Evaluation of climate model aerosol seasonal and spatial variability over Africa using AERONET

Hannah M. Horowitz¹, Rebecca M. Garland²,³, Marcus Thatcher⁴, Willem A. Landman²,⁵, Zane Dedekind², Jacobus van der Merwe², Francois A. Engelbrecht²,⁶

¹Department of Earth & Planetary Sciences, Harvard University, Cambridge, MA, 02138, USA
²Natural Resources and the Environment Unit, Council for Scientific and Industrial Research, Pretoria, 0001, South Africa
³Climatology Research Group, North West University, Potchefstroom, 2520, South Africa
⁴Marine and Atmospheric Research, Commonwealth Scientific and Industrial Research Organisation, Melbourne, 3195, Australia
⁵Department of Geography, Geoinformatics and Meteorology, University of Pretoria, Hatfield, 0028, South Africa
⁶School of Geography, Archaeology and Environmental Studies, University of the Witwatersrand, Johannesburg, 2000, South Africa

Correspondence to: hmhorow@post.harvard.edu

Abstract. The sensitivity of climate models to the characterization of African aerosol particles is poorly understood. Africa is a major source of dust and biomass burning aerosols and this represents an important research gap in understanding the impact of aerosols on radiative forcing of the climate system. Here we evaluate the current representation of aerosol particles in the Conformal Cubic Atmospheric Model (CCAM) with ground-based remote retrievals across Africa, and additionally provide an analysis of observed aerosol optical depth at 550 nm (AOD$\text{550nm}$) and Ångström exponent data from thirty-four Aerosol Robotic Network (AERONET) sites. Analysis of the 34 long-term AERONET sites confirms the importance of dust and biomass burning emissions to the seasonal cycle and magnitude of AOD$\text{550nm}$ across the continent and the transport of these emissions to regions outside of the continent. In general, CCAM captures the seasonality of the AERONET data across the continent. The magnitude of modeled and observed multi-year monthly average AOD$\text{550nm}$ overlap within ± 1 standard deviation of each other for at least 7 months at all sites except the Reunion Island site. The timing of modeled peak AOD$\text{550nm}$ in southern Africa occurs one month prior to the observed peak, which does not align with the timing of maximum fire counts in the region. For the western and northern African sites, it is evident that CCAM currently overestimates dust in some regions while others (e.g., the Arabian Peninsula) are better characterized. This may be due to overestimated dust lifetime, or that the characterization of the soil for these areas needs to be updated with local information. The CCAM simulated AOD$\text{550nm}$ for the global domain is within the spread of previously published results from CMIP5 and AeroCom experiments for black carbon, organic carbon and sulfate aerosols. The model’s performance provides confidence for using the model to estimate large-scale regional impacts of African aerosols on radiative forcing, but local feedbacks between dust aerosols and climate over northern Africa and the Mediterranean may be overestimated.
1 Introduction

Africa contains the largest single sources of biomass burning emissions and dust globally (Crutzen and Andreae, 1990; van der Werf et al., 2010; Schütz et al., 1981; Prospero et al., 2002). Dust aerosols and carbonaceous aerosols produced from biomass burning are known to impact climate through direct scattering and absorption of radiation, and indirectly through their effects on cloud formation and properties. Black carbon is estimated to be second only to CO₂ in contributing to warming globally (Bond et al., 2013). Currently, the largest uncertainty in climate models is the impact of aerosols on the radiative balance of the Earth (Boucher et al., 2013).

Mineral dust emitted into the atmosphere primarily originates in topographic depressions (Prospero et al., 2002), consistent with the acceleration of winds in between mountains and plateaus (Evan et al., 2016). Meteorology plays a key role in the seasonality of dust emissions and transport in Africa. Latitudinal changes in the large-scale circulation, including the intertropical convergence zone (ITCZ) and the African monsoon, shift the location of maximum dust activity and transport of dust northward (~5°N to ~20°N) from winter through summer (Jankowiak and Tanre, 1992; Moulin et al., 1997; Prospero et al., 2002; Schepanski et al., 2009; Leon et al., 2009). The movement of the ITCZ also determines the seasonality of precipitation, and so determines the onset and severity of dry season biomass burning in Africa. Most fires in Africa are set by humans during the dry season for agricultural practices, when there is a near absence of convection and lightning (e.g., Swap et al., 2003; Archibald, 2016). Maximum biomass burning activity thus shifts from June–September in southern Africa, to December–February in sub-Saharan northern Africa (Haywood et al., 2008; Duncan et al., 2003; Cooke et al., 1996). The magnitude of emissions in a given biomass burning season is largely determined by the amount of rainfall preceding burning (which is affected by climate variability including the El Niño Southern Oscillation), as this impacts the amount of vegetation that grows and can be burned (Swap et al., 2003; Anyamba et al., 2003; van der Werf et al., 2004). Biomass burning emissions in southern Africa contribute an estimated 86% of total carbonaceous aerosols emitted in Africa, which is a higher percentage than other regions worldwide (Bond et al., 2004). In many places, biomass burning aerosols dominate the seasonal cycle of the aerosol column in the region (Tesfaye et al., 2011; Queface et al., 2011; Sivakumar et al., 2010; Eck et al., 2003), which in turn can have a significant impact on the regional climate (Abel et al., 2005; Winkler et al., 2008; Tummon et al., 2010). Although these two sources dominate total column aerosol in Africa, fine anthropogenic aerosols are also observed, including at sites in the Sahara desert and off the coast of northern Africa (Rodriguez et al., 2011; Guirado et al., 2014).

In addition to the local and regional effects of African dust and biomass burning aerosols near emission sources, these aerosol particles can also be transported long distances to impact other regions. Saharan dust is exported over the Atlantic Ocean, cooling the tropical North Atlantic and influencing Atlantic climate variability (Evan et al., 2011; Doherty and Evan, 2014). Climate change may reduce future dust emissions, thus leading to a positive warming feedback over the North Atlantic (Evan et al., 2016). Saharan dust significantly enhances nutrient transport to regions like the Amazon rainforest, which may also have a feedback on climate (e.g., Bristow et al., 2010; Yu et al., 2015). Over southern Africa, massive
aerosol plumes during peak biomass burning are exported in a so-called “river of smoke” off the southeastern coast of southern Africa to the Indian Ocean, as well as over the southwestern coast over Angola out to the Atlantic Ocean (Garstang et al., 1996; Tyson et al., 1996a; Tyson et al., 1996b; Swap et al., 2003). This latter exit pathway aligns with the stratocumulus cloud deck that forms off of the southwestern coast and has motivated multiple recent and on-going ground-based and aircraft campaigns (Zuidema et al., 2016). The simulation of this cloud deck with the AeroCom intercomparison of global models was found to differ significantly between models, and to be the area of highest uncertainty in modeling aerosol radiative forcing (Stier et al., 2013). An assessment of the first phase of AeroCom showed that the largest model differences were from dust and carbonaceous aerosols (Kinne et al., 2006), the dominant aerosol constituents over Africa. Additionally, this AeroCom experiment highlighted an overestimation of dust at northern African sites in winter (Kinne et al., 2006). An accurate representation of African aerosols is critical in climate models to understand the regional and global radiative forcing and climate impacts of dust and biomass burning aerosols, at present and under future climate change, and is currently a major challenge.

This study performs the first evaluation of the representation of African aerosols in the Conformal Cubic Atmospheric Model (CCAM) (McGregor, 2005). The CCAM aerosol parameterizations are based on the CSIRO Mk3.6 climate model used in the fifth Coupled Model Intercomparison Project (CMIP5) to estimate radiative forcing for the Intergovernmental Panel on Climate Change AR5. CCAM will be included as part of a coupled earth system model, the Variable Resolution Earth System Model (VRESM), towards the South African Council of Scientific and Industrial Research (CSIR) submission to CMIP6. We evaluate CCAM using the CMIP5 emissions inventory against long-term aerosol optical depth (AOD) retrievals across Africa and outflow regions off the coast from the Aerosol Robotic Network (AERONET) (Holben et al., 1998). A particular emphasis is placed on evaluating the long-term seasonal variability at sites heavily impacted by dust and biomass burning aerosol particles. CCAM simulates four prognostic aerosol species (organic carbon (OC), black carbon (BC), sulfate, and dust) and diagnostic (i.e. prescribed) sea salt aerosols, and their individual contributions to total AOD. Detailed case studies at six sites across Africa are used to examine the modeled source distribution of AOD and to understand the model processes, determining how well CCAM represents the observational data. The evaluation of aerosols in CCAM against observations has implications for its estimates of radiative forcing.

2 Methods

2.1 CCAM model description

CCAM is a global atmospheric circulation model, and was run at a quasi-uniform resolution of 50 km in the horizontal and with 27 levels in the vertical. The simulations applied in this study form part of the CSIR’s contribution to the Coordinated Regional Downscaling Experiment (CORDEX) of the World Climate Research Programme (WCRP). Horizontal wind and temperature upwards of 900 hPa and the surface pressure in CCAM were nudged towards the ERA-Interim reanalysis data (Dee et al., 2011). This nudging was applied every 6 hours at a length scale of ~2250 km using the
digital filter of Thatcher and McGregor (2009). The sea-surface temperature and sea-ice from ERA-Interim were used as lower boundary forcing; these values were interpolated to the CCAM grid with the differences in the land-sea mask taken into account. For this study, 6-hourly model output was regridded to 0.5° x 0.5° resolution over the African continent (40°N to 40°S, 20°W to 60°E) from 1999 to 2012, the period for which most AERONET observations are available for comparison. The simulation was initialized in 1979 such that prognostic soil variables like temperature and moisture, in addition to aerosol fields, were sufficiently spun up.

The aerosol parameterization in CCAM has been documented in detail elsewhere (Rotstayn et al., 2007; Rotstayn et al., 2010; Rotstayn et al., 2011; Rotstayn et al., 2012). In summary, the aerosol scheme is a bulk / mass scheme (i.e. single moment) to represent the sulfur cycle, carbonaceous aerosols, dust, and diagnosed sea salt. Carbonaceous aerosols are represented by separate prognostic species for organic carbon (OC) and black carbon (BC). Sea salt concentrations above the ocean surface are diagnosed (i.e. prescribed) at each time step as a function of the 10-m wind speed. It is assumed that sea salt aerosols are well-mixed in the marine boundary layer (MBL), and that the concentration is zero above the MBL. There are two size bins of sea salt aerosols (mode radii of 0.035 µm and 0.35 µm). As the sea salt concentrations are prescribed at each time step, they are not actively emitted, transported or removed, and thus no sea salt is transported over land (Rotstayn et al., 2007).

The atmospheric model determines the transport of the prognostic aerosol species (sulfate, carbonaceous, and dust aerosols), including turbulent mixing in the boundary layer and transport due to convection. Wet scavenging processes are included, with linked to warm rain and frozen precipitation processes in the cloud microphysics parameterizations and the convection scheme (Rotstayn et al., 2007). The model also accounts for both direct and indirect aerosol effects, representing an important feedback into the atmospheric simulation. The semi-direct effect is also included in CCAM; however, as the vertical temperatures upwards of 900 hPa are nudged towards the ERA-Interim reanalysis data every six hours in accordance with CORDEX, the semi-direct impact on the simulation presented here is diminished.

The size distribution of the sulfate, OC and BC aerosol particles is represented by a mode radius with a geometric standard deviation. Dust is represented by four size bins with radii of 0.1-1, 1-2, 2-3 and 3-6 µm, with the parameterization of aeolian dust emissions closely based on Ginoux et al. (2001) and Ginoux et al. (2004) (see also Rotstayn et al., 2011). Specifically, dust emissions are described by the expression,

\[
F_p = CS_p u_{10m}^2 (u_{10m} - u_t) \text{ (if, } u_{10m} > u_t)
\]

where \(F_p\) is the flux (µg s\(^{-1}\) m\(^{-2}\)), \(C\) is a dimensional factor set to 0.5 µg s\(^{-2}\) m\(^{-2}\), \(S_p\) is a fraction for each dust size bin following Ginoux et al. (2001), \(u_{10m}\) is the horizontal wind speed (m s\(^{-1}\)) and \(u_t\) (m s\(^{-1}\)) is the threshold velocity, which accounts for soil moisture and the particle size. If \(u_{10m}\) is not greater than \(u_t\), then \(F_p = 0\). For this study, the dimensional factor \(C\) was set to be smaller than that used by Ginoux et al. (2001), which has the effect of reducing the dust emissions for the same wind speed and soil moisture.

Emissions of OC, BC and SO\(_2\) from anthropogenic and biomass burning sources are from the CMIP5 recommended historical emissions datasets through the year 2000 (Lamarque et al., 2010) and extend through 2012 using emissions from
the RCP4.5 modest mitigation scenario (Moss et al., 2010; Riahi et al., 2007). Aerosol emissions across the RCP scenarios through the latest year simulated here (2012) are similar (van Vuuren et al., 2011). Within CCAM, of the SO$_2$ emissions from fossil fuel and smelting, 3% are emitted as sulfate directly (Rotstayn and Lohmann, 2002); a similar fraction is assumed in other global models to represent rapid in-plume transformation of SO$_2$ to sulfate (Liu et al., 2005; Chin et al., 2000; Koch et al., 1999). The model has three prognostic variables to represent the sulfur cycle: Dimethylsulfide (DMS), SO$_2$ and sulfate. Additional minor sources of model sulfate aerosol are volcanic SO$_2$ emissions and biogenic DMS emissions, which can be oxidized to sulfate (Rotstayn and Lohmann, 2002). Concentrations of sulfur oxidants (OH, NO$_3$, H$_2$O$_2$ and O$_3$) are prescribed, with the amount of SO$_2$ dissolved into cloud water described by Henry’s Law.

Within the CMIP5 emissions used, anthropogenic and biomass burning sources vary decadally, and during the 2005–2012 period forced by RCP4.5 they vary every 5 years. Biomass burning emissions also have a monthly varying annual cycle, while non-biomass burning anthropogenic emissions remain constant annually. Thus, changes in modeled aerosol loading using the CMIP5 emissions on smaller than monthly temporal scales for OC, BC, and sulfate, and interannual variability within a given decade, are not due to changes in sources, but instead changes in transport and deposition sinks resulting from meteorological variability. An earlier study over southern Africa during the biomass burning season found that a chemical transport model was able to reproduce day-to-day variability in AOD using time-invariant emissions, suggesting meteorological variability is more important on this timescale than emissions (Myhre et al., 2003).

Prognostic aerosol species for hydrophobic and hydrophilic forms of OC and BC are transported separately in CCAM. Hydrophobic OC and BC are non-hygroscopic, while hydrophilic species’ hygroscopic growth is based on Köhler theory. The model assumes fossil fuel emissions are 50% hydrophilic, and biomass and biofuel burning are 100% hydrophilic. Conversion from hydrophobic to hydrophilic follows Cooke et al. (1999) with an e-folding lifetime of 1.15 days. Secondary organic aerosol (SOA) formation is not treated in the model. All prognostic aerosol species are removed via wet and dry deposition, while dust is additionally removed through gravitational settling (Rotstayn and Lohmann, 2002; Lohmann et al., 1999; Ginoux et al., 2001).

### 2.2 AERONET observational data

The global network of AERONET stations measure aerosol optical properties at multiple wavelengths ranging from the UV to shortwave infrared using a ground-based Cimel sun-photometer (Holben et al., 1998). For this work, the retrieved AOD at 440 nm (AOD$_{440\text{nm}}$), and the Ångström exponent of extinction for 440 nm to 870 nm ($\alpha_{\text{ext}(440/870)}$) from AERONET were used.

The Ångström exponent of extinction is the negative slope of the natural log of AOD with wavelength. The AOD$_{440\text{nm}}$ was adjusted to 550 nm using the $\alpha_{\text{ext}(440/870)}$ for comparison to modeled AOD at 550 nm following Eq. 2, where $\tau_{440}$ is AOD at 440nm retrieved by AERONET, and $\tau_{550}$ is AOD at 550nm:

$$\tau_{550} = \tau_{440} \left( \frac{550}{440} \right)^{-\alpha_{\text{ext}(440/870)}}$$
A climatology of AOD$_{550nm}$ and $\alpha_{\text{ext}(440/870)}$ observations from 34 sites in Africa and the Middle East outside of heavily urbanized areas with at least 1 full year of level 2.0 data (cloud-screened, and manually inspected for quality assurance; (Smirnov et al., 2000) (see Fig. 1 and Tables 1 and 2) is developed. Sites were selected in southern Africa that could characterize the model performance in regions dominated by biomass burning aerosol were selected, and in northern and western Africa and the Middle East that could characterize the model representation of Saharan and Sahelian dust sources and outflow. This analysis includes sites in the Mediterranean and Europe influenced by North African dust outflow (Basart et al., 2009; Toledano et al., 2007a; Toledano et al., 2007b; Querol et al., 2009; Pace et al., 2006).

For the comparison with model outputs, sites with multiple years of complete data for most of the annual cycle (see Sect. 2.3 and Fig. 3a and 3b) were selected. Where multiple sites were proximal to each other and showed similar features, the site with the longest data record was selected to be representative of the sites and was used for comparison to the model (see site names in bolded font in Fig. 1, and Tables 1 and 2). This selection results in 23 sites being chosen and used in the comparison with model outputs. Daily average values, calculated for days with at least 3 measurements, were downloaded from the AERONET website (http://aeronet.gsfc.nasa.gov) and used in this analysis.

### 2.3 Model-observation comparisons

Monthly-average time series and multi-year monthly mean climatology of AOD$_{550nm}$ were calculated for each site for observed and modeled data. The 550 nm wavelength is representative of the model AOD output. Modeled Ångstrom exponent is not available. The AERONET monthly average AOD$_{550nm}$ was calculated from the daily averages using a 70% data completeness rule (i.e., if more than 30% of the daily values were missing, a monthly average was not calculated for that time period). A multi-year mean seasonal cycle was also calculated from daily averages for each month for all available years of data at each site, following the same data coverage exclusions. This is to ensure that the observed monthly averages were representative of the entire month to provide a relevant comparison for modeled output, as it is difficult for climate models to represent specific days individually (e.g., Magi et al., 2009), and as CCAM used CMIP5 emissions that do not vary daily.

Daily average AOD from AERONET is calculated for a minimum of 3 time points from sun photometer measurements, which can only be made during daytime, while modeled AOD is reported at 6-hourly resolution. Therefore, only CCAM AOD between 06:00 and 18:00 UTC was averaged for monthly and multi-year means (similar to other AERONET-model comparison studies; (e.g., Tegen et al., 2013). Model monthly means were, however, insensitive to the choice of daylight cut-off (see Fig. 2), which gives confidence that the instantaneous 6-hourly values from CCAM can represent the range of daytime hours sampled by AERONET. Multi-year CCAM seasonal cycles were calculated from daily averages at each site from 1) only the specific months with valid observational data, and 2) all months of all model years (1999–2012). As many of the observational sites do not have continuous data, nor are the sampling times across sites always overlapping, the two calculations of modeled multi-year seasonal cycles were compared to test whether the entire model time period (1999–2012) for each month could be used to evaluate modeled spatial patterns against all available sites (Section 4.2.5).
Modeled and observed AOD$_{550\text{nm}}$ at each site were compared on a monthly timescale using a variety of metrics to quantify how well the model captures seasonal and interannual variability, and overall magnitude. To this end, the Pearson’s correlation coefficient between the model and observations ($r$), Normalized Mean Bias (NMB) of the model as a percentage of the observed values, and the Mean Absolute Error (MAE) of the model in units of AOD$_{550\text{nm}}$ were calculated.

We also compare modeled daily average AOD$_{550\text{nm}}$, using the same daylight hours previously described, to observed AERONET daily average AOD$_{550\text{nm}}$ for the specific days with available data at each site. As described in Section 2.1, outside of the dust parameterization, the experimental setup of the model following CMIP5 does not take daily variations in emissions into account, and thus the daily variation in modeled AOD from all other aerosol types will be due to daily variations in transport and removal only. Even with these limitations, the daily comparison is useful for further investigating model biases.

3 Climatology of AERONET AOD and $\alpha_{\text{ext}}$ observations over Africa: seasonal variability and drivers

AERONET and CCAM AOD are all reported at 550 nm. Additionally, the Ångström exponent of extinction ($\alpha_{\text{ext}}$) from AERONET reported here is from the 440 nm and 870 nm wavelength pair. Figures 3a and 3b show a compilation of multi-year monthly mean observed AOD and Fig. 4a and 4b $\alpha_{\text{ext}}$ for the 34 study sites, ordered by region from north to south. The symbols are the multi-year mean values, and the whiskers represent $\pm1$ standard deviation. The number of years of AERONET data used per month is shown at the top of each plot. The Ångström exponent is an empirical proxy related to the relative contribution to optical thickness from coarse vs. fine aerosols, with values varying between approximately 0 for pure coarse dust particles to 2 for predominantly fine particles (Leon et al., 2009; Hamonou et al., 1999). In Fig. 4a and 4b, values of $\alpha_{\text{ext}}$ below 0.4 are indicative of aerosols dominated by coarse particles (e.g., mineral dust or coarse sea salt particles) (shaded gray area), while higher values show a contribution from predominantly fine, submicron aerosols, indicative of biomass burning or anthropogenic sources (Holben et al., 2001; Ogunjobi et al., 2008; Rajot et al., 2008).

Table 1 displays the multi-year daily average and median AOD and $\alpha_{\text{ext}}$, which were calculated using all available data points per site. Table 2 displays the maximum and minimum multi-year monthly average per site together with the month when that value was observed. The amount of data is not equal at all sites, nor were the sampling periods at all sites overlapping, and thus detailed comparisons of the sites are not possible. Instead, we focus on overall regional patterns, including timing of peaks and minima.

3.1 Northern Africa and Middle East AERONET AOD and $\alpha_{\text{ext}}$ observations

The mean AERONET AOD in the northern African and Middle East sites (Table 1, blue in Fig. 1 and Fig. 3a) range from 0.06–0.49 and medians range from 0.02 – 0.39. The maximum multi-year monthly average values range 0.15–0.69, and minima range 0.015–0.36 (Table 2). The average $\alpha_{\text{ext}}$ range 0.49–1.04 and the medians range 0.41-1.05. The multi-year
monthly average maxima in $\alpha_{\text{ext}}$ range 0.73–1.59 and minima range 0.17–0.96. The spread of $\alpha_{\text{ext}}$ values suggests a mixture of fine and coarse aerosols at these sites.

The impact of coarse particles on the aerosol loading is observed in this region. Ras El Ain, Ouarzazate, La Laguna, Dahkla, Solar Village, Mezaira, Hamim, and Tamanrasset INM have multi-year monthly average $\alpha_{\text{ext}}$ below the 0.4 “coarse particle” threshold, and all other sites pass this threshold within the standard deviation from the multi-year mean except for El Arenosillo (Fig. 4a). This may be due to the influence of local industrial pollution sources at El Arenosillo (Toledano et al., 2007a; Toledano et al., 2009). While low values of $\alpha_{\text{ext}}$ could represent other coarse particles besides dust like sea salt, previous work has indicated sea salt is a minor contributor to aerosols at island sites to the north of Africa, including Izaña (Rodríguez et al., 2011; Putaud et al., 2000; Querol et al., 2009). The correspondence of the seasonality in $\alpha_{\text{ext}}$ and AOD with known dust events suggests mineral dust is the primary contributor to extinction from coarse particles. The maximum AOD occurs across most sites during June–August, and coincides with a decrease in $\alpha_{\text{ext}}$. This is later than the AOD peak at the western African sites (Sect. 3.2). This delay and corresponding change in $\alpha_{\text{ext}}$ suggest that transported dust from the Sahara leads to the higher observed AOD. Thus, the seasonal variation in the location of the ITCZ and associated northward shift in dust transport may be responsible for the shift in timing of maximum AOD between the western and northern African sites. AOD at most of the Middle Eastern sites (Eilat, Sede Boker, IASBS, KAUST, and Solar Village) peaks earlier, in March through May, indicative of different seasonality of the local dust sources in the Arabian peninsula (Basart et al., 2009).

The greatest seasonal differences in $\alpha_{\text{ext}}$ occur at Hamim, where in addition to high local dust emissions in spring and summer, regional circulation transports dust from deserts in Iraq and Southern Iran during summer and a mixture of fine pollution aerosols from the Persian Gulf throughout the year (Eck et al., 2008; Basart et al., 2009). The Izaña site has a different seasonal pattern in $\alpha_{\text{ext}}$ than its neighboring two sites, La Laguna and Santa Cruz, on the same island. It is, however, the highest elevation site in our study at 2391 m, 1800–2300 m higher than La Laguna and Santa Cruz (see Table 1). Local topography, meteorology, or transport patterns affecting the sinks and sources reaching Izaña may lead to a different aerosol size distribution.

### 3.2 Western Africa AERONET AOD and $\alpha_{\text{ext}}$ observations

The highest AERONET AOD across all sites is observed in western Africa (denoted in red in Fig. 1 and Fig. 3b). The overall mean AOD ranges 0.44–0.67 and the median values range 0.37–0.56 (Table 1). AOD peaks at 0.62–1.10, and minimum AOD range 0.26–0.38 (Table 2). The minimum AOD values seen here are similar to the maximum AOD values seen in northern and southern Africa. The western African sites also have low average $\alpha_{\text{ext}}$ (0.29–0.66) and median values (0.22–0.59) (Table 1). The maximum $\alpha_{\text{ext}}$ ranges 0.52–0.96, and minia range 0.092–0.33 (Table 2). The maximum multi-year monthly average $\alpha_{\text{ext}}$ occurs in December across all western Africa sites, while the minimum values vary in timing (Table 2).

In general, as AOD increases, $\alpha_{\text{ext}}$ decreases (Figs. 3b and 4b), which would suggest that the variation in the AOD is dominated by the variation in coarse aerosol particles, most likely dust. A similar relationship was found previously for Banizoumbou (Holben et al., 2001; Ogunjobi et al., 2008; Rajot et al., 2008). This relationship is prominent at Agoufou,
Banizoumbou, Zinder Airport, Maine Soroa, and Ougadougou. In addition, this relationship is seen in January–June in Djougou, while in October–December the increase in AOD at this site corresponds to an increase in $\alpha_{\text{ext}}$. In Ilorin, which is south of the other sites, the AOD peaks in January–March, while the $\alpha_{\text{ext}}$ is at a minimum value in March–May. Previous work found that minimum values of $\alpha_{\text{ext}}$ are related to dust storms at Ouagadougou, Dakar, and Agoufou, and clearly linked to dust at Ilorin and Banizoumbou based on air mass back trajectories and observed seasonality (Ogunjobi et al., 2008). While Dakar is frequently influenced by air transported over the Atlantic Ocean (Ogunjobi et al., 2008), analysis off the coast of Dakar at Cape Verde found the AOD and aerosol mass loading were dominated by desert dust, with sea salt minimally contributing to AOD (6%) in part due to its small extinction (Chiapello et al., 1999) which would also imply a minor influence on $\alpha_{\text{ext}}$.

The timing of peak monthly-mean AOD varies between February–March for the Banizoumbou, Ouagadougou, Djougou, and Ilorin sites, and May–June for the Agoufou, Dakar, Zinder Airport, and DMN Maine Soroa sites, approximately following a south to north gradient. The latitudinal movement of dust transport northward from winter (i.e. February–March) to summer (i.e. May–June), thus appears to dictate the seasonal cycle in AOD at these sites, consistent with a previous regional dust model-AERONET comparison at Dakar, Agoufou, and Banizoumbou (Tegen et al., 2013).

Ilorin and Djougou, the most southerly sites in this region, have slightly higher $\alpha_{\text{ext}}$ on average (0.66 ± 0.36 and 0.52 ± 0.34, respectively), especially during late fall to early winter (peaking at ~0.9 in December). This coincides with the sub-Saharan Northern Africa biomass burning season (December–February) (e.g., (Roberts et al., 2009; Giglio et al., 2006). The highest AOD during December–February out of the western African sites is also observed at Ilorin and Djougou (up to a peak of 1.10 in February at Ilorin), which are closer to the primary area of biomass burning during this time (Lioussse et al., 2010; Pinker et al., 2010). This suggests that biomass-burning aerosols could make up a larger fraction of total AOD at Ilorin and Djougou than elsewhere during this time period, and explains the different relationship in the seasonality of $\alpha_{\text{ext}}$ and AOD at these two sites.

Dakar has the smallest month-to-month variability in AOD, ranging from 0.30–0.62. Leon et al. (2009) find that Dakar is subject to transport of both dust and biomass burning aerosols, depending on the season, as well as poorly constrained anthropogenic emissions from the city and other nearby urban centers. This variety of sources, the site’s greater distance from dust and biomass burning aerosol sources, and proximity to anthropogenic emissions that have lower seasonal variability, may explain its observed seasonal cycle.

### 3.3 Southern Africa AERONET AOD and $\alpha_{\text{ext}}$ observations

The average AERONET AOD in the southern African sites ranges 0.064–0.21 and the medians range 0.06–0.14, with multi-year monthly maximum AOD peaking at 0.095–0.50 and minimum AOD ranging 0.046–0.13 (Tables 1 and 2). The region has larger $\alpha_{\text{ext}}$, with averages ranging 0.7–1.6 and medians ranging 0.66–1.75. The maximum monthly average $\alpha_{\text{ext}}$ ranges 1.12–1.85 and the minima range 0.28–1.14. Mongu and Skukuza in southern Africa have the highest observed $\alpha_{\text{ext}}$, respectively.
indicating little influence from coarse aerosols and confirming the importance of biomass burning as an aerosol source in this region.

Previous studies have shown AOD is highest in this region during the biomass burning season, from AERONET AOD through the year 2007 at Mongu and Skukuza (Queface et al., 2011) and MISR satellite data over South Africa (Tesfaye et al., 2011). Mongu is situated in Zambia in the middle of the biomass burning source region in southern Africa (e.g., Swap et al., 2003; Eck et al., 2003; Edwards et al., 2006; Queface et al., 2011). For Southern Hemisphere Africa, peak fire activity typically occurs in June through October, with a shift in general toward later months moving from north to south, except in the winter rain areas of southwestern South Africa (Archibald et al., 2010; Giglio et al., 2006). At Ascension Island, the transport of biomass burning aerosols from southern Africa west over the Atlantic Ocean is observed in the seasonal cycle of \( \alpha_{\text{ext}} \) and AOD (Figs. 3b and 4b), as both peak in September, which is the timing of climatological peak AOD and peak biomass burning at Mongu (Giglio et al., 2006). This known transport pathway off the coast of Angola (Garstang et al., 1996) is also seen in the AOD and \( \alpha_{\text{ext}} \) observed at Etosha Pan, but peak values occur in October as opposed to September. However, these values at Etosha Pan may not represent a long-term mean seasonal cycle as only one year of data was available at this site during the time period of our study.

The AOD at Skukuza also peaks in September, indicating transport of biomass burning aerosols southeast over the site and exiting the continent toward the Indian Ocean, consistent with the so-called “river of smoke” or major export pathway off the coast of southeastern South Africa (e.g., Swap et al., 2003). The eventual transport of biomass burning aerosols from southern Africa over Reunion Island is indicated in the seasonal cycle of \( \alpha_{\text{ext}} \) and AOD, which increase toward an October peak.

The continental sites closest to the region of burning have sustained and relatively constant high values of \( \alpha_{\text{ext}} \) during April–October (Fig. 4b). This is especially evident at Mongu. The \( \alpha_{\text{ext}} \) at all southern Africa sites declines in austral spring and summer. While these small variations in \( \alpha_{\text{ext}} \) alone are not enough to distinguish aerosol size distributions, they are consistent with results from MISR for the central South African region (including Skukuza) that showed an increase in the coarse mode fraction in summer due to dust from the Northern Cape and Namibian desert regions (Tesfaye et al., 2011).

4. Model evaluation

4.1 Annual model aerosol budgets

Annual burdens, deposition, wet deposition fraction, lifetime, and emissions for each of the four prognostic aerosol species in 2010 are shown in Table 3 for the globe and the Africa domain (40°S to 40°N, 20°W to 60°E), separately. These values are compared to estimates from other present-day models and the CMIP5 and AeroCom experiments in Fig. 5. CCAM is within the range of global present-day annual aerosol burden estimates from models in the CMIP5 and AeroCom experiments for BC, OC, and sulfate. In addition, in Fig. 5b–c, CCAM is within the range of estimates for total deposition, wet deposition fraction, burden, and lifetime of organic aerosols (OA) and BC (Tsigeridis et al., 2014; Allen and
Landuyt, 2014). CCAM modeled OC emissions and burden is converted to OA by multiplying by a factor of 1.4 for a consistent comparison (Tsigaridis et al., 2014). In general the CCAM values for BC burden and lifetime are higher than the CMIP5 median values, but are well within the range of models. For OA, CCAM is close to median estimates from the AeroCom Phase II models with the exception of OA/OC lifetime, which is at the high end of all models.

While CCAM performs well compared to other models for BC, OC and sulfate, CCAM has a dust burden (68 Tg) ~2–7 times higher than AeroCom Phase I models (Huneeus et al., 2011) and all available dust modeling results summarized in a recent review (Kinne et al., 2006; Zender et al., 2004) (see Fig. 5a, 5d). In the CCAM model, annual dust emissions over the Africa region alone (40°S to 40°N, 20°E to 60°W) in 2010 are 2320 Tg yr⁻¹, contributing 83% of global total modeled dust emissions. The range from AeroCom models is 35–77.9% of global dust emissions (Huneeus et al., 2011). Global dust emissions (Fig. 5) are above the mean, but within the range of AeroCom models. This together with an overestimation of dust in Africa would lead to a large percentage contribution of global dust emissions from Africa.

The global dust emissions, burden, wet deposition, dry deposition and sedimentation, and lifetime are compared to AeroCom experiments in Fig. 5d (Huneeus et al., 2011). The modeled dust lifetime (8.9 days) is longer than models examined in Zender et al. (2004) that range from 2.8 to 7.1 days, and AeroCom Phase I that range from 1.6 to 7.1 days (Huneeus et al., 2011), indicating the sinks of dust in the model may be too low, contributing to a high global dust burden. The wet deposition (1571 Tg a⁻¹) is higher than AeroCom results (range of 295 to 1382 Tg a⁻¹, median 357 Tg a⁻¹), however the dry deposition and sedimentation (1209 Tg a⁻¹) are similar to the AeroCom median (753 Tg a⁻¹) in spite of the much higher dust burden. This overestimation of dust is discussed more in Sect. 4.2.2 and Sect. 4.2.3 below.

4.2 Evaluation of model against AERONET AOD observations: Multi-year mean seasonal cycle comparison

Figure 6 shows the same multi-year mean seasonal cycle for observed AERONET AOD as in Fig. 3 (here in red triangles), overlaid with CCAM results for all model years (dark blue) and only those months with corresponding AERONET data that met the 70% completeness cutoff (yellow). The shaded red areas are within ±1 standard deviation from the observed values, and the shaded blue areas are within ±1 standard deviation from the all model years CCAM output. In this comparison, only AERONET sites with multiple years of complete data for most of the annual cycle are included in order to compare multi-year monthly cycles from observations and the model.

The monthly cycle from CCAM considering the full model period (dark blue line) and only those years with observational data (yellow line) are similar across all sites, with only minor differences that are within ±1 standard deviation of the full model period. Thus, the full model time period (1999–2012) can be used to evaluate modeled spatial patterns against all available AERONET sites, even though the observations at different sites are from disparate time periods. All following analyses are presented using the full model time period.

For most sites, the monthly cycle (i.e. timing of peak and minimum AOD) is well-captured by CCAM, indicating the seasonality in CMIP5 emissions and the model parameterization of dust emissions is adequate. A few notable exceptions
(e.g., timing of maxima at Mongu and Ascension Island, missing winter minima in western African sites, and missing summertime peaks after observed springtime maxima at Sede Boker and Solar Village) will be investigated in Sect. 4.2.1–4.2.3 below. The magnitude of modeled and observed multi-year monthly average AOD overlap within ±1 standard deviation of each other for at least 7 months at all sites except Reunion Island, and for all observed months at 8 sites that span all three regions (Granada, Blida, Zinder Airport, Banizoumbou, Ouagadougou, Djougou, Ilorin, and Skukuza). The differences in magnitude per region will also be detailed in Sect. 4.2.1–4.2.3 below.

Figure 7 highlights two representative sites each from the northern, western, and southern regions with the most observational data available in greater detail, comparing multi-year monthly mean observed and modeled AOD, with the modeled contribution of each aerosol type (sea salt, large size bin dust (radius ≥ 1 µm), small size bin dust (radius < 1 µm), BC, OC, sulfate) to total AOD shown. Further investigation of model performance, by region, follows.

4.2.1 Southern Africa

In comparison to the other regions, the model better represents the magnitude of AOD at the southern African sites (except for Reunion Island) with a smaller normalized mean bias and mean absolute error (see Fig. 6 and Table 4). However, the timing of the modeled peak AOD at two of the sites where maximum AOD is dominated by biomass-burning (Ascension Island and Mongu) occurs 1 month too early (in August, instead of September as highlighted in Table 2). Modeled AOD at both Mongu and Skukuza remain relatively constant between August and September (Fig. 7). This is consistent with the observations at Skukuza, likely due to the greater influence of anthropogenic aerosol sources at this site. Figure 7, shows the modeled sulfate contribution (emitted from both anthropogenic and biomass burning) to total AOD is higher and that of OC (primarily emitted from biomass burning) is lower at Skukuza relative to Mongu, indicating the breakdown of model emissions sources is consistent with this explanation. There is a larger observed increase in AOD between August and September at the biomass burning source region (Mongu) and the more remote Ascension Island whose seasonality is impacted by transported biomass burning aerosol as seen in the $\alpha_{ext}$ (Fig. 4a and 4b).

This mismatch in timing of the peaks is a long-standing issue in understanding southern African biomass burning, first noted during the SAFARI-2000 measurement campaign (Swap et al., 2003). In a study of Southern Hemisphere biomass burning observed by satellite, Edwards et al. (2006) found that in southern Africa alone, peak CO and AOD lagged peak fire counts by ~1 month (late September to October vs. late August, respectively). Using a chemical transport model, they found that the residence time of CO over the region was much too short for transport patterns to explain the 1 month time lag (Edwards et al., 2006). Two recent modeling studies also found that peak AOD over Southern Hemisphere Africa lagged peak fire counts and estimates of peak biomass burning emissions using either the GFEDv2 or AMMA inventories by 1–2 months (Magi et al., 2009; Tummon et al., 2010). The CMIP5 emissions used in our CCAM model study are from GFEDv2 for year 2000 onward (van der Werf et al., 2006; Lamarque et al., 2010), which at the source region of Mongu peak in August leading to the maximum modeled AOD. The GFED inventory is based on estimates of burned area from burn scars and thermal signatures of active fires viewed by the MODIS satellite, combined with land cover data and meteorological
parameters to estimate emissions for different vegetation types (van der Werf et al., 2006; van der Werf et al., 2010). This type of method would only capture large fires that produce satellite-detectable burn scars. A recent study updated the GFED inventory to include a parameterization of fire counts, burned area, and emissions from previously missing small fires, but this did not change the seasonality in biomass burning emissions over Southern Hemisphere Africa (Randerson et al., 2012).

Burned area still peaked in August, as it increased more early in the biomass burning season than late in the season when small fires were included, and higher fuel load burns (e.g., from dense, wooded vegetation) late in the season did not lead to a compensating change in emissions (Randerson et al., 2012). The small fires parameterization still relies on detection of thermal anomalies (Randerson et al., 2012).

The observed AOD peak in September aligns with the peak in fire intensity found in the generalized fire regime of savanna-woodland in Archibald et al. (2010). The peak in fire intensity in southern Africa as well as fire size occurs later in the season than the peak in fire number, though the increase in these is not large over the season (Archibald et al., 2010). However, this does suggest that fire intensity may be an important factor to consider in modeling emissions from biomass burning in southern Africa, e.g., through the new initiative FireMIP (Hantson et al., 2016).

Table 4 displays a summary of model-observation comparison by site. The normalized mean bias of the model is negative at Mongu (-21.2%) and positive at the three other southern Africa sites, showing that overall AOD is underestimated at the biomass burning source while overestimated at receptor regions (Table 4). Figure 6 suggests the model overestimates transport of biomass burning emissions to receptor sites in particular for the months of June through August. Because the AOD in both the model and observations are smaller here than in other regions, the mean absolute error is very low (0.07–0.09) and is the lowest of all sites in this model comparison. At all sites except Reunion Island, the model captures some of the temporal variability, with highly statistically significant correlation coefficients ranging from 0.48 to 0.67. Relative to other regions, the model performs best over southern Africa in terms of mean AOD magnitude, but overestimates the transport of biomass burning aerosols to Reunion Island in June through September.

### 4.2.2 Western Africa

At the western African sites, which in the observations are dominated by dust (Fig. 4b), the model captures the overall seasonal cycle in AOD except between September and December, where the observations show a decrease at all sites except the two southernmost (Djougou and Ilorin) while the model increases (see Fig. 6). As a result, the modeled minimum AOD occurs between August and October, instead of in November–December as in the observations at Agoufou, Dakar, Zinder Airport, Banizoumbou, DMN Maine Soroa, and Ouagadougou.

Figure 7 shows in a case study for two sites, Dakar and Banizoumbou, the strong influence of dust on these sites. The increase in modeled AOD from September through December, which is not seen in the observations, is due to increases in the large dust (orange bars) and small dust (red bars) contribution. This could be due to the systematic overestimation of 10m wind speed during the dry season in the ERA-interim reanalysis, a problem common to several meteorological re-analyses in the Sahelian region (Largeron et al., 2015). Although the ERA-Interim reanalysis used in this study was found to perform
best overall against wind speed observations, it also exhibited a strong positive bias during Northern Hemisphere winter (Largeron et al., 2015). Given that the CCAM simulations are nudged within the ERA reanalysis data, this may contribute to an overestimation of wind-driven dust emissions into the CCAM atmosphere during this season (September–December).

The remainder of the shape of the seasonal cycle is captured relatively well at western African sites, with the peaks in AOD in CCAM occurring within 1 month of the peak in AERONET AOD. Only at Ilorin is the timing of the peak the same in the model and the observations. Correlation coefficients between the modeled and observation AOD are statistically significant ($r$ ranges 0.27–0.61) at all sites except Djougou (Table 4). The lack of statistically significant correlation at Djougou may in part be due to a lack of data with only 24 individual months. In most of the western African sites, the model has an overall positive normalized mean bias (ranging from 29% to 103%). The exceptions are Djougou and Ilorin, which are the two southernmost sites. Djougou and Ilorin are slightly farther away from major dust sources originating in topographic depressions (Evan et al., 2015), which are represented in the CCAM dust emissions scheme (Rotstayn et al., 2011), and have relatively small, but negative normalized mean biases (-1.3%, -12.6%, respectively). The mean absolute error for all sites ranges from 0.20–0.48, which are higher than southern Africa, but lower than northern Africa, which has lower AOD on average compared to the western African sites.

The model overestimations in AOD at western African sites closer to the dust source regions may be due to an overestimation of wind speeds. Largeron et al. (2015) found that on an annual mean scale, ERA-Interim overestimates observed 10m wind speeds by 0.27 m s$^{-1}$ in the Sahel, but this was largely a result of the wintertime overestimate mentioned previously. In fact, wind speeds during springtime and the monsoon season were underestimated in the ERA-Interim because the reanalysis did not represent large increases in wind speed from boundary layer free convection and deep convection (Largeron et al., 2015). Previously, the CSIRO Mk3 coupled global climate model (GCM) accounted for this by estimating sub-grid gustiness from both boundary layer and deep convection to increase the effective 10m wind speed used in the model dust emission parameterization (Ginoux et al., 2004). In the case of CCAM, it was found that the effective sub-grid scale winds were too high (compared to the CSIRO Mk3 simulations), possibly due to differences in vertical and horizontal resolution, as well as changes in the model physical parameterizations. This led to an overestimation of global total dust emissions that were far outside the range suggested by observations (Rotstayn et al., 2011; Rotstayn et al., 2012). Therefore, these sub-grid gustiness terms have been removed from the model version presented here. In spite of this, it is still possible that 10m winds in the model may be too high. Part of the determination of surface wind speeds in CCAM relies on the Community Atmosphere-Biosphere Land Exchange (CABLE) model estimate of surface roughness. Dust emissions additionally depend on local soil moisture and soil texture from the CABLE land surface model. Issues with modeled precipitation and wet deposition, the response of soil moisture to precipitation, and how recent changes to soil texture implemented in CABLE from the Harmonized World Soil Database affect the atmospheric simulation could all contribute to an overestimate in dust emissions and atmospheric dust concentrations.
4.2.3 Northern Africa and Middle East

Potential issues with dust emissions and transport in CCAM become more apparent when comparing to northern African AOD observations. There are substantial overestimates of the multi-year monthly mean AOD in northern Africa (see Fig. 6) of up to a factor of 8 to 42 for individual months at each site. This region has the highest normalized mean biases, with NMB over 200% at 6 of the 11 sites (see Table 4). As shown in Fig. 7 for two of the northern sites, Saada and Santa Cruz Tenerife, almost all modeled AOD in this region comes from dust. However, the observational data indicate that Saada and Santa Cruz Tenerife rarely experience low values of $\alpha_{ext}$ reaching the threshold representative of coarse dust (Fig. 4a). Thus, CCAM overestimates the contribution of dust to AOD over Saada and Santa Cruz Tenerife. The global dust burden in CCAM (67 Tg) is more than twice that of the high end of values in a recent review of global dust models as well as AeroCom and CMIP5 models (Zender et al., 2004). Global dust emissions are higher than the median but are well within the range of estimates from Zender et al. (2004) and AeroCom models (Huneeus et al., 2011) (see Fig. 5). It is possible that an overestimate of dust lifetime combined with an overestimate of dust emissions plays a major role in this issue (see Section 4.1). At the same time, over the Arabian Peninsula (Dhadnah, Solar Village, Hamim) the model performs better with the lowest mean biases across sites in northern Africa and the Middle East (Table 4), suggesting dust emissions and transport may be better characterized in this region.

However, the model does capture the monthly trends in observed AOD, with a strong peak in boreal summer and relatively lower values through rest of the year. At Saada, Santa Cruz Tenerife and Dakhla, CCAM AOD peaks in August, while the observations peak in July. Modeled and observed AOD peaks in June at Hamim and July at Blida and Dhadnah. At Tamanrasset INM, CCAM AOD also peaks in July, however there are no data for July at that site. The model output shows a higher proportion of dust AOD relative to total AOD in the summer months, especially July and August (Fig. 7), which is consistent with the observed decrease in $\alpha_{ext}$ and known northward movement of Saharan dust transport in summer from the shifting ITCZ (Jankowiak and Tanre, 1992; Moulin et al., 1997; Leon et al., 2009; Schepanski et al., 2009). The model also reproduces the increase in fine aerosol (e.g., BC and SO$_4^-$) relative to coarse dust in winter months at the two sites (Fig. 7) as implied by the increasing observed $\alpha_{ext}$ (Fig. 4a). A small impact of simulated sea salt can be seen at the Santa Cruz Tenerife site (Fig. 7) (mean AOD of 0.04). The sea salt contribution to simulated monthly AOD at 550nm from AeroCom Phase III-CTRL2015 (AeroCom Phase II Interface, 2017) ranges from negligible to greater than 0.1 at Santa Cruz Tenerife.

In spite of the high model bias, all sites in northern Africa and the Middle East have statistically significant correlations, including some of the highest correlation coefficient values (ranging from 0.23 to 0.89). At Sede Boker, which has the lowest correlation coefficient in this region, the model predicts an increase in AOD from June to August, similar to other Northern African sites, which is not observed. This discrepancy may be caused by an overestimate of Saharan dust transported to the site during summer.
4.2.4 Daily variability in modeled and AERONET AOD

Figure 8 shows probability densities for daily average AOD at each of the 23 evaluation sites, with that observed by AERONET in black and modeled in red. In general, the model at most sites has a wider and smoother distribution of AOD than that observed. This is consistent with modeling limitations from the spatial resolution, 6-hourly time resolution of nudging to reanalysis meteorological data, low time resolution of anthropogenic and biomass burning emissions, and highly parameterized dust emissions (see Section 2.1). The modeled daily AOD distribution is particularly more broad and smooth than that observed for sites in northern Africa and the Middle East, where CCAM had the largest positive model biases against observed monthly mean AOD (see Section 4.2.3; Table 4). Very low AERONET AOD is frequently observed, and high AOD events associated with dust are sporadic. Modeled dust events appear to be too frequent in this region. In addition, the model is unable to capture the very low AOD days at Lampedusa, Sede Boker, Santa Cruz Tenerife, and Dahkla. These sites had the largest modeled NMB against monthly mean observed AOD after Tamanrasset (Table 4). The modeled monthly mean AOD performs relatively well against AERONET observations at Hamim and Dhadnah relative to other sites in North Africa and the Middle East, with high correlation coefficients and low NMB (Table 4). These two sites have the highest median observed daily AERONET AOD after Tamanrasset in this region (Table 1) and in Figure 8 have a broader peak at higher AOD that the model is better able to represent. At most sites in Western Africa, modeled AOD has a bimodal distribution that is not present in the observations. The secondary peak at low values of AOD suggests the model may be able to simulate aerosol removal processes and/or the absence of dust events, but this does not occur with enough frequency. The model represents the observed daily AOD distribution well at Mongu and Skukuza, near biomass burning source, in spite of modeled emissions varying only on a monthly timescale. This is similar to previous work which suggested daily variability in observed and modeled AOD in this region was impacted mostly by meteorology (Myhre et al., 2003). However, the AERONET observations show occasional high (>1) AOD events that the model cannot represent likely due to the time averaging of emissions.

4.2.5 Spatial patterns

Figure 9 shows the multi-year monthly mean climatology of modeled (background) and observed (filled circles) AOD for March (Fig. 9a), representing high AOD at many western African sites, and September (Fig. 9b), the peak observed AOD at many southern Africa sites impacted by biomass burning (note the different scales for the two months). Panels showing the 5th and 95th percentiles of 6-hourly CCAM AOD highlight the modeled variability and additional spatial features. The scales are consistent across the maps within each month to aid in comparison and as such some high AOD values are saturated in the color scale (see legend in Fig. 9).

We take advantage of the high temporal and spatial resolution of the model to show how significantly an individual 6-hourly output, in this example within the months of March and September (Fig. 9), can depart from the multiyear monthly mean AOD. Given that the emissions of all aerosol species and their precursors (with the exception of dust) vary only on a
monthly to multi-annual timescale in CMIP5 emissions (see Sect. 2.1), the variability at the 6-hourly timescale must be a result of transport and aerosol sinks in the model (and dust emissions for western and Northern Africa). This confirms the importance of model processes driven by meteorology to modeled AOD. In Southern Hemisphere Africa, where aerosols are dominated by emissions that in the CMIP5 inventory are constant within a given month for a 5 to 10 year period, Fig. 9b shows that fine-scale temporal variability can still be represented in spite of limitations in emissions inventories, consistent with previous work in this region (Myhre et al., 2003).

In March (Fig. 9a), the discrepancy in the modeled location of maximum AOD in dust-dominated northern and western Africa and the Middle East is clear, as CCAM overestimates mean AOD at all sites in this region except Djougou and Ilorin, the two southernmost sites, which are underestimated. Given that the large-scale circulation in the model is constrained to reanalysis data, it seems unlikely that issues with large-scale transport would lead to this spatial pattern in the misrepresentation of AOD. It is more likely that the overall overestimate in total dust emissions varies regionally due to regional discrepancies in precipitation, soil texture, and soil moisture, which contribute to the surface roughness (affecting surface wind speeds, feeding into the magnitude of dust emissions) and dryness (which determines the likelihood of erosion and dust emission). Dust emissions may be especially overestimated towards the north and northwest of Africa, and may even be locally underestimated in the southern Sahel. Figure 9a also shows modeled AOD over the Arabian Peninsula is more consistent with observations, suggesting a better model representation of local dust emissions in this region (see also Sect. 4.2.3). In September, when AOD is less impacted by dust, CCAM better captures the mean AOD at the available sites in western Africa along a similar latitude band, but still significantly overestimates AOD at more northern sites. This also points to a regional overestimate in dust emissions. Modeled dust lifetime, which is longer than CMIP5 and AeroCom models (see Fig. 5 and Sect. 4.1), may also play a role and could lead to dust transported too far off the northern coast of Africa. Another climate modeling study found that a non-prognostic dust scheme resulted in dust shifted too far north, while prognostic dust simulations had too much dust transport off the coast of western Africa (Mulcahy et al., 2014), suggesting the interaction between dust and meteorology in the model may be important in the current study as well. This is an area of on-going research in CCAM.

Modeled AOD at the biomass burning source region, Mongu, is underestimated in September, as seen in Fig. 9b, but the mean modeled AOD at receptor regions like Ascension Island and Skukuza are similar to that observed. The transport in CCAM of biomass burning aerosols off the coast of Angola and southeastern Africa is visible with small enhancements in the mean modeled AOD, but is more apparent in the 5th and 95th percentile results, showing that the model captures known exit pathways for southern African biomass burning (Garstang et al., 1996; Swap et al., 2003). Figure 9b also illustrates that the transport of biomass burning aerosols from southern Africa eastward toward Reunion Island is overestimated. Overall, from this analysis and given that the lifetime of OC and BC aerosols in CCAM is more consistent with other global models from AeroCom and CMIP5 (Figure 5), biomass burning aerosol emissions and transport are relatively well represented in CCAM driven by the CMIP5 emissions inventory.
5 Conclusions

The compilation of long-term AERONET observations across Africa indicates different regimes of source types and their seasonality for northern, western, and southern Africa. The importance of dust and biomass burning aerosols in the regions, as well as the transport and long-range impact of these aerosol sources, are evident in the AOD and $\alpha_{ext}$ trends across sites.

The prognostic aerosol scheme in CCAM is a key feature in the coupled earth system model VRESM currently under development for inclusion in the CMIP6 intercomparison. An accurate representation of African aerosols is critical in climate models and this current evaluation to understand how well the scheme performs in the present-day when forced with CMIP-style emissions is essential to interpreting any future climate predictions using the model. CCAM aerosol output for OC, BC and sulfate compares well with output other CMIP5 models and AeroCom model global experiments. CCAM captures the seasonal cycle of the AOD well at most sites, with statistically significant correlation coefficients between the model and observed monthly mean timeseries of AOD at all but two sites of the 23 sites studied. The seasonal cycle at these sites is strongly influenced by dust and biomass burning aerosols, and thus CCAM is able to capture the general seasonal cycle of the emissions of dust, and the transport of dust, carbonaceous and sulfate aerosol types.

This analysis has also highlighted areas within CCAM and the emissions inventory that need further work. There is a notable shift in peak AOD one month earlier than observations in biomass burning regions. This shift has been seen in previous modeling studies, and is likely due to missing processes in the emissions inventory. Comparing to CMIP5 models and AeroCom global experiments, CCAM overestimates many dust parameters including burden and lifetime. This overestimate is also seen in the comparisons to AERONET at northern and western African sites. At the northern African sites in particular, the model has large positive normalized mean biases. The model attributes large AOD primarily to dust where the observations of the Ångström exponent and AOD suggest there is very little dust present. This is likely a combination of an overestimate of dust lifetime leading to longer-range transport of dust and higher dust burdens, and overestimated dust emissions in the northwestern Sahara. The increase in AOD in the boreal winter at western African dust-influenced sites is likely due to a high bias in ERA-Interim reanalysis wind speeds in the Sahel during this season (also present in other reanalyses). The simulation of local soil parameters and injection height in CCAM could also lead to emissions biases; testing and improvement of these fields in the CABLE land surface model in the development of VRESM may help to improve the representation of dust aerosols in Africa.

The CCAM results are consistent with state-of-the-art CMIP5 GCMs, providing confidence for using the model to study the regional impacts and linkages between African aerosols and climate change under different scenarios. In addition, CCAM can be used to downscale the CMIP5 GCMs to finer spatial scales with its variable resolution global grid, and therefore refine our understanding of aerosols in this important region.
Author contribution

H. Horowitz, R. Garland, M. Thatcher and F. Engelbrecht developed the research question and designed the experiment. H. Horowitz performed the analyses of AERONET and CCAM data. M. Thatcher developed the prognostic aerosol scheme in CCAM. W. Landman provided input into the model-observation comparisons and needed statistical test. J. van der Merwe and Z. Dedekind extracted and formatted CCAM data. H. Horowitz prepared the manuscript with input from all authors.

Competing interests

The authors declare they have no conflict of interest.

Acknowledgements

This work was supported by NRF CSUR Grant Number 9157, and a CSIR Parliamentary Grant. HH was funded through the NSF GROW with USAID RI Fellowship. We thank the PIs and their staff for establishing and maintaining the 34 AERONET sites used in this study.
References


AeroCom Phase II Interface: [http://aerocom.met.no/cgi-bin/aerocom/surfobs_annualrs.pl](http://aerocom.met.no/cgi-bin/aerocom/surfobs_annualrs.pl), Last accessed 8 September 2017


Tegen, I., Schepanski, K., and Heinold, B.: Comparing two years of Saharan dust source activation obtained by regional modelling and satellite observations, Atmos Chem Phys, 13, 2381-2390, 10.5194/acp-13-2381-2013, 2013.


van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Mu, M., Kasibhatla, P. S., Morton, D. C., DeFries, R. S., Jin, Y., and van Leeuwen, T. T.: Global fire emissions and the contribution of deforestation, savanna, forest, agricultural, and peat fires (1977-2009), Atmos Chem Phys, 10, 11707-11735, 10.5194/acp-10-11707-2010, 2010.


Table 1: AERONET site information (site names in bold font indicate those sites used in model comparison). The average (± 1 standard deviation) and median (25th and 75th percentile) values for AOD$_{550}$ and $\alpha_{440-870}$ per site are shown.

<table>
<thead>
<tr>
<th>Site</th>
<th>Lat (° N)</th>
<th>Lon (° E)</th>
<th>Elevation (m)</th>
<th>Years of data used in study</th>
<th>Multi-year daily average ± 1 sd</th>
<th>Median (25th, 75th percentile)</th>
<th>Multi-year daily average ± 1 sd</th>
<th>Median (25th, 75th percentile)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Western Africa Sites</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agoufou</td>
<td>15.35</td>
<td>-1.48</td>
<td>305</td>
<td>2003-2009</td>
<td>0.51 ± 0.41</td>
<td>0.40 (0.25, 0.65)</td>
<td>0.29 ± 0.23</td>
<td>0.22 (0.13, 0.38)</td>
</tr>
<tr>
<td>Dakar</td>
<td>14.39</td>
<td>-16.96</td>
<td>0</td>
<td>1999-2012</td>
<td>0.44 ± 0.30</td>
<td>0.38 (0.25, 0.55)</td>
<td>0.36 ± 0.25</td>
<td>0.29 (0.17, 0.50)</td>
</tr>
<tr>
<td>Zinder Airport</td>
<td>13.78</td>
<td>8.99</td>
<td>456</td>
<td>2009-2012</td>
<td>0.52 ± 0.41</td>
<td>0.40 (0.25, 0.67)</td>
<td>0.35 ± 0.25</td>
<td>0.30 (0.16, 0.49)</td>
</tr>
<tr>
<td>Banizoumbou</td>
<td>13.54</td>
<td>2.67</td>
<td>250</td>
<td>1999-2012</td>
<td>0.52 ± 0.42</td>
<td>0.40 (0.26, 0.64)</td>
<td>0.35 ± 0.25</td>
<td>0.29 (0.17, 0.48)</td>
</tr>
<tr>
<td>DMN Maine Soroa</td>
<td>13.22</td>
<td>12.02</td>
<td>350</td>
<td>2005-2010</td>
<td>0.48 ± 0.39</td>
<td>0.37 (0.24, 0.62)</td>
<td>0.37 ± 0.30</td>
<td>0.28 (0.15, 0.53)</td>
</tr>
<tr>
<td>Ouagadougou</td>
<td>12.20</td>
<td>-1.40</td>
<td>290</td>
<td>1999-2007</td>
<td>0.52 ± 0.44</td>
<td>0.40 (0.27, 0.62)</td>
<td>0.40 ± 0.24</td>
<td>0.35 (0.21, 0.54)</td>
</tr>
<tr>
<td>Djougou</td>
<td>9.76</td>
<td>1.60</td>
<td>400</td>
<td>2004-2007</td>
<td>0.66 ± 0.44</td>
<td>0.56 (0.38, 0.82)</td>
<td>0.52 ± 0.34</td>
<td>0.41 (0.26, 0.71)</td>
</tr>
<tr>
<td>Ilorin</td>
<td>8.32</td>
<td>4.34</td>
<td>350</td>
<td>1999-2012</td>
<td>0.67 ± 0.49</td>
<td>0.54 (0.32, 0.87)</td>
<td>0.66 ± 0.36</td>
<td>0.59 (0.35, 0.93)</td>
</tr>
<tr>
<td>Southern Africa Sites</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ascension Island</td>
<td>-7.98</td>
<td>-14.42</td>
<td>30</td>
<td>1999-2012</td>
<td>0.16 ± 0.10</td>
<td>0.13 (0.10, 0.20)</td>
<td>0.70 ± 0.37</td>
<td>0.65 (0.42, 0.94)</td>
</tr>
<tr>
<td>Mongu</td>
<td>-15.25</td>
<td>23.15</td>
<td>1107</td>
<td>1999-2009</td>
<td>0.21 ± 0.19</td>
<td>0.14 (0.08, 0.28)</td>
<td>1.60 ± 0.43</td>
<td>1.75 (1.39, 1.89)</td>
</tr>
<tr>
<td>Reunion St. Denis</td>
<td>-20.88</td>
<td>31.59</td>
<td>150</td>
<td>2007-2012</td>
<td>0.064 ± 0.036</td>
<td>0.06 (0.04, 0.08)</td>
<td>0.70 ± 0.36</td>
<td>0.66 (0.42, 0.96)</td>
</tr>
<tr>
<td>Skukuza</td>
<td>-24.99</td>
<td>31.59</td>
<td>150</td>
<td>1999-2011</td>
<td>0.18 ± 0.14</td>
<td>0.14 (0.08, 0.23)</td>
<td>1.34 ± 0.42</td>
<td>1.42 (1.09, 1.64)</td>
</tr>
</tbody>
</table>

Note: AOD$_{550}$ indicates the optical depth at 550 nm, $\alpha_{440-870}$ is the Angstrom exponent between 440 nm and 870 nm.
<table>
<thead>
<tr>
<th>Site</th>
<th>AOD$_{550nm}$ (± σ)</th>
<th>$a_{ext(440/870)}$ (± σ)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Western Africa Sites</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agoufou</td>
<td>0.77 ± 0.41 (Jun)</td>
<td>0.53 ± 0.25 (Dec)</td>
</tr>
<tr>
<td>Dakar</td>
<td>0.62 ± 0.29 (Jun)</td>
<td>0.62 ± 0.30 (Dec)</td>
</tr>
<tr>
<td>Zinder Airport</td>
<td>0.67 ± 0.04 (Aug)</td>
<td>0.51 ± 0.29 (Dec)</td>
</tr>
<tr>
<td>Banizoumbou</td>
<td>0.89 ± 0.57 (Mar)</td>
<td>0.54 ± 0.29 (Dec)</td>
</tr>
<tr>
<td>DMN Maine Soroa</td>
<td>1.01 ± 0.75 (May)</td>
<td>0.62 ± 0.35 (Dec)</td>
</tr>
<tr>
<td>Ouagadougou</td>
<td>0.88 ± 0.70 (Mar)</td>
<td>0.56 ± 0.26 (Dec)</td>
</tr>
<tr>
<td>Djougou</td>
<td>0.97 ± 0.48 (Mar)</td>
<td>0.96 ± 0.30 (Dec)</td>
</tr>
<tr>
<td>Ilorin</td>
<td>1.10 ± 0.56 (Feb)</td>
<td>0.91 ± 0.30 (Dec)</td>
</tr>
<tr>
<td><strong>Southern Africa Sites</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ascension Island</td>
<td>0.32 ± 0.14 (Sep)</td>
<td>1.34 ± 0.17 (Sep)</td>
</tr>
<tr>
<td>Mongu</td>
<td>0.50 ± 0.26 (Sep)</td>
<td>1.85 ± 0.16 (Aug)</td>
</tr>
<tr>
<td>Etoisha Pan</td>
<td>0.40 ± 0.17 (Oct)</td>
<td>1.80 ± 0.16 (Oct)</td>
</tr>
<tr>
<td>Reunion St. Denis</td>
<td>0.095 ± 0.044 (Oct)</td>
<td>1.12 ± 0.28 (Oct)</td>
</tr>
<tr>
<td>Skukuza</td>
<td>0.27 ± 0.18 (Sep)</td>
<td>1.46 ± 0.28 (Sep)</td>
</tr>
</tbody>
</table>

Table 2: Maximum and minimum multi-year monthly averages of AERONET AOD$_{550nm}$ and $a_{ext(440/870)}$ per site. The month of the reported max/min value is indicated in parenthesis. Site names in bold font are used in model comparison.
Table 3. Global and Africa-only annual average burdens, lifetimes, total deposition fluxes, and fraction wet deposition of four prognostic aerosol species in CCAM for the year 2010.

<table>
<thead>
<tr>
<th>Species</th>
<th>Burden (Tg)</th>
<th>Total deposition (Tg a(^{-1}))</th>
<th>Fraction wet deposition of total</th>
<th>Lifetime (days)</th>
<th>Emissions (Tg yr(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Global</td>
<td>Africa</td>
<td>Global</td>
<td>Africa</td>
<td>Global</td>
</tr>
<tr>
<td>BC</td>
<td>0.187</td>
<td>0.0465</td>
<td>6.84</td>
<td>1.56</td>
<td>0.844</td>
</tr>
<tr>
<td>OC</td>
<td>1.11</td>
<td>0.305</td>
<td>44.1</td>
<td>12.1</td>
<td>0.819</td>
</tr>
<tr>
<td>Sulfate</td>
<td>0.961</td>
<td>0.161</td>
<td>65.1</td>
<td>7.18</td>
<td>0.865</td>
</tr>
<tr>
<td>Dust</td>
<td>67.7</td>
<td>26.9</td>
<td>2780</td>
<td>1460</td>
<td>0.565</td>
</tr>
</tbody>
</table>
Table 4. Summary of model-observation comparison of monthly-average AOD$_{550nm}$.
The significance of the Pearson’s correlation is indicated by ‘*’ for $p<0.05$, ‘**’ for $p<0.01$, and ‘***’ for $p<0.001$; NS is not significant at 0.05 level.

<table>
<thead>
<tr>
<th>Site</th>
<th>Correlation coefficient (r)</th>
<th>Normalized Mean Bias*</th>
<th>Mean Absolute Error*</th>
<th>Number of months</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Northern African and Middle Eastern Sites</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Granada</td>
<td>0.47 ***</td>
<td>176.6 %</td>
<td>0.27</td>
<td>50</td>
</tr>
<tr>
<td>Blida</td>
<td>0.70 ***</td>
<td>220.0 %</td>
<td>0.54</td>
<td>33</td>
</tr>
<tr>
<td>Lampedusa</td>
<td>0.58 ***</td>
<td>278.2 %</td>
<td>0.51</td>
<td>46</td>
</tr>
<tr>
<td>Saada</td>
<td>0.60 **</td>
<td>231.7 %</td>
<td>0.50</td>
<td>74</td>
</tr>
<tr>
<td>Sede_Boker</td>
<td>0.23 *</td>
<td>245.5 %</td>
<td>0.43</td>
<td>129</td>
</tr>
<tr>
<td>Santa Cruz Tenerife</td>
<td>0.44 ***</td>
<td>339.1 %</td>
<td>0.54</td>
<td>60</td>
</tr>
<tr>
<td>Dhadnah</td>
<td>0.81 ***</td>
<td>125.1 %</td>
<td>0.45</td>
<td>50</td>
</tr>
<tr>
<td>Solar Village</td>
<td>0.51 ***</td>
<td>121.1 %</td>
<td>0.42</td>
<td>128</td>
</tr>
<tr>
<td>Dahkla</td>
<td>0.49 *</td>
<td>242.2 %</td>
<td>0.75</td>
<td>19</td>
</tr>
<tr>
<td>Hamim</td>
<td>0.82 ***</td>
<td>115.2 %</td>
<td>0.37</td>
<td>28</td>
</tr>
<tr>
<td>Tamanasset_INM</td>
<td>0.89 **</td>
<td>253.6 %</td>
<td>0.51</td>
<td>19</td>
</tr>
<tr>
<td><strong>Western Africa Sites</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agoufou</td>
<td>0.51 **</td>
<td>89.7 %</td>
<td>0.47</td>
<td>58</td>
</tr>
<tr>
<td>Dakar</td>
<td>0.33 **</td>
<td>103.2 %</td>
<td>0.48</td>
<td>95</td>
</tr>
<tr>
<td>Zinder_Airport</td>
<td>0.61 **</td>
<td>59.3 %</td>
<td>0.35</td>
<td>30</td>
</tr>
<tr>
<td>Banizoumbou</td>
<td>0.50 **</td>
<td>58.6 %</td>
<td>0.34</td>
<td>126</td>
</tr>
<tr>
<td>DMN_Maine_Soroa</td>
<td>0.52 **</td>
<td>94.5 %</td>
<td>0.46</td>
<td>41</td>
</tr>
<tr>
<td>Ouagadougu</td>
<td>0.27 *</td>
<td>29.3 %</td>
<td>0.28</td>
<td>61</td>
</tr>
<tr>
<td>Djougou</td>
<td>0.29 NS</td>
<td>-1.3 %</td>
<td>0.20</td>
<td>24</td>
</tr>
<tr>
<td>Ilorin</td>
<td>0.59 **</td>
<td>-12.6 %</td>
<td>0.22</td>
<td>61</td>
</tr>
<tr>
<td><strong>Southern Africa Sites</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ascension_Island</td>
<td>0.51 **</td>
<td>41.8 %</td>
<td>0.09</td>
<td>53</td>
</tr>
<tr>
<td>Mongu</td>
<td>0.67 **</td>
<td>-21.2 %</td>
<td>0.09</td>
<td>77</td>
</tr>
<tr>
<td>Reunion – St Dénis</td>
<td>0.21 NS</td>
<td>135.0 %</td>
<td>0.09</td>
<td>84</td>
</tr>
<tr>
<td>Skukuza</td>
<td>0.48 **</td>
<td>24.6 %</td>
<td>0.07</td>
<td>72</td>
</tr>
</tbody>
</table>

* $NMB = \frac{1}{N} \sum_{i=1}^{N} \frac{M_i - O_i}{O_i} \times 100\%$; $MAE = \frac{1}{N} \sum_{i=1}^{N} |M_i - O_i|$ (N is number of points, $M$ are modeled values and $O$ are observed values)
Figure 1: Map of long-term AERONET sites used in this study. Sites are color-coded by general geographic area and aerosol source type. Site names in bold italics are used in the model comparison.
Figure 2: Comparison of methods to compute mean modeled AOD$_{550\text{nm}}$ for an example in January 2000: red bars include model output only for 6am to 12pm UTC; yellow bars for 6am to 6pm UTC; and blue bars for 24 hours. Whiskers are ±1 standard deviation across the 6-hourly model values within each time range.
Figure 3a: Multi-year mean seasonal cycle of observed AERONET AOD$_{550}$ at long-term sites in Northern Africa and the Middle East. The number of years of data used for each month is shown at the top of the plot area, and the total range of years of observations used is listed under each site name. Whiskers are ±1 standard deviation across daily means within a given month.
Figure 3b: Multi-year mean seasonal cycle of observed AERONET AOD$_{550nm}$ at long-term sites in Western and Southern Africa. The number of years of data used for each month is shown at the top of the plot area, and the total range of years of observations used is listed under each site name. Whiskers are ±1 standard deviation across daily means within a given month.
Figure 4a: Same as Fig. 3a but for observed $\alpha_{\text{ext}}(440/870)$ from AERONET. Gray shaded region represents $\alpha_{\text{ext}}(440/870)$ values typical of aerosols dominated by coarse particles (Holben et al., 2001; Ogunjobi et al., 2008). Whiskers are ±1 standard deviation across daily means within a given month.
Figure 4b: Same as Fig. 3b but for observed $\alpha_{\text{ext}(440/870)}$ from AERONET. Gray shaded region represents $\alpha_{\text{ext}(440/870)}$ values typical of aerosols dominated by coarse particles (Holben et al., 2001; Ogunjobi et al., 2008). Whiskers are ±1 standard deviation across daily means within a given month.
Figure 5: Comparison of present-day model results for CCAM (blue triangles) against ranges from other models (shaded gray area), for a) global burdens of major aerosol constituents, b) characteristics of OA aerosol, c) characteristics of BC aerosol, and d) characteristic of dust aerosol. Reference model ranges in a) are from Kinne et al. (2006) with additional models provided from Jathar et al. (2011) for OC, Liu et al. (2005) for sulfate, and Zender et al. (2004) for dust. AeroCom Phase II model ranges and medians (black crosses) in b) are from Tsigaridis et al. (2014); CCAM modeled OC is converted to OA by multiplying by a factor of 1.4 for a consistent comparison (Tsigaridis et al., 2014). CMIP5 model ranges and medians (black circles) in c) are from Allen and Landuyt (2014). AeroCom Phase I model ranges and medians (black crosses) in d) are from Huneus et al. (2011).
Figure 6: Multi-year mean seasonal cycle of AOD$_{550nm}$ for observed (red) and modeled with all CCAM outputs, 1999–2012 (blue), and only those months with AERONET data meeting the 70% completeness cutoff (yellow). ± 1 standard deviation for the observations and CCAM 1999–2012 output across daily means within a given month is shaded.
Figure 6, continued: Multi-year mean seasonal cycle of AOD$_{550nm}$ for observed (red) and modeled with all CCAM outputs, 1999–2012 (blue), and only those months with AERONET data meeting the 70% completeness cutoff (yellow). ± 1 standard deviation for the observations and CCAM 1999–2012 output across daily means within a given month is shaded.
Figure 7: Multi-year mean observed vs. modeled seasonal cycle of AOD\textsubscript{550nm} at six AERONET sites. Modeled AOD\textsubscript{550nm} is broken down into the contribution from each aerosol species (sea salt, large size bin dust (radius $\geq$ 1 $\mu$m), small size bin dust (radius < 1 $\mu$m), BC, OC, sulfate (SO\textsubscript{4})).
Figure 8. Comparison of the probability densities of modeled (red) and observed (black) daily average AOD$_{550\text{nm}}$ at each site.
Figure 9: Seasonal variation in multi-year monthly 5th percentile, mean, and 95th percentile modeled AOD$_{550nm}$ (map background) for the full model climatology (1999–2012), with observed multi-year means (points) for all available AERONET data, for a) September and b) March. The number of years and range of years used for each site is the same as in Fig. 3.