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Key Points:

- The low single scattering albedo in biomass burning aerosol (BBA) over southern West Africa results from the presence of refractory black carbon
- The brown carbon contribution to the aerosol absorption is negligible in these well-aged BBA plumes
- Accounting for the diversity in black carbon mixing state is needed to accurately estimate BBA optical properties

Supporting Information:

- Supporting Information S1

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Unexpected Biomass Burning Aerosol Absorption Enhancement Explained by Black Carbon Mixing State

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Abstract Direct and semi-direct radiative effects of biomass burning aerosols (BBA) from southern and central African fires are still widely debated, in particular because climate models have been unsuccessful in reproducing the low single scattering albedo in BBA over the eastern Atlantic Ocean. Using state-of-the-art airborne in situ measurements and Mie scattering simulations, we demonstrate that low single scattering albedo in well-aged BBA plumes over southern West Africa results from the presence of strongly absorbing refractory black carbon (rBC), whereas the brown carbon contribution to the BBA absorption is negligible. Coatings enhance light absorption by rBC-containing particles by up to 210%. Our results show that accounting for the diversity in black carbon mixing state by combining internal and external configurations is needed to accurately estimate the optical properties and henceforth the shortwave direct radiative effect and heating rate of BBA over southern West Africa.

Plain Language Summary Extensive seasonal fires over southern and central Africa result in the transport of massive amounts of biomass burning aerosols over huge areas of the eastern Atlantic Ocean. Recent field observations highlight that biomass burning aerosols transported from the coast of southern Africa to the far north over southern West Africa were characterized by low single scattering albedo. This finding is of paramount interest, because radiative heating within the absorbing aerosol layer is hypothesized to affect the low cloud deck over this specific region and may ultimately influence the large-scale circulation. However, debate remains about the causes of the low single scattering albedo by biomass burning aerosols, causing ambiguous parameterizations of their optical properties in climate models. Here we present simultaneous airborne measurements of the composition and optical properties of biomass burning aerosols transported over southern West Africa. We show that black carbon particles dominated the light absorption by biomass burning aerosols at mid-visible wavelengths. Our findings indicate that the black carbon mixing state plays a significant role in the aerosol optical properties and may be an important modulator to be considered in climate models for simulating direct and semi-direct radiative effects of biomass burning aerosol over southern West Africa.

1. Introduction

Biomass burning aerosols (BBAs) are an important component of the climate system due to their strong absorption of solar radiation and the associated impact on the atmospheric energy budget and cloud fraction (i.e., direct and semi-direct aerosol radiative effects, DRE and SDE) (Myhre, Shindell, et al., 2013; Wilcox, 2012). Extensive seasonal fires result in the transport of massive amounts of BBA from southern and central Africa over huge areas of the eastern Atlantic Ocean (Haslett et al., 2019; Menut et al., 2018; Zuidema et al., 2018), which constitutes the Earth's region with the largest aerosol optical depths above clouds (Peers et al., 2016; Waquet et al., 2013). Over this specific region, shortwave heating within the absorbing aerosol layer is hypothesized to affect the low cloud deck through modification of the atmospheric stability (Adebiyi & Zuidema, 2016; Herbert et al., 2020; Sakaeda et al., 2011; Wilcox, 2010) and may ultimately influence the large-scale circulation (Randles & Ramaswamy, 2010). A number of field campaigns—including Dynamics-Aerosol-Chemistry-Clouds Interactions in West Africa (DACCIWA),

Aerosols, Radiation and Clouds in southern Africa (AEROCLO-SA), Layered Atlantic Smoke Interactions with Clouds (LASIC), Observations of Aerosols above Clouds and their InteractionS (ORACLES), and Cloud and Aerosols Radiative Impact and Forcing (CLARIFY)—have recently been carried out over the eastern Atlantic Ocean, with the aim to better quantify the BBA radiative effects (Flamant, Knippertz, et al., 2018; Formenti et al., 2019; Zuidema et al., 2016, 2018). Both in situ and remote-sensing observations acquired during these campaigns have all emphasized a strong shortwave (SW) absorption (low single scattering albedo, SSA) in BBA transported from the coast of southern Africa to the far north over southern West Africa (SWA) (Chylek et al., 2019; Denjean et al., 2020; Pistone et al., 2019; Wu et al., 2020; Zuidema et al., 2018). This means that BBA over this climatically important region is more absorbing than is currently represented in climate models (Mallet et al., 2019; Peers et al., 2016; Stier et al., 2013). However, debate remains about the causes of this low SSA in these BBA plumes.

The absorption of sunlight by BBA particles is largely driven by refractory black carbon (rBC) (Bond et al., 2004). Coating of rBC-containing particles with light scattering material can enhance the absorption of the rBC core due to the so-called “lensing effect” and can thus amplify their impact (Bond et al., 2013). Absorption enhancement (E_{abs}) is defined as the absorption by mixed rBC-containing particles relative to absorption by the same amount of rBC in pure, uncoated particles. Currently, there are only few—and partly contradictory—studies reporting E_{abs} of ambient BBA. Negligible enhancements ($E_{abs} \sim 1$) have been reported in BBA originating from forest fires in North America, even when substantial coatings were present (Cappa et al., 2012; Healy et al., 2015), whereas E_{abs} values up to 1.6 have been observed in the Amazonia region (Lack et al., 2012; Liu et al., 2017). Several hypotheses have been put forward to explain this variability: (1) A restructuring of rBC-containing particles from fractal morphologies towards a compact core could counteract the lensing effect (Kahnert & Kanngießner, 2020; Shingler et al., 2016); (2) rBC-containing particles are not entirely coated but instead exhibit an aggregate structure with non-rBC material (Liu & Mishchenko, 2007), thus leading to overall little change in absorption by rBC-containing particles when internally mixed (Wu et al., 2018); (3) BBA absorption enhancement is partly attributed to the presence of light-absorbing organic aerosols termed brown carbon (BrC) (Brown et al., 2018; Chylek et al., 2019); (4) accounting for mixing-state diversity within the rBC-containing particle population (i.e., the distribution of coating material across the ensemble of rBC-containing particles), as has been done by Fierce et al. (2016) and Liu et al. (2017), is necessary to accurately quantify the average absorption. In part because of the paucity of comprehensive ambient measurements of rBC mixing state in BBA plumes, the mechanisms responsible for the large variation of E_{abs} remain elusive. In addition, previous studies lacked assessing the effect of rBC mixing state on the SSA , which is a key parameter for the quantification of DRE and SDE. As a result, the representation of BBA optical properties in current climate models is uncertain, and many models assume that all rBC-containing particles are externally mixed (Chung et al., 2012; Myhre, Samset, et al., 2013) or use a constant E_{abs} (typically 1.5 as recommended in Bond & Bergstrom, 2006) to account for coating by other species. Here we use airborne in situ measurements to demonstrate how the rBC mixing state impacts key-climate relevant optical properties in BBA plumes. Our results highlight that the mixing state of rBC-containing particles may be an important modulator to be considered in climate models for simulating BBA absorbing properties, DRE and SDE over the eastern Atlantic Ocean.

2. Methods

2.1. Biomass Burning Source and Transport to SWA

As part of the DACCIWA project (Knippertz et al., 2015), the ATR-42 airborne platform of SAFIRE performed 20 flights over the Gulf of Guinea and inland from 29 June to 16 July 2016 (Flamant, Knippertz, et al., 2018). Aerosol and trace gas measurements revealed two distinct BBA layers over SWA (Denjean et al., 2020; Flamant, Knippertz, et al., 2018; Haslett et al., 2019), namely, (i) a lower layer located in the marine boundary layer (BL-BBA) and (ii) an upper, about 2 km-thick layer centered at ~ 3 km above sea level (FT-BBA). In the DACCIWA region, the widespread low level clouds commonly lead to a decoupled BL which top is located at about 500 m above sea level deep away from the coast and about 800 m at the coast (Flamant, Deroubaix, et al., 2018). Denjean et al. (2020) examined measurements of BBA size distribution and optical properties from straight level runs (SLRs) and vertical profiles. Here

we use the same 13 SLRs measurements as in Denjean et al. (2020). Supporting information Figure S1 and Table S1 summarize the geographical location, date, time, and altitude of these 13 SLRs.

The source locations of BBA have been analyzed using the chemistry-transport models CHIMERE, COSMO-ART, GEOS-Chem, and WRF (Flamant, Knippertz, et al., 2018; Haslett et al., 2019; Menut et al., 2018). The results indicate that all BBA originated from fires in the tropical and subtropical grasslands, savannas, and scrubland areas of southern and central Africa. Additional assessments using the backward trajectories from the HYSPLIT model in conjunction with the fire spot data from the MODIS satellite confirm this origin, as shown in Figure S1. The BBA plumes were sampled by the aircraft after estimated aging larger than 1 week.

2.2. Aircraft Instrumentation and Data Analysis

The aircraft was extensively equipped to measure gas and aerosol properties. Onboard aerosol instruments sampled ambient air via stainless steel tubes through the Community Aerosol Inlet (CAI). This is an isokinetic and isoaxial inlet with a 50% sampling efficiency for particles with a diameter of 5 μm (Denjean et al., 2016).

Here, we use in situ data obtained using a single-particle soot photometer (SP2). The SP2 measures the mass of rBC within single aerosol particles in the size range of 70–600 nm based on laser-induced incandescence. The mass equivalent diameter of rBC cores ($D_{\text{core rBC}}$) was derived from the mass of rBC per particle assuming the density of rBC as 1.8 g cm^{-3} based on the recommendations of Bond and Bergstrom (2006). SP2 data were also used to estimate the optical diameter of rBC-containing particles ($D_{\text{total rBC}}$), using a slightly modified version of the leading-edge-only method described by Gao et al. (2007) as implemented by Laborde et al. (2012). A detailed description of this method, including a description of the SP2 calibration and the method used for data quality control, is shown in section S1. The rBC mixing state of each particle can be expressed as a mass mixing ratio, that is, the mass of non-rBC material in a rBC-containing particle divided by the mass of rBC. The bulk mass mixing ratio ($M_{R,\text{bulk}}$) for a given time window is then calculated as follows:

$$M_{R,\text{bulk}} = \left(\frac{\sum_i D_{\text{total rBC},i}^3}{\sum_i D_{\text{core rBC},i}^3} - 1 \right) \times \frac{\rho_{\text{coating}}}{\rho_{\text{rBC}}}$$

where ρ_{coating} is the coating density assumed to be 1.5 g cm^{-3} .

The aerosol chemical composition for non-refractive compounds was measured using an Aerodyne compact Time-of-Flight Aerosol Mass Spectrometer (HR-ToF-AMS). For a more detailed description of the instrument and data analysis, the reader is referred to Brito et al. (2018). The aerosol number size distribution was measured with a custom-built scanning mobility sizer spectrometer (SMPS, 20–485 nm), an ultra-high sensitivity aerosol spectrometer (UHSAS, 0.04–1 μm), and an optical particle counter (OPC GRIMM model 1.109, 0.3–32 μm). The aerosol absorption coefficient (σ_{abs}) was measured using a three-wavelength particle soot absorption photometer (PSAP) with an uncertainty of 8% (Davies et al., 2019). The PSAP measures changes of filter attenuation due to the collection of aerosol deposited on the filter, which were corrected for the scattering artifacts according to the Virkkula (2010) method. The particle scattering coefficient (σ_{scat}) was measured using a TSI 3563 3-Wavelength Integrating Nephelometer and corrected for angular truncation error by using Lorenz-Mie scattering model, number size distributions, and the complex refractive index provided in Denjean et al. (2020) for the same sampled BBA. Uncertainty in σ_{scat} measured with the nephelometer is estimated to be 5% (Muller et al., 2011). The concentration of carbon monoxide (CO) was measured with an ultraviolet and infrared analyzer (Picarro).

The mass absorption cross section of rBC (MAC_{rBC}) and the enhancement factor of light absorption of rBC due to the lensing effect (E_{abs}) are defined, respectively, as follows:

$$MAC_{\text{rBC}}(\lambda) = \frac{\sigma_{\text{abs}}(\lambda)}{M_{\text{rBC}}}$$

and

$$E_{\text{abs}}(\lambda) = \frac{MAC_{\text{rBC}}(\lambda)}{MAC_{\text{uncoated rBC}}(\lambda)}$$

where λ is the wavelength. The uncertainty in MAC_{rBC} and E_{abs} is $\pm 12\%$ by propagating the uncertainties of M_{rBC} and σ_{abs} . The reference $MAC_{\text{uncoated rBC}}$ is determined by extrapolating the observed relationship

between coating thicknesses ($M_{R,bulk}$) and MAC_{rBC} to the limit of no coating material (Figure S2). The resulting $MAC_{uncoated\ rBC}$ value for BBA during DACCIWA is $7.7\text{ m}^2\text{ g}^{-1}$ at 550 nm. This value agrees with the value of $7.5 \pm 1.2\text{ m}^2\text{ g}^{-1}$ reported in the review of Bond and Bergstrom (2006).

The absorption Angstrom exponent (AAE), which describes the wavelength dependence of aerosol absorption, is calculated between 450 and 660 nm as follows.

$$AAE = -\frac{\ln\left(\frac{\sigma_{abs}(450)}{\sigma_{abs}(660)}\right)}{\ln\left(\frac{450}{660}\right)}$$

The uncertainty in AAE is estimated to be 5% (Davies et al., 2019).

The SSA and asymmetry factor (g) are defined respectively as follows:

$$SSA = \frac{\sigma_{scat}(\lambda)}{\sigma_{scat}(\lambda) + \sigma_{abs}(\lambda)}$$

and

$$g(\lambda) = \frac{1}{2} \int_0^\pi \cos(\theta) \sin(\theta) P(\theta, \lambda) d(\theta)$$

with $P(\theta, \lambda)$ the scattering phase function and θ the scattering angle. g was calculated using Lorenz-Mie scattering model and measured size distribution (Denjean et al., 2020). The uncertainties in SSA and g are estimated to be 9% and 13%, respectively, by propagating the uncertainties of σ_{scat} and σ_{abs} .

The aerosol properties were measured at dry conditions ($RH < 30\%$). Data obtained in clouds were excluded from analysis to avoid inlet artifacts. Aerosol data are reported at standard temperature and pressure conditions (273.15 K and 1013.25 hPa).

2.3. Modeling BBA Optical and Radiative Properties

2.3.1. Lorenz-Mie Scattering Calculations

The influence of observed coatings of rBC-containing particles on BBA optical properties relevant to parameterizing aerosol radiative effects in models (MAC , SSA , and g) was investigated using a Lorenz-Mie scattering model with the core-shell assumption (Bohren & Huffman, 1983). The rBC-containing particles are assumed to have concentric-sphere morphology with a spherical rBC core embedded in a shell of internally mixed nonabsorbing material. The use of Mie theory, which assumes that the particles are spherical, is supported by the overall range of AAE (see section 3.2) that is typical for rBC particles in a core-shell morphology, whereas much larger AAE would be expected for more complex morphologies (Liu & Mishchenko, 2007). The calculations were performed at the wavelength of 550 nm for each BBA plume using observations, namely, the size distributions of rBC-containing and rBC-free particles and the mass fraction of rBC-containing particles in the aerosol population (f_{rBC}). The size distribution of rBC-free particles was derived by subtracting the SP2-measured size distribution of rBC-containing particles (which includes both coated and noncoated particles) from the total aerosol size distribution measured using the SMPS, UHSAS, and OPC GRIMM. The refractive index of the rBC core at the wavelength of 550 nm was assumed to be $1.95-0.79i$ (Bond & Bergstrom, 2006). Because the analysis of aerosol chemical composition indicates the predominance of organic and sulfate species in BBA (see section 3.1), the refractive index of the coating material and BC-free particles was assumed to be $1.50-0i$ (Fuller et al., 1999). Densities of 1.8, 1.2, and 1.7 g cm^{-3} were used for rBC, organics, and inorganics, respectively, for the conversion of mass fractions to volume fraction.

2.3.2. ecRad Radiative Transfer Model

The offline version of the radiative code ecRad (Hogan & Bozzo, 2018), operationally used at the European Center for Medium-Range Weather Forecasts (ECMWF), was used to compute the SW radiative forcing of BBA in clear-sky and cloudy-sky conditions. To this end, an average atmospheric profile was built from the airborne measurements taken during the campaign, including vertical profiles of pressure, temperature,

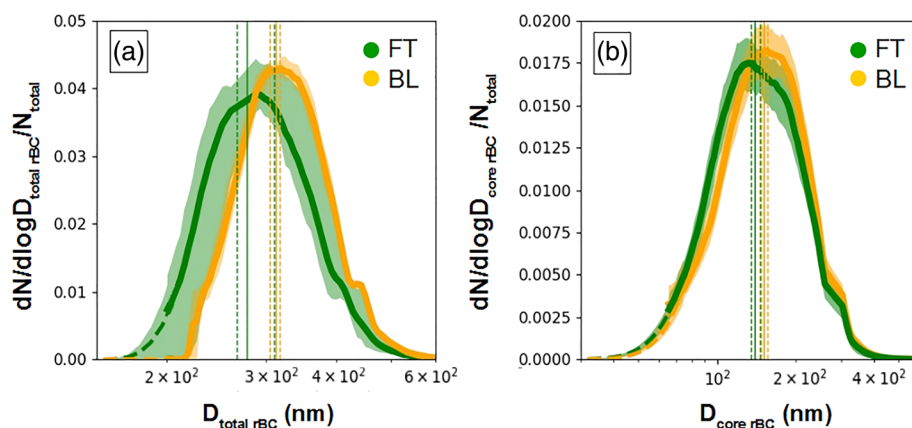


Figure 1. Number size distributions normalized by the total number concentration of (a) coated rBC particles and (b) rBC cores. The dashed lines are Gaussian distributions fitted to the mass distribution and converted to the corresponding number distributions. Colored areas represent the 25th and 75th percentiles of the number size distributions. The vertical bars show the median values (solid lines) and the 25th and 75th percentiles (dashed lines) of the geometric diameter for the two BBA categories.

relative humidity, and CO_2 . Vertical distributions of BBA mass concentration measured in situ during vertical profiles (Denjean et al., 2020) were averaged over the campaign and used as input in ecRad (Figure S2). BBA optical properties (MEC , SSA , and g) at 550 nm were constrained either from our observations (Denjean et al., 2020) or from the three optical models presented in section 2.3.1. These input parameters were extended to the full SW spectrum assuming the same spectral variations as the rBC natively implemented in ecRad. The SW DRE at the top of the atmosphere (TOA) and atmospheric heating rates due to BBA were computed as the difference between simulations including aerosols and reference aerosol-free simulations. The simulations were performed assuming a surface albedo of 0.25, typical of the rainforest above which the measurements were taken (Moody et al., 2005). The cosine of the solar zenith angle was set to 0.8, which corresponds to the weighted daily average value during the campaign. The simulations conducted under cloudy-sky conditions assumed a cloud optical thickness of 10 located at 2.4 km, at the top of the boundary layer (2.4 km). The cloud optical properties were computed according to the parameterization of Edwards and Slingo (1996). To this end, the cloud droplet effective radius was assumed to be $5 \mu\text{m}$, which is consistent with the observations by Taylor et al. (2019) during DACCIWA.

3. Results

3.1. rBC Mixing State

Figure 1 shows size distributions of rBC-containing particles (panel A) and rBC cores only (panel B) for the boundary layer (BL) and free troposphere (FT). The modal $D_{total \text{ rBC}}$ decreases from ~ 310 in the BL to ~ 280 nm in the FT. This is mainly attributed to the increasing coating thickness in BL-BBA, since the geometric diameters of rBC cores are ~ 140 and ~ 152 nm, respectively. The coating thicknesses translate into an average $M_{R,bulk}$ of ~ 4.7 and ~ 6.0 in FT-BBA and BL-BBA, respectively. This indicates that rBC-containing particles were mostly constituted of non-rBC materials, making up more than four times the rBC mass in these rBC-containing particles. In addition, the percentage of coated rBC particles and the average coating thicknesses, respectively, as observed in the BL (91% and 159 nm), are larger than those measured in the FT (88% and 137 nm).

The variability of rBC coating thickness could result either from differences in the sources or from aging processes affecting BBA plumes during the long-range transport of BBA over SWA. We have compared the correlation between M_{rBC} and ΔCO ($\Delta\text{CO} = \text{plume CO} - \text{background CO}$) to gain information about the BBA sources (Spackman et al., 2008). In addition, the M_{rBC} to ΔCO ratio is useful for evaluating the potential influence of wet removal processes on BBA, as precipitation scavenging affects rBC-containing particles but does not affect CO (Moteki et al., 2012; Taylor et al., 2014). The background value of CO was

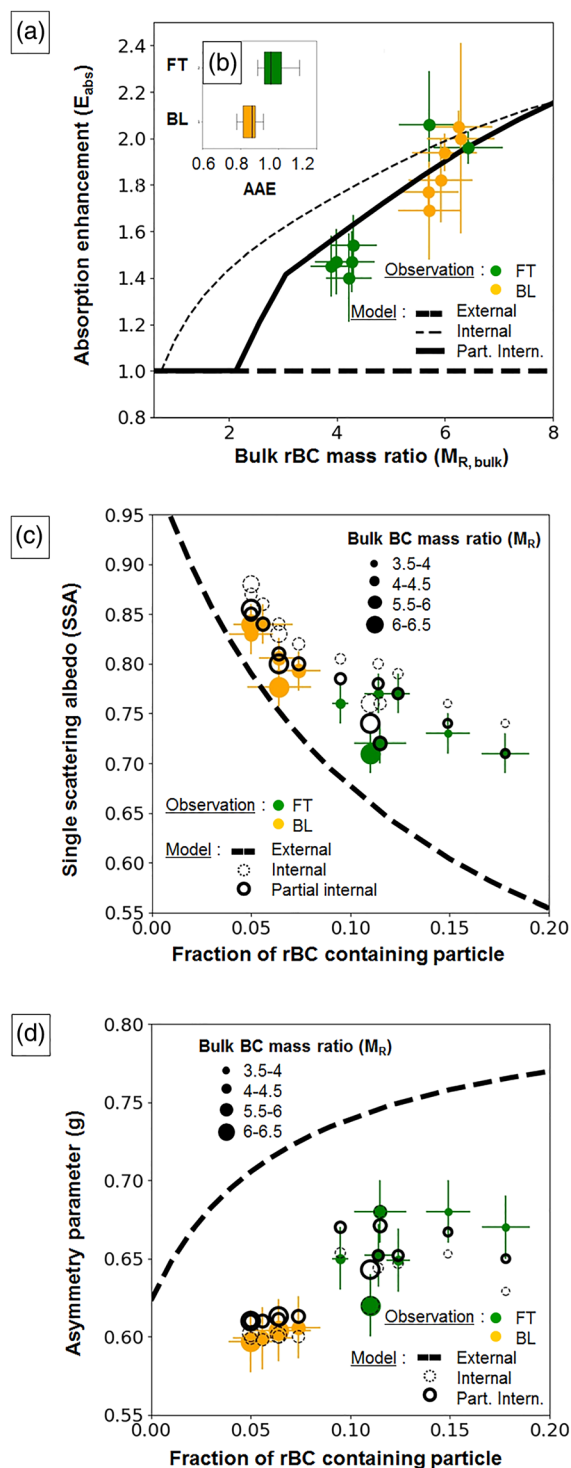


Figure 2. Sensitivity of BBA optical properties to rBC mixing state. Measured and modeled employing different rBC population mixing state (external, internal, and partial internal) of (a) absorption enhancement at 550 nm as a function of $M_{R,bulk}$, (b) Angstrom absorption exponent calculated between 450 and 660 nm, (c) single scattering albedo at 550 nm as a function of fraction of rBC containing particle in the total aerosol mass, and (d) asymmetry parameter at 550 nm as a function of mass fraction of rBC-containing particles in the total aerosol mass. Error bars are one standard deviation over the mean.

estimated with the intercept of rBC versus CO using all data points available within the plume. It ranged from 80 to 95 ppb depending on the altitude of the BBA plume. The correlations between M_{rBC} and ΔCO were similar for BL-BBA and FT-BBA (Figure S4), with mean values of the M_{rBC} to ΔCO ratio of 7.4 and 7.6 $\text{ng m}^{-3} \text{ppb}^{-1}$, respectively. These values are comparable to those previously reported in BBA from southern and central African fires near emission sources (7.4 $\text{ng m}^{-3} \text{ppb}^{-1}$ in the review of Andreae & Merlet, 2001) and SWA (7.4 $\text{ng m}^{-3} \text{ppb}^{-1}$ by Capes et al., 2008). This suggests that BBA-BL and BBA-FT in the present work had similar source locations and had not been influenced by wet removal processes during transport.

Measurements of the aerosol composition during DACCIWA revealed a larger contribution of sulfate in BL-BBA (~27%) (Figure S5), which was traced back to a strong influence of marine emissions (Haslett et al., 2019). In addition, given the large loadings of organic aerosol in BBA over SWA (greater than 50%), we expect organic aerosol to contribute significantly to the coating over rBC cores. The organic aerosol within BBA plumes over SWA was highly oxidized, being dominated by a peak at m/z 44 and with low contribution of m/z 60 towards the total organic aerosol mass (Figure S6), as discussed in Haslett et al. (2019). Although both BBA layers were found to be highly oxidized, the f_{44}/f_{43} (the fraction of the organic aerosol mass spectrum signal at m/z 44 and 43, respectively) is negatively correlated with the $M_{R,bulk}$ (Figure S7). In general, f_{44}/f_{43} increases with altitude. There is an exception for two FT-BBA plumes with high $M_{R,bulk}$, but these coating thicknesses were also correlated with high f_{44}/f_{43} . Two hypotheses could explain the tendency of decreasing f_{44}/f_{43} ratio with increasing coating thickness: (i) the condensation of less-oxidized material onto preexisting highly aged organics favored by the lower temperature in the FT and/or (ii) fragmentation of highly oxidized organics induced by enhanced photochemical aging in the FT and subsequent evaporation (e.g., Ng et al., 2010; Palm et al., 2018). It should be noted here that the chemical composition was measured for bulk aerosols. It cannot be ruled out that the bulk f_{44}/f_{43} ratio is a poor indicator of the composition of the coating material on individual rBC-containing particles. While the exact reason for the variability in $M_{R,bulk}$ remains unclear, our results strongly suggest that atmospheric aging alters rBC mixing state through processes that are dynamic- and altitude-dependent.

3.2. Exploring the Sensitivity of BBA Optical Properties to rBC Mixing State

In Figure 2a, a large absorption enhancement in both BL-BBA and FT-BBA is evident. E_{abs} values range between 1.4 and 2.1, which correspond to MAC_{rBC} values between 10.8 and 16.2 $\text{m}^2 \text{g}^{-1}$. Our measured E_{abs} in BL-BBA is larger than the value of 1.5 commonly adopted in climate models when internal mixing of rBC is accounted for (Bond et al., 2013; Chung et al., 2012; Myhre, Samset, et al., 2013). The observed increase of E_{abs} with increasing $M_{R,bulk}$ indicates that a lensing-driven absorption enhancement is substantial in BBA. In field observations, significant absorption and strong spectral dependence (values of $AAE > 1.7$) in BBA have frequently been attributed to the presence of BrC (Chakrabarty et al., 2010; Kirchstetter et al., 2004; Pokhrel et al., 2016; Romonosky et al., 2019; Sandradewi et al., 2008). Our measurements suggest that the contribution of BrC to light absorption is negligible, since

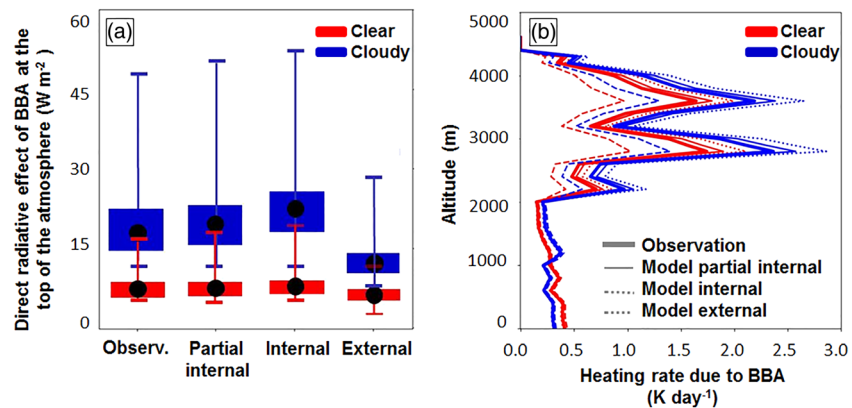


Figure 3. (a) Direct radiative effect (DRE) at the top of the atmosphere and (b) heating rate profile due to the transport of both BL-BBA and FT-BBA, calculated from measured and modeled BBA optical properties assuming different representation of rBC mixing state. On the left, the boxes mark the respective median values (middle points), the 25th and 75th percentiles (left and right box edges), and the minimum and maximum values (error bars). On the right, heating rate profiles are averaged over the campaign.

AAE values range from 0.8 to 1.1 (Figure 2b). These results imply that BBA absorption properties were largely dominated by the rBC mixing state. Figures 2c and 2d show decreases (increases) in the *SSA* (*g*) of the overall BBA population as f_{rBC} increases. When the air mass was transported in the FT, the *SSA* reached very low values (down to ~ 0.71), and rBC-containing particles contributed to the largest aerosol mass fraction ($\sim 18\%$). Despite this general trend, a large spread in *SSA* and *g* is observed independent of f_{rBC} .

To further investigate the variability on the observed optical properties, the core-shell Mie model was used to calculate E_{abs} , *SSA*, and *g* using the observed f_{rBC} , $D_{core\ rBC}$, and $D_{total\ rBC}$ and by assuming different rBC population mixing state. In the external mixing case, rBC-containing particles were assumed to be entirely uncoated, whereas the coating material is equally distributed across the population of rBC-containing particles in the internal mixing case. In the partial internal mixing case, the diversity in rBC mixing state is accounted for by combining internal and external configurations, parameterized using a simplified function of the observed $M_{R,bulk}$ (Figure S8). When an external mixing is considered, we find that E_{abs} and *SSA* are on average underestimated by $\sim 42\%$ and $\sim 9.8\%$ compared to observations (Figures 2a and 2c, respectively) and *g* overestimated by $\sim 11\%$ (Figure 2d). The fully rBC internal mixing assumption overestimates E_{abs} and *SSA* by $\sim 13\%$ and $\sim 5.2\%$, respectively, and reduces the *g* bias to $\sim 3.6\%$. When both mixing states coexist with the partial internal mixing case, the E_{abs} and *SSA* biases are reduced to $\sim 5.1\%$ and $\sim 2.0\%$, respectively, which are in the range covered by the measurements uncertainties (i.e., 12% and 9%, respectively). This points out that accounting for the diversity of rBC mixing state is essential to properly model aerosol optical properties and that combining external and internal mixtures leads to more representative optical properties of BBA.

4. Implications for Atmospheric Radiation

Assuming that BBA absorption comes exclusively from spherical rBC-containing particles, the Mie scattering model successfully reproduces the observed BBA optical properties. Along with low *AAE* values, this suggests that BrC makes a negligible contribution to BBA optical properties at mid-visible wavelengths for well-aged air masses (larger than 1 week) that have undergone extensive oxidation. A plausible explanation is that BrC is removed during transport through photobleaching, volatilization, and/or cloud-phase reactions (Clarke et al., 2007; Forrister et al., 2015; Lewis et al., 2008), which leads to a higher fraction of rBC-containing particles in the total aerosol mass, lower *AAE*, and lower *SSA* values.

Our findings indicate that the treatment of rBC mixing state plays a significant role in resolving the BBA optical properties over the SWA region. To estimate the radiative importance of resolving the diversity of rBC mixing state in BBA plumes, radiative fluxes were computed with the radiative transfer model ecRad. In Figure 3, we examine the SW DRE at TOA and heating rates in clear-sky and cloudy-sky conditions

caused by BBA over SWA using measured and modeled BBA optical properties. BBA induces an intense positive SW DRE at TOA over the continental SWA in all-sky conditions, with the highest magnitude of the instantaneous forcing reaching $+49 \text{ W m}^{-2}$ for the more intense FT-BBA event. A sensitivity test of DRE to surface albedo suggests that, with a surface albedo of 0.1 closer to that of the ocean, BBA is able to exert either a positive or negative SW DRE at TOA depending on the presence of clouds (Figure S9), which is consistent with a large body of literature focused on this region (De Graaf et al., 2014; Feng & Christopher, 2015; Mallet et al., 2019; Meyer et al., 2013; Zuidema et al., 2016). In addition, the FT-BBA plumes lead to an $\sim 2.4 \text{ K d}^{-1}$ increase in the average instantaneous SW heating rate in cloudy conditions, which can reach $\sim 6.0 \text{ K d}^{-1}$ during extreme cases (Table S2). This increase in SW heating can reduce local relative humidity, implying a negative SDE over the ocean where absorbing BBA layers frequently exist above persistent stratus cloud decks (Ajoku et al., 2020; Herbert et al., 2020; Wilcox, 2012). In addition, the significant radiative impact of BBA highlighted in this study could also have some large implications on the stratus-to-cumulus cloud transition as shown by Deetz et al. (2018) during DACCIWA. Using unrealistic mixing state assumption for rBC-containing particles leads to notable systematic biases in the resulting DRE (heating rates) of $\sim -26\%$ ($\sim -40\%$) for externally and $\sim +11\%$ ($\sim -21\%$) for internally mixing, arising from underestimation and overestimation of the rBC absorption enhancement by the coating, respectively (Figures 3a and 3b). The range of extreme DRE values doubles from ~ 4.5 (11) W m^{-2} in the external mixing case to ~ 7.5 (22) W m^{-2} in the internal mixing case in clear-sky (cloudy) conditions (Figure 3a), reflecting the high sensitivity of BBA DRE to rBC coating thickness. The DRE and heating rates of BBA were successfully reproduced when the diversity of rBC mixing state is accounted for, which is currently unaccounted for in climate models (Chung et al., 2012; Mallet et al., 2019; Myhre, Samset, et al., 2013). Missing or incorrect representation of the internal rBC mixing state can thus contribute to a large underestimation or overestimation of BBA DRE and SW heating rates calculated by climate models in this region, depending on the mixing state assumption used.

5. Concluding Remarks

Using state-of-the-art airborne in situ measurements in conjunction with Mie scattering simulations, we demonstrate that the low SSA in BBA plumes over SWA results from the presence of strongly absorbing rBC-containing particles, whereas the BrC contribution to the BBA absorption is negligible. To date, climate models have been unsuccessful in reproducing the SSA in BBA plumes over eastern Atlantic Ocean (Lacagnina et al., 2015; Mallet et al., 2019; Peers et al., 2016; Stier et al., 2013). Our results show that accounting for the complex rBC mixing-state diversity (fraction of coated and uncoated rBC-containing particle and amount of scattering material on the rBC-containing particles) is needed to accurately estimate BBA optical properties. The assumed mixing state of rBC-containing particles controls the magnitude of the SSA and henceforth the SW DRE at TOA and heating rate of BBA. In addition, our results advise against the use of fixed timescales to represent aging of rBC-containing particles, as $M_{R,bulk}$ has been shown to vary with the altitude of BBA plumes, thus depending on local environmental conditions. Even though computational resources do not permit explicit calculation of aging by condensation and coagulation, some climate models have been moving towards aerosol modules that represent aerosol aging using several interacting modes (Aquila et al., 2011; Bauer et al., 2008; Fierce et al., 2016, 2017; Matsui et al., 2013; Stevens & Dastoor, 2019; Wilson et al., 2001). Accounting for a more realistic representation of rBC mixing state should significantly decrease the ambiguity in the radiative and climatic effects of BBA. Further studies testing the sensitivity of regional and global climate models to rBC mixing state representation are needed. Our measurements of rBC mixing state and vertically resolved abundances are broadly applicable to BBA over SWA for improvement in radiative transfer calculations, which would be very helpful to capture the full picture of how BBA modifies the atmospheric energy budget and cloud properties over SWA.

Data Availability Statement

All data used in this study are publicly available on the AERIS Data and Service Center, which can be found online (<http://baobab.sedoo.fr/DACCIWA>). Information on how to obtain ecRad for noncommercial education or research is available at the ecRad website (<https://confluence.ecmwf.int/display/ECRAD>).

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