

# Secondary Organic Aerosol (SOA) yields from NO<sub>3</sub> radical + isoprene based on nighttime aircraft power plant plume transects

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## Abstract

Nighttime reaction of nitrate radicals (NO<sub>3</sub>) with biogenic volatile organic compounds (BVOC) has been proposed as a potentially important but also highly uncertain source of secondary organic aerosol (SOA). The southeast United States has both high BVOC and nitrogen oxide (NO<sub>x</sub>) emissions, resulting in a large model-predicted NO<sub>3</sub>-BVOC source of SOA. Coal-fired power plants in this region constitute substantial NO<sub>x</sub> emissions point sources into a nighttime atmosphere characterized by high regionally widespread concentrations of isoprene. In this paper, we exploit nighttime aircraft observations of these power plant plumes, in which NO<sub>3</sub> radicals rapidly remove isoprene, to obtain field-based estimates of the secondary organic aerosol yield from NO<sub>3</sub> + isoprene. Observed in-plume increases in nitrate aerosol are consistent with organic nitrate aerosol production from NO<sub>3</sub> + isoprene, and these are used to determine molar SOA yields, for which the average over 9 plumes is 9% (+/- 5%). Corresponding mass yields depend on the assumed molecular formula for isoprene-NO<sub>3</sub>-SOA, but the average over 9 plumes is 27% (+/- 14%), on average larger than those previously measured in chamber studies (12 – 14% mass yield as ΔOA/ΔVOC after oxidation of both double bonds). Yields are larger for longer plume ages. This suggests that ambient aging processes lead more effectively to condensable material than typical chamber conditions allow. We discuss potential mechanistic explanations for this difference, including longer ambient peroxy radical lifetimes and heterogeneous reactions of NO<sub>3</sub>-isoprene gas phase products.

43 More in-depth studies are needed to better understand the aerosol yield and oxidation  
44 mechanism of NO<sub>3</sub> radical + isoprene, a coupled anthropogenic – biogenic source of SOA that  
45 may be regionally significant.

## 46 **1 Introduction**

47 Organic aerosol (OA) is increasingly recognized as a globally important component of the fine  
48 particulate matter that exerts a large but uncertain negative radiative forcing on Earth's climate  
49 (Myhre et al., 2013) and adversely affects human health around the world (Lelieveld et al.,  
50 2015). This global importance is complicated by large regional differences in OA concentrations  
51 relative to other sources of aerosol such as black carbon, sulfate, nitrate and sea salt. OA  
52 comprises 20 – 50% of total fine aerosol mass at continental mid-latitudes, but more in urban  
53 environments and biomass burning plumes, and up to 90% over tropical forests (Kanakidou et  
54 al., 2005; Zhang et al., 2007). Outside of urban centers and fresh biomass burning plumes, the  
55 majority of this OA is secondary organic aerosol (SOA) (Jimenez et al., 2009), produced by  
56 oxidation of directly emitted volatile organic compounds followed by partitioning into the aerosol  
57 phase. Forests are strong biogenic VOC emitters, in the form of isoprene (C<sub>5</sub>H<sub>8</sub>), monoterpenes  
58 (C<sub>10</sub>H<sub>16</sub>), and sesquiterpenes (C<sub>15</sub>H<sub>24</sub>), all of which are readily oxidized by the three major  
59 atmospheric oxidants, OH, NO<sub>3</sub>, and O<sub>3</sub>. The total global source of biogenic SOA from such  
60 reactions remains highly uncertain, with a review estimating it at 90 +/- 90 Tg C yr<sup>-1</sup> (Hallquist et  
61 al., 2009), a large fraction of which may be anthropogenically controlled (Spracklen et al.,  
62 2011; Goldstein et al., 2009; Carlton et al., 2010; Hoyle et al., 2011). As most NO<sub>3</sub> arises from  
63 anthropogenic emissions, OA production from NO<sub>3</sub> + isoprene is one mechanism that could  
64 allow for the anthropogenic control of biogenic SOA mass loading.

65  
66 Isoprene constitutes nearly half of all global VOC emissions to the atmosphere, with a flux of  
67 ~600 Tg yr<sup>-1</sup> (Guenther et al., 2006). As a result, accurate global biogenic SOA budgets depend  
68 strongly on yields from isoprene oxidation. Recent global modeling efforts find that isoprene  
69 SOA is produced at rates from 14 (Henze and Seinfeld, 2006; Hoyle et al., 2007) to 19 TgC yr<sup>-1</sup>  
70 (Heald et al., 2008), which implies that it could constitute 27% (Hoyle et al., 2007) to 48%  
71 (Henze and Seinfeld, 2006) to 78% (Heald et al., 2008) of total SOA (based also on varying  
72 estimates of total SOA burden in each study). More recent observational constraints on SOA  
73 yield from isoprene find complex temperature-dependent mechanisms that could affect vertical  
74 distributions (Worton et al., 2013) and suggest that isoprene SOA constitutes from 17% (Hu et  
75 al., 2015) to 40% (Kim et al., 2015) up to 48% (Marais et al., 2016) of total OA in the  
76 southeastern United States. This large significance comes despite isoprene's low SOA mass  
77 yields – two recent observational studies estimated the total isoprene SOA mass yield to be  
78 ~3% (Marais et al., 2016; Kim et al., 2015), and modeling studies typically estimate isoprene  
79 SOA yields to be 4 to 10%, depending on the oxidant, in contrast to monoterpenes' yields of 10  
80 to 20% and sesquiterpenes' yields of >40% (Pye et al., 2010). Furthermore, laboratory studies  
81 of SOA mass yields may have a tendency to underestimate these yields, if they cannot access  
82 the longer timescales of later-generation chemistry, or are otherwise run under conditions that  
83 limit oxidative aging of first-generation products (Carlton et al., 2009).

84

85 Laboratory chamber studies of SOA mass yield at OA loadings of  $\sim 10 \mu\text{g m}^{-3}$  from isoprene  
86 have typically found low yields from  $\text{O}_3$  (1% (Kleindienst et al., 2007)) and OH (2% at low  $\text{NO}_x$  to  
87 5% at high  $\text{NO}_x$  (Kroll et al., 2006; Dommen et al., 2009); 1.3% at low  $\text{NO}_x$  and neutral seed  
88 aerosol pH but rising to 29% in the presence of acidic sulfate seed aerosol due to reactive  
89 uptake of epoxydiols of isoprene (IEPOX) (Surratt et al., 2010)). One recent chamber study on  
90 OH-initiated isoprene SOA formation focused on the fate of second-generation  $\text{RO}_2$  radical  
91 found significantly higher yields, up to 15% at low  $\text{NO}_x$  (Liu et al., 2016), suggesting that omitting  
92 later-generation oxidation chemistry could be an important limitation of early chamber  
93 determinations of isoprene SOA yields. Another found an increase in SOA formed with  
94 increasing  $\text{HO}_2$  to  $\text{RO}_2$  ratios, suggesting that  $\text{RO}_2$  fate could also play a role in the variability of  
95 previously reported SOA yields (D'Ambro et al., 2017).

96  
97 For  $\text{NO}_3$  oxidation of isoprene, early chamber experiments already pointed to higher yields (e.g.,  
98 12% (Ng et al., 2008)) than for OH oxidation. Ng et al. (Ng et al., 2008) also observed chemical  
99 regime differences: SOA yields were approximately two times larger when chamber conditions  
100 were tuned such that first-generation peroxy radical fate was  $\text{RO}_2+\text{RO}_2$  dominated than when it  
101 was  $\text{RO}_2+\text{NO}_3$  dominated. In addition, Rollins et al. (Rollins et al., 2009) observed a significantly  
102 higher SOA yield (14%) from second-generation  $\text{NO}_3$  oxidation than that when only one double  
103 bond was oxidized (0.7%). This points to the possibility that later-generation,  $\text{RO}_2+\text{RO}_2$   
104 dominated isoprene +  $\text{NO}_3$  chemistry may be an even more substantial source of SOA than  
105 what current chamber studies have captured. Schwantes et al. (Schwantes et al., 2015)  
106 investigated the gas-phase products of  $\text{NO}_3$  + isoprene in the  $\text{RO}_2+\text{HO}_2$  dominated regime and  
107 found the major product to be isoprene nitrooxy hydroperoxide (INP, 75-78% molar yield), which  
108 can photochemically convert to isoprene nitrooxy hydroxyepoxide (INHE), a molecule that might  
109 contribute to SOA formation via heterogeneous uptake similar to IEPOX. Here again, multiple  
110 generations of chemistry are required to produce products that may contribute to SOA.

111  
112 Because the SOA yield appears to be highest for  $\text{NO}_3$  radical oxidation, and isoprene is such an  
113 abundantly emitted BVOC, oxidation of isoprene by  $\text{NO}_3$  may be an important source of OA in  
114 areas with regional  $\text{NO}_x$  pollution. Since the SOA yield with neutral aerosol seed appears to be  
115 an order of magnitude larger than that from other oxidants, even if only 10% of isoprene is  
116 oxidized by  $\text{NO}_3$ , it will produce comparable SOA to daytime photo-oxidation. For example,  
117 Brown et al. (Brown et al., 2009) concluded that  $\text{NO}_3$  contributed more SOA from isoprene than  
118 OH over New England, where  $> 20\%$  of isoprene emitted during the previous day was available  
119 at sunset to undergo dark oxidation by either  $\text{NO}_3$  or  $\text{O}_3$ . The corresponding contribution to total  
120 SOA mass loading was 1 – 17% based on laboratory yields (Ng et al., 2017). Rollins et al.  
121 (Rollins et al., 2012) concluded that multi-generational  $\text{NO}_3$  oxidation of biogenic precursors was  
122 responsible for one-third of nighttime organic aerosol increases during the CalNex-2010  
123 experiment in Bakersfield, CA. In an aircraft study near Houston, TX, Brown et al. (Brown et al.,  
124 2013) observed elevated organic aerosol in the nighttime boundary layer, and correlated vertical  
125 profiles of organic and nitrate aerosol in regions with rapid surface level  $\text{NO}_3$  radical production  
126 and BVOC emissions. From these observations, the authors estimated an SOA source from  
127  $\text{NO}_3$  + BVOCs within the nocturnal boundary layer of  $0.05 - 1 \mu\text{g m}^{-3} \text{h}^{-1}$ . Carlton et al. (Carlton  
128 et al., 2009) note the large scatter in chamber-measured SOA yields from isoprene

129 photooxidation and point throughout their review of SOA formation from isoprene to the likely  
130 importance of poorly understood later generations of chemistry in explaining field observations.  
131 We suggest that similar differences in multi-generational chemistry could explain the variation  
132 among the (sparse) chamber and field observations of  $\text{NO}_3$  + isoprene yields described in the  
133 previous paragraph, and summarized in a recent review of  $\text{NO}_3$  + BVOC oxidation mechanisms  
134 and SOA formation (Ng et al., 2017).

135  
136 The initial products of  $\text{NO}_3$  + isoprene include organic nitrates, some of which will partially  
137 partition to the aerosol phase. Organic nitrates in the particle phase ( $\text{pRONO}_2$ ) are challenging  
138 to quantify with online methods, due to both interferences and their often overall low  
139 concentrations in ambient aerosol. Hence, field datasets to constrain modeled  $\text{pRONO}_2$  are  
140 sparse (Ng et al., 2017; Fisher et al., 2016). One of the most used methods in recent studies,  
141 used also here, is quantification with the Aerodyne Aerosol Mass Spectrometer (AMS). Organic  
142 nitrates thermally decompose in the AMS vaporizer and different approaches have been used to  
143 apportion the organic fraction contributing to the total nitrate signal. Allan et al. (Allan et al.,  
144 2004a) first proposed the use of nitrate peaks at  $m/z$  30 and 46 to distinguish various nitrate  
145 species with the AMS. Marcolli et al. (Marcolli et al., 2006), in the first reported tentative  
146 assignment of aerosol organic nitrate using AMS data, used cluster analysis to analyze data  
147 from the 2002 New England Air Quality Study. In that study, cluster analysis identified two  
148 categories with high  $m/z$  30 contributions. One of these peaked in the morning when  $\text{NO}_x$  was  
149 abundant and was more prevalent in plumes with lowest photochemical ages, potentially from  
150 isoprene oxidation products. The second was observed throughout the diurnal cycle in both  
151 fresh and aged plumes, and contained substantial  $m/z$  44 contribution (highly oxidized OA). A  
152 subsequent AMS laboratory and field study discussed and further developed methods for  
153 separate quantification of organic nitrate (in contrast to inorganic nitrate) (Farmer et al., 2010). A  
154 refined version of one of these separation methods, based on the differing  $\text{NO}_2^+/\text{NO}^+$   
155 fragmentation ratio for organic vs. inorganic nitrate, was later employed to quantify organic  
156 nitrate aerosol at two forested rural field sites where strong biogenic VOC emissions and  
157 relatively low  $\text{NO}_x$  combined to make substantial organic nitrate aerosol concentrations ((Fry et  
158 al., 2013; Ayres et al., 2015)). Most recently, Kiendler-Scharr et al. (Kiendler-Scharr et al., 2016)  
159 used a variant of this method to conclude that across Europe, organic nitrates comprise ~40%  
160 of submicron organic aerosol. Modeling analysis concluded that a substantial fraction of this  
161 organic nitrate aerosol is produced via  $\text{NO}_3$  radical initiated chemistry. Chamber studies have  
162 employed this fragmentation ratio method to quantify organic nitrates (Boyd et al., 2015; Bruns et  
163 al., 2010; Fry et al., 2009; Fry et al., 2011; Rollins et al., 2009), providing the beginnings of a  
164 database of typical organonitrate fragmentation ratios from various BVOC precursors.

165  
166 Measurements conducted at the SOAS ground site in Centreville, Alabama in 2013 found  
167 evidence of significant organonitrate contribution to SOA mass loading. Xu et al. (Xu et al.,  
168 2015) reported that organic nitrates constituted 5 to 12% of total organic aerosol mass from  
169 AMS data applying a variant of the  $\text{NO}_2^+/\text{NO}^+$  ratio method. They identify a nighttime-peaking  
170 “LO-OOA” AMS factor which they attribute to mostly  $\text{NO}_3$  oxidation of BVOC (in addition to  $\text{O}_3$  +  
171 BVOC). They estimated that the  $\text{NO}_3$  radical oxidizes 17% of isoprene, 20% of  $\alpha$ -pinene, and  
172 38% of  $\beta$ -pinene in the nocturnal boundary layer at this site. However, applying laboratory-

173 based SOA yields to model the predicted increase in OA, Xu et al. predict only  $0.7 \mu\text{g m}^{-3}$  of  
174 SOA would be produced, substantially lower than the measured nighttime LO-OOA production  
175 of  $1.7 \mu\text{g m}^{-3}$ . The more recent analysis of Zhang et al. (Zhang et al., 2018) found a strong  
176 correlation of monoterpene SOA with the fraction of monoterpene oxidation attributed to  $\text{NO}_3$ ,  
177 even for non-nitrate containing aerosol, suggesting an influence of  $\text{NO}_3$  even in pathways that  
178 ultimately eliminate the nitrate functionality from the SOA, such as hydrolysis or  $\text{NO}_2$   
179 regeneration. Ayres et al. (Ayres et al., 2015) used a correlation of overnight organonitrate  
180 aerosol buildup with calculated net  $\text{NO}_3$  + monoterpene and isoprene reactions to estimate an  
181 overall  $\text{NO}_3$  + monoterpene SOA mass yield of 40 – 80%. The factor of two range in this  
182 analysis was based on two different measurements of aerosol-phase organic nitrates. These  
183 authors used similar correlations to identify specific CIMS-derived molecular formulae that are  
184 likely to be  $\text{NO}_3$  radical chemistry products of isoprene and monoterpenes, and found minimal  
185 contribution of identified first-generation  $\text{NO}_3$  + isoprene products to the aerosol phase (as  
186 expected based on their volatility). Lee et al. (Lee et al., 2016) detected abundant highly  
187 functionalized particle-phase organic nitrates at the same site, with apparent origin both from  
188 isoprene and monoterpenes, and both daytime and nighttime oxidation, and estimated their  
189 average contribution to submicron organic aerosol mass to be between 3 – 8 %. For the same  
190 ground campaign, Romer et al. (Romer et al., 2016) found evidence of rapid conversion from  
191 alkyl nitrates to  $\text{HNO}_3$ , with total alkyl nitrates having an average daytime lifetime of 1.7 hours.

192  
193 Xie et al. (Xie et al., 2013) used a model constrained by observed alkyl nitrate correlations with  
194  $\text{O}_3$  from the INTEX-NA/ICARTT 2004 field campaign to determine a range of isoprene nitrate  
195 lifetimes between 4 and 6 hours, with 40-50% of isoprene nitrates formed by  $\text{NO}_3$  + isoprene  
196 reactions. Laboratory studies show that not all organic nitrates hydrolyze to  $\text{HNO}_3$  equally  
197 rapidly: primary and secondary organic nitrates were found to be less prone to aqueous  
198 hydrolysis than tertiary organic nitrates (Darer et al., 2011;Hu et al., 2011;Fisher et al.,  
199 2016;Boyd et al., 2015). This suggests that field-based estimates of the contribution of organic  
200 nitrates to SOA formation could be a lower limit, if they are based on measurement of those  
201 aerosol-phase nitrates. This is because if hydrolysis is rapid, releasing  $\text{HNO}_3$  but leaving behind  
202 the organic fraction in the aerosol phase, then that organic mass would not be accurately  
203 accounted for as arising from nitrate chemistry. This was addressed in a recent modeling study  
204 of SOAS (Pye et al., 2015) in which modeled hydrolysis products of particulate organic nitrates  
205 of up to  $0.8 \mu\text{g m}^{-3}$  additional aerosol mass loading in the southeast U.S. were included in the  
206 estimate of change in OA due to changes in  $\text{NO}_x$ . Another recent GEOS-Chem modeling study  
207 using of gas- and particle-phase organic nitrates observed during the SEAC<sup>4</sup>RS and SOAS  
208 campaigns similarly finds  $\text{RONO}_2$  to be a major sink of  $\text{NO}_x$  across the SEUS region (Fisher et  
209 al., 2016;Lee et al., 2016).

210  
211 Complementing these SOAS ground site measurements, the NOAA-led SENEX (Southeast  
212 Nexus) aircraft campaign conducted 18 research flights focused in part on studying the  
213 interactions between biogenic and anthropogenic emissions that form secondary pollutants  
214 between 3 June and 10 July 2013 (Warneke et al., 2016). Flight instrumentation focused on  
215 measurement of aerosol precursors and composition enable the present investigation of SOA  
216 yields using this aircraft data set. Edwards et al. (Edwards et al., 2017) used data from the

217 SENEX night flights to evaluate the nighttime oxidation of BVOC, observing high nighttime  
218 isoprene mixing ratios in the residual layer that can undergo rapid NO<sub>3</sub> oxidation when sufficient  
219 NO<sub>x</sub> is present. These authors suggest that past NO<sub>x</sub> reductions may have been uncoupled  
220 from OA trends due to NO<sub>x</sub> not having been the limiting chemical species for OA production, but  
221 that future reductions in NO<sub>x</sub> may decrease OA if NO<sub>3</sub> oxidation of BVOC is a substantial  
222 regional SOA source. Because isoprene is ubiquitous in the nighttime residual layer over the  
223 southeastern United States and the NO<sub>3</sub> + isoprene reaction is rapid, NO<sub>3</sub> reaction will be  
224 dominant relative to O<sub>3</sub> in places with anthropogenic inputs of NO<sub>x</sub> (Edwards et al. (Edwards et  
225 al., 2017) concludes that when NO<sub>2</sub>/BVOC > 0.5, NO<sub>3</sub> oxidation will be dominant). Hence, a  
226 modest NO<sub>3</sub> + isoprene SOA yield may constitute a regionally important OA source.

227  
228 Several modeling studies have investigated the effects of changing NO<sub>x</sub> on global and SEUS  
229 SOA. Hoyle et al. (Hoyle et al., 2007) found an increase in global SOA production from 35 Tg yr<sup>-1</sup>  
230 to 53 Tg yr<sup>-1</sup> since preindustrial times, resulting in an increase in global annual mean SOA  
231 mass loading of 51%, attributable in part to changing NO<sub>x</sub> emissions. Zheng et al. (Zheng et al.,  
232 2015) found only moderate SOA reductions from a 50% reduction in NO emissions: 0.9 – 5.6 %  
233 for global NO<sub>x</sub> or 6.4 – 12.0% for southeast US NO<sub>x</sub>, which they attributed to buffering by  
234 alternate chemical pathways and offsetting tendencies in the biogenic vs. anthropogenic SOA  
235 components. In contrast, Pye et al. (Pye et al., 2015) find a 9% reduction in total organic aerosol  
236 in Centreville, AL for only 25% reduction in NO<sub>x</sub> emissions. A simple limiting-reagent analysis of  
237 NO<sub>3</sub> + monoterpene SOA from power plant plumes across the United States found that between  
238 2008 and 2011, based on EPA-reported NO<sub>x</sub> emissions inventories, some American power  
239 plants shifted to the NO<sub>x</sub>-limited regime (from 3.5% to 11% of the power plants), and showed  
240 that these newly NO<sub>x</sub>-limited power plants were primarily in the southeastern United States (Fry  
241 et al., 2015). The effect of changing NO<sub>x</sub> on SOA burden is clearly still in need of further study.

242  
243 Here, we present aircraft transects of spatially discrete NO<sub>x</sub> plumes from electric generating  
244 units (EGU), or power plants (PP), as a method to specifically isolate the influence of NO<sub>3</sub>  
245 oxidation. These plumes are concentrated and highly enriched in NO<sub>x</sub> over a scale of only a  
246 few km (Brown et al., 2012), and have nitrate radical production rates ( $P(\text{NO}_3)$ ) 10 – 100 times  
247 greater than those of background air. The rapid shift in  $P(\text{NO}_3)$  allows direct comparison of air  
248 masses with slow and rapid oxidation rates attributable to the nitrate radical, effectively isolating  
249 the influence of this single chemical pathway in producing SOA and other oxidation products.  
250 Changes in organic nitrate aerosol (pRONO<sub>2</sub>) concentration and accompanying isoprene  
251 titration enable a direct field determination of the SOA yield from NO<sub>3</sub> + isoprene.

## 252 **2 Field campaign and experimental and modeling methods**

253 The Southeast Nexus (SENEX: <http://esrl.noaa.gov/csd/projects/senex/>) campaign took place 3  
254 June through 10 July 2013 as the NOAA WP-3D aircraft contribution to the larger Southeast  
255 Atmospheric Study (SAS: [http://www.eol.ucar.edu/field\\_projects/sas/](http://www.eol.ucar.edu/field_projects/sas/)), a large, coordinated  
256 research effort focused on understanding natural and anthropogenic emissions, oxidation  
257 chemistry and production of aerosol in the summertime atmosphere in the southeastern United  
258 States. The NOAA WP-3D aircraft operated 18 research flights out of Smyrna, Tennessee,

259 carrying an instrument payload oriented towards elucidating emissions inventories and reactions  
260 of atmospheric trace gases, and aerosol composition and optical properties (Warneke et al.,  
261 2016). One of the major goals of the larger SAS study is to quantify the fraction of organic  
262 aerosol that is anthropogenically controlled, with a particular focus on understanding how OA  
263 may change in the future in response to changing anthropogenic emissions.

264

265 The subset of aircraft instrumentation employed for the present analysis of nighttime  $\text{NO}_3$  +  
266 isoprene initiated SOA production includes measurements used to determine  $\text{NO}_3$  radical  
267 production rate ( $P(\text{NO}_3) = k_{\text{NO}_2+\text{O}_3}(\text{T}) [\text{NO}_2] [\text{O}_3]$ ), isoprene and monoterpene concentrations,  
268 other trace gases for plume screening and identification, aerosol size distributions, and aerosol  
269 composition. The details on the individual measurements and the overall aircraft deployment  
270 goals and strategy are described in Warneke et al. (Warneke et al., 2016). Briefly,  $\text{NO}_2$  was  
271 measured by UV photolysis and gas-phase chemiluminescence (P-CL) and by cavity ringdown  
272 spectroscopy, (CRDS), which agreed within 6%.  $\text{O}_3$  was also measured by both gas-phase  
273 chemiluminescence and CRDS and agreed within 8%, within the combined measurement  
274 uncertainties of the instruments. Various volatile organic compounds were measured with  
275 several techniques, including for the isoprene and monoterpenes of interest here, proton  
276 reaction transfer mass spectrometry (PTR-MS) and canister whole air samples and post-flight  
277 GC-MS analysis (iWAS/GCMS). A comparison of PTR-MS and iWAS/GCMS measurements of  
278 isoprene during SENEX has high scatter due to imperfect time alignment and isoprene's high  
279 variability in the boundary layer, but the slope of the intercomparison is 1.04 ((Warneke et al.,  
280 2016); for more details on the VOC intercomparisons, see also Lerner et al., (Lerner et al.,  
281 2017)). Acetonitrile from the PTRMS was used to screen for the influence of biomass burning.  
282 Sulfur dioxide ( $\text{SO}_2$ ) was used to identify emissions from coal-fired power plants. All gas-phase  
283 instruments used dedicated inlets, described in detail in the supplemental information for  
284 Warneke et al. (Warneke et al., 2016).

285

286 Aerosol particles were sampled downstream of a low turbulence inlet (Wilson et al., 2004), after  
287 which they were dried by ram heating, size-selected by an impactor with 1  $\mu\text{m}$  aerodynamic  
288 diameter size cut-off, and measured by various aerosol instruments (Warneke et al., 2016). An  
289 ultra-high-sensitivity aerosol sizing spectrometer (UHSAS, Particle Metrics, Inc., Boulder, CO  
290 (Brock et al., 2011; Cai et al., 2008)) was used to measure the dry submicron aerosol size  
291 distribution down to about 70 nm. Data for the UHSAS are reported at 1 Hz whereas AMS data  
292 were recorded roughly every 10 seconds. The ambient (wet) surface areas were calculated  
293 according to the procedures described in Brock et al., 2016 (Brock et al., 2016). A pressure-  
294 controlled inlet (Bahreini et al., 2008) was employed to ensure that a constant mass flow rate  
295 was sampled by a compact time-of-flight aerosol mass spectrometer (C-ToF-AMS) which  
296 measured the non-refractory aerosol composition (Drewnick et al., 2005). The aerosol volume  
297 transmitted into the AMS was calculated by applying the measured AMS lens transmission  
298 curve (Bahreini et al., 2008) to the measured particle volume distributions from the UHSAS. For  
299 the entire SENEX study, the mean, calculated fraction of aerosol volume behind the 1 micron  
300 impactor that was transmitted through the lens into the AMS instrument was 97% (with  $\pm 4\%$   
301 standard deviation), indicating that most of the submicron aerosol volume measured by the  
302 sizing instruments was sampled by the AMS.

303

304 After applying calibrations and the composition-dependent collection efficiency following  
305 Middlebrook et al. (Middlebrook et al., 2012), the limits of detection for the flight analyzed here  
306 were  $0.05 \mu\text{g m}^{-3}$  for nitrate,  $0.26 \mu\text{g m}^{-3}$  for organic mass,  $0.21 \mu\text{g m}^{-3}$  for ammonium, and  $0.05$   
307  $\mu\text{g m}^{-3}$  for sulfate, determined as three times the standard deviation of 10-second filtered air  
308 measurements obtained for 10 minutes during preflight and 10 minutes during postflight (110  
309 datapoints). Note that the relative ionization efficiency for ammonium was 3.91 and 3.87 for the  
310 two bracketing calibrations and an average value of 3.9 was used for the flight analyzed here.  
311 An orthogonal distance regression (ODR-2) of the volume from composition data (AMS mass  
312 plus refractory black carbon) using a mass weighted density as described by Bahreini et al.  
313 (Bahreini et al., 2009) versus the volume based on the sizing instruments (after correcting for  
314 AMS lens transmission as above) had a slope of 1.06 for the entire SENEX study and 72% of  
315 the data points were within the measurements' combined uncertainties of  $\pm 45\%$  (Bahreini et al.,  
316 2008). For the flight analyzed here, however, the same regression slope was 1.58, which is  
317 slightly higher than the combined uncertainties. It is unclear why the two types of volume  
318 measurements disagree more for this flight. This does not change the conclusions of this work  
319 because this has been incorporated into the error in aerosol organic nitrate, which still show  
320 positive enhancements in pRONO<sub>2</sub> for these plumes (see Figure 4 below). These complete  
321 error estimates are also used in Figure 5 to clearly show the uncertainties in the yields. The  
322 volume comparison is discussed further in the Supplemental Information and shown for the  
323 plumes of interest in Fig. S1.

324

325 The C-ToF-AMS is a unit mass resolution (UMR) instrument and the mass spectral signals that  
326 are characteristic of aerosol nitrate at  $m/z$  30 and 46 ( $\text{NO}^+$  and  $\text{NO}_2^+$ ) often contain interferences  
327 from organic species such as  $\text{CH}_2\text{O}^+$  and  $\text{CH}_2\text{O}_2^+$ , respectively. Here, the  $m/z$  30 and 46 signals  
328 have been corrected for these interferences by using correlated organic signals at  $m/z$  29, 42,  
329 43, and 45 that were derived from high-resolution AMS measurements during the NASA  
330 SEAC<sup>4</sup>RS campaign that took place in the same regions of the SE US shortly after SENEX (see  
331 Supplemental Information and Fig. S2). The corrections were applied to the individual flight  
332 analyzed here from July 2. All of the corrections were well correlated with each other for the  
333 SEAC<sup>4</sup>RS dataset and we used the organic peak at  $m/z$  29 (from  $\text{CHO}^+$ ) and the peak at  $m/z$  45  
334 (from  $\text{CHO}_2^+$ ), respectively, since those corrections were from peaks closest (in  $m/z$ ) to those  
335 being corrected. Once corrected, the nitrate mass concentrations in the final data archive for  
336 this flight were reduced by  $0\text{--}0.24 \mu\text{g sm}^{-3}$ , an average reduction of  $0.11 \mu\text{g sm}^{-3}$  or 32% from  
337 the initial nitrate mass concentrations. The organic interferences removed from the  $m/z$  30 and  
338  $m/z$  46 signals are linearly correlated with the total organic mass concentrations, corresponding  
339 to an average 1.3% increase in the total organic mass.

340

341 The ratio of the corrected  $\text{NO}_2^+/\text{NO}^+$  signals was then used to calculate the fraction of aerosol  
342 nitrate that was organic (pRONO<sub>2</sub>) or inorganic (ammonium nitrate) based on the method  
343 described first in (Fry et al., 2013). Here we used an organic  $\text{NO}_2^+/\text{NO}^+$  ratio that was equal to  
344 the ammonium nitrate  $\text{NO}_2^+/\text{NO}^+$  ratio from our calibrations divided by 2.8. This factor was  
345 determined from multiple datasets (see discussion in Supplemental Information). The  
346 ammonium nitrate  $\text{NO}_2^+/\text{NO}^+$  ratio was obtained from the two calibrations on 30 June and 7 July



347 that bracketed the flight on 2 July, which is analyzed here. It was 0.514 and 0.488, respectively,  
348 and for all of the data from both calibrations it averaged 0.490. Hence, the organic nitrate  
349  $\text{NO}_2^+/\text{NO}^+$  ratio was estimated to be 0.175. This is the first time, to our knowledge, that UMR  
350 measurements of aerosol nitrate have been corrected with HR correlations and used to  
351 apportion the corrected nitrate into inorganic or organic nitrate species.

352

353 The time since emission of intercepted power plant plumes was estimated from the slope of a  
354 plot of  $\text{O}_3$  against  $\text{NO}_2$ . For nighttime emitted  $\text{NO}_x$  plumes that consist primarily of NO (Peischl  
355 et al., 2010),  $\text{O}_3$  is negatively correlated with  $\text{NO}_2$  due to the rapid reaction of NO with  $\text{O}_3$  that  
356 produces  $\text{NO}_2$  in a 1:1 ratio:

357



359

360 Reaction R1 goes rapidly (NO pseudo first order loss rate coefficient of  $0.03 \text{ s}^{-1}$  at 60 ppb  $\text{O}_3$ ) to  
361 completion, so that all  $\text{NO}_x$  is present as  $\text{NO}_2$ , as long as the plume NO does not exceed  
362 background  $\text{O}_3$  after initial mixing of the plume into background air. Subsequent oxidation of  
363  $\text{NO}_2$  via reaction (R2) leads to an increasingly negative slope of  $\text{O}_3$  vs  $\text{NO}_2$ :

364



366

367 Equation (1) then gives plume age subsequent to the completion of (R1) in terms of the  
368 observed slope,  $m$ , of  $\text{O}_3$  vs  $\text{NO}_2$  (Brown et al., 2006).

369

$$370 t_{\text{plume}} = \frac{\ln[1-S(m+1)]}{Sk_1\bar{\text{O}}_3} \quad (1)$$

371

372 Here  $S$  is a stoichiometric factor that is chosen for this analysis to be 1 based on agreement of  
373 plume age with elapsed time in a box model run initialized with SENEX flight conditions (see  
374 below);  $k_1$  is the temperature dependent bimolecular rate constant for  $\text{NO}_2 + \text{O}_3$  (R2) and  $\bar{\text{O}}_3$  is  
375 the average  $\text{O}_3$  within the plume.

376

377 We calculate plume ages using both a stoichiometric factor of 1 (loss of  $\text{NO}_3$  and  $\text{N}_2\text{O}_5$   
378 dominated by  $\text{NO}_3$  reactions) and 2 (loss dominated by  $\text{N}_2\text{O}_5$  reactions), although we note that  
379 the chemical regime for  $\text{NO}_3+\text{N}_2\text{O}_5$  loss may change over the lifetime of the plume, progressing  
380 from 1 to 2 as the BVOC is consumed. We use  $S=1$  values in the analysis that follows. Because  
381 the more aged plumes are more likely to have  $S$  approach 2, this means that some of the older  
382 plumes may have overestimated ages. Fig. S3 in the Supplemental Information shows the  
383 plume age calculated by Eq. 1 using modeled  $\text{NO}_x$ ,  $\text{NO}_y$  and  $\text{O}_3$  concentrations for  $S=1$  and  
384  $S=2$ , from nighttime simulations of plume evolution using an observationally constrained box  
385 model. This confirms that for nighttime plumes,  $S=1$  plume ages match modeled elapsed time  
386 well. The model used for this calculation, and those used to assess peroxy radical lifetimes and  
387 fates in Section 4.3, was the Dynamically Simple Model of Atmospheric Chemical Complexity  
388 (DSMACC (Emmerson and Evans, 2009)) containing the Master Chemical Mechanism v3.3.1  
389 chemistry scheme (Jenkin et al., 2015). More details on the model approach are provided in the  
390 SI.

### 391 **3 Nighttime flight selection**

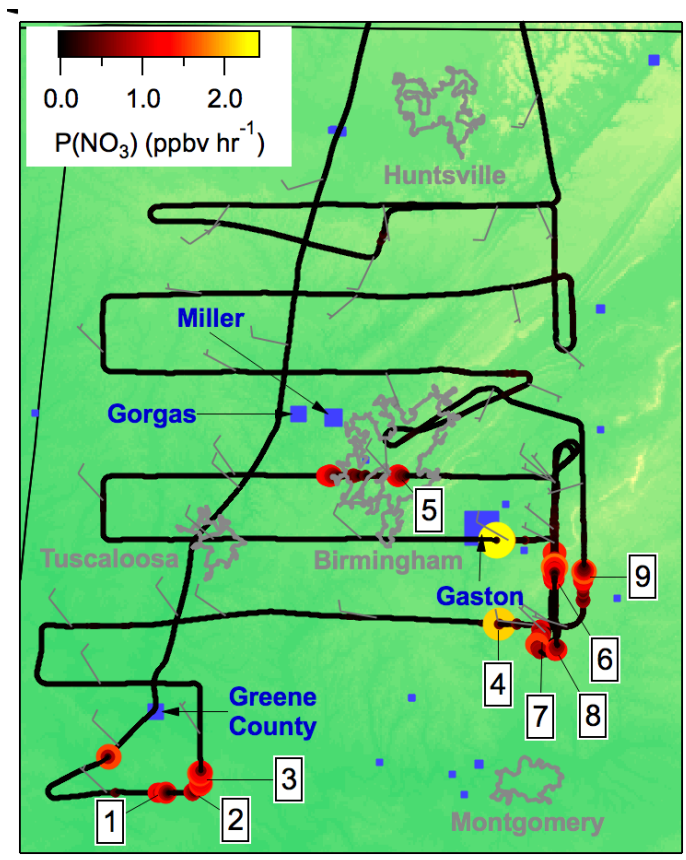
392 There were three nighttime flights (takeoffs on the evenings of 19 June, 2 July, and 3 July,  
393 2013, local time) conducted during SENEX, of which one (2 July) surveyed regions surrounding  
394 Birmingham, Alabama, including multiple urban and power plant plume transects. As described  
395 in the introduction, these plume transects are the focus of the current analysis since they  
396 correspond to injections of concentrated NO (and subsequently high  $P(\text{NO}_3)$ ) into the regionally  
397 widespread residual layer isoprene. The nighttime flight on 3 July, over Missouri, Tennessee  
398 and Arkansas sampled air more heavily influenced by biomass burning than biogenic emissions.  
399 The 19 June night flight sampled earlier in the evening, in the few hours immediately after  
400 sunset, and sampled more diffuse urban plume transects that had less contrast with background  
401 air. Therefore, this paper uses data exclusively from the 2 July flight, in which 9 transects of  
402 well-defined  $\text{NO}_x$  plumes from power plants emitted during darkness can be analyzed to obtain  
403 independent yields measurements.

404  
405 A map of the 2 July flight track is shown in Fig. 1a. After takeoff at 8:08 pm local Central  
406 Daylight Time on 2 July, 2013 (1:08 am UTC 3 July, 2016), the flight proceeded towards the  
407 southwest until due west of Montgomery, AL, after which it conducted a series of east-west  
408 running tracks while working successively north toward Birmingham, AL. Toward the east of  
409 Birmingham, the aircraft executed overlapping north-south tracks at six elevations to sample the  
410 E. C. Gaston power plant. During the course of the flight, concentrated  $\text{NO}_x$  plumes from the  
411 Gaston, Gorgas, Miller and Greene City power plants were sampled. Around 1:30 and 2:30 AM  
412 Central Daylight Time (5:30 and 6:30 am UTC), two transects of the Birmingham, AL urban  
413 plume were measured prior to returning to the Smyrna, TN airport base.

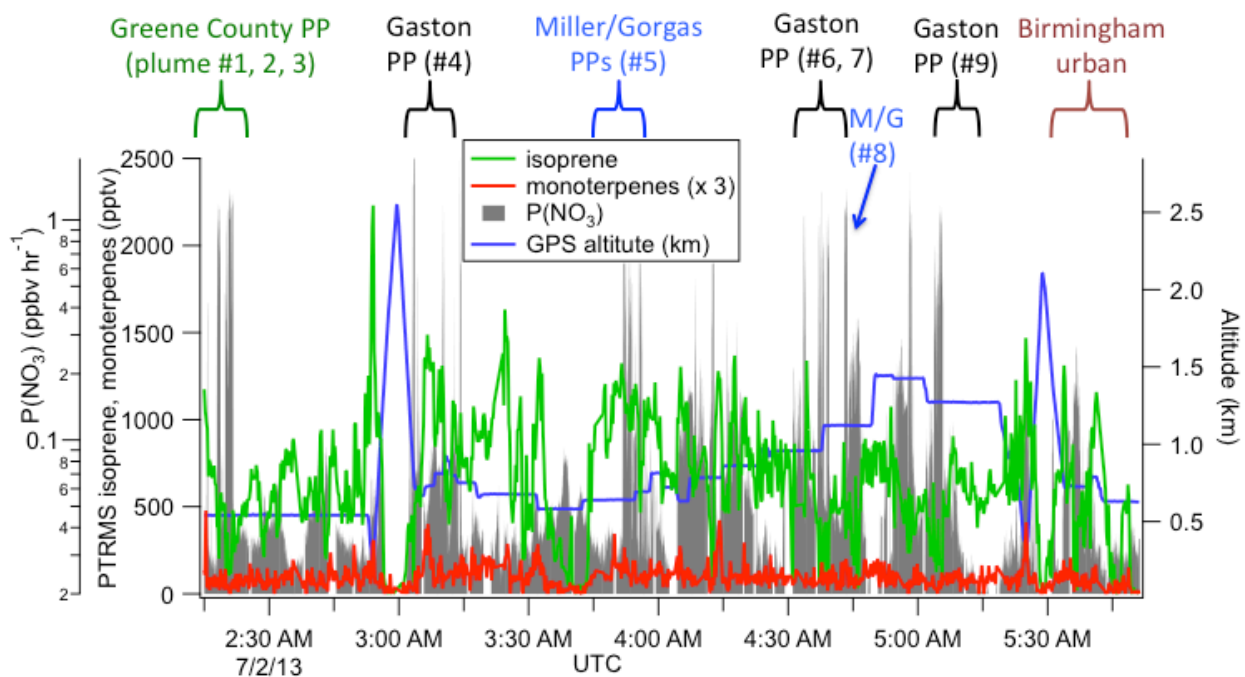
414  
415 The flight track is shown colored by the nitrate radical production rate,  $P(\text{NO}_3)$ , to show the  
416 points of urban and/or power plant plume influence:

$$417 \quad P(\text{NO}_3) = k_2(T) [\text{NO}_2][\text{O}_3] \quad (2)$$

418  
419 Here,  $k_2$  is again the temperature-dependent rate coefficient for reaction of  $\text{NO}_2 + \text{O}_3$  (Atkinson  
420 et al., 2004), and the square brackets indicate concentrations. Fig. 1b further illustrates the  
421 selection of power plants plumes: sharp peaks in  $P(\text{NO}_3)$  are indicative of power plant plume  
422 transects, during which isoprene mixing ratios also are observed to drop from the typical  
423 regional residual layer background values of  $\sim 1$  ppb, indicative of loss by  $\text{NO}_3$  oxidation (an  
424 individual transect is shown in more detail below in Fig. 2). Also shown in Fig. 1b are measured  
425 concentrations of isoprene and monoterpenes throughout the flight, showing substantial residual  
426 layer isoprene and supporting the assumption that effectively all  $\text{NO}_3$  reactivity is via isoprene  
427 (see calculation in next section). Residual layer concentrations of other VOCs that could  
428 produce SOA (e.g., aromatics) are always below 100 pptv, and their reaction rates with  $\text{NO}_3$  are  
429 slow. Edwards *et al.* (Edwards et al., 2017) have shown that  $\text{NO}_3$  and isoprene mixing ratios for  
430 this and other SENEX night flights exhibit a strong and characteristic anticorrelation that is  
431 consistent with nighttime residual layer oxidation chemistry.  
432  
433



434 1a  
435 1b



436  
437  
438  
439

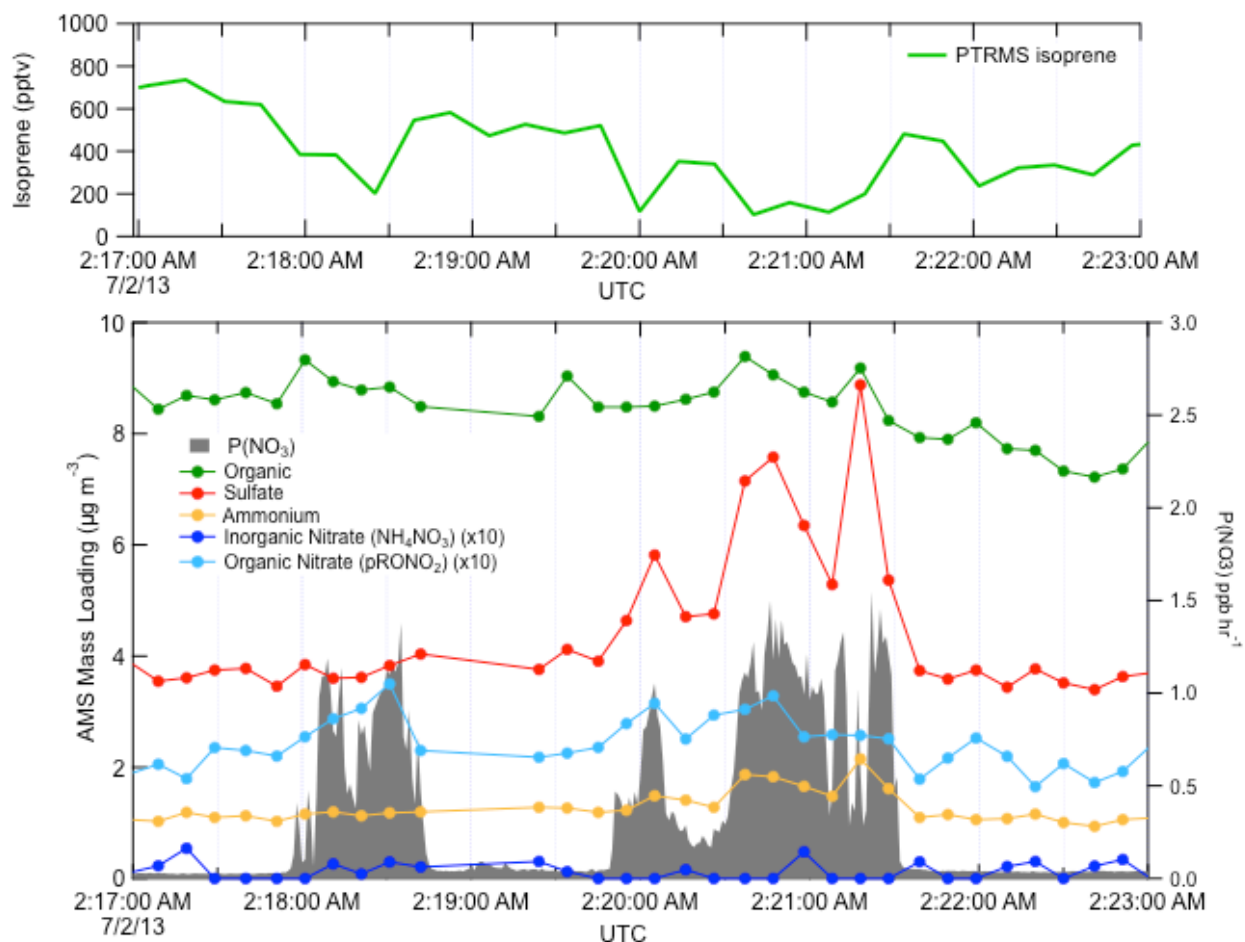
**Figure 1a.** Map of northern Alabama, showing the location of the flight track of the 2 July 2013 night flight used in the present analysis, with plume numbers labeled and wind direction shown. Although the wind direction changed throughout the night, these measurements enable us to

440 attribute each plume to a power plant source (see labels in Figure 1b and Table 2). Color scale  
441 shows  $P(\text{NO}_3)$  based on aircraft-measured  $[\text{NO}_2]$  and  $[\text{O}_3]$ , while power plants discussed in the  
442 text are indicated in blue squares with marker size scaled to annual  $\text{NO}_x$  emissions for 2013  
443 (scale not shown). Isoprene emissions are widespread in the region (Edwards et al., 2017).  
444 **Figure 1b** shows time series data from the same flight, with plume origins and numbers labeled,  
445 showing aircraft-measured isoprene and monoterpene concentrations, altitude, and  $P(\text{NO}_3)$   
446 determined according to Eq. 2 (log scale), showing that the isoprene was uniformly distributed  
447 (mixing ratios often in excess of 1 ppbv), while the more reactive monoterpenes were present at  
448 mixing ratios below 100 ppt except at the lowest few hundred meters above ground in the  
449 vertical profiles (not used in the present analysis). Figure 1b also shows that sharp peaks in  
450 nitrate radical production rate occur both at the lowest points of these vertical profiles, when the  
451 aircraft approached the surface, but also frequently during periods of level flight in the residual  
452 layer, which correspond to the power plant plume transects analyzed in this paper.

## 453 **4 Results**

### 454 **4.1 Selection of plumes**

455 Figure 2 shows a subset of the July 2 flight time series data, illustrating three  $\text{NO}_x$  plumes used  
456 for analysis. The large  $\text{NO}_3$  source and isoprene loss was accompanied by an increase in  
457 organic nitrate aerosol mass, which we attribute to the  $\text{NO}_3 + \text{isoprene}$  reaction based on prior  
458 arguments. We observed each plume as a rapid and brief perturbation to background  
459 conditions, of order 10 – 50 sec., or 1 – 5 km in spatial scale. Each plume's perturbed  
460 conditions can correspond to different plume ages, depending on how far downwind of the  
461 power plant the plume transect occurred.



463  
 464 **Figure 2.** Three representative plume transect observations from the 2 July 2013 flight (plumes  
 465 are identified by the peaks in  $P(\text{NO}_3)$ , listed in Table 1 at times 02:18, 02:20, and 02:21 UTC).  
 466 Note the difference in sulfate enhancement in the three plumes, which is largest in the third  
 467 plume, and is accompanied by increases in ammonium. In all three cases, the isoprene  
 468 concentration drops in the plumes, accompanied by a clear increase in organic nitrate, no  
 469 changes in the inorganic nitrate, and a modest changes in organic aerosol mass concentrations.

470  
 471 Candidate plumes were initially identified by scanning the time series flight data for any period  
 472 where the production rate of nitrate radical ( $P(\text{NO}_3)$ ) rose above  $0.5 \text{ ppbv hr}^{-1}$ . This threshold  
 473 was chosen to be above background noise and large enough to isolate only true plumes (see  
 474 Fig. 1a). The value is thus subjectively chosen, but was consistently applied across the dataset.  
 475 For each such period, a first screening removed any of these candidate plumes that occurred  
 476 during missed approaches or other periods where radar altitude above ground level (AGL) was  
 477 changing, because in the stratified nighttime boundary layer structure, variations in altitude may  
 478 result in sampling different air-masses, rendering the adjacent out of plume background not  
 479 necessarily comparable to in-plume conditions. A second criterion for rejection of a plume was  
 480 missing isoprene or AMS data during brief plume intercepts. No selected plumes on July 2  
 481 showed enhanced acetonitrile or refractory black carbon, indicating no significant biomass

482 burning influence. Finally, two plumes downwind of the Gaston power plant (at 03:10 and  
483 03:14) were removed from the present analysis, because (03:10) the background isoprene was  
484 changing rapidly, preventing a good baseline measurement, and (03:14) there was no observed  
485 decrease in isoprene concentration in-plume (as well as no increase in nitrate aerosol). The  
486 03:14 plume was apparently too recently emitted to have undergone significant nighttime  
487 reaction; its  $O_3/NO_2$  slope was unity to within the combined measurement error of  $O_3$  and  $NO_2$   
488 (Eq. 1). After this filtering, there are 9 individual plume observations for determination of  $NO_3$  +  
489 isoprene SOA yields (see Table 1). The rapid increases in  $P(NO_3)$  appeared simultaneously with  
490 significant decreases in isoprene and increases in aerosol nitrate. The aerosol and isoprene  
491 measurements (taken at data acquisition rates  $< 1$  Hz) were not exactly coincident in time which  
492 leads to some uncertainty in the yield analysis below.

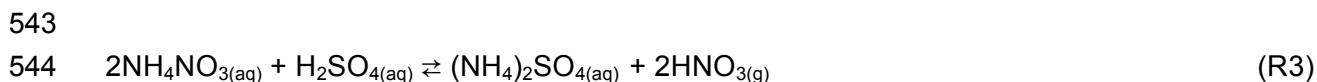
493  
494 Derivation of SOA yields from observed changes in isoprene and aerosol mass in plumes  
495 depends on two conditions, and has several caveats that will be discussed in the text that  
496 follows (see Table 3 below for a summary of these caveats). The two conditions are: (1) that the  
497 majority of VOC mass consumed by  $NO_3$  in plumes is isoprene (rather than monoterpenes or  
498 other VOC), and then either or both (2a) that the change in aerosol organic mass concentration  
499 during these plumes is due to  $NO_3$  + isoprene reactions, and/or (2b) that the change in aerosol  
500 nitrate mass concentration is due to  $NO_3$  + isoprene reactions. There are separate  
501 considerations for each of these conditions.

502  
503 For the first condition, we note that the isoprene to monoterpenes ratio just outside each plume  
504 transect was always high (a factor of 10 to 70, on average 26). With the 298 K  $NO_3$  rate  
505 constants of  $\sim 5 \times 10^{-12} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$  for monoterpenes and  $6.5 \times 10^{-13} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$  for  
506 isoprene (Calvert et al., 2000), isoprene ( $\sim 2$  ppb) will always react faster with nitrate than  
507 monoterpenes ( $\sim 0.04$  ppbv). At these relative concentrations, even if all of the monoterpene is  
508 oxidized, the production rate of oxidation products will be much larger for isoprene. Contribution  
509 to aerosol by  $N_2O_5$  uptake is also not important in these plumes. Edwards et al. (Edwards et al.,  
510 2017) calculated the sum of  $NO_3$  and  $N_2O_5$  loss throughout this flight and showed that it is  
511 consistently  $NO_3$ +BVOC dominated (Fig. S4 of that paper). As isoprene depletes,  $N_2O_5$  uptake  
512 will increasingly contribute to  $NO_3$  loss, but as shown below, we are able to rule out a  
513 substantial source of inorganic nitrate for most plumes. We also know that despite increased  
514 OH production in-plume, the isoprene loss is still overwhelming dominated by  $NO_3$  (Fig. S5 in  
515 Edwards, et al. (Edwards et al., 2017)) .

516  
517 The second condition requires that we can find an aerosol signal that is attributable exclusively  
518 to  $NO_3$  + isoprene reaction products, whether it be organic aerosol (OA) or organic nitrate  
519 aerosol ( $pRONO_2$ ) mass loading, or both. We note that the ratio of in-plume aerosol organic  
520 mass increase to  $pRONO_2$  mass increase is noisy (see discussion below at Fig. 6), but indicates  
521 an average in-plume  $\Delta OA$  to  $\Delta pRONO_2$  ratio of about 5. The large variability is primarily due to  
522 the fact that the variability in organic aerosol mass between successive 10-second data points  
523 for the entire flight is quite large (of order  $0.75 \mu\text{g m}^{-3}$ ) and comparable to many of the individual  
524 plume  $\Delta OA$  increases, far exceeding the expected organonitrate driven increases in OA, which  
525 are roughly twice the  $pRONO_2$  mass increases. It is also possible that in these plumes, where

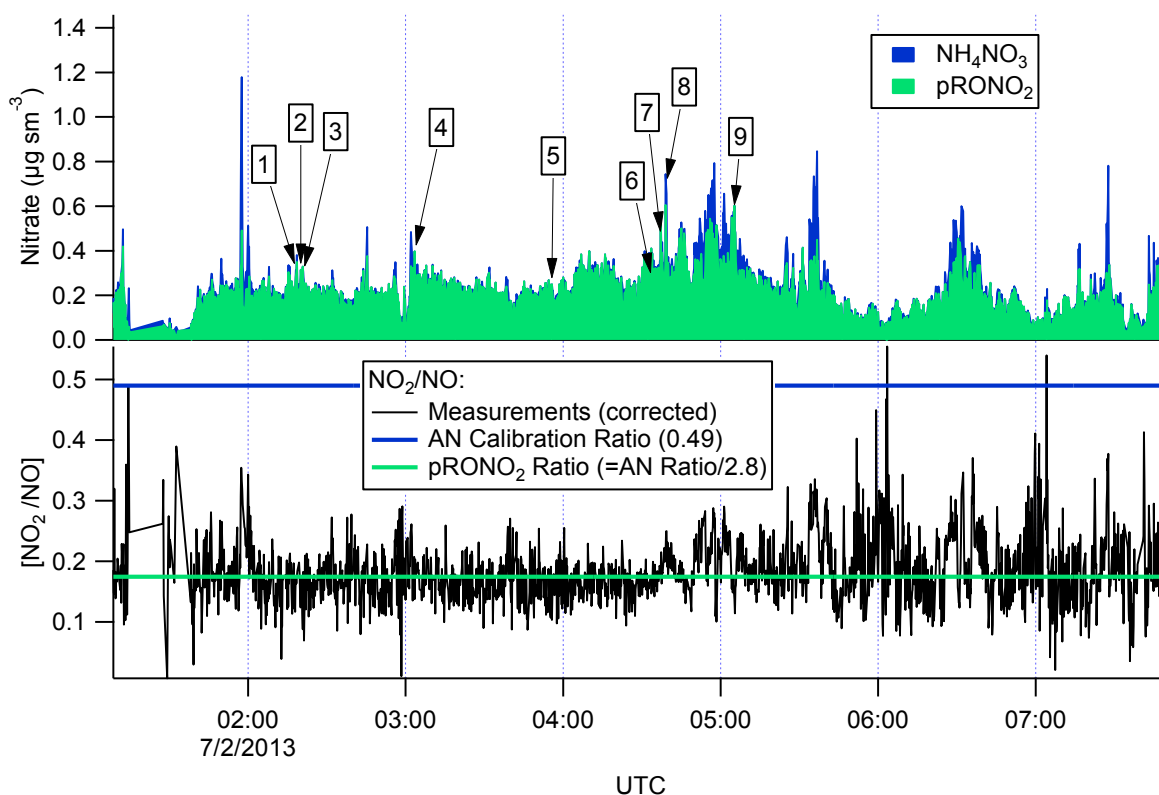
526 total aerosol mass is elevated, semivolatile organic compounds may re-partition to the aerosol  
527 phase, contributing a non-pRONO<sub>2</sub> driven variability in ΔOA. For example, if some gas phase  
528 IEPOX is present in the residual layer, it may be taken up into the highly acidic aerosol from the  
529 power plants. Alternatively, very polar gas-phase compounds could partition further into the  
530 higher liquid water associated with the sulfate in the plume. Therefore, in-plume organic aerosol  
531 increases cannot be attributed clearly to NO<sub>3</sub> + isoprene SOA production, so we do not use  
532 them in the SOA yield calculations.

533  
534 This leaves consideration 2b, whether all increase in nitrate mass is due to NO<sub>3</sub> + isoprene  
535 reactions. Here we must evaluate the possibility of inorganic nitrate aerosol production in these  
536 high-NO<sub>x</sub> plumes. Fine-mode aerosol inorganic nitrate can be formed by the (reversible)  
537 dissolution of HNO<sub>3(g)</sub> into aqueous aerosol. In dry aerosol samples, inorganic nitrate is typically  
538 in the form of ammonium nitrate (NH<sub>4</sub>NO<sub>3</sub>), when excess ammonium is available after  
539 neutralization of sulfate as (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and NH<sub>4</sub>(HSO<sub>4</sub>). Because of the greater stability of  
540 ammonium sulfate salt relative to ammonium nitrate, in high-sulfate plumes with limited  
541 ammonium, inorganic nitrate aerosol will typically evaporate as HNO<sub>3(g)</sub> (Guo et al., 2015)  
542 (reaction R3):



546 Inorganic nitrate can also form when crustal dust (e.g. CaCO<sub>3</sub>) or seasalt (NaCl) are available.  
547 Uptake of HNO<sub>3</sub> is rendered favorable by the higher stability of nitrate mineral salts, evaporating  
548 CO<sub>2</sub> or HCl. Inorganic nitrate can also be produced by the heterogeneous uptake of N<sub>2</sub>O<sub>5</sub> onto  
549 aqueous aerosol; Edwards et al. (2017) demonstrated that this process is negligible relative to  
550 NO<sub>3</sub> + BVOC for the July 2 SENEX night flight considered here.

551  
552 There are several lines of evidence that the observed nitrate aerosol is organic and not  
553 inorganic. First, examination of the NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> (interference-corrected *m/z* 46:*m/z* 30) ratio  
554 measured by the aircraft AMS (Fig. 3) shows a ratio throughout the July 2 flight, including the  
555 selected plumes, that is substantially lower than that from the bracketing ammonium nitrate  
556 calibrations. This lower AMS measured NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> ratio has been observed for organic nitrates  
557 (Farmer et al., 2010), and some mineral nitrates (e.g. Ca(NO<sub>3</sub>)<sub>2</sub> and NaNO<sub>3</sub>, (Hayes et al.,  
558 2013)), which are not important in this case because aerosol was dominantly submicron. As  
559 described above, we can separate the observed AMS nitrate signal into pRONO<sub>2</sub> and inorganic  
560 nitrate contributions. These mass loadings are also shown in Fig. 3, indicating dominance of  
561 pRONO<sub>2</sub> throughout the flight.



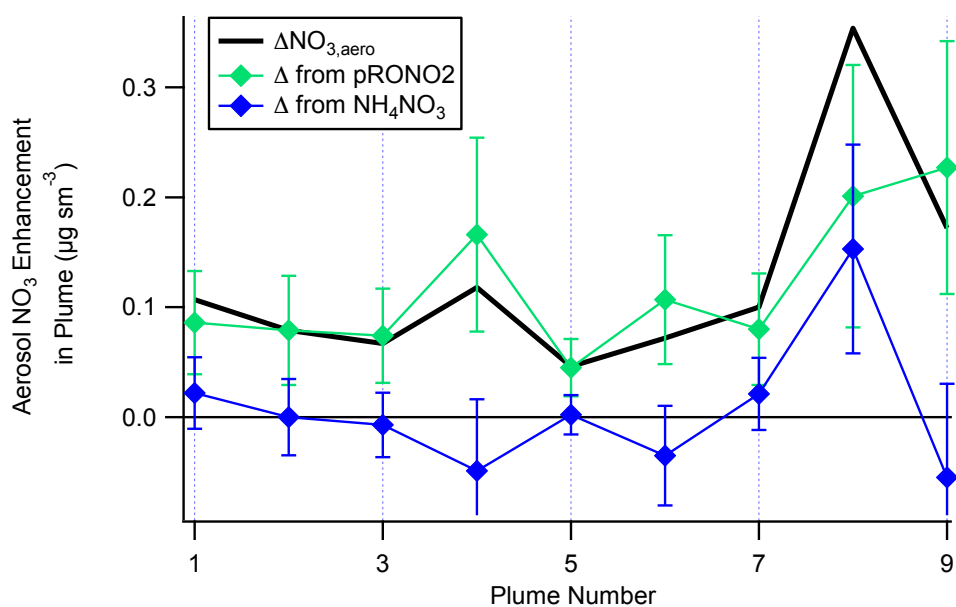
563  
 564 **Figure 3.** For the flight under consideration, the estimated relative contributions of ammonium  
 565 and organic nitrate to the total corrected nitrate signal (top panel) was calculated from the ratios  
 566 of the corrected peaks at  $m/z$  30 and 46 (lower panel). Each of the plumes is identified here by  
 567 plume number. The ratios of  $\text{NO}_2^+/\text{NO}^+$  (black data in the lower panel) from the corrected peaks  
 568 at  $m/z$  46 and 30, respectively, are compared to the ratios expected for ammonium nitrate (AN  
 569 Calibration Ratio, blue horizontal line at 0.49) or organic nitrate (pRONO<sub>2</sub> Ratio, green  
 570 horizontal line at 0.175 which is estimated from the AN calibration ratio using multiple data sets  
 571 (see discussion in Supplemental Information). The measured ratio for most of the flight is more  
 572 characteristic of organic nitrate than ammonium nitrate.

573  
 574 We can also employ the comparison of other AMS-measured aerosol components during the  
 575 individual plumes to assess the possibility of an inorganic nitrate contribution to total measured  
 576 nitrate. Fig. S5a shows that the in-plume increases in sulfate are correlated with increases in  
 577 ammonium with an  $R^2$  of 0.4. The observed slope of 5.4 is characteristic of primarily  $(\text{NH}_4)\text{HSO}_4$ ,  
 578 which indicates that the sulfate mass is not fully neutralized by ammonium. We note, however,  
 579 that if the largest observed aerosol nitrate increase is due solely to ammonium nitrate, the  
 580 ammonium increase would be only  $0.11 \mu\text{g m}^{-3}$ , which would be difficult to discern from the  $\text{NH}_4$   
 581 variability of order  $0.11 \mu\text{g m}^{-3}$ . However, the slope is consistent with incomplete neutralization  
 582 of the sulfate by ammonium, which would make  $\text{HNO}_{3(\text{g})}$  the more thermodynamically favorable  
 583 form of inorganic nitrate. The ion balance for the ammonium nitrate calibration particles and the  
 584 plume enhancements are shown in Fig. S5b. Complete neutralization of the calibration aerosols  
 585 is nearly always within the gray 10% uncertainty band for the relative ionization efficiency of



586 ammonium (Bahreini et al., 2009). In contrast, many of the plume enhancements are near the  
587 1:2 line (as primarily ammonium bisulfate) within the combined 10% ammonium and 15%  
588 sulfate uncertainty error bars or without ammonium (sulfuric acid). Thus,  $\text{NH}_4\text{NO}_3$  is unlikely to  
589 be stable in the aerosol phase under the conditions of these plumes, consistent with the AMS  
590 observations.

591  
592 A plot of the calculated plume enhancements from the derived apportionment into organic  
593 ( $\text{pRONO}_2$ ) and inorganic (ammonium) nitrate is shown in Fig. 4. The increases in aerosol nitrate  
594 for nearly all of the plumes appear to be mostly due to enhancements in  $\text{pRONO}_2$ . Based on  
595 these considerations, we conclude that in-plume  $\text{pRONO}_2$  mass increases are a consequence  
596 (and thus a robust measure) of organic nitrate aerosol produced from  $\text{NO}_3 + \text{isoprene}$ . Since  
597 each isoprene molecule condensing will have one nitrate group, the ratio of these increases to  
598 isoprene loss is a direct measure of the molar organic aerosol yield from  $\text{NO}_3$ -isoprene  
599 oxidation.



600

601

602 **Figure 4.** The contribution of each species to the nitrate enhancements in each of the plumes,  
603 showing that the enhancements in most of the plumes are mainly due to enhancements in  
604 organic nitrate, with the exception of Plume 8 which had enhancements in both organic and  
605 ammonium nitrate. Error bars are estimated from the measurement variability, the UMR  
606 corrections to the nitrate signals, apportionment between organic and inorganic nitrate, and the  
607 total nitrate uncertainty (see Supplemental Information).

608

609 Table 1 shows the selected plumes to be used for yield analysis. Wherever possible, multiple  
610 points have been averaged for in-plume and background isoprene and nitrate aerosol  
611 concentrations; in each case the number of points used is indicated and the corresponding  
612 standard deviations are reported. In two cases (2:20 and 3:03 plumes), the plumes were so  
613 narrow that only a single point was measured in-plume at the 10 s time resolution of the PTR-

614 MS and AMS; for these “single-point” plumes it is not possible to calculate error bars. Error bars  
615 were determined using the standard deviations calculated for in-plume and background  
616 isoprene and nitrate aerosol concentrations, accounting also for the additional uncertainty in the  
617 AMS measurement described in the caption to Figure 4, and propagated through the yield  
618 formula detailed in the following section.  
619

620 **Table 1.** List of plumes used in this NO<sub>3</sub> + isoprene SOA yield analysis. For each plume, the  
 621 delta-values listed indicate the difference between in-plume and outside-plume background in  
 622 average observed concentration, and the standard deviations (SD) are the propagated error  
 623 from this subtraction. (For ΔNO<sub>3</sub> from pRONO<sub>2</sub>, the standard deviations also include error  
 624 propagated as described in the caption for Figure 4) After each plume number, the numbers of  
 625 points averaged for isoprene (10 s resolution) and AMS (10 s resolution), respectively, are  
 626 listed. Because the isoprene data were reported at a lower frequency, these numbers are  
 627 typically lower to cover the same period of time. Plume numbers annotated with \* indicate brief  
 628 plumes for which only single-point measurements of in-plume aerosol composition were  
 629 possible. Additional AMS and auxiliary data from each plume is included in the Supplemental  
 630 Information, Table S3.

plume number [#isop/#AMS]	7/2/13 plume time (UTC)	P(NO <sub>3</sub> ) (ppbv hr <sup>-1</sup> )	ΔISOP (ppt) [± SD]	ΔNO <sub>3,aero</sub> (μg m <sup>-3</sup> ) [± SD]	ΔNO <sub>3</sub> from pRONO <sub>2</sub> (μg m <sup>-3</sup> ) [± SD]	ΔNO <sub>3</sub> from NH <sub>4</sub> NO <sub>3</sub> (μg m <sup>-3</sup> ) [± SD]
Typical variability (μg m <sup>-3</sup> ):				0.05	0.05	0.05
1 [2/3]	2:18	0.9	-335 [128]	0.107 [0.039]	0.086 [0.047]	0.022 [0.012]
2 [*]	2:20	0.8	-404	0.079	0.079 [0.049]	0
3 [4/5]	2:21	1.2	-228 [121]	0.067 [0.039]	0.074 [0.043]	-0.007 [0.027]
4 [*]	3:03	1.4	-453	0.118	0.166 [0.088]	-0.049
5** [3/4]	3:55	1.0	-255 [251]	0.046 [0.019]	0.045 [0.026]	0.002 [0.015]
6 [2/2]	4:34	0.6	-713 [219]	0.072 [0.031]	0.107 [0.059]	-0.035 [0.029]
7 [5/6]	4:37	0.8	-298 [197]	0.100 [0.082]	0.080 [0.051]	0.021 [0.034]
8*** [2/3]	4:39	0.9	-443 [75]	0.354 [0.058]	0.201 [0.120]	0.153 [0.057]
9 [7/8]	5:04	0.6	-293 [131]	0.172 [0.048]	0.227 [0.115]	-0.055 [0.042]

631 \*\*Plume 5 has the smallest ΔNO<sub>3,aero</sub> and may be affected by background pRONO<sub>2</sub> variability.

632 \*\*\*Plume 8 has a measurable increase in inorganic nitrate as well as organic.

## 633 4.2 SOA yield analysis

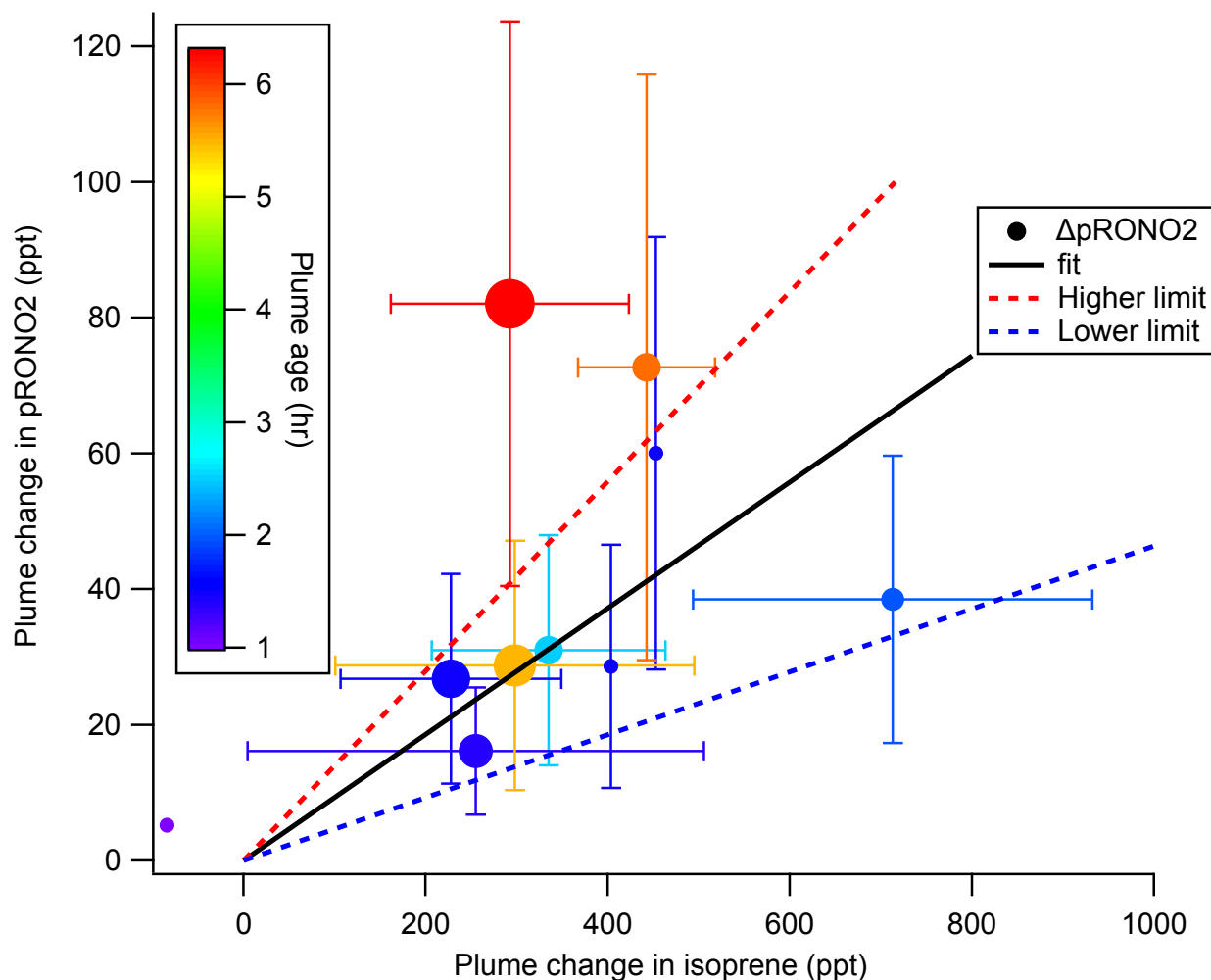
634 A **molar** SOA yield refers to the number of molecules of aerosol organic nitrate produced per  
 635 molecule of isoprene consumed. In order to determine molar SOA yields from the data  
 636 presented in Table 1, we convert the aerosol organic nitrate mass loading differences to mixing  
 637 ratio differences (ppt) using the NO<sub>3</sub> molecular weight of 62 g mol<sup>-1</sup> (the AMS organic nitrate

638 mass is the mass only of the  $-\text{ONO}_2$  portion of the organonitrate aerosol). At standard  
 639 conditions of 273 K and 1 atm (all aerosol data are reported with this STP definition), 1000 ppt  
 640  $\text{NO}_3 = 2.77 \mu\text{g m}^{-3}$ , so each  $\Delta M_{\text{pRONO}_2}$  is multiplied by  $361 \text{ ppt } (\mu\text{g m}^{-3})^{-1}$  to determine this molar  
 641 yield:

$$642 \quad Y_{\text{SOA,molar}} = \frac{(p\text{RONO}_2_{\text{plume}} \pm SD_{p\text{RONO}_2_{\text{plume}}}) - (p\text{RONO}_2_{\text{bkg}} \pm SD_{p\text{RONO}_2_{\text{bkg}}})}{-[(isop_{\text{plume}} \pm SD_{isop_{\text{plume}}}) - (isop_{\text{bkg}} \pm SD_{isop_{\text{bkg}}})]} \times \frac{361 \text{ ppt } \text{NO}_3}{\mu\text{g m}^{-3}} \quad (3)$$

644 The SOA molar yields resulting from this calculation are shown in Table 2, spanning a range of  
 645 5-28%, with uncertainties indicated based on the SDs in measured AMS and isoprene  
 646 concentrations. In addition to this uncertainty based on measurement precision and ambient  
 647 variability, there is an uncertainty of 50% in the AMS derived-organic nitrate mass loadings (see  
 648 SI) and 25% in the PTR-MS isoprene concentrations (Warneke et al., 2016). The average molar  
 649 pRONO<sub>2</sub> yield across all plumes, with each point weighed by the inverse of its standard  
 650 deviation, is 9%. (As noted below, the yield appears to increase with plume age, so this average  
 651 obscures that trend.) An alternate graphical analysis of molar SOA yield from all nine plumes  
 652 plus one 'null' plume (03:14, in which no isoprene had yet reacted and thus not included in  
 653 Tables 1 and 2) obtains the same average molar yield of 9% (Fig. 5). Here, the molar yield is  
 654 the slope of a plot of plume change in pRONO<sub>2</sub> vs plume change in isoprene. The slope is  
 655 determined by a linear fit with points weighted by the square root of the number of AMS data  
 656 points used to determine in-plume pRONO<sub>2</sub> in each case. This slope error gives a rather narrow  
 657 uncertainty range for the slope (0.0930 +/- 0.0011); to obtain an upper limit in the uncertainty of  
 658 this molar yield we apply the combined instrumental uncertainties, based on adding in  
 659 quadrature the PTR-MS uncertainty of 5% and the AMS uncertainty of 50%. This gives an  
 660 overall uncertainty of 50.2%, resulting in upper and lower limit slopes of 0.140 and 0.046,  
 661 respectively; we use this maximum uncertainty estimate to report the average molar yield as 9%  
 662 (+/- 5%). We have not corrected the calculated yields for the possibility of NO<sub>3</sub> heterogeneous  
 663 uptake, which could add a nitrate functionality to existing aerosol. Such a process could be rapid  
 664 if the uptake coefficient for NO<sub>3</sub> were 0.1, a value characteristics of unsaturated substrates (Ng  
 665 et al., 2017), but would not contribute measurably at more conventional NO<sub>3</sub> uptake coefficients  
 666 of 0.001 (Brown and Stutz, 2012).

667  
 668  
 669

671  
672

673 **Figure 5.** SOA molar yield can be determined as the slope of  $\Delta p\text{RONO}_2$  vs.  $\Delta \text{isoprene}$ , both in  
 674 mixing ratio units. The linear fit is weighted by square root of number of points used to  
 675 determine each in-plume  $p\text{RONO}_2$ , with intercept held at zero. The slope coefficient  $\pm$  one  
 676 standard deviation is  $0.0930 \pm 0.0011$ . Larger “outside” high and low limits of the slope (shown  
 677 as dashed red and blue lines) are obtained by adding and subtracting from this slope the  
 678 combined instrumental uncertainties, based on adding in quadrature the PTR-MS uncertainty of  
 679 5% and the AMS uncertainty of 50%. This gives an overall uncertainty of 50.2%, resulting in  
 680 upper and lower limit slopes of 0.140 and 0.046, respectively. Points are colored by plume age,  
 681 and size scaled by square root of number of points (the point weight used in linear fit). This plot  
 682 and fit includes the nine plumes listed in Tables 1 and 2, as well as the 03:14 “unreacted” plume  
 683 (at  $\Delta \text{isoprene} = -84$  ppt). Error bars on isoprene are the propagated standard deviations of the  
 684 (in plume - out plume) differences, for plumes in which multi-point averages were possible. Error  
 685 bars on  $p\text{RONO}_2$  are the same as in Figure 4, converted to ppt. The points without error bars  
 686 are single-point plumes.

687

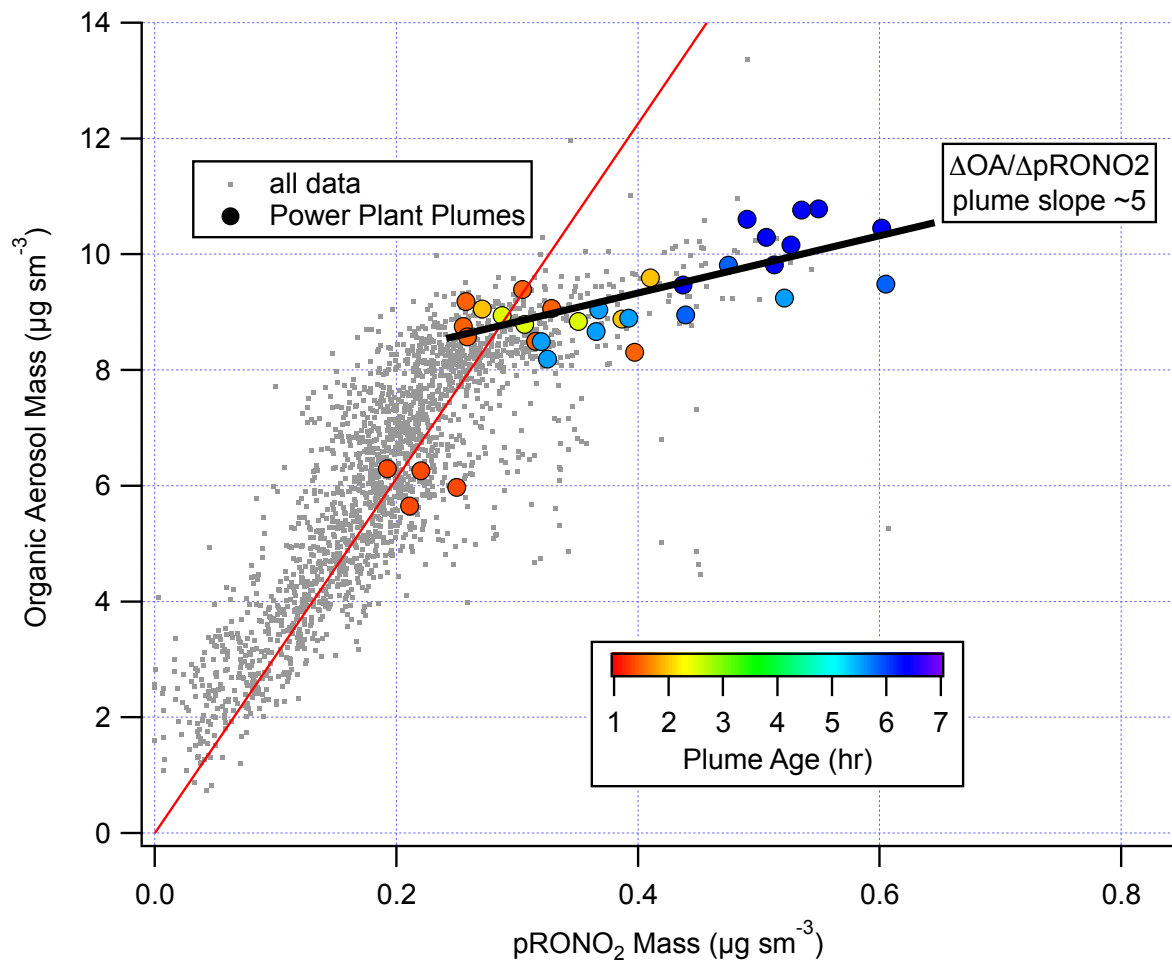
688 To estimate SOA **mass** yields, we need to make some assumption about the mass of the  
689 organic molecules containing the nitrate groups that lead to the observed nitrate aerosol mass  
690 increase. The observed changes in organic aerosol are too variable to be simply interpreted as  
691 the organic portion of the aerosol organic nitrate molecules. We conservatively assume the  
692 organic mass to be approximately double the nitrate mass ( $62 \text{ g mol}^{-1}$ ), based on an “average”  
693 molecular structure of an isoprene nitrate with 3 additional oxygens: e.g. a tri-hydroxynitrate  
694 (with organic portion of formula  $\text{C}_5\text{H}_{11}\text{O}_3$ ,  $119 \text{ g mol}^{-1}$ ), consistent with 2<sup>nd</sup>-generation oxidation  
695 product structures suggested in Schwantes, et al. (Schwantes et al., 2015). Based on this  
696 assumed organic to nitrate ratio, all plumes’ expected organic mass increases would be less  
697 than the typical variability in organic of  $0.75 \mu\text{g m}^{-3}$ . This assumed structure is consistent with  
698 oxidation of both double bonds, which appears to be necessary for substantial condensation of  
699 isoprene products, and which structures would have calculated vapor pressures sufficiently low  
700 to partition to the aerosol phase (Rollins et al., 2009). Another possible route to low vapor  
701 pressure products is intramolecular H rearrangement reactions, discussed below in Section 4.3,  
702 which would not require oxidant reactions at both double bonds. In the case of oxidant reactions  
703 at both double bonds, it is difficult to understand how the second double bond would be oxidized  
704 unless by another nitrate radical, which would halve these assumed organic to nitrate ratios  
705 (assuming the nitrate is retained in the molecules). On the other hand, any organic nitrate  
706 aerosol may lose  $\text{NO}_3$  moieties, increasing the organic to nitrate ratio. Given these uncertainties  
707 in both directions, we use the assumed “average” structure above to guess an associated  
708 organic mass of double the nitrate mass. Thus, to estimate SOA mass yield, we multiply the  
709 increase in organic nitrate aerosol mass concentration by three (i.e.,  $2 \times \Delta M_{p\text{RONO}_2} + \Delta M_{p\text{RONO}_2}$ ),  
710 and divide by the observed decrease in isoprene, converted to  $\mu\text{g m}^{-3}$  by multiplying by 329 ppt  
711 ( $\mu\text{g m}^{-3}$ )<sup>-1</sup>, the conversion factor based on isoprene’s molecular weight of  $68.12 \text{ g mol}^{-1}$ .

$$713 Y_{SOA, mass} = \frac{(p\text{RONO}_2_{plume} \pm SD_{p\text{RONO}_2_{plume}}) - (p\text{RONO}_2_{bkg} \pm SD_{p\text{RONO}_2_{bkg}})}{-[(isop_{plume} \pm SD_{isop_{plume}}) - (isop_{bkg} \pm SD_{isop_{bkg}})]} \times 3 \times \frac{329 \text{ ppt}}{\mu\text{g m}^{-3}} \quad (4)$$

714 Note that the SOA mass yield reported here is based on the (assumed) mass of organic aerosol  
715 plus the (organo)nitrate aerosol formed in each plume. If instead the yield were calculated using  
716 only the assumed increase in **organic** mass (i.e.,  $2 \times \Delta M_{p\text{RONO}_2}$  instead of  $3 \times \Delta M_{p\text{RONO}_2}$ ), which  
717 would be consistent with the method used in Rollins, et al. (Rollins et al., 2009) and Brown et al.  
718 (Brown et al., 2009), the mass yields would be 2/3 the values reported here. However, since  
719 SOA mass yield is typically defined based on the total increase in aerosol mass, we use the  
720 definition with the sum of the organic and nitrate mass here. This results in an average SOA  
721 mass yield of 27%, with propagated instrumental errors (see caption to Fig. 5) giving a range of  
722 27% +/- 14%.

724 We note also that correlation of in-plume increases in OA with  $p\text{RONO}_2$  (Fig. 6) point to a  
725 substantially larger 5:1 organic-to-nitrate ratio; if this were interpreted as indicating that the  
726 average molecular formula of the condensing organic nitrate has 5 times the organic mass as  
727 nitrate, this would increase the SOA mass yields reported here. However, due to the  
728 aforementioned possibility of additional sources of co-condensing organic aerosol, which led us  
729 to avoid using  $\Delta\text{OA}$  in determining SOA yields, we do not consider this to be a direct indication  
730

731 of the molecular formula of the condensing organic nitrate. Including OA in the SOA yield  
732 determination, based on this 5:1 slope rather than the assumed 2:1 OA:pRONO<sub>2</sub>, would give  
733 2.5 times larger SOA mass yields than reported here.  
734



735  
736 **Figure 6.** Correlation of organic aerosol mass concentration with pRONO<sub>2</sub> mass concentration  
737 for the full 2 July flight (grey points and red fit line, fitted slope and thus average OA/pRONO<sub>2</sub>  
738 mass ratio of ~30) and for the points during the selected plumes (colored points, colored by  
739 plume age, average OA/pRONO<sub>2</sub> mass ratio of ~ 5).

740  
741

742 **Table 2.** SOA Yields for each plume observation, estimated plume age, and likely origin. See  
 743 text for description of uncertainty estimates. For the mass yields, the calculated SOA mass  
 744 increase includes both the organic and (organo)nitrate aerosol mass; the measurements for OA  
 745 increases shown in Figure 6 do not include the nitrate mass.

plume number	plume time (UTC)	SOA molar yield (fraction) [± SD]	SOA mass yield (fraction) [± SD]	plume age from O <sub>3</sub> / NO <sub>2</sub> clock assuming S=1 (hours)	Likely NO <sub>x</sub> origin & altitude (m)
1	7/2/13 2:18	0.09 [0.06]	0.25 [0.17]	2.5	Greene County @ 540 m
2	7/2/13 2:20	0.07	0.21	1.5	<i>ibid</i>
3	7/2/13 2:21	0.12 [0.10]	0.32 [0.25]	1.5	<i>ibid</i>
4	7/2/13 3:03	0.13	0.36	1.5	Gaston @ 720 m
5	7/2/13 3:55	0.06 [0.07]	0.17 [0.20]	1.4	Miller / Gorgas @ 690 m
6	7/2/13 4:34	0.05 [0.03]	0.15 [0.09]	2	<i>ibid</i>
7	7/2/13 4:37	0.10 [0.09]	0.26 [0.24]	5.5	<i>ibid</i>
8	7/2/13 4:39	0.16 [0.10]	0.45 [0.28]	5.8	Miller / Gorgas @ 1120 m
9	7/2/13 5:04	0.28 [0.19]	0.77 [0.52]	6.3	Gaston @ 1280 m

746  
747



748 **Table 3.** Several caveats to the present SOA yields analysis are listed below, alongside the  
 749 expected direction each would adjust the estimated yields. Because we do not know whether or  
 750 how much each process may have occurred in the studied plumes, we cannot quantitatively  
 751 assess the resulting uncertainties, so we simply list them here. See text above for more detailed  
 752 discussion.

Process	Effect on determined SOA yield
Organic nitrate aerosol loses NO <sub>3</sub> functional group	Larger, because the non-nitrate OA would not be counted in this analysis
Both double bonds in isoprene are oxidized by NO <sub>3</sub> : two nitrates per condensing molecule	Smaller, because the assumed organic to nitrate mass ratio assumes one nitrate per molecule
NO <sub>3</sub> oxidizes daytime isoprene oxidation products (e.g. ISOPOOH) to make new aerosol	Smaller, because this would produce organic nitrate aerosol without corresponding decrease in isoprene, so that some of existing SOA production is mis-attributed to isoprene + NO <sub>3</sub>
Assumed organic to nitrate mass ratio is incorrect	Unknown direction of effect, depends on whether assumed ratio is high or low
Daytime-produced IEPOX uptake onto acidic particles	No effect (only changes ΔOA, not nitrate)
Suppression of O <sub>3</sub> + monoterpene or O <sub>3</sub> + isoprene SOA in plumes	No effect (only changes ΔOA, not nitrate)

753  
 754 Finally, the large range in observed yields can be interpreted by examining the relationship to  
 755 estimated plume age. Using the slope of O<sub>3</sub> to NO<sub>2</sub> (Eq. 1) to estimate plume age as described  
 756 above, a weak positive correlation is observed (Table 2, Fig. S4), suggesting that as the plume  
 757 ages, later-generation chemistry results in greater partitioning to the condensed phase of NO<sub>3</sub> +  
 758 isoprene organonitrate aerosol products. This is consistent with the observation by Rollins et al.  
 759 (Rollins et al., 2009) that 2nd-generation oxidation produced substantially higher SOA yields  
 760 than the oxidation of the first double bond alone, but we note that these mass yields (averaging  
 761 27%, would be 18% using the organic mass only) are higher than even the largest yield found in  
 762 that chamber study (14%, used organic mass only).

763  
 764 We observe increasing SOA yield, from a molar yield of around 10% at 1.5 hours up to 30% at 6  
 765 hours of aging. The lowest yields observed are found in the most recently emitted plumes,  
 766 suggesting the interpretation of the higher yields as a consequence of longer aging timescales  
 767 in the atmosphere.

### 768 **4.3 Mechanistic considerations**

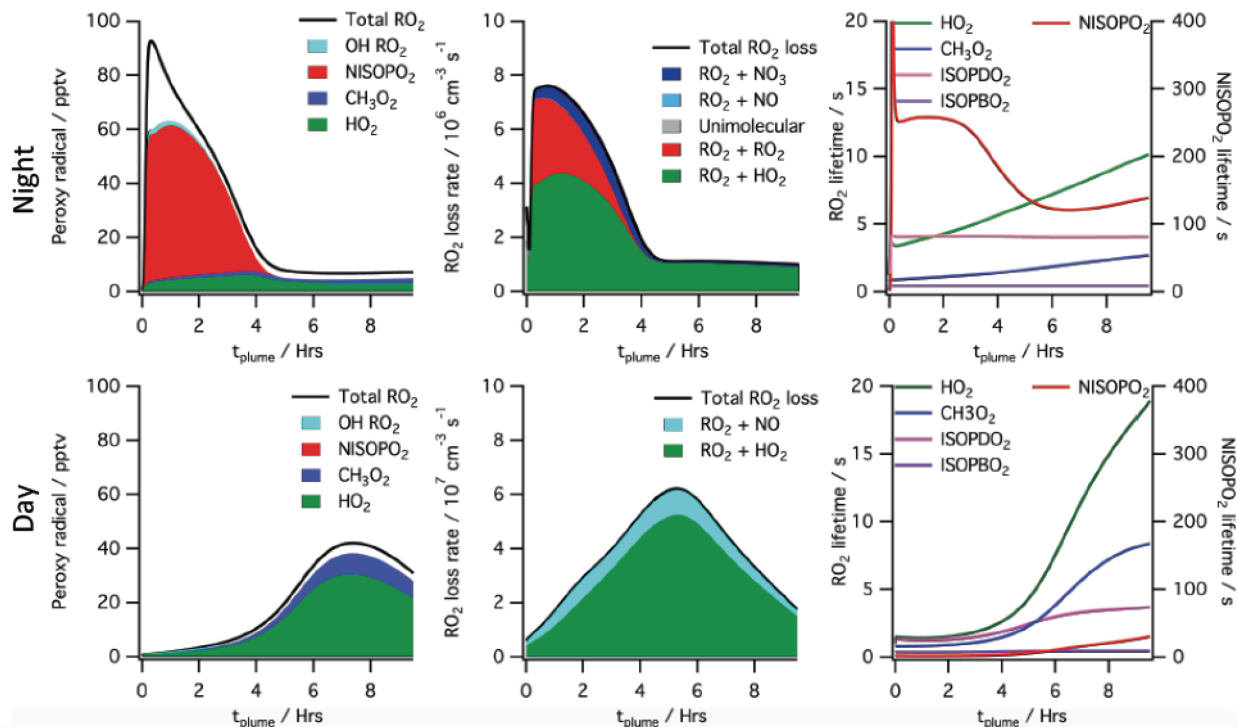
769  
 770 These larger SOA mass yields from field determinations (average 27%) relative to chamber  
 771 work (12 – 14%, see introduction) may arise for several reasons. We first assess the volatility of  
 772 assumed first- and second-generation products using group contribution theory in order to  
 773 predict partitioning. After a single oxidation step, with a representative product assumed to be a

774 C<sub>5</sub> hydroperoxynitrate, the saturation vapor pressure estimated by group contribution theory  
775 (Pankow and Asher, 2008) at 283 K would be  $2.10 \times 10^{-3}$  Torr ( $C^* = 1.7 \times 10^4 \mu\text{g m}^{-3}$  for MW =  
776  $147 \text{ g mol}^{-1}$ ), while a double-oxidized isoprene molecule (assuming a C<sub>5</sub> dihydroxy dinitrate) has  
777 an estimated vapor pressure of  $7.95 \times 10^{-8}$  Torr ( $C^* = 1.01 \mu\text{g m}^{-3}$  for MW =  $226 \text{ g mol}^{-1}$ ). This  
778 supports the conclusion that while the first oxidation step produces compounds too volatile to  
779 contribute appreciably to aerosol formation, oxidizing both double bonds of the isoprene  
780 molecule is sufficient to produce substantial partitioning, consistent with Rollins et al. (Rollins et  
781 al., 2009). This is also true if the second double bond is not oxidized by nitrate (group  
782 contribution estimate  $P_{\text{vap}}$  for a C<sub>5</sub> tri-hydroxy nitrate is  $7.7 \times 10^{-8}$  Torr,  $C^* = 0.79 \mu\text{g m}^{-3}$  for MW =  
783  $181 \text{ g mol}^{-1}$ ). These C\* saturation concentration values suggest that no dimer formation or  
784 oligomerization is *required* to produce low-enough volatility products to condense to the aerosol  
785 phase; however, such oligomerization would result in more efficient condensation. The fact that  
786 Rollins et al. (Rollins et al., 2009) did not observe larger mass yields may indicate that it takes  
787 longer than a typical chamber experiment timescale to reach equilibrium, or that this absorptive  
788 partitioning model did not accurately capture those experiments, or that substantial loss of  
789 semivolatiles to the chamber walls (e.g. (Krechmer et al., 2016)) suppressed apparent yields.

790  
791 Determination of yields from ambient atmospheric data differs from chamber determinations in  
792 several additional respects. First, ambient measurements do not suffer from wall loss effects,  
793 such that no corrections are necessary for loss of aerosol or semi-volatile gases (Matsunaga  
794 and Ziemann, 2010;Krechmer et al., 2016). Second, ambient measurements take place on the  
795 aging time scale of the atmosphere rather than a time scale imposed by the characteristics of  
796 the chamber or the choice of oxidant addition. Third, the typical lifetime of the initially produced  
797 nitrooxy-isoprene-RO<sub>2</sub> radical is more representative of the ambient atmosphere rather than a  
798 chamber. The unique conditions of a high NO<sub>x</sub> power plant plume affect lifetime and fates of  
799 peroxy radicals, as described below.

800  
801 To help interpret these in-plume peroxy radical lifetimes, a box model calculation using the  
802 MCM v3.3.1 chemistry scheme was run (see details in Supplemental Information). This box  
803 model shows substantially longer peroxy radical lifetimes during nighttime than daytime,  
804 initializing with identical plume-observed conditions. These long peroxy radical lifetimes may  
805 have consequences for comparison to chamber experiments: for example, in Schwantes'  
806 (Schwantes et al., 2015) chamber experiment on the NO<sub>3</sub> + isoprene reaction mechanism, the  
807 HO<sub>2</sub>-limited nitrooxy-RO<sub>2</sub> lifetime was at maximum 30 s. In the plumes investigated in this study,  
808 peroxy radical lifetimes are predicted to be substantially longer (>200 s early in the night, see  
809 Fig. 7), allowing for the possibility of different bimolecular fates, or of unimolecular  
810 transformations of the peroxy radicals that may result in lower-volatility products (e.g., auto-  
811 oxidation to form highly oxidized molecules (Ehn et al., 2014)).

812  
813



814  
 815 **Figure 7.** Simulated peroxy radical concentration (left), loss rates (middle), and lifetime (right),  
 816 using the MCM v3.3.1 chemical mechanism, for conditions typical of a nighttime intercepted  
 817 power plant plume (top) and the same plume initial conditions run for daytime simulation  
 818 (bottom, local noon occurs at 5 hrs). Included are total peroxy radical concentration and losses,  
 819 as well as the highlighted subclasses HO<sub>2</sub>, CH<sub>3</sub>O<sub>2</sub>, total nitrooxy-isoprene-RO<sub>2</sub>, and the total  
 820 hydroxy-isoprene-RO<sub>2</sub> produced from OH oxidation. The righthand panels show HO<sub>2</sub>, CH<sub>3</sub>O<sub>2</sub>  
 821 and the dominant hydroxy-isoprene-RO<sub>2</sub> ISOPBO<sub>2</sub> and ISOPDO<sub>2</sub> ( $\beta$ -hydroxy-peroxy radicals  
 822 from OH attack at carbons 1 and 4 respectively) lifetime on the left axis and nitrooxy-isoprene-  
 823 RO<sub>2</sub> on the right axis, showing nighttime lifetimes an order of magnitude longer than daytime for  
 824 this NO<sub>3</sub> + isoprene derived RO<sub>2</sub> radical (NISOPO<sub>2</sub>).

825  
 826 The typically assumed major fate of nighttime RO<sub>2</sub> in the atmosphere is reaction with HO<sub>2</sub> to  
 827 yield a hydroperoxide, NO<sub>3</sub>-ROOH. This is shown in the model output above as the green  
 828 reaction, and is responsible for half of early RO<sub>2</sub> losses in the MCM modeled plume. Schwantes  
 829 *et al.* (Schwantes *et al.*, 2015) proposed reaction of these nighttime derived hydroperoxides with  
 830 OH during the following day as a route to epoxides, which in turn can form SOA via reaction  
 831 with acidic aerosol. Reaction of hydroperoxides with nighttime generated OH may similarly  
 832 provide a route to SOA through epoxides, albeit more slowly than that due to photochemically  
 833 generated OH.

834  
 835 The predicted longer nighttime peroxy radical lifetimes may enable unique chemistry. For  
 836 example, if nitrooxy-isoprene-RO<sub>2</sub> self-reactions are substantially faster than assumed in the  
 837 MCM, as suggested by Schwantes *et al.* (Schwantes *et al.*, 2015), RO<sub>2</sub>+RO<sub>2</sub> reactions may  
 838 compete with the HO<sub>2</sub> reaction even more than shown in Fig. 7, and dimer formation may be  
 839 favored at night, yielding lower volatility products. The 5:1 AMS Organic:Nitrate ratio observed in

840 the SOA formed in Rollins et al. (Rollins et al., 2009) , and consistent with aggregated  
841 observations reported here, may suggest that in some isoprene units the nitrate is re-released  
842 as NO<sub>2</sub> in such oligomerization reactions. We note that this larger organic to nitrate ratio would  
843 mean higher SOA mass yields than estimated in Table 2.

844  
845 Alternatively, longer nighttime peroxy radical lifetimes may allow sufficient time for  
846 intramolecular reactions to produce condensable products. This unimolecular isomerization  
847 (auto-oxidation) of initially formed peroxy radicals is a potentially efficient route to low-volatility,  
848 highly functionalized products that could result in high aerosol yields. For OH-initiated oxidation  
849 of isoprene, laboratory relative rate experiments found the fastest 1,6-H-shift isomerization  
850 reaction to occur for the hydroxy-isoprene-RO<sub>2</sub> radical at a rate of 0.002 s<sup>-1</sup> (Crouse et al.,  
851 2011), meaning that peroxy radicals must have an ambient lifetime of >500 s for this process to  
852 be dominant. As shown in Fig. 7, the simulated power plant plume peroxy radical lifetimes are  
853 long (>200 s), so an isomerization reaction at this rate may play a significant role. However, a  
854 recent study has demonstrated that OH-initiated and NO<sub>3</sub>-initiated RO<sub>2</sub> radicals from the same  
855 precursor VOC can have very different unimolecular reactive fates due to highly structurally  
856 sensitive varying rates of reactions of different product channels (Kurtén et al., 2017). A similar  
857 theoretical study on the rate of unimolecular autooxidation reactions of nitrooxy-isoprene-RO<sub>2</sub>  
858 radicals would be valuable to help determine under what conditions such reactions might occur,  
859 and this knowledge could be applied to comparing chamber and field SOA yields.

#### 860 **4.4 Atmospheric implications and needs for future work**

861 Because this paper proposes higher SOA yield for the NO<sub>3</sub> + isoprene reaction than measured  
862 in chamber studies, we conclude with some discussion of the implications for regional aerosol  
863 burdens, and further needs for investigation in the NO<sub>3</sub> + isoprene system.

864  
865 Using an isoprene + NO<sub>3</sub> yield parameterization that gave a 12% SOA mass yield at 10 μg m<sup>-3</sup>,  
866 Pye et al. (2010) found that adding the NO<sub>3</sub> + isoprene oxidation pathway increased isoprene  
867 SOA mass concentrations in the southeastern United States by about 30%, increases of 0.4 to  
868 0.6 μg m<sup>-3</sup>. The larger NO<sub>3</sub> + isoprene SOA mass yields suggested in this paper, with average  
869 value of 30%, could double this expected NO<sub>3</sub> radical enhancement of SOA production.  
870 Edwards et al. (2017) concluded that the southeast U.S. is currently in transition between NO<sub>x</sub>-  
871 independent and NO<sub>x</sub>-controlled nighttime BVOC oxidation regime. If NO<sub>3</sub>-isoprene oxidation is  
872 a larger aerosol source than currently understood, and if future NO<sub>x</sub> reductions lead to a  
873 stronger sensitivity in nighttime BVOC oxidation rates, regional SOA loadings could decrease by  
874 a substantial fraction from the typical regional summertime OA loadings of 5 +/- 3 μg m<sup>-3</sup> (Saha  
875 et al., 2017).

876  
877 Analysis of the degree of oxidation and chemical composition of NO<sub>3</sub> + isoprene SOA would  
878 help to elucidate mechanistic reasons for the different field and lab SOA yields. For example,  
879 the potential contribution of the uptake of morning-after OH + NISOPOOH produced epoxides,  
880 discussed above in section 4.3, onto existing (acidic) aerosol could be quantified by  
881 measurement of these intermediates or their products in the aerosol phase. Assessment of  
882 degree of oxidation could help determine whether auto-oxidation mechanisms are active. Future

883 similar field studies would benefit from the co-deployment of the complementary tool of a  
884 Chemical Ionization Mass Spectrometer (CIMS) to detect NO<sub>3</sub> + isoprene products such as  
885 organic nitrates (Slade et al., 2017; Lee et al., 2016). Because of the potentially large effect on  
886 predicted SOA loading in regions of high isoprene emissions, a better mechanistic  
887 understanding of these observed yields is crucial.

888

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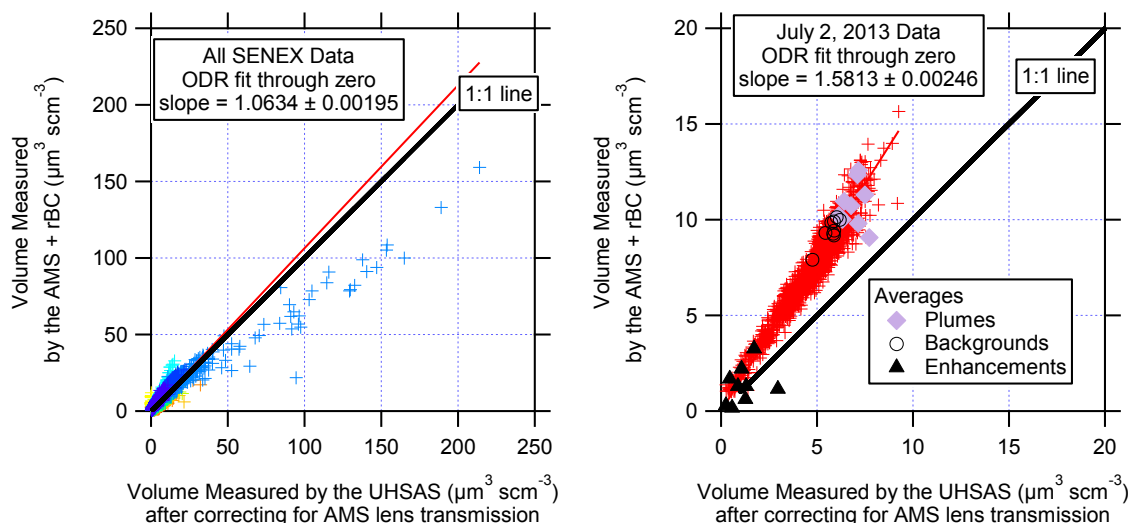
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1341 **Supplemental Information**

1342 In the main text, we noted a discrepancy between overall average aerosol volume estimates  
1343 based on size measurements vs. AMS for the flight analyzed here (see Figure S1). We checked  
1344 to see if this bias was also present in the individual plumes studied here by calculating the  
1345 volume changes from the sizing instruments and the derived volume changes from the  
1346 AMS+rBC mass. There is quite a bit of scatter in the volume enhancements, with with some of  
1347 the points falling along the same line as the data for this flight and some falling significantly  
1348 below the line. It is unclear why the two types of volume measurements disagree more for this  
1349 flight. Therefore, the bias in volume changes introduces additional uncertainty in the magnitude  
1350 of the plume enhancements, which is not included in the uncertainty propagation.  
1351



1352 **Figure S1.** Aerosol volume measured using the total aerosol mass from the AMS plus refractory  
1353 black carbon (rBC) and mass-weighted densities versus the aerosol volume measured by  
1354 optical size with the UHSAS after correcting for AMS lens transmission. The procedure for  
1355 calculating the mass-weighted density is described by Bahreini et al. (2009). On average, the  
1356 measured aerosol volume from composition is roughly equal to the measured aerosol volume  
1357 from size for the entire SENEX study (left hand panel) and is higher than one for the flight  
1358 analyzed here (July 2, 2013, right hand panel).  
1359  
1360

1361 **Corrections for AMS UMR nitrate data and applicability to pRONO<sub>2</sub> estimation**

1362 Nitrate in the AMS is quantified in unit mass resolution mode (UMR) as the sum of the estimated  
1363 NO<sup>+</sup> at *m/z* 30 and NO<sub>2</sub><sup>+</sup> at *m/z* 46, with a correction factor to account for the smaller ions (N<sup>+</sup>  
1364 and HNO<sub>3</sub><sup>+</sup>, mostly) produced from nitrate (Allan et al., 2004b). The default AMS UMR  
1365 quantification algorithm (documented in the AMS “fragmentation table”) estimates NO<sup>+</sup> as the  
1366 total signal at *m/z* 30 minus a small (2.2% of OA at *m/z* 29, “Org29” in AMS parlance)  
1367 subtraction to account for organic interferences and an isotopic correction for naturally-occurring  
1368 <sup>15</sup>N<sub>2</sub> from nitrogen in air. The default UMR fragmentation table was developed for mixed ambient  
1369 aerosols, in particular in urban studies, and it is the responsibility of each AMS user to correct it  
1370 as needed for each study. In environments with high biogenic contributions to total OA, and/or  
1371



1372 low total nitrate concentrations, the contribution of the  $\text{CH}_2\text{O}^+$  ion can be much larger than the  
1373 default subtraction at  $m/z$  30. Similarly, the  $\text{CH}_2\text{O}_2^+$  ion at  $m/z$  46 becomes non-negligible, and  
1374 hence nitrate reported from AMS data with UMR resolution will frequently be overestimated in  
1375 these situations. The poor performance of the default AMS correction is likely due to the initial  
1376 focus on urban OA with high nitrate fractions when deriving those corrections (Allan et al.,  
1377 2004b;Zhang et al., 2004).

1378  
1379 Here we derive a set of corrections based on an aircraft high-resolution (HR) dataset acquired  
1380 with the University of Colorado HR-AMS (Dunlea et al., 2009) on the NASA DC-8 during the  
1381 SEAC<sup>4</sup>RS campaign (Toon et al., 2016). SEAC<sup>4</sup>RS took place with a strong emphasis on the  
1382 SEUS 6 weeks after the SENEX flight analyzed in this manuscript. Based on an initial screening  
1383 of the correlations of the  $\text{CH}_2\text{O}^+$  and  $\text{CH}_2\text{O}_2^+$  ions with UMR signals, 10 potential UMR  $m/z$   
1384 between  $m/z$  29 and  $m/z$  53 were selected as viable for deriving suitable corrections. Further  
1385 analysis using three specific SEAC<sup>4</sup>RS flights (RF11 on 30 Aug 30<sup>th</sup>, 2013, RF16 on Sep 11<sup>th</sup>,  
1386 2013 and RF18 on Sep 16<sup>th</sup>, 2013) that covered a wide range of OA composition with both  
1387 strong biogenic contributions and fresh and aged biomass plumes showed that only four  $m/z$   
1388 (29, 42, 43 and 45) had good enough S/N and robust enough correlations to be used as  
1389 corrections. Table S1 summarizes the correction coefficients obtained in this analysis, and  
1390 Figure S2 shows the ability of matching the actual  $\text{NO}^+$  and  $\text{NO}_2^+$  signals (as obtained from  
1391 high-resolution analysis of these flights) with the corrected UMR procedure. These corrections  
1392 are applied as:

$$\text{UMR NO} = \text{Signal}(m/z30) - a_i \cdot \text{Signal}(\text{Variable}_i)$$

$$\text{UMR NO}_2 = \text{Signal}(m/z 46) - b_i \cdot \text{Signal}(\text{Variable}_i)$$

1393  
1394  
1395  
1396  
1397 with the coefficients  $a_i$  and  $b_i$  as reported in Table S1. It should be noted that in all cases the  
1398 contributions of  $\text{C}^{18}\text{O}^+$  to  $m/z$  30 need to be subtracted first before applying the correction (which  
1399 is constrained to the organic  $\text{CO}_2^+$  signal, measured at  $m/z$  44, by the naturally-occurring  
1400 isotopic ratio and assuming that OA produces  $\text{CO}^+ = \text{CO}_2^+$  (Zhang et al., 2005;Takegawa et al.,  
1401 2007). Likewise, the contribution of  $^{13}\text{CO}^+$  to Org29 needs to be subtracted first. It is hence very  
1402 important for this analysis that the corrections to the AMS frag table to suitably estimate the  
1403 contribution of gas phase  $\text{CO}_2^+$  to total UMR  $m/z$  44 as well as the baseline correction for  $m/z$  29  
1404 be properly applied first (Allan et al., 2004b). Finally, also note that the corrections using  $m/z$  29  
1405 and 43 are rather based on Org29 and Org43, which are standard AMS products that take the  
1406 OA relative ionization efficiency (RIE) into account.

1407  
1408 For the SEAC<sup>4</sup>RS dataset, the corrections amounted to on average subtracting 55% from UMR  
1409  $m/z$  30 and 33% from UMR  $m/z$  46. Despite this large subtraction, the corrected data correlates  
1410 very well with the HR AMS results, with less than 5% deviation in the regression slope between  
1411 the two datasets.

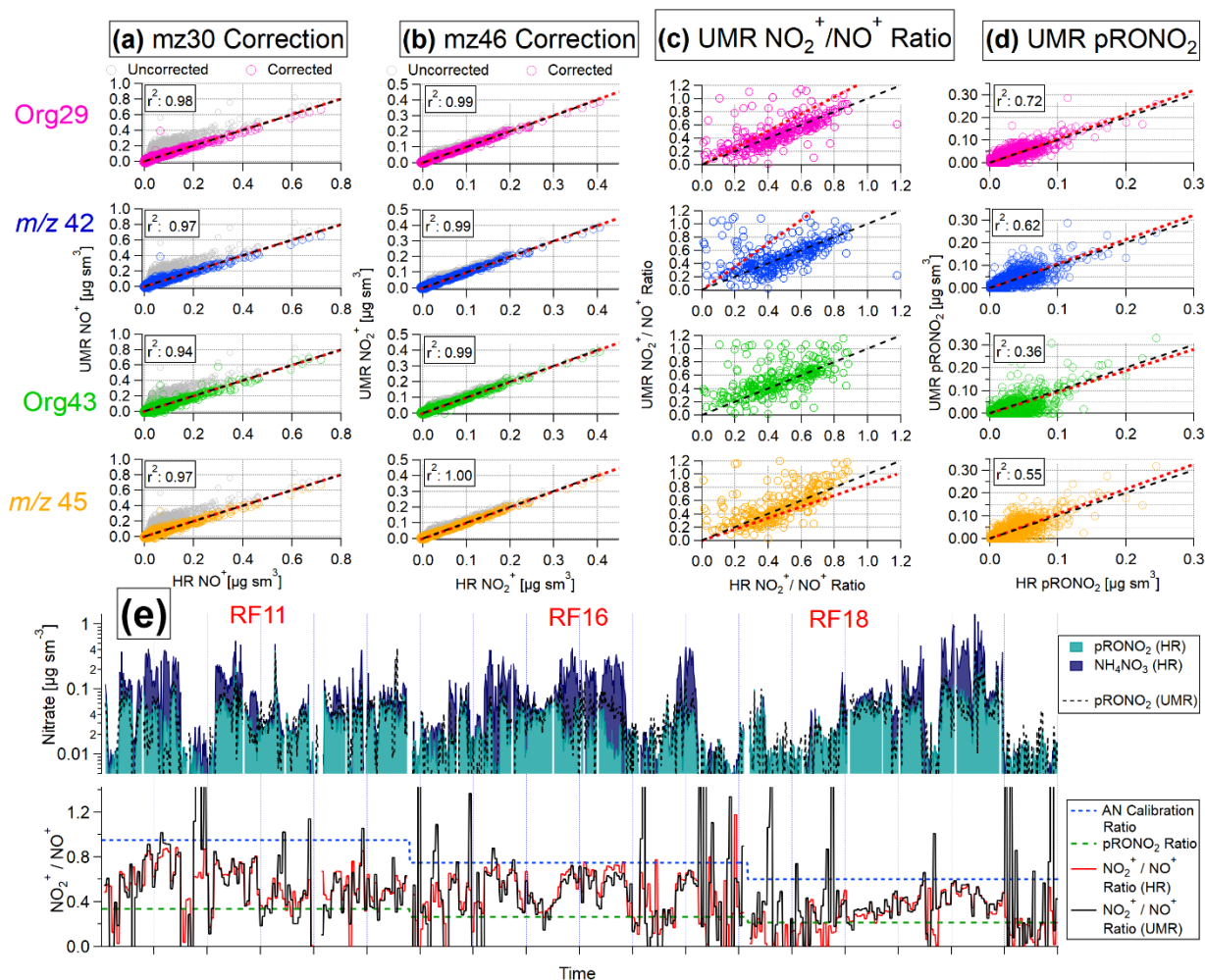
1412  
1413 Although all of the corrections in Table S1 were valid for the SEAC<sup>4</sup>RS data set, for the flight  
1414 analyzed here we chose Org29 to correct  $m/z$  30 and  $mz$  45 correction to correct  $m/z$  46  
1415 because they were the closest organic signals to the UMR nitrate peaks with organic

1416 interferences and may be more valid for other field studies where different types of OA are  
1417 sampled. After these UMR signals were corrected and the appropriate RIEs and CE were  
1418 applied, the nitrate mass concentrations in the final data archive for the flight analyzed here  
1419 were reduced by 0-0.24  $\mu\text{g sm}^{-3}$ , averaging 0.11  $\mu\text{g sm}^{-3}$  or 32%. The corresponding increase in  
1420 OA due to the organic interferences in the UMR nitrate had linear dependence on the reported  
1421 OA mass concentrations ( $r^2 = 0.89$ ) with a slope of 1.3%.

1422  
1423 To estimate the fraction of nitrate that is organic nitrate (pRONO<sub>2</sub>) the use of the NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> ratio  
1424 with an empirically determined pRONO<sub>2</sub> calibration ratio has been successfully used previously  
1425 with HR-AMS data (Farmer et al., 2010; Fry et al., 2013; Ayres et al., 2015; Fisher et al.,  
1426 2016; Day et al., 2017; Palm et al., 2017; Lee et al., 2016). Figure S2 summarizes how well the  
1427 ratio of the corrected UMR *m/z* 30 and 46 signals correlate with the NO<sub>2</sub><sup>+</sup> and NO<sup>+</sup> (and ratios)  
1428 determined using HR data. As expected, there is considerable scatter at very low nitrate  
1429 concentrations (which is a considerable part of the dataset, as the time series shows, since the  
1430 free troposphere was sampled extensively). However, for the predicted pRONO<sub>2</sub> (which is  
1431 mass-weighted), most of this scatter disappears, and for concentrations above 0.1  $\mu\text{g sm}^{-3}$  of  
1432 nitrate there is good agreement between the HR results and the UMR-corrected pRONO<sub>2</sub>,  
1433 regardless of the correction chosen. For lower concentrations the scatter is considerable larger,  
1434 with the Org29 correction providing the best overall agreement. Based on the variability in this  
1435 dataset for this correction (Org29), we estimate the uncertainty in pRONO<sub>2</sub> fraction  
1436 apportionment using UMR to be about 30%, in addition to an estimated uncertainty for the  
1437 apportionment method using HR of 20%. From the comparison of UMR-corrected total nitrate  
1438 to HR nitrate (not shown), we estimate an additional error of 5% for total nitrate error using  
1439 these corrections.

1440  
1441 As mentioned in the main text, the empirically determined pRONO<sub>2</sub> calibration ratio used for the  
1442 flight data analyzed here was the ratio of NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> from the ammonium nitrate calibration  
1443 aerosols divided by 2.8. This factor was determined as the average of several literature studies  
1444 (Fry et al., 2009; Farmer et al., 2010; Rollins et al., 2009; Sato et al., 2010; Fry et al., 2011; Boyd et  
1445 al., 2015) and applied according to the “ratio of ratios” method (Fry et al., 2013). The ammonium  
1446 nitrate NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> ratio was obtained from the two calibrations on 30 June and 7 July that  
1447 bracketed the flight on 2 July, as described above. This ratio averaged 0.490. Hence, the  
1448 organic nitrate NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> ratio was estimated to be 0.175. The ratio of NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> from the flight  
1449 data was then used with the pRONO<sub>2</sub> and ammonium nitrate NO<sub>2</sub><sup>+</sup>/NO<sup>+</sup> calibration ratios to  
1450 estimate the fraction of the total corrected nitrate mass concentrations that was organic  
1451 (pRONO<sub>2</sub>) or inorganic (nitrate associated with ammonium or NH<sub>4</sub>NO<sub>3</sub>). Propagating the 30%  
1452 UMR vs HR uncertainty and 20% apportionment (see above) error on top of the 34% AMS total  
1453 nitrate measurement uncertainty results in  $\pm 50\%$  uncertainties in the derived organic nitrate  
1454 mass concentrations (and similar for NH<sub>4</sub>NO<sub>3</sub>; however it will depend on the relative  
1455 contributions of pRONO<sub>2</sub> and NH<sub>4</sub>NO<sub>3</sub> to total nitrate since the absolute concentration errors  
1456 associated with pRONO<sub>2</sub> - NH<sub>4</sub>NO<sub>3</sub> apportionment should be similar [64]).

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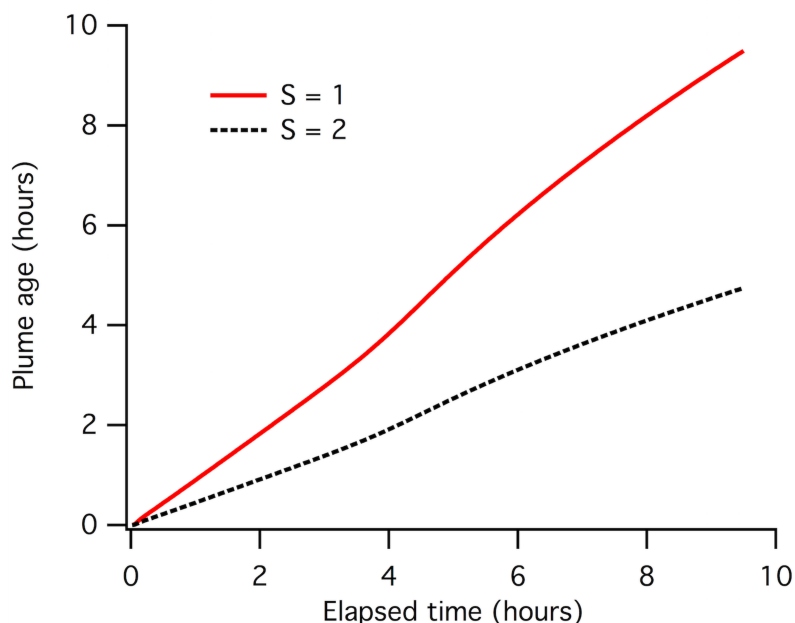


1458  
 1459 **Figure S2.** (a and b) Comparison of  $m/z$  30 and 46 with the  $\text{NO}^+$  and  $\text{NO}_2^+$  signals from the high  
 1460 resolution analysis of the AMS data before and after applying the four different corrections listed  
 1461 in Table S1. The Pearson  $r^2$  for the corrected dataset is shown as well. (c) Comparison of the  
 1462  $\text{NO}_2^+/\text{NO}^+$  ratio obtained from HR analysis with the ratios of the corrected UMR NO and  $\text{NO}_2$   
 1463 variables (d) Comparison of the  $\text{pRONO}_2$  concentrations derived using the HR and UMR  $\text{NO}_2^+/\text{NO}^+$   
 1464 ratios. (e) Time series of the total and speciated nitrate as reported from HR analysis of the  
 1465 SEAC<sup>4</sup>RS data (DOI: 10.5067/Aircraft/SEAC4RS/Aerosol-TraceGas-Cloud) compared to the  
 1466 speciation using the Org29 correction (note the logarithmic scale). The bottom time series  
 1467 shows the  $\text{NO}_2^+/\text{NO}^+$  ratio that the speciation is based on, again for the HR and corrected UMR  
 1468 case.

1469 **Table S1.** Coefficients used to correct  $m/z$  30 and 46 to estimate total nitrate.

AMS Variable	Correction coefficient for $m/z$ 30 (a <sub>i</sub> )	Correction coefficient for $m/z$ 46 (b <sub>i</sub> )
Org29	0.215	0.037
$m/z$ 42	0.51	0.092
Org43	0.215	0.037
$m/z$ 45	0.72	0.127

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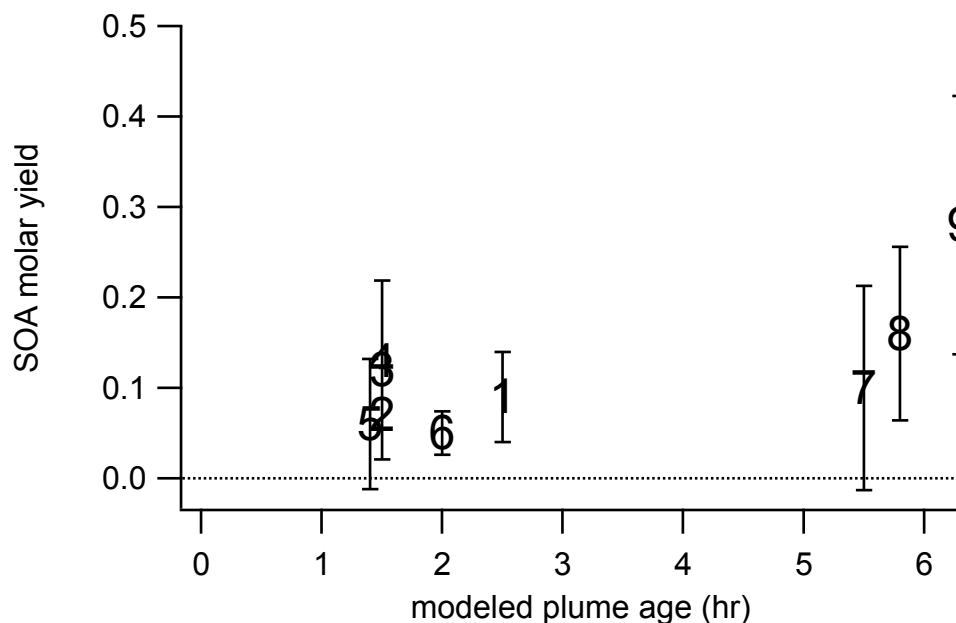
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**Figure S3.** Calculated plume age vs. elapsed time in a box model run for a single representative night. Plume ages on the y-axis are calculated based on Equation 1 in the main text but using model  $\text{NO}_2$  and  $\text{O}_3$  data. Time since sunset on the x-axis is the model elapsed time (i.e., run time of the model during darkness).



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**Figure S4.** SOA molar yield is positively correlated with estimated plume age. This SOA molar yield is based on Eq. 3, with error bars determined by propagation of observed variability in  $\text{pRONO}_2$  and isoprene, where multiple point averaging was possible. Markers correspond to

1482 plume numbers. ). Based on the box model described in more detail below, the first-generation  
1483 isoprene products peak at a approximately 4 hours plume age and then begin to decay.

1484

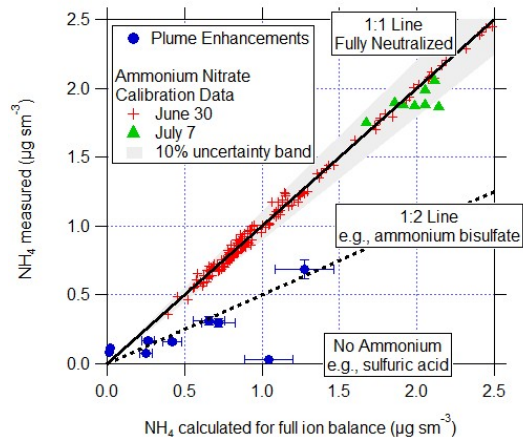
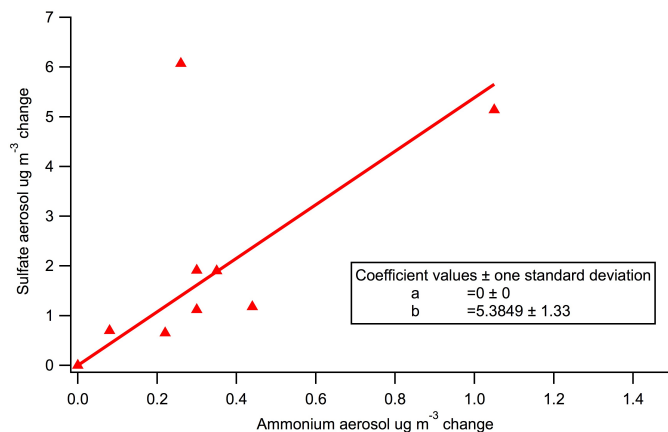
1485 **Table S2.** Peak ambient (wet) aerosol surface area during each plume used in the yield  
1486 analysis (plume numbers 1 – 9), and for the two longer urban plumes transected at the end of  
1487 the flight.

plume number	7/2/13 plume time (UTC)	Peak aerosol surface area ( $\mu\text{m}^2 \text{cm}^{-3}$ )
1	2:18	280
2	2:20	370
3	2:21	470
4	3:03	340
5	3:55	800
6	4:34	470
7	4:37	370
8	4:39	420
9	5:04	490
Urban plume	5:36	340
Urban plume	6:37	300

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**Figure S5.** (a) In-plume change in sulfate mass concentration vs. change in ammonium aerosol mass concentration is generally well correlated, with a slope of 5.4. The masses of the cations and anions would give an ion balance for pure  $(\text{NH}_4)_2\text{SO}_4$  of  $\text{MW}(\text{SO}_4)/(2 \times \text{MW}(\text{NH}_4)) = 2.7$ , and for  $(\text{NH}_4)\text{HSO}_4$  of  $\text{MW}(\text{SO}_4)/(\text{MW}(\text{NH}_4)) = 5.4$ . Hence, this slope provides support for a mix of these two ammonium sulfate salts, with sometimes exclusively  $(\text{NH}_4)\text{HSO}_4$ . This is consistent with incomplete neutralization of the sulfate mass by ammonium. The one clear outlier (sulfate increase of  $6 \mu\text{g m}^{-3}$  for Plume #5) suggests excess sulfate, rendering ammonium or other inorganic nitrate formation even less likely. Points with ammonium aerosol below  $0.1 \mu\text{g m}^{-3}$  are within the variability of that measurement; their omission does not change the slope. (b) Measured vs. calculated (ion balanced)  $\text{NH}_4$  for calibration data and plume enhancements. This also shows that plumes are acidic than ammonium sulfate, ruling out the possibility of inorganic nitrate formation.

1508 **Additional AMS and auxiliary data from plumes**

1509

1510 **Table S3.** Additional information for the list of plumes used in this NO<sub>3</sub> + isoprene SOA yield  
 1511 analysis, for which key yield-related data is presented in Table 1. For each plume, the delta-  
 1512 values listed indicate the difference between in-plume and outside-plume background in  
 1513 average observed concentration. After each plume number, the numbers of points averaged for  
 1514 isoprene and AMS, respectively, are listed. Plume numbers annotated with \* indicate brief  
 1515 plumes for which only single-point measurements of in-plume aerosol composition were  
 1516 possible. Also shown are the plume changes in isoprene used in the present analysis ( $\Delta$ isop,  
 1517 the difference between in-plume and background isoprene concentration, reproduced from  
 1518 Table 1), alongside for comparison the  $\Delta$ isop determined as the difference between in-plume  
 1519 isoprene and the modeled sunset (initial) concentration of isoprene present at that location  
 1520 outside of the plume, determined using an iterative box model (Edwards et al., 2017). The  
 1521 similarity between these two values for most points suggests that the isoprene just outside of  
 1522 each plume transect was largely unperturbed from the sunset initial value.

plume number [#isop/#AMS]	7/2/13 plume time (UTC)	$\Delta$ ORG <sub>aero</sub> ( $\mu\text{g m}^{-3}$ )	$\Delta$ NH <sub>4,aero</sub> ( $\mu\text{g m}^{-3}$ )	$\Delta$ SO <sub>4,aero</sub> ( $\mu\text{g m}^{-3}$ )	Temp (C)	%RH	$\Delta$ isop (pptv)	$\Delta$ isop from model (pptv)	Isop:M1 Mole Rat
Typical variability ( $\mu\text{g m}^{-3}$ ):		0.75	0.1	0.5					
1 [2/3]	2:18	0.35	0	0	23.6	66.5	-335	-327	36.5
2 [*]	2:20	0.89	0.3	1.91	23.6	65	-404	-453	71.4
3 [4/5]	2:21	1.25	1.05	5.14	23.6	65.2	-228	-337	16.6
4 [*]	3:03	0.16	0.08	0.7	21.2	68.1	-453	-391	50.6
5 [3/4]	3:55	0.32	0.26	6.07	21.9	65.5	-255	-376	34.2
6 [2/2]	4:34	0.57	0.3	1.12	19.9	74.6	-713	-233	17.3
7 [5/6]	4:37	1.05	0.22	0.65	19.7	76.2	-298	-221	14.2
8 [2/3]	4:39	1.26	0.44	1.18	18.3	82.2	-443	-353	11.0
9 [7/8]	5:04	1.45	0.35	1.9	17.2	84.8	-293	-434	17.8

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1526 **Box model calculations**

1527 Box model simulations were performed using the Dynamically Simple Model of Atmospheric  
 1528 Chemical Complexity (DSMACC, [http://wiki.seas.harvard.edu/geos-](http://wiki.seas.harvard.edu/geos-chem/index.php/DSMACC_chemical_box_model)  
 1529 [chem/index.php/DSMACC\\_chemical\\_box\\_model](http://wiki.seas.harvard.edu/geos-chem/index.php/DSMACC_chemical_box_model)), containing the Master Chemical Mechanism  
 1530 v3.3.1 chemistry scheme (<http://mcm.leeds.ac.uk/MCM/>). The model approach is similar to that  
 1531 described in detail in Edwards et al. 2017, and the accompanying supplement, with the model  
 1532 run over a 9.5 hour night to simulate the nocturnal residual layer. For the nocturnal simulation  
 1533 used in this work (for both the plume lifetime calculation and the peroxy radical lifetime analysis  
 1534 in Sect. 4.3) the model was initialized with concentrations of the constraining species  
 1535 representative of the SENEX observations (Table S4). As the model is simulating power plant  
 1536 plume evolution from point of emission, a starting NO mixing ratio of 10 ppb was used to  
 1537 constrain NO<sub>x</sub>, and the chemistry scheme was subsequently allowed to partition the reactive  
 1538 nitrogen. The top panels in Figure S7 show the evolution of key species during this nocturnal  
 1539 simulation.

1540 **Table S4:** Species constrained (MCM v3.3.1 names) during model simulations and constraining  
 1541 values. Constraint column indicates if species concentrations were held at the constrained value  
 1542 throughout the simulation (Fixed) or allowed to vary after initialization (Initial).

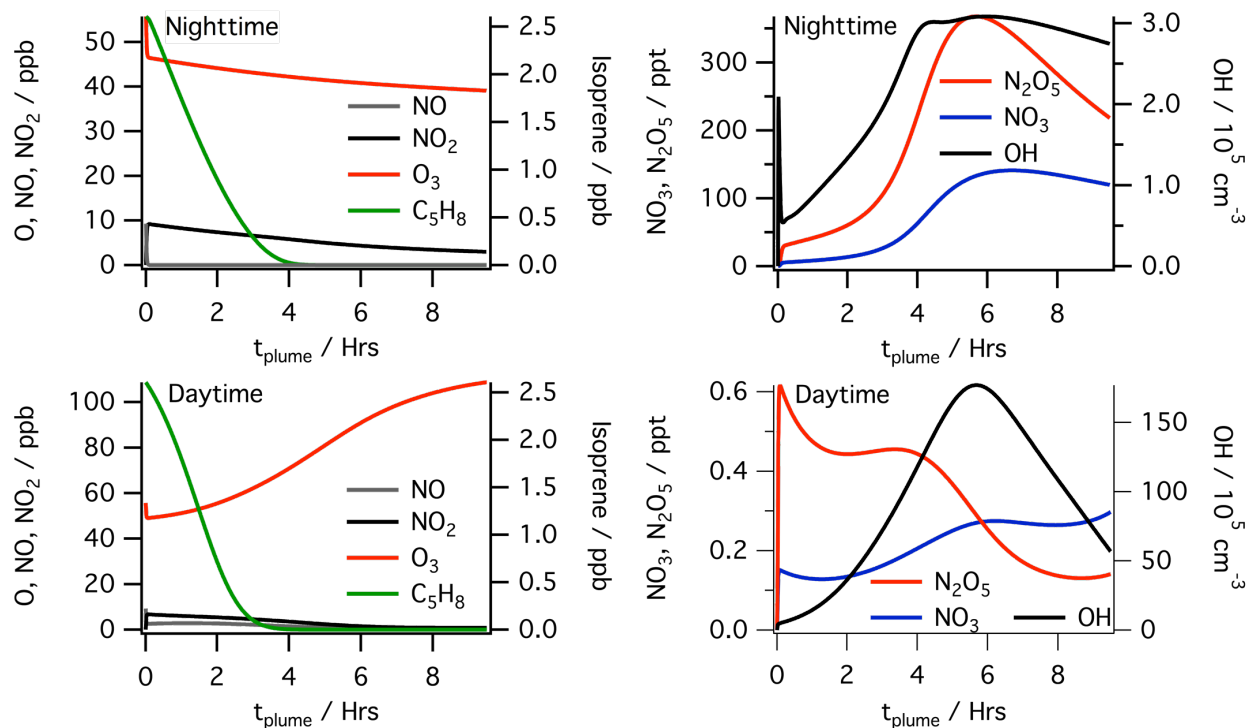
Species	Mixing ratio	Units	Constraint
NO	9.28	ppb	Initial
O3	55.72	ppb	Initial
CO	134.00	ppb	Fixed
CH4	1920.00	ppb	Fixed
C5H8	2606.80	ppt	Initial
APINENE	38.87	ppt	Initial
BPINENE	195.50	ppt	Initial
LIMONENE	12.42	ppt	Initial
MACR	454.13	ppt	Initial
MVK	1006.00	ppt	Initial
IC4H10	47.00	ppt	Fixed
NC4H10	128.00	ppt	Fixed
C2H6	1199.00	ppt	Fixed
C2H4	117.00	ppt	Fixed
C2H2	145.00	ppt	Fixed
NC6H14	20.00	ppt	Fixed
IC5H12	120.00	ppt	Fixed
NC5H12	76.00	ppt	Fixed
C3H8	344.00	ppt	Fixed
C3H6	26.00	ppt	Fixed
CH3COCH3	2556.00	ppt	Fixed
BENZENE	35.90	ppt	Fixed
C2H5OH	2239.00	ppt	Fixed
MEK	309.00	ppt	Fixed
CH3OH	5560.00	ppt	Fixed

1543 The daytime simulation used for comparison in Sect. 4.3 of the main manuscript (lower panels  
 1544 of Figure S7) uses the same initialization as the nocturnal simulation; with the only difference  
 1545 being the model is run during the daytime. Photolysis rates are calculated using TUV  
 1546 (<https://www2.acom.ucar.edu/modeling/tropospheric-ultraviolet-and-visible-tuv-radiation-model>).  
 1547 The daytime simulation does not accurately simulate daytime mixing ratios of species such as  
 1548 O<sub>3</sub> representative of SENEX observations. However, the intent of this simulation is to compare  
 1549 model daytime peroxy radical fate and lifetime with the nocturnal simulation. The presence of



1550 intense convective mixing in the daytime planetary boundary layer of the Southeast US makes  
1551 accurately modeling these concentrations difficult with a zero dimensional model.

1552



1553

1554 **Fig. S6.** Model calculated NO, NO<sub>2</sub>, O<sub>3</sub>, and isoprene (left) and NO<sub>3</sub>, N<sub>2</sub>O<sub>5</sub> and OH (right for the  
1555 nocturnal (top) and daytime (bottom) simulations shown in Sect. 4.3.

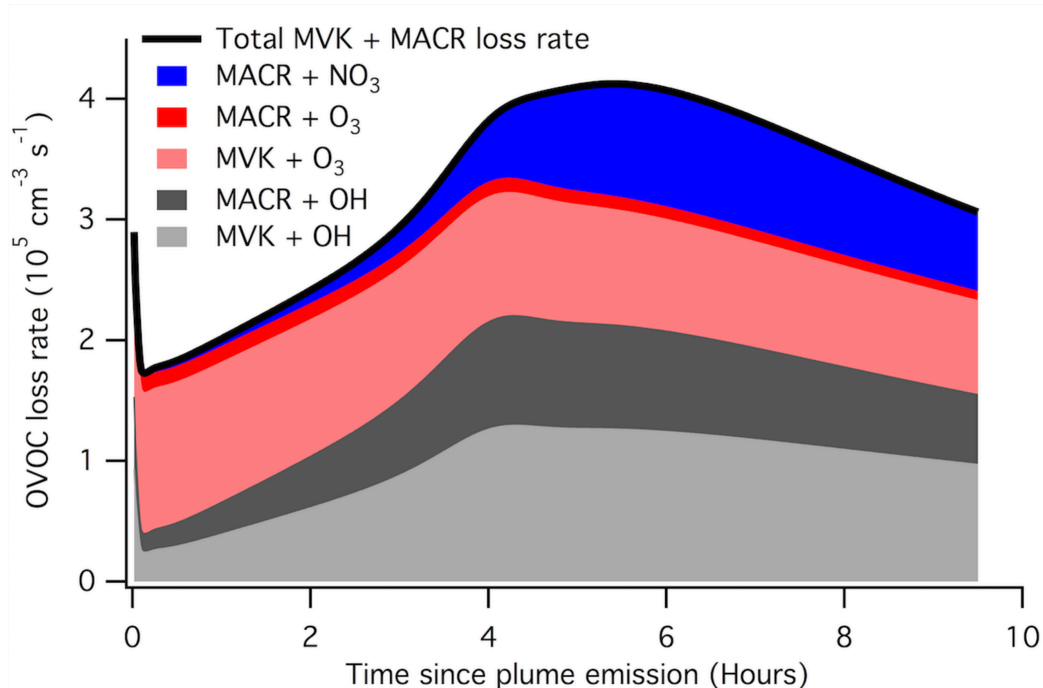
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### 1557 **Additional considerations investigated via RO<sub>2</sub> fate box modeling**

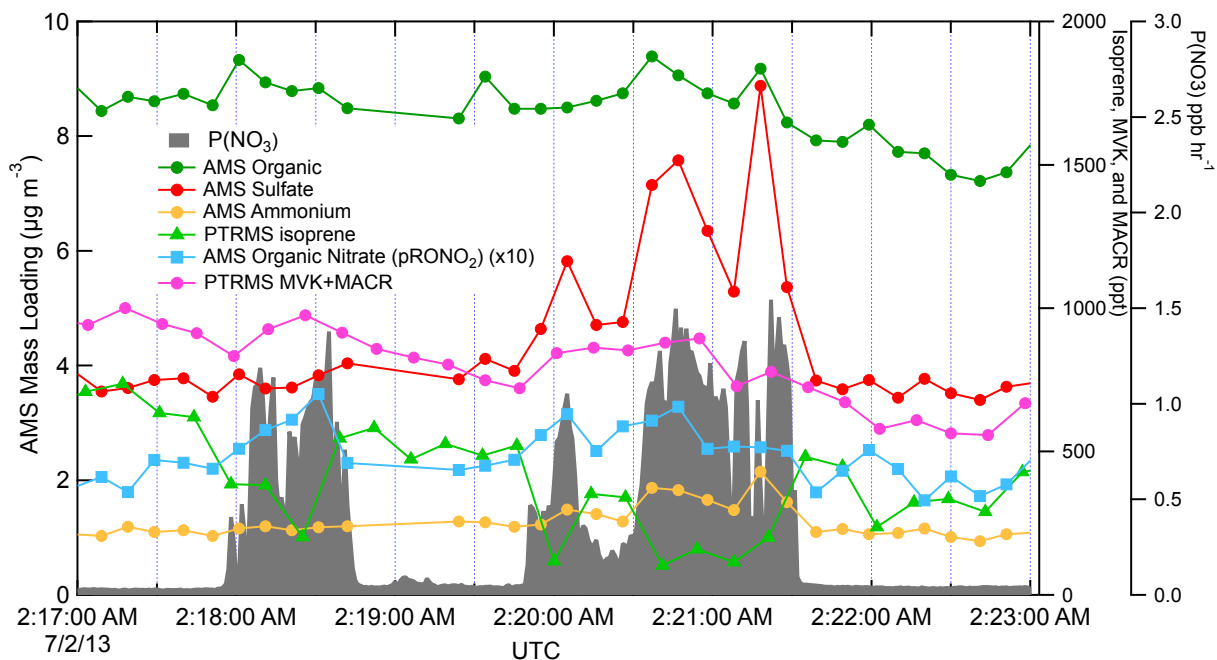
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1559 Based on the potentially larger than previously estimated contribution of RO<sub>2</sub>+RO<sub>2</sub> reactions at  
1560 night, we considered a related possible source of a high bias in the determined SOA yields. If  
1561 NO<sub>3</sub> reaction with the major daytime isoprene oxidation products MVK and/or MACR produces  
1562 RO<sub>2</sub> radicals that can cross-react with NO<sub>3</sub> + isoprene products to produce condensable  
1563 products, this would be a mechanism of recruiting isoprene-derived organic mass into the  
1564 aerosol, but that original isoprene oxidation would not be counted in the denominator of the yield  
1565 calculation, since its interaction with NO<sub>3</sub> began as MACR or MVK. In the box model, substantial  
1566 MVK and MACR are available in the plume at nighttime, but only MACR reacts with NO<sub>3</sub>, and a  
1567 maximum fraction of one-quarter of MVK+MACR losses go to reaction with NO<sub>3</sub> overnight (see  
1568 Figure S8). In addition, in our power plant plume observations, MVK+MACR are not observed to  
1569 be appreciably depleted by the large NO<sub>3</sub> injection, further suggesting that this chemistry is not  
1570 a substantial additional source of SOA (see Figure S9).

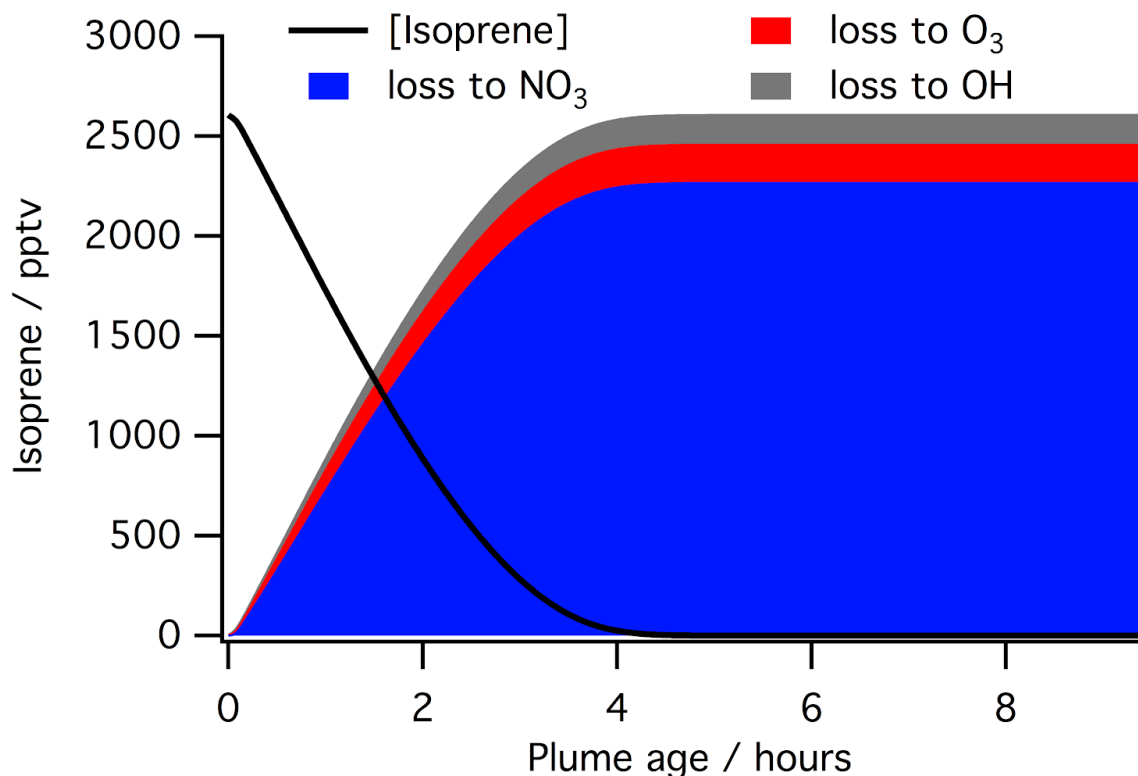
1571



1572  
 1573 **Figure S7.** Calculated (via MCM) loss rate contributions for the daytime isoprene products  
 1574 methyl vinyl ketone (MVK) and methacrolein (MACR) in the simulated nighttime plume used in  
 1575 the text. Only MACR reacts with  $\text{NO}_3$ , and the contribution of this process to total losses (green  
 1576 stack) is relatively minor.



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 1578 **Figure S8.** MVK and MACR are not titrated on the timescale of these yield estimates in power  
 1579 plant plumes.  
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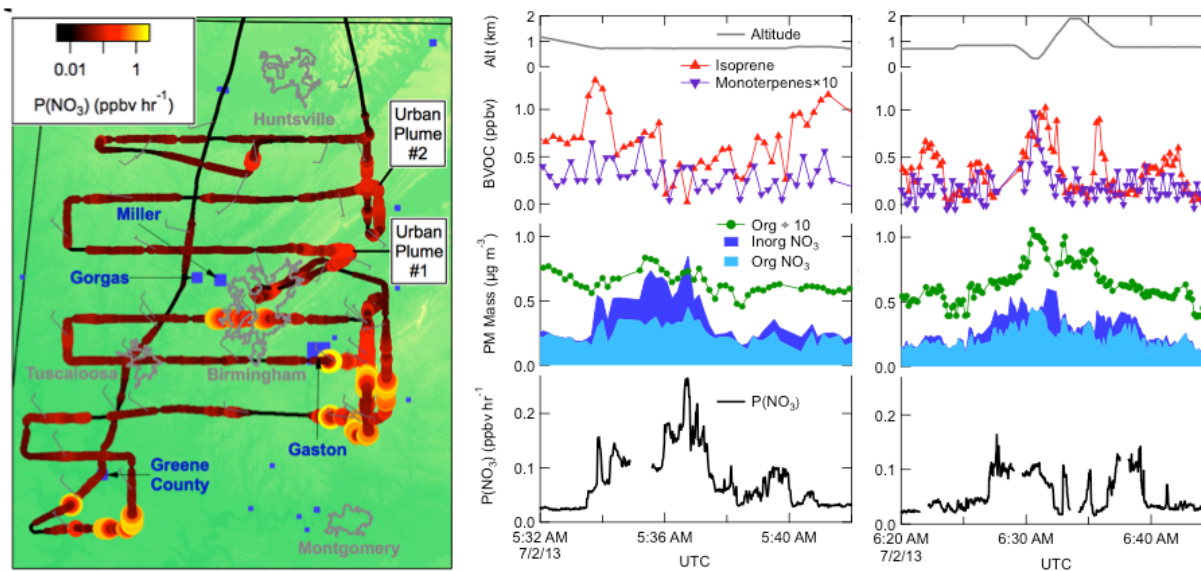
**Figure S9.** Model simulation of typical in-plume consumption of isoprene (black line), and stacked plot showing the contributions to this from the NO<sub>3</sub>, O<sub>3</sub>, and OH. Modeled plume was emitted at sunset, so this represents nocturnal processing under power plant plume conditions.

## 1587 Two urban plume case studies

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1589 In addition to the nine power plant plumes analyzed above to determine the NO<sub>3</sub> + isoprene  
1590 SOA molar yield, towards the end of the July 2 flight, the Birmingham urban plume was  
1591 intercepted twice (around 5:36 am and 6:37 am UTC, Fig. 8). These downwind urban plumes  
1592 are among the most aged plumes (estimated at 5.2 and 5.8 hours, respectively), but are also  
1593 substantially more diffuse than the narrow power plant plume intercepts and have lower peak  
1594  $P(\text{NO}_3)$ . Nevertheless, we note that these two plumes contain periods of apparent anti-  
1595 correlation of isoprene and organic nitrate aerosol time series and high apparent SOA molar  
1596 yields (23%, 19%) and mass yields (62%, 51%), if calculated by the same method as above and  
1597 omitting the period of vertical profiling in the second plume. Potentially complicating these urban  
1598 SOA yield determinations is the fact that the inorganic fraction of nitrate was much larger than in  
1599 the power plant plumes (see Fig. 8). The background isoprene is also somewhat lower in these  
1600 urban plumes, potentially shifting the NO<sub>3</sub>/N<sub>2</sub>O<sub>5</sub> fate to reactions other than NO<sub>3</sub> + isoprene (see  
1601 Fig. S4 in Edwards et al. (Edwards et al., 2017)). The aerosol surface area is not noticeably  
1602 higher in these urban plumes, which one might expect to lead to a larger contribution of N<sub>2</sub>O<sub>5</sub>  
1603 uptake and hydrolysis. In the more complex mix of gases characteristic of an urban plume, we  
1604 hesitate to attribute these apparent yields exclusively to the NO<sub>3</sub> + isoprene reaction.

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**Figure S10.** Flight map and time series of two urban plume intercepts, showing anticorrelation of organic nitrate and isoprene. These more diffuse plumes, with lower  $P(\text{NO}_3)$  and larger inorganic nitrate contribution, make yield determination more uncertain, so we do not include them in the overall yield determination. However, using the same methodology as for the power plant plumes would give similarly high yields for these very aged plumes.