



1 **Calibrating adsorptive and reactive losses of monoterpenes and sesquiterpenes in**
2 **dynamic chambers using deuterated surrogates**

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10 **Abstract**

11 Accurately measuring the emissions of monoterpenes (MTs) and sesquiterpenes (SQTs) using dynamic
12 chambers requires careful consideration of their adsorptive and reactive losses, which are often overlooked
13 and difficult to assess in situ. This study evaluated the effectiveness of deuterated surrogates, α -pinene-d3
14 and β -caryophyllene-d2, in tracing these losses in a dynamic chamber system. Using standard gas mixtures
15 of 10 MTs and 10 SQTs, we characterized adsorptive losses across varying concentrations, temperatures, and
16 humidity levels, as well as reactive losses with ozone. Results indicated that adsorptive losses were
17 significantly influenced by concentration and temperature, with species-specific variations particularly under
18 low concentrations and low temperatures, while relative humidity had negligible impact. Reactive losses with
19 ozone exhibited substantial species-specific variability. Key MTs (α -pinene, β -pinene, 3-carene, limonene,
20 and 1,8-cineole) and SQTs (β -caryophyllene and α -humulene) demonstrated consistent adsorptive and
21 reactive behavior with their respective deuterated surrogates α -pinene-d3 and β -caryophyllene-d2, suggesting
22 that these surrogates are effective for correcting losses in in-situ emission measurements using dynamic
23 chambers. However, due to varied adsorptive and reactive losses, additional deuterated MTs and SQTs are
24 recommended, particularly selected according to their O₃ reactivities, to cover a broader range of MTs and
25 SQTs for loss correction. A strong correlation between adsorptive capacity and ozone reactivity was observed,
26 underscoring the need to carefully address losses of highly reactive MTs and SQTs during emission
27 measurements. This study also emphasizes that ozone-free circulating air should be used for accurately
28 measuring emissions of highly reactive SQTs, such as β -caryophyllene and α -humulene, especially when loss
29 correction methods are unavailable.

30



31 **1 Introduction**

32 Monoterpenes (MTs) and sesquiterpenes (SQTs) emitted from terrestrial vegetation account for about 18%
33 of global biogenic volatile organic compound (BVOC) emissions, with even higher contributions in certain
34 ecosystems (Guenther et al., 2012). These reactive organic gases can rapidly react with atmospheric oxidants
35 such as ozone (O₃), hydroxyl radicals (OH), and nitrate radicals (NO₃) (Di Carlo et al., 2004; Jardine et al.,
36 2011; Edwards et al., 2017; He et al., 2021), influencing atmospheric oxidation capacity and leading to the
37 formation of organic aerosols, which, as significant components of fine particles, can scatter solar radiation
38 and act as cloud condensation nuclei (Yli-Juuti et al., 2021), directly and indirectly affecting air quality and
39 climate (Peñuelas and Staudt, 2010; Unger, 2014; Scott et al., 2017).

40 Despite their importance, current BVOC emission models carry large uncertainties, largely due to inaccurate
41 emission factors (Guenther et al., 2006, 2012). Common BVOC emission measurement techniques, such as
42 leaf cuvettes and dynamic branch chambers (Niinemets et al., 2011; Šimpraga et al., 2019), are prone to
43 adsorptive and reactive losses, particularly for the less volatile MTs and SQTs (Ortega et al., 2008). Their
44 adsorptive losses, which can vary with conditions like temperature and concentration, may lead to significant
45 underestimation of emission rates (Niinemets et al., 2011; Zeng et al., 2022a). For example, Helmig et al.
46 (2004) reported that, after a 5-hour equilibrium, only 80% of SQTs and 60% of oxygenated SQTs were
47 recovered in a leaf cuvette, requiring 5-10 hours to purge to background levels. Additionally, MTs and SQTs
48 have short atmospheric lifetimes (from minutes to hours) due to their high reactivity with OH and O₃
49 (Atkinson and Arey, 2003). Field studies have shown significant in-canopy losses of SQTs (up to 61%) due
50 to O₃ in the Amazonian rainforest (Jardine et al., 2011). These findings highlight the risk of underestimating
51 the emissions of reactive terpenes using existing methods, especially in dynamic chambers where adsorptive
52 and reactive losses remain a challenge (Niinemets et al., 2011).

53 Dynamic chambers, commonly made from chemically inert materials like Teflon or Tedlar, minimize but do
54 not eliminate these losses (Niinemets et al., 2011; Materic et al., 2015). Previous studies have reported that
55 adsorptive losses of MTs and SQTs in such chambers can exceed 30%, even with residence times of the
56 chamber under one minute (Zeng et al., 2022a). Field studies also revealed that adsorptive losses in Tedlar
57 bags reached 20-30% for MTs and ranged 10-80% for SQTs (Ortega et al., 2008). Temperature plays a critical
58 role, with more substantial adsorptive losses at lower temperatures (Schaub et al., 2010). The adsorbed
59 compounds may re-emit at higher temperatures, leading to potential overestimation (Ruuskanen et al., 2005;
60 Schaub et al., 2010). Meanwhile, the presence of O₃ complicates measurements, as it both influences BVOC
61 emissions from plants and reacts with MTs and SQTs (Feng et al., 2019; Zeng et al., 2022b). Some studies
62 have used O₃-free circulating air to mitigate the reactive losses (Helmig et al., 2007; Ortega et al., 2008;
63 Hellén et al., 2021), but this may neglect the impact of O₃ on BVOC emissions. In contrast, the use of ambient
64 air can reflect real-world conditions (Kuhn et al., 2002; Bourtsoukidis et al., 2012; Zeng et al., 2022b), but
65 the reactive loss existed. Despite reducing residence time by increasing flow rate of circulating air can lower



66 the reactive losses (Zeng et al., 2022a), it may still not be suitable for highly reactive MTs and SQTs
67 (Pollmann et al., 2005; Niinemets et al., 2011).

68 Addressing these issues is crucial for improving BVOC emission measurements. While internal surrogates
69 like aromatic compounds have been used to assess adsorptive losses, their suitability as surrogates for MTs
70 and SQTs remains debated (Helmig et al., 2007; Ortega et al., 2008). More importantly, the reactivity of these
71 compounds with O₃ in dynamic chambers has not been thoroughly evaluated (Pollmann et al., 2005; Helin et
72 al., 2020; Zeng et al., 2022a), and this reactive loss for individual MT and SQT species cannot be assessed
73 by using surrogates like aromatic compounds with quite different reactivity with O₃.

74 This study investigates the potential of using deuterated α -pinene-d₃ and β -caryophyllene-d₂ to quantify the
75 adsorptive and reactive losses of individual MTs and SQTs in dynamic chambers, taking advantage of the
76 almost identical adsorption and reaction behaviors of these deuterated compounds as their counterpart MTs
77 and SQTs. We evaluated these losses of α -pinene-d₃ and β -caryophyllene-d₂, along with target MTs and
78 SQTs, under varying concentration, temperature, and humidity, and analyzed the gas-phase reactive losses
79 with O₃ at different concentrations.

80 **2 Materials and methods**

81 **2.1 Lab evaluation**

82 A cylindrical dynamic chamber, which is made of polymethyl methacrylate with inner surface coated with
83 fluorinated ethylene propylene (FEP) Teflon film (FEP 100, Type 200A, DuPont, Witton, USA), was used
84 for lab evaluation. Detailed information about the design and initial characterization of this dynamic chamber
85 was described in our previous study (Zeng et al., 2022a). Here, we used standard gas mixtures of α -pinene-
86 d₃, β -caryophyllene-d₂, ten species of MTs, and ten species of SQTs to simulate their emissions from
87 enclosed leaves in the chamber. Detailed description of the preparation of the standard gas mixtures can be
88 found in Zeng et al. (2022a), and the information of standards were provided in Table S1. As shown in Fig.
89 S1, when the chamber reached a steady state offline samples were collected simultaneously from inlet (C_0)
90 and outlet (C_1) by commercial adsorbent cartridges (Tenax TA/Carbograph 5TD, Markes International Ltd,
91 Bridgend, UK) connected onto a portable dual-channel sampler (ZC-QL, Zhejiang Hengda Instrumentation
92 Ltd., Zhejiang, China) at a rate of 200 mL min⁻¹ for 5 min. Therefore, the recovery was expressed as C_1/C_0
93 in percentage (Fig. S1). More detail descriptions of the experiment are provided in Text S1.

94 The effect of concentration on adsorptive losses was evaluated under four concentration levels, with total
95 mixing ratios of MTs and SQTs of 0.23 and 0.26, 0.37 and 0.47, 0.68 and 0.94, and 2.26 and 3.16 ppb,
96 respectively. These experiments were conducted in O₃-free air at a temperature of approximately 25 °C and
97 a relative humidity of about 50%. The temperature effect was tested at two MTs and SQTs concentration
98 levels: a low concentration of 0.22 and 0.28 ppb and a high concentration of 2.26 and 3.16 ppb. In these



99 experiments, the relative humidity was about 50% but varied slightly with temperature, while the circulating
100 air was O₃-free. For the low-concentration experiments, the temperature ranged from 15 °C to 40 °C in 5 °C
101 increments, whereas the high-concentration experiments started at 10 °C and ended at 45 °C, also in 5 °C
102 increments. Humidity effect was tested under both low and high concentration levels, similar to the
103 temperature experiments. These humidity experiments were conducted at low (5%), mid (50%), and high
104 (95%) relative humidity (RH) levels, with a constant temperature of about 25 °C and O₃-free circulating air.
105 Additionally, the ozone effect was assessed at six ozone mixing ratios of 0, 30, 45, 60, 100, and 200 ppb,
106 with temperature of about 25 °C and RH of approximately 50%.

107 2.2 Lab analysis

108 Cartridge samples were analyzed by an automatic thermal desorption system (TD-100, Markes International
109 Ltd, Bridgend, UK) coupled to a mode 7890 gas chromatography (GC) with a model 5975 mass selective
110 detector (MSD) (Agilent Technologies, Inc., CA, USA). The adsorbent cartridges were thermally desorbed
111 by the TD-100 at 280 °C for 10 minutes and then the desorbed analytes were transferred by pure helium into
112 a cryogenic trap (U-T11PGC-2S, Markes International Ltd, Bridgend, UK) at -10 °C. Then the trap was
113 rapidly heated to transfer the analytes to the GC/MSD system with an HP-5MS capillary column (30 m ×
114 0.25 mm × 0.25 μm, Agilent Technologies, Inc., CA, USA). The GC oven temperature was programmed to
115 be initially at 35 °C (held for 3 minutes), increased to 100 °C at 5 °C min⁻¹ and held for 1 minute, then to
116 120 °C at 10 °C min⁻¹ and held for 12 minutes, and then to 260 °C with a final hold time of 2 minutes. The
117 MSD was operated simultaneously under scan mode and selected ion monitoring mode with electron
118 impacting ionization at 70 eV. Target compounds were identified by comparing their retention times with
119 authentic standards (Table S1), and were quantified with the standard calibration curves. More detailed
120 descriptions of the identification and quantification can refer to our previous studies (Zeng et al., 2022b; Zeng
121 et al., 2023; Zeng et al., 2024). Method detection limits (MDLs) were determined by seven parallel analyses
122 of the lowest concentration of calibration standards. Based on a sample volume of 1 L, the MDLs varied from
123 8 to 92 ng m⁻³ for MTs and from 2 to 92 ng m⁻³ for SQTs, and the measurement precision ranged from 2.3 to
124 4.8 % for MTs and from 1.1 to 3.4 % for SQTs (Table S2).

125 3 Results and discussion

126 3.1 Concentration effect

127 The adsorptive behaviors of different MTs and SQTs varied significantly across concentration levels (Fig. 1).
128 At low concentrations, recoveries ranged from 59% for γ-terpinene to 81% for α-pinene among MTs and
129 from 40% for α-humulene to 71% for longicyclene among SQTs (Fig. 1a). In contrast, under high
130 concentration levels, recoveries were more consistent, ranging from 75% for β-myrcene to 87% for β-pinene
131 among MTs and from 74% for β-chamigrene to 84% for α-longipinene for SQTs (Fig. 1d). Overall, recoveries



132 were higher at elevated concentrations (Figs. S3 and S4), with average recoveries of $68\pm 9\%$, $75\pm 7\%$, $78\pm 5\%$,
133 and $84\pm 2\%$ for all MTs, and $58\pm 8\%$, $63\pm 5\%$, $71\pm 4\%$, and $82\pm 1\%$ for all SQTs across four concentration
134 levels (Fig. S5).

135 Despite using chemically inert materials in chamber construction, significant adsorptive losses of MTs and
136 SQTs persist (Ortega et al., 2008; Niinemets et al., 2011; Kolari et al., 2012; Zeng et al., 2022a). Our findings
137 corroborate previous studies, demonstrating that low concentrations lead to higher relative losses (Fig. 1). In
138 steady-state conditions, the amount of adsorbed MTs and SQTs is constant, implying that lower concentration
139 results in a greater proportion of total loss (Figs. S3 and S4). This suggests that low-emitting plants, especially
140 for SQTs, may experience substantial adsorptive losses.

141 We observed species-specific recovery differences at low concentrations (Fig. 1a), likely due to their
142 physicochemical properties (Niinemets et al., 2004, 2011). Less reactive MTs like α -pinene-d3, α -pinene, β -
143 pinene, and 1,8-cineole exhibited stable recoveries regardless of concentration changes. In contrast, more
144 reactive MTs, such as β -myrcene, α -phellandrene, γ -terpinene, and terpinolene, showed significant recovery
145 variations (Fig. S3). This pattern also applies to SQTs, with highly reactive compounds, like β -caryophyllene,
146 β -caryophyllene-d2, and α -humulene, being more affected than others (Fig. S4). This compound-specific
147 adsorptive losses indicate that α -pinene-d3 and β -caryophyllene-d2 cannot serve as surrogates for all MTs
148 and SQTs, respectively.

149 3.2 Temperature effect

150 Overall recoveries of MTs and SQTs were higher at elevated temperatures, with a smaller absolute effect
151 observed at high concentrations compared to low concentrations (Figs. 2 and S6). At the high concentration,
152 mean recoveries increased from $80\pm 4\%$ to $91\pm 1\%$ for MTs and from $73\pm 3\%$ to $89\pm 3\%$ for SQTs as
153 temperature rose from 10 to 45 °C (Figs. S7 and S8). At the low concentration, temperature influenced
154 recoveries more significantly (Fig. 2), with mean values rising from $70\pm 12\%$ to $85\pm 4\%$ for MTs and from
155 $50\pm 21\%$ to $77\pm 6\%$ for SQTs as temperature increased from 15°C to 40°C. At low concentration and low
156 temperatures (e.g., 15 °C), the standard deviation of recoveries was greater, such as 12% for MTs and 21%
157 for SQTs, indicating variability among species (Fig. 2). However, at low concentration but high temperatures
158 (e.g., 40°C), recoveries became more consistent, with reduced standard deviation of 4% for MTs and 6% for
159 SQTs (Fig. 2).

160 The temperature effect on recovery was compound-specific. Among MTs, β -myrcene, α -phellandrene, γ -
161 terpinene, terpinolene, and linalool showed greater sensitivity (Fig. 2a), while β -caryophyllene, β -
162 caryophyllene-d2, and α -humulene showed greater sensitivity among SQTs (Fig. 2b). We categorized MTs
163 and SQTs into two groups based on their adsorptive behaviors (Fig. 3). MTs in group 1 included α -pinene-
164 d3, α -pinene, β -pinene, 3-carene, limonene, and 1,8-cineole, while MTs in group 2 consisted of β -myrcene,



165 α -phellandrene, γ -terpinene, terpinolene, and linalool. SQTs in group 1 comprised α -longipinene,
166 longicyclene, α -copaene, α -gurjunene, thujopsene, aromadendrene, alloaromadendrene, and β -chamigrene,
167 whereas SQTs in group 2 included β -caryophyllene, β -caryophyllene-d2, and α -humulene. Under the high
168 concentration, temperature changes minimally affected recoveries, with slopes of $0.37\pm 0.07\%$ °C⁻¹ for all
169 MTs and $0.43\pm 0.08\%$ °C⁻¹ for all SQTs (Fig. 3). However, under the low concentration, sensitivity varied
170 significantly between groups, with slopes of $0.27\pm 0.06\%$ °C⁻¹ for MTs and $0.53\pm 0.05\%$ °C⁻¹ for SQTs in
171 group 1 much lower than those $1.12\pm 0.12\%$ °C⁻¹ for MTs and $2.51\pm 0.19\%$ °C⁻¹ for SQTs in group 2 (Fig. 3).

172 Our results demonstrate that adsorptive losses of MTs and SQTs are influenced by both concentration and
173 temperature. Elevated temperatures facilitate evaporation and diffusion, reducing adsorptive losses
174 (Niinemets et al., 2004; Ortega and Helmig, 2008). Field measurements on *Picea pungens* have similarly
175 shown increased recoveries at high temperatures (Ortega et al., 2008). At low concentrations, the pronounced
176 sensitivity to temperature results in greater recovery differences among species particularly at lower
177 temperatures (Fig. 3), indicating that α -pinene-d3 and β -caryophyllene-d2 may not effectively calibrate
178 adsorptive losses across all MTs and SQTs. Since the inter-group adsorptive behaviors (e.g., MTs in group 1
179 vs. in group 2) differ significantly but the intra-group adsorptive behaviors for both MTs and SQTs remain
180 consistent (Fig. 3), suggesting that α -pinene-d3 and β -caryophyllene-d2 instead can be used to calibrate the
181 adsorptive losses for MTs in group 1 (α -pinene, β -pinene, 3-carene, limonene, and 1,8-cineole) and SQTs in
182 group 2 (β -caryophyllene and α -humulene), respectively.

183 3.3 Humidity effect

184 The impact of humidity on the adsorptive losses of MTs and SQTs was examined at three RH levels: 5%,
185 50%, and 95%. Overall recoveries were slightly higher at 95% RH compared to lower RH levels (Fig. 4). For
186 low-concentration experiments, mean recoveries of MTs and SQTs were $72\pm 9\%$ and $64\pm 20\%$ at 5% RH,
187 $72\pm 10\%$ and $62\pm 13\%$ at 50% RH, and $80\pm 6\%$ and $70\pm 15\%$ at 95% RH, respectively (Fig. 4a), while for
188 high-concentration experiments, they were $77\pm 2\%$ and $76\pm 2\%$ at 5% RH, $79\pm 5\%$ and $82\pm 2\%$ at 50% RH,
189 and $82\pm 2\%$ and $82\pm 2\%$ at 95%, respectively (Fig. 4b), with recoveries more consistent across RH levels.

190 The results suggests that humidity has a minor impact on adsorptive loss, aligning with previous findings
191 (Hohaus et al., 2016; Helin et al., 2020; Zeng et al., 2022a). The relationship between adsorptive losses and
192 humidity is complex, as water molecules can compete with MTs and SQTs for adsorption sites, potentially
193 reducing the adsorption of hydrophobic MTs and SQTs (Kolari et al., 2012; Zeng et al., 2022a). These results
194 indicate that the effect of humidity on the adsorptive loss of MTs and SQTs in dynamic chambers can be
195 considered negligible during field studies.

196 3.4 Ozone effect

197 The impact of O₃ on the reactive loss of MTs and SQTs was evaluated at six mixing ratio levels: 0, 30, 45,



198 60, 100, and 200 ppb. Recoveries of α -pinene-d3, α -pinene, β -pinene, 3-carene, and 1,8-cineole remained
199 stable across O₃ levels. However, recoveries of β -myrcene, limonene, and γ -terpinene declined slightly with
200 elevated O₃ levels, and more pronounced losses were observed for α -phellandrene and terpinolene, with
201 recoveries of 70±3% and 78±2% at an O₃ mixing ratio of 60 ppb, respectively (Fig. S9). For SQTs,
202 longicyclene, α -copaene, aromadendrene, and alloaromadendrene exhibited relatively stable recoveries,
203 while α -longipinene, α -gurjunene, thujopsene, and β -chamigrene suffered significant losses (Fig. 5). Notably,
204 the recoveries of β -caryophyllene, β -caryophyllene-d2, and α -humulene dropped sharply with increasing O₃
205 levels, with recoveries as low as 41±3%, 36±6%, and 30±4%, respectively, even at a low O₃ level of 30 ppb
206 (Fig. 5).

207 Given ozone's role in reactive losses, some studies have used O₃-free circulating air during field
208 measurements to mitigate these effects (Ortega et al., 2008; Helmig et al., 2013). However, it is also critical
209 to understand ozone's role in affecting emissions of MTs and SQTs (Bourtsoukidis et al., 2012; Feng et al.,
210 2019; Zhang et al., 2022). Since the measured emissions may not reflect real-world emissions if O₃ is cleaned,
211 some studies have used ambient air as circulating air during field measurements (Kuhn et al., 2002;
212 Bourtsoukidis et al., 2012; Zeng et al., 2022b). The relationship between reactive loss and O₃ concentration
213 can be modelled by:

$$214 \quad C_t = C_0 \cdot \exp(-k \cdot t \cdot [\text{O}_3]) \quad (1)$$

215 where C_0 (molecule cm⁻³) and C_t (molecule cm⁻³) are the initial and time t concentrations in the chamber,
216 respectively; k (cm³ molecule⁻¹ s⁻¹) is the reaction rate constant; t (s) is the reaction time of the compound in
217 the chamber; [O₃] (molecule cm⁻³) represents the concentration of O₃. In this study, when the C_t reached to
218 equilibrium concentration (C_s) with a reaction time of the residence time, Eq. 1 can be rewritten as:

$$219 \quad -\ln(C_s/C_0) = k \cdot \frac{V}{F} \cdot [\text{O}_3] \quad (2)$$

220 where V (L) is the volume of the chamber and F (L s⁻¹) is the flow rate of circulating air. As a result, through
221 the application of a linear regression between $-\ln(C_s/C_0)$ and [O₃], we can derive k values for MTs and SQTs,
222 as illustrated in Fig. S10. However, k values for some less reactive MTs and SQTs cannot be obtained this
223 way and Table 1 presents the k values of some highly reactive species. In present study, the k values of β -
224 myrcene, α -phellandrene, limonene, and terpinolene were 5.93±0.17, 28.15±1.11, 3.06±0.14, and 19.21±1.20
225 (×10⁻¹⁶) cm³ molecule⁻¹ s⁻¹, respectively, which are in agreement with those reported in literatures (Table 1).
226 For SQTs, the k values were 20.17±0.98, 14.56±0.15, 115.33±3.83, 129.14±4.42, 8.64±0.16, 147.04±3.87,
227 and 15.14±0.16 (×10⁻¹⁶) cm³ molecule⁻¹ s⁻¹ for α -longipinene, α -gurjunene, β -caryophyllene, β -
228 caryophyllene-d2, thujopsene, α -humulene, and β -chamigrene, respectively. The k values for the most
229 important β -caryophyllene and α -humulene were consistent with those reported previously (Table 1). The



230 good consistency of k values with literatures proves that the assessment of reactive losses with O_3 in the
231 dynamic chamber used here is reliable.

232 The compound-specific reactive losses due to their varying reactivity with O_3 (Table 1) necessitates their
233 consideration during field measurements, particularly for highly reactive species like β -caryophyllene and
234 α -humulene. Reactive loss caused by O_3 can be reduced by shortening the residence time through increasing
235 the flow rate of circulating air. For the chamber in this study, with a flow rate of 9 L min^{-1} for circulating air
236 (~ 1.5 minutes of residence time), the reactive losses of all tested MTs, except for α -phellandrene and
237 terpinolene, were less than 10%, even at a high O_3 mixing ratio of 100 ppb (Fig. S9). This suggests that the
238 effect of O_3 on the reactive losses of main MTs (e.g., α -pinene, β -pinene, limonene) can be disregarded during
239 field measurements. For SQTs, although the reactive losses for longicyclene, α -copaene, aromadendrene, and
240 alloaromadendrene were less than 10% at 100 ppb O_3 , more than 60% of the critical species like β -
241 caryophyllene and α -humulene were consumed by O_3 even at a low mixing ratio of 30 ppb (Fig. 5). As shown
242 in Fig. S11, to limit the reactive loss of β -caryophyllene within 15%, the flow rate needs to exceed 42 L min^{-1}
243 while O_3 mixing ratio should be kept below 30 ppb. Although the detection of instrument is not a problem
244 even under this high flow rate (Zeng et al., 2022a), the requirement for an O_3 mixing ratio below 30 ppb
245 poses practical challenges, as this level is already lower than those observed in most background regions
246 worldwide (Lu et al., 2018; Wang et al., 2019). This suggests that using O_3 -free circulating air may be more
247 suitable when measuring emissions of highly reactive SQTs in the field, as a short-term O_3 scavenge during
248 measurements could have negligible impact on the BVOC emissions as demonstrated by Niinemets et al.
249 (2011).

250 3.5 Implications for field measurements

251 As discussed above and illustrated in Fig. 6, the effects of concentration, temperature, and O_3 on the losses
252 of MTs and SQTs are compound-specific. The adsorptive and reactive losses in group 2 are generally greater
253 than those in group 1, reflecting their different physicochemical properties (Niinemets et al., 2011). Notably,
254 we observed a positive correlation between adsorptive capacity and O_3 reactivity (Fig. 7), suggesting that
255 compounds with higher O_3 reactivity experience greater temperature- and concentration-driven adsorptive
256 losses. These findings underscore the importance of accounting for both reactive and adsorptive losses in
257 accurately measuring emissions of highly reactive MTs and SQTs.

258 Furthermore, findings here demonstrate temperature as a crucial factor in influencing the adsorptive losses
259 of terpenes especially SQTs in dynamic chambers. For instance, assuming leaves with a dry mass of 10 g are
260 enclosed in the chamber, the lowest concentration evaluated here, equivalent to emission rates of $0.07 \mu\text{g g}^{-1}$
261 h^{-1} for MTs and $0.12 \mu\text{g g}^{-1} \text{h}^{-1}$ for SQTs, represents a low MT-emitting but high SQT-emitting plant (Duhl et
262 al., 2008; Ortega et al., 2008). Despite being a low MT emitter, at temperatures below $30 \text{ }^\circ\text{C}$, the adsorptive
263 losses for MTs in group 1 are only about 20%, while those in group 2 exceed 30%. In contrast, adsorptive



264 losses for SQTs in groups 1 and 2 can be greater than 30% and 45%, respectively, even for a high SQT-
265 emitting plant. This suggests that in high-latitude regions with relatively cold weather, while adsorptive losses
266 of some key MTs, such as those in group 1, may be acceptable, losses can be significant for SQTs especially
267 the most important β -caryophyllene and α -humulene, thus needing loss corrections during measurements.
268 Moreover, in tropical and subtropical regions characterized by hot weather, high daytime temperatures and
269 strong sunlight can cause temperatures in dynamic chambers to exceed 35 °C, even surpassing 45 °C during
270 heat events (Zeng et al., 2022b; Doughty et al., 2023). Under these conditions, adsorptive losses can be
271 reduced to within 20% for most MTs and 25% for most SQTs, regardless of changes in concentration and
272 humidity. This indicates that adsorptive losses of MTs and SQTs may not pose a significant issue for dynamic
273 chambers used in hot regions; however, the reactive losses of highly reactive compounds must still be
274 carefully considered to ensure accurate measurement of their emission rates.

275 Although species-specific variabilities in adsorptive and reactive losses existed, we demonstrated that MTs
276 in group 1, including α -pinene, β -pinene, 3-carene, limonene, and 1,8-cineole, exhibit consistent adsorptive
277 behavior with α -pinene-d3, regardless of changes in concentration, temperature, or humidity, and the reactive
278 losses of these MTs are minimally affected by ambient-level O₃. This consistency makes α -pinene-d3 a
279 suitable surrogate. Similarly, β -caryophyllene-d2 aligns well with β -caryophyllene and α -humulene, and is a
280 suitable surrogate for these compounds. However, MTs in group 2 and SQTs in group 1 exhibit significant
281 differences in adsorptive and reactive behaviors compared to α -pinene-d3 and β -caryophyllene-d2,
282 respectively, and therefore cannot be calibrated using these two deuterated surrogates. Given the strong
283 correlation between adsorptive capacity and O₃ reactivity (Fig. 7), we recommend considering other
284 deuterated MTs and SQTs that share similar adsorptive and reactive behaviors with those in group 2 and
285 SQTs in group 1 as internal surrogates for loss correction.

286 **4 Conclusions**

287 In this study, lab evaluations identified significant species-specific variability in the adsorptive and reactive
288 losses of MTs and SQTs in dynamic chambers, influenced by their distinct physicochemical properties.
289 Adsorptive losses were notably influenced by concentration and temperature, with pronounced variability at
290 low concentrations and low temperatures. The reactive losses were also species-specific, with highly reactive
291 compounds like β -caryophyllene and α -humulene exhibiting significant losses. This emphasizes the
292 importance of using O₃-free circulating air during emission measurement to minimize these losses for highly
293 reactive SQTs.

294 This study supports the use of α -pinene-d3 and β -caryophyllene-d3 as effective surrogates for loss correction.
295 These surrogates showed consistent adsorptive and reactive behaviors with corresponding target compounds
296 (e.g., α -pinene, β -pinene, 3-carene, limonene, and 1,8-cineole for MTs, and β -caryophyllene and α -humulene
297 for SQTs), enabling more accurate calibration of losses during in-situ measurements. Additionally, the



298 observed correlation between adsorptive capacity and ozone reactivity indicates that other deuterated MTs
299 and SQTs could be employed to expand coverage for loss correction.

300 Our study demonstrates that adsorptive losses in dynamic chambers are highly temperature dependent. In
301 tropical and subtropical regions, higher temperatures potentially reduce adsorptive losses of MTs and SQTs
302 to acceptable levels. Conversely, low temperatures typical of high-latitude regions could lead to significant
303 adsorptive losses, particularly for SQTs, necessitating careful correction.

304 To ensure accurate emission measurements, it is crucial to account for both adsorptive and reactive losses of
305 MTs and SQTs, especially those with high reactivity. For highly reactive SQTs like β -caryophyllene and α -
306 humulene, O₃-free circulating air should be used to avoid substantial reactive losses. Furthermore, selecting
307 internal surrogates that closely match the adsorptive and reactive properties of target compounds is vital for
308 precise loss correction.

309

310 *Data availability*

311 All data used in this study are provided in the manuscript and/or the supplement.

312

313 *Supplement*

314 The supplement related to this article may be available online.

315

316 *Author contributions*

317 JZ designed and carried out the experiments with the support of XW, YZ, WS, and HG. JZ, HR, WP, and ZM
318 analyzed the samples in the lab. JQZ analyzed the data and prepared the original manuscript. XMW and YLZ
319 revised the manuscript.

320

321 *Competing interests*

322 The authors declare that they have no conflict of interest.

323



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330

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- 464



465 **Figure captions**

466 Figure 1. Recoveries of MT and SQT species under four different concentration levels. Total MTs and SQTs
467 mixing ratios are 0.23 and 0.26 ppb (a), 0.37 and 0.47 ppb (b), 0.68 and 0.94 ppb (c), and 2.26 and 3.16 ppb
468 (d), respectively.

469 Figure 2. Recoveries of different MT (a) and SQT (b) species under different temperatures under the low
470 concentration level.

471 Figure 3. Variations of recoveries of MTs and SQTs along with temperature under low and high concentration
472 levels. MTs group 1 includes α -pinene-d3, α -pinene, β -pinene, 3-carene, limonene, and 1,8-cineole, while
473 MTs group 2 includes β -myrcene, α -phellandrene, γ -terpinene, terpinolene, and linalool. α -Longipinene,
474 longicyclene, α -copaene, α -gurjunene, thujopsene, aromadendrene, and alloaromadendrene belong to SQTs
475 group 1, while β -caryophyllene, β -caryophyllene-d2, and α -humulene belong to SQTs group 2.

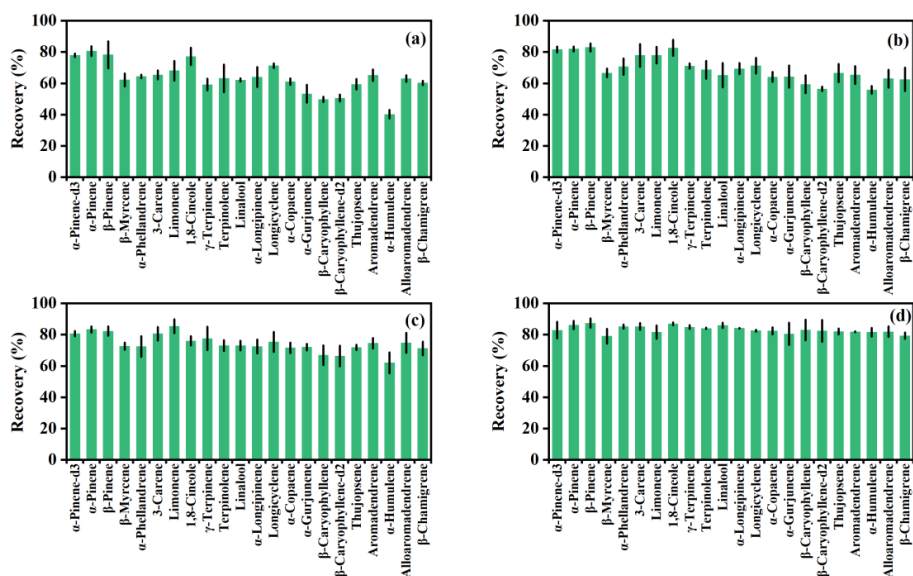
476 Figure 4. Recoveries of different MT and SQT species under different relative humidity levels; (a) low
477 concentration level, (b) high concentration level.

478 Figure 5. Reactive losses of SQT species under different O₃ mixing ratios.

479 Figure 6. The relationships between loss ratio of different MTs and SQTs groups with concentration,
480 temperature, and ozone mixing ratio.

481 Figure 7. Relationship of ozone reactivity with the effects of concentration (a) and temperature (b); The conc.
482 effect represents the absolute change of adsorptive loss per ppb ($|a|$) by fitting the loss ratio and the total
483 mixing ratios of MTs or SQTs using $y = a \cdot \ln(x) + b$ as in Fig. 6; The temp. effect represents the absolute change
484 of adsorptive loss per degree Celsius ($|a|$) by fitting the loss ratio and the temperature using $y = a \cdot x + b$ as in
485 Fig. 6; Each point represents an individual MT or SQT species.

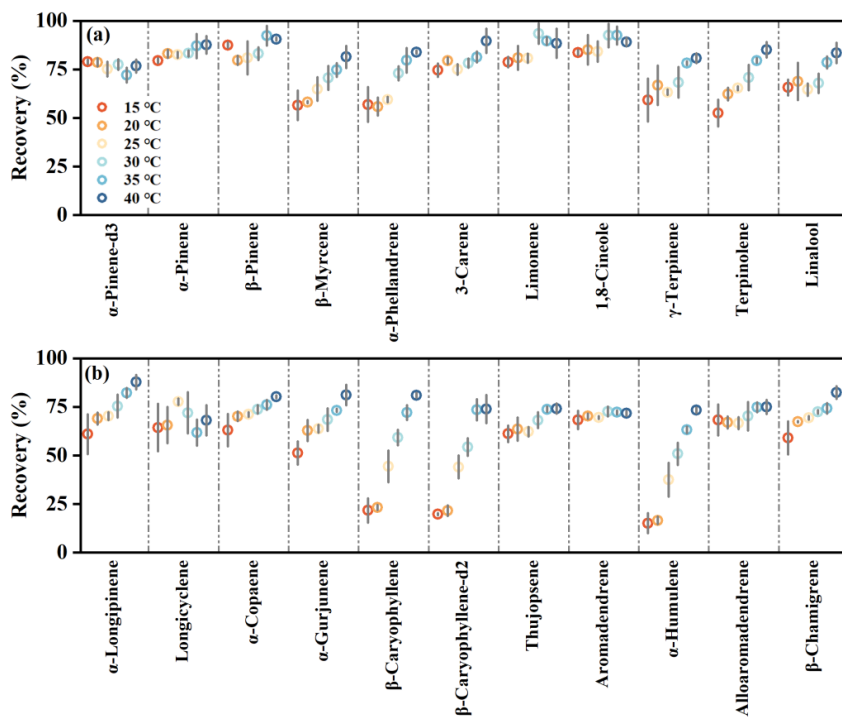
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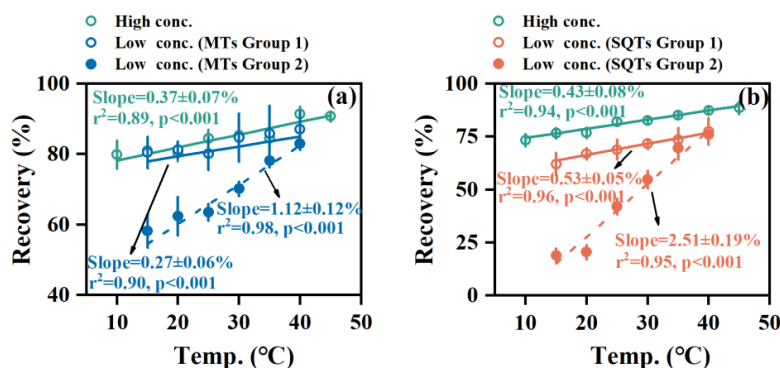
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493 **Figure 2.** Recoveries of different MT (a) and SQT (b) species under different temperatures under the low
494 concentration level.

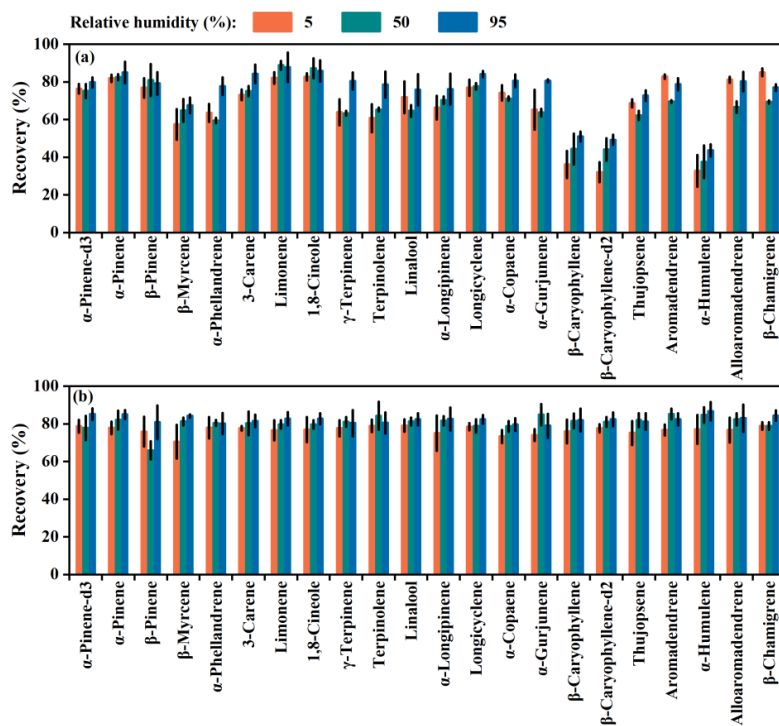
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496

497 **Figure 3.** Variations of recoveries of MTs and SQTs along with temperature under low and high concentration
498 levels. MTs group 1 includes α -pinene-d3, α -pinene, β -pinene, 3-carene, limonene, and 1,8-cineole, while
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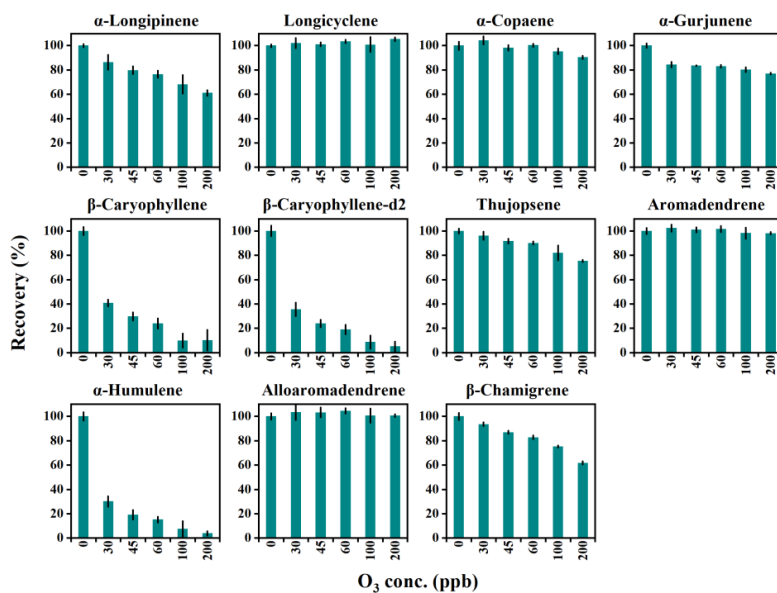
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506

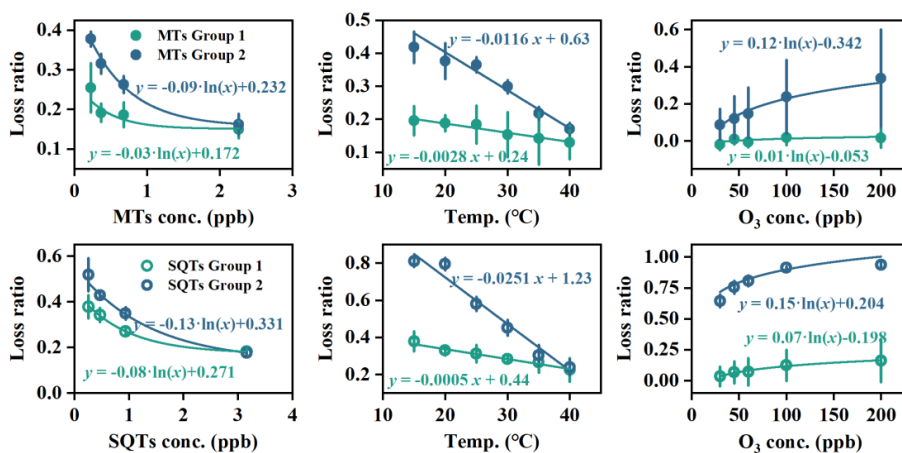


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Figure 5. Reactive losses of SQT species under different O₃ mixing ratios.

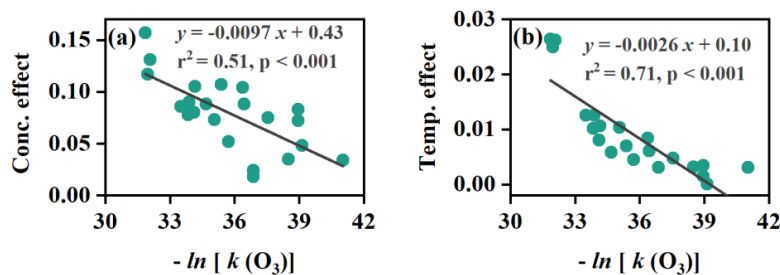
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513



514

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518 of adsorptive loss per degree Celsius ($|\Delta|$) by fitting the loss ratio and the temperature using $y = a \cdot x + b$ as in
519 Fig. 6; Each point represents an individual MT or SQT species.

520



521 **Table 1.** The measured and literature reported rate constants (k , $\times 10^{-16}$ cm³ molecule⁻¹ s⁻¹) of MTs and SQTs
522 with ozone.

Species	This study	Atkinson, 1997	Hoffmann et al., 1997	IUPAC* (298K)	Pollmann et al., 2005
β -Myrcene	5.93±0.17	4.70		4.7	
α -Phellandrene	28.15±1.11	29.8		29	
Limonene	3.06±0.14	2	2	2.2	
Terpinolene	19.21±1.20	18.8		16	
α -Longipinene	20.17±0.98				
α -Gurjunene	14.56±0.15				
β -Caryophyllene	115.33±3.83	116	116	120	110±5.1
β -Caryophyllene-d2	129.14±4.42				
Thujopsene	8.64±0.16				
α -Humulene	147.04±3.87	117		120	140±8.8
β -Chamigrene	15.14±0.16				

523 *IUPAC Task Group on Atmospheric Chemical Kinetic Data Evaluation (<http://iupac.pole-ether.fr>)