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Key Points:

- Multisite observations constrain both urban and rural CH₄ emissions
- California total emissions are likely 1.2–1.8 times state inventory
- More efforts are needed to constrain emissions by both sector and region

Supporting Information:

Supporting Information S1

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Estimating methane emissions in California's urban and rural regions using multitower observations

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Abstract We present an analysis of methane (CH_4) emissions using atmospheric observations from 13 sites in California during June 2013 to May 2014. A hierarchical Bayesian inversion method is used to estimate CH₄ emissions for spatial regions (0.3° pixels for major regions) by comparing measured CH₄ mixing ratios with transport model (Weather Research and Forecasting and Stochastic Time-Inverted Lagrangian Transport) predictions based on seasonally varying California-specific CH₄ prior emission models. The transport model is assessed using a combination of meteorological and carbon monoxide (CO) measurements coupled with the gridded California Air Resources Board (CARB) CO emission inventory. The hierarchical Bayesian inversion suggests that state annual anthropogenic CH₄ emissions are 2.42 ± 0.49 Tg CH₄/yr (at 95% confidence), higher (1.2–1.8 times) than the current CARB inventory (1.64 Tq CH₄/yr in 2013). It should be noted that undiagnosed sources of errors or uncaptured errors in the model-measurement mismatch covariance may increase these uncertainty bounds beyond that indicated here. The CH₄ emissions from the Central Valley and urban regions (San Francisco Bay and South Coast Air Basins) account for ~58% and 26% of the total posterior emissions, respectively. This study suggests that the livestock sector is likely the major contributor to the state total CH₄ emissions, in agreement with CARB's inventory. Attribution to source sectors for subregions of California using additional trace gas species would further improve the quantification of California's CH₄ emissions and mitigation efforts toward the California Global Warming Solutions Act of 2006 (Assembly Bill 32).

1. Introduction

California has committed to an ambitious plan to reduce greenhouse gas (GHG) emissions to 1990 levels by 2020 through Assembly Bill 32 (AB-32), which requires accurate accounting of CH₄ emissions for effective mitigation planning and verification of future emission reductions. The state official GHG inventory reports that California currently emits a total of approximately 459.3 Tg CO₂ (1 Tg = 10^{12} g) equivalent GHGs each year [*California Air Resources Board (CARB)*, 2015]. The CARB GHG inventory is produced in support of AB-32, thus only includes anthropogenic emission sources. Among the reported GHGs, ~9% of the total GHG emissions are attributed to methane (CH₄), which is the second largest contributor to climate forcing emissions in California behind carbon dioxide (CO₂) [*CARB*, 2015]. Moreover, as shown in previous studies [e.g., *Jeong et al.*, 2013, 2014] CH₄ emissions in California are relatively uncertain compared to those of CO₂ due to lack of activity data and incomplete understanding of emission processes, and top-down studies can be complicated by California's diverse emission sources, complex topography, and weather patterns.

Several recent studies have estimated CH₄ emissions in different regions of California using measurements from ground towers, aircrafts, and satellites. At the regional scale, *Zhao et al.* [2009] and *Jeong et al.* [2012a, 2013] estimated CH₄ emissions using towers in the Central Valley. In particular, *Jeong et al.* [2013] conducted the first multisite analysis of CH₄ emissions in California based on measurements from five ground sites and across seasons (10 months during 2010–2011) and estimated a state total of 2.03–2.71 Tg CH₄/yr (at 68% confidence). *Wecht et al.* [2014] used airborne measurements during a short-period campaign

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Site	Location	Latitude	Longitude	Inlet Height (m agl) ^a	Data Availability
ARV	Arvin	35.24	-118.79	10	June 2013 to May 2014
CIT	Caltech, Pasadena	34.14	-118.12	10	June 2013 to May 2014
LVR	Livermore	37.67	-121.71	27	June 2013 to May 2014
MAD	Madera	36.87	-120.01	10	June 2013 to May 2014
STB	Sutter Buttes	39.21	-121.82	10	June 2013 to May 2014
STR	San Francisco	37.76	-122.45	232	June 2013 to May 2014
THD	Trinidad Head	41.05	-124.15	20	June 2013 to August 2013
TRA	Tranquility	36.63	-120.38	10	June 2013 to April 2014
TSB	Tuscan Buttes	40.26	-122.09	10	June 2013 to May 2014
VTR	Victorville	34.61	-117.29	90	June 2013 to August 2013
WGC	Walnut Grove	38.27	-121.49	91	June 2013 to May 2014
SBC	San Bernardino	34.09	-117.31	58	June 2013 to May 2014
SIO	Scripps Institution of Oceanography	32.87	-117.26	10	June 2013 to May 2014

Table 1. GHG Sites Information Across California

^aInlet heights used in the inversion.

(May–June 2010) and estimated a total of 2.65–3.07 Tg CH₄/yr (at 68% confidence) based on a different prior emission model that resulted in a different source apportionment from that of *Jeong et al.* [2013], attributing significantly higher emissions to landfill and wastewater.

At the subregional scale, most studies focused on the urban regions of Southern California [e.g., *Wunch et al.*, 2009; *Hsu et al.*, 2010; *Wennberg et al.*, 2012; *Peischl et al.*, 2013]. Although the urban studies relied on different analysis methods (e.g., ratio of CH₄ to CO (carbon monoxide)) and measured data from different years, the focus region for each study generally covered the South Coast Air Basin (SoCAB) of California. For SoCAB, the estimated CH₄ emissions ranged from 280 to 700 Gg CH₄/yr (based on the reported uncertainty estimates; $1 \text{ Gg} = 10^9 \text{ g}$). In another study, *Jeong et al.* [2014] estimated statewide CH₄ emissions from petroleum production and the natural gas system, taking a unique approach of combining a bottom-up inventory with results from a field campaign.

Here we expand on previous work by Jeong et al. [2012a, 2013, 2014] to quantify both urban and rural CH₄ emissions from California, presenting the first analysis of full annual CH₄ emissions from California using atmospheric observations from 13 tower sites covering all major CH₄-emitting regions of California. In particular, this study uses the hierarchical Bayesian approach introduced by Ganesan et al. [2014] for the purpose of GHG emission quantification. The hierarchical approach has the advantage of being less reliant on choices of parameters such as the prior uncertainty and model-measurement mismatch uncertainty because they are estimated from probability distributions using additional parameters (i.e., hyperparameters). Moreover, these parameters are fully propagated in the inverse optimization for estimating emission fluxes. The hierarchical approach is a preferred choice for pixel-based (i.e., high resolution) inversions as in this study because hierarchical models are more favorable with the case where there are more parameters than data points [Gelman et al., 2014]. Hierarchical models have been widely applied to other similar fields [e.g., Clark, 2003; Riccio et al., 2006; Moraes et al., 2014], and this study demonstrates the utility of the hierarchical approach for GHG emission estimation. In particular, we illustrate how uncertainty in the inversion can be treated by a combination of our best a priori knowledge of error sources (e.g., transport error) and statistical inference and how groundbased multitower measurements can be effectively used to constrain regional emissions. In section 2, we describe the methods we employed, including atmospheric measurements, a priori CH₄ emissions, transport modeling, and the hierarchical Bayesian inverse method. Section 3 presents results, including the inferred CH₄ emissions from California for different regions and sources. Section 4 further discusses the results, and section 5 presents the conclusions for CH₄ emissions in California.

2. Data and Methods

2.1. CH₄ Measurements and Background

CH₄ measurements were made at the collaborative 13-site GHG network across California during June 2013 to May 2014. The information of sites and data availability is summarized in Table 1 (see Figure 1 for site locations). Detailed information regarding measurement methods for the Central Valley sites are summarized in



Figure 1. (a) CALGEM total (1.7 Tg CH₄/yr, 1 Tg = 10^{12} g) prior emissions (nmol/m²/s) with locations of measurement sites across California and (b) region classification (California air basins).

Jeong et al. [2012a, 2013] and Andrews et al. [2014]. Here we briefly describe measurements as a component of the inverse modeling framework. All sites are operated with temperature- and pressure-controlled cavity ring-down CH₄ gas analyzers (Picarro Inc.), permeation-tube gas sample driers, and periodic calibrations using either primary National Oceanic and Atmospheric Administration (NOAA) CH₄ gas standards or secondary gas standards. For this study, we added four new sites in Southern California: Caltech, San Bernardino, Scripps Institution of Oceanography, and Victorville (VTR), and two sites for the San Francisco Bay Area (SFBA): Livermore and Sutro Tower (STR). All new sites except STR had similar instrumentation to existing sites, while STR employed daily flask samples collected for approximately 2 min near 14:00 local standard time (LST) for subsequent analysis at NOAA Earth System Research Laboratory. Measurements at Trinidad Head (THD) were made by a flame ionization gas chromatography (FIGC) system as part of the Advanced Global Atmospheric Gases Experiment (AGAGE) network [Prinn et al., 2000]. The Tohoku University calibration scale used by AGAGE is indistinguishable from the NOAA04 calibration scale used for our Picarro measurements, with a relative scale factor of 1.0003 [Hall et al., 2014]. Thus, no corrections for scale differences were applied. In addition, we assume that the isotopic effect in transferring the NOAA standards that are calibrated by FIGC measuring all CH₄ isotopologues to the Picarro instrument (measuring only the predominant CH₄ isotopologue) is negligible. For continuous measurement sites, calibrated data were averaged to hourly intervals and then 3 hourly intervals for inversions following the procedure in Jeong et al. [2012a, 2013]. All sites are expected to provide measurement precision that is smaller than the CH₄ synoptic variations typically observed in the ambient air and with absolute accuracy sufficient to provide negligible bias in estimating the scaling relationship between observed and predicted CH₄ signals.

Following previous work [e.g., Haszpra, 1999; Matross et al., 2006; Gourdji et al., 2010; Miller et al., 2013; Jeong et al., 2013], we selected measurements that coincided with periods when the atmospheric boundary layer was well mixed to avoid the impact of large uncertainties associated with modeling the nocturnal boundary layer. For the Walnut Grove Tower (WGC) we explicitly evaluated atmospheric mixing using measured vertical CH₄ profiles. As in Jeong et al. [2012a, 2013], WGC data from 91 m were selected in the time window between 12:00 and 17:00 LST, subject to the requirement that the CH₄ mixing ratio difference ($C_{91}-C_{483}$) between 91 and 483 m fell within the range of -1 SD < ($C_{91}-C_{483}$) < 3 SD, where SD is the standard deviation of the 91–483 m difference. This additional requirement retained approximately 80% of data in the 12:00–17:00 LST window. We selected all data in the afternoon time window (12:00–17:00 LST) for other sites without profile information.

The predicted CH_4 upstream boundary values were estimated using a similar method to the one used in *Jeong et al.* [2012b, 2013]. The details for estimating the boundary values are described in *Jeong et al.* [2013], and only a summary is provided here. CH_4 boundary values were estimated using data from the

Pacific coast aircraft network CH₄ profiles (http://www.esrl.noaa.gov/gmd/ccgg/aircraft/) and remote Pacific marine boundary layer sampling sites (http://www.esrl.noaa.gov/gmd/ccgg/flask.html) within the NOAA Earth System Research Laboratory Cooperative Air Sampling Network. The data were smoothed and interpolated to create a three-dimensional (3-D) curtain, varying with latitude, height, and time. To quantify the errors associated with the 3-D curtain, we fit a smooth curve through the data and computed the seasonal cycle of the root mean square of the residuals from the curve. Predicted background values were computed for each hourly footprint simulation by sampling the curtain at each of the 500 particle trajectory endpoints (near the domain boundary at 130°W) and calculating the average value.

2.2. A Priori CH₄ Emission Model

This work used the California Greenhouse Gas Emission Measurements (CALGEM) project a priori CH₄ emission model (henceforth CALGEM model, available at calgem.lbl.gov) described by *Jeong et al.* [2012a, 2013, 2014] with some modifications. The CALGEM emission model provides emissions by sector at a high spatial resolution $(0.1^{\circ} \times 0.1^{\circ})$ for California. The CALGEM model has seasonal components for wetlands and crop agriculture only, and these seasonal emissions are combined with nonseasonal emissions to construct monthly emission maps for inversions. The inversion approach using nonseasonal prior emissions is widely used [e.g., *Zhao et al.*, 2009; *Jeong et al.*, 2012a, 2012b, 2013; *Wecht et al.*, 2014; *Cui et al.*, 2015; *Turner et al.*, 2015]. In particular, *Jeong et al.* [2012a, 2012b, 2013] showed that nonseasonal priors can provide information on seasonality in the posterior emission.

In this study, the CALGEM prior emission distributions are scaled to match 2012 CARB state totals for anthropogenic emission sectors [*CARB*, 2014], with small (<50 Gg CH₄/yr) adjustments for some regions and sectors (per ARB staff private communication). The spatial distribution of the dairy livestock emissions was revised by incorporating the 2012 county-level dairy statistics from U.S. Department of Agriculture (http://www.nass. usda.gov/Statistics_by_State/California/Publications/County_Estimates/) to the spatial distribution from *Jeong et al.* [2013]. This revision changed the dairy livestock emissions for each region due to recent changes in the number of dairy cows, in particular for SoCAB. The current dairy livestock emissions in SoCAB (Table 2) decreased by ~50% compared to those (~80 Gg CH₄/yr) of *Jeong et al.* [2013], which was based on the 2004 statistics reported in *Salas et al.* [2009], reflecting the decrease in the number of dairy cows in the region (see Figure S1 in the supporting information for the trend of dairy cows in SoCAB). For natural wetlands, we used the prior emission map from *Jeong et al.* [2013].

Table 2 provides annual CALGEM prior emissions used in this study by source and region, and Figure 1 shows the annual total emission map for the CALGEM prior emission model along with the subregion classification. The regions in this study are different from those in *Jeong et al.* [2013] and follow the California Air Basins (http://www.arb.ca.gov/ei/maps/statemap/abmap.htm). Also, the prior emissions used in this study are 1.3 times higher than those used in *Jeong et al.* [2013] in both the state and Central Valley annual totals (see Figure S2). In section 3, we compare inversion results between *Jeong et al.* [2013] and this study where different prior emissions and inversion methods are used. Inversion results are summarized by region to be compared with the prior emissions. Based on the prior emission estimates, the Central Valley (Regions 3 and 8, Sacramento Valley (SV) and San Joaquin Valley (SJV), respectively) accounts for 55% of the total statewide CH₄ emissions and the two major urban regions (Regions 7 and 12, SFBA and SoCAB) account for 29% of the total. In terms of source sectors, livestock emissions represent 52% of the state total emission followed by landfills (20%) and natural gas (17%; petroleum production included). Livestock emissions are concentrated in Region 8 (San Joaquin Valley), where 86% (667 Gg CH₄/775 Gg CH₄) of the region's total emissions are from livestock. This is consistent with a recent study by *Gentner et al.* [2014] that suggests that the majority of CH₄ emissions in the San Joaquin Valley are from dairy operations.

2.3. Atmospheric Transport Modeling

We used the coupled Weather Research and Forecasting and Stochastic Time-Inverted Lagrangian Transport (WRF-STILT) model for particle trajectory simulations [*Lin et al.*, 2003; *Skamarock et al.*, 2008; *Nehrkorn et al.*, 2010]. The WRF-STILT model has been used to constrain GHG emissions in many studies including airborne measurement-based [e.g., *Gerbig et al.*, 2003; *Kort et al.*, 2008] and tower measurement-based [e.g., *Zhao et al.*, 2009; *Jeong et al.*, 2012a, 2012b, 2013; *Newman et al.*, 2013] applications. We adopt the setup used in *Jeong et al.* [2013] to run the STILT model. In this setup, an ensemble of 500 STILT particles are run backward in time

Source ^a /			-		-											
Region ^b	GBV (6)	LC (5)	LT (15)	MC (4)	MD (10)	NC (2)	NCC (9)	NEP (1)	SoCAB (12)	SCC (11)	SD (14)	SFBA (7)	SJV (8)	SS (13)	SV (3)	Total
DLS	0.1	0.0	0.1	2.5	21.1	21.7	2.8	2.2	37.9	1.0	2.6	14.3	598.1	3.8	30.1	738.3
LF	1.1	1.5	0.0	2.6	8.2	2.3	9.0	1.1	157.0	14.3	26.3	53.9	28.7	3.1	26.4	335.4
NDLS	2.0	0.4	0.1	8.3	3.5	8.2	5.2	13.3	5.8	8.3	1.9	10.5	68.5	1.6	19.8	157.4
NG	0.2	0.4	0.2	2.3	5.5	1.7	4.0	0.4	112.2	17.4	16.2	38.8	51.1	3.1	29.8	283.3 ^c
RM	0.1	0.1	0.0	2.1	2.5	0.4	0.9	0.1	12.0	1.2	2.2	10.0	4.3	0.5	3.3	39.7
WW	0.0	0.1	0.0	0.5	1.0	0.3	1.6	0.4	23.6	13.1	2.8	11.0	9.0	0.9	2.8	67.1
WL	0.3	0.0	0.0	0.5	0.1	0.2	0.2	9.9	0.9	0.5	0.2	4.1	14.0	0.1	7.1	38.1
CP	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	1.5	0.0	47.8	49.2
Total	3.8	2.4	0.5	18.8	41.8	34.8	23.7	27.5	349.3	55.8	52.2	142.5	775.2	13.2	167.0	1708.6

Table 2. Annual CALGEM CH_4 Emissions by Region and Sector (Gg CH_4)

^aSectors include dairy livestock (DLS), landfill (LF), nondairy livestock (NDLS), natural gas including petroleum production and local processing (NG), petroleum refining and mobile sources (RM), wastewater (WW), wetland (WL), and crop (CP; largely rice).

^bThe number in the parentheses shows the region number shown in Figure 1.

^cIncludes 24 Gg CH₄/yr from petroleum seeps (CARB staff private communication).

for 7 days driven with meteorology from the WRF model (version 3.5.1) [*Skamarock et al.*, 2008]. Hourly predicted signals based on WRF-STILT are aggregated into 3 hourly averages for inverse modeling.

The WRF model simulations closely follow those described in *Jeong et al.* [2012a, 2012b, 2013] with some modifications. Here we use version 3.5.1 of the WRF model [*Skamarock et al.*, 2008]. As in *Jeong et al.* [2013], we simulated meteorology for four different horizontal resolutions of 36, 12, 4, and two 1.3 km (vertical levels = 50) using initial and boundary meteorological conditions provided by the North American Regional Reanalysis (NARR) data set [*Mesinger et al.*, 2006]. In this study, the 1.3 km domain for the metropolitan area of Los Angeles was extended to better resolve outflow from the SoCAB region into eastern valleys that



Figure 2. WRF simulation domains and locations of wind profiler sites used for the evaluation of WRF meteorology: CCO (Chico), SAC (Sacramento), LVR (Livermore), CCL (Chowchilla), LHS (Lost Hills), LAX (Los Angeles Airport), ONT (Ontario Airport), and MRV (Moreno Valley). The black dashed lines show the 4 km (d03) and 1.3 km (d04 and d05) domains for WRF simulations.

include the VTR site (see d04 in Figure 2). As in *Jeong et al.* [2013] we applied two-way coupling between domains and 3-D analysis nudging at the outer domain every 3 h using the NARR product.

For surface physics, we use two different land surface models (LSMs) depending on the location of each site as in Jeong et al. [2013]. As pointed out in Jeong et al. [2013], this sitedependent approach to the model configuration is to avoid potential biases in transport modeling. For the Central Valley, we use the five-layer thermal diffusion LSM (5-L LSM) to account for irrigation in the land surface process during summer while using the Noah LSM [Chen and Dudhia, 2001] for other seasons. This is because the Noah LSM overestimates the planetary boundary layer (PBL) in the Central Valley without considering irrigation properly (dry surface leads to overestimation in PBL) [Jeong et al., 2013]. For the urban areas (e.g., SoCAB), we used the Noah LSM following Newman et al. [2013]. Note that as in Jeong et al. [2013] we run WRF multiple times to implement

different model configurations appropriate to individual measurement sites mostly characterized by the air basin each site belongs to.

We also use different PBL schemes depending on the location of the GHG site. As a default for urban areas, we use the Mellor-Yamada-Nakanishi-Niino 2.5 level (MYNN2) PBL scheme [*Nakanishi and Niino*, 2006] coupled with the Noah LSM. This is because we found that the Mellor-Yamada-Janjic (MYJ) scheme [*Mellor and Yamada*, 1982; *Janjić*, 1990] often underestimates nighttime PBL, although it represents daytime PBL well. For the Central Valley region we also use the MYNN2 PBL scheme except for summer, for which we used the MYJ scheme as in *Jeong et al.* [2013] coupled with the 5-L LSM for the Central Valley site. Some sites required improved representation of topographic influences on boundary layer meteorology during winter. Based on the transport evaluation using predicted and measured CO data, we apply the Yonsei University scheme [*Hong et al.*, 2006] with additional parameterization that corrects for surface wind biases at sites with complex topography (e.g., winter season in the southern San Joaquin Valley) [*Jiménez and Dudhia*, 2012]. Based on these configurations for the WRF-STILT model, we computed footprints (sensitivity of concentrations to surface emission fluxes) for each measurement site to be used in the inversion (see equation (2)), and seasonal mean footprints across the measurement sites are provided in Figure S3.

A more complete evaluation of the WRF model simulations and transport errors associated are described in Bagley et al. (submitted to J. Geophy. Res.; henceforth, Bagley et al.), including a comparison of measured and predicted CO for the same period as this study (June 2013 to May 2014). However, we note here that because CO is emitted from sources with different spatial and temporal variations than CH_4 , the results of the CO comparison need to be interpreted with care in ascribing uncertainties to CH_4 emission estimates. In this study, we apply previous results from *Jeong et al.* [2013] to parameterize the prior probability distribution (instead of fixed values) for transport uncertainty and then update the prior uncertainty estimates using the hierarchical Bayesian method as described below.

2.4. Bayesian Inverse Model

We used a hierarchical Bayesian inversion (HBI) [*Ganesan et al.*, 2014] method to estimate regional CH_4 emissions in California. In this work we develop an HBI method with more complex structure in representing the model-measurement mismatch matrix than *Ganesan et al.* [2014] for regional CH_4 emission quantification.

We start with Bayes' rule and describe each probability distribution in the hierarchical structure of parameters that include the scaling factor (a set of factors used to adjust prior emissions, denoted as λ). Generally, Bayes' rule can be applied to multiple parameters at different levels as

$$p(\boldsymbol{\phi}, \boldsymbol{\theta} | \boldsymbol{D}) \propto p(\boldsymbol{D} | \boldsymbol{\phi}, \boldsymbol{\theta}) p(\boldsymbol{\phi}, \boldsymbol{\theta}) = p(\boldsymbol{D} | \boldsymbol{\phi}) p(\boldsymbol{\phi} | \boldsymbol{\theta}) p(\boldsymbol{\theta})$$
(1)

where Φ and θ represent the generic parameters in vector form and **D** is the data used to estimate the parameters. The first line in equation (1) simply states the posterior probability is proportional to the likelihood function and prior distribution for the parameters. The refactorization in equation (1) holds because the data **D** depend only on the parameter ϕ (thus, θ is factored out) and the values of ϕ depend on the values of θ , constructing a hierarchical structure. The transition of $p(\phi, \theta)$ to $p(\phi|\theta)p(\theta)$ is by the property of a conditional probability, given the dependence of ϕ on θ . Any probabilistic model that can be factorized in chains as shown in equation (1) is a hierarchical model [*Kruschke*, 2015].

The general model in equation (1) can be applied to estimate surface emissions and their uncertainties. For GHG applications, the parameter vector ϕ can be scaling factors for emission adjustment (or surface emission itself). The vector θ can be a set of parameters including the hyperparameters (e.g., mean) that determine the distribution for the scaling factor or surface emissions.

We use the following linear model for estimating scaling factors for regional emissions [*Zhao et al.*, 2009; *Jeong et al.*, 2012a, 2012b, 2013; *Wecht et al.*, 2014]

$$\mathbf{y} = \mathbf{K}\boldsymbol{\lambda} + \mathbf{v} \tag{2}$$

where **y** is the measurement vector $(n \times 1)$, which represents 3 hourly local mixing ratio time series after subtracting background values, **K** = **FE** (an $n \times k$ matrix), **F** is the footprint $(n \times m)$, **E** is the prior emissions $(m \times k)$, λ is a $k \times 1$ vector for scaling factors with a covariance matrix **Q** $(k \times k)$, and **v** is a vector representing the model-measurement mismatch with a covariance matrix **R** $(n \times n)$. In this study we solve for a vector of 195 for λ , which includes $0.3^{\circ} \times 0.3^{\circ}$ grid cells (a total of 183) within the major regions (i.e., Regions 3, 7, 8, and 12). We aggregated grid cells from other 12 regions at the subregion scale so that the number of parameters can be reduced for those regions with low prior emissions and weak sensitivity to the measurement sites. Thus, after solving for λ using the HBI method and multiplying it by **E**, we can obtain posterior emissions (a vector of *m*).

For the model in equation (2), the joint parameters we need to estimate are

$$\boldsymbol{\Theta} = \{\boldsymbol{\lambda}, \boldsymbol{\mu}_{\boldsymbol{\lambda}}, \boldsymbol{\sigma}_{\boldsymbol{\lambda}}, \boldsymbol{\sigma}_{\boldsymbol{R}}, \boldsymbol{\eta}, \tau\}$$
(3)

where λ is the scaling factor, μ_{λ} is the prior (i.e., hyperparameter) mean for λ_{i} and σ_{λ} is the uncertainty for λ (i.e., square root of diagonal elements of **Q**). In HBI using a sampling method, λ is sampled from a probability distribution with mean μ_{λ} and standard deviation σ_{λ} , which are also estimated (as part of Θ) instead of being prescribed as in previous work [e.g., Jeong et al., 2013] (see supporting information for details]. $\sigma_{\mathbf{R}}$, η , and τ are the parameters used to construct the model-measurement mismatch matrix R (see below for the representation of **R**). The diagonal elements of **R** represent the total model-measurement mismatch errors that are propagated through the inversion, while Q is used to define the uncertainty level for the prior emission. These two quantities need to be either prescribed with known values or estimated. In HBI we estimate the joint parameter set simultaneously, using the measurements only once [Ganesan et al., 2014]. This joint estimation is different from previous approaches [e.g., Jeong et al., 2013], where the covariance matrix R was prescribed via explicit estimation without using atmospheric measurements. It is also different from other methods where atmospheric measurements were used to optimize \mathbf{R} , and measurements were thereafter also used for inversions [e.g., Michalak et al., 2005]. Depending on the number of measurement sites with data available for each monthly inversion (σ_R is estimated for each of 10–13 sites), the total number of parameters to be estimated ranges from 597 to 600 (195 for each of $\{\lambda, \mu_{\lambda}, \sigma_{\lambda}\}$ and a single estimate for each of $\{\tau, \eta\}$). The number of parameters is usually more than the number of measurements in our pixel-based inversion. We note that it is often sensible to apply hierarchical models for cases where there are more parameters than observed data points because the additional structure provided reduces problems with overfitting [Gelman et al., 2014].

With the parameter set identified, we need to write out the posterior probability up to the likelihood function and prior probability densities. We apply the identified joint parameter (i.e., Θ) to the general formulation of a hierarchical model in equation (1) to express the posterior probability as

$$p(\lambda, \mu_{\lambda}, \sigma_{\lambda}, \sigma_{R}, \eta, \tau | \mathbf{y}) \propto p(\mathbf{y} | \lambda, \sigma_{R}, \eta, \tau) p(\lambda | \mu_{\lambda}, \sigma_{\lambda}) p(\mu_{\lambda}) p(\sigma_{\lambda}) p(\sigma_{R}) p(\eta) p(\tau)$$

$$(4)$$

where the right-hand side shows the likelihood function and the prior distribution for each parameter. Note that in equation (4) all variables are in vector form except for η and τ . To build Markov chain Monte Carlo (MCMC) samplers for the posterior distribution in equation (4), the JAGS system (just another Gibbs sampler) [*Plummer*, 2003] is used together with the R statistical language (https://cran.r-project.org/). JAGS has been widely used for statistical inference studies in many fields including ecology and genetics [*Korner-Nievergelt et al.*, 2015; *McKeigue et al.*, 2010]. To demonstrate the flexibility of our inversion system, we derive a simplified model compared to that in equation (4) and compare inversion results between the two models (see Text S3 in the supporting information for more details). The individual probability distributions (i.e., probability density functions) in equation (4) are described below.

First, for the likelihood function we use

$$p(\mathbf{y}|\boldsymbol{\lambda}, \boldsymbol{\sigma}_{\boldsymbol{R}}, \boldsymbol{\eta}, \tau) \sim \mathcal{N}(\boldsymbol{K}\boldsymbol{\lambda}, \boldsymbol{R})$$
(5)

where *N* is the normal distribution with mean $\mathbf{K}\lambda$ ($n \times 1$) and covariance \mathbf{R} ($n \times n$). Note that \mathbf{y} is conditionally independent of all other parameters given λ , $\sigma_{\mathbf{R}}$, η , and τ .

In order to estimate parameter values with Bayesian inference, prior uncertainty needs to be specified. In the hierarchical model, we need to include prior uncertainty for the joint parameter set Θ using a series of distributions. The scaling factor λ is sampled from a normal distribution (here truncated normal to avoid negative emission fluxes [*Miller et al.*, 2014; *Michalak*, 2008]) instead of a fixed value [e.g., *Jeong et al.*, 2013; *Wecht et al.*, 2014] as

$$p(\lambda) \sim N(\mu_{\lambda}, \sigma_{\lambda})$$
 (6)

where μ_{λ} itself is sampled from a truncated normal distribution [*Miller et al.*, 2014; *Michalak*, 2008] with a mean of 1 and a standard deviation of 0.5 so that 68% of the samples are within 50~150% from the mean,

which is a similar setup to that of *Ganesan et al.* [2014]. Parameter σ_{λ} is modeled using a half-Cauchy distribution, which is one of the recommended distributions for model variances [*Gelman and Hill*, 2007; *Gelman et al.*, 2014; *Korner-Nievergelt et al.*, 2015]. The hyperparameterization ("hyper" meaning the upper level in the hierarchy) for σ_{λ} can formally be expressed as

 σ

$$\mathbf{z} \sim hCauchy(0, 1) \tag{7}$$

where *hCauchy* is the half-Cauchy distribution. Note that we take the absolute value from the Cauchy distribution so that we consider the positive values only (i.e., half-Cauchy). Equation (7) suggests that if we generate random samples (large enough) from equation (7) we get a median value close to 1. Thus, the use of 1 for the half-Cauchy scale parameter (the larger the scale parameter is, the more spread out the distribution is) is similar to assuming that the uncertainty for λ is 100% in the classical Bayesian inversion [e.g., *Zhao et al.*, 2009; *Jeong et al.*, 2013]. The difference is that in this study σ_{λ} is sampled from a distribution with a heavy tail (see Figure S4 for an example of half-Cauchy distribution) so that σ_{λ} can be optimized from a broad distribution (instead of being a fixed value such as 50% of the mean emission).

For the model-measurement mismatch covariance matrix \mathbf{R} , we use an exponential covariance function [*Rasmussen and Williams*, 2006]

$$R_{i,j} = \eta^2 \exp\left(-\frac{1}{\tau} |t_i - t_j|\right) + \delta_{i,j} \sigma_{R_s}^2$$
(8)

where η , τ , and σ_{R_s} are the parameters that define the covariance function; t is the measurement time; and δ is the Kronecker delta function (value of 1 if i = j, otherwise zero). We use two terms in equation (8) to ensure the positive definiteness of **R** [*Stan Development Team*, 2015]. The second term in equation (8) (i.e., with the Kronecker delta function) is analogous to the noise variance in the regression equation [*Rasmussen and Williams*, 2006]. Note that here we use the L_1 norm (i.e., $|t_i - t_j|$) as in *Ganesan et al.* [2014]. The subscript *s* in σ_{R_s} indicates that σ_R is estimated for each site as was done in *Jeong et al.* [2013] for their multitower analysis. This set of multiple parameters for σ_R adds more complexity to the model (than estimating a single value for σ_R) but also reflects the fact that model-measurement errors are not uniform across California.

We model σ_{R_s} using the half-Cauchy distribution as in σ_{λ} [Gelman and Hill, 2007; Gelman et al., 2014; Korner-Nievergelt et al., 2015]. The scale parameter (in the hyperparameter sense) for the half-Cauchy distribution for σ_{R_s} is calculated using the first-order approximation method following Jeong et al. [2012a, 2012b, 2013] and used as

$$p(\sigma_{R_s}) \sim hCauchy(0, \sigma_{Rp|s})$$
 (9)

where $\sigma_{Rp|s}$ is the first-order estimate for σ_{Rs} and includes errors from several sources (e.g., transport and background errors) combined in quadrature. A more detailed discussion of the methodology for the first-order estimation of σ_{Rs} can be found in Text S1 [*Bianco and Wilczak*, 2002; *Bianco et al.*, 2008; *Dye et al.*, 1995; *Gerbig et al.*, 2003; *Göckede et al.*, 2010; *Jeong et al.*, 2012a, 2012b, 2013; *Lin and Gerbig*, 2005; *Newman et al.*, 2013; *Wyngaard and LeMone*, 1980; *Zhao et al.*, 2009]. More details for $\sigma_{Rp|s}$ are also described in the following section.

For η , we use noninformative prior (i.e., uniform distribution) as

$$-unif(0, L) \tag{10}$$

where η is allowed to vary from 0 to *L* with an equal probability of 1/*L*. In this study we use $\sigma_{R\rho|s}$ as an upper limit for *L* because in our choice of the covariance function η is estimated to be smaller than σ_R , and this ensures the positive definiteness of the **R** covariance matrix, which is strictly checked in the JAGS sampler (version 3.4) [*Plummer*, 2003].

 η^{-}

Following Ganesan et al. [2014], we use the exponential distribution for τ as

$$\tau \sim \exp\left(\frac{1}{\tau_p}\right) \tag{11}$$

where τ_p is the hyperparameter for τ , which is assumed to be 7 days (typical synoptic time scale for transport [*Ganesan et al.*, 2014]). The posterior distributions for η and τ are provided in Figure S5 (midmonth for each season), and the complete structure of the **R** covariance matrix is also available in Figure S6. A more detailed

discussion of the methodology for the convergence and accuracy of the estimated parameters including those for the **R** matrix can be found in Text S2 [*Gelman and Rubin*, 1992; *Kass et al.*, 1998; *Plummer*, 2003; *Kruschke*, 2015].

2.5. Uncertainty Matrix

The posterior distribution in equation (4) is used to generate MCMC samples for the parameters (i.e., Θ) in equation (3), which include the components of the error covariance matrices \mathbf{R} and \mathbf{Q} . In other words, we estimate the model-measurement mismatch covariance matrix (i.e., R) and prior emission uncertainty (Q) simultaneously with λ and other parameters (using the measurements just once) instead of using fixed values. In the case of the R matrix, it can be estimated without assuming prior knowledge (e.g., uniform distribution) or from a simple assumption for the hyperparameter as in Ganesan et al. [2014], where for the hyperparameter of the variance component of \mathbf{R} they used the sum of the fixed instrument uncertainty and the uncertainty associated with propagating the calibration scale (0.05 pmol/mol, respectively). Here we take a more informed approach by using site-specific values in constructing \mathbf{R} , drawing on the model-measurement mismatch uncertainties reported in Jeong et al. [2013] for the sites included in that study: Arvin, Madera, Tranquility, WGC, and Sutter Buttes. For other sites, we estimated the modelmeasurement uncertainty for summer of 2013 following the method from Jeong et al. [2012a, 2012b, 2013] (see Text S1 for details). For other seasons, we scaled the summer uncertainty estimates in proportion to the monthly background-subtracted mean mixing ratio signal. We use these uncertainty values (i.e., σ_{Rols}) as the hyperparameter for σ_{Rs} in the covariance function for **R**. As described above, $\sigma_{Rp|s}$ is used as the scale parameter in the half-Cauchy distribution in equation (9) (see Table S1 in the supporting information for $\sigma_{Rp|s}$). As shown in equation (8), the diagonal elements of **R** were then calculated as the sum of squares of σ_{Rs} and η .

Figure 3 shows the (optimized) posterior model-measurement mismatch uncertainty (i.e., diagonal elements of **R**) given the atmospheric measurements for several measurements sites that constrain the major emission regions (SV, SJV, and SoCAB regions; see Figure S7 for the correlation between posterior and prior values across the sites). As described, the HBI approach allows for simultaneous estimation of model-measurement mismatch uncertainty values while inferring posterior emissions, using the measurements only once. This means that the model-measurement mismatch uncertainty has posterior estimates given the prior ($\sigma_{Rp|s}$) and data (i.e., model predictions and measurements). Overall, the posterior values follow the trend of the prior in seasonality and magnitude (Figure 3), although there are some differences between the prior and posterior estimates depending on the site and season. This difference suggests that observations drive the optimization constraining the uncertainty parameters (away from assumed values by our prior knowledge) for some sites and seasons. In most sites, both the prior and posterior uncertainties are large during the winter season when boundary layer heights are low and predicted mixing ratios are very sensitive to the simulated boundary layer [*Jeong et al.*, 2012a, 2013].

In our inverse model, the uncertainty in the prior emissions is expressed in terms of uncertainty in the scaling factors (i.e., σ_{λ} , diagonal terms in **Q**). Here as with $\sigma_{\mathbf{R}}$, the posterior values of σ_{λ} are also sampled from a half-Cauchy distribution with a scale parameter of 1 (equation (7)). For the major emitting regions (3, 7, 8, and 12), the region average of prior uncertainties for individual pixels is estimated to be ~150% (see Figure S8), which is higher than the prescribed 70% in *Jeong et al.* [2013]. It is reasonable to expect this result because the pixel-based inversions have many more degrees of freedom and hence larger per pixel uncertainties than aggregate regions as in *Jeong et al.* [2013].

3. Results

3.1. State Total Emissions

State total emissions were estimated by optimizing 195 scaling factors each month (i.e., dimension of $\lambda = 195 \times 1$) given the multisite measurements and multiplying them by the CALGEM prior emissions, which were essentially the same as the CARB inventory at the subregion scale (see Figure 1 for each subregion). As described, we estimate a scaling factor for each 0.3° pixel within the major emission regions (i.e., SV, SFBA, SJV, and SoCAB), which account for 84% of the CALGEM total emission. For other regions, we estimated a scaling factor for each region. Figure 4 compares predicted and background-subtracted measured mixing ratios using all data (used in the inversion) available for each season and also shows linear regression analysis



Figure 3. Estimated diagonal elements of the model-measurement mismatch matrix **R** for CH₄ inversions. The posterior values were estimated using 25,000 MCMC samples, and the error bar represents the 95% confidence interval. The prior values were estimated using the method described in *Jeong et al.* [2012a, 2012b, 2013] (see Text S1). For May at TRA and September at WGC, the posterior values were not estimated because most of the measurements were not available.

results. Before inversion, the regression analysis estimates best fit slopes to be 0.41-0.75 (predicted versus measured). This simple analysis without full consideration of errors suggests that CH₄ emissions are underestimated by the CARB inventory. After inversion, the best fit slope, root-mean-square error, and coefficient of determination (R^2 ; using the Spearman's method) for each season are significantly improved.



Figure 4. Comparison of predicted and measured CH₄ mixing ratios before (prior) and after (posterior) inversion for each season. The relatively low best fit slopes in the prior comparison (left plot in each season) suggest that prior emissions are underestimated. The filled circles represent individual 3 h data points across different sites used in the inversion. The gray dashed line indicates the 1:1 line, and the black solid line represents the best fit slope for the data shown. The regression coefficients in the posterior plot were calculated based on the median values of the 25,000 MCMC samples. The gray shaded area in the posterior plot represents the 95% uncertainty region for the regression analysis using 25,000 MCMC samples.

The HBI analysis estimates that the state total annual emission is 2.42 ± 0.43 Tg CH₄/yr at 95% confidence not including the (median) posterior estimate for natural wetlands (0.07 Tg CH_{4}/yr). This estimate is equivalent to 1.2-1.7 times the anthropogenic CH₄ emissions in CARB's current official inventory for the year 2013 (1.64 Tg CH₄/yr) [CARB, 2015]. Note that the state total in CARB's current official inventory for 2013 is only slightly different from the prior total in Table 2 after excluding the wetland emission. The state total emission estimate from HBI is consistent with the annual emission estimate from Jeong et al. [2013], 2.38 ± 0.67 Tg CH₄/ yr (at 95% confidence), which combined inverse model estimates for the Central Valley with urban emissions estimated by Wennberg et al. [2012]. As described in section 2, the prior emissions used in this study are 1.3 times higher than those in Jeong et al. [2013]. Also, Jeong et al. [2013] used a simpler inversion method solving for 13 parameters only (i.e., emissions for 13 subregions in California) compared to this study's pixel-based inversion. The good agreement between the two studies for the state annual total emissions indicates that the atmospheric observations drive the inversion system, given the difference in both the prior emissions and the inversion methods. In addition, as described in section 2.2 we performed the inversion using a simplified model compared to the more complex one in equation (4). The simplified model uses prescribed values (i.e., first-order estimates in Table S1) for the model-measurement mismatch uncertainty. The state total emission estimated by the simplified model, described in Text S3 and Figure S9 (see supporting information for the details [National Research Council, 2010; Jeong et al., 2012a, 2013]), differs by only 6% from the HBI estimate, but it has larger uncertainties. Also, we performed a sensitivity test of estimated emissions to well-mixed conditions, showing that there is no significant difference in the emission estimate between



Figure 5. Inferred CH₄ emissions using measurements from 13 sites for four seasons: summer (JJA), fall (SON), winter (DJF), and spring (MAM). The error bar represents the 95% confidence interval around the median value of the posterior emission estimate.

the cases with and without the application of data filtering based on the well-mixed criteria in section 2.2 (see Figure S10 for more information).

As noted in the method section (section 2), transport model error could affect the estimate of CH₄ emissions. Comparison of predicted and measured CO mixing ratios at the four towers during June 2013 to May 2014 (same period as this study) yields near-unity slopes for the majority of sites and seasons (Bagley et al.), suggesting that the WRF-STILT simulations are sufficient to estimate emissions of CO and likely other GHGs across California to within 10% \pm 10% (at 95% confidence) on annual time scales. Based on this result, we add a mean transport bias uncertainty of 10% in quadrature to our Bayesian statistical uncertainty estimates to estimate total uncertainty in annual state total CH₄ emissions from 0.43 to \pm 0.49 Tg CH₄/yr (95% confidence; see Table S2 for the details). If the potential bias in transport modeling that was found for CO is also applicable to CH₄, then the state annual anthropogenic CH₄ emission becomes 10% lower, 2.18 \pm 0.49 Tg CH₄/yr, but still 1.0–1.6 times the CARB inventory (see Table S2 for the application of the CO-based bias). Undiagnosed sources of uncertainty may increase these error bounds beyond that indicated here.

We estimate statewide CH₄ emissions for each season because our measurements are available for a full annual analysis (June 2013 to May 2014). This is the first analysis to estimate full seasonal CH₄ emissions using multitower measurements across California. Although *Jeong et al.* [2013] estimated seasonal CH₄ emissions in California using multitower measurements, they analyzed 10 month data only (not including July and August data) and did not constrain emissions from the Southern California region. Figure 5 shows the estimated mean seasonal emissions for the state, which are the average of the monthly emissions belonging to the season. Note that the prior emissions in Figure 5 only partially account for seasonality because CALGEM has monthly emissions for crop agriculture (largely rice) and wetlands but not other sources. Across seasons, the posterior emissions are greater than the prior emissions without strong evidence for seasonality, similar to previous work by *Jeong et al.* [2013].

3.2. Emissions in Rural and Urban Regions

The hierarchical Bayesian inversion using multiple sites across California constrains CH₄ emissions from a significant portion of both rural and urban regions in California. In particular, the inverse analysis in this study yields a large reduction in the posterior uncertainty for the urban regions of California (e.g., SoCAB) compared to the inverse analysis by *Jeong et al.* [2013], where urban regions were undersampled. We first examine the emissions for the rural regions of California, focusing on the Central Valley because it accounts for ~90% of the total rural emissions based on the CALGEM prior emission.



Figure 6. Estimated annual CH₄ emissions for the major emission regions (at 95% confidence). Regions 3, 7, 8, and 12 represent the Sacramento Valley (SV), San Francisco Bay Area (SFBA), San Joaquin Valley (SJV), and South Coast (SoCAB) air basins, respectively.

Figure 6 shows the comparison between prior and posterior emissions for the major emission regions that account for 84% of the state total in the CALGEM prior emission, including the Central Valley of California (see Table 3 for all regions). We estimate that the Central Valley (Regions 3 and 8) emissions are 1.02-1.74 Tg CH₄/yr (at 95% confimedian = $1.38 \text{ Tg CH}_4/\text{yr}$). dence. These estimates are consistent with the annual emission for the Central Valley estimated by Jeong et al. $[2013], 1.57 \pm 0.20 \text{ Tg CH}_4/\text{yr}$ (95%) confidence). Similarly, Wecht et al. [2014] estimated 1.23 Tg CH₄/yr for the Central Valley using a different transport model, although it was only during the early summer period (May-June 2010). These results sug-

gest that emissions from the Central Valley are underestimated in the CALGEM prior emissions (0.94 Tg CH₄/yr). The spatial distribution of posterior emissions is shown in Figure 7 along with comparison with the CALGEM prior field. As can be seen in the figure, the posterior emissions for some of the pixels in the Central Valley are significantly larger than the prior. However, it should be noted that the uncertainty range for those pixels is also significantly large. This result shows that although the emissions at the subregional scale are well constrained in the Central Valley (aggregated error at 95% confidence is ~25% of the posterior total of the valley), the emission uncertainties for many of the individual pixels are still high. *Bergamaschi et al.* [2005] and *Jeong et al.* [2012a, 2013] reported that posterior emissions between regions. In this study, using pixel-based inversion for major emitting regions we have significantly reduced the anticorrelation in the posterior emissions at the subregional scale (e.g., between Region 3 and Region 7) to 0–20%, compared to those (up to 60% depending on the season) of *Jeong et al.* [2012a] (see Figure S11). This indicates that our total emission for each subregion is much more independent than those of *Jeong et al.* [2012a, 2013].

For urban emissions of California, we focus on emissions from the two major urban regions (SoCAB and SFBA). According to the CALGEM prior, the two urban regions account for 25% of the state total emissions. The HBI analysis estimates that the posterior emissions are 301-490 Gg CH₄/yr (median = 380, 95% confidence) for Region 12 (SoCAB), which are 0.9–1.4 times the prior (349 Gg/yr). This suggests that the prior inventory for SoCAB is consistent with our posterior estimate. Our posterior estimate is also consistent with the results of most of the recent studies that were conducted in SoCAB [Wunch et al., 2009; Hsu et al., 2010; Wennberg et al., 2012; Peischl et al., 2013; Wecht et al., 2014; Wong et al., 2015; Cui et al., 2015]. Figure 8 shows the comparison of estimated CH₄ emissions for SoCAB among eight different recent studies including this study. The estimate (600 Gg CH₄/yr) by Wunch et al. [2009] using the CH₄/CO₂ ratio is likely the upper limit for SoCAB CH₄ emissions and is not included in this comparison. Although the estimated emissions are consistent among the different studies given the reported uncertainty, there are some differences in the mean/median estimates. These differences may arise from different assumptions and undiagnosed uncertainties (e.g., spatial distribution of bottom-up emissions, transport model errors, and different seasonal coverage). For example, most of the studies in SoCAB rely on the combination of measured CH₄ to CO₂ or CO ratios and the bottom-up inventory of CO₂ or CO, with uncertainties that assume that those inventories are relatively well known (e.g., 10% uncertainty assumption in CO₂ inventory by Wong et al. [2015]).

This study constrains CH_4 emissions for SFBA with a significant reduction in the posterior uncertainty, compared to *Jeong et al.* [2012a, 2013]. We estimate the posterior emissions for SFBA to be 159–340 Gg CH_4 /yr (median = 245, at 95% confidence). These emission estimates are consistent with those reported by *Fairley*

Table 3. Posterior A	3. Posterior Annual Emission Estimates (Gg CH ₄ /yr) by Region														
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
Regions ^a	(NEP)	(NC)	(SV)	(MC)	(LC)	(GBV)	(SFBA)	(SJV)	(NCC)	(MD)	(SCC)	(SoCAB)	(SS)	(SD)	(LT)
Prior	28	35	167	19	2	4	143	775	24	42	56	349	13	52	1
HBI	186	144	360	84	20	23	340	1486	180	243	162	490	68	145	2
posterior (upper) ^b															
HBI posterior (lower)	² 1	1	164	1	0	0	159	859	1	1	1	301	1	37	0

^aRegion abbreviations are shown in the parentheses. 97.5th percentile.

^c97.5th percentile.

and Fischer [2015], where they reported a total of 240 ± 60 Gg/yr (at 95% confidence) for the recent period of 2009 to 2012 using CH₄:CO enhancement ratios from 14 air quality sites in SFBA. For SFBA, we have two bottom-up estimates to be compared with our inverse analysis: CALGEM emission model (143 Gg CH₄/yr; see Table 2) and the Bay Area Air Quality Management District (BAAQMD) inventory (126 Gg CH₄/yr [*Bay Area Air Quality Management District*, 2015]). Compared to bottom-up estimates, actual CH₄ emissions in the SFBA are likely 1.1–2.4 and 1.3–2.7 times larger than the CALGEM prior and BAAQMD's inventory, respectively, suggesting that both inventories are lower than our posterior estimate.

3.3. Source Attribution of Emissions

We investigate the likely sources of emissions in the rural and urban regions of California. We estimate CH_4 emissions from different sources assuming the spatial distribution of the CALGEM emission model. Based



Figure 7. Estimated annual CH₄ emissions from the HBI analysis: (a) posterior (median) annual emissions (Gg/yr), (b) ratio of posterior to prior, (c) ratio of estimated 97.5th percentile to prior, and (d) ratio of estimated 2.5th percentile to prior.



Figure 8. Comparison of the CALGEM prior (total for SoCAB = 349 Gg CH₄/yr) and estimated CH₄ emissions for SoCAB in the eight different recent studies including the posterior emission from this study. The value from *Wunch et al.* [2009] shows the CO-based estimate. Originally, *Hsu et al.* [2010] reported LA County emissions (at 200 Gg CH₄/yr), and *Wennberg et al.* [2012] expanded the Hsu et al. results to the full SoCAB. The uncertainty estimates are 68% confidence intervals reported by the individual studies.

on this assumption, we scale individual sector prior emissions at each pixel or region by the inferred scaling factors from the HBI analysis. Figure 9 (left) shows the posterior annual emissions for the HBI analysis by sector. The posterior emissions (804-1410 Gg CH₄/yr, median = 1070 Gg) for the dairy livestock are 1.1–1.9 times larger than the prior emissions. Assuming the distribution of the prior, the posterior estimates for the nondairy livestock (199–345 Gg CH₄/yr, median = 263 Gg) are also 1.3-2.2times larger than the prior. The combined total emissions for dairy and nondairy livestock emissions (1050-1699 Gg CH₄/yr) are 1.2-1.9 times higher than the CALGEM prior. The underestimate in prior livestock emissions agrees with the results described in the region analysis that posterior emissions in the Central Valley (Regions 3 and 8) are larger than

the CALGEM prior. This is also consistent with the reported livestock emissions (1265–1805 Gg CH₄/yr, at 95% confidence) by *Jeong et al.* [2013]. A recent global study suggests a similar underestimation for manure management in a bottom-up inventory. Based on published data on field-scale measurements of GHG emissions, *Owen and Silver* [2015] report that predicted CH₄ emissions by the Intergovernmental Panel on Climate Change Tier 2 method are lower than the mean estimates using the field measurements for most manure management practices. However, we caveat the source attribution above because the spatial distribution of sources by sector may not be perfectly captured in the CALGEM model. In terms of seasonality by sector, Figure 9 (right) suggests that except for wetland (WL) and crop emission (CP), the seasonal variation in the emissions is small, showing similar seasonal posterior emissions within error (Figure 9).



Figure 9. Posterior (left) annual and (right) seasonal emissions (Gg CH₄/yr) estimated from the HBI analysis by sector: dairy livestock (DLS), nondairy livestock (NDLS), landfill (LF), natural gas including petroleum production (NG), petroleum refining and mobile sources (RM), wastewater (WW), crop agriculture (CP; largely rice), and wetland (WL). The error bar represents the 95% confidence interval.

Our inverse analysis also suggests that actual natural gas (NG; includes petroleum production) and landfill (LF) emissions are likely larger than the prior emissions. Our posterior NG emissions (305-502 Gg CH₄/yr) are higher than the prior (283 Gg; see Figure 9) used in this study but consistent with that (331 Gg CH₄/yr) estimated by *Jeong et al.* [2014], where they find that their spatially explicit bottom-up inventory for NG itself is generally lower than those of top-down analyses [e.g., *Peischl et al.*, 2013; *Wennberg et al.*, 2012]. The result for seasonal emissions by sector in Figure 9 (right) shows that the seasonal variation for NG and LF is small, consistent among seasons within error. Other sources, including petroleum refining and mobile (RM), wastewater, crop (rice) emissions (CP), and wetlands (WL) are generally similar between prior and posterior emissions. The rice emissions are 39-101 Gg CH₄/yr (at 95% confidence), which are consistent with those of *Jeong et al.* [2013] (68 ± 18 Gg, at 95% confidence) and *Peischl et al.* [2012] (~85 Gg).

4. Discussion

We further discuss likely source emissions by comparing our estimates with results from previous studies. *Jeong et al.* [2013] estimated annual CH₄ emissions from the livestock source sector in the San Joaquin Valley (Region 8) to be 1.13 ± 0.42 Tg CH₄/yr (at 95% confidence), significantly higher than all other sources combined in the region. This is consistent with the finding by *Gentner et al.* [2014], who concluded that the "vast majority" of the total emissions in San Joaquin Valley are due to dairy operations. In another similar study, *Guha et al.* [2015] used collocated measurements of CO and various volatile organic compounds (VOCs; e.g., alkanes) and a positive matrix factorization technique to estimate the contribution of regional sources to observed enhancements of CH₄. The results in *Guha et al.* [2015] indicate that the livestock emissions account for a majority of the CH₄ (70–90%, uncertainty = 29%) enhancements based on measurements near Bakersfield, CA, during May–June 2010. The reported 29% uncertainty is calculated from the standard deviation in the mass fraction of CH₄ attributed to the dairy source factor profile as estimated from a bootstrapping method. Although these two studies do not report estimated emissions by mass, they suggest that a significant portion of the total CH₄ emission in the San Joaquin Valley (Region 8) is attributed to the livestock sector.

More quantitatively, Jeong et al. [2014] estimated CH₄ emissions from the natural gas sector (petroleum production included) for the state based on activity data and reported emission factors (mostly from U.S. Environmental Protection Agency). They estimated the emission from the natural gas sector to be 128 Gg CH₄/yr for the San Joaquin Valley; the majority of which was from petroleum and natural gas production. After adjusting this bottom-up estimate based on the result in SoCAB by Peischl et al. [2013], they estimated the natural gas emission in San Joaquin Valley to be $162.6 \text{ Gg CH}_4/\text{yr}$, with the San Joaquin Valley accounting for 30% of the state total natural gas emissions. The adjusted natural gas emission (i.e., 162.6 Gg) by Jeong et al. [2014] is 11–19% of the annual total emissions (0.86-1.49 Tg CH₄) in the San Joaquin Valley estimated in this study, which is consistent with Gentner et al. [2014], Guha et al. [2015], and Jeong et al. [2013]. Note that, based on the CALGEM prior, the San Joaquin Valley emits 82% of the total CH₄ emissions in the Central Valley; 86% of which are from the livestock sector. These results suggest that our a priori assumption about the ratio of livestock emissions to the total in the San Joaquin Valley is likely similar to the source attribution of the actual emissions in Region 8. Furthermore, our source analysis indicates that the posterior emissions for landfill, natural gas, and wastewater are generally consistent with or slightly higher than our CALGEM prior, and livestock emissions are higher than the prior, although this is a statewide result (see Figure 9, left). Given this source analysis result, the higher posterior emissions in the San Joaquin Valley (Region 8) from our region analysis (1.1-1.9 times the CALGEM prior) are likely mainly due to livestock sources.

We also examine the emissions in SoCAB for possible source attributions by combining the results from this study and other previous work. *Wennberg et al.* [2012] suggest that the majority of the CH₄ enhancements observed are likely due to natural gas activities, while *Peischl et al.* [2013] estimate 192 ± 54 Gg CH₄ for the combination of emissions from natural gas transmission and distribution plus local seeps and 32 ± 7 Gg CH₄ for oil and gas production and processing. Hence, the total of fossil fuel-related activities from *Peischl et al.* [2013] is 224 ± 55 Gg CH₄, assuming uncorrelated errors in the above estimates. This estimate is larger than our CALGEM prior for the combined total from the natural gas (NG) and refining and on-road mobile (RM) sectors of 124 Gg (see Table 2) by a factor of 1.4–2.3, suggesting an underestimate for total fossil fuel-related emissions in the CALGEM prior for SoCAB. *Lyon et al.* [2015] reported a similar result in a recent

subregional-scale study for the Barnett Shale region, where they estimated higher CH₄ emissions from the oil and gas sector than three inventories by factors of 1.5–4.3. It should be noted that the emission estimates for SoCAB in most of recent studies [e.g., *Wennberg et al.*, 2012] including ours include emissions from petroleum seepage and abandoned wells in the total without distinguishing these as nonanthropogenic emissions. For landfill, wastewater, and livestock sectors, the CALGEM prior estimates 224 Gg CH₄/yr for SoCAB, which is consistent with that (182 ± 54 Gg CH₄/yr) of *Peischl et al.* [2013]. For livestock, *Cui et al.* [2015] estimates emissions in SoCAB to be 52 ± 15 Gg CH₄/yr, which is consistent with the CALGEM prior (44 Gg CH₄/yr). Last, *Cui et al.* [2015] also estimated a combined CH₄ emission of 347 ± 71 Gg CH₄/yr for the landfill and natural gas sectors. This also indicates that natural gas emissions are likely larger than the CALGEM natural gas prior, because their minimum estimate (276 Gg) for the landfill and natural gas sectors is larger than that of the CALGEM prior for natural gas and landfills together (268 Gg). Taken together, these results suggest that while the prior emissions (SoCAB total of 349 Gg) are toward the low end of most of the top-down estimates in SoCAB, underestimation in NG emissions from the CALGEM prior model is possible as indicated by the higher top-down estimates from *Peischl et al.* [2013] and *Cui et al.* [2015].

5. Conclusions

We presented the first analysis of a full annual CH₄ emission budget from California using atmospheric observations from 13 tower sites covering all major CH₄-emitting regions of California. Our analysis suggests that California's actual emissions are likely 1.2–1.8 times larger than a recent state inventory [*CARB*, 2015]. Our measurement network across California constrains CH₄ emissions from California's urban and rural emissions, and the added measurement sites to the CH₄ network significantly reduced the posterior uncertainty estimates. In particular, this study constrains the San Francisco Bay Area's CH₄ emissions for the first time using a Bayesian inverse analysis. This study also shows that the inverse framework based on the measurement network can be an effective approach to quantifying emissions at the regional scale and monitoring long-term spatial and temporal changes in emissions. Although the CO comparison (Bagley et al.) appears largely consistent with expectation, it should be noted that undiagnosed sources of errors or uncaptured errors in the model-measurement mismatch covariance may increase these uncertainty bounds beyond that indicated here.

In the future, a combination of improved prior emissions and meteorological models, expanded multigas measurements, and inverse model analyses will reduce uncertainty in California's GHG emissions. Also, more efforts are needed to constrain emissions by both sector and region. For example, while our results and other studies indicate both livestock and natural gas emissions appear to be underestimated, attribution of the magnitude of errors to specific sectors is difficult. A recent study on CH₄ emissions from the Aliso Canyon blowout in Los Angeles emphasizes the utility of tracers (e.g., ethane) for source speciation [*Conley et al.*, 2016]. Using both methane and ethane measurements, *Conley et al.* [2016] reported that at its peak the Aliso Canyon event doubled SoCAB emissions during the 3 month period, producing a total of 97 Gg CH₄, which is 28% of the SoCAB total CH₄ emission (349 Gg/yr) from our CALGEM prior model. Given the importance of distinguishing the regional variations in dominant CH₄ sources (e.g., Central Valley versus SoCAB) and large-scale events such as the Aliso Canyon blowout, a combination of facility specific emission measurements and regionally representative measurements of source-specific tracers (e.g., CO, VOCs, and potentially CH₄ isotopes) [*Townsend-Small et al.*, 2012; *Peischl et al.*, 2013; *Guha et al.*, 2015] are likely to prove useful in the future.

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