

Dear Dr. Andreas Petzold,

We want to thank you for the review of the manuscript. Based on your comments, we have made revisions to the original manuscript as follows:

- Papers of Sandradewi et al. (2008), Formenti et al (2011), Petzold et al. (2013) have been incorporated.
- Technical information on the CPC, UHSAS, GRIMM and SMPS have been added.
- A brief comparison of AAE in biomass burning plumes with literature has been included.
- Figure 4, Figure 7 and Appendix 2 have been improved.
- A number of typographical corrections have been made to the text.

We hope that these changes will fulfill your suggestions. The article with tracked changes can be found below.

Yours sincerely,
Cyrielle Denjean

1 Overview of aerosol optical properties over southern 2 West Africa from DACCWA aircraft measurements

3

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18

19 **Abstract.** Southern West Africa (SWA) is an African pollution hotspot but a relatively poorly
20 sampled region of the world. We present an overview of *in-situ* aerosol optical measurements
21 collected over SWA in June and July 2016 as part as of the DACCWA (Dynamics-Aerosol-
22 Chemistry-Clouds Interactions in West Africa) airborne campaign. The aircraft sampled a wide
23 range of air masses, including anthropogenic pollution plumes emitted from the coastal cities,
24 long-range transported biomass burning plumes from Central and Southern Africa and dust plumes
25 from the Sahara and Sahel region, as well as mixtures of these plumes. The specific objective of
26 this work is to characterize the regional variability of the vertical distribution of aerosol particles
27 and their spectral optical properties (single scattering albedo: *SSA*, asymmetry parameter,
28 extinction mass efficiency, scattering Ångström exponent and absorption Ångström exponent:
29 *AAE*). First findings indicate that aerosol optical properties in the planetary boundary layer were
30 dominated by a widespread and persistent biomass burning loading from the Southern
31 Hemisphere. Despite a strong increase of aerosol number concentration in air masses downwind of
32 urban conglomerations, spectral *SSA* were comparable to the background and showed signatures of
33 the absorption characteristics of biomass burning aerosols. In the free troposphere, moderately to
34 strongly absorbing aerosol layers, dominated by either dust or biomass burning particles, occurred

35 occasionally. In aerosol layers dominated by mineral dust particles, *SSA* varied from 0.81 to 0.92 at
36 550 nm depending on the variable proportion of anthropogenic pollution particles externally mixed
37 with the dust. For the layers dominated by biomass burning particles, aerosol particles were
38 significantly more light absorbing than those previously measured in other areas (e.g. Amazonia,
39 North America) with *SSA* ranging from 0.71 to 0.77 at 550 nm. The variability of *SSA* was mainly
40 controlled by variations in aerosol composition rather than in aerosol size distribution.
41 Correspondingly, values of *AAE* ranged from 0.9 to 1.1, suggesting that lens-coated black carbon
42 particles were the dominant absorber in the visible range for these biomass burning aerosols.
43 Comparison with literature shows a consistent picture of increasing absorption enhancement of
44 biomass burning aerosol from emission to remote location and underscores that the evolution of
45 *SSA* occurred a long time after emission.
46 The results presented here build a fundamental basis of knowledge about the aerosol optical
47 properties observed over SWA during the monsoon season and can be used in climate modelling
48 studies and satellite retrievals. In particular and regarding the very high absorbing properties of
49 biomass burning aerosols over SWA, our findings suggest that considering the effect of internal
50 mixing on absorption properties of black carbon particles in climate models should help better
51 assessing the direct and semi-direct radiative effects of biomass burning particles.

52

53 **1. Introduction**

54 Atmospheric aerosols play a crucial role in the climate system by altering the radiation budget
55 through scattering and absorption of solar radiation and by modifying cloud properties and
56 lifetime. Yet considerable uncertainties remain about the contribution of both natural and
57 anthropogenic aerosol to the overall radiative effect (*Boucher et al., 2013*). Large uncertainties are
58 related to the complex and variable properties of aerosol particles that depend on the aerosol
59 source and nature as well as on spatial and temporal variations. During transport in the
60 atmosphere, aerosol particles may undergo physical and chemical aging processes altering the
61 composition and size distribution and henceforth the optical properties and radiative effects. The
62 capability of reproducing this variability in climate models represents a real challenge (*Myhre et*
63 *al., 2013; Stier et al., 2013; Mann et al., 2014*). Therefore, intensive experimental observations in
64 both aerosol source and remote areas are of paramount importance for constraining and evaluating
65 climate models.

66

67 Key parameters from a climate perspective are the aerosol vertical distribution and respective
68 spectral optical properties. Radiative transfer codes commonly incorporated in climate models and

69 in satellite data retrieval algorithms use single scattering albedo (*SSA*), mass extinction efficiency
70 (*MEE*) and asymmetry factor (*g*) as input parameters. These parameters depend on the aerosol size
71 distribution, the real and imaginary parts of the refractive index (*n-ik*), and the wavelength of
72 incident light, λ . The knowledge of the vertical distribution of these fundamental parameters is
73 crucial to accurately estimate the direct and semi-direct radiative effects of aerosols as well as the
74 vertical structure of atmospheric heating rates resulting from absorption by particles. Above
75 information is also required to retrieve aerosol properties (aerosol optical depth, size distribution)
76 from remote sensing data.

77

78 Southern West Africa (SWA) is one of the most climate-vulnerable region in the world , where the
79 surface temperature is expected to increase by $\sim 3^{\circ}\text{K}$ at the end of the century (2071-2100) in the
80 Coupled Model Intercomparison Project Phase 5 (CMIP5) (Roehrig *et al.*, 2013). It is
81 characterized by a fast-growing population, industrialization and urbanization (Lioussé *et al.*,
82 2014). This is particularly the case along the Guinea Coast where several already large cities are
83 experiencing rapid growth (Knippertz *et al.*, 2015a). Despite these dramatic changes, poor
84 regulation strategies of traffic, industrial and domestic emissions lead to a marked increase of
85 anthropogenic aerosol loading from multiple sources including road traffic, industrial activities,
86 waste burning, ship plumes, domestic fires, power plants, etc. Tangible evidence for regional
87 transport of anthropogenic pollutants associated with urban emissions has altered air pollution
88 from a local issue to a regional issue and beyond (Deetz *et al.*, 2018; Deroubaix *et al.* 2019). This
89 is particularly the case during summer when land-sea breeze systems can develop and promote the
90 transport of pollutants inland, away from the urbanized coastal strip of SWA (Flamant *et al.*,
91 2018a). In addition to this anthropogenic regional pollution, SWA is impacted by a significant
92 import of aerosols from remote sources. Biomass burning mainly from vegetation fires in Central
93 Africa are advected to SWA in the marine boundary layer and aloft (Mari *et al.*, 2008; Menut *et al.*
94 2018; Haslett *et al.*, 2019). The nearby Sahara desert and the Sahel are large sources of natural
95 wind-blown mineral dust aerosol throughout the year with a peak in springtime (Marticorena and
96 Bergametti, 1996). Biomass burning, dust and anthropogenic pollution aerosols can be mixed
97 along their transport pathways (Flamant *et al.*, 2018a; Deroubaix *et al.* 2019), resulting in
98 complex interactions between physical and chemical processes and even meteorological
99 feedbacks.

100

101 In West Africa, most of the aerosol–radiation interaction studies focused on optical properties of
102 dust and biomass burning aerosols in remote regions far from major sources of anthropogenic

103 pollution aerosol. They include ground-based and airborne field campaigns such as DABEX (Dust
104 and Biomass Experiment, *Haywood et al., 2008*), AMMA (Analysis Multidisciplinary of African
105 Monsoon, *Lebel et al., 2010*), DODO (Dust Outflow and Deposition to the Ocean, *McConnell et*
106 *al., 2008*), SAMUM-1 and SAMUM-2 (Saharan Mineral Dust Experiment, *Heintzenberg, 2009;*
107 *Ansmann et al., 2011*) and AER-D (AERosol Properties – Dust, *Ryder al. 2018*). These projects
108 concluded that the influence of both mineral dust and biomass burning aerosols on the radiation
109 budget is significant over West Africa, implying that meteorological forecast and regional/global
110 climate models should include their different radiative effects for accurate forecasts and climate
111 simulations. Over the Sahel region, *Solmon et al. (2008)* have highlighted the high sensitivity of
112 mineral dust optical properties to precipitation changes at a climatic scale. However, the optical
113 properties of aerosols particles in the complex chemical environment of SWA are barely studied.
114 This is partly due to the historically low level of industrial developments of the region. Motivated
115 by the quickly growing cities along the Guinea Coast, the study of transport, mixing, and feedback
116 processes of aerosol particles is therefore very important for better quantification of aerosol
117 radiative impact at the regional scale and improvement of climate and numerical weather
118 prediction models.

119

120 In this context, the DACCIWA (Dynamics-Aerosol-Chemistry-Clouds Interactions in West Africa,
121 *Knippertz et al., 2015b*) campaign, designed to characterize both natural and anthropogenic
122 emissions over SWA, provides important and unique observations of aerosols in a region much
123 more affected by anthropogenic emissions than previously thought. A comprehensive field
124 campaign took place in June–July 2016 including extensive ground-based (*Kalthoff et al., 2018*)
125 and airborne measurements (*Flamant et al., 2018b*). In this study, we present an overview of *in-*
126 *situ* airborne measurements of the vertical distribution of aerosol particles and their spectral optical
127 properties acquired with the ATR-42 French research aircraft over the Guinea Coast.

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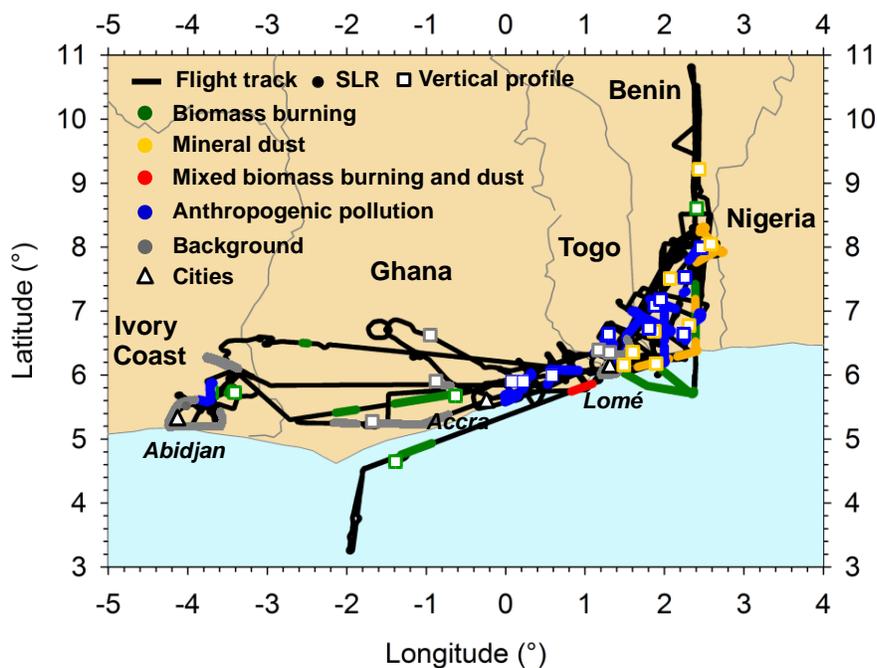
129 Section 2 presents the flight patterns, instrumentation and data analysis. Section 3 provides an
130 overview of the aerosol microphysical and optical properties. The impact of aging and mixing
131 processes on aerosol optical properties is discussed in section 4 before conclusions are presented in
132 section 5.

133

134 **2. Methodology**

135 **2.1. ATR-42 measurements overview**

136 This analysis focuses on flight missions conducted by the ATR-42 aircraft of SAFIRE (Service des
137 Avions Français Instrumentés pour la Recherche en Environnement - the French aircraft service
138 for environmental research) over the Gulf of Guinea and inland. A full description of flight
139 patterns during DACCIWA is given in *Flamant et al. (2018b)*. Here we present results from 15
140 flights focused on the characterization of anthropogenic pollution, dust and biomass burning
141 plumes. The flight tracks are shown in Figure 1 and a summary of flight information is provided in
142 Appendix 1. The sampling strategy generally consisted of two parts: first, vertical soundings were
143 performed from 60 m up to 8 km above mean sea level (amsl) to observe and identify interesting
144 aerosol layers. Subsequently, the identified aerosol layers were probed with the *in-situ* instruments
145 by straight levelled runs (SLR) at fixed flight altitudes.



146

147 **Figure 1. Tracks of the 15 flights analyzed in this study. The colors indicate aircraft flight**
148 **sampling layers dominated by biomass burning (green), mineral dust (orange), mixed dust-**
149 **biomass burning (red), anthropogenic pollution (blue) and background particles (grey) from**
150 **both vertical profiles (squares) and straight and level runs (SLRs; dots).**

151

152 The ATR-42 aircraft was equipped with a wide variety of instrumentation performing gas and
153 aerosol measurements. The measured meteorological parameters include temperature, dew point
154 temperature, pressure, turbulence, relative humidity, as well as wind speed and direction. Gas
155 phase species were sampled through a rear facing ¼ inch Teflon tube. Carbon monoxide (CO) was

156 measured using ultra-violet and infrared analysers (PICARRO). The nitric oxide (NO) and
157 nitrogen dioxide (NO₂) measurements were performed using an ozone chemiluminescence
158 instrument (Thermo Environmental Instrument TEi42C with a Blue Light Converter for the NO₂
159 conversion). On-board aerosol instruments sampled ambient air via stainless steel tubing through
160 the Community Aerosol Inlet (CAI). This is an isokinetic and isoaxial inlet with a 50 % sampling
161 efficiency for particles with a diameter of 5 μm (*Denjean et al., 2016*).

162

163 The total number concentration of particles larger than 10 nm (N_{tot}) was measured by a **butanol-**
164 **based conductive cooling type** condensation particle counter (CPC, model MARIE built by
165 University of Mainz; *Mertes, Schröder, and Wiedensohler, 1995; Russell et al., 1996;*
166 *Wiedensohler et al., 1997*). The aerosol size distribution was measured using an ultra-high
167 sensitivity aerosol spectrometer (UHSAS, DMT, **0.04 – 1 μm**), a custom-built scanning mobility
168 sizer spectrometer (SMPS, **20 - 485 nm**) and an optical particle counter (OPC, GRIMM model
169 1.109, **0.3 – 32 μm**). Instrument calibration was performed with PSL nanospheres and oil particles
170 size-selected by a differential mobility analyser (DMA) for diameters from 90 nm to 20 μm. The
171 SMPS data acquisition system failed after two-third of the campaign and could not be repaired. We
172 found the UHSAS to show false counts in the diameters below 100 nm. Therefore, these channels
173 were disregarded in the data analysis.

174

175 The particle extinction coefficient (σ_{ext}) at the wavelength of 530 nm was measured with a cavity
176 attenuated phase shift particle light extinction monitor (CAPS-PMex, Aerodyne Research). The
177 particle scattering coefficients (σ_{scat}) at 450, 550 and 635 nm were measured using a **TSI 3563 3-**
178 **Wavelength Integrating Nephelometer** and corrected for angular truncator error in the data
179 inversion procedure using a Mie code (i.e. Section 2.3.1 and Appendix 2). The absorption
180 coefficients (σ_{abs}) at 467, 520 and 660 nm were measured by a Radiance Research Particle Soot
181 Absorption Photometer (PSAP). The PSAP measures changes of filter attenuation due to the
182 collection of aerosol deposited on the filter, which were corrected for the scattering artifacts
183 according to the *Virkkula (2010)* method. Prior to the campaign, the CAPS-PMex was evaluated
184 against the combination of the **integrating** nephelometer and the PSAP. An instrument
185 intercomparison was performed with purely scattering ammonium sulfate particles and with
186 strongly absorbing black carbon particles (BC). Both types of aerosol were generated by
187 nebulizing a solution of the respective substances and size-selected using a DMA. For instrument
188 intercomparison purposes, σ_{ext} from the combination of **integrating** nephelometer and PSAP was
189 adjusted to that for 530 nm by using the scattering and absorption Ångstrom exponent (SAE and

190 AAE, respectively). The instrument evaluation showed an excellent accuracy of the CAPS-PMex
191 measurements by comparison to the **integrating** nephelometer and PSAP combination. **The results**
192 **were within the ± 3 % uncertainty reported by Massoli et al. (2010) and Petzold et al. (2013) for**
193 **the same instrument configuration.**

194

195 **2.2. Ancillary products**

196 In order to determine the history of air masses prior aircraft sampling, backward trajectories and
197 satellite images were used. The trajectories were computed using the Hybrid Single Particle
198 Lagrangian Integrated Trajectory Model (HYSPLIT) and the National Centers for Environmental
199 Prediction (NCEP) Global Data Assimilation System (GDAS) data with 0.5° horizontal resolution
200 for sequences and times of interest. We compared the backward trajectory heights with information
201 of fire burning times (e.g. MODIS Burnt Area Product) and dust release periods (e.g. Meteosat
202 Second Generation (MSG) dust RGB composite images) to assess the aerosol source regions of the
203 investigated air masses. The air masses represented by the trajectory are assumed to obtain their
204 aerosol loading from source regions when the trajectory passes over regions with significant dust
205 activation and/or fire activity at an altitude close to the surface. Trajectory calculations with
206 slightly modified initial conditions with respect to the arrival time, location and altitude were
207 performed to check the reliability of the location of source regions. Uncertainties in this approach,
208 caused by unresolved vertical mixing processes, and by general uncertainties of the trajectory
209 calculations are estimated to be in the range of 15–20 % of the trajectory distance (*Stohl et al.,*
210 *2002*).

211

212 **2.3. Data analysis**

213 In the following, extensive aerosol parameters (concentrations, scattering, absorption and
214 extinction coefficients) are converted to standard temperature and pressure (STP) using $T = 273$ K
215 and $P = 1013.25$ hPa. The STP concentration data correspond to mixing ratios, which are
216 independent of ambient pressure and temperature during the measurement. In the analysis, the data
217 were averaged over sections of SLR with homogeneous aerosol conditions outside of clouds.

218

219 **2.3.1. Derivation of aerosol microphysical and optical properties**

220 Appendix 2 and Table 1 show the iterative procedure and the equations used to calculate the
221 aerosol microphysical and optical parameters as briefly explained below.

222

223 The particle number concentration in the coarse mode (N_{coarse}) was calculated by integrating the
224 OPC size distributions over the range 1 to 5 μm . The signal to noise ratio of the OPC for particles
225 in this size range was higher than 3, which makes the instrument well suited to quantify variations
226 in N_{coarse} . The number concentration of particles in the fine mode (N_{fine}) was obtained as the
227 difference between total number concentration (N_{tot} particle diameter range above 5 nm) measured
228 by the CPC and N_{coarse} .

229

230 For optical calculations, the 3λ - σ_{abs} from the PSAP were adjusted at the 3 wavelengths measured
231 by the **integrating** nephelometer using the AAE calculated from the 3λ measured σ_{abs} . Once σ_{scat}
232 and σ_{abs} obtained at the same wavelength, an optical closure study estimated the complex refractive
233 index based on optical and size data. Optical calculations were performed using Mie theory,
234 implying a sphericity assumption, because it facilitates a quantitative comparison with past data,
235 mostly using this simplification and because most climate models assume spherical properties. The
236 retrieval algorithm consists of iteratively varying the real part of the complex refractive index (n)
237 from 1.33 to 1.60 and the imaginary part of the complex refractive index (k) from 0.000 to 0.080 in
238 steps of resolution of 0.001. n and k were fixed when the difference between calculated values of
239 σ_{scat} and σ_{abs} and measurements was below 1%. Given that the size distribution measured by the
240 UHSAS and the OPC depends on m , the optical-to-geometrical diameter conversion was
241 recalculated at each iteration based on the assumed m . The resulting number size distributions
242 from SMPS, UHSAS and OPC were parameterized by fitting four log-normal distributions and
243 used as input values in the optical calculations. Once n and k were obtained at 3λ , we estimated the
244 following optical parameters:

245 - SAE depends on the size of the particles. Generally, it is lower than 0 for aerosols dominated by
246 coarse particles, such as dust aerosols, but it is higher than 0 for fine particles, such as
247 anthropogenic pollution or biomass burning aerosol (*Seinfeld and Pandis, 2006; Schuster et al.,*
248 *2006*).

249 - AAE provides information about the chemical composition of atmospheric aerosols. BC absorbs
250 radiation across the whole solar spectrum with the same efficiency, thus it is characterized by AAE
251 values around 1. Conversely, mineral dust particles show strong light absorption in the blue to
252 ultraviolet spectrum leading to AAE values up to 3 (*Kirchstetter et al., 2004; Petzold et al., 2009*).

253 - SSA describes the relative importance of scattering and absorption for radiation. Thus, it indicates
254 the potential of aerosols for cooling or warming the lower troposphere.

255 - g describes the probability of radiation to be scattered in a given direction. Values of g can range
256 from -1 for entirely backscattered light to $+1$ for complete forward scattering light.

257 - *MEE* represents the total light extinction per unit mass concentration of aerosol. The estimates of
258 *MEE* assume mass densities of 2.65 g cm³ for dust aerosol, 1.35 g cm³ for biomass burning
259 aerosol, 1.7 g cm³ for anthropogenic aerosol and 1.49 g cm³ for background aerosol (*Hess et al.*,
260 *1998; Haywood et al., 2003a*).

Aerosol parameters	Symbol	λ (nm)	Method
Aerosol microphysical properties			
Total number concentration	N_{tot}	-	Measured by a CPC in the particle diameter range above 5 nm
Number concentration in the coarse mode	N_{coarse}	-	GRIMM size distributions integrated on the range 1 to 5 μm .
Number concentration in the fine mode	N_{fine}	-	Difference N_{tot} and N_{coarse} .
Number size distribution	$dN/d\log D_p$	-	$dN/d\log D_p = \sum_{i=1}^4 (N_{tot,i} \exp(-(\log D_p - \log D_{p,g,i})^2 / (2 \log \sigma_i)) / (\sqrt{2 \log \sigma_i}))$ with $N_{tot,i}$ the integrated number concentration, $D_{p,g,i}$ the geometric median diameter and σ_i geometric standard deviation for each mode i
Volume size distribution	$dV/d\log D_p$	-	$dV/d\log D_p = \sum_{i=1}^4 (N_{tot,i} D_p^3 \pi/6 \exp(-(\log D_p - \log D_{p,g,i})^2 / (2 \log \sigma_i)) / (\sqrt{2 \log \sigma_i}))$
Aerosol optical properties			
Scattering coefficient	σ_{scat}	450, 550, 635	Measured by the integrating nephelometer and corrected for truncator error
Absorption coefficient	σ_{abs}	467, 520, 660	Measured by the PSAP and corrected for filter based artefacts
Extinction coefficient	σ_{ext}	530	Measured by the CAPS-PMex
Scattering Ångström exponent	SAE	450 to 700	Calculated from the integrating nephelometer measurements : $SAE = -\ln(\sigma_{scat}(450)/\sigma_{scat}(700)) / \ln(450/700)$
Absorption Ångström exponent	AAE	440 to 660	Calculated from the PSAP measurements : $AAE = -\ln(\sigma_{abs}(467)/\sigma_{abs}(660)) / \ln(467/660)$
Complex refractive index	n	450, 550, 660	Inversion closure study using Mie theory (Fig. A2) $m(\lambda) = n(\lambda) - ik(\lambda)$
Single scattering albedo	SSA	450, 550, 660	Inversion closure study using Mie theory (Fig. A2) $SSA(\lambda) = \sigma_{scat}(\lambda) / \sigma_{ext}(\lambda)$
Mass extinction efficiency	MEE	450, 550, 660	Inversion closure study using Mie theory (Fig. A2) $MEE(\lambda) = \sigma_{ext}(\lambda) / C_m$ with C_m the aerosol mass concentration
Asymmetry parameter	g	450, 550, 660	Inversion closure study using Mie theory (Fig. A2) $g(\lambda) = 1/2 \int_0^\pi \cos(\Theta) \sin(\Theta) P(\Theta, \lambda) d(\Theta)$ with $P(\Theta, \lambda)$ the scattering phase function and Θ the scattering angle.

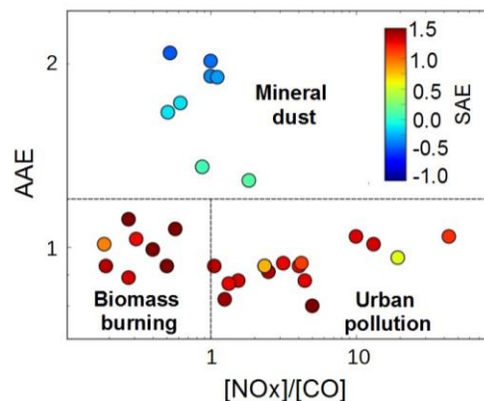
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Table 1. Aerosol microphysical and optical properties derived in this work

263

264 2.3.2. Classification of aerosols plumes

265 Data were screened in order to isolate plumes dominated by anthropogenic pollution from urban
266 emissions, biomass burning and mineral dust particles, resulting in a total number of 19, 12 and 8
267 genuine plume interceptions, respectively, across the 15 flights. As shown in Figure 2,
268 identification of the plumes was based on a combination of CO and NO_x (sum of NO and NO_2)
269 concentrations, as well as AAE and SAE that have been shown to be good parameters for
270 classifying aerosol types (Kirchstetter et al., 2004; Petzold et al., 2009). The classification was
271 then compared with results from the back trajectory analysis (Figure 3) and satellite images
272 described in section 2.2.



273

274 **Figure 2. Absorption Ångström Exponent (AAE) as a function of the ratio NO_x to CO. The**
275 **markers are colored according to the Scattering Ångström Exponent (SAE). Classification of**
276 **mineral dust, biomass burning and urban pollution particles has been added to the figure.**
277

278 The guidelines for classification are as follows:

279 - *Anthropogenic pollution*: SAE was beyond threshold 0, indicating a large number fraction of
280 small particles in urban plumes, and CO and NO_x concentrations 2 times higher than the
281 background concentrations. During the DACCIWA campaign background CO and NO_x values
282 were around 180 ppb and 0.28 ppb, respectively. The trajectories show large differences in the
283 flow patterns and source regions with urban plumes originating from the polluted cities of Lomé,
284 Accra and Abidjan. The aircraft sampling over land mostly followed the north-eastward direction
285 (Figure 3d).

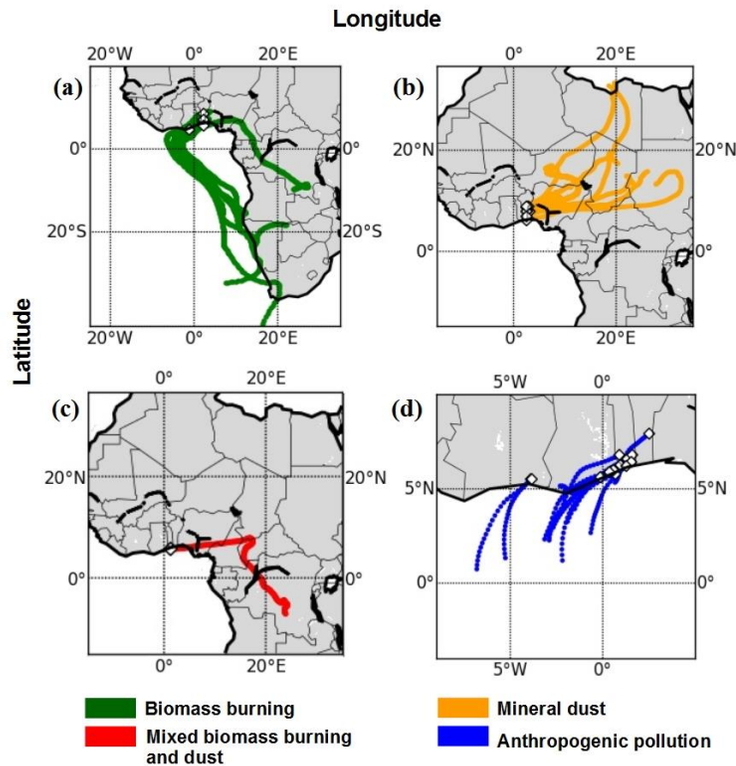
286 - *Biomass burning*: The criteria are the same as for urban pollution plumes except that trajectories
287 track these plumes back to active fire hotspots as observed by MODIS and the ratio NO_x to CO
288 was set below 1. CO and NO_x are byproducts of combustion sources but CO is preserved longer
289 along the plume when compared with NO_x , which makes the ratio NO_x to CO a good indicator for
290 distinguishing fresh anthropogenic pollution plumes from biomass burning plumes transported

291 over long distances (Wang *et al.*, 2002; Silva *et al.*, 2017). During this time of the year, most of the
292 forest and grassland fires were located in Central and Southern Africa (Figure 3a).

293 - *Mineral dust*: AAE higher than 1 indicates a large mass fraction of mineral dust and a SAE below
294 0 indicate a high effective particle diameter. The source region of the dust loaded air masses was
295 located in the Saharan desert and in the Sahel (Figure 3b).

296 - *Dust and biomass burning mixing*: Combining remote sensing observations and model
297 simulations, Flamant *et al.* (2018a) identified a biomass burning plume mixed with mineral dust.
298 This agrees well with the measured AAE of 1.2 and SAE of 0.3 observed in this layer. Menut *et al.*
299 (2018) have shown that one of the transport pathways of biomass burning aerosols from Central
300 Africa was associated with northward advection towards Chad and then westward displacement
301 linked to the African Easterly Jet. The plume originated from a broad active biomass burning area
302 including Gabon, the Republic of Congo and the Democratic Republic of Congo and passed over
303 areas with strong dust emissions further north within 1–3 days before being sampled by the aircraft
304 (Figure 3c).

305 - *Background*: We refer to background conditions as an atmospheric state in the boundary layer
306 without the detectable influence of mineral dust, biomass burning or local anthropogenic sources.
307 Most back trajectories originated from the marine atmosphere and coastal areas south of the
308 sampling area.

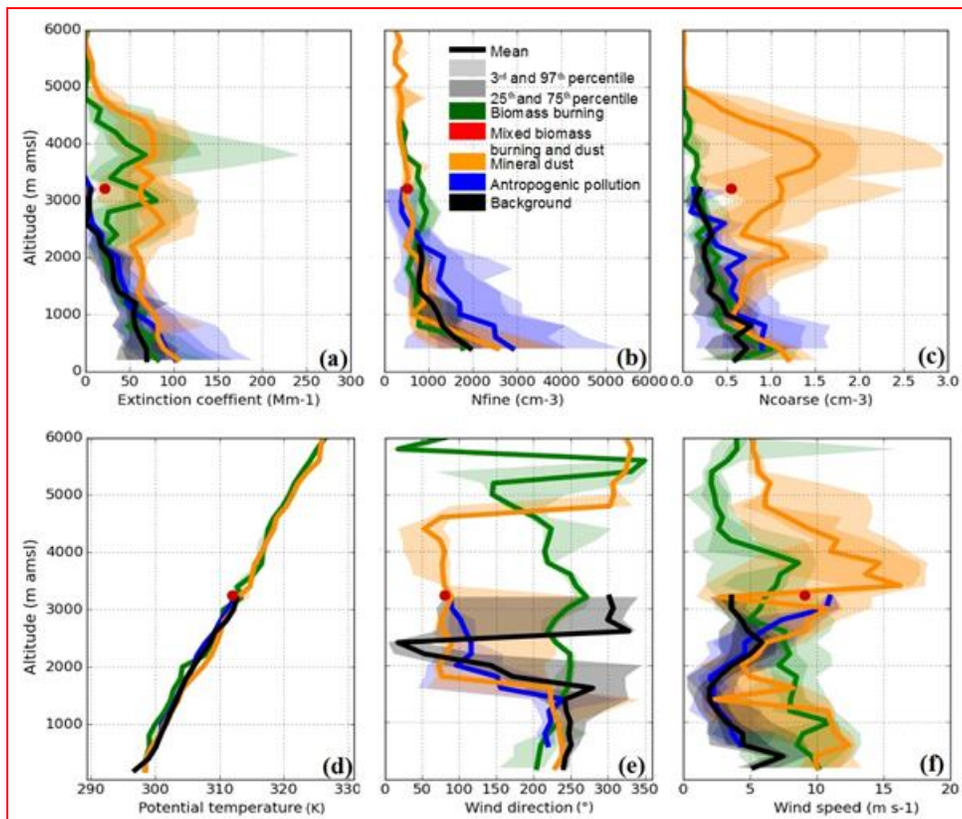


309 **Figure 3. Backward trajectories for the analyzed aerosol layers. Trajectories date back 10**
 310 **days for (a), 5 days for (b) and (c), and 1 day for (d).**

311

312 3.1. Aerosol vertical distribution

313 Figure 4 shows a statistical analysis of N_{fine} , N_{coarse} and σ_{ext} derived from the *in-situ* measurements
 314 of vertical profiles. The aerosol vertical structure is strongly related to the meteorological structure
 315 of the atmosphere (see Knippertz *et al.*, 2017 for an overview of the DACCIWA field campaign).
 316 Therefore wind vector and potential temperature profiles acquired with the aircraft have been
 317 added to Figure 4 as a function of the dominating aerosol composition, introduced in Figure 1. The
 318 data from individual vertical profiles were merged into 200 m vertical bins from the surface to 6
 319 km amsl. The profiles were calculated using only individual profiles obtained outside of clouds.



320

321 **Figure 4. Vertical layering of aerosols and meteorological variables for profiles for which**
 322 **aerosols dominated by biomass burning (green), dust (orange), mixed dust-biomass burning**
 323 **(red), anthropogenic pollution (blue) and background particles (black) were detected. The**
 324 **panels show profiles of (a) the extinction coefficient at 530 nm, (b) the particle number**
 325 **concentration in the range $0.005 < D_p < 1 \mu\text{m}$, (c) the particle number concentration in the**
 326 **range $1 < D_p < 5 \mu\text{m}$, (d) potential temperature, (e) the wind direction and (f) the wind speed.**
 327 **The colored areas represent the 3th, 25th, 75th and 97th percentiles of the data. The mixed dust-**
 328 **biomass burning plume is represented by a dot because it is derived from measurements**
 329 **during a SLR.**

330

331 The observed wind profiles highlight the presence of several distinct layers in the lower
332 troposphere. For cases related to dust, urban pollution and background condition, we clearly
333 observe the monsoon layer up to 1.5 km amsl which is characterized by weak to moderate wind
334 speeds (2 to 10 m s⁻¹, the later corresponding to dust cases) and a flow from the southwest (220-
335 250°). In all three air mass regimes, the monsoon layer is topped by a 500 to 700 m deep layer
336 characterised by a sharp wind direction change (from south-westerly to easterly). Weak wind
337 speeds (less than 5 m s⁻¹) are observed in urban pollution and background conditions, while higher
338 wind speeds are observed in the dust cases. Above 2.5 km amsl, the wind speed increases in the
339 urban pollution and dust cases and the wind remains easterly, indicating the presence of the
340 African easterly jet with its core typically farther north over the Sahel (Figure 8 in *Knippertz et al.*,
341 2017 for the latitudinal variations of the African easterly jet during the DACCIWA field phase).
342 The maximum easterlies are observed in the dust cases slightly below 3.5 km amsl (> 15 m s⁻¹).
343 For the background cases, the wind above the shear layer shifts to north-westerly and remains
344 weak (~i.e. 5 m s⁻¹). Overall, the wind profile associated with the biomass burning cases is quite
345 different from the other three cases, with a flow essentially from the south-southwest below 5 km
346 amsl and higher wind speeds in the lower 2 km amsl than above, and a secondary maximum of 7
347 m s⁻¹ at 4 km amsl.

348

349 The vertical distribution of aerosol particles was very inhomogeneous, both across separate
350 research flights and between individual plumes encountered during different periods of the same
351 flight. Measurements of aerosols within this analysis cover a broad geographic region, as shown in
352 Figure 1, which may explain some of the variability. SWA is subject to numerous anthropogenic
353 emission sources (e.g. road traffic, heavy industries, open agriculture fires, etc.) coupled to
354 biogenic emissions from the ocean and forests. These resulting large emissions are reflected in the
355 high variability of σ_{ext} , N_{fine} and N_{coarse} in the lower troposphere over SWA. Below 2.5 km amsl, σ_{ext}
356 showed a large heterogeneity with values ranging from 35 to 188 Mm⁻¹ between the 3rd and 97th
357 percentile and a median value of 55 Mm⁻¹. The variability of σ_{ext} values was slightly enhanced near
358 the surface and was correlated to N_{fine} and N_{coarse} which ranged from 443 to 5250 cm⁻³ and from
359 0.15 to 1.6 cm⁻³, respectively. Maximum surface σ_{ext} was recorded in the anthropogenic pollution
360 plume of Accra where high N_{fine} was sampled. The aerosol vertical profile is strongly modified
361 during biomass burning and dust events. The dust plume extends from 2 to 5 km amsl, and is
362 associated with transport from the dust sources in Chad and Sudan (see Figure 3) with the midlevel
363 easterly flow. The biomass burning plume extends from 1.5 to 5 km amsl and is associated with
364 transport from the southwest in a layer of enhanced wind speed just below 4 km amsl as discussed

365 above. Both layers showed enhanced σ_{ext} with median values of 68 Mm^{-1} ($p_{03} = 12 \text{ Mm}^{-1}$; $p_{97} = 243$
366 Mm^{-1}) in biomass burning plumes and 78 Mm^{-1} ($p_{03} = 45 \text{ Mm}^{-1}$; $p_{97} = 109 \text{ Mm}^{-1}$) in dust plumes.
367 As expected, the extinction profile was strongly correlated to N_{fine} for biomass burning layers and
368 N_{coarse} for dust layers.

369

370 A prominent feature in the vertical profiles is the presence of fine particles up to 2.5 km amsl
371 outside of biomass burning or dust events. σ_{ext} , N_{fine} and N_{coarse} continuously decrease with altitude,
372 most likely due to vertical mixing of local emissions from the surface to higher levels. Therefore,
373 the regional transport of locally emitted aerosols was not limited to the surface but occurred also at
374 higher altitude. Recently, numerical tracer experiments performed for the DACCIWA airborne
375 campaign period have demonstrated that a combination of land–sea surface temperature gradients,
376 orography-forced circulation and the diurnal cycle of the wind along the coastline favor the
377 vertical dispersion of pollutants above the boundary layer during daytime (*Deroubaix et al., 2019*;
378 *Flamant et al., 2018a*). Because of these complex atmospheric dynamics, aerosol layers transiting
379 over the Gulf of Guinea in the free troposphere could be contaminated by background or urban
380 pollution aerosols from the major coastal cities.

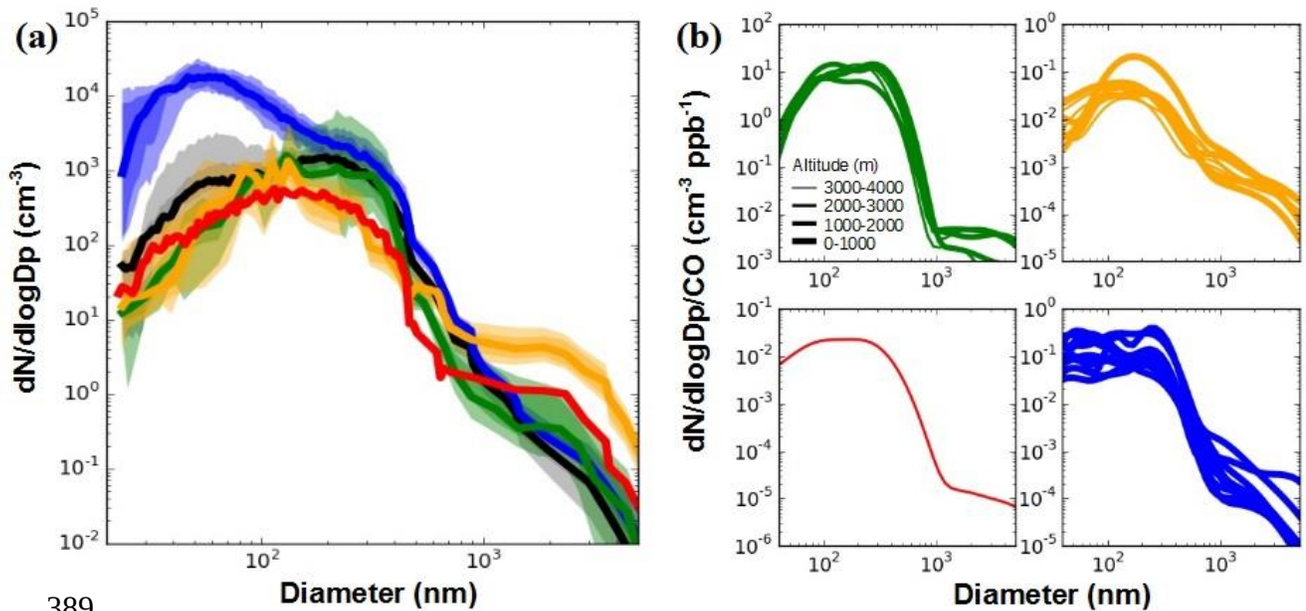
381

382 **3.2. Aerosol size distribution**

383 Figure 5a shows the range of variability of the number and volume size distributions measured
384 during DACCIWA. These are extracted from the SLRs identified in Fig.1. Figure 5b shows the
385 same composite distribution normalized by CO concentration in order to account for differences in
386 the amount of emissions from combustion sources.

387

388



389
 390 **Figure 5. (a) Statistical analysis of number size distributions with colored areas representing**
 391 **the 3th, 25th, 75th and 97th percentiles of the data and (b) number size distributions normalized**
 392 **to CO for plumes dominated by biomass burning (green), dust (orange), mixed dust-biomass**
 393 **burning (red), anthropogenic pollution (blue) and background particles (black). In panel b,**
 394 **the line thickness is scaled by the altitude of the aerosol plume.**
 395

396 Considerable variability in the number concentration of the size distributions, up to approximately
 397 2 orders of magnitude, was observed for a large fraction of the measured size range. The size
 398 distributions varied both for different aerosol types and for a given aerosol class. This reflects the
 399 relative wide range of different conditions that were observed over the region, both in terms of
 400 sources, aerosol loading, and lifetimes of plumes.

401

402 In particular for ultrafine particles with diameters below 100 nm, large differences were observed,
 403 with an increase as large as a factor of 50 in urban plumes, which reflects concentration increase
 404 from freshly formed particles. Interestingly elevated number concentrations of these small-
 405 diameter particles were also observed in some dust layers. Comparing the particle size distribution
 406 of the different dust plumes sampled during the field campaign, a variation as large as a factor of
 407 20 in the number concentration of ultrafine particles is found (i.e. Figure 4). Their contribution
 408 decreased with height as reflected by higher small particle number recorded in dust plumes below
 409 2.5 km amsl (Figure 4b and 5b). As the composite urban size distributions showed a relatively
 410 similar ultrafine mode centered at 50 nm, dust layers have most likely significant contributions
 411 from anthropogenic pollution aerosol freshly emitted in SWA. The ultrafine mode was not

412 observed in biomass burning size distributions, even though dust and biomass burning plumes
413 were sampled in the same altitude range. We interpret this observation with dust plumes
414 transported below 2.5 km amsl that were sampled over the region of Savè (8°01'N, 2°29'E; Benin)
415 near the identified urban air mass transported northeastwards from Lomé and/or Accra and which
416 may have collected significant fresh pollution on their way, whereas biomass burning plumes
417 collected at the same altitude and sampled over Ivory Coast south of the Abidjan pollution plumes
418 may not have been affected by significant direct pollution (Figure 1).

419

420 The accumulation mode was dominated by two modes centered at $D_{p,g} \sim 100$ and 230 nm
421 depending on the aerosol plume. The particle size distributions for biomass burning plumes were
422 generally dominated by an accumulation mode centered at $D_{p,g} \sim 230$ nm. Despite the relative wide
423 range of sources and lifetimes of the biomass burning plumes sampled throughout the campaign
424 (Figure 3), the $D_{p,g}$ in the accumulation mode showed little variation ($D_{p,g}$ from 210 to 270 nm)
425 between the plumes. Similarly, previous field studies found accumulation mode mean diameters
426 from 175 to 300 nm for aged biomass burning plumes, regardless of their age, transport time and
427 source location (Capes *et al.*, 2008; Janhäll *et al.*, 2010; Weinzierl *et al.*, 2011; Sakamoto *et al.*,
428 2015; Carrico *et al.* 2016). The coagulation rate can be very high in biomass burning plumes and
429 can shape the size distribution over a few hours (Sakamoto *et al.*, 2016). It is worth noting that in
430 the biomass burning and dust size distributions there is a persistent particle accumulation mode
431 centered at ~ 100 nm that exceeds the amount of particles centered at 230 nm in some layers. This
432 small mode is unlikely to be related to long-range transport of biomass burning and Saharan dust
433 emissions, as it would be expected that particles in this size range would grow to larger particles
434 through coagulation relatively quickly. As similar concentrated accumulation modes of particles
435 have been observed in background plumes, it suggests the entrainment of background air from the
436 boundary layer in dust and biomass burning plumes.

437

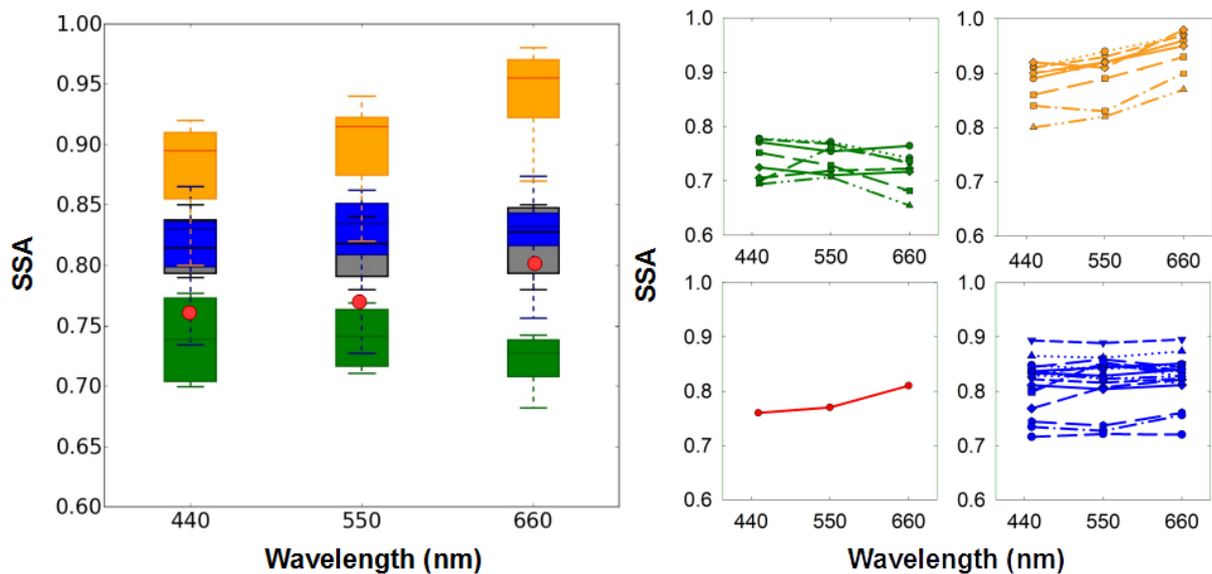
438 The number concentration of large super-micron particles was strongly enhanced in the mineral
439 dust layers. The peak number concentration displayed a broad shape at $D_{p,g} \sim 1.8 \mu\text{m}$, which is
440 comparable to literature values of other long-range transported dust aerosols (Formenti *et al.*,
441 2011a; Weinzierl *et al.*, 2011; Ryder *et al.*, 2013; Denjean *et al.*, 2016; Liu *et al.*, 2018). The super-
442 micron mode of the dust plume is expected to be impacted by the mixing with other particles in
443 case of an internal mixing, which should somewhat increase the particle size. The relatively
444 homogeneous $D_{p,g}$ in the coarse mode of dust reported here ($D_{p,g}$ from 1.7 to 2.0 μm) suggests low
445 internal mixing with other atmospheric species. Besides, the volume size distribution in urban

446 plumes showed significant presence ($\sim 65\%$ of the total aerosol volume) of large particles with
 447 diameters of $\sim 1.5 - 2 \mu\text{m}$, which were also observed in background conditions. We measured
 448 *AAE* in the range 0.7-1 in anthropogenic pollution plumes (Figure 2), which suggests negligible
 449 contribution of mineral dust in these plumes. This coarse mode has most likely significant
 450 contributions from sea salt particles, as plumes arriving from the cities were transported at low
 451 altitude over the ocean (Fig. 3).

452

453 3.3. Aerosol optical properties

454 *SSA* is one of the most relevant intensive optical properties because it describes the relative
 455 strength of the aerosol scattering and absorption capacity and is a key input parameter in climate
 456 models (Solmon *et al.*, 2008). Figure 6 shows the spectral *SSA* for the different SLRs considered in
 457 this study.



458

459 **Figure 6. (a) Statistical analysis of single scattering albedo at 450, 550 and 660 nm for**
 460 **plumes dominated by biomass burning (green), dust (orange), mixed dust-biomass burning**
 461 **(red), anthropogenic pollution (blue) and background particles (black). The boxes enclose the**
 462 **25th and 75th percentiles, the whiskers represent the 5th and 95th percentiles and the**
 463 **horizontal bar represents the median. (b) Spectral SSA for the different individual plumes**
 464 **considered in this study. The mixed dust-biomass burning plume is represented by a dot**
 465 **because it is derived from measurements during only one SLR.**
 466

467 The highest absorption (lowest *SSA*) at all three wavelengths was observed for biomass burning
 468 aerosols. *SSA* values ranged from 0.69–0.78 at 440 nm, 0.71–0.77 at 550 nm and 0.65–0.76 at 660
 469 nm. This is on the low side of the range of values (0.73–0.93 at 550 nm) reported over West Africa

470 during DABEX for biomass burning plumes mixed with variable proportion of mineral dust
471 (*Johnson et al., 2008*). No clear tendency was found for the spectral dependence of *SSA*, which in
472 some of the cases decreased with wavelength and in others were very similar to each other at all
473 three wavelengths.

474

475 *SSA* values of anthropogenic pollution aerosols were generally intermediate in magnitude with
476 median values of 0.81 at 440 nm, 0.82 at 550 nm and 0.82 at 660 nm. Our data show that the value
477 of *SSA* varied significantly for the different plumes. Some pollution aerosols absorb almost as
478 strongly as biomass burning aerosols with *SSA(550nm)* values as low as 0.72, whereas the highest
479 *SSA(550nm)* value observed was 0.86. In addition, the absorption properties of urban aerosol
480 varied greatly between the sampled plumes for smoke of apparent same geographic origin. For
481 example, we measured *SSA(550nm)* values from 0.72 to 0.82 in the Accra pollution outflow. The
482 variability in *SSA* values may be due to the possible contribution of emissions from different cities
483 to the sampled pollution plumes (*Deroubaix et al., 2019*), thus having different combustion
484 sources and chemical ages. ~~Past in-situ measurements of aerosol optical properties over SWA cities~~
485 ~~appear, unfortunately, to be absent from literature. However, t~~The flat spectral dependence of *SSA*
486 appears to be anomalous for anthropogenic pollution aerosols, as *SSA* has been shown to decrease
487 with increasing wavelength for a range of different urban pollution plumes ~~over European,~~
488 ~~American and Asian cities~~ (*Dubovick et al., 2002; Petzold et al., 2011; Di Biagio et al., 2016; Shin*
489 *et al., 2019*).

490

491 The magnitude of *SSA* increased at the three wavelengths when dust events occurred. Large
492 variations in *SSA* were obtained with values ranging from 0.76–0.92 at 440 nm, 0.81–0.94 at 550
493 nm and 0.81–0.97 at 660 nm. The measurement of *SSA* is highly dependent on the extent to which
494 the coarse mode is measured behind the aerosol sampling inlet. *Denjean et al. (2016)* found that
495 the absolute error associated with *SSA*, *g* and *MEE* of dust aerosols due to the CAI inlet is in the
496 range covered by the measurement uncertainties. However, different aerosol inlet systems were
497 used during previous field campaigns, which makes comparison of our results with previous
498 measurements difficult. Overall, compared with the literature for transported dust, lower values
499 were obtained in the present study for few cases. For example, *Chen et al. (2011)* reported
500 *SSA(550 nm)* values of 0.97 ± 0.02 during NAMMA (a part of AMMA operated by NASA) using
501 an inlet with a comparable sampling efficiency. The lower values from DACCIWA reflect
502 inherently more absorbing aerosols in some dust plumes. In contrast to fire plumes, the *SSA* of
503 dust aerosol showed a clear increasing trend with wavelength. This behavior is likely due to the

504 domination of large particles in dust aerosol, which is in agreement to similar patterns observed in
 505 dust source regions (*Dubovik et al., 2002*). Moreover, an increase of SSA is observed with
 506 wavelength for mixed dust-smoke aerosol, suggesting that the aerosol particles were
 507 predominantly from dust, albeit mixed with a significant loading of biomass burning.
 508

		<i>SSA(450)</i>	<i>SSA(550)</i>	<i>SSA(660)</i>	<i>MEE(450)</i>	<i>MEE(550)</i>	<i>MEE(660)</i>	<i>g(450)</i>	<i>g(550)</i>	<i>g(660)</i>	<i>SAE</i>
Mineral dust	median	0.88	0.90	0.93	0.74	0.68	0.66	0.74	0.72	0.69	-0.35
	3 th	0.82	0.82	0.86	0.38	0.38	0.39	0.69	0.67	0.65	-0.56
	25 th	0.85	0.87	0.90	0.43	0.43	0.43	0.73	0.72	0.67	-0.48
	75 th	0.91	0.93	0.96	0.94	0.85	0.85	0.75	0.74	0.72	-0.25
	97 th	0.92	0.95	0.97	1.57	1.37	1.21	0.78	0.76	0.72	-0.12
Biomass burning	median	0.74	0.76	0.72	1.91	1.62	1.34	0.69	0.68	0.61	1.07
	3 th	0.70	0.72	0.66	0.94	1.45	1.22	0.64	0.65	0.59	0.59
	25 th	0.70	0.76	0.71	1.67	1.48	1.27	0.69	0.65	0.60	0.83
	75 th	0.77	0.77	0.74	1.86	1.65	1.55	0.72	0.68	0.62	1.15
	97 th	0.78	0.77	0.76	2.38	1.92	1.58	0.73	0.68	0.63	1.64
Mixed dust- Biomass burning	median	0.76	0.77	0.81	1.58	1.40	1.30	0.73	0.66	0.64	0.38
	3 th	-	-	-	-	-	-	-	-	-	-
	25 th	-	-	-	-	-	-	-	-	-	-
	75 th	-	-	-	-	-	-	-	-	-	-
	97 th	-	-	-	-	-	-	-	-	-	-
Anthropogenic Pollution	median	0.83	0.84	0.85	2.60	2.49	1.90	0.60	0.61	0.62	0.75
	3 th	0.78	0.79	0.81	0.70	1.24	0.54	0.60	0.59	0.54	0.30
	25 th	0.80	0.82	0.83	2.14	2.25	1.53	0.62	0.60	0.56	0.65
	75 th	0.84	0.86	0.85	3.51	2.96	2.53	0.69	0.62	0.67	0.89
	97 th	0.87	0.88	0.90	3.70	4.83	2.74	0.73	0.64	0.70	0.94

509

510 **Table 2. Single scattering albedo, mass extinction efficiency (in $\text{m}^2 \text{g}^{-1}$), asymmetry parameter**
 511 **and scattering Ångstrom exponent for the dominant aerosol classification.**

512

513 As shown in Table 2, the observed variability of SSA reflects a large variability for *MEE* at 550nm,
 514 which spans a wide range from 0.38 to 1.37 $\text{m}^2 \text{g}^{-1}$, 1.45 to 1.92 $\text{m}^2 \text{g}^{-1}$ and 1.24 to 4.83 $\text{m}^2 \text{g}^{-1}$ for
 515 dust, biomass burning for anthropogenic polluted aerosols, respectively. *MEE* is heavily influenced
 516 by the mass concentrations in the accumulation mode where the aerosol is optically more efficient
 517 in extinguishing radiation. We found *MEE* to be positively correlated with *SAE* (not shown), which
 518 was expected because of the dependence of *MEE* on particle size. In contrast, the values of *g*
 519 appear to differ only little between the sampled plumes for a given aerosol class. We found *g* in the
 520 range of 0.67–0.76 for dust, 0.65–0.68 for biomass burning and 0.59–0.64 for anthropogenic
 521 polluted aerosols at 550 nm. *g* values in dust plumes were high, which is expected due to the
 522 presence of coarse particles contributing to forward scattering.

523

524 This analysis includes sampled aerosols originating from different source regions and having
525 undergone different aging and mixing processes, which could explain some of the variability. The
526 impact of these factors on the magnitude and spectral dependence of optical parameters will be
527 investigated in the following section.

528

529 **4. Discussion**

530 **4.1. Contribution of local anthropogenic pollution on aerosol absorption properties**

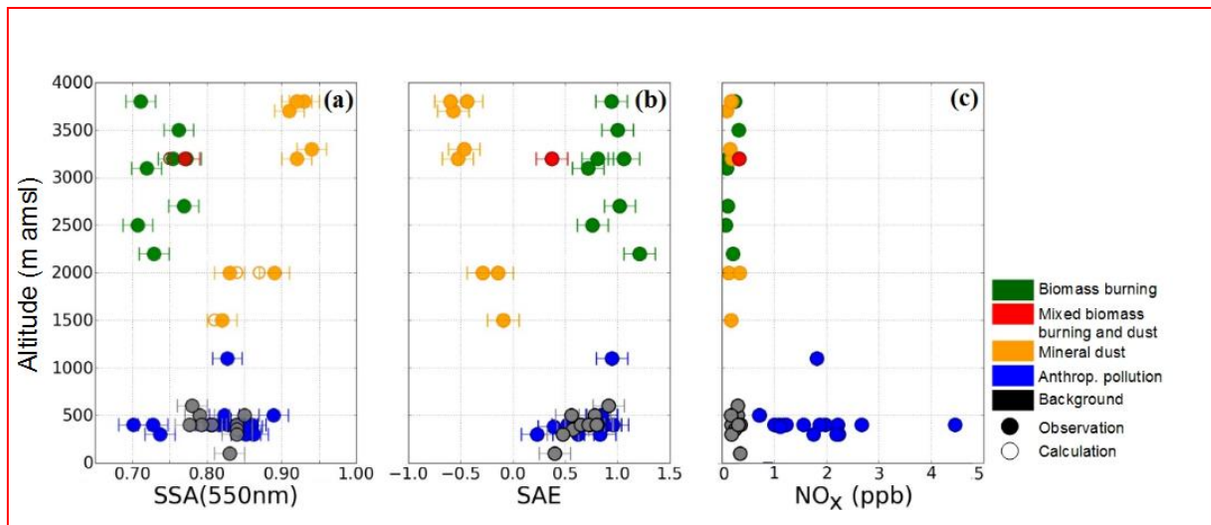
531 Figure 7 shows the vertical distribution of *SSA*, *SAE* and NO_x mixing ratio for the dominant
532 aerosol classification. We exclusively consider measurements acquired during SLRs, since only
533 during these phases the whole set of aerosol optical properties were measured. In dust plumes, if
534 we exclude the case of mixing with biomass burning aerosol, *SSAs* were fairly constant above 2.5
535 km amsl with values ranging between 0.90 and 0.93 at 550 nm, in agreement with values reported
536 over dust source regions (*Schladitz et al., 2009; Formenti et al., 2011b; Ryder et al., 2013, 2018*).
537 Despite the range of sources identified during DACCIWA, dust absorption properties do not seem
538 to be clearly linked to particle origin or time of transport. Aerosols were more absorbing within the
539 low-altitude dust plumes with *SSA* values dropping to 0.81. *SAE* values exhibited simultaneously a
540 sharp increase close to zero below 2.5 km amsl. This is consistent with a higher concentration of
541 fine particles, though the value of *SAE* was still much lower than for pollution or background
542 aerosol (i.e. where it is typically > 0.2), which means that scattering was still dominated by larger
543 particles. Based on **the whole sets of** observations, the strong variation in the light-absorption
544 properties of dust-dominated aerosol over SWA could be attributed to the degree of mixing into the
545 vertical column with either freshly emitted aerosols from urban/industrial sources or long-range
546 transported biomass burning aerosol.

547

548 One of the critical factors in the calculation of aerosol direct and semi-direct radiative effects is the
549 mixing state of the aerosols, which can significantly affect absorbing properties. There were no
550 direct observational constraints available on this property during the DACCIWA airborne
551 campaign. However, we investigated the probable aerosol mixing state by calculating composite
552 *SSA* from the aerosol size distribution. On the basis of Figure 5, dust size distributions showed
553 only minor discrepancies in the mean and standard deviation of the coarse mode but significant
554 differences in the balance between fine and coarse modes, which suggests low internal mixing of
555 dust with other atmospheric species. The size distributions of mixed dust-pollution have been
556 deconvoluted by weighting the size distributions of mineral dust and anthropogenic pollution

557 aerosol averaged over the respective flights. This assumes that dust was externally mixed with the
 558 anthropogenic pollution particles and assumes a homogeneous size distribution for the dust and
 559 anthropogenic pollution aerosol throughout a flight. σ_{scat} and σ_{abs} were then calculated using Mie
 560 theory from each composite size distributions and the corresponding k and m . The refractive
 561 indices at 550 nm were assumed to be 1.52-0.002i and 1.60-0.040i for dust and anthropogenic
 562 pollution particles, respectively, which are the mean values deduced from the data inversion
 563 procedure (i.e. section 2.3.1) throughout the campaign. The resulting σ_{scat} and σ_{abs} were used to
 564 calculate a composite SSA. A similar calculation was performed for the mixed dust-biomass
 565 burning case. Figure 7 shows a good agreement with the observations of SSA, implying that
 566 external mixing appears to be a reasonable assumption to compute aerosol direct and semi-direct
 567 radiative effects in these dust layers for modeling applications. This is consistent with the filter
 568 analysis performed during AMMA and SAMUM-2, which did not reveal any evidence of internal
 569 mixing in both mixed dust-biomass burning and dust-anthropogenic pollution layers (*Chou et al.*,
 570 2008; *Lieke et al.*, 2011; *Petzold et al.*, 2011).

571



572 **Figure 7. Vertical distribution of (a) the single scattering albedo at 550 nm, (b) the scattering**
 573 **Ångström exponent and (c) NO_x mixing ratio for the dominant aerosol classification. In panel**
 574 **(a), full circles represent SSA measurements and empty circles represent composite SSA**
 575 **calculated by deconvoluting size distribution measurements in mixed dust layers and**
 576 **assuming an external mixing state.**

578
 579 SSA, SAE and NO_x of biomass burning plumes did not significantly vary with height from 2.2 to
 580 3.8 km amsl. Moreover, the size distribution of biomass burning aerosols for the observed cases
 581 did not show significant contribution of ultrafine particles (Figure 5). These observations seem to
 582 indicate that the absorption properties of biomass burning plumes were not affected by direct
 583 pollution emissions/

584

585 In the boundary layer, the similar *SSA* and *SAE* in anthropogenic pollution and background plumes
586 suggests that background aerosol may be rather called background pollution originating from a
587 regional background source in the far field. Our analysis of the spectral dependence of *SSA*
588 showed no apparent signature of anthropogenic pollution aerosols (see section 3.3) despite a strong
589 increase of aerosol number concentrations in air masses crossing urban centers (see section 3.2).
590 This can be explained by two factors: First, the majority of accumulation mode particles were
591 present in the background, while the large proportion of aerosols emitted from cities resided in the
592 ultrafine mode particles that have less scattering efficiencies (Figure 5). Second, large amounts of
593 absorbing aerosols in the background can minimize the impact of further increase of absorbing
594 particles to the aerosol load. The high CO values (~180 ppb) observed in background conditions
595 further indicates a strong contribution of combustion emissions at the surface. Recent studies
596 showed a large background of biomass burning transported from the Southern Hemisphere in SWA
597 that dominated the aerosol chemical composition in the boundary layer (*Menut et al., 2018*;
598 *Haslett et al., 2019*). The high absorbing properties (*SSA*~0.81 at 550nm) and the presence of
599 particles both in the accumulation and super-micron modes (i.e. section 3.2.) in background
600 plumes are consistent with being a mixture of aged absorbing biomass burning and Atlantic marine
601 aerosol. These results highlight that aerosol optical properties at the surface were dominated by the
602 widespread biomass burning particles at regional scale.

603

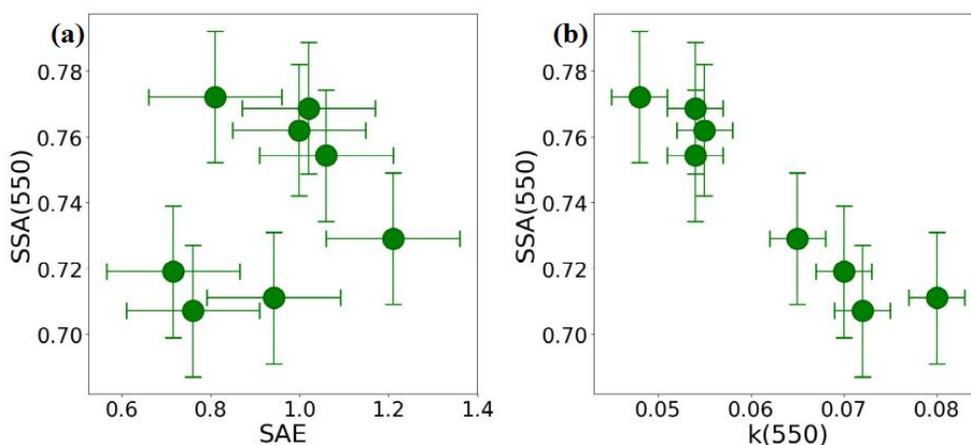
604 **4.2. Aging as a driver for absorption enhancement of biomass burning aerosol**

605 The optical properties of aerosols are determined by either the aerosol chemical composition, the
606 aerosol size distribution, or both. Changes in the size distribution of biomass burning aerosol due
607 to coagulation and condensation have been shown to alter the *SSA*, as particles increase towards
608 sizes for which scattering is more efficient (*Laing et al., 2016*). Variations in particle chemical
609 composition, caused by source emissions and aging processes associated with gas-to-particle
610 transformation and internal mixing, has been shown to change the *SSA* (*Abel et al., 2003*; *Petzold*
611 *et al., 2011*).

612

613 In order to determine the contributions from size distribution and chemical composition to the
614 variation of *SSA* in biomass burning plumes, *SSA* is presented as a function of *SAE* and *k* in Figure
615 7a and b, respectively. *k* was iteratively varied to reproduce the experimental scattering and
616 absorption coefficients, as described in section 2.3.1. It appears that the variation of the size
617 distribution (assessed via *SAE* in Figure 8a) had minimal impact in determining the variability of

618 SSA. Thus, the observations suggest that there was no effect of plume age on the size distribution,
 619 consistent with previous observations of size distribution in aged North American biomass
 620 burning plumes (Sakamoto *et al.*, 2015; Carrico *et al.*, 2016; Laing *et al.*, 2016). Using a
 621 Lagrangian microphysical model, Sakamoto *et al.* (2015) have shown a rapid shift to larger sizes
 622 for biomass burning plumes within the first hours of aging. Less drastic but similarly rapid growth
 623 by coagulation was seen by Capes *et al.* (2008) in their box model. Given that the biomass burning
 624 plumes sampled during DACCIWA had more than 5 days in age, the quick size-distribution
 625 evolution within the early plume stages might explain the limited impact of the size distribution on
 626 the SSA.



627
 628 **Figure 8. Contribution to single scattering albedo (a) from particle size (assessed via SAE)**
 629 **and (b) from composition (assessed via k) in biomass burning plumes.**
 630

631 In contrast, Figure 8b shows that there was a consistent decrease in SSA with increasing k ,
 632 although there is some variability between the results from different plumes. The observed
 633 variability of SSA is reflected in a large variability of k , which is estimated to span the large range
 634 0.048–0.080 at 550 nm. k depends both on the aerosol chemical composition and size distribution
 635 (Mita and Isono, 1980). Given that SSA was found to be independent of the aerosol size
 636 distribution (Figure 8a), Figure 8b suggests that SSA variability was strongly influenced by the
 637 variability in composition of biomass burning aerosol, implying a high contribution for light-
 638 absorbing particles. No clear tendency was found for the wavelength dependence of k , which in
 639 some of the cases increases with wavelength and in others decreases (not shown). **In field**
 640 **observation, significant absorption and strong spectral dependence (values of AAE >1.7) in**
 641 **biomass burning plumes have been frequently attributed to the presence of brown carbon (BrC)**

642 [Kirchstetter et al., 2004; Sandradewi et al., 2008; Romonosky et al., 2019; Chakrabarty et al.,
643 2010; Pokhrel et al., 2016]. Contrary to the current understanding, our measurements show that the
644 contribution of BrC to light absorption is negligible as AAE values ranged from 0.9 to 1.1 with a
645 median value of 1.0. (Figure 2B). Theoretically, fine-mode aerosol with absorption determined
646 exclusively by BC would have AAE equal to 1.0, since BC is expected to have a spectrally
647 constant k (Bond et al., 2013). Therefore, the low SSA values observed in biomass burning plumes
648 over SWA and the small spectral variation of k both suggest that BC is the dominant absorber in
649 the visible and near-IR wavelengths for these biomass burning aerosols.

650

651 Compared with past in-situ measurements of aged biomass burning aerosol, SSA values over SWA
652 (0.71–0.77 at 550 nm) are at the lower end of those reported worldwide (0.73–0.99 at 550 nm)
653 (Maggi et al., 2003; Reid et al. 2005; Johnson et al., 2008; Corr et al. 2012; Laing et al. 2016).
654 This can be attributed in part to the high flaming versus smoldering conditions of African smoke
655 producing more BC particles (Andreae and Merlet, 2001; Reid et al., 2005), which inherently have
656 low SSA compared to other regions (Dubovick et al. 2002). However, SSA values over SWA are
657 significantly lower than the range reported near emission sources in sub-Saharan Africa and over
658 the southeast Atlantic, where values span over 0.84–0.90 at 550 nm (Haywood et al., 2003b;
659 Pistone et al., 2019). Recent observations carried out on Ascension Island to the south-west of the
660 DACCWA region showed that smoke transported from Central and South African fires can be
661 very light absorbing over the July-November burning season but SSA values were still higher
662 (0.80±0.02 at 530 nm; Zuidema et al., 2018) than those reported over SWA. A possible cause of
663 the lower SSA in SWA is that Ascension Island is much closer to the local sources and the aerosol
664 is therefore less aged.

665

666 Currently there are few field measurements of well-aged biomass burning emissions. Our
667 knowledge of biomass burning aerosol primarily comes from laboratory experiments and near-
668 field measurements taken within a few hours of a wildfire (Abel et al., 2003; Yokelson et al., 2009;
669 Adler et al., 2011; Haywood et al., 2003b; Vakkari et al., 2014; Zhong and Jang, 2014; Forrister
670 et al., 2015; Laing et al., 2016; Zuidema et al., 2018). Exception made of the study by Zuidema et
671 al. (2018) over the southeast Atlantic, it is generally found that the aged biomass burning aerosol
672 particles are less absorbing than freshly emitted aerosols due to a combination of condensation of
673 secondary organic species and an additional increase in size by coagulation. This is in contrast to
674 our results showing that SSA of biomass burning aerosols were significantly lower than directly
675 after emission and that the evolution of SSA occurred long time after emission.

676

677 There are three possible explanations for these results. First, one must consider sample bias. As
678 regional smoke ages, it can be enriched by smoke from other fires that can smolder for days
679 producing large quantities of non-absorbing particles, thereby increasing the mean *SSA* (*Reid et al.*,
680 2005; *Laing et al.*, 2016). However, during DACCIWA, biomass burning plumes were transported
681 over the Atlantic Ocean and were probably less influenced by multiple fire emissions. Second,
682 there is evidence that fresh BC particles become coated with sulfate and organic species as the
683 plume ages in a manner that enhances their light absorption (*Lack et al.*, 2012; *Schwarz et al.*,
684 2008). Finally, organic particles produced during the combustion phase can be lost during the
685 transport through photobleaching, volatilization and/or cloud-phase reactions (*Clarke et al.*, 2007;
686 *Lewis et al.*, 2008; *Forrister et al.*, 2015), which is consistent with the low *SSA* and *AAE* values we
687 observed. Assessing whether these aging processes impact the chemical components and
688 henceforth optical properties of transported biomass burning aerosol would need extensive
689 investigation of aerosol chemical composition that will be carried out in a subsequent paper.

690

691 **5. Conclusions**

692 This paper provides an overview of *in-situ* airborne measurements of vertically resolved aerosol
693 optical properties carried out over SWA during the DACCIWA field campaign in June-July 2016.
694 The peculiar dynamics of the region lead to a chemically complex situation, which enabled
695 sampling various air masses, including long-range transport of biomass burning from Central
696 Africa and dust from Sahelian and Saharan sources, local anthropogenic plumes from the major
697 coastal cities, and mixtures of these different plumes. This work fills a research gap by providing,
698 firstly, key climate relevant aerosol properties (*SSA*, *MEE*, *g*, *SAE*, *AAE*) and secondly,
699 observations of the impact of aging and mixing processes on aerosols optical properties.

700

701 The aerosol vertical structure was very variable and mostly influenced by the origin of air mass
702 trajectories. While aerosol extinction coefficients generally decreased with height, there were
703 distinct patterns of profiles during dust and biomass burning transport to SWA. When present,
704 enhanced values of extinction coefficient up to 240 Mm^{-1} were observed in the 2–5 km amsl range.
705 These elevated aerosol layers were dominated by either dust or biomass burning aerosols, which is
706 consistent with what would be expected on the basis of the atmospheric circulations during the
707 monsoon season (*McConnell et al.*, 2008; *Knippertz et al.*, 2017). However, during one flight a
708 mixture of dust and biomass burning was found in a layer at around 3 km amsl, implying that there
709 may be substantial variability in the idealized picture. In the lower troposphere, the large

710 anthropogenic pollution plumes extended as far as hundreds of kilometers from the cities emission
711 sources and were not limited to the boundary layer but occurred also at higher levels up to 2.5 km
712 amsl, which is explained by vertical transport and mixing processes, partly triggered by the
713 orography of SWA (*Deroubaix et al., 2019; Flamant et al., 2018a*). The analysis of the aerosol size
714 distributions, *SAE* and NO_x suggests a strong mixing of dust with anthropogenic pollution particles
715 in dust layers transported below 2.5 km amsl, whereas biomass burning plumes that were
716 transported more northward were not affected by this mixing. Both transport pathways and vertical
717 structures of biomass burning and dust plumes over SWA appear to be the main factors affecting
718 the mixing of anthropogenic pollution with dust and biomass burning particles.

719

720 The aerosol light absorption in dust plumes was strongly enhanced as the result of this mixing. We
721 find a decrease of *SSA(550nm)* from 0.92 to 0.81 for dust affected by anthropogenic pollution
722 mixing compared to the situation in which the dust plumes moved at higher altitudes across SWA.
723 Comparison of the particle size distributions of the different dust plumes showed a large
724 contribution of externally mixed fine mode particles in mixed layers, while there was no evidence
725 for internal mixing of coarse particles. Concurrent optical calculations by deconvoluting size
726 distribution measurements in mixed layers and assuming an external mixing state allowed to
727 reproduce the observed *SSAs*. This implies that an external mixing would be a reasonable
728 assumption to compute aerosol direct and semi-direct radiative effects in mixed dust layers.

729

730 Despite a strong increase of aerosol number concentration in air masses crossing urban
731 conglomerations, the magnitude of the spectral *SSAs* was comparable to the background.
732 Enhancements of light absorption properties were seen in some pollution plumes, but were not
733 statistically significant. A persistent spectral signature of biomass burning aerosols in both
734 background and pollution plumes highlights that the aerosol optical properties in the boundary
735 layer were strongly affected by the ubiquitous biomass burning aerosols transported from Central
736 Africa (*Menut et al., 2018; Haslett et al., 2019*). The large proportion of aerosols emitted from the
737 cities of Lomé, Accra and Abidjan that resided in the ultrafine mode particles have limited impact
738 on already elevated amounts of accumulation mode particles having a maximal absorption
739 efficiency. As a result, in the boundary layer, the contribution from local city emissions to aerosol
740 optical properties were of secondary importance at regional scale compared with this large
741 absorbing aerosol mass. While local anthropogenic emissions are expected to rise as SWA is
742 currently experiencing major economic and population growth, there is increasing evidence that
743 climate change is increasing the frequency and distribution of fire events (*Joly et al., 2015*). In

744 terms of future climate scenarios and accompanying aerosol radiative forcing, whether the large
745 biomass burning events that occur during the monsoon season would limit the radiative impact of
746 increasing anthropogenic emissions, remains an open and important question.

747

748 The *SSA* values of biomass burning aerosols transported in the free troposphere were very low
749 (0.71–0.77 at 550 nm) and have only rarely been observed in the atmosphere. The variability in
750 *SSA* was mainly controlled by the variability in aerosol composition (assessed via *k*) rather than by
751 variations in the aerosol size distribution. Correspondingly, values of *AAE* ranged from 0.9 to 1.1,
752 suggesting that BC particles were the dominant absorber in the visible for these biomass burning
753 aerosols. In recent years the southern Atlantic Ocean, especially the area of the west coast of
754 Africa, became an increasing focus in the research community, through the ORACLES/LASIC
755 (ObseRvations of Aerosols above CLouds and their intEractionS/Layered Atlantic Smoke
756 Interactions with Clouds), AEROCLO-sA (AErosol RadiatiOn and CLOuds in Southern Africa –
757 AEROCLO-SA) and CLARIFY (Cloud and Aerosols Radiative Impact and Forcing) projects
758 (*Zuidema et al.; 2016; Zuidema et al.; 2018; Formenti et al.; 2019*). Comparison with literature
759 showed a consistent picture of increasing absorption enhancement of biomass burning aerosol
760 from emission to remote locations. Further, the range of *SSA* values over SWA was slightly lower
761 than that reported on Ascension Island to the south-west of the DACCIWA region, which
762 underscores that the evolution of *SSA* occurred long time after emission. While the mechanism
763 responsible for this phenomenon warrants further study, our results support the growing body of
764 evidence that the optical parameters used in regional/global climate modeling studies, especially
765 absorption by biomass burning aerosols, have to be better constrained using these recent
766 observations to determine the direct and semi-direct radiative effects of smoke particles over this
767 region (*Mallet et al. 2019*). In particular and regarding the very high absorbing properties of
768 smoke, specific attention should be dedicated to the semi-direct effect of biomass burning aerosols
769 at the regional scale and its relative contribution to the indirect radiative effect.

770

771 We believe the set of DACCIWA observations presented here is representative of the regional
772 mean and variability in aerosol optical properties that can be observed during the monsoon season
773 over SWA, as the main dynamical features were in line with climatology (*Knippertz et al., 2017*).
774 This is why results from the present study will serve as input and constraints for climate modeling
775 to better understand the impact of aerosol particles on the radiative balance and cloud properties
776 over this region and also will substantially support remote sensing retrievals.

777 *Data availability.*

778 All data used in this study are publicly available on the AERIS Data and Service Center, which can
779 be found at <http://baobab.sedoo.fr/DACCIWA>.

780

781 *Author contributions.*

782 CD conducted the analysis of the data and wrote the paper. CD, TB, FB, NM, AC, PD, JB, RD, KS
783 and AS operated aircraft instruments and processed and/or quality-controlled data. MM provided
784 expertise on aerosol-climate interaction processes. CF and PK were PIs, who led the funding
785 application and coordinated the DACCIWA field campaign. All co-authors contributed to the
786 writing of the paper.

787

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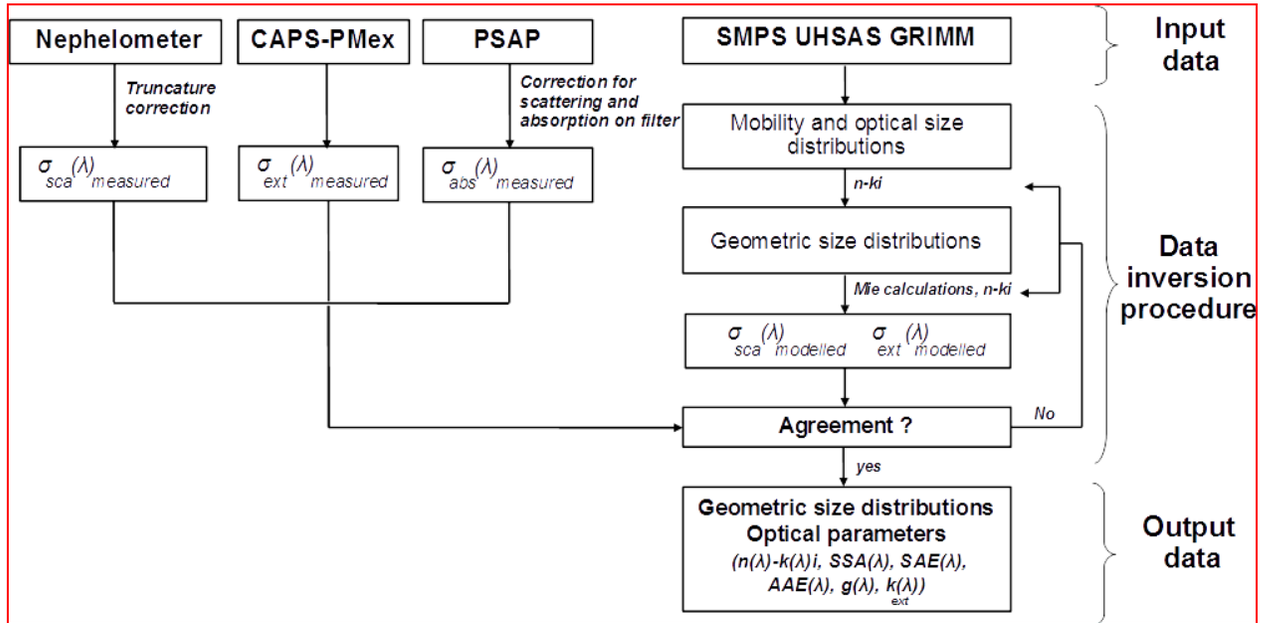
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798 **Appendix 1. Summary of flight information. All flights were conducted during 2016.**

Flight number	Date	Take off time (UTC)	Landing time (UTC)	Events observed
F17	29 June	14:17	17:10	Export of pollution from Lomé
F18	30 June	12:52	16:29	Export of pollution from Lomé
F19	1 July	10:35	14:06	Export of pollution from Accra
F20	2 July	09:53	13:21	Export of pollution from Lomé Dust outbreak
F21	2 July	15:04	18:29	Export of pollution from Lomé Biomass burning outbreak Mixed dust-biomass burning outbreak
F22	3 July	09:54	13:29	Export of pollution from Lomé
F24	6 July	07:17	11:03	Export of pollution from Abidjan
F27	8 July	05:52	09:28	Export of pollution from Accra
F28	8 July	10:53	14:22	Dust outbreak
F29	10 July	10:31	14:11	Export of pollution from Lomé Dust outbreak
F30	11 July	07:19	11:01	Export of pollution from Abidjan Biomass burning outbreak
F31	11 July	13:48	16:42	Biomass burning outbreak
F32	12 July	13:56	17:20	Export of pollution from Accra
F33	13 July	12:40	16:11	Biomass burning outbreak
F35	15 July	09:32	13:00	Export of pollution from Lomé

799

800 **Appendix 2. Data inversion procedure to calculate the aerosol microphysical and optical**
 801 **parameters.**



802

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