Chemical Tomography in a Fresh Wildland Fire Plume: a Large Eddy Simulation (LES) Study

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November 23, 2022

Abstract

Wildland fires involve complicated processes that are challenging to represent in chemical transport models. Recent airborne measurements reveal remarkable chemical tomography in fresh wildland fire plumes, which remain yet to be fully explored using models. Here we present a high-resolution large eddy simulation (LES) model coupled to chemistry to study the chemical evolution in fresh wildland fire plume. The model is configured for a large fire heavily sampled during the Fire Influence on Regional to Global Environments and Air Quality (FIREX-AQ) field campaign, and a variety of airborne measurements are used to evaluate the chemical heterogeneity revealed by the model. We show that the model captures the observed cross-transect variations of a number of compounds quite well, including ozone (O_3) , nitrous acid (HONO), and peroxyacetyl nitrate (PAN), etc. The combined observational and modeling results suggest that the top and edges of fresh plume drive the photochemistry, while dark chemistry is also present but in the lower part of the plume. The model spatial resolution is shown to be very important as it may shift the chemical regime, leading to biases in O_3 and NO_x chemistry. Based on findings in this work, we speculate that the impact of small fires on air quality may be largely underestimated in models with coarse spatial resolutions.

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2 Simulation (LES) Study

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31	KEY POINTS		
32	1.	Photochemistry is active at the edges of thick fire plumes, while dark chemistry is present	
33		in the lower part and below thick plumes	
34	2.	Hydroxyl radicals formed from nitrous acid drive the plume oxidation. Nitrous acid may	
35		be produced on aerosols	
36	3.	Model spatial resolution affects chemical regime; sufficiently high spatial resolution (e.g.,	
37		1 km) is needed to capture the impacts of wildfires on air quality	
38			

39 ABSTRACT

40 Wildland fires involve complicated processes that are challenging to represent in chemical transport models. Recent airborne measurements reveal remarkable chemical tomography in 41 fresh wildland fire plumes, which remain yet to be fully explored using models. Here we present 42 43 a high-resolution large eddy simulation (LES) model coupled to chemistry to study the chemical 44 evolution in fresh wildland fire plume. The model is configured for a large fire heavily sampled 45 during the Fire Influence on Regional to Global Environments and Air Quality (FIREX-AQ) field 46 campaign, and a variety of airborne measurements are used to evaluate the chemical 47 heterogeneity revealed by the model. We show that the model captures the observed cross-48 transect variations of a number of compounds quite well, including ozone (O_3), nitrous acid (HONO), and peroxyacetyl nitrate (PAN), etc. The combined observational and modeling results 49 50 suggest that the top and edges of fresh plume drive the photochemistry, while dark chemistry is 51 also present but in the lower part of the plume. The model spatial resolution is shown to be very important as it may shift the chemical regime, leading to biases in O₃ and NO_x chemistry. Based 52 53 on findings in this work, we speculate that the impact of small fires on air quality may be largely 54 underestimated in models with coarse spatial resolutions.

56 PLAIN LANGUAGE SUMMARY

57 Recent fire seasons in the United States have been record-setting for many states. Several 58 large wildfires raged across the entire west coast, and lofted smoke plumes spread to the 59 majority of the continental U.S. From a scientific perspective, wildland fires are fascinating due 60 to their complexity. Fires emit heat, creating a plume of hot and turbulent air. The fire plume also 61 contains many gases and aerosol particles produced from the burning and baking of a variety of 62 fuels on the ground (trees, grasses, leaf letter and other fallen debris, etc). Many of these gases 63 and aerosol particles can impact climate, air quality, and human health. For this reason, most modern air quality and climate models now consider wildland fires. However, wildland fires are 64 65 fundamentally challenging for these models, because many fine-scale and large-scale processes are entangled at the same time. In this work, we use a high resolution turbulence-resolving 66 67 numerical model to study the fine details in a wildland fire plume, with implications for large-68 scale air quality and climate models.

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70 KEYWORDS

Biomass burning, wildland fire, large eddy simulation, airborne measurements, ozone, nitrous
 acid, peroxyacetyl nitrate

74 **1. INTRODUCTION**

75 Wildland fires are a natural phenomenon and an important component of the global 76 ecosystem. Although the global trends of wildland fires remain a subject of debate (Doerr & 77 Santín, 2016), the projected burned area and the fire season length are expected to increase in 78 the 21st century, especially in the Northern Hemisphere (Doerr & Santín, 2016; Flannigan et al., 79 2013). In the United States, wildland fires are a growing concern. Records have shown that large 80 wildfire activity in the United States increased suddenly in the mid-1980s, with higher large-fire 81 frequency, longer wildfire durations, and longer wildfire seasons (Westerling et al., 2006), largely 82 driven by human-caused climate change (Abatzoglou & Williams, 2016).

83 Wildland fires often pose considerable risks to human health and property, with major impacts on air quality and the climate system. A 2010 survey showed that 32% of the U.S. 84 85 population reside in "wildland-urban interface areas", which are defined as areas where 86 structures and other human development meet or intermingle with undeveloped wildland and where wildfires have their greatest impacts on people (Martinuzzi et al., 2015). Wildland fires 87 88 emit a wide range of pollutants and greenhouse gases, such as particulate matters, carbon 89 monoxide (CO), carbon dioxide (CO₂), volatile organic compounds (VOCs), and reactive nitrogen 90 species (Liu et al., 2017), many of which produce secondary pollutants such as ozone (O₃) and 91 secondary organic aerosols in the atmosphere (Hodshire et al., 2019; Jaffe et al., 2020; Lu et al., 92 2016; Palm et al., 2020). Growing evidence indicates that wildfire smoke exposure is associated 93 with respiratory morbidity and potentially cardiovascular morbidity as well (Colleen et al., 2016). 94 Efforts to improve air quality in the U.S. show some promising trends as evidenced from 95 improvements in fine particulate matter pollution in the majority of the U.S. over the past

decades, except in the wildfire-prone regions where air quality has been worsening due to
wildfire events (Jaffe et al., 2020; McClure & Jaffe, 2018). Given the broad impacts they have on
the atmosphere and the climate system, wildland fires are a vital component in most modern
chemical transport models and climate models.

The scope of wildland fires ranges from microscale to mesoscale, with influence from both 100 101 microphysics and chemistry. Wildland fires also involve complicated feedback mechanisms, e.g., 102 the fire source and plume alter the flow field and the radiation in the atmosphere; in turn, the 103 meteorology, terrain, and the fuels also affect the fire behavior. As a result, wildland fires are 104 challenging to model. State-of-the-art fire models have been developed to explicitly solve the two-way coupling between weather and fire behavior (Coen, 2013; Mandel et al., 2011). 105 106 However, models with realistic representations of the two-way coupling with the land/fuel 107 models and the fire-weather interactions are often highly complex and rather computationally 108 demanding, even more so if coupled with chemistry. Most air quality and climate models are not 109 designed to accommodate such complexity. Because of the computational concerns, many air 110 quality/climate models are operated at coarse spatial resolutions and with longer integration 111 time steps. As a result, many key physical and chemical processes (e.g., heat release and plume 112 rise) cannot be explicitly modeled, and hence rely on subgrid parameterizations. This presents a 113 gap between "what is happening" and "what is coded in air quality/climate models".

Another gap emerges when studying wildland fires using observations. In the past decades, a number of field campaigns have revealed remarkable insights into the physical and chemical processes in wildland fire plumes. For logistic and safety reasons, the early stage of the plume (e.g., first hour or younger) remains poorly sampled. Controlled laboratory experiments

enable detailed characterizations of the fuel and the initial smoke (e.g., Hodshire et al., 2019a;
Roberts et al., 2020), however, these laboratory studies also only provide limited information on
the chemical evolution in the early stages of a wildfire plume. The early stage of the fire plume
(e.g., first hour or so) often involves plume rise and dilution, as well as rapid changes in
microphysical properties (Hodshire et al., 2019b; Konovalov et al., 2019; Schwarz et al., 2008).
The early stage of fire plumes remains poorly studied and a better understanding is needed.

124 Numerical models are a valuable tool to understand the chemical evolution in wildland 125 fire plumes. For instance, Lagrangian models have been used to track the plume-scale chemical 126 evolution (e.g., Alvarado et al., 2015; Alvarado & Prinn, 2009; Coggon et al., 2019; Hodshire et al., 127 2019; Mason et al., 2006; Trentmann et al., 2005), and have greatly advanced our understanding of chemical transformation and gas-aerosol interactions in wildfire plumes. However, such 128 129 models are not designed to resolve the spatial heterogeneity in wildfire plumes, which may 130 largely drive the variability in observations within wildfire plumes. In light of this, high resolution 131 Eulerian type plume models have been used to study the chemical evolution in wildfire plumes 132 (Alvarado et al., 2009; Trentmann et al., 2002; Trentmann et al., 2003). In particular, Trentmann 133 et al. (2003) revealed remarkable chemical heterogeneity within the fire plume, with OH radical 134 enhanced in the edges of the plume, and O_3 suppressed in the early stage of the plume but 135 enhanced in the later stage (Trentmann et al., 2003). Similar spatial heterogeneity in OH and O₃ 136 has been inferred for power plant plumes as well (Brock et al., 2002). However, such Eulerian 137 type plume models are often computationally demanding, posing a limit to the spatial resolution and/or the domain size, both of which affect the capability of capturing turbulent processes 138 (Agee & Gluhovsky, 1999; Roode et al., 2004). Moreover, the observational datasets available to 139

evaluate the chemical heterogeneity revealed in such Eulerian models were not readily availablein previous studies.

142 The motivation of this work is to use a high-resolution turbulence-resolving model (large eddy simulation, LES) to study the chemical evolution in the early stages of a wildland fire plume, 143 144 and evaluate the results using airborne observations. The LES model within the Weather 145 Research and Forecasting (WRF) package (Moeng et al., 2007) is coupled to a simple yet 146 representative chemical mechanism. The plume dynamics and interactions with the flow field are 147 resolved; the fire source characteristics, terrain, and variations in the meteorology, are highly 148 idealized, allowing to focus on the chemical evolution. The modeling experiment in this work is configured for the Williams Flats Fire (47.9392°N, 118.6183°W), a large wildfire ignited on 2 149 150 August 2019 in the State of Washington, U.S. This fire was extensively sampled by a number of 151 research aircraft and satellites during the Fire Influence on Regional to Global Environments and 152 Air Quality (FIREX-AQ) field campaign (July-September 2019). FIREX-AQ was a comprehensive 153 airborne mission jointly funded by National Oceanic and Atmospheric Administration (NOAA) and National Aeronautics and Space Administration (NASA), focusing on the trace gases and aerosols 154 155 emitted from wildfires and their impacts on air quality and climate in the United States. Airborne measurements collected from the NASA DC-8 aircraft are used to evaluate the chemical 156 157 heterogeneity predicted by the model.

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159 2. WRF-LES-CHEMISTRY MODEL

160 In this work, the LES model in the WRF package (version 4.0) is run in an idealized 161 configuration, with a domain size of 70 km × 30 km × 6 km. The horizontal grid resolution is 100

162 m, and the vertical resolution is ~78 m near the surface and up to 138 m near the top of the 163 model. The total number of grid boxes is 126,000,000, and the time step is 1 second. More details 164 of the LES model configuration are provided in the Supporting Information (Section S1). This LES 165 model is coupled with a simple yet representative chemical mechanism, consisting of 39 166 transported tracers and 69 reactions, covering major tropospheric chemical reactions involving 167 key NO_x/VOC/O₃ chemistry (Table S1). This chemical mechanism is condensed based on the 168 widely used MOZART T1 mechanism in the WRF package (Knote et al., 2014). Photolysis 169 frequencies are calculated using the Fast Tropospheric Ultraviolet-Visible scheme (Tie et al., 170 2003). Two inert aerosol tracers, black carbon (BC) and organic carbon (OC), are included in this 171 mechanism, to account for the aerosol impacts on radiation and chemistry, via both photolysis frequencies and heterogeneous chemistry. The LES model is initialized using the vertical 172 173 soundings (potential temperature, water vapor, and wind) and vertical profiles of chemical 174 species from a mesoscale WRF-Chem simulation for the Williams Flats fire. More details of this mesoscale WRF-Chem simulation are given in the Supporting Information (Section S2, Figures S1 175 and S2). The LES model is spun-up for two hours (from 20:00:00 to 22:00:00 3 August 2019, all 176 177 times are in UTC unless otherwise noted. UTC = Coordinated Universal Time) and simulated for three more hours (22:00:00 3 August 2019 to 01:00:00 4 August 2019). The fire plume in this LES 178 179 model is driven by surface fluxes of sensible heat and chemicals, characterized based on 180 measurements: the fire size is 0.66 km² and the sensible heat flux is 10⁴ W/m², both 181 approximated from the GOES-16 fire detection product (Li et al., 2020) for the Williams Flats Fire. The emission fluxes of chemical species are adjusted until a reasonable agreement between the 182 183 modeling results and the airborne measurements is reached (Section 3). The adjusted emission

fluxes are given in Table S2. This LES configuration does not have terrain, posing another limitation to the capability of resolving the observed plume transport and dispersion.

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3. AIRBORNE OBSERVATIONS USED FOR MODEL EVALUATION

188 On 3 August 2019, the DC-8 aircraft sampled the Williams Flats fire plume in two flight 189 patterns (Figure 1): Firstly, an overpass (bird's eye view) above the plume from 21:45 UTC to 190 21:59 UTC (3 August 2019) over the plume provides the structure of the plume along the direction 191 of the spread of the fire plume using the downward-looking NASA Langley Airborne Differential 192 Absorption Lidar (DIAL). Secondly, the aircraft descended and flew through the plume in a 193 "lawnmower" pattern, providing a number of semi-Lagrangian transects. The one second merge dataset provided by the NASA Langley Research Center is used in this work, including high time 194 195 response measurements from a number of instruments, except for the NCAR Trace Organic Gas 196 Analyzer (TOGA) with a time resolution of ~105 seconds (with a sampling window of ~32 seconds). 197 Table S3 summarizes all measurements used in this work.



Figure 1. (Left) Flight tracks and (right) 532 nm backscatter ratio measured using NASA DIAL. The flight altitudes during Segments A and B (solid black lines in the Right panel) are overlaid on top

of the "bird's eye view" curtain in the Right panel, but Segments A and B occurred at differenttimes than the "bird's eye view" flight.



Figure 2. Selected airborne measurements collected during the "in-plume" Segments A and B, two transects each. The full view of Segments A and Segments B is provided in Figure S3.

The second stage (semi-Lagrangian transects) consists of two major segments: Segment 207 A, from 22:20-23:15 UTC (3 August 2019), and Segment B, from 00:30-01:24 UTC (4 August 2019). 208 209 Segments A and B both sampled the first ~50 km of the plume, however, there are several marked differences: (i) The GOES satellite image suggests that, before the DC-8 aircraft sampling, the 210 211 Williams Flats Fire region was dominated by a north wind in the boundary layer, transporting the 212 Williams Flats fire towards the south. Starting from ~17:00 UTC the low-level wind direction 213 shifted to the west-southwest, transporting the smoke to the east-northest. Therefore Segment 214 A sampled some aged/diffused smoke (due to the shift in wind direction), while the smoke 215 sampled during Segment B was relatively well defined "fresh" smoke. (ii) Based on the

216 combination of remote sensing and in situ measurements, the DC-8 aircraft likely flew through 217 the dense "core" of the plume during Segment B, while Segment A likely skimmed the upper edge 218 of the plume. During Segment B, the sampled plume was rather optically opaque such that the 219 DIAL laser signal was severely attenuated at the center of each transect (Figure 2). Accordingly, 220 the measured photolysis frequencies, such as $j-NO_2$, were suppressed at the center of each 221 transect (Figure 2). O₃ was also suppressed at the center of the plume due to the rapid reaction 222 with NO (Figure 2). During Segment A, however, measured CO and NO_x levels inside the plume 223 were lower than that during Segment B, and the $j-NO_2$ reduction inside the plume was also 224 weaker than that during Segment B (Figure 2), implying that the Segment A perhaps mostly 225 sampled the upper part of the plume, which is optically thinner and with lower levels of primary 226 pollutants. Interestingly, O₃ during Segment A was always enhanced at the center of each 227 transect (Figure 2), similar to the O_3 measured when the DC-8 aircraft was entering and leaving 228 the plume in Segment B (i.e., the edges of the plume sampled in Segment B), providing additional 229 evidence that the aircraft sampled the top part of the plume during Segment A. The modified 230 combustion efficiency, defined as $\Delta CO_2/(\Delta CO + \Delta CO_2)$ (Yokelson et al., 2008), did not vary 231 dramatically between the two Segments (both 0.89-0.92), implying the average burning 232 condition remained relatively consistent during these two Segments. To sum up, the DC-8 233 measurements during Segment A may not reflect the center of the plume, and thus Segment B 234 is mainly used for model evaluation.

A variety of merged files have been created for each research flight. In this work, the 1-Hz merge files are used unless otherwise noted. More details of the merging algorithm can be found elsewhere (https://www-air.larc.nasa.gov/missions/etc/onlinemergedoc.pdf).



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239 Figure 3. Modeled plume-wide averages and standard deviations (red lines and shadings) of 240 selected compounds and parameters, compared to the airborne measurements averages and 241 standard deviations (black symbols and error bars) during each transect during Segment B at X 242 km downwind. The measurements are averaged using the smoke flag provided in the 1-Hz merge, 243 while the modeled plume-wide averages are calculated for each plume cross section at X km 244 downwind. OC is calculated from organic aerosol and OA/OC ratio measured using the CU HR-AMS. The measurement-derived plume age is estimated from the measured winds and distance 245 246 from the fire location (provided in the 1-Hz merge), while the plume age in the model is estimated 247 based on Equation 1. Total measured volatile organic compounds (TVOCs) are listed in the main 248 text and Table S3. Total OH reactivity is calculated for TVOCs as well as CO and methane. 249

Figure 3 compares the model chemical species and parameters to the airborne observations collected during Segment B, providing a semi-quantitative comparison of the plume chemical evolution as a function of distance in km from the fire source (X). As shown, the model captures the chemical characteristics within the plume reasonably well, as the modeled plumewide means of gaseous (CO, NO_x, HONO, HCHO, O₃, and PAN) and aerosol constituents (BC and

255 OC) are both in good agreement with observations, although the observations do show larger 256 variations than the modeled. Total VOCs were calculated as the sum of the mixing ratios of 257 ethane, propane, i-butane, n-butane, i-pentane, n-pentane, i-butene + 1-butene, isoprene, MVK, 258 MACR, benzene, toluene, formaldehyde, acetaldehyde, methanol, ethanol, acrolein, acetone + 259 propanal, furan, 2-methylfuran + 3-methylfuran, furfural, phenol, and styrene, reported from 260 several different instruments on board the DC-8 (Table S3). Notably, the simplified mechanism 261 captures roughly about half of the measured total VOCs, yet the modeled total OH reactivity is 262 comparable to the calculated total OH reactivity based on measurements available (Figure 3).

263 In the model, two fire tracers are added to track the physical age of any air parcel in the 264 plume: one tracer with 1-hour fixed lifetime, and the other inert one with infinite lifetime. The 265 plume physical age (in hours) is given by:

266 Physical age =
$$-\ln\left[\frac{Tracer \ 1 \ (\tau=1 \ hr)}{Tracer \ 2 \ (\tau=\infty)}\right]$$
 Equation 1

267 As shown in Figure 3, the modeled physical age also shows good agreement with that estimated 268 using airborne measurements (estimated from measured winds and distance from center of 269 transect to the fire source), implying that the meteorological conditions (especially wind) are 270 decently represented in this simple model configuration. The total OH reactivity in the model 271 (including CO and methane) is also in reasonable agreement with the total OH reactivity 272 calculated using airborne measurements, despite the fact that the chemical mechanism used in 273 this model is highly condensed. Overall, this figure reveals that the highly simplified model 274 configuration and chemical mechanism captures the major chemical characteristics of the plume. 275 We discuss the detailed plume physics and chemistry in the following sections.

4. PLUME DYNAMICS IN THE EARLY STAGE OF THE PLUME

278 The high-resolution LES model resolves the turbulence induced by the fire source and the 279 interactions with the background flow field. Due to the strong heat release from the fire, the plume rapidly rises in the early stage, characterized by high updraft speed in the early stage 280 281 (modeled mean updraft: 3 m s⁻¹ in Figure 4; maximum updraft in the model reaches up to 12 m 282 s⁻¹). The strong updraft injects the plume into a lofted layer that is 1.5-2.5 km above the surface, 283 consistent with the DIAL measurements (~1.5-3 km above ground level, Figures 1 & 2). The rapid 284 updraft at the center of the plume leads to downdraft and hence small circulations near the 285 plume, as indicated by the streamlines in the cross section plots (Figure 4). The rapid plume rise and the circulations also lead to dilution. In the model, the plume 1st order dilution rate can be 286 quantified by tracking the decay of the inert fire tracer: 287

288 Dilution rate
$$(s^{-1}) = \frac{dln([Tracer 2])}{dt}$$
 Equation 2

Where [*Tracer 2*] is the concentration of the inert fire tracer and *t* is the plume physical age. As shown in Figure 4, the plume dilution rate is on the order of 0.001 s⁻¹ in the early stage of the plume, which is decreased to and maintained at the order of 0.0001 s⁻¹ in the late stage of the plume. The plume dilution leads to entrainment of the background air.

293 Different processes drive the plume dilution. In the uprising stage (e.g., X = 1 km), the 294 dilution is mostly driven by the circulations induced by the plume-rise, therefore faster at the 295 bottom of the plume, where the entrainment flows enter the plume. Later on, the plume rise has 296 ceased (e.g., X = 10 and 30 km, with mean vertical velocity (w) close to zero), the plume-wide 297 mean dilution is nearly an order of magnitude slower than that in the uprising stage, and is still 298 faster at the edges of the plume.

Notably, the model predicts the lower part of the plume may be strongly mixed throughout the boundary layer, due to the convective mixing within the boundary layer. As a result, the horizontal dispersion in the lower part of the plume is slower, compared to the upper part (Figure 4). This is typical for daytime wildland fire plumes with low-intermediate plume injection, which is also consistent with the DIAL images that diffused smoke extends from right below the dense smoke all the way to the surface (Figures 1 & 2).



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Figure 4. (Top) Streamlines at the centerline of the plume (curtain); (middle) plume-wide average updraft speed and dilution rate (shadings: standard deviations); (bottom) dilution rate cross section at X = 1, 10, and 30 km. The thickness of the streamlines roughly represents the wind speed. The vertical black lines represent the locations of the cross sections. The plume (brown colored area) is defined as CO > 150 ppb.

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312 5. PHYSICAL AND CHEMICAL EVOLUTION IN THE EARLY STAGE OF THE PLUME

The LES configuration coupled with chemistry reveals how the plume evolves over the course of dispersion. Figure 5 briefly shows how the plume ages physically, and also provides an

overview of the chemical evolution in fresh plume via two important reactive species: OH radicals

for photochemistry, and NO₃ radicals for dark chemistry. The physical age shows considerable heterogeneity vertically - the lower part of the plume is generally older than the upper part of the plume (Figure 5), which is a combined result of the vertical wind gradient, and enhanced mixing within the boundary layer.

OH radicals are the major driver of atmospheric oxidation. The model predicts very 320 321 interesting heterogeneity within the fire plume: in the early stage of the plume (first ~8 km or 322 so), OH is severely suppressed in the bulk of the plume, but enhanced at the upper most edge of 323 the plume (Figure 5). This is mainly because the solar radiation is severely attenuated in the 324 plume due to the presence of high levels of aerosols. Further downwind, OH radicals become 325 enhanced in the upper part of the plume (~8-30 km), or even the entire vertical extent of the plume in the later stage (~30+ km, Figure 5). NO₃ radical, a major oxidant in the atmosphere 326 327 under dark conditions, is also present in the modeled plume at the same time, but only in the 328 lower-most part of the plume and, to a lesser extent, below the thick plume (Figure 5). In 329 particular, the modeled NO₃ mixing ratio reached ~3 ppt in the lower-most part of the plume in 330 the early stage. With the dilution and entrainment, the NO₃ oxidation products in the lower-most 331 part of the plume or below the plume may be mixed into the bulk of the plume. A recent study 332 reported rapid aging of biomass burning aerosols under dark conditions (driven by NO₃ radical), 333 a potential source of organic aerosols that has not been widely considered (Kodros et al., 2020). 334 The chemical heterogeneity within fresh wildland fire plumes warrants further investigation.





Figure 5. Modeled curtains of the plume physical age and mixing ratios of OH, NO₃, CO, HONO,
and the photolysis frequency of HONO (j-HONO). The curtains are at the centerline of the plume.
The black lines represent the roughly defined the plume boundary (CO = 150 ppb)

340 6. CHEMICAL TOMOGRAPHY IN THE PLUME

In this section, we discuss the modeled chemical tomography inside the plume, as well as 341 the observational evidence. We focus on the cross-transect variations of a few species: CO, O₃, 342 PAN, maleic anhydride (a product from furan oxidation), and HONO. Again because of the 343 344 difficulties in providing realistic initial and boundary conditions to the LES model, also the fact 345 that this simplified LES configuration does not consider terrain impact, it is unrealistic to compare 346 the modeling outputs to the DC-8 measurements precisely. Therefore, the modeled fields are not 347 sampled along the flighttrack. Instead, the comparison is focusing on whether the model is capable of reproducing the broad features in the observations. In this section, several aircraft 348 349 transects showing distinct cross-transect variations are selected, and compared to cross sections 350 of the modeled plume sampled at the same downwind distances at 2 km above the model surface 351 (aircraft altitude: ~2.8 km during Segment B, mean terrain height 0.85 ± 0.32 km).

352 Figure 6 shows the observed cross-transect variability of CO, a (mostly) primarily emitted 353 fire tracer with fairly long atmospheric lifetime. As shown, the observed CO is generally elevated 354 at the center of the plume, except at X = 30 km the observed CO shows two distinct "modes", 355 possibly due to the influence of very aged/diffused smoke. The modeled cross sections at the 356 same transect distances show very good agreement with observations at X = 5-15 km. At X = 30357 km, the model is able to capture one of the "modes". Because of the simple model configuration 358 (e.g., short simulation time, small domain, no smoke in the boundary condition), it is challenging 359 for this model to capture aged/diffused smoke. Overall, Figure 6 demonstrates that the model 360 captures the plume dispersion and transport quite well (at least in the first ~15 km).

361 Similar cross-transect variations for j-HONO, NO_x, HCHO, BC, and OC are provided in the Supporting Information. In general, the model well captures the cross-transect variations of these 362 363 constituents in the first ~15 km (Figure S4-S8), and they are all enhanced inside the plume relative 364 to that at the edges. The model captures j-HONO outside the plume but overestimates j-HONO 365 inside the plume by ~20% (relative to the background j-HONO, Figure S4). In addition, the *in situ* measurements show that plume "optical width" (the plume width characterized by the measured 366 j-values) appears to be wider than the plume "chemical width" (the plume width characterized 367 by the chemical constituents). Although the modeled plume "chemical width" is generally well 368 369 captured by the model in the first ~15 km (Figures 6, S4-S8), the plume "optical width" is 370 underestimated by the model (Figure S3). The model underestimation of j-value suppression 371 inside the plume and the underestimation of the plume "optical width" may be due to the 3-D 372 radiation effect (Trentmann, et al., 2003) that is not captured by FTUV (Tie et al., 2003), although we cannot rule out the possibility of diffused or high level smoke/cloud not captured in the model. 373

Moreover, only BC and OC are considered in this work. Although CU HR-AMS measurements suggest that organic aerosols alone accounts for 93% of measured PM₁ mass, recent studies found that brown carbon may also contribute to light absorption in biomass burning plumes (Palm et al., 2020). In summary, primarily fire-emitted long-lived tracers such as CO, BC show Gaussian-like cross-transect profiles. In the next few sections we will examine a few compounds with drastically different cross-transect profiles



381 Figure 6. Measured CO (top) during Segment B and modeled O_3 cross sections (bottom) at 10 km,

15 km, and 30 km downwind of the fire source. Also shown in the top panel are the modeled CO
sampled at the same distance from the fire, at ~2 km above the model surface.



Figure 7. Measured O_3 (top) during Segment B and modeled O_3 cross sections (bottom) at 10 km, 15 km, and 30 km downwind of the fire source. Also shown in the top panel are the modeled O_3 sampled at the same distance from the fire, at ~2 km above the model surface.

390 6.1. CROSS-TRANSECT VARIATION OF OZONE

391 As shown in Figure 7, O₃ shows considerable heterogeneity within the modeled plume. In 392 the early stage (X = 10 km), the model predicts that O_3 is suppressed inside the plume (up to 30 393 ppb lower than the background O_3 at the same height), because of fire-emitted NO. Normally, 394 NO₂ produced from NO-O₃ reaction would undergo photolysis and produce O₃ back; but j-NO₂ is 395 strongly suppressed within thick fire plumes, and NO₂ will not photolyze and hence serves as a 396 temporary reservoir for O_3 . The modeled O_3 during this transect compares well to the 397 measurements. At X = 15 km, measured O_3 is recovered at the center of the transect and is 398 elevated at the edges, and the model shows similar broad features. At X = 30 km, the measured 399 O_3 is well mixed throughout the plume, and is enhanced relative to the background by 10-40 ppb. 400 The model predicts a consistent O_3 trend at X = 30 km that O_3 is enhanced throughout the plume, 401 compared to the background air.

402 As discussed in Section 3, Segment B is used to examine the cross-transect variations of O₃, since the aircraft flew through the core of the plume multiple times during Segment B. 403 404 However, the edge effect on O₃ is also visible in Segment A, when the aircraft likely sampled the 405 upper edge of the plume. The measured O_3 is enhanced relative to the background in all transects 406 during Segment A (Figure 2) by up to 100 ppb, roughly 2 times greater than the O₃ sampled at 407 the edges of the plume during Segment B. This may indicate possibly stronger O₃ enhancement 408 at the upper edge of the plume than the sides of the plume. The model also shows enhanced O_3 levels at the upper edge of the plume (Figure 7) relative to the sides of the plume, but the 409 410 magnitude of the O₃ enhancement is again underestimated by the model.

411 Overall, the model qualitatively captures the observed O₃ variations in these transects, 412 but seems to underestimate the O₃ enhancement at the edges. The plume-wide average of O₃+NO₂ reaches ~40 ppb higher than that in the background air after ~2 hours of aging, indicating 413 net O₃ formation in this modeled plume. Note that in previous studies, O₃ is not always found to 414 415 be enhanced relative to the background air especially in more aged (e.g., a few days or older) 416 biomass burning plumes (Alvarado et al., 2010; Parrington et al., 2013). Because of the highly 417 simplified VOC chemistry used in this model, we do not intend to further investigate O₃ formation 418 and variability, which is beyond the scope of this study. But we show that the observed cross-419 transect variations in O₃ can be qualitatively explained by including simple chemistry with a 420 turbulence-resolving model.



Figure 8. Measured PAN (top) during Segment B and modeled PAN cross sections (bottom) at 10
km, 15 km, and 30 km downwind of the fire source. Also shown in the top panel are the modeled
PAN sampled at the same distance away from the fire, at ~2 km above the model surface.

426 6.2. ELEVATED PAN LEVELS AT THE PLUME EDGES

PAN is an important reservoir of NO_x. Previous studies have reported rapid PAN formation
in biomass burning plumes (Alvarado et al., 2010, 2015; Fischer et al., 2018; Jaffe et al., 2013; Liu
et al., 2017; Singh et al., 2012). Because of the limited capability in resolving the transport and

chemistry of reactive nitrogen species in early biomass burning plumes, large scale models
sometimes partition a fraction of the reactive nitrogen emissions as PAN (Fischer et al., 2014),
and this process is subject to potential uncertainties because of non-linear dependency of PAN
formation on VOCs and NO_x (somewhat similar to O₃). In this section, we examine PAN formation
in the modeled fresh plume as the plume ages.

435 As shown in Figure 3, the modeled plume-wide mean PAN is in reasonable agreement 436 with observations. Figure 8 shows the cross-transect comparison for PAN. Interestingly, at X = 10437 and 15 km, measured PAN is elevated at the edges of the plume compared to that at the center. 438 In particular, at X = 15 km, the measured PAN is nearly doubled at the edges. The model 439 reproduces this feature at X = 15 km but tends to overestimate PAN by 2-3 ppt. The edge 440 enhancement of PAN occurs because in the early stage of the plume, PAN formation is limited by 441 the initial OH attack on its VOC precursors, and OH production is more active at the edges of the 442 plume. At X = 30 km, however, it appears that the measured PAN is spread out in a wider range than the model predicts, similar to a few other compounds (Figures 6 & S5-S8). Overall, PAN 443 444 formation accounts for 10-70% of the NO_x loss in this modeled plume. In summary, the observed 445 PAN shows interesting core-edge differences in the early stage of the plume, corroborating faster 446 photochemical aging at the edges of the plume. This effect may be largely diminished in the later 447 stages of the plume, because dilution further lowers the aerosol loading inside the plume, and 448 hence the suppression of photochemistry inside the plume becomes weaker. Notably, elevated 449 PAN is often found outside the studied plume (i.e., other pollutants such as CO are close to the 450 background levels), providing additional evidence of possible diffused/aged smoke.

452 6.3. FURANS AS A FIRE-TRACER AND INDICATOR OF PHOTOCHEMICAL AGING

453 Furans are an important VOC class emitted from biomass burning, contributing 454 substantially to the total VOC reactivity in biomass burning plumes (Coggon et al., 2019). In this section, we discuss the simplified furan chemical mechanism included in the LES model (Table 455 S1), as well as the photochemical aging tracked by furan and one of its unique oxidation products, 456 457 maleic anhydride. For simplicity, "furan" in this model is a lumped species representing total 458 furan species, and the modeled furan is compared to the TOGA measurements (sum of furan, 2-459 methlyfuran, 3-methlyfuran, and furfural). We do not compare the modeled furans to the PTR-460 MS measurements during Segment B since secondary products and lesser-reactive isomers likely result in measurement biases after several hours of oxidation (Coggon et al., 2019; Koss et al., 461 462 2018).



463

Figure 9. TOGA measured total furans (top) during Segment B and modeled furan cross sections (bottom) at 15 km, 20 km, and 30 km downwind of the fire source. Also shown in the top panel are the modeled furan sampled at the same distance from the fire, at ~2 km above the model surface.



469 470

Figure 10. PTR-MS measured maleic anhydride (top) during Segment B and modeled maleic anhydride cross sections (bottom) at 10 km, 15 km, and 30 km downwind of the fire source. Also shown in the top panel are the modeled maleic anhydride sampled at the same distance from the fire, at ~2 km above the model surface.

475

476 Furans are primarily emitted from fire, and their major sink is the reaction with OH 477 radicals, and to a lesser extent, with O₃ and NO₃ (Coggon et al., 2019). Figure 9 shows the 478 measured and modeled furans. As shown, the modeled and measured furan show reasonable 479 agreement at X = 15 and 20 km, but not at 30 km. The modeled cross-transect variability of furan at X = 15 is similar to other primary compounds such as CO (Figure 6), i.e., concentrated at the 480 center/core of the plume but reduced at the edges. Its oxidation product, maleic anhydride, 481 482 however, is enhanced at the edges according to the measurements at X = 15 km. Such core-edge discrepancy is consistent with PAN (Section 6.2), and also implies faster photochemical aging at 483 the plume edges. This core-edge discrepancy at X = 15 km is qualitatively captured by the model 484 (Figure 10). At X = 10 km, the model also predicts similar core-edge discrepancy. The measured 485 maleic anhydride at X = 10 km, however, shows a peak at the center of the transect in addition 486 to the two peaks at the edges. This might imply that perhaps a small fraction of maleic anhydride 487 488 may be directly emitted from the fire source. Maleic anhydride emissions have not been reported 489 in the literature, but a reanalysis of measurements conducted during the FIREX Firelab 2016 study

490 (Coggon et al., 2019; Koss et al., 2018) shows that PTR-MS measurements of maleic anhydride
491 represent 1-2% of furan emissions, suggesting that emissions likely contribute to higher plume
492 center mixing ratios at X = 10 km.

493 The impact of plume chemical heterogeneity on furan decay is also examined using the 494 modeling results. We discussed in Section 5 and Section 6.1 that ozonolysis and NO_3 oxidation 495 may be active in optically dense smoke, and furan can be oxidized by O₃, and NO₃ as well. We 496 found that OH oxidation is the dominant furan loss pathway, accounting for 78-89% of total furan 497 loss. Ozonolysis and NO₃ oxidation account for up to 16% and 7% of total furan loss, respectively. 498 The relative importance of the three oxidation pathways evolves with time: OH oxidation 499 accounts for >90% in the early stage, while ozonolysis and NO₃ oxidation together contribute to <10% furan oxidation in the early stage. Further downwind, with the gradual recovery of O₃, the 500 501 contribution of ozonolysis increases to up to \sim 30% at X = \sim 30 km, where NO₃ oxidation also 502 contributes to another \sim 20% of total furan loss. NO₃ oxidation mostly occur in the lower part of 503 the plume, while furan mostly exists in the bulk of the plume aloft. Therefore, the NO₃ oxidation 504 is partially affected by the plume dynamics. The impact of chemical heterogeneity on the overall 505 VOC reactivity is challenging to represent in small-scale Lagrangian type models or large-scale 506 models with coarser spatial resolutions.

507

508 6.4. HONO: SECONDARY PRODUCTION, IMPACT OF PLUME DYNAMICS

509 HONO is an important precursor of OH radicals in the atmosphere and is emitted from 510 wildland fires (Peng et al., 2020; Theys et al., 2020). In this modeling work, we include both 511 primary HONO emission and secondary HONO formation from NO₂ heterogeneous reaction on

512 BC and OC aerosols (both reasonably well captured by the model, Figures S7-S8). The reactive uptake coefficient of NO₂ is mostly on the order of 10^{-5} - 10^{-4} in the literature (Ammann et al., 513 2013). Therefore, we assume the NO₂ uptake coefficient on aerosols is 5×10⁻⁵ in this work. Model 514 515 sensitivity tests are also performed to examine the possible secondary HONO production. As for 516 the primary HONO emission, it depends on the fuel type, which is primarily grassland with tall 517 grass, timber, and brush in this region (https://inciweb.nwcg.gov/), and the average HONO/NO 518 emission ratio (molar basis) for savannah/grassland and temperate forest are 12% and 7%, 519 respectively (Andreae, 2019). This is consistent with a recent laboratory study in which HONO 520 accounts for 13% of NO produced from stack fires (Roberts et al., 2020). In this work, we found a 521 fire HONO emission of 11% of fire NO emission (molar basis) yields the best agreement when 522 compared to airborne HONO measurements (Figure 3): modeled plume-wide mean HONO 523 exceeded 10 ppb in the very beginning stage of the plume, which then decreased to 2-4 ppb at 524 10-25 km downwind, and eventually to 1-2 ppb at 30-40 km downwind.

Figure 11 shows the cross transect comparison for HONO. As shown, the measurements 525 526 suggest that HONO is concentrated at the center of the plume, and the modeled HONO levels 527 across the transects show good agreement with observations, especially in the early stage (X =528 10 and 15 km). The model also shows that HONO is always depleted at the edges, especially the 529 upper edge of the plume (Figure 11). The cross transect comparison for HONO/NO₂ ratio is given 530 in Figure S10. This ratio remains fairly constant at the center of the modeled plume (with 531 secondary HONO production), but rapidly decreases at the edges of the plume. At X = 30 km 532 downwind, the model drastically overestimates HONO, mainly due to the overestimation in NO_x (Figure S5). This is likely because the sink of NO_x is not well captured by this highly condensed 533

534 chemical mechanism. Notably, we discussed previously that the model fails to predict the plume 535 profile at X = 30 km, since the observations show two distinct "modes" at X = 30 km and model 536 captures only one of them (Figures 6, 8, 10, S5-S8). Figures 11 and S5 show that the observed 537 HONO and NO_x are higher in the other "mode", consistent with the modeled HONO and NO_x, 538 respectively. This may also imply a change in meteorological condition which alters the plume 539 transport that is not captured by the simple model configuration.



Figure 11. Measured HONO (top) during Segment B and modeled HONO cross sections (bottom)
at 10 km, 15 km, and 30 km downwind of the fire source. Also shown in the top panel are the
modeled HONO sampled at the same distance away from the fire (solid black line: with secondary
HONO production from NO₂ heterogeneous reaction; dashed black line: with only primary HONO).



546

540

- 547 Figure 12. Measured plume-wide averages of HONO compared to modeled in different model 548 scenarios: with both primary and secondary HONO (dark red, same as that in Figure 3), with only 549 primary HONO and no secondary HONO production (blue), primary HONO emission increased by 550 a factor of 5 and no secondary HONO production (green).
- 551

We now discuss the potential secondary HONO production from NO₂ heterogeneous

reactions on aerosols in the modeled plume. As shown in Figure 11 and Figure 12, the model

553	simulation with secondary HONO production yields best agreement with observations. Without
554	secondary HONO production, in order to capture the detected HONO levels at $X = 10-15$ km,
555	primary HONO emission needs to be increased by a factor of \sim 5 (i.e., about half of NO emission),
556	which still underestimates HONO in the later stage (X > 20 km, Figure 12). The primary HONO
557	emission contributes substantially to the total HONO levels in the early stage: the modeling
558	results suggest that >93% HONO in the early stage is from primary emission, which decreases to
559	55% at 40 km downwind (observed HONO: 1.2 ± 1.1 ppb at X = 37 km, Figure 3). Note that the
560	modeled simulation with heterogeneous conversion from NO_2 to HONO may not necessarily
561	mean a considerable increase in HONO/NO $_2$ ratio (Figure S9). To sum up, although no direct
562	evidence can be provided in this work, our combined observational and modeling analysis does
563	suggest that a secondary HONO source better explains the observed HONO levels and variability
564	in this particular plume.



565

Figure 13. Modeled apparent HONO decay rate (first order) at the center of the plume, as well as the modeled dilution rate and local HONO photolysis rate also at the center. This plot is created using the model simulation without secondary HONO production. Plume center is defined with top 5% modeled CO.

570 We now take HONO as an example to demonstrate the impact of plume dynamics on the

571 chemical evolution. We showed in Figure 4 that the modeled plume dilution (mixing) is faster in

572 the early stage (driven by strong updraft and entrainment) but is maintained relatively constant

573 in the later stage. The physical mixing within the plume may pose a limiting factor for slow 574 chemical reactions. At the center of the plume, solar radiation is suppressed and hence HONO 575 photolysis is low, leading to a local photolysis lifetime of half an hour or longer. However, as 576 shown in Figure 13, the apparent decay rate of HONO at the center of the plume is nearly 9 times faster than the local photolysis rate, but is very close to the physical dilution rate. That is, 577 578 although in the early stage photochemistry is slow in the majority of the plume, HONO still decays 579 fairly fast – with an effective lifetime of several minutes at the dark center of the plume. The 580 rapid mixing bring HONO from the center of the plume to the edges and upper part, where HONO 581 is quickly destroyed via photolysis. Even at X = 40 km, physical mixing still contributes to half of 582 apparent HONO decay at the center of the modeled plume. This exercise demonstrates the 583 importance of plume dynamics in the chemical evolution in fresh biomass burning plumes.

584 HONO photolysis is the most important primary OH source in this modeled plume, while 585 other primary OH sources (such as the photolysis of O_3 and aldehydes) are 1-2 orders of 586 magnitude weaker (Figure S10). In addition to the primary OH sources, secondary OH sources (i.e., recycled from HO₂) also plays a key role in the simulated plume, which is on average an 587 588 order of magnitude higher than the primary OH source (Figure S10), implying an efficient HO_{x} recycling in this modeled plume mediated by NO_x. This secondary HO_x source also efficiently 589 590 propagates HO_x chemistry into the darker interior of the plume, while photons (that can trigger 591 photochemical reactions) are quickly quenched at the edges and do not reach the bulk of 592 optically dense plumes (e.g., Figure 5). In summary, this section highlights the importance of 593 HONO in wildland fire plumes. The reactive uptake coefficient of NO₂ as well as the dependency on particle type and other parameters or conditions warrant further investigation. 594

595

596 **7. IMPLICATIONS FOR MODELS WITH COARSER HORIZONTAL RESOLUTIONS**

597 In principle, LES schemes resolve major (large) eddies, through which momentum, energy, 598 and other passive scalars are transported. This sets the upper limit of the horizontal grid size in 599 LES models to about 0.1 km. Most planetary boundary layer schemes (widely used in regional 600 and global chemical transport models) are not designed to resolve the energy-baring eddies, 601 which are parameterized as sub-grid processes instead. This sets the lower limit of the horizontal 602 grid size of most planetary boundary layer schemes to about 1 km. Moreover, operating with small grid sizes (or high horizontal resolution) often also requires shorter integration time steps, 603 which also leads to an increase in computational cost. Currently, most regional chemical 604 605 transport models are operated at horizontal grid resolutions of a few to tens of km. In this section, 606 we discuss the impact of grid resolution on the chemical evolution of wildland fire plumes.



607

Figure 14. Sensitivities of maximum O_3 formation as a function of fire-emitted NO_x and nonmethane VOCs. The thick black line represents the transition between the NO_x -saturated and the NO_x -limited regimes. The circle/error bars represent the airborne measurements averaged during the Segment B of Williams Flats Fire (3-4 August 2019). The thin dashed line is the 1:1 dilution line for Williams Flats fire.

614 Biomass burning emission inventories report the emissions of pollutants on a mass basis. 615 When using an emission inventory in a particular model, the emissions need to be re-gridded into 616 the horizontal resolution at which the model is operated. Usually, an area weighted interpolation 617 method would be used, since the total amount of emitted pollutants need to be conserved. If the 618 model horizontal grid size is larger than the fire size, to maintain mass conservation, all fire 619 emitted tracers are immediately diluted within the grid cell, leading to an instantaneous 620 numerical dilution. For instance, the area of the fire source in this model is 0.6 km², and if 621 mimicking this particular fire using a model with 4-km grid resolution (with same vertical 622 resolution), the emission fluxes are reduced by a factor of 26.7 ($4^2/0.6$); if the model grid 623 resolution is 1 km, the emission fluxes are still reduced by a factor of 1.5 ($1^2/0.6$). The atmospheric 624 chemical system is highly non-linear, especially the O₃ chemistry, such that dilution may lead to 625 a shift in chemical regimes. Figure 14 shows the sensitivity of maximum O₃ formation as a 626 function of fire-emitted NO_x and VOCs. This plot is created using a box model with a near-explicit 627 chemical mechanism (MCM v3.3.1), including furan chemistry (Coggon et al., 2019), and 628 constrained to airborne measurements collected from Segment B (in-plume only) from the 629 Williams Flats Fire. Also shown on this plot is the dashed line representing the impact of dilution 630 - numerical or otherwise. The dilution always leads to a reduction in the maximum O₃ formation 631 potential, but the magnitude of such reduction depends on the chemical regime. As shown, the 632 dilution line for the Williams Flats Fire is close to the "ridge" (transition between NO_x/VOC-633 sensitive regimes), and hence dilution would lead to a rapid decrease in O_3 formation potential, 634 and perhaps a change in the chemical regime as well (from NO_x -saturated/VOC-sensitive regime 635 to the VOC-saturate/ NO_x -sensitive regime).





Figure 15. Modeled lower tropospheric column densities of O₃ and NO₂ with horizontal resolution
of 0.1 km, 1km, and 4 km. All column densities are integrated from the surface to 3 km. The 0.1
km resolution model is the LES model that is discussed throughout this work, while the 1 km and
the 4 km models are using YSU scheme and are driven by the same fire emissions but regridded
accordingly. Other configurations in the 1km and 4 km models are identical to the 0.1 km model.
See main text for more details.

644	Figure 15 illustrates the impact of model grid resolution on fire-induced column O_3 and
645	NO ₂ , by comparing the results from the 0.1-km LES model to that from two model configurations
646	with coarser horizontal resolutions: 1 km and 4 km. The 1-km and 4-km models shown in Figure
647	15 are configured in the same way as the 0.1-km LES model, except the fire emissions are re-
648	gridded to 1 km and 4 km grids accordingly (mass conserved), and the Yonsei University (YSU)
649	planetary boundary layer (PBL) scheme (Hong et al., 2006) are used in the 1 km and 4 km models.

As discussed before, the underlying principle of LES sets the upper limit to the grid spacing toabout 0.1 km, therefore a grid spacing of 1 km or 4 km is inappropriate for LES.

652 As shown, the 0.1 km model reveals O₃ suppression in the first ~14 km or so, and O₃ 653 enhancement is visible starting from \sim 14 km. At X = 40-50 km, the predicted O₃ column reaches \sim 3.71×10¹⁷ molecule cm⁻² in the 0.1 km model, or \sim 0.70×10¹⁷ molecule cm⁻² higher than the 654 655 background O₃ column (~ 3.01×10^{17} molecule cm⁻²). Both the 1 km and 4 km models capture the 656 O₃ suppression in the first ~10 km or so. However, the maximum O₃ enhancement in the 4 km model appears at X = 16-28 km at \sim 3.27 \times 10¹⁷ molecule cm⁻², or 0.27 \times 10¹⁷ molecule cm⁻² higher 657 658 than the background, and the O₃ enhancement quickly diminished beyond $X = \sim 32$ km. That is, 659 the 4 km model does not efficiently transport the fire-emitted O_3 precursors, and the column O_3 660 enhancement is underestimated by 61%. The 1 km model better captures the maximum O3 enhancement (~3.47×10¹⁷ molecule cm⁻², or ~0.46×10¹⁷ molecule cm⁻² above the background 661 662 level, 34% lower than the 0.1 km LES model). NO₂ column shows similar trends as well: the 4 km 663 model shows a fire-induced NO₂ column enhancement only in the first ~20 km or so, while the 664 0.1 km model efficiently transports the fire-induced NO₂ column to > 40 km. The 1 km model also better captures the spatial variation of the NO₂ column in the downwind direction. Interestingly, 665 666 all three models are driven by the same external forcing, yet the plume in the 4 km model is much 667 wider than that in the 0.1 km LES model, possibly due to numerical diffusion (a known issue in 668 finite volume Eulerian grids). The 1 km model appears to predict a narrower plume width than 669 the 0.1 km LES model. This is mainly because of the alignment of the fire source. In the 0.1 km 670 LES model, in order to match the observed plume width (Figure 6 and Figure S4-S8), the fire 671 source is tuned into a rectangular shape of 0.3 km (X) \times 2.2 km (Y), while in the 1 km model the

fire source is just one 1 km × 1 km grid. The high horizontal resolution of the 0.1 km LES model allows a better representation of the shape and the orientation of the fire source as well. Overall, this exercise shows results that are consistent with those in Figure 14, that the numerical dilution leads to a decrease in O₃ formation potential, with a reduced efficiency of transporting smoke to the downwind direction. The 1 km model performs better than the 4 km model in terms of efficiently transporting the smoke and predicting the O₃ formation.

678

679 8. WHAT HORIZONTAL RESOLUTION SHOULD AIR QUALITY MODELS TARGET?

680 The effects of spatial resolution on the chemical evolution in biomass burning plumes or similar point-/area-sources have been discussed (Alvarado et al., 2009; Valin et al., 2011; van 681 Wees & van der Werf, 2019). In particular, Valin et al. (2011) pointed out that in order to capture 682 683 the NO_2 removal in the urban plumes, the model resolution has to be in the range of 4-12 km. 684 Most active wildland fires areas are smaller than the urban scale. According to the Fire INventory from NCAR (FINN) version 2 (Wiedinmyer et al., 2011), a widely used biomass burning inventory, 685 90% of wildfires in 2019 were smaller than 1 km² (fire size). Therefore, the impacts of small fires 686 687 on O₃ formation may be largely reduced (possibly entirely missing) in models with coarser spatial 688 resolutions, which is caused by numerical dilution, a bias purely caused by grid resolution.

To what degree small fires affect air quality on the regional scale remains unclear. Yu et al. (2016) examined the model grid resolution effect on NO_x and O₃ in the Southeast U.S., with horizontal resolution varying from $0.25^{\circ} \times 0.3125^{\circ}$ to $4^{\circ} \times 5^{\circ}$, and concluded that *"the good agreement of simulated and observed concentration variances implies that smaller-scale nonlinearities (urban and power plant plumes) are not important on the regional scale"* (Yu et al.,

694 2016). Although Yu et al. (2016) did not specifically target wildfire influence, from the modeling
695 point-of-view, wildland fire sources are conceptually similar to urban and power plant sources.
696 Based on the sensitivity test in Figure 15, we argue that impacts of small wildfires on the regional
697 scale may not be captured in their 0.25° × 0.3125° model (horizontal resolution: ~20 km).

698 To represent the smoke transport and O_3 formation associated with wildland fires, in 699 principle, the model grid resolution should be comparable to the active fire source. Although 700 plausible, it remains computationally challenging to perform simulations with LES schemes at 701 regional scale. Considering the computational cost associated with model resolution, we propose 702 that air quality forecast models should target ~1 km horizontal resolution in the near future. We 703 show in this work that a 1 km model can reasonably well capture the transport of NO_x and O_3 704 formation associated with a fire area that is ~60% of the grid size. Based again on FINN version 2, 705 in 2019, approximately 17% of wildfires in the U.S. are larger than 0.6 km², and these top 17% 706 wildfires contribute 62% total fire-NO_x emission, 60% total fire-CO emission, and 61% total non-707 methane VOC emissions from fires nationwide. Therefore, an air quality forecast model with ~1 708 km horizontal resolution can greatly improve our ability to predict wildfire impacts on air quality, 709 which could potentially affect nearly half of the U.S. population (Figure S11). Additional benefit 710 of 1 km horizontal resolution is that the impacts of urban emissions will also be better resolved 711 (Valin et al., 2011).

712

713 9. CONCLUSIONS AND REMARKS

In this work, a high-resolution turbulence-resolving model, WRF-LES, is coupled to
chemistry to study the chemical evolution in a wildland fire plume. The results are evaluated

using airborne measurements collected from the Williams Flats Fire during the FIREX-AQ field
campaign. The model reveals remarkable chemical heterogeneity that is supported by
observations. Major findings are summarized as follows:

1. The strong heat released from the fire source drives rapid plume rise in the very early
stage, which also leads to rapid dilution with background air. Dilution and physical mixing greatly
affects chemistry inside the plume.

2. O_3 shows considerable heterogeneity inside the plume: suppressed at the center (due to NO reaction) but may be enhanced at the edges. To our knowledge, our model is the first to capture the observed cross-transect O_3 variations in optically thick biomass burning plumes.

3. PAN and maleic anhydride (from the oxidation of furan, a distinct class of compounds emitted from fires) are also enhanced at the edges of the plume in the early stage, implying faster OH oxidation at the edges. The OH enhancement at the plume edges is largely diminished in the later stage, mainly because the suppression of photochemistry inside the plume becomes weaker due to dilution.

4. Primary OH production is dominated by HONO photolysis, and HO_x cycling extends OH radicals deeper into the plume. Although OH oxidation is the major sink of VOCs in daytime plumes, we show that dark chemistry driven by O_3 and NO_3 oxidation may proceed at the same time but mostly in the lower part of (or below) thick plumes.

5. Although this modeling work cannot provide direct evidence of secondary HONO
production in the studied plume, we find that observed HONO variability can be best explained
by including a secondary HONO production from NO₂ heterogeneous reaction on aerosols.

737 We acknowledge that the model configuration is highly idealized, and much of the 738 observations remain unexplained by the model. Nevertheless, there are several key take-away 739 messages worth mentioning. Firstly, Lagrangian-type models that explore the chemical evolution 740 in dense fire plumes must specify the plume region being modeled, since it is the edges that 741 largely drive the photochemistry in thick fire plumes. Secondly, dilution and mixing within the 742 plume are important, which are not constant and may greatly affect how fast chemistry proceeds 743 inside the plume. Thirdly, we found that HONO and the photons that can trigger HONO photolysis 744 (represented by j_{HONO}) are always segregated in the modeled plume, i.e., HONO is always 745 depleted in the edges and the upper part, while j_{HONO} is exclusively elevated in the edges and 746 upper most part of the optically dense smoke (Figure 5). This has implications for satellite HONO 747 retrievals. Recently Theys et al. (2020) reported the detection of HONO in biomass burning 748 plumes from TROPOspheric Monitoring Instrument (TROPOMI), showcasing the remarkable 749 capability of retrieving reactive trace gases such as HONO from this space-borne instrument. High 750 resolution models with chemistry such as the LES-chemistry model used in this work can better 751 resolve the plume structure and the distributions of trace gases and aerosols, therefore may be 752 used to test remote sensing retrievals for optically dense plumes.

Lastly, we showed that model spatial resolution may shift the chemical regime and lead to reduced O_3 formation potential. For the Williams Flats Fire studied in this work, a model with ~1 km horizontal resolution may decently capture the fire-induced O_3 and NO_2 enhancement in the downwind direction, but a model with ~4 km horizontal resolution may be insufficient to capture the impacts on these short-lived pollutants. We thus propose that regional or national air quality models in the U.S. should target ~1 km horizontal resolution in the near future, which

- may greatly improve our capability to predict the wildfire impacts on air quality for nearly half of
- the U.S. population.
- 761

762 ACKNOWLEDGEMENTS AND DATA AVAILABILITY

763 S.-Y. W. is partially supported by the FIREX-AQ project (80NSSC18K0681). S.R.H. and K.U. 764 and partially supported by the NASA FIREX-AQ project (80NSSC18K0638). The computing and 765 data storage resources, including the Cheyenne supercomputer (doi:10.5065/D6RX99HX), were 766 provided by the Computational and Information Systems Laboratory (CISL) at NCAR. NCAR is a 767 major facility sponsored by the NSF under Cooperative Agreement No. 1852977. We thank the 768 FIREX-AQ Science Team for their exceptional professionalism in support of this mission. We thank 769 Glenn Wolfe, Thomas Hanisco (NASA Goddard), Joshua P. Schwarz, Joseph M. Katich (NOAA/CSL), 770 Jose-Luis Jimenez, Pedro Campuzano-Jost, Douglas Day, Hongyu Guo, Demetrios Pagonis, Mindy 771 Schueneman (CU Boulder), and Alan Fried (CU Boulder) for their efforts in collecting the 772 measurements. Stuart McKeen, Rebecca Schwantes (CIRES/NOAA CSL), Brian McDonald, and 773 Gregory Frost (NOAA CSL) are acknowledged for helpful discussion. We thank Jimy Dudhia and 774 Mary Barth (NCAR) for the support regarding the LES modeling. The FIREX-AQ campaign data is 775 available from NASA archived and Langley Research Center (https://www-776 air.larc.nasa.gov/missions/firex-aq/). The WRF model is available from github 777 (https://github.com/wrf-model/WRF).

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779 **REFERENCES**

- Abatzoglou, J. T., & Williams, A. P. (2016). Impact of anthropogenic climate change on wildfire
 across western US forests. *Proceedings of the National Academy of Sciences*, 113(42),
 11770–11775. https://doi.org/10.1073/pnas.1607171113
- Agee, E., & Gluhovsky, A. (1999). LES Model Sensitivities to Domains, Grids, and Large-Eddy
 Timescales. *Journal of Atmospheric Sciences*, *56*(4), 599–604.
- 785 https://doi.org/10.1175/1520-0469(1999)056<0599:LMSTDG>2.0.CO;2
- Alvarado, M. J., Logan, J. A., Mao, J., Apel, E., Riemer, D., Blake, D., et al. (2010). Nitrogen oxides
 and PAN in plumes from boreal fires during ARCTAS-B and their impact on ozone: an
 integrated analysis of aircraft and satellite observations. *Atmospheric Chemistry and Physics*, 10(20), 9739–9760. https://doi.org/10.5194/acp-10-9739-2010
- Alvarado, M. J., Lonsdale, C. R., Yokelson, R. J., Akagi, S. K., Coe, H., Craven, J. S., et al. (2015).
 Investigating the links between ozone and organic aerosol chemistry in a biomass
 burning plume from a prescribed fire in California chaparral. *Atmospheric Chemistry and*
- 793 *Physics*, *15*(12), 6667–6688. https://doi.org/10.5194/acp-15-6667-2015
- Alvarado, Matthew James, & Prinn, R. G. (2009). Formation of ozone and growth of aerosols in
 young smoke plumes from biomass burning: 1. Lagrangian parcel studies. *Journal of*
- 796 *Geophysical Research: Atmospheres, 114*(D9). https://doi.org/10.1029/2008JD011144

797 Alvarado, Matthew James, Wang, C., & Prinn, R. G. (2009). Formation of ozone and growth of 798 aerosols in young smoke plumes from biomass burning: 2. Three-dimensional Eulerian 799 studies. Journal of Geophysical Research: Atmospheres, 114(D9). 800 https://doi.org/10.1029/2008JD011186 801 Ammann, M., Cox, R. A., Crowley, J. N., Jenkin, M. E., Mellouki, A., Rossi, M. J., et al. (2013). 802 Evaluated kinetic and photochemical data for atmospheric chemistry: Volume VI -803 heterogeneous reactions with liquid substrates. Atmospheric Chemistry and Physics, 804 13(16), 8045–8228. https://doi.org/10.5194/acp-13-8045-2013 805 Andreae, M. O. (2019). Emission of trace gases and aerosols from biomass burning – an 806 updated assessment. Atmospheric Chemistry and Physics, 19(13), 8523-8546. 807 https://doi.org/10.5194/acp-19-8523-2019 Brock, C. A., Washenfelder, R. A., Trainer, M., Ryerson, T. B., Wilson, J. C., Reeves, J. M., et al. 808 809 (2002). Particle growth in the plumes of coal-fired power plants. Journal of Geophysical 810 Research: Atmospheres, 107(D12), AAC 9-1-AAC 9-14. 811 https://doi.org/10.1029/2001JD001062 812 Coen, J. (2013). Modeling wildland fires : A description of the Coupled Atmosphere-Wildland 813 Fire Environment model (CAWFE). https://doi.org/10.5065/D6K64G2G 814 Coggon, M. M., Lim, C. Y., Koss, A. R., Sekimoto, K., Yuan, B., Gilman, J. B., et al. (2019). OH 815 chemistry of non-methane organic gases (NMOGs) emitted from laboratory and 816 ambient biomass burning smoke: evaluating the influence of furans and oxygenated 817 aromatics on ozone and secondary NMOG formation. Atmospheric Chemistry and 818 Physics, 19(23), 14875–14899. https://doi.org/10.5194/acp-19-14875-2019 819 Doerr, S. H., & Santín, C. (2016). Global trends in wildfire and its impacts: perceptions versus 820 realities in a changing world. Philosophical Transactions of the Royal Society B: Biological 821 Sciences, 371(1696), 20150345. https://doi.org/10.1098/rstb.2015.0345 822 Fischer, EV, Jacob, D. J., Yantosca, R. M., Sulprizio, M. P., Millet, D. B., Mao, J., et al. (2014). 823 Atmospheric peroxyacetyl nitrate (PAN): a global budget and source attribution. Atmos. 824 Chem. Phys., 14(5), 2679–2698. https://doi.org/10.5194/acp-14-2679-2014 825 Fischer, Emily V., Zhu, L., Payne, V. H., Worden, J. R., Jiang, Z., Kulawik, S. S., et al. (2018). Using 826 TES retrievals to investigate PAN in North American biomass burning plumes. 827 Atmospheric Chemistry and Physics, 18(8), 5639–5653. https://doi.org/10.5194/acp-18-828 5639-2018 829 Flannigan, M., Cantin, A. S., de Groot, W. J., Wotton, M., Newbery, A., & Gowman, L. M. (2013). 830 Global wildland fire season severity in the 21st century. Forest Ecology and 831 Management, 294, 54–61. https://doi.org/10.1016/j.foreco.2012.10.022 832 Hodshire, A. L., Bian, Q., Ramnarine, E., Lonsdale, C. R., Alvarado, M. J., Kreidenweis, S. M., et al. 833 (2019). More Than Emissions and Chemistry: Fire Size, Dilution, and Background Aerosol Also Greatly Influence Near-Field Biomass Burning Aerosol Aging. Journal of Geophysical 834 835 Research: Atmospheres, 124(10), 5589–5611. https://doi.org/10.1029/2018JD029674 836 Hodshire, Anna L., Akherati, A., Alvarado, M. J., Brown-Steiner, B., Jathar, S. H., Jimenez, J. L., et 837 al. (2019). Aging Effects on Biomass Burning Aerosol Mass and Composition: A Critical 838 Review of Field and Laboratory Studies. Environmental Science & Technology, 53(17), 839 10007–10022. https://doi.org/10.1021/acs.est.9b02588

840 Hong, S.-Y., Noh, Y., & Dudhia, J. (2006). A New Vertical Diffusion Package with an Explicit 841 Treatment of Entrainment Processes. Monthly Weather Review, 134(9), 2318–2341. 842 https://doi.org/10.1175/MWR3199.1 Jaffe, D. A., Wigder, N., Downey, N., Pfister, G., Boynard, A., & Reid, S. B. (2013). Impact of 843 844 Wildfires on Ozone Exceptional Events in the Western U.S. Environmental Science & 845 Technology, 47(19), 11065–11072. https://doi.org/10.1021/es402164f 846 Jaffe, D. A., O'Neill, S. M., Larkin, N. K., Holder, A. L., Peterson, D. L., Halofsky, J. E., & Rappold, 847 A. G. (2020). Wildfire and prescribed burning impacts on air quality in the United States. 848 Journal of the Air & Waste Management Association, 70(6), 583–615. 849 https://doi.org/10.1080/10962247.2020.1749731 850 Knote, C., Hodzic, A., Jimenez, J. L., Volkamer, R., Orlando, J. J., Baidar, S., et al. (2014). 851 Simulation of semi-explicit mechanisms of SOA formation from glyoxal in aerosol in a 3-852 D model. Atmospheric Chemistry and Physics, 14(12), 6213–6239. 853 Kodros, J. K., Papanastasiou, D. K., Paglione, M., Masiol, M., Squizzato, S., Florou, K., et al. 854 (2020). Rapid dark aging of biomass burning as an overlooked source of oxidized organic 855 aerosol. Proceedings of the National Academy of Sciences. 856 https://doi.org/10.1073/pnas.2010365117 857 Konovalov, I. B., Beekmann, M., Golovushkin, N. A., & Andreae, M. O. (2019). Nonlinear 858 behavior of organic aerosol in biomass burning plumes: a microphysical model analysis. 859 Atmospheric Chemistry and Physics, 19(19), 12091–12119. https://doi.org/10.5194/acp-19-12091-2019 860 861 Koss, A. R., Sekimoto, K., Gilman, J. B., Selimovic, V., Coggon, M. M., Zarzana, K. J., et al. (2018). 862 Non-methane organic gas emissions from biomass burning: identification, 863 quantification, and emission factors from PTR-ToF during the FIREX 2016 laboratory 864 experiment. Atmospheric Chemistry and Physics, 18(5), 3299–3319. 865 https://doi.org/10.5194/acp-18-3299-2018 866 Li, F., Zhang, X., Kondragunta, S., Schmidt, C. C., & Holmes, C. D. (2020). A preliminary 867 evaluation of GOES-16 active fire product using Landsat-8 and VIIRS active fire data, and 868 ground-based prescribed fire records. Remote Sensing of Environment, 237, 111600. 869 https://doi.org/10.1016/j.rse.2019.111600 870 Liu, X., Huey, L. G., Yokelson, R. J., Selimovic, V., Simpson, I. J., Müller, M., et al. (2017). Airborne 871 measurements of western U.S. wildfire emissions: Comparison with prescribed burning and air quality implications. Journal of Geophysical Research: Atmospheres, 122(11), 872 873 6108–6129. https://doi.org/10.1002/2016JD026315 874 Lu, X., Zhang, L., Yue, X., Zhang, J., Jaffe, D. A., Stohl, A., et al. (2016). Wildfire influences on the 875 variability and trend of summer surface ozone in the mountainous western United 876 States. Atmospheric Chemistry and Physics, 16(22), 14687–14702. https://doi.org/10.5194/acp-16-14687-2016 877 878 Mandel, J., Beezley, J. D., & Kochanski, A. K. (2011). Coupled atmosphere-wildland fire modeling 879 with WRF 3.3 and SFIRE 2011. Geoscientific Model Development, 4(3), 591–610. 880 https://doi.org/10.5194/gmd-4-591-2011 881 Martinuzzi, S., Stewart, S. I., Helmers, D. P., Mockrin, M. H., Hammer, R. B., & Radeloff, V. C. 882 (2015). The 2010 wildland-urban interface of the conterminous United States. Research 883 Map NRS-8. Newtown Square, PA: U.S. Department of Agriculture, Forest Service,

884 Northern Research Station. 124 p. [Includes Pull-out Map]., 8, 1–124. 885 https://doi.org/10.2737/NRS-RMAP-8 886 Mason, S. A., Trentmann, J., Winterrath, T., Yokelson, R. J., Christian, T. J., Carlson, L. J., et al. 887 (2006). Intercomparison of Two Box Models of the Chemical Evolution in Biomass-888 Burning Smoke Plumes. Journal of Atmospheric Chemistry, 55(3), 273–297. 889 https://doi.org/10.1007/s10874-006-9039-5 890 McClure, C. D., & Jaffe, D. A. (2018). US particulate matter air quality improves except in 891 wildfire-prone areas. Proceedings of the National Academy of Sciences, 115(31), 7901-892 7906. https://doi.org/10.1073/pnas.1804353115 893 Moeng, C.-H., Dudhia, J., Klemp, J., & Sullivan, P. (2007). Examining Two-Way Grid Nesting for 894 Large Eddy Simulation of the PBL Using the WRF Model. Monthly Weather Review, 135(6), 2295–2311. https://doi.org/10.1175/MWR3406.1 895 896 Palm, B. B., Peng, Q., Fredrickson, C. D., Lee, B. H., Garofalo, L. A., Pothier, M. A., et al. (2020). 897 Quantification of organic aerosol and brown carbon evolution in fresh wildfire plumes. 898 Proceedings of the National Academy of Sciences, 117(47), 29469–29477. 899 https://doi.org/10.1073/pnas.2012218117 900 Parrington, M., Palmer, P. I., Lewis, A. C., Lee, J. D., Rickard, A. R., Di Carlo, P., et al. (2013). 901 Ozone photochemistry in boreal biomass burning plumes. Atmospheric Chemistry and 902 Physics, 13(15), 7321–7341. https://doi.org/10.5194/acp-13-7321-2013 903 Peng, Q., Palm, B. B., Melander, K. E., Lee, B. H., Hall, S. R., Ullmann, K., et al. (2020). HONO 904 Emissions from Western U.S. Wildfires Provide Dominant Radical Source in Fresh 905 Wildfire Smoke. Environmental Science & Technology, 54(10), 5954–5963. 906 https://doi.org/10.1021/acs.est.0c00126 907 Reid Colleen E., Brauer Michael, Johnston Fay H., Jerrett Michael, Balmes John R., & Elliott 908 Catherine T. (2016). Critical Review of Health Impacts of Wildfire Smoke Exposure. 909 Environmental Health Perspectives, 124(9), 1334–1343. 910 https://doi.org/10.1289/ehp.1409277 911 Roberts, J. M., Stockwell, C. E., Yokelson, R. J., de Gouw, J., Liu, Y., Selimovic, V., et al. (2020). 912 The nitrogen budget of laboratory-simulated western US wildfires during the FIREX 2016 913 Fire Lab study. Atmospheric Chemistry and Physics, 20(14), 8807–8826. 914 https://doi.org/10.5194/acp-20-8807-2020 915 Roode, S. R. de, Duynkerke, P. G., & Jonker, H. J. J. (2004). Large-Eddy Simulation: How Large is 916 Large Enough? Journal of Atmospheric Sciences, 61(4), 403–421. 917 https://doi.org/10.1175/1520-0469(2004)061<0403:LSHLIL>2.0.CO;2 918 Schwarz, J. P., Gao, R. S., Spackman, J. R., Watts, L. A., Thomson, D. S., Fahey, D. W., et al. 919 (2008). Measurement of the mixing state, mass, and optical size of individual black 920 carbon particles in urban and biomass burning emissions. Geophysical Research Letters, 921 35(13). https://doi.org/10.1029/2008GL033968 922 Singh, H. B., Cai, C., Kaduwela, A., Weinheimer, A., & Wisthaler, A. (2012). Interactions of fire 923 emissions and urban pollution over California: Ozone formation and air quality 924 simulations. Atmospheric Environment, 56, 45–51. 925 https://doi.org/10.1016/j.atmosenv.2012.03.046

926	Theys, N., Volkamer, R., Müller, JF., Zarzana, K. J., Kille, N., Clarisse, L., et al. (2020). Global
927	nitrous acid emissions and levels of regional oxidants enhanced by wildfires. Nature
928	<i>Geoscience</i> , 13(10), 681–686. https://doi.org/10.1038/s41561-020-0637-7
929	Tie, X., Madronich, S., Walters, S., Zhang, R., Rasch, P., & Collins, W. (2003). Effect of clouds on
930	photolysis and oxidants in the troposphere. Journal of Geophysical Research:
931	Atmospheres, 108(D20). https://doi.org/10.1029/2003JD003659
932	Trentmann, J., Andreae, M. O., Graf, HF., Hobbs, P. V., Ottmar, R. D., & Trautmann, T. (2002).
933	Simulation of a biomass-burning plume: Comparison of model results with observations.
934	Journal of Geophysical Research: Atmospheres. 107(D2), AAC 5-1-AAC 5-15.
935	https://doi.org/10.1029/2001JD000410
936	Trentmann, Jörg, Andreae, M. O., & Graf, HF. (2003). Chemical processes in a young biomass-
937	burning plume. Journal of Geophysical Research: Atmospheres, 108(D22).
938	https://doi.org/10.1029/2003JD003732
939	Trentmann, Jörg, Früh, B., Boucher, O., Trautmann, T., & Andreae, M. O. (2003). Three-
940	dimensional solar radiation effects on the actinic flux field in a biomass-burning plume.
941	Journal of Geophysical Research: Atmospheres, 108(D17).
942	https://doi.org/10.1029/2003JD003422
943	Trentmann, Jörg, Yokelson, R. J., Hobbs, P. V., Winterrath, T., Christian, T. J., Andreae, M. O., &
944	Mason, S. A. (2005). An analysis of the chemical processes in the smoke plume from a
945	savanna fire. Journal of Geophysical Research: Atmospheres, 110(D12).
946	https://doi.org/10.1029/2004JD005628
947	Valin, L. C., Russell, A. R., Hudman, R. C., & Cohen, R. C. (2011). Effects of model resolution on
948	the interpretation of satellite NO ₂ observations. Atmospheric Chemistry and Physics.
949	11(22), 11647–11655, https://doi.org/10.5194/acp-11-11647-2011
950	van Wees, D., & van der Werf, G. R. (2019). Modelling biomass burning emissions and the effect
951	of spatial resolution: a case study for Africa based on the Global Fire Emissions Database
952	(GFED). Geoscientific Model Development, 12(11), 4681–4703.
953	https://doi.org/10.5194/gmd-12-4681-2019
954	Westerling, A. L., Hidalgo, H. G., Cayan, D. R., & Swetnam, T. W. (2006). Warming and Earlier
955	Spring Increase Western U.S. Forest Wildfire Activity. Science, 313(5789), 940–943.
956	https://doi.org/10.1126/science.1128834
957	Wiedinmyer, C., Akagi, S. K., Yokelson, R. J., Emmons, L. K., Al-Saadi, J. A., Orlando, J. J., & Soja,
958	A. J. (2011). The Fire INventory from NCAR (FINN): a high resolution global model to
959	estimate the emissions from open burning. <i>Geosci. Model Dev.</i> , 4(3), 625–641.
960	https://doi.org/10.5194/gmd-4-625-2011
961	Yokelson, R. J., Christian, T. J., Karl, T. G., & Guenther, A. (2008). The tropical forest and fire
962	emissions experiment: laboratory fire measurements and synthesis of campaign data.
963	Atmospheric Chemistry and Physics, 8(13), 3509–3527. https://doi.org/10.5194/acp-8-
964	3509-2008
965	Yu, K., Jacob, D. J., Fisher, J. A., Kim, P. S., Marais, E. A., Miller, C. C., et al. (2016). Sensitivity to
966	grid resolution in the ability of a chemical transport model to simulate observed oxidant
967	chemistry under high-isoprene conditions. Atmospheric Chemistry and Physics, 16(7).
968	4369–4378. https://doi.org/10.5194/acp-16-4369-2016
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AGUPUBLICATIONS

1 2 Journal of Geophysical Research 3 Supporting Information for Chemical Tomography in a Fresh Wildland Fire Plume: a Large Eddy 4 Simulation (LES) Study 5 Siyuan Wang,^{1,2,3,*} Matthew M. Coggon,^{1,2} Georgios I. Gkatzelis,^{1,2,†} Carsten Warneke,^{1,2} Ilann 6 Bourgeois,^{1,2} Thomas Ryerson,² Jeff Peischl,^{1,2} Patrick R. Veres,² J. Andrew Neuman,^{1,2} Johnathan Hair,⁴ 7 Taylor Shingler,⁴ Marta Fenn,⁴ Glenn Diskin,⁴ L. Greg Huey,⁵ Young Ro Lee,⁵ Eric C. Apel,³ Rebecca S. 8 Hornbrook,³ Alan J. Hills,³ Samuel R. Hall,³ Kirk Ullmann,³ Megan M. Bela,^{1,2} Michael K. Trainer,^{1,2} Rajesh 9 Kumar,⁶ John J. Orlando,³ Frank M. Flocke,³ Louisa K. Emmons,³ 10 11 ¹ Cooperative Institute for Research in Environmental Sciences (CIRES), University of Colorado, Boulder, 12 Colorado 13 ² National Oceanic and Atmospheric Administration (NOAA), Chemical Sciences Laboratory (CSL), Boulder, Colorado 14 ³ National Center for Atmospheric Research (NCAR), Atmospheric Chemistry Observations and Modeling 15 16 Laboratory (ACOM), Boulder, Colorado 17 ⁴ National Aeronautics and Space Administration, Langley Research Center, Hampton, Virginia 18 ⁵ School of Earth and Atmospheric Sciences, Georgia Institute of Technology, Atlanta, Georgia 19 ⁶ National Center for Atmospheric Research (NCAR), Research Applications Laboratory (RAL), Boulder, 20 Colorado 21 22 * Corresponding author: Siyuan Wang (siyuan.wang@noaa.gov) 23 Cooperative Institute for Research in Environmental Sciences (CIRES), University of Colorado, Boulder, 24 Boulder, Colorado 25 NOAA, Chemical Sciences Laboratory (CSL) 325 Broadway, Boulder, CO 26

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34

35 Section S1. WRF-LES Technical Updates.

The publicly released WRF v4.0 is used in this work. Several new updates have been released in newer versions of WRF. Two of them are relevant for this work: (1) A bug fix that improves the potential temperature tendency, which is recently implemented in WRF v4.2.2 (https://github.com/wrf-model/WRF/pull/1259/files#diff-

40 d6fd08d956dbe380e8c1094cfc668bd4); and (2) another bug fix that improves the momentum

flux and sub-grid scale stress, which is implemented in WRF v4.2.1 (https://github.com/wrf-

42 model/WRF/pull/1214/files/323fc93b19465163ef6529e79eec5a82292e5eb1). These two new

43 implementations are merged into the WRF model version 4.0 used in this work.

This idealized WRF-LES model is evaluated against another widely used LES model, 44 45 NCAR-LES (Patton et al., 2005). In this comparision, both WRF-LES and NCAR-LES are configured in the same way (i.e., same initial conditions, driven by the same surface heat and moisture 46 fluxes). It drew our attention that the friction velocity (U_*) calculated using this WRF-LES (with 47 48 the aforementioned bug fixes) is roughly a factor of two lower than that calculated using NCAR-49 LES. The reason may be that, in the default WRF, the friction velocity is calculated using the bulk Richardson number, which may not be properly updated in idealized WRF-LES when the 50 51 surface heat flux is prescribed (personal communication: Jimy Dudhia, National Center for Atmospheric Research). Therefore, a new friction velocity scheme is developed in this work for 52 the idealized WRF-LES to solve U* directly: 53

54
$$U_* = \frac{\kappa U}{\ln(\frac{z}{z_0}) - \Psi_m(\frac{z}{L})}$$
 Equation S1
55 $L = -\frac{\rho C_p \theta U_*^3}{kgH}$ Equation S2

Where κ is the von Kármán constant, U is the surface horizontal wind speed (m s⁻¹), z_0 is the 56 roughness length scale (m), Ψ_m is the stability correction function for momentum (in the WRF 57 surface layer scheme), L is the Obukhov length (m), ρ is the air density (kg m⁻³), C_{ρ} is the heat 58 capacity (1005 J kg⁻¹ K⁻¹), ϑ is the potential temperature (K), H is the surface heat flux (W m⁻²), 59 and g is the acceleration due to gravity (9.8 m s⁻²), Equations S1 and S2 are solved implicitly 60 using the Newton iteration method. Using this new scheme, the friction velocity calculated in 61 62 this idealized WRF-LES driven by surface heat fluxes is greatly improved and in very good agreement with the NCAR-LES. 63

64 Other major physical and dynamical settings of the WRF-LES model are summarized as 65 follows: microphysics option: Morrison 2-moment scheme (mp_physics = 10);

- 66 longwave/shortwave radiation option: Rapid Radiative Transfer Model for GCMs (RRTMG;
- ra lw physics/ra sw physics = 4); turbulence and mixing option: evaluates mixing terms in
- 68 physical space (stress form; diff_opt = 2); eddy coefficient option: 1.5 order TKE closure
- 69 (km_opt = 2).
- 70

71 Section S2. Mesoscale WRF-Chem

- 72 A mesoscale WRF-Chem simulation (12 km horizontal resolution) was used to generate
- 73 the initial conditions for the WRF-LES model (Figures S1 and S2). Some of the major settings of
- this 12 km WRF-Chem simulation are summarized as follows: microphysics option: Morrison 2-
- 75 moment scheme (mp_physics = 10); longwave/shortwave radiation option: Rapid Radiative
- 76 Transfer Model for GCMs (RRTMG; ra_lw_physics/ra_sw_physics = 4); turbulence and mixing
- option: evaluates mixing terms in physical space (stress form; diff_opt = 2); eddy coefficient
- option: horizontal Smagorinsky first order closure (km_opt = 4); surface-layer option: MYNN
- 79 surface layer (sf_sfclay_physics = 5); land-surface option: Noah Land-Surface Model
- 80 (sf_surface_physics = 2); boundary-layer option: MYNN 2.5 level TKE scheme (bl_pbl_physics =
- 5); cumulus option: Grell-Devenyi ensemble scheme (cu_physics = 3); chemistry option:
- 82 NOAA/ESRL RACM chemical mechanism (chem_opt = 108); photolysis option: Madronich
- 83 photolysis (TUV; phot_opt = 1).

- **Table S1.** The highly simplified chemical mechanism used in the WRF-LES model. This
- 86 mechanism is condensed based on T1 MOZART in the WRF package (Knote et al., 2014).

Reactions	Rate coefficients
O3 = 2 OH	O3_hv_H2O(T, [M], [H2O], j_O1D), note 1
NO2 + hy = NO + O3	FTUV
NO2 + aerosols = 0.5 HNO2	$v = 5 \times 10^{-4}$. note 2
HNO2 + hv = OH + NO	FTUV
H2O2 + hv = 2 OH	FTUV
HCHO + hv = 2 HO2 + CO	FTUV
HCHO + hv = CO	FTUV
OH + HO2 = H2O	T1 MOZART
OH + O3 = HO2	T1 MOZART
HO2 + HO2 = H2O2	T1 MOZART
HO2 + O3 = OH	T1 MOZART
NO + O3 = NO2	T1 MOZART
NO2 + OH =	T1 MOZART
NO2 + O3 = NO3	T1 MOZART
NO + HO2 = NO2 + OH	T1 MOZART
NO2 + NO3 = N2O5	T1 MOZART
NO + NO3 = NO2 + NO2	T1 MOZART
HO2 + NO3 = OH + NO2	T1 MOZART
N2O5 = NO2 + NO3	T1 MOZART
N2O5 =	T1 MOZART
NO3 + hv = 0.11 NO + 0.89 NO2 + 0.89 O3	FTUV
CH3O2 + NO = HCHO + 0.99 NO2 + HO2	T1 MOZART
CH3O2 + HO2 =	T1 MOZART
CO + OH = HO2	T1 MOZART
CH4 + OH = CH3O2	T1 MOZART
HCHO + OH = HO2 + CO	T1 MOZART
C3H6 + OH = PO2	T1 MOZART
PO2 + NO = HO2 + HCHO + ALDS + NO2	T1 MOZART
PO2 + HO2 = POOH	T1 MOZART
POOH + OH = 0.5 PO2 + 0.5 OH + 0.5 ALDS	T1 MOZART
C3H6 + O3 = 0.5 HCHO + 0.5 ALDS + 0.28 CH3O2 + 0.28	T1 MOZART
HO2 + 0.36 OH	
C3H6 + NO3 = NO2 + ALDS	T1 MOZART
ISOP + OH = ISOPO2	T1 MOZART
ISOP + 03 = 0.6 MVKMACR + 0.27 OH + 0.06 HO2 + 0.6	T1 MOZART, note 3
HCHU + 0.3 CU + 0.1 U3 + 0.2 RCU3	T1 MOZADT
SOPO2 + NO2 - ISOPNO2 + 0.02 NO2 + 0.55	
MVKMACR + 0.37 ALDS + 0.55 HCHO + HO2	T1 MOZART
ISOPO2 + HO2 = ISOPOOH	T1 MOZART

ISOPO2 + CH3O2 = HO2 + 1.2 HCHO + 0.45 MVKMACR +	
0.3 ALDS	TI WOZARI
ISOPOOH + OH = 0.5 XO2 + 0.5 ISOPO2	T1 MOZART
ISOPOOH + hv = 0.69 MVKMACR + 0.69 HCHO + HO2	FTUV
MACRO2 + NO = NO2 + 0.47 HO2 + 0.25 HCHO + ALDS +	T1 MOZART
0.53 KCO3 + 0.22 CO MACRO2 + HO2 -	
MACRO2 + CH3O2 = 0.73 HO2 + 0.88 HCHO + 0.11 CO + 0.000 HCHO + 0.0000 HCHO	
0.73 ALDS + 0.26 RCO3	T1 MOZART
MVKMACR + hv = 0.165 OH + 0.335 HO2 + 0.65 RCO3 +	
0.335 HCHO + 0.685 CO + 0.15 CH3O2	T1 MOZART
MVKMACR + OH = 0.75 MACRO2 + 0.25 RCO3	T1 MOZART
MVKMACR + O3 = 0.75 HCHO + 0.895 ALDS + 0.148 OH	
+ 0.2 O3 + 0.168 HO2 + 0.125 CO	TI WOZARI
ONITR + hv = HO2 + CO + NO2 + HCHO	T1 MOZART
ONITR + OH = ALDS + HO2 + 0.4 NO2	T1 MOZART
ALDS + OH = RCO3	T1 MOZART, note 4
ALDS + hv = CH3O2 + CO + HO2	j-CH3CHO in FTUV
XO2 + NO = NO2 + 1.5 HO2 + CO + 0.75 ALDS	T1 MOZART
XO2 + HO2 =	T1 MOZART
XO2 + CH3O2 = HO2 + 0.7 HCHO + 0.4 CO + 0.3 ALDS	T1 MOZART
RCO3 + NO = NO2 + HCHO	T1 MOZART
RCO3 + HO2 = 0.25 O3	T1 MOZART
RCO3 + CH3O2 = 2 HCHO + HO2	T1 MOZART
RCO3 + NO2 = PAN	T1 MOZART
PAN = RCO3 + NO2	T1 MOZART
PAN + hv = 0.8 RCO3 + 0.8 NO2 + 0.2 CH3O2 + 0.2 NO3	j-PAN in FTUV
ISOPNO3 + NO = 1.206 NO2 + 0.794 HO2 + 0.072 HCHO	Τ1 ΜΟΖΔΡΤ
+ 0.206 MVKMACR + 0.794 ONITR	
ISOPNO3 + HO2 = 0.206 NO2 + 0.206 OH + 0.206 HCHO	T1 MOZART
+ 0.206 MVKMACR + 0.794 ONTR	74 1407407
BENZENE + OH = 0.83 HO2 + 0.18 CO + 0.35 BENZO2	
BENZO2 + HO2 = BENZOOH	
BENZO2 + NO = NO2 + 0.5 ALDS + HO2	
	11 MOZARI
10LUENE + OH = 0.65 10L02 + 0.05 RC03 + 0.38 H02 + 0.15 CO	T1 MOZART
TOLO2 + HO2 = TOLOOH	T1 MOZART
TOLO2 + NO = NO2 + 0.6 ALDS + HO2	T1 MOZART
TOLOOH + OH = TOLO2	T1 MOZART
FURAN + OH = 0.39 DIAL + 0.49 HO2 + 0.51 FURANO2	Coggon et al. 2019
DIAI + OH = DIAI O2	Coggon et al., 2019
DIALO2 + NO = NO2 + 0.49 MALANHY + 0.96 HO2 + 0.04	
CH3O2	Coggon et al., 2019
DIALO2 + NO2 =	Coggon et al., 2019
DIAL + hv = 0.74 DIALO2 + 0.74 HO2 + 0.34 CO	0.098*j _{NO2} , Coggon et al., 2019

FURANO2 + NO = NO2 + FURANONE + 0.81 HO2 + 0.19	Coggon et al., 2019
FURANONE + OH = 0.69 MALANHY + 0.66 HO2 + 0.34	
CH3O2 + 0.43 CO	Coggon et al., 2019
FURAN + O3 =	Coggon et al., 2019
FURAN + NO3 =	Coggon et al., 2019

87

88 Note 1: O3_hv_H2O(T, [M], [H2O], j_O1D) = 1.63E-10*EXP(60/T)*[H2O]* j_O1D / (1.63E-10*EXP(60/T)*

89 [H2O] + 2.15E-11*EXP(110/T)*0.79*[M] + 3.30E-11*EXP(55/T)*0.21*[M]), where T is temperature (K),

and [M] and [H2O] are air and water vapor concentrations (molecules cm⁻³), respectively. This approach
 assumes steady-state for O¹D.

92 Note 2: the reactive uptake coefficient of NO2 is estimated from based on the literature values

93 (Ammann et al., 2013). Aerosol surface area in this work considers only BC and OC.

94 Note 3: MVKMACR is the sum of MVK and MACR.

95 Note 4: ALDS is a lumped species for aldehydes including acetaldehyde, glycolaldehyde, hydroxyacetone,

96 as well as lumped unsaturated hydroxycarbonyl from T1 MOZART. Of all these lumped species,

97 acetaldehyde is the most dominant one in fire plumes, therefore the kinetics of acetaldehyde (e.g.,

98 photolysis frequency, OH rate coeffcieint) are used for the removal of ALDS.

100	Table S2. Fire emissions used in the WRF-LES model. These emissions a	are adjusted until
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101 reasonable agreements are achieved when compared to airborne measurements (Figure 3).

Species	Fire emission	Unit
NO	1097	ppb m s ⁻¹
CO	77780	ppb m s⁻¹
ISOP	238	ppb m s⁻¹
MVKMACR	153	ppb m s⁻¹
EC	297	µg m ⁻² s ⁻¹
OC	19649	µg m ⁻² s ⁻¹
C3H6	802	ppb m s⁻¹
ALDS	560	ppb m s⁻¹
HNO2	123	ppb m s⁻¹
НСНО	930	ppb m s⁻¹
TOLUENE	129	ppb m s⁻¹
BENZENE	259	ppb m s⁻¹
FURAN	387	ppb m s⁻¹

102

104 **Table S3.** NASA DC-8 airborne measurements used in this work. The NASA Langley Research

105 Center Online Custom Data Merging Tool (beta 1.4, <u>https://www-air.larc.nasa.gov/cgi-</u>

106 <u>bin/Driver.cgi?operation=doMerge</u>) is used to generate the 1-Hz merge dataset, except for the

107 DIAL data.

Measurements	Instrument	Reference
532 nm backscatter ratio	NASA Differential Absorption LIDAR (DIAL)	(Hair et al., 2008)
Ozone (O₃), nitrogen oxides (NO _x)	NOAA NOyO3 4-channel chemiluminescence	(Ryerson et al., 2000)
Photolysis frequencies (j-values)	NCAR CCD Actinix Flux Spectroradiometers (CAFS)	(Shetter & Müller, 1999)
Carbon monoxide (CO)	NASA Differential Absorption Carbon monOxide Measurement (DACOM)	(Warner et al., 2010)
Nitrous acid (HONO)	NOAA lodide Time-of-Flight Chemical lonization Mass Spectrometer (ToF CIMS)	(Neuman et al., 2016; Veres et al., 2020)
Peroxyacetyl nitrate (PAN)	GeorgiaTech chemical ionization mass spectrometer (CIMS)	(Huey, 2007)
Formaldehyde (HCHO)	NASA In Situ Airborne Formaldehyde (ISAF)	(Cazorla et al., 2015)
Black carbon (BC)	NOAA Single-Particle Soot Photometer (SP2)	(Schwarz et al., 2006)
Organic aerosol (OA), Organic Aerosol-to-organic carbon ratio (OA/OC Ratio)	CU High Resolution Time-of- Flight Aerosol Mass Spectrometer (HR-AMS)	(DeCarlo et al., 2006)
Ethane	CU Compact Atmospheric Multispecies Spectrometer (CAMS)	(P. Weibring et al., 2006; Petter Weibring et al., 2007)
Propane, i-butane, n-butane, i- pentane, n-pentane, i- butene/1-butene, isoprene, MVK, MACR, MEK, furan, 2- methylfuran,3-methylfuran, furfural	NCAR Trace Organic Gas Analyzer (TOGA)	(Apel et al., 2015; Wang et al., 2019)
Benzene, toluene, acetaldehyde, methanol, ethanol, acrolein, acetone/propanal, maleic anhydride, phenol, styrene	NOAA Proton-Transfer Reaction Time-of-Flight Mass Spectrometry (PTR-ToF-MS)	(Yuan et al., 2016)

108



- 111 Figure S1. The vertical profiles of potential temperature (θ), water vapor, and wind used to
- initialize the LES model. These profiles are derived from the 12-km mesoscale WRF-Chem
- simulation (Section S2).



- 116 Figure S2. The vertical profiles of chemicals used to initialize the LES model. These profiles are
- derived from the 12-km mesoscale WRF-Chem simulation (Section S2).
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122 Figure S3. Selected airborne measurements collected during the "in-plume" Segments A and B.

123



125 Figure S4. Measured j-HONO (top) during Segment B and modeled j-HONO across sections

126 (bottom) at 10 km, 15 km, and 30 km downwind of the fire source. Also shown in the top panel

are the modeled j-HONO sampled at the same distance from the fire, at 2 km above the model

128 surface.

129







- 142 Figure S8. Same as Figure S4 but for OC. OC is calculated from CU HR-AMS measured OA and
- 143 OA/OC Ratio.
- 144

141



- 147 Figure S9. Same as Figure S4 but for the ratio between HONO and NO₂ (ppb ppb⁻¹).
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- 150 Figure S10. Modeled plume cross sections of HONO photolysis, heterogeneous HONO
- 151 production, other primary OH source (mostly photolysis of O₃ and aldehydes), and secondary
- 152 OH source, at X = 5, 10, and 30 km.

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155 Figure S11. U.S. wildfires in 2019 larger than 1 km² (fire size) with 50 km buffer zones (grey

- patches) overlaid on the U.S. population density map. The wildfire sizes are reported in FINN
- 157 version 2. The gridded U.S. population density map is from (Center for International Earth
- 158 Science Information Network, Columbia University, 2020).

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161 References

162 Ammann, M., Cox, R. A., Crowley, J. N., Jenkin, M. E., Mellouki, A., Rossi, M. J., et al. (2013). Evaluated 163 kinetic and photochemical data for atmospheric chemistry: Volume VI – heterogeneous 164 reactions with liquid substrates. Atmospheric Chemistry and Physics, 13(16), 8045–8228. 165 https://doi.org/10.5194/acp-13-8045-2013 166 Apel, E., Hornbrook, R., Hills, A., Blake, N., Barth, M., Weinheimer, A., et al. (2015). Upper tropospheric 167 ozone production from lightning NOx-impacted convection: Smoke ingestion case study from 168 the DC3 campaign. Journal of Geophysical Research: Atmospheres, 120(6), 2505–2523. 169 https://doi.org/10.1002/2014JD022121 170 Cazorla, M., Wolfe, G. M., Bailey, S. A., Swanson, A. K., Arkinson, H. L., & Hanisco, T. F. (2015). A new 171 airborne laser-induced fluorescence instrument for in situ detection of formaldehyde 172 throughout the troposphere and lower stratosphere. Atmos. Meas. Tech., 8(2), 541–552. 173 https://doi.org/10.5194/amt-8-541-2015 174 Center for International Earth Science Information Network, Columbia University. (2020). Gridded 175 Population of the World, Version 4 (GPWv4): Population Density Revision UN WPP Country 176 Totals, Revision 11. Palisades, NY: NASA Socioeconomic Data and Applications Center (SEDAC). 177 Retrieved from https://doi.org/10.7927/H4F47M65 178 Coggon, M. M., Lim, C. Y., Koss, A. R., Sekimoto, K., Yuan, B., Gilman, J. B., et al. (2019). OH chemistry of 179 non-methane organic gases (NMOGs) emitted from laboratory and ambient biomass burning 180 smoke: evaluating the influence of furans and oxygenated aromatics on ozone and secondary 181 NMOG formation. Atmospheric Chemistry and Physics, 19(23), 14875–14899. 182 https://doi.org/10.5194/acp-19-14875-2019 183 DeCarlo, P. F., Kimmel, J. R., Trimborn, A., Northway, M. J., Jayne, J. T., Aiken, A. C., et al. (2006). Field-184 deployable, high-resolution, time-of-flight aerosol mass spectrometer. Analytical Chemistry, 185 78(24), 8281-8289. 186 Hair, J. W., Hostetler, C. A., Cook, A. L., Harper, D. B., Ferrare, R. A., Mack, T. L., et al. (2008). Airborne 187 High Spectral Resolution Lidar for profiling aerosol optical properties. Applied Optics, 47(36), 188 6734-6752. https://doi.org/10.1364/AO.47.006734 189 Huey, L. G. (2007). Measurement of trace atmospheric species by chemical ionization mass spectrometry: Speciation of reactive nitrogen and future directions. Mass Spectrometry Reviews, 190 191 26(2), 166–184. https://doi.org/10.1002/mas.20118 Knote, C., Hodzic, A., Jimenez, J. L., Volkamer, R., Orlando, J. J., Baidar, S., et al. (2014). Simulation of 192 193 semi-explicit mechanisms of SOA formation from glyoxal in aerosol in a 3-D model. Atmospheric 194 Chemistry and Physics, 14(12), 6213–6239. 195 Neuman, J. A., Trainer, M., Brown, S. S., Min, K.-E., Nowak, J. B., Parrish, D. D., et al. (2016). HONO 196 emission and production determined from airborne measurements over the Southeast U.S. 197 Journal of Geophysical Research: Atmospheres, 121(15), 9237–9250. 198 https://doi.org/10.1002/2016JD025197 Patton, E. G., Sullivan, P. P., & Moeng, C.-H. (2005). The Influence of Idealized Heterogeneity on Wet and 199 200 Dry Planetary Boundary Layers Coupled to the Land Surface. Journal of Atmospheric Sciences, 201 62(7), 2078–2097. https://doi.org/10.1175/JAS3465.1 Ryerson, T., Williams, E., & Fehsenfeld, F. (2000). An efficient photolysis system for fast-response NO2 202 203 measurements. Journal of Geophysical Research: Atmospheres, 105(D21), 26447–26461. 204 https://doi.org/10.1029/2000JD900389

- Schwarz, J. P., Gao, R. S., Fahey, D. W., Thomson, D. S., Watts, L. A., Wilson, J. C., et al. (2006). Single particle measurements of midlatitude black carbon and light-scattering aerosols from the
 boundary layer to the lower stratosphere. *Journal of Geophysical Research: Atmospheres*,
 111(D16). https://doi.org/10.1029/2006JD007076
- Shetter, R. E., & Müller, M. (1999). Photolysis frequency measurements using actinic flux
 spectroradiometry during the PEM-Tropics mission: Instrumentation description and some
 results. *Journal of Geophysical Research: Atmospheres, 104*(D5), 5647–5661.
 https://doi.org/10.1029/98JD01381
- Veres, P. R., Neuman, J. A., Bertram, T. H., Assaf, E., Wolfe, G. M., Williamson, C. J., et al. (2020). Global
 airborne sampling reveals a previously unobserved dimethyl sulfide oxidation mechanism in the
 marine atmosphere. *Proceedings of the National Academy of Sciences*, *117*(9), 4505–4510.
 https://doi.org/10.1073/pnas.1919344117
- Wang, S., Hornbrook, R. S., Hills, A., Emmons, L. K., Tilmes, S., Lamarque, J.-F., et al. (2019). Atmospheric
 Acetaldehyde: Importance of Air-Sea Exchange and a Missing Source in the Remote
 Troposphere. *Geophysical Research Letters*. https://doi.org/10.1029/2019GL082034
- Warner, J. X., Wei, Z., Strow, L. L., Barnet, C. D., Sparling, L. C., Diskin, G., & Sachse, G. (2010). Improved
 agreement of AIRS tropospheric carbon monoxide products with other EOS sensors using
 optimal estimation retrievals. *Atmospheric Chemistry and Physics*, *10*(19), 9521–9533.
 https://doi.org/10.5194/acp-10-9521-2010
- Weibring, P., Richter, D., Fried, A., Walega, J. G., & Dyroff, C. (2006). Ultra-high-precision mid-IR
 spectrometer II: system description and spectroscopic performance. *Applied Physics B*, 85(2),
 207–218. https://doi.org/10.1007/s00340-006-2300-4
- Weibring, Petter, Richter, D., Walega, J. G., & Fried, A. (2007). First demonstration of a high performance
 difference frequency spectrometer on airborne platforms. *Optics Express*, *15*(21), 13476–13495.
 https://doi.org/10.1364/OE.15.013476
- Yuan, B., Koss, A., Warneke, C., Gilman, J. B., Lerner, B. M., Stark, H., & de Gouw, J. A. (2016). A high resolution time-of-flight chemical ionization mass spectrometer utilizing hydronium ions (H₃O⁺
 ToF-CIMS) for measurements of volatile organic compounds in the atmosphere. *Atmospheric*
- 233 *Measurement Techniques, 9*(6), 2735–2752. https://doi.org/10.5194/amt-9-2735-2016
- 234