,3-Butadiene | 50 | 51.6 | 3.2 | Pass |
| | 12.22 | Ethylopo | 10 | 11.2 | 12.0 | Pacc |
| | 15.52 | Bropylopo | 10 | 11.2 | 2.0 | Pass |
| | | 1-Butene | 10 | 10.2 | 2.0 | Pass |
| | | 1-Pentene | 10 | 10.7 | 3.0 | Pass |
| | | 1-Hexene | 10 | 10.4 | 4.0 | Pass |
| | | 1.3-Butadiene | 10 | 10.1 | 1 | Pass |
| | | | | | | |
| | 13:49 | HCN | 50 | 49.7 | -0.6 | Pass |
| | 13:51 | HCN | 25 | 26.9 | 7.6 | Pass |
| | 13:54 | HCN | 10 | 10.1 | 1.0 | Pass |
| | | | | | | |
| | 14:11 | H ₂ S | 125 | 122 | -2.4 | Pass |
| | 14:14 | H ₂ S | 25 | 25.5 | 2.0 | Pass |
| | 14:16 | H ₂ S | 10 | 10.1 | 1.0 | Pass |
| | 14:18 | H ₂ S | 5 | 5.12 | 2.4 | Pass |
| | | | | | | |
| İ | 14:29 | Propane | 500 | 482 | -3.6 | Pass |
| | | Butane | 500 | 465 | -7.0 | Pass |
| | | Pentane | 500 | 475 | -5.0 | Pass |
| | | Hexane | 500 | 451 | -9.8 | Pass |
| | | Heptane | 500 | 459 | -8.2 | Pass |
| | | | | | | |
| | 14:31 | Propane | 250 | 242 | -3.2 | Pass |
| | | Butane | 250 | 248 | -0.8 | Pass |
| | | Pentane | 250 | 225 | -10.0 | Pass |
| | | Hexane | 250 | 228 | -8.8 | Pass |
| | | пертапе | 250 | 242 | -3.2 | Pass |
| | 1/1-22 | Pronano | 100 | ۵۵ | -20 | Dace |
| | 14.33 | Butane | 100 | 30 | -2.0 | Pace |
| | | Pentane | 100 | Q1 | -10.0 | Pass |
| | | Hexane | 100 | 93 | -7.0 | Pass |
| | | Heptane | 100 | 91 | -9.0 | Pass |
| | | Teptune | 100 | J. | 5.0 | 1 4 5 5 |
| | 14:36 | Propane | 25 | 24.3 | -2.8 | Pass |
| | | Butane | 25 | 23.7 | -5.2 | Pass |
| | | Pentane | 25 | 23.9 | -4.4 | Pass |
| | | Hexane | 25 | 24.2 | -3.2 | Pass |
| | | Heptane | 25 | 22.1 | -11.6 | Pass |



| | | | Instrument Calibratio | on Check | | |
|------------|-------|------------------|--------------------------|----------|--------------|-----------|
| | | Calibration | Calibration Value | Response | Difference | |
| Date | Time | Gas Component | (ppb v) | (ppb v) | (% of value) | Pass/Fail |
| 11/14/2022 | 8:09 | Ethylene | 50 | 50.8 | 1.6 | Pass |
| | | Propylene | 50 | 51.9 | 3.8 | Pass |
| | | 1-Butene | 50 | 47.2 | -5.6 | Pass |
| | | 1-Pentene | 50 | 50.4 | 0.8 | Pass |
| | | 1-Hexene | 50 | 48.3 | -3.4 | Pass |
| | | 1,3-Butadiene | 50 | 51.7 | 3.4 | Pass |
| | | | | | | |
| | 8:15 | Benzene | 100 | 103 | 3.0 | Pass |
| | | Toluene | 100 | 102 | 2.0 | Pass |
| | | Xylenes | 200 | 213 | 6.5 | Pass |
| | | Benzene | 10 | 9.86 | -1.4 | Pass |
| | | Toluene | 10 | 9.42 | -5.8 | Pass |
| | | Xylenes | 20 | 19.3 | -3.5 | Pass |
| | | | | | | |
| | 8:35 | HCN | 25 | 26.4 | 5.6 | Pass |
| | | | | | | |
| | 8:44 | H ₂ S | 100 | 98.6 | -1.4 | Pass |
| | 8:48 | H ₂ S | 20 | 20.1 | 0.5 | Pass |
| | | | | | | |
| | 8:51 | Propane | 150 | 147 | -2.0 | Pass |
| | | Butane | 150 | 136 | -9.3 | Pass |
| | | Pentane | 150 | 139 | -7.3 | Pass |
| | | Hexane | 150 | 141 | -6.0 | Pass |
| | | Heptane | 150 | 143 | -4.7 | Pass |
| | | | | | | |
| | | | | | | |
| | 15:12 | HCN | 25 | 25.5 | 2.0 | Pass |
| | | | | | | |
| | 15:23 | H ₂ S | 20 | 20.5 | 2.5 | Pass |
| | | | | | | |
| | 15:25 | Propane | 150 | 142 | -5.3 | Pass |
| | | Butane | 150 | 143 | -4.7 | Pass |
| | | Pentane | 150 | 139 | -7.3 | Pass |
| | | Hexane | 150 | 139 | -7.3 | Pass |
| | | Heptane | 150 | 137 | -8.7 | Pass |
| | | | | | | |
| | 15:29 | Benzene | 100 | 103 | 3.0 | Pass |
| | | Toluene | 100 | 97.6 | -2.4 | Pass |
| | | Xylenes | 200 | 203 | 1.5 | Pass |
| | | | | | | |
| | 15:31 | Ethylene | 50 | 53.1 | 6.2 | Pass |
| | | Propylene | 50 | 48 | -4.0 | Pass |
| | | 1-Butene | 50 | 44.1 | -11.8 | Pass |
| | | 1-Pentene | 50 | 45.7 | -8.6 | Pass |
| | | 1-Hexene | 50 | 46.7 | -6.6 | Pass |
| | | 1,3-Butadiene | 50 | 44.1 | -11.8 | Pass |
| | | | | | | |



| | | - | Instrument Calibratio | n Check | | |
|------------|-------|------------------|-----------------------|----------|--------------|-----------|
| | | Calibration | Calibration Value | Response | Difference | |
| Date | Time | Gas Component | (ppb v) | (ppb v) | (% of value) | Pass/Fail |
| 11/15/2022 | 11:08 | Ethylene | 50 | 47.3 | -5.4 | Pass |
| | | Propylene | 50 | 53.3 | 6.6 | Pass |
| | | 1-Butene | 50 | 48 | -4.0 | Pass |
| | | 1-Pentene | 50 | 47.2 | -5.6 | Pass |
| | | 1-Hexene | 50 | 52 | 4.0 | Pass |
| | | 1,3-Butadiene | 50 | 49.8 | -0.4 | Pass |
| | | | | | | |
| | 11:10 | Benzene | 100 | 102 | 2.0 | Pass |
| | | Toluene | 100 | 98.4 | -1.6 | Pass |
| | | Xylenes | 200 | 206 | 3.0 | Pass |
| | 11:17 | Benzene | 20 | 19.1 | -4.5 | Pass |
| | | Toluene | 20 | 19.3 | -3.5 | Pass |
| | | Xylenes | 40 | 38.3 | -4.3 | Pass |
| | | | | | | |
| | 11:20 | HCN | 25 | 26 | 4.0 | Pass |
| | | | | | | |
| | 11:29 | H ₂ S | 100 | 97.8 | -2.2 | Pass |
| | 11:31 | H ₂ S | 20 | 21.6 | 8.0 | Pass |
| | | | | | | |
| | 11:33 | Propane | 150 | 139 | -7.3 | Pass |
| | | Butane | 150 | 142 | -5.3 | Pass |
| | | Pentane | 150 | 142 | -5.3 | Pass |
| | | Hexane | 150 | 149 | -0.7 | Pass |
| | | Heptane | 150 | 139 | -7.3 | Pass |
| | | | | | | |
| | | | | | | _ |
| | 17:13 | HCN | 25 | 25.1 | 0.4 | Pass |
| | | | | | | |
| | 17:22 | H ₂ S | 50 | 46.7 | -6.6 | Pass |
| | | | | | | |
| | 17:24 | Propane | 150 | 145 | -3.3 | Pass |
| | | Butane | 150 | 142 | -5.3 | Pass |
| | | Pentane | 150 | 138 | -8.0 | Pass |
| | | Hexane | 150 | 141 | -6.0 | Pass |
| | | Heptane | 150 | 147 | -2.0 | Pass |
| | | - | 105 | | | |
| | 17:27 | Benzene | 100 | 99.4 | -0.6 | Pass |
| | | loluene | 100 | 97.3 | -2.7 | Pass |
| | | Xylenes | 200 | 196 | -2.0 | Pass |
| | 45.00 | Ealer 1 | F0 | 56.2 | 12.0 | D- |
| | 15:29 | Etnylene | 50 | 56.3 | 12.6 | Pass |
| | | Propylene | 50 | 54.3 | 8.6 | Pass |
| | | T-Butene | 50 | 52.2 | 4.4 | Pass |
| | | 1-Pentene | 50 | 51 | 2.0 | Pass |
| | | 1.2 Pute diana | 50 | 54.5 | 9.0 | Pass |
| | | T'2-RALAGIENE | 50 | 53.4 | δ.σ | Pa55 |
| | | | | | | |



| | | | Instrument Calibratio | on Check | | |
|------------|-------|------------------|-----------------------|----------|--------------|-----------|
| | | Calibration | Calibration Value | Response | Difference | |
| Date | Time | Gas Component | (ppb v) | (ppb v) | (% of value) | Pass/Fail |
| 11/16/2022 | 8:52 | Ethylene | 50 | 51.5 | 3.0 | Pass |
| | | Propylene | 50 | 55.9 | 11.8 | Pass |
| | | 1-Butene | 50 | 49.1 | -1.8 | Pass |
| | | 1-Pentene | 50 | 52.6 | 5.2 | Pass |
| | | 1-Hexene | 50 | 55.5 | 11.0 | Pass |
| | | 1,3-Butadiene | 50 | 51.3 | 2.6 | Pass |
| | | | | | | |
| | 9:04 | Benzene | 100 | 102 | 2.0 | Pass |
| | | Toluene | 100 | 99 | -1.0 | Pass |
| | | Xylenes | 200 | 210 | 5.0 | Pass |
| | 9:15 | Benzene | 20 | 19.2 | -4.0 | Pass |
| | | Toluene | 20 | 18.3 | -8.5 | Pass |
| | | Xylenes | 40 | 38.9 | -2.8 | Pass |
| | | | | | | |
| | 8:59 | HCN | 25 | 25.5 | 2.0 | Pass |
| | | | | | | |
| | 9:24 | H ₂ S | 100 | 99.9 | -0.1 | Pass |
| | 9:26 | | 20 | 19.4 | -3.0 | Pass |
| | | | | | | |
| | 8:48 | Propane | 150 | 151 | 0.7 | Pass |
| | | Butane | 150 | 139 | -7.3 | Pass |
| | | Pentane | 150 | 153 | 2.0 | Pass |
| | | Hexane | 150 | 146 | -2.7 | Pass |
| | | Heptane | 150 | 152 | 1.3 | Pass |
| | | | | | | |
| | | | | | | |
| | 15:28 | HCN | 25 | 26.4 | 5.6 | Pass |
| | | | | | | |
| | 15:36 | H ₂ S | 50 | 49.1 | -1.8 | Pass |
| | | | | | | |
| | 15:39 | Propane | 150 | 139 | -7.3 | Pass |
| | | Butane | 150 | 135 | -10.0 | Pass |
| | | Pentane | 150 | 132 | -12.0 | Pass |
| | | Hexane | 150 | 140 | -6.7 | Pass |
| | | Heptane | 150 | 138 | -8.0 | Pass |
| | | | | | | |
| | 15:51 | Benzene | 100 | 105 | 5.0 | Pass |
| | | Toluene | 100 | 97.2 | -2.8 | Pass |
| | | Xylenes | 200 | 213 | 6.5 | Pass |
| | | | | | | |
| | 15:59 | Ethylene | 50 | 42.9 | -14.2 | Pass |
| | | Propylene | 50 | 47.6 | -4.8 | Pass |
| | | 1-Butene | 50 | 45.7 | -8.6 | Pass |
| | | 1-Pentene | 50 | 50.2 | 0.4 | Pass |
| | | 1-Hexene | 50 | 45.2 | -9.6 | Pass |
| | | 1,3-Butadiene | 50 | 52.7 | 5.4 | Pass |
| | | | | | | |



| | | | Instrument Calibrati | on Check | | |
|------------|-------|------------------|--------------------------|----------|--------------|-----------|
| | | Calibration | Calibration Value | Response | Difference | |
| Date | Time | Gas Component | (ppb v) | (ppb v) | (% of value) | Pass/Fail |
| 11/17/2022 | 9:45 | Ethylene | 50 | 50.1 | 0.2 | Pass |
| | | Propylene | 50 | 50.2 | 0.4 | Pass |
| | | 1-Butene | 50 | 46.2 | -7.6 | Pass |
| | | 1-Pentene | 50 | 50.9 | 1.8 | Pass |
| | | 1-Hexene | 50 | 47.7 | -4.6 | Pass |
| | | 1,3-Butadiene | 50 | 50.4 | 0.8 | Pass |
| | | | | | | |
| | 9:49 | Benzene | 100 | 100 | 0.0 | Pass |
| | | Toluene | 100 | 94.7 | -5.3 | Pass |
| | | Xylenes | 200 | 191 | -4.5 | Pass |
| | | | | | | |
| | 9:51 | HCN | 25 | 24.3 | -2.8 | Pass |
| | | | | | | |
| | 9:57 | H ₂ S | 100 | 97.3 | -2.7 | Pass |
| | | | | | | |
| | 9:59 | Propane | 150 | 142 | -5.3 | Pass |
| | | Butane | 150 | 146 | -2.7 | Pass |
| | | Pentane | 150 | 145 | -3.3 | Pass |
| | | Hexane | 150 | 137 | -8.7 | Pass |
| | | Heptane | 150 | 146 | -2.7 | Pass |
| | | | | | | |
| | | | | | | |
| | 17:08 | HCN | 25 | 22.9 | -8.4 | Pass |
| | | | | | | |
| | 17:12 | H ₂ S | 50 | 51.4 | 2.8 | Pass |
| | | | | | | |
| | 17:15 | Propane | 150 | 141 | -6.0 | Pass |
| | | Butane | 150 | 138 | -8.0 | Pass |
| | | Pentane | 150 | 145 | -3.3 | Pass |
| | | Hexane | 150 | 137 | -8.7 | Pass |
| | | Heptane | 150 | 149 | -0.7 | Pass |
| | | | | | | |
| | 17:18 | Benzene | 100 | 99.6 | -0.4 | Pass |
| | | Toluene | 100 | 94.9 | -5.1 | Pass |
| | | Xylenes | 200 | 198 | -1.0 | Pass |
| | | | | | | |
| | 17:21 | Ethylene | 50 | 48.8 | -2.4 | Pass |
| | | Propylene | 50 | 48.1 | -3.8 | Pass |
| | | 1-Butene | 50 | 45.9 | -8.2 | Pass |
| | | 1-Pentene | 50 | 46.7 | -6.6 | Pass |
| | | 1-Hexene | 50 | 47.5 | -5.0 | Pass |
| | | 1,3-Butadiene | 50 | 48.2 | -3.6 | Pass |
| | | | | | | |



| | | | Instrument Calibration | on Check | | |
|------------|-------|------------------|--------------------------|----------|--------------|-----------|
| | | Calibration | Calibration Value | Response | Difference | |
| Date | Time | Gas Component | (ppb v) | (ppb v) | (% of value) | Pass/Fail |
| 11/18/2022 | 8:58 | Ethylene | 50 | 51.8 | 3.6 | Pass |
| | | Propylene | 50 | 55.7 | 11.4 | Pass |
| | | 1-Butene | 50 | 52.7 | 5.4 | Pass |
| | | 1-Pentene | 50 | 51.8 | 3.6 | Pass |
| | | 1-Hexene | 50 | 51.1 | 2.2 | Pass |
| | | 1,3-Butadiene | 50 | 51.6 | 3.2 | Pass |
| | | | | | | |
| | 9:02 | Benzene | 100 | 104 | 4.0 | Pass |
| | | Toluene | 100 | 98.1 | -1.9 | Pass |
| | | Xylenes | 200 | 208 | 4.0 | Pass |
| | 9:07 | Benzene | 20 | 19.3 | -3.5 | Pass |
| | | Toluene | 20 | 19.2 | -4.0 | Pass |
| | | Xylenes | 40 | 38.4 | -4.0 | Pass |
| | | | | | | |
| | 9:09 | HCN | 25 | 24.3 | -2.8 | Pass |
| | | | | | | |
| | 9:17 | H ₂ S | 100 | 99.9 | -0.1 | Pass |
| | 9:20 | | 20 | 20.9 | 4.5 | Pass |
| | | | | | | |
| | 9:23 | Propane | 150 | 149.1 | -0.6 | Pass |
| | | Butane | 150 | 143.4 | -4.4 | Pass |
| | | Pentane | 150 | 147.1 | -1.9 | Pass |
| | | Hexane | 150 | 146.2 | -2.5 | Pass |
| | | Heptane | 150 | 151.2 | 0.8 | Pass |
| | | | | | | |
| | | | | | | |
| | 14:04 | HCN | 25 | 25.4 | 1.6 | Pass |
| | | | | | | |
| | 14:08 | H ₂ S | 50 | 48.1 | -3.8 | Pass |
| | | | | | | |
| | 14:10 | Propane | 150 | 142 | -5.3 | Pass |
| | | Butane | 150 | 144 | -4.0 | Pass |
| | | Pentane | 150 | 139 | -7.3 | Pass |
| | | Hexane | 150 | 148 | -1.3 | Pass |
| | | Heptane | 150 | 154 | 2.7 | Pass |
| | | | | | | |
| | 14:14 | Benzene | 100 | 99.8 | -0.2 | Pass |
| | | Toluene | 100 | 95.4 | -4.6 | Pass |
| | | Xylenes | 200 | 197 | -1.5 | Pass |
| | | | | | | |
| | 14:16 | Ethylene | 50 | 52.4 | 4.8 | Pass |
| | | Propylene | 50 | 46.4 | -7.2 | Pass |
| | | 1-Butene | 50 | 43.8 | -12.4 | Pass |
| | | 1-Pentene | 50 | 47.2 | -5.6 | Pass |
| | | 1-Hexene | 50 | 46.9 | -6.2 | Pass |
| | | 1,3-Butadiene | 50 | 50.1 | 0.2 | Pass |
| | | | | | | |



Suncor 4th Quarter PTR Parameters 11/13 – 11/18/22 Initial Calibration Checks

| <u> </u> | | • • | | |
|----------------------------------|--|--|--|--|
| Setting | Odor | V 1 | | |
| Primary Ion | H3O+ | × 🥖 | | |
| ansmission | DC | 🖉 | | |
| | Man/Ctrl | Ctrl | | |
| PC | 349.9 🚔 | 349.90 mbar | | |
| p Drift | 2.30 📮 | 2.29 mbar | | |
| TofLens | | 4.70E-5 mbar | | |
| TOF | | 1.03E-6 mbar | | |
| E/N | | 120 Td | | |
| Temps | 80.20 °C | 80.10 °C | | |
| SrcValve | 50.0 | | | |
| H2O | 6.0 | 6.00 sccm | | |
| 02 | 0.0 | 0.00 sccm | | |
| NO | 0.0 | 0.00 sccm | | |
| lhc | 4 | 4.0 mA | | |
| | On/Off | On | | |
| FCinlet | 60.0 | 59.97 sccm | | |
| J FU | ℃ 🕞 | D* | | |
| Us | 150 🚍 | 145.0 V | | |
| Uso | 80 🚍 | 78.6 V | | |
| Udrift | 525 | 526.1 V | | |
| | | | | |
| | 0 | | | |
| | | | | |
| Hex1 | | CP | | |
| Hex1 OFF | | COP CON | | |
| Hex1 Freque | 1 — 1 ➡ /ON ☑ ency 6.00 | OP ON €.00Mhz | | |
| Hex1 Freque Ampli | 1 — /ON ☑ ency 6.00 tude 95.0 | OP OP ON € 57.1V | | |
| Hex1 OFF, Freque Offset | I /ON ✓ ency 6.00 tude 95.0 - 0.70 | OP ON 6.00Mhz 57.1V → -0.67V | | |

Production Settings



| Lens 1 | 14.0 | \$ | 14.0 V | | All on 🖂 |
|------------|--------|----|----------|-----------|----------|
| Lens 2 | 30.0 | + | 30.0 V | | Lenses 🖂 |
| 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 | • | 17.0 V | | |
| Push L | 16.5 | + | 16.0 V | | 3 mA |
| Push H | 790.0 | + | 790.0 V | \square | 3 mA |
| Pull L | 80.0 | + | 80.0 V | \square | 3 mA |
| Pull H | 680.0 | \$ | 680.0 V | \square | 3 mA |
| Grid | 2400.0 | + | 2282.0 V | \square | 1 µA |
| Cage | 5020.0 | • | 4766 V | \square | 99 µA |
| Refl. Grid | 667.0 | * | 634.0 V | \square | 75 µA |
| efl. Back | 900.0 | • | 855.0 V | | 167 µA |
| MCP F | 5400 | + | 5134 V | \square | 17 µA |
| MCP B | 2570 | + | 2471 V | | 225 µA |

TOF Settings



| Acquisition | | | | AC | Q active |
|---|---|---|-------|--------|------------------|
| 6 8 | | | | | |
| Single Spec T | me (ms) | 1000 | ÷ | | |
| Extraction | time (µs) | 5.0 | - | 37 | 2.7 amu |
| max Flight | time(µs) | 32.0 | - | 3 | 1.25 kHz |
| Data Save Se | ettings | | | | |
| Spec 🖉 | \square | Trace | | | Raw |
| Time Duratio | in | | | | × . |
| 02:00:00 💠 🥸 | Single File | e Duratio | n. | | |
| 12 🔤 1 | Vumber | of Files To | s Sto | re | |
| C:\lonicon\c | lata | | | | Contr. |
| Add File C | ount Ext for new f | ension file | | | |
| <year>_<mo Data_<hour></hour></mo </year> | onth>_ <d •_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond</td><td></td><td>1</td></sec<></td></minut<></d | lay>\ te>_ <sec< td=""><td>ond</td><td></td><td>1</td></sec<> | ond | | 1 |
| 2022_11_12\[|)ata_18_0 | 6_25_par | t_XX | × | |
| | libration | | | | |
| Mass Axis Ca | | | | | 1.41 |
| Mass Axis Ca | 👗 Cal | E | 2 | 30 sec | |
| Mass Axis Ca 🥶 日 峙 Mass | Cal TimeBin | | 2 | 30 sec | |
| Mass Axis Ca Mass 21.0220 | Cal TimeBin 15984 | | 2 | a a | 15008 |
| Mass Axis Ca Mass 21.0220 203.9400 | Cal TimeBin 15984 161505 |] 1 1 | | a b | 15008 52822.1 |

Acquisition Settings

| 🛞 H | 100 | | × |
|-----------|------|---|---------|
| RBB | ł | | |
| Hex1 | | | OP |
| OFF/ON | | | ON |
| Frequency | 6.00 | - | 6.00Mhz |
| Amplitude | 95.0 | - | 57.4V |
| Offset - | 0.70 | 4 | -0.67V |
| < | | | |

Hexapole Settings



11-14-22 Western Hills Neighborhood PTR Operating Parameters



Production Settings



| 1 | MCP | | | | 3 🕝 🗠 |
|----------|-----------|----------|----|--------|------------|
| All on 🖂 | | 15.0 V | \$ | 14.0 | Lens 1 |
| Lenses 🗹 | | 30.0 V | * | 30.0 | Lens 2 |
| | | 20.0 V | - | 20.0 | Lens 3 |
| | | 60.0 V | - | 60.0 | Lens 4 |
| | | 69.0 V | + | 70.0 | Lens 5 |
| | | 80.0 V | ÷ | 80.0 | Lens 6 |
| | | 17.0 V | ÷ | 17.0 | Lens 7 |
| 3 mA | | 16.0 V | \$ | 16.5 | Push L |
| 2 mA | | 790.0 V | * | 790.0 | Push H |
| 3 mA | | 80.0 V | ÷ | 80.0 | Pull L |
| 3 mA | | 680.0 V | - | 680.0 | Pull H |
| 1 µA | | 2283.0 V | ÷ | 2400.0 | Grid |
| 99 µA | \square | 4768 V | * | 5020.0 | Cage |
| 75 µA | | 634.0 V | + | 667.0 | Refl. Grid |
| 167 µA | | 855.0 V | \$ | 900.0 | lefl. Back |
| 17 µA | | 5134 V | * | 5400 | MCP F |
| 221 µA | | 2461 V | \$ | 2570 | MCP B |

TOF Voltages



| Acquisition | | | | A | CQ act | ive |
|--|--|--|--------|-----|----------|-----|
| | 1 | | | | | |
| Single Spec | Time (ms) | 1000 | * | | | |
| Extraction | n time (µs) | 5.0 | * | | 372.9 ar | nu |
| max Flig | httime(µs) | 32.0 | * | | 31.25 k | Hz |
| Data Save | Settings | | | | | |
| Spec 🗹 | | Trace | | E | Raw | |
| Time Durat | tion | | | | \sim | |
| 02:00:00 🖨 | Single File | e Duratio | n | | | |
| 12 🗘 | Number | of Files T | o Stoi | e | | |
| C:\lonicon | \data | | | | 6 | |
| Add File | Count Ext Q for new 1 | ension file | | | | |
| <year>_<n Data_<hou< td=""><td>nonth>_<c ir>_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond></td><td></td><td>22</td><td></td></sec<></td></minut<></c </td></hou<></n </year> | nonth>_ <c ir>_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond></td><td></td><td>22</td><td></td></sec<></td></minut<></c | lay>\ te>_ <sec< td=""><td>ond></td><td></td><td>22</td><td></td></sec<> | ond> | | 22 | |
| 2022_11_13 | \Data_21_0 | 9_30_par | t_XX) | C | | |
| Mass Axis (| alibration | | | | | |
| 💕 🖬 🖬 | Cal | | 3 | 0 s | ec | ÷ |
| Mass | TimeBin | | | | | |
| 21.0220 | 15956 | Ċ | ^ | а | 1500 | 5.5 |
| | 161452 | 前 | i l | b | -52838 | 3.7 |
| 203.9400 | 101436 | 2.000 | 1 | | | |

Acquisition Settings

| 🛞 H | | | \times |
|-----------|------|---|----------|
| i A B B | 4 | | |
| Hex1 | | | OP |
| OFF/ON | | | ON |
| Frequency | 6.00 | * | 6.00Mhz |
| Amplitude | 95.0 | + | 59.1V |
| Offset - | 0.70 | 4 | -0.67V |

Hexapole Settings



| 7 55 . 7 | | 21.022 *(H2O)H+ | 6.78E+7 | ccps |
|-----------|---------|------------------------|---------|--------|
| /.JE+/- | Ma | 28.015 (HCN)H+ | 2.85E+4 | ccps |
| 7E+7- | | 28.026 (C2H4)+ | 1.41E+4 | ccps |
| 6.5E+7- | | 34.995 (H2S)H+ | 2.91E+3 | ccps |
| 6E+7- | | 43.047 Propylene[C3H6 | 794 | ccps |
| | | 55.054 [1,3 BDE] | 319 | ccps |
| 5.5C+7- | | 57.070 Butenes | 695 | ccps |
| 5E+7- | | 79.054 Benzene | 22.5 | ccps |
| 4.5E+7- | <u></u> | 86.090 [13BDE][O2]+ | 10.1 | ccps |
| 4E+7- | | 93.070 Toluene | 146 | ccps |
| | | 105.07 Styrene | 50.6 | ccps |
| 3.5E+ / - | | 107.11 Xylenes, EB | 262 | ccps |
| 3E+7- | | 44.040 Propane[O2] | 6.75E+3 | 3 ccps |
| 2.5E+7- | | 58.050 Butanes[O2] | 1.46E+3 | ccps |
| 2E+7- | | 71.086 Pentenes[H+] | 468 | ccps |
| | | 72.080 Pentanes[O2] | 1.81E+3 | s ccps |
| 1.5E+7- | | 85.105 Hexenes[H+] | 469 | ccps |
| 1E+7- | | 86.090 Hexanes[O2] | 1.12E+3 | 3 ccps |
| 5E+6- | | 100.12 Heptanes[O2] | 915 | ccps |
| 0- | | 166.85 Tetrachloroethy | 6.32 | ccps |

757555.0 AM 75820.0 AM 75845.0 AM 75935.0 AM 75935.0 AM 80025.0 AM 80025.0 AM 80025.0 AM 801:15.0 AM 80140.0 AM 80205.0 AM 80235.0 AM 80225.0 AM 80325.0 AM 80325.0 AM 80345.0 AM 804:10 AM 80435.0 AM 11/14/2022 11/14/202

Hydronium Ion Trace



Western Hills



End Western Hills



PTR Operating Parameters 11-15-22 Dupont

| | l | | 0 | \odot |
|--|--|--------|--------------------------------|----------------------|
| Setting | Odor | | ~ | 1 |
| rimary lor | H3O+ | | ~ | 9 |
| nsmission | n DC | | ~ | 9 |
| | Man/Ct | trl | Ctrl | |
| PC | 350.9 | | 350.97 m | bar |
| p Drift | t 2.30 | 4 | 2.30 m | bar |
| TofLens | s | in and | 4.70E-5 m | bar |
| TOF | - | | 1.00E-6 m | bar |
| E/N | 1 | | 12 | 0 Td |
| Temp | s 80.10 | °C | 80.0 | 0 °C |
| SrcValve | e 50.0 | 1 | | |
| H2C | 6.0 | 9 | 6.01 sc | cm |
| 02 | 2 0.0 | | 0.00 sc | cm |
| NC | 0.0 | | 0.00 sc | cm |
| lh | c 4 | - | 4.0 | mA |
| | On/Of | f | On | |
| FCinle | t 60.0 | | 60.07 so | cm |
| FU | °C [| .> | D* | |
| Us | 150 | 3 | 145.0 | v |
| Uso | 80 | | 78.6 | V |
| Udrift | 525 | 3 | 526.1 | v |
| | | | | |
| | | | | |
| 8 | H — | | 12 | × |
| (| H — | | 12 | × |
| () Hext | H — | | 10 | × |
| (Second second s | H — 1 F/ON 🗹 | | 40 (| × |
| Hext Frequ | H — I F/ON 🗹 Jency 6.00 | | OF 1001 | × ⊃; |
| Hext OFI Frequ Amp | H — I F/ON 🗹 Jency 6.00 litude 95.0 | | € 6.00M € 56.1V | × A |
| Hext Frequ Amp Offse | H — F/ON 🗹 Jency 6.00 litude 95.0 t – 0.70 |) | € 6.00M 56.1V € -0.67 | × vi ilhz / |

Production Settings



| j 🙆 🗠 | | | MCP | a a |
|------------|----------|----------|-----------|----------|
| Lens 1 | 14.0 🖨 | 14.0 V | | All on 🖂 |
| Lens 2 | 30.0 韋 | 30.0 V | | Lenses 🔽 |
| Lens 3 | 20.0 ≑ | 20.0 V | | |
| Lens 4 | 60.0 韋 | 60.0 V | | |
| Lens 5 | 70.0 ≑ | 69.0 V | | |
| Lens 6 | 80.0 韋 | 80.0 V | | |
| Lens 7 | 17.0 韋 | 17.0 V | | |
| Push L | 16.5 韋 | 16.0 V | | 3 mA |
| Push H | 790.0 韋 | 790.0 V | | 3 mA |
| Pull L | 80.0 ≑ | 80.0 V | \square | 3 mA |
| Pull H | 680.0 韋 | 680.0 V | | 3 mA |
| Grid | 2400.0 🖨 | 2282.0 V | | 1 μA |
| Cage | 5020.0 ≑ | 4766 V | | 99 µA |
| Refl. Grid | 667.0 韋 | 634.0 V | | 75 µA |
| Refl. Back | 900.0 🖨 | 855.0 V | \square | 167 µA |
| MCP F | 5400 韋 | 5134 V | | 17 µA |
| MCP B | 2570 🖨 | 2473 V | M | 227 µA |

TOF Lens Voltages



| | | | | A | CQ activ |
|--|--|--|---------------------|--|---|
| | | | | | |
| Single Spec 1 | līme (ms) | 1000 | 4 | | |
| Extraction | time (µs) | 5.0 | 4 | | 372.4 am |
| max Fligh | ittime(µs) | 32.0 | * | | 31.25 kH |
| Data Save S | ettings | | | | |
| Spec 🖉 | | Trace | | E | Raw |
| Time Durati | on | | | | 4 |
| 02:00:00 🚖 | Single File | E Duratio | n | | |
| 12 🔤 | Number | of Files To | Sto | re | |
| C:\lonicon\ | data | | | | 6 |
| Add File | Count Ext | ension | | | |
| New ACC | I tor new 1 | ne | | | |
| New ACC <year>_<m Data_<hour< td=""><td>onth>_<c >_<minut< td=""><td>iay>\ te>_<sec< td=""><td>ond></td><td>2</td><td>62</td></sec<></td></minut<></c </td></hour<></m </year> | onth>_ <c >_<minut< td=""><td>iay>\ te>_<sec< td=""><td>ond></td><td>2</td><td>62</td></sec<></td></minut<></c | iay>\ te>_ <sec< td=""><td>ond></td><td>2</td><td>62</td></sec<> | ond> | 2 | 62 |
| New ACC <year>_<m Data_<hour 2022_11_15\</hour </m </year> | onth>_ <c >_<minul Data_07_4</minul </c | iay>\ te>_ <sec 1_06_par</sec | ond> t_XX) | (| 63 |
| New ACC <year>_<m Data_<hour 2022_11_15\ Mass Axis Ca</hour </m </year> | onth>_ <c onth>_<c >_<minul Data_07_4 alibration</minul </c </c | lay>\ te>_ <sec l1_06_par</sec | ond> t_XX) At | < | CAL done |
| New ACC <year>_<m Data_<hour 2022_11_15\ Mass Axis Ca</hour </m </year> | onth>_ <c >_<minut Data_07_4 alibration</minut </c | iay>\ te>_ <sec 1_06_par</sec | ond> t_XX) Au | (1 to(30 si | CAL done |
| New ACC <year>_<m Data_<hour 2022_11_15 Mass Axis Ca Mass Mass</hour </m </year> | onth>_ <c >_<minut Data_07_4 alibration</minut </c | iay>\ te>_ <sec 1_06_par</sec | ond> t_XX Au | (1 to(10 si | CAL done |
| New ACC <year>_<m Data_<hour 2022_11_15\ Mass Axis Ca Mass 21.0220</hour </m </year> | onth>_ <commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth>_<commonstants onth onth>_<commonstants onth onth onth onth onth onth onth onth</commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants </commonstants | iay>\ te>_ <sec I1_06_par</sec | ond> t_XX0 Au | (1 to (30 si a | CAL done ec 🗧 |
| New ACC <year>_<m Data_<hour 2022_11_15\ Mass Axis Ca Mass 21.0220 203.9400</hour </m </year> | onth>_ <c >_<minut Data_07_4 alibration Cal TimeBin 16021 161611</minut </c | iay>\ te>_ <sec 1_06_par</sec | ond> t_XXX Au | x Ito(30 si a b | CAL done ec 4 15015.7 |
| New ACC <year>_<m Data_<hour 2022_11_15\ Mass Axis Ca Mass 21.0220 203.9400 330.8500</hour </m </year> | onth>_ <c >_<minut Data_07_4 alibration Cal TimeBin 16021 161611 220295</minut </c | iay>\ te>_ <sec 11_06_par</sec | ond> t_XX0 Au | (100 (0 5) (0 5) | CAL done ec 4 15015.' -52817.9 |

Acquisition Settings





Dupont Neighborhood



11-16-2 PTR Parameters Pioneer Park

| | 5 | 00 | | | | |
|--|---|--|--|--|--|--|
| Setting | Odor | ~ / | | | | |
| Primary lor | n H3O+ | H3O+ V | | | | |
| ansmissior | DC | × 🥜 | | | | |
| | Man/Ctrl | Ctrl | | | | |
| PC | 352.0 | 352.00 mbar | | | | |
| p Drift | t 2.30 🖨 | 2.29 mbar | | | | |
| TofLens | 5 | 4.85E-5 mbar | | | | |
| TOF | - | 7.75E-7 mbar | | | | |
| E/N | 1 | 120 Td | | | | |
| Temp | s 80.10 °C | 80.00 °C | | | | |
| SrcValve | = 50.0 🚍 |] | | | | |
| H2C | 6.0 | 6.00 sccm | | | | |
| 02 | 2 0.0 🌲 | 0.00 sccm | | | | |
| NC | 0.0 | 0.00 sccm | | | | |
| Jh | c 4 | 4.0 mA | | | | |
| | On/Off | On | | | | |
| FCinle | t 60.0 | 60.00 sccm | | | | |
| J FU | °C □ | D* | | | | |
| Us | 150 🗐 | 145.0 V | | | | |
| Uso | 80 🚔 | 78.6 V | | | | |
| 11.000 | 525 | 526 1 V | | | | |
| Udrift | [(*) [| 520.11 | | | | |
| Udnft | H | | | | | |
| Udnft | H — | | | | | |
| Udnft () Hext | H — | | | | | |
| Udrift () Hext | H — I E/ON 🗹 | | | | | |
| Udrift | H — F/ON Juency 6.00 | OP ON 6.00Mhz | | | | |
| Udrift Went | H – I F/ON V Jency 6.00 litude 95.0 | OP ON €.00Mhz \$1V | | | | |
| Udrift Wext OFF Frequ Amp Offse | H – F/ON Z uency 6.00 litude 95.0 tt – 0.70 | OP ON 6.00Mhz € 58.1V € -0.67V | | | | |

Production Settings



| Lens 1 | 14.0 韋 | 14.0 V | | All on 🗹 |
|------------|----------|----------|-----------|----------|
| Lens 2 | 30.0 韋 | 30.0 V | | Lenses 🖂 |
| Lens 3 | 20.0 🖨 | 20.0 V | - | |
| Lens 4 | 60.0 韋 | 60.0 V | 1 | |
| Lens 5 | 70.0 韋 | 70.0 V | | |
| Lens 6 | 80.0 韋 | 80.0 V | | |
| Lens 7 | 17.0 | 17.0 V | | |
| Push L | 16.5 | 16.0 V | | 3 mA |
| Push H | 790.0 韋 | 790.0 V | | 3 mA |
| Pull L | 80.0 | 80.0 V | \square | 3 mA |
| Pull H | 680.0 🖨 | 680.0 V | | 3 mA |
| Grid | 2400.0 | 2282.0 V | | 1 µA |
| Cage | 5020.0 韋 | 4766 V | | 99 µA |
| lefl. Grid | 667.0 韋 | 634.0 V | \square | 75 µA |
| efl. Back | 900.0 韋 | 855.0 V | | 167 µA |
| MCP F | 5400 韋 | 5134 V | \square | 17 µA |
| MCP B | 2570 🖨 | 2465 V | | 224 µA |

TOF Voltages



| Acquisition | 1 | | | ACQ active |
|--|--|---|-------|------------|
| 🗃 🖬 层 | L. | | | |
| Single Spec | Time (ms) | 1000 | | |
| Extraction | n time (µs) | 5.0 | | 372.3 amu |
| max Flig | httime(µs) | 32.0 | + | 31.25 kHz |
| Data Save | Settings | | | |
| Spec 🖉 | \sim | Trace | | Raw |
| Time Durat | ion | | | 4 |
| 02:00:00 | Single File | Duration | ē | |
| 12 🗳 | Number | of Files To | Store | e) |
| C:\lonicon | \data | | | 6 |
| Add File | Count Ext Q for new f | ension file | | |
| <year>_<n Data_<hou< td=""><td>nonth>_<c r>_<minut< td=""><td>lay>\ te>_<secc< td=""><td>nd></td><td>62</td></secc<></td></minut<></c </td></hou<></n </year> | nonth>_ <c r>_<minut< td=""><td>lay>\ te>_<secc< td=""><td>nd></td><td>62</td></secc<></td></minut<></c | lay>\ te>_ <secc< td=""><td>nd></td><td>62</td></secc<> | nd> | 62 |
| 2022_11_16 | \Data_08_3 | 4_32_part | _XXX | |
| Mass Axis (| alibration | | | |
| | Cal | E | 30 |) sec 😫 |
| Mass | TimeBin | | | |
| 21.0220 | 16021 | 1 | ^ | a 15017.1 |
| 203.9400 | 161630 | 1 | | b -52827 |
| 220 9500 | 220322 | 龠 | 22 | |

Acquisition Settings







Pioneer Park



11-17-22 Parameters Globeville and Elyra-Swansea

| tanta and task and the second | | • 🕗 | |
|--------------------------------|--|---|--|
| Setting | Odor | Ø | |
| Primary Ion | H3O+ | ~ <i>9</i> | |
| Transmission | DC | | |
| | Man/Ctrl | Ctrl | |
| PC | 352.6 | 352.58 mbar | |
| p Drift | 2.30 | 2.29 mbar | |
| TofLens | | 4.79E-5 mbar | |
| TOF | | 7.06E-7 mbar | |
| E/N | | 120 Td | |
| Temps | 80.00 °C | 80.10 °C | |
| SrcValve | 50.0 🚍 | | |
| H2O | 6.0 | 6.00 sccm | |
| 02 | 0.0 | 0.00 sccm | |
| NO | 0.0 | 0.00 sccm | |
| lhc | 4 | 4.0 mA | |
| | On/Off | On | |
| FCinlet | 60.0 | 59.95 sccm | |
| U FU | °C D+ | C+ | |
| Us | 150 🚍 | 145.0 V | |
| Uso | 80 🚔 | 78.6 V | |
| Udrift | 525 | 526.1 V | |
| | | | |
| | | | |
| | + — | 0 × | |
| () Hex1 | + — | © × | |
| Hex1 | + —] []] /ON ☑ | OP ON | |
| Hex1 OFF Frequ | + — | OP OP ON 6.00Mhz | |
| Hex1 OFF, Frequ Ampli | 4 — /ON ✓ ency 6.00 tude 95.0 | OP ON 6.00Mhz \$8.4V | |
| Hex1 Frequ Offset | /ON / ency 6.00 tude 95.0 | OP ON €.00Mhz \$8.4V \$8.4V | |

Production Settings



| 3 🝺 🗖 | 18 | | | MC | P 2 |
|------------|--------|----|----------|-----------|----------|
| Lens 1 | 14.0 | • | 14.0 V | | All on 🗹 |
| Lens 2 | 30.0 | \$ | 30.0 V | | Lenses 🖂 |
| 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 | \square | 3 mA |
| Push H | 790.0 | \$ | 790.0 V | \square | 2 mA |
| Pull L | 80.0 | + | 80.0 V | \square | 3 mA |
| Pull H | 680.0 | ÷ | 680.0 V | \square | 3 mA |
| Grid | 2400.0 | + | 2283.0 V | \square | 1 μA |
| Cage | 5020.0 | \$ | 4766 V | \square | 99 µA |
| Refl. Grid | 667.0 | \$ | 634.0 V | \square | 75 µA |
| Refl. Back | 900.0 | + | 855.0 V | \square | 167 µA |
| MCP F | 5400 | \$ | 5134 V | \square | 17 µA |
| MCP B | 2570 | \$ | 2464 V | | 223 µA |

TOF Voltages



| Acquisition | | | | AC | Q active |
|--|--|---|-------|-------|----------|
| 🍯 🖬 层 | | | | | |
| Single Spec | Time (ms) | 1000 | ÷ | | |
| Extraction | n time (µs) | 5.0 | 4 | 3 | 72.1 amu |
| max Flig | httime(µs) | 32.0 | 4 | 3 | 1.25 kHz |
| Data Save S | Settings | | | | |
| Spec 🗹 | | Trace | | | Raw |
| Time Durat | ion | | | | \sim |
| 02:00:00 🖨 | Single File | e Duratio | n | | |
| 12 🖨 | Number | of Files To | o Sto | re | |
| C:\lonicon | data | | | | (|
| Add File | Count Ext Q for new 1 | ension file | | | |
| <year>_<n Data_<hou< td=""><td>nonth>_<c r>_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond</td><td></td><td></td></sec<></td></minut<></c </td></hou<></n </year> | nonth>_ <c r>_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond</td><td></td><td></td></sec<></td></minut<></c | lay>\ te>_ <sec< td=""><td>ond</td><td></td><td></td></sec<> | ond | | |
| 2022_11_17 | Data_09_2 | 7_35_par | t_XX | X | |
| Mass Axis C | alibration | | | | |
| | Cal | I | | 30 se | c 😫 |
| Mass | TimeBin | | | 1 | |
| 21.0220 | 16038 | Û | ^ | а | 15021.7 |
| 203,9400 | 161691 | Û | | b | -52831.5 |
| 20010 100 | | | | | |

Acquisition Settings



Field Zero Counts





Globeville



Elyra-Swansea



11-18-22 PTR Parameters Adams City



Production Settings



| 🚰 📴 🗠 | | | MCP | 1 and 1 |
|------------|--------|----------|-----|----------|
| Lens 1 | 14.0 | 14.0 V | | All on 🖂 |
| Lens 2 | 30.0 | 30.0 V | | Lenses 🖂 |
| 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 | 17.0 V | | |
| Push L | 16.5 韋 | 16.0 V | | 3 mA |
| Push H | 790.0 | 790.0 V | | 2 mA |
| Pull L | 80.0 | 80.0 V | | 3 mA |
| Pull H | 680.0 | 680.0 V | | 3 mA |
| Grid | 2400.0 | 2283.0 V | | 1 µA |
| Cage | 5020.0 | 4766 V | | 99 µA |
| Refl. Grid | 667.0 | 634.0 V | | 75 µA |
| Refl. Back | 900.0 | 855.0 V | | 167 µA |
| MCP F | 5400 😂 | 5134 V | | 17 µA |
| MCP B | 2570 | 2463 V | | 225 µA |

| 🛞 н | - | | × |
|-----------|--------|---|---------|
| r P B | ł | | |
| Hex1 | | | OP |
| OFF/ON | \sim | | ON |
| Frequency | 6.00 | 1 | 6.00Mhz |
| Amplitude | 95.0 | - | 58.4V |
| Offset - | 0.70 | - | -0.67V |
| < | | |) |

TOF Voltages



| Acquisition | | | | ACQ ac | tive |
|--|--|---|------------|-------------------|--------------|
| ii 🖓 🕞 🕞 | | | | | |
| Single Spec | Time (ms) | 1000 | * | | |
| Extraction | time (µs) | 5.0 | | 372.4 a | mu |
| max Flig | nttime(µs) | 32.0 | | 31.25 | kHz |
| Data Save S | Settings | | | | |
| Spec 🗹 | \leq | Trace | | Raw | |
| Time Durat | ion | | | ~ | |
| 02:00:00 \$ | Single File | e Duratio | n | | |
| 12 🤹 | Number | of Files To | o Stor | e | |
| C:\lonicon\ | data | | | 0 | |
| Add File | Count Ext Q for new 1 | ension file | | | |
| <year>_<n Data_<hou< td=""><td>nonth>_<c r>_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond></td><td>B23</td><td></td></sec<></td></minut<></c </td></hou<></n </year> | nonth>_ <c r>_<minut< td=""><td>lay>\ te>_<sec< td=""><td>ond></td><td>B23</td><td></td></sec<></td></minut<></c | lay>\ te>_ <sec< td=""><td>ond></td><td>B23</td><td></td></sec<> | ond> | B 23 | |
| 2022_11_17 | Data_09_2 | 7_35_par | t_XXX | | |
| Mass Axis C | alibration | | | | |
| BEE | K Cal |] | ∠ 3 | 0 sec | - |
| | | | | | |
| Mass | TimeBin | <u>19</u> | | | |
| Mass 21.0220 | TimeBin 16012 | đ | ^ | a 1501 | 15.6 |
| Mass 21.0220 203.9400 | TimeBin 16012 161607 | 1 | | a 1501 b -5282 | 15.6 28.8 |

Acquisition Settings



Field Zero





Adams City

Initial Calibration Screen Shots 11-13-22



BTEX 250, 50, 20 and 5 ppb Cal



100, 50 and 10 ppb Alkenes Cal





125, 25, 10 and 5 ppb H_2S Cals

| 550- | | | | | | | | | | | | | | |
|------------|----------------------------------|----------------------------|----------------------------|----------------------------|----------------------------|----------------------------|----------------------------|----------------------------|----------------------------|----------------------------|---|-------------------------------|--|----------------------------|
| 500- | en warder | Mar Shan | Here and | | | | | | | | | | | |
| 400- | exclaiming | u de la Mar | ACCER AND | | | | | | | | | | | |
| 350- | | | | | | | | | | | | | | |
| 300- | | | | | | | | | | | | | | |
| 250- | | | | . attituente a lui | and a second designed | and a state of a state of | | | | | | | | |
| 200- | | | | hadcaean | san san sa | a the second | 4 | | | | | | | |
| 150- | | | | | | | | | | | | | | |
| 100- | | | | | | | prosperso | Cranth (Paris and a | the second second | | | | | |
| 50- | | | | | | | Ţ | | | L | an the second | angunangu jaujan makananganga | and a state of a state | |
| 2:26:40.01 | PM 2:27:30.0 PM 22 11/13/2022 | 2:28:20.0 PM 11/13/2022 | 2:29:10.0 PM 11/13/2022 | 2:30:00.0 PM 11/13/2022 | 2:30:50.0 PM 11/13/2022 | 2:31:40.0 PM 11/13/2022 | 2:32:30.0 PM 11/13/2022 | 2:33:20.0 PM 11/13/2022 | 2:34:10.0 PM 11/13/2022 | 2:35:00.0 PM 11/13/2022 | 2:35:50.0 PM 11/13/2022 | 2:36:40.0 PM 11/13/2022 | 2:37:30.0 PM 11/13/2022 | 2:38:20.0 PM 11/13/2022 |

500, 250, 100 and 25 ppb Alkanes



APPENDIX E CALIBRATION GAS CERTIFICATION SHEETS



CERTIFICATE OF ANALYSIS

Grade of Product: CERTIFIED STANDARD-SPEC

Customer: M Part Number: Xi Cylinder Number: C Laboratory: 12 Analysis Date: D Lot Number: 12

MONTROSE AIR QUALITY SERVICES LLC X02NI99C15W0061 CC519990 124 - La Porte Mix - TX Dec 14, 2021 126-402278540-1 Expiration Date: Dec 14, 2024

Reference Number: Cylinder Volume: Cylinder Pressure: Valve Outlet: 126-402278540-1 144.3 CF 2015 PSIG 330

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 | Analytical Uncertainty | |
| HYDROGEN SULFIDE NITROGEN | 1.000 PPM Balance | 1.084 PPM | */-5% | |

Notes: MONTROSE AIR QUALITY SERVICES LLC PO3: PO018078



Signature on file Approved for Release

Page 1 of 1





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: Part | *CRYSTAL LAKE , IL* MONTROSE AIR QUALITY SERVICES X06NI99C15A00A3 | Reference Number: | 126-402159020-1 |
|------------------------------------|--|-------------------------------------|------------------|
| Number: Cylinder | CC344804 | Cylinder Volume: | 144.3 CF |
| Number: Laboratory: Analysis | 124 - La Porte Mix - TX Jul 30, 2021 | Cylinder Pressure: Valve Outlet: | 2015 PSIG 350 |
| Date: Lot Number: | 126-402159020-1 Expiration Date: Jul 30, 2024 | | |
| D d at an | monthing undfied by direct comparison to calibration standards trans | his to NICT which | and/or NICT |

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 | Analytical | |
| | | (Mole %) | 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



Approved for Release

Page 1 of 126-402159020-1





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

CERTIFICATE OF BATCH ANALYSIS

Grade of Product: ZERO

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

| | 1 84 92 883 8 | Here Rester | | |
|---------------------|---------------|---------------------|----------------------------|--|
| 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

Page 1 of 152-402047887-1





Airgan USA, LLC 6141 Easton Road Bidg 1 Plumsteadwille, PA 18949 Airgas.com

CERTIFICATE OF ANALYSIS Grade of Product: CERTIFIED STANDARD-SPEC

Customer: Part Number: Cylinder Number: Laboratory: Analysis Date: Lot Number: MONTROSE ENVIRONMENTAL GROUP X02AI99C15AH586 ALM060589 124 - Plumsteadville - PA Feb 19, 2020 160-401735121-1 Expiration Date: Feb 19, 2023

Reference Number: Cylinder Volume: Cylinder Pressure: Valve Outlet:

160-401735121-1 129.3 CF 2016 PSIG 590

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.

| | AN | VALYTICAL RESULTS | |
|-----------|-----------|----------------------|------------|
| Component | Req Conc | Actual Concentration | Analytical |
| BENZENE | 1.000 PPM | 1.055 PPM | +/- 5% |
| AIR | Balance | | |



Chil





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 Number: Cylinder CC164840. Number: Laboratory: 124 - La Porte Mix - TX Analysis Aug 09, 2021 Date: Lot Number: 126-402159021-1 Expiration Date: Aug 09, 2023

| Reference Number: | 126-402159021-1 |
|-------------------------------------|------------------|
| Cylinder Volume: | 144.3 CF |
| Cylinder Pressure: Valve Outlet: | 2015 PSIG 350 |

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 Analytic | | | | | |
| | | (Mole %) | 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 | Belance | | | | |

Notes:

MONTROSE AIR QUALITY SERVICES LLC PO#: PO-011307 NITROGEN BALANCE : 99.99939022%



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Page 1 of 126-402169021-1



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