

# 1 **Understanding aerosol composition in a tropical inter-Andean valley** 2 **impacted by agro-industrial and urban emissions**

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## 14 **Abstract.**

15 Agro-industrial areas are frequently affected by various sources of atmospheric pollutants that have a negative impact on public  
16 health and ecosystems. However, air quality in these areas is infrequently monitored because of their smaller population  
17 compared to large cities, especially in developing countries. The Cauca River Valley (CRV) is an agro-industrial region in  
18 Southwest Colombia, where a large fraction of the area is devoted to sugarcane and derivative production. The CRV is also  
19 affected by road traffic and industrial emissions. This study aims to elucidate the chemical composition of particulate matter  
20 fine mode (PM<sub>2.5</sub>) and to identify the main pollutant sources before source attribution. A sampling campaign was carried out  
21 at a representative site in the CRV region, where daily-averaged mass concentrations of PM<sub>2.5</sub> and the concentrations of water-  
22 soluble ions, trace metals, organic and elemental carbon, and various fractions of organic compounds (carbohydrates, n-  
23 alkanes, and polycyclic aromatic hydrocarbons – PAHs) were measured. The mean PM<sub>2.5</sub> was  $14.4 \pm 4.4 \mu\text{g m}^{-3}$ , and the most  
24 abundant constituent was organic material ( $52.7\% \pm 18.4\%$ ), followed by sulfate ( $12.7\% \pm 2.8\%$ ), and elemental carbon ( $7.1\%$   
25  $\pm 2.5\%$ ), which indicates secondary aerosol formation and incomplete combustion. Levoglucosan was present in all samples  
26 with a mean concentration of ( $113.8 \pm 147.2 \text{ ng m}^{-3}$ ) revealing biomass burning as a persistent source. Mass closure using the  
27 EC tracer method explained 88.4% of PM<sub>2.5</sub>, whereas the organic tracer method explained 70.9% of PM<sub>2.5</sub>. We attribute this  
28 difference to the lack of information of specific organic tracers for some sources, both primary and secondary. Organic material

29 and inorganic ions were the dominant groups of species (79% of  $PM_{2.5}$ ).  $OM_{prim}$  and  $OM_{sec}$  contribute 24.2% and 28.5% to  
30  $PM_{2.5}$ . Inorganic ions as sulfate, nitrate and ammonia constitute 19.0%; EC, 7.1%; dust, 3.5%; PBW, 5.3%; and TEO, 0.9% of  
31  $PM_{2.5}$ . The aerosol was acidic, with a pH of  $2.5 \pm 0.4$ , mainly because of the abundance of organic and sulfur compounds.  
32 Diagnostic ratios and tracer concentrations indicate that most  $PM_{2.5}$  was emitted locally and had contributions of both  
33 pyrogenic and petrogenic sources, that biomass burning was ubiquitous during the sampling period and was the main source  
34 of PAHS, and that the relatively low  $PM_{2.5}$  concentrations and mutagenic potentials are consistent with low-intensity, year-  
35 long BB and sugarcane PHB in CRV.

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38 Keywords: agro-industry; pre-harvest burning;  $PM_{2.5}$ ; chemical speciation; Northern South America

39

## 40 1. Introduction

41 Urban and suburban locations, with moderate to high population densities, are exposed to air pollutant emissions, including of  
42 fine particulate matter (PM) from industry, road traffic, and other anthropogenic activities. Suburban areas may also be  
43 impacted by emissions from agricultural activities (Begam et al., 2016). Air quality in areas under these conditions is  
44 infrequently monitored, particularly in developing countries, despite the extensive use of highly emitting practices, including  
45 intensive use of insecticides and pesticides, fire for land and crop management, and diesel-based mechanization (Aneja et al.,  
46 2008, 2009). Agricultural sources emit pollutants, such as volatile organic compounds (VOC), which are precursors of  
47 tropospheric ozone (Majra, 2011) and secondary organic aerosols (SOA) (Majra, 2011). Most agricultural activities also emit  
48  $PM_{2.5}$  (solid and liquid particles with aerodynamic diameters smaller than  $2.5 \mu m$ ), which may contain black carbon (BC) and  
49 toxic and carcinogenic pollutants, e.g., polycyclic aromatic hydrocarbons (PAHs). Other agricultural activities, including  
50 mechanized land preparation, sowing and harvesting, consume significant volumes of fossil fuels, particularly diesel, and emit  
51 trace gases (including  $CO_2$ , CO,  $SO_2$ ,  $NO_x$ ,  $NH_3$ , VOC) that also generate  $O_3$  and SOA, all of which affect human health and  
52 climate (Yadav and Devi, 2019). Furthermore, agricultural operations are a significant source of nitrogen-containing trace  
53 gases ( $NO_2$ , NO,  $NH_3$ ,  $N_2O$ ) that are released from fertilizers, livestock waste, and farm machinery into the atmosphere (Sutton  
54 et al., 2011). Also, poultry and pig farming are high emitters of sulfur compounds, particularly  $H_2S$ .

55

56 The Cauca River Valley (CRV) is an inter-Andean valley in Southwest Colombia with a flat area of  $5287 km^2$  (248 km long  
57 by 22 km mean width), a mean altitude of 985 m MSL (Figure 1). CRV is bounded by the Colombian Andes Western and  
58 Central Cordilleras, and is located at  $\sim 120 km$  from the Pacific Ocean. CRV encompasses the cities of Cali, Colombia's third-  
59 largest city with 2.2 million inhabitants (inhab), Yumbo (129 thousand inhab), an important industrial hub, and Palmira (313

60 thousand inhab), an important agro-industry center. Industry is also present in the other major CRV cities (Tuluá, Cartago,  
61 Jamundí, and Buga).

62

63 CRV hosts a highly efficient, resource-intensive sugarcane agro-industry with one of the highest biomass yields (up to 120 ton  
64  $\text{ha}^{-1}$ ) and the highest sugar productivities in the world ( $\sim 13 \text{ ton sugar ha}^{-1}$ ) (Asocaña, 2018, 2019). Sugarcane sowing,  
65 harvesting, and transport to mills are all mechanized and use diesel as fuel. Besides, all the sugarcane bagasse is used, either  
66 to produce heat and electric power (cogeneration) or as feedstock to the local paper industry. Moreover, although pre-harvest  
67 burning is being phased out in CRV, one-third of the sugarcane area in 2018 was burned prior to harvesting. CRV is also the  
68 third largest poultry producer (351,104 ton  $\text{yr}^{-1}$ ), and the first egg producer (4,559 million units per year) in Colombia  
69 (Min.Agricultura, 2020). In addition, CRV produces 15.1% of Colombia's pork meat (over 1 million pigs in stock)  
70 (Min.Agricultura, 2019) and 1.8% of national beef production (467,782 heads in stock) (Min.Agricultura, 2018). Poultry and  
71 livestock production are significant sources of  $\text{H}_2\text{S}$  and  $\text{NH}_3$ . Besides a long-time established energy-intensive industry, there  
72 are also a variety of smaller industries, including brick kilns. Regarding mobile sources, there are nearly 2 million vehicles  
73 (1,951,638 vehicles) registered in CRV (RUNT, 2021). These include the standard urban categories along with off-road  
74 unregulated farming machinery. The sugarcane agroindustry uses multi-car trailers towed by diesel-powered tractors, with  
75 enough annual activity to be considered an independent source (the activity of which is proportional to the sugarcane harvested  
76 area and the distance to sugar mills). Overall, CRV mobile sources consumed 772 million L of gasoline and 590 million L of  
77 diesel in 2018 (SICOM, 2018). Moreover, the local airport, the most important in southwest Colombia, located very close to  
78 Palmira, handled 1.3 million passengers in 2019 (Aerocivil, 2019). Also, 1657 ha of sugarcane and corn were fumigated in  
79 2020 using small aircraft (Aerocivil, 2020).

80

81 For this research, we prepared a preliminary, aggregated  $\text{PM}_{10}$  emission inventory for CRV by putting together disparate source  
82 data, including from the stationary source emission inventories of CRV's six largest cities (Cali, Tuluá, Cartago, Jamundí,  
83 Palmira, Yumbo and Buga), Cali's and other cities' mobile source emission inventories, and our estimation of sugarcane pre-  
84 harvest burning (PHB) and other point, linear and area sources (Table S1). Our preliminary inventory indicates that the  
85 manufacturing industry is by far the main  $\text{PM}_{10}$  emitter in CRV, with annual emissions of  $\sim 11.4 \text{ Gg PM}_{10}$ .  $\text{PM}_{10}$  emissions  
86 from mobile sources ( $\sim 1.4 \text{ Gg PM}_{10} \text{ yr}^{-1}$ ) and open-field sugarcane pre-harvest burning ( $1.7 \text{ Gg PM}_{10} \text{ yr}^{-1}$ ) are a factor  $\sim 5$   
87 smaller. The emissions of inorganic and organic secondary aerosol precursors are also significant. We estimate that 30.1 Gg  
88 of  $\text{SO}_2$  are annually emitted in CRV (41% from sugar mills and other agro-industries, 32% from food industries, and 9% from  
89 cement, ceramic, and asphalt production). Emissions of volatile organic compounds (VOCs) are very similar ( $34.7 \text{ Gg yr}^{-1}$ ).  
90 Although a significant number of coal-fired boilers have been converted to natural gas, CRV's sulfur-rich coal (1.4-4% total  
91 S) is still an important industrial fuel. It must be stressed that this is a preliminary, not fully updated, regional inventory. The

92 available information was insufficient for disaggregating the fine-mode PM emissions ( $PM_{2.5}$ ). The multiplicity, disparity, and  
93 uncertainty of sources are indicative of the complexity of the  $PM_{2.5}$  source identification, quantification, and location tasks.  
94

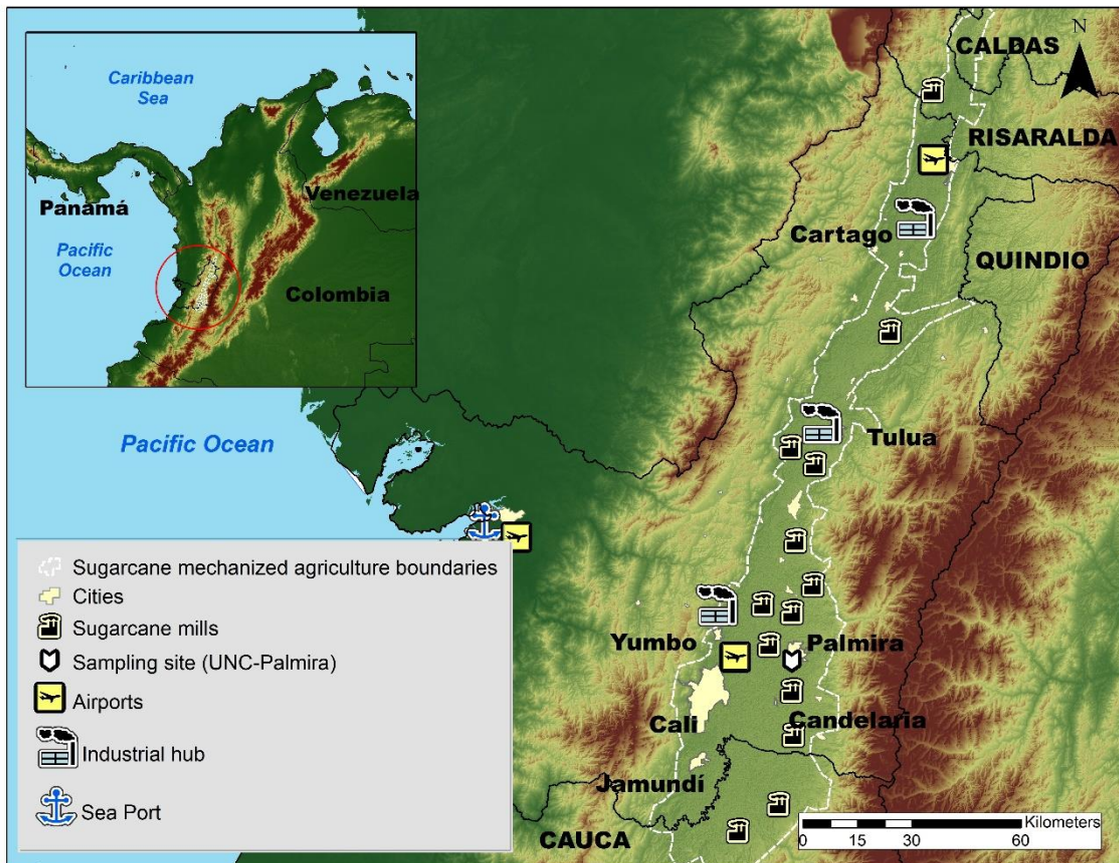
95 The determination of the particulate matter (PM) chemical composition is instrumental for the apportionment of pollutant  
96 sources. Most field measurement-based studies have been conducted in North America, Europe, and Asia (Karagulian et al.,  
97 2015). The number of studies in Latin America and the Caribbean (LAC) is much smaller and have focused on the chemical  
98 composition of  $PM_{10}$  (Pereira et al., 2019; Vasconcellos et al., 2011), as well as the PM source apportionment in urban areas  
99 of Colombia (Ramírez et al., 2018; Vargas et al., 2012), Chile (Jorquera and Barraza, 2012, 2013; Villalobos et al., 2015),  
100 Costa Rica (Murillo et al., 2013) and Brazil (de Andrade et al., 2010). The number of studies that involve agro-industrial  
101 sources and their impact on suburban areas is smaller. These include the Indo-Gangetic plain (Alvi et al., 2020), the Sao Paulo  
102 State in Brazil (Gonçalves et al., 2016; Urban et al., 2016), Ouagadougou in Burkina Faso (Boman et al., 2009), the Anhui  
103 Province in China (Li et al., 2014), for which the chemical composition of  $PM_{2.5}$  and some of its sources have been identified.  
104 Likewise, regions in South America with sugarcane agroindustry, such as Mexico (Mugica-Alvarez et al., 2015; Mugica-  
105 Álvarez et al., 2016, 2018) and Brazil (de Andrade et al., 2010; De Assuncao et al., 2014; Lara et al., 2005; Pereira et al., 2017)  
106 have also reported on their agro-industry impact on  $PM_{2.5}$  levels at nearby population centers. They are very few studies on air  
107 pollution in agro-industrial areas of Colombia. Most notably, Romero et al., (2013) measured PAHs and metals in  $PM_{10}$ . Most  
108 of the studies above identified biomass burning and fossil fuel combustion as significant PM sources, and some also identified  
109 industrial and fertilizer as relevant.  
110

111 This research aimed to characterize the chemical composition of  $PM_{2.5}$  at a representative location in the CRV, including EC,  
112 primary and secondary OC, ions, trace metals, and specific molecular markers, such as PAHS, n-alkanes, and carbohydrates,  
113 as well as the relationships among these components and with emission sources. Diagnostic ratios were used to identify the  
114 most important  $PM_{2.5}$  components and as a tool for preliminary pollutant source attribution, including primary and secondary  
115 aerosols generated by or associated with sugarcane pre-harvest burning PHB. We believe that in the CRV case, this analysis  
116 is needed prior to source apportionment with receptor models for three reasons: 1) This is the first comprehensive investigation  
117 of PM composition in the CRV (prior studies included two types of components at most); 2) There are no suitable chemical  
118 profiles for some pollutant sources, particularly sugarcane PHB; 3) Our measurements dataset is just barely large enough for  
119 profile-free receptor modeling (positive matrix factorization). We expect that this study also motivates future research on  
120 source apportionment in the region. Our results are particularly relevant for urban communities and atmospheres impacted by  
121 large-scale intensive agriculture and industrial emissions, particularly in developing countries, especially in Latin America  
122 where PM composition information is still scarce (Liang et al., 2016).  
123

## 124 2. Methods

### 125 2.1. Description of the sampling site

126 The sampling site was located on the rooftop of an 8-story administrative building at the Palmira Campus of Universidad  
 127 Nacional de Colombia ( $3^{\circ}30'44.26''$  N;  $76^{\circ}18'27.40''$  W, 1065 m altitude), about 27 m above the ground. The campus is located  
 128 on the western outskirts of Palmira's urban area and is surrounded by short buildings on the east, and extensive sugarcane  
 129 plantations, several sugar mills, and other industries elsewhere. Palmira is located at ~27 km northeast of Cali and ~22 km  
 130 southeast of Yumbo, an important industrial hub. The Pacific Ocean coastline stretches at ~120 km across the Western  
 131 Cordillera, as shown in Figure 1, where operates one of the busiest international trade seaports in Colombia (López, 2017).  
 132 Most of the freight is transported by diesel-powered trucks. Road traffic is also substantial within the CRV, with Bogota and  
 133 along the Pan-American highway that connects Colombia with other South American countries (Orozco et al., 2012).



134

135 Figure 1. Map of the Cauca River Valley (CRV). The inset shows the location of CRV in Colombia and in Northern South  
136 America. The map shows the main cities in CRV, including Palmira (312 thousand inhabitants), our measurement site, Cali,  
137 the largest city in the southwest of Colombia, Yumbo, an industrial hub, and the main highways. Sugar mills, which produce  
138 sugar, bio-ethanol, and electric power are also shown. The dashed-line defined area is CRV's flattest (slope < 5%) bottomland,  
139 where mechanized, intensive sugarcane agriculture takes place. Significant diesel combustion emissions occur along the  
140 Buenaventura highway because it is one of the busiest ports in Colombia.

141

142 The Andes Cordillera splits into three south-to-north diverging mountain ranges (Western, Central, and Eastern Cordilleras)  
143 near the Colombia-Ecuador border (see Figure 1). The Western Cordillera separates the CRV from the Colombian Pacific  
144 Ocean watershed, the rainiest region on Earth (Hernández and Mesa, 2020). The elevated precipitation in this basin (Mesa and  
145 Rojo, 2020) is due to the presence of a Walker cell convergence zone at the surface, persistent under neutral and La Niña  
146 conditions. This synoptic feature is one of the most important determinants of atmospheric circulation in Colombia, with  
147 prevailing east-to-west winds in the lower troposphere along with upper troposphere return winds (Mesa and Rojo, 2020). The  
148 Andean Cordilleras are nevertheless effective barriers to the Walker circulation near the CRV surface (Lopez and Howell,  
149 1967; Mesa S. and Rojo H., 2020). The elevated humidity in the Pacific Ocean watershed and the closeness of the two Andes  
150 branches drive a zonal regional circulation pattern, consisting ~~in~~ of west-to-east anabatic winds over the Pacific slope of the  
151 Western Cordillera during the daytime followed by rapid katabatic winds in the late afternoon (Lopez and Howell, 1967).  
152 These winds rapidly ventilate the CRV during the late afternoon – early evening period on an almost regular basis. CRV is  
153 wide (~22 km) and long (~248 km) enough to develop a valley-mountain wind circulation pattern during the daytime. Winds  
154 are very mild during this time period and expected to be highly dispersive, i.e. with high turbulence intensities (Ortiz et al.,  
155 2019). The arrival of the katabatic “tide” in the late afternoon wipes the valley-mountain wind pattern out (Lopez and Howell,  
156 1967).

## 157 **2.2. Sampling protocols**

158 The sampling campaign was conducted between July 25<sup>th</sup> and September 19<sup>th</sup>, 2018. PM<sub>2.5</sub> aerosol particles (aerodynamic  
159 diameter < 2.5 μm) were collected on Teflon and quartz fiber filters simultaneously for 23 h (from 12:00 local time – LT – to  
160 the next day at 11:00 LT), using 2 in-tandem low-volume samplers (ChemComb speciation samplers, R&P). Each sampler  
161 used an independent pump set at a flow rate of 14 L min<sup>-1</sup>. For both types of filters, three lab blank filters without exposure  
162 were analyzed. Quartz filters were pre-baked at 600 °C for 8 h before sampling to eliminate contaminant trace hydrocarbons.  
163 In total, 45 samples were collected. Prior to and after exposure, the filters were conditioned at constant humidity (36±1.5%  
164 relative humidity) and temperature (24 ± 1.2 °C) for 24 h before being weighing on a microbalance (Sartorius, Mettler Toledo)  
165 with a 199.99 g capacity and 10 μg resolution. PM<sub>2.5</sub>-loaded filters were saved at Petri boxes previously prepared to avoid  
166 cross-contamination of organic species. The filters were subsequently stored at –20°C until analysis to reduce the volatilization

167 of species such as ammonium nitrate and semi-volatile organic compounds. Blank quartz filters were pre-baked and stored  
168 following an identical procedure to exposed filters to collect samples. Blank Teflon filters were treated under the same  
169 conditions of storage, transport, and analysis as PM<sub>2.5</sub>-loaded filters. By differential weighing, mass concentrations were  
170 determined from the Teflon filters.

171

172 Several frequent challenges can affect compound measurements in particle matter, including: 1) The absorption of some gases  
173 in the inlet's galvanic steel, which alters the gas and particle balance of the HNO<sub>3</sub> ⇌ NO<sub>3</sub><sup>-</sup> system of particles collected. During  
174 the collecting of the samples for this study, no denuders were utilized. 2) Significant temperature changes during sampling and  
175 then in the conditioning before to filter weighing can cause ammonium nitrate to volatilize. Because the samples were collected  
176 at temperatures ranging from 17 to 33 C and then conditioned to 25 C, the equilibrium of the HNO<sub>3</sub> ⇌ NO<sub>3</sub><sup>-</sup> system could be  
177 a source of ambiguity in the data reported here. The vaporization of some semi-volatile organic species throughout the sampling  
178 and storage period, as well as the absorption of organic gases over the filter material, are two additional sources of uncertainty.

179

180 It's worth mentioning that during the sampling period, 1888 sugarcane PHB episodes occurred. This register was made by the  
181 regional environmental agency (CVC, as per its acronym in Spanish), using information from sugar mills about PHB events.  
182 The vast majority of these events were intentional, controlled, size-limited (~6 ha median area), and brief (~25-minute median  
183 duration) (Fig S1).

### 184 **2.3. Analytical methods**

185 The quartz-fiber filter samples were analyzed for ions, metals, elemental and organic carbon, and speciation of the  
186 carbonaceous fraction. The Teflon-membrane filter samples were analyzed for metals.

187

188 Two circular pieces with an 8 mm diameter (100.5 mm<sup>2</sup>) were punched from each quartz and Teflon filter, following the  
189 method described by Wadinga Fomba et al., (2020), and extracted using 1 mL of ultrapure water (18 MΩ) in a shaker at 400  
190 rpm for 120 min. The extracts were filtered through 0.45 μm syringe filters (Acrodisc Pall). An aliquot of the solution was  
191 analyzed for inorganic (K<sup>+</sup>, Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, NO<sub>2</sub><sup>-</sup>, PO<sub>4</sub><sup>3-</sup>, Br<sup>-</sup>, F<sup>-</sup>) and some organic ions (C<sub>2</sub>O<sub>4</sub><sup>2-</sup>,  
192 CH<sub>3</sub>O<sub>3</sub>S<sup>-</sup>, and CHO<sub>2</sub><sup>-</sup>) by ion chromatography (IC690 Metrohm; ICS3000, Dionex). Another aliquot was analyzed for  
193 carbohydrates, including levoglucosan, mannosan, and galactosan, as described by Iinuma et al. (2009a). Organic and  
194 elemental carbon were determined from 90.0 mm<sup>2</sup> filter pieces following the EUSAAR 2 protocol (Cavalli et al., 2010), with  
195 a thermal-optical method using a Sunset Laboratory dual carbonaceous analyzer.

196

197 Seventeen metals, including K, Ca, Ti, V, Cr, Mn, Fe, Ni, Cu, As, Se, Sr, Ba, Pb, Sn, Sb, and Cu, were analyzed from Teflon  
198 (22 samples) and quartz (23 samples) filters by total reflection X-Ray Fluorescence Spectroscopy – TXRF (TXRF, PICOFOX

199 S2, Bruker). Si was not determined as this element is part of the quartz filter substrate. Metals were analyzed from three 8-mm  
200 circular pieces punched from Teflon filters, which were digested a nitric and chloride acid solution for 180 min at 180 °C.  
201 After this, 20- $\mu$ l aliquots of the digested solution were placed on the surface of polished TXRF quartz substrates along with  
202 10  $\mu$ l of Ga solution, which served as an internal standard. This solution was left to evaporate at 100°C. The samples were  
203 measured at two angles with a difference of 90° between them to ensure complete excitation of metals. More details on the  
204 analytical technique can be found in Fomba et al. (2013).

205

206 Alkanes and PAHs were determined from two circular filter punches (6 mm diameter, 56.5 mm<sup>2</sup>), using a Curie-point pyrolyzer  
207 (JPS-350, JAI) coupled to a GC-MS system (6890 N GC, 5973inert MSD, Agilent Technologies). The chemical identification  
208 and quantification of the C<sub>20</sub> to C<sub>34</sub> n-alkanes, as well as the following organic species were performed using the following  
209 external standards (Campro, Germany): pristane, phytane, fluorene (FLE), phenanthrene (PHEN), anthracene (ANT),  
210 fluoranthene (FLT), pyrene (PYR), retene (RET), benzo(b)naphtho(1,2-d)thiophene (BNT(2,1)), cyclopenta(c,d)pyrene  
211 (CPY), benz(a)anthracene (BaA), chrysene(+Triphenylene) (CHRY), 2,2-binaphtyl (BNT(2,2)), benzo(b)fluoranthene (BbF),  
212 benzo(k)fluoranthene (BkF), benzo(e)pyrene (BeP), benzo(a)pyrene (BaP), indeno (1,2,3-c,d)pyrene (IcdP),  
213 dibenz(a,h)anthracene (DahA), and benzo(g,h,i)perylene (BghiP), coronene (COR), 9H-Fluorenone (FLO(9H)), 9,10-  
214 Anthracenedione (ANT (9,10)) and 1,2-Benzanthraquinone (BAQ (1,2)). Four deuterated PAHs, (acenaphthene-d10,  
215 phenanthrene-d10, chrysene-d12, and perylene-d12), and two deuterated alkanes (tetracosane-d50 and tetratriacontane-d70)  
216 were used as internal standards, following the analytical method described by (Neusüss et al., 2000). For each analyzed  
217 compound, the sample concentration was calculated by subtracting the average concentration of three blank filters from the  
218 measured concentration.

#### 219 **2.4. Diagnostic ratios and mass closure**

220 The main PM<sub>2.5</sub> components were estimated from the concentrations of EC, OC, water-soluble ions (NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, NH<sub>4</sub><sup>+</sup>, and  
221 Na<sup>+</sup>), and tracer metal concentrations (Ca, Ti, Fe, Ni, Cu, Zn, As, Se, Sb, Ba, and Pb) as follows: organic material (OM), EC,  
222 ammonium sulfate ((NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>), ammonium nitrate (NH<sub>4</sub>NO<sub>3</sub>), crustal material (dust), other trace elements oxides (TEOs),  
223 and particle-bounded water (PBW). PM<sub>2.5</sub> closure is described by Eq 1 (Dabek-Zlotorzynska et al., 2011). We used the  
224 Interagency Monitoring of Protected Visual Environment (IMPROVE) equations (Chow et al., 2015) to quantify the  
225 concentrations of main compounds (Table 1). The aerosol particle bounded water content was estimated from the measured  
226 ionic composition, relative humidity, and temperature, following the aerosol inorganic model (AIM) described by (Clegg et  
227 al., 1998), which is available for running online at <http://www.aim.env.uea.ac.uk/aim/model2/model2a.php>. The  
228 thermodynamic equilibrium of the system H<sup>+</sup> - NH<sub>4</sub><sup>+</sup> - Na<sup>+</sup> - SO<sub>4</sub><sup>2-</sup> - NO<sub>3</sub> - Cl<sup>-</sup> - H<sub>2</sub>O is described by AIM.

229

$$230 \quad PM_{2.5}(\text{mass closure estimated}) = OM_{pri} + OM_{sec} + EC + NH_4SO_4 + NH_4NO_3 + Dust + TEO + SS + PBW \quad \text{Eq (1)}$$

231 Table 1. Equations used to estimate the main components of  $PM_{2.5}$

Component	Equation	Reference
<b>OM<sub>prim</sub></b>	$= f_1 OC_{prim}$	(Chow et al., 2015) (Turpin and Lim, 2010)
<b>OM<sub>sec</sub></b>	$= f_2 OC_{sec}$	(El-Zanan et al., 2005)
<b>SO<sub>4</sub></b>	$= SO_4^{2-}$	(Chow et al., 2015)
<b>NO<sub>3</sub></b>	$= NO_3^-$	(Chow et al., 2015)
<b>Dust</b>	$= 1.63Ca + 1.94Ti + 2.42Fe$ (Assuming CaO, Fe <sub>2</sub> O <sub>3</sub> , FeO (in equal amounts) and TiO <sub>2</sub> )	(Chow et al., 2015)
<b>PBW</b>	$= k (SO_4^{2-} + NH_4^+)$	(Clegg et al., 1998)
<b>TEO</b>	$= 1.47[V] + 1.27[Ni] + 1.25[Cu] + 1.24[Zn] + 1.32[As] +$ $1.2[Se] + 1.07[Ag] + 1.14[Cd] + 1.2[Sb] + 1.12[Ba] +$ $1.23[Ce] + 1.08[Pb]$	(Snider et al., 2016)

232  $f_1 = 1.6$ . This factor was estimated considering the predominant sources.

233  $f_2 = 2.1$ . This factor was estimated by subtracting the non-carbon component of  $PM_{2.5}$  from the measured mass.

234  $k = 0.32$  was calculated using the Aerosol Inorganic Model.

235

236 The EC tracer method was applied to estimate primary ( $OC_{prim}$ ) and secondary ( $OC_{sec}$ ) organic carbon (Lee et al., 2010). This  
 237 method utilizes EC as a tracer for primary OC, which implies that  $OC_{prim}$  from non-combustion sources is deemed negligible.  
 238 Primary and secondary OC can be estimated by defining a suitable primary OC to EC ratio ( $[OC/EC]_{prim}$ ). See Eq (2) and Eq  
 239 (3). We estimated the  $[OC/EC]_{prim}$  ratio as the slope of a Deming linear fit between EC and OC measurements. The term  $b$   
 240 corresponds to the linear fit intercept, which can be interpreted as the emitted  $OC_{prim}$  that is not associated with EC emissions.  
 241 This method is limited by the following assumptions: 1)  $[OC/EC]_{prim}$  is deemed constant, despite the reality that it may change  
 242 throughout the day depending on factors such as wind direction and the location of the dominant emission sources. Our 23-h  
 243 sampling is expected to smooth this variability source out; 2) It neglects  $OC_{prim}$  from non-combustion sources; and 3) It assumes  
 244 that  $OC_{prim}$  is nonvolatile and nonreactive. Departure from these assumptions implies that the estimation of  $OC_{prim}$  and  $OC_{sec}$   
 245 might be biased, likely underestimating  $OC_{sec}$ .

246

$$247 \quad OC_{prim} = [OC/EC]_{min} * EC + b \quad \text{Eq (2)}$$

$$248 \quad OC_{sec} = OC - OC_{prim} \quad \text{Eq (3)}$$

249  $OC_{prim}$  was also estimated by using an organic tracers method from three sources significant in the CRV, namely fossil fuel  
 250 combustion ( $OC_{FF}$ ), biomass burning ( $OC_{BB}$ ), and vegetable detritus ( $OC_{det}$ ).  $OC_{FF}$ ,  $OC_{BB}$  and  $OC_{det}$  were estimated using a  
 251 fitted linear model by robust regression with a M estimator with bisquare function, which were find the coefficients X, Y and  
 252 Z to multiply the tracers concentrations of each source. The tracers used were the sum of the BghiP and IcdP for fossil fuel  
 253 ( $T_{FF}$ ); levoglucosan for biomass burning ( $T_{BB}$ ); and the sum of the highest molecular weight alkanes ( $C_{27} - C_{33}$ ) for vegetable  
 254 detritus ( $T_{det}$ ). The sum of each tracers multiply by X, Y and Z, respectively, Eq (5), corresponding to  $OC_{prim}$  attributed to  
 255 known sources present in CRV. Th substration of  $OC_{prim}$  attributed to OC total is named  $OC_{rest}$ , which corresponding to another  
 256 sources of OC primary and OC secondary.

$$257 \quad OC_{prim} = (T_{FF} * X) + (T_{BB} * Y) + (T_{det} * Z) \quad \text{Eq (4)}$$

$$258 \quad OC_{prim} = OC_{FF} + OC_{BB} + OC_{det} \quad \text{Eq (5)}$$

$$259 \quad OC_{rest} = OC - OC_{prim} \quad \text{Eq (6)}$$

260 According to Table 1, OM was estimated from OC using conversion factors  $f_1$  and  $f_2$  (Chow et al., 2015), which are dependent  
 261 on the OM oxidation level and the secondary organic aerosol formation and aging during transportation. Turpin and Lim  
 262 (2001a) recommended an OM/OC ratio of  $1.6 \pm 0.2$  for urban aerosols, and  $2.1 \pm 0.2$  for non-urban aerosols, values comparable  
 263 with those found by Aiken et al. (2008), of 1.71 (1.41 – 2.15), where lower values (1.6 – 1.8) are attributed to ground  
 264 measurements in the morning, and higher values (1.8 – 1.9) to aircraft sample measurements. BB aerosols can have even higher  
 265 f values (2.2-2.6), due to the presence of organic components with higher molecular weights, e.g., levoglucosan. However,  
 266 Andreae (2019) recommends a factor of 1.6 for fresh BB aerosol, which is consistent with Hodshire et al (2019). We believe  
 267 that traffic and biomass burning are the dominant  $OC_{prim}$  sources at our site. Therefore, we used  $f_1 = 1.6$  to estimate  $OM_{pri}$ . We  
 268 used a factor of 2.1 to estimate  $OM_{sec}$  from the  $OC_{sec}$  fraction. This factor was chosen based on recommended ratios of  $2.1 \pm 0.2$   
 269 for aged aerosols (Schauer, 1998). Some of the global climate models used to estimate direct radiative forcing from organic  
 270 material present in the aerosols employ OM/OC ratios without separating the sources, while others change the ratio depending  
 271 on type of source using values ranging from 1.4 - 1.6 for fossil fuel and biofuel, and 2.6 for biomass burning. Other set of  
 272 models use specific molecules as tracers to follow the OM, such as monoterpenes, isoprene, aromatics and alkanes. Tsigaridis  
 273 et al., (2014) present a list of tracers than haven been used in various models to quantify OM in the aerosols.

274

275 Concentration ratios among distinct species were used to chemically characterize and infer the main sources of fine particle  
 276 matter at Palmira. As a preliminary proxy for  $PM_{2.5}$  acidity, the cation/anion equivalent ratio and the  $[NH_4^+]/[SO_4^{2-}]$  molar  
 277 ratio were used. The first one is based on electroneutrality and assumes that  $H^+$  balances the excess of anions in the solution  
 278 considered, and the second one ratio is an indicator of acidity attributable to those two ions, which are usually the most

279 abundant cation and anion contained in the PM<sub>2.5</sub>. The cation equivalent to anion equivalent ratio was calculated using Eq (7)  
 280 and Eq (8) for each term.

281

282 However, these approaches to inferring the PM<sub>2.5</sub> acidity can result in challenging interpretations, incomplete and incorrect  
 283 results due to an indirect connection to the system's acidity (Pye et al., 2020). Therefore, the E-AIM (Extended Aerosol  
 284 Inorganics Model) was used to determine the equilibrium state of a system containing water and the following ions: SO<sub>4</sub><sup>2-</sup>,  
 285 NH<sub>4</sub><sup>+</sup>, NO<sub>3</sub><sup>-</sup>, Na<sup>+</sup> and Cl<sup>-</sup>, with an atmosphere of known temperature and relative humidity, without information on gas-phase  
 286 concentrations (NH<sub>3</sub>, HNO<sub>3</sub> and SO<sub>2</sub>), which were not available in this study. The H<sup>+</sup> mole fraction concentration from E-AIM  
 287 IV (Friese and Ebel, 2010), was used to calculate pH following Eq (9). E-AIM requires that the input data for ionic composition  
 288 be balanced on an equivalent basis, which means that the sums of the charges on the cations and anions considered in the  
 289 system do balance, accordingly [SO<sub>4</sub><sup>2-</sup>] + [NO<sub>3</sub><sup>-</sup>] + [Cl<sup>-</sup>] = [NH<sub>4</sub><sup>+</sup>] + [Na<sup>+</sup>]. The disadvantage of this approach is that it does not  
 290 allow for the partitioning of trace gases into the vapor phase. The model is available to run on the following website:  
 291 <http://www.aim.env.uea.ac.uk/aim/model4/model4a.php> (last access: 22 January 2022).

292

$$293 \quad AE = \frac{[SO_4^{2-}]}{48} + \frac{[NO_3^-]}{62} + \frac{[C_2O_4^{2-}]}{44} + \frac{[Cl^-]}{35} + \frac{[PO_4^{3-}]}{31.3} + \frac{[NO_2^-]}{46} + \frac{[Br^-]}{79.9} + \frac{[F^-]}{18.9} + \frac{[CH_3O_3S^-]}{95} + \frac{[CHO_2^-]}{45} \quad \text{Eq (7)}$$

$$294 \quad CE = \frac{[Na^+]}{23} + \frac{[K^+]}{39} + \frac{[NH_4^+]}{18} + \frac{[Mg^{2+}]}{12} + \frac{[Ca^{2+}]}{20} \quad \text{Eq (8)}$$

$$295 \quad pH_x = -\log_{10}(H^+) \quad \text{Eq (9)}$$

296

297 Parent PAH ratios are widely used to identify combustion-derived PAHS (Khedidji et al., 2020; Szabó et al., 2015; Tobiszewski  
 298 and Namieśnik, 2012), although some of them are photochemically degraded in the atmosphere (Yunker et al., 2002).  
 299 Additionally, n-alkanes are employed as markers of fossil fuel or vegetation contributions to PM<sub>2.5</sub>. Carbon number maximum  
 300 concentration (C<sub>max</sub>), carbon preference index (CPI), and wax n-alkanes percentage (WNA%) were the criteria utilized to  
 301 determine the n-alkane origin. Table 2 summarizes the diagnostic ratio equations and the expected dominating source based  
 302 on the ratio value.

303

304 Table 2. Diagnostic ratios of organic compounds used to infer the sources of PM<sub>2.5</sub> in this study.

Diagnostic ratios	Equation	Value	Source	References
BeP/(BeP+BaP)		~0.5 < 0.5	Fresh particles Photolysis	(Tobiszewski and Namieśnik, 2012)
IcdP/(IcdP+BghiP)		<0.2 0.2 - 0.5 >0.5	Petrogenic Petroleum combustion Grass, wood and coal combustion	(Yunker et al., 2002) (Tobiszewski and Namieśnik, 2012)
BaP/BghiP		<0.6 >0.6	Non-traffic emissions Traffic emissions	(Tobiszewski and Namieśnik, 2012) (Szabó et al., 2015)
IcdP/BghiP		>1.25 <0.4	Brown coal* Gasoline	(Ravindra et al., 2008)
LMW/(MMW+HMW)		<1 >1	Pyrogenic Petrogenic	(Tobiszewski and Namieśnik, 2012)
C <sub>max</sub>		< C <sub>25</sub> C <sub>27</sub> - C <sub>34</sub>	Anthropogenic Vegetative detritus	(Lin et al., 2010)
CPI	$CPI = 0.5 * \left[ \frac{\sum_{19}^{33} C_i}{\sum_{20}^{32} C_k} + \frac{\sum_{19}^{33} C_i}{\sum_{22}^{34} C_k} \right]$	CPI ~1 CPI > 1	Fossil carbon Biogenic	(Marzi et al., 1993) (Kang et al., 2018)
WNA%	$\sum WNA_{C_n} = [C_n] - \left[ \frac{(C_{n+1}) + (C_{n-1})}{2} \right]$ $WNA\% = \frac{\sum WNA_{C_n}}{\sum Total\ n - alkanes}$ $PNA\% = 100 - WNA\%$	WNA ~ 100 PNA ~ 100	Biogenic Anthropogenic	(Lyu et al., 2019)

305 \*Used for residential heating and industrial operation.

306

307 As all measured variables were subject to analytical uncertainty and temporal variability, linear fitting parameters were  
 308 obtained from Deming regressions as recommended for atmospheric measurements (Wu and Zhen Yu, 2018). The Spearman  
 309 coefficient was selected instead of Pearson's as an indicator of statistical correlation between chemical components to reduce  
 310 the effect of outliers. Derived ratios and other parameters were considered statistically significant when p-values < 0.05. The  
 311 statistical analysis was conducted using R version 4.0.2, 24 including the packages corr (0.4.2), mcr (1.2.1), cluster (2.1.0),  
 312 tidyverse (1.3.0), ggplot (3.3.2), MASS (7.3-53.1) and openair (2.7-4).

### 313 3. Results and discussions

#### 314 3.1. Meteorology

315

316 One year prior to the sampling period, we monitored the local meteorology, first at 14.5 m above the ground, a few meters  
 317 over the mean canopy level, and then at 32.5 m above the ground during the sampling campaign. The box-and-whisker plot in

318 Fig 2 shows katabatic tide winds of up to ~8 m/s at the sampling site elevation, peaking at ~17:00 LT. Wind speeds were a  
319 factor ~2-3 slower at ground level. The wind runs at the sampling height were typically above ~200 km per day (Fig S3)  
320 indicating that the samples had substantially broader spatial coverage of the CRV, much larger than it would have been at  
321 ground level. This also implies that the samples were frequently and significantly influenced by emissions coming from  
322 Yumbo's industrial hub (northwest of Palmira), and also by Palmira and Yumbo urban and highway emissions, as well as  
323 sugarcane PHB and sugarcane mill emissions. The wind rose (Fig 2a) suggests that the influence of urban emissions from Cali,  
324 CRV's largest city by far, was minor. Other meteorological variables are reported in the Supplementary Material (SM) (Fig  
325 S2). Temperature (24.2°C on average) and relative humidity (71.6%) were very likely controlled by solar radiation (350 W m<sup>-2</sup>  
326 on average). The late-afternoon katabatic tide is fast enough to temporarily reduce temperature. The daily pressure profile  
327 (~763 hPa on average) clearly showed the influence of the katabatic tide, with a ~3 hPa drop during its arrival in the late  
328 afternoon. Overall, we believe our measurements at the Palmira site are reasonably representative of the regional air quality.

329



### 336 3.2. Bulk PM<sub>2.5</sub> concentration and composition

337

338 The daily PM<sub>2.5</sub> concentration measured in this study ranged from 6.73 to 24.45  $\mu\text{g m}^{-3}$  with a campaign average of  $14.38 \pm$   
339  $4.35 \mu\text{g m}^{-3}$  (23 h-average,  $\pm 1$ -sigma). Although these concentrations may appear comparatively low, it is worth stressing that  
340 samples were collected at more than 30 m height, with hourly wind speeds frequently above  $4 \text{ m s}^{-1}$ . However, most days  
341 during this study, PM<sub>2.5</sub> concentration exceeded the  $5 \mu\text{g m}^{-3}$  annual mean and  $15 \mu\text{g m}^{-3}$  24-h mean guidelines by World Health  
342 Organization, (2021). Nevertheless, the Colombian standards are less demanding, thus observed concentrations comply with  
343 the  $37 \mu\text{g m}^{-3}$  24-h mean (MADS, 2017).

344

345 Previous studies conducted in rural areas of Brazil impacted by open field sugarcane burning reported significantly higher  
346 (mean  $22.7 \mu\text{g m}^{-3}$ ; Lara et al., 2005), similar (mean  $18 \mu\text{g m}^{-3}$  Souza et al., 2014), and significantly lower PM<sub>2.5</sub> concentrations  
347 (mean  $10.88 \mu\text{g m}^{-3}$ ; Franzin et al., 2020). Comparable measurements in Mexico during harvest periods showed much higher  
348 concentrations, from  $29.14 \mu\text{g m}^{-3}$  (Mugica-Alvarez et al., 2015) up to  $51.3 \mu\text{g m}^{-3}$  (Mugica-Álvarez et al., 2016). Our PM<sub>2.5</sub>  
349 concentration measurements in the CRV are thus substantially lower than those usually reported in Mexico and Brazil during  
350 sugarcane burning periods. Major differences among sugarcane PHB practices in Colombia, Brazil and Mexico must be  
351 considered while comparing concentrations. First,  $\sim 1/3$  of the sugarcane harvested area is burned before harvest at CRV. This  
352 fraction is much larger in Mexico and Brazil (FAO, 2020). Second, sugarcane is harvested year-round in CRV, as opposed to  
353 Brazil and Mexico, where harvest is limited to a  $\sim 6$ -month period (known in Spanish as *zafra*, “the harvest”). Third, the size  
354 of the individual plots burned in CRV is typically  $\sim 6$  ha (median burned area; Cardozo-Valencia et al., 2019), compared to  
355 much larger plots and total areas in Brazil and Mexico (FAO, 2020).

356

357 OC was the most abundant measured PM<sub>2.5</sub> component with a mean daily concentration of  $3.97 \pm 1.31 \mu\text{g m}^{-3}$ , whereas the  
358 mean EC concentration was only  $0.96 \pm 0.31 \mu\text{g m}^{-3}$ . These two components contributed to  $29.1 \pm 8.3\%$  and  $7.2 \pm 2.3\%$  of the  
359 PM<sub>2.5</sub> mass, respectively (carbonaceous fractions were thus  $4.93 \pm 1.58 \mu\text{g m}^{-3}$ , i.e.  $36.31 \pm 10.41\%$  of PM<sub>2.5</sub>).

360

361 The most abundant water-soluble ions found in Palmira’s PM<sub>2.5</sub> were  $\text{SO}_4^{2-}$ ,  $\text{NH}_4^+$ , and  $\text{NO}_3^-$ , with average concentrations of  
362  $2.15 \pm 1.39 \mu\text{g m}^{-3}$ ,  $0.67 \pm 0.62 \mu\text{g m}^{-3}$ , and  $0.51 \pm 0.30 \mu\text{g m}^{-3}$ , respectively ( $12.7 \pm 2.8\%$ ,  $3.7 \pm 1.1\%$  and  $2.6 \pm 1.3\%$  of mass  
363 concentration, respectively). Other water-soluble ions, such as  $\text{Na}^+$ ,  $\text{Ca}^+$ , and  $\text{C}_2\text{O}_4^{2-}$ , had mean concentrations of around  $0.1$   
364  $\mu\text{g m}^{-3}$ , while those of  $\text{K}^+$ ,  $\text{PO}_4^{3-}$ ,  $\text{CH}_3\text{O}_3\text{S}^-$ ,  $\text{Mg}^{2+}$ , and  $\text{Cl}^-$  had concentrations ranging from  $10$ - $80 \text{ ng m}^{-3}$  (Table 3).

365

366 The predominant elements were Ca ( $0.42 \pm 0.33 \mu\text{g m}^{-3}$ ), K ( $0.13 \pm 0.08 \mu\text{g m}^{-3}$ ), and Fe ( $88 \pm 65 \text{ ng m}^{-3}$ ), followed by Zn ( $34$   
367  $\pm 33 \text{ ng m}^{-3}$ ), Pb ( $18 \pm 19 \text{ ng m}^{-3}$ ), Sn ( $52 \pm 37 \text{ ng m}^{-3}$ ), Ti ( $5 \pm 4 \text{ ng m}^{-3}$ ), Ba ( $9 \pm 13 \text{ ng m}^{-3}$ ), Sr ( $2 \pm 5 \text{ ng m}^{-3}$ ). Mn, Ni, Cr, and

368 Se concentrations were below  $2 \pm 1 \text{ ng m}^{-3}$ . Trace metals such as Ti, Cr, Mn, K, Ca, Fe, Ni, Cu, Zn Sr, Pb and Se were found  
 369 in all  $\text{PM}_{2.5}$  samples, while V was found only in a few samples. Other trace metals such as As and Sb were detected only at a  
 370 reduced number of samples with concentrations below  $20 \text{ ng m}^{-3}$ . Table 3 shows the mean, standard deviation, minimum, and  
 371 maximum concentration of the carbonaceous fraction, soluble ions, and metals found in the  $\text{PM}_{2.5}$  samples collected in the  
 372 CRV.

373 Table 3. Mean, 1 standard deviation, minimum and maximum concentrations of carbonaceous fraction, soluble ions, and  
 374 metals in samples of  $\text{PM}_{2.5}$  collected in Palmira.

Species	# of samples	Mean	SD	Min	Max	Units
$\text{PM}_{2.5}$	22	14.38	4.35	6.73	24.45	$\mu\text{g m}^{-3}$
OC	45	3.97	1.31	2.31	8.35	
EC	45	0.96	0.31	0.52	2.15	
$\text{SO}_4^{-2}$	45	2.15	1.39	0.98	10.27	
$\text{NH}_4^+$	45	0.67	0.62	0.18	4.29	
$\text{NO}_3^-$	45	0.51	0.30	0.11	1.45	
$\text{Na}^+$	19	0.21	0.16	0.02	0.45	
$\text{Ca}^{+2}$ (Water soluble ion)	45	0.14	0.06	0.06	0.28	
$\text{C}_2\text{O}_4^{-2}$	45	0.11	0.06	0.04	0.36	
$\text{K}^+$ (Water soluble ion)	45	0.09	0.06	0.02	0.30	
Ca (Trace metal)	42	0.42	0.33	0.01	1.95	
K (Trace metal)	43	0.13	0.08	0.02	0.46	
Formate	13	82	88	0	217	$\text{ng m}^{-3}$
$\text{PO}_4^{-3}$	21	66	42	10	148	
Methansulfonate	45	50	36	13	256	
Cl-	30	20	19	0	75	
$\text{Mg}^{+2}$	45	19	10	2	52	
$\text{NO}_2^-$	45	3	1	1	6	
Fe	42	88	64	2	293	
Sn	23	52	37	9	137	
Zn	42	34	33	0	153	
Pb	42	18	19	0	84	
Ba	20	9	13	2	72	
Sb	19	8	5	3	22	
Cu	42	6	5	1	22	
Ti	42	5	4	0	17	
As	5	2	4	0	10	
Mn	42	2	1	0	5	

Ni	42	2	1	0	9
Sr	42	2	5	0	28
Cr	41	1	1	0	4
Se	41	1	1	0	6
V	20	0	1	0	3

375

376 **3.3. Ions**

377  $\text{SO}_4^{2-}$  and  $\text{NH}_4^+$  were the most abundant anion and cation in the  $\text{PM}_{2.5}$  samples. The molar ratio  $[\text{NH}_4^+]/[\text{SO}_4^{2-}]$  was  $1.6 \pm 0.3$   
378 (min: 0.8 and max: 2.3), suggesting that  $\text{PM}_{2.5}$  is acid. The pH of  $\text{PM}_{2.5}$  samples was determined using the IV E-AIM  
379 thermodynamic model, which estimates the activity coefficient of these species in aqueous phase equilibrium using the  $\text{H}^+$ -  
380  $\text{NH}_4^+$ - $\text{Na}^+$ - $\text{SO}_4^{2-}$ - $\text{NO}_3^-$ - $\text{Cl}^-$ - $\text{H}_2\text{O}$  system. As a result, the pH was  $2.5 \pm 0.4$ . The correlation between the ratio  $[\text{NH}_4^+]/[\text{SO}_4^{2-}]$  and  
381 the pH was strong ( $r^2 = 0.96$ , as plot in Figure S3), suggesting that the molar concentrations of those ions can significantly  
382 explain the particle acidity. Other studies have found similar  $[\text{NH}_4^+]/[\text{SO}_4^{2-}]$  values for pH lower than the estimated in CRV.  
383 Xue et al., (2011), for example, shows molar ratios in ranging from 1.32 to 1.71 and pH values between -0.45 and 0.59. ). Pye  
384 et al., (2020) showed that fine particles have a bimodal distribution of pH, with one mode around a pH of 1–3, and another  
385 mode around a pH of 4–5, the latter influenced by dust, sea spray, and potentially biomass burning). In this study, only one  
386  $\text{PM}_{2.5}$  sample exceed a pH value of 4. Overall, this is an indicator of the abundance of sulfate and organics compounds in  
387 samples collected in the CVR.

388

389 The pH affects the partitioning of total nitrate ( $\text{NO}_3^- + \text{HNO}_3$ ) and total ammonium ( $\text{NH}_4^+ + \text{NH}_3$ ) between the gas and  
390 particulate phases. Lower pH values favor the partitioning of total nitrate toward the gaseous phase ( $\text{HNO}_3$ ) rather than the  
391 particulate phase ( $\text{NO}_3^-$ ). In contrast, the partitioning of total ammonium is favored toward the particulate phase, remaining as  
392  $\text{NH}_4^+$  in the aerosol, whereas  $\text{SO}_4^{2-}$  is a nonvolatile species that remained in the particulate phase. Acidity conditions in the  
393 samples collected in this study are consistent with concentrations of  $\text{SO}_4^{2-}$ ,  $\text{NH}_4^+$ , and  $\text{NO}_3^-$ , corresponding to  $2.5 \mu\text{g m}^{-3}$ ,  $0.7$   
394  $\mu\text{g m}^{-3}$ , and  $0.5 \mu\text{g m}^{-3}$ , respectively. Ammoniated sulfate and ammonium nitrate are generally considered the predominant  
395 forms of nitrate and sulfate in the inorganic fraction in fine particles. In limited environmental ammonium conditions, ammonia  
396 reacts preferentially with  $\text{H}_2\text{SO}_4$  to form ammonium sulfate ( $[\text{NH}_4]_2\text{SO}_4$ ), letovicite ( $[\text{NH}_4]_3\text{H}[\text{SO}_4]_2$ ) or ammonium bisulfate  
397 ( $[\text{NH}_4\text{HSO}_4]$ ) (Lee et al., 2008). Although the correlation coefficient between  $\text{SO}_4^{2-}$  and  $\text{NH}_4^+$  concentrations was high ( $R^2 =$   
398  $0.98$ ), the amount of ammonium contained in the samples was not high enough to neutralize sulfate completely and form  
399  $[\text{NH}_4]_2\text{SO}_4$ . In environmental with limited concentrations of ammonium, is expected the formation of sulfate salts not  
400 completely neutralized, as  $[\text{NH}_4]_3\text{H}[\text{SO}_4]_2$  and  $[\text{NH}_4\text{HSO}_4]$  (Ianniello et al., 2011). Thus, based on the limited ammonium  
401 concentrations found in  $\text{PM}_{2.5}$  of CRV, the stoichiometric molar ratio between  $[\text{NH}_4^+]/[\text{SO}_4^{2-}]$  of 3:2 for letovicite and 1:1 for  
402 ammonium bisulfate, and the results of the E-AIM model, it is possible to indicate that there is a mixture of sulfate salts, such  
403 as, ammonium bisulfate, letovicite, and ammonium sulfate, which is going to form progressively, according to ammonia

404 availability. The E-AIM model presents the saturation ratio of each solid species, which usually forms before ammonium  
405 bisulfate than letovicite and ammonium sulfate. For a molar ratio of 1.5, the aerosol phase consists almost exclusively of  
406 letovicite and to form ammonium sulfate, the ratio should be over 2.0 (Seinfeld and Pandis, 2006). As result of the  
407  $[\text{NH}_4^+]/[\text{SO}_4^{2-}]$  ratios observed in the samples collected in CRV and the pH estimated from the IV E-AIM model, there is no  
408 reason to assume that nitrate is present as ammonium nitrate in the PM<sub>2.5</sub>.

409 .  
410 Instead of this,  $\text{NO}_3^-$  might be bound to cations contained in sea salt and dust particles to form relative nonvolatile salts, as  
411  $\text{KNO}_3$ ,  $\text{NaNO}_3$  and  $\text{Ca}(\text{NO}_3)_2$ .  $\text{NO}_3^-$  showed correlation with  $\text{Na}^+$ ,  $\text{Ca}^{2+}$  and  $\text{K}^+$  ( $r^2 = 0.6, 0.2$  and  $0.2$ , respectively), indicating  
412 possible formation of those salts. The correlation between  $\text{Na}^+$  and  $\text{NO}_3^-$  could be explained by the impact of sea salt aerosol  
413 that comes from air mass origin in the Pacific Ocean. However, the amount of  $\text{Na}^+$  is not enough to neutralize the total of  $\text{NO}_3^-$   
414 , while  $\text{Ca}^{2+}$  showed to be enough amount to neutralize the  $\text{NO}_3^-$ . The molar ratio observed in PM<sub>2.5</sub> samples of CRV for  $[\text{NO}_3^-]$   
415  $[\text{Ca}^{2+}]$  was  $2.6 \pm 1.4$ ,  $[\text{NO}_3^-]/[\text{Na}^+]$  was  $1.7 \pm 1.3$ , and  $[\text{NO}_3^-]/[\text{K}^+]$  was  $5.0 \pm 3.2$ , overcoming the stoichiometric molar ratio  
416 required to form  $\text{Ca}(\text{NO}_3)_2$ ,  $\text{NaNO}_3$ , and  $\text{KNO}_3$ .

417  
418 While, In this study, the abundance of  $\text{SO}_4^{2-}$  in PM<sub>2.5</sub> can be attributed to oxidation of  $\text{SO}_2$  and  $\text{SO}_3$  emitted by from coal fired  
419 in power plants and industrial facilities (Wang et al., 2016), biomass burning activities (Song et al. (2006)) and the emission  
420 of  $\text{H}_2\text{S}$  in poultry production (Casey et al., 2006). The  $\text{H}_2\text{S}$  emission from poultry and pork production is estimated using the  
421 factor emission given by animal units (AU) and the time that it stays in the housing, where one AU corresponding to 500 Kg  
422 of body mass.  $\text{H}_2\text{S}$  emissions from swine and poultry housing trend to be under 5 g  $\text{H}_2\text{S AU}^{-1} \text{ d}^{-1}$  Casey et al., (2006), which  
423 can reach a 3.5 Ton  $\text{H}_2\text{S d}^{-1}$  by poultry and 5 Ton  $\text{H}_2\text{S d}^{-1}$  by pork production. Ammonia emissions factors by poultry and  
424 livestock vary from 0.09 to 12.9  $\text{AU}^{-1} \text{ d}^{-1}$  which represents 9.05 Ton  $\text{NH}_3 \text{ d}^{-1}$  by poultry housing and 12. Ton  $\text{d}^{-1}$  by pork  
425 production.

426  
427 PM<sub>2.5</sub> consistently contained methanesulfonate, with an average concentration of  $50 \text{ ng} \pm 13 \text{ m}^{-3}$ . This ion is produced by the  
428 aqueous oxidation of dimethyl sulfide (DMS), one of the most prevalent biogenic sulfur compounds in the troposphere. DMS  
429 oxidation is a major source of non-sea salt sulfate aerosols in marine aeras (Tang et al., 2019), but also can have origin in  
430 continental origins, such as biomass burning, (Gondwe, 2004; Meinardi et al., 2003; Sorooshian et al., 2015; Stahl et al., 2020),  
431 Methanesulfonate was mainly correlated to the ions sulphate and ammonia ( $r^2 = 0.88$ ) and  $\text{C}_2\text{O}_4^{2-}$  ( $r^2 = 0.66$ ), the metals Se ( $r^2$   
432  $= 0.74$ ) and Fe ( $r^2 = 0.41$ ) and the carbonaceous fraction EC ( $r^2 = 0.56$ ) and OC ( $r^2 = 0.49$ ) in this study. Knowing the origin  
433 of this ion in PM<sub>2.5</sub> in CRV, which is not directly coastal area, prompts future studies with a higher time resolution (6-12  
434 hours) to establish the connection with changes in the wind pattern and the impact of the katabatic circulation, especially  
435 because biomass burning, mainly from sugarcane burnt, is an activity developing during all year in CRV.

436

437 The measured average ratio of  $[\text{SO}_4^{2-}]/[\text{NO}_3^-] = 4.5 \pm 2.9$ . This ratio is higher than the one obtained by Souza et al. (2014) at  
438 Piracicaba ( $3.6 \pm 1.0$ ) and Sao Paulo ( $1.8 \pm 1.0$ ), Brazil. The strong correlations between  $\text{SO}_4^{2-}$  and  $\text{NH}_4^+$  ( $r^2 = 0.84$ ),  $\text{SO}_4^{2-}$  and  
439 methanesulfonate ( $\text{CH}_3\text{O}_3\text{S}^-$ ) ( $r^2 = 0.88$ ), and  $\text{SO}_4^{2-}$  and oxalate dianion ( $\text{C}_2\text{O}_4^{2-}$ ) ( $r^2 = 0.71$ ) allow us to infer that inorganic  
440 secondary aerosol formation is a significant  $\text{PM}_{2.5}$  source in the CRV. In addition, the presence of potassium cation ( $\text{K}^+$ ) in  
441 submicron particles is recognized as a biomass burning tracer (Andreae, 1983; Ryu et al., 2004).  $\text{K}^+$  showed a moderate  
442 correlation with nitrite anion ( $\text{NO}_2^-$ ) ( $r^2 = 0.44$ ) and  $\text{C}_2\text{O}_4^{2-}$  ( $r^2=0.43$ ) in the CRV, which suggests that biomass burning  
443 influences secondary aerosol formation.  $\text{Mg}^{2+}$  and  $\text{Ca}^{2+}$  ions, usually considered crustal metals, exhibited a moderate  
444 correlation of  $r^2 = 0.59$  (Li et al., 2013). Also,  $\text{Mg}^{2+}$  and  $\text{C}_2\text{O}_4^{2-}$  moderate correlation ( $r^2 = 0.26$ ) points to a link between crustal  
445 species and secondary aerosols. Such an association could be plausibly explained by soil erosion induced by pyro-convection  
446 during sugarcane pre-harvest burning (Wagner et al., 2018). Our study full species correlation matrix is shown in Fig 4S.  
447

### 448 3.4. Metals

449  
450 The measured total  $\text{PM}_{2.5}$  trace metal concentration was  $706 \pm 462 \text{ ng m}^{-3}$  ( $101.3 \text{ ng m}^{-3}$  to  $2638 \text{ ng m}^{-3}$ ). Trace metals can  
451 originate from non-exhaust and exhaust emissions. Non-exhaust emissions come from brake and tire wear, road surface  
452 abrasion, wear/corrosion of other vehicle components, and the resuspension of road surface dust (Pant and Harrison, 2013).  
453 Metals in exhaust emissions are related to fuel, lubricant combustion, catalytic converters, and engine corrosion. As shown by  
454 Kundu and Stone (2014), many of these sources share some metals in their chemical composition profile, thus an unambiguous  
455 specific source attribution is non-trivial. In this study, we found a significant correlation among Fe, Mn and Ti ( $r^2 \approx 0.72$ ),  
456 which is typically associated with a high abundance of crustal material (Fomba et al., 2018), indicating that soil dust is a  
457 significant source in the CRV. Also, tire and brake wear tracer metals, including Zn and Cu, showed weaker but still significant  
458 correlations among them ( $r^2 \approx 0.32$ ). Ca concentrations were quite high ( $405 \pm 334 \text{ ng m}^{-3}$  ( $1.6 \text{ ng m}^{-3}$  to  $1952 \text{ ng m}^{-3}$ ). These  
459 levels can be attributed to dust generation by agricultural practices, particularly land planning, liming and tilling, PHB pyro-  
460 convection-induced soil erosion, and traffic-induced soil resuspension on unpaved rural roads. One of the very few previous  
461 investigations into on PM composition in the CRV (Criollo and Daza, 2011) analyzed trace metals in  $\text{PM}_{10}$  at 4 CRV locations,  
462 including Palmira. They found significant enrichment of Fe and K metals at locations exposed to PHB. It must be kept in mind  
463 that  $\text{PM}_{10}$  samples included coarse mode aerosols, of which dust might have been a significant fraction. Also, environmental  
464 regulations have been successful in steadily reducing the sugarcane burned area in the CRV since 2009. The Burned area  
465 dropped from 72% in 2011 to 35.46% in 2018, our year of measurements (Cardozo-Valencia et al., 2019).  
466

467 Cd, Pb, Ni, Hg and As, and other metals and metalloids are considered carcinogenic (WHO Regional Office for Europe, 2020).  
468 Measured concentrations of Pb and Ni in  $\text{PM}_{2.5}$  at Palmira were  $18 \text{ ng m}^{-3}$  (+/-19) and  $2 \text{ ng m}^{-3}$  (+/-1), respectively. These

469 mean values were below the EU target values of ( $0.5 \mu\text{g m}^{-3}$  and  $20 \text{ ng m}^{-3}$  respectively) (WHO, 2013a), and below the annual  
470 average limit of the Colombian national ambient air quality standard ( $0.5 \mu\text{g m}^{-3}$  and  $0.18 \mu\text{g m}^{-3}$  respectively) (MADS, 2017).  
471 Nevertheless, these concentrations are significantly higher than those reported for other suburban areas in Midwestern United  
472 States and remote sites in the northern tropical Atlantic (Fomba et al., 2018; Kundu and Stone, 2014). Pb concentrations are  
473 similar to those reported for Bogota and other large urban areas (SDA, 2010; Vasconcellos et al., 2007). Pb has been long  
474 banned as a fuel additive in Colombia, thus the observed levels might be associated with metallurgical industry and waste  
475 incineration. Information on ambient air hazardous metal concentrations in Latin America's urban and rural areas is still scarce.  
476

### 477 3.5. Carbohydrates

478

479 Levoglucosan is a highly specific biomass burning organic tracer (Bhattarai et al., 2019). Along with  $\text{K}^+$ , OC and EC, it can  
480 be used to effectively identify the relevance of biomass burning as an aerosol source. The relative contribution of levoglucosan  
481 to the PM carbohydrate burden, and especially the levoglucosan to mannosan ratio, can be used as indicators of the type of  
482 biomass burned (Engling et al., 2009). In this study, the following carbohydrates were quantified: levoglucosan, mannosan,  
483 glucose, galactosan, fructose and arabitol. Levoglucosan was by far the most abundant ( $113.8 \pm 147.2 \text{ ng m}^{-3}$ ), reaching values  
484 of up to  $904.3 \text{ ng m}^{-3}$ , followed by glucose ( $10.4 \pm 6.1 \text{ ng m}^{-3}$ ), mannosan ( $7 \pm 6.1 \text{ ng m}^{-3}$ ), and arabitol ( $4.1 \pm 3.5 \text{ ng m}^{-3}$ ).  
485 Levoglucosan and mannosan were detected in all  $\text{PM}_{2.5}$  samples, while galactosan and fructose were detected only in 9 and 11  
486 samples, respectively. Levoglucosan was  $3.5 \pm 2.3\%$  of OC and  $0.96\% \pm 0.81\%$  of  $\text{PM}_{2.5}$ .

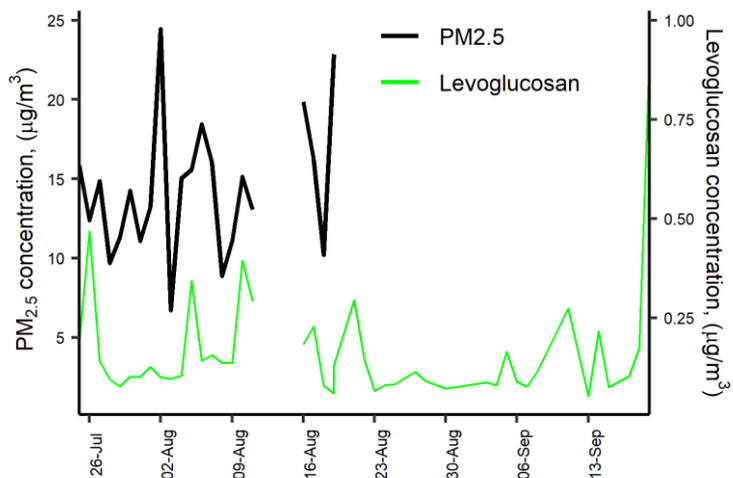
487

488 The levoglucosan concentration found in this study was quite similar to that reported in areas of Brazil where sugarcane  
489 production and processing are important economic activities, Figure 3. For instance, during the harvest (*zafra*) period in  
490 Araraquara, the levoglucosan mean concentration was  $138 \pm 91 \text{ ng m}^{-3}$ , although during the non-harvest period it was  
491 unexpectedly high ( $73 \pm 37 \text{ ng m}^{-3}$ ) (Urban et al., 2014). Likewise, the levoglucosan average concentration at Piracicaba during  
492 a reduced fire period was  $66 \text{ ng m}^{-3}$  (Souza et al., 2014). The measured mean levoglucosan/mannosan ratio in Palmira was  $17.6$   
493  $\pm 13.0$  (min: 8.1 – max: 58.1). Chemical profile studies found a levoglucosan/mannosan ratio of  $\sim 10$  for sugarcane leaves  
494 burned in stoves (Hall et al., 2012; Dos Santos et al., 2002) and of  $\sim 54$  for burned bagasse (Dos Santos et al., 2002). Leaves  
495 constitute the largest fraction (20.8%, Victoria et al., 2002) of pre-harvest burned sugarcane. Consistently and expectably, the  
496 levoglucosan/mannosan ratio at Palmira is much closer to the chemical profile ratio of leaves than that of bagasse. Moreover,  
497 ambient air samples in Araraquara and Piracicaba showed levoglucosan/mannosan ratios of  $9 \pm 5$  and  $\sim 33$ , respectively. For  
498 comparison, the levoglucosan/mannosan ratio in PM from rice straw and other crops burned were  $\sim 26.6$  and  $\sim 23.8$ , respectively  
499 (Engling et al., 2009). This indicates that the levoglucosan/mannosan ratio is sensitive to the type of biomass burned but also  
500 to burning conditions. The large levoglucosan/mannosan ratio in our study suggests that Palmira was impacted by sugarcane  
501 PHB most of the time, and, to a lesser extent, by bagasse combustion in sugar mills. We hypothesize that, even if these were

502 very small, levoglucosan and mannosan combustion emissions might not be negligible as the CRV sugarcane biomass yields  
 503 are very high and most of the harvested sugarcane bagasse is combusted for electric power and steam production.

504

505



506

507

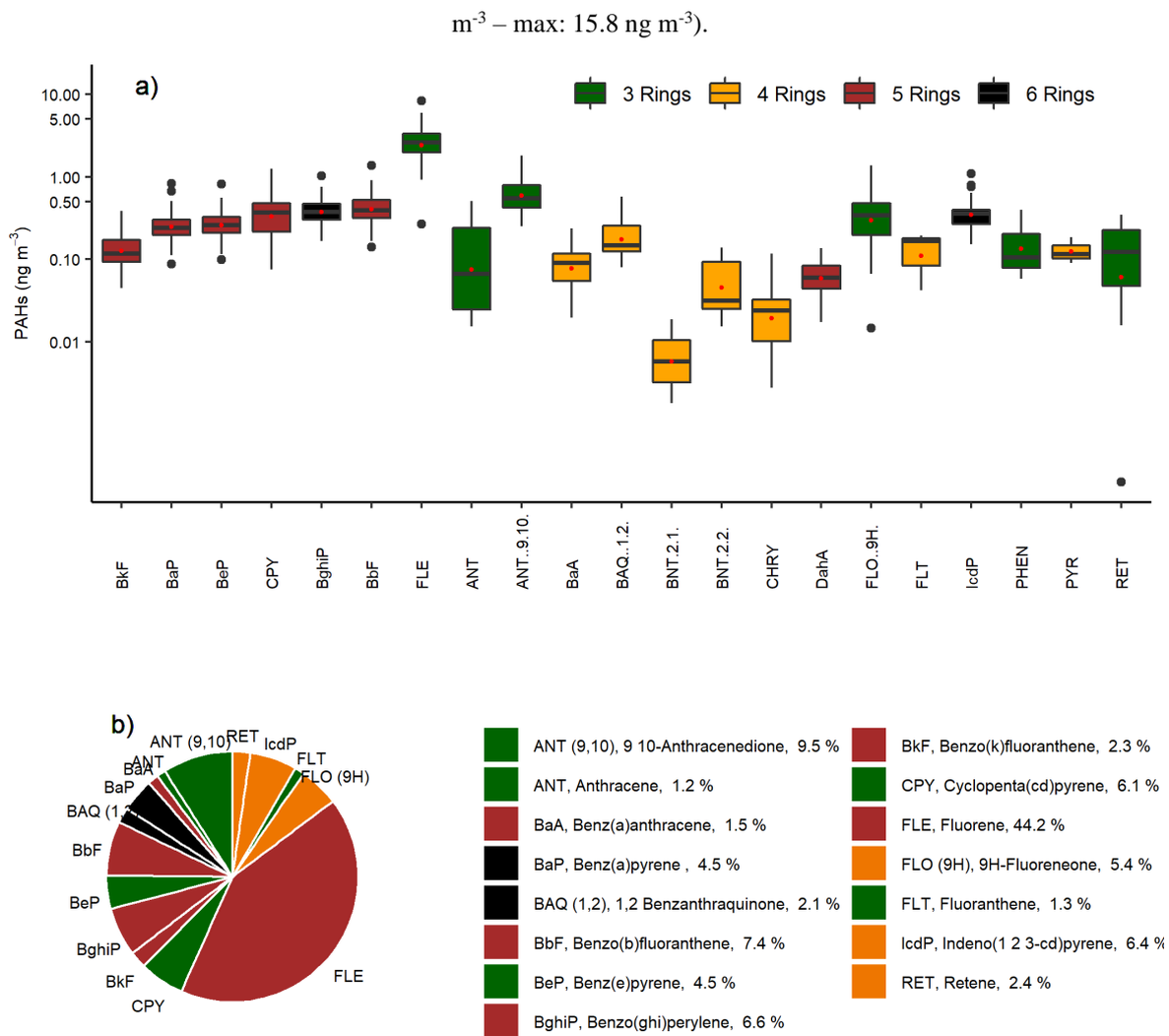
Figure 3. Daily variation of Levoglucosan and PM<sub>2.5</sub> concentration at CRV.

### 508 3.6. Polycyclic Aromatic Hydrocarbons (PAHs)

509

510 A total of 22 PAHs were measured in each sample collected at Palmira, including the 16 PAHs listed as human health  
 511 priority pollutants by WHO and US-EPA (Yan et al., 2004). The total PAHs concentration was  $5.6 \pm 2.9 \text{ ng m}^{-3}$  (min: 2.3 ng

512

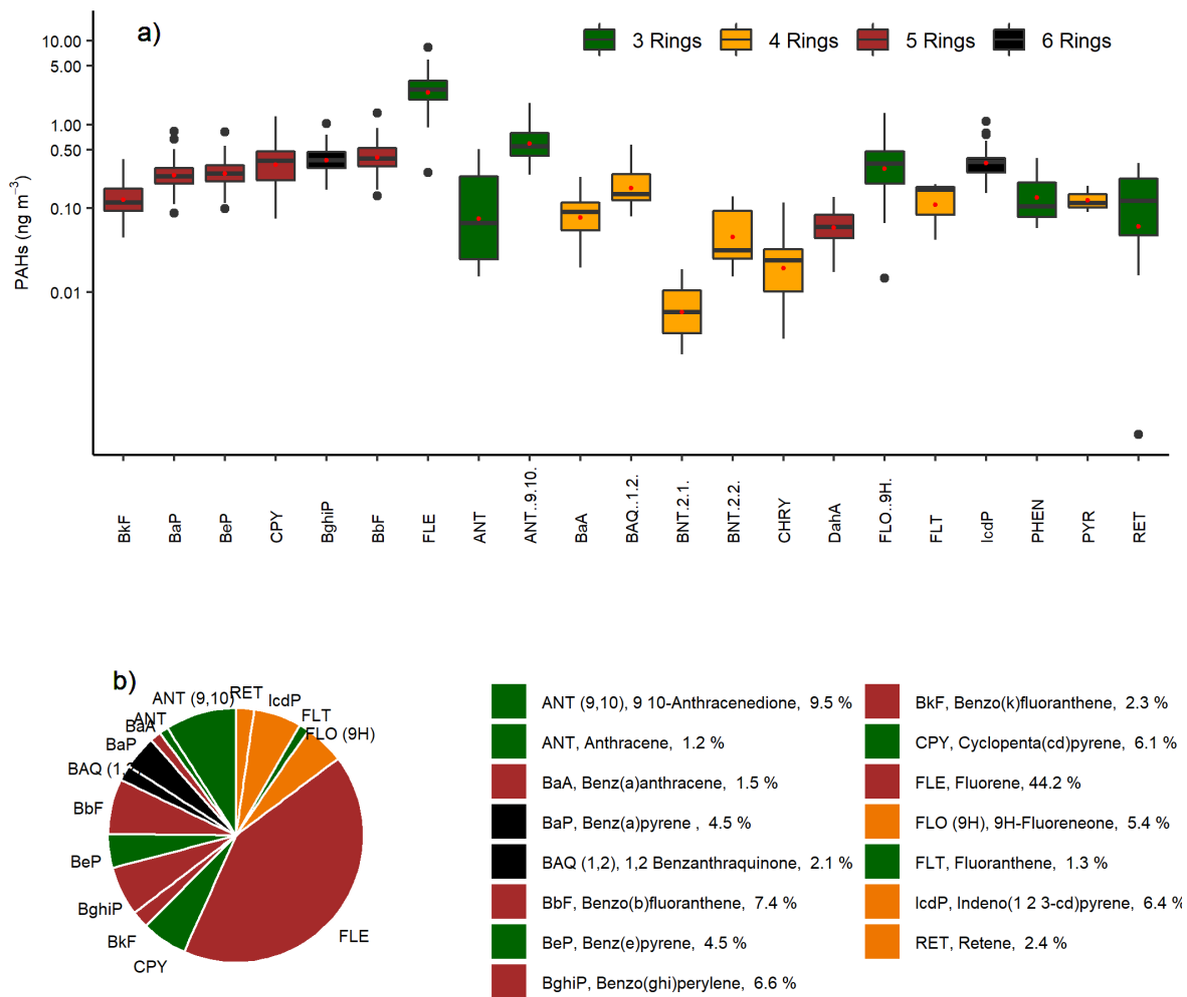


513

514 Figure 4a shows the PAHs concentration variability during the sampling campaign (mean and standard deviation are  
 515 available in Table S2). The most abundant PAH were FLE ( $44.2\% \pm 11.9\%$  total concentration share), ANT (9,10)  
 516 ( $10.0\% \pm 4.5\%$ ), BbF ( $7.4\% \pm 2.3\%$ ), BghiP ( $6.7\% \pm 2.4\%$ ), IcdP ( $6.4\% \pm 1.9\%$ ), CPY ( $6.0\% \pm 2.3\%$ ), FLO (9H) ( $5.4\% \pm 3.1\%$ ),

517

BeP(4.6%±1.3%), and BaP(4.4%±1.6%), which accounted for 95.1% of the total PAH concentration (



518

519 Figure 4b). Three-ring PAHs were the most abundant (59.04% of total PAH). Put together, five- and six-ring PAHs accounted  
 520 for an additional 38.44%. The less abundant PAH group was the four-ring (2.52%). A previous study in CRV, carried out on  
 521 PM<sub>10</sub> samples by Romero et al. (2013), showed higher FLT, PYR, and PHE concentrations in areas highly exposed to sugarcane  
 522 PHB compared to other locations. In contrast, PM<sub>2.5</sub> FLE concentrations in this research were significantly higher than those  
 523 in PM<sub>10</sub> by Romero et al. (2013), while PYR and PHE levels were similar.

524

525 The carcinogenic species BaP, BbF, BkF, BaA, BghiP, FLE, CPY and BeP were identified in all the PM<sub>2.5</sub> samples. BaP is a  
 526 reference for PAH carcinogenicity (WHO, 2013a) that is used as a PAH exposure metric, known as the Benzo(a)Pyrene-  
 527 equivalent carcinogenic potency (BaPE). We calculated BaPE using the toxic equivalent factors (TEF) proposed by Nisbet

528 and LaGoy (1992) and (Malcolm and Dobson, 1994). PAH concentrations were multiplied by TEF and then added to estimate  
529 the carcinogenic potential of PM<sub>2.5</sub>-bound PAHs. The mean carcinogenicity level at Palmira, expressed as BaP-TEQ, was  
530  $0.4 \pm 0.2 \text{ ng m}^{-3}$  (min:  $0.1 \text{ ng m}^{-3}$  - max:  $1.4 \text{ ng m}^{-3}$ ). Only one sample exceeded the Colombian annual limit of  $1 \text{ ng m}^{-3}$  but  
531 most of them exceeded the WHO reference level of  $0.12 \text{ ng m}^{-3}$ . The mutagenic potential of PAHs (BaP-MEQ) was estimated  
532 using the mutagenic equivalent factors (MEF) reported by Durant et al., (1996). The average BaP-MEQ was  $0.5 \pm 0.3 \text{ ng m}^{-3}$   
533 (min:  $0.2 \text{ ng m}^{-3}$  - max:  $1.8 \text{ ng m}^{-3}$ ). These levels are comparable to those measured in PM<sub>2.5</sub> by Mugica-Álvarez et al., (2016)  
534 in Veracruz (Mexico) but during the sugarcane non-harvest period. PM<sub>10</sub> BaP-MEQ levels in Araraquara (Brazil) (de Andrade  
535 et al., 2010; De Assuncao et al., 2014) were twice as high as those found in this study. This suggests that year-long sugarcane  
536 PHB in the CRV leads to lower mutagenic potentials compared to those at locations where the harvesting period (*zafra*) is  
537 shorter, thus with higher burning rates. We estimated the average BaP-TEQ and BaP-MEQ concentrations in the CRV  
538 according to their exposure to sugarcane burning products from Romero et al., (2013) data and used them as a benchmark to  
539 our measurements. PM<sub>10</sub>-bound BaP-TEQ and BaP-MEQ levels for areas not directly exposed to sugarcane burning were  $0.16$   
540  $\text{ng m}^{-3}$  and  $=0.21 \text{ ng m}^{-3}$ , respectively. Toxicity and mutagenicity due to PM<sub>10</sub>-bound PAHs were 4 times as high as those at  
541 areas directly exposed to sugarcane burning. It is reasonable to assume that PAHs are largely bound to fine aerosol ( $<2.5 \mu\text{m}$ ),  
542 thus that our measurements are comparable to (Romero et al., 2013). If so, our site at Palmira would be at an intermediate  
543 exposure condition, higher than areas not directly exposed to sugarcane burning but lower than directly exposed areas.

544

545 Ratios among different PAHs have been extensively used to distinguish between traffic and other PAH sources. We used the  
546 diagnostic ratios presented by Ravindra et al. (2008) and Tobiszewski and Namieśnik (2012a) to better understand the  
547 contribution of sources to PM<sub>2.5</sub> in the CRV. The ratio benzo(e)pyrene to the sum of benzo(e)pyrene and benzo(a) pyrene is  
548 used as an indicator of aerosol aging. Local or “fresh” aerosols have  $[\text{BeP}]/([\text{BeP}]+[\text{BaP}])$  ratios around 0.5, while aged  
549 aerosols can have ratios as low as zero as a result of photochemical decomposition and oxidation. The  $[\text{BeP}]/([\text{BeP}]+[\text{BaP}])$   
550 ratio at Palmira was  $0.51 \pm 0.04$ , with a majority (84.4%,  $n = 38$ ) of fresh samples a minor fraction (15.6%,  $n=7$ ) of  
551 photochemically-degraded samples.

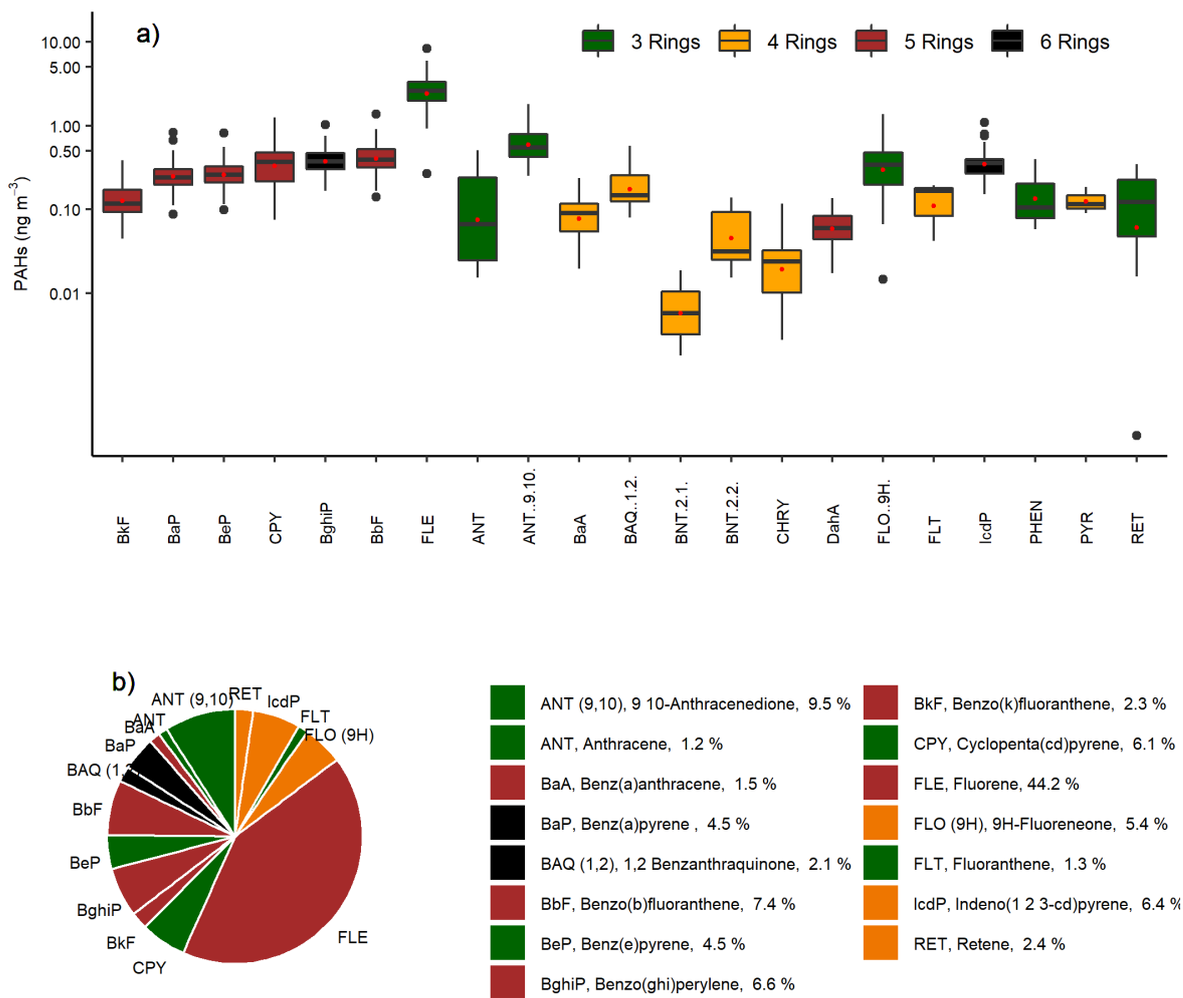
552

553 Other two diagnostic ratios were used to assess the prevalence of traffic as a PM<sub>2.5</sub> source. The first ratio used IcdP BghiP, two  
554 automobile emissions markers (Miguel and Pereira, 1989). Values higher than 0.5 for the ratio  $[\text{IcdP}]/([\text{IcdP}]+[\text{BghiP}])$   
555 indicates aged particles (Tobiszewski and Namieśnik, 2012) generated by coal, grass or wood burning (Yunker et al., 2002).  
556 The second ratio is  $[\text{BaP}]/[\text{BghiP}]$ . Ratios higher than 0.6 are indicative of traffic emissions (Tobiszewski and Namieśnik,  
557 2012). At Palmira, the  $[\text{IcdP}]/([\text{IcdP}]+[\text{BghiP}])$  and  $[\text{BaP}]/[\text{BghiP}]$  ratios were  $0.48 \pm 0.04$  and  $0.69 \pm 0.13$ , which indicates  
558 that ~63% of the samples originated from combustion of oil products ( $n = 30$ ), and ~36% came from non-traffic sources, like  
559 wood, grass, or coal ( $n = 15$ ).

560

561 Also, the structure and size of PAHs are indicative of their sources. PAHs of low molecular weight (LMW) (two or three  
562 aromatic rings) have been reported as tracers of wood, grass, and fuel oil combustion, while those of medium molecular weight  
563 (MMW) (four rings) and high molecular height (HMW) (five and six rings) are associated with coal combustion and vehicular  
564 emissions. The ratio between LMW and the sum of MMW and HMW,  $LMW/(MMW+HMW)$ , is used for source identification.  
565 Ratios lower than one are indicative of oil products combustion, while ratios larger than one are associated with coal and  
566 biomass combustion (Tobiszewski and Namieśnik, 2012). The ratio at Palmira,  $LMW/(MMW+HMW) = 1.43 \pm 1.00$ , was  
567 rather variable but suggests that a large fraction of PAHs in CRV (82.2% of samples) were generated by biomass burning or  
568 combustion, as well as coal combustion in brick kilns. Just one in five samples (17.8%) had PAHs attributable to oil product  
569 combustion.  
570

571 Sugarcane-burning emitted PAH are mainly LMW, especially of two (~66% of PAHs) and three rings (~27%), among which  
 572 FLE, PHE and ANT are the most emitted, according to Hall et al. (2012) chemical profile. The relative abundance of 3-ring



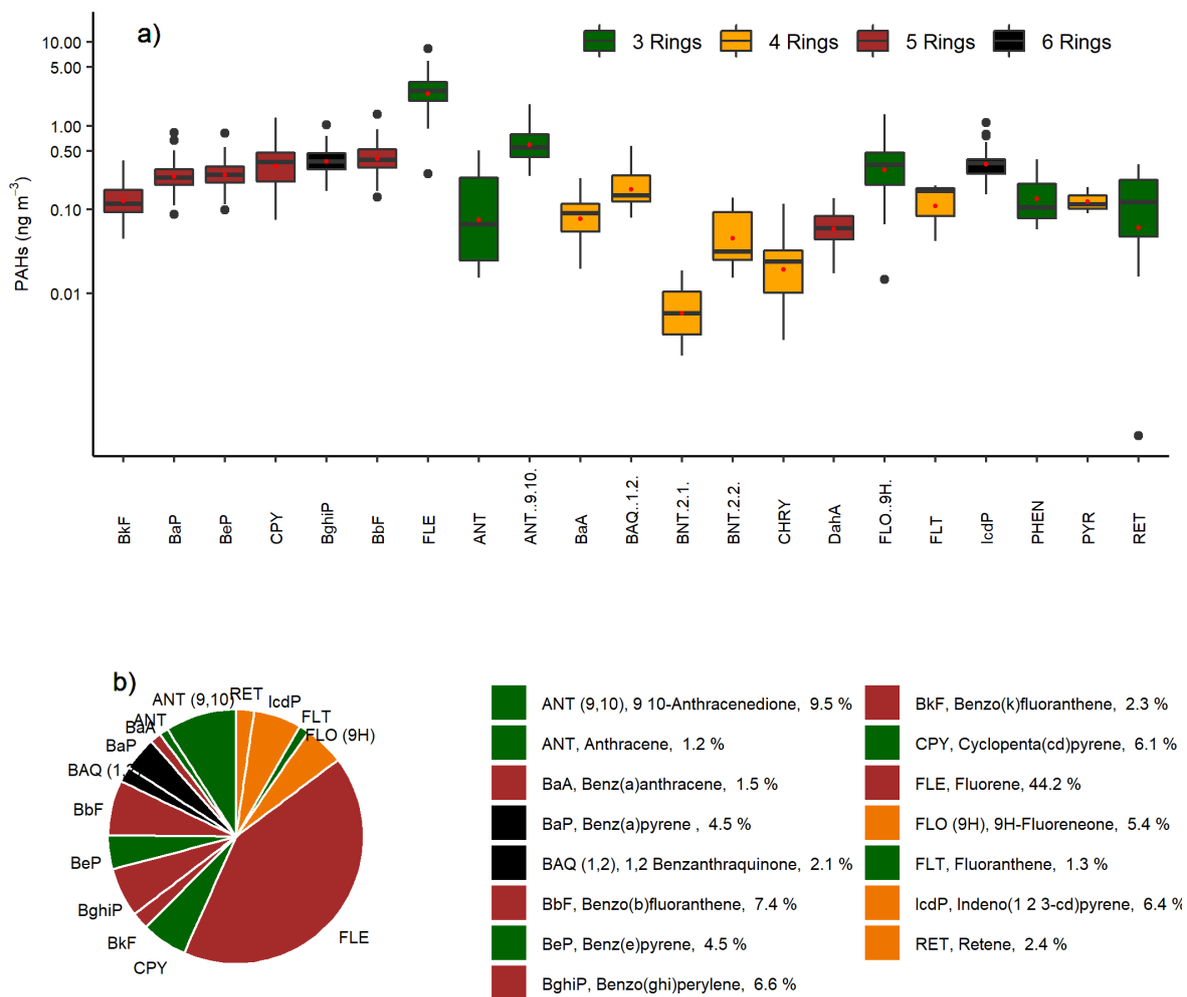
573 PAHs (

574 Figure 4) in CRV's PM<sub>2.5</sub> is likely due to open-field sugarcane PHB to a major extent, and to controlled bagasse combustion  
 575 for electric power and steam production; to a lesser extent.

576

577 The highest PAH concentrations were observed on 10<sup>th</sup> August and 11<sup>th</sup> September 2018, with levels of 15.8 ng m<sup>-3</sup> and 14.4  
 578 ng m<sup>-3</sup>, respectively (Fig 5S). Elevated concentrations of 5 and 6 ring PAHs were observed on 10<sup>th</sup> August 2018. A change in  
 579 the wind circulation pattern was observed on the previous day (Fig S2), with a wind speed reduction and a predominance of  
 580 winds from the north. Later, on 11<sup>th</sup> September 2018, we observed an increase in 3-ring PAHs and winds from the NW at the  
 581 average wind speed at the sampling location. This indicates that there were at least two types of sources. The abundance of

582 HMW PAHs indicates fossil fuel combustion sources, and LMW PAHs suggest that parts of these come from non-fossil fuel  
 583 combustion sources.  
 584



585

586 Figure 4. The abundance of PAHs measured in PM<sub>2.5</sub> samples collected in CRV, represented by colors according to the number  
 587 of rings of each PAHs, green (tree rings), yellow (four rings), brown (five rings), and black (six rings). a) Boxplot of  
 588 concentrations in ng m<sup>-3</sup>, red dots represent mean concentrations of each PAHs. b) pie-plot of the relative abundance of PAHs  
 589 in PM<sub>2.5</sub> samples.

### 590 3.7. Alkanes

591

592 A total of 16 alkanes ranging from C<sub>20</sub> up to C<sub>34</sub> were analyzed in this study and used to identify the presence of fossil fuel  
593 combustion and plant fragments in the PM<sub>2.5</sub> samples. The abundance of total n-alkanes during the whole sampling period was  
594 in the range of 13.0 to 88.45 ng m<sup>-3</sup> with an average concentration of 40.36 ng m<sup>-3</sup> ± 18.82 ng m<sup>-3</sup>. In general, the high molecular  
595 weight n-alkanes such as C<sub>29</sub> – C<sub>31</sub> were the most abundant. These are characteristic of vegetative detritus corresponding to  
596 plant fragments in airborne PM (Lin et al., 2010). The most abundant n-alkanes were C<sub>29</sub>, C<sub>30</sub>, and C<sub>31</sub> (Fig 6.). Likewise, the  
597 carbon number maximum concentration (C<sub>max</sub>) was C<sub>29</sub> in 43% of samples and C<sub>31</sub> in 28% of them. This result is consistent  
598 with the chemical profile of sugarcane burning reported by (Oros et al., 2006) with a C<sub>max</sub> of C<sub>31</sub>.

599

600 The carbon preference index (CPI) and wax n-alkanes percentage (WNA%) are parameters used to elucidate the origin of the  
601 n-alkanes and infer whether emissions come from biogenic or anthropogenic sources. The CPI represents the ratio between  
602 odd and even carbon number n-alkanes. The equation used to calculate CPI in the present study is shown in Table 2, following  
603 the procedure reported by (Marzi et al., 1993). Values of CPI ≤ 1 (or close to 1) indicate that n-alkanes are emitted from  
604 anthropogenic sources, while values higher than 1 indicate the influence of vegetative detritus and biomass burning in the  
605 PM<sub>2.5</sub> samples (Mancilla et al., 2016). In this study, the mean CPI was always greater than 1, with an average value of 1.22 ±  
606 0.18 (min:1.02 – max:1.8) that is between the CPI for fossil fuel emissions of ~1.0 (Caumo et al., 2020) and sugarcane burning  
607 of 2.1 (Oros et al., 2006), revealing the influence of several sources over the PM<sub>2.5</sub> in the CRV.

608

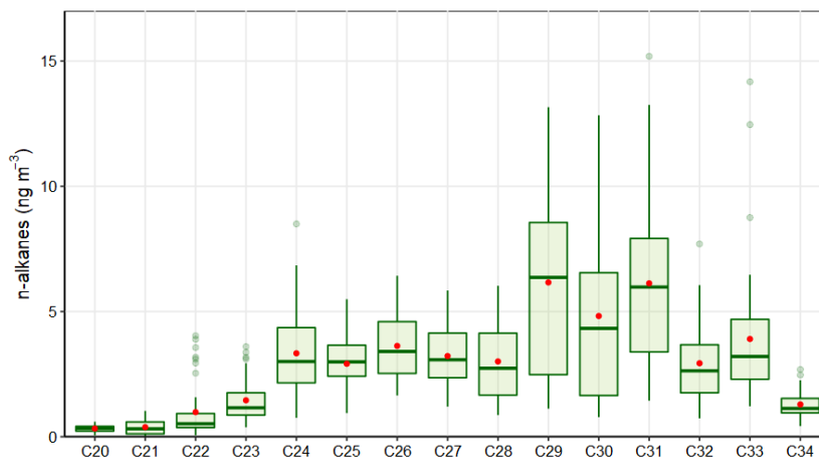
609 Likewise, WNA% represents the preference of odd n-alkanes in the sample. The odd n-alkanes, especially of higher molecular  
610 weight, are representative of plant wax related emissions. The waxes are present on the surface of plants, especially on the  
611 leaves, and they become airborne by a direct or indirect mechanism like wind action or biomass burning (Kang et al., 2018;  
612 Simoneit, 2002). In this research, the samples analyzed showed a preference for odd carbon on C<sub>27</sub>, C<sub>29</sub>, C<sub>31</sub> and C<sub>33</sub>, which  
613 have higher concentrations than the next higher and lower even carbon number homologs, proving the biogenic contribution  
614 over the PM<sub>2.5</sub> in the CRV. The WNA% was calculated using the equation shown in Table 2, described by Yadav et al. (2013).  
615 A larger WNA% represents the contribution from emissions of plant waxes or biomass burning. Otherwise, a smaller value  
616 represents n-alkanes from petrogenic sources, known as petrogenic n-alkanes (PNA)%. The mean WNA% calculated for the  
617 PM<sub>2.5</sub> samples collected from the CRV was 12.65% ± 5.21% (min: 4.71% – max: 29.92%) and can be defined as petrogenic  
618 inputs (PNA%) that were 87.35% during the sampling period. The correlation between CPI and WNA was moderate (r<sup>2</sup>=0.53)  
619 supporting a consistent meaning between these two parameters, and they are useful for assessing the plant wax contribution to  
620 PM<sub>2.5</sub>.

621

622 Overall, the total concentration of n-alkanes in the PM<sub>2.5</sub> in the CRV was lower than those reported in areas where sugarcane  
623 is often burned in Brazil (Urban et al., 2016), although the behavior of the parameters of CPI and C<sub>max</sub> is similar. Compared  
624 with other urban areas in Latin America, the n-alkane concentration in the CRV was similar to that reported in the metropolitan

625 zone of the Mexican valley (MZMV) for  $PM_{2.5}$  (Amador-Muñoz et al., 2011), Bogota for  $PM_{10}$  and slightly lower than reported  
 626 in Sao Paulo for  $PM_{10}$  (Vasconcellos et al., 2011). However, the CPI and WNA in these cities were smaller than in the CRV,  
 627 because of the strong influence of vehicular emissions in these densely populated cities. The OC/EC ratio was moderately  
 628 associated with WNA values ( $r^2 = 0.41$ ), indicating that an increase in this ratio can be explained by the vegetative detritus  
 629 contribution to  $PM_{2.5}$ , while the levoglucosan concentrations did not show correspondence to the CPI and WNA values;  
 630 therefore, the levoglucosan levels did not explain the preference of odd carbon number homologs. These results indicated that  
 631 the n-alkanes found in this study came from several sources, with a noticeable contribution from plant wax emissions. The  
 632 parameters used to assess the source contribution of  $PM_{2.5}$  through n-alkanes such as CPI and WNA%, were characteristic of  
 633 aerosols collected in urban areas.

634



635

636

Figure 5. Average n-alkanes concentrations in  $PM_{2.5}$  samples

### 637 3.8. $PM_{2.5}$ mass closure

638

639 Mass closure (Figure 6) shows the crucial contribution of organic material ( $52.66 \pm 18.44\%$ ) and inorganic fraction, represented  
 640 by sulfate ( $12.69 \pm 2.84\%$ ), ammonium ( $3.75 \pm 1.05\%$ ), nitrate ( $2.56 \pm 1.29\%$ ). EC constituted  $7.13 \pm 2.44\%$  of  $PM_{2.5}$ . The  
 641 mineral fraction corresponded to dust ( $3.51 \pm 1.35\%$ ) and TEO ( $0.85 \pm 0.42\%$ ). Mass closure of  $88.42 \pm 24.17\%$  was achieved.  
 642 Although  $PM_{2.5}$  concentrations observed in the CRV were not so high as compared with those registered in Brazil and Mexico  
 643 during the preharvest season, the EC percentage is in a similar range or slightly lower than those observed in other urban areas  
 644 (Snider et al., 2016), showing the key role of incomplete combustion processes in the area.

645

646 The average (OC/EC) ratio found in CRV was  $4.2 \pm 0.72$ , from which we can infer that secondary aerosol formation had a  
 647 relevant role. The segregation of OC into the primary and secondary fractions was carried out using two methods. The first

648 was the EC tracer method applied in previous studies (Pio et al., 2011; Plaza et al., 2011), and the second was the organic  
649 tracer method, which is based on the lineal regression between OC and organic tracers from primary sources. In the EC tracer  
650 method, the  $(OC/EC)_{\min}$  ratio selected to differentiate  $OC_{\text{prim}}$  from  $OC_{\text{sec}}$  was the minimum ratio observed, equivalent to 2.12.  
651 Still, this value could induce the overestimation of  $OC_{\text{prim}}$  due to the distance between the emission sources and the sampling  
652 site (27 m aboveground), and the local meteorological conditions that favor the volatilization and oxidation of organic  
653 components into particles before being collected. As a result,  $OC_{\text{prim}}$  was estimated at 50.3% and  $OC_{\text{sec}}$  at 49.7% of the total  
654 OC, with a minimum variability of 3.8%. The estimated  $OM_{\text{pri}}$  concentration was  $3.22 \pm 1.09 \mu\text{g m}^{-3}$  and the  $OM_{\text{sec}}$   
655 concentration was  $4.01 \pm 1.78 \mu\text{g m}^{-3}$ , which represented 24.2% and 28.5% of  $PM_{2.5}$  respectively.

656

657 In the organic tracer method, the contribution of fossil fuel combustion - mainly derived from transport -, biomass burning,  
658 and vegetative detritus to  $OC_{\text{prim}}$  was estimated from a linear model by robust regression using an M estimator with bisquare  
659 function between organic tracers and OC. Resulting contributions were as follows:  $OC_{\text{ff}}$ : 16.38%,  $OC_{\text{bb}}$ : 15.19%, and  $OC_{\text{det}}$ :  
660 1.45% of total OC measured. Overall, the use organic tracer method to estimate  $OC_{\text{prim}}$  indicates that this carbonaceous fraction  
661 represents  $32.68\% \pm 11.02\%$  of total OC, and it may fluctuate between 17.61% and 68.60%.

662

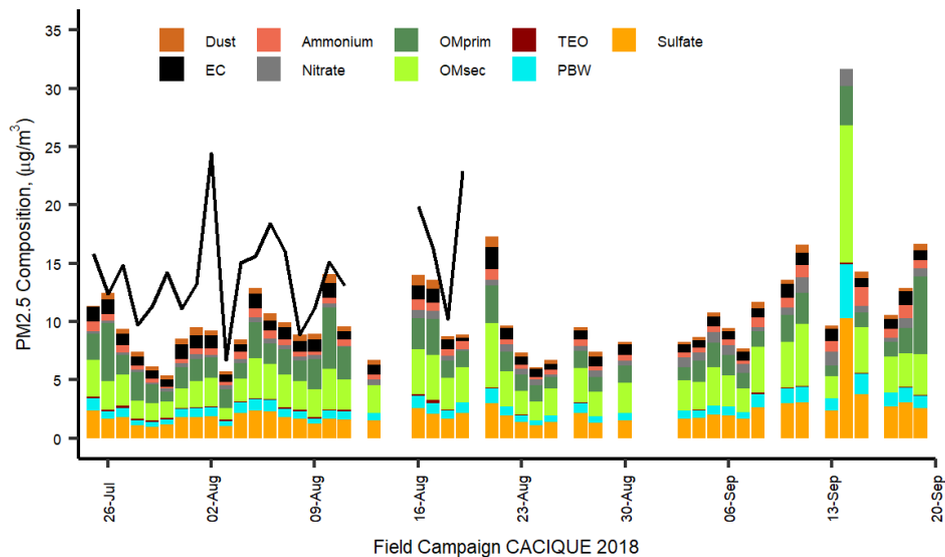
663 The difference between  $OC_{\text{prim}}$  from the organic tracer method and that obtained from the EC tracer method can be associated  
664 to the fact that the organic tracer method may not be representative of all sources. Industrial coal and fuel oil burning, garbage  
665 burning, cooking, charcoal production and other sources may not be accounted for by this method, since we did not have  
666 specific organic tracers for each of these activities.

667

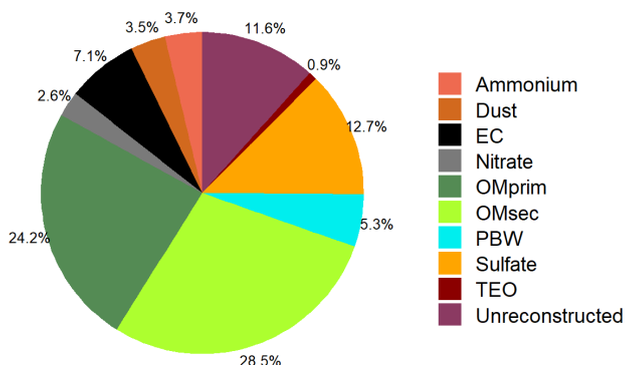
668 The mineral fraction, quantified as the sum of the oxides present in the crustal material (dust) and other TEO contributed  $3.51 \pm$   
669  $1.35\%$  and  $0.85 \pm 0.42\%$ , respectively. Despite the non-quantification of highly abundant mineral dust elements such as Si,  
670 the concentrations of Ca, Ti, and Fe indicated the impact of soil resuspension on the  $PM_{2.5}$  mass concentration.

671

672 PBW depends on the concentration of hygroscopic compounds embodied in the PM and the relative humidity of the weighing  
673 room where  $PM_{2.5}$  mass collected on the filters was determined. In this study, it was assumed that (i)  $\text{NH}_4^+$ ,  $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$   
674 were the main compounds responsible for absorbed water and (ii) thermodynamic equilibrium is dominated by these ions that  
675 allow calculating the  $\text{H}^+$  molar fraction as a difference between  $(\text{SO}_4^{2-} + \text{NO}_3^-)$  and  $\text{NH}_4^+$ , which is required to establish charge  
676 neutrality. Polar organic compounds and other water-soluble ions were not considered in the present study. The PBW content  
677 was estimated using the mean measured concentrations of  $\text{NH}_4^+$ ,  $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  in the AIM Model, where a multiplier factor  
678 of 0.32 was found as a proportion between the concentrations of the sum of these ions and the water fraction contained in  
679  $PM_{2.5}$ . As a result, PBW was 5.3% of the  $PM_{2.5}$  mass concentration.



680



681

682 Figure 6. Mass reconstruction of PM<sub>2.5</sub> collected in CRV. Figure in upper corresponding to timeseries of PM<sub>2.5</sub> gravimetric  
 683 mass measured and reconstructed mass from the chemical speciation in CRV during July – September 2018 and lower is the  
 684 to pie plot the relative mean contributions (%) of major chemical components of gravimetric PM<sub>2.5</sub> based on chemical  
 685 speciation.  
 686

#### 687 4. Conclusions

688 PM<sub>2.5</sub> samples collected in the Cauca River Valley, Colombia, were analyzed to determine the main chemical components of  
689 fine aerosol particles and to qualitatively identify aerosol sources using its chemical composition and diagnostic ratios. PM<sub>2.5</sub>  
690 during the campaign was  $14.4 \pm 4.4 \mu\text{g m}^{-3}$ . Its main components were OC ( $4.0 \pm 1.3 \mu\text{g m}^{-3}$ ), sulfate ( $2.2 \pm 1.4 \mu\text{g m}^{-3}$ ), and  
691 EC ( $1.0 \pm 0.3 \mu\text{g m}^{-3}$ ), ammonium ( $0.7 \pm 0.6 \mu\text{g m}^{-3}$ ), and nitrate ( $0.5 \pm 0.3 \mu\text{g m}^{-3}$ ). OM was estimated using the EC tracer  
692 method and the organic tracer method. Mass closure using the EC tracer method explained 88.4% of PM<sub>2.5</sub>, whereas the organic  
693 tracer method explained 70.9% of PM<sub>2.5</sub>. We attribute this difference to the lack of information of specific organic tracers for  
694 some sources, both primary and secondary. Organic material and inorganic ions were the dominant groups of species,  
695 constituting almost 79% of PM<sub>2.5</sub>. OM<sub>prim</sub> and OM<sub>sec</sub> from the EC tracer method contribute 24.2% and 28.5% to PM<sub>2.5</sub>.  
696 Inorganic ions made up 19.0%, EC 7.1%, dust 3.5%, PBW 5.3%, and TEO 0.9% of PM<sub>2.5</sub>.

697

698 Aerosol acidity was evaluated using three methods. The first, using the nitrate/sulfate ratio; the second using the anion/cation  
699 equivalent ratio; and the third, estimating the pH with the E-AIM thermodynamic model. All methods showed that the aerosol  
700 was acidic, with a pH of  $2.5 \pm 0.4$ , mainly because of the abundance of organic and sulfur compounds.

701

702 Diagnostic ratios applied to organic compounds indicate that most PM<sub>2.5</sub> was emitted locally and had contributions of both  
703 pyrogenic and petrogenic sources. In addition, levoglucosan and mannosan levels showed that biomass burning was ubiquitous  
704 during the sampling period. Fluoranthene (FLE) was the most abundant PAH, confirming the strong influence of BB associated  
705 with agro-industry. Five- and six-ring PAH associated with vehicular emissions were also abundant in PM<sub>2.5</sub>. Our  
706 measurements point to BB as the main source of PAHs in CRV. Relatively low PM<sub>2.5</sub> concentrations and mutagenic potentials  
707 are consistent with low-intensity, year-long BB and sugarcane PHB in CRV, which leads to lower atmospheric pollutant  
708 burdens and mutagenic potentials compared to those at locations where the harvesting period is shorter (*zafra*) thus with higher  
709 burning rates.

710 *Author contribution:* RJ, GR-S, and NR conceived and managed the project. LM-F, ACV-B, GR-S, and RJ set the instruments  
711 up and performed the aerosol sampling. LM-F carried out the sample chemical analysis at TROPOS with the guidance and  
712 support of DvP, MvP, KW, and HH. LM-F and ACV-B analyzed the measurement results, including PCA and other techniques  
713 with the support of DvP and LM-F, RJ, NR and ACV-B prepared the manuscript with substantial contributions from all the  
714 authors.

715 *Competing interests:* The authors declare that they have no conflict of interest.

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