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### ► To cite this version:

J. Kammer, P.-M. Flaud, A. Chazeaubeny, Raluca Ciuraru, K. Le Ménach, et al.. Biogenic volatile organic compounds (BVOCs) reactivity related to new particle formation (NPF) over the Landes forest. *Atmospheric Research*, 2020, 237, pp.1-11. 10.1016/j.atmosres.2020.104869 . hal-02553094

HAL Id: hal-02553094

<https://hal.inrae.fr/hal-02553094>

Submitted on 21 Jul 2022

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1 **Title: Biogenic Volatile Organic Compounds (BVOCs)**  
2 **reactivity related to New Particle Formation (NPF) over the**  
3 **Landes forest**

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24        **Abstract**

25        Atmospheric particles play a major role in both air quality and climate change. Formation of  
26        secondary particles in the atmosphere has been observed over many different environments  
27        and is believed to provide up to half of the atmospheric cloud condensation nuclei (CCN) at a  
28        global scale. However, high uncertainties are still remaining in the description of mechanisms  
29        involved in new particle formation (NPF). Especially, more evidences of the implication of  
30        biogenic volatile organic compounds (BVOCs) in NPF from field studies are still needed. To  
31        investigate this question, two field campaigns have been set up during July 2014 and July  
32        2015, in the French Landes forest (south west of France). Summer 2015 was characterised  
33        by a strong hydric stress, whereas summer 2014 was rainy. In 2015, frequent nocturnal NPF  
34        was observed, reaching a frequency of occurrence of ~55% of the nights, while only one  
35        event was observed in 2014. In July 2015, monoterpene mixing ratios (dominated by  $\alpha$ - and  
36         $\beta$ -pinene) were higher, mostly due to high ambient temperatures and drought. A focus was  
37        made on the 2015 field campaign, where NPF was mostly observed. The mean diurnal  
38        variation of the ratio between  $\alpha$ - and  $\beta$ -pinene mixing ratios highlighted in-canopy reactivity of  
39        monoterpenes with ozone in the early night. This hypothesis was reinforced by the increasing  
40        gas phase levels of pinonaldehyde and nopinone, the main first-generation products arising  
41        from  $\alpha$ - and  $\beta$ -pinene ozonolysis, at night, before NPF started. It strongly suggests that  
42        monoterpene oxidation further generated very-low volatility gases involved in NPF. This  
43        finding is also supported by the high concentrations of the SOA traditional biogenic tracers,  
44        *e.g.* pinic and pinonic acids, quantified in the particulate phase. The role of BVOCs in NPF is  
45        thus highlighted, as well as the importance of nighttime NPF.

46

47 **Keywords:** Monoterpenes, BVOCs, oxidation products, nighttime NPF, Landes forest

48 **Highlights**

- 49 • High monoterpene mixing ratios were related to high nighttime NPF frequency.
- 50 • In canopy  $\alpha$ - and  $\beta$ -pinene ozonolysis was highlighted in the evening.
- 51 • Nopinone and pinonaldehyde levels increase before NPF started.
- 52 • Biogenic tracer (pinic and pinonic acids) concentration levels were high in the
- 53 particulate phase.
- 54 • BVOC oxidation was strongly involved in nighttime NPF in the Landes forest.

55

## 56 **1 Introduction**

57 Forest ecosystems are key components of our environment, representing around 30% of the  
58 Earth land surface (United Nations food and agricultural organization, 2015). Also, forests  
59 represent a large source of Volatile Organic Compounds (VOCs). It is widely recognized that  
60 90% of VOCs are emitted by biogenic sources, isoprene and monoterpenes being the most  
61 emitted biogenic VOCs (BVOCs) in the atmosphere (Guenther et al., 1995; Sindelarova et  
62 al., 2014). Once oxidized in the atmosphere, some of their products have sufficiently low  
63 volatility to partition between the gas and the particle phases. Through this process, forests  
64 act as a source of secondary organic aerosol (SOA), which play a crucial role in air quality  
65 and climate change (Hallquist et al., 2009; IPCC, 2013; Kazil et al., 2010; Pope III and  
66 Dockery, 2006).

67 New Particle Formation (NPF) is a process that consists of the secondary formation of  
68 atmospheric particles (Dal Maso et al., 2005). Currently, NPF is estimated to contribute  
69 around half of the global Cloud Condensation Nuclei (CCN) at a global scale (Merikanto et  
70 al., 2009). NPF was observed in a large number of locations worldwide (Kulmala et al.,  
71 2004). However, processes governing NPF remain quite uncertain (Andreae, 2013; Kulmala  
72 et al., 2013; Nallathamby et al., 2014). Until recently, it was accepted that NPF could not  
73 occur without the involvement of sulphuric acid in the first step, *i.e.* nucleation (Andreae,  
74 2013; Kulmala, 2003; Kulmala et al., 2013; Nallathamby et al., 2014). In the atmosphere, gas  
75 phase sulphuric acid is mainly generated from SO<sub>2</sub> reaction with OH radicals in the presence  
76 of water. It may also be produced in the gas phase by oxidation of SO<sub>2</sub> with H<sub>2</sub>O<sub>2</sub> or by  
77 reaction with Criegee intermediates (Sipilä et al., 2014). Due to its very low vapour pressure,  
78 sulphuric acid may easily form clusters with other molecules, *i.e.* water or oxygenated  
79 molecules (Elm et al., 2014; Kulmala, 2003; Neitola et al., 2015; Praplan et al., 2015;  
80 Riccobono et al., 2014). Then, if this cluster is stable, it can grow through the condensation  
81 of extremely low volatility organic compounds (ELVOCs), produced from autoxidation (Ehn et  
82 al., 2014; Kulmala et al., 2013; Praplan et al., 2015; Riccobono et al., 2014). Recently, it was

83 demonstrated from laboratory studies that NPF can be caused solely by  $\alpha$ -pinene ozonolysis  
84 (Kirkby et al., 2016; Tröstl et al., 2016). It was highlighted from the CLOUD chamber  
85 experiments that highly oxidized molecules (like HOMs) can be rapidly formed, and induce  
86 NPF, following autoxidation mechanisms (Ehn et al., 2014; Jokinen et al., 2014). Direct  
87 evidence of such implication of HOMs in the real atmosphere was highlighted very recently in  
88 the free troposphere, where one of the main driver of NPF could be anthropogenic VOCs  
89 (Bianchi et al., 2016). More, Rose *et al.* (2018) observed that biogenic ions can induce stable  
90 cluster formation. But identification of biogenic NPF still needs more investigations in real  
91 atmosphere, especially at pristine locations. As a consequence, new measurements in  
92 ecosystems poorly or not yet investigated are needed.

93 In this context, the goal of the present study was to reinforce our knowledge on the relation  
94 between BVOCs and NPF. For that purpose, two field measurement campaigns took place in  
95 the summers of 2014 and 2015 in the Landes forest, in the south west of France. A first  
96 paper by our group recently evidenced frequent nocturnal NPF in 2015, together with high  
97 monoterpene mixing ratios (Kammer et al., 2018). This new paper first presents a  
98 comparison between 2014 and 2015 field campaigns. Then, the focus is made on the main  
99 emitted BVOCs at the measurement site, their reactivity and their oxidation products both in  
100 the gas and the particulate phases. Finally, considering the recent findings, the link between  
101 in canopy BVOC reactivity and NPF is discussed.

## 102 **2 Methodology**

### 103 **2.1 Site description**

104 As part of the LANDEX-episode zero project, two intensive field campaigns were conducted,  
105 in the French Landes forest. The first campaign started from the 5<sup>th</sup> of July until the 18<sup>th</sup> of  
106 July 2014, and the second started from the 5<sup>th</sup> to the 23<sup>rd</sup> of July 2015. The measurement  
107 site, located at Bilos (44°29'39.69"N, 0°57'21.75"W, and 37 m above sea level) in the  
108 northern part of the Landes forest, is a measurement site of the European Integrated Carbon

109 Observation System (ICOS). The nearest urban area is Bordeaux and surroundings, at 40  
110 km northeast, and the Atlantic Ocean is 23 km west of the site. The measurement area is a  
111 maritime pine (*Pinus pinaster* Aiton) stand sowed in 2004, with an understorey mostly  
112 composed of gorse (*Ulex europaeus* L.), grass (*Molinia caerulea* (L.) Moench) and heather  
113 (*Calluna vulgaris* (L.) Hull). Tree height was around 7 m in 2014 and 8 m in 2015, over a  
114 sandy hydromorphic podzol. A more complete description of Bilos site can be found  
115 elsewhere (Moreaux *et al.*, 2011; Kammer *et al.*, 2018). Due to the proximity of the Atlantic  
116 Ocean, the local climate is oceanic, and winds are consequently frequently originating from  
117 west-north west.

## 118 **2.2 Field measurements**

119 Meteorological parameter measurements have been performed on a mast during each  
120 campaign, at 15 m above ground level (a.g.l.). Air temperature and relative humidity (HMP  
121 155, VAISALA HUMICAP), wind speed and direction (Windsonic 1, Campbell scientific),  
122 global solar radiation (CMP22 and CNR4, Campbell scientific) and rainfall was continuously  
123 measured at a half hourly scale. To complete these meteorological measurements, air mass  
124 backward trajectories have been computed using the NOAA HYSPLIT model (Stein *et al.*,  
125 2015). Daily backward trajectories were calculated for each day of the campaign using the  
126 GDAS (Global Data Assimilation System) meteorological dataset, indicating the origin of the  
127 analysed air mass at Bilos. Besides, eddy covariance fluxes of momentum, heat, water  
128 vapour and CO<sub>2</sub> were measured at 15 m a.g.l., in order to evaluate the physiological state of  
129 the ecosystem. The details of the eddy covariance methods have been fully described  
130 elsewhere (Aubinet *et al.*, 2000; Burba and Anderson, 2010).

131 A mobile laboratory truck (the “Barracuda” facility), located in the centre of the area, was  
132 used for physico-chemical measurements with air tubing devoted to aerosol and gas  
133 sampling being located at 4 m a.g.l., corresponding to the canopy level. A 5 m Teflon tubing  
134 (4.5 mm inner diameter) was used for the gas phase measurements, whereas a 4 m long

135 conductive silicone tubing (4.5 mm inner diameter) was used to sample particles. Both gas  
136 and particle sampling inlets were located right next (about 20 cm) to ensure the collection of  
137 the same air masses. Preliminary tests were performed by measuring the same air with and  
138 without tubing. As no difference was observed, it is assumed that no significant wall losses of  
139 terpenes occurred in tubing. During the campaign, ozone and nitrogen oxides ( $\text{NO}_x = \text{NO} +$   
140  $\text{NO}_2$ ) mixing ratios were monitored through a UV absorption analyzer (APOA 370, HORIBA,  
141 detection limit (DL) = 0.5 ppb) and a chemiluminescence analyzer (APNA 370, HORIBA, DL  
142 = 0.5 ppb), respectively. Sulphur dioxide ( $\text{SO}_2$ ) measurements were performed only during  
143 the 2015 field campaign using a UV fluorescence monitor (API 100 E, Teledyne, DL = 0.4  
144 ppb). All the three monitors ( $\text{O}_3$ ,  $\text{NO}_x$  and  $\text{SO}_2$ ) have been calibrated before the campaigns,  
145 and leak checks were frequently performed using activated charcoal air filters. A scanning  
146 mobility particle sizer (SMPS, TSI model 3080) was used to characterize particle number  
147 size distribution every 10 minutes. It combines a differential mobility analyser (DMA, TSI  
148 model 3081) and an optical counter (CPC, TSI model 3772). Aerosol flow and sheath flow  
149 were set respectively to  $0.5 \text{ L min}^{-1}$  and  $5 \text{ L min}^{-1}$  to ensure a 1:10 ratio in the DMA. Under  
150 these flow conditions, particle number size distribution and concentration were measured for  
151 particles having an electric mobility diameter between 10.9 nm and 487 nm. Particle wall  
152 losses were calculated following the methodology described in Baron and Willeke (2001).  
153 The estimated particle loss rates were estimated to be 34.7% for the smallest particles (with  
154 a 10.9 nm diameter) analysed. The particle number and size distribution data were corrected  
155 to account for the calculated wall losses. Then, a proton transfer reaction - time of flight -  
156 mass spectrometry (PTR-TOF-MS, Kore Technology) was used to study the composition of  
157 the gaseous phase, especially the evolution of BVOCs and their related oxidation products.  
158 This instrument is based on the proton transfer reaction from  $\text{H}_3\text{O}^+$  to the analytes, a soft  
159 ionisation which prevents from strong fragmentation. Basically, the air mass composition  
160 could thus be analyzed by recording the temporal evolution of the  $\text{MH}^+$  ions. During the  
161 campaign, the PTR-TOF-MS was operated at 600 V drift voltage and 1.3 mbar reactor  
162 pressure, ensuring an E/N ratio around 130 Td ( $1 \text{ Townsend} = 10^{-17} \text{ V cm}^{-2}$ ), with E being the



163 electric field strength in the reactor and  $N$  the gas number density. The PTR-TOF-MS was  
164 tuned to work at this  $E/N$  ratio as it is a good compromise to prevent from high fragmentation  
165 in the mass spectrometer and ambient relative humidity effect (Pang, 2015; Tani et al.,  
166 2003). PTR-TOF-MS blanks and calibrations were daily performed using the vapour pressure  
167 (at 0°C and diluted in  $N_2$  flow) of a pure  $\alpha$ -pinene solution, regulated by a mass flow controller  
168 (Millipore, Coastal Instruments). Typical day to day sensitivity variations were about 10 %.  
169 The variation of the sensitivity may be sometimes larger, when the PTR-TOF-MS parameters  
170 (for example, the voltage applied to extraction lenses) were optimized during the campaign.  
171 For non-calibrated compounds (other than  $\alpha$ -pinene), calibration coefficients determined for  
172  $\alpha$ -pinene were used to consider the day-to-day sensitivity variations of the instrument. But as  
173 we could not consider that these calibration coefficients provide a real estimation of the  
174 concentration of these other compounds, then each signal was normalized by its maximum  
175 count number.

176 As monoterpenes  $C_{10}H_{16}$  present many isomers and as the PTR-TOF-MS was only able to  
177 measure their total concentrations, an online gas chromatograph coupled to a flame  
178 ionization detector (GC-FID, airmoVOC, Chromatotec) was used to discriminate their  
179 speciation. The GC-FID sampled air at  $0.11 \text{ L min}^{-1}$ , and concentrated VOCs on a cold air  
180 trap composed of adsorbent Tenax TA (20–35 mesh, Chrompack). The desorbed VOCs to  
181 analyse were then separated using a Chrompack Sil 8CB low bleed capillary column, and  
182 finally detected with the FID. A complete description of the GC-FID can be found elsewhere  
183 (Staudt and Lhoutellier, 2011). In 2015, the GC-FID was only available during the last five  
184 days of the campaign.

185 The chemical composition of particles collected during the 2015 field campaign was  
186 investigated using high performance liquid chromatography coupled to mass spectrometry.  
187 Briefly, a high volume sampler (DA-80, Digitel, with an air flow set to  $30 \text{ m}^3 \text{ h}^{-1}$ ) was used to  
188 collect  $PM_{2.5}$  (particles with an aerodynamical diameter smaller than  $2.5 \mu\text{m}$ ) onto quartz fiber  
189 filters ( $\varnothing = 150 \text{ mm}$ ). Different laboratory tests showed that ultrasonic extraction provides the

190 highest and more reproducible extraction yields. If there are still some uncertainties related to  
191 this technique, it is widely used to measure pinic acid and pinonic acid in aerosol samples  
192 (Anttila et al., 2005; Feltracco et al., 2018; Kourtchev et al., 2014; Nozière et al., 2015). As a  
193 result, particles were extracted two times by placing a 47 mm diameter disc in 3 mL of a  
194 mixed acetonitrile/water solution (70/30) under ultrasonication for 15 min. Then, each extract  
195 was centrifugated and the supernatant was transferred in a 15 mL vial and concentrated to  
196 about 400  $\mu$ L under a gentle pure nitrogen flow (99.995 % purity, Linde Gas SA) at 40 °C.  
197 Samples were weighted using gravimetry (precision:  $10^{-4}$  g), then stored at -18°C until  
198 analysis. LC-HRMS analyses were carried out using an Agilent 1290 HPLC system coupled  
199 to an Agilent 6540 QToF mass spectrometer equipped with an Agilent Jet Stream  
200 electrospray ionization source (ESI) operating in the negative mode. The chromatographic  
201 separation was performed on a Zorbax Eclipse XDB-C18 (2.1 $\times$ 150 mm; 3.5  $\mu$ m) regulated at  
202 30°C. Gradient LC elution was performed at a flow rate of 0.5 mL min<sup>-1</sup> using 0.2% formic  
203 acid in purified water as mobile phase A and 0.2% formic acid in acetonitrile as mobile phase  
204 B. Sample injection volume was set at 5  $\mu$ L. In this study, pinic acid and pinonic acid were  
205 quantified using authentic standards of the two targeted compounds (pinic acid 98 % and cis-  
206 pinonic acid 98 %, Sigma-Aldrich) as external standards. More details about LC-QTOF-MS  
207 parameters can be found in the supplementary information (**Table S1**).

### 208 **3 Results and discussion**

#### 209 **3.1 Meteorological conditions during 2014 and 2015 field campaigns**

210 Meteorological parameters are crucial as they may greatly influence BVOC emissions and  
211 the oxidative capacity of the atmosphere. The Bilos site is for instance frequently subject to  
212 periods of large hydric stress during summer. Detailed meteorological conditions during 2014  
213 and 2015 field campaigns can be found in **Figures S1** and **S2**. For each campaign, the  
214 conditions were very different. The 2015 field campaign was very hot, with few rain events  
215 (see **Figures S1.a, d** and **Figures S2.a, d**). During this last campaign, the air temperature

216 frequently exceeded 30°C during the day and was hardly ever below 15°C at night, which  
217 was not the case in July 2014. Such high temperature values were already reported to  
218 induce a thermic stress on trees (Laothawornkitkul et al., 2009). One particularity of the site  
219 was that relative humidity increased to reach its maxima (*i.e.* 100%) almost every night, in  
220 both July 2014 and 2015.

221 No rainfall was observed from the middle of June 2015 (meteorological parameters outside  
222 the campaign range are available from the ICOS network) and only few mm of rain was  
223 recorded between the 5<sup>th</sup> and the 22<sup>nd</sup> July in 2015. As a result, a very weak water vapour  
224 flux was observed in July 2015, whereas the sensible heat flux was very high (**Figure S2.b**),  
225 especially comparing to 2014 fluxes (**Figure S1.b**). The Bowen ratio (between sensible heat  
226 fluxes and water vapour fluxes) is a powerful tool to evaluate the hydric stress level. During  
227 drought, the water vapour fluxes decrease whereas the heat fluxes increase (by  
228 compensation to conserve the energy balance), making the Bowen ratio particularly  
229 elevated. Indeed, the 2015 campaign was characterized by Bowen ratio values about 10,  
230 which is very high for a forest stand (Gu et al., 2006). Such high Bowen ratios were not  
231 observed during the 2014 field campaign, with values only around 0.5 (Kammer et al., 2019).  
232 Thus, a thermic stress (due to temperatures over 30°C) coupled to a hydric stress occurred  
233 during the 2015 campaign. The presence of such a stress situation may increase BVOC  
234 emissions (Kesselmeier and Staudt, 1999; Laothawornkitkul et al., 2009; Loreto et al., 1998;  
235 Peñuelas and Staudt, 2010). Wind directions indicated that air masses were mostly coming  
236 from west-northwest direction in 2014 as well as in 2015 (**Figures S1.e** and **S2.e**), which was  
237 confirmed by backward air mass trajectories calculations. Air masses were thus mostly  
238 influenced by the Atlantic Ocean in both campaigns.

### 239 **3.2 General conditions of the campaigns**

240 A previous study reported by our group (Kammer et al., 2018) showed that NPF has been  
241 frequently observed during the 2015 field campaign, mostly during the night, although a few

242 events (only 2) were also recorded during daytime. NPF events were defined as an increase  
243 of particle concentration in the nucleation mode followed by a subsequent growth during at  
244 least 2 hours (Dal Maso et al., 2005). In 2014, only one NPF event was observed, occurring  
245 during nighttime. This leads to a NPF frequency of occurrence (i.e. the number of nights  
246 where NPF events were observed divided by the total number of nights during the field  
247 campaign) of about 8% during the 2014 field campaign. In 2015, 12 NPF events were  
248 reported, reaching a high frequency of occurrence of around 55% (Kammer et al., 2018).  
249 NPF thus occurred more frequently in a summer influenced by a hydric stress (according to  
250 the high Bowen ratio values measured during the 2015 campaign). This observation may  
251 further suggest that the presence of a hydric stress is indirectly inducing more NPF events  
252 because of stronger monoterpene emissions. Nevertheless, more studies about the effect of  
253 hydric stress on NPF will be needed to confirm such hypothesis. All the observed NPF  
254 events are listed in **Table S2**. A more complete analysis of NPF events (growth rates,  
255 nucleation rates, etc.) can be found in Kammer et al. (2018).

256 Analyses of wind directions and air mass back trajectories have shown that air mass origin  
257 did not affect NPF. During both campaigns, NPF were always recorded along (i) high relative  
258 humidity (i.e.  $RH \geq 75\%$ ), (ii) temperatures higher than  $16^\circ\text{C}$  and (iii)  $1/u^*$  (where  $u^*$  is the  
259 friction velocity) over  $2.5 \text{ s m}^{-1}$ , indicating vertical stratification (this parameter was usually  
260 below  $2 \text{ s m}^{-1}$  during the day where turbulence occurred, see the meteorological parameters  
261 in **Figures S1** and **S2** during NPF events). It shows that NPF started once the stable  
262 nocturnal boundary layer was taking hold. As suggested by the air mass analysis, nocturnal  
263 NPF were very likely influenced by local emissions, limiting the impact of transported  
264 compounds.

265  $\text{NO}$ ,  $\text{NO}_2$ ,  $\text{SO}_2$  (only for 2015 campaign) and ozone mixing ratios are represented on **Figures**  
266 **S3** and **S4**.  $\text{NO}_x$  mixing ratios were very low during both campaigns, confirming the very rural  
267 feature of the Bilos site. The mixing ratios of  $\text{SO}_2$ , which is recognized as a key species for  
268 daytime nucleation, was always around the DL value of the instrument (only available in

269 2015, **Figure S4.b**). As a consequence, and taking into account the nocturnal nature of NPF,  
270 it is highly probable that sulphuric acid was not responsible for nocturnal NPF at Bilos,  
271 conversely to classical daytime NPF. Ozone mixing ratios presented a typical diurnal cycle  
272 with maximal values reached during the afternoon, and frequently dropped under the  
273 analyser limit (*i.e.* 0.5 ppb) in the middle of the night. The nights of July 15<sup>th</sup> and 16<sup>th</sup> in 2014  
274 and 12<sup>th</sup> to 15<sup>th</sup>, 17<sup>th</sup>, 18<sup>th</sup> and 20<sup>th</sup> in 2015 are perfect examples of this ozone drop (**Figures**  
275 **S3.c** and **S4.c**). The meteorological conditions observed during these nights indicate that  
276 these ozone drops occurred under stratified atmosphere conditions (low wind speed and high  
277  $1/u^*$ ).

### 278 **3.3 Monoterpene mixing ratios**

279 The sum of monoterpene as well as ozone mixing ratios are represented in Figure 1 for both  
280 field campaigns. A strong diurnal cycle was observed in July 2014 as well as in July 2015 for  
281 both ozone and monoterpenes. A typical ozone diurnal cycle was observed, with higher  
282 values in the afternoon (Figure 1). The ozone cycle was mirrored by the monoterpenes cycle,  
283 with higher monoterpene mixing ratios values during the night, as previously reported in the  
284 Landes forest (Simon et al., 1994). Monoterpenes were dominated from far by  $\alpha$ - and  $\beta$ -  
285 pinene, accounting for around 90% of total monoterpenes, in accordance with previous  
286 studies (Riba et al., 1987; Simon et al., 1994). Maritime pines are well-known to be strong  
287 temperature driven BVOC emitters (Kesselmeier and Staudt, 1999; Lathiere et al., 2006).  
288 Although emissions are higher during the day, the larger dilution of emissions in the surface  
289 layer and the higher oxidant levels conduct to lower BVOC mixing ratios during light hours.  
290 More explanation about the diurnal cycle and the effect of the change in the boundary layer  
291 height on monoterpene mixing ratios can be found in Kammer *et al.* (2018). In 2015,  
292 monoterpene mixing ratios frequently (*i.e.* almost half of the nights) reached around 15 ppb,  
293 whereas they were always lower than 10 ppb in 2014, except the night of July 16<sup>th</sup> to 17<sup>th</sup>  
294 (**Figure 1**). The highest values were reached the 21<sup>st</sup> of July 2015, where the sum of  
295 monoterpene mixing ratios exceeded 35 ppb (**Figure 1.b**). The very high mixing ratios of

296 monoterpenes observed in 2015 were assumed to be the result of the meteorological  
297 conditions, and more specifically the thermic/hydric stress. The high frequency of occurrence  
298 of NPF in 2015 occurred for the highest monoterpene mixing ratios, emphasizing the role of  
299 monoterpenes at night at Bilos.

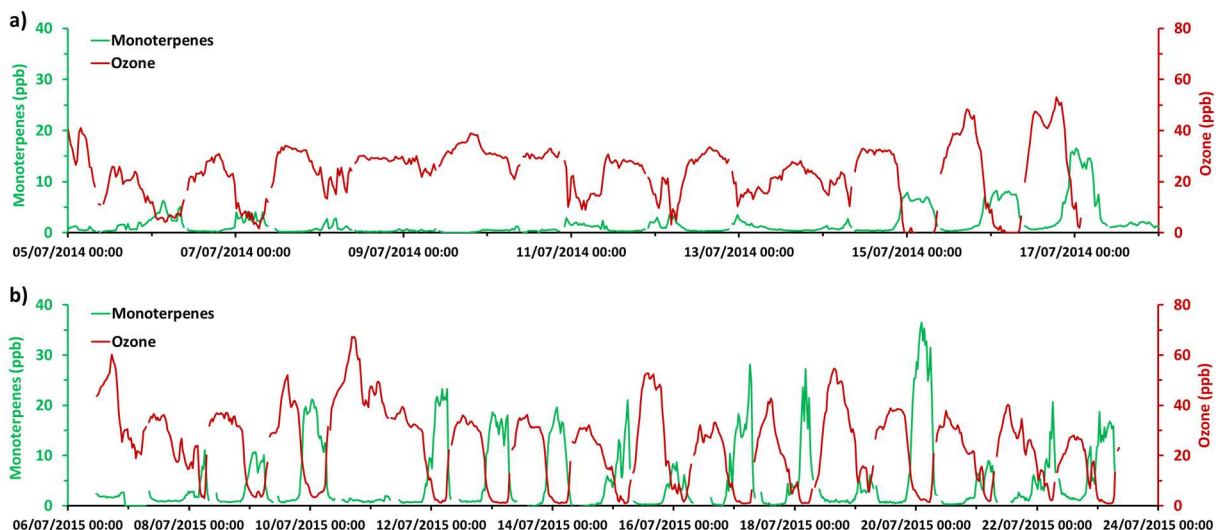


Figure 1: Time series of the sum of monoterpene mixing ratios (in ppb, green line) and ozone mixing ratios (in ppb, red line) for **a)** 2014 and **b)** 2015 field campaigns.

### 300           3.4       Monoterpene/O<sub>3</sub> reactivity

301 Time series of ozone and monoterpene mixing ratios (**Figure 1**) suggest that their respective  
302 diurnal cycles were closely anti-correlated. The monoterpene mixing ratios increased every  
303 night, as reported on **Figure 1**, when it is likely that no ozone was produced (because of its  
304 dependence on solar radiation), nor ozone advection from higher layers of the atmosphere  
305 (because of the nocturnal stratified boundary layer). The anti-correlation between ozone and  
306 monoterpenes can be observed on **Figure 2.a**. Interestingly, when ozone was close to 0,  
307 monoterpene mixing ratios values were at least of 5 ppb during the corresponding nights  
308 (**Figure 2.a**). Reaction between ozone and monoterpenes in the early night could thus be  
309 expected. Such reactions could only partially explain the ozone drop to levels close to zero,  
310 as their kinetic rate constants are quite low (Atkinson and Arey, 2003). A rough estimation,  
311 assuming pseudo-first order reactions between ozone and monoterpenes, led to a maximal

312 contribution to the ozone decrease being around 50 %. Thus, there is also a possibility for  
313 other reactive BVOCs to be co-emitted with monoterpenes and react very fast with ozone  
314 (like  $\beta$ -caryophyllene, for example). Further modelling of ozone deposition and chemical  
315 reactions between ozone and BVOCs will be needed for a detailed estimation.

316 As monoterpene emissions come from the diffusion of resin stored in pools, the ratio  
317 between  $\alpha$ - and  $\beta$ -pinene in maritime pine emissions is assumed to be constant (Simon et al.,  
318 1994). Hence, changes in the  $\alpha/\beta$ -pinene ratio can mostly be assigned to differences in  
319 oxidation processes, as the reactivity of  $\alpha$ - and  $\beta$ -pinene with the main atmospheric oxidants  
320 differs (see **Table 1**). The daily evolution of the  $\alpha/\beta$ -pinene ratio may provide an interesting  
321 tool to evaluate the importance of each oxidation processes. When the OH-initiated  
322 monoterpene photooxidation is the dominant process,  $\beta$ -pinene mixing ratio should decrease  
323 faster than that of  $\alpha$ -pinene, because of its larger reactivity with OH (**Table 1**). In the morning,  
324 the  $\alpha/\beta$ -pinene ratio measured at Bilos was increasing, meaning that photooxidation was  
325 increasingly dominated by OH radicals (**Figure 2-b**). In the afternoon, the  $\alpha/\beta$ -pinene ratio  
326 was found to drop, during both 2014 and 2015 field campaigns. Such decrease reflects that  
327 the dominating oxidation process was changing, as the reaction rate constants of  $\alpha$ -pinene  
328 with ozone and  $\text{NO}_3$  are known to be larger than for  $\beta$ -pinene (**Table 1**). It is recognized that  
329  $\text{NO}_3$  reactivity is mostly significant during nighttime, as nitrate radicals are photolysed by  
330 solar radiations (Seinfeld and Pandis, 2006). As a result, the drop of the  $\alpha/\beta$ -pinene ratio  
331 highlighted the dominant role of ozone oxidation processes in the afternoon.

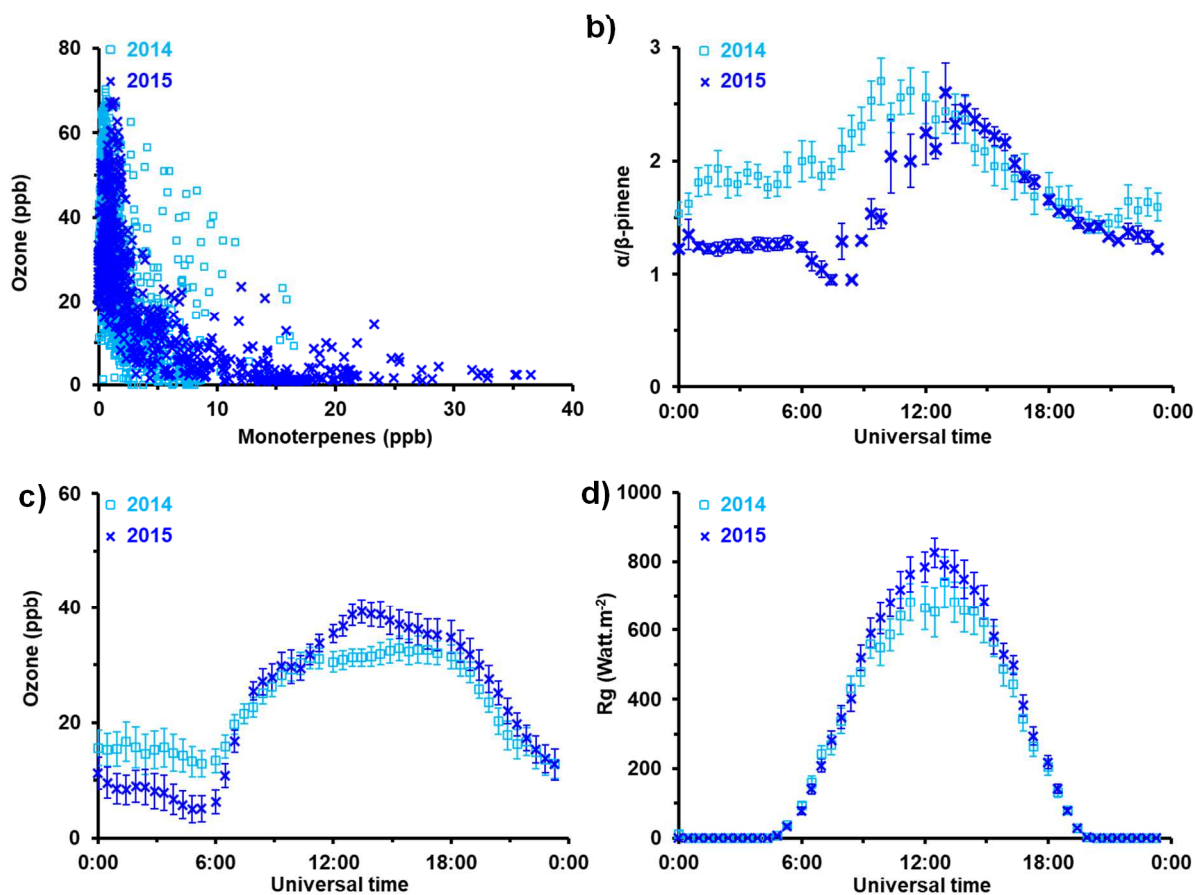
**Table 1:** Chemical kinetic rate constants of reactions of  $\alpha$ - and  $\beta$ -pinene with the main atmospheric oxidants (OH, O<sub>3</sub> and NO<sub>3</sub>)

	Rate constant (at 298K, in cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup> )		
	OH	O <sub>3</sub>	NO <sub>3</sub>
$\alpha$ -pinene	$(5.3\pm 0.2) \times 10^{-11}$	$(9.0\pm 0.2) \times 10^{-17}$	$(6.2\pm 0.1) \times 10^{-12}$
$\beta$ -pinene	$(7.6\pm 0.1) \times 10^{-11}$	$(1.9\pm 0.3) \times 10^{-17}$	$(2.5\pm 0.1) \times 10^{-12}$

*IUPAC recommended values (Atkinson et al., 2006)*

332 The **Figure 2.b** shows that the  $\alpha/\beta$ -pinene ratio mean diurnal cycle was more pronounced in  
333 2015. Solar radiation, taken as a simple proxy for OH concentration, and ozone mixing ratios,  
334 were higher in 2015 than in 2014 (**Figures 2.c** and **2.d**). Higher OH concentration or ozone  
335 mixing ratios led to a more pronounced increase or drop in the  $\alpha/\beta$ -pinene ratio values,  
336 respectively. The **Figure 2.b** illustrates that the mean diurnal cycle of the  $\alpha/\beta$ -pinene ratio  
337 logically followed the solar radiation and ozone diurnal profiles. The analysis of the mean  
338 diurnal evolution of the  $\alpha/\beta$ -pinene ratio also supports the hypothesis of the importance of  
339 ozone reactions with monoterpenes in the afternoon/evening. To check this hypothesis, the  
340 presence (or absence) of monoterpene oxidation products in the gas phase was  
341 investigated.  
342





**Figure 2:** **a)** Anti-correlation between ozone and monoterpene mixing ratios **b)** mean diurnal profile of the  $\alpha/\beta$ -pinene ratio **c)** mean diurnal profile of the ozone mixing ratios **d)** mean diurnal profile of global radiation, during each campaign. The error bars represent the standard errors of each variable. For each graphics, light blue squares represent 2014 dataset, and dark blue crosses represent 2015 dataset. (Universal time = local time – 2h)

343

### 344 3.5 BVOC oxidation products

345 As the PTR-TOF-MS has a good mass resolution, and considering the high mixing ratios of  
 346 both  $\alpha$ - and  $\beta$ -pinene at the Bilos site, several detected ions were able to be attributed to  
 347 biogenic oxidation products. The identification of the following ions was proposed based on  
 348 the proximity of the peak with the theoretical mass, and considering the characteristics of the  
 349 Bilos site (very rural, high monoterpene mixing ratios, etc.). However, a contribution of a  
 350 compound with a  $m/z$  very closed to the proposed compound could not be fully excluded.

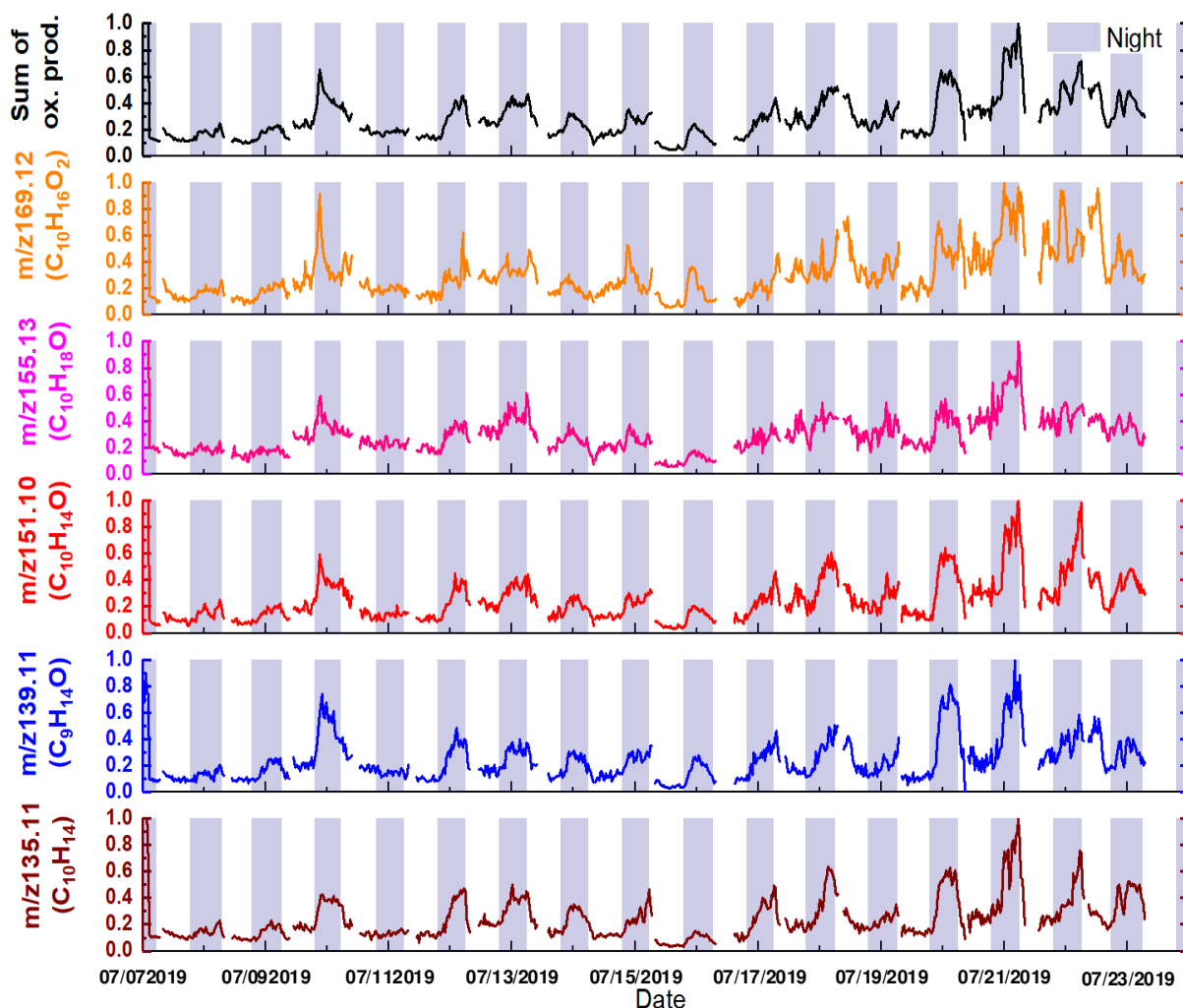
351 First, m/z 139.11 was proposed to be attributed to nopinone (C<sub>9</sub>H<sub>14</sub>O) and m/z 169.12 to  
352 pinonaldehyde (C<sub>10</sub>H<sub>16</sub>O<sub>2</sub>), as both compounds were frequently reported at rural field sites  
353 (Bourtsoukidis et al., 2014; de Gouw and Warneke, 2007; Park et al., 2014, 2013; Rantala et  
354 al., 2014; Ruuskanen et al., 2011). Relevant signals were also observed at m/z 135.11, m/z  
355 151.10 and m/z 155.13, but their attribution was more difficult. The m/z 135.11 protonated  
356 ion corresponds to compounds of the molecular formula C<sub>10</sub>H<sub>14</sub>. Among several potential  
357 hydrocarbons, the m/z 135.11 ion may correspond to *p*-cymene, which has already been  
358 reported by Gratien *et al.* (2011) as a secondary product of  $\alpha$ -pinene photo-oxidation. Even if  
359 *p*-cymene has not yet been reported among monoterpene emissions from maritime pines,  
360 some works have already pointed out its emission from different plant species (Bäck et al.,  
361 2012). *P*-cymene could thus have been emitted by understorey species (for which BVOC  
362 emissions remain unknown or poorly investigated), or result from  $\alpha$ -pinene photo-oxidation.  
363 *P*-cymene levels were found to be highly correlated with pinonaldehyde and nopinone ( $R^2 =$   
364 0.8), suggesting the secondary origin of this compound.

365 The molecular formula [C<sub>10</sub>H<sub>14</sub>O]H<sup>+</sup> was proposed to be the major contributor of m/z 151.10  
366 ion. This ion is recognized as a monoterpene oxidation product, and some studies attributed  
367 this mass to a significant pinonaldehyde fragment (de Gouw and Warneke, 2007; Holzinger  
368 et al., 2007; Park et al., 2014; Rantala et al., 2014). Note that *i*) other compounds such as  
369 myrtenal or verbenone, that may have primary sources, could also be important contributors  
370 to the m/z 151.10 signal (Kim et al., 2010) and *ii*) OH initiated oxidation of nopinone was also  
371 reported to yield products with m/z 151.10. Finally, the PTR-TOF-MS signal recorded at  
372 m/z 155.13 has been identified as C<sub>10</sub>H<sub>18</sub>O compounds. Several isomers can potentially  
373 contribute to this signal. Notably, primary OVOCs such as linalool or 1,8-cineol have already  
374 been reported at m/z 155.13 elsewhere (Bourtsoukidis et al., 2014; Park et al., 2013; Rantala  
375 et al., 2014). However, a contribution of secondary oxygenated compounds cannot be  
376 excluded. The presence of all such monoterpene oxidation products confirms that  
377 monoterpenes were oxidized at the Bilos site. Especially, pinonaldehyde and nopinone are

378 known to be exclusively formed from the oxidation (by ozone or OH radical) of  $\alpha$ - and  $\beta$ -  
379 pinene, respectively, supporting the occurrence and the importance of the oxidation of  
380 monoterpenes in the Landes forest. Indeed, OH can be formed as a secondary product of  
381 the ozonolysis of monoterpenes. Any observed OH reactivity during nighttime would thus be  
382 an indication of a primary monoterpene ozonolysis process. It was finally assumed that the  
383 increase of nopinone and pinonaldehyde during the night was arising from monoterpene  
384 ozonolysis.

385 The temporal evolution of the compounds of interest measured during the 2015 field  
386 campaign is represented on **Figure 3**. In this section, we make the choice to focus on the  
387 2015 data as *i*) oxidation product concentration values were frequently around the DL of the  
388 instrument in 2014 and *ii*) there was only one NPF event in 2014. This is consistent with  
389 lower monoterpene mixing ratios observed during this first campaign. As oxidation products  
390 were not quantified in this study, their corresponding signals were normalized to their  
391 maximum to allow the comparison of their temporal evolution profiles and diurnal dynamics.  
392 Interestingly, all the oxidation products globally followed the same diurnal cycle, similar to  
393 that observed for monoterpenes (**Figure 3**). Their levels started to increase with the sunset,  
394 then decreased when the first rays of the sunlight appeared (**Figure 3**). The decrease of the  
395 monoterpene oxidation product could be explained by *i*) a higher dilution due to the increase  
396 of the surface layer height as soon as the first rays break the stable nocturnal boundary layer  
397 and *ii*) potential photolysis, especially in the case of pinonaldehyde and nopinone, as both  
398 are carbonyl compounds prone for photolysis. Although oxidation products are quite volatile  
399 and thus are not expected to directly contribute significantly to NPF, they highlight a strong  
400 reactivity between ozone and monoterpenes, also yielding (at night) to potential very low  
401 volatile oxidation products that may contribute to nocturnal NPF at Bilos.

402



**Figure 3:** Half hourly evolution of m/z 135.11 ( $C_{10}H_{14}$ ), m/z 139.11 ( $C_9H_{14}O$ , nopinone), m/z 151.10 ( $C_{10}H_{14}O$ ), m/z 155.13 ( $C_{10}H_{18}O$ ), m/z 169.12 ( $C_{10}H_{16}O_2$ , pinonaldehyde) and their sum. Y axes are unit less as signals (number of counts) have been normalized to their maximum. Missing data are due to calibrations and blanks of the instrument. The blue shaded areas denote the nighttime periods (defined as the time when global radiation was below  $50 \text{ W m}^{-2}$ ).

403

### 3.6 Nighttime reactivity of BVOC oxidation products

404

In order to go further in the understanding of the role of oxidation products observed in the forest, it was decided to particularly focus on a short period, on m/z 139.11 and m/z 169.12

405

signals, corresponding to nopinone and pinonaldehyde respectively (**Figure 4**). This is of

406

particular interest as they are directly related to  $\alpha$ -pinene (for pinonaldehyde) and  $\beta$ -pinene

407

408 (for nopinone) oxidation. To the best of our knowledge, no direct comparison of diurnal

409 behaviours of both nopinone and pinonaldehyde from online field measurements has ever  
410 been reported. The **Figure 4** presents the temporal evolution of nopinone, pinonaldehyde,  
411 total monoterpenes and ozone mixing ratios during two nights (12/13 and 13/14 July 2015),  
412 representative of the entire campaign. It can be observed that during both nights,  
413 pinonaldehyde signal first increased faster than that of nopinone, reaching its maximum  
414 within a few hours and then decreased throughout the night. At the end of the night, a new  
415 increase was observed for pinonaldehyde mixing ratios, which finally returned to its daily  
416 level in the morning. The nocturnal dynamics of nopinone was observed to be simpler.  
417 Nopinone level started to increase at the sunset and decreased in the morning as turbulence  
418 reappeared. Such a strong difference between the nocturnal profiles of these two  
419 compounds probably results from their different physico-chemical properties and related  
420 reactivity. The formation yields of pinonaldehyde and nopinone from ozonolysis of  $\alpha$ - and  $\beta$ -  
421 pinene have been reported to be highly dependent on the physico-chemical conditions (Lee  
422 et al., 2006). More, the range of formation yields for both compounds are quite similar and  
423 very large (**Table 2**). If formation yields did not directly allow elucidating the faster increase of  
424 pinonaldehyde, the higher reaction rate constant of  $\alpha$ -pinene ozonolysis (compared to  $\beta$ -  
425 pinene) may at least partially explain such trend.

**Table 2:** Formation yields of nopinone and pinonaldehyde from  $\beta$ - and  $\alpha$ -pinene ozonolysis, reaction rate constants with the main atmospheric oxidants OH, O<sub>3</sub> and NO<sub>3</sub> and saturation vapour pressures (P<sub>s</sub> in Torr).

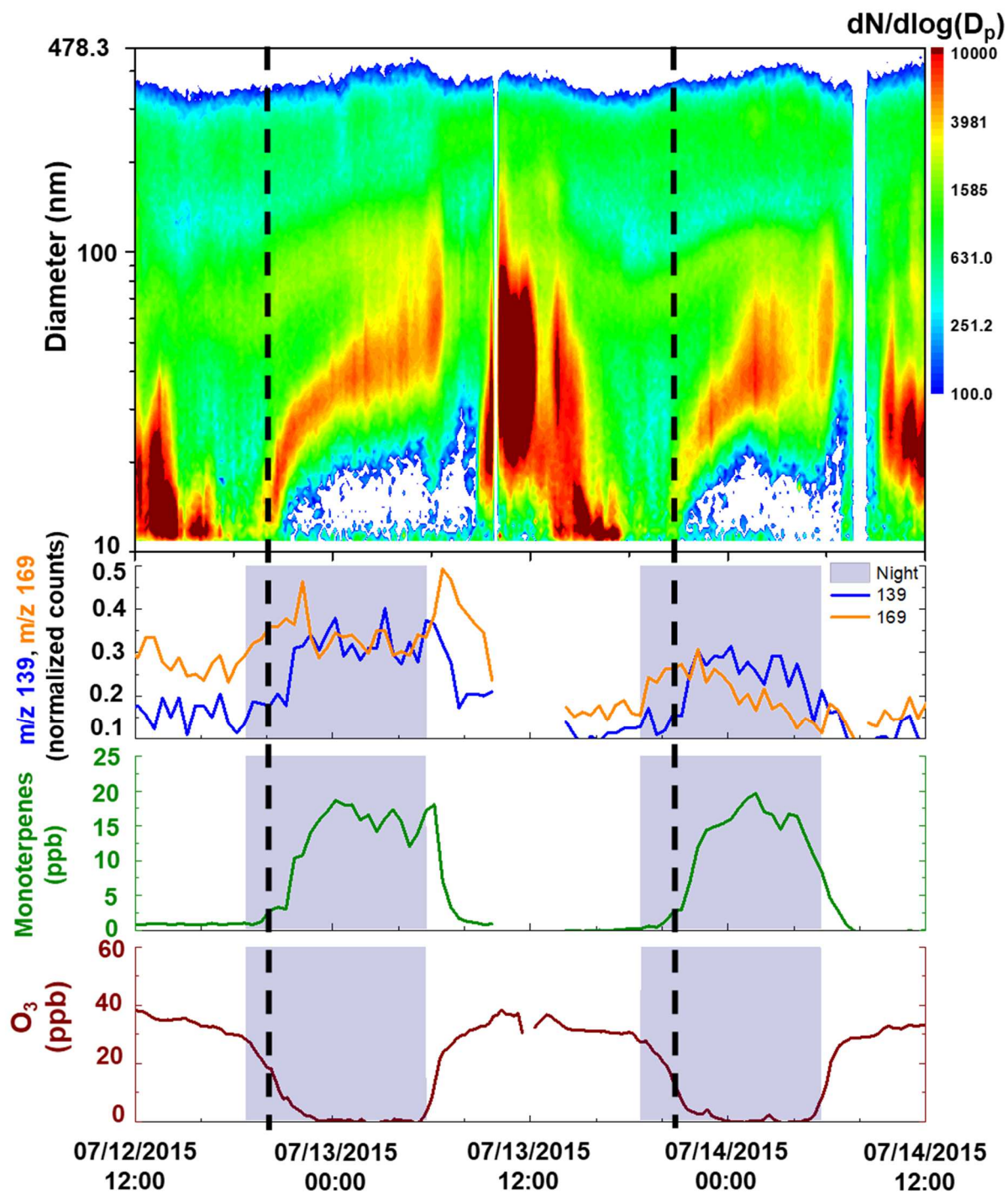
	Compounds	
	Nopinone	Pinonaldehyde
Formation yields (%)	16-40 <sup>a</sup>	6-53 <sup>a</sup>
k <sub>OH</sub> (cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup> )	(1.7±0.2) × 10 <sup>-11</sup> <sup>b</sup>	(3.9±0.2) × 10 <sup>-11</sup> <sup>c</sup>
k <sub>O<sub>3</sub></sub> (cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup> )	< 5 × 10 <sup>-21</sup> <sup>b</sup>	< 2 × 10 <sup>-20</sup> <sup>c</sup>
k <sub>NO<sub>3</sub></sub> (cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup> )	(1.1±0.3) × 10 <sup>-15</sup> <sup>b</sup>	(2.0±0.3) × 10 <sup>-14</sup> <sup>c</sup>
P <sub>s</sub> at 298K (Torr)	0.4	0.038 <sup>d</sup>

<sup>a</sup> Lee *et al.* (2006) and references therein <sup>b</sup> Calogirou *et al.* (1999) <sup>c</sup>recommended IUPAC values (Atkinson *et al.*, 2006) <sup>d</sup>Hallquist *et al.* (2009)

426 The second difference between the two nighttime product time profiles was the decrease of  
 427 pinonaldehyde during the night, not (or less) observable in the case of nopinone, highlighting  
 428 a larger sink for pinonaldehyde than for nopinone (**Figure 4**), assumed to be due to its faster  
 429 reactions with atmospheric oxidants (**Table 2**). The reaction of pinonaldehyde with ozone is  
 430 known to be very slow, and may not be responsible of such a decrease. NO<sub>3</sub>-initiated  
 431 reaction might be an important removal process for pinonaldehyde but would have required  
 432 larger concentrations of nitrate radicals that those expected at the very low NO<sub>x</sub> levels  
 433 measured during the campaign (**Figures S3.a** and **S4.a**). In addition, NO<sub>3</sub> reaction rate  
 434 constants with  $\alpha$ - and  $\beta$ -pinene are at least 100 times faster than nopinone and  
 435 pinonaldehyde (**Table 1**). As a result, any NO<sub>3</sub> radicals present at Bilos would first react with  
 436 monoterpenes, not allowing to explain the decrease of pinonaldehyde.

437 Many studies have reported the secondary formation of OH radicals following the  
438 monoterpene ozonolysis, with yields up to 1 (Rickard *et al.*, 1999; Aschmann *et al.*, 2002;  
439 Fick *et al.*, 2002; Atkinson and Arey, 2003; Forester and Wells, 2011). Moreover, Faloon *et al.*  
440 *al.* (2001) reported unexpected high OH concentrations at a deciduous forest site during  
441 nighttime. Therefore, it is reasonable to hypothesize that the nighttime pinonaldehyde  
442 decrease was due to its reaction with OH (this reaction being 5 times faster than that of  
443 nopinone) (**Table 2**). Obviously, future measurement of OH radicals at the Bilos site will be  
444 required to deeply answer to this question.

445 Another hypothesis can be put forward to tentatively explain the different profiles of  
446 pinonaldehyde and nopinone: their potential condensation on the particle phase. Indeed, the  
447 saturation vapour pressures of the two compounds differ by one order of magnitude,  
448 pinonaldehyde being 10 times less volatile than nopinone at ambient temperature (**Table 3**).  
449 But Kavouras *et al.* (1999) have shown that both compounds are usually mostly present in  
450 the gas phase. As a result, gas/particle partitioning might only weakly impact their  
451 concentration in the gas phase. Lastly, heterogeneous reactions (as for example, reactions  
452 with amines) could not be excluded from this discussion, and may play an important role in  
453 the fate of both products (Duporté *et al.*, 2016), but the investigation of such reactions was  
454 beyond the scope of the present study and might be considered in the further measurement  
455 campaigns at Bilos.



**Figure 4:** Size distribution of particles between 10 nm and 478.3 nm, during two typical NPF events observed at Bilos in July 2015. On lower panels: the half hourly evolution of m/z 139.21 ( $C_9H_{14}O$ , nopinone) and m/z 169.24 ( $C_{10}H_{16}O_2$ , pinonaldehyde) normalized signals, total monoterpene and ozone mixing ratios are represented. Intensities of oxidation product signals (number of counts) have been normalized to their maximum. Blue shaded areas denote the nighttime periods (defined as the time when global radiation was below  $50 \text{ W}\cdot\text{m}^{-2}$ ).



### 457           **3.7       Contribution of BVOC oxidation products to NPF**

458   This work raises the question of the link between the in-canopy reactivity of monoterpenes  
459   and the NPF events observed at night. In a recent paper (Kammer et al., 2018), we reported  
460   that NPF always occurred *i)* when the condensation sink was low, *ii)* after the nocturnal  
461   boundary layer was taking hold and *iii)* for the nights where monoterpene mixing ratios were  
462   high, suggesting their contribution to nocturnal NPF. In the **section 3.5**, we relate that the  
463   mixing ratios of monoterpene oxidation products increased at night. The **Figure 4** presents  
464   two consecutive nights (12/13 and 13/14 July 2015) where NPF occurred, as a typical  
465   example of what was observed during the campaign. At night, the sum of monoterpene  
466   mixing ratios started to increase about 1-2 hours before NPF took place, at the same time as  
467   nopinone and pinonaldehyde. This time might corresponded to a switch from condensation to  
468   nucleation, resulting from the drop of the condensation sink and the higher abundance of low  
469   volatile intermediates. To go further, a better knowledge of nanometer particle formation and  
470   size-resolved chemical composition of aerosols will be necessary. **Figure 4** shows that  
471   monoterpenes and their oxidation products levels increased simultaneously, which means  
472   that the main drivers that are triggering NPF are probably physical. However, the time lag  
473   between the simultaneous increase and the beginning of the NPF event may be due to *i)* the  
474   inability of the SMPS to catch the first steps of NPF and *ii)* the required time for condensable  
475   vapors to reach their saturation limit. This last hypothesis suggests a fast oxidation process,  
476   able to generate very low volatile gases in a short time. Auto-oxidation process has been  
477   shown to be able to produce highly oxidized multifunctional organic compounds (HOMs) very  
478   fast, which can then promote particle nucleation and growth (Jokinen et al., 2014; Kirkby et  
479   al., 2016; Tröstl et al., 2016). Hence, simultaneously to nopinone and pinonaldehyde  
480   generations, HOMs may have been produced by fast auto-oxidation processes, participating  
481   *in fine* to nocturnal NPF (Jokinen et al., 2014). In addition, OH radicals were very probably  
482   produced from the ozonolysis of monoterpenes during the night. This is supported by the  
483   time evolution of pinonaldehyde, which decreased in the middle of the night, probably due to

484 its reaction with OH. Thus, secondary reactions involving OVOCs might also be involved in  
485 NPF. The lag time described above might also be the time required for these secondary  
486 reactions to take place. But, even if the reactions between OH and OVOCs could be  
487 important for NPF at Bilos, the main source of OH during nighttime was arising from the  
488 primary ozonolysis of monoterpenes, highlighting the importance this process as an initial  
489 step. To go further, the contribution of each oxidation pathways (*i.e.* ozone and OH) to NPF  
490 will need more investigations. Considering that most of NPF events occurred during  
491 nighttime and the very low SO<sub>2</sub> mixing ratios observed at Bilos, it was not expected that  
492 sulfuric acid was involved, emphasizing again the role of BVOCs. The recent studies about  
493 NPF from pure BVOC ozonolysis, and the high monoterpene mixing ratios at our site let us  
494 hypothesise that ozonolysis of monoterpenes - and potentially other BVOCs - may be one of  
495 the main drivers of nocturnal NPF at the Bilos site (Kirkby *et al.*, 2016; Tröstl *et al.*, 2016).

### 496 **3.8 Molecular tracers of BVOC oxidation in the particle phase**

497 To support our hypothesis about the contribution of BVOCs to the nocturnal NPF, SOA  
498 tracers from  $\alpha$ - and  $\beta$ -pinene oxidation were investigated in the PM<sub>2.5</sub> particulate fraction  
499 collected during the 2015 field campaign. Even if the chemical composition of PM<sub>2.5</sub> and  
500 newly formed particle could be different, it is well-recognized that pinic and pinonic acids are  
501 good tracers for SOA formed from  $\alpha$ - and  $\beta$ -pinene ozonolysis (Claeys *et al.*, 2009; Faiola *et al.*,  
502 2014; Hallquist *et al.*, 2009; Jaoui *et al.*, 2005; Kristensen *et al.*, 2016; Nozière *et al.*,  
503 2011; Yasmeen *et al.*, 2011). Thus, the presence of such markers in PM<sub>2.5</sub> would support our  
504 hypothesis. During July 2015, pinic and pinonic acids were detected and quantified in all  
505 samples. Both products were found in large quantities in the particles collected at Bilos, in  
506 every 24 hours filters. Mean concentration of pinic acid was 18.3 ng m<sup>-3</sup>, and reached the  
507 maximum value of 33.8 ng m<sup>-3</sup> (**Table 3**). These values are in the same order of magnitude  
508 as those previously reported in studies conducted in other rural forested areas around the  
509 world. For example, the concentration of pinic acid measured in a boreal forest (Hyytiälä,  
510 Finland) during summertime was 7.2 ng m<sup>-3</sup>, and reached a maximal concentration of 31 ng

511 m<sup>-3</sup>. Pinic acid concentrations reported at other sites (**Table 3**) were reported to be between 4  
512 and 8 ng m<sup>-3</sup>, slightly lower than those measured at Bilos. This is consistent with the high  
513 monoterpene mixing ratios presented above. The mean concentrations of pinonic acid during  
514 the campaign was 10.2 ng m<sup>-3</sup>, also close to the reported values in **Table 3**. The high  
515 concentrations of these markers arising from  $\alpha$ - and  $\beta$ -pinene oxidation measured in this  
516 work support the high reactivity between monoterpenes and ozone, which is an important  
517 oxidation pathway for monoterpenes during nighttime at the Bilos site. This result attests that  
518 oxidation products of pinenes are well transferred to the particle phase, resulting from either  
519 gas/particle transfer during NPF or condensation. Further measurements of the size-resolved  
520 chemical composition of aerosols will be necessary to differentiate these processes.

521 In the present study, the concentrations of markers in the particle phase were in the same  
522 range as the reported values (or around twice as high), whereas monoterpene mixing ratios  
523 were at least one order of magnitude higher (**Table 3**). Such differences in the ratio between  
524 particulate SOA tracers concentrations and gaseous monoterpene mixing ratios suggest that  
525 different physico-chemical processes may occur at our site.

526 To further investigate the link between in-canopy monoterpene reactivity and nocturnal NPF,  
527 night and day filter sampling should be performed. Mochizuki *et al.* (2014) observed strong  
528 differences in the concentration of night and day particulate SOA tracers. In the case of  
529 nocturnal NPF identified in this study, such information could help to unravell the detailed  
530 processes. the This will be considered in a next field campaign implemented at Bilos. Note  
531 that a large panel of other biogenic SOA tracers were detected in the particles collected. The  
532 detailed chemical composition of atmospheric particles collected during the campaign will be  
533 the subject of a future publication.

534

535

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**Table 3: Comparison of pinic and pinonic acid concentrations measured at the Bilos site in**

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**July 2015 and those reported in the literature**


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Field site	Pinic acid		Pinonic acid		Monoterpenes	Ref.
	Mean	Min – Max	Mean	Min - Max	Mixing ratios (ppb)	
<b>Concentrations (ng m<sup>-3</sup>)</b>						
<b>Landes forest (Bilos, France) July 2015</b>	18.3	4.3 – 33.8	10.2	2.2 – 31.7	10 – 30	This work
<b>Boreal forest (Hyytiälä, Finland) July/August 2010/2011</b>	7.2	0.64 – 31	10.7	0.1 - 80	<1	(Vestenius et al., 2014)
<b>Deciduous forest (Jülich, Germany) July 2003</b>	4.2	0.9 – 9.9	-	-	-	(Kourtchev et al., 2014)
<b>Pine plantation (California, USA) Sept 2007 July 2009</b>	7.6 7.1	- -	6.1 11	- -	0.7 - 2.1	(Kristensen et al., 2013)
<b>Larch forest (Fujiyoshida, Japan) July 2012</b>	Day Night	0 – 18.2 0.8 – 14.5	12.7 3.1	1.7 – 28 0.9 – 6.8	0.5 - 4.4	(Mochizuki et al., 2014)

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## 536 Conclusions

537 Two intensive field measurement campaigns were conducted in the French Landes forest, at  
538 Bilos, in the summers of 2014 and 2015. Frequent nocturnal NPF events were observed  
539 during periods of strong thermic and/or hydric stress (dry and hot summer). In parallel,  
540 monoterpene mixing ratios (dominated by  $\alpha$ - and  $\beta$ -pinene) reached very high levels almost  
541 every night, frequently over 20 ppb in summer 2015. A strong anti-correlation between ozone  
542 and monoterpene mixing ratios was highlighted, suggesting a high chemical reactivity. This  
543 hypothesis was reinforced by the mean diurnal variation of the ratio between  $\alpha$ - and  $\beta$ -pinene  
544 mixing ratios. Products from the oxidation of  $\alpha$ - and  $\beta$ -pinene, namely pinonaldehyde and  
545 nopinone, were measured in the gas phase. The diurnal profiles of these two oxidation  
546 products is consistent with the high reactivity between  $\alpha$ - and  $\beta$ -pinene mostly with ozone in  
547 the evening. Furthermore, pinonaldehyde and nopinone presented different nocturnal

548 profiles, suggesting a possible secondary source of OH radicals from monoterpene  
549 ozonolysis during the night. Further work will be necessary to address this issue. The particle  
550 phase analysis allowed to identify and quantify pinic and pinonic acids, confirming the  
551 oxidation of monoterpenes observed at Bilos as well as the contribution of these precursors  
552 to SOA. Considering the differences between 2014 and 2015 field campaigns in the  
553 frequency of occurrence of NPF, the monoterpene mixing ratios, and the observed  
554 monoterpene in-canopy reactivity, monoterpene ozonolysis is proposed to very likely  
555 contribute to nocturnal NPF at our site. Measurements of size-resolved aerosol composition  
556 will be required to determine more specifically to which extent monoterpene oxidation  
557 products are involved in nucleation and/or aerosol growth in the Landes pine tree forest.

558 The high NPF frequency of occurrence observed in 2015 may be associated to the direct  
559 consequence of the hydric/thermic stress, which promotes strong BVOC emissions.  
560 Considering that the number of hydric stress periods will continue to increase due to the  
561 global warming, it can be hypothesized that NPF events may be more and more important in  
562 southwestern France. More studies will thus be required to confirm the effect of hydric stress  
563 on NPF.

564

## 565 **Acknowledgements**

566 This work was supported by the French Environment and Energy Management Agency  
567 (ADEME), the CNRS-INSU LEFE-CHAT research program, the NEEDS action, and the  
568 PEPS IdEx Bordeaux. The authors want to acknowledge the Bilos ICOS team (Christophe  
569 Chipeaux, Sébastien Lafont and Denis Loustau) for meteorological data and site availability  
570 and Michael Staudt (CEFE-CNRS) for the loan of the GC-FID during the campaign. We also  
571 thank Didier Garrigou (ISPA-INRA) for his precious help during the field campaigns,  
572 especially in the installation and the maintenance of instruments.

573 **Competing interests:** The authors declare no competing of interest.

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