#### 1 Characterization of a Real-Time Tracer for Isoprene Epoxydiols-Derived Secondary

#### **Organic Aerosol (IEPOX-SOA) from Aerosol Mass Spectrometer Measurements** 2

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#### 52 Abstract

53 Substantial amounts of secondary organic aerosol (SOA) can be formed from isoprene

- 54 epoxydiols (IEPOX), which are oxidation products of isoprene mainly under low-NO conditions.
- 55 Total IEPOX-SOA, which may include SOA formed from other parallel isoprene oxidation
- 56 pathways, was quantified by applying Positive Matrix Factorization (PMF) to aerosol mass
- 57 spectrometer (AMS) measurements. The IEPOX-SOA fractions of OA in multiple field studies
- across several continents are summarized here and show consistent patterns with the
- 59 concentration of gas-phase IEPOX simulated by the GEOS-Chem chemical transport model.
- 60 During the Southern Oxidant and Aerosol Study (SOAS) study, 78% of PMF-resolved IEPOX-
- 61 SOA is accounted by the measured IEPOX-SOA molecular tracers (methyltetrols, C5-Triols and
- 62 IEPOX-derived organosulfate and its dimers), making it the highest level of molecular
- 63 identification of an ambient SOA component to our knowledge. Enhanced signal at  $C_5H_6O^+$  (m/z
- 64 82) is found in PMF-resolved IEPOX-SOA spectra. To investigate the suitability of this ion as a
- tracer for IEPOX-SOA, we examine  $f_{C_5H_6O}$  ( $f_{C_5H_6O}$  = C<sub>5</sub>H<sub>6</sub>O<sup>+</sup>/OA) across multiple field, chamber
- and source datasets. A background of  $\sim 1.7\pm0.1\%$  (%=parts per thousand) is observed in studies
- 67 strongly influenced by urban, biomass-burning and other anthropogenic primary organic aerosol
- 68 (POA). Higher background values of  $3.1\pm0.6\%$  are found in studies strongly influenced by
- 69 monoterpene emissions. The average laboratory monoterpene SOA value  $(5.5\pm2.0\%)$  is 4 times
- 10 lower than the average for IEPOX-SOA ( $22\pm7\%$ ), which leaves some room to separate both
- 71 contributions to OA. Locations strongly influenced by isoprene emissions under low-NO levels
- had higher  $f_{C_5H_6O}$  (~6.5±2.2‰ on average) than other sites, consistent with the expected IEPOX-
- 73 SOA formation in those studies.  $f_{C_5H_6O}$  in IEPOX-SOA is always elevated (12–40‰) but varies
- substantially between locations, which is shown to reflect large variations in its detailed
- molecular composition. The low  $f_{C_5H_6O}$  (<3‰) reported in non IEPOX-derived isoprene-SOA
- from chamber studies indicates that this tracer ion is specifically enhanced from IEPOX-SOA,
- and is not a tracer for all SOA from isoprene. We introduce a graphical diagnostic to study the
- presence and aging of IEPOX-SOA as a "triangle plot" of  $f_{CO_2}$  vs.  $f_{C_5H_6O}$ . Finally, we develop a
- real simplified method to estimate ambient IEPOX-SOA mass concentrations, which is shown to
- 80 perform well compared to the full PMF method. The uncertainty of the tracer method is up to a
- factor of ~2 if the  $f_{C_5H_6O}$  of the local IEPOX-SOA is not available. When only unit mass
- resolution data is available, as with the aerosol chemical speciation monitor (ACSM), all
- methods may perform less well because of increased interferences from other ions at m/z 82.
- 84 This study clarifies the strengths and limitations of the different AMS methods for detection of
- 85 IEPOX-SOA and will enable improved characterization of this OA component.

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### 1. Introduction

Isoprene (2-methyl-1,3-butadiene,  $C_5H_8$ ) emitted by vegetation is the most abundant non-87 methane hydrocarbon emitted to the Earth's atmosphere (~440–600 TgC/year) (Guenther et al., 88 89 2012). It is estimated to contribute substantially to the global secondary organic aerosol (SOA) 90 budget (Paulot et al., 2009b;Guenther et al., 2012). Higher SOA yields from isoprene are observed under low-NO<sub>x</sub> conditions (Surratt et al., 2010). Under low-NO conditions, i.e. when a 91 92 substantial fraction of the peroxy radicals do not react with NO, gas-phase isoprene epoxydiols 93 (IEPOX) are produced with high yield through a HO<sub>x</sub>-mediated mechanism (Paulot et al., 94 2009b). Note that some IEPOX can also be formed from isoprene in high NO region via 95 oxidation of the product 4-hydroxy-3-nitroxy isoprene (Jacobs et al., 2014), however this pathway is thought to be much smaller than the low-NO pathway. Subsequently, IEPOX can be 96 97 taken up by acidic aerosols (Gaston et al., 2014), where IEPOX-SOA can be formed through 98 acid-catalyzed oxirane ring-opening of IEPOX (Cole-Filipiak et al., 2010;Eddingsaas et al., 99 2010;Lin et al., 2012;Nguyen et al., 2014), which is thought to be the main pathway to form 100 IEPOX-SOA (Surratt et al., 2010;Pye et al., 2013;Worton et al., 2013). Although the complete 101 molecular composition of IEPOX-SOA has not been elucidated, several molecular species that 102 are part of IEPOX-SOA have been identified through gas chromatography/mass spectrometry 103 (GC/MS), liquid chromatography/mass spectrometry (LC/MS) and particle analysis by laser 104 mass spectrometry (PALMS). They include 2-methyltetrols (and oligomers that contain them) 105 (Surratt et al., 2010;Lin et al., 2014), C5-alkene triols (Wang et al., 2005), 3-106 methyltetrahydrofuran-3,4-diols (Lin et al., 2012), and an IEPOX-organosulfate (Froyd et al., 107 2010; Liao et al., 2014). These molecular species account for a variable fraction of the IEPOX-SOA reported, e.g., 8% in a chamber study (Lin et al., 2012) or 26% in a field study at Look 108 109 Rock, TN (Budisulistiorini et al., 2015). An estimate of total IEPOX-SOA can also be derived

from an IEPOX-SOA molecular tracer(s) via multiplying the tracer concentration by the total
IEPOX-SOA to tracer ratio. However, that method is hindered by the limited information on
these molecular tracers and the reported variability of IEPOX-SOA to tracer ratios. IEPOX-SOA
may include SOA formed from other parallel isoprene low-NO oxidation pathways (Liu et al.,
2014; Krechmer et al. 2015). In addition, the IEPOX-SOA molecular tracers are typically
measured with slow time resolution (12/24 h).

116 Multiple field studies, supported by chamber studies, have shown that the total amount of IEPOX-SOA can be obtained by factor analysis of organic spectra from an aerosol mass 117 118 spectrometer (AMS) or the aerosol chemical speciation monitor (ACSM) (Robinson et al., 119 2011;Lin et al., 2012;Budisulistiorini et al., 2013;Nguyen et al., 2014). Robinson et al. (2011) first reported an SOA factor with pronounced  $f_{82}$  (= m/z 82/OA) in the mass spectra acquired 120 121 above a forest with high isoprene emissions in Borneo, and hypothesized that the elevated  $f_{82}$ 122 may have arisen from methylfuran (C<sub>5</sub>H<sub>6</sub>O), consistent with C<sub>5</sub>H<sub>6</sub>O<sup>+</sup> being the major ion at m/z82 in isoprene-influenced areas. Lin et al. (2012) demonstrated that the 3-MeTHF-3,4-diols 123 124 associated with IEPOX-SOA result in enhanced  $f_{82}$  in AMS spectra, presumably through the 125 formation methylfuran-like structures during thermal desorption. Electron-impact ionization of 126 aerosols formed by atomizing a solution containing IEPOX ( $C_5H_{10}O_3$ ) can also yield  $C_5H_6O^+$ 127 signals in an AMS via two dehydration reactions (Lin et al., 2012). However, because gas-phase IEPOX has high volatility, non-reactive gas-to-particle partitioning of IEPOX into OA is 128 negligible under typical ambient concentrations in forest areas  $(1-10 \ \mu g \ m^{-3})$  (Worton et al., 129 130 2013).

131 IEPOX-SOA was estimated to account for 33% of ambient OA in summertime Atlanta from
132 PMF analysis of ACSM spectra. The source apportionment result was supported by the

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pronounced  $f_{82}$  peak in the factor spectrum and good temporal correlation of the factor with sulfate and 2-methyltetrols (Budisulistiorini et al., 2013). Sulfate is often strongly correlated with the acidity of an aerosol, and might also play a direct role in the chemistry, e.g. via direct reaction or nucleophilic effects (Surratt et al., 2007;Liao et al., 2014;Xu et al., 2014). While discussing the results of a recent aircraft campaign from Brazil, Allan et al. (2014) also used  $f_{82}$ as a tracer for IEPOX-SOA.

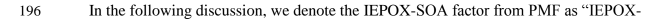
If  $f_{82}$  in AMS spectra (and/or  $f_{C_5H_6O}$  in HR-AMS spectra) is dominated by IEPOX-SOA,  $f_{82}$ 139 140 would be a convenient, high-time-resolution, and potentially quantitative tracer for IEPOX-SOA. 141 Thus, it will be very useful for investigating the impacts of SOA formation from isoprene with AMS/ACSM measurements, which have become increasingly common in recent years including 142 143 some continental-scale continuous networks (Fröhlich et al., 2015). However, no studies to date 144 have systematically examined whether enhanced  $f_{82}$  is unique to IEPOX chemistry or whether it 145 could also be enhanced in other sources. Nor has the range of  $f_{82}$  been determined for IEPOX-SOA. Questions also have been raised about the uniqueness of this tracer and potential 146 147 contributions from monoterpene SOA (Anonymous\_Referee, 2014). 148 In this study, the IEPOX-SOA results reported in various field campaigns are summarized 149 and compared to predicted gas-phase IEPOX concentrations from a global model to help confirm 150 the robustness of the AMS identification of this type of SOA. We then investigate the usefulness and limitations of the IEPOX-SOA tracers  $f_{C_5H_6O}$  (= C<sub>5</sub>H<sub>6</sub>O<sup>+</sup>/OA) and  $f_{82}$  by combining AMS 151 152 data from multiple field and laboratory studies including a new dataset from the 2013 Southern Oxidant and Aerosol Study (SOAS). We compare the tracer levels in different OA sources 153 (urban, biomass burning and biogenic), characterizing the background levels and interferences 154 155 on this tracer for both high-resolution (HR) and unit mass resolution (UMR) data. We also

provide a simplified method to rapidly estimate IEPOX-SOA from  $f_{C_5H_6O}$  and  $f_{82}$ . While this method is no substitute for a detailed IEPOX-SOA identification via PMF, it is a simple method to estimate IEPOX-SOA concentrations (or its absence) in real-time from AMS or ACSM measurements or under conditions in real-time, or where PMF analysis is not possible or is difficult to perform.

#### 161 **2 Experimental**

162 We classify the field datasets used in this study into three categories: (1) studies strongly 163 influenced by urban and biomass-burning emissions: Los Angeles area, US and Beijing, China 164 (urban); Changdao island, downwind of China and Barcelona area, Spain (urban downwind); flight data from biomass-burning plumes and continental areas (NW and western, US) in 165 166 SEAC4RS and DC3 campaigns; and biomass burning lab emissions (FLAME-3 study). (2) 167 Studies strongly influenced by isoprene emissions, including a SE US forest site (SOAS campaign); Two pristine forest site and one forest site partially impacted by urban plumes in the 168 169 Amazon rain forest (Brazil). The latter site is classified in this category because (i) high isoprene concentrations (e.g. 3 ppb in average peaks in the afternoon) were observed during the study; (ii) 170 the impact of biogenic SOA formed during 1000 km where the air travels over the pristine forest 171 172 upwind of Manaus; (iii) PMF results indicate an important impact of IEPOX-SOA at this site (de 173 Sá et al., 2015); (iv) PTRMS results indicate a substantial concentration of the isoprene 174 hydroperoxyde formed by low-NO chemistry. Borneo rain forest in Malaysia; and flight data 175 from SE US flights from aircraft campaign (SEAC4RS); (3) Studies strongly influenced by monoterpene emissions in a pine forest in the Rocky Mountains and a European boreal forest. 176 177 Locations and additional detailed information about these studies can be found in Fig. 1 and 178 Table 1.

179	With the exception of SOAS, all of the campaigns included in this analysis have been
180	previously described elsewhere (Table 1). The SOAS campaign took place in a forested area of
181	the SE US during June and July, 2013 (Fig. 1) and has several ground sites. The new dataset
182	introduced below was acquired at the SEARCH supersite, Centreville (CTR), AL (32.95° N,
183	87.13°W). Some results from a different SOAS site (Look Rock, TN) are also discussed later
184	(Budisulistiorini et al., 2015). Relatively high average isoprene and monoterpene concentrations
185	of $3.3\pm2.4$ ppb and $0.7\pm0.4$ ppb, respectively, were observed in SOAS-CTR by on-line GC/MS.
186	Measurements of non-refractory aerosol components of submicron particles (PM1) were made
187	using an Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS,
188	"AMS" hereafter) (DeCarlo et al., 2006). By applying positive matrix factorization (PMF) to the
189	time series of organic mass spectra (Ulbrich et al., 2009), we separated contributions from
190	IEPOX-SOA and other sources/components of OA. The AMS PMF results used here are very
191	consistent with those from a separate HR-ToF-AMS operated by another group at the same site
192	(Xu et al., 2014). The global gas-phase IEPOX concentrations in 2013 were modeled at as
193	resolution of 2 x 2.5 degrees as described in Nguyen et al. (2015). The gas-phase chemistry of
194	isoprene in GEOS-Chem is based on Paulot et al (2009a;2009b) as described by Mao et al.
195	(2013).



197 SOA<sub>PMF</sub>" and IEPOX-SOA from lab studies as "IEPOX-SOA<sub>lab</sub>" for clarity. If we use "IEPOX-

198 SOA" in the paper, it refers to a broad concept of IEPOX-SOA. We use a superscript to clarify

199 the type of OA for which  $f_{C_5H_6O}$  is being discussed:  $f_{C_5H_6O}^{OA}$  refers to  $f_{C_5H_6O}$  in total OA,

200  $f_{C_5H_6O}^{IEPOX-SOA}$  to  $f_{C_5H_6O}$  in IEPOX-SOA<sub>PMF</sub> or IEPOX-SOA<sub>lab</sub>,  $f_{C_5H_6O}^{MT-SOA}$  to the  $f_{C_5H_6O}$  value in pure

201 MT-SOA and  $f_{C_5H_6O}^{OA-Bkg-UB}$  and  $f_{C_5H_6O}^{OA-Bkg-MT}$  refer to background  $f_{C_5H_6O}^{OA}$  from areas strongly

influenced by urban+biomass-burning emissions and by monoterpene emissions, respectively. If we refer to  $f_{C_5H_6O}$  in general, we will still use  $f_{C_5H_6O}$ . When we report the average  $f_{C_5H_6O}^{OA}$  in each campaign, as shown in the Table 1, we used the average from the time series of  $f_{C_5H_6O}^{OA}$  at their raw time resolution (secs to mins). During this process, we exclude points whose OA mass concentrations are below twice the detection limit of OA in AMS (typically 2× 0.26 µg m<sup>-3</sup>=0.5 µg m<sup>-3</sup>). When averaging  $f_{C_5H_6O}^{OA}$  values across datasets, we counted each dataset as one data point.

209 **3 Results and Discussion** 

## 210 **3.1 IEPOX-SOA in a SE US forest during SOAS, 2013**

We use the SOAS-CTR field study (SE US-CTR) as an example for the determination of 211 IEPOX-SOA from AMS data via PMF analysis. The time series and mass spectrum of this 212 213 component are shown in Fig. 2. The IEPOX-SOAPMF mass concentration is the sum of mass 214 concentrations of all the ions in the IEPOX-SOAPMF mass spectra. The "mass concentration" of 215 an ion is used to represent the mass of the species whose detection resulted in the observed ion 216 current of that ion, based on the properties of electron ionization (Jimenez et al., 2003). An uncertainty (standard deviation) of IEPOX-SOA<sub>PMF</sub> mass concentration of ~9% was estimated 217 from 100 bootstrapping runs in PMF analysis (Ulbrich et al., 2009) (Fig. S1). This uncertainty 218 concerns only the PMF separation method. In practice the uncertainty in IEPOX-SOAPMF 219 concentration is dominated by the larger uncertainty on the AMS concentrations arising from the 220 221 collection efficiency and relative ionization efficiency (Middlebrook et al., 2012).

A strong correlation is found between AMS IEPOX-SOA<sub>PMF</sub> and 2-methyltetrols (R=0.79) and sulfate (R = 0.75) as expected (Surratt et al., 2010;Lin et al., 2012;Nguyen et al., 2014;Xu et

224	al., 2014). The diurnal variation of IEPOX-SOA <sub>PMF</sub> is also similar to gas-phase IEPOX and
225	isoprene measured in SOAS-CTR. 2-Methyltetrols, measured on-line by GC-EI/MS with the SV-
226	TAG instrument (Isaacman et al., 2014), comprise 26% of IEPOX-SOA <sub>PMF</sub> in SOAS-CTR on
227	average, as shown in Fig. 2b. A similar ratio (29%) is found between 2-methyltetrols measured
228	by offline analysis of filter samples using GC-EI/MS and LC/MS (Lin et al., 2014) and IEPOX-
229	SOA <sub>PMF</sub> . Other IEPOX-SOA tracers, such as C5-alkene triols, IEPOX-organosulfates, and
230	dimers containing them, can also be measured by offline GC-EI/MS and LC/MS (Lin et al.,
231	2014;Budisulistiorini et al., 2015), and they account for 28% and 24% in total IEPOX-SOA <sub>PMF</sub>
232	in SOAS (R=0.7), respectively (Fig. S2). The total IEPOX-SOA tracers measured in SOAS
233	account for ~78 $\pm$ 42% of the total IEPOX-SOA <sub>PMF</sub> mass concentration. The uncertainty of the
234	fraction of IEPOX-SOA molecular tracers in IEPOX-SOA <sub>PMF</sub> in SOAS study (42%) is estimated
235	by combining the overall uncertainty from IEPOX-SOA molecular tracer measurement (24%),
236	linear regression between tracer vs IEPOX-SOA <sub>PMF</sub> (17%, see Fig. 2b and Fig. S2), IEPOX-
237	SOA <sub>PMF</sub> in PMF separation method (9%) and the quantification of IEPOX-SOA <sub>PMF</sub> based on
238	AMS calibration (30%) (Middlebrook et al., 2012). This is a remarkably high value compared to
239	the tracer to total SOA ratios for other SOA systems (e.g., SOA from monoterpenes or aromatic
240	hydrocarbons) (Lewandowski et al., 2013). A total tracers to IEPOX-SOAPMF ratio of 26% was
241	reported for the Look Rock site in SOAS (SOAS-LR) (Budisulistiorini et al., 2015). Thus, the
242	measured total molecular tracer fraction in total IEPOX-SOA appears to be quite variable (a
243	factor of 3) even if the same or similar techniques are used. Although the calibration
244	methodology between different campaigns may result in some uncertainties, this value likely
245	changes significantly between different times and locations, potentially due to changes in

particle-phase reaction conditions such as sulfate and water concentrations, acidity, and theidentity and concentrations of oligomerization partners.

248	IEPOX-SOA <sub>PMF</sub> accounts for 17% of the total OA mass concentration at SOAS-CTR. This is
249	shown in Fig. 1 along with the IEPOX-SOA <sub>PMF</sub> fraction from several previous studies (Robinson
250	et al., 2011;Slowik et al., 2011;Budisulistiorini et al., 2013;Hayes et al., 2013;Hu et al.,
251	2013; Chen et al., 2014; Hu et al., 2015). Fig. 1 also shows the surface gas-phase IEPOX
252	concentrations for July, 2013 as simulated with GEOS-Chem. At all sites with at least ~30 ppt
253	predicted average IEPOX concentration, IEPOX-SOAPMF is identified in AMS data. IEPOX-
254	$SOA_{PMF}$ accounts for 6% – 36% of total OA in those studies, signifying the importance of
255	IEPOX-SOA for regional and global OA budgets. No IEPOX-SOAPMF factor (i.e. below the
256	PMF detection limit of ~5% of OA, Ulbrich et al., 2009) was found in areas strongly influenced
257	by urban emissions where high NO concentrations suppress the IEPOX pathway, even in the
258	presence of substantial isoprene concentrations (e.g. Hayes et al., 2013). GEOS-Chem indeed
259	predicts negligible modeled gas-phase IEPOX concentrations in those areas, where isoprene
260	peroxy radicals are expected to react primarily with NO. Some IEPOX can also be formed via
261	high NO chemistry (Jacobs et al., 2014), however this pathway is thought to be much smaller
262	than the low-NO pathway, consistent with the lack of observation of IEPOX-SOA $_{PMF}$ in the
263	polluted studies included here. The fraction of IEPOX-SOAPMF positively correlates with
264	modeled gas-phase IEPOX, as shown in the inset of Fig. 1.
265	The mass spectrum of IEPOX-SOA during SOAS-CTR is similar to those from other studies
266	as seen in Fig. S3 – S4 (Robinson et al., 2011;Lin et al., 2012;Budisulistiorini et al., 2013;Chen

et al., 2014;Nguyen et al., 2014;Xu et al., 2014), and also exhibits a prominent  $C_5H_6O^+$  peak at

268 m/z 82. We investigated the correlation between the time series of IEPOX-SOA<sub>PMF</sub> and each ion

in the OA spectra. The temporal variation of ion  $C_5H_6O^+$  correlates best (R=0.96) with IEPOX-SOA<sub>PMF</sub> among all measured OA ions (Table S1). This result suggests that  $C_5H_6O^+$  ion may be the best ion tracer for IEPOX-SOA among all OA ions.  $C_5H_5O^+$  (m/z 81),  $C_4H_5^+$  (m/z 53),  $C_4H_6O^+$  (m/z 70) and  $C_3H_7O_2^+$  (m/z 75) also correlate well with IEPOX-SOA<sub>PMF</sub> in SOAS-CTR and could be potential tracers for IEPOX-SOA<sub>PMF</sub>. Scatter plots between these four ions and  $C_5H_6O^+$  at different campaigns indicate they either have higher background values or lower signal-to-noise compared to  $C_5H_6O^+$  (Fig. S5).

 $f_{C_5H_6O}^{IEPOX-SOA}$  from SOAS and other field and laboratory studies (Table 1) ranges from 12‰ to 40‰ (‰=parts per thousand) and have an average value of 22±7‰. The average  $f_{C_5H_6O}^{IEPOX-SOA}$ value shown here also includes  $f_{82}$  data from four UMR IEPOX-SOA<sub>PMF</sub> spectra. This is justified since C<sub>5</sub>H<sub>6</sub>O<sup>+</sup> accounts for over 95% of m/z 82 in IEPOX-SOA based on results from SOAS-CTR and other lab studies (Kuwata et al., 2015). Indeed the average does not change if the UMR studies are removed from the average. These values are substantially higher than those from other types of OA or from locations with little impact from IEPOX-SOA, as discussed below.

## 3.2 $f_{C_5H_60}$ in areas with strong influence from urban and biomass burning emissions

We next examine whether POA or SOA from field studies in areas strongly influenced by urban and biomass-burning emissions and without substantial predicted gas-phase IEPOX concentrations or IEPOX-SOA contributions can lead to enhanced  $f_{C_5H_6O}^{OA}$ . Figure 3a shows the distribution of  $f_{C_5H_6O}^{OA}$  in this category of studies peaks at 1.7±0.1‰ (range 0.02 – 3.5‰). Data from continental air masses sampled from aircraft over the western and northwest US (where isoprene emissions are low) are shown in Fig. 3b and show a similar range as the polluted ground sites. Biomass burning emissions and plumes sampled over multiple studies show a similar range to the pollution studies, with some slightly higher values. The peak of the distribution of  $f_{C_5H_6O}^{OA}$ from fresh biomass-burning smoke across many different biomasses during the FLAME-3 study is 2.0‰. During the SEAC4RS aircraft campaign, many biomass burning plumes were sampled, where OA concentrations varied over a wide range (several tens to more than one thousand µg m<sup>-3</sup>). The average  $f_{C_5H_6O}^{OA}$  across these biomass-burning plumes was 1.75‰ with low variability (~20%), see Fig. S6.

298 We also explore whether other anthropogenic primary OA (POA) emission sources could elevate  $f_{C_5H_6O}$  above the observed background levels of ~1.7‰. Figure 3c shows  $f_{C_5H_6O}$  for POA 299 spectra from vehicle exhaust, cooking, coal combustion, and multiple pure chemical standards 300 (e.g., some alcohols; di- or poly acids) (Canagaratna et al., 2015). Almost all the values are 301 302 below 2‰, with exceptions for one type of cooking POA at 3‰, the polyol xylitol (4.2‰), and 303 some acids (5-Oxoazelaic acid= 4.8‰, Gamma ketopimelic acid = 5.2‰, ketopimelic acid = 304 6.5‰, 3-Hydroxy-3-Methylglutaric acid = 11.8‰, Adipic acid = 16.4‰). All the tracers resulting in elevated  $f_{C_5H_6O}$  contain multiple hydroxyl groups, and may result in furan-like 305 306 structures via facile dehydration reactions (Canagaratna et al., 2015). Xylitol has been proposed 307 as a tracer of toluene SOA (Hu et al., 2008). It has a similar structure to 2-methyltetrols, with 5 -308 OH groups instead of 4. In the AMS, xylitol may form the methylfuran structure through dehydration reactions like 2-methytetrols. However,  $f_{C_5H_6O}$  in other toluene SOA tracers in our 309 dataset show background levels of  $f_{C_5H_6O}$  (<2‰). Given the small fraction of xylitol in toluene 310 SOA (Hu et al., 2008), xylitol is unlikely to increase  $f_{C_5H_6O}$  in anthropogenic SOA, consistent 311 312 with our results.

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In summary, in the absence of strong impacts from biogenic SOA, the AMS high resolution ion C<sub>5</sub>H<sub>6</sub>O<sup>+</sup> has a clear and stable background, spanning a small range (0.02 - 3.5%) with an average values around 1.7±0.1‰ ( $f_{C_5H_6O}^{OA-Bkg-UB}$ ), about an order of magnitude lower than the average value ( $22\pm7\%$ ) of  $f_{C_5H_6O}^{IEPOX-SOA}$ .

## 317 **3.3 Enhancements of** $f_{C_5H_60}$ in areas strongly influenced by isoprene emissions

GEOS-Chem predicts much higher surface gas-phase IEPOX concentrations over the SE US 318 319 and Amazon rainforest than those in temperate urban areas (Fig. 1). This is expected from high 320 isoprene concentrations (e.g. 3.3 ppb in SOAS-CTR and 4 ppb in the Amazon) under low 321 average NO concentrations (~0.1 ppb) (Karl et al., 2009; Ebben et al., 2011). Probability distributions of  $f_{C_5H_6O}^{OA}$  during both campaigns are shown in Fig. 4a, and are very similar with 322 323 averages of 5 - 6% (range 2.5% - 11%). The Amazon forest downwind of Manaus and a 324 Borneo tropical forest study show even higher averages of 7‰ and 10‰, respectively (Robinson et al., 2011; de Sá et al., 2015). During the SEAC4RS aircraft campaign, the average  $f_{C_5H_6O}^{OA}$ 325 326  $(4.4\pm1.6\%)$  from all SE US flights is also enhanced compared to levels observed in the 327 northwest and western US continental air masses (1.7±0.3‰) where isoprene emissions are 328 much smaller (Guenther et al., 2012). Thus, campaigns in locations strongly influenced by isoprene emissions under lower NO conditions show systematically higher  $f_{C_5H_6O}^{OA}$  values (with 329 330 an average peak of  $6.5\%\pm2.2\%$ ) than background levels found in other locations (1.7%). The fact that  $f_{C_5H_60}^{OA}$  (6.5±2.2‰) in these studies is lower than the values in IEPOX-SOA (22‰±7‰) 331 is expected, since ambient datasets also include OA from other sources, and confirms that 332 333 IEPOX-SOA is not an overwhelmingly dominant OA source at most of those locations (See Fig. S7). 334

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## 335 **3.4** Values of $f_{C_5H_6O}$ in laboratory studies of non IEPOX-derived isoprene SOA

We also investigate  $f_{C_5H_6O}$  in laboratory SOA from isoprene in Fig. 4a. For SOA produced 336 by chamber isoprene photooxidation under high NO<sub>x</sub> conditions, low  $f_{C_5H_6O}$  (<2‰) within the 337 338 background level is observed (Kroll et al., 2006;Chen et al., 2011). SOA from oxidation of 339 isoprene hydroxyhydroperoxide (ISOPOOH, a product of low-NO oxidation of isoprene) under low-NO conditions, when formed under conditions that are not favorable for the reactive uptake 340 of IEPOX into aerosols also has low  $f_{C_5H_6O}$  of 2‰ (Krechmer et al., 2015). Low values of 341  $f_{C_5H_6O}$  (<3‰) are also observed in SOA from isoprene + NO<sub>3</sub> radical reactions without acid 342 seeds (Ng et al., 2008). The low  $f_{C_5H_6O}$  (<3‰) observed in non IEPOX-derived isoprene SOA 343 indicate that  $f_{C_5H_6O}$  is specifically enhanced from IEPOX-SOA, and is not a tracer for all SOA 344 from isoprene. 345

## 346 **3.5 Enhancements of** $f_{C_5H_6O}$ in areas strongly influenced by monoterpene emissions

347 The BEACHON-RoMBAS campaign was carried out in a Rocky Mountain pine forest with high monoterpene emissions that account for 34% in daytime and 66% at night of the total VOC 348 mixing ratios (on average peaking at 0.15 ppb during day and 0.7 ppb at night) (Fry et al., 2013) 349 350 but lower isoprene emissions (peaking at 0.35 ppb during daytime) (Kaser et al., 2013;Karl et al., 2014). One-third of the RO<sub>2</sub> radicals react via the low-NO route (i.e. via  $RO_2 + HO_2$ ) at this site 351 352 (Fry et al., 2013). The isoprene/monoterpene ratio at the Rocky Mountain site is 0.48, and is  $\sim 10$ 353 -20 times lower than the value (4.7) in SOAS-CTR and (8.3) in Amazon studies (Chen et al., 2014), suggesting that  $f_{C_5H_6O}^{OA}$  may be near background levels because of the very low potential 354 contribution of IEPOX-SOA at the Rocky Mountain site. However, the average  $f_{C_5H_6O}^{OA}$  at the 355 Rocky Mountain site is 3.7±0.5‰ (Fig. 4a), which although lower than the average  $f_{C_5H_6O}^{OA}$ 356

357 (6.5‰) found in the SE US-CTR, Amazon and Borneo forests, it is still twice the

358  $f_{C_5H_6O}^{OA-Bkg-UB}$  values of 1.7‰ observed in pollution and smoke-dominated locations.

Three circumstances may lead to such an enhanced  $f_{C_5H_6O}^{OA}$  at the Rocky Mountain site, which 359 360 we examine here. (1) A small amount of IEPOX-SOA may be formed from the limited isoprene present at the Rocky Mountain site and surrounding region. However, the average isoprene 361 362 concentration in this pine forest area is only 0.2 ppb, which is around 16 times less than that (3.3 ppb) at the SE US site in SOAS. The conditions at the Rocky Mountain site were less favorable 363 364 for IEPOX-SOA formation due to a higher fraction (70% in daytime) of the RO2 radicals reacting with NO and less acidic aerosols (Fry et al., 2013;Levin et al., 2014). Thus we can 365 estimate an upper limit contribution of IEPOX-SOA to the  $f_{C_5H_6O}^{OA}$  tracer at the Rocky Mountain 366 site assuming the same ratio of IEPOX-SOA to isoprene in both campaigns. In this case, we 367 would expect  $f_{C_5H_60}^{OA}$  at the Rocky Mountain site to be the background level (1.7‰) plus 1/16<sup>th</sup> of 368 the enhancement above the background observed in SOAS (5% - 1.7% = 3.3%) multiplied by 369 the ratio of OA concentrations at both sites (4.8 µg m<sup>-3</sup> in SE US site vs 1.8 µg m<sup>-3</sup> in Rocky 370 Mountain site). This calculation results in an expected upper limit  $f_{C_5H_6O}^{OA} \sim 2.25\%$  at the Rocky 371 372 Mountain site due to the IEPOX-SOA contribution. This estimate is much lower than the observed average 3.7‰. Thus the elevated  $f_{C_5H_6O}^{OA}$  in Rocky Mountain pine forest is very unlikely 373 to be due to IEPOX-SOA. 374 (2) The second explanation of high  $f_{C_5H_6O}^{OA}$  observed at Rocky Mountain site is that SOA from 375 monoterpene oxidation (MT-SOA) may have a higher  $f_{C_5H_6O}$  than background OA from other 376

377 sources. Several chamber studies show that MT-SOA, e.g., SOA from ozonolysis (Chhabra et al.,

- 378 2011; Chen et al., 2014) or photooxidation (Ng et al., 2007) of  $\alpha$ -pinene, or NO<sub>3</sub> reaction with  $\alpha$ -
- pinene, or NO<sub>3</sub> reaction with α-pinene, β-pinene and  $\Delta^3$ -Carene (Fry et al., 2014;Boyd et al.,

2015) can result in higher  $f_{C_5H_6O}$  (average 5.5±2.0‰) than background levels of ~1.7‰ (Fig. 4a). 380 We note that the average lab-generated MT-SOA value  $(f_{C_5H_6O}^{MT-SOA})$  is still 4 times lower than the 381 average  $f_{C_5H_6O}^{IEPOX-SOA}$  for IEPOX-SOA<sub>PMF</sub> and IEPOX-SOA<sub>lab</sub> (Fig. S8), and thus there is some 382 room to separate both contributions. Oxidation of monoterpenes can lead to species with multiple 383 384 -OH groups, which may result in the production of methylfuran (or ions of similar structure) 385 upon AMS analysis. We do not observe enhanced  $f_{C_{cH_6O}}$  in SOA from sesquiterpene oxidation (<2‰) (Chen et al., 2014). The values of  $f_{C_5H_6O}^{MT-SOA}$  in chamber studies, together with the finding 386 387 of a substantial contribution of monoterpenes to SOA at this Rocky Mountain site (Fry et al., 2013) suggest that MT-SOA may explain the values of  $f_{C_5H_6O}^{OA}$  observed there. 388 Two other field studies support the conclusion that ambient MT-SOA may have slightly 389 enhanced  $f_{C_5H_6O}$ . Fig. 6 shows data from a DC3 aircraft flight in the areas around Missouri and 390 Illinois. Ambient  $f_{C_5H_6O}^{OA}$  increases from background levels (~1.7‰) to ~4.1‰ in a highly 391 correlated manner to monoterpene concentration increases (with an average of 3.0% during the 392 393 enhanced period). Meanwhile, isoprene and gas-phase IEPOX stay at low levels similar to the rest of the flight, indicating that enhanced  $f_{C_5H_6O}^{OA}$  in the periods with higher MT concentrations 394 395 should arise from MT-SOA and not IEPOX-SOA. Fig. 4a includes AMS measurements at a MT-396 emission dominated European boreal forest (Hyytiälä in Finland) (Robinson et al., 2011). Average  $f_{C_5H_6O}^{OA}$  is ~2.5‰ at this site, which is again higher than the  $f_{C_5H_6O}^{OA-Bkg-UB}$  value of 1.7‰. 397 The slightly lower  $f_{C_5H_6O}^{OA}$  in the Boreal forest vs. the Rocky Mountain site may be partially 398 explained by a small contribution from IEPOX-SOA at the latter (estimated above to increase 399  $f_{C_5H_6O}^{OA}$  up to 2.25‰ at the Rocky Mountain site), as well as by differences of the MT-SOA/OA 400

ratio at both sites (Corrigan et al., 2013) and the relative importance of different MT species and
oxidation pathways.

(3) The enhanced  $f_{C_5H_6O}^{OA}$  at the Rocky Mountain site may have arisen from oxidation 403 products of 2-methyl-3-buten-2-ol (MBO, C<sub>5</sub>H<sub>10</sub>O) emitted from pine trees. MBO, with a 404 daytime average of 2 ppb accounts for ~50% of the total VOC mixing ratio during the day (Karl 405 et al., 2014). MBO has been shown to form aerosol with a 2-7 % yield in chamber studies, 406 407 which is thought to proceed via the uptake of epoxide intermediates ( $C_5H_{10}O_2$ , vs. IEPOX 408 C<sub>5</sub>H<sub>10</sub>O<sub>3</sub>) under acidic aerosol conditions (Zhang et al., 2012;Mael et al., 2014;Zhang et al., 409 2014). Some aerosol species formed by MBO-derived epoxides have similar structures (e.g.,  $C_5H_{12}O_3$ ) to the IEPOX oxidation products in SOA and thus they might contribute to  $f_{C_5H_6O}^{OA}$ . No 410 pure MBO-derived epoxides or their oxidation products in the aerosol phase have been measured 411 by AMS so far, to our knowledge. 412 To attempt to differentiate whether MT-SOA or MBO-SOA dominate the higher  $f_{C_5H_6O}^{OA}$  at 413 the Rocky Mountain site, average diurnal variations of ambient  $f_{C_5H_6O}^{OA}$ , monoterpene and 414 isoprene+MBO are plotted in Fig. S9.  $f_{C_5H_6O}^{OA}$  shows a diurnal pattern that increases at night and 415 peaks in the early morning, similar to the diurnal variation of monoterpenes. Monoterpenes 416 continue to be oxidized during nighttime at this site by NO<sub>3</sub> radical and O<sub>3</sub> with a lifetime of ~30 417 418 min (with 5 ppt of NO<sub>3</sub> and 30 ppb of O<sub>3</sub>) (Fry et al., 2013). In contrast only a decrease and later a plateau of  $f_{C_5H_6O}^{OA}$  are observed during the period with high MBO concentration and higher 419 oxidation rate of MBO due to high OH radical in daytime (as MBO reacts slowly with O<sub>3</sub> and 420 NO<sub>3</sub>) (Atkinson and Arey, 2003). While MBO-SOA may or may not have  $f_{C_5H_6O}$  above 421 background levels, the diurnal variations point to MT-SOA playing a dominant role in  $f_{C_5H_6O}^{OA}$  at 422 this site. 423

18

The average  $f_{C_5H_6O}^{OA}$  in areas strongly influenced by monoterpene emissions is 3.1±0.6‰, obtained by averaging the values from the Rocky mountain forest (3.7‰), European boreal forest (2.5‰), and DC3 flight (3.0‰). Note that the difference between  $f_{C_5H_6O}^{OA}$  in areas strongly influenced by monoterpene emissions (3.1±0.6‰) and isoprene emissions (6.5±2.2‰) is reduced, compared to a factor of 4 differences between pure MT-SOA (5.5±2.0‰) and IEPOX-SOA (22±7‰). This is likely due to the physical mixing of OA from different sources and in different proportions at each location.

## 431 **3.6** $f_{C_5H_60}$ vs OA oxidation level ( $f_{CO_2}$ ) "triangle plot" – background studies

432 In AMS spectra, the  $CO_2^+$  ion is a marker of aging and oxidation processes (Alfarra et al., 2004;Ng et al., 2011a). To evaluate whether oxidation plays a role on the observed  $f_{C_5H_6O}$  for 433 different types of OA, in this section we use plots of  $f_{CO_2}$  (= CO<sub>2</sub><sup>+</sup>/OA) vs.  $f_{C_5H_6O}$  as a graphical 434 diagnostic of this process, similar to graphical diagnostics ("triangle plots") used for other 435 purposes with AMS data (Cubison et al., 2011;Ng et al., 2011a). For studies strongly influenced 436 by urban and biomass-burning emissions in Fig. 3d we observe a wide range of  $f_{CO_2}^{OA}$  values from 437 0.001 to 0.3 (= 30% or 300‰). The wide range of  $f_{CO_2}^{OA}$  is due to variable fractions of POA and 438 439 SOA (mixing effect) and a variable oxidation level of POA and SOA (oxidation effect) in the different studies. In fact, to our knowledge, these studies encompass the values of  $f_{CO_2}^{OA}$  observed 440 in all ambient AMS studies to date (Ng et al., 2011a). Several studies when urban and forest air, 441 442 or biomass burning smoke were aged by intense OH oxidation with an oxidation flow reactor (OFR) (Kang et al., 2007;Li et al., 2013;Ortega et al., 2013) are also included. However, despite 443 the wide range of  $f_{CO_2}^{OA}$ ,  $f_{C_5H_6O}^{OA}$  changes little, staying in the range 0.02 – 3.5‰, and with little 444 apparent dependence on  $f_{CO_2}^{OA}$  for the ambient studies. A linear regression to quantiles from this 445

446 dataset results in an intercept of 1.7‰ and a very weak decrease with increasing  $f_{CO_2}^{OA}$ . A stronger 447 decrease is observed when aging urban air (Los Angeles) by intense OH exposure in flow 448 reactor, as shown in Fig. 3d.

Ambient  $f_{C0_2}^{OA}$  at the Rocky Mountain forest site shows a moderate oxidation level (0.1 – 0.15), similar to the SE US-CTR (Fig. 5).  $f_{C_5H_6O}^{OA}$  in the Rocky mountain site decreases linearly when  $f_{C0_2}^{OA}$  increases. During the Rocky Mountain study, the intense OH aging of ambient air in a flow reactor shows a continuation of the trend observed for the ambient data, where  $f_{C_5H_6O}^{OA}$ decreases as  $f_{C0_2}^{OA}$  increases. A linear regression to the combined ambient and OFR datasets  $(f_{C_5H_6O}^{OA} = -0.013 \times f_{C0_2}^{OA} + 0.0054)$  will be used below to estimate background  $f_{C_5H_6O}^{OA}$  in areas with strong monoterpene and low isoprene emissions.

456  $f_{C_5H_6O}$  in ambient SOA from other studies catalogued in the HR-AMS spectral database are 457 also shown in Fig. 5. Most urban oxygenated OA (OOA) are within  $f_{C_5H_6O}^{OA-Bkg-UB}$  (average 1.7%; 458 range: 0.02 – 3.5%), which is consistent with the  $f_{C_5H_6O}$  (<3%) in lab aromatic SOA and other 459 urban OA in Fig. 5. However, some ambient SOA spectra do show higher  $f_{C_5H_6O}$  (3 – 10%) than 460 the  $f_{C_5H_6O}^{OA-Bkg-UB}$  (0.02 – 3.5%), which we will discuss in the next section.

# 461 3.7 $f_{C_5H_60}$ vs. OA oxidation level ( $f_{CO_2}$ ) – IEPOX-SOA influenced Studies

462  $f_{CO_2}^{OA}$  vs.  $f_{C_5H_6O}^{OA}$  in studies impacted by IEPOX-SOA are shown in Fig. 5. Consistent with the 463 distributions discussed above, the bulk of points from these areas all show distinctively enhanced 464  $f_{C_5H_6O}^{OA}$  when compared to background  $f_{C_5H_6O}^{OA}$  points of similarly moderate or higher oxidation 465 levels. The  $f_{C_5H_6O}^{OA}$  measurements with lower  $f_{CO_2}^{OA}$  values are more broadly distributed than the 466  $f_{C_5H_6O}^{OA}$  points with higher  $f_{CO_2}^{OA}$  values in SE US-CTR, SEAC4RS, Borneo forest and Amazon

forest downwind of Manaus. However, increased  $f_{C_5H_6O}^{OA}$  with higher  $f_{CO_2}^{OA}$  was observed in the 467 468 Amazon. Both oxidation and mixing of airmasses with different OA can influence these observations.  $f_{C_5H_6O}^{IEPOX-SOA}$  in IEPOX-SOA usually will decrease with oxidative aging. E.g., 469  $f_{C_5H_60}^{OA}$  from the SOAS oxidation flow reactor decreases continuously as OA becomes more 470 oxidized than ambient OA in SOAS-CTR ( $f_{CO_2}^{OA}$  increases from 0.15 to 0.3). Airmass mixing 471 effects are more complex. Depending on the  $f_{CO_2}^{OA}$  in the airmasses mixed with,  $f_{C_5H_6O}^{OA}$  in IEPOX-472 SOA-rich air can show positive, neutral or negative trends with increasing  $f_{CO_2}^{OA}$ . E.g., in pristine 473 Amazon forest, points with both lower  $f_{CO_2}^{OA}$  (<0.08) and  $f_{C_5H_6O}^{OA}$  (<8‰) values are thought to be 474 475 mainly caused by advection of POA from occasional local pollution. The overall trend for the ambient measurements in studies strongly influenced by isoprene 476 emissions (Fig. 5) is that those points cluster in a triangle shape and  $f_{C_5H_6O}^{OA}$  decreases as  $f_{CO_2}^{OA}$ 477 increases, as illustrated in Fig. S10. This "triangle shape" indicates that as the ambient OA 478 oxidation increases, the IEPOX-SOA signature is reduced, potentially by the ambient oxidation 479 480 processes or by physical mixing with airmasses containing more aged aerosols. Finally, points with higher  $f_{C_5H_6O}$  in OOA/aged OA are labeled with numbers in Fig. 5. The 481 sources of those labeled points are summarized in Table S2. OA from those studies are all 482 partially influenced by biogenic emissions. For example, during measurements of ambient OA in 483 the Central Valley of California (number 2), high isoprene emissions and acidic particles were 484 485 observed (Dunlea et al., 2009), suggesting that potential IEPOX-SOA formed in this area may explain the higher  $f_{C_5H_6O}^{OA}$  there. 486

487 **3.8 Best estimate of**  $f_{C_5H_60}$  in IEPOX-SOA

488 IEPOX-SOA from different field campaigns and chamber studies lay towards the right and on the bottom half of Fig. 5. IEPOX-SOA from chamber studies show systematically lower 489  $f_{CO_2}^{IEPOX-SOA}$  than ambient studies. This is likely explained by the lack of additional aging in the 490 laboratory studies, because all the lab IEPOX-SOA were measured directly after uptake gas-491 phase IEPOX onto acidic aerosol without undergoing substantial additional oxidation. 492 A wide range (12 – 40‰) of  $f_{C_5H_6O}^{IEPOX-SOA}$  is observed with an average of 22‰±7‰ in 493 ambient and lab IEPOX-SOA.  $f_{C_5H_6O}^{IEPOX-SOA}$  did not show a trend vs.  $f_{CO_2}^{IEPOX-SOA}$ . The IEPOX-494 SOA molecular tracer 3-MeTHF-3,4-diols has been shown to enhance the  $f_{C_5H_6O}$  in OA (Fig. 5) 495 (Lin et al., 2012; Canagaratna et al., 2015). Except 3-MeTHF-3,4-diols none of the other pure 496 IEPOX-derived polyols standards have been atomized and injected into the AMS system so far, 497 498 to our knowledge. We suspect other polyols such as 2-methyltetrols may also lead to such an 499 enhancement through dehydration reactions in the AMS vaporizer leading to methylfuran-type structures. The diversity of  $f_{C_5H_6O}^{IEPOX-SOA}$  in different studies is related with the variable content of 500 specific IEPOX-SOA molecular species that enhance  $f_{C_5H_6O}^{IEPOX-SOA}$  differently. The fractions of 501 molecular IEPOX-SOA species in total IEPOX-SOA<sub>PMF</sub> is plotted vs  $f_{C_5H_6O}^{IEPOX-SOA}$  in three 502 503 different studies in Fig. 7, which show a strong correlation between each other. The strong simultaneous variation of both quantities indicates that the diversity of  $f_{C_5H_6O}^{IEPOX-SOA}$  is very likely 504 explained by the variability of the molecules comprising IEPOX-SOA among different studies. 505 During one day in SOAS (June 26th, 2013), IEPOX-SOA<sub>PMF</sub> comprised 80 - 90% of total 506 OA (Fig. S11), possibly due to high sulfate concentrations favoring IEPOX-SOA formation. 507  $f_{C_5H_6O}^{OA}$  reached 25‰, which is similar to the 22‰ for the IEPOX-SOA<sub>PMF</sub> from this study, and 508 consistent with a slightly lower value for the average vs. freshest ambient IEPOX-SOA. Among 509 510 the chamber studies, the study of reactive uptake of isoprene-oxidation products into an acidic

seed is most similar to the full chemistry in real ambient environments (Liu et al., 2014), and reports similar  $f_{C_5H_6O}^{IEPOX-SOA}$  values (19‰). Hence, we propose an average  $f_{C_5H_6O}^{IEPOX-SOA}$  (22‰)

513 from both studies as the typical value of fresh IEPOX-SOA.

## 514 **3.9 Proposed Method for Real-Time Estimation of IEPOX-SOA**

515 So far, PMF of AMS spectra is the only demonstrated method for quantifying total IEPOX-

516 SOA concentrations. However, the PMF method is labor-intensive and requires significant

517 expertise, and may fail to resolve a certain factor when present in lower mass fractions (<5%). A

simpler, real-time method to estimate IEPOX-SOA would be useful in many studies, including

519 ground-based and aircraft campaigns.

520 We propose an estimation method for IEPOX-SOA based on the mass concentration of its 521 tracer ion  $C_5H_6O^+$ . To do this, we express the mass concentration of  $C_5H_6O^+$  as

522 
$$C_5 H_6 O_{total}^+ = C_5 H_6 O_{IEPOX-SOA, ambient}^+ + C_5 H_6 O_{background}^+.$$
(1)

523 Where,  $C_5H_6O_{total}^+$  is measured total  $C_5H_6O^+$  signal in AMS;  $C_5H_6O_{IEPOX-SOA,ambient}$  and 524  $C_5H_6O_{background}^+$  are the  $C_5H_6O^+$  signals contributed by IEPOX-SOA in ambient OA and other 525 background OA (non IEPOX-SOA).

526 Then, 
$$C_5H_6O_{IEPOX-SOA,ambient}$$
 and  $C_5H_6O^+_{background}$  can be calculated as:

527 
$$C_5 H_6 O^+_{IEPOX-SOA, ambient} = IEPOX-SOA \times f^{IEPOX-OA}_{C_5 H_6 O}.$$
 (2)

528 
$$C_5 H_6 O_{background}^+ = (OA_{mass} - IEPOX - SOA) \times f_{C_5 H_6 o}^{OA - Bkg}.$$
 (3)

529 Where,  $f_{C_5H_6O}^{IEPOX-OA}$  is the fractional contribution of C<sub>5</sub>H<sub>6</sub>O<sup>+</sup> to the total ion signal in the

- 530 spectra of IEPOX-SOA from IEPOX-SOA<sub>lab</sub> or IEPOX-SOA<sub>PMF</sub> factors.  $f_{C_5H_6O}^{OA-Bkg}$  is the
- background  $f_{C_5H_6O}$  in other non-IEPOX-SOA, e.g., values from OA strongly influenced by urban

and biomass-burning emissions 
$$(f_{C_5H_6O}^{OA-Bkg-UB})$$
.

533 Then, by combining Eq. (1) – (3), we can express  $C_5H_6O_{total}^+$  as:

534 
$$C_5 H_6 O_{total}^+ = IEPOX - SOA \times f_{C_5 H_6 O}^{IEPOX - OA} + (OA - IEPOX - SOA) \times f_{C_5 H_6 O}^{OA - Bkg}.$$
(4)

535 Finally, IEPOX-SOA can be estimated as:

536 
$$IEPOX-SOA = \frac{C_{5}H_{6}O_{total}^{\dagger} - OA \times f_{C_{5}H_{6}O}^{OA-Bkg}}{f_{C_{5}H_{6}O}^{IEPOX-OA} - f_{C_{5}H_{6}O}^{OA-Bkg}}.$$
 (5)

537 In Eq. (5),  $C_5 H_6 O_{total}^+$  and OA mass are measured directly by AMS.  $f_{C_5 H_6 O}^{OA-Bkg}$  and 538  $f_{C_5 H_6 O}^{IEPOX-OA}$  are two parameters that must be determined by other means.

As discussed above, the background value in the absence of a substantial impact of MT-539 SOA is ~1.7‰. In studies influenced by monoterpene emissions, the background value may be 540 elevated by MT-SOA.  $f_{C_5H_6O}^{OA}$  at the Rocky Mountain site estimated by  $f_{C_5H_6O}^{OA} = (0.41 - 10^{-10})^{-10}$ 541  $f_{CO_2}^{OA}$  × 0.013 (Fig. 5) can be used as  $f_{C_5H_6O}^{OA-Bkg}$  for areas with strong MT-SOA contributions 542  $(f_{C_{5}H_{6}O}^{OA-Bkg-MT})$ . There is some uncertainty in this value, due to possible contributions of a small 543 amount of IEPOX-SOA, MBO-SOA, and other OA sources at this site. An alternative estimate 544 for  $f_{C_5H_6O}^{OA-Bkg-MT}$  would be ~ 1.7‰ + 3×MT<sub>avg</sub> (ppb), which is also approximately consistent with 545 our ambient data, but may have higher uncertainty. Further characterization of the background 546  $f_{C_5H_6O}$  in areas with MT-SOA impact is of interest for future studies. Finally, we have decided to 547 use  $f_{C_5H_6O}^{OA}$  estimated from the Rocky Mountain site as  $f_{C_5H_6O}^{OA-Bkg-MT}$  in the following calculation. 548 As discussed above, we use average  $f_{C_5H_6O}^{IEPOX-OA} = 22\%$  in Eq. (3) as a representative value of 549 ambient IEPOX-SOA. Several scenarios based on different  $f_{C_5H_6O}^{OA}$  values to use this tracer-based 550 method are addressed in the supporting information. The justification from users on using this 551 method is needed. 552

553

## 3.10 Application of the Real-Time Estimation Method of IEPOX-SOA

554	To test the proposed estimation method, we use SE US forest (SOAS) data as an example in
555	Fig. 8, applying both background estimates (urban & biomass burning, and monoterpene
556	emissions). Since there are high monoterpene concentrations (~1 ppb during the night) in SOAS,
557	we expect the MT-influenced background to be more accurate. The IEPOX-SOA estimated by
558	subtracting the MT-SOA background (IEPOX-SOA $_{MT}$ ) is indeed better correlated with IEPOX-
559	$SOA_{PMF}(R=0.99)$ than that (R = 0.96) when the urban & biomass-burning background is applied
560	(IEPOX-SOA <sub>urb&amp;bb</sub> ). The intercept of regression line between IEPOX-SOA <sub>MT</sub> and IEPOX-
561	SOA <sub>PMF</sub> is zero, indicating the background of IEPOX-SOA contributed by MT-SOA is clearly
560	de duete d
562	deducted.
563	The regression slope between IEPOX-SOA <sub>MT</sub> and IEPOX-SOA <sub>PMF</sub> is 0.95, suggesting that
563	The regression slope between IEPOX-SOA <sub>MT</sub> and IEPOX-SOA <sub>PMF</sub> is 0.95, suggesting that
563 564	The regression slope between IEPOX-SOA <sub>MT</sub> and IEPOX-SOA <sub>PMF</sub> is 0.95, suggesting that $C_5H_6O^+$ in SE US CTR site (SOAS) may be slightly overcorrected by minimizing $C_5H_6O^+$ from
563 564 565	The regression slope between IEPOX-SOA <sub>MT</sub> and IEPOX-SOA <sub>PMF</sub> is 0.95, suggesting that $C_5H_6O^+$ in SE US CTR site (SOAS) may be slightly overcorrected by minimizing $C_5H_6O^+$ from monoterpene emissions. This underestimation may be associated with higher MT-SOA
563 564 565 566	The regression slope between IEPOX-SOA <sub>MT</sub> and IEPOX-SOA <sub>PMF</sub> is 0.95, suggesting that $C_5H_6O^+$ in SE US CTR site (SOAS) may be slightly overcorrected by minimizing $C_5H_6O^+$ from monoterpene emissions. This underestimation may be associated with higher MT-SOA contribution to $C_5H_6O^+$ in Rocky Mountain pine forest site than SE US forest site, or interference

570 Among all the datasets introduced in this study, the SOAS-CTR dataset should be the best 571 case scenario since  $f_{C_5H_6O}^{IEPOX-OA} = 22\%$  is coincidently the same value in the spectrum of IEPOX-572 SOA<sub>PMF</sub> in SOAS-CTR and a large fraction (17%) of IEPOX-SOA existed in SOAS-CTR as 573 well. Given the spread of values of  $f_{C_5H_6O}^{IEPOX-OA}$  (12 – 40‰) in different studies, if no additional 574 local IEPOX-SOA spectrum is available for a given site, the estimation from this method should be within a factor of ~2 of the actual concentration, as illustrated in Fig. S13-S14. Further information concerning the estimation method using unit mass resolution m/z 82 (or  $f_{82}$ ) can be found in the Appendix.

## 578 4. Conclusions

To investigate if the ion  $C_5H_6O^+$  (at m/z 82) in AMS spectra is a good tracer for IEPOX-SOA, 579 tens of field and lab studies are combined and compared, including the SOAS 2013 campaign in 580 the SE US. The results show that  $f_{C_5H_6O}^{OA}$  is clearly elevated when IEPOX-SOA is present, and 581 thus has potential usefulness as a tracer of this aerosol type. The average  $f_{C_5H_6O}^{IEPOX-OA}$  in 582 chamberand ambient studies is  $22\pm7\%$  (range 12% - 40%). No dependence of  $f_{C_5H_6O}^{IEPOX-OA}$  on 583 oxidation level ( $f_{CO_2}^{IEPOX-SOA}$ ) was found. Background  $f_{C_5H_6O}$  in OA strongly influenced by urban 584 or biomass-burning emissions or pure anthropogenic POAs averages 1.7±0.1‰ (range 0.02 -585 3.5‰). 586

In ambient OA that is strongly influenced by isoprene emissions under lower NO, we observe systematically higher  $f_{C_5H_6O}^{OA}$  (with an average of ~6.5±2.2‰), consistent with presence of IEPOX-SOA. Low tracer values ( $f_{C_5H_6O}$ <3‰) are observed in non IEPOX-derived isoprene-SOA from laboratory studies, indicating that the tracer ion is specifically enhanced from IEPOX-SOA, and is not a tracer for all SOA from isoprene.

Higher background values of  $f_{C_5H_6O}^{OA}$  (3.1±0.6‰ in average) were found in area strongly impacted by monoterpene emissions.  $f_{CO_2}^{MT-SOA}$  is 5.5±2.0‰, which are substantially lower than for IEPOX-SOA (22±7‰), and thus they leave some room to separate both contributions. A  $f_{C_5H_6O}^{OA-Bkg-MT}$  as a function of  $f_{CO_2}^{OA}$  in monoterpene emissions is determined by linear regressing 596 the  $f_{C_5H_6O}^{OA}$  and  $f_{CO_2}^{OA}$  at a Rocky Mountain pine forest site.

597	A simplified method to estimate IEPOX-SOA based on measured ambient $C_5H_6O^+$ , $CO_2^+$ and
598	OA in AMS is proposed. Good correlations (R>0.96) between estimated IEPOX-SOA and
599	IEPOX-SOA <sub>PMF</sub> are obtained for SOAS, confirming the potential usefulness of this estimation
600	method. Given the observed variability in IEPOX-SOA composition, the method is expected to
601	be within a factor of $\sim 2$ of the true concentration if no additional information about the local
602	IEPOX-SOA is available for a given study. When only unit mass resolution data is available as
603	in ACSM data, all methods may perform less well because of increased interferences from other
604	ions at $m/z$ 82.

## 606 ACKNOWLEDGMENT

607 This study was partially supported by NSF AGS-1243354 and AGS-1360834, NASA NNX12AC03G, DOE (BER/ASR) DE-SC0011105, and NOAA NA13OAR4310063. B. Palm 608 and J. Krechmer are grateful for fellowships from EPA STAR (FP-91761701-0 and FP-609 610 91770901-0) and CIRES. A. Ortega is grateful for a CU-Boulder Chancellor's and DOE SCGF (ORAU/ORISE) fellowship. A. Wisthaler and T. Mikoviny were supported by the Austrian 611 Federal Ministry for Transport, Innovation and Technology (BMVIT) through the Austrian 612 613 Space Applications Programme (ASAP) of the Austrian Research Promotion Agency (FFG), and the Visiting Scientist Program at the National Institute of Aerospace (NIA). G. Isaacman-614 VanWertz is grateful for an NSF Fellowship (DGE-1106400). UC Berkeley was supported by 615 NSF AGS-1250569. We acknowledge the logistical support from the LBA Central O ce at INPA 616 (Instituto Nacional de Pesquisas da Amazonia). P. Artaxo acknowledges support from FAPESP 617 grants 2013/05014-0 and 2014/05238-8 and CNPq support from grants 457843/2013-6 and 618 307160/2014-9. We acknowledges this work was funded by the U.S. Environmental Protection 619 620 Agency (EPA) through grant number 835404. The contents of this publication are solely the responsibility of the authors and do not necessarily represent the o cial views of the U.S. EPA. 621 Further, the U.S. EPA does not endorse the purchase of any commercial products or services 622 mentioned in the publication. The U.S. EPA through its O ce of Research and Development 623 collaborated in the research described here. It has been subjected to Agency review and approved 624 625 for publication, but may not necessarily reflect ocial Agency policy. The authors would also like 626 to thank the Electric Power Research Institute (EPRI) for their support. M. Riva and J. D. Surratt wish to thank the Camille and Henry Dreyfus Postdoctoral Fellowship Program in 627 Environmental Chemistry for their financial support. We thank J. Crounse and P. Wennberg 628 629 from Caltech for gas-phase IEPOX data in SOAS-CTR and DC3, under support from NASA NNX12AC06G. We thank Lu Xu and Nga Lee Ng from Georgia Tech for providing data from 630 631 their studies. We acknowledge funding from the UK Natural Environment Research Council

through the OP3 and SAMBBA projects (Grant refs. NE/D002117/1 and NE/J010073/1).

#### 633 APPENDIX

In addition to the preceding high resolution  $C_5H_6O^+$  data analysis, we also investigated unit mass 634 resolution (UMR) m/z 82 as a tracer of IEPOX-SOA. In addition to C<sub>5</sub>H<sub>6</sub>O<sup>+</sup> (m/z 82.0419), the 635 reduced ion  $C_6H_{10}^+$  and oxygenated ion  $C_4H_2O_2^+$  often contribute signal to UMR m/z 82. The 636 average background level of  $f_{82}^{OA}$  (= m/z 82/OA) is from 4.3±0.9‰ (0.01 to 10‰) in studies 637 638 strongly influenced by urban, biomass-burning and other anthropogenic POA, as shown in Fig. A1a – c. This value is higher than the high-resolution  $f_{C_5H_6O}^{OA-Bkg-UB}$  (1.7‰) in the same studies. 639 Background  $f_{82}^{OA}$  increases when OA is fresher (lower  $f_{44}$ ,  $f_{44}^{OA} = m/z$  44/OA) as shown in Fig A1d, 640 and can be estimated as  $f_{82}^{OA} = 5.5 \times 10^{-3} - 8.2 \times 10^{-3} \times f_{44}^{OA}$  in areas strongly impacted by urban and 641 642 biomass-burning emissions. The uncertainty of calculated  $f_{82}$  can be as high as 30% in the lower fresh OA plumes by considering the uncertainties from quantile average and linear regression. 643 There are also some pure chemical species that exhibit high  $f_{82}$  values, as shown in Fig. A1c. 644 These species include docosanol, eicosanol and oleic acid. However, none of these pure chemical 645 646 species alone contributes substantially to ambient aerosol.

The probability density distributions of  $f_{82}^{OA}$  in studies strongly influenced by isoprene emissions are shown in Fig. A2a. The peaks (~8.7±2.5‰) are similar in SE US, pristine, polluted Amazon forest, Borneo forest to high resolution  $f_{C_5H_6O}^{OA}$  (~6.5±2.2‰), indicating C<sub>5</sub>H<sub>6</sub>O<sup>+</sup> is the dominant ion at UMR *m/z* 82 in these studies. Compared to the studies with strong urban and biomassburning emissions, clear enhancements of  $f_{82}^{OA}$  in studies strongly influenced by isoprene emissions are still observed, but with less contrast than for in high resolution datasets (Fig. A2 – A3).

Figure 2Aa also shows the probability density distributions of  $f_{82}^{OA}$  at Rocky Mountain and European boreal forests (strongly influenced by monoterpene emissions). Those distributions peak at ~5‰, which are within the range (0.01 - 10‰) of  $f_{82}^{OA}$  in aerosols strongly influenced by urban and biomass-burning emissions. In the lab studies, most of  $f_{82}^{MT-SOA}$  (average 6.7±2.2‰; range 4 – 11‰) observed in the spectra of MT-SOA are also comparable to background  $f_{82}^{OA}$ levels (average 4.3±0.9‰; range 0.01–10‰), and tend to be in the higher  $f_{82}^{OA}$  region from urban and biomass-burning emissions. A linear regression line of  $f_{44}^{OA}$  vs  $f_{82}^{OA}$  for the Rocky Mountain site ( $f_{82}^{OA}$ =7.7×10<sup>-3</sup>–0.019×  $f_{44}^{OA}$ ) is used to estimate the background  $f_{82}^{OA}$  from areas strongly influenced by monoterpene emissions.

In summary, elevated  $f_{82}^{OA}$  in studies with high isoprene-emissions is observed. Pronounced  $f_{82}^{IEPOX-SOA}$  should be a key feature of IEPOX-SOA spectra. Thus IEPOX-SOA can be estimated as Eq. (6) here:

where  $f_{82}^{IEPOX-SOA}$  is 22‰ as obtained average (Fig. A3). In Eq. (4),  $f_{82}^{OA-Bkg}$  can be calculated as a function of  $f_{44}^{OA}$  in studies strongly influenced by urban and biomass-burning emissions  $(f_{82}^{OA}=5.5\times10^{-3}-8.2\times10^{-3}\times f_{44}^{OA})$  or monoterpene emissions  $(f_{82}^{OA}=7.7\times10^{-3}-0.019\times f_{44}^{OA})$ , as discussed earlier.  $m82_{total}$  and  $OA_{mass}$  are the measured ambient m/z 82 and OA mass concentrations by AMS. Because  $f_{82}$  in MT-SOA and OA from urban and biomass-burning emissions cannot be separated, only one background value of  $f_{82}^{OA-Bkg}$  will be used in the UMR method.

To test this UMR empirical method, we apply Eq. (6) to SOAS-CTR dataset, see Fig. A4. The

estimated IEPOX-SOA in SOAS-CTR from both background corrections (urban+biomass

burning vs monoterpene) both correlates well with IEPOX-SOA<sub>PMF</sub> with R=0.97 and R=0.98,

677 respectively. The regression slopes between estimated fresh IEPOX-SOA vs IEPOX-SOA<sub>PMF</sub> are 1.11 and 0.94, which are within 15% of 1:1 line. The deviation of estimated IEPOX-SOA from 678 UMR by subtracting the background of MT-SOA influences is similar to that from HR in the 679 680 SOAS dataset, indicating the UMR-based IEPOX-SOA estimation may perform as well as HR in areas with high IEPOX-SOA fractions. For areas with small IEPOX-SOA fractions, more 681 uncertainties may exist in UMR calculation, e.g., there are wider variations of  $f_{82}^{OA-Bkg}$  from 682 urban and biomass-burning emissions with oxidation level, whereas a smaller and less variable 683  $f_{C_5H_6O}^{OA-Bkg}$  is found in HR. Overall, m/z 82 in unit mass resolution data is also useful to estimate 684 IEPOX-SOA. The different methods to estimate IEPOX-SOA may perform less well because of 685 increased interferences from other ions at m/z 82, however at locations with very high fractions 686 687 of IEPOX-SOA such as SOAS-CTR, the UMR-based method performs well.

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**Table 1**. Datasets used in this study<sup>a</sup>. Ranges or average plus standard deviation of  $f_{C_5H_6O}$  (high resolution) and  $f_{82}$  (unit mass 1117

resolution) in different studies are also included. 1118

Name of datasets	Time Period	Site locations and descriptions	Campaign name	Ranges or average±std.de v. f <sub>C5H60</sub> (‰)	Ranges or average±std.de v. f <sub>82</sub> (‰)	References		
		Studies strongly-influenced by	isonrono omissions r	5 0	v. <i>J</i> 82 (700)			
SE US forest-CTR site	Jun-Jul, 2013	Centreville, AL,	SOAS	6.2±2.4	7.6±2.2	(1)		
Pristine Amazon	Feb-Mar,	Centrevine, AL,	SUAS	0.2±2.4	7.0±2.2	(1)		
forest 2008, Brazil	2008	Pristine rain forest site, TT34,	AMAZE-08	5.0±2.3	7.9±1.7	(2)		
Amazon forest downwind Manaus, Brazil	Feb-Mar, 2014	T3 site, near Manacapuru	GoAmazon2014/5	6.9±1.6	7.1±1.0	(3)		
Pristine Amazon forest 2014, Brazil	Aug- Dec,2014	T0 site, ~150 km northeast of Manaus	GoAmazon2014/5	N/A	5.6±1.7	(4)		
SE US	Aug-Sep, 2013	Aircraft measurement:	SEAC4RS	4.3±1.6	N/A	(5)		
Borneo forest, Malaysia	Jun-Jul, 2008	Rain forest GAW station, Sabah, Malaysia	OP3	10±0.3	12.4±0.4	(6)		
Atlanta, US	Aug-Sep, 2011	Urban JST site, Atlanta, Georgia, US	N/A	N/A	3.7±1.9	(7)		
Atlanta (JST), US	May, 2012	Urban JST site, Atlanta, Georgia, US	N/A	3.3±0.9	N/A	(8)		
Atlanta (GT), US	Aug, 2012	Urban Georgia Tech site, Georgia, US	N/A	5.4±1.9	N/A	(8)		
Yorkville, US	July, 2012	Rural sites, 80km northwest of JST site, Georgia, US	N/A	7.7±2.2	N/A	(8)		
Harrow, Canada	Jun-Jul, 2007	Harrow site, rural sites surrounded by farmland, Canada	BAQSMET	N/A	N/A	(9)		
Bear Creek, Canada	Jun-Jul, 2007	Bear Creek site, wetlands area surrounded by farmland, Canada	BAQSMET	N/A	N/A	(9)		
	Studies strongly-influenced by monoterpene emissions							

Rocky mountain pine	Jul-Aug,	Manitou Experimental Forest	BEACHON-	3.7±0.5	5.1±0.5	(10)
forest, CO, USA European Boreal forest,	2011	Observatory, CO, Hyytiala site in Pine forest,	RoMBAS EUCAARI			
Finland	2008-2009	Finland	campaign	2.5±0.1 <sup>b</sup>	4.8±0.1 <sup>b</sup>	(11)
Studies mixed-influenced by isoprene and monoterpene emissions						
North American temperate, US	Aug-Sep, 2007	Blodgett Forest Ameriflux Site, CA, US	BEARPEX	4.0±<0.1 <sup>b</sup>	4.0±<0.1 <sup>b</sup>	(11)
Studies strongly-influenced by urban emissions						
Los Angeles area , CA, USA	May-Jun, 2010	Pasadena, US	CalNex	1.6±0.2	3.6±0.5	(12)
Beijing, China	Nov-Dec, 2010	Peking University, in NW of Beijing city, China	N/A	1.5±0.3	4.6±0.7	(13)
Changdao island, Downwind of China	Mar-Apr, 2011	Changdao island, China	CAPTAIN	1.6±0.2	3.8±0.5	(14)
Barcelona area, Spain	Feb-Mar, 2009	Montseny, Spain	DAURE	1.6±0.2	4.8±0.9	(15)
Studies of biomass-burning smokes						
BB Chamber study	Sep– Oct, 2009	Missoula, MO, USA	FLAME-3	1.9±0.6	5.9±1.4	(16)
Biomass burning plumes	Aug-Sep, 2013	All over US, aircraft measurement	SEAC4RS	1.8±0.5	N/A	(6)
Biomass burning plumes	May-Jun, 2011	All over US, aircraft measurement	DC-3	1.8±0.4	N/A	(17)
		Conti	nental plumes			
NW US	Aug-Sep, 2013	Aircraft measurement	SEAC4RS	1.7±0.3	N/A	(16)
Western US	May-Jun, 2011	Aircraft measurement	DC-3	1.9±0.6	N/A	(17)
		OA from	specific sources			
IEPOX-SOA from ambient PMF factors and chamber studies.			22±7	22±7	(18)	
Isoprene derived non-IEPOX SOA (reaction with OH under conditions of high NO or low NO without seed not favorable for the reactive-uptake of IEPOX, reaction with NO3 without seed)				<3	<3	(19)

Monoterpene-derived SOA	$5.5 \pm 2.0$	$6.7 \pm 2.0$	(20)
Other SOA (not from isoprene and mononterpene)	$2.2 \pm 0.9$	6.1±2.1	(21)
Cooking	$1.5 \pm 0.8$	$8.2{\pm}1.1$	(22)
Coal combustion	1.4-2.0	N/A	(23)
Vehicle emission	1.1±0.6	5.1±1.1	(24)
Biomass burning	2.3±0.7	4.3±1.5	(25)
Pure chemical species	$0.7{\pm}1.0$	$4.0 \pm 5.5$	(26)

a- HR-ToF-AMS was used for all the campaigns except the Atlanta, US and Pristine Amazon forest 2014, Brazil using ACSM.
 b- Standard error

1121 (1) This study; (2) (Chen et al., 2014); (3) (de Sá et al., 2015); (4) (Carbone et al., 2015); (5)(Liao et al., 2014); (6) (Robinson et

1122 al., 2011); (7) (Budisulistiorini et al., 2013); (8) (Xu et al., 2014;Xu et al., 2015) (9) (Slowik et al., 2011); (10) (Ortega et al., 2014);

(11) (Robinson et al., 2011); (12) (Hayes et al., 2013); (13) (Hu et al., 2015); (13) (Hu et al., 2013); (15) (Minguillón et al., 2011);

1124 (16) (Ortega et al., 2013); (17) (Barth et al., 2014); (18) (Chhabra et al., 2011; Robinson et al., 2011; Budisulistiorini et al., 2013; Chen

1125 et al., 2014;Liu et al., 2014;Kuwata et al., 2015); (19) (Kroll et al., 2006;Ng et al., 2008;Krechmer et al., 2015); (20) (Bahreini et al.,

1126 2005; Chen et al., 2014; Boyd et al., 2015); (21) (Bahreini et al., 2005; Liggio et al., 2005; Chhabra et al., 2011; Loza et al., 2012); (22)

(Lanz et al., 2007; Mohr et al., 2009; He et al., 2010; Huang et al., 2010; Mohr et al., 2012; Crippa et al., 2013; Hu et al., 2015); (23)

(Hu et al., 2013;Hu et al., 2015); (24) (Canagaratna et al., 2004;Lanz et al., 2007;Sage et al., 2008;Aiken et al., 2009;Mohr et al.,

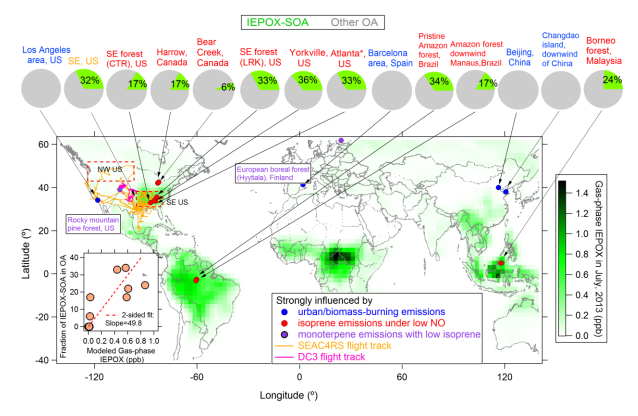
1129 2009; Chang et al., 2011; Docherty et al., 2011; Hersey et al., 2011; Ng et al., 2011b; Coggon et al., 2012; Mohr et al., 2012; Saarikoski

1130 et al., 2012;Setyan et al., 2012;Crippa et al., 2013); (25) (Schneider et al., 2006;Weimer et al., 2008;Aiken et al., 2009;He et al.,

1131 2010;Ng et al., 2011b;Schneider et al., 2011;Mohr et al., 2012;Saarikoski et al., 2012;Crippa et al., 2013;Hu et al., 201

1132 2015); (26) (Alfarra, 2004;Katrib et al., 2004;Phinney et al., 2006;Dzepina et al., 2007;Takegawa et al., 2007;Aiken et al., 2009;Li et

1133 al., 2011;Schneider et al., 2011)



## 1136

**Figure 1**. Locations of field campaigns used in this study. The IEPOX-SOA fractions of OA in different studies are shown in the pie charts on the top of graph. Site names are color-coded with

site types. Detailed information these studies can be found in Table 1. Note that the Atlanta pie

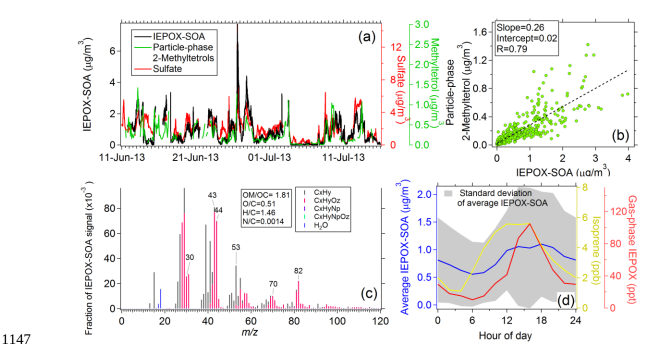
1140 chart was averaged by three urban datasets in Budisulistiorini et al. (2013) and Xu et al. (2015).

1141 The green background is color coded with modeled global gas-phase IEPOX concentrations for

1142July, 2013 from the GEOS-Chem model. The insert shows as scatter plot of observed average

1143 fraction of IEPOX-SOA in OA vs. GEOS-Chem modeled gas-phase IEPOX in various field

1144 campaigns.



1148 Figure 2. Results from the SOAS campaign in a SE US forested site. (a) Time series of IEPOX-

1149 SOA<sub>PMF</sub>, sulfate and particle-phase 2-methylterols (a key IEPOX uptake product) from on-line

1150 GC/MS; (b) Scatter plot between particle-phase 2-methylterols and IEPOX-SOA. (c) Mass

1151 spectrum of IEPOX-SOA; (d) Diurnal cycle of IEPOX-SOA, isoprene and gas-phase IEPOX

1152 (the latter measured by  $CF_3O^- CIMS$ ).

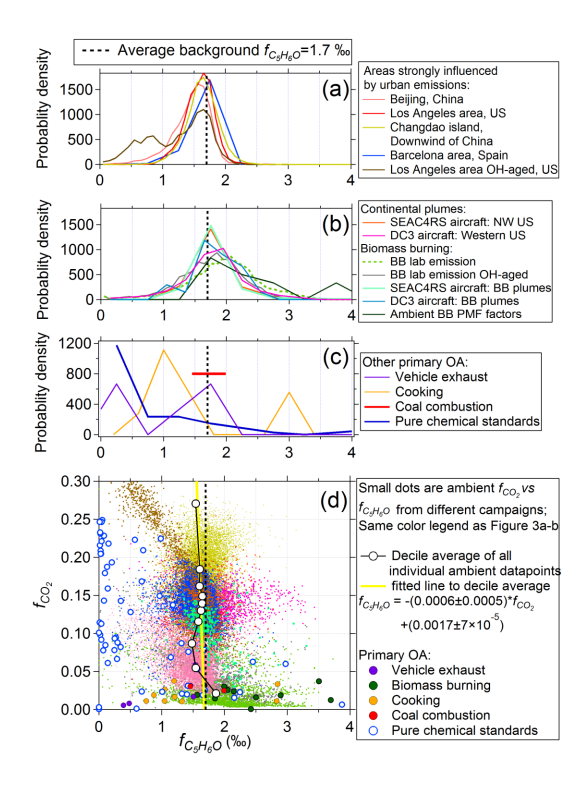
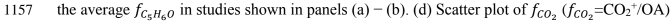
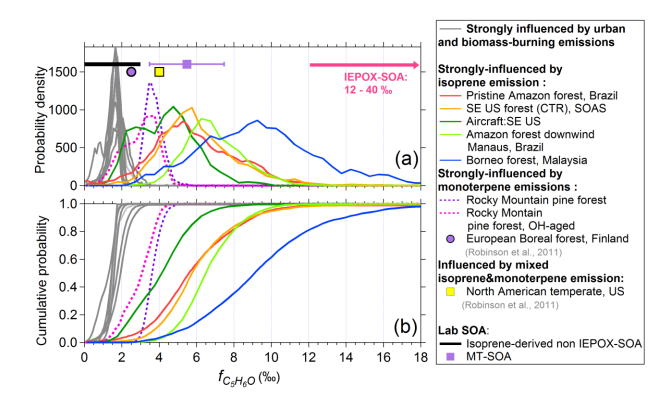


Figure 3. Probability density distributions of  $f_{C_5H_6O}$  in studies (a) strongly influenced by urban emissions; (b) continental air masses sampled from aircraft and biomass-burning emissions; (c) other anthropogenic primary OA sources and pure chemical standards. The dashed line (1.7‰) is



- 1158 vs.  $f_{C_5H_6O}$  for all studies shown in panels (a) (c), using the same color scheme. Quantile
- 1159 averages of  $f_{C_5H_6O}$  across all studies sorted by  $f_{CO_2}$  are also shown, as is a linear regression line to
- 1160 the quantile points.

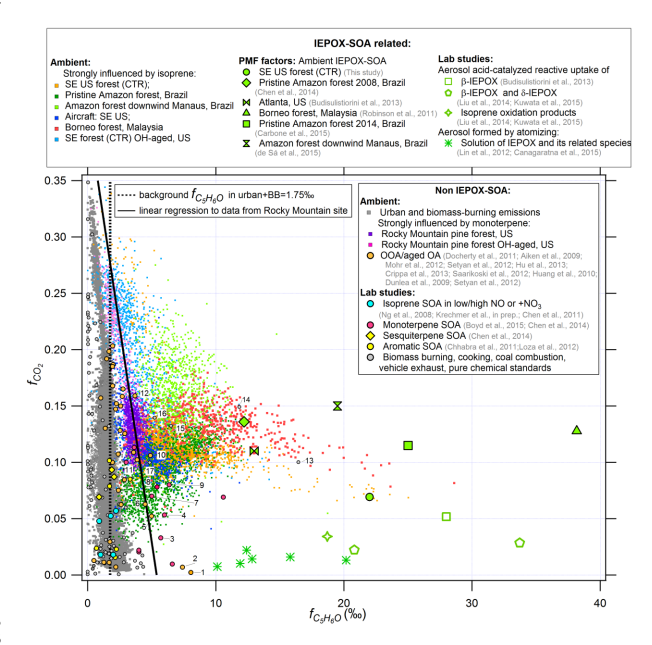


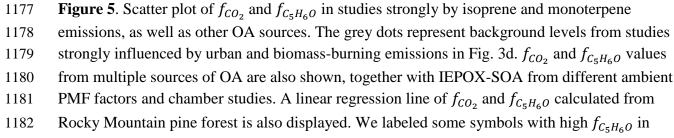
1164

**Figure 4**. (a) Probability density and (b) cumulative probability distributions of  $f_{C_5H_6O}^{OA}$  in studies strongly influenced by isoprene and/or monoterpene emissions. The ranges of  $f_{C_5H_6O}$  from other non IEPOX-derived isoprene-SOA and MT-SOA are also shown. The background grey lines are from studies strongly influenced by urban and biomass-burning emissions and are the same data from Fig. 3a – b. The arrow in Fig. 4a indicates the range of  $f_{C_5H_6O}^{IEPOX-SOA}$  between 12‰ (start of the arrow) to 40‰ which is beyond the range of x-axis scale.



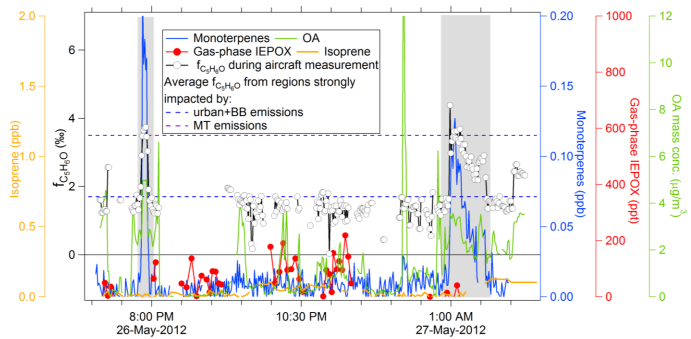






1183 numbers. From number 1 - 12 are all OAs with biogenic influences. Number 13 - 17 are some

- 1184 pure chemical standards (acids) as discussed above. For detailed information on the meaning of
- 1185 the numbered symbols see supporting information Table S2.



**Figure 6**. Time series of ambient  $f_{C_5H_6O}^{OA}$ , gas-phase IEPOX, monoterpenes and isoprene in DC3

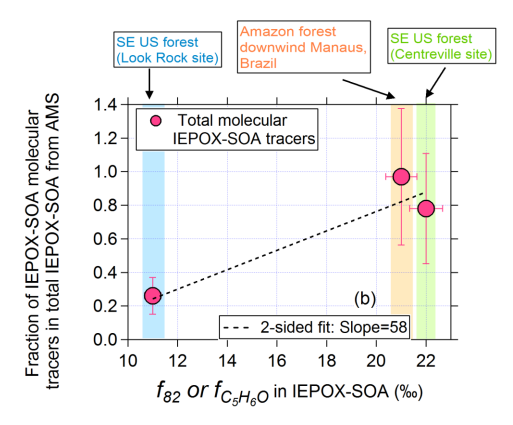
1190 aircraft measurement. Average  $f_{C_5H_6O}$  from regions strongly impacted by urban and biomass-

burning emissions and MT emissions are also shown for reference. Two areas with grey

1192 background indicate the periods when  $f_{C_5H_6O}^{OA}$  increases when monoterpene concentrations

1193 increase.

1194



1195

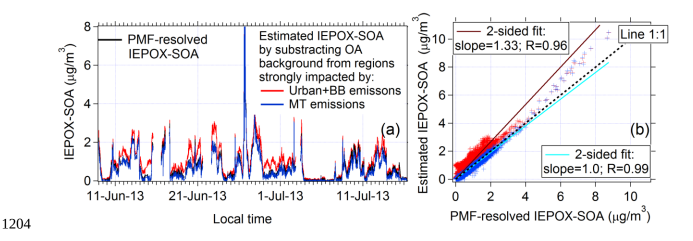
1196Figure 7. Scatter plot between total IEPOX-SOA molecular tracers (=Methyltetrol + C5-alkene1197triols +IEPOX-derived organosulfates and dimers) in IEPOX-SOA<sub>PMF</sub> and  $f_{82}^{IEPOX-SOA}$ . Besides1198SOAS, the other two datasets in the graph are from Budisulistiorini et al. (2015) and de Sá et

al.(2015). The relative uncertainty value estimated for the SOAS study is applied to the other two

1200 datasets.

1201





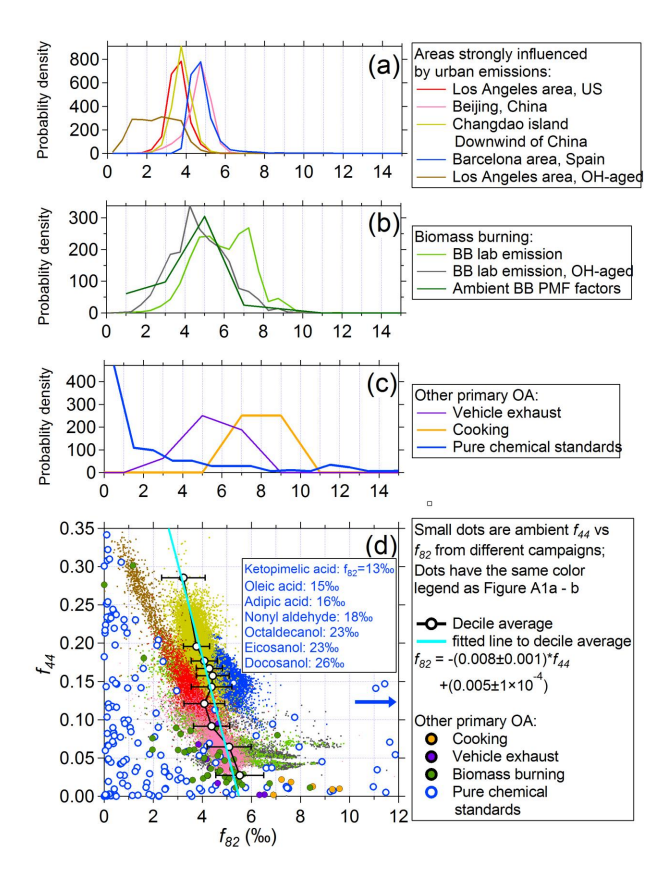
1205 **Figure 8**. (a) Time series of IEPOX-SOA<sub>PMF</sub> and estimated IEPOX-SOA based on  $C_5H_6O^+$  for

1206 the SOAS data in SE US. Two different estimates of background  $C_5H_6O^+$  are shown, using

values from regions strongly impacted by urban and biomass-burning emissions vs. regions with
 strong monoterpene emissions. (b) Scatter plot of estimated IEPOX-SOA vs. IEPOX-SOA<sub>PMF</sub>.

Note that the largest IEPOX-SOA plume on 26-Jun-13 had a slightly higher  $f_{C_5H_6O}^{OA}$  of 24‰,

1210 resulting in a slight overestimation of IEPOX-SOA for those data points.



- 1212 **Figure A1**. Probability density distributions of  $f_{82}$  in studies (a) strongly influenced by urban
- 1213 emissions; (b) biomass-burning emissions; (c) other anthropogenic primary OA sources and pure
- 1214 chemical standards. Several pure chemical species showing higher  $f_{82}$  between 15 30% are
- 1215 labeled with arrow. (d) Scatter plot of  $f_{44}$  ( $f_{44}=m/z$  44/OA) vs.  $f_{82}$  for all studies shown in panels
- 1216 (a) (c), using the same color scheme. Quantile averages of  $f_{82}$  across all studies sorted by  $f_{44}$
- 1217 are also shown, as is a linear regression line to the quantile points.
- 1218

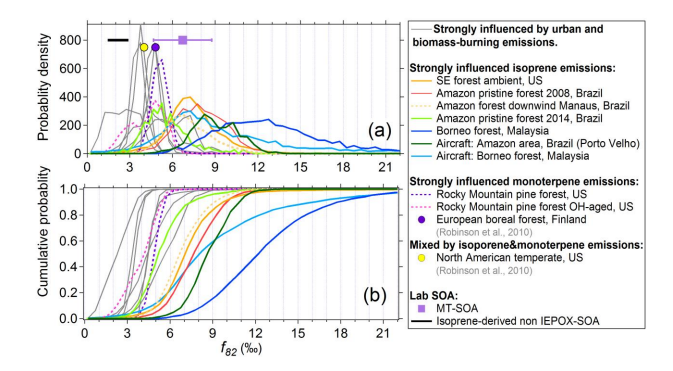


Figure A2. (a) Probability density and (b) cumulative probability distributions of  $f_{82}$  in studies strongly influenced by isoprene and/or monoterpene emissions. The ranges of  $f_{82}$  from other non IEPOX-derived isoprene-SOA and MT-SOA are also shown. The background grey lines are from studies strongly influenced by urban and biomass-burning emissions and are the same data from Fig. A1a – b.

1227

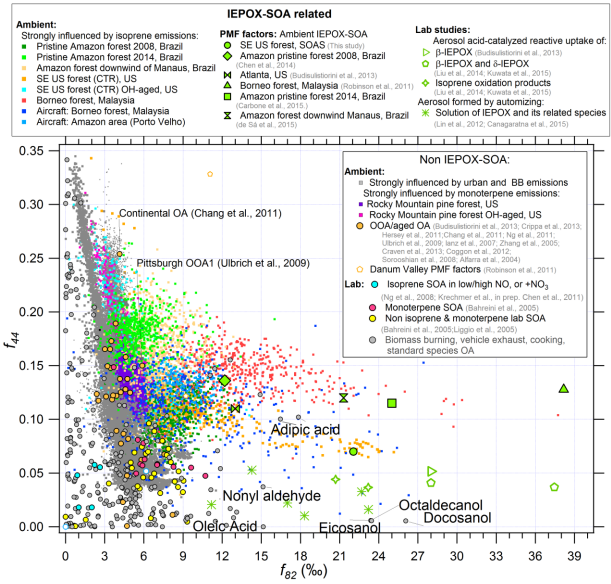
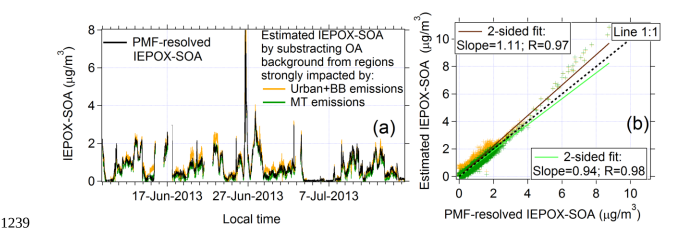




Figure A3. Scatter plot of  $f_{44}$  and  $f_{82}$  in studies strongly by isoprene and monoterpene emissions, as well as other OA sources. The grey dots represent background levels from studies strongly influenced by urban and biomass-burning emissions in Fig. A1d.  $f_{44}$  and  $f_{82}$  values from multiple sources of OA (Jimenez-Group, 2015) are also shown, together with IEPOX-SOA from different ambient PMF factors and chamber studies. 







1241 the SOAS-CTR data in SE US forest. Two different estimates of background m/z 82 are shown,

1242 using values from regions strongly impacted by urban and biomass-burning emissions vs.

regions with strong monoterpene emissions. (b) Scatter plot of estimated IEPOX-SOA vs.

1244 IEPOX-SOA<sub>PMF</sub>. Note that the largest IEPOX-SOA plume (> 4  $\mu$ g m<sup>-3</sup>) on 26-Jun-13 had a

1245 slightly higher  $f_{82}^{OA}$  of 24‰, resulting in a slight overestimation of IEPOX-SOA for those data

- 1246 points.
- 1247
- 1248