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Key Points:

- Spatial gradients are observed in dilution-corrected mixing ratios of nitrous acid, ozone, and reactive organics in fresh wildfire plumes
- Gradients evolve with time of day, as intensity of chemical aging varies relative to plume transport and mixing
- Gradients should be considered when comparing plume models with measurements and estimating fire emissions from in situ observations

Abstract

Wildfire emissions affect downwind air quality and human health. Predictions of these impacts using models are limited by uncertainties in emissions and chemical evolution of smoke plumes. Using high-time-resolution aircraft measurements, we illustrate spatial variations that can exist horizontally and vertically within a plume due to differences in the photochemical environment. Dilution-corrected mixing ratio gradients were observed for reactive compounds and their oxidation products, such as nitrous acid, catechol, and ozone, likely due to faster photochemistry in optically-thinner plume edges relative to darker plume cores. Mixing ratio gradients in midday plumes, driven by j_{HONO} gradients, are often steepest in the freshest transects, and become flatter with chemical aging. Gradients in plumes emitted close to sunset are characterized by titration of O_3 in the plume and little to no gradient formation. We show how gradients can lead to underestimated emission ratios for reactive compounds and overestimated emission ratios for oxidation products.

Plain Language Summary

Wildfire emissions have large impacts on air quality and health in downwind communities. Previous research has shown that chemical reactions in fire plumes

can be remarkably fast, which modifies the impact the smoke has downwind. In this work, we show how this chemistry happens faster on the edges versus the core of plumes emitted during midday. For plumes emitted near or after sunset, oxidation chemistry generally slows and remains nearly uniform across the plume. These variations in plume chemistry will impact how plumes are modeled, and how well we can predict downwind air quality impacts.

1. Introduction

Wildfire emissions are becoming increasingly important drivers of degraded air quality, especially in the Western U.S. (McClure & Jaffe, 2018; O’Dell et al., 2019). Wildfire season is growing longer and fires are growing larger due to several factors, including climate change, a legacy of fire suppression, and an increasing wildland-urban interface (Abatzoglou & Williams, 2016; Balch et al., 2017; Dennison et al., 2014; Higuera et al., 2021; Higuera & Abatzoglou, 2021; Radeloff et al., 2018; Westerling, 2016). A better understanding of the composition and evolution of wildfire emissions is needed to predict the impacts on downwind communities, atmospheric dynamics, and climate.

In aircraft studies of wildfires and other point sources, emissions are typically evaluated by sampling plume composition in horizontal transects of the plume perpendicular to the wind direction, where each transect corresponds to a similar physical plume age. A regional background abundance from outside the plume is subtracted from the average crosswind transect abundance to determine a normalized excess mixing ratio (NEMR) for each transect (e.g., Akagi et al., 2012; Liu et al., 2017; Palm et al., 2020; Yokelson et al., 2009). The NEMR values in the youngest parts of the plume are often used as estimates of the emission ratio (ER), which is meant to represent the relative emission flux of a species from the fire prior to any substantial physical or chemical changes with aging. ERs and related emission factors (EF; emission per mass of fuel burned) are essential components of biomass burning emissions inventories used in air quality and earth system models (e.g., Andreae, 2019; van der Werf et al., 2017; Wiedinmyer et al., 2011; Yokelson et al., 1999).

Spatial variations in plume chemistry can make NEMRs and ERs dependent upon which portion of the crosswind transects are used in the calculation (e.g., entire transect versus plume core), especially for chemically reactive components. Spatial variation in wildfire plume chemistry has only rarely been investigated in detail across plume transects (Garofalo et al., 2019; Hodshire et al., 2021; Juncosa Calahorrano et al., 2021; Peng et al., 2020). Furthermore, vertical gradients in plume chemistry have not been investigated to our knowledge. We use high-time-resolution measurements to investigate horizontal and vertical spatial variation in wildfire plume chemistry. We show that gradients can be anywhere from steep to nonexistent depending on factors such as physical age or time of day at emission and sampling. Impacts on NEMR calculations are investigated for all crosswind transects sampled during a recent flight campaign. This work shows the importance of considering spatial gradients in oxidation chemistry of fresh wildfire plumes.

1. Materials and Methods

(a) **WE-CAN campaign**

The Western Wildfire Experiment for Cloud Chemistry, Aerosol Absorption, and Nitrogen (WE-CAN) field campaign took place in the Western U.S. from July–September 2018. Measurements were taken using a suite of instruments installed aboard a C-130 aircraft operated by the National Center for Atmospheric Research (NCAR) and the National Science Foundation (NSF). The C-130 sampled nearly two dozen individual wildfire plumes, targeting fresh emissions and aging up to several hours. Special focus is given to measurements from the Taylor Creek Fire and the South Sugarloaf Fire smoke plumes.

1. Plume measurements

The main instrument used in this analysis was an iodide-adduct high-resolution time-of-flight mass spectrometer (I⁻ CIMS), measuring a suite of gas-phase oxidized organic and inorganic compounds. The I⁻ CIMS operation has been described in detail in Palm et al. (2020). More details of the I⁻ CIMS and other instruments used in this analysis can be found in Sect. S1. For this analysis, data from all instruments were smoothed by a 5 s average. Plumes were defined as any data where the enhancement of CO above regional background was greater than 250 ppbv.

NEMR values across transects were calculated by taking the ratio of background-subtracted abundances of each compound to background-subtracted CO using 5 s averaged data. See Sect. S1 for more details of background subtraction.

1. Results and Discussion

(a) Taylor Creek fire: narrow plume with high [HONO]

i. Gradients in reactive compound NEMRs

Early on 30 July 2018, the Taylor Creek fire in southern Oregon emitted smoke into a smoke-filled boundary layer with other nearby fires. Soon after the research aircraft arrived in the area, the fire intensity and environmental conditions supported the injection of a fresh plume above the boundary layer (3.3–3.7 km above sea level; ASL) in the relatively clean free troposphere (see pictures in Fig. S1). The plume was narrow (~3–5 km across) with a dense core tapering on either side. The plume was first sampled shortly (~10 min) after it was observed to be injecting into the free troposphere, with an estimated physical age of about 21 min after emission. The aircraft was able to continually resample the leading edge of the plume as it aged from 15:45–18:00 local time (LT), providing a near-Lagrangian experiment.

Crosswind gradients in the Taylor Creek plume can be seen in the map of NEMRs in Fig. 1. With the effects of dilution removed by normalizing to excess CO, the strong horizontal crosswind gradients observed for this suite of reactive compounds and oxidation products were a result of variable rates of chemical loss/production across the plume. For the fastest reacting compounds such as

$C_6H_6O_2$ (likely catechol), HONO (nitrous acid), and $C_7H_8O_2$ (likely methyl catechol, shown in Fig. S2), high mixing ratios were observed only in the core of the first several transects closest to the fire. At these locations, the plume core was at its darkest, as confirmed by j_{HONO} . In the earliest two transects, the dilution-corrected production rate of OH from HONO photolysis was faster on the plume edges relative to the core (Fig. S2). However, even on the edges of the first several transects, HONO was substantially photolyzed and compounds such as $C_6H_6O_2$ (lifetime of 33 min at $[OH] = 5 \times 10^6$ molecules cm^{-3} ; Olariu et al., 2000) were already mostly reacted away. Slower reacting compounds such as $C_6H_6O_1$ (likely phenol, lifetime of 2.1 h at $[OH] = 5 \times 10^6$ molecules cm^{-3} ; Atkinson et al., 1989) were depleted more gradually with downwind transport, and smaller gradients were observed in the crosswind transects. For compounds that were not reactive on this time scale of several hours of downwind transport, gradients were not observed (or expected) in either the crosswind or downwind directions (e.g., HCN shown in Fig. S2).

For compounds produced from oxidation chemistry, reversed gradients were observed where the compounds were enhanced relative to CO on the plume edges versus the core. This group includes organic oxidation products such as $C_3H_4O_3$ (pyruvic acid and other isomers, as an example of a variety of small organic oxidation products) and ozone (O_3), as shown in Fig. 1. A general trend of increasing NEMRs with increasing age downwind was observed for this subset of compounds. In the Taylor Creek plume, near-field O_3 formation was likely influenced by rapid OH chemistry from HONO photolysis during the mid-day sampling. The Taylor Creek plume contained one of the highest measured HONO ERs of any fire during the campaign (Peng et al., 2020).

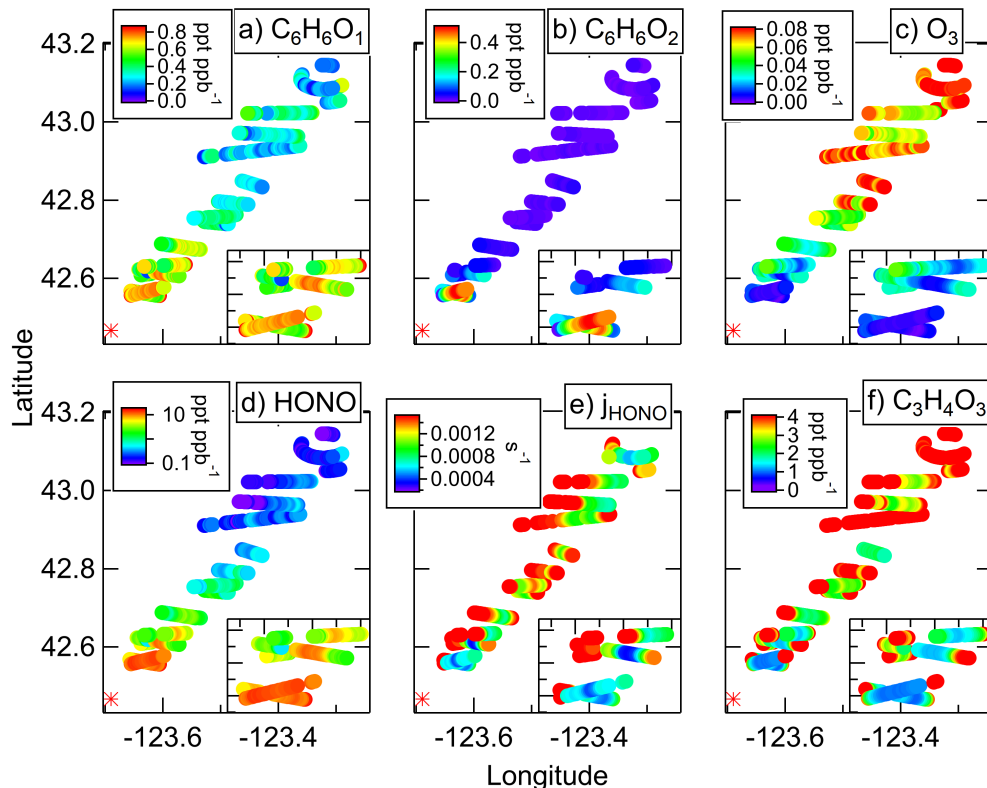


Figure 1. Plume transects of the Taylor Creek fire, showing spatial variations in NEMRs of a) $\text{C}_6\text{H}_6\text{O}_1$, b) $\text{C}_6\text{H}_6\text{O}_2$, c) O_3 , d) HONO , e) j_{HONO} , and f) $\text{C}_3\text{H}_4\text{O}_3$. The emission source is marked with an asterisk. Insets show the first five transects in greater detail.

1. Inferred gradients of average OH concentrations in fresh plumes

The gradients observed in the NEMRs of reactive compounds and their oxidation products indicate that the rate of oxidation, and therefore oxidant concentrations, have gradients as well. In the Taylor Creek plume, the large source of OH from HONO photolysis in this relatively bright midday plume meant that OH was likely the dominant oxidant for most compounds. Here, we use the gradients of reactive compounds to infer spatial variations in photochemical aging due to OH oxidation at different locations within the plume.

The average OH radical concentration ($[\text{OH}]_{\text{avg}}$) experienced by air sampled at any point within the plume since emission can be calculated using Eq. 1, derived from de Gouw et al. (2005) and Roberts et al. (1984). We use the known reaction rates of compounds A and B with OH ($k_{\text{OH}+\text{A}}$ and $k_{\text{OH}+\text{B}}$; see

Table S1 for all rates used), the initial emission ratio of the compounds (at $t=0$), and the estimated physical time since emission (Δt).

$$[\text{OH}]_{\text{avg}} = \frac{1}{t(k_{\text{OH}+\text{A}} - k_{\text{OH}+\text{B}})} \times \left(\ln \left(\frac{[\text{A}]}{[\text{B}]} \right) \Big|_{t=0} - \ln \left(\frac{[\text{A}]}{[\text{B}]} \right) \right) \quad (1)$$

This equation can be rearranged to give OH exposure as well ($\text{OH}_{\text{exp}} = [\text{OH}]_{\text{avg}} \times \Delta t$). Equation 1 is valid when OH is the only relevant oxidant for the chosen compounds during Δt . Compounds A and B are chosen such that they react with OH at different rates (and aren't produced as oxidation products) and their ratio of emissions can be estimated. When $k_{\text{OH}+\text{B}}$ is slow enough that B does not appreciably react during Δt (as is the case for CO), then the sole purpose of compound B is to account for dilution, and Eq. 1 can be simplified to the exponential decay of dilution corrected compound A. The methodology is further described in Sect. S2.

We used $\text{C}_6\text{H}_6\text{O}_2$ and CO as compounds A and B in Eq. 1 to estimate $[\text{OH}]_{\text{avg}}$ experienced by the plume between emission and the point of sampling across each transect, as shown in Fig. 2. For $\text{C}_6\text{H}_6\text{O}_2$, the nitrate radical (NO_3) is also a relevant oxidant. Since $k_{\text{OH}+\text{C}_6\text{H}_6\text{O}_2}$ and $k_{\text{NO}_3+\text{C}_6\text{H}_6\text{O}_2}$ are nearly equal (1.0×10^{-10} and $9.9 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, respectively; Olariu et al., 2000, 2004), Eq. 1 provides an estimate of $[\text{OH}+\text{NO}_3]_{\text{avg}}$. Box modeling (Sect. S2) suggests approximately 25% of $\text{C}_6\text{H}_6\text{O}_2$ reacts with NO_3 (and negligible reaction with O_3) in the Taylor Creek Fire plume. Thus, we multiplied the result of Eq. 1 by 0.75 to estimate $[\text{OH}]_{\text{avg}}$ whenever using $\text{C}_6\text{H}_6\text{O}_2$.

In the several transects closest to the fire with estimated physical age of about 20 min, steep cross-plume gradients were observed for $[\text{OH}]_{\text{avg}}$ in Fig. 2. On the plume edges where HONO photolysis was faster, we inferred $[\text{OH}]_{\text{avg}}$ on the order of $1.0 \times 10^7 \text{ molecules cm}^{-3}$, while the plume core was much lower at approximately $4 \times 10^6 \text{ molecules cm}^{-3}$. In transects further downwind, with physical ages > 45 min, the HONO in the center of the transects had also photolyzed, and the $[\text{OH}]_{\text{avg}}$ inferred in the plume center was essentially the same as that of the plume edges near $1.0 \times 10^7 \text{ molecules cm}^{-3}$. The inferred $[\text{OH}]_{\text{avg}}$ (and the gradients of other compounds in Fig. 1) was essentially flat across each transect as the plume aged further, decreasing with increasing est. physical age, from $1.0 \times 10^7 \text{ molec cm}^{-3}$ closer to the fire to $4 \times 10^6 \text{ molec cm}^{-3}$ after two hours of physical aging. These values are the $[\text{OH}]_{\text{avg}}$ between emission and sampling, so the actual [OH] in the latter part of this plume was lower. As shown in Fig. S4, OH_{exp} increased rapidly with steep cross-plume gradients over the first several transects, and then increased gradually without cross-plume gradients beyond that age.

The choice of initial ratio of $\text{C}_6\text{H}_6\text{O}_2$ to CO is a potential source of uncertainty. For the calculations above, we used an initial ratio of 1. A sensitivity analysis, discussed in Sect. S3, found that the gradients in $[\text{OH}]_{\text{avg}}$ were robust for different initial ratios. Analysis of other compound ratios (Sect. S3) illustrates how crosswind $[\text{OH}]_{\text{avg}}$ gradients are only quantifiable because $\text{C}_6\text{H}_6\text{O}_2$ reacts

fast enough to form measureable crosswind gradients, while compounds such as $C_6H_6O_1$ and C_7H_8 (toluene) are not as useful.

Since the OH source from HONO in this plume was one of the largest measured during WE-CAN, our estimates likely fall near the upper end of [OH] in fresh plumes from fires in this region. Larger and darker plumes, or plumes emitted near or after sunset, may have stronger influence from other oxidants such as NO_3 or O_3 . However, crosswind gradients were observed in multiple wildfire plumes sampled during WE-CAN, as shown for the South Sugarloaf fire in the next section and for the broader WE-CAN measurements in Sect. 3.3.

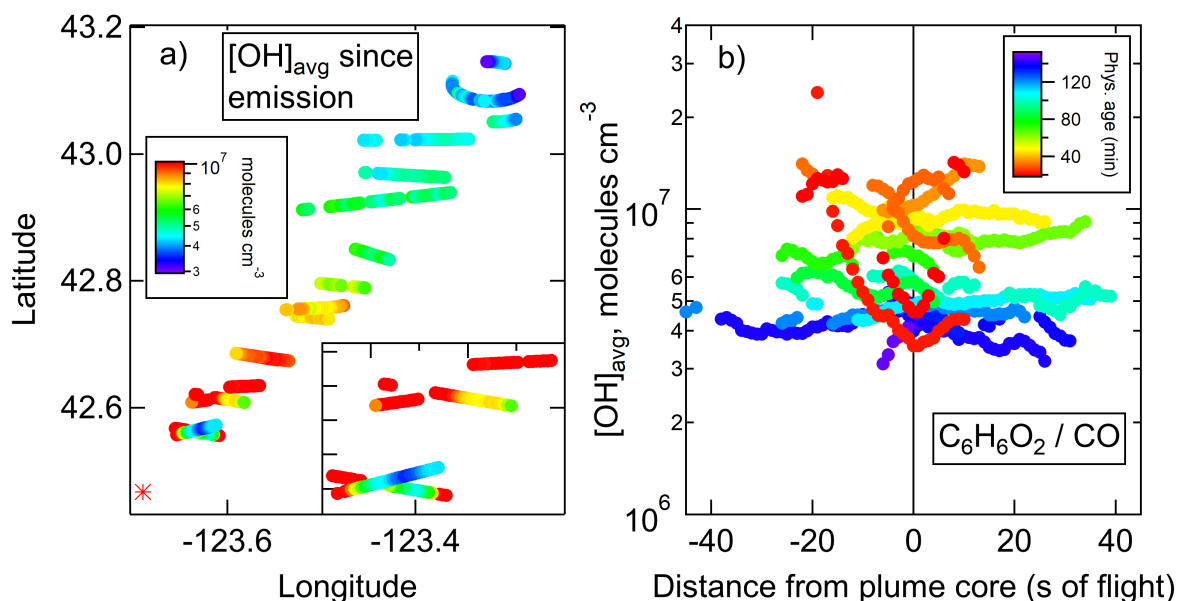


Figure 2. In the Taylor Creek Fire plume, a) $[OH]_{avg}$ between emission source (marked with an asterisk) and each sampled location, and b) $[OH]_{avg}$ between emission source and each sampled location versus distance from plume core defined as maximum [CO] in each transect. Data are colored by estimated physical age.

1. South Sugarloaf Fire: vertical resolution of a wide plume

The South Sugarloaf Fire plume, sampled in northern Nevada near the Idaho border on 26 August, 2018, differed from the Taylor Creek plume in several ways. The fire was active over a larger area, leading to a plume approaching 100 km wide in some locations. Moreover, the South Sugarloaf plume remained in the boundary layer and formed a plume that was more than one km in height from roughly 3.6–4.6 km ASL, which made it possible to sample vertically-stacked crosswind transects to investigate vertical NEMR gradients (see Fig. S7). The maximum HONO mixing ratios were not as high as Taylor Creek, but were more

in line with other fires sampled during WE-CAN (Peng et al., 2020). Also, the HONO photolysis rates were generally slower, especially in the darker plume core, in part because this plume was sampled when clouds were present aloft. The clouds were not interacting with the smoke but led to slower photolysis rates within and around the plume. While the Taylor Creek plume was sampled only during midafternoon, the South Sugarloaf fire was sampled from midday through nearly sunset (14:00-20:00 LT) due to the closer proximity to the aircraft base of operations. This more extended sampling allowed an investigation of how vertical and horizontal crosswind gradients evolved as day transitioned to night.

Dilution-corrected gradients were observed between the cores and edges of vertically stacked crosswind transects in the South Sugarloaf fire, four of which are shown in Fig. 3. These transects were sampled during midday (1440-1540 LT) at an estimated physical age of 76 min. Compounds such as $\text{C}_6\text{H}_6\text{O}_1$, $\text{C}_6\text{H}_6\text{O}_2$, and HONO were depleted on the edges of each transect, while O_3 and $\text{C}_3\text{H}_4\text{O}_3$ were enhanced. Gradients can be observed by plotting NEMRs as a function of excess CO, where the densest part of the plume has the highest excess CO. An example is shown in Fig. S8 for a fresh transect. Vertically stacked measurements of HCN, CO, and calculated $[\text{OH}]_{\text{avg}}$ are shown in Fig. S9 for comparison. $[\text{OH}]_{\text{avg}}$ is again calculated from $\text{C}_6\text{H}_6\text{O}_2$ and CO, shown for comparison with Fig. 2 though it may be less reliable due to a larger possible influence from non-OH oxidation chemistry in the darker plume conditions.

NEMR gradients in the vertical are observed as well. HONO was most depleted in the top transect and along the edges, owing to the higher j_{HONO} values in those locations. The highest HONO mixing ratios were found not in the lowest altitude transect as j_{HONO} might suggest, but in the upper middle transect where both j_{HONO} was low and $[\text{CO}]$ was high. Either the smoke in the lowest crosswind transects experienced higher j_{HONO} values along their trajectory between emission and being sampled, or the HONO emission factor was higher for smoke that rose to a higher altitude, consistent with more intense combustion conditions being a source of HONO (Peng et al., 2020). While the narrow Taylor Creek plume might reasonably be assumed to have a single emission factor for each compound, the much broader South Sugarloaf Fire likely resulted from a more diverse mix of emission processes that may not have been well-mixed during transport. Some variation in dilution corrected HCN was observed, though generally it remained constant across each transect (Fig. S9). Reactive organic compounds such as $\text{C}_6\text{H}_6\text{O}_1$ and $\text{C}_6\text{H}_6\text{O}_2$ followed the trend of HONO of being most concentrated in the core of the plume with highest $[\text{CO}]$. These compounds were depleted above, below, and to the sides of that core, while O_3 was enhanced in those same locations relative to the core. $\text{C}_3\text{H}_4\text{O}_3$ was enhanced more towards the lower altitudes and on the edges.

Several other vertically stacked transects of the South Sugarloaf fire plume were conducted at various physical ages and times of day, which further illustrate the variety of possible gradients. These examples, discussed in Sect. S4, include two sets of transects sampled at the same estimated physical age of 47 min. One

set was sampled during midday and captured strong vertical gradients in the top-most part of the plume (Fig. S10), while the other set was sampled near sunset and showed conditions with O_3 depletion and the lack of any gradients due to slow HONO photolysis (Fig. S11). These examples highlight the need to consider the time of day when wildfire emissions occur.

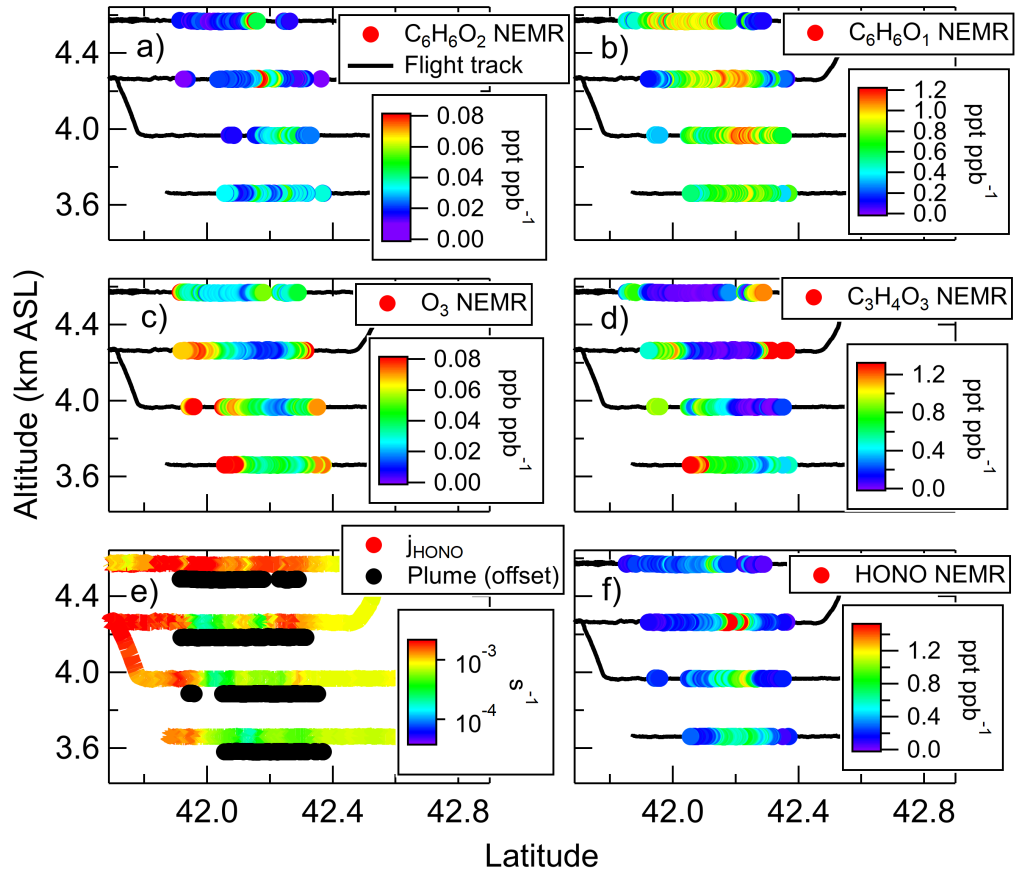


Figure 3. Measurements in four vertically stacked transects of the South Sugarloaf fire, sampled at an average est. physical age of 76 min from 1450-1550 LT. Gradients are observed in both horizontal and vertical directions. The plume center is defined by enhanced a) $C_6H_6O_2$, b) $C_6H_6O_1$, and f) HONO. c) O_3 and d) $C_3H_4O_3$ are enhanced on the edges, top, and bottom of the plume. Measured e) j_{HONO} is shown along the entire flight track, with the plume location

indicated with offset black dots for comparison.

1. Gradient strength depends on time of day at emission and sampling

The case studies above illustrate how observed NEMR gradients are dependent on the specific photochemical conditions experienced within each plume. To investigate the representativeness of those examples, we performed an analysis of gradients observed in all crosswind transects in coherent plumes (at least three physical ages sampled) sampled during the WE-CAN field campaign with estimated physical ages up to six hours. The gradient magnitude was defined as the difference between the NEMR of the plume core (defined as where [CO] was above the 90th percentile in each crosswind transect) versus the transect average NEMR. Since the above examples suggested that gradients are more tied to the extent of chemical aging rather than physical age, this analysis used the ratio of estimated physical age to the lifetime of HONO against photolysis in the plume core ($\tau_{\text{HONO,core}}$), calculated as the inverse of the measured $j_{\text{HONO,core}}$ in each transect. Values less than one indicate chemically fresh plumes where HONO largely has not yet been photolyzed, while larger values indicate more chemically aged plumes. This calculation does not take into account the history of $j_{\text{HONO,core}}$ between emission and measurement (generally increasing with dilution while decreasing as the sun sets), but still provides a reasonable estimate of relative chemical age for these plumes in the first several hours after emission.

In Fig. 4, we show the gradient magnitude as a function of this chemical age metric for several compounds, with data also colored by the solar zenith angle (SZA) at the time of emission. Broadly, fast reacting compounds like $\text{C}_6\text{H}_6\text{O}_2$, HONO, and $\text{C}_7\text{H}_8\text{O}_2$ (Fig. S12) showed positive gradients (depleted on edges versus core) in chemically fresh plumes and near zero gradients in aged plumes where the compounds were depleted throughout the plumes. Oxidation products like O_3 and $\text{C}_3\text{H}_4\text{O}_3$ tended to have negative gradients (enhanced on edges versus core). Compounds that were reactive but not enough to be fully depleted in the first six hours of physical aging, like $\text{C}_6\text{H}_6\text{O}_1$, showed positive gradients across the range of chemical ages, while compounds that were unreactive on these time scales like HCN showed no gradients (as expected; Fig. S12).

Focusing on the chemically fresh plumes where the estimated physical age was less than $\tau_{\text{HONO,core}}$, the gradient strengths depended on SZA at emission. The plumes emitted closest to midday with the lowest SZA showed the strongest gradients, positive for reactive compounds and negative for oxidation products. When plumes were emitted late in the day at higher SZA, little to no gradients were observed for all compound types.

Trends are also observed when comparing gradient magnitudes to absolute NEMR magnitudes, as shown in Fig. S13 as a function of physical age. For the fastest reacting compounds with photolysis or reaction lifetimes on the order of 30 min or less (e.g., $\text{C}_6\text{H}_6\text{O}_2$, HONO, and $\text{C}_7\text{H}_8\text{O}_2$), the core NEMR was up to 50% higher than the transect average NEMR in the chemically freshest transects. The core NEMR values in the chemically freshest plumes will most

closely represent the true ER values. For compounds that react or form on the time scale of one to several hours (e.g., $\text{C}_6\text{H}_6\text{O}_2$, $\text{C}_3\text{H}_4\text{O}_3$, O_3 , etc.), the core NEMR was typically up to 20% higher than the transect average and the gradients lasted out to longer physical ages.

We expect there are other fast reacting or fast forming compounds that are not measureable in this campaign because, e.g., the Γ^- CIMS cannot separate isomers with different gradients, or an oxidation product is formed that rapidly partitions to the unspiciated particle phase.

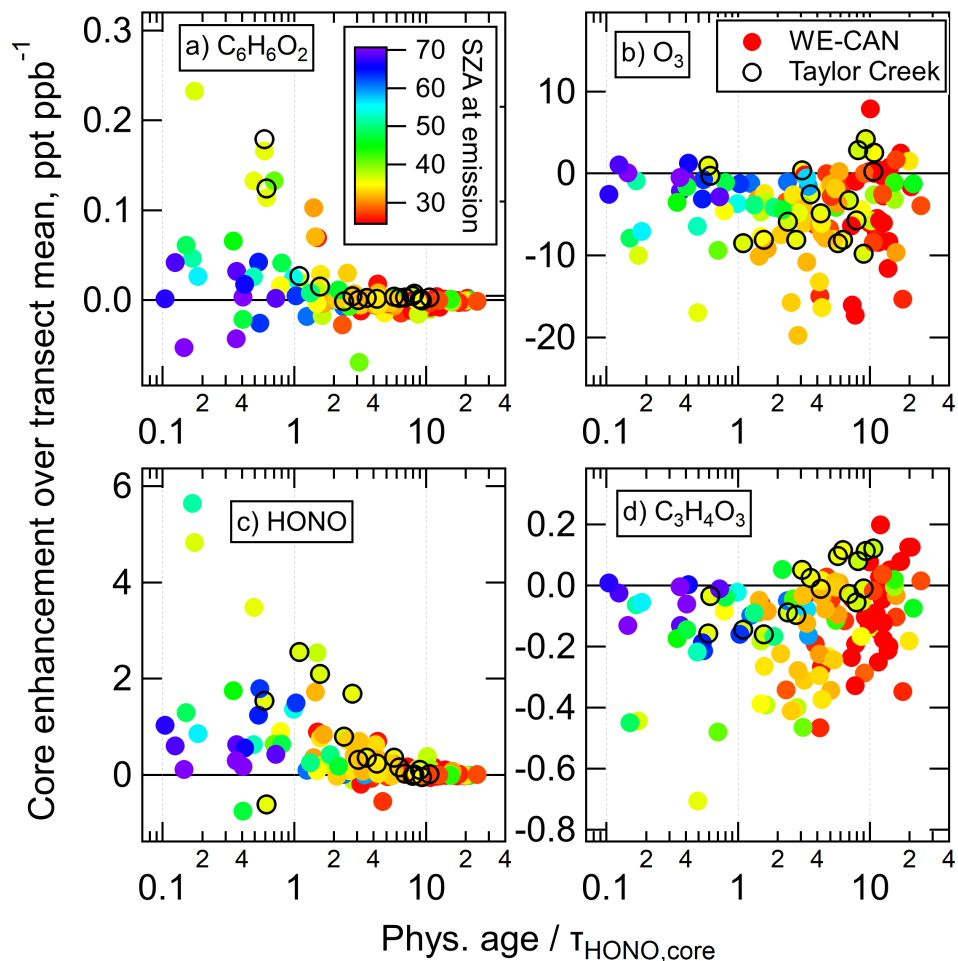


Figure 4. Enhancements of NEMRs in plume cores (average where $[\text{CO}] > 90^{\text{th}}$

percentile) relative to transect mean NEMRs for all WE-CAN transects, for a) $\text{C}_6\text{H}_6\text{O}_2$, b) O_3 , c) HONO, and d) $\text{C}_3\text{H}_4\text{O}_3$. The Taylor Creek plume transects are highlighted.

1. Conclusions

By sampling crosswind transects of wildfire plumes at different physical ages and altitudes with high spatial resolution, spatial variations in plume oxidation chemistry were observed during the WE-CAN aircraft campaign. NEMR gradients were strongest for the most reactive compounds such as HONO and $\text{C}_6\text{H}_6\text{O}_2$ in the chemically freshest plume transects during midday. In the South Sugarloaf fire, vertical gradients were also observed around the plume core. Gradients appeared to be driven by variations in j_{HONO} on diffuse plume edges or tops versus concentrated plume cores during midday plumes. During late evening plumes (and likely nighttime plumes) when the photolysis lifetime of HONO became long even on plume edges, gradients were no longer observed as the OH oxidation chemistry slowed down. This suggests that the fate of wildfire emissions may be dramatically different depending on the time of day at emission.

These measurements indicate that averaging entire transects into single NEMR values may lead to up to 50% underestimated ERs for the most reactive compounds (smaller amounts for less reactive compounds), and overestimated ERs for oxidation products in fresh wildfire plumes. Using screening methods such as using the top percentile of data may be more accurate for some applications, but should be combined with other information about the chemical age of the smoke. Plume gradients should also be considered when comparing *in situ* measurements with 0D plume models.

Acknowledgments and Data

B.B.P., Q.P., and J.A.T. were supported by the US National Science Foundation (NSF; AGS-1652688) and National Oceanic and Atmospheric Administration (NOAA; NA17OAR4310012). This research was supported by the U.S. NSF (AGS-1650786 and AGS-1650275). WE-CAN data (merge R4) is available at <https://www-air.larc.nasa.gov/cgi-bin/ArcView/firexaq?MERGE=1>. We thank everyone involved in planning and operations during WE-CAN. This material is based upon work supported by the National Center for Atmospheric Research, which is a major facility sponsored by the NSF under Cooperative Agreement No. 1852977. The data were collected using NSF’s Lower Atmosphere Observing Facilities, which are managed and operated by NCAR’s Earth Observing Laboratory. The operational and scientific support from NCAR’s Earth Observing Laboratory and Research Aircraft Facility is gratefully acknowledged.

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