Calibration and Field Testing of Cavity Ring-Down Laser 1 Spectrometers Measuring CH₄, CO₂, and δ^{13} CH₄ Deployed on 2 **Towers in the Marcellus Shale Region** 3

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18 Abstract. Four in-situ cavity ring-down spectrometers (G2132-i, Picarro, Inc.) measuring methane dry mole 19 fraction (CH₄), carbon dioxide dry mole fraction (CO₂) and the isotopic ratio of methane (δ^{13} CH₄) were deployed at 20 four towers in the Marcellus Shale natural gas extraction region of Pennsylvania. In this paper, we describe 21 laboratory and field calibration of the analyzers for tower-based applications, and characterize their performance in 22 the field for the period January – December 2016. Prior to deployment, each analyzer was tested using bottles with 23 various isotopic ratios, from biogenic to thermogenic source values, which were diluted to varying degrees in zero 24 air, and an initial calibration was performed. Furthermore, at each tower location, three field tanks were employed, 25 from ambient to high mole fractions, with various isotopic ratios. Two of these tanks were used to calibrate the 26 analyzers on a daily basis. A method to correct for cross interference from ethane is also described. Using an 27 independent field tank for evaluation, the standard deviation of 4-hour means of the isotopic ratio of methane 28 difference from the known value was found to be 0.26 $\% \delta^{13}$ CH₄. Following improvements in the field tank testing 29 scheme, the standard deviation of 4-hour means was 0.11 ‰, well within the target compatibility of 0.2 ‰. Round 30 robin style testing using tanks with near ambient isotopic ratios indicated mean errors of -0.14 to 0.03 % for each of 31 the analyzers. Flask to in-situ comparisons showed mean differences over the year of 0.02 and 0.08 ‰, for the East 32 and South towers, respectively. 33

34 Regional sources in this region were difficult to differentiate from strong perturbations in the background. During

35 the afternoon hours, the median differences of the isotopic ratio measured at three of the towers, compared to the

36 background tower, were -0.15 to 0.12 ‰ with standard deviations of the 10-min isotopic ratio differences of 0.8 ‰.

- In terms of source attribution, analyzer compatibility of 0.2 $\% \delta^{13}$ CH₄ affords the ability to distinguish a 50 ppb CH₄ 37
- 38 peak from a biogenic source (at -60 ‰, for example) from one originating from a thermogenic source (-35 ‰), with

- 1 the exact value dependent upon the source isotopic ratios. Using a Keeling plot approach for the non-afternoon data
- 2 at a tower in the center of the study region, we determined the source isotopic signature to be -31.2 ± 1.9 ‰, within
- 3 the wide range of values consistent with a deep-layer Marcellus natural gas source.

1 1 Introduction

2 Quantification of regional greenhouse gas emissions resulting from natural gas extraction activities is critical for 3 determining the climate effects of natural gas usage compared to coal or oil. Studies have shown that the emission 4 rates as a percentage of production vary significantly from reservoir to reservoir. An aircraft-based mass balance 5 study in the Uintah basin in Utah (Karion et al., 2013; Rella et al., 2015) found a methane emission rate of 6.2–11.7 6 % of production, exceeding the 3.2 % threshold for natural gas climate benefits compared to coal determined by 7 Alvarez et al. (2012). In the Denver-Julesburg basin in Colorado, Pétron et al. (2014) found an emissions rate of 4 8 % of production, again using an aircraft mass balance approach. The Barnett Shale, one of the largest production 9 basins in the United States with 8 % of total U.S. natural gas production, was found to exhibit a lower emission rate 10 of 1.3–1.9 % (Karion et al., 2015). Using a model optimization approach for aircraft data, Barkley et al. (2017) 11 found the weighted mean emission rate from unconventional natural gas production and gathering facilities in the 12 Marcellus region in northeastern Pennsylvania, a region with mostly dry natural gas, to be only 0.36 % of total gas 13 production.

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15 Aircraft-based studies cover large areas, but the temporal coverage is limited. Tower-based networks offer a 16 complementary approach, making continuous measurements over long periods of time. At the Boulder Atmospheric 17 Observatory (BAO) tall tower, daily flask measurements are found to contain enhanced levels of methane and other 18 alkanes, compared to the other tall towers in the National Oceanic and Atmospheric Administration (NOAA) 19 network (Pétron et al., 2012). Tower measurements allow for continuous measurements in the well mixed boundary 20 layer which are influenced by both nearby sources and the integrated effect of the upstream emissions. While towers 21 provide near continuous coverage of regional emissions, specific emissions sources with specific isotopic signatures 22 are often diluted by mixing, making the differences from background very small.

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24 Differentiating CH_4 emissions from natural gas activities from other sources (e.g., wetlands, cattle, landfills) is key 25 to documenting the greenhouse gas impact of natural gas production and to evaluate the effectiveness of emissions 26 reduction activities. The isotopic ratio of methane (δ^{13} CH₄) is particularly useful in this regard (Coleman et al., 27 1995). In general, heavy isotope ratios are characteristic of thermogenic CH_4 sources (i.e., fossil-fuel based) and 28 light isotope ratios are characteristic of biogenic sources (Dlugokencky et al., 2011). Schwietzke et al. (2016) 29 compiled a comprehensive database of isotopic methane source signatures, indicating signatures of -44.0 ‰ for 30 globally averaged fossil-fuel sources of methane, $-62.2 \ \%$ for globally averaged microbial sources such as wetlands, 31 ruminants, and landfills, and -22.2 ‰ for globally averaged biomass burning sources. Atmospheric measurements 32 of δ^{13} CH₄ have been used to partition emissions of CH₄ into source categories (e.g., Mikaloff Fletcher et al., 2004a,b; 33 Kai et al., 2011). It is important to note, however, that for fossil-fuel sources of methane, isotopic ratios of methane 34 vary significantly from reservoir to reservoir (e.g., Townsend-Small et al., 2015; Rella et al., 2015), and with depth 35 in a single reservoir (Molofsky et al., 2011; Baldassare et al., 2014). 36

1 The isotopic ratio of methane has traditionally been measured in the laboratory with continuous flow gas 2 chromatography/ isotope ratio mass spectrometry, with repeatability of $\pm 0.05\%$ (Fisher et al., 2006). Röckmann et 3 al. (2016) recently compared continuous in-situ measurements of methane isotopic ratio using a dual isotope mass 4 spectrometric system (IRMS) and a quantum cascade laser absorption spectroscopy (QCLAS)-based technique at the 5 Cabauw tower site in the Netherlands. They showed that high-temporal-resolution methane isotopic ratio data can 6 be used in conjunction with a global and a mesoscale model to evaluate CH₄ emission inventories. Röckmann et al. 7 (2016) also used a moving Keeling plot approach to identify source isotopic ratios.

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9 Cavity ring-down spectroscopy (CRDS) is another technique for measurement of continuous in-situ isotopic ratio of 10 methane (Rella et al., 2015). CRDS is a laser-based technique in which the infrared absorption loss caused by a gas 11 in the sample cell is measured to quantify the mole fraction of the gas. The analyzers utilize three highly reflective 12 mirrors such that the flow cell has an effective optical path length of 15–20 km, allowing highly precise 13 measurements. The temperature and pressure of the sample cell is tightly controlled, improving the stability of the 14 measurements (Crosson 2008). Rella et al. (2015) documented the operation of CRDS (Picarro, Inc., model G2132-15 i) analyzers, including cross-interference from other gases, and general calibration approach.

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Furthermore, Rella et al. (2015) described the use of two tanks to correct for analyzer drift of the isotopic ratio measured by the G2132-i analyzers. In this approach, the variables of interest, i.e., the total methane mole fraction and the isotopic ratio, are directly calibrated. The drift terms in the calibration equations have differing dependence on mole fraction, requiring the use of at least two tanks for calibration. For this study, three field tanks were deployed at each tower location, two for the field calibration and one as an independent test.

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23 In this paper, we describe a network of four tower-based atmospheric observation locations, measuring CH_4 and 24 CO₂ dry mole fractions and δ^{13} CH₄ using CRDS (Picarro, Inc., model G2132-i) analyzers in the Marcellus shale 25 region in north-central Pennsylvania at towers referred to here as the North, South, East and Central towers. We 26 focus on the specific application of tower-based measurements of isotopic methane using CRDS analyzers. Instead 27 of describing the methods and results separately, we combine these for each topic. First, we describe laboratory 28 calibration of the G2132-i analyzers, field calibration approach, and calibration results. We determine the 29 compatibility achieved for the isotopic measurements in the current field deployment, using an independent field 30 tank, round-robin style testing and comparisons to flasks as our primary metrics. We also evaluate the performance 31 of the G2132-i analyzers in terms of CH_4 and CO_2 mole fractions measurements by comparing to a G2301 analyzer. 32 We then describe the tower locations and compare differences in CH₄ mole fraction and isotopic ratio observed at 33 the towers and use a Keeling plot approach to determine source isotopic signatures. Finally, we describe 34 recommendations for future isotopic methane CRDS tower-based networks.

1 2 Compatibility goals

Because this is the first network of multiple isotopic ratio of methane continuous analyzers to date, the needed compatibility has not yet been defined. Thus, our compatibility goals for CO_2 and CH_4 mole fractions follow the WMO compatibility recommendation for global studies: 0.1 ppm for CO_2 (in the Northern Hemisphere) and 2 ppb for CH_4 (GAW Report No. 229, 2016). Here we use the term compatibility, as advised in the GAW Report No. 229 (2016), to describe the difference between two measurements, rather than the absolute accuracy of those measurements.

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9 For $\delta^{13}CH_4$ we set our target compatibility at 0.2 ‰, thought to be a reasonable goal based on laboratory testing 10 prior to deployment and the results shown in Rella et al. (2015). This goal corresponds to the WMO extended 11 compatibility goal for the isotopic ratio of methane, which was deemed sufficient for regionally focused studies with 12 large local fluxes. The measured signal at the towers is a mixture of the source and the background (Pataki et al., 13 2003), and the ability to distinguish between a biogenic and thermogenic source depends on the difference of the 14 source isotopic signature from background and the peak strength in terms of methane mole fraction. Equating the 15 slope of a source and the background with the slope of a mixture and the background on a Keeling plot (Keeling, 16 1961), the measured isotopic ratio difference ($\Delta\delta$) is given by

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18
$$\Delta \delta = (\delta_{\rm src} - \delta_{\rm back}) \frac{\Delta CH_4}{CH_{4,\rm meas}} , \qquad (1)$$

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20 where δ_{src} and δ_{back} are the isotopic ratios of the source and the background, $CH_{4,meas}$ is the measured methane 21 mole fraction, and ΔCH_4 is the difference between the measured mole fraction and the background. This equation is 22 represented graphically in Fig. 1. If there are two possible sources in a region, a biogenic source at -60 ‰ and a 23 thermogenic source at -35 ‰, for example, the difference in isotopic ratio difference is at least three times the 24 compatibility goal of 0.2 % (and thus distinguishable) for a peak strength of 50 ppb CH₄ or greater, assuming a 25 measured CH_4 mole fraction of 2000 ppb and a background isotopic ratio of -47.5 %. In this case, the biogenic 26 source would measure 0.3 % above the background, as opposed to the thermogenic source measuring 0.3 % below 27 the background. As shown in Fig. 1, sources closer to the background in isotopic ratio require a larger peak in CH_4 28 and those further from the background can be attributed with a smaller peak in CH₄.

29 3 Allan standard deviation testing

Allan standard deviation testing (Allan, 1966) is a useful tool for testing the noise and drift response of instrumentation. The Allan standard deviation for each averaging interval is proportional to the range of values for each averaging interval. This range typically decreases for increasing averaging interval, as the noise is reduced through averaging. As the averaging interval increases, however, analyzer drift may contribute, placing an upper bound on the optimal averaging interval. Thus, the Allan deviation results are critical for defining the minimum averaging time required for a given target compatibility.

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To calculate the Allan standard deviation of the G2132-i analyzers used in this study, one tank containing an ambient mole fraction of CH_4 (1.9 ppm), and CO_2 (~400 ppm) mole fraction and one tank containing high mole fraction of CH_4 (9.7 ppm) and an ambient mole fraction of CO_2 (~400 ppm) were tested with an analyzer for 24 hours. For simplicity, we call these the "high" and "low" tanks, respectively, and they are described further in Section 5.1. We tested both as the noise is known to be less for higher mole fractions, and at least one tank with higher CH_4 mole fraction is necessary for the isotopic ratio calibration (Rella et al., 2015).

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The resulting Allan standard deviations for δ^{13} CH₄, CH₄ and CO₂ are shown in Fig. 2. For the high tank, the Allan deviation for δ^{13} CH₄ (Fig. 2A) was < 0.2 ‰ (our target compatibility) for an averaging interval of 2 min (the averaging interval used each field calibration cycle of the high tank). To reduce the noise to < 0.1 ‰, an averaging interval of 4 min is sufficient (in addition to the time required for the transition between gases). For the low tank, in order for the Allan standard deviation to be < 0.2 ‰, 32 min were required and 64 min for 0.1 ‰ noise. Note that for much of the deployment, the near ambient mole fraction target tank was not sampled sufficiently within each day

- 17 for the desired compatibility goals.
- 18

19 For CH_4 (Fig. 2B), both the high and low tank Allan deviation were < 1 ppb for even a 1-min averaging interval.

20 The CO₂ levels in the high and low tanks were similar (~400 ppm), and an averaging interval of 6 min corresponded

21 to Allan standard deviations of 0.3 ppm, and 64 min were necessary for 0.1 ppm (Fig. 2C). The performance of the

22 G2132-i analyzers in terms of CO₂ precision is worse than that of the G2301/G2401 analyzers primarily because a

23 weaker spectral line is used (Rella et al., 2015).

24 4 Laboratory calibration

25 4.1 Experimental set-up

26 Prior to field deployment, each analyzer was calibrated for CH₄ and CO₂ mole fraction. Four NOAA-calibrated

tertiary standards (traceable to the WMO X2004 scale for CH_4 and the WMO X2007 scale for CO_2) were used for

28 the linear mole fraction calibration, as described in Richardson et al. (2017). These NOAA tertiary standards ranged

between 1790 and 2350 ppb CH_{4} , and between 360 and 450 ppm CO_2 .

30 To calibrate the $\delta^{13}CH_4$ measurement prior to deployment, four different target mixing ratios, each at four different

- 31 known isotopic ratios were tested by the four analyzers using the experimental setup in Fig. 3. Commercially-
- 32 available isotopic standard bottles (Isometric Instruments, Inc., product numbers L-iso1, B-iso1, T-iso1 and H-iso1)
- 33 were diluted with zero air to produce mixtures with varying CH₄ mixing ratios and δ^{13} CH₄. The gravimetrically-

1 determined zero air (Scott Marrin, Inc.) was natural ultra-pure air, containing no methane or other alkanes but 2 ambient levels of CO₂. The isotopic calibration standard bottles each contained approximately 2500 ppm of CH_4 at 3 -23.9, -38.3, -54.5, and $-66.5 \ \% \ \delta^{13}$ CH₄, with uncertainty of $\pm 0.2 \ \%$ reported by the supplier. These isotopic ratios 4 were tied to the Vienna Pee Dee Belemnite (VPDB) scale. Mass flow-controllers (MC-1SCCM and MC-500SCCM, 5 Alicat Scientific, Inc.) and a 6-port rotary valve (EUTA-2SD6MWE, Valco Instruments Co., Inc.) were used to 6 direct the standard bottle air for each isotopic calibration standard bottle into a mixing volume (~ 4 m of 1/8 in, 0.32 7 cm OD stainless steel tubing; TSS285-120F, VICI Precision Sampling, Inc.) at 0.400 sccm and mixed with zero CH₄ 8 air at 137, 161, 303, and 555 sccm to create target CH_4 mole fractions of 7.3, 6.2, 3.3, and 1.8 ppm, respectively. 9 Thus 16 CH₄ mole fraction/isotopic ratio pairs were produced. The accuracy of the mass flow controllers can be a 10 significant source of error in making mixtures. Here the nominal range of the mass flow controllers was 1 sccm for 11 the standard bottle line and 500 sccm for the zero air line, and the accuracy was ± 0.2 % of full scale. To avoid 12 isotopic fractionation at the head of the low-flow mass flow controller, the flow of the zero air was varied rather than 13 the isotope standard. It is possible that fractionation did occur due to the tees used to direct gas into the individual 14 analyzers. For this reason, it would have been preferable to set up the analyzers to each sample directly from the 15 mixing volume.

The first mixture of each isotopic standard was tested for 60 minutes to flush out the span gas line and to avoid isotopic fractionation at the head of the span mass flow controller. Subsequent dilutions using the same isotopic standard were tested for 20 minutes each and each dilution was repeated twice. With the flow rate of 0.400 sccm for the isotopic standard bottles, the total volume of standard gas used was 88 cc. Observations were collected at ~0.5 Hz and the final 5 minutes of data for each dilution were averaged to compare against the target value. The standard deviation of the raw data collected during these tests (Fig. 4) decrease exponentially with increasing mole fraction.

Averaged methane isotopic ratios *prior to calibration* are shown in Fig. 5. There is an offset in the measured isotopic ratio as a function of the changing known isotopic ratio. For higher mole fractions, this offset is fairly constant, but for near ambient mole fractions it is analyzer-specific. We note that the precision of these results could be improved by averaging over longer periods. We now describe the calibration technique to remove these offsets.

26 4.2 Application of calibration equations

The first step in the calibration process for the analyzers is to remove the nearly linear error that is a function of isotopic ratio. We applied methods leading from the theoretical framework developed by Rella et al. (2015) to calibrate the isotopic ratio data. Applying a linear fit to highest mole fraction values (7.3 ppm) measured in the laboratory for known δ^{13} CH₄ values (-23.9, -38.3, -54.5, -66.5 ‰) for each analyzer, we determined the linear

- 31 calibration coefficients p_1 and p_0 .
- 32

33
$$[\delta^{13}CH_4]_{intermediate} = p_1[\delta^{13}CH_4]_{measured} + p_0.$$
(2)

For this step, we used only the highest mole fraction values because δ^{13} CH₄ is more precise for higher mole fractions (Fig. 4). We note that these laboratory tests were completed prior to the Allan standard deviation testing and that the averaging times were not sufficient to achieve the desired compatibility at ambient mole fractions. Ambient mole fractions could be used for this step if measured for sufficient durations.

5

To correct for the CH₄ mole fraction dependence of the measured δ^{13} CH₄, the two time-dependent drift parameters described in Rella et al. (2015) c_0 and χ must be determined. Here c_0 varies because of spectral variations in the optical loss of the empty cavity and χ varies because of errors in the temperature or pressure of the gas, or changes in the wavelength calibration. These parameters are defined in Eq. (15) of Rella et al. (2015). A coefficient describing the changes in the crosstalk between the two methane isotopologues was ignored, following Rella et al. (2015). For the laboratory calibration, we determined c_0 and χ using measurements at -23.9 ‰ for a high mole fraction (7.3 ppm) and a low mole fraction (1.8 ppm). We then applied Eq. (12) of Rella et al. (2015)

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$$[\delta^{13}CH_4]_{calibrated} = [\delta^{13}CH_4]_{intermediate} + \frac{c_0}{c_{12}} + \chi([\delta^{13}CH_4]_{intermediate} - B),$$
 (3)

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16 to correct for the CH₄ mole fraction dependence of δ^{13} CH₄. Here c_{12} is the measured [¹²CH₄] and

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$$18 \qquad B = p_1 B_{default} + p_0,$$

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with $B_{default}$ being -1053.59 ‰. $B_{default}$ is the intercept of the fit of the isotopic ratio to the ratio of the absorption peak heights for the standard calibration and *B* is the updated value, specific to the analyzer. We followed Rella et al. (2015) and ignored the contribution of an additional offset term that depends on neither mole fraction nor isotopic ratio. Note that the slope of the linear calibration was the only component of the calibration that was not adjusted in the field using field tanks (Section 5.4).

(4)

25 5 Methods: Field deployment

26 5.1 In-situ field tanks

At each tower site, three field tanks were utilized, as listed in Table 1. One tank at each tower site was calibrated by the National Oceanic and Atmospheric Administration (NOAA) for CH_4 and CO_2 mole fractions and by the Institute of Arctic and Alpine Research (INSTAAR) for $\delta^{13}CH_4$. This tank was tested quasi-daily (every 21 hours) and used to adjust the intercept for the CH_4 and CO_2 mole fraction calibrations (Richardson et al., 2017). The constituents of

31 this tank were at typical ambient levels (as listed in Table 1), and for the purposes of this paper, we call it the

32 "target", although it was not independent.

- 1 Two additional tanks were tested at each of the tower sites (Table 1). These tanks were filled using ultra-pure air
- 2 and spiked (using Isometric Instruments, Inc bottles) by Scott Marrin, LLC, (one at 1.9–2.1 ppm CH_4 and –23.9
- 3 $\% \delta^{13}$ CH₄) and one at 9.7–10.5 ppm CH₄ at –38.3 $\% \delta^{13}$ CH₄). Recall that these are called the "low" and "high"
- 4 tanks, for simplicity. These tanks contained ambient levels of CO_2 (368 407 ppm). The choice of the CH_4 mole
- 5 fraction of the high tank is based on the optimal determination of the calibration coefficients c_0 and χ , rather than
- 6 the expected range of ambient CH_4 mole fractions. The effect of c_0 on the calibrated isotopic ratio is largest at low
- 7 mole fractions, whereas the effect of χ is independent of mole fraction. Thus the ratio of the high and low tank mole
- 8 fractions determines how separable the two effects are. We therefore chose the high tank mole fraction to be as high
- 9 as possible without introducing other nonlinearities into the system.
- 10 The high and low tanks for each tower were calibrated for δ^{13} CH₄ in the laboratory prior to deployment. First we 11 applied a linear calibration for δ^{13} CH₄ using measurements from each of four Isometric Instruments bottles (-23.9, -12 38.3, -54.5, -66.5 ‰), diluted with zero air to 10.3 - 10.4 ppm CH₄. A 3-way solenoid valve (091-0094-900, 13 Parker Hannifin Corp.) was used just downstream of the mixing volume in the laboratory calibration system to stop 14 flow from the zero air tank and Isometric Instrument bottles and allow flow from the working standards. Then a 15 mole fraction correction was applied using the -23.9% bottle diluted to 10.4 ppm CH₄ and the -38.3% bottle 16 diluted to 1.9 ppm. These calibration results are shown in Table 1. The values assigned to the tanks differed slightly 17 (with the differences ranging in magnitude from 0.01 to 0.38 %) from the bottles used for spiking. Possible reasons 18 for these slight differences include noise in the measurement, fractionation upon tank-filling, bottle assignment error 19 with the 0.2 ‰ uncertainty reported by the supplier (Isometric Instruments, Inc.) and insufficient testing times for 20 the tanks at ambient mole fractions (5 min). We note that it would have been preferable to utilize calibration tanks 21 closer to the observed air samples in terms of isotopic ratio. In particular, the low tank could have been spiked with 22 the -38.3 ‰ bottle, or a mixture of the -38.3 and -54.5 ‰ bottles.

23 5.2 In-situ field calibration gas sampling system

24 The flow diagram of the field calibration system is shown in Fig. 6. Polyethylene/aluminum composite tubing (1/4 25 in, 0.64 cm OD, Synflex 1300, Eaton Corp.) was used to sample from the top of each tower for the CRDS analyzer 26 and a separate sample line made from 3/8 in (0.95 cm) OD Synflex 1300 tubing was used for the flask sampling 27 packages. The top end of each tube was equipped with a rain shield to prevent liquid water from entering the 28 sampling line. For the CRDS analyzer, air was drawn down the tube at 1 L/min, with 30 cc/min flow into the 29 analyzer and the remainder purged. The residence time in the tube was about 1 min. Separate tubes were used for 30 the CRDS and flask sampling lines because of the differing flow rates required for the flask samples (varying 31 between 0.29 and 3.8 liters per minute) (Turnbull et. al., 2012) and to ensure independence of the CRDS and flask 32 measurements.

2 using a 6-port rotary valve (EUTA-2SD6MWE, Valco Instruments Co, Inc.). Stainless steel tubing (1/8 in, 0.32 cm 3 OD, TSS285-120F, VICI Precision Sampling, Inc.) and single-stage regulators (Y11-C444B590, Airgas, Inc.) were 4 used for testing the field tanks. Rella et al. (2015) noted that the effect of water vapor on the isotopic ratio of 5 methane measurement is up to 1 % and nonlinear, and recommended drying to less than 0.1% H₂O mole fraction. 6 Thus we used a Nafion dryer (MD-070-96S-2, PermaPure) in the reflux configuration, with an additional pump 7 (ME1, Vacuubrand, Inc.) on the outlet of the Nafion dryer (Fig. 6). The sample air was dried to ~ 0.06 % H₂O and 8 the calibration gases were humidified to 0.02 % H_2O_2 , in a manner similar to Andrews et al. (2014). The CH₄ mole 9 fraction was corrected for water vapor following Rella et al. (2015 supp), and the CO₂ mole fraction following Chen 10 et al. (2010).

For the continuous in-situ measurement system, switching between sample and calibration gases was accomplished

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12 A cycle including 90 min of ambient sampling, 6 min testing the high mole fraction field tank, and 10 min testing 13 the low mole fraction field tank was repeated 12 times, then the target tank was tested for 10 min (occurring every 14 \sim 21 hours, to test for diurnal effects). Thus, there were 13.5 calibration cycles for the high and low tanks each day, 15 on average. The first 4 min of data were discarded each time after switching gases to ensure sufficient flushing of 16 the sample cell. After this time, the CO_2 and CH_4 mole fractions stabilized. The ideal calibration tank testing time 17 is a balance between minimizing calibration gas usage (and consequently maximizing ambient air sampling time) 18 and achieving sufficient precision. Note that the Allan standard deviation results indicate that testing for 4 min for 19 the high tank and for 32 min for the low and target tanks is required to achieve our target compatibility of 0.2 ‰ 20 δ^{13} CH₄. Thus, this averaging time was achieved in two calibration cycles for the high tank (excluding flushing 21 time), but in 5.3 calibration cycles for the low and target tanks (completed in about 10 hours in the case of the 22 sampling scheme utilized for most of the deployment). An improved sampling strategy was implemented on 3 23 December 2016 and is discussed in Section 5.4.

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The flow rate of the instruments was 35 cc/min, and the 150A tank size was used, corresponding to 4.021×10^6 cc at standard pressure and temperature. Thus there was sufficient gas to test each tank for about one hour per day for

about five years, as a general guideline.

28 5.3 Cross-interference from other species

29 5.3.1 Overview

30 The effects of cross-interference from other species must be considered for spectroscopic measurements. Rella et al.

31 (2015) give proportional relationships for cross-interference from various species for the G2132-i analyzers. Listed

32 in Table 2 are species with potential to affect the isotopic methane calibration, and their estimated effects for tower-

- 33 based applications. We based these estimates on typical maximum values determined by flask (level at which 99 %
- 34 of flask measurements at the South and East towers were below; for carbon monoxide, propane, butane, ethylene,

and ethane), by in-situ measurements at the towers in this deployment (for water vapor and carbon dioxide), and by
 typical values (Warneck and Williams, 2012; for ammonia and hydrogen sulfide). There are no known ambient

3 estimates for methyl mercaptan (Barnes, 2015), so the odor threshold (Devos et al., 1990) was used as a maximum

4 5

value.

For the Picarro G-2132i analyzers, ethane contributed the largest interference and a correction to the isotopic ratio
was applied (Section 4.4.2). Because of water vapor effects, the sample was dried and the calibration gases
humidified. The effects of other species were neglected.

9 5.3.2 Ethane correction

Ethane (C_2H_6) is co-emitted with methane during natural gas extraction and its cross-interference with the isotopic ratio of methane is significant. The magnitude of the effect of ethane on the isotopic methane is proportional to its mole fraction and inversely proportional to the methane mole fraction. The two Scott-Marrin field tanks at each site were scrubbed of alkanes (including ethane), but the one NOAA/INSTAAR field tank at each site contained ambient levels of these species. Typical mole fractions of C_2H_6 (1.3 ppb) compared to the Scott-Marrin tanks containing no ethane would lead to a 0.04‰ bias, if uncorrected. Furthermore, flask measurements at the South and East towers indicated ethane up to 8 ppb, which corresponds to a 0.23 ‰ error.

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18 The G2132-i analyzers reported an ethane measurement, but were not designed for high-compatibility $C_{2}H_{6}$ 19 measurements at levels near background. In this deployment, 99 % of the flask measurements, which were taken in 20 the afternoon, were less than 8.0 ppb C_2H_6 . In comparison, the drives near natural gas sources conducted by Rella et 21 al. (2015) indicated C_2H_6 mole fractions up to 13 ppm (note unit change). The ethane signal is subject to strong 22 cross-interference from water vapor, methane and carbon dioxide. Rella et al. (2015; Eq. (S20)) report coefficients 23 for these corrections. These coefficients indicate corrections larger in magnitude than the ethane mole fractions 24 measured in this deployment. We have thus not attempted to analyze the ethane results themselves. The ethane 25 output was however used to correct the isotopic methane data. To do so, we first developed a linear calibration using 26 the Scott-Marrin high field tank containing zero ethane and the NOAA/INSTAAR target tank which we assumed 27 contained a background level of 1.5 ppb ethane (Peischl et al., 2016). This calibration is clearly a rough estimate. 28 Note that we determined the linear relationship between the reported ethane of each analyzer and its calibrated value 29 initially, and assumed that this relationship does not change throughout the deployment. Newer models of the 30 δ^{13} CH₄ analyzer (G2210-i, Picarro Inc.) measure C₂H₆ at ppb levels, simplifying this correction process.

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We then corrected the isotopic methane for the effects of ethane cross-interference. For example, 1.3 ppb of ethane in an air sample of 2 ppm CH₄ would, if uncorrected, shift the δ^{13} CH₄ measurement higher by [+58.56 ‰ ppm

34 $CH_4(ppm C_2H_6)^{-1} \ge [0.0013 ppm C_2H_6]/[2 ppm CH_4] = +0.04 \%$. Note that the calibration coefficient for ethane has

- 1 been updated from that indicated in Rella et al. (2015). The correction to compensate for this error was applied to
- 2 all data, using the estimated ethane and measured methane values.

3 5.3.3 Water vapor and carbon dioxide

Water vapor can have a significant effect on the measurements of isotopic methane (up to ± 1 ‰ for up to 2.5 % H₂O) (Rella et al., 2015). Thus, the sample air was dried and the calibration gases slightly humidified such that this effect is minimized (estimated to be < 0.02 ‰). For the range of ambient CO₂ observed in this study (~375 - 475 ppm), the difference from the calibration gases was ~100 ppm, and the effect was estimated to be < 0.03 ‰ (Table 2). The isotopic ratio of methane was thus not corrected for CO₂ effects.

9 5.3.4 Oxygen, argon, and carbon monoxide

10 The ambient variability in oxygen, argon, and carbon monoxide is expected to have a negligible effect on the 11 isotopic ratio measurements (Rella et al., 2015) and no corrections for these constituents were applied to the isotopic 12 methane data.

13 5.3.5 Other species

14 Ammonia, hydrogen sulfide, methyl mercaptan, propane, butane, ethylene are components of natural gas, but their 15 cross-interference effects were small for our tower-based application for which the sources are relatively far from 16 the measurement location. The effects of these species may be significant for other applications, such as 17 automobile-based measurements. Like for ethane, the magnitude of the effect of these gases on the isotopic methane 18 is proportional to the mole fraction of the contaminant species and inversely proportional to the methane mole 19 fraction. In Table 2, maximum mole fractions from the flasks if available, or typical mole fractions from the 20 literature, were used to estimate the effect of these species for our application. The cross-interference from these 21 species was insignificant for our application, < 0.01 ‰.

22

23 5.4 Field calibration

The linear calibration was determined in the laboratory as described in Section 4.2. We then used the daily average of the high and target field tanks to adjust the mole fraction correction (terms c_0 and χ) for the field data. The low tank was used as an independent test. For October 2016, the mean errors for the low tank at the South tower are 0.2 \pm 0.7 ‰, for example (Table 3, SCHEME B). Here the standard deviation was calculated using all of the calibration cycles during the month. The errors near the isotopic ratio of the target tank are likely less in magnitude. Instead using the low tank in the calibration and keeping the target tank independent yielded similar magnitudes of errors

(Table 3, SCHEME A), but minimized bias near the low tank (about -23.9 ‰) rather than near the target tank (about -47.2 ‰). Therefore, despite increased testing of the low tank throughout the majority of the deployment, we chose to use the target tank in the calibration to minimize errors near ambient isotopic ratios.

4

5 On 3 December 2016, an improved tank testing strategy was implemented, in which the target tank testing time was 6 increased from 6 min/day to 54 min/day (excluding transition times), achieved by sampling for 20 min every 420-7 min cycle (3.4 times/day, on average). The calibration times were completed using multiple cycles in order to avoid 8 not sampling the atmosphere for long periods and to measure possible changes in analyzer response throughout each 9 day. The low tank was tested using an identical strategy (20 min every 420-min cycle), with the total amount of 10 testing time per day changing from 81 min to 54 min. The high tank was tested on average 1.7 times per day (every 11 840 min) for 10 min. Excluding the transition times, the high tank testing time was thus reduced from 26 min/day 12 to about 10 min/day. Following the implementation of the improved strategy, the mean error of the independent low 13 tank at the sites was similar but the standard deviation was reduced from 0.5 to 1.3 ‰ to 0.3 to 0.9 ‰ (Table 3).

14

As an example of the effects of calibration, the tank results (differences from known values) using only the laboratory calibration for isotopic ratio, and following the SCHEME B are shown in Fig. 7 for the period September – December 2016. For the results using only the laboratory calibration, analyzer drift is apparent for all three tanks. Without a field calibration, the isotopic ratio was biased by up to 2 ‰. The target tank measurement was used in the calibration; hence the apparent drift following final calibration was necessarily zero. The noise apparent in Fig. 7B prior to 3 December 2016 when the calibration scheme was improved is at least partially due to insufficient sampling times of the target tank.

22

23 The relative effects of the calibration terms are illustrated in Fig. 8. The terms c_0 (Fig. 8A) and χ (Fig. 8B) in Eq. 24 (3) are time-dependent drift terms. These terms vary because of spectral variations in the optical loss of the empty 25 cavity (c_0) , and because of errors in the temperature or pressure of the gas, or changes in the wavelength calibration 26 (χ). Recall that the parameters c_0 and χ were calculated following Eq. (15) in Rella et al. (2015). The calculation of 27 the parameter c_0 used measurements from the high and target tank. The calculation of the parameter χ used 28 measurements of the high tank and was not independent from p_0 . The largest calibration effect was from the c_0 29 term, which increased the calibrated isotopic ratios by -0.5 to 4 ‰ during September to December 2016. The χ 30 term increased the final calibrated isotopic ratios by a smaller amount, -0.6 to 0.2 %. Thus over this period, there 31 were large changes in the calibration effect of these terms, although no software or hardware changes were applied. 32 Considering shorter term changes, the day to day changes in the calibration were less than 0.5 % for December 33 2016. Less frequent calibrations, e.g., twice per week, could be considered, but the reduction in field tank use is not 34 large considering the low flow rates of the instruments and steady changes up to 2 ‰ in the raw data over the time 35 scale of days were observed in Rella et al. (2015).

1 6 Evaluation of the compatibility of in-situ tower measurements

2 6.1 Independent low tank

The low tank was treated as an ambient sample, independent of the calibration. To evaluate the noise in the calibrated ambient samples that results from noise in the calibration, we calculated the standard deviation over the period September 1 – December 2 of the individual low-tank calibration cycles (6 min each), of the calibration cycles averaged over 1 day (81 min total), and of the calibration cycles averaged over 3 days (4.1 hours total). These results are a proxy for the noise in the calibrated ambient samples over those testing periods.

8

9 The low tank differences from known values, averaged over differing intervals, are shown in Fig. 9. The standard 10 deviation of individual low-tank calibration cycles (6 min each) over the period September 1 – December 2 is 0.62 11 . During this period, the calibration used 6 min/day measurements of the target tank. The standard deviation of 12 the low tank calibration cycles was similar to expectations based on the Allan standard deviation (Fig. 2). The low 13 tank was tested a total of 81 min (1.35 hour) per day. Thus calculating the standard deviation of the low tank values 14 averaged over each day is a measure of the noise due to the calibration scheme for hourly averages of sample data. 15 The standard deviation of daily averages for the low tank (81 min total) was 0.40 %. Based on this result, 16 differences in the hourly average between towers of less than 0.40 ‰ were likely not significant. For 3-day means 17 (a total of 4.1 hours), the standard deviation over the three-month period was 0.26 %. For the period after the 18 calibration tank sampling scheme was improved (primarily by sampling the target tank for 54 min/day instead of 6 19 min/day), December 3 – December 31, the standard deviation of the individual cycles reduced substantially, to 0.2520 ‰, and that of the 81-min (4.1 hour) mean of the cycles was 0.18 ‰ (0.11 ‰). Therefore, according to this metric, 21 after the improved calibration scheme was implemented, differences in the hourly average between towers of greater 22 than 0.18 ‰ were significant.

23 6.2 Round-robin testing

24 Post-deployment round-robin style tests were completed in the laboratory in March 2017 for three of the analyzers, 25 to assess the compatibility achievable via our calibration method. The analyzer deployed at the South tower was not 26 included in these tests, as it was still in the field. Two NOAA/INSTAAR tanks (JB03428: -46.82 $\% \delta^{13}$ CH₄, 27 1895.3 ppb CH₄ and 381.63 ppm CO₂; and JB03412: -45.29 $\% \delta^{13}$ CH₄, 2385.2 ppb CH₄ and 432.71 ppm CO₂) were 28 tested and treated as unknowns. The uncertainty for these NOAA tertiary standards was 0.1 ppm CO₂, including 29 scale transfer (Hall 2017; Zhao and Tans 2006), and 1 ppb CH₄ (GAW Report No. 185, 2009). The reproducibility 30 based on the calibration results was 0.06 ppm CO₂ and 0.4 ppb CH₄. The isotopic ratio was tied to the VPDB scale 31 but was not an official calibration (Michel and Vaughn, personal communication, 2015). The precision of the 32 determined values assigned to the tanks was 0.04‰ (https://instaar.colorado.edu/research/labs-groups/stable-33 isotope-laboratory/services-detail/). High, low, and target tanks were tested, with the calibration applied as in the field for ambient samples (as described in Section 5.4). The high mole fraction tank was tested for 20 min and the all ambient mole fraction tanks were tested for 70 min, with 8 min ignored after each gas transition. Four to six tests were completed for each analyzer. We used these tests as a means of evaluating the compatibility of the analyzers, in terms of both mole fractions and the isotopic ratio.

5

6 The results for the round-robin style laboratory testing are shown in Fig. 10. The mean of the errors (measured – 7 NOAA known value) for each analyzer/tank pair was -0.08 to 0.04 ppm CO₂ within the 0.1 ppm WMO 8 compatibility recommendation for global studies of CO2 (GAW Report No. 229, 2016). The standard error, 9 indicating an estimate of how far the sample mean is likely to be from the true mean, for the means of the CO₂ tests 10 were 0.03 - 0.10 ppm. The mean difference was -0.03 to 0.02 ppm CO₂ for the analyzers, averaged over the two 11 round-robin tanks (analogous to averaging over the entire range of CO₂ during the flask comparison, for example). 12 For CH₄, the means of the errors were 0.03 - 0.07 ppb CH₄, for the NOAA/INSTAAR tank measuring 2385.2 ppb, 13 and -0.83 to - 0.70 ppb CH₄ for the NOAA/INSTAAR tank measuring 1895.3 ppb CH₄. Therefore, there was a 14 slight error in the slope of the linear calibration, possibly attributable to tank assignment errors. However, the error 15 was well within the WMO recommendations for global studies of 2 ppb CH₄ (GAW Report No. 229, 2016), and the 16 range of NOAA/INSTAAR tanks encompassed the majority of the CH₄ mole fraction observed during the study. 17 We also note that the standard error for the means of the CH_4 tests were 0.07 - 0.12 ppb. Averaging over the two 18 round-robin tanks, the mean difference was -0.40 to -0.32 ppm CH₄ for the analyzers. For δ^{13} CH₄, the mean errors 19 for each analyzer/tank pair were -0.33 to 0.24 ‰ for these tanks within the range of ambient isotopic ratio and the 20 standard errors were 0.05 - 0.10 ‰. The mean errors were -0.14 to 0.03 ‰ for each analyzer.

21 6.3 Side-by-side testing

- 22 The precision and drift characteristics are not optimized for CO₂ for the G2132-i analyzers, compared to the G2301 23 and G2401 analyzers, which measure mole fractions and not isotopic ratios. Whereas the spectral line for CH₄ is the 24 same between the two types of analyzers (Rella et al., 2014), for CO₂, the absorbance of the spectral line used in the 25 G2132-i analyzers is a factor of 11 times less, meaning the precision is dramatically reduced. Although not central 26 to the primary results of this project, the performance of the analyzers in terms of CO_2 is important if the data are to 27 be used as part of the continental-scale CO₂ network. To test the performance of the G2132-i analyzers for 28 consideration of the data for this use, G2301 and G2132-i (Picarro, Inc.) analyzers were run side-by-side for one 29 month (June 2016) at the South tower. The sampling system for the G2132-i was as described in Section 5.2. A 30 separate 1/4" (0.64 cm) tube was used for the G2301 analyzer and an intercept calibration using the target tank was 31 applied daily. The sample air for the G2301 analyzer was not dried and the internal water vapor correction was 32 used.
- 33

This testing resulted in mean differences of 0.06 ± 0.41 ppm CO₂ and 0.9 ± 1.5 ppb CH₄, with the G2132-i analyzer measuring slightly lower for both species. Here the standard deviation was based on the 10-min average calibrated

- 1 values for the month for all times of the day. The standard error of the differences was $0.01 \text{ ppm } \text{CO}_2$ and 0.02 ppb
- 2 CH_4 . These results indicate that the performance of the G2132-i is similar for CO_2 and CH_4 mole fractions, at least
- 3 in terms of the long-term mean. In terms of utilizing the mole fraction data in atmospheric inversions, the multi-day
- 4 mean afternoon differences are most appropriate. The five-day mean afternoon difference for the month was
- 5 0.05 ± 0.08 ppm CO₂ and -0.7 ± 0.1 ppb CH₄. The G2132-i analyzers are thus appropriate for use in the atmospheric
- 6 inversions and in the global network where 0.1 ppm CO₂ and 2.0 ppb CH₄ have been identified as criteria. For these
- 7 results, recall that the target tank was tested for a total of 30 min in five days. To optimize results on a daily time
- 8 scale, sampling the target tank for 60 min per day would be preferable for improving CO₂ results. We also note that
- 9 round robin testing of these instruments requires 60 min sampling per tank.

10 6.4 Flask to in-situ comparison

11 In addition to the continuous G2132-i analyzers, the East and South towers were also equipped with NOAA flask 12 sampling systems (Turnbull et al. 2012). These flask measurements were used for independent validation and error 13 estimation of the continuous CO₂, CH₄ and δ^{13} CH₄ in-situ measurements. In addition, the flasks were measured for 14 a suite of species including N₂O, SF₆, CO, H₂ (Dlugokencky et al., 2017), halo- and hydro-carbons (Montzka et al., 15 1993) and stable isotopes of CH_4 (Vaughn et al., 2004). The flasks were filled over a 1-hour time period in the 16 afternoon (1400–1500 LST), thereby yielding a more representative measurement compared to most flask sampling 17 systems, which collect nearly instantaneous samples (e.g., ~10 sec). Samples were collected only when winds were 18 blowing steadily out of the west or north (\sim 45–225°) to ensure that the samples were sensitive to and representative 19 of the broader Marcellus shale gas production region that is the focus of this study. For the in-situ data, ten-minute 20 segments were reported. These were averaged over the hour for comparison with the flask measurements. For CH₄, 21 data points with high temporal variability (standard deviation of the 10-min means within the hour > 20 ppb) were 22 excluded, on the basis that the ambient variability was large, making comparisons difficult.

23

24 For January – December 2016, the mean flask to in-situ CH₄ difference at the East tower was -1.2 ± 2.2 ppb CH₄, 25 and at the South tower was -0.9 ± 1.4 ppb CH₄ (Fig. 11A). Here the standard deviation reported is that of the hourly 26 flask to in-situ differences. Thus, at the South tower, for example, on 67% of the sampled afternoons indicated 27 differences for CH_4 within 1.4 ppb of the mean of -0.9 ppb. The standard error was 0.24 ppb at the East tower and 28 0.14 ppb at the South tower. Thus, there is high confidence that the difference between the in-situ and flask 29 measurements at both towers is more compatible than the WMO recommendation. As for the side-by-side testing, 30 the G2132-i analyzers were slightly lower than the "known", in this case, the flask results. The difference, was 31 however, less than the target compatibility, and the flasks could in theory be biased.

32

Although CO₂ is not the focus of this paper, the differences were -0.21 ± 0.31 ppm for the East tower and 0.21

- ± 0.35 ppm for the South tower (Fig. 11B). The standard error was 0.03 ppm at the East tower and 0.04 ppm at the
- 35 South tower. The magnitude of CO₂ differences was somewhat larger in the growing season. The mean flask to in-

situ differences were thus larger than the WMO recommendation of 0.1 ppm, but at the extended compatibility goal
 of 0.2 ppm CO₂ (GAW Report No. 229, 2016).

3

For the isotopic ratio of methane, the mean flask to in-situ differences were $0.08 \pm 0.54\%$ and $0.02 \pm 0.38\%$ at the East and South towers, respectively (Fig. 11C). The standard error of the differences was 0.06% and 0.04% at the East and South towers, respectively. The range of δ^{13} CH₄ throughout the project (including day and night) was relatively small: one standard deviation (67%) of the data points are between 46.7 – 48.2 ‰, a range of 1.5 ‰. Errors for isotopic ratios outside the calibration range (further from the high and target calibration tanks) would likely be larger. For example, the mean error of the independent low tanks (averaging over all calibration cycles during a one month period) at the towers (Table 3) were 0.2 - 0.7 ‰.

12 7 Network comparisons

13 7.1 Study area

Four CRDS isotopic CH₄ analyzers (G2132-i, Picarro, Inc.) were deployed on commercial towers 46–61 m AGL in northeast Pennsylvania (Fig. 12). The South and North towers were located on the southern and northern edges of the unconventional gas well region, respectively, and were intended to measure background values depending on the wind direction. Measurements began in May 2015, but a complete set of field tanks necessary for calibration of δ^{13} CH₄ was not deployed until January 2016. The Central tower measured only mole fractions for the period June – December 2016. For inter-tower comparisons, we focused on the period January – May 2016 when all sites measured both CH₄ and δ^{13} CH₄

21 7.2 Inter-network differences in CH₄ and δ^{13} CH₄

22 A background value is required to calculate differences in CH₄ and δ^{13} CH₄. For this simple analysis, we chose a 23 single tower to represent the background for the entire period. The predominant wind direction for the Marcellus 24 region is from the west (Fig. 13). For westerly winds, the South tower is a reasonable choice for a background 25 tower. The South tower measured the lowest overall mean afternoon methane mole fraction (1960.2 ppb CH₄). The 26 mean afternoon methane mole fractions of the other towers, averaged only when data for the South tower exist, were 27 8.7, 7.0, and 2.9 ppb higher, at the North, Central, and East towers, respectively. For future analysis, a wind 28 direction-dependent background tower (South or North) could be considered, but the North tower did have the 29 largest mean enhancement in CH₄ mole fraction compared to the South tower. As noted by Barkley et al. (2017), 30 the area encompassing southwestern Pennsylvania and northeastern West Virginia contains large sources of CH₄, 31 with emissions from conventional gas, unconventional gas, and coal mines all having significant contributions to the 32 total. These large sources complicated the interpretation of the signals, as does changing wind direction. For this

overview analysis, we calculated differences above the South background tower to determine overall signal strength to compare with our target compatibility. We first examine the afternoon (defined here are 1700 – 2059 UTC), when the atmospheric is well mixed, allowing simpler interpretation of the measurements and more tractable modeling. We then consider non-afternoon hours, when the atmosphere is less mixed and signals are typically larger.

6 In the first set of plots, we focus on the majority of the afternoon data points by truncating the scale for the 7 probability distribution functions of methane mole fraction and isotopic ratio (Fig. 14A, B, D, E, G, and H). The 8 averaging interval of the individual data points was 10 min and the data were afternoon only (1700-2059 UTC, 9 1200–1559 LST) for the time period January – May 2016. The median differences for both isotopic ratio (-0.15 to 10 0.12 ‰) and methane mole fraction (less than 1 ppb) were less in magnitude than the compatibility of the analyzers. 11 This result is generally consistent with the results of Barkley et al. (2017), who found the emission rate of methane 12 due to natural gas extraction activities to be very low, 0.36 % of total production. The standard deviation of 10-min 13 segments of isotopic ratio differences was 0.8 ‰ at each of the towers. We note that the Allan standard deviation 14 for 10-min averaging times for ambient levels of methane was 0.4 $\% \delta^{13}$ CH₄. The standard deviation of the daily 15 afternoon averages (rather than 10-min averages) was 0.6 - 0.7 %. Thus the observed width of the distribution 16 appears to be persistent throughout the afternoon and not merely measurement noise. For isotopic ratio, 43 - 54 %, 17 depending on the tower, of the 10-min segments were greater than 0.6 ‰ in magnitude (3 times the target 18 compatibility) (Fig. 14A, D, and G) and are thus detectable by the analyzers. The standard deviations of the 19 methane mole fraction differences were 60.7, 30.0 and 33.8 ppb for the North, Central, and East towers, respectively 20 (Fig. 14 B, E, and H). 57 - 66 % of the data points indicated differences greater than 6 ppb CH₄ in magnitude (3 21 times the target compatibility) for the North, Central, and East towers, respectively (Fig. 14 B, E, and H) and are 22 thus detectable. The majority of afternoon data points indicated relatively few local sources of contamination.

23

There are however a few outliers during the time period with large values above the background tower during the afternoon hours (up to 1500 ppb enhancement at the North tower). The isotopic as a function of inverse methane mole fraction at each non-background tower are shown in Fig 13C, F, and I. While the range of measured isotopic ratios is large, the majority of the 10-min means lie close to the ambient values: the standard deviation of the 10min means of the measured isotopic ratios during the afternoon were 0.6 - 0.8 ‰.

29

During non-afternoon hours (0000–1659 and 2100–2359 UTC), the median isotopic ratio difference from the South tower were still indistinguishable from zero (Fig. 15A, D, and G). The median methane mole fraction enhancement was slightly higher than during the afternoons, at 3.5, 6.8, and 9.8 ppb for the North, Central, and East towers, respectively (Fig. 15B, E, and H). There were however more outliers, particularly at the Central tower (Fig. 15C, F, and I). Applying a best fit line to all of the data shown in Fig 14F gave a poor correlation coefficient (r²=0.22) because there were many data points with no local sources.

- 36
- 37 7.3 Keeling plots

Keeling plots (Keeling 1961; Röckmann et al., 2016) are used to infer the isotopic ratio of the methane source as the intercept of the best fit line of the isotopic ratio as a function of the inverse methane mole fraction. We used this approach to estimate the source isotopic ratio of the eight largest peaks observed during non-afternoon hours at the Central tower. The time series of CH₄ encompassing the peak observed on DOY 55 is shown in Fig. 16, as an example. The time during which the tower was in the plume was clear (lasting about 1.5 hours) and only those points were included in the calculation of the linear fit.

8

9 The Keeling plots for each of the eight largest peaks in the non-afternoon methane data are shown in Fig. 17. The 10 intercepts of the best fit lines for the peaks indicate that the sources contributing to the peaks have a mean isotopic 11 ratio of -31.2 ± 1.9 ‰. The correlation coefficients were high ($r^2=0.92 - 1.0$) except for one peak, which was 12 excluded from the statistics. Propagating a potential error (attributable of analyzer uncertainty) of 0.2 ‰ at the 13 heavy end of the Keeling plots and -0.2 ‰ at the light end, and vice versa, the potential range of the mean is from – 14 32.0 to -30.4 ‰.

15

16 Compared to mobile measurements near the ground, for example, the footprints of towers are large, which is ideal 17 for determining regional emissions. But the emissions sources with specific isotopic signatures are diluted by 18 mixing, making the enhancements above background small, particularly for this region/time period with small 19 leakage rates. For these eight non-afternoon peaks at the Central tower, the enhancements over background were 20 334.1 - 2007.8 ppb CH₄ and the differences of isotopic ratio were -2.5 to -8.7 ‰.

21 8 Discussion

22 In this paper, we present the methods used to calibrate a network of four CRDS methane isotopic ratio analyzers 23 (Picarro G-2132i). Evaluation of the calibration results using an independent tank, round-robin style testing and 24 flask comparisons showed that the analyzers are compatible within 0.2 ‰. The calibration required consideration of 25 1) the isotopic ratio linear calibration, 2) the mole fraction dependence of the isotopic ratio calibration (using high 26 and ambient mole fraction tanks), 3) the correction due to ethane cross interference (using one tank without ethane 27 and one tank with ambient ethane), and 4) drift in the CO_2 and CH_4 mole fractions (using at least one tank near 28 ambient isotopic ratio and mole fraction). The isotopic ratios and CH_4 mole fractions of these tanks as used in the 29 present deployment are graphically represented in Fig. 18A. Prior to implementation of the improved field tank 30 testing strategy, the high and low tanks were tested for 26 and 52 min/day (excluding transition time between gases), 31 as listed in Table 4. The testing times throughout each day for the high and low standards are sufficient for Allan 32 deviation < 0.1‰, but neither of those tanks were at ambient ranges of δ^{13} CH₄. If the calibrations and analyzer 33 response were both linear, we would expect negligible errors in the target tank if kept independent, but we found a 34 bias between -0.3 and -0.8 %, which is very large compared to the ambient differences observed (one standard 35 deviation of the tower measurements at all times of day were between -48.2 and -46.7 ‰). Thus, we instead chose to minimize mean error at ambient values (target tank) rather than at the isotopic ratios of the low tanks (-23.9 ‰).
This procedure added noise to the ambient data because the daily sampling time for the target tank was only 6 min/day. On 3 December 2016, we implemented an improved tank testing strategy, primarily by increasing the testing time for the target tank to 54 min/day.

5

6 Our recommendation for future similar studies is to choose both target and low tanks closer to the expected range of 7 isotopic ratios, in addition to being near ambient CH₄ mole fractions. For example, suggested values for the low and 8 target tanks are 2.1 ppm CH₄ at -46.5 ‰ and 1.9 ppm CH₄ at -47.5 ‰ (Table 4 and Fig. 18B). The testing time 9 required is dependent upon the compatibility goals. After implementing our improved tank testing time strategy, we 10 tested each target and low tank for about an hour per day, to achieve Allan deviations of 0.2 %. Source attribution 11 using mobile measurements, rather than tower measurements, for example, is less demanding in terms of 12 compatibility needed, due to the relatively large ambient signals typically encountered. The estimated testing time required to achieve Allan deviations less than 0.4 ‰, for example, can be achieved in 8 min. In general, it is 13 14 desirable to distribute the tank testing time throughout the time period, in our case, one day. In this case, persistent 15 changes in analyzer response over the day, if any, would be averaged over rather than an extreme value used in the 16 calibration. This procedure also avoids not sampling the ambient air for extended periods. We did not find any 17 evidence of variability in the calibrations on scales less than one day, compared to the precision possible given our 18 tank testing times, but this possibility could be further explored by testing the field tanks for longer periods of time. 19

20 The high tanks used in this network contained methane with about -38.3 % CH₄. This specific isotopic ratio is 21 available commercially, and depending on the compatibility goals of the project, may not require laboratory 22 calibration of the tank. For our case, however, it may have been beneficial to utilize isotopic ratios closer to the 23 observed range, perhaps -44 ‰ (Table 4). Another possibility is to add an additional high tank (Fig. 18B) in the 24 range of -54.5 % to -52 % (with -52 % more closely bracketed the observed isotopic ratios in the present study). 25 In this case, laboratory linear calibration of the analyzers is not necessarily required. Both the slope and intercept of 26 the linear calibration can be adjusted in field, rather than just the intercept, which may improve the calculated 27 accuracy and precision. However, the laboratory calibration in the present study utilized four different isotopic 28 ratios, rather than two, and it is unknown which is more important - improving linear calibration frequency or 29 avoiding over-constraining the calibration.

30

In this paper, we calibrated the total CH₄ and the isotopic ratio of methane. An alternative calibration approach is to separately calibrate the individual isotopologues (in this case, ¹³CH₄ and ¹²CH₄ dry mole fractions), as has been applied to Fourier Transform infrared and isotope ratio infrared spectrometers measuring δ^{13} C and δ^{18} O of CO₂ in air (Griffith et al., 2012; Wen et al., 2013; Flores et al., 2017). This approach has the advantage of simple calibration equations, but has the disadvantage that the quantities of interest (e.g., total mole fraction and isotopic ratio) are calculated rather than directly calibration. Like the approach applied in this paper, it also requires at least two standard tanks, and could utilize an independent tank for testing. Rella et al. (2015) list further practical reasons 1 to calibrate δ^{13} CH₄, including the lack of primary standards for 13 CH₄. However, a comparison of performance using 2 each of these techniques on the same dataset would be beneficial.

3

4 The signals observed in the study region were generally small, but the isotopic ratio differences were larger than 5 would be expected based on the methane mole fraction enhancements from local sources. For afternoon hours at the 6 Central tower, for example, 43 % of the differences in δ^{13} CH₄ were detectable above background with magnitudes > 7 0.6 %, 3 times the analyzer compatibility. For a thermogenic source with isotopic ratio of -35 %, a background 8 isotopic ratio of -47 ‰, and assuming a measured CH₄ mole fraction of 2000 ppb, a measured isotopic ratio 9 difference of -0.6 % corresponds to a 100 ppb peak in CH₄ above background, following Eq. (1). Enhancements in 10 CH_4 of 100 ppb were rarely encountered, however (Fig. 14B, E, and H). Using Eq. (1) to predict differences of 11 isotopic ratio based on the observed methane mole fraction enhancements corresponded to only 3 % of the isotopic 12 ratio differences expected to be > 0.6 % in magnitude. Thus during the afternoon hours, most of the deviations 13 from background were not likely directly from local sources. These larger than expected differences in isotopic 14 ratio are not primarily attributable to analyzer noise. The Allan Deviation (Fig. 2) is 0.4 % for 10-min means at 15 ambient mole fractions of 2 ppm CH₄. We also note that we focused on the period January – May 2016 in this work. 16 Larger differences were observed in the latter half of 2016.

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18 During the morning hours, however, several peaks resulting from local sources were observed. The mean source 19 isotopic signal indicated by Keeling plot analysis of the eight largest peaks at the Central tower was -31.2 ± 1.9 %, 20 fairly heavy even for oil/natural gas sources. In general, the isotopic signature for natural gas sources varies from 21 region to region, and even within one region. The mean isotopic ratio of methane in gas wells in the northeastern 22 Pennsylvania section of the Marcellus region has been shown to vary by depth, from -43.42 ‰ with a standard 23 deviation of 6.84 ‰ for depths of 0 to 305 m, to -32.46 ‰ with a standard deviation of 3.84 ‰ for depths greater 24 than 1524 m (Baldassare et al., 2014). Similarly, Molofsky et al. (2011) found that the isotopic signatures of gases 25 from the deeper layers of the Marcellus Shale in Susquehanna County, Pennsylvania, to be heavier than the 26 shallower Middle and Upper Devonian deposits, with values for the deep layers ranging from -30 to -21 ‰. Thus, 27 the source signature determined here is consistent with a natural gas source originating from deep wells in the 28 Marcellus region. The peaks occurred during the morning hours, when the boundary layer is typically stable, 29 making modeling more difficult, and the winds prior to the peaks were not from a consistent direction. 30 Determination of the location of the specific emitter(s) contributing to these peaks is thus beyond the scope of this 31 paper. Based on the lack of consistent wind direction, it seems likely that more than one location (with potentially 32 different source signatures) contributed to these peaks. We note that the Keeling plot approach to determine source 33 isotopic signatures far from the point of emission will be difficult to apply in regions without sources that are 34 significantly depleted or enriched in ¹³CH₄ compared to ambient.

For determination of the source signature for a specific known location, the tower-based approach is not ideal.Instead the strength of the tower-based approach lies in covering larger areas and many potential source locations,

1 and for longer periods of time than is feasible by other approaches. The instrumental performance demonstrated 2 here could be used to disaggregate methane sources in areas of stronger enhancements and differing source isotopic 3 signatures. Networks of high-temporal-resolution methane isotopic ratio data have the potential to constrain 4 regional methane budgets when used within a modeling framework. 5 6 7 **Data Availability** 8 Miles, N.L., D.K. Martins, S.J. Richardson, T. Lauvaux, K.J. Davis, B.J. Haupt, and C. Rella, 2017. In-situ tower 9 atmospheric methane mole fraction and isotopic ratio of methane data, Marcellus Shale Gas Region, Pennsylvania, 10 USA, 2015-2016. Data set. Available on-line [http://datacommons.psu.edu] from The Pennsylvania State University 11 Data Commons, University Park, Pennsylvania, USA. http://dx.doi.org/10.18113/D3SG6N. 12 13 **Competing Interests** 14 TL, SJR, NLM, and KJD are co-owners of a related company, Carbon Now Cast, LLC. 15 16 17 Acknowledgments. The authors thank B. Vaughn and S. Englund Michel (Institute of Arctic and Alpine Research, 18 University of Colorado) for providing analysis of methane isotopic ratios of the flask data and for advice regarding 19 gas handling techniques involving isotopic ratios. The authors also acknowledge R.P. Barkley (Tunkhannock Area 20 Middle School) for his contributions to maintaining instrumentation at the tower sites. This work was funded by the 21 Department of Energy National Energy Technology Laboratory (DE-FOA-0000894). 22

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Tables

Table 1. Field tanks used at the tower locations. The high and target tanks were used for the field calibration of δ^{13} CH₄. Only the target tank is used for field adjustment of the CH₄ and CO₂ mole fraction calibration. The CH₄ and CO₂ mole fractions for the high and low tanks are less certain than that of the target tanks.

*Determined via laboratory measurements. **NOAA/INSTAAR calibration (WMO X2004A scale for CH₄ and WMO X2007 for CO₂). *** Field calibration – values not used.

Tank number	Deployment location	Measured isotopic ratio δ ¹³ CH ₄ (‰)	CH4 mole fraction (ppb)	CO ₂ mole fraction (ppb)	Used for field calibration of δ ¹³ CH4	Independent test of δ ¹³ CH ₄ calibration	Used for field adjustment of CH ₄ and CO ₂ mole fraction calibration (intercept only)	Used for ethane correction
CA06418	North-High	-38.31*	9701*	397.75***	~	,		✓
CA05551	North-Low	-23.67*	1926.8*	402.70***		✓		
CB10825	North- Target	-47.26**	1867.59**	399.71**	~		✓	~
CA05419	Central- High	-38.48*	10534*	399.66***	\checkmark			\checkmark
CA06438	Central-Low	-23.80*	2064.6*	397.82***		\checkmark		
CB10734	Central- Target	-47.25**	1878.53**	397.09**	✓		~	✓
CA05330	South-High	-38.68*	10152*	403.10***	\checkmark			\checkmark
CC114999	South-Low	-23.72*	1999.2*	402.58***		\checkmark		
CB10727	South- Target	-47.24**	1868.33**	399.68**	\checkmark		\checkmark	~
CA06410	East-High	-38.52*	10414*	407.45***	\checkmark			✓
CA06357	East-Low	-24.02*	2079.7*	368.47***		√		
CB10718	East-Target	-47.26**	1867.94**	399.67**	\checkmark		\checkmark	~

Table 2. Maximum error estimate attributable to cross-interference due to direct absorption on δ^{13} CH₄. These estimates were based on typical values for this tower-based application and estimated effects on CRDS measurements (Rella et al., 2015), and assumed 2 ppm ambient CH₄ mole fraction. For water vapor and carbon dioxide, the interferences are independent of CH₄ mole fraction for 1 – 15 ppm. For the other species listed, the interferences are inversely proportional to CH₄ mole fraction. Typical maximum values determined by flask^f (level at which 99 % of (afternoon) flask measurements at the South and East towers are below), by in-situ measurements at Marcellus towersⁱ, or by typical values^t (Warneck and Williams, 2012). ^aNo known ambient estimates (Barnes, 2015) / odor threshold (Devos et al., 1990).

Gas Species	Typical maximum value or range	Estimated maximum error
Carbon monoxide	Range ^f : 107.5-200.7 ppb	0.01‰
Water vapor, dried sample	Range ⁱ : 0.02 – 0.06%	0.02‰
Water vapor, ambient moisture	Range: 0 – 2.5%	±1‰ (Rella et al., 2015)
Carbon dioxide	Range ⁱ : 375 – 475 ppm	0.03‰
Propane	Max ^f 3.6 ppb	0.01‰
Butane (i-Butane + n-Butane)	Max ^f 1788 ppt	0.01‰
Ammonia	Typical ^t 90 ppt	0.01‰
Hydrogen sulfide	Typical ^t 30 ppt	0.01‰
Methyl mercaptan	Odor threshold ^a : 1 ppb	0.01‰
Ethylene	13.0 ^f ppt	0.01‰
Ethane	Max ^f 8.0 ppb (typical background ^t : 1.3 ppb)	0.23‰ (0.04‰ typical)

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Table 3. Results for the four Marcellus towers using two possible calibration schemes. Tank errors are shown for the using the high and low tank in the calibration (SCHEME A) and using the high and target tank in the calibration (SCHEME B). The third set of results are for SCHEME B, but following the change in field tank testing times on 3 Dec 2016. Results fare from October 2016 for the South, East and North towers, but are from May 2016 for the Central tower, as the analyzer was at the manufacturer for repairs during October 2016. Note that the daily means of the field tanks are used in the calibrations.

	Tower	High tank error (‰)	Low tank error (‰)	Target tank error (‰)	
		mean \pm standard	mean \pm standard	mean \pm standard deviation	
		deviation for one	deviation for one	for one month (standard	
		month (standard	month (standard	error)	
		error)	error)		
SCHEME A	South	Used in cal	Used in cal	-0.3±0.4 (0.1)	
SCHEME A	East	Used in cal	Used in cal	-0.8±0.5 (0.1)	
SCHEME A	Central	Used in cal	Used in cal	-0.5±0.3 (0.1)	
SCHEME A	North	Used in cal	Used in cal	-0.4±0.7 (0.1)	
SCHEME B	South	Used in cal	0.2±0.7 (0.0)	Used in cal	
SCHEME B	East	Used in cal	0.7±0.6 (0.0)	Used in cal	
SCHEME B	Central	Used in cal	0.4±0.5 (0.0)	Used in cal	
SCHEME B	North	Used in cal	0.3±1.3 (0.1)	Used in cal	
(following change in					
field tank testing times					
on 3 December 2016)					
SCHEME B	South	Used in cal	0.3±0.3 (0.0)	Used in cal	
SCHEME B	East	Used in cal	0.6±0.5 (0.0)	Used in cal	
SCHEME B	Central	Used in cal	0.4±0.3 (0.0)	Used in cal	
SCHEME B	North	Used in cal	$-0.4\pm0.9(0.0)$	Used in cal	

Table 4. Possible field tanks and sampling strategies, including those employed in the present study. The "Improved strategy" column suggests a possible strategy in which three field tanks and one independent tank are employed, and thus laboratory calibration is not required. Estimated tank testing times (excluding transition times) are listed for various compatibility requirements.

	Present study prior	Present study 3	Improved strategy
	to 3 December 2016	December 2016	
		and thereafter	
Laboratory	Yes, for linear	Yes, for linear	No
calibration needed?	calibration	calibration	
High CH ₄ mole	HIGH (10 ppm,	HIGH (10 ppm,	HIGH (10 ppm, -38.3‰ to -44‰, 8 min/day for 0.1‰
fraction tank(s)	-38.3‰, 26 min/day)	-38.3‰, 10	Allan deviations, 1 for 0.2 ‰, 1 for 0.4 ‰)
		min/day)	
	-	-	HIGH (10 ppm, -54.5‰ to -52‰, 8 min/day for 0.1‰
			Allan deviations, 1 for 0.2 ‰, 1 for 0.4 ‰)
Low CH ₄ mole	LOW (2 ppm,	LOW (2 ppm, -	LOW (2.1 ppm, -46.5‰ (ambient), 120 min/day for 0.1‰
fraction tanks	-23.9‰, 81 min/day)	23.9‰, 54	Allan deviations, 60 for 0.2 ‰, 8 for 0.4 ‰)
	independent	min/day)	
		independent	
	TARGET (2 ppm,	TARGET (2	TARGET (1.9 ppm, -47.5‰ (ambient), 120 min/day for
	-47.2‰, 6 min/day)	ppm, –47.2‰,	0.1‰ Allan deviations, 60 for 0.2 ‰, 8 for 0.4 ‰)
		54 min/day)	independent
Notes		Reduced noise in	Does not necessarily require laboratory calibration of
		calibration due to	analyzers. Range of ideal isotopic ratios for the high tanks
		increased target	is given. Utilizing the isotopic ratios of commercially
		tank sampling	available bottles for spiking (i.e., -38.3‰ and -54.5‰) may
		time	avoid the need for laboratory calibration of these tanks.
			Using low/target tanks near ambient isotopic ratio range (but
			not exactly the same isotopic ratio, and preferably not
			exactly the same mole fraction) is more accurate reflection
			of compatibility and range of the isotopic ratio of the high
			tanks better encompasses expected values. For applications
			with reduced compatibility requirements (e.g., 0.4 %),
			utilizing low/target tanks at commercially available -38.3‰
			and -54.5‰ may be sufficient. It is advantageous to
			distribute field tank testing throughout the day, to avoid not
			sampling ambient air for long periods and to measure
			potential changes in analyzer response.





Figure 1. Isotopic ratio difference from background ($\Delta\delta$) resulting of a mixture of background and source signatures, as a function of source isotopic ratio (δ_{source}) and CH₄ mole fraction enhancement above background (Δ CH₄). Here the source end members are -60 ‰ and -35 ‰. Background CH₄ mole fraction was assumed to be 2000 ppb and background isotopic ratio -47.5 ‰ (vertical solid line). Dashed lines indicate -0.3 ‰ and 0.3 ‰ difference from background.





Figure 2. Allan standard deviation for (A) δ^{13} CH₄, (B) CH₄, and (C) CO₂ for a high CH₄ mole fraction tank (9.7 ppm CH₄, ~400 ppm CO₂, -38.3 ‰ δ^{13} CH₄) (orange) and a low (1.9 ppm CH₄, ~400 ppm CO₂, -23.7 ‰ δ^{13} CH₄) tank (blue). The x-axis is truncated to focus on minimum averaging times required to achieve the desired compatibility goals.





Figure 3. Flow diagram of the experimental setup used for the laboratory calibration of the analyzers and the field tanks (working standars). At standard pressure and temperature, the gas volume of the zero air and working standard tanks

6 was 4021 L and that of the Isometric Instruments bottles was 28 L.



11 Figure 4. Standard deviation of the CH₄ isotopic ratio during the test results shown in Fig. 5.



3 4 5 6 7 8 9 Figure 5. Measured isotopic ratio error as a function of known isotopic ratio for each of the four analyzers (A - D), prior to calibration. The colors indicate the ¹²CH₄ mole fraction, as shown in the legend. The serial numbers (FCDS2046, FCDS2047, FCDS2048, and FCDS2049) of the analyzers are indicated as well. These analyzers were deployed at the South, Central, North and East towers, respectively. Interpolating from the Allan standard deviation results (Fig. 2), the estimated precision is 0.40 ‰ for the 1.80-1.82 ppm CH₄ tests, 0.34 ‰ for 3.28-3.32 ppm CH₄ tests, 0.24 ‰ for 6.10-6.16 ppm CH₄ tests, and 0.20‰ for 7.26–7.32 ppm CH₄ tests.



Figure 6. Flow diagram of the field calibration system. At standard pressure and temperature, the gas volume of the

field tanks was 4021 L.





Figure 7. Results following isotopic ratio laboratory calibration only (black) and following calibration (blue) for the South tower for September - December 2016 for the "high" CH_4 mole fraction tank (A), "low" CH_4 mole fraction tank (B), and target tank (C). The target tank was used in the isotopic ratio calibration, whereas the low tank was independent. An improved calibration tank sampling strategy was implemented on 3 December 2016 (indicated by vertical dashed lines). The Allan deviation for time period used for each calibration cycle was, for the period prior to the improved tank sampling strategy, 0.2 ‰ for the high tank, and 0.5 ‰ for the low and target tanks. Following the high tank, and 0.3 ‰ for the low and target tanks.





4 5 6 7 Figure 8. Effect of each of the calibration coefficient terms for the South tower for September - December 2016 for the optimized calibration scheme. The terms c_0 (A) and χ (B) in Eq. (3) are time-dependent drift terms. Note the differing scales. An improved calibration tank sampling strategy was implemented on 3 December 2016 (indicated by vertical dashed lines).

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12 13 14 Figure 9. Low tank methane isotopic ratio differences from known value, for the individual calibration cycles (blue), and for 1-day (red) and 3-day (black) means of the calibration cycles, for the South tower for September - December 2016. An improved calibration tank sampling strategy was implemented on 3 December 2016 (indicated by the vertical dashed line). The low tank is independent of the isotopic ratio calibration.



Figure 10. Results from round-robin style testing using two NOAA/INSTAAR tanks (JB03428: -46.82 ‰ δ^{13} CH₄, 1895.3 ppb CH₄ and 381.63 ppm CO₂; and JB03412: -45.29 ‰ δ^{13} CH₄, 2385.2 ppb CH₄ and 432.71 ppm CO₂) for CO₂ (top row), CH₄ (middle row), and δ^{13} CH₄ (bottom row), for the analyzer deployed at the North tower (serial number FCDS2048; left column), at the Central tower (serial number FCDS2047; middle column), and at the East Tower (serial number FCDS2049; right column). These tests were completed in the laboratory, post deployment (March 2017). The analyzer deployed at the South tower (serial number FCDS2046) was not included in these tests. Open circles are individual tests and filled circles are the means of the individual tests for each analyzer/constituent. The mean error for each analyzer/tank/constituent is indicated in the plots.



Figure 11. Afternoon in-situ to flask differences for January – December 2016 for the East (blue) and the South towers (orange) for A) CO₂, B) CH₄, and C) δ^{13} CH₄. For CH₄, data points with high temporal variability (standard deviation of raw ~2sec data within the 10-min segments > 20 ppb) are indicated by '+' symbols and have been excluded. The standard deviation of the in-situ to flask differences are shown in parentheses on each plot. The standard errors, indicating an estimate of how far the sample mean is likely to be from the true mean, is 0.24 ppb CH₄, 0.03 ppm CO₂ and 0.06 ‰ at the East tower and 0.14 ppb CH₄, 0.04 ppm CO₂ and 0.04 ‰ at the South tower.



Figure 12. Map of Pennsylvania with permitted unconventional natural gas wells (magenta dots) and network of towers with methane and stable isotope analyzers (Picarro G2132-i). The East and South towers were also equipped with NOAA flask sampling systems. The Binghamton Airport is also indicated.



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5 Figure 13. Wind rose for surface station at Binghamton, NY airport for the period April 2015 – April 2016 (using the 6 mean of the afternoon hours for each day). The magnitude of wedges indicates relative frequency for each wind direction 7 and the wind speeds are indicated by color. These afternoon means were based on hourly reported measurements. For 8 the hourly measurements, calm winds (< 1.6 m s⁻¹) were not categorized by direction and thus were not included in the 9 afternoon mean. For the hourly measurements, calm winds (< 1.6 m s⁻¹) were reported as zero and were included in the 10 afternoon mean.



6 7 8 9 10 11 12 13 14 15 16 Figure 14. Probability distribution function of measured isotopic ratio differences from the background South tower (Δ δ^{13} CH₄) for the A) North, D) Central, and G) East towers for afternoon hours (1700–2059 UTC, 1200–1559 LST). The averaging interval of the individual data points for all plots is 10 min and the time period is January - May 2016. The bin size for A), D) and G) is 0.2 ‰. The median and standard deviation of the differences are indicated on the plots. Probability distribution function of measured methane mole fraction enhancements (Δ CH₄) for the B) North, E) Central, and H) East towers. Note that the scale for B), E, and H) has been truncated to focus on majority of the data points. The bin size is 10 ppb CH₄. Keeling plots for the C) North, F) Central, and I) East towers. The black box in each plot indicates the approximate scale of the corresponding isotopic ratio difference and methane mole fraction enhancement plots. The median and standard deviation of the isotopic ratios at each tower are indicated on the plots. Note that the Allan deviation for 10-min means at ambient mole fractions was 0.4 ‰ and this decreases with increasing mole fraction.



Figure 15. Probability distribution function of measured isotopic ratio differences from the background South tower ($\Delta \delta^{13}$ CH₄) for the A) North, D) Central, and G) East towers for all times of data excluding the afternoon hours shown in Fig. 14. The averaging interval of the individual data points for all plots is 10 min and the time period is January – May 2016. The bin size for A), D) and G) is 0.2 ‰. The median and standard deviation of the differences are indicated on the plots. Probability distribution function of methane mole fraction enhancements (Δ CH₄) for the B) North, E) Central, and H) East towers. Note that the scale for B), E, and H) has been truncated to focus on majority of the data points. The bin size is 10 ppb CH₄. Keeling plots for the C) North, F) Central, and I) East towers. The black box in each plot indicates the approximate scale of the corresponding isotopic ratio difference and methane mole fraction enhancement plots. Note that the Allan deviation of the isotopic ratios at each tower are indicated on the plots. Note that the Allan deviation for 10-min means at ambient mole fractions was 0.4 ‰ and this decreases with increasing mole fraction.





9 10 11 12 13 Figure 16. Time series of CH₄ encompassing one of the eight peaks in CH₄ at the Central tower (DOY 55) for which the Keeling plot approach was applied. The averaging interval of the individual points was 10 min, and periods during which field tanks were sampled were excluded from the plot. The linear fit was calculated using the points clearly within the plume (black dots).

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Figure 17. Keeling plots for the Central tower for the eight largest peaks in the non-afternoon methane time series. Black lines indicate the best-fit lines. Correlation coefficients (r²), day of year (DOY) and y-intercepts are indicated in the plots. FIX subscripts.





Figure 18. Graphical representation of the field tanks used in the present study (A), and for an improved strategy (as in Table 4) (B). Orange 'H' symbols indicate high mole fraction tanks, blue 'L' symbols indicate low mole fraction tanks, and red 'T' symbols indicate target tanks. Lines in (B) indicate range of isotopic values desirable for the high tanks. Ideally, calibration tanks are near the range of ambient values to be measured, but in this case, specific values are more easily obtained (-54.5 ‰, -38.3 ‰, -23.9 ‰, from Isometric Instruments, Inc.)