- 1 Reconciling the total carbon budget for boreal forest wildfire emissions using airborne
- 2 observations
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17 Abstract

18 Wildfire impacts on air quality and climate are expected to be exacerbated by climate change with the

19 most pronounced impacts in the boreal biome. Despite the large geographic coverage, there is a lack

20 of limited information on boreal forest wildfire emissions, particularly for organic compounds, which are

21 critical inputs for air quality model predictions of downwind impacts. In this study, airborne

22 measurements of 193 250 compounds from 15 instruments, including 173228 non-methane organics

23 compounds (NMOG), were used to provide the most detailed characterization, to date, of boreal forest

24 wildfire emissions. Highly speciated measurements showed a large diversity of chemical classes

25 highlighting the complexity of emissions. Using measurements of the total NMOG carbon (NMOG<sub>T</sub>), the

26  $\Sigma$ NMOG was found to be 46.250±3 to 53±3 % of NMOG<sub>T</sub>, of which, the intermediate- and semi-volatile

27 organic compounds (I/SVOCs) were estimated to account for 7.47 to 10 %. These estimates of I/SVOC

28 emission factors expand the volatility range of NMOG typically reported. Despite extensive speciation, a

substantial portion of NMOG<sub>T</sub> remained unidentified ( $46.447\pm15$  to  $50\pm15$  %), with expected

30	contributions from more highly-functionalized VOCs and I/SVOCs. The emission factors derived in this
31	study improve wildfire chemical speciation profiles and are especially relevant for air quality modelling
32	of boreal forest wildfires. These aircraft-derived emission estimates were further linked with those
33	derived from satellite observations demonstrating their combined value in assessing variability in
34	modelled emissions. These results contribute to the verification and improvement of models that are
35	essential for reliable predictions of near-source and downwind pollution resulting from boreal forest
36	wildfires.
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#### 43 **1 Introduction**

44 Wildfires play a natural role in maintaining forest health and diversity through the release of 45 nutrients, seed germination, removal of aging vegetation, and reducing the spread of forest diseases. 46 Wildfires are, however, one of the largest global sources of trace gases and aerosols to the atmosphere 47 (Andreae, 2019; Yu et al., 2019) and can have deleterious impacts on human health (Cascio, 2018; Cherry 48 and Haynes, 2017; Reid et al., 2016; Finlay et al., 2012), air quality (Landis et al., 2018; Miller et al., 49 2011; Rogers et al., 2020), ecosystems (Kou-Giesbrecht et al., 2019; Campos et al., 2019; Kallenborn et 50 al., 2012; Johnstone et al., 2010) and climate (Randerson et al., 2006). Not only can wildfire pollutants 51 fumigate local source areas, they can be transported over long distances resulting in degraded air quality 52 in locations far from fire sources (Miller et al., 2011; Rogers et al., 2020), and pose threats to downwind 53 ecosystems through wet and dry deposition processes (Kou-Giesbrecht et al., 2019; Kallenborn et al., 54 2012; Campos et al., 2019).

55 The severity and frequency of wildfires is expected to increase in response to climate change 56 (Bush and Lemmen, 2019; Seidl et al., 2017; Whitman et al., 2019) with evidence to suggest that Wildfire 57 impacts on air quality and climate are expected to be exacerbated by climate change (Bush and Lemmen, 58 <del>2019; Seidl et al., 2017; Whitman et al., 2019) and</del> such impacts are expected to be most pronounced in 59 the boreal biome (Seidl et al., 2017; Whitman et al., 2019). The boreal forest zone is the most northerly 60 of all forest biomes accounting for 1.2 billion ha of mostly coniferous forest and comprising about 61 3033 % of the global forest area, or 11-14 % of the earth's land surface (https://www.nrcan.gc.ca/our-62 natural-resources/forests/sustainable-forest-management/boreal-forest/8-facts-about-canadas-boreal-63 forest/17394). On a global basis, boreal forest wildfires are responsible for an estimated 20 % of yearly 64 global biomass burning emissions (van der Werf et al., 2006). Canada's boreal forests account for 65  $\sim 2830$  % of the global boreal zone area and encompasses 75 % of Canada's 347 million ha of forested 66 land (Fig. S1) (https://www.nrcan.gc.ca/our-natural-resources/forests/sustainable-forest-67 management/boreal-forest/8-facts-about-canadas-boreal-forest/17394. In the past decade, Canada has

68 experienced unprecedented fire seasons, with large numbers of evacuations, major property damage, poor

- 69 air quality and significant economic impacts (NRCan, 2018; Landis et al., 2018; McGee et al., 2015).
- 70 Model predictions have suggested that Canadian fire occurrences will increase by 25 % by 2030 from a

71 1975 to 1990 baseline scenario (Wotton et al., 2010).

72 To adequately assess and mitigate the risks of wildfire emissions to human and ecosystem health, 73 reliable pollutant predictions are required which depend on accurate and detailed fire emissions data. 74 Such emissions data are developed by multiplying emission factors and ratios with the mass of biomass 75 burned (Chen et al., 2019). In Canada, Environment and Climate Change Canada (ECCC) provides 76 predictions of particulate matter (PM) (<2.5  $\mu$ m in diameter) from wildfire smoke to the public using the 77 FireWork modelling system that combines forecast meteorology, emissions inputs (e.g. emission factors), 78 forest fire and fuel data (e.g. fuel maps, plume height parameterization), and a regional air quality model, 79 GEM-MACH (details in Chen et al., 2019). FireWork is also used for air quality research studies with 80 significantly more complex chemical mechanisms for emissions characterization and detailed physical 81 processes. Wildfire field studies, as well as prescribed burns and laboratory work, have resulted in 82 valuable global databases of fire emission factors covering a broad range of ecosystems and geographic 83 areas (e.g. Andreae, 2019; Akagi et al., 2011), however, they are primarily concentrated on the temperate 84 forests of the American mid-west and savannah/grasslands of Africa (e.g. Andreae 2019; Permar et al., 85 2021; Palm et al., 2020; Lindaas et al., 2020; Roberts et al., 2020; Juncosa-Calaharrano et al., 2021; 86 Coggon et al., 2019; Koss et al., 2018; Hatch et al., 2017). Until now, the most complete characterization 87 of boreal forest wildfire emissions in Canada was provided by Simpson et al. (2011) which relied on 88 whole air canisters with offline analysis for organic compounds. Due to a lack of limited comprehensive 89 emission data specific for boreal wildfires, air quality models for northern regions face significant 90 challenges resulting in uncertain predictions of emissions, exposure and associated impacts. 91 In the summer 2018, a research aircraft was deployed to measure emissions and subsequent

92 transformation processes from an active-boreal forest wildfire in western Canada (Fig. 1; Fig. S1). In this

93 paper, measurements of a comprehensive suite of detailed emissions information is provided from an

94 active, near field boreal forest wildfire using a detailed measurement suite of over 200 gas- and particle-95 phase compounds are used to. Emissions of highly speciated non-methane organic gases (NMOG) are 96 provide a detailed characterization of smoldering wildfire edemissions. The highly speciated non-97 methane organic gas (NMOG) measurements are described by broad chemical classes and across a range 98 of volatilities extending from VOCs to SVOCs. The wide range of measured Speciated NMOGs, along 99 with concurrent total NMOG carbon (NMOG<sub>T</sub>) measurements, provides a unique opportunity to reconcile 100 the total carbon budget. Emission factors are derived for all measured 193 compounds which represents 101 the most extensive chemical speciation of wildfire emissions to date, almost tripling the number of 102 reported values for the boreal forest ecosystem in the Andreae (2019) compilation paper. Emission 103 estimates are also combined resulting in more relevant emissions information for boreal forest wildfires 104 and improved emission quantification and chemical speciation representations in air quality models. 105 Combining aircraft derived emissions with those from satellite observations demonstrates usefulness to 106 evaluate modelled emissions diurnal variability. The purpose of this work is to provide relevant The 107 emissions information for boreal forest wildfires to ultimately in this work will contribute towards 108 improved emissions quantification and chemical speciation representations in air quality models. 109 verification and improvements of models that are essential for reliable predictions of boreal forest 110 wildfires pollutants. 111 112 2 Methods 113 **2.1 Aircraft measurements** 114 The NRC's Convair-580 research aircraft was deployed on June 25, 2018 to sample a wildfire 115 detected to the east of the Alberta/Saskatchewan border (56.4°N, 109.7°W) (Fig. 1). Measurements of a 116 comprehensive suite of trace gases, particles and meteorology were made with high time resolution. 117 Meteorological measurements including relative humidity, temperature, wind direction and speed, as well 118 as aircraft state parameters such as altitude (masl) and geographic coordinates were conducted at 1 sec 119 intervalstime resolution. A detailed description of the various measurements methods with references is

provided in the supporting information (SI Sect. 21.1, Table S1, S2), with only a brief description
provided here.

122 **2.1.1 Trace gas measurements** In-situ measurements of NO, NO<sub>2</sub>, NO<sub>y</sub>, O<sub>3</sub> and SO<sub>2</sub> were conducted 123 using commercial instruments (Thermo Scientific Inc.) modified to measure at 1 sec time resolution. 124 Ammonia (NH<sub>3</sub>) measurements were made at 1 sec time resolution using a Los Gatos Research (LGR) 125 NH<sub>3</sub>/H<sub>2</sub>S Analyzer, model 911-0039. Calibrations were conducted periodically throughout the 126 measurement study using NIST certified standards. Instrument zeros were performed for all these 127 instruments 3.5 times per flight for a duration of ~3.5 minutes each time at the beginning, during and 128 after each flight. Gas phase elemental Hg (GEM) was measured with a Tekran 237X instrument (Tekran 129 Instruments Corporation) modified to allow a reduced sampling time of 2 min (McLagan et al., 2021; 130 Cole et al., 2014). CO, CO<sub>2</sub> and CH<sub>4</sub> were measured with a Cavity Ring Down spectroscopy instrument 131 (Picarro G2401-m). A second Picarro G2401-m instrument was used to measure Total Carbon (TC, in 132 units of ppm C) by passing the sample air through a catalyst to convert all carbon species to  $CO_2$ . through 133 a platinum catalyst (Shimadzu) which was placed at the external rear facing inlet assembly and 134 maintained at 650 °C, adapted from Stockwell et al. (2018) and Veres et al., (2010 TTotal non-methane 135 organic gases (NMOG<sub>T</sub>), in mixing ratios units of ppm C, were was quantified by subtracting the ambient 136 CH<sub>4</sub>, CO and CO<sub>2</sub> measurements (instrument without the upstream catalyst) from the TC measurements 137 (see SI Methods for more details).

138 Individually speciated NMOGs (as well as some inorganic species) were measured with a 139 Chemical Ionization Mass Spectrometer (CIMS), a Proton Transfer Time-of-Flight Mass Spectrometer 140 (PTRMS), and through whole air sampling using canisters (Advanced Whole Air Sampler; AWAS). In 141 addition, integrated cartridge-based samples were taken. The CIMS (a modified Tofwerk/Aerodyne Api-142 ToF) was operated using iodide as the reagent ion providing 1 sec time resolved measurements for 30 143 compounds (Table S2). The PTRMS (Ionicon Analytik GmbH, Austria) used chemical ionization with 144  $H_3O^+$  as the primary reagent ion providing 1 sec measurements for a suite of organic compounds. For 145 those compounds with no available gas standard, a relative response factor was calculated with reaction

146 rate constants using the method described in Sekimoto et al. (2017) and guided by the work of Koss et al. 147 (2018) ('calculated' compounds). Integrated 'grab' samples (20-30 sec) were collected from the aircraft 148 using the Advanced Whole Air Sampler (AWAS) with offline analysis. The AWAS provided speciated 149 measurements of hydrocarbons ( $\leq < C10$ ), but no oxygenates. Overlapping compounds/isomers that were 150 measured by both the PTRMS and AWAS, as well as between the PTRMS and CIMS, were handled as 151 described in SI Sect. 12.1.4. Integrated gas phase samples were collected using an automated adsorbent 152 tube (i.e. cartridge) sampling assembly with offline analysis (Ditto et al., 2021; Sheu et al., 2018; Khare et 153 al., 2019). These samples provided targeted measurements of gas-phase compounds ranging in volatility 154 from C<sub>10</sub> volatile organic compounds (VOCs) to C<sub>25</sub> semivolatile organic compounds (SVOCs) including 155 hydrocarbons (CH), and functionalized compounds containing 1 oxygen atom (CHO<sub>1</sub>), and 1 sulfur atom 156  $(CHS_1)$ .

### 157 **2.1.2 Particle measurements**

Particle chemistry was obtained with a high resolution aerosol mass spectrometer (AMS)
(Aerodyne) providing mass concentrations of particle species including total organics (OA), NO<sub>3</sub>, SO<sub>4</sub>
and NH<sub>4</sub> for particles less than ~1 µm. Particle size distributions were measured between 60 and 1000
nm at 1 sec time resolution using the Ultra High Sensitivity Aerosol Spectrometer (UHSAS; Droplet
Measurement Technologies). Refractory black carbon (rBC) was measured using a single particle soot
photometer (SP2; Droplet Measurement Technologies).

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#### 165 **2.2 Flight and fire description**

A wildfire located near Lac La Loche in Saskatchewan (56.40°N 109.90°W) was detected by satellite on June 23 (Fig. 1; Fig. S1). The fire was ignited by lightning on June 23, 2018 at 19:45 UTC and lasted 50 hrs to June 25 21:41 UTC burning an estimated 10,000 ha before being extinguished by rain. The area burned was mostly mature Jack pine and boreal spruce forest with a smaller fraction of boreal mixed-wood forest. Satellite images from the VIIRS spectroradiometer on the Suomi NPP and NOAA-20 satellites taken on June 25 showed merged fire hot spots with a visible smoke plume moving 172 in a north-westerly direction (Fig. 1; see SI Sect. 2.02 for more details). Lagrangian flight tracks were 173 flown downwind of the wildfire to follow the fire plumes. Multiple horizontal transects, vertically 174 stacked and perpendicular to the plume direction were made at different altitudes from 640 to 1460 m asl 175 (~220 – 1040 m agl, based on 420 m asl at Lac La Loche) forming virtual screens. Five screens were 176 completed over two flights with the closest screen  $\sim 10$  km and the farthest screen 164 km downwind of 177 the fire, with the screens spaced such that the instruments sampled the same air parcels as they were 178 transported downwind. A vertical profile which typically reached ~2500 m asl was conducted in the 179 plume at each screen to gather information on its vertical structure and the height of the plume. As 180 demonstrated by the elevated CO mixing ratios in Fig. 2, two distinct plumes were identified - a south 181 plume (SP) and north plume (NP), that were transported in parallel in a northwesterly direction. The SP 182 is estimated to be ~42 min old based on the measured wind speed at Screen 1 and the distance from the 183 closest edge of the VIIRS fire hot spots (~10 km). The NP is estimated to be an additional 30 min older 184 than the SP (further details in SI Sect. 2.02). For the purposes of this investigation, only data from Screen 185 1 are used to characterize the direct emissions from this fire. Evaluation of emissions of photolabile 186 species could be influenced by photochemical and depositional losses that may take place between the 187 time of emission and the time of measurement. sprimary at 10 km (<1 hr) away from the fire source, 188 Screen 1 measurements represent some of the freshest emissions ever measured under wildfire conditions. 189 There are were no other significant anthropogenic sources like upwind urban or industrial areas, 190 impacting the Screen 1 measurements. Plume evolution during transport from Screen 1 to downwind 191 Screens 2 to 5 is discussed in other papers (Liu et al., 2022; Ditto et al., 2021; McLagan et al., 2021). 192 193 2.3 Emission ratios, emission factors and combustion efficiency

194 **2.3 Emission ratios** 

Emission ratios (ERs) were calculated using an integration method (e.g. Yokelson et al., 2009; Garofalo et al., 2019) with-using the in-plume measurements for the SP and NP. The integration method was carried out for the real-time measurements by first subtracting a background from the in-plume

198 measurements. Background measurements were defined as the average over short time segments (~30 199 sec) outside and at the same altitude as inside the plume, and typically selected at the ends of the 200 horizontal transects. The background-subtracted plume measurements yielded enhanced plume values 201 (e.g.  $\Delta X(t)$ ) which were then integrated using the plume start and end times guided by when CO mixing 202 ratios were above the CO background. Nominal plume time periods are indicated by the vertical grey 203 bars in Fig. 3 which shows time series for CO, NMOG, OA and acetonitrile for the first 4 of 5 transects 204 on Screen 1. Integrated pollutant values were subsequently normalized by the integrated values of CO 205 (Eq. 1) to account for changes due to dilution producing emission ratios (ER) for the SP and NP for each 206 transect on Screen 1.

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$$ER = \frac{\int_{start}^{end} \Delta X(t) (dt)}{\int_{start}^{end} \Delta CO(t) (dt)}$$
(1)

209

CO is known to be a suitable dilution tracer as it has a long atmospheric lifetime of 1-4 months (Seinfeld and Pandis, 1998), is unreactive on the time scale of the measurements. , and is a particularly good tracer for smoldering fires (e.g. Simpson et al., 2011). In this study, ERs were calculated using CO as it was well-enhanced above a background of ~0.119±0.005 ppmv for the plumes measured, there were no other significant CO sources in the study area, and CO is a particularly good tracer for smoldering fires (e.g. Simpson et al., 2011).and co varied well with the majority of measurements.

ERs for the AWAS compounds were determined using the average mixing ratio of 3 samples taken in the SP and two in the NP, and the average mixing ratio of two background samples. CO mixing ratios were averaged across the AWAS sample time period. For the integrated cartridges, samples were collected over the lower set of aircraft transects ('LOW') and higher set of transects ('HIGH'), resulting in two integrated cartridge samples for each screen. The HIGH sample was used as the background. The HIGH sample was collected largely outside the wildfire plume, but may have been influenced to some extent from emissions. However, this impact is expected to be minimal as average CO mixing ratios 223 during the HIGH sample were at background levels (~0.14 ppmv). Nevertheless, to address the potential 224 for influence of the plume in the HIGH sample, the ERs are presented as ranges with the lower estimates 225 derived by subtracting the HIGH background sample, and the upper estimates without subtracting the 226 HIGH sample. This calculation is described in Eq. 2 where  $Cartridge_{LOW}$  represents the LOW cartridge 227 sample measurements,  $Cartridge_{BKGD}$  is the background derived from the HIGH cartridge sample 228 measurements, and  $CO_{LOW}$ ,  $CO_{BKGD}$  are the average CO concentrations during the respective LOW and 229 HIGH cartridge integration time periods. The uncertainty with this bounding analysis is acknowledged, 230 but the I/SVOCs ERs within a plume are likely to vary similar to other work (Hatch et al., 2018).

$$ER = \frac{Cartridge_{LOW} - Cartridge_{BKGD}}{CO_{LOW} - CO_{BKGD}} \ to \ \frac{Cartridge_{LOW} - 0}{CO_{LOW} - CO_{BKGD}}$$
(2)

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231

#### 233 **2.4 Emission factors**

234 Emission factors (EFs) were determined as the mass of species X emitted per unit mass of dry 235 fuel burned in g kg<sup>-1</sup> assuming that all of the carbon in the fuel was released into the atmosphere and 236 measured (Ward and Radke, 1993; Yokelson et al., 2007), and that the mass fraction of carbon in the fuel 237 is constant. EFs were determined using Eq. 32 where  $F_c$  is the mass fraction of carbon in the fuel and 238 estimated to be 0.5 (de Groot et al., 2009 and references therein),  $mm_x$  is the molar mass of the compound 239 of interest, and mm<sub>c</sub> is the molar mass of carbon, 12 g mol<sup>-1</sup>,  $\Delta X$  is the integrated background-subtracted 240 mixing ratio or concentration of the species of interest,  $\Delta TC$  is the integrated background-subtracted total 241 carbonTC. TC otal Carbon (TC) (see Sect. 2.1) was directly measured and includes all the carbon mass in 242 CO<sub>2</sub>, CO, CH<sub>4</sub>, and NMOG<sub>T</sub>, as well as that from particulate black carbon (rBC) and particulate organic 243 carbon (OC) (which were added to the TC), for a complete accounting of all the emitted carbon. For 244 species measured in mass concentration units, Eq. 32 was modified by converting TC to mass 245 concentrations using the measured temperature and pressure, and removing the molar mass ratio term. 246 The EFs for the AWAS and the cartridge samples were derived using the average measurements as 247 discussed for the ER, but with TC as the denominator.

$$EF\left(\frac{g}{kg}\right) = F_c x \ 1000 \ \left(\frac{g}{kg}\right) x \ \frac{mm_X}{mm_c} \ x \ \frac{\int_{start}^{end} \Delta X(t) \ (dt)}{\int_{start}^{end} \Delta TC(t) \ (dt)} \frac{\Delta X}{\Delta TC}$$
(32)

250

EFs were determined for the SP and NP for each transect, and then averaged to obtain screen-averaged
EFs for the SP and the NP, as well as for both plumes together. There is a potential for inherent
uncertainties with this approach for calculating EFs and ERs as the ratios derived this way represent the
average plume composition and ignore the spatial heterogeneity in wildfire plumes (Liu et al., 2022;
Decker et al., 2021; Peng et al., 2020; Garofalo et al., 2019), chemical transformation processes, and can
also be affected by changing background levels.

# 257 **2.5 Emissions Uncertainties**

258 There is the potential for inherent uncertainties using a plume integration method for calculating 259 EFs and ERs as the ratios derived this way represent the average plume composition and ignore the 260 spatial heterogeneity in wildfire plumes (Palm et al., 2021; Decker et al., 2021; Garofalo et al., 2019), 261 chemical transformation processes, and can also be affected by changing background levels. Pollutants 262 released by wildfires can be influenced by photochemical and physical changes that may take place 263 between the time of emission and the time of measurement, particularly for more reactive compounds 264 (e.g. Palm et al., 2021; Lindaas et al., 2020; Peng et al., 2020; Akagi et al., 2011). Although controlled 265 laboratory studies are well suited to examine direct emissions with minimal aging, they cannot reproduce 266 realistic burning conditions. Field measurements are critical to understand emissions that are impacted by 267 factors such as complex burning dynamics, fuel moisture, temperature and winds (Andreae 2019). 268 Recognizing the challenges of measuring primary emissions by aircraft, at 10 km (<1 hr) away from the 269 fire source, Screen 1 measurements represent some of the freshest emissions measured under wildfire 270 conditions, thus providing best estimates of initial conditions.

Uncertainties in the EFs and ERs are estimated by summing in quadrature the standard error of
the average EF (or ER) and the propagated measurement uncertainties. The standard error is used as

273 description of the uncertainty on the average EF (and ER) characterizing repeated transects across the SP 274 and NP for a total of 20 min of in-plume sampling. The standard error is expected to at least partially 275 capture uncertainties associated with plume aging and vertical plume heterogeneity. As many compounds 276 exhibited significant in-plume enhancements above background levels, uncertainties in the integrated  $\Delta X$ , 277  $\Delta CO$  and  $\Delta TC$  values were assumed to be dominated by instrumental (measurement) uncertainties (Table 278 S1, S2). Emissions are not reported for compounds where the average mixing ratios were within  $1\sigma$  of 279 the background average. The low and high I/SVOCs EFs (and ERs) are provided as estimates of their 280 uncertainties (as described in Sect. 2.3). The derivation of AWAS and cartridge EFs (and ERs) may have 281 potential limitations as they rely on a limited number of samples, with the potential of the AWAS discrete 282 samples capturing only part of a plume.

283

### 284 **2.6** Combustion efficiency

285 Combustion efficiency (CE) is a useful indicator of the relative proportion of flaming vs 286 smoldering stages of combustion which has a significant influence on the chemical composition of the 287 smoke (see SI Sect. 3.04 for further details). Flaming fires have CE > 0.90 (Yokelson et al., 1996) and 288 smoldering fires are typically ~0.8 with a range of 0.65 to 0.85 reported in the literature (Akagi et al., 289 2011; Yokelson et al., 2003). A modified combustion efficiency (MCE) is commonly calculated 290 assuming that CO<sub>2</sub>+CO adequately represents all of the fuel carbon that has been volatilized and detected 291 in ambient air. Here, as the TC in the plume was directly measured,  $\Delta$ TC was used in Eq. 43 to improve 292 on the estimation of the CE by accounting for all the sources of carbon.  $\Delta CO_2$  and  $\Delta TC$  in Eq. 43 are the 293 integrated, background-subtracted mixing ratios.

294

$$295 \quad CE = \frac{\Delta CO_2}{\Delta TC} \tag{43}$$

#### **3 Results and Discussion**

298

#### **3.1 Fire combustion state**

299 The plume-averaged CE for the SP (transects 1 to 4) was  $0.84\pm0.04$  and for the NP (transects 1 to 300 3)  $0.82\pm0.01$ . Transect 4 was excluded from the calculations for the NP because only a portion of the 301 plume was detectable at this altitude (Fig. 3). The derived CE indicates that the fire was predominantly in 302 a low intensity smoldering phase which is consistent with the satellite-derived fire intensities during the 303 flight (see Fig. 10) and ground-based meteorological observations, and may reflect some residual 304 smoldering combustion (RSC). It is estimated that emissions from this fire were sampled 14 hrs post 305 flaming. Other chemical measurements from this flight also support that the fire was largely smoldering 306 including the detection of elevated  $C_2H_4O_2^+$  (levoglucosan fragment from the AMS), low NO<sub>\*</sub> levels 307 (Lapina et al., 2008) (Fig. S32), and no detectable K<sup>+</sup> (from the AMS) (Lee et al., 2010). Significant 308 spatial variability in the concentrations of many of the measured species were observed closest to the fire 309 source, while the plumes became more well-mixed as they were transported downwind (Fig. S63). This 310 highlights the complexities of assessing wildfire combustion processes (Ward and Radke, 1993), and in 311 particular, boreal forests have been observed to exhibit greater variability in combustion efficiencies than 312 for other vegetation types (Urbanski et al., 2009).

# 313 **3.2 General plume features**

314 Most pollutants were strongly concentrated in the fire plumes Table A1 shows mixing ratios (or 315 concentrations) and background levels of 193200 pollutants that were enhanced in the fire plumes. with 316 the exception of sThe quantification of this suite of compounds provides new and additional emission 317 estimates to those reported in Simpson et al. (2011) and compiled in Andreae (2019) for the boreal forest 318 ecosystem. Several sulphur-containing compounds and a few other VOCs were not detected (Table S6), 319 and although not part of the measurement suite in the present study, Simpson et al. (2011) did not observe 320 emissions of anthropogenic halocarbons from wildfires in the same boreal forest ecosystem. In Fig. 3, the 321 in-plume portions are highlighted by the grey vertical bars and the SP and NP are indicated as the aircraft 322 flew at increasing altitudes to complete five horizontal transects. The lowest 4 transects showed enhanced pollutant levels while the 5<sup>th</sup> transect (not shown) was predominantly above the height of the plumes. 323

324 Higher concentrations were generally observed in the SP compared to the NP, possibly because of some 325 plume dilution in the NP resulting from a change in wind direction prior to sampling. The SP and NP 326 were distinctly separated from each other, with pollutants typically dropping to background levels 327 between the plumes. NMOG<sub>T</sub> mixing ratios varied between background levels of  $\sim 100-375$  ppbv to near 328 10 ppmv in-plume. CO and acetonitrile, often used as tracers of biomass burning (e.g. Wiggins et al., 329 2021; Landis et al., 2018; Simpson et al., 2011; de Gouw et al., 2006), reached 6.6 ppmv and 20 ppbv, 330 respectively in the SP, while maximum OA concentrations reached  $276 \,\mu g \, m^3$ , above a background level 331 of  $\sim$ 912.5±0.83 µg m<sup>-3</sup>. OA was the largest contributor to particulate mass (PM) comprising over 90 % of 332 the measured submicron mass with remaining portion comprised of BC, NO<sub>3</sub>, NH<sub>4</sub>, and SO<sub>4</sub> (Fig. S64). 333 Integrated filter samples taken from the aircraft across Screen 1 also showed the presence of a diverse set 334 of functionalized particle-phase organic compounds (Ditto et al., 2021). 335 The most abundant reactive nitrogen compounds ( $N_r$ ) were in the forms of reduced nitrogen (85-79) 336 %) with NH<sub>3</sub> comprising 421.7 % of  $\Sigma N_r$  (Fig. 4) and substantially lower nitrogen oxides i.e. NO<sub>x</sub> < 1 337 ppbv. A large portion of unmeasured nitrogen containing compounds found in these plumes was likely 338 dominated by peroxyacetyl nitrate (PAN) (Liu et al., 2022). These observations are consistent with 339 emissions from smoldering fires (Burling et al., 2011; Goode et al., 2000; McMeeking et al., 2009; 340 Yokelson et al., 1996). Dominant proportions of reduced nitrogen in biomass burning emissions were 341 also reported previously (Lindaas et al., 2020; Burling et al., 2011; Yokelson et al., 1996). Nitrogen-342 containing organics were detected in the present study totalling 3.9 ppbv and 18 % of  $\Sigma N_r$  (Fig. 4), 343 however, other such compounds that were not included with the instrument suite used in this study were 344 also likely emitted. Such compounds could include organic nitrates, amines, amides, heterocyclic 345 compounds, nitriles and nitro compounds that have been found in biomass burning emissions (Roberts et 346 al., 2020; Lindaas et al., 2020; Andreae 2019; Koss et al., 2018; Tomaz et al., 2018; Stockwell et al., 347 2015). Alkyl nitrates have been identified in biomass burning emissions, but their contributions to total  $N_r$ 348 appeared to be small (Juncosa-Calahorrano et al., 2021; Roberts et al., 2020; Lindaas et al., 2020; 349 Simpson et al., 2011; Alvarado et al., 2010; Singh et al., 2010).

### 350 **3.3 Total carbon budget**

#### 351 3.3.1 NMOG chemical classes – PTRMS, CIMS, AWAS

352 In-plume mixing ratios and the relative contribution of individually measured NMOG species to 353 the sum of those species ( $\Sigma$ NMOG) are shown for 13 chemical classes in Fig. 5. (See Fig. S75 for 354 separate SP and NP chemical classes). The largest chemical classes include carbonyls (acids, aldehydes 355 and ketones), alcohols, hydrocarbons (alkanes, alkenes, alkynes), aromatics (including furans, phenol, 356 benzene and toluene), and nitriles. Hydrocarbons (i.e. C<sub>x</sub>H<sub>y</sub>, including some aromatics)- were responsible 357 for just over half of the  $\Sigma$ NMOG (52.853 %) (Fig. S86), with 27.229 % identified as alkenes such as 358 ethene, propadiene, and propene, 19.319 % alkanes, predominantly ethane, and 3.13 % alkynes, almost 359 entirely acetylene. Non-aromatic oxygenates accounted for an additional  $\frac{36.236}{9}$  % of the  $\Sigma$ NMOG with 360 roughly equal contributions ( $\frac{10.1 \text{ to } 11.09}{12}$  to 12 %) from acids, aldehydes and alcohols, and a smaller 361 fraction from ketones (4.85 %). Including other oxygenated compounds such as furanoids and 362 phenol/phenol derivatives, all oxygenates ( $C_xH_yO_z$ ) comprised 41.442 % (Fig. S86), of the  $\Sigma NMOG$ .

363 A similar range of compound classes has been observed in previous field and laboratory studies, 364 noting that the measured compound suite between studies varies to some extent. For example, Simpson 365 et al. (2011) found a similar distribution of compound classes with 57 measured NMOG species, based on 366 discrete canister samples, in boreal forest wildfires. In that study, oxygenates (non-aromatic) comprised a 367 smaller portion of NMOG (29 %) as major emitted species like acetaldehyde and acetic acid (Fig. 8) were 368 not included. For example, some hydrocarbons, like 1-butene, ethane, propane, and isobutene measured 369 in the present study were not included in Koss et al., (2018) results. Other studies have also found 370 oxygenates to be a large portion of NMOG emissions across multiple fuel types, including those similar 371 to the current study, ranging from 51 - 68 % (Permar et al., 2021; Koss et al., 2018; Gilman et al., 2015; 372 Akagi et al., 2011) with a range of 25 – 55 % reported in Hatch et al. (2017). Comparisons between 373 studies are influenced by differences in study measurement suites and variations in fuel composition. The 374 fraction of NMOG oxygenates in the present study (41.442 %) was closer to those reported in Hatch et al. 375 (2017) when only the most relevant fuel types of pine and spruce were considered (55 % and 43 %,

376 respectively). Similar to previous work (Koss et al., 2018, Stockwell et al., 2015; Hatch et al., 2015),
377 emissions of substituted oxygenates like furanoids (furans+derivatives) and phenolic compounds were
378 observed. Furanoids contributed 4 % of the ΣNMOG mostly due to furfural, furan and methyl furan
379 while phenolic compounds eg. guaiacol, methyl guaiacol, contributed 0.5 % of the ΣNMOG (Fig. S97).
380 Although their-these emissions were less abundant in the present study, they represent important OH
381 reactants (Coggon et al., 2019; Koss et al., 2018; Gilman et al., 2015) with phenols being implicated as
382 precursors to brown carbon formation in secondary organic aerosol (SOA) (Palm et al., 2020).

Biogenic emissions of terpenoids including isoprene, monoterpenes, carvone, sesquiterpenes, camphor/isomers and terpine-4-ol/cineole/isomers were elevated in the plumes collectively reaching ~2.4 ppbv, and contributing ~1-2 % to the  $\Sigma$ NMOGs (Fig. S97). Isoprene was ~70-66 % of these compounds with an additional 29-32 % from monoterpenes. Emissions of isoprene from biomass burning has been observed from a wide range of fuel types (Hatch et al., 2019). As isoprene is not stored by plants and the measurements were taken ~14 hrs post flaming, it was likely emitted as a combustion product.

389 In this study, furfural was the most abundant oxygenated aromatic compound and a factor of 5 390 times higher than that of phenol. , whereas Hatch et al. (2015) and Although Koss et al. (2018) found 391 that phenol and furfural emissions were slightly larger than that of furfural-similar for all-most fuels tested 392 in the laboratory, furfural emissions derived from multiple wildfires sampled in Permar et al. (2021) were 393 similar to those in the present study, and a factor of 1.6 higher for phenol. As phenol emissions are 394 associated with lignin pyrolysis (Stockwell et al., 2015; Simoneit et al., 1999), the lower emissions in the 395 current study could be because the lignin content in the fuel mixture was lower than fuels used in previous 396 laboratory studies or that most of the phenolic compounds were emitted during the earlier phases of the 397 fire. Several modelling studies have indicated that aromatics and terpenes are insufficient to explain SOA 398 formation in biomass burning plumes (e.g. Hodshire et al., 2019) suggesting the importance of inclusion 399 of other aromatic species such as phenolics and furanoid compounds. However, models typically do not 400 include reactions involving phenolic and furanoids species, especially substituted compounds like 401 furfural, guaiacol, and methyl guaiacol. Box model simulations have also shown that incorporation of

402 OH oxidation of furan, 2-methyfuran, 2,5-dimethylfuran, furfural, 5-methylfurfural, and guaiacol, leads to
403 10 % more O<sub>3</sub> formed (Coggon et al., 2019).

### 404 **3.3.2 Intermediate-volatility and semivolatile organic compounds (I/SVOCs)**

405 Offline analysis of cartridge samples showed a wider range of hydrocarbons and functionalized 406 gas-phase organic compounds not observed in the PTRMS, CIMS, and AWAS measurements, including 407 I/SVOC compounds in the wildfire plume. ERs (Table S7) for species containing carbon and, hydrogen, 408 and with either sulfur and or oxygen (i.e. CH (hydrocarbons), CHS<sub>1</sub> and CHO<sub>1</sub> type molecules)) 409 accounted for a sizeable fraction of carbon in the  $\frac{1}{10}$  to C<sub>25</sub> range. , with expected eAdditional 410 cContributions are expected from more highly functionalized organics in the gas (and particle) phase not 411 reflected in the CH, CHO<sub>1</sub>, and CHS<sub>1</sub> compound classes (e.g., gas-phase species with multiple oxygen 412 atoms like vanillic acid or acetovanillone, and gas-phase species containing combinations of oxygen and 413 nitrogen atoms (CHON) (Ditto et al., 2021; 2022)). ERs in the plume varied across the carbon number 414 range; in general, the highest ratios were observed for the complex mixture of hydrocarbons (i.e. CH 415 compounds) broadly peaking at  $C_{200}$ - $C_{235}$  in the SVOC range, with a larger contribution from  $C_{10}$ 416 compounds including monoterpenes. By comparison, the complex mixture of CHO<sub>1</sub> compounds was 417 slightly lower in abundance than CH with contributions from  $C_{10}$  monoterpenoid emissions or 418 monoterpene oxidation products. CHS<sub>1</sub> IVOC-SVOCs were the lowest abundance species quantified. 419  $CHN_1$  compounds represent another observed contributor of IVOCs-SVOCs; the sum of all  $CHN_1$ 420 compound ion abundances was two orders of magnitude smaller than the sum of all CHO<sub>1</sub> species. We 421 note that for CHN<sub>1</sub>, this qualitative comparison is in terms of ion abundances only, given a lack of 422 appropriate standards to calibrate for the mass spectrometer's response to the complex mixture of reduced 423 nitrogen-containing I/SVOCs. 424 EFs were estimated for CH, CHO<sub>1</sub>, and CHS<sub>1</sub>-I/SVOCs based on Table S7 ERs (to CO) and the

425 average EF of CO (115.7  $\pm$  7.5 g kg<sup>-1</sup>, Appendix A). It was not possible to directly calculate EFs due to 426 the lack of a background sample upwind of the fire. EFs were estimated to be 1.4 $\pm$ 0.037 – 2.4 $\pm$ 0.063 g 427 kg<sup>-1</sup> for CH, 0.81 $\pm$ 0.078 – 0.81 $\pm$ 0.079 g kg<sup>-1</sup> for CHO<sub>1</sub>, and 0.21  $\pm$ 0.0033 – 0.22 $\pm$ 0.0060 g kg<sup>-1</sup> for CHS<sub>1</sub>

species, for a total EF of  $2.4\pm0.12 - 3.5\pm0.15$  g kg<sup>-1</sup> (Table A1). EFs were estimated to be  $1.6\pm0.04$  g kg<sup>-1</sup> 428 429 for CH, 0.9±0.09 g kg<sup>+</sup> for CHO<sub>1</sub>, and 0.1±0.003 g kg<sup>+</sup> for CHS<sub>1</sub> species, for a total EF of 2.6±0.14 g kg<sup>+</sup> 430 (Table A1). Here, the uncertainty represents measurement uncertainty associated with the conversion 431 from signal to mass, and the reported ranges show lower and upper limit EF values that account for a 432 contaminated background and that assume no background concentrations, respectively (as described 433 above). These estimates accounted for  $C_{11}$ - $C_{25}$  species and focused on I/SVOCs to avoid double counting 434 the monoterpenes and  $C_{10}$  monoterpenoid species, as they were already accounted for in the PTRMS data. 435 It is noted that the concentrations estimated for the cartridge samples may be sensitive to variations in 436 sampling efficiency within the under-wing sampling pod across  $C_{10}$ - $C_{25}$  though these effects are expected 437 to be minimal for the adsorbent tubes used in this study (Ditto et al., 2021; Sheu et al. 2018). These 438 emission estimates expanded the characterized spectrum of organic species to include IVOC/SVOCs in 439 boreal forest fire emissions, which until now, had only been available from laboratory measurements 440 (Hatch et al., 2018). However, the observed emissions of the complex mixture of hydrocarbons and 441 functionalized species may include contributions from the re-volatilization of compounds previously 442 emitted from upwind oil sands operations and deposited in the forest ecosystem, as noted in Ditto et al. 443 (2021).

444

445 **3.3.3 Accounting for the observed carbon** 

446 Measurements of TC, along with the speciated measurements from the PTRMS, CIMS, AWAS 447 and cartridges, provided a unique opportunity to reconcile the TC budget in a wildfire. Fig. 6 shows the 448 TC partitioning based on derived EFs (Sect. 3.5); overlapping compounds from the individual 449 measurement methods were handled as described in SI Sect. 12.1.4. The total EF for all carbon-450 containing compounds was 1652 g-C kg<sup>-1</sup> and, as expected, CO<sub>2</sub> was the dominant contributor comprising 451 >90 % of TC. CO contributed 7.0 % followed by a contribution from NMOG<sub>T</sub> of 1.9 % with even 452 smaller contributions observed from CH<sub>4</sub> (0.5 %) followed by OC and BC (not shown) at <0.5 %. The 453 magnified pie chart (right side) displays the two magnified pie charts (right side), representing the low

454 and high I/SVOC EF estimates, show the percent breakdown of the measured NMOGs, and the remaining 455 unidentified portion of NMOG<sub>T</sub>. The EF values (g C kg<sup>-1</sup>) are identified in the box below. The  $\Sigma$ NMOG EFs (for PTRMS+CIMS+AWAS measurements), totalling 14.413.6±3.20.9 g C kg<sup>-1</sup>, which accounted 456 457 for  $46.243\pm3$  % of the NMOG<sub>T</sub> EF of  $31.2\pm4.73.8$  g C kg<sup>-1</sup> (refer to Fig. S108 for the individual SP and 458 NP breakdowns). The  $\Sigma$ NMOG uncertainties were estimated by summing in quadrature the individual 459 compound EF uncertainties for the SP and NP separately, with these uncertainties subsequently summed 460 in quadrature to derive the average  $\Sigma$ NMOG uncertainty (Fig. 6). The cartridge data showed the presence 461 of a range of larger molecular weight I/SVOC compounds between  $C_{10}$  and  $C_{25}$  representing with an additional 2.1 to 3.0  $3\pm0.08$  g C kg<sup>-1</sup> representing and 7.47 to 10 % of NMOG<sub>T</sub>. Together, all of the 462 463 speciated NMOG measurements in this study accounted for  $\frac{53.650\pm3}{53.650\pm3}$  % to  $53\pm3$  % of NMOG<sub>T</sub>. The 464 remaining carbon mass was unidentified comprising  $46.447\pm15$  % to  $50\pm15$  % of NMOG<sub>T</sub>. Despite using 465 four state-of-the-art measurement techniques resulting in an extensive measurement suite, almost half of 466 NMOG<sub>T</sub> remained unidentified. This is consistent with previous work estimating  $\sim$ 50 % of NMOG<sub>T</sub> by 467 mass as unidentified (Akagi et al., 2011). It is noted, however, that the magnitude of the unidentified 468 portion is partly affected by uncertainties in the speciated measurements. For example, many of the 469 'calculated' PTRMS compounds are uncertain by an estimated factor of ~2 (SI Sect. 21.1.1, Table S1). 470 Nevertheless, a portion of the unidentified species likely consisted of challenging-to-measure-VOCs and 471 larger I/SVOCs that were highly functionalized or contained molecular features like reduced nitrogen 472 groups (e.g. amines) that have been observed in the gas and particle phase at various sites (Ditto et al., 473 2020; Ditto et al., 2022). While a complex mixture of I/SVOCs were observed from this fire (Table S7), 474 it is likely that other functionalized gas phase species containing nitrogen and/or multiple oxygens (e.g. 475 CHO<sub>>1</sub>, CHON, CHN) were also emitted, similar to particle phase observations in the fire plume via 476 tandem MS in Ditto et al. (2021). The presence of I/SVOCs in biomass burning emissions has been 477 previously observed in laboratory experiments (e.g. Koss et al., 2018; Hatch et al., 2018; Hatch et al., 478 2017; Bruns et al., 2016) with smoldering more likely to emit a higher fraction of compounds with low 479 volatility than higher temperature processes (Koss et al., 2018). The unidentified portion may also have

been comprised of nitrogen-containing organics (Sect. 3.1). Studies that included measurements of a
larger range of nitrogen-containing organics in biomass burning emissions estimated that they comprised
< 5-6 % of the total nitrogen budget (Lindaas et al. 2020; Gilman et al., 2015), and thus, an even smaller</li>
fraction of NMOG<sub>T</sub>. Advancing analytical techniques to expand the suite of NMOG speciation will
enable further reconciliation of the TC budget which is important for assessing secondary formation
processes in the atmosphere.

486

### 5 3.3.4 Volatility distribution of NMOG

487 Volatility distributions can help track the full range of organic species to assess their partitioning 488 between the condensed and gas phases (Donahue et al., 2011). Fig. 7 shows the fractional sum of all NMOG EFs within each volatility bin in terms of saturation concentration ranges ( $\log_{10}C_0$ ,  $\mu g m^{-3}$ ) for the 489 490 low I/SVOC EF estimate. C<sub>o</sub> values were estimated using the parameterization developed by Li et al. 491 (2016). NMOG emissions from this fire spanned a large range of volatilities from  $\log_{10}C_0$  of -2 to 10 µg 492  $m^{-3}$  across SVOCs to VOCs categories. The bin-averaged O/C ratio based on the measurements increased 493 with reduced volatility reflecting the presence of compounds with additional oxygen-containing 494 functional groups. The highest fraction of emissions was present as VOCs with  $\frac{63.381}{63.381}$  % having  $\log_{10}C_{o}$ 495  $> 6 \mu g m^{-3}$ , and 11.69 % as IVOCs having  $4 < \log_{10}C_0 \mu g m^{-3} < 6 \mu g m^{-3}$  and 107.9 % as SVOCs having 496  $\log_{10}C_0 < 3 \ \mu g \ m^{-3}$ . These results align with laboratory studies showing that oxygenates comprised more 497 than > 75 % of IVOCs across a range of biomass types with IVOCs accounting for ~11 % of the  $\Sigma$ NMOG 498 (Hatch et al.; 2018). Fig. 7 encompasses the range of volatilities based on all the identified NMOGs in 499 this study that is expected to represent initial emission conditions for modelling downwind chemistry. 500 However, improved speciation, particularly of lower volatility compounds, are is needed to further 501 expand the range of volatilities and advance knowledge in gas to particle partitioning processes. 502

### 503 **3.4 Emission factors and comparisons with other studies**

504 Emission factors (EFs) (and emission ratios (ERs)) in this study are derived for 250-193 505 compounds from 15 instruments of which 228-173 are NMOG species (Table A1). This dataset

506 represents the most extensive range of field-based EFs ever determined for a wildfire in the boreal forest 507 ecosystem. In Fig. 8 average EFs are shown for compounds grouped by a) particles, b) gas-phase 508 inorganics, and c) gas-phase organics. Separate EFs and ERs for the SP and NP are shown in the SI 509 (Figs. S119 to S134). In Fig. 9a-c, EFs are compared with those from other relevant studies. Fig. 9a 510 shows a comparison with boreal forest field measurements largely taken from a compilation by Andreae 511 (2019) referred to as BFF19, as well as values from Akagi et al. (2011) and Liu et al. (2017). This results 512 in a comparison for 50 compounds (35 organics and 15 inorganics/particulate species) with the largest 513 suite of EFs from one study conducted in a similar boreal region as the present study (Simpson et al., 514 2011). EFs are also compared with laboratory-derived EFs for lodgepole pine Koss et al. (2018; referred 515 to as LAB18) (Fig. 9b), a dominant fuelsimilar fuel type in the current study, with a total of 99 NMOGs 516 and 3 inorganics in common. In Fig. 9c, EFs are compared with those recently reported in Permar et al. 517 (2021) (referred to as TFF21) based on aircraft measurements of temperate forest wildfires in areas 518 mostly dominated by pine, fir and spruce trees, which provides the closest suitable comparison with 519 similar speciated NMOGs under wildfire conditions. Comparisons include 111 NMOGs, and 4 520 inorganics/black carbon. While the Permar et al. (2021) study was conducted in a temperate forest 521 region, it was at high elevation locations with similar vegetation types as the current study. 522 523 **3.4.1 Particle species** The PM<sub>1</sub> EF (6.8 $\pm$ 0.8 $\pm$ 1.1 g kg<sup>-1</sup>) represents the total of all particle component

524 species as measured by the AMS. The PM<sub>1</sub> EF of  $6.8\pm1.1$  g kg<sup>-1</sup> (Fig. 8a) (accounting for estimated mass 525 differences due to particle diameters (SI Sect. 1.1.2)) falls in the lower end of the large range previously 526 observed for boreal forest wildfires (18.7 $\pm$ 15.9 g kg<sup>-1</sup>; Fig. 9b). The few PM EFs for BFF19 (n=5) over a 527 limited range of MCEs (i.e. 0.89 to 0.93) shows significant variability consistent with previous work 528 (Jolleys et al., 2015; Akagi et al., 2011; Cubison et al., 2011; Hosseini et al., 2013). OA, accounting for 529 90 % of PM<sub>1</sub>, has the largest EF, accounting for 90 % of PM<sub>1</sub>, with comparatively lower EFs for  $PNO_3$ , 530 **#**BC, **p**NH<sub>4</sub>, and **p**SO<sub>4</sub> (Fig. 8a, Fig. S64). This reflects the dominant particle-phase organic carbon 531 content of the burned fuel and correspondingly lower fractions of nitrogen and sulphur-containing

532 compounds. Similar high organic fractions have been previously observed in biomass burning emissions 533 (Liu et al., 2017; May et al., 2014; Hecobian et al., 2011). ERs similarly highlight the dominant OA 534 emissions. Although tThe magnitude of EFs and ERs are generally similar between the SP and NP are 535 within their derived uncertainties, the ERs showed differences by up to 70 % for NH<sub>4</sub> (Fig. S12) 536 suggesting some differences in photochemistry between the two plumes. EFs and ERs for chemically-537 speciated particle species derived in this study represent the first such measurements under boreal forest 538 wildfire conditions. In Fig. 9a, EFs for chemically speciated compounds are not found in BFF19 (except 539 BC), but when compared with available values for U.S. temperate forest wildfires (Liu et al., 2017) are 540 found to be lower for OA (Fig. 9a), SO<sub>4</sub>, NO<sub>3</sub> and NH<sub>4</sub> by factors of 32.7, 5.0, 5.3, and 3.04, respectively. 541 Although differences in fuel type burned between the present study (mature Jack pine, boreal spruce, 542 boreal mixed-wood) and Liu et al. (2017) (mixed conifer, grass, brush and chaparral) may influence the 543 chemical composition of emissions, these large differences suggest the importance of other factors in 544 controlling OA emissions. The lower OA emissions under smoldering conditions in the current study 545 compared to Liu et al. (2017) with higher combustion efficiencies (0.877 to 0.935) conflicts with some 546 findings showing increased OA emissions with lower fire intensities (Liu et al., 2017, Burling et al., 547 2011). However, the relationship between  $EF_{OA}$  and combustion efficiency can be impacted by multiple 548 factors such as OA loading, gas-particle partitioning related to dilution, and fuel moisture content (May et 549 al., 2014). The EF<sub>OA</sub> in the current study ( $6.6\pm2.6 \text{ g kg}^{-1}$ ) lies in the range of EF<sub>OA</sub> reported for prescribed 550 burns across three temperate ecosystems  $(2.8\pm1.6 \text{ to } 11.2\pm2.7 \text{ g kg}^{-1})$  (May et al., 2014). This may imply 551 that the low intensity, surface, smoldering wildfire conditions in the present study (Sect. 3.1) may be 552 similar to prescribed burn conditions which are typically low intensity fires that are restricted to the forest 553 floor and understory, and conducted under controlled and consistent meteorological and fuel moisture 554 conditions (Yokelson et al., 2013; Carter and Foster, 2004). The lower iInorganic particulate-PM 555 emissions, however, are likely more dependent in the present study may reflect differences in on fuel 556 elemental composition between temperate and boreal forest ecosystems than combustion efficiency (Liu et 557 al., 2017). Differences in fuel composition between boreal and temperate forest ecosystems areis inferred

558 through comparisons of NO<sub>x</sub> and SO<sub>2</sub> emissions. For example, the average NO<sub>x</sub> and SO<sub>2</sub> EFs for boreal 559 forests, are lower than the average EFs for temperate forests by factors of 2.5 and 3.0, respectively. The 560 lower  $NO_x$  and  $SO_2$  emissions from boreal vs temperate forest wildfires are likely reflective of the 561 reduced S and N content in boreal biomass (Bond-Lamberty et al., 2006) relative to conifer (Misel, 2012) 562 fuels in the western U.S., as well as the possible influence of lower anthropogenic sources of nitrogen and 563 sulphur atmospheric deposition in boreal forests (Jia et al., 2016). The PM<sub>1</sub> EF of  $6.85\pm1.09$  g kg<sup>-1</sup> 564 derived in the present study is a factor of 2.8 lower than the PM<sub>2.5</sub> EF of 18.76±15.90 g kg<sup>-1</sup> that is 565 available for BFF19 (Fig. 9b). The lower PM emissions in the present study, despite accounting for 566 particle diameter differences (Sect. 2.1.2), - PM EFs for BFF19 (n=5) over a limited range of MCEs (i.e. 567 0.89 to 0.93) show significant variability. is somewhat surprising given emissions of PM are typically 568 higher from smoldering compared to flaming fires (Liu et al., 2017; Akagi et al., 2012). However, there 569 are few PM EFs for BFF19 (n=5) over a limited range of MCEs (i.e. 0.89 to 0.93) showing significant 570 variability. The PM<sub>1</sub> EF derived in the present study falls within the range previously observed for boreal 571 forest wildfires and underscores the significant variability in PM emissions.

572

573 **3.4.2 Gas-phase inorganic species** The largest average EFs for inorganic gases (Fig. 8b; separate NP 574 and SP Fig. S11) were from reduced nitrogen compounds dominated by  $NH_3$  (0.63±0.149 g kg<sup>-1</sup>) and 575 followed by HCN  $(0.31\pm0.07 \text{ g kg}^{-1})$ , with lower EFs for oxidized nitrogen compounds such as NO<sub>2</sub> 576  $(0.151\pm0.0437 \text{ g kg}^{-1})$  and HONO  $(0.021\pm0.012 \text{ g kg}^{-1})$ . This is consistent with previous work 577 identifying elevated emissions of NH<sub>3</sub> and HCN during smoldering conditions, whereas emissions of 578 HONO and NO<sub>x</sub> are primarily associated with flaming combustion (e.g. Roberts et al., 2020; Akagi et al., 579 2013; Yokelson et al., 1997; Griffith et al., 1991). The EFs for CO<sub>2</sub> and CO from the present study are 580 very close comparable within uncertainties of to that previously reported for BFF19 (Table A1). However, 581 EFs for most other gaseous inorganic species were lower than the BFF19 EF average including  $NH_{3,r}$ 582 HONO,  $\frac{SO_2}{(n=2)}$  and  $NO_x$  (n=11), by factors of 4.03.9, 2041, 4.7 and 7.114.9, respectively-(Fig. 583 9a).(Fig. 9a).- There are only a limited number of studies reporting EFs for these compounds in the

584 BFF19 category. For example, the HONO EF can only be compared with one other BFF19 study, but is 585 also lower compared to LAB18 (Fig. 9b). There are also only 4 previously reported BFF19 EFs for  $NH_3$ 586  $(2.56\pm1.87 \text{ g kg}^{-1})$  showing a large range of values. Although these comparisons are limited by the few 587 reported values in the literature, -the differences indicateing a strong sensitivity towards factors like fire 588 intensity, -and-chemical reactivity, fuel type and moisture, and meteorology. In contrast, EFs for HCN 589 derived in the current study  $(0.31\pm0.0728 \text{ g kg}^{-1})$  compare fairly well within the range of BFF19, 590 LAB18 and TFF21 values (0.28±0.06 to 0.53±0.30 g kg<sup>-1</sup>), (Figs 9a, b, c, respectively) and does not vary 591 widely suggesting that HCN may be less sensitive to burning characteristics. HCN is of concern due to 592 its impacts on human health particularly since biomass burning emissions are responsible for the majority 593 of the global HCN (Moussa et al., 2016 and references therein).

594

595 **3.4.3 Gas-phase organic species** In Fig. 8c, the top 25 average EFs for gas-phase organic species are 596 shown in decreasing order of magnitude. The most abundant emissions were from the lower molecular 597 weight compounds; such trends are generally in agreement with previous field-based measurements for a 598 range of fuel types (e.g. Permar et al., 2021; Andreae, 2019; Liu et al., 2017; Simpson et al., 2011; 599 Urbanski et al., 2009). Excluding CH<sub>4</sub>, the largest EFs were associated with methanol, followed by 600 ethene, ethane, acetic acid,  $C_5$  oxo-carboxylic acids, acetaldehyde, formaldehyde, and acetone ranging from  $1.9\pm0.45$  g kg<sup>-1</sup> to  $0.82\pm0.22$  g kg<sup>-1</sup> for these compounds. Noting some variations related to 601 602 differences in measurement methods, other studies have identified many of these same species as 603 dominating biomass burning emissions (e.g. Permar et al., 2021; Simpson et al., 2011; Akagi et al., 2011). 604 For example, Simpson et al. (2011) found that 5 of the same compounds in the present study including 605 formaldehyde, methanol, ethene, ethane and acetone were in the top 10 NMOG EFs from aircraft-based 606 measurements made of boreal forest wildfires in northern Saskatchewan, Canada, and within ~300 km of 607 the current study. In the present study, the top 24 NMOG compounds accounted for just over half (57 608  $\frac{149}{100}$  % of the  $\Sigma$ NMOG by total molecular mass with lower lower emissions from the remaining 149

609 measured compounds. In western U.S. wildfires, small emissions from 151 species were found to 610 account for almost half of  $\Sigma$ NMOG (Permar et al., 2021).

EFs for the NP and SP generally agreed within their uncertainties with larger differences for some of the more reactive species like isoprene, monoterpenes, and furan. For example, the SP EF for isoprene was a factor of 3.4 lower than that for the NP ( $0.64\pm0.34$  g kg<sup>-1</sup>) (Fig. S13). Although the reasons for these differences are not yet known, observations of higher O<sub>3</sub> in the SP ( $52.4\pm3.0$  ppbv) compared to the NP ( $44.7\pm3.6$  ppbv) suggest the influence of higher oxidant chemistry in the SP emissions compared to the NP.

617 To compare the total NMOG derived in the present study with those from previous studies that 618 typically sum up their speciated measurements i.e.  $\Sigma NMOG$ , estimates were made using two methods: 1. 619 increasing the  $\Sigma$ NMOG to account for the unidentified portion of NMOG<sub>T</sub>; and 2. adjusting the NMOG<sub>T</sub> 620 to reflect the total molecular mass (not just the carbon portion). For method 1, the  $\Sigma NMOG EF$ 621 (including the I/SVOCs) in this study  $(\frac{25.8\pm3.2}{24.5\pm1.6} \text{ to } 25.6\pm1.6 \text{ g kg}^{-1})$  was increased by 46.450 and 622 47 % (Fig. 6), respectively, equalling  $\frac{37.8}{36.8\pm11.3}$  to  $37.6\pm12.2$  g kg<sup>-1</sup>. This estimate assumes that the 623 carbon distribution is the same as the identified, speciated measurements. For method 2, based on the 624 speciated measurements, the average molecular mass was 100 g mol<sup>-1</sup> and the average carbon number was 625 6 resulting in  $\sim 28\pm 24$  % of the molecular fraction represented by atoms other than carbon. Adjusting the NMOG<sub>T</sub> of 31.2 $\pm$ 3.8 g C kg<sup>-1</sup> upwards by 28 $\pm$ 24 % to reflect the additional molecular mass results in a 626 627 NMOG<sub>T</sub> of 39.9±5.8 g kg<sup>-1</sup>. The resulting estimated NMOG<sub>T</sub> in this study of  $\frac{37.836.8\pm11.3}{37.836.8\pm11.3}$  to 39.9±5.8 g 628 kg<sup>-1</sup> lies between the estimated average of 58.7 g kg<sup>-1</sup> for the BFF19 (Fig. 9a) and those estimated from 629 the  $\Sigma$ NMOG EFs of 25.0 g kg<sup>-1</sup> (LAB18) (Fig. 9b), and 26.1±6.9 g kg<sup>-1</sup> (TFF21) (Fig. 9c) derived from 630 laboratory- and field-based studies (Table A1). In contrast to the current work, previous estimates of 631  $NMOG_T$  are likely to underestimate total NMOG emissions as they typically represent the sum of 632 measured species only. Some studies have attempted to account for NMOG<sub>T</sub> by including the sum of 633 measured plus estimates of 'unknown' portions of NMOGs (*Σ*NMOGs) (Permar et al., 2021; Koss et al., 634 2018; Stockwell et al., 2015; Gilman et al., 2015). The BFF19 EF was recently doubled from  $29.3 \pm 10.1$  g

 $kg^{-1}$  to 58.7 g kg<sup>-1</sup> to account for unidentified NMOGs where the ΣNMOGs were measured by FTIR, GC and PTRMS (Andreae, 2019; Akagi et al., 2011). These results support that doubling the ΣNMOG provides a reasonable estimate the NMOG<sub>T</sub>. It is noted, however, that the average BFF19 NMOG EF is ~1.5 times higher than that derived in the present study, however, this may reflect variability in NMOG emissions even within the same boreal biome.

640

641 Although it is known that acidic compounds are emitted from biomass burning, few studies have 642 quantified their emissions, particularly under field conditions (Andreae, 2019; Veres et al., 2010; 643 Yokelson et al., 2009; Goode et al.; 2000). In this study, EFs for 31-22 organic acidic compounds were 644 derived (Table A1) representing the most detailed set of organic acid EFs from biomass burning for any 645 ecosystem (Andreae, 2019). The largest EFs for these compounds include acetic acid, C5 oxo carboxylic 646 acids, C4 oxo-carboxylic acids, and pyruvic acid, all of which are found among the top 24 NMOGs (Fig. 647 8c). For those measurements that are available for comparison, EFs in the present study were lower for 648 formic acid and acetic acid compared to, than in BFF19, and were also lower than in LAB18, and TFF21, 649 ranging from factors of 1.7 to 8.8 (Figs. 9a, b, c, d). A total of nine Oorganic acids that were in common 650 with TFF21 and LAB18 (Table A1) hadve lower EFs, with the exception of pyruvic acid, which was 651 substantially higher (> factor of 37) in the present study. Differences in fuel type may be an important 652 factor in the variability of these comparisons. Based on laboratory experiments, Veres et al. (2010) found 653 a large range (factor of 5 to 13) of organic acid emissions with different fuel types suggesting that the 654 lignin content of the fuel could be a source of biomass burning organic acid emissions. Emissions for an 655 additional 1023 organic acids that have not previously been reported, as well as several inorganic acids 656 including nitrous acid, isocyanic acid, and peroxynitric acid, are included in Table A1. These acids, 657 representing 10.39 % of the  $\Sigma$ NMOGs (Fig. 5), are an important class of oxygenates as they can form 658 additional PM (Reid et al., 2005) and influence the hygroscopicity of smoke particles (Rogers et al., 1991; 659 Kotchenruther and Hobbs, 1998).

Isoprene and monoterpenes, with similar EFs  $\sim 0.410\pm0.190$  g kg<sup>-1</sup>, represented 167<sup>th</sup> and 1820<sup>th</sup>, 660 661 respectively, of the top 24 NMOG EFs in this study. Terpenes are known to be emitted from a range of 662 biomass burning fuels (Andreae, 2019 and references therein), but there have been few measurements in 663 boreal forest wildfire plumes (Simpson et al., 2011; Andreae, 2019). It is noted that PTRMS 664 measurements of IVOCs like sesquiterpenes likely represent lower limits as they tend to be easily lost to sample inlet lines due to their low volatility. The isoprene average EF of  $0.421\pm0.2610$  g kg<sup>-1</sup> was more 665 666 than a factor of 5 higher, while the monoterpenes EF,  $0.4139\pm0.034\cdot19$  g kg<sup>-1</sup>, was substantially lower 667 than the only reported EF for boreal forest wildfires (Simpson et al., 2011). The difference in EFs for 668 isoprene would be even greater if only the NP EF  $(0.64\pm0.34 \text{ g kg}^{-1})$  is compared (if it is assumed that 669 isoprene emissions were influenced by photochemical losses in the SP). As the present study and the 670 Simpson et al. (2011) study were conducted in similar locations (i.e. boreal forest region within ~300 km 671 of each other), with similar average MCEs, and comparable background levels, these differences are 672 likely driven by fire stage sampled combustion state, despite having similar study-averaged MCEs. The 673 majority of monoterpenes are stored in plant tissues (resin stores) for long periods of time, but isoprene is 674 synthesized and immediately released by plants, and can also be emitted as a combustion product 675 (Ciccioli et al., 2014; Akagi et al., 2013). Hatch et al. (2019) found that a wide range of terpenoids are 676 released across a variety of biomass types with variable emissions that were dependent on plant species, 677 and specifically related to their fuel resin stores. In the present study, monoterpenes may have 'boiled-678 off' through distillation processes in the early stages of the fire resulting in lower monoterpenes emissions 679 at the aircraft sampling time, ~14 hrs post-flaming. In contrast, the Simpson et al. (2011) study sampled 680 comparatively earlier and more intense fire stages where higher monoterpene emissions were likely 681 released from live or recently fallen trees that still contained significant resin stores. The monoterpenes 682 EF reported by Simpson et al. (2011) was likely even higher given only two monoterpenes were speciated 683 and emissions of other terpenes were likely (Hatch et al., 2019). Higher isoprene emissions in the present 684 study compared to Simpson et al. (2011) could be related to the comparatively larger smoldering 685 component. Although limited data exist on the release of isoprene as a function of fire intensity, negative

relationships between isoprene and MCE were observed in Australian temperate forest fires (Guérette etal., 2018) and wheat fields (Kumar et al., 2018).

688 Several furanoid compounds also exhibited significant emissions (Fig. 8c) including furfural, 689 furan, and methyl furan ranking 12<sup>th</sup>, 19<sup>th</sup>, and 22<sup>nd</sup> of the top 24 NMOG-organic compounds, 690 respectively. Emissions of furanoids have been observed for a wide range of fuel types (Hatch et al., 691 2017; Simpson et al., 2011). Fairly good aAgreement within uncertainties was found with BFF19 for furfural, and furan (Fig 9a). The EFs for furan  $(0.39\pm0.1920 \text{ g kg}^{-1})$  and furfural  $(0.65\pm0.31\text{-g kg}^{-1})$  were 692 693 also similar to that in LAB18 (Fig. 9b), and TFF21 (Fig. 9c), as well as other ecosystems (Andreae, 2019) 694 suggesting their emissions were relatively insensitive to fire intensity and fuel mixture. Overall, tThe 695 comparisons in Fig. 9 indicate that for the higher emitting species, the current results are fairly similar, 696 but for the lower emitting species, these results are lower than previous reported values. These 697 comparisons provide context for the emissions reported in the present study and moves towards improved 698 statistics to better constrain wildfire emissions. Additional factors are considered to explain variability in 699 emissions between this study and other reported values, as well as within this study (NP vs SP). 700 Differences and variability in burn conditions (e.g. fire intensity, winds, fuel density, flame dynamics, 701 fuel moisture) likely influence these comparisons; the Screen 1 measurements in the present study were 702 taken from 9-10 am LT when the fire was in a low intensity, smoldering state, while those in Permar et al. 703 (2021) and Simpson et al. (2011) took place during mid-day under active wildfire conditions. Aircraft 704 measurements in general have a higher probability of sampling variable burn conditions compared to 705 laboratory studies (Hodshire et al., 2019), and as such, aircraft-derived EFs are likely to reflect variability 706 for reactive species as speculated earlier with isoprene. Particularly for reactive species that can exhibit 707 complex variation across plumes, EFs (and ERs) derived by integrating across plumes can be biased low, 708 (Sect. 2.5; Peng et al., 2021; Decker et al., 2021). Also, EFs derived using TC in this study may result in 709 lower, albeit small, EFs compared to reported values that do not account for all the carbon (estimated to 710 be 1-2 % (Akagi et al., 2011)).

### 712 **3.5 Evaluation of emissions models**

714

#### 713 **3.5.1** Comparison of EFs with the model emissions speciation profile

715 EFs derived in the present study are compared with those that are currently incorporated into the 716 emissions component of the FireWork modelling system using the Forest Fire Emissions Prediction 717 System (CFFEPS). CFFEPS uses EFs allocated for 3 combustion states (flaming, smoldering and 718 residual) and for 8 species including lumped non-methane hydrocarbons (NMHC) based on United States 719 vegetation data compiled in Urbanski et al. (2014) (Table 3 in Chen et al., 2019). Fig. 9d (bolded 720 compounds) shows that the smoldering EFs in the present study were comparable for CO, -and-CH<sub>4</sub> and 721 **NMOG**,-but lower for PM<sub>1</sub> (PM<sub>2.5</sub>), NH<sub>3</sub>, SO<sub>2</sub> and NO<sub>x</sub> by factors of 3.4, 2.4, 6.6 and 17, respectively. In 722 the present study, additional mass between PM<sub>1</sub> and PM<sub>25</sub> accounted for only an additional 10 % of 723 aerosol mass (SI Sect. 2.1.2). The lower EFs for these species implies that the CFFEPS EFs would not 724 adequately capture their total emissions under smoldering conditions for the boreal fuel in the current 725 measurement study.

726 For incorporation into numerical air quality models, total organic gas (TOG=NMOG+CH<sub>4</sub>) 727 emissions are typically split into detailed chemical components using chemical mass speciation profiles, 728 and converted to lumped chemical mechanism species. In the FireWork modelling system, the 729 smoldering combustion TOG is split into components based on EPA's SPECIATEv4.5 profile (#95428) 730 (US EPA 2016, Urbanski et al.; 2014 - supplement Table A.2, Boreal Forest Duff/Organic soil). This 731 profile is ultimately compiled using laboratory data from Yokelson et al. (2013), Bertschi et al. (2003), 732 and Yokelson et al. (1997) based entirely on U.S. fuel types. EFs in the present study were found to be 733 generally lower than the laboratory-based EFs for 74 species in common ranging from factors of 1.7 to 734 8.5 including for monoterpenes, formic acid, phenol, and furan (Fig. 9d)<del>and acetonitrile (Fig. 9d)</del>. The 735 largest differences (factors of 49-57) were observed for sesquiterpenes, benzofuran, and naphthalene. A 736 few species including furfural, propane nitrile and ethyl styrene are comparable, while isoprene, and 737 pyruvic acid, acetylene and cyclohexene are notably higher by factors 2 to 5.3.

738 For a research version of the FireWork system, the component speciation is mapped to the 739 SAPRC-11 chemical mechanism species (Carter and Heo, 2013) with detailed oxygenated compounds 740 and aromatic species, largely to better represent SOA formation processes. For comparison with the 741 measurement derived speciation profile in this study, EFs were first mapped to SAPRC-11 species and then-normalized by the total identified mass species fraction without unknowns to obtain mass fractions 742 743 of relevant model mechanism species (Table S9). Comparing the normalized mass fractions for similar 744 mechanism species (Fig. S142) showed a substantially much-lower fractions of reactive alkaenes (ALK5) 745 and aromatics (ARO2) and a slightly higher acetic acid group (CCOOH) with an estimated 5 % in this 746 study compared to 28 % in the SPECIATEv4.5 wildfire smoldering profile. Mass fractions in this study 747 are notably higher for the ACYL, ETHE, and ISOP lumped model species by factors of 13, 7 and 51. The 748 mass fraction of CH<sub>4</sub> is also different with 2413 % of TOG in this study compared to 4 % from the 749 SAPRC 11SPECIATE4.5 profile. The measurement derived chemical speciation profile is expected to be 750 slightly different from the average speciation profile from EPA's SPECIATEv4.5 due to differences in 751 chemical species identification, fuel type, fire and measurement conditions, chemical species 752 identification and uncertainties on how measured compounds are mapped to lumped mechanism mapping 753 schemespecies. The emissions profile developed in the present study is considered a more can be used to 754 improve representative predictions of wildfire smoldering emissions profile specific to the wildfire characterization for the Canadian boreal forest-fuel. 755

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3.5.2 Linking aircraft and satellite observations to evaluate modelled emissions diurnal variability
Wildfires generally exhibit a diurnal cycle with fire intensities maximizing late afternoon and
diminishing at night having important implications for fire emissions (Chen et al., 2019). Evaluating
modelled emissions throughout the diurnal cycle with observations is a critical step in verifying smoke
predictions. Emissions models mostly parameterize diurnal fire emissions with prescribed profiles that
distribute daily total emissions to hourly. In CFFEPSs, a diurnal profile is applied to allocate daily burn
area to hourly intervals, with highest activity in the late afternoon. The actual fuel consumed, and thus,

764 hourly emissions, is then calculated with depth of burn estimates driven by hourly meteorology (Chen et 765 al., 2019). In Fig. 10, for the wildfire in the present study, the hourly CFFEPS-predicted emissions 766 (orange dots) for selected compounds are shown between 2018-06-24 17:00 UTC and 2018-06-25 21:00 767 UTC, spanning the aircraft sample time (red arrow at 15:00 UTC). After 21:00 UTC, the discrepancy 768 between the CFFEPS-predicted emissions and FRP increased as a result of rain that passed through the 769 area that is not considered in the model bottom-up emission estimates (not shown in the figure). The 770 burning phases are outlined in the figure where flaming (light pink background) is assumed to occur when 771 the atmospheric conditions alongside fire behaviour and emissions model outputs infer a fireline intensity 772 >4,000 kW m<sup>-1</sup> (Forestry Canada Fire Danger Rating Group, 1992), and a smoldering fire (blue 773 background) for intensity <4000 kW m<sup>-1</sup>. The fire intensity distinction between flaming and smoldering 774 roughly aligns with the observed minimum for this particular fire with the fire radiative power (FRP, grey 775 dots) retrieval from the GOES-16 satellite sensor of 500 MW where smoldering occurs <500 MW and 776 flaming for >500 MW. The 500 MW threshold over the approximately 1,700 ha of actively smoldering 777 area observed by overnight VIIRS thermal detections gives an estimated energy density of 0.29 MW ha<sup>-1</sup>. 778 This energy density threshold for smoldering < 0.29 MW ha<sup>-1</sup> found in this study is in agreement with FRP 779 per unit area corresponds with observed FRP for flaming combustion of >0.4 MW ha<sup>-4</sup> from lower 780 intensity flaming fires by O'Brien et al. (2015) who found flaming combustion at >0.4 MW ha<sup>-1</sup> for lower 781 intensity flaming fires and smoldering combustion at lower energy densities. The FRP represents the sum 782 over all hotspots of this fire for each 15-min observation period. Emission rates in metric tonnes per hour 783 (t h<sup>-1</sup>) were derived from selected aircraft measurements using a mass balance method that was designed 784 to estimate pollutant transfer rates through virtual screens using aircraft flight data (Gordon et al., 2015) 785 (see SI Methods). Emission rates were-and estimated to be  $29\pm2.1$  t h<sup>-1</sup> for PM<sub>1</sub>,  $433\pm26.7$  t h<sup>-1</sup> for CO, 786  $0.65\pm0.03$  t h<sup>-1</sup> for NO<sub>x</sub> (as NO), and  $2.7\pm0.16$  t h<sup>-1</sup> for NH<sub>3</sub> (red arrows). Emission rates were also 787 derived from satellite observations (black arrows) for CO, NO<sub>x</sub>, and NH<sub>3</sub>. Emissions of CO were 788 estimated using a flux method as described in Stockwell et al. (2021) using TROPOMI satellite 789 observations yielding  $1670\pm670$  t h<sup>-1</sup> at 19:06 UTC and  $4050\pm1620$  t h<sup>-1</sup> at 20:48 UTC. NO<sub>x</sub> emissions

790  $(9.1\pm3.4;$  scaled to t NO h<sup>-1</sup> at 19:06 UTC (not enough high-quality observations for the 20:48 UTC 791 overpass) were derived from the TROPOMI NO2 dataset using an Exponentially Modified Gaussian 792 approach (Griffin et al., 2021). NH<sub>3</sub> emission rates  $(5.6\pm3.9 \text{ th}^{-1})$  were derived from CRIS satellite 793 observations at the satellite overpass time of 19:00 UTC by applying a flux method (Adams et al., 2019). 794 The aircraft measurements were taken when the FRP was low reflecting a smoldering surface 795 fire. However, the satellite overpass occurred ~4 hrs later than the aircraft measurements close to the 796 FRP daily maximum, after which rain passed through the area. The CFFEPS model, exhibiting a 797 prescribed diurnal pattern, captures the increase in  $NO_x$  and  $NH_3$  emissions between that derived from the 798 aircraft and satellites transitioning from a smoldering to predominantly flaming fire; NO<sub>x</sub> emissions 799 increased by a factor >10, whereas the NH<sub>3</sub> emissions increased by a factor of approximately 2. This is in 800 agreement with recent laboratory measurements that found that the release of  $NO_x$  is favoured during the 801 flaming stage and the release of reduced forms of nitrogen, such as  $NH_3$ , is favoured during the 802 smoldering phase (Roberts et al., 2020) (also see Fig. 4). However, the CFFEPS CO emission rates do 803 not track the increase in CO emissions between the aircraft-derived value and the two TROPOMI values, 804 indicating that the CO EF for flaming is low in the model. This highlights the need to validate model 805 emission rates with measurements to adjust and update the EFs accordingly. 806 Using tThe aircraft- and satellite-derived emission rates for CO, NO<sub>x</sub> and NH<sub>3</sub> were each ratioed 807 relative to FRP (in units of t  $h^{-1}$  MW<sup>-1</sup>, referred to as R<sub>species/FRP</sub>) to represent the the-two end burning states 808 ie. smoldering and flaming conditions., estimates of total emissions from this fire were made for CO, 809 NO<sub>\*</sub> and NH<sub>3</sub>.-R<sub>species/FRP</sub> values were estimated for the flaming and smoldering phases of the fire and it 810 was assumed that flaming occurred for FRP >500 MW and smoldering for FRP < 500 MW. Total 811 emissions were estimated by integrating the GOES FRP over the period June 24, 20182018-06-24 17:00 812 UTC to June 252018 06 25 23:00 UTC (after which no more hot spots were detected by GOES and the 813 fire presumably extinguished), and applying the derived smoldering and flaming emission 814 ratioscoefficients. It was assumed that flaming occurred for FRP > 500 MW and smoldering for FRP < 815 500 MW. Emission rates coefficients were estimated with respect to the FRP for the flaming and

816 smoldering phases of the fire. The CO R<sub>species/FRP</sub> emission rates coefficients are-values were roughly 817 twice as large during smoldering compared to flaming. For the satellite emission estimates from the two 818 satellite overpasses during the flaming phase of the fire, the CO-R<sub>CO/FRP</sub> values emission rates are were 819 very similar and well-within the uncertainties (19:060 UTC  $R_{CO/FRP}ER_{CO} = 0.47 \pm 0.25 - 4.7$  t h<sup>-1</sup> MW<sup>-1</sup>; 820 20:4800 UTC  $R_{CO/FRP}ER_{CO} = 0.43\pm0.2343$  t h<sup>-1</sup> MW<sup>-1</sup>). The  $R_{NOx/FRP}$  valuecoefficientratio for NO<sub>x</sub> is also 821 twice as large for flaming compared to smoldering, and for NH<sub>3</sub>, the ratio R<sub>NH3/FRP</sub> value coefficient is ~5 822 times larger for smoldering than flaming. Total emissions were then estimated by integrating the GOES 823 FRP over the period 2018-06-24 17:00 UTC to 2018-06-25 23:00 UTC (after which no more hot spots 824 were detected by GOES and the fire presumably extinguished), and applying the derived smoldering and 825 flaming R<sub>species/FRP</sub> values. Assuming that the fire went out when GOES did not observe any hot spots, 826 total emissions for this fire of CO, NO<sub>x</sub> and NH<sub>3</sub> are estimated at  $\frac{21,808}{22,000\pm8700}$ ,  $104.1\pm42$ , and 827  $83.74\pm33$  tonnes, respectively. If the fire is assumed to have continued burning when GOES did not 828 detect any fire hot spots (between 22:00 - 04:00 UTC and 07:00 - 15:00 UTC, with an FRP of 150 MW 829 (~GOES detection limit; Roberts et al., 2015), the emissions increase to 24,000±96003,986, 106±43.4 and 830  $97.798\pm39$  tonnes, respectively, providing an upper limit of emissions. The combination of aircraft and 831 satellite-derived emission estimates for multiple species helps to obtain the diurnal variability of 832 emissions and to obtain more complete details on the emission information across different burning 833 stages.

# 834 **4. Summary and Implications**

This study provides detailed emissions information for boreal forest wildfires under a-smoldering combustion processconditions. Consistent with previous results, hHighly speciated airborne measurements showed a large diversity of chemical classes highlighting the complexity of emissions. Despite extensive speciation across a range of NMOG volatilities, a substantial portion of NMOG<sub>T</sub> remained unidentified (46.447±15 to 50±15 %) and is expected to be comprised of more highly functionalized VOCs and I/SVOCs. Although these compounds are challenging to measure, their characterization is necessary to more fully understand particle-gas partitioning processes related to the

842 formation of SOA. Methodological advancements to achieve higher time resolution speciated 843 measurements of I/SIVOCs would move towards further NMOG<sub>T</sub> closure and span a more complete 844 range of volatilities. A detailed suite of EFs that were derived in this study builds on previous work (e.g. 845 Simpson et al., 2011; Andreae 2019) and can be used to improve chemical speciation profiles that are 846 relevant for air quality modelling of boreal forest wildfires. Aircraft-derived emission estimates were 847 paired with those from satellite observations demonstrating their combined usefulness in assessing 848 modelled emissions diurnal variability. As satellite instrumentation and methodologies advance, linking 849 emissions derived from aircraft (and ground) observations for additional compounds will improve the 850 ability to simulate and predict the diurnal variation in wildfire emissions. 851 Although the measurements from this study provide a detailed characterization of a wildfire, tThe 852 results presented here represent only one smoldering boreal forest wildfire with limited in-plume 853 sampling times. Additional measurements are needed under a variety of fire conditions (combustion 854 state, fire stage, biomass mixtures, time of day, etc) in order to elucidate the major controlling factors and 855 improve statistical representation for constraining and modelling these sources. For example, 856 measurements are needed to assess dark chemistry reactions in biomass burning emissions which have 857 been shown to be important in the formation of OA (Kodros et al., 2020) and brown carbon (Palm et al.; 858 2020). In addition, reduced actinic flux associated with high particle loadings in biomass burning 859 emissions can influence plume chemistry (e.g. Juncosa-Calahorrano et al., 2021; Parrington et al., 2013). 860 The emissions information in this work can be used for will contribute to the evaluation and 861 improvements of models that are essential for reliable predictions of boreal forest wildfire pollutants and 862 their downwind chemistry. 863 864 865 866 Acknowledgements

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890	All data used in this publication are available upon request.
891	

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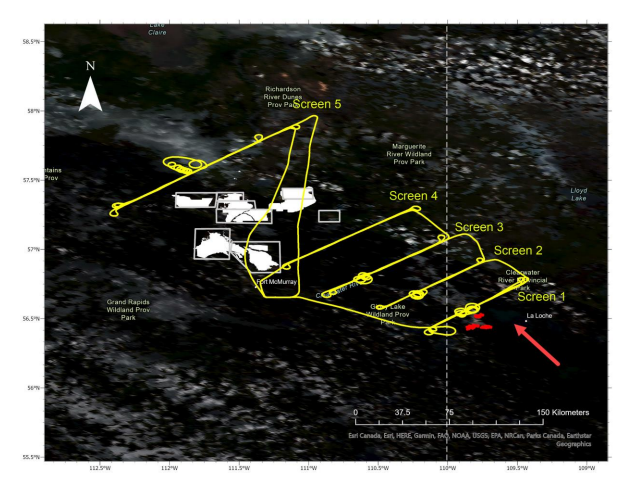


Figure 1. Corrected reflectance satellite image from the VIIRS spectroradiometer on the Suomi
NPP and NOAA-20 satellites taken on June 25, 2018. The fire hot spots for the wildfire of
interest are indicated by the red dots. Flight tracks were flown at Lagrangian distances

downwind of the wildfire. Multiple transects at varying altitudes perpendicular to the plume
 direction formed 5 virtual screens. Plume direction of travel is indicated by the large-red

1525 arrow. The location of the Alberta oil sands mining facilities are shown in white.

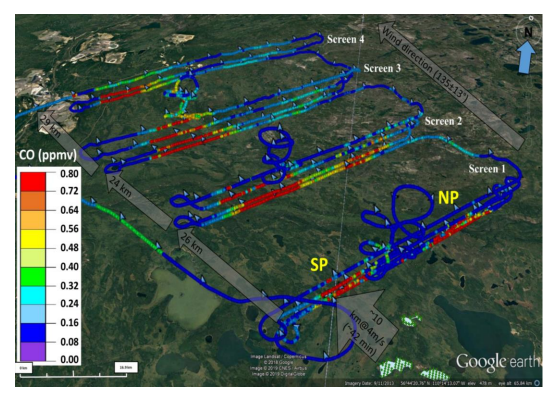
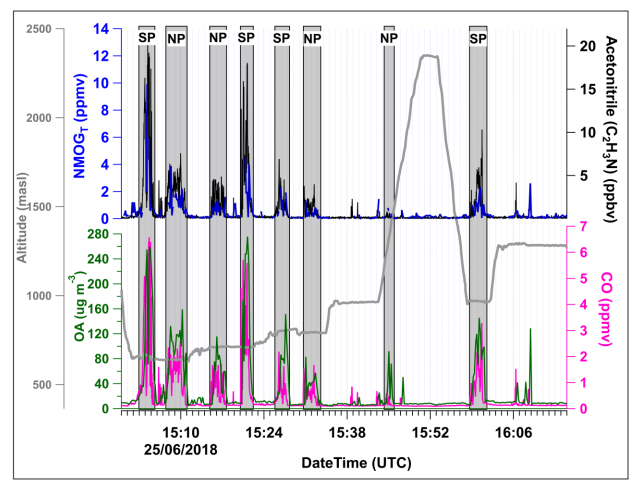


Figure 2. Flight tracks coloured by CO mixing ratio (ppmv) for Screens 1 to 4. The two plumes are identified as south plume (SP) and north plume (NP). The fire perimeter surrounding the detected MODIS-derived 'hot spots' on June 25, 2018 is shown in the green hatched area. The source of the NP is expected to be the same hot spots as the SP but ~ 30 min older; see SI-Sect.
2.2. The small blue arrows along the flight tracks indicate the aircraft measured wind direction

1534 with the average wind direction depicted with the large gray arrow. Distances between screens

- 1535 are shown in the grey arrows.
- 1536



1538 Figure 3. Time series of NMOGs (ppmv), acetonitrile (C<sub>2</sub>H<sub>3</sub>N; ppbv) and CO (ppmv), as well as OA concentrations (µg m<sup>-3</sup>) and altitude for Screen 1. The in-plume portions are indicated by the vertical grey bars. The aircraft flew back and forth across the plumes at increasing altitudes to complete five transects; a transect represents one pass across the SP and NP at the same altitude. 

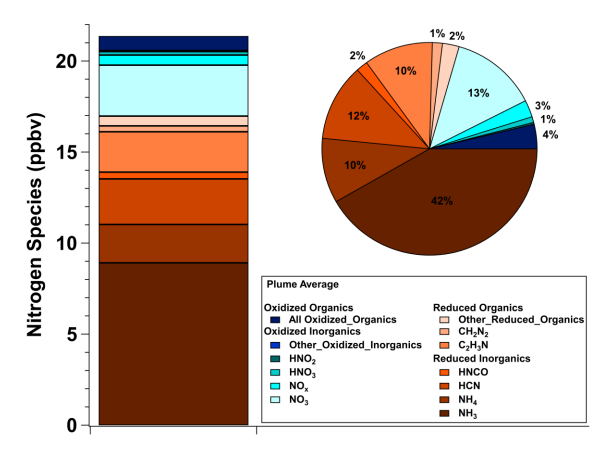
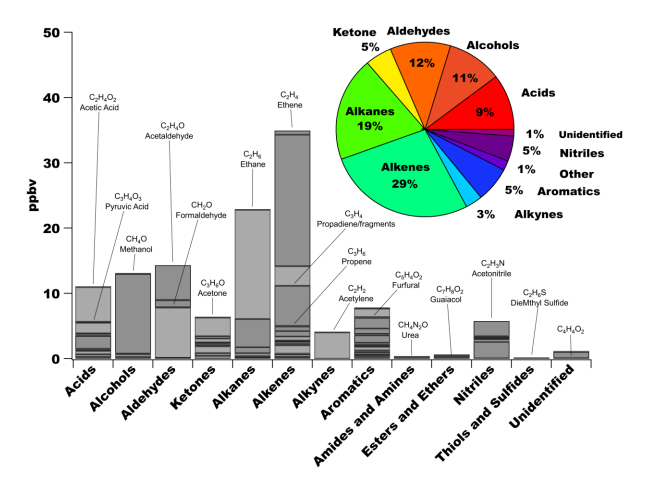




Figure 4. Background-subtracted average Screen 1 in-plume mixing ratios of measured gas- and particle-phase N-containing species ( $N_r$ ) and their fractional contribution to the total summed  $N_r$ species. The  $N_r$  species are grouped into categories of reduced inorganics, reduced organics, oxidized inorganics and oxidized organics with reduced species in shades of red and oxidized species in shades of blue.



1559 Figure 5. Background-subtracted average mixing ratios of individually measured NMOGs

shown for thirteen chemical classes. In some cases, compounds are double- (or triple-) counted if

they can be identified in more than one category. For example, phenol is an alcohol + an
aromatic; guaiacol is an alcohol + an ether + an aromatic. In the pie chart, the *Other* category

1562 includes amides, amines, ethers, thiols and sulfides. The unidentified category contains

1564 molecular formulas detected, but the compound(s) could not be identified.

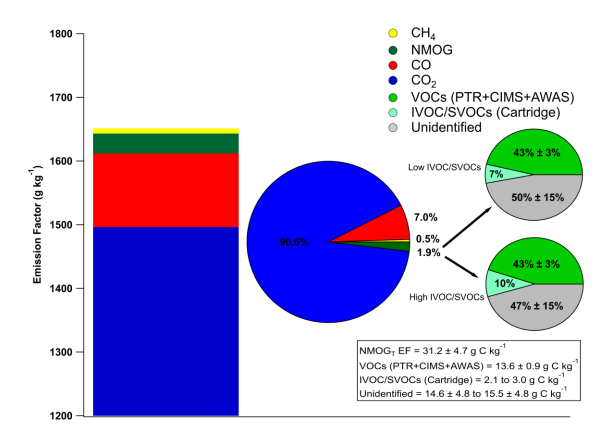


Figure 6. Total carbon (TC) partitioning based on EFs (carbon fraction). The bar chart shows the stacked EFs for carbon-containing compounds with the middle pie chart showing their percent contributions to the TC. The two magnified pie charts (right side), representing the low and high I/SVOC EF estimates, show the percent breakdown of the measured NMOGs and the remaining unidentified portion. The EF values (g C kg<sup>-1</sup>) are provided in the box. The pie chart on the right show the percent breakdown of the measured NMOGs with the remaining unidentified portion in terms of g C kg<sup>-1</sup>. Note that all the EFs shown in Table A1 were converted to g C kg<sup>-1</sup>. for this breakdown.

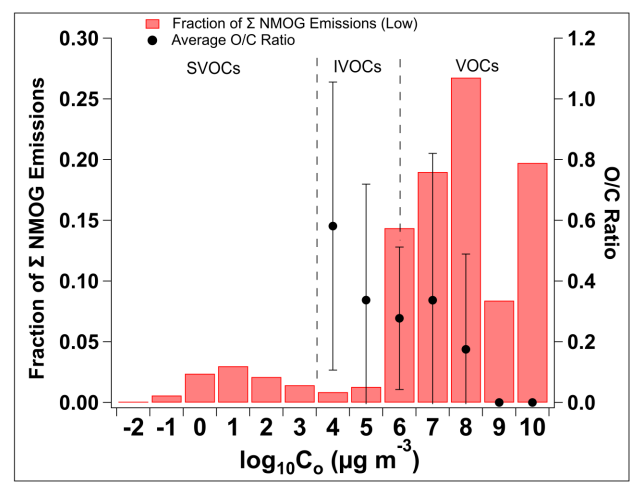


Figure 7. Fraction of total  $\Sigma$ NMOG emissions in each volatility bin, as well as the bin-averaged 1579 O/C ratio spanning VOCs, IVOCs and SVOCs. Data is included from PTRMS, CIMS, AWAS 1580 and cartridge measurements. The O/C ratio is derived for only the PTRMS, CIMS and AWAS measurements and the errors bars indicate the standard deviation of the average O/C ratio. 1581 1582

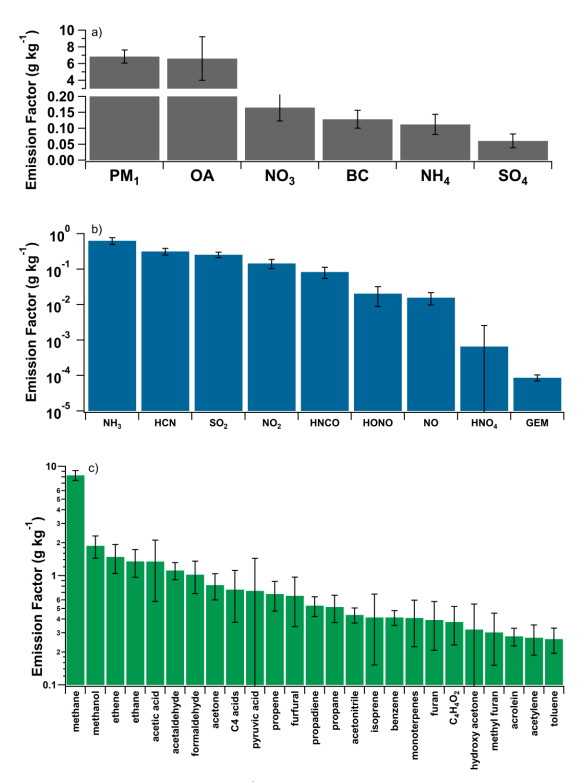
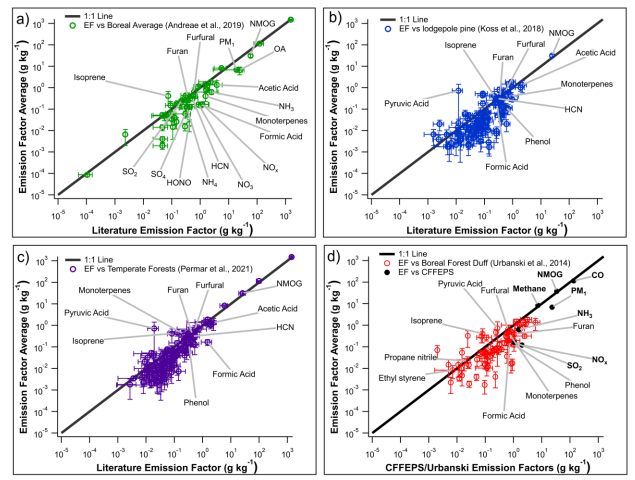




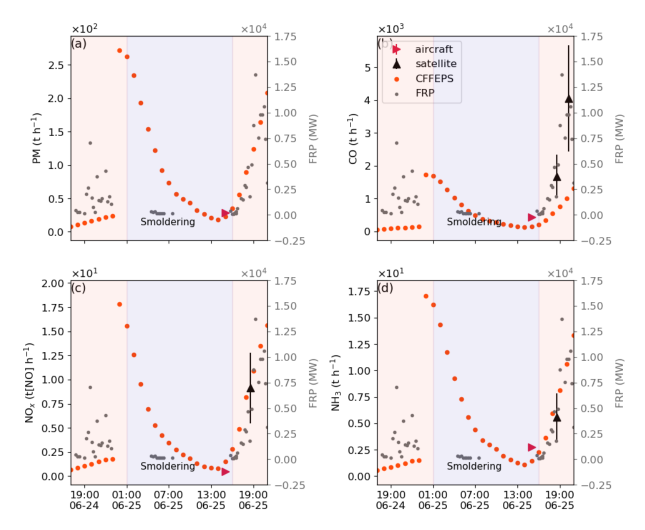
Figure 8. Average emission factors (g kg<sup>-1</sup>) of a) particle species; b) inorganic gas-phase species, and c) 1586 1587 the top 25 measured gas-phase organic species.  $C_{5} = C_{5} = C_{5} = C_{5} = C_{5} = C_{4} = C_{4}$ carboxylic acids; propadiene = fragments/propadiene; hydroxy acetone = hydroxy acetone/ ethyl formate. 1588 Organic species measurements are from the PTRMS, CIMS and AWAS.



1589

1590 Figure 9. Comparison of averaged emission factors with a) boreal forest field-based

- 1591 measurements (Andreae, 2019; Akagi et al., 2011; Liu et al., 2017), b) laboratory-based 1592 measurements of lodgepole pine (Koss et al., 2018), c) temperate forest field-based
- 1593 measurements (Permar et al., 2021), and d) those used in CFFEPS (Urbanski et al., 2014). See
- 1594 Table S8 for compound comparisons that don't have exact matches.
- 1595
- 1596
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- 1599



1602 Figure 10. Fire radiative power (FRP; in MW) from GOES-R (grey dots) and emissions from 1603 the CFFEPS model (orange dots) from 2018-06-24 17:00 UTC to 2018-06-25 21:00 UTC. Local 1604 time = UTC - 6 hrs. Aircraft-derived emission rates are shown for a) PM<sub>1</sub>, b) CO, c) NO<sub>x</sub> (as 1605 NO) and d) NH<sub>3</sub> (in t h<sup>-1</sup>; red arrow) at 15:00 UTC when the aircraft flew closest to the fire. The 1606 corresponding TROPOMI satellite-derived emission rates are also shown (in t h<sup>-1</sup>; black arrows). Note, the aircraft flight time occurred when the fire intensity reflected a surface, smoldering fire 1607 1608 and the satellite overpass time took place when the fire had transitioned to a crown (flaming) 1609 fire. The smoldering and flaming time periods are coloured in blue and pink, respectively. 1610

# 1611 **Table A1**

## 1612

1613 Table S7. Summary of in-plume and background average mixing ratios (or concentrations),

emission factors (g kg<sup>-1</sup>) (EF) and emission ratios (ppby ppmv<sup>-1</sup> except CO<sub>2</sub> which is in units of  $\frac{1}{2}$ 

1615 ppmv ppmv<sup>-1</sup>; particulates in  $\mu$ g m<sup>-3</sup> ppm<sup>-1</sup> and TGM-GEM in ng m<sup>-3</sup> ppmv<sup>-1</sup>) (ER) for the SP, 1616 NP, and the EF average of the two plumes. In-plume and background averages are in units of ug

 $m^{-3}$  for particulates, ppbv for gas-phase compounds, except GEM which is ng  $m^{-3}$ , and CO<sub>2</sub> is

1618 ppmv. Compounds are, grouped by particulate species, and inorganic and organic gas-phase

1619 species and (sorted by increasing molecular weight).  $PM_1$  is the sum of all-the AMS-derived

particulate species. –The CE was 0.84±0.04 and 0.82±0.01 for the SP and NP, respectively. For

1621 comparison, EFs are also included from previously published literature including: Andreae

1622  $(2019)^1$ , Liu et al.  $(2017)^a$ ; Akagi et al.  $(2011)^b$ ; and Simpson et al.  $(2011)^e$ , Koss et al.  $(2018)^2$ , 1623 and Permar et al.  $(2021)^3$ , and Liu et al.  $(2017)^4$ . The Andreae (2019) PM EF represents

 $PM_{2.5}$ . See Table S8 for compounds that did not have exact matches for comparison to literature

values. To derive the EF fFor species measured in mass concentration units, Eq. 32 was

1626 modified by converting TC to mass concentrations using the measured temperature and pressure,

and removing the molar mass ratio term. \* Indicates that the compound was 'calculated' (SI Sect

1628 12.1.1) while the remaining compounds were calibrated. \*\*Estimated, see text in Sect. 3.4.3.

1629 Uncertainties y reflects the standard deviation of the calibration. were estimated by summing in

quadrature the standard error of the average EF (or ER) and the measurement uncertainties (see Sect. 2.5).

1632

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg⁻¹)	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
Particulate	es											
	PM₁	particulat e matter (<1µm)	AMS	112±35	75.5±29. 3	13.2±0.9	6.8±0.8	7.1±0.3	6.6±1.1	18.7±15.9 <sup>1</sup> 26.0±6.2 <sup>4</sup>	58.8±1. 0	65.1±7.3
	F IVI1	black	AIVIS	112-35	0.74±0.3	0.11±0.0	0.13±0.0	0.11±0.0	0.0±1.1 0.14±0.0	0.43±0.21 <sup>1</sup>	0.55±0.	05.117.5
	BC	carbon	SP2	1.3±0.4	0	6	3	2	4	0.39±0.17 <sup>3</sup>	08	0.58±0.19
	NH4	p- ammoniu m	AMS	2.3±1.0	1.2±0.4	0.21±0.0 3	0.11±0.0 3	0.11±0.0 3	0.12±0.0 4	0.34±0.15 <sup>4</sup>	1.1±0.3	1.9±0.6
				2 2 4 5	4.4.0.5	0.078±0.	0.17±0.0	0.14±0.0	0.19±0.0	0.0710.424	0.90±0.	1 2 1 0 2
	NO <sub>3</sub>	p-nitrate p-	AMS	3.2±1.5	1.4±0.5 0.98±0.3	017 0.39±0.0	4 0.060±0.	3 0.066±0.	5 0.055±0.	0.87±0.134	16 0.035±	1.2±0.3
	SO <sub>4</sub>	p- sulphate	AMS	1.7±0.7	0.98±0.3 1	0.39±0.0 3	0.060±0. 022	0.066±0. 020	0.055±0. 023	0.30±0.164	0.035± 0.011	0.054±0.020
	OA	p-total organics	AMS	101±34	72.6±27. 0	12.5±0.8 3	6.6±2.6	6.9±2.4	6.3±2.8	24.3±0.214	57.5±1 9.4	61.7±27.1
Gas Inorganic												
17.031	NH₃	ammonia	LGR	15.4±9. 6	5.2±2.1	- 0.039±2. 2	0.63±0.1 4	0.45±0.0 4	0.82±0.1 9	2.5±1.8 <sup>1</sup> 0.68±0.19 <sup>2</sup>	5.8±0.6	12.6±2.5
		hydrogen				0.18±0.0	0.31±0.0	0.34±0.0	0.29±0.0	0.53±0.30 <sup>1</sup> 0.28±0.060		
27.026	HCN	cyanide	CIMS	3.2±1.6	2.3±1.0	0.18±0.0 7	0.51±0.0 7	0.34±0.0 6	0.29±0.0 7	0.43±0.17 <sup>3</sup>	2.8±0.5	2.9±0.7
	-	carbon monoxid		991±44						121±47 <sup>1</sup>		
28.01	СО	е	Picarro	3	819±327	119±5	116±6	127±4	104±7	99.3±19.7 <sup>3</sup>	108±39	126±52
30.006	NO	nitric oxide	TECOs	0.14±0. 05		0.0014±0 .086	0.016±0. 006	0.016±0. 006	0.14±0.0 5	0.291		0.14±0.046

Molecu- lar	Comp- ound	Compou nd Name	Instru- ment	SP Averag	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF	NP ER	SP ER
Weight				е						(g kg⁻¹)		
		isocyanic		0.52±0.	0.44±0.1	0.068±0.	0.083±0.	0.091±0.	0.076±0.	0.57±0.24 <sup>2</sup>	0.46±0.	
43.025	HNCO	acid	CIMS	17	3	024	029	027	031	0.16±0.04 <sup>3</sup>	13	0.47±0.92
		carbon		414±0.		405.0	1 100 100	1481±10	1511.00	1529±135 <sup>1</sup>	7 4 9 5	0.4.0.45
44.009	CO <sub>2</sub>	dioxide	Picarro	4 0.88±0.	411±0.2	405±0.4 0.39±0.1	1496±92 0.15±0.0	3 0.15±0.0	1511±80 0.88±0.1	1413±61 <sup>3</sup>	7.4±0.5	9.4±0.45
46.005	NO <sub>2</sub>	nitrogen dioxide	TECOs	0.88±0. 17		0.39±0.1 9	0.15±0.0 4	0.15±0.0 4	0.88±0.1 7	1.0 <sup>1</sup>		0.83±0.21
10.005	1102	sum	12005	17		5			,	1.0		0.0010.21
		(NO+NO <sub>2</sub>				0.39±0.2	0.17±0.0	0.17±0.0				
46.005	NOx	)	TECOs	1.0±0.2		0	4	4	1.0±0.2	1.2±0.91		0.97±0.58
47 01 0		nitrous	CINAC	0.22±0.		0.098±0.	0.020±0.	0.020±0.	0.22±0.0	0.0010.202		0.1110.001
47.013	HONO	acid sulphur	CIMS	04		038 0.19±0.4	012 0.26±0.0	012 0.26±0.0	4	0.60±0.20 <sup>2</sup>		0.11±0.061
64.064	SO <sub>2</sub>	dioxide	TECOs	1.3±0.3		6	5	5	1.3±0.3	$0.22\pm0.31^{1}$		1.1±0.16
											0.0008	
		pernitric		0.036±	0.032±0.	0.020±0.	0.0010±0	0.00047±	0.00085±		9±0.00	0.0028±0.003
79.011	HNO <sub>4</sub>	acid	CIMS	0.0049	0043	007	.0019	0.0025	0.001		68	3
		gaseous elementa					0.000087 ±0.00001	0.000082 ±0.00001	0.000092 ±0.00001	0.00023±0.	0.0006 8±0.00	0.00091±0.00
200.59	GEM	Imercury	Tekran	1.6±0.2	1.4±0.1	1.2±0.03	10.00001 7	7	10.00001 6	0.00023±0. 00030 <sup>1</sup>	8 <u>1</u> 0.00 014	0.0009110.00
Gas							1 -		1 -		1 •	
Organic					-			_				
			PTRMS									
			+CIMS									
		non methane	+AWAS +				24.5±1.6					
	ΣΝΜΟ	organic	cartrid				to					
	G	gases	ges				25.6±1.6	26.2±2.1	25.4±5.8			
			PTRMS									
	Estim.		+CIMS									
	NMOG ⊤(see	non methane	+AWAS +				36.8±11.			58.7 <sup>1</sup>		
	Sect	organic	cartrid				3 to			25.0 <sup>2</sup>		
	3.4.3)	gases	ges				39.9±5.8			26.1 <sup>3</sup>		
		carbon										
	NMOG	fraction		936±34							680±11	
	Т	of NMOG	Picarro	1 2026±5	649±225	375±85	31.2±3.8	36.8±5.1	25.5±5.6	5 5 1 2 5 1	1	580±92
16.043	CH4	methane	Picarro	2020±5 4	1982±35	1911±8	8.3±0.9	7.8±0.4	8.7±1.1	5.5±2.5 <sup>1</sup> 5.9±1.8 <sup>3</sup>	107±5	146±16
10.015	0114	meenane	Ticuito	•	1502255	0.34±0.0	0.27±0.0	0.20±0.0	0.34±0.1	5.521.0	10725	110110
26.038	$C_2H_2$	acetylene	AWAS	6.9±2.5	1.8±0.7	041	8	5	1	0.31±0.17 <sup>3</sup>	2.2±0.9	4.0±1.1
				32.4±1		0.64±0.0					12.9±3.	
28.054	C <sub>2</sub> H <sub>4</sub>	ethene	AWAS	4.0	9.3±3.8	1	1.5±0.4	1.3±0.3	1.7±0.5	1.5±1.0 <sup>3</sup>	5	18.3±5.0
		formalde		13.9±4.					0.93±0.3	1.8±0.4 <sup>1</sup> 1.9±0.7 <sup>2</sup>		
30.026	CH <sub>2</sub> O	hyde	PTR	13.9±4. 9	10.1±2.8	4.4±2.0	1.0±0.3	1.1±0.3	0.95±0.5	1.9±0.7 1.9±0.4 <sup>3</sup>	8.1±2.2	8.9±3.2
•			l	27.1±1		1.9±0.01			1		12.5±2.	-
30.07	C <sub>2</sub> H <sub>6</sub>	ethane	AWAS	2.4	10.1±3.4	6	1.3±0.4	1.3±0.3	1.4±0.5	1.1±0.84 <sup>3</sup>	7	13.8±3.8
										2.3±1.0 <sup>1</sup>		
32.042	CH₄O	methanol	PTR	21.9±7. 9	15 0+4 0	6 010 0	1 0+0 4	2.2±0.4	1 6+0 4	$0.90\pm0.35^2$	14.9±2.	12 4+2 6
52.04Z		methanol	FIR	3	15.9±4.9	6.8±0.9	1.9±0.4	2.2±0.4	1.6±0.4	1.5±0.4 <sup>3</sup> 0.060±0.03	9	13.4±3.6
		fragment								0.000±0.03		
		s/propadi				0.39±0.3	0.53±0.1	0.64±0.1	0.42±0.1	0.088±0.04		
40.065	$C_3H_4$	ene*	PTR	3.7±1.6	3.0±1.3	5	1	1	1	1 <sup>2</sup>	3.5±0.6	2.8±0.7
										$0.31\pm0.10^{1}$		
		acetonitri				0.10±0.0	0.44±0.0	0.48±0.0	0.40±0.0	0.086±0.02 7 <sup>2</sup>		
41.053	C <sub>2</sub> H <sub>3</sub> N	le	PTR	2.8±1.4	2.0±0.8	0.10±0.0 6	0.44±0.0 7	0.48±0.0 6	0.40±0.0 8	0.31±0.15	2.6±0.3	2.6±0.5
. 2.000		cyanamid	· · · ·	0.55±0.	0.40±0.1	0.10±0.2	, 0.064±0.	0.067±0.	0.061±0.			
42.041	$CH_2N_2$	e*	PTR	22	2	0	042	042	042		1.4±0.9	1.3±0.9

Molecu-	Comp-	Compou	Instru-	SP	NP	Backgr-	Average	NP EF (g	SP EF (g	Literature	NP ER	SP ER
lar Weight	ound	nd Name	ment	Averag e	Average	ound	EF (g kg <sup>-1</sup> )	kg⁻¹)	kg⁻¹)	EF (g kg <sup>-1</sup> )		
						0.12±0.0	0.68±0.2	0.62±0.1	0.73±0.2			
42.081	C₃H <sub>6</sub>	propene	AWAS	9.7±4.8	2.9±1.3	1	1	5	5	0.74±0.62 <sup>3</sup>	4.2±1.0	5.2±1.5
		acetalde				0.96±0.2				0.81±0.23 <sup>1</sup> 0.92±0.32 <sup>2</sup>		
44.053	C <sub>2</sub> H <sub>4</sub> O	hyde	PTR	7.3±3.2	5.2±2.0	0	1.1±0.2	1.2±0.2	1.0±0.2	1.7±0.4 <sup>3</sup>	6.3±0.9	6.3±1.2
44.097	C₃H <sub>8</sub>	propane	AWAS	6.7±3.2	2.5±1.0	0.28±0.0 3	0.52±0.1 4	0.53±0.1 2	0.50±0.1 6	0.46±0.18 <sup>3</sup>	3.4±0.7	3.4±1.0
11.057	03118	propune	/	0.7 20.2	2.5_1.0	5		-	0	1.0±0.9 <sup>1</sup>	5.120.7	5.121.0
		formic					0.17±0.0	0.17±0.0	0.17±0.0	0.28±0.14 <sup>2</sup>		
46.025	CH <sub>2</sub> O <sub>2</sub>	acid	CIMS	3.0±1.2	2.8±0.9	2.4±0.1	5	4	6	1.5±0.6 <sup>3</sup>	1.2±2.6	0.56±1.3
48.103	CH₄S	methane thiol*	PTR	0.074± 0.039	0.049±0. 022	0.0024±0 .028	0.014±0. 0086	0.015±0. 009	0.013±0. 008	0.011±0.00 6 <sup>2</sup>	0.068± 0.043	0.073±0.043
50.057	CH <sub>6</sub> O₂	methanol hydrate*	PTR	0.25±0. 10	0.16±0.0 6	0.062±0. 057	0.028±0. 020	0.034±0. 023	0.022±0. 017		0.15±0. 10	0.12±0.09
50.057		buten-	FIN	10	0	037	020	025	017		10	0.1210.09
52.076	C4H4	yne/frag ments*	PTR	0.11±0. 05	0.080±0. 034	0.011±0. 044	0.018±0. 010	0.020±0. 011	0.016±0. 010	0.052±0.01 8 <sup>3</sup>	0.086± 0.046	0.081±0.050
										0.025±0.01		
		acrylonitr		0.17±0.	0.12±0.0	0.0024±0	0.036±0.	0.040±0.	0.032±0.	2 <sup>2</sup> 0.044±0.01	0.17±0.	
53.064	C <sub>3</sub> H <sub>3</sub> N	ile*	PTR	0.17±0. 08	6	.013	0.030±0. 018	0.040±0. 018	0.03210.	5 <sup>3</sup>	0.17±0. 07	0.16±0.09
	-3 5	-								0.034±0.01	-	
						-				4 <sup>2</sup>		
54.048	C <sub>3</sub> H <sub>2</sub> O	propynal *	PTR	0.053± 0.033	0.023±0. 0054	0.013±0. 019	0.0087±0 .0053	0.0045±0 .0031	0.013±0. 007	0.037±0.01 5 <sup>3</sup>	0.018± 0.013	0.062±0.032
54.048			PIR	0.033	0054	019	.0053	.0031	007	0.089±0.03	0.013	0.062±0.032
		butadien				-				01		
		e/fragme		0.74±0.	0.47±0.2	0.070±0.	0.15±0.0	0.15±0.0	0.15±0.0	0.34±0.18 <sup>2</sup>	0.62±0.	
54.092	C <sub>4</sub> H <sub>6</sub>	nts*	PTR	38	4	17	8	8	8	0.27±0.10 <sup>3</sup>	30	0.73±0.37
		1,3-								0.089±0.03 0 <sup>1</sup>		
		butadien		0.74±0.	0.20±0.0	0.0041±0	0.065±0.	0.055±0.	0.075±0.	0.34±0.18 <sup>2</sup>	0.29±0.	
54.092	C <sub>4</sub> H <sub>6</sub>	е	AWAS	38	9	.0006	022	016	026	0.27±0.10 <sup>3</sup>	09	0.41±0.12
										0.012±0.00 5 <sup>2</sup>		
		propane		0.11±0.	0.080±0.	0.0097±0	0.022±0.	0.025±0.	0.019±0.	0.037±0.01	0.10±0.	
55.08	C₃H₅N	nitrile*	PTR	05	032	.019	012	012	012	8 <sup>3</sup>	05	0.094±0.057
										0.34 <sup>1</sup>		
56.064	C <sub>3</sub> H <sub>4</sub> O	acrolein	PTR	1.5±0.6	1.0±0.4	0.17±0.0 9	0.28±0.0 5	0.29±0.0 4	0.26±0.0 6	0.97±0.50 <sup>2</sup> 0.40±0.18 <sup>3</sup>	0.82±0. 12	0.83±0.15
50.004	C3114O	cis-2-	1 11	0.16±0.	1.0±0.4	0.016±0.	0.015±0.	4	0.015±0.	0.4010.10	12	0.85±0.15
56.108	$C_4H_8$	butene	AWAS	08		008	006		006			0.078±0.023
56 400	<u> </u>	isobuten		0.94±0.	0.34±0.1	0.062±0.	0.084±0.	0.082±0.	0.086±0.		0.41±0.	0.45+0.40
56.108	C <sub>4</sub> H <sub>8</sub>	e t-2-	AWAS	49 0.13±0.	2	0022 0.010±0.	023 0.012±0.	008	032 0.012±0.		03	0.45±0.13
56.108	C <sub>4</sub> H <sub>8</sub>	butene	AWAS	0.13±0. 07		0.010±0.	0.01210.		0.012±0.			0.063±0.018
					0.41±0.1	0.014±0.	0.13±0.0	0.12±0.0	0.14±0.0		0.60±0.	
56.108	C <sub>4</sub> H <sub>8</sub>	1-butene	AWAS	1.4±0.7	7	005	3	1	4		05	0.74±0.12
	C <sub>2</sub> H <sub>3</sub> N	hydroxy acetonitri		0.021±	0.0078±0	0.00035±	0.0035±0	0.0025±0	0.0044±0	0.033±0.00	0.0095	
57.052	0	le	CIMS	0.0211	.014	0.000331	.0031	.0028	.0034	9 <sup>3</sup>	±0.011	0.021±0.016
		methyl									1	
F7 050	C <sub>2</sub> H <sub>3</sub> N	isocyanat	DTD	0.074±		0.0067±0	0.0052±0		0.0052±0	0.033±0.00		0.024/0.045
57.052	0	e*	PTR	0.029		.06	.0032		.0032	9 <sup>3</sup> 1.6±1.6 <sup>1</sup>		0.024±0.015
							0.82±0.2	0.99±0.2	0.65±0.1	$1.6\pm1.6^{2}$ 0.34±0.12 <sup>2</sup>	0.065±	
58.08	C₃H <sub>6</sub> O	acetone	PTR	6.0±1.8	4.7±1.2	2.5±0.3	2	5	9	0.84±0.22 <sup>3</sup>	0.018	0.072±0.028
										0.11±0.06 <sup>1</sup>		
F0 404		n hut	A)A/AC	1 5:0 7	0.62±0.2	0.098±0.	0.15±0.0	0.16±0.0	0.14±0.0	0.12±0.06 <sup>3</sup>	0.79±0.	0.72+0.20
58.124	C <sub>4</sub> H <sub>10</sub>	n-butane	AWAS	1.5±0.7	2	013	4	4	5	I	17	0.73±0.20

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
		acetic								3.8±2.0 <sup>1</sup>		
60.052	C <sub>2</sub> H <sub>4</sub> O <sub>2</sub> CH <sub>4</sub> N <sub>2</sub>	acid	CIMS	8.8±7.5 0.44±0.	6.0±3.9 0.28±0.0	2.1±0.8 0.067±0.	1.3±0.8 0.078±0.	1.1±0.5 0.079±0.	1.6±0.9 0.076±0.	2.4±0.6 <sup>3</sup>	7.4±3.5 0.29±0.	8.9±5.2
60.056	0	Urea*	PTR	18	8	13	0.078±0. 052	0.079±0. 054	0.07010.		20	0.34±0.21
										0.074±0.03		
61.04	CH₃NO ₂	nitromet hane*	PTR	0.055± 0.023	0.038±0. 020	0.0051±0 .023	0.011±0. 007	0.010±0. 007	0.011±0. 007	0 <sup>2</sup> 0.078±0.00 9 <sup>3</sup>	0.036± 0.024	0.048±0.030
		ethylene		0.023±		0.0036±0	0.0036±0		0.0036±0			
62.068	$C_2H_6O_2$	glycol*	PTR	0.0077		.018	.0023		.0023	0.001010.0		0.015±0.010
62.13	C₂H <sub>6</sub> S	dimethyl sulfide	PTR	0.051± 0.022		0.011±0. 034	0.0067±0 .0047		0.0067±0 .0047	0.0016±0.0 008 <sup>2</sup> 0.080±0.08 3 <sup>3</sup> 0.0047 <sup>4</sup>		0.029±0.020
					0.40.00	0.005.0	0.000.0		0.000.0	0.011±0.00		
66.103	C₅H <sub>6</sub>	cyclopent andiene*	PTR	0.13±0. 05	0.12±0.0 4	0.025±0. 039	0.032±0. 019	0.041±0. 022	0.023±0. 016	5 <sup>3</sup>	0.14±0. 07	0.096±0.064
00.105	03110	undicite		03		000	015	022	010	0.054±0.02	0,	0.05020.001
67.091	C₄H₅N	pyrrole*	PTR	0.10±0. 06	0.067±0. 033	- 0.00073± 0.013	0.026±0. 014	0.027±0. 014	0.025±0. 014	9 <sup>2</sup> 0.039±0.02 1 <sup>3</sup>	0.090± 0.046	0.098±0.055
07.051	C41131V	pyrroic	1 110	00	000	-	014	014	014	0.36±0.44 <sup>1</sup>	0.040	0.05010.055
co 075		c *		4.5.0.0	4.4.9.5	0.0083±0	0.39±0.1	0.43±0.1	0.35±0.2	0.36±0.11 <sup>2</sup>		
68.075	C <sub>4</sub> H <sub>4</sub> O	furan*	PTR	1.5±0.8	1.1±0.5	.035	9	7	0	0.43±0.19 <sup>3</sup> 0.074 <sup>1</sup>	1.4±0.6	1.4±0.8
						0.52±0.2	0.42±0.2	0.64±0.3	0.19±0.1	0.074 <sup>2</sup> 0.22±0.11 <sup>2</sup> 0.082±0.09		
68.119	C₅H <sub>8</sub>	isoprene	PTR	1.7±0.7	1.8±0.8	5	6	4	5	5 <sup>3</sup>	2.1±1.1	0.47±0.47
				0.82±0.	0.35±1.6		0.30±0.1	0.40±0.1	0.20±0.1	0.074 <sup>1</sup> 0.22±0.11 <sup>2</sup> 0.082±0.09		
68.119	C₅H <sub>8</sub>	isoprene	AWAS	46	5	1.3±0.4	8	7	9	5 <sup>3</sup>	2.0±1.0	0.18±0.18
69.083	C₄H₅O	*	PTR	0.017± 0.0064	0.013±0. 0063	0.00079± 0.01	0.0043±0 .0025	0.0047±0 .0028	0.0038±0 .0022		0.015± 0.009	0.015±0.008
09.085	C4H5O	butane	FIN	0.030±	0.022±0.	0.001	0.0077±0	0.0088±0	0.0065±0	0.011±0.00 5 <sup>2</sup> 0.020±0.01	0.028±	0.01310.008
69.107	$C_4H_7N$	nitrile*	PTR	0.014	0.02210.	.011	.0051	.0059	.0042	0 <sup>3</sup>	0.019	0.025±0.016
		MVK, methacro lein, crotonald				0.91±0.1	0.19±0.0	0.20±0.0	0.18±0.1	0.11±0.12 <sup>1</sup> 0.34±0.15 <sup>2</sup>	0.66±0.	
70.091	C <sub>4</sub> H <sub>6</sub> O	ehyde	PTR	1.3±0.3	1.3±0.3	7	9	8	1	0.39±0.15 <sup>3</sup>	26	0.68±0.38
				Ī		1			1	0.046±0.02	I	
70.135	C₅H10	c-2- pentene	AWAS	0.040± 0.017	0.013±0. 0034	0.0040±0 .0006	0.0040±0 .0012	0.0033±0 .0007	0.0048±0 .0016	5 <sup>1</sup> 0.015±0.00 8 <sup>3</sup>	0.013± 0.004	0.021±0.0057
										0.046±0.02		
70.135	C₅H <sub>10</sub>	cyclopent ane	AWAS	0.031± 0.013	0.015±0. 0022	0.0052±0 .0003	0.0035±0 .0009	0.0038±0 .0009	0.0031±0 .0009	5 <sup>1</sup> 0.015±0.00 8 <sup>3</sup>	0.016± 0.004	0.014±0.005
	-		_	-						0.046±0.02		
70.135	C₅H <sub>10</sub>	1- pentene	AWAS	0.42±0. 21	0.15±0.0 6	0.0053±0 .0012	0.052±0. 015	0.053±0. 013	0.052±0. 018	5 <sup>1</sup> 0.015±0.00 8 <sup>3</sup>	0.21±0. 05	0.22±0.06
										0.046±0.02		
70.135	C₅H10	t-2- pentene	AWAS	0.13±0. 12	0.068±0. 020	0.0094±0 .0058	0.018±0. 013	0.0049±0 .0040	0.031±0. 018	5 <sup>1</sup> 0.015±0.00 8 <sup>3</sup>	0.013± 0.010	0.063±0.035

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
										0.046±0.02		
70.135	C₅H <sub>10</sub>	2-me-1- butene	AWAS	0.12±0. 061	0.047±0. 018	0.0088±0 .0017	0.014±0. 005	0.014±0. 004	0.015±0. 005	5 <sup>1</sup> 0.015±0.00 8 <sup>3</sup>	0.056± 0.013	0.062±0.018
										0.046±0.02 5 <sup>1</sup>	0.0068	
70.135	C5H10	2-me-2- butene	AWAS	0.019± 0.0071	0.0077±0 .0016	0.0034±0 .0006	0.0019±0 .0006	0.0017±0 .0004	0.0022±0 .0008	0.015±0.00 8 <sup>3</sup>	±0.003 0	0.0095±0.002 7
										0.046±0.02 5 <sup>1</sup>		
70.135	C5H10	3-me-1- butene	AWAS	0.10±0. 10	0.045±0. 032	0.029±0. 012	0.016±0. 010	0.0078±0 .0037	0.025±0. 013	0.015±0.00 8 <sup>3</sup>	0.045± 0.018	0.058±0.028
72.063	C <sub>3</sub> H <sub>4</sub> O <sub>2</sub>	acrylic acid	CIMS	0.28±0. 24	0.21±0.1 5	0.060±0. 046	0.096±0. 048	0.13±0.0 6	0.062±0. 035	0.22±0.08 <sup>3</sup>	0.25±0. 11	0.35±0.20
70 107	<u> </u>	MEK, 2- methyl acetate, ethyl	DTD	0.80±0.	0.57±0.2 3	0.097±0. 051	0.18±0.0	0.22±0.0 8	0.14±0.0 6		0.67±0.	
72.107	C <sub>4</sub> H <sub>8</sub> O	formate n-	PTR	35 0.59±0.	3 0.26±0.0	0.035±0.	7 0.078±0.	8 0.086±0.	6 0.070±0.	0.057±0.02	23 0.34±0.	0.54±0.20
72.151	C <sub>5</sub> H <sub>12</sub>	pentane 2-	AWAS	28	9	005	021	019	023	8 <sup>3</sup>	07	0.29±0.08
72.151	C5H12	2- methylbu tane	AWAS	0.21±0. 08	0.11±0.0 1	0.051±0. 001	0.022±0. 006	0.024±0. 0049	0.021±0. 007	0.057±0.02 8 <sup>3</sup>	0.097± 0.019	0.086±0.027
74.079	C <sub>3</sub> H <sub>6</sub> O <sub>2</sub>	propanoi c acid	CIMS	0.81±0. 24	0.70±0.1 5	0.49±0.1 7	0.13±0.0 8	0.12±0.0 8	0.14±0.0 9	0.57±0.20 <sup>3</sup>	0.35±0. 24	0.51±0.30
74.079	C <sub>3</sub> H <sub>6</sub> O <sub>2</sub>	hydroxy acetone/ ethyl formate *	PTR	1.5±0.6 2	1.1±0.4	0.30±0.1 1	0.32±0.2 3	0.35±0.2 5	0.30±0.2 0		1.0±0.8	1.1±0.7
78.114	C <sub>6</sub> H <sub>6</sub>	benzene	PTR	1.4±0.6 9	1.0±0.5	0.054±0. 045	0.41±0.0 6	0.47±0.0 6	0.36±0.0 7	$\begin{array}{c} 0.57 \pm 0.21^1 \\ 0.42 \pm 0.25^2 \\ 0.50 \pm 0.14^3 \end{array}$	1.3±0.2	1.2±0.2
80.086	C₅H₄O	cyclopent andienon e/isomer s*	PTR	0.049± 0.026	0.024±0. 012	- 0.00054± 0.0061	0.011±0. 006	0.0093±0 .0047	0.012±0. 007	0.027±0.01 7 <sup>3</sup>	0.026± 0.013	0.040±0.023
		cyclohex adiene/m onoterpe ne										
90.12	C II	fragment *	DTD	0.45±0.	0.40±0.2	0.040±0.	0.14±0.0	0.17±0.0	0.10±0.0		0.48±0.	0.24+0.20
80.13	C <sub>6</sub> H <sub>8</sub>	pentene nitriles/ methyl	PTR	18 0.018±	0 0.013±0.	040 0.0015±0	7 0.0047±0	7 0.0050±0	6 0.0044±0	0.020±0.01	19 0.014±	0.34±0.20
81.118	$C_5H_7N$	pyrrole*	PTR	0.0181	0053	.0055	.0032	.0036	.0028	1 <sup>3</sup>	0.010	0.015±0.009
82.102	C₅H₀O	methyl furan*	PTR	1.1±0.5	0.69±0.3 2	0.042±0. 065	0.30±0.1 5	0.31±0.1 3	0.29±0.1 7	0.32±0.11 <sup>2</sup> 0.28±0.13 <sup>3</sup>	0.84±0. 34	0.96±0.55
82.146	C <sub>6</sub> H <sub>10</sub>	cyclohex ene*	PTR	0.14±0. 06	0.093±0. 044	0.015±0. 035	0.054±0. 030	0.075±0. 038	0.033±0. 020	0.015±0.01 1 <sup>3</sup>	0.20±0. 10	0.11±0.07
83.09	C₄H₅N O	methylox azole*	PTR	0.0096 ±0.004 4		0.00012± 0.0083	0.0020±0 .0011		0.0020±0 .0011			0.0066±0.003 7
		pentanen itriles*		0.049±	0.037±0.	0.0024±0 .0088	0.016±0.	0.019±0.	0.013±0. 008	0.021±0.01 1 <sup>3</sup>	0.049± 0.024	0.042±0.025
83.134	C₅H9N	iu iles "	PTR	0.025	017 0.79±0.2	.0088 0.29±0.1	008 0.38±0.1	009 0.23±0.0	0.52±0.2	1 <sup>3</sup> 0.32±0.11 <sup>3</sup>	0.024 0.61±0.	0.042±0.025
84.074	$C_4H_4O_2$	*	CIMS	1.7±0.7	4	3	4	4	0		20	1.3±0.6

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
		furanone		0.54±0.	0.37±0.1	- 0.0030±0	0.16±0.0	0.16±0.0	0.15±0.0		0.42±0.	
84.074	$C_4H_4O_2$	* cyclopent	ptr	25	6	.041	9	9	8		23	0.48±0.26
84.118	C₅HଃO	anone/is omers*	PTR	0.23±0. 11	0.16±0.0 7	0.017±0. 026	0.069±0. 036	0.073±0. 032	0.065±0. 039	0.087±0.03 8 <sup>3</sup>	0.19±0. 09	0.21±0.12
				0.029±	0.021±0.	0.0013±0	0.015±0.	0.020±0.	0.0098±0	0.008±0.01	0.052±	
84.162	C <sub>6</sub> H <sub>12</sub>	hexene*	PTR	0.013	0065	.015	011	014	.0065	4 <sup>3</sup>	0.037 0.0069	0.031±0.019
84.162	$C_6H_{12}$	c-2- hexene	AWAS	0.019± 0.012	0.0079±0 .0024	0.0031±0 .0002	0.0020±0 .0014	0.0021±0 .0016	0.0020±0 .0011	0.008±0.01 4 <sup>3</sup>	±0.004 4	0.0064±0.005 5
84.162	C <sub>6</sub> H <sub>12</sub>	cyclohex ane	AWAS	0.022± 0.0077	0.0089±0 .0010	0.0051±0 .0008	0.0022±0 .0016	0.0019±0 .0015	0.0026±0 .0016	0.008±0.01 4 <sup>3</sup>	0.0064 ±0.005 3	0.0097±0.005 8
		butanedi one/isom		0.57±0.	0.39±0.1	0.089±0.	0.13±0.0	0.13±0.0	0.13±0.0		0.33±0.	
86.09	C <sub>4</sub> H <sub>6</sub> O <sub>2</sub>	ers	ptr	23	4	077	9	9	9		23 0.0095	0.41±0.26
86.134	$C_5H_{10}O$	pentanon e	PTR	0.14±0. 07	0.11±0.0 4	0.013±0. 020	0.046±0. 025	0.053±0. 026	0.038±0. 024	0.062±0.02 3 <sup>3</sup>	±0.004 6	0.0080±0.004 9
86.178	$C_6H_{14}$	n-hexane	AWAS	0.31±0. 14	0.13±0.0 5	0.013±0. 0012	0.049±0. 020	0.053±0. 019	0.044±0. 021	0.050±0.03 6 <sup>3</sup>	0.17±0. 06	0.16±0.07
86.178	C <sub>6</sub> H <sub>14</sub>	2,3-DMB	AWAS	0.017± 0.0091	0.013±0. 001	0.0048±0 .0001	0.0031±0 .0012	0.004±0. 0012	0.0022±0 .0013	0.050±0.03 6 <sup>3</sup>	0.014± 0.004	0.0066±0.003 9
001270	001114	2,3- methylpe		0.090±	0.026±0.	0.011±0.	0.010±0.	0.0089±0	0.011±0.	0.050±0.03	0.032±	
86.178	$C_6H_{14}$	ntane	AWAS	0.047	004	0003	005	.0041	006	6 <sup>3</sup> 0.012±0.00	0.011	0.039±0.021
							0 70 0 7	0.56.0.5		5 <sup>2</sup>	0.000.	
88.062	$C_3H_4O_3$	pyruvic acid	CIMS	4.4±2.4	3.4±1.0	2.3±0.3	0.72±0.7 1	0.56±0.5 4	0.89±0.8 5	0.019±0.00 8 <sup>3</sup>	0.022± 0.022	- 0.0025±0.010
88.106	C4H8O2	methyl propanoa te *	ptr	0.24±0. 11	0.17±0.0 7	0.021±0. 040	0.070±0. 043	0.075±0. 047	0.065±0. 040		0.19±0. 12	0.20±0.12
88.168	C4H8OS	oxathian e*	PTR	0.012± 0.0049	0.0090±0 .0030	- 0.00061± 0.0073	0.0031±0 .0024	0.0023±0 .0021	0.0040±0 .0026		0.0058 ±0.005 3	0.012±0.008
90.125	C7H6	*	PTR	0.026± 0.014		0.0012±0 .018	0.0074±0 .0011		0.0074±0 .0011			0.022±0.013
		diethyl sulfide, butaneth		0.31±0.	0.22±0.1	0.036±0.	0.077±0.	0.083±0.	0.071±0.		0.20±0.	
90.184	C <sub>4</sub> H <sub>10</sub> S	iol	PTR	13 0.62±0.	0 0.48±0.2	12 0.034±0.	015 0.26±0.0	014 0.26±0.0	015 0.26±0.0	0.35±0.11 <sup>1</sup> 0.25±0.13 <sup>2</sup>	04 0.63±0.	0.21±0.05
92.141	C <sub>2</sub> H <sub>8</sub>	toluene	PTR	30 0.0071 ±0.003	1	037 0.0012±0	7 0.0025±0	4	9 0.0025±0	0.42±0.16 <sup>3</sup>	08	0.71±0.23 0.0070±0.004
93.082	O <sub>3</sub>	*	PTR	4		.0046	.0012		.0012	0.0026±0.0		8
93.085	C₅H₃N O	furancarb onitrile*	PTR	0.056± 0.031	0.038±0. 017	0.00022± 0.0040	0.020±0. 011	0.022±0. 010	0.018±0. 011	010 <sup>2</sup> 0.0088±0.0 037 <sup>3</sup>	0.053± 0.025	0.053±0.030
94.113	C <sub>6</sub> H <sub>6</sub> O	phenol*	PTR	0.42±0. 22	0.27±0.1 3	0.0026±0 .030	0.12±0.0 6	0.12±0.0 5	0.12±0.0 7	3.0 <sup>1</sup> 0.57±0.36 <sup>2</sup> 0.33±0.13 <sup>3</sup>	0.28±0. 11	0.35±0.20
94.157	C7H10	cyclohept adiene*	PTR	0.080± 0.035	0.061±0. 028	0.005±0. 020	0.021±0. 011	0.023±0. 011	0.020±0. 012		0.053± 0.025	0.056±0.033
94.19	C <sub>2</sub> H <sub>6</sub> S <sub>2</sub>	dimethyl disulfide*	PTR	0.012± 0.0071		- 0.0012±0 .012	0.0039±0 .0022		0.0039±0 .0022	0.0024±0.0 009 <sup>2</sup>		0.011±0.006

Molecu-	Comp-	Compou	Instru-	SP	NP	Backgr-	Average	NP EF (g	SP EF (g	Literature	NP ER	SP ER
lar Weight	ound	nd Name	ment	Averag e	Average	ound	EF (g kg <sup>-1</sup> )	kg⁻¹)	kg⁻¹)	EF (g kg⁻¹)		
											0.0099	
05 077		*	DTD	0.014± 0.0085	0.0094±0	0.0012±0	0.0030±0	0.0043±0	0.0038±0		±0.006 3	0.011+0.007
95.077	$C_5H_3O_2$		PTR	0.0085	.0043	.0070	.0021	.0028	.0012	0.0099±0.0	3	0.011±0.007
	C₅H₅N	pyridinol		±0.002	0.0045±0	0.00087±	0.0022±0	0.0021±0	0.0023±0	0.0055±0.0	±0.0048	0.0063±0.003
95.101	0	*	PTR	6	.0026	0.0041	.0015	.0017	.0012		1	2
				0.0068		-						
		C2		±0.003		0.000091	0.0021±0		0.0021±0			0.0060±0.003
95.145	C <sub>6</sub> H <sub>9</sub> N	pyrrole*	PTR	4		±0.0054	.0012		.0012	0.61 <sup>1</sup>		3
						- 0.0040±0	0.65±0.3	0.67±0.2	0.64±0.3	0.54±0.17 <sup>2</sup>		
96.085	$C_5H_4O_2$	furfural*	PTR	2.0±1.0	1.3±0.6	.029	1	6	6	$0.53\pm0.21^3$	1.5±0.6	1.8±1.0
						-						
		C2-		0.32±0.	0.20±0.0	0.00016±	0.087±0.	0.086±0.	0.087±0.	$0.20\pm0.10^{3}$	0.20±0.	
96.129	C <sub>6</sub> H <sub>8</sub> O	furan*	PTR	16	9	0.024	044	037	050		09	0.24±0.14
96.173	C7H12	cyclohept ene*	PTR	0.042± 0.02	0.035±0. 013	0.0049±0 .017	0.022±0. 015	0.033±0. 020	0.011±0. 007		0.076± 0.047	0.031±0.020
50.175	C/112	ene	TIK	0.02	015	.017	015	020	007		0.0098	0.031±0.020
	$C_4H_3N$			0.010±	0.0093±0	0.0012±0	0.0030±0	0.0044±0	0.0036±0		±0.007	0.0096±0.007
97.073	O <sub>2</sub>	*	PTR	0.005	.0026	.0075	.0021	.0027	.0012		3	1
										0.0088±0.0	0.0093	
97.161	$C_6H_{11}N$	hexaneni trile*	PTR	0.011± 0.0053	0.0077±0 .0041	0.00031± 0.0040	0.0040±0 .0026	0.0041±0 .0028	0.0040±0 .0023	047 <sup>3</sup>	±0.006 3	0.011±0.006
97.101	C6H11N	maleic	PIK	0.0053	.0041	0.0040	.0026	.0028	.0023		3	0.011±0.006
		anhydrid		0.21±0.	0.16±0.0	0.011±0.	0.070±0.	0.072±0.	0.068±0.	0.14±0.07 <sup>3</sup>	0.16±0.	
98.057	$C_4H_2O_3$	e* ,	PTR	08	5	032	036	031	040		07	0.18±0.11
		furan										
		methanol				0.004.0	0.050.0	0.054.0	0.054.0	0.38±0.15 <sup>2</sup>		
98.101	C₅H <sub>6</sub> O <sub>2</sub>	/isomers *	PTR	0.28±0. 13	0.20±0.0 8	0.021±0. 047	0.058±0. 030	0.061±0. 025	0.054±0. 034	0.090±0.04 3 <sup>3</sup>	0.14±0. 06	0.15±0.09
56.101	C5116O2	methyl	1 11	15	0	047	050	025	034	0.022±0.00	00	0.15±0.05
		cyclopent								9 <sup>2</sup>		
		anone/is		0.052±	0.036±0.	0.0027±0	0.015±0.	0.017±0.	0.013±0.	0.034±0.01	0.038±	
98.145	C <sub>6</sub> H <sub>10</sub> O	omers*	PTR	0.023	015	.015	008	009	008	5 <sup>2</sup>	0.019	0.035±0.020
		unsaturat ed C5										
		carboxyli		0.20±0.	0.25±0.0	0.078±0.	0.072±0.	0.10±0.0	0.045±0.	0.11±0.04 <sup>3</sup>	0.22±0.	
100.117	$C_5H_8O_2$	c acids	CIMS	03	7	043	027	3	021		07	0.13±0.06
		methyl										
		methacry										
100 117	C 11 O	late/isom ers*	atr	0.15±0.	0.12±0.0 4	0.017±0.	0.036±0.	0.035±0.	0.037±0.		0.078±	0.008+0.062
100.117	C <sub>5</sub> H <sub>8</sub> O <sub>2</sub>	ers	ptr	06	4	041	023	022	024	0.0046±0.0	0.049	0.098±0.062
		hexanal/								029 <sup>2</sup>		
		hexanon		0.022±	0.018±0.	0.003±0.	0.0065±0	0.0074±0	0.0057±0	0.013±0.00	0.016±	
100.161	C <sub>6</sub> H <sub>12</sub> O	e*	PTR	0.0077	008	010	.0043	.0049	.0035	6 <sup>3</sup>	0.011	0.015±0.009
		C4 oxo-					0.7410.0	0.57.0.0	0.0210.4	0.044±0.02		
102.089	C <sub>4</sub> H <sub>6</sub> O <sub>3</sub>	carboxyli c acids	CIMS	4.7±1.2	3.8±0.7	2.3±0.2	0.74±0.3 7	0.57±0.3 0	0.92±0.4 3	0 <sup>3</sup>	1.2±0.7	2.4±1.1
102.005	C41 16 U3	acetic	CINIS	7.7 ±1.2	5.0±0.7	2.3±0.2	,		5		1.2.10.7	£.7±1.1
		anhydrid		0.033±	0.022±0.	0.0020±0	0.0075±0	0.0078±0	0.0072±0		0.017±	
102.089	$C_4H_6O_3$	e*	ptr	0.016	008	.018	.0046	.0046	.0045		0.010	0.019±0.012
100	$C_5H_{10}O$	valeric		0.083±	0.057±0.	0.0037±0	0.024±0.	0.027±0.	0.020±0.		0.059±	0.050-5-5
102.133	2	acid*	ptr	0.042	025	.016	014	016	012		0.035	0.052±0.031
103.121	C4H9N O2	*	PTR	0.017± 0.0063	0.013±0. 005	0.0012±0 .0032	0.0030±0 .0020	0.0074±0 .0026	0.0064±0 .0011		0.016± 0.009	0.016±0.010
103.121	02		1.11	0.0003	005	.0052	.0020	.0020	.0011	0.021±0.00	0.005	0.01010.010
										4 <sup>2</sup>		
		benzonitr		0.15±0.	0.11±0.0	0.00028±	0.060±0.	0.065±0.	0.054±0.	0.055±0.02	0.14±0.	
103.124	C7H5N	ile*	PTR	08	5	0.0051	029	026	031	2 <sup>3</sup>	06	0.14±0.08

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
	C <sub>5</sub> H <sub>12</sub> O	pentaned		0.0073 ±0.003	0.0052±0	- 0.00087±	0.0029±0	0.0033±0	0.0024±0		0.0069 ±0.006	0.0060±0.003
104.149	2	iol*	PTR	4	.0016	0.0057	.0025	.0033	.0013		9	2
										0.088±0.05		
				0.053±	0.041±0.	0.000051	0.039±0.	0.056±0.	0.022±0.	6 <sup>2</sup> 0.018±0.01	0.12±0.	
104.152	C <sub>8</sub> H <sub>8</sub>	styrene*	PTR	0.027	020	±0.0092	021	026	013	2 <sup>3</sup>	06	0.058±0.033
	C <sub>4</sub> H <sub>10</sub> O	Diethylen		0.014±		0.0016±0	0.0036±0		0.0036±0			0.0088±0.006
106.121	3	e glycol*	PTR	0.0035		.011	.0026		.0026	0.095±0.05		1
										3 <sup>2</sup>		
		benzalde		0.10±0.	0.079±0.	0.011±0.	0.036±0.	0.042±0.	0.030±0.	0.084±0.02	0.087±	
106.124	C <sub>7</sub> H <sub>6</sub> O	hyde* C8	PTR	05	037	015	019	018	019	6 <sup>3</sup>	0.038	0.077±0.047
		aromatic		0.19±0.	0.13±0.0	0.0082±0	0.075±0.	0.082±0.	0.068±0.		0.17±0.	
106.168	C <sub>8</sub> H <sub>10</sub>	S	PTR	09	6	.020	012	012	013	0.21±0.08 <sup>3</sup>	02	0.17±0.03
	6 H N	pyridine		0.0035		-	0.0015+0		0.0045+0			0.0000.0000
107.112	C <sub>6</sub> H₅N O	aldehyde *	PTR	±0.002 1		0.00051± 0.0036	0.0015±0 .0008		0.0015±0 .0008			0.0038±0.002 0
10/1112	0	dimethyl		-		0.0000	10000		10000			
		pryidine/		0.0048								
107.156	C7H9N	heptyl nitriles*	PTR	±0.001 4		0.000012 ±0.0040	0.0018±0 .001		0.0018±0 .0010	0.0050±0.0 033 <sup>2</sup>		0.0043±0.002 4
107.150	C/Hgiv	menes	1 110	-		10.0040	.001		.0010	0.084±0.02		-
		benzoqui								4 <sup>2</sup>		
108.096	C <sub>6</sub> H <sub>4</sub> O <sub>2</sub>	none/qui none*	PTR	0.093± 0.043	0.061±0. 023	0.0025±0 .019	0.025±0. 013	0.024±0. 011	0.025±0. 015	0.077±0.02 0 <sup>3</sup>	0.049± 0.022	0.062±0.035
108.096	C6H4U2	methyl	PIR	0.043	023	.019	013	011	015	0-	0.022	0.062±0.035
		phenol/a										
		nisol/cres		0.13±0.	0.083±0.	0.00068±	0.040±0.	0.040±0.	0.040±0.	0.41±0.17 <sup>2</sup>	0.083±	0.000.00.057
108.14	C <sub>7</sub> H <sub>8</sub> O	ol* cycloocta	PTR	07 0.036±	043 0.029±0.	0.0094 0.0038±0	020 0.015±0.	017 0.017±0.	023 0.013±0.	0.23±0.11 <sup>3</sup>	0.035 0.034±	0.099±0.057
108.184	C <sub>8</sub> H <sub>12</sub>	diene*	PTR	0.016	013	.017	0089	010	008		0.020	0.032±0.020
				0.019±	0.013±0.	0.0012±0	0.0030±0	0.0055±0	0.0055±0		0.011±	
109.104	C <sub>6</sub> H <sub>5</sub> O <sub>2</sub>	* benzene	PTR	0.0095	005	.0058	.0020	.0026	.0011		0.007	0.014±0.008
		diol/met				-						
		hyl		0.34±0.	0.22±0.1	0.00028±	0.11±0.0	0.11±0.0	0.11±0.0	0.68±0.29 <sup>2</sup>	0.21±0.	
110.112	$C_6H_6O_2$	furfural*	PTR	17	1	0.016	5	4	6	0.25±0.12 <sup>3</sup>	08	0.27±0.15
		norcamp								0.079±0.03 2		
		hor/C3		0.096±	0.062±0.	0.0014±0	0.032±0.	0.030±0.	0.034±0.	0.046±0.02	0.059±	
110.156	C <sub>7</sub> H <sub>10</sub> O	furan*	PTR	0.049	030	.016	018	015	020	4 <sup>3</sup>	0.029	0.083±0.048
110.2	C <sub>8</sub> H <sub>14</sub>	cycloocte ne*	PTR	0.017± 0.0083	0.012±0. 004	0.0011±0 .0085	0.0088±0 .0071	0.012±0. 009	0.0053±0 .0034		0.024± 0.019	0.014±0.009
	-0.14	dihydrox										
		У,										
		piridine/ methyl		0.0074							0.0061	
	C₅H₅N	maleimid		±0.003	0.0062±0	0.00025±	0.0026±0	0.0031±0	0.0022±0	0.024±0.00	±0.004	0.0051±0.003
111.1	O <sub>2</sub>	e*	PTR	1	.0021	0.0040	.0018	.0022	.0015	8 <sup>3</sup>	3	4
		furoic acid/hydr										
		oxy		0.18±0.	0.14±0.0	0.071±0.	0.041±0.	0.044±0.	0.038±0.		0.087±	
112.084	$C_5H_4O_3$	furfural*	PTR	06	4	058	036	038	035	0.12±0.03 <sup>3</sup>	0.074	0.089±0.080
		cyclohex		0.055	0.000	-	0.01110	0.01115	0.01115		0.000	
112.128	C <sub>6</sub> H <sub>8</sub> O <sub>2</sub>	anedione *	PTR	0.057± 0.027	0.044±0. 021	0.0017±0 .017	0.014±0. 0072	0.014±0. 007	0.014±0. 007	0.12±0.06 <sup>3</sup>	0.028± 0.014	0.033±0.018
112.120	C61 1802	ethylcycl	1 11	0.027	021	.017	0072	007	007	0.12±0.00	0.014	0.03310.010
		oheptano		0.019±		0.0019±0	0.0070±0		0.0070±0	0.014±0.00		
112.172	C7H12O	ne*	PTR	0.0074		.0087	.0045		.0045	7 <sup>3</sup>	1	0.016±0.010

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
114.144	C <sub>6</sub> H <sub>10</sub> O 2	sum of cyclic saturated and n- unsaturat ed C5 carboxyli c acids	CIMS		0.19±0.0 7	0.074±0. 086	0.12±0.0 4	0.12±0.0 44		0.039±0.01 7 <sup>3</sup>	0.11±0. 04	
		Caprolact one/c6 esters/ c6										
114.144	C <sub>6</sub> H <sub>10</sub> O	diketone isomers*	ptr	0.033± 0.011	0.029±0. 011	0.0065±0 .019	0.0068±0 .0048	0.0082±0 .0057	0.0053±0 .0037		0.016± 0.011	0.013±0.009
114.188	C7H14O	heptanon e/heptan al/isomer s*	PTR	0.010± 0.006		0.00080± 0.0078	0.0039±0 .0025		0.0039±0 .0025	0.0072±0.0 025 <sup>3</sup>		0.0090±0.005 7
116.116	C₅H8O3	C5 oxo- carboxyli c acids	CIMS	0.18±0. 04	0.15±0.0 3	0.10±0.0 1	0.031±0. 017	0.026±0. 013	0.037±0. 020	0.034±0.01 9 <sup>3</sup>	0.050± 0.025	0.083±0.045
116.16	C <sub>6</sub> H <sub>12</sub> O	butyl acetate/c 6 esters *	ptr	0.023± 0.010	0.019±0. 009	0.00086± 0.0081	0.0073±0 .0045	0.0094±0 .0054	0.0052±0 .0033		0.018± 0.010	0.012±0.007
116.222	C <sub>6</sub> H <sub>12</sub> S	cyclohex anethiol*	PTR	0.0094 ±0.003 7	0.0073±0 .0017	- 0.00048± 0.0064	0.0032±0 .0028	0.0040±0 .0037	0.0025±0 .0016		0.0075 ±0.006 9	0.0056±0.003 6
118.088	$C_4H_6O_4$	succinic acid*	PTR	0.0026 ±0.001 1	0.0025±0 .0011	- 0.0012±0 .0031	0.0017±0 .0007	0.0026±0 .0009	0.00081± 0.00034		0.0048 ±0.001 6	0.0018±0.000 7
118.135	CଃH₅O	benzofur an*	PTR	0.039± 0.022	0.025±0. 016	- 0.0018±0 .0060	0.017±0. 009	0.018±0. 010	0.017±0. 009	$\begin{array}{c} 0.037 \pm 0.02 \\ 0^2 \\ 0.041 \pm 0.01 \\ 5^3 \end{array}$	0.034± 0.018	0.038±0.021
118.179	C9H10	methylst yrene/pr openyl benzenes *	PTR	0.022± 0.010	0.016±0. 007	0.0021±0 .0079	0.018±0. 012	0.024±0. 016	0.011±0. 007	0.037±0.02 0 <sup>3</sup>	0.046± 0.030	0.025±0.015
119.167	C8H∂N	*	PTR	0.0039 ±0.002 2		0.0012±0 .0017	0.0016±0 .0015		0.0016±0 .0015			0.0035±0.002 9
120.151	C8H8O	methylbe nzaldehy de/tolual dehyde*	PTR	0.064± 0.031	0.039±0. 019	0.0026±0 .013	0.025±0. 014	0.024±0. 012	0.026±0. 016	0.13±0.08 <sup>2</sup> 0.082±0.03 0 <sup>3</sup>	0.044± 0.022	0.058±0.034
		trimethyl benzene/ C9								0.051 <sup>1</sup> 0.051±0.02 2		
120.195	C9H12	aromatic s*	PTR	0.070± 0.031	0.056±0. 022	0.0078±0 .017	0.052±0. 029	0.075±0. 037	0.029±0. 018	0.069±0.03 1 <sup>3</sup>	0.14±0. 07	0.064±0.039
		benzoic acid/hydr oxybenza		0.068±	0.053±0.	0.0092±0	0.020±0.	0.021±0.	0.019±0.	0.079±0.03 5 <sup>2</sup> 0.065±0.02	0.037±	
122.123	C <sub>7</sub> H <sub>6</sub> O <sub>2</sub>	ldehyde* xylenol/C	PTR	0.025	016	.018	011	010	012	3 <sup>3</sup>	0.018	0.040±0.026
122.167	C <sub>8</sub> H <sub>10</sub> O	2 phenol/ methylan isole*	PTR	0.033± 0.019	0.025±0. 014	0.00069± 0.0081	0.015±0. 0082	0.016±0. 009	0.013±0. 008	0.11±0.04 <sup>2</sup> 0.10±0.06 <sup>3</sup>	0.029± 0.015	0.029±0.017

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg <sup>-1</sup> )	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
		cyclohexy		0.019±	0.014±0.	0.0027±0	0.0076±0	0.0083±0	0.0068±0		0.015±	
122.211	C <sub>9</sub> H <sub>14</sub>	lallene*	PTR	0.0074	005	.0083	.0051	.0059	.0043	0.070+0.04	0.011	0.015±0.009
		hydroxy benzoqui		0.014±	0.011±0.	- 0.00040±	0.0032±0	0.0029±0	0.0035±0	0.073±0.01 8 <sup>2</sup> 0.045±0.02	0.0051 ±0.003	0.0075±0.004
124.095	$C_6H_4O_3$	none*	PTR	0.008	005	0.0051	.0022	.0021	.0023	6 <sup>3</sup>	7	7
124.139	C7H8O2	guaiacol*	PTR	0.15±0. 09	0.093±0. 054	- 0.0013±0 .0073	0.052±0. 026	0.051±0. 020	0.053±0. 031	0.37±0.12 <sup>2</sup> 0.27±0.17 <sup>3</sup>	0.091± 0.037	0.12±0.07
124.183	C <sub>8</sub> H <sub>12</sub> O	acetylcycl ohexene *	PTR	0.018± 0.009	0.013±0. 005	0.00032± 0.0056	0.0078±0 .0045	0.0087±0 .0051	0.0068±0 .0039		0.015± 0.009	0.015±0.008
126.111	C <sub>6</sub> H <sub>6</sub> O <sub>3</sub>	hydroxy methylfu rfural*	PTR	0.033± 0.014	0.025±0. 010	0.00066± 0.0087	0.0096±0 .0053	0.0094±0 .0049	0.0098±0 .0056	$0.27\pm0.10^{2}$ $0.064\pm0.02$ $6^{3}$	0.016± 0.009	0.021±0.012
		unsaturat ed C6 cyclic										
126.155	C7H10O	carboxyli c acid*	CIMS	0.052± 0.005	0.043±0. 011	0.0088±0 .0096	0.012±0. 005	0.015±0. 0057	0.0087±0 .0044		0.026± 0.010	0.019±0.009
120.135		cyclohex ene	CINIS									0.01510.005
126.155	C7H10O	carboxyli c acid *	ptr	0.015± 0.006	0.013±0. 003	0.0026±0 .0091	0.0064±0 .0045	0.0080±0 .0056	0.0048±0 .0032		0.014± 0.010	0.010±0.007
126.199	C <sub>8</sub> H <sub>14</sub> O	octenone *	PTR	0.0064 ±0.002 9		0.00012± 0.0051	0.0027±0 .0016		0.0027±0 .0016			0.0057±0.003 3
120.199	C811140	trimethyl	FIN	5		0.0051	.0010		.0010			5
126.217	C7H10S	thiophen e* di	PTR	0.013± 0.004	0.011±0. 005	0.00044± 0.0084	0.011±0. 011	0.016±0. 015	0.0054±0 .0030		0.028± 0.026	0.012±0.007
128.127	C₀H8O3	hydroxy methyl furan*	PTR	0.024± 0.013	0.020±0. 007	- 0.0017±0 .0087	0.0044±0 .0026	0.0059±0 .0032	0.0029±0 .0017		0.010± 0.006	0.0063±0.003
	C7H12O	cyclohex anoic acid *		0.014±	007	0.0028±0	0.0050±0	.0032	0.0050±0		0.000	
128.171	2	naphthal	ptr	0.004 0.033±	0.025±0.	.011 0.00075±	.0034 0.017±0.	0.018±0.	.0034 0.015±0.	0.078±0.05	0.031±	0.010±0.007
128.174	C <sub>10</sub> H <sub>8</sub>	ene*	PTR	0.013	011	0.0086	011	012	009	6 <sup>2</sup>	0.021	0.031±0.018
128.215	C <sub>8</sub> H <sub>16</sub> O	octanone *	PTR	0.0072 ±0.003 5		0.0014±0 .0055	0.0028±0 .0020		0.0028±0 .0020			0.006±0.0042
130.187	C7H14O	amyl acetate *	ptr	0.0074 ±0.004 6	0.0056±0 .0021	0.00025± 0.0046	0.0031±0 .0019	0.0034±0 .0020	0.0028±0 .0018		0.0056 ±0.003 3	0.0058±0.003 6
132.159	C <sub>6</sub> H <sub>12</sub> O 3	C6 hydroxy- carboxyli c acids	CIMS	0.0090 ±0.000 6	0.010±0. 002	0.0062±0 .0014	0.0017±0 .0008	0.0027±0 .0011	0.00075± 0.00034		0.0045 ±0.001 8	0.0016±0.001 0
		methyl benzo		0.023±	0.016±0.	0.00018±	0.010±0.	0.010±0.	0.011±0.	0.055±0.03 0 <sup>2</sup> 0.046±0.02	0.017±	
132.162	C <sub>9</sub> H <sub>8</sub> O	furans*	PTR	0.0231	0.010±0. 0085	0.0050	006	0.01010.	006	1 <sup>3</sup>	0.009	0.021±0.012
		ethyl styrene/ methyl		0.010	0.01410	0.0012+0	0.0002+0	0.000010	0.000010	0.041±0.01 9 <sup>2</sup>	0.014	
132.206	$C_{10}H_{12}$	propenyl benzene*	PTR	0.019± 0.009 0.0074	0.014±0. 007	0.0013±0 .0063	0.0083±0 .0050	0.0083±0 .0050	0.0083±0 .0050	0.040±0.02 6 <sup>3</sup>	0.014± 0.008 0.0071	0.017±0.010
134.134	C <sub>8</sub> H <sub>6</sub> O <sub>2</sub>	phthalic acid*	PTR	±0.002 8	0.0071±0 .0025	0.0011±0 .0051	0.0039±0 .0029	0.0044±0 .0035	0.0033±0 .0022		±0.005 7	0.0065±0.004 2

Molecu- lar Weight	Comp- ound	Compou nd Name	Instru- ment	SP Averag e	NP Average	Backgr- ound	Average EF (g kg <sup>-1</sup> )	NP EF (g kg⁻¹)	SP EF (g kg <sup>-1</sup> )	Literature EF (g kg <sup>-1</sup> )	NP ER	SP ER
-												
		methylac				-				0.053±0.03 1 <sup>2</sup>		
		etopheno		0.012±	0.0085±0	0.00032±	0.0059±0	0.0062±0	0.0056±0	0.045±0.01	0.010±	
134.178	C <sub>9</sub> H <sub>10</sub> O	ne* C10	PTR	0.007	.0041	0.004	.0035	.0036	.0033	9 <sup>3</sup>	0.006	0.011±0.006
		Aromatic		0.030±	0.024±0.	0.0024±0	0.024±0.	0.035±0.	0.013±0.	0.040±0.02	0.058±	
134.222	C <sub>10</sub> H <sub>14</sub>	s*	PTR	0.013	010	.0095	014	019	008	1 <sup>3</sup>	0.031	0.026±0.016
		methyl								0.081±0.03 0 <sup>2</sup>		
		benzoic		0.027±	0.018±0.	0.00058±	0.013±0.	0.014±0.	0.012±0.	0.066±0.02	0.022±	
136.15	C <sub>8</sub> H <sub>8</sub> O <sub>2</sub>	acid *	PTR	0.015	009	0.007	007	0069	007	9 <sup>3</sup>	0.011	0.023±0.014
		monoter		0.68±0.	0.65±0.3	0.057±0.	0.41±0.1	0.49±0.2	0.33±0.1	0.41±0.06 <sup>1</sup> 0.87±0.72 <sup>2</sup>	0.80±0.	
136.238	C <sub>10</sub> H <sub>16</sub>	penes*	PTR	28	1	055	9	0	7	0.21±0.15 <sup>3</sup>	32	0.65±0.38
		hydroxyb		0.0080	0.0076.0	-	0.0000.0	0.0000.0			0.0061	0.0000.0000
138.122	C <sub>7</sub> H <sub>6</sub> O <sub>3</sub>	enzoic acid*	PTR	±0.003 0	0.0076±0 .0033	0.000093 ±0.0050	0.0026±0 .0017	0.0039±0 .0023	0.0014±0 .0008		±0.003 6	0.0028±0.001 5
	-, 0-3	creosol/		-							-	
139 166	C <sub>8</sub> H <sub>10</sub> O	methyl	DTD	0.016±	0.012±0.	0.000003	0.0073±0	0.0077±0	0.0069±0	0.14±0.11 <sup>3</sup>	0.012±	0.012+0.009
138.166	2	guiacol* isophoro	PTR	0.0093 0.027±	006 0.025±0.	8±0.0069 0.0079±0	.0046 0.0092±0	.0049 0.0086±0	.0042 0.0098±0	0.14±0.113	0.008 0.014±	0.013±0.008
138.21	C <sub>9</sub> H <sub>14</sub> O	ne*	PTR	0.0075	009	.01	.0064	.0053	.0074		0.009	0.019±0.014
		dimethyl benzofur								0.043±0.01		
		an/ethyl		0.0098		-				0.045±0.01 8 <sup>2</sup>	0.0078	
	$C_{10}H_{10}$	benzofur		±0.004	0.0072±0	0.00041±	0.0048±0	0.0052±0	0.0045±0	0.051±0.02	±0.005	0.0083±0.004
146.189	0	an*	PTR	4 0.0064	.0033	0.0037	.0029	.0034	.0024	8 <sup>3</sup>	1	4
				±0.003		0.0012±0	0.0034±0		0.0034±0			0.0061±0.004
146.233	$C_{11}H_{14}$	*	PTR	5		.0041	.0013		.0013			4
		cinnamic		0.0040 ±0.003		- 0.00094±	0.0021±0		0.0021±0			0.0037±0.002
148.161	C9H8O2	acid*	PTR	3		0.0048	.0013		.0013			2
				0.0047						0.027±0.01		
	C <sub>10</sub> H <sub>12</sub>	benzylac etone/es		0.0047 ±0.002	0.0043±0	0.00045±	0.0023±0	0.0022±0	0.0024±0	2 <sup>2</sup> 0.025±0.01	0.0033 ±0.002	0.0044±0.002
148.205	0	tragole*	PTR	1	.0019	0.0030	.0016	.0017	.0015	5 <sup>3</sup>	6	7
		C11								0.014+0.00		
		aromatic s/pentam		0.0074						0.014±0.00 8 <sup>2</sup>	0.0064	
		ethylben		±0.003		0.00054±	0.0041±0		0.0038±0	0.014±0.00	±0.004	0.0069±0.004
148.249	C <sub>11</sub> H <sub>16</sub>	zene* ethyl	PTR	2	.0027	0.0047	.0028	.0032	.0023	7 <sup>3</sup>	8	1
		benzoate		0.0059						0.14±0.08 <sup>2</sup>	0.0043	
	C <sub>9</sub> H <sub>10</sub> O	/vinyl		±0.002	0.0044±0	0.00039±	0.0028±0	0.0029±0	0.0028±0	0.036±0.02	±0.003	0.0049±0.003
150.177	2	guaiacol*	PTR	9 0.0040	.0020	0.0034	.0020	.0023	.0017	5 <sup>3</sup>	4 0.0039	0
	$C_{10}H_{14}$			±0.001	0.0039±0	0.00055±	0.0021±0	0.0027±0	0.0015±0		±0.003	0.0027±0.001
150.221	0	carvone*	PTR	2	.0017	0.0028	.0016	.0021	.0009	0.00710.01	0	6
		camphor								0.027±0.01 7 <sup>2</sup>		
	$C_{10}H_{16}$	/isomers		0.022±	0.023±0.	0.0063±0	0.011±0.	0.013±0.	0.0087±0	0.025±0.01	0.020±	
152.237	0	*	PTR	0.0074	010	.0086	007	008	.0066	4 <sup>3</sup>	0.011	0.015±0.011
	C <sub>9</sub> H <sub>14</sub> O	norborna neacetic		0.0036 ±0.001		- 0.00050±	0.0022±0		0.0022±0			0.0038±0.002
154.209	2	acid*	PTR	5		0.0041	.0012		.0012			0
		acenapht		0.0054 ±0.002	0.0040±0	- 0.00026±	0.0029±0	0.0033±0	0.0025±0		0.0046 ±0.004	0.0042±0.002
	1	acenapric	PTR	±0.002 0	0.0040±0 .0015	0.00026± 0.0031	0.0029±0 .0022	.0028	.0013		±0.004 0	0.0042±0.002

Molecu-	Comp-	Compou	Instru-	SP	NP	Backgr-	Average	NP EF (g	SP EF (g	Literature	NP ER	SP ER
lar	ound	nd Name	ment	Averag	Average	ound	EF (g kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )	EF		
Weight				е						(g kg <sup>-1</sup> )		
		terpine-										
		4-								0.0056±0.0		
		ol/cineol		0.0024		-				021 <sup>2</sup>	0.0029	
	$C_{10}H_{18}$	e/isomer		±0.000	0.0020±0	0.00044±	0.0018±0	0.0019±0	0.0017±0	0.0027±0.0	±0.002	0.0028±0.001
154.253	0	s*	PTR	9	.00068	0.0022	.0014	.0018	.0009	017 <sup>3</sup>	8	5
				0.0027						0.15±0.07 <sup>2</sup>		
		sesquiter		±0.001		0.00030±	0.0017±0		0.0017±0	0.029±0.02		0.0022±0.001
204.357	C15H24	penes*	PTR	1		0.0021	.0011		.0011	8 <sup>3</sup>		4
						5.2	1.4±0.03					
							7 to					
	C <sub>11</sub> to	I/SVOCs	cartrid				2.4±0.06					
239±61	C <sub>25</sub>	– CH	ge				3					
						4.6	0.81±0.0					
							78 to					
	C <sub>11</sub> to	I/SVOCs	cartrid				0.81±					
255±61	C <sub>25</sub>	- CHO1	ge				0.079					
						0.2	0.21±0.0					
							033 to					
	C <sub>11</sub> to	I/SVOCs	cartrid				0.22±0.0					
271±61	C <sub>25</sub>	- CHS <sub>1</sub>	ge				060					