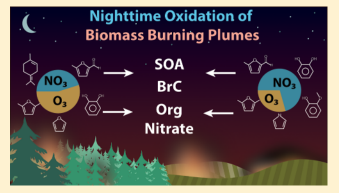


Nighttime Chemical Transformation in Biomass Burning Plumes: A Box Model Analysis Initialized with Aircraft Observations

Zachary C. J. Decker, †,‡,\$® Kyle J. Zarzana,†,\$,◆ Matthew Coggon,†,\$® Kyung-Eun Min,†,\$,# Ilana Pollack,†,® Thomas B. Ryerson, Jeff Peischl,†,\$ Pete Edwards, William P. Dubé,†,\$ Milos Z. Markovic,†,○ James M. Roberts,†,\$ Patrick R. Veres, Martin Graus,†,∇ Carsten Warneke,†,\$ Joost de Gouw, †,‡6 Lindsay E. Hatch, Kelley C. Barsanti, and Steven S. Brown*,‡,§

Supporting Information

ABSTRACT: Biomass burning (BB) is a large source of reactive compounds in the atmosphere. While the daytime photochemistry of BB emissions has been studied in some detail, there has been little focus on nighttime reactions despite the potential for substantial oxidative and heterogeneous chemistry. Here, we present the first analysis of nighttime aircraft intercepts of agricultural BB plumes using observations from the NOAA WP-3D aircraft during the 2013 Southeast Nexus (SENEX) campaign. We use these observations in conjunction with detailed chemical box modeling to investigate the formation and fate of oxidants (NO₃, N₂O₅, O₃, and OH) and BB volatile organic compounds (BBVOCs), using emissions representative of agricultural burns (rice



straw) and western wildfires (ponderosa pine). Field observations suggest NO₃ production was approximately 1 ppbv hr⁻¹ while NO₃ and N₂O₅ were at or below 3 pptv, indicating rapid NO₃/N₂O₅ reactivity. Model analysis shows that >99% of NO₃/ N₂O₅ loss is due to BBVOC + NO₃ reactions rather than aerosol uptake of N₂O₅. Nighttime BBVOC oxidation for rice straw and ponderosa pine fires is dominated by NO₃ (72, 53%, respectively) but O₃ oxidation is significant (25, 43%), leading to roughly 55% overnight depletion of the most reactive BBVOCs and NO₂.

INTRODUCTION

Wildfire size and frequency in the Western U.S. has increased over the last 20 years, and these trends are projected to continue due to factors such as forest management practices, elevated summer temperatures, earlier snowmelt, and drought.1,2 Biomass burning (BB), including wildfires, prescribed burning, and agricultural burning, represents a large, imperfectly characterized, and chemically complex source of reactive material to the troposphere. BB releases reactive species and particulate matter that impact the radiative balance of the atmosphere, air quality, and human health on local to global scales. 3-7 The gas-phase components of BB plumes include volatile organic compounds (BBVOCs) as well as nitrogen oxides (NO_x= NO + NO₂ and higher oxides such as peroxyacyl and alkyl nitrates), oxidants, and oxidant precursors. The air quality and climate effects of BB emissions are defined in part by the oxidative processes and atmospheric chemical cycles that occur as the smoke is transported, diluted, and exposed to oxidants over the hours and weeks following emission.

The photochemistry of BB plumes has been studied previously in a number of field and laboratory studies. Daytime BB plumes can have OH concentrations 5-10 times higher than background air, and daytime reactions of NO_x, BBVOCs, and OH involve complex pathways that generally lead to O₃ formation, but in some cases to near-field O₃ titration. 9-14 Much less is known about nighttime BB plume oxidative processes, which are expected to be dominated by nitrate radicals (NO₃) and O₃. NO₃ is formed by O₃ oxidation of

September 24, 2018 Received: January 22, 2019 Revised: Accepted: January 30, 2019 Published: January 30, 2019

[†]Cooperative Institute for Research in Environmental Sciences, University of Colorado, Boulder, Colorado 80309, United States *Department of Chemistry, University of Colorado, Boulder, Colorado 80309-0215, United States

[§]NOAA Earth System Research Laboratory (ESRL), Chemical Sciences Division, Boulder, Colorado 80305, United States

Wolfson Atmospheric Chemistry Laboratories, Department of Chemistry, University of York, York YO10 5DD, United Kingdom

¹Department of Chemical and Environmental Engineering and College of Engineering – Center for Environmental Research and Technology (CE-CERT), University of California, Riverside, California 92507, United States

 NO_x (eq R1 and Figure 1) but is rapidly (τ < 10 s) destroyed in the daytime by NO and photolysis. ^{15,16} NO₃ is a precursor for N₂O₅ (eq R2), a NO_x reservoir. N₂O₅ may undergo heterogeneous uptake to form ClNO₂ and HNO₃ (eq R3). The former is a daytime Cl radical precursor affecting both marine and continental environments and influencing next-day O₃ production. ^{17–20} NO₃ can also be directly taken up onto aerosol (eq R4).

$$NO_2 + O_3 \rightarrow NO_3 \tag{R1}$$

$$NO_3 + NO_2 \rightleftharpoons N_2O_5 \tag{R2}$$

$$N_2O_{S(g)}$$
 + aerosol $\rightarrow \phi CINO_2 + (2 - \phi)HNO_3$ (R3)

$$NO_3$$
 + aerosol \rightarrow products (R4)

Mixing of background or smoke-derived 14 O3 with NO, in a BB plume leads to the production of NO₃, which may be rapid (>0.5 ppbv h⁻¹). Recent laboratory measurements conducted during both the Fire Lab at Missoula Experiment (FLAME-4) and the ongoing Fire Influence on Regional and Global Environments Experiment (FIREX) have provided detailed identification and quantification of emissions for a range of BBVOCs. 4,5,21-23 Emissions inventories from these experiments indicate that the compounds emitted and their relative concentrations depend on the fuel type (e.g., pine vs grass), combustion process (e.g., smoldering or flaming), ignition procedure (e.g., fast or slow), and pyrolysis temperature (e.g., high or low). 4,21,24,25 Generally, primary BBVOC emissions include oxygenated hydrocarbons and aromatics (e.g., phenols) as well as unsaturated hydrocarbons, biogenic and heteroaromatic species. 4,5,21 Many such compounds are very reactive toward NO₃²⁶⁻³³ and may significantly limit its lifetime, promote secondary organic aerosol formation (SOA), 34,35 and alter nighttime oxidative budgets.

The coemission of NO_x highly reactive VOCs, and aerosol particles leads to the potential for significant nighttime chemical transformations. Despite this potential, there has

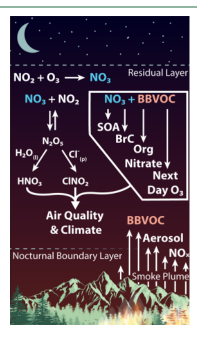


Figure 1. Schematic of nighttime NO_3 and N_2O_5 chemical processing in a biomass burning plume.

been only one aircraft campaign to date from which sampling of nighttime biomass burning plumes has been reported. 36,37 The Southeast Nexus (SENEX) campaign in 2013 included 20 research flights of an instrumented NOAA WP-3D aircraft, and one of the goals was to study the interactions between anthropogenic and biogenic emissions. 38 A night flight on July 2–3 targeted the emissions and nighttime chemistry from a power plant plume near the Mississippi river. During this flight, the WP-3D also targeted and intercepted agricultural BB plumes yielding the first airborne study of nighttime smoke that included NO₃ and N₂O₅ measurements. 36 Even so, there has been no previous analysis of BB NO₃ chemistry using nighttime aircraft intercepts.

Here, we present the first analysis of nighttime smoke oxidation based on aircraft intercepts of fire plumes using data from this flight. With these observations, we initiate a detailed chemical box model to understand the chemical evolution of oxidants (NO₃, N₂O₅, O₃, and OH) and BBVOCs over one night (10 h) using emissions for rice straw to model a generic agricultural burning plume. We then use this analysis to model nighttime chemistry in western wildfires using emissions for a ponderosa pine fire.

Field and Laboratory Measurements. Field data for this study were taken from multiple instruments deployed on the NOAA WP-3D aircraft during the SENEX 2013³⁸ flight on July 2–3, 2013 (20:00–03:00 CDT). Our analysis utilizes data from the NOAA nitrogen oxide cavity ring-down spectrometer (CRDS) for NO₂, NO₃, N₂O₅, and O₃^{39–42} as well as the NO₂O₃ chemiliuminescence instrument (CL) for NO, NO₂, O₃, and NO₂⁴³ with 1 Hz acquisition resolution. Within the plume regions we study, the measurements of NO₂ and O₃ from the CRDS and CL instruments agree within 7%. We also use data from an ultrahigh sensitivity aerosol spectrometer (UHSAS) for aerosol size measurements (1 Hz)^{44,45} and a proton-transfer-reaction mass spectrometer (PTR-MS) for VOC measurements (1 s every 17 s).⁴⁶

BB intercepts were identified by the enhancement above background of four species: black carbon (BC), glyoxal (CHOCHO), CO, and acryloyl peroxynitrate (APAN). BB identifier data were provided by the NOAA airborne cavity enhanced spectrometer (ACES) for glyoxal, iodide chemical ionization mass spectrometer (I CIMS) for APAN, single particle soot photometer (SP2) for black carbon, and vacuum ultraviolet fluorimeter for CO. Power plant plumes were identified by the above background enhancements of NO_x and N₂O₅. While CO is also present in the power plant plumes, the three other BB identifiers were not. Information on background and plume measurements is included in the Supporting Information (SI) (Tables S1 and S2).

Five VOCs (toluene, isoprene + furan, methylvinylketone + methacrolein (MVK + MACR), and methylethylketone (MEK)) as well as acetonitrile were measured by the PTR-MS during SENEX and overlap with our inventory. However, we explain in the SI that we do not use these observations because we do not know the fire source, number of fires, or fuel, and plume age estimates are highly uncertain (Figure S5).

Our detailed chemical box model uses emission inventories from Hatch et al. 5 and Koss et al. 4 for the ponderosa pine and rice straw fuels. The BBVOC emissions from Hatch et al. 5,21 were measured during FLAME-4 using the following instruments: two-dimensional gas chromatography—time-of-flight mass spectrometry, open-path Fourier-transform infrared spectroscopy, 22 whole-air sampling with one-dimensional gas

chromatography—mass spectrometry, and PTR time-of-flight mass spectrometry (PTR-ToF). BBVOC emissions from Koss et al. Were measured by PTR-ToF during FIREX. Details regarding how the two inventories were merged are included in the SI. In general, for compounds shared between both inventories, the emission ratios (E1) agree within an order of magnitude with some exceptions (Figure S6). We propagate this variability into our model results (SI).

Analysis and Modeling Methods. We report our emissions in the form of laboratory-derived emission ratios (ER), which is the background subtracted emitted compound (x) normalized to background subtracted CO.^{4,21}

$$ER_{x} = \frac{x(ppbv)}{CO(ppmv)}$$
(E1)

These emissions are integrated over the entirety of the laboratory fires and therefore contain emissions from all stages of the fire.

The modified combustion efficiency (MCE) was calculated for each plume.

$$MCE = \frac{CO_2 - CO_{2bkg}}{(CO_2 - CO_{2bkg}) + (CO - CO_{bkg})}$$
(E2)

During plume intercepts, the average MCE was 95 \pm 6%, which is consistent with previous MCE calculations of the July 2–3 night flight.³⁶

The total NO₃ reactivity toward BBVOCs is given by

$$k_{\text{NO}_3}^{\text{BBVOC}} = \sum k_{\text{NO}_3 + \text{BBVOC}_i} [\text{BBVOC}_i]$$
 (E3)

where $k_{\mathrm{NO_3+BBVOC}_i}$ is the bimolecular rate coefficient for $\mathrm{NO_3}$ + BBVOC_i and $k_{\mathrm{NO_3}}^{\mathrm{BBVOC}}$ is the pseudo-first order rate coefficient. The bimolecular rate coefficients for $\mathrm{NO_3}$, $\mathrm{O_3}$, or OH + BBVOC were taken from literature where available and estimated by structure—activity relationships 31,53 or structural similarity where unavailable (SI).

Due to limited literature on NO_3 + BBVOC rate coefficients, our inventory excludes many nitriles, amines, alkynes, acids, and other compounds whose rate coefficients were unavailable and could not be estimated. We also removed saturated hydrocarbons because they are generally unreactive toward NO_3 . Despite this, our merged inventory retains about 87% of the total inventory carbon mass, or 96% by mass, with 235 compounds from Hatch et al. and 171 compounds from Koss et al. with 103 compounds shared in both inventories for a total of 303 unique compounds.

To calculate the observed NO_3 reactivity during SENEX BB plume intercepts we determined BBVOC concentration using background corrected CO measured on the WP-3D.

$$BBVOC(ppbv) = ER_{BBVOC}(CO - CO_{bk\sigma})$$
(E4)

As shown below, BBVOC is likely the main sink of NO₃; therefore, the extent of BBVOC oxidation by NO₃ will be limited by the NO_x/BBVOC ratio as NO_x is the source for NO₃ (eq R1). Furthermore, the relative oxidative importance between O₃ and NO₃ depends on the NO_x/BBVOC ratio as explained by Edwards et al. Therefore, in contrast to the method used for calculating BBVOC concentration in SENEX fire plume intercepts described above, we initiate our box model with fire emissions scaled to NO_x in order to preserve

the NO_x/BBVOC ratio observed during the fire lab experiments.

To estimate the emitted NO_x at the fire source, we assume that the total reactive nitrogen (NO,, which does not include NH_3) is equivalent to the emitted NO_x . The NO_x/NO_y ratio as measured during SENEX fire plume intercepts in Figure 2 was 0.84. We calculated the observed NO_{ν} emission ratio using NO_{v} (13.2 ± 3.1 ppbv) and CO (543.4 ± 87.7 ppbv) enhancements above background. The calculated NO, emission ratio, which we assume to be the NO_x emission ratio at the fire source, was determined to be 24.3 ± 6.4 ppbv NO_v/ppmv CO for the plume intercept. We compared the estimated observed NO_x emission ratio to the NO_x emission ratios reported by Selimovic et al. for rice straw (43.9 ppbv $NO_x/ppmv$ CO) and ponderosa pine (26.9 \pm 4.3 ppbv $NO_x/$ ppmv CO).²³ We then scaled the BBVOC emissions by this ratio (E5), effectively scaling the fire emissions to the NO_x of the observed fire plume.

$$[BBVOC]^{model} = [BBVOC]^{inventory} \frac{ER_{NO_y}^{obsd}}{ER_{NO_x}^{inventory}}$$
(E5)

The NO_x emission ratio observed during the SENEX fire plume intercepts in Figure 2 was 45% and 11% lower than the laboratory-derived NO_x emission ratio for rice straw and ponderosa pine fires, respectively. To correctly model the NO_3

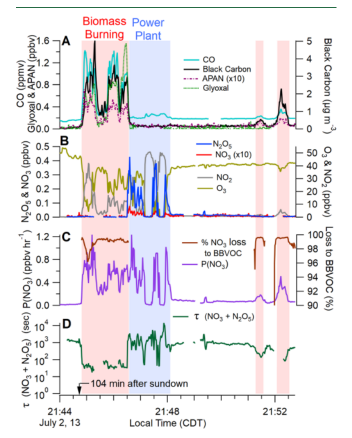


Figure 2. Time traces during representative sections of BB (red) and power plant (blue) plume intercepts made 104 min after sundown (SZA = 90°): (A) BB tracers, (B) NO₃ N₂O₅, NO₂, and O₃ mixing ratio, (C) production rate of NO₃ and the percentage of NO₃ reactivity toward BBVOCs, (D) lifetime of NO₃ and N₂O₅.

Environmental Science & Technology

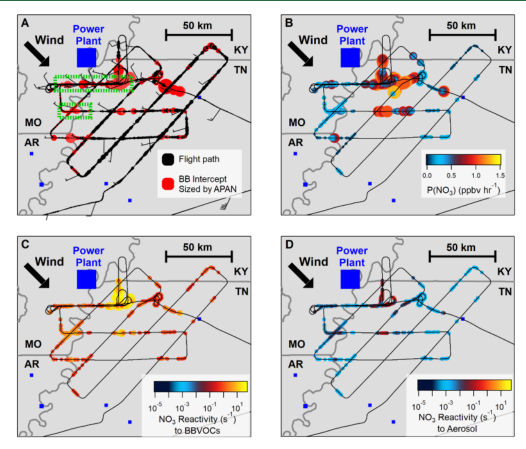


Figure 3. Flight maps of the SENEX July 2–3, 2013, night flight. (A) BB intercepts colored by red markers, sized by APAN (0.01–0.1 ppbv), and green dashes indicate sections shown in Figure 2. (B) Production rate of NO₃. (C, D) Comparisons of NO₃ reactivity toward BBVOCs (C) and toward aerosol (D) on the same color and log scale.

oxidation of these fires we reduced our BBVOC emissions by a factor of 55% for rice straw and 89% for ponderosa pine.

Model background and initial concentrations of NO_x , CO_y , and O_3 were taken from the SENEX observations shown in Figure 2. We estimate the NO/NO_2 ratio at the fire source using the NO and NO_2 emission ratios from FIREX for each fuel. The NO/NO_2 ratios used are 5.3 and 2.8 for rice straw and ponderosa pine, respectively.²³ The background NO_2 mixing ratio was taken to be 0.9 ppbv. The background O_3 mixing ratio, 43.9 ppbv, was used as the starting O_3 mixing ratio and is representative of the background O_3 in the region where BB plumes were intercepted (Figure S3).

Box modeling was performed using the Framework for 0-D Atmospheric Modeling (F0AM)⁵⁵ to investigate the evolution of oxidized mass and oxidant fractions over 10 h (the approximate duration of one night in July in the Southeastern U.S.). Chemical mechanisms were adopted from the MCM (v3.3.1,⁵⁶⁻⁶⁰ via http://mcm.york.ac.uk), and published mechanisms for methylguaiacol, syringol, *o*-guaiacol, and 3-methylfuran were added (Table S4).⁶¹⁻⁶³ Compounds not included in the above references were modeled as a one-step reaction of BBVOC + NO₃, BBVOC + O₃, or BBVOC + OH to form a single oxidation product.

All models were run at 298 K, typical experimental conditions for most published rate coefficients. Temperatures during flight ranged between 288 and 290 K (SI). In order to account for dilution processes, as well as entrainment of O_3 , we apply a first order dilution of $k_{\rm dil} = 1.16 \times 10^{-5} \ \rm s^{-1}$ or a 24 h lifetime. The sensitivity of this assumption is shown in Figure

S2 and discussed in the SI. We report a base case model result with upper and lower bound uncertainties based on the emission and rate coefficient uncertainties, although, as discussed in the SI, the bounds do not provide information on the error distribution.

RESULTS AND DISCUSSION

In panel A of Figure 2 the power plant plume intercepts (blue background) are distinguished from the fire plume intercepts (red background) by CO, black carbon, APAN, and glyoxal. Intercepts shown in Figure 2 were at an altitude between 700 and 900 m. Relative to the BB plume intercepts, the power plant plume intercepts exhibited elevated levels of NO_3 and N_2O_5 (Figure 2B). Figure 3A shows a flight map of the July 2–3 flight colored red during BB plume intercepts and sized by the APAN mixing ratio. Roughly 97% of the indicated BB plumes do not show signs of power plant plume mixing (SI). Green dashed boxes indicate sections of data shown in Figure 2.

The flight covered the intersection of Missouri, Kentucky, Tennessee, and Arkansas at the Mississippi river. According to the USDA CropScape database, this land is mainly agricultural, and therefore, the fire plume is most likely the result of burning crop residue and stubble. Hume intercepts occurred near winter wheat crops and rice straw crops are situated roughly 70 km northwest. Still, rice straw is the best available fuel proxy for agricultural burning emissions. The wind direction was roughly northwesterly with most BB plume intercepts occurring in the northwest corner of Tennessee.

Environmental Science & Technology

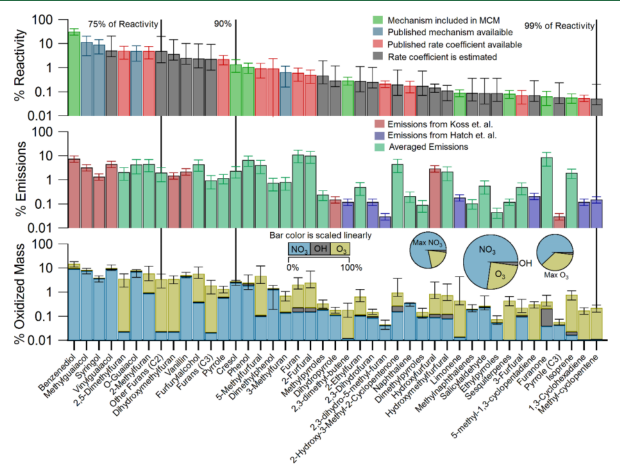


Figure 4. Rice straw fuel. The top panel shows the ranked order of the compounds that account for 99% of the rice straw initial NO₃ reactivity. The color scale describes the origin of the mechanisms or rate coefficient used. The middle panel is the relative BBVOC emission ratio normalized to the total BBVOC emission ratio, and the color scale describes the origin of the emissions data. The bottom panel is the relative nighttime reacted mass (10 h) normalized to total reacted mass. While the bar height is on a log scale, the color scale is linear and indicates the fraction of oxidation by NO₃ (blue), O₃ (gold), and OH (gray). The center pie chart shows the fraction of reacted mass in the base case with the maximum NO₃ oxidation case to the left and maximum O₃ oxidation case to the right. All panels sum to 100%.

To illustrate the NO $_3$ chemistry within a BB plume, we use previously published NO $_3$ and N $_2$ O $_5$ analysis metrics. The NO $_3$ production rate, P(NO $_3$), is the instantaneous source of NO $_3$ from the reaction of NO $_2$ with O $_3$ and is given in (E6). The NO $_3$ + N $_2$ O $_5$ lifetime (τ) is the ratio of NO $_3$ and N $_2$ O $_5$ concentration to the NO $_3$ production rate (E7). The summed lifetime is useful because NO $_3$ and N $_2$ O $_5$ reach an equilibrium state that is typically more rapid than the individual sink reactions for either, such that they can be regarded as a sum.

$$P(NO_3) = k_{NO_3}[NO_2][O_3]$$
 (E6)

$$\tau(NO_3 + N_2O_5) = \frac{NO_3 + N_2O_5}{P(NO_3)}$$
 (E7)

 $P(NO_3)$ was large and of similar magnitude in both the power plant plume and BB plume (Figure 2C). Figure 3B is colored by NO_3 production during BB intercepts only, and shows that large NO_3 production rates, near 1 ppbv hr^{-1} , were observed during multiple BB plume intercepts. Despite the large NO_3 radical production, the NO_3 and N_2O_5 concentrations within the BB plume were below the 3 pptv³⁸ stated detection limit of the instrument (Figure 2B), yielding short $NO_3 + N_2O_5$ lifetimes. Indeed, as shown in Figure 2D, τ is roughly a factor of 100 lower within the BB plume as

compared to the power plant plume and background air. Because the NO_3 and N_2O_5 were below stated detection limits in the BB plumes, the corresponding lifetimes shown in Figure 2D are upper limits, and the actual lifetimes may be considerably shorter.

The high production rate and short lifetime of $NO_3 + N_2O_5$ within the BB plume is evidence for rapid NO_3 or N_2O_5 loss pathways. BB plumes contain large quantities of both aerosol and BBVOCs, which provide two efficient NO_3/N_2O_5 loss pathways. To understand the competition between these loss processes, we calculated an instantaneous NO_3 reactivity toward aerosol and toward BBVOCs. The total NO_3 loss to BBVOC is calculated using the sum of BBVOC reactivity normalized to CO (E3). The total NO_3 loss to aerosol uptake is given as the sum of both NO_3 and N_2O_5 uptake rate coefficients. By assuming a steady state for both NO_3 and N_2O_5 , we estimate the total aerosol uptake, and therefore NO_3 reactivity toward aerosol, as

$$k_{\text{NO}_3}^{\text{aerosol}} = K_{\text{eq}}[\text{NO}_2]k_{\text{N}_2\text{O}_3+\text{aerosol}} + k_{\text{NO}_3+\text{aerosol}}$$
(E8)

where $k_{\mathrm{NO_3}}^{\mathrm{aerosol}}$ is a first-order rate coefficient, K_{eq} is the equilibrium constant between NO₃ and N₂O₅ (R2), and $k_{x+\mathrm{aerosol}}$ is the first order rate coefficient for N₂O₅ or NO₃ aerosol uptake expressed below.

Environmental Science & Technology

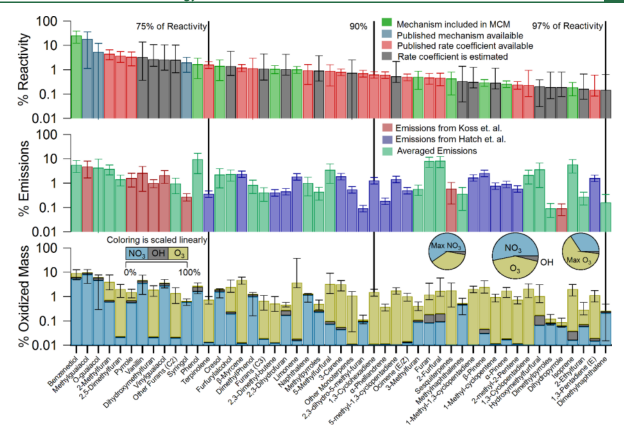


Figure 5. Same as Figure 4 but for ponderosa pine fuel. In the bottom panel the bar height is on a log scale, but the color scale is linear and indicates the fraction of oxidation by NO₃ (blue), O₃ (gold), and OH (gray).

$$k_{x+\text{aerosol}} = \frac{\gamma \overline{c} SA}{4} \tag{E9}$$

Here, γ is the aerosol uptake coefficient, \overline{c} is the mean molecular speed, and SA is the aerosol surface area. Calculations use uptake coefficients of $\gamma_{\rm N_2O_5}=10^{-2}$ for ${\rm N_2O_5}^{19}$ and $\gamma_{\rm NO_3}=10^{-3}$ for ${\rm NO_3}$. However, $\gamma_{\rm NO_3}$ values have a wide range; therefore, we include calculations with $\gamma_{\rm NO_3}=1$ in the SI but find similar results. ¹⁵

Parts C and D of Figure 3 compare the NO_3 reactivity toward BBVOCs and aerosol uptake during BB plume intercepts, respectively. In all BB intercepts, the calculated NO_3 reactivity toward BBVOCs is a factor of 100-1000 greater than aerosol uptake. Figure 2C shows the percentage of NO_3 reactivity dominated by BBVOC with a median >99%.

To understand which BBVOCs may be responsible for the rapid initial loss of NO_3 , we calculated the relative NO_3 reactivity for 303 compounds in rice straw and ponderosa pine burning emissions. The top panel of Figure 4 shows the ranked order of the compounds that account for 99% of the rice straw initial NO_3 reactivity. Eight furan or phenol compounds are responsible for 75% of the initial NO_3 reactivity. Most of, the initial NO_3 reactivity for a rice straw fire is accounted for by phenols $(60^{+20}_{-14}\%)$ and furans $(23^{+20}_{-6}\%)$, as well as pyrroles and furfurals $(8^{+9}_{-3}\%)$ combined).

The top panel of Figure 5 shows the ranked order of the compounds that account for 97% of the ponderosa pine initial NO₃ reactivity. The top 75% of initial NO₃ reactivity is distributed among 13 compounds with phenols $(62^{+27}_{-23}\%)$, furans $(18^{+12}_{-4}\%)$, and pyrrole and furfural $(8^{+8}_{-3}\%)$ combined) again dominating the total reactivity. Unlike rice straw, a

ponderosa pine fire plume has significant reactivity toward terpenes $(8^{+2}_{-1}\%)$. The initial NO_3 reactivity toward terpenes and unsaturated hydrocarbons in a rice straw plume is <1%. These differences in reactivity are due to differences in emissions between the two fuels as explained below.⁵

The middle panels of Figures 4 and 5 show the emission ratios for each compound normalized to total emissions. The color indicates the origin of the emission ratio. The rice straw fire emissions for compounds included in Figure 4 are mainly furans $(33 \pm 8\%)$, phenols $(27 \pm 4\%)$, and furfurals $(24 \pm 6\%)$, while unsaturated hydrocarbon and terpene emissions account for only $3 \pm 1\%$. In contrast, the ponderosa pine fire emissions have a larger representation of terpenes $(18 \pm 4\%)$ and unsaturated hydrocarbons $(10 \pm 2\%)$, but phenols $(33 \pm 10\%)$, furans $(17 \pm 4\%)$, and furfurals $(18 \pm 6\%)$ are all still significant.

To better understand smoke plume evolution and to determine the amount of BBVOC mass oxidized during one night (10 h), we ran a 0-D box model for both rice straw and ponderosa pine fire emissions. NO₃ and N₂O₅ remained below 3 pptv (Figure S1), consistent with field observations (Figure 2B). Figure S1 illustrates that the summed concentrations of the most reactive BBVOCs are comparable to NO₂, suggesting there is approximately as much NO₃ precursor available as there is BBVOC to be oxidized. For both fuels, roughly 50–60% of NO₂ and the BBVOC compounds listed in Figure 4 and Figure 5 are depleted by chemistry (excluding dilution) in one night. Our box-model suggests several abundant BBVOCs survive the night with more than 50% of their initial starting concentration, such as phenol, furan, furfural, and hydroxymethylfurfural (SI).

HNO₃ production is complex within the model, and both maximum and minimum uncertainty bounds on HNO₃ concentrations are the result of higher bound BBVOC emissions, but lower and higher bound BBVOC rate coefficients, respectively. HNO₃ is the product of reactions of phenolic compounds with NO₃, which proceeds by H-abstraction. HNO₃ production is dominated by catechol + NO₃ (~60%) within the first few hours, but as the more reactive compounds are depleted, the lesser reactive compounds like methyl guaiacol, guaiacol, and syringol react with NO₃ and dominate in the last 2 h. HNO₃ may be lost to the particle phase with concurrent NH₃ emission or other nitrogen species; however, this loss mechanism is not included in our model.

For both fuels, catechol is the most reactive compound, and accounts for $32 \pm 9\%$ and $26 \pm 13\%$ of initial NO₃ reactivity at the start of the simulation for rice straw and ponderosa pine plumes, respectively. However, Koss et al.4 were unable to distinguish between catechol and methylfurfural at m/z =110.1 We assume a 50/50 contribution here, which yields catechol emission ratios of 2.5 ± 0.8 ppbv ppmv⁻¹ CO for rice straw and 1.5 \pm 0.6 ppbv ppmv⁻¹ CO for ponderosa pine. Still, the high reactivity is mainly due to the large catechol rate coefficient $(9.9 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}),^{67}$ which is the third greatest among the emitted compounds. Catechol is known to react with NO3 by H-abstraction, with subsequent addition of NO₂ to the aromatic peroxy radical to form 4nitrocatechol with a near-unity molar yield of 0.91 ± 0.06 . Further, 4-nitrocatechol is expected to almost completely (96%) partition to the particle phase.⁶⁸ Recently, Hartikainen et al.²⁵ investigated dark oxidation of residential wood combustion and found strong correlations between the depletion of phenolic compounds and the formation of NO₃initiated SOA. In wintertime BB events, 4-nitrocatechol and other derivatives have been detected in aerosol and are considered important light-absorbing components of brown carbon (BrC).3

SOA yields are a function of mass loadings.⁷⁷ Using a catechol mass loading of 300 μ g m⁻³ from Finewax et al.⁶⁸ as well as a total observed aerosol plume measurement of 58.7 μ g m⁻³, we estimate a 4-nitrocatechol SOA mass yield of 120%. Assuming 0.6 ppbv of catechol in ponderosa pine and 0.8 ppbv in rice straw (initial model conditions) with 44 ppbv O₃, 13 ppbv of NO_x and $k_{\rm dil} = 1.16 \times 10^{-5} \, {\rm s}^{-1}$, we estimate the SOA produced from catechol to be 3.8 \pm 1.0 μ g m⁻³ in 8 h and 4.0^{+1.1}_{-1.0} μ g m⁻³ in 8.5 h for a rice straw and ponderosa pine plume, respectively. Further, there is evidence to suggest furans and furfurals may also be a source of SOA precursors.^{5,25}

The bottom panel of Figure 4 shows the reacted mass per compound normalized to the total reacted mass. The bar height is on a log scale, but the bar color is linearly scaled and indicates the fraction of nighttime oxidation by NO_3 (blue), O_3 (gold), and OH (gray) after 10 h for each compound. The center pie chart in Figure 4 and 5 represents the base case fraction of reactant mass oxidized by each oxidant. The left and right pie charts show results for the estimated maximum possible NO_3 and maximum possible O_3 oxidation, respectively. Uncertainty in the fraction of oxidized mass is calculated from the uncertainties in individual compound emissions and rate coefficients. For the compounds comprising a rice straw BB plume, the majority of mass is oxidized by NO_3 ($72^{+6}_{-11}\%$). This is expected because the rice straw fuel emissions are rich in oxygenated aromatic and heteroaromatic emissions, which

are generally less reactive toward O_3 . Terpenes and unsaturated hydrocarbons, which are a small fraction of emissions in Figure 4, are relatively more reactive toward O_3 . Even so, O_3 still has a significant oxidative impact and is responsible for 26^{+11}_{-6} % of oxidized BBVOC mass.

The relative amount of oxidized mass for ponderosa pine is shown in the bottom panel of Figure 5. Almost half of the oxidized mass for compounds included in Figure 5 is due to O_3 (43 $^{+21}_{-6}$ %) for our base case. The phenolic compounds mainly undergo NO_3 oxidation while terpenes and unsaturated hydrocarbons are mainly oxidized by O_3 . Furans and the heteroaromatics are oxidized approximately evenly by O_3 and NO_3 . The increased fraction of O_3 oxidation is the result of the increased fraction of unsaturated hydrocarbon and terpenes in the ponderosa pine emissions when compared to rice straw.

The nighttime chemical evolution and oxidation products of a biomass burning plume will depend on the relative NO₃ and O₃ reactivity. Neglecting the small contribution from OH oxidation, Edwards et al. 54 show the competition between NO₃ and O₃ oxidation of biogenic VOCs (BVOC) is dependent on the NO_x/BVOC ratio. We scaled our BBVOC emissions to maintain the NO_x/BBVOC ratio expected for rice straw (0.4 \pm 0.1) or ponderosa pine (0.3 ± 0.1) emissions. However, because fires are highly variable, the NO_x/BBVOC ratio for any given fuel may vary from fire to fire. For rice straw, a factor of 2 increase in NO_x increases the fraction of NO₃ oxidation from 72% to 84%, while a factor of 2 decrease in NO. decreases relative NO₃ oxidation to 55%. Similarly, for ponderosa pine, doubling NO_x increases the fraction of NO₃ oxidation from 53% to 66%, while halving NO_x decreases relative NO₃ oxidation to 37% and increases O₃ to 57%. Furthermore, we find that a factor of 2 change in ambient O₃ concentration has little effect on the relative NO3 and O3 reactivity (see the SI).

Our reactivity calculations and box-model results are most limited by a lack of kinetic and mechanistic studies for O_3 , NO_3 , and OH + BBVOCs reactions. Kinetic and mechanistic studies of furan, furfural, phenol, and pyrrole analogues reacting with NO_3 will be most critical to understanding nighttime BB processes, which we highlight in the SI.

The time of day in which a fire is active will determine the fate of its emissions. This paper presents the first nighttime aircraft intercepts of a BB plume combined with an inventory of 303 BBVOC emissions and an oxidation model to predict the lifetime and fate of BB emissions in the dark. Fire emissions at times near sunset will undergo the chemistry we have detailed here, which suggests a roughly 60% depletion (for both rice straw and ponderosa pine) of fire-derived NO_x. We find that nighttime chemistry is likely to proceed by NO₃, rather than N_2O_5 , further slowing the loss of NO_x (eqs R1 and R2). Our model applies to chemistry at the center of a plume and does not include dispersion. Dispersion mixes NOx with background O₃ at the edges of the plume leading to faster depletion, and therefore, the values we report are likely lower limits. Even so, 18-19% of BBVOC mass, out of the total BBVOC mass that we model, will be oxidized in one night. That is roughly a 55% depletion of the BBVOCs that are reactive toward NO₃. There is evidence that many of these NO₃ reactive compounds can form secondary BrC aerosol, 35,69-76 suggesting nighttime oxidation may be a significant source of BB derived BrC. Furthermore, future BB photochemical models should consider that these reactive phenolic-, furan-, and furfural-like compounds are not only reactive

toward NO_3 , but also O_3 and OH, thus affecting next-day BB photochemistry.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.8b05359.

Figure S1, box model time traces of key species; Figure S2, box model sensitivity to the dilution rate coefficient; Figure S3, correlation of O_3 and NO_2 from aircraft observations; Figure S4, altitude profiles of key species and potential temperature; Figure S5, plume age estimates; Figure S6, variability in emission ratios; Table S1, BB plume and background values; Table S2, plume and background times; Table S3, list of reactions excluded from the MCM; Table S4, mechanisms added to the MCM (PDF)

AUTHOR INFORMATION

Corresponding Author

*E-mail: steven.s.brown@noaa.gov.

ORCID

Zachary C. J. Decker: 0000-0001-9604-8671 Matthew Coggon: 0000-0002-5763-1925 Joost de Gouw: 0000-0002-0385-1826 Kelley C. Barsanti: 0000-0002-6065-8643

Present Addresses

- ◆(K.J.Z.) Department of Chemistry, University of Colorado, Boulder, Colorado 80309-0215, United States.
- *(K.-E.M.) Gwangju Institute of Science and Technology, School of Earth Sciences and Environmental Engineering, Gwangju 61005, South Korea.
- [⊗](I.P.) Colorado State University, Atmospheric Science Department, Ft. Collins, CO 80523-1371.
- O(M.Z.P.) Picarro, Inc., Santa Clara, CA 95054..
- V(M.G.) Department of Atmospheric and Cryospheric Sciences, University of Innsbruck, Innsbruck 6020, Austria.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We thank Charles A. Brock for aerosol surface area measurements.

■ REFERENCES

- (1) Dennison, P. E.; Brewer, S. C.; Arnold, J. D.; Moritz, M. A. Large Wildfire Trends in the Western United States, 1984–2011. *Geophys. Res. Lett.* **2014**, *41*, 2928.
- (2) Westerling, A. L.; Hidalgo, H. G.; Cayan, D. R.; Swetnam, T. W. Warming and Earlier Spring Increase Western U.S. Forest Wildfire Activity. *Science (Washington, DC, U. S.)* **2006**, 313 (5789), 940–943.
- (3) Pope, C. A.; Dockery, D. W. Health Effects of Fine Particulate Air Pollution: Lines That Connect. *J. Air Waste Manage. Assoc.* **2006**, 56 (6), 709–742.
- (4) Koss, A. R.; Sekimoto, K.; Gilman, J. B.; Selimovic, V.; Coggon, M. M.; Zarzana, K. J.; Yuan, B.; Lerner, B. M.; Brown, S. S.; Jimenez, J. L.; Krechmer, J.; Roberts, J. M.; Warneke, C.; Yokelson, R. J.; de Gouw, J. Non-Methane Organic Gas Emissions from Biomass Burning: Identification, Quantification, and Emission Factors from PTR-ToF during the FIREX 2016 Laboratory Experiment. *Atmos. Chem. Phys.* 2018, 18 (5), 3299–3319.

- (5) Hatch, L. E.; Yokelson, R. J.; Stockwell, C. E.; Veres, P. R.; Simpson, I. J.; Blake, D. R.; Orlando, J. J.; Barsanti, K. C. Multi-Instrument Comparison and Compilation of Non-Methane Organic Gas Emissions from Biomass Burning and Implications for Smoke-Derived Secondary Organic Aerosol Precursors. *Atmos. Chem. Phys.* **2017**, *17* (2), 1471–1489.
- (6) Rotstayn, L. D.; Penner, J. E. Indirect Aerosol Forcing, Quasi Forcing, and Climate Response. *J. Clim.* **2001**, *14* (13), 2960–2975.
- (7) Andreae, M. O.; Merlet, P. Emission of Trace Gases and Aerosols from Biomass Burning. *Global Biogeochem. Cycles* **2001**, *15* (4), 955–966.
- (8) Yokelson, R. J.; Crounse, J. D.; DeCarlo, P. F.; Karl, T.; Urbanski, S.; Atlas, E.; Campos, T.; Shinozuka, Y.; Kapustin, V.; Clarke, A. D.; Weinheimer, A.; Knapp, D. J.; Montzka, D. D.; Holloway, J.; Weibring, P.; Flocke, F.; Zheng, W.; Toohey, D.; Wennberg, P. O.; Wiedinmyer, C.; Mauldin, L.; Fried, A.; Richter, D.; Walega, J.; Jimenez, J. L.; Adachi, K.; Buseck, P. R.; Hall, S. R.; Shetter, R. Emissions from Biomass Burning in the Yucatan. *Atmos. Chem. Phys.* **2009**, *9* (15), 5785–5812.
- (9) Hennigan, C. J.; Sullivan, A. P.; Collett, J. L.; Robinson, A. L. Levoglucosan Stability in Biomass Burning Particles Exposed to Hydroxyl Radicals. *Geophys. Res. Lett.* **2010**, 37 (9), 1–4.
- (10) Hennigan, C. J.; Miracolo, M. A.; Engelhart, G. J.; May, A. A.; Presto, A. A.; Lee, T.; Sullivan, A. P.; McMeeking, G. R.; Coe, H.; Wold, C. E.; Hao, W. M.; Gilman, J. B.; Kuster, W. C.; De Gouw, J.; Schichtel, B. A.; Collett, J. L.; Kreidenweis, S. M.; Robinson, A. L. Chemical and Physical Transformations of Organic Aerosol from the Photo-Oxidation of Open Biomass Burning Emissions in an Environmental Chamber. *Atmos. Chem. Phys.* **2011**, *11* (15), 7669–7686
- (11) Ortega, A. M.; Day, D. A.; Cubison, M. J.; Brune, W. H.; Bon, D.; De Gouw, J. A.; Jimenez, J. L. Secondary Organic Aerosol Formation and Primary Organic Aerosol Oxidation from Biomass-Burning Smoke in a Flow Reactor during FLAME-3. *Atmos. Chem. Phys.* 2013, 13 (22), 11551–11571.
- (12) Bruns, E. A.; El Haddad, I.; Slowik, J. G.; Kilic, D.; Klein, F.; Baltensperger, U.; Prévôt, A. S. H. Identification of Significant Precursor Gases of Secondary Organic Aerosols from Residential Wood Combustion. *Sci. Rep.* **2016**, 6 (June), 1–9.
- (13) Bruns, E. A.; El Haddad, I.; Keller, A.; Klein, F.; Kumar, N. K.; Pieber, S. M.; Corbin, J. C.; Slowik, J. G.; Brune, W. H.; Baltensperger, U.; Prévôt, A. S. H. Inter-Comparison of Laboratory Smog Chamber and Flow Reactor Systems on Organic Aerosol Yield and Composition. *Atmos. Meas. Tech.* **2015**, 8 (6), 2315–2332.
- (14) Jaffe, D. A.; Wigder, N. L. Ozone Production from Wildfires: A Critical Review. *Atmos. Environ.* **2012**, *51*, 1–10.
- (15) Brown, S. S.; Stutz, J. Nighttime Radical Observations and Chemistry. *Chem. Soc. Rev.* **2012**, *41*, 6405–6447.
- (16) Wayne, R.; Barnes, I.; Biggs, P.; Burrows, J.; Canosa-Mas, C.; Hjorth, J.; Le Bras, G.; Moortgat, G.; Perner, D.; Poulet, G.; Restelli, G.; Sidebottom, H. The Nitrate Radical: Physics, Chemistry, and the Atmosphere. *Atmos. Environ., Part A* **1991**, 25 (1), 1–203.
- (17) Osthoff, H. D.; Roberts, J. M.; Ravishankara, A. R.; Williams, E. J.; Lerner, B. M.; Sommariva, R.; Bates, T. S.; Coffman, D.; Quinn, P. K.; Dibb, J. E.; Stark, H.; Burkholder, J. B.; Talukdar, R. K.; Meagher, J.; Fehsenfeld, F. C.; Brown, S. S. High Levels of Nitryl Chloride in the Polluted Subtropical Marine Boundary Layer. *Nat. Geosci.* **2008**, *1* (5), 324–328.
- (18) Thornton, J. A.; Kercher, J. P.; Riedel, T. P.; Wagner, N. L.; Cozic, J.; Holloway, J. S.; Dubé, W. P.; Wolfe, G. M.; Quinn, P. K.; Middlebrook, A. M.; Alexander, B.; Brown, S. S. A Large Atomic Chlorine Source Inferred from Mid-Continental Reactive Nitrogen Chemistry. *Nature* **2010**, *464* (7286), 271–274.
- (19) Chang, W. L.; Bhave, P. V.; Brown, S. S.; Riemer, N.; Stutz, J.; Dabdub, D. Heterogeneous Atmospheric Chemistry, Ambient Measurements, and Model Calculations of N2O5: A Review. *Aerosol Sci. Technol.* **2011**, *45* (6), 665–695.
- (20) Ahern, A.; Goldberger, L.; Jahl, L.; Thornton, J.; Sullivan, R. C. Production of N2O5 and ClNO2 through Nocturnal Processing of

- Biomass-Burning Aerosol. Environ. Sci. Technol. 2018, 52 (2), 550–559.
- (21) Hatch, L. E.; Luo, W.; Pankow, J. F.; Yokelson, R. J.; Stockwell, C. E.; Barsanti, K. C. Identification and Quantification of Gaseous Organic Compounds Emitted from Biomass Burning Using Two-Dimensional Gas Chromatography—time-of-Flight Mass Spectrometry. *Atmos. Chem. Phys.* **2015**, *15* (4), 1865—1899.
- (22) Stockwell, C. É.; Yokelson, R. J.; Kreidenweis, S. M.; Robinson, A. L.; Demott, P. J.; Sullivan, R. C.; Reardon, J.; Ryan, K. C.; Griffith, D. W. T.; Stevens, L. Trace Gas Emissions from Combustion of Peat, Crop Residue, Domestic Biofuels, Grasses, and Other Fuels: Configuration and Fourier Transform Infrared (FTIR) Component of the Fourth Fire Lab at Missoula Experiment (FLAME-4). Atmos. Chem. Phys. 2014, 14 (18), 9727–9754.
- (23) Selimovic, V.; Yokelson, R. J.; Warneke, C.; Roberts, J. M.; de Gouw, J. A.; Griffith, D. W. T. Aerosol Optical Properties and Trace Gas Emissions from Laboratory-Simulated Western US Wildfires during FIREX. *Atmos. Chem. Phys.* **2018**, *18*, 2929–2948.
- (24) Sekimoto, K.; Koss, A. R.; Gilman, J. B.; Selimovic, V.; Coggon, M. M.; Zarzana, K. J.; Yuan, B.; Lerner, B. M.; Brown, S. S.; Warneke, C.; Yokelson, R. J.; Roberts, J. M.; de Gouw, J. High- and Low-Temperature Pyrolysis Profiles Describe Volatile Organic Compound Emissions from Western US Wildfire Fuels. *Atmos. Chem. Phys. Discuss.* 2018, 1–39.
- (25) Hartikainen, A.; Yli-Pirilä, P.; Tiitta, P.; Leskinen, A.; Kortelainen, M.; Orasche, J.; Schnelle-Kreis, J.; Lehtinen, K.; Zimmermann, R.; Jokiniemi, J.; Sippula, O. Volatile Organic Compounds from Logwood Combustion: Emissions and Transformation under Dark and Photochemical Aging Conditions in a Smog Chamber. *Environ. Sci. Technol.* **2018**, *52*, 4979–4988.
- (26) Atkinson, R.; Arey, J. Gas-Phase Tropospheric Chemistry of Biogenic Volatile Organic Compounds: A Review. *Atmospheric Environment*; Pergamon, 2003; Vol. 37, pp 197–219. DOI: 10.1016/S1352-2310(03)00391-1.
- (27) Atkinson, R.; Arey, J. Atmospheric Degradation of Volatile Organic Compounds. *Chem. Rev.* **2003**, *103* (3), 4605–4638.
- (28) Atkinson, R. Kinetics and Mechanisms of the Gas-Phase Reactions of the NO3 Radical with Organic Compounds. *J. Phys. Chem. Ref. Data* **1991**, *20*, 459–507.
- (29) El Zein, A.; Coeur, C.; Obeid, E.; Lauraguais, A.; Fagniez, T. Reaction Kinetics of Catechol (1,2-Benzenediol) and Guaiacol (2-Methoxyphenol) with Ozone. *J. Phys. Chem. A* **2015**, *119* (26), 6759–6765.
- (30) Martínez, E.; Cabañas, B.; Aranda, A.; Martín, P.; Salgado, S. Absolute Rate Coefficients for the Gas-Phase Reactions of NO3 Radical with a Series of Monoterpenes at T = 298 to 433 K. *J. Atmos. Chem.* 1999, 33 (3), 265–282.
- (31) Kerdouci, J.; Picquet-Varrault, B.; Doussin, J. F. Structure-Activity Relationship for the Gas-Phase Reactions of NO3radical with Organic Compounds: Update and Extension to Aldehydes. *Atmos. Environ.* **2014**, 84 (3), 363–372.
- (32) Grosjean, D.; Williams, E. L. Environmental Persistence of Organic Compounds Estimated from Structure-Reactivity and Linear Free-Energy Relationships. Unsaturated Aliphatics. *Atmos. Environ., Part A* **1992**, 26 (8), 1395–1405.
- (33) Cabañas, B.; Baeza, M. T.; Salgado, S.; Martín, P.; Taccone, R.; Martínez, E. Oxidation of Heterocycles in the Atmosphere: Kinetic Study of Their Reactions with NO3 Radical. *J. Phys. Chem. A* **2004**, *108* (49), 10818–10823.
- (34) Harrison, M. A. J.; Barra, S.; Borghesi, D.; Vione, D.; Arsene, C.; Iulian Olariu, R. Nitrated Phenols in the Atmosphere: A Review. *Atmos. Environ.* **2005**, 39, 231–248.
- (35) Laskin, A.; Laskin, J.; Nizkorodov, S. A. Chemistry of Atmospheric Brown Carbon. *Chem. Rev.* **2015**, *115* (10), 4335–4382.
- (36) Zarzana, K. J.; Min, K. E.; Washenfelder, R. A.; Kaiser, J.; Krawiec-Thayer, M.; Peischl, J.; Neuman, J. A.; Nowak, J. B.; Wagner, N. L.; Dubè, W. P.; St. Clair, J. M.; Wolfe, G. M.; Hanisco, T. F.; Keutsch, F. N.; Ryerson, T. B.; Brown, S. S. Emissions of Glyoxal and Other Carbonyl Compounds from Agricultural Biomass Burning

- Plumes Sampled by Aircraft. *Environ. Sci. Technol.* **2017**, *51* (20), 11761–11770.
- (37) Neuman, J. A.; Trainer, M.; Brown, S. S.; Min, K. E.; Nowak, J. B.; Parrish, D. D.; Peischl, J.; Pollack, I. B.; Roberts, J. M.; Ryerson, T. B.; Veres, P. R. HONO Emission and Production Determined from Airborne Measurements over the Southeast U.S. J. Geophys. Res. 2016, 121 (15), 9237–9250.
- (38) Warneke, C.; Trainer, M.; De Gouw, J. A.; Parrish, D. D.; Fahey, D. W.; Ravishankara, A. R.; Middlebrook, A. M.; Brock, C. A.; Roberts, J. M.; Brown, S. S.; Neuman, J. A.; Lerner, B. M.; Lack, D.; Law, D.; Hübler, G.; Pollack, I.; Sjostedt, S.; Ryerson, T. B.; Gilman, J. B.; Liao, J.; Holloway, J.; Peischl, J.; Nowak, J. B.; Aikin, K. C.; Min, K. E.; Washenfelder, R. A.; Graus, M. G.; Richardson, M.; Markovic, M. Z.; Wagner, N. L.; Welti, A.; Veres, P. R.; Edwards, P.; Schwarz, J. P.; Gordon, T.; Dube, W. P.; McKeen, S. A.; Brioude, J.; Ahmadov, R.; Bougiatioti, A.; Lin, J. J.; Nenes, A.; Wolfe, G. M.; Hanisco, T. F.; Lee, B. H.; Lopez-Hilfiker, F. D.; Thornton, J. A.; Keutsch, F. N.; Kaiser, J.; Mao, J.; Hatch, C. D. Instrumentation and Measurement Strategy for the NOAA SENEX Aircraft Campaign as Part of the Southeast Atmosphere Study 2013. Atmos. Meas. Tech. 2016, 9 (7), 3063–3093.
- (39) Brown, S. S.; Stark, H.; Ravishankara, A. R. Cavity Ring-down Spectroscopy for Atmospheric Trace Gas Detection: Application to the Nitrate Radical (NO3). *Appl. Phys. B: Lasers Opt.* **2002**, *75* (2–3), 173–182.
- (40) Dubé, W. P.; Brown, S. S.; Osthoff, H. D.; Nunley, M. R.; Ciciora, S. J.; Paris, M. W.; McLaughlin, R. J.; Ravishankara, A. R. Aircraft Instrument for Simultaneous, *in Situ* Measurement of NO3 and N2O5 via Pulsed Cavity Ring-down Spectroscopy. *Rev. Sci. Instrum.* 2006, 77 (3), 034101.
- (41) Fuchs, H.; Dubé, W. P.; Ciciora, S. J.; Brown, S. S. Determination of Inlet Transmission and Conversion Efficiencies for in Situ Measurements of the Nocturnal Nitrogen Oxides, NO3, N 2O5 and NO2, via Pulsed Cavity Ring-down Spectroscopy. *Anal. Chem.* 2008, 80 (15), 6010–6017.
- (42) Wild, R. J.; Edwards, P. M.; Dube, W. P.; Baumann, K.; Edgerton, E. S.; Quinn, P. K.; Roberts, J. M.; Rollins, A. W.; Veres, P. R.; Warneke, C.; Williams, E. J.; Yuan, B.; Brown, S. S. A Measurement of Total Reactive Nitrogen, NOy, Together with NO2, NO, and O3 via Cavity Ring-down Spectroscopy. *Environ. Sci. Technol.* 2014, 48 (16), 9609–9615.
- (43) Ryerson, T. B.; Huey, L. G.; Knapp, K.; Neuman, J. A.; Parrish, D. D.; Sueper, D. T.; Fehsenfeld, F. C. Design and Initial Characterization of an Inlet for Gas-Phase NOy Measurements from Aircraf. *J. Geophys. Res. Atmos.* 1999, 104 (D5), 5483–5492.
- (44) Brock, C. A.; Schröder, F.; Kärcher, B.; Petzold, A.; Busen, R.; Fiebig, M. Ultrafine Particle Size Distribution Measured in Aircraft Exhaust Plumes. *J. Geophys. Res.* **2000**, *105* (Vi), 26555–26567.
- (45) Brock, C. A.; Cozic, J.; Bahreini, R.; Froyd, K. D.; Middlebrook, A. M.; McComiskey, A.; Brioude, J.; Cooper, O. R.; Stohl, A.; Aikin, K. C.; De Gouw, J. A.; Fahey, D. W.; Ferrare, R. A.; Gao, R. S.; Gore, W.; Holloway, J. S.; Hübler, G.; Jefferson, A.; Lack, D. A.; Lance, S.; Moore, R. H.; Murphy, D. M.; Nenes, A.; Novelli, P. C.; Nowak, J. B.; Ogren, J. A.; Peischl, J.; Pierce, R. B.; Pilewskie, P.; Quinn, P. K.; Ryerson, T. B.; Schmidt, K. S.; Schwarz, J. P.; Sodemann, H.; Spackman, J. R.; Stark, H.; Thomson, D. S.; Thornberry, T.; Veres, P.; Watts, L. A.; Warneke, C.; Wollny, A. G. Characteristics, Sources, and Transport of Aerosols Measured in Spring 2008 during the Aerosol, Radiation, and Cloud Processes Affecting Arctic Climate (ARCPAC) Project. Atmos. Chem. Phys. 2011, 11 (6), 2423–2453.
- (46) De Gouw, J.; Warneke, C. Measurements of Volatile Organic Compounds in the Earth's Atmosphere Using Proton-Transfer-Reaction Mass Spectrometry. *Mass Spectrometry Reviews*; Wiley-Blackwell, 2007; pp 223–257. DOI: 10.1002/mas.20119.
- (47) Bond, T. C.; Doherty, S. J.; Fahey, D. W.; Forster, P. M.; Berntsen, T.; Deangelo, B. J.; Flanner, M. G.; Ghan, S.; Kärcher, B.; Koch, D.; Kinne, S.; Kondo, Y.; Quinn, P. K.; Sarofim, M. C.; Schultz, M. G.; Schulz, M.; Venkataraman, C.; Zhang, H.; Zhang, S.; Bellouin, N.; Guttikunda, S. K.; Hopke, P. K.; Jacobson, M. Z.; Kaiser, J. W.;

- Klimont, Z.; Lohmann, U.; Schwarz, J. P.; Shindell, D.; Storelvmo, T.; Warren, S. G.; Zender, C. S. Bounding the Role of Black Carbon in the Climate System: A Scientific Assessment. *J. Geophys. Res. Atmos.* **2013**, *118* (11), 5380–5552.
- (48) Min, K.-E.; Washenfelder, R. a.; Dubé, W. P.; Langford, a. O.; Edwards, P. M.; Zarzana, K. J.; Stutz, J.; Lu, K.; Rohrer, F.; Zhang, Y.; Brown, S. S. A Broadband Cavity Enhanced Absorption Spectrometer for Aircraft Measurements of Glyoxal, Methylglyoxal, Nitrous Acid, Nitrogen Dioxide, and Water Vapor. *Atmos. Meas. Tech.* **2016**, 9 (2), 423–440.
- (49) Zheng, W.; Flocke, F. M.; Tyndall, G. S.; Swanson, A.; Orlando, J. J.; Roberts, J. M.; Huey, L. G.; Tanner, D. J. Characterization of a Thermal Decomposition Chemical Ionization Mass Spectrometer for the Measurement of Peroxy Acyl Nitrates (PANs) in the Atmosphere. *Atmos. Chem. Phys.* **2011**, *11* (13), 6529–6547.
- (50) Schwarz, J. P.; Gao, R. S.; Fahey, D. W.; Thomson, D. S.; Watts, L. A.; Wilson, J. C.; Reeves, J. M.; Darbeheshti, M.; Baumgardner, D. G.; Kok, G. L.; Chung, S. H.; Schulz, M.; Hendricks, J.; Lauer, A.; Kärcher, B.; Slowik, J. G.; Rosenlof, K. H.; Thompson, T. L.; Langford, A. O.; Loewenstein, M.; Aikin, K. C. Single-Particle Measurements of Midlatitude Black Carbon and Light-Scattering Aerosols from the Boundary Layer to the Lower Stratosphere. J. Geophys. Res. 2006, 111 (16), 1–15.
- (51) Holloway, J. S.; Jakoubek, R. O.; Parrish, D. D.; Gerbig, C.; Volz-Thomas, A.; Schmitgen, S.; Fried, A.; Wert, B.; Henry, B.; Drummond, J. R. Airborne Intercomparison of Vacuum Ultraviolet Fluorescence and Tunable Diode Laser Absorption Measurements of Tropospheric Carbon Monoxide. *J. Geophys. Res. Atmos.* 2000, 105 (D19), 24251–24261.
- (52) Stockwell, C. E.; Veres, P. R.; Williams, J.; Yokelson, R. J. Characterization of Biomass Burning Emissions from Cooking Fires, Peat, Crop Residue, and Other Fuels with High-Resolution Proton-Transfer-Reaction Time-of-Flight Mass Spectrometry. *Atmos. Chem. Phys.* **2015**, *15* (2), 845–865.
- (53) Kerdouci, J.; Picquet-Varrault, B.; Doussin, J. F. Prediction of Rate Constants for Gas-Phase Reactions of Nitrate Radical with Organic Compounds: A New Structure-Activity Relationship. *ChemPhysChem* **2010**, *11* (18), 3909–3920.
- (54) Edwards, P. M.; Aikin, K. C.; Dube, W. P.; Fry, J. L.; Gilman, J. B.; De Gouw, J. A.; Graus, M. G.; Hanisco, T. F.; Holloway, J.; Hübler, G.; Kaiser, J.; Keutsch, F. N.; Lerner, B. M.; Neuman, J. A.; Parrish, D. D.; Peischl, J.; Pollack, I. B.; Ravishankara, A. R.; Roberts, J. M.; Ryerson, T. B.; Trainer, M.; Veres, P. R.; Wolfe, G. M.; Warneke, C.; Brown, S. S. Transition from High- to Low-NO x Control of Night-Time Oxidation in the Southeastern US. *Nat. Geosci.* 2017, 10 (7), 490–495.
- (55) Wolfe, G. M.; Marvin, M. R.; Roberts, S. J.; Travis, K. R.; Liao, J. The Framework for 0-D Atmospheric Modeling (F0AM) v3.1. *Geosci. Model Dev.* **2016**, *9* (9), 3309–3319.
- (56) Jenkin, M. E.; Saunders, S. M.; Pilling, M. J. The Tropospheric Degradation of Volatile Organic Compounds: A Protocol for Mechanism Development. *Atmos. Environ.* **1997**, *31* (1), 81–104.
- (57) Jenkin, M. E.; Saunders, S. M.; Wagner, V.; Pilling, M. J. Protocol for the Development of the Master Chemical Mechanism, MCM v3 (Part B): Tropospheric Degradation of Aromatic Volatile Organic Compounds. *Atmos. Chem. Phys.* **2003**, 3 (1), 181–193.
- (\$8) Bloss, C.; Wagner, V.; Jenkin, M. E.; Volkamer, R.; Bloss, W. J.; Lee, J. D.; Heard, D. E.; Wirtz, K.; Martin-Reviejo, M.; Rea, G.; Wenger, J. C.; Pilling, M. J. Development of a Detailed Chemical Mechanism (MCMv3.1) for the Atmospheric Oxidation of Aromatic Hydrocarbons. *Atmos. Chem. Phys.* **2005**, *5* (3), 641–664.
- (59) Jenkin, M. E.; Wyche, K. P.; Evans, C. J.; Carr, T.; Monks, P. S.; Alfarra, M. R.; Barley, M. H.; McFiggans, G. B.; Young, J. C.; Rickard, A. R. Development and Chamber Evaluation of the MCM v3.2 Degradation Scheme for β -Caryophyllene. *Atmos. Chem. Phys.* **2012**, 12 (11), 5275–5308.
- (60) Jenkin, M. E.; Young, J. C.; Rickard, A. R. The MCM v3.3.1 Degradation Scheme for Isoprene. *Atmos. Chem. Phys.* **2015**, *15* (20), 11433–11459.

- (61) Yang, B.; Zhang, H.; Wang, Y.; Zhang, P.; Shu, J.; Sun, W.; Ma, P. Experimental and Theoretical Studies on Gas-Phase Reactions of NO3 Radicals with Three Methoxyphenols: Guaiacol, Creosol, and Syringol. *Atmos. Environ.* **2016**, *125* (3), 243–251.
- (62) Tapia, A.; Villanueva, F.; Salgado, M. S.; Cabañas, B.; Martínez, E.; Martín, P. Atmospheric Degradation of 3-Methylfuran: Kinetic and Products Study. *Atmos. Chem. Phys.* **2011**, *11* (7), 3227–3241.
- (63) Lauraguais, A.; El Zein, A.; Coeur, C.; Obeid, E.; Cassez, A.; Rayez, M.-T.; Rayez, J.-C. Kinetic Study of the Gas-Phase Reactions of Nitrate Radicals with Methoxyphenol Compounds: Experimental and Theoretical Approaches. *J. Phys. Chem. A* **2016**, *120* (17), 2691–2699.
- (64) CropScape NASS CDL Program https://nassgeodata.gmu.edu/CropScape/ (accessed Aug 6, 2018).
- (65) Brown, S. Absorption Spectroscopy in High Finesse Cavities for Atmospheric Studies. *Chem. Rev.* **2003**, *103*, 5219.
- (66) Brown, S. S.; Stark, H.; Ravishankara, A. R. Applicability of the Steady State Approximation to the Interpretation of Atmospheric Observations of NO 3 and N 2 O 5. *J. Geophys. Res.* **2003**, *108* (D17), 4539.
- (67) Olariu, R. I.; Bejan, I.; Barnes, I.; Klotz, B.; Becker, K. H.; Wirtz, K. Rate Coefficients for the Gas-Phase Reaction of NO3 Radicals with Selected Dihydroxybenzenes. *Int. J. Chem. Kinet.* **2004**, 36 (11), 577–583.
- (68) Finewax, Z.; De Gouw, J. A.; Ziemann, P. J. Identification and Quantification of 4-Nitrocatechol Formed from OH and NO3R-adical-Initiated Reactions of Catechol in Air in the Presence of NOx: Implications for Secondary Organic Aerosol Formation from Biomass Burning. *Environ. Sci. Technol.* **2018**, 52 (4), 1981–1989.
- (69) Desyaterik, Y.; Sun, Y.; Shen, X.; Lee, T.; Wang, X.; Wang, T.; Collett, J. L. Speciation of "Brown" Carbon in Cloud Water Impacted by Agricultural Biomass Burning in Eastern China. *J. Geophys. Res. Atmos.* **2013**, *118* (13), 7389–7399.
- (70) Claeys, M.; Vermeylen, R.; Yasmeen, F.; Gómez-González, Y.; Chi, X.; Maenhaut, W.; Mészáros, T.; Salma, I. Chemical Characterisation of Humic-like Substances from Urban, Rural and Tropical Biomass Burning Environments Using Liquid Chromatography with UV/Vis Photodiode Array Detection and Electrospray Ionisation Mass Spectrometry. *Environ. Chem.* **2012**, *9* (3), 273–284.
- (71) Lin, P.; Bluvshtein, N.; Rudich, Y.; Nizkorodov, S. A.; Laskin, J.; Laskin, A. Molecular Chemistry of Atmospheric Brown Carbon Inferred from a Nationwide Biomass Burning Event. *Environ. Sci. Technol.* **2017**, *51* (20), 11561–11570.
- (72) Iinuma, Y.; Böge, O.; Gräfe, R.; Herrmann, H. Methyl-Nitrocatechols: Atmospheric Tracer Compounds for Biomass Burning Secondary Organic Aerosols. *Environ. Sci. Technol.* **2010**, 44, 8453–8459.
- (73) Gaston, C. J.; Lopez-Hilfiker, F. D.; Whybrew, L. E.; Hadley, O.; McNair, F.; Gao, H.; Jaffe, D. A.; Thornton, J. A. Online Molecular Characterization of Fine Particulate Matter in Port Angeles, WA: Evidence for a Major Impact from Residential Wood Smoke. *Atmos. Environ.* **2016**, *138*, 99–107.
- (74) Mohr, C.; Lopez-Hilfiker, F. D.; Zotter, P.; Prévoît, A. S. H.; Xu, L.; Ng, N. L.; Herndon, S. C.; Williams, L. R.; Franklin, J. P.; Zahniser, M. S.; Worsnop, D. R.; Knighton, W. B.; Aiken, A. C.; Gorkowski, K. J.; Dubey, M. K.; Allan, J. D.; Thornton, J. A. Contribution of Nitrated Phenols to Wood Burning Brown Carbon Light Absorption in Detling, United Kingdom during Winter Time. *Environ. Sci. Technol.* **2013**, 47 (12), 6316–6324.
- (75) Xie, M.; Chen, X.; Hays, M. D.; Lewandowski, M.; Offenberg, J.; Kleindienst, T. E.; Holder, A. L. Light Absorption of Secondary Organic Aerosol: Composition and Contribution of Nitroaromatic Compounds. *Environ. Sci. Technol.* **2017**, *51* (20), 11607–11616.
- (76) Hinrichs, R. Z.; Buczek, P.; Trivedi, J. J. Solar Absorption by Aerosol-Bound Nitrophenols Compared to Aqueous and Gaseous Nitrophenols. *Environ. Sci. Technol.* **2016**, *50* (11), 5661–5667.
- (77) Odum, J. R.; Hoffmann, T.; Bowman, F.; Collins, D.; Flagan, R. C.; Seinfeld, J. H. Gas Particle Partitioning and Secondary Organic Aerosol Yields. *Environ. Sci. Technol.* **1996**, 30 (8), 2580–2585.