Constraining Black Carbon Aerosol over Asia using OMI Aerosol Absorption Optical Depth and the Adjoint of GEOS-Chem

Li Zhang\textsuperscript{1,2}, Daven K. Henze\textsuperscript{1}, Georg A. Grell\textsuperscript{2}, Gregory R. Carmichael\textsuperscript{3}, Nicolas Bousserez\textsuperscript{1}, Qiang Zhang\textsuperscript{4}, Omar Torres\textsuperscript{5}, Changwook Ahn\textsuperscript{6}, Zifeng Lu\textsuperscript{7}, Junji Cao\textsuperscript{8}, Yuhao Mao\textsuperscript{9,10}

\textsuperscript{1}Department of Mechanical Engineering, University of Colorado, Boulder, CO, USA
\textsuperscript{2}Global Systems Division, Earth System Research Laboratory, NOAA, Boulder, CO, USA
\textsuperscript{3}Department of Chemical and Biochemical Engineering, University of Iowa, Iowa, IA, USA
\textsuperscript{4}Center for Earth System Science, Tsinghua University, Beijing, China
\textsuperscript{5}NASA Goddard Space Flight Center, Greenbelt, MD, USA
\textsuperscript{6}Science Systems and Applications, Inc., Lanham, MD, USA
\textsuperscript{7}Decision and Information Sciences Division, Argonne National Laboratory, Argonne, IL, USA
\textsuperscript{8}Key Lab of Aerosol Chemistry & Physics, Institute of Earth Environment, Chinese Academy of Sciences, Xi'an, China
\textsuperscript{9}Department of Atmospheric and Oceanic Sciences, University of California, Los Angeles, CA, USA
\textsuperscript{10}State Key Laboratory of Atmospheric Boundary Layer Physics and Atmospheric Chemistry, Institute of Atmospheric Physics, Chinese Academy of Sciences, Beijing, China

Submitted to Atmospheric Chemistry and Physics

May 2015

*Corresponding author:
daven.henze@colorado.edu
Dept. of Mechanical Engineering
1111 Engineering Drive ECES 114
University of Colorado, Boulder
Boulder, CO 80309
Abstract

Accurate estimates of the emissions and distribution of black carbon (BC) in the region referred to here as Southeastern Asia (70°E–150°E, 11°S–55°N) are critical to studies of the atmospheric environment and climate change. Analysis of modeled BC concentrations compared to in situ observations indicates levels are underestimated over most of Southeast Asia when using any of four different emission inventories. We thus attempt to reduce uncertainties in BC emissions and improve BC model simulations by developing top-down, spatially resolved, estimates of BC emissions through assimilation of OMI observations of aerosol absorption optical depth (AAOD) with the GEOS-Chem model and its adjoint for April and October of 2006. Overwhelming enhancements, up to 500%, in anthropogenic BC emissions are shown after optimization over broad areas of Southeast Asia in April. In October, the optimization of anthropogenic emissions yields a slight reduction (1~5%) over India and parts of southern China, while emissions increase by 10~50% over eastern China. Observational data from in situ measurements and AERONET observations are used to evaluate the BC inversions and assess the bias between OMI and AERONET AAOD. Low biases in BC concentrations are improved or corrected in most eastern and central sites over China after optimization, while the constrained model still underestimates concentrations in Indian sites in both April and October, possibly as a consequence of low prior emissions. Model resolution errors may contribute up to a factor of 2.5 to the underestimate of surface BC concentrations over northern India. We also compare the optimized results using different anthropogenic emission inventories and discuss the sensitivity of top-down constraints on anthropogenic emissions with respect to biomass burning emissions. In addition, the impacts of different observation operators and a priori constraints on the optimization are investigated. Overall, despite these limitations and uncertainties, using OMI AAOD to constrain BC sources improves model representation of BC distributions, particularly over China.
1. Introduction

Black carbon (BC) is a product of incomplete combustion of carbonaceous fuels, enhanced concentrations of which have led to a present-day overall positive radiative forcing and climate warming [Charlson and Pilat, 1969; Satheesh and Ramanathan, 2000; Bond et al., 2013]. More than ten years ago, Jacobson [2000] and Hansen et al. [2000] recognized that preindustrial to present increases in BC might warm the atmosphere about one third as much as CO$_2$. Recently, an assessment report by Bond et al. [2013] indicates that the global average preindustrial to present radiative forcing from BC is +1.1 W/m$^2$ with 90% uncertainty bounds of +0.17 to +2.1 W/m$^2$, which is more than two thirds that of CO$_2$ (+1.56 W/m$^2$). Additionally, BC aerosols constitute up to 10-15% of the mass concentration of fine particulate matter (PM$_{2.5}$) over continental regions, exposure to which is known to adversely effect human health [e.g., Janssen et al., 2005; Schwartz et al., 2008; Janssen et al., 2011]. Given the magnitude of BC climate effects and health impacts, a number of studies have investigated its direct effect [Forster 2007; Ramanathan and Carmichael, 2008], semi-direct effect [Ackeman et al., 2000; Johnson et al., 2004], indirect effect [Cozic et al., 2007; Liu et al., 2009; Oshima et al., 2009], and the albedo effect when deposited on snow [Hansen and Nazarenko, 2004; Hansen et al., 2005; Flanner et al., 2007; Qian et al., 2009] using various numerical models and observations.

Central estimates of global annual emissions of BC are 8.0 Tg, of which 38% comes from fossil fuel, 20% from biofuel and 42% from open burning [Bond et al., 2004]. At the same time, estimates of BC emissions are recognized as having large uncertainties -- 50% at global scales and a factor of two to five at regional scales.
[Bond et al., 2004; Ramanathan and Carmichael, 2008]. The Asian region referred to here as Southeast Asia (70°E–150°E, 11°S–55°N) is the major anthropogenic BC source region in the world, with growth in BC emissions of 21% over China and 41% over India from 1996 to 2010 associated with rapid economic and industrial development [Lu et al., 2011]. BC emissions from both energy-related combustion and biomass burning that occur largely in Asia and Africa currently appear underestimated [Bond et al., 2013]. A global top-down estimate of BC emission using AERONET observation by Cohen and Wang [2014] indicated that commonly used global BC emissions datasets may be underestimated by a factor of two or more. Sixteen models from the AeroCom aerosol model intercomparisons underestimated the Southeast Asian BC surface concentrations by a factor of 2~3 [Koch et al., 2009]. The GEOS-Chem model also underestimated monthly BC concentrations at almost all rural sites in China, particularly in January 2006, which indicated a regional underprediction of carbonaceous aerosol sources associated with anthropogenic activities [Fu et al., 2012; Wang et al., 2013]. In addition, the global atmospheric absorption attributable to BC is too low in many global aerosol models by a factor of almost three on a global mean basis, which can be attributed to the models lacking treatment of enhanced absorption caused by mixing of BC with other constituents and the amount of BC in the atmosphere [Koch et al., 2009; Bond et al., 2013]. On the other hand, a typical fresh particle mass absorption cross section (MABS, essentially the column BC absorption divided by the load) of about 7.5 m² g⁻¹ recommended by Bond and Bergstrom [2006] is not represented in most models, which should
probably increase as particles age [Koch et al., 2009]. This bias would also impact simulated AAOD, and inferences about emissions based on such comparisons would likewise be biased.

To reduce uncertainties in BC emissions and improve poor representation of BC in model simulations, different top-down approaches have been used to constrain bottom up BC emissions, such as the linear constraints between concentrations and emissions [Park et al., 2003; Kondo et al., 2011; Fu et al., 2012; Wang et al., 2013], inverse modeling using the decoupled direct method [Hu et al., 2009a; Hu et al., 2009b], the Kalman filter technique [Cohen and Wang 2014], and the adjoint based 4D variational approach [Hakami et al., 2005]. These studies have exclusively used in situ measurements or airborne observations, which can provide accurate observations of aerosol properties. However, they are often incomplete in their spatial or temporal coverage. Satellite measurements of aerosol optical depth (AOD) have much broader temporal and spatial coverage, and have also been used to constrain BC sources [Huneeus et al., 2003; Xu et al., 2013]. However, AOD reflects the contribution from all aerosol components, making it difficult to distinguish and quantify different aerosol species, especially their relative fractions.

The OMI aerosol absorption optical depth (AAOD), the non-scattering part of the AOD, is an atmospheric column measurement of absorbing aerosol particles, i.e., absorbing carbon and mineral dust, which provides a different perspective to constrain BC sources [Torres et al., 1998; Koch et al., 2009]. In this study, the GEOS-Chem adjoint model and satellite observations of OMI AAOD are used to
constrain spatially resolved BC emissions. Our study will focus on April and October to compare times when the dust loading is relatively large and small over Southeast Asia. Section 2 describes the observations, emissions, and forward and inverse model used in this study. Then we quantify discrepancies between observations and model estimates based on different BC anthropogenic emissions in Section 3. Section 4 describes how formulation of the inverse problem affects the results; evaluation of the inversion results with different prior emission inventories and independent observations are presented in Section 5, and we end with discussion and conclusions in Section 6.

2. Data and Models

2.1 Observations

2.1.1 OMI AAOD

The Ozone Monitoring Instrument (OMI) aboard Aura is a nadir-viewing, wide-swath hyper-spectral imaging spectrometer that provides daily global coverage with high spectral resolutions and spatial resolution of $13 \times 24$ km$^2$ at nadir [Levelt et al., 2006a]. It detects backscattered solar radiance in the ultraviolet-visible wavelengths (0.27 to 0.5 µm) to measure aerosols, clouds, surface UV irradiance, and trace gases [Levelt et al., 2006b]. OMI takes advantage of the greater sensitivity of radiances measured at the top-of-atmosphere in the near-UV region to the varying load and type of aerosols to derive extinction AOD, single scattering albedo (SSA), and AAOD using an inversion procedure at 354, 388 and 500 nm generated by the near-UV
(OMAERUV) algorithm [Torres et al., 2007]. The optical depths at 388 nm are inverted from radiance observations while the 354 and 500 nm results are obtained by conversion of the 388 nm retrievals. The OMAERUV retrieval algorithm is particularly sensitive to carbonaceous and mineral aerosols. It assumes that the column aerosol load can be represented by one of three types of aerosols and uses a set of aerosol models to account for the presence of these aerosols: carbonaceous aerosol from biomass burning, desert dust, and weakly absorbing sulfate-based aerosols. Each aerosol type is represented by seven aerosol models of varying single scattering albedo, for a total of twenty-one models. The twenty-one aerosol models used by OMAERUV are based on long-term statistics of ground-based observations by the AERONET. The major factor affecting the quality of aerosol products is sub-pixel cloud contamination, while AAOD is probably less affected by cloud contamination due to a partial cancellation of cloud effects on the retrieved AOD and SSA co-albedo. Due to the large sensitivity of the OMI near UV observations to particle absorption, the AAOD is the most reliable quantitative OMAERUV aerosol parameter, especially over land. The root-mean-square error for AAOD is estimated to be ~0.01\(^1\). In this study, we used the OMAERUV Level-2 aerosol data product that includes the quality assurance flag, thus only the most reliable retrievals minimally affected by sub-pixel cloud contamination are used [Ahn et al., 2014]. Important algorithm improvements have been implemented in the current OMAERUV algorithm. The carbonaceous aerosol model was replaced with a new model that

\(^1\)daac.gsfc.nasa.gov/Aura/data-holdings/OMI/documents/v003/OMAERUV_README_V003.doc
accounts for the presence of OC while the previous aerosol model only assumed black
carbon as the absorbing component [Jethva and Torres, 2011]. In the revised
algorithm, the identification of aerosol type has been improved by taking advantage
of the Atmospheric Infrared Sounder (AIRS) carbon monoxide (CO) observations in
conjunction with OMI UV-AI. The aerosol layer height (ALH) value is taken from a
climatology derived from CALIOP (Cloud-Aerosol Lidar with Orthogonal
Polarization) observations specifically produced for this purpose [Torres et al., 2013].
The Level 2 OMI AAOD data reports a set of retrieved parameters for different
assumptions of the altitude of the aerosol center of mass: at the surface, and at 1.5,
3.0, 6.0 and 10.0 km above the surface [Torres et al., 2005]. A best-guess set of
retrieved values of AOD, AAOD and SSA associated with the climatological ALH
value from the CALIOP-based climatology is reported as the standard OMAERUV
aerosol product. When the aerosol layer height is not available from CALIOP
climatology, the height is obtained as in the previous version of the algorithm based
on a climatology of GOCART model simulated aerosol heights. For carbonaceous and
desert dust particles, the aerosol load is assumed to be vertically distributed following
a Gaussian function characterized by peak (aerosol layer height) and half-width
(aerosol layer geometric thickness) values [Torres et al., 2005; Torres et al., 2013].
The retrieval values of AAOD are much larger if using the aerosol layer altitude
where more absorbing aerosols are loaded. In general, when comparing satellite
retrievals of trace gases with other measurements or model simulations, it is essential
to take into account the different sensitivities of the instruments by applying
averaging kernels [Luo et al., 2007; Worden et al., 2007]. However, there is no
averaging kernel for OMI AOD/AAOD retrievals. It is thus important to consider
differences in aerosol properties and distributions used in the retrieval algorithm with
those in the assimilation model (e.g., GEOS-Chem). The retrieval “Final AAOD”
products (OMI_Final) are interpolated values using the aerosol layer height value
given by the CALIOP climatology [Torres et al., 2013].

OMAERUV retrievals of AOD and SSA have been evaluated by comparison to
independent ground-based observations provided by the world-wide Aerosol Robotic
Network (AERONET). OMAERUV AOD retrievals at 380 nm were compared to
AERONET observations [Ahn et al., 2014]. Over 10,000 matched OMAERUV-
AERONET AOD pairs at 44 globally distributed land-locations were analyzed. The
AERONET-OMAERUV analysis reported a high level of agreement between the two
datasets, yielding a correlation coefficient of 0.81, y-intercept of 0.1, and slope of
0.79. Sixty five percent of the analyzed OMAERUV AOD data agreed with
AERONET measurements within OMAERUV’s stated uncertainty (largest of 0.1 or
30%). The OMAERUV SSA product has also been evaluated using AERONET
retrievals. Jethva et al [2014] compared OMAERUV and AERONET SSA retrievals
using all available AERONET data at 269 sites for the 2005-2013 period. After
accounting for the wavelength difference (AERONET’s 440 nm versus OMAERUV’s
388 nm), it was shown that 50% of the satellite SSA retrievals agree with
AERONET’s values within 0.03, whereas 75% of the matched pairs agree within 0.05
for all aerosol types. The most important source of uncertainty is the effect of sub-
pixel cloud contamination, related to the sensor’s coarse spatial resolution, that causes
AOD and SSA overestimates for cases of low aerosol load, and severely limits the
overall retrieval yield of the algorithm.

In order to obtain a consistent vertical profile between the OMI retrieval and GEOS-
Chem, we use the GEOS-Chem simulated aerosol layer height instead of the
CALIOP-based aerosol layer height climatology to calculate a GEOS-Chem-based
observed AAOD (referred as OMI_GC AAOD) as a linear interpolation of the OMI
observed AAOD values corresponding to different assumed peak heights. Figure 1
shows the differences between OMI_Final and OMI_GC AAOD over Southeast Asia
for April and October 2006. In April, the enhancements from applying the GEOS-
Chem aerosol layer height are quite significant, with 30-50% increases over eastern
China and downwind areas while 20-30% increases over India and southeastern Asia,
since the simulated aerosol layer heights are much lower than those based on
CALIOP. The increases even exceed 60% across broad areas over the tropical ocean.
Some reductions are shown over parts of western China and northern Asia in the
OMI_GC AAOD. In October, the patterns of enhancement and reduction are similar
to those in April, with smaller changes (less than 20%) over broad continental areas.
The most significant differences occur near the major aerosol source regions, such as
eastern China and South Asia. We also evaluate the linearity of the relationship
between aerosol layer height and AAOD from OMI retrievals. We find (not shown)
that there is less than 30% error in linearly interpolating AAOD corresponding to a
specific aerosol layer height from the AAODs corresponding to two other aerosol
2.1.2 AERONET AAOD

The Aerosol Robotic Network (AERONET) is a ground-based instrument network providing a long-term, continuous and readily accessible public domain database of aerosol optical, microphysical and radiative properties [Holben et al., 1998].

AERONET inversion code provides aerosol optical properties (including size distribution, refractive index, and single scattering albedo) in the total atmospheric column derived from the direct and diffuse radiation measured by Cimel sun/sky-radiometers [Dubovik and King, 2000; Dubovik et al., 2000, 2002a, 2002b; Dubovik et al., 2006; Sinyuk et al., 2007].

We use Level 2.0 quality-assured AERONET aerosol inversions data of AAOD at 440 nm. The prefield and postfield calibrations have been applied in these measurements and they were cloud cleared and manually inspected [Omar et al., 2013]. The total uncertainty in the AERONET AOD for field instruments is ±0.1 to ±0.2 and is spectrally dependent with the higher errors (±0.2) in the UV spectral range [Eck et al., 1999]. The retrieved single scattering albedo uncertainties were within 0.03, estimated by Dubovik et al., [2000], with the exception of the 0.44 µm retrievals for the desert dust case when they increased by ~0.09 and 0.07 for low and high aerosol loadings, respectively [Sinyuk et al., 2007]. In this study, only the AAOD data corresponding to AOD values greater than 0.4 are include.

2.1.3 In situ measurements

For the monthly surface BC observation over Southeast Asia, we combine the in situ
measurements of BC concentration based on several published studies [Zhang et al., 2008; Beegum et al., 2009; Moorthy et al., 2013]. Over China, the monthly surface BC concentrations are from 12 sites, including urban sites and rural sites for April and October, 2006, which were based on results of Zhang et al. [2008]. The locations of these 12 sites are shown in Fig. 2. The BC concentrations are analyzed using thermo-chemical analysis from PM$_{10}$ aerosols, which were collected by air sample [Zhang et al., 2008]. The daily BC measurements are only available at the site of Xi’an (XIA). The PM$_{2.5}$ BC concentrations were measured continuously as 5-min averages by quartzfiber filter tape transmission at an 880 nm wavelength with an aethalometer [Hansen et al., 1984]. More details about the measurement methods are described by Cao et al. [2007; 2009].

The measurements of monthly surface BC concentrations for 2006 using aethalometers over India were based on Beegum et al. [2009] and Moorthy et al. [2013], which were carried out in eight sites (see Table 1) covering India and adjacent oceanic regions. Locations of these sites are indicated in Fig. 2. More details about the measurements and sites are described by Beegum et al. [2009]. DEL and KGP represent urban and semi-urban sites in the Indo-Gangetic Plain (IGP). HYD and PUN represent urban locations. TVM is a semi-urban coastal station in the south India; NTL is a high altitude location in the central Himalayas, and MCY and PBR are two island locations representing the Arabian Sea and Bay of Bengal, respectively.

2.2 GEOS-Chem
GEOS-Chem is a global three-dimension chemical transport model driven by assimilated meteorological observations from the Goddard Earth Observing System (GEOS) of the NASA Global Modeling and Assimilation Office (GMAO) [Bey et al., 2001]. We use the nested-grid GEOS-Chem model [Wang et al., 2004; Chen et al., 2009] driven by GEOS-5 meteorological fields with 6-hour temporal resolution (3-hour for surface variables and mixing depths), 0.5° (latitude) × 0.667° (longitude) horizontal resolution over the window of Southeast Asia (70°E–150°E, 11°S–55°N), and 47 vertical layers between the surface and 0.01 hPa. A global simulation with lower resolution of 4° (latitude) × 5° (longitude) provides the lateral boundary conditions to the higher resolution nested-grid simulation every 3 hours.

The original carbonaceous aerosol simulation in GEOS–Chem was developed by Park et al. [2003]. It assumes that 80% of BC and 50% of OC emitted from primary sources are hydrophobic and that hydrophobic aerosols become hydrophilic with an e-folding time of 1.15 days [Park et al., 2003; Chin et al., 2002; Cooke et al., 1999]. Dust in GEOS-Chem is distributed across four size bins (radii 0.1–1.0, 1.0–1.8, 1.8–3.0, and 3.0–6.0 µm) following Ginoux et al. [2004]. The smallest size bin is further divided equally into four sub-micron size bins (with effective radii centered at 0.15, 0.25, 0.4 and 0.8µm) for calculation of optical properties and heterogeneous chemistry [Fairlie et al., 2010; Ridley et al., 2012]. Due to the significant positive biases identified in GEOS-Chem dust simulations both in surface concentration and dust AOD [Fairlie et al., 2010, Ku and Park, 2011; Ridley et al., 2012; Wang et al., 2012], a new emitted dust particle size distribution (PSD) based upon scale-invariant
fragmentation theory [Kok, 2011] with constraints from in situ measurements [Zhao et al., 2010] is implemented in GEOS-Chem to improve the dust simulation [Zhang et al., 2013]. Large discrepancies are reduced between the simulated surface-level fine dust concentration and measurements from the IMPROVE network in the western US during March to May of 2006 [Zhang et al., 2013]. The new PSD also improves the positive biases of AOD over the Asian and African dust source region in April 2006 (See Fig. S1 in supplemental). The wet deposition scheme [Liu et al., 2001] includes scavenging in convective updrafts as well as in-cloud and below-cloud scavenging from convective and large-scale precipitation. Dry deposition is based on the resistance-in-series scheme of Wesely [1989] as implemented by Wang et al. [1998]. The aerosol optical depth at 400 nm is calculated online assuming log-normal size distributions of externally mixed aerosols and is a function of the local relative humidity to account for hygroscopic growth [Martin et al., 2003]. The AAOD of each aerosol species is calculated as [Ma et al., 2012; Cohen and Wang, 2014; Cohen, 2014]

$$\text{AAOD} = \text{AOD} \times (1 - \text{SSA})$$

(1)

where SSA is the single scattering albedo.

2.3 BC Emission Inventories

Emissions of BC from biomass burning sources are taken from version 2 of the GFED 8-day inventory [van der Werf et al., 2006; Randerson et al., 2006]. GFED v2 is derived using satellite observations of active fire counts and burned areas in conjunction with the Carnegie-Ames-Stanford-Approach (CASA) biogeochemical
model. Carbon emissions are calculated as the product of burned area, fuel load and combustion completeness. Burned area is derived using the active fire and 500-meter burned area datasets from the Moderate Resolution Imaging Spectroradiometer (MODIS) as described by Giglio et al. [2006]. We also use a newer version of GFED v3 daily emissions for sensitivity analysis [van der Werf et al., 2010]. Compared to GFED v2, the main update in GFED v3 is the spatial resolution of the global grid is quadrupled from 1° to 0.5°, the native 500-m MODIS daily burned area maps are applied [Giglio et al., 2010], the regional regression trees of GFEDv2 are replaced by a local regression approach in producing the indirect, active-fire based estimates of burned area, and a revised version of Carnegie-Ames Stanford Approach (CASA) biogeochemical model is used.

Global anthropogenic emissions for carbonaceous aerosols (BC/OC) in GEOS-Chem are originally from Bond et al. [2004, 2007], which contain both biofuel and fossil fuel emissions. The estimated BC emissions uncertainties are $-36\%$ to $149\%$ over China and $38\%$ to $-119\%$ for India [Bond et al., 2004; Lu et al., 2011]. In this study, we evaluate three additional carbonaceous anthropogenic emission inventories over Southeast Asia and China: the Streets regional inventory for Intercontinental Chemical Transport Experiment - Phase B (INTEX-B), the Southeast Asia Composition, Cloud, Climate Coupling Regional Study (SEAC4RS) emission inventory, and the Multi-resolution Emission Inventory for China (MEIC, http://www.meicmodel.org/). Anthropogenic emissions are all classified into four major sectors: power generation, industry, residential and transport. The INTEX-B
inventory is based on 2006 and contains monthly variations with $0.5^\circ \times 0.5^\circ$ horizontal resolution over Southeast Asia (Zhang et al., 2009). The SEAC$^4$RS inventory is an annual, finer resolution inventory based on 2012, with $0.1^\circ \times 0.1^\circ$ horizontal resolution over Southeast Asia [Lu et al., 2011]. The average uncertainties of BC are estimated to be $-43\%$ to $90\%$ over China, which are much lower than those of the INTEX-B between $-68\%$ to $308\%$ [Zhang et al., 2009; Lu et al., 2011]. The MEIC emission inventory over China also includes monthly variations and is provided at the $0.5^\circ \times 0.5^\circ$ horizontal resolution. These four anthropogenic emission inventories are regridded to the GEOS-Chem resolution of $0.5^\circ \times 0.667^\circ$, and their annual emissions are shown in Fig. 3. The differences in these inventories exceed 100% across broad areas, especially over India and eastern China. The anthropogenic emission inventory of INTEX-B is comparable to that of MEIC over eastern China while lower than that of Bond and SEAC$^4$RS over western China and India. Both Bond and SEAC$^4$RS inventories are lower over central and eastern China compared to those of INTEX-B and MEIC inventories. With much finer resolution, the SEAC$^4$RS emission inventory indicates more hot spots spread across eastern and central China and the IGP and eastern India where rural population densities are high and residential coal and biofuel combustion are prevalent [Lu et al., 2011].

2.4 GEOS-Chem Adjoint and Inverse Modeling

An adjoint model is a set of equations auxiliary to a forward model that are used to efficiently calculate the gradient of a scalar model response function with respect to all model parameters simultaneously [Lions, 1971]. The adjoint of GEOS-Chem was
developed specifically for inverse modeling including explicit treatment of gas-phase chemistry, heterogeneous chemistry, black and organic primary aerosol, as well as the treatment of the thermodynamic couplings of the sulfate-ammonium-nitrate-formation chemistry [Henze et al., 2007; 2009], with code updates following the relevant parts of the GEOS-Chem forward model up through version v9. The GEOS-Chem adjoint model has been developed and widely used to constrain sources of emission such as dust [Wang et al., 2012], ammonia [Zhu et al., 2013], CO [Kopacz et al., 2009; Kopacz et al., 2010; Jiang et al., 2011], CH$_4$ [Wecht et al., 2012; Wecht et al., 2014], and to investigate pollution transport [e.g., Zhang et al., 2009, Kopacz et al., 2011].

The 4D variational data assimilation technique is used with the GEOS-Chem adjoint model to combine observations and models to calculate an optimal estimate of emissions. A range of emissions are constructed using control variables, $\sigma$, to adjust the vector of model emissions via application as scaling factors with elements $\sigma = \frac{E}{E_a}$, where $E$ and $E_a$ are posterior and prior BC emission vectors, respectively. This method of inverse modeling seeks $\sigma$ that minimizes the cost function, $J$, presented by:

$$J = \frac{1}{2} \sum_{c \in \Omega} (Hc - c_{obs})^T S_{obs}^{-1} (Hc - c_{obs}) + \frac{1}{2} y_r (\sigma - \sigma_a)^T S_a^{-1} (\sigma - \sigma_a)$$  \hspace{1cm} (2),

where $c$ is the vector of species concentrations mapped to the observation space by $H$, the observation operator, $c_{obs}$ is the vector of species observations, $\sigma_a$ is the prior estimate of the scaling factors, $S_{obs}$ and $S_a$ are error covariance estimates of the observations and scaling factors, respectively, and $\Omega$ is the domain over which observations are available. The first term of the cost function in Eq. (2) is the
observation term, which is the total prediction error incurred for departure of model
predictions from the observations. The second term, the a priori term or penalty
(background) term, is the penalty incurred for departure from the prior emissions.
Here $S_\alpha$ is assumed to be diagonal, and the significance of the prior information is
more of a smoothness constraint than a rigorous estimate of prior uncertainty
[Rodgers, 2000]. $\gamma_r$ is a regularization parameter, which used to balance the two terms
[Hansen 1998; Henze et al., 2009]. We will discuss the contributions of the penalty
term in Section 4.2.

Overall, the minimum value of the cost function balances the objectives of improving
model performance while ensuring the model itself remains within a reasonable range
(as dictated by $S_\alpha^{-1}$) of the initial model. The minimum of the cost function is sought
iteratively using the quasi-Newton L-BFGS-B algorithm [Zhu et al., 1994; Byrd et al.,
1995]. This approach requires the gradients of the cost function with respect to the
emission scaling factors at each iteration, which are calculated with the GEOS-Chem
adjoint model.

2.5 Cost function and adjoint forcing

OMI_GC AAOD column observations represent the combined absorption of all
aerosols species (dominated by BC, dust, and to a lesser extent OC). Similarly,
modeled total column AAOD, $T_{GC}$, is the sum of modeled column absorption from
BC ($T_{GC,BC}$), OC ($T_{GC,OC}$) and dust ($T_{GC,Dust}$):

$$T_{GC} = T_{GC,BC} + T_{GC,OC} + T_{GC,Dust} \quad (3).$$

In order to use AAOD observations to develop constraints on BC alone, we must
formulate the observation term of the cost function to isolate the impacts of BC on the
difference between simulated and observed AAOD. Here we consider four
approaches: methods (a) – (d). The first two methods use modeled ratios of BC to
total absorption (either in each layer (a), or the total column (b)) to derive an
“observed” BC AAOD. Method (c) makes a direct comparison between total AAOD
in the model and measurements. Lastly, in method (d), we also consider using a
subset of the OMI data that has been flagged in the retrieval process as being
impacted by carbonaceous aerosol. These different approaches to constructing a cost
function, and the gradient of these cost functions with respect to the vertically
resolved modeled BC concentration (i.e., the adjoint forcing) are presented below.
Here we do not consider the penalty term in the cost function in order most clearly
assess how formulation of the observation term impacts the inversion. The
consequences of the different cost function formulations are described in Section 4.1.
(a): In this method, the observation term of the cost function can be written as:
\[
J = \frac{1}{2} \sum_{i=1}^{N} \left( \tau_{BC_{i}} - \tau_{OMI_{i}} \right)^2 \cdot S^{-2}_{OMI_{i}} \quad (4),
\]
where \( L \) is the top of atmosphere, \( N \) is the total number of observations, and \( \tau_{BC_{i}} \)
and \( \tau_{OMI_{i}} \) are the modeled and observed BC AAODs at layer \( l \) for the \( i^{th} \)
observation, respectively. The latter is calculated for any \( i \) from the OMI column
AAOD (\( \tau_{OMI_{i}} \)) using the ratio of vertically resolved BC AAOD to column AAOD in
the prior model,
\[
\tau_{OMI_{i}} = \tau_{OMI_{i}} \frac{\tau_{GC_{i}}^{a}}{\tau_{GC_{i}}} \quad (5),
\]
where superscript $\alpha$ indicates the prior model estimates. Since the ratio $\frac{\tau_{\text{GC,BC},i}}{\tau_{\text{GC},i}}$ is a constant throughout the inversion, the $i^{th}$ adjoint forcing is

$$\frac{\partial J}{\partial \tau_{\text{BC},i}} = \frac{\partial \tau_{\text{GC,BC},i}}{\partial \tau_{\text{BC},i}} * \left( \tau_{\text{GC,BC},i} - \frac{\tau_{\text{GC,BC},i}}{T_{\text{GC,i}}} \right) * S_{\text{OML},i}^2 \quad (6).$$

(b) In this method, the cost function is based on BC AAOD column differences:

$$J = \frac{1}{2} \sum_i^N \left( \tau_{\text{GC,BC},i} - \tau_{\text{OML,BC},i} \right)^2 * S_{\text{OML},i}^2 \quad (7).$$

The observed BC AAOD column is calculated from the OMI_GC AAOD column and the ratio of modeled column BC AAOD to total column AAOD from the prior simulation:

$$\tau_{\text{OML,BC},i} = \frac{\tau_{\text{OML},i}}{\tau_{\text{GC,i}}} \quad (8).$$

The $i^{th}$ adjoint forcing is thus

$$\frac{\partial J}{\partial \tau_{\text{BC},i}} = \frac{\partial \tau_{\text{GC,BC},i}}{\partial \tau_{\text{BC},i}} * \left( \tau_{\text{GC,BC},i} - \frac{\tau_{\text{GC,BC},i}}{T_{\text{GC,i}}} \right) * S_{\text{OML},i}^2 \quad (9).$$

(c) The observation term of the cost function can be written in terms of total column absorption as:

$$J = \frac{1}{2} \sum_i^N \left( \tau_{\text{GC},i} - \tau_{\text{OML},i} \right)^2 * S_{\text{OML},i}^2 \quad (10).$$

In this case, the adjoint forcing is

$$\frac{\partial J}{\partial \tau_{\text{BC},i}} = \frac{\partial \tau_{\text{GC,BC},i}}{\partial \tau_{\text{BC},i}} * \left( \tau_{\text{GC,BC},i} + \tau_{\text{GC,OC},i} + \tau_{\text{GC,Dust},i} - \tau_{\text{OML},i} \right) * S_{\text{OML},i}^2 \quad (11).$$

(d) The OMI OMAERUV retrievals algorithm also flags instances for which the retrieval algorithm relied upon the presence of carbonaceous aerosols. Using only these retrievals, the observation term of the cost function can be written in terms of the direct difference between simulated columns BC AAOD and BC flagged OMI AAOD observations:
where $T_{OMI,BC,Flag}$ is the OMI AAOD flagged for the presence of carbonaceous aerosols ($OMI_{GC,AAOD,BC}$, which is different than Eq. 5 or 8 which depend upon prior model ratios). In this case, the gradient of the cost function with respect to BC concentration at the layer $l$ will be

$$\frac{\partial J}{\partial BC_1} = \frac{\partial T_{GC,BC,l}}{\partial BC_1} \cdot (T_{GC,BC,l} - T_{OMI,BC,Flag,l}) \cdot S_{OMI,BC,l}^2$$

(13).

The implications of the different cost function formulations will be described in Section 4.1.

3 Impacts of BC anthropogenic emission uncertainties

In this section, we quantify the extent to which differences in anthropogenic emission inventories contribute to uncertainties in simulated surface BC and AAOD. Here, the SEAC4RS emission inventory is appended to the MEIC emission inventory outside of China for the Southeast Asian nested simulation (MEIC_SEAC4RS). Figure 4 shows the impact of different BC anthropogenic emission inventories on simulated surface BC concentrations and comparisons to in situ measurements over China [Zhang et al., 2008, Cao et al., 2009]. The monthly and daily ground-based measurements at sites representative of four different regions are shown: northern China (Gucheng, GUC), northeastern China (Longfengshan, LFS), southern China (Nanning, NAN), and midwestern China (XiAn, XIA). Generally, the modeled and observed BC concentrations are higher in winter than in summer. In addition to enhanced anthropogenic emissions during the winter [Fu et al., 2012], the Asian summer monsoon plays an important role in this seasonal cycle by reducing aerosol
concentrations in the summer over China [Zhang et al., 2010]. Though the model simulation is able to capture the seasonal variability, it underestimates surface BC concentration at the urban sites, such as GUC, NAN, and XIA, with all of these anthropogenic emission inventories, except at NAN, where the SEAC4RS inventory leads to values as high or higher than observed, but the seasonal variation has not yet been reproduced. With the INTEX-B and MEICS inventory, though the surface BC concentrations are underestimated at some background and rural sites [Fu et al., 2012; Wang et al., 2013], the simulated BC surface concentrations at the rural site of LFS are quite comparable to the observation, especially the seasonal variations. The INTEX-B and MEIC inventories improve the BC concentrations in winter with the inclusion of monthly variability over China compared to the inventories of Bond and SEAC4RS.

The spatial distributions of simulated surface BC concentrations using MEIC_SEAC4RS and INTEX-B inventories are compared to the in situ observation at 20 sites over Southeast Asia for April and October 2006 in Fig. 5. The east to west gradient in China and the north to south gradient in India are not well reproduced by the model, where the simulated BC concentrations are much lower over eastern China and the IGP for both April and October, especially for the urban areas since the model is unable to resolve individual urban hot spots [Fu et al., 2012].

Figure 6a shows the differences in monthly average AAOD between the model using the MEIC_SEAC4RS inventory and OMI (former minus latter) for April and October 2006. GEOS-Chem underestimates AAOD compared to OMI across broad areas of
Southeast Asia in April, especially eastern China and the IGP. In October, AAOD is underpredicted over northern China while it is over predicted over eastern China and most of South Asia. Corresponding OMI data counts towards the monthly average at each grid cell are shown in Fig. 6b. In general, more data are available over northern China and India. We note that the data counts are much lower in October compared to April over southern China and the Indo China Peninsular, where the observations are overestimated. Sparse OMI observations over these areas may result in apparent high or low biases. If we only take into account the OMI_GC AAOD_BC retrievals, the differences and corresponding OMI data counts for April and October are shown in Fig. 7. The spatial distributions are quite similar to those using all AAOD observations shown in Fig. 6, but with much larger negative differences over Asia in April and over northern China and IGP in October. The data counts are also smaller when only considering the OMI_GC AAOD_BC observations, especially over the dust source regions and downwind areas in April and broad areas over South Asia in October.

We also compared the observed to simulated AAOD using different emission inventories (figures not shown here). The simulated AAOD is comparable using INTEX-B and MEIC emission inventories over eastern China, while it is much lower than the OMI column retrieval using the inventories of Bond and SEAC4RS. With the SEAC4RS inventories, the simulated AAOD over the IGP shows enhancements compared to that using Bond and INTEX-B inventories.

4. Uncertainties of observation and penalty terms
4.1 Adjoint forcing

As described in Section 2.5, there are four methods to formulate the observation term of the cost function owning to different approaches of deriving an “observed” BC AAOD. We perform sensitivity experiments to quantify the impact of using these different formulations. For these tests, only the observation term is considered in the cost function (i.e., the penalty term is not included), and we use the same anthropogenic emission inventory (MEIC_SEAC4RS) as the prior emissions for each test. Figure 8 shows the results of the differences between optimized and prior anthropogenic BC emissions based on the four approaches.

Qualitatively, there are many noticeable differences between the optimization results using the different formulations of the observation operator. In April, enhanced anthropogenic BC emissions are shown over broad areas using all four methods. However, slight reductions appear over eastern China and southern India when using method (b), (c) and (d). In particular, method (c) results in lower posterior emissions over China. The results of methods (c) and (d) are quite consistent except the enhancements of posterior emissions over southern India occur using method (d).

Similarly, although the four optimized patterns are quite consistent in October, much larger areas of BC emissions reduction result from using method (c). The reductions of method (d) are similar to that of method (c) over eastern China, while quite different over India with significantly enhanced posterior emissions.

The differences in results are related to different assumptions implicit in the various forms of the cost function considered. Both method (a) and method (b) depend on the
relative ratio of BC to other absorbing aerosol (e.g. dust, OC) in the model. Further, method (a) introduces a stronger dependency on the GEOS-Chem prior vertical distribution, since the observation operator includes three dimensions with all vertical layers, compared to the column based method (b). Since there are more observations over IGP and northeastern China in April, this stronger constraint may enhance the negative forcing due to the model underestimation, which leads to increasing emissions. Since, through the adjustment of the OMI data to generate the OMI_GC product, we have already used the GEOS-Chem prior information on the aerosol vertical distribution, it seems preferable to adopt a column-based approach for the assimilation. Though both method (b) and method (c) are based on the column AAOD, the former assumes that the relative contributions of BC to total AAOD in the model is correct, while the latter assumes that absolute contributions of OC and dust are correct. The simulated total AAOD might not be equivalent to the observed AAOD after optimization in both method (a) and method (b) since the adjoint forcing only accounts for the BC AAOD. In addition, the results would highly depend on the model performance in simulating the ratio between BC and other absorbing aerosol. There are no significant biases for the GEOS-Chem simulated faction of coarse model dust mass [Wang et al., 2012, Philip et al., 2014], which suggests that the simulated dust AAOD fraction is likely unbiased. However the simulated mass of both BC and OC in GEOS-Chem are biased low [Heald et al., 2005; Fu et al., 2012]. We thus adopt method (c), since the strength of the adjoint forcing with respect to BC sources depends upon the BC absolute contribution in AAOD rather than the relative
contribution of method (b), which may have less model dependency in simulating the
distribution of other aerosols. The major differences between method (c) and method
(d) are the available observation data counts, as the data counts of the latter are much
greater than the former. In April, the pattern of optimized emissions using method (c)
and method (d) are quite consistent, suggesting that BC AAOD play a dominant role
in contributing to the total AAOD. We will adopt method (c) for the following
experiments and also discuss method (d) in section 5.4 for comparison.

4.2 Penalty Term

The inclusion of a penalty, or background term, in the cost function is a key factor for
inverse modeling. It is specified through the prior (background) error covariance
matrix, $S_\alpha$, and a regularization parameter $\gamma_r$. In the absence of rigorous statistical
information on the error covariance of the emissions, we assume the errors are
uncorrelated and use an L-curve selection criterion to identify an optimal value of $\gamma_r$
[Hansen, 1998; Henze et al., 2009]. The uncertainties of BC are assumed to be 100% of
the maximum BC emissions over the simulation domain. Thus, the optimal values
of $\gamma_r$ are selected to be 0.5 for April and 1.0 for October based on the
MEIC_SEAC4RS emission and the cost function in Eq. (10). The contribution of the
penalty term results in smaller adjustments to emissions, as the regularized results
prefer smoother solutions than those of the unconstrained inversion tests in Fig. 8.
Here we assume a single constant value for $S_\alpha$ along the diagonal and no off-diagonal
terms.
5. Analysis of Optimizations

We next proceed to constrain Southeast Asian BC sources using OMI_GC AAOD. The OMI_GC AAOD observations are compared to model estimates from GEOS-Chem nested simulation for April and October 2006 using the difference between simulated total AAOD and observed OMI_GC AAOD (i.e., Eq. (10)). Tens of thousands of OMI retrievals per month are available for the assimilation, but not all of the retrievals are usable. In the presence of cirrus clouds, retrievals errors are significant. The effect of optically thin cirrus is similar to that of subpixel cloud contamination. As plumes of dust or smoke aerosol drift away from their source regions, they become mixed with clouds. This problem is particularly evident over the oceans, which are frequently covered with thin cirrus and fair-weather cumulus clouds. Generally, the retrieved AAOD shows a reduced coverage especially over the oceans due to cloud contamination and the effects of sun glint [Torres et al., 2007]. Thus, quality and diagnostic flags are defined to classify and filter the retrievals. In October, only observations north of 5°N are included for data assimilation to minimize contributions of biomass burning from Indonesian fires.

5.1 Optimized emissions

Considering the performances of the four emission inventories discussed in Section 2.3, the following optimized results will mainly focus on using the MEIC_SEAC4RS and INTEX-B inventories. The prior and posterior (optimized) BC emissions from anthropogenic sources are shown in Fig. 9. Overall, the results show an enhancement in BC emissions over broad areas of Southeast Asia, with adjustments that are
seasonally and spatially heterogeneous. This is consistent with the top-down
constraints on BC emissions based on ground-base measurements by Fu et al., [2012],
which also show that the BC emissions are greatly enhanced across broad areas of
China, in particular northern and central China and the megacity clusters. In April,
either using MEIC_SEAC4RS or INTEX-B inventories, large increases of up to a
factor of 3-5 are shown after optimization. The largest enhancements occur sharply in
eastern China and the IGP in April by up to a factor of five (Fig. 9). Other large
increases are located in South Asia, northeastern and northwestern China. There is a
small decrease in anthropogenic BC in part of southwestern China. That is quite
different from the inversion results based on AOD by Xu et al. [2013], wherein the
optimized anthropogenic BC emissions are reduced by 9.1% over China, even though
the prior BC anthropogenic emissions that they used are from Bond et al., [2004,
2007], which much lower than what we used. The dust scheme had not yet been
updated and modified in Xu et al., [2013] following the revised particle size
distribution suggested in Zhang et al. [2013]. Thus it is possible that overestimated
dust and AOD projected a model bias onto adjustments of emissions of all type of
aerosols over dust regions and downwind areas, such as eastern China. Considering
the dust season in April, we also perform a sensitivity experiment to quantify the
uncertainty of dust impacts on the inversion results by doubling the dust emission in
April. The general pattern of the optimized anthropogenic BC emissions are
consistent with that of the standard inversion, with a maximum differences less than
20%. 
However, the adjustments of anthropogenic BC emissions before and after optimization in October are different than those in April (Fig. 10). The optimization of anthropogenic emissions yields a slight reduction (1~5%) over central India and part of southern China and an increase by 10~50% over eastern and northern China, as well as northwestern India.

Though the adjusted patterns of optimized BC emission are basically comparable by using MEIC_SEAC4RS and INTEX-B inventories, significant differences are located over India and eastern China (Fig. 11). We also note that the differences in the optimized results are almost the same as those of the prior emissions between MEIC_SEAC4RS and INTEX-B inventories. The ratio between their posterior differences and prior differences (see Fig. 11, right column) shows that the optimization increases their differences, relative to the prior, over broad areas over China and India up to a factor of three in April, with only slight decreases over south India. In October, optimization decreases the posterior differences between MEIC_SEAC4RS and INTEX-B emission inventories relative to the prior by 10-20% over southern and most of India. Areas where prior differences are increased/reduced are consistent with the areas where the emissions increase/decrease after optimization (see Fig. 10). This suggests that absolute errors in the prior emissions may be larger than the relative prior uncertainty percentages considered here.

In addition to reducing the bias of the emissions, it is important to consider how much the inversion has reduced uncertainty in the emissions. A new method based on the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm is used to estimate the
posterior uncertainty [Bousserez et al., 2014]. The posterior error reductions are up to
30% and 15% in April and October over the IGP and eastern China, where the
anthropogenic emission enhancements were the largest (Figure 9 and 10). The prior
errors do not change across broad areas, where the changes of optimized emissions
are relatively smaller.

While the most substantial adjustments are made to anthropogenic emissions, biomass
burning emission are also adjusted. The most significant increases are over South
Asia and eastern Europe in April, especially, the indo-China peninsula and eastern
Russia (figures not shown). The optimized biomass burning emissions in October
have the largest enhancements are over south Borneo and Sumatra. Similar to the
optimized anthropogenic emission, there is also not much change for the optimized
biomass burning emission throughout India and and indo-China peninsula in October.

To examine the impacts of different prior anthropogenic inventories on optimized
biomass burning emissions, we consider the following ratios:

\[
\frac{\Delta \text{MEIC,SEACARSGFED}_3 - \Delta \text{MEIC,SEACARSGFED}_2}{\text{GFED}_3 - \text{GFED}_2}
\]  (14).

Eq. 14 shows how changes in anthropogenic emissions during the optimization
compare when using two different biomass burning inventories, relative to the
difference in these biomass burning inventories themselves. Large values of this ratio
indicate regions where our top-down constrains on anthropogenic emissions are more
sensitive to errors in the prior biomass burning inventories, such as over eastern China
and the southern IGP (Fig. 12).

5.2 Optimized BC AAOD
The largest uncertainty reductions are obtained over eastern China and the IGP, so here we consider AAOD in these regions alone. Fig. 13 shows the observed and simulated BC AAOD over eastern China (105°-125°E, 20°-45°N) before and after optimization in green along with linear line slope equation and correlation $R^2$. Here the observed BC AAOD is derived from the OMI_GC AAOD and the prior ratio of simulated BC AAOD versus total AAOD. The prior BC AAOD is misrepresented and underestimated compared to observation over eastern China, especially in April. The low biases of the prior slopes are improved after optimization in April and October by 132% and 11%, respectively. Similar to the optimized BC concentrations, the improvements in October after optimization are less significant than in April. There are only slight changes in correlation coefficients, which may due to the large number of samples in both spatial and temporal dimensions across which variations are not in the same directions. In the IGP area, which we define as (70°-90°E, 23°-32°N), the low biases of prior BC AAOD are much larger than those in eastern China (Fig. 14). The values of most observed BC AAOD are lower than 0.3 and the slopes are 0.22 and 0.28 in April and October. After optimization, the slope increase by 155% and the correlation coefficients change from 0.2 to 0.25 in April. In October, there is a 32% increase in slope and the correlation coefficient doubles but still remains small (from 0.06 to 0.12).

Though slopes improve after optimization for both eastern China and India, they still show considerable lower biases. This results, in part, from constraints of the penalty term. Additionally, we note that many prior AAOD values are very small and close to
These are hard for the optimization routine to adjust significantly in the areas where the values of prior emission are very small or close to zero. Since the optimization scheme is based on the use of emissions scaling factors, large gradients with respect to BC concentrations will result in small gradients with respect to emissions scaling factors in locations with small emissions. To test how much this formulation restricts the inversion, a sensitivity experiment was performed assuming uniform prior emissions in all grid boxes. This facilitates adjustments to prior emissions throughout the domain, resulting in larger posterior AAOD after optimization. However, the resulting spatial distributions and gradients of anthropogenic emissions are not realistic (e.g., large emissions are not placed in known source areas). Alternatively, instead of adjusting emissions through application of scaling factors, $\sigma$, to the a priori emissions, the BC emissions themselves could be treated as the control variables in the cost function (Eq. 15). Another sensitivity experiment is performed for April 2006, inverting for the emissions themselves rather than the emissions scaling factors. Figure S2 in supplemental shows the total emissions (summed across sectors) after optimization using different inversion approaches. Fig. S2a is result based on the scaling factor as describe by Eq. 2 in Section 2.4 that the range of emissions are constructed using scaling factors as control variables to adjust the vector of model emissions. Fig. S2b shows the results when emissions are constrained directly as the control variables in the penalty term as:

$$ J = \frac{1}{2} \sum_{c \in \Omega} (Hc - c_{obs})^T S_{obs}^{-1}(Hc - c_{obs}) + \frac{1}{2} \gamma_r (E - E_a)^T S_a^{-1}(E - E_a) $$

(15).

This formulation allows the inversion to place significant emissions in areas where
the prior emissions are very small or close to zero. The optimized emissions over the larger prior source areas, such as northeastern China and the middle IGP, are smaller than when optimizing scaling factors. These sensitivity tests demonstrate the value of using the prior emissions inventories, either explicitly or implicitly through scaling factors, in terms of constraining the magnitude of known sources, and the downside in terms of the difficulty in introducing new sources through the inversion.

We also evaluate (Fig. 15) the prior and posterior simulated AAOD against the OMI and AERONET daily average AAOD at 4 sites where there are available measurements during the periods of April and October, 2006 (see the red sites in Fig. 2): Beijing (BJ) in China, Kanpur (KP) and Gandhi College (GH) in India, and Mukdahan (MD) in Thailand. The daily average GEOS-Chem model results and OMI_GC AAOD are sampled according to the AERONET observations at the locations of the 4 sites. At the Beijing site, the prior model AAOD estimates driven either by MEIC_SEAC4RS or INTEX-B inventories are underestimated by a factor of ~2, while the posterior AAOD are more comparable to the observations in April. In terms of temporal variability, the model is able to capture some features of peaks after optimization. At the two sites in India, only a few measurements are available in late April, but the magnitudes are close to OMI observation. The optimized results using the MEIC_SEAC4RS inventory shows great improvements compared to the prior AAOD. However, the optimized AAOD using the INTEX-B inventory still shows negative biases. The differences in optimized AAOD between using INTEX-B and MEIC_SEAC4RS come from their prior differences in AAOD. This again
demonstrates that the posterior optimization results are not independent of the prior emission inventories, consistent with the estimated reduction in posterior error shown in Fig 10. At the site of Gandhi_College (GH) and Mukdahan (MK) there are large differences between the OMI and AERONET AAODs; the magnitudes of the OMI_GC AAODs are much lower than those from AERONET, even close to zero on some days. Koch et al. [2009] compared the AERONET and OMI retrievals of AAOD at AERONET sites. The results showed that the two retrievals broadly agree with each other, but that the OMI_GC AAOD is much smaller over Asia. In our study, only a few OMI observed AAOD pixels are available in Thailand site (MK) (Fig. 6); these limited and sparse observations do not provide enough information to robustly constrain emissions in this region.

5.3 Optimized surface BC concentrations

As mentioned before, the prior surface BC concentrations are underestimated in most of the urban and rural sites over China. Figure 16 shows the spatial distribution of optimized surface BC concentrations compared to in situ measurements at 20 sites in Southeast Asia. The largest in situ BC concentrations observed over eastern China and the IGP, which are densely populated, industrialized areas, are now reproduced well by the optimized simulation. After optimization, the spatial gradients of the observed BC concentrations are captured by the model: high in the east and low in the west for China, and high in the north and low in the south for India. Using the MEIC_SEAC4RS inventory for the prior emissions, the optimized spatial distributions are better simulated than when than using the INTEX-B inventory. In particular, the
simulated BC concentrations are much closer to the observations over the IGP after optimization. The performance of simulated surface BC concentrations in the WRF-Chem (Weather Research and Forecasting model coupled with Chemistry) model with GOCART aerosol scheme using our optimized INTEX-B inventory has also been tested (see supplemental Fig. S3). A low bias using the prior INTEX-B inventory have been significantly reduced, and the simulated surface BC concentrations have increased by a factor of 1.5-2. The scatter plots in Fig. 17 show the correlations of BC concentrations from surface observations and GEOS-Chem before (blue) and after (red) optimization. Initial negative biases are shown in both April and October. The linear regression slope increases by more than a factor of four in April. However, the modeled BC concentrations at most of the sites only slightly change after the optimization in October, which result in a much smaller improvement in the regression slope (21%). The correlation coefficients increase by 0.04 to 0.08 after optimization, such small improvement may be owing to the sparse spatial distributions of the observational sites.

More specific site-by-site comparisons between model and observations are shown in Fig. 18. Although the optimized BC surface concentrations are enhanced in April, overestimation occurs in some eastern sites over China. In October, the low biases are corrected both in the urban sites and rural sites, especially the eastern rural sites in China. However, there is a persistent negative bias in most sites after optimization in October. Due to the very low prior emissions, the optimization has less impact on the western sites over China. The GEOS-Chem prior simulation underestimates surface
BC concentrations in all the urban sites and coastal sites over India in April (Fig. 16).

While the optimization enhances the BC sources and surface concentration, it still shows a negative bias in most of sites over India, especially the urban sites. The smaller improvement in coastal sites is not only due to the low prior emissions but also the large uncertainties of AAOD retrieval for low aerosol amounts over the ocean.

Given the stark contrast between the inversion results in April and October, we also conducted the optimization for two additional months in winter (January) and summer (July) season using MEIC_SEAC4RS as the prior inventory. In January, the anthropogenic emissions show enhancements over the IGP and parts of western and northern China and slight decreases over southern India and eastern and southern China (figures not shown here), which results in increasing the surface BC concentrations in XIA and LFS sites while decreasing concentrations in the sites of GUC and NAN (see Fig. 4). In July, there is no significantly change for the surface BC concentrations after optimization owing to very sparse observation in July over eastern China. From this seasonal comparison, it appears that the BC anthropogenic emissions are not always underestimated during the year. The largest underestimations across the whole region of Southeast Asia occur in April. The underestimated regions are mainly over IGP and northern China in both January and October. The slight overestimates are indicated over southern India and part of eastern China in January as well as northern China in July.

Discrepancies versus surface observations might also relate to model representational
error incurred by comparing ~50 km gridded estimates to in situ BC measurements, which likely have finer length-scales of variability [Wang et al., 2013; Cohen and Prinn, 2011; Cohen et al., 2011]. Considering the coarse resolution of the model when comprising with the ground-based measurements, we investigate the impacts of model resolution by considering approaches for downscaling the model simulations. One approach is to use high-resolution population datasets to redistribute primary aerosol concentrations [e.g., Krol et al., 2005; UNEP, 2011; Silva et al., 2013]. Based on a finer resolution population density dataset, a parameterization of the urban increment for non-reactive primary emitted anthropogenic BC and organic matter has been developed and tested for coarse resolution air quality model. This method does not alter concentrations at rural sites since it assumes that results at coarse resolution only represent the rural (background) sites. According to this method, we used a high-resolution (1/24° x 1/24°) population dataset of Gridded Population of the World, Version 3 (GPWv3, http://sedac.ciesin.columbia.edu/data/set/gpw-v3-population-density-future-estimates) to downscale and adjust the simulated BC concentration at urban sites (defined locations where population density exceeding 600/km²). The scatter plots (Fig. 17b) show that, on average, the application of population downscaling improves the performance of the modeled results compared to the non-adjusted BC concentrations in April for both the prior and posterior simulations, although low biases remain in each. It does not make any change in the slope in October after applying the population parameterization, and correlation is degraded. Downscaled estimates at only two sites (LIA and NAN) show enhancements, the rest
are not impacted.

To more directly investigate the impact of model resolution, it would be ideal to compare the results of the present simulations to higher resolution simulations with the same model [e.g., Punger and West, 2013]. While this is not currently an option for this model version, we can conduct GEOS-Chem simulations at a coarser resolution (2° latitude × 2.5° longitude) and make inferences about the role of resolution errors. Fig. 19 shows the resolution errors in estimated surface BC concentrations in the coarse resolution results (2° × 2.5°) with respect to fine resolution simulations (0.5° × 0.667°). The resolution error exceeds 20% across broad areas, and even up to 300% over the IGP and part of Southeastern Asia. The surface BC concentrations are much lower using coarse resolution over the major source regions, in particular the IGP, where the resolution error is more than 3. This is likely owing to coarse grid boxes not describing the sharp gradient between high concentrations in the valley and low concentrations in the mountain. The optimized surface BC concentrations from our 0.5° × 0.667° simulations are underestimated by a factor of 2-3 at the IGP sites compared to in situ measurements. Punger and West [2013] show that the percent difference between all-cause mortality estimates at 12 km resolution and at coarser resolutions of 36 km and 96 km for BC is ~9% and ~23% respectively. Assuming that model skill at estimating variations in concentrations at the scales of the in situ measurements is similar to that for estimating exposure based on highly resolved populations distribution, we can extrapolate from the results of Punger and West [2013] that the resolution errors in the
0.5°x0.667° simulation, relative to the scale of the measurements, is a bit less than the
resolution error in the 2°x2.5° simulation relative to the 0.5°x0.667° simulation
Thus, the former may be as large as a factor of ~2.5 in individual grid cells.

5.4. Comparisons using OMI_GC AAOD_BC

A subset of the OMI retrievals (OMI_GC AAOD_BC) represents the presence of
carbonaceous aerosols. Using only these retrievals for the inversion, the differences
between prior and posterior (later minus former) BC anthropogenic emissions using
MEIC_SEAC4RS inventory are shown in Fig. 20. Compared to Fig. 9 and Fig. 10,
there are similar signs of emissions adjustments over most of Southeast Asia except in
October over India where reductions are not shown in the posterior emissions due to
fewer available observations in the OMI_AADO_BC data subset. Moreover, the
magnitudes of enhanced emissions in April are much larger if we use only the
OMI_GC AAOD_BC retrievals. This also results in larger posterior surface BC
concentrations (figures not shown) in some area and AAOD that improve the
underestimates in a few sites when compared to the ground-base measurements and
AERONET observation. However, the differences are not obvious in October and the
improvements in April are neither significant nor widespread. Considering there are
less observations available using OMI_GC AAOD_BC, especially in October and
other summer month (e.g. July), and that it does not change the major conclusions
compared to using OMI_GC AAOD, using OMI_GC AAOD is recommended.

6. Summary and Discussions
In this study, we used space-based observations of absorbing aerosol optical depth (AAOD) from the OMI instrument to constrain BC monthly average emissions for April and October 2006, with the GEOS-Chem model and its adjoint. First, we evaluated the model simulated BC concentrations using four different anthropogenic emission inventories. The differences in these inventories exceeded 100% across broad areas of Southeast Asia. For each of the four emission inventories, the simulated surface BC concentrations had low biases compared to the available surface observations in most urban sites in Southeast Asia.

The adjoint model was used to perform 4D-Var inverse modeling to constrain BC emissions. After optimization, both anthropogenic and biomass burning emissions were adjusted. Either using the MEIC\_SEAC4\_RS or INTEX-B inventory, the optimized anthropogenic emissions for BC were significantly enhanced over broad areas of Southeast Asia in April compared to the prior emission, with the largest enhancements in eastern China and India IGP of up to a factor of five. From analysis of inversions using different prior biomass burning inventories it was shown that optimized anthropogenic emissions was most sensitive to the prior biomass burning over eastern China and southern IGP. The adjustments in October were smaller than those in April. Inverse modeling in additional months indicated that BC anthropogenic emissions were not always underestimated throughout the year. The largest underestimates occurred in April throughout Southeast Asia. Only slight overestimates were indicated over southern India and eastern China for both January in July. Inversion results were in general similar using either all OMI observed
AAOD or just the OMI_GC AAOD_BC. In October, the posterior anthropogenic emissions yielded a slight reduction (1−5%) over central India and part of southern China while they increased by 10−50% over eastern and northern China, as well as northwestern India. The uncertainty of the posterior emissions over the IGP and eastern China were estimated to have reduced up to 30% and 15% in April and October. Though April is the Asian dust season, the impact of doubling dust emissions on the posterior anthropogenic emissions is less than 20%.

After optimization, the low model biases for BC AAOD improved by 132% and 11% over Southeast Asia in April and October, respectively. In eastern China, these improvements were more significant (143% and 30% in April and October). The remaining residual error in the simulated AAOD, which was significant in October, particularly in India, may be a consequence of the inverse modeling framework, which had difficulty introducing emissions in locations where the prior emissions were close to zero. This downside may be overcome by performing inversions directly for the emissions, rather than emissions scaling factors.

Results of the inversion were also compared to remote and in situ measurements that were not assimilated. The posterior AAOD were quite comparable to AERONET AAOD observations in April in China; however, large discrepancies remained at the sites over India and Thailand after data assimilation. These residual errors compared to AERONET may be associated with the limited and sparse observations of OMI observed AAOD in these regions, which themselves were not very consistent with the AERONET AAOD. Jethva et al., [2014] also pointed out that much of the observed
inconsistency of SSA between OMI and AERONET is found to occur at moderate to lower aerosol loading (AOD 440nm<0.7) for which both inversion techniques might have issues related to signal-to-noise ratio and algorithmic assumptions. Low biases of surface BC concentrations were improved or corrected at urban sites and eastern rural sites over China in April, with the linear regression slope between model and observed values increasing by more than a factor of four. However, the adjustments were not strong enough in most sites over India in April and October and over China in October. Moreover, the optimization had less impact on the western sites over China and costal sites over India due to the very low prior emissions and the large uncertainties in AAOD retrieval for low aerosol amounts over ocean. Model resolution error was also an important factor contributing to discrepancies of BC concentrations compared to in situ measurements. Comparison to coarser model simulations and the results of Punger and West [2013] indicates that the resolution errors may be up to a factor of 2.5 in grid cells in regions such as the IGP and part of southeastern Asia. Nevertheless, the results found here are not exclusively germane to GEOS-Chem, as we find that implementing the optimized INTEX-B inventory in WRF-Chem improved simulated surface BC concentrations by a factor of 1.5-2 relative to simulations with the prior INTEX-B inventory. Overall, this work was the first attempt to formally use the absorbing aerosol products from satellite observation for a BC emissions inversion. Both the simulated AAOD and surface BC concentration showed significant improvements spatially and temporally after data assimilation, especially in April. However, there were still
several sources of uncertainty and limitations of this work worth considering. Aspects such as model error and assumptions made regarding the observations and uncertainties in the observations and prior emissions inventories contributed greatly to uncertainties in the optimization results.

Our estimate that the errors in the prior emissions were only 100% restricted the magnitude of the emissions adjustments allowed by the inversion. One might conclude that such restrictions were too strict; however, uncertainties in emissions were also not likely the only source of the discrepancy between observed and predicted BC concentrations and AAOD. Textor et al. [2007] noted that inter-model differences were only partially explained by differences in emission inventories; removal processes also play an important role in affecting the lifetime and concentrations of BC in the free troposphere. Although the 1 day aging from hydrophobic BC to hydrophilic BC in GEOS-Chem is typical for this type of model [Koch et al., 2009], aerosol internal mixing that includes effects of various physical, chemical, and meteorological processing can also significantly impact BC concentrations and aerosol absorptions [Stier et al., 2006; Cohen and Prinn 2011; Cohen et al., 2011; Buchard et al., 2014], in some cases even more so than uncertainties in emissions [Shen et al., 2014]. The scheme used in our study for aerosol scavenging was based on Liu et al., [2001], which did not distinguish between rain and snow. The recent updates by Wang et al. [2011] included corrections to below-cloud and in-cloud scavenging that improved the overestimation of integrated scavenging [Dana and Hales, 1976]. Corresponding updates to the wet scavenging in
the GEOS-Chem adjoint might also be helpful for improving the optimized results.

The optimizations were sensitive to how model information was used to calculate BC component of the measured AAOD, which alone provided only a constraint on the column concentrations of all absorbing aerosol (i.e., including dust and OC). We have adjusted the OMI observed AAOD by applying the GEOS-Chem simulated aerosol layer height to reduce the differences in the vertical profiles between the model and observation, referred to as OMI_GC AAOD. However, there could be inconsistent treatment of microphysical and optical properties used in the AAOD calculation between the model and OMI retrievals. The results of the optimization may be biased by error in the model’s vertical distribution of BC, which has been adjusted in other studies [van Donkelaar et al., 2013]. To evaluate the magnitude of this potential source of error, we also repeated the inversions using the OMI retrieval “Final AAOD” products (OMI_Final) based on the CALIOP and GOCART aerosol layer height. The difference in the optimized anthropogenic BC emissions are less than 30% in April and 10% in October compared to inversions using OMI_GC AAOD which is based on GEOS-Chem aerosol layer height.

It is important to realize that BC from most emission sources contained not only elemental and organic fractions [Chow et al., 2009], but also non-soot OC, i.e., brown carbon, that has a significant absorbing component at short wavelengths comparable to elemental carbon absorption [Jacobson, 1999; Kirchstetter et al., 2004; Andreae and Gelencser, 2006; Hoffer et al., 2006; Magi et al., 2009]. However, absorbing aerosols in GEOS-Chem only include BC, OC and dust, while the brown carbon has
not yet been taken into account. While the attribution of ambient aerosol absorption to BC may be a reasonable approximation in areas dominated by fresh soot emissions, it may lead to misleading estimates of the AAOD when other light absorbing particles were present since the brown carbon contributed 28% on average of the total absorption at the wavelength of 440 nm [Bahadur et al., 2012]. It undoubtedly resulted in overestimation of BC emissions after optimization in the areas where brown carbon and other absorbing aerosols were considered in the observed AAOD. Lastly, it is well known that the quality of the observation data plays the most important role in data assimilation. Although the OMI observed AAOD retrieval provided much better spatial and temporal coverage than the remote sensing measurements, such as AERONET, we noted that there were large discrepancies between OMI_GC AAOD and AERONET observation in some areas, especially in October (See Fig. 15). Normally, the OMAERUV retrievals were more reliable over land than over water since the ocean surface reflectance show distinct angular and spectral variations. The major factor affecting the quality of the OMI aerosol product was sub-pixel cloud contamination due to the relatively large footprint of the OMI observations [Torres et al., 1998]. Satheesh et al. [2009] demonstrated the potential of multisatellite analysis of A-train data to improve the accuracy of retrieved aerosol products and suggested that a combined OMI-MODIS-CALIPSO retrieval had potential to further improve assessments of aerosol absorption, which would possible enhance the observation quality in data assimilation. Recently, other improvements included the development of CALIOP-based aerosol layer height climatology and the
use of AIRS carbon monoxide real time observations to distinguish smoke from dust type aerosols, which improved the retrieval performance by 5-20% [Torres et al., 2013]. Using the updated OMAERUV when it becomes available will likely improve the optimization results in future work.

Acknowledgement.

This work was supported from Environmental Protection Agency-STAR grant RD-83503701-0. Although the research described in the article has been funded wholly or in part by the U.S. EPA's STAR program through grant (RD-83503701-0), it has not been subjected to any EPA review and therefore does not necessarily reflect the views of the Agency, and no official endorsement should be inferred. We thank the OMI team (http://disc.sci.gsfc.nasa.gov/Aura/data-holdings/OMI/omaeruv_v003.shtml) and AERONET team (http://aeronet.gsfc.nasa.gov/cgi-bin/webtool_opera_v2_inv) for providing the data and establishing and maintaining the sites used in this study.

Table and Figures

Table 1 Comparison of BC anthropogenic emissions over eastern China (105°-125°E, 20°-45°N) and IGP (70°-90°E, 23°-32°N), unit: Tg.

<table>
<thead>
<tr>
<th>Domain</th>
<th>Prior emissions (MEIC_SEAC4RS)</th>
<th>Posterior emissions (with penalty term)</th>
<th>Posterior emissions (without penalty term)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>April</td>
<td>October</td>
<td>April</td>
</tr>
<tr>
<td>Eastern China</td>
<td>0.11</td>
<td>0.11</td>
<td>0.30</td>
</tr>
<tr>
<td>IGP</td>
<td>0.04</td>
<td>0.04</td>
<td>0.14</td>
</tr>
</tbody>
</table>

Figure captions.
Figure 1. Absolute and relative differences in AAOD between OMI_Final and OMI_GC AAOD for April and October, 2006.

Figure 2. Twenty sites of ground measurements (black dots) and four sites of AERONET observation (red cross dots). Also shown are terrain heights (color shaded contours, unit: m).

Figure 3. Annual anthropogenic emission of BC regridded into GEOS-Chem resolution of 0.5° × 0.667° from the inventories of (a) Bond, (b) INTEX-B, (c) SEAC4RS, and (d) MEIC.

Figure 4. Comparison of the observed and simulated surface BC concentrations using four emission inventories at the site of GUC, LFS, NAN, XIA. The orange dots are the monthly mean posterior surface BC concentrations at these sites using MEIC inventory over China.

Figure 5. Spatial distributions of prior surface BC concentrations using INTEX-B and MEIC_SEAC4RS inventories overlaid with BC in situ measurements of 20 sites.

Figure 6. (a) Differences of monthly average AAOD between model using MEIC_SEAC4RS inventory and the OMI observation (former minus latter) and (b) corresponding OMI monthly data in each grid cell for April and October, 2006.

Figure 7. The same as Figure 6, but for OMI_AAOD_BC.

Figure 8. Differences between optimized and prior anthropogenic BC emissions based on four methods of adjoint forcing (a) vertically resolved BC AAOD base on model, (b) column BC AAOD based on model, (c) total OMI_GC AAOD and (d) column OMI_GC AAOD_BC for April and October, 2006.

Figure 9. Anthropogenic BC emissions for April, 2006. The first column shows the prior inventory, the second the optimized inventory, the third the differences between the prior and optimization, and the last column the relative changes of posterior error, based on the inventories of (a) INTEX-B and (b) MEIC_SEAC4RS.

Figure 10. The same as Figure 9, but for October 2006.

Figure 11. Differences of anthropogenic BC emissions between using the inventories of MEIC_SEAC4RS and INTEX-B for April and October 2006. The left column shows the prior inventory, the center the optimized inventory, and right column the between their posterior differences and prior differences.

Figure 12. The sensitivities of optimized anthropogenic emission based on GFED2
and GFED3 relative to the differences between GFED2 and GFED3.

**Figure 13.** Comparison of BC AAOD over eastern China (105°-125°E, 20°-45°N) from OMI measurements and GEOS-Chem before and after the assimilation for April and October, 2006.

**Figure 14.** Comparison of BC AAOD over IGP (70°-90°E, 23°-32°N) from OMI measurements and GEOS-Chem before and after the assimilation for April and October, 2006.

**Figure 15.** Comparison of total daily AAOD from OMI, AERONET and GEOS-Chem before and after the assimilation at the four AERONET sites for April and October, 2006.

**Figure 16.** Spatial distributions of optimized surface BC concentrations using INTEX-B and MEIC_SEAC4RS inventories overlaid with BC in situ measurements of 20 sites.

**Figure 17.** Comparison of monthly surface BC concentration for April and October, 2006, from in situ measurements and GEOS-Chem before and after the assimilation (a) without and (b) with population density downscaling.

**Figure 18.** Comparison of monthly surface BC concentration from in situ measurements and GEOS-Chem over (a) China and (b) India before and after the assimilation using the inventories of MEIC_SEAC4RS and INTEX-B for April and October, 2006.

**Figure 19.** The resolution errors of surface BC between the simulations of coarse resolution (2°x2.5°) and fine resolution (0.5°x0.667°).

**Figure 20.** The differences between the prior and posterior anthropogenic BC emissions for April and October, 2006, using OMI_GC AAOD_BC as the observation.
References


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1999.


Henze, D. K., Hakami, A., and Seinfeld, H. J: Development of the adjoint of GEOS-
air quality influences of inorganic PM2.5 precursor emissions using the adjoint
of GEOS-Chem, Atmos. Chem. Phys., 9, 5877–5903, doi: 10.5194/acp-9-5877-

Hoffer, A., Gelencser, A., Guyon, Kiss, P., G., Schmid, O., Frank, P. G., Artaxo, P.,
and Andreae, O. M.: Optical properties of humic-like substances (HULIS) in

Holben, B. N., Eck, F. T., Slutsker, I., Tanré, D., Buis, P. J., Setzer, A., Vermote, E.,
Reagan, A. J., Kaufman, J. Y., Nakajima, T., Lavenu, F., Jankowiak, I., Smirnov,
A.: AERONET--A federated instrument network and data archive for aerosol

estimation of 2004 elemental carbon emissions inventory in the United States to
the choice of observational networks, Geophys. Res. Lett., 36, L15806,

emissions inventory in the United States by inverse modeling using Community
Multiscale Air Quality model with decoupled direct method (CMAQ-DDM), J.

Huneeus, N., Boucher, O., and Chevallier, F.: Atmospheric inversion of SO2 and
primary aerosol emissions for the year 2010, Atmos. Chem. Phys., 13, 6555-

Jacobson, M. Z.: A physically-based treatment of elemental carbon optics:
Implications for global direct forcing of aerosols, Geophys. Res. Lett., 27(2),

Jacobson, M. Z.: Isolating nitrated and aromatic aerosols and nitrated aromatic gases
as sources of ultraviolet light absorption, J. Geophys. Res.-Atmos., 104(D3),
3527-3542, 1999

Janssen N.A., Hoek G., Simic-Lawson M., Fischer P., van Bree L., ten Brink H.,
Keuken, M; Atkinson, R. W., Anderson, H. R., Brunekreef, B., Cassee, F. R.:
Black Carbon as an Additional Indicator of the Adverse Health Effects of
Airborne Particles Compared with PM10 and PM2.5. Environ Health Perspect

Janssen NAH, Lanki, T., Hoek, G., Vallius, M., de Hartog, J. J., Van Grieken, R.,
Pekkanen, J., Brunekreef, B.: Associations between ambient, personal and
indoor exposure to fine particulate matter constituents in Dutch and Finnish

Jethva, H. and Torres, O.: Satellite-based evidence of wavelengthdependent aerosol
absorption in biomass burning smoke inferred from Ozone Monitoring
Instrument, Atmos. Chem. Phys., 11, 10541–10551, doi:10.5194/acp-11-10541-

Jethva, H., Torres, O., and Ahn, C.: Global assessment of OMI aerosol single-
scattering albedo using ground-based AERONET inversion, J. Geophys. Res.


Fig. 2
Fig. 3
Fig. 4
Fig. 5
Fig. 6
Figure 7

(a) GEOS-Chem minus OMI

(b) Data counts

April

October
Fig. 8
Fig. 9

(a) INTEX-B
(b) MEIC_SEAC'RS

Prior  Posterior  Posterior - Prior  Relative changes of posterior error

0.1 1 2.5 5 10 20 30 50 10^4 kg

-30 -20 -10 -5 -1 0.01 0.01 5 10 20 50 10^4 kg

-30 -20 -15 -10 -5 1 5 10 15 20 30 %
Fig. 10

Prior

Posterior

Posterior - Prior

Relative changes of posterior error

(a) INTEX-B

(b) MEIC_SEAC'RS

Prior

Posterior

Posterior - Prior

Relative changes of posterior error

Fig. 10
Fig. 11
Fig. 12
Fig. 15
Fig. 16
Fig. 1

Obervation
Prior-MEIC_SEAC
Posterior-MEIC_SEAC
Prior-INTEX-B
Posterior-INTEX-B
Fig. 19
Fig. 20