

1 **In situ measurements and modeling of reactive trace gases**  
2 **in a small biomass burning plume**

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21

22 **Abstract**

23 An instrumented NASA P-3B aircraft was used for airborne sampling of trace gases in a  
24 plume that had emanated from a small forest understory fire in Georgia, USA. The plume was  
25 sampled at its origin for deriving emission factors and followed ~ 13.6 km downwind for  
26 observing chemical changes during the first hour of atmospheric aging. The P-3B payload  
27 included a proton-transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS), which  
28 measured non-methane organic gases (NMOGs) at unprecedented spatio-temporal resolution  
29 (10m/0.1s). Quantitative emission data are reported for CO<sub>2</sub>, CO, NO, NO<sub>2</sub>, HONO, NH<sub>3</sub> and  
30 16 NMOGs (formaldehyde, methanol, acetonitrile, propene, acetaldehyde, formic acid,  
31 acetone plus its isomer propanal, acetic acid plus its isomer glycolaldehyde, furan, isoprene  
32 plus isomeric pentadienes and cyclopentene, methyl vinyl ketone plus its isomers  
33 crotonaldehyde and methacrolein, methylglyoxal, hydroxy acetone plus its isomers methyl  
34 acetate and propionic acid, benzene, 2,3-butanedione and 2-furfural) with molar emission ratios  
35 relative to CO larger than 1 ppbV ppmV<sup>-1</sup>. Formaldehyde, acetaldehyde, 2-furfural and  
36 methanol dominated NMOG emissions. No NMOGs with more than 10 carbon atoms were  
37 observed at mixing ratios larger than 50 pptV per ppmV CO emitted. Downwind plume  
38 chemistry was investigated using the observations and a 0-D photochemical box model  
39 simulation. The model was run on a near-explicit chemical mechanism (MCM v3.3) and  
40 initialized with measured emission data. Ozone formation during the first hour of atmospheric  
41 aging was well captured by the model, with carbonyls (formaldehyde, acetaldehyde, 2,3-  
42 butanedione, methylglyoxal, 2-furfural) in addition to CO and CH<sub>4</sub> being the main drivers of  
43 peroxy radical chemistry. The model also accurately reproduced the sequestration of NO<sub>x</sub> into  
44 PAN and the OH-initiated degradation of furan and 2-furfural at an average OH concentration  
45 of  $7.45 \pm 1.07 \times 10^6 \text{ cm}^{-3}$  in the plume. Formaldehyde, acetone/propanal, acetic  
46 acid/glycolaldehyde and maleic acid/maleic anhydride (tentatively identified) were found to  
47 be the main NMOGs to increase during one hour of atmospheric plume processing, with the  
48 model being unable to capture the observed increase. A mass balance analysis suggests that  
49 about 50% of the aerosol mass formed in the downwind plume is organic in nature.

50

51

## 52 1 Introduction

53 Understanding and predicting the impacts of biomass burning emissions on air quality is a  
54 challenging but important task. Fire emissions include a plethora of inorganic and organic  
55 species, both in the gas and the particulate phase, and many of them undergo rapid chemical  
56 transformations and phase changes after their release to the atmosphere (e.g. Simoneit, 2002).  
57 These processes are the focus of intense research efforts, both in the laboratory and in the  
58 field. Over the last decade, many airborne field studies have been undertaken for  
59 characterizing emissions and evolution of gases and particles in the aging plume (e.g. Akagi et  
60 al., 2012, 2013; Yokelson et al., 2009). In general, these studies have targeted emissions from  
61 medium and large-scale fires. Small fires (< 500 m diameter of burned area) have been  
62 undersampled although they may contribute 35% or more to global biomass burning carbon  
63 emissions (Randerson et al., 2012). Emissions from small fires are often not included in  
64 emission inventories and local and regional air quality assessments seldom include emissions  
65 from small fires. In addition, the chemical complexity of emissions poses a major challenge to  
66 modeling efforts. Lumped mechanisms are thus typically used in chemical models to predict  
67 the evolution of trace gases in biomass burning plumes. Lumping of species may, however,  
68 result in an oversimplification of the involved chemistry, which will ultimately yield  
69 erroneous model predictions.

70 In this work, we present the results from an airborne study, in which inorganic and organic  
71 trace gases emanating from a small forest understory fire were measured with state-of-the-art  
72 analytical tools. A proton-transfer-reaction time-of-flight mass spectrometry (PTR-ToF-MS)  
73 instrument delivered non-methane organic gas (NMOG) data at unprecedented spatio-  
74 temporal resolution. We sampled the plume at its origin for deriving emission factors and  
75 followed it downwind for observing chemical changes during the first hour of atmospheric  
76 aging. We also found that a 0-D photochemical box model, run on a near-explicit chemical  
77 mechanism and properly initialized with the measured emission data, adequately described  
78 key chemical processes (ozone and radical formation, NO<sub>x</sub> sequestration) in the aging plume.  
79

80 **2 Methods**

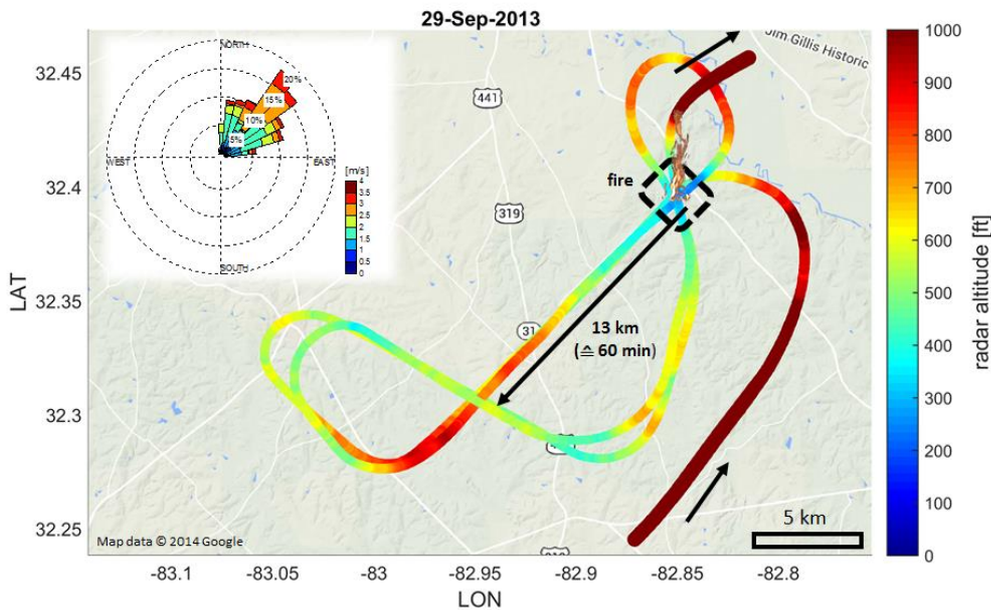
81 **2.1 Sampling strategy and conditions**

82 A small biomass burning plume was intercepted by the NASA P-3B research aircraft in  
83 Laurens County near Dublin, GA, USA on September 29<sup>th</sup>, 2013, during a flight from  
84 Houston, TX, to Wallops Island, VA. The plume emanated from a managed forest understory  
85 fire located at 32° 23' 42" N and 82° 51' 7.2" W which had been applied after logging and  
86 forest clearance activities. Historic Google Earth imagery shows that the area to the SW of the  
87 fire location had undergone intense forest clearing between 2011 and 2014. After the flight,  
88 the burned area was inspected by a local official who identified residual tree logs (pine, oak)  
89 and weeds as fire fuels. Figures 1a and 1b are two frames from the P-3B front camera  
90 showing the fire and the emanating plume at 17:33:32 UTC (UTC = local time + 4 hours) and  
91 17:42:51 UTC, respectively.



93 **Figure 1.** NASA P-3B front camera frames showing the forest understory fire and the  
94 emanating biomass burning plume at 17:33:32 (a) and UTC 17:42:51 (b), respectively.

95 Figure 2 depicts the P-3B flight pattern color-coded in radar altitude, with blue lowest and red  
96 highest. The flight direction is indicated by black arrows. Winds steadily blew from the NE at  
97 an average speed of  $3.5 \text{ ms}^{-1}$  (Figure 2, wind rose inset in the upper left corner). The average  
98 temperature during the sampling period (17:30-17:55 UTC) as measured by the P-3B met  
99 sensors was  $26.5 \pm 5.3 \text{ }^\circ\text{C}$  and the average relative humidity was  $60.4 \pm 2.3 \%$ . The average  
100 vertical temperature gradient was  $-1.34 \text{ }^\circ\text{C}/100 \text{ m}$ , causing the plume to slowly rise downwind  
101 of the source. The turbulence condition of the boundary layer was neutral to slightly unstable.



102

103 **Figure 2.** Flight pattern of the NASA P-3B to obtain four point source emission profiles, two  
 104 longitudinal plume transects (source to 1 hour downwind) and two transverse downwind  
 105 plume transects (1 hour downwind from source). The inset shows wind rose data obtained  
 106 during the two longitudinal plume transects when wind measurements are most accurate.

107 The fire was sighted and approached from the SW. Following a 180° turn, the aircraft  
 108 overflew the fire for the first time at 125 m altitude (Figure 1a) at 17:33:35 UTC (source  
 109 emission profile 1). The plume was then followed downwind in southwesterly direction for  
 110 approximately 2 minutes, slowly climbing in altitude to reach a radar altitude of 190 m at a  
 111 13.6 km downwind location (longitudinal plume transect 1). The underlying terrain was  
 112 forested and agricultural land. At an average wind speed of  $3.5 \text{ ms}^{-1}$ , the plume travel time for  
 113 a 13.6 km distance is approximately one hour. Following a horizontal loop maneuver, the ~ 8  
 114 km broad plume was sampled transversely at 160 m radar altitude at the 13.6 km downwind  
 115 location (transverse downwind plume transect 1). Subsequently, the P-3B returned to the fire  
 116 intercepting the freshly emitted plume at 17:42:57 UTC (Figure 1b) and at 17:45:38 UTC, at  
 117 110 m and 80 m altitude respectively (source emission profiles 2 and 3). The downwind  
 118 pattern was repeated with longitudinal plume transect 2 reaching 220 m altitude at the 13.6  
 119 km downwind location. The second transverse downwind plume transect was at 160 m  
 120 altitude at the 13.6 km downwind location. The fourth and final fire overflight was at 75 m

121 altitude at 17:54:25 UTC (source emission profile 4). By implementing this sampling strategy,  
 122 we obtained i) four source emission profiles within 21 minutes, ii) two longitudinal plume  
 123 transects (source to 1 hour downwind) and iii) four plume characterizations at 1 hour  
 124 downwind distance from source (two longitudinal “spot” samples and two “integrated” cross-  
 125 plume samples). The results (see Section 3) indicate near-stable source conditions during the  
 126 sampling period. This implies that the observed downwind differences were mostly due to  
 127 dilution and photochemistry.

## 128 2.2 Analytical instrumentation

129 The NASA P-3B was returning from a DISCOVER-AQ deployment ([http://discover-](http://discover-aq.larc.nasa.gov/)  
 130 [aq.larc.nasa.gov/](http://discover-aq.larc.nasa.gov/)) in Houston, which had it equipped with a payload for *in-situ* atmospheric  
 131 chemistry measurements. The data used in this study were obtained using the analytical  
 132 instruments listed in Table 1.

133 **Table 1.** Excerpt of the P-3B analytical chemistry payload.

Instrument acronym	Measurement principle	Analyte*	Accuracy	Reference
PTR-ToF-MS	chemical ionization	NMOGs, HONO, NH <sub>3</sub>	5-40%	Müller et al., 2014
NO <sub>x</sub> O <sub>3</sub>	chemiluminescence	NO	10 pptV + 10%	Ridley and Grahek, 1990
		NO <sub>2</sub>	20 pptV + 10%	
		NO <sub>v</sub>	50 pptV + 20%	
		O <sub>3</sub>	0.1 ppbV + 5%	
AVOCET	non-dispersive IR spectroscopy	CO <sub>2</sub>	0.25 ppmV	Vay et al., 2011
DACOM	differential absorption spectroscopy	CO	< 1 ppbV	Sachse et al., 1987
		CH <sub>4</sub>		
UHSAS	laser-based optical-scattering	sub- $\mu$ m particle size distribution	20%	Cai et al., 2008

134 \* Measurement frequency was 1 Hz for instruments except PTR-ToF-MS (10 Hz).

135 This work focuses on NMOGs as measured by the PTR-ToF-MS instrument described in  
 136 detail by Müller et al. (2014). The data presented herein were acquired at a frequency of 10  
 137 Hz, which makes the PTR-ToF-MS instrument ideally suited for airborne NMOG  
 138 measurements at high spatio-temporal resolution. However, only the elemental composition of  
 139 organic analytes can be determined and not their structure. In other words, the PTR-ToF-MS  
 140 instrument does not resolve isomeric NMOGs (e.g. acetic acid and glycolaldehyde). The PTR-  
 141 TOF Data Analyzer Toolbox (<https://sites.google.com/site/ptrtof/>) was used for data analysis  
 142 (Müller et al., 2013). Accurate *m/z* information, element restriction to C, H, N and O atoms

143 and isotopic pattern analyses were used to determine the elemental composition ( $C_wH_xN_yO_z$ )  
144 of detected analyte ions. It has been shown in previous work that accurate  $m/z$  information can  
145 be obtained even at a moderate mass resolution  $m/\Delta m$  in the range of 1000 to 1500 (Müller et  
146 al., 2011, 2014). The assignment of observed  $m/z$  signals to specific chemical compounds was  
147 based on the literature (see paragraph 3.1.2).

148 Methanol, acetonitrile, acetaldehyde, acetone, isoprene, methyl ethyl ketone, benzene,  
149 toluene, m-xylene, 1,3,5-trimethylbenzene and monoterpenes ( $\alpha$ -pinene) were calibrated  
150 externally using a dynamically diluted certified standard. The measurement accuracy is  $\pm 5\%$   
151 for pure hydrocarbons and  $\pm 10\%$  for oxygenates. Formic acid and acetic acid were calibrated  
152 ( $\pm 10\%$ ) in a post-campaign study using a liquid standard nebulization device (LCU, Ionicon  
153 Analytik, Austria). The protonated formaldehyde ion signal was cross-calibrated to  
154 formaldehyde data collected by a Difference Frequency Absorption Spectroscopy (DFGAS,  
155 Weibring et al. (2007)) instrument during the same flight and at the same humidity conditions.  
156 Although less accurate ( $\pm 10\%$ ), PTR-ToF-MS formaldehyde data were used instead of  
157 DFGAS observations because of a higher data density in the plume. Instrumental response  
158 factors to furan, methylglyoxal and 2-furfural were calculated from ion-molecule collision  
159 theory (Cappellin et al., 2012). The estimated measurement accuracy for these species is  
160  $\pm 25\%$ . Peroxyacetyl nitrate (PAN) was quantified ( $\pm 40\%$ ) using a calibration factor obtained  
161 in a previous study (unpublished data). All other organic signals were corrected for  
162 instrumental mass discrimination effects and converted to volume mixing ratios by using the  
163 acetone sensitivity as a proxy. Mixing ratios in acetone-equivalents are estimated to be  
164 accurate to within  $\pm 40\%$ . This is also the maximum error we must assume for the total  
165 NMOG mass calculated by summing all individual signals calibrated as specified above.

166 The PTR-ToF-MS instrument also detects a few inorganic gases, nitrous acid (HONO) and  
167 ammonia ( $NH_3$ ) being two prominent examples. Given the importance of HONO for fire  
168 plume photochemistry, we made an attempt to quantify HONO emissions. HONO dehydrates  
169 upon protonation forming  $NO^+$  ions, which are observed at  $m/z$  29.997. The excess  $NO^+$   
170 signal in the plume was assigned to HONO. The contribution from organic nitrites was  
171 assumed to be minor. A positive measurement artifact from  $NO_2$ -to-HONO conversion (1% of  
172  $NO_2$ ) on instrumental surfaces was subtracted. The instrumental response to HONO and  
173 HONO inlet artifacts have been characterized in previous studies (Metzger et al., 2008;

174 Wisthaler et al., 2003). Given that different inlet and drift tube configurations were used in  
175 those studies, the 1% NO<sub>2</sub>-to-HONO conversion efficiency is to be considered an upper limit  
176 estimate. Still, the NO<sub>2</sub>-artifact only accounts for 10.4% of the NO<sup>+</sup> signal measured at the  
177 source. The estimated accuracy of the reported HONO data is ±30%. NH<sub>3</sub> measurements  
178 suffered from a high intrinsic background signal generated in the ion source of the instrument.  
179 This deteriorated the detection limit to 12 ppbV for 1-Hz measurements.

### 180 **2.3 Data processing**

181 Volume mixing ratios (VMRs) were obtained as described in section 2.2. When referring to  
182 the VMR of a species X, the italic style, *X*, is used throughout this work.

183 Given that the P-3B spent about two seconds in the plume during fire overflights and that CO  
184 was only measured at 1 Hz, it was not possible to perform linear regression analyses, *X* vs.  
185 *CO*, on data from individual plume intercepts. For each plume intercept, we calculated the  
186 excess mixing ratio of X in the fire plume, Δ*X*, as the average mixing ratio of X inside the  
187 plume,  $\bar{X}_{plume}$ , minus the average mixing ratio of X outside the plume,  $\bar{X}_{background}$ :

$$\Delta X = \bar{X}_{plume} - \bar{X}_{background}$$

188  $\bar{X}_{background}$  was calculated from the data obtained immediately before plume interception.  
189 Background mixing ratios of all species discussed herein were stable in the investigated  
190 domain. This analysis was performed for each of the fire overflights resulting in four data  
191 points, Δ*X* vs. Δ*CO*, for source emission characterization~~This analysis resulted in four data~~  
192 ~~points, Δ*X* vs. Δ*CO*, for characterizing source emission profiles.~~ A linear least-square  
193 regression analysis was then applied to these four data points, with the slope of the regression  
194 line describing the molar emission ratio (ER) of the species X relative to CO, ER<sub>X/CO</sub>, in ppbV  
195 ppmV<sup>-1</sup>. The precision of the CO data is better than ±1 ppbV which justifies the use of a  
196 univariate regression method. The standard error of the slope reflects both the natural  
197 variability in the plume and the measurement imprecision. A delayed instrument response was  
198 observed for formic acid and acetic acid. In-plume concentrations of these acids were derived  
199 as discussed in the Supplement.

200 The dilution-corrected molar excess mixing ratio of a species X,  $\Delta_{dil}X$  (in ppbV), at a  
201 downwind location was calculated from [the excess mixing ratio of CO observed at the fire](#)  
202 [source,  \$\Delta CO\_{source}\$ , and](#) the locally observed  $\Delta X$  and  $\Delta CO$  using the following equation:

$$\Delta_{dil}X = \Delta X \frac{\Delta CO_{source}}{\Delta CO}$$

203 By introducing this parameter, we are able to study loss or formation processes in the plume  
204 without confounding contributions from dilution. On a one-hour timescale, no photochemical  
205 loss of CO occurs and the contribution from photochemically formed CO to the large CO  
206 levels already present in the plume is negligible. Reported  $\Delta_{dil}X$  are average values from two  
207 longitudinal plume transects for which data were binned at 1 km spatial resolution.

208 The emission factor of a species X,  $EF_X$ , in g/kg was calculated according to Yokelson et al.  
209 (1999):

$$EF_X = F_c \times 1000 \times \frac{MM_X}{MM_C} \times \frac{C_X}{C_T}$$

210 with  $F_c$  being the mass fraction of carbon of the fuel,  $MM_X$  and  $MM_C$  the molecular masses of  
211 the species X and of carbon, and  $C_X/C_T$  the fraction of moles emitted as species X relative to  
212 the total number of moles carbon emitted.  $F_c$  was not measured during this study but 0.50 is a  
213 typical value for biomass (Burling et al., 2010). The accuracy of  $C_T$  is limited by unmeasured  
214 carbon. This fraction is assumed to be less than 2%. EFs were calculated as averages from the  
215 four fire overflights.

216 The oxygen-to-carbon (O:C) ratio of all detected NMOGs was calculated as follows:

$$\frac{O}{C} = \frac{\sum_i n_{O,i} X_i}{\sum_i n_{C,i} X_i}$$

217 with  $n_{O,i}$  and  $n_{C,i}$  being the number of oxygen atoms and carbon atoms in the species  $X_i$ ,  
218 respectively.

219 The modified combustion efficiency (MCE) was calculated as follows (Ferek et al., 1998):

$$MCE = \frac{\Delta CO_2}{\Delta CO + \Delta CO_2}$$

220 Aerosol mass was calculated from the 60-1000 nm integrated optical aerosol volume as  
221 measured by the UHSAS instrument assuming an average biomass burning secondary organic  
222 aerosol density of 1.3 g cm<sup>-3</sup> (Aiken et al., 2008).

## 223 2.4 Chemical box model calculations

224 We used the University of Washington Chemical Box Model (UWCM) (Wolfe and Thornton,  
225 2011) run on Master Chemical Mechanism (MCM) v3.3 chemistry (Jenkin et al., 1997, 2003,  
226 2015; Saunders et al., 2003) to simulate the downwind processing of trace gases in the  
227 biomass burning plume. The model was initialized using measured source concentrations of  
228 NO, NO<sub>2</sub>, HONO, O<sub>3</sub>, CO, CH<sub>4</sub> and of the 16 most abundant NMOGs detected by PTR-ToF-  
229 MS (ER<sub>X/CO</sub> > 1.0 ppbV ppmV<sup>-1</sup>; compounds identified in previous studies as detailed in  
230 paragraph 3.1.2). The model was run using the measured meteorological parameters (pressure,  
231 temperature, relative humidity, solar zenith angle) and the observed NO<sub>2</sub> photolysis rate. The  
232 model calculates dilution from the simple equation

$$\frac{dX}{dt} = -k_{dil}(X(t) - \bar{X}_{background})$$

233 where  $k_{dil}$  represents the dilution rate coefficient obtained from the decrease of  $\Delta CO$  versus  
234 plume travel time.  $k_{dil}$  was calculated in 285 s time bins (equivalent to 1 km distance bins).  
235 More information on the UWCM and the underlying theory can be found in Dillon et al.  
236 (2002), Wolfe and Thornton (2011) and Wolfe et al. (2012). ~~The dilution rate was obtained~~  
237 ~~from the measured molar excess mixing ratios of CO.~~ MCM v3.3 chemistry does not include  
238 the degradation of furan and 2-furfural, two highly reactive compounds with significant  
239 primary emissions from fires. We included these species in our chemical mechanism using the  
240 photolysis rates reported by Colmenar et al. (2015) and the OH reaction rates reported by  
241 Bierbach et al. (1992). We assumed that butenedial is the only primary reaction product of the  
242 reaction of furan with OH radicals (Aschmann et al., 2014). The atmospheric oxidation  
243 products of 2-furfural are unknown.

## 244 3 Results and discussion

### 245 3.1 Emissions

#### 246 3.1.1 Inorganic gases

247 Table 2 summarizes  $ER_{X/CO}$  and  $EF_X$  values of major inorganic gases as obtained from four  
248 source emission profiles. An MCE of  $0.90 \pm 0.02$  was derived from the measured CO and  
249 CO<sub>2</sub> data indicating stable burning conditions and roughly equal amounts of biomass  
250 consumption by flaming and smoldering combustion.

251 **Table 2.** Molar emission ratios (ER) relative to CO and emission factors (EF) of the major  
252 inorganic gases as obtained from four fire overflights.

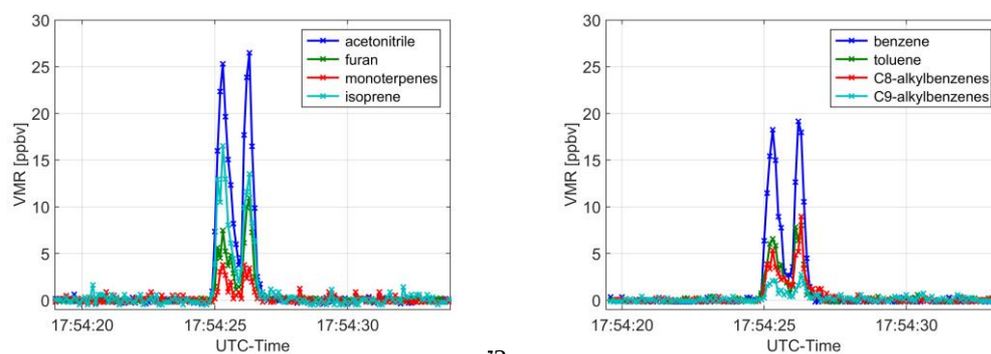
compound	$ER_{X/CO}$	$EF_X$ (g/kg)
CO <sub>2</sub>	--	$1623 \pm 68$
CO	--	$94.6 \pm 31.3$
NO	$10.4 \pm 5.2$	$0.63 \pm 0.51$
NO <sub>2</sub>	$9.4 \pm 2.0$	$1.24 \pm 0.06$
HONO	$2.0 \pm 0.7$	$0.15 \pm 0.05$
NH <sub>3</sub>	< 5.2	< 0.73

253 ERs and EFs of NO and NO<sub>2</sub> are within typical ranges reported in the literature (Akagi et al.,  
254 2011). The observed  $ER_{HONO/CO}$  of  $2.0 \pm 0.7$  ppbV ppmV<sup>-1</sup> is also in good agreement with  
255 previously reported values (e.g. Veres et al., 2010) increasing our confidence in the tentative  
256 identification and quantification of HONO emissions by PTR-ToF-MS. Excess mixing ratios  
257 of NH<sub>3</sub> in the plume were below the detection limit so that only an upper limit for  $ER_{NH_3/CO}$   
258 and  $EF_{NH_3}$  is reported.

#### 259 3.1.2 Organic gases

260 Methane (CH<sub>4</sub>) was the main organic gas emitted from the fire.  $ER_{CH_4/CO}$  and  $EF_{CH_4}$  are  $108.4$   
261  $\pm 13.4$  ppbV ppmV<sup>-1</sup> and  $6.25 \pm 2.86$  g/kg, respectively. This work, however, focuses on  
262 NMOG emissions. Figure 3a shows the 10 Hz time series of acetonitrile (CH<sub>3</sub>CN), furan  
263 (C<sub>4</sub>H<sub>4</sub>O), sum of monoterpene isomers (C<sub>10</sub>H<sub>16</sub>) and isoprene (C<sub>5</sub>H<sub>8</sub>) as measured during the  
264 overflight at 17:54:25 UTC (source emission profile 4). Figure 3b shows the time series of  
265 benzene (C<sub>6</sub>H<sub>6</sub>), toluene (C<sub>7</sub>H<sub>8</sub>), C<sub>8</sub>-alkylbenzene isomers (C<sub>8</sub>H<sub>10</sub>) and C<sub>9</sub>-alkylbenzene

266 isomers ( $C_9H_{12}$ ) for the same time period. The data demonstrate that the airborne PTR-ToF-  
267 MS instrument generates high-precision NMOG data even for very localized emission  
268 sources. The two small plumes discernible in Figures 1a and 1b are well resolved in the PTR-  
269 ToF-MS data shown in Figure 3. All signals instantly drop to background levels outside the  
270 plume confirming the excellent time response of the airborne PTR-ToF-MS instrument for  
271 analytes that do not adhere to instrumental surfaces.

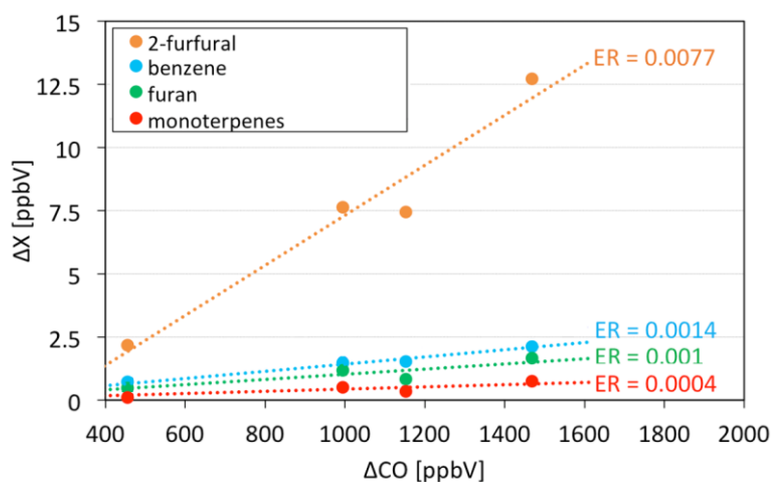


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\_ 13

274 **Figure 3.** 10 Hz time-series of (a) acetonitrile, furan, the sum of monoterpene isomers and  
275 isoprene and (b) benzene, toluene,  $C_8$ -alkylbenzene isomers and  $C_9$ -alkylbenzene isomers as  
276 measured during the fourth fire overflight at 17:54:25 UTC.

277 It is currently not possible to fully exploit these highly time resolved NMOG data for  
278 determining  $ER_{X/CO}$  because CO is only measured at 1-second time resolution.  $ER_{X/CO}$  values  
279 were thus obtained from average values for each source emission profile as described in  
280 section 2.3.



281  
 282 **Figure 4.** Average excess VMRs of 2-furfural, benzene, furan, and monoterpenes versus  
 283 average excess VMRs of CO. Each data point represents data from one fire overflight (source  
 284 emission profile). The slopes of the least-square regressions (dotted lines) correspond to the  
 285 initial molar emission ratios ( $ER_{X/CO}$ , in ppbV/ppbV).

286 Figure 4 shows  $\Delta X$  vs.  $\Delta CO$  as obtained for 2-furfural, benzene, furan, and monoterpenes  
 287 during each of the four fire overflights. The compounds were selected as representatives of  
 288 different chemical classes (including furans, aromatics, aldehydes, terpenes) that can have  
 289 different production mechanisms in the fire, e.g. furan being formed by pyrolysis and  
 290 monoterpenes just being evaporated (Yokelson et al., 1996). A strong linear relationship was  
 291 found not only for the species shown here but for all detected NMOGs indicating that source  
 292 emissions were near-stable during the 21 minute sampling period. This important finding will  
 293 later allow us to draw conclusions from analyte ratios measured downwind.

294 In total, 57  $m/z$  signals ( $NO^+$ ,  $NO_2^+$  and 55 C-containing ions) in the PTR-ToF-MS spectrum  
 295 showed an enhancement in the source emission profiles. Table 3 lists  $ER_{X/CO}$  and  $EF_X$  of the  
 296 18 ion signals that contain carbon atoms and that were observed with an  $ER_{X/CO} > 1$  ppbV  
 297  $ppmV^{-1}$ . These signals contribute 93% of the total NMOG emissions as detected by PTR-ToF-  
 298 MS. Emissions are dominated by formaldehyde, methanol, acetaldehyde and 2-furfural ( $EF >$   
 299  $1 \text{ g kg}^{-1}$ ). The complete list of all detected ion signals is given in Table S1 in the Supplement.

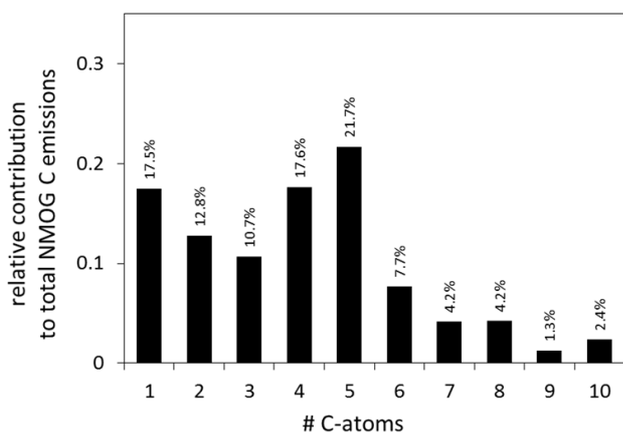
300 **Table 3.** Measured accurate  $m/z$ , elemental composition  $C_wH_xN_yO_z^+$  of the detected ion,  
 301 neutral precursor assignment based on literature information (significant interferants in

302 parentheses, tentative assignments in italic), emission factor (EF) and standard deviation (SD),  
 303 emission ratio (ER) and standard deviation for all detected NMOGs with  $ER_{X/CO} > 1$  ppbV  
 304  $\text{ppmV}^{-1}$ .

m/z	elemental composition	neutral precursor	EF	SD	ER	SD
			[g/kg]		[ppbV/ppmV]	
31.018	$\text{CH}_3\text{O}^+$	formaldehyde	2.31	0.57	22.7	1.3
33.034	$\text{CH}_5\text{O}^+$	methanol	2.25	1.06	19.6	2.0
42.034	$\text{C}_2\text{H}_4\text{N}^+$	acetonitrile	0.19	0.06	1.5	0.2
43.055	$\text{C}_3\text{H}_7^+$	propene (other unknown precursors)	0.64	0.25	4.5	0.2
45.034	$\text{C}_2\text{H}_5\text{O}^+$	acetaldehyde	1.52	0.50	10.4	0.3
47.020	$\text{CH}_3\text{O}_2^+$	formic acid	$\leq 0.13$	0.38	$\leq 1.4$	0.6
59.050	$\text{C}_3\text{H}_7\text{O}^+$	acetone (propanal)	0.83	0.31	4.1	0.1
61.029	$\text{C}_2\text{H}_5\text{O}_2^+$	acetic acid (glycolaldehyde)	0.47	0.18	2.7	0.3
69.034	$\text{C}_4\text{H}_5\text{O}^+$	furan	0.25	0.12	1.0	0.1
69.070	$\text{C}_5\text{H}_9^+$	isoprene (pentadienes, cyclopentene)	0.23	0.14	1.1	0.1
71.050	$\text{C}_4\text{H}_7\text{O}^+$	MVK (crotonaldehyde, MACR)	0.33	0.12	1.4	0.0
73.024	$\text{C}_3\text{H}_5\text{O}_2^+$	methylglyoxal	0.27	0.07	1.2	0.1
75.044	$\text{C}_3\text{H}_7\text{O}_2^+$	hydroxy acetone (methyl acetate, propionic acid)	0.28	0.15	1.1	0.1
79.055	$\text{C}_6\text{H}_7^+$	benzene	0.40	0.15	1.4	0.0
85.027	$\text{C}_4\text{H}_5\text{O}_2^+$	<i>dioxin, furanone</i>	0.39	0.12	1.5	0.1
87.043	$\text{C}_4\text{H}_7\text{O}_2^+$	2,3-butandione	0.44	0.18	1.6	0.1
97.029	$\text{C}_5\text{H}_5\text{O}_2^+$	2-furfural	2.31	1.07	7.7	0.6
111.041	$\text{C}_6\text{H}_7\text{O}_2^+$	<i>benzenediols, methylfurfural</i>	0.39	0.21	1.2	0.1

305 It is beyond the scope and possibilities of this work to make an independent assignment of  $m/z$   
 306 signals to specific neutral precursors. The P-3B payload did not include any NMOG analyzer  
 307 with higher analytical selectivity than the PTR-ToF-MS instrument. Our assignment of  $m/z$   
 308 signals to specific chemicals in Tables 3 and S2 thus exclusively relies on two recent studies  
 309 and the references used therein. Yokelson et al. (2013) used results from multiple analytical  
 310 techniques for assigning  $m/z$  peaks. Stockwell et al. (2015) used a high mass resolution PTR-  
 311 ToF-MS instrument for elemental composition determination and open-path FTIR data  
 312 together with literature reports for mass spectral interpretation. In the case of multiple neutral  
 313 precursors for a specific  $m/z$  signal, we considered only species with a relative contribution  $>$   
 314 10 % to the total signal. Two ion signals ( $m/z$  85.027 and  $m/z$  111.041) were not reported  
 315 previously. The assignment made is tentative and the compounds (in italic) were not included  
 316 in the modeling study. The reader is cautioned that this is still an evolving field of research  
 317 and some signals may be misassigned or suffer from yet unknown interferences.

318 Total observed carbon emitted as NMOGs (55 ion signals) was 10,472 ppbC. The O/C ratio at  
319 the fire source was 0.41. Figure 5 shows the relative contribution of C<sub>1</sub> to C<sub>10</sub> compounds to  
320 total NMOG emissions on a carbon atom basis.



321  
322 **Figure 5.** Relative contributions of C<sub>1</sub>-C<sub>10</sub> compounds to total NMOG carbon emissions. C<sub>1</sub>  
323 to C<sub>5</sub> compounds each have relative contributions > 10%, and in sum contribute ~ 80% of the  
324 total NMOG carbon emissions.

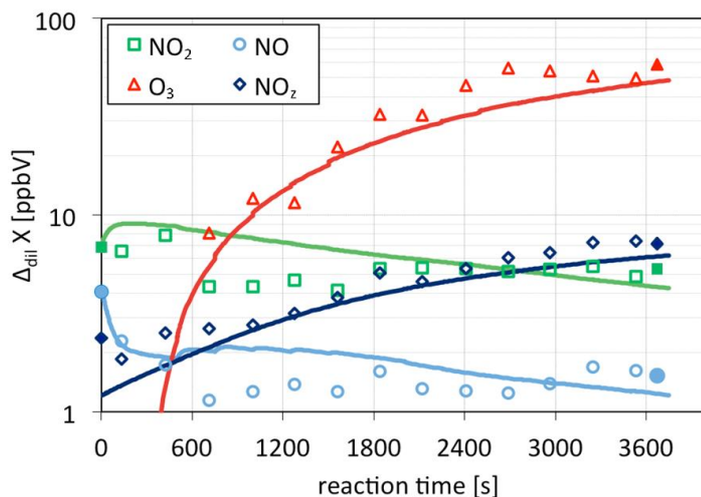
325 The dominant contribution to NMOG carbon emissions came from the C<sub>5</sub>-compound 2-  
326 furfural. Significant carbon emissions ( $ER_{X/CO} > 50 \text{ pptV ppmV}^{-1}$ ) were detected only up to  
327 C<sub>10</sub> (monoterpenes).

### 328 3.2 Plume evolution

329 The NASA P-3B sampled the downwind plume for approximately 2 minutes of flight time. At  
330 an average wind speed of  $3.5 \text{ m s}^{-1}$ , this corresponds to approximately one hour of  
331 atmospheric plume processing. Volume mixing ratios of inert tracers (CO<sub>2</sub>, CO, acetonitrile  
332 and benzene) consistently decreased by a factor of ~ 13.5 during the two longitudinal plume  
333 transects. We used this decrease to derive dilution-corrected molar excess mixing ratio of  
334 reactive trace gas species X,  $\Delta_{dil}X$  (see paragraph 2.3).  $\Delta_{dil}X$  were used to investigate  
335 downwind plume chemistry by observations and by a 0-D photochemical box model  
336 simulation initialized with measured emission data.

337 3.2.1 Ozone formation and sequestration of nitrogen oxides

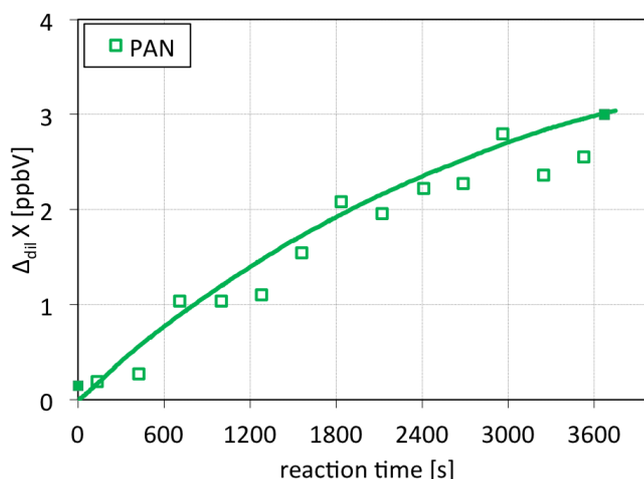
338 Figure 6 shows dilution-corrected molar excess mixing ratios of O<sub>3</sub>, NO, NO<sub>2</sub> and NO<sub>z</sub> (=NO<sub>y</sub>-  
 339 NO-NO<sub>2</sub>) during one hour of atmospheric plume processing. Point symbols refer to the  
 340 measured data; solid lines represent the output of the UWCM based on MCM v3.3 chemistry.



341  
 342 **Figure 6.** Dilution-corrected molar excess mixing ratios of O<sub>3</sub>, NO, NO<sub>2</sub> and NO<sub>z</sub> (= NO<sub>y</sub>-  
 343 NO-NO<sub>2</sub>) during one hour of plume evolution (in one kilometer bins). Point symbols refer to  
 344 the measured data; solid lines represent the output of the UWCM based on MCM v3.3  
 345 chemistry.

346 Ozone is efficiently formed in the plume in the presence of NO<sub>x</sub> and NMOGs. Close to the  
 347 source (t < 600 s), ambient O<sub>3</sub> reacts with abundantly emitted NO resulting in negative O<sub>3</sub>  
 348 excess mixing ratios (not displayed on the logarithmic ordinate of Figure 6). After ~ 10  
 349 minutes of plume processing net ozone formation starts, resulting in a dilution-corrected  
 350 increase of O<sub>3</sub> on the order of 50-60 ppbV during the first hour the plume resides in the  
 351 atmosphere. The UWCM (MCM v3.3 chemistry; initialized with measured emissions of NO,  
 352 NO<sub>2</sub>, HONO, O<sub>3</sub>, CO, CH<sub>4</sub> and 16 NMOGs) simulates the evolution of O<sub>3</sub>, NO and NO<sub>2</sub> well.  
 353 An even better agreement in the ozone evolution is obtained if the model is constrained to  
 354 measured formaldehyde values which slightly exceed the modeled values at t > 1500 s (see  
 355 see paragraph 3.2.2). O<sub>3</sub> formation is fueled by HO<sub>2</sub>/CH<sub>3</sub>O<sub>2</sub>+NO reactions. The model  
 356 indicates that HO<sub>2</sub> radicals are primarily generated in the CO+OH, 2-furfural+OH and  
 357 formaldehyde+OH reactions. CH<sub>3</sub>O<sub>2</sub> radicals are primarily formed in the CH<sub>3</sub>C(O)O<sub>2</sub>+NO

358 and  $\text{CH}_4 + \text{OH}$  reactions; the main precursors of  $\text{CH}_3\text{C}(\text{O})\text{O}_2$  radicals are acetaldehyde, 2,3-  
359 butanedione and methylglyoxal.



360

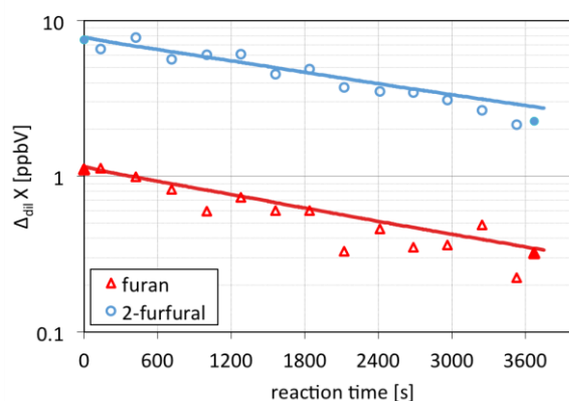
361 **Figure 7.** Dilution-corrected molar excess mixing ratios of PAN during one hour of plume  
362 evolution (in one kilometer bins). Point symbols refer to the measured data; the solid line  
363 represents the output of the UWCM based on MCM v3.3 chemistry.

364 The model also accurately captures the net formation of  $\text{NO}_z$  ( $= \text{NO}_y - \text{NO} - \text{NO}_2$ ). Modelled  
365  $\text{NO}_z$  sums all species in the MCM v3.3 degradation scheme that include nitro or nitroso  
366 groups. The main contributors to  $\text{NO}_z$  being formed are peroxyacetyl nitrate (PAN) and nitric  
367 acid ( $\text{HNO}_3$ ). The model simulates  $\Delta_{\text{dil}} \text{PAN} = 3$  ppbV and  $\Delta_{\text{dil}} \text{HNO}_3 = 2.4$  ppbV after one  
368 hour of plume evolution which accounts for  $\sim 90\%$  of all  $\text{NO}_z$  formed. Under the operating  
369 conditions used in this study, PAN is predominantly detected at  $m/z$  45.992 ( $\text{NO}_2^+$ ) by the  
370 PTR-ToF-MS instrument (Hansel and Wisthaler, 2000). Using a PAN calibration factor  
371 obtained in a previous study, we obtain an excellent agreement between measured and  
372 modeled PAN concentrations (Figure 7).

### 373 3.2.2 Evolution of NMOGs

374 Fire emissions include many NMOGs that quickly react with OH radicals. OH radicals are  
375 abundantly formed in biomass burning plumes causing highly reactive NMOGs to disappear  
376 even on the one-hour time scale investigated in this study (Akagi et al., 2012, 2013; Hobbs et  
377 al., 2003). Figure 8 shows dilution-corrected mixing ratios of furan and 2-furfural during one

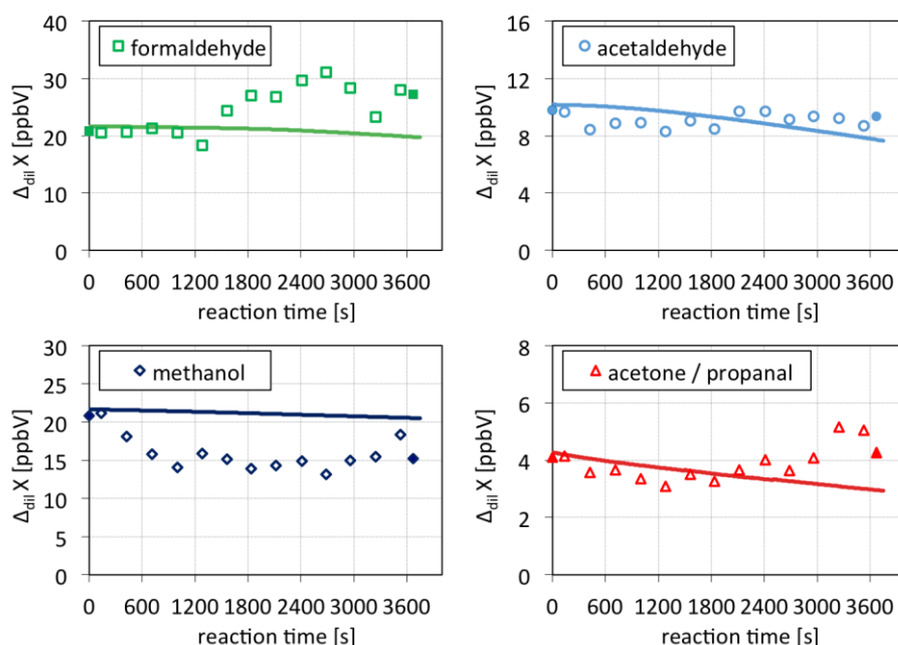
378 hour of plume evolution. Point symbols refer to the dilution-corrected experimental data; solid  
379 lines represent the output of the UWCM. Measured and modeled data are in excellent  
380 agreement confirming that we observed the OH-initiated degradation of furan and 2-furfural.  
381 The influence of interfering isomers (or fragment ions), if any, is small. The box model output  
382 indicates near-stable OH radical concentrations of  $7.45 \pm 1.07 \times 10^6 \text{ cm}^{-3}$  along the 13 km  
383 downwind transect. Other studies (eg. Yokelson et al., 2009) have reported similarly high  
384 average OH levels in biomass burning plumes.



385  
386 **Figure 8.** Dilution-corrected molar excess mixing ratios of furan and 2-furfural during one  
387 hour of plume evolution. Point symbols refer to the measured data (one kilometer bins); solid  
388 lines represent the output of the UWCM fed with MCM v3.3 chemistry.

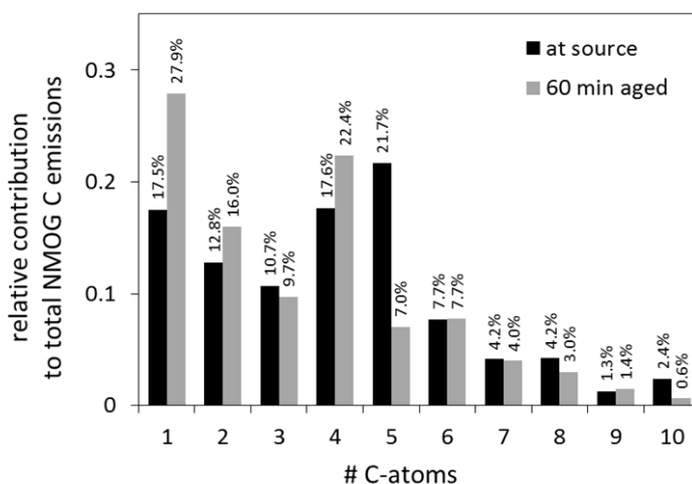
389 Figure 9 shows dilution-corrected mixing ratios of four important oxygenated NMOGs,  
390 formaldehyde, acetaldehyde, methanol and acetone/propanal. Point symbols again refer to the  
391 dilution-corrected experimental data; solid lines represent the output of the UWCM.  
392 Formaldehyde and acetone/propanal show a distinct increase after half an hour of plume  
393 processing, which is not captured by the model simulation based on MCM v3.3 degradation  
394 chemistry of the 16 most abundant NMOGs (as detected by PTR-ToF-MS). Interestingly, the  
395 experimental data indicate a significant loss of methanol during the initial 15 minutes of  
396 plume processing. This sink is also not included in MCM v3.3 chemistry and heterogeneous  
397 loss processes should be investigated. The observed initial drop could, however, also be  
398 caused by an unknown highly reactive compound that interferes with the detection of  
399 methanol. In addition to the carbonyls discussed above, acetic acid/glycolaldehyde and the  
400  $\text{C}_4\text{H}_3\text{O}_3^+$  signal, which is tentatively assigned to maleic acid/maleic anhydride, exhibited

401 dilution-corrected increases of  $\sim 1.5$  ppbV and  $\sim 1$  ppbV, respectively. The model was unable  
402 to capture the observed increase. This does not come as a surprise since these species are  
403 typical higher-order degradation products that are not included in MCM v3.3 degradation  
404 schemes.



405  
406 **Figure 9.** Dilution-corrected molar excess mixing ratios of formaldehyde (a), acetaldehyde  
407 (b), methanol (c), and acetone/propanal (d) during one hour of plume evolution. Point  
408 symbols refer to the measured data (one kilometer bins); solid lines represent the output of the  
409 UWCM fed with MCM v3.3 chemistry.

410 Figure 10 compares the relative contributions of  $C_1$  to  $C_{12}$  compounds to total NMOG carbon  
411 measured at the fire source and at the 1-hour downwind location.  $C_1$ ,  $C_2$  and  $C_4$  compounds  
412 exhibited the largest relative increase. The observed O/C ratio at the 1-hour downwind  
413 location source was 0.56, compared to 0.41 observed at the source. This is consistent with the  
414 conceptual picture of a photochemical breakdown of NMOGs into smaller, more oxidized  
415 species.



416

417 **Figure 10.** Relative contributions of C<sub>1</sub> to C<sub>12</sub> compounds to total NMOG carbon measured  
 418 at the fire source and at the 1-hour downwind location.

### 419 3.2.3 Gas-to-particle conversion

420 A dilution-corrected mass balance analysis reveals that 40.8 μg cm<sup>-3</sup> of the mass initially  
 421 emitted as NMOGs was lost during one hour of atmospheric processing. This equals 24% of  
 422 the carbon initially emitted as NMOGs. At the same time, the dilution-corrected total particle  
 423 mass concentration as derived from UHSAS measurements increased by ~78 μg m<sup>-3</sup>. These  
 424 mass concentration calculations are only approximate (for details see paragraph 2.2), but this  
 425 analysis suggests that about 50% of the aerosol mass formed in the downwind plume is  
 426 organic in nature. This agrees with findings from previous studies that observed significant  
 427 organic and inorganic aerosol formation in aging biomass burning plumes (Cubison et al.,  
 428 2011; Yokelson et al., 2009). Given that photo-oxidation of 2-furfural has the highest mass  
 429 turnover, secondary organic aerosol formation from the 2-furfural + OH reaction should be  
 430 investigated in laboratory experiments.

## 431 4 Summary and Conclusion

432 A plume emanating from a small forest understory fire was investigated in an airborne study.  
 433 High spatio-temporal resolution data were obtained for inorganic and organic trace gases, the  
 434 latter being sampled for the first time at 10 Hz using a PTR-ToF-MS instrument. We  
 435 generated quantitative emission data for CO<sub>2</sub>, CO, NO, NO<sub>2</sub>, HONO, NH<sub>3</sub> and 16 NMOGs

436 with  $ER_{X/CO} > 1.0$  ppbV ppmV<sup>-1</sup>. NMOG emissions were dominated by formaldehyde,  
437 acetaldehyde, 2-furfural and methanol. No NMOGs with more than 10 carbon atoms were  
438 observed at mixing ratios larger than 50 pptV per ppmV CO emitted. Downwind plume  
439 chemistry was investigated both by observations and by a model simulation using near-  
440 explicit MCM v3.3 chemistry. The observed dilution-corrected O<sub>3</sub> increase on the order of 50-  
441 60 ppbV was well captured by the model, which indicated carbonyls (formaldehyde,  
442 acetaldehyde, 2,3-butanedione, methylglyoxal, 2-furfural) in addition to CO and CH<sub>4</sub> as the  
443 main drivers of peroxy radical chemistry. The model also accurately reproduced the  
444 sequestration of NO<sub>x</sub> into PAN and the degradation of furan and 2-furfural at average OH  
445 plume concentrations of  $7.45 \pm 1.07 \times 10^6$  cm<sup>-3</sup>. Formaldehyde, acetone/propanal, acetic  
446 acid/glycolaldehyde and maleic acid/maleic anhydride (tentative identification) were found to  
447 increase during one hour of atmospheric plume processing, with the model being unable to  
448 capture the increase. A dilution-corrected mass balance analysis suggests that about 50% of  
449 the aerosol mass formed in the downwind plume is secondary organic in nature.

450 We conclude that the PTR-ToF-MS instrument is a powerful analytical tool for airborne  
451 plume studies. The generated data are highly valuable in characterizing point source  
452 emissions and near-field chemical transformations. Key chemical processes (ozone and  
453 radical formation, NO<sub>x</sub> sequestration) in an aging biomass burning plume were accurately  
454 simulated using a 0-D photochemical box model run with up-to-date and near-explicit MCM  
455 v3.3 chemistry.

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473

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