Spatial patterns and source attribution of urban methane in the Los Angeles Basin

Francesca M. Hopkins¹,², Eric A. Kort³, Susan E. Bush⁴, James R. Ehleringer⁴,⁵, Chun-Ta Lai⁶, Donald R. Blake⁷, and James T. Randerson¹

¹Department of Earth System Science, University of California, Irvine, California, USA, ²Now at Jet Propulsion Laboratory, California Institute of Technology, Pasadena, CA, USA, ³Department of Atmospheric, Ocean, and Space Sciences, University of Michigan, Ann Arbor, Michigan, USA, ⁴Department of Biology, University of Utah, Salt Lake City, Utah, USA, ⁵Global Change and Sustainability Center, University of Utah, Salt Lake City, Utah, USA, ⁶Department of Biology, San Diego State University, San Diego, California, USA, ⁷Department of Chemistry, University of California, Irvine, California, USA

Abstract

Urban areas are increasingly recognized as a globally important source of methane to the atmosphere; however, the location of methane sources and relative contributions of source sectors are not well known. Recent atmospheric measurements in Los Angeles, California, USA, show that more than a third of the city’s methane emissions are unaccounted for in inventories and suggest that fugitive fossil emissions are the unknown source. We made on-road measurements to quantify fine-scale structure of methane and a suite of complementary trace gases across the Los Angeles Basin in June 2013. Enhanced methane levels were observed across the basin but were unevenly distributed in space. We identified 213 methane hot spots from unknown emission sources. We made direct measurements of ethane to methane (C₂H₆/CH₄) ratios of known methane emission sources in the region, including cattle, geologic seeps, landfills, and compressed natural gas fueling stations, and used these ratios to determine the contribution of biogenic and fossil methane sources to unknown hot spots and to local urban background air. We found that 75% of hot spots were of fossil origin, 20% were biogenic, and 5% of indeterminate source. In regionally integrated air, we observed a wider range of C₂H₆/CH₄ values than observed previously. Fossil fuel sources accounted for 58–65% of methane emissions, with the range depending on the assumed C₂H₆/CH₄ ratio of source end-members and model structure. These surveys demonstrated the prevalence of fugitive methane emissions across the Los Angeles urban landscape and suggested that un inventoried methane sources were widely distributed and primarily of fossil origin.

1. Introduction

Methane (CH₄) is an important atmospheric pollutant; the second largest contributor to global warming and a key constituent regulating CO and O₃ [Cicerone and Oremland, 1988]. Most sources of CH₄ to the atmosphere have been identified; however, their relative importance to the global budget is uncertain [Kirschke et al., 2013]. CH₄ source budgets are even more uncertain at continental and regional scales [e.g., Kort et al., 2008; Miller et al., 2013]. The majority (50–65%) of CH₄ emissions globally come from anthropogenic sources, with a flux of approximately 330 Tg CH₄ per year [Kirschke et al., 2013]. Reduction of CH₄ emissions has been suggested to be an effective short-term strategy to reduce global warming because of CH₄’s high radiative forcing relative to CO₂, around 28 times on a mass basis over a 100 year time horizon [Shindell et al., 2012; Myhre et al., 2013]. However, mitigation of anthropogenic CH₄ emissions requires an accurate CH₄ budget, including knowledge of location and sectoral contributions of different CH₄ emitters, particularly at scales where mitigation policies may be enacted [Hsu et al., 2010; Jeong et al., 2013].

Observations of elevated CH₄ levels in cities demonstrate that significant emissions of anthropogenic CH₄ are derived from urban areas [Blake et al., 1984; Wunch et al., 2009]. According to inventory estimates, 35% of the anthropogenic CH₄ in North America is emitted from urban regions [Marcotullio et al., 2013]. However, recent atmospheric studies at the state and city levels in California suggest a 30–80% underestimation of CH₄ emissions in the state greenhouse gas inventory, using stationary and airborne trace gas measurements [Wunch et al., 2009; Hsu et al., 2010; Wennberg et al., 2012; Jeong et al., 2013; Peischl et al., 2013; Wong et al., 2015]. Uncounted fugitive emissions, such as leaks from natural gas pipelines, are hypothesized to account for this mismatch between bottom-up inventories and top-down measurements [Brandt et al., 2014]. On-road surveys in major cities such as Boston and Washington, DC have revealed large fugitive leaks from natural gas distribution...
pipelines [Phillips et al., 2013; Jackson et al., 2014]. In addition to extensive natural gas pipeline networks, cities have a variety of other CH4 sources, including landfills, water treatment plants, natural gas vehicles, and infrastructure, and in the case of Los Angeles, fossil fuel extraction and refining and dairy agriculture. Fugitive emissions may also originate from these sectors. The heterogeneous mixture of source sectors in the urban environment complicates stationary and aircraft measurements of trace gases that cannot resolve fine-scale structure at the source level. Extensive road networks in cities enable vehicle coverage over large areas, providing a method for measuring the spatial distribution of CH4 emissions. On-road sampling of surface trace gas enhancement can locate CH4 emission hot spots and attribute CH4 enhancements to source sectors and aid in interpretation of stationary or remotely sensed measurements [e.g., Petron et al., 2012; Leifer et al., 2013].

In the U.S., the two largest sources of urban CH4 emissions are waste disposal and natural gas systems [U.S. Environmental Protection Agency (US EPA), 2014]. These two sources represent the two primary pathways by which CH4 is produced—biogenic and thermogenic. Waste disposal, in landfills and wastewater treatment plants, produces biogenic CH4 as a result of microbial decomposition of organic matter under anaerobic conditions. Biogenic CH4 is also produced in the gut of livestock and from manure. In contrast, thermogenic CH4 originates from the geologic processes that create all fossil fuels and is present in fossil fuel deposits including coal beds, oil fields, and geologic seeps [Etiope and Ciccioli, 2009]. Thermogenic CH4 is also emitted through intentional venting and fugitive leaks in the extraction, storage, refining, transport, and use of natural gas. Incomplete combustion of fuels represents a third pathway for CH4 production; pyrogenic sources are also a minor component (<2%) of CH4 emissions [US EPA, 2014]. Both thermogenic and pyrogenic sources of CH4 also emit more complex hydrocarbons, including ethane (C2H6), whereas biogenic sources do not [Rudolph, 1995; Kirchstetter et al., 1996; Etiope and Ciccioli, 2009]. Hence, elevated CH4 accompanied by elevated C2H6 values can be used as a tracer of fossil fuel sources of CH4 [e.g., Aydin et al., 2011].

In the Los Angeles Basin, evidence from stable isotopes of CH4 and measurements of higher hydrocarbons (e.g., C2H6, propane, and butane) suggest that fossil emissions are the predominant source of CH4 [Townsend-Small et al., 2012; Wennberg et al., 2012; Peischl et al., 2013]. Specifically, leakage from natural gas infrastructure [Wennberg et al., 2012] and from local fossil CH4 sources [Peischl et al., 2013] are thought to be the most important contributors. However, the complex geologic setting and intense human impact within the basin complicate CH4 source attribution in the Los Angeles area. The Los Angeles region is known for naturally occurring geologic seeps, such as the La Brea Tar Pits, as well as extensive oil drilling taking place during the last century and continuing to the present day [Bilodeau et al., 2007]. In addition, Los Angeles is a major industrial and shipping center, with more than 10 oil refineries and storage facilities. The number of producing wells in the basin has decreased since the 1960s; however, the number has increased anew over the past decade with application of enhanced oil recovery techniques such as hydraulic fracturing [Cardno ENTRIX, 2012; Gautier et al., 2012; Tennyson, 2005]. Industrial centers, such as the Port of Los Angeles, the Harbor area, the surroundings of Los Angeles International Airport, and downtown, were constructed near or over major oil fields (Figure 1a) meaning that many anthropogenic sources of CH4 are colocated with each other and with potential geologic CH4 sources.

Missing from stationary or airborne measurements is detailed spatial information about the distribution of CH4 sources that is needed for developing monitoring and mitigation strategies for urban CH4 emissions. Here we present data from an extensive on-road survey of CH4 levels in the Los Angeles Basin during June 2013. Our goals were twofold: to describe the spatial patterns of CH4 and other trace gases with urban sources in the Los Angeles Basin and to use local measurements of the C2H6 to CH4 ratio to attribute sources of fugitive CH4 emissions, measured as hot spots and in local background air. We demonstrate the utility of mobile surveys to locate and attribute urban CH4 hot spots, and of complementary measurements of C2H6, to perform a regional source apportionment.

2. Methods
2.1. Instrumentation

We used a mobile observatory system to continuously survey trace gas mole fractions on-road while also recording global position and winds. The platform was a 2011 Ford Transit Connect van with a modified
electrical system and sampling mast that extended to 3.5 m above the road surface located just behind the driver’s seat of the vehicle [Bush et al., 2015].

The observatory was equipped with two cavity ring-down spectrometers (Picarro Inc., Sunnyvale, California), one measuring CH₄, CO₂, and H₂O (G1301) and the other measuring CO, CO₂, and H₂O (G1302). The instruments were plumbed in serial, sampling air from a common inlet located at the top of the sampling mast. Outside air was pumped continuously through the line at a rate of 4.2 L/min. The G1301 reported measurements every 2–3 s and the G1302 every 3–4 s. The time delay for measurement of outside air ranged from 0.7 to 3 s, depending on the variable instrument sampling rate. During this campaign, the laboratory also had an Aerodyne Ethane Mini Monitor quantum cascade laser spectrometer on board to measure C₂H₆ [Yacovitch et al., 2014]. The Ethane Mini Monitor sampled from a separate inlet line to enable instrument background scans by purging with N₂ for 30 s every 15 min. In addition, the instrument measured a reference C₂H₆ cell for 3 s every 2 min to maintain a line lock on the C₂H₆ spectral feature. Apart from reference and background scan times, data were collected once a second, with a 1 s response time.

Position data were collected every 5 s by a GPS16X-HVS receiver (Garmin International, Inc., Olathe, Kansas) mounted on the vehicle’s roof, and wind direction and speed were measured with a Weather Station WS-200WM (Airmar Technology Corp., Milford, New Hampshire) mounted to the top of the sampling mast.

2.2. Calibration

We calibrated measurements made by cavity ring-down trace gas spectrometer using two NOAA-certified air standards that contained known amounts of CO₂, CO, and CH₄. Standards were measured for several minutes before and after each transect, with each transect defined as a set of on-road measurements made over the course of several hours. The relationship between known and measured standard values was applied as a linear correction to the data collected during that transect. Measurement precision was 1.7 ppm for CO₂, 2.1 ppb for CH₄, and 8 ppb for CO based on the standard deviation of all measured standards for the campaign. For CH₄, G1301 performance was found to be linear over a range of standard values from 1.7 to 10 ppm. Reported CH₄ values greater than 10 ppm hence represent an approximation (<0.01% of data reported here).
We report C$_2$H$_6$ data as spectroscopic mole fraction, which has an expected 1 s precision of 0.1 ppb on a moving mobile laboratory platform [Yacovitch et al., 2014]. We checked the accuracy of spectroscopic C$_2$H$_6$ by comparing it to C$_2$H$_6$ measured in the laboratory on five simultaneous whole air samples that were taken during periods when atmospheric C$_2$H$_6$ was relatively constant. The average of spectroscopic C$_2$H$_6$ from the whole air sampling period was within 10% of reported C$_2$H$_6$ values by gas chromatography-flame ionization detector [Colman et al., 2001].

2.3. Data Processing

Original data were collected at varying frequencies for each instrument ranging from once every second (Aerodyne Mini Monitor) to once every 3–4 s (Picarro G1302). We used correlation analysis of the two CO$_2$ data streams to adjust for time delays between the two Picarro instruments and CH$_4$ correlations to adjust for time delays between the Aerodyne Mini Monitor and Picarro G1301. All data were subsequently averaged to 5 s intervals.

For spatial analyses, 5 s averaged data were spatially gridded along 150 m road intervals by averaging observations taken within this increment. The 150 m segments were chosen to maximize the information content of gridded data; less than 3% of the 150 m segments required linear interpolation of the nearest neighbors to fill in gaps in data.

2.4. Background Trace Gas Mole Fractions

We determined urban excess values for each trace gas species by subtracting an estimated clean air background mole fraction from trace gas-enhanced measurements made in polluted urban air, following Blake et al. [1984]. We estimated the clean air background value for each trace gas by selecting the minimum observation for each transect in the study period and then averaging the lowest 20% of these transect minimums (i.e., the lowest four out of 21 transects). We estimated the uncertainty in background values as the standard deviation of the minimum trace gas measurement from all transects within the basin (n = 21). The background value for CH$_4$ was 1821 ± 24 ppb, CO was 80 ± 24 ppb, CO$_2$ was 396.3 ± 3.2 ppm, and C$_2$H$_6$ was 0.6 ± 0.2 ppb. Background values measured in the basin were all within 1 standard deviation of values reported at Kumukahi, Hawaii in June 2013 (June 2009 for C$_2$H$_6$) by NOAA’s Global Greenhouse Reference Network [Helmig et al., 2011; Dlugokencky et al., 2013a, 2013b; Novelli and Masarie, 2013]. At Kumukahi, CH$_4$ was 1836 ± 7 ppb, CO was 85 ± 5 ppb, and CO$_2$ was 399.8 ± 0.7 ppm from the average of flask samples on eight measurement days in June 2013, and C$_2$H$_6$ was 0.6 ± 0.1 ppb from the average of flask samples from five measurement days in June 2009. The background value for C$_2$H$_6$ was similar to observations made from aircraft above the Los Angeles Basin, 0.6 ± 0.3 ppb for the lowest values on three measurement days that coincided with on-road sampling.

2.5. Observations

We sampled approximately 1900 km of road in the Los Angeles Basin, in June and early July of 2013 (Figure 1). We conducted 21 transects, with each transect defined as a set of on-road measurements made within a specific time period (e.g., midday on 15 June). Most routes were repeated a minimum of two times, at midday (10 A.M. to 4 P.M.) and after nightfall (9 P.M. to 1 A.M.).

Transect routes were designed to characterize the C$_2$H$_6$/CH$_4$ ratios of known biogenic and fossil CH$_4$ sources, including landfills, wastewater treatment facilities, livestock, oil fields, refineries, natural gas storage, distribution infrastructure, and geologic sources (Figure 1a: map image of known CH$_4$ emitters in Los Angeles Basin). We also designed transect routes to cover common land use types within the Los Angeles Basin, with varying degrees of human influence (urbanization).

On 17–19 June, we coordinated on-road sampling to cover the same locations at approximately the same time as a concurrent aircraft campaign. The aircraft was a DC-8 flown out of NASA’s Dryden Flight Research Center over the Los Angeles Basin as part of the NASA Student Airborne Research Program (www.nserc.und.edu/sarp/sarp-2009-2013/2013). During flight, whole air samples were taken in evacuated 2 L stainless steel canisters. Canister samples were analyzed in the laboratory for CH$_4$, C$_2$H$_6$, CO, and CO$_2$ mole fractions and along with other constituents by gas chromatography-flame ionization detector [Colman et al., 2001]. We included only samples taken over land in the Los Angeles Basin at less than 3000 ft (914 m) above ground level in this analysis.
2.6. Hot Spot Identification

Plumes of air containing CH₄ values that exceeded the 95th percentile of CH₄ observations for that transect were considered indicators of a CH₄ hot spot. Some hot spots coincided with known emission sources (shown in Figure 1a), while others were of an unknown origin. We defined hot spots as road segments where at least one 150 m segment had a CH₄ value that exceeded the 95th percentile threshold (132 to 360 ppb above the local background level). The spatial extent of each hot spot was defined by the number of adjacent 150 m road segments that had CH₄ values above the local background level. Local background CH₄ levels varied over the course of each transect due to spatial variability and diurnal changes in boundary layer height and were thus determined by visual inspection of each transect. We approximated the amount of local CH₄ enhancement by summing excess CH₄ above the local background level (i.e., area under the curve) for each hot spot.

2.7. Source Apportionment

We used the range of mole fraction ratios of excess C₂H₆ to excess CH₄ from known CH₄ emission sources to apportion CH₄ in hot spots of unknown origin, and for local background air measured away from CH₄ hot spots, representing a regional mix of CH₄ source sectors. We used linear regression on excess C₂H₆ and excess CH₄ observations from CH₄ hot spots of known emitters to determine the range of C₂H₆/CH₄ ratios that characterized biogenic and fossil sources. Regression slope estimates included uncertainty in background mole fractions, estimated as 1 standard deviation of background levels [York et al., 2004]. In theory, biogenic sources

Table 1. Summary of Biogenic CH₄ Hot Spots Associated With Individual Facilities or Sites

<table>
<thead>
<tr>
<th>Source Type</th>
<th>Facility/Site</th>
<th>Date(s)</th>
<th>Time of Day</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Hot Spot Extent (km)</th>
<th>Average Excess CH₄ (ppm)</th>
<th>Maximum Excess CH₄ (ppb)</th>
<th>Maximum Excess C₂H₆ (ppb)</th>
<th>C₂H₆/CH₄ Slope (%)</th>
<th>Sample Size (n)</th>
<th>R² Value</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active</td>
<td>Puente Hills</td>
<td>22 June</td>
<td>A.M. and P.M.</td>
<td>34.02°</td>
<td>-118.00°</td>
<td>10.14</td>
<td>1.5 ± 2.2</td>
<td>22/25</td>
<td>3.19</td>
<td>0.4</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
</tr>
<tr>
<td>Scholl Canyon</td>
<td>22 June</td>
<td>A.M.</td>
<td>34.16°</td>
<td>-118.19°</td>
<td>0.92</td>
<td>0.8 ± 0.4</td>
<td>3.15</td>
<td>0.27</td>
<td>1.39</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>UCI landfill</td>
<td>24 June</td>
<td>A.M.</td>
<td>33.65°</td>
<td>-117.86°</td>
<td>0.44</td>
<td>0.12 ± 0.03</td>
<td>2.86</td>
<td>0.50</td>
<td>1.92</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>Palos Verdes landfill</td>
<td>24 June</td>
<td>A.M.</td>
<td>33.79°</td>
<td>-118.35°</td>
<td>1.39</td>
<td>0.05 ± 0.5</td>
<td>1.32</td>
<td>0.50</td>
<td>1.92</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>Cal Compact landfill</td>
<td>26 June</td>
<td>A.M. and P.M.</td>
<td>33.94°</td>
<td>-118.27°</td>
<td>1.93</td>
<td>0.4 ± 2.2</td>
<td>1.81</td>
<td>0.50</td>
<td>1.92</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>Calinlandfill</td>
<td>18 and 27 June</td>
<td>A.M. and P.M.</td>
<td>33.59°</td>
<td>-117.63°</td>
<td>3.86</td>
<td>3.9 ± 3.9</td>
<td>13.26</td>
<td>0.40</td>
<td>3.9 ± 3.9</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>Water treatment</td>
<td>14, 19, 22, 23, 24, 26, and 27 June</td>
<td>A.M. and P.M.</td>
<td>33.67°</td>
<td>-117.91°</td>
<td>5.3</td>
<td>0.7 ± 0.5</td>
<td>7.76</td>
<td>0.40</td>
<td>3.9 ± 3.9</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>Mesa Water District colored water treatment plant</td>
<td>14, 19, 22, 23, 24, 26, and 27 June</td>
<td>A.M. and P.M.</td>
<td>33.67°</td>
<td>-117.91°</td>
<td>5.3</td>
<td>0.7 ± 0.5</td>
<td>7.76</td>
<td>0.40</td>
<td>3.9 ± 3.9</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td>Orange County Sanitation District reclamation plant</td>
<td>14, 19, 22, 23, 24, 26, and 27 June</td>
<td>A.M. and P.M.</td>
<td>33.67°</td>
<td>-117.91°</td>
<td>5.3</td>
<td>0.7 ± 0.5</td>
<td>7.76</td>
<td>0.40</td>
<td>3.9 ± 3.9</td>
<td>0.00 ± 0.01</td>
<td>46</td>
<td>0.04</td>
<td>0.21</td>
</tr>
<tr>
<td><strong>Reported error is one standard deviation of the mean.</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Summary of Natural Gas CH4 Hot Spots Associated With Individual Facilities or Sites

<table>
<thead>
<tr>
<th>Source Type</th>
<th>Facility/Site</th>
<th>Date(s)</th>
<th>Time of Day</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Hot Spot Extent (km)</th>
<th>Average Excess CH4 (ppm)(^a)</th>
<th>Maximum Excess CH4 (ppb)</th>
<th>Maximum Excess C2H6 (ppb)</th>
<th>C2H6/CH4 Slope (%)(^b)</th>
<th>Sample Size (n)</th>
<th>R(^2) value</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power plants</td>
<td>Haynes steam plant</td>
<td>14 and 27 June</td>
<td>A.M. and P.M.</td>
<td>33.76</td>
<td>-118.096</td>
<td>15.0</td>
<td>0.07 ± 0.02</td>
<td>312</td>
<td>4.5</td>
<td>3.1 ± 0.5</td>
<td>19</td>
<td>0.24</td>
<td>0.03</td>
</tr>
<tr>
<td></td>
<td>AES Alamitos</td>
<td>22 June</td>
<td>P.M.</td>
<td>33.77</td>
<td>-118.10</td>
<td>0.8</td>
<td>0.27</td>
<td>482</td>
<td>9.7</td>
<td>3.8 ± 0.3</td>
<td>50</td>
<td>0.36</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>NRG</td>
<td>23 June</td>
<td>A.M.</td>
<td>33.91</td>
<td>-118.425</td>
<td>0.4</td>
<td>0.29</td>
<td>1,256</td>
<td>29.2</td>
<td>2.5 ± 0.7</td>
<td>5</td>
<td>0.61</td>
<td>0.12</td>
</tr>
<tr>
<td>CNG fueling stations</td>
<td>Clean Energy headquarters</td>
<td>17, 23, 24,</td>
<td>A.M. and P.M.</td>
<td>33.774</td>
<td>-118.077</td>
<td>1.9</td>
<td>0.3 ± 0.4</td>
<td>1,110</td>
<td>30.5</td>
<td>3.1 ± 0.1</td>
<td>38</td>
<td>0.94</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Clean Trucks, Port of Long Beach</td>
<td>26 June</td>
<td>A.M.</td>
<td>33.78</td>
<td>-118.222</td>
<td>0.9</td>
<td>2.1 ± 3.0</td>
<td>58,425</td>
<td>479.7</td>
<td>0.87 ± 0.03</td>
<td>141</td>
<td>0.81</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Clean Energy Newport Beach</td>
<td>14 June</td>
<td>A.M.</td>
<td>33.632</td>
<td>-117.927</td>
<td>0.6</td>
<td>0.1</td>
<td>203</td>
<td>1.8</td>
<td>1.2 ± 0.3</td>
<td>12</td>
<td>0.35</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>PH(^c) landfill truck gas fueling</td>
<td>22 June</td>
<td>A.M.</td>
<td>34.023</td>
<td>-118.028</td>
<td>0.6</td>
<td>2.2</td>
<td>11,365</td>
<td>117.0</td>
<td>0.9 ± 0.1</td>
<td>32</td>
<td>0.49</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Natural gas pipelines under roads</td>
<td>Pacific Coast Hwy at Superior, NB(^d)</td>
<td>5 July</td>
<td>A.M. and P.M.</td>
<td>33.623</td>
<td>-117.939</td>
<td>1.3</td>
<td>0.05 ± 0.01</td>
<td>315</td>
<td>9.5</td>
<td>2.8 ± 0.5</td>
<td>50</td>
<td>0.25</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Santa Ana Ave., Costa Mesa</td>
<td>15 June and 5 July</td>
<td>A.M. and P.M.</td>
<td>33.637</td>
<td>-117.911</td>
<td>3.0</td>
<td>0.06 ± 0.03</td>
<td>490</td>
<td>31.1</td>
<td>1.8 ± 0.1</td>
<td>74</td>
<td>0.27</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Goldenwest St., Huntington Beach</td>
<td>15 June</td>
<td>A.M.</td>
<td>33.74</td>
<td>-118.007</td>
<td>0.3</td>
<td>0.29</td>
<td>720</td>
<td>16.9</td>
<td>2.1 ± 0.1</td>
<td>56</td>
<td>0.80</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Campus and Carlson, Irvine</td>
<td>16 June</td>
<td>A.M.</td>
<td>33.664</td>
<td>-117.851</td>
<td>0.5</td>
<td>0.18</td>
<td>248</td>
<td>3.0</td>
<td>2.1 ± 0.5</td>
<td>5</td>
<td>0.78</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>91 Fwy at Buchanan, Corona</td>
<td>16 June</td>
<td>A.M.</td>
<td>33.895</td>
<td>-117.50</td>
<td>0.9</td>
<td>0.3</td>
<td>316</td>
<td>4.1</td>
<td>1.9 ± 0.6</td>
<td>8</td>
<td>0.96</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

\(^a\)Reported error is one standard deviation of the mean.

\(^b\)Slope is calculated by orthogonal distance regression; reported error is the largest of error estimates for slope from orthogonal distance regression and ordinary linear regression.

\(^c\)PH: Puente Hills.

\(^d\)NB: Newport Beach.
should have a ratio of 0 (i.e., no concurrent C2H6 production). In practice, many primarily biogenic sources had small but significant positive ratios because of measurement uncertainties and a small amount of contribution from nearby or colocated fossil sources. We used the observed range of excess C2H6 to excess CH4, hereafter denoted C2H6/CH4, from known sources to determine the likely origins of unknown hot spots based on their C2H6/CH4 slopes. For regional source apportionment, we included only observations of local background air measured away from CH4 hot spots, whereas we assumed that the air is well mixed with respect to CH4 and hence representative of a mixture of urban CH4 sources across the Los Angeles Basin.

3. Results

3.1. Spatial Distribution of CH4 and C2H6 Across the Los Angeles Basin

CH4 mole fractions were highly variable across the Los Angeles Basin, ranging from near background levels to 58 ppm measured at the Clean Trucks compressed natural gas (CNG) fueling station in the Port of Long Beach (Figure 1b and Map S1 in the supporting information). We observed local CH4 enhancement (hot spots) in the vicinity of many known CH4 emission sources, including active and closed landfills, cattle operations, water treatment facilities, geologic seeps, oil extraction and refining facilities, natural gas infrastructure, and gas-fired power plants (Tables 1–3). We also observed CH4 mole fractions highly variable across the Los Angeles Basin, ranging from near background levels to 58 ppm measured at the Clean Trucks compressed natural gas (CNG) fueling station in the Port of Long Beach. City air measured away from CH4 hot spots based on their C2H6/CH4 slopes. For regional source apportionment, we included only observations of local background air measured away from CH4 hot spots, whereas we assumed that the air is well mixed with respect to CH4 and hence representative of a mixture of urban CH4 sources across the Los Angeles Basin.

Table 3. Summary of Geologic CH4 Hot Spots Associated With Individual Facilities or Sites

<table>
<thead>
<tr>
<th>Source Type</th>
<th>Facility/Site</th>
<th>Date(s)</th>
<th>Time of Day</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Hot Spot Extent (km)</th>
<th>Average Excess CH4 (ppm)</th>
<th>Maximum Excess CH4 (ppm)</th>
<th>Maximum Excess C2H6 (ppb)</th>
<th>C2H6/CH4 Slope (%) ± Error</th>
<th>Sample Size (n)</th>
<th>R2 Value</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oil refineries</td>
<td>Conoco Phillips</td>
<td>24 June</td>
<td>A.M.</td>
<td>33.778</td>
<td>-118.29</td>
<td>0.3</td>
<td>0.24</td>
<td>1,781</td>
<td>38.9</td>
<td>2.8 ± 0.5</td>
<td>11</td>
<td>0.54</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>BP</td>
<td>23 and 26 June</td>
<td>A.M. and P.M.</td>
<td>33.82</td>
<td>-118.24</td>
<td>2.6</td>
<td>0.24 ± 0.01</td>
<td>347</td>
<td>8.8</td>
<td>3.5 ± 0.4</td>
<td>24</td>
<td>0.45</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Exxon</td>
<td>17 June</td>
<td>A.M.</td>
<td>33.85</td>
<td>-118.32</td>
<td>2.4</td>
<td>0.13</td>
<td>559</td>
<td>17.3</td>
<td>4.0 ± 0.2</td>
<td>116</td>
<td>0.64</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Chevron</td>
<td>19 and 23 June</td>
<td>A.M.</td>
<td>33.91</td>
<td>-118.40</td>
<td>3.1</td>
<td>0.2 ± 0.2</td>
<td>1,101</td>
<td>9.6</td>
<td>1.1 ± 0.1</td>
<td>69</td>
<td>0.35</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Oil fields</td>
<td>Newport Beach</td>
<td>14, 15, and 17 June and 5 July</td>
<td>A.M. and P.M.</td>
<td>33.626</td>
<td>-117.946</td>
<td>0.9</td>
<td>0.09 ± 0.03</td>
<td>223</td>
<td>1.7</td>
<td>-0.1 ± 0.1</td>
<td>35</td>
<td>0.00</td>
<td>0.79</td>
</tr>
<tr>
<td></td>
<td>Huntington Beach</td>
<td>15 June</td>
<td>A.M.</td>
<td>33.687</td>
<td>-118.005</td>
<td>2.8</td>
<td>0.09 ± 0.03</td>
<td>352</td>
<td>3.2</td>
<td>1.1 ± 0.2</td>
<td>20</td>
<td>0.47</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Seal Beach</td>
<td>14, 17, and 24 June</td>
<td>A.M. and P.M.</td>
<td>33.76</td>
<td>-118.11</td>
<td>1.4</td>
<td>0.2 ± 0.03</td>
<td>623</td>
<td>39.0</td>
<td>7.2 ± 0.6</td>
<td>38</td>
<td>0.48</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Long Beach</td>
<td>17, 24, and 26 June</td>
<td>A.M. and P.M.</td>
<td>33.81</td>
<td>-118.17</td>
<td>3.4</td>
<td>0.3 ± 0.3</td>
<td>855</td>
<td>19.4</td>
<td>2.7 ± 0.1</td>
<td>79</td>
<td>0.75</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Santa Fe Springs</td>
<td>22 June</td>
<td>A.M.</td>
<td>33.943</td>
<td>-118.065</td>
<td>1.5</td>
<td>0.08 ± 0.02</td>
<td>1,111</td>
<td>9.9</td>
<td>0.67 ± 0.05</td>
<td>118</td>
<td>0.48</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Inglewood</td>
<td>26 June</td>
<td>P.M.</td>
<td>34.00</td>
<td>-118.37</td>
<td>0.2</td>
<td>0.16</td>
<td>653</td>
<td>39.2</td>
<td>9 ± 5</td>
<td>3</td>
<td>0.58</td>
<td>0.45</td>
</tr>
<tr>
<td>Geologic seeps</td>
<td>Holmwood and Broad</td>
<td>5 July</td>
<td>A.M. and P.M.</td>
<td>33.626</td>
<td>-117.924</td>
<td>0.4</td>
<td>0.14 ± 0.03</td>
<td>1,352</td>
<td>2.3</td>
<td>0.01 ± 0.02</td>
<td>68</td>
<td>0.01</td>
<td>0.45</td>
</tr>
<tr>
<td></td>
<td>Newport Beach</td>
<td>19 and 23 June</td>
<td>A.M. and P.M.</td>
<td>33.973</td>
<td>-118.421</td>
<td>5.3</td>
<td>5 ± 9</td>
<td>27,201</td>
<td>65.6</td>
<td>0.14 ± 0.01</td>
<td>405</td>
<td>0.70</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>La Brea tarpits</td>
<td>19 and 26 June</td>
<td>A.M. and P.M.</td>
<td>34.063</td>
<td>-118.355</td>
<td>1.8</td>
<td>1 ± 1</td>
<td>5,021</td>
<td>23,757</td>
<td>4.04 ± 0.08</td>
<td>615</td>
<td>0.58</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

a Reported error is one standard deviation of the mean.
bSlope is calculated by orthogonal distance regression; reported error is the largest of error estimates for slope from orthogonal distance regression and ordinary linear regression.
cNB: Newport Beach.
of atmospheric C2H6 were also observed in many, but not all locations where CH4 enhancements were observed (Figure 1c).

Local CH4 background levels, excluding known and unknown CH4 hot spots, varied across the region (Figure 1b). In general, local CH4 background levels were higher in inland parts of the basin. For example, local background CH4 levels in Riverside were twice as high as they were in Irvine for the same day, three times as high in Ontario, and four times as high in Pasadena. Inland areas tend to accumulate polluted air generated in the basin due to prevalent onshore winds during spring and summer [e.g., Vutukuru et al., 2006; Peischl et al., 2013]. Local CH4 levels tended also to be higher in more urbanized areas of the basin, such as near downtown Los Angeles.

C2H6 also increased along a coast-to-inland gradient in a manner similar to CH4, with large areas of enhancement located in downtown Los Angeles, around the Long Beach oil field, and in the city of Ontario (Figure 1c). Local CH4 levels tended also to be higher in more urbanized areas of the basin, such as near downtown Los Angeles.

C2H6 mole fractions ranged from background levels similar to those observed in remote ocean regions to 2370 ppb measured at the La Brea Tar Pits. Local background levels of C2H6 varied even more than for CH4, with levels fourfold higher in Riverside and sixfold higher in Pasadena than in Irvine and up to 20 times higher in Ontario. The greater urban enhancements of C2H6 compared to CH4 were likely driven by significant urban C2H6 emissions and lower remote background mole fractions as a consequence of a shorter lifetime (approximately 2 months for C2H6 versus >10 years for CH4) and little C2H6 production in remote ocean regions [Xiao et al., 2008]. The ratio of C2H6 to CH4 was also highly variable across the region. In particular, C2H6/CH4 was markedly higher along surface streets in dense, highly urbanized areas near downtown Los Angeles and central Orange County (Figure 1d and Map S1).

### 3.2. C2H6/CH4 Ratio of Known Emitters

We made measurements of CH4 and C2H6 at known CH4 emission sources to characterize the C2H6/CH4 ratios of biogenic and fossil end-members and to enable subsequent use of this information to apportion CH4 emission sources in the Los Angeles Basin. We calculated C2H6/CH4 ratios for each known CH4 emitter and subsequently for each source sector as the slope of a line fit through a plot of excess C2H6 versus excess CH4 using a minimization of orthogonal distance and including measurement error in both variables (Tables 1–3). The C2H6/CH4 ratio for biogenic sources ranged from −0.05 ± 0.08% for former landfills to 0.02 ± 0.01% for water treatment facilities (Table 4). Most biogenic CH4 emitters did not have statistically significant C2H6/CH4 slopes (i.e., slopes were not statistically different from 0), except for the Puente Hills

---

**Table 4. Sectoral CH4 Hot Spot Characteristics**

<table>
<thead>
<tr>
<th>Source Category</th>
<th>Emitters Sampled</th>
<th>C2H6/CH4 Slope (%)</th>
<th>Percent of the Total Campaign Distance Driven</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biogenic Sources</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Active landfills</td>
<td>2</td>
<td>0.01 (0.01)</td>
<td>0.4</td>
</tr>
<tr>
<td>Former landfills</td>
<td>3</td>
<td>-0.05 (0.08)</td>
<td>0.5</td>
</tr>
<tr>
<td>Cattle</td>
<td>1d</td>
<td>0.00 (0.00)</td>
<td>1.5</td>
</tr>
<tr>
<td>Water treatment</td>
<td>1d</td>
<td>0.02 (0.01)</td>
<td>1.1</td>
</tr>
<tr>
<td>Biogenic average</td>
<td>7d</td>
<td>0.05 (0.03)</td>
<td>3.5</td>
</tr>
<tr>
<td><strong>Fossil Sources: Fugitive Natural Gas Leaks</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power plants</td>
<td>3</td>
<td>3.1 (0.9)</td>
<td>0.1</td>
</tr>
<tr>
<td>CNG fueling stations</td>
<td>4</td>
<td>1.5 (0.3)</td>
<td>0.4</td>
</tr>
<tr>
<td>NG pipeline leaks</td>
<td>5</td>
<td>2.1 (0.9)</td>
<td>0.3</td>
</tr>
<tr>
<td>Natural gas average</td>
<td>12</td>
<td>2.2 (0.4)</td>
<td>0.8</td>
</tr>
<tr>
<td><strong>Fossil Sources: Geologic Sources and Fossil Fuel Production</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oil refineries</td>
<td>4</td>
<td>2.9 (0.7)</td>
<td>0.4</td>
</tr>
<tr>
<td>Oil drilling</td>
<td>6</td>
<td>3.4 (5.0)</td>
<td>0.6</td>
</tr>
<tr>
<td>Geologic leaks</td>
<td>3</td>
<td>1.4 (0.1)</td>
<td>0.7</td>
</tr>
<tr>
<td>Geologic source average</td>
<td>13</td>
<td>2.8 (1.4)</td>
<td>1.7</td>
</tr>
<tr>
<td>Fossil average</td>
<td>25</td>
<td>2.5 (1.1)</td>
<td>2.5</td>
</tr>
</tbody>
</table>

aSectoral averaged (e.g., biogenic, fossil) shown in bold; source category sub-averages (e.g., natural gas, geologic source) shown in italic.
bSlope is calculated as the average of C2H6/CH4 for each emitter sampled; error on slope is propagated from error reported in Table 1.
cSingle region of cattle influence includes multiple individual dairies.
dExcludes one water treatment plant with suspected fossil emissions (Orange County Sanitation District Plant 1).
landfill and several water treatment facilities (Table 1). Significant C2H6/CH4 slope values were likely caused by collocated fossil emissions. For example, fossil-derived natural gas and biogenic CH4 may be combined to fuel natural gas vehicles. Both the Puente Hills landfill and the Orange County Sanitation District water treatment plant had CNG fueling stations on site.

Known fossil sources of CH4 were grouped into two categories based on their expected C2H6/CH4 ratios: (1) fugitive leaks of pipeline-quality natural gas, which include emissions from gas-fired power plants, CNG fueling stations, and natural gas pipelines (Table 2), and (2) unprocessed geologic CH4 sources, including oil fields, oil refineries, and geologic seeps (Table 3). Previous studies have shown a relatively narrow range of C2H6/CH4 ratios for pipeline natural gas compared to geologic CH4 sources [Jeffrey et al., 1991; Wennberg et al., 2012]. Fugitive natural gas leaks for sources sampled in this campaign had C2H6/CH4 ratios ranging from 1.5 ± 0.3% for CNG fueling stations to 3.1 ± 0.9% for natural gas-fired power plants (Table 4). Among pipeline gas sources, all C2H6/CH4 slopes were statistically significant at the p < 0.05 level except for one power plant sample with few (n = 5) measurements (Table 2). This range is consistent with direct measurements of C2H6/CH4 in pipeline gas in Southern California, 1.33–2.59%, reported by Wennberg et al. [2012]. C2H6/CH4 ratios for geologic CH4 sources tended to be higher than those for pipeline gas, ranging from 1.4 ± 0.1% for geologic seeps to 3.4 ± 5.0% for oil fields (Table 4). However, the range of C2H6/CH4 ratios for individual geologic sources was very broad, spanning −0.1 ± 0.1% to 9 ± 5% for oil fields and 0.01 ± 0.02% to 4.0 ± 0.1% for geologic seeps (Table 3). These C2H6/CH4 values were consistent with previous observations in oil fields, 0.7–12.0%, and geologic seeps, 0.8–7.5%, in the Los Angeles Basin [Jeffrey et al., 1991]. Several geologic sources had C2H6/CH4 ratios similar to biogenic sources (e.g., Newport Beach oil field, geologic seeps in Newport Beach, and Playa Vista) and may be due to biodegradation of higher hydrocarbons or mixing of thermogenic and biogenic natural gases, both of which have been observed in the Los Angeles Basin [Jeffrey et al., 1991].

3.3. Distribution of CH4 and Other Trace Gases

Atmospheric levels of trace gases were distributed unevenly across the basin, with high CH4 (and C2H6) values concentrated in CH4 hot spots (Figure 1). We compared the spatial distributions of the four measured gases with urban sources by their Gini coefficients, where 0 represents a perfectly equal distribution and 1 represents a perfectly unequal distribution [Gini, 1912]. CH4 was the most unevenly distributed trace gas over the basin, with a Gini coefficient of 0.55, followed by C2H6, 0.47; CO, 0.42; and CO2, 0.39.

To better understand patterns of emission variability, particularly for CH4 hot spots of unknown origin, we removed data collected in CH4 hot spots associated with known emission sources from further analysis. This subset of data comprised about 5.9% of the total distance covered in the campaign. We then sorted and ranked the remaining 94.1% of the observations according to their contribution to the total excess measured during the campaign (Figure 2). A considerable amount of excess CH4 measured in the Los Angeles Basin was associated with CH4 hot spots from unknown sources. Just 1% of the total distance traveled, excluding CH4 hot spots of known emission sources, accounted for 8% of excess CH4 measured across the basin. Similarly, the top 5% of distance traveled by CH4 mole fraction was responsible for 21% of the total excess CH4. Hot spots also contributed disproportionately to trace gas excess for C2H6 and CO; the top 1% of transect increments accounted for 7% of the total excess C2H6 and 8% of the total excess CO. CO2 was the most evenly distributed trace gas, with the top 1% of transect distance with respect to CO2 accounting for just 4% of the total excess CO2.

3.4. Variability in Spatial Patterns Over Time

We used repeated measurements of a section of the Pacific Coast Highway between Newport Beach and Seal Beach to determine how spatial patterns in atmospheric trace gas levels varied with time and under different wind conditions. We observed many CH4 hot spots in the same location across four different sampling days at four different times of day, despite differences in wind speed and direction (Figure 3).

We calculated the correlation coefficient for trace gas mole fractions between the four complete transects to quantify the repeatability of trace gas observations. CH4 was highly correlated in space among all four transect runs (correlation p values < 0.004), with an average correlation coefficient of 0.24 and a maximum correlation of 0.51 between the afternoon and night transects. These correlations were highly significant, given the large number of 150 m road segments on each transect (n = 174). CO2 was also well correlated among transects, with an average of 0.21 and a maximum correlation coefficient of 0.47 between the noon and
evening transects. CO had the lowest average correlation across transect pairs, with significant correlation observed for only three of six transect combinations \((p < 0.05)\). The high spatial correlation of \(\text{CH}_4\) values on repeated transects was consistent with emissions from persistent point sources, whereas lower correlations for \(\text{CO}_2\) and CO were consistent with more variable traffic emissions.

Trace gas excess mole fractions varied over the course of the day, with a different pattern observed for \(\text{CH}_4\) than for CO and \(\text{CO}_2\). \(\text{CH}_4\) mole fractions were lowest during midday and highest at night. This pattern was consistent with a higher planetary boundary layer and a well-mixed atmosphere in the afternoon that reduced local \(\text{CH}_4\) levels and more stable atmospheric conditions at night that trapped \(\text{CH}_4\) emissions near the surface. Diurnal patterns of CO and \(\text{CO}_2\) were subject to the same boundary layer effect; however, the highest values of CO and \(\text{CO}_2\) along this transect were observed in the 5–6 P.M. period, suggesting a time-varying emission source that was consistent with increased vehicle emissions during the evening rush hour.

The lowest trace gas excess values in this section of the Pacific Coast Highway were consistently measured while passing the Bolsa Chica Ecological Reserve, a salt marsh nature preserve (located between km 6 and km 12 of this road section and marked in orange in Figure 3). Reductions in trace gas levels around the marsh were particularly pronounced for \(\text{CH}_4\). No \(\text{CH}_4\) hot spots were observed in this area, suggesting that \(\text{CH}_4\) hot spots originated from built-up urban areas. Despite differences in wind direction and speed among measurement time periods, the consistently low \(\text{CH}_4\) mole fractions measured in the vicinity of Bolsa Chica demonstrate the sensitivity of the measurement technique to local emissions and provided qualitative evidence that the measurement footprint of observed hot spots in other parts of the basin was on the order of several kilometers or less.

### 3.5. Trace Gas Ratios of Unknown Hot Spots and Local Background Air

Both \(\text{C}_2\text{H}_6\) and \(\text{CH}_4\) were unevenly distributed in space; however, some of the locations with the highest \(\text{CH}_4\) values were unmatched by \(\text{C}_2\text{H}_6\) (Figure 4). The top 3% of transect distance with respect to \(\text{CH}_4\) values contained 15% of excess \(\text{CH}_4\) including unknown hot spots and local background air, but only 10% of excess \(\text{C}_2\text{H}_6\), suggesting a spatially varying combination of sources across the basin. To visualize this mixture, we calculated the ratio between \(\text{C}_2\text{H}_6\) and \(\text{CH}_4\), and between CO and \(\text{CO}_2\) for every 150 m transect segment.
Figure 3. Trace gas excess along a repeated coastal transect. (first panel) Map of route along Pacific Coast Highway beginning at 33.61°N, 117.89°W and ending at 33.76°N, 117.11°W (second panel) Excess CH₄, (third panel) excess CO₂, (fourth panel) excess CO, and (fifth panel) winds, corresponding to the locations shown in the first panel. Trace gas measurements along this transect were made at five different times. Orange highlighted section of map and plots shows section of road passing through a natural reserve (salt marsh), while the remainder of road passed through urbanized land.

Figure 4. Spatial distribution of CH₄ compared to distribution of C₂H₆ at the same locations. Distribution includes excess CH₄ values in unknown CH₄ hot spots and local background air. Height of bars represents percent of excess CH₄ for each 1% of distance traveled (blue bars) and corresponding percent of excess C₂H₆ for the same locations (red bars).
and plotted the ratios as a fraction of the total CH<sub>4</sub> or CO<sub>2</sub> excess (Figure 5). The bimodal C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> distribution suggests two different source processes contributing to atmospheric CH<sub>4</sub> observations, with a small peak centered on a C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> mole ratio less than 0.5%, reflecting biogenic CH<sub>4</sub> sources, and a second broad peak centered on 1.2%, reflecting fossil CH<sub>4</sub> sources (Figure 5a). Much of the distribution falls along the expected ratio for natural gas in the Los Angeles Basin, 1.33–2.59%, reported by Wennberg et al. [2012].

In contrast, a plot of the ratio of CO to CO<sub>2</sub> had only one peak, centered between 0.44% and 0.66% (Figure 5b). This distribution is similar to CO/CO<sub>2</sub> ratios measured in Irvine during the winter of 2007–2008 by Djuricin et al. [2010] (0.65–1.0%) and the range observed in Pasadena during summer of 2010 by Newman et al. [2013] (0.2–2.0%). Very little excess CO<sub>2</sub> (1%) had a CO/CO<sub>2</sub> ratio less than 0.2%, as would be expected from biogenic CO<sub>2</sub> sources. Hence, the CO/CO<sub>2</sub> ratio suggests that the vast majority of CO<sub>2</sub> measured during the on-road campaign is not representative of terrestrial ecosystem sources, and observed variation was likely driven by combustion sources.

### 3.6. Source Apportionment of Unknown Hot Spots

We used C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> ratios to apportion excess CH<sub>4</sub> measured at hot spots of unknown origin to biogenic and fossil sources (Figure 6). Unknown CH<sub>4</sub> hot spots, defined as spatially contiguous observations where at least one CH<sub>4</sub> observation exceeded the 95th percentile for that transect with no obvious emission source, comprised 5.8% of the total distance traveled. We used the highest C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> ratio observed for known biogenic sources, 0.50%, as the upper limit for possible C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> ratios from unknown biogenic sources. C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> slopes for CH<sub>4</sub> hot spots falling beneath this value were classified as biogenic. Similarly, we used the lowest C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> excess ratio measured from fugitive natural gas sources, 0.87%, as the lower limit for identifying fossil-derived CH<sub>4</sub> sources. Forty of 213 unknown hot spots were biogenic, 161 were fossil, and 11 had an intermediate slope and hence were considered indistinguishable.

### 3.7. Source Apportionment of Urban Background Air

The remaining 88.3% of distance traveled, excluding known and unknown CH<sub>4</sub> hot spots, was assumed to represent a more integrated measurement of excess CH<sub>4</sub> sources across the basin. The C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub> ratio measured in these samples was similar to that measured by aircraft over the same locations during three days of coincident sampling (Figure 7), 3.1 ± 0.8% for aircraft versus 3.2 ± 0.1% for on-road on 17 June, 1.6 ± 0.3% for aircraft versus 1.4 ± 0.2% for on-road on 18 June, and 1.7 ± 1.2% for aircraft versus 2.7 ± 0.6% for on-road on 19 June. C<sub>2</sub>H<sub>6</sub>/CH<sub>4</sub>
Figure 6. C$_2$H$_6$/CH$_4$ relationship for source apportionment. Excess C$_2$H$_6$/CH$_4$ values for all (a) known hot spots, (b) unknown hot spots, and (c) well-mixed local background air. For known hot spots, data collected at biogenic CH$_4$ sources are indicated by red dots, and data collected at fossil CH$_4$ sources are indicated by blue circles. Solid lines show criteria slope values, showing the maximum biogenic slope (red) and minimum fossil slope (blue). Dashed lines are best fit lines for known biogenic (red) and fossil (blue) hot spots. For unknown hot spots and local background air, data are plotted with blue circles. Values attributed to biogenic sources are marked with red dots, and values attributed to fossil sources are marked with green dots. Insets show same data for Figures 6a–6c but at a reduced scale.
values of this well-mixed, local background air were often higher than the minimum C_{2}H_{6}/CH_{4} ratio observed for natural gas sources, suggesting that most of excess CH_{4} present in urban air was fossil derived (Figure 7).

To apportion CH_{4} sources in local background air, we compared observed C_{2}H_{6}/CH_{4} ratios to simulated C_{2}H_{6}/CH_{4} values for varying proportions of biogenic and fossil CH_{4} inputs. We represented C_{2}H_{6}/CH_{4} values of biogenic sources as a random, normal distribution with a mean of 0% and a standard deviation of 0.03%, based on the average and standard error of C_{2}H_{6}/CH_{4} values observed at known biogenic hot spots. Similarly, we constructed a normal distribution of C_{2}H_{6}/CH_{4} values with a mean of 2.5% and standard deviation of 1.1% for fossil sources (Table 4). We determined the fossil fraction that best simulated the observed local background C_{2}H_{6}/CH_{4} distribution by performing 15,000 mixing trials for each fossil fraction, ranging from 0 to 1, with
end-member ratios randomly selected using the mean and standard deviations of biogenic and fossil emission ratio data described above. We found that a mean fossil fraction of 62% (59–64%) best matched \( \text{C}_2\text{H}_6/\text{CH}_4 \) values in local background air according to Kolmogorov-Smirnov and Cramer-von Mises test statistics. The range of simulated fossil fraction values includes uncertainty in measured and background \( \text{CH}_4 \) and \( \text{C}_2\text{H}_6 \) values, by refitting simulations to \( \text{C}_2\text{H}_6/\text{CH}_4 \) observations plus or minus propagated error from those sources.

In addition to measurement error, variation in \( \text{C}_2\text{H}_6/\text{CH}_4 \) end-member values introduces uncertainty into the inferred fossil fraction. We examined the role of this variation by performing similar simulations to those described above but assuming different fossil end-member \( \text{C}_2\text{H}_6/\text{CH}_4 \) values (we do not expect biogenic end-members to differ from 0). Modifying the fossil fraction to reflect only natural gas sources (mean and standard deviation of 2.2 ± 0.4%), the optimal fossil fraction increased to 64%. For the case of only geologic sources (mean and standard deviation of 2.8 ± 1.4%), the inferred fossil fraction decreased to 58%. Nevertheless, neither of these scenarios were able to match the data as well as the fossil end-member that includes both sources (Figure 8). We also separately simulated three hypothesized \( \text{CH}_4 \) sources: biogenic, natural gas, and geologic, with the same \( \text{C}_2\text{H}_6/\text{CH}_4 \) values used in the two-source simulations. The combination of the three sources had a best fit fossil fraction of 65% (Figure 8) and was better able to reproduce the high frequency of \( \text{C}_2\text{H}_6/\text{CH}_4 \) values clustered around 1.2%, as well as the large variability in \( \text{C}_2\text{H}_6/\text{CH}_4 \) greater than 3%. However, the three-source mixture case is poorly constrained because of the need to fit additional parameters.

4. Discussion

4.1. Methane Source Apportionment in Los Angeles

Mobile laboratory on-road sampling enabled us to make high-frequency, local-scale measurements of multiple trace gases across the Los Angeles Megacity region. We directly measured emission ratios of different sources, at the scale of individual facilities, and simultaneously obtained regionally integrated measurements of multiple trace gases in the urban air mass. Highly spatially resolved \( \text{CH}_4 \) and \( \text{C}_2\text{H}_6 \) data allowed us to determine the relative contributions of different emission sectors to regional urban \( \text{CH}_4 \) excess. These data offer further evidence for an under inventoried and dispersed set of fossil \( \text{CH}_4 \) sources to the Los Angeles atmosphere. We also identified targets for future work and potential mitigation of \( \text{CH}_4 \) emissions.

We measured \( \text{CH}_4 \) enrichment at 33 individual point sources to determine the range of \( \text{C}_2\text{H}_6/\text{CH}_4 \) ratios for biogenic or fossil sources. As expected, observed \( \text{C}_2\text{H}_6/\text{CH}_4 \) of known biogenic sources tended to be very low compared to other observations, despite a few locations with higher than expected \( \text{C}_2\text{H}_6 \) levels (e.g., Orange County Sanitation District Water Reclamation Plant). Known fossil \( \text{CH}_4 \) emission sources had varied \( \text{C}_2\text{H}_6/\text{CH}_4 \) signatures, reflecting the wide range of \( \text{C}_2\text{H}_6/\text{CH}_4 \) signatures of geologic sources in the basin. In contrast, local background air had a \( \text{C}_2\text{H}_6/\text{CH}_4 \) signature distinct from hot spot biogenic and fossil sources, clearly showing contributions of natural gas and oil sources of \( \text{CH}_4 \).
Our attribution of 62% of Los Angeles CH₄ to fossil sources, with a possible range of 58–65%, is within the range of other studies using atmospheric C₂H₆/CH₄ measurements in Los Angeles, with estimates of 56% by Peischl et al. [2013] and 70% by Wennberg et al. [2012]. Our analysis gives insight into why such a large range has been inferred by previous work. First, we found that the inferred contribution of fossil sources depends strongly on the choice of the C₂H₆/CH₄ ratio of the fossil end-member. Wennberg et al. [2012] hypothesized that the majority of CH₄ in the Los Angeles Basin originated from leaks of pipeline natural gas and may have overestimated the fossil contribution by assuming that all fossil CH₄ had a C₂H₆/CH₄ ratio representative of natural gas. In contrast to aircraft measurements, our sample of urban air had a much wider range of C₂H₆/CH₄ values that are outside the range of natural gas sources measured here or in previous work in the region [Wennberg et al., 2012], demonstrating that fossil CH₄ sources in the Los Angeles Basin must be a mixture of pipeline and unprocessed gas emissions.

Most of the variability in fossil C₂H₆/CH₄ ratios we observed is likely to stem from the large variation in geologic CH₄ sources, which ranged from 0 to 9%. We found that geologic seeps had particularly low C₂H₆/CH₄ values, with two out of three surveyed geologic seeps with C₂H₆/CH₄ values <0.15%. Hence, geologic sources may be confounded for biogenic sources without specific knowledge of the emission source in a particular location. It is possible that the analysis of Peischl et al. [2013] underestimated geologic emissions, as some geologic seeps have C₂H₆/CH₄ ratios that are indistinguishable from biogenic sources.

4.2. Strengths and Weaknesses of Mobile Laboratory Technique

Discrepancies between top-down measurements and bottom-up inventories present a major challenge for greenhouse gas mitigation policy and planning, particularly for CH₄. Mobile laboratories are particularly suited for addressing this challenge, having the ability to make complementary regional top-down CH₄ measurements [Petron et al., 2012] and facility-level bottom-up measurements [Jackson et al., 2014]. In this study, regional C₂H₆/CH₄ apportionment of local background air demonstrated that the predominant source of CH₄ emissions is fossil. We also found that the majority of unknown CH₄ hot spots were of fossil origin, providing further evidence for the importance of dispersed, fugitive fossil CH₄ emissions in the Los Angeles Basin. These CH₄ hot spots of unknown source represent an important target for future research. Repeated measurements of these unknown hot spots are needed to determine whether they persist in time and with tools that can pinpoint the precise locations of CH₄ leaks for source attribution.

Our regional apportionment attributed a large portion of CH₄ in the Los Angeles atmosphere to emissions of pipeline-quality natural gas; however, we observed few discrete leaks from natural gas pipelines. This contrasts with recent observations of frequent pipeline leaks detected by on-road sampling in Boston and Washington, DC [Phillips et al., 2013; Jackson et al., 2014] and is consistent with recent work showing relatively fewer pipeline gas leaks in the western U.S. [Lamb et al., 2015]. Fugitive gas emissions in those cities were found in areas with cast iron gas mains, which are not present in Los Angeles [Southern California Gas Company, 2011]. The most significant fugitive emissions of natural gas we observed came from compressed natural gas fueling stations. Fugitive emissions associated with natural gas fueling infrastructure are not currently included in the California Air Resources Board inventory or in prominent life cycle assessment models [California Air Resources Board, 2011; A. Burnham, personal communication, 2014]. A better understanding of which components of CNG fueling are leaking is needed for emission quantification and mitigation.

Closed landfill sites also had surprisingly large emissions, with measureable increases in atmospheric CH₄ levels near most sites surveyed (Table 1). Atmospheric CH₄ enhancements were also observed at active and several closed landfill sites that had landfill gas mitigation systems in place, suggesting that the effectiveness of these mitigation activities needs to be verified by atmospheric measurement.

While we were able to make facility-level measurements to identify sources and C₂H₆/CH₄ ratios of urban CH₄ emissions, we acknowledge that a major drawback to on-road sampling is the challenge of making representative measurements. Local atmospheric CH₄ levels strongly depend on proximity to emission source; however, we were not always able to directly access CH₄ emitting facilities, such as inside oil refineries or on landfill surfaces. Here we used measurements away from CH₄ emission sources (local background air) to overcome the limitations of our sampling approach. For this reason, quantitative flux estimates from mobile laboratory sampling in urban areas are extremely challenging. Future studies can reduce this bias by combining multiple techniques along with on-road sampling, including continuous measurement from...
tall towers, aircraft, and total column trace gas measurements along with direct flux measurements such as chambers or eddy covariance. Another potential bias in our sample may be due to more extensive coverage in the western portions of the basin, which are densely populated and have the most concentrated oil and gas infrastructure.

Mobile laboratory sampling has significant cost and logistic advantages over aircraft. Unlike aircraft, mobile measurements are also able to target specific CH$_4$ sources and observe a variety of source mixtures. Nevertheless, proximity to emission sources can also be a disadvantage. Aircraft and remote sensing studies can use the relationship between CH$_4$ and CO or CO$_2$ in regionally representative air masses, along with inventories of those gases to estimate CH$_4$ emissions [e.g., Wennberg et al., 2012; Wong et al., 2015]. With on-road sampling, however, we found poor relationships between CH$_4$ and CO, and CH$_4$ and CO$_2$, even in well-mixed local background air, likely due to our proximity to on-road sources of CO and CO$_2$. We found no evidence of vehicle emissions on our CH$_4$ and C$_2$H$_6$ measurements. No CH$_4$ hot spots coincided with areas of high traffic emissions, including roadway tunnels.

4.3. Urban Pattern of CH$_4$ (and C$_2$H$_6$) Distinct From CO and CO$_2$

The differing spatial patterns of these four long-lived trace gases demonstrate that different measurements, monitoring, and mitigation approaches are needed for different urban greenhouse gases. Excess CH$_4$ was relatively concentrated in space compared to CO$_2$, and large CH$_4$ enhancement measured in the vicinity of the strongest hot spots suggests that they contribute significantly to basin-wide CH$_4$ emissions. It may be easier for mitigation efforts to target these point emission sources to achieve equivalent reduction in radiative forcing as reducing more diffuse CO$_2$ emissions from a whole system. More work is needed to quantify the relative importance of emissions from these CH$_4$ hot spots.

In addition to monitoring hot spots, the discrepancy between inventory and atmospheric measurement also suggests that wide-ranging measurements must also be made to capture the effects of distributed CH$_4$ sources. We observed basin-wide enhancement of atmospheric CH$_4$ levels, a so-called urban dome, but found that the distribution of CH$_4$ within the city is more strongly controlled by proximity to myriad emission sources. The mobile laboratory approach demonstrated that these points of emission can be linked with individual emitters and attributed to anthropogenic sources. The repeatability of hot spot locations suggests that there are significant, discrete CH$_4$ emission sources that can be targeted by mitigation efforts. Future work is needed to determine the mechanisms of these leaks (e.g., pipeline seam weld leaks versus fitting leaks), the cost of repair, and an effective strategy for reducing the most critical CH$_4$ sources. Finally, the fine spatial scales of 10 s to 100 s of meters at which CH$_4$ hot spots occur suggest that a mobile sampling strategy should be an integral part of a city- or regional-scale greenhouse gas measurement effort.

Acknowledgments

Original geolocated, time-stamped trace gas measurements made by mobile laboratory are provided as Data Set S1. This research is funded by the U.S. Department of Energy’s Office of Science (BER) under grant DE-SC0005266. This work was supported in part by the W. M. Keck Institute for Space Studies. We thank Valerie Carranza, Joshua Miu, Mariela Ruacho, Kristal Verhulst, Josette Marrero, and Tianyang Zhu for help with field work and Simone Meinardi for help in the laboratory.

References


Cardno ENTRIX (2012), Hydraulic fracturing study, PXP Inglewood oil field report, 206 pp., Plains Exploration & Production Company and Los Angeles County, Department of Regional Planning, Los Angeles, Calif.


Southern California Gas Company (2011), Pipeline safety activities performed by Southern California gas company within the County of Los Angeles report, County of Los Angeles Department of Public Works, Los Angeles, Calif.


