Lidar detection of high concentrations of ozone and aerosol transported from Northeast Asia over Saga, Japan

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4 Osamu Uchino^{1, 2}, Tetsu Sakai², Toshiharu Izumi², Tomohiro Nagai², Isamu Morino¹,
5 Akihiro Yamazaki², Makoto Deushi³, Keiya Yumimoto², Takashi Maki², Taichu Y.

6 Tanaka², Taiga Akaho⁴, Hiroshi Okumura⁴, Kohei Arai⁴, Takahiro Nakatsuru¹, Tsuneo

7 Matsunaga¹, Tatsuya Yokota¹

8

⁹ ¹National Institute for Environmental Studies, 16-2 Onogawa, Tsukuba, Ibaraki 305-8506, Japan

10 ²Meteorological Research Institute, 1-1 Nagamine, Tsukuba, Ibaraki 305-0052, Japan

³Japan Meteorological Agency, 1-3-4 Otemachi, Chiyoda-ku, Tokyo 100-8122, Japan

12 ⁴Saga University, 1 Honjou, Saga, Saga 840-8502, Japan

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14 Correspondence to: O. Uchino (uchino.osamu@nies.go.jp)

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Abstract. To validate products of the Greenhouse gases Observing SATellite (GOSAT), we observed 1718vertical profiles of aerosols, thin cirrus clouds, and tropospheric ozone with a mobile lidar system that 19 consisted of a two-wavelength (532 and 1064 nm) polarization lidar and a tropospheric ozone Differential 20Absorption Lidar (DIAL). We used these lidars to make continuous measurements over Saga (33.24°N, 21130.29°E) during 20-31 March 2015. High ozone and high aerosol concentrations were observed almost 22simultaneously in the altitude range 0.5-1.5 km from 03:00 to 20:00 Japan Standard Time on 22 March 232015. The maximum ozone volume mixing ratio was ~110 ppby. The maxima of the aerosol extinction coefficient and optical depth at 532 nm were 1.2 km⁻¹ and 2.1, respectively. Backward trajectory analysis 24and the simulations by the Model of Aerosol Species IN the Global AtmospheRe (MASINGAR) mk-2 2526and the Meteorological Research Institute Chemistry-Climate Model, version 2 (MRI-CCM2) indicated 27that mineral dust particles originated from the Gobi Desert and an air mass with high ozone and aerosol 28(mainly sulfate) concentrations originated from the North China Plain could have been transported over 29the measurement site within about two days. These high ozone and aerosol concentrations impacted 30 surface air quality substantially in the afternoon of 22 March 2015. After some modifications of its 31physical and chemical parameters, MRI-CCM2 approximately reproduced the high-ozone volume-mixing 32ratio. The MASINGAR mk-2 successfully predicted high aerosol concentrations, but the predicted peak 33 aerosol optical thickness was about one-third of the observed value.

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37 1 Introduction

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39 Tropospheric ozone is a major air pollutant and impacts human health and vegetation (HTAP, 2010; Yue 40 and Unger, 2014). It is also an important greenhouse gas (IPCC, 2013). Tropospheric aerosols are also air 41 pollutants and aggravate respiratory conditions (HTAP, 2010). Tropospheric aerosols also enhance 42radiative forcing in a negative (sulfuric acid particles) or positive (black carbon) way (IPCC, 2013), and 43they affect remote sensing such as the measurement of greenhouse gases from space (Houweling et al., 44 2005; Uchino et al., 2012a). It is therefore very important to monitor tropospheric ozone and aerosols and 45to understand their temporal and spatial variations. The aerosols transported from the East Asia to the 46 western Japan were observed by lidar and their vertical distributions were reported (Iwasaka et al., 1988; 47Murayama et al., 2001; Hara et al., 2009). On the other hand the ozone pollutions from the Asia were 48 mainly studied by the surface measurements (Akimoto et al., 1996; Yamaji et al., 2006). Continuous 49 ozone vertical distributions by ozone DIAL are very useful for studying the transport process and the 50origin. 51To validate products of the Greenhouse gases Observing SATellite (GOSAT), we developed a two-52wavelength (532 and 1064 nm) polarization lidar (hereafter abbreviated as Mie lidar) to observe vertical

profiles of tropospheric and stratospheric aerosols and thin cirrus clouds at the National Institute for Environmental Studies (NIES), Tsukuba (36.05°N, 140.13°E), Japan in 2009. In 2010 we also developed a DIfferential Absorption Lidar (DIAL) to measure tropospheric ozone profiles (hereafter abbreviated as ozone DIAL). The ozone DIAL was installed in a container with the Mie lidar. In March 2011, we moved the lidar container to Saga (33.24°N, 130.29°E) in the Kyushu district of western Japan at a location 2.6 m above sea level. The ozone DIAL was modified in September 2012 (Uchino et al., 2014).

59 Mie lidar has been used to demonstrate the influence of high-altitude aerosols and cirrus clouds on the 60 GOSAT product of the column-averaged dry air mole fraction of carbon dioxide (XCO₂) retrieved from 61 the Thermal And Near infrared Sensor for carbon Observation-Fourier Transform Spectrometer (TANSO-62 FTS) Short-Wavelength InfraRed (SWIR) spectral data onboard GOSAT. The XCO₂ data were improved 63 by taking the vertical profiles of aerosols and cirrus clouds measured by Mie lidar into account (Uchino et 64 al., 2012a). The increases of stratospheric aerosols caused by the 2009 Sarychev eruption and the 2011 65 Nabro eruption were observed by Mie lidar (Uchino et al., 2012b).

66 Ozone DIAL has been used to validate the GOSAT ozone product retrieved from TANSO-FTS 67 Thermal InfraRed (TIR) spectral data (Ohyama et al., 2012), to observe ozone concentrations in the lower 68 troposphere, and to compare the observed concentrations with those predicted by the Meteorological 69 Research Institute Chemistry-Climate Model, version 2 (MRI-CCM2) (Deushi and Shibata, 2011). Use of 70 Mie lidar and ozone DIAL will facilitate satellite product validation not only for GOSAT but also for 71upcoming satellites such as the TROPOspheric Monitoring Instrument (TROPOMI, Veefkind et al., 2012) 72and the Geostationary Environment Monitoring Spectroscopy (GEMS, Bak et al., 2013). High ozone 73episodes in the lower troposphere have been observed by lidar (Banta et al., 1998; Koutidis et., 2002; 74Ancellet et al., 2005; Eisele and Trickl, 2005; Kuang et al., 2011). These observation records were limited to one week at most. We made an 11-day continuous record on 20–31 March 2015.

76 In this paper we report an event during which high concentrations of ozone and aerosols were observed 77 almost simultaneously below an altitude of 1.5 km over Saga on 22 March 2015 by Mie lidar and ozone

- 78 DIAL, which substantially impacted surface air quality. We also compared the observational results with
- those simulated by the models.
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82 2 Characteristics of lidar system and observation parameters

- Mie lidar and ozone DIAL were installed in a container with dimensions of about 228 cm (width), 683 cm (length), and 255 cm (height), as shown in Fig. 1. Mie lidar is a two-wavelength (532 and 1064 nm) polarization lidar based on a neodymium-doped yttrium-aluminum-garnet (Nd:YAG) laser; the characteristics are summarized in Table 1. The output energy at 532 and 1064 nm was 130 mJ, with a pulse repetition rate of 10 Hz. The diameter of the receiving telescope was 30.5 cm. The output signals from the photomultiplier tubes (PMT) and a silicon avalanche photodiode (APD) were processed by transient recorders with a 12-bit analog/digital converter and a photon counter.
- 91 The data analysis methods of Mie lidar and ozone DIAL have been described by Uchino et al. (2012b)
 92 and Uchino et al. (2014), respectively. We summarize the observation parameters obtained by Mie lidar.
 93 The backscattering ratio *R* is defined as
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95 R = (BR + BA)/BR,

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97 where *BR* and *BA* are the Rayleigh and Mie backscattering coefficients, respectively. Backscattering ratio 98 profiles were derived by the inversion method (Fernald, 1984). The reference altitude was usually set 99 between 9 and 12 km where only molecular backscattering could be assumed when there were no clouds. 100 We assumed the lidar ratio LR (extinction-to-backscatter ratio) for aerosols to be 50 sr at 532 nm and 45 101 sr at 1064 nm based on the lidar ratios for Asian dust and pollution aerosols summarized by Sakai et al. 102(2003), Anderson et al. (2003) and Cattrall et al. (2005). Their summaries are as follows: Sakai et al. 103 (2003): Asian dust 47 \pm 18 sr, Cattral et al. (2005): dust (spheroids) 42 \pm 4 sr, South East Asia pollution 104 58 \pm 10 sr, Anderson et al. (2003): ACE-Asia pollution (fine-dominated, submicron portion) 50 \pm 5 sr, 105dust (coarse-dominated, dust-like chemistry, supermicron portion) 46 ± 8 sr. As a simplification, we used 106 the same value for both species. To calculate BR, we used the atmospheric molecular density profiles 107obtained by operational radiosondes at the Fukuoka District Meteorological Observatory (33.58°N, 108130.38°E), Japan Meteorological Agency (JMA). The aerosol extinction coefficient was calculated by 109multiplying *BA* by *LR*.

110 The total volume depolarization ratio *D* was defined as

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112 $D = S / (P + S) \cdot 100 (\%),$

(1)

114 where *P* and *S* are the parallel and perpendicular components of the backscattered signals, respectively. 115 The particle depolarization ratio D_p was obtained from the equation

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$$D_p = (D \cdot R - D_m)/(R - 1),$$
 (3)

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119 where D_m is the atmospheric molecular depolarization ratio. We used a D_m value of 0.37% for this lidar 120 system; we calculated D_m from the spectral transmission data of the interference filter at 532 nm and the 121 Rayleigh backscattering cross sections (Sakai et al., 2003). The value of D_p indicates whether the particles 122 are spherical or non-spherical; large values indicate the presence of non-spherical particles. The 123 backscatter-related Ångström exponent Alp, the qualitative indicator of aerosol particle size, is defined by

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$$BA(\lambda) \propto \lambda^{-Alp}$$
, (4)

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127 where λ is the wavelength. Larger values of *Alp* indicate the predominance of smaller (i.e.,

128 submicrometer-sized) particles. The vertical resolution of these observation parameters was 150 m, and

129 the time resolution was set to be 1 h for comparison with the Model of Aerosol Species in the Global

130 Atmosphere (MASINGAR)-mk2 (Yukimoto et al., 2012). The lowest altitude of Mie lidar measurement

131 was 225 m due to the non-perfect overlap of the transmitter-receiver optical axes of the lidar system.

The ozone DIAL consisted of a Nd:YAG laser and a 2-m-long Raman cell filled with CO_2 gas that generated four Stokes lines from stimulated Raman scattering by CO_2 ; the characteristics are summarized in Table 2. In this study, we used three Stokes lines (276, 287, and 299 nm). The output energies of these Stokes lines were about 8–9 mJ per pulse, with a pulse repetition rate of 10 Hz. The receiving telescope diameters were 10 cm for boundary layer ozone measurements and 49 cm for free tropospheric ozone measurements. The Mie lidar and ozone DIAL were synchronized by two pulse-delay generators.

The 276/287 nm and 287/299 nm wavelength pairs were used for ozone DIAL measurements in the altitude ranges of 0.57–2.0 km and 2.0–6.0 km, respectively. The effective vertical resolutions were 270 m for 0.57–2.0 km and 540 m for 2.0–6.0 km, respectively (Uchino et al., 2014). The time resolution was set to 1 h to facilitate comparison with the MRI-CCM2. The aerosol correction was not made for the ozone retrieval. Next, we report the continuous lidar observational results made at Saga from 20 March to 31 March 2015.

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146 **3 Ozone DIAL data**

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Figure 2a shows a time-altitude cross-section of ozone volume mixing ratios observed by DIAL at Saga
from 11:10 JST on 20 March to 14:33 JST on 31 March 2015. Lidar observations were not obtained from
15:56 JST on 27 March to 21:58 JST on 29 March 2015, mainly because of rainy or cloudy conditions.

We made quality checks of the DIAL data. The gray regions in Fig. 2a correspond to areas where there were no observational data or the errors were larger than 10%. The errors were computed from the lidar signal-to-noise ratios by use of Poisson statistics. Regions surrounded by a black rectangle are areas where the data were affected by aerosols and/or clouds with *R* larger than 2 at 299 nm, which were calculated assuming LR=50 sr without correcting attenuation by ozone absorption. In the lowest row of

- 156 Fig. 2a, we show hourly data of surface oxidant volume mixing ratios (Ox) at Takagimachi in Saga city
- 157 measured by the Saga Prefectural Environmental Research Center (https://www.pref.saga.lg.jp/web/at-158 contents/kankyo1/shisetsu/ 40810/ 41304/ 67819.html). Takagimachi is located about 2.8 km northeast
- 159 from the ozone DIAL site. Because the surface Ox was observed by an UV photometer, the contribution 160 of other components such as peroxyacetyle nitrate (PAN) to oxidant concentrations was extremely low, 161 and the oxidant volume mixing ratio was considered to be that of ozone.

162Figure 2a indicates that the ozone volume mixing ratios measured by DIAL were usually about 50-70 163ppbv during the study period. Comparatively high ozone concentrations, >75 ppbv, were detected at 164 altitudes of 0.57-3 and 0.57-2 km on 20-23 March and 30-31 March, respectively. Notably high ozone 165volume mixing ratios of 90–110 ppbv at altitudes of 0. 57–1.5 km were observed from 03:00 to 20:00 JST 166on 22 March. These high ozone concentrations were also seen in the surface photochemical oxidants data, 167 i.e., the Ox equaled 92-101 ppbv from 15:00 to 21:00 JST on 22 March, as shown in the lowest row in 168Fig. 2a. The maximum concentration of Ox was 101 ppbv at 16:00 JST. This maximum value was far 169 above the environmental quality standard of 60 ppbv for hourly photochemical oxidants in Japan 170(https://www.env.go.jp/en/air/aq/aq.html).

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172 **3.1 Comparison of DIAL data with MRI CCM-2**

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174The MRI-CCM2 is a global model that simulates chemical and physical processes that affect the 175distribution and evolution of ozone and other trace gases from the surface to the stratosphere (Deushi and 176Shibata, 2011). Uchino et al. (2014) have provided an outline of MRI-CCM2. The vertical resolution of 177the model increases from about 100 to 600 m from the surface to 6 km. The time step of the transport 178(chemistry) scheme is 30 (15) min. We used hourly model output data. The horizontal resolution is about 179110 km. We examined whether or not the model could simulate DIAL observational results. The MRI-180CCM2 simulated the DIAL observations reasonably well. However, the MRI-CCM2 predicted the high 181 ozone concentrations of 50-60 ppbv and could not reproduce those of 90-110 ppbv observed with DIAL 182below an altitude of 1.5 km during 03:00-20:00 JST on 22 March 2015.

We therefore performed some simulations in which we changed the emission inventory data and the term that forced the reanalysis wind field. The most reasonable results which were shown in Fig.2b were obtained when the following changes were made. The e-folding time of the nudging term was changed from 18 hours to 12 hours to more strongly force the simulated wind fields toward the reanalysis data. In addition, we changed the emission inventory of Regional Emission inventory in Asia version 1.1 (REAS

188 1.1) (Ohara et al., 2007) to the REAS 2.1 emission inventory in 2007 (Kurokawa et al., 2013) and the

- 189 NO₂/NO_x emissions ratio from 5% to 15% by volume, which is within the range of uncertainty (Carslaw,
- 190 2005). The emission inventory of NO_x increased about 50% from REAS 1.1 to REAS 2.1. Figure 2c
- 191 shows the differences between the observed and simulated ozone mixing ratios. Simulated ozone volume
- mixing ratios were about 60–70 ppbv below an altitude of 1.5 km from 14:00 JST on 21 March to 21:00
- 193 JST on 22 March 2015, lower by about 20-50 ppbv compared with the DIAL results. And the MRI-
- 194 CCM2 predicted the high ozone concentration a half day earlier than the DIAL observation.
- 195 The maximum bias (systematic error) of ozone DIAL data caused by aerosols was estimated to be 20%
- 196 (15 ppbv) at 0.57 km, and the mean bias and the standard deviation were $7\% \pm 5\%$ in the altitude range
- 197 0.57–2.0 km at 11:00 JST. These biases were estimated from *Alp* observed at the same time by Mie lidar
- and assuming LR = 50 sr in the wavelength range 276–299 nm, based on the equations of (6) and (7) in
- 199 Uchino and Tabata (1991). These biases were not large since the 276/287 nm and 287/299 nm wavelength
- 200 pairs were suitable for measurements of ozone in the boundary layer and the free troposphere respectively
- 201 (Nakazato et al., 2007). As mentioned earlier, the ozone DIAL data with the statistical error smaller than
- 202 10% was used in this study. Therefore the uncertainty of the ozone DIAL data was estimated to be smaller
- than 22% and the mean value of the uncertainty was 12%. A model with higher horizontal resolution
- might be necessary to more realistically simulate high surface ozone concentration events in the planetaryboundary layer.
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208 4 Mie lidar data

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Figures 3a, 3b, and 3c show time-altitude cross-sections of the backscattering ratio (R), the total volume depolarization ratio (D), and the particle depolarization ratio (D_p), respectively, observed by Mie lidar at Saga from 11:10 JST on 20 March to 14:33 JST on 31 March 2015. Mie lidar data were not obtained from 15:56 JST on 27 March to 21:58 JST on 29 March 2015, mainly because of rainy or cloudy conditions. We made quality checks of Mie lidar data. Gray regions are areas where there were no observational data or the data were affected by clouds.

Aerosol layers with *R* in the range 2–4 almost always existed below an altitude of 2.5 km for 20–31

217 March 2015. An event of high aerosol loading with large values of R (>8) was observed below altitudes of

- 218 1.5 km for 03:00–21:00 JST on 22 March, when the values of *D* were small (the mean and the standard
- deviation: $3.9 \pm 2.1\%$) compared with those before and after the event, when the values of D were larger
- 220 than $7.9 \pm 2.1\%$ during 15:00 JST on 21 through 15:00 JST on 23, except for 03:00-21:00 JST on 22.
- 221 The main aerosol component during the event might be submicrometer-sized spherical particles, because
- 222 D_p was small (4 ± 2%), and the wavelength exponent Alp was large (1.3 ± 0.3). In contrast, the main
- 223 aerosol particles before and after the event could be supermicrometer-sized, nonspherical mineral dust
- particles because D_p was comparatively large (13 ± 3%) and Alp was 1.0 ± 0.2 (Sakai et al., 2003; Cattrall
- et al., 2005). When there were no clouds above, R at 1064 nm was estimated assuming Alp=1.5 at the
- reference altitude where very small amount of aerosols was expected to be present, i.e., $R=1.06 \pm 0.06$

- $(D=1.2 \pm 0.5)$ at 532 nm, in the altitude range 3–6 km. If the value of *Alp* was changed from 1.0 to 2.0 at the reference altitude, the uncertainty in *Alp* was estimated to be ± 0.2 . *Alp* was 0.3–2.0 in the 11-day Mie
- lidar record. The maximum errors of D and D_p were 0.1% and 2% for R>2 at 532 nm.
- During the same time period, high aerosol concentrations were also observed at the surface (Fig.4).
- $231 \qquad \text{Hourly values of the mass concentrations of particulate matter with a diameter of 2.5 \ \mu\text{m or less} (PM_{2.5}) \ \text{at}}$
- Takagimachi measured by the Saga Prefectural Environmental Research Center were 23 μ g m⁻³ at 10:00 JST and increased up to a maximum value of 110 μ g m⁻³ at 15:00 JST on 22 March; the concentrations
- were greater than 82 μ g m⁻³ during 13:00–16:00 JST and decreased to 17 μ g m⁻³ at 01:00 JST on 23
- 235 March. The daily mean value of $PM_{2.5}$ was 50.6 µg m⁻³ for 24 hours on 22 March at Takagimachi, larger

than the environmental quality standard of 35 μ g m⁻³ in Japan (https://www.env.go.jp/en/air/ag/ag.html).

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4.1 Comparison of Mie lidar data with MASINGAR mk-2

- 240The MASINGAR-mk2 is an improved version of the MASINGAR aerosol model (Tanaka et al., 2003); it 241treats five aerosol species: sulfate, black and organic carbon, sea salt, and soil dust. We used emission 242data for sulfur dioxide and for black and organic carbon from MACCity (Granier et al., 2011). Soil dust 243and sea salt were represented by 10 bins with particle diameters of 0.2-20 µm. The model was coupled 244online with the atmospheric general circulation model MRI-AGCM3 (Yukimoto et al., 2012). The 245Meteorological fields were from JMA Global Analysis data (GANAL). The horizontal resolution of the 246MASINGAR-mk2 was about 60 km, and the number of vertical layers was 40 from the surface to 0.1 247hPa. The vertical resolutions were 100, 300, and 600 m at the lowest level and altitudes of 1 and 6 km, 248respectively. The time step of the transport (chemistry) scheme was 450 seconds, and we used hourly 249model output data.
- Figures 4a and 4b show the time-height cross sections of aerosol extinction coefficients observed by Mie lidar and simulated by MASINGAR-mk2, respectively. Figure 4c represents the difference between the observed and simulated extinction coefficients. The model was able to capture the general characteristics of the observational results rather well. A close look at Fig. 4c reveals that the model underestimated the aerosol extinction coefficients of the anthropogenic pollutant event on 22 March but slightly overestimated the extinction coefficients associated with particles having larger total volume depolarization ratios on 30 and 31 March (i.e., dust-dominant case).
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258 **4.2 Comparison of aerosol optical depths**

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Figure 5 shows temporal variations of the aerosol optical depths (AOD) measured by Mie lidar at 532 nm

- and sky radiometer at 500 nm (Kobayashi et al., 2006, Uchino et al., 2012a) and simulated at 550 nm by
- 262 MASINGAR-mk2 from 20 to 31 March. To estimate AODs from the lidar data, the extinction coefficient
- at 225 m was extrapolated to the ground, the extinction coefficient from 15 to 35 km was observed at
- night on the same day, and *S* was assumed to be 50 sr for all altitudes. When clouds and thick aerosols

- were present, AODs were not obtained. The sky radiometer was positioned on the roof of the building,
- which is four stories high and located to the west of the container (brown building in Fig. 1). Although it
- 267 must be noted that the measured and simulated wavelengths differed slightly, the AODs were almost the
- same, except for the high aerosol and ozone event on 22 March. The mean bias and the standard deviation
- of AOD between Mie lidar and sky radiometer was 0.029 ± 0.051 , and that between MASINGAR mk-2
- and sky radiometer was -0.07 ± 0.24 for 20–31 March, except for 12:00–14:00 JST on 22 March. The
- 271 maximum values of the AODs were 2.1 at 12:00 JST by lidar, 1.92 at 13:00 JST by sky radiometer, and
- 272 0.53 at 13:00 JST by MASINGAR-mk2. One possible reason for the large difference in AOD (~0.2)
- 273 between Mie lidar and Sky radiometer data is that we set the reference altitudes 8.2 km and 2.8 km at
- 12:00 and 13:00 JST on 22 March respectively for the lidar because the backscattered signals were
- strongly attenuated by the dense aerosol layers below 2 km. This might cause error in AODs for the Mie
- lidar data. The sky radiometer could have different sight than the Mie lidar. This might be also a possiblereason for the difference.
- The model underestimated the AODs by factors of about 3.6–4 compared to the sky radiometer and
- 279 lidar observations. One plausible reason for that is that the model resolution (about 60 km) was
- insufficient to reproduce the observed prominent peak in which the observed AOD increased from 1.0 to
- 2.0 in 6 hours. The other plausible reason for the underestimation is the uncertainty of the emissions
 inventories of aerosol precursors. Grainer et al. (2011) collected various emission inventories and
 compared them in global scales. They found that differences in Chinese sulfur dioxide (SO₂) emissions in
- 284 2000 reached 66% between the lowest and highest emissions and concluded that there was no consensus285 among the different inventories for the emissions of Chinese SO₂ This large variation among the
- 286 inventories indicates that estimate of SO₂ emission in China has large error. In their comparison, the
- 287 MACCity emission which was used in MASINGAR-mk2 simulation, showed the lower amount of
- 288 Chinese SO_2 emission among the inventories. This might be responsible for the underestimation of
- pollution aerosol (sulfate) concentrations. In MASINGAR-mk2, dust emission flux is estimated by a
 parameterized dust emission scheme and has strong dependency upon various parameters (i.e., soil
- texture, soil wetness, land use, snow cover fraction, vegetation cover, surface wind speed, etc.). The dust
- 292 model intercomparison project (DMIP; Uno et al., 2006) reported that simulated dust emission amounts
- 293 over East Asia among eight dust models (including the former version of MASINGAR) differed
- sometimes by a factor of ten. These facts indicate that estimate of dust emission also causes large errors.
- 295 To solve this problem, for example, it might be better to use the near real-time satellite data of SO_2 and
- nitrogen dioxide (NO₂) provided by the Ozone Monitoring Instrument (OMI) onboard NASA's Aura
 satellite (Krotkov et al., 2016)-, and/or to use a data assimilation technique that integrates model
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- 301 5 Discussion: origin and transport pathways of ozone and aerosol plumes

simulation and observation data (Yumimoto et al., 2016).

303 Figure 7 shows the time-altitude cross sections of total aerosol extinction coefficients at 550 nm, and the

- 304 ratios of dust extinction coefficients to total aerosol extinction coefficients simulated by MASINGAR-
- 305 mk2 with potential temperatures over Saga for 20–31 March 2015. For the event on 22 March, the model
- 306 predicted the dust particles (about 60–100%) in the altitude range 1–3 km, and sulfate (about 40–60%) 307 and dust (about 30–40%) particles below 1 km. The number of the parenthesis represents the ratio of each
- 308 component's extinction coefficient to the total extinction coefficient. The dust particles descended to the
- 309 surface in the afternoon (Fig.7b). For the event on 30 March, MASINGAR mk-2 predicted the dust
- 310 particles (about 50-100%) for 1-6 km, and sulfate (about 50-80%) and dust (about 0-20%) particles

below 1 km in the morning. Mie lidar data support the model prediction because D_p is high (17 ± 6%) for 1–3 km and low (10 ± 3%) below 1 km. For both events, small amounts of organic carbon, black carbon

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- and sea salt particles were predicted.
- 314To identify the origin of the aerosols and related transport processes, three-dimensional backward 315trajectories of air parcels were calculated with the NOAA Hybrid Single Particle Lagrangian Integrated 316 Trajectory (HYSPRIT) model (Draxler and Hess, 1998; Stein et al., 2015). An air parcel was initially left 317at altitudes of 1500 m (Fig. 8a) and 500 m (Fig.9a) over the lidar site at Saga. The trajectories were 318calculated for three days from 21:00 UTC on 21 (06:00 JST on 22) March 2015. Figures 8b and 9b show 319 the time-altitude cross sections of dust and sulfate extinction coefficients simulated by MASINGAR mk-2 320 along the trajectory paths of Figs. 8a and 9a, respectively. Based on the results of the backward 321trajectories and the model simulations, the dust and sulfate particles on 22 March could have been 322transported within about two days from the Gobi Desert and the North China Plain (NCP), respectively, to 323the measurement site. The MASINGAR mk-2 simulation suggested that the dust particles emitted during 32418:00-24:00 UTC on 19 March around 40°N and 130.29°E were responsible for the dust storm captured 325by the Mie lidar observation. The highest concentrations of SO_2 and NO_2 in the world were observed in 326 NCP for 2013–2015 by the Ozone Monitoring Instrument (OMI) onboard NASA's Aura satellite, as 327 shown in Fig.5 by Krotkov et al. (2016). These gases are important precursors of sulfate particles and 328 ozone. Figure 10 represents the horizontal maps of ozone volume mixing ratios at 925 hPa (about 760 m 329 altitude) simulated by MRI-CCM2 at 21:00 JST on 19, 20, 21 March and at 03:00 JST on 22 March 2015. 330These maps indicate that the high ozone could be transported from NCP to the Yellow Sea and then Saga 331within about two days.
- 332Because it was difficult to obtain observational data of surface ozone and sulfate particles in NCP 333 including Beijing on 19-20 March, we refer to the following papers related to those data. According to the 334ozonesonde measurements made by Wang et al. (2012), ozone concentrations ≥ 90 ppby were observed 335over Beijing, China in late March. Ma et al. (2016) reported a significant increase of surface ozone from 336 2003 to 2015 at Shangdianzi (40.65°N, 117.10°E), which is located about 100 km northeast of suburban 337 Beijing, and the maximum daily average 8-h concentrations of ozone appear to have been >100 ppbv in 338 March 2015 based on Fig. 2 in their paper. High PM_{2.5} and submicron aerosol concentrations have been 339observed in Beijing (Zhang et al., 2013; Sun et al., 2015). Ozone and aerosol concentrations may 340therefore have been high in March 2015 over NCP.

To investigate the vertical transport processes of the aerosol and ozone in the lower troposphere over the measurement site, we show in Fig. 11 the time variations of the top altitudes of the mixed layers from two hours after sunrise to two hours before sunset during 11:10 JST on 20 March through 14:33 JST on 31 March 2015 which were estimated from the 1064 nm range-corrected backscatter signals with a range

resolution of 15 m using the wavelet covariance transform method (Baars et al., 2008; Izumi et al., 2016),

and those obtained from the radiosonde data at Fukuoka and the JMA Meso-scale Analysis (MA) data

347 over Saga using the parcel method (Holzworth, 1964). When the mixed layers developed in the afternoon,

348 the tops of the mixed layers (1.5–2 km) estimated by Mie lidar were almost consistent with those by MA.

Although the radiosonde data at 9:00 JST on 22 March found the top of the mixed layer was 117 m (Stull,

1988), it was difficult for Mie lidar to detect the mixed layer because the lowest altitude of the Mie lidarmeasurement was 225 m.

352The dust particles originated from the Gobi Desert arrived at 1-3 km altitudes over the lidar site at 35306:00 JST on 22 March. When the mixed layer developed to 1.5-2 km at 11:00-15:00 JST on 22, the dust 354particles were supposed to be mixed into the boundary layer and then reached the surface by the 355entrainment, as simulated in Fig.7b. This could result in the sharp increase in PM2.5 concentrations at the 356surface after 11:00 JST, as shown in Fig.4. The similar phenomenon was observed over the northern 357Kyushu area during the dust event in late May-early June 2014 (Uno et al., 2016). A similar high-surfaceozone event was observed by eight ozonesonde measurements during 6-9 June 2003 over the Seoul 358359metropolitan region (Oh et al., 2010).

360 361

362 6. Concluding remarks

363

By using ozone DIAL and a two-wavelength polarization (Mie) lidar, we made continuous measurements of ozone and aerosol concentrations over Saga during 20–31 March 2015. High ozone and high aerosol concentrations that occurred nearly simultaneously were observed in the altitude range 0.5–1.5 km from 03:00 to 20:00 JST on 22 March 2015. The ozone volume mixing ratio was larger than 100 ppbv. The aerosol extinction coefficient and AOD at 532 nm were larger than 0.5 km⁻¹ and 1.5, respectively.

369 Backward trajectory analysis and the simulations by the MASINGAR mk-2 and the MRI-CCM2 370 models indicated that the mineral dust particles originated from the Gobi Desert and an air mass with high 371ozone and aerosol (mainly sulfate) concentrations originated from the North China Plain could have been 372transported over the lidar site within about two days. Based on the lidar and surface measurement data 373and the simulation by MASINGAR-mk2, there is a possibility that the air mass with high ozone and 374 aerosol concentrations could have been transported from the lower troposphere to the surface by vertical 375mixing when the planetary boundary layer developed in the afternoon of 22 March 2015. The 376 combination of ozone DIAL measurements with surface in-situ ozone measurements is very useful for 377 studying the process of descent of high ozone concentrations in the lower troposphere to the surface and 378 the impacts on surface air quality. Such measurements of pollution plumes that descend from the free troposphere to the surface are highly recommended (HTAP, 2010).

380 The MRI-CCM2 could approximately reproduce the high-ozone volume-mixing ratios after some 381 modifications of physical and chemical parameters. MASINGAR mk-2 successfully predicted high 382 aerosol concentration events, but the predicted peak AOD was about one-third of the observed AOD. For 383 further improvement of these models, it will be important to continue comparing these models with ozone

- 384 DIAL, Mie lidar, and surface in-situ ozone and particle measurements.

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 concentrations of surface oxidant and PM_{2.5} measured by the Saga Prefectural Environmental Research
 Center. The NOAA Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPRIT) model was used
 to calculate backward trajectories of air parcels.

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 $\frac{580}{581}$

Fransmitter				
Laser	Nd:YAG			
Wavelength	532 nm 1064 nm			
Pulse energy	130 mJ	130 mJ		
Pulse repetition rate	10 Hz			
Pulse width	8 ns			
Beam divergence	0.2 mrad	0.2 mrad		
Receiver	· · · · · · · · · · · · · · · · · · ·			
Telescope type	Schmidt Cassegrain			
Telescope diameter	30.5 cm			
Focal length	304	8 mm		
Field of view	1 m	rad		
Polarization	P and S	None		
Number of channels	3	1		
Interference filter				
Center wavelength	532.0 nm	1064.1 nm		
Bandwidth (FWHM)	0.29 nm	0.38 nm		
Transmission	0.66	0.58		
Detectors	PMT	APD		
	(Hamamatsu R3234	4-01) (EG&G C30956EH)		
Signal processing	12bit A/D + Photon	counting		
Time resolution	1 min			
Vertical resolution	7.5 m			

Table 1. Characteristics of Mie lida

Transmitter						
Pump laser	Nd:YAG					
Wavelength	266 nm					
Pulse energy	107 mJ					
Pulse repetition rate	10 Hz					
Pulse width	8 ns					
Raman active gas	CO_2					
Stokes lines	276 nm	287 nm	299 nm	312 nm		
Pulse energy	7.5 mJ	9.1 mJ	8.4 mJ	No. meas.		
Beam divergence	0.1 mrad					
Receiver						
Telescope type	Newtonian			Prime focus (fiber coupled)		
Telescope diameter	49 cm			10 cm		
Focal length	1750 mm			320 mm		
Field of view	1 mrad			3 mrad		
Interference filter						
Center wavelength	287.2 nm	299.0 nm	312.0 nm	276.1 nm	287.2 nm	
Bandwidth (FWHM)	1.02 nm	1.15 nm	0.82 nm	1.07 nm	1.05 nm	
Transmission	0.18	0.32	0.36	0.17	0.21	
Detectors	PMT (Hamamatsu R3235-01)					
Signal processing	12bit A/D + Photon counting					
Time resolution	1 min					
Vertical resolution	7.5 m					

Table 2. Characteristics of tropospheric ozone DIAL system



 $\begin{array}{c} 685\\ 686 \end{array}$

Figure 1. Mie lidar and ozone DIAL (right) were installed in the container at the left on the ground (left).





719Figure 2. Time-altitude cross-sections of (a) ozone volume mixing ratios observed by DIAL over Saga 720 from 11:10 JST on 20 March to 14:33 JST on 31 March 2015, (b) the ratios simulated by a modified 721MRI-CCM2 for 20-31 March 2015, and (c) the difference between the observed and simulated ozone 722volume mixing ratios (a-b). Gray regions indicate areas where there were no observational data or the 723statistical errors were larger than 10%. Regions enclosed with black rectangles are areas where the data 724were affected by aerosols and/or clouds. The lowest row in Fig. 2a shows photochemical oxidant (ozone) 725volume mixing ratios at Takagimachi in Saga city as measured by the Saga Prefectural Environmental 726 Research Center.



Figure 3. Time-altitude cross-sections of (a) backscattering ratios, (b) total volume depolarization ratios, and (c) particle depolarization ratios for *R* larger or equal to 2.0 at 532 nm observed by Mie lidar at Saga from 11:10 JST on 20 March to 14:33 JST on 31 March 2015. Lidar observations were not available from 15:56 JST on 27 March to 21:58 JST on 29 March 2015 mainly because of rainy or cloudy conditions. Gray regions are areas where there were no observational data or where the observations were affected by clouds.

PM2.5 and Ox surface observation





Figure 4. Hourly (JST) data of surface PM2.5 (red line) and Ox (blue line) measured by the Saga
Prefectural Environmental Research Center for 20-31 March 2015. The volume mixing ratio of Ox was
considered to be that of ozone.



Figure 5. Time-altitude cross sections of (a) aerosol extinction coefficients observed by Mie lidar at 532 nm over Saga from 11:10 JST on 20 March to 14:33 JST on 31 March 2015, (b) the coefficients simulated by MASINGAR-mk2 at 550 nm for 20-31 March 2015, and (c) the difference between the Mie lidar observations and the simulation (a-b). Gray regions represent areas where there were no observational data.

Aerosol optical deapth





Figure 6. Temporal variation of the aerosol optical depth (AOD) measured by Mie lidar at 532 nm (red
circles), by sky radiometer at 500 nm (blues circles), and simulated at 550 nm by MASINGAR-mk2
(green circles).





Figure 7. Time (JST)-altitude cross sections of (a) total aerosol extinction coefficients at 550 nm (color shading) and (b) ratios of dust extinction coefficient to total aerosol extinction coefficient (color shading) simulated by MASINGAR-mk2 with potential temperatures (black contours) over Saga for 20-31 March 2015. The gray regions in Fig. 7b indicate that the simulated total aerosol extinction coefficient is less than 0.02.







Figure 8. (a) 72-h HYSPLIT-model backward trajectory (red line) and terrain height (black line) from Saga at 1500 m above ground level (AGL) ending at 06:00JST on 22 May 2015. (b) Time-altitude cross section of dust extinction coefficient simulated by MASINGAR mk-2 along the trajectory path.

Dust Extinction Coefficient (1/km)

0.2

0.25

NOAA HYSPLIT MODEL Backward trajectory ending at 2100 UTC 21 Mar 15 GDAS Meteorological Data 33.24 N 130.29 E **f**16 at Source 🖈 -34 -32 Meters MSL
 18
 12
 06
 00
 18
 12
 06
 00
 18
 12
 06

 Job ID: 12514
 Job Start: Fri Sep 9 01:32:38 UTC 2016
 Job Start: Fri Sep 9 01:32:38 UTC 2016
 Source 1
 lat.: 33.240000
 lon.: 130.290000
 height: 500 m AGL
 03/19 Trajectory Direction: Backward Duration: 72 hrs Vertical Motion Calculation Method: Model Vertical Velocity Meteorology: 0000Z 15 Mar 2015 - GDAS1



(a)



- - -

Figure 9. Same as Fig. 8 but for 500 m AGL.





Figure 11. Time-altitude cross section of (color shaded) range-corrected backscatter signal at 1064 nm
and the tops of the mixed layers estimated by Mie lidar (closed black circles), radiosonde (open triangles),
and JMA Meso-Scale Meteorological Analysis (open squares) data over Saga from 11:10 JST on 20
March to 14:33 JST on 31 March 2015.