Atmospheric benzene observations from an oil and gas field in the Denver Julesburg basin in July and August 2014

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Abstract

High time resolution measurements of volatile organic compounds (VOCs) were collected using a quadrupole PTR-MS at the Platteville Atmospheric Observatory (PAO) to investigate how oil and natural gas (O&NG) development impacts air quality within the Wattenburg Gas Field (WGF) in July and August 2014 as part of NASA’s DISCOVER-AQ Colorado field campaign. The PTR-MS measurements at PAO were supported by pressurized whole air canisters collected at the surface, vertical VOC measurements made using an airborne PTR-MS Time Of Flight (TOF) instrument installed on the NASA P-3B aircraft, and VOC measurements collected during the Front Range Air Pollution and Photochemistry Experiment (FRAPPÉ) on the NSF/NCAR C-130. Unexpectedly high benzene mixing ratios were observed at PAO (mean benzene = 0.53 ppbv, maximum benzene = 29.3 ppbv), primarily at night, and this high benzene is associated with SW flow over the observation site. Vertical diurnal profiles indicate that the benzene is originating from the nearby environment, and source investigations conducted with tracers measured in the pressurized canister samples implicate O&NG rather than urban benzene sources. A regional toluene/benzene ratio analysis associated southerly flow over PAO with the Denver urban plume, and the benzene/CO correlations are weak over all wind quadrants, indicating the urban emissions of benzene are not responsible for the high observations at PAO. These results allow us to put a spatial constraint on the source of the WGF benzene that has been measured in prior work at the Boulder Atmospheric Observatory (BAO).

Introduction

Oil and natural gas (O&NG) extraction has been rapidly expanding in the Denver-Julesburg Basin (DJB) in NE Colorado. In 2014 Colorado was ranked as the 5th highest state in number of gas wells, with 7.4% of the nation’s wells located in Colorado. Colorado produced 5.2% of the total US natural gas production in 2014. There are more than 25,000 active gas wells currently operating in the DJB, with the greatest density operating in the Wattenburg Gas Field (WGF) in Weld County (COGCC, 2015). The area has been drilled since the 1970’s, but recent high natural gas prices have created a larger market and incentivized natural gas development in the region. Increased unconventional extraction of oil and gas from shale deposits in particular is a concern in the public health community (Colborn et al., 2011) and has been shown to have an adverse effect on air quality in areas with heavy O&NG development (McKenzie et al., 2012).

Ozone is a secondary pollutant that is produced from reactions between reactive nitrogen (NO + NO₂ = NO₃) and VOCs under sunlight. The air quality impact of hydraulic fracturing has received
considerable attention from researchers and air quality managers because the Denver metro area and Northern Front Range were categorized as non-compliant with National Ambient Air Quality Standards (NAAQS) for ozone pollution in 2007, and the region has remained in marginal nonattainment through 2015 (Allison, 2015). Increased mixing ratios of volatile organic compounds (VOCs) have been attributed to oil and gas activity in natural gas extraction basins, and these increased VOCs have been connected to increased ozone (Katzenstein et al., 2003; Kemball-Cook et al., 2010; Petron et al., 2012; Gilman et al., 2013; Moore et al., 2014; Warneke et al., 2014). The O&NG contribution to VOCs in the front range is significant; the 2011 Environmental Protection Agency (EPA) National Emissions Inventory (http://www.epa.gov/air-emissions-inventories/national-emissions-inventory) facility emissions reports that 64% of the total point source VOC emissions in Colorado were attributable to O&NG extraction activity, while in Weld County the O&NG contributions account for 90% of the total point source VOC emissions.

Studies conducted within the DJB since 2011 have found a clear oil and gas signature in the VOCs measured in the region, with enhancement in the alkanes and aromatics without corresponding increases in combustion tracers (Petron et al., 2012; Gilman et al., 2013; LaFranchi et al., 2013; Swarthout et al., 2013; Thompson et al., 2014). In addition to the effect of degrading air quality, high VOC concentrations are a concern from a public health perspective. VOCs can be toxic to human health in their own right, especially as emissions increase from human activities. Increasing mixing ratios of VOCs in heavy O&NG development regions has been directly attributable to O&NG extraction, with sources coming from infrastructure and transport, fugitive emissions, and deliberate venting and flaring (Adgate, 2014). Increasing concentrations of methane, light alkanes, and aromatics in the DJB have been connected to O&NG development (Petron et al., 2012; Gilman et al., 2013; La Franchi et al., 2013; Thompson et al., 2014). Aromatics in particular are a public health concern, and the mix of benzene, toluene, ethylbenzene and xylenes (BTEX) has been well characterized. BTEX has a wide variety of industrial uses as solvents and feedstocks, and is released into the environment primarily through O&NG production and gasoline use and storage (Bolden et al, 2015).

Exposure to aromatic VOCs has been linked to cancer, respiratory effects, and endocrine disruption (McKenzie et al., 2012; Bolden et al. 2015). Benzene, the primary aromatic VOC, has a resonance stabilized ring structure and an atmospheric lifetime of approximately 10 days, with the primary atmospheric sink owing to oxidation by OH radicals. Benzene is classified as an air toxic under the USA Clean Air Act in section 112, and is a known human carcinogen (IARC Group 1). Health effects associated with benzene exposure include leukemia, anemia and other blood disorders and cancers, immune system impairment, decreased respiratory function and neural tube defects in newborn babies (Bolden et al. 2015; McKenzie et al. 2012). The effects on human health have been well documented in the literature, and benzene is one of the better characterized air toxics. Although benzene emissions are not controlled through national standards, exposure standards and guidelines have been set. However, exposure limits have generally been defined for industry and acute exposure, which leads to high limits. The Occupational Safety and Health Administration (OSHA) sets a general industrial permissible exposure limit of 1 ppm 8 hour time weighted average, with a 5 ppm short term exposure limit. The National Institute for Occupational Safety and Health (NIOSH) sets a lower short term 8 hour time weighted average of 0.1 ppm, with a short term exposure limit of 5 ppm. The California Air Resources Board (CARB) literature sets the acceptable exposure limits much lower, with an 8 ppb 6 hour exposure limit, and a 2 ppb 8 hour and chronic exposure limit. However, there has been no safe threshold value for benzene exposure, and the EPA Integrated Risk Information System (IRIS) assessment of benzene inhalation puts the excess carcinogenic risk from chronic exposure to benzene
at 1 in 100,000 for a chronic exposure of 0.41 to 1.41 ppbv benzene (1.3 to 4.5 µg/m³ benzene). Since there are uncertainties in the carcinogenic risk from benzene, the mean value of 0.877 ppb benzene for this excess risk factor is used (2.8 µg/m³ benzene).

Benzene is used as an anthropogenic tracer in VOC studies because it has a small number of well characterized sources, with 78% of the benzene emissions in the EPA’s 1996 National Toxic’s Inventory estimates attributed to mobile sources, mostly on-road. The remaining national benzene emissions are attributed to the evaporation of fuels at service stations, solvent use, and oil and gas operations and processing (Wallace, 1989; Bolden et al., 2015). Densely populated urban areas are the primary regional sources of benzene due to traffic and industry emissions. Urban outflow studies have used benzene as a tracer for urban influence in the rural environment (Roberts et al., 1984; Bravo et al., 2002; Seigneur et al., 2003; de Gouw et al., 2005; Warneke et al., 2007; Apel et al., 2010). However, increasing oil and gas production of aromatics means that while benzene mixing ratios have been dropping nationally (Fortin et al., 2005), there is some evidence that benzene concentrations have been increasing outside of the urban centers (Thompson et al., 2014). These gas field aromatic emissions have not been well characterized or unambiguously identified. In Colorado a top down estimate of benzene over the WGF found that the official inventories were too low by a factor of seven, and that the oil and gas emissions of benzene are on the same order as the vehicle emissions for the region (Petron et al., 2014). A specific source for the natural gas benzene was not definitively identified.

The present investigation, part of the 2014 NASA Deriving Information on Surface conditions from Column and Vertically Resolved Observations Relevant to Air Quality (DISCOVER-AQ) field campaign in July and August 2014, presents the first detailed analysis of benzene measurements made continuously at a sampling site in the midst of O&NG activity in the WGF. The benzene data were collected along with other VOCs by a PTR-MS at the Platteville Atmospheric Observatory, CO, one of six instrumented ground stations in DISCOVER-AQ. Vertical benzene measurements were made using an airborne PTR-MS Time Of Flight (TOF) instrument installed on the NASA P-3B aircraft which conducted regular spirals over Platteville. Additional VOC measurements were collected concurrently during the Front Range Air Pollution and Photochemistry Éxperiment (FRAPPÉ) on the NSF/NCAR C-130. From both surface and aircraft we found unexpectedly high, possibly toxic, benzene levels under nighttime conditions. In the following sections, we present the Platteville and aircraft benzene observations in detail to more completely understand emissions from O&NG activities in the WGF. We address two main points in this study. First, we examine benzene variability, which is characterized by comparison of aircraft data across the FRAPPÉ region, over DISCOVER-AQ sampling sites, and at the Platteville surface site on a diurnal basis and over three weeks of sampling. Second, we investigate potential benzene sources at the Platteville ground station, using benzene data measured in canister samples and by the PTR-MS. Tracer correlations with benzene and wind direction-benzene relationships measured at the site are used to identify general source locations and likely O&NG activities contributing to the elevated benzene.

Methods

Methods: Site Description

NASA’s DISCOVER-AQ field campaign was conducted from mid-July to mid-August of 2014 in the Colorado Front Range region. The measurements encompassed the Denver urban region and the rural gas field environment to the north of the Denver urban region bounded by Fort Collins, CO and the Platteville Atmospheric Observatory (PAO). A map of the study area is shown in Figure 1. The campaign measurements included six instrumented ground sites and two instrumented NASA aircraft (the P-3B and
the B200). This study primarily analyzes a subset of these measurements collected at the Platteville Atmospheric Observatory (PAO) (40.1815 °N, 104.7268 °W). PAO was selected due to its central location in the WGF, allowing the characterization of the gas field emissions. The site is located 9 km SE of the City of Platteville (Lat: 40.214 °N, Long: 104.823 °W) and 25 km S of Greeley CO (40.422 °N, 104.709 °W). The site elevation is 1520 m above mean sea level (ASL).

Air quality relevant trace gas sampling was made with a common inlet 4 meters above ground level (AGL) within the Nittany Atmospheric Trailer and Integrated Validation Experiment (NATIVE) mobile air quality research station (see Martins et al. (2012) for a full description). The NATIVE instrument package provides one minute averaged in situ measurements of ozone, NO/NO₂/NOₓ, total reactive nitrogen (NOₓ), sulfur dioxide (SO₂), carbon monoxide (CO), carbon dioxide (CO₂), and methane (CH₄). Additionally, one-minute averaged meteorological data was also collected, including wind speed and direction, temperature, relative humidity, pressure, and NO₂ photolysis rates.

Methods: PTR-MS VOC Collection

A conventional proton-transfer-reaction quadrupole mass spectrometer (PTR-QMS; Ionicon Analytik, Innsbruck, Austria) was used for VOC measurements at the PAO site. The measurement principle and technical details of this instrument have been reviewed in great detail elsewhere (DeGouw and Warneke, 2007). Routine operating parameters and procedures were applied in this study. The drift tube was operated at a pressure of 2.25 mbar, an electric field strength of 57 V/cm and a temperature of 50 °C. The QMS was run in the multiple ion detection (MID) mode. It sequentially measured 51 m/z-signals over the course of one minute. Benzene was detected as protonated benzene (m/z 79) with a 1-second signal integration time during the 1-minute measurement cycle. The ethylbenzene interference with benzene measurements was minimized (~25 % fragmentation on m/z 79). The interference from acetic acid (m/z 79 from hydrated protonated acetic acid) was found to be negligible.

A dynamically diluted certified multi-component gas mixture (Apel Riemer Environmental Inc., Broomfield, CO, USA) containing 1 ppmV of benzene was used for calibration. The instrumental background was determined by supplying catalytically (Pt/Pd at 325 °C) cleaned air to the instrument. The accuracy of the reported benzene volume mixing ratios is ±5.2%.

The PTR-QMS was housed at the PAO site in a separate trailer located 5 m from the NATIVE trailer. Ambient air was sampled at a height of 5.6 m above ground (2.4 m above trailer roof) from a mast attached to the trailer. The inlet line was a 6.25 mm outer diameter, 4.3 m long Teflon PFA tube that was capped with a 5-6 μm pore size Teflon PTFE filter. The tube was heated to 40 °C inside the air-conditioned trailer to prevent condensation. The flow rate through this main inlet line was 7 standard liters per minute (slpm). The PTR-QMS subsampled from this flow.

A proton-transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS) was used for airborne benzene measurements. This instrument and its operating procedures have been described in great detail by Müller et al. (2014).

Methods: FRAPPÉ TOGA VOC Collection

The NCAR Trace Organic Gas Analyzer (TOGA), a fast on-line gas chromatograph/mass spectrometer (GC/MS), was installed on the NSF/NCAR C-130 for the FRAPPÉ field study and made continuous measurements of over 70 VOCs including benzene and toluene during the study. Integrated air samples 35-s in duration were taken every two minutes and analyzed by TOGA in flight. For a detailed description of TOGA, see Apel et al. (2015). The reported TOGA benzene and toluene data have an overall uncertainty of ±15%, with detection limits of 0.5 pptV and 1 pptV for benzene and toluene, respectively.
Methods: Canister Sample Whole Air VOC Collection

The whole air canister samples were collected and analyzed by the University of California at Irvine (UCI) VOC canister collection program. Modules of 12 stainless steel 0.9 liter canisters connected in series with remotely operated valves. The fill time was ~30 seconds and the evacuated canisters reached final pressures of ~10 psig. After the canisters were filled the modules were shipped to the UCI laboratory where they underwent GC analysis for ~30 halogenated hydrocarbons and alkyl nitrates and ~50 C2-C10 nonmethane hydrocarbons (NMHCs). The limit of detection (LOD) was 3 pptV for all NMHCs while the halogenated gases/alkyl nitrate LODs ranged from 0.05 pptV to 10 pptV. For more details see Colman et al., 2001.

Methods: Ozonesonde Boundary Height Calculations

Planetary boundary layer height (PBLH) calculations were made from ozonesonde profiles that were collected at the PAO site. One to three ozonesonde-radiosonde profile pairs were launched each day for a total of 42 total profiles. Boundary layer height was calculated from these profiles using the method outlined in Martins et al. (2012). The profiles were smoothed by averaging the vertical data into 50 m bins, and the lapse rate for defining the inversion was set at a minimum of 5 K/km. In addition we added the criteria that the relative humidity did not change with altitude for these samples. If multiple possible heights were selected with these metrics we used the lowest realistic value by inspection.

Results: Spatial Distribution of Benzene

Six sites collected ground measurements during the campaign. Vertical measurements were made by the P-3B flying spirals over the sites; each measurement location had two to three spirals per flight day. VOC measurements were made with a PTR-MS Time of Flight (TOF) instrument, including benzene. To determine the spatial differences of benzene measured over the sites we used the bottom one km AGL over each site to calculate benzene statistics for the campaign. Figure 1 shows the mean benzene calculated for each site are shown in context of the study area including the flight track taken by the aircraft and the boundaries of the WGF. The anthropogenic sources of benzene lead to high mixing ratios at low altitudes, so confining the statistical calculations to a single vertical km for each site allows us to assume that we are comparing measurements that were primarily collected within the planetary boundary layer.

PAO shows the highest mean benzene over all spirals (0.140 ppbv), followed by Denver (0.119 ppbv) and the Boulder Atmospheric Observatory (BAO) (0.102 ppbv). PAO and BAO are both gasfield sites, while Denver was the urban core. In comparison, the three sites closest to the mountains (Fort Collins, Golden, and Chatfield) have low values relative to the gas field and urban sites. Fort Collins has the lowest mean value during the campaign (0.068 ppbv) while the two sites expected to be most influenced by the Denver Plume (Chatfield (0.080 ppbv) and Golden (0.087 ppbv) are slightly elevated relative to Fort Collins but still low compared to Denver.

Table 1 shows the aggregate statistics for the six measurement sites. Putting the mean values into context with the quartile values for each site shows that the means are relatively uniform, with Denver and PAO in particular having close values despite the difference in environment. The two locations with the highest observed values, BAO and PAO, are both gas field sites and are within the boundary of the WGF. Benzene is expected to be high in the city center where vehicle emissions are the driver of anthropogenic VOCs. Thus the high values in measured at the rural gas field sites are unexpected and indicate different source influences on the observed benzene.
Results: PAO Benzene Observations

The time series of ground site benzene measured by PTR-MS during the campaign is shown in Figure 2A. Figure 2B shows the daily benzene statistics, calculated from midnight to midnight local time (MDT). The campaign diurnal cycle is shown in Figure 2C, and shows the extreme difference between the daytime and nighttime data. The high nighttime benzene drives the high statistics over all measured days in the campaign, Table 2. The diurnal split for the statistics is based on the sun's position relative to the horizon; the sun above the horizon was defined as daytime, and below the horizon was defined as nighttime. During the daytime benzene mixing ratios are reduced and become less variable, while the nighttime benzene shows much greater variability and the largest observed mixing ratios of the campaign. Beyond the day/night diurnal split, Figure 2B also shows that there is some temporal variability in the observed benzene, with the later third of the campaign from 31 July to the end of the observations on 08 August having higher nighttime benzene.

The benzene measured at PAO was higher than expected for the region. Canister samples collected in the City of Platteville in 2013 saw peak benzene of 0.96 ppbv (Thompson et al. 2014), well below the 2014 PAO maximum of 29 ppbv. While some of this disparity is explained by the differences in sample techniques and the fact that the samples in Thompson et al. (2014) were collected between 0600 and 0900 MST, the difference in the benzene range was unexpected. The benzene levels measured at PAO during the 2014 campaign are high enough to be problematic from a health and welfare standpoint. The fact that these high benzene mixing ratios mostly occurred at night, when canister samples are not routinely collected in this region, means that the canister studies previously conducted in the WGF did not capture the full range of the benzene exposure.

Figure 3 shows the benzene relationship to PBLH, calculated from both the ozonesonde-radiosonde profile pairs and MPL aerosol data. The surface benzene concentrations are sensitive to the planetary boundary layer height (PBLH), especially in the early morning and late evening when the PBLH is low but the sun is still above the horizon. Under these conditions the boundary layer has a stable inversion structure, which prevents vertical mixing and keeps any ground level emissions of VOCs close to the earth’s surface, driving up mixing ratios at the surface. Benzene mixing ratios are enhanced when the PBLH is shallow, but as convective mixing begins and the PBLH begins to grow, the benzene mixing ratios drop off. Once the PBLH is greater than 0.5 km above ground level (AGL) the benzene mixing ratios level off and approach the daytime median benzene value.

The MPL PBLH measurements generally agreed with the ozonesonde PBLH measurements collected at the site. The mean and median MPL PBLH begin growing at 0900 MDT and level off between 1700 and 1900 MDT (Supplementary Figure 1). At 1000 MDT both the mean and the median of the MPL PBLH are \( \geq 0.5 \) km AGL, which is the PBLH height where benzene stops being sensitive to PBLH. After 1900 MDT the MPL PBLH can continue to increase, but since this method uses aerosols to define the boundary layer, these measurements can be unreliable. Aerosols lofted into the troposphere as the PBLH grows will remain in the troposphere even when the PBLH collapses, which makes determining the true nighttime PBLH difficult or impossible using the MPL.

Results: Aircraft Comparisons

The ground observations at PAO were supported by vertical profiles sampled by PTR-MS of the P3B aircraft, collected by spirals conducted in a top-down approach over the site, with the aircraft starting high and ending at low altitude. An overview comparison between the benzene measured on the P3-B and the ground instrument is shown in Supplementary Figure 2. Flights occurred over the PAO between the
hours of 0800 and 1600 MDT throughout the campaign with two to three full profiles scheduled on each flight day. A total of 32 full vertical profiles were completed over PAO during the campaign.

Figure 4 shows the diurnal pattern of benzene measured in the vertical columns and on the ground, faceted by hour of day binned vertically by 50 m sections. The overall mean vertical profile is shown in Supplementary Figure 3. In the early morning hours the benzene measured on the ground is high, with low benzene in the aircraft measured column. As the day progresses and mixing in the boundary layer begins two trends begin: first, the ground measured benzene decreases, second, the benzene in the column increases. A clear benzene boundary layer can be observed in the vertical benzene measurements growing throughout the morning and midday. By midafternoon the troposphere is well mixed with respect to benzene.

The high ground benzene at PAO coupled with the low benzene mixing ratios in the early morning spirals indicates that a pool of benzene builds overnight at the surface. The benzene emitted into the low boundary layer overnight is mixed up into the troposphere as the boundary layer grows, causing the mean benzene to fall with height. Similar decreases of benzene concentration with height were observed at BAO in 2011 (Swarthout et al. 2013), and attributed to local emissions of gas field benzene. The diurnal profiles indicate that the high benzene comes from the surface, and this indicates advection aloft is not a significant contributor to the benzene measured in the columns.

**Results: Benzene Polar Frequency Analysis**

To get more information on potential sources, we used polar frequency analysis of the surface measured benzene at PAO. These plots use wind speed and direction information to determine how wind regimes are associated with the high benzene observations. The wind speed and direction data was binned by 10 degrees wind direction and 0.5 m/s wind speed, and statistics on the associated benzene observations are calculated for each bin independently. The full campaign polar frequency plots are presented in Figure 5. Benzene is associated with W and SW wind flow, with the highest mixing ratios originating from the southwest in a confined wind regime.

Separating the polar frequency plots into median (Fig. 5a), mean (Fig. 5b) and maximum (Fig. 5c) allows us to investigate the directional dependence of the enhanced benzene at different mixing ratio levels – low and background through the median, mid-level through the mean, and the wind regimes associated with the highest benzene through the maximum. In all three polar frequency plots the benzene is associated with SW flow. A westerly enhancement in the median plot is most likely due to mountain-valley circulation over the site, but the persistent SW enhancement of benzene to the SW in all three polar frequency plots indicates a primary benzene source SW of PAO. Although there is not a distinct point source, the high benzene observed at PAO has a directional component; it is not simply a general elevation of mixing levels throughout the region.

**Results: Canister Correlations and VOC Influences at PAO**

To assess the general influence of VOC sources at PAO we use four O&NG VOCs that correlate with the O&NG and urban tracers, propane and acetylene: i-pentane, n-pentane, n-butane, and ethane. All four of the tracer species are primarily emitted from natural gas, but can also be sourced from traffic combustion in small amounts (Gentner et al. 2013; Thompson et al. 2014). The results are shown in Figure 6a and 6b. The canister sample correlations of the four tracer species with propane are strong, with i-pentane (R² = 0.985), n-pentane (R² = 0.971), n-butane (R² = 0.992) and ethane (R² = 0.936), showing excellent correlation over all 35 samples. In contrast, the correlations with acetylene are much weaker; i-pentane (R² = 0.492), n-pentane (R² = 0.434), n-butane (R² = 0.504) and ethane (R² = 0.424). The tracer correlations
presented in our study agree very well with the correlations of benzene and propane that are reported in Thompson et al (2014), a study conducted with canister samples collected by the CDPHE in the city of Platteville in 2013. Strong correlations with acetylene would indicate that the Denver plume is the most important VOC source due to the heavy traffic influence in the urban center, but the propane correlations are exceptional and indicate that PAO was heavily influenced by O&NG activity in 2014 during DISCOVER-AQ.

The comparison of Thompson et al (2014) to the PAO canister samples is presented in Table 4; this comparison study is currently the best published comparison of VOCs and correlations for non-BAO measurements in the WGF. Our reported correlations with acetylene agree in general with Thompson et al (2014), although our $R^2$ values are much higher. There is far less acetylene measured in the canister samples at the PAO, which is isolated from major traffic, compared to the City of Platteville, which straddles US 85, a N-S highway with heavy truck traffic. Conversely, propane is expected to have higher mixing ratios at PAO due to its location centered in the gas field; this is verified in the observations.

While the VOC/tracer correlations in Figure 6a and 6b indicate that an O&NG source important for benzene at PAO, we wanted to quantify this relationship for the benzene samples. The high benzene measured at PAO coupled with the vertical diurnal profile, Figure 4, and the directional dependence, Figure 5, indicates that benzene originates from some kind of anthropogenic activity from the SW of PAO. To better identity the primary benzene source, tracer species well characterized for atmospheric sources are analyzed. Commonly used compounds in O&NG studies include the VOCs propane, an O&NG tracer, and acetylene, a traffic and urban tracer. These tracer compounds are not measured by PTR-MS, but pressurized canister samples collected at PAO during the P3-B spirals were analyzed for these tracers. Note, however, that the canister samples are confined to daylight observations.

Benzene was collected in both the canister sample and by the PTR-MS, and these two techniques had excellent agreement over the measurement period (Supplementary Figure 4). Using a two minute average centered at the time of canister opening to accounting for the slight separation of the instruments returned a correlation slope of slope of 1.007 ppbv PTR-MS benzene/ppbv canister benzene and an $R^2$ value is 0.921. Figure 6c and 6d show the results of correlating canister and PTR-MS benzene to propane and acetylene measured in the canisters. The correlations with propane (Figure 6c) are moderately correlated (canister benzene to canister propane, $R^2 = 0.782$; PTR-MS benzene to canister propane, $R^2 = 0.700$), and indicate that there may be some common source. In contrast the correlations with acetylene (Figure 6d) are very weak (canister benzene to canister acetylene, $R^2 = 0.375$; PTR-MS benzene to canister acetylene, $R^2 = 0.272$). The benzene correlations with acetylene and propane are comparable to literature values collected in the City of Platteville reported in Thompson et al. (2014), which found that benzene correlated with propane ($R^2 = 0.73$) much more strongly than to acetylene ($R^2 = 0.29$).

In emissions budgets benzene is primarily attributed to combustion emissions. The CDPHE keeps a bottom-up emission inventory based on permit data and empirical calculations, and the 2011 inventory assessments assigned nearly 90% of the total front range benzene emissions to vehicles, both highway and nonroad (Petron et al. 2014). Benzene has been quantified in the WGF by three papers: two studies at BAO (Petron et al. 2012; Gilman et al. 2013), and the Thompson et al. (2014) study in the City of Platteville. These studies indicate that benzene Platteville has contributions from both traffic and non-combustion evaporative sources. While these studies did not quantify the source contributions on the ground, a top down estimate of benzene over Weld County was seven times higher than the CDPHE inventory estimates of benzene emissions for the same region (Petron et al. 2014). They conclude that the benzene emissions from evaporative sources over the WGF are on the same order of magnitude as the vehicle emissions, and
that mismatch between the top down measurements and the state inventory assessment is consistent with missing benzene emissions in the Colorado Front Range Region. The assertion that the benzene evaporative emissions could be equal or greater than the vehicle emissions is substantiated by the correlation results that we see in our work, with the propane tracer having a stronger correlation to benzene than acetylene.

The isomers of pentane, i-pentane and n-pentane can be used as additional tracers for distinguishing traffic from natural gas sources. Tailpipe emissions enhance i-pentane, so the i-pentane/n-pentane is used as a fingerprint to discriminate urban vs natural gas VOC sources. High ratios of i-pentane/n-pentane indicate urban and traffic sources, whereas lower ratios are a fingerprint of O&NG sources. Furthermore, these are affected equally by atmospheric mixing and dilution and by photochemical processing, so the source ratios remain constant (Gilman et al. 2013). A comparison of the PAO observations to published values is shown in Figure 7. The canisters collected at PAO in July and August 2014 have a i-pentane/n-pentane ratio of 0.888 ($R^2 = 0.9885$), a value close to the ratio of 0.86 reported for raw natural gas in the WGF (CDPHE, 2007). Other studies in NE Colorado have found similarly low ratios: 0.885 at the BAO in 2011 (Gilman et al. 2013), 0.965 in Longmont CO (Thompson et al. 2014), and 1.00 at BAO (Swarthout et al. 2013). In contrast, fresh vehicle emissions have a ratio of 2.96 (Broderick and Marnane, 2002). Measurements taken in Denver, CO, and Boulder, CO, show a greater influence from the fresh vehicle emissions, with Denver having a reported ratio of 1.87 (Baker 2008), and Boulder recording 1.10 (Gilman 2013). PAO is dominated by O&NG influence during the day when both urban/vehicle and O&NG emissions can influence the site.

A final correlation between i-butane and n-butane was calculated, which returned a campaign ratio of 0.405 ($R^2 = 0.993$). This ratio is in general agreement with the i-butane to n-butane ratios measured by Swarthout et al. (2013) and Thompson et al. (2014), respectively 0.41 and 0.43, and to measurements collected by the CDPHE (2012), $4.2 \pm 0.03$. These regional ratios are comparable to measurements in well mixed urban and rural regions (Parrish et al., 1998; Watson et al., 2001), but lower than the ratio expected for vehicle emissions or gasoline (Conner et al., 1995; Jobson et al., 2004). Emissions from commercial natural gas have measured i-butane/n-butane at 0.60 (Fujita, 2001), but butane ratios this high have not been recorded in the WGF, indicating that the natural butane ratio for this gas field are naturally around 0.4.

**Results: Toluene/Benzene Ratios**

Toluene is co-emitted with benzene with source dependent ratios. Toluene photolyses more rapidly than benzene under ambient conditions and the ratio of these two compounds can be used to assess the photochemical processing of an airmass; a lower ratio of toluene to benzene indicates an aged airmass (Roberts et al. 1984). In idealized urban outflow scenarios a photochemical clock calculation performed with benzene and toluene can be used to calculate an effective airmass age. However, these calculations don’t work reliably in areas where there are additional non-vehicle emissions of the aromatic compounds (Gelencser et al. 1997), such as in the WGF. However, these calculations can still be useful for identifying and separating observations at PAO. Fresh emissions from gasoline evaporation and vehicles without catalysts have a toluene/benzene ratio $\geq 2$ (Barletta et al. 2002; Heeb et al. 2000; Bravo et al. 2002), and the mean ratio for the Denver urban region was measured as 2.1 (Baker, et al 2009). A spatial map of the toluene/benzene ratio calculated from the C-130 TOGA instrument, Figure 8, shows that using a toluene/benzene ratio cutoff of 2 or greater is appropriate for differentiating the urban environment from the surrounding rural environment.
The spatial toluene/benzene analysis presented in Figure 8 offers context on both the ratio analysis as well as the spatial distribution of benzene in the study area. This plot encompasses all data collected during the Frappé field campaign (27 July 2014 to 18 August 2014) to offer a broad context for the observations. The points are colored by the toluene/benzene ratio cut off value of 2, but the points are sized by the magnitude of the benzene observation for each point. The highest benzene of the campaign is measured within the boundary of the WGF, but these high observations occur on the W side of the Greeley urban area as well as in the SW corner of the WGF just to the north of the Denver urban boundary. While PAO has high benzene measurements through the P-3B observations and ground observations, the TOGA data shows that PAO is associated with comparatively low values. This is both evidence of the value of ground measurements, especially at night, as well as some additional evidence for a benzene source located to the SW of PAO. Interestingly, the highest benzene observations are not associated with NE flow over the site despite the high benzene values over Greeley, CO. This is explained by the flow characteristics over PAO, where SW flow is more likely (Figure 4D), and the Greeley, CO, emissions are not advected strongly to the SW to affect PAO.

Figure 9A shows the correlation of benzene to toluene for all PAO PTR-MS observations taken during the campaign; the data with a toluene/benene ratio ≥ 2 has been colored. This high ratio of toluene/benzene only accounts for 5% of the total benzene observations; furthermore, these observations are a distinct lobe of the correlate plot. When the maximum values viewed spatially in a polar frequency plot (Figure 9B) it is clear that these high values are originating from due south of PAO, an area that is not associated with the elevated benzene.

A previous study done at BAO used the toluene/benzene ratio separated by wind sector to attribute emissions types in each sector (Swarthout, 2013). This study also found the highest mean toluene/benzene ratios to the south (1.09 ± 0.40), and attributed the source to regular impact from the Denver plume. Lower ratios of toluene/benzene in the NE sector (0.76 ± 0.25) are attributed to O&NG emissions. When the mean toluene/benzene ratio is calculated by wind sector for PAO (Table 3), we also see the highest ratios associated with southerly winds (1.41 ± 1.86). This result, coupled with the polar frequency analysis of the max toluene/benzene ratio, Figure 9B, indicates that a recently emitted urban signal is sourced from south of PAO and the high benzene signal from the SW is not associated with the Denver plume. This is an important result. The high benzene measured in the WGF at PAO clearly originates from local gas field sources, and is not due to urban outflow.

The full list of the toluene/benzene ratios are in Table 3. The ratios show a distinct diurnal pattern from all wind directions, with higher ratios observed at night in all sectors. The lowest mean ratios are observed in the E and SE wind sectors but these two sectors are also associated with the lowest mean values of both benzene (E = 0.28 ± 0.36 ppbv, SE = 0.20±0.25 ppbv) and toluene (E = 0.35 ± 0.55 ppbv, SE = 0.25 ± 0.37 ppbv). This is in contrast to Swarthout (2013), who observed their depressed toluene/benzene ratios occurring in the NE in association with increased aromatics. The PAO observations show approximately equal value toluene/benzene ratios from all sectors, including during calm winds (wind speeds < 1 m/s). These high ratio values compared to Swarthout et al. (2013) and the uniformity with respect to wind direction indicates that there is considerable directional impact from the Denver urban plume at PAO. However, when it does occur it is associated with S rather than SW flow.

Results: Benzene and CO Relationship

The previous WGF benzene studies at BAO used correlations with CO to differentiate benzene sources (Petrone et al., 2012; Swarthout et al., 2013). Urban emissions of benzene are correlated with CO
due to their major common source in vehicle exhaust; at BAO benzene originating from the southern wind sector correlates well with CO ($R^2 > 0.8$ for S wind sector). Both of these studies also identified enhanced aromatics in the NE wind sector; the benzene/CO slopes were enhanced while the two species were still correlated ($R^2 = 0.58 – 0.86$).

When the benzene/CO correlations are calculated for the PAO data over all observations, the $R^2$ value of 0.099 suggests no correlation. Both benzene and CO have a strong directional component (Figures 4 and 10), and both species have their highest correlations from SW of the site. However, there is no directional dependence in the correlation of benzene and CO over the campaign; even in the SW wind sector, high benzene occurs during low CO measurements.

High benzene without a corresponding CO source is observed from all wind sectors, which can be explained by an O&NG benzene source. In contrast, elevated CO is associated exclusively with the SW wind regime (Figure 10). While benzene and CO do track occasionally, as expected for a vehicle or urban source, often these two species occur independently. The directionally dependent correlations of CO and benzene are shown in Figure 11. There are three major benzene/CO regimes observed during the campaign: benzene and CO tracking from a common traffic or urban source, benzene increases without associated CO increases, and CO increases without associated benzene.

The highest independent CO and benzene observations are observed from the same SW wind sector, but they are independent in time, indicating that there are two separate sources for these important molecules that are both located in the SW. The tightly constrained wind regimes associated with high CO and the lack of correlation with benzene indicates that it’s unlikely these high CO values originated from either the Denver urban area or from major freeways to the W and SW of PAO. Instead, it seems more likely that we observe a stationary CO source and a separate temporally variable benzene source, both located in the same general region to the SW of PAO.

In Colorado CO emissions have been attributed gasoline vehicles (66%) and non-road gasoline powered equipment (26%) (LaFranchi et al. 2013). The remaining emissions are attributed to a variety of processes, including industrial activity and natural gas processing. The extremely high observed mixing ratios of CO coupled with the lack of a correlation with benzene or other tracers indicates that a close range CO production is not associated with vehicle emissions.

**Discussion: Potential Sources of Benzene**

If the high benzene observed at PAO was primarily from the Denver metropolitan area we would expect to see it correlate strongly to the combustion tracers, acetylene and CO (Bourbon et al. 2013; Apel et al. 2010). However, we see weak and non-existent correlations over the full campaign for these species. In addition, the highest benzene occurs with low CO mixing ratios. High benzene that is not correlated with vehicle combustion tracers has been observed in areas with heavy industry, such as Houston TX (Gilman et al. 2009). O&NG emissions of benzene are expected to correlate better with other O&NG VOCs, such as propane and ethane. Studies in areas of heavy O&NG production have shown this behavior, especially for sites associated with natural gas production (Helmig et al. 2014; Petron et al. 2014, Warneke et al. 2014).

Observational work conducted at BAO has found anomalously large benzene mixing ratios from the NE, which they attributed to general O&NG production in Weld county (Petron, et al. 2012; LaFranchi et al. 2013, Swarthout et al. 2013). However, further specification of the exact source of the high benzene has not been made, except to assume that there is a mix of automotive and natural gas sources in the WGF. A separate source of urban benzene was measured from S and W of BAO. The measurements taking at PAO for our 2014 work are based much deeper in the WGF and are NE of the BAO tower. At PAO the
highest measurements of benzene come from the SW, indicating that the source of high benzene measured
from the NE in the BAO papers is located somewhere in between the BAO and PAO sites. The SW quadrant
is influenced by a mix of air sourced from vehicle exhaust as well as from this unidentified benzene source,
resulting in the highly uncorrelated benzene/CO behavior that we observe.

Our observations, along with the BAO observations (Petron, et al. 2012; LaFranchi et al. 2013,
Swarthout et al. 2013), confine the high benzene to a smaller geographic area within the WGF. In addition
to a narrowed source area we also know that the benzene mixing ratios are highly variable, especially at
night, indicating that a transient source related to O&NG production is operating near PAO. Benzene
emissions have been a concern in upstream gas production, and VOCs have been identified as emissions for
various parts of the natural gas extraction process. Specifically, these processes are 1) diesel vehicle
emissions; 2) gas flaring and deliberate venting; 3) drilling flowback of waste mud and water; 4) fugitive
emissions from infrastructure and maintenance; 5) condensate tanks (Adgate et al. 2014; Verma et al. 2000;
Buzcu-Guven et al. 2013).

The benzene observations do not correlate with formaldehyde measurements ($R^2 = 0.003$) at PAO;
these formaldehyde measurements were made with the PTR-MS on the surface at the observation site. A
correlation could have indicated a flaring source (Knighton et al. 2012), which has been identified as a
benzene driver in Texas (Buzcu-Guven et al. 2013). Our tracer correlations eliminated traffic and urban
sources as well. This leaves the most likely potential O&NG sources as one of four potential options:
drilling flowback, gas venting during production or pipeline maintenance, fugitive emissions from
infrastructure, or emissions from condensate tanks.

Drilling flowback and waste water is stored in temporary waste pits at the drilling sites (Colborn et
al. 2011), the locations of which are monitored and reported by the Colorado Oil and Gas Conservation
Commission (COGCC). Regulatory advances have phased out this type of storage in favor of closed
systems (CDPHE, 2012), but active locations are still recorded in the WGF by the COGCC. There is a
dense group of these pits located between PAO and BAO (Supplementary Figure 5). While many of these
are inactive locations, meaning decommissioned and buried, a large number of them have an unknown
status. Additionally, a number of the active pits are located west of PAO, which is associated with high
mean and median benzene mixing ratios and low mixing ratios of CO. However, produced water and
drilling waste has not been identified with high atmospheric benzene concentrations in previous studies
(Bloomdahl et al., 2014; Bunch et al., 2014), although published atmospheric measurements of these waste
pits are difficult to locate. Environmental impacts from these waste pits are more likely to be associated
with poor water quality (Eiceman et al., 1986; Gross et al., 2013)

While we’ve rejected flaring as a significant source of benzene based on the correlations between
benzene and formaldehyde, the proximity of O&NG facilities to air quality monitoring stations has been
shown to increase measured benzene levels; gas plants < 50 km from the measurement sites were shown to
have a statistical increase for benzene, although a mechanism was not hypothesized (Bursyn et al. 2007).
PAO was located 4.5 km to the NE of the Platte Valley Gas Plant, which is owned by DCP Midstream LP
and was operational through our study period. Fugitive VOC emissions, including benzene and toluene, are
expected in refining facilities, mostly from storage tanks and poor seals, but also from conversion and
treatment processes (Rao et al., 2005). In oil refineries the coker unit has specifically been implicated (the
process cracks long chain hydrocarbons into shorter molecules), along with the storage areas for the raw
feedstock and final product tanks (Chambers et al. 2008). Natural gas processing does not have a coking
process, but leaks from storage containers may be a significant source of VOCs from the processing plant
(Bar-Ilan et al., 2008).
Condensate tanks have been implicated in gas field VOC emissions in previous studies (Moore et al., 2014). In exposure studies of upstream natural gas, the gas dehydrator units are implicated as the largest single factor of exposure for workers (Capleton and Levy, 2005). Flashing emissions from these units have been estimated to produce about 65% of the total natural gas extraction VOCs in Colorado (Pertron et al. 2012). A study using a mobile platform in the Uintah Basin found a clear aromatic source from the condensate and water tanks on the gas well pads (Warneke et al. 2014).

Petron et al. (2012) implicated venting and condensate tanks in their assessment of benzene emissions from O&NG operations, but noted that the benzene to propane ratios modeled from condensate tank flashing and vented emissions was a factor of two lower than the benzene to propane ratios they measured at BAO. The measured ratio BAO (10.1 ppt/ppb benzene/propane) is similar to the ratio of benzene and propane measured in the canister samples from this work (0.011 ppb/ppb benzene/propane) (Figure 7), an indication that the modeled condensate benzene emissions are too low and are not capturing the full range of benzene being emitted during this process. This indicates that the O&NG processes responsible for benzene in the WGF are still not well understood.

Conclusions

A PTR-MS quadrupole instrument collected continuous VOC time series data at PAO in the WGF as part of the 2014 July-August DISCOVER-AQ field campaign. The interpretation of VOC measurements is supported by meteorology and general AQ monitoring collected at minute resolution, as well as compressed air canister samples collected during flight days. In addition to the ground based measurements 32 vertical profiles of VOCs measured by PTR-MS were collected over the site.

Unexpectedly high mixing ratios of benzene were observed at PAO during this study. The lowest mixing ratios occurring the daytime, while the general nighttime increases in benzene was attributed to a low nocturnal boundary layer. The high nighttime mixing ratios greatly increased the statistics for the site, and indicate a much larger benzene pollution problem than was suspected in the area. VOC monitoring in the WGF has primarily been conducted only during the daytime, and the high nighttime mixing ratios are pushing up the benzene to levels that are a concern for public health. In addition the ground measurements, aircraft measurements provided profiles of benzene and related compounds. These vertical diurnal profiles over PAO indicate that the benzene originates from a local ground source rather than being transported to the site, indicating that at short range O&NG benzene sources are more important than the more remote Denver signal.

Canister samples collected at the surface during the aircraft profiles allowed for general source signatures to be examined. Acetylene and propane were used as tracers for urban and natural gas sources. The PAO canisters showed an overall strong natural gas signature, similar to canister results collected at a nearby site in 2013; this was inferred from the i-pentane to n-pentane ratio. Additionally, acetylene and propane used as a tracer against benzene indicate that, whereas there are multiple sources for benzene measured in the canister samples at PAO, natural gas is a stronger source than the traffic sources.

Polar frequency analysis of the benzene measured in the PTR-MS shows that there is a directional component to the benzene mixing ratios. The highest mixing ratios are associated with the SW wind regime, while elevated mean and median benzene originates from the W and SW. This is a notable result because it indicates we are observing a directional source rather than an increase in background benzene in the gas field. Additionally, the high benzene measured from the SW at PAO corresponds to high benzene measured in previous studies at BAO, which saw elevated benzene mixing ratios from the NE at BAO. The
observation of this SW benzene source is a more certain spatial constraint on the benzene source in the
WGF.

In prior work based in the WGF the major benzene sources in the gas field have been attributed to
a combination of combustion engines and natural gas extraction. However, the benzene to CO ratios
measured at PAO are unexpectedly high, even compared to the high ratio of benzene to CO measured at
BAO and attributed to gas field benzene production (Petron et al., 2012; Swarthout et al., 2013). The most
likely sources of benzene at PAO are condensation tanks and fugitive emissions of benzene from natural
gas operations in the Platteville area; the most likely source of CO is a non-combustive industrial process
to the SW of the site. However, to better resolve the benzene source question we need more directed
measurements around the natural gas infrastructure in the region and long term stationary measurements
deep in the WGF.

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References


Air emissions case study related to oil and gas development in Erie, Colorado, 2012. Air Pollution Control Division Technical Services Program. Colorado Department of Public Health and Environment (CDPHE).


Swarthout, R.F., Russo, R.S., Zhou, Y., Hart, A.H., Sive, B.C., 2013. Volatile organic compound distributions during the NACHTT campaign at the Boulder Atmospheric Observatory: Influence of


### Table 1. Spatial statistics for benzene measured over the collection sites from 3 km ASL and below.

<table>
<thead>
<tr>
<th></th>
<th>Fort Collins</th>
<th>Chatfield</th>
<th>NREL/Golden</th>
<th>BAO</th>
<th>Denver/La Casa</th>
<th>PAO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>0.068</td>
<td>0.080</td>
<td>0.087</td>
<td>0.102</td>
<td>0.119</td>
<td>0.140</td>
</tr>
<tr>
<td>Median</td>
<td>0.064</td>
<td>0.071</td>
<td>0.085</td>
<td>0.095</td>
<td>0.110</td>
<td>0.115</td>
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<tr>
<td>25&lt;sup&gt;th&lt;/sup&gt;</td>
<td>0.036</td>
<td>0.038</td>
<td>0.045</td>
<td>0.053</td>
<td>0.067</td>
<td>0.062</td>
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<tr>
<td>75&lt;sup&gt;th&lt;/sup&gt;</td>
<td>0.095</td>
<td>0.110</td>
<td>0.122</td>
<td>0.139</td>
<td>0.166</td>
<td>0.190</td>
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<tr>
<td>Max</td>
<td>0.269</td>
<td>0.284</td>
<td>0.260</td>
<td>0.674</td>
<td>0.466</td>
<td>1.004</td>
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### Table 2. Quantitative statistics for benzene measured at PAO, units in ppbv.

<table>
<thead>
<tr>
<th></th>
<th>Campaign</th>
<th>Day</th>
<th>Night</th>
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<tbody>
<tr>
<td>Mean</td>
<td>0.53</td>
<td>0.37</td>
<td>0.74</td>
</tr>
<tr>
<td>Median</td>
<td>0.33</td>
<td>0.18</td>
<td>0.54</td>
</tr>
<tr>
<td>25&lt;sup&gt;th&lt;/sup&gt;</td>
<td>0.14</td>
<td>0.10</td>
<td>0.30</td>
</tr>
<tr>
<td>75&lt;sup&gt;th&lt;/sup&gt;</td>
<td>0.66</td>
<td>0.45</td>
<td>0.89</td>
</tr>
<tr>
<td>Max</td>
<td>29.3</td>
<td>20.6</td>
<td>29.3</td>
</tr>
</tbody>
</table>

### Table 3. Mean PAO toluene/benzene ratios by wind sector, with diurnal information.

<table>
<thead>
<tr>
<th>Wind Sector</th>
<th>Total</th>
<th>Day</th>
<th>Night</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>1.26 ± 0.62</td>
<td>1.16 ± 0.81</td>
<td>1.37 ± 0.28</td>
</tr>
<tr>
<td>NE</td>
<td>1.32 ± 0.52</td>
<td>1.03 ± 0.40</td>
<td>1.55 ± 0.49</td>
</tr>
<tr>
<td>E</td>
<td>1.03 ± 0.42</td>
<td>0.88 ± 0.36</td>
<td>1.25 ± 0.40</td>
</tr>
<tr>
<td>SE</td>
<td>1.09 ± 0.71</td>
<td>0.97 ± 0.70</td>
<td>1.36 ± 0.65</td>
</tr>
<tr>
<td>S</td>
<td>1.41 ± 1.86</td>
<td>1.13 ± 1.22</td>
<td>1.61 ± 2.19</td>
</tr>
<tr>
<td>SW</td>
<td>1.34 ± 0.48</td>
<td>1.06 ± 0.71</td>
<td>1.41 ± 0.38</td>
</tr>
<tr>
<td>W</td>
<td>1.29 ± 0.43</td>
<td>0.95 ± 0.64</td>
<td>1.39 ± 0.26</td>
</tr>
<tr>
<td>NW</td>
<td>1.33 ± 0.46</td>
<td>1.21 ± 0.41</td>
<td>1.42 ± 0.27</td>
</tr>
<tr>
<td>Calm</td>
<td>1.32 ± 0.41</td>
<td>1.02 ± 0.56</td>
<td>1.39 ± 0.32</td>
</tr>
</tbody>
</table>

### Table 4. PAO canister correlations compared to Platteville, CO, literature correlation values

<table>
<thead>
<tr>
<th></th>
<th>Propane</th>
<th>Acetylene</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-butane</td>
<td>0.95</td>
<td>0.99</td>
</tr>
<tr>
<td>i-pentane</td>
<td>0.91</td>
<td>0.99</td>
</tr>
<tr>
<td>n-pentane</td>
<td>0.94</td>
<td>0.97</td>
</tr>
<tr>
<td>Benzene</td>
<td>0.73</td>
<td>0.70</td>
</tr>
</tbody>
</table>
Figure 1. Map of the 2014 DISCOVER-AQ study area. The urban areas are shown in grey (data courtesy of the United States Census Bureau, http://www.census.gov/geo/maps-data/data/tiger.html). The boundary of the WGF is shown in black along with the gas wells (yellow points) (data courtesy of the Colorado Oil and Gas Conservation Commission, http://cogcc.state.co.us/). The DISCOVER-AQ ground sites are plotted and colored using the mean benzene measured from the aircraft spiral observations for each site. The benzene statistics were calculated using the bottom 1 km AGL for each site.
Figure 2. PTR-MS Benzene measurements collected at PAO during the campaign. A. The ground level benzene time series for the full campaign. B. Box and whisker benzene mixing ratios organized by day, midnight to midnight MDT. The outlier points are not shown. The open points are the daily means. C. The diurnal cycle of benzene over all observations. The outlier points are not shown for clarity, the open points are the hours mean values. All dates and times are in local (MDT) time.
Figure 3. The Benzene relationship to PBLH high (AGL). The purple triangles are PBLH calculated from
the ozonesondes launched at PAO, using 15 minute mean benzene data beginning from the launch time.
The gold points are the mean aerosol MPL calculated PBLH binned by 50 m, collected between 0700 and
1900 MDT. The grey boxes are the statistical box and whisker plots for the MPL PBLH data. The outlier
points for the box and whisker plots are excluded for clarity.
Figure 4. Vertical diurnal profiles of benzene over PAO. The aircraft data was binned into 50 m boxes and plotted as a box and whiskers plot. The plot is further separated by hour of day (local time, MDT). All benzene mixing ratios are cut off a 0.60 ppbv for clarity. A total of 29 total spirals and three partial spirals were used to construct this plot.
Figure 5. Total campaign polar frequency plots of benzene measured at PAO. The wind observations are binned by 10 degrees wind direction and 0.5 m/s wind speed. The wind speed and direction bins are plotted on a polar plot and colored based on the calculated statistic. Enhanced benzene has directional dependence and is associated with SW flow over the site, especially in higher mixing ratios (mean, maximum).
Figure 6. Canister VOC measurements of VOCs correlated with propane, an O&NG tracer, and acetylene, a traffic and combustion tracer. The combustion tracers i-pentane, n-pentane, n-butane and ethane show a stronger correlation with propane (A) than with acetylene (B), indicating a strong O&NG influence at PAO. Benzene also shows a strong correlation with propane (C) over acetylene (D), although the correlations with benzene in general have lower correlation constants compared to the combustion tracers.
Figure 7. Canister VOC measurements of i-pentane and n-pentane during the campaign, and literature ratios from NE Colorado studies. (\textsuperscript{a}Baker et al., 2009; \textsuperscript{b}Gilman et al., 2013; \textsuperscript{c}Swarthout et al. 2013; \textsuperscript{d}Thompson et al., 2014; \textsuperscript{e}This work, 2016)
Figure 8. Toluene to benzene spatial ratios calculated from the FRAPPE C-130 TOGA data. The data points over the map are colored by the toluene to benzene ratio, and sized by the measured benzene mixing ratio at each observation. PAO is marked with a cyan square.
Figure 9. Toluene to benzene correlation from the PTR-MS data. A. The correlation plot of benzene to toluene for the full campaign. The points are color coded by the toluene/benzene ratio. A toluene/benzene ratio $\geq 2$ is associated with fresh vehicle emissions and/or vehicles without catalytic converters. B. A polar frequency plot of the toluene/benzene ratio shows that these high ratio observations are associated with southerly winds.
Figure 10. Total campaign polar frequency plots of CO measured at PAO. The wind observations are binned by 10 degrees wind direction and 0.5 m/s wind speed. Enhanced CO has directional dependence associated with SW flow over the site, similar to the results for benzene (Figure 4). CO has a stronger wind direction dependence, indicative of a point source located in close proximity to PAO.
Figure 11. Benzene-CO correlations for all observations, faceted by general wind direction. Calm winds had a measured wind speed less than 1 m/s for the minute average.
Supplementary Figure 1. PBLH diurnal profile. The box and whisker plots are created from the MPL PBLH calculations from 22 July 2014 through the end of 11 August 2014. The open points show mean MPL PBLH for each hour. The red points show mean ozonesonde PBLH for each hour. At 1000 MDT both the mean and the median of the PBLH calculated from the MPL are ≥ 0.5 Km AGL, making 1000 MDT the beginning of the daytime-type boundary layer.
Supplementary Figure 2. Vertical benzene profiles and ground benzene measurements collected at PAO during the study. Each point is plotted based on the time stamp and GPS altitude data from the P3-B spirals. The altitudes are reported above ground level (AGL). Each point is colored by measured benzene mixing ratio. Low aircraft measurements of benzene contrast with the high ground benzene measured throughout the campaign.
Supplementary Figure 3. Benzene mean profile over PAO from the vertical aircraft spirals. The points are the mean mixing ratios binned by 50 m bins. The error bars show the standard deviation of the mean for each vertical bin.
Supplementary Figure 4. Canister benzene correlated with 2min mean benzene, with the mean benzene window centered on the canister opening time. The grey dashed line shows the 1-to-1 line. The black lines shows the linear regression best fit line, which has a slope of 1.003, and $R^2 = 0.9205$. The points are colored by their hour of collection, in local time (MDT).
Supplementary Figure 5. Waste pit locations and statuses provided by COGCC. The cyan points show the locations of the PAO and BAO sampling sites for the campaign. The black outline shows the political boundary of the WGF as defined by the COGCC. The red triangles show the locations of the active waste pit locations as reported from the COGCC data.