

1 Retrieval Algorithm for Aerosol Effective Height from the
2 Geostationary Environment Monitoring Spectrometer (GEMS)

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4 Sang Seo Park^{1,*}, Jhoon Kim², Yeseul Cho², Hanlim Lee³, Junsung Park³, Dong-Won
5 Lee⁴, Won-Jin Lee⁴, Deok-Rae Kim⁴

6 ¹ *Department of Urban and Environmental Engineering, Ulsan National Institute of Science*
7 *and Technology, Ulsan, Korea*

8 ² *Department of Atmospheric Sciences, Yonsei University, Seoul, Korea*

9 ³ *Division of Earth and Environmental System Sciences, Pukyong National University, Busan,*
10 *South Korea*

11 ⁴ *Environment Satellite Center, National Institute of Environmental Research, Incheon, Korea*

12
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15 *Corresponding author. Sang Seo Park (sangseopark@unist.ac.kr)

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Abstract

An algorithm for aerosol effective height (AEH) was developed for operational use with observations from the Geostationary Environment Monitoring Spectrometer (GEMS). The retrieval technique uses the slant column density of the oxygen dimer ($\text{O}_2\text{-O}_2$) at 477 nm, which is converted into AEH after retrieval of aerosol and surface optical properties from GEMS operational algorithms. The AEH retrieval results show significant AEH values and continuously monitor aerosol vertical height information in severe dust plumes over East Asia, and the collection of plume height information for anthropogenic aerosol pollutants over India. Compared to the AEH retrieved from Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP), the retrieval results show bias of -0.03 km with a standard deviation of 1.4 km for the AEH difference over the GEMS observation domain from January to June, 2021. The AEH difference depends on aerosol optical properties and surface albedo. Compared to the aerosol layer height obtained from the tropospheric monitoring instrument (TROPOMI), differences of 1.50 ± 1.08 km, 1.59 ± 1.22 km, and 1.71 ± 1.24 km were obtained for pixels with single scattering albedo (SSA) < 0.90 , $0.90 < \text{SSA} < 0.95$, and $\text{SSA} > 0.95$, respectively.

Keywords: aerosol effective height, aerosol optical depth, environmental satellite, GEMS

1. Introduction

Since the launch of the Total Ozone Mapping Spectrometer (TOMS) on Nimbus-7, ultraviolet (UV)-visible satellite measurements have been used for environmental monitoring of the distribution and reaction processes of pollutants (e.g., anthropogenic aerosols, tropospheric ozone, NO₂, and SO₂). Measurements from environmental satellites have been used to estimate gaseous species in the atmosphere, resulting in vertical column integrated amounts. However, these column-integrated amounts and associated surface concentrations have uncertainty due to simultaneous changes in optical path length associated with the vertical distribution of target species and amounts of scattering materials (clouds and aerosols) present. In addition, aerosol vertical information is also important information for the application for tropospheric concentration of aerosols. For example, aerosol height information in the free troposphere is particularly important for aviation safety by affecting the visibility. Also, scientific applications including radiative forcing studies, long-range transport modelling and studies of cloud formation processes have been used aerosol vertical information as an input parameter.

Environmental satellite sensors, that measure UV-visible wavelength, have been used the UV aerosol index (UVAI) for aerosol detection (e.g., Buchard *et al.*, 2015; Herman *et al.*, 1997; Torres *et al.*, 1998, 2002; Prospero *et al.*, 2000; de Graaf *et al.*, 2005). Furthermore, scattering radiative index values were investigated for the possibility of the cloud signal detection (Penning de Vries *et al.*, 2009, 2015; Kooreman *et al.*, 2020; Kim *et al.*, 2018). However, these indices only have qualitative characteristics and limitations to identify aerosol amounts.

For the quantitative estimation, measurements of aerosol optical depth (AOD) and

radiative cloud fraction have also been retrieved from pixel-based radiance data in UV-visible wavelength range. Recently, various aerosol retrieval algorithms have been developed to be applied in passive satellite sensors. These algorithms focus on improved trace gas retrieval as well as direct monitoring of aerosol properties, such as AOD and single scattering albedo (SSA) (e.g., Ahn *et al.*, 2014; Kim *et al.*, 2020; Torres *et al.*, 2020).

Although the algorithms developed for environmental satellite sensors indicate the presence and amount of scattering materials, the accuracy of these retrieval algorithms for trace gases is affected by the relative vertical distributions between trace gases and scattering materials (e.g., Lorente *et al.*, 2017; Hong *et al.*, 2017). For this reason, estimating cloud vertical parameters is important. For cloud vertical information, cloud height information has been estimated simultaneously with cloud optical depth and radiative cloud fraction data using the rotational Raman scattering (Joiner and Vasilkov, 2006; Vasilkov *et al.*, 2008; Joiner and Bhartia, 1995) and absorption intensity of the oxygen dimer (O_2-O_2) (Accarreta *et al.*, 2004; Vasilkov *et al.*, 2018; Choi *et al.*, 2021) combined with normalized radiance.

Similarly, the aerosol vertical distribution can be estimated using the oxygen absorption bands, such as the O_2-O_2 (Park *et al.*, 2016; Chimot *et al.*, 2017; Choi *et al.*, 2019, 2020), O_2-A (Dubisson *et al.*, 2009; Geddes and Boesch, 2015; Sanders *et al.*, 2015; Zeng *et al.*, 2020), and O_2-B (Ding *et al.*, 2016) bands, as well as combinations of these bands (Sanghavi *et al.*, 2012; Chen *et al.*, 2021). In addition, an algorithm for aerosol vertical information has been developed based on hyperspectral UV-visible radiance from satellite observation. Nanda *et al.* (2018) demonstrated the possibility of aerosol height retrieval from the O_2-A band developed an algorithm using Tropospheric

Monitoring Instrument (TROPOMI) (Sanders and de Haan, 2016; Nanda *et al.*, 2020) and implemented the algorithm operationally.

However, the vertical distribution of aerosol is difficult to assess because of its large spatio-temporal variability. Although the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) provided the aerosol vertical distribution with high vertical resolution (Omar *et al.*, 2009), other passive satellite sensors are only able to estimate the representative parameter of aerosol height. Veihelmann *et al.* (2007) showed that the number of degrees of freedom of signal for aerosol is 2~4 for most of satellite observation conditions by the ozone monitoring instrument (OMI). In addition, the number of degrees of freedom is not exceeded to 3 from the shortwave satellite measurements (e.g., Rao *et al.*, 2019; Choi *et al.*, 2021). It means that the amount of information for aerosol vertical distribution has a limitation for satellite sensor. Because of limitation for describing the aerosol vertical information, aerosol layer height (ALH) (Nanda *et al.*, 2018) or aerosol effective height (AEH) (Park *et al.*, 2016) were defined to retrieve the aerosol vertical information from the passive satellite sensors.

The Geostationary Environment Monitoring Spectrometer (GEMS), which was launched by South Korea in February 2020, retrieves data related to major trace gases and aerosol properties (Kim *et al.*, 2020). The main purpose of GEMS is to monitor air quality, and aerosol properties are targets of such monitoring over East Asia. For this reason, the GEMS aerosol algorithm was developed as multiple operational products. Aerosol properties are obtained for the purposes of monitoring surface air quality and aerosol effects for the air mass factor (AMF) calculation. In addition to the aerosol optical property algorithm, the GEMS aerosol product is applied to the aerosol vertical information, AEH. For the possibility for development of an AEH retrieval algorithm,

Park *et al.* (2016) conducted theoretical sensitivity testing of AEH retrieval using solely the O₂-O₂ absorption band along with aerosol and surface properties. Overall, the sensitivity of AEH retrieval was strongly affected by SSA, AOD, and aerosol types including optical and size properties, and the error budget for AEH retrieval using the O₂-O₂ band was 739 ~ 1276 m. In addition, case studies of AEH during dust transport over East Asia were conducted using radiance data from the Ozone Monitoring Instrument (OMI) and aerosol optical properties from the Moderate Resolution Imaging Spectroradiometer (MODIS).

Based on theoretical considerations and case results of previous studies, we introduce an operational retrieval algorithm for AEH. Section 2 introduces the details of satellite sensors for the comparison and collocation method in this study. Section 3 describes the details of the AEH retrieval algorithm for GEMS and provides a list of the detailed input parameters. Section 4 reports retrieval results based on case studies of aerosol transport, and section 5 contains long-term validation results based on CALIOP and TROPOMI data. Finally, in section 6, the summary and main conclusions are presented.

2. Data

2.1 GEMS

The GEMS is onboard the Geostationary Korea multipurpose satellite 2B (GK2B) as orbiting at 128.2°E, and scans from 145°E to 75°E with north-south coverage of 5°S~45°N. The GK2B observation schedule shares the GEMS and the Geostationary Ocean Color Imager 2 (GOCI2), and the GEMS scan the 30 minutes duration from every hour from 45 minutes to 15 minutes during daytime. The standard spatial resolution of GEMS is 7 km × 8 km. The spectral resolution and sampling are

respectively 0.6 nm with full-width and half-maximum (FWHM) and 0.2 nm with spectral range of 300~500 nm.

The GEMS Level 2 aerosol operational algorithm (L2AERAOD) retrieves the aerosol index (AI) values for UV and visible wavelengths, as well as AOD and SSA with considering the aerosol types (National Institute of Environmental Research, 2020a). The aerosol types are defined as absorbing, non-absorbing, and dust types by using the classification methods based on the UV and visible AIs (e.g., Go et al., 2020). Park *et al.* (2016) noted that the error budget of AEH is significantly affected by uncertainty in AOD and SSA and by the misclassification of aerosol types, which is directly related to the optical property and size information. Overall, the error for AEH is ranged from 739~1276 m under the AOD error of 0.2, particle size error of 20%, SSA error of 10%, and surface albedo error of 0.02 (Park et al., 2016). The main variables causing errors for AEH retrieval can be obtained from the L2AERAOD results. Therefore, the L2AERAOD results for AOD at 550 nm and SSA at 443 nm were adopted as input data for aerosol properties.

Although L2AERAOD retrieved their own surface reflectance for accurate separation of surface signals from total reflectance at the top of the atmosphere (TOA), the standard product for surface reflectance (L2SFC) (National Institute of Environmental Research, 2020b) was also independently retrieved from GEMS radiance/irradiance data with specific temporal periods. L2SFC is the reference product for spectral surface reflectance. The L2SFC retrieves the surface reflectivity in multiple spectral channels and provides the black surface reflectivity (BSR) and bi-directional reflectance distribution function (BRDF) based on the original pixel resolution. Recently, L2SFC accurately estimated surface reflectance in near real time in operation. For this reason,

L2SFC was used as reference data for the surface products for the air mass factor estimation to all trace gas retrieval algorithms. Similarly, the AEH retrieval algorithm also uses L2SFC as a reference surface reflectance in operation. Specifically, the BSR value at 477 nm is used as the surface reflectance input for AEH retrieval. However, this study used the minimum reflectance under the Lambertian assumption to retrieve AOD and AEH to coincide with the use of surface information on L2AERAOD and AEH retrieval.

2.2. TROPOMI

TROPOMI is a nadir-viewing spectrometer, the only payload of the Sentinel-5 Precursor (S5P), measuring radiance in the UV, visible, near-infrared, and the shortwave IR (Veefkind et al., 2012). The S5P crosses the equator at 13:30 local time in a polar orbit with ascending node providing near-global daily coverage. The aerosol layer height product from TROPOMI (AER_LH) retrieves vertically localized aerosol layers in free troposphere with cloud free condition by using the level 1b earth radiance measurements from 758 to 770 nm (de Graaf *et al.*, 2022). The definition of ALH from TROPOMI is the optical centroid layer height of the plume above sea level. Spectral fit estimation of reflectance around the O₂-A band is based on a neural network for the forward model calculation for simulated condition. After cloud masking to avoid the cloud affected pixels, an optimal estimation method was used to retrieve the aerosol layer height parameters for the inversion method from observation. During the radiance fitting, the ALH and AOD are fitted parameters, but other aerosol parameters, such as SSA, layer thickness, and scattering phase function, are assumed to be fixed values (Nanda et al., 2020). Furthermore, the ALH retrieval has limitation to the aerosol plume

with higher than 12 km, because the ALH neural network method is currently adopted to the plume pressure range of 75~1000 hPa (Michailidis et al., 2023).

Main purpose of the AER_LH product is the retrieval of aerosol layers in the free troposphere (desert dust, biomass burning, and volcanic ash) (Michailidis et al., 2023). The target requirement on the accuracy and precision is 0.5 km or 50 hPa, and the threshold requirement is 1 km or 100 hPa under the elevated aerosol plumes with cloud-free conditions (de Graaf et al., 2022, Veefkind et al., 2012). However, the TROPOMI ALH product has strong dependence of the surface albedo, especial to the bright surfaces (Sanders et al., 2015). From Michailidis et al. (2023), a mean bias of -0.51 ± 0.77 and -2.27 ± 1.17 km is estimated over ocean and land, respectively. In this study, we use version 02.04.00 of the TROPOMI offline level 2 AER_LH product (European Space Agency, 2021) with the spatial resolution is $3.5 \text{ km} \times 5.5 \text{ km}$ at nadir viewing geometry.

2.3. CALIOP

The CALIOP is a spaceborne lidar onboard the Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) to measure the vertical information of aerosol and cloud with estimating the optical properties. The CALIOP has two different wavelength channels (532 and 1064 nm) by using the Nd:YAG laser to generate the signals (Winker et al., 2009). The orbit for CALIPSO is Sun synchronous orbit constellated to the A-train with period of 98.3 minutes by ascending node. It crosses the equator at 13:30 local time. To cover the GEMS observation area, we used CALIOP data of 3~4 orbits per day. For the vertical information, the resolution for vertical sampling is 30 m below 8 km altitude, and 60 m from 8 to 20 km altitude, respectively.

Although the CALIOP retrieves the data with extremely high horizontal and vertical resolutions, the spatial coverage is narrow because the footprint of the CALIOP is about 90m at the Earth surface. In this study, the data of Level 2 aerosol profile product (APro, version 3.41) was used (Tackett et al., 2018). The aerosol profile product simultaneously includes both aerosol and cloud optical depth. In addition, extinction quality flag (Extinction_QC_Flag) data shows the quality of extinction profile. In this study, we checked and used to the quality flag value at 532 nm for the aerosol extinction coefficient profile data. The AOD from CALIOP is vertically integrated aerosol extinction coefficient from surface to top of atmosphere, and representative layer height parameters (ALH and AEH) are directly estimated by using the vertical profile of aerosol extinction coefficient at 532 nm to minimize the spectral discrepancy of aerosol extinction. For the ALH retrieval, we adopted mean extinction height from the CALIOP extinction coefficient profile (e.g., Koffi et al., 2012; Xu et al., 2019). Furthermore, the AEH adopted the same method as CALIOP for estimating the ALH, with only changes in the height definition.

2.4 Data Selection and Colocation

In this study, we used the data within 75°E~145°E for east-west and 5°S~45°N for north-south direction, which is observation area of GEMS. For spatial colocation, we selected pixels for which distance between GEMS and CALIOP (or TROPOMI) observations was less than 50 km. From Park et al. (2020), the spatial scales for AOD validation are 30~40 km. To ensure that the number of retrieved observation pixels, we relax the spatial limits for collocation matching. In addition, only the closest 10% of pixels were used. Given the different orbital characteristics of CALIOP (or TROPOMI)

and GEMS, temporal colocation was also considered. During the period of image scanning from east to west over Asia by GEMS, CALIOP and TROPOMI pass through the GEMS observation area from south to north every 98.3 minutes. On average, two low earth orbit (LEO) satellites pass three to four orbits through the GEMS scan area during a single day of daytime observation. To consider these different orbital characteristics, only observations taken within ± 1 hour of the GEMS observation time was selected for temporal colocation. As GEMS observes hourly, collocated pixels between the two satellites shift from east to west over time.

To ensure the accuracy of ALH from TROPOMI, only pixels with quality assurance (QA) values of 1.0 were used. To minimize the cloud contamination, the TROPOMI ALH product uses the VIIRS cloud mask information and cloud parameters from the Fast Retrieval Scheme for Clouds from the Oxygen A-band (FRESCO). To consider the cloud contamination for the aerosol products, in addition, the VIIRS cirrus cloud reflectance ($\text{viirs_cirrus_reflectance} < 0.4$), VIIRS cloud mask ($\text{viirs_cloud_mask} < 0.1$), and cloud fraction from the FRESCO ($\text{cloud_fraction} < 0.1$) are considered in this study (Michailidis et al., 2023). However, de Graaf *et al.* (2022) showed that respective cloud masking method have difficulty detecting various clouds. For this reason, accuracy problem of ALH by the cloud contamination is still remained. From the previous studies, the UVAI is used as the threshold to define the absorbing aerosol pixels (e.g., Chen et al., 2021; Griffin et al., 2020; Michailidis et al., 2023; Sanders et al., 2015). However, the GEMS aerosol product is retrieved not only the absorbing aerosols, but also the non-absorbing aerosols. For this reason, the UVAI is not used to the threshold of aerosol pixel identification. Instead, we adopted the larger than 0.3 for AOD at 443 nm from GEMS to identify both absorbing and non-absorbing aerosol loading.

3. AEH retrieval algorithm

AEH is a layer height parameter that considers the penetration of photons into the aerosol layer. In this study, the AEH product from GEMS is defined as the height with aerosol extinction integrated from the surface of $(1-e^{-1}) \times \text{AOD}$, and a detailed definition of AEH was introduced by Park *et al.* (2016). Numerous previous studies have used the aerosol top layer height (Kohkanovsky and Rozanov, 2010) or middle layer height (i.e. ALH or centroid height) (e.g., Sanders *et al.*, 2015; Nanda *et al.*, 2020) as the aerosol vertical layer parameter. AEH is similar to the aerosol top layer height but with a slight bias.

For AEH retrieval, the vertical distribution assumption is also important. The Gaussian Density Fitting (GDF) distribution, which is a modified Gaussian distribution structure, is assumed for AEH retrieval. The full-width at half-maximum (FWHM) of the aerosol layer is 1 km. Schematic description of AEH and other aerosol vertical parameters are shown in Figure 1. Based on the assumptions about the aerosol vertical distribution, the AEH value is higher than the peak height of the Gaussian distribution and lower than the aerosol top layer height. Otherwise, aerosol layer height (ALH) in this study is defined as the height integrated aerosol extinction from the surface reaching half of AOD (i.e., $0.5 \times \text{AOD}$). Therefore, the ALH is same to the peak height for the vertical profile condition as shown in Figure 1.

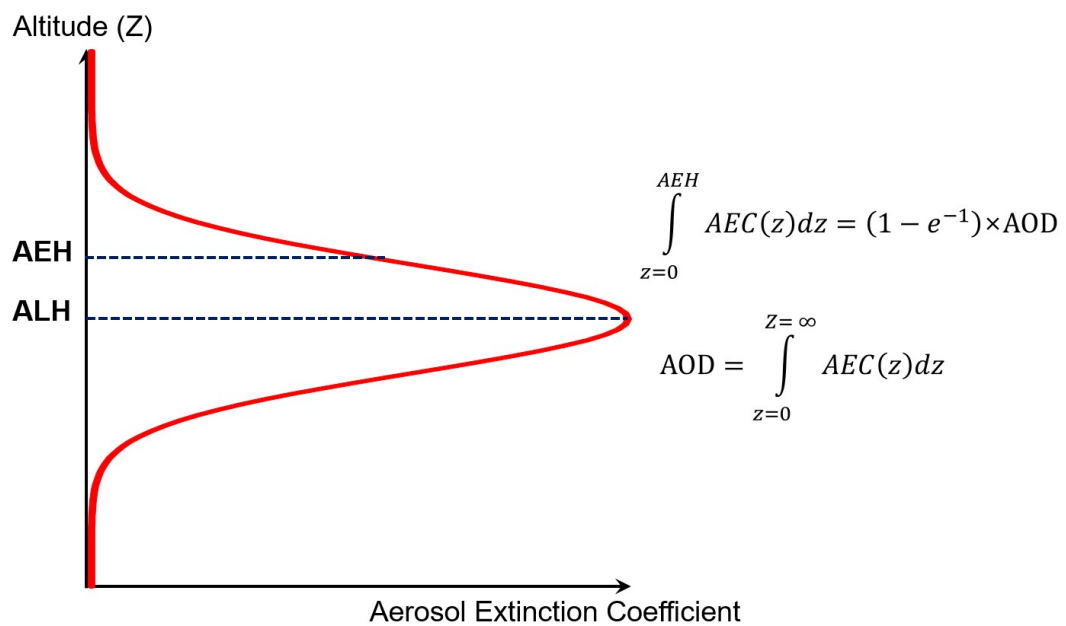


Figure 1. A schematic illustration of AEH and ALH definitions in an idealized Gaussian shape of aerosol vertical distribution.

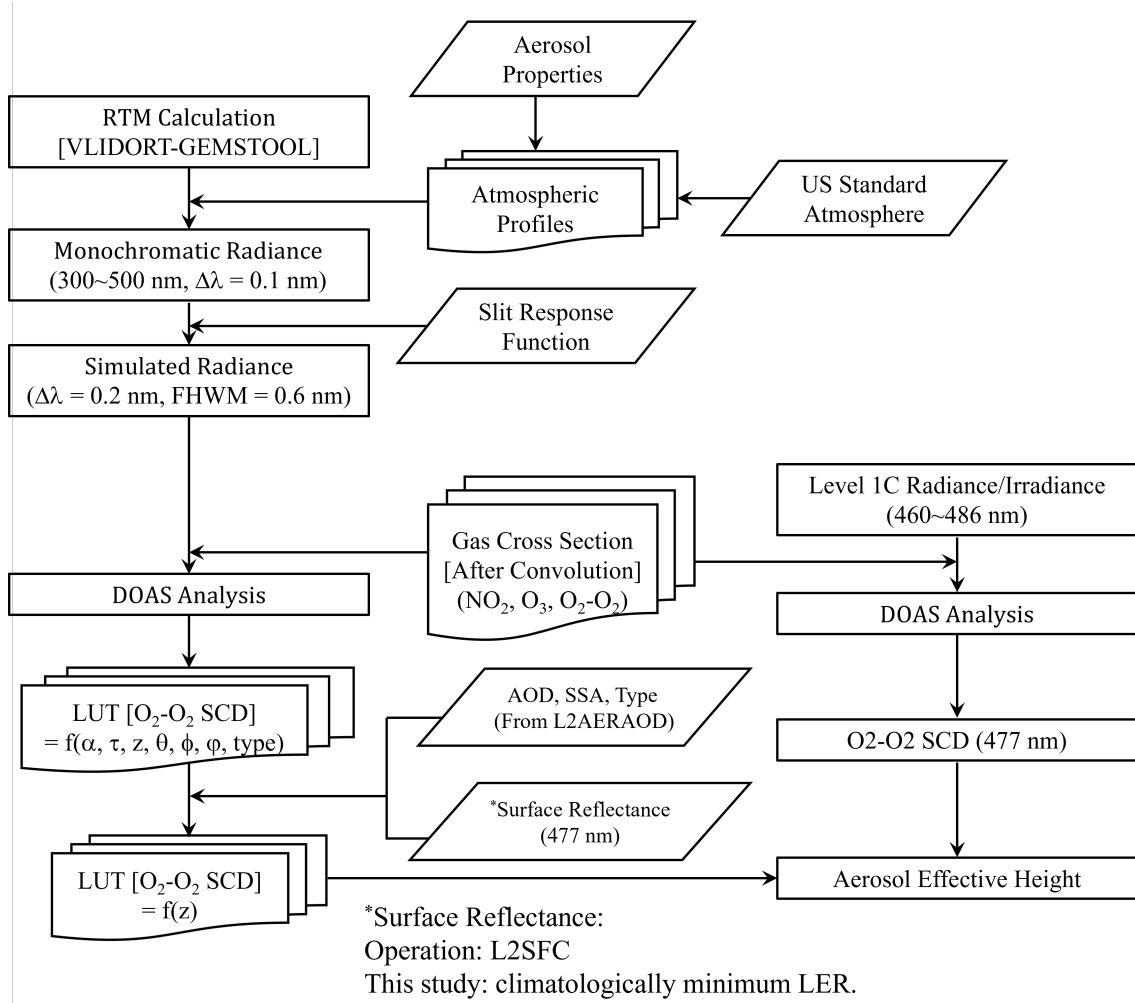


Figure 2. Flowchart of the AEH retrieval algorithm for GEMS satellite observation.

Figure 2 shows the overall flowchart of the AEH algorithm for GEMS satellite. The AEH algorithm for GEMS composed a look-up table (LUT) development between AEH and O₂-O₂ SCD by the radiance simulation and SCD estimation from the satellite radiance. The LUT is a function of observation geometries [solar zenith angle (SZA; θ), viewing zenith angle (VZA; Φ), relative azimuth angle (RAA; ϕ)], surface altitude (z), surface albedo (α), AOD (τ), and aerosol type. During the radiance simulation, the radiance is monochromatically simulated and simulated monochromatic radiance is

convolved as considering the spectral response of GEMS instruments. For AEH estimation, the radiance information is finally converted to the AEH values by using the differential optical absorption spectroscopy (DOAS) method. DOAS method is identification technique for the spectral absorption signals from radiance information and detailed principle and information is explained by Platt (1994). DOAS method has been frequently used to estimate the amount of trace gases (i.e., SCD of trace gas) from ground (e.g., Cheng et al., 2023; Irie et al., 2008; Platt and Stutz, 2008; Wagner et al., 2011; Wang et al., 2017) and satellite (e.g., Kwon et al., 2019; Li et al., 2023; Wagner et al., 2007, 2010) measurements.

For AEH retrieval, the basic method is the identification of changes in optical path length caused by effective aerosol layer height variation. To measure the optical path length change, O₂-O₂ slant column density (SCD) retrieved by the DOAS method was used because the spectral coverage is limited to 300-500 nm (Park et al., 2016, Kim et al., 2020). In the GEMS product, the O₂-O₂ SCD at 477 nm absorption band is most useful absorption band because this absorption band is strongest absorption band within the GEMS spectral observation range. Detailed DOAS fitting parameter and setting information is provided in Table 1 for the estimation of O₂-O₂ SCD from both the simulation and observation data. For the O₂-O₂ SCD estimation at 477 nm, the fitting window is ranged from 460 to 486 nm to cover the full absorption structure of O₂-O₂. Within the fitting window, the absorptions of NO₂ and O₃ is significant. To describe these two absorbing species, temperature dependent cross section information are adopted. The temperature dependent cross section setting considers the stratosphere and troposphere, simultaneously.

Table 1. Details of fitting parameter for O₂-O₂ SCD estimation via the DOAS method.

Parameter	
Fitting window	460 – 486 nm
Absorption	NO ₂ at 220 and 294 K (Vandaele <i>et al.</i> , 1998)
cross section	O ₃ at 223, 243 and 293K (Bogumil <i>et al.</i> , 2001)
	O ₂ -O ₂ at 293 K (Thalman and Volkamer, 2013)
	Ring

Table 2. Ratio between SCD error and the SCD of O₂-O₂ according to the polynomial order and offset settings used for DOAS fitting.

Polynomial	Offset = none	Offset = 0 th
2 nd order	6.06 ± 2.07	6.79 ± 2.31
3 rd order	6.32 ± 2.20	6.79 ± 2.32
4 th order	7.86 ± 2.78	7.34 ± 2.85

To minimize the noise effect and improve DOAS fitting quality, the optimal settings for fitting were also analyzed. Table 2 shows ratios of O₂-O₂ SCD error to the O₂-O₂ SCD for various polynomial and bias orders from observed radiance. The polynomial and offset are basic fitting parameters for the DOAS fitting. Two parameters describe the broadband spectral feature of radiance before identifying the gas absorption structure. The ratio between SCD error and the SCD of O₂-O₂ is important to determine the AEH retrieval quality. When the fitting error increase, the uncertainty of AEH is also enhanced during the retrieval. For this reason, the setting with 2nd order of polynomial and none offset was used for the O₂-O₂ SCD estimation from the GEMS radiance due to the smallest fitting error.

In AEH estimation, other aerosol characteristics, including aerosol load and optical properties, affect retrieval accuracy. From Park et al. (2016), uncertainty of AEH retrieval result is largest by the SSA uncertainty. In addition, the AEH retrieval uncertainty by the aerosol optical properties (e.g., AOD and phase function) and surface albedo has dependence of observation geometries. After the estimation of O₂-O₂ SCD, for this reason, conversion from O₂-O₂ SCD to AEH is an essential process. Table 3 shows the dimension of the LUT for the AEH retrieval algorithm. To calculate the LUT, a linearized pseudo-spherical vector discrete ordinate radiative transfer model (VLIDORT) version 2.6 was used (Spurr, 2013). During the radiative transfer model simulation, reference wavelength for the SSA and AOD is assumed to be 440 nm. The aerosol type is considered by the radiative absorptivity and size information, which is based on the method from Lee et al. (2010). Based on the Lee et al. (2010), the aerosol type is classified to absorbing, dust, and non-absorbing aerosol. Absorbing and non-absorbing aerosol types are assumed to the fine-mode dominant particles. For the spectral conversion of AOD, the angstrom exponent of 1.186, 0.222, and 1.179 are used for absorbing, dust, and non-absorbing aerosol, respectively. Otherwise, the SSA is assumed as the fixed value within the spectral range for O₂-O₂ estimation. Although the center of O₂-O₂ absorption is 477 nm, the spectral discrepancy between model assumed wavelength and center wavelength of O₂-O₂ absorption is assumed to be ignored in this study. After calculating spectral radiance with 0.1 nm sampling, we performed the slit response function of GEMS and sampling specification prior to the DOAS fitting. For O₂-O₂ absorption, the absorption cross section used for the radiative transfer model calculation is considered the temperature dependent absorption cross section (e.g., Park et al., 2017).

Table 3. The dimension of the LUT for the GEMS AEH retrieval algorithm used to estimate AEH from O₂-O₂ SCD. (SZA: solar zenith angle, VZA: viewing zenith angle, RAA: relative azimuth angle, SUR: surface reflectance).

Variable [unit]	No. of entries	Entries	
Spectral range [nm]	-	455~491 nm (0.1 nm interval)	
SZA [°]	7	0.01, 10, 20, 30, 40, 50, 60	
VZA [°]	7	0.01, 10, 20, 30, 40, 50, 60	
RAA [°]	10	0.01, 20, 40, 60, 80, 100, 120, 140, 160, 180	
SUR	3	0.0, 0.05, 0.2	
AOD at 440 nm	11	0.04, 0.2, 0.4, 0.7, 1.0, 1.3, 1.6, 2.0, 2.5, 3.0, 5.0	
Refractive Index (Imaginary) at 440 nm	3×3	Absorbing (Real: 1.45)	0.000, 0.0074, 0.0314
		Dust (Real: 1.53)	0.0, 0.0030, 0.0080
		Non-Absorbing (Real: 1.41)	0.0, 0.0040, 0.0156
AEH [km]	13	0.0 (Extrapolate), 0.2, 0.5, 1.0, 1.3, 1.6, 2.0, 2.3, 2.7, 3.0, 3.5, 5.0, 10.0 (Extrapolate)	
Terrain Height [km]	2	0.0, 2.0	

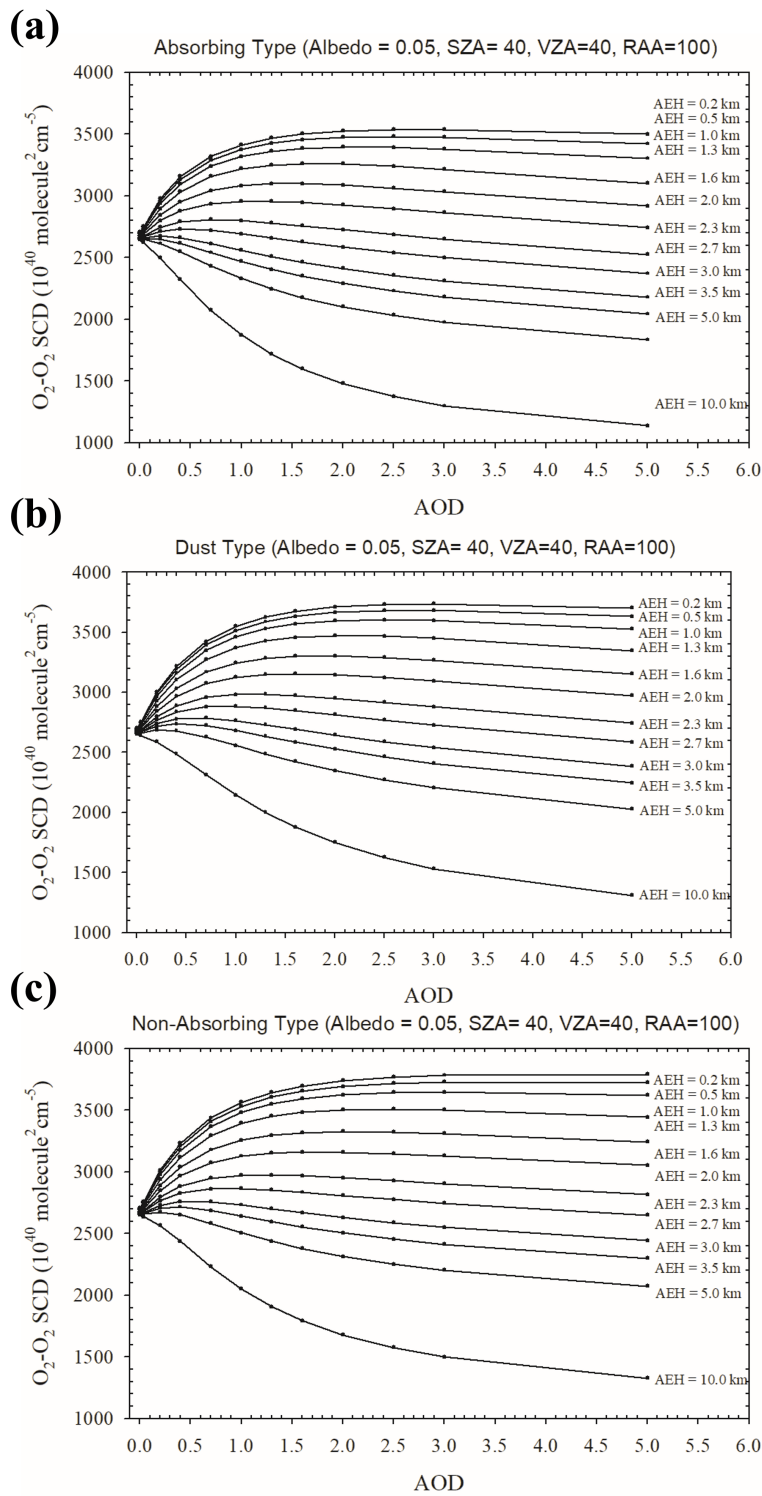


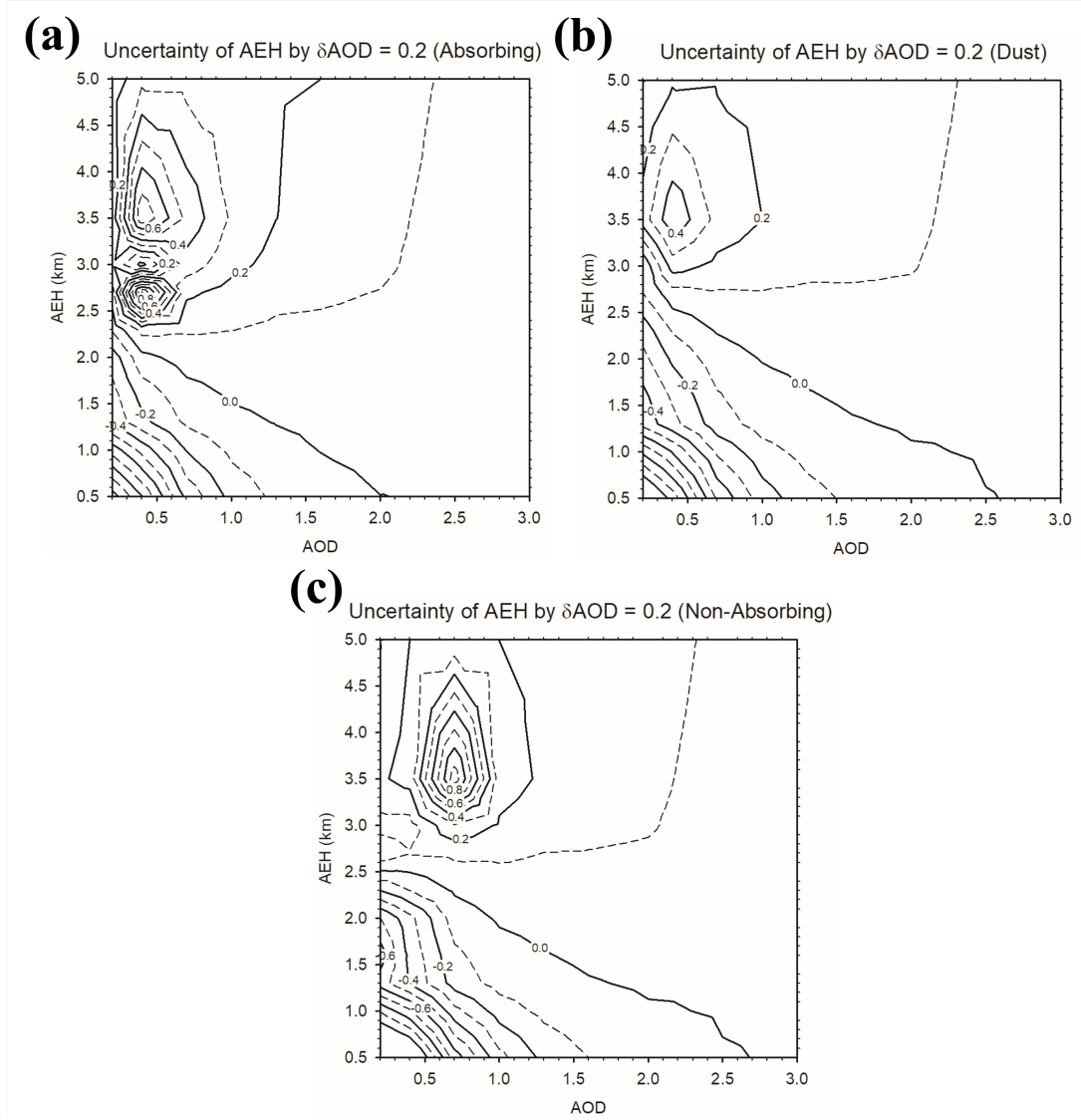
Figure 3. Example of LUT to retrieve the AEH according to (a) Absorbing, (b) Dust, and (c) Non-Absorbing aerosol types.

Figure 3 shows the example of LUT to retrieve the AEH from O₂-O₂ SCD according to the respective aerosol types and AOD. O₂-O₂ SCD decreases with increasing AEH for all aerosol types and AOD (Park *et al.*, 2016). Similar to the previous study, the O₂-O₂ SCD sensitivity is enhanced at high AOD and absorbing aerosol cases from GEMS LUT. In addition, the contrast of O₂-O₂ SCD is greater for absorbing aerosols than non-absorbing aerosols. During the radiance passing through the aerosol layer, the absorbing aerosol is more efficiently absorbed the radiance. For this reason, the effective optical path length is significantly shorter for absorbing aerosols.

Although the surface reflectance and AOD are retrieved by the GEMS operational products, retrieved values are always included the uncertainties. To consider the retrieval error, the retrieval uncertainty of AEH was evaluated as the relative ratio of SCD changes with respect to input variables. This ratio is defined by the relationship between changes in SCD per unit AEH and changes in SCD due to uncertainties in input variables. Figure 4 shows the AEH retrieval uncertainty based on the O₂-O₂ SCD LUT with an AOD error of 0.2. The AOD error of 0.2 is twice the target accuracy of the GEMS standard algorithm product. Because the O₂-O₂ SCD change is small in high AOD, the AEH retrieval uncertainty is relatively small in high AOD condition. In addition, the SCD sensitivity in high AEH is weak due to the vertical distribution characteristics of O₂-O₂. Thus, the AEH uncertainty is up to 0.9 km at high AEH case in moderate AOD cases (around 0.5~1.0).

Figure 5 shows the AEH retrieval uncertainty with the surface reflectance error of 0.02. Because surface reflection and aerosol scattering simultaneously affect the increase in optical path length, surface reflectance uncertainty have a significant impact especial to low AOD. The AEH retrieval error due to uncertainty in surface reflectance

384 is larger than 1 km in $AOD < 0.4$. In addition, the AEH error by the surface reflectance
 385 uncertainty linearly decreases with increasing AOD, indicating relatively small impact
 386 with aerosol height change.



387

388 **Figure 4.** AEH retrieval uncertainty by the AOD error of 0.2 according to (a)
 389 Absorbing, (b) Dust, and (c) Non-Absorbing aerosol types.

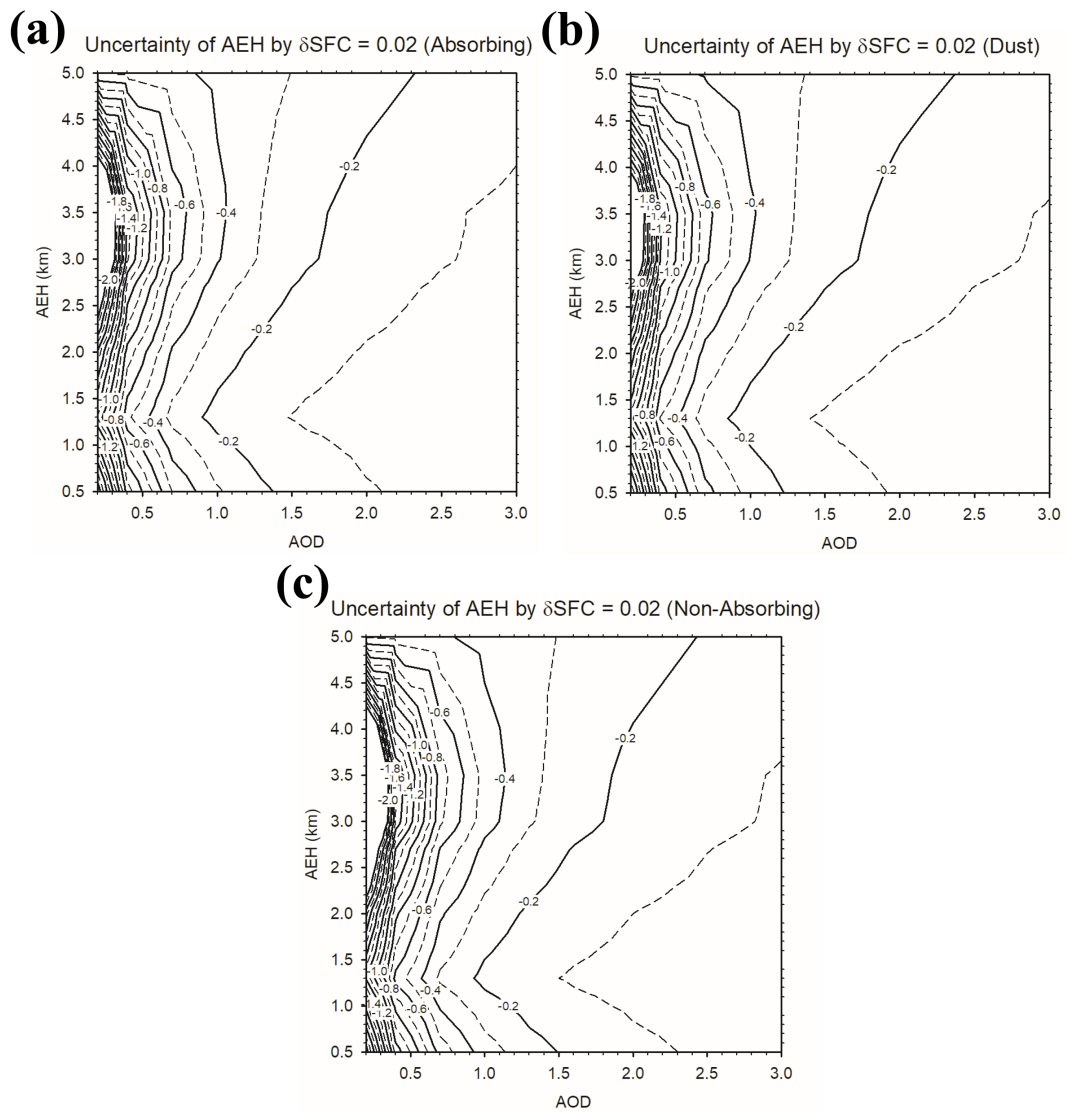


Figure 5. AEH retrieval uncertainty by the surface reflectance error of 0.02 according to (a) Absorbing, (b) Dust, and (c) Non-Absorbing aerosol types.

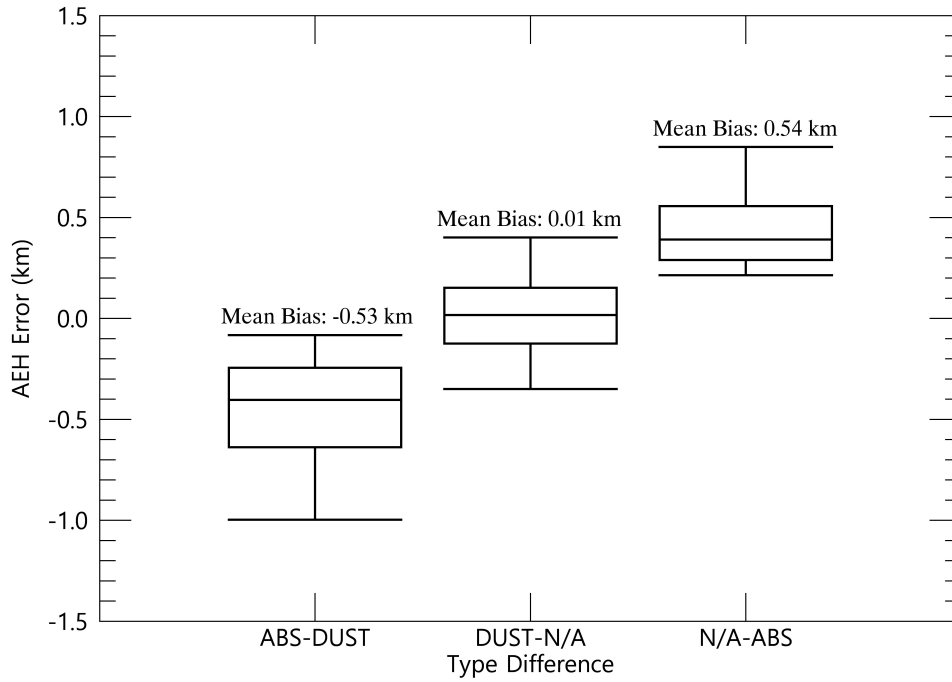


Figure 6. AEH retrieval uncertainty caused by the aerosol type misclassification.

Figure 6 shows the retrieval uncertainty caused by misclassification of the aerosol types. On average, the AEH retrieval uncertainty has an impact of up to 0.5 km, but this uncertainty shows larger than 2 km uncertainty under low AOD and short optical path length conditions. Based on the changes in sensitivity observed for optical path length, aerosol type (in terms of SSA) and AOD, and surface reflectance are carefully considered as input parameters for AEH retrieval. In the AEH retrieval, the AOD, aerosol type, and SSA are obtained from the L2AERAOD, which is standard aerosol product of GEMS. As shown in Section 2.1, the L2SFC product is used in operation, but this study used the climatological minimum Lambertian surface reflectance. However, the aerosol vertical distribution is always fixed as GDF function profiles as shown in Spurr and Christi (2014). The data quality of AEH retrieval results are considered the qualities of aerosol optical property and DOAS fitting results.

4. Case studies

Figure 7 shows retrieval results for AEH from GEMS on March 29 over East Asia. Because the operational schedule is hourly during the daytime, the GEMS retrieval results are shown at 1-hour intervals from 01:00 to 07:00 Universal Time Coordinated (UTC). AOD and SSA are also shown in Figures S1 and S2, respectively. From Park et al. (2016), pixels with low AOD values have large AEH uncertainty due to weak aerosol scattering information. For this reason, only AEH retrieval results with AOD greater than 0.3 are shown in this study. In addition, the AEH retrieval results are also adopted to the quality flag for “Reliable” from L2AERAOD, which considers the significant averaging kernel for optimal estimation for aerosol property retrievals. During this case study, a dust plume was located along the coast of China and South Korea with AOD at 443 nm of 0.8~1.2. Simultaneously, another plume was also present over the northeastern Korean Peninsula with AOD of 1.0~2.0 at 443 nm. SSA at 443 nm was 0.90~0.93 for the plume over South Korea and 0.87~0.90 for the plume over the northeastern Korean Peninsula. Retrieved AEH results from these different plumes show similar ranges. For both detected plumes, the AEH shows similar pattern ranging between 1.0 to 2.0 km in this case.

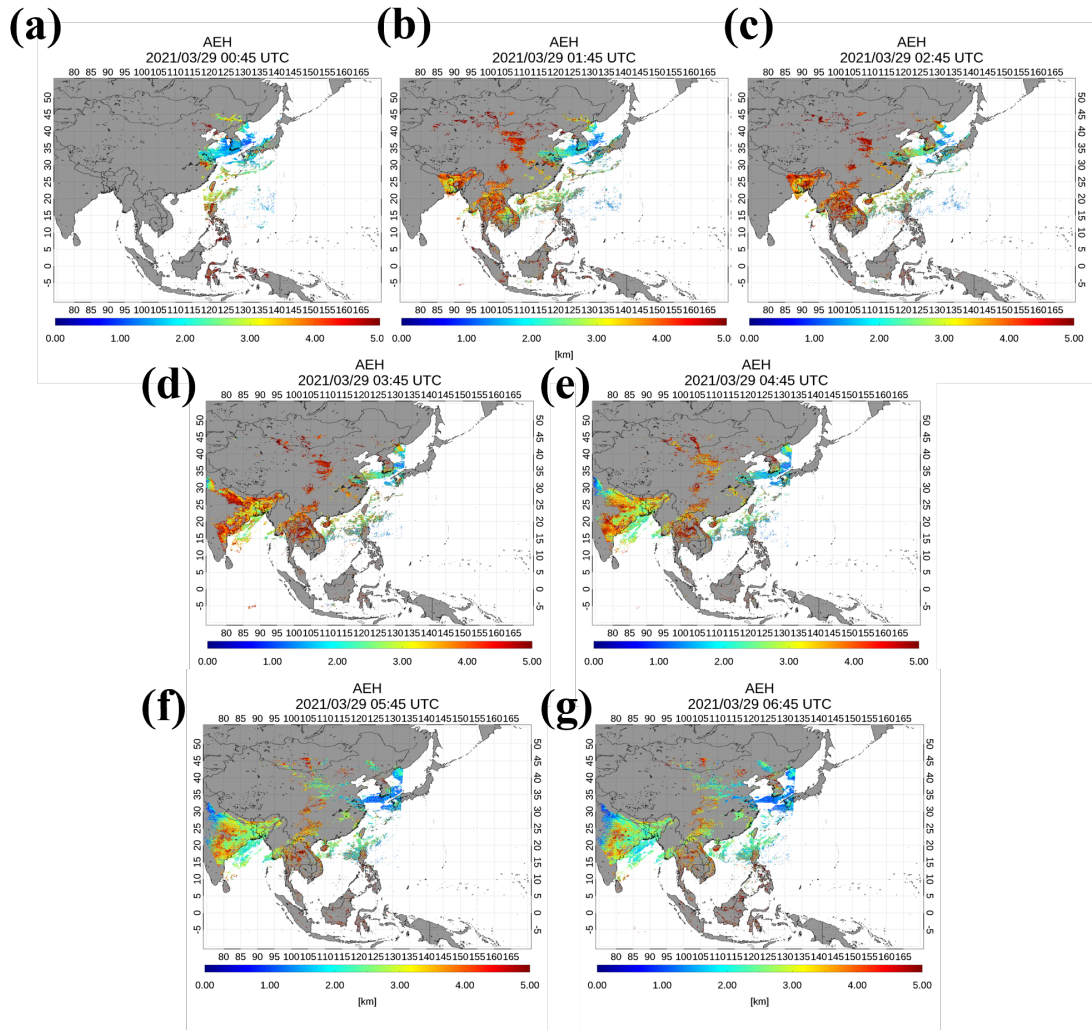


Figure 7. Case study results for AEH based on GEMS observations on March 29, 2021 [(a) 00:45 UTC, (b) 01:45 UTC, (c) 02:45 UTC, (d) 03:45 UTC, (e) 04:45 UTC, (f) 05:45 UTC, and (g) 06:45 UTC].

An additional severe aerosol plume was present over northeastern India, with AOD at 443 nm of 1.0~2.0 and SSA at 443 nm of 0.85~0.90. From Rana *et al.* (2019), metropolitan cities and industrial cluster in India are heavy emitters of black carbon, and high concentrations of black carbon are distributed over the Indo-Gangetic Plain (IGP). Therefore, the aerosol plume with high AOD and low SSA (high absorbing) was a result that actually exists, and it was not a result with high uncertainty due to edge of GEMS

observation field. Except for the inland parts of India, AEH in high AOD pixels ranged from 1.5 to 3.5 km.

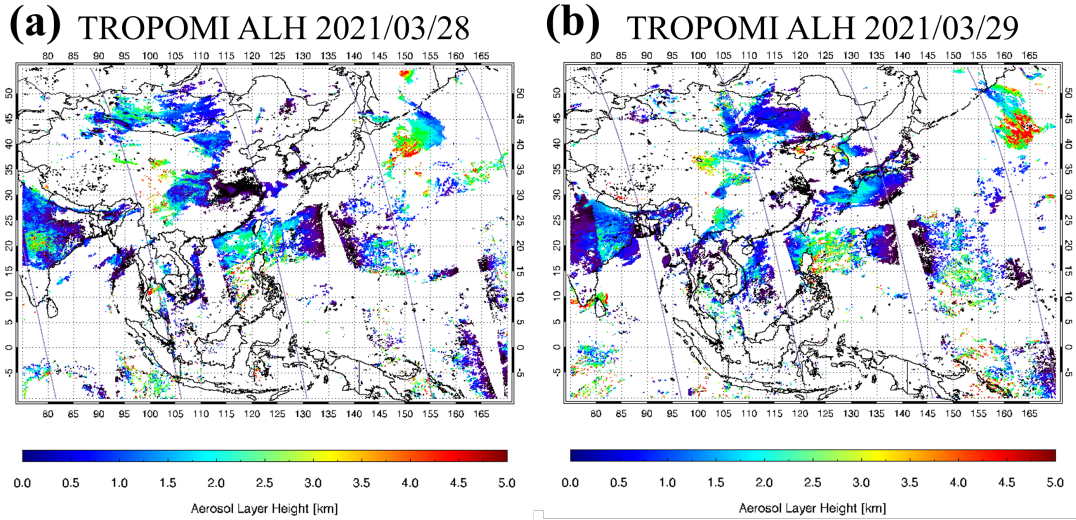
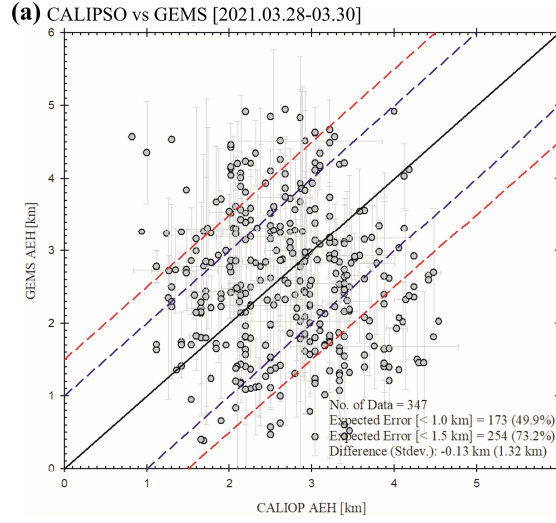
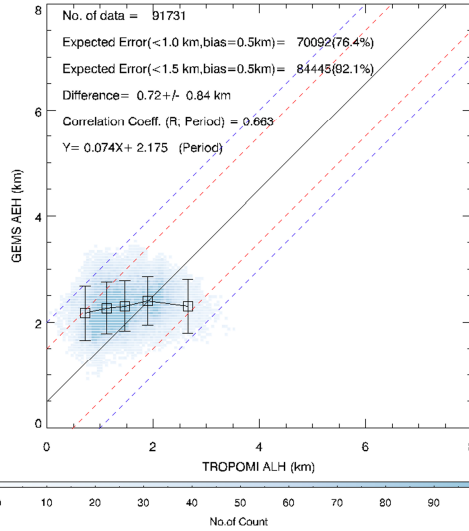


Figure 8. ALH retrieved from TROPOMI and orbit path of CALIOP on (a) March 28 and (b) March 29, 2021 (Unit: km).

For comparison of the retrieval, Figure 8 shows the ALH retrieved from TROPOMI on March 28 and 29, 2021 over East Asia. A dust plume was transported from China to South Korea during this period, then split into two distinct plumes over northeastern China and the coastal area of South Korea. The ALH retrieved from TROPOMI for both plumes were 0.5~1.5 km. Given the difference in definition for the aerosol height parameters between ALH and AEH, relatively high height values were retrieved from GEMS compared to TROPOMI. In an ideal case under symmetric gaussian distribution with a width of 1 km, the AEH from GEMS was around 0.5 km higher than the peak height of aerosol layer. The ALH expresses the center (or peak) height, thus, the AEH from GEMS was overestimated by around 0.5 km relative to the ALH from TROPOMI. Although AEH had higher values than ALH from TROPOMI, the GEMS AEH retrievals for the dust transport case study were successfully retrieved.



(b) TROPOMI vs GEMS [2021.03.28-03.30], Ocean



(c) TROPOMI vs GEMS [2021.03.28-03.30], Land

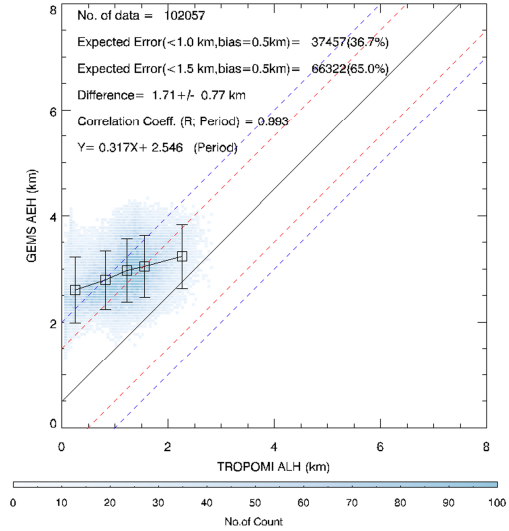


Figure 9. Intercomparison of (a) AEH between CALIOP and GEMS and (b) ALH from TROPOMI and AEH from GEMS over ocean and (c) over land (Black box and error bar is mean and standard deviation in 20% interval of each TROPOMI ALH) over the period from March 28 to 30, 2021.

Figure 9 shows intercomparison results for aerosol plume height among GEMS, CALIOP, and TROPOMI during the case study of dust transport in East Asia from March 28 to 30, 2021. For the direct comparison shown in Figure 9a, the difference in AEH between GEMS and CALIOP was -0.13 ± 1.32 km. Nanda *et al.* (2020) reported

that the difference in ALH between TROPOMI and CALIOP was 0.53 km for 4 cases of thick Saharan dust plumes. In addition, 49.9% and 73.2% of the total pixels showed differences less than 1.0 and 1.5 km, respectively. Large AEH uncertainty occurred mostly over the inland area of China. Because AEH from GEMS uses only the O₂-O₂ absorption band, the accuracy of AEH is sensitive to uncertainty in surface reflectance and AOD. From Park et al. (2016), total error budget of AEH is 0.74~1.28 km, and the total error budget considered the uncertainty of AOD, SSA, aerosol particle size, and surface albedo in the aerosol retrieval process. The total error budget amount from the previous study is similar value of standard deviation of AEH difference between GEMS and CALIOP.

Figures 9b and 9c illustrate the comparison results between GEMS AEH and TROPOMI ALH, for the period of March 28~30, 2021 over land and ocean surface, respectively. The difference between GEMS AEH and TROPOMI ALH was 0.72 ± 0.84 km and 1.71 ± 0.77 km over ocean and land in this case, respectively. In addition, 82.4% and 37.3% of all pixels had differences less than 1.5 km over ocean and land, respectively. However, the ALH from TROPOMI is generally lower than the AEH from GEMS because of the discrepancy in definitions. Based on the assumption of aerosol vertical distribution for AEH retrieval, the difference between AEH and center height of aerosol extinction profile is around 0.5 km. To consider the inconsistency of definition between ALH and AEH, the difference between two retrieval results decreased to 0.5 km bias. After consideration of definition inconsistency, the proportion of pixels within the expected error ranges of 1.5 km are enhanced to 92.1% and 65.0% over ocean and land, respectively.

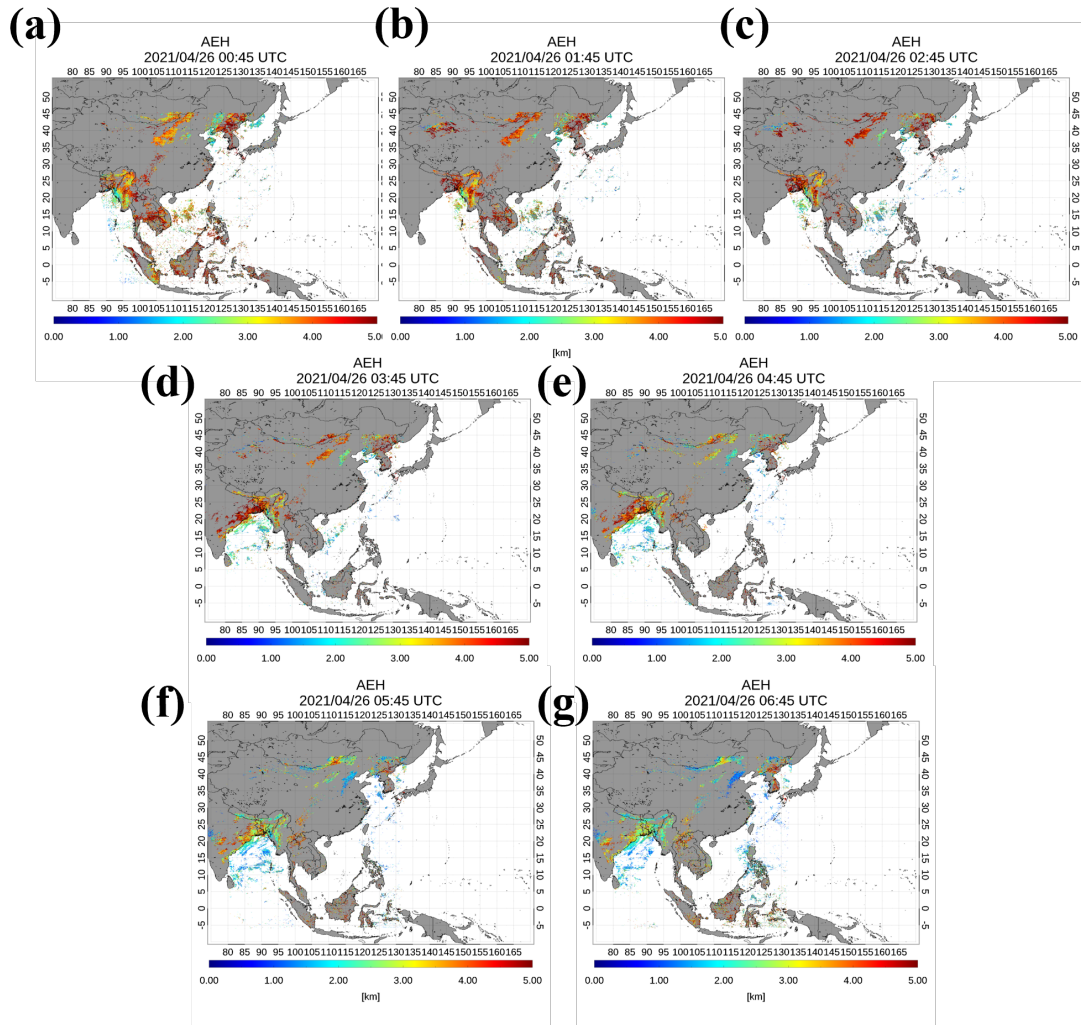


Figure 10. Case study results for AEH based on GEMS observations on April 26, 2021 [(a) 00:45 UTC, (b) 01:45 UTC, (c) 02:45 UTC, (d) 03:45 UTC, (e) 04:45 UTC, (f) 05:45 UTC, and (g) 06:45 UTC].

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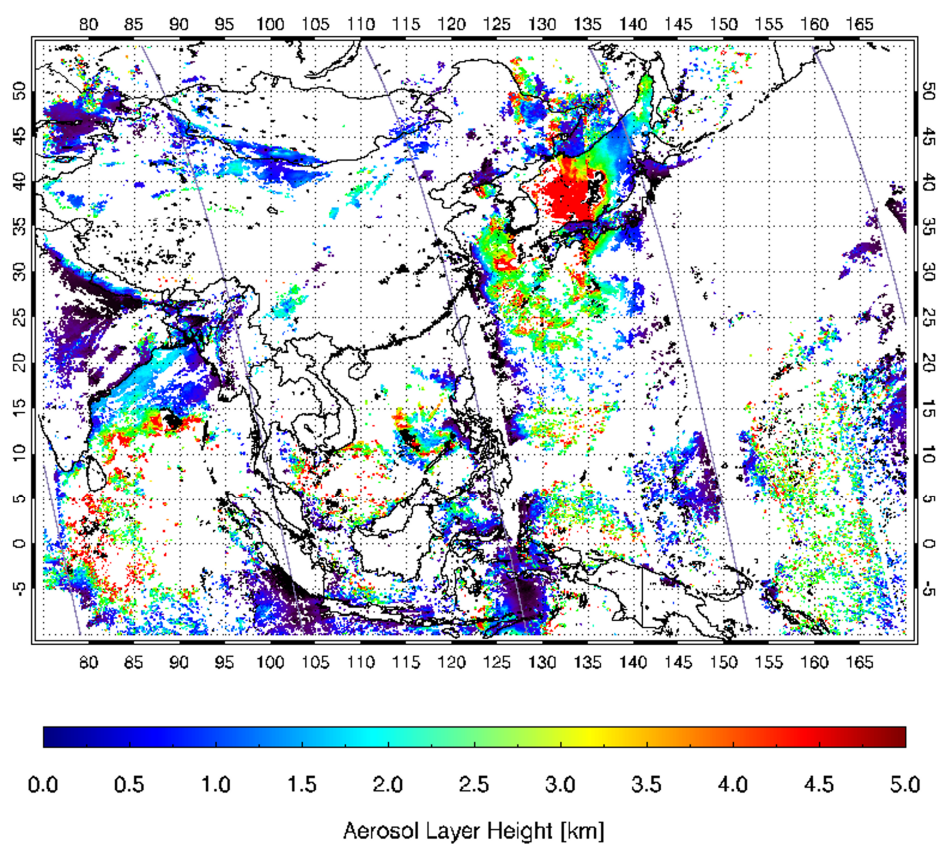


Figure 11. ALH retrieved from TROPOMI and orbit path of CALIOP on April 26, 2021 (Unit: km).

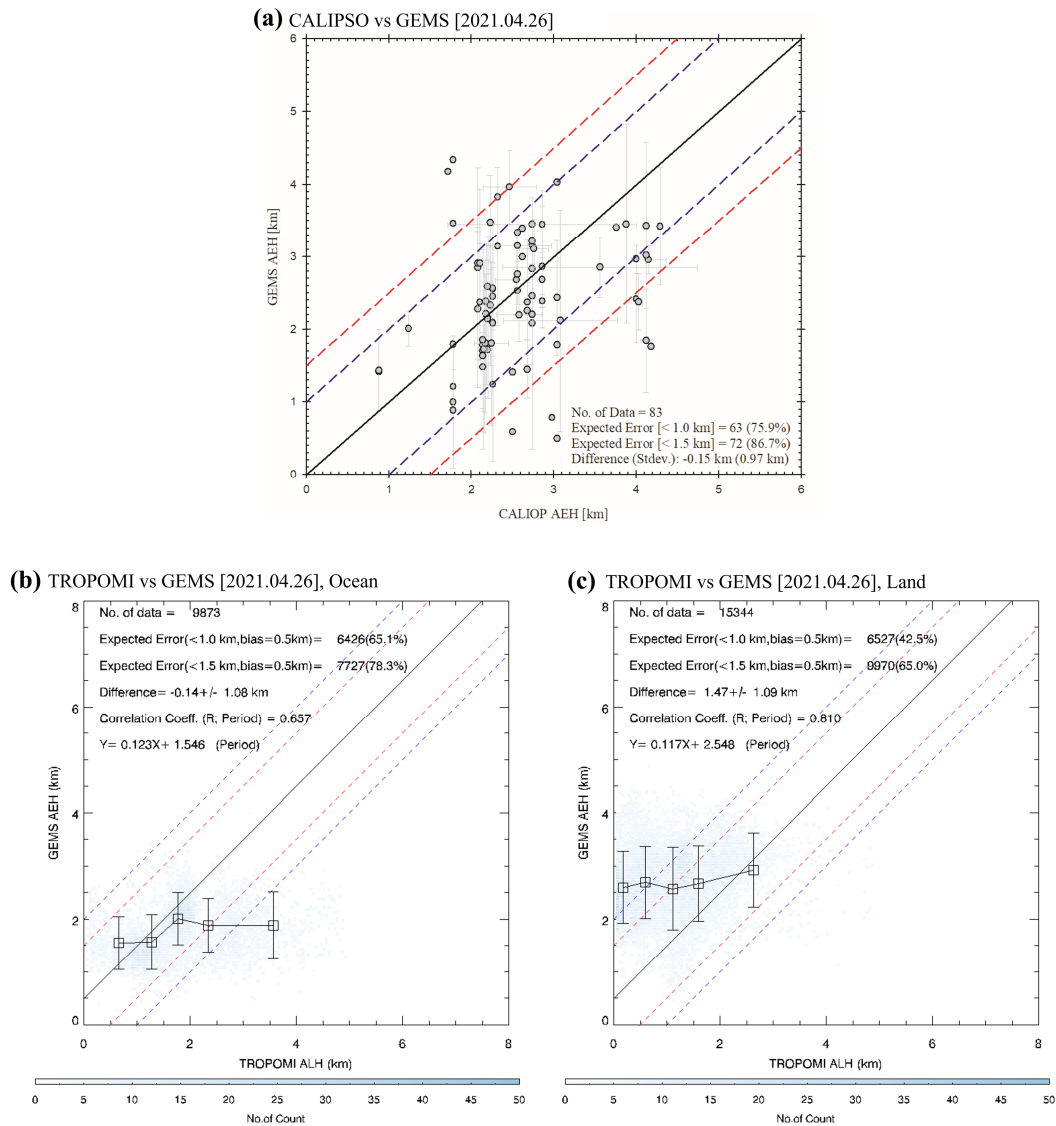


Figure 12. Intercomparison of (a) AEH between CALIOP and GEMS, and (b) ALH from TROPOMI and AEH from GEMS over ocean and (c) over land (black dot and error bar is mean and standard deviation in 20% interval of each TROPOMI ALH) on April 26, 2021.

An additional intercomparison case of April 26, 2021, is shown in Figures 10 (GEMS) and 11 (TROPOMI). During the transport of the dust plume from inland China to the coastal area, AEH changed from 4.0 km at 02:00 UTC to 2.0 km at 06:00 UTC. By contrast, ALH from TROPOMI only observed the 1.5~2.5 km layer height over East

Asia around 04:00 UTC. Although the AEH from GEMS had spatio-temporal uncertainty, this case demonstrates the advantage of AEH retrieval from GEMS for continuous monitoring of changes in plume height, in particular during dust transport. As shown in Figure 12, AEH from GEMS showed differences in height of -0.15 ± 0.97 km (compared to CALIOP). In addition, the differences in height of -0.14 ± 1.06 and 1.47 ± 1.09 km over ocean and land as compared to TROPOMI ALH.

From two different case results, proportion values within 1.0 km (or 1.5 km) height difference between TROPOMI and GEMS have strong dependence of surface types. The proportion over land (over ocean) was lower (higher) than the corresponding result from the comparison of GEMS and CALIOP. The TROPOMI ALH from version 2 is strong surface type dependence as compared to the ground lidar data (Michailidis et al., 2023). However, the relationship between TROPOMI ALH and GEMS AEH in 20% interval of each TROPOMI ALH have high correlation coefficients. In the case of March 28~30, the correlation coefficients between TROPOMI and GEMS are 0.663 and 0.993 over ocean and land, respectively. In the case of April 26, the correlation coefficients are 0.657 and 0.810 over ocean and land, respectively.

5. Long-term validation

For long-term validation, we used the AEH retrieval results from January to June, 2021. The CALIOP and TROPOMI satellites passed over the study area around 13:30 local time, which is around 04:30 UTC for East Asia and around 06:30 UTC for India. Most temporal collocation pixels aligned with observation times of 04:00~06:00 UTC, respectively. To check the dependence of several retrieval variables, the UVAI, AOD, SSA, and dominant aerosol type in each pixel (TYPE) were obtained from the

L2AERAOD. Although the GEMS algorithm retrieved AEH in the range of 0~10 km, the sensitivity of O₂-O₂ SCD was weak in cases of high AEH because of the vertical distribution of air molecules. To ensure sufficient quality of retrieved data, therefore, the AEHs from GEMS and CALIOP, and the ALH from TROPOMI were used only in pixels where the AEH from GEMS were lower than 5 km.

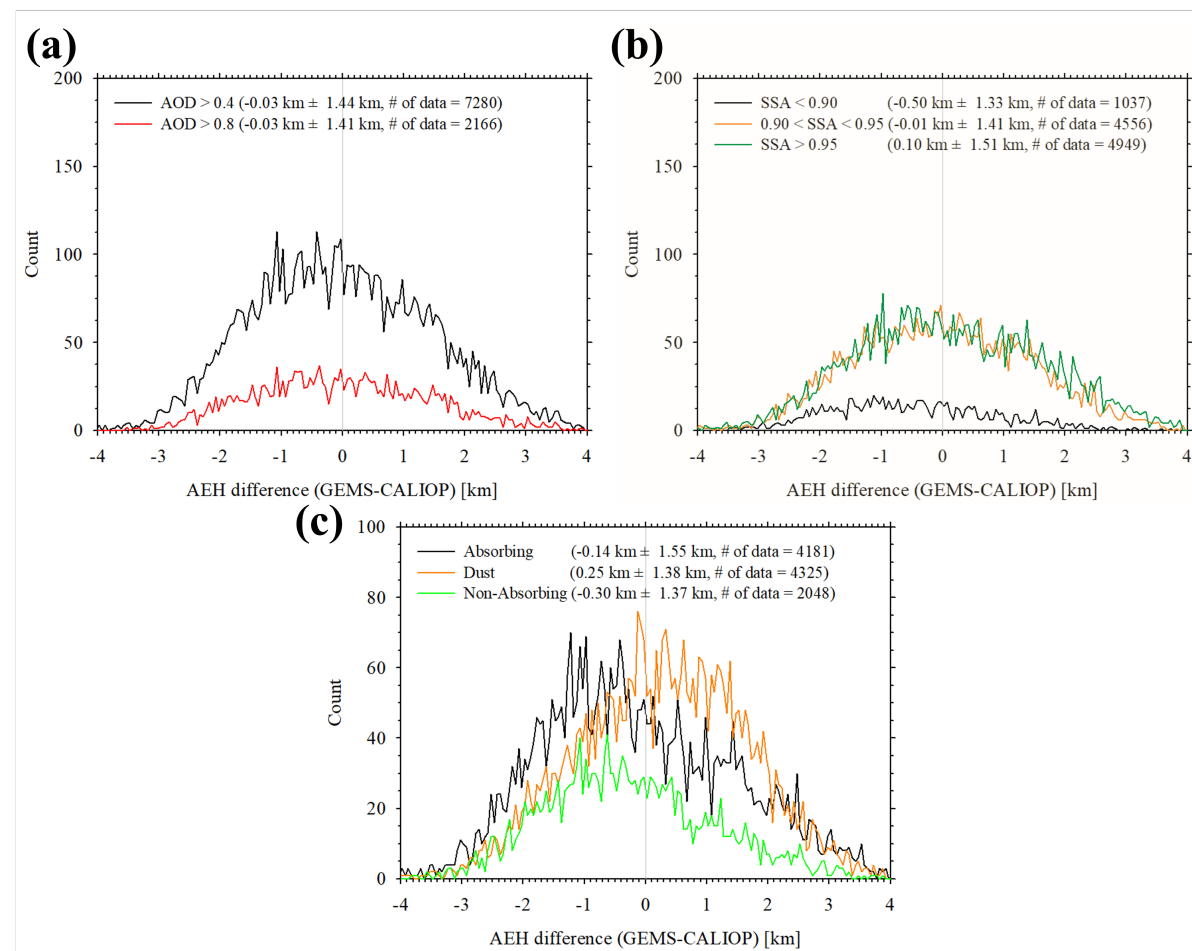


Figure 13. Histogram of AEH difference between CALIOP and GEMS with respect to (a) AOD, (b) SSA, and (c) TYPE from GEMS over the period from January 1 to June 30, 2021.

Figure 13 shows histograms of difference in AEH between GEMS and CALIOP according to AOD at 443 nm, SSA at 443 nm, and TYPE from GEMS. From Figure 13a, the dependence on AOD threshold was insignificant; the average estimated AEH difference was -0.03 km, but the variation in AEH difference was around 1.4 km based on the standard deviation for AOD > 0.4. Because of uncertainty in GEMS aerosol operational products, AEH from GEMS exhibits large variability. Although L2AERAOD from GEMS retrieved the AOD, SSA, and aerosol types, the retrieved results from L2AERAOD include significant uncertainty. Go et al. (2020) reported that the root-mean square error (RMSE) of AOD between MODIS and OMI UV aerosol algorithm is 0.276~0.341.

In addition, fitting error perturbs the fitting signals and tends to result in the underestimation of SCD. Although the fitting error of O₂-O₂ SCD from GEMS radiance was minimized, the fitting error has still remained around 6%, as indicated in Table 2. The discrepancy in fitting condition between the simulated and observed radiance biased the SCD estimation, which in turn led to bias and variation in the AEH retrieval. Combined with the high sensitivity of AEH errors to aerosol optical properties, uncertainty arising from L2AERAOD causes AEH variability.

The variation in AEH difference between observation platforms is shown in Figure 13b as a histogram according to SSA threshold. Across the entire SSA threshold range, the standard deviation of the AEH difference was 1.33~1.51 km. In particular, this standard deviation decreased slightly with decreasing SSA. The aerosol height parameter is more sensitive to absorbing-dominant aerosols than scattering-dominant aerosols (e.g., Park *et al.*, 2016; Nanda *et al.*, 2020). For this reason, the variability of AEH is smaller in absorbing-dominant aerosols than scattering-dominant aerosols, if the

uncertainty of other aerosol parameters (AOD, SSA, and TYPE) is the same conditions.

Figure 13c shows the dependence of AEH difference on TYPE. The TYPE product included dependence on the aerosol size and optical absorptivity. For this reason, the AEH difference graphs for the “Dust” and “Absorbing” types differ, despite both types being absorbing-dominant aerosols. The AEH difference for the “Absorbing” type showed a negative bias with a large standard deviation, whereas a positive bias with a small standard deviation was obtained for the “Dust” type. The AEH difference for the “Non-Absorbing” aerosol type showed the largest negative bias in this comparison. These results suggest that the aerosol size distribution of fine particles affects the negative bias of AEH. Combined with the AEH difference bias illustrated in Figure 8b, these findings indicate that the bias in AEH difference for “Absorbing” aerosols is weakened by their absorbing-dominant property.

Figure 14 shows means and standard deviations for AEH difference between CALIOP and GEMS according to AOD and AI values from GEMS. For AOD, the mean AEH difference ranged from -0.13 to 0.03 km with a standard deviation of approximately 1.45 km. Similar to Figure 13a, the variation in AEH difference with AOD change was insignificant. For the AI, the lowest AEH difference was -0.19 km, obtained for the AI range of 1.5~2.0. The largest AEH difference was 0.24 km for the AI range of 4.0~4.5. Although the AEH difference varied slightly, no consistent tendency in AEH variation with AI was observed. Overall, the standard deviation of AEH difference ranged from 1.49 km ($0.0 < AI < 0.5$) to 1.18 km ($4.5 < AI < 5.0$), and a consistent tendency of decreasing variance in AEH difference was found with increasing AI.

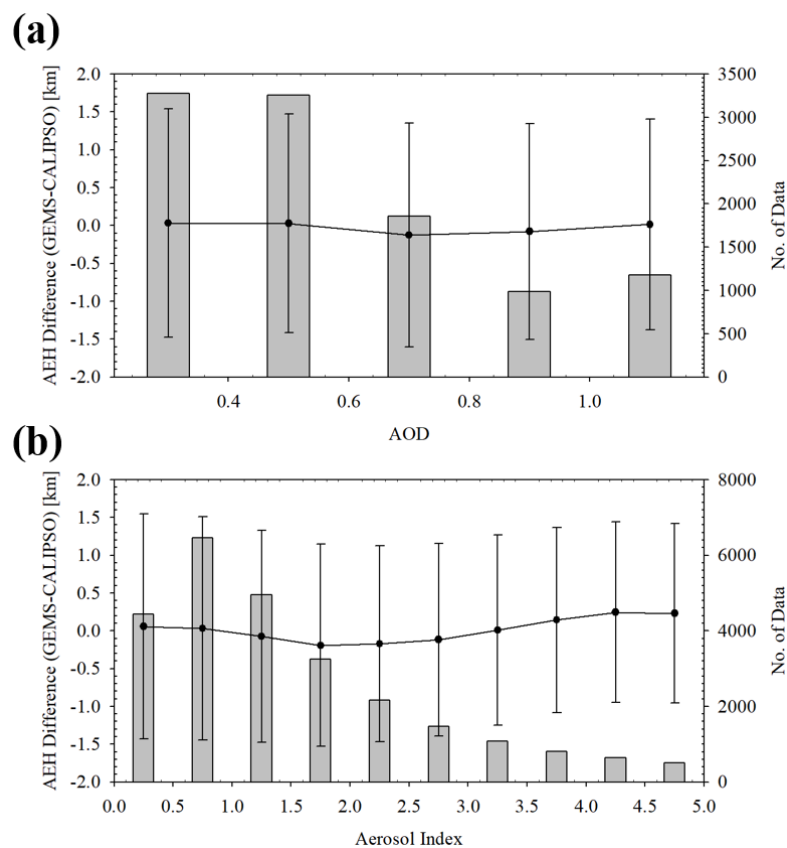


Figure 14. AEH difference between CALIOP and GEMS with respect to ranges of (a) AOD and (b) AI obtained from GEMS from January 1 to June 30, 2021 (Black dots and error bars denote the mean and standard deviation of AEH difference, while the gray-box indicates the number of data).

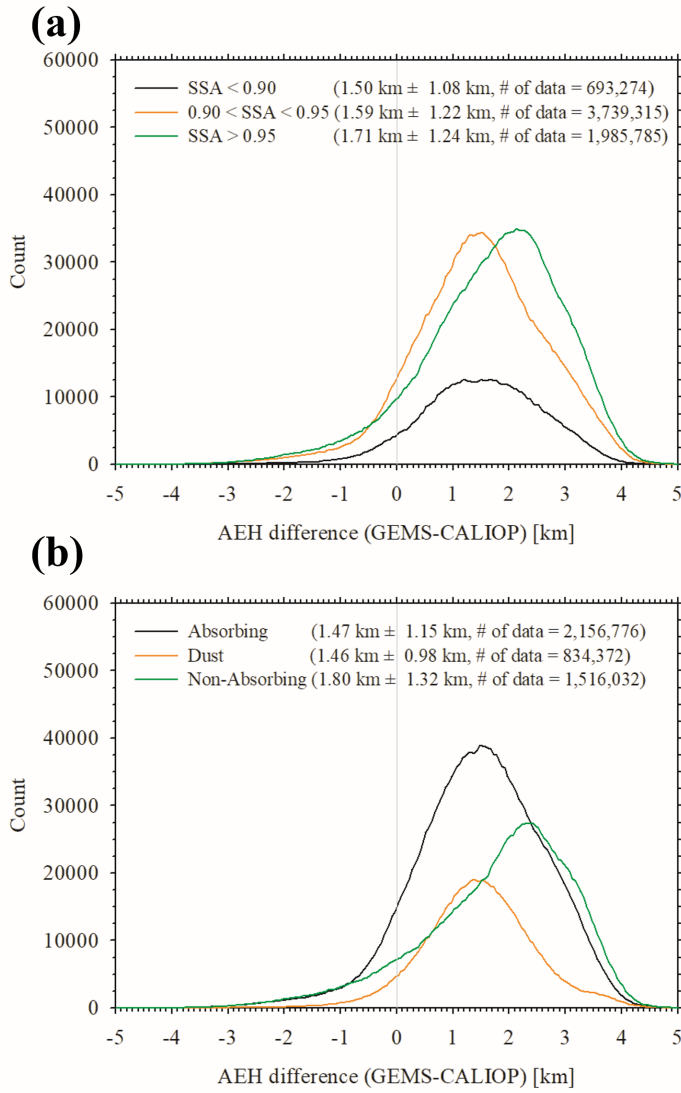


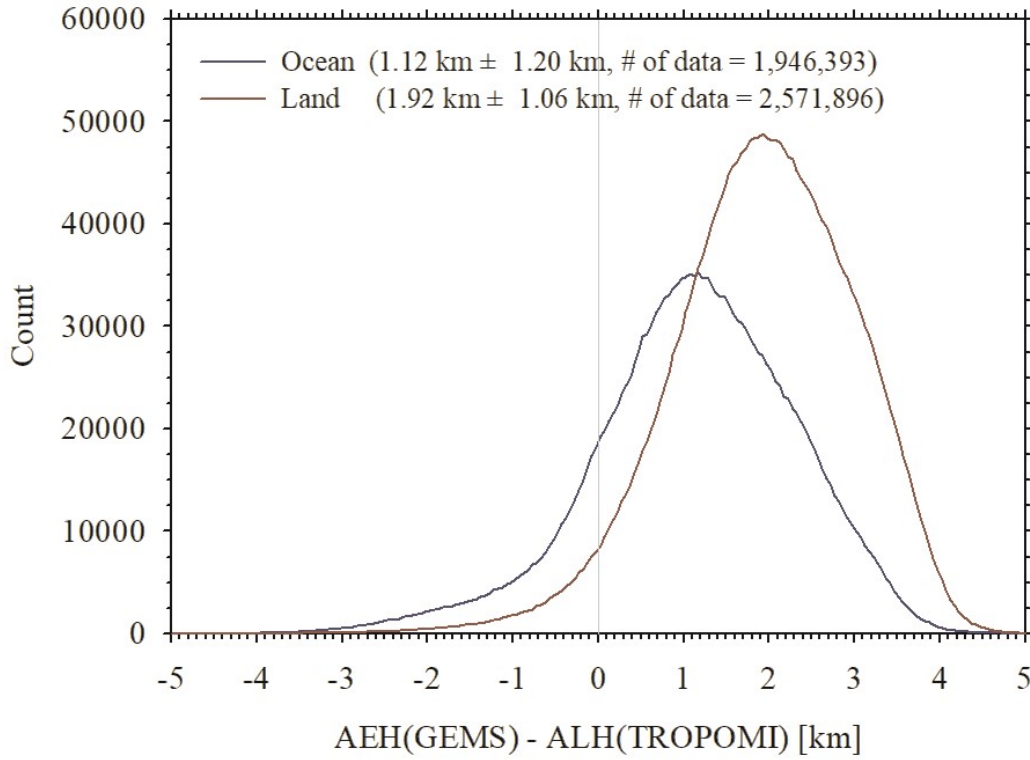
Figure 15. Histograms of differences between TROPOMI ALH and GEMS AEH with respect to (a) SSA, and (b) TYPE from GEMS in the period from January 1 to June 30, 2021.

Figure 15 shows histograms of differences between GEMS AEH and TROPOMI ALH according to the SSA and TYPE obtained from GEMS. The difference between GEMS AEH and TROPOMI ALH depends on both SSA and TYPE. The mean difference value between GEMS AEH and TROPOMI ALH decreased as the aerosol

absorptivity increased. This difference was 1.50 ± 1.08 , 1.59 ± 1.22 , and 1.71 ± 1.24 km for pixels of $SSA < 0.90$, $0.90 < SSA < 0.95$, and $SSA > 0.95$, respectively. Comparing these results to Figure 13b, we find that the standard deviation of the comparison with TROPOMI was approximately 75% of the corresponding value for CALIOP. It is because both TROPOMI and GEMS are passive sensors that use similar retrieval methods for oxygen absorption bands. Nanda *et al.* (2020) showed that the operational algorithm of TROPOMI operational algorithm can provide ALH pixel retrievals only for scenes dominated by absorbing aerosol particles. In addition, Griffin *et al.* (2020) reported that the pixels with low positive UVAI values (weak absorbing cases) are identified with low QA values ($QA \leq 0.5$) in the offline product of ALH. Although the TROPOMI ALH algorithm updated for the expansion of retrieval range, the contrast of O_2 - O_2 SCD to the aerosol layer height change is fundamentally weak sensitivity in scattering dominant aerosols (e.g., Park *et al.*, 2016). For this reason, the bias and standard deviation of height difference between GEMS and ALH is generally larger in high SSA.

In addition, difference between GEMS AEH and TROPOMI ALH depends on TYPE, as shown in Figure 15b. The difference was 1.47 ± 1.15 , 1.46 ± 0.98 , and 1.80 ± 1.32 km for “Absorbing”, “Dust”, and “Non-Absorbing” type aerosols, respectively. Similar to Figure 13c, the TYPE dependence of aerosol height information was influenced by both absorptivity and size information. “Dust” type of aerosol is mainly transported in the free troposphere with gaussian-like vertical distribution, and the associated plume thickness is highly variable. However, “Absorbing” aerosols mainly originate from anthropogenic emissions in East Asia and mostly distributed near the surface with homogeneous concentration (e.g., Gao *et al.*, 2014; Wang *et al.*, 2012; Peng *et al.*, 2016).

620 Transport patterns and vertical distribution shape depending on aerosol types, can affect
 621 the accuracy of aerosol height retrieval results.



622
 623 **Figure 16.** Histogram of the difference between TROPOMI ALH and GEMS AEH
 624 over land and ocean pixels, respectively, from January 1 to June 30, 2021.

625
 626 The non-Lambertian effect on the land surface impacted surface albedo uncertainty
 627 during AEH retrieval, and this effect led to bias and variance in AEH. In this study, the
 628 minimum Lambertian equivalent reflectance was used as the reference reflectance value.
 629 However, surface reflectivity has geometric dependence due to non-Lambertian effects,
 630 which leads to a bias of 0.01-0.02 for surface reflectance over the land surface (e.g., Qin
 631 *et al.*, 2019). To identify the sensitivity of surface property, a histogram was constructed
 632 of difference between GEMS AEH and TROPOMI ALH after classification into land

and ocean surface types, as shown in Figure 16. From the statistical results, the mean differences were estimated to be 1.12 ± 1.20 and 1.92 ± 1.06 km for ocean and land pixels, respectively. The bias has significant difference between two different surface types. Michailidis et al. (2023) explained that the experimental retrieval range of ALH from TROPOMI is 0.27~6.5 km and 0.06~2.15 km over ocean and land, respectively. Constrained retrieval range over land by the TROPOMI influences the negative bias of aerosol height retrieval and increases the mean difference of aerosol height between GEMS and TROPOMI. In addition, the non-Lambertian effect for surface reflectance is also affecting the increasing the discrepancy of aerosol height information.

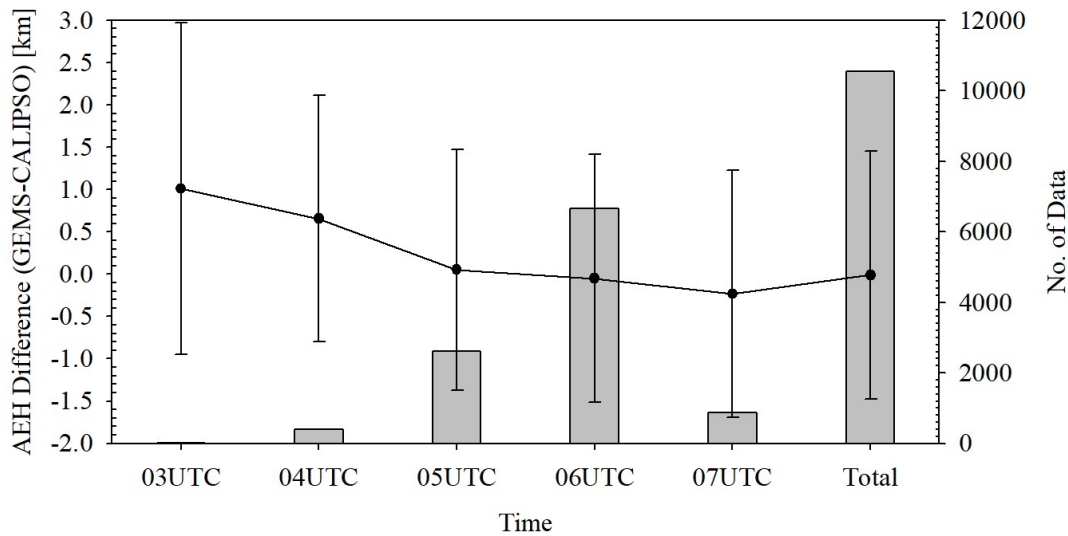


Figure 17. Diurnal dependence of AEH difference between CALIOP and GEMS from January 1 to June 30, 2021 (Black dots and error bars denote the mean and standard deviation of AEH difference, while the gray-box indicates the number of data).

The results of hourly statistical analyses are presented in Figure 17. Because they use

a consistent definition of AEH, we show only a comparison of GEMS and CALIOP. The diurnal variation in AEH difference ranged from -0.23 ± 1.45 km (07:00 UTC, Number of Data = 867) to 1.01 ± 1.96 km (03:00 UTC, Number of Data = 23). However, the number of pixels observed at 03:00 UTC was insufficient for the identification of diurnal variation. The AEH difference of 0.66 ± 1.45 km was the next highest value obtained at 04:00 UTC (Number of Data = 395). The inhomogeneous number of data is mainly due to the lack of spatial homogeneity among retrieval pixels. Over India, very high AOD values were consistently observed during the comparison period. Otherwise, the AEH was only retrieved under conditions of severe anthropogenic emissions over East Asia. In addition, the diurnal variation in AEH difference was caused by spatial characteristics of AEH difference. From 03:00 to 05:00 UTC, CALIOP mainly passed over East Asia, which has numerous sources of aerosol emissions, including biomass burning, dust, and industrial activity. In addition, GEMS observed only the eastern part of India, which is dominated by anthropogenic aerosols. The spatial distribution of the dominant aerosol types may impact the diurnal variation in AEH difference.

6. Summary & Conclusions

Based on the possibility of retrieving AEH from environmental satellite sensors, an AEH retrieval algorithm for GEMS was developed that solely uses the O_2-O_2 absorption band with considering aerosol and surface properties. Because the sensitivity of AEH retrieval is strongly affected by optical amounts and properties of aerosols, as well as surface reflectivity, an AEH retrieval algorithm for GEMS was developed after retrieval of the GEMS operational algorithms, L2AERAOD and L2SFC. With the newly developed retrieval algorithm, GEMS can be used to monitor aerosol vertical

information with high temporal and spatial resolution.

For dust plumes over East Asia, AEH retrieval results from GEMS indicated appropriated aerosol vertical information. After spatial and temporal colocation, the AEH from GEMS aligned well with the AEH information obtained from CALIOP. The differences in AEH between GEMS and CALIOP for dust plume cases were -0.13 ± 1.32 and -0.15 ± 0.97 km, with 49.9~75.9% and 73.2~86.7% of all pixels showing differences less than 1.0 and 1.5 km, respectively. Large AEH uncertainty was found mostly over inland China due to uncertainty in surface reflectance and AOD over the land surface. In addition, AEH from GEMS was overestimated compared to the TROPOMI ALH results. The overestimation is partially caused by different definitions of ALH from TROPOMI and AEH from GEMS.

In long-term intercomparison with CALIOP, the average AEH difference was estimated to be -0.03 km, with a standard deviation of 1.4 km under the scenario of $AOD > 0.4$. The large variation in AEH difference between GEMS and CALIOP was caused by uncertainty in the input parameters estimated from L2AERAOD and L2SFC. In the long-term intercomparison against TROPOMI, this difference was dependent on both SSA and TYPE. The difference was 1.50 ± 1.08 km, 1.59 ± 1.22 km, and 1.71 ± 1.24 km for pixels with $SSA < 0.90$, $0.90 < SSA < 0.95$, and $SSA > 0.95$, respectively. In addition, differences of 1.47 ± 1.15 km, 1.46 ± 0.98 km, and 1.80 ± 1.32 km were obtained for the “Absorbing”, “Dust”, and the “Non-Absorbing” types of aerosols, respectively. The AEH difference also has a diurnal dependence, which is ranged from -0.23 ± 1.45 km to 1.01 ± 1.96 km, due to the spatial characteristics of dominant aerosol types.

The case studies and results of the long-term validation show that AEH retrieved

from GEMS can provide information on aerosol vertical distribution, with applications in diverse research fields. The AEH results with the long-term statistical accuracy make possible to use the application study for AMF calculation of GEMS trace gas retrieval. In addition, AEH considerably affects the surface particulate matter (PM) concentration obtained from satellite-based AOD because PM estimation is significantly affected by the mixing layer height of aerosols. For this reason, the AEH can provide the effective mixing layer height of aerosols for anthropogenic aerosols, and also provide the vertical patterns for long-range transport of aerosols.

Although several fields of study may apply the AEH retrieval results, retrieval uncertainty in AEH remains due to the uncertainty of retrieved AOD and SSA. In addition, the uncertainty in surface reflectance and the discrepancy in O_2-O_2 SCD values between the simulation results and observations can be affected to the potential error sources of AEH from GEMS. To minimize the AEH retrieval uncertainty, further analysis related to the optimized input parameters of AOD, SSA, and aerosol type information is essential. For this reason, the quantitative analysis AEH uncertainty due to aerosol and surface property is important for the improvement of AEH retrieval algorithm. In addition, aerosol optical property retrieval by the visible channel will be needed for the further study to improve the aerosol type determination. Although the aerosol indices of UV and visible provide the aerosol type information, developing the aerosol type classification algorithm is necessary to make synergy with AEH retrieval. AEH provides representative layer height information as only one variable because of its sole reliance on O_2-O_2 SCD for direct estimation of aerosol height information. This method is limited to the consideration of aerosol vertical structures (i.e., Gaussian or exponential vertical distribution structures). Rather than using the GEMS sensor alone,

721 using another absorption band for oxygen-based materials would provide additional
722 scattering information about aerosols.

723

Data Availability

The TROPOMI ALH product is available from <http://doi.org/10.5270/S5P-7g4iapn>, and the CALIOP aerosol extinction profile product is available from https://doi.org/10.5067/CALIOP/CALIPSO/CAL_LID_L2_05kmAPro-Prov-V3-41.

The GEMS AEH and AERAOD products are available from the Environmental Satellite Center in National Institute of Environmental Research (NIER) of the Republic of Korea.

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