Top-down estimates of biomass burning emissions of black carbon in the Western United States

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Abstract. We estimate biomass burning and anthropogenic emissions of black carbon (BC) in the western US for May–October 2006 by inverting surface BC concentrations from the Interagency Monitoring of PROtected Visual Environment (IMPROVE) network using a global chemical transport model. We first use active fire counts from the Moderate Resolution Imaging Spectroradiometer (MODIS) to improve the spatiotemporal distributions of the biomass burning BC emissions from the Global Fire Emissions Database (GFEDv2). The adjustment primarily shifts emissions from late to middle and early summer (a 33 % decrease in September–October and a 56 % increase in June–August) and leads to appreciable increases in modeled surface BC concentrations in early and middle summer, especially at the 1–2 and 2–3 km altitude ranges. We then conduct analytical inversions at both 2\(^\circ\) \times 2.5\(^\circ\) (nested over North America) horizontal resolutions. The a posteriori biomass burning BC emissions for July–September are 31.7 Gg at 2\(^\circ\) \times 2.5\(^\circ\) (an increase by a factor of 4.7) and 19.2 Gg at 0.5\(^\circ\) \times 0.667\(^\circ\) (an increase by a factor of 2.8). The inversion results are rather sensitive to model resolution. The a posteriori biomass burning emissions at the two model resolutions differ by a factor of \sim 6 in California and the Southwest and by a factor of 2 in the Pacific Northwest. The corresponding a posteriori anthropogenic BC emissions are 9.1 Gg at 2\(^\circ\) \times 2.5\(^\circ\) (a decrease of 48 %) and 11.2 Gg at 0.5\(^\circ\) \times 0.667\(^\circ\) (a decrease of 36 %). Simulated surface BC concentrations with the a posteriori emissions capture the observed major fire episodes at most sites and the substantial enhancements at the 1–2 and 2–3 km altitude ranges. The a posteriori emissions also lead to large bias reductions (by \sim 30 % on average at both model resolutions) in modeled surface BC concentrations and significantly better agreement with observations (increases in Taylor skill scores of 95 % at 2\(^\circ\) \times 2.5\(^\circ\) and 42 % at 0.5\(^\circ\) \times 0.667\(^\circ\)).

1 Introduction

Black carbon (BC), as a component of fine particulate matter, has deleterious effects on human health (e.g., Anenberg et al., 2011, 2012; Smith et al., 2009). BC is also known as the agent to cause both degraded air quality (e.g., Anenberg et al., 2011, 2012) and warming due to its strong absorption of solar radiation (e.g., Ramanathan and Carmichael, 2008; Horvath, 1993). BC thus has considerable impacts on global climate (Fuglestvedt et al., 2010; Shindell et al., 2008; Levy II et al., 2008; Reddy et al., 2007; IPCC, 2007; Jacobson, 2001, 2004). BC deposited on snow and ice can significantly decrease the surface albedo (Warren and Wiscombe, 1980) and quicken surface melt (e.g., Flanner et al., 2007, 2009; Hansen and Nazarenko, 2004; Zwally et al., 2002). BC thus has a significant contribution to the observed rapid glacier retreats (Painter et al., 2013; Xu et al., 2009). Because of its shorter lifetime relative to long lived greenhouse gases such as carbon dioxide, BC shows a much stronger regional
warming effect and its reduction may provide an efficient solution to mitigate near-term climate change and to improve air quality and human health simultaneously (Bond et al., 2013; Shindell et al., 2012; Kopp and Maurerall, 2010; Ramana et al., 2010; Jacobson, 2002, 2010; Bond and Sun, 2005; Hansen et al., 2005).

The transport and subsequent deposition of BC is known to impact the regional climate and hydrological cycle in the western US mountain ranges. Mountain snowmelt accounts for over 70% of the annual stream flows in this region (Qian et al., 2009). Qian et al. (2009) showed that the deposition of BC on snow over the western US mountain ranges reduced snow accumulation in winter, snowmelt in spring, and runoff between April and June. The ongoing and most severe drought on record during the last 5 decades in California (e.g., Mirchi et al., 2013) and the dwindling water level of the Colorado River (e.g., Vano et al., 2013) both add even more urgency to better understand the sources, transport, and deposition of BC in the western US mountain ranges.

The uncertainty in current BC emission estimates ranges from at least ±50% on global scales to a factor of 2–5 on regional scales (Ramanathan and Carmichael, 2008; Streets et al., 2001, 2003). The uncertainty of biomass burning BC emissions in the western US is equally large, not only in the absolute magnitudes of fire emissions but also in the timing and location of fires (Mao et al., 2011). Part of the large uncertainty stems from estimates of burned area and fuel load (Langmann et al., 2009), especially of small fires, including agricultural burning (Randerson et al., 2012). Giglio et al. (2006, 2010) have shown that small fires can lead to high relative errors of 50–100% in the burned area estimates. The uncertainty is likely because of the lack of detection of small fires in the burned area algorithms (Randerson et al., 2012; van der Werf et al., 2010; McCarty et al., 2009; Roy and Boschetti, 2009; Korontzi et al., 2006). In addition, the increases of fire frequency, fire season length, and annual burned area observed in the western US in recent decades have been linked to the warm climate (Peterson and Marcinkowski, 2014; Westerling et al., 2006). The annual burned area in the western US is also projected to increase by 25–125% in the 2050s relative to the present under future warming (Yue et al., 2013; Spracklen et al., 2009). It is therefore conceivable that, biomass burning emissions as an important source to BC will have an even larger contribution to BC in the western US in the 21st century.

Understanding the distributions of a chemical species in the atmosphere depends on the information of the emissions. Bottom-up emission estimates generally rely on emission factors using socioeconomic, energy, land use, or environmental data (Bond et al., 2007, 2013; Lu et al., 2011). In recent years, there has been an increasing emphasis on the use of inverse methods to characterize the temporal and spatial variability of emissions. Top-down inversions have been widely used for estimating emission fluxes of long-lived trace gases such as carbon dioxide (e.g., Pickett-Heaps et al., 2011; Chevallier et al., 2007; Gloor et al., 1999), methane (e.g., Wecht et al., 2012; Meirink et al., 2008; Hein et al., 1997), and carbon monoxide based on observations from surface stations (e.g., Bergamaschi et al., 2000; Kasibhatla et al., 2002), aircraft (e.g., Palmer et al., 2003, 2006), and satellites (Jiang et al., 2011, 2013; Jones et al., 2009; Stavrakou and Müller, 2006; Arellano et al., 2004, 2006), when the atmospheric concentrations are linearly or weakly non-linearly dependent on their emissions (Müller and Stavrakou, 2005). Top-down inversions, by nature, are a way to examine the consistency of model results with observations. It is important to note that the abovementioned studies have consistently demonstrated that the bottom-up and top-down approaches are not mutually exclusive. Improving estimate of emissions is inherently an iterative process and it is imperative to optimally integrate both bottom-up and top-down approaches.

Several studies have used multiple regressions to estimate annual mean sources of primary carbonaceous aerosols over the US (Park et al., 2003) and in China (Wang et al., 2013; Fu et al., 2012). The goal of the present study is to improve our understanding of sources of BC in the western US mountain ranges, with a particular focus on biomass burning emissions during May–October 2006, broadly encompassing the fire season in the region. We first improve the spatial distributions and the seasonal and interannual variations of BC emissions using high spatial resolution active fire counts. We then apply top-down inversions of surface BC concentration measurements using a global 3-dimensional chemical transport model (CTM). We briefly describe the observations in Sect. 2 and the model in Sect. 3. We then discuss in Sect. 4 improvements to the spatiotemporal distributions of biomass burning emissions of BC. In Sect. 5, we describe the analytical inversion method. The inversion results and discussions are presented in Sect. 6. We give a summary and draw conclusions in Sect. 7.

2 Observations

2.1 IMPROVE

The Interagency Monitoring of PROtected Visual Environment (IMPROVE) surface network in the US was created for the protection of visibility in Class I remote areas (Malm et al., 1994). Long-term measurements of aerosols with chemical species including BC are available from the network since 1987 (data available at http://vista.cira.colostate.edu/improve/). We use surface BC concentration data for 2006 from 69 sites across the western US (Fig. 1 and Supplement Table S1). These are mostly remote sites located predominantly in national parks. BC mass concentration is measured by the thermal optical reflectance (TOR) combustion method based on the preferential oxidation of organic carbon (OC) and BC at different temperatures (Chow et al., 2004). The
Figure 1. The 69 IMPROVE sites (black dots) used in this study (data available at http://vista.cira.colostate.edu/improve/). Also shown are the three biomass burning regions used to define the state vector of the inversion analyses: the Rocky Mountains (RM), California and the Southwest (CSW), and the Pacific Northwest (PNW). Terrain heights are indicated by color contours.

2.2 MODIS

The Moderate Resolution Imaging Spectroradiometer (MODIS) active fire product (available at ftp://fuoco.geog.umd.edu) detects fires in 1 km pixels that are burning at the time of overpass under relatively cloud-free conditions using a contextual algorithm (Giglio et al., 2003). The detection utilizes the strong emission of mid-infrared radiation from fires and applies thresholds to the observed middle-infrared and thermal-infrared brightness temperatures. The algorithm examines each pixel of the MODIS swath, and ultimately assigns to each one of the following classes: missing data, cloud, water, non-fire, fire, or unknown. We use here MODIS active fire counts at the spatial resolution of 0.5° x 0.5° and a temporal resolution of 8 days.

3 Model description and simulations

We apply the GEOS-Chem global 3-dimensional CTM (Bey et al., 2001, with many updates thenceforth) to analyze IMPROVE surface BC observations. GEOS-Chem is driven by assimilated meteorological observations from the Goddard Earth Observing System (GEOS) of the NASA Global Modeling and Assimilation Office (GMAO). Here we use GEOS-Chem version 8-03-02 (available at http://geos-chem.org) driven by GEOS-5 meteorological data with a temporal resolution of 6 h (3 h for surface variables and mixing depths), horizontal resolutions of 2° (latitude) x 2.5° (longitude) and 0.5° (latitude) x 0.667° (longitude), and 47 hybrid eta levels in the vertical from the surface to 0.01 hPa. The lowest model levels are centered at approximately 60, 200, 300, 450, 600, 700, 850, 1000, 1150, 1300, 1450, 1600, and 1800 m above the local surface. GEOS-Chem simulations at the finer horizontal resolution are typically nested over a continental-scale domain within the global domain. Lateral boundary conditions for the nested-grid simulations are archived from a global simulation. The nested-grid simulation employs the same meteorology, dynamics, and chemistry as the global simulation, thus allows for consistent propagation of features from the global to the nested domain via one-way nesting.

Tracer advection is computed every 15 minutes with a flux-form semi-Lagrangian method (Lin and Rood, 1996). Tracer moist convection is computed using GEOS convective, entrainment, and detrainment mass fluxes as described by Allen et al. (1996a, b). The deep convection in GEOS-5 is parameterized using the relaxed Arakawa-Schubert scheme (Moorthi and Suarez, 1992; Arakawa and Schubert, 1974), and the shallow convection treatment follows Hack (1994). Simulation of aerosol wet and dry deposition follows Liu et al. (2001). Wet deposition includes contributions from scavenging in convective updrafts, rainout from convective anvils, and rainout and washout from large-scale precipitation. Dry deposition of aerosols uses a resistance-in-series model (Walcek et al., 1986) dependent on local surface type and meteorological conditions.

GEOS-Chem simulation of carbonaceous aerosols has been reported previously by Park et al. (2003). Eighty percent of BC and 50 % of OC emitted from primary sources are assumed to be hydrophobic, and hydrophobic aerosols become hydrophilic with an e-folding time of 1.2 days (Park et al., 2003; Chin et al., 2002; Cooke et al., 1999). BC aerosols in the model are assumed to be externally mixed. Global anthropogenic (fossil fuel and biofuel) emissions of BC are based upon Bond et al. (2007) with imposed seasonality following Park et al. (2003). Biomass burning emissions of BC are from the Global Fire Emissions Database version 2 (GFEDv2) emissions with an 8-day temporal resolution (Randerson et al., 2007; van der Werf et al., 2006). We include for comparison BC emissions from GFEDv3 (van der Werf et al., 2010) and the Fire Locating and Monitoring of Burning Emissions (FLAMBE) inventory (Reid et al., 2009).

GFEDv2 was derived using satellite observations including active fire counts and burned areas in conjunction with a biogeochemical model (Randerson et al., 2007; van der Werf et al., 2006). Burned area was derived using monthly 1° x 1° active fire and 500 m burned area data from MODIS (Giglio et al., 2006). Total carbon emissions were then calculated
as the product of burned area, fuel load, and combustion completeness. Fuel load depends on vegetation type, climate, soil type, and time passed since the last fire, while combustion completeness (the fraction of the available fuel combusted during a fire) depends on the type of fire, fuel (e.g., stems, leaves, and litter) and its moisture content (Langmann et al., 2009). The Carnegie–Ames–Stanford Approach (CASA) biogeochemical model was used to estimate combustion completeness as well as fuel load and the associated spatial variability (van der Werf et al., 2006, and references therein). BC emissions were then derived from the total carbon emissions and BC emission factors. GFEDv3 includes several updates over GFEDv2: a finer spatial resolution (0.5◦), a better mapping method in estimating 500 m global burned area, a local (in lieu of regional) regression in producing the indirect, active-fire based estimates of burned area, and an improved CASA model.

FLAMBE provides carbon emissions at a 1° × 1° spatial resolution and hourly temporal resolution based on both MODIS and Geostationary Operational Environmental Satellites (GOES) fire counts. Hourly emissions are available from 2005. Fire pixel detections and subpixel burning characterizations were computed based on the operational NOAA/NESDIS GOES Wildfire Automated Biomass Burning Algorithm (WF_ABBA) for most of the Western Hemisphere. For the rest of the globe, the near real time University of Maryland/NASA MODIS fire products from Terra and Aqua were used. The Advanced Very High Resolution Radiometer (AVHRR) Global Land Cover Characteristics (GLCC) data (v2.0) was used to assign surface emissivity and to screen for false alarms. Based on the thermal anomaly and fire radiative power in a subpixel fire, FLAMBE provides hourly burned areas and carbon emissions. Following Fisher et al. (2010), we calculate BC emissions using emission factors from Andreade and Merlet (2001).

For computational expediency, we conduct GEOS-Chem “offline” simulations of carbonaceous aerosols following Mao et al. (2011). We conduct simulations for 2006 at both 2° × 2.5° and 0.5° × 0.667° resolutions. The first 3 months are used for initialization and we focus our analysis on May through October. The finer resolution is nested over North America (40–140° W longitudes, 10–70° N latitudes, cf. Fig. 1 in Wang et al., 2004). Wang et al. (2004) reported the first nested-grid GEOS-Chem simulations of carbon monoxide over North America. Chen et al. (2009) provided a detailed description of the one-way nesting in the model.

Model results are sampled at the corresponding locations of the IMPROVE sites. IMPROVE observations are 24 h averages sampled every 3 days and we sample the model accordingly. It is known that comparing localized observations such as the IMPROVE data with model results that are representative of a much larger area is inherently problematic (Mao et al., 2011; Fairlie et al., 2007). Furthermore, many of the IMPROVE sites are at high elevations and the associated up-slope flow is difficult to represent in a coarse-resolution model.

4 Spatiotemporal distributions of biomass burning BC emissions

Mao et al. (2011) pointed out that the GFEDv2 inventory not only underestimated the magnitude but also inadequately captured the temporal (and presumably spatial) distribution of biomass burning emissions in the western US (WUS, defined hereinafter as 30–50° N, 100–125° W for clarity). Ample evidence has suggested that the root cause of these deficiencies was likely the lack of detection of small fires, for example, prescribed and agricultural burning (e.g., Randerson et al., 2012; Giglio et al., 2010). Randerson et al. (2012) pointed out that current global burned area products largely missed small fires because such fires were typically well below the detection limit of the burned area algorithms. For instance, GFEDv2 burned area was derived from MODIS 500 m surface reflectance, suited for detecting large fires with fire scars greater than 500 m (Giglio et al., 2006, 2010). Active fire data has long been used as a proxy for burned area due to the lack of long-term global burned area data (Giglio et al., 2006, 2010). MODIS active fire data, based on thermal anomalies, allows for the detection of fires that are an order of magnitude smaller in size than those detected by MODIS 500 m surface reflectance. MODIS active fire data thus preserves detailed aspects of the spatial distribution and seasonality of burning (Randerson et al., 2012, and references therein).

Here we use MODIS active fire counts from 2005–2007 to improve the spatiotemporal distribution of GFEDv2 BC emissions in North America. The adjusted emissions will be the a priori for the inversions presented hereinafter. There are large uncertainties in relating fire counts to actual burned area because of inadequate temporal sampling, variability in fuel conditions and cloud cover, differences in fire behavior, and issues related to spatial resolution (Giglio et al., 2006; Kasischke et al., 2003). We choose the 3-year period in part to minimize such uncertainties. We adjust the emissions for three zones: boreal North America, temperate North America, and Mexico and Central America, following the geographic regions defined in the GFED inventory (Giglio et al., 2006, 2010). We first sum up for each zone the monthly carbon emissions and MODIS active fire counts from 2005–2007. We then redistribute the emissions according to the ratio of the active fire counts for each 8-day period within each grid box (0.5° × 0.5°) to the total fire counts. We note that the adjusted GFEDv2 emissions have the same total emissions as the original GFEDv2. We would like to point out that, fuel loading and combustion completeness are also basic factors to derive emissions and have large spatial and temporal variations. The emission adjustment presented here therefore would introduce some uncertainties without considering
these issues. We make another emission adjustment with the same calculation method but using daily active fire counts and carbon emissions only in the WUS for 2006 to partially measure these uncertainties. The resulting emissions are essentially the same as the former adjusted emissions and we thus focus our analysis on the former adjusted emissions hereinafter.

Figure 2 shows the active fire counts and the biomass burning emissions of BC (middle and bottom panels) summed over the western US from 2005 to 2007. Of the 3 years, 2006 is a relatively large fire year. Fire seasons in the WUS typically last from late June to October even November, as evident in both the fire counts and the emissions. The fire season in 2006 is from July to September primarily. The adjustment largely shifts emissions from late to mid-summer. Take 2006 for example, the adjustment results in a 33% decrease in September–October and a 56% increase in June–August. The spatial distributions of monthly BC emissions for July–September 2006, before and after the adjustments, are shown in Fig. 3. There are significant increases in the agricultural areas in August and September, e.g., Columbia River Basin in Washington, Oregon, and Idaho, and Snake River Basin in Idaho. These increases reflect improved spatial allocation of biomass burning emissions using MODIS active fire counts, which now presumably pick up (small) agricultural burning. The adjusted emissions track spatiotemporally MODIS active fire counts (Fig. 3a), as expected.

The spatiotemporal shift of the emissions manifests in higher BC concentrations in some regions and lower concentrations in others. Figure 4 compares modeled surface BC concentrations against observations for May–October 2006 at six IMPROVE sites. Model results are from simulations at $2^\circ \times 2.5^\circ$ or $0.5^\circ \times 0.667^\circ$ horizontal resolutions. There are substantial increases of surface BC concentrations (up to $\sim 100\%$) after the emission adjustment, at Bend, MT ($47.6^\circ$ N, 108.7$^\circ$ W, 0.89 km) and North Cheyenne, MT ($45.7^\circ$ N, 106.6$^\circ$ W, 1.28 km) in early and middle summer, while concentrations are appreciably lower (up to $\sim 10\%$ at $2^\circ \times 2.5^\circ$) at Starkey, OR ($45.2^\circ$ N, 118.5$^\circ$ W, 1.26 km) and Mt. Cabinet, MT ($48.0^\circ$ N, 115.7$^\circ$ W, 1.44 km) in September and October. The resulting changes to modeled surface BC concentration are largely consistent between the two resolutions but are considerably more pronounced at $2^\circ \times 2.5^\circ$ than at $0.5^\circ \times 0.667^\circ$ at Starkey and Mt. Cabinet.

Figure 5 shows modeled vs. observed surface BC concentrations averaged for sites at the 0–1, 1–2, 2–3, and 3–4 km altitude ranges for May–October 2006. The adjusted emissions lead to small yet significant relative enhancements of model surface BC concentrations (up to $\sim 0.05$ μg m$^{-3}$) at both model resolutions from late June to August. These enhancements are particularly evident at the 1–2 and 2–3 km
Based on Bayes’ theorem and the assumption of Gaussian uncertainties, the posterior (MAP) solution to Eq. (1) is equivalent to finding the minimum of an error-weighted least squares cost function $J(X)$ (Rodgers, 2000).

$$J(X) = (Y - KX)^T S_X^{-1} (Y - KX) + (X - X_a)^T S_a^{-1} (X - X_a),$$

where $X_a$ and $S_a$ are the a priori emissions and the associated error covariance, $S_X$ is the observational error covariance, and $K = \nabla_X F$ is the Jacobian matrix. Minimization of $J(X)$ yields

$$\hat{X} = X_a + (K^T S_X^{-1} K + S_a^{-1})^{-1} K^T S_X^{-1} (Y - KX_a),$$

$$\hat{S} = (K^T S_X^{-1} K + S_a^{-1})^{-1} = (I - A) S_a (I - A)^T + GS_G G^T,$$

where $\hat{X}$ is the a posteriori emissions, $\hat{S}$ is the a posteriori error covariance, $I$ is the identity matrix, $G = \frac{1}{3} \hat{X}^T$ is the gain matrix (the sensitivity of the retrieval to the observations), and $A = \frac{1}{3} X = G K = I - \hat{S} S_a^{-1}$ is the averaging kernel matrix. In a successful inversion, the cost function should be of the same order as the number of observations, provided that errors are properly specified (Palmer et al., 2003).

The matrix of averaging kernel and the number of degrees of freedom for signal (DOFs) are useful metrics for inspecting the ability of the observing system to uniquely constrain individual element of the state vector (Kasibhatla et al., 2002). The averaging kernel matrix represents the sensitivity of the a posteriori estimates to the unknown true state. Averaging kernels peaking at their own state vector element denote a well constrained source, which shows the inversion system has enough information to constrain the source categories independently. DOFs is the trace of the averaging kernel matrix.

5 Analytical solution to the inverse problem

Analytical inversion methods are applicable for linear or weakly non-linear conditions between emissions and atmospheric concentrations (Müller and Stavrakou, 2005). Previous studies have shown that the GEOS-Chem simulations of BC were linear with respect to BC emissions (e.g., Wang et al. 2011, 2013; Kopacz et al., 2011). Considering the problem of inferring emissions (state vector $X$) given observations (observation vector $Y$) with error $\epsilon$, the two are related via a forward model $F$ as follows (Rodgers, 2000):

$$Y = F(X) + \epsilon,$$

where $X$ is monthly BC emissions (Fig. 1) in the present study, $Y$ the 24-h average BC surface concentrations (Fig. 1 and Supplement Table S1), and $F$ the GEOS-Chem model. Based on Bayes’ theorem and the assumption of Gaussian error distributions (Rodgers, 2000), the optimal or maximum...
We assume a 50% uncertainty for the anthropogenic BC emissions in North America, within the 30–60% range suggested by Bond et al. (2004). GFEDv2 fuel loading has an uncertainty of 22% for woody biomass and 44% for herbaceous biomass, while combustion completeness has an uncertainty of ∼50% (van der Werf et al., 2010). GFEDv2 burned area has an uncertainty of 10–40% for large fires and 50–100% for small fires (Giglio et al., 2006). BC emission factor has an uncertainty of ∼40% for savanna, grasslands, and extra-tropical forest (Andreae and Merlet, 2001). Mao et al. (2011) suggested that GFEDv2 BC emissions were likely biased low by a factor of 2–3 in the WUS. We assume for separate inversions presented here uncertainty of 200, 300, or 500% for biomass burning emissions of BC in the WUS. We assume that the a priori errors are spatially uncorrelated.

The observation error includes contributions from predominantly model transport error, representation error, and measurement error (Heald et al., 2004). We estimate the transport error by computing the variance of the relative difference between the observations and the corresponding model BC concentrations, following Palmer et al. (2003) and Heald et al. (2004). The mean model bias, as diagnosed by the mean relative difference, is a result of errors in the a priori sources, while the variance of the relative residual error is chiefly because of errors in the transport. The resulting model transport error is ∼20%, consistent with previous studies (Kopacz et al., 2009; Heald et al., 2004; Palmer et al., 2003). Jiang et al. (2011) used the differences in GEOS-Chem simulated carbon monoxide distributions with two different GEOS meteorological fields (GEOS-3 vs. GEOS-4) as an alternative measure of model transport error. They found that such transport error introduced an error of up to 20% on the source estimates. The representation error describes the mismatch between model results and the corresponding observations. This error arises because model results are averaged over a large model grid scale (e.g., ∼200 × 250 km² at 2° × 2.5°), while as observations are typically point measurements. Palmer et al. (2003) estimated a representation error (for carbon monoxide) of 5–10% of observations for GEOS-Chem simulations at 2° × 2.5°. Their estimate was based upon aircraft measurements, mostly 1 km over the ocean, and corresponding model simulations over the western Pacific. It is likely that the representation error in the WUS should be larger than 5–10% merely because of the complex terrain, which is more difficult (relative to remote oceans) to resolve in coarse-resolution models. We thus assume a 5–10% representation error as a lower limit in the analyses presented here. We further assume a measurement error of 5–10% for IMPROVE BC measurements. The total observation error is set at 30 or 50%.

Inspection of the averaging kernels and DOFs shows that an a priori biomass burning error of 500% and an observation error of 30% provide the best overall inversion results, with higher resolution in the averaging kernels (Fig. 7) and highest values of DOFs (Table 1). The averaging kernels at the two

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**Figure 6.** Number frequency of model bias for May–October 2006. Model bias is defined as the difference between simulated and observed 24 h average surface BC concentrations (µg m⁻²) at IMPROVE sites. Model results are from simulations at 2° × 2.5° and 0.5° × 0.667° horizontal resolutions and with the standard, the adjusted (see Figs. 2 and 3), and a posteriori emissions. Also shown are the mean, median, standard deviation, and fitted Gaussian distribution curve of the biases.

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kernel matrix (Rodgers, 2000). The number of pieces of information from a perfect inversion system is expected to be the size of the state vector. Following Jaffe et al. (2008), we initially include in the state vector five regions of biomass burning: the northern Rocky Mountains, the southern Rocky Mountains, California, the Southwest, and the Pacific Northwest. Upon examining the averaging kernel and DOFs, we find that the inversion system is unable to distinguish the biomass burning from the northern vs. the southern Rockies. Therefore, we lump these two regions. We also lump California and the Southwest for the same reason. Our analyses show that biomass burning emissions in the WUS account for ∼85% of the total biomass burning contribution to surface BC in the region for May–October 2006. The corresponding contribution from Canadian fire is ∼7%. We thus define a four-component state vector (Fig. 1): biomass burning emissions in the Rockies (BBRM), California and the Southwest (BBCSWM), and the Pacific Northwest (BBPNW), and anthropogenic emissions in the WUS (ANTHWUS).
Figure 7. The averaging kernels for inversions (May–October 2006) of BC emissions in the western US, with each line corresponding to an emission source or source region: biomass burning emissions in the Rockies (BBRM, black line), biomass burning emissions in California and the Southwest (BBCSW, red line), biomass burning emissions in the Pacific Northwest (BBPNW, green line), and anthropogenic emissions in the western US (ANTHWUS, blue line). Results here are from simulations at $2^\circ \times 2.5^\circ$ (solid line) and $0.5^\circ \times 0.667^\circ$ (dotted line) horizontal resolutions and with best set of error characterizations (30 % for observations and 500 % for biomass burning emissions; bottom two rows in Table 1).

Table 1. Number of degrees of freedom for signal (DOFs) for inversions (May–October 2006) using different error characterizations and at $2^\circ \times 2.5^\circ$ and $0.5^\circ \times 0.667^\circ$ horizontal resolutions.

<table>
<thead>
<tr>
<th>Error specification</th>
<th>DOFs</th>
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<tr>
<td></td>
<td>May</td>
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<tr>
<td>$2^\circ \times 2.5^\circ$</td>
<td>S$_\Sigma$ = 50%</td>
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<td>S$_a$BB = 300%</td>
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<td>S$_\Sigma$ = 30%</td>
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<td>S$_\Sigma$ = 30%</td>
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<td></td>
<td>S$_a$BB = 500%</td>
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<tr>
<td>$0.5^\circ \times 0.667^\circ$</td>
<td>S$_\Sigma$ = 30%</td>
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Model resolutions are essentially indistinguishable. Overall the retrievals are slightly better constrained at $0.5^\circ \times 0.667^\circ$ than at $2^\circ \times 2.5^\circ$, as evident in the higher DOFs. Figure 7 shows that the inversion system is able to constrain each element of the state vector independently, especially during July to September and to a lesser degree in October. However, the system has insufficient information to distinguish anthropogenic from biomass burning BC emissions in May and June.

The cost function provides yet another metric for appraising the quality of an inversion. The inversions presented here see large cost function reductions (42 % at $2^\circ \times 2.5^\circ$ and 38 % at $0.5^\circ \times 0.667^\circ$). In addition, the cost function values are of the same order as the number of observations ($\sim 690$). Palmer et al. (2003) showed that the value of the cost function in a successful inversion should be of the same order as the number of observations, provided that the errors are properly specified.
6 Results and discussions

6.1 A posteriori estimates of BC emissions

Figure 8 shows the a priori (i.e., adjusted GFEDv2, see Sect. 4) and the a posteriori estimates of monthly BC emissions in the WUS for May–October 2006. The anthropogenic emissions are divided by 3 in the figures, for the sake of clarity, because the anthropogenic emissions are considerably larger than biomass burning emissions. The a posteriori biomass burning emissions increase (by a factor of 3–5 on average) dramatically and consistently, while the a posteriori anthropogenic emissions decrease (by ∼50%) substantially. The retrievals also largely reduce the uncertainties of the emissions by more than 50%. An a priori biomass burning emissions error of 500% and an observational error of 30% provide the best overall inversion results, with larger reductions in uncertainties (Fig. 8), larger DOFs (Table 1), and better-resolved averaging kernels (Fig. 7). By the same metrics, the inversion results are appreciably better at 0.5° × 0.667° than at 2.5° × 2.5°. As expected, model simulations with the former also result in better agreement with observations.

Table 2 presents the a priori and a posteriori monthly biomass burning and anthropogenic BC emissions in the WUS for May–October 2006. The a posteriori biomass burning BC emissions are 31.7 Gg at 2.5° × 2.5° (4.7 times the a priori) and 19.2 Gg at 0.5° × 0.667° (2.8 times the a priori) for July–September. The a posteriori biomass burning emissions show large regional and month-to-month differences between the two model resolutions. For example, the a posteriori biomass burning emissions at 2.5° × 2.5° for July–September increase by factors of 7.0 in the Rockies and 4.3 in the CSW, relative to the corresponding a priori. The a posteriori biomass burning emissions in the WUS increase by factors of 6.3 (4.1) in July and 3.6 (2.2) in August at 2.5° × 2.5° (0.5° × 0.667°). The a posteriori anthropogenic BC emissions are 9.1 Gg at 2.5° × 2.5° (a 48% reduction relative to the a priori) and 11.2 Gg at 0.5° × 0.667° (a 36% reduction relative to the a priori) for July–September 2006. These decreases in anthropogenic BC emissions estimates are consistent with the findings by Rao and Somers (2010), who showed that BC emissions in the US had declined by ∼30% from 1990 to 2005 and were expected to decline by an additional 80% by 2030. Murphy et al. (2011) have shown that observed BC concentrations from 50 IMPROVE sites decreased by more than 25% on average from 1990 to 2004, reflecting effective anthropogenic BC emission control in the US.

It is useful to compare biomass burning BC emissions in the WUS between the a priori, a posteriori, GFEDv3, and FLAMBE. The total BC emissions for 2006, however, are lower in GFEDv3 than in GFEDv2 (and the a priori emissions) (van der Werf et al., 2010). Here we focus on July–September 2006. The emissions are 8% higher in GFEDv3 than in the a priori. These differences are largest in August (20%) and September 2006 (15%). GFEDv3 emissions are considerably lower than the a posteriori estimates at 2.5° × 2.5° (by a factor of 4.4) and at 0.5° × 0.667° (by a
Table 2. Monthly biomass burning BC emissions from three regions (see Fig. 1) and anthropogenic BC emissions in the western US for May–October 2006 (unit: Gg). A priori and a posteriori biomass burning emissions from inversions at $2^\circ \times 2.5^\circ$ and $0.5^\circ \times 0.667^\circ$ are shown. Also shown are the Global Fire Emissions Database version 3 (GFEDv3) and the Fire Locating and Monitoring of Burning Emissions (FLAMBE) for comparison.

<table>
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<tr>
<th>Source region</th>
<th>Emissions</th>
<th>BC Emissions (Gg), 2006</th>
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<tbody>
<tr>
<td></td>
<td>May</td>
<td>Jun</td>
</tr>
<tr>
<td>BBRM</td>
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<td>$0.5^\circ \times 0.667^\circ$</td>
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<td></td>
<td>FLAMBE</td>
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<td>$2^\circ \times 2.5^\circ$</td>
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<td>$0.5^\circ \times 0.667^\circ$</td>
<td>(0.36)</td>
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</table>

* Uncertainties in parentheses.

The emissions are substantially higher (by a factor of 6) in FLAMBE than in the a priori emissions. We note that the a priori emissions have the same total emissions as the original GFEDv2 emissions and are thus known to underestimate biomass burning BC emissions in the WUS (Randerson et al., 2012; Mao et al., 2011). The difference between FLAMBE and the a posteriori emissions, however, is substantially smaller. The emissions are 26% higher in FLAMBE than in the a posteriori at $2^\circ \times 2.5^\circ$. The difference is a factor of two between FLAMBE and the a posteriori at $0.5^\circ \times 0.667^\circ$. Previous studies have indicated substantial reductions to the FLAMBE inventory for Russia (53%, Wang et al., 2011) and for Southeast Asia (45%, Fisher et al., 2010).

6.2 Evaluation of the a posteriori estimates with IMPROVE observations

Figure 9 shows model simulated and observed surface BC concentrations for May–October 2006 at 10 sites in the Rockies (a–d), the Pacific Northwest (e–g), and California and the Southwest (h–j). Model results are from simulations at $2^\circ \times 2.5^\circ$ and $0.5^\circ \times 0.667^\circ$ and with the a priori and the a posteriori emissions. The a posteriori emissions significantly improve model agreement with observations, not only at individual sites (Fig. 9) but also on average over each of the aforementioned three regions (Fig. 10) and at four altitude ranges (below 1, 1–2, 2–3, and above 3 km) (Fig. 11). The improvements are most pronounced for July–September, the strongest part of the fire season, reflecting the large increases...
and 13 % at 0.5 emissions lead to mean bias reductions of 32 % at 2 dens, AZ (36.1°W, 1.87 km), Three Sisters, OR, and Indian Gardens, AZ (36.1°N, 112.1°W, 1.17 km). The a posteriori emissions lead to mean bias reductions of 32 % at 2° × 2.5° and 13 % at 0.5° × 0.667° for May–October (Fig. 6). The bias reduction is ∼30 % on average in the simulated surface BC concentrations at the 1–2 km altitude range.

Figure 10. Observed (red line) and simulated 24 h average surface BC concentrations (µg m⁻³) averaged at IMPROVE sites in the Rockies (top left panel, 31 sites), in California and the Southwest (top right panel, 28 sites), and in the Pacific Northwest (bottom panel, 10 sites) for May–October 2006. Model results are from simulations at 2° × 2.5° (solid line) and 0.5° × 0.667° (dotted line) horizontal resolutions and with the a priori (black line; see Figs. 2 and 3) and a posteriori (green line) emissions.

in biomass burning emissions in the a posteriori inventory (Sect. 6.1 and Table 2). The improvements are especially evident in the Rockies and to a lesser degree in the Pacific Northwest (Fig. 10), a result of the large increases in the a posteriori biomass burning emissions in these regions. Figure 9 shows that model results with the a posteriori emissions now reproduce the observed synoptic variability in surface BC and capture the major fire episodes, for example, at Craters of the Moon, ID (43.5°N, 113.6°W, 1.82 km), North Cheyenne, MT (45.7°N, 106.6°W, 1.28 km), Jarbridge Wild, NV (41.9°N, 115.4°W, 1.87 km), Three Sisters, OR (44.3°N, 122.0°W, 0.89 km), Starkey, OR, and Indian Gardens, AZ (36.1°N, 112.1°W, 1.17 km). The a posteriori emissions lead to mean bias reductions of 32 % at 2° × 2.5° and 13 % at 0.5° × 0.667° for May–October (Fig. 6). The...

Figure 9. Observed (red line) and simulated 24 h average surface BC concentrations (µg m⁻³) at 10 IMPROVE sites in the Rocky Mountains (a–d), in the Pacific Northwest (e–g), and in California and the Southwest (h–j) for May–October 2006. Model results are from simulations at 2° × 2.5° (solid line) and 0.5° × 0.667° (dotted line) horizontal resolutions and with the a priori (black line; see Figs. 2 and 3) and a posteriori (green line) emissions.
addition, the differences in the simulated surface BC concentrations averaged at those 10 sites are rather small (<7 % at 2° × 2.5° and <4 % at 0.5° × 0.667°) between the two sets (one with 59 sites and the other with 69 sites) of a posteriori BC emissions (Supplement Fig. S1).

6.3 Sensitivity to model resolution

In this section, we compare and contrast the a posteriori emissions and the resulting model surface BC concentrations at both 2° × 2.5° and 0.5° × 0.667° resolutions. GEOS-Chem simulations of surface BC with the a priori and a posteriori emissions are very sensitive to the model horizontal resolution. The 0.5° × 0.667° resolution generally provides better results (compared with the observations) than the 2° × 2.5° resolution does. This is expected because the finer resolution allows for better resolved topography, meteorological fields, transport and emissions, and therefore provides better agreement with observations, as shown in previous modeling studies (e.g., Zhang et al., 2012; Chen et al., 2009; Wang et al., 2004). The surface BC simulations at 0.5° × 0.667° with the a priori emissions show better agreement with the observations compared with those at 2° × 2.5°, as evident in both a lower (by 44 % on average, Fig. 6) model bias and a higher (by 71 %, Fig. 13) Taylor skill score. Inversions at 0.5° × 0.667° also lead to larger reductions in the uncertainties of emissions, larger values of DOFs, and higher-resolution averaging kernels, compared with those at 2° × 2.5°. The resulting simulated surface BC concentrations at 0.5° × 0.667° show improved agreement with observations, too. The mean biases of the simulated surface BC concentrations are 29 % lower at 0.5° × 0.667° than at 2° × 2.5° (Fig. 6) and the corresponding Taylor skill score is higher by 24 % on average (Fig. 13).

The difference in the two a posteriori biomass burning emissions (at the two resolutions) is most evident from July through September 2006. The a posteriori biomass burning emissions are substantially higher (by 65 % in the WUS, 48 % in the RM, and nearly a factor of six in the CSW; Table 2) at 2° × 2.5° than at 0.5° × 0.667°. In contrast, the resulting surface BC concentrations averaged at the IMPROVE sites are higher at the finer resolution and in better agreement with the observations. The larger discrepancies between the coarser-resolution model results and the observations to a large degree reflect the considerably larger smearing out (“dilution”) of emissions at the coarse resolution than at the nested finer resolution (Chen et al., 2009). Consequently, these larger discrepancies (for the coarser-resolution model results) dictate that the inversion system, which seeks to minimize the very differences, imposes larger emissions at the coarser resolution than at the finer resolution.

7 Summary and conclusions

We have applied Bayesian linear inversions to derive top-down estimates of biomass burning and anthropogenic emissions of BC in the WUS for May–October 2006 by inverting surface BC concentrations from the IMPROVE network. We conducted analytical inversions using the GEOS-Chem chemical transport model at both 2° × 2.5° and 0.5° × 0.667° (nested over North America) horizontal resolutions. Model simulated surface BC concentrations with both the a priori and the a posteriori emissions were compared with observations.

We first used MODIS active fire counts to improve the spatiotemporal distributions of the GFEDv2 biomass burning emissions of BC. The adjustment primarily shifted emissions from late to middle and early summer (a 33 % decrease in September-October and a 56 % increase in June–August) and led to significant enhancements in model simulated surface BC concentrations in early and middle summer,
particularly evident at the 1–2 and 2–3 km altitude ranges and at 0.5° × 0.667°. The adjustment also reduced the median model bias by 64% at 0.5° × 0.667° but had little to no effect on the model bias at 2° × 2.5°. The resulting emissions were then used as the a priori for the inversions.

The examination of the averaging kernels and DOFs at both 2° × 2.5° and 0.5° × 0.667° indicated that the inversion system had sufficient information to uniquely constrain the four elements of the state vector and our retrievals were reliable with specified errors, especially for July–September 2006. The inversions led to large reductions in cost functions (42% at 2° × 2.5° and 38% at 0.5° × 0.667°) and the uncertainties of the emissions (>50%). An a priori biomass burning emissions error of 500% and a total observation error of 30% provided the best overall inversion results, with larger reductions in uncertainties, larger values of DOFs, and better-resolved averaging kernels.

The a posteriori biomass burning BC emissions for July–September are 31.7 Gg at 2° × 2.5° (an increase by a factor of 4.7) and 19.2 Gg at 0.5° × 0.667° (an increase by a factor of 2.8), relative to the a priori. The corresponding a posteriori anthropogenic BC emissions were 9.1 Gg at 2° × 2.5° (a decrease of 48%) and 11.2 Gg at 0.5° × 0.667° (a decrease of 36%). The GFEDv3 BC emissions were higher than the a priori estimates (8%) and lower than the a posteriori estimates (by factors of 4.4 at 2° × 2.5° and 2.7 at 0.5° × 0.667°) in the WUS for July–September 2006. FLAMBE emissions were higher than the a priori emissions (by a factor of 5.9) and similar to the a posteriori emissions (factors of 1.3 at 2° × 2.5° and 2.1 at 0.5° × 0.667°) in the WUS during July to September 2006. Model surface BC concentrations with the a posteriori emissions captured the major fire episodes at many IMPROVE sites, especially at the 1–2 and 2–3 km altitude ranges. The a posteriori estimates significantly improved the model agreement with observations, especially at 0.5° × 0.667°, in the Rockies, and for July–September. The a posteriori emissions substantially reduced the bias of the simulated surface BC concentrations (~30% on average) and increased the resulting Taylor skill scores (95% at 2° × 2.5° and 42% at 0.5° × 0.667°).

The inversion results are rather sensitive to the model horizontal resolution. The a posteriori biomass burning emissions showed large regional and month-to-month differences between the two model resolutions. The biomass burning emissions were generally lower at 0.5° × 0.667° than at 2° × 2.5°. The large differences in the two a posteriori emissions suggested that the inversion system likely imposed larger emissions at the coarser resolution than at the finer resolution to minimize the larger differences between the coarser-resolution model results and the observations, because of the larger smearing out of emissions at the coarse resolution.

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References


