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Evaluation of improved land use and canopy representation in BEIS v3.61 with biogenic VOC measurements in California

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Abstract

Biogenic volatile organic compounds (BVOC) participate in reactions that can lead to secondarily formed ozone and particulate matter (PM) impacting air quality and climate. BVOC emissions are important inputs to chemical transport models applied on local to

- ⁵ global scales but considerable uncertainty remains in the representation of canopy parameterizations and emission algorithms from different vegetation species. The Biogenic Emission Inventory System (BEIS) has been used to support both scientific and regulatory model assessments for ozone and PM. Here we describe a new version of BEIS which includes updated input vegetation data and canopy model formulation
- for estimating leaf temperature and vegetation data on estimated BVOC. The Biogenic Emission Landuse Database (BELD) was revised to incorporate land use data from the Moderate Resolution Imaging Spectroradiometer (MODIS) land product and 2006 National Land Cover Database (NLCD) land coverage. Vegetation species data is based on the US Forest Service (USFS) Forest Inventory and Analysis (FIA) version 5.1 for
- ¹⁵ years from 2002 to 2013 and US Department of Agriculture (USDA) 2007 census of agriculture data. This update results in generally higher BVOC emissions throughout California compared with the previous version of BEIS. Baseline and updated BVOC emissions estimates are used in Community Multiscale Air Quality Model (CMAQ) simulations with 4 km grid resolution and evaluated with measurements of isoprene and 20 monoterpenes taken during multiple field campaigns in northern California. The up-
- ²⁰ monoterpenes taken during multiple field campaigns in northern California. The updated canopy model coupled with improved land use and vegetation representation resulted in better agreement between CMAQ isoprene and monoterpene estimates compared with these observations.

1 Introduction

²⁵ Volatile organic compounds (VOC) are known to contribute to ozone (O_3) and particulate matter less than 2.5 microns in diameter $(PM_{2.5})$ formation in the troposphere.





Elevated concentrations of O_3 and $PM_{2.5}$ have known deleterious health effects (Bell et al., 2004; Pope and Dockery, 2006; Pope et al., 2006) and climate implications. Biogenic VOC (BVOC) are highly reactive and contribute to local and continental scale O_3 and $PM_{2.5}$ (Carlton et al., 2009; Chameides et al., 1988; Wiedinmyer et al., 2005).

- ⁵ Terrestrial biogenic emissions are an important input to photochemical transport models which are used to quantify the air quality benefits and climate impact of emission control plans. Despite the important role of BVOC in atmospheric chemistry, the spatial representation of vegetation species, their emission factors, and canopy parameterization remain highly uncertain.
- ¹⁰ Isoprene, a highly reactive BVOC, contributes to O_3 (Chameides et al., 1988) and influence secondary organic aerosol (SOA) formation (Carlton et al., 2009). Monoterpenes and sesquiterpenes are BVOCs known to react in the atmosphere to form SOA (Sakulyanontvittaya et al., 2008). The impact of BVOC emissions on these pollutants is significant enough that model simulations have been conducted to explicitly quantify
- their impact (Fann et al., 2013; Kwok et al., 2013; Lefohn et al., 2014). The Biogenic Emission Inventory System (BEIS) (Pierce and Waldruff, 1991; Schwede et al., 2005) estimates these and other BVOC species and has been used extensively to support scientific (Carlton and Baker, 2011; Fann et al., 2013; Kelly et al., 2014; Simon et al., 2013; Wiedinmyer et al., 2005) and regulatory (US Environmental Protection Agency, 2010, 2011, 2012b, a) model applications.

BVOC emissions are highly variable among different types of vegetation, therefore the representation of vegetative coverage is critically important for accurate spatial distribution of emissions. Northern California has a large gradient in high isoprene emitting vegetation extending from the Sacramento valley eastward toward the Sierra Nevada

(Dreyfus et al., 2002; Karl et al., 2013; Misztal et al., 2014). Many counties in California have been designated as non-attainment of both the 8 h O₃ and PM_{2.5} National Ambient Air Quality Standards (NAAQS). Recent field studies measuring BVOC concentrations in this area provide a unique opportunity to evaluate photochemical model estimated BVOC ambient concentrations using an existing (BEIS version 3.14) and up-





dated version of BEIS (version 3.61) and input vegetation data. Ground measurements of BVOC concentrations were made during the Carbonaceous Aerosols and Radiative Effects Study (CARES) campaign in an urban area (Sacramento) and at a site downwind from Sacramento (Cool, CA) that is located near vegetation known for high isoprene emissions (Zaveri et al., 2012). The Biosphere Effects on Aerosols and Photochemistry Experiment (BEARPEX) 2009 campaign provides BVOC measurements at a remote location in the Sierra Nevada foothills to the east of Sacramento and Cool

(Beaver et al., 2012), an area of high monoterpene emitting vegetation.

In this manuscript, BVOC emissions estimated with the existing, version 3.14 (Schwede et al., 2005), and updated version of BIES, version 3.61, are input to the Community Multiscale Air Quality (CMAQ) photochemical transport model (Hutzell et al., 2012; Byun and Schere, 2006; Foley et al., 2010) and estimated BVOC ambient concentrations are compared to surface observations at these field campaigns in central and northern California. Canopy coverage and vegetation species data has been

- ¹⁵ updated with the FIA 5.1 and 2006 NLCD data sets using more spatially explicit techniques for tree species allocation. BEIS 3.61 has been updated with new a canopy model of leaf temperature for emissions estimation. Canopy leaf temperature estimates are also compared with infrared skin temperature measurements over a grass canopy made at Duke Forest. BVOC estimates from the Model of Emissions of Gases and Acreacia from Nature (MECAN). (Querther et al. 2010) are also input to CMAO and
- Aerosols from Nature (MEGAN) (Guenther et al., 2012) are also input to CMAQ and model predictions are compared with field study measurements to provide additional context for BEIS updates.

2 Methods

2.1 Land cover and vegetation speciation

²⁵ BEIS 3.14 used the BELD 3 landuse dataset relied on combined US county level USDA-USFS Forest Inventory and Analysis (FIA) vegetation speciation circa 1992





information with the 1992 USGS landcover information (Kinnee et al., 1997). A new land cover dataset (BELD 4) integrating multiple data sources has been generated at 1 km resolution covering North America. Landuse categories are based on the 2001 to 2011 National Land Cover Dataset (NLCD), 2002 and 2007 USDA census of agriculture county level cropping data, and Moderate Resolution Imaging Spectroradiometer

 fure county level cropping data, and Moderate Resolution Imaging Spectroradiomete (MODIS) satellite products where more detailed data was unavailable.

Fractional tree canopy coverage is based on the 30 m resolution 2001 NLCD canopy coverage (http://nationalmap.gov/landcover.html: Homer et al., 2004) and land cover is based on 30 m resolution 2006 NLCD Land Cover data. The 2001 canopy data was used because there was no canopy product developed for the 2006 NLCD. Land cover

¹⁰ used because there was no canopy product developed for the 2006 NLCD. Land cover for areas outside the conterminous United States is based on 500 m MODIS land cover data for 2006 (https://lpdaac.usgs.gov/products/modis_products_table; MCD12Q1) using the International Geosphere Biosphere Programme classification.

Vegetation speciation is based on multiple data sources. Tree species are based on 2002 to 2013 Forest Inventory and Analysis (FIA) version 5.1 and crop species information is based on 2002 and 2007 USDA census of agriculture data. The FIA includes approximately 250 000 representative plots of species fraction data that are within approximately 75 km of one another in areas identified as forest by the NLCD tree canopy coverage. USDA census of agriculture data is available on a State and 20 County level only and has been used to refine the agricultural classes to the NLCD

agricultural land use categories.

FIA version 5.1 location data has been degraded to enhance landowner privacy in accordance with the Food Security Act of 1985 (O'Connell et al., 2012). The provided locations are accurate within approximately 1.6 km with most plots being within 0.8 km

of the reported coordinates and have accurate State and County identification codes (O'Connell et al., 2012). BELD 3 FIA vegetation specie fractions were aggregated to county level based on national above ground biomass estimates for deciduous, pine, juniper, fir, and hemlock species. In the BELD 4 data set, FIA plot level forest biomass (kg ha⁻¹) and specific leaf area (g m⁻²) were estimated using the allometric scaling





methods of (Jenkins et al., 2003) and (Chojnacky et al., 2014). Plot level tree biomass estimates were corrected for sampled bole biomass and scaled to a per hectare bases following (O'Connell et al., 2012). The plot level total and foliage biomass estimates are then extrapolated to the continental United States by spatial kriging using the plots

- longitude, latitude and elevation as predictors and weighted by the NLCD canopy fraction. If elevation was not reported at the plot then elevation was supplied by a digital elevation model from WRF. Kriging was done in 140 by 140 km windows with a 50% overlap to address regional differences in spatial gradients. A buffer that extended beyond this window was determined by a semivariogram. Similarly, tree species biomass
 information was kriged with the additional constraint of the NLCD land use categories
 - (deciduous, evergreen or mixed forest) applied as weights.

The fractional species composition of the NLCD canopy coverage was then calculated and the FIA 5.1 species were aggregated to the BELD 4 species (Table S1 and Fig. S1 in the Supplement). The NLCD land cover defines trees as greater than 5 m tall,

- forest refers to greater than 20 % canopy coverage, with deciduous forests have more than 75 % foliage shed in winter and evergreen forests have more than 75 % of foliage retained in winter (http://www.mrlc.gov/nlcd06_leg.php). These tolerances were used constraining the kriging processes. Total kriged biomass estimates were quantitatively evaluated against the independent estimates of (Blackard et al., 2008). Species spe-
- cific data in BELD 4 were qualitatively evaluated against the range maps of Critchfield and Little (1966) and Little Jr. (1971, 1976). This kriging approach provides an estimate of vegetation speciation for land cover categories where information is not readily available such as urban, grassland, and shrublands. While this kriging approach may provide better spatial estimates of biomass and vegetation type for most areas of the
- ²⁵ continental United States, it is possible that small areas with vegetation and biomass dramatically different than the surrounding region (e.g. some urban areas) will likely need further refinement.





2.2 Biogenic emissions

MEGAN and BEIS are both used to support regional to continental scale O_3 and $PM_{2.5}$ photochemical model applications (Carlton and Baker, 2011). Both modeling systems estimate emissions based on vegetation type, meteorological variables, and canopy

⁵ characteristics (Carlton and Baker, 2011). MEGAN and BEIS have similar governing equations but differ in vegetation characterization, emission factors, meteorological adjustments, and canopy treatment. These models have been evaluated against BVOC measurements in the central United States (Carlton and Baker, 2011) and Texas (Warneke et al., 2010) but little evaluation of both models has been done for California. BEIS version 3.14 provides a baseline for comparison of BEIS version 3.61 that includes enhancements described here.

BEIS version 3.61 estimates emissions for 33 volatile organic compounds, carbon monoxide, and nitric oxide. Table 1 shows the complete list of compounds estimated by BEIS with mapping to contemporary gas phase chemical mechanisms SAPRC07T

- and CB6. BEIS estimates isoprene, 14 unique monoterpene compounds, and total sesquiterpenes. In addition, emissions are estimated for 16 other volatile organic compounds and an aggregate group of other unspeciated VOC. All biogenic VOC emissions are a function of leaf temperature while only isoprene, methanol, and MBO are a function of both leaf temperature and photosynthetically activated radiation (PAR). All
 species emissions have small indirect impacts from PAR via the canopy module.
- Inputs to BEIS include normalized emissions for each vegetation species, gridded vegetation species, temperature, and PAR. Temperature and PAR can be provided from prognostic meteorological models such as WRF or other sources such as satellite products (Pinker and Laszlo, 1992; Pinker et al., 2002) or ambient measurements.
- The BELD 4 database contains vegetation specie information for 275 different vegetation categories (Table S1). Table 2 shows emission rates for each emitted compound by aggregated vegetation type to illustrate variability in emissions. These vegetation types include 20 MODIS and 21 NLCD land cover categories, and 20 different types





of crops both irrigated and non-irrigated (40 total). The remaining categories include tree species, much of which are broadleaf (e.g. oak) and needle leaf (e.g. fir) species. A gridded file indicating leaf-on based on the 2009 modeled meteorology, bioseasons file, is also provided as input to BEIS. In the future leaf out and leaf fall dates will be metabed with LAL data. However, it is unlikely the current simple leaf on parameteric

⁵ matched with LAI data. However, it is unlikely the current simple leaf-on parameterization will impact typical regulatory assessments since elevated O_3 and $PM_{2.5}$ organic carbon events often happen outside the spring and fall seasons.

For various sensitivity studies presented here, BEIS 3.14 is applied with BELD 3 vegetation data, WRF temperature, and both WRF and satellite derived estimates of

- PAR. BEIS 3.61 is applied similarly but with BELD 3 and BELD 4 vegetation data to isolate the impact of the updates to the canopy model. A gridded 0.5 by 0.5° resolution satellite estimate of PAR from 2009 was processed to match the model domain specifications and input to both BEIS and MEGAN. The satellite estimates are based on the GEWEX Continental Scale International Project and GEWEX Americas Pre-
- ¹⁵ diction Project Surface Radiation Budget (www.atmos.umd.edu/~srb/gcip/gcipsrb.htm) (Pinker et al., 2002). MEGAN version 2.1 (Guenther et al., 2014, 2012) with version 2011 North America Leaf Area Index and Plant Functional Type (Guenther et al., 2014) was applied with WRF estimated temperature and PAR and also with satellite derived PAR.

20 2.3 Canopy Model – Leaf temperature update

BEIS 3.61 includes a two layer canopy model. Layer structure varies with light intensity and solar zenith angle. Both layers of the canopy model include estimates of sunlit and shaded leaf area based on solar zenith angle and light intensity, direct and diffuse solar radiation, and leaf temperature. BEIS 3.14 previously used 2 m temperature to
 ²⁵ represent canopy temperature for emissions estimation even though BVOC emission factors are typically based on leaf temperature (Niinemets et al., 2010). The canopy model has been updated to use land surface physics from the Weather and Research Forecasting model and air-surface exchange algorithms from the CMAQ model to ap-





proximate leaf temperature using an energy balance for the sunlit and shaded portion of each canopy layer. Emissions are estimated for sunlit and shaded fractions of the canopy and summed over the two layers for total canopy emissions.

A simple two big leaf (sun and shade) temperature model was developed based on 5 a radiation balance. The leaf radiation balance is solved for both the sun (Eq. 1) and shaded (Eq. 2) leaf sides in each layer.

sun leaf

$$R_{abs} + IR_{in} - IR_{out} - H - \lambda E + G = 0$$

shade leaf

П

25

 $R_{\text{shade}} + IR_{\text{in}} - IR_{\text{out}} - H - \lambda E + G = 0,$

where IR_{in} is the incoming infrared radiation, IR_{out} is the outgoing infrared radiation, λ is the latent heat of evaporation, E_{sun} and E_{shade} are the latent heat flux from sun and shade leaves respectively, H is the sensible heat flux, and G is the soil heat flux. To maintain the same energy balance as WRF it was assumed that E scales linearly with sunlit and shaded fractions of the canopy. Note, that conventionally G is positive when 15 the soil is being heated and negative when the soil is cooling while the sign convention of the other variables are relevant to heating and cooling of the atmosphere. $R_{\rm abs}$ is the total incoming solar radiation from the meteorological model and R_{shade} is modeled using the attenuation, scattering and diffuse radiation from Weiss and Norman (1985). The infrared budget is parameterized as 20

$$IR_{in} = \varepsilon_{atm} \sigma T_{atm}^4$$

$$IR_{out} = \varepsilon_{leaf} \sigma T_{leaf}^4,$$

where $arepsilon_{\rm atm}$ and $arepsilon_{\rm leaf}$ are the emissivities of the atmosphere and leaf respectively, σ is the Stephan Bolzman constant and T_{atm} and T_{leaf} are the atmospheric and leaf temperatures respectively.



(1)

(2)

(3)

(4)



E is parameterized as

$$E = \frac{e_{\rm s}(T_{\rm leaf}) - e_{\rm a}}{R_{\rm w, \rm leaf}P_{\rm atm}},$$

15

where $e_s(T_{\text{leaf}})$ is the saturation vapor pressure at the leaf, e_a is the atmospheric vapor pressure, $R_{w,\text{leaf}}$ is the resistance to water vapor transport from the leaf to atmosphere and P_{atm} is the atmospheric pressure at the surface.

The saturation vapor pressure of the leaf is defined as

 $e_{\rm s}(T_{\rm leaf}) = ae^{\frac{b(T_{\rm leaf}-273.15)}{T_{\rm leaf}-c}},$

where the empirical coefficients are a = 611.0 Pa, b = 17.67, and c = 29.65 °C.

H is parameterized following the WRF Pleim-Xiu (PX) land surface model (Skamarock et al., 2008) as

$$H = \frac{\rho_{\rm atm} C_{\rho} \left(\frac{P_0}{P_{\rm atm}}\right)^{R_{\rm atm}/C_{\rho}} (T_{\rm leaf} - T_{\rm air})}{R_{\rm h, leaf}},$$

where ρ_{atm} is the atmospheric density, C_p is the specific heat of air, P_0 is the STP pressure, R_{atm} is the gas constant for dry air, and $R_{\text{h,leaf}}$ is the resistance to heat advection between the atmosphere and leaf.

The T_{leaf}^4 variable and Eq. (6) prevents an analytical solution. Thus the approximation from Campbell and Norman (1998) is used.

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The T_{leaf}^4 term is simplified as follows:

$$\varepsilon_{\text{leaf}} \sigma T_{\text{leaf}}^4 \approx \varepsilon \sigma T_{\text{atm}}^4 + \frac{\rho_{\text{atm}} C_{\rho} \left(\frac{P_0}{P_{\text{atm}}}\right)^{R_{\text{atm}}/C_{\rho}} (T_{\text{leaf}} - T_{\text{air}})}{R_{\text{r,leaf}}},$$

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(6)

(7)

(8)

where $R_{r,leaf}$ is the atmospheric radiative resistance ~ 230 s m⁻¹ (Monteith and Unsworth, 2013).

Equation (6) is then further simplified:

$$\lambda \rho_{\rm atm} \frac{e_{\rm s}(T_{\rm leaf}) - e_a}{R_{\rm w,leaf} P_{\rm atm}} \approx \lambda S(T_{\rm atm}) \frac{\left[T_{\rm leaf} - T_{\rm atm}\right]}{R_{\rm w,leaf}} + \lambda \rho_{\rm atm} \frac{e_{\rm s}(T_{\rm atm}) - e_a}{P_{\rm atm} R_{\rm w,leaf}},\tag{9}$$

5 where

$$S = \frac{de_{\rm s}(T)}{dT}.$$
(10)

Equations (1), (3), (5), (7), (8), and (9) are algebraically combined to estimate the sunlit leaf temperature assuming that $\varepsilon_{\text{atm}} = \varepsilon_{\text{leaf}}$.

$$T_{\text{sun,leaf}} \approx T_{\text{atm}} + \frac{R_{\text{sun}} + G - \lambda \rho_{\text{atm}} \frac{e_{\text{s}}(T_{\text{atm}}) - e_{a}}{P_{\text{atm}}R_{\text{w,leaf}}}}{\rho_{\text{atm}} \left[\left(\frac{P_{0}}{P_{\text{atm}}}\right)^{R/C_{p}} C_{p} \left(\frac{1}{R_{\text{h,leaf}}} + \frac{1}{R_{\text{r,leaf}}}\right) + \lambda S \left(\frac{1}{R_{\text{w,leaf}}}\right) \right]}.$$
(11)

Equations (2), (3), (5), (7), (8), and (9) are combined to estimate the shaded leaf temperature:

$$T_{\text{shade,leaf}} \approx T_{\text{atm}} + \frac{R_{\text{shade}} + G - \lambda \rho_{\text{atm}} \frac{e_{\text{s}}(T_{\text{atm}}) - e_{a}}{P_{\text{atm}}R_{\text{w,leaf}}}}{\rho_{\text{atm}} \left[\left(\frac{P_{0}}{P_{\text{atm}}} \right)^{R/C_{p}} C_{p} \left(\frac{1}{R_{\text{h,leaf}}} + \frac{1}{R_{\text{r,leaf}}} \right) + \lambda S \left(\frac{1}{R_{\text{w,leaf}}} \right) \right]}.$$
(12)

The sunlit leaf area index, ${\rm LAI}_{\rm Sun},$ is estimated following (Campbell and Norman, 1998)

¹⁵
$$LAI_{Sun} = \int_{0}^{LAI} e^{-k_{be}(\Psi)L} dL,$$
 (13)



where LAI is the total canopy leaf area index, k_{be} is the extinction coefficient for direct beam incoming solar radiation as a function of the solar zenith angle, Ψ following Campbell and Norman (1998). The shaded leaf area index, LAI_{Shade}, is then estimated as follows:

 $5 \text{ LAI}_{\text{Shade}} = \text{LAI} - \text{LAI}_{\text{Sun}}.$

BVOC emission fluxes, F_i , are estimated similar to Guenther et al. (2006) for sunlit and shaded fractions of the canopy

 $F_{i,j} = E_i \gamma_{\mathsf{PAR},i,j} \gamma_{T,i,j} \mathsf{LAI}_j,$

Ι ΔΙ-

where E_i is the emission factor or BVOC species *i*, γ_{PAR} is the emission activity factor for PAR (currently only applied to isoprene, methanol and MBO), γ_T is the emission activity factor for leaf temperature following Guenther et al. (1993), and *j* is the index for sunlit or shaded leaves. γ_{PAR} integrates the PAR emissions activity factor of Guenther et al. (1993) for sunlit and shaded layers following Niinemets et al. (2010).

$$\gamma_{\text{PAR},i,\text{Sunlit}} = \text{PAR}C_{L} \int_{0}^{L \cap \text{Sun}} \frac{e^{-2k_{\text{dd}}L}}{\sqrt{1 + \alpha^2 \text{PAR}^2 e^{-2k_{\text{dd}}L}}} dL$$
(16)

- 1	-
	10

10

$$\gamma_{\text{PAR},i,\text{Shaded}} = \text{PAR} C_L \int_{\text{LAI}_{\text{Sun}}}^{\text{LAI}} \frac{e^{-2k_{\text{dd}}L}}{\sqrt{1 + \alpha^2 \text{PAR}^2 e^{-2k_{\text{dd}}L}}} dL, \qquad (17)$$

where k_{dd} is the net attenuation coefficient for direct and diffuse PAR and α and C_L are empirical coefficient, 0.0027 and 1.066 respectively, defined in Guenther et al. (1993).



(14)

(15)



2.4 Photochemical model background, inputs, and application

Chemical species are estimated using the Community Multiscale Air-Quality Model (CMAQ) version 5.0.2 (www.cmaq-model.org) photochemical grid model. CMAQ was applied with SAPRC07TB gas phase chemistry (Hutzell et al., 2012), ISORROPIA II inorganic chemistry (Fountoukis and Nenes, 2007), secondary organic aerosol treatment (Carlton et al., 2010) and aqueous phase chemistry that oxidizes sulfur, glyoxal, and methyglyoxal (Carlton et al., 2008; Sarwar et al., 2013). The Weather Research and Forecasting (WRF) Advanced Research WRF core (ARW) version 3.3 (Skamarock et al., 2008) was used to generate gridded meteorological inputs for CMAQ and emissions models. While not coincident with this study, this WRF configuration compared well with mixing layer height and surface measurements of temperature and winds in central California during the summer of 2010 (Baker et al., 2013). For model performance evaluation presented here, model estimates are paired with measurements using the grid cell where the measurement was located. Measurements are paired in time with hourly model estimates with the closest model hour (Simon et al., 2012).

The model domain covers central and northern California with 4 km square sized grid cells. The surface to 50 mb is resolved with 34 layers. Layers nearest the surface are most finely resolved with an approximate height of 38 m for layer 1. The modeling period extends from 3 June through 31 July 2009 to be coincident with the BEARPEX field campaign and minimize the influence of initial conditions on model estimates. Initial conditions and boundary inflow are from a coarser CMAQ simulation covering the continental United States. Inflow to the coarser simulation is from a global 2009 application of the GEOS-CHEM (v8-03-02) model (http://acmg.seas.harvard.edu/geos/)

Stationary point sources are based on 2009 specific emissions where available and the 2008 National Emission Inventory (NEI) version 2 otherwise. Mobile emissions are interpolated between 2007 and 2011 estimates provided by the California Air Resources Board (CARB) and allocated spatially and temporally using the Spare Ma-

(Henderson et al., 2014).





trix Operator Kernel Emissions (SMOKE) model (http://www.cmascenter.org/smoke). Other non-point and commercial marine emissions are based on the 2008 NEI version 2 (http://www.epa.gov/ttn/chief/net/2008inventory.html).

2.5 Field study measurements

- ⁵ Between 15 June and 31 July 2009, the BEARPEX study was conducted to study photochemical reactions and products in areas downwind of urban areas with large biogenic influences. The study was located at a managed ponderosa pine plantation in the foothills of the Sierra Nevada (38.90° N, 120.63° W), located near the University of California's Blodgett Research Forest Station. The measurement site was near
- Georgetown, CA, approximately 75 km from Sacramento, CA. Two research towers housed meteorology and atmospheric composition measurements and inlets during BEARPEX 2009. Meteorological measurements were made on the south, 12.5 m tower, including photosynthetically active radiation (PAR) measured by a LI-COR LI190. The second tower (17.8 m) was located approximately 10 m north of the meteorological
- tower and housed most of the atmospheric composition measurements. The inlet used to sample BVOCs was located at the top of the north tower, approximately 9 m above the ponderosa pine canopy level. BVOCs including isoprene, monoterpenes, methyl vinyl ketone, and methacrolein were quantified using an online gas chromatograph with a flame ionization detector (GC-FID) (Park et al., 2010, 2011). BVOC samples
 were collected during the first 30 min of every hour, then subsequently analyzed with the GC-FID.

During June 2010, the CARES study was conducted to study the formation of organic aerosols and the subsequent impacts on climate. The study was composed of two surface monitoring sites: T0 and T1. The T0 was located in Sacramento, CA at the

²⁵ American River College campus (38.65° N, 121.35° W), and the T1 site was in Cool, CA on the campus of Northside School (38.87° N, 121.02° W). The T0 site was approximately 14 km northeast of downtown Sacramento, and the T1 site was surrounded by the forested foothills of the Sierra Nevada. Isoprene and monoterpene measurements





at the Sacramento (TO) and Cool (T1) CARES ground sites were made with GC-MS and PTRMS, respectively (Zaveri et al., 2012), and sampled via inlets at approximately 10 m above the surface. PTRMS data were reported as 1 s measurements approximately every 30 s. GC-MS data were 10 min collections every 30 min. All observation data was averaged to hourly concentrations before comparison with model estimates.

The sunlight leaf temperature in MEGAN 2.1 and the revised canopy model in BEIS 3.61 were evaluated against observations taken in 2008 at the Blackwood Division of the Duke Forest in Orange County, North Carolina, USA (35.97° N, 79.09° W). Details regarding the site (FLUXNET, 2014), measurements, and species composition are available elsewhere (Almand-Hunter et al., 2015). Leaf temperature measurements were taken using an infrared temperature sensor (IRTS-P, Apogee Instruments Inc, Logan, UT) mounted on the grassland tower.

3 Results

3.1 Leaf temperature algorithms compared to observations

- ¹⁵ The canopy model updates for leaf temperature estimation are evaluated by comparing canopy model output with infrared skin temperature measurements of a grass canopy at the Duke Forest field site in central North Carolina (Fig. 1). BEIS 3.61 canopy model inputs are based on field measurements taken at this location coincident with the skin temperature data collection. The infrared skin temperature measurements do not rep-
- 20 resent a mean canopy leaf temperature but rather the temperature of the portion of the canopy exposed to the atmosphere. The infrared skin temperature measurement should be warmer than the mean leaf temperature during periods of solar irradiation and cooler during periods of radiative cooling due to the insulating effect of the unexposed portion of the canopy. Only the estimated exposed leaf temperature (Eq. 12)
- was used in the evaluation to account for this discrepancy between measurements and canopy model output. Figure 1 shows observed and predicted estimates of leaf temper-





ature and difference between leaf and ambient temperature. The average temperature estimated by the BEIS 3.61 canopy model for the top of the canopy compares well with observations (mean bias of 0.3 K and mean error 1.2 K). Top of the canopy leaf temperature estimated by MEGAN 2.1 are comparable to BEIS 3.61 and the observations at the Duke Forest site.

3.2 Evaluation of the BELD 4 land use data

BELD 4 total forest biomass estimates were evaluated against the independent estimates of Blackard et al. (2008). Figure 2 shows the BELD 4 and (Blackard et al., 2008) estimates of forest biomass for this model domain at 4 km resolution. The (Blackard et al., 2008) 250 m grid resolution data set was projected and aggregated to the CMAQ 4 km grid resolution projection using rgdal and raster libraries in R (Bivand et al., 2014). The BELD 4 estimates evaluated well against those of (Blackard et al., 2008) with a Pearson's correlation coefficient of 0.872 (p < 0.001) and a mean and median difference in tree biomass in areas where the NLCD data indicated canopy coverage was -13 kg ha^{-1} (-32 %) and $-0.004 \text{ kg ha}^{-1}$ (0 %) respectively. BELD 4 estimates of forest biomass were greater than those of Blackard et al. (2008) in the densely forested areas in the high Sierras and lower in the lower elevation areas of the domain, primarily in the basin and range areas in the Sacramento valley. The prevalence of the lower elevation

- areas with lower biomass estimates drives the difference between the forest biomass estimates. The biomass estimates of Blackard et al. (2008) under predicted the full range of the biomass variability with over predictions in areas with low biomass and under predictions in areas of high biomass compared to the FIA tree survey biomass observations. The total biomass estimates presented here have a larger range, 0– 661 kg ha⁻¹ versus 0–499 kg ha⁻¹ with a median absolute deviation of 2.9 kg ha⁻¹ versus 2.5 kg ha⁻¹ for areas with NLCD canopy coverage. The lower biomass estimates
- ²⁵ sus 2.5 kg ha⁻¹ for areas with NLCD canopy coverage. The lower biomass estimates here and those of Blackard et al. (2008) may be due to our use of 30 m grid NLCD canopy data rather than their use of 250 m grid MODIS canopy data or due to the gen-





eral underestimation of 2001 NLCD canopy fraction product (Nowak and Greenfield, 2012).

There are currently no databases to quantitatively evaluate the fractional tree species data coverage developed here. However the species range maps of Critchfield and Little (1966) and Little Jr. (1971, 1976) can be used for a qualitative evaluation. The tree species that constituted the largest fraction of biomass observations in the FIA data base generally fell within the tree species range maps (Fig. 3). Note that the maps represent a binary distribution of the tree species natural range and the BELD 4 estimates represent a gradient of species density. Species that did not constitute a large fraction in FIA observations typically had a much smaller estimated spatial range than indicated by the range maps. This could partially be due to the criteria, e.g. tree height greater than 5 m, etc., for trees carried over from the NLCD classification scheme or due to sparse sampling of these tree species in the FIA data base due to the species scarcity. However, these species likely represent a small fraction of the forest encourse and the species scarcity.

¹⁵ coverage in the domain and a small fraction of the domain wide BVOC emissions. Also, it is possible that tree coverage has changed in California since the 1970s when the trees were surveyed due to urban planning, plantations, fire, forest growth and climate change.

3.3 Describing changes in modeled BVOC estimates in Northern California

- ²⁰ Biogenic VOC emissions estimated with BEIS using the new canopy model (BEIS 3.61) and updated vegetation data (BELD 4) are shown for the northern California region in Fig. 4. A similar Figure of spatial biogenic emissions estimated with BEIS 3.14 and BELD 3 are shown in Fig. 5. In this model domain, isoprene emissions are highest in the foothills of the Sierra Nevada where high emitting isoprene vegetation (e.g. oak
 ²⁵ trees) are located. Monoterpene emissions are highest in the Sierra Nevada Mountains
- where high emitting needle leaf trees are located. Sesquiterpene emissions are highest in the Sacramento and San Joaquin valleys where grasses are common. Most other





biogenic VOC emissions show similar spatial patterns as isoprene or monoterpenes (Fig. 4).

The fractional coverage of oak (high isoprene emitting species) and needle leaf trees (high monoterpene emitting species) are shown using BELD 3 and BELD 4 in Fig. S2.

- ⁵ The BELD 4 representation shows a higher intensity of fractional coverage in much of the Sierra Nevada as county level information is allocated more spatially explicitly than in BELD 3. Smearing out vegetation coverage, as in BELD 3, will lead to lower emissions estimates where narrow features such as the band of oak trees in the western Sierra Nevada foothills exist and over predictions in areas that get allocated vegetation
- that does not exist in that area. Changes in oak and needle leaf fractional coverage between BELD 3 and BELD 4 are notable for both the Cool and Blodgett Forest sites meaning the observation data available at these locations is useful for evaluating the methodology used to generate BELD 4 (Fig. S2).

The updated leaf canopy module increases biogenic VOC emissions throughout Cal¹⁵ ifornia (Fig. 5). The changes to the vegetation input data show increases and decreases in isoprene and monoterpene emissions related to changing spatial allocation of high emitting vegetation species and changes to leaf area estimates. Sesquiterpene emissions generally decrease due to the changes in landuse and vegetation for this region (Fig. 5). The new vegetation allocation approach employed here for BELD 4 provides
²⁰ more detailed sub-County level representation of emitting species compared to BELD 3 and those changes are reflected in biogenic VOC emissions differences.

3.4 CMAQ estimates compared with CARES and BEARPEX measurements

The most recent publicly available version of BEIS (version 3.14) and BELD 3 vegetation input were used to provide biogenic emissions for a 4 km CMAQ simulation ²⁵ covering northern and central California for the period of time coincident with the 2009 BEARPEX field study. Additional simulations were done to illustrate the impact of updating the leaf canopy module in BEIS 3.61 and also how updating vegetation input data have on biogenic VOC model performance. Model runs were also done using



satellite derived PAR as input to BEIS rather than WRF estimated solar radiation. The MEGAN 2.1 model was also run using WRF and satellite estimates of PAR for the same domain and period.

- Temperature and solar radiation used for the biogenic emissions models were com pared to measurements at these field sites (Sacramento, Cool, and Blodgett Forest) to determine how meteorological inputs may bias model estimated BVOC. WRF model evaluation against meteorological variables is summarized in Table 3. The WRF model does well at capturing daytime high temperatures at Blodgett Forest and slightly over-estimates daily peak PAR. Daytime minimum temperatures at Blodgett Forest are
 largely overestimated by WRF. Temperature maximums and minimums are well characterized at Sacramento and Cool. The satellite estimated PAB underestimates the
- acterized at Sacramento and Cool. The satellite estimated PAR underestimates the ground measurements at Blodgett Forest on certain days but does better at capturing daytime peaks than WRF. In general, meteorological model performance at Blodgett Forest should result in overestimated emissions of isoprene and monoterpenes due to model overestimates in PAR and nighttime ambient temperature.

Field study measurements of isoprene and monoterpenes taken in 2010 at Sacramento and Cool and 2009 at Blodgett Forest provide an opportunity to better understand if the changes to BEIS and BELD better reflect the biogenic VOC gradient seen over these sites. Figure 6 shows the observed distribution of isoprene concentrations at Sacramento and Cool from 2010, Blodgett Forest in 2009, and model estimates from

- ²⁰ at Sacramento and Cool from 2010, Blodgett Forest in 2009, and model estimates from 2009 for the baseline CMAQ/BEIS simulation (BEIS 3.14 and BELD 3), canopy model updates (BEIS 3.61), vegetation data updates (BELD 4), and using satellite PAR with all formulation and other input data updates. Measured isoprene concentrations are lowest in Sacramento and highest at Cool where a high density of Oak trees exist. The
- ²⁵ baseline simulation predicts the highest isoprene at Blodgett Forest rather than Cool, but when canopy parameterization updates and vegetation data inputs are used the modeling system captures the gradient in concentration well across these three sites and also the distribution in observations at each site (Fig. 6).





Measured monoterpenes are highest at Blodgett Forest and lowest at Sacramento (Fig. 7). The baseline model captured this gradient but notably overestimated monoterpenes at Cool. When BELD 4 is used as input the modeling system compares much closer to observations at Cool and begins to slightly underestimate at Blodgett For-

sest. The use of satellite PAR rather than solar radiation estimated by WRF does little to change model performance of isoprene. Monoterpenes are not directly sensitive to PAR input and change little due to indirect use of PAR in the canopy model.

The MEGAN 2.1 model generally captures the gradient in observations between sites for isoprene and monoterpenes, but predicts much higher isoprene concentra-

- tions at each site compared to observations (see Fig. 6). This is consistent with other studies comparing MEGAN 2.1 isoprene flux with measurements in the Sierra Nevada of northern California (Misztal et al., 2014) and also with modeling systems using MEGAN 2.1 isoprene emissions compared with ambient isoprene concentrations in Texas (Kota et al., 2015) and southern Missouri (Carlton and Baker, 2011). Using the
- MEGAN model estimates of monoterpenes resulted in overestimates at Cool and underestimates at Blodgett Forest (Fig. 7). Estimates of isoprene using MEGAN improved when using satellite PAR as input rather than WRF solar radiation. This is consistent with similar evaluation in other parts of the United States (Carlton and Baker, 2011). The use of satellite PAR with MEGAN exacerbated monoterpene overestimates at Cool
- and increased model estimates at Blodgett Forest reducing the model underestimate. First generation oxidation products of isoprene (methacrolein and methyl vinyl ketones) were also measured at Blodgett Forest in 2009. Model performance is similar to isoprene where BEIS estimates compare favorably with measurements and MEGAN 2.1 emissions result in notable overestimates (Fig. S3) similar to previous studies (Kota et
- al., 2015). Methacrolein can further react in the atmosphere to form methacryloyl peroxynitrate (MPAN) which can form methacrylic acid epoxide (MAE) and subsequently secondary organic aerosol including aerosol methylglyceric acid, organic sulfates, and organic nitrates (Worton et al., 2013). CMAQ over-estimates MPAN at Blodgett Forest using either biogenic emisisons model, but overestimates are greater when using





MEGAN. Model performance for isoprene propagates through secondary reactions and could lead to similar over or under estimates of SOA.

4 Future direction

The updated biomass and tree species vegetation characterization in BELD would
⁵ benefit from additional evaluation for other parts of the conterminous United States. It is critically important to evaluate biogenic emissions models with field experiments designed for biogenic model evaluation or those that provide robust measurements of key biogenic VOC species such as those used for this assessment. Future work is planned to evaluate BEIS against a larger field study in California designed for biogenic
¹⁰ emissions model evaluation (2011 California Airborne BVOC Emission Research in Natural Ecosystem Transects; CABERNET) (Karl et al., 2013; Misztal et al., 2014) and also with a field study done in the southeast United States during the summer of 2013 (Southern Oxidant and Aerosol Study; SOAS). Evaluation of the model in urban areas would be useful although little field data exists for urban areas making this type of assessment difficult.

Code availability

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BEIS 3.61 code is available upon request prior to the public release of CMAQ v5.1 and available now in SMOKE 3.6.5 (https://www.cmascenter.org/smoke/). Please contact Jesse Bash at Bash.Jesse@epa.gov for more information. Additional model output, comparison with measurements and formulas used for data pairing are provided in the Supplement.

The Supplement related to this article is available online at doi:10.5194/gmdd-8-8117-2015-supplement.





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Table 1. Species emissions estimated by BEIS and mapping to the SAPRC07T and CB6r2 gas phase chemical mechanism lumped species.

Emitted Specie	BEIS Abbreviation	SAPRC07 Species	CB6r2 Species
ethene	ETHE	ETHENE	ETH
ethane	ETHA	ALK1	ETHA
methanol	METH	MEOH	MEOH
ethanol	ETHO	ALK3	ETOH
formaldehyde	FORM	НСНО	FORM
acetaldehyde	ACTAL	ССНО	ALD2
formic acid	FORAC	НСООН	FACD
acetic acid	ACTAC	ССООН	AACD
propene	PROPE	OLE1	33.3 % PAR + 66.7 % OLE
hexenol	HEXE	OLE1	33.3 % PAR + 66.7 % IOLE
hexenylacetate	HEXY	OLE1	37.5 % PAR + 50 % IOLE + 12.5 % NR
butenone	BUTO	OLE1	50 % PAR + 50 % OLE
2-methyl-3-buten-2-ol	MBO	OLE2	60 % PAR + 40 % OLE
butene	BUTE	OLE2	50 % PAR + 50 % OLE
acetone	ACET	ACETONE	ACET
hexanal	HEXA	RCHO	66.7 % PAR + 33.3 % ALDX
Other Reactive VOCs	ORVOC	10 % OLE2 + 85 % ALK2 + 5 % NR	80 % PAR + 20 % OLE
Isoprene	ISOP	ISOPRENE	ISOP
α-pinene	APIN	TRP1	TERP
β -pinene	BPIN	TRP1	TERP
$\delta - 3$ -carene	D3CAR	TRP1	TERP
δ – limonene	DLIM	TRP1	TERP
camphene	CAMPH	TRP1	TERP
myrcene	MYRC	TRP1	TERP
α -terpinene	ATERP	TRP1	TERP
β -phellandrene	BPHE	TRP1	TERP
sabinene	SABI	TRP1	TERP
<i>p</i> −cymene	PCYM	TRP1	TERP
ocimene	OCIM	TRP1	TERP
α -thujene	ATHU	TRP1	TERP
terpinolene	TRPO	TRP1	TERP
γ-terpinene	GTERP	TRP1	TERP
Sesquiterpines	SESQ	SESQ	SESQ
Carbon Monoxide	CO	CO	CO
Nitric Oxide	NO	NO	NO
	Emitted Specie ethene ethane methanol formaldehyde acetia acid propene hexenol hexenylacetate butenone 2-methyl-3-buten-2-ol butene acetone hexanal Other Reactive VOCs Isoprene β -pinene β -pinene δ -3-carene δ -limonene camphene myrcene α -terpinene β -phellandrene sabinene ρ -cymene ocimene α -thujene terpinolene γ -terpinene Sesquiterpines Carbon Monoxide Nitric Oxide	Emitted SpecieBEIS AbbreviationetheneETHEethaneETHAmethanolMETHethanolETHOformaldehydeFORMacetia acidACTALformic acidFORACacetia acidACTACpropenePROPEhexenolHEXEhexenylacetateHEXYbutenoneBUTO2-methyl-3-buten-2-olMBObuteneBUTOacetic acidACEThexanalHEXAOther Reactive VOCsORVOClsopreneISOP α -pineneAPIN β -pineneBPIN δ - 3-careneD3CAR δ -limoneneDLIMcampheneCAMPHmyrceneMYRC α -terpineneATERP β -phellandreneBPHEsabineneSABI ρ -cymenePCYMocimeneOCIM α -thujeneATHUterpinoleneTRPO γ -terpineneSESQCarbon MonoxideCONitric OxideNO	Emitted SpecieBEIS AbbreviationSAPRC07 SpeciesetheneETHEETHENEethaneETHAALK1methanolMETHMEOHethanolETHOALK3formaldehydeFORMHCHOacetaldehydeACTALCCHOformic acidFORACHCOOHacetic acidACTACCCOOHacetic acidACTACCCOOHpropenePROPEOLE1hexenolHEXEOLE1butenoneBUTOOLE12-methyl-3-buten-2-olMBOOLE2buteneBUTEOLE2acetora active VOCSORVOC10% OLE2 + 85% ALK2 + 5% NRIsopreneISOPISOPRENE α -pineneAPIINTRP1 β -pineneDIACARTRP1 β -pineneDIACARTRP1 β -pineneDACARTRP1 β -phellandreneDLIMTRP1 β -phellandreneSABITRP1 β -phellandreneSABITRP1 ρ -cymeneOCIMTRP1 α -tupineneSABITRP1 ρ -cymeneCOIMTRP1 α -tupineneSABITRP1 ρ -cymeneGEYMTRP1 ϕ -cymeneGEYMTRP1 ϕ -tupineneSABITRP1 ϕ -cymeneGCIMTRP1 ϕ -tupineneGTERPTRP1 φ -tupineneGERPTRP1 φ -tupineneGERPTRP1 φ -tupineneGERPTRP





Table 2. Emissions $(ug m^{-2} h^{-1})$ for each specie estimated by BEIS. Median, minimum, and maximum emission rates for each aggregated land cover/vegetation group are shown. Emission rates are uniform for some vegetation categories resulting in the same value for median, minimum, and maximum.

		Pine			Fir			Spruce			Oak			Maple			Other			Crops			Grass	
																	Deciduous							
Number of species	40	40	40	12	12	12	9	9	9	44	44	44	13	13	13	684	684	684	42	42	42	2	2	2
Metric	Median	Min	Max	Median	Min	Max	Median	Min	Max	Median	Min	Max	Median	Min	Max	Median	Min	Max	Median	Min	Max	Median	Min	Max
Isoprene (ug m ⁻² h ⁻¹)	79	79	79	170	170	170	11900	1700	11900	29750	29750	29750	43	43	43	43	43	29750	10	1	102	56	56	56
Sesquiterpenes	70	70	210	150	150	150	150	150	150	37	37	37	37	37	37	37	37	150	29	29	29	29	29	29
Nitric Oxide	4	4	4	4	4	4	4	4	4	4	4	4	4	4	4	4	4	4	160	0	774	58	58	58
MBO	76	0	52675	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	11	11	11
apinene	840	28	2100	1038	239	1472	881	449	1176	26	26	26	127	127	127	15	0	1839	8	0	102	9	9	9
bpinene	420	0	1134	519	346	929	322	75	716	5	5	5	26	26	26	8	0	580	3	0	51	5	5	5
d3carene	57	0	867	260	0	260	229	0	730	0	0	0	150	150	150	3	0	280	2	0	26	2	2	2
dlimonene	48	0	1290	260	107	792	260	2	688	10	10	10	78	78	78	3	0	233	2	0	26	2	2	2
camphene	7	0	406	260	62	260	260	57	748	6	6	6	31	31	31	3	0	210	2	0	26	2	2	2
myrcene	37	0	611	260	39	260	218	54	1340	0	0	0	48	48	48	3	0	74	2	0	26	2	2	2
aterpinene	0	0	96	0	0	324	0	0	78	0	0	0	3	3	3	0	0	18	0	0	0	0	0	0
bphellandrene	0	0	221	0	0	779	78	0	488	0	0	0	0	0	0	0	0	35	0	0	0	0	0	0
sabinene	0	0	263	0	0	260	0	0	86	0	0	0	129	129	129	0	0	61	0	0	0	0	0	0
pcymene	0	0	462	0	0	221	2	0	173	8	8	8	0	0	0	0	0	162	0	0	0	0	0	0
ocimene	0	0	20	0	0	0	0	0	0	10	10	10	0	0	0	0	0	248	0	0	0	0	0	0
athujene	0	0	82	0	0	26	0	0	0	0	0	0	5	5	5	0	0	91	0	0	0	0	0	0
terpinolene	0	0	37	0	0	75	2	0	10	9	9	9	0	0	0	0	0	34	0	0	0	0	0	0
gterpinene	0	0	7	0	0	70	2	0	8	0	0	0	5	5	5	0	0	28	0	0	0	0	0	0
methanol	1120	1120	1120	2400	2400	2400	2400	2400	2400	600	600	600	600	600	600	600	600	2400	480	480	480	480	480	480
ethene	74	74	74	158	158	158	158	158	158	40	40	40	40	40	40	40	40	158	32	32	32	32	32	32
propene	74	74	74	158	158	158	158	158	158	40	40	40	40	40	40	40	40	158	32	32	32	32	32	32
ethanol	121	121	121	259	259	259	259	259	259	65	65	65	65	65	65	65	65	259	52	52	52	52	52	52
acetone	102	102	102	218	218	218	218	218	218	55	55	55	55	55	55	55	55	218	44	44	44	44	44	44
hexanal	38	38	38	82	82	82	82	82	82	20	20	20	20	20	20	20	20	82	16	16	16	16	16	16
hexenol	156	156	156	333	333	333	333	333	333	83	83	83	83	83	83	83	83	333	67	67	67	67	67	67
hexenylacetate	166	166	166	355	355	355	355	355	355	89	89	89	89	89	89	89	89	355	71	71	71	71	71	71
formaldehyde	70	70	70	150	150	150	150	150	150	38	38	38	38	38	38	38	38	150	30	30	30	30	30	30
acetaldehyde	51	51	51	110	110	110	110	110	110	28	28	28	28	28	28	28	28	110	22	22	22	22	22	22
butene	33	33	33	70	70	70	70	70	70	18	18	18	18	18	18	18	18	70	14	14	14	14	14	14
ethane	18	18	18	38	38	38	38	38	38	10	10	10	10	10	10	10	10	38	8	8	8	8	8	8
formic_acid	54	54	54	115	115	115	115	115	115	31	31	31	31	31	31	31	31	115	23	23	23	23	23	23
acetic_acid	35	35	35	75	75	75	75	75	75	20	20	20	20	20	20	20	20	75	15	15	15	15	15	15
butenone	20	20	20	44	44	44	44	44	44	12	12	12	12	12	12	12	12	44	9	9	9	9	9	9
Carbon monoxide	490	490	490	1050	1050	1050	1050	1050	1050	264	264	264	264	264	264	264	264	1050	210	210	210	210	210	210
Other reactive VOC	57	0	57	122	122	122	122	122	122	31	31	31	31	31	31	31	31	122	25	25	25	25	25	25



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	Scenario	Location	Units	N	Average	Average	Meadian	Median	Average	Average	Fractional	Fractional
					Observation	Prediction	Bias (%)	Error (%)	Bias	Error	Bias (%)	Error (%)
Isoprene	BEIS v3.14	Blodgett Forest	ppb	155	1.4	2	26.0	56.0	0.5	1.1	-0.4	73.9
Isoprene	BEIS v3.6 WRF par	Blodgett Forest	ppb	155	1.4	1.5	-6.0	49.0	0.1	0.8	-22.3	70.3
Isoprene	BEIS v3.61 SAT par	Blodgett Forest	ppb	155	1.4	1.4	-18.0	49.0	0.0	0.9	-34.4	76.3
Isoprene	MEGAN v2.1 WRF par	Blodgett Forest	ppb	153	1.4	4.6	203.0	203.0	3.2	3.5	60.3	108.6
Isoprene	MEGAN v2.1 SAT par	Blodgett Forest	ppb	153	1.4	3.4	97.0	110.0	2.0	2.5	26.3	101.5
Monoterpenes	BEIS v3.14	Blodgett Forest	ppb	855	0.7	0.8	-10.0	43.0	0.1	0.4	-13.8	58.0
Monoterpenes	BEIS v3.61 WRF par	Blodgett Forest	ppb	855	0.7	0.6	-20.0	40.0	-0.1	0.3	-31.2	57.2
Monoterpenes	BEIS v3.61 SAT par	Blodgett Forest	ppb	855	0.7	0.6	-21.0	41.0	-0.1	0.3	-33.2	58.6
Monoterpenes	MEGAN v2.1 WRF par	Blodgett Forest	ppb	855	0.7	0.4	-42.0	44.0	-0.3	0.4	-64.1	69.2
Monoterpenes	MEGAN v2.1 SAT par	Blodgett Forest	ppb	855	0.7	0.5	-32.0	39.0	-0.2	0.3	-45.8	58.5
MVK+MACR	BEIS v3.14	Blodgett Forest	ppb	157	1.3	0.9	-29.0	33.0	-0.4	0.5	-44.5	60.8
MVK+MACR	BEIS v3.61 WRF par	Blodgett Forest	ppb	157	1.3	1.4	-4.0	43.0	0.1	0.7	-21.9	65.2
MVK+MACR	BEIS v3.61 SAT par	Blodgett Forest	ppb	157	1.3	1.3	-9.0	47.0	0.0	0.7	-31.8	69.3
MVK+MACR	MEGAN v2.1 WRF par	Blodgett Forest	ppb	155	1.3	2.5	69.0	83.0	1.2	1.6	28.3	82.7
MVK+MACR	MEGAN v2.1 SAT par	Blodgett Forest	ppb	155	1.3	1.6	12.0	61.0	0.4	1.0	-11.4	77.7
Wind Speed	WRF	Cool	m s ⁻¹	920	2.1	2.8	37.0	40.0	0.7	0.9	30.4	39.3
Wind Speed	WRF	Sacramento	m s ⁻¹	1266	2.1	2.8	38.0	41.0	0.8	0.9	34.0	41.8
Wind Speed	WRF	Blodgett Forest	m s ⁻¹	1035	1.5	2.9	104.0	104.0	1.3	1.4	63.9	66.9
Temperature	WRF	Cool	C	1786	22.2	23.1	5.0	7.0	0.9	1.6	5.3	8.1
Temperature	WRF	Sacramento	Ċ	1721	22.2	22.5	2.0	5.0	0.2	1.4	1.6	6.4
Temperature	WRF	Blodgett Forest	c	1035	18.4	22.6	28.0	29.0	4.2	5.6	28.4	34.1
PAR	WRF	Blodgett Forest	watts m ⁻²	1056	148.3	167.6	0.0	47.0	19.2	45.5	-11.3	52.3
PAR	Satellite estimate	Blodgett Forest	watts m ⁻²	1056	148.3	131.5	0.0	30.0	-16.8	44.3	-39.5	58.0
DM organia corbon	PEIC v2 14		ug m ⁻³	141	17	1.1	24.0	40.0	0.6	1.0	42.0	60.6
DM arganic carbon	DEIG VO. 14		ug m 	141	1.7		-34.0	45.0	-0.0	1.0	-43.2	70.0
PINI2.5 organic carbon	BEIS V3.61 WHF par	INPROVE siles	ug m	141	1.7	1.1	-35.0	50.0	-0.6	1.0	-44.9	70.9
PM _{2.5} organic carbon	BEIS V3.61 SAI par	IMPROVE sites	ug m -	141	1.7	1.1	-35.0	50.0	-0.6	1.0	-45.6	/1.5
PM _{2.5} organic carbon	MEGAN v2.1 WRF par	IMPROVE sites	ug m ⁻³	141	1.7	1.8	8.0	43.0	0.1	1.2	-0.8	57.9
PM _{2.5} organic carbon	MEGAN v2.1 SAT par	IMPROVE sites	ug m ⁻³	141	1.7	2.2	11.0	47.0	0.5	1.4	9.1	62.5
O ₃ greater than 60	BEIS v3.14	AQS sites	ppb	7125	70.9	64.8	-8.0	13.0	-6.1	11.2	-10.1	16.9
O ₃ greater than 60	BEIS v3.61 WRF par	AQS sites	ppb	7125	70.9	64.7	-8.0	13.0	-6.2	11.0	-10.1	16.7
O ₃ greater than 60	BEIS v3.61 SAT par	AQS sites	ppb	7125	70.9	64.3	-9.0	13.0	-6.6	11.0	-10.8	16.8
O ₃ greater than 60	MEGAN v2.1 WRF par	AQS sites	ppb	7125	70.9	65.4	-9.0	14.0	-5.5	12.0	-9.5	17.8
O ₃ greater than 60	MEGAN v2.1 SAT par	AQS sites	ppb	7125	70.9	62.1	-12.0	14.0	-8.8	11.9	-14.1	18.3
O ₃ less than 60	BEIS v3.14	AQS sites	ppb	48 939	32.0	41.0	29.0	33.0	8.9	11.2	30.2	36.6
O ₃ less than 60	BEIS v3.61 WRF par	AQS sites	ppb	48 939	32.0	40.8	29.0	32.0	8.8	11.1	29.8	36.4
O ₃ less than 60	BEIS v3.61 SAT par	AQS sites	ppb	48 939	32.0	40.7	29.0	32.0	8.7	11.0	29.4	36.2
O ₃ less than 60	MEGAN v2.1 WRF par	AQS sites	ppb	48 939	32.0	41.7	32.0	34.0	9.7	11.8	31.9	37.9
O ₃ less than 60	MEGAN v2.1 SAT par	AQS sites	ppb	48 939	32.0	40.7	29.0	32.0	8.7	11.0	30.0	36.4

Table 3. Model evaluation against field campaigns and network observations.



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Figure 1. Diurnal observed, MEGAN 2.1 and BEIS 3.61 estimated leaf temperatures (top left); MEGAN 2.1 and BEIS 3.61 leaf temperature estimates plotted against skin temperature observations (top right); observed, MEGAN 2.1, and BEIS 3.61 estimated gradient between leaf and ambient temperatures (bottom left); MEGAN 2.1 and BEIS 3.61 estimated leaf temperature biases (model-observed) (bottom right).







Figure 2. Total above ground forest biomass (Mg ha⁻¹) estimates for BELD 4 (left) and Blackard et al. (2008) (right) projected onto the 4 km California model domain.





Figure 3. BELD 3 spatial allocation of Ponderosa Pine (Pinus ponderosa, top left), BELD 4 spatial allocation, (top center), and the absolute difference between the BELD 4 and BELD 3 spatial allocation (top right). BELD 3 spatial allocation of Canyon Live Oaks (Quercus chrysolepis, bottom left), BELD 4 spatial allocation, (bottom center), and the absolute difference between the BELD 4 and BELD 3 spatial allocation (bottom right). The natural range maps of Critchfield and Little (1966) and Little (1971, 1976) are represented by the dashed red lines.







Figure 4. BEIS 3.61/BELD 4 estimated total emissions (tons) for the modeling period.







Figure 5. Baseline BEIS 3.14/BELD 3 emissions (tons; left column) and difference between canopy update and baseline BEIS 3.61/BELD 3 (center column) and between the canopy update and landuse/vegetation species updates BEIS 3.61/BELD 4 (right column).







Figure 6. Distribution of observed and modeled isoprene. Observations at Sacramento and Cool represent June 2010. Observations at Blodgett Forest match the modeled period.







Figure 7. Distribution of observed and modeled monoterpenes. Observations at Sacramento and Cool represent June 2010. Observations at Blodgett Forest match the modeled period.



