

# Diurnal aging of biomass burning emissions: Impacts on secondary organic aerosol formation and oxidative potential

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## 1 Abstract

2 Residential biomass burning is an important wintertime source of aerosols. These particles are  
3 subjected to complex diurnal aging processes in the atmosphere, contributing to urban and  
4 regional air pollution. The cumulative impact of these aging cycles on aerosol composition and  
5 oxidative potential, a key toxicity metric, remains unclear. This study examined the oxidation  
6 cycles of biomass burning emissions during day-to-night and night-to-day transitions in the  
7 FORTH (Foundation for Research and Technology – Hellas) atmospheric simulation chamber,  
8 focusing on emissions from burning of olive wood. The final high-resolution AMS spectra of  
9 biomass burning organic aerosol (bbOA) after either oxidation cycle were almost identical ( $R^2$   
10 > 0.99,  $\theta = 3^\circ$ ). This indicates transformation into similar biomass burning secondary organic  
11 aerosol (bbSOA) regardless of the initial step of the diurnal cycle. A 56% average increase in  
12 the bbOA oxygen-to-carbon (O:C) ratio was observed during both cycle cases (from  $0.38 \pm$   
13 0.06 for the fresh to  $0.59 \pm 0.07$  after aging). Additional OA mass was produced after the two  
14 cycles, varying from 35 to 90 % of the initial OA. The aging of the emissions led to a final  
15 water-soluble oxidative potential (WS-OP) increase of 60% to  $68 \pm 18 \text{ pmol min}^{-1} \mu\text{g}^{-1}$  for both  
16 cycles, but with notably different transient values that depend on the order of the oxidation  
17 regimes. The effect of each oxidation regime on the WS-OP of the bbOA depends on the  
18 airmass history. The evolution of the WS-OP was not well correlated with that of the O:C.

19 **1 Introduction**

20 Biomass burning for residential heating has significantly increased over the past two decades  
21 in several countries, primarily driven by rising energy costs and efforts to reduce the use of  
22 fossil fuels (Alper et al., 2020). Alongside contributions from wildfires, residential biomass  
23 burning has emerged as a major source of urban and regional pollution worldwide (Zauqi-  
24 Sajani et al., 2024). Solid biomass currently represents nearly 45% of the total bioenergy supply  
25 in the EU, 40% of which is allocated to residential heating, with an anticipated 20% increase  
26 projected by 2050 (IEA, 2019, 2021; Reid et al., 2020). This upward trend in the residential  
27 burning of solid biomass, particularly wood, has raised serious concerns regarding air quality  
28 and human health (Cincinelli et al., 2019; Guercio et al., 2021; Pardo et al., 2024).

29 Particles emitted from biomass burning consist of organic compounds, elemental carbon  
30 (EC), sulfates, nitrates, ammonium, and ash (Jiang et al., 2024). Biomass burning emissions  
31 also include a range of gases; carbon monoxide (Shen et al., 2020), volatile organic compounds  
32 (VOCs) such as aldehydes, ketones, and organic acids (Zhang et al., 2021; Huang et al., 2022),  
33 carcinogenic polycyclic aromatic hydrocarbons (PAHs and oxy-PAHs) (Tsiodra et al., 2021,  
34 2024; Lim et al., 2022), as well as nitrogen oxides and ammonia (Bray et al., 2021). The emitted  
35 VOCs contribute to the formation of biomass burning secondary organic aerosol (bbSOA) and  
36 can have direct health effects (Fang et al., 2021). The emission profile of these pollutants is  
37 variable, influenced by factors such as fuel type and quality (e.g., logs vs. pellets; hardwood  
38 vs. softwood; certified vs. non-certified wood, moisture content etc.), burning conditions (e.g.,  
39 flaming vs. smoldering, air/oxygen supply, and dilution), and the type of combustion appliance  
40 (Fachinger et al., 2017; Nyström et al., 2017; Price-Allison et al., 2021; Trubetskaya et al.,  
41 2021).

42 After their release, biomass burning emissions are subject to chemical transformations  
43 through homogeneous or heterogeneous reactions, that differ between daytime and nighttime  
44 (Donahue et al., 2012; Hodshire et al., 2019; Yazdani et al., 2023). During these reactions, a  
45 significant amount of SOA (Yazdani et al., 2023) and reactive oxygen species (ROS) (Wang  
46 et al., 2023) can be generated. Hennigan et al. (2011) reported significant variability in bbSOA  
47 formation during the photo-oxidation of different emissions. Yazdani et al. (2023) reported that  
48 after 6 to 10 hours of daytime exposure, up to 30% (with an average of 15%) of the primary  
49 bbOA (bbPOA) mass was oxidized, forming bbSOA that was predominantly composed of  
50 acids. The coupled gas-particle partitioning, and reaction of semi-volatile vapors (SVOCs) may  
51 play an important role in the processing of bbPOA (Hennigan et al., 2011; Srivastava et al.,

52 2022). Li et al. (2024) demonstrated that intermediate volatility species (IVOCs) can contribute  
53 approximately 70% of the formed bbSOA, more than twice the contribution from VOCs.

54 The nighttime oxidation of biomass burning emissions by the nitrate radical ( $\text{NO}_3$ ) also  
55 leads to rapid aerosol changes (Kodros et al., 2020), but to a lesser extent compared to OH  
56 oxidation (Yazdani et al., 2023). In some cases, a doubling of bbOA levels compared to the  
57 initial primary bbOA has been observed. This increase has been attributed to gas-phase  
58 reactions between the  $\text{NO}_3$  radical and mainly phenolic compounds or furanic aldehydes  
59 (Hartikainen et al., 2018). Moreover, a substantial increase (7-100%) in the aerosol oxygen-to-  
60 carbon (O:C) ratio, as well as in the mass of organic nitrates in bbOA has been reported, as  
61 result of nocturnal aging (Kiendler-Scharr et al., 2016; Kodros et al., 2022; Yazdani et al.,  
62 2023).

63 To date, field and atmospheric simulation chamber studies have focused on the oxidation  
64 of biomass burning emissions during either daytime or nighttime oxidation regimes, driven  
65 respectively by OH and  $\text{NO}_3$  radicals (Hennigan et al., 2011; Fry et al., 2014; Hodshire et al.,  
66 2019; Jorga et al., 2021; Kodros et al., 2022; Wang et al., 2023; Yazdani et al., 2023). While  
67 such investigations have significantly advanced our understanding of the individual effects of  
68 these oxidation regimes, they do not fully capture the real-world evolution of biomass burning  
69 aerosols, which undergo multiple repeated cycles of daytime and nighttime chemistry during  
70 their atmospheric lifetime. Studies on successive aging from daytime and nighttime cycling do  
71 exist, but have focused on the changes of the optical and chemical properties of bbOA and the  
72 gas-particle phase partitioning of semi- and intermediate-volatility organic compounds (Tiitta  
73 et al., 2016; Hartikainen et al., 2018; Cappa et al., 2020; Che et al., 2022; Desservetaz et al.,  
74 2023; Yazdani et al., 2023). These alternating oxidation regimes cause successive changes in  
75 chemical composition, reactivity, and toxicity (Li et al., 2021, 2023; Tomlin et al., 2022; He et  
76 al., 2024) that are not well understood. Consequently, the timing of atmospheric BB emissions,  
77 being released during the day or night, may also influence the chemical trajectory of BB aerosol  
78 aging and therefore affect its composition and properties, including toxicity.

79 Biomass burning particles are significant sources of reactive oxygen species (ROS),  
80 including free radicals (e.g., OH,  $\text{RO}_2$ ,  $\text{HO}_2^-$ ) and non-radicals (e.g.,  ${}^1\text{O}_2$ ,  $\text{H}_2\text{O}_2$ ). Upon  
81 inhalation, these species interact with biological tissues and can disrupt cellular redox balance,  
82 triggering (or propagating) oxidative stress and systemic health effects (Costabile et al., 2023).  
83 The ability of particulate matter (PM) to catalyze ROS production, known as oxidative  
84 potential (OP), is a critical metric linking aerosol exposure to health outcomes (Zhang et al.,  
85 2022; Dominutti et al., 2025). Among the various in vivo and in vitro methods developed to

86 quantify OP (Ng et al., 2019), the abiotic dithiothreitol (DTT) assay is the most well established  
87 one, providing a measure of the water-soluble OP (WS-OP) of aerosols through the depletion  
88 of surrogate DTT in aerosol extracts (Cho et al., 2005). In this assay, DTT acts as a surrogate  
89 biological reducing agent that is oxidized by redox-active PM components. The remaining DTT  
90 is determined via a colorimetric reaction with DTNB (5,5'-dithiobis-(2-nitrobenzoic acid)),  
91 producing a light-absorbing compound measured spectrophotometrically at 412 nm. Blank-  
92 corrected depletion rates are typically normalized to aerosol mass or organic carbon content to  
93 provide a per mass health-relevant measure of WS-OP. The broad sensitivity of this method to  
94 diverse sources of ROS in aerosols with long lifetimes (Gao et al., 2020a; Rao et al., 2020),  
95 along with its optimization over the years (Fang et al., 2015; Puthussery et al., 2020) to provide  
96 more rapid measurements of water-soluble OP (WS-OP), makes it highly suitable for large-  
97 scale studies. In this study, we focus on WS-OP as a partial measure of aerosol toxicity,  
98 acknowledging that the DTT assay reflects only one aspect of oxidative potential, and its direct  
99 link to health outcomes remains uncertain. Recent studies emphasize that no single OP assay  
100 can fully represent particle toxicity and that complementary approaches (e.g., thiol- and  
101 hydroxyl radical-based assays) are needed to capture the full range of oxidative mechanisms  
102 and to strengthen links with health-relevant outcomes (Dominutti et al., 2025). Studies using  
103 the DTT assay have identified bbOA and SOA as dominant contributors to DTT activity,  
104 accounting respectively for 35% and 30% of total OP in ambient aerosols in the Southeastern  
105 USA (Verma et al., 2015). More recent studies confirm that biomass burning is a significant  
106 source of OP in diverse environments, highlighting the importance of understanding diurnal  
107 variations in OP from biomass burning (Paraskevopoulou et al., 2019, 2022; Mylonaki et al.,  
108 2024).

109 Photochemical aging during daytime oxidation promotes particle-bound ROS  
110 production, enhancing the OP of the aged aerosols (Li et al., 2021; Wang et al., 2023). For  
111 bbOA, the OP was found to increase by a factor of two ( $2.1 \pm 0.9$ ) after multiple days (68 h) of  
112 atmospheric aging (Wong et al., 2019). This implies that the health impacts of bbOA may  
113 extend far from its sources, as it ages and becomes part of the background aerosol  
114 (Vasilakopoulou et al., 2023; Mylonaki et al., 2024).

115 While it is well-established that bbOA ages rapidly at night, the effects of its nocturnal  
116 aging on aerosol OP are poorly understood. Moreover, to our knowledge no studies have yet  
117 investigated how the oxidation sequence (day-to-night and night-to-day) affects aerosol  
118 chemical composition, aging trajectory, and toxicity (i.e., evolution of OP). This study aims to  
119 address these knowledge gaps through controlled chamber experiments simulating realistic

120 diurnal oxidation cycles. In these experiments, fresh biomass burning emissions undergo  
121 sequential aging, either through daytime oxidation followed by nighttime oxidation or the  
122 reverse. By comparing day-to-night and night-to-day sequences, we aim to elucidate the  
123 interplay of oxidation regimes on aerosol chemical evolution and OP, providing novel insights  
124 into the health impacts of diurnally aged biomass burning aerosols.

## 125 **2 Methods**

### 126 **2.1 Atmospheric simulation chamber experiments**

127 Emission aging experiments took place at the FORTH-ASC chamber facility at Patras,  
128 Greece. Figure 1 illustrates the setup used for conducting the experiments. Fresh biomass  
129 burning emissions were produced in the combustion facility beneath FORTH-ASC by a  
130 residential wood stove, fed with commercially available olive wood logs and branches. This  
131 type of hardwood is widely used as a fuel in Greece. The emissions were diluted before their  
132 injection into the smog chamber, using a custom-made dilution device that was located at the  
133 chamber inlet.

134 The FORTH-ASC consists of 10 m<sup>3</sup> squared Teflon chamber, located inside a 30 m<sup>3</sup>  
135 reflective room (polished interior aluminium walls), which is temperature-regulated and  
136 equipped with ultraviolet lights (Osram, L 36W/73 UV lamps). This setup yields a maximum  
137 NO<sub>2</sub> photo-dissociation rate coefficient ( $J_{NO_2}$ ) of 0.5 min<sup>-1</sup> when all lights are on. In this study  
138 1/3 to 2/3 of the ultraviolet lights were used during photooxidation, resulting in a NO<sub>2</sub> photo-  
139 dissociation rate coefficient ( $J_{NO_2}$ ) of 0.17 to 0.33 min<sup>-1</sup>.

140 Eight day-to-night (denoted as DN) and eight night-to-day (denoted as ND) aging  
141 experiments were performed under dry (12-24% RH) conditions. Because a small amount of  
142 water vapor is inherently present in biomass burning emissions, achieving extremely low RH  
143 (<5%) would require complete removal of this water, which would result in losses of organic  
144 vapors and particles and compromise experimental quality. The selected RH range therefore  
145 ensured stable experimental conditions and is consistent with previous chamber studies  
146 (Kodros et al., 2022; Li et al., 2023). Table 1 summarizes the initial aerosol composition and  
147 experimental conditions for all the conducted experiments. To investigate the impact of fire  
148 starter on biomass burning emissions characteristics, pine kindling mixed with olive logs was  
149 used in two of the ND experiments (ND7, ND8). Pine, which is a softwood, has chemically  
150 distinct characteristics compared to olive wood (hardwood) and is used as a kindling material  
151 because it burns quickly due to its high resin content. While all night-to-day (ND) and day-to-night

152 (DN) experiments were conducted under the same general initial chamber conditions (temperature,  
153 relative humidity, and sampling protocol), there were the unavoidable in these chamber experiments  
154 differences in the initial aerosol and gas-phase composition, including the starting organic aerosol (OA)  
155 mass, black carbon (BC) content, and oxidant concentrations (e.g., O<sub>3</sub>, NO<sub>2</sub>) (Table 1).

156 The smog chamber was first flushed with clean air overnight at a rate of 20 L min<sup>-1</sup>.  
157 Approximately 30 min after the combustion ignition in the wood stove, when flaming  
158 conditions had been achieved, a fraction of the fresh emissions was diluted with clean air  
159 (dilution ratio ranging from 1:5 to 1:10) and was injected into the chamber, which was pre-  
160 filled with clean air and regulated to the desired RH level. This resulted in additional dilution  
161 (dilution ratio ~1:30) of the emissions. Two high precision mass flow controllers (Bronkhorst  
162 EL-FLOW Prestige FG-201CVP), operating at flow rates ranging from 0 to 20 L min<sup>-1</sup>, were  
163 used; one to supply clean air to the smog chamber during its filling and cleaning stages, and  
164 the other to supply clean air to the dilution system. The initial PM<sub>1</sub> concentration achieved in  
165 the chamber was 112 ± 56 µg m<sup>-3</sup> on average (Table 1). The fresh emissions were left to  
166 equilibrate and were characterized for about 2 hours. 30-90 ppb of d<sub>9</sub>-butanol (98%, Cambridge  
167 Isotope Laboratories) was also injected in the chamber as a tracer to determine the  
168 concentration of OH radicals (Barmet et al., 2012).

169 Subsequently, in DN experiments, the UV lights were turned on, initiating the daytime  
170 aging of fresh biomass burning emissions by OH radicals for at least 2 hours, without the  
171 addition of further oxidants. This oxidation step was then followed by at least 2 hours of aging  
172 with NO<sub>3</sub> radicals under dark conditions. To initiate NO<sub>3</sub> radical formation, NO<sub>2</sub> (50-150 ppb)  
173 was first injected into the chamber and allowed to mix for approximately 10 min, followed by  
174 a 1-min O<sub>3</sub> injection (60-280 ppb). This sequence ensured uniform NO<sub>2</sub> distribution and  
175 prevented its immediate consumption, enabling accurate concentration control. The injected  
176 NO<sub>2</sub> concentrations are consistent with polluted urban environments, while the O<sub>3</sub> levels  
177 correspond to those observed during daytime pollution episodes and in the residual layer above  
178 the nocturnal boundary layer, from where they can gradually mix downward and react with  
179 NO<sub>2</sub> to form NO<sub>3</sub> radicals (Kodros et al., 2020). Homogeneous mixing was confirmed through  
180 real-time gas monitoring, and measurements commenced only after full mixing to minimize  
181 artifacts. During ND cycling experiments, the same oxidation steps were performed but in  
182 reverse order. In DN experiments, “time zero” was defined as the moment when the UV lights  
183 were turned on, whereas in ND experiments, it was the point at which O<sub>3</sub> was injected.

184 Particle wall losses were also characterized for each experiment. After the completion of  
185 the two oxidation stages ammonium sulfate ((NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> ≥99 %, Sigma Aldrich) was injected

186 into the chamber and its loss rate was monitored for at least 3 hours. The dry seeds were  
187 produced by atomizing a  $(\text{NH}_4)_2\text{SO}_4$  solution using a TSI atomizer (model 3076) and drying  
188 the resulting droplets with a diffusion silica gel dryer (Fig. 1), as described in Wang et al.  
189 (2018). Particle wall loss corrections were applied to all aerosol data. Organic vapor wall losses  
190 were neglected over the corresponding experimental timescales. Wall losses of inorganic gases,  
191 such as  $\text{NO}_2$  and  $\text{O}_3$ , were evaluated during preliminary chamber characterization and found to  
192 be minimal, typically only a few percent or less (<5%) over the course of the experiments.

## 193 **2.2 Online instrumentation**

194 A suite of instrumentation was used for the online characterization of both particle and gas-  
195 phase pollutants (Fig. 1). A scanning mobility particle sizer (SMPS; Classifier) model 3080;  
196 DMA, model 3081, TSI) coupled to a butanol-based condensation particle counter (CPC,  
197 model 3775 high, TSI), was used for the measurement of the number and volume size  
198 distributions (mobility diameter in the range of 13–700 nm) of the aerosol particles. The SMPS  
199 sampled every 3 min with its sheath flow rate set at  $3 \text{ L min}^{-1}$  and the sample flow rate at  $0.6 \text{ L min}^{-1}$ . A high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne  
200 Research Inc.), working in V mode with vaporizer temperature set at  $600^\circ\text{C}$  and sampling flow  
201 rate of approximately  $0.1 \text{ L min}^{-1}$ , was used for monitoring the time evolution of the non-  
202 refractory organic and inorganic  $\text{PM}_1$  aerosol composition with time resolution of 3 min.  
203 Aerosol absorption and black carbon (BC) concentration were measured with a seven-  
204 wavelength aethalometer (Magee Scientific, Model AE33-7), sampling at  $2 \text{ L min}^{-1}$ . VOCs  
205 were measured using a proton transfer reaction mass spectrometer (PTR-QMS 500, Ionicon  
206 Analytik), sampling at  $0.5 \text{ L min}^{-1}$ . The drift tube was maintained at 2.3 mbar and operated at  
207 600 V. A detailed explanation of the PTR-MS operational parameters and the calibration  
208 procedure using VOC standards can be found in in Kaltsonoudis et al. (2016). Concentrations  
209 of carbon monoxide (CO) and dioxide ( $\text{CO}_2$ ), sulfur dioxide ( $\text{SO}_2$ ), ozone ( $\text{O}_3$ ) and nitrogen  
210 oxides ( $\text{NO}_x$ ) were measured using the corresponding monitors; CO (Teledyne model 300E),  
211  $\text{CO}_2$  (Teledyne model T360), (Thermo model 43i-TLE),  $\text{O}_3$  (Teledyne model 400E), NO and  
212  $\text{NO}_2$  (Teledyne model T201). The total sampling flow rate of all monitors was  $3.8 \text{ L min}^{-1}$ .

## 214 **2.3 Online data analysis methodology**

215 The initial combustion conditions in the chamber were characterized by calculating the  
216 modified combustion efficiency (MCE) as the ratio of the carbon dioxide ( $\text{CO}_2$ ) to the sum of  
217  $\text{CO}_2$  and carbon monoxide (CO) (Yokelson et al., 1996).

218 The HR-ToF-AMS data were analyzed using the packages SQUIRREL (Sequential Igor  
219 data Retrieval; v1.57) and PIKA (Peak Integration by Key Analysis; v1.16) incorporated in  
220 Igor Pro software (WaveMetrics; version 6.37). The method described in Canagaratna et al.  
221 (2015) was used to estimate of elemental O:C ratio. The AMS collection efficiency (CE) and  
222 the corresponding OA density have been determined using the algorithm proposed by  
223 Kostenidou et al. (2007). This approach combines the volume distributions obtained from the  
224 SMPS and the mass distributions of the main  $PM_1$  components from the AMS. The BC  
225 concentration obtained by the aethalometer was also included in the calculation, assuming a  
226 size distribution for BC similar to that of OA.

227 SMPS measurements were corrected using size-dependent wall loss rate constants,  
228 estimated by monitoring the decline in the mass concentration of  $(NH_4)_2SO_4$  particles injected  
229 into the chamber at the end of each experiment. Practically size independent first-order wall  
230 loss rates were observed for particle diameters ranging from 60 to 700 nm. Based on this, the  
231 concentrations of the non-refractory  $PM_1$  aerosol species measured by the AMS were corrected  
232 using one experiment-specific, size-independent wall loss rate constant that was  $0.15 \pm 0.05$   
233  $h^{-1}$  on average.

234 The total OA was split into primary (bbPOA) and secondary (bbSOA) following the  
235 approach proposed by Jorga et al. (2020) and applied for bbOA by Kodros et al. (2022). To  
236 quantify the variation between primary and secondary bbOA mass spectra, obtained by the  
237 AMS, the theta angle ( $\theta$ ) was estimated (Kostenidou et al., 2009). This angle represents the  
238 inner product of the two spectra (i.e., fresh and aged one), considered as n-dimensional vectors  
239 (n is to the number of the mass-to-charge ( $m/z$ ) ratios). Theta angles less than  $10^\circ$  imply high  
240 similarity, while major differences between two compared spectra correspond to  $\theta$  values  
241 higher than  $25^\circ$  (Florou et al., 2023). The approach described in Kiendler-Scharr et al. (2016)  
242 was used to quantify the particulate organic nitrate (ON). In the present study the minimum  
243 measured  $NO_2^+/NO^+$  ratio in all experiments was 0.04. The corresponding measured ratio for  
244 pure  $NH_4NO_3$ , determined through calibration, was equal to 0.56.

245 Prior to each experiment, background VOC levels in the chamber were measured using  
246 the PTR-MS for at least 1 hour. The PTR-MS was unavailable during experiments DN2–DN7.  
247 The PTR-MS measurements of the protonated VOCs were background-corrected and averaged  
248 at the end (over the last 1 h) of the fresh emissions' stabilization period, as well as at the end  
249 (over the last 1 h) of each oxidation step. The final values are summarized in Table S1 of the  
250 supplement, along with a classification of the identified VOCs by chemical structure and  
251 functional groups.

252 Following the work of Barmet et al. (2012), the average OH radical concentration was  
253 estimated from the decline/reduction in the concentration of the *m/z* 66 (protonated mass of d<sub>9</sub>-  
254 butanol). A d<sub>9</sub>-butanol reaction rate coefficient equal to  $3.4 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$  (at 295  
255 K) was assumed (Allani et al., 2021).

## 256 **2.4 Collection of samples for offline analysis**

257 To investigate the WS-OP of both fresh and aged BB aerosol, as well as to measure their  
258 organic (OC) and elemental carbon (EC) content, filter samples (Whatman Tissuquartz  
259 2500QAO-UP, 47 mm, 0.45 *pose* size) were collected for 1 h at the end of the emissions'  
260 equilibration period as well as at the end of each oxidation step. Prior to each experiment, blank  
261 filter samples were also collected. Sampling was conducted using a filter holder coupled with  
262 a PM<sub>2.5</sub> cyclone positioned at the chamber exit. An external vacuum pump (Becker VT 4.10,  
263 150 Mbar), operating at a flow rate of 16.7 L min<sup>-1</sup>, was used, with its exhaust connected to a  
264 HEPA filter (Whatman 6702-9500). Prior to sampling, the quartz filters were baked at 500 °C  
265 for 10 h and left in the oven overnight, to remove any absorbed organic material. Each filter  
266 was wrapped in prebaked aluminum foil and was kept before and after sampling in sterile  
267 polystyrene petri dishes (50 mm, Pall Laboratory). After sampling all filters were stored at a  
268 temperature of -20 °C, until WS-OP and OC/EC analysis.

269 Tenax sorbent tubes (stainless steel 3.5 x 1/4 in tubes, filled with Tenax TA, Markes  
270 International) were used to collect VOC samples at specific time intervals. The custom-made  
271 sampling system used included a mass flow controller (Alicat Scientific MC-500SCCM-  
272 D/5M), the sampling tube, and a diaphragm vacuum pump (AIRPO, Model D2028B 12VDC),  
273 operating at a flow of 0.3 L min<sup>-1</sup> for 1 to 1.5 h, resulting in total collected sample volumes  
274 ranging from 18-27 L. After sampling all sorbent tubes were capped with long-term storage  
275 brass caps containing PTFE ferrules and were stored in a freezer at -18 °C (Harshman et al.,  
276 2016).

## 277 **2.5 TD-GCMS measurements**

278 The offline determination of VOCs/IVOCs involved a two-step desorption process. The  
279 compounds adsorbed in the Tenax tubes were first desorbed using a thermal desorber (UNITY-  
280 Air Server-xr, Markes International Ltd.). During thermal desorption (TD), the sorbent tube  
281 underwent heating up to 280 °C for 10 min to release all its contents. Subsequently, the  
282 desorbed VOCs were captured using Helium (as the carrier gas) and then deposited onto a  
283 sorption cold trap at 20 °C. Subsequently, the temperature of the cold trap was gradually

increased from 20 °C to 300 °C at a rate of 100 °C s<sup>-1</sup>, where it remained for 6 min. The retained analytes were then injected into a single quadrupole gas chromatograph-mass spectrometer (GSMS, Shimadzu model QP2010, with helium as carrier gas). The GC-MS system was equipped with an inert capillary column (MEGA-5MS, 30 m length, 0.25 mm inner diameter, 0.25 µm film thickness). The oven temperature of the GC column remained at 32°C for approximately 5 min, increasing to 320 °C at 5 °C min<sup>-1</sup>. MS data acquisition was conducted in full scan mode, scanning within the *m/z* range of 35 to 300 amu. After the analysis, both the Tenax tubes and the GC column were cleaned. Calibration of the system was performed using standards of specific VOCs (EPA labelled) loaded in clean tubes. The species detected by TD-GCMS for a typical experiment (DN4) are presented in Table S2.

## 2.6 Oxidative potential (OP) measurements

The water-soluble oxidative potential (WS-OP) of redox-active aerosol components was measured using a DTT assay system (Fig. S1) at FORTH/ICE-HT in Patras, Greece, which is based on the semi-automated method of Fang et al. (2015). A detailed description of the system components, operation, measurement protocol, and data treatment, is provided in the Supplementary Information Section S1. Briefly, the fresh and aged aerosol samples (1.5 cm<sup>2</sup> punches of the collected quartz filters) are extracted, filtered, and incubated with DTT, in excess, under controlled conditions. The DTT is gradually oxidized by ROS in the sample, with its consumption rate (DTT activity, in nmol min<sup>-1</sup>) determined spectrophotometrically by measuring the absorbance of 2-nitrobenzoic acid (TNB), the derivatization product of DTT with DTNB reagent, at 412 nm at specific time intervals. The WS-OP was calculated by correcting for blank samples and was normalized to the OC mass of the sample, yielding net DTT consumption rates (mass-normalized DTT activity – DTT<sub>m</sub>) in pmol min<sup>-1</sup>µg<sup>-1</sup> (Table S3). OC was quantified via thermal-optical analysis (NIOSH-870 protocol), with an estimated relative standard deviation of 15 ± 5% for replicate measurements.

## 3 Results and Discussion

### 3.1 Characterization of fresh olive wood emissions

Flaming conditions predominated in all experiments, as indicated by the estimated modified combustion efficiency (MCE) that ranged from 0.91 to 0.99 (Table 1) (Li et al., 2015; Briggs et al., 2016). The initial PM<sub>1</sub> concentration of the fresh olive wood burning emissions in the chamber varied from 47 to 177 µg m<sup>-3</sup> (considering experiments DN1-DN8 and ND1-ND6).

315 This range of concentrations is representative of light to severe biomass burning pollution  
316 episodes in polluted urban areas during wintertime (Chen et al., 2022; Luo et al., 2022; Othman  
317 et al., 2022). The average AMS collection efficiency (CE) of the fresh emissions averaged 0.8  
318  $\pm$  0.2, while the mean OA density, calculated following the approach of Kostenidou et al.  
319 (2007), was  $1.11 \pm 0.12 \text{ g cm}^{-3}$ . Estimating the OA density from measured O:C and hydrogen-  
320 to-carbon (H:C) ratios, following the Kuwata et al. (2012) approach, yielded an average of  $1.18 \pm 0.03 \text{ g cm}^{-3}$ .

322 The fresh aerosol primarily consisted of organics ( $95 \pm 3 \%$ ) with OA concentrations  
323 ranging from 46 up to  $174 \mu\text{g m}^{-3}$  (Table 1). The rest of the aerosol consisted of BC ( $2.4 \pm 2.4\%$ ), nitrates ( $1.4 \pm 0.7\%$ ), sulfates ( $0.7 \pm 0.4\%$ ), chloride ( $0.4 \pm 0.2\%$ ) and ammonium ( $0.2 \pm 0.1\%$ ). In experiment DN3, ammonium sulfate seeds were also present explaining the higher  
326 initial sulfate (28%) and ammonium (10%) content.

327 The initial mass ratio of the organic aerosol to black carbon (OA/BC) ranged from 13 to  
328 263. The OA/BC differs significantly depending on the combustion conditions. When MCE  
329 values exceed 0.9, the OA/BC ratio can range between 0.3 to  $10^5$  (McClure et al., 2020), with  
330 higher values indicating more efficient combustion (Novakov et al., 2005). The relatively low  
331 primary BC concentrations in experiments DN3, DN5, DN6, and ND2, despite high MCEs of  
332 0.96 – 0.99, likely reflect variability in combustion conditions and fuel composition during our  
333 biomass burning experiments. Such variability can lead to differences in OA/BC ratios even  
334 under high MCE conditions. Overall, our OA/BC values indicate relatively efficient wood  
335 stove operation.

336 The average initial oxygen to carbon ratio (O:C) of the bbOA in all olive wood burning  
337 experiments was  $0.39 \pm 0.04$ . The average initial hydrogen to carbon ratio (H:C) was  $1.67 \pm 0.04$  ranging from 1.62 to 1.76. These values are consistent with previously reported field and  
338 smog chamber O:C and H:C observations for fresh biomass burning aerosols (Ng et al., 2010;  
339 Sun et al., 2016; Lim et al., 2019; Kodros et al., 2020; He et al., 2024). The relatively low AMS  
340  $f_{44}/f_{60}$  ratios ( $1.56 \pm 0.52$ ) observed in the experiments are representative of fresh biomass  
341 burning emissions from wildfires and laboratory wood burning chamber studies (Li et al.,  
342 2023).

344 The average high-resolution (HR) fresh bbOA mass spectrum obtained by AMS for the  
345 olive wood burning experiments (Fig. S2a) showed predominant fragments at  $m/z$  29 ( $\text{CHO}^+$ ,  
346  $\text{C}_2\text{H}_5^+$ ), 41 ( $\text{C}_2\text{HO}^+$ ,  $\text{C}_2\text{H}_3\text{N}^+$ ,  $\text{C}_3\text{H}_5^+$ ), 43 ( $\text{C}_2\text{H}_3\text{O}^+$ ,  $\text{C}_3\text{H}_7^+$ ), 55 ( $\text{C}_3\text{H}_3\text{O}^+$ ,  $\text{C}_4\text{H}_7^+$ ), 57 ( $\text{C}_3\text{H}_5\text{O}^+$ ),  
347 69 ( $\text{C}_5\text{H}_9^+$ ,  $\text{C}_4\text{H}_5\text{O}^+$ ) and 73 ( $\text{C}_3\text{H}_5\text{O}_2^+$ ), suggesting a significant presence of alkenes, alkanes,  
348 and fatty acids. The observed signals at  $m/z$  44 ( $\text{CO}_2^+$ ) and  $m/z$  60 ( $\text{C}_2\text{H}_4\text{O}_2^+$ ), are typical tracer

349 fragments for OOA and bbOA, respectively. The obtained fresh bbOA spectrum profile is quite  
350 similar to those reported in previous biomass burning chamber studies that examined wood or  
351 pellets burning (He et al., 2010; Kodros et al., 2020, 2022; Florou et al., 2023). The average  
352 theta angle  $\theta$  of the fresh bbOA spectra, calculated for all possible pairs of the olive wood  
353 burning experiments in the present study, was on average  $9^\circ \pm 7^\circ$  (Fig. S3), indicating a  
354 generally similar composition of fresh bbOA.

355 Based on PTR-MS measurements, oxygen-containing compounds contributed the largest  
356 portion of the protonated VOCs identified in the fresh emissions (Fig. 2a). Aldehydes,  
357 including acetaldehyde ( $m/z$  45;  $12.9 \pm 3.7$  ppb), formaldehyde ( $m/z$  31;  $1.6 \pm 0.7$  ppb), acrolein  
358 ( $m/z$  57;  $3.5 \pm 1.5$  ppb), and hexenal ( $m/z$  99;  $2.1 \pm 1.5$  ppb), along with saturated ketones like  
359 acetone ( $m/z$  59;  $4.7 \pm 2.0$  ppb) and unsaturated ones such as ethyl vinyl ketone ( $m/z$  85;  $2.1 \pm$   
360  $1.4$  ppb), contributed a total of 32.5 ppb, accounting for 19.7% of the measured VOCs.  
361 Carboxylic acids, such as formic ( $m/z$  47) and acetic ( $m/z$  61) acids, averaged a total  
362 concentration of 8.2 ppb, comprising 5% of the total VOCs (Fig. 2a). The main identified  
363 alcohol was 1-butanol ( $m/z$  75), which accounted for 3% of the VOC composition, with  
364 concentrations varying from 2.1 ppb to 9.5 ppb across experiments (Table S1). Furans and their  
365 derivatives ( $m/z$  69, 83, 113, 147) had an average concentration of 7.9 ppb, accounting for 5%  
366 of the total measured VOCs (Fig. 2a).

367 Cyclic and heterocyclic aromatic compounds (with 1-ring or 2-ring structure) contributed  
368 approximately 10% to the total VOCs. This includes benzene ( $m/z$  79;  $1.5 \pm 1.0$  ppb) and its  
369 substituted forms ( $m/z$  139, 151, 155; 3.3 ppb), toluene ( $m/z$  93;  $1.2 \pm 0.8$  ppb), phenol ( $m/z$  95;  
370  $2.1 \pm 1.7$  ppb) and its substituted forms ( $m/z$  121, 135, 149, 169; 2.4 ppb in total), and C8  
371 aromatics, including xylenes, ( $m/z$  107;  $3.2 \pm 2.4$  ppb). Other minor contributors, with varying  
372 concentrations across experiments, included terpenes and terpenoids ( $m/z$  81 and  $m/z$  137),  
373 averaging 2.9 ppb, and naphthalene ( $m/z$  129), averaging  $1.4 \pm 1.1$  ppb. The presence of these  
374 aromatic species is corroborated by the Tenax samples, along with compounds like  
375 benzonitrile, trimethoxy- benzene, methylindene and benzofurans. For a typical sample of fresh  
376 emissions, chromatographic analysis yielded a variety of phenolic species other than phenol,  
377 with functional groups including several alkyl groups (methyl-, dimethyl-,ethyl), but also with  
378 oxygenated functional groups (methoxy-, dimethoxy-) as presented in Table S2. Furans  
379 comprised approximately 11% of the identifiable VOCs in the offline analysis, with the most  
380 prominent being furfural, followed by methyl-furans and methyl- furancarboxaldehyde. In  
381 terms of polyaromatic species, similarly to the PTR-QMS observations, the most abundant was  
382 naphthalene, while there were several alkyl-substituted naphthalenes present in comparable

383 concentrations. Trace amounts of higher ring number PAHs (e.g., phenanthrene) were also  
384 observed. Most of these compounds have been previously reported in biomass burning ambient  
385 and laboratory studies (Stockwell et al., 2014; Bruns et al., 2017; Sun et al., 2019; Desservetaz  
386 et al., 2023; Florou et al., 2023).

387 The average WS-OP of the fresh olive wood burning aerosol was  $42.9 \pm 16.1 \text{ pmol}$   
388  $\text{min}^{-1} \mu\text{g}^{-1}$ , comparable to toxicity levels reported in literature for the water- and methanol-  
389 soluble portion of freshly emitted bbOA, which were also estimated using the acellular DTT  
390 assay protocol (Cao et al., 2021; Wang et al., 2023). The WS-OP values ranged from  $21.2 \pm$   
391  $5.7 \text{ pmol min}^{-1} \mu\text{g}^{-1}$  (in DN3) to  $79 \pm 11.3 \text{ pmol min}^{-1} \mu\text{g}^{-1}$  (in DN7) (Table S3).

### 392 3.2 Effect of pine kindling on fresh olive wood emissions

393 In experiments ND7 and ND8, where pine kindling sticks were mixed with olive wood logs,  
394 the  $\text{PM}_1$  concentration during the characterization period was  $126 \mu\text{g m}^{-3}$  and  $276 \mu\text{g m}^{-3}$ ,  
395 respectively (Table 1). High amounts of BC ( $67 \mu\text{g m}^{-3}$  and  $190 \mu\text{g m}^{-3}$ ) were produced in these  
396 experiments, constituting more than half (53% and 69%) of the total fresh  $\text{PM}_1$  mass. Given  
397 the efficient combustion conditions (MCE ranged from 0.96 to 0.98), these elevated BC levels  
398 were likely related to the properties of the pine (e.g., higher moisture, ash, and carbon content)  
399 compared to the olive logs (Nyström et al., 2017; Trubetskaya et al., 2021). The initial O:C of  
400 the fresh bbOA was 0.23 in experiment ND7 and 0.36 in ND8. The O:C in ND7 was the lowest  
401 of all experiments.

402 Comparison of the average fresh bbOA mass spectrum from olive-pine mixed emissions  
403 with that of olive logs burning (Fig. S4a) reveals significantly higher peaks at  $m/z$  28 ( $\text{CO}^+$ ;  
404 +69%), 41 (+36%), 44 (+40%), and 73 (+39%), indicating an increase in certain oxygenated  
405 organic species. Additionally, the stronger fractional signals at  $m/z$  91 ( $\text{C}_7\text{H}_7^+$ ; 104%), and at  
406 higher masses, such as  $m/z$  105 ( $\text{C}_8\text{H}_9^+$ ; 154%), 129 ( $\text{C}_{10}\text{H}_9^+$ ; +166%), suggest a higher relative  
407 contribution of cyclic hydrocarbons, PAHs, and other aromatic compounds. The theta angle of  
408 the two average fresh spectra was approximately  $20^\circ$ , implying distinct chemical composition  
409 of olive-pine mixed emissions. For the VOCs, while most aromatic compound concentrations  
410 were lower in the mixed fuel emissions, their relative contribution to the total VOCs was higher  
411 (17.7% vs. 9.6% in olive wood alone), suggesting differences in pyrolysis pathways and  
412 thermal degradation mechanisms between the two wood types (Fig. 2a,b). Additionally,  
413 monoterpenes ( $m/z$  137 and their fragment  $m/z$  81) showed a significant increase in the mixed  
414 emissions, rising from 2.9 ppb to 9.1 ppb, highlighting the influence of pine higher terpene  
415 content on VOC composition (Fig. 2). The variations observed in aldehydes, ketones, and

416 heavier PAHs were within the experimental uncertainty. A more detailed breakdown of the  
417 absolute and CO<sub>2</sub>-normalized VOC concentrations, including experiment-specific observations  
418 and comparisons, is provided in the Supplement (Fig. S5, Table S1).

419 No changes were observed in the WS-OP of the fresh olive-pine mixed emissions  
420 compared to fresh olive wood emissions. The corresponding DTT<sub>m</sub> values in experiments ND7  
421 and ND8 were  $44.7 \pm 4.0 \text{ pmol min}^{-1} \mu\text{g}^{-1}$  and  $41.1 \pm 3.4 \text{ pmol min}^{-1} \mu\text{g}^{-1}$ , respectively (Table  
422 S3). Similar WS-OP values (25 to 45 pmol min<sup>-1</sup> μg<sup>-1</sup>) were reported by Wang et al. (2023) for  
423 fresh bbOA from pine combustion under smoldering conditions (MCE=0.61). These values are  
424 comparable to the average WS-OP measured in this study for olive wood emissions ( $42.9 \pm$   
425  $16.1 \text{ pmol min}^{-1} \mu\text{g}^{-1}$ ).

### 426 3.3 Typical day-to-night (DN) aging experiment

427 During a typical dry DN oxidation experiment (DN1), two hours before the start of oxidation  
428 (at  $t = -2 \text{ h}$ ),  $70 \pm 0.4 \mu\text{g m}^{-3}$  of fresh olive wood burning PM<sub>1</sub> (91 % OA) were injected into  
429 the chamber along with approximately 14 ppb of O<sub>3</sub> (Fig. 3). During the emissions equilibration  
430 period (-2 to 0 h), the average O:C was 0.43, H:C was 1.67, OA/BC was 17, and the  $f_{44}/f_{60}$   
431 ratio was 1.37 (Table 1), and remained quite stable. The WS-OP of the fresh aerosol was  
432 estimated at  $51.4 \pm 4.7 \text{ pmol min}^{-1} \mu\text{g}^{-1}$  (Table S3).

433 At time zero ( $t = 0 \text{ h}$ ), daytime oxidation of the emissions was initiated by turning on the  
434 UV lights of the chamber, without adding any oxidants, and allowing the process to proceed  
435 for 2 h. Under the given experimental conditions, each hour of UV exposure in the simulation  
436 chamber corresponds to approximately 2 hours of atmospheric photochemical oxidation,  
437 assuming an average OH concentration of  $1.5 \times 10^6 \text{ molecule cm}^{-3}$  (Liu et al., 2018; Nault et al.,  
438 2018). In DN1, the average OH concentration during this 2-h oxidation period, estimated from  
439 the decay of d9-butanol, was  $3.2 \times 10^6 \text{ molecules cm}^{-3}$ , corresponding to an equivalent daytime  
440 exposure of 4.3 h. The average O<sub>3</sub> concentration was  $33 \pm 14 \text{ ppb}$ .

441 During this 2-h period the OA (wall loss corrected) increased by  $22 \mu\text{g m}^{-3}$  (34%).  
442 Organic nitrates also increased by 54% and O<sub>3</sub> reached 56 ppb. The H:C decreased by 4% while  
443 the  $f_{44}/f_{60}$  more than doubled (3 times higher). The O:C increased from 0.43 to 0.58 (35%),  
444 consistent with previous observations (Tiitta et al., 2016). The change in the HR-AMS  
445 spectrum of the day-aged OA was modest ( $\theta = 8^\circ$ ). The photochemical processing resulted in  
446 an 51% increase of WS-OP ( $77.6 \pm 6.3 \text{ pmol min}^{-1} \mu\text{g}^{-1}$ ) of the bbOA (Table S3). Similar  
447 increases of OP have also been reported in previous studies (Wong et al., 2019; Lei et al., 2023;  
448 Wang et al., 2023).

449 Furans, terpenes and cyclic aromatic hydrocarbons, major precursors of SOA production,  
450 were significantly reduced during daytime (Fig. S6). Aromatic hydrocarbons including toluene  
451 (*m/z* 93), phenol (*m/z* 95), styrene (*m/z* 105), C8 aromatics (*m/z* 107), C9 aromatics (*m/z* 121),  
452 and creosol/2-methoxy-4-methylphenol (*m/z* 139) reacted and their levels were reduced (Fig.  
453 S6b). Daytime aging also led to small changes (1 ppb or less) in the concentrations of  
454 formaldehyde, acetaldehyde, acetone, acetic acid, and heptanal which however could also be  
455 attributed to chamber background effects. According to TD-GCMS analysis, maleic anhydride  
456 was also identified at *m/z* 99 in the aged emissions (Table S2).

457 Reaction with OH radicals was estimated to be the dominant daytime oxidation pathway  
458 for most of the examined VOC species. For methyl vinyl ketone (*m/z* 71), benzene (*m/z* 79),  
459 monoterpenes fragment (*m/z* 81), methyl furan (*m/z* 83), toluene (*m/z* 93), phenol (*m/z* 95), and  
460 C8 aromatics (*m/z* 107, assuming o-xylene), the observed reductions in concentration were  
461 close to the theoretically expected values (Table S4). Lower than predicted reductions, due to  
462 OH oxidation, were observed for furan/isoprene (*m/z* 69; 32% less), ethyl vinyl ketone (*m/z* 85;  
463 21% less), styrene (*m/z* 105; ~30% less), C9 aromatics (*m/z* 121; assuming 1,2,3  
464 trimethylbenzene; 23% less), monoterpenes (*m/z* 137; assuming a-pinene; 58% less), and  
465 creosol (*m/z* 139; 60% less). This discrepancy from theoretical predictions is likely due to the  
466 presence of other compounds at the same *m/z* signal, including isomers, that react more slowly.  
467 Ozone-induced oxidation was a minor consumption mechanism for most of the VOCs ( $k_{O_3}$   
468 ranged from  $10^{-17}$  to  $10^{-22}$  molecule $^{-1}$  cm $^3$  s $^{-1}$ ) (Table S5), with the exception of monoterpenes  
469 and their fragments (*m/z* 137 and 81).

470 At the end of the daytime oxidation ( $t = 2$  h), the UV lights were turned off, and nighttime  
471 oxidation of the already aged emissions was conducted for two hours (2–4 h) by injecting  
472 additional 80 ppb of O<sub>3</sub> and 130 ppb of NO<sub>2</sub> into the chamber. The reaction of NO<sub>2</sub> and O<sub>3</sub>  
473 resulted in the decrease of their levels along with production of NO<sub>3</sub> radical (Fig. 3d). Although  
474 the NO<sub>3</sub> radical concentration was not directly measured in this study, it was estimated to range  
475 between 1 and  $5 \times 10^8$  molecule cm $^{-3}$  (typical for nighttime urban environments) based on  
476 previous dark aging experiments conducted with the same chamber setup under similar  
477 conditions (Kodros et al., 2022; Florou et al., 2023). This corresponds to approximately 4–7  
478 hours of equivalent atmospheric exposure.

479 Nighttime aging led to further bbSOA production, with OA increasing by 17%, reaching  
480 100  $\mu\text{g m}^{-3}$  (Fig. 3a). Organic nitrate increased by 0.62  $\mu\text{g m}^{-3}$  (72%) and total nitrate by 0.94  
481  $\mu\text{g m}^{-3}$  (53%) compared to their daytime levels. Nighttime enhancement of organic nitrate has

482 been also reported in other studies (Kodros et al., 2020, 2022; Florou et al., 2023). The O:C  
483 ratio slightly increased from 0.58 to 0.61 (5%). A small decrease (<1%) in H:C was observed,  
484 while the  $f_{44}/f_{60}$  increased further by 19% due to the nighttime oxidation. At the end of the DN  
485 oxidation cycle the theta angle of the aged aerosol compared to the fresh one was 23° (Fig. 3e),  
486 suggesting significant differences. The final DTT<sub>m</sub> of the aged emissions was  $68.7 \pm 6.0$  pmol  
487 min<sup>-1</sup>  $\mu\text{g}^{-1}$ , higher by 11% compared to the daytime measured WS-OP and 34% higher than  
488 the fresh one (Table S3).

489 The most notable VOC increases were observed for formaldehyde (*m/z* 31), which  
490 increased from 1.8 to 2.2 ppb (22%); hexenal/maleic anhydride (*m/z* 99), which increased from  
491 2.7 to 3.7 ppb (37%), and 2,3-benzofurandione (*m/z* 149), that increased from 0.2 to 0.3 ppb  
492 (50%) (Fig. S6). Further decreases in the gas-phase concentrations of furan (*m/z* 69) by 0.4 ppb  
493 (61%), methylfuran (*m/z* 95) by 0.35 ppb (26%), phenol (*m/z* 95) by 1.1 ppb (86%), and styrene  
494 (*m/z* 105) by 0.4 ppb (56%), were observed (Fig. S6).

#### 495 3.4 Typical night-to-day (ND) aging experiment

496 ND1 is as a typical night to day oxidation experiment (Fig. 4). The initial PM<sub>1</sub> concentration  
497 injected into the chamber was  $121 \mu\text{g m}^{-3}$ , with OA contributing 97%. The transition from fresh  
498 emissions to nighttime (0-2 h) and then daytime (2-4 h) oxidation resulted in significant  
499 changes in both the particle and gas phase. OA concentration increased by  $78 \mu\text{g m}^{-3}$  (65%  
500 increase) during the nighttime oxidation and by  $34 \mu\text{g m}^{-3}$  (an additional 17% increase) during  
501 daytime oxidation. During nighttime, total nitrate increased from 0.66 to  $5.4 \mu\text{g m}^{-3}$ , driven by  
502 production of organic nitrate. During daytime, organic nitrate levels decreased slightly (8%)  
503 compared to nighttime. The ND cycle also led to increases in ammonium levels first by  $0.9 \mu\text{g}$   
504  $\text{m}^{-3}$  (from 0.3 to  $1.2 \mu\text{g m}^{-3}$ ) and then by  $0.3 \mu\text{g m}^{-3}$  (from 1.2 to  $1.5 \mu\text{g m}^{-3}$ ).

505 The theta angle between the HR-AMS spectra of fresh and night-aged OA was 13° while  
506 at the end of the ND oxidation cycle the overall change of spectrum of the aged aerosol  
507 compared to the fresh one was 24° (Fig. 4e), similar to that observed during the DN cycle. The  
508  $f_{44}/f_{60}$  ratio increased from 1.7 to 5.5 during night and from 5.5 to 9 during the day, while the  
509 H:C decreased from 1.67 to 1.61 and 1.58, respectively (Tables 1 and 2). The O:C increased  
510 by 34% (from 0.41 to 0.55) at night with a further 20% enhancement (from 0.55 to 0.66)  
511 observed after the day aging. DTT<sub>m</sub> increased from  $31.8 \pm 2.8$  pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  for the fresh  
512 aerosol to  $42.5 \pm 3.1$  pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  for night-aged aerosol (33 % increase) and to  $71.0 \pm 5.7$   
513 pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  for day-aged aerosol (67% increase) (Table S3). Unlike experiment DN1,

514 which exhibited an initial increase (daytime) followed by a decrease (nighttime) in oxidative  
515 potential, experiment ND1 showed a monotonic increase with aging (Fig. 5).

516 During daytime oxidation, the OH concentration was  $4.2 \times 10^6$  molecule cm<sup>-3</sup>, which  
517 corresponds to approximately 5.3 h of equivalent photochemical atmospheric aging. O<sub>3</sub> levels  
518 increased by 88 ppb, rising from 96 ppb to 184 ppb by the end of daytime oxidation.

519 Similar trends and concentration levels were observed for most identified VOCs in  
520 experiments ND1 and DN1 (Fig. S6a and Fig. S7a). The observed differences in the percentage  
521 reduction of key bbSOA precursors, such as aromatic compounds and furans, between DN1  
522 and ND1 (Fig. S6b and Fig. S7b) suggest that the variability in precursor depletion dynamics  
523 is primarily influenced by differences in oxidant availability, photochemical reactivity, and the  
524 chemical composition of the emissions.

### 525 **3.5 Results of all dry DN and ND experiments**

526 The average OA production (including organic nitrate) observed in the DN (DN1-DN8) and  
527 ND (ND1-ND6) experiments at the end of a complete diurnal aging cycle was  $51 \pm 22 \mu\text{g m}^{-3}$ ,  
528 ranging from 19 to  $136 \mu\text{g m}^{-3}$  (Fig. 6a). These values correspond to a total OA mass increase  
529 ranging from 35% to 91% compared to the fresh OA. In both oxidation cycles the majority of  
530 the produced OA was formed during the first stage of oxidation. This is consistent with the  
531 higher availability of precursor VOCs initially. No significant differences were observed in the  
532 levels of OA produced in experiments ND7 and ND8 compared to the rest (Fig. 6a). This  
533 suggests that, although the use of pine kindling resulted in a different composition of fresh  
534 wood emissions, its overall impact on SOA production was less significant compared to the  
535 influence of oxidation conditions.

536 In some cases, the ND oxidation cycle resulted in higher SOA production (Fig. 6a and  
537 Fig. 7b). For instance, in experiment ND1, an OA mass increase of over 90% was observed at  
538 the end of the ND cycle. Similarly, in ND5, the OA increased by 78% compared to fresh. The  
539 enhanced SOA observed in ND1 and ND5 appears to be linked to their experimental  
540 conditions. Both experiments had high initial OA concentration in the fresh emissions (Table  
541 1) and higher initial O<sub>3</sub> levels (at around 30 ppb) compared to the rest of the ND experiments,  
542 which had an average level of  $14 \pm 2$  ppb. These conditions suggest that the enhanced SOA  
543 formation in ND1 and ND5 was driven by the higher availability of precursors, oxidants, and  
544 existing OA mass, which together facilitated higher secondary production. However, statistical  
545 analysis across the full dataset did not confirm that the ND oxidation cycle generally leads to  
546 higher SOA production compared to the DN cycle.

547 The experiments in this study were performed under relatively low RH conditions (12–  
548 24%) to minimize condensation and artifacts during sampling. While this approach enables  
549 comparison across oxidation scenarios, it does not capture the effects of higher RH and  
550 associated aerosol liquid water content, which are often significant during atmospheric  
551 nighttime aging. Aerosol water can enhance multiphase chemistry and SOA formation, and  
552 thus our results should be interpreted in the context of this limitation.

553 During both DN and ND oxidation cycles, the average density of the aged aerosol  
554 increased from 1.17 to 1.33 g cm<sup>-3</sup>, corresponding to approximately 13% increase in both cases  
555 (Table 2). Similar increases in bbSOA density, in the range of 1.31–1.34 g cm<sup>-3</sup>, have been also  
556 reported in other chamber studies during dark aging (Li et al., 2015; Florou et al., 2023).

557 At the end of the DN oxidation cycle, the average O:C was  $0.61 \pm 0.04$ , 56% higher than  
558 the average O:C ( $0.39 \pm 0.03$ ) of the fresh bbOA in our experiments. Almost 90% of this  
559 increase occurred during daytime (O:C increased from 0.39 to 0.59;  $\Delta$ O:C= 0.2) (Fig. 7c), while  
560 the subsequent nighttime oxidation resulted in an additional 10% increase in O:C (from 0.59  
561 to 0.61). For the ND cycle, the O:C increased from  $0.40 \pm 0.06$  for the fresh emissions to  $0.61$   
562  $\pm 0.06$  (a 54% increase) at the end of the cycle (Fig. 7d). In this case, the contributions of the  
563 nighttime and daytime oxidation stages to the increase in O:C were almost equal, at 55% and  
564 45%, respectively. In both oxidation cycles the final O:C is similar, but the importance of each  
565 oxidation stage depends on the order (oxidation sequence).

566 In all experiments, the OA AMS spectra changed progressively with aging. The  
567 predominant differences between the average fresh and aged bbSOA spectra at the end of DN  
568 cycle were found for *m/z* 28 (more than 2-fold increase) and 44 (1.5-fold increase) (Fig. S2).  
569 Significant decreases were observed for *m/z* 60 (37%) and 91 (36%), 115 (38%) and 137 (42%).  
570 The same changes were observed comparing the fresh and the ND aged bbSOA (Fig. S2).  
571 During the DN cycles the main changes in the OA spectrum occurred during the first (daytime)  
572 oxidation regime, with a theta angle of  $26 \pm 4^\circ$  on average (Fig. 7e). The further change in the  
573 second step (nighttime) was  $4 \pm 2^\circ$  on average. In contrast, a more balanced change was  
574 observed in the evolution of the theta angle over time during the ND cycle (Fig. 7f). The  
575 average OA spectrum shifted by  $19 \pm 4^\circ$  on average during nighttime, followed by an additional  
576  $10^\circ$  shift during UV exposure. Overall, at the end of both cycles, regardless of the followed  
577 oxidation path, the final average bbSOA spectra were almost identical ( $\theta < 3^\circ$ ) (Fig. 8).

578 To evaluate the environmental relevance of the chamber-produced bbSOA, the final DN  
579 and ND bbSOA spectra from this study were compared to the spectra of oxidized OA, that was  
580 measured at a remote site in Greece (Pertouli) during the summer of 2022 (Vasilakopoulou et

581 al., 2023). Most of this aged OA was aged emissions of wildfires from different regions of  
582 Europe. Two oxygenated OA (OOA) factors; a more-oxidized OOA (MO-OOA) and a less-  
583 oxidized OOA (LO-OOA) were needed to reproduce the observed OA spectra. Our final  
584 bbSOA spectra showed greater similarity to the LO-OOA factor, with a theta angle of  
585 approximately 16°, and were more distinct ( $\theta$  at around 30°) from the MO-OOA spectra  
586 measured in Pertouli. This suggests that our experiments simulated the earlier stages of  
587 atmospheric aging, while additional aging processes likely occur under ambient conditions (see  
588 also Fig. S8).

589 Changes in VOC levels of aged emissions across all DN (Fig. S9) and ND (Fig. S10)  
590 experiments were consistent with those observed in the typical experiments DN1 and ND1.  
591 Both aging cycles resulted in a significant decrease in the concentration of furans and their  
592 derivatives, cyclic and polycyclic aromatic hydrocarbons and terpenes. The day aged Tenax  
593 samples indicated a moderate decrease in aromatic species like toluene (~20%) and benzene,  
594 which is consistent with their lower reaction rates compared to higher carbon number  
595 aromatics. Rapid decrease in concentration was noted for species like phenol (~45%) and  
596 furfural (~75%), as well as their structurally related compounds. Related products, including  
597 2-nitro-phenol, 4-methyl-2-nitro-phenol, maleic anhydride, and 3-methyl-2,5-furandione, were  
598 also detected. p-Benzoquinone was also formed, possibly as a result of the reacted aromatics.  
599 Benzofuran was absent from the aged samples; instead, 2,3-benzofurandione was detected. At  
600 the same time a progressive increase in aldehydes and ketones was observed, along with  
601 significant increases in carboxylic acids, such as formic ( $m/z$  47) and acetic ( $m/z$  61). The  
602 benzaldehyde concentration increased, accompanied by the formation of benzeneacetaldehyde,  
603 2-hydroxy-benzaldehyde, 3-ethyl-benzaldehyde. A notable increase in butanol was also  
604 observed in the Tenax samples, along with the formation of straight-chain aldehydes (hexanal  
605 to undecanal). The GC-MS measurements for the night-aged samples following daytime  
606 processing were consistent with those of the PTR-MS. Furfural was no longer detected, while  
607 a further decrease in phenol and increases in benzaldehyde and butanol were noted. A  
608 cumulative depiction of the experiment's progression in terms of oxidation and VOCs detected  
609 by the GC-MS, is provided in Fig. S11. Similar results were obtained for the other experiments.

### 610 3.6 Effect of DN and ND oxidation cycle on WS-OP

611 The water-soluble oxidative potential (WS-OP) of fresh emissions ranged from 21 pmol min<sup>-1</sup>  
612  $\mu\text{g}^{-1}$  to 79 pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  and that of aged wood-burning emissions from 39 pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$   
613 to 127 pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  (Table S3). These values fall within the range reported in literature for

614 fresh bbOA and aged bbOA (Verma et al., 2015; Tuet et al., 2017; Bates et al., 2019;  
615 Daellenbach et al., 2020; Wang et al., 2023).

616 Figure 6b presents the absolute ( $\Delta DT_{m}$ , pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$ ) and percent changes in  $DT_{m}$   
617 activity for each experiment. In all DN experiments, WS-OP followed a consistent pattern of  
618 increasing during daytime oxidation and decreasing during subsequent nighttime aging, except  
619 for DN6 and DN8, where it continued to rise, suggesting experiment-specific chemical  
620 variability. Similarly, in the ND experiments, WS-OP generally increased during both  
621 nighttime and daytime oxidation, except for ND4 and ND7, which exhibited a reduction during  
622 the second daytime step, indicating that the extent of oxidative enhancement can depend on the  
623 initial emission composition and oxidation conditions.

624 The evolution of average WS-OP of fresh and aged emissions, considering all  
625 experiments and both oxidation cycles, (Fig. 9) was similar to that observed in experiments  
626 DN1 and DN1 (Fig. 5). The average WS-OP values for the DN cycle were  $47.9 \pm 17.7$  pmol  
627 min<sup>-1</sup>  $\mu\text{g}^{-1}$  for fresh emissions,  $93 \pm 27$  pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  for daytime-aged emissions,  
628 representing a 94% increase compared to fresh aerosol, and  $73.4 \pm 13.3$  pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  for  
629 nighttime-aged emissions, indicating a 21% reduction compared to daytime-aged WS-OP (Fig.  
630 9a). For the ND oxidation cycle, the average WS-OP of the fresh emissions were  $37.8 \pm 10.6$   
631 pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$ . After nighttime aging, it increased by 44% to  $54.4 \pm 13.6$  pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$ , and  
632 following daytime aging, it further increased by  $62.9 \pm 20.4$  pmol min<sup>-1</sup>  $\mu\text{g}^{-1}$  (Fig. 9b).

633 Statistical analysis (t-test) showed that aged WS-OP values were significantly higher than  
634 those of fresh emissions in all experiments, for both oxidation cycles. Additionally, for the DN  
635 oxidation cycle, a statistically significant difference was observed between the WS-OP of  
636 nighttime-aged (NO<sub>3</sub>-oxidized) and daytime-aged (UV-oxidized) emissions. Further details on  
637 the statistical analysis are provided in Supplementary Section S4 and Table S9.

638 The overall increase in WS-OP at the end of the two oxidation cycles was  $53 \pm 34\%$  for  
639 the DN cycle and  $66 \pm 8\%$  for the ND cycle, indicating that both daytime and nighttime aging  
640 of biomass burning emissions consistently enhanced their oxidative potential. Our results  
641 suggest that the sequence of chemical processes – whether the emissions are first oxidized by  
642 OH or NO<sub>3</sub> – can significantly affect the temporal evolution of OP. This, in turn, may also  
643 influence the health impacts associated with exposure to biomass burning plumes, depending  
644 on the time of day when the emissions occur. Although daytime boundary layer dynamics  
645 generally favour mixing and dilution of pollutants, daytime burning in urban environments may  
646 actually be as or more aggravating than nighttime burning, owing to the enhanced oxidative  
647 processing of the emissions occurring in the former stage of the diurnal cycle.

648 The correlation of WS-OP with produced OA and degree of oxidation (O:C) were also  
649 investigated. Three OA types were considered (fresh, day-aged, and night-aged). WS-OP was  
650 not well correlated with either the O:C ratio ( $R^2 < 30\%$ ) of the organic aerosol or its fresh and  
651 aged fractions ( $R^2$  up to 34%) (Fig. S12). This implies that the link between bbOA aging, and  
652 WS-OP change is complex and cannot be just described by one variable. Weak correlations  
653 ( $R^2 < 0.3$ ) between WS-OP (both fresh and aged) and most of the rest of the aerosol components  
654 (Tables 1 and 2) were also observed (Figs. S13–S17). An exception was the nighttime WS-OP,  
655 which showed a stronger correlation with inorganic nitrate ( $R^2 > 0.3$ ) in both oxidation cycles.  
656 In addition, after one complete diurnal cycle, WS-OP showed a stronger correlation with OC,  
657 with  $R^2$  values reaching up to 0.65 (DN cycle) and 0.53 (ND cycle) (Figs. S15 and S17),  
658 highlighting the growing influence of organic components. These results point to a complex  
659 interplay of chemical processes governing WS-OP variability.

660 It is important to note that our study focused exclusively on the water-soluble fraction of  
661 OP and thus does not capture contributions from relatively water insoluble components (e.g.,  
662 brown carbon and metals), which can significantly influence total oxidative properties (Gao et  
663 al., 2020b). Atmospheric aging usually tends to increase OA solubility through oxidative  
664 functionalization, which may move compounds with high OP to the aqueous phase during the  
665 extraction. Previous field observations support this hypothesis. Wong et al. (2019) reported  
666 that the OP of bbOA increased during the first day of atmospheric transport, even as OC and  
667 water soluble organic carbon (WSOC) concentrations decreased, indicating that aging removes  
668 less reactive compounds while producing more DTT-active compounds. The majority of this  
669 activity (around 80%) was associated with the water-soluble fraction, highlighting its dominant  
670 role in driving the oxidative properties of aged bbOA. These findings emphasize the crucial  
671 role of water-soluble aerosol components in driving OP, while also underscoring the need for  
672 studies that simultaneously capture both soluble and insoluble contributions.

673 The observed WS-OP trends could be also linked to the VOC composition and oxidation  
674 processes in the DN and ND cycles. The WS-OP of fresh emissions exhibited positive  
675 correlations ( $0.2 < R^2 < 0.6$ ) with several VOCs, including aldehydes ( $m/z$  45, 57), benzene  
676 ( $m/z$  79), monoterpenes ( $m/z$  81), toluene ( $m/z$  93), phenol ( $m/z$  95) and its substituted forms  
677 ( $m/z$  121, 135), styrene ( $m/z$  105), xylenes ( $m/z$  107), and naphthalene ( $m/z$  129) (Fig. S13).  
678 Among these, the strongest correlations were observed for benzene ( $R^2 = 0.56$ ), phenol ( $R^2 =$   
679  $0.56$ ), and styrene ( $R^2 = 0.56$ ), suggesting that the products of aromatic hydrocarbons and  
680 phenolic compounds may contribute significantly to the WS-OP of fresh bbOA. The DN cycle  
681 exhibits a high daytime WS-OP due to the OH oxidation of VOCs such as furans, aromatics,

682 and phenolic compounds, leading to the formation of reactive species like, 4-methyl-2-  
683 nitrophenol, and highly reactive p-benzoquinone. In contrast, the ND cycle shows a gradual  
684 increase in WS-OP, with the exception of experiments ND4 and ND7 (Fig. 6b). These  
685 contrasting trends likely reflect differences in the subsequent aging of oxidation products. In  
686 the DN case, the decrease in intrinsic DTT activity during nighttime oxidation may result from  
687 the formation of compounds with lower OP (less redox-active), lower solubility, or both, when  
688 daytime OH oxidation products are further processed by  $\text{NO}_3$  radicals. In some specific cases  
689 (DN6, DN8; Fig. 6b), WS-OP continued to increase at night, suggesting potential ongoing  
690 nighttime formation of soluble or redox-active compounds under certain oxidation conditions.  
691 Conversely, in the ND case, the reverse sequence of reactions appears to generate products  
692 with higher OP or solubility or both. Importantly, this divergence does not seem to be  
693 controlled by the bulk O:C ratio, but rather by differences in chemical structure of the later-  
694 generation products. For example, after nighttime oxidation in the ND cycle the WS-OP was  
695 positively correlated with formic acid ( $m/z$  47;  $R^2 = 0.24$ ), toluene ( $m/z$  93;  $R^2 = 0.28$ ),  
696 naphthalene ( $m/z$  129;  $R^2 = 0.25$ ) and substituted forms of phenol ( $m/z$  121, 135;  $R^2 = 0.25$ ) and  
697 benzene ( $m/z$  151, 155;  $R^2$  up to 0.2) (Fig. S18). After further daytime oxidation WS-OP was  
698 only correlated with  $m/z$  121 (trimethylbenzene;  $R^2 = 0.20$ ) (Fig. S19). Oxidized derivatives of  
699 phenol and benzene formed during atmospheric aging tend to exhibit higher water solubility  
700 than their parent compounds, as the functionalization increases molecular polarity and  
701 hydrogen-bonding capacity. More detailed work is needed to identify these later generation  
702 products and to quantify both their OP and water solubility.

703 After one complete diurnal cycle, WS-OP values in both cycles converge, indicating that  
704 oxidative processes in both pathways ultimately lead to similar levels of oxidation products.  
705 This convergence highlights the role of both fast and slow oxidation mechanisms in  
706 determining aerosol OP and suggests that even VOCs with lower reactivity can significantly  
707 contribute to aerosol toxicity over extended atmospheric aging.

## 708 4 Conclusions

709 This study investigated how different diurnal oxidation sequences - daytime-first (DN)  
710 and nighttime-first (ND)- affect the formation of OA, the gas-phase composition, and the  
711 oxidative potential of emissions produced by burning olive wood and olive wood mixed with  
712 pine throughout a complete diurnal aging cycle. Both DN and ND oxidation cycles resulted in  
713 enhancement in OA levels by 35%-90%, demonstrating efficient SOA production under both  
714 OH- and  $\text{NO}_3$ -dominated oxidation conditions. The mixture of olive wood with pine kindling

715 resulted in a different composition of fresh emissions, however its overall impact on SOA  
716 production was less significant compared to the influence of oxidation conditions.

717 The DN cycle favoured rapid daytime oxidation, producing highly oxygenated species  
718 and increasing the O:C ratio of the fresh emissions from  $0.39 \pm 0.04$  to  $0.59 \pm 0.04$  during  
719 daytime, reaching finally at  $0.61 \pm 0.03$  during nighttime. The ND cycle showed a gradual  
720 (two-steps) oxidation increase with a similar final O:C ratio of  $0.61 \pm 0.06$ . DN cycle exhibited  
721 rapid spectral changes during daytime oxidation, while ND cycles showed a more balanced  
722 two-step evolution. The convergence of O:C ratios after one diurnal cycle imply that multiple  
723 oxidation pathways can yield comparable end products through distinct chemical routes. At the  
724 end of both cycles, the final average bbSOA spectra were nearly identical ( $R^2 > 0.99$ ;  $\theta < 3^\circ$ ),  
725 indicating that the aerosol was transformed into similar aged OA regardless of the initial  
726 oxidation step (daytime or nighttime) at the start of the cycle. The chamber-produced bbSOA  
727 resembled the less-oxidized OOA in a field campaign in Greece, with the corresponding OA  
728 dominated by aged bbOA, suggesting that the present study has addressed only part of the  
729 aging that occurs in the atmosphere.

730 Both the DN and ND oxidation cycles effectively reduced the concentration of bbSOA  
731 precursors (e.g., furans, aromatic hydrocarbons, terpenes). Concurrently, a progressive increase  
732 in aldehydes and ketones was observed in both cycles, alongside increases in carboxylic acids,  
733 such as formic and acetic acids. The higher daytime WS-OP in the DN cycle likely results from  
734 rapid OH oxidation of reactive VOCs (e.g., furans, aromatics, phenolics), producing redox-  
735 active species such as nitrophenols and quinones. The subsequent nighttime decrease in DTT  
736 activity suggests formation of less soluble, less redox-active products. In contrast, the ND cycle  
737 accumulated moderately oxidized intermediates during nighttime  $\text{NO}_3^-$  oxidation, which were  
738 further transformed into more soluble, higher-OP compounds during daytime processing. Thus,  
739 variations in molecular structure and solubility, rather than bulk oxidation state (O:C),  
740 primarily drive differences in WS-OP.

741 From an atmospheric perspective, these findings imply that emissions released at  
742 different times of day may contribute differently to local air toxicity, depending on oxidant  
743 availability and photochemical activity. For example, daytime burning in urban or suburban  
744 environments may enhance the formation of ROS-generating compounds more rapidly than  
745 nighttime burning, even if total SOA mass is comparable. This temporal dependence suggests  
746 that biomass burning – related health impacts effects may vary with exposure timing, a factor  
747 rarely considered in air quality or exposure models. The DN cycle resulted in  $53 \pm 34\%$  increase  
748 in WS-OP of aerosol while the ND cycle showed a slightly higher increase of  $66 \pm 8\%$ . The

749 final WS-OP values of the DN ( $73 \pm 14 \text{ pmol min}^{-1} \mu\text{g}^{-1}$ ) and ND ( $63 \pm 20 \text{ pmol min}^{-1} \mu\text{g}^{-1}$ )  
750 cycles were statistically similar. This convergence suggests a self-organizing tendency in  
751 atmospheric aging toward similar chemical and toxicological endpoints.

752 Although this study focused on water-soluble OP, insoluble components such as  
753 transition metals and brown carbon (not assessed here) can also contribute significantly to total  
754 aerosol reactivity. Oxidative functionalization during atmospheric aging generally increases  
755 OA solubility, which may shift OP from the insoluble to the soluble phase, either altering or  
756 maintaining total OP, depending on chemical composition. Consequently, measuring only the  
757 water-soluble fraction likely underestimates the total oxidative capacity of aged bbOA. Future  
758 studies combining complementary assays (e.g., ascorbic acid or AA and glutathione or GSH)  
759 and phase-resolved analyses are necessary to better constrain the contributions of soluble and  
760 insoluble components to overall aerosol toxicity.

761 A few additional limitations of the present work must be also acknowledged. The  
762 experiments were performed at relatively low relative humidity (12–24%) to reduce  
763 condensation artifacts, and therefore the potential role of multiphase chemistry enhancement  
764 and promotion of SOA formation was not investigated. Furthermore, the aging period  
765 examined here corresponds approximately to a single diurnal cycle, representing only the first  
766 stages of atmospheric processing, whereas ambient bbOA typically undergoes multi-day  
767 oxidation cycles that may further modify its composition and OP. In addition, the controlled  
768 combustion setup, while suitable and effective for mechanistic interpretation, inevitably  
769 simplifies real-world burning conditions, where variations in fuel moisture, temperature  
770 gradients, meteorology, and interactions with co-emitted anthropogenic pollutants play  
771 important roles in shaping emissions chemistry.

772 Despite these constraints, the present study provides a mechanistic framework for  
773 understanding how diurnal oxidation sequences regulate the chemical and redox activity  
774 evolution of bbOA. The findings underscore the importance of considering oxidation  
775 sequences and timing when assessing the environmental fate and health impacts of biomass  
776 burning emissions. They highlight the complex and dynamic nature of atmospheric aging  
777 processes and emphasize the need for time-resolved approaches to better predict the  
778 transformation and OP evolution of bbOA under realistic atmospheric conditions.

## 779 **Author contributions**

780 M.P.G., K.F., and A.M. contributed to investigation, conducted the experiments and performed  
781 the laboratory measurements.; M.P.G. and G.S. performed the offline measurement of the

782 water-soluble oxidative potential of the collected aerosol samples.; A.M. performed the offline  
783 TD-GCMS analysis of the Tenax samples; C.K. contributed to chamber set-up optimization.;  
784 A.N. conceived and supported the research project; S.N.P. supported and directed this  
785 research.; M.P.G. and K.F. interpreted the results and contributed to formal data analysis;  
786 M.P.G. wrote the original manuscript with contributions from all co-authors.; All authors  
787 contributed to the review and editing of the manuscript and have approved the final submitted  
788 version.

## 789 **Conflicts of interest**

790 The authors declare that there are no conflicts to declare.

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1189

1190 **Table 1:** Initial biomass burning aerosol composition and initial chamber conditions for all the  
 1191 conducted experiments.

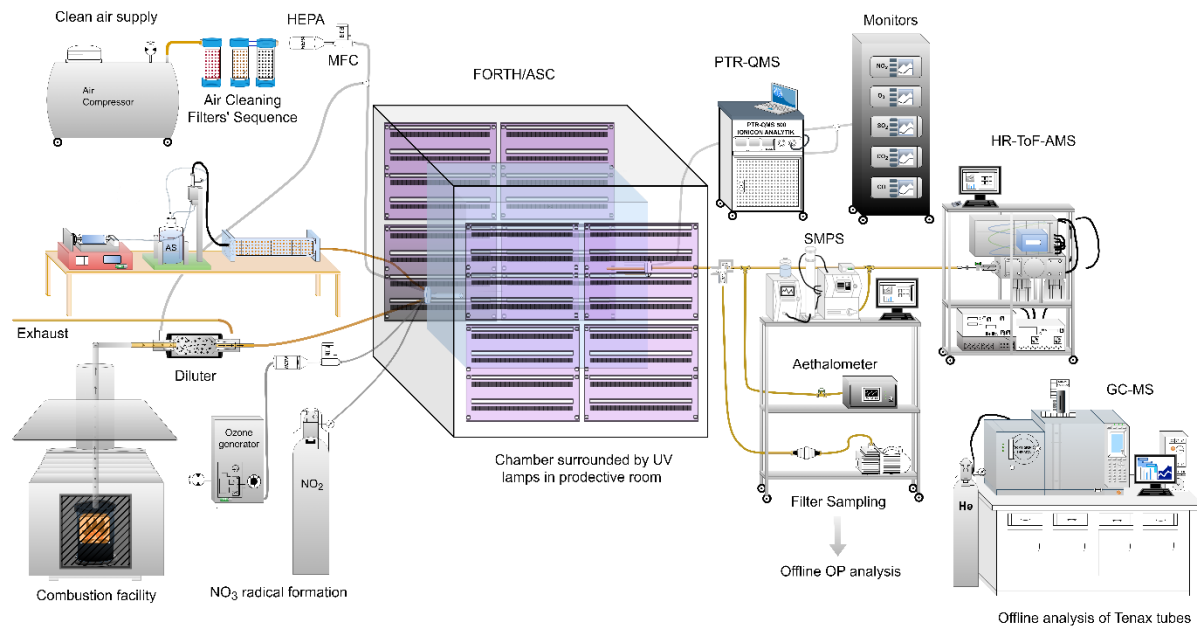
Exp.	PM <sub>1</sub> [ $\mu\text{g m}^{-3}$ ]	BC [ $\mu\text{g m}^{-3}$ ]	Amm onium [ $\mu\text{g m}^{-3}$ ]	Sulfat e [ $\mu\text{g m}^{-3}$ ]	Orga nics [ $\mu\text{g m}^{-3}$ ]	Nitrat e [ $\mu\text{g m}^{-3}$ ]	Org. Nitrat e [ $\mu\text{g m}^{-3}$ ]	Inorg. Nitrat e [ $\mu\text{g m}^{-3}$ ]	Chlor ide [ $\mu\text{g m}^{-3}$ ]	$f_{44}/f_{60}$	O:C	H:C	T <sub>init</sub> °C	RH <sub>init</sub> [%]	MCE*
DN1	70	3.7	0.06	0.62	63.6	1.18	0.56	0.62	0.45	1.37	0.43	1.67	16.4	13	0.96
DN2	114	7.9	0.13	0.86	102	2.01	1	1.01	0.42	2.21	0.39	1.66	N/A	N/A	0.92
DN3	79.1	N/A	8.1	22.1	48.2	0.52	0.23	0.29	0.11	1.82	0.38	1.62	N/A	N/A	0.99
DN4	177	1.1	0.24	1.11	172	2.02	0.91	1.11	0.62	1.11	0.39	1.71	N/A	N/A	0.91
DN5	102	0.4	0.09	0.47	99.8	0.65	0.26	0.39	0.19	1.00	0.35	1.72	N/A	N/A	0.99
DN6	53.6	0.9	0.07	1.02	49.6	1.66	1.02	0.64	0.41	1.50	0.36	1.76	N/A	N/A	0.96
DN7	74.8	0.8	0.06	0.54	72.1	1.15	0.43	0.72	0.16	1.34	0.41	1.67	N/A	N/A	0.91
DN8	85.5	5.4	0.09	0.49	78.0	1.22	0.28	0.94	0.33	1.02	0.44	1.65	16.8	13	0.92
ND1	121	N/A	0.28	1.01	118	1.21	0.66	0.55	0.58	1.72	0.41	1.65	16.5	13	0.92
ND2	72.0	0.5	0.21	0.45	69.4	1.13	0.76	0.37	0.24	2.53	0.47	1.67	N/A	N/A	0.96
ND3	47.2	0.4	0.16	0.12	45.8	0.67	0.34	0.33	0.05	2.37	0.29	1.67	16.5	14	0.94
ND4	93.3	1	0.22	0.54	90.2	0.99	0.50	0.49	0.34	1.61	0.40	1.66	15.9	15	0.92
ND5	176	N/A	0.25	0.34	174	1.35	0.62	0.73	0.18	1.27	0.37	1.66	16.6	13	0.91
ND6	124	3	0.18	0.41	120	0.72	0.31	0.41	0.38	0.92	0.43	1.65	16.7	14	0.90
ND7	126	67	0.06	0.07	58.7	0.15	0.10	0.05	0.05	2.47	0.23	1.61	17.1	12	0.98
ND8	276	190	0.12	0.62	83.4	1.22	0.85	0.37	0.25	1.88	0.36	1.65	17	24	0.96

1192 \*Modified combustion efficiency (MCE) calculated based on equation:  $([\Delta\text{CO}_2]/([\Delta\text{CO}] + [\Delta\text{CO}_2]))$ .

1193 **Table 2:** Composition of aged biomass burning aerosol, averaged over the last 30 minutes of  
 1194 each oxidation state, for both DN and ND experiments.

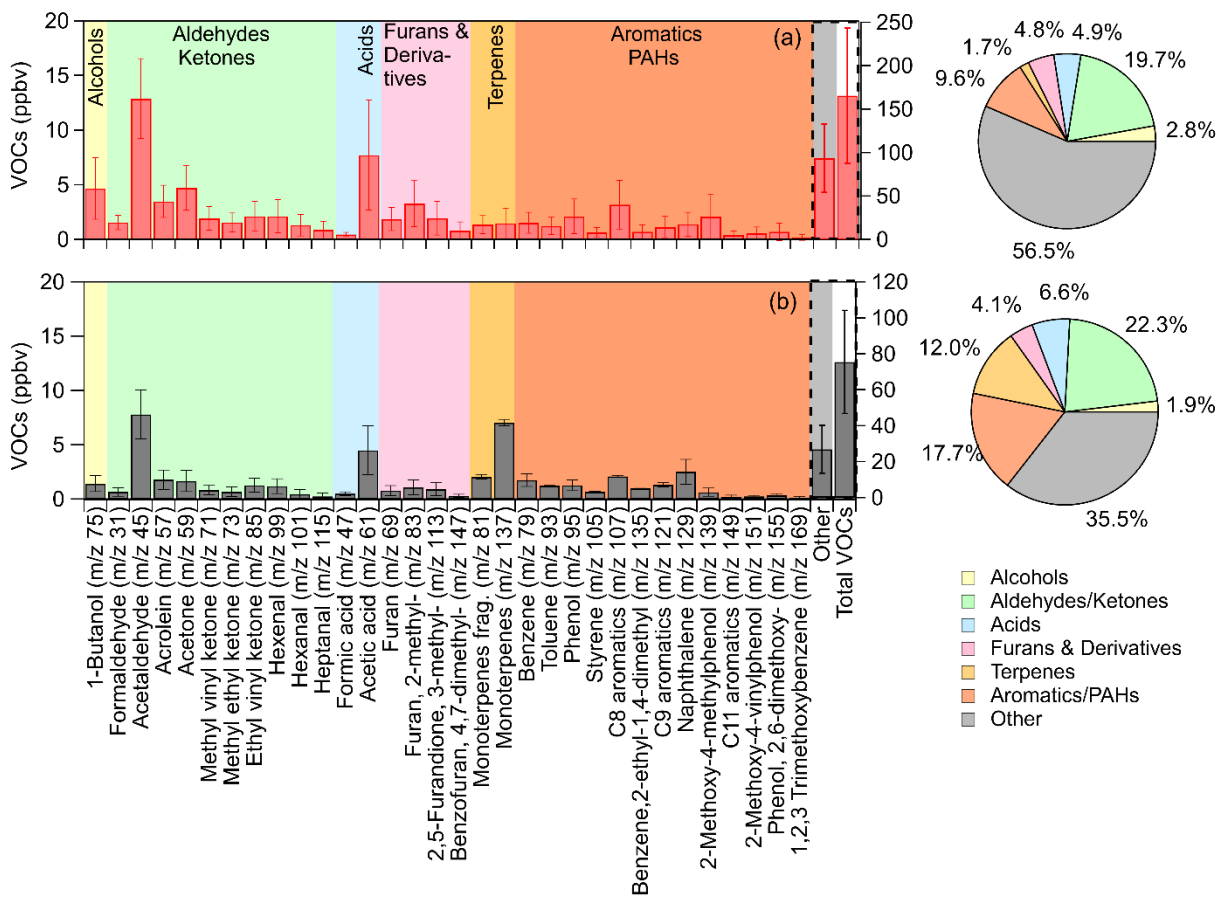
Exp.	Oxid.	PM <sub>1</sub> [ $\mu\text{g m}^{-3}$ ]	Amm onium [ $\mu\text{g m}^{-3}$ ]	Sulfate [ $\mu\text{g m}^{-3}$ ]	Organic [ $\mu\text{g m}^{-3}$ ]	Nitrate [ $\mu\text{g m}^{-3}$ ]	Org. Nitrate [ $\mu\text{g m}^{-3}$ ]	Inorg. Nitrate [ $\mu\text{g m}^{-3}$ ]	Chlor ide [ $\mu\text{g m}^{-3}$ ]	$\rho^*$ [g $\text{cm}^{-3}$ ]	$f_{44}/f_{60}$	O:C	H:C
<b>Day - Night</b>													
DN1	Day	88	0.1	0.7	85	1.76	0.86	0.90	0.38	1.34	4.3	0.58	1.61
	Night	104	0.15	0.83	100	2.74	1.48	1.26	0.4		5.1	0.61	1.6
DN2	Day	157	0.1	0.9	154	2.42	1.75	0.67	0.26	1.34	15.3	0.64	1.52
	Night	185	0.47	1.46	179	4.21	2.98	1.23	0.3		16.5	0.67	1.53
DN3	Day	102	10.3	28.9	62	0.63	0.36	0.27	0.13	1.37	8.5	0.63	1.55
	Night	107	10.8	30.3	65	0.94	0.61	0.33	0.15		9.7	0.64	1.55
DN4	Day	240	1.1	1.2	232	4.92	2.29	2.63	0.49	1.32	3.7	0.58	1.64
	Night	262	1.26	1.35	252	6.14	2.82	3.32	0.52		3.9	0.6	1.64
DN5	Day	134	0.1	0.5	132	0.97	0.52	0.45	0.18	1.29	3.4	0.52	1.66
	Night	156	0.19	0.59	153	2.19	1.51	0.68	0.2		4.3	0.56	1.65
DN6	Day	83	0.2	1.3	78	2.80	1.55	1.25	0.38	1.33	6.5	0.59	1.63
	Night	90	0.22	1.48	85	3.27	1.78	1.49	0.4		7.1	0.61	1.61
DN7	Day	105	0.1	0.6	103	1.43	0.69	0.74	0.15	1.34	5.4	0.6	1.61
	Night	122	0.17	0.79	119	1.94	1.03	0.91	0.17		5.8	0.62	1.61
DN8	Day	111	0.1	0.6	108	1.68	0.59	1.09	0.33	1.33	3	0.58	1.61
	Night	134	0.19	0.74	130	2.95	1.37	1.58	0.36		3.4	0.6	1.6
<b>Night – Day</b>													
ND1	Night	199	1.2	1.1	188	9.0	5.4	3.6	0.28	1.36	5.5	0.55	1.61
	Day	233	1.5	1.3	222	8.7	5.0	3.7	0.29		9	0.66	1.58
ND2	Night	110	0.5	0.5	102	7.0	5.5	1.5	0.15	1.39	8.1	0.62	1.6
	Day	119	0.6	0.6	112	6.3	4.8	1.5	0.16		11.1	0.7	1.57
ND3	Night	61	0.2	0.2	57	2.9	2.2	0.7	0.06	1.28	7.3	0.42	1.64
	Day	66	0.3	0.2	63	2.5	1.9	0.6	0.07		12.2	0.52	1.61
ND4	Night	133	0.4	0.8	127	4.3	3.1	1.2	0.29	1.31	3.9	0.49	1.62
	Day	148	0.6	1	142	4.2	2.9	1.3	0.3		6.4	0.58	1.6
ND5	Night	278	0.5	0.5	270	7.0	4.8	2.2	0.14	1.31	3.8	0.48	1.63
	Day	315	0.7	0.6	307	6.6	4.0	2.6	0.14		6.5	0.58	1.61
ND6	Night	192	0.7	0.6	184	6.8	4.1	2.7	0.18	1.33	2.6	0.51	1.62
	Day	203	0.8	0.7	195	6.0	3.4	2.6	0.19		4.8	0.61	1.61
ND7	Night	182	0.1	0.1	93	4.6	3.1	1.5	0.06	1.2	5	0.36	1.65
	Day	211	0.1	0.2	105	4.7	3.0	1.7	0.08		6.5	0.41	1.62
ND8	Night	400	0.3	1.2	149	9.3	5.4	3.9	0.26	1.29	4.7	0.48	1.64
	Day	451	0.5	1.7	163	9.1	5.2	3.9	0.29		7.6	0.54	1.6

1195 \*Density calculated based on O:C and H:C ratios, following the approach of Kuwata et al. (2012).



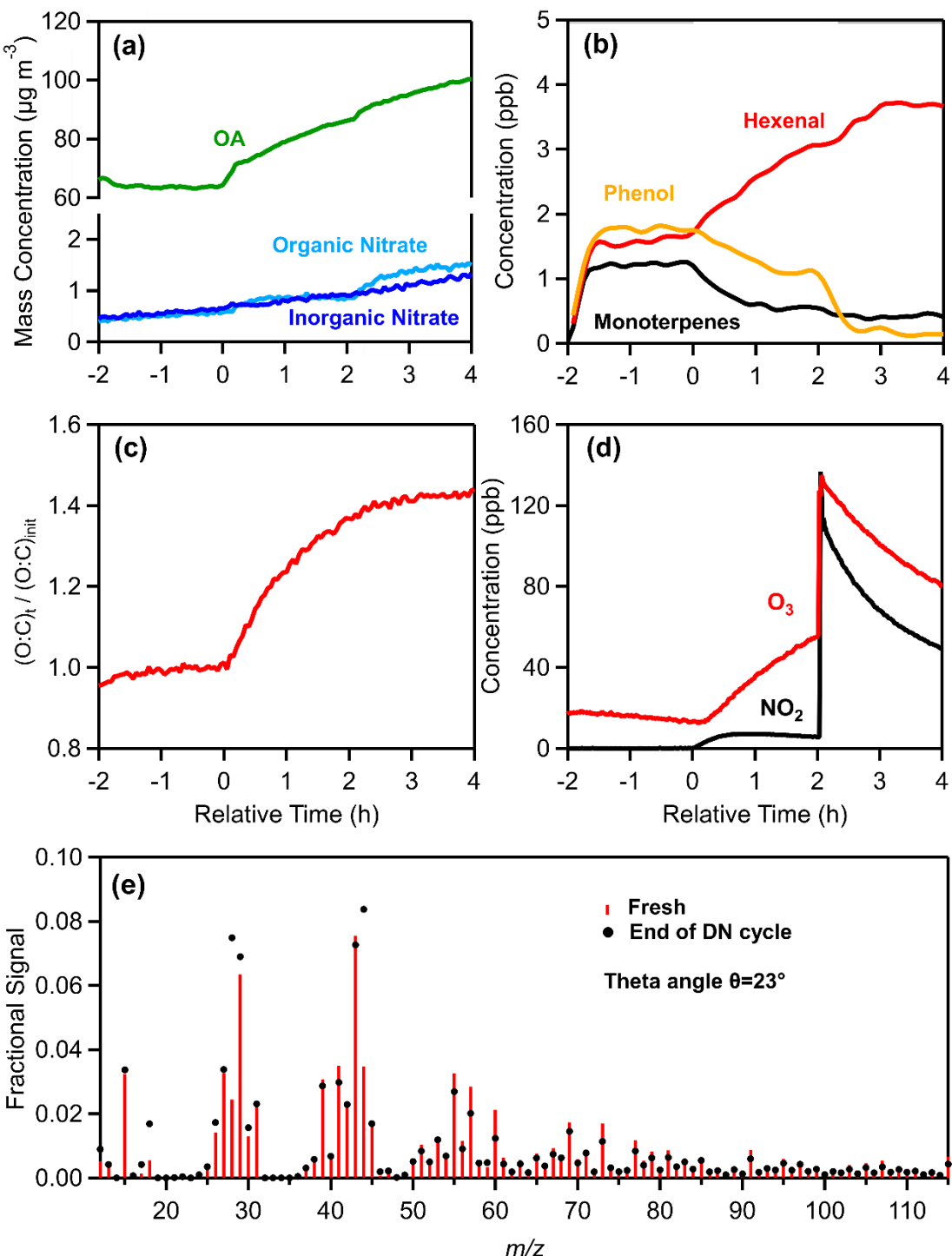
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1197 **Figure 1:** Experimental setup of the FORTH-ASC facility, illustrating the surrounding  
 1198 instrumentation and the combustion facility.

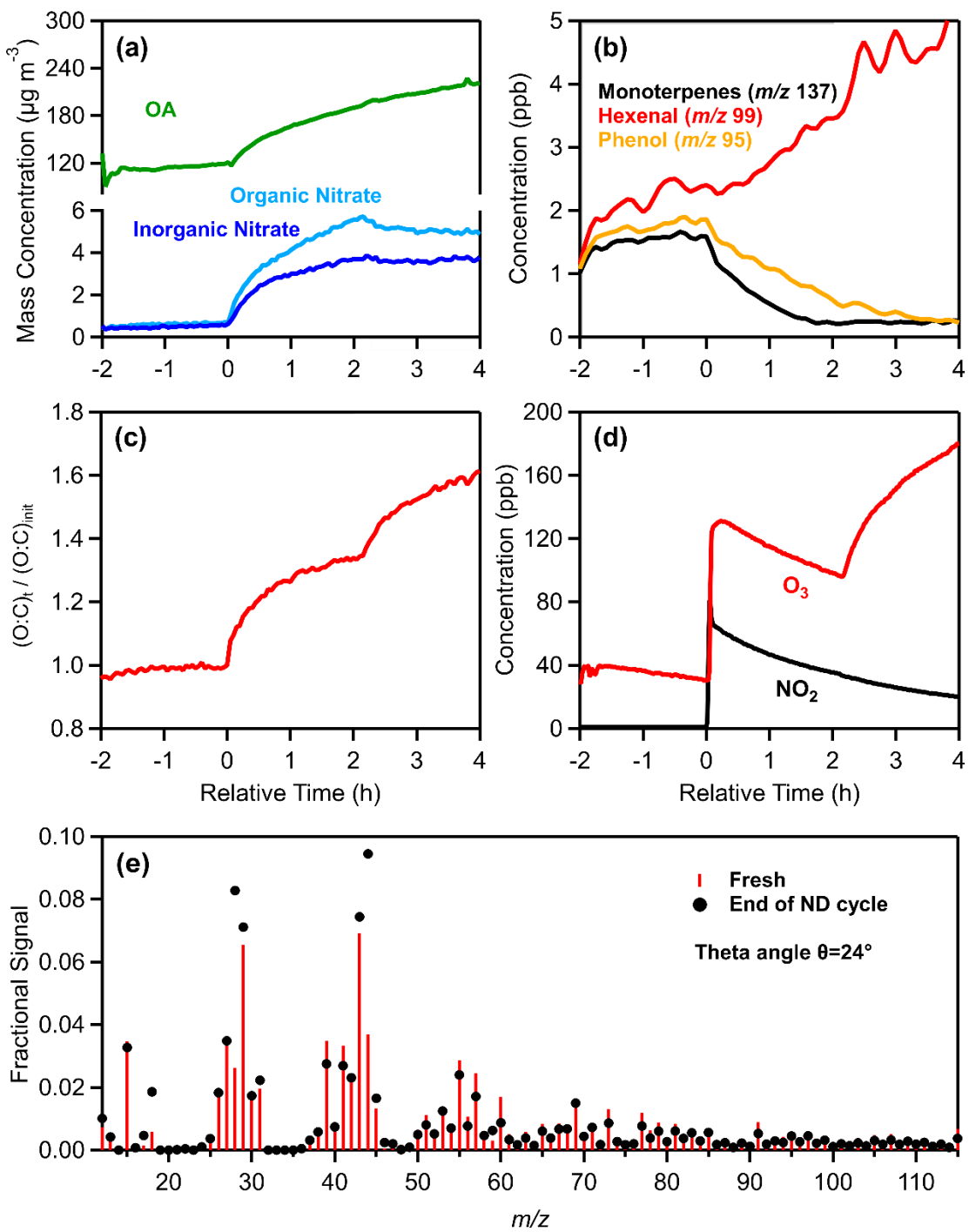


1199

1200 **Figure 2:** Average concentrations, in ppb, of the identified VOCs in (a) fresh olive wood  
 1201 burning emissions (red bars) and (b) fresh olive-pine mixed emissions (grey bars), along with  
 1202 their percentage contribution to the total VOCs concentration measured by PTR-QMS. The  
 1203 protonated *m/z* for each compound is shown in parentheses on the x-axis. The left y-axis shows  
 1204 the concentrations of identified VOCs, while the right y-axis displays the concentrations of the  
 1205 sum of the unidentified (other) and the total measured VOCs.

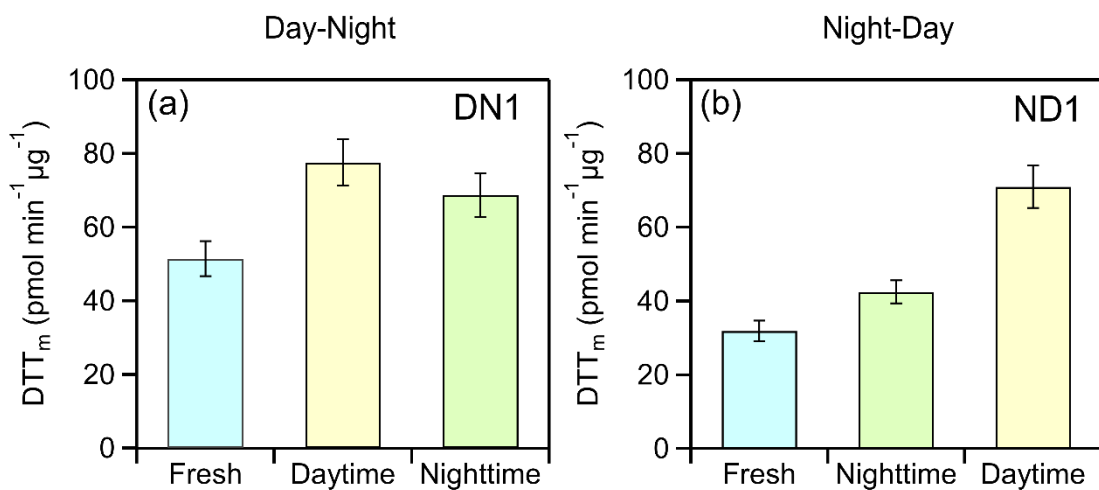


1208 **Figure 3:** Measurements from the experiment DN1, showing the time evolution of: (a) wall-  
 1209 loss-corrected organic aerosol, particulate organic and inorganic nitrate, (b) selected VOCs,  
 1210 including monoterpenes ( $m/z$  137), hexenal ( $m/z$  99), and phenol ( $m/z$  95), (c) normalized O:C  
 1211 ratio, (d)  $\text{O}_3$  and  $\text{NO}_2$ , and (e) a comparison of the fresh (red sticks) and nighttime (black  
 1212 markers) oxidized aerosol mass spectra at the end of the DN oxidation cycle.



1215 **Figure 4:** Measurements from the experiment ND1, showing the time evolution of: (a) wall-  
 1216 loss-corrected organic aerosol, particulate organic and inorganic nitrate, (b) selected VOCs,  
 1217 including monoterpenes ( $m/z$  137), hexenal ( $m/z$  99), and phenol ( $m/z$  95), (c) normalized O:C  
 1218 ratio, (d)  $\text{O}_3$  and  $\text{NO}_2$ , and (e) a comparison of the fresh (red sticks) and daytime (black  
 1219 markers) oxidized aerosol mass spectra at the end of the ND oxidation cycle.

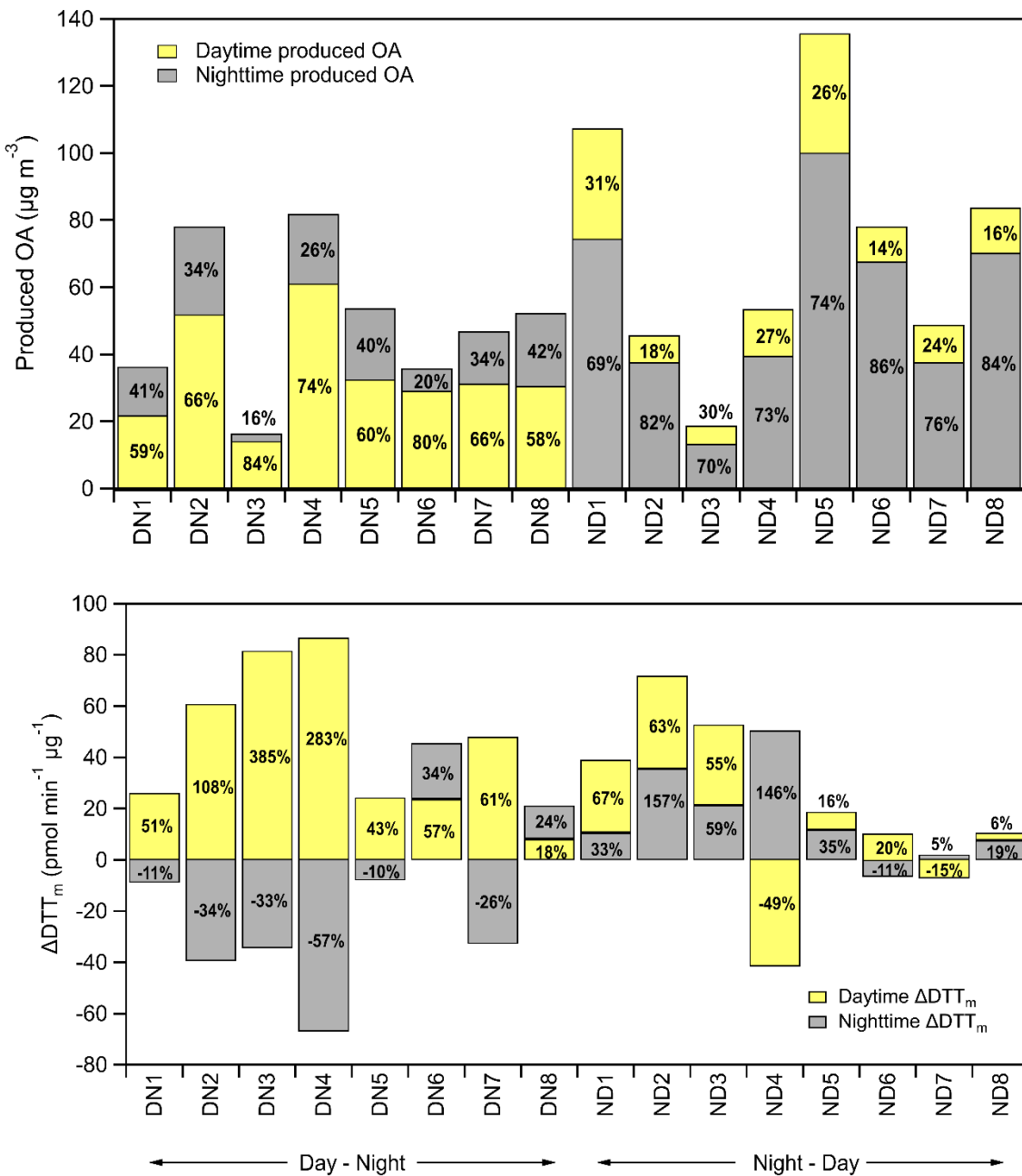
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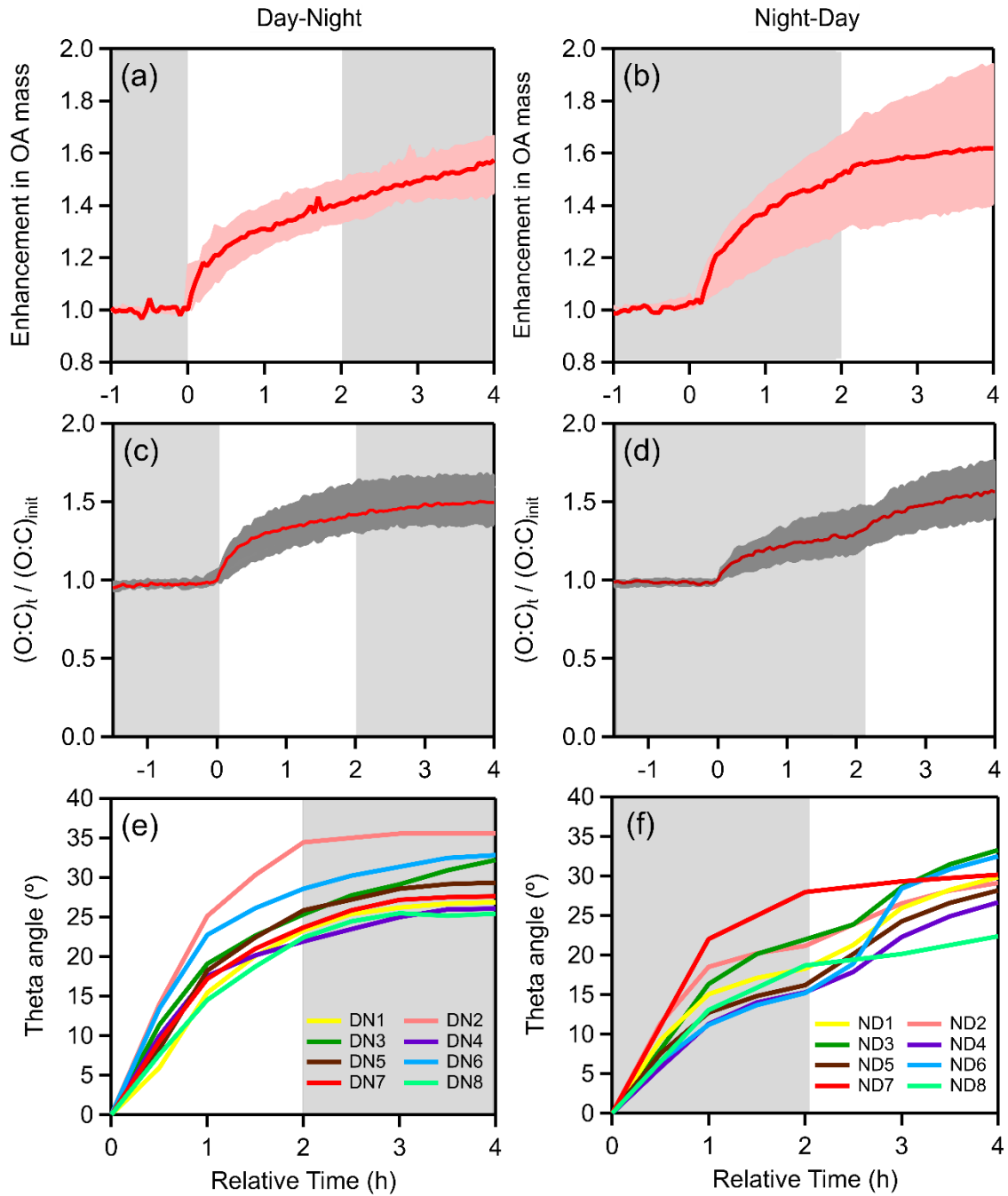
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1222 **Figure 5:** Category plots illustrating the evolution in water-soluble oxidative potential (WS-  
1223 OP), expressed as per OC mass normalized DTT<sub>m</sub> activity (pmol min<sup>-1</sup> μg<sup>-1</sup>), in case of typical  
1224 experiment (a) DN1 and (b) ND1.

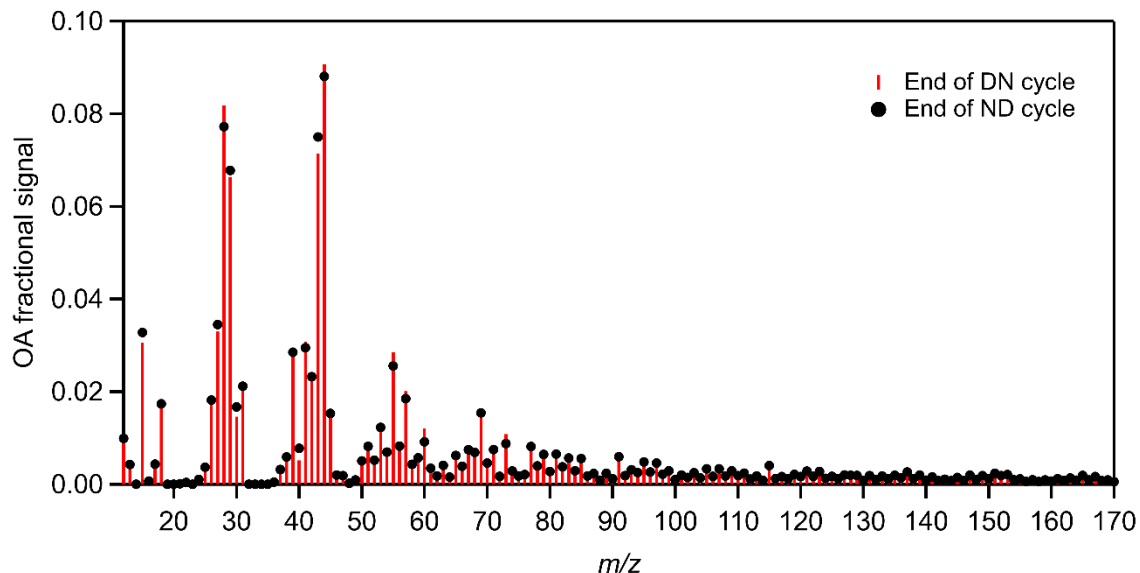
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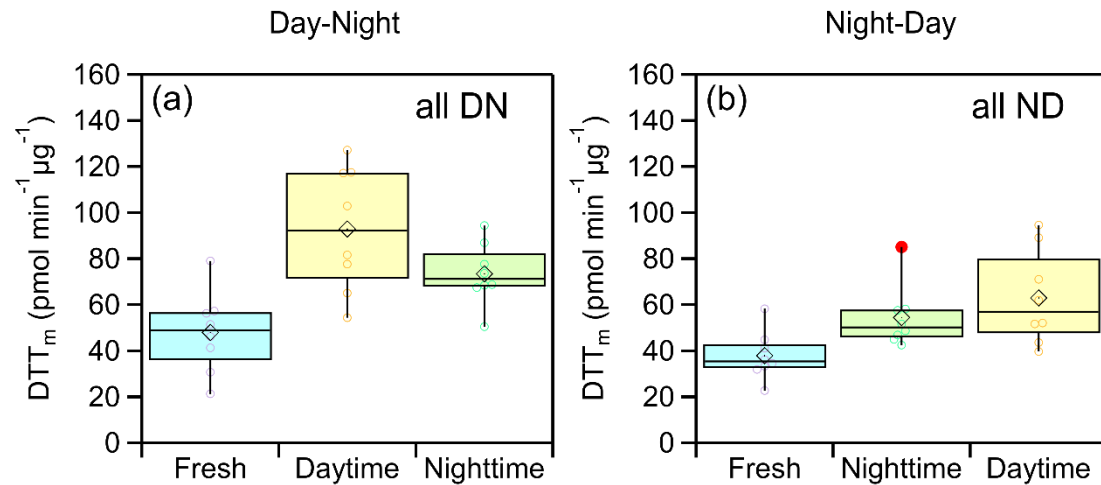
1228 **Figure 6:** (a) Absolute mass (in  $\mu\text{g m}^{-3}$ ) and percentage increase (%) of OA (including organic  
 1229 nitrate) per oxidation regime (daytime, nighttime); (b) absolute change in DTT<sub>m</sub> activity  
 1230 ( $\Delta\text{DTT}_m$ , in  $\text{pmol min}^{-1} \mu\text{g}^{-1}$ ) and percentage change (%) per oxidation regime (daytime,  
 1231 nighttime) for both DN and ND cycles, for all conducted experiments.



1234 **Figure 7:** Evolution over time of: OA enhancement during (a) DN and (b) ND oxidation cycle;  
1235 O:C ratio enhancement during (c) DN and (d) ND oxidation cycle; theta angle during (e) DN  
1236 and (f) ND oxidation for experiments conducted under dry initial conditions using only olive  
1237 wood logs as burning fuel (DN1-DN8, ND1-ND6). In experiments ND3, ND4, and ND6, the  
1238 change in spectrum occurred slightly later, as the first-step oxidation extended to 3 h compared  
1239 to 2 h lasted in the other ND experiments.



1242 **Figure 8:** Relative differences in the average spectra obtained at the end of DN (red sticks) and  
 1243 ND (black circles) oxidation cycle, respectively, for experiments conducted using olive wood  
 1244 logs as burning fuel. The theta angle between the averaged DN and ND aged spectra was 3°  
 1245 (identical).



1248 **Figure 9:** Box plots illustrating the changes in WS-OP, expressed as per OC mass normalized  
 1249 DTT<sub>m</sub> activity (pmol min<sup>-1</sup> µg<sup>-1</sup>), considering all performed experiments, in case of (a) DN  
 1250 oxidation cycle and (b) ND oxidation cycle.