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A multi-model evaluation of aerosols over South Asia: Common problems and possible causes

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19 Abstract

- Atmospheric pollution over South Asia attracts special attention due to its effects on regional climate, water cycle and human health. These effects are potentially growing owing to rising
- trends of anthropogenic aerosol emissions. In this study, the spatio-temporal aerosol
- 23 distributions over South Asia from seven global aerosol models are evaluated against aerosol
- 24 retrievals from NASA satellite sensors and ground-based measurements for the period of
- 25 2000-2007. Overall, substantial underestimations of aerosol loading over South Asia are found
 26 systematically in most model simulations. Averaged over the entire South Asia, the annual
- 27 mean aerosol optical depth (AOD) is underestimated by a range 15% to 44% across models
- compared to MISR, which is the lowest bound among various satellite AOD retrievals (from
- 29 MISR, SeaWiFS, MODIS Aqua and Terra). In particular during the post-monsoon and
- 30 wintertime periods (i.e. October-January), when agricultural waste burning and anthropogenic
- emissions dominate, models fail to capture AOD and aerosol absorption optical depth (AAOD)
 over the Indo-Gangetic Plain (IGP) compared to ground-based AERONET supphotometer
- 33 measurements. The underestimations of aerosol loading in models generally occur in the lower
- 34 troposphere (below 2km) based on the comparisons of aerosol extinction profiles calculated by
- 35 the models with those from CALIOP data. Furthermore, surface concentrations of all aerosol
- 36 components (sulfate, nitrate, organic aerosol and black carbon) from the models are found
- 37 much lower than in-situ measurements in winter. Several possible causes for these common
- 38 problems of underestimating aerosols in models during the post-monsoon and wintertime
- 39 periods are identified: the aerosol hygroscopic growth and formation of secondary inorganic
- 40 aerosol are suppressed in the models because relative humidity is biased far too low in
- 41 boundary layer, the nitrate aerosol is either missing or inadequately accounted for, and
- 42 emissions from agricultural waste burning and biofuel usage are too low in the emission

- 43 inventories. These common problems and possible causes found in multiple models point out
- 44 directions for future model improvements in this important region.
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46 **1. Introduction**

47 South Asia, particularly the Indo-Gangetic Plain (IGP) bounded by the towering Himalaya that is conducive to trapping both anthropogenic and dust aerosols (Fig. 1), is one of the global 48 49 hotspots with persistent high aerosol optical depth (AOD) routinely observed by satellite remote sensors (e.g. Moderate Resolution Imaging Spectroradiometer or MODIS, Multi-angle 50 51 Imaging SpectroRadiometer or MISR and Sea-Viewing Wide Field-of-View Sensor or 52 SeaWiFS), as well as from ground-based measurements (e.g. Aerosol Robotic Network or 53 AERONET). The potential influence of aerosols on the climate and water cycle in this region (e.g. Indian summer monsoon) via surface dimming and atmospheric warming has been widely 54 discussed in the literature (e.g. Ramanathan et al., 2005; Lau et al., 2006). The atmospheric 55 heating due to absorbing aerosols (mainly from black carbon i.e. BC) is estimated to be large 56 especially in the wintertime, about 50-70 W m⁻² (Ganguly et al., 2006). Recent studies have 57 shown that the depositions of absorbing aerosols such as BC and dust over Himalava are 58 linked to snow albedo reduction and accelerated snow/ice melt in Himalaya during the pre-59 60 monsoon season (Lau et al., 2010; Qian et al., 2011; Yasunari et al., 2010; Gautam et al., 61 2013).

62 Besides these climate impacts, fine aerosol particles (PM_{2.5}) are known to affect public health, especially over IGP where large portions of the Indian population live. At Delhi, for 63 example, PM_{2.5} concentration in 2007 was 97±56 µg/m³ (Tiwari et al., 2009), nine times the air 64 65 quality guidelines recommended by the World Health Organization in 2005. Increases in anthropogenic aerosol emissions and loading in South Asia in recent decades have been well 66 documented (Ohara et al., 2007; Hsu et al., 2012; Kaskaoutis et al., 2012; Babu et al., 2013), 67 in contrast with the decreasing emission trends over Europe and North America (Granier et al., 68 69 2011; Diehl et al., 2012). Therefore, it is critical to accurately represent aerosol sources, distributions and properties in models over this heavily polluted region in order to project the 70 71 future climate and air quality changes in South Asia with confidence.

72 Previous studies, however, reported that global models generally underestimated 73 aerosol loading over South Asia, especially over the IGP in winter (Reddy et al., 2004; Chin et 74 al., 2009; Ganguly et al., 2009; Henriksson et al., 2011; Goto et al., 2011; Cherian et al., 2013; 75 Sanap et al., 2014). Among them, Ganguly et al. (2009) reported that the GFDL-AM2 model 76 largely underestimated the AOD over the IGP during winter by a factor of 6. Recently, AOD simulated by the regional climate model (RegCM4) showed higher correlation with AERONET 77 78 AOD at stations over dust-dominated areas in south Asia than over the regions dominated by 79 anthropogenic aerosols, i.e. 0.71 vs. 0.47 (Nair et al., 2012). Eleven out of twelve models participating in the Aerosol Comparisons between Observations and Models (AeroCom) phase 80 81 I exercise were also found to underestimate the aerosol extinction over South Asia, especially 82 under 2 km, in comparison with the space-borne lidar measurements from the Cloud-Aerosol 83 Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) satellite (Koffi et al., 2012).

84 The ability to capture surface BC concentrations over South Asia for models has also been

- found to be limited, with the low biases that tend to be larger in winter (Ganguly et al., 2009;
- Menon et al., 2010; Nair et al., 2012; Moorthy et al., 2013). A very recent study evaluating the
- 87 latest generations of quasi-operational aerosol models participating in International
- 88 Cooperative for Aerosol Prediction (ICAP) has shown that the models have very low skill
- 89 scores in reproducing AERONET measured AOD at Kanpur, an urban city in northern India
- 90 (Sessions et al., 2015). These studies underscore great challenges for current global aerosol
 91 models to adequately represent aerosols in South Asia.

Extending from previous studies and utilizing the recent model outputs from the AeroCom Phase II multi-model experiments, the present work systematically evaluates aerosol simulations in South Asia by seven global aerosol models with observations from satellites and ground-based measurements, and strives to characterize the model deficiency in reproducing observations. The outcomes of this study will help us understand the discrepancies between models and observations, thus providing directions for future model improvements in this important region.

99 The description of models is given in Section 2, followed by the introduction of 100 observational data from satellites and ground-based measurements in Section 3. The model 101 results are compared with observations in Section 4, including the spatial and temporal 102 distribution of AOD and aerosol absorption optical depth (AAOD), vertical profile of aerosol 103 extinction coefficient, and the surface BC concentration. The diversity among models is 104 discussed in Section 5, and possible causes for the model underestimations of aerosol 105 amounts are investigated in Section 6. Major findings are summarized in Section 7.

106

107 **2. Model description**

108 2.1 Models

109 Aerosol simulations for the period of 2000-2007 from seven models, including six models that participated in AeroCom Phase II hindcast experiment (i.e. AeroCom II HCA) and one 110 111 additional model, GEOS5, are analyzed in this paper (see Table 1 for details). Note that the 112 model outputs related to aerosol optical properties, such as AOD, AAOD and extinction 113 coefficient, are at the wavelength of 550 nm. Given that MODIS and MISR are available only 114 after 2000, we chose the years 2000-2007 in this study although longer time period of 115 simulations (starting from 1980) are available from the AeroCom models (note that ECHAM5-116 HAMMOZ ended in 2005 and HadGEM2 in 2006). Aerosol modules in GEOS5 are based on GOCART with some modifications (Colarco et al., 2010). More detailed descriptions about 117 these models can be found in previous studies (see references listed in Table 1 and Myhre et 118 al., 2013). All models include sulfate (SO₄²⁻), BC, organic aerosol (OA), dust (DU) and sea salt 119 120 (SS). Nitrate (NO_{3⁻}) is included only in three models (GISS-modelE, GISS-MATRIX and 121 HadGEM2). The secondary organic aerosol (SOA) chemistry is resolved in two models, GISS-122 modelE and HadGEM2, whereas simple parameterizations of SOA are used in the remaining 123 models. There are some differences among the seven models on aerosol optical properties 124 (see refractive indices listed in Table 1). In comparison with satellite retrievals and AERONET 125 observations that are available only under clear-sky conditions, it is desirable to use the

126 modeled AOD for clear-sky as well; however only two GISS models provide such output (other

models just provide all-sky results). In general, clear-sky AOD is lower than all-sky AOD, for

example, by 20% globally based on the GEOS-Chem model (Yu et al., 2012). All seven

- models use the assimilated wind fields although from different datasets. The horizontal
- resolutions vary from 2.8° by 2.8° (ECHAM5-HAMMOZ) to 1.1° by 1.1°(SPRINTARS) and the
- vertical levels range from 30 (GOCART-v4) to 72 (GEOS5) intervals. More information is givenin Table 1.
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134 **2.2 Emissions**

For anthropogenic emissions, which are mainly from consumption of fossil fuel and biofuel, the models use either A2-ACCMIP or A2-MAP emission dataset that are provided for the AeroCom Phase II model experiments (Diehl et al., 2012). Both A2-ACCMIP and A2-MAP were

- 137 constructed by combining multiple inventories but in different ways. The annual anthropogenic
- emissions from A2-MAP are yearly emission dataset with inter-annual variability, while those
- from A2-ACCMIP are without actual inter-annual variability, simply generated by linear
- 140 interpolation between decadal endpoints except for biomass burning (Granier et al., 2011;
- 141 Interpolation between decadal endpoints except for biomass burning (Granier et al., 2011; 142 Diehl et al., 2012). Over South Asia, the spatial distribution and total emission amount are
- 142 Diehl et al., 2012). Over South Asia, the spatial distribution and total emission amount are 143 somewhat different between the two emission datasets, with higher emission amount in A2-
- ACCMIP. Detailed information on both emission datasets can be found in Diehl et al. (2012).
- 145 Figure 2 shows the averaged annual mean (2000-2007) anthropogenic BC, organic 146 aerosol (OC), SO₂, NH₃ and NO_x emissions in South Asia from A2-ACCMIP anthropogenic 147 emission dataset (A2-MAP is not shown and it does not provide NH₃ and NO_x emissions). In 148 this study, we define the South Asia domain as 60°E–95°E longitude and 5°N –36°N latitude. 149 Note that the seasonal cycle of anthropogenic emission is not resolved in either emission 150 datasets, which could be problematic especially for biofuel emission in this region (discussed 151 in Section 6.3). The anthropogenic emissions display high spatial heterogeneities over South 152 Asia, coinciding with the population density distribution as reported by previous studies (e.g. 153 Girolamo et al., 2004). Densely populated regions are usually associated with heavy 154 anthropogenic emissions in South Asia, especially over IGP. The annual mean anthropogenic aerosols emission in South Asia for the period of 2000-2007 from A2-ACCMIP (A2-MAP) are 155 7.46 (5.33) Tg yr⁻¹ of SO₂, 5.94 Tg yr⁻¹ of NH₃, 4.50 Tg yr⁻¹ of NO_x, 2.18 (1.71) Tg C yr⁻¹ of OC, 156 157 and 0.69 (0.65) Tg C yr⁻¹ of BC. The ratio of OC/BC anthropogenic emissions (fossil fuel and 158 biofuel) is 3.2 (2.6) over South Asia.
- 159 Open biomass burning including the agricultural residue burned in the field and forest fires contributes to 25% of total BC (and OC) emissions over India based on the estimation by 160 161 Venkataraman et al. (2006) with the difference between the total crop waste and that used as 162 fuel and animal fodder. Figure 3 shows the seasonal BC biomass burning emission based on monthly Global Fire Emissions Database Version 2 (GFED2), which is used by both A2-163 164 ACCMIP and A2-MAP emission datasets. The open biomass burning displays strong spatial 165 and seasonal variations. Pre-monsoon period is the most active open biomass burning season with an emission amount of 0.22 Tg C yr⁻¹ of BC over South Asia, concentrated over 166

northeastern India associated with the Jhum cultivation to clear the forest and create fields
(Vadrevu et al., 2013). Seasonal practices of biomass burning of agricultural crop residues
associated with rice-wheat crop rotation over the western IGP, such as Punjab, Haryana and
western Uttar Pradesh, could explain the high aerosol loading during the post-monsoon of
October-November (Badarinath et al., 2009a; Sharma et al., 2010; Vadrevu et al., 2011;
Vadrevu et al., 2013) with a total emission amount of 0.001 Tg C yr⁻¹ BC over South Asia in
GFED2. The ratio of OC/BC open biomass burning emission is 8.0 averaged over South Asia.

174 The major natural aerosol over South Asia is the wind-blown mineral dust from the arid 175 and semi-arid regions of southwest Asia, such as Iran, Afghanistan, Pakistan, Arabian 176 Peninsula, and Thar Desert in the northwestern India. The dust emissions are calculated by 177 each model and show a large diversity varying from 10.6± 3.3 (ECH) to 185.8± 33.6 (SPR) Tg 178 yr⁻¹ over South Asia (averaged for 2000-2007). This model diversity is attributed to differences 179 in the model size bins of dust aerosols, parameterization of source strength, and wind fields 180 and soil properties over source regions (see more detailed discussions in Section 5). Sea salt 181 emission is negligible for the study area.

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183 **3. Observational datasets**

184 **3.1 Satellite data**

185 In this study, five satellite products are used to characterize aerosol distribution and evaluate the model simulations. MODIS Terra and Aqua level-3 monthly mean AOD products 186 187 at 550nm wavelength (Collection 5.1) are used by averaging the daily aerosol products at 188 1°×1° grid. The MODIS AOD is a composite of the Dark Target (Levy et al., 2007 and 2010) 189 and Deep Blue retrieval products (Hsu et al., 2006), as the latter is able to retrieve AOD over 190 bright surfaces such as the Thar Desert in South Asia. SeaWiFS level-3 monthly AOD 191 products at 550nm (V003) are obtained by averaging the daily aerosol products at 1°×1° grid. 192 SeaWiFS retrieval adopts the Deep Blue algorithm over land (Hsu et al., 2006, 2012) and 193 Ocean Aerosol Retrieval (SOAR) algorithm over ocean (Saver et al., 2012 and 2013). MISR 194 level-3 monthly AOD products at 555nm (V004) are used by averaging the weekly aerosol products at 0.5°×0.5° grid. MISR retrieves aerosol properties over a variety of terrain including 195 196 bright surface like deserts (Martonchik et al., 2004; Kahn et al., 2007 and 2010). In spite of the 197 fact that the satellite data are instantaneous observations at local overpass times (varying 198 between 10:30AM to 1:30PM for MODIS, MISR, and SeaWiFS) while models outputs are 199 diurnally varying, any bias caused by diurnal vs. instantaneous sampling is expected to be 200 small for monthly mean AOD. The study by Colarco et al. (2010) compared model simulated 201 AOD sampled at MODIS/MISR overpass times with those averaged over diurnal time steps 202 and found the differences to be small for monthly mean AOD, with only about 10% difference 203 in south America and southern Africa (i.e. biomass burning regions) and smaller elsewhere. 204 The climatology (averaged over the period of June 2006-December 2011) of vertical 205 extinction profiles from the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) layer 206 product version 3.01(onboard CALIPSO satellite) is used to evaluate the model simulated 207 aerosol vertical distribution in 2006 (CALIPSO 2011; Koffi et al., 2012). Only the CALIOP

208observations in 532 nm channel for nighttime are used because of their better signal-to-noise209compared to daytime observations. Three aerosol parameters are used to inter-compare210model simulations with CALIOP, namely AOD, Z_a (km) and F_{2km} (%). AOD is the integral of211extinction coefficient within the entire column (Eq.1). Z_a is defined as the averaged aerosol212layer height (Eq. 2), and F_{2km} is defined as the percentage of AOD located in the lowest 2 km

- 213 (Eq. 3) in the column.
- 214

215
$$AOD = \sum_{i=1}^{n} EXT_i \times \Delta Z_i$$
 (1)
216 $Z_a = \frac{\sum_{i=1}^{n} EXT_i \times Z_i}{\sum_{i=1}^{n} EXT_i \times Z_i}$ (2)

217
$$F_{2km} = \frac{\sum_{i=1}^{level of \ 2km} EXT_i \times \Delta Z_i}{\sum_{i=1}^{n} EXT_i \times \Delta Z_i}$$
(3)

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219 3.2 AERONET

In this study, we use AOD and AAOD data from the ground-based AERONET (Holben et al.,
1998) sites in South Asia. Monthly mean AOD and AAOD were analyzed over Kanpur, Lahore
and Karachi. Level-2 (version 2) data are used, which are cloud-screened and quality-assured
aerosol products with a low uncertainty of 0.01–0.02. Locations of the three stations are shown
in Fig.1 along with eleven in-situ measurement sites as described in the following Section 3.3.
The information of all fourteen ground-based measurement sites is given in Table 2.

226227 3.3 In-situ measurements

Modeled BC concentrations are also evaluated with the surface in-situ measurements from the
Integrated Campaign for Aerosols gases and Radiation Budget (ICARB) field campaign in
India over eight stations, which spread over Indian mainland and islands for the entire year of
2006. The BC data from ICARB field campaign were measured by inter-compared
aethalometers following a common protocol. More details of ICARB measurements can be
found in previous publications (e.g. Beegum et al., 2009 and Moorthy et al., 2013).

234 In order to examine the aerosol chemical composition (such as surface concentrations 235 of nitrate, sulfate, organic aerosol and black carbon) and meteorological conditions (such as 236 surface relative humidity and temperature) of winter haze over IGP in multi-models, we refer to 237 measurements from the Indian Space Research Organization- Geosphere Biosphere 238 Programme (ISRO-GBP) campaign which provided valuable information about aerosol physical, optical and chemical properties along the IGP during the wintertime of December 239 2004. For this study, four stations in IGP are selected because of their relatively complete 240 241 measurements. They are Hisar (Ramachandran et al., 2006; Rengarajan et al., 2007; Das et 242 al., 2008), Agra (Safai et al., 2008), Kanpur (Tripathi et al., 2006; Tare et al., 2006) and 243 Allahabad (Ram et al., 2012a), from western to eastern IGP. Note that the in-situ data used in 244 this study are obtained from the aforementioned references.

- 245
- 246 **4. Results**

- In this section, the aerosol simulations by multi-models are evaluated in comparison to satellite
 data and ground-based measurements in terms of temporal variation and spatial distribution
 (horizontally and vertically) over South Asia.
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251 **4.1 Interannual variability of AOD**

252 Figure 4a shows the annual averaged mean AOD over the entire South Asia domain (land only, 253 shown in gray shaded area) for the period of 2000-2007. AODs are 0.270±0.008 and 254 0.273±0.012 from MISR and SeaWiFS (SeaW) retrievals respectively, and 0.326±0.010 and 255 0.332±0.018 from MODIS Agua (MODIS-a) and Terra (MODIS-t) respectively. MISR AOD is 256 the lowest bound of four satellite retrievals. The difference in AODs among satellite data is 257 significant and could be up to 0.062 or 22% of MISR. Six out of seven models (except for 258 HAD) consistently underestimated AOD by 0.043-0.119 or 15%-44% relative to MISR. As 259 shown in Fig. 4b, over the central IGP region (77-83°E/25-28°N, denoted by the red box in Fig. 260 4a) where the hotspot of AOD is observed from satellites, the performance of the same six 261 models are even worse, with the annual averaged mean AOD underestimated by 20-57% 262 relative to MISR. Unlike other models, HAD shows comparable AOD with MISR and SeaWiFS 263 over the entire South Asia (Fig.4a), but exceeds all satellite data over the central IGP (Fig.4b), 264 higher than SeaWiFS and MISR by 47% and 58% respectively, and higher than MODIS-Terra 265 and Agua by 16% and 20% respectively. As shown in Figure 4a, the peak AOD in 2003 and 266 the low AOD in 2005 appear in all satellite data (except MODIS Agua in 2003), which are 267 associated with the strength of dust emissions during the dry season in the same years 268 (Kaskaoutis et al., 2012; Hsu et al., 2012; Ramachandran et al., 2013). However, all models 269 fail to reproduce the peak AOD in 2003, whereas only two models (GE5 and SPR) indicate the 270 low AOD in 2005.

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4.2 Seasonal cycle of AOD and AAOD over 3 AERONET stations

To further examine the details of underestimations occurring in most models, we compare the 274 275 model-simulated monthly variations of AOD and AAOD with the AERONET data at three selected sites in South Asia (Fig. 5). These locations represent different aerosol environments 276 277 in South Asia: Kanpur, an industrial city located in the central IGP, is influenced by high 278 anthropogenic emissions throughout the year and by the transported dust during pre-monsoon 279 (MAM) and early monsoon periods (JJ); Lahore, an urban city located in the western IGP, is 280 directly influenced by biomass burning in the pre-monsoon (MAM) and post-monsoon (ON) 281 seasons; and Karachi, an urban coastal city in Pakistan, is influenced by frequent dust 282 outbreaks, especially from the Arabian peninsula around early summer monsoon season (JJ). 283 A two-year period is chosen for each site based on the availability of AERONET 284 measurements. Three satellite datasets, namely MODIS-Agua, MISR, and SeaWiFS, are also 285 displayed to draw inter-comparison of AOD with AERONET data.

At Kanpur (first row of Fig. 5), strong seasonal variation of AERONET AOD (left column in Fig. 5) is evident with two peaks, one in May-July associated with dust outbreaks and the other in October-January associated with active open biomass burning as well as high 289 anthropogenic emissions. However, most models (except for HAD) only show the peak in May-290 July but miss the peak in October-January. Although the HAD model simulates two seasonal 291 maxima, they disagree with the peak months observed from AERONET. Overall, AOD from all 292 models have weak or negative correlation coefficients with AERONET data (from -0.34 to 0.34). 293 with four models anti-correlated with AERONET data (ECH, GIM, GOC and HAD), and one 294 with no correlation (GIE). AODs from six models are lower than those from AERONET as 295 indicated by the relative biases ranging from 0.31 to 0.74. In contrast, HAD model 296 overestimate the AOD by 44% (relative bias of 1.44). As for AAOD (right column in Fig. 5), 297 models are much lower than the AERONET data by a factor of 2 on average, suggesting the 298 underestimation of BC loading or weak aerosol absorption strength in models (see more 299 analysis of BC in Section 4.5).

300 At Lahore (second row of Fig. 5), AERONET data are mostly available in the year 2007, 301 when only five model results are available (no HAD and ECH for 2007, see Table 1). Lahore is 302 located in the Punjab region, which is an agriculture region known as the "breadbasket" for 303 Pakistan and India. The enhanced AERONET AOD and AAOD are evident at Lahore during 304 October-November, which is linked to the agricultural waste burning after harvest. However, all 305 five models largely underestimate AOD and AAOD in the October-November period. This 306 suggests that emissions from agriculture waste burning are likely underestimated in GFED2 307 that are used by the models (discussed in Section 6.4). Compared to observations, HAD again 308 showed abnormal seasonal variation at Lahore, similar to that at Kanpur, with extreme high 309 AOD in October though.

310 At Karachi (third row of Fig. 5), a unimodal seasonal distribution is revealed in 311 AERONET AOD data, in contrast to the bimodal seasonal variation at Kanpur. The maximum 312 AOD around July is associated with the wind-driven mineral dust from the Arabian Peninsula, 313 which is captured by the models as indicated by relatively strong correlation from 0.58 to 0.91 314 (except HAD. Note ECH is not available for 2006-2007). However, similar to other sites, AOD 315 from all models are too low in late autumn to winter. Models also fail to capture the relatively 316 higher AAOD around November that is associated with smoke transported from agriculture 317 waste burning in northwestern IGP (i.e. the area around Lahore) (Badarinath et al., 2009a,b). 318 Overall, in comparison with AERONET at three sites, most models tend to significantly 319 underestimate AOD in October-January when aerosols from agriculture waste burning and 320 anthropogenic activities are dominant. On the other hand, the monthly variations and 321 magnitudes of AOD from the satellites are in general similar to those from AERONET. As an 322 exception, MODIS-Agua is biased high (up to a factor of 2) during pre-monsoon and monsoon 323 months. This overestimation of AOD partially results from low bias of surface reflectance under 324 dusty conditions in the MODIS Dark-Target aerosol retrieval algorithm (Jethva et al., 2009).

In order to diagnose the discrepancies between models and AERONET data, the
 individual component AOD from four models (HAD, GE5, SPR and GOC, unavailable from
 other three models) are examined at Kanpur for 2004 in Fig. 6. We choose the year of 2004
 because ISRO-GBP campaign took place in the same year (see Section 3.3 and Section 6), so
 that we can intercompare AERONET data with that in ISRO-GBP campaign. In December and

330 January, AOD from AERONET data is around 0.7, dominated by anthropogenic contributions 331 (about 75%, estimated by Tripathi et al. 2006). All four models have difficulties to capture the 332 magnitude of AOD in December and January. Among them, AOD from HAD (upper left panel 333 in Fig. 6) matches relatively well with AERONET data, capturing about half of the observed 334 value. Interestingly, nitrate (NO₃) AOD is the major component in HAD, contributing to 50% of 335 total modeled AOD. In contrast, three other models (SPR, GE5 and GOC) largely 336 underestimate the peak in the winter (December and January) by up to a factor of 7. As a 337 common problem, these three models do not include the nitrate aerosol component. During the 338 months of May to July, coarse mode aerosol (i.e. dust) contributes mostly to total AOD (> 60%) 339 based on studies with ground based sun/sky radiometer data (e.g. Srivastava et al., 2012a). 340 SPR and GE5 capture this feature while HAD and GOC underestimate the contribution of dust. 341 In the HAD model, AODs from nitrate alone during April and October are comparable to 342 column total AERONET AOD, indicating a problem in representing seasonal variation of nitrate 343 in HAD, as shown in Fig. 5. Instead, nitrate aerosol is expected to peak in winter because of 344 high relative humidity and low temperature over IGP that favor the formation of NH₄NO₃ (Feng and Penner 2007; Ram et al., 2010b; Ram et al., 2012b). 345

Overall, Fig. 6 demonstrates that the magnitudes and seasonal cycles of aerosol
compositions are quite different across the models. Further examination of the model
diversities will be discussed in Section 5.

349

4.3 Spatial distribution of AOD in different seasons

351 In this section, we compare the spatial distributions of AOD over the entire South Asia and 352 neighboring oceans among four satellite products (MODIS-Terra, MODIS-Agua, MISR, and 353 SeaWiFS) and seven model simulations during the winter monsoon (DJF), pre-monsoon 354 (MAM), summer monsoon (JJAS) and post monsoon (ON) phases averaged over 2000-2007, 355 shown in Fig. 7a-b. Locations of the three aforementioned AERONET stations are also labeled 356 in the maps for reference. In general, the spatial distribution of AOD is closely associated with 357 the emission source over South Asia, and the aerosol abundance in the atmosphere is 358 modulated by meteorological conditions, such as efficient atmospheric dispersion associated 359 with the strengthened westerly flow in March-July, high wet removal associated with the 360 monsoon rainfall in June-September, and stable atmospheric conditions and thus less efficient 361 atmospheric dispersion in December-February.

362 During the winter season (DJF), local anthropogenic sources dominate over dust, 363 contributing as much as 80% (±10%) to the aerosol loading (Ramanathan et al., 2001; Tripathi 364 et al., 2006). The maximum AOD is found in the central and eastern IGP based on four 365 satellite datasets as shown in Fig. 7a, which coincides with clusters of coal-based large 366 thermal power plants (capacity >1970 MW) (Prasad et al., 2006). The natural topography (i.e. 367 gradually decreased elevation eastward but narrow opening to the Bay of Bengal as shown in 368 Fig. 1) is conducive to the accumulation of aerosol over central and eastern IGP. Additionally, 369 the winter season is characterized by relatively stable atmospheric conditions that traps 370 pollutants in the shallow atmospheric boundary layer (ABL), leading to strengthened hazy

371 conditions in the IGP (Girolamo et al., 2004; Gautam et al., 2007). The outflow of aerosols to 372 the Bay of Bengal is clearly depicted by satellite data. As shown in the first column of Fig. 7b, 373 however, only the HAD model shows the observed spatial pattern and magnitude of AOD, 374 although it overestimates AOD over eastern IGP. Other models greatly underestimate the high 375 AOD over IGP by 50% on average. In addition, the observed north-south gradient of AOD is 376 not captured by most models, with SPR showing no gradient and ECH and GIM showing 377 opposite gradient. The model underestimation over the Indian subcontinent in winter is 378 probably owing to missing aerosol species such as nitrate aerosol suggested by Fig. 6, 379 incorrect meteorological fields such as air temperature and relative humidity, or the 380 underestimation of anthropogenic emissions (discussed in more details in Section 6).

381 Starting from the pre-monsoon season (MAM), the entire South Asia is characterized by 382 high AOD mainly due to the mineral dust transported from the arid and desert regions in 383 southwest Asian dust sources by westerly winds, with maximum AOD over western IGP seen 384 from most satellites (Fig. 7a). As shown in the second column of Fig. 7b, five models (GOC, 385 SPR, GIM, GIE and GE5) partially capture this observed spatial distribution and magnitude. However, the HAD model shows high biases of AOD over northern India due to nitrate (refer to 386 387 Fig. 6). A higher nitrate concentration than dust is unrealistic because the contribution of dust 388 to the total AOD has been reported to be over 60% during pre-monsoon season by Srivastava 389 et al. (2012a) based on the ground based sun/sky radiometer data. The dust source in the 390 northwestern parts of South Asia is weak in HAD (Fig. 7b). Additionally, the ECH model shows 391 very low AOD and little dust over IGP associated with its small dust size in coarse mode (Table 392 1). Despite these deficiencies, model simulations over South Asia during the pre-monsoon 393 season are still closer to the satellite data than those during winter, with the model-averaged 394 AOD capturing 65% of the satellite data in the pre-monsoon season compared to only 50% in 395 winter.

396 During the monsoon season (JJAS), dust transported from the Arabian Peninsula by the 397 strong southwesterly winds explains the high AOD over northwestern India. High AOD over the 398 Arabian Sea and southwest Asia is evident in MODIS and MISR (Fig. 7a). As shown in the 399 third column of Fig. 7b, most models reproduce both the spatial distribution and the magnitude 400 of AOD during this season, implying that these models capture dust emission over the Arabian 401 Peninsula and its transport to South Asia. However, it should be noted that during the 402 monsoon season the monthly mean AOD from MODIS is likely to be biased high as shown 403 earlier in Fig 5, partly due to underestimated surface reflectance.

404 During the post-monsoon season (ON), the southwesterly flow significantly weakens, 405 and thus dust transported to the Indian subcontinent is lower compared to the pre-monsoon 406 and monsoon seasons. Based on the spatial distributions from satellite data (Fig. 7a), high 407 AOD is found along IGP with maxima over western IGP including Punjab, Haryana and 408 western Uttar Pradesh that are associated with biomass burning from agriculture waste fires. 409 With the aid of northwesterly winds, aerosols are transported to the central IGP along the 410 valley as well as the region to the south (Badarinath et al., 2009a, b). However, none of the 411 models capture these features (the fourth column of Fig. 7b), indicating the biomass burning

emissions are severely underestimated in the current inventory based on GFED2, which will be
discussed further in Section 6.4. In contrast to the underestimations by other models, HAD
overestimated AOD over IGP due to the high amount of nitrate (Fig. 6).

415

416 **4.4 Aerosol vertical distribution**

417 Figure 8 shows the comparison of aerosol extinction profile among models and with CALIOP 418 data in four seasons. In order to represent the latitudinal gradient of aerosol vertical profiles, 419 two locations are chosen, Kanpur in northern India and Hyderabad in central India (refer two locations to Fig. 1). The CALIOP aerosol extinction profile over Kanpur (Fig. 8a, 2° x 2° box 420 averaged around the station location) reaches a maximum value of 0.4 km⁻¹ at the altitude < 1 421 km during winter (Z_a =1.18 km) but decreases rapidly upward and diminishes around 4 km. 422 423 Note that low values near the surface (within 180 meters) in CALIOP profiles are likely due to 424 the contamination by the surface return (CALIPSO, 2011; Koffi et al., 2012). In contrast with 425 the relatively stable lower troposphere in winter, boundary layer mixing, convection, and 426 transport are strengthened in pre-monsoon season. As a result, aerosols are more efficiently mixed vertically, with Z_a from CALIOP almost doubled from the season of DJF to MAM (from 427 428 1.18 to 2.18 km). The aerosol vertical mixing is relatively uniform within the lowest 2 km and 429 extends to higher altitude around 6 km in MAM. The aerosol extinction near the surface in 430 MAM is only 60% of its DJF values with the fraction of AOD in the lowest 2 km reducing from 431 84% in DJF to 52% in MAM. The aerosol profile during monsoon season (JJAS) is similar to 432 that in pre-monsoon period but with a slightly lower value of Z_a as 2.02 km; and the profile 433 during the post-monsoon is similar to that in the winter but with a slightly higher value of Z_a 434 as1.24 km.

435 Most models, especially GE5, capture the observed seasonal variation of Z_a (and F_{2km}) 436 over Kanpur, with lower Z_a (higher F_{2km}) during wintertime (DJF) and post-monsoon season 437 (ON), while higher Z_a (lower F_{2km}) during the pre-monsoon (MAM) and monsoon seasons 438 (JJAS). The profiles and magnitude in models, however, are guite different from those of 439 CALIOP. At Kanpur in DJF, most models (except for HAD and GIE) largely underestimate 440 AOD by 59% (ECH) to 85% (SPR), consistent with the preceding results (Fig. 5-7). In particular, 441 the extinction coefficient in the lowest 2 km is largely underestimated, with F_{2km} varying from 442 68% (GIM) to 87% (GE5) among these five models in contrast to 84% in CALIOP (Fig. 8a). At 443 Hyderabad in central India (Fig.8b), models agree better with the CALIOP during the winter 444 (DJF) and post-monsoon (ON) seasons. At both stations, models agree better with CALIOP 445 during the dust-laden pre-monsoon (MAM) and monsoon (JJAS) seasons than during two 446 other seasons, consistent with the results in Fig.7a-b. There are some extremes of model 447 simulated vertical profiles. For example, HAD produces extremely high extinction coefficients 448 close to the surface at Kanpur throughout all seasons that are a factor of two greater than 449 CALIOP in the season of DJF and a factor of ten greater in ON; GIE and GIM are greater than 450 CALIOP by a factor of four and seven close to the surface in JJAS, respectively; and GIE 451 exhibits extremely large extinction coefficients between 2 and 3 km in all seasons, which is not found in CALIOP. 452

454 **4.5 Monthly BC surface concentration**

455 Figure 9 shows the observed and modeled monthly surface BC concentration in the year of 456 2006 (2005 from model ECH) at eight ICARB stations (refer the locations to Fig.1). In general, 457 the magnitude of BC surface concentrations is closely related to the strength of emission 458 source, with higher values in northern India where higher BC anthropogenic emissions are 459 located (refer the spatial pattern to Fig.2). The highest BC surface concentration is particularly found in the largest Indian city Delhi, with a value of 27µg m⁻³ in January. In contrast, BC 460 surface concentration is lower in the remote sites, such as the island sites (Minicoy and Port 461 Blair) and mountain site (Nainital), not exceeding 2.8µg m⁻³. The observed surface BC 462 concentration exhibits pronounced seasonal variation with higher values found in the winter 463 464 and post-monsoon seasons and lower values in the spring and summer, which can be 465 attributed to the seasonal variations of emission, atmospheric boundary layer (ABL) depth 466 (affecting vertical mixing), and rainfall (removing BC from the atmosphere). It was reported by 467 previous studies that total BC loading over South Asia mainly resulted from biofuel emissions 468 in winter along with coal burning in the vicinity of the measurement location (e.g. Ali et al., 469 2004; Singh et al., 2008; Beegum et al., 2009; Srivastava et al., 2012b). In comparison with 470 observation, modeled BC surface concentrations at all stations except Nainital (a mountain 471 site) and Kharapgpur are too low, especially in winter. In particular, at Delhi and Hyderabad -472 two very large cities with populations of 16.75 and 6.81 million respectively (Table 1), all 473 models show a pronounced low bias in the winter, capturing only 3%-19% of the observed 474 values. As a matter of fact, the models have difficulties to reproduce the observed high 475 pollution levels only near the emission sources such as urban cities (e.g. Delhi and 476 Hyderabad), but also in more remote locations (e.g. over the mountain site of Nainital and the 477 island sites of Minicov and Port Blair). At Minicov and Port Blair, where the observed BC 478 concentrations are relatively low, models capture only about 10%-38% of the observed values. In addition to the fact that modeled AODs were also found to be significantly low in comparison 479 with both AERONET point observations and with the multiple gridded satellite data from 480 481 MODIS, SeaWiFS (both 1°x1° resolution) and MISR (0.5°x0.5° resolution) on regional scales, 482 as shown earlier in Fig. 5 and Fig. 7, the underestimations of modeled BC and AOD in winter 483 are more likely due to other factors than coarse model resolution, which will be discussed in 484 details in Section 6. As an exception, the simulated BC surface concentrations are found to 485 have a better agreement at Kharagpur, a semi-urban city with populations less than 1 million, where models capture 20%-100% of the observed value. This contrast with other stations is 486 487 possibly attributed to the fact that BC loading at Kharagpur mainly comes from coal-fired power 488 plants (Nair et al., 2007), which are likely well represented in the emission data (discussed 489 further in Section 6.3).

490

491 **5. Model diversity**

492 Clearly, there is a large diversity existing among models in simulating AOD and BC

493 concentrations as shown in Fig. 4-9, despite similar emission datasets used in these models

494 (see Section 2.2 and Table 1). It is seen that models with the same emissions datasets

495 produce guite different results. For example, at Kharagpur, shown in the upper right panel of Fig.9, the surface concentration of BC from the SPR model is four times as large as that from 496 497 GIM, although both models use the same anthropogenic emission (A2-ACCMIP) and biomass 498 burning emission (GFED2). Similarly, surface concentration of BC in the HAD model is twice 499 as that of GOC, although the same emissions (A2-MAP and GFED2) are used in both models. 500 Such substantial differences indicate that the large diversity among model simulations is due to 501 factors other than the differences in emissions. Textor et al. (2007) also found that the 502 differences in the model treatment of atmospheric processes (e.g., wet removal, dry deposition, 503 cloud convection, aqueous-phase oxidation and transport), assumptions of particle size, 504 mixture, water uptake efficiency, and optical properties are more responsible than emission for 505 the model diversity. More recently, Bond et al. (2013) pointed out that large differences in 506 modeled horizontal and vertical transport are mostly responsible for the inter-model diversity of 507 BC distributions.

508 The multi-model diversity (defined as the percentage of the standard deviation to the mean 509 of results from the seven models) over South Asia in 2006 (2005 from the model ECH) is 510 summarized in Table 3 (also demonstrated in supplement Fig. S1-3). In general, on an annual 511 basis, we found the following features: (1) For aerosols with anthropogenic origin (i.e. BC, OA 512 and SO₄), the diversity of dry deposition among models is large, with diversity ranging from 513 41% to 46% across these three species. Correspondingly, the fraction of dry deposition to total 514 deposition shows 29-40% diversity for the same three species. In contrast, the diversity of wet 515 deposition is relatively smaller with a range from 15 to 22% across these three species. The 516 chemical production of sulfate in gas phase among models (four models) has large diversity 517 about 66%. (2) For mineral dust, the emission itself has very large diversity among the models 518 about 124%, leading to a similarly large diversity of dry deposition (aerodynamic dry deposition) 519 + gravitational settling) of 115%. The difference of treatment of dust size bin in models 520 contributes significantly to these diversities (see Table 1). In contrast, diversity of dust mass 521 loading and AOD are much smaller at 45% and 22% respectively. (3) BC has the largest 522 model diversity of mass extinction efficiency (MEE) at 51%, compared to 25% and 27% for 523 SO₄ and OA respectively.

524 We further examine the aerosol refractive index at 550nm for each species as listed in 525 Table 1. The real part of refractive indices at 550nm is similar among the seven models, but 526 the imaginary part (representing light absorption) is different. In the case of BC, the most 527 absorbing aerosol, its imaginary refractive index is 0.44 in four models and 0.71 in three 528 models, which are lower than the value of 0.79 recommended by Bond and Bergstrom (2006). 529 For dust, the light absorption at 550nm is significantly less than that of BC. The imaginary 530 refractive index of dust ranges from 0.001 (ECH) to 0.008 (GE5), a range that is much wider 531 than that of BC. The differences in the absorption properties, together with the differences of 532 model simulated BC and dust amount, contribute to the diversity of model calculated AAOD at 533 37%.

534

535 6. Possible causes of model underestimation of aerosols over South Asia

As shown in Section 4, AOD, AAOD and BC surface concentration over South Asia are consistently underestimated in seven global models used in this study, in particular during winter and the post-monsoon season. Such underestimation seems to be a common problem in other models as well (e.g. Reddy et al., 2004; Ganguly et al., 2009; Nair et al., 2012). AOD and surface BC concentrations are most severely underestimated over the IGP (the main region of anthropogenic emissions). Several possible causes for these underestimations are suggested below.

543

544 6.1 Wintertime relative humidity (RH) over the IGP

Foggy days with high near-surface relative humidity are very common during wintertime over
IGP (Gautam et al., 2007). For example, Kanpur was subjected to heavy fog or haze for about
>65% days in December 2004, with averaged surface RH of about 75% and the surface
temperature about 14.6°C (Tripathi et al., 2006). Low precipitation and thus low wet removal in
winter further contributes to accumulation of aerosols (Tripathi et al., 2006).

550 Figure 10 shows comparisons between models and in-situ measurements (ISRO-GBP land campaign) at four stations located in the IGP region in December 2004. Comparisons are 551 shown for surface meteorological conditions (RH and temperature); surface aerosol 552 concentrations of SO₄²⁻, NO₃⁻, OA and BC; and columnar AOD and AAOD. AODs from the 553 554 models are only 10% to 50% of the observed values at Kanpur. Interestingly, we found that RH 555 in six of the seven models (except for HAD) only range from 11 to 35% at Kanpur, much lower 556 than the measured RH of 75% (first row, Fig. 10). This large underestimation of RH could be 557 partly due to the warm bias of air temperature by 1.7-7.5 °C across models (second row, Fig. 558 10) and thus high bias of saturation water vapor pressure and low bias of RH. Under such dry 559 conditions in models, the hygroscopic growth of soluble aerosols is consequently suppressed. 560 Averaged over these four IGP stations, for example, if RH is improved from the modeled 21% to the observed 66%, mass extinction efficiencies (MEE) of SO_4^{2-} would be doubled, and those 561 of OC and NO₃⁻ would be enhanced by 50% (Fig. 11). It is interesting that the HAD model, in 562 563 which the simulated AOD matches observed one relatively better, is the only model with high 564 bias of RH.

565 In addition to favor hygroscopic growth, foggy conditions also favor the formation of 566 secondary inorganic aerosol through the aqueous-phase reactions. This phenomenon was supported by the observations of increased aerosol number concentration and surface SO₄²⁻ 567 568 concentrations under foggy conditions at Kanpur (Tare et al., 2006), Hisar and Allahabad (Ram 569 et al., 2012a). High RH and lower temperature in winter also favor the formation of NH₄NO₃ by 570 the reaction of nitric acid (HNO₃) with NH₃ (Feng and Penner 2007; Ram et al., 2010b; Ram et 571 al., 2012b). However, the lack of representing foggy conditions in current models, which is 572 indicated by the low bias of RH, would suppress these reactions in winter. Hence, it is not surprising that the surface mass concentrations of SO_4^{2-} and NO_3^{-} in models are found to be 573 574 much lower than the observed values. As shown in Fig. 10, all models underestimate the surface concentration of SO_4^{2-} , with capturing merely from 5% (GIE and GIM) to 50% (GE5) of 575 the observed value. SO₄²⁻ concentration, however, is found low in HAD as well although with 576

- 577 high relative humidity. The specific reason is unclear yet. Among three models that include
- NO_3^{-} , GIE and GIM produce extremely low NO_3^{-} concentrations that are only 0.1% of the
- observed amount, whereas HAD captures about 38% of the observation. The model
- underestimations of surface aerosol concentrations might be caused by other factors as well,
- such as unaccounted for anthropogenic emissions (see section 6.3) or insufficient oxidant
- amounts (H_2O_2 and OH); however, the lack of representing foggy conditions or the low bias of RH in the models appears to be a critical factor contributing to the overall underestimation of aerosols.
- 585

586 6.2 Nitrate component

- As shown in Fig. 10, the observed surface concentrations of NO_3^- are comparable to or even 587 higher than those of SO₄²⁻ at four stations (e.g. 14.9 μ g m⁻³ of SO₄²⁻ and 15.7 μ g m⁻³ of NO₃⁻ at 588 Kanpur, and 14.1 μ g m⁻³ of SO₄²⁻ and 31.4 μ g m⁻³ of NO₃⁻ at Agra). However, NO₃⁻ is either 589 590 missing in the models (GOC, ECH, SPR, GE5) or much too low (especially in GIE and GIM). 591 Interestingly, AOD is closer to observations in the HAD model than in other models, which is 592 not only apparent at 4 stations in IGP (Kanpur, Agra, Allahabad and Hisar) (Fig. 10) but also 593 over entire South Asia (Fig. 7b). Such agreement is partly associated with its inclusion of NO₃⁻ 594 (Fig. 6) and aforementioned high relative humidity in winter (Section 6.1). This study 595 underscores the importance of NO_3^- to adequately represent the total AOD over South Asia.
- 596

597 **6.3 Anthropogenic/Biofuel emission amounts and seasonal variation**

- 598 The uncertain and inadequate representations of aerosol emissions over South Asia have 599 been pointed out by previous studies (e.g. Sahu et al., 2008; Ganguly et al., 2009; Nair et al., 2012; Lawrence and Lelieveld, 2010). The results in this study further prove this issue. At 600 Kanpur, the models underestimate not only surface concentrations of SO_4^{2-} and NO_3^{-} as 601 discussed earlier but also those of OA and BC, with capturing only 8% (GIE and GIM) to 602 603 75% (SPR) of the observed OA values, and 8% (GIE and GIM) to 46% (SPR) of the observed BC values, respectively. At other stations in the IGP such as Agra, Allahabad and Hisar (Fig. 604 10), the surface concentrations of OA, BC, SO_4^{2-} and NO_3^{-} are underestimated in a similar 605 606 degree by all models, although these stations are less populated than Kanpur. AOD and 607 AAOD, indicating columnar aerosol loading, are also underestimated by all these models. It is well known that air pollutants are confined to near surface in winter due to the low ABL, 608 609 thereby the results above suggest that the anthropogenic emissions used by the models (i.e., A2-ACCMIP and A2-MAP) are likely biased low. BC emissions in year 2000 over India from 610 A2-ACCMIP and A2-MAP are 0.5Tg yr⁻¹, which is at the low end of a group of emission 611 612 inventories, for instance, lower than those considered by REAS and GAINS-2008 emission inventories (Fig. 5a in Granier et al., 2011) by 40% or 0.3Tg yr⁻¹. With the REAS emission 613 inventory, Nair et al. (2012) reported that the simulated BC surface concentration agreed better 614 615 with observations at Kharagpur.
- 616 Different from other regions in northern hemisphere where fossil fuel burning and industrial 617 processes tend to dominate, biofuel and open biomass burning in South Asia contribute two-

618 thirds of carbon-containing aerosols to form the dense brown clouds in winter (Gustafsson et 619 al., 2009). Over India, 42% of total BC emission is from biofuel, which is believed to be the 620 largest source of BC, with the remaining 33% from open biomass burning and 25% from fossil 621 fuel (Venkataraman et al., 2005). The percentage of biofuel is high because residential heating 622 and cooking (burning of wood, paper or other solid wastes) is guite common in South Asia, especially among the underprivileged, leading to large amount of smoke comprised mainly of 623 624 black carbon and condensed semi-volatile organics. Based on in-situ measurements, the ratios 625 of OC/BC surface concentrations were reported as high as 8.0±2.2 at Allahabad (Ram et al., 2012a) and 8.5±2.2 at Hisar (Rengarajan et al., 2007) in December 2004, indicating a major 626 627 emission source from biomass combustion including biofuel and open biomass burning 628 (Husain et al., 2007). However, in this study, fossil fuel are the dominant emission sources 629 instead, because the ratio of OC/BC anthropogenic emission (from combination of fossil fuel 630 and biofuel) in A2-ACMMIP (A2-MAP) emission database is 3.2 (2.6) (see Section 2.2) over 631 South Asia, and thus it is not surprising that the ratios of OC/BC surface concentrations are 632 found only varying 0.4-4.0 across models at Allahabad and 0.6-3.8 at Hisar. Although the ratio 633 of OC/BC in open biomass burning emission database is higher with a value of 8.0, open 634 biomass burning emissions are very low in winter, only 4% of anthropogenic emissions (see 635 Fig.2 and Fig.3). Furthermore, we found that the simulated BC surface concentrations by most 636 models agree better with the observations at Kharagpur than at other stations (Fig. 9). As 637 reported by Prasad et al. (2006), the sources of BC at Kharagpur located in eastern IGP were 638 mainly linked to the clusters of the coal-based industries there. Therefore, this contrast 639 suggests that the fossil fuel emissions are likely better represented than the biofuel emissions 640 in the A2-MAP and A2-ACCMIP emission inventories. In addition, the lack of seasonal 641 variation in anthropogenic emission datasets would amplify the underestimation of aerosol 642 amount during the winter when biofuel emissions are prevalent. In sum, the model 643 underestimation of anthropogenic OA and BC concentrations in winter is mostly due to the 644 underestimation of biofuel emissions.

645

646 6.4 Agriculture waste burning emissions

647 During the post-monsoon season (October-November), the extensive agriculture waste 648 burning after harvest in northwest India (e.g., Punjab) makes a large contribution to the dense 649 haze over South Asia based on previous observational studies (Vadrevu et al., 2011; Sharma 650 et al., 2010). The agricultural fires in this area are evident in the MODIS fire count product. 651 Smoke plumes from Punjab also impact the downwind regions by eastward transport along IGP and southward to central-south India (Sharma et al., 2010; Badarinath et al., 2009a, b). 652 653 Over India, the contribution from open biomass burning to the total BC emission is 654 significant, about half of anthropogenic emissions (i.e. biofuel plus fossil fuel emissions) 655 (Venkataraman et al., 2005). The biomass burning contribution is evident based on the AERONET data at Lahore, where AAOD enhances by 70% in November (after harvest) from 656 657 previous months (Fig. 5), and its contribution is also clearly seen in the MODIS-Terra and 658 Aqua data with the maximum AOD found near Lahore in the post-monsoon season (the fourth column of Fig. 7a). BC emission from open biomass burning (based on GFED2) used by the
models, however, is less than 1% of that from anthropogenic sources (comparing Fig. 2 and
Fig. 3) during the post-monsoon season, both on regional average and in areas around Lahore,
Therefore, it is not surprising that all models fail to capture high AAOD and AOD in this season
(Fig. 5 and Fig. 7b). The underestimation of BC emission from agriculture waste burning also
implies a similar degree of underestimation of OC from the same source.

The open biomass burning emission from GFED2 is derived from MODIS burned area products. It was previously reported that the small fires such as agricultural waste burning were largely missing in the GFED product (e.g. van der Werf et al., 2010; Randerson et al., 2012). The agricultural waste burning area is usually underestimated or overlooked in MODIS because the size of agriculture fires is too small to generate detectable burn scars in the 500 meter pixel resolution of MODIS product (van der Werf et al., 2010; Randerson et al., 2012).

672 6.5 Other factors

671

673 Other factors can also cause the models to underestimate AOD. For example, the observed 674 ratio of secondary organic carbon (SOC) to primary OC is 30% - 40% in several stations 675 located in North India, suggesting a significant contribution from SOC (Rengarajan et al., 2007; 676 Ram and Sarin et al., 2010a). However, only two models include a resolved SOC chemistry. In 677 addition, although the dust emission is minimal in winter compared to anthropogenic emission, 678 dust sources from road traffic, soil re-suspension, and construction activity in the urban regions 679 of the IGP (Tripathi et al., 2006; Tiwari et al., 2009) could be important, which are not 680 considered in the current models.

Some difficulties with the models might be associated with the coarse spatial resolution (at $1.1^{\circ} - 2.8^{\circ}$, see Table 1). Considering the complex terrain variations over South Asia, especially the valley-type topography of the IGP region with the towering Himalaya in the north (Fig. 1), the aerosol processes may not be adequately represented at such coarse spatial resolution. In addition, because of the non-linearity of wind-dependent dust emission and RHdependent aerosol hygroscopic growth, a finer model spatial resolution would result in a higher dust emission and AOD (Bian et al., 2009).

688 Another important factor contributing to high surface aerosol concentrations in winter 689 over South Asia is the shallow wintertime ABL that suppresses ventilation thereby trapping 690 pollutants near the surface. At Kanpur, ABL height is about 200 m in winter according to the 691 observations (Tripathi et al., 2006; Nair et al., 2007). However, the averaged ABL in GOC and 692 GE5 models are 400-500 m in the study region (other models did not provide this information), 693 allowing more efficient vertical mixing to dilute the surface concentrations and thus contributing 694 to the low bias of surface aerosol concentration (Fig. 9 and 10). Therefore, a better-695 constrained ABL would be helpful to reduce the model bias of surface concentrations. Here we 696 would like to iterate, however, that the columnar AOD and AAOD during wintertime is 697 underestimated by the models as well, despite to a lesser degree than the underestimation of 698 surface concentration (for example, model-simulated BC concentrations are too low by a factor 699 of about 10, compared to the underestimation of AAOD by a factor of \sim 3). Considering the 700 results that both aerosol surface concentration and columnar loading are underestimated, the

- dominant factor in underestimating aerosol surface concentrations by these models is likely the
 underestimation of the emissions in wintertime, as addressed in Section 6.3.
- 703

704 **7. Conclusions**

- In this study, the aerosol simulations for 2000-2007 from seven global aerosol models are
- evaluated with satellite data and ground-based measurements over South Asia, in particular
- over IGP, one of the heavily polluted regions in the world. The high AOD over IGP is
- associated with persistent high aerosol and precursor gas emissions (such as dust, SO_2 , NO_x ,
- NH₃, OA and BC) from local and upwind regions, and with its valley-type topography (bounded
- 510 by the towering Himalaya) that is conducive to trapping both anthropogenic and dust aerosols
- in this region. The main results of this study are summarized below.
- 1. Averaged over the entire South Asia for 2000-2007, the annual mean AOD is about 0.27-
- 0.33 from satellites retrievals. Six out of seven global models consistently underestimate
- the annual mean AOD by 15%-44% compared to MISR, the lowest bound of four satellite
- datasets used in the present study. The model performances are worse over northern India.
- In general, the underestimation of aerosol loading is mainly found during the winter and
 post-monsoon months when anthropogenic and open biomass burning emissions are
- dominant.
- 7192. During wintertime (DJF), six out of seven models largely underestimate columnar AOD and720AAOD over Indian subcontinent, and the underestimations of aerosol extinction generally721occur in the lower troposphere (below 2 km). The simulated surface mass concentrations of722 SO_4^{2-} , NO_3^{-} , OA and BC are as small as 0.1-60% of the observed values in winter. Several723possible causes for the common underestimations are identified: (a) the wintertime near-
- surface relative humidity is too low (e.g., about 20% in IGP in six out of seven models,
 compared to the observed value of > 60%) such that the hygroscopic growth of soluble
- 726 aerosols and formation of secondary inorganic aerosol (NO₃⁻ and SO₄²⁻) are suppressed:
- 727 (b) NO₃⁻ is either missing or inadequately accounted for; (c) anthropogenic emission,
- especially from biofuel in winter, is underestimated in the emission datasets. The lack of
 seasonal variation of emissions amplifies the discrepancies in winter.
- 3. During the post-monsoon season (ON), none of the models capture the observed high
 AOD over western and central IGP. AAOD and BC surface concentrations are
 underestimated at the stations in IGP as well. Such discrepancy is attributed largely to the
 underestimation of open biomass burning in the satellite-based emission inventory
 (GFED2). It is likely due to missing small agricultural waste burning that is difficult to be
 retrieved by the satellite remote-sensors.
- 4. As for the inter-model diversity, the results show that the largest diversity occurs in the treatment of dry deposition, with diversity of dry deposition amount ranging from 41 to 46% for BC, OA, and $SO_4^{2^-}$. In contrast, the diversity of wet deposition is smaller, from 15 to 22% across three species. For mineral dust, the emission itself has very large diversity among the models (about 124%), leading to a similar diversity of dry deposition (aerodynamic dry deposition + gravitational settling) as of 115%, although the diversity of dust AOD is much
- smaller at 22%.

To sum up, we have identified the major discrepancies of seven state-of-the-art global aerosol models in simulating aerosol loading over South Asia. Results from this study suggest directions to improve model simulations over this important region, including improving

- meteorological fields (particularly relative humidity), revising biofuel and agriculture fire
- emission inventories, and adding/improving NO₃. Currently, we are working on quantifying the
- factors that cause the model underestimation by ranking their importance via a series of model
- sensitivity experiments using the GEOS5 model. Our ongoing work includes adjusting the
- model spatial resolution, emission strength, and meteorological variables and adding nitrate,
- which will be presented in subsequent publications. Here, we also would like to suggest to
- establish more systematic measurements, especially long-term (at least one year-around)
- surface and vertical characterization of aerosol composition, precursor gases, optical
- properties, and meteorological fields (e.g. temperature, winds, and relative humidity), because
 they are essential for understanding the aerosol physical and chemical characterization.
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185 **Tables**

186

187 Table1. General information of models used in this study.

				-			-	
Model		HadGEM2	GOCART-	ECHAM5-	GISS-	GISS-	SPRIN-	GEOS5-
			v4	HAMMOZ	modelE	MATRIX	TARS	GOCART
ID		HAD	GOC	ECH	GIE	GIM	SPR	GE5
Time range		2000-2006	2000-2007	2000-2005	2000-2008	2000-2007	2000-2008	2000-
Time Tunge								2008
Snatial		1 8x1 2x38	2 5×2×30	2 8x2 8x31	25x2x40	25×2×40	1 1×1 1×56	2 5×2×72
Resolution a		10 11 00	10 1 00	10 10 01	1.0 1 10	210 2 10	111 111 00	
Anthrn Emi b		A2-MAP	A2-MAP	A2-MAP	A2-ACCMIP	A2-ACCMIP	A2-ACCMIP	A2-ACCMIP
manp. Lini.			112 1.111		112 110000111		112 110000111	112 11001-111
BB Emi. ^c		GFED2	GFED2	GFED2	GFED2	GFED2	GFED2	GFED2
Met. Field		ERA- Interim	GEOS-DAS	ECMWF	NCEP wind	NCEP-wind	NCEP/	MERRA
				analysis			NCAR	
Refrac	SO42-	1.53 – 1e-7 <i>i</i>	1.43-1e-8 i	1.43-1e-8 <i>i</i> ^e	1.528-1e-7 <i>i</i>	1.528-1e-7 <i>i</i>	1.43-1e-8 <i>i</i>	1.43-1e-8 <i>i</i>
-tive	BC	1.75 – 0.44 <i>i</i> (FF) ^d	1.75-0.44 <i>i</i>	1.85-0.71 <i>i</i>	1.85-0.71 <i>i</i>	1.85-0.71 <i>i</i>	1.75-0.44 <i>i</i>	1.75-0.44 <i>i</i>
index	OA	1.54 – 0.006i (FF)	1.53-0.006 <i>i</i>	1.53-0.0055 <i>i</i>	1.527-0.014 <i>i</i>	1.527-0.014 <i>i</i>	1.53-0.006 <i>i</i>	1.53-0.006i
550nm	Dust	1.52 – 0.0015i	1.53-0.0055 <i>i</i>	1.517-0.0011 <i>i</i>	1.564-0.002 <i>i</i>	1.564-0.002 <i>i</i>	1.53-0.002 <i>i</i>	1.53-0.008i
	SS	1.55 – 1e-7 <i>i</i>	1.50-1e-8 i	1.49-1e-8 <i>i</i>	1.45-0 <i>.i</i>	1.45-0 <i>.i</i>	1.38-4.26e-	1.50- 1e-8 <i>i</i>
		Aged BB: 1.54 –					9i	
		0.018 <i>i</i>						
Additional		NO ₃ -	-	-	NO ₃ -	NO ₃ -	-	-
Species f								
Dust Size		6 bins 0.0316-0.1-	8 bins 0.1-	Accum. mode:	5 bins 0.1-1-	4 bins	6 bins 0.1-	8 bins 0.1-
distribution		0.316-	0.18-0.3-0.6-	0.05 <rm<0.5< td=""><td>2-4-8-16</td><td>0-1-2-4-8</td><td>0.22-0.46-</td><td>0.18-0.3-</td></rm<0.5<>	2-4-8-16	0-1-2-4-8	0.22-0.46-	0.18-0.3-
(um) g		1.0-3.16-10-31.6	1.0-1.8-3.0-	coarse mode:			1.0-2.15-	0.6-1.0-1.8-
(r.)			6.0-10.0	r _m >0.5			4.64-10.0	3.0-6.0-10.0
References		Bellouin et al	Chin et al	Pozzoli et al.	Tsigaridis et	Bauer et al	Takemura	Colarco et
		2011	2002.2014	2011	al 2013	2008.2010	et al., 2005	al., 2010
			,		, = 0 ±0		2009	,

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^a Spatial resolutions (°latitude × °longitude × number of vertical levels).

^b Anthropogenic emission data are from either A2-ACCMIP or A2-MAP (refer to Diehl et al. 2012).

^c Biomass burning emission data (refer to Diehl et al. 2012).

192 ^d FF is fossil fuel and BB is biomass burning.

^eAs for EHCAM5-HAMMOZ model with a mixed aerosol scheme, the refractive index for each of the 7 modes is

194 calculated as the volume weighted average of the refractive indices of the components of the mode, including the 195 diagnosed aerosol water.

^f Additional aerosols besides commonly included aerosol species, i.e. SO_{4²⁻} (sulfate), Dust, SS (sea salt), BC (black
 carbon), and OA (organic aerosol). Here NO_{3⁻} is nitrate.

^g Listed is the edges of size bins in all models except for ECH, in which r_m is modal radii.

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Table 2. Summary of stations in South Asia used in this study

Туре	Station ^a	Lat	Lon	Alt (<i>m</i>)	Popul- ation ^b	Data Source ^c	Data Category	Main Feature
	Delhi	28.58° N	77.20° E	260	16.75	ICARB	BC	In western IGP, the largest city in India
	Karachi	24.87° N	67.03° E	49	13	AERONET	AOD AAOD	Coastal location in southern Pakistan
Urban	Lahore	31.54° N	74.32° E	270	9	AERONET	AOD AAOD	In western IGP, major agricultural region
	Hyderabad	17.48° N	78.40° E	545	6.81	ICARB	BC	In central Indian Peninsula
	Pune	18.52° N	73.85° E	559	5.05	ICARB	BC	In western plateau
	Kanpur	26.51° N	80.23 °E	123	2.77	AERONET/ ISRO-GBP	Misc. ^d	In central IGP
	Agra	27.06° N	78.03º E	169	1.75	ISRO-GBP	Misc. ^d	Between Delhi and Kanpur
	Allahabad	25.45° N	81.85º E	98	1.22	ISRO-GBP	Misc. ^d	In central-eastern IGP
	Kharagpur	22.52° N	87.52° E	28	0.37	ICARB	BC	In eastern IGP-outflow region to Bay of Bengal
Semi- Urban	Hisar	29.09° N	75.42° E	41	0.3	ISRO-GBP	Misc. ^d	Surrounded by agricultural field in western IGP
	Trivandrum	8.55° N	76.90° E	3	0.75	ICARB	BC	A coastal station in southern India
	Port Blair	11.63° N	92.70° E	60	0.1	ICARB	BC	Island in Bay of Bengal
Remote	Nainital	29.20° N	79.30° E	1950	0.04	ICARB	BC	High altitude remote location in the Himalayan foothills
	Minicoy	8.30° N	70.00° E	1	0.009	ICARB	BC	Island in Arabian Sea

^{a.} In decreasing order of the population ^{b.} Statistics in 2011 from wikipedia

211 212 ^{c.} Details in section 3.2 and 3.3

^{d.} Miscellaneous, including meteorological fields, AOD, AAOD and aerosol surface concentration.

Parameter	Unit	#	Mean	Median	Min	Max	Stdev	Diversity ^a
				SO4				
Emi ^b	$Tg(SO_2) vr^{-1}$	7	7.36	7.39	5.81	8.61	0.86	12%
Cheag	$Tg(SO_4) vr^{-1}$	4	0.27	0.28	0.15	0.35	0.10	36%
Chegd	$Tg(SO_4) vr^{-1}$	4	0.24	0.18	0.12	0.46	0.16	66% ^h
Wet	$Tg(SO_4) vr^{-1}$	7	4.38	3.98	3.64	6.21	0.93	21%
Drv	$Tg(SO_4) vr^{-1}$	7	0.78	0.77	0.27	1.26	0.35	44%
Dry/Dry+Wet	%	7	19	20	8	29	8	40%
Life time	Davs	7	5.02	4.81	3.22	8.50	1.73	34%
Load	Tg(SO ₄)	7	0.06	0.05	0.04	0.08	0.02	26%
MEE ^e	$m^2 g^{-1}(SO_4)$	4	8.56	8.99	5.58	10.68	2.15	25%
AOD	Unitless	4	0.07	0.07	0.04	0.08	0.02	27%
				BC				
Emi	Tg yr ⁻¹	7	0.70	0.71	0.59	0.78	0.06	9%
Wet	Tg vr ⁻¹	7	0.27	0.28	0.21	0.31	0.04	15%
Dry	Tg yr ⁻¹	7	0.15	0.19	0.05	0.21	0.07	46%
Dry/Dry+Wet	%	7	33	37	15	41	10	29%
Life time	Days	7	7.67	6.56	4.13	15.82	3.84	50%
Load	Tg	7	0.008	0.007	0.005	0.014	0.003	39%
MEE	$m^2 g^{-1}$	4	7.07	7.56	2.77	10.40	3.63	51%
AOD	Unitless	4	0.008	0.010	0.003	0.011	0.004	45%
				OA				
Emi ^f	Tg vr ⁻¹	7	3.69	3.58	2.77	4.46	0.61	16%
Wet	Tg vr ⁻¹	7	1.68	1.62	1.26	2.31	0.37	22%
Drv	Tg yr ⁻¹	7	0.78	0.82	0.31	1.21	0.32	41%
Drv/Drv+Wet	%	7	35	38	20	44	10	29%
Life time	Davs	7	5.60	5.25	4.44	7.09	1.07	19%
Load	Tg	7	0.05	0.04	0.03	0.07	0.01	25%
MEE	$m^2 g^{-1}$	4	5.17	4.99	3.69	7.00	1.39	27%
AOD	Unitless	4	0.023	0.022	0.018	0.030	0.005	21%
				DUST				
Emi	Tg vr ⁻¹	7	103.84	43.34	6.43	367.28	128.23	124%
Wet	Tg yr ⁻¹	7	43.43	41.07	11.82	92.55	24.47	56%
Drv + Sed ^g	Tg vr ⁻¹	7	98.50	46.92	1.34	316.87	113.42	115%
Drv/Drv+Wet	%	7	56	68	12	84	26	47%
Life time	Davs	7	3.86	4.17	1.08	6.92	1.98	51%
Load	Tg	7	0.87	0.91	0.16	1.43	0.39	45%
MEE	$m^2 g^{-1}$	4	0.64	0.59	0.50	0.89	0.17	27%
AOD	Unitless	4	0.10	0.09	0.08	0.12	0.02	22%
		-		TOTAL				/0
AOD	Unitless	7	0.21	0.18	0.16	0.33	0.06	28%
AAOD	Unitless	7	0.02	0.02	0.01	0.02	0.01	37%

Table 3. The statistics of the aerosol parameters over South Asia (60°E–95°E; 5°N–36°N. Land only) in 2006.

²¹⁹ ^a. The diversity is defined as the ratio of standard deviation and mean (i.e. stdev/mean). The largest and second

220 largest diversities in each species are highlighted in bold.

^b. The emission of SO₂, including anthropogenic and biomass burning emission.

222 c. The chemical production of SO_4 in aqueous phase reaction (i.e. SO_2 reacts with H_2O_2).

⁴. The chemical production of SO_4 in gaseous phase reaction (i.e. SO_2 reacts with OH).

^e. Mass extinction efficiency, defined as the ratio of AOD and load (i.e. AOD/load).

225 f. Sum of anthropogenic emission, biomass burning emissions and secondary organic aerosol.

226 g. Dry deposition plus sedimentation.

²²⁷ ^h. The top two largest diversities in each species are highlighted in bold.

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229	Figures
230	Captions
231	
232	Fig. 1. Topography of South Asia and the locations of the stations used in this study. Three
233	AERONET stations are labeled in blue, eight ICARB stations in red, and four ISRO-GBP stations in
234	black except for Kanpur. The topography map is obtained from
235	http://mapofasia.blogspot.com/2013/02/map-of-south-asia-area-pictures.html.
236	
237	Fig. 2. Spatial distribution of anthropogenic emissions of BC, OC, SO ₂ , NH ₃ and NO _x averaged for
238	2000-2007 from A2-ACCMIP emission dataset (units: g m ⁻² yr ⁻¹) over South Asia (60°E–95°E; 5°N
239	–36°N). The annual mean emission amount over South Asia (land only) is shown at the bottom.
240	
241	Fig. 3. Spatial distribution of biomass burning emission of BC based on GFED2 for each season
242	averaged for 2000-2007 (units: g C m ⁻² yr ⁻¹) over South Asia (60°E–95°E; 5°N –36°N). The
243	seasonal mean emission amount over South Asia (land only) is shown at the bottom. Note that the
244	color scale is the same as that of BC in the Fig. 2 for the purpose of comparison.
245	
246	Fig. 4. The annual averaged mean AOD for 2000-2007 over region: (a) South Asia (60°E–95°E; 5°N
247	–36°N, averaged over land only, i.e. the gray area in the map); (b) Central IGP (77°E-83°E; 25°N-
248	28°N, averaged over the red box on the map shown in Fig. 4a). Thin lines with symbols represent
249	seven models, and thick lines represent four satellite datasets. Multi-year averaged mean AOD and
250	the standard deviation is listed on each panel.
251	
252	Fig. 5. Monthly mean AOD (left column) and AAOD (right column) at three AERONET stations in
253	South Asia. The gray bar represents data from AERONET, the thin lines represent results from
254	seven models, and symbols represent the data from three satellite retrievals. On each panel, corr is
255	correlation coefficient of a model with AERONET, bias is relative mean bias, i.e. Σ
256	$(AOD_MODEL_i)/\Sigma$ $(AOD_AERONET_i)$, and rmse is root-mean-square error relative to AERONET.
257	
258	Fig. 6. Monthly AOD of total aerosol (aer) and components (ss, so ₄ , bc, oa, dust, no ₃ , soa and bb) at
259	Kanpur in 2004 from four models, HAD (upper left), GOC (upper right), GE5 (lower left), and SPR
260	(lower right). The gray bar represents total AOD from AERONET, and the lines represent the
261	model results of total AOD (black line) and component AODs (colored lines). The corresponding
262	annual mean values are also listed. NOTE: For the HAD model, bc and oa are only from fossil fuel
263	sources; the biomass burning aerosol is labeled "bb".
264	
265	Fig. 7a. Spatial distribution of AOD over South Asia in four seasons averaged for 2000–2007 from
266	four satellite datasets (MODIS-Terra, MODIS-Aqua, MISR, and SeaWiFS). The corresponding area
267	averaged seasonal mean AOD value over land is listed on each panel. Three AERONET stations
268	used in this study are labeled in the maps for references. Area in white means no retrieval
269	available due to the presence of bright surface or frequent cloud cover.
270	
271	Fig. 7b. Spatial distribution of AOD over South Asia in four seasons averaged for 2000–2007 from
272	seven models (the first three models using the anthropogenic emissions from A2-MAP and the rest
273	using A2-ACCMIP). The area averaged seasonal mean AOD value over land is listed on each panel.
274	Three AERONET stations used in this study are shown on the maps for references.
275	
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- Fig. 8. Seasonal mean of vertical profile of extinction coefficient (units: km⁻¹) at (a) Kanpur, and (b)
 Hyderabad from CALIOP and seven models. The corresponding seasonal mean AOD, Za (units: km)
 and F_{2km} are listed after each symbol name. The gray shaded area in CALIOP is one standard
 deviation relative to the average of 2006-2011.
- Fig. 9. Monthly mean surface BC concentration at eight ICARB stations in 2006 (units: μg m⁻³).
 Gray bar represents measurement from ICARB and thin lines represent seven models.
- Fig.10. Comparisons of seven models against ISRO-GBP campaign measurements at four IGP
 stations (Hisar, Agra, Kanpur, Allahabad from western to eastern IGP) in December 2004. The
 variables include meteorological fields of surface relative humidity (1st row) and surface
 temperature (2nd row), aerosol species mass concentrations of SO₄²⁻ (3rd row), NO₃⁻ (4th row),
 BC (5th row), and OA (6th row), and columnar AOD (7th row) and AAOD (8th row) at 550nm.
- Fig. 11. Mass extinction efficiency (MEE) at 550nm for individual aerosol components (units: m² g⁻
- ²⁹¹ ¹) as a function of relative humidity (RH). For SO_4^{2-} , OC and BC, MEE is calculated using the
- relationship of RH and size growth based on optical properties of aerosols and clouds (OPAC)
- 293 (Hess et al., 1998). For NO_3^- , MEE is calculated according to the work by A. Lacis
- 294 (http://gacp.giss.nasa.gov/data_sets/lacis/introduction.pdf).





Fig. 2.





0 0.20 0.30 0.40 0.50 1.0 1.5 2.0 5.0 10 20 105



Annual mean AOD (2000-2007)



321 Fig. 4.322







0 0.05 0.10 0.20 0.30 0.40 0.50 0.60 0.70 0.80 1.0 5.8





Fig.7b.

0.05 0.10 0.20 0.30 0.40 0.50 0.60 0.70 0.80 1.0 5.8

(a) Kanpur

(b) Hyderabad



351 Fig. 8.





