Insight into Global Trends in Aerosol Composition over 2005-2015 Inferred from the OMI Ultraviolet Aerosol Index

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Abstract

Observations of aerosol scattering and absorption offer valuable information about aerosol composition. We apply a simulation of the Ultraviolet Aerosol Index (UVAI), a method of detecting aerosol absorption from satellite observations, to interpret UVAI values observed by the Ozone Monitoring Instrument (OMI) over 2005-2015 to understand global trends in aerosol composition. We conduct our simulation using the vector radiative transfer model VLIDORT with aerosol fields from the global chemical transport model GEOS-Chem. We examine the 2005-2015 trends in individual aerosol species from GEOS-Chem, and apply these trends to the UVAI simulation to calculate the change in simulated UVAI due to the trends in individual aerosol species. We find that global trends in the UVAI are largely explained by trends in absorption by mineral dust, absorption by brown carbon, and scattering by secondary inorganic aerosol. Trends in absorption by mineral dust dominate the simulated UVAI trends over North Africa, the Middle-East, East Asia, and Australia. The UVAI simulation well resolves observed negative UVAI trends over Australia, but underestimates positive UVAI trends over North Africa and Central Asia near the Aral Sea, and underestimates negative UVAI trends over East Asia. We find evidence of an increasing dust source from the desiccating Aral Sea, that may not be well represented by the current generation of models. Trends in absorption by brown carbon dominate the simulated UVAI trends over biomass burning regions. The UVAI simulation reproduces observed negative trends over central South America and West Africa, but underestimates observed UVAI trends over boreal forests. Trends in scattering by secondary inorganic aerosol dominate the simulated UVAI
trends over the eastern United States and eastern India. The UVAI simulation slightly overestimates the observed positive UVAI trends over the eastern United States, and underestimates the observed negative UVAI trends over India. Quantitative simulation of the OMI UVAI offers new insight into global trends in aerosol composition.

1. Introduction

Atmospheric aerosols have significant climate impacts due to their ability to scatter and absorb solar radiation and to their indirect effect through modification of cloud properties. The exact magnitude of the direct radiative forcing remains highly uncertain (IPCC, 2014), although most studies agree it is significant (Andreae and Gelencsér, 2006; Mann and Emanuel, 2006; Mauritsen, 2016). Storelvmo et al. (2016) estimate that changes in global aerosol loading over the past 45 years have caused cooling (direct and indirect) that masks about one third of the atmospheric warming due to increasing greenhouse gas emissions. Aerosol absorption has been estimated to be the second largest source of atmospheric warming after carbon dioxide (Ramanathan and Carmichael, 2008; Bond et al., 2013; IPCC, 2014), although considerable uncertainty remains regarding the exact magnitude (Stier et al., 2007). The large uncertainty regarding the direct radiative impacts of aerosols on climate is driven by the large variability in aerosol physical and chemical properties, as well as their various emission sources, making it extremely difficult to fully understand their interactions with radiation (Pöschl, 2005; Moosmüller et al., 2009; Curci et al., 2015; Kristiansen et al., 2016). Global observations of trends in aerosol scattering and absorption would offer valuable constraints on trends in aerosol sources and composition.

The emissions of aerosols and their precursors have changed significantly over the past decade. In North America and Europe, the anthropogenic emissions of most aerosol species (e.g. black carbon, organic aerosols) and aerosol precursors (e.g. sulfur dioxide and nitrogen oxides) have decreased due to pollution controls (Leibensperger et al., 2012; Klimont et al., 2013; Curier et al., 2014; Simon et al., 2014; Xing et al., 2015; Li et al., 2017a). By contrast, emissions of aerosols and aerosol precursors have increased in developing countries due to increased industrial activity, particularly in China and India. Chinese emissions of black carbon (BC), organic carbon (OC), and nitrogen oxides (NOx) have been increasing over the past decade (Zhao et al., 2013; Cui et al., 2015), although in the most recent years NOx emissions have been declining, driven by
denitration devices at power plants (Liu et al., 2016). Due to the wide implementation of flue-gas
desulfurization equipment on most power plants in China, emissions of sulfur dioxide (SO₂) in
some regions have been decreasing since about 2006-2008 (Lu et al., 2011; Wang et al., 2015;
Fioletov et al., 2016). Indian emissions of anthropogenic aerosols and their precursors have been
increasing over the past decade (Lu et al., 2011; Klimont et al., 2017). There have also been
significant changes in global dust and biomass burning emissions. Shao et al. (2013) use synoptic
data to estimate a global decrease in dust emissions between 1974 and 2012, driven largely by
reductions from North Africa with weaker contributions from Northeast Asia, South America, and
South Africa. By examining trends in burned area, Giglio et al. (2013) estimate a decrease in global
biomass burning emissions between 2000 and 2012. Trends in aerosol composition produced by
these changing emissions may be detectable from satellite observations of aerosol scattering and
absorption.

Detection of aerosol composition from passive nadir satellite observations is exceedingly
difficult; few methods exist. The aerosol-type classification provided by retrievals from the MISR
instrument, enabled by multi-angle viewing, is one such source of information about aerosol
composition from constraints on particle size, shape, and single scattering albedo (SSA) (Kahn
and Gaitley, 2015). MISR retrievals have been used to classify particles relating to events such as
biomass burning, desert dust, volcanic eruptions, and pollution events (e.g. Liu et al., 2007;
Kalashnikova and Kahn, 2008; Dey and Di Girolamo, 2011; Scollo et al., 2012; Guo et al., 2013).
The most commonly used satellite product for aerosol information is aerosol optical depth (AOD),
the columnar extinction of radiation by atmospheric aerosols. AOD can be retrieved from satellite
measurements of top of atmosphere radiance in combination with prior knowledge of aerosol
optical properties. Several studies have examined trends in satellite AOD. Following trends in
emissions, over the past decade positive trends in satellite AOD have been observed over Asia and
Africa corresponding to regions experiencing industrial growth (de Meij et al., 2012; Chin et al.,
2014a; Mao et al., 2014; Mehta et al., 2016), while negative trends in satellite AOD have been
observed over North America and Europe, largely due to pollution controls (Hsu et al., 2012; de
Meij et al., 2012; Chin et al., 2014b; Mehta et al., 2016). Studies such as these demonstrate the
information about the evolution of aerosol abundance offered by total column AOD retrievals,
however measurements of absorption would complement the scattering information in AOD
retrievals by providing independent information on aerosol composition.
The Ultraviolet Aerosol Index (UVAI) is a method of detecting aerosol absorption from satellite measured radiances (Herman et al., 1997; Torres et al., 1998). Because the UVAI is calculated from measured radiances, a priori assumptions about aerosol composition are not required for its calculation, thus yielding independent information on aerosol scattering (Herman et al., 1997; Torres et al., 1998, 2007; de Graaf et al., 2005; Penning de Vries et al., 2009) and absorption. The UVAI has been widely applied to examine mineral dust (Israelevich et al., 2002; Schepanski et al., 2007; Badarinath et al., 2010; Huang et al., 2010) and biomass burning aerosols (Duncan et al., 2003; Guan et al., 2010; Torres et al., 2010; Kaskaoutis et al., 2011; Mielonen et al., 2012), including brown carbon (Jethva and Torres, 2011; Hammer et al., 2016). The UVAI is not typically used to examine scattering aerosol, however aerosol scattering causes a net decrease in the overall value of the UVAI, meaning that the UVAI could be used to detect changes due to both aerosol absorption and scattering. Prior interpretation of the UVAI has been complicated by its dependence on geophysical parameters, such as aerosol layer height (Herman et al., 1997; Torres et al., 1998; de Graaf et al., 2005). Examining trends in the UVAI would provide an exciting opportunity to investigate the evolution of aerosol absorption and scattering over time, if the multiple parameters affecting the UVAI could be accounted for through simulation.

In this work, we apply a simulation of the UVAI, which was developed and evaluated regionally and seasonally in Hammer et al. (2016), to interpret trends in recently reprocessed OMI UVAI observations for 2005-2015 to understand global changes in aerosol composition. We interpret observed UVAI values by using a radiative transfer model (VLIDORT) to calculate UVAI values as a function of simulated aerosol composition from the global 3-D chemical transport model GEOS-Chem. By using scene-dependent OMI viewing geometry together with scene-dependent modelled atmospheric composition we enable quantitative comparison of model results with observations. Comparison of trends in observed OMI UVAI values to the trends in simulated UVAI values, which are calculated using known aerosol composition, enables qualification of how changes in aerosol absorption and scattering could influence the observed UVAI trends and identification of model development needs. We conduct our analysis at the global scale to understand trends worldwide. Section 2 describes the OMI UVAI observations and our UVAI simulation. Section 3 examines the trends in emissions of GEOS-Chem aerosols and their precursors for 2005-2015 to provide context for the trends in our simulated UVAI. Section 4 compares the mean values over 2005-2015 of the OMI UVAI and our simulated UVAI. Section 5
compares the 2005-2015 trends in OMI and simulated UVAI values. In section 6 we examine the
sensitivity of the UVAI to changes in the abundance of individual aerosol species. Trends in our
UVAI simulation are interpreted by applying the trends in the GEOS-Chem aerosol species to
calculate the associated change in UVAI. Section 7 reports the conclusions.

2. Methods

2.1 OMI Ultraviolet Aerosol Index

The OMI Ultraviolet Aerosol Index is a method of detecting absorbing aerosols from
satellite measurements in the near-UV wavelength region and is a product of the OMI Near-UV
algorithm (OMAERUV) (Herman et al., 1997; Torres et al., 1998, 2007). The OMAERUV
algorithm uses the 354 nm and 388 nm radiances measured by OMI to calculate the UVAI as a
measure of the deviation from a purely Rayleigh scattering atmosphere bounded by a Lambertian
reflecting surface. Positive UVAI values indicate absorbing aerosol while negative values indicate
non-absorbing aerosol. Near-zero values occur when clouds and Rayleigh scattering dominate.
Negative UVAI values due to aerosol scattering are often weak and have historically been affected
by noise in previous datasets (Torres et al., 2007; Penning de Vries et al., 2015). Because UVAI
values are calculated from top of atmosphere (TOA) radiance which contains total aerosol effects,
the presence (or lack) of scattering aerosol along with absorbing aerosol can either weaken (or
strengthen) the absorption signal. Therefore the UVAI could be used to detect changes over time
due to both aerosol absorption and scattering.

The main source of error affecting a trend analysis of the UVAI is the OMI row anomaly
which has reduced the sensor viewing capability for specific scan angles since 2008
(http://projects.knmi.nl/omi/research/product/rowanomaly-background.php). The sudden
suppression of observations for specific viewing geometries (i.e. the row anomaly), could cause
an additional spurious trend in the UVAI trend calculation. We address this concern by considering
only scan positions 3-23 which remain unaffected by the row anomaly, and also by using the
recently reprocessed OMAERUV UVAI that is less sensitive to scan-angle dependent cloud
artifacts due to the implementation of a Mie-scattering based water cloud model (Torres et al.,
2018). We focus on cloud-filtered observations by excluding scenes with OMI UVAI radiative
cloud fraction exceeding 5% to further reduce uncertainty due to clouds. Furthermore, we focus
on 10-years of observations so that multiple observations can reduce the random error of UVAI observations.

Because the OMI UVAI is calculated directly from OMI measured radiances, instrument degradation over time could be a significant source of uncertainty (Povey and Grainger, 2015). Schenkeveld et al. (2017) found that the OMI radiances in the channel used for the UVAI have changed by only ~1-1.15% over the entire OMI record. Applying this change to the radiances results in a change in the absolute UVAI of ~10^{-4}, which is negligible. Schenkeveld et al. (2017) also calculated the trend in the ratio of the 354/380 nm radiances measured by OMI for pixels unaffected by the OMI row anomaly and over the Tropical Pacific where the presence of aerosol is expected to be minimal, to assess the change in the spectral dependence of OMI’s overall radiance calibration over the course of the mission. They found that the trend in the 354/380 nm radiance ratio over the entire OMI record was < 0.5 % per decade. We estimate the effect of instrument degradation on our trend analysis by calculating the change in UVAI associated with the 0.5 % per decade trend in the 354/380 nm radiance ratio. Applying the trend in 354/380 nm radiance ratio to the UVAI calculation globally resulted in a negligible change in the UVAI of ~2×10^{-4} yr^{-1}. To avoid the influence of any possible spurious trends due to instrument degradation on our trend analysis, we subtract the trend in global mean UVAI from the cloud-filtered UVAI prior to interpretation.

We perform trend analysis on monthly mean time series data for the years 2005-2015 using Generalized Least Squares (GLS) regression, as described by Boys et al. (2014). Prior to regression, the data are aggregated to monthly mean values, and the monthly time series data are deseasonalized by subtracting the monthly mean for the period 2005-2015 to focus on the long-term trend. Deseasonalization is a recommended method to accurately calculate a long-term trend in a seasonally-varying time series (Weatherhead et al., 1998, 2002; Wilks, 2011), and is widely employed for the trend analysis of geophysical data including temperature, chemical species concentrations, relative humidity, cloud cover, and aerosol parameters (Reynolds and Reynolds, 1988; Prinn et al., 1992; Pelletier and Turcotte, 1997; Zhang et al., 1997; Dai, 2006; Norris and Wild, 2007; Hsu et al., 2012b; Boys et al., 2014; Li et al., 2014; Ma et al., 2016). Each pixel is required to have data for at least 60% of the time-period before regression is performed. In the following section, we discuss our UVAI simulation and the implementation of the new UVAI algorithm in the simulation.
2.2 Simulated UVAI

We simulate the UVAI using the VLIDORT radiative transfer model (Spurr, 2006), following Buchard et al. (2015) and Hammer et al. (2016). We calculate the top of atmosphere radiances at 354 nm and 388 nm needed for the UVAI calculation by supplying VLIDORT with the OMI viewing geometry for each scene, as well as the GEOS-Chem simulation of vertical profiles of aerosol extinction, spectrally dependent single scattering albedo, and the corresponding spectrally dependent scattering phase function. Thus these parameters account for the sensitivity of the UVAI to aerosol layer height and spectrally dependent aerosol optical properties.

We introduce to the UVAI simulation a Mie-scattering based water cloud model (Deirmendjian, 1964) for consistency with the reprocessed OMI UVAI dataset. Following Torres et al. (2018), we compute the radiances used in the UVAI calculation as a combination of clear and cloudy sky conditions. We use the same cloud fractions and cloud optical depths used in the OMI UVAI algorithm for coincident OMI pixels. We avoid cloudy scenes by considering only pixels with OMI radiative cloud fraction of less than 5%. For the UVAI calculation we use the surface reflectance fields provided by OMI. We calculated the 2005-2015 trends in these surface reflectance fields, and found that they were statistically insignificant globally and on the order of $10^{-5}$ yr$^{-1}$. We calculated the change in UVAI due to a change in surface reflectance of this order of magnitude, and found that the change in UVAI was negligible. We also calculated the change in UVAI due to changes in simulated aerosol altitude, but found that the trends in aerosol altitude were negligible (order $10^{-5}$ hPa yr$^{-1}$). Therefore we focus our analysis on trends in aerosol composition which have a larger effect on the UVAI as demonstrated below.

We use the GEOS-Chem model v11-01 (http://geos-chem.org) as input to the UVAI simulation, and to calculate the sensitivity of the UVAI simulation to aerosol composition. The simulation is driven by assimilated meteorological data from MERRA-2 Reanalysis of the NASA Global Modeling and Assimilation Office (GMAO). Our simulation is conducted at a spatial resolution of 2° x 2.5° with 47 vertical levels for the years 2005-2015. We supply VLIDORT with GEOS-Chem aerosol fields coincident with OMI observations.

GEOS-Chem contains a detailed oxidant-aerosol chemical mechanism (Bey et al., 2001; Park et al., 2004). The aerosol simulation includes the sulfate-nitrate-ammonium system (Fountoukis and Nenes, 2007; Park et al., 2004; Pye et al., 2009), primary carbonaceous aerosol (Park et al., 2003), mineral dust (Fairlie et al., 2007), and sea salt (Jaeglé et al., 2011). Semivolatile
primary organic carbon and secondary organic aerosol formation is described in Pye et al. (2010). We update the original semi-volatile partitioning of secondary OA (SOA) formed from isoprene with the irreversible uptake scheme in Marais et al. (2016). HNO$_3$ concentrations are reduced following Heald et al. (2012). Aerosol optical properties are based on the Global Aerosol Data Set (GADS) (Koepke et al., 1997) as originally implemented by Martin et al. (2003), with updates for organics and secondary inorganics from aircraft observations (Drury et al., 2010), for mineral dust (Lee et al., 2009; Ridley et al., 2012), and for absorbing brown carbon (Hammer et al., 2016). Here we update the mineral dust optics at ultraviolet wavelengths using a refractive index that minimizes the difference between the mean simulated and OMI UVAI values to allow focus on trends. Aerosols are treated as externally mixed.

Anthropogenic emissions are from the EDGARv4.3.1 global inventory (Crippa et al., 2016) with emissions overwritten in areas with regional inventories for the United States (NEI11; Travis et al., 2016), Canada (CAC), Mexico (BRAVO; Kuhns et al., 2005), Europe (EMEP; http://www.emep.int/), China (MEIC v1.2; Li et al., 2017a) and elsewhere in Asia (MIX; Li et al., 2017a). Emissions from open fires for individual years from the GFED4 inventory (Giglio et al., 2013) are included. The long-term concentrations from this simulation have been extensively evaluated versus ground-based PM$_{2.5}$ composition measurements where available, and versus satellite-derived PM$_{2.5}$ trends (Li et al., 2017b).

The Supplement evaluates trends in simulated SO$_2$, NO$_2$, and AOD versus satellite retrievals from multiple instruments and algorithms. We find broad consistency between our simulated NO$_2$ and SO$_2$ column trends with those from OMI (Figures S1 and S2). Our simulated AOD trends are generally consistent with the trends in satellite AOD retrievals, except for positive trends in AOD over western North America and near the Aral Sea in most retrieval products, and a negative trend in AOD over Mongolia/Inner Mongolia in all retrieval products (Figure S3).

We filter our GEOS-Chem aerosol simulated fields based on the coincident OMI pixels, which are regridded to the model resolution of 2° x 2.5°. This allows for the direct comparison between our GEOS-Chem simulation and the OMI UVAI observations.

3. Trend in emissions of GEOS-Chem aerosols and their precursors

Figure 1 shows the trends in emissions of aerosols and their precursors from our GEOS-Chem simulation calculated from the GLS regression of monthly time series values for 2005-2015.
Cool colors indicate negative trend values, warm colors indicate positive trend values, and the opacity of the colors indicates the statistical significance of the trends. The trends in emissions of sulfur dioxide (SO$_2$) and nitrogen oxides (NO$_x$) follow similar patterns (Figure 1a and 1b, respectively). Negative trends (−1 to -0.01 kg km$^{-2}$ yr$^{-1}$) are present over North America and Europe, corresponding to pollution controls (Leibensperger et al., 2012; Klimont et al., 2013; Curier et al., 2014; Simon et al., 2014; Xing et al., 2015; Li et al., 2017). Positive trends (0.5 to 1 kg km$^{-2}$ yr$^{-1}$) in both species are present over India and eastern China, however the positive trends in emissions of SO$_2$ over eastern China are interspersed with negative trends (-1 to -0.5 kg km$^{-2}$ yr$^{-1}$) in SO$_2$ emissions, corresponding to the deployment of desulfurization equipment on power plants in recent years (Lu et al., 2011; Klimont et al., 2013; Wang et al., 2015). Ammonia (NH$_3$) emissions (Figure 1c) have positive trends (0.001 to 0.05 kg km$^{-2}$ yr$^{-1}$) over most of South America, North Africa, the Middle-East, and most of Asia with larger trends (0.1 to 0.5 kg km$^{-2}$ yr$^{-1}$) over India and eastern China. There are positive trends (0.001 to 0.05 kg km$^{-2}$ yr$^{-1}$) in black carbon (BC) emissions (Figure 1d) over North Africa, Europe, the Middle-East, India, and China, and negative trends (-0.05 to -0.001 kg km$^{-2}$ yr$^{-1}$) over North America, Europe, West Africa, and central South America. The trends in primary organic aerosol (POA) emissions (Figure 1e) follow a similar pattern as the trends in BC emissions, except there are negative trends (-0.1 to -0.05 kg km$^{-2}$ yr$^{-1}$) over eastern China, and the negative trends (-1 to -0.1 kg km$^{-2}$ yr$^{-1}$) over West Africa and central South America are larger in magnitude reflecting regional changes in fire activity (Chen et al., 2013; Andela and van der Werf, 2014). There are also positive trends (0.001 to 0.05 kg km$^{-2}$ yr$^{-1}$) over the northern United States and Canada. The trends in dust emissions (Figure 1f) show the largest magnitude of all the various species, although many have low statistical significance, with areas of positive and negative trends (> 1 and < -1 kg km$^{-2}$ yr$^{-1}$) over North Africa, positive trends (> 1 kg km$^{-2}$ yr$^{-1}$) parts of the Middle-East, and negative trends (< -1 kg km$^{-2}$ yr$^{-1}$) over northern China and southern Australia.


We examine the seasonal long-term mean UVAI values for insight into the spatial distribution of the aerosol absorption signals. Figures 2 and 3 show the seasonal mean UVAI values for 2005-2015 for OMI and the simulation, respectively. Positive UVAI values between 0.2 and 1.5 indicating aerosol absorption are present over major desert regions globally for both OMI and
the simulation, particularly over the Saharan, Iranian, and Thar deserts. These positive signals are driven by the absorption by mineral dust (Herman et al., 1997; Torres et al., 1998; Buchard et al., 2015). The simulation underestimates some of the smaller dust features captured by OMI, such as over western North America, South America, Australia, and parts of Asia, perhaps reflecting an underestimate in the simulated mineral dust lifetime (Ridley et al. 2012) and missing dust sources (Ginoux et al., 2012; Guan et al., 2016; Huang et al., 2015; Philip et al., 2017). The seasonal variation in the observed and simulated UVAI is similar albeit with larger simulated values in spring (MAM) over North Africa. In all seasons, the UVAI values driven by absorption by dust in the simulation are concentrated mostly over North Africa, while for OMI the UVAI values are more homogeneous over the Middle-East and Asia as well. Positive UVAI values of ~0.2-1 over West and central Africa appearing in both the OMI and simulated values correspond to absorption by brown carbon from biomass burning activities in these regions (Jethva and Torres, 2011; Hammer et al., 2016). Over ocean most data are removed by our strict cloud filter.

5. Trend in UVAI values between 2005-2015

Figure 4 shows the trend in OMI and simulated UVAI values (coincidently sampled from OMI) calculated from the GLS regression of monthly UVAI time series values over 2005-2015. Several regions exhibit consistency between the OMI and simulated UVAI trends. There are statistically significant, positive trends in both OMI and simulated UVAI values over the eastern United States (OMI: 1.0 x10^{-5} to 2.5x10^{-4} yr^{-1}, simulated: 2.5x10^{-4} yr^{-1} to 5.0 x10^{-4} yr^{-1}), and Canada and parts of Russia (OMI: 1.0 x10^{-5} to 2.5x10^{-4} yr^{-1}, simulated: 5.0 x10^{-4} to 2.0x10^{-3} yr^{-1}). Positive UVAI trends (1.0 x10^{-5} to 2.5x10^{-4} yr^{-1}) in both OMI and simulated values are present over Europe, although the simulated trends have low statistical significance. Statistically significant, positive UVAI trends (5.0 x10^{-4} to 2.0 x10^{-3} yr^{-1}) in OMI values are apparent over North Africa, which generally are captured by the simulation but with low statistical significance. Negative UVAI trends (-1.5x10^{-3} yr^{-1} to -1.0x10^{-5} yr^{-1}) in both OMI and simulated values are apparent over most of South America, southern Africa, and Australia. Negative UVAI trends (-2x10^{-3} to -5.0x10^{-4} yr^{-1}) in both OMI and simulated values are present over West Africa, with low statistical significance that could be related to the filtering of persistent clouds. OMI and simulated UVAI values show negative trends (-2x10^{-3} to -5.0x10^{-4} yr^{-1}) over India, although the simulated trends have lower statistical significance.
Some regions have trends in OMI UVAI values which are not captured by the simulation. Statistically significant, positive UVAI trends \((2.5 \times 10^{-4} \text{ yr}^{-1} \text{ to } 1.5 \times 10^{-3} \text{ yr}^{-1})\) over the western United States are apparent in the OMI values but not in the simulation. Zhang et al. (2017) found positive trends in aerosol absorption optical depth from OMI retrievals that they attributed to positive trends in mineral dust over the region, which were not captured by their GEOS-Chem simulation. Statistically significant, positive UVAI trends \((5.0 \times 10^{-4} \text{ to } 2.0 \times 10^{-3} \text{ yr}^{-1})\) in OMI values exist over the Middle-East, while the simulation has negative trends with low statistical significance. The OMI UVAI reveals a region of statistically significant, negative trends \((-2 \times 10^{-3} \text{ to } -5.0 \times 10^{-4} \text{ yr}^{-1})\) over Mongolia/Inner Mongolia which is not captured by the simulation. There is also a small area of statistically significant, positive UVAI trends \((1.5 \times 10^{-3} \text{ to } 2.0 \times 10^{-3} \text{ yr}^{-1})\) in OMI values of over Central Asia between the Caspian Sea and the Aral Sea which is not captured by the simulation. Trends in surface reflectance from the diminishing Aral Sea cannot solely explain the UVAI trends since they extend over the Caspian Sea. Trends in mineral dust are a more likely explanation as discussed further below.

Figures 5 and 6 show the seasonality of the OMI and simulated UVAI trends respectively. The positive UVAI trends over the eastern United States is strongest in summer (JJA) for both OMI and the simulation. The positive UVAI trends over North Africa and the Middle-East are present for all seasons for OMI and for most seasons in the simulation, except in JJA for North Africa and spring (MAM) for the Middle-East. The simulation underestimates the observed UVAI trend over North Africa in SON, perhaps related to an underestimate in trends in mineral dust emissions in the simulation during this season. He et al. (2014) examined the 2000-2010 trends in global surface albedo using the Global Land Surface Satellites (GLASS) dataset and found no significant trends over this region during SON. The negative trend in UVAI values over West Africa is most apparent in the fall (SON) and winter (DJF) for both OMI and the simulation. The negative OMI UVAI trends over Mongolia/Inner Mongolia and the positive OMI UVAI trends near the Aral Sea are strongest in JJA and weakest in DJF, providing evidence for a mineral dust source. The OMI UVAI trend over Mongolia/Inner Mongolia may be part of a longer term trend. Guan et al. (2017) examined dust storm data over northern China (including Inner Mongolia) for the period 1960-2007, and found that dust storm frequency has been declining over the region due to a gradual decrease in wind speed. The current generation of chemical transport models is unlikely to represent the source near the Aral Sea without an explicit parameterization of the drying
The desiccation of the Aral Sea over recent decades has resulted in a steady decline in water coverage over the area (Shi et al., 2014; Shi and Wang, 2015) and has led to the dried up sea bed becoming an increasing source of dust activity in the region (Spivak et al., 2012). Indoitu et al. (2015) found that most dust events are directed towards the west, consistent with the OMI observations. An increase in surface reflectance due to the drying up of the sea bed could also positively influence trends in UVAI. He et al. (2014) found a positive trend over 2000-2010 in surface albedo over the region in JJA and SON, corresponding to when the OMI UVAI trends are strongest.

6. Contribution of individual aerosol species to the simulated UVAI

To further interpret the UVAI trends, we examine the trends in aerosol concentrations from our GEOS-Chem simulation (Figure 7). Figure 7a shows the trends in secondary inorganic aerosol (SIA). There are statistically significant, negative trends over the eastern United States (-1 to -0.05 µg m⁻² yr⁻¹) and statistically significant, positive trends over the Middle-East (0.05 to 0.5 µg m⁻² yr⁻¹), India (0.05 to 1 µg m⁻² yr⁻¹), South America, and southern Africa (0.05 to 0.25 µg m⁻² yr⁻¹). Figure 7b shows the trends in dust. Similar to the trends in emissions, the trends in dust concentrations are of the largest magnitude of the various species, however often with low statistical significance. There are positive trends over the Middle-East (> 2 µg m⁻² yr⁻¹), India (0.05 to 2 µg m⁻² yr⁻¹), and north west China (1 to 2 µg m⁻² yr⁻¹). There are also positive trends (0.05 to 0.25 µg m⁻² yr⁻¹) with low statistical significance over the United States, northern South America, southern Africa, and northern Australia. There is a combination of positive and negative trends (> 2 and < -2 µg m⁻² yr⁻¹) over North Africa, and negative trends over China and Mongolia (< -2 µg m⁻² yr⁻¹) and Australia (-1 to -0.5 µg m⁻² yr⁻¹). Figures 7c and 7d show the trends in total organic aerosol (OA) and the absorbing brown carbon (BrC) component of OA, respectively. Positive trends over Canada and parts of Russia (0.05 to 0.5 µg m⁻² yr⁻¹) in total OA are mainly due to the positive trend in BrC. Statistically significant, negative trends in total OA (-1 to -0.05 µg m⁻² yr⁻¹) over the eastern United States are dominated by scattering organic aerosol. Statistically significant, negative trends (-2 to -0.05 µg m⁻² yr⁻¹) over West Africa and South America for total OA are dominated by the trend in absorbing BrC. Figures 5e and 5f show the trends in black carbon (BC)
and salt, respectively. There are positive trends (0.05 to 0.25 µg m⁻² yr⁻¹) in BC with low statistical significance over India and China. Sea salt trends are negligible.

To gain further insight into how changes in aerosols effect the trends in simulated UVAI, we examine the sensitivity of the UVAI to changes in individual aerosol species. Figure 8 shows the change in annual mean UVAI due to doubling the concentration of individual aerosol species. This information facilitates interpretation of the observed UVAI trends by identifying the chemical components that could explain the observed trends. Doubling scattering SIA concentrations (Figure 8a) decreases the UVAI between -0.25 and -0.1 over most of the globe, with the largest changes over the Eastern United States, Europe, parts of the Middle-East, India, and south east China. Doubling dust concentrations (Figure 8b) produces the largest changes in UVAI, causing increases between 0.5 and 1 over North Africa, and smaller increases between 0.2 and 0.5 over the Middle-East, Europe, and parts of Asia and Australia. Figures 8c and 8d show the changes in UVAI due to doubling total OA concentrations and the absorbing BrC component, respectively. The doubling of BrC increases the UVAI between 0.1 and 0.5 over Canada, West and central Africa, India, parts of Russia, eastern China, and central South America. Doubling total OA concentrations over central South America causes a net decrease of ~ -0.1 as the scattering component of total OA cancels out the absorption by BrC. Doubling BC concentrations (Figure 8e) increases the UVAI of 0.1 over central Africa, India, and south east China, while doubling sea salt concentrations (Figure 8f) has negligible effect on the UVAI.

Figure 9 shows the change in simulated UVAI due to the 2005-2015 trends in individual aerosol species from our GEOS-Chem simulation. The change for each species is calculated by applying the aerosol concentration trends for the individual aerosol type while leaving the concentrations unchanged for the other aerosol species, then taking the difference between this perturbed UVAI simulation and an unperturbed simulation. Negative trends in scattering SIA (Figure 9a) increase the UVAI by 1.0x10⁻⁴ to 7.5x10⁻³ yr⁻¹ over the eastern United States and by 1.0x10⁻⁴ to 2.5x10⁻³ yr⁻¹ over Europe, corresponding to regions of positive UVAI trends in both OMI and the simulation (Figure 4). Increasing SIA decreases the UVAI by -2.5x10⁻³ yr⁻¹ to -1.0x10⁻⁴ yr⁻¹ over the Middle-East, India, and east China. Trends in dust concentrations (Figure 9b) cause the largest change in UVAI with regional increases > 1x10⁻² yr⁻¹ and regional decreases < -1x10⁻² yr⁻¹. Simulated UVAI trends due to mineral dust are mostly negative over North Africa, East Asia, and Australia, while mostly positive over the Middle-East. Noisy trends in regional...
meteorology cause heterogeneous trends in dust and in the UVAI, with low statistical significance.

Figures 9c and 9d show the change in UVAI due to the trends in total OA and the absorbing BrC component of total OA, respectively. Most of the changes in UVAI due to the trends in total OA are caused by the trends in the absorbing BrC component, with increases in the UVAI between $2.5 \times 10^{-3}$ and $1 \times 10^{-2}$ yr$^{-1}$ over Canada and parts of Russia, corresponding to regions of positive UVAI trends for both OMI and the simulation (Figure 4). There are decreases in the UVAI < $1 \times 10^{-2}$ yr$^{-1}$ over central South America and West Africa due to the negative trends in BrC, corresponding to regions of negative UVAI trends for both OMI and the simulation (Figure 4).

Over the eastern United States there is a mixture of increases and decreases in the UVAI due to the trends in scattering organic aerosol. Positive trends in BC increase the UVAI (Figure 9e) by $1.0 \times 10^{-4}$ to $2.5 \times 10^{-3}$ yr$^{-1}$ over India and China. There are no obvious changes in the UVAI due to the trends in sea salt (Figure 9f).

7. Conclusions

Observations of aerosol scattering and absorption offer valuable information about aerosol composition. We simulated the Ultraviolet Aerosol Index (UVAI), a method of detecting aerosol absorption using satellite measurements, to interpret trends in OMI observed UVAI over 2005-2015 to understand global trends in aerosol composition. We conducted our simulation using the vector radiative transfer model VLIDORT with aerosol fields from the global chemical transport model GEOS-Chem.

We demonstrated that interpretation of the OMI UVAI with a quantitative simulation of the UVAI offers information about trends in aerosol composition. We found that global trends in the UVAI were largely explained by trends in absorption by mineral dust, absorption by brown carbon, and scattering by secondary inorganic aerosols. We also identified areas for model development, such as dust emissions from the desiccating Aral Sea.

We examined the 2005-2015 trends in individual aerosol species from GEOS-Chem, and applied these trends to the UVAI simulation to calculate the change in simulated UVAI due to the trends in individual aerosol species. The two most prominent positive trends in the observed UVAI were over North Africa and over Central Asia near the desiccating Aral Sea. The simulated UVAI attributes the positive trends over North Africa to increasing mineral dust, despite an underestimated simulated trend in fall (SON) that deserves further attention. The positive trends
in the observed UVAI over Central Asia near the shrinking Aral Sea are likely due to increased
dust emissions, a feature that is unlikely to be represented in most chemical transport models. The
most prominent negative trends in the observed UVAI were over East Asia, South Asia, and
Australia. The simulation attributed the negative trends over East Asia and Australia to decreasing
mineral dust, despite underestimating the trend in East Asia. The simulation attributed the negative
trend over South Asia to increasing scattering secondary inorganic aerosols, a trend that the
observations imply could be even larger. We found the positive trends in the UVAI over the eastern
United States that were strongest in summer (JJA) in both the observations and the simulation were
driven by negative trends in scattering secondary inorganic aerosol and organic aerosol. Observed
negative trends in winter (DJF) were less well simulated. Over West Africa and South America,
negative trends in UVAI were explained by negative trends in absorbing brown carbon. Thus,
trends in the observed UVAI offer valuable information on the evolution of global aerosol
composition that can be understood through quantitative simulation of the UVAI.

Looking forward, the availability of the UVAI observations from 1979 to the present offer
a unique opportunity to understand long-term trends in aerosol composition. The recent launch of
the TROPOspheric Monitoring Instrument (TROPOMI; Veefkind et al., 2012) and the
forthcoming geostationary constellation offer UVAI observations at finer spatial and temporal
resolution. The forthcoming Multi-Angle Imager for Aerosols (MAIA; Diner et al., 2018) satellite
instrument offers an exciting opportunity to derive even more information about aerosol
composition by combining measurements at ultraviolet wavelengths with multi-angle observations
and polarization sensitivity.

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Figure 1: Trend in emissions of a) sulfur dioxide (SO$_2$) (kg SO$_2$ km$^{-2}$ yr$^{-1}$), b) nitrogen oxides (NO$_x$) (kg NO km$^{-2}$ yr$^{-1}$), ammonia (NH$_3$) (kg NH$_3$ km$^{-2}$ yr$^{-1}$), black carbon (BC) (kg C km$^{-2}$ yr$^{-1}$), primary organic carbon (POA) (kg C km$^{-2}$ yr$^{-1}$), and dust (kg km$^{-2}$ yr$^{-1}$) used in our GEOS-Chem simulation. The trends are calculated from the Generalized Least Squares regression of monthly time series values over 2005-2015.
**Figure 2:** Seasonal mean UVAI values for the 2005-2015 period as observed by OMI for MAM (May, April, March), JJA (June, July August), SON (September, October, November), and DJF (December, January, February). Gray indicates persistent cloud fraction greater than 5%.
Figure 3: Seasonal mean UVAI values for the 2005-2015 period from our simulation coincidently sampled from OMI for MAM (May, April, March), JJA (June, July August), SON (September, October, November), and DJF (December, January, February). Gray indicates persistent cloud fraction greater than 5%.
Figure 4: Trends in OMI (top panel) and simulated (bottom panel) UVAI values coincidently sampled from OMI calculated from the Generalized Least Squares regression of monthly time series values over 2005-2015. The opacity of the colors indicates the statistical significance of the trend. Gray indicates persistent cloud fraction greater than 5%.
Figure 5: Seasonality of the trends in OMI UVAI values calculated from the Generalized Least Squares regression of monthly time series values over 2005-2015 for MAM (May, April, March), JJA (June, July August), SON (September, October, November), and DJF (December, January, February). The opacity of the colors indicates the statistical significance of the trend. Gray indicates persistent cloud fraction greater than 5%.
Figure 6: Seasonality of the trends in simulated UVAI values coincidently sampled from OMI calculated from the Generalized Least Squares regression of monthly time series values over 2005-2015 for MAM (May, April, March), JJA (June, July August), SON (September, October, November), and DJF (December, January, February). The opacity of the colors indicates the statistical significance of the trend. Gray indicates persistent cloud fraction greater than 5%.
Figure 7: Trend in GEOS-Chem aerosol concentrations for a) secondary inorganic aerosol (SIA), b) dust, c) total organic aerosol (OA), d) brown carbon (BrC), e) black carbon (BC), and f) sea salt. The trends are calculated from the GLS regression of monthly aerosol concentration time series values over 2005-2015. The opacity of the colors indicates the statistical significance of the trend. Gray indicates persistent cloud fraction greater than 5%.
Figure 8: Annual mean change in simulated UVAI values for 2008 due to the doubling of concentrations of a) secondary inorganic aerosol (SIA), b) dust, c) total organic aerosol (OA), d) brown carbon (BrC), e) black carbon (BC), and f) sea salt from the GEOS-Chem simulation. Gray indicates persistent cloud fraction greater than 5%.
Figure 9: Change in simulated UVAI values due to the 2005-2015 trends in a) secondary inorganic aerosols (SIA), b) dust, c) total organic aerosol (OA), d) brown carbon (BrC), e) black carbon (BC), and f) sea salt from the GEOS-Chem simulation. Gray indicates persistent cloud fraction greater than 5%.