We thank the editor (Dr. Xiaobin Xu), Dr. Jing Ming, and the anonymous referee for their careful review of the revised manuscript and valuable comments. Below are our point-by-point responses (blue color) and note of further changes we’ve made to the manuscript (in italic).

Editor Decision: Reconsider after minor revisions (Editor review) (11 Dec 2014) by Xiaobin Xu

Comments to the Author:

Dear authors,

There are some remaining issues in your revised manuscript. Please make further revision according to the referee’s reports.

Response: we have addressed all the remaining issues raised by the two referees. Please see our detailed responses as follows.

I have some additional points:

1. It is not easy for me to understand Fig. 4. Do the plots really present fractional source contributions to the BC deposition flux at the drilling site? If yes, the sum of the data from different regions surrounding the site should be 100%. However, the sum of the “fractional contributions” from South Asia and East Asia seems to be much higher than 100% if the color scale is correct, not to mention contributions from other regions. How can you obtain the data in Table 1?

Response: Yes, the plots do show fractional contributions to BC deposition flux from four major source regions (outlined by the big black box in each panel in the same row), but not just for the drilling site. For example, the upper-right panel shows the spatial distribution of contributions from emissions in South Asia to BC deposition at any location on the map during monsoon season. For the drilling site, only the color in the small black box, greenish (30-40%) in this South Asia monsoon case, should be counted. Then percentages indicated by colors in the small black box in all four panels of the same row should add up to a number very close to 100%. The small deficit would be the sum of contributions from the rest of the world that are not shown in the figures. We have double-checked the colors and the numbers in Table 1 and they match. To clarity, we have added the following sentence to the figure caption. “Color in the small black box in each panel corresponds to the fraction contribution to BC deposition at the sampling site. Exact percentage contributions are provided in Table 1.”

2. Lines 187-193, I think “meteorological conditions (and associated transport and wet removal processes)” and “local precipitation rate” are not independent of each other, so the logic here seems to be obscure.
Response: Thanks for pointing this out. We meant to say “meteorological conditions that determine transport pathways (and associated wet removal processes during the transport)” and “precipitation rate at the sampling location”. The sentences have been reworded.

3. Line 48, delete the “in” after “absorbs”.

Response: Done.

4. Line 136, “the global distribution of aerosols” cannot be “improved”. Please reword the sentence.

Response: Reworded to “…improved the model prediction of global distribution of aerosols…”

5. Fig. 6, please use another color for the India emissions given by Lu et al. (2011). It should also be mentioned that the flux data are based on ice-core measurements or modeling.

Response: Figure 6 has been revised as suggested. We also noted in the figure caption that the deposition fluxes were based on the ice-core measurements.

Referee #2 (Dr. Jing Ming)

The authors have addressed all the issues raised by me in the previous comments. However, some the comments are not correctly addressed, possibly due to my unclear requests. Please clarify them before the work can be published in ACP.

1. Citing the previous comment, “This work introduced a new concept, OC’s forcing, which has not been widely recognized by the societies. The introduction of a new concept must be previously supported by measurement, as we all know. The OC’s forcing is just like an aerolite in the whole paper. The authors should firstly list some literatures that clarified OC has the characteristic of radiation absorption in snow as well as it does in atmosphere claimed by Bond and Bergstrom (2006) and Kirchstetter et al. (2004). The online SNICAR model only simulates the reductions of snow albedo caused by BC and dust, but cannot have the ability to simulate the forcing of OC. The mass absorption cross-section of OC in the atmosphere cannot be directly used for it in snow.”
   - For the issue of OC’s forcing, the author list some studies concerning the OC’s radiation absorption in the atmosphere. My question is whether the forcing still exists in snow, and how to apply the SNICAR model to correctly simulate the forcing, for the model builder himself does not mention this. I strongly suggest the author to consult Dr. Mark Flanner, the model builder to get suggestions.
Response:

We truly appreciate the referee for his concern on the OC-in-snow forcing. We agree that most studies on light-absorbing properties of OC in the literature focused on airborne OC (or brown carbon), but only a few have investigated OC in snow (see below for a brief literature review). To the best of our knowledge, however, light-absorbing OC particles in snow can also exert a positive forcing like BC particles do. Co-authors (H. Wang, Y. Qian and PJ Rasch) have had extensive collaborations with Dr. Mark Flanner in improving the representation of light-absorbing particles in the SNICAR model and its coupling to the CAM5 aerosol-climate model. They have long recognized the importance of including OC absorption in SNICAR, in which OC in snow is currently inactive by default (e.g., switched off) because of large uncertainties in OC optical properties. A large amount of work is still needed to develop OC optical properties that span the dimensions of snow grain size and OC particle size, as Dr. Flanner has done for BC in SNICAR (personal communication with Dr. Mark Flanner). This was the reason why we simply used BC’s spectral dependence of light absorption for OC but scaled down the mass absorption cross-section in the radiative forcing calculation using SNICAR, and we clearly stated the limitations of the crude estimates.

To further address the referee’s concern, we have decided to separate the OC-in-snow forcing estimation from that for BC and put up front the discussions on the limitation of the methodology and potential biases in the forcing estimates. We have also condensed the literature review on airborne OC but expanded the review on OC-in-snow studies, according to the referee’s suggestion. For the convenience of the referee and editor, we make a note of the newly added sentences here as follows.

A growing number of studies (e.g., Kirchstetter et al., 2004; Andreae and Gelencsér, 2006; Hoffer et al., 2006; Yang et al., 2009; Kirchstetter and Thatcher, 2012) have reported that airborne brown carbon can contribute significantly to aerosol light absorption in the atmosphere, although there are still substantial uncertainties in quantifying optical properties of brown carbon, which makes the model estimation of OC radiative forcing difficult. Similarly, the importance of OC absorption in snow has been recognized and suggested for inclusion in modeling aerosol snow-albedo effect (e.g., Flanner et al., 2009; Aoki et al., 2011). Observational analysis of light-absorbing particles in Arctic snow reported that the main non-BC component is brown carbon, which accounted for 20-50% of the visible and ultraviolet absorption (Hegg et al., 2009, 2010; Doherty et al., 2010). In the rural area of central north China, brown carbon in winter snow also played an important role in visible light absorption, which contributed about 60% to light absorption at 450 nm and about 40% at 600 nm (Wang et al., 2013). A more recent observational study by Dang and Hegg (2014) qualified the light absorption by different light-absorbing particulates in snow, and suggested that
humic-like substances and polar OC contributed 9% and 4% to the total light absorption respectively. Despite the substantial uncertainties in brown carbon optical properties, a recent global modeling study (Lin et al., 2014), in which a range of optical properties of brown carbon taken from the literature were applied to OC-in-snow concentrations simulated in a global chemical transport model, showed that the global OC forcing in land snow and sea ice is up to 24% of that caused by BC.

The SNICAR model currently does not support the calculation of OC-in-snow forcing in the same way as that for BC due to a lack of reliable OC optical properties that span the dimensions of snow grain size and OC particle size (personal communication with Mark Flanner, 2014). We take a MAC value of 0.6 m² g⁻¹ at 550 nm for OC (Kirchstetter et al., 2004), and assume a constant factor of 0.08 (i.e., 0.6/7.5) to scale down MAC values of BC at all wavelengths to obtain a first-order guess of OC-in-snow forcing using SNICAR. The estimated OC forcing has a 4-fold increase from 0.2 W m⁻² (for mean OC concentration of 13.8 ng g⁻¹ during 1956-1979) to 0.84 W m⁻² (for mean OC concentration of 61.3 ng g⁻¹ in 2006), which are 27% and 43% of corresponding BC-in-snow forcing, respectively. The BC/OC forcing ratios based on our simple guesses are larger than the upper bound of estimates (i.e., 24%) by Lin et al. (2014).

Two main assumptions could have caused our first-order estimate of OC forcing to have large biases. First, the MAC value of 0.6 m² g⁻¹ (at 550 nm) was based on OC extracted from biomass burning samples that tends to have higher absorption efficiency than OC emitted from fusil fuel combustion (Kirchstetter et al., 2004). This may cause an overestimation of OC forcing. Second, we treated all the water-insoluble OC from the ice-core measurements as light-absorbing brown carbon in the forcing estimation, which also likely results in an overestimation of OC forcing if a significant fraction of OC is non-absorbing. However, water-soluble part, accounting for about half of OC observed in Manora peak and northwest India (Ram et al., 2010; Rajput et al., 2013), can also contribute to some absorption of UV and visible light (Chen and Bond, 2010; Beine et al., 2011). Thus the absorption by water-soluble OC that was not included in the forcing estimate may compensate for the high bias to some extent. According to a laboratory study by Chen and Bond (2010), a large fraction of absorbing OC from hard wood burning is water-insoluble. As water-insoluble OC recorded in the ice core herein was very likely dominated by biomass burning emissions (Section 3.4), the second assumption we used here may not cause a huge bias in estimating OC forcing in snow.

2. Citing the previous comment, "Paragraph 2 in Page 19721. Ming et al. (2013) in Adv. Water. Resources suggest BC deposited in Himalayan and High Asian glaciers cannot significantly affect their energy balances, which is a very minority but different viewpoint from most literatures listed here, which should not be neglect here.”
The author just put the reference in the context, without making any notes. Please address the original intention raised by the reviewer in the paragraph.

Response: The original concern has been addressed, and another paper with a similar opinion has been added too.

Referee #3

The manuscript has been improved and the authors have addressed my comments carefully. I have only two minor comments.

1. In section 2.2 Model and experiment setup (page 6 line 130-133): Please indicate which modal aerosol model was used in this CAM5 simulation. CAM5 can run at least two aerosol microphysics models: 7 mode and 3 mode. Only the 3 mode model allows BC and POC emitted into an accumulation size mode, while the 7 mode model gets the emission into primary carbon mode.

Response: We used the default 3-mode aerosol model, and have now noted it in the manuscript.

2. Page 11 line 267-268 and table 1: Please check the unit (Tg/a) and emission amount of BC in Table 1. If the BC emission unit is Tg/a, the annual emission should be the sum of the emissions over monsoon and non-monsoon periods.

Response: We have double-checked the units and numbers in Table 1 and they are correct. The original emissions are monthly means, and then further averaged over 4 months (June-September) for the monsoon period, 8 months (October-May) for the non-monsoon period and 12 months (January-December) for the entire year. Therefore, the BC emissions in Table 1 still represent monthly mean rather than total. Units (in Tg/a) are made consistent with those used in the literature. The annual emission rate times 12 should equal to the sum of emission rate over monsoon and non-monsoon periods times 4 and 8, respectively. We have now added “mean” before “BC emissions” in the table title for clarity.
Carbonaceous Aerosols Recorded in a Southeastern Tibetan Glacier: Analysis of Temporal Variations and Model Estimates of Sources and Radiative Forcing

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Abstract. High temporal resolution measurements of black carbon (BC) and organic carbon (OC) covering the time period of 1956-2006 in an ice core over the southeastern Tibetan Plateau show a distinct seasonal dependence of BC and OC with higher respective concentrations but lower OC/BC ratio in the non-monsoon season than during the summer monsoon. We use a global aerosol-climate model, in which BC emitted from different source regions can be explicitly tracked, to quantify BC source-receptor relationships between four Asian source regions and the southeastern Tibetan Plateau as a receptor. The model results show that South Asia has the largest contribution to the present-day (1996-2005) mean BC deposition at the ice core drilling site during the non-monsoon season (October to May) (81%) and all year round (74%), followed by East Asia (14% to the non-monsoon mean and 21% to the annual mean). The ice-core record also indicates stable and relatively low BC and OC deposition fluxes from late 1950s to 1980, followed by an overall increase to recent years. This trend is consistent with the BC and OC emission inventories and the fuel consumption of South Asia (as the primary contributor to annual mean BC deposition). Moreover, the increasing trend of OC/BC ratio since the early 1990s indicates a growing contribution of coal combustion and/or biomass burning to the emissions. The estimated radiative forcing induced by BC and OC impurities in snow has increased since 1980, suggesting an increasing potential influence of carbonaceous aerosols on the Tibetan glacier melting and the availability of water resources in the surrounding regions. Our study indicates that more attention to OC is merited because of its non-negligible light absorption and the recent rapid increases evident in the ice core record.

Keywords

Carbonaceous aerosol, Tibetan glacier, Emissions, Radiative forcing

1. Introduction
Carbonaceous aerosol, released from fossil fuel, biofuel and/or biomass combustion, contains both black carbon (BC, a.k.a. elemental carbon, EC), a strong light absorber, and organic carbon (OC), which also absorbs the near infrared, but more weakly than BC (Kirchstetter et al., 2004; Bond et al., 2006). Often mixed with other aerosol species, BC impacts human health, crop yields and regional climate (Auffhammer et al., 2006; Tie et al., 2009), and is believed to be the second strongest climate warming forcing agent after carbon dioxide (Jacobson, 2001; IPCC, 2013).

Because of their high population density and relatively low combustion efficiency, developing countries in South and East Asia such as India and China are hotspots of carbonaceous aerosol emissions (Ramanathan and Carmichael, 2008). During the cold and dry winter season, haze (heavily loaded with carbonaceous aerosols) builds up over South Asia, and exerts profound influences on regional radiative forcing (Ramanathan et al., 2007; Ramanathan and Carmichael, 2008), hydrologic cycles (Menon et al., 2002; Ramanathan et al., 2005), and likely Himalaya-Tibetan glacier melting that could be accelerated by the absorption of sunlight induced by BC in the air and deposited on the ice and snow surfaces (Ramanathan et al., 2007; Hansen and Nazarenko, 2004). Although BC deposited in snow and glaciers at some locations may not significantly affect the energy balance (Ming et al., 2013; Kaspari et al., 2014).

Due to the lack of long-term observations of emissions and concentrations of atmospheric carbonaceous aerosols, it is difficult to evaluate the effects of BC and OC on historical regional climate and environment before the satellite era. Some studies have evaluated historical anthropogenic emissions based on the consumption of fossil fuels and biofuels (Novakov et al., 2003; Ito and Penner, 2005; Bond et al., 2007; Fernandes et al., 2007). While fossil fuel is the major energy source in the urban areas of South Asia and East Asia, biomass combustion, such as fuel wood, agricultural residue and dung cake, is prevalent in rural areas (Revelle, 1976;
Venkataraman et al., 2010; Street and Waldhoff, 1998). Biomass burning has been considered as the major source of black carbon emissions (Reddy and Venkataraman, 2002; Venkataraman et al., 2005). However, as reliable biomass consumption data are hard to obtain, estimates of BC and OC emissions from biomass burning are ambiguous and incomplete.

Measurements of carbonaceous aerosol concentrations in glacier ice are an ideal means to reconstruct historical emissions and reveal long-term trends of anthropogenic aerosol impacts on local climate. Greenland ice core measurements were previously used to reconstruct the North American BC emission history and its effects on surface radiative forcing back to the 1880s (McConnell et al., 2007). Himalayan ice cores retrieved from the Tibetan Plateau have revealed the mixed historical emissions from South Asia, Central Asia and the Middle East and also been used to evaluate radiative forcing from BC in snow (Ming et al., 2008; Kaspari et al., 2011). Using the Snow, Ice, and Aerosol Radiative (SNICAR) model, Flanner et al. (2007) estimated an instantaneous regional forcing of exceeding 20 W m$^{-2}$ by BC in snow/glaciers over the Tibetan Plateau during the spring season.

By using five ice core records, Xu et al. (2009a) elucidated an important contribution of BC to the retreat of Tibetan glaciers in addition to greenhouse gases. Due to the short atmospheric lifetime of carbonaceous aerosols compared to greenhouse gases, emission reductions may be an effective way to mitigate their warming effects. Thus it is particularly important to identify the source regions and the source types of carbonaceous aerosols observed in Tibetan glaciers. Xu et al. (2009a) suggested that BC deposited on Tibetan Plateau was broadly from Europe and Asia. However, they didn’t perform in-depth analysis on emissions from more specific source regions and the source types. In this study, we use the ice core retrieved from the southeastern Tibetan Plateau, also known as the Zuoqiupu ice core in Xu et al. (2009a), to reconstruct the history of atmospheric deposition of carbonaceous aerosols in this glacier, and to characterize emissions and
source-receptor relationships with the help of a global climate model in which BC emitted from different source regions can be explicitly tracked. We also estimate the respective contributions from BC and OC to radiative forcing in the Zuoqipu glacier using the ice core measurements and the SNICAR model.

2. Methods

2.1 Measurements of carbonaceous aerosols in ice core

Zuoqipu glacier is in the southeastern Kangri Karpo Mountains, located at the southeastern margin of the Tibetan Plateau (Figure 1). In 2007, an ice core of 97 meters in depth (9.5 cm in diameter) was retrieved within the accumulation zone of Zuoqipu glacier at 96.92°E, 29.21°N, 5600 m a.s.l. The ice core was kept frozen and transported to laboratory facilities at the Institute of Tibetan Plateau Research (Lhasa branch) for analysis. The annual accumulation of snow/ice at the drill site was around 2 meters on average. The oxygen isotope ($\delta^{18}$O) samples were cut at 10 cm internals, and BC and OC samples at 10-25 cm, resulting in 18 and 9 samples per year on average, respectively. Thus this ice core provided a high temporal-resolution of $\delta^{18}$O, and BC and OC concentrations. BC and OC concentrations were measured by using a Desert Research Institute (DRI) Model 2001 Thermal/Optical Carbon Analyzer following the IMPROVE TOR protocol (Chow et al. 1993; Chow and Watson 2002; Cao et al. 2008). Note that according to the thermal/optical measurement method, the analytical result is technically called “EC”. Herein we use “BC” to be consistent with the notation in our model simulations and in the literature. The reported OC concentrations from the ice-core measurements can only account for water-insoluble part of OC in the ice samples because most of the water-soluble part cannot be captured by the filter-based method applied to liquid samples (melted from the ice). Further details on the analysis methods, ice core dating and calculation of BC and OC seasonal deposition fluxes can be found in Xu et al.
2.2 Model and experimental setup

We use the Community Atmosphere Model version 5 (CAM5; Neale et al., 2012) to help understand the emissions, transport and dry/wet deposition of carbonaceous aerosols in the atmosphere. In the default 3-mode modal aerosol scheme of CAM5 used for this study, BC and primary OC are emitted into an accumulation size mode, where they immediately mix with co-existing hygroscopic species such as sulfate and sea salt (Liu et al., 2012). Hygroscopic aerosol particles in the accumulation mode are subject to wet removal by precipitation. Recent model improvements to the representation of aerosol transport and wet removal in CAM5 by Wang et al. (2013) have substantially improved the model prediction of global distribution of aerosols, particularly, over remote regions away from major sources. To minimize the model biases in simulating meteorological conditions and, particularly, circulations that are critical to aerosol transport, we configure the CAM5 model to run in an offline mode (Ma et al., 2013) with wind, temperature, surface fluxes and pressure fields constrained by observations. However, cloud/precipitation fields and interactions between aerosol and clouds are allowed to evolve freely. A source tagging technique has been recently implemented in the CAM5 model to allow for explicitly tracking aerosols emitted from individual source regions and, therefore, assists in quantitatively characterizing source-receptor relationships (Wang et al., 2014). This tagging technique along with the CAM5 model is used in the present study to do source attribution for carbonaceous aerosols deposited to the Zuoqipu glacier.

We conducted an 11-year (1995-2005) CAM5 simulation at horizontal grid spacing of 1.9° × 2.5° and 56 vertical levels, with prescribed sea surface temperatures and sea ice distribution. Reanalysis products from NASA Modern Era Retrospective-Analysis for Research and Applications (MERRA) (Rienecker et al.,
2011) are used to constrain the meteorological fields of CAM5. For aerosols (including OC, BC and other important species), we use the year-2000 monthly mean emissions described by Lamarque et al. (2010) that have been used in many global climate models for present-day climate simulations, included in the fifth assessment report (AR5) by the Intergovernmental Panel on Climate Change (IPCC). The monthly mean emissions are repeatedly used for each year in the 11-year simulation. Note that we do not intend to design the model experiment to simulate the whole historical record of BC in the ice core, but rather for a period of time to demonstrate the impact of meteorology (and associated transport and removal of aerosols) on the seasonal dependence of BC deposition in the target region and the lack of longer-term trend in deposition without considering the temporal variation of emissions.

As the ice core drill site was located at a remote and elevated area over the southeastern Tibetan Plateau, where local emissions are minimal. Deposition of carbonaceous aerosols is most likely contributed by the non-local major emission sources (e.g., distributions of mean BC emissions during non-monsoon and monsoon seasons shown in Figure 2) in South Asia and East Asia. These two regions, along with Southeast Asia and Central Asia, are identified as the potential source contributors. Thus BC emissions from the four regions and the rest of the world are explicitly tracked in the CAM5 simulation.

3. Results and Discussion

3.1 Seasonal dependence of carbonaceous aerosols

BC and OC concentrations in the Zuoqipu ice core both exhibit statistically significant seasonal variations at the 0.05 level corresponding to the stable oxygen isotope variability, which shows high values during the winter and low values during the summer (Xu et al., 2009a). As shown in Figure 3, concentrations of BC
and OC have distinct differences between the summer monsoon and non-monsoon seasons. Seasonally varying emissions and meteorological conditions that determine the transport pathways of BC and OC emitted from major sources, removal during the transport, and local precipitation rate can cause the seasonal variations of BC and OC in ice at the sampling site. The seasonal dependence of BC and OC in ice core is consistent with available observations of atmospheric aerosols in the south slope of the Himalayas and the southeastern Tibetan Plateau, where the high concentration of carbonaceous aerosols during the cold and dry season was suggested to associate with the South Asian haze (Cong et al., 2009; Marinoni et al., 2010; Kaspari et al., 2011; Zhao et al., 2013a; Zhao et al., 2013b). The consistency between the seasonal dependence of airborne BC and OC concentrations and the seasonal variation of ice-core measurements indicates that seasonal differences in precipitation rate at the sampling location is less likely to be the determining factor. Our model results (details discussed in the section 3.2) suggest that the seasonal dependence of BC deposition flux in the target region could be mainly due to meteorological conditions that determine the transport pathways (and associated wet removal processes during the transport). The small seasonal contrasts in BC emissions from the major source regions (see Table 1) that are used in the model simulation do not seem to be able to explain the large seasonal difference in BC deposition, although the BC emissions are known to have large uncertainties.

Our further analysis shows that the ratio of OC to BC also has clear seasonal dependence. In Figure 3, the slope of the fitted line to measured OC versus BC concentrations during monsoon season is ~6.3, which is twice the slope for non-monsoon season (~3.2). The analysis of covariance (ANCOVA) for slope differences of single linear regressions of OC against BC between monsoon and non-monsoon seasons indicates that the seasonal dependence of the relationship between the concentrations of OC and BC is significant (at the 0.05 significance level). This also agrees with measurements derived from the ice core drilled from
the Palong-Zanbu No. 4 Glacier (Xu et al., 2009b) and in atmospheric samples collected from Lulang, southeastern Tibetan Plateau (Zhao et al., 2013b). The seasonal dependence of the OC/BC ratio can possibly be derived from the seasonal sources of carbonaceous particles, circulation strength, transport pathways, and/or atmospheric deposition processes. Compared to the respective BC and OC concentrations, the seasonal dependence of OC/BC ratio is less straightforward to understand. Circulation patterns together with wet removal processes still determine the transport pathways of emissions from major BC and OC source regions to the sampling site, which however are less likely to change OC/BC ratio from certain sources. Therefore, it is more plausible due to seasonally dependent contributions from source regions and/or emission sectors (including fuel types, quantity, and combustion conditions). Cao et al. (2005) found that the average OC/BC ratios measured from plumes of residential biomass burning and coal combustion are substantially higher than from vehicle exhaust. Higher OC/BC ratio during summer monsoon might indicate more contributions from biomass and/or coal burning than fossil fuel combustion.

3.2 Source attribution

To quantitatively attribute the source of BC at the drilling site (as a receptor region), we use the CAM5 model with the BC source tagging capability to conduct an 11-year simulation, with the last 10 years (1996-2005) used for analysis. The surrounding area is divided into four source regions (see Table 1 and Figure 4): South Asia, East Asia, Southeast Asia and Central Asia. BC emissions from each of the four regions and the rest of the world are explicitly tracked, so that the fractional contributions by emissions from the individual source regions to BC deposition at the receptor region can be explicitly calculated. Figure 4 shows the spatial distribution of fractional contribution from the four source regions. BC deposition at the drilling site (indicated by the black box in Figure 4), which has a consistent
seasonal dependence (i.e., more during the non-monsoon season; Figure 5) with ice
core measurements, is predominately (over 95%) from South Asia and East Asia.
The seasonal dependence of BC deposition is also consistent with a recent regional
climate modeling study on BC deposition on the Himalayan snow cover from 1998
to 2008 (Ménégoz et al., 2014).

The 10-year (1996-2005) average wind fields (at the surface and 500 hPa from
MERRA reanalysis datasets), as shown in Figure 2, indicate distinct circulation
patterns during summer monsoon (June-September) and non-monsoon
(October-May) season, which in part determine the seasonal dependence of transport
of aerosols emitted from the different major sources. During the non-monsoon
season, strong westerly dominates the transport from west to east at all levels.
Emissions from northern India and central Asia can have influence on BC in the
direct downwind receptor region over southeastern Tibetan Plateau. During the
summer monsoon season, the westerly moves northward and the monsoon flow from
Bay of Bengal at the surface and middle levels (e.g., 500hPa), coupled with the
monsoon from Indochina peninsula and South China Sea, exert influence on BC in the
receptor area. The strong monsoon precipitation removes BC from the
atmosphere during the transport. The high Himalayas can partly block the further
transport of emissions from South Asia to Tibetan Plateau, although small local
topographical features such as the Yarlung Tsangpo River valley can provide a gate
for the pollution to enter the inner Tibetan Plateau (Cao et al., 2010). Elevated
emissions from the west (or northern part of South Asia) can take the pathways at
middle and upper levels but they have minimal contribution to deposition. Therefore,
BC emissions from East Asia play a relatively more important role affecting
deposition at the Zuoqipu site during the monsoon season.

The fractional contributions to 10-year mean BC deposition at the drilling site
from the four tagged regions are summarized in Table 1. Results show that South
Asia is the dominant contributor (~81%) during the non-monsoon season with ~14%
from East Asia, while the contribution of East Asia (~56%) is larger than that of South Asia (~39%) during the monsoon season. For the annual mean BC deposition, South Asia (~75%) is the biggest contributor, followed by East Asia (~21%). Emissions from the central Asia and Southeast Asia regions have much smaller contributions (<3%) for all seasons. These results agree well with the short-term source attribution study by Lu et al. (2012) using the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model.

For comparison, seasonal and annual mean BC emissions from the individual tagged source regions are also included in Table 1. Apparently, the contrast in strengths of regional emissions alone cannot explain their relative contributions to BC deposition at the sampling site, and the small seasonal variations in emissions are unlikely the cause of seasonal dependence of source attribution. Note that the BC emission inventory (Lamarque et al., 2010) used in CAM5 doesn’t consider seasonal variations in anthropogenic emissions, which is likely to have introduced biases in the quantitative model estimates of seasonal dependence of contributions, but the relative importance of source regions should be robust.

### 3.3 Interannual variations and long-term trend

Based on annual snow accumulation and BC and OC concentrations derived from the ice record, the annual BC and OC deposition fluxes can be estimated, which are then used to examine the interannual variations and long-term trend in the fluxes and the ratio of OC/BC, as well as the relationship with emissions from the major contributor. As illustrated in Figure 6, from late 1950s to 1980, the BC and OC fluxes in the Zuoqipu ice core are relatively low and stable in comparison to those after 1980. During the period 1956 to 1979, average fluxes are 9.1 and 28.7 mg m\(^{-2}\) a\(^{-1}\) for BC and OC, respectively. Both BC and OC fluxes began to show increasing trends from early 1980s. These trends continued in the early 1990s but started to drop in the mid-1990s, reaching a minimum in 2002 followed by a rapid
increase. In 2006, BC and OC fluxes are 19.2 and 93.9 mg m$^{-2}$ a$^{-1}$, respectively, which are two and three times the respective average fluxes before 1980. The five-year average OC/BC flux ratio is steady before 1990; however, it shows a continual increase afterwards and has been higher than the average value (3.2) for the period of 1956-1979 since mid-1990s (Figure 6). The 10-year CAM5 model simulation, in which annual emissions are fixed but meteorological conditions vary, shows no increasing trend in BC and OC deposition fluxes (BC deposition shown in Figure 5), indicating that the increasing trend seen in the observations was not due to changes in meteorology.

As shown in the CAM5 model simulation, the annual mean atmospheric deposition of BC over southeastern Tibetan Plateau is mostly contributed by emissions from South Asia, particularly, in the non-monsoon season. The BC and OC deposition fluxes derived from the ice-core measurements may reflect changes in South Asian emissions to some extent. The temporal variations of BC and OC deposition fluxes (see Figure 6) are compared with the primary BC and OC emissions from fossil fuel and biofuel combustion in South Asia during 1955-2000 (Bond et al., 2007). BC and OC emissions during 1996-2010 from Lu et al. (2011) are also illustrated in Figure 6 to extend the emission data to cover the entire time period that the ice core data span. Note that the emission data from Lu et al. (2011) are only for India, which is the largest energy consumer and carbonaceous aerosol-emitting country in South Asia. There are differences between the emissions of Bond et al. and Lu et al. during the overlap time period (1996-2000). However, good agreements on the increasing trend can be found in the respective deposition fluxes and emissions of BC and OC (Figure 6). The OC/BC emission ratio also shows an increasing trend from the late 1990s to 2003, which is consistent with that of OC/BC ratio in the ice core record. The annual mean aerosol index over industrial and populated cities in the northern part of India increased from 1982-1993 and more significantly from 2000-2003 (Sarka et al., 2006). This trend is similar to that
of carbonaceous aerosols in the ice core record, and it might indicate a causal relationship between BC and OC over southeastern Tibetan Plateau and emissions from north part of South Asia.

3.4 Emission source analyses

BC and OC in the atmosphere are co-emitted from a variety of natural and anthropogenic sources, including combustion of fossil fuel, biofuel and/or biomass burning. In general, open biomass burning typically produces more abundant OC (i.e., larger OC/BC ratio) compared to fossil fuel combustion due to a lower process temperature (Ducret and Cachier, 1992). The OC/BC ratio has often been used to discriminate fossil fuel combustion and biomass burning emissions in the atmosphere and in precipitation (Novakov et al., 2000; Stone et al., 2007; Ducret and Cachier, 1992; Xu et al., 2009b). For example, Cao et al. (2005) collected particulate matter samples from the plumes of residential biomass burning, coal combustion, and motor-vehicle exhaust sources, and analyzed OC and BC with DRI Thermal/Optical Carbon Analyzer (Model 2001). They reported average OC/BC ratios of 60.3, 12.0, and 4.1 for biomass burning, coal-combustion and vehicle exhaust, respectively. The increasing OC/BC ratios based on the ice core measurements since the early 1990s (Figure 6) suggest an expanded coal consumption and/or usage of biomass fuel, although the ratios might have been underestimated because water-soluble OC was not captured in the sample analyses. However, such bias would have occurred to all the samples and had little impact on the trend, unless including water-soluble OC could dominate the temporal variation of OC/BC ratio. Otherwise, our results indicate that the relative contribution of coal combustion and biomass burning to the carbonaceous particles deposited into the ice core in southeastern Tibetan Plateau has been increasing faster than the contribution of fossil fuel combustion since early 1990s. Improved combustion technologies may have reduced both BC and OC emissions from the combustion of the same amount
of fuels, but the influence on OC/BC ratio is unclear. Presumably improved combustion technologies after 1990 in South and East Asia did not dominate the OC/BC ratio.

The temporal variations of BC and OC in the Zuoqiupu ice core, along with the source attribution analysis of the CAM5 model results, suggest an increasing trend in emissions and altered emission sources in South Asia during the late 20th century. Coal has been the primary energy source in South Asia. For example, in India coal accounted for 41% of the total primary energy demand in 2007, followed by biomass (27%) and oil (24%) (IEA, 2009). The consumption data of coal and crude oil in South Asia (BP Group, 2009) is compared with the BC and OC fluxes in Figure 6 (bottom right). Coal consumption had an increasing trend from 1965 to 2008, particularly in the two time periods 1980-1995 and 2003-2008 after a level off during 1996-2002. This trend is consistent with the variations of BC and OC deposition fluxes in the Zuoqiupu ice core. The correlations between coal consumption and BC ($R^2 = 0.43, p < 0.001$) and OC ($R^2 = 0.62, p < 0.001$) in the ice core are both statistically significant. The oil consumption had a comparable increasing trend as coal before it slowed down during 2000-2006.

Biomass is the second largest energy resource in South Asia, and it is essential in rural areas. In India, 70% of the population lives in rural areas, and depends substantially on solid fuels (i.e., firewood, animal dung, and agriculture residues) for cooking and heating (Heltberg et al., 2000). Even in urban areas, biomass contributes to 27% of the household cooking fuel (Venkataraman et al., 2010). Although the consumption of biomass is lower than coal, the OC/BC emission ratio for biomass burning is much higher than from coal combustion (60.3 vs. 12.0) (Cao et al., 2005). BC emission factor for biomass burning (varying from 0.48 ± 0.18 g kg$^{-1}$ for savanna and grassland burning to 1.5 g kg$^{-1}$ for charcoal burning) is also generally higher than that for coal (0.2 g kg$^{-1}$ for most combustion conditions) and oil combustion (0.3 g kg$^{-1}$ on average, varying from 0.08 g kg$^{-1}$ for heavy fuel oil to
Therefore, it is very likely that the OC/BC ratio of atmospheric carbonaceous aerosols and in the ice-core samples (Figure 6) was dominated by biomass burning emissions. Previous studies have concluded that carbonaceous aerosol emissions from biomass burning are the largest source in South Asia (Venkataraman et al., 2005; Gustafsson et al., 2009). A general increase in energy-intensive life-styles associated with the accelerated growth of population and economy put pressure on energy resources, and induced energy transitions and use of non-sustainable biomass in South Asia (Sathaye and Tyler, 1991; Pachauri, 2004; Fernandes et al., 2007). For instance, biofuel consumption in South Asia increased by 21% per decade on average during 1950-2000 (Bond et al., 2007; Fernandes et al., 2007). In addition, fuel wood, a more desirable biofuel option, contributed 68% in 1978 to total energy demand by rural populations in India, and increased to 78% in 2000 (Fernandes et al., 2007).

3.5 Radiative forcing induced by carbonaceous aerosols in Tibetan Glaciers

BC is often the most important light-absorbing impurity in surface snow because of its strong absorption of solar radiation. Effect of BC in snow on surface albedo reduction and resultant positive radiative forcing have been widely addressed and reported (e.g., Warren and Wiscombe, 1980; Clarke and Noone, 1985; Hansen and Nazarenko, 2004; Hadley and Kirchstetter, 2012; Flanner et al., 2007; 2009; McConnell et al., 2007; Ming et al., 2008; Kaspari et al., 2011; Qian et al., 2011, 2014, 2015). In contrast, the impact of OC in snow has not been widely assessed because of its relatively weak light-absorption over the entire spectrum compared to BC, and because of large uncertainties associated with OC light-absorbing properties and measurements of OC in snow. However, there have been increasing interests in light-absorbing OC (a.k.a. brown carbon) and its radiative effect in both the atmosphere and snow. A growing number of studies (e.g., Kirchstetter et al., 2004; Andreae and Gelencsér, 2006; Hoffer et al., 2006; Moosmüller et al., 2009; Yang et al., 2009; Lack and Cappa, 2010; Cheng et al., 2011). Hoffer et al. (2006) estimated that humus-like substances as part of OC from biomass burning contribute ~7% to the absorption over the entire spectrum, which is not negligible. Yang et al. (2009) highlighted that as the contribution to absorption from BC decreases towards the ultra violet wavelengths, absorption due to brown carbon and dust becomes more significant, and they reported that at an observation site near Beijing brown carbon contributes over 10% to total absorption at mid-visible wavelengths.
2004; Andreae and Gelencsér, 2006; Hoffner et al., 2006; Yang et al., 2009; Kirchstette and Thatcher, 2012) have reported that airborne brown carbon can contribute significantly to aerosol light absorption in the atmosphere, although there are still substantial uncertainties in quantifying optical properties of brown carbon, which makes the model estimation of OC radiative forcing difficult. Similarly, the importance of OC absorption in snow has been recognized and suggested for inclusion in modeling aerosol snow-albedo effect (e.g., Flanner et al., 2009; Aoki et al., 2011). Observational analysis of light-absorbing particles in Arctic snow reported that the main non-BC component is brown carbon, which accounted for 20-50% of the visible and ultraviolet absorption (Hegg et al., 2009, 2010; Doherty et al., 2010). In the rural area of central north China, brown carbon in winter snow also played an important role in visible light absorption, which contributed about 60% to light absorption at 450 nm and about 40% at 600 nm (Wang et al., 2013). A more recent observational study by Dang and Hegg (2014) qualified the light absorption by different light-absorbing particulates in snow, and suggested that humic-like substances and polar OC contributed 9% and 4% to the total light absorption respectively. Despite the substantial uncertainties in brown carbon optical properties, a recent global modeling study (Lin et al., 2014), in which a range of optical properties of brown carbon taken from the literature were applied to OC-in-snow concentrations simulated in a global chemical transport model, showed that the global OC forcing in land snow and sea ice is up to 24% of that caused by BC. Thus the contribution of OC in snow to the surface albedo reduction is likely to be important, which has also been considered in recent climate modeling studies (Qian et al., 2015).

In this study, we use the SNICAR-online model (available at http://snow.engin.umich.edu/; Flanner et al., 2007) to estimate radiative forcing induced by the observed BC as if they were present in snow. Detailed description of the SNICAR model has been documented by Flanner and Zender (2005, 2006) and...
Flanner et al. (2007). Here we only briefly describe the setup of input parameters required for running the SNICAR model. A mass absorption cross-section (MAC) of 7.5 m² g⁻¹ at 550 nm for uncoated BC particle (Bond and Bergstrom, 2006) is assumed to be same as the default value, and thus one of the input parameters for adjusting the MAC value in the online SNICAR model, MAC scaling factor, is set to 1. According to the previous studies (Cuffey and Paterson, 2010; Wiscombe and Warren 1980) and measurements in Qiyi glacier and Zuoqiupu glacier, an effective radius of 100 µm with density of 60 kg m⁻³ for new snow, and the effective radius of 400 µm with density of 400 kg m⁻³ for aged snow are adopted for the forcing calculation. As we focus on the estimation of radiative forcing by carbonaceous particles, other impurity contents, such as dust and volcanic ash, are set to be zero. The annual mean BC concentration during 1956-1979 was 4.4 ng g⁻¹, and increased to 12.5 ng g⁻¹ in 2006. As a consequence, the annual mean radiative forcing induced by BC in snow, as calculated by the SNICAR model, nearly proportionally increased from 0.75 W m⁻² to 1.95 W m⁻². Our estimate of mean BC forcing is lower than the estimated Eurasian radiative forcing (2.7 W m⁻²) in spring (Flanner et al., 2009), but it’s comparable to that in the East Rongbuk glacier over Himalayas, which was in the range of 1-2 W m⁻² (Ming et al., 2008). Kaspari et al. (2009) reported a three-fold increase in radiative forcing from BC in snow over Himalayas after 1975, which is consistent with the increasing trend in our results.

The SNICAR model currently does not support the calculation of OC-in-snow forcing in the same way as that for BC due to a lack of reliable OC optical properties that span the dimensions of snow grain size and OC particle size [personal communication with Mark Flanner, 2014]. We take a MAC value of 0.6 m² g⁻¹ at 550 nm for OC (Kirchstetter et al., 2004), and assume a constant factor of 0.08 (i.e., 0.6/7.5) to scale down MAC values of BC at all wavelengths to obtain a first-order guess of OC-in-snow forcing using SNICAR. The estimated OC forcing has a 4-fold increase from 0.2 W m⁻² (for mean OC concentration of 13.8 ng g⁻¹ during...
1956-1979) to 0.84 W m\(^{-2}\) (for mean OC concentration of 61.3 ng g\(^{-1}\) in 2006), which are 27% and 43% of corresponding BC-in-snow forcing, respectively. The BC/OC forcing ratios based on our simple guesses are larger than the upper bound of estimates (i.e., 24%) by Lin et al. (2014).

Two main assumptions could have caused our first-order estimate of OC forcing to have large biases. First, the MAC value of 0.6 m\(^2\) g\(^{-1}\) (at 550 nm) was based on OC extracted from biomass burning samples that tends to have higher absorption efficiency than OC emitted from fusil fuel combustion (Kirchstetter et al., 2004). This may cause an overestimation of OC forcing. Second, we treated all the water-insoluble OC from the ice-core measurements as light-absorbing brown carbon in the forcing estimation, which also likely results in an overestimation of OC forcing if a significant fraction of OC is non-absorbing. However, water-soluble part, accounting for about half of OC observed in Manora peak and northwest India (Ram et al., 2010; Rajput et al., 2013), can also contribute to some absorption of UV and visible light (Chen and Bond, 2010; Beine et al., 2011). Thus the absorption by water-soluble OC that was not included in the forcing estimate may compensate for the high bias to some extent. According to a laboratory study by Chen and Bond (2010), a large fraction of absorbing OC from hard wood burning is water-insoluble. As water-insoluble OC recorded in the ice core herein was very likely dominated by biomass burning emissions (Section 3.4), the second assumption we used here may not cause a huge bias in estimating OC forcing in snow.

It is also important to note that we didn't consider variations in chemical compounds of OC, the changes of OC during sample filtration, and the different spectral dependence of OC and BC absorption. Although such uncertainties can also cause bias in the estimation of OC radiative forcing herein, the increasing trend should be robust.

BC and OC concentrations in the ice core increased rapidly since 1980, and the
induced radiative forcing rose as a consequence. According to the estimates using the SNICAR model, the average BC radiative forcing had increased 43% after 1980, and OC radiative forcing had an increase of 70%. These numbers are by no means accurate, but the stronger increasing trend in the ice core recorded OC than BC during 1990-2006 (Figure 6) suggests that the contribution of OC to the total radiative forcing in the glacier induced by snow/ice impurities deserves more attention.

4. Summary and Conclusions

Light-absorbing carbonaceous aerosols can induce significant warming in the atmosphere and in snow and glaciers, which likely accelerates the melting of glaciers over Himalayas and Tibetan Plateau. Ice-core measurement of carbonaceous aerosols is a useful mechanism for evaluating historical emission inventories and revealing long-term changes in anthropogenic aerosols and their impacts on regional climate. In this study, we analyze carbonaceous aerosols recorded in an ice core (97 meters in depth and 9.5 cm in diameter) retrieved from the Zuoqiiupu glacier (96.92°E, 29.21°N, 5600 m above sea level) in the southeastern Tibetan Plateau for their seasonal dependence and long-term trend. The glacier has a unique geographical location that is in close proximity to major Asian emission sources. With the help of a global climate model (CAM5) in which black carbon (BC) emitted from different source regions can be explicitly tracked, we are able to characterize BC source-receptor relationships between four Asian source regions (i.e., South Asia, East Asia, Southeast Asia and Central Asia) and the Zuoqiiupu glacier area as a receptor. We also estimate the radiative forcing in snow due to BC and OC using the ice core measurements and an offline snow-ice-aerosol-radiation model (called SNICAR).

BC and OC concentrations in small segments of the Zuoqiiupu ice core were measured using a thermal-optical method. Ice core dating based on significant
seasonal variations of oxygen isotope ratios ($\delta^{18}$O) was used to construct the time series of BC and OC concentrations, which turned out to span the time period of 1956–2006. Not only do the concentrations of OC and BC in the ice core exhibit significant differences between the summer monsoon and non-monsoon seasons, which is likely due to changes in transport pathways and wet removal, but also the ratio of OC to BC shows a clear seasonal dependence that might be due to seasonal change in contributions from source regions and/or emission sectors. The CAM5 results show a similar seasonal dependence of BC and OC deposition to the glacier.

The MERRA reanalysis products used to drive the CAM5 model simulation show distinct circulation patterns during summer monsoon (June-September) and non-monsoon (October-May) seasons. Both the circulation patterns (and associated aerosol transport and wet removal) and seasonal variation of emissions in major source regions influence the seasonal deposition of aerosol at the Zuoqiupu site. The CAM5 simulation with tagged BC regional sources shows that South Asia is the dominant contributor (81%) to the 10-year mean BC deposition at the Zuoqiupu site during the non-monsoon season with 14% from East Asia, while the contribution of East Asia (56%) is larger than that of South Asia (39%) during the monsoon season. For the annual mean BC deposition, South Asia (75%) is the biggest contributor, followed by East Asia (21%).

The annual mean BC and OC deposition fluxes into the ice core are also estimated to explore the interannual variations and long-term trends. Results show stable and relatively low BC and OC fluxes from late 1950s to 1979, followed by a steady increase through the mid-1990s. A more rapid increase occurred after the minimum in 2002. The BC and OC deposition fluxes in 2006 were two and three times the respective average before 1980.

The overall increasing trend in deposition fluxes since 1980 is consistent with the BC and OC emissions in South Asia as the major contributor. Moreover, the increasing trend of OC/BC ratio since early 1990s indicates a growth of the
contribution of coal combustion and/or biomass burning to the carbonaceous aerosol emissions in the major contributing source regions, which is consistent with the trends in the consumption of coal, oil and biomass in South Asia.

Our offline calculation using the SNICAR model shows a significant increase of radiative forcing induced by the observed BC and OC in snow after 1980, which has implications for the Tibetan glacier melting and availability of water resources in the surrounding regions. More attention to OC is merited because of its non-negligible light absorption and the recent rapid increases evident in the ice core record.

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Table 1. Source regions (South Asia, East Asia, Southeast Asia, and Central Asia) and corresponding monthly mean BC emissions (Tg a\(^{-1}\)) and fractional contributions (%) to BC deposition flux at the Zuoqiupu site in monsoon (June-September), non-monsoon (October-May), and all months during 1996-2005.

<table>
<thead>
<tr>
<th>Source Regions</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Monsoon Contribution</th>
<th>Emission</th>
<th>Non-monsoon Contribution</th>
<th>Emission</th>
<th>Annual Contribution</th>
<th>Emission</th>
</tr>
</thead>
<tbody>
<tr>
<td>South Asia</td>
<td>5-35°N</td>
<td>50-95°E</td>
<td>38.51</td>
<td>0.65</td>
<td>81.26</td>
<td>0.74</td>
<td>74.48</td>
<td>0.71</td>
</tr>
<tr>
<td>East Asia</td>
<td>15-50°N</td>
<td>95-150°E</td>
<td>56.24</td>
<td>1.75</td>
<td>13.91</td>
<td>1.90</td>
<td>20.66</td>
<td>1.85</td>
</tr>
<tr>
<td>Southeast Asia</td>
<td>0-15°N</td>
<td>95-130°E</td>
<td>0.05</td>
<td>0.28</td>
<td>0.16</td>
<td>0.33</td>
<td>0.15</td>
<td>0.31</td>
</tr>
<tr>
<td>Central Asia</td>
<td>35-50°N</td>
<td>50-95°E</td>
<td>2.62</td>
<td>0.11</td>
<td>0.86</td>
<td>0.09</td>
<td>1.14</td>
<td>0.10</td>
</tr>
</tbody>
</table>
Figure 1. Site location of Zuoqipu Glacier (top): black circle represents the location of Zuoqipu Glacier, and warm colors indicate high elevations over the Tibetan Plateau. Detailed elevation contours of the Zuoqipu Glacier are shown in the bottom panel. Red circle marks the ice core drill site.
Figure 2. 10-year (1996-2005) mean wind vectors (denoted by arrows) at 500hPa (a, b) and the surface (c, d) during summer monsoon (June-September; a, c) and non-monsoon season (October-May; b, d) from MERRA reanalysis datasets used to drive the CAM5 simulation. 500 hPa Geopotential height (units: 10 m) contours with an interval of 60 m and mean sea-level pressure (units: hPa) contours with an interval of 4 hPa are superimposed on panels (a, b) and (c, d), respectively. The background colors show mean BC emission rates based on the IPCC present-day scenario for the corresponding months. The small black box marks the model grid-cell in which the ice core drill site resides.
Figure 3. Scatter plots for yearly monsoon and non-monsoon mean OC and BC concentrations during 1956-2006, obtained from the ice core measurements, and corresponding linear regressions.

OC = 6.31BC - 3.41
$R^2 = 0.72$

OC = 3.17BC + 0.29
$R^2 = 0.69$
Figure 4. Spatial distributions of fractional contribution from the four source regions (South Asia, East Asia, Southeast Asia, and Central Asia) to monsoon, non-monsoon, and annual mean BC deposition fluxes during 1996-2005. The large black boxes indicate the boundary of source regions, and the small black box marks the model grid-cell where the Zuoqiu site is located. Color in the small black box in each panel corresponds to the fraction contribution to BC deposition at the sampling site. Exact percentage contributions are provided in Table 1.
Figure 5. Seasonal dependence ("NM" for non-monsoon and "M" for monsoon season) of BC deposition flux at the Zuoqiupu site from 1995 to 2005 simulated in CAM5. The dash line represents a linear regression of all data points.
Figure 6 Time series of annual (dotted line with circles) and 5-year averaged (solid line) OC/BC ratios (top-left), BC (top-right) and OC deposition fluxes (bottom-left) based on the Zuoqipu ice core measurements for the time period of 1956-2006. The average values of OC/BC ratio, BC and OC during 1956-1979 are marked by dashed lines. BC and OC emissions in South Asia (Bond et al., 2007) and corresponding OC/BC emission ratios are illustrated with gray triangles, and with red diamonds for emissions in India (Lu et al., 2011). Coal and oil consumption data are shown in the bottom-right panel (BP Group, 2009).