# Assessing formic and acetic acid emissions and chemistry in western U.S. wildfire smoke: implications for atmospheric modeling

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# 1 Abstract

2 Formic acid (FA) and acetic acid (AA), two of the most abundant organic acids in the

atmosphere, are typically underestimated by atmospheric models. Here we investigate their

4 emissions, chemistry, and measurement uncertainties in biomass burning smoke sampled during

5 the WE-CAN and FIREX-AQ aircraft campaigns. Our observed FA emission ratios (ERs) and

6 emission factors (EFs) were generally higher than the 75<sup>th</sup> percentile of literature values, with

7 little dependence on fuel type or combustion efficiency. Rapid in-plume FA production was

8 observed (2.7 ppb  $ppm_{CO}^{-1}h^{-1}$ ), representing up to ~20 % of the total emitted carbon being

9 converted to FA within half a day.

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- 11 AA ERs and EFs showed good agreement with the literature, with little or no secondary
- 12 production observed within < 8 hours of plume aging. Observed FA and AA trends in the near-
- 13 field were not captured by a box model using the explicit Master Chemical Mechanism nor
- simplified GEOS-Chem chemistry, even after tripling the model's initial VOC conditions.
- 15 Consequently, the GEOS-Chem chemical transport model underestimates both acids in the

western U.S. by a factor of > 4. This is likely due to missing secondary chemistry in biomass 16

burning smoke and/or coniferous forest biogenic emissions. This work highlights uncertainties in 17 measurements (up to 100%) and even large unknowns in the chemical formation of organic acids

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in polluted environments, both of which need to be addressed to better understand their global 19

20 budget.

#### 21 **1** Introduction

Formic acid (FA) and acetic acid (AA) are the two most prevalent organic acids in the 22 troposphere, affecting aqueous-phase chemistry<sup>1</sup> and gas-aerosol partitioning<sup>2</sup> by regulating pH 23 levels in cloud droplets and aerosols.<sup>3–6</sup> Multiple studies have shown that various models 24 continuously underestimate both FA and AA abundance compared to ground, airborne, and 25 satellite observations. This low model bias is most pronounced in biogenic source regions,<sup>7,8</sup> 26 including United States (U.S.) deciduous forests,<sup>9,10</sup> boreal forests,<sup>11</sup> tropical forests,<sup>6</sup> and in the 27 Arctic tundra.<sup>12</sup> Additionally, models typically fail to capture FA and AA enhancements in 28 plumes from mixed anthropogenic sources<sup>13,14</sup> and in biomass burning (BB) impacted regions,<sup>15–</sup> 29 <sup>17</sup> indicating potential missing primary and/or secondary sources in smoke. In this work, we 30 investigate emissions, secondary productions, and model representations of FA and AA in the 31 western U.S. during two wildfire seasons, using measurements made during the WE-CAN 32 (Western Wildfire Experiment for Cloud Chemistry, Aerosol Absorption, and Nitrogen) and 33 FIREX-AQ (Fire Influence on Regional to Global Environments and Air Quality) field 34 campaigns. 35

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FA and AA are two of the most abundantly emitted volatile organic compounds (VOCs) from 37

BB, accounting for 16 % of the average VOC emissions by mass in western U.S. wildfires.<sup>18</sup> As 38

smoke plumes age, substantial secondary production of FA and AA may occur, <sup>19–23</sup> resulting in 39

these compounds together being one of the largest OH sinks in smoke aged more than 3 days, 40

accounting for up to ~25 % of plume OH reactivity.<sup>24</sup> Similarly, FA and AA can account for up 41

to 15 % of the VOC OH reactivity in urban atmospheres as well as the clean free troposphere in 42 the western U.S. during wildfire season.<sup>24</sup> 43

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Globally, top-down estimates suggest FA sources could reach 100–120 Tg y<sup>-1</sup>,<sup>6</sup> which is two to 45

three times higher than the sum of its known sources.<sup>6,8,16</sup> Of this, photochemical production 46

from biogenic sources has been estimated to contribute up to 90 % of the global FA budget.<sup>6</sup> 47 Biomass burning may account for up to 16 Tg  $y^{-1}$  (~13–16 %) of FA globally.<sup>16</sup> though such 48 estimates for BB are mostly based on direct emissions. The secondary production of FA from BB 49 precursors is poorly known due in part to the high uncertainty in BB emissions and a large 50 amount of reactive BB precursors, such as furan containing species, not being implemented in 51 current chemical transport models (CTMs).<sup>24</sup> Similarly, global AA sources have been estimated 52 using a bottom-up approach to be 85 Tg  $y^{-1}$ ,<sup>8</sup> which is likely a lower bound.<sup>7</sup> Despite BB being a 53 major source of AA,<sup>23-26</sup> the contribution of BB to the global AA budget is rarely discussed in 54 the literature and is not 55

56 well constrained.

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The primary sinks of atmospheric FA and AA include wet and dry deposition, photochemical oxidation by OH radicals, and the irreversible uptake on dust resulting in atmospheric lifetimes of 2–4 days for FA and ~2 days for AA.<sup>6–8,27</sup> Consequently, their relatively short atmospheric lifetimes coupled with the localized and seasonal nature of fires likely means BB alone cannot close the global FA and AA budgets.<sup>8</sup> However, in regions heavily impacted by BB it is likely that fires play an important role in their regional abundance and a more detailed understanding of their emissions and chemistry in wildfire smoke is needed.

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66 As the two simplest organic acids, FA and AA may be produced from the oxidation of many

67 different VOCs and are known photochemical products of isoprene, terminal alkenes,

68 monoterpenes, glycolaldehyde, aromatics, acetone, and acetaldehyde.<sup>7,8,14,28–31</sup> Heterogeneous

69 formation of FA in aerosols and cloud droplets has also been identified as a potential major

source, which, when included in the global chemistry-climate model ECHAM5/MESSy

71 (EMAC), has been found to largely reconcile the global FA budget.<sup>32,33</sup> However, regional

72 discrepancies remain. For example, FA abundances were still underestimated in boreal forested

regions, likely due to low emissions of FA precursors from BB.<sup>32</sup>

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Analytical challenges measuring FA and AA,<sup>34</sup> along with an incomplete understanding of

chemical processes in smoke,<sup>35</sup> has made it difficult to accurately model their evolution in BB

plumes.<sup>16,36</sup> For example, it has been well documented that the GEOS-Chem CTM underpredicts

FA and AA abundances. Missing secondary production from biogenic precursors is thought to be 78 one of the most significant reasons for the low model bias,<sup>7</sup> though in some ecosystems there 79 may still be missing primary emissions and/or in-canopy sources.<sup>11,37</sup> The overall model sink 80 may also be too large.<sup>38</sup> By updating the model chemistry to reflect photochemical FA 81 production from alkynes, monoterpenes, isoprene, methyl peroxy radical (CH<sub>3</sub>O<sub>2</sub>), ozonolysis of 82 terminal alkenes, keto-enol tautomerization, and phototautomerization of acetaldehyde,<sup>7,8,39</sup> Chen 83 et al.<sup>17</sup> were able to improve GEOS-Chem representation of the remote free troposphere relative 84 to observations during ATOM. Despite the updated chemistry, the model underestimated the 85 median FA:CO ratio by a factor of >2 and the 95th percentile by a factor of >4,  $^{17}$  suggesting that 86 there are still significant missing secondary sources in smoke. 87

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89 In this work, we examine FA and AA emissions and chemistry in wildfire smoke to better understand the role of BB in their regional budgets. Using observations from the WE-CAN and 90 91 FIREX-AQ aircraft campaigns, we first assess FA measurements made by two commonly used chemical ionization mass spectrometers, PTR-ToF and I<sup>-</sup> CIMS (proton-transfer-reaction time-92 93 of-flight mass spectrometer and iodide adduct chemical-ionization mass spectrometer). Emissions for FA and AA are then compared with literature values before examining their 94 95 chemistry during WE-CAN in five pseudo-Lagrangian sampled smoke plumes. Finally, we assess GEOS-Chem representation of both acids across two fire seasons, first using observations 96 97 made during the WE-CAN field campaign and then FIREX-AQ as an additional test for year-toyear variability and regional representativeness. 98

#### 99 **2 Methods**

#### 100 2.1 WE-CAN and FIREX-AQ campaign overviews and sampling approach

Comprehensive gas and aerosol measurements were made in wildfire smoke plumes across seven
western U.S. states from 24 July to 13 September 2018 during the WE-CAN aircraft campaign
(https://www.eol.ucar.edu/field\_projects/we-can). *In situ* smoke plume sampling was carried out
aboard the NSF/NCAR C-130 research aircraft based out of Boise, ID, typically between 14:00
and 19:00 local time when burning conditions were most active. Figure S1 depicts the C-130

106 flight tracks during WE-CAN, colored by the observed formic and acetic acid mixing ratios.

Upon arriving at a fire, the C-130 would typically sample fire emissions by flying perpendicular 107 transects through the plume, as near to the source as was allowed by firefighting operations and 108 plane safety constraints. To investigate plume aging, most plumes were subsequently sampled 109 using a pseudo-Lagrangian approach where perpendicular transects were performed in a stepwise 110 pattern starting near a fire and continuing as far downwind as possible (seen as the zig-zag flight 111 pattern in Figure S1). In total, WE-CAN sampled more than 22 hours of wildfire smoke, 112 including 31 emission transects of 24 unique fires<sup>18</sup> and 1.2 hours of smoke estimated to have 113 aged >3 days, along with 4.8 hours of the clean free troposphere.<sup>24</sup> 114 115

The FIREX-AQ aircraft campaign sampled BB plumes across the western and southeastern U.S. 116 from 22 July to 5 September 2019 (https://csl.noaa.gov/projects/firex-aq) following a similar 117 sampling approach as WE-CAN.<sup>40</sup> In this work, we separate FIREX-AQ data into its western and 118 southeastern U.S. portions (FIREX-AQ-W and FIREX-AQ-SE), delimitated by the 105th 119 meridian west, for a more accurate regional comparison. This allows us to assess FA and AA 120 representation in the GEOS-Chem CTM across multiple fire seasons and regions. As the total 121 122 VOC emissions in the western U.S. during the 2018 WE-CAN campaign were  $\sim 10 \times$  higher, with  $\sim 2 \times$  more area burned, than during the 2019 FIREX-AQ campaign (190 GgC vs. 20 GgC, 123  $3.5 \times 10^6$  ha vs.  $1.9 \times 10^6$  ha), <sup>41,42</sup> these two datasets provide complementary representation of a 124 wide range of seasonal fire activity allowing the model to be assessed under varying real-world 125 126 conditions.

# 127 **2.2 Measurements of formic acid, acetic acid, and organic aerosol**

128 FA and AA were both measured by two different proton-transfer-reaction time-of-flight mass

- 129 spectrometers (PTR-ToF) and an iodide adduct chemical-ionization mass spectrometers (I-
- 130 CIMS) during the WE-CAN and FIREX-AQ campaigns. The PTR-ToF<sup>18</sup> and I<sup>-</sup>CIMS<sup>43–46</sup>

operated during WE-CAN and FIREX-AQ<sup>47,48</sup> and referenced in this work have been extensively

- described by the cited literature, while here we include those details most relevant to their
- 133 measurements of FA and AA.
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- 135 Organic aerosol (OA) was measured by high-resolution aerosol mass spectrometry (HR-AMS;
- 136 Aerodyne Inc.), described in detail by Garofalo et al.<sup>49</sup> During WE-CAN, the HR-AMS

measured OA with 5 s time resolution, vacuum aerodynamic diameter of  $\sim$ 70–1,000 nm, and uncertainty of 35%. In this work we primarily use the fractional component of OA attributed to

the  $CO_2^+$  ion (f<sub>44</sub>), an OA oxidation marker.<sup>50,51</sup> For plume transects, f<sub>44</sub> averages are weighted

140 by the measured OA mass. $^{52}$ 

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#### 142 **2.2.1 PTR-ToF**

During WE-CAN, the PTR-ToF measured at 2 or 5 Hz frequency with drift tube conditions
maintained at 3.00 mbar, 810 V, and 60°C, resulting in an E/N of 130 Td. Sampling was done by
drawing ambient air into the cabin at 10–15 lpm through ~3 meters of 3.175 mm I.D.

146 perfluoroalkoxy (PFA) tubing, maintained at  $\sim$ 55°C. This sample stream was then subsampled

by the PTR-ToF drift tube through  $\sim 100$  cm of 1.588 mm O.D. PEEK tubing (60°C), resulting in

a total inlet residence of less than 2 s. Three-minute instrument zeroes were performed every

hour by sampling VOC free air generated via a platinum bead catalyst heated to 375°C.

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Calibrating FA and AA is analytically challenging due to their instability in gas standards and 151 known humidity-dependent sensitivities in PTR-ToF measurements.<sup>53</sup> To overcome these 152 challenges, humidity-dependent FA and AA sensitivities were determined in the laboratory post-153 154 campaign using a commercial liquid calibration unit (LCU; Ionicon Analytik). Analytical grade FA and AA were volatilized in the LCU and dynamically diluted into zero air where the 155 156 humidity was varied within the range observed during WE-CAN as determined by the internal humidity proxy of H<sub>2</sub>O•H<sub>3</sub>O<sup>+</sup> to H<sub>3</sub>O<sup>+</sup> ([m/z 39]/[m/z 21], 0-6 %).<sup>53,54</sup> The resulting calibration 157 curves for FA and AA sensitivities as a function of the percent [m/z 39]/[m/z 21] are shown in 158 Figure S2 and applied to all WE-CAN FA and AA PTR-ToF measurements. Note that similar 159 160 calibrations and humidity corrections were applied to PTR-ToF measurements during FIREX-AQ. Over the 0–6 % [m/z 39]/[m/z 21] range, sensitivities for both species were observed to 161 decrease with increased humidity, ranging ~9–4 ncps/ppb, similar to the sensitivity change 162 reported by Baasandorj et al.<sup>53</sup>. During WE-CAN the PTR-ToF FA and AA uncertainties are 163 conservatively estimated as 50 %, mostly due to 40 % potential instrument drift between WE-164 CAN and the laboratory calibrations as determined from the observed instrument sensitivity 165

change of other gas standards. The detection limits are 1.0 ppb for FA and 0.5 ppb for AA for 166

our 1 Hz measurements, defined as  $3\sigma$  for the inflight instrument zeros. 167

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In PTR-ToF, FA (HCOOH) and AA (CH<sub>3</sub>COOH) are detected at their protonated masses, m/z169 47.013 and m/z 61.028 respectively. The corresponding mass resolution during WE-CAN is 2120 170  $m/\Delta m$  at m/z 47 and 3060  $m/\Delta m$  at m/z 61, where  $\Delta m$  is the full width at half maximum for the 171 ion peak. FA has three major potential interfering ions: dimethyl ether (DME, m/z 47.077), 172 ethanol (m/z 47.050), and N<sub>2</sub>H<sub>3</sub>O<sup>+</sup> (m/z 47.024).<sup>25,53</sup> The mass resolution during WE-CAN was 173 high enough to separate DME and ethanol signals from FA, with ethanol abundance also 174 expected to be  $\sim 4 \times \text{lower}$  than FA in BB smoke with an instrumental sensitivity  $\sim 10 \times \text{lower}$ 175 than FA.<sup>25</sup> The  $N_2H_3O^+$  signal, which was not fully resolved from FA, was observed to stay 176 177 constant regardless of emission source strength throughout the campaign and was therefore classified and corrected as instrumental background. Consequently, we treat the m/z 47 signal as 178 being primarily FA in agreement with previous literature.<sup>53–55</sup> 179 180 181 Potential interferences of AA in PTR-TOF measurements include 2-propanol and n-propanol (m/z 61.065), peroxyacetic acid (PAA) fragments (m/z 61.028), ethyl acetate fragments (m/z 61.028)182 61.028), methyl formate (m/z 61.028), and glycolaldehyde (m/z 61.028).<sup>25,53,55-59</sup> Propanol was 183 resolved from AA during WE-CAN, while PAA fragments, ethyl acetate fragments, methyl 184 185 formate, and glycolaldehyde are all isomeric with AA. PAA is formed by the reaction of CH<sub>3</sub>C(O)O<sub>2</sub> radicals with HO<sub>2</sub>, which may be important in low NO<sub>x</sub> conditions<sup>53</sup> but is ~100 × 186

less abundant than AA in fresh BB smoke (~ $20 \times \text{less}$  abundant after 1.5 hours aging)<sup>22</sup> making 187 its fragment unlikely to be a significantly contributor to m/z 61.<sup>25</sup> Ethyl acetate is used in

coatings, adhesives, cosmetics, and as a process solvent,<sup>60</sup> resulting in it being most prevalent in 189

190 anthropogenically polluted areas, while it has not been reported in significant quantities in BB

emissions.<sup>25,61</sup> For methyl formate, a small peak can be seen in the GC-MS during the FIREX-191

AQ laboratory burning experiment, but FTIR comparison suggests its contribution is 192

negligible.<sup>25</sup> Due to interference from these isomers being minimal in BB smoke, we do not 193

attempt to correct for their presence, and assume m/z 61 to be predominantly AA and 194

glycolaldehyde in wildfire emissions.<sup>62</sup> 195

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Based on the FIREX-AQ Missoula fire laboratory burning experiments, the m/z 61 signal is on 197 average 67 % AA and 33 % glycolaldehyde ( $\pm$  45 % of value) in fresh BB emissions.<sup>25,63</sup> 198 However, the glycolaldehyde contribution in aged smoke is not well described. As 199 glycolaldehyde's atmospheric lifetime of 1 day<sup>64</sup> is approximately half of that of  $AA^8$ , it is likely 200 that m/z 61 becomes more predominantly AA in aged air masses, though glycolaldehyde 201 production could offset its loss. Given that the relative contribution of glycolaldehyde to m/z 61 202 was not constrained during WE-CAN, we do not attempt to correct for the potential 203 204 glycolaldehyde interference and apply only the humidity dependent AA sensitivity to m/z 61. Though we treat and discuss the PTR-ToF m/z 61 as AA in this work, the reported values reflect 205 the combined AA and glycolaldehyde isomers and therefore likely represent an upper bound for 206 AA. 207

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#### 209 **2.2.2 I**<sup>-</sup> CIMS

<sup>210</sup> I<sup>-</sup> CIMS operates by colliding iodide ions (I<sup>-</sup>) with neutral analytes inside an ion-molecule <sup>211</sup> reaction region (IMR), forming clusters which are then analyzed by a time-of-flight mass <sup>212</sup> spectrometer. During WE-CAN, ambient air was sampled at 20 lpm through a 40 cm long, 18 <sup>213</sup> mm O.D. PTFE tube before being subsampled into the IMR. Between the inlet and IMR, the <sup>214</sup> residence time was < 0.7 seconds. Humidity in the IMR was controlled to maintain a constant <sup>215</sup> iodide-water (m/z 145) to iodide (m/z 127) ratio thereby reducing the instruments humidity <sup>216</sup> dependence.

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The I<sup>-</sup> CIMS employed a fast-zeroing approach described in Palm et al.<sup>45</sup> where background 218 concentrations were found by sampling ultra-high purity N<sub>2</sub> into the IMR for 6 seconds every 219 minute. The fast zeros were used to determine the background-subtracted signal by isolating the 220 effects of adsorption and desorption of 'sticky' molecules on the internal IMR surfaces. This 221 zero occurred both in and out of smoke plumes to account for the changes in background signal 222 with varying sampled concentrations. A full inlet zero was also performed for 10 seconds every 223 20 minutes to determine the combined background signal from inlet tubing plus IMR surfaces, 224 which confirmed that the dominant source of the background was from the IMR and not inlet 225 tubing.45 226

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The I<sup>-</sup> CIMS detects FA as a cluster with iodide at m/z 172.911. For WE-CAN, FA was 228 calibrated in the laboratory prior to the campaign by flowing pure air over heated permeation 229 tubes with gravimetrically determined permeation rates. Although I<sup>-</sup> CIMS measures AA, due to 230 its low sensitivity and apparent interference from an unknown compound during WE-CAN, we 231 only report AA from PTR-ToF. Recent work has shown that one potential source of uncertainty 232 for the I<sup>-</sup>CIMS FA measurement is that its sensitivity to FA decreases with increasing IMR 233 temperature.<sup>48</sup> The I<sup>-</sup> CIMS deployed during WE-CAN did not directly regulate temperature in 234 the IMR. During smoke sampling periods the C-130 cabin temperatures measured near the I<sup>-</sup> 235 CIMS ranged from 20–32 °C (10<sup>th</sup> and 90<sup>th</sup> percentiles: 22–26 °C). Robinson et al.<sup>48</sup> showed that 236 a 10 °C change in IMR temperature could correspond to a 50 % change in sensitivity, though 237 238 differences in pressures and tuning between instruments makes applying this uncertainty to the WE-CAN deployment highly uncertain. It is also unlikely that the IMR temperature fluctuated as 239 240 widely as the cabin temperature, though a lack of data makes it difficult to constrain the actual IMR temperature during WE-CAN or during the laboratory calibrations. Consequently, we 241 242 conservatively estimate the I<sup>-</sup> CIMS FA measurement to be a likely upper bound, with 60 % uncertainty and 30 ppt detection limit for 1 Hz data, based on calibration uncertainties and 243 244 potential variation in IMR temperature.

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### 246 **2.2.3 FIREX-AQ**

the PTR-ToF and I<sup>-</sup>CIMS deployed during FIREX-AQ had a few notable configuration 247 differences relative to ones used during the WE-CAN deployment. For the PTR-ToF, the inlet 248 was only  $\sim 1$  m in length and was comprised of 3.175 mm I.D. PTFE heated to  $\sim 50-60$  °C. The 249 residence time is estimated as less than 1s. The PTR-ToF was calibrated against FA and AA 250 using total carbon methods as described by Veres et al.<sup>65</sup> with similar humidity dependencies 251 determined following the methods described in Section 2.2.1. The FIREX-AQ PTR-ToF 252 instrument uncertainty is 30 % for FA and 50 % for AA. Note that in this work, we remove all 253 FIREX-AQ PTR-ToF FA observations above ~4.9 km ASL (above sea level, <550 hPa) from 254 our analysis due to a known background issue in the high-altitude FA measurements. 255

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For the I<sup>-</sup>CIMS, ambient air was sampled at 6 slpm through a mass-flow-controlled PFA inlet 257 (70 cm length, 6.4 mm I. D.) maintained at 40 °C. A pressure control region upstream of a 258 critical orifice at the entrance to the IMR was maintained at 140 mbar, and thus a constant flow 259 of 1.2 slpm ambient air entered the IMR to mix with the 1 slpm ion source flow. Similar to the I-260 CIMS deployed during WE-CAN, the IMR was humidified to minimize instrument humidity 261 dependence. The instrument background signal was determined in flight by overflowing the inlet 262 with scrubbed ambient air for 30 s every 10 min through a port located 2 cm downstream of the 263 inlet entrance.<sup>47</sup> IMR temperature was not controlled during FIREX-AO, but a post campaign 264 temperature correction was applied to the data as described in Robinson et al.<sup>48</sup> The instrument 265 uncertainty for FA is 15% + 30 ppt with a 3-sigma detection limit of 6 ppt. 266

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#### 268 **2.3 GEOS-Chem chemical transport model**

GEOS-Chem nested grid simulations (version 12.1.1)<sup>66,67</sup> over North America were run for the 269 WE-CAN and FIREX-AQ periods using the model conditions described in Chen et al.<sup>17</sup> 270 Simulations were carried out using Goddard Earth Observation System Forward Processing 271 (GEOS-FP) assimilated meteorology data with detailed HO<sub>x</sub>, NO<sub>x</sub>, VOC, ozone, halogen, and 272 aerosol chemistry. Model runs were conducted at  $0.25^{\circ} \times 0.3125^{\circ}$  (~25 km) resolution with time 273 steps of 5 min (transport/convection) and 10 min (chemistry/emission). Emissions follow Chen 274 et al.<sup>17</sup> with the notable exception that we use Global Fire Assimilation System version 1.2 275 (GFAS) BB emissions with FA and AA emission ratios updated based on Permar et al.,<sup>18</sup> which 276 in turn reflect the WE-CAN averages discussed in Section 4. GEOS-Chem was subsequently 277 278 sampled along both campaign flight tracks for comparison to the observations.

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The GEOS-Chem simulations also reflect updated FA chemistry including photochemical FA production based on OH initiated oxidation of alkynes, monoterpenes, isoprene, and CH<sub>3</sub>O<sub>2</sub>, ozonolysis of terminal alkenes (e.g. ethene, propene, isoprene), keto-enol tautomerization,<sup>7,8</sup> and phototautomerization of acetaldehyde.<sup>39</sup> The model does not included the aerosol chemistry proposed by Franco et al.<sup>32</sup> Based on these updates, Chen et al.<sup>17</sup> found that GEOS-Chem accurately simulated FA concentrations in the remote free troposphere during the Atmospheric Tomography (ATom) aircraft campaign, indicating that GEOS-Chem is not missing any

- significant FA sources in the remote free troposphere. The model was found to significantly
- underestimate FA mixing ratios in 1–10 day aged plumes attributed to both anthropogenic and
- BB sources. In this work, we investigate how well GEOS-Chem, with the Chen et al.<sup>17</sup> treatment
- of FA and AA chemistry, represents these acids in the western U.S. under heavily smoke
- 291 impacted conditions.

#### **3 Formic acid measurement intercomparison**

- 293 Formic acid is analytically challenging to measure due to its 'stickiness' in sample inlets and its
- humidity/temperature dependent sensitivities in PTR-ToF $^{34,53}$  and I<sup>-</sup> CIMS $^{43,48}$ . Figure 1 shows
- 295 the 1 Hz time series and cumulative mixing ratios of FA measured by PTR-ToF and I<sup>-</sup> CIMS
- during five plume transects (< 20 km downwind) of the Taylor Creek (TC) fire sampled during
- 297 WE-CAN (Research Flight #3). When corrected for inlet residence times, the two measurements
- show good temporal agreement, capturing the real-time plume variability. However, the PTR-
- 299 ToF consistently measures  $\sim 2 \times$  lower maximum FA concentrations than the I<sup>-</sup> CIMS during the
- 300 plume transects likely representing sample retention in the inlet, a baseline offset due to
- 301 background correction differences, and/or calibration errors.
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Figure 1. Time series of 1 Hz PTR-ToF and I<sup>-</sup> CIMS formic acid mixing ratios (bottom panel)
 and cumulative mixing ratios for each plume through the following background period (top
 panel) during 5 plume transects made < 20 km downwind from the Taylor Creek Fire, OR during</li>
 WE-CAN.

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The TC fire was sampled shortly after injection into the free troposphere with little to no regional 309 310 smoke impacts, resulting in clearly defined plume edges that can be seen in Figure 1 by the rapid FA enhancement upon entry into the smoke. However, when exiting the plumes, the PTR-ToF 311 trace shows a distinct tail indicative of FA being initially retained in the inlet before flushing out 312 in the 60–90 seconds after returning to background air. This is further illustrated by the upper 313 314 panel in Figure 1, where the cumulative mixing ratios for each plume through the subsequent background sampling periods are shown for both instruments. For all plumes shown in Figure 315 316 the I<sup>-</sup>CIMS and PTR-ToF integrated FA mixing ratios agree within <50 % after accounting for residual FA in the inlet. This indicates that the two measurements agree within their stated 317 318 uncertainty given sufficient time to recapture FA from the PTR-ToF inlet. However, Plumes 2 and 3 also demonstrate how FA may wash out of the inlet and increase the signal in subsequent 319 transects. Due to most other sampling periods having either more poorly defined plume edges, 320 elevated background signals from regional smoke, and/or not having enough time between 321

consecutive transects, we are unable to accurately extend this analysis to other fires. It is likely
 though that inlet retention decreases the maximum PTR-ToF measured FA in most plumes
 sampled during WE-CAN.

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The lack of a similar inlet artifact in the I<sup>-</sup> CIMS measurement is likely explained by a few 326 characteristics. First, inlet sizes and materials differ slightly between the two instruments. The I<sup>-</sup> 327 CIMS inlet is significantly shorter than the one used by the PTR-ToF, resulting in a shorter 328 residence time in I<sup>-</sup> CIMS (< 0.7 vs. 2 s). The I<sup>-</sup> CIMS inlet was also comprised of only PTFE, 329 while the PTR-ToF used PFA and PEEK tubing. Second, the I<sup>-</sup> CIMS fast zeroing strategy (seen 330 as the data gaps in the I<sup>-</sup> CIMS trace in Figure 1) results in a FA background-subtracted signal 331 that minimizes the effects of adsorption and desorption from walls and surfaces in the 332 instrument.<sup>45</sup> Consequently, this points to the importance of the instrument's inlet configuration 333 and background correction procedures for the most accurate measurement of FA in environments 334 335 with rapid concentration changes.

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337 Although inlet retention explains a large part of the disagreement of FA measured by the two instruments while sampling smoke plumes with high, rapidly changing concentrations, the 338 339 average flight integrated I CIMS ( $120 \pm 61$  ppm) to PTR-ToF ( $66 \pm 28$  ppm) formic acid ratio of 2.1 and total least squares regression (TLS; slope = 2.06,  $r^2 = 0.82$ ), indicates that the I<sup>-</sup> CIMS 340 341 measured  $\sim 2 \times$  more FA than the PTR-ToF across all research flights (Figure S3). Although the exact reason for this disagreement is unknown, it likely stems from both a baseline offset and 342 calibration uncertainty. For example, the PTR-ToF inflight zeros may have contained residual 343 FA due to desorption from the instrument/inlet surfaces and/or incomplete oxidation in the 344 catalyst- generated zero air. Consequently, an excessive background signal was subtracted, 345 346 resulting in the mixing ratios being biased slightly low, especially when sampling relatively clean air. 347

348

The FA sensitivity in I<sup>-</sup> CIMS is also strongly dependent on IMR temperature,<sup>48</sup> which was not directly controlled or logged during WE-CAN. The IMR used during WE-CAN<sup>45</sup> was different than the IMR used in Robinson et al.,<sup>48</sup> with each also operated at different pressures (100 mbar vs 40 mbar). Although variability in IMR temperature may influence the I<sup>-</sup> CIMS FA sensitivity during WE-CAN, instrument differences likely change the temperature dependence between

studies in ways that have not been tested. Consequently, the full extent of temperature effects on

355 the I<sup>-</sup> CIMS sensitivity during WE-CAN is unknown and future work should focus on controlling

356 IMR temperature while further characterizing the sensitivity dependence on temperature under

357 different instrument conditions.

358

We repeat a similar analysis with FIREX-AQ FA measurements and find that in contrast to WE-359 360 CAN, the PTR-ToF generally measured slightly lower FA mixing ratios than the I CIMS (TLS slope = 0.87,  $r^2 = 0.78$ ) for all FIREX-AQ data at altitudes below ~4.9 km, with the two agreeing 361 well within their stated instrument uncertainty. When compared between regions, the FIREX-362 AQ-W FA measurements shows better agreement (slope = 0.89,  $r^2 = 0.82$ ) than FIREX-AQ-SE 363 (slope = 0.69,  $r^2 = 0.67$ ), possibly representing uncertainty in the instrument sensitivity due to the 364 higher humidity typical of the southeastern U.S. relative to the western U.S. (Figure S3). 365 Additionally, neither instrument shows significant inlet retention, with FA mixing ratios for both 366 instruments generally returning to background levels at the same rate after exiting plumes 367 368 (Figure S4).

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370 The observed disagreement between the two instruments during WE-CAN is likely due to a combination of factors including uncertainty in the FA sensitivity due to its humidity and 371 372 temperature dependence, potential instrument drift between the laboratory calibrations and field measurements, inlet losses, and differences in background correction procedures. FIREX-AQ FA 373 measurements largely corrected for these issues by using a shorter PTR-ToF inlet along with 374 correcting for the I<sup>-</sup> CIMS IMR temperature variations. Consequently, to improve future FA 375 measurements made by I<sup>-</sup> CIMS and PTR-ToF special attention should be given to shortening 376 377 inlet residence times and minimizing sensitivity dependencies on temperature and humidity. 378

In this work, we primarily use I<sup>-</sup> CIMS FA measurements for most analysis and discussion due to its lower detection limits and lack of apparent inlet artifacts. This is likely an upper bound for WE-CAN data and therefore we include PTR-ToF FA observations to further constrain the measurement uncertainty where appropriate. We note that despite the high uncertainty in the FA measurements (up to 100%), model underestimates discussed in Section 6 are much greater than
 the measurement uncertainty.

#### **4 Emissions of formic and acetic acid from WE-CAN sampled fires**

Emission ratios (ERs) and emission factors (EFs) were calculated for 31 WE-CAN emission 386 transects of 24 individual fires as described in Permar et al.<sup>18</sup> Here, emission transects are 387 defined as the nearest transect of a well-defined smoke plume that is traceable to a single 388 389 emission source sampled 27–130 minutes downwind from the fire, as calculated by wind speeds measured aboard the C-130 and fire locations reported by the U.S. Forest Service.<sup>18</sup> Although 390 these transects represent the freshest smoke sampled during the campaign, this is sufficient time 391 for substantial secondary formation to have occurred.<sup>19,46,68,69</sup> Consequently, FA and AA ERs 392 and EFs during WE-CAN represent their combined production and loss before being sampled by 393 the research aircraft, which may be more appropriate for the spatial and temporal resolution of 394 many CTMs.<sup>70</sup> Normalized excess mixing ratios (NEMRs) were calculated using the background 395 corrected plume integrated mixing ratios of a VOC to CO (ppb VOC ppm<sub>CO</sub><sup>-1</sup>) for each emission 396 and subsequent downwind plume transect. ERs were used to calculate EFs, expressed as grams 397 of VOC emitted per kilogram of burned fuel, using the carbon mass balance method<sup>23,71</sup> with the 398 total emitted carbon as the sum of CO<sub>2</sub>, CO, CH<sub>4</sub>, organic carbon, black carbon, and 161 399 VOCs.<sup>18</sup> Fuels burned during WE-CAN were primarily those characteristic of mixed conifer 400 forests. 401

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Literature values were compiled from 16 different papers, reporting 330 FA and 289 AA ERs 403 and EFs.<sup>72</sup> Approximately half of the EFs were also recorded and retrieved from the Smoke 404 Emissions Reference Application (SERA), which may include some additional values 405 recalculated to match their fuel types.<sup>73</sup> Average FA and AA ERs and EFs for the literature 406 described in this work are summarized in Table S1 and represent a variety of burned fuels. Pre 407 408 2007 FA data measured by Fourier transform infrared spectroscopy (FTIR) have also been corrected by a factor of 2.1 following Yokelson et al.<sup>74</sup> Similar to vegetation classifications in 409 global BB emissions inventories, we broadly categorize these ERs and EFs as conifer forest (147 410 FA and 122 AA EFs), mixed hardwood forest (17, 23), shrubland (53, 38), grassland (36, 31), 411 crop residue (46, 47), and organic soil/peat (31, 28). Table S1 also includes modified combustion 412

efficiency (MCE) when available, instrumentation used, region of fuels burned, and whether the
data are from a laboratory or field study.<sup>19,20,23,25,26,34,75–78</sup>

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Figure 2 shows FA and AA ERs for each of the 24 fires sampled during WE-CAN (green 416 points), 16 fires sampled during FIREX-AQ, and literature values for all fuel types in our review 417 (box-and-whisker plots). During WE-CAN, the average formic acid ER calculated from I<sup>-</sup> CIMS 418 data was found to be  $9.5 \pm 4.2$  ppb ppm<sub>CO</sub><sup>-1</sup> (Table 1), which is 3.5 times higher than the 419 literature average of 2.7  $\pm$  2.6 ppb ppm<sub>CO</sub><sup>-1</sup> calculated from 168 data points reported for 10 of the 420 16 studies in Table S1. We note that although FA ERs calculated from PTR-ToF measurements 421 (average 6.6  $\pm$  2.5 ppb ppm<sub>CO</sub><sup>-1</sup>) are slightly lower than those from I<sup>-</sup> CIMS, both are generally 422 higher than the 75<sup>th</sup> percentile of literature values. Similarly, though FA ER during FIREX-AQ 423 (average  $3.31 \pm 2.0$  ppb ppm<sub>CO</sub><sup>-1</sup>) are lower than during WE-CAN, half are still above the 75<sup>th</sup> 424 percentile of the literature. Additionally, constraining ERs to only western U.S fuels has little 425 426 effect on this comparison as discussed in more detail below.

427

428 One possible explanation for the higher ERs observed during WE-CAN and FIREX-AQ is that a significant amount of FA has been produced in the plumes prior to being intercepted by the C-429 430 130 (Section 5). To approximate how much FA may have been formed before being sampled, we estimate t<sub>0</sub> emission ratios from the least squares regression of WE-CAN NEMRs vs. physical 431 432 age for three of the five pseudo-Lagrangian sampled smoke plumes discussed in Section 5. Assuming a constant production rate, projected FA NEMRs from I CIMS measurements at  $t_0$ 433 range from 5.7–7.4 ppb  $ppm_{CO}^{-1}$ , which is still approximately 2–3 times higher than the literature 434 average. Consequently, while many of the FA ERs measured during WE-CAN likely reflect 435 some plume aging (which, although hard to quantify, may also be the case in literature values), 436 437 near-field production alone is not enough to explain the disagreement. Given that WE-CAN ERs calculated using both I<sup>-</sup> CIMS (9.5 ppb ppm<sub>CO</sub><sup>-1</sup>) and PTR-ToF (6.6 ppb ppm<sub>CO</sub><sup>-1</sup>) measurements 438 agree within their stated uncertainty, it is likely that the ERs observed during WE-CAN generally 439 represent higher FA emissions from the wildfires sampled that season than the literature average 440 (Figure S6). As FIREX-AQ FA ERs are also generally higher than the literature, this may in part 441 reflect the bias of these two datasets towards sampling relatively large wildfires, which could 442 produce different FA emissions than laboratory burns and the smaller fires predominantly 443

represented in the literature. We recommend that future studies report their estimated aging when 444 reporting FA ERs. 445

- 446
- In contrast, AA ERs measured during WE-CAN and FIREX-AQ mostly fall within the 25th-75th 447

percentiles of literature values (Figure 2 and Table 1), with good agreement between their 448

- averages (WE-CAN 11.5  $\pm$  2.1 ppb ppm<sub>CO</sub><sup>-1</sup>, FIREX-AQ 8.9  $\pm$  1.5 ppb ppm<sub>CO</sub><sup>-1</sup>, literature 15.5  $\pm$ 449
- 14.2 ppb  $ppm_{CO}^{-1}$ ). 450
- 451
- Figure S5 shows that FA EFs follow the same trend as the ERs, with the WE-CAN average EF 452
- of 1.5  $\pm 0.60$  g kg<sup>-1</sup> (PTR-ToF = 0.96  $\pm 0.39$  g kg<sup>-1</sup>) and FIREX-AQ average EF of 0.6  $\pm 0.42$  g 453
- kg<sup>-1</sup> approximately 5 and 2 times higher than the literature average of  $0.35 \pm 0.48$  g kg<sup>-1</sup> 454
- Similarly, both campaign AA EFs are within the 25<sup>th</sup>-75<sup>th</sup> percentile of literature values, with 455
- good agreement between their averages (WE-CAN 2.4  $\pm$  6.1 g kg<sup>-1</sup>, FIREX-AQ 2.1  $\pm$  6.3 g kg<sup>-1</sup>, 456
- literature  $2.5 \pm 2.6 \text{ g kg}^{-1}$ ). 457
- 458
- 459



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Figure 2: Emission ratios of formic and acetic acid for literature values (box-and-whisker), WE-461 CAN PTR-ToF observations (green points), I<sup>-</sup> CIMS FA (green squares), and FIREX-AQ PTR-462 ToF (blue points). The box and whisker plots reported include literature ERs from all studies in

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Table S1, representing a variety of fuels (204 data points for formic acid and 196 for acetic acid). 464

Boxes represent the 25<sup>th</sup> and 75<sup>th</sup> percentiles, with vertical lines as median, whiskers as  $1.5 \times$  the interquartile range, and black points as >  $1.5 \times$  interquartile range of literature values.

468

To examine if the observed organic acid emission variability is related to burning condition, we compare the derived EFs from WE-CAN and literature coniferous forests to the modified combustion efficiency, which is a simple proxy used to describe the degree of flaming versus smoldering combustion in a fire. MCE is defined as,

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$$MCE = \frac{\Delta CO_2}{\Delta CO_2 + \Delta CO} \tag{1}$$

where  $\Delta CO_2$  and  $\Delta CO$  are the excess CO<sub>2</sub> and CO mixing ratios. An MCE near 1 corresponds to 474 pure flaming combustion, while MCEs of 0.65–0.85 represents pure smoldering.<sup>79</sup> During WE-475 CAN, MCEs ranged between 0.86–0.94, while those for mixed conifer forests in our literature 476 review have a larger range of 0.76–0.98. We note that MCEs span 0.68–0.99 when including all 477 fuel types in our literature review, with most MCEs < 0.84 corresponding to combustion of peat 478 and organic soils. Figure 3 shows FA and AA EFs vs MCE for both WE-CAN sampled fires and 479 coniferous forest literature values. The WE-CAN and literature EFs for FA have only a weak 480 negative dependence on MCE, with slopes of -4.8 ( $r^2 = 0.03$ ) and -6.4 ( $r^2 = 0.35$ ) respectively. 481 Using PTR-ToF FA data makes little difference, with the WE-CAN  $r^2 = 0.05$ . AA EFs have a 482 stronger negative correlation with MCE during WE-CAN (slope -20.7,  $r^2 = 0.52$ ) and for 483 literature values (slope = -20.2,  $r^2 = 0.14$ ). Expanding this analysis to include all fuel types in our 484 literature review results in a lower slope and  $r^2$  for the literature FA (-3.0, 0.071) and a larger 485 slope and  $r^2$  for AA (-27.3, 0.26). The poor correlation of FA EFs with MCE suggests that its 486 emissions variability is driven by factors other than combustion efficiency. Conversely, AA 487 488 emissions likely have some MCE dependence that should be accounted for when reporting and using EFs. 489

490



491

**Figure 3**: Correlations of FA (left) and AA (right) EFs versus MCE for both WE-CAN (red points) and literature reported coniferous forest values (blue points). The least squares regression for each group is shown in corresponding colors. For FA, the line of best fit for WE-CAN data is y = -4.8x + 5.8 ( $r^2 = 0.03$ ) and y = -6.4x + 6.3 ( $r^2 = 0.35$ ) for literature values. For AA, the line of best fit for WE-CAN data is y = -20.7x + 21.1 ( $r^2 = 0.52$ ) and y = -20.2x + 20.8 ( $r^2 = 0.14$ ) for literature values. Detailed statistics including PTR-ToF FA are shown in Table 1.

499

To determine if the type of fuel burned influenced FA or AA emissions, we compare WE-CAN 500 and literature EFs between the six fuel categories described above. For each organic acid we use 501 a Tukey's range test to evaluate if the 95 % confidence interval (CI) of emission factors for each 502 503 fuel type overlap. For FA, the Tukey range test p-values are > 0.05 for comparisons between all fuel types except with shrubland, indicating an overlap in the 95 % CI for most fuels. This 504 suggests that FA EFs for shrubland, mainly consisting of chaparral vegetation types in our 505 506 literature review, have statistically significant differences from the other 5 fuel categories. 507 Alternatively, no statistical difference was found between any of the other categories. Coupled with the lack of correlation with MCE, this suggests that a single FA ER of  $3.5 \pm 3.4$  ppb ppm<sub>CO<sup>-</sup></sub> 508 <sup>1</sup> and EF of 0.42  $\pm$  0.56 g kg<sup>-1</sup> (average of WE-CAN, FIREX-AQ, and literature values  $\pm$  1 $\sigma$ , 509 Table 1) best describe most BB emissions, though a fuel-specific EF for shrubland fuels (0.11  $\pm$ 510  $0.09 \text{ g kg}^{-1}$ ) may be more accurate. 511

512

513 AA EFs between coniferous forests, mixed hardwood forests, shrubland, and grassland similarly

- show no statistically significant fuel related difference, and an average ER of  $14.5 \pm 12.8$  ppb
- 515  $ppm_{CO}^{-1}$  and EF of 2.0 ± 1.9 g kg<sup>-1</sup> may best describe most fuel types. However, organic soil/peat

- and crop residue both have p-values < 0.05 when compared to the other four fuels, suggesting
- that MCE and fuel dependent EFs may be needed to best describe AA EFs. Given that AA shows
- some dependence on MCE, it is possible that the differences between crop residue and organic
- soil/peat EFs compared to EFs for the other fuel categories is in part due to combustion
- 520 efficiency. For example, organic soil/peat combustion is generally dominated by smoldering
- 521 (MCE = 0.68-0.92 in this work), which would result in higher EFs (Figure 3). Box plots of FA
- and AA EFs for each fuel category are shown in Figure S6.
- 523

**Table 1:** Emission factors (g kg<sup>-1</sup>) and emission ratios (ppb ppm<sub>CO</sub><sup>-1</sup>) for formic and acetic acid

reported in this work and in the literature. Note that the recommended average for FA is

526 calculated from WE-CAN I<sup>-</sup> CIMS, FIREX-AQ PTR-ToF, and all literature values. The

527 recommended average for AA are calculated from WE-CAN PTR-ToF, FIREX-AQ PTR-ToF,

528 and all literature values excluding crop residue and organic soil/peat.

529

		WE-CAN	FIREX-AQ	Literature average	Recommended Average
<u>Formic acid</u>	ER±1σ (range)	9.5 ± 4.2 (3.4–18.8)	3.3 ± 2.0 (0–6.5)	2.7 ± 2.6 (0.17–13.4)	3.5 ± 3.4
	EF±1σ (range)	1.5 ± 0.60 (0.55–2.5)	0.60 ± 0.42 (0–1.6)	0.35 ±0.48 (0.002–4.2)	0.42 ± 0.56
	n. obs.	20	16	168 ERs, 330 EFs	204, 366
	Eq. with MCE	y = -4.8x + 5.8 $r^2 = 0.03$		y = -6.4x + 6.3 $r^2 = 0.35$	y = -9.7x + 9.4 r <sup>2</sup> = 0.31
<u>Acetic acid</u>	ER ± 1σ (range)	11.5 ± 2.1 (6.4–16.7)	8.9 ± 1.5 (6.0–11.7)	15.5 ± 14.2 (0.9–85.6)	14.5 ± 12.8
	EF±1σ (range)	2.4 ± 6.1 (1.2–3.3)	2.1 ± 0.63 (1.1–3.4)	2.5 ±2.6 (0.14–14.0)	2.0 ± 1.9
	n. obs.	24	16	156 ERs, 289 EFs	196, 254
	Eq. with MCE	<i>y</i> = -20.7 <i>x</i> + 21 r <sup>2</sup> = 0.52		y = -20.2x + 20.8 r <sup>2</sup> = 0.14	y = -19.9x + 20.5 r <sup>2</sup> = 0.15

530

# 531 5 Near-field acid production during WE-CAN

532 FA and AA concentrations varied widely during WE-CAN with maximum mixing ratios of 98

ppb and 89 ppb, respectively. The highest FA NEMR of 71 ppb  $ppm_{CO}^{-1}$  was observed in smoke

aged ~13 hours. ERs were not measured for this fire; however, this is  $\sim 7 \times$  higher than campaign

average ER (9.5 ± 4.2 ppb ppm<sub>CO</sub><sup>-1</sup>), and 4 × higher than the maximum ER (18.8 ppb ppm<sub>CO</sub><sup>-1</sup>).

- 536 This suggests a maximum FA production rate of 4.0–4.7 ppb  $ppm_{CO}^{-1}h^{-1}$  in aged smoke sampled
- 537 during WE-CAN. This NEMR is approximately half of the maximum observed during ATom in
- smoke sampled off the African coast estimated to have been aged 1-10 days (140 ppb  $ppm_{CO}^{-1}$ ),
- though this latter value is similar to many other plumes intercepted during that campaign.<sup>17</sup>
- 540
- Figure 4 shows FA and AA NEMRs as a function of smoke plume age for 5 fires with more than 541 10 plume transects sampled in a pseudo-Lagrangian fashion during WE-CAN, while NEMRs for 542 all sampled plumes are shown in Figure S7 for reference. In the first 8 hours of plume aging FA 543 is rapidly produced at an average rate of 2.7 ppb  $ppm_{CO}^{-1}$  h<sup>-1</sup>. This is in good agreement with FA 544 production seen in other studies including 2.6–3.3 ppb  $ppm_{CO}^{-1} h^{-1}$  in smoke from Alaskan boreal 545 forest fires,<sup>20</sup> 1.6 ppb ppm<sub>C0</sub><sup>-1</sup> h<sup>-1</sup> from BB in the Yucatan, Mexico,<sup>22</sup> and 0.9 ppb ppm<sub>C0</sub><sup>-1</sup> h<sup>-1</sup> in 546 chaparral fires in California.<sup>19</sup> We hypothesize that the rapid FA production observed during 547 548 WE-CAN is at least in part responsible for the higher FA EFs and ERs discussed in Section 3.
- 549

550 During WE-CAN, AA NEMRs remain relatively constant in the first 8 hours of plume aging, increasing by a statistically insignificant 0.3 ppb  $ppm_{CO}^{-1}$  h<sup>-1</sup> (p = 0.13, r<sup>2</sup> = 0.03, Figure 4). 551 Additionally, the maximum AA NEMR was observed to be 17 ppb  $ppm_{CO}^{-1}$  in the same ~13 hour 552 aged plume discussed above, which is within  $3\sigma$  of the campaign average ER. The extent that 553 554 AA is produced in BB plumes is not well understood. For example, no net AA production has similarly been observed in smoke aged 1.4 hour over the Mexican Yucatán Peninsula<sup>22</sup> nor 555 inferred in BB plumes measured across Alaska and western Canada.<sup>80</sup> However, multiple other 556 studies have observed rapid AA production in the first few hours of plume aging including: 2.3 557 ppb  $ppm_{CO}^{-1}h^{-1}$  in smoke from a Californian chaparral fire aged 4.5 hours, <sup>19</sup> 1.5 ppb  $ppm_{CO}^{-1}h^{-1}$ 558 in 1 hour aged smoke from southeast U.S. prescribed agricultural burning,<sup>34</sup> 1.5–2.0 ppb ppm<sub>CO</sub><sup>-1</sup> 559  $h^{-1}$  in smoke aged < 1 hour from African Savanah fires, <sup>26</sup> and 7.2 ppb ppm<sub>CO</sub><sup>-1</sup>  $h^{-1}$  in smoke aged 560 1 hour from Alaskan boreal forest fires.<sup>20</sup> Future work is needed to better characterize AA 561 production, especially in smoke that has aged more than half a day. 562 563

564 One potential explanation for the lack of observed AA production during WE-CAN is that the 565 removal of glycolaldehyde offsets the formation of AA. This is possible because AA measured

by PTR-ToF may be ~30 % glycolaldehyde in fresh emissions,<sup>25,63</sup> both species have similar 566 sensitivities in PTR-MS,<sup>53</sup> and glycolaldehyde is  $\sim 20 \times$  more reactive than AA with OH ( $k_{OH}$ 567  $1.1 \times 10^{-11}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup> vs.  $7.4 \times 10^{-13}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup>; NIST chemical kinetics database 568 average). Additionally, of the studies that observed AA production listed above, only Muller et 569 al.<sup>34</sup> used a PTR-ToF while the others used FTIR, which does not have isomeric interferences for 570 AA. To test this hypothesis, we attribute only 66 % of the PTR-ToF m/z 61 to AA to calculate a 571 corrected campaign average ER of 7.6 ppb  $ppm_{CO}^{-1}$ . Assuming there is negligible glycolaldehyde 572 formation downwind, such that the PTR-ToF is only measuring AA in aged smoke, the 573 maximum AA production rate over ~13 hours of plume aging would be 0.7 ppb  $ppm_{CO}^{-1} h^{-1}$ . 574 Although this production rate is still lower than in the literature, the 30 % reduction in the 575 campaign average AA ER brings it outside of the observed NEMR variance for most aged smoke 576 577 samples. This suggests that AA production could be statistically significant if these assumptions are true, and more detailed characterization of the glycolaldehyde interference in aged smoke is 578 579 needed.

580

581 It is also possible that most of the AA formation happens in the first hours of plume aging, which is then averaged out in our analysis of plumes that are aged 8 hours during WE-CAN. For 582 example, in one plume, Goode et al.<sup>20</sup> observed an AA increase of 11 ppb  $ppm_{CO}^{-1}$  in the first 1.5 583 hours of plume aging followed by no net production relative to emissions in 2.8 hour old smoke. 584 585 However, limiting WE-CAN observations to those with <2 hours aging in Figure 4 still results in a non-statistically significant NEMR-vs-age slope of 0.23 ppb  $ppm_{CO}^{-1}$  h<sup>-1</sup> (p = 0.80, r<sup>2</sup> = 0.002). 586 Similarly, if each of the 5 fires in Figure 4 are treated individually, only two have statistically 587 significant correlations (p = 0.02) with plume age, both with negative slopes (-0.2 and -0.3 ppb 588  $ppm_{CO}^{-1} h^{-1}$ ). Given the relatively long atmospheric lifetime of AA (~2-3 days),<sup>8,38</sup> it is unlikely 589 590 that a significant amount was removed from the plume in the 8 hours of aging shown here. Consequently, these results suggest that most of the observed AA in the near-field during WE-591 CAN is from primary emissions, though photochemical production may still be an important 592 source in some fires and over longer plume aging times which should be investigated further. 593 594 Previous work has used the Framework for 0-D modeling (F0AM)<sup>81</sup> to simulate the Taylor Creek

595 Previous work has used the Framework for 0-D modeling (F0AM)<sup>81</sup> to simulate the Taylor Creek
596 (TC) fire sampled during WE-CAN due to it being a well isolated plume with pseudo-

Lagrangian samples performed just after injection into the free troposphere.<sup>24,46,49,82–85</sup> Here, we 597 use the same F0AM model run as originally described in Peng et al.<sup>85</sup> with updated VOC 598 emissions per Permar et al.<sup>24</sup> Briefly, F0AM was initialized using 49 VOCs, plus NO, NO<sub>2</sub>, 599 HONO, O<sub>3</sub>, and CO (Table S2). For VOCs measured by PTR-ToF, potential interfering isomers, 600 including glycolaldehyde, were removed so that the model was initialized based on the 601 proportion of the mass attributed only to the given species following Koss et al.<sup>25</sup> Physical 602 parameters such as photolysis frequencies, temperature, and pressure were constrained to 603 604 measured values at each model step with a dilution correction factor based on CO observations. Model chemistry was simulated using the explicit Master Chemical Mechanism (F0AM+MCM) 605 including recently developed furans and phenolic chemistry,<sup>86,87</sup> with an additional sensitivity 606 test run using  $3 \times \text{VOC}$  initial values for all gases except FA and AA (F0AM+MCM×3). We also 607 608 run the same box model driven with the GEOS-Chem chemical mechanisms (F0AM+GC) to test if recent updates by Chen et al.<sup>17</sup> significantly impact the modeled FA production in fresh smoke. 609 610

Figure 4 shows that neither F0AM+MCM, F0AM+MCM×3, nor F0AM+GC can reproduce the rapid FA formation observed in the TC plume, with modeled FA instead decreasing slightly with plume age. The MCM predicted loss rate for FA in the base run to be ~0.3 ppb h<sup>-1</sup> by reaction with OH, while there is minimal production (~0.01 ppb h<sup>-1</sup>) from C<sub>4</sub>H<sub>6</sub>O<sub>3</sub> and CH<sub>2</sub>OO Criegee intermediates. Coupled with the MCM being insensitive to increased initial values, this represents more FA being removed in the model than is being produced and indicates that both the MCM and GC are missing a substantial amount of secondary FA production in BB smoke.

The total emitted VOC carbon during WE-CAN averaged  $367.3 \pm 29.6 \text{ ppb}_{\text{C}} \text{ ppm}_{\text{CO}^{-1}}$ .<sup>18</sup> As the average FA enhancement shown in Figure 4 is 21 ppb ppm<sub>CO</sub><sup>-1</sup>, approximately 5.7 % of the VOC carbon oxidation would need to go to FA to explain the observed production. Additionally, the maximum observed FA NEMR of 71 ppb ppm<sub>CO</sub><sup>-1</sup> after ~13 hours of plume aging represents a 61 ppb ppm<sub>CO</sub><sup>-1</sup> enhancement relative to the campaign average ER, indicating that up to 17 % of the total emitted carbon could be converted to FA within half a day.

AA NEMRs in the TC plume are highly variable, likely representing changes in fire emissions or
 sampling different parts of the plume. Figure 4 shows that F0AM+MCM and F0AM+GC

generally have good agreement with the observed AA NEMRs. Similar to FA, AA is also mainly lost in the MCM through reaction with OH at ~0.4 ppb h<sup>-1</sup>, with negligible production (<0.01

 $^{630}$  ppb h<sup>-1</sup>) from CH<sub>3</sub>CHOO Criegee intermediates and CH<sub>3</sub>C(O)O<sub>2</sub> peroxy acetyl radicals.

631



Figure 4: NEMRs of FA and AA for 5 research flights with more than 10 pseudo-Lagrangian 633 transects. Blue triangles highlight plume transects of the Taylor Creek (TC) fire and correspond 634 to the red dashed F0AM+MCM, orange F0AM+MCM×3, and green F0AM-GC predicted 635 NEMRs for the same fire. Black points correspond to the other 4 fires. Least squares regression 636 lines for the aggregated data are shown in gray. During the first 8 hours of plume aging FA 637 NEMR increased on average 2.7 ppb  $ppm_{CO}^{-1}$  per hour (r<sup>2</sup> = 0.58, intercept = 9.3 ppb  $ppm_{CO}^{-1}$ ), 638 while AA has a statistically insignificant increase of 0.3 ppb  $ppm_{CO}^{-1}$  per hour ( $r^2 = 0.03$ , 639 intercept = 8.4 ppb  $ppm_{CO}^{-1}$ ). 640

641

A current lack of understanding of the major FA and AA precursors is one of the largest hurdles

to accurately modeling their evolution in smoke. To evaluate potential VOC precursors, NEMRs

for both acids measured in the same 5 wildfires as described above were compared to NEMRs of

- 645 152 VOCs measured during WE-CAN using least squares regression. FA was found to have
- statistically significant negative correlations (p-value < 0.05,  $r^2 > 0.10$ ) with 94 VOCs. Over the
- 647 8 hours of plume aging the oxidation of these 94 species collectively accounts for 127 ppb<sub>C</sub>
- 648  $ppm_{CO}^{-1}$  that is reacted away. This indicates that those species lose 6 × more carbon than is
- needed to account for the observed FA production, though the exact chemical pathways are often
- unknown. For example, C<sub>3</sub>H<sub>4</sub>O<sub>2</sub> (methyl glyoxal + acrylic acid), styrene, and furanoid
- compounds such as 3-methylfuran and furfural are among the species with strongest correlations

to FA in fresh smoke during WE-CAN (Figure 5;  $r^2 > 0.40$ ). However, the MCM and recent chemical mechanism developments do not show any chemical production of FA from these compounds.<sup>86,88,89</sup>

655

Figure 5 similarly shows that FA is well correlated  $(r^2 > 0.4)$  with isoprene, ethene, and 656 acetaldehyde, consistent with the current understanding of these species being known FA 657 precursors. Additionally, FA was found to have a strong correlation with peroxyacetyl nitrate 658 (PAN;  $r^2 = 0.46$ ) and a modest correlation with ozone ( $r^2 = 0.23$ ), further indicating that FA 659 production follows the overall plume gas phase oxidization. While the correlations of FA 660 NEMRs with the VOCs in Figure 5 do not directly indicate that they are FA precursors in smoke 661 plumes, when coupled with FA being well corelated to PAN, ozone, and 94 different VOCs, they 662 do demonstrate that FA is likely being produced through the oxidation of many different species, 663 most of which are currently not well studied in the literature. 664

665

Heterogenous formation is also likely to be an important FA source in smoke via a multiphase 666 pathway where methanediol (HOCH<sub>2</sub>OH) is off gassed from aerosols and is rapidly oxidized by 667 OH to form HCOOH.<sup>32,33</sup> The WE-CAN dataset does not have sufficient data to fully examine 668 how this pathway may contribute to the FA production observed during the campaign. Instead, 669 we explore whether the FA NEMRs show dependence on OA aging during WE-CAN in Figure 670 5j by comparing FA NEMRs with the OA oxidation marker  $f_{44}$ .<sup>50,51</sup>  $f_{44}$  is the fractional 671 component of OA attributed to the  $CO_2^+$  ion which is ascribed to fragments of acids or acid-672 derived species.<sup>90</sup> Consequently, f<sub>44</sub> is generally well correlated with the OA elemental O:C 673 ratio,<sup>91</sup> where both increase as the bulk aerosol becomes more oxidized. During WE-CAN f<sub>44</sub> 674 was found to increase with smoke plume age, while the dilution-adjusted OA mass generally 675 remained unchanged over ~8 hours of plume aging.<sup>49</sup> Figure 5j shows that FA NEMRs are 676 positively correlated with  $f_{44}$  ( $r^2 = 0.42$ ) as well as with the OA O:C ratio ( $r^2 = 0.32$ , not shown). 677 This suggests that FA production follows the bulk aerosol oxidation during WE-CAN. 678 Additionally, the increasing OA oxidation with the constant downwind dilution-adjusted OA 679 mass reported by Garofalo et al.<sup>49</sup> requires a balance between evaporation and condensation of 680 semivolatile species. This indicates that FA could be formed as part of this OA mass balance and 681

more detailed laboratory and field studies are needed to better understand this potentially

683 significant FA formation pathway in BB smoke.





685

Figure 5: FA NEMRs compared to various gas phase species NEMRs and aerosol f<sub>44</sub> ratios 686 measured in 5 smoke plumes with more than 10 pseudo-Lagrangian plume transects. Slope and 687  $r^2$  for the least squares regression of each species are shown at the bottom of each panel, while 688 the gray lines represent the best fit. Panels a, b, and c show the three VOCs with the strongest 689 correlation to FA. Panels d and e show two of the largest OH radical sinks (ranked by OH 690 reactivity from individual VOC)<sup>24</sup> that are highly correlated with FA in wildfire emissions. 691 Panels f, and g show known FA precursors, while quantities plotted in h, i and j are 692 representative of the overall plume oxidation. Note, methyl glyoxal is measured with acrylic acid 693  $(C_{3}H_{4}O_{2})$ . PAN = peroxyacetyl nitrate.  $f_{44}$  = ratio of m/z 44 to the total signal in the aerosol 694 component spectrum with higher ratios indicating more aged organic aerosol and higher O:C. 695 696

- 697 Similar analysis with AA is shown in Figure S8, with AA NEMRs plotted against a similar
- grouping of gases as in Figure 5. The three species with the strongest correlation against AA are
- shown in Figure S8a, b and c:  $C_3H_6O_2$  (hydroxyacetone + methyl acetate + ethyl formate;  $r^2 =$
- 0.62), C<sub>5</sub>H<sub>8</sub>O<sub>3</sub> (5-hydroxymethyl tetrahydro 2-furanone;  $r^2 = 0.48$ ), and methyl propionate ( $r^2 = 0.48$ )
- 0.45). Like AA, NEMRs for these three species are not well correlated with the physical plume
- age. Figure S8 also shows that AA has only modest correlation with the reactive VOCs furfural

and isoprene as well as with acetaldehyde and ozone ( $r^2 = 0.14-0.32$ ) but is poorly correlated

with ethene and PAN ( $r^2 < 0.1$ ). The fact that a) AA is most strongly correlated with other VOCs

whose NEMRs remain mostly unchanged with plume age and b) has poor negative correlations

with the plume oxidation indicators such as PAN, ozone, and  $f_{44}$ , further supports the observation

that little AA is produced and instead is mainly from primary emissions in the WE-CAN sample

708 wildfire plumes.

#### 709 6 GEOS-Chem representation of FA and AA during WE-CAN and FIREX-AQ

Global chemical transport models typically have difficulty simulating formic and acetic acid

mixing ratios, particularly in the presence of BB smoke. Section 5 suggests that the GEOS-Chem

chemistry underestimates a significant amount of secondary production of FA in fresh smoke.

Here we investigate how the GEOS-Chem CTM, with the most recent updates for FA

<sup>714</sup> implemented by Chen et al.<sup>17</sup> (Section 2.3), represents FA and AA in different environments

sampled during the WE-CAN, FIREX-AQ-W, and FIREX-AQ-SE. All WE-CAN and FIREX-

AQ measurements have also been averaged to 5 minutes to match the model resolution. GEOS-

717 Chem was sampled along the plane flight tracks at the time of each corresponding research

718 flight.

719

Figure 6 shows that GEOS-Chem generally underestimates the vertical distribution of FA 720 observed during WE-CAN (-92 %; normalized mean bias to I<sup>-</sup> CIMS, NMB) and in the middle to 721 lower troposphere (>450 hPa or below ~7.2 km above sea level) during FIREX-AQ-W (-76 % 722 NMB) and FIREX-AQ-SE (-37 %). This corresponds to the model underestimating the average 723 724 measured FA by nearly a factor of 13 during the WE-CAN deployment, while also underestimating FA by a factor of 4 and 2 in the lower altitude FIREX-AQ-W and FIREX-AQ-725 SE samples. However, GEOS-Chem does significantly better simulating FA in the middle to 726 upper troposphere (<450 hPa; -27 % NMB in FIREX-AQ-W), consistent with findings by Chen 727 728 et al.<sup>17</sup> Interestingly, GEOS-Chem overestimates FA mixing ratios compared to I<sup>-</sup> CIMS measurements at higher altitudes (< 450 hPa) in the southeastern U.S. (213 %), though the 729 measured FA is reaching the stated I<sup>-</sup> CIMS detection limit (~30 ppt). Figure S9 shows a similar 730 underestimation for acetic acid mixing ratios with NMB ranging -92 % to -99 % in both high and 731 732 low altitude WE-CAN and FIREX-AQ-W samples, and slightly better agreement (-80 %) with

lower altitude FIREX-AQ-SE periods. We note that this significant underestimate of FA by
GEOS-Chem holds true regardless of the high uncertainty in FA measured during the WE-CAN
deployment as the difference between the modeled and measured values is much greater than the
instrument uncertainty.

737



Figure 6. Vertical profiles of the median formic acid mixing ratios measured during the full WE-739 CAN and FIREX-AQ field campaigns, binned at every 33 hPa. Black and gray lines correspond 740 to the measurements made by I<sup>-</sup>CIMS and PTR-ToF, with error bars representing the 25<sup>th</sup> and 741 75<sup>th</sup> percentile of I<sup>-</sup> CIMS measurements at each pressure bin. Red dashed lines correspond to 742 GEOS-Chem with GFAS BB emissions (GC), orange dashed lines represent GEOS-Chem with 3 743  $\times$  GFAS BB emissions (GC $\times$ 3), and the pink dotted lines show GEOS-Chem with BB emissions 744 turned off (GC NoBB). The number of samples in each pressure bin are shown on the right of the 745 plots, while the normalized mean bias (NMB) to the I<sup>-</sup> CIMS measurement for lower altitude 746 observations (> 450 hPa) are shown at the top. 747

748

749 There are a few possible explanations for why GEOS-Chem underestimates FA and AA during

the two campaigns, including: incorrect or missing emissions, sampling bias, and/or missing

secondary chemistry from BB (Section 5) and biogenic precursors. Recent model developments

- <sup>752</sup> have improved the GEOS-Chem representation of the free troposphere,<sup>17</sup> chemistry, and
- biogenic sources.<sup>7</sup> Subsequently, we hypothesize that missing secondary production from BB
- and western U.S. specific biogenic precursors are likely key reasons for GEOS-Chem
- underestimating FA mixing ratios during WE-CAN and FIREX-AQ-W. The exact reason for the
- underestimation of AA mixing ratios is uncertain, though we speculate it may in part be due to

the model sink being too large<sup>38</sup> and/or secondary production in BB smoke aged over greater processing times than discussed in Section 5 (i.e., > 8 hours).

759

#### 760 Model BB emissions and sampling bias

Recent work has shown that commonly used global emission inventories, including GFAS, 761 GFED4, QFED, and FINNv1.5, underestimate BB emissions by a factor of three or more in the 762 western U.S. when compared to aircraft and ground-based measurements.<sup>41,92</sup> Jin et al.<sup>41</sup> 763 attributes this mostly to the significant underestimation of the dry biomass burned in the BB 764 765 emission inventories. To explore if underestimated BB emissions can explain the low FA and AA model bias, GEOS-Chem was also initiated with 3 × GFAS BB emissions as a sensitivity 766 767 test, in which the BB VOC and CO emissions are tripled from the base run. Figure S10 shows that GEOS-Chem with base GFAS emissions underestimate CO during WE-CAN, FIREX-AQ-768 W, and FIREX-AQ-SE. Model representation in the western U.S. is improved by the  $3 \times GFAS$ 769 model run, in good agreement with Jin et al.<sup>41</sup> Similarly, though benzene and acetone are better 770 771 represented by the base model in this work, the  $3 \times GFAS$  emission simulation further improves their model agreement. Despite this, Figures 6 and S9 show that increasing BB emissions by a 772 773 factor of 3 only slightly increases the model FA and AA mixing ratios, decreasing the NMB by  $\sim$ 5 % in all cases. Given 1) that tripling BB emissions has minimal impact on the modeled FA or 774 AA. 2) that the GFAS inventory is not missing the location/timing of the fires sampled during 775 both campaigns,<sup>41</sup> and 3) that FA and AA BB emissions were implemented per observed ERs,<sup>41</sup> 776 777 underestimation of primary BB emissions of either acid or their known precursors in GEOS-Chem alone cannot account for the low model bias. This reflects that the contribution of primary 778 BB emissions to ambient FA during WE-CAN and FIREX-AQ in the western US is small. 779 Additionally, the  $3 \times \text{GFAS}$  run also increases emissions for all BB implemented species.<sup>24</sup> thus 780 pointing to missing secondary formation pathways from either implemented and/or unknown 781 precursors in the model (see Section 5). Given the lack of evidence for near-field AA production 782 during WE-CAN, the model being largely insensitive to a 3-fold increase in AA emissions 783 suggests that AA production in BB plumes aged greater than the 8 hours observed during WE-784 CAN may still be significant and/or the overall model sink is too large. 785

786

The WE-CAN and FIREX-AQ aircraft campaigns were focused on sampling and tracking BB 787 smoke whenever possible. As GEOS-Chem was run at  $0.25^{\circ} \times 0.3125^{\circ}$  (~25 km) resolution, the 788 low model bias may in part also reflect the dilution of narrow smoke plumes over the model grid. 789 Though some error is inherent in the model comparisons due to this sampling bias, using GEOS-790 Chem run with the same WE-CAN and FIREX-AO datasets. Jin et al.<sup>41</sup> demonstrated that the 791 model also had difficulty simulating smoke impacts at longer term ground measurement sites 792 across the western U.S. This is indicative that the low model biases cannot be explained by the 793 model resolution alone. Similarly, Jin et al.<sup>41</sup> showed that fire detection products across emission 794 inventories did well capturing the large fires sampled during WE-CAN and that GEOS-Chem is 795 fairly insensitive to plume injection heights for the averaged WE-CAN campaign, likely due to 796 efficient vertical mixing during the summer months.<sup>93,94</sup> However, because of these issues when 797 798 comparing fire plumes sampled by aircraft to global CTMs, the GEOS-Chem evaluation here further focuses on the campaign averages across two different years, in smoke impacted, no/low 799 800 smoke, and clean free troposphere environments.

801

#### 802 **Representation in different environments**

803 To investigate potential model deficiencies over broad regions, we further examine the model performance in different environments sampled during the campaigns as described in our 804 previous work.<sup>24</sup> Here, smoke-impacted sampling periods for both campaigns are defined as 805 those with hydrogen cyanide (HCN) > 250 ppt and acetonitrile (CH<sub>3</sub>CN) > 200 ppt, while 806 periods below this threshold are discussed as low/no-smoke. However, due to widespread 807 regional smoke during the fire season, the low/no-smoke samples likely still represent some BB 808 influence. In addition to this coarse filter, clean free troposphere samples were also defined for 809 both campaigns based on HCN < 250 ppt, CH<sub>3</sub>CN < 150 ppt, and pressure < 624 hPa (~4 km 810 above sea level, representing the maximum boundary layer height as determined from vertical 811 temperature profiles). 812

813

Figure 7 shows the vertical profiles for the median observed and modeled FA in the three

815 different environments. We find GEOS-Chem underestimates the median FA mixing ratio most

significantly in smoke impacted samples, doing slightly better during low/no smoke periods in

the western U.S. Alternatively, GEOS-Chem does well simulating FA mixing ratios in the free 817 troposphere during all three periods, in good agreement with Chen et al.<sup>17</sup> This is particularly 818 evident in FIREX-AQ-W free troposphere samples, which agree nearly 1:1 with the model. 819 Similarly, the model also does very well simulating median FA mixing ratios in the low/no 820 smoke southeast U.S. samples (NMB -36 %). As this profile reflects minimal smoke impact 821 during the period, it suggests that the model is accurately simulating FA from biogenic sources in 822 the southeast U.S., as reflected in recent model developments including production from 823 stabilized Criegee intermediates and acetaldehyde tautomerization as implemented by Millet et 824 al.<sup>7</sup> and Chen et al.<sup>17</sup> 825 826

Although GEOS-Chem does better simulating FA during low/no smoke samples than in smoke 827 in the western U.S., the improvement is only modest with NMB decreasing by < 10 %. This may 828 in part reflect the widespread smoke impacts in the western U.S. during fire season, where a pool 829 of longer-lived oxygenated species could persist in the region.<sup>24</sup> However, it also suggests that 830 the model may be missing a FA source or secondary chemistry from biogenic precursors unique 831 to coniferous forests,<sup>11</sup> which are likely different than those most responsible for FA in the 832 southeastern U.S. For example, isoprene oxidation is thought to be one of the main contributors 833 to FA formation above deciduous forests,<sup>7</sup> while in coniferous forests emissions are typically 834 dominated by monoterpenes and 2-methyl-3-buten-2-ol (MBO),<sup>95</sup> whose potential contribution 835 836 to FA formation is unclear.

837



**Figure 7.** Vertical profiles of the median formic acid mixing ratios measured during the WE-CAN field campaign for smoke impacted, low/no smoke, and free troposphere sampling periods. Pressures are binned at every 33 hPa. Black and gray lines correspond to the measurements made by I<sup>-</sup>CIMS and PTR-ToF. Red dashed lines correspond to GEOS-Chem with GFAS BB emissions (GC), orange dashed lines are GEOS-Chem with  $3 \times$  GFAS BB emissions (GC×3), and the pink dotted lines are GEOS-Chem with BB emissions turned off (GC NoBB). Error bars are the 25<sup>th</sup> and 75<sup>th</sup> percentile of the I<sup>-</sup>CIMS measurement at each pressure bin.

846

To explore the regional sources of FA using the two campaign datasets, Figure 8 shows how FA

correlates with CO, methanol, acetone, and MVC+MACR (methyl vinyl ketone and

849 methacrolein) in the three regions and environments shown in Figure 7. The plot of FA vs CO

shows two distinct populations between the smoke and low/no smoke environments. As CO is

mainly from BB in the WE-CAN and FIREX-AQ datasets, the correlation of FA with CO in

- smoke samples indicates FA coming from BB sources, while the spread likely represents FA
- enhancement relative to primary emissions. In low/no smoke samples, the FA:CO slope is
- steeper than in the smoke samples, suggesting a FA source that is independent of the combustion
- process thus pointing to photochemical origin.
- 856

Interestingly, Figure 8 also shows that FA is well correlated with both methanol ( $r^2 = 0.55-0.75$ ) 857 and acetone (0.42-0.72), with generally similar slopes in both smoke and low/no smoke samples 858 (methanol = 0.5-0.9, acetone = 0.2-0.4) during WE-CAN and FIREX-AQ-W periods. In 859 contrast to FA, neither methanol nor acetone measured during WE-CAN see net production in 860 the 5 smoke plumes and ~8 hours of aging discussed in Section 5 (Figure S11). This suggests 861 that their correlations in Figure 8 are not due to near-field production in BB, though 862 enhancement in much more aged plumes (> 2 days) has been observed.<sup>96,97</sup> As methanol and 863 acetone are known to be major primary emissions and secondary products from biogenic 864 sources,<sup>12,96,98–101</sup> we hypothesize that their strong correlation with FA during WE-CAN and 865 FIREX-AO indicates that a portion of the observed FA may be of biogenic origin, though the 866 long atmospheric lifetimes of all three species (> 2 days) likely also play a role in why they are 867 868 well correlated with each other.

869

870 As isoprene is known to be the major FA precursor in deciduous forests, Figure 8 also shows FA vs. MVK+MACR, an important isoprene oxidation product. During WE-CAN and FIREX-AO, 871 FA has a weak positive correlation with MVK+MACR in most environments ( $r^2 = 0.11-0.24$ ), 872 further indicating that some of the observed FA is indeed related to biogenic species. Some of 873 the agreement between the two is also likely due to both being primary BB emissions,<sup>18</sup> while 874 the large spread in the correlations also points to MVK+MACR being lost as the plumes age 875 876 (Figure S11). Consequently, the model underestimate of FA in the western U.S. is likely due to both missing secondary chemistry from BB and biogenic sources, pointing to a need for more 877 detailed studies of FA production from both BB and coniferous forest emissions. 878

879



880

Figure 8. Correlations of FA with CO, methanol, acetone, and MVK+MACR (methyl vinyl
ketone and methacrolein) in WE-CAN and FIREX-AQ observations. Orange points represent
smoke-impacted data, blue points indicate low/no smoke impact, and green points show clean
free troposphere measurements (see main texts for definitions). The data have been averaged to 5
minutes. Lines show the least squares regression corresponding to each set of colored points.
Note that acetone is also measured with its isomer propanal.

887

A similar trend can be seen for acetic acid in Figure S12, where GEOS-Chem underestimates AA in both smoke-impacted and low/no smoke environments during all three sampling periods, with

- NMB improving by < 10 % between smoke low/no smoke conditions. Additionally, AA is better
- captured by the model in the clean free troposphere during WE-CAN (NMB -44 %), though the
- disparity is larger for both portions of FIREX-AQ (NMB -92 %). Figure S13 shows that AA is
- well correlated with CO across all WE-CAN and FIREX-AQ-W samples, with a slope in the
- range of reported ERs (WE-CAN slope = 15.8 ppb  $ppm_{CO}^{-1}$ ,  $r^2 = 0.84$ ; FIREX-AQ-W slope =
- 11.0 ppb  $ppm_{CO}^{-1}$ ,  $r^2 = 0.92$ ). This further indicates that AA and CO in the western U.S. come

from the same source, likely BB. Given the lack of evidence for production of AA in the fresh 896 BB plumes sampled during WE-CAN (Section 5), near-field production is unlikely to explain the 897 low model bias, though production in plumes aged longer than those sampled during WE-CAN 898 is still possible. Additionally, the underrepresentation cannot be accounted for by BB emission 899 alone for two reasons: 1) the AA (and FA) emission ratio in the model was implemented using 900 the WE-CAN observations per Permar et al.<sup>18</sup> and 2) the GEOS-Chem +  $3 \times$  GFAS, which 901 should account for the underestimated BB emissions per Jin et al.<sup>41</sup> only slightly increases the 902 modeled AA. Consequently, the exact reason behind the low model bias for AA is unknown, 903 though it may be due to too large of a model sink and/or missing secondary production from long 904 lived biogenic and BB precursors. 905

#### 906 4 Conclusions

Using detailed formic acid and acetic acid measurements made during the WE-CAN and FIREX-907 908 AQ aircraft campaigns, we assess their emissions, chemistry, and model representation in the western and southeastern U.S. FA measured by two commonly used mass spectrometers, PTR-909 910 ToF and I<sup>-</sup>CIMS, was found to have high measurement uncertainty during the WE-CAN deployment (up to 100%) due to its humidity and temperature dependent sensitivities, inlet 911 artifacts, and instrument baseline issues. However, FA measured by two different PTR-ToF and 912 I<sup>-</sup> CIMS instruments during the FIREX-AQ campaign were found to agree within their 913 measurement uncertainty. Accuracy for the FA measurement could be greatly improved by 914 reducing inlet losses via shorter sampling lines, increased flow rate, and/or reduced sampling line 915 diameter, thus reducing sample residence time in the instrument. In addition, regulating reaction 916 917 chamber temperatures, and performing more frequent humidity dependent calibrations and instrument zeroing checks are key to improving the FA measurement quality in both instrument 918 types. Despite the high uncertainty in FA measured during WE-CAN, model underestimates of 919 FA mixing ratios were found to be much greater than the measurement uncertainty. 920

921

During WE-CAN FA ERs and EFs were found to be  $9.5 \pm 4.2 (1\sigma)$  ppb ppm<sub>CO<sup>-1</sup></sub> and  $1.5 \pm 0.60$  g kg<sup>-1</sup> respectively, which are 3.5 times higher than literature values. FIREX-AQ AA EFs and ERs agree better with the literature, though are still often higher than the 75<sup>th</sup> percentile of literature values. As FA was found to have little to no dependence on MCE or fuel type. The exact reason for this discrepancy is currently unknown, though may reflect differences in emissions between the larger wildfires sampled during WE-CAN and FIREX-AQ than from laboratory BB studies and the smaller fires typically reported in the literature. It may also reflect some extent of early plume production as the WE-CAN flights sampled 27–130 minutes downwind from the source. However, extrapolating FA NEMRs measured downwind from these fires to t<sub>0</sub> does little to close the gap between WE-CAN and literature ERs and EFs.

932

933 Analysis of 5 smoke plumes sampled in a pseudo-Lagrangian fashion finds that FA is rapidly produced at 2.7 ppb  $ppm_{CO}^{-1}$  h<sup>-1</sup> during the first 8 hours of plume aging, in good agreement with 934 previous studies. However, F0AM run with explicit MCM or simplified GEOS-Chem chemistry 935 was unable to capture the observed production due to missing secondary sources. Observed FA 936 937 production was found to have statistically significant correlations (p-value < 0.5) with 94 VOCs measured during WE-CAN. The oxidation of these 94 species collectively accounts for 127 ppb<sub>C</sub> 938  $ppm_{CO}^{-1}$  that is reacted away over the 8 hours of plume aging. This indicates that those species 939 could lose  $6 \times$  more carbon than is needed to account for the observed FA production, though the 940 941 exact chemical pathways are often unknown.

942

AA ERs and EFs were found to fall within the 25<sup>th</sup> and 75<sup>th</sup> percentiles of the literature-reported values, exhibiting a modest negative dependence on MCE and some fuel types. In contrast to some previous studies, AA was not found to have any statistically significant production during the first 8 hours of plume aging during WE-CAN, with downwind NEMRs generally in the range of observed ERs. Consequently, most of the observed AA in the nearfield is likely from primary emissions, though photochemical production may still be important for certain fires/fuels and in more aged smoke.

950

GEOS-Chem simulations with updated FA and AA chemistry and emissions were performed for
the WE-CAN and FIREX-AQ campaigns and compared to field observations. For both
campaigns, FA and AA were found to be biased low in the model by ~90 %. The model does
slightly better simulating FA mixing ratios in no/low smoke impacted western U.S. samples, and
significantly better in no/low smoke periods over southeast U.S. forests. It is likely that much of
the low model bias for FA is due to missing secondary production from both BB and coniferous

957 forest specific biogenic sources. The factors leading the underestimate of AA are unknown, but

may reflect too large of a model sink and secondary production in smoke aged longer than

observed during WE-CAN.

960

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