Radiative Heating Rate Profiles over the Southeast Atlantic Ocean during the 2016 and 2017 Biomass Burning Seasons

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Abstract. Marine boundary layer clouds, including the transition from stratocumulus to cumulus, are poorly represented in numerical weather prediction and general circulation models. Further uncertainties in the cloud structure arise in the presence of biomass burning carbonaceous aerosol, as is the case over the southeast Atlantic Ocean where biomass burning aerosol is transported from the African continent. As the aerosol plume progresses across the southeast Atlantic Ocean, radiative heating within the aerosol layer has the potential to alter the thermodynamic environment and therefore the cloud structure; however, this has yet to be quantified. The deployment of the First Atmospheric Radiation Measurement Mobile Facility (AMF1) in support of the Layered Atlantic Smoke Interactions with Clouds (LASIC) field campaign provided a unique opportunity to collect observations of cloud and aerosol properties during two consecutive biomass burning seasons during July through October of 2016 and 2017 over Ascension Island (7.96 S, 14.35 W). Using observed profiles of temperature, humidity, and clouds from the LASIC field campaign, alongside aerosol optical properties from the Modern Era Retrospective analysis for Research and Applications, version 2 (MERRA-2) as input for the Rapid Radiation Transfer Model (RRTM), profiles of the radiative heating rate due to aerosols and clouds were computed. Radiative heating is also assessed across the southeast Atlantic Ocean using an ensemble of back trajectories from the Hybrid Single Particle Lagrangian Integrated Trajectory Model (HYSPLIT). Idealized experiments using RRTM with and without aerosols and a range of values for the single scattering albedo demonstrate that shortwave (SW) heating within the aerosol layer above Ascension Island can locally range between 2 and 8 K per day, though impacts of the aerosol can be felt elsewhere in the atmospheric column. SW radiative heating due to biomass burning aerosol is not balanced by additional longwave cooling, and the net radiative impact results in a stabilization of the lower troposphere. However, these results are extremely sensitive to the single scatter albedo and the height of the aerosol plume with respect to the inversion layer.

1 Introduction

Marine stratocumulus and trade wind cumulus are prominent cloud types over the Atlantic Ocean with regional and global impacts on the energy budget (Bony and Dufresne, 2005). Despite their importance, models struggle to accurately...
represent these clouds and their properties. Within the southeast Atlantic and other subsidence regions, general circulation models and reanalyses tend to underestimate the cloud fraction (Klein et al., 2013; Dolinar et al., 2015) and optical thickness of warm marine stratocumulus due to underestimates in cloud albedo and liquid water path (Lin et al., 2014; Noda and Satoh, 2014; Rapp. 2015). Furthermore, models struggle to properly link environmental conditions to cloud properties of trade wind cumuli (Nuijens et al., 2015). The uncertainty and discrepancy among models within the region is further complicated by the presence of biomass burning aerosol (Stier et al., 2013; Peers et al., 2016). Using global model simulations, it was shown by Brown et al. (2018) that the largest radiative impact from brown carbon occurs off the west coast of southern Africa. Biomass burning aerosol that gets entrained into marine stratocumulus in the southeast Atlantic has a larger impact on the radiation budget than the direct radiative effect of the aerosol itself (Lu et al., 2018). The determination to answer questions and resolve uncertainties surrounding this topic in the southeast Atlantic Ocean led to an international effort termed COLOCATE (Clarify-Oracles-Lasic-aerOClo-seAls Team Experiment), with overlapping field campaigns and modeling studies from the United Kingdom, France, South Africa, Namibia, and the United States (Zuidema et al., 2016). The focus here is a combination of radiation transfer modeling and observations from DOE’s Layered Atlantic Smoke Interactions with Clouds (LASIC) campaign.

Originating in the savannas of southwestern Africa, biomass burning aerosol is lofted to between 3.5 to 4.5 km and transported via the Southern Africa Easterly Jet over the southeast Atlantic Ocean where the aerosol plume begins to descend (Adebiyi et al., 2015; Adebiyi and Zuidema, 2016; Das et al., 2017). Fires and the associated aerosol in this region are typical during the months of July through October. When compared to satellite observations, models commonly allow for the biomass burning aerosol to descend too rapidly once over the ocean (Das et al., 2017; Gordon et al., 2018), which can have implications on the thermodynamic structure and can indicate dynamical deficiencies. While over the ocean, observations indicate that the aerosol plume is primarily above the boundary layer. Over Ascension Island, the aerosol tends to be in the boundary layer during the beginning of the biomass burning season but is located above the cloud layer towards the end in September and October (Zuidema et al., 2018b).

Biomass burning aerosol in the Southeast Atlantic region and its impact on heating within the column has been investigated through recent modeling experiments (Chang and Christopher, 2017; Lu et al., 2017, Gordon et al., 2018; Mallet et al., 2019). Heating rate profiles within the region were calculated by Chang and Christopher (2017) using the Santa Barbara DISORT Atmospheric Radiative Transfer (SBDART) model and fixed values for aerosol and cloud properties corresponding to Southern AFRicAn Regional science Initiative (SAFARI 2000) observations. Chang and Christopher (2017) noted that with fixed aerosol and cloud properties, the radiative heating rate increased throughout the biomass burning season due to the decreasing solar zenith angle. This study also determined the solar zenith angle (54°) at which the direct radiative effect of aerosols located above liquid clouds is maximized. Lu et al. (2017) used large eddy simulations nested within Weather Research and Forecasting with Chemistry (WRF-Chem) to quantify the microphysical, direct, and semi-
direct effects of aerosol within the Southeast Atlantic. A total cooling of roughly 8 W m$^{-2}$ in the shortwave (SW) was found at the top of the atmosphere with a large component of that from the microphysical effects of biomass burning aerosols on clouds as a result of the Twomey effect, higher liquid water path, and higher cloud fraction before noon (Lu et al., 2017). Another recent study by Gordon et al. (2018) quantified radiative heating within the atmospheric column by switching biomass burning aerosols and absorption due to biomass burning aerosols on and off in a hybrid of the regional configuration of the UK Chemistry and Aerosol Model and HadGEM. While Gordon et al. (2018) established the use of the hybrid model combination for aerosol studies and identified discrepancies between the model and observations, only the first ten days of August 2016 were analyzed.

The primary goal of this study is to quantify the individual impact of clouds, aerosols, and black carbon on heating within the atmospheric column above Ascension Island in the Southeast Atlantic, where the First Atmospheric Radiation Measurement (ARM; Mather and Voyles, 2013) Mobile Facility (AMF1; Miller et al., 2016) was deployed during the biomass burning seasons of 2016 and 2017. Previous studies of the radiative heating rate within the column in the Southeast Atlantic are expanded upon by employing varying thermodynamic, cloud, and aerosol properties using observations from the AMF1 and observation-constrained aerosol profiles from the Modern-Era Retrospective analysis for Research and Applications, Version 2 (MERRA-2; Gelaro et al., 2017; GMAO, 2015a; GMAO, 2015b) throughout the biomass burning seasons of 2016 and 2017 over Ascension Island. Our approach uses observations and analyses of aerosol and cloud properties as input to a radiative transfer model. Aerosol impacts on cloud properties resulting in changes in the cloud radiative properties, i.e. aerosol indirect effects, will be captured through the observed cloud properties. Furthermore, heating rates are explored along a back trajectory originating at Ascension Island. Section 2 describes the observational and reanalysis data sets that are used in this study as well as the methodology for idealized radiation transfer simulations. Results beginning with an evaluation of aerosol optical depth (AOD) and ending with radiative heating rates due to atmospheric constituents are detailed in Section 3, while a discussion and conclusions can be found in Section 4.

2 Data and Methodology

2.1 ARM Mobile Facility and Value-Added Products

Observations of thermodynamic profiles, clouds, and aerosols used in this study are from the AMF1, which was located on Ascension Island (7.7 S, 14.35 W, 340.77 m) from 1 June 2016 through 31 October 2017 with the objective of observing two consecutive biomass burning seasons. While the AMF1 was stationed on the windward side of the island, radiosondes were launched at the airport on the southeastern side of the island near an existing Aerosol Robotic Network (AERONET) site (Zuidema et al., 2018a). The interpolated sounding (INTERPSONDE) value-added product (VAP) is used for temperature and humidity profiles (ARM Climate Research Facility, 2016a; ARM Climate Research Facility 2016b). INTERPSONDE is anchored by six hourly radiosonde launches and a linear interpolation is used to fill in time steps
between launches (Toto and Jensen, 2016). Microwave radiometer retrievals (MWRRET; Gaustad et al. 2011) of precipitable water vapor are used to further constrain the humidity profiles. The resulting INTERPSONDE data has a temporal resolution of one minute and vertical resolution ranging from 20 m to 500 m, depending on the height above ground level. Aerosol optical depth (AOD) was observed using a multifilter rotating shadowband radiometer (MFRSR) and calculated using the 1st Michalsky algorithm (Koontz et al., 2013). Additional measurements of AOD from AERONET were taken using a Cimen sun photometer (Holben et al., 2001; Giles et al., 2019).

Cloud properties used in the radiation transfer simulations were determined using a Ka-band cloud radar, micropulse lidar, and laser ceilometer, with the data combined into the Active Remote Sensing of Clouds (ARSCL) VAP (Clothiaux et al., 2000). Cloud properties including cloud liquid/ice water content, ice and liquid/ice cloud droplet effective were determined using the method presented in Dunn et al. (2011) and currently used in MICROBASE, which is a retrieval algorithm utilizing constrained data from ARSCL as well as the microwave radiometer and INTERPSONDE profiles. The accuracy of this retrieval algorithm has been evaluated using radiative closure experiments and it is known to be accurate enough to adequately represent radiation transfer through clouds. It has been used in past studies (e.g. Mather et al., 2007) to estimate tropical heating rate profiles. Complete validation of such an algorithm is not possible using in-situ measurements, but its reliance upon cloud liquid water path and its use in the tropical atmosphere are consistent with its capability. All clouds that are colder than -16°C were considered to be comprised entirely of ice, while all clouds above 0°C were liquid. A linear fractionation scheme was used to partition particle phase in the region between 0 and 16°C. It is worth noting however that clouds over Ascension Island are primarily liquid. Thorough comparisons to other retrieval algorithms, and evaluations of the relative performance of the MICROBASE algorithm are presented by Zhao et al. (2012) and Huang et al. (2012).

2.2 MERRA-2

The vertical profile of aerosols and their integrated properties can be difficult to continuously observe, especially during cloudy conditions. Throughout the LASIC campaign, there were numerous hours without observations of aerosol optical depth. In order to maximize time steps when heating rate profiles could be calculated given the near constant partly cloudy skies over Ascension Island, aerosol properties from the Modern-Era Retrospective analysis for Research and Applications, Version 2 (MERRA-2; Gelaro et al., 2017; GMAO, 2015a; GMAO, 2015b) were instead used for the radiation transfer simulations. MERRA-2 is the latest contemporary reanalysis from NASA that has the advantage of assimilated aerosol optical depth (AOD), a feature that is not present in other reanalysis products. The decision to use MERRA-2 was made such that we would have a self-consistent data source of aerosols, clouds, and thermodynamic profiles to use for heating rate profiles not just over Ascension Island, but along the entire trajectory of biomass burning aerosol transport originating from southern Africa. MERRA-2 data is available at a spatial resolution of roughly 50 km and 72 vertical levels from the surface through 0.1 hPa and a temporal resolution of one hour for single level variables and three hours for three dimensional variables.
The dominant observational source of AOD that is assimilated into MERRA-2 is Collection 5 bias-corrected Moderate Resolution Imaging Spectroradiometer (MODIS) AOD (Randles et al., 2017). Other aerosol datasets are assimilated into MERRA-2 however they are not applicable for the time period of the LASIC campaign. Daily emissions of biomass burning aerosol come from the Quick Fire Emissions Dataset (QFED) version 2.4-r6 (Darmenov and da Silva, 2015). Within MERRA-2, aerosols are simulated using the Goddard Chemistry, Aerosol, Radiation, and Transport Model (GOCART), which separates the AOD into five species, sea salt, dust, sulfate, organic carbon, and black carbon, and defines the vertical distribution of aerosols. Further details on the assimilation of aerosols in MERRA-2 can be found in Randles et al. (2017), while an evaluation with respect to independent observations can be found in Buchard et al. (2017). MERRA-2 aerosols during the LASIC campaign are further evaluated in Section 3.1.

2.3 Rapid Radiative Transfer Model

The Rapid Radiative Transfer Model (RRTM) was used to perform idealized experiments to calculate the SW heating within the column due to black carbon, all aerosols, and clouds. A user-specified vertical profile was used with the temperature and humidity profiles coming from INTERPSONDE and cloud properties from MICROBASE. Prior to insertion into RRTM, the INTERPSONDE profiles were interpolated onto the MERRA-2 vertical profile based on height above ground level. RRTM runs were performed every four seconds to match the temporal resolution of MICROBASE, while solar zenith angle was updated every fifteen minutes, the temperature and humidity profiles hourly, and aerosols every three hours due to the temporal resolution of vertical profiles in MERRA-2. Aerosol optical properties, including AOD, angstrom exponent, and single scatter albedo (SSA) are from MERRA-2, and were scaled in the vertical by the profile of mixing ratio for the individual species (GMAO, 2015; GMAO, 2015b). The value for SSA at 550 nm from MERRA-2 was used and assumed to be spectrally independent. Asymmetry parameter was assumed to be 0.756, the value given by Hess et al. (1998) for a polluted maritime air mass. Other values of asymmetry parameter were tested but did not impact the results. A total of six sets of experiments were completed to quantify the individual and combined contribution of clouds and aerosols: 1) Clean and clear sky without clouds or aerosols, 2) Clear sky with all aerosols, 3) Clear sky with all aerosols except black carbon, 4) Clean and cloudy sky, 5) Cloudy sky with all aerosols, and 6) Cloudy sky with all aerosols except black carbon. It is worth noting that a true assessment of heating due to biomass burning aerosol should isolate brown carbon, however that is not an aerosol species available in MERRA-2 at this time.

3 Results

3.1 Evaluation of Aerosols in MERRA-2

Previous evaluations of aerosol properties in MERRA-2 have been limited so it is therefore essential to ensure that MERRA-2 is representative of the observations that are available from the AMF1 when it was stationed in Ascension Island. Aside from observations from the AMF1, there are also AOD observations from an existing AERONET site located near the
airport where the soundings were launched (Holben et al., 2001). Daily mean AOD from the two observational sources as well as MERRA-2 for August and September 2016 and 2017 can be seen in Figure 1. It can readily be seen that observations from the AMF1 are limited in all four months. Therefore, correlation coefficients and biases presented in Figure 1 were calculated for MERRA-2 with respect to AERONET observations only including days when observations were available.

When it was cloudy, AERONET was not able to measure AOD. The highest aerosol loading over Ascension Island was present in the middle and end of August 2016, with daily values of AOD ranging from 0.1 the first couple days of the month to a maximum of 0.73 on 13 August 2016, followed by additional periods of elevated AOD during September 2017. These values for AOD are similar to those presented by Zuidema et al. (2016) using AERONET observations over the period of 2000 through 2013. A periodicity can be seen in each of the four months as the aerosol plume drifts overhead of Ascension Island. Correlations between AOD in the observations and MERRA-2 exceed 0.8 in all four months. The largest bias of 0.04 with respect to AERONET occurs in August 2017 however the AERONET observations are also generally higher than those from the AMF1.

Observations of SSA during LASIC were presented by Zuidema et al. (2018b) and monthly mean values of 0.78 and 0.81 were specified during August and September at a wavelength of 529 nm. When all of the aerosol species are considered in MERRA-2, the SSA tends to be a bit higher, with monthly mean values of 0.92 and 0.93 for August and September respectively. There are a few possible explanations for this discrepancy. In reality, much of the organic biomass burning aerosol can be considered brown carbon, a species that is not represented in GOCART and the Goddard Earth Observing System (GEOS), the underlying model and data assimilation system in MERRA-2. Brown carbon tends to be more absorbing than organic carbon and therefore if included, the SSA could be lower. In addition, the optical properties for aerosols in MERRA-2 are defined by a look up table as a function of relative humidity. Differences in the thermodynamic profile will therefore result in a different SSA. On the contrary, the SSA in MERRA-2 is more aligned with those presented for the region by Pistone et al. (2019) from the ORACLES and previous field campaigns. It is also possible that the observations presented by Zuidema et al. (2018b) are representative of volcanic dust from the island itself, in addition to the plume of biomass burning aerosol, and therefore not the same as the aerosol properties in the MERRA-2 50 km grid box.

The impact of the discrepancy in SSA on the heating rate profile due to aerosols will be further discussed in Section 3.3.

Only AOD is assimilated in MERRA-2 and therefore GOCART is used to distribute the aerosol within the atmospheric column. The vertical profile of the mixing ratio of black and organic carbon in MERRA-2 is shown in Figure 2, alongside contours of cloud fraction with a value of 0.25. From an initial glance, it can be seen that larger values for the mixing ratio of black and organic carbon correspond to the dates with elevated AOD in Figure 1. The majority of the aerosol loading is located between 850 and 650 hPa, which corresponds to roughly 1500 to 3750 km in height in MERRA-2. In agreement with Zuidema et al. (2018), the black and organic carbon in MERRA-2 is located above the cloud layer, but perhaps extends higher in the atmosphere than indicated by lidar observations. Qualitatively, MERRA-2 is also able to capture the thinning of the vertical extent of the aerosol as the loading decreases following the maximum in the middle of August. Also evident is a decrease in the height of the boundary layer as the season progresses.
The AOD at Ascension Island is a function of both the large-scale transport and also the timing and location of fires in Southern Africa. To determine the origin of the aerosol and how it differs between the 2016 and 2017 biomass burning seasons, the HYbrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model was used to compute ten day back trajectories for a parcel originating at Ascension Island at 12z on each day in August and September 2016 and 2017, driven by the large scale meteorology from MERRA-2 (Stein et al., 2015). Based on prior results for the height of the aerosol plume, the parcel originated at a height of 2 km. Some similarities can be seen between the back trajectories and the magnitude of the AOD at Ascension Island (Figure 3). The highest values of AOD were observed during August 2016 and September 2017. Both of these months have back trajectories that extend well into the African continent (Figures 3a and d), which is hardly the case for August 2017 when the subtropical highs over the southern Indian and Atlantic Oceans were shifted further to the east compared to 2016 (Figure 3c). Excluding the day with the highest AOD in August 2016, days with an elevated AOD had a back trajectory that originated from the south of Ascension Island, crossing the land-ocean boundary of the African coast between 10 and 15 S (Figure 3a). On the contrary, days in August 2016 that observed an AOD below 0.3 tended to have back trajectories that originate further north. The variance in daily AOD was not as large in September 2017, with most of the back trajectories having a more easterly path.

Given that the observed aerosol loading over Ascension Island is highest during August 2016, we have elected to only focus on that month. The same analysis has been completed for August 2017 as well as September 2016 and 2017 and monthly mean maximum SW heating rates within the atmospheric column due to clouds and aerosols for all months are presented in Table 1.

3.2 Thermodynamic Profiles over Ascension Island

A key characteristic of the atmosphere over Ascension Island is an inversion-topped marine boundary layer (MBL) as seen in the temperature time-height section (top panel of Figure 4) as a warmer (yellow) region bounded above and below by cooler (more orange) regions at ~700 hPa. Beneath the thermal inversion relative humidity is generally much higher and more hospitable for cloud development as seen is Figure 5a. A thorough treatment of the thermodynamic structure during the biomass burning seasons of 2016 and 2017 can be found in Zhang and Zuidema (2019).

MBLs of this depth typically accommodate transition cloud structure, which is characterized by single layer stratocumulus when the MBL is relatively shallow and trade cumulus when it is deeper. Intermediate stages in this deepening-warming MBL structure are characterized by hybrid cloud configurations consisting of a mix of layered stratocumulus and cumulus that intermingle in complex ways. Deeper MBL’s tend to contain two or more internal boundary layers that are separated by a weak inversion, a process known as decoupling, which leads to the development of cumulus convection that rejoins the two-layers leading to a “cumulus-coupled” MBL. Manifestations of decoupling are best observed in the bottom panel of Figure 4, which exhibits a subtle, intermittent sub-layer at ~900 hPa, and in Figure 13 of Zhang and Zuidema (2019). Above the MBL where most of the biomass-burning aerosol is located, there are intermittent bursts of moist air, potentially a result of weak easterly waves. Occasionally these waves may be accompanied by mid-level cloud
cover, for example at ~600 hPa around August 25, 2016, but these clouds are too thin and contain small enough droplets that they are not detectable using a cloud radar (see Figure 5a).

### 3.3 Heating Rate Profiles over Ascension Island

Idealized radiative transfer calculations were used to quantify the heating rates of aerosols and clouds within the atmospheric column over Ascension Island. Given the discrepancy in SSA between MERRA-2 and the observations, a sensitivity test was performed to determine the role of SSA on heating due to aerosols within the column to quantify the uncertainty due to the SSA used. Three different values of the SSA were used to represent the original SSA in MERRA-2, and potential deficiencies related to the vertical profile in relative humidity and the lack of brown carbon. In order to adjust for relative humidity, the SSA was determined by the lookup table used in MERRA-2 for the scattering and extinction properties of black and organic carbon at 550 nm as a function of the observed relative humidity. Adjusting for the humidity alone does not fully explain the difference in SSA between MERRA-2 and the observations. To account for the lack of brown carbon, the SSA for organic carbon was multiplied by 0.85, which is the mean percent difference between MERRA-2 and the observations presented by Zuidema et al. (2018b). A summary of the monthly mean maximum SW radiative heating and where it occurred within the column for the entire 2016 and 2017 biomass burning seasons can be seen in Table 1 however the figures with more detailed information are only shown for August 2016 as that was the month with the highest aerosol loading.

Results for the SW aerosol radiative effect using these three sets of values for SSA under clear-sky conditions can be seen in the left column of Figure 6. Within the atmospheric column, the majority of the heating due to aerosols occurs in the layer around 800 hPa, though the impact of aerosols can be felt to a lesser extent down to the surface regardless of the SSA (Figure 6a, b, and c). There is minimal heating due to aerosols the first few days of August 2016 as the AOD is only around 0.1. Throughout the rest of the month the radiative heating rate profile follows the periodicity of aerosol loading as seen in Figure 2. Aerosols are spread within a deeper layer beginning 25 August 2016 and, as such, the heating within the column occurs in a thicker layer than earlier in the month. Although the highest AOD occurs on 13 August 2016, the maximum heating rate on that day is just shy of the largest heating rate of ~2.7 K day\(^{-1}\) (~6.25 K day\(^{-1}\) when the SSA for black and organic carbon is reduced) within the month that occurs on 30 August 2016 and 31 August 2016. A likely explanation for this is that there is more swelling of the aerosol aloft the last two days in August as a result of increased humidity.

As expected, heating rates are smaller when the original SSA from MERRA-2 is used. Though somewhat difficult to see with the color bar, SW heating rates are slightly larger in magnitude when the SSA is scaled based on the observed relative humidity (Figure 6 a and c). SW heating rates can actually double or triple if the SSA for organic carbon is reduced to simulate the role of brown carbon and to be more in line with the observed SSA (Figure 6b). This finding furthers the importance of an accurate representation of aerosol optical properties in models within the Southeast Atlantic already stressed in the literature (Mallet et al., 2019; Pistone et al., 2019).
Unlike in other regions, heating due to clouds, generally located below 900 hPa, is underwhelming and of similar order of magnitude as the heating due to aerosol with the original MERRA-2 SSA (Figure 6d). There is less day-to-day variability in the magnitude of SW heating within the cloud layer compared to heating from aerosols. This is somewhat expected due to the consistent nature of the cloud water path and effective radius. There is however some variability in the location of the heating in connection with fluctuations in the height of the boundary layer. By comparing the results for cloudy to clear conditions, it can be seen that in the presence of clouds, radiative heating within the aerosol layer is embellished (Figure 6d, e, and f).

For simplicity, from this point forward heating rates due to clouds and aerosols are discussed using the relative humidity scaled SSA for organic and black carbon. To isolate the absorption due to black carbon itself, the heating rates from the simulations without black carbon are subtracted from those that incorporate all aerosol species (Figure 7). Note that the color bar in Figure 7 has been reduced compared to Figure 6 in order to emphasize the day-to-day variability in SW heating. Most of the SW heating within the atmospheric column due to aerosols is indeed a result of black carbon. On days when the aerosol plume is overhead, the daytime local heating as a result of black carbon can exceed 2 K per day under clear skies (Figure 7a). Considering the monthly mean maximum heating within the clear sky column is roughly 2.5 K per day (Table 1), this is not insignificant, but only accounts for a portion of the SW heating due to all aerosols. The remaining SW heating within the aerosol plume and down to the surface is likely due to the extinction of radiation from organic carbon that is not scattered within the plume. Above the primary aerosol plume and extending upward in the column to around 600 hPa, heating due to aerosol on the order of ~0.5 K per day as seen in Figure 6 is not associated with black carbon, but rather is absorption from dust. As was the case with all aerosols, heating due to black carbon is enhanced when clouds are present and can approach 3 K per day (Figure 7b). Our results are quite similar to those presented by Gordon et al. (2018), who showed a mean SW heating of 1.9 K day\(^{-1}\) due to biomass burning aerosol for the period of 6 August 2016 through 10 August 2016 over the Southeast Atlantic. For the same five day period, we see a mean daytime SW heating due to black carbon of 1.86 K day\(^{-1}\) within the layer between 760 and 840 hPa. Gordon et al. (2018) took a similar approach by turning aerosols and black carbon off in a model simulation but this was done using global and regional simulations with HadGEM. However, the authors stated their results might not be representative of the heating that actually occurred as the aerosols in their simulations were too low in altitude.

There is an interplay between clouds and aerosols when they are considered together as opposed to individually. Photons scattered by clouds reenter the aerosol layer and have an additional opportunity to be absorbed within the atmosphere as opposed to reaching the surface. The enhancement of heating within the aerosol layer due to clouds is displayed in Figure 8 and is on the order of tenths of a K per day. On most days with sufficient aerosol loading the enhancement is a few K per day but when all aerosols are considered the majority of the enhancement is located within the aerosol layer (Figure 8a). Additional heating due to aerosols in the presence of clouds occurs below the aerosol layer and down to the surface, however this is limited to the morning and evening hours when the sun angle is low and not present when only black carbon is considered (Figure 8b). Within the aerosol layer itself, between 900 and 700 hPa, black carbon is
mostly responsible for the additional heating. The amount of enhancement in SW heating within the aerosol layer due to clouds is variable depending on the location and thickness of the cloud as well the AOD. The greater the AOD and cloud water path, the greater the interaction between the two. While the largest local heating rate under clear skies occurs on 30 and 31 August 2016, when cloudy, the largest local heating rate occurs on 13 August 2016. The aerosol heating rate is further enhanced due to clouds on 13 August when not only the AOD is higher, but the cloud water content is also higher compared to the end of the month.

Aerosols tend to have a minimal direct impact in the longwave (LW) part of the spectrum but they can indirectly impact the LW radiation within the atmospheric column. Heating within the atmospheric column can be lost due to LW radiational cooling. At the present time, aerosols are not a direct input for RRTM LW. In an effort to quantify the LW radiational cooling associated with SW aerosol heating, the hourly mean heating rates, as shown in Figure 6c, were used to modify the observed temperature profile and this was used as the input temperature profile for RRTM LW; the humidity profile was not adjusted. The temperature profile was adjusted each hour, however, any SW heating that was not lost due to additional radiational cooling from the increased temperature was allowed to persist through the following hour. This methodology is somewhat extreme as heating due to aerosol can be transferred to other forms of energy such as latent heat and transported through advection. However, it can be used to determine whether the SW heating due to aerosols is offset by increased radiational cooling.

Results from this exercise are displayed in Figure 9a. The LW heating rate due to aerosols was determined by subtracting a control run using the observed temperature profile from the runs with the modified temperature profile due to SW aerosol heating. Radiational cooling occurs throughout the aerosol layer and is maximized at the bottom of the layer, where at times it can locally reach near 5 K/day. Heat is transferred above and below the aerosol layer when the radiational cooling occurs, with a larger magnitude of the heat being displaced toward the surface. It is evident that without an atmospheric circulation or other processes occurring in the atmosphere, additional heat due to aerosol absorption remains in the column. This is demonstrated by the fact that radiational cooling still occurs through mid to late August despite a suppressed aerosol loading (Figure 9a). During the daytime hours, LW radiational cooling never offsets the absorption due to aerosols (Figure 9b). Even at night, the magnitude of the LW cooling never reaches the magnitude of the daytime SW heating. There is, however, a redistribution of heat as a result of aerosols. The largest magnitude of warming due to aerosol occurs during the daytime hours in the middle of the aerosol layer, and this daytime heating extends vertically in both directions. At night, cooling is maximized at the bottom of the aerosol layer, though is present to some extent within the entire aerosol layer, and some heating occurs above and below the aerosol. The thermodynamic structure of the atmospheric column is therefore altered on a diurnal cycle when aerosols are present and this can have implications for other atmospheric processes such as the development, maintenance, and transition of marine stratocumulus and trade cumulus clouds (Zhang and Zuidema, 2019).

While it is informative to investigate the heating rate profile due to biomass burning aerosol above Ascension Island, it is imperative that such an analysis is also be completed along the trajectory of the aerosol plume as it makes its way
from southern Africa and over the Atlantic Ocean. A case study has been completed for the seven day HYSPLIT back trajectory originating at 2 km above Ascension Island at 13z on 13 August 2016. This date was chosen as it had the highest observed and MERRA-2 analyzed AOD among the 2016 and 2017 biomass burning seasons. As indicated by the spread of the trajectories in panels a-d of Figure 5 from Zuidema et al. (2018b) and Figure 10, there is some uncertainty regarding the exact path of the biomass burning aerosol plume. In order to account for this, HYSPLIT was forced by the meteorology from the 27 ensemble members of NCEP’s Global Data Assimilation System (GDAS) at 0.5 degree spatial resolution. Clear-sky radiative heating rate profiles were then calculated along each of the trajectories using the same methodology as for over Ascension Island except using the temperature and humidity profiles from MERRA-2. Shinozuka et al. (2019) demonstrated good agreement in the SSA between GEOS and aircraft observations over the Southeast Atlantic Ocean so the original MERRA-2 SSA was used. Only clear sky was evaluated as MERRA-2 does not provide the necessary cloud microphysical parameters for RRTM. While there is the potential to gain this information from satellite observations, these observations would lack an appropriate vertical resolution and there would likely be inconsistencies between the thermodynamic profiles in MERRA-2 and the cloud structure in the observations. Given the RRTM results over Ascension Island, SW heating rates due to aerosols along the back trajectories are likely larger than what is presented for the clear sky scenario.

The ensemble mean heating rate due to aerosols is shown in Figure 11a and the ensemble standard deviation in Figure 11b. It is readily apparent that the SW heating due to aerosols is lower in magnitude than over Ascension Island using the observed thermodynamic profiles. Additionally, the location of the heating is limited to roughly below 900 hPa (Figure 11a), which is the height of the temperature inversion and an abrupt change in relative humidity in MERRA-2. Contrary to the observations over Ascension Island, the primary aerosol layer along the trajectory is located entirely above the inversion layer as opposed to being within or below for the RRTM runs above the site. It is known that the boundary layer is too deep over Ascension Island in MERRA-2, and thus minimal SW heating occurs in the last few hours of the back trajectory before reaching Ascension Island. Radiatively, this has a large impact on the results. SW heating due to aerosol is no longer maximized within the aerosol layer but rather at the surface. A secondary maxima in the SW heating due to aerosol, around 0.5 K day\(^{-1}\) is present aloft two, three, and five days prior to aerosol reaching Ascension Island. Though small in magnitude, this heating has the potential to alter the thermodynamic profile and weaken the temperature inversion. Further out in time, AOD decreases and therefore there is minimal heating due to aerosol through the result of the week prior to the trajectory reaching Ascension Island.

As expected based on the trajectories, there is minimal spread in the SW heating due to aerosols within the first few days before arriving at Ascension Island. Prior to this, four to five days before reaching Ascension Island, some ensemble members have a more northerly component and loop around the equator while others continue to have a northeasterly trajectory. It is at this point that there is a discrepancy in the temperature, humidity, and aerosol profiles and thus the standard deviation is maximized near the surface at just under 1 K day\(^{-1}\) (Figure 11b). Greater than five days out, the location of the back trajectories are so varied that the standard deviation exceeds the mean SW heating rate.
4 Summary and Conclusions

The interplay between clouds, aerosols, and radiation is a source of uncertainty within the atmospheric science community and within general circulation models, particularly in the southeast Atlantic region. In this study, an idealized approach was used to quantify the contribution of clouds and biomass burning aerosol to heating within the atmospheric column located above Ascension Island in connection with the LASIC campaign conducted by DOE’s ARM program. The field campaign included the deployment of the AMF1 on Ascension Island that spanned two biomass burning seasons with the highest aerosol loading present during August 2016 followed by September 2017. An assessment of aerosols within the MERRA-2 reanalysis revealed good agreement in AOD compared to AMF1 and AERONET observations, likely due to the assimilation of AOD from MODIS. However, the SSA was too high in MERRA-2, impacting the absorption of SW radiation, and therefore heating, within the atmospheric column. This was mitigated in the radiation transfer experiments by adjusting the SSA to be aligned with the observed relative humidity and reducing the SSA for organic carbon based upon observations to mock that of brown carbon. Given the greater depth of the boundary layer in MERRA-2, it is also possible that the vertical distribution of aerosol in MERRA-2 is not completely realistic.

Due to the uncertainty of the SSA, a range of possible SW heating rates due to aerosols were calculated. On average, the maximum local aerosol SW heating within the column over the course of the biomass burning season likely ranges from 2 to 4 K per day. Local heating rates are sensitive to the thickness of the aerosol plume, and when integrated across the atmospheric column heating due to aerosols can be just as important on days that have a thick but not dense aerosol layer. There is variability in the heating due to aerosol as a result of day-to-day and seasonal fluctuations in aerosol loading and cloud cover as the large-scale circulation and presence of wildfires in southern Africa influence the AOD over Ascension Island. Overall, black carbon is indeed responsible for most of the SW absorption, though clouds also contribute. Biomass burning aerosols and clouds are typically located in distinct layers though at times, biomass burning can extend to the surface. On days with the biomass burning aerosol plume overhead, interaction between SW radiation, clouds, and aerosols result in an enhancement of heating within the aerosol plume on the order of 0.5 K per day. Any heating with the atmospheric column due to aerosol is not offset by additional LW radiational cooling.

There are a few limitations to this study that are worth noting and perhaps expanding upon in future work. 1) Despite the fact that other processes within the atmosphere can respond to the presence of aerosol and the resulting heating, radiation transfer was isolated here. It was assumed that all SW heating would go into altering the temperature profile when in reality, some energy could be lost to other processes such as latent heating. Furthermore, vertical mixing was not accounted for, which can also alter the temperature profile. 2) A single grid point was used to represent the entire southeast Atlantic. The sensitivity experiments for SSA and the difference in the heating rate profiles when using the observed thermodynamic profile as opposed to MERRA-2 demonstrate how sensitive the heating due to biomass burning aerosol is. As the aerosol plume travels, the aerosol ages and tar balls form. This changes the scattering versus absorbing properties and indicates that the heating right off the coast of Africa can be very different from over Ascension Island. 3) A simplified
representation of aerosols in the radiation transfer experiments. At the present time, RRTM only allows for one aerosol type to be characterized in each vertical layer. Not only did this result in a weighting of aerosol properties based on the species, but it also eliminated the ability to characterize the SSA based on wavelength. While a sensitivity study could be completed to quantify the impact of wavelength on the SSA, and therefore the heating rate, the results would likely not yield information that is any more realistic than what was presented here given the dominance of SW radiation centered around 550 nm.

One important implication of the present study is the dependence of the clear-sky (and presumably cloudy-sky) heating rates upon the exact trajectory experienced by the biomass burning plume as it moves from its source region to the Southeastern Atlantic. Clear-sky heating rates varied considerably depending upon trajectories dictated by the large-scale flow, which suggests that there may be an important scale interaction operating in this region. The length of the trajectory from the source region coupled with the loading of black carbon may be an important parameter in facilitating changes in the cloud structure across the Southeastern Atlantic. The most significant anthropogenic alterations to the natural stratocumulus and transition stratocumulus offshore might result from a plume that possesses a large amount of black carbon and follows a long trajectory across the stratocumulus region as it moves away from the African coast.

Ultimately, the goal is to determine how heating due to biomass burning aerosol influences the formation and transition of marine stratocumulus to trade cumulus. Future work includes relating the clear and cloudy sky heating rates along the back trajectories to cloud structure. The goal is ultimately to determine how aerosols impacted the thermodynamic environment and the formation and transition of marine stratocumulus to trade cumulus. A Lagrangian approach to study the interaction between clouds and aerosols in this region was also recommended by Diamond et al. (2018).

Data Availability

MERRA-2 data (GMAO, 2015a, 2015b) are available at https://disc.gsfc.nasa.gov/ and AMF1 data are available at https://www.archive.arm.gov/discovery/.

Author Contribution

ABMC and MAM developed the methodology, completed the analysis, and wrote the manuscript. ABMC performed the RRTM runs and created all figures. LCT provided quality controlled INTERPOSONDE data and MPJ and MW processed and provided data for the cloud fields. All authors except MW edited the manuscript.
Competing Interests

The authors declare that they have no conflict of interest.

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References


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### Tables

**Table 1:** Monthly mean maximum heating rate within the column due to clouds and aerosols in K day$^{-1}$ and the pressure where the maximum occurs. Italicized values in parentheses for all aerosols are results with the decreased SSA.

<table>
<thead>
<tr>
<th></th>
<th>August 2016</th>
<th>August 2017</th>
<th>September 2016</th>
<th>September 2017</th>
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<tr>
<td>M2 SSA</td>
<td>2.39, 840 hPa</td>
<td>2.05, 870 hPa</td>
<td>2.15, 870 hPa</td>
<td>1.99, 663 hPa</td>
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<tr>
<td>M2 OC SSA * 0.85</td>
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<td>2.32, 840 hPa</td>
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<td>1.99, 663 hPa</td>
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<td><strong>Clouds</strong></td>
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<td>2.20, 870 hPa</td>
<td>2.20, 901 hPa</td>
<td>2.13, 870 hPa</td>
</tr>
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</table>
Figure 1: Daily mean aerosol optical depth from the AMF1 (black), AERONET (AER, red), and MERRA-2 (M2, blue) at Ascension Island during (a) August 2016, (b) September 2016, (c) August 2017, and (d) September 2017. Correlation and bias for MERRA-2 is with respect to AERONET observations.
Figure 2: Vertical profile of the mixing ratio of black and organic carbon for the (a) 2016 and (b) 2017 biomass burning seasons from MERRA-2. Black contours indicate a cloud fraction of 0.25.
Figure 3: 10 day back trajectories of a parcel originating at 2 km over Ascension Island colour coded based on the AOD on the start date for (a) August 2016, (b) September 2016, (c) August 2017, and (d) September 2017.
Figure 4: Hourly vertical profiles of (a) temperature and (b) relative humidity over Ascension Island during August 2016.
Figure 5: (a) Cloud water path and (b) liquid effective radius over Ascension Island during August 2016 as calculated by the MICROBASE algorithm.
Figure 6: SW heating due to aerosols based on the single scattering albedo (SSA) in (a, d) MERRA-2, (b, e) the SSA for organic carbon in MERRA-2 multiplied by 0.85, and the SSA albedo in MERRA-2 rescaled based on the observed humidity profile over Ascension Island during August 2016 under (a, b, c) clear and (d, e, f) cloudy skies.
Figure 7: SW heating due to black carbon under (a) clear and (b) cloudy skies Ascension Island during August 2016.
Figure 8: Enhancement of SW heating due to (a) all aerosols and (b) black carbon in the presence of clouds over Ascension Island during August 2016.
Figure 9: (a) LW cooling as a result of increased temperature from SW heating due to aerosols and (b) the net heating rate due to aerosols over Ascension Island during August 2016.
Figure 10: Seven day HYSLPIT back trajectories forced with meteorology from the 27 ensembles of the GDAS, originating at 2 km above Ascension Island at 13z on 13 August 2016.
Figure 11: (a) The mean SW heating rate profile due to aerosols along the ensemble of back trajectories displayed in Figure 10 and (b) the standard deviation.