

1 **Strong influence of Black Carbon on aerosol optical properties in**  
2 **Central Amazonia during the fire season**

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13 **Abstract.** During the dry season, the Amazonian atmosphere is strongly impacted by fires, even in remote areas. However,  
14 there are still knowledge gaps regarding how each aerosol type affects the aerosol radiative forcing. This work characterizes  
15 the chemical composition of submicrometer aerosols and source apportionment of Organic Aerosols (OA) and Equivalent  
16 Black Carbon (eBC) to study their influence on light scattering and absorption at a remote site in central Amazonia during the  
17 dry season (August-December 2013). We applied Positive Matrix Factorization (PMF) and multi-linear regression models to  
18 estimate chemical-dependent mass scattering (MSE) and extinction (MEE) efficiencies. Mean PM1 aerosol mass loading was  
19  $6.3 \pm 3.3 \mu\text{g m}^{-3}$ , with 77% of organics, grouped into 3 factors: Biomass Burning OA (BBOA), Isoprene derived Epoxydiol-  
20 Secondary OA (IEPOX-SOA) and Oxygenated OA (OOA). The bulk scattering and absorption coefficients at 637 nm were  
21  $17 \pm 10 \text{ Mm}^{-1}$  and  $3 \pm 2 \text{ Mm}^{-1}$ , yielding a single scattering albedo of  $0.87 \pm 0.03$ . Although eBC represented only 6% of the PM1  
22 mass loading, MSE was highest for the eBC ( $13.58 \pm 7.62 \text{ m}^2 \text{ g}^{-1}$  at 450-700 nm), followed by BBOA ( $7.96 \pm 3.10 \text{ m}^2 \text{ g}^{-1}$ ) and  
23 ammonium sulfate (AS,  $4.79 \pm 4.58 \text{ m}^2 \text{ g}^{-1}$ ). MEE was dominated by eBC (30.8%), followed by the OOA (19.9%) and AS  
24 (17.6%). The dominance of eBC over light scattering, in addition to absorption, depicts a ~~surprisingly~~ remarkably high role of  
25 this important climate agent, ~~indicating the need with~~ potentially broad implications for more precise radiative forcing  
26 quantification, increasing climate modelling precision, representing deep contributions, to further investigate the chemical  
27 processing and interaction between natural and anthropogenic aerosol sources over remote tropical forested areas. Earth's  
28 climate system comprehension.

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1 Introduction

The Amazon is the largest hydrologic basin and contiguous tropical forest area of the planet, representing important carbon reserves, sources of freshwater, and housing a large biodiversity (Andreae et al., 2015; Artaxo et al., 2013; Davidson et al., 2012; Pöhlker et al., 2016). The strong coupling between climate and the biological functioning of the forest is a key factor in the maintenance of the Amazonian ecosystem (Martin et al., 2010b; Pöhlker et al., 2012). The Amazonian atmosphere is considered an important reactor, regulating its physical properties and chemical composition due to the high insolation and humidity (Andreae, 2001). Despite the high temperatures, precipitation rates, and insolation during most of the year, an annual cycle can be observed, with a wetter and less warm season and a drier and hotter season, whose length and period vary depending on the region of the Amazon (Marengo et al., 2001).

During the wet season, the Amazonian atmosphere represents one of the few continental regions with episodic atmospheric composition near pristine conditions (Andreae et al., 2015; Martin et al., 2010b; Pöhlker et al., 2018; Pöschl et al., 2010). Particle number concentration during the cleanest period is a few hundred particles  $\text{cm}^{-3}$ , very similar to remote oceanic regions (Andreae et al., 2015; Artaxo et al., 2013; Martin et al., 2010). However, during the dry season, forest fires are provoked in order to clear forest areas for agriculture, and also as part of pasture and cropland management (Aragão et al., 2016; Berenguer et al., 2021; Davidson et al., 2012). During these periods, forest fire emissions coupled with smaller rates of aerosol scavenging lead to particle number concentration increases by a factor of 10 in remote forest areas (Artaxo et al., 2013; Pöhlker et al., 2018). These stark seasonal differences in aerosol loading and composition have the potential to significantly modify the coupling biosphere-atmosphere (Zaveri et al., 2022), which is expected to be exacerbated in the future due to extreme climatic events in Amazonia (Flores et al., 2024).

Atmospheric aerosol particles influence climate through scattering and absorption of solar radiation (aerosol-radiation interactions, ARI) and by affecting cloud formation and lifetime (aerosol-cloud interactions, ACI) (Forster et al., 2021). However, the magnitude and the signal of global radiative forcing of aerosols still represent one of the largest uncertainties in global climate models (Kuhn et al., 2010; Rizzo et al., 2013; Szopa et al., 2023). Uncertainties on the radiative forcing of individual aerosol components are even higher (Myhre et al., 2013), with a direct impact on the accuracy of future climatic scenarios (Forster et al., 2021). Carbonaceous aerosols (i.e. organic and black carbon) dominate the Amazonian atmosphere (Artaxo et al., 2013). While Organic Aerosol (OA) originates from both primary emissions, as well as secondary formation from gaseous precursors (Martin et al., 2010), black carbon is mostly primarily emitted from incomplete combustion, and in remote areas of the Amazon it is associated with Amazonian or transatlantic forest fires (Artaxo et al., 2013; Holanda et al., 2020). The sign and magnitude of the ARI forcing are dependent on several parameters such as particle origin, size distribution, mixture and age, notably affecting the light-absorbing component of OA, termed brown carbon (Laskin et al., 2015; Saturno et al., 2018b).

The secondary component of OA (SOA) in the Amazon has been shown to be a major component, notably during the wet season (Chen et al., 2015; Shrivastava et al., 2019). Isoprene (2-methyl-1,3-butadiene,  $\text{C}_5\text{H}_8$ ) is the most abundant VOC emitted

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globally, mostly located in tropical forests such as the Amazon (Marais et al., 2016; Yáñez-Serrano et al., 2015). The formation of isoprene-derived Secondary Organic Aerosol (SOA) is a sequence of complex reactions and depends on different factors, such as low concentrations of NO (a typical byproduct of fossil fuel combustion), and pre-existing aerosol particles where isoprene can condense on (Brito et al., 2018; Caravan et al., 2024; Marais et al., 2016; Nah et al., 2019). One of the dominating isoprene SOA pathways in the Amazon is through the OH attack, leading to hydroperoxy radicals and subsequently via the HO<sub>2</sub> pathway (Shrivastava et al., 2019; Wennberg et al., 2018). This pathway can lead to different low-volatility products generally termed IEPOX-SOA (Isoprene EPOXdiols-Secondary Organic Aerosol) (Allan et al., 2014; Hu et al., 2015; Surratt et al., 2010). Isoprene oxidation product mixing ratios were previously shown to be higher during the dry season above the forest canopy, likely due to the higher insolation and temperature during this period, which favors the oxidative capacity of the atmosphere and leaf emission potential (Yáñez-Serrano et al., 2015). IEPOX-SOA mass concentrations have been shown to be significantly reduced during polluted conditions, associated with suppression due to urban NO emissions (de Sá et al., 2017). Alternatively, a study in polluted urban plumes in West Africa found that SO<sub>4</sub> increase plays a larger role in enhancing IEPOX-SOA loadings than NO in suppressing it (Brito et al., 2018). Secondary oxidized aerosol particles have been demonstrated to be more efficient at scattering radiation than primary particles (Kleinman et al., 2020; Paredes-Miranda et al., 2009; Reid et al., 2005; Smith et al., 2020).

Chemical composition, processes, and sources of atmospheric aerosol particles in the Amazon have been widely studied during both the wet and dry seasons, in sites representing pristine conditions (Andreae et al., 2015; Chen et al., 2015; Martin et al., 2010a) as well as strongly impacted by fires and urban pollution (Brito et al., 2014; Ponczek et al., 2021; de Sá et al., 2018; Zaveri et al., 2022). Physical properties of radiation absorption and scattering were described (Artaxo et al., 2013; Nascimento et al., 2021; Palácios et al., 2020; Rizzo et al., 2013; de Sá et al., 2019; Sena et al., 2013). However, the intrinsic optical properties of each aerosol species are still rare (Velázquez-García et al., 2023), notably associated with OA origins (Ponczek et al., 2021). Our study details the chemical properties of submicrometer aerosol particles in a forest site in central Amazonia during the dry season and their influence on radiation scattering and absorption. We applied positive matrix factorization (PMF) to the organic fraction, associated mass extinction, absorption, and scattering efficiencies to different aerosol components via multi-linear regression to improve our comprehension of their intrinsic properties, as well as estimate their role on aerosol-radiation interaction in Central Amazonia during the dry season.

The strong coupling between climate and the biological functioning of Amazonia is a key factor in the maintenance of its ecosystem (Martin et al., 2010b; Pöhlker et al., 2012). The Amazonian atmosphere is considered an important reactor, regulating its physical properties and chemical composition due to the high insolation and humidity (Andreae, 2001). However, during the dry season, forest fire emissions coupled with smaller rates of aerosol scavenging lead to particle number concentration increases by a factor of 10 in remote forest areas compared to near pristine conditions episodes during the wet season (Andreae et al., 2015; Artaxo et al., 2013; Pöhlker et al., 2018). These stark seasonal differences in aerosol loading and composition have the potential to significantly modify the biosphere-atmosphere coupling (Zaveri et al., 2022). These seasonal differences are expected to be exacerbated in the future due to extreme climatic events in Amazonia (Flores et al., 2024).

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96 Carbonaceous aerosols (i.e. organic aerosols, OA and black carbon, BC) dominate the atmosphere particles chemical classes  
97 in Amazonia (Artaxo et al., 2013). The secondary component of OA (SOA) has been shown to have a major contribution,  
98 notably during the wet season (Chen et al., 2015; Shrivastava et al., 2019). In remote regions of Amazonia, aged and highly  
99 processed oxygenated particles originated from multiple sources (forest fires, derived from volatile organic compounds –  
100 VOCs...) are a major component of basin-wide haze observed during biomass burning season (Darbyshire et al., 2019).  
101 Isoprene (2-methyl-1,3-butadiene, C<sub>5</sub>H<sub>8</sub>) is the most abundant VOC emitted globally, mostly in tropical forests (Marais et al.,  
102 2016; Yáñez-Serrano et al., 2015). The formation of isoprene-derived Secondary Organic Aerosol (SOA) is a sequence of  
103 complex reactions and depends on different factors, such low concentrations of NO and pre-existing aerosol particles where  
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107 EPOXydiols-Secondary Organic Aerosol) (Allan et al., 2014; Hu et al., 2015; Surratt et al., 2010). While OA originates from  
108 both primary emissions, as well as secondary formation from gaseous precursors (Martin et al., 2010b), BC is mostly primarily  
109 emitted from incomplete combustion, and in remote areas of Amazonia it is associated with regional or transatlantic forest  
110 fires (Artaxo et al., 2013; Holanda et al., 2020; Saturno et al., 2018a).  
111 Atmospheric aerosol particles influence climate through scattering and absorption of solar radiation (aerosol-radiation  
112 interactions, ARI) and by affecting cloud formation and lifetime (aerosol-cloud interactions, ACI) (Forster et al., 2021).  
113 However, the magnitude and the signal of global radiative forcing of aerosols still represent one of the largest uncertainties in  
114 global climate models (Szopa et al., 2023). Uncertainties on the radiative forcing of individual aerosol components are even  
115 higher, with a direct impact on the accuracy of future climate scenarios (Forster et al., 2021). The sign and magnitude of the  
116 ARI forcing are dependent on several parameters such as particles size distribution, mixture, aging processes and  
117 meteorological conditions, as well as the particle chemical composition and its effect on the complex refractive index, based,  
118 among other factors, on the origin of the particles (Laskin et al., 2015; Li et al., 2024; Saturno et al., 2018a). Aerosol particles  
119 known for efficiently absorbing radiation - such as BC - often also exhibit significant scattering efficiencies, which are strongly  
120 influenced by their size, chemical composition, and the extent and nature of their atmospheric aging and coatings (Bond and  
121 Bergstrom, 2006; Schwarz et al., 2006; Yu et al., 2010). Although chemical aging has shown to enhance light absorption due  
122 to the coating of the BC core by condensing semi- and intermediate volatility organic compounds or coagulation with other  
123 particles (Darbyshire et al., 2019; Metcalf et al., 2013; Saturno et al., 2018b; Tasoglou et al., 2017; Wang et al., 2016), primary  
124 biomass burning aerosols have also been associated with high scattering efficiencies (Hand and Malm, 2007; Malm et al.,  
125 2005). Coating by non-absorbing material, such as Organics (Romshoo et al., 2021), has been shown to increase BC scattering  
126 by a factor of 3-24 depending on the size, morphology, aging stage, coating thickness and composition of the BC particles (He  
127 et al., 2015). Conversely, sulfate and water coating have also shown to increase elemental carbon particle diameter, playing a  
128 stronger role on its scattering efficiency, more than absorption (Cheng et al., 2008; Yu et al., 2010). Precisely quantifying  
129 distinct ARI for each chemical species, and especially decreasing uncertainties on the ones with high potential to both absorb

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130 and scatter radiation such as BC is critical to improve our understanding and prediction of the atmospheric system and improve  
131 climate models.  
132 Chemical composition, processes, and sources of atmospheric aerosol particles in Amazonia have been widely studied during  
133 both the wet and dry seasons, in sites representing pristine conditions (Andreae et al., 2015; Cheng et al., 2015; Martin et al.,  
134 2010a) as well as strongly impacted by fires and urban pollution (Brito et al., 2014; Palm et al., 2018; Ponczek et al., 2021;  
135 Zaveri et al., 2022). Physical properties of radiation absorption and scattering have been described for the whole particles mass  
136 loading, regardless of the specific chemical groups (Artaxo et al., 2013; Nascimento et al., 2021; Palácios et al., 2020; Rizzo  
137 et al., 2013; de Sá et al., 2019; Sena et al., 2013). However, intensive optical properties of each aerosol species are still rare  
138 (Velazquez-Garcia et al., 2023), notably associated with OA origins (Ponczek et al., 2021) and with BC behaviour. Our study  
139 details the chemical properties of submicrometer aerosol particles in a forest site in central Amazonia during the dry season  
140 and their influence on radiation scattering and absorption. We applied positive matrix factorization (PMF) to the organic  
141 fraction, and associated mass extinction, absorption, and scattering efficiencies to different aerosol components via multi-  
142 linear regression (MLR) to improve our comprehension of their intrinsic properties, as well as estimate their role on ARI in  
143 Central Amazonia during the dry season.

## 144 2 Material and Methods

### 145 2.1 Sampling site

146 The measurements site is located in Central Amazonia, 60 km northwest of the city of Manaus, Brazil, in the Cuieiras biological  
147 reserve (2°35'39.24''S, 60°12'33.42''W), and referred to in this study as T0z (Martin et al., 2015; Whitehead et al.,  
148 2016)(Martin et al., 2015; Whitehead et al., 2016). The vegetation is characterized as *terra firme* (upland forest, not impacted  
149 by seasonally flooded seasonal flooding), and the canopy is between 30 m and 35 m high (Martin et al., 2010)(Martin et al.,  
150 2010a, 2010). As a result of steady northeasterly-easterly winds (Andreae et al., 2015; Araújo et al., 2002)(Andreae et al.,  
151 2015; Araújo et al., 2002), only rarely the site is impacted by Manaus emissions (Chen et al., 2015). The seasonality at the  
152 region of this site in central Amazonia has been previously defined as the wet season in this region is typically from 1 December  
153 – 14 June, and the dry season from 15 June – 30 November (Andreae et al., 2015). During the wet season, air masses reaching  
154 the site pass over more than 1,500 km of undisturbed forest (Andreae et al., 2015; Pöschl et al., 2010)(Pöschl et al., 2010).  
155 However, during the dry season, regional biomass burning pollution can be detected at the site (Artaxo et al., 2013; Rizzo et  
156 al., 2013)(Artaxo et al., 2013; Rizzo et al., 2013), as well as aerosol plumes advected from African wildfires (Holanda et al.,  
157 2023). Our observations comprise from 1 August until 10 December 2013, sampling the atmosphere at 38.8 meters above  
158 ground level. The instrumentation was located inside an air-conditioned container at the base of the tower. A cyclone (50 %  
159 cut-off at 10 µm) was used at the entrance of the inlet. An automatic diffusion dryer (Tuch et al., 2009)(Tuch et al., 2009) kept  
160 the relative humidity of the sampled air between 20% and 50%. Lodging for scientists/staff and a diesel generator were located

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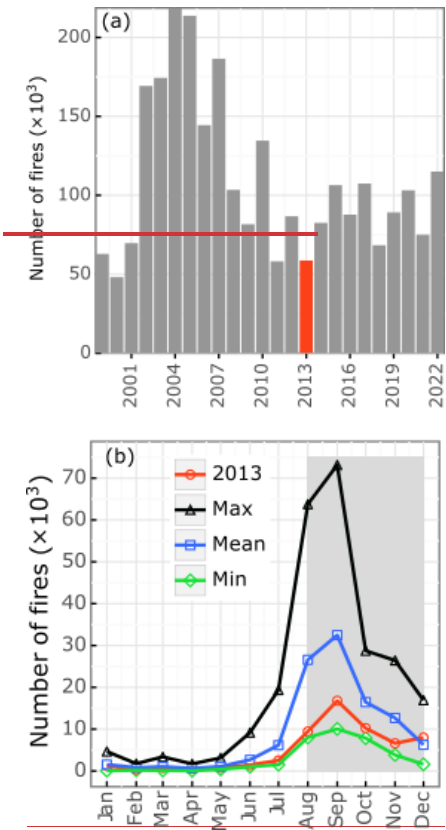
161 330m and 720m downwind (west) from the tower, respectively. The measurement tower has been shown to be practically  
162 unaffected by the generator (Whitehead et al., 2016). The year of this study (2013) was characterized by a historical minimum  
163 of fire detection over ~~the last~~ 20 years (Figure 1, ~~(F. G. Assis et al., 2019)~~(F. G. Assis et al., 2019)), providing an interesting  
164 outlook to assess the best scenarios for a dry season in recent times, and thus evaluate atmospheric composition within targets  
165 and goals for the ~~Amazon forest~~Amazonia rainforest preservation. The observation period here has been considered to fit  
166 entirely within dry-season atmospheric conditions. The previous transitional (wet-dry) period occurred in June-July  
167 ~~(Whitehead et al., 2016)~~(Whitehead et al., 2016), and the subsequent (dry-wet) soon after the end of our measurements.

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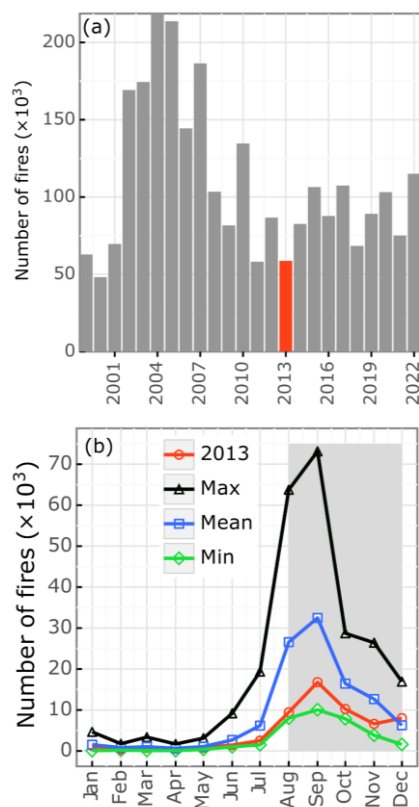
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**Figure 1: a) Number of fires in the Brazilian Amazon forest/Amazonia rainforest, from 1999 to 2022, showing how 2013 (marked in red) is the lowest in the past 20 years in terms of total number of fires in 20 years, and b) mean (blue), maximum (black), and minimum (green), and for 2013 (red) monthly fires between 1999-2022 in the Amazon/Amazonian Basin. The year of 2013 is marked in red, and it is evident how it was very close to the minimum (green) line. The gray area in b) marks the period of measurements in our study (01/Aug – 10/Dec) (Instituto Nacional de Pesquisas Espaciais, 2024).**

## 2.2 Instrumentation

Non-refractory submicrometer aerosol composition was measured using a quadrupole Aerosol Chemical Speciation Monitor (ACSM, Aerodyne Research Inc) (Ng et al., 2011), which is a compact version of the Aerosol Mass Spectrometer (AMS).

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Instrument calibration consisted of injecting monodispersed (300 nm) aerosol particles of ammonium nitrate (AN) and ammonium sulfate (AS). Aerosol particles were generated using an atomizer and subsequently dried, and size selected using a Differential Mobility Analyzer. A collection efficiency of 0.5 has been adopted (Middlebrook et al., 2012), yielding a very good agreement in of particle mass considering measurements from different collocated instruments (S1). This method was successfully used in previous studies (Brito et al., 2014; Sun et al., 2010)(Brito et al., 2014; Sun et al., 2010), and the value of 0.5 agrees with other studies in the Amazon Amazonia during the dry season (de Sá et al., 2019) and during the transition from wet to dry season (Ponczek et al., 2021)(Ponczek et al., 2021; de Sá et al., 2019). The measured ammonium (NH<sub>4</sub>) mass concentrations were close to, or often lower than the detection limit of 0.28 μm<sup>-3</sup> (Pöhlker et al., 2018; Whitehead et al., 2016) μg m<sup>-3</sup> (Pöhlker et al., 2018; Whitehead et al., 2016), and were therefore calculated based on nitrate (NO<sub>3</sub>) and sulfate (SO<sub>4</sub>) molar masses and their mass concentrations, assuming neutralization as in Equation 1:

$$NH_{4, predicted} = 18 \times \left( \frac{SO_4}{96} \times 2 + \frac{NO_3}{62} \right) \quad (1)$$

Furthermore, the SO<sub>4</sub> and NO<sub>3</sub> ions were used to estimate AS and AN (Equations 2 and 3) for the chemical-dependent optical properties analyses (Section 3.3), assumed here to be their most abundant form given the very low NH<sub>4</sub> levels:

$$AS = 132 \times \frac{SO_4}{96} \quad (2)$$

$$AN = 80 \times \frac{NO_3}{62} \quad (3)$$

Size-resolved particle number size distribution from 10 to 450 nm was measured with a Scanning Mobility Particle Sizer (SMPS, model 3081, TSI Inc.) coupled to a Condensation Particle Counter (CPC, model 3772, TSI Inc.) to provide equivalent mobility particle diameter for singly charged particles (D<sub>pg</sub>, (Wiedensohler et al., 2012)). Aerosol light scattering coefficient (σ<sub>s</sub>) at 450 nm, 550 nm, and 700 nm (Anderson and Ogren, 1998) was measured using a Nephelometer (model 3563, TSI Inc.). Calibration was performed using CO<sub>2</sub> as the high-span gas and filtered air as the low-span gas. The averaging time applied was 60 minutes, and therefore the detection limits, (defined as a signal-to-noise ratio of 2,) for scattering coefficients are 0.08, 0.03, and 0.05 Mm<sup>-1</sup> for 450, 550, and 700 nm, respectively (Anderson and Ogren, 1998). Since a PM10 inlet was used, the “no-cut” factors were used for the truncation corrections (Anderson and Ogren, 1998). Scattering coefficients at 637 nm were calculated from interpolation, assuming a power law spectral dependency. A Since a PM10 inlet was used, the “no-cut” factors were used for the truncation corrections. We used a Multi Angle Absorption Photometer (MAAP, model 5012, Thermo Electron Group, Waltham, USA) (Müller et al., 2011) measured to measure aerosol light absorption coefficient (σ<sub>a</sub>) at 637 nm, and was used to estimate equivalent Black Carbon (eBC) concentration, assuming an absorption cross-section value of 6.6 m<sup>2</sup> g<sup>-1</sup>. Considering the conditions of the experiment, the MAAP detection limit for σ<sub>a</sub> was of 0.13 Mm<sup>-1</sup> (Petzold et al., 2005).

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Episodes of possible contamination from the city of Manaus and from the diesel generator were removed by filtering the datapoints when either the wind direction was between 270-340° (from our local wind direction measurements) or when the calculated backtrajectories from the Hysplit model (Draxler and Hess, 1998) passed over Manaus coordinates, as in (Whitehead et al., 2016) (Supplement S2).

### 2.3 Optical properties

Single Scattering Albedo (SSA, Equation 4) is a measure of the ratio of  $\sigma_s$  to the total radiation extinction coefficient ( $\sigma_e = \sigma_s + \sigma_a$ ) by aerosol particles (Rizzo et al., 2013). Since the MAAP instrument only measures the  $\sigma_a$  at 637 nm wavelength, the  $\sigma_e$  was calculated only for this wavelength using  $\sigma_s$  at 637 nm from the nephelometer, estimated using the scattering Angstrom exponent ( $\alpha_s$ , Equation 5):

$$SSA = \frac{\sigma_s}{\sigma_s + \sigma_a} \quad (4)$$

The  $\alpha_s$  is a measure of the dependence of radiation scattering on the light wavelength ( $\lambda$ ), and it is an indication of particle size (Rizzo et al., 2013; Saturno et al., 2018b; Schuster et al., 2006):

$$\ln \sigma_s = -\alpha_s \ln \lambda + \ln(\text{constant}) \quad (5)$$

Scattering coefficients at 637 nm were calculated from interpolation using the scattering Angström exponent ( $\alpha_s$ , Equation 4), assuming a power-law spectral dependency. The  $\alpha_s$  is a measure of the dependence of radiation scattering on the light wavelength ( $\lambda$ ), and it is an indication of particle size (Rizzo et al., 2013; Saturno et al., 2018b; Schuster et al., 2006):

$$\ln \sigma_s = -\alpha_s \ln \lambda + \ln(\text{constant}) \quad (4)$$

Single Scattering Albedo (SSA, Equation 5) is a measure of the ratio of  $\sigma_s$  to the total radiation extinction coefficient ( $\sigma_e = \sigma_s + \sigma_a$ ) by aerosol particles (Rizzo et al., 2013). Since the MAAP instrument only measures the  $\sigma_a$  at a wavelength of 637 nm, the  $\sigma_e$  was calculated using  $\sigma_s$  at 637 nm interpolated from the nephelometer.

$$SSA = \frac{\sigma_s}{\sigma_s + \sigma_a} \quad (5)$$

After rain events and other moments when the atmosphere is very clean, both  $\alpha_s$  for  $\sigma_e$  values all the optical parameters are very low close to zero, and therefore the ratio between them (SSA, Equation 4) becomes unrealistically high. We therefore calculated SSA for  $\alpha_s$  for only when  $\sigma_e$  and  $\sigma_e > 1 \text{ Mm}^{-1}$ .

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245 2.4 Statistical Analyses

246 2.4.1 Positive Matrix Factorization (PMF)

247 ~~The PMF (Positive Matrix Factorization) was used on the submicrometer non-refractory organic mass spectra in order to group~~  
248 ~~m/z ratios with similar temporal variability, supporting the identification of sources and processes that formed and transformed~~  
249 ~~atmospheric particles. A detailed description of the PMF can be found in (Paatero & Tapper, 1994; Ulbrich et al., 2009), and~~  
250 ~~it is an established methodology for aerosol source apportionment (Zhang et al., 2011). The model can be represented by the~~  
251 ~~following Equation. (6):~~

252 We used Positive Matrix Factorization (PMF) in order to group the submicrometer non-refractory organic mass spectra (m/z  
253 ratios) with similar temporal variability, supporting the identification of sources and processes that formed and transformed  
254 atmospheric particles (Paatero and Tapper, 1994; Ulbrich et al., 2009; Zhang et al., 2011). The model can be represented by  
255 the following Equation. (6):

256 
$$\mathbf{X}_{(m \times n)} = \sum_{p=1}^p \mathbf{G}_{(m \times p)} \mathbf{F}_{(p \times n)} + \mathbf{E} \quad (6)$$

259 Where **X** is the input matrix of *n* (elements – m/z ratios) lines and *m* (number of samples) columns (Ulbrich et al., 2009).  
260 (Ulbrich et al., 2009). In this study, the **X** matrix had 2901 lines (1-hour averages for more than 4 months of measurements)  
261 and 70 columns (m/z ratios). The receptor model aims to determine the number of *p* factors, representing sources or processes,  
262 their chemical composition, and the relative contribution of each factor. **G** is a matrix in which columns are the time series of  
263 the factors. **F** is a matrix in which lines are the profiles of the factors (mass spectra). **E** represents the residuals, the part of the  
264 data that was not modeled by any factor *p*. We used an IGOR™-based interface to apply the PMF analysis (Ulbrich  
265 et al., 2009). The PMF ions were normalized to the organics concentration.

266 2.4.2 Multilinear Regression (MLR)

267 ~~We used a Multilinear Regression (MLR) model to estimate the contribution of each aerosol chemical component to scattering,~~  
268 ~~absorption, and extinction coefficients, deriving the corresponding efficiencies (MSE, MAE, and MEE, respectively). The  $\sigma_s$~~   
269 ~~and  $\sigma_a$  were the dependent variables (response) and the species/factors were the independent (predictor) variables.~~

270 We used a Multilinear Regression (MLR) model to estimate the contribution of each aerosol chemical component to scattering,  
271 absorption, and extinction coefficients, deriving the corresponding efficiencies (MSE, MAE, and MEE, respectively) (Yu et  
272 al., 2010). The scattering ( $\sigma_s$ ) and extinction ( $\sigma_a$ ) coefficients (Mm<sup>-1</sup>) were the dependent variables (response) and the ACSM  
273 species/PMF factors were the independent (predictor) variables.

274 A generalization of the mass efficiency (ME) calculation is presented in Equation 7:

275 
$$ME = \sum_i a_i x_i + r \quad (7)$$

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Where ME can be MSE, MAE or MEE;  $x$  is the chemical species mass concentration;  $a_i$  is the efficiency of each component, and  $r$  are the residuals. We used NNLS (Non-Negative Least Squares) from Python package Scipy version 1.5.2 (Virtanen et al., 2020). To constrain the model to produce results with physical meaning, the coefficients  $a_i$  were constrained to be positive, as in (Velazquez-Garcia et al., 2023). Since eBC is assumed to be the only absorbing component measured in this study with the MAAP, a MLR could not be applied (Virtanen et al., 2020). To constrain the model to produce results with physical meaning, the coefficients  $a_i$  were constrained to be positive, as in (Velazquez-Garcia et al., 2023). Since eBC is assumed to be the only absorbing component measured in this study with the MAAP, a MLR could not be applied for  $\sigma_{a\lambda}$  and the MAE was considered to be equivalent to the cross-section value ( $6.6 \text{ m}^2 \text{ g}^{-1}$ , Section 2.2).

**3 Results and Discussion**  
**3.1 Aerosol chemical composition**

The concentrations of organics and inorganics aerosols follow similar variation patterns during the measurement period (Figure 2a). This can be an indication that the total mass loading consists of well-mixed biomass burning and secondary aerosols, associated with large and regional-scale processes (Darbyshire et al., 2019). The total submicrometer ( $\text{PM1} = \text{Organics} + \text{NO}_3 + \text{NH}_4 + \text{SO}_4 + \text{eBC}$ ) mean mass concentration during the observation period was  $6.3 \pm 3.3 \text{ } \mu\text{g m}^{-3}$  (Table 1, Figure 2a). This represents about half of what was measured during the dry season of the following year (2014, with much more fires, Figure 1) at a nearby site (ATTO tower, Amazon Tall Tower Observatory), with similar conditions (Central central Amazonia, isolated from major biomass burning focus or Manaus urban plume) ( $10.5 \text{ } \mu\text{g m}^{-3}$  (ATTO, (de Sá et al., 2019)) and regions directly impacted by fires during the dry season ( $12.4 \text{ } 13.7 \text{ } \mu\text{g m}^{-3}$  (Brito et al., 2014; Ponczek et al., 2021)). The  $\text{PM1}$  aerosol composition was dominated by the organic fraction, representing a mass fraction of  $77 \pm 5\%$  (Table 1, Figure 2b). This is very similar to the wet season and wet to dry season transition in the same site ( $> 80\%$  (Artaxo et al., 2013; Chen et al., 2015; Whitehead et al., 2016)), but high compared to other continental urban areas, such as across Europe ( $\sim 30\text{--}50\%$ , and regions directly impacted by fires during the dry season (Brito et al., 2014; Ponczek et al., 2021). The  $\text{PM1}$  aerosol composition was dominated by the organic fraction ( $77 \pm 5\%$ , Table 1, Figure 2b), similar to what was found in the same site in wetter conditions (Artaxo et al., 2013; Chen et al., 2015; Whitehead et al., 2016), but lower than what was found in a region highly impacted by biomass burning in southwestern Amazonia (Brito et al., 2014). In continental urban areas, such as across Europe, the organic particles represented a much lower fraction of the total particles mass (Chen et al., 2022) and lower than the strongly impacted by biomass burning region in Southwestern Amazonia (90%, (Brito et al., 2014). Sulfate is the main soluble inorganic component of the aerosol mass fraction in the Amazon. Sulfate is the main soluble inorganic component of the aerosol mass fraction in Amazonia, both during the wet and dry seasons (Fuzzi et al., 2007; Yamasoe et al., 2000)(Fuzzi et al., 2007; Yamasoe et al., 2000). In our study, the mean  $\text{SO}_4$  mass fraction was  $9 \pm 3\%$  (Table 1, Figure 2b), which is

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308 comparable to the ATTO site (Andreae et al., 2015). However, in Southwestern Amazonia, in areas impacted by fresh biomass  
309 burning, the average SO<sub>4</sub> mass fraction was significantly lower (2-3% (Brito et al., 2014; Ponczek et al., 2021)).(Brito et al.,  
310 2014; Ponczek et al., 2021)).  
311 The eBC mass fraction was 6±2% (Table 1, Figure 2b), which is half than of the fraction observed in the wet season at the same  
312 site (11%), despite the much lower total submicrometer aerosol mass loading (Chen et al., 2015), despite the much lower total  
313 submicrometer aerosol mass loading. Occasional urban pollution from Manaus, long-range transport from Africa, and potential  
314 artifacts combined with much lower overall aerosol loading are possible causes for this relatively higher eBC mass fraction  
315 found in the wet season (Chen et al., 2015). In Southwestern Amazonia, during the transition from dry to wet season, the  
316 contribution of eBC to PM<sub>1</sub> reached 15% (Ponczek et al., 2021). Nitrate had a minor contribution during our observations  
317 (3±1%), with concentration levels comparable to the ATTO site (Pöhlker et al., 2018), as well as Southwestern Amazonia  
318 (Brito et al., 2014; Ponczek et al., 2021)). The concentrations of organics and inorganics follow similar patterns during the  
319 measurement period (Figure 2a). This can be an indication that the total mass loading consists of well-mixed biomass burning  
320 and secondary aerosols, associated with large and regional scale processes (Darbyshire et al., 2019). In Southwestern  
321 Amazonia, highly impacted by fresh biomass burning, the contribution of eBC to PM<sub>1</sub> reached 15% (Ponczek et al., 2021).  
322 Nitrate had a minor contribution during our observations (3±1%), with concentration levels comparable to the ATTO site  
323 (Pöhlker et al., 2018), as well as Southwestern Amazonia (Brito et al., 2014; Ponczek et al., 2021).  
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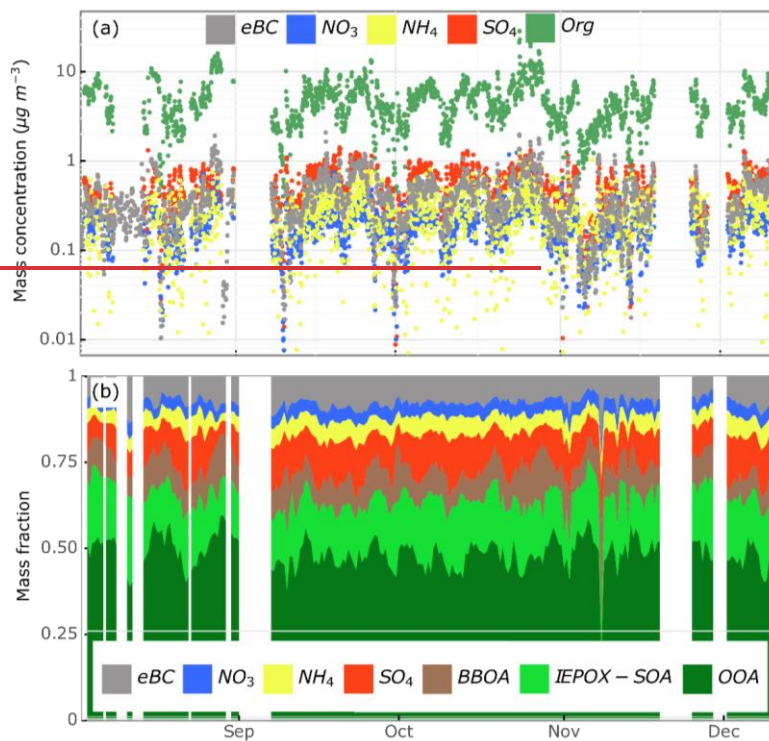
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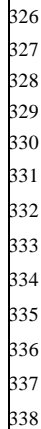
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The PMF analysis yielded 4 factors, although 2 statistical factors, although 2 of them were closely related to the Oxygenated Organic Aerosol (OOA) fraction, and the. Their mass spectra, diurnal/daily profile, and time series of these factors did not present enough differences to justify their separation (Supplements S3 and S4). Therefore, these 2 factors were manually summed in order to generate a 3 factors solution, which was different from the factors found in the 3 factors solution presented by the PMF. In (Ulbricht et al., 2009), the authors describe how one PMF resulting statistical factor can split into various other factors which, after added, represent the real factor. The recombination often considers similarities between the statistical factors in the mass spectra, diurnal/daily profile and time series (Carbone et al., 2013). In our study, the identification of the factors was further confirmed with the correlation between the PMF statistical factors and the inorganic aerosols, with the eBC

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(Supplement S4) and ~~diurnal~~daily profile analyses. The 3 factors were identified as BBOA (Biomass Burning Organic Aerosol), OOA (Oxygenated Organic Aerosol) and IEPOX-SOA (Isoprene derived Epoxydiol–Secondary Organic Aerosol), and they represent together 99% of the measured submicrometer organic aerosol mass, with 1% of residuals. The correlation between BBOA and NO<sub>3</sub> and SO<sub>4</sub> is comparable with findings in Southwestern Amazonia (Brito et al., 2014).

**Table 1 - Dry season (01/Aug – 10/Dec) mean mass concentration (µ m<sup>-3</sup>), standard deviations, and percentile range (10-90, in parenthesis) of the species measured by the ACSM and MAAP (eBC).**

	Mass concentration (µg m <sup>-3</sup> )	Mass fraction
Total PM1	6.3±3.3 (2.7-10.3)	100%
Organics	4.9±2.7 (2.1-7.9)	77±5%
BBOA	0.6±0.6 (0.2-1.1)	9±5%
IEPOX-SOA	1.0±0.5 (0.4-1.7)	17±5%
OOA	3.2±1.5 (1.3-5.5)	51±6%
NO <sub>3</sub>	0.2±0.1 (0.1-0.3)	3±1%
NH <sub>4</sub>	0.3±0.1 (0.0-0.5)	4±1%
SO <sub>4</sub>	0.5±0.3 (0.2-0.9)	9±3%
eBC	0.4±0.3 (0.1-0.7)	6±2%

The ~~largest contribution~~dominant PMF-derived statistical factor in our study was OOA, which contributed to 51±6% of the PM1 (Table 1), and 65% of the organic mass. ~~It presented the largest m/z 44 fraction (Figure 3), and therefore, this is the factor~~ This agrees with the highest estimated O:C ratio (Chen et al., 2015). The high oxidation level indicates highly aged particles and may lose some of their original chemical signatures, in terms of elementary ratios, during the aging process (Jimenez et al., 2009). The m/z 44 is formed mainly by the fragment CO<sub>2</sub><sup>+</sup>, typical of the thermal decarboxylation of the organic acids groups (Alfarra et al., 2004)-previous studies showing that highly processed SOA are a ~~The more aged the aerosols, the more chemically similar they become, which makes the task of separating them into different factors with distinctive characteristics very difficult. Therefore, the OOA factor probably groups aerosol particles from different sources, and their common characteristic is that they are probably originating relatively distant from the sampling site. This factor is a~~major component of basin-wide haze observed during biomass burning season, associated with the transport from eastern Amazonia into the atmospheric particles in remote central regions at the center of the basinAmazonia (Darbyshire et al., 2019). This factor has the highest estimated O:C ratio, which is evident in the observed m/z 44 fraction (Figure 3, note the different scales). The high oxidation level indicates highly aged particles and may lose some of their original chemical signatures, in terms of elementary ratios, during the aging process (Jimenez et al., 2009). The m/z 44 signal predominantly arises from the CO<sub>2</sub><sup>+</sup> ion fragment, which is typically generated by thermal decarboxylation of carboxylic acid functional groups in organic aerosols (Alfarra et al., 2004). Therefore, m/z 44 serves as a valuable marker for the extent of aerosol oxidation and the presence of oxygenated

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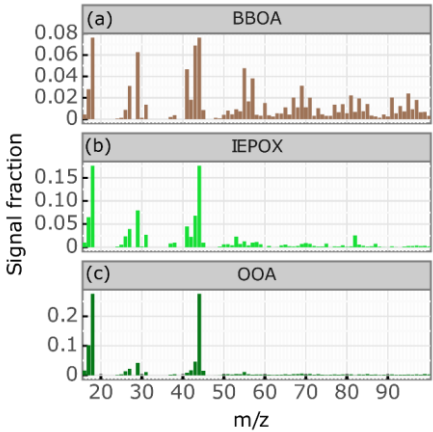
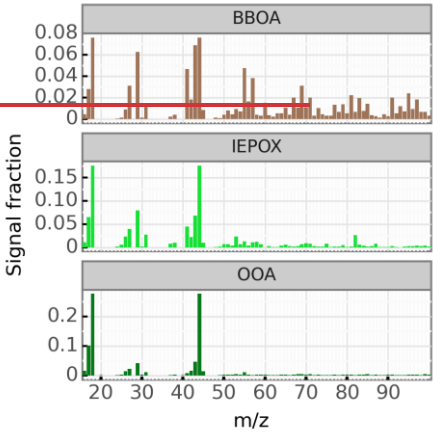
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362 organic compounds, providing insight into aerosol aging and secondary organic aerosol formation processes. The more aged  
363 the aerosols, the more chemically similar they become, which makes the task of separating them into different factors with  
364 distinctive characteristics very difficult. Therefore, the OOA factor probably groups aerosol particles from different sources,  
365 and their common characteristic is that they are probably originating relatively distant from the sampling site.

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368 **Figure 3:** PMF Mass spectra composition of each statistical factor and its relative contribution to the total submicrometer organic  
369 aerosol mass. It is possible to observe typical tracer ions such as m/z 60 and m/z 73 that characterize the BBOA factor, and also the  
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371 m/z 82 in the IEPOX-SOA factor. The scale of the Y axis is different in order to facilitate visualisation of m/z signal fractions mainly  
372 of (a) and (b).

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374 The production of IEPOX-SOA generally leads to the production of markers in the atmosphere, such as the 2-methylthetrol  
375 and the 3-methylfuran (m/z 82, C<sub>5</sub>H<sub>6</sub>O<sup>+</sup>). These markers may not originally exist in the IEPOX-SOA molecule due to their  
376 extremely high volatility, but they can be formed during the decomposition of some IEPOX-SOA species, such as 3-  
377 methyltetrahydrofuran-3,4-diols (3-MeTHF-3,4-diols) (Lin et al., 2012). The organic fraction in the m/z  
378 82 is therefore important for the identification of the IEPOX-SOA factor (Figure 3b), despite its low contribution to the  
379 submicrometer organic aerosol mass fraction (usually below 4%). Beyond that, most of submicrometer organic aerosol.  
380 Most of the other m/z are common to other factors, making the m/z 82 distinctive of the IEPOX-SOA, which can also be  
381 identified by the m/z 53 (C<sub>4</sub>H<sub>5</sub><sup>+</sup>) and m/z 75 (C<sub>3</sub>H<sub>7</sub>O<sub>2</sub><sup>+</sup>) (Allan et al., 2014; Lin et al., 2012; Xu et al., 2015a). The lifetime of  
382 IEPOX-SOA in the boundary layer is estimated to be about 2 weeks (Hu et al., 2016). The IEPOX-SOA mean mass  
383 concentration in our study was 1.0±0.5 µg m<sup>-3</sup> (Table 1, Figure 4). Previous studies reported 0.26 µg m<sup>-3</sup> during the wet  
384 season (Figure 3b) (Allan et al., 2004; Lin et al., 2012; Xu et al., 2015). IEPOX-SOA mean mass concentration in our study  
385 was 1.0±0.5 µg m<sup>-3</sup> (Table 1). Previous studies reported 0.26 µg m<sup>-3</sup> during the wet season at the same site (Chen et al., 2015),  
386 while downwind of Manaus it was around 0.5 µg m<sup>-3</sup> during background conditions, and 0.1 µg m<sup>-3</sup> during polluted conditions  
387 (de Sá et al., 2017). It is important to note that at T0z, while  
388 While the organic particle loading typically increases by an order of magnitude from the wet to the dry season (Artaxo et al.,  
389 2013), we estimated that IEPOX-SOA increases about a factor ~3. The relative contribution of IEPOX-SOA to organics during  
390 the wet season was estimated to be 34%, in contrast to the 17% found in this study (Table 1), which IEPOX-SOA increases by  
391 about a factor ~3 (Table 1 and (Chen et al., 2015), while its relative contribution to the organic aerosols drops by half (from  
392 34% (Chen et al., 2015) to 17%, Table 1). This is likely the result of a complex balance between increased isoprene emissions  
393 (Yáñez-Serrano et al., 2015) sulfate abundance and increased pollution levels (including NO<sub>x</sub>  
394 from forest fires, and biomass-burning related aerosol particles). The relative contribution of IEPOX-SOA to the total PM1  
395 mass was relatively constant during the whole measurement period (Figure 2b), as well as most of the other species (with the  
396 exception of some episodes). This indicates that an atmospheric dynamics of rain/dilution controlling the chemical composition  
397 could be more important than the influence of local sources of particles, confirming the regional haze hypothesis raised by  
398 (Darbyshire et al., 2019).

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399 The correlation (Pearson coefficient = 0.7) observed between the IEPOX-SOA factor and AS (Supplement S4.1) is similar to  
400 the correlation measured in regions affected by urban pollution in the Amazon, Africa and USA (R<sup>2</sup> = 0.37-0.48 (Brito et al.,  
401 2018; Budisulistiorini et al., 2013; de Sá et al., 2017)). Sulfate is the main aqueous-phase particle in which isoprene products  
402 use to condense on (Budisulistiorini et al., 2013; Kroll et al., 2006; Lin et al., 2012; Marais et al., 2016; Surratt et al., 2010;  
403 Xu et al., 2015b), and therefore a positive and moderate-high correlation is expected. eBC presents a similar correlation with  
404 IEPOX-SOA as AS (Supplement S4), but the correlation is even higher with the other PMF statistical factors, especially OOA

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(Pearson coefficient = 0.85, Supplement S4). Since OOA corresponds to more than half of PM1 (Table 1), this high correlation indicates that most of the submicrometer aerosols measured during the dry season in Central Amazonia are, in general, influenced by biomass burning emissions, since eBC is an important combustion tracer. However, the fact that the eBC correlation is higher with the OOA factor than with BBOA (which constitutes only 9% of PM1, Table 1, Supplement S4) indicates that long-range transport of aged and internally well mixed biomass burning plumes plays a more important role than nearby fire sources. The correlation (Pearson coefficient = 0.7) observed between the IEPOX-SOA factor and AS (Supplement S4.1) is similar to the correlation measured in regions affected by urban pollution in Amazonia, Africa and USA (Brito et al., 2018; Budisulistiorini et al., 2013; de Sá et al., 2017). Sulfate is the main aqueous phase particle in which isoprene products condense on, and therefore a positive and moderate-high correlation is expected (Budisulistiorini et al., 2013; Kroll et al., 2006; Lin et al., 2012; Marais et al., 2016; Surratt et al., 2010; Xu et al., 2015). eBC presents a similar correlation with IEPOX-SOA as AS (Supplement S4), but the correlation is even higher with the other PMF factors, especially OOA (Pearson coefficient = 0.85, Supplement S4). This suggests that a significant fraction of the aged submicrometer aerosols measured during the dry season in Central Amazonia is largely influenced by biomass burning emissions, in combination with other combustion sources such as sporadic urban plumes transported from Manaus. In addition, co-variability between aerosol species is expected due to strong washout events that, although less frequent, can still occur during the dry season and impact multiple aerosol components simultaneously. The fact that the eBC correlation is higher with the OOA factor than with BBOA (which constitutes only 9% of PM1, Table 1, Supplement S4) indicates that long-range transport of aged and internally well mixed biomass burning plumes plays a more important role than nearby sources (Darbyshire et al., 2019). The BBOA factor can be identified by the presence of the m/z 60 and m/z 73 (Figure 33a), which are dominated by the  $C_2H_4O_2^+$  and the  $C_3H_5O_2^+$  fragments. These fragments are originated from levoglucosan and other similar anhydro-sugars (such as manosan and galactosan). Levoglucosan (1,6- $\alpha$ -D-anhydroglucopyranose,  $C_6H_{10}O_5$ ) is known as a biomarker of biomass burning emissions due to its production from the pyrolysis of carbohydrates as cellulose (Alfarra et al., 2007; Artaxo et al., 2013; Chen et al., 2009; Lee et al., 2010). The signal fraction of m/z 60 for the BBOA factor in our study was 1.5%, which is 5 times higher than the 0.3% threshold typically used as an appropriate background fraction for biomass burning (Cubison et al., 2011). OOA presented a m/z 60 signal fraction of 0.2%, while IEPOX-SOA presented a negligible signal. The BBOA diurnal profile is different from that of the other PMF factors (Figure 4). While the OOA and IEPOX-SOA mass concentrations increase during the daytime due to photochemical oxidation processes, the BBOA mass concentration is fairly constant (Figure 4). Since the BBOA is a biomass-burning indicator, it is composed of mostly primary particles, so its concentration does not depend on photochemical activity. This pattern is different than the stark decrease in mass concentrations of fresh biomass burning particles during daytime, observed. The daily profile of BBOA differs significantly from other factors (Figure 4). While OOA and IEPOX-SOA mass loadings increase during the day, likely due to photochemically driven oxidation processes, BBOA remains relatively constant throughout the day, despite the daytime dilution effect of a rising boundary layer (Andreae et al., 2015). Interestingly, this pattern contrasts with the pronounced

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439 daytime decrease in fresh biomass-burning aerosol concentrations reported in southwestern Amazonia (Brito et al., 2014),  
440 where there were constant local sources of fires and the diurnal cycle was mostly determined by the boundary layer increasing  
441 and diluting the particles in a bigger area during daytime (Andreae et al., 2015). The lack of a clear diurnal pattern in our study  
442 for BBOA seems to confirm the regional origin of the aerosol particles, likely transported from distant biomass burning sources  
443 in the eastern parts of the basin, and long-range transport with complex local fire emissions were more prevalent. The absence  
444 of a clear diurnal cycle for BBOA in our study corroborates a regional, rather than local, origin—likely from biomass-burning  
445 sources located in the eastern Amazon. The flat variability of this primary factor reflects transport over long distances and the  
446 influence of complex vertical mixing, including interactions between residual and nocturnal layers (Darbyshire et al., 2019).  
447 An additional confirmation of the long-range transport  
448 Further supporting this hypothesis is the relatively flat pattern of the eBC diel cycle of eBC, although there is a small but  
449 noticeable slight daytime increase in the eBC diurnal mass concentration during daytime is observed (Figure 4), which may  
450 indicate some possibly due to lensing effect due to the increase in the particle coating effects as particles acquire coatings  
451 during transport (Denjean et al., 2020). While the diel cycle of the Unlike eBC,  $\text{NH}_4$  and  $\text{NO}_3$  show practically nominal  
452 diurnal variation, the while  $\text{SO}_4$  indicates the influence of exhibits a daytime increase, consistent with secondary production via  
453 photochemical processes (Figure 4) reactions from biogenic sources, or atmospheric transport processes. The higher rise in the  
454 boundary layer during the afternoon (Fisch et al., 2004) may favor the downward transport of long-distance particles (Fisch  
455 et al., 2004) may facilitate the entrainment of particles from above the boundary layer (Darbyshire et al., 2019). An additional  
456 possible explanation for this observed increase in  $\text{SO}_4$  during the afternoon is biogenic sources of  $\text{SO}_4$  precursors.  
457 As the OOA and the IEPOX SOA factors represent together around 68% of the total mass fractions of the submicron particles  
458 during our study, and conversely, eBC and BBOA represent only 15%, the importance of the atmospheric photochemical  
459 activity in Central Amazonia becomes evident. Well-preserved parts of the Amazon are strongly affected by the regional  
460 transport of well-processed biomass burning plumes, overwhelming the local biogenic processes that usually modulate the  
461 diurnal behavior of secondary aerosol development (Artaxo et al., 2013; Darbyshire et al., 2019).

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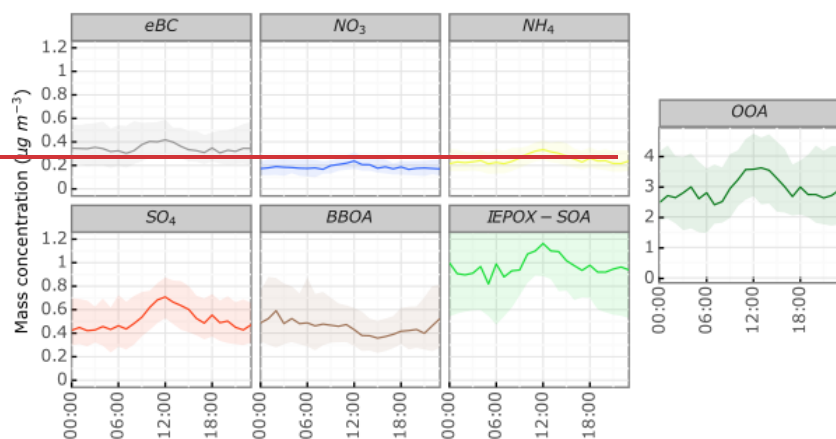
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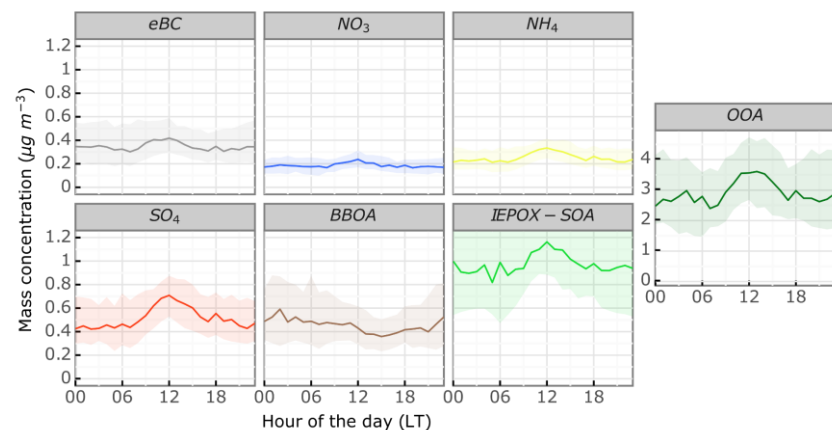
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As the OOA and the IEPOX-SOA factors represent together around 68% of the total mass fractions of the submicron particles during our study (Table 1), and conversely, eBC and BBOA represent only 15%, the importance of the atmospheric photochemical activity in Central Amazonia becomes evident. Well-preserved parts of Amazonia are strongly affected by the regional transport of well-processed biomass burning plumes, overwhelming the local biogenic processes that usually modulate the daily behavior of secondary aerosol development (Artaxo et al., 2013; Darbyshire et al., 2019).



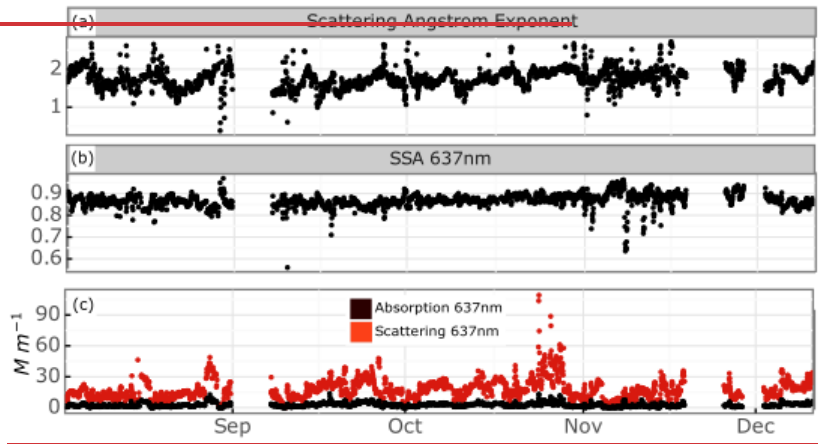
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469 **Figure 4: Diurnal profile (local time) of the PMF derived statistical factors, the inorganic chemical species and eBC mass**  
470 **concentrations for the whole period of measurements (1 August – 10 December, 2013). The lines represent mean values, and the**  
471 **shaded areas represent the standard deviations. The OOA factor, shown separately, has a different vertical scale to improve**  
472 **visualisation.**

473 **3.2 Physical properties**

474 The mean scattering coefficient at 637 nm in our study was  $17 \pm 10 \text{ Mm}^{-1}$  (Table 2), which is similar to the values reported for  
475 the same site and at the ATTO site during the dry season in previous years (Rizzo et al., 2013) and lower than observations  
476 close to biomass burning sources ( $32\text{--}80 \text{ Mm}^{-1}$  (Artaxo et al., 2013; Ponczek et al., 2021)). In the dry season, fine mode particles  
477 predominate and are more efficient at scattering radiation than coarse mode dominated biogenic particles in the wet season  
478 (Rizzo et al., 2013). The absorption coefficient mean value was  $3 \pm 2 \text{ Mm}^{-1}$  (Table 2, Figure 5, Section 2.2) (Rizzo et al., 2013).  
479 The  $\sigma_a$  mean value was  $3 \pm 2 \text{ Mm}^{-1}$  (Table 2, Figure 5c, Section 2.2), in accordance with low values previously reported for  
480 aged biomass burning haze (Formenti et al., 2003).



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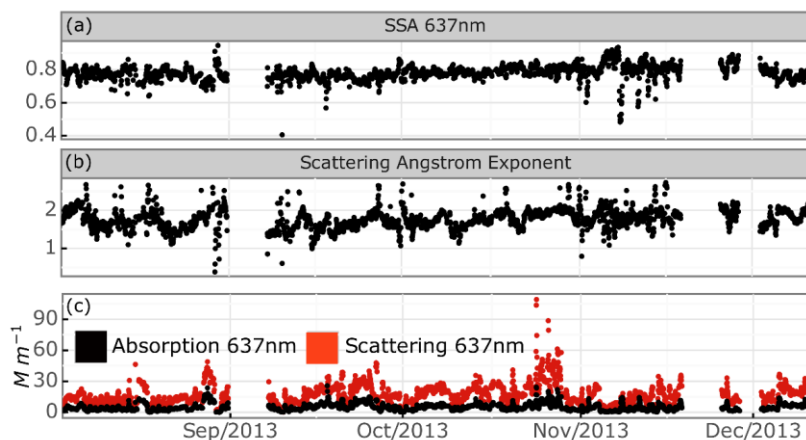
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**Figure 5: Time series of: a) the Scattering Angstrom Exponent (unitless), b) Single Scattering Albedo (SSA) at 637 nm (unitless), and c) absorption and scattering coefficients at 637 nm ( $Mm^{-1}$ ).**

The mean SSA observed in our study ( $0.87 \pm 0.03$ , Table 2) was very similar to the SSA reported for the nearby ATTO site (Saturno et al., 2018b). In previous years (2009–2012) at the same site as in our study, SSA varied from 0.84 to 0.91 from the wet to the dry season respectively. The dominance of organics and the relatively high  $SO_4$  fraction in our study (9%, Table 1) are probably important factors for the high SSA (Artaxo et al., 2013; Rizzo et al., 2013). SSA was lower ( $0.77 \pm 0.08$  at 637 nm) during the transition from dry to wet season in a site highly impacted by fires in southwestern Amazonia (Ponczek et al., 2021).

The mean value for the scattering Angstrom The mean SSA observed in our study ( $0.87 \pm 0.03$ , Table 2) was very similar to the SSA reported for a nearby site in Amazonia, as well as sites impacted by fires or urban pollution (Carrico et al., 2003; Deng et al., 2016; Kim, 2015; Kleinman et al., 2020; Nakayama et al., 2010; Saturno et al., 2018b; Wang et al., 2017; Zhu et al., 2015). The dominance of organics and the relatively high  $SO_4$  fraction in our study (9%, Table 1) are probably important factors contributing to the high SSA (Artaxo et al., 2013; Rizzo et al., 2013), and aged biomass burning plumes have been demonstrated to be more efficient in scattering radiation than freshly emitted particles (Formenti et al., 2003). Mean SSA was lower ( $0.77 \pm 0.08$  at 637 nm) in a site highly impacted by fires in southwestern Amazonia (Ponczek et al., 2021). In urban environments impacted by pollution, SSA was 0.92–0.89 (Tian et al., 2022), and 0.75–0.84 when the urban pollution was mixed with biomass burning (Pani et al., 2023). Slightly higher SSA values are related to urban haze episodes with high AS contributions, rural areas dominated by dust plumes, or high altitude regions influenced by clean maritime air masses (Fan et al., 2010; Han et al., 2015; Park et al., 2019). Lower SSA values were influenced by highly absorbing urban pollution (Andreae

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et al., 2008; Cho et al., 2017; Gao et al., 2015; Jing et al., 2015; Ma et al., 2011; Ram et al., 2016; Soni et al., 2010; Titos et al., 2012) or maritime regions impacted by biomass burning from Africa (Dobracki et al., 2023). The mean value for the scattering Angström exponent in our study was  $1.76 \pm 0.26$  (Table 2), which is very similar to the  $1.70 \pm 0.41$  and the  $1.71 \pm 0.24$  measured in the dry seasons of previous years at the same site and at the nearby ATTO station (Rizzo et al., 2013; Saturno et al., 2018b)(Rizzo et al., 2013; Saturno et al., 2018b), and the  $1.65 \pm 0.37$  measured during the dry season at a site more impacted by forest fires (Ponczek et al., 2021)(Ponczek et al., 2021). However, it was higher than the  $1.48 \pm 1.12$  (although within the high variability range) and the  $1.29 \pm 0.50$  measured in wet seasons of previous years at the same station and the ATTO site (Rizzo et al., 2013; Saturno et al., 2018b)(Rizzo et al., 2013; Saturno et al., 2018b). Higher scattering Angström exponent values are usually related to a greater proportion of fine mode particles in the aerosol population (Andreae et al., 2015), and in our case, it is probably related to the occurrence of fresh biomass burning particles.

**Table 2 – Optical properties mean and standard deviation (in parentheses) for the whole study period for different wavelengths.**

Optical property	Wavelength (nm)	Mean
Scattering coefficient ( $\text{Mm}^{-1}$ )	450	$32 \pm 19$
Scattering coefficient ( $\text{Mm}^{-1}$ )	550	$22 \pm 13$
Scattering coefficient ( $\text{Mm}^{-1}$ )	637	$17 \pm 10$
Scattering coefficient ( $\text{Mm}^{-1}$ )	700	$14 \pm 8$
Absorption coefficient ( $\text{Mm}^{-1}$ )	637	$3 \pm 2$
Single Scattering Albedo ( $\text{Mm}^{-1}$ )	637	$0.87 \pm 0.03$
Scattering Angström Exponent		$1.76 \pm 0.26$

### 3.3 Chemical-dependent optical properties

We applied the multiple linear regression (Section 2.4.2) to our dataset and the resulting coefficients successfully predicted the observed scattering (Figure 6), confirming the validation of this methodology to estimate the specific contribution of each chemical group to the optical properties. All the coefficients of the multilinear regression of all the wavelengths were statistically significant ( $p < 0.001$ ) for both MSE and MEE. The MSE of the PMF factors decreased as a function of wavelength (Figure 7), in agreement with a previous study (Ponczek et al., 2021). Regarding the inorganic species, this decrease was also observed in the AN, but absent in the AS, which showed no wavelength dependency (Figure 7). The highest MSE values were attributed to eBC (Table 3, Figure 7a), followed by the BBOA. The MSE values of the eBC, BBOA and OOA components calculated in our study at 637 nm were circa 180%, 67% and 43% respectively compared to (Ponczek et al., 2021). MSE of

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AS in our study was between  $4.58\text{--}4.79\text{ m}^2\text{ g}^{-1}$  (considering wavelengths from 450–700 nm, Table 4), which is almost double of the  $2.5\pm0.6\text{ m}^2\text{ g}^{-1}$  fine mode average of 93 observations of a variety of regions (urban, remote, rural continental, ocean/marine) and seasons, and using different methods, mostly at 550 nm (Hand and Malm, 2007). Despite the small variability, lower MSE was reported for drier and cleaner environments, and higher MSE was found in more polluted regions and larger particle sizes (Hand and Malm, 2007), which may partially explain the higher values found in our study—in a wet climate and during the forest fires season when the atmosphere can resemble that of highly polluted regions (Artaxo et al., 2013). A recent study in an area highly impacted by urban pollution in France reported MSE for AS ranging from  $4.8\text{--}7.1\text{ m}^2\text{ g}^{-1}$  (450–635 nm) (Velazquez-Garcia et al., 2023), resembling more similar results to our study for the lower wavelengths. Concerning the contribution of AN to the PM1 mass concentration, we tested the MLR removing AN, and the results were comparable, especially for eBC (Supplement Table S5.1). We also tested the robustness of the method by running 100 times MLR on random 50% of the data, yielding similar results (Supplement Table S5.2).

Aerosol particles which typically absorb radiation are also known to have significant scattering efficiencies, highly dependent on their sizes (Bond and Bergstrom, 2006). Biomass-burning aerosols have been previously associated with high scattering efficiencies (Hand and Malm, 2007; Malm et al., 2005). However, no clear particle size dependency was observed for the radiation scattering in our study (Figure 6). Our result of a pronounced MSE of the eBC represents an opposite trend than the observed in the transition from the dry to the wet season at a site more impacted by the fires, where the MSE of BBOA was higher than eBC's (Ponczek et al., 2021).

When the mass concentration is considered, the relative contribution of eBC to the scattering in all the measured wavelengths is about 20–25% or the total scattering (Figure 7b), comparable to Southwestern Amazonia (Ponczek et al., 2021), despite our site having a significantly lower eBC concentration. The contribution of AS and AN to MSE was from 20% to 30% with increasing wavelength (Figure 7b), less than half of that in urban sites in Europe (e.g. 67% in Northern France, (Velazquez-Garcia et al., 2023), but about twice as high as during an extreme pollution haze episode in Beijing (Wang et al., 2015).

We applied the multiple linear regression (Section 2.4.2) to our dataset, and the resulting coefficients successfully predicted the observed scattering ( $R^2 = 0.86$ , Figure 6), confirming the validation of this methodology to estimate the specific contribution of each chemical group to the optical properties. We tested the MLR removing AN (due to its low contribution to the PM1 mass concentration, close to the ACSM detection limit, and therefore, possible artifacts), and the results were comparable, especially for eBC (Supplement Table S5.1). We also tested the robustness of the method by running 100 times MLR on randomly selected 50% of the data, yielding similar results (Supplement Table S5.2). All standard errors were small (Table 3), and the Variance Inflation Factor was around 3 for IEPOX-SOA, BBOA, AS and AN; 5.20 for OOA, and 6.19 for eBC. The abovementioned tests suggest that typical MLR caveats such as collinearity had minimal effect on the observed final results. No clear particle size dependency was observed for the radiation scattering in most of the cases (regression fitting under typical conditions, of aerosol sizes in the range of 100–150 nm), except at events dominated by ultra fine particles, at around 50 nm (Figure 6). This is notably an underestimation of observed scattering at lower particle diameters.

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561 The highest MSE values were attributed to eBC (Table 3, Figure 7a), followed by the BBOA. Previous studies on plumes  
562 dominated by either urban pollution or mixed with biomass burning presented MSE around 4.4 m<sup>2</sup> g<sup>-1</sup>, dominated by organic  
563 particles (Cheng et al., 2015; Pani et al., 2023; Tao et al., 2019). The MSE values of the eBC, BBOA and OOA components  
564 calculated in our study at 637 nm were circa 180%, 67% and 43% respectively compared to previous measurements in  
565 Amazonia, highly impacted by biomass burning (Ponczek et al., 2021). MSE of AS in our study was between 4.58-4.79 m<sup>2</sup> g<sup>-1</sup>  
566 (Table 3, Figure 7a), in very good agreement with the MSE described for fine-mode ambient AS particles in an urban  
567 environment (Tao et al., 2019). Our result is in the lower range of the MSE described in regions impacted by urban pollution  
568 (4.8-7.1 m<sup>2</sup> g<sup>-1</sup>, (Velazquez-Garcia et al., 2023), probably due to the smaller mean diameter found in our study (Figure 6).  
569 However, other regions (urban, remote, rural continental, ocean/marine) presented much smaller MSE values for AS (Cheng  
570 et al., 2015; Hand and Malm, 2007). MSE for AN at 550 nm in our study (4.79 m<sup>2</sup> g<sup>-1</sup>, Table 3) is in very good agreement with  
571 the MSE found in AN in a urban pollution plume (Tao et al., 2019), and within the range previously described in regions highly  
572 impacted by urban pollution (Cheng et al., 2015; Tian et al., 2022) and a mixture of urban pollution and biomass burning (Pani  
573 et al., 2023).  
574 The organic particles presented higher MSE for freshly emitted aerosols (BBOA) than for oxygenated particles (OOA) (Table  
575 3), an opposite trend to what was found at PM<sub>2.5</sub> in a region impacted by urban pollution (Tian et al., 2022). However, previous  
576 studies in Amazonia demonstrated that the size distribution of the particles is mainly below 200 nm, and even aging processes  
577 do not appear to cause an overall increase in total particles diameter, probably due to the type of the vegetation, the precursors  
578 of SOA, disintegration of larger particles, and other factors (Artaxo et al., 2013; Brito et al., 2014). Fresh biomass burning  
579 plumes at 532 nm presented a MSE range of 1.5-5.7 m<sup>2</sup> g<sup>-1</sup>, depending on the fuel type, and plume age (Levin et al., 2010),  
580 and the MSE of BBOA found in our study for 550 nm is 5.33 m<sup>2</sup> g<sup>-1</sup> (Table 3). A review of MSE biomass burning plumes  
581 revealed higher MSE values for more aged plumes (Reid et al., 2005). Fine-mode organic aerosols in an urban environment  
582 presented a mean MSE of 4.6 m<sup>2</sup> g<sup>-1</sup> at wavelength 550 nm (Tao et al., 2019), closer to our BBOA MSE (Table 3).  
583 The pronounced MSE of the eBC (7.62-13.58 m<sup>2</sup> g<sup>-1</sup>, Table 3) is strongly corroborated by other studies which found remarkably  
584 high scattering efficiency related to BC, especially when the particles undergo atmospheric processing and aging, such as in  
585 the case of our study (Bond and Bergstrom, 2006; He et al., 2015; Malm et al., 2005; Pitchford et al., 2007; Romshoo et al.,  
586 2021; Schwarz et al., 2006). It has been demonstrated that while aerosol scattering efficiency increases with increasing size,  
587 age and distance from the source, the absorption efficiency remains nearly constant (Kleinman et al., 2020; Zhang et al., 2020).  
588 MSE of elemental carbon in a rural area ranged from 5.4-66.2 m<sup>2</sup> g<sup>-1</sup>, and the high increase was found to be related to sulfate  
589 addition during cloud processing (Yu et al., 2010). Recently, on a comparable method, MSE for eBC has been estimated at 6  
590 m<sup>2</sup> g<sup>-1</sup> in a site located in Western Amazonia. Located within the deforestation arc, the site is strongly impacted by fresh,  
591 sometimes local emissions, in contrast to regional or long-range transport of fires impacting Central Amazonia (Ponczek et  
592 al., 2021). In regions impacted by urban pollution MSE of eBC was 2.6 m<sup>2</sup> g<sup>-1</sup> (Tao et al., 2019), and found not to influence  
593 MSE for coarse mode particles (Titos et al., 2012).

594 When the mass concentration is considered, the relative contribution of eBC to the scattering in all the measured wavelengths  
595 in our study is about 20-25% or the total scattering (Figure 7b), comparable to Southwestern Amazonia (Ponczek et al., 2021),  
596 despite our site having a significantly lower eBC concentration (but higher MSE for eBC). In this same Southwestern  
597 Amazonia site, the contributions of the OOA and BBOA to MSE were about twice as high than in our study. The contribution  
598 of AS and AN to MSE was from 20% to 30% with increasing wavelength (Figure 7b), less than half of that in urban sites in  
599 Europe (e.g. 67% in Northern France, (Velazquez-Garcia et al., 2023), but about twice as high as during an extreme pollution  
600 haze episode (Wang et al., 2015)). As shown in Figure 7a, the MSE of all components except AS decreases with increasing  
601 wavelength, which is consistent with the typical behavior of submicrometric aerosols. This spectral dependence can be  
602 attributed to Mie scattering theory, where smaller particles scatter shorter wavelengths more efficiently (Hand and Malm,  
603 2007; Malm et al., 2005). Nonetheless, the variability in the MSE slopes among the different components reflects a complex  
604 interplay between aerosol mixing state, refractive index, and size distribution dynamics—particularly the diurnal evolution of  
605 each factor's contribution to the total aerosol population (Figure 4). It is particularly interesting that AS exhibits a distinct  
606 spectral behavior, typically associated with coarse-mode aerosols, denoting stark differences in its sources and atmospheric  
607 processing compared to the other components. Sulfate in Amazonia has been associated with secondary production from  
608 biogenic emissions and mixing with primary biogenic organic aerosols (PBOA) (Martin et al., 2010b; Pöhlker et al., 2012), as  
609 well as with coarse-mode particles such as dust and sea salt transported over long distances (Brito et al., 2014; Wu et al., 2019).  
610 It is remarkable that the MLR analysis captured this behavior, considering that the ACSM is limited to non-refractory species  
611 in the submicron range and is not particularly efficient at detecting the sources likely involved. This highlights the sensitivity  
612 of the MLR approach to broader aerosol population dynamics, which were captured by the optical instruments operating with  
613 a PM<sub>10</sub> inlet, suggesting the influence of coarse-mode aerosol sources.

614 While the MSE of eBC does decrease with increasing wavelength (Figure 7a), its slope (or more precisely, its SAE) is lower  
615 than that of other aerosol components. As a result, eBC retains a relatively higher fractional contribution to total scattering at  
616 longer wavelengths compared to components with steeper MSE declines (Figure 7b). The absolute contribution to scattering  
617 is determined by both MSE and mass concentration, and although eBC mass concentrations are generally lower, its weaker  
618 wavelength dependence allows it to contribute proportionally more at longer wavelengths.

619 When considering the total light extinction (scattering + absorption), the relative contribution of eBC reaches about 30.31%  
620 (Figure 8), which is comparable to the work in highly urbanized region in Europe (Velazquez-Garcia et al., 2023), however  
621 significantly lower than the 76±20% observed in urban pollution in China (Yu et al., 2010), and an episodic biomass burning  
622 event in a rural area (Yu et al., 2010). However, it is less than half of the MEE relative contribution of BC observed during  
623 urban pollution episodes (Tian et al., 2022; Yu et al., 2010). The comparison with urban pollution particles contribution to  
624 MEE reveals that the contribution of highly oxygenated particles is very similar (circa 20%, Figure 8, (Tian et al., 2022)), and  
625 the most evident difference is the nitrate-based particles, with a much larger contribution in the urban pollution region (Figure  
626 8, (Tian et al., 2022)). MEE has been shown to increase by a factor of 3 while freshly emitted smoke from fires ages in the  
627 atmosphere, reaching up to 7m<sup>2</sup> g<sup>-1</sup> at 532 wavelength (Saide et al., 2022). The OOA factor presented a relatively high

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628 contribution to MSE and MEE (Figures 8b and 9) due to its high fraction of the total PM1 mass (Table 1), although its MSE  
629 is relatively low (Figure 7a). The contribution of AS to MSE increases with increasing wavelength (from 10% to 20%, Figure  
630 7b), while OOA decreased (from 30% to 20%, Figure 7b).  
631 By using the MSE and MEE ratios, we calculated specific SSA for the eBC, obtaining a value of 0.57. This means that 57%  
632 of the light extinction provoked by the eBC is scattered rather than absorbed, which is higher than the eBC specific SSA of  
633 0.46 based on previous studies in more polluted conditions (Ponczek et al., 2021; Velazquez-Garcia et al., 2023). SSA of eBC  
634 has been described as typically ranging from 0.3 to 0.4, with higher values associated to heavy coating (Luo et al., 2020).  
635 Evidence of coating increasing elemental carbon scattering efficiency has been found in the past (He et al., 2015; Yu et al.,  
636 2010), however other studies found that thick coating in eBC lowered scattering coefficients (Darbyshire et al., 2019), and  
637 below 0.3 for pure black carbon (Wang et al., 2021).  
638 A previous study found that eBC absorption cross-section for the Amazon was  $12.3 \text{ m}^2 \text{ g}^{-1}$  (Saturno et al., 2018b), and we  
639 tested our dataset applying this value. The result is that eBC mass concentrations would become half of what they are reported,  
640 with no change in  $\sigma_a$ , SSA, but MSE would double, while eBC contribution to MEE (Figure 8) would remain unchanged. Due  
641 to some methodology differences between our study and (Saturno et al., 2018a) (they measured refractory Black Carbon using  
642 a single particle soot photometer SP2, with a higher cut-off, possibly leading to a sub-estimation of the mass), and the fact that  
643 applying the absorption cross-section value they found would make MSE of eBC be an order of magnitude higher than the  
644 others (Supplement Figure S6), we opted to remain with the more established value of  $6.6 \text{ m}^2 \text{ g}^{-1}$ .  
645 The OOA factor presented a relatively high relative contribution to MSE and MEE (Figures 8b and 9) due to its high mass  
646 relative contribution to the total PM1 mass (Table 1), although its MSE is relatively low (Figure 7a). Our results show an  
647 increase in AS relative fraction of contribution to MSE with increasing wavelength (from 10% to 20%, Figure 7b), while OOA  
648 decreased (from 30% to 20%, Figure 7b). The relative contributions to MSE of the OOA and BBOA were about twice as high  
649 in a site more directly impacted by fires during the dry season than in our study (Ponczek et al., 2021).  
650 Table 3—MEE, MAE and MSE for different wavelengths and aerosol components with standard errors. All the  
651 coefficients were statistically significant for all the wavelengths for MSE and MEE ( $p < 0.001$ ).———  
652 A previous study found an eBC absorption cross-section in Amazonia of  $12.3 \text{ m}^2 \text{ g}^{-1}$  (Saturno et al., 2018b), and we tested our  
653 dataset applying this value (Supplement, Figure S6.1). The result is that eBC mass concentrations would decrease by half, with  
654 no change in  $\sigma_a$  and SSA, but MSE would double, while eBC contribution to MEE (Figure 8, Figure S6.2) would remain  
655 unchanged. Due to some methodology differences between our study and (Saturno et al., 2018b) (they measured refractory  
656 Black Carbon using a single-particle soot photometer SP2, with a higher cut-off, possibly leading to a sub-estimation of the  
657 mass), and the fact that applying the absorption cross-section value they found would make MSE of eBC be an order of  
658 magnitude higher than the others (Supplement Figure S6.1), we opted to remain with the more established value of  $6.6 \text{ m}^2 \text{ g}^{-1}$ .  
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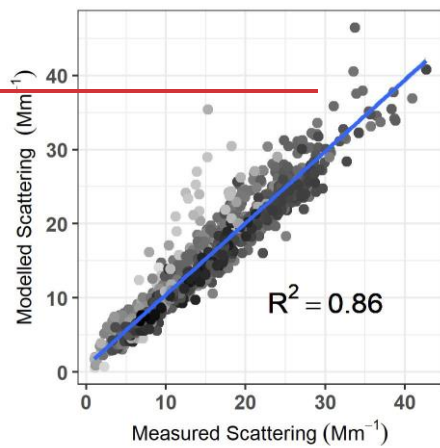
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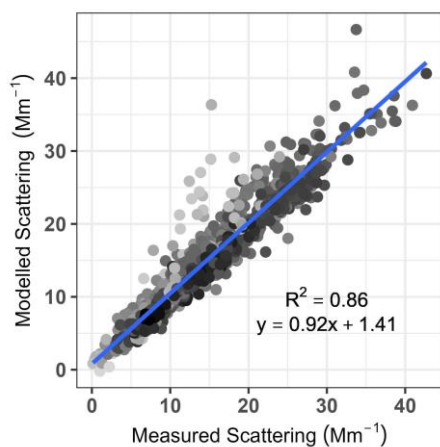
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$D_{pg}$  (nm) 50 100 150 200



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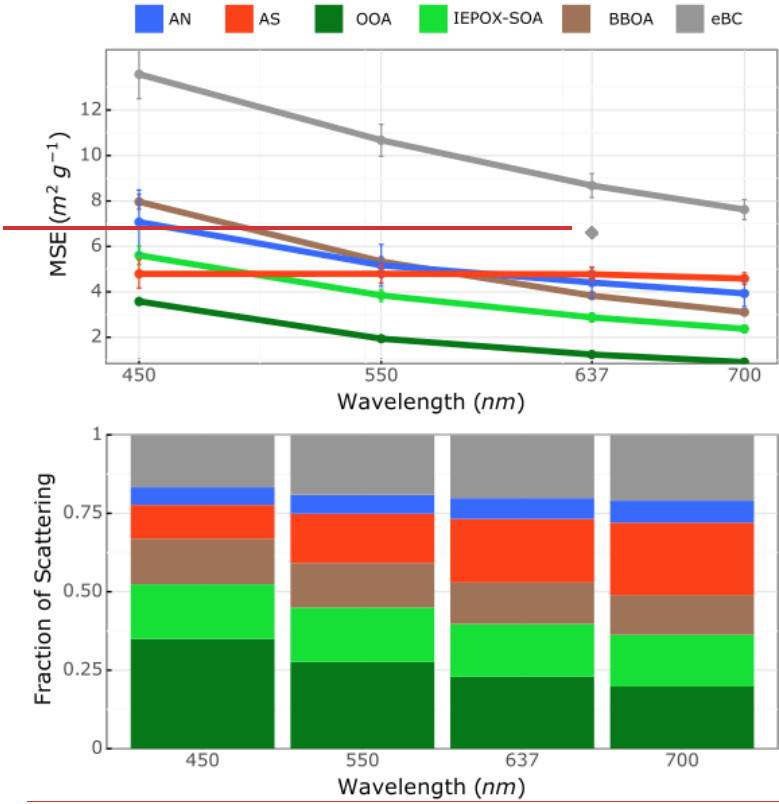


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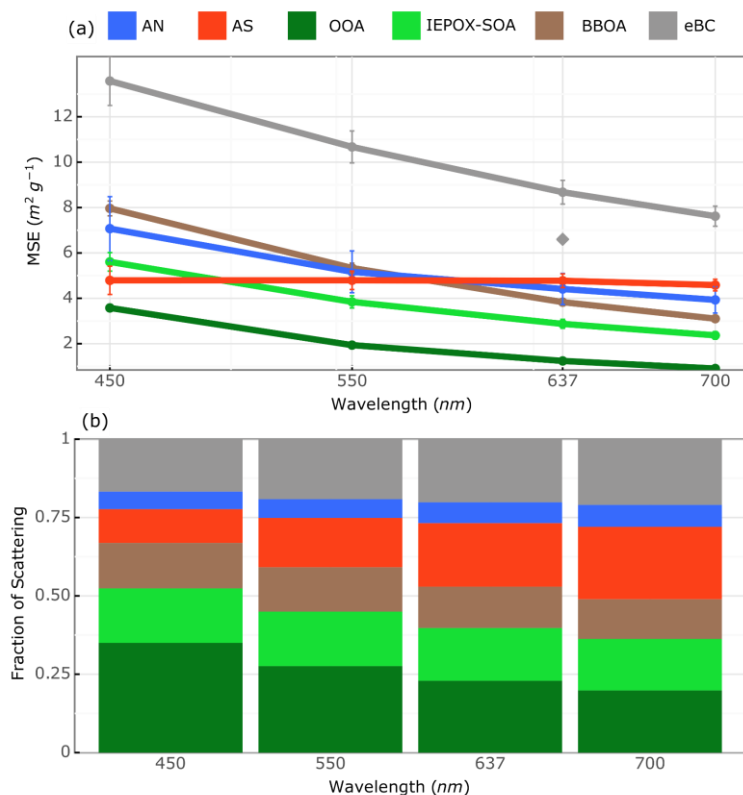
Figure 6: Measured radiationys modelled aerosol light scattering at 637 nm vs modeled scattering (MSE). The gray scale corresponds to the equivalent mobility particle diameter for singly charged particles (D<sub>pg</sub>, section 2.2). The blue line indicates ashows the linear correlation regression (slope =0.92).



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**Figure 7:** a) Mass scattering efficiencies (MSE) for each of individual chemical component (color components (colored lines) at each wavelength, where the dots are the coefficients obtained from the multi-linear regression, the with bars are indicating the standard errors, and the diamond shape is denotes the value of absorption efficiency of the eBC at 637 nm; and b) the fraction-fractional contribution of each component to total light scattering of each component at each wavelength, considering its mass fraction to the total mass of submicrometer particles.

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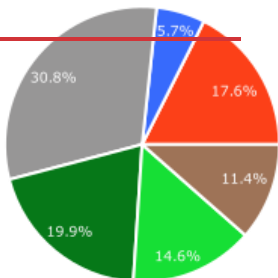
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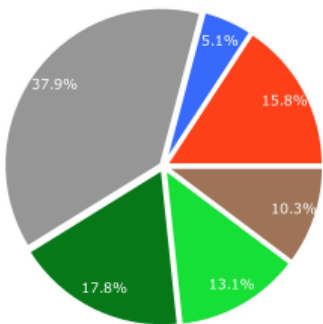
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OOA IEPOX-SOA BBOA  
AN AS eBC



OOA IEPOX-SOA BBOA  
AN AS eBC



**Figure 8: The relative contribution (%) of extinction-coefficient (scattering + absorption)MEF of each component considering its mass fraction to the total mass of submicrometer particles: (Table 1).**

#### 4 Conclusions

The results have shown that the submicrometer particle mass concentration during the dry season ( $6.3\pm3.3\text{ }\mu\text{g m}^{-3}$ ), which is about an order of magnitude higher than typically observed at this site during the wet season, is highly dominated by the organic fraction ( $77\pm5\%$ ). The organic aerosols were separated into 3 PMF statistical factors, identified as BBOA, OOA, and IEPOX-SOA. The OOA, associated with highly processed and oxidized particles is the dominant factor ( $51\pm6\%$  of the  $\text{PM}_{10}$ ),

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690 followed by IEPOX-SOA (isoprene SOA following a low-NO route,  $17\pm 5\%$ ), while the factor more directly associated with  
691 fresh biomass burning emissions (BBOA) represents  $9\pm 5\%$  of PM<sub>1</sub>.  
692 The mean radiation scattering coefficient at 637 nm was  $17\pm 10$  Mm<sup>-1</sup>, and the mean absorption coefficient was  $3\pm 2$  Mm<sup>-1</sup>,  
693 which ~~are-is higher than what was observed in the wet season, but~~ lower than ~~the respective coefficients measured at~~ sites  
694 much more impacted by the proximity to fresh forest fires ~~but exceed those observed during the wet season~~. The SSA of  
695  $0.87\pm 0.03$  was elevated compared to values previously described in the wet season, but generally in good agreement with SSA  
696 values of previous dry seasons ~~- likely an indication of the scattering efficiency of eBC following atmospheric processing and~~  
697 ~~coating~~. The mean scattering ~~Angstrom~~Angström exponent was  $1.76\pm 0.26$ , surpassing the values previously measured during  
698 the wet season, likely due to the greater relative proportion of smaller particles in the aerosol population measured in our study,  
699 compared to the primary biogenic aerosols, Saharan dust and sea salt typical of the wet season in central Amazonia.  
700 An MLR analysis of optical properties and different species/factors yielded the highest MSE for eBC in all wavelengths (~~7.62-~~  
701 ~~13.58 m<sup>2</sup> g<sup>-1</sup>~~), followed by the BBOA and AN in the shorter wavelengths (450 and 550 nm), ~~with the addition of and by~~ AS in  
702 the longer wavelengths (637 and 700 nm). Despite having a small mass contribution (6%), eBC dominated the MEE relative  
703 contribution (~~30.837.9%~~), followed by the OOA (~~49.917.8%~~). The high MSE of eBC (and a calculated specific SSA of 0.57)  
704 is ~~surprising~~remarkable, and more studies concerning the chemical processing of those particles after their emission are needed  
705 in order to understand the processes behind the outstanding efficiency of light scattering of this highly absorbing particle. A  
706 special focus should be given to the role of aerosol coating in this process.  
707 ~~While previous studies show a significant contribution of organic particles to light~~Aerosol observations at the heart of  
708 ~~Amazonia are extremely challenging. Although our analysis during the dry season yielded robust results, limitations remain —~~  
709 ~~particularly concerning the size distribution of BC and its absorption (brown carbon) our study shows that the eBC can also~~  
710 ~~have a significant contribution to light across multiple wavelengths. To advance our understanding, future studies should~~  
711 ~~prioritize extensive field and laboratory observations aimed at better constraining aerosol coating formation mechanisms and~~  
712 ~~their impact on the radiative scattering. The properties of BC particles in Amazonia. Increasing the precision of the~~  
713 quantification of the eBC contribution to light scattering has the potential to improve models and decrease uncertainties in  
714 global radiative forcing estimations.

715 **Code availability**

716 **Codes and Data availability**

717 Data will be madeCodes and data are currently available upon acceptance of the article in this link;  
718 <https://zenodo.org/records/15345166>; DOI: 10.5281/zenodo.15345166

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**Author Contribution**

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**Competing Interests**

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**Acknowledgments**

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