

1 **Organic peroxides gas-particle partitioning and rapid**
2 **heterogeneous decomposition on secondary organic**
3 **aerosol**

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11 **Abstract**

12 Organic peroxides, important species in the atmosphere, promote secondary organic aerosols
13 (SOA) aging, affect HO_x radicals cycling, and cause adverse health effects. However, the
14 formation, gas-particle partitioning, and evolution of organic peroxides are complicated and
15 still unclear. In this study, we investigated in the laboratory the production and gas-particle
16 partitioning of peroxides from the ozonolysis of α -pinene, which is one of the major biogenic
17 volatile organic compounds in the atmosphere and an important precursor for SOA at a global
18 scale. We have determined the molar yields of hydrogen peroxide (H₂O₂), hydromethyl
19 hydroperoxide (HMHP), peroxyformic acid (PFA), peroxyacetic acid (PAA) and total
20 peroxides (TPO, including unknown peroxides) and the fraction of peroxides in α -pinene/O₃
21 SOA. Comparing the gas-phase with the particle-phase peroxides, we find that gas-particle
22 partitioning coefficients of PFA and PAA are 10⁴ times higher than the values from theoretical
23 prediction, indicating that organic peroxides play a more important role in the SOA formation
24 than previously expected. Here, the partitioning coefficients of TPO were determined to be as
25 high as $(2-3) \times 10^{-4} \text{ m}^3 \mu\text{g}^{-1}$. Even so, more than 80% of the peroxides formed in the reaction

26 remain in the gas phase. Water changes the distribution of gaseous peroxides, while it does not
27 affect the total amount of peroxides in either the gas or the particle phase. Approx. 18% of
28 gaseous peroxides undergo rapid heterogeneous decomposition on SOA particles in the
29 presence of water vapor, resulting in the additional production of H₂O₂. This process can
30 partially explain the unexpectedly high H₂O₂ yields under wet conditions. Transformation of
31 organic peroxides to H₂O₂ also preserves OH in the atmosphere, helping to improve the
32 understanding of OH cycling.

33 **1 Introduction**

34 Organic peroxides are important trace components in the atmosphere, serving as reservoirs of
35 HO_x and RO_x radicals, participating in the formation of secondary organic aerosols (SOA), and
36 causing adverse health effects as reactive oxygen species (ROS). Recently, peroxides are found
37 to play a key role in the aging of SOA. The particle-bound organic peroxides undergo
38 atmospheric photolysis with a lifetime of about 6 days (Epstein et al., 2014), and decline
39 significantly within the mean SOA age of 4–7 days (Rudich et al., 2007). A laboratory
40 experiment on the photolysis of SOA shows a high yield of hydroxyl radicals (OH), which are
41 considered to form from the decomposition of peroxides (Badali et al., 2015). This OH may
42 cause the in-particle oxidation of SOA.

43 Model studies have tried to simulate the SOA formation in chamber experiments, but great
44 discrepancies still exist between predicted and observed results (Camredon et al., 2010;
45 Hoffmann et al., 1997; Griffin et al., 1999; Cocker III et al., 2001; Saathoff et al., 2009; Presto
46 et al., 2005; Pye and Seinfeld, 2010; Farina et al., 2010). Jenkin (2004) added the formation of
47 dimers and improved the simulation, especially at the beginning of SOA formation. Organic
48 peroxides were found to be highly abundant in SOA (Ziemann, 2005; Docherty et al., 2005;
49 Surratt et al., 2006; Nguyen et al., 2010; Bateman et al., 2011; Mertes et al., 2012; Epstein et
50 al., 2014), possibly in the form of oligomers, which are even more important than carboxylic
51 acids (Bonn et al., 2004). In order to improve the simulation of the production of the SOA mass
52 within the chamber, explicit parameters about gas-particle partitioning of organic peroxides are
53 urgently needed.

54 The reactions and processes that generate or remove peroxides have been studied for many
55 years. Cross-reactions of organic peroxy radicals (RO₂) and the hydroperoxy radical (HO₂) and

56 self-reactions of HO₂ are thought to be major sources of organic peroxides and hydrogen
57 peroxide (H₂O₂), respectively, in the atmosphere. Ozonolysis of biogenic volatile organic
58 compounds (VOC) also produces H₂O₂ in high yields although its mechanism is unknown
59 (Zhang et al., 2009; Huang et al., 2013). Hydrolysis and reaction with OH are the main removal
60 pathways for both organic peroxides and H₂O₂, and dry/wet deposition removes only a small
61 portion of peroxides (Khan et al., 2015). However, existing theories about sources of and
62 removal of peroxides cannot account for the field observation results. Model simulations
63 showed an overestimation on total peroxides (TPO) and a underestimation on H₂O₂ as
64 compared with field records in the airborne GABRIEL (Guyanas Atmosphere-Biosphere
65 exchange and Radicals Intensive Experiment with the Learjet) field campaign (Kubistin et al.,
66 2010), indicating the existence of possible underestimated or new removal paths for organic
67 peroxides and overestimated or new formation paths for H₂O₂. Field observations and
68 laboratory experiments showed that particulate components, possibly particle-bound organic
69 peroxides, could be transformed to H₂O₂. Arellanes et al. (2006) found that H₂O₂ in ambient
70 SOA solution was 200–1000 times greater than expected levels based on the gas-liquid
71 partitioning, implying that almost all H₂O₂ is generated from SOA solution. Wang et al. (2011)
72 investigated several kinds of SOA derived from the oxidation of α -pinene, β -pinene, and
73 toluene, and came to the similar conclusion that more than 97.5% H₂O₂ arose from SOA
74 formation rather than from gas-liquid partitioning. However, this process happens in SOA
75 solution and the amount of H₂O₂ produced by such a pathway is too small to account for the
76 large discrepancy between observations and simulations for the gas-phase H₂O₂.

77 The effect of water on peroxides is complex. Laboratory experiments suggested that yields of
78 particle-phase total peroxides in the ozonolysis of alkenes are not influenced by water vapor
79 (Docherty et al., 2005). Unlike the total peroxides, yields of individual peroxides depend on
80 relative humidity (RH). Yield of H₂O₂ increases under wet conditions (Becker et al., 1990;
81 Hewitt and Kok, 1991; Simonaitis et al., 1991; G ěb et al., 1995; Huang et al., 2013), while the
82 yields of bis-hydroxymethyl hydroperoxide and three unknown organic peroxides decrease
83 under wet conditions (Huang et al., 2013). Theoretical studies suggest that water helps both the
84 formation and decomposition of organic peroxides. Water can react with stabilized Criegee
85 Intermediates (SCIs) and generate hydroxyalkyl hydroperoxides (HAHPs). It has been
86 proposed that not only isolated water molecules, but also water dimers react with SCI, and the

87 latter path could even be more important (Ryzhkov and Ariya, 2004). Numerous laboratory
88 experiments support this proposal (Chao et al., 2015; Lewis et al., 2015; Berndt et al., 2014).
89 As a result, the reaction with water dimers will be the largest sink for CH₂OO. However, the
90 quantum-chemical calculations predict that the larger SCIs react more slowly with water, both
91 the water monomer and dimer (Vereecken et al., 2014). Water also helps gas-phase
92 decomposition of HAHPs, although the decomposition rate constant is small according to the
93 theoretical calculations (Crehuet et al., 2001; Aplincourt and Anglada, 2003).

94 This study investigates the ozonolysis of α -pinene, which is considered as one of the largest
95 contributors to SOA and a dominant source of organic peroxides on a global scale (Khan et al.,
96 2015), focusing on the formation of peroxides in both the gas and the particle phase. Gas-
97 particle partitioning and water effect are examined carefully.

98 **2 Material and methods**

99 **2.1 Chemicals**

100 α -Pinene (Aldrich, 99%), cyclohexane (Sigma-Aldrich, $\geq 99.7\%$), potassium iodide (Alfa
101 Aesar, 99.9%), hydrogen peroxide (Alfa Aesar, 35 wt %), *ortho*-phosphoric acid (Fluka,
102 85–90%), hemin (Sigma, $\geq 98.0\%$), 4-hydroxyphenylacetic acid (Alfa Aesar, 99%), ammonia
103 solution (NH₃·H₂O, Beijing Tongguang Fine Chemicals Company, 25.0–28.0%), ammonium
104 chloride (NH₄Cl, Beijing Chemical Works, $\geq 99.5\%$), ultrapure water (18 M Ω , Millipore), N₂
105 ($\geq 99.999\%$, Beijing Haikeyuanchang Practical Gas Company Limited, Beijing, China), O₂ (\geq
106 99.999%, Beijing Haikeyuanchang Practical Gas Company Limited, Beijing, China), and
107 polytetrafluoroethylene (PTFE) filter membrane (Whatman Inc., 47 mm in diameter) were
108 used in this study.

109 **2.2 Apparatus and procedures**

110 A flow tube reactor (2 m length, 70 mm inner diameter, quartz wall) equipped with a water
111 jacket for controlling temperature was used to investigate the ozonolysis of α -pinene. All the

112 experiments were conducted at 298 ± 0.5 K and in dark. O_3 was generated by the photolysis of
113 O_2 in a 2 L quartz tube with low-pressure Hg lamp, and the detailed quantification method of
114 O_3 was described in our previous study (Chen et al., 2008). O_3 (~25 ppmv) was used in the
115 experiments. α -Pinene gas was generated by passing a flow of N_2 over liquid α -pinene in a
116 diffusion tube at the selected controlled temperature. The initial concentration of α -pinene,
117 determined by gas-chromatography-flame ionization detector (GC-FID, Agilent 7890A, USA),
118 was ~273 ppbv in the experiments. Water vapor was generated by passing N_2 through a water
119 bubbler. The mixing gases, including α -pinene, O_3 , and dry or wet synthetic air (80% N_2 and
120 20% O_2), were continuously introduced into the reactor with a total flow rate of 4 standard L
121 min^{-1} (standard liters per minute) and a residence time of 120 s. The relative humidity (RH)
122 was controlled at two levels: <0.5% RH (dry conditions) and 60% RH (wet conditions). Gas
123 from the reactor (2 standard L min^{-1}) was directed into a coil collector and scrubbed by H_3PO_4
124 stripping solution (5×10^{-3} M, pH 3.5) for hydroperoxides analysis. SOA produced from the
125 ozonolysis of α -pinene were collected onto a PTFE filter for 4 h at a flow rate of 4 standard L
126 min^{-1} , and the mass of SOA on the filter was immediately measured by a semi-micro balance
127 (Sartorius, Germany). After that, each loaded filter was extracted with 20 mL H_3PO_4 solution
128 (5×10^{-3} M, pH 3.5) using a shaker (Shanghai Zhicheng ZWY 103D, China) at 180 rpm and 4
129 $^\circ\text{C}$ for 15 min, and then the SOA solution was immediately analyzed to determine the particle-
130 phase peroxides. Each SOA solution was analyzed 7 times at different times to investigate the
131 evolution of SOA solution.

132 To explore the effect of water vapor on the formation of peroxides in the ozonolysis, two-stage
133 reaction experiments were designed and carried out. In the first stage, dry synthetic air (2
134 standard L min^{-1}) with α -pinene (~275 ppbv) and O_3 (~42 ppmv) entered the first 2 L flow-tube
135 reactor; in the second stage, the gas passed through the second 2 L flow-tube reactor but with
136 the addition of dry or wet synthetic air (2 standard L min^{-1}). The residence time was 68 s in the
137 first reactor and 34 s in the second reactor. The concentration of α -pinene at the outlet of the
138 first reactor was found to be below the GC-FID detection limit (< 5 ppbv), meaning that α -
139 pinene was almost completely consumed before the gas entered the second reactor. Thus, water
140 vapor appearing in the second reactor only affected the products from the first reactor. A filter
141 was placed at the outlet of the first reactor or second reactor to collect SOA when necessary.

142 2.3 Peroxides analysis

143 The low-weight molecular peroxides were measured using high-performance liquid
144 chromatography (HPLC, Agilent 1100, USA) coupled with a post-column derivatization
145 module and fluorescence detection, and the concentration of total peroxides was determined by
146 an iodometric spectrophotometer method. Details about the HPLC-fluorescence method have
147 been reported in our previous study (Hua et al., 2008). Briefly, this method is based on the
148 reaction of *p*-hydroxyphenylacetic acid (POPHA) with organic hydroperoxides or hydrogen
149 peroxide in the catalysis of the hemin forming POPHA dimer (2,2'-bisphenol-5,5'-diacetic acid),
150 which is a fluorescent substance, and then is quantified by fluorescence detector. The separation
151 of peroxides was implemented by column chromatography before the peroxides were reacted
152 with POPHA. The synthetic method for organic peroxides standards is described in our previous
153 study (Huang et al., 2013).

154 The iodometric spectrophotometric method is used to quantify all classes of peroxides (ROOR',
155 ROOH and H₂O₂), with the exception of tertiary dialkyl peroxides, in the aqueous phase without
156 distinction (Banerjee and Budke, 1964). Peroxyhemiacetals formed in α -pinene ozonolysis can
157 be measured with this method. Excess potassium iodide reacts with peroxides producing I₃⁻ ion
158 (R1), which can be quantified by UV/VIS spectrophotometry.



160 α -Pinene SOA is freely soluble in polar solvents, e.g., water, acetonitrile and methanol, but it
161 is poorly soluble in nonpolar solvents, e.g., chloroform and toluene (Nguyen et al., 2010).
162 Hence, a H₃PO₄ solution, as a kind of polar solvent, could entirely extract SOA from filters.
163 The HPLC fluorescence method uses H₃PO₄ solution as a solvent for peroxides such as H₂O₂,
164 hydromethyl hydroperoxide (HMHP), performic acid (PFA), and peracetic acid (PAA) which
165 are more stable in acidic solution than in pure water (Zhou and Lee, 1992). In order to be
166 comparable with the HPLC fluorescence method, SOA loaded filters were also extracted by
167 H₃PO₄ solution. The influence of pH on extraction efficiency is discussed in the Supplement.
168 In this study, SOA solution (2.5 mL) was added into a 10 mL airtight micro reaction vessel
169 (Supelco, USA). Each solution was then purged of oxygen by bubbling with N₂ for 5 min. After
170 purging, an aqueous solution of KI (250 μ L, 0.75 M) was added into the vessel. The vessel was
171 then capped tightly, covered with aluminium foil, and allowed to stand in the dark for 12–24 h.

172 The solution absorbance was then measured at 420 nm by an UV/VIS spectrophotometer
173 (SHIMADZU UV-1800, Japan). The efficiency of peroxide measurements was discussed in the
174 supplement.

175 **3 Results and discussion**

176 **3.1 Gas-particle partitioning of peroxides**

177 **3.1.1 Particle-phase peroxides**

178 We measured the different mass values of SOA produced from the ozonolysis of α -pinene at
179 different RHs in the presence or absence of the OH scavenger cyclohexane, and found that the
180 typical in-reactor SOA concentration was 450–650 $\mu\text{g m}^{-3}$. A comparison of the aerosol mass
181 yields (Y_{SOA}), defined as the ratio of the formed aerosol mass to the consumed α -pinene mass,
182 (Table 1) indicated that while the SOA yields were independent on the presence of water vapor,
183 they decreased in the presence of OH scavenger.

184 Organic peroxides are considered to be one of the major constituents in SOA (Docherty et al.,
185 2005; Ziemann, 2005; Surratt et al., 2006; Nguyen et al., 2010; Mertes et al., 2012; Kidd et al.,
186 2014; Badali et al., 2015) (Table S1). The sensitivity of the iodometric method to ROOR is
187 critical to obtain an accurate concentration of total peroxides since peroxyhemiacetals are a
188 significant component (Docherty et al., 2005). In the present study, we determined the total
189 molar concentration of peroxides in SOA using the iodometric method. Stability of peroxides
190 in SOA stored on-filter was also tested, and the results show that peroxides concentration
191 decrease with increasing sitting time (Figure S4). Hence, the peroxides in SOA were determined
192 immediately after collection. Here, the mass fraction of peroxides in SOA ($F_{\text{peroxides}}$) is defined
193 as the ratio of mass of particle-bound peroxides to SOA mass, which is defined as follows:

$$194 \quad F_{\text{peroxides}} = \frac{m_{\text{peroxides}}}{m_{\text{SOA}}} \quad (1)$$

195 where $m_{\text{peroxides}}$ is the mass of particle-bound peroxides, such as PFA, PAA and TPO, and m_{SOA}
196 is the mass of SOA. Assuming that the average molecular weight of peroxides is 300, we
197 obtained the mass fraction of total peroxides in SOA (F_{TPO}) as ~0.21 (Table 1), which is

198 consistent with 0.22 reported by Epstein et al. (2014), but less than 0.47 reported by Docherty
199 et al. (2005) and 0.34 reported by Mertes et al. (2012). Several factors, such as the presence of
200 OH scavengers, reactor type, extraction method, SOA mass measurements, and SOA density
201 assumptions, may cause these discrepancies. In addition to the concentration of total peroxides,
202 we measured the concentration of two small organic peroxides peroxyformic acid (PFA) and
203 peroxyacetic acid (PAA) in SOA and calculated the contribution of PFA (F_{PFA}) and PAA (F_{PAA})
204 to SOA mass (Table 1). Under dry conditions, the F_{PFA} and F_{PAA} were 0.35 ± 0.06 and 0.11 ± 0.04
205 $\text{ng } \mu\text{g}^{-1}$, respectively, without the OH scavengers and 0.14 ± 0.00 and 0.09 ± 0.01 $\text{ng } \mu\text{g}^{-1}$,
206 respectively, with cyclohexane. After adding water vapor, F_{PFA} did not significantly change,
207 but F_{PAA} approached 0.

208 3.1.2 Gas-phase peroxides

209 In addition to the particle phase peroxides, we measured the gas-phase peroxides generated in
210 the ozonolysis of α -pinene. Here, the molar yield of gaseous peroxides ($Y_{\text{peroxides}}$) is defined in
211 eq (2):

$$212 \quad Y_{\text{peroxides}} = \frac{\Delta_{\text{peroxides}}}{\Delta_{\alpha\text{-pinene}}} \quad (2)$$

213 where $\Delta_{\text{peroxides}}$ are moles of formed gaseous peroxides, such as HMHP, PFA, PAA and TPO,
214 and $\Delta_{\alpha\text{-pinene}}$ are moles of consumed α -pinene. The molar yield of total peroxides (Y_{TPO}) were
215 estimated to be nearly the same under both dry conditions and wet conditions in the absence of
216 OH scavengers (Table 1), indicating that total yield of peroxides was unaffected by water vapor.
217 Moreover, when we employed the MCM v3.1 mechanism to simulate the present reaction
218 system, the modelled yield of total peroxides was about 0.25, consistent with our experimental
219 result. The model results also suggested that hydroperoxides account for more than 99% of total
220 peroxides. The yields of HMHP (Y_{HMHP}), PFA (Y_{PFA}) and PAA (Y_{PAA}) are shown in Table 1.
221 Compared with dry conditions, Y_{HMHP} and Y_{PFA} doubled under wet conditions, while Y_{PAA}
222 increased only slightly. However, yields of these three organic peroxides were all lower in the
223 presence of OH scavengers, indicating the importance of OH in the formation of small organic
224 peroxides.

225 Considering that all the peroxides originally existed in the gas phase at the beginning of the
 226 ozonolysis of α -pinene, we estimated the fraction of peroxides that entered the particulate phase
 227 from the gas phase through gas-particle partition based on measured peroxides in the particle
 228 and gas phases. The fraction of particulate TPO in gaseous and particulate TPO
 229 (TPO(p)/TPO(g+p)) were essentially the same (Table 1) under both wet and dry conditions. To
 230 the best of our knowledge, for the ozonolysis of α -pinene, this is the first report of the yield of
 231 gas-phase total peroxides (including hydrogen peroxide and organic peroxides) and the gas-
 232 particle partitioning fraction.

233 The gas-particle partitioning coefficient (K_p) describes the partitioning ability of a given species,
 234 calculated as follows (Odum et al., 1996):

$$235 \quad K_p = \frac{C_a}{C_g C_{om}} \quad (3)$$

236 where C_a is the concentration of this species in the aerosol phase, $\mu\text{g m}^{-3}$; C_g is the concentration
 237 of this species in the gas phase, $\mu\text{g m}^{-3}$; and C_{om} is the total concentration of condensed organic
 238 matter, $\mu\text{g m}^{-3}$. Based on the gas-phase peroxides concentration, particle-phase peroxides
 239 concentration, and the aerosol yields summarized in Table 1, we can obtain the observed K_p
 240 (Table 2).

241 The Pankow absorption model (Pankow, 1994) is the most widely accepted mechanism to
 242 explain the gas-particle partitioning, and has been used to predict aerosol yields in chamber
 243 experiments (Cocker III et al., 2001; Jenkin, 2004; Yu et al., 1999). Theoretical K_p can be
 244 calculated by the following equation:

$$245 \quad K_p = \frac{7.501 \times 10^{-9} RT}{MW_{om} \zeta p_L^\circ} \quad (4)$$

246 where R is the ideal gas constant, $\text{J K}^{-1} \text{mol}^{-1}$; T is the temperature, K ; MW_{om} is the mean
 247 molecular weight of the condensed organic material, g mol^{-1} . In the present study, MW_{om} is
 248 estimated to be 130 g mol^{-1} ; ζ is the activity coefficient of the given species in the condensed
 249 organic phase, and here, is assumed to be unity; p_L° is the liquid vapour pressure of this species,
 250 Torr. The theoretical p_L° can be calculated by an expanded, semi-empirical form of Clausius-

251 Clapeyron equation (Baum, 1997). Theoretical gas-particle partitioning coefficients of PFA and
252 PAA are shown in Table 2.

253 The observed gas-particle partitioning coefficients of PFA, PAA and TPO were $(3-9)\times 10^{-5}$, $(2-$
254 $4)\times 10^{-5}$ and $(2-3)\times 10^{-4} \text{ m}^3 \mu\text{g}^{-1}$, respectively (Table 2), which, to the best of our knowledge,
255 are reported here for the first time. However, the long time collection for SOA do have effects
256 on gas- and particle-phase constituents possibly due to re-partitioning of species between the
257 two phases. Collected SOA mass and peroxides amount per unit time decreases slightly with
258 increasing collection time, and peroxides amount decreases faster than SOA mass (Supplement).
259 Hence, the gas-particle partitioning coefficients of peroxides given here are underestimated by
260 about 21%. Compared with the observed K_p values, theoretical K_p values of PFA and PAA,
261 $2\times 10^{-9} \text{ m}^3 \mu\text{g}^{-1}$ and $4\times 10^{-9} \text{ m}^3 \mu\text{g}^{-1}$, respectively, were lower by a factor of 10^4 . This large
262 difference between observed and theoretical K_p values has also been reported previously
263 (Cocker III et al., 2001; Jenkin, 2004; Kamens and Jaoui, 2001). Jenkin (2004) considered the
264 existence of a significant systematic error, which is independent of key parameters involved in
265 prediction of K_p values, or the inability to interpret the Pankow absorption model. After
266 inducing a species-independent scaling factor of ca. 120 for all partitioning species, Jenkin
267 obtained a reasonable simulation of the final experimental aerosol concentration, but was still
268 unable to interpret the early stages of aerosol accumulation. In addition to the absorptive
269 partitioning mechanism, the participation of bi- and multifunctional acid dimers in the aerosol
270 formation process was also considered, resulting in the presentation of simulated results.
271 Organic peroxides are also important compounds in dimer formation, for instance,
272 hydroperoxides can react with aldehydes subsequently producing peroxyhemiacetals (Tobias
273 and Ziemann, 2000; Tobias and Ziemann, 2001; Ziemann, 2005). The vapor pressures of
274 hydroperoxides decreased in the formation of peroxyhemiacetal by an additional factor of $\sim 10^2-$
275 10^5 (Tobias and Ziemann, 2000), which could partially explain the large discrepancy between
276 the observed and theoretical gas-particle partitioning coefficient. However, related
277 thermodynamic and kinetic parameters need further study to resolve the problem.

278 **3.2 Evolution of SOA in the aqueous phase**

279 We investigated the evolution of SOA in the aqueous phase, focusing on the change of H_2O_2 in
280 SOA. The initial concentration of H_2O_2 was low, but it increased rapidly in the first 3.5 h,

281 approaching the peak at ~7.5 h, and then decreased slowly, meaning the existence of a sustained
282 release of H₂O₂ in the SOA solution at room temperature (298 K) (Figure 1). The molar fraction
283 of peak-H₂O₂ to total peroxides in SOA (H₂O₂(p)/TPO(p)) under dry conditions was twice as
284 high as that observed under wet conditions in the absence of OH scavengers (Table 1). The
285 contribution of H₂O₂ to the SOA mass ($F_{\text{H}_2\text{O}_2}$) was 1.9 times higher under dry conditions
286 ($5.09 \pm 0.99 \text{ ng } \mu\text{g}^{-1}$) than under wet conditions ($2.67 \pm 0.17 \text{ ng } \mu\text{g}^{-1}$) (Table 1). In the presence
287 of cyclohexane, H₂O₂(p)/TPO(p) was slightly higher under dry conditions than wet conditions
288 (0.09 ± 0.00 and 0.06 ± 0.00 , respectively) while $F_{\text{H}_2\text{O}_2}$ values were not significantly affected by
289 RH (Table 1). Our observations consist with those of Wang et al. (2011) who studied α -pinene-
290 derived SOA, and measured the $F_{\text{H}_2\text{O}_2}$ as $2.01 \pm 0.76 \text{ ng } \mu\text{g}^{-1}$, which was unchanged by a variety
291 of oxidants (NO/light, O₃ and O₃/cyclohexane) over the range of 14–43% RH.

292 The sustained release of H₂O₂ coupled with the attenuation of total peroxides provided
293 experimental evidence for the hypothesis that the decomposition/hydrolysis of organic
294 peroxides generates H₂O₂. The decay of H₂O₂ in SOA solution after 18 h is a comprehensive
295 phenomenon including formation and decomposition, and the rate was estimated to be 0.06 μM
296 h^{-1} and 0.03 $\mu\text{M } \text{h}^{-1}$ for SOA produced under dry and wet conditions, respectively. To assess
297 the formation, we determined the decomposition rates of pure H₂O₂ at different concentrations.
298 When the H₂O₂ concentration was ~5 μM (~2.5 μM), the rate of decomposition was 0.11 μM
299 h^{-1} (0.05 $\mu\text{M } \text{h}^{-1}$). The H₂O₂ concentrations were equivalent with those of H₂O₂ in SOA solution.
300 Thus, we can estimate H₂O₂ formation in the SOA solution after 18 h to have a rate of 0.05 μM
301 h^{-1} under dry conditions and 0.02 $\mu\text{M } \text{h}^{-1}$ under wet conditions. As shown in Figure 2, the
302 peroxy-carboxylic acids (PCAs) (PFA and PAA), decayed quickly while the HAHP (HMHP)
303 decayed slowly. Hence, the formation of H₂O₂ after 18 h could be attributed to the
304 decomposition of HAHPs. However, in the first 7.5 h period, H₂O₂ increased rapidly which is
305 more consistent with the decay of PCAs rather than HAHPs. Not all the organic peroxides
306 decayed during the observation time, since the attenuation of TPO almost stopped after 40 h.
307 The residual peroxides were more stable, possibly due to the formation of ROOR by
308 oligomerization.

309 3.3 Unexpectedly high levels of H₂O₂ in the gas phase

310 Table 1 shows the molar yields of gas-phase H₂O₂. The H₂O₂ yield in the absence and presence
311 of cyclohexane was essentially the same in dry conditions, but under conditions of high RH
312 increased to 0.16±0.01 in control studies and to 0.14±0.02 in the presence of cyclohexane. Thus,
313 the presence of water vapor elevated the H₂O₂ yield, while the presence of a radical scavenger
314 had no effect. Previous studies on gas-phase H₂O₂ yields of the ozonolysis of α -pinene are
315 reviewed (Table S2). Becker et al. (1990) firstly reported that the presence of water vapor will
316 significantly promote the H₂O₂ yield, and our work confirmed this observation. However, our
317 measured values of H₂O₂ were ten times higher than those reported by others under both dry
318 and wet conditions, except that by Simonaitis et al. (1991). Differences in reactant concentration,
319 reactor type and measuring methods account for these discrepancies. Worth noting, the
320 reactants concentrations used in these previous and our experiments are very high, therefore,
321 yields of peroxides may not represent actual yields of peroxides in oxidation of α -pinene in
322 nature (Supplement).

323 The source of gas-phase H₂O₂ remains unclear. We suggest that further ozonolysis and OH
324 oxidation of gaseous products and reactants in the aqueous phase during and after gas collection
325 are not likely to be the main sources of H₂O₂. In this study, the online GC-FID test showed that
326 α -pinene was completely consumed in the gas phase. Hence, the contribution of aqueous-phase
327 α -pinene ozonolysis to the measured H₂O₂ in the coil collector should be negligible. The main
328 gas-phase non-peroxy organic products of α -pinene ozonolysis are carbonyls and organic acids,
329 e.g., pinonaldehyde, formaldehyde, acetone and pinic acid; these compounds without carbon-
330 carbon double bonds cannot be oxidized by O₃. The ozonolysis of α -pinene produces the OH
331 radical in high yield (0.68–0.91) (Berndt et al., 2003), which potentially oxidizes carbonyls and
332 organics. However, we observed no difference of $Y_{\text{H}_2\text{O}_2}$ in the absence and presence of OH
333 scavengers, indicating that OH oxidation in the aqueous phase may not be a source of H₂O₂.

334 Decomposition/hydrolysis of organic peroxides in the aqueous phase during and after gas
335 collection are also found to be a minor source of gas-phase H₂O₂. HAHPs and PCAs, two kinds
336 of organic peroxides, are the probable candidate for generating H₂O₂. HAHPs are the main
337 products of the reaction of SCI with water molecules and dimers (Ryzhkov and Ariya, 2004),
338 and they can decompose to H₂O₂ plus the corresponding aldehyde or H₂O plus the

339 corresponding organic acid (Hellpointner and G äb, 1989). The hydrolysis of PCAs, which are
340 generated from the reaction of RC(O)OO with HO₂, is another possible source of H₂O₂. Several
341 kinds of PCAs have been qualitatively observed in the ozonolysis of α -pinene (Venkatachari
342 and Hopke, 2008). In this study, we quantitatively observed PFA and PAA in the gas phase
343 (Table 1), and simulated the formation of PCAs using MCM v3.1 mechanism. Model results
344 showed that the yield of total PCAs was extremely low, 0.0005, and PAA contributed more
345 than half of the yield while the formation pathway of PFA was not included. The large
346 discrepancy between modelled and experimental results indicates that PCAs play a more
347 important role than was expected previously. We estimate the H₂O₂ generated from organic
348 peroxides in the aqueous phase by measuring the decomposition/hydrolysis rate of organic
349 peroxides. Considering the effects of concentration, coexisting components, and ionic strength,
350 we conducted the measurements with coil collection solutions rather than with synthesized
351 samples. The decomposition/hydrolysis of organic peroxides is a pseudo-first order reaction
352 due to the excess of the other reactant, i.e., water. The decay rate constants of HMHP, PFA and
353 PAA were determined to be 0.09, 1.06 and 0.64 h⁻¹, respectively (Figure 2). Larger HAHPs
354 were less active compared with HMHP and should have lower decay rate constants. If all the
355 TPO are composed of HAHPs and the production of H₂O₂ plus aldehydes is the only
356 decomposition pathway of HAHPs, the upper bound of H₂O₂ formed in the aqueous phase
357 within 8 min may be estimated to be 1.2% of TPO. However, the observed ratio of gas-phase
358 H₂O₂ to TPO was 28%–78%, indicating that the aqueous-phase decomposition of HAHPs is
359 insignificant. Compared with HMHP, the decay rates of PFA and PAA were quite high.
360 Assuming that all the TPO, except for PFA, are PCAs and their decay rates were the same as
361 PAA's, H₂O₂ formed in aqueous phase within 8 min is estimated to be 13.2% of TPO, which
362 can partially explain the observed H₂O₂ level. However, the yield of PCAs in the ozonolysis of
363 α -pinene is predicted to be low in the MCM v3.1 model. The experimental results mentioned
364 above concluded that the aqueous-phase formation of H₂O₂ is not important, for the decay rate
365 of HAHPs was too slow, and the amount of PCAs was too low although their decay rate was
366 higher.

367 Whether the self-reaction of HO₂ and decomposition of HAHP in the gas phase are the main
368 sources of H₂O₂ is discussed here. The self-reaction of HO₂ is considered to be the main source
369 of ambient H₂O₂ (Lee et al., 2000; Reeves and Penkett, 2003) and occurs in the ozonolysis of

370 α -pinene. When we estimated the contribution of this pathway to the observed H₂O₂ from α -
371 pinene ozonolysis using MCM v3.1 mechanisms, the yield was less than 0.001 under both dry
372 and wet conditions, meaning that this pathway is negligible. Chamber experiments showed that
373 SCI mainly reacts with water vapor even under dry conditions (Jenkin, 2004), and the major
374 product is HAHP. Aplincourt and Anglada (2003) considered that the unimolecular
375 decomposition of gaseous HAHPs was unlikely to occur, and only the water-assisted
376 decomposition was efficient in the gas phase. They estimated the water-assisted decomposition
377 rate constant of 2-propenyl α -hydroxy hydroperoxide to be $1.5 \times 10^{-30} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ by
378 quantum chemical calculation. Based on their work, the gas-phase decomposition fraction of
379 HAHP in 2 min can be calculated to be less than 0.01%, which is too small to account for the
380 H₂O₂ observed in our experiments.

381 In summary, the high H₂O₂ yields in the gas phase cannot be explained by the bias caused by
382 measuring method and the current formation mechanism of H₂O₂. There may exist an unknown
383 or underestimated pathway producing H₂O₂. In section 3.4, we propose that gaseous organic
384 peroxides can undergo rapid heterogeneous decomposition in the presence of water and
385 producing H₂O₂.

386 **3.4 Rapid heterogeneous decomposition of gaseous organic peroxides**

387 Our results demonstrate that water vapor has no significant effect on either the yield of total
388 peroxides (combining gaseous and particulate peroxides) or the contribution of peroxides to
389 SOA mass. However, water vapor does change the concentrations of H₂O₂, PFA and PAA in
390 the gas phase and particle phase in an opposite manner (Table 1). In the presence of water vapor,
391 H₂O₂ yield increased dramatically by ~300%, and gas-phase H₂O₂/TPO increased from 0.26 to
392 0.78. Yields of HMHP, PFA and PAA also increased with the presence of water vapor. These
393 results clearly indicate that water vapor can change the formation and distribution of peroxides.

394 We carried out a series of two-stage experiments using two reactors under various scenarios to
395 further study the effect of water vapor on peroxides (Table 3). In scenario 1d, no water vapor
396 was added and a filter was used to intercept SOA entering the coil collector, which is similar to
397 measuring H₂O₂ under dry conditions with one reactor. The concentration of H₂O₂ observed in
398 the coil collection solution under this condition was considered to be the baseline value, 100%.

399 When the filter was placed at the outlet of the first reactor (scenario 2d) instead of at the second
400 reactor, the concentration of H₂O₂ was 103 ±6%, almost the same as the baseline, indicating that
401 the coexistence of gaseous products and SOA will not lead to the formation of H₂O₂. In scenario
402 2w, a filter was placed at the outlet of the first reactor and water vapor was added to the second
403 reactor, resulting in the coexistence of gaseous products and water vapor (50% RH) in the
404 second reactor. The concentration of H₂O₂ observed in this scenario was 87%, slightly lower
405 than the baseline, possibly due to losses on the wall of the reactor under wet conditions, which
406 has been reported to be 5% for H₂O₂ at 50% RH (Huang et al., 2013). When we maintained the
407 water vapor and moved the filter to the outlet of the second reactor (scenario 1w), the H₂O₂
408 concentration increased to 165 ±6% of baseline. In scenario 3w, with water vapor added to the
409 second reactor and without a filter in the gas flow, the H₂O₂ concentration was 172 ±5%, almost
410 the same as that in scenario 1w. In scenario 3d, no water vapour and no filter were used, but a
411 high H₂O₂ concentration, 164 ±9%, was also observed. For scenarios 1w, 3w, and 3d, where
412 H₂O₂ increased by ~67%, gaseous products, SOA, and water were all present in the second
413 reactor or coil collector. The coexistence of gaseous products and water vapor (see scenario 2d
414 and 2w), the coexistence of gaseous products and SOA (see scenario 1d and 2d), and the
415 coexistence of SOA and water (see section 3.3) did not result in a high yield of H₂O₂. We
416 therefore concluded that the presence of three components together, the gaseous products, SOA,
417 and water, was necessary for a high yield of H₂O₂. Once the gaseous products and SOA had
418 been in contact with water vapor in the second reactor, the levels of H₂O₂ were increased to the
419 same extent, whether or not these compounds were mixed with condensed water (see scenario
420 1w and 3w), indicating that the process producing H₂O₂ in the gas phase is quite rapid.

421 When we measured the total peroxides formed from gaseous products and SOA in scenarios 1d
422 and 1w, the results showed that for these two scenarios, the levels of the total peroxides in both
423 gaseous products and SOA were not significantly different, indicating that SOA does not
424 change in the presence of water vapor and no new peroxides formed in the gas phase. This
425 outcome supports the idea that the increment of H₂O₂ comes from the redistribution of gaseous
426 peroxides, which is induced by the heterogeneous decomposition of gaseous products in the
427 presence of both SOA and water. Based on the measured increment of H₂O₂ and concentration
428 of gaseous total peroxides, we concluded that 18% of the gaseous total peroxides undergo rapid
429 heterogeneous decomposition.

430 Heterogeneous reactions of trace gases on the surface of particles relevant to the atmosphere
431 have been studied for many years. The investigated trace gases, including nitrogen oxides (e.g.,
432 HNO₃, NO₂ and N₂O₅), SO₂, O₃, H₂O₂ and oxygenated VOCs (Liggio et al., 2005; Kroll et al.,
433 2005; Prince et al., 2007; Zhao et al., 2010; Zhao et al., 2011; Zhao et al., 2014; Huang et al.,
434 2015), could react with the active sites on the surfaces of mineral dust (Goodman et al., 2001;
435 Fu et al., 2007). Unlike mineral dust, however, SOA has no such active sites. The elucidation
436 of the mechanism of the rapid heterogeneous decomposition of organic peroxides on SOA
437 particles remains a great challenge and needs urgent study.

438 **4 Conclusions and atmospheric implications**

439 Our laboratory study has provided more evidence that organic peroxides are important
440 components of SOA derived from the ozonolysis of alkenes. In the case of α -pinene, organic
441 peroxides account for ~21% of the SOA mass and this fraction is not affected by RH and the
442 presence of OH scavengers. More interestingly, the gas-particle partitioning coefficients of
443 organic peroxides have been estimated for the first time based on the measurements of both
444 gaseous and particulate peroxides. Due to the long-time collection for SOA, these coefficients
445 reported here are lower bounds in this study. For PFA and PAA, the observed values were 10⁴
446 times higher than that of the theoretical value calculated by the Pankow absorption model. This
447 discrepancy indicates a more important role of peroxides in SOA formation than expected
448 previously and the existence of yet undefined mechanisms in addition to the absorption. The
449 reaction of organic hydroperoxides with carbonyls forming peroxyhemiacetals may explain part
450 of the enhancement of the partitioning of peroxides. However, the kinetic parameters of
451 peroxyhemiacetal formation are lacking. The explicit mechanisms of gas-particle partitioning
452 and the determination of gas-particle partitioning coefficients of larger organic peroxides
453 deserve further study to improve the simulation of SOA mass.

454 We also examined gas-phase peroxides. The yield of gaseous total peroxides was ~0.22, which
455 was independent of RH and OH scavengers. The MCM v3.1 mechanism predicted this yield
456 but failed to explain the yields of individual peroxides, i.e., H₂O₂, HMHP, PFA and PAA,
457 indicating that our previous understanding of α -pinene ozonolysis was insufficient. For H₂O₂
458 with a yield of 0.048 under dry conditions and 0.16 under wet conditions, the known pathways,
459 including dissolution of SOA, aqueous oxidation of gaseous compounds, and

460 decomposition/hydrolysis of organic peroxides in the aqueous phase, cannot explain such an
461 unexpectedly high yield of H₂O₂. The presence of both water and SOA leads to the rapid
462 transformation of gaseous organic peroxides into H₂O₂. This heterogeneous process increases
463 the H₂O₂ yield by ~67%. Our results also show that water vapor affects the distribution of
464 gaseous peroxides, although it cannot change the yield of total peroxides.

465 The rapid heterogeneous transformation of organic peroxides to H₂O₂ helps to explain the
466 differences between modelled and observed levels of peroxides and OH in the forest area. In
467 the airborne GABRIEL field campaign in equatorial South America (Surinam) in October 2005
468 (Kubistin et al., 2010), two issues arose: (1) organic peroxides were overestimated while H₂O₂
469 was underestimated and (2) OH and HO₂ were also underestimated, especially when
470 concentrations of VOCs were high. These investigators suggested the occurrence of additional
471 recycling from HO₂ to OH or the contributions of additional direct OH sources. Our finding
472 that organic peroxides can transform to H₂O₂ by rapid heterogeneous reactions can address the
473 first discrepancy directly and the second indirectly. Peroxides influence OH through the
474 removal pathways:



479 Predominant removal paths for organic peroxides in the atmosphere are reaction with OH (95%)
480 and photolysis (4.4%) (Khan et al., 2015), while for H₂O₂, these two paths are almost equally
481 important. The OH oxidation process consumes OH while photolysis process produces OH.
482 Obviously, H₂O₂ plays a different role in the OH cycling compared with organic peroxides. One
483 molecule of organic peroxides transformed into H₂O₂ yields ~1.4 molecules of OH. Thus, the
484 rapid transformation of organic peroxides to H₂O₂ by the heterogeneous process would increase
485 OH levels. However, not all the organic peroxides could be transformed to H₂O₂ by the
486 heterogeneous process. Further studies are needed to clarify this process in the atmosphere and
487 unveil the features of the peroxides undergoing heterogeneous transformation.

488 **Acknowledgements**

489 We gratefully acknowledge the National Natural Science Foundation of China (grants
490 41275125, 21190053, 21477002) and the State Key Laboratory of Environment Simulation and
491 Pollution Control (special fund) for financial support.

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689 **Table 1.** Peroxide content in the gas phase and particle phase of α -pinene ozonolysis as affected by the OH
 690 radical scavenger cyclohexane^a.

OH scavenger RH	None		Cyclohexane	
	<0.5%	60%	<0.5%	60%
Gas phase				
Y ^b _{H₂O₂}	0.048±0.012	0.16±0.01	0.048±0.010	0.14±0.02
Y ^b _{HMHP}	0.0030±0.0003	0.0062±0.0005	0.0024±0.0005	0.0037±0.0006
Y ^b _{PFA}	0.0057±0.0020	0.012±0.002	0.0020±0.0003	0.005±0.001
Y ^b _{PAA}	0.0067±0.0006	0.009±0.001	0.0022±0.0002	0.0024±0.0001
Y ^b _{TPO}	0.18±0.01	0.20±0.01	0.27±0.02	0.25±0.02
H ₂ O ₂ (g)/TPO(g) (ppbv ppbv ⁻¹)	0.26±0.02	0.78±0.04	0.18±0.02	0.50±0.04
Particle phase				
Y ^c _{SOA}	0.41±0.01	0.39±0.02	0.28±0.02	0.27±0.01
F ^d _{H₂O₂} (ng µg ⁻¹)	5.09±0.99	2.67±0.17	1.61±0.11	1.41±0.21
F ^d _{PFA} (ng µg ⁻¹)	0.35±0.06	0.27±0.02	0.14±0.01	0.16±0.04
F ^d _{PAA} (ng µg ⁻¹)	0.11±0.04	0	0.09±0.01	0
F ^d _{TPO} (µg µg ⁻¹)	0.23±0.01	0.25±0.01	0.16±0.01	0.20±0.01
H ₂ O ₂ (p)/TPO(p) (µM µM ⁻¹)	0.20±0.03	0.10±0.01	0.09±0.01	0.06±0.01
TPO(p)/TPO(g+p) ^e	0.19±0.02	0.18±0.03	0.07±0.01	0.09±0.01

691 ^a ~275 ppbv α -pinene, ~1300 ppmv cyclohexane and ~42 ppmv O₃ were used in these experiments. The data
 692 represent the mean ± s.d. of 3 observations.

693 ^b molar yield of peroxides

694 ^c mass yield of SOA

695 ^d contribution of peroxides to SOA mass

696 ^e fraction of particulate TPO in gaseous and particulate TPO

697

698 **Table 2.** Comparison of observed and theoretical gas-particle partitioning coefficients (K_p) of PFA, PAA and
 699 TPO at different scenarios (298K).

	K_p (observed)				K_p (theoretical)
	$(\text{m}^3 \mu\text{g}^{-1})^a$				$(\text{m}^3 \mu\text{g}^{-1})$
	Sc1	Sc2	Sc3	Sc4	
PFA	8.06×10^{-5}	2.95×10^{-5}	9.19×10^{-5}	4.20×10^{-5}	2×10^{-9}
PAA	1.76×10^{-5}	0	4.38×10^{-5}	0	4×10^{-9}
TPO	3.47×10^{-4}	3.39×10^{-4}	1.61×10^{-4}	2.17×10^{-4}	–

700 ^a The four scenarios represent four reaction conditions: Sc1 (<0.5% RH, no OH scavenger), Sc2 (60% RH,
 701 no OH scavenger), Sc3 (<0.5% RH, with cyclohexane) and Sc4 (60% RH, with cyclohexane). Cyclohexane
 702 used here was ~1300 ppmv.
 703

704 **Table 3.** Hydrogen peroxide in the coil collector at different scenarios in the two-stage experiments.

Scenarios	Filter position ^a	Water vapor ^b	H ₂ O ₂	Species in second reactor
1d	Second reactor	No	100±1%	Gaseous products, SOA
1w	Second reactor	Yes	165±6%	Gaseous products, SOA, water vapor
2d	First reactor	No	103±6%	Gaseous products
2w	First reactor	Yes	87±7%	Gaseous products, water vapor
3d	No filter	No	164±9%	Gaseous products, SOA ^c
3w	No filter	Yes	172±5%	Gaseous products, SOA, water vapor

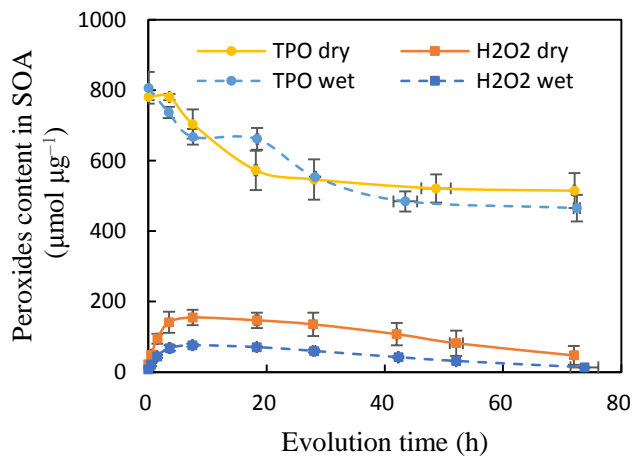
705 ^a The filters were placed at the outlet of the first reactor or second reactor.

706 ^b Water vapor was induced into the second reactor.

707 ^c Although gaseous products and SOA did not contact water vapor in the second reactor, they were in contact
 708 with the condensed water in the coil collector.

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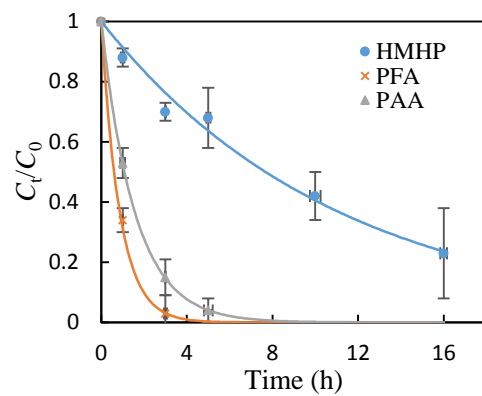
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712 **Figure 1.** Evolution of total peroxides and H₂O₂ contents in SOA produced under dry (<0.5% RH) and wet
713 (60% RH) conditions at 298K. Circles and diamonds represent total peroxides and H₂O₂ contents in SOA,
714 respectively; and solid lines and dash lines represent that obtained under dry and wet conditions, respectively.
715 The data represent the mean ±s.d. of 3 observations.

716



717

718 **Figure 2.** Decomposition/hydrolysis of organic peroxides in the aqueous phase. C_t/C_0 is the ratio of peroxides
 719 concentration at time = t h to peroxides concentration at time = 0 h. Lines are exponential fits for HMHP,
 720 PFA and PAA. The decay rate constants of HMHP, PFA and PAA are 0.09, 1.06 and 0.64 h^{-1} , respectively.
 721 The data represent the mean \pm s.d. of 3 observations.