

1 **Physico-chemical characterization of urban aerosols from specific combustion**
2 **sources in West Africa at Abidjan in Côte d'Ivoire and Cotonou in Benin in the frame**
3 **of the DACCIWA program**

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5 Aka Jacques Adon¹, Catherine Lioussé¹, Elhadji Thierno Doumbia², Armelle Baeza-Squiban³,
6 Hélène Cachier¹, Jean-Francois Léon¹, Veronique Yoboué⁴, Aristique Barthel Akpo⁵, Corinne
7 Galy-Lacaux¹, Benjamin Guinot¹, Cyril Zouiten⁶, Hongmei Xu^{1,7}, Eric Gardrat¹, Sekou. Keita⁸.

8

9 ¹ Laboratoire d'Aérodynamique, Université de Toulouse, CNRS, UPS, Toulouse, France

10 ² Centre National de Recherche Météorologique (CNRM)/Groupe d'étude de l'Atmosphère
11 Météorologique, CNRS-Météo-France, Toulouse, France

12 ³ Université Paris Diderot, Unité de Biologie Fonctionnelle et Adaptative-RMCX, CNRS, UMR
13 8251, Paris, France

14 ⁴ Laboratoire de Physique de l'Atmosphère, Université Félix Houphouët-Boigny, Abidjan BPV
15 34, Côte d'Ivoire

16 ⁵ Laboratoire de Physique du Rayonnement, Université d'Abomey-Calavi, Abomey-Calavi,
17 Bénin

18 ⁶ Géosciences Environnement Toulouse, Université de Toulouse, CNRS, UPS, Toulouse,
19 France

20 ⁷ Department of Environmental Science and Engineering, Xi'an Jiaotong University, Xi'an,
21 China

22 ⁸ Université de Khorogo, Khorogo, Côte d'Ivoire.

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24 Correspondence to: adonjacks@gmail.com (J. Adon) and lioc@aero.obs-mip.fr (C. Lioussé)

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31 **Abstract**

32 Urban air pollution in West Africa has yet to be well characterized. In the frame of
33 DACCIWA (Dynamics-Aerosol-Chemistry-Cloud Interactions in West Africa) program,
34 intensive measurement campaigns were performed in Abidjan (Côte d'Ivoire) and Cotonou
35 (Benin), in dry (January 2016 and 2017) and wet (July 2015 and 2016) seasons, at different
36 sites chosen to be representative of African urban combustion sources (i.e. domestic fires
37 (ADF), traffic (AT) and waste burning (AWB) sources in Abidjan and traffic source in Cotonou
38 (CT)). Both the size distribution of particulate matter (PM) and their chemical composition
39 including elemental carbon (EC), organic carbon (OC), water-soluble organic carbon (WSOC),
40 water-soluble inorganic ions (WSI) and trace metals were examined. Results show very high
41 PM concentrations at all sites and a well-marked seasonality as well as a strong spatial variation.
42 The average PM_{2.5} mass concentrations during the wet season are 517.3, 104.1, 90.3 and 69.1
43 $\mu\text{g}\cdot\text{m}^{-3}$ at the ADF, CT, AT and AWB sites, respectively. In the dry season, PM_{2.5} concentrations
44 decrease to 375.7 $\mu\text{g}\cdot\text{m}^{-3}$ at the ADF site, while they increase to 269.7, 141.3 and 175.3 $\mu\text{g}\cdot\text{m}^{-3}$
45 at the CT, AT and AWB sites, respectively. The annual PM_{2.5} levels at almost all sites are
46 significantly higher than the WHO guideline level of 10 $\mu\text{g}\cdot\text{m}^{-3}$. As for PM mass, (EC) and
47 (OC) concentrations are also maximum at ADF site, accounting up to 69% of the total PM mass.
48 Such high content is mainly linked to wood burning for domestic cooking and commercial food
49 smoking activities. Dust contributions are dominant at CT (57-80%), AT (20-70%) and AWB
50 (30-69%) sites and specially in the coarse and fine particle modes at CT and in the coarse
51 fraction at AT site, which may be explained by the impact of long-range desert-dust transport
52 and re-suspended particles from the roads, in addition to anthropogenic sources. The
53 contributions of WSI to the total PM mass, mainly driven by chloride, nitrate and calcium in
54 the fine and/or large particles are highly variable according to the sites but remain less than
55 30%. Values are generally 1-3 times higher in the wet season than the dry season. This is due
56 to anthropogenic emissions but also to nitrate formation by reaction processes and natural
57 emissions. The concentrations of trace element well reflect the trends of dust at the traffic and
58 AWB sites, with a predominance of Al, Na, Ca, Fe and K, keys markers of crustal dust. This
59 study constitutes an original database that characterizes specific African combustion sources.

60

61 **Keywords:** atmospheric pollution, chemical composition, physicochemical characterization,
62 Particulate matter, traffic, waste burning, domestic fires.

63 1. Introduction

64 The impact of anthropogenic pollution on environment and health has been demonstrated by
65 numerous studies in Europe and North America, which have contributed to the implementation
66 of emission reduction policies. By contrast, air pollution in Africa is far from being well
67 characterized, although it is suspected to be responsible for negative health outcomes (*WHO*,
68 2014). This is a major problem since Africa is an intense emitter of pollution from
69 anthropogenic sources that includes domestic fires, vehicular traffic, waste burning as well as
70 growing oil and mining industries. It has also one of the fastest growing urban populations in
71 the world, especially in West and East Africa (UN, 2019). As a consequence, it has been shown
72 that massive urbanization and rapid economic growth could be responsible for tripling
73 anthropogenic emissions in Africa between 2000 and 2030 (*Lioussse et al.*, 2014). Moreover, it
74 is important to recall the impact of biomass burning and dust sources in the African atmospheric
75 composition, especially occurring during the dry season. All of this results in a major
76 degradation of urban air quality and an impact on the health of exposed populations. Only a few
77 studies on this subject have been conducted in West Africa (*Val et al.*, 2013; *Dieme et al.*, 2012;
78 *Kouassi et al.*, 2009) despite the high atmospheric pollutant concentrations already measured
79 to be of the same order as in Asian megacities and well above WHO international standards
80 (*WHO*, 2014).

81 West Africa is then an "unique laboratory" to study urban pollution. Previous studies conducted
82 under the framework of the AMMA (Analyses Multidisciplinaires de la Mousson Africaine)
83 and POLCA (POLlution des Capitales Africaines) programs, have revealed very high average
84 particulate mass concentrations in Cotonou (Benin), Bamako (Mali), Dakar (Senegal) and
85 Yaoundé (Cameroun) during the dry season (*Doumbia et al.*, 2012; *Val et al.*, 2013), suggesting
86 that the population may be affected by negative health outcomes. For example, during the dray
87 season in Bamako (Mali) and Dakar (Senegal), *Val et al.* (2013) showed that the inflammatory
88 impact of combustion aerosol depends on the type of emission sources and determined the
89 predominant role of particulate organic matter. This is consistent with global findings showing
90 that fine and ultrafine aerosol fractions, as well as their content in trace metals and organic
91 compounds, induce biological effects due to their ability to reach the distal lung (*Cassee et al.*,
92 2013). Such reasons highlight the need to better understand the size-speciation of aerosol
93 chemical composition for the main West African anthropogenic sources during the different
94 seasons. Within this context, the DACCIIWA (Dynamics-Aerosol-Chemistry-Cloud
95 Interactions in West Africa) program, dedicated a specific work package to "Air Pollution and

96 Health” dealing with pollutant characterization related to health issues through toxicological
97 studies and epidemiological studies.

98 Campaigns have been organized from December 2014 to March 2017 in Abidjan and Cotonou.
99 The strategy was to measure aerosol chemical composition in different sites, representative of
100 the main prevailing urban sources in West Africa following *Liousse et al. (2014)* and *Keita et*
101 *al. (2018)*. Two typical traffic-sampling sites were chosen, one in Abidjan (Côte d’Ivoire) and
102 another one in Cotonou (Benin), to take into account differences in terms of fleets, type of fuel
103 used and quality of roads. Indeed, in Cotonou, the majority of population uses two-wheel
104 vehicles susing gasoline fuel or gasoline and oil fuel, whereas in Abidjan, the vehicle fleet is
105 dominated by four-wheel engines using diesel fuel. Measurements were also performed at
106 domestic fire and waste burning sites, both located in Abidjan.

107 For a period of two years, PM_{2.5} mass and carbonaceous aerosol were weekly measured and
108 results are discussed in *Djossou et al. (2018)*. In this paper, we focus on the results from the
109 intensive campaigns. We present measurements obtained at each site during the wet and dry
110 seasons of the studied periods: (i) PM size distribution and mass concentrations and (ii) PM
111 chemical composition including carbonaceous aerosol, water-soluble organic carbon, water-
112 soluble inorganic ions, dust and trace elements in different size fractions. Experimental method
113 including description of sites, types of measurements and analyses, meteorological conditions,
114 will be presented in the section 2, whereas results and discussion are discussed in the sections
115 3 and 4 of the paper, respectively.

116

117 **2. Experimental method**

118 **2.1. Description of sites**

119 Measurement campaigns have been performed in wet season (July 20-26, 2015 and July
120 4-13, 2016) and dry seasons (January 7-15, 2016 and January 5-14, 2017) at three sites in
121 Abidjan (Côte d’Ivoire), representative of different sources, i.e. ADF for Abidjan Domestic
122 Fires, AWB for Abidjan Waste Burning and AT for Abidjan Traffic (Figure 1), as well as one
123 traffic site in Cotonou (Benin) (Figure 2).

124 As shown in Figure 3 which presents pictures of the different sampling sites, the ADF site (5°
125 19' 44 "N, 4° 06' 21" W) is situated on a platform, 5 m above ground level, in Yopougon

126 Bracody district near a market (Figure 1). This geographical area is highly populated with
127 various small commercial activities such as a fish and meat-smoking by women. There are also
128 many formal and informal settlements, which mainly use wood and charcoal as a source of fuel
129 for private and professional combustion activities. Other sources of concern contributing to the
130 mix of pollutant emissions in the area include transportation-related emissions, biomass
131 burning, garbage bins or small landfills and various other fugitive sources. The AT site (5° 21'
132 14" N, 4° 01' 04" W) is located in Adjamé, on the roof of « 220 pharmacie logement building »,
133 about 7 m above ground level and roughly 10 m away from the main road. This site, close to
134 the Adjamé market and to a bus station, is highly affected by traffic (Gbaka, bus, taxi, woro-
135 woro, private cars...). The AWB site (5° 21' 12" N, 3° 56' 16" W) is located at Akouédo in the
136 district of Cocody, on the roof of « Talafigué », a building 15 m above ground level. This site
137 close to the big municipal landfill of Abidjan operational since 1965 and now covering an area
138 of 153 ha, is submitted to frequent waste burning pollution. The Cotonou Traffic (CT) (6° 22'
139 19" N, 2° 26' 5" E) site is located in Cotonou, on the «Sogema» building roof, about 7 m above
140 ground level. This site is close to the Dantokpa market and also to the biggest crossroad of
141 Cotonou (intersection of 4 main roads). This site is highly influenced by intense traffic
142 activities. As previously mentioned, such a site is interesting because the vehicle fleet and fuels
143 are different in Cotonou compared to Abidjan: (1) there are many two-wheel vehicles in
144 Cotonou whereas a few only in Abidjan; (2) in Cotonou, gasoline is of poor quality due to the
145 illegal fuel transport from Nigeria and (3) the roads are in worse conditions in Cotonou than in
146 Abidjan.

147 **2.2. Measurements**

148 During each intensive campaign and on each site, two 3-hour samples collected with
149 cascade impactors operating in parallel are obtained for three consecutive days (i.e. six size-
150 resolved samples per site during each campaign), to allow size-speciated characterization of the
151 aerosol chemical composition. Note that the choice of the 3-hour periods is linked to the period
152 of maximum pollution for each site as shown by preliminary studies (e.g. morning at ADF site
153 (7-10am), afternoon at AT site (4-7pm, morning at CT site (7-10am)). There is no specific
154 period at AWB site since the activities are roughly the same during the day. The first impactor
155 with 4 stages ($PM_{>2.5}$; $PM_{2.5-1}$; $PM_{1-0.2}$; $PM_{<0.2}$) includes 4 quartz fiber filters (QMA, Whatman)
156 for mass and carbonaceous aerosol (EC, OC and WSOC analysis). The second impactor with 3
157 stages ($PM_{>2.5}$; $PM_{2.5-1}$; $PM_{1-0.1}$) is equipped with three Teflon filters (Zefluor, Pall
158 Corporation®), dedicated for water-soluble ions species and trace elements. Due to operational

159 problems in July 2016, this second 3-stage cascade impactor is replaced by another 3-stage
160 cascade impactor with different size cuts ($PM_{>10}$, $PM_{10-2.5}$, $PM_{2.5-1}$). For consistency, results will
161 be presented as an ultrafine (UF), fine (F) and coarse (C) classification. The two first stages
162 ($PM_{>2.5}$ and $PM_{2.5-1}$) being considered as the coarse particulate fraction, the $PM_{1-0.1}$ or $PM_{1-0.2}$
163 stage, the fine particulate fraction and the $PM_{<0.2}$ stage, the ultrafine fraction.
164 All the filters are prepared and analyzed at the Laboratoire d'Aerologie in Toulouse under
165 different protocols described in the following paragraphs. Note that the quartz filters are
166 prefired before sampling.

167 **2.3. Analyses**

168 **2.3.1. Gravimetric analyses**

169 Aerosol mass concentrations are obtained using a high-precision balance (SARTORIUS
170 MC21S), placed under a controlled temperature and humidity atmosphere (Person and Tymen,
171 2005). Before weighing, the filters are kept 24 hours in the weighing room at an ambient relative
172 humidity of $30 \pm 15\%$. The filters are weighed before and after sampling. Result of a gravimetric
173 measurement consists of the average of 2 to 4 weighing whose differences do not exceed $5 \mu\text{g}$.
174 The standard error on a gravimetric measurement is therefore less than $10 \mu\text{g}$, typically
175 representing less than 5% of the particles mass.

176 **2.3.2. Carbonaceous aerosols**

177 Carbonaceous aerosol is determined with thermal analysis with a two-step method
178 adapted from Cachier et al. (1989). Two aliquots of the same filter are separately analyzed.

179 One portion is directly analyzed for its total carbon content (TC). The other portion is
180 first submitted to a pre-combustion step (2 h at 340°C under pure oxygen) in order to eliminate
181 Organic Carbon (OC), and then analyzed for its Elemental Carbon (EC) content. Organic
182 carbon (OC) concentrations are calculated as the differences between TC and EC. Note that the
183 aerosol carbon content is quantified by a non-dispersive infrared (NDIR) detector with G4
184 ICARUS instrument with a detection limit of the order of $2 \mu\text{gC}\cdot\text{cm}^{-2}$. Uncertainty is in the
185 order of 5% for TC, while being in the range of 5-20%, for EC and OC.

186 **2.3.3 Water Soluble Organic Carbon analysis**

187 WSOC measurements are performed using a total organic carbon analyzer (Sievers M9).
188 A detailed description of this technique is reported in Favez et al. (2008). Briefly, the full

189 oxidation of total organic carbon into CO₂ is obtained by coupling chemical oxidation (with
190 ammonium persulphate) and UV light. CO₂ is then quantified by conductivity. Analyses are
191 conducted on 20 ml of solution extracts. For UF samples, solutions to be analyzed are obtained
192 using a total filter surface of 3cm² (6x0.5 cm² punches symmetrically taken out of each QMA
193 filter), whereas, for C and F sizes, due to the geometry of the spots at the surface of the filters,
194 samples are divided into equivalent parts (1/2 or 1/4 of 47 mm filters, rest of the filters being
195 used for carbonaceous analysis). The extraction protocol consists of 16h soaking under soft
196 shaking in an Erlen-Meyer containing 20mL of ultra-pure water. Prior to WSOC analysis, water
197 extracts are filtered through Teflon (PTFE) filters (0.2µm pore size diameter) in order to remove
198 any suspended particle. Measurement uncertainty, given by the manufacturer, is of the order of
199 7%. The overall calculated blank value is of the order of $2.27 \pm 0.33 \mu\text{gC}\cdot\text{cm}^{-2}$, which represents
200 $16.4 \pm 8.5\%$ of the mean WSOC content. For each sample, duplicate analyses show a good
201 reproducibility.

202 **2.3.4. Water-soluble ionic species**

203 Water-soluble ionic species (Na⁺, NH₄⁺, K⁺, Mg²⁺, Ca²⁺, SO₄²⁻, NO₃⁻ and Cl⁻) are
204 analyzed using ion chromatograph (IC), following the analytical protocol described in Adon et
205 al. (2010). Briefly, the aerosol water-soluble fraction is first extracted from half-sampled Teflon
206 filter (the other part being used for trace element analysis), with a 10-min sonication in plastic
207 vials including 6 ml or 10 ml of purified water with a controlled resistivity of 18.2MΩ. Then
208 these vials are subjected to ionic chromatograph analysis or stored at +4°C if not analyzed
209 immediately. Cations are analyzed with Dionex DX-100 and anions with Dionex DX-500 with
210 a detection limit of 1 to 6 ppb depending on ionic species. Uncertainties in the range of 1-50%
211 is found depending on ionic species.

212 **2.3.5. Trace elements**

213 The protocol to measure trace element concentrations is developed and performed at the
214 Laboratory of Environmental Geosciences of Toulouse. Half-sampled Teflon filters (the other
215 part being used for water-soluble ionic species, see below) are mineralized by acid digestion
216 with a 10 ml concentrated HNO₃ and 0.5 ml HF solution (Lamaison, 2006) using a closed vessel
217 microwave accelerated reaction system (MARS 5, CEM Corporation) at high pressure (700 psi)
218 (Celo et al. 2010). The digestion is realized in 3 steps: a rise in temperature at 130°C in 3min
219 and holding for 1 min, then, a second rise at 160°C in 1 min and holding for 30 seconds and
220 finally a third rise to 180°C in 1 min and holding for 3min. After a 12 h cooling period, the

221 solutions are evaporated at 80°C, and concentrated in 7 ml of 2% concentrated HNO₃ solution,
222 before analysis by ICP-MS which are performed with a 7500 ce Agilent Technologies
223 instrument equipped with a collision cell, and using In and Re as internal standards. The
224 detection limit is less than 10 ppt. For all the samples, the final blank values and detection limit
225 on filters are taken into account for final concentrations calculations. 13 trace metals are
226 considered in this work: Al, Ti, Cr, Mn, Fe, Ni, Cu, Zn, Ba, La, Th, Pb and Cd.

227 **2.3.6. Dust calculation**

228 Many methods can be used to quantify dust concentrations. We have selected three
229 methods (Sciare et al. 2005, Guinot et al. 2007, Terzi et al. 2010) to underline the uncertainties
230 linked to dust estimates.

231 (1) Sciare et al. (2005) method consists of using soluble calcium data obtained with Ionic
232 Chromatography (IC), to estimate the dust concentrations following the relationship:
233 $\text{dust} = 10.96 * \text{nss-Ca}^{2+}$, where $\text{nss-Ca}^{2+} (=1.02 * \text{Ca}^{2+} - 0.038 * \text{Na}^+)$ refers to non-sea-salt
234 calcium concentration.

235 (2) Guinot et al. (2007) method is based on a chemical closure where fine and coarse
236 particle aerosols are separated in 4 components (EC, POM, WSI and dust). EC, WSI,
237 and total aerosol mass are directly experimentally determined (see below paragraphs).
238 POM concentrations are obtained from OC concentrations experimentally determined
239 and k, the OC/POM conversion factor. Dust concentrations are obtained from measured
240 Ca^{2+} concentrations and f, the abundance of calcium in dust. The k and f values are
241 obtained from a linear regression (L) between the reconstructed and the weighed aerosol
242 mass concentrations. Briefly, first step consists of focusing on the aerosol coarse
243 fraction. k is fixed to 1.8 and as a result of (L) just mentioned, f is obtained to be in the
244 range of 0.012 to 0.15 depending on our sites. Second step deals with the aerosol fine
245 fraction. The f values just obtained for the aerosol coarse fraction are applied to the fine
246 fraction and k ratios are estimated using (L) to be in the range of 1.2 to 2.1. Note that at
247 all of our sites, the correlation between Ca^{2+} and the missing mass between the weighed
248 and the reconstructed aerosol mass is sufficiently good ($r^2=0.9$) to support the
249 consistency of this simple approach for the evaluation of dust. Also, f and k values are
250 included in the range of values provided in the literature (He et al. 2001; Sun et al. 2004;
251 Guinot et al. 2007). However, it is important to mention that the range of f and k
252 coefficients are large which is due to the source mixing observed in this study.

253 (3) In Terzi et al. (2010) method, dust is obtained with the following relationship: dust =
254 $1.89[\text{Al}] + 1.21[\text{K}] + 1.95[\text{Ca}] + 1.66[\text{Mg}] + 1.7 [\text{Ti}] + 2.14[\text{Si}] + 1.42[\text{Fe}]$. In our study,
255 all these elements are determined except Silica (Si). Consequently, we used mean Si
256 values obtained from different relationships available in the literature ($\text{SiO}_2 = 3 * \text{Al}_2\text{O}_3$
257 for Alastuey et al., 2005, $\text{Si} = 4.0 * \text{Al}$ for Zhang et al., 2003 and $\text{Si} = 2.03 * \text{Al}$ for
258 Chiapello et al., 1997).

259 The results of dust concentrations estimated from the three methodologies above described are
260 summarized in the Table 1 for wet season (WS) 2016 and dry season (DS) 2017. Indeed, Ca,
261 Al, and Fe concentrations measured by ICP-MS are only available in WS2016 and DS2017 due
262 to experimental problems, whereas Ca^{2+} concentrations measured by IC are available for all
263 campaigns. As shown in table 1, the dust obtained from Ca^{2+} measured by IC (Sciare et al.,
264 2005) and by the Guinot et al. (2007) method is lower than that obtained from trace elements
265 (Terzi et al., 2010) for DS2017 whereas in the same order of magnitude in WS2016. Such
266 results are in agreement with methodological aspects. Indeed, Al, Fe, Ca ... obtained by ICP-
267 MS include both soluble and insoluble particles whereas Ca^{2+} measured by IC only include
268 soluble particles. During the dry season, comparison of Ca measured by ICP-MS (not shown
269 here) is higher than that of the IC, by a factor of 1.7, 1.8, 2.2 and 1.1, at the ADF, AWB, AT,
270 and CT sites respectively. By contrast, this factor is low and constant (1.3) in the wet season
271 for all the sites. In our study, due to the lack of trace element data for WS2015 and DS2016,
272 dust estimations are performed from Guinot et al. (2007) method. This choice globally implies
273 an underestimate of dust concentrations by a factor of 1.5 to 3.5 in DS2017 as shown in Table
274 1.

275

276 **2.3.7. Aerosol chemical closure methodology**

277 As previously mentioned and detailed, aerosol chemical closure is performed following
278 the Guinot et al. (2007) methodology.

279

280 **2.4. Meteorological conditions**

281 In Figure 4, meteorological data (surface temperature, wind directions and speed) issued
282 from the NOAA Integrated Surface database (ISD; see [https:// www.ncdc.noaa.gov/isd](https://www.ncdc.noaa.gov/isd)) and the
283 ASECNA (Agence pour la Sécurité de la Navigation Aérienne en Afrique et à Madagascar) are
284 presented for the South-West Africa region including Abidjan and Cotonou. As expected, this

285 area is under the influence of the Convergence Zone of two air masses of a different nature, i.e.
286 Harmattan (hot and dry continental trade winds) from the north and Monsoon (humid maritime
287 trade winds) from the south (Figure 4). Ground contact between these two air masses constitutes
288 the intertropical front (ITF) of which the fluctuations during the year determine the seasons in
289 the Gulf of Guinea (Tapsoba, 1997). During the dry season (from November to March),
290 temperatures are relatively high with maximum around 30°C on the coast. The humidity is low,
291 since the prevailing Harmattan wind blows from the desert, usually bringing dust (Figure 4,
292 lower line). The period from June to September, especially in July is the wet season period
293 when daytime temperatures are slightly lower, with maximum around 26/28°C on the coast
294 (Figure 4, upper line). At this season, the humidity level is high across the region. On the coast,
295 rains may occur from March to November.

296 During our campaigns (not shown here), temperatures are roughly the same at Abidjan and
297 Cotonou, reaching 28°C and 26°C in the dry and wet seasons, respectively. Gentle to moderate
298 wind speeds are observed during the measurement campaigns at the two cities, with average
299 values of 15-20 and 15-22 km.h⁻¹ at Abidjan and Cotonou, respectively. There is no
300 precipitation at CT site during the studied periods. In Abidjan on the contrary, low rains occur
301 both in wet and dry periods with cumulative precipitation higher in DS2017 (7mm), than in
302 WS2016 (4.7mm) and WS2015 (2mm). There is no rain in DS2016 ([https://www.historique-](https://www.historique-meteo.net/afrique/)
303 [meteo.net/afrique/](https://www.historique-meteo.net/afrique/)).

304 **2.5. Backward trajectories**

305 The Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) modelling
306 system (Air resources laboratory, Draxler and Rolph, 2012) is used for the trajectory analysis.
307 HYSPLIT model is run to compute 120 h back trajectories ending at Abidjan and Cotonou at
308 50 m a.g.l. (Figure 5). Global Data Assimilation System reanalysis database is used as
309 meteorological input, with a 0.25 × 0.25 degrees horizontal resolution. Results presented in
310 Figure 5, confirm that air masses mainly come from the north with a few from the south-west
311 in dry season (January), whereas from the south-west and the south in wet season (July).
312 Therefore, in January, Abidjan and Cotonou are mainly impacted by polluted air masses from
313 surrounding areas and northern countries with possible dust and west African biomass burning
314 influences, whereas in July, the impact of oceanic sources possibly polluted by long-range south
315 African biomass burning aerosols may be observed.

316

317 **3. Results**

3.1. Aerosol size distribution and mass concentration

In Figure 6, the relative mass distribution of PM for Coarse (C), Fine (F) and Ultra-Fine (UF) particle sizes in percentages are presented with bulk mass concentration averages indicated in the black boxes for each site and for each campaign. As it may be seen, bulk concentrations vary widely from site to site and from campaign to campaign. During the wet season, the average total concentrations range from 82 to 676 $\mu\text{g.m}^{-3}$ in 2015 and 56 to 358 $\mu\text{g.m}^{-3}$ in 2016, with the maximum at the Abidjan Domestic Fire (ADF) site. While during the dry season, values range from 168 to 269 $\mu\text{g.m}^{-3}$ in 2016 and from 114 to 559 $\mu\text{g.m}^{-3}$ in 2017, with maximum concentration obtained at the Cotonou Traffic (CT) and ADF sites. In terms of size distribution, concentration peaks may be observed for all aerosol size-fractions which are found to exhibit different seasonal patterns. UF particles ($<0.2 \mu\text{m}$) represent the highest contributor to the bulk mass at the ADF site, by up to 60 % (335.3 $\mu\text{g.m}^{-3}$) in DS2017. F particles (1-0.2 μm) are the second most important contributor and both combined particle sizes account for more than 85 % of the total mass at the ADF site. In this site, ultra-fine and fine fractions are also found to be maximum during WS2015 and WS2016 by up to 90 and 83%, respectively. Let us note that C particle contribution in bulk is relatively higher in the traffic and waste burning sites than in ADF site (40%) whereas F and UF particle contributions are on the order of 60%.

In terms of PM_{2.5}, the results of this work are presented in Figure 7. The mass concentration of PM_{2.5} averaged over DS2016 and DS2017 are 154 ± 74 , 144 ± 42 , 134 ± 7 and $211 \pm 51 \mu\text{g.m}^{-3}$ at the ADF, AWB, AT and CT sites, respectively and 338 ± 24 , 45 ± 3 , 52 ± 4 and $70 \pm 1 \mu\text{g.m}^{-3}$ over the wet seasons (2015-2016). The increase in PM_{2.5} is of the order of 54% at ADF from dry to wet season, whereas a sharp reduction (more than 60%) is obtained at AWB, AT, and CT sites.

3.2. Carbonaceous aerosol

3.2.1. EC and OC concentrations

In Figure 8, EC relative mass contributions are presented for each size, site and campaign: wet season 2015 (WS2015), wet season 2016 (WS2016), dry season 2016 (DS2016) and dry season 2017 (DS2017). Mean EC bulk mass concentrations are added in the black boxes for each size and for each campaign. The most striking feature is that the ADF site concentrations are higher than at the other sites in WS2016 and in DS2017, whereas of the same

349 order of CT site concentrations in the other seasons. Mean concentration at the CT site ($16\mu\text{g}\cdot\text{m}^{-3}$)
350 ³) is slightly higher than at the AT site ($10\mu\text{g}\cdot\text{m}^{-3}$), whereas the lowest concentrations are found
351 at the AWB site. Results of the EC size distribution are very consistent among the different
352 sites (Figure 8). Whatever the site and the season, higher EC concentrations are found in C
353 (42%) and UF (43%) particles compared to F particles.

354 Same data are presented for OC concentrations in Figure 9. It may be underlined that
355 ADF OC values are always higher than in the other sites by a factor ranging from 6 to 30, for
356 all seasons and particle sizes, with highest and lowest values respectively in DS2017 and DS
357 2016. In terms of size distribution, maximum OC concentrations at the ADF site may be found
358 in UF (53%), then F (29%) and finally C (18%) particles. The same distribution is observed for
359 the traffic sites in DS2016, however, for the other campaigns, OC size distribution looks like
360 the EC ones with higher concentrations in UF and C particles than in F particles.

361 As shown in Figure 10, the highest OC/EC ratios are always obtained at the ADF site
362 with a value as high as 25 for F particles in WS2016 whereas the lowest values are found in
363 DS2017. This is the same feature for the other sites with ratios lower than 2 in DS2017. OC/EC
364 ratios in AWB site are higher than in the traffic sites. Note that values at AT site are higher than
365 CT values in the wet season whereas lower in the dry season. Finally, it is interesting to
366 underline that linear correlations between EC and OC are obtained in the ultrafine and fine
367 modes in all campaigns, particularly in DS2017 ($r^2 = 0.8, 0.8, 0.9$ and 0.9) at the ADF, AWB,
368 AT and CT sites, respectively. This suggests that different studied sources can be assessed as
369 significant sources of both EC and OC.

370

371 **3.2.2. Water-Soluble Organic Carbon**

372 Concentrations of WSOC and WSOC/OC ratios are presented in Table 2 for each size
373 (UF, F, C and $\text{PM}_{2.5}$) and campaign. As seen, WSOC are always higher at the ADF site than
374 in other sites, at least by a factor of 12. Maximum values are obtained in WS2016 with an
375 average of 16.47, 17.08 and $79.68\mu\text{gC}\cdot\text{m}^{-3}$ for coarse, fine and ultra-fine fractions, respectively,
376 followed by WS2015 and DS2017. WSOC concentrations are the lowest in DS2016, with an
377 average of 4.14, 6.95 and $21.89\mu\text{g}\cdot\text{m}^{-3}$ for coarse, fine and ultrafine fractions, respectively. In
378 terms of seasonality, there is not a clear trend in WSOC values at the AWB and AT sites,
379 whereas at the CT and ADF sites, WSOC values are found to be respectively higher and lower

380 in dry seasons compared to wet seasons. It is also interesting to note that WSOC are maximum
381 in UF sizes in the AT, ADF and AWB sites. At the CT site, the highest values are found in the
382 coarse particulate fractions, except in DS2016.

383 As expected, WSOC is strongly correlated with OC ($r=0.7$ at ADF, 0.8 at AT, 0.5 at
384 AWB and 0.7 at CT), whereas correlations with EC are weaker, especially at the AWB and CT
385 sites with values ranging from 0.1 to 0.4 , respectively. Finally, when looking at WSOC/OC
386 ratios (Table 2), maximum values are obtained at the ADF site with PM_{2.5} ratios as high as
387 43% , followed by the AT and AWB sites with 32% . The lowest value (23%) is found at the CT
388 site. Also, Table 2 shows that there is no clear seasonality in WSOC/OC values, excepted at
389 ADF where maximum values occur during the wet season. Note as for WSOC, that ratios are
390 maximum in UF and F fractions for all sites except at the CT site where the ratio for coarse
391 fraction is the highest.

392 3.3. Water-soluble ionic species

393 Figure 11 shows the relative contribution of the major ions to the total concentration
394 (also given) of the ions in the different particle modes (C and F) at the ADF, AWB, AT and CT
395 sites for the different measurement campaigns. Let us recall here that only C and F fractions
396 may be documented due to the our experimental protocol. Total concentrations present
397 maximum values in ADF and CT sites. Values in AWB and AT sites are of the same order of
398 magnitude and lower by a factor of 2 than in ADF and CT sites. The contribution of different
399 ions show significant variations from site to site. The dominant ionic species at the ADF site
400 over all campaigns is chloride (Cl^-), with a 26% contribution, followed by nitrate (NO_3^-) (16%),
401 calcium (Ca^{2+}) (13%) and potassium (K^{2+}) (12%). Sulfate (SO_4^{2-}), ammonium (NH_4^+), sodium
402 (Na^+) and to a lesser extent magnesium (Mg^{2+}) contributions are lower, ranging from 4 to 7%
403 of the total ion species. The lowest contribution is for organic acids with their total value lower
404 than 5% . NO_3^- is the major ionic component at the AWB and AT sites, representing 24% and
405 29% of the total water soluble inorganic concentration, respectively. The second major
406 contributor in AWB and AT is SO_4^{2-} , accounting for 21% and 17% of the ion mass, respectively
407 followed by Ca^{2+} (12% and 15%) and Cl^- (15% and 13%). In CT, Ca^{2+} is predominant with a
408 relative abundance of 24% , followed by NO_3^- (23%), SO_4^{2-} (19%) and Cl^- (13%). Na^+ , NH_4^+
409 and K^+ contributions are lower and in the same order of magnitude in AT, AWB and CT sites,
410 ranging from 4 to 9% of the total ion species. Note that organic ion contributions at AT, CT and
411 AWB is of the same order than in ADF, with lower values at CT. It is interesting to underline
412 in the Figure 11, that NO_3^- contribution is always higher in the coarse than in the fine size.

413 Conversely, K^+ is always higher in the fine than in the coarse size. In CT, Ca^{2+} in the fine
414 fraction is as high as in the coarse fraction whereas in AT, AWB and ADF, Ca^{2+} coarse fraction
415 is predominant. Fine particle contribution may be noticed for Cl^- in ADF whereas in the other
416 sites, Cl^- is most likely dominated by coarse particles. Finally, SO_4^{2-} is mainly found in the fine
417 mode at the AT, AWB and CT sites, but in the coarse mode in ADF site.

418 In terms of seasonal variations, it may be shown in Figure 11 that higher Cl^- values are found
419 in wet seasons than in the dry seasons everywhere, except in ADF site where there is no marked
420 difference between seasons. For example, the mean relative total percentages of Cl^- at the CT
421 site are 38 and 24% in the WS2015 and WS2016, respectively, while these percentages decrease
422 significantly to 18 and 13% in the DS2016 and DS2017, respectively. The Cl^-/Na^+ ratios are
423 about 1.5 everywhere in both seasons, in agreement with the typical sea water ratio (1-1.2)
424 (Hara et al.,2004), except at the ADF site where these ratios increase to 4 and 5 in wet and dry
425 season, respectively and at the AWB site in the dry season (2). K^+ and Ca^{2+} are always higher
426 in dry season than in wet season except for Ca^{2+} in ADF where values are of the same order.
427 Finally, the same trend is observed for NO_3^- and SO_4^{2-} with higher values in dry than in wet
428 seasons at AWB and CT sites whereas values at ADF and AT sites are of the same order of
429 magnitude for the two seasons.

430 **3.4. Trace element concentrations**

431 Table 3 shows the mean values of the major trace elements in bulk aerosol at the
432 different studied sites in WS2016 and DS2017, with their corresponding relative abundances in
433 the total aerosol mass into brackets. Let us recall that data are not available in WS2015 and
434 DS2016. The concentrations of trace elements span a wide range, from 0.2 to 25.2 $\mu g.m^{-3}$.
435 Among the measured elements, Al, K, Na and Ca are the most abundant, followed by Fe and
436 Mg. In DS2017, Al and Na concentrations are higher in AWB than in the other sites. The
437 minimum value for these species is found in ADF site. Values in traffic sites are of the same
438 order of magnitude and higher than in ADF site. Maximum of Ca and K values may be found
439 in CT and ADF site respectively. It is interesting to note that Al, K, Na concentrations are higher
440 in the dry season than in the wet season. Such feature is less clear for Ca, whose seasonal
441 variability is less marked except in AWB and AT sites. In terms of Mg, maximum values are
442 observed in ADF site and of the same order of magnitude whatever the season. Fe abundance
443 is higher in AWB and CT sites than in ADF and AT sites and higher in DS2017 than in WS2016
444 everywhere. The other metals (Ti, P, Zr, Zn, Cr, Mn, Pb and Ni) represent less than 0.5% and

445 2% of the total mass in WS2016 and DS2017, respectively, at all sites, with Cr, Mn, Pb and Ni
 446 exhibiting less seasonal variability compared to the rest of the metal elements.

447 To assess the relative contribution of crustal and non-crustal origin of elemental aerosol
 448 loadings, source enrichment factor (EF) of a trace element X have been first calculated with the
 449 following formula using both literature data of the typical elemental composition of the upper
 450 continental crust (Mason and Moore, 1982; Taylor, 1964), measured elemental composition
 451 from this study and Al as a reference element :

$$452 \quad EF_X = \frac{\frac{[X]_{atm}}{[Al]_{atm}}}{\frac{[X]_{soil}}{[Al]_{soil}}}$$

453 Where $[X]_{atm}$ and $[Al]_{atm}$ are the concentrations of the chemical element X and Al in the
 454 atmosphere, respectively, and $[X]_{soil}$ and $[Al]_{soil}$ are the typical concentrations of the element
 455 X and Al in the earth's crust, respectively. Al is frequently used as a reference element assuming
 456 that its anthropogenic sources in the atmosphere are negligible (Gao et al., 2002; Cao et al.,
 457 2005; Xu et al., 2012). In all sampling sites, EF values typically lower than 5 are obtained for
 458 several trace elements (Be, Sc, Ti, V, Fe, Ga, Sr, Nb, Rh, Ba, La, Ce, Pr, Nd, Sm, Eu, Gd, Tb,
 459 Dy, Ho, Er, Tm, Yb, Lu, Ta, Th and U). This suggests a natural origin of these species (Freitas
 460 et al., 2007; Gao et al., 2002). The most enriched elements ($EF > 100$) are Sb, Sn, Zn, Se, Te,
 461 Cd, Pb, Bi and Mo at nearly all of the sites, indicating significant anthropogenic origin (Wang
 462 et al., 2006). These elements are mainly emitted into the atmosphere through fossil fuel
 463 combustion, traffic emission, wear of brake lining materials and industrial processes (Watson
 464 and Chow, 2001; Samara and al., 2003). Secondly, source contributions have been estimated
 465 from these EF values following the method described by Arditsoglou and Samara (2005). Note
 466 that this study refers to ratios for a limited list of sources, perhaps not including the African
 467 source specificities. As a result, it may be seen that 30% of trace element concentrations is of
 468 anthropogenic origin at ADF site whereas about 17 % at the others sites.

469 **3.5. Dust**

470 Figure 12 shows dust concentrations calculated from Guinot et al. (2007) methodology
 471 (see paragraph 2.3.6) for C and F particle sizes at the different sites for each season. Note that
 472 as for WSI and trace element and due to our sampling procedure, there are values for fine and
 473 coarse particles for all seasons excepted for WS2016 with values for coarse particles only.

474 During the wet season, coarse dust concentrations range from 5 to 25 $\mu\text{g}\cdot\text{m}^{-3}$ in 2015 and 9 to
 475 37 $\mu\text{g}\cdot\text{m}^{-3}$ in 2016, with higher values at the CT and ADF sites in 2015 and at AT, CT and ADF

476 in 2016. In WS2015, fine dust concentrations range from 12 to 49 with maximum values at
477 ADF and CT sites also. During the dry season, values range from 38 to 156 $\mu\text{g}\cdot\text{m}^{-3}$ in 2016 and
478 from 41 to 116 $\mu\text{g}\cdot\text{m}^{-3}$ in 2017, with maximum concentrations obtained at the CT site, followed
479 by AWB site. When considering mean values of the dry seasons, total dust at CT is 2.4 times
480 the values found at AT, 1.6 times at AWB and 3.4 times at ADF. Seasonal comparison shows
481 that total dust concentration is higher in the dry seasons than in WS2015 by a factor of 3 in AT,
482 2.6 in CT and 4 in AWB, but of the same order of magnitude in ADF site.

483 **3.6. Aerosol chemical closure**

484 The aerosol chemical closure obtained using the Guinot et al. (2007) method (see below)
485 at the different sites for each season is presented in Figure 13. Results show clear intra- and
486 inter-annual variations at all of the sites, as well as significant differences among the sites. In
487 total, dust accounts for 39 to 75% of the bulk PM mass at both traffic sites, with no clear
488 seasonal cycle and higher contributions in Cotonou (Figures 13c and 13d). These percentages
489 vary from 32 to 64% at the AWB site, and from 18 to 35% at the ADF site, with percentages
490 1.8 times higher in the dry season than in the wet season in AWB and no clear seasonal
491 difference in ADF (Figures 13a and 13b). Carbonaceous aerosol, the sum of EC and POM,
492 show large contributions at the ADF site (from 49 to 69% of the total PM mass), with relatively
493 similar proportions in each season (Figure 13a). The absence of a clear seasonal pattern is also
494 observed in CT whereas carbonaceous aerosol is slightly higher in WS than in DS in AWB (23
495 and 16% respectively) and AT (37 and 21% respectively) (Figures 13b-d). Carbonaceous
496 aerosol contribution accounts for about 11- 49% of the total mass at both traffic sites with higher
497 values in AT (mean of 30%) than in CT (13%). The ion percentages in PM fractions present
498 the same pattern at AT, CT and AWB sites with higher values in wet than in dry seasons. In
499 these sites, we may notice that mass concentration in coarse particles is larger in the wet season
500 whereas of the same order of magnitude than the one in fine particles in the dry season. In ADF,
501 no marked difference may be found between the seasons and the sizes (Figure 13).

502

503 **4. Discussion**

504 A discussion of the results site by site (Abidjan domestic fire site, traffic sites both together and
505 waste burning site) will be first proposed. We will scrutinize (1) the proximity between the sites
506 and the sources; (2) the source specificity with more or less incomplete combustion (e.g. wood
507 combustion and two-wheel vehicle emission factors are higher than gasoline emission factors

508 (Keita et al., 2018); (3) the relative influence of other local sources or transported sources to
509 the studied sites such as dust and biomass burning; (4) the occurrence of continental air masses;
510 (5) the variation of the boundary layer height (as reported by Colette et al., 2007); and (6) the
511 meteorological parameters (e.g. temperature, relative humidity and wet deposition) to explain
512 the differences of pollutant concentrations and their seasonal and inter-annual variabilities.
513 In a second part, we will present comparison of our values with other DACCIWA values and
514 also with literature values for other intensive campaigns in Africa.

515

516 **4.1. Abidjan Domestic Fires (ADF)**

517 As shown in the above paragraphs, maximum values are obtained at the ADF site, for aerosol
518 mass, EC, OC, WSOC, water-soluble ionic species (e.g. Cl^- , NO_3^- , Ca^{2+} and K^+) and some trace
519 elements such as Mg and K (whereas Al, Na and Fe are lower than in the other sites). Also,
520 aerosol PM_{2.5} values are well above the annual and daily WHO guidelines of 25 and 10 $\mu\text{g}\cdot\text{m}^{-3}$
521 respectively, whatever the season.

522 Such pattern is due to the proximity of the ADF site to the studied combustion source: in that
523 area, the use of wood combustion is very active due to commercial activities of women drying
524 fish and meat and domestic cooking. This is also confirmed by the high relative importance of
525 total carbon in aerosol mass whatever the size (49 to 69%) and by values of source enrichment
526 factor. Indeed, at least 30% of trace element concentrations are of anthropogenic origin at ADF
527 site. In addition, wood combustion is well known to be highly polluting due to incomplete
528 combustion: this is shown here by the measurements of very high OC/EC ratios at ADF, on the
529 order of the one measured at the source level by Keita et al. (2018). This is also shown by
530 WSOC relative importance which is expected for wood burning following Yu et al. (2018),
531 Tang et al. (2016), Feng (2006) and Saxena and Hildemann (1996) and by the strong correlation
532 of WSOC with biomass burning K^+ tracer.

533 Chloride is most likely associated with sea salt origin (normal chloride concentrations represent
534 at least 55% of marine aerosols following Goldberg (1963)) or secondary aerosol production
535 (Li et al., 2016). Since chloride relative concentration at the ADF site remains lower than the
536 that of sea-sal aerosols, the secondary production source would be the better explanation for
537 high chloride concentrations observed at ADF. The size distributions of Cl^- , K^+ , NH_4^+ and SO_4^{2-}
538 support this conclusion : the predominance of these elements in fine particle mode at the ADF
539 site would be associated with anthropogenic emissions, particularly biomass combustion and
540 domestic fires, or with secondary inorganic aerosols origin. This is confirmed by Cl^-/Na^+ ratio
541 values as shown earlier. Contrarily, Ca^{2+} and NO_3^- contributions to the total ions at the ADF

542 site peak mainly in the large particle fraction and may be attributed to quasi natural origin,
543 primarily to dust emissions and nitrate formation by reaction processes, respectively. In
544 addition, Na^+ and Mg^{2+} display similar size distributions at the ADF site, with the major
545 contribution in the coarse particle fraction, suggesting the common sea salt origin of these two
546 elements (Belis et al., 2013).

547 As we have shown above, the lower proportion of metal elements at the ADF site (6.5% of the
548 bulk concentration) can be explained by the less dominant influence of re-suspended dust
549 particles compared to traffic sources. Elements such as Cr, Mn, Pb and Ni have less seasonal
550 variability than other metallic elements. These small proportions of these non-crust elements
551 suggest a low contribution of elements emitted mainly by anthropogenic activities such as
552 industrial processes (Viana et al., 2007 and 2008; Minguillón et al., 2014). Finally, the Zn/Cd
553 ratio has been also examined. A value of 29 close to ratio reported for gasoline vehicle (27, Qin
554 et al., 1997) is obtained for the ADF site, indicating that this site is also impacted by traffic
555 sources.

556 High values of WSOC/OC ratios are expected to be harmful to health (Ramgolam et al., 2009,
557 Val et al., 2013). This effect is being enhanced by the particulate size measured at this site (Kim
558 et al., 2003; Wilson et al., 2002). Indeed, the relative mass distribution of PM and OC particle
559 sizes shows a major contribution of particles less than $1\ \mu\text{m}$ (as high as 85% of PM). This could
560 be due to the fact that carbonaceous aerosols are formed near emission sources and are mainly
561 of submicron size (Boucher, 2012). Nevertheless, note that EC also presents large coarse
562 particle contribution. This could be due to the importance of wood burning at this site with less
563 efficient combustion and large particle emissions (Watson et al., 2011).

564 In terms of seasonality, higher concentrations of aerosol mass, OC, WSOC, EC and total water
565 soluble ionic species (SO_4^{2-} , NH_4^+ and NO_3^-) are observed in WS2015 and WS2016 than in
566 DS2016. This may be explained by a more incomplete combustion in the wet seasons than in
567 DS2016 due to the use of moist wood for cooking and smoking fish, which leads to large
568 amount of smoke and higher particulate emission factor values. Note that DS2017 values are
569 as important as the ones of wet seasons, which will be explained later in the text. With regard
570 to WSOC, their variabilities may be also linked to meteorological factors, such as solar radiation
571 (Tang et al., 2016; Favez et al., 2008) and relative humidity (Liang et al., 2016). At ADF site,
572 temperatures are roughly similar in both seasons. However, RH variability may play a role since
573 it is higher in wet season than in dry season. Finally, our results indicate no clear seasonal cycle
574 for Cl^- , which confirms its anthropogenic origin, as previously shown.

575

576 **4.2. Traffic sites (Abidjan traffic and Cotonou Traffic sites)**

577 Let us recall first that the two traffic sites have been chosen since they are representative of the
578 traffic diversity in West Africa. At CT site, both personal cars, taxis and an important two-
579 wheel fleet may be found whereas at AT site, there are buses, taxis and personal cars. Also, the
580 distance between the site and the traffic sources is the same for the two traffic sites, slightly
581 larger than the distance between the site and the wood burning sources at ADF site.

582 In these two sites, concentrations are high with PM_{2.5} values well above the WHO guidelines.
583 Average aerosol mass, EC, OC, dust and water soluble ionic concentrations (with NO₃⁻ and
584 Ca²⁺ maximum at AT and CT sites respectively) are higher at CT than at AT site by a factor of
585 1.5 to 2. Note that this poor air quality found in Cotonou has been reported by Cachon et al.
586 (2014). The higher values found in Cotonou could be due to more intense traffic in Cotonou
587 than in Abidjan. Also in Cotonou, this traffic is associated with the lack of public transportation
588 and the use of highly polluted mopeds (aged over 15 years) (Gounougbe, 1999; Avogbe et al.,
589 2011), despite the effort in the last 10 years to restrict their use. Several studies such as MMEH
590 (2002) have shown that more than 94,000 mopeds and 350,000 second-hand vehicles are in
591 circulation in Cotonou. Other factors contributing to the local pollution include outdoor
592 restaurants using charcoal and motorcycle garages, which are more present around the Cotonou
593 traffic site compared to Abidjan site. It also includes anthropogenic dust. Indeed, at Cotonou,
594 the lack of road infrastructure favours the resuspension of dust particles. Finally, other sources
595 may potentially influence aerosol seasonal composition in these two sites, including marine
596 aerosols, transported dust and biomass burning particles as well as anthropogenic aerosols from
597 the surrounding countries (Figure 5). Note also that source enrichment factor values show that
598 about 17% of trace element concentrations are of anthropogenic origin at both traffic sites and
599 that the relative importance of total carbon in mass is higher at AT than at CT sites.

600 Aerosol mass, composition and size depend on the season and the two traffic sites are differently
601 affected. The EC and OC concentrations measured in both traffic sites and averaged per season
602 are higher in dry than in wet season. Such variations may be explained by several factors:
603 particulate wet deposition occurring during the wet season, reduction of traffic flow due to
604 school vacations and meteorological influence. Higher EC and OC concentrations are obtained
605 at CT than at AT sites in dry seasons whereas no statistical difference may be found between
606 the two sites in wet seasons. Such a result is mainly explained by figures 5. In wet seasons,
607 similar backtrajectory pattern may be observed for both sites whereas in dry seasons, CT traffic
608 site only would be influenced by Nigerian anthropogenic sources.

609 In terms of WSOC concentrations, concentrations at the AT site are on average higher than
610 those recorded at the CT site in the wet season, but lower in dry season. The presence of dust
611 can produce semi-volatile organic gas scavenging and therefore WSOC and OC enhancement.
612 Such a phenomenon can explain the highest WSOC concentrations observed in dry season at
613 the CT site where dust concentrations are highest (see dust paragraph). Moreover, this can also
614 explain why the maximum WSOC are in coarse particles at CT, while at AT maximum values
615 are in ultra-fine particles.

616 Total WSI concentrations are larger at AT site in the wet than in the dry season with higher
617 values in coarse particles. At CT site, total WSI concentrations in fine particles are higher in
618 the dry than in the wet season whereas same values are obtained in coarse particles for both
619 seasons. Note that CT values are generally higher than AT values with a more important
620 contribution of fine particles in the dry season. These WSI variations can be explained by the
621 relative importance of Ca^{2+} , SO_4^{2-} and NO_3^- in both sites.

622 First, Ca^{2+} contribution to total WSI is higher in CT site than in AT site with no clear seasonal
623 variation at CT site and higher values in dry season than in wet season at AT site. Also at CT,
624 fine and coarse Ca^{2+} particles are in the same range, whereas coarse Ca^{2+} particles are
625 predominant at AT site. Such feature may be explained by the impact of dust sources including
626 long-range dust transport at Abidjan and a combination of long-range dust transport and road
627 resuspension at Cotonou.

628 Second, the relative contribution of SO_4^{2-} , NH_4^+ and NO_3^- as a percentage of total WSI in the
629 different particle modes is reduced in the wet season. During the wet season, the clean winds
630 surrounding the ocean before reaching the measurement sites could contribute to lower the
631 proportion of these species, in addition to the scavenging processes during the rainy days.
632 Unlike the wet season, a relatively good correlation with r^2 of 0.87 (SO_4^{2-} versus NH_4^+), 0.73
633 (NO_3^- versus NH_4^+) and 0.87 (SO_4^{2-} versus NO_3^-) has been found in coarse particles, indicating
634 similar sources for these three species during the dry season. In order to try to identify these
635 sources, the ratio of $\text{SO}_4^{2-}/\text{Ca}^{2+}$ and $\text{NO}_3^-/\text{Ca}^{2+}$ has been determined. The average $\text{SO}_4^{2-}/\text{Ca}^{2+}$
636 and $\text{NO}_3^-/\text{Ca}^{2+}$ ratios in combined coarse particles (1.07 and 2.58 during the wet season and 0.33
637 and 1.60 during the dry season) are higher than the corresponding ratios for typical soil (0.026
638 and 0.003, respectively). On the other hand, the $\text{SO}_4^{2-}/\text{Ca}^{2+}$ ratio increases in the fine particles
639 (5.07 during the wet season and 2.53 during the dry season), while that of $\text{NO}_3^-/\text{Ca}^{2+}$ remains
640 almost constant (2.86 during the wet season and 1.65 during the dry season). This implies that
641 the atmosphere at AT and CT sites is enriched by SO_4^{2-} formed as anthropogenic secondary
642 particles, possibly from sulfur containing pollution sources (Seinfeld and Pandis, 1998),

643 particularly in fine particle mode, and by NO_3^- mostly coming from nitrogen containing sources
644 in all particle sizes. The higher contributions of these elements during the dry season could
645 result from a combination of several factors: 1) an atmosphere loaded with dust favoring
646 heterogeneous chemistry to obtain secondary aerosol and the rise of biomass burning emissions;
647 2) the increase of photochemical activity and higher concentrations of hydroxyl radicals in the
648 dry season, which can oxidize SO_2 from combustion (Arndt et al., 1997) to SO_4^{2-} (Li et al.,
649 2014) ; and 3) the wind transport of anthropogenic secondary particles from the industrial zone
650 located upstream from our sites. Finally, the proportion of Cl^- relative to the total mass of ions
651 is highest for coarse particles at both traffic sites especially during the wet season, suggesting
652 that Cl^- at AT and CT sites is from natural origin and probably from sea salt emissions.

653 If we focus now on dust during the two wet seasons, concentrations are higher in 2016 than in
654 2015 at CT and AT sites for coarse particles (no data of fine particles are available in WS2016).
655 This is consistent with observed aerosol optical depth (AOD) values at CT, which increased by
656 a factor of 2 between 2015 and 2016. No AOD value is given by Léon et al. (2019) at Abidjan
657 in WS2015 to allow such comparison in Abidjan. Moreover, during the wet season, an
658 Angström coefficient (AE) on the order of 1 has been found at CT site, indicating smaller
659 particles that could be due to road resuspension. It is interesting to note that during WS2016,
660 AOD and AE are respectively higher and lower at Abidjan than at Cotonou. Again, this is
661 consistent with our dust concentrations at CT site. In Abidjan, we could assume that another
662 source of Ca^{2+} , which is not taken into account in our dust calculations, may explain our dust
663 concentration data. That may be the result of anthropogenic Ca^{2+} emissions from residential
664 combustion, more important in 2016 than in 2015 as shown earlier
665 (http://naei.beis.gov.uk/overview/pollutants?pollutant_id=84).

666 The relative contribution of dust generally peaks in the coarse mode and, to a lesser extent, in
667 the fine mode, reflecting their natural origin. It is interesting to note that the dust contribution
668 observed in this study for the year 2016 at the Abidjan site is in agreement with the results of
669 Xu et al. (2019) which show a $\text{PM}_{2.5}$ dust contribution of 35-50% compared to our values of
670 18-52%.

671

672 **4.3. Abidjan Waste Burning site**

673 Concentrations measured at AWB site are slightly lower than values found in the other sites.
674 This can be explained by the larger distance of the site to the main studied source (here waste
675 burning source) than in the other sites. However, $\text{PM}_{2.5}$ values are also higher than WHO
676 guidelines. +

677 Aerosol mass, EC and OC concentrations are higher in dry than in wet season, which suggests
678 less waste burning activities during the wet season or impacts of other local anthropogenic
679 sources or long-range biomass burning sources. Highest values are found in DS2017 with the
680 lowest OC/EC ratio, as at AT site. OC/EC ratio is highly variable at AWB (1-10) which
681 confirms that AWB site may be impacted by different types of sources as well as by secondary
682 aerosol organic formation which can be detected for OC/EC higher than 2 (Turpin et al. 1990;
683 Hildermann et al. 1991; Chow et al. 1996). Note that OC/EC typical for waste burning source
684 is of the order of 8 (Keita et al., 2018).

685 It is also observed that at the AWB site, PM mass concentrations are mainly distributed in C
686 mode (30-44%) over the entire period of study, excepted during the WS2015, and to a lesser
687 extend in F mode (21-44%). EC and OC being mainly distributed in C and UF modes. Water-
688 soluble fraction of organic carbon is important (32%) and on the order of the one found at AT
689 site. Same for WSI concentrations and WSI composition. At AWB, WSI values are globally
690 slightly higher in wet than in dry season. However, it is interesting to underline that Ca^{2+} is
691 much higher in dry season than in wet season, especially in DS2017. This is in agreement with
692 dust concentrations and trace element concentrations, which have been found to be maximum
693 at AWB, reaching 35.8% of the total PM mass in the dry season. These maximum percentages
694 are due to the large contribution of both Al and Na crustal elements which account for about
695 26%. Also note that Cu/Sb of 0.08 in DS2017, which indicate an influence of re-suspended
696 particles. A Zn/Cd value of 56 is obtained for the AWB site which is in close agreement with
697 values reported for oil burning (Watson et al., 2001, Samara et al., 2003). That could indicate
698 that oil might be one of the waste burning materials.

699 Our result suggests that AWB aerosol mass is influenced by a mix of sources, including fuel
700 combustion and mineral salt from sources around the measurement site, associated to long-
701 range source impact of dust and biomass burning which will be further discussed in the next
702 paragraph.

703

704 **4.4. Interannual variability of aerosols in Abidjan and Cotonou**

705 EC and OC concentrations are generally higher in DS2017 than in DS2016 for all the sites. This
706 is not due to the meteorological condition, which is similar in both years. This is also not due
707 to biomass burning impacts. Indeed, when looking at MODIS burnt areas for our period of study
708 (<http://www.aeris-data.fr/redirect/MODIS-MCD64A1>), burnt areas of west African savannas
709 are higher in 2016 than in 2017. Therefore, carbonaceous aerosol concentrations should be
710 higher in 2016. Then, this could be due to a counter effect between biomass burning emission

711 strength and air mass transport efficiency. As a result, biomass burning impact could not explain
712 the difference in EC and OC during the dry season between 2016 and 2017. Rather, this is due
713 to the variability of local sources. In DS2016 in Abidjan, there was a general strike of civil
714 servants of the State with important consequences on urban activities. Lower activities were
715 observed (lower fish smoking emissions, lower traffic ..) in DS2016 compared to DS2017, thus
716 explaining the lower EC and OC concentrations at Abidjan sites. In Cotonou, highest
717 carbonaceous aerosol values in DS2017 may be explained by backtrajectory patterns: Cotonou
718 would be impacted by air masses coming from the high polluted Lagos (Nigeria) area in that
719 period whereas from less polluted northern areas in DS2016. Such an assumption is validated
720 by the AOD values at 550nm from MODIS satellite images ([http://www.aeris-](http://www.aeris-data.fr/redirect/MODIS-MCD64A1)
721 [data.fr/redirect/MODIS-MCD64A1](http://www.aeris-data.fr/redirect/MODIS-MCD64A1)), which show very high particulate concentrations in the
722 Guinean Golf (Figure 14).

723 This figure also shows the AOD difference between Cotonou and Abidjan for DS2017, with
724 higher values at Cotonou than in Abidjan for the campaign period, in agreement with our
725 measurements of aerosol mass, EC, OC and dust. This is confirmed by the DACCIWA
726 sunphotometer AOD and Angström coefficient (AE) measurements at Abidjan and Cotonou
727 (Léon et al., 2019; Djossou et al., 2018). Indeed, in DS2017, during our period of
728 measurements, mean AOD in Cotonou is of the order of 1.3 versus 0.9 in Abidjan for an AE of
729 0.6 for both sites, which clearly indicates the presence of coarse dust particles.

730 Finally, aerosol mass and dust concentrations have been seen to be higher in DS2016 than in
731 DS2017 in Abidjan whereas values are on the same order of magnitude at Cotonou. Such high
732 values at Abidjan in DS2016 can be explained by the back-trajectory pattern with air-masses
733 all coming from northern dusty areas in DS2016 (Bodélé depression in Tchad, Prospero et al.
734 (2002), Washington et al., (2003), Knippertz et al. (2011), Balarabe et al., (2016)) and/or from
735 northern dusty countries (Mali, Niger) (Ozer, 2005), whereas in DS2017, contribution of
736 southern marine clean air-masses may also be noted.

737 In the wet season, aerosol mass, EC and OC are higher in WS2015 than in WS2016. This may
738 be due to particulate wet deposition, more efficient in WS2016 which have been seen earlier to
739 be more rainy (4.7mm) than in WS2015 (2mm). Moreover, at AT site, dust concentrations are
740 higher for coarse particles in WS2016 than in WS2015. Such variations may be explained by
741 long-range dust sources and/or road dust resuspension processes. As no dust event has been
742 noticed, local source explanation seems to be more evident.

743 In AT, CT and AWB, OC/EC ratios are globally on the same order for WS2015, WS2016 and
744 DS2016, with values lower than for DS2017. This could be due to lower traffic activities linked

745 to the DS2016 strike and the wet season vacation periods. Indeed, much higher OC/EC ratios
746 measured in DS2017 are typical of those of diesel vehicles (Mmari et al., 2013; Keita et al.,
747 2018). Finally, it is interesting to note that OC/EC ratios measured in this study are in the range
748 of those previously reported for other megacities such as Agra in India with 6.7 (Pachauri et al.,
749 2013), Helsinki in Finland with 2.7 (Viidanoja, 2002), Cairo in Egypt with 2.9 (Favez, 2008),
750 Paris in France with 3.5 (Favez, 2008), and Milan in Italy with 6.6 (Lonati et al., 2007).

751

752 **4.5. Comparison with literature data**

753 Firstly, the comparison between our data and other DACCIWA results including other time
754 sampling focuses on PM_{2.5} levels, since these particle sizes are relevant for health impact
755 studies (Xing et al., 2016). In addition to our values, Figure 7 presents data from Xu et al. (2019)
756 using personal samplers collected in the same area and at the same dates in 2016 during 12h on
757 women at the ADF site, students at the AWB site and drivers at the CT site, and from Djossou
758 et al. (2018) study based on one week exposed filters collected at the same areas and for the
759 same periods as this study. We note that PM_{2.5} directly measured on women are 2.3 and 0.9
760 times our values obtained at the ADF site in dry and wet seasons, respectively, and 3.4 and 4.9
761 times higher on students than at the AWB site, and 1.6 and 2.1 times higher on drivers than at
762 the CT site. Also, our values are on average 1.6, 3, 5 and 8 times higher than weekly-integrated
763 values of Djossou et al. (2018) including our 3 days of measurements at the AWB, ADF, AT
764 and CT sites respectively. As it may be seen, the lowest concentrations are observed in Djossou
765 et al. (2018), whereas the highest concentrations are recorded in Xu. et al. (2019). This is valid
766 for all sites, seasons and campaigns. Differences between our values and Djossou values may
767 be explained by the sampling times of the two studies. Indeed, as recalled, Djossou
768 measurements are weekly integrated, taking into account diurnal activities during all the week,
769 including week-end and nights which have expected lower PM_{2.5} concentrations. Our study
770 includes only maximum pollution conditions for each site. The highest differences occur for
771 the traffic sites. This may be clearly understood since diurnal and weekly variations of traffic
772 sources are the most variable. Comparison between our values and Xu et al. (2019) values is
773 also interesting. Indeed, it is at the ADF site that on-site and women PM_{2.5} concentrations are
774 the closest, which shows that this site is the most representative of the pollution exposure to
775 women. The biggest differences are found at the AWB site. As already mentioned, distance
776 from the site to the waste burning source is more important than for other sites, which explains
777 why concentrations obtained on students who are leaving close to the sources are much higher
778 than on-site concentrations. At the Cotonou traffic site, measurements taken from people are

779 also higher than on site measurements. Such differences can be explained by additional
780 pollution exposure as people move around. Note that the sampling technique may also play a
781 role in such a comparison. In terms of seasonal variation, our results are in agreement with long-
782 term EC measurements obtained by Djossou et al. (2018) for the same sites and period. Finally,
783 Table 4 compares our mean PM_{2.5} results obtained from 3-hour sampling for three consecutive
784 days to literature data for different traffic sites in the world given at a daily scale. It is interesting
785 to note that our values are situated at the higher end of the range of PM_{2.5} data observed from
786 the other sites.

787 Secondly, Table 5 compares our OC and EC values to those obtained by Djossou et al. (2018)
788 and Xu et al. (2019) as previously described for the same period and the same sites. Again, it is
789 interesting to note that Djossou's values are in general lower than ours. Indeed, for the wet and
790 dry seasons, our OC measurements are 4 and 1.4 times higher than Djossou's at the AT site,
791 2.1 and 5.7 times higher at the CT site, and 2.5 and 2.5 times higher at the ADF site,
792 respectively. As for PM_{2.5}, this can be explained by the different sampling times between our
793 experiments that were performed at the peak of urban activities, while Djossou's dataset
794 represents weekly integrated values. Differences at the ADF site are largely explained by the
795 temporal pattern of fish smoking activities which take place every day, only in the morning, as
796 such the associated pollution is not well represented in the weekly sampling. Finally, there are
797 less differences at the AWB site between both datasets. As above explained, there are no marked
798 temporal variations of concentrations at AWB site. The predominant waste burning emissions
799 impacting our site can occur night and day, week-days and week-end since origin of such
800 burning can be either anthropogenic or from spontaneous combustion. It may be also recalled
801 that another reason of agreement between both datasets may come from the large distance
802 between the site and the local and regional sources. Comparisons made between our values and
803 those of Xu's personal data show that both OC and EC are of the same order at the ADF site,
804 whereas Xu values are higher than ours at the CT and AWB sites. This result is in agreement
805 with what we found with PM_{2.5} concentrations as detailed above. Finally, Table 6 presents OC
806 and EC for the PM_{2.5} comparison between our values and other recent studies dealing with
807 traffic sites in other regions of the world and with similar operational conditions. We find that
808 our values are situated in the middle of the range observed in these different studies. Briefly, as
809 presented in Table 7, it is interesting to compare our WSOC concentrations to literature data
810 for different traffic sites of the world. We note that our values are on the same order as values
811 found in Asia and higher than those found in Europe.

812 Thirdly, the percentages of the total WSI to PM mass (15-20%) at the three Abidjan sites (ADF,
813 AWB and AT) are in the same order of magnitude than the data from PM_{2.5} personal exposure
814 samples collected at the same locations in 2016 by Xu et al. (2019). Our results also are very
815 close to the ionic contribution of 9% of the PM₁₀ mass found at the urban curbside site in Dar
816 es Salaam in Tanzania during the wet season 2005 by Mkoma (2008).

817

818 **5. Conclusion**

819 This paper presents the mass and the size-speciated chemical composition of particulate matter
820 (PM) obtained during the dry and wet seasons in 2015, 2016 and 2017. During each campaign,
821 3-hour sampling at the peak period of pollution for three consecutive days was performed at
822 three sites in Abidjan, representative of domestic fire (ADF), waste burning (AWB) and traffic
823 (AT) sources, and at one traffic site in Cotonou (CT).

824 It is important to underline that our results and their temporal variations are very sensitive to
825 (1) the source activities whose pollution levels are highly linked to socio-economic status of
826 each city; (2) the impact of imported pollution (sea-salt, biomass burning, dust, anthropogenic
827 emissions from neighboring countries), according to air mass origins; and (3) the particle wet
828 deposition.

829 The comparison between our results and other DACCIWA measurements underlines the
830 importance of the distance of the chosen site to the sources. At the source level (such as ADF),
831 pollution results at the site are in agreement with exposure of people living at this site. However,
832 at the other sites, comparison is more difficult since the sites are under the influence of a mix
833 of transported sources. That shows the key importance of exposure studies to estimate air
834 quality and health impacts. That shows also the need of long-term studies to really understand
835 role of imported sources in urban air quality.

836 The main striking feature is that PM_{2.5} values are well above the annual and daily WHO
837 guidelines of 25 and 10 $\mu\text{g}\cdot\text{m}^{-3}$, respectively, whatever the site and the season. Also, measured
838 concentrations from this study are situated in the middle to the high part of the range of
839 worldwide urban aerosol concentrations given at a daily scale. In addition, we have stressed the
840 importance of ultra-fine and fine particles in the studied aerosol and of species such as
841 particulate organic matter and water soluble organic carbon, which are well known to be
842 particularly harmful. This is again a warning signal for pollution levels in African capitals if
843 nothing is done to reduce emissions in the future.

844 Our study constitutes an original database to characterize urban air pollution from
845 specific African combustion sources. The next step will be to cross such an exhaustive aerosol

846 chemical characterization to biological data in order to evaluate the impact of aerosol size and
847 chemical composition on aerosol inflammatory properties.

848

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854

855 **Author Contributions**

856 J.A. and C.L. conceived and designed the study. J.A., C.L. and E.T.D. contributed to the
857 literature search, data analysis/interpretation and manuscript writing. J.A., C.L., A.B. and
858 E.T.D. contributed to manuscript revision. J.A., C.L., J.F.L, H.C, V.Y., A.A, C.G, C.Z, E.C and
859 S.K. carried out the particulate samples collection and chemical experiments, analyzed the
860 experimental data.

861

862 **Additional Information**

863 Figure S1 and Appendix A-D accompany this manuscript can be found in
864 SupplementaryInformation.

865

866 **Competing financial interests**

867 The authors declare no competing financial interests

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1251 Figure caption

1252 Figure 1: Map of the city of Abidjan reporting the geographical location of DACCIWA urban
1253 sampling sites.

- 1254 Figure 2: Map of the city of Cotonou reporting the geographical location of DACCIWA urban
1255 sampling site.
- 1256 Figure 3: Pictures of the different sampling sites: (a) Traffic in Cotonou (Benin, CT station),
1257 (b) Waste burning in Abidjan (Côte d'Ivoire, AWB station), (c), Domestic fire, showing
1258 smoking activity in Yopougon, Abidjan (Côte d'Ivoire, ADF station), (d) " woro-woro
1259 and Gbaka " traffic in Abidjan (Côte d'Ivoire, AT station).
- 1260 Figure 4: Wind, pressure and temperature diagram at Abidjan and Cotonou during the different
1261 campaigns.
- 1262 Figure 5: Back trajectories arriving at Abidjan (a) and Cotonou (b) for each season (WS2015,
1263 WS2016, DS2016 and DS2017).
- 1264 Figure 6: Aerosol Mass concentrations at the different study sites for each campaign and for the
1265 different sizes (C in black, Fine in light Grey, Ultra-fine in grey). Bulk aerosol mass is
1266 indicated in boxes.
- 1267 Figure 7: Comparison of PM_{2.5} mass concentrations in $\mu\text{g}\cdot\text{m}^{-3}$ at the four sites with those
1268 obtained by Djossou et al. (2018) and Xu et al. (2019) for the same sites and periods.
1269 Data for the following weeks were then selected in Djossou et al. (2018): 20-27/07/2015
1270 for WS15, 4-11/01/2016 for DS16, 4-11/07/2016 for WS16 and 9-16/01/2017 for DS17.
- 1271 Figure 8: EC relative concentrations in each size classes (C in black, Fine in light grey, Ultra-
1272 fine in grey) at the different study sites for each campaign. Bulk EC concentration for
1273 each site is indicated in boxes.
- 1274 Figure 9: OC relative concentrations in each size classes (C in black, Fine in light grey, Ultra-
1275 fine in grey) at the different study sites for each campaign. Bulk OC concentration for
1276 each site is indicated in boxes.
- 1277 Figure 10: OC/ EC ratio for the different campaigns and sites for each aerosol size (C in black,
1278 Fine in light grey, Ultra-fine in grey). Each box shows the median and the first and the
1279 third quartiles.
- 1280 Figure 11: Water-soluble ionic species speciation for each site, each campaign and each aerosol
1281 size.
- 1282 Figure 12: Dust concentrations at the different study sites for each campaign and for the
1283 different sizes (C in black, Fine in light grey, Ultra-fine in grey).

1284 Figure 13: Size-specified aerosol chemical composition for each site, for each campaign and
1285 each aerosol size.

1286 Figure 14: MODIS Aerosol optical depth regional distribution over West Africa. Data are for
1287 2017, focusing on our campaign date at Abidjan (a-c 01/11-12-left part) and Cotonou
1288 (b-d 01/6-7, right part).

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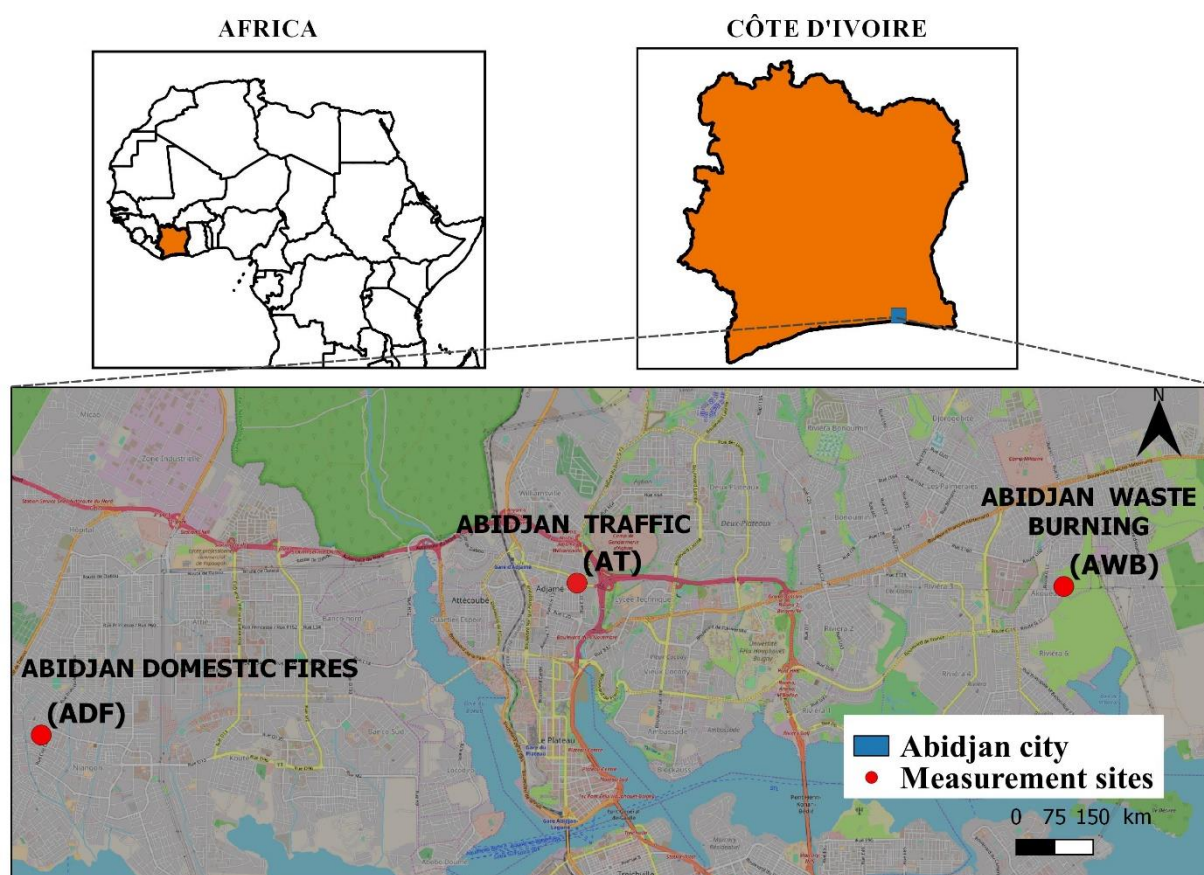
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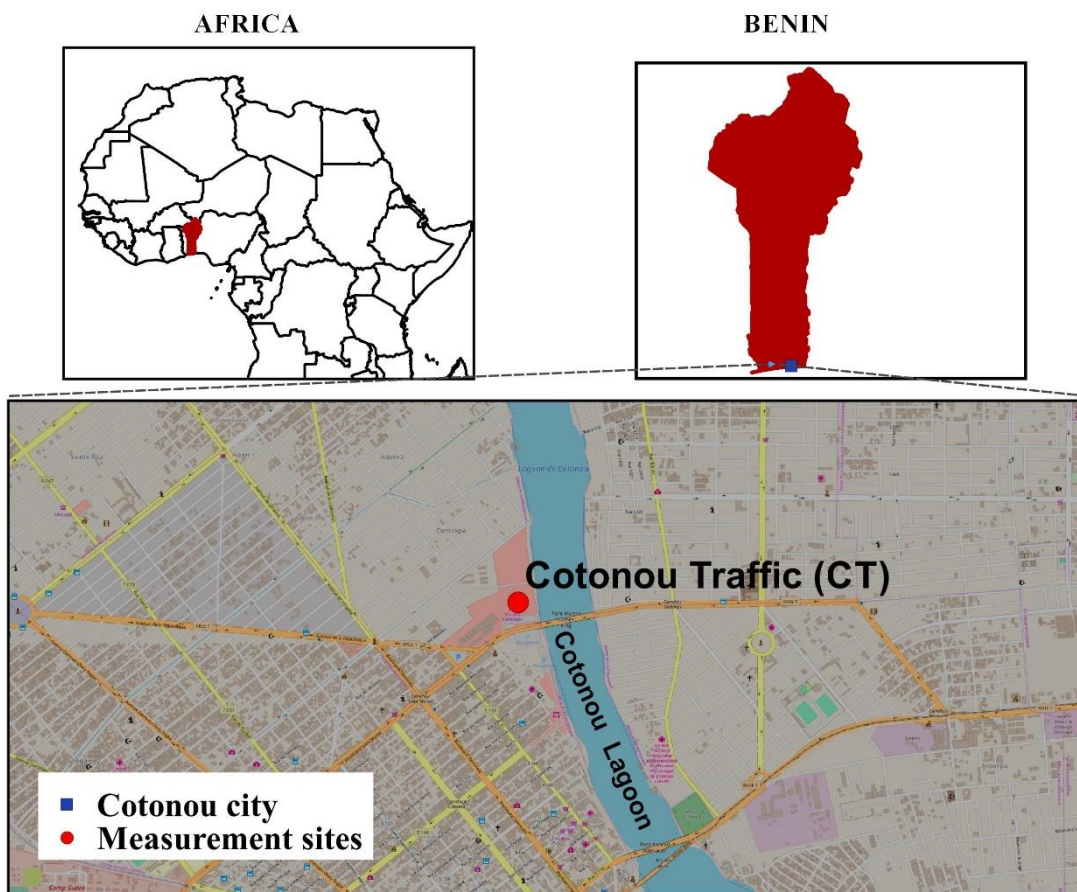
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Figure 1



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Figure 2



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1314 **Figure 3**

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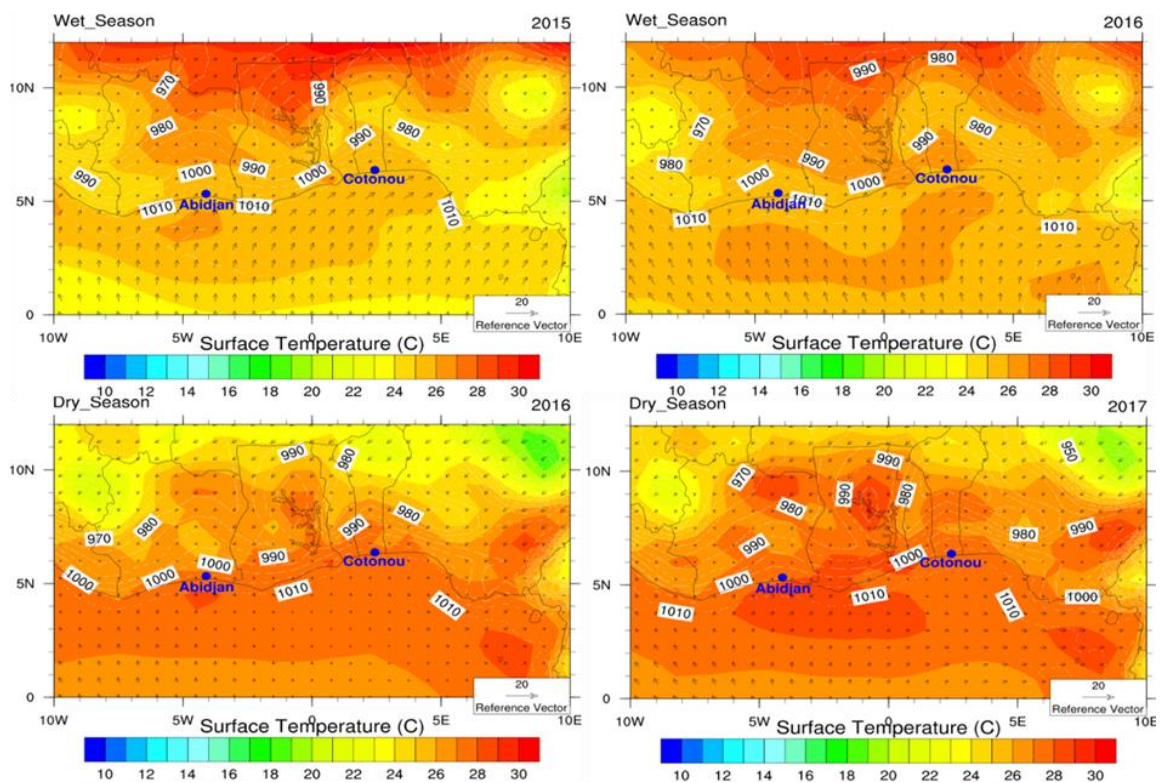
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1326 **Figure 4**

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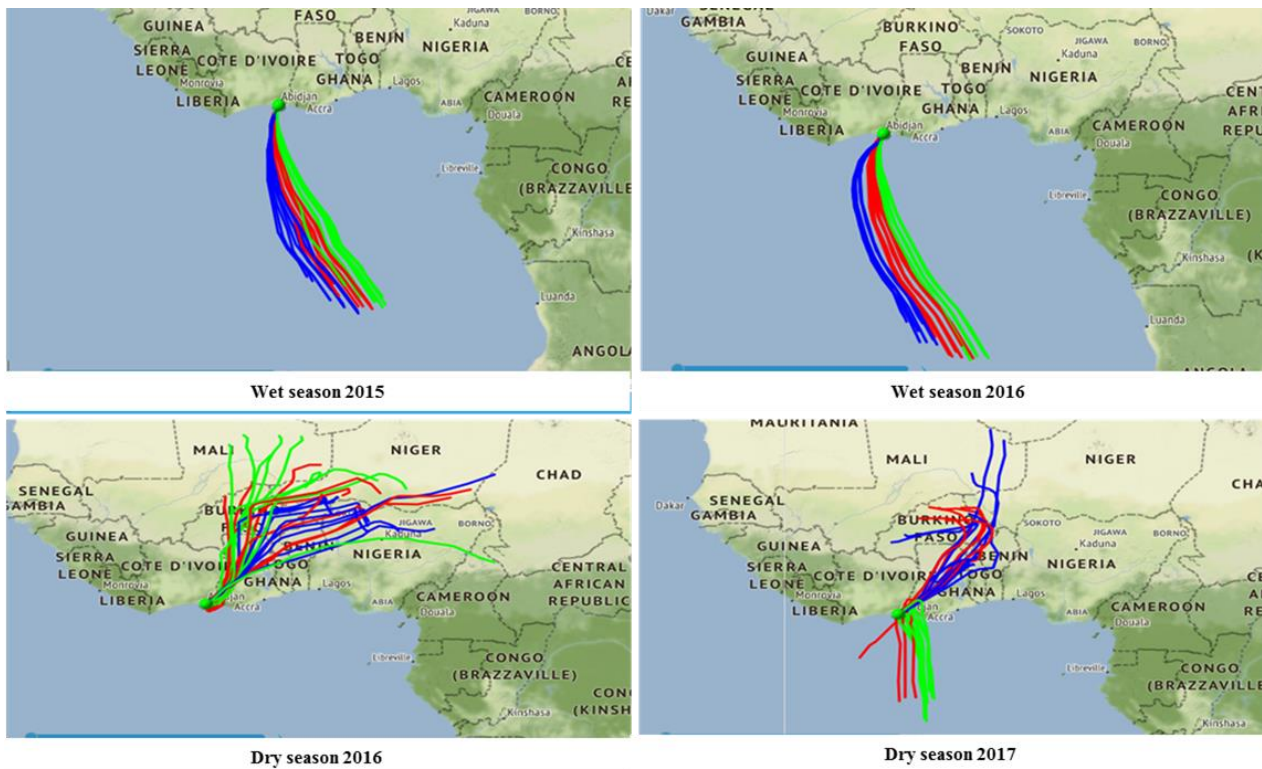
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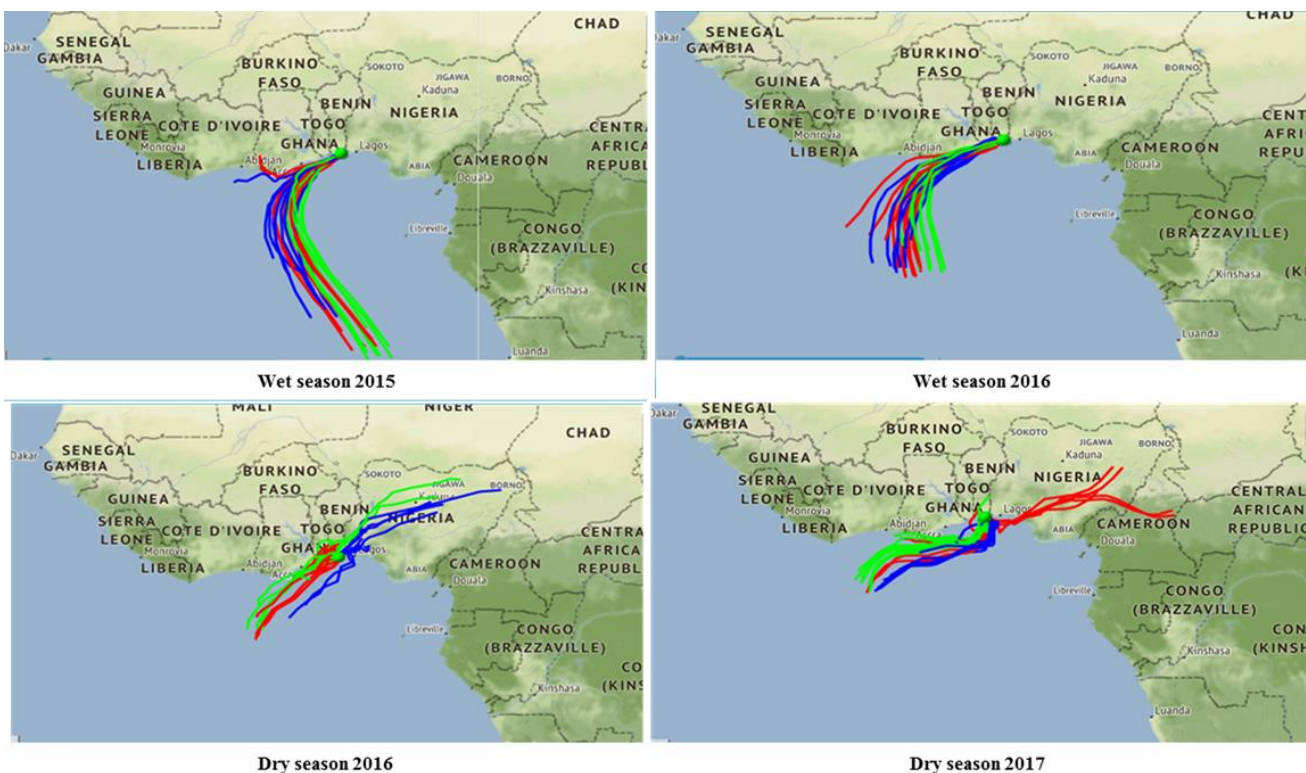
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1344 **Figure 5-a**



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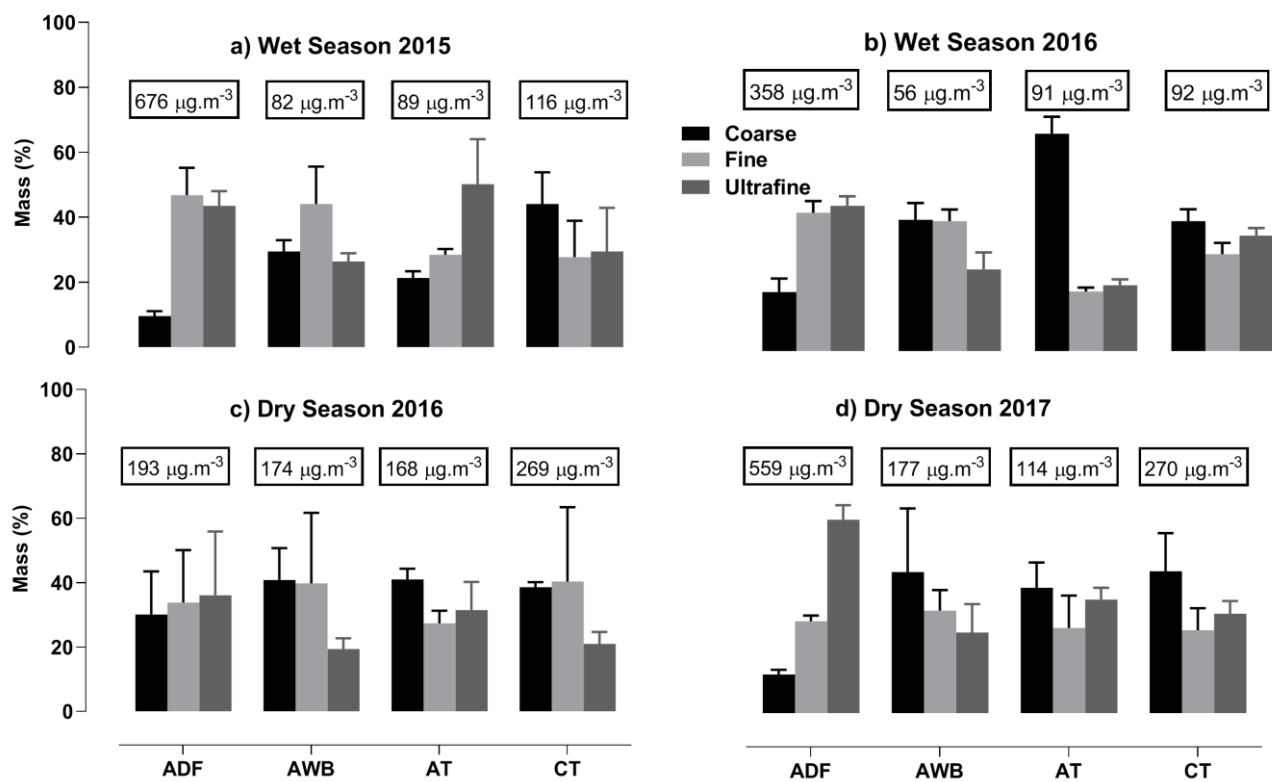
1346 **Figure 5-b**

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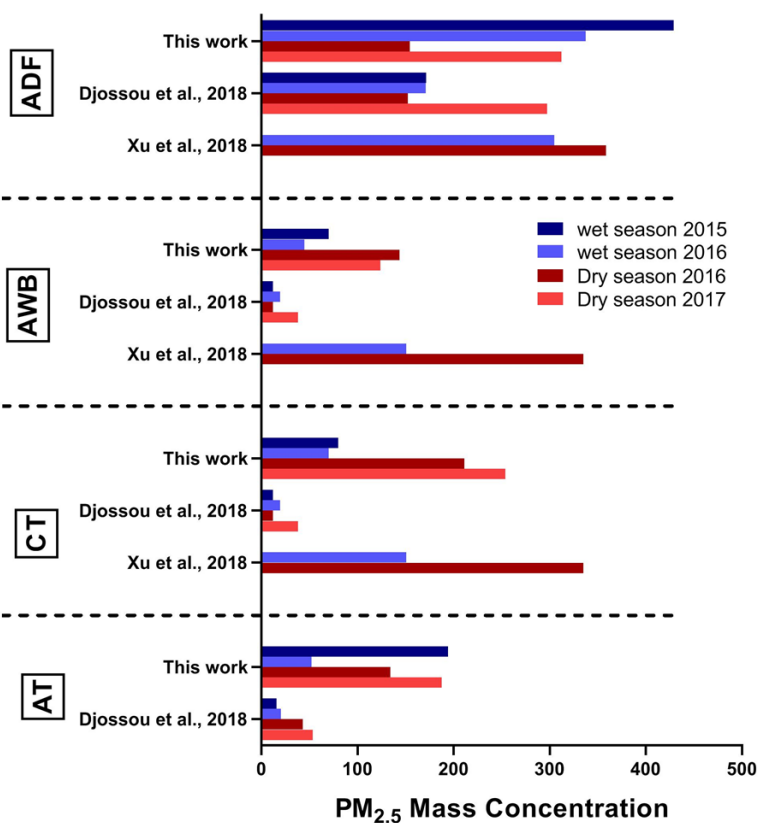


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1352 **Figure 6**

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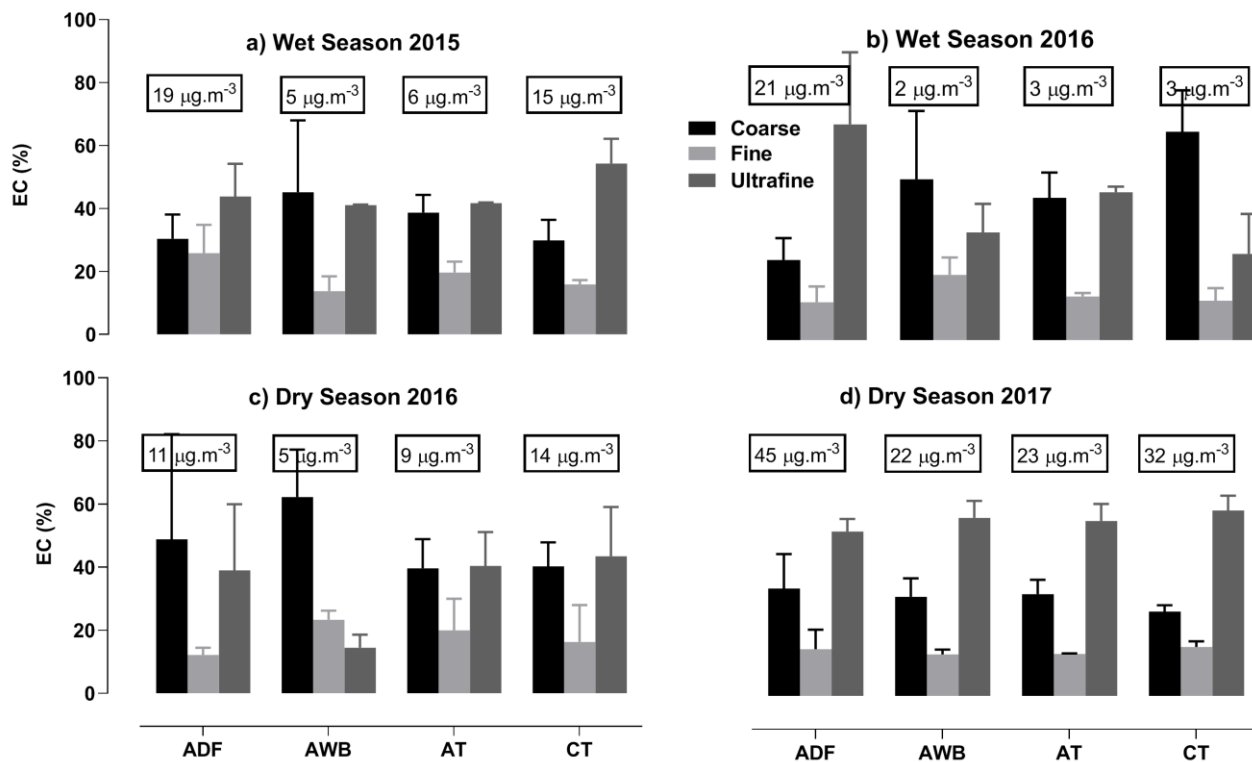
1356 **Figure 7**

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1362 **Figure 8**

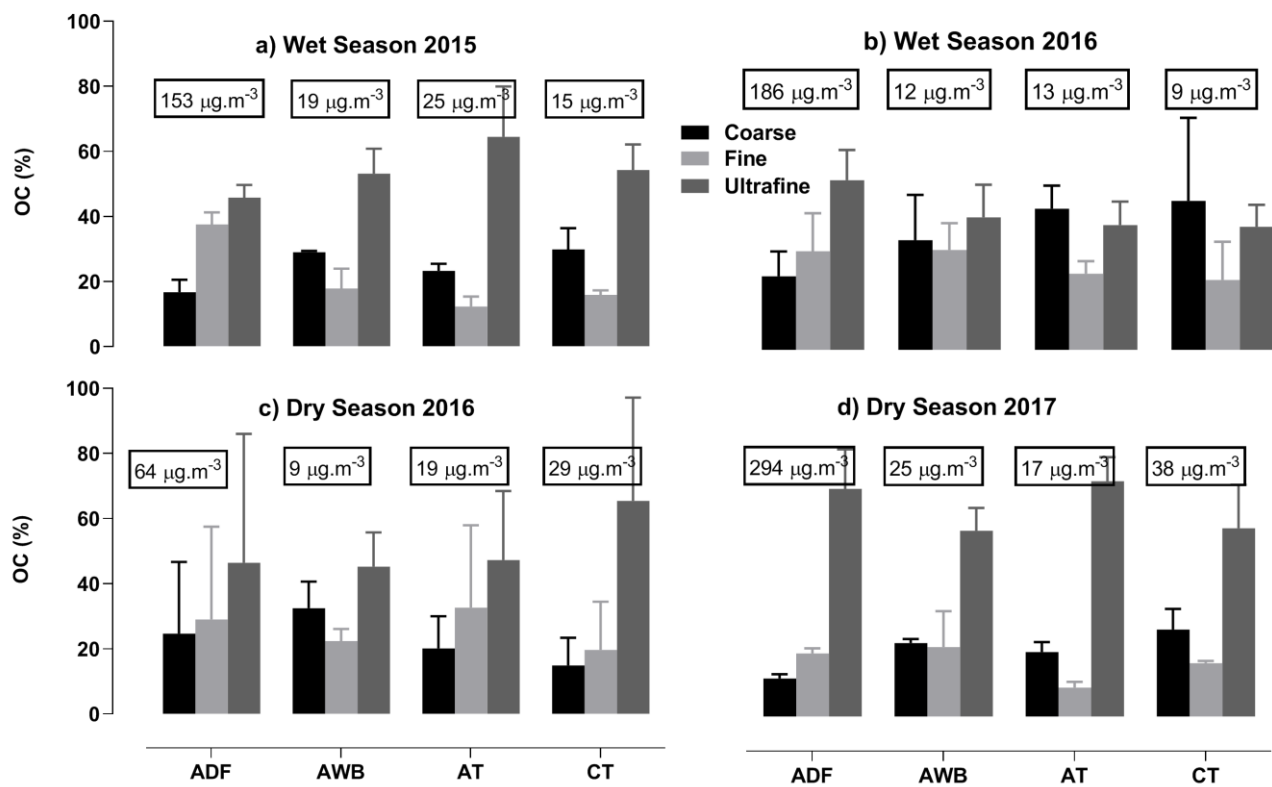
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1369 **Figure 9**

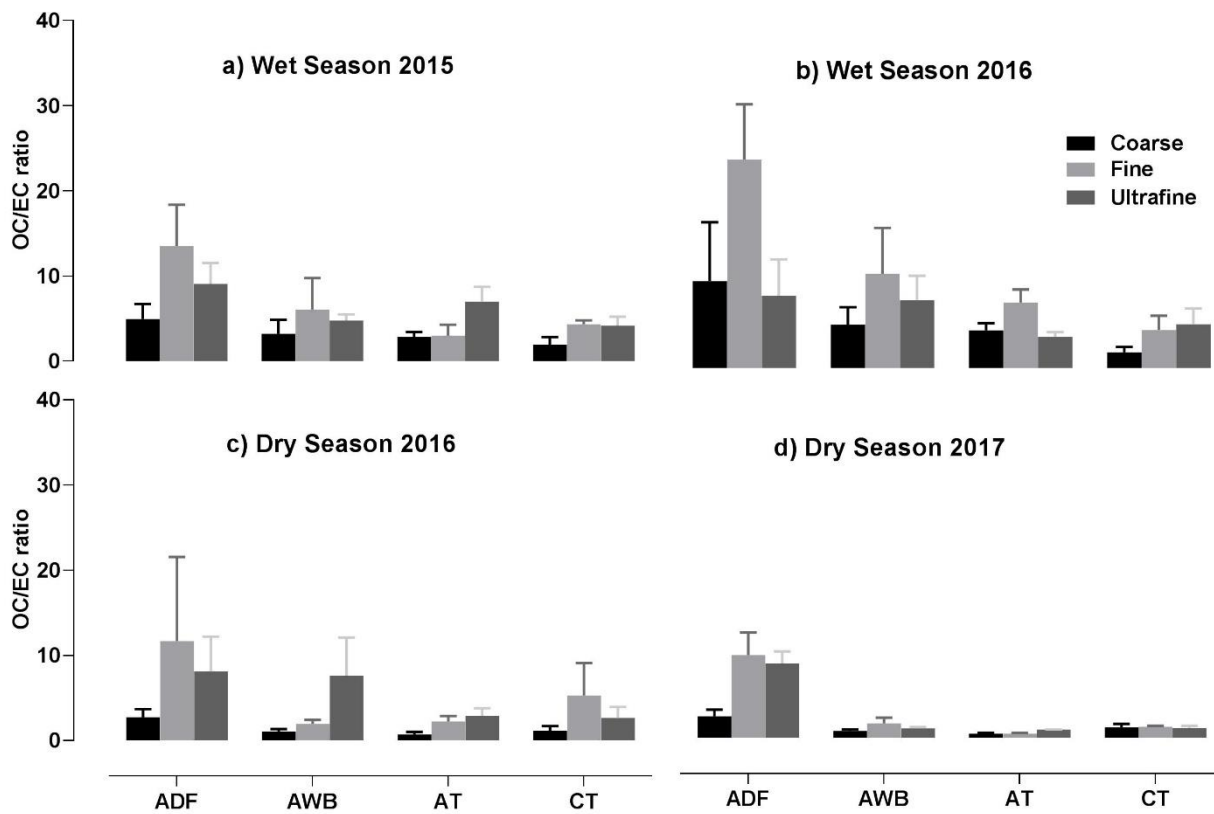
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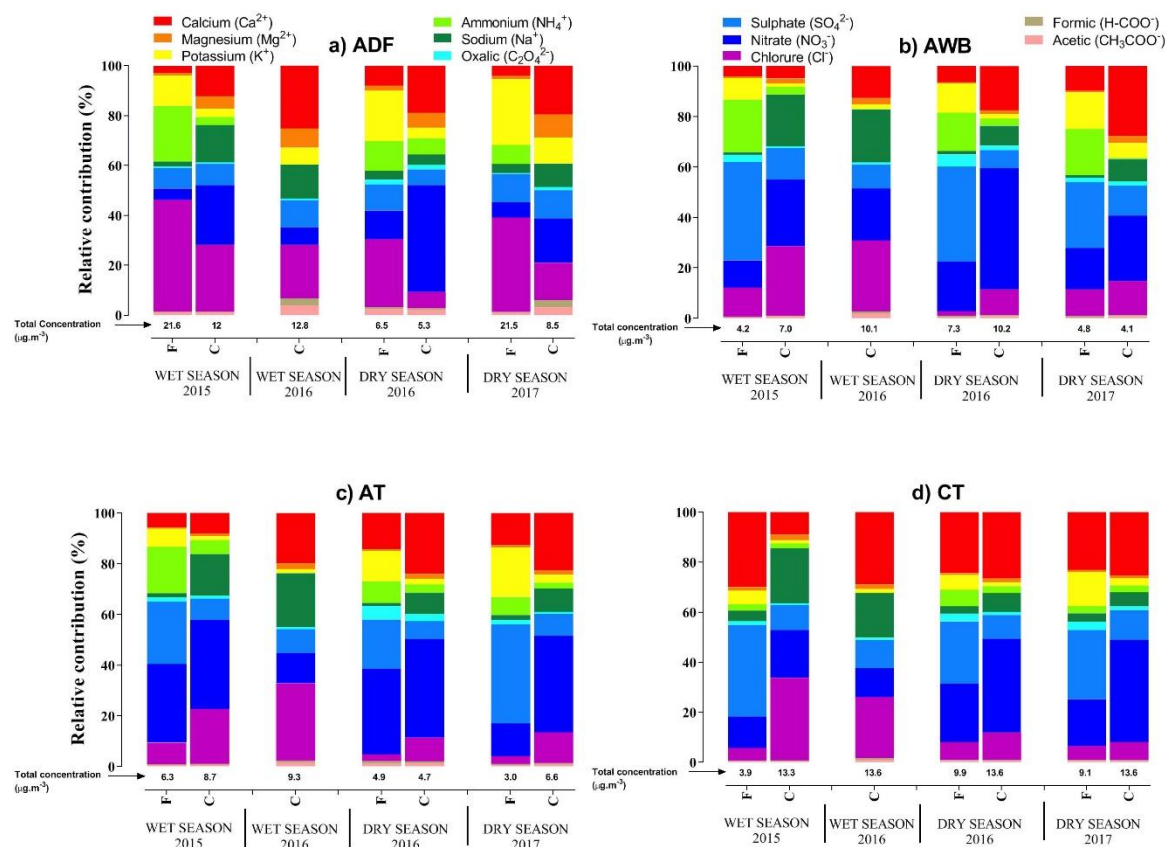


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1376 **Figure 10**

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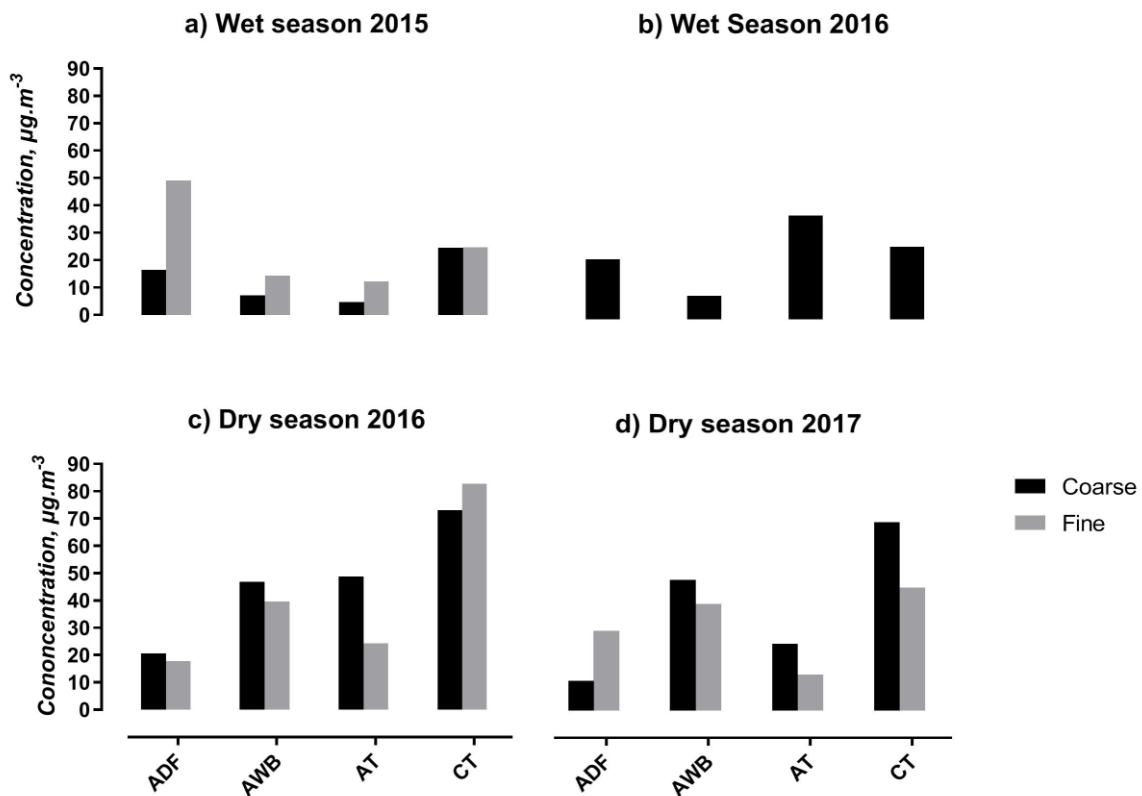


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1380 **Figure 11**

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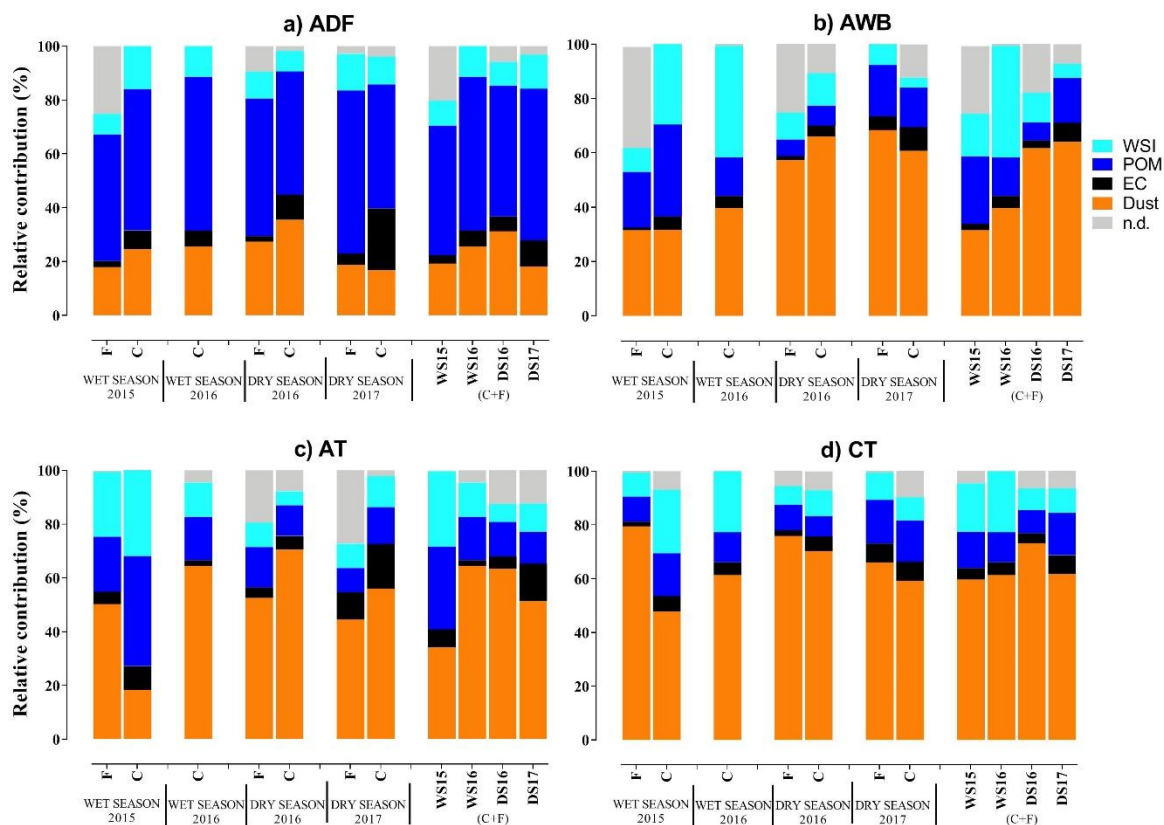
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1384 **Figure 12**

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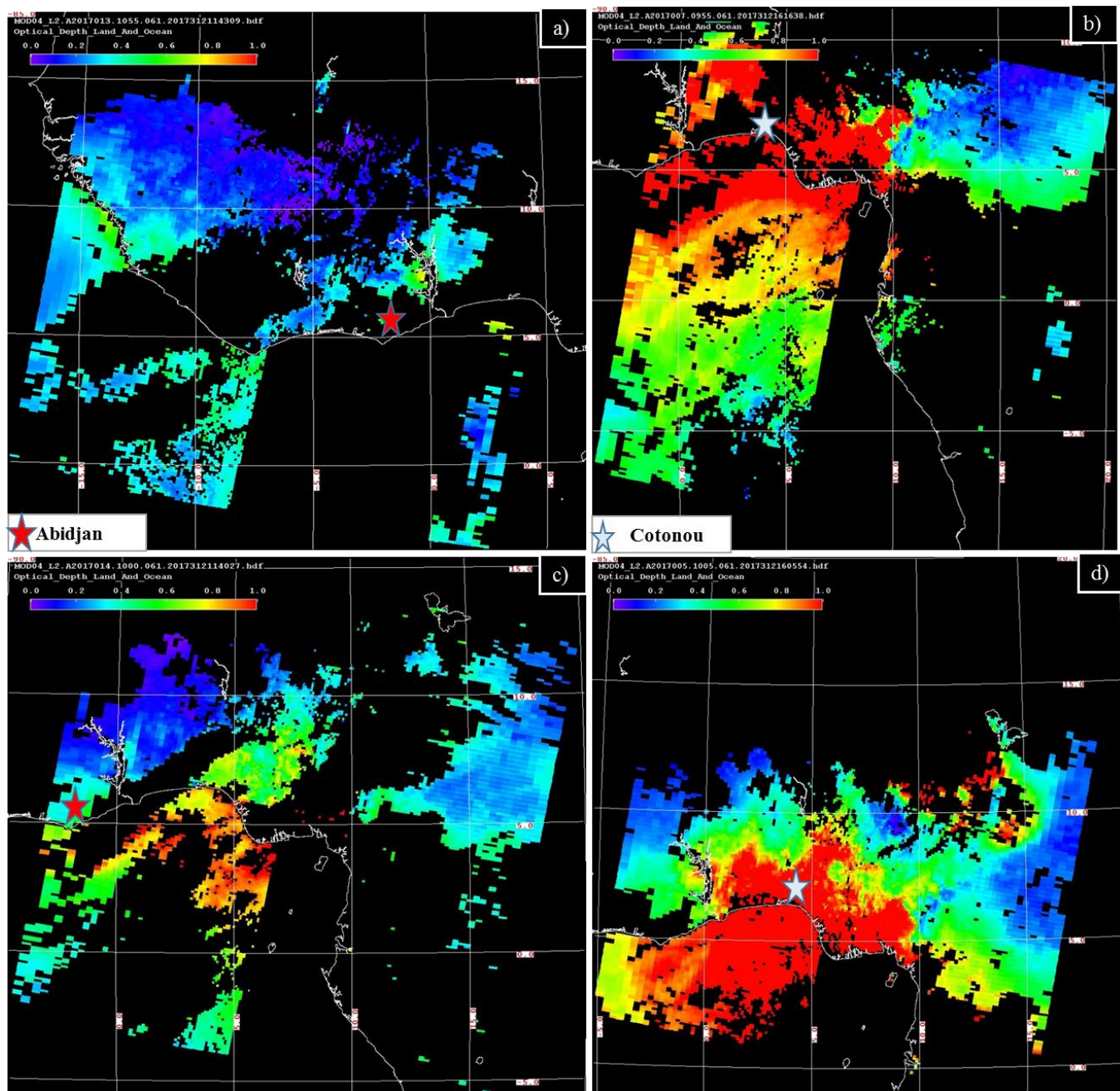
1387 **Figure 13**

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1393 **Figure 14**

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1416 scale have been selected.

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1440 Table 1: Comparison of dust concentrations obtained from different methodologies in $\mu\text{g}\cdot\text{m}^{-3}$

Dry 2017		Sciare et al. (2005)	Guinot et al. (2007)	Terzi et al. (2010)
ADF	C	18.5	11.2	86.9
	F	9.3	29.7	22.2
	bulk	27.7	40.9	109.1
AWB	C	12.3	48.5	126.4
	F	5.2	39.7	106.4
	bulk	17.6	88.2	232.8
AT	C	16.4	24.8	98.5
	F	4.3	13.4	34.2
	bulk	20.7	38.2	132.7
CT	C	37.9	70.0	98.4
	F	23.4	45.6	55.8
	bulk	61.3	115.6	154.2

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Wet 2016		Sciare et al., 2005	Guinot et al., 2007	Terzi et al.(2010)
ADF	C	35.34	21.5	27.9
AWB	C	13.46	8.6	21.1
AT	C	19.65	37.5	21.4
CT	C	42.98	26.2	52.5
	Bulk	42.98	26.2	52.5

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1450 Table 2: WSOC concentrations ($\mu\text{g.m}^{-3}$) and WSOC/OC ratios (%) for each site, each campaign and
 1451 each aerosol size

Site		Abidjan Waste Burning		Abidjan Domestic Fire	
Period	Size	WSOC	WSOC/OC	WSOC	WSOC/OC
Wet season 2015	Coarse	1.3	24.6	8.2	32.5
	Fine	0.7	19.9	12.8	22.7
	Ultra fine	4.1	43.6	51.3	72.5
	PM2.5	5.5	33.7	69.5	47.2
Dry season 2016	Coarse	0.4	12.3	4.4	18.8
	Fine	0.9	46.9	7.0	20.4
	Ultra fine	1.5	38.4	21.9	61.5
	PM2.5	2.7	32.7	31.0	32.0
Wet season 2016	Coarse	1.3	42.5	16.5	44.3
	Fine	0.8	26.3	17.1	33.0
	Ultra fine	2.0	41.2	79.7	84.5
	PM2.5	3.5	37.1	106.0	52.0
Dry season 2017	Coarse	1.9	32.9	12.1	36.0
	Fine	1.4	38.4	19.9	35.0
	Ultra fine	1.6	11.5	38.6	19.0
	PM2.5	4.0	30.0	65.8	29.0

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1454 Table 2 (suite): WSOC concentrations ($\mu\text{g.m}^{-3}$) and WSOC/OC ratios (%) for each site, each campaign
 1455 and each aerosol size

Site		Abidjan Traffic		Cotonou Traffic	
Period	Size	WSOC	WSOC/OC	WSOC	WSOC/OC
Wet season 2015	Coarse	2.4	39.6	1.1	23.3
	Fine	1.3	46.7	0.5	22.1
	Ultra fine	4.7	29.0	0.4	12.7
	PM2.5	6.9	34.0	2.2	18.0
Dry season 2016	Coarse	1.4	43.0	2.3	64.1
	Fine	1.9	59.0	0.6	10.5
	Ultra fine	4.9	62.0	6.3	42.9
	PM2.5	7.5	49.4	8.0	29.0
Wet season 2016	Coarse	1.1	23.1	1.2	34.7
	Fine	0.5	16.8	0.5	32.2
	Ultra fine	1.4	34.8	0.9	23.0
	PM2.5	2.4	26.0	1.9	28.0
Dry season 2017	Coarse	0.9	24.0	3.5	37.8
	Fine	0.3	24.3	2.4	39.6
	Ultra fine	1.8	14.8	1.9	10.4
	PM2.5	2.6	16.0	6.0	18.2

1456 Table 3: Trace element concentrations for bulk aerosol for each site and for DS2017 and WS2016.

	Bulk ng.m ⁻³ (%)							
	DRY 2017				WET 2016			
	ADF	AWB	AT	CT	ADF	AWB	AT	CT
Al	10050.8 (1.8)	25186.1 (13.7)	14015.8 (12.26)	15480.4 (5.7)	1370.5 (0.4)	1990.1 (3.5)	2191.4 (2.4)	4010.5 (4.4)
K	8634.3 (1.5)	6093.7 (3.3)	3677.7 (3.22)	5068.9 (1.9)	1105.0 (0.3)	472.0 (0.8)	275.9 (0.3)	1076.0 (1.2)
Na	6847.8 (1.2)	23430.5 (12.8)	15372.1 (13.44)	11529.3 (4.3)	2070.6 (0.6)	3735.4 (6.6)	2861.5 (3.1)	5310.2 (5.8)
Ca	4321.2 (0.8)	2923.7 (1.6)	4117.6 (3.60)	6233.5 (2.3)	4124.7 (1.1)	447.5 (0.8)	374.7 (0.4)	4954.02 (5.4)
Mg	1940.6 (0.3)	384.0 (0.2)	410.3 (0.36)	823.2 (0.3)	1524.7 (0.4)	294.9 (0.5)	283.5 (0.3)	619.2 (0.7)
Fe	1709.9 (0.3)	3807.9 (2.1)	1628.1 (1.42)	3406.8 (1.3)	1314.0 (0.4)	709.3 (1.3)	987.3 (1.1)	1549.4 (1.7)
P	1521.9 (0.3)	696.0 (0.4)	147.8 (0.13)	207.4 (0.1)	605.4 (0.2)	8.6	13.2	81.4 (0.1)
Ti	488.9 (0.1)	2270.3 (1.2)	282.8 (0.25)	457.9 (0.17)	170.8 (0.05)	75.7 (0.13)	96.8 (0.11)	154.7 (0.17)
Zn	189.7 (0.03)	80.9 (0.04)	57.9 (0.05)	149.4 (0.06)	60.3 (0.02)	1.9	41.1 (0.04)	36.2 (0.04)
Zr	172.1 (0.03)	390.3 (0.21)	217.9 (0.19)	145.3 (0.05)	-	22.4 (0.04)	36.7 (0.04)	31.2 (0.03)
Pb	87.1 (0.02)	11.0 (0.01)	4.8	11.5	8.3	2.1	2.3	9.3 (0.01)
Sn	79.7 (0.01)	38.4 (0.02)	21.6 (0.02)	37.4 (0.01)	0.77	0.09	0.0006	9.ç (0.01)
Mn	74.2 (0.01)	35.2 (0.02)	33.7 (0.03)	160.6 (0.06)	48.9 (0.01)	12.01 (0.02)	9.1 (0.01)	41.41 (0.05)
Rb	52.4 (0.01)	8.7	5.9 (0.01)	8.5	4.47	0.71	0.85	1.9
Sb	59.9 (0.01)	201.2 (0.11)	123.6 (0.11)	149.04 (0.06)	24.4 (0.01)	0	0.0006	2.9
Ba	37.3 (0.01)	53.3 (0.03)	47.4 (0.04)	65.8 (0.02)	18.5	8.02 (0.01)	9.9 (0.01)	32.0 (0.03)
Ni	36.5 (0.01)	34.5 (0.02)	27.9 (0.02)	50.2 (0.02)	18.00	33.1 (0.06)	9.7 (0.01)	14.9 (0.02)
Cr	29.4 (0.01)	53.8 (0.03)	35.8 (0.03)	28.6 (0.01)	41.9 (0.01)	47.7 (0.08)	24.3 (0.03)	29.7 (0.03)
Sr	28.1 (0.01)	15.5 (0.01)	21.2 (0.02)	34.02 (0.01)	17.02	0	0.19	8.1 (0.01)
Cu	24.0	12.3 (0.01)	3.6	9.6	3.99	0.26	0.87	2.8)
Sr	12.6	-	-	-	17.1	-	0.22	8.9 (0.01)
Li	7.3	15.5 (0.01)	7.8 (0.01)	7.39	0.36	0.32	0.23	0.75
Cd	6.1	1.6	1.0	0.83	1.18	0.05	0.02	0.17
V	5.5	12.4 (0.01)	5.1	10.62	2.14	1.84	2.0	3.35
Mo	5.5	8.0	4.9	3.19	4.56	6.84 (0.01)	2.04	3.2
Cs	5.4	0.9	1.2	0.94	0.11	0.12	0.01	0.17
Hf	4.5	10.8 (0.01)	6.8 (0.01)	4.63	0	0.67	1.03	0.97
As	4.2	4.5	3.1	1.22	0	0 (0)	0.05	0.60
Li	4.0	9.8	5.9 (0.01)	5.82	0.27	0.37	0.16	0.93
Co	3.8	1.1	2.1	35.67 (0.01)	0.86	0.49	0.13	0.33
Ce	3.7	6.8	6.0 (0.01)	9.85	1.06	0.50	0.42	2.03
La	1.8	3.5	2.9	4.78	0.54	0.24	0.25	0.92
Nb	1.5	2.6)	1.4	2.48	0.98	0.46	0.5	0.63
Nd	1.5	2.5	2.4	4.15	0.05	0 (0)	-	0.40
Sc	0.69	1.4	1.1)	1.31		0.00		0.02
Be	0.13	0.19	0.2	0.28	0.003	-	-	0.03
Ga	0.61	1.15	0.8	0.98	0.2	0.11	0.12	0.37
Ge	0.42	1.02	0.8	0.68	0.01	0.07	0.02	0.11
Se	0.91	-	-	0.02	0.59	-	0.20	0.18
Rh	0.02	0.02	0.00002	0.002	-	0.002		0.0002
Te	0.06	0.08	0.08	0.05	0.02	0.02	0.02	0.00
Pr	0.40	0.74	0.7	1.13	0.06	0.01	0.004	0.15
Sm	0.27	0.46	0.45	0.76	0.01	0.00	-	0.07
Eu	0.05	0.08	0.08	0.15	0.01	0.003	-	0.0
Gd	0.31	0.57	0.54	0.86	0.05	0.02	0.001	0.15
Tb	0.04	0.07	0.07	0.10	0.005	-	-	0.02
Dy	0.24	0.42	0.40	0.57	0.001	-	-	0.05
Ho	0.05	0.09	0.09	0.12	0.01	0.008	0.01	0.03
Er	0.16	0.31	0.28	0.35	0.02	0.03	0.02	0.09

Tm	0.02	0.05	0.05	0.05	0.002	0.003	0.001	0.01
Yb	0.18	0.38	0.32	0.34	0.01	0.02	0.03	0.09
Lu	0.03	0.06	0.08	0.06	0.003	0.005	0.005	0.02
Ta	0.07	0.14	0.09	0.16	0.06	0.02	0.02	0.03
W	0.80	1.63	0.69	0.54	0.26	0.41	0.4	0.3
Tl	0.22	0.01	0.03	0.06		0.009		
Bi	0.32	0.26	0.02	0.08	0.06	-	-	0.08
Th	0.41	0.88	0.79	1.29	0.15	0.09	0.09	0.24
U	0.22	0.43	0.49	0.51	0.03	0.03	0.02	0.09
Total	36459.9	65817.6	40312.2	44159.2	12562.9	7874.7	7227.2	18001.1
Mass($\mu\text{g}/\text{m}^3$)	558.8	183.6	114.4	270.0	374.7	56.3	91.6	91.9

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1484 Table 4: Comparison of PM_{2.5} concentrations with literature data. Only literature data given at a daily
 1485 scale have been selected.

Location	PM _{2.5} ($\mu\text{g}\cdot\text{m}^{-3}$)	Reference
Abidjan, Côte d'Ivoire	142	This work
Cotonou, Benin	154	This work
Beijing, China	81.4	Xie et al., 2019
Christchurch, New Zealand	9.2	Tunno et al., 2019
Pune, India	98 ± 28	Pipal et al., 2019
Delhi, India	123	Guttikunda and Calori, 2013
Lahore, Pakistan	91	Colbeck et al., 2011
Ahvaz, Iran	69	Shahsavani et al., 2012
Hong Chong, Hong Kong	54.7 ± 25.6	Cheng et al., 2015
Lecce, Italia	16	Cesari et al., 2016
Libreville, Gabon	35.8	Ngo et al., 2019
Port Gentille, Gabon	60.9	
Kenitra, Morocco	51.3	Zghaid et al., 2009
Bilecik, Turkey	247	Gaga et al., 2018
Algiers, Algeria	34.8	Bouhila et al., 2015
Shobra, Egypt	216	Lowenthal et al., 2015

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1505 Table 5: PM_{2.5}-EC and PM_{2.5}-OC comparison with Djossou et al. (2018) and Xu et al. (2019) values.
 1506 Units are $\mu\text{gC.m}^{-3}$.

Location	Period	PM _{2.5} OC	PM _{2.5} EC	References
Traffic Abidjan. Cote d'Ivoire	July 2015	22.6 ± 3.4	4.3 ± 0.2	This Work
	January 2016	15.2 ± 5.3	7.0 ± 2.6	
	July 2016	9.3 ± 1.3	2.2 ± 0.1	
	January 2017	16.1 ± 1.7	18.9 ± 1.4	
	Djossou et al. 2018	July 2015	3.3 ± 0.2	2.3 ± 0.2
		January 2016	7.7 ± 0.0	3.9 ± 0.0
		July 2016	7.6 ± 0.2	4.9 ± 0.0
		January 2017	19.1 ± 6.2	13.9 ± 5.5
Traffic Cotonou. Benin	July 2015	13.1 ± 1.2	3.5 ± 0.7	This Work
	January 2016	27.8 ± 11.3	10.9 ± 2.6	
	July 2016	6.7 ± 1.9	2.0 ± 0.5	
	January 2017	33.1 ± 4.6	27.3 ± 0.9	
	Djossou et al. 2018	July 2015	4.2 ± 0.7	1.5 ± 0.1
		January 2016	3.0 ± 0.3	1.5 ± 0.2
		July 2016	6.7 ± 0.2	1.6 ± 0.1
		January 2017	14.5 ± 0.8	4.4 ± 0.7
Domestic fire Abidjan. Cote d'Ivoire	January 2016	49.5 ± 12.5	13.6 ± 3.6	Xu et al. 2019
	July 2016	37.0 ± 3.5	9.3 ± 0.8	
	This Work	July 2015	147.2 ± 14.5	16.1 ± 1.6
		January 2016	56.5 ± 51.5	7.4 ± 3.1
		July 2016	172.3 ± 39.0	17.9 ± 4.8
		January 2017	283.9 ± 34.9	37.9 ± 4.3
	Djossou et al. 2018	July 2015	80.5 ± 1.1	32.2 ± 1.6
		January 2016	76.3 ± 13.7	11.4 ± 0.2
July 2016		68.4 ± 16.5	17.4 ± 2.1	
January 2017		66.4 ± 7.5	21.1 ± 6.6	
Waste Burning Abidjan. Cote d'Ivoire	January 2016	72.4 ± 24.6	19.5 ± 7.3	Xu et al. 2019
	July 2016	189.3 ± 197.8	11.5 ± 10.8	
	This Work	July 2015	14.8 ± 1.1	4.4 ± 0.1
		January 2016	7.7 ± 1.3	3.0 ± 0.3
		July 2016	10.0 ± 2.4	1.5 ± 0.3
		January 2017	21.9 ± 4.2	19.2 ± 2.4
	Djossou et al. 2018	July 2015	3.7 ± 2.2	4.3 ± 0.3
		January 2016	13.9 ± 9.0	3.6 ± 1.8
July 2016		9.8 ± 4.4	2.8 ± 0.9	
January 2017		22.4 ± 7.8	8.7 ± 3.0	
Xu et al. 2019	January 2016	85 ± 57.4	15 ± 4.7	
	July 2016	65.2 ± 65.2	12.3 ± 11.4	

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1510 Table 6: EC and OC comparison with literature values. Only literature data given at a daily scale have
 1511 been selected.

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Location	OC ($\mu\text{g}\cdot\text{m}^{-3}$)	BC ($\mu\text{g}\cdot\text{m}^{-3}$)	Reference
Abidjan (Côte d'Ivoire)	16	8.1	This study
Cotonou (Benin)	20.2	11	This study
Bilecik (Turkey)	49.6-62.8	38.8-58.1	Gaga et al., 2018
Pune (India)	30	5	Pipal et al., 2019
Shanghai (China)	4.9-13.1	1.9-5	Ding et al., 2017
Lahore (Pakistan)	85.7-152	13.8-21	Stone et al., 2010
Agra (India)	25.4-70	3.3-9.5	Satsangi et al., 2012, Pipal et al., 2014
Delhi (India)	34.1-50	5.3-10.6	Bisht et al., 2015a, Pipal et al., 2014
Ahmedabad (India)	18.3	3	Rengarajan et al., 2011
Yokohama (Japan)	4	2	Khan et al., 2010
Beijing (China)	2.9-28.2	1.2-16.3	Guinot et al., 2007

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1534 Table 7: Comparison of WSOC concentrations with literature data. Only literature data given at a daily
 1535 scale have been selected.

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Location	WSOC ($\mu\text{g}\cdot\text{m}^{-3}$)	Reference
Abidjan, Côte d'Ivoire	2-8	This work
Cotonou, Benin	2-8	This work
Beijing, China	9-27	Yu et al., 2018
Beijing, China	4-6	Xiang et al., 2017
Beijing, China	8-12	Tang et al., 2016
Beijing, China	7	Du et al., 2014
Beijing, China	6-8	Feng et al., 2006
Shanghai, China	2-7	Feng et al., 2006, Huang et al., 2012
Guangzhou, Hong Kong	2	Huang et al., 2012
Guangzhou, Hong Kong	5-10	Feng et al., 2006
Gwangju, Korea	2-3.5	Park et al., 2015
Tokyo, Japan	3-23	Sempere and Kawamura, 1994
Cairo, Egypt	3	Favez et al., 2008
Amsterdam, Netherland	1-2	Feng et al., 2007
Barcelone, Spain	1-2	Viana et al., 2007 and 2008
Brindisi, Italy	1.5	Genga et al., 2017
Saint Jean de Maurienne, France	1-5	Sullivan et al., 2004, Jaffrezo et al., 2005a

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