Wet deposition of fossil and non-fossil derived particulate carbon:

Insights from radiocarbon measurement

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1 Abstract

Radiocarbon (¹⁴C) measurements of both organic carbon (OC) and elemental carbon (EC) 2 3 allow a more detailed source apportionment, leading to a full and unambiguous distinction 4 and quantification of the contributions from non-fossil and fossil sources. A thermal-5 optical method with a commercial OC/EC analyzer to isolate water-insoluble OC (WIOC) and EC for their subsequent ¹⁴C measurement was applied for the first time to filtered 6 7 precipitation samples collected at a costal site in Portugal and at a continental site in 8 Switzerland. Our results show that WIOC in precipitation is dominated by non-fossil 9 sources such as biogenic and biomass-burning emissions regardless of rain origins and 10 seasons, whereas EC sources are shared by fossil-fuel combustion and biomass burning. 11 In addition, monthly variation of WIOC in Switzerland was characterized by higher 12 abundance in warm than in cold seasons, highlighting the importance of biogenic 13 emissions to particulate carbon in rainwater. Samples with high particulate carbon 14 concentrations in Portugal were found to be associated with increased biogenic input. 15 Despite the importance of non-fossil sources, fossil emissions account for approximately 16 20% of particulate carbon in wet deposition for our study, which is in line with fossil 17 contribution in bulk rainwater dissolved organic carbon as well as aerosol WIOC and EC estimated by the ¹⁴C approach from other studies. 18

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20 Keywords: Radiocarbon, Particulate carbon, Precipitation, Fossil, Non-fossil

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23 **1. Introduction**

24 Carbonaceous particles (total carbon, TC) are of great importance due to effects 25 on human health and earth's climate (Pöschl, 2005; IPCC, 2013). Particulate carbon in 26 precipitation has been investigated in various locations across the globe, including sites 27 in urban, rural, mountain, coastal and marine areas. Wet deposition is known to be a key 28 scavenging (removing) process of carbonaceous particles (Fig. 1). Much of this research 29 has been focused on the abundance of different carbon fractions (elemental carbon, EC, 30 and water-insoluble organic carbon, WIOC), their spatial distribution, incorporation of 31 particles in hydrometeors and wet deposition fluxes (Cadle et al., 1980; Ducret and Cachier, 1992; Chylek et al., 1999; Cerqueira et al., 2010). In contrast, rather less attention 32 33 has been paid to the identification of sources (e.g. fossil and non-fossil emissions) and the 34 assessment of their relative contribution to particulate carbon.

Radiocarbon (¹⁴C) measurements have been widely used to distinguish contemporary and fossil carbon in ambient air aerosol samples in either bulk TC (Szidat et al., 2013; Zotter et al., 2014b) or in different carbonaceous fractions (Szidat et al., 2009; Zhang et al., 2014a, 2014b). Source attribution is based on a simple model which assumes that emissions from biogenic processes or biomass combustion are labelled with the ambient ¹⁴C/¹²C ratio and emissions from fossil-fuel combustion contain no ¹⁴C because of the large age of this material (Currie, 2000).

To the best of our knowledge this approach has never been used in source apportionment studies of WIOC and EC in precipitation samples (rain and snow), although it was applied in some cases to deposited snow of ice core research (Jenk et al., 2006; Cao et al., 2013) as well as total organic carbon (TOC) and dissolved organic carbon (DOC) in rain (Raymond, 2005; Avery et al., 2006, 2013). One of the most challenging problems of the method lies in lower concentration of particulate organic

carbon (POC) compared to DOC and offline separation methods for ¹⁴C measurements in 48 49 water-insoluble total carbon (WITC) and, especially, different carbon fractions such as 50 EC and WIOC. Thermal-optical analysis (TOA) is one of the most commonly used 51 methods for measuring the OC/EC concentration in aerosol and precipitation samples. In 52 most TOA methods such as NIOSH, IMPROVE and EUSAAR protocols, the filter 53 sample is heated sequentially in an inert (He) and oxidative (He/O₂ mixture) atmosphere 54 (Cavalli et al., 2010). Ideally, OC and EC are released in each step separately. However, 55 part of OC is formed as charring (EC-like material that is refractory and light-absorbing), 56 which occurs in the first step and is mistakenly detected as EC in the second step. 57 Therefore, a laser is used to monitor the filter transmittance throughout analysis to 58 account for the charring contribution. The charring correction is then carried out to 59 quantify OC/EC mass concentration by assigning a split point at the time when the 60 transmittance returns to its initial value during the He/O₂ step (Cavalli et al., 2010). 61 However, the charring correction is not feasible for ¹⁴C analysis of OC and EC, since this 62 requires a physical separation of the two fractions and the split point is unknown before 63 measurement. This was recently improved by a thermaloptical method utilizing both the 64 special thermal and optical properties of OC and EC, thereby optimizing both the 65 quantification and isolation of OC and EC (Zhang et al., 2012).

Aerosol pollution from wood burning has recently been recognized in large-scale areas of Europe including Scandinavia, Portugal and alpine regions, as well as in major European cities such as London, Paris and Berlin despite the importance of fossil-fuel emissions (Szidat et al., 2009; Borrego et al., 2010; Fuller et al., 2013, 2014; Herich et al., 2014; Zotter et al., 2014a). However, current understanding on the impact of fossil versus biomass burning to aerosols is incomplete, since the source of WITC in wet deposition, a major removal process of particulate carbon in the atmosphere (Ducret and 73 Cachier, 1992), are still not well constrained. Therefore, the motivation for the present study was (1) to adapt a method that has been successfully applied for ¹⁴C analysis of 74 atmospheric aerosols to the quantification of ¹⁴C in particulate carbon extracted from 75 76 rainwater and (2) to quantify fossil and non-fossil contributions to particulate carbon 77 including OC and EC in precipitation from samples collected in Portugal and Switzerland. 78 This study also provides new insights into the wet deposition of particulate carbon emitted 79 by both fossil and non-fossil sources, and improves our understanding of the role of 80 atmospheric particulate carbon in the models of the global carbon cycle.

81 2. Experimental

82 2.1 Measurement sites and analytical method

83 Investigations were carried out at two different sites, Aveiro in Portugal (40° 38' 07''N 84 and 8° 39' 35''W) and Dübendorf (47° 24' 11''N and 8° 36' 48''W) in Switzerland. 85 Sampling at Aveiro was conducted at the University of Aveiro campus, located at the 86 southwestern rim of the city of Aveiro, western coast of Portugal. The campus is about 7 87 km from the sea-side and there are no significant pollution sources in the area between it 88 and the ocean. The roof of a small container placed near the campus meteorological 89 station was used as a platform to collect rain samples, at about 4 m above ground level. 90 Sampling at Dübendorf was performed on the premises of EMPA located outside of 91 Dübendorf, a city with 25,000 inhabitants and a suburb of the city of Zurich (400,000 92 inhabitants). The site therefore is of suburban type located 432 m a.s.l. in the northern 93 part of Switzerland. The rain samples have been collected at 2 m above ground level.

94 2.2 Sampling method

Rainwater samples in Portugal were collected on an event basis from mid-February until
the end of April 2011 with an Eigenbrodt model UNS130/E automatic wet-only collector.
The sampler consisted of a glass funnel with an open area of 500 cm², connected to a 5 L

98 glass storage bottle, a movable lid and a precipitation sensor to control start and stop of 99 each collection period. Prior to use, all the collector components that could come in 100 contact with samples were cleaned with soap and water, followed by rinsing of them with 101 distilled and deionized water. Rainwater samples at Dübendorf, Switzerland were 102 collected on a daily basis from April 2012 to March 2013 with a Digitel model DRA-92 103 HK wet-only collector. The sampler consists of a Teflon-coated funnel with an open area 104 of 500 cm². The rain samples were collected in storage bottles (polyethylene, 250 mL) 105 which were automatically changed at midnight or when they were full. The funnel and 106 the subsequent Teflon tube have monthly been cleaned with ultrapure water, the storage 107 bottles have been rinsed with ultrapure water before they were used. One field blank at 108 each site was prepared by pouring ultrapure water (50 mL) through the sampling device, 109 collecting the rinsate in a storage bottle and, after staying one day inside the closed 110 collector, processed under the same conditions as samples.

111 2.3 Filtration

112 The particulate carbon in the rainwater samples were collected on prebaked quartz fibre 113 filter (Pallflex Tissuguartz, 2500QAO-UP) with a diameter of 10-11 mm by filtration 114 using vacuum suction. Prior to the use of containers and the filtration funnel, they were 115 rinsed three times with ~25 mL ultra-pure water with low carbon impurity in order to 116 reduce the blank. A series of three filters were used to collect particles to increase 117 filtration efficiency. The filtration efficiency with a similar approach was reported to be 118 $92 \pm 7\%$ using prepared standards of known quantities of soot in water (Hadley et al., 119 2008). Filtered water volumes were typically in the range of 250-700 mL. For samples 120 from Portugal (n = 7), daily rain samples were filtered at the University of Aveiro before sending to the University of Bern for ¹⁴C analysis by express mail in a cooled container. 121

122 For samples from Switzerland (n = 12), rain events were pooled per month for filtration,

123 which was conducted at the University of Bern.

124 2.4 Determination and separation of different carbonaceous particle fractions

125 OC and EC fractions extracted from precipitation samples were isolated by the Swiss 4S protocol for subsequent ¹⁴C measurement. This method was first described in Zhang et 126 127 al. (2012) for analyzing ambient aerosols samples and recently applied for particulate 128 carbon extracted from firn/ice samples by Cao et al. (2013). In brief, WIOC is combusted 129 in pure oxygen at 375 °C for 180 s in step one and subsequently EC is combusted in pure 130 oxygen at 760 °C for 120 s in step four after complete OC removal in pure oxygen at 475 131 °C for 120 s (step two) and in helium at 650 °C for 180 s (step three). The evolving CO₂ 132 is quantified by a non-dispersive infrared (NDIR) detector, separated from interfering reaction gases, cryotrapped and sealed in glass ampoules for ¹⁴C measurements. By using 133 134 the Sunset OC/EC analyzer, which monitors the filter optical properties during the 135 combustion with a laser, the quantification of the EC losses before step four is also 136 possible. Therefore, mass concentration of WIOC, EC and WITC can be quantified as 137 well. A good agreement of carbon mass determination by the Swiss 4S and the 138 EUSAAR 2 protocol was previously reported for aerosol samples (Zhang et al., 2012). 139 The extent of charring formation during the thermal treatment can be reduced by using 140 pure oxygen instead of helium, which is evident by the comparison of thermograms 141 applying different protocols (see Supplemental material, Figure S1). However, substantial 142 OC charring still occurs, sometimes larger than 10% of EC mass, which results in a positive bias of the ¹⁴C measurement of the EC fraction, as OC and EC often differ in 143 144 their ¹⁴C content (Szidat et al., 2009; Zhang et al., 2013). Since some charred material is very likely removed during step three before the EC step, we assume that $50 \pm 15\%$ of 145

146 the charred OC remains in the EC fraction. Therefore, the fraction of modern (f_M, see Sec

147 2.4) of EC ($f_{M,EC}$) as corrected for charring ($f_{M,EC,final}$) is as follows:

148 $f_{M, EC corr} = (mC_{EC} * f_{M, EC} - mC_{charring} * f_{M, charring}) / (mC_{EC} - mC_{charring}),$

- $149 \qquad \text{where } f_{M,EC} \text{ and } f_{M,charring} \text{ denote the } f_M \text{ measured and } mC_{EC} \text{ and } mC_{charring} \text{ the carbon}$
- 150 mass for EC and charred OC, respectively.
- 151 f_{M,charr} is assumed to equate to the measured f_{M,OC}, as charred OC is part of the OC
- 152 fraction. The typical uncertainties of source apportionment results arise from analytical
- 153 uncertainties (i.e. OC/EC mass and ¹⁴C measurements), blank/charring correction and
- 154 the variability of reference $f_{M,nf}$ values (see Sec 2.5) approximately amounted to $\pm 20\%$,
- $\pm 20\%, \pm 15\%$ and $\pm 10\%$ for fossil EC, non-fossil EC, fossil OC and non-fossil OC,
- 156 respectively.
- 157 $2.5^{14}C$ analysis

¹⁴C analyses of CO₂ produced samples from OC/EC analyzer were measured with the 158 159 Mini Carbon Dating System (MICADAS) at the Laboratory for the Analysis of 160 Radiocarbon with AMS and at ETH Zurich, both of which using a gas ion source, which 161 allows direct CO₂ injection after dilution with He (Wacker et al., 2013; Szidat et al., 2014). 162 14 C results are usually expressed as fractions of modern (f_M), which were further 163 converted to fraction of non-fossil (fNF) by dividing by fM reference values for 164 contemporary carbon for the source apportionment of non-fossil and fossil carbon. fM 165 reference value of OC ($f_{M,ref}$) were estimated as 1.08 ± 0.05 for non-fossil continental 166 sources for both sites according to (Zhang et al., 2014a). For samples influenced by 167 marine sources at the Portugal sampling site, this reference value was estimated as 0.98 168 ± 0.04 (see Sec 3.1.1, and Figure S2) following the concept of (Ceburnis et al., 2011), 169 which is currently the best available approach for this determination. It should be noted 170 that the uncertainty of source-specific contributions are dependent on the selected f_M

- 171 reference values. By varying reference f_M by ± 0.05 , the uncertainty is typically within 172 5%. A mass-dependent blank correction is applied to the measured f_M values for OC 173 according to an isotopic mass balance approach. The f_M and OC value of the blank was 174 0.59 ± 0.02 and $1.2 \pm 0.8 \mu gC$ per filtration, respectively. No blank correction was carried 175 out for ¹⁴C measurement of EC as carbon amount of blank EC ($0.4 \pm 0.4 \mu gC$) was below 176 the detection limit (~2-3 μgC) of the ¹⁴C analysis.
- 177 **3. Results and discussion**
- 178 3.1 Particulate OC and EC levels in precipitation
- 179 *3.1.1 Aveiro, Portugal*

180 Seven rain samples were characterized for their WITC concentrations and the 181 corresponding ¹⁴C values. Details about rain events are reported in Table S1 and 182 analytical results are presented in Fig. 2. Backward air mass trajectories arriving at Aveiro 183 during the rain events generated with the HYSPLIT model are shown in Figure S2. 184 According to air mass origins, 7 samples were classified into 2 types: marine (TYPE1: 185 samples AVE1 to AVE5) where the airmass travelling over the ocean and continental 186 influenced rainwater (TYPE2: samples AVE6 and AVE7) with a mixture of continental 187 and marine influences. The average WITC was $128.4 \pm 87.7 \,\mu$ g/L and $272.1 \pm 42.3 \,\mu$ g/L 188 for marine and continental influenced rainwater, respectively. The lowest WITC 189 concentrations were found in rain samples AVE1, AVE3 and AVE5, with a mean of 66 190 \pm 12 µg/L. With a similar air mass origin (Figure S2), WITC in sample AVE2 and AVE4 191 was significantly increased. These two samples were likely influenced by additional local 192 emissions. Indeed, plant debris (or detritus) was visible after filtration of the sample 193 AVE2, explaining the high WITC concentration. This sample was taken under conditions 194 with elevated wind speed compared to the other samples (Table S1), which might have 195 been the cause of a significant input of plant debris into local air. On the other hand,

196 pollen grains were observed after filtration of sample AVE4. Marine pine (Pinus pinaster) 197 is the dominant tree in north-western Portugal and airborne pollen from this species is 198 known to have a seasonal variation with a peak during March (Sousa et al., 2008). Pollen 199 grains of marine pine deposit intensely at this time of the year and are frequently observed 200 as a yellow dust accumulated on outside surfaces. In addition, the average WIOC/EC ratio 201 (5.7) of samples AVE2 and AVE4 were significantly higher than that (3.2) in samples 202 AVE1, AVE3 and AVE5, indicating that the larger TC abundance for samples AVE2 and 203 AVE4 is rather due to biogenic/biomass sources than anthropogenic sources. The higher 204 WITC concentrations detected in TYPE2 were probably due to higher aerosol pollution 205 in continental air-mass origins. Actually, during the rain event of 18-19 of April (sample 206 AVE6), air masses originated from the western coast of Africa and were transported over 207 densely populated areas in coastal Portugal before arriving at the sampling site. Sample 208 AVE7 reflects the influence of two types of transport: whereas air masses arrived from 209 northwest, after crossing France and Spain, during the first half of the sampling period, 210 the second half was dominated by air-mass transport over northern Africa, southern Spain 211 and inland Portugal. However, the source of the continental-influenced samples remains 212 unknown, which we will discuss below (Sec 3.2.1).

213 3.1.2 Dübendorf, Switzerland

WITC concentration levels in the precipitation samples from Dübendorf ranged from 73 μ g/L to 228 μ g/L, with an annual average of 147±58 μ g/L (Figure 3). The monthly WIOC concentrations were lower in cold seasons than in warm seasons, associated with larger biogenic emissions in both spring and summer. Although wood burning in winter is a dominant contributor of airborne OC in Switzerland (Zotter et al., 2014a), this may not affect particulate OC fraction in precipitation significantly, as a large fraction of woodburning OC may be dissolved into the rainwater due to high water solubility and 221 thus mainly exist in the rainwater DOC fraction. Previous studies reported higher levels 222 of DOC in rainwater in warm seasons in other regions, such as coastal New Zealand and 223 North Carolina, USA, as biogenic emissions from vegetation is largely controlled by 224 seasonality (Willey et al., 2000; Kieber et al., 2002). However, this seasonality was not 225 found for EC, as this fraction is only produced by fossil-fuel combustion and biomass 226 burning, but not by biogenic emissions. Actually, monthly EC concentrations showed 227 relatively small variations throughout the year in contrast with the fact that EC abundance 228 in ambient aerosol samples has been reported to be highest in winter due to accumulation 229 during winter-smog episodes and enhanced residential wood burning in Switzerland 230 (Herich et al., 2011). It should be noted that precipitation seldom occurs during winter-231 smog episodes and most precipitation events in winter are snow events so that different 232 wet scavenging efficiencies between rain and snow events may play a role on wet 233 deposition of EC. Correlation with the amount of precipitation was neither observed for 234 WITC, WIOC, or EC concentrations, and also not for their ¹⁴C level, probably because of 235 pooling monthly averaged samples.

236 3.2 Sources of particulate carbon in precipitation: fossil vs. non-fossil sources

237 *3.2.1 Aveiro, Portugal*

238 In general, both natural and anthropogenic sources including biogenic emissions, biomass 239 burning and fossil-fuel combustion can contribute to rainwater particulate carbon. In this 240 work, the source apportionment of particulate carbon in precipitation was made based on 241 14 C measurements, assuming non-fossil carbon has the same f_M as the contemporary atmosphere and fossil fuel carbon is devoid of ¹⁴C. As samples AVE2 and AVE4 were 242 influenced by large primary biogenic particulate matter, ¹⁴C measurement has only been 243 244 performed on bulk WITC, because large primary biogenic particulate matter may 245 significantly affect the WIOC/EC separation. The average non-fossil contribution to

246 WIOC is 84% and 91% for marine and continental influenced rainwater, respectively. 247 The fossil components found in rainwater at this coastal site likely reflect the export of 248 fossil emissions from the North American continent, the input of emissions from ship and aircraft emissions as well as local traffic emissions. ¹⁴C measurement of samples with the 249 250 highest WITC content (samples AVE2 and AVE4) confirm, however, that these two 251 samples are indeed influenced by nonfossil emissions (i.e. primary biogenic emissions). 252 Non-fossil contributions (i.e. biomass burning) to EC were on average $52\% \pm 2\%$ and 58%253 \pm 2% for TYPE1 and TYPE2, respectively, which was lower than the corresponding non-254 fossil fraction in OC with average contributions of $84\% \pm 2\%$ and $90\% \pm 2\%$. Biomass 255 burning EC mainly originates from residential wood burning for heating and/or open 256 biomass burning. Wood burning has been reported to be a considerable contributor to EC 257 by many previous studies conducted in Europe (Szidat et al., 2009; Fuller et al., 2014; 258 Zotter et al., 2014a). The MODIS fire counts map 259 (https://firms.modaps.eosdis.nasa.gov/firemap/, Figure S3) during April shows that 260 intensive open biomass-burning activities took place in upwind direction of and/or in 261 surrounding regions of the sampling site, which could be another explanation of the 262 enrichment of non-fossil carbon in both OC and EC fractions of samples AVE6 and 263 AVE7. Therefore, high abundance particulate OC found in continental influenced 264 rainwater is not due to larger urban fossil emissions, but due to enhanced biogenic and biomass-burning emissions according to ¹⁴C results in both WIOC and EC fractions. 265

266 3.2.2 Dübendorf, Switzerland

267 On average, non-fossil sources contributed $84 \pm 5\%$ to OC on an annual average, which 268 was always higher than non-fossil EC with a mean contribution of $48 \pm 7\%$ (Fig. 3). This 269 finding agrees with the fact that OC/EC emissions ratios are higher in biogenic or biomass 270 burning sources than those in fossil fuel emissions (i.e. mainly vehicular emissions in 271 Europe) (Pio et al., 2011). A notable seasonality was not found for relative fossil and non-272 fossil contributions to OC, whereas non-fossil EC fraction was slightly higher in winter 273 than other seasons due to larger wood burning emission for heating during cold period 274 (Zhang et al., 2013; Zotter et al., 2014a). The high non-fossil fractions in rainwater WIOC 275 are attributed to biogenic and wood burning emissions. Based on a carbon isotope mass 276 balance (i.e. $f_{NF}(WITC)$ * mC_{WITC} = $f_{NF}(OC)$ *mC_{OC} + $f_{NF}(EC)$ *mC_{EC}), the annual-average non-fossil contribution to WITC was $79 \pm 5\%$ with a consistent non-fossil predominance 277 278 throughout the year, which dominated the wet deposition of atmospheric organic aerosols. Previous studies also found similar ¹⁴C values for bulk rain DOC, although there are no 279 studies reporting ¹⁴C measurements on POC in rainwater to the best of our knowledge. 280 For example, ¹⁴C measurements of rain DOC from two sites in the northeast USA suggest 281 282 approximately 78% and 85% was non-fossil origin, respectively (Raymond, 2005; Avery 283 et al., 2006).

284 3.3 Summary and discussions

This paper reports the first successful application of radiocarbon (¹⁴C) analysis to water-285 insoluble particles extracted from precipitation. Based on ¹⁴C measurement of particulate 286 287 OC and EC of rainwater samples at a costal site in Portugal and a continental site in 288 Switzerland, particulate carbon in precipitation is dominated by non-fossil sources, such 289 as biogenic and biomass-burning emissions regardless of season as well as air-mass and 290 rain origins. It should be worth noting that non-fossil contributions in total particulate 291 carbon in precipitation in both studied sites were higher than those reported in airborne 292 aerosol samples. For example, Gelencsér et al. (2007) showed that non-fossil contribution 293 fraction was on average 73% and 81% of TC in PM2.5 aerosol samples collected during 294 summer and winter at Aveiro, Portugal, respectively, (Gelencsér et al., 2007), slightly 295 lower than our finding (averaged to 81%) in rainwater samples. Recently, a yearly cycle

of ¹⁴C measurement of OC and EC of PM₁₀ aerosol samples was investigated at an urban 296 297 background station in Zurich, Switzerland (Zotter et al., in preparation). Non-fossil 298 contributions to TC is on average 15% lower compared to those extracted from rainwater samples collected at Dübendorf, 6 km north of Zurich. In addition, they found a very clear 299 300 seasonal cycle for relative non-fossil contribution of EC (ranging from 9 to 50%) in PM₁₀ 301 aerosol samples, whereas in the precipitation analysed in this study, less seasonal 302 variation (ranging from 37 to 59%) was observed. We hypothesize that particles emitted 303 from biomass-burning processes may be coated with more hydrophilic compounds such 304 as inorganic substances and oxidized organic aerosols, which may increase the wet 305 removal efficiency of non-fossil particles. By examining aerosol size distributions and 306 BC coating properties sampled in three Canadian boreal biomass burning plumes, Taylor 307 et al. (2014) have also demonstrated that less coated EC particles are less effectively 308 removed by wet deposition (Taylor et al., 2014). Moreover, it should be emphasized that 309 precipitation includes larger particulate matter than PM₁₀ aerosol, which may have an influence on the outcome of the ¹⁴C source apportionment, as we recently showed for fine 310 311 and coarse aerosol fractions (Zhang et al., 2013). Further experiments with simultaneous 312 measurements on both aerosol and precipitation samples are required to study wet 313 scavenging processes and mechanisms for carbonaceous particles from both fossil and 314 non-fossil origins.

The findings from our study suggest that wet deposition preferentially removes non-fossil derived carbonaceous particles. Despite of the importance of non-fossil sources, fossil emissions account for approximately 20% of total particulate carbon in wet deposition for our study, which is in line with fossil contribution in bulk rainwater DOC as well as aerosol WIOC and EC estimated by the ¹⁴C approach from other studies (Raymond, 2005; Avery et al., 2006; Szidat et al., 2009; Zotter et al., 2014a). Based on literature studies 321 and our observations, a very gross estimation of wet-deposited fossil-derived carbon was 322 also carried out. By assuming 430 Tg C/y is removed from the atmosphere by rainwater 323 DOC (Willey et al., 2000) and multiplying it by the average POC percentages (15%) in 324 TOC (the sum of POC and DOC) from previous studies (Sempere and Kawamura, 1996; 325 Willey et al., 2000) as well as an average fossil contribution (20%) to POC from our 326 current study could approximately yield a net removal of 12.9 Gt fossil-derived C per 327 year in the particulate phase via wet deposition. This gross estimation is within the upper 328 range of bottom-up primary organic aerosols from fossil fuel emissions of 5-15 TgC/y 329 (Bond et al., 2004; Hallquist et al., 2009), suggesting that the majority of fossil-derived 330 primary organic aerosols are removed by wet deposition. Therefore, fossil-derived 331 particulate carbon can be transferred between different ecosystem (e.g. from land to ocean 332 or from ocean to land) via wet deposition, and it can play an important role on the Earth's environment and human health via biogeochemical carbon cycle. 333

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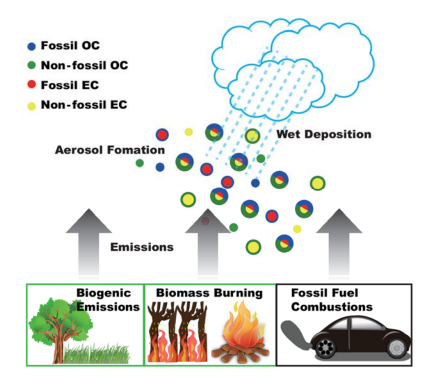
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- **Figure 1**: A simplified sketch showing the sources and wet deposition of carbonaceous
- 481 particles (organic and elemental carbon).

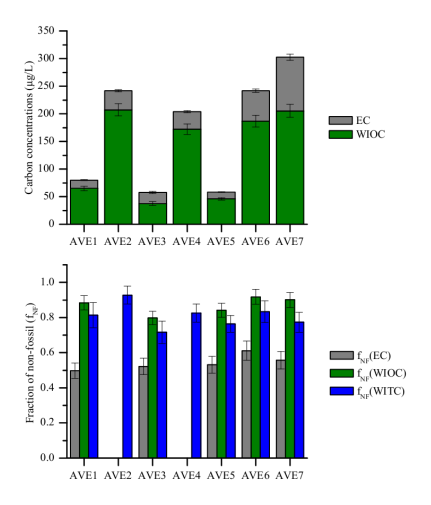


Figure 2. WIOC and EC concentrations and fraction of non-fossil (f_{NF}) values of

485 WIOC, EC and WITC for rain samples collected at Aveiro, Portugal.

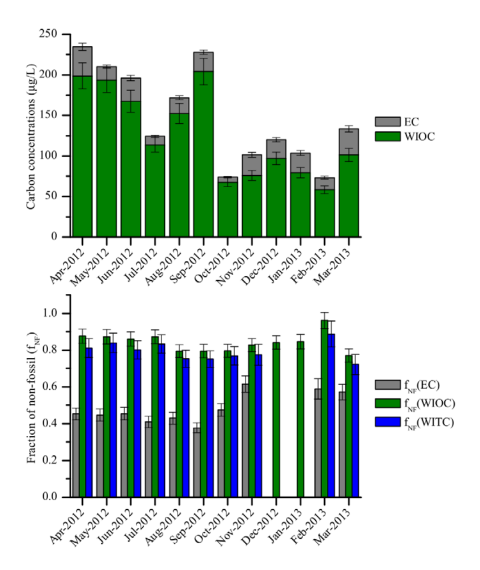


Figure 3. Monthly WIOC and EC concentrations and fraction of non-fossil (f_{NF}) values
of WIOC, EC and WITC for precipitation samples collected at Dübendorf, Switzerland.