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A global 3-D CTM evaluation of black carbon in the Tibetan Plateau

C. He1,2, Q. B. Li1,2, K. N. Liou1,2, J. Zhang3, L. Qi1,2, Y. Mao1,2, M. Gao1,2, Z. Lu4, D. G. Streets4, Q. Zhang5, M. M. Sarin6, and K. Ram7
1Department of Atmospheric and Oceanic Sciences, University of California, Los Angeles, CA, USA
2Joint Institute for Regional Earth System Science and Engineering, University of California, Los Angeles, CA, USA
3Department of Atmospheric and Oceanic Sciences, School of Physics, Peking University, Beijing, China
4Decision and Information Sciences Division, Argonne National Laboratory, Argonne, Illinois, USA
5Center for Earth System Science, Tsinghua University, Beijing, China
6Department of Geosciences, Physical Research Laboratory, Ahmedabad, India
7Institute of Environment and Sustainable Development, Banaras Hindu University, Varanasi, India

Correspondence to: C. He (cenlinhe@atmos.ucla.edu)

Abstract. We systematically evaluate the black carbon (BC) simulations for 2006 over the Tibetan Plateau by a global 3-D chemical transport model (CTM) (GEOS-Chem) driven by GEOS-5 assimilated meteorological fields, using in situ measurements of BC in surface air, BC in snow, and BC absorption aerosol optical depth (AAOD). Using improved anthropogenic BC emission inventories for Asia that account for rapid technology renewal and energy consumption growth (Zhang et al., 2009; Lu et al., 2011) and improved global biomass burning emission inventories that account for small fires (van der Werf et al., 2010; Randerson et al., 2012), we find that model results of both BC in surface air and in snow are statistically in good agreement with observations (biases <15%) away from urban centers. Model results capture the seasonal variations of the surface BC concentrations at rural sites in the Indo-Gangetic Plain, but the observed elevated values in winter are absent. Modeled surface-BC concentrations are within a factor of 2 of the observations at remote sites. Part of the discrepancy is explained by the deficiencies of the meteorological fields over the complex Tibetan terrain. We find that BC concentrations in snow computed from modeled BC deposition and GEOS-5 precipitation are spatiotemporally consistent with observations ($r = 0.85$). The computed BC concentrations in snow are a factor of 2–4 higher than the observations at several Himalayan sites because of excessive BC deposition. The BC concentrations in snow are biased low by a factor of 2 in the central plateau, which we attribute to the absence of snow aging in the CTM and strong local emissions unaccounted for in the emission inventories. Modeled BC AAOD is more than a factor of 2 lower than observations at most sites, particularly to the northwest of the plateau and along the southern slopes of the Himalayas in winter and spring, which is attributable in large part to underestimated emissions and the assumption of external mixing of BC aerosols in the model. We find that assuming a 50% increase of BC absorption associated with internal mixing reduces the bias in modeled BC AAOD by 57% in the Indo-Gangetic Plain and the northeastern plateau and to the northeast of the plateau, and by 16% along the southern slopes of the Himalayas and to the northwest of the plateau. Both surface BC concentration and AAOD are strongly sensitive to anthropogenic emissions (from China and India), while BC concentration in snow is especially responsive to the treatment of BC aerosol aging. We find that a finer model resolution ($0.5^\circ \times 0.667^\circ$ nested over Asia) reduces the bias in modeled surface-BC concentration from 15 to 2%. The large range and non-homogeneity of discrepancies between model results and observations of BC across the Tibetan Plateau undoubtedly undermine current assessments of the climatic and hydrological impact of BC in the region and thus warrant imperative needs for more extensive measurements of BC, including its concentration in surface air and snow, AAOD, vertical profile and deposition.

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1 Introduction

Black carbon (BC) is the most important light-absorbing aerosol formed during incomplete combustion (Bond et al., 2013, and references therein), with major sources from fossil fuel and biofuel combustion and open biomass burning (Bond et al., 2004). BC warms the atmosphere by strongly absorbing solar radiation in the visible and the near-infrared (Ramanathan and Carmichael, 2008), influences cloud formation as cloud condensation nuclei (Jacobson, 2006), and accelerates snow and ice melting by significantly reducing snow and ice albedo (i.e., the snow-albedo effect) (Hansen and Nazarenko, 2004; Flanner et al., 2007). With an estimated global climate forcing of $+1.1 \text{ W m}^{-2}$, BC is now considered the second most important human emission in terms of its climate forcing in the present-day atmosphere after carbon dioxide (Ramanathan and Carmichael, 2008; Bond et al., 2013). The regional warming effect of BC can be even stronger, particularly over snow-covered regions (Jacobson, 2004; Flanner et al., 2007, 2009). There is ample evidence that BC aerosols deposited on Tibetan glaciers have been a significant contributing factor to observed rapid glacier retreat in the region (e.g., Xu et al., 2009). It has also been proposed that the radiative forcing from ever-increasing deposition of BC in snow was an important cause for the retreat of Alpine glaciers from the last Little Ice Age through the mid-19th century (Painter et al., 2013).

The Tibetan Plateau is the highest plateau in the world with the largest snow and ice mass outside the polar regions (Xu et al., 2009). The Tibetan glaciers and the associated snowmelt are the primary source of freshwater supply for drinking, agricultural irrigation, and hydropower for more than 1 billion people in Asia (Immerzeel et al., 2010). The plateau also plays a critical role in regulating the Asian hydrological cycle. Changes of snow cover affect heat flux and water exchange between the surface and the atmosphere, and further disturb the formation of the Asian monsoon (Lau and Kim, 2006).

Observations have shown remarkable warming and accelerated glacier retreat in the Tibetan Plateau in the past decades (Qin et al., 2006; Prasad et al., 2009). Ramanathan et al. (2005, 2007) argued that the ever-increasing amount of BC transported to the Himalayas accounts for half of the observed warming in the region, comparable to the warming attributable to greenhouse gases (Barnett et al., 2005). Recent studies reaffirmed a strong BC-induced regional warming over the plateau that results in more than 1% decrease of snow/ice cover (Lau et al., 2010; Menon et al., 2010), 2–5% reduction of snow albedo (Yasumari et al., 2010), and an increase of runoff in early spring (Qian et al., 2011). Surrounded by the world’s two largest BC source regions, south and east Asia (Lamarque et al., 2010), the plateau has received an increasing BC deposition from 1951 to 2000, particularly after 1990 (Ming et al., 2008). Recent studies have shown that the amount of BC transported to the plateau has increased by 41% from 1996 to 2010, with south and east Asia accounting for 67% and for 17% on an annual basis (Lu et al., 2012). The modeling study by Kopacz et al. (2011) suggested that long-range transport from Middle East, Europe, and northern Africa also contributes to the BC deposition over the plateau.

The climatic effects of BC over the Tibetan Plateau are not well understood, with large uncertainties in the estimates of BC radiative forcing (e.g., Flanner et al., 2007; Kopacz et al., 2011; Ming et al., 2013). Accurate assessment of BC-related radiative forcing in the Tibetan Plateau critically depends on reliable model simulations of BC emissions, transport and subsequent deposition, and vertical distribution over the plateau. Previous modeling studies have found invariably large discrepancies with observations. For example, the simulations of surface BC at several sites in the southern slope of the Himalayas are biased low by more than a factor of 2, particularly in winter and spring in regional, multi-scale and global models (Nair et al., 2012; Moorby et al., 2013). Fu et al. (2012) showed that surface-BC concentrations simulated by a global chemical transport model (CTM) are more than 50% lower than observations in China in general and across the Tibetan Plateau in particular. A global CTM study (Kopacz et al., 2011) and a global climate model (GCM) study (Qian et al., 2011) both showed large differences between modeled and observed BC concentration in snow over the plateau. Sato et al. (2003) and Bond et al. (2013) pointed out large underestimates of BC absorption aerosol optical depth (AAOD) in previous models compared with Aerosol Robotic Network (AERONET) retrievals.

In this study we seek to understand the capability of a global 3-dimensional CTM (GEOS-Chem) in simulating BC in the Tibetan Plateau and the associated discrepancies between model results and observations. The GEOS-Chem model has been widely used in previous studies to understand BC emissions, transport and deposition in the plateau (Kopacz et al., 2011), in China (Fu et al., 2012), over Asia (Park et al., 2005), in the Arctic (Wang et al., 2011) and globally (Wang et al., 2014). To our knowledge, this is the first attempt to systematically evaluate a global simulation of BC in the Tibetan Plateau using all three types of available in situ measurements: BC in surface air, BC in snow, and BC AAOD. We further delineate the effects of anthropogenic BC emissions from China and India, BC aging process and model resolution on the simulation. Potential factors driving model-versus-observation discrepancies are also examined, which gives implications for improving the estimate of BC climatic effects. Observations and model description are presented in Sect. 2. Simulations of surface BC, BC in snow and BC AAOD are discussed in Sect. 3. Sensitivity and uncertainty analyses are in Sects. 4, 5 and 6. Finally, summary and conclusions are given in Sect. 7.
2 Method

2.1 Observations

For the sake of clarity, we define here the Tibetan Plateau roughly as the region in 28–40° N latitudes and 75–105° E longitudes. We also define several sub-regions of the plateau and adjacent regions (Fig. 1): the central plateau (30–36° N, 82–95° E), the northeastern plateau (34–40° N, 95–105° E), the southeastern plateau (28–34° N, 95–105° E), to the north of the plateau (40–50° N, 85–95° E), to the northwest of the plateau (40–50° N, 70–85° E), to the northeast of the plateau (40–50° N, 95–105° E), and the Himalayas. There are rather limited measurements of BC in the Tibetan Plateau. Figure 1 shows sites with measurements of BC surface concentration, concentration in snow, and AAOD in the region.

2.1.1 BC surface concentration

There are 13 sites with monthly or seasonal measurements of surface BC concentration (Table 1 and Fig. 1). Observations are available for 2006 at nine of the sites. Four sites provide observations for 1999–2000, 2004–2005 or 2008–2009. We distinguish these sites as urban, rural, or remote sites based upon annual mean surface-BC concentration, following Zhang et al. (2008). The concentration is typically higher than 5 µg m⁻³ at urban sites (within urban centers or near strong local residential and vehicular emissions), in the range of 2–5 µg m⁻³ at rural sites, and less than 2 µg m⁻³ at more remote, pristine sites.

Ganguly et al. (2009b) retrieved surface BC concentration at Gandhi College (25.9° N, 84.1° E; 158 m a.s.l.) by combining aerosol optical properties from AERONET measurements and aerosol extinction profiles from Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) observations. The retrieval is rather sensitive to errors in the aerosol single scattering albedo, size distribution and vertical profiles derived from the observations (Ganguly et al., 2009a). Measurements at Delhi (28.6° N, 77.2° E; 260 m a.s.l.), Digharg (27.3° N, 94.6° E; 111 m a.s.l.), Kharagpur (22.5° N, 87.5° E; 22 m a.s.l.) and Nepal Climate Observatory at Pyramid (NCP, 28.0° N, 86.8° E; 5079 m a.s.l.) used an aethalometer (Beegum et al., 2009; Pathak et al., 2010) or a multi-angle absorption photometer (Bonasoni et al., 2010; Nair et al., 2012). The uncertainties of these measurements stem mainly from the interference from other components in the aerosol samples (Bond et al., 1999; Petzold and Schonlinner, 2004) and the shadowing effects under high filter loads (Weingartner et al., 2003). BC concentrations at the other sites were derived from measurements of the thermal–optical reflectance or thermal–optical transmittance (Carrico et al., 2003; Qu et al., 2008; Zhang et al., 2008; Ming et al., 2010; Ram et al., 2010a,b). These measurements are strongly influenced by the temperature chosen to separate BC and organic carbon (OC) (Schmid et al., 2001; Chow et al., 2004).

2.1.2 BC concentration in snow

There are 16 sites with monthly or seasonal measurements of BC concentration in snow during 1999–2007 and two with annual measurements (Xu et al., 2006, 2009; Ming et al., 2009a, b, 2012, 2013). These sites are at high-elevation (> 3500 m a.s.l.), remote locations in the Himalayas and other parts of the plateau (Table 2 and Fig. 1). The snow and ice samples taken from these sites were heated and filtered through fiber filters in the laboratory. Thermal techniques (Cachier and Pertuisot, 1994; Chow et al., 2004) were then used to isolate BC from other constituents (especially OC) in the filters, followed by analysis using carbon analyzers including heating-gas chromatography (Xu et al., 2006), optical carbon analysis (Chow et al., 2004) and coulometric titration-based analysis (Cachier and Pertuisot, 1994). The
The absorption by fine-mode aerosols is primarily from BC while the absorption by larger particles (diameter >1 µm) is principally from dust. Dust AAOD is estimated from the super-micron part of aerosol size distribution provided by the AERONET retrieval method and a refractive index of 1.55 + 0.0015i (Bond et al., 2013). BC AAOD is then the difference between the total and dust AAOD. This process attributes all fine-mode aerosol absorption to BC. Because of the contributions from OC and fine dust particles to fine-mode AAOD, the inferred BC AAOD is likely biased high. Bond et al. (2013) estimated that the uncertainty from the impact of dust and OC on the fine-mode AAOD could be as large as 40–50%. The limited AERONET sampling in this region is another source of uncertainty (Bond et al., 2013).

### 2.1.4 Model description and simulations

The GEOS-Chem model is driven by assimilated meteorology from the Goddard Earth Observing System (GEOS) of the NASA Global Modeling and Assimilation Office (GMAO). We use here GEOS-Chem version 9-01-03 (available at http://geos-chem.org), driven by GEOS-5 data assimilation system (DAS) meteorological fields. The meteorological fields have a native horizontal resolution of 0.5° × 0.667°, 72 vertical layers, and a temporal resolution of 6 h (3 h for surface variables and mixing depths). The spatial resolution is degraded to 2° × 2.5° in the horizontal and 47 layers in the vertical (from the surface to 0.01 hPa) for computational expediency. The lowest model levels are centered at approximately 60, 200, 300, 450, 600, 700, 850, 1000, 1150, 1300, 1450, 1600, and 1800 m a.s.l.

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...
Table 2. Observed and simulated BC concentrations in snow over the Tibetan Plateau (see also Fig. 1).

| Region            | Site              | Lat. (° N) | Long. (° E) | Elev. (km) | Time        | BC in snow (µg kg⁻¹) | Obs.ᵃ | Exp. Aᵇ | Exp. Bᶜ | Exp. Cᵈ | Exp. Dᵉ |
|-------------------|-------------------|------------|-------------|------------|-------------|----------------------|-------|--------|--------|--------|--------|
| The Himalayas     | Zuoqiu   | 29.21      | 96.92       | 5.50       | monsoon 2006 | 7.9[²]            | 22.5 (13.6–43.0) | 18.4 | 25.5   | 24.6   |
|                   | Qiangyang | 28.83      | 90.25       | 5.40       | summer 2001  | 43.1[¹]           | 66.1 (42.2–122.8) | 49.7 | 63.7   | 18.3   |
|                   | Noijn Kangsang  | 29.04      | 90.20       | 5.95       | annual 2005  | 30.6[²]           | 39.5 (21.1–60.8) | 34.6 | 52.3   | 22.5   |
|                   | East Rongbuk    | 28.02      | 86.96       | 6.50       | monsoon 2001 | 35.0[³]           | 26.4 (16.8–48.9) | 22.7 | 29.2   | 22.4   |
|                   | Noijn Kangsang  | 28.02      | 86.96       | 6.50       | non-monsoon 2001 | 21.0[³]            | 32.8 (21.6–55.6) | 31.1 | 59.4   | 42.7   |
|                   | East Rongbuk    | 28.02      | 86.96       | 6.50       | summer 2002  | 20.3[⁴]           | 26.5 (16.8–49.3) | 22.8 | 28.9   | 23.0   |
|                   | Kangwure        | 28.47      | 85.82       | 6.00       | summer 2001  | 21.8[¹]           | 26.5 (16.8–49.3) | 22.8 | 28.9   | 18.0   |
|                   | Namunani        | 30.45      | 81.27       | 5.90       | summer 2004  | 4.3[¹]            | 24.8 (15.8–45.2) | 21.2 | 26.6   | 19.6   |
| Northwestern      | Mt. Muztagh     | 38.28      | 75.02       | 6.35       | summer 2001  | 37.2[¹]           | 31.0 (23.3–52.8) | 36.6 | 31.9   | 32.9   |
| Tibetan Plateau   | 38.28          | 75.10      | 6.30        | 1999       |             | 26.6[¹]           | 33.0 (26.6–48.6) | 36.4 | 45.8   | 42.1   |
| Northeastern      | Laohugou #12    | 39.43      | 96.56       | 5.05       | Oct. 2005    | 35.0[⁴]           | 54.4 (34.9–97.2) | 60.0 | 60.3   | 65.0   |
| Tibetan Plateau   | Qiyi             | 39.23      | 97.06       | 4.85       | Jul. 2005    | 22.0[⁴]           | 25.7 (18.5–67.0) | 30.3 | 27.4   | 48.9   |
|                   | 1 July glacier  | 39.23      | 97.75       | 4.60       | summer 2001  | 52.6[¹]           | 59.2 (32.8–122.8) | 68.8 | 61.0   | 106.2  |
| Central           | Meikiang         | 35.67      | 94.18       | 5.20       | summer 2001  | 44.0[¹]           | 24.4 (15.4–47.0) | 24.8 | 27.2   | 32.9   |
| Tibetan Plateau   | 35.67          | 94.18      | 5.20        | Nov. 2005  |             | 81.0[⁵]           | 40.9 (18.8–50.0) | 43.3 | 50.5   | 38.6   |
|                   | Tanglela        | 33.11      | 92.09       | 5.80       | 2003        | 53.1[²]           | 16.1 (10.4–28.6) | 14.5 | 25.6   | 12.0   |
|                   | Dongkemadi      | 33.10      | 92.08       | 5.60       | summer 2001  | 18.2[¹]           | 19.6 (12.2–36.8) | 17.7 | 22.1   | 15.0   |
|                   | 33.10          | 92.08      | 5.60        | year 2005  |             | 36.6[⁷]           | 15.8 (10.2–28.1) | 14.2 | 23.7   | 11.8   |
|                   | La’ngong        | 30.47      | 90.57       | 5.85       | Jun. 2005    | 67.0[⁴]           | 39.1 (25.8–72.9) | 35.9 | 37.8   | 22.9   |
|                   | Zhadang         | 30.47      | 90.50       | 5.80       | Jul. 2006    | 87.4[⁴]           | 27.9 (17.0–53.4) | 21.7 | 30.3   | 19.3   |
| North of the       | Haxilegen River | 43.73      | 84.46       | 3.76       | Oct. 2006    | 46.9[⁴]           | 36.1 (34.1–45.2) | 37.9 | 36.7   | 71.4   |
| plateau            | Urumqi Riverhead| 43.10      | 86.82       | 4.05       | Nov. 2006    | 141[⁵]            | 131.9 (71.8–270.4)| 155.2 | 118.4  | 127.9  |
|                   | Miao’ergou #3   | 43.06      | 94.32       | 4.51       | Aug. 2005    | 111[⁴]            | 98.8 (59.3–158.2) | 113.2 | 103.7  | 113.8  |

ᵃ References: [¹]Xu et al. (2006), [²]Xu et al. (2009), [³]Ming et al. (2008), [⁴]Ming et al. (2009a), [⁵]Ming et al. (2009b), [⁶]Ming et al. (2012), [⁷]Ming et al. (2013).
ᵇ Values from Experiment A (Table 5) for 2006. See text for details. Values in parentheses are the same Experiment but using instead the upper and lower bounds of anthropogenic BC emissions in China and India.
ᶜ Values from experiment B (Table 5) for 2006. See text for details.
ᵈ Values from experiment C (Table 5) for 2006. See text for details.
ᵉ Values from experiment D (Table 5) for 2006. See text for details.

2.1.5 BC emissions

The global anthropogenic BC emissions are from Bond et al. (2007), with an annual emission of 4.4 Tg C for the year 2000. Anthropogenic BC emissions in Asia, chiefly in China and India, have increased significantly since 2000 (Griener et al., 2011). Zhang et al. (2009) developed an Asian anthropogenic BC emissions (for China and India and the rest of Asia) for 2006 for the Intercontinental Chemical Transport Experiment-B (INTEX-B) field campaign (Singh et al., 2009), with considerable updates to a previous inventory developed by Streets et al. (2003). They employed a dynamic methodology that accounts for rapid technology renewal and updated the fuel consumption data. Fu et al. (2012) pointed out that Zhang et al. (2009) underestimate anthropogenic BC emissions in China by a factor of 1.6 compared with the top-down estimates. Lu et al. (2011) further updated the activity rates, technology penetration data and emission factors in China and India, and reported anthropogenic BC emissions only in these two countries for 1996–2010. Table 4 is a summary of the two inventories. Anthropogenic BC emissions in India are lower in the Zhang et al. (2009) inventory (hereinafter the INTEX-B inventory) than in the Lu et al. (2011) inventory (hereinafter the LU inventory) by a factor of 2, while emissions in China are 10% higher in the INTEX-B inventory than in the LU inventory (Table 4). The higher emissions in India in the LU inventory are primarily a result of the updated biofuel emission factors and the new method used to estimate biofuel consumptions. The biofuel emissions, which

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Table 3. Observed and simulated annual mean BC AAOD at AERONET sites over the Tibetan Plateau (see also Fig. 1).

| Region                        | Site             | Lat. (°N) | Long. (°E) | Alt. (m) | Time        | Obs. \(^a\)  | Model\(^b\) | Ratio\(^c\) |
|-------------------------------|------------------|-----------|------------|----------|-------------|--------------|-------------|-------------|
| The Indo-Gangetic Plain      | Lahore           | 31.54     | 74.33      | 270      | 2007–2012   | 0.0434       | 0.0162      | 2.7         |
|                              | Kanpur           | 26.51     | 80.23      | 123      | 2006–2012   | 0.0426       | 0.0221      | 1.9         |
|                              | Gandhi College   | 25.87     | 84.13      | 60       | 2006–2012   | 0.0443       | 0.0282      | 1.6         |
|                              | Gual_Pahari      | 28.43     | 77.15      | 384      | 2008–2010   | 0.0511       | 0.0222      | 2.3         |
|                              | Jaipur           | 26.91     | 75.81      | 450      | 2009–2012   | 0.0202       | 0.0168      | 1.2         |
| The Himalayas                | Jomsom           | 28.78     | 83.71      | 2803     | 2012        | 0.0231       | 0.0136      | 1.7         |
|                              | Pantnagar         | 29.05     | 79.52      | 241      | 2008–2009   | 0.0507       | 0.0114      | 4.4         |
|                              | Nainital          | 29.36     | 79.46      | 1939     | 2008–2010   | 0.0204       | 0.0125      | 1.6         |
|                              | Pokhara           | 28.15     | 83.97      | 807      | 2010–2012   | 0.0524       | 0.0056      | 9.4         |
|                              | Kathmandu Univ.  | 27.60     | 85.54      | 1510     | 2009–2010   | 0.0406       | 0.0057      | 7.1         |
| Northeastern Tibetan Plateau| SACOL            | 35.95     | 104.14     | 1965     | 2007–2011   | 0.0163       | 0.0100      | 1.6         |
| Northeast of the Tibetan Plateau| Dalanzadgad   | 43.58     | 104.42     | 1470     | 2006, 2012  | 0.0038       | 0.0025      | 1.5         |
| Northwest of the Tibetan Plateau| Issyk-Kul     | 42.62     | 76.98      | 1650     | 2008–2010   | 0.0196       | 0.0020      | 9.8         |
|                              | Dushanbe         | 38.55     | 68.86      | 821      | 2011–2012   | 0.0131       | 0.0030      | 4.4         |

\(^a\) AERONET retrieved BC AAOD (Bond et al., 2013). Values are multi-year averages.
\(^b\) Values from Experiment A (Table 5) for 2006. See text for details. Values in parentheses are from the same experiment but use the upper and lower bounds of anthropogenic BC emissions in China and India instead.
\(^c\) The ratio of AERONET retrieved to GEOS-Chem modeled BC AAOD.

Table 4. Anthropogenic BC emissions in China and India in 2006.

| Emissions (Gg yr\(^{-1}\)) | China          | India         |
|---------------------------|----------------|---------------|
| Lu et al. (2011)          | Zhang et al. (2009) | Lu et al. (2011) | Zhang et al. (2009) |
| Industry                  | 509            | 575           | 201          | 47           |
| Power plants              | 15             | 36            | 1            | 8            |
| Residential               | 971            | 1022          | 608          | 268          |
| Transportation            | 178            | 205           | 75           | 80           |
| Total                     | 1673 (954–3229\(^a\)) | 1838 (884–3823) | 885 (522–1655) | 404 (112–1454) |

\(^a\) Uncertainties (in parentheses).
are dominated by residential burning, account for more than 50 % of total BC emissions in India (Lu et al., 2011). There are large uncertainties in both inventories. Lu et al. (2011) used a Monte Carlo method to show that the 95 % uncertainty ranges of BC emissions are from −43 to 93 % for China and from −41 to 87 % for India. The uncertainties in the INTEX-B inventory are ±208 % for China and ±360 % for India (Zhang et al., 2009). A recent study by Qin and Xie (2012) showed slightly (5–10 %) lower total anthropogenic BC emissions in China than those from Lu et al. (2011) for 2006, but the emissions in the northeastern and northwestern China were higher by a factor of 2. Kurokawa et al. (2013) further updated BC emissions in Asia and found 10 % lower anthropogenic BC emissions for China and 30 % lower for India compared with those from Lu et al. (2011), yet with a similar spatial distribution. Wang et al. (2014) developed a new global BC emission inventory, where anthropogenic BC emissions are 20 % higher in China and 30 % lower in India than those from Lu et al. (2011). They found that the use of the new inventory reduces model biases of surface BC concentrations in Asia by 15–20 %. However, the abovementioned three latest inventories are still associated with large uncertainties (95 % confidence intervals), which are more than 100 % for anthropogenic BC emissions in China and India (Qin and Xie, 2012; Kurokawa et al., 2013; Wang et al., 2014).

Global biomass burning emissions are from the Global Fire Emissions Database version 3 (GFEDv3) (van der Werf et al., 2010). Kaiser et al. (2012) showed that GFEDv3 underestimates carbon emissions by a factor of 2–4 globally because of undetected small fires. Randerson et al. (2012) reported an updated GFEDv3 inventory that accounts for small fire emissions. Small fires increase carbon emissions by 50 % in southeast Asia and equatorial Asia (Randerson et al., 2012). We use the GFEDv3 emissions with a monthly temporal resolution in the present study. The uncertainty of the GFEDv3 emissions is at least 20 % globally and higher in boreal regions and equatorial Asia (van der Werf et al., 2010). The major uncertainty lies in insufficient data on burned area, fuel load and emission factor (van der Werf et al., 2010; Randerson et al., 2012).

2.1.6 BC deposition

Simulation of aerosol dry and wet deposition follows Liu et al. (2001). Dry deposition of aerosols uses a resistance-in-series model (Walcek et al., 1986) dependent on local surface type and meteorological conditions. There have been many updates. A standard resistance-in-series scheme (Weck, 1989) has been implemented in the non-snow/non-ice regions (Wang et al., 1998) with a constant aerosol dry deposition velocity of 0.03 cm s\(^{-1}\) prescribed over snow and ice (Wang et al., 2011). This velocity is within the range (0.01–0.07 cm s\(^{-1}\)) employed in Liu et al. (2011) to improve the BC simulation in the Geophysical Fluid Dynamics Laboratory (GFDL) Atmospheric Model version 3 (AM3) global model (Donner et al., 2011). We found that dry deposition accounts for 20 % of the total BC deposition over the Tibetan Plateau in winter and 10 % in summer.

Liu et al. (2001) described the wet scavenging scheme for aerosols in the GEOS-Chem. Wang et al. (2011) implemented in the model a new below-cloud scavenging parameterization for individual aerosol mode, which distinguishes between the removal by snow and by rain drops for aerosol washout. They also applied different in-cloud scavenging schemes to cold and to warm clouds, and with an improved areal fraction of a model grid box that experiences precipitation. These changes are included in the GEOS-Chem version used for the present study.

The GEOS-Chem model does not directly predict BC (or any aerosols for that matter) in snow at the surface in the absence of a land–surface model that explicitly treats snow including its aging. As an approximation, we estimate BC concentration in snow in the model as the ratio of total BC deposition to total precipitation, following Kopacz et al. (2011) and Wang et al. (2011). Although the use of total precipitation here is reasonable considering the low temperature typical over the Tibetan Plateau (Wu and Liu, 2004), it introduces uncertainties to the calculation of snow BC concentration. Bonasoni et al. (2010) found that precipitation can be partly in the form of rain even at altitudes of 5 km in the Himalayas. Thus, the use of total precipitation may overestimate both snow precipitation and BC removed by snow. Besides, rain also results in the melting of snowpack (Marks et al., 2001), which further affects BC concentration in snow. Additional uncertainties exist in the GEOS-5 precipitation fields because of the coarse model resolution and the complex topography in the plateau (see Sect. 3.2). Ménéguz et al. (2013) showed that using a higher resolution model (0.2\(^\circ\) x 0.2\(^\circ\)) improves the simulation of the spatial variability of precipitation in the Himalayas, but the bias in total precipitation remains high. The uncertainty in precipitation is thus propagated to the BC concentration in snow computed from model results. Our calculation of BC concentration in snow assumes a well mixing of BC and snow. However, BC content is not uniform throughout a snow column. Thus, an ideal comparison of modeled and observed BC concentrations in snow should be for the same depth of a snow column. We also neglect the aging of surface snow and the internal mixing of snow and BC, which conceivably contribute to the underestimation of BC concentration in snow computed here. This may be an especially important issue for comparisons in the central plateau and to the north of the plateau, where snowmelt has been suggested to strongly increase BC concentration in snow (Zhou et al., 2007; Ming et al., 2013).
2.1.7 BC aging

Freshly emitted BC is mostly (80 %) hydrophobic (Cooke et al., 1999). Hydrophobic BC becomes hydrophilic typically on the timescale of a few days (McMeeking et al., 2011, and references therein) because of coating by soluble materials like sulfate and organic matter (Friedman et al., 2009; Khalizov et al., 2009). The internal mixing of BC and other aerosol constituents significantly changes the morphology, hygroscopicity and optical properties of BC particles (Zhang et al., 2008). This further influences BC absorption efficiency (Bond et al., 2006) and lifetime against deposition (Mikhailov et al., 2001). However, the aging process is not explicitly simulated in the GEOS-Chem, where an e-folding time of 1.15 days for the conversion of hydrophobic to hydrophilic BC is simply assumed (Park et al., 2005; Kopacz et al., 2011; Wang et al., 2011). Liu et al. (2011) proposed a condensation–coagulation parameterization for BC aging where the conversion time is not uniform but varies. Specifically, the conversion is assumed to be primarily a result of sulfuric acid deposition (condensation) onto BC particles, and the mass deposition rate is proportional to the concentration of gaseous sulfuric acid and to the BC particle surface area. Gaseous atmospheric sulfuric acid is a product of sulfurous acid oxidation by the hydroxyl radical (OH). Consequently its steady-state concentration is linearly linked to OH concentration. Thus, in the absence of nucleation, which is a slow process when there exists plenty of primary particles as found in urban and biomass burning plumes (Seinfeld and Pandis, 2006), the BC aging rate can be parameterized as a linear function of OH concentration, where the coefficient of OH concentration controls a fast aging process (i.e., condensation) and the constant term governs a slow aging process (e.g., coagulation). Huang et al. (2013) further combined the Liu et al. (2011) parameterization with a chemical oxidation aging mechanism from chamber study results (Poschl et al., 2001) in GEOS-Chem. They found that the chemical aging effects on surface BC concentrations are strongest in the tropical regions but negligible over the Tibetan Plateau.

2.1.8 Model simulations

For the present study, we conducted four GEOS-Chem simulations for 2006 (Table 5). Detailed discussions and justifications for these model experiments are provided below where appropriate. Model results are sampled at the corresponding locations of the measurement sites. Model results presented here are monthly averages. As pointed out in previous studies (Fairlie et al., 2007; Mao et al., 2011), comparing localized observations with model results that are representative of a much larger area is inherently problematic. The mountainous sites and the complex terrain in the Tibetan Plateau further complicate the comparison.

In Experiment A, we replace the Bond et al. (2007) emissions in China and India with the LU inventory and use the INTEX-B inventory for the rest of Asia. This is our standard simulation, and the results are used for all model evaluations presented here unless stated otherwise. We also provide the model results from Experiment A but instead using the lower and upper bounds of anthropogenic BC emissions from China and India estimated by Lu et al. (2011). We find that wet deposition accounts for 83 % of the global annual BC deposition, consistent with the previous results of 78.6 ± 17 % from the Aerosol inter-Comparison project (AeroCom) multi-model study (Textor et al., 2006). The tropospheric lifetime of BC against deposition is 5.5 days, at the lower end of the range (5–11 days) reported by Koch et al. (2009). The difference between Experiments B and A is that we replace the Bond et al. (2007) emissions in China and India with the INTEX-B inventory in Experiment B. By contrasting model results from these two experiments, we aim to assess the sensitivity of BC in the Tibetan Plateau to changes in the anthropogenic emissions from India and China, the two largest source regions of BC to the plateau (Kopacz et al., 2011; Lu et al., 2012), as will be discussed in further details in Sect. 4. Both Experiments A and B use an e-folding time of 1.15 days for BC aging. Experiment C applies the Liu et al. (2011) parameterization for BC aging instead. We used monthly mean OH concentrations with diurnal variations in the parameterization, which is derived from the offline GEOS-chem simulation with the same spatial resolution as BC simulations. The resulting e-folding time is 2.5 days on average globally and 2 days in Asia. The longer e-folding time results in longer atmospheric lifetime, larger deposition and higher hydrophobic fraction of BC over the Tibetan Plateau (not shown). We discuss further in Sect. 5 the differing results between Experiments C and A, which allow us to appraise the effect of a variable BC aging time on BC in the plateau. In Experiment D, we replace the model resolution of 2°×2.5° used in Experiment A with a finer resolution of 0.5°×0.667° nested over Asia (11° S–55° N, 70–150° E). The differences between the results from Experiments D and A will be discussed in Sect. 6 for the purpose of evaluating the impact of model resolution on BC in the plateau. In all model experiments, we use a BC mass absorption cross section (MAC) of 7 m² g⁻¹ from observations (Clarke et al., 2004) for calculating modeled BC AAOD. We note that BC MAC is associated with large uncertainties, which varies from 3 to 25 m² g⁻¹ depending on atmospheric conditions (Bond and Bergstrom, 2006).
Table 5. GEOS-Chem simulations of BC.

| Model experiment | A         | B         | C         | D         |
|------------------|-----------|-----------|-----------|-----------|
| Resolution       | $2^\circ \times 2.5^\circ$ | $2^\circ \times 2.5^\circ$ | $2^\circ \times 2.5^\circ$ | $0.5^\circ \times 0.667^\circ$ (Asia) $2^\circ \times 2.5^\circ$ (global) |
| Anthropogenic emissions | China & India | Rest of Asia | Rest of Asia | Rest of world |
|                   | Lu et al. (2011) | Zhang et al. (2009) | Lu et al. (2011) | Lu et al. (2011) |
| Biomass burning emissions | GFEDv3 (van der Werf et al., 2010), with updates from Randerson et al. (2012) |
| BC aging (hydrophobic to hydrophilic) | e-folding time | e-folding time | Liu et al. (2011) | e-folding time |
|                   | 1.15 days | 1.15 days | 1.15 days | 1.15 days |
| Deposition        | Dry deposition | Wet deposition | Wesely (1989) as implemented by Wang et al. (1998) | Liu et al. (2001) with updates from Wang et al. (2011) |

Figure 2. Observed (black curve) and GEOS-Chem simulated (colored curves: red – Experiment A; blue – Experiment B; green – Experiment C; yellow – Experiment D; grey dashed curves – Experiment A using upper/lower bounds of anthropogenic BC emissions in China and India) monthly mean surface-BC concentration ($\mu g m^{-3}$) at three rural sites (a–c) and six remote sites (d–i) over the Tibetan Plateau in 2006 (see Table 1 and Fig. 1). Only seasonal mean observations are available at sites e and f. Also shown are standard deviations for observations (error bars). See text for details.
3 Results

3.1 BC in surface air

Figure 2 shows surface BC concentrations at three rural sites: Kharagpur (22.5° N, 87.5° E; 28 m a.s.l.; Fig. 2a), Gandhi College (25.9° N, 84.1° E; 158 m a.s.l.; Fig. 2b) and Kanpur (26.4° N, 80.3° E; 142 m a.s.l.; Fig. 2c). Model results reproduce the observed BC concentrations with the exception of winter, when the model underestimates the concentrations by 50%. The high wintertime concentrations are primarily because of emissions from agricultural waste and wood fuel burning that is dominant over the Indo-Gangetic Plain during winter (Ram et al., 2010b). Model results using the upper bound of BC emissions capture the observed high concentrations in winter (Fig. 2a–c). The wintertime low biases in the model therefore clearly call for enhanced emission estimates. Moorthy et al. (2013) found that modeled surface-BC concentrations in this region are underestimated by more than a factor of 2 during winter when the planetary boundary layer (PBL) is convectively stable, while model underestimates are smaller in summer when the PBL is unstable. They suggested that the overestimate of wintertime PBL height in chemical transport models is an important contributor to model underestimates of surface pollutant concentrations. Lin and McElory (2010) pointed out that the assumption of full PBL mixing (instantaneous vertical mixing throughout the mixing depth) in the GEOS-Chem tends to overestimate vertical mixing under a stable PBL condition. They proposed and implemented in GEOS-Chem a non-local PBL mixing scheme (Holtslag and Boville, 1993; Lin et al., 2008), where the mixing states are determined by static instability. They used a local K-theory scheme (Louis, 1979) for a stable PBL and added a “non-local” term for an unstable PBL to account for the PBL-wide mixing triggered by large eddies. Our results show that the non-local boundary layer mixing increases surface BC concentrations by up to 25% in winter and spring, a significant improvement. Nair et al. (2012) showed that the non-local boundary layer mixing still tends to overestimate the vertical mixing during winter in the Indo-Gangetic Plain.

Model results are within ±50% of the observations at two remote sites, Zhuazhang (28.0° N, 99.7° E; 3583 m a.s.l.; Fig. 2h) and NCOS (30.8° N, 91.0° E; 4730 m a.s.l.; Fig. 2i), where observations are available for only fall and winter. Model results are lower than the observations at NCOP (28.0° N, 86.8° E; 5079 m a.s.l.; Fig. 2g) and Nagarkot (27.7° N, 85.5° E; 2150 m a.s.l.; Fig. 2e) by a factor of 2 in spring. Using the upper bound of BC emissions captures the springtime high concentrations at Nagarkot but not at NCOP (Fig. 2g, e). The two sites are influenced by emissions from nearby Nepal valleys transported by the mountain-valley wind (Carrico et al., 2003; Bonasoni et al., 2010). In contrast, model results capture the relatively high concentrations in winter and spring observed at Manora Peak (29.4°, 79.5° E; 1950 m a.s.l.; Fig. 2d) and Langtang (28.1° N, 85.6° E; 3920 m a.s.l.; Fig. 2f), but overestimate the summertime concentrations by a factor of 2. Using the lower bound of BC emissions is still not able to capture the observed low values in summer (Fig. 2d, f). Part of the discrepancies is explained by the inherent difficulty in simulating the meteorological fields over the complex Himalayan terrain. Chen et al. (2009) showed that the terrain effects and meteorological features in the Tibetan Plateau are not entirely reproduced by the GEOS-5 meteorological fields. Such difficulty is not unique to the Himalaya region. Emery et al. (2012) also showed that the transport of chemical species is not well simulated over the complex terrain in the western US using GEOS-Chem driven by GEOS-5 meteorological fields 2° × 2.5°.

Surface concentrations of BC at Lhasa and Delhi, two urban sites (see Table 1), are strongly affected by emissions from city traffic and industries (Zhang et al., 2008; Beegum et al., 2009). The BC concentrations at Dibrugarh are highly impacted by the emissions from the oil wells upwind and vehicular emissions from national highways nearby (Pathak et al., 2010). The concentrations at Dunhuang, a well-known tourist attraction and archaeological site, likely reflect vehicular emissions associated with tourist traffic including tour buses. All four sites are characterized by strong local emissions. Model results reproduce the seasonal trends at these “urban” sites (sites that are near urban centers or heavily influenced by local emissions), but are low by an order of magnitude (Fig. 3). Using the upper bound of BC emissions results in doubling BC concentrations (Table 1), which by itself still cannot fully explain the model versus observation discrepancies. We exclude these four urban sites from analysis hereinafter.
There is a small negative bias of $-0.3\, \mu g\, m^{-3}$ in model-simulated surface-BC concentrations (Fig. 4, left column), and the difference between model results and the observations is statistically insignificant. We note that the residual errors at very low BC concentrations may not be particularly meaningful. Overall, model results reproduce the spatiotemporal variation of surface BC concentration throughout the Tibetan Plateau ($r = 0.9$, root-mean-square error (RMSE) = 1.3 $\mu g\, m^{-3}$) with the exception of peak values (Fig. 4).

### 3.2 BC in snow

BC deposition and precipitation together determine BC concentration in snow, which we approximate as the ratio of total BC deposition to total precipitation (see Sect. 2.2.2). Figure 5 shows GEOS-Chem simulated annual mean BC deposition and GEOS-5 precipitation over Asia. The largest BC deposition over the Tibetan Plateau is in the Himalayas and the southeastern plateau (Fig. 5a), reflecting the proximity of strong BC sources in northern India and southwestern China (Lu et al., 2012) and the intense precipitation in the region (Fig. 5b). The northern plateau is heavily influenced by BC transported in the westerlies (Kopacz et al., 2011; Lu et al., 2012), but the lack of strong precipitation (Fig. 5b) results in considerably smaller BC deposition (Fig. 5a).

Table 2 shows BC concentrations in snow at 18 sites across the plateau. The concentrations are 30% lower during the monsoon season (June–September) than during the non-monsoon seasons (October–May), both in the observations and in the model. Here the monsoon and non-monsoon seasons are defined following Xu et al. (2009). The lowest BC concentrations in snow (minimum of 4.3 $\mu g\, kg^{-1}$) are in the northern slope of the Himalayas, while the highest values (maximum of 141 $\mu g\, kg^{-1}$) are to the north of the plateau. Such spatial variation largely reflects the varying elevations of the sites. Ming et al. (2009a, 2013) have shown that observed BC concentration in snow over the Tibetan Plateau is inversely correlated with the elevation of a site, with lower concentrations at higher elevations. Our model
results capture this spatial variation, but deviate from the observed concentrations by more than a factor of 2 at several sites in the Himalayas and the central plateau (Table 2).

Model results overestimate BC concentrations in snow during the monsoon season by a factor of 2–4 at three Himalayan sites: Zuoqiu (29.2°N, 96.9°E; 5500 m a.s.l.), East Rongbuk (28.0°N, 87.0°E; 6500 m a.s.l.), and Namu-nani (30.4°N, 81.3°E; 5900 m a.s.l.) (Fig. 6). Model results using the lower bound of BC emissions still overestimate the concentrations at these three sites (Table 2). Wet scavenging accounts for more than 80% of the BC deposition over the Tibetan Plateau during the monsoon season in the model. The large overestimate implies either excessive wet deposition or inadequate precipitation or both in the Himalayas, given that BC concentration in snow is approximated here as the ratio of BC deposition to precipitation (see Sect. 2.2.2). Figure 7 shows the monthly precipitation over different parts of the Tibetan Plateau from the Global Precipitation Climatology Project (GPCP, Huffman et al., 2001), the NOAA Climate Prediction Center (CPC) Merged Analysis of Precipitation (CMAP, Xie and Arkin, 1997), the University of East Anglia Climate Research Unit (CRU, Harris et al., 2014) and GEOS-5. GPCP precipitation is generally consistent with that from CMAP in most parts of the plateau except the southeastern plateau, where it is stronger by more than a factor of 2. CRU precipitation tends to be much stronger than those from GPCP and CMAP during the monsoon season, particularly in the southeastern plateau and the Himalayas (Fig. 7). Previous studies have shown that the monsoon precipitation in the Himalayas is too weak in both GPCP and CMAP data (Kitoh and Kusunoki, 2008; Voisin et al., 2008) yet too strong in the CRU data (Zhao and Fu, 2006; Xie et al., 2007). The scarcity of observational sites and the complex terrain of the Himalayas are two of the principle reasons for large uncertainties in different precipitation data sets and apparent inconsistencies among them (Ma et al., 2009; Anderman et al., 2011). Figure 7 shows that GEOS-5 precipitation is stronger than GPCP and CMAP data by a factor of 2 in the Himalayas during the monsoon season. To probe the sensitivity of BC deposition and our calculated BC concentration in snow in the Himalayas to precipitation, we conducted a GEOS-Chem simulation where we reduced GEOS-5 precipitation in the region by 20% during the monsoon season. The resulting BC wet deposition is only slightly lower (up to 5%), rather insensitive to changes in presumably already intense precipitation during the monsoon season in the region. This lack of strong sensitivity reflects an already efficient wet scavenging of BC in the intense monsoon precipitation. The resulting BC concentrations in snow are higher by 18% on average in the region, because the reduced precipitation tends to concentrate BC in snow. Therefore, model overestimates of BC concentration in snow in the region are more likely resulting from excessive BC deposition than from excessive monsoon precipitation. This is likely a result of over-long BC lifetime due to insufficient wet removal. Wang et al. (2014) compared GEOS-Chem simulated atmospheric BC concentrations with the HIAPER Pole-to-Pole Observations (HIPPO) aircraft measurements and concluded that wet scavenging in the model is too weak. This is in part because of the underestimated scavenging efficiency of BC in the model. In addition, the relatively long BC aging time used in the model also potentially contributes to the weak wet scavenging. Recent observations suggest that the e-folding time of about 1 day, for the hydrophobic-to-hydrophilic conversion of BC, typically used in global models is too long (Akagi et al., 2012). The uncertainties in wet scavenging in the model reduce the accuracy in the estimates of snow BC concentrations. Excessive BC deposition can in part result from excessively strong PBL mixing in the source regions and consequently excessive BC being transported into the free troposphere. Our model results show that the non-local boundary layer mixing (Lin and McElory, 2010) reduces BC wet deposition by up to 5% on average in the Himalayas during the monsoon season.

Figure 6 shows that our calculated BC concentrations in snow are lower than observations by a factor of 2 across the central plateau. Model results using the upper bound of BC emissions are able to reproduce the high BC concentrations in snow at La’pong but miss those high values at the other sites (Table 2). Ming et al. (2009a) pointed out that this region is predominantly influenced by biofuel burning (residential cooking and heating) and biomass burning from religious activities. These local emissions are largely unaccounted for in current emission inventories (Wang et al., 2012). In addition, it is likely that the lack of consideration of snow aging also lowers the BC concentration in snow computed here (Xu et al., 2006). We choose to exclude Meikuang and Zhadang from the comparison here on account of local emissions from coal-containing rock strata at the former (Xu et al., 2006) and strong snow melting at the latter (Zhou et al., 2007). GEOS-5 precipitation in the central plateau is in general agreement with those from CMAP and CRU during the non-monsoon season and that from GPCP during the monsoon season (Fig. 7e).

Model results are consistent with observations at the elevated sites in the northwestern and northeastern plateau and to the north of the plateau (Fig. 6), where free tropospheric BC is primarily northern mid-latitude pollution transported by the westerlies (Kopacz et al., 2011; Lu et al., 2012). Regional emissions from western and central China also contribute to BC deposition in these regions (Lu et al., 2012). Although precipitation in these regions is weaker in GEOS-5 than in GPCP and CMAP (Fig. 7b, d), previous studies have shown that GPCP and CMAP precipitation is likely too strong there (Voisin et al., 2008; Ma et al., 2009).

Overall model results of BC concentration in snow have a small negative bias but a large RMSE (Fig. 4, middle column); the latter results from the large discrepancies in the Himalayas and the central plateau. Model results are
statistically in good agreement with observations and reproduce the observed spatiotemporal variation ($r = 0.85$).

### 3.3 BC AAOD

Modeled BC AAOD is consistently lower than AERONET retrievals at most sites on both a monthly (Fig. 4, right column) and an annual bases (Table 3). The annual mean modeled BC AAOD over the Tibetan Plateau is 0.002 (Fig. 8), considerably lower than the observations. Model results somewhat capture the observed spatial and seasonal trends ($r = 0.53$), but to varying degrees underestimate the magnitudes (Fig. 9). Forty percent of the data points are too low by more than a factor of 2 in the model, particularly in the Himalayas and to the northwest of the plateau during winter.
and spring when emissions are larger (relative to emissions during the rest of the year). Most AERONET measurements in and around the plateau are after 2006, whereas our model results are for 2006. BC emissions in India have increased by 3.3 % yr\(^{-1}\) since 2006 (Lu et al., 2011). Therefore, the large low bias in part reflects the abovementioned temporal (hence emissions) mismatch. Using the upper bound of BC emissions reduces the model versus observations discrepancies at six sites (Table 3), but model results are still lower by a factor of 2 than the observed high AAODs at the other sites. We also note that there are large uncertainties in the AERONET AAOD retrieval (Bond et al., 2013), and that BC AAOD data are only scarcely available in the plateau and adjacent regions.

Another equally important factor contributing to the large discrepancy is the assumption of external mixing of BC in the model, which leads to a weaker BC absorption (Jacobson, 2001). Previous studies have found that BC absorption is enhanced by 50 % because of internal mixing (Bond et al., 2006). We find that a 50 % increase of BC absorption (using a MAC of 11 m\(^2\) g\(^{-1}\) in our calculation) would reduce the model bias by 57 % in the Indo-Gangetic Plain and the northeastern plateau and to the northeast of the plateau, and by 16 % along the southern slopes of the Himalayas and to the northwest of the plateau (Fig. 9, right panel). There is evidence that the enhancement of BC absorption due to internal mixing may be considerably smaller than previously thought (Cappa et al., 2012). It is clear that the large discrepancy (more than a factor of 2) in the Himalayas and to the northwest of the plateau cannot be fully explained by the lack of BC internal mixing consideration (and the associated larger absorption) in the model. Bond et al. (2013) pointed out that current models significantly underestimate BC AAOD, particularly in south and southeast Asia, primarily because of the absence of internal mixing and underestimated emissions. They recommended scaling up modeled BC AAOD to AERONET observations in order to accurately estimate BC radiative effects.

Therefore, although surface BC concentration is relatively well captured by model results (see Sect. 3.1), more measurements of vertical profiles over the Tibetan Plateau are imperative for evaluating column quantities such as BC AAOD.

4 Sensitivity to BC emissions

Figure 2 shows that model-simulated surface-BC concentrations are considerably lower in Experiment B (using the INTEX-B inventory) relative to Experiment A (using the LU inventory) at rural sites. The difference in surface BC concentration is more than 30 % at rural sites and 10–20 % at remote sites, decreasing with distance from the source region. Such varying difference in surface BC concentration largely reflects the spatially non-uniform differences between the two emission inventories. The difference in BC concentration in snow between the two sets of results is less than 20 %. The relatively smaller difference is because the sites with measurements of BC concentration in snow are invariably remote high-elevation sites, further away from the source regions. BC concentrations in snow are higher over the northwestern and northeastern plateau and to the north of the plateau but lower in the Himalayas and the central plateau in Experiment B than in A. This is because the BC emissions in the central plateau and India are lower, and the emissions in northwestern and central China are higher in the INTEX-B than in the LU inventories. BC AAOD values are higher (15 %) to the northeast and northwest of the plateau and lower (10–60 %) in the Indo-Gangetic Plain in Experiment B than in A. Therefore, both surface BC concentration and AAOD along the southern slope of the Himalayas are strongly sensitive to Indian emissions, while the high-altitude remote sites are less affected by the emission changes in the source regions. Overall, Experiment B results show larger negative bias and root-mean-square error (RMSE) (Table 6) and lower Taylor score (Fig. 10) relative to Experiment A. As such, our results suggest that the INTEX-B inventory considerably underestimates anthropogenic BC emissions in India, which is also implied by some latest estimates of BC emissions in Asia (Kurokawa et al., 2013; Wang et al., 2014).

5 Sensitivity to BC aging parameterization

Compared with model results from the standard simulation (Experiment A, Table 5), the use of Liu et al. (2011) parameterization for BC aging in the model (Experiment C, Table 5) results in increased surface-BC concentrations, BC concentrations in snow, and BC AAOD, because of the longer BC atmospheric lifetime against wet scavenging (see Sect. 2.2.3). The increase in surface BC concentration is 1 % on average.
Figure 9. Observed and GEOS-Chem simulated monthly mean BC AAOD at AERONET sites over the Tibetan Plateau (see Table 3 and Fig. 1). Left panel: assuming external mixing of BC. Right panel: assuming a 50 % increase of BC absorption associated with internal mixing. Regions are color-coded: Indo-Gangetic Plain (red), the Himalayas (green), the northeastern plateau (blue), northeast of the plateau (grey), northwest of the plateau (magenta). Solid line is a 1 : 1 ratio line; dashed lines are 1 : 2 (or 2 : 1) ratio lines; dashed-dotted lines are 1 : 4 (or 4 : 1) ratio lines. Also shown are the correlation coefficient ($r$) and $p$ value. Values are for 2006 unless stated otherwise. See text for details.

Figure 10. Taylor diagram of GEOS-Chem simulated versus observed BC concentration in surface air ($\text{BC}_{\text{air}}$) and in snow ($\text{BC}_{\text{snow}}$), and BC AAOD at sites over the Tibetan Plateau (see Tables 1, 2 and 3 and Fig. 1). Red asterisk is the observation. Triangles, circles and squares, respectively, indicate modeled $\text{BC}_{\text{air}}$, $\text{BC}_{\text{snow}}$, and BC AAOD from Experiments A (blue), B (green), C (magenta) and D (yellow). See Table 5 and text for more details on the model experiments. Also shown are the Taylor scores (S). Values are for 2006 unless stated otherwise. See text for details.

Sensitivity to model resolution

Compared with model results from the standard simulation (Experiment A, Table 5), the use of a finer model resolution ($0.5^\circ \times 0.667^\circ$) nested over Asia (Experiment D, Table 5) reduces the bias in modeled surface-BC concentrations from 15 to 2 % but increases the RMSE (Table 6). Wang et al. (2014) found that replacing a coarse-resolution model ($1.27^\circ \times 2.5^\circ$) with a finer-resolution one ($0.51^\circ \times 0.66^\circ$) reduces the bias of surface BC simulation in Asia by 30 %. Figure 2 shows (maximum 3 %) at rural sites and 10 % on average (maximum 30 %) at remote sites (Fig. 2). This is consistent with the results from Huang et al. (2013), where the use of Liu et al. (2011) parameterization only changes GEOS-Chem simulated surface-BC concentrations by less than 0.01 $\mu g \text{ m}^{-3}$ in the Tibetan Plateau. Liu et al. (2011) showed that their aging parameterization significantly improves seasonal variations of modeled surface-BC concentrations in the Arctic. This is different from the present study, where the aging parameterization has a minor impact on the seasonality of both surface BC concentrations and AAOD (Fig. 2 and Table 6). The increase in BC AAOD is 10 % on average (maximum 30 %) at most AERONET sites but leads to lower spatiotemporal correlations with observations (Table 6). The aging parameterization has a much stronger impact on modeled BC concentration in snow than on surface BC concentration and BC AAOD. The increase in BC concentration in snow is more than 30 % at a number of sites in the Himalayas and the central plateau (Table 2). Compared with the standard simulation (Experiment A), the use of the Liu et al. (2011) parameterization results in an overestimate of BC concentration in snow relative to observations, which increases the absolute bias by a factor of 2 (Table 6) and decreases the Taylor score (Fig. 10). This suggests that the Liu et al. (2011) aging parameterization may result in excessively long BC-conversion times (from hydrophobic to hydrophilic) and hence excessively long atmospheric lifetimes.
that the nested model simulation slightly improves seasonal variations of surface BC concentrations at several remote sites, whereas it does not improve those at urban sites both in magnitude and in temporal variation (Fig. 3). This is similar to the results in Fu et al. (2012), where the GEOS-Chem nested model underestimates surface BC concentrations by an order of magnitude at Dunhuang and Lhasa sites, even using the enhanced BC emissions from top-down estimates. Compared with the standard simulation, the nested model simulation increases the absolute bias by 57 % in modeled snow-BC concentration and by 5 % in modeled BC AAOD (Table 6). The nested model results also show a lower spatiotemporal correlation with observations of snow BC concentration and BC AAOD (Fig. 10). Our results suggest that the finer model resolution alone cannot explain model versus observation discrepancies on the simulation of snow BC concentration and BC AAOD in the Tibetan Plateau.

7 Summary and conclusions

This study sought to understand the capability of GEOS-Chem in simulating BC over the Tibetan Plateau and the potential factors driving model versus observation discrepancies. We used GEOS-Chem version 9-01-03 driven by GEOS-5 assimilated meteorological fields and systematically evaluated the model simulations against in situ measurements of BC in surface air, BC in snow, and BC AAOD for 2006. We also examined the effects of anthropogenic BC emissions from China and India, BC aging process and model resolution on BC simulations.

Model results captured the seasonal variation of surface BC concentrations at rural sites, but the observed wintertime high values were absent in the model, which calls for improved emission estimates particularly in the Indo-Gangetic Plain. The use of non-local PBL mixing scheme reduced part of the discrepancy between observed and modeled surface-BC concentrations in winter. Modeled surface-BC concentrations at remote sites were within a factor of 2 of the observations. Part of the discrepancy is explained by the inherent difficulty in simulating the meteorological fields over the complex Himalayan terrain. Surface BC concentrations at urban sites are significantly underestimated by model results.

Modeled BC concentrations in snow were spatiotemporally consistent with observations ($r = 0.85$). The highest snow BC concentrations were seen north of the plateau (40–50° N), while the lowest values were found in the northern slope of the Himalayas. However, model results were a factor of 2–4 higher than the observations at three Himalayan sites during the monsoon, primarily because of the excessive BC deposition resulted from overlong BC lifetime. Model results underestimated snow BC concentration by a factor of 2 in the central plateau, due to the lack of snow aging in the CTM and the strong local emissions unaccounted for in the emission inventories. Model results are consistent with the observations at the elevated sites in the northwestern northeastern plateau and to the north of the plateau. Model results of both BC in snow and in surface air showed no statistically significant difference with observations with biases less than 15%.

Modeled BC AAOD is consistently biased low at most AERONET sites over the plateau, especially to the northwest of the plateau and in the Himalayas in winter and spring. The large model versus observation discrepancies were mainly because of underestimated emissions and the assumption of external mixing of BC in the model. This suggests that modeled BC AAOD should be scaled to AERONET observations in order to accurately estimate BC climatic effects. More measurements of vertical profiles over the Tibetan Plateau are imperative for evaluating modeled column quantities such as BC AAOD.

Sensitivity simulations showed that both surface BC concentration and BC AAOD along the southern slope of the Himalayas were strongly sensitive to Indian emissions, while the elevated remote sites were less affected by the change of emissions in source regions. The BC aging parameterization from Liu et al. (2011) resulted in a large increase of BC concentration in snow, but only had a minor impact on surface BC concentration and AAOD. The use of a finer model resolution nested over Asia reduced the bias in modeled surface-BC concentration from 15 to 2 %, but increased

Table 6. Error statistics of GEOS-Chem simulations of BC in the Tibetan Plateau for 2006.

| Statistical quantities* | BC in surface air | BC in snow | BC AAOD |
|------------------------|------------------|------------|---------|
|                        | A     | B     | C     | D     | A     | B     | C     | D     | A     | B     | C     | D     |
| Mean error             | -0.29 | -0.82 | -0.26 | -0.03 | -1.23 | -0.75 | 3.39  | 1.94  | -0.019 | -0.023 | -0.020 | -0.020 |
| Mean absolute error    | 0.59  | 0.93  | 0.59  | 0.74  | 13.39 | 12.74 | 16.09 | 16.09 | 0.020  | 0.023  | 0.020  | 0.020  |
| Fractional gross error | 0.39  | 0.46  | 0.39  | 0.46  | 0.43  | 0.40  | 0.48  | 0.58  | 0.78   | 0.93   | 0.80   | 0.77   |
| Root-mean-square error (RMSE) | 1.34  | 1.95  | 1.33  | 1.33  | 16.73 | 16.65 | 18.67 | 24.54 | 0.026  | 0.029  | 0.027  | 0.027  |
| Bias-corrected RMSE    | 1.31  | 1.77  | 1.30  | 1.33  | 16.69 | 16.63 | 18.36 | 24.46 | 0.017  | 0.018  | 0.018  | 0.019  |
| Correlation coefficient ($p$ value < 0.001) | 0.90  | 0.87  | 0.90  | 0.87  | 0.85  | 0.86  | 0.81  | 0.70  | 0.53   | 0.46   | 0.45   | 0.37   |

* Units for mean error, mean absolute error, RMSE and bias-corrected RMSE are µg m$^{-3}$ for BC in surface air and µg kg$^{-1}$ for BC in snow.
the bias in modeled snow-BC concentration and BC AAOD by 57% and 5%, respectively. More quantitative analyses are required to investigate the uncertainties in different model processes of BC simulations.

Therefore, accurate model simulations of BC in surface air, BC in snow and BC AAOD over the Tibetan Plateau require improvements in BC emission inventories particularly for China and India, model meteorological fields over the complex Himalayan terrain, PBL mixing scheme particularly under a stable PBL condition, model representation of atmospheric BC aging process including the conversion of hydrophobic to hydrophilic BC and the enhancement of BC absorption resulted from internal mixing, model parameterization of BC wet scavenging, and model simulation of snow processes such as snow aging, snow melting and BC transfer among different snow layers.

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