- Relating geostationary satellite measurements of aerosol optical
- 2 depth (AOD) over East Asia to fine particulate matter (PM_{2.5}):
- 3 insights from the KORUS-AQ aircraft campaign and GEOS-
- 4 Chem model simulations
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60 Abstract. Geostationary satellite measurements of aerosol optical depth (AOD) over East Asia from the GOCI and 61 AHI instruments can augment surface monitoring of fine particulate matter (PM2.5) air quality, but this requires 62 better understanding of the AOD-PM2.5 relationship. Here we use the GEOS-Chem chemical transport model to 63 analyze the critical variables determining the AOD-PM2.5 relationship over East Asia by simulation of observations 64 from satellite, aircraft, and ground-based datasets. This includes the detailed vertical aerosol profiling over South 65 Korea from the KORUS-AQ aircraft campaign (May-June 2016) with concurrent ground-based PM2.5 composition, 66 PM₁₀, and AERONET AOD measurements. The KORUS-AQ data show that 550 nm AOD is mainly contributed by 67 sulfate-nitrate-ammonium (SNA) and organic aerosols in the planetary boundary layer (PBL), despite large dust 68 concentrations in the free troposphere, reflecting the optically effective size and high hygroscopicity of the PBL 69 aerosols. We updated SNA and organic aerosol size distributions in GEOS-Chem to represent aerosol optical 70 properties over East Asia by using in-situ measurements of particle size distributions from KORUS-AQ. We find 71 that SNA and organic aerosols over East Asia have larger size (number median radius of 0.11 µm with geometric 72 standard deviation of 1.4) and 20% larger mass extinction efficiency as compared to aerosols over North America 73 (default setting in GEOS-Chem). Although GEOS-Chem is successful in reproducing the KORUS-AQ vertical 74 profiles of aerosol mass, its ability to link AOD to PM2.5 is limited by under-accounting of coarse PM and by a large 75 overestimate of nighttime PM2.5 nitrate. The GOCI/AHI AOD data over East Asia in different seasons show 76 agreement with AERONET AODs and a spatial distribution consistent with surface PM2.5 network data. The AOD 77 observations over North China show a summer maximum and winter minimum, opposite in phase to surface PM2.5. 78 This is due to low PBL depths compounded by high residential coal emissions in winter, and high relative humidity 79 (RH) in summer. Seasonality of AOD and PM2.5 over South Korea is much weaker, reflecting weaker variation of 80 PBL depth and lack of residential coal emissions.

1 Introduction

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82 PM_{2.5} (particulate matter with aerodynamic diameter less than 2.5 μm) in surface air is a severe public health 83 concern in East Asia, but surface monitoring networks are too sparse to thoroughly assess population exposure. 84 Satellite observations of aerosol optical depth (AOD) can provide a valuable complement (Van Donkelaar et al., 85 2015). Geostationary satellite sensors, including the Geostationary Ocean Color Imager (GOCI) launched by the 86 Korea Aerospace Research Institute (KARI) in 2011 (Choi et al., 2016, 2018, 2019) and the Advanced Himawari 87 Imager (AHI) launched by the Japanese Meteorological Agency (JMA) in 2014 (Lim et al., 2018, 2021), offer the 88 potential for high-density mapping of PM_{2.5} over East Asia (Chen et al., 2019; Wei et al., 2021a). However, more 89 confidence is needed in relating AOD to PM2.5. Here we evaluate the capability of the GEOS-Chem chemical 90 transport model (CTM) to simulate AOD-PM2.5 relationships over East Asia, exploiting in-situ aircraft 91 measurements of vertical aerosol profiles and optical properties from the joint NASA-NIER Korea - United States 92 Air Quality (KORUS-AQ) field study in May-June 2016 (Crawford et al., 2021; Peterson et al., 2019; Jordan et al., 93 2020) together with GOCI/AHI geostationary satellite data and surface measurement networks. This enables us to 94 identify critical variables and uncertainties for inferring PM2.5 from satellite AOD data.

96 The standard geophysical approach has been to use a CTM, such as GEOS-Chem, to compute the PM2.5/AOD ratio 97 (Liu et al., 2004; van Donkelaar et al., 2006; van Donkelaar et al., 2015; Xu et al., 2015; Geng et al., 2017), with 98 recent applications correcting for CTM biases using available PM2.5 surface network data (Brauer et al., 2016; Van 99 Donkelaar et al., 2016; van Donkelaar et al., 2019; Hammer et al., 2020). An alternative approach is to use artificial 100 intelligence algorithms to relate satellite AOD to PM2.5 by training on the surface network data (Hu et al., 2017; Deleted: machine-learning 101 Chen et al., 2018; Xiao et al., 2018; Wei et al., 2021a; Wei et al., 2021b; Pendergrass et al., 2021), and sometimes 102 including CTM values as predictors (Di et al., 2019; Xue et al., 2019). Yet another approach is to assimilate the 103 satellite-measured AODs in a CTM and correct in this manner the PM2.5 simulation, although this requires 104 attribution of model AOD errors to specific model parameters (Kumar et al., 2019; Saide et al., 2014; Sekiyama et 105 al., 2010; Cheng et al., 2019). In all of these approaches, a better physical understanding of the AOD-PM2.5 106 relationship as simulated by CTMs can greatly enhance the capability to infer PM2.5 from AOD data. 107 AOD measures aerosol extinction (scattering and absorption) integrated over the atmospheric column, so that its 108 relationship to 24-hr average surface PM2.5 (the standard air quality metric) depends on the aerosol vertical 109 distribution and optical properties, ambient relative humidity (RH), diurnal variation of PM2.5, and contribution from 110 coarse particulate matter to AOD, Airborne measurements of aerosol vertical profiles (without species information) Deleted: Little study of these factors has been conducted for East Asia. 111 in East Asia are limited (Zhang et al., 2006; Liu et al., 2009; Zhang et al., 2009; Sun et al., 2013; Li et al., 2017), and Deleted: very 112 speciated vertical profiles are rarer. AOD is highly sensitive to RH (Brock et al., 2016; Latimer and Martin et al., Deleted: that 113 2019; Saide et al., 2020), but the impact from RH uncertainty on AOD simulation lacks evaluation. In addition, Deleted: of aerosol species 114 because the AOD is a daytime measurement that needs to be related to 24-h average PM_{2.5}, the diurnal variation of Deleted: even rare 115 PM_{2.5} needs to be understood (Guo et al., 2017; Lennartson et al., 2018). Finally, although there are studies on the 116 optical depth of coarse mode desert dust (Eck et al., 2010; Ridley et al., 2016), there has been to our knowledge no 117 study of how coarse anthropogenic PM may contribute to the AOD measurements. Coarse anthropogenic PM 118 (distinct from desert dust) is known to be high over East Asia (Chen et al., 2015; Dai et al., 2018). 119 2 Data and methods 120 2.1 Observations 121 We use observations over China and South Korea from multiple platforms including surface sites, aircraft, and 122 satellites (Table 1 and 2). Surface data (Table 1) include PM2.5 from national observation networks in China (Zhai et 123 al., 2019) and South Korea (Jordan et al., 2020), speciated PM_{2.5} at 7 supersites in South Korea during KORUS-AQ 124 (Choi et al., 2019), and ground-based AODs from the AERONET network at 5 sites in East, China and 10 sites in Deleted: North

South Korea (21 sites during KORUS-AQ). We use total and fine-mode AODs at 500 nm wavelength from the

Ångström Exponents at 500 nm for consistency with the satellite AOD data.

AERONET Version 3; Spectral Deconvolution Algorithm (SDA) Version 4.1 Retrieval Level 2.0 database (Giles et

al., 2019; O'Neill et al., 2003). The AERONET AODs at 500 nm are converted to 550 nm using total and fine mode

A number of past studies have used satellite AOD data to infer surface PM2.5 using physical and statistical models.

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137 Table 1. Surface site observations used in this work (2016)

|142 |143 |144 |145 |146 |147 |150 |151 |152 |153 |154 |155 |156

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Variable	Number of sites		
PM _{2.5} in East China ^a	<u>598</u>	Deleted: North	
PM _{2.5} in South Korea ^b	130	Deleted: 117	
PM _{2.5} composition in South Korea (May-June 2016) ^c	7		
AERONET total and fine mode AOD in East China d	5	Deleted: North	
AERONET total and fine mode AOD in South Korea ^d	10-21 °		
^a Hourly PM _{2.5} from the China National Environmental Mon	itoring Centre (CNEMC; quotsoft.net/air/) in East	Deleted: North	
China, including only sites with more than 90% data coverage CNEMC dataset is described in our previous study (Zhai et a reference $RH \le 35\%$.		Deleted: (115.5-122° E, 34.5-40.5° N)	
b Hourly PM _{2.5} from the AirKorea network (airkorea.or.kr), v The PM _{2.5} measurements are made at reference RH \leq 35%.	with the same data selection criteria as for East China.	Deleted: North	
^c Major PM _{2.5} components including sulfate, nitrate, ammoni South Korea during KORUS-AQ (May-June 2016; Choi et al converted to that of organic aerosol with a multiplicative fact al., 2018).	1., 2019). The mass concentration of organic carbon is		
$^{\rm d}$ AODs are from the AERONET Version 3 Level 2.0 all-poin at the XuZhou site in East China are from the Version 3 Level converted to 550 nm (AOD _{550nm}) using Ångström Exponent a AOD _{500nm} ($^{550}_{500}$)-AE _{500nm} .	el 1.5 database. AOD at 500 nm (AOD _{500nm}) is	Deleted: North	
$^{\rm c}{\rm AERONET}$ AODs in South Korea are from 10 sites for the	full year of 2016 and 21 sites during KORUS-AQ.		
The KORUS-AQ campaign (Table 2) includes 20 flights over	r the Korean peninsula and the surrounding ocean from		
May 2 to June 10, 2016, with vertical profiling up to 8 km al	titude. We use the aircraft observations of remote and in		
situ aerosol extinction (scattering + absorption) coefficients,	dry aerosol number size distributions, sub-micron non-		
refractory aerosol composition, bulk aerosol ionic composition	on, black carbon (BC), and relative humidity (RH).		
Geostationary satellite AOD at 550 nm are retrieved by the YGOCI (Choi et al., 2016, 2018) and AHI (Lim et al. 2018) in	struments, with GOCI covering East China and South		
Korea and AHI covering the broad East Asia region. AOD fr	*		
resolution and 2.5-minute (AHI) to 1-hour (GOCI) temporal		Deleted: to 2.5-minute (AHI)	
time). We use the fused AOD product generated from the Yo	nsei GOCI and AHI AOD retrievals, each using two		
different surface reflectance methods (Lim et al., 2021). Fusi	on of this four-member ensemble is done by the		
maximum likelihood estimate (MLE) method, with weighting	g and averaging based on errors determined by		
comparison to AERONET AOD. The fused satellite AOD pr	oduct is shown by Lim et al. (2021) to have higher		

accuracy than its member products in comparison with AERONET data during the KORUS-AQ campaign. We will

refer to it as the 'GEO satellite AOD' product in what follows.

Table 2. KORUS-AQ aircraft observations used in this work (May-June 2016).

Variable	Instrument
Aerosol extinction profile at 532 nm	HSRL ^a
Aerosol scattering coefficient at 550 nm	TSI nephelometers ^b
Aerosol absorption coefficient at 532 nm	PSAPs ^c
Aerosol dry size distribution	TSI LAS ^d
Bulk aerosol ionic composition	SAGA ^e
Sub-micron non-refractory aerosol composition	HR-ToF-AMS ^f
Black carbon concentration	HDSP2 ^g
Relative humidity	DLH h

- 176 a NASA Langley airborne High Spectral Resolution Lidar (HSRL) (Hair et al., 2008; Scarino et al., 2014).
- 177 b NASA Langley TSI-3563 nephelometers (Ziemba et al., 2013).
- 178 c Radiance Research 3-wavelength particle soot absorption photometers (PSAPs; Ziemba et al., 2013).
- 179 definition in Institute and Institute a
- 180 (LAS) Model 3340.

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- The cutoff aerodynamic diameter of the inlet is around 4 μm , corresponding to a geometric particle diameter of 2.5
- $183~\mu m$ (McNaughton et al., 2007; McNaughton et al., 2009).
- $184 \qquad {}^{\rm f} \, \text{University of Colorado Boulder High-Resolution Time-of-Flight Aerosol Mass Spectrometer (HR-ToF-AMS;}$
- 185 DeCarlo et al., 2006; Nault et al., 2018; Guo et al., 2020).
- 186 g NOAA Humidified-Dual-Single-Particle Soot Photometer (HDSP2; Lamb et al., 2018).
- 187 h NASA Diode Laser Hygrometer (DLH; Podolske et al., 2003).

2.2 GEOS-Chem simulation

- We use GEOS-Chem version 12.7.1 (DOI: 10.5281/zenodo.3676008) in a nested-grid simulation at a horizontal
- resolution of $0.5^{\circ} \times 0.625^{\circ}$ over East Asia (100-145 °E, 20-50 °N). GEOS-Chem simulates detailed tropospheric
- 191 oxidant-aerosol chemistry and is driven here by GEOS-FP assimilated meteorological data from the NASA Global
- 192 Modeling and Assimilation Office (GMAO). Boundary layer mixing uses the non-local scheme implemented by Lin
- 193 and McElroy (2010). Dry deposition of gases and particles follows a standard resistance-in-series scheme (Zhang et
- 194 al., 2001; Fairlie et al., 2007; Fisher et al., 2011; Jaeglé et al., 2018). Wet deposition of gases and particles includes
- 195 contributions from rainout, washout, and scavenging in convective updrafts (Liu et al., 2001; Amos et al., 2012; Q.
- Wang et al., 2011; Q. Wang et al., 2014) with recent updates by Luo et al. (2019, 2020). We use pre-archived initial

197 conditions from Zhai et al. (2021) and run the model from December 1, 2015 to December 31, 2016. The first month 198 is used for spin-up and the year 2016 is used for analysis.

GEOS-Chem has been used extensively to simulate PM2.5 and its composition in East Asia (Geng et al., 2017; Li et al., 2016; Choi et al., 2019; Jeong et al., 2008; Park et al., 2021; Zhai et al., 2021). Here we use the bulk representation of aerosols including sulfate (Park et al., 2004; Alexander et al., 2009), nitrate (Jaeglé et al., 2018),

primary and secondary organics (Pai et al., 2020), BC (Q. Wang et al., 2014), natural dust in four advected size

203 ranges (Fairlie et al., 2007), anthropogenic fine dust (Philip et al., 2017), and sea salt in two size ranges (Jaeglé et

al., 2011). Heterogeneous sulfate formation on aqueous aerosols is represented by a simplified parameterization

scheme (Y. Wang et al., 2014), where the SO₂ uptake coefficient (γ) linearly increases from 1 × 10⁻⁵ at RH \leq 50% to 2 × 10⁻⁵ at RH = 100%. The thermodynamic equilibrium of sulfate-nitrate-ammonium (SNA) aerosols with the gas

phase is computed with ISORROPIA II (Fountoukis and Nenes, 2007; Pye et al., 2009) assuming an aqueous

aerosol. We include reactive uptake on dust of acid gases (HNO₃, SO₂, and H₂SO₄), limited by consumption of dust

alkalinity (Fairlie et al., 2010). The alkalinity of emitted dust is estimated by assuming 7.1% Ca²⁺ and 1.1% Mg²⁺ as

alkaline cations by dust mass (Shah et al., 2020).

- 211 Monthly anthropogenic emissions are from the Multi-resolution Emission Inventory in 2016 for China (MEIC;
- Zheng et al., 2018; http://meicmodel.org) and from the KORUSv5 emission inventory at base year 2015 (Woo et al.,
- 213 2020; http://aisl.konkuk.ac.kr/#/emission_data/korus-aq_emissions) for other Asian countries and shipping
- 214 emissions. MEIC over China applies weekly and diurnal scaling factors for all anthropogenic emissions (Zheng et
- al., 2018). The KORUSv5 agricultural NH3 emissions apply the diurnal scaling factors from MEIC. Natural
- emissions include NO_x from lightning (Murray et al., 2012) and soil (Hudman et al., 2012), MEGANv2 biogenic
- volatile organic compounds (VOCs) (Guenther et al., 2012), dust (Meng et al., 2020), and sea salt (Jaeglé et al.,
- 218 2011). Open fire emissions are from the Global Fire Emissions Database version 4 (GFED4; van der Werf et al.,
- 219 2017).

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2.3 AOD simulation

- AOD in GEOS-Chem is diagnosed by integrating vertically the aerosol scattering and absorption coefficients
- 222 obtained with a standard Mie calculation applied to assumed size distributions, hygroscopicity, refractive indices,
- and densities for individual aerosol components, and summing over all components (Martin et al., 2003). Optical
- properties are listed in Table 3. Sulfate, nitrate, and ammonium share the same optical properties and are lumped as
- an SNA aerosol component for the purpose of optical calculations. All aerosol components except dust are assumed
- 227 and 4.0 µm) for optical calculations, with the smallest four bins partitioned by mass from the first advected dust bin
- $228 \qquad (<2.5~\mu m~in~geometric~diameter)~following~L.~Zhang~et~al.~(2013).~Dust~particles~follow~a~gamma~size~distribution$
- 229 within their optical size bins (Curci, 2012). The BC absorption enhancement from coating is as given by X. Wang et
- 230 al. (2014).

231 Our initial simulations indicated that aerosol extinction coefficients from the standard GEOS-Chem version 12.7.1 232 underestimated in situ measured extinction coefficients during KORUS-AQ by 20% on average (Figure S1). We 233 traced this problem to bias in the assumed size distributions for SNA and organic aerosol, as shown in Section 3. 234 Therefore, we re-computed the diagnostic AOD using updated log-normal size distributions for SNA and organic 235 aerosol with number median radius $R_{N,med} = 0.11~\mu m$ and geometric standard deviation $\sigma = 1.4$ based on KORUS-236 AQ observations, instead of $R_{N,med} = 0.058 \, \mu \text{m}$ and $\sigma = 1.6$ in the standard model version 12.7.1, which is derived 237 from IMPROVE network measurements of aerosol mass scattering efficiency over North America (Latimer and 238 Martin, 2019).

239 Table 3. Aerosol optical properties a.

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Aerosol component	R _{N,med} , μm	σ	Hygroscopicity b	Refractive index	ρ , g cm ⁻³
SNA °	0.11	1.4	$\kappa = 0.61$	$1.53 - 6.0 \times 10^{-3}i$	1.7
Organic ^c	0.11	1.4	$\kappa = 0.1$	$1.53 - 6.0 \times 10^{-3}i$	1.3
BC	0.020	1.6	GADS	$1.75 - 4.4 \times 10^{-3}i$	1.8
Sea salt (fine)	0.085	1.5	GADS	$1.5 - 1.0 \times 10^{-3}i$	2.2
Sea salt (coarse)	0.40	1.8	GADS	$1.5 - 1.0 \times 10^{-3}i$	2.2
Dust	7 size bins	NA	$\kappa = 0$ d	$1.558 - 1.4 \times 10^{-3}i$	2.5-2.65 e

240 ^a Aerosol optical properties used in this work for computing aerosol scattering and absorption coefficients. Values 241 are from the standard GEOS-Chem model version 12.7.1, except for the size distributions of SNA and organic 242 aerosol which are based on KORUS-AQ observations (see text). All aerosol components except dust have log-243 normal dry size distributions where $R_{N,med}$ is the number median radius and σ is the geometric standard deviation. 244 Refractive indices are for 550 nm wavelength. ρ is the dry aerosol mass density.

^b Hygroscopic growth for SNA and organic aerosol as a function of relative humidity (RH, %) is computed from κ -Kohler theory as a diameter growth factor GF = $(1 + \kappa * RH/(100-RH))^{1/3}$ (Latimer and Martin, 2019). Hygroscopic growth factors for other aerosol components are from the Global Aerosol Data Set (GADS) as tabulated in Chin et al. (2002) and Martin et al. (2003).

- 249 ^c R_{N,med} and σ are fit to KORUS-AQ observations as described in the text. Standard GEOS-Chem v12.7.1 assumes 250 $R_{N,med} = 0.058 \mu m$, $\sigma = 1.6$ (Latimer and Martin, 2019).
- 251 ^d Hygroscopic growth of dust particles is assumed negligible.
- 252 e Sub-micron dust particles have a density of 2.5 g cm⁻³ while coarse mode dust particles have a density of 2.65 g 253 cm⁻³. Dust size distribution is described in the text.

Aerosol concentrations and optical properties during KORUS-AQ 3

Here we use the KORUS-AQ aircraft observations and their simulation with GEOS-Chem to better understand the vertical distributions of different aerosol components contributing to AOD over South Korea. We begin with the mean vertical profile of aerosol mass and go on to examine the aerosol optical properties. This provides the basis for analyzing the observed vertical profile of aerosol extinction, its simulation by GEOS-Chem, and the consistency with GEO satellite and AERONET AOD measurements over South Korea during the KORUS-AQ period.

3.1 Vertical profile of aerosol mass

Figure 1 shows the mean aircraft vertical profiles of aerosol mass observed during KORUS-AQ and their simulation by GEOS-Chem. The KORUS-AQ aircraft sampled during the daytime, mainly between 9 am and 3 pm local time. Here and elsewhere, the model is sampled along the flight tracks and at the flight times. The observed vertical distribution of aerosol mass concentrations (Figure 1a) shows that 58% of column aerosol mass is below 2 km altitude, which we define as the average planetary boundary layer (PBL) during KORUS-AQ, and 34% is at 2-5 km altitude, which we define as the lower free troposphere (FT). The model has a similar vertical distribution (Figure 1b), with 57% of aerosol mass in the PBL and 36% in the lower FT. SNA, organic, and dust each contribute about a third of aerosol mass in the PBL while dust dominates in the lower FT both in the observations and in the model. The enhanced dust in the lower FT is driven by a few dust events, which the model reproduces (Figure S2). Black carbon and sea salt (not shown) make only minor contributions to aerosol mass. The model underestimates sulfate by 28% in the PBL, which leads to a 20% overestimate of nitrate, with canceling effect on the SNA mass simulation.

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The GEOS-Chem simulation of organic aerosol in this work uses the simple scheme of Pai et al. (2020) and underestimates aircraft observations by 16% in the PBL. Over 90% of GEOS-Chem organic aerosol is secondary, consistent with observations (Figure S4; Nault et al., 2018; Pai et al., 2020). GEOS-Chem simulation of the KORUS-AQ aerosol component profiles for different meteorological regimes is presented in Park et al. (2021).

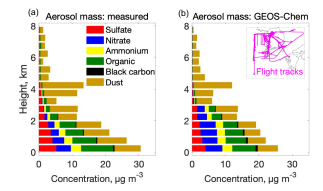


Figure 1. Vertical profiles of aerosol mass during KORUS-AQ. Panel (a) shows the mean vertical distributions of observed mass concentrations of major aerosol components at ambient temperature and pressure. Panel (b) is the same as (a) but from the GEOS-Chem model sampled along the flight tracks (inset). We derive dust concentration from SAGA Ca^{2+} and Na^+ following Shah et al. (2020) by assuming that non-sea salt Ca^{2+} accounts for 7.1% of dust mass: [dust] = $([Ca^{2+}] - 0.0439 \ [Na^+]/2) / 0.071$ where the brackets denote mass concentration. Modeled dust is shown for particles with geometric diameter < 2.5 μ m, to be consistent with SAGA measurements (Table 2 footnote e). Measured sulfate, nitrate, ammonium, and organic aerosol concentrations are from the AMS instrument (values from the SAGA instrument are shown in Figure S4). All data are averaged over 500-m vertical bins. Here and elsewhere, we excluded pollution plumes diagnosed by either NO_2 or $SO_2 > 10$ ppbv (3.4% of all the data).

3.2 Aerosol size distributions

Figure 2a shows the normalized dry aerosol number size distributions on each of the 20 flights and in 3 altitude bands: < 1.5 km, 3-5 km, and 6-7 km (60 lines). The spread in the size distributions above 1 μ m in diameter reflects dust influence. We select measurements below 1.5 km altitude when SNA + organic aerosol mass concentrations are more than 4 times that of dust as defining the SNA + organic aerosol size distributions (green lines in Figure 2a). Conditions dominated by SNA + organic aerosols define the lower envelopes of the ensemble of size distributions at diameter > 1 μ m. SNA and organics were observed to have similar size distributions during KORUS-AQ (Kim et al., 2018).

Figure 2b converts the SNA + organic dominated number size distributions to volume size distributions. The observed SNA + organic dominated aerosol size distribution is shifted toward larger sizes relative to the standard GEOS-Chem. The secondary maximum in the coarse mode could be due to dust. We fitted the observed SNA + organic aerosol size distributions to a lognormal distribution with volume median radius $R_{V,med} = 0.15 \mu m$ and geometric standard deviation $\sigma = 1.4$. The number median radius is derived from the volume median radius following Seinfeld and Pandis (2016):

$$\ln R_{N,med} = \ln R_{V,med} - 3\ln^2 \sigma \tag{1}$$

which yields $R_{N,med} = 0.11~\mu m$. In comparison, the standard GEOS-Chem size distribution from Latimer and Martin (2019) has $R_{N,med} = 0.058~\mu m$ and $\sigma = 1.6$. We adopt the observed log-normal size distribution parameters in what follows (Table 3).

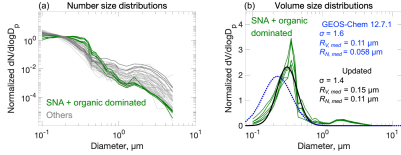


Figure 2. Aerosol dry size distributions measured in the KORUS-AQ aircraft campaign. Panel (a) shows mean normalized number size distributions measured on each of the 20 flights and for 3 altitude bins: < 1.5 km, 3-5 km, and 6-7 km (60 lines total). The SNA + organic dominated size distribution profiles are highlighted in color. Panel (b) shows normalized volume size distributions for conditions dominated by SNA + organic aerosols (green lines), along with a least-square fit to a lognormal distribution (black line), and the standard GEOS-Chem v12.7.1 size distribution from Latimer and Martin (2019) (blue dashed line). Normalization imposes an arbitrary value of unit area below each line. Lognormal

distribution parameters are inset in panel (b) including volume median radius ($R_{V,med}$), number median radius ($R_{N,med}$), and geometric standard deviation (σ). 3.3 Aerosol extinction and relation to AOD Figure 3 shows the vertical profiles of ambient aerosol extinction coefficients and RH during KORUS-AQ. Vertical profiles of aerosol extinction were measured on the aircraft both remotely with the HSRL instrument (above and below the aircraft) and in situ with TSI-3563 nephelometers (for scattering) and PSAPs (for absorption). The two agree well, as shown in Figure 3a. They indicate that 76-90% of column aerosol extinction is in the PBL at 0-2 km altitude and 9-19% is in the lower FT at 2-5 km. Both measurements show that aerosol extinction is much more strongly weighted to the PBL than aerosol mass (Figure 1). Also shown in Figure 3a are the contributions of individual aerosol components to the extinction profile, as computed from the GEOS-Chem optical properties (Table 3) applied to the observed mass concentrations. The sum shows a good match to the measured extinction coefficient profiles. The much larger contribution of the PBL to column aerosol extinction than to column mass is because aerosol mass in the lower FT is mainly composed of dust, whose mass extinction efficiency is much smaller than SNA and organics due to its coarse size and lack of hygroscopic growth (Figure S5). The mean AOD inferred from the aircraft data is 0.36 and is contributed 59% by SNA, 27% by organic aerosol, 12% by dust, and 2% by BC. It is consistent with the mean AODs measured at AERONET stations in South Korea during KORUS-AQ (Figure S6). Figure 3b shows the GEOS-Chem simulation of aerosol extinction profiles for comparison to the observations in Figure 3a. The model underestimates extinction coefficients by 20% below 1 km altitude, leading to a 10%underestimate of aircraft inferred AOD, although there is no such underestimate in aerosol mass. This is caused by a negative RH bias in the GEOS-FP meteorological data used to drive GEOS-Chem, particularly at high RH conditions (Figure 3c) and is corrected if we apply the observed RH rather than the GEOS-FP RH to the GEOS-

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Chem aerosol mass concentrations (Figure 3d).

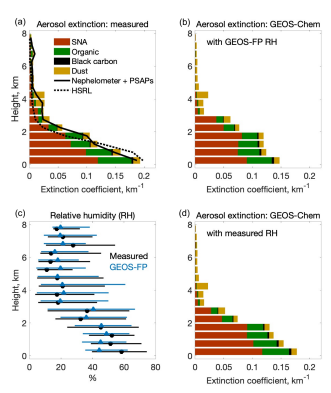


Figure 3. Vertical profiles of aerosol extinction coefficients and relative humidity (RH) during KORUS-AQ. Panel (a) shows the mean observed vertical distributions of 550 nm extinction coefficients measured in situ (nephelometer + PSAPs; at ambient RH) and remotely (HSRL), along with an independent calculation (colored horizontal bars) from the measured mass concentrations of major aerosol components, measured RH, and GEOS-Chem optical properties as given in Table 3. Panel (b) shows the mean aerosol extinction profile in GEOS-Chem and the contributions from the different model components. Panel (c) is the median vertical profile of RH (horizontal bars are 25-75th percentiles) from aircraft measurements and the GEOS-FP assimilated meteorological data used to drive GEOS-Chem. Panel (d) is the same as (b) but calculated using measured RH.

4 AOD and surface particulate matter over South Korea during KORUS-AQ

Our analysis of Section 3 used the KORUS-AQ aircraft data together with GEOS-Chem to attribute AOD over South Korea to individual aerosol components and altitudes. We now take the next step of evaluating the capability of GEOS-Chem to independently simulate observed AODs and surface particulate matter concentrations.

351 during the KORUS-AQ period with AERONET total AOD added as circles. The GEO satellite AOD shows high 352 values (0.5-0.6) along the west coast of South Korea, significantly correlated with AERONET total AOD with a 353 spatial correlation coefficient (R) of 0.7. GEO satellite AOD is biased low at sites in the Seoul Metropolitan Area 354 (SMA) and is biased high on the Yellow Sea islands, resulting in an overall -10% bias. The low biases in the SMA 355 could be due to high-concentration aerosol pixels mis-identified as clouds and/or possible issues with the aerosol 356 type assumption in the aerosol retrieval, while the high biases on the Yellow Sea islands could result from 357 uncertainties in the assumption of ocean surface reflectance, as has been discussed by Choi et al. (2016, 2018) and 358 Lim et al. (2018, 2021). Sampling the AODs at or near the seven PM2.5 supersites operating during KORUS-AQ 359 shows no significant bias (inset values in Figure 4a). 360 Figure 4b-e shows the spatial distributions of GEOS-Chem AOD, surface PM₁₀ (particulate matter with aerodynamic 361 diameter less than 10 µm), surface PM_{2.5}, and surface coarse PM (PM₁₀ minus PM_{2.5}; particulate matter with 362 aerodynamic diameter less than 10 µm and larger than 2.5 µm), with surface observations shown as circles and 363 median values at the measurement sites inset. GEOS-Chem reproduces the satellite AOD enhancements along the 364 west coast of South Korea but the values are lower than observed, which we attribute to unaccounted coarse PM and 365 negative RH bias as discussed below. Comparison of AERONET total and fine mode AOD shows a 13% 366 contribution of coarse particles to total AOD. Comparison of GEOS-Chem to the fine-mode AERONET AOD, as 367 shown in Figure 4b, finds a 15% underestimate that could be attributed to the low-RH bias (Figure 3c). Concurrent 368 measurements of PM₁₀ and PM_{2.5} at AirKorea sites show that coarse PM (median 21 μg m⁻³) accounts for 41% of total PM₁₀ (50 µg m⁻³), while coarse PM in GEOS-Chem is much lower (3.5 µg m⁻³; Figure 4e). Therefore, about 369 Deleted: 1 370 half of the GEOS-Chem underestimate of total AOD can be attributed to missing coarse PM, with the other half Deleted: 4 371 comes from negative RH bias. Coarse PM has a concentration larger than 10 µg m⁻³ across South Korea, with higher 372 concentration in the SMA ($\sim 30~\mu g~m^{-3}$) than in rural areas ($\sim 15~\mu g~m^{-3}$), implying an origin from both 373 anthropogenic and natural sources (Figure 4e). 374 GEOS-Chem overestimates surface PM2.5 by 43% over South Korea (Figure 4d), in contrast to the simulation of 375 AERONET fine mode AOD (Figure 4b). Figure 4f-j shows the spatial distributions of major PM2.5 components in 376 GEOS-Chem (background) and measurements (filled circles). GEOS-Chem is not significantly biased relative to the 377 observations for organic aerosol and BC, and underestimates sulfate by 22%. We find that the model bias for PM2.5 378 is largely driven by nitrate, which is overestimated by a factor of 3 and leads to a 56% overestimate of ammonium. 379 By contrast, comparison to the KORUS-AQ data below 1-km altitude showed only a 20% overestimate of nitrate 380 (Figure 1). This is because the model bias is mainly driven by nighttime conditions (Figure 5), while aircraft 381 samples in the daytime during KORUS-AQ. The cause of this large model bias is analyzed by K. R. Travis et al. Deleted: as shown in Figure 5 382 (manuscript in preparation) and is attributed to nighttime nitrate chemistry and deposition in the stratified boundary 383 layer.

Figure 4a shows the spatial distribution of the fused geostationary satellite (GOCI/AHI) AOD (GEO satellite AOD)

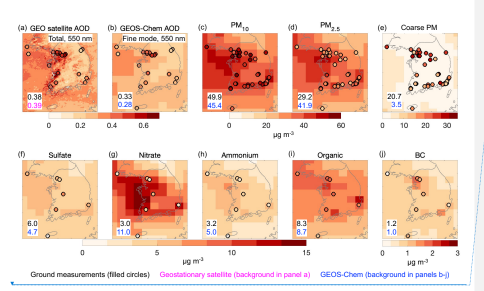
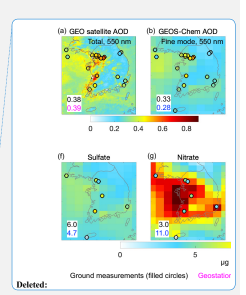


Figure 4. Spatial distributions of AOD and surface PM₁₀, PM_{2.5}, coarse PM (PM₁₀ minus PM_{2.5}), and major PM_{2.5} components over South Korea averaged during KORUS-AQ (May 9 - June 10, 2016). Panel (a) shows the fused geostationary (GEO) 550 nm AOD from the GOCI and AHI satellites (background) and AERONET 550 nm total AOD (filled circles). Panel (b) shows GEOS-Chem 550 nm AOD sampled at hourly GEO satellite AOD (GEOS-Chem clear-sky AOD; background) and AERONET 550 nm fine mode AOD (filled circles). Panel (c) shows surface PM₁₀ modelled by GEOS-Chem (background) and measured at ground sites (filled circles). Panels (d-j) are the same as panel (c) but respectively for PM_{2.5}, coarse PM (PM₁₀ minus PM_{2.5}), and sulfate, nitrate, ammonium, organic, and BC PM_{2.5} components. Values inset are median values from ground-based measurements (black) and sampled from GEO satellite (magenta) and GEOS-Chem (blue). Measured PM₁₀, PM_{2.5}, and coarse PM in panels (c-e) are shown for a random selection of 50% of AirKorea sites to visualize spatial distribution, and inset values are for the seven supersites where PM_{2.5} composition was measured. Median AOD values inset are sampled at or near the seven supersites to avoid biasing by the large number of sites in the Seoul Metropolitan Area. Modelled total PM_{2.5} concentrations are calculated at 35% RH (Table 3). Modelled PM₁₀ is the sum of PM_{2.5}, coarse dust, and coarse sea salt.



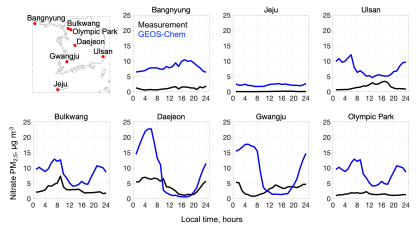


Figure 5. Median diurnal variations of $PM_{2.5}$ nitrate concentrations at the seven supersites (top left panel) operated in South Korea during KORUS-AQ (May 9 - June 10, 2016). Values are medians binned by hour. GEOS-Chem model values are sampled to coincide with the measurements.

5 AOD and its relationship to PM_{2.5} over East Asia

We build on our analysis of the KORUS-AQ period for a broader interpretation of the distribution of AOD over Korea and China and its relationship to surface PM_{2.5}, acknowledging that the conditions sampled in KORUS-AQ may not be representative of other seasons or of China. Figure 6 shows the spatial distributions of 2016 annual and seasonal mean geostationary (GEO) satellite AODs, the corresponding GEOS-Chem clear-sky AODs, and GEOS-Chem surface PM_{2.5}. The Figure gives normalized mean biases (*NMB*s) relative to ground-based measurements from AERONET and from the PM_{2.5} surface networks (shown as circles) over the North China region (115.5-122° E, 34.5-40.5° N) and South Korea. The North China region is defined to overlap with the domain of the geostationary satellite AOD, and to ensure consistent seasonal variations within its narrow latitude.

On an annual mean basis, AOD over North China (~ 0.5 -0.6) is about 50% larger than over South Korea (~ 0.3 -0.4). AOD over South Korea shows higher values (> 0.4) in the Seoul Metropolitan Area, consistent with that during the KORUS-AQ period (Figure 4a). Transport from the Asian continent is strongest in spring when the frequency of cold front passages is highest (Liu et al., 2003). AERONET total AOD in spring (0.4-0.6) is twice as large as fine-mode AOD (0.2-0.3), reflecting a large contribution of dust. In seasons other than spring, 80-90% of AERONET total AOD is contributed by the fine mode. There is large seasonality in AODs over North China, and weaker seasonality over South Korea, which will be discussed below.

The GEOS-Chem clear-sky AODs show the same spatial and seasonal patterns as GEO satellite AODs but tend to be low in spring and summer. Comparison of the model to AERONET AODs confirms this bias and shows better

agreement with fine-mode AOD in spring (*NMB* of -1%), implying an underestimate of coarse dust that is consistent with our comparisons to the AirKorea network data (Figure 4e). Comparison of clear-sky and all-sky AODs in GEOS-Chem shows no significant difference on an annual and seasonal mean basis, except for winter (Figure S7). Winter has larger all-sky AOD than clear-sky AOD and the lowest rate of successful satellite retrievals (Figure S7), which may be due in part to misclassification of heavy wintertime PM_{2.5} pollution as clouds (Zhang et al., 2020).

The spatial distributions of PM_{2.5} in GEOS-Chem in different seasons match closely the observations (Figure 6, bottom row). We see also a close coincidence between the spatial distributions of PM_{2.5} and AODs, both in the observations and the model. On an annual mean basis, GEOS-Chem overestimates PM_{2.5} by 16% in North China and by 14% in South Korea, even though it underestimates AERONET fine mode AODs by 15%. The overestimate of PM_{2.5} in South Korea is worst in spring (27%), consistent with KORUS-AQ results which we previously attributed to excessive nighttime nitrate build-up in the model. Over North China, the overestimate of PM_{2.5} is worst in summer (33%), consistent with the nitrate overestimate in summer shown in our previous study (Zhai et al., 2021), which could also be due to model overestimate of nighttime nitrate (Miao et al., 2020).

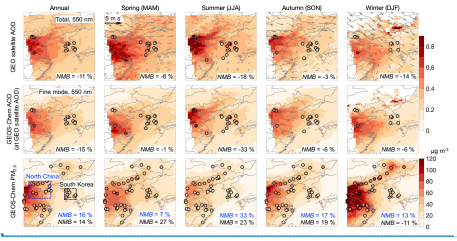
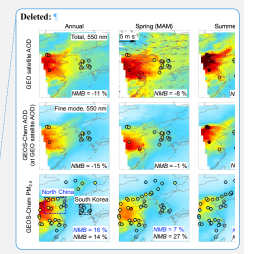


Figure 6. Spatial distributions of 2016 annual and seasonal mean AOD (550 nm) and surface $PM_{2.5}$. The top row shows the observed GOCI/AHI geostationary satellite AOD (GEO satellite AOD) on the GEOS-Chem $0.5^{\circ} \times 0.625^{\circ}$ grid with superimposed 925 hPa GEOS-FP wind fields and AERONET total AODs (circles). The middle row shows clear-sky GEOS-Chem AOD, with AERONET fine mode AOD added as circles. The bottom row shows GEOS-Chem surface $PM_{2.5}$ (background) with surface network measurements (circles). AERONET AODs are shown only when more than 10 months of data are available for the annual mean and all 3 months data are available for each season. The $PM_{2.5}$ observations shown are for a random selection of 7% of network sites for visual clarity. GEOS-Chem $PM_{2.5}$ is calculated

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at 35% RH (Table 3). Normalized mean biases (NMBs) inset are for the comparisons of GEO satellite and GEOS-Chem values to the corresponding ground measurements.

Figure 7 shows daily correlations of the regional average series between AERONET total AOD and GEO satellite AOD, between AERONET fine mode AOD and GEOS-Chem AOD, as well as between measured PM2.5 and GEOS-Chem PM2.5. Correlations in Figure 7 are all statistically significant with correlation coefficients (R) ranging from around 0.7 to more than 0.9 and normalized mean biases (NMB) within \pm 30%. The correlations of these three pairs are similar over South Korea and North China, except that GEOS-Chem overestimates springtime PM2.5 in South Korea but not over North China, possibly due to a model overestimate of the long-range transport of PM2.5 from China to South Korea in spring.

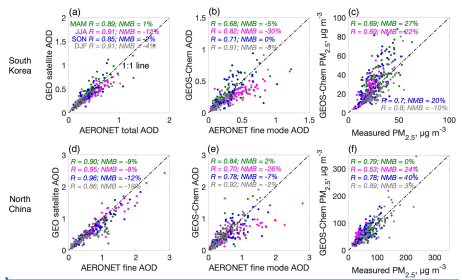
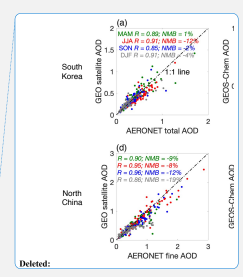


Figure 7. Scatter plots of regional mean daily (a and d) GEO satellite AOD vs. AERONET total AOD, (b and e) GEOS-Chem AOD vs. AERONET fine-model AOD, and (c and f) GEOS-Chem PM_{2.5} vs. measured PM_{2.5} over South Korea (a-c) and North China (d-f). Different colors represent different seasons. Values inset are correlation coefficients (R) and normalized mean biases (NMB) between surface measurements and GEO satellite or GEOS-Chem values.



Figure 8 compares the seasonalities of AOD and PM_{2.5} over the North China and South Korea regions. The GEO satellite AOD over North China peaks in July and is minimum in winter. Most of AOD is attributed by GEOS-Chem to SNA aerosol, same as in South Korea. AOD over South Korea also has a summer maximum and winter minimum



but with weaker amplitude than over North China. The GEOS-Chem AOD is $\sim 20\%$ biased low in summer and this is largely due to a low RH bias (Figure S8), as seen previously in the KORUS-AQ comparisons but amplified by the high RH in summer that drives hygroscopic growth (Latimer and Martin, 2019).

Surface $PM_{2.5}$ in the observations over North China and South Korea shows opposite seasonality to AOD, with minimum values in summer and maximum values in winter-spring. GEOS-Chem reproduces the strong seasonality of $PM_{2.5}$ in North China and the much weaker seasonality in South Korea. The high $PM_{2.5}$ values over North China in winter in the model are mostly driven by organic aerosol, reflecting the large residential coal burning source (Figure S9; Zheng et al., 2018). In South Korea, by contrast, household energy is mainly from natural gas and electricity (Lee et al., 2020; Woo et al., 2020). GEOS-FP daytime PBL height also shows a stronger seasonality over North China than over South Korea (Figure S8), generally consistent with the CALIPSO daytime PBL height (Su et al., 2018). Previous studies have shown opposite seasonality between MODIS AOD and surface $PM_{2.5}$ over North China and attributed this to the seasonality in PBL height and RH (Qu et al., 2016; Xu et al., 2019). The mean $PM_{2.5}$ /AOD ratio over North China in winter (236 μ g m⁻³) is 8 times that in summer (29 μ g m⁻³), with autumn (94 μ g m⁻³) and spring (89 μ g m⁻³) in between, while over South Korea, the $PM_{2.5}$ /AOD ratio in winter (62 μ g m⁻³) is only 70% larger than in summer (36 μ g m⁻³).

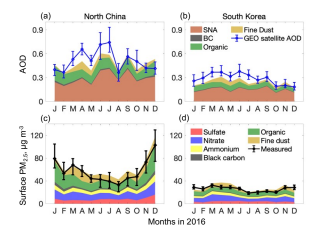


Figure 8. Seasonality of AOD and PM $_{2.5}$ over North China and South Korea, and contributions from individual aerosol components. Lines show regional medians (error bars: 25^{th} and 75^{th} percentiles) for the ensemble of monthly averaged observations in the regions (Figure 6) in 2016. GEOS-Chem values are shown as stacked contours for individual components and are sampled in the same way as the observations.

488 Conclusions 489 Geostationary satellite observations of aerosol optical depth (AOD) over East Asia may usefully complement PM2.5 490 air quality networks if the local relationship between AOD and PM2.5 can be inferred from a physical and/or 491 statistical model. Here we analyzed the ability of the GEOS-Chem chemical transport model to provide this 492 relationship by using a new fused GOCI/AHI geostationary satellite product together with AERONET ground-based 493 AOD measurements, aerosol vertical profiles over South Korea from the KORUS-AQ aircraft campaign (May-June 494 2016), and surface network observations. This allowed us to identify the critical features and limitations of the 495 model for successful representing the AOD-PM_{2.5} relationship. 496 The KORUS-AQ observations show that total aerosol extinction (550 nm) in the vertical column is dominated by 497 sulfate-nitrate-ammonium (SNA) and organic aerosol in the planetary boundary layer (PBL), despite large 498 concentrations of dust in the free troposphere. This reflects the optically effective size and high hygroscopicity of 499 the PBL aerosols. We find that GEOS-Chem aerosol optical properties based on measurements over the North 500 America (default model setting) underestimate KORUS-AQ aerosol mass extinction efficiency by around 20%. In 501 addition, a low bias in GEOS-FP RH below 1 km leads to a 10% underestimate of AOD inferred from the aircraft 502 profile. Adjustments of GEOS-Chem aerosol optical properties and RH enable a successful simulation of the aerosol 503 extinction profile. SNA aerosol contributes 59% of column aerosol extinction in the KORUS-AQ data, while 504 organic aerosol contributes 27% and dust contributes 12%. 505 Comparison of GOCI/AHI geostationary (GEO) satellite AOD to AERONET AODs over South Korea shows good 506 agreement, with high values along the west coast. GEOS-Chem is more consistent with the fine-mode AERONET 507 AOD because of its insufficient accounting of coarse particles, which account for 13% of AERONET AOD. The 508 remaining 15% underestimate of AERONET fine-mode AOD by GEOS-Chem can be attributed to the RH low bias. 509 GEOS-Chem overestimates 24-h surface PM2.5 over South Korea by 43% during the KORUS-AQ period, despite its 510 successful simulation of the aircraft data and fine-mode AERONET AOD, and we find that this is due to a large 511 overestimate of nighttime nitrate. 512 Broader examination of the GOCI/AHI AOD satellite data over East Asia shows spatial distributions and 513 magnitudes consistent with AERONET and featuring in particular strong Asian outflow in spring that includes a 514 large dust component. We find that AODs and PM2.5 have similar large-scale spatial distributions but opposite 515 seasonality. PM2.5 in North China has a strong winter maximum and summer minimum, while AOD shows the 516 opposite. GEOS-Chem simulates successfully the seasonality of measured PM2.5 but is ~ 20% biased low in summer 517 for AOD, due again to RH low bias like that during KORUS-AQ, amplified by the high RH in summer that drives 518 hygroscopic growth (Latimer and Martin, 2019). We find that the opposite AOD and PM2.5 seasonality is mainly 519 driven by residential coal heating sources and low PBL depths in winter, and high RH in summer. Observations of 520 PM_{2.5} and AOD in South Korea show the same seasonal phases as in North China but with much weaker amplitude,

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reflecting the lack of residential coal burning in winter and a weaker seasonal amplitude of PBL depth.

522 In summary, we find that the geostationary GOCI/AHI satellite AOD data provide high-quality information for 523 monitoring of PM_{2.5} over East Asia but that physical interpretation requires accurate information on aerosol size 524 distributions, PBL depths, RH, the role of coarse particles, and diurnal variation of PM2.5, all of which are subject to 525 large uncertainties in chemical transport models. Addressing these uncertainties should be a target of future work. 526 We have used results from our study in a recent machine-learning reconstruction of daily 2011-present PM2.5 over East Asia from GOCI AOD data by identifying critical variables for the machine-learning algorithm and providing 527 528 blended gap-filling data for cloudy scenes (Pendergrass et al., 2021). Besides the factors discussed in this study, 529 topography might be another important factor influencing surface PM2.5 and its vertical mixing (Su et al., 2018), and 530 this also requires future investigation. 531 532 Data availability. Aircraft data during KORUS-AQ are available at: www-air.larc.nasa.gov/cgi-533 bin/ArcView/korusaq. PM2.5 data over China are from: quotsoft.net/air/. PM2.5 data over South Korea are from: 534 www.airkorea.or.kr/web. AERONET data can be found at: aeronet.gsfc.nasa.gov. The MEIC emission inventory are 535 at: www.meicmodel.org/. The KORUSv5 emission inventory is developed by Konkuk University, available at: 536 http://aisl.konkuk.ac.kr/#/emission_data/korus-aq_emissions. 537 538 Author contributions. SZ and DJJ designed the study. SZ performed the data analysis and model simulations with 539 contributions from JFB, KL, HCL, SKK, XW, PL, KRT, and Hong Liao. JK, SL, and Hyunkwang Lim provided 540 satellite AOD data. RJP and JIJ contributed to AirKorea data processing. JM and RM provided the dust emission 541 inventory. GL, FY, and JMM updated wet deposition simulation. JWH, BEA, JED, JLJ, PCJ, and BAN contributed 542 to KORUS-AQ campaign measurements. JHW and YK provided the KORUSv5 emission inventory. QZ provided 543 the MEIC emission inventory. SZ and DJJ wrote the paper with input from all authors. 544 545 Acknowledgement. This work was funded by the Samsung Advanced Institute of Technology and the Harvard-546 NUIST Joint Laboratory for Air Quality and Climate (JLAQC). JLJ, PCJ, and BAN acknowledge NASA grant 547 NNX15AT96G and 80NSSC19K0124 for support. 548 549 Competing interests. The authors declare that they have no conflict of interest.

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