



*Supplement of*

## **Photochemical evolution of the 2013 California Rim Fire: synergistic impacts of reactive hydrocarbons and enhanced oxidants**

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### **Text S1. Adjustment of Lab-Reported ERs**

As noted in Sect. 2.4 of the main text, we adjust the emission ratios of acrolein and biacetyl downward by factors of 2.3 and 10, respectively, relative to the values reported by Koss et al. (2018). Here we provide some justification for these modifications.

For acrolein, instrument inter-comparisons during and after FIREX-AQ revealed a factor of 2.3 error in the quantification of the NOAA acrolein gas standard (personal communication, A. Wisthaler and M. Coggon, 2021). This is the same standard used in Koss et al. (2018).

For biacetyl (2,3-butanedione), it is likely that the work of Koss et al. (2018) did not account for all potential isomers in the PTR-ToF-MS interpretation. The molecular formula for this compound is  $C_4H_6O_2$ . Using GC-CIMS data, Koss et al. (2018) inferred contributions to this PTR-ToF-MS signal from biacetyl (87%), methacrylate (5%), and other unidentified compounds (8%). Previous work has suggested the presence of additional isomers that are not easily detected by GC. In one study of pine burning emissions, 2-oxobutanal emissions were 3 times greater than those of biacetyl (Schauer et al., 2001). 1,4-butanedial has also been observed in significant amounts in tobacco smoke (personal communication, A. Wisthaler, 2021). Based on the likely presence of these compounds, we conservatively reduce biacetyl by a factor of 10.

These adjustments reduce model over-prediction for APAN (produced solely from acrolein oxidation) and PAN (where biacetyl is a major precursor) in sensitivity simulations described in the main text.

### **Text S2. Other oVOC**

Figure S12 shows the age progression of several other oVOC. Methanol is long-lived, and variability may reflect changing emissions or background conditions (Fig. S12a). A sharp rise in the methanol NEMR at 2 h may be another indicator of biogenic influence. Acetone and propanal are isomers ( $C_3H_6O$ ) and are reported as a sum in the SEAC<sup>4</sup>RS dataset. Acetone is likely the dominant isomer given the short lifetime of propanal, and this is consistent with the small NEMR variability as the lifetime of acetone against oxidation is weeks (Fig. S12b). The hydroxyacetone NEMR is relatively constant with age, and model values agree with observations within uncertainties (Fig. S12c). The sum of MVK and MACR tells a story similar to acetaldehyde, with a biogenic signature at ~2 h and an over-rapid decline in the base simulation (Fig. S12d). Results from other simulations are discussed in the main text when relevant.

### **Text S3. Additional NO<sub>y</sub> Details**

Several studies have noted potential positive artifacts in NO<sub>2</sub> measurements due to decomposition of thermally unstable gases in the sample inlet or instrument (Browne et al., 2011; Silvern et al., 2018; Nault et al., 2015). This is unlikely to explain the discrepancy between observed and modeled NO<sub>x</sub> in simulations M0 and M1 (Fig. 2i) for several reasons. First, such an interference would need to affect both the TDLIF and chemiluminescence NO<sub>2</sub> measurements similarly, as these two measurements are strongly correlated:  $NO_2(\text{TDLIF}) = 1.2 * NO_2(\text{CL}) - 0.12$  ppbv,  $r^2 = 1.00$ . Second, if the artifact were due to known NO<sub>x</sub> reservoirs, the conversion efficiency would need to be substantial. The difference between observed and

modeled  $\text{NO}_2$  in simulation M1 is  $260 \pm 100$  pptv at ages beyond 5 h. Mean observed PAN and total PNs are 1.3 and 2.1 ppbv, respectively. Thus a conversion efficiency of 10% or more would be required to fully explain the model-measurement difference, and this is unlikely given typical aircraft cabin and inlet temperatures ( $< 40$  °C). Modeled  $\text{HO}_2\text{NO}_2$  and  $\text{CH}_3\text{O}_2\text{NO}_2$  are  $< 5$  pptv and  $< 1$  pptv, respectively. We cannot rule out the potential influence of yet-unidentified  $\text{NO}_x$  reservoirs, though previous work suggests such artifacts are likely limited to the upper troposphere (Silvern et al., 2018).

In addition to PAN (discussed in the main text), the SEAC<sup>4</sup>RS dataset includes observations of several other speciated peroxy nitrates (PNs) and a total PN measurement. Other speciated PNs, shown in Fig. S14, include peroxypropionyl nitrate (PPN), peroxyacryloyl nitrate (APAN), and peroxyisobutyryl nitrate (PiBN). In the base simulation, early PPN NEMR growth is under-predicted, but the model and observations converge after 2h. APAN and PiBN are generally under-predicted, due in part to a lack of VOC precursors in the base simulation. Changes in model PNs in simulation M1 reflect increases in VOC precursors. In particular, APAN is produced solely through oxidation of acrolein. All PNs increase upon addition of initial HONO or  $\text{pNO}_3^-$  photolysis due to more  $\text{RO}_2$  and  $\text{NO}_2$ . Conversely, heterogeneous  $\text{NO}_2$  conversion to HONO has essentially no effect on PN NEMRs. In this case, decreasing  $\text{NO}_2$  and increasing NO offsets the increase in  $\text{RO}_2$ .

Model-measurement comparison with the  $\Sigma\text{PN}$  observations tell a qualitatively similar story to the speciated data (Fig. S15a). This measurement (via thermal dissociation and laser-induced fluorescence detection of  $\text{NO}_2$ ) is typically higher than the sum of speciated PN measurements (via thermal dissociation and detection of the peroxyacyl radicals), and in the first few hours this difference exceeds the combined uncertainty of the measurements. The reasons for this difference are unclear.

Alkyl nitrates (ANs) are minor products of the reaction of organic peroxy radicals ( $\text{RO}_2$ ) with NO. The observed  $\Sigma\text{AN}$  NEMR is variable with no clear trend (Fig. S15b). The simulated  $\Sigma\text{AN}$  NEMR is relatively constant throughout each simulation, and all simulations fall within the variability of observed NEMRs.

#### **Text S4. Co-optimization of Unmeasured VOC and HONO**

Additional reactive VOC and HONO chemistry collectively improve model-measurement agreement for most species. HONO sensitivity simulations presented in Sect. 3.3 of the main text utilize the same initial VOC as simulation M1. Uncertainties in initial VOC concentrations stem from the age distribution and history of sampled smoke, adaptation of laboratory-derived emission factors to ambient conditions, potential errors in the emission factors themselves, and translation of unmeasured VOC to MCM species. Uncertainties in primary and secondary HONO are also significant, as we lack a HONO measurement for comparison and putative aerosol mechanisms are not well understood.

In an extended set of simulations, we simultaneously tune initial unmeasured VOC, initial HONO, and  $\text{pNO}_3^-$  photolysis. For these runs, default values for initial unmeasured VOC concentrations (M1), initial HONO (M2c), and particulate nitrate photolysis rate (M3b) are each independently scaled by factors of 0, 0.25, 0.5, 0.75, and 1. Iterating over all combinations yield 125 simulations. Results from these simulations are analyzed in terms of normalized mean bias (NMB) (Gustafson and Yu, 2012) relative to observed NEMR age profiles. Heterogeneous  $\text{NO}_2$  uptake is not included, as tests with this mechanism indicate that it degrades agreement with observed ozone and other species.

Figure S26a-e shows the NMB of the ozone NEMR for each of the 125 simulations. Consistent with the examples presented in the main text, this bias trends negative with increasing VOC and positive with increasing initial or secondary HONO. Multiple scaling combinations produce a net O<sub>3</sub> NEMR NMB near zero (Fig. S27a). In general, increasing initial unmeasured VOC concentrations necessitates higher HONO to maintain agreement with observed O<sub>3</sub>. Initial and secondary HONO similarly influence the overall O<sub>3</sub> NEMR NMB, though this simplified metric masks age-dependent differences (Fig. S28).

Figures S26f-o and S27b-c show analogous NMB gradients and minimum-value isopleths for NO<sub>x</sub> and PAN NEMRs. Compared to the results for ozone, higher HONO is required to close the NO<sub>x</sub> budget for any given value of unmeasured VOC. PAN bias can be minimized at higher VOC, but only with low initial HONO and relatively slow pNO<sub>3</sub><sup>-</sup> photolysis. No combination of scaling factors optimizes agreement among all observations.

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Table S1. SEAC<sup>4</sup>RS measurement details.

Measurement	Instrument <sup>a</sup>	Accuracy
Pressure Temperature	MMS	< 1%
H <sub>2</sub> O	DLH	5%
CO	DACOM	2%
Photolysis frequencies	CAFS	12 – 20% <sup>b</sup>
O <sub>3</sub>	NOyO3	3%
NO		4%
NO <sub>2</sub>		7%
NO <sub>2</sub>	TDLIF	5%
ΣPN		10%
ΣAN		15%
H <sub>2</sub> O <sub>2</sub> HNO <sub>3</sub> HCN Peroxyacetic acid Hydroxyacetone Hydroxymethyl hydroperoxide Ethene hydroxynitrate Propene hydroxynitrate Butene hydroxynitrate Ethanal nitrate Propanone nitrate Isoprene hydroxynitrate	CIT-CIMS	30%
		30%
		50%
		50%
		40%
		50%
		50%
		30%
		50%
		50%
		50%
		30%
PAN PPN APAN PiBN		GT-CIMS
	20%	
	40%	
	40%	
VOC <sup>c</sup>	WAS	5%
Acetonitrile Methanol Acetaldehyde Acetone + propanal MVK + MACR Isoprene + Furan	PTR-MS	15%
		15%
		15%
		5%
		10%
		5%
HCHO	ISAF CAMS	10% 4%
Particulate nitrate	AMS	17%
Aerosol surface area	LAS	20%
Solar irradiance	BBR	5%

<sup>a</sup>See Toon et al. (2016) for details.

<sup>b</sup>Varies based on uncertainties in recommended cross sections and quantum yields.

<sup>c</sup>Methyl nitrate, ethyl nitrate, isopropyl nitrate, n-propyl nitrate, 2-butyl nitrate, 3-methyl-2-butyl nitrate, 3-pentyl nitrate, 2-pentyl nitrate, methane, ethane, propane, n-butane, isobutene, n-pentane, isopentane, n-hexane, 2-methyl pentane, 3-methyl pentane, 2,3-dimethylbutane, n-heptane, ethene,

propene, 1-butene, cis-2-butene, trans-2-butene, isobutene, 1,3-butadiene, 1-pentene, propadiene, benzene, toluene, ethyl benzene, o-xylene, m-xylene + p xylene (measured as sum, assumed 50%/50% distribution), isoprene,  $\alpha$ -pinene,  $\beta$ -pinene.

Table S2. MCM assignments for unmeasured VOC.

Koss ID	Koss Formula	MCM Name	Emission Ratio ( $\Delta X/\Delta CO$ )
<b>Species with direct MCM analogues</b>			
Acetic acid + glycolaldehyde	C2H4O2H	CH3CO2H	10.7633
Acetic acid + glycolaldehyde	C2H4O2H	HOCH2CHO	5.3013
HONO	HNO2H	HONO	3.2765
2-furfural + 3-furfural + other HCO2	C5H4O2H	FURFURAL2	3.2625
Formic acid	CH2O2H	HCOOH	2.3024
2-(3H)Furanone	C4H4O2H	BZFUONE	1.8951
5-Methyl furfural +Benzene diols (=catechol, resorcinol)	C6H6O2H	MFURFURAL	1.8749
5-Methyl furfural +Benzene diols (=catechol, resorcinol)	C6H6O2H	CATECHOL	1.8749
Guaiacol (=2-methoxyphenol)	C7H8O2H	GUAIACOL	1.8696
Acrolein	C3H4OH	ACR	1.8446
2-Methylphenol (=o-cresol) + anisol	C7H8OH	CRESOL	1.7222
Phenol	C6H6OH	PHENOL	1.7124
2-Methoxy-4-methylphenol (= creosol)	C8H10O2H	MGUAIACOL	1.1605
2-methylfuran + 3-methylfuran + general HCO	C5H6OH	M2F	0.9074
methyl acetate + ethyl formate + hydroxyacetone	C3H6O2H	METHACET	0.9060
Pyruvaldehyde (=methyl glyoxal) + acrylic acid	C3H4O2H	MGLYOX	0.7083
Pyruvaldehyde (=methyl glyoxal) + acrylic acid	C3H4O2H	ACO2H	0.7083
Glyoxal	C2H2O2H	GLYOX	0.7048
MEK + butanal + 2-methylpropanal	C4H8OH	MEK	0.5282
MVK + methacrolein + crotonaldehyde	C4H6OH	C4ALDB	0.5053
Quinone (=p-Benzoquinone)	C6H4O2H	PBZQONE	0.4795
Ethanol	C2H6OH	C2H5OH	0.4481
2,5-dimethyl furan + 2-ethylfuran + other C2 substituted furans	C6H8OH	DIM25FURAN	0.4226
C2 Phenols + methyl anisol	C8H10OH	OXYLOL	0.4222
methyl acetate + ethyl formate + hydroxyacetone	C3H6O2H	ETHFORM	0.3624
Acetic anhydride	C4H6O3H	METHCOACET	0.3198
Benzaldehyde	C7H6OH	BENZAL	0.1962
2-methylfuran + 3-methylfuran + general HCO	C5H6OH	M3F	0.1779
2,3-butanedione + methyl acrylate + other HCO2	C4H6O2H	BIACET	0.1650
Styrene	C8H8H	STYRENE	0.1625
2-furfural + 3-furfural + other HCO2	C5H4O2H	FURFURAL3	0.1554
syringol	C8H10O3H	SYRINGOL	0.1485
Tolualdehyde	C8H8OH	PXYLAL	0.0900
Tolualdehyde	C8H8OH	MXYLAL	0.0900
3-methyl-2-butanone + 2-methylbutanal+3-methylbutanal+2-pentanone +3-pentanone	C5H10OH	MIPK	0.0877

MEK + butanal + 2-methylpropanal	C4H8OH	IPRCHO	0.0870
Tolualdehyde	C8H8OH	OXYLAL	0.0772
3-methyl-2-butanone + 2-methylbutanal+3-methylbutanal+2-pentanone +3-pentanone	C5H10OH	MPRK	0.0653
Methyl benzoic acid	C8H8O2H	PXYLCO2H	0.0652
Methyl benzoic acid	C8H8O2H	MXYLCO2H	0.0652
Sesquiterpenes	C15H24H	BCARY	0.0651
Methyl benzoic acid	C8H8O2H	OXYLCO2H	0.0559
3-methyl-2-butanone + 2-methylbutanal+3-methylbutanal+2-pentanone +3-pentanone	C5H10OH	DIEK	0.0428
Pyruvic acid	C3H4O3H	CH3COCO2H	0.0407
heptanal + 2,4-dimethyl-3-pentanone + heptanone	C7H14OH	C6H13CHO	0.0317
heptanal + 2,4-dimethyl-3-pentanone + heptanone	C7H14OH	HEPT3ONE	0.0186
Dimethyl sulfide	C2H6SH	DMS	0.0116
hexanal + hexanones	C6H12OH	C5H11CHO	0.0110
hexanal + hexanones	C6H12OH	HEX2ONE	0.0070
MEK + butanal + 2-methylpropanal	C4H8OH	C3H7CHO	0.0062
hexanal + hexanones	C6H12OH	HEX3ONE	0.0054
3-methyl-2-butanone + 2-methylbutanal+3-methylbutanal+2-pentanone +3-pentanone	C5H10OH	BUT2CHO	0.0041
3-methyl-2-butanone + 2-methylbutanal+3-methylbutanal+2-pentanone +3-pentanone	C5H10OH	C3ME3CHO	0.0041

**Species mapped to MCM using OH reaction rate coefficient and molecular formula**

2-furanmethanol + other HCO2	C5H6O2H	MEKAOH	1.7325
5-(hydroxymethyl)-2-furfural	C6H6O3H	C512OOH	1.0774
5-hydroxymethyl-2[3H]-furanone	C5H6O3H	C512OOH	0.9055
2-hydroxy-3-methyl-2-cyclopenten-1-one	C6H8O2H	HEX3ONDOOH	0.7753
Product of levoglucosan dehydration (pyrolysis)	C6H8O4H	M3HEXANO3	0.6436
2,5-di(hydroxymethyl)furan + Methyl hydroxy dihydrofurfural	C6H8O3H	CO1M22CHO	0.5730
Methyl methacrylate + other HCO2	C5H8O2H	HO2CO4CHO	0.5689
3-methyl-3-butene-2-one + cyclopentanone + HCO1 isomers	C5H8OH	PEBOH	0.5685
5-Hydroxy 2-furfural/2-furoic acid	C5H4O3H	C4DBDIKET	0.4838
2,4-Cyclopentadiene-1-one + 2 other HCO isomers	C5H4OH	HO25C6	0.4779
Vanillin	C8H8O3H	C7CO4EDB	0.4712
Methyl propanoate	C4H8O2H	MAE	0.4602
Vinyl guaiacol	C9H10O2H	LIMKET	0.3705
Acetamide	C2H5NOH	ACO2H	0.3637
5-hydroxymethyl tetrahydro 2-furanone + 5-hydroxy tetrahydro 2-furfural	C5H8O3H	CO2M33CO3H	0.3543
C3 furan + various HCO	C7H10OH	HO25C7	0.3052
1-Buten-3-yne	C4H4H	ACR	0.2894
pyrrole + butene nitrile isomers	C4H5NH	C5H8	0.2417

Eugenol + isoeugenol	C10H12O2H	LIMKET	0.2415
Nitromethane	CH3NO2H		0.2264
2-propynal	C3H2OH	ACR	0.2091
Dihydro furandione	C4H4O3H	HMACO3H	0.2083
Hydroxy benzoquinone	C6H4O3H	M3HEXANO3	0.2058
2-hydroxybenzaldehyde (=Salicylaldehyde)	C7H6O2H	HO3C5CHO	0.1526
Pyridine + pentadienenitriles	C5H5NH	M23C4	0.1432
Naphthalene	C10H8H	UDECOH	0.1388
C9 Aromatics	C9H12H	DECOH	0.1255
methane thiol	CH4SH	CHCL2CHO	0.1243
Methyl benzofuran	C9H8OH	NOPINAOH	0.1191
C6 Diones + C6 1-DBE esters	C6H10O2H	IEB4CHO	0.1141
Acrylonitrile	C3H3NH	DICLETOH	0.1138
Methyl thiophenes	C5H6SH	ETBE	0.1098
1,3-Cyclopentadiene	C5H6H	ME2BUT2ENE	0.1055
dimethylbenzofuran	C10H10OH	NOPINAOH	0.1045
Methyl cyclopentanone + cyclohexanone + other ketones	C6H10OH	M3PECOOH	0.0927
methyl isocyanate + hydroxyacetonitrile	C2H3NOH	ETHOX	0.0871
3-methylacetophenone	C9H10OH	C8BCCO	0.0870
Methyl propenyl benzene + ethyl styrene	C10H12H	C7MOCOCO3H	0.0860
Formamide	CH3NOH	CCL3CHO	0.0842
Indane + methyl styrenes + propenyl benzenes	C9H10H	APINENE	0.0820
Benzofuran	C8H6OH	NOPINAOH	0.0819
Propane nitrile	C3H5NH	CH3CCL2OH	0.0797
C10 Aromatics	C10H14H	NC9H2O	0.0792
Methyl pyrrole isomers + Pentene nitrile isomers	C5H7NH	ME2BUT1ENE	0.0743
C6 esters	C6H12O2H	EMPHCOME	0.0690
Benzonitrile	C7H5NH	MC6OTKETOH	0.0685
Thiophene	C4H4SH	IBUTOL	0.0610
2-methyl pyridine + 3-methylpyridine	C6H7NH	MIPK	0.0603
Dihydronaphthalene	C10H10H	C108NO3	0.0574
Dihydropyrrole + butane nitrile	C4H7NH	CL12PRCHO	0.0570
Methyl naphthalene	C11H10H	C129CO	0.0552
Propiolic acid	C3H2O2H	ALLYLOH	0.0542
Methyl chavicol (estragole)	C10H12OH	PINAL	0.0522
Ethylcyclopentanone	C7H12OH	HM33C4OH	0.0497
1,3-dimethylnaphthalene	C12H12H	NC1313OH	0.0493
Indene + propynyl benzene isomer	C9H8H	BPINENE	0.0488
4-pyridinol	C5H5NOH	TBUACET	0.0427
Camphor + other oxygenated monoterpenes	C10H16OH	C828PAN	0.0396
2-ethenyl benzofuran	C10H8OH	NOPINAOH	0.0379

2,5-dimethyl pyrrole + 1-ethylpyrrole + other C2 substituted pyrroles	C6H9NH	CYHXONAOOH	0.0366
Ethenamine	C2H5NH	C2H6	0.0346
Nitrobenzene	C6H5NO2H	ACECOCOCH3	0.0325
Pentanenitriles	C5H9NH	C5PAN6	0.0314
Phenylacetylene	C8H6H	MC6OTKETOH	0.0276
nitrotoluene	C7H7NO2H	ACCOPRONE	0.0246
dihydroxy pyridine + methyl maleimide	C5H5NO2H	M3PEAOH	0.0225
pyridine aldehyde + methylfuronitrile + nitrosobenzene	C6H5NOH	H25M2C6	0.0211
benzeneacetonitrile	C8H7NH	NC71CO	0.0207
C11 aromatics	C11H16H	C129CO	0.0197
ethylindene	C11H12H	BPINENE	0.0194
2-furancarboxitrile + 3-furancarboxitrile	C5H3NOH	H2M3C4CHO	0.0189
Trimethylamine	C3H9NH	MEPROPENE	0.0181
dimethyl pyridine + ethylpyridine + heptylnitriles	C7H9NH	M2PEDOH	0.0172
C7 acrylonitrile	C7H11NH	C6CO2OHPAN	0.0169
Acenaphthylene	C12H8H	DDEC3ONE	0.0164
Propene amine	C3H7NH	PXYL	0.0158
Cineole + other oxygenated monoterpenes	C10H18OH	HO36C10	0.0137
C12 aromatics	C12H18H	C126CHO	0.0113
4-methylpentanenitrile	C6H11NH	MPRK	0.0109
Carbon suboxide	C3O2H	EOX2COMEOH	0.0104
Methyl benzeneacetonitrile	C9H9NH	C920PAN	0.0104
Dimethyl disulfide	C2H6S2H	PXYCATECH	0.0098
C13 aromatics	C13H20H	BCKET	0.0097
butene nitrates	C4H7NO3H	NPRACBOOH	0.0078
Vinylpyridine	C7H7NH	THEX2ENE	0.0066
decanal	C10H20OH	NC11H24	0.0065
Nitrofuran	C4H3NO3H	MALDALCO3H	0.0064
Methanimine	CH3NH	C2H6	0.0062
Propiolonitrile (=propyne nitrile)	C3HNNH	DICLETOH	0.0058
Butene amine	C4H9NH	MVK	0.0055
nitroethene	C2H3NO2H	PROPACID	0.0054
Ethylpyrrole	C6H6N	CYHXONAOOH	0.0046
methane diol	CH4O2H	ETHOXOOH	0.0041
Nitroethane or ethane nitrite	C2H5NO2H	ETHOX	0.0035
Ethylamine + dimethylamine	C2H7NH	CRESOL	0.0030
Dihydro pyridine	C8H9NH	C5PAN6	0.0029
Nitropropanes	C3H7NO2H	PROPACID	0.0018
Cyanoallene isomers	C4H3NH	IPECOH	0.0013
C8 nitriles	C8H15NH	HEPT3ONE	0.0012
Dimethyl trisulfide	C2H6S3H	PXYCATECH	0.0011
n-sulfinyl methanamine	CH3NOSH	C2H6	0.0003

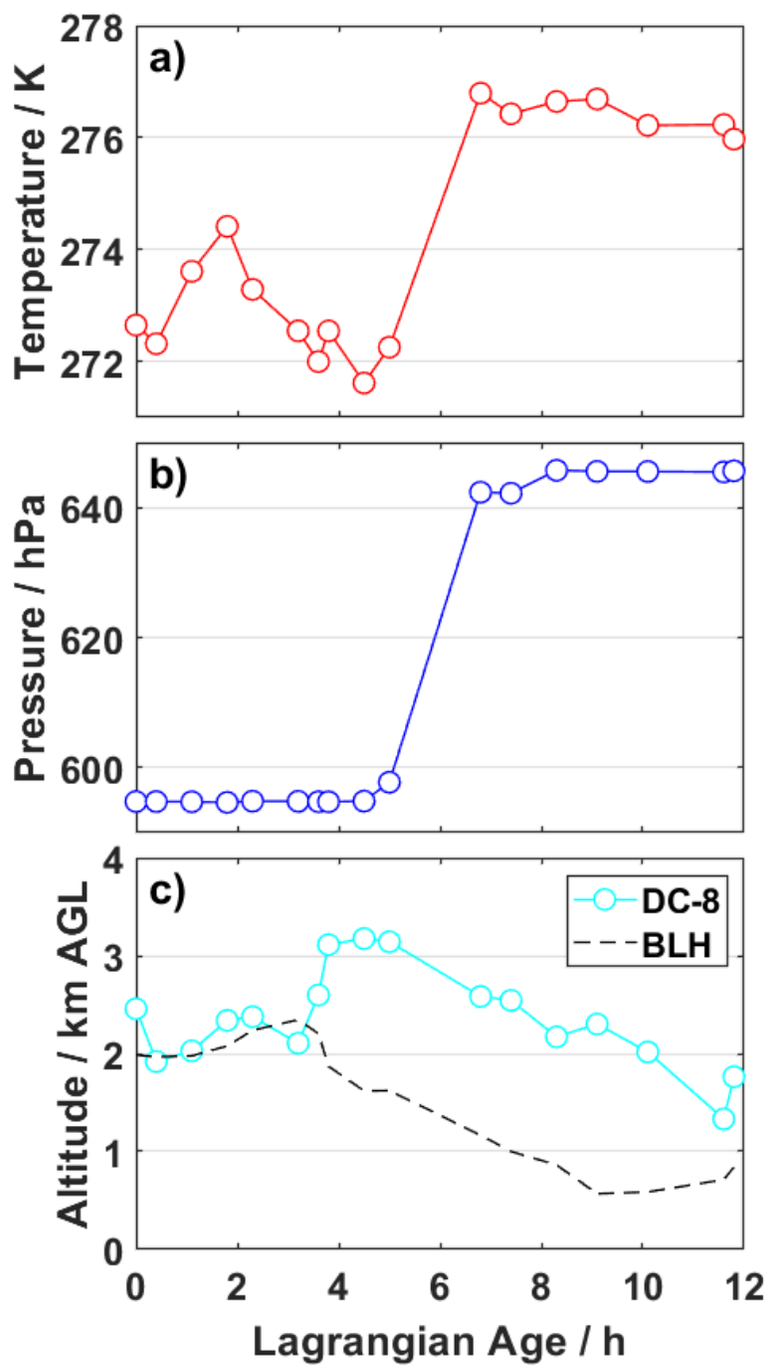


Figure S1. DC-8 sampling temperature (a), atmospheric pressure (b), and altitude above ground level (c, cyan circles) as a function of plume Lagrangian Age. The dashed line in (c) denotes the boundary layer height relative to ground level based on output from the two meteorological datasets used for trajectory analysis.

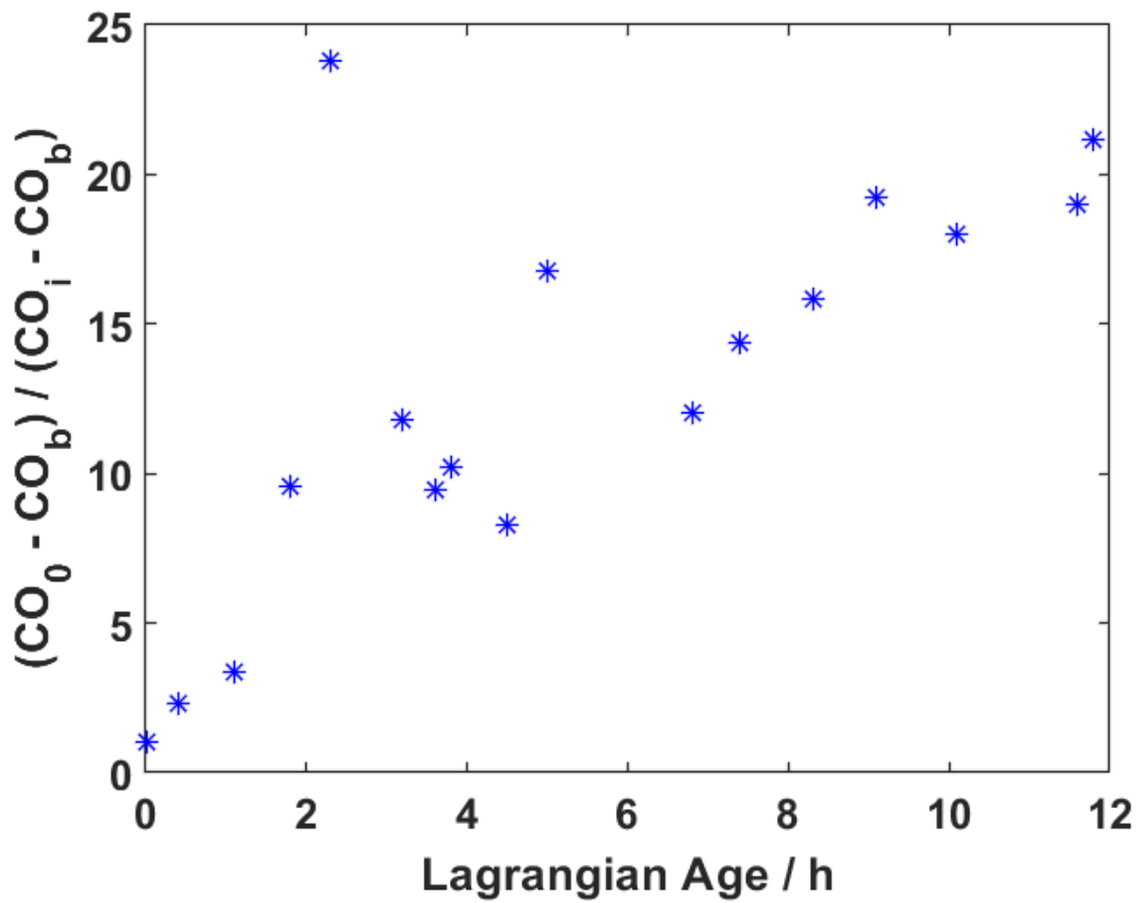


Figure S2. Dilution factor for each WAS plume sample, calculated as the ratio of initial to sample-time background-corrected CO.

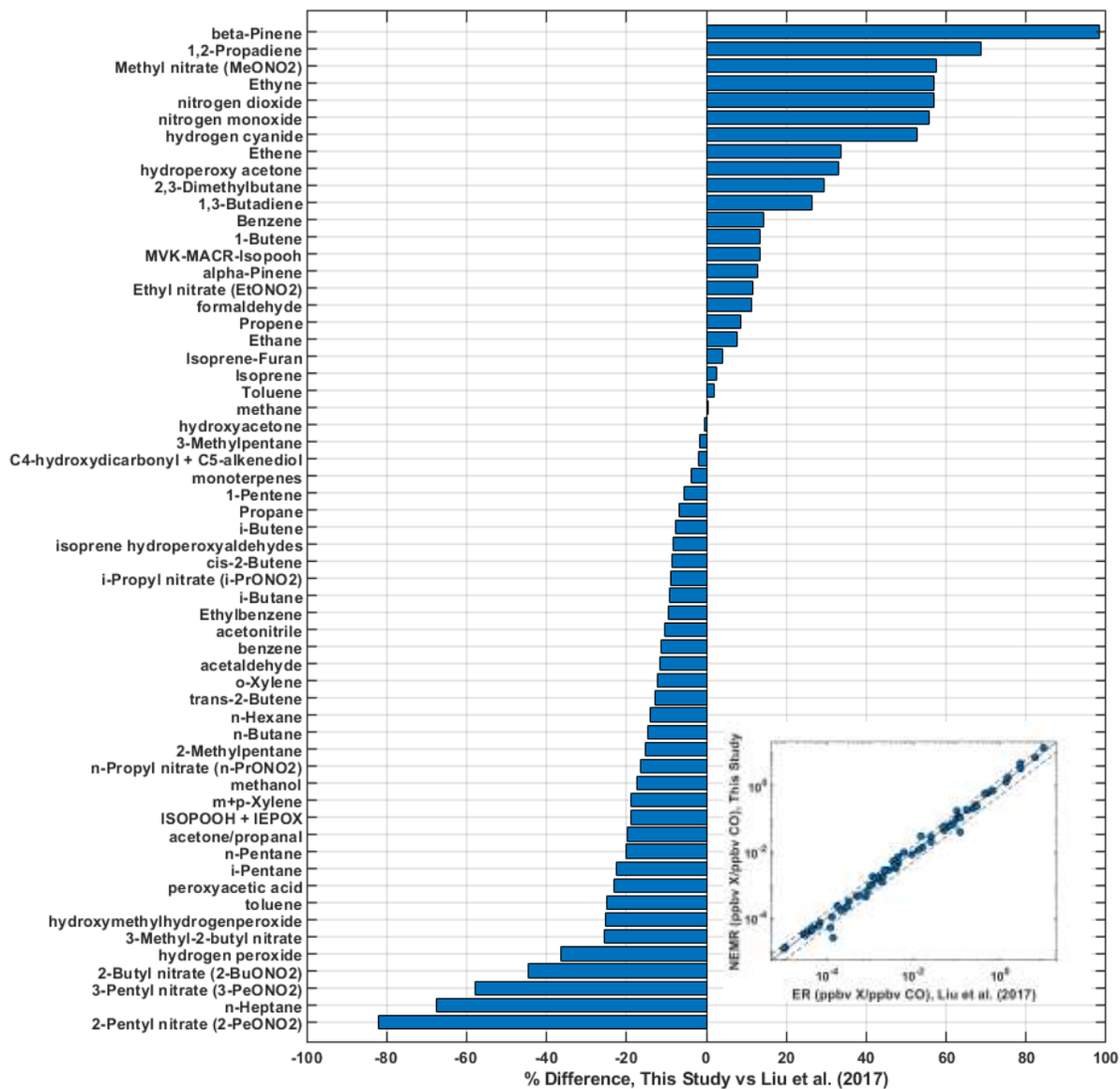


Figure S3. Comparison of normalized excess mixing ratios (NEMR) from the “source” sample of this study and the Rim Fire emission ratios (ERs) reported by Liu et al. (2017). Both NEMR and ER values are normalized to excess CO. In the species-specific plot, positive values correspond to species with a higher ratio in long-axis source sample, and values with ER < 10<sup>-4</sup> ppbv / ppbv are excluded. In the inset, the solid line is the 1:1 relationship and dashed lines are ±50%.

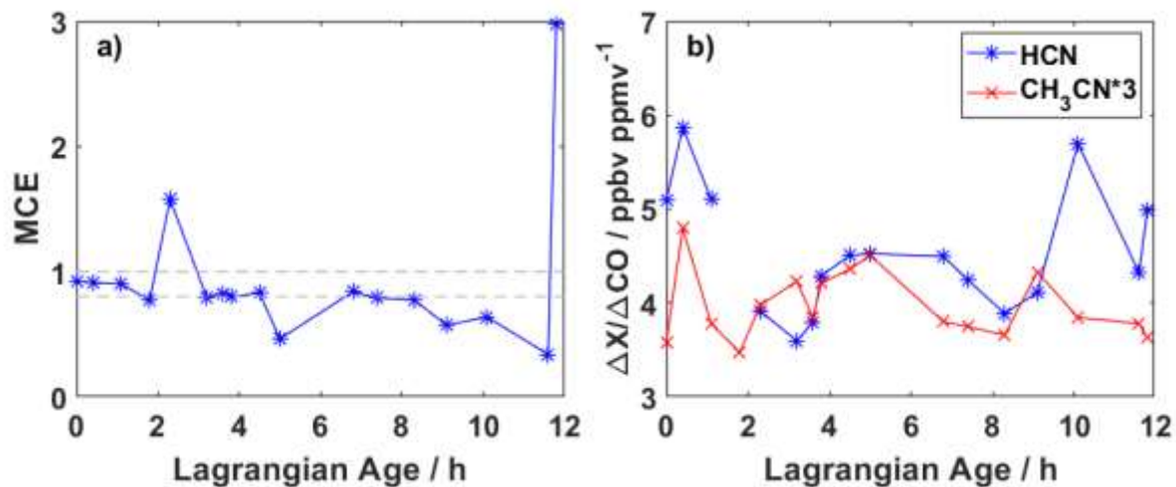


Figure S4. Modified combustion efficiency (MCE) (a) and NEMRs for formonitrile and acetonitrile (b). MCE is defined as  $\Delta\text{CO}_2 / (\Delta\text{CO} + \Delta\text{CO}_2)$ . Gray dashed lines in (a) denote the range of 0.8 – 1 typical of wildfires (Akagi et al., 2011).

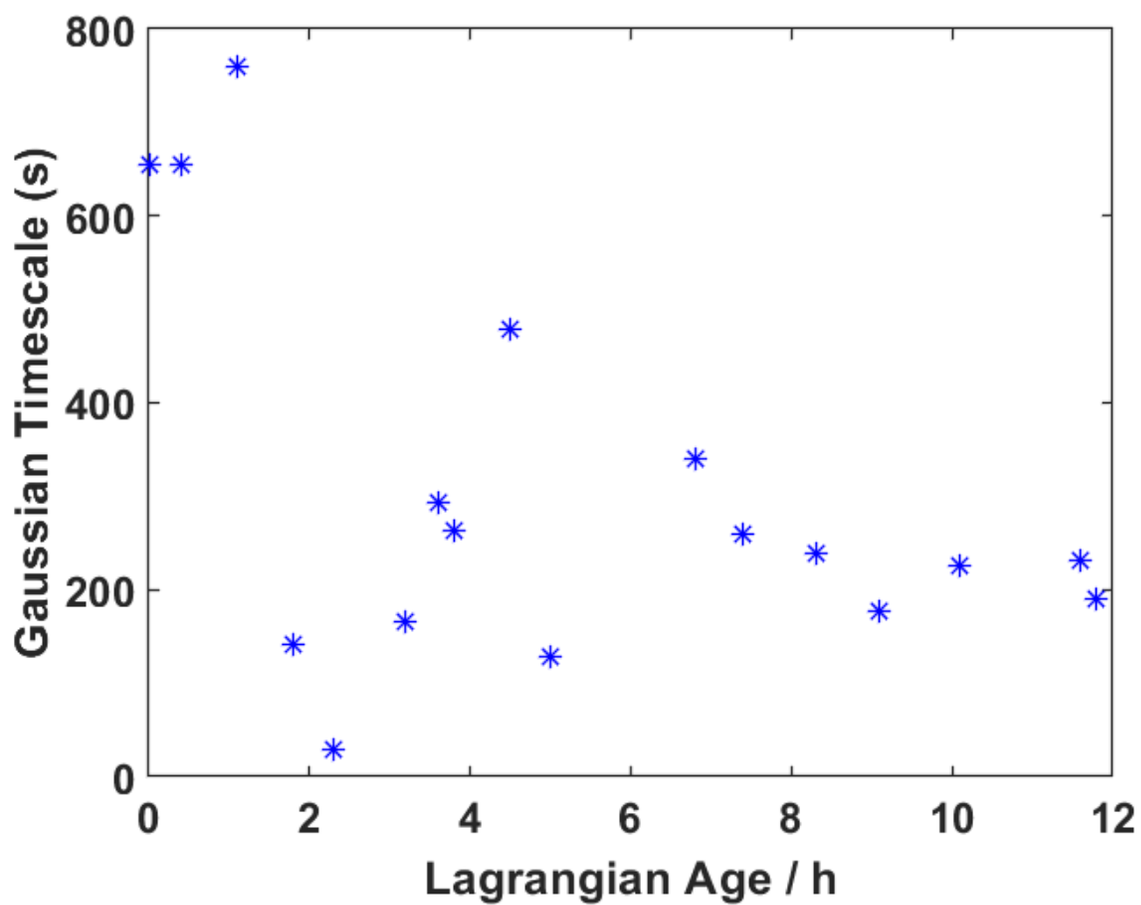


Figure S5. Gaussian dilution timescale for each model puff, calculated from observations of the decay of CO and Eqn. (2) as described in the main text.

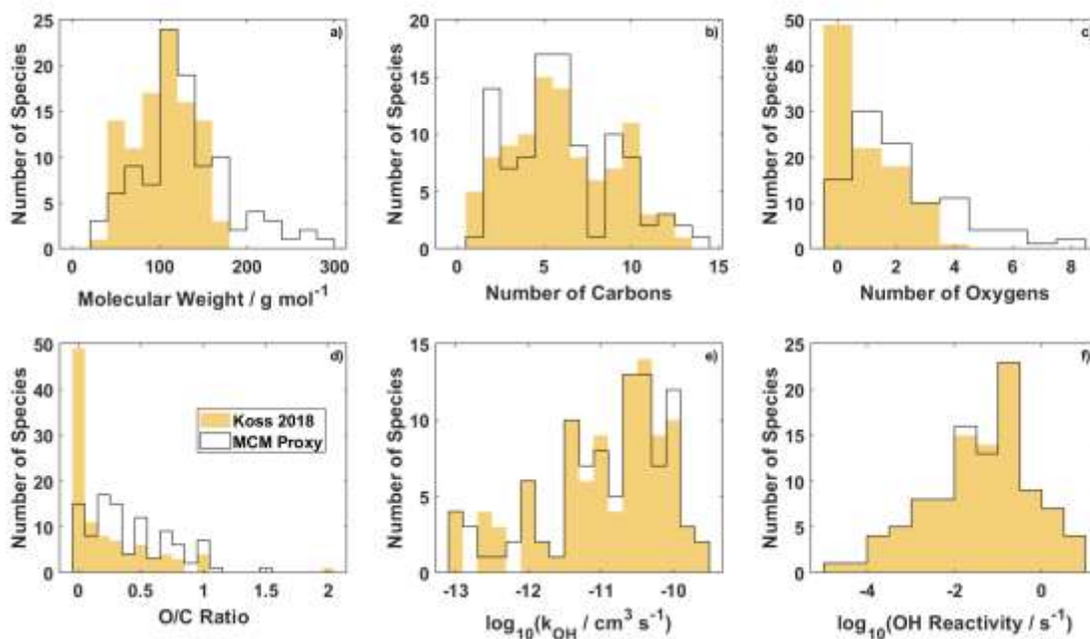


Figure S6. Comparison of chemical metrics for non-MCM unmeasured VOC from Koss et al. (2018) and MCM proxies (see Sect. 2.4.1 and Table S2). (a) Molecular weight, (b) number of carbons per molecule, (c) number of oxygens per molecule, (d) oxygen/carbon ratio per molecule, (e) OH reaction rate coefficient, and (f) initial OH reactivity. Note that (e) and (f) are on a log scale.

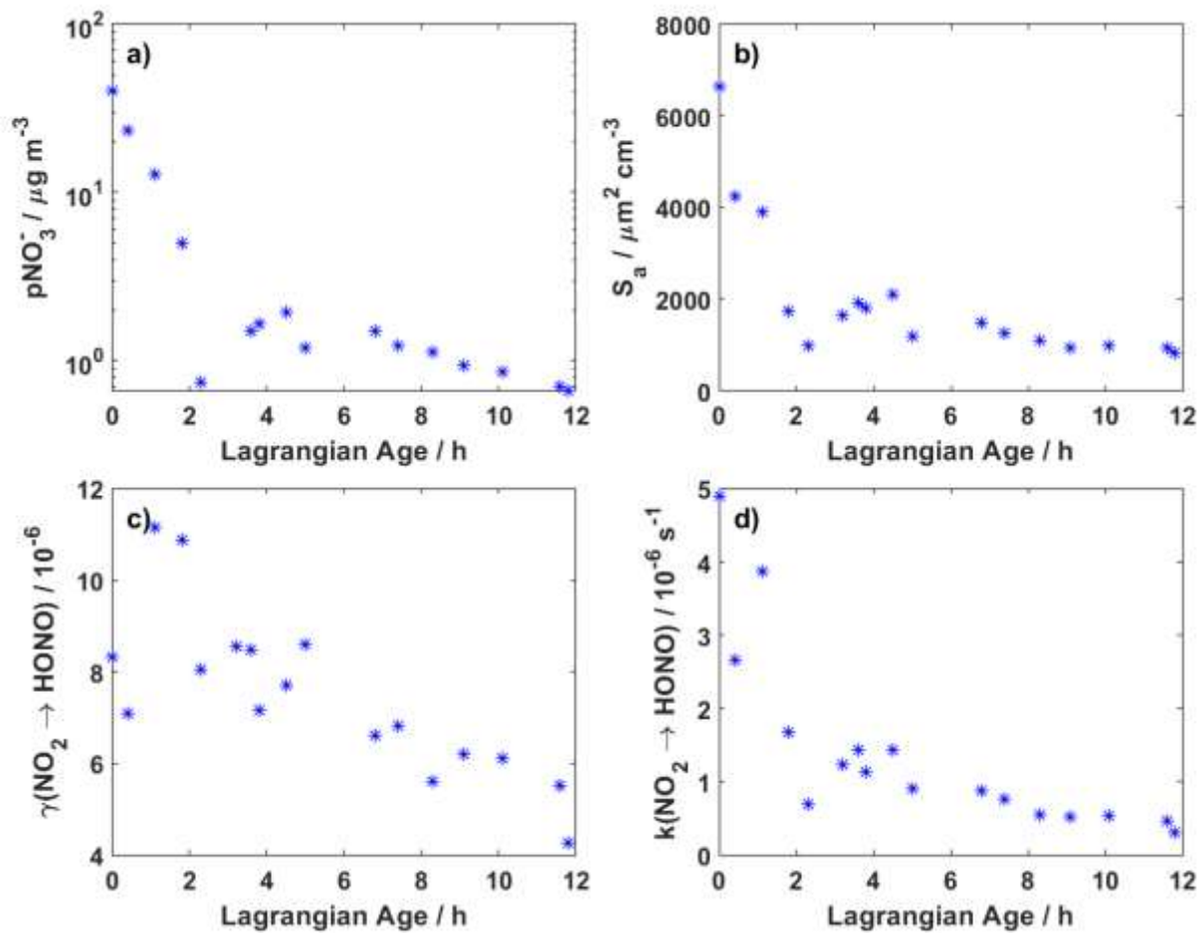


Figure S7. Observed and derived aerosol-related properties as a function of plume age: AMS-observed particulate nitrate mass concentration (a), LAS-observed aerosol surface area (b), calculated reactive uptake coefficient for  $\text{NO}_2$  conversion to HONO (c), and calculated first order rate coefficient for the same (d).

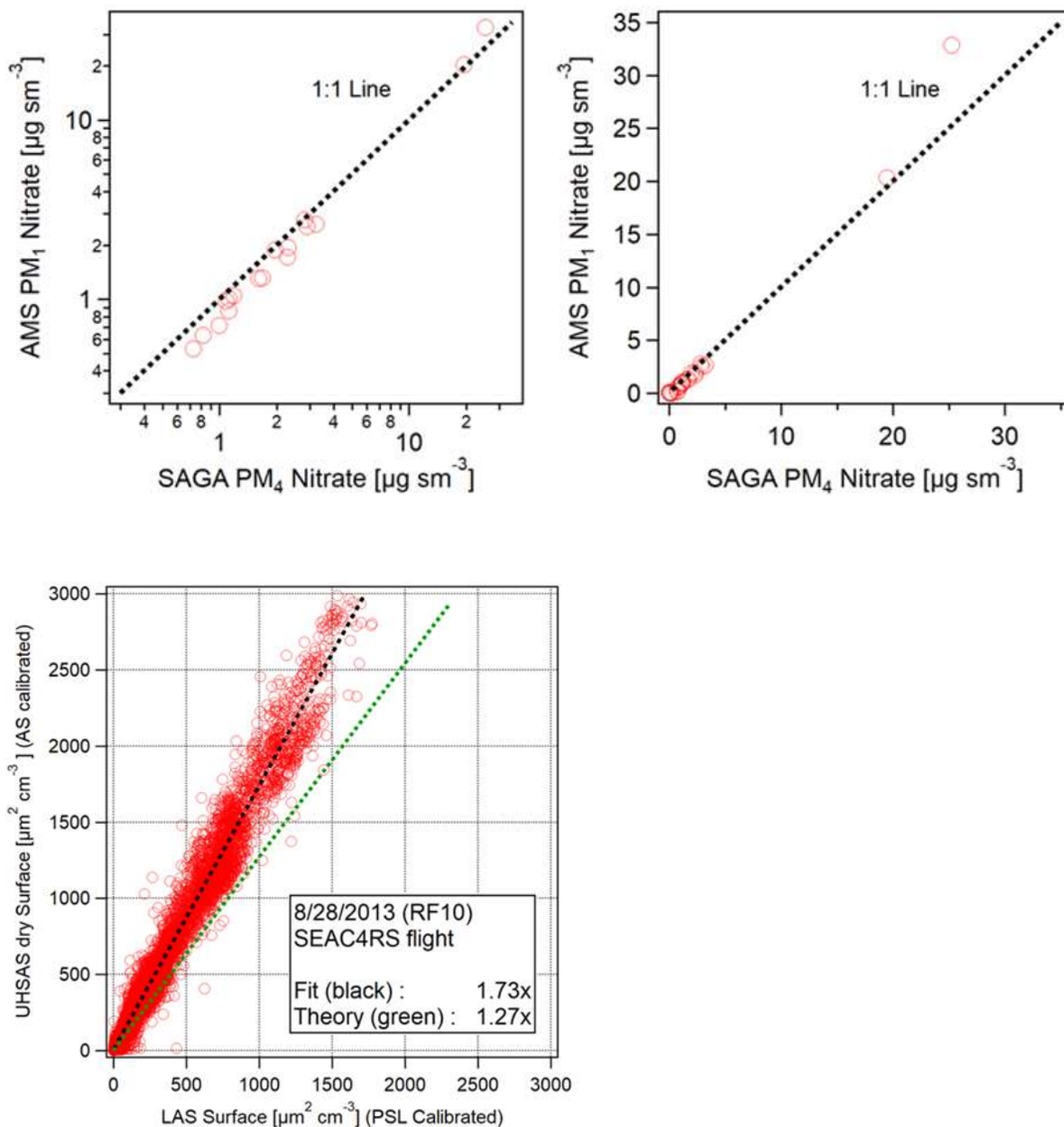


Figure S8. (top) Comparison of particulate nitrate observed by the AMS and SAGA instruments. The AMS has a size cut of  $\sim 1$  micron, while SAGA samples up to 4 microns. AMS data are averaged over the SAGA sampling interval ( $\sim 5$  minutes) for all Rim Fire observations. Data is shown on both a log (left) and linear (right) scale. (bottom) Comparison of aerosol surface area observed by the LAS and UHSAS instruments. Low bias in the LAS results from the use of PSLs for size calibration instead of ammonium sulfate (P. Campuzano-Jost, personal communication, 2021).

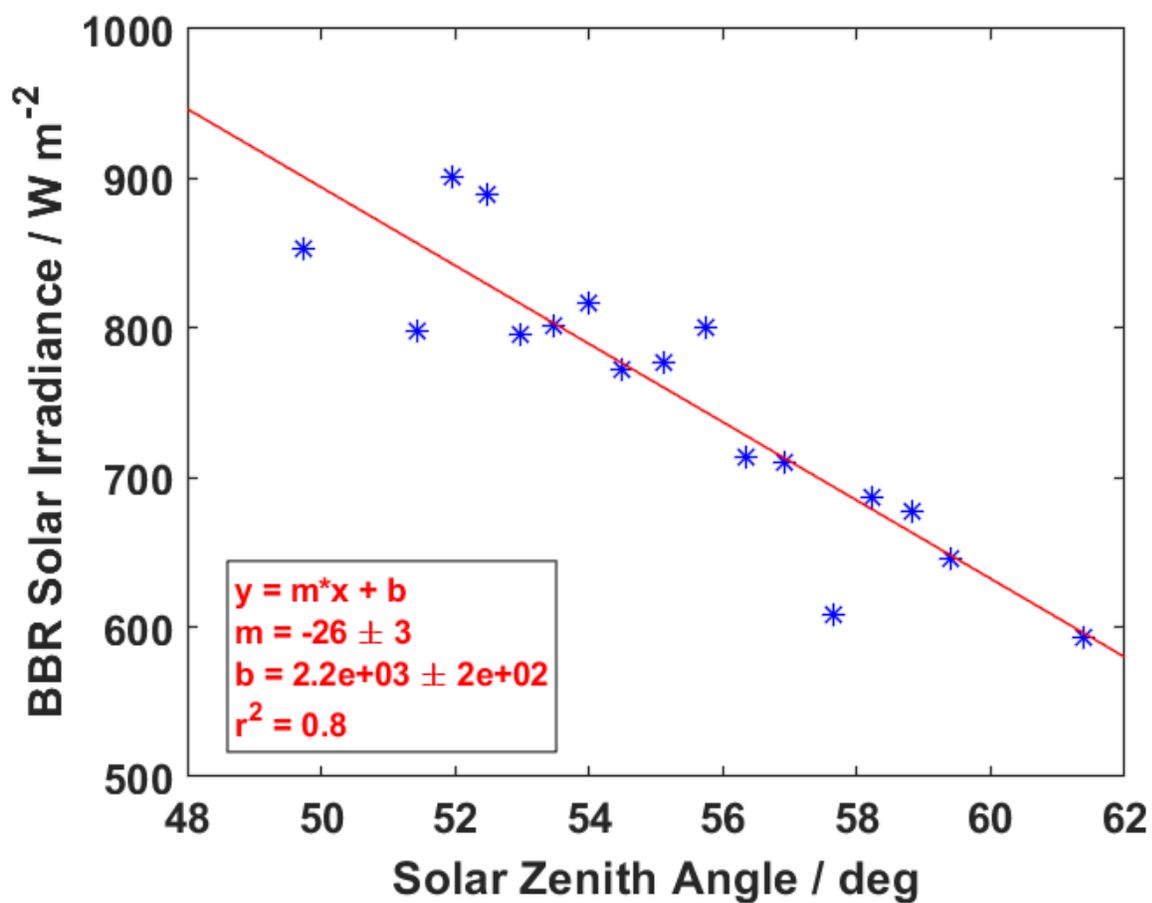


Figure S9. Linear relationship between solar zenith angle and total (up + down) solar irradiance from the broadband radiometer (BBR) instrument. The red line represents an ordinary least-squares fit, used to estimate irradiance for the parameterization of NO<sub>2</sub> reactive uptake.

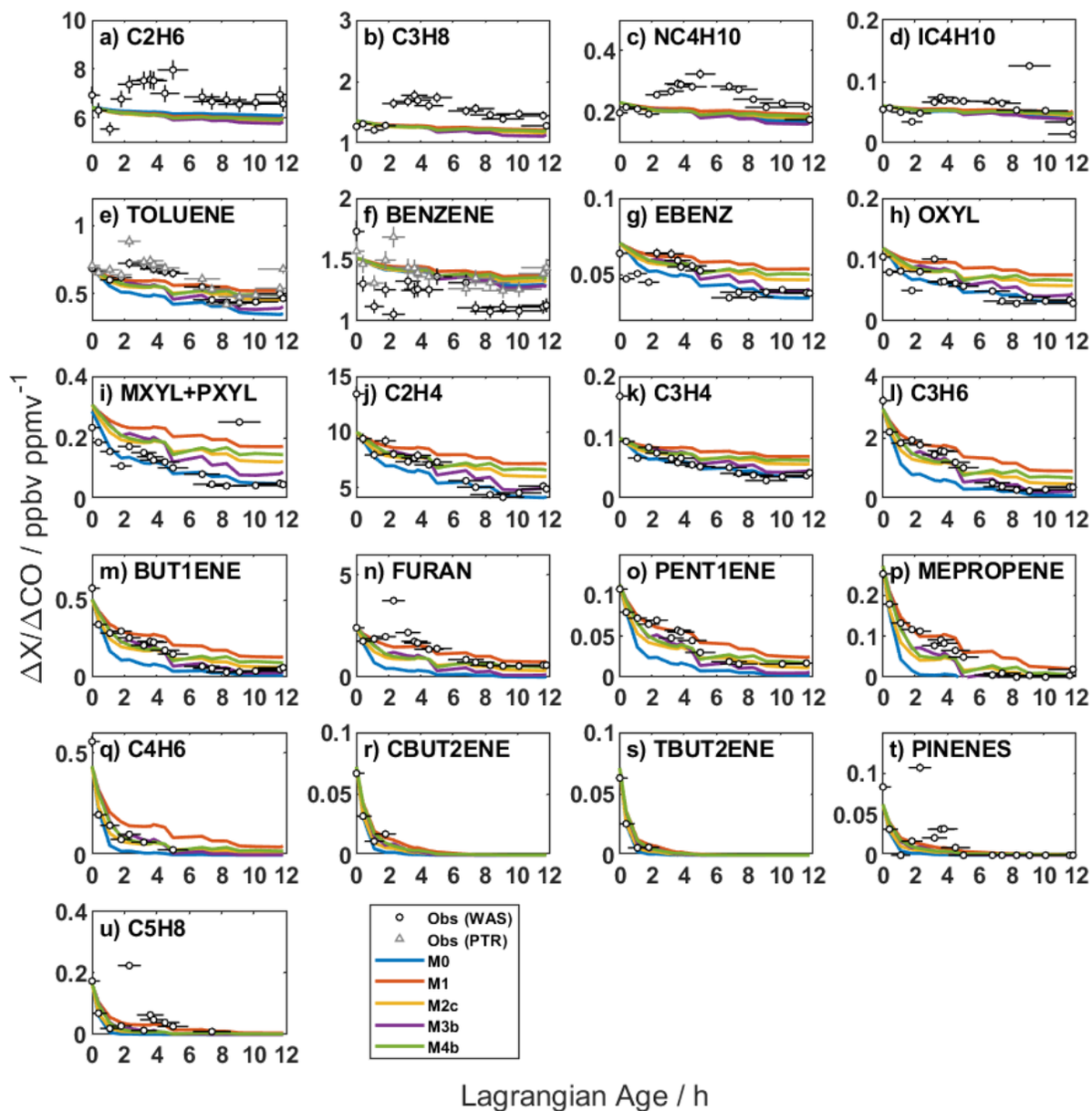


Figure S10. Age evolution of NEMRs for all observed VOC. Black circles and gray triangles are observations from the WAS and PTR-MS, respectively, with their corresponding uncertainty due to measurement accuracy and age. Species, in order from a) to u), are: ethane, propane, n-butane, i-butane, toluene, benzene, ethyl benzene, o-xylene, m-xylene + p-xylene, ethene, propadiene, propene, 1-butene, furan, 1-pentene, methyl propene (isobutene), 1,3-butadiene, cis-2-butene, trans-2-butene,  $\alpha$ -pinene +  $\beta$ -pinene, and isoprene. Colored lines are model output from the base simulation (M0, blue), addition of unmeasured VOC (M1, red), and addition of unmeasured VOC and primary HONO (M2c, yellow), secondary HONO via  $\text{pNO}_3^-$  photolysis (M3b, purple), or  $\text{NO}_2$  heterogeneous uptake (M4b, green). Note that the furan observation is the difference between PTR-MS (furan + isoprene) and WAS isoprene.

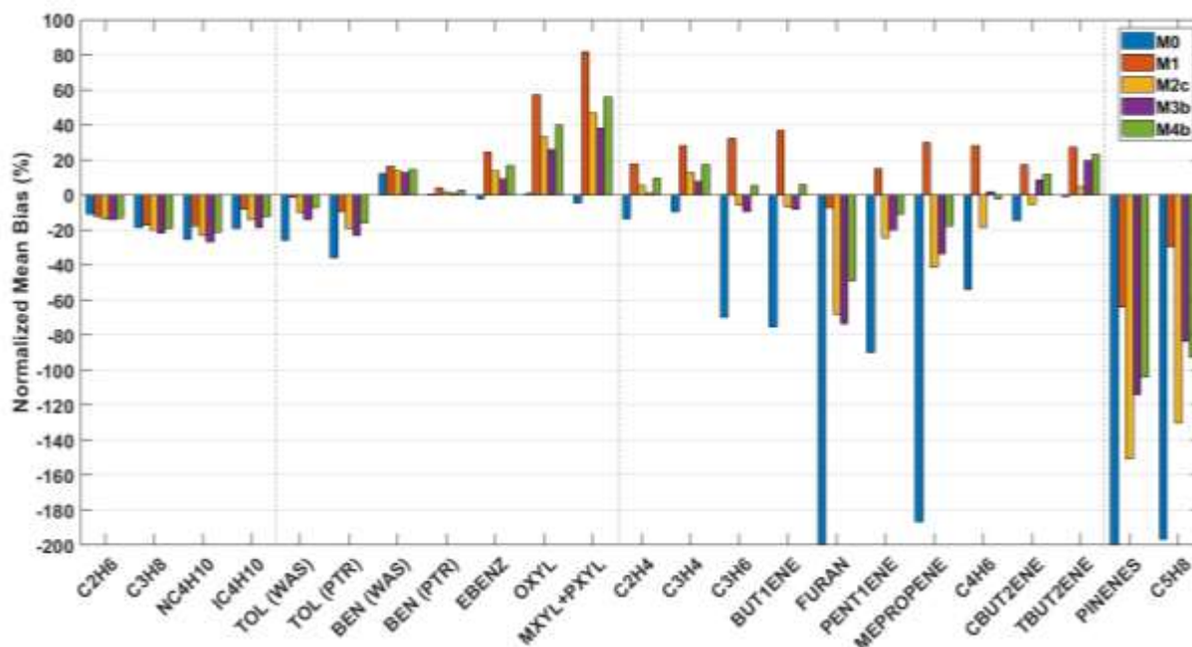


Figure S11. Normalized mean bias (NMB) of modeled VOC profiles compared to observations. For each simulation and each VOC, NMB is computed with model output shown in the previous figure following Gustafson and Yu (2012). Negative bias means that the model is lower than observations on average. Vertical dotted lines demarcate the four groups discussed in the main text.

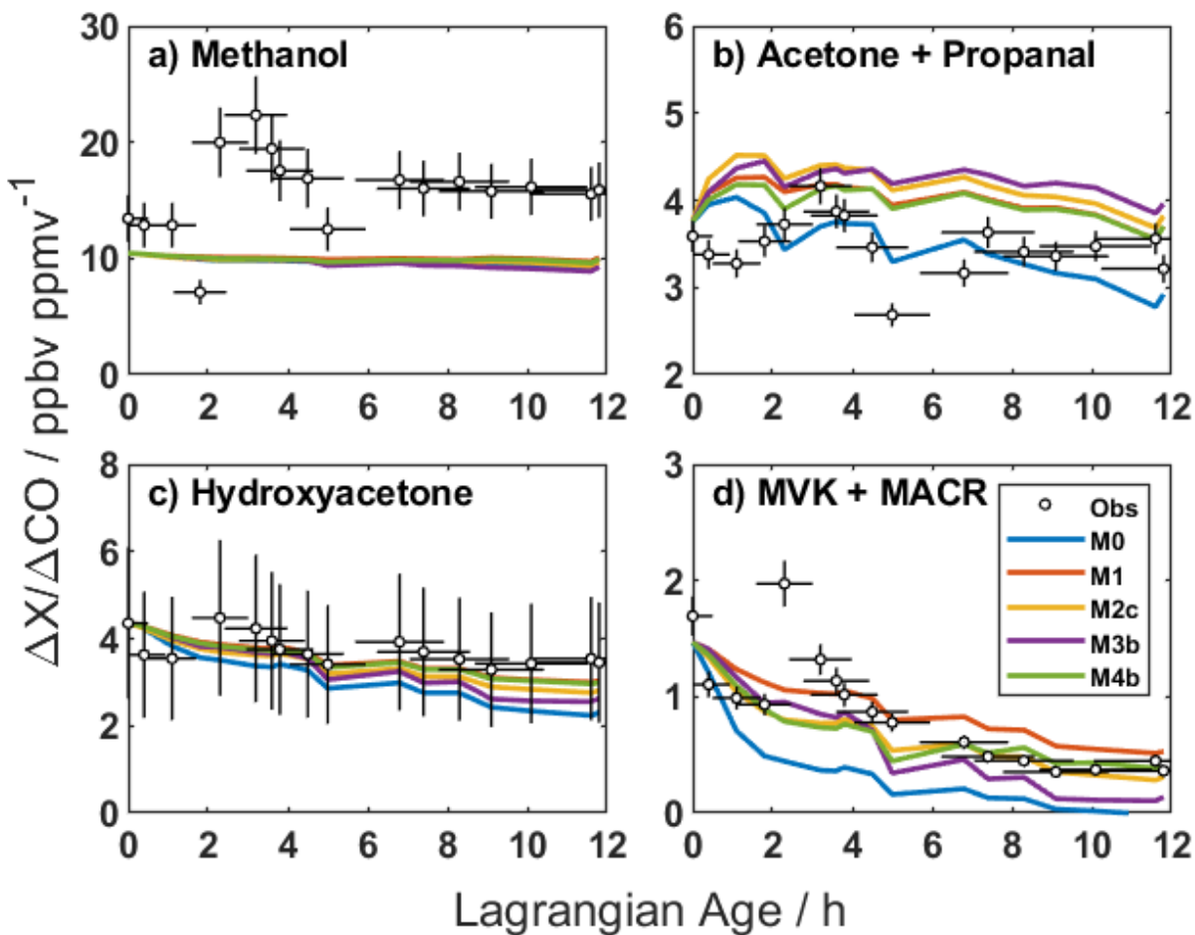


Figure S12. Age evolution of NEMRs for oxygenated VOC. Black circles are observations with their corresponding uncertainty due to measurement accuracy and age. Colored lines are as described in Fig. S10.

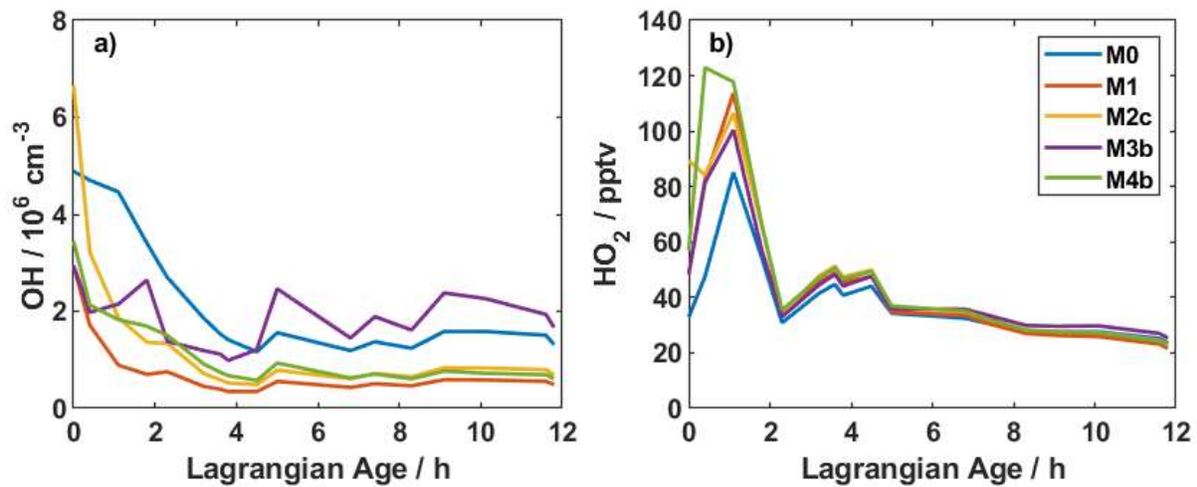


Figure S13. Age evolution of model-predicted OH concentration (a) and HO<sub>2</sub> mixing ratio (b). Colors are as described in Fig. S10.

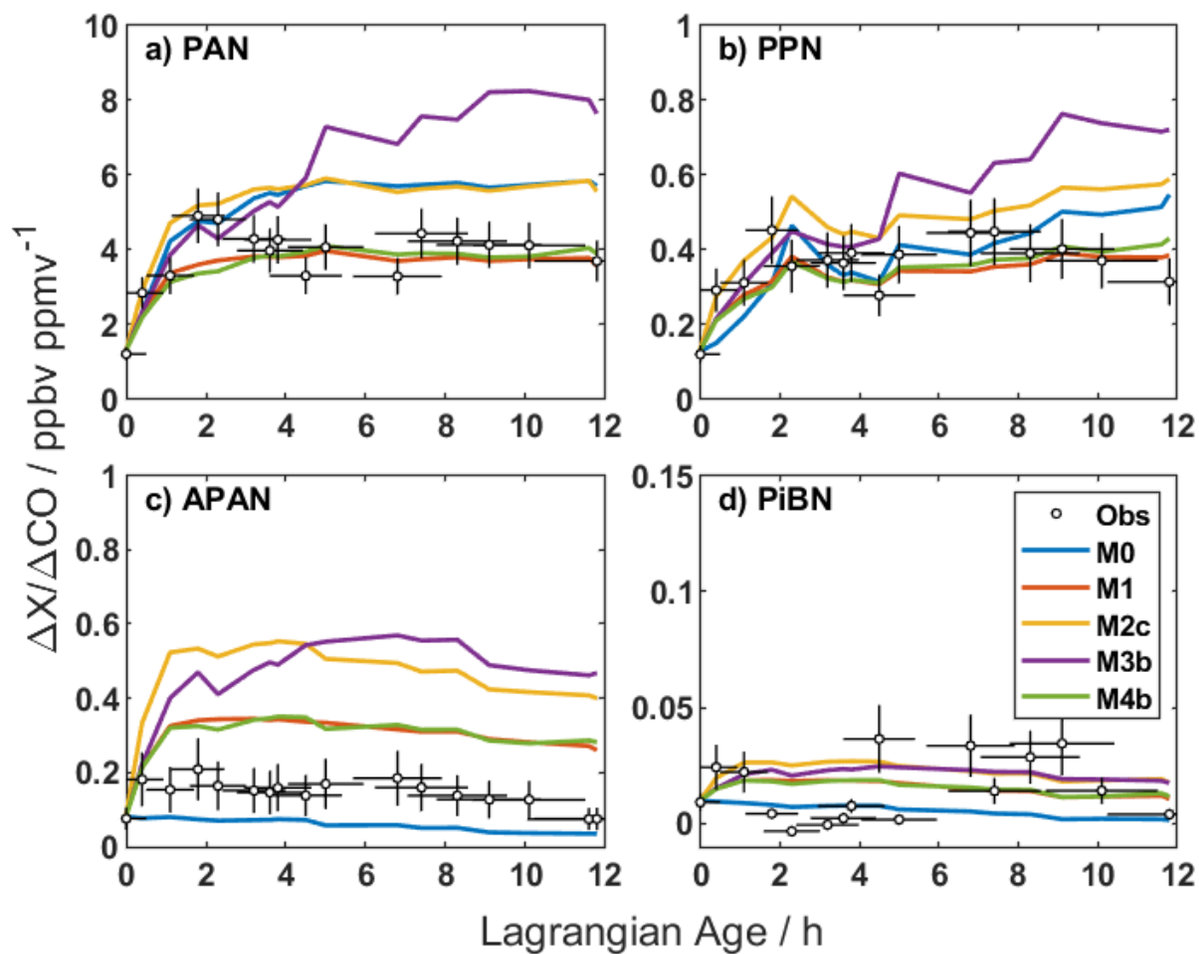


Figure S14. Age evolution of NEMRs for speciated peroxy nitrates. Black circles are observations with their corresponding uncertainty due to measurement accuracy and age. Colored lines are as described in Fig. S10.

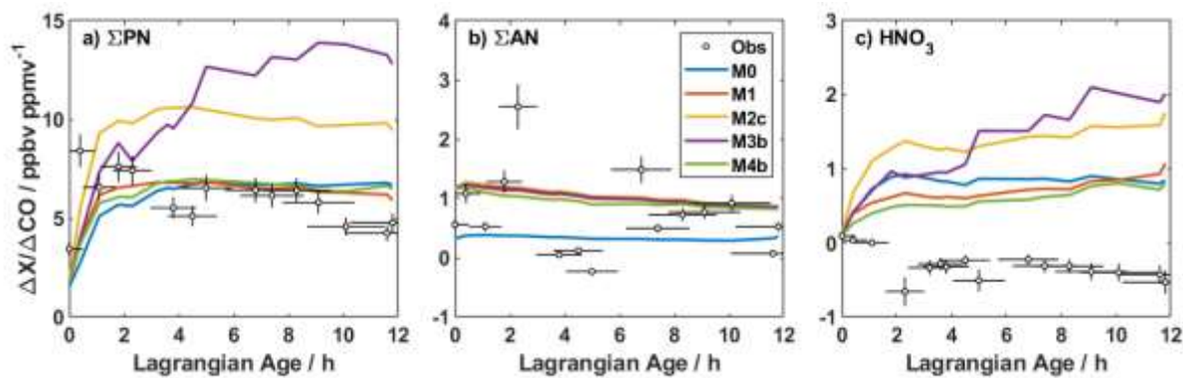


Figure S15. Age evolution of NEMRs for total peroxy nitrates (a), total alkyl nitrates (b), and nitric acid (c). Black circles are observations with their corresponding uncertainty due to measurement accuracy and age. Colored lines are as describe in Fig. S10. MCM PN and AN species are identified using simplified molecular-input line-entry system (SMILES) strings and SMILES filtering code provided with FOAM. Model  $\text{HNO}_3$  NEMRs deviate significantly from observations because the model does not account for gas-to-particle nitrate partitioning.

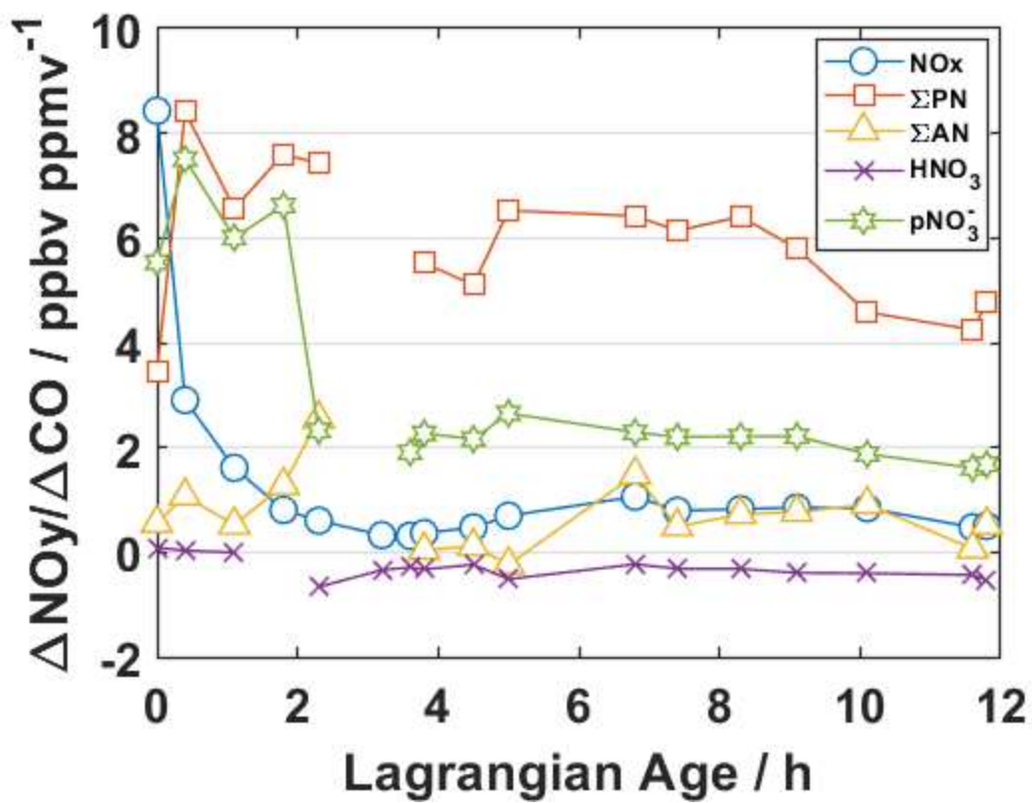


Figure S16. Comparison of age trends for different components of observed NO<sub>y</sub>, including NO<sub>x</sub> (blue circles), total peroxy nitrates (red squares), total alkyl nitrates (yellow triangles), nitric acid (purple X), and particulate nitrate (green stars).

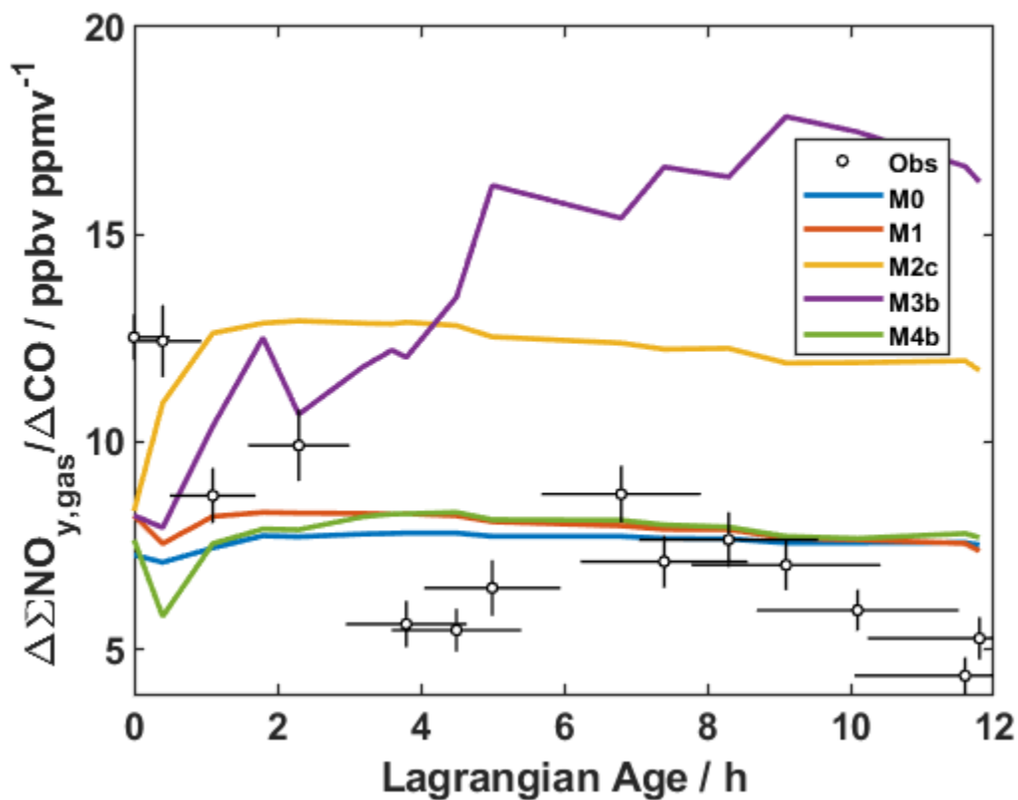


Figure S17. Age evolution of modelled gas-phase  $\text{NO}_y$ . Symbols and lines are as described Fig. S10. Observed values represent the sum of  $\text{NO}_x$ ,  $\Sigma\text{PN}$ ,  $\Sigma\text{HN}$ , and  $\text{HNO}_3$ . Model values represent the sum over the same modelled species and thus exclude  $\text{HONO}$ ,  $\text{HO}_2\text{NO}_2$ , and nitroaromatics.

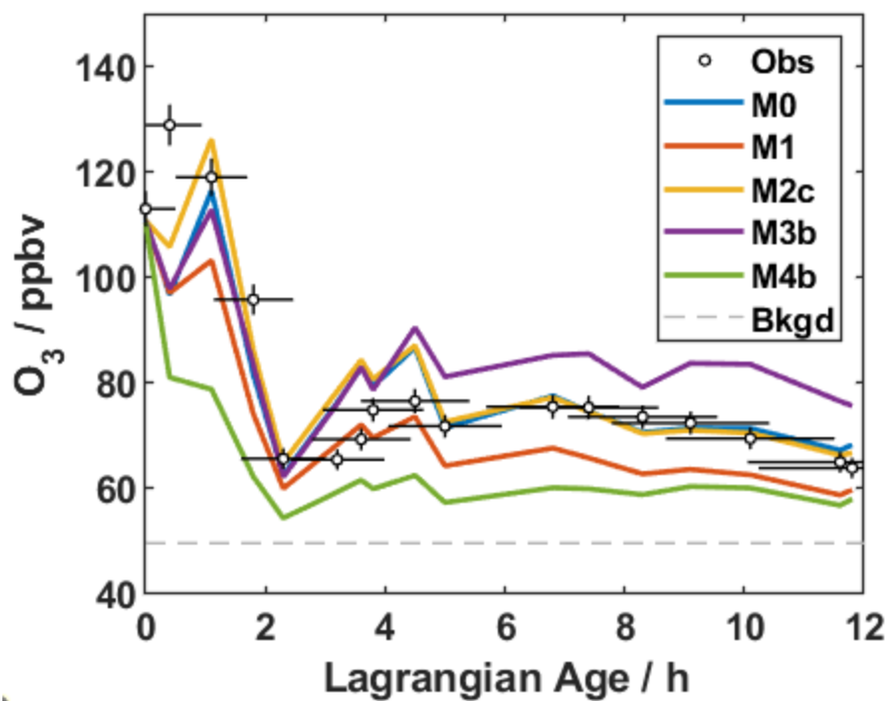


Figure S18. Age evolution of absolute ozone mixing ratio. Symbols and lines are as described Fig. S10. The grey dashed line denotes the estimated  $O_3$  background mixing ratio.

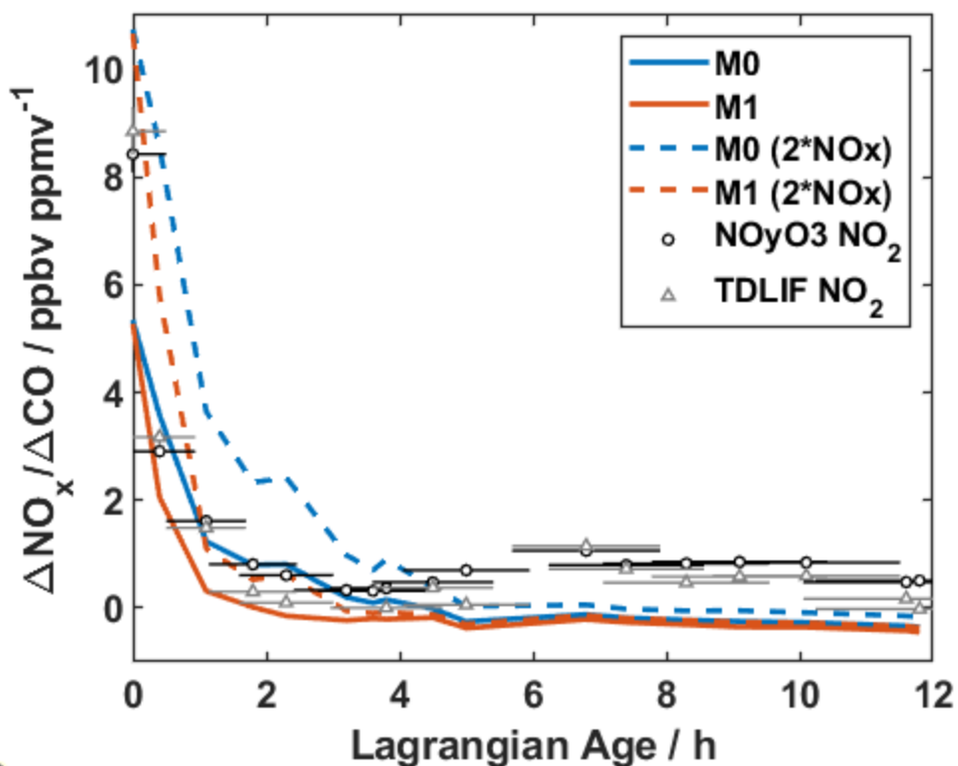


Figure S19. Comparison of NO<sub>x</sub> NEMRS for observations (symbols), simulations M0/M1 (solid lines), and sensitivity perturbations where initial NO<sub>x</sub> is doubled (dashed lines). For observations, black circles and gray triangles represent NO<sub>x</sub> calculated with two different NO<sub>2</sub> measurements and the same NO measurement (from the NO<sub>y</sub>O<sub>3</sub> instrument). Error bars denote uncertainty due to age estimation. Uncertainty due to measurement accuracy is small (4%).

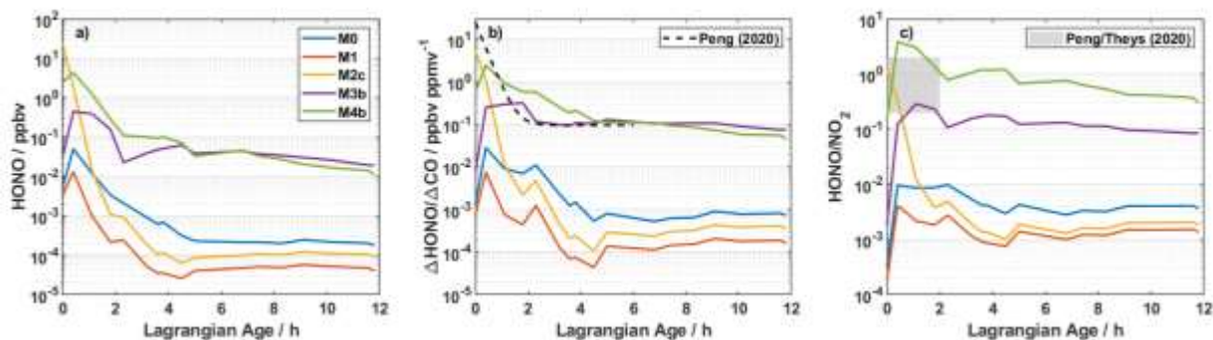


Figure S20. Age evolution of simulated absolute HONO mixing ratios (a), HONO NEMRs (b), and the ratio of HONO to  $\text{NO}_2$  (c). Colored lines are as described in Fig. S10. In (b), the black dashed line shows the fitted line from Fig. 3 of Peng et al. (2020). In (c), the shaded gray area is the range of values reported by Peng et al. (2020) and Theys et al. (2020).

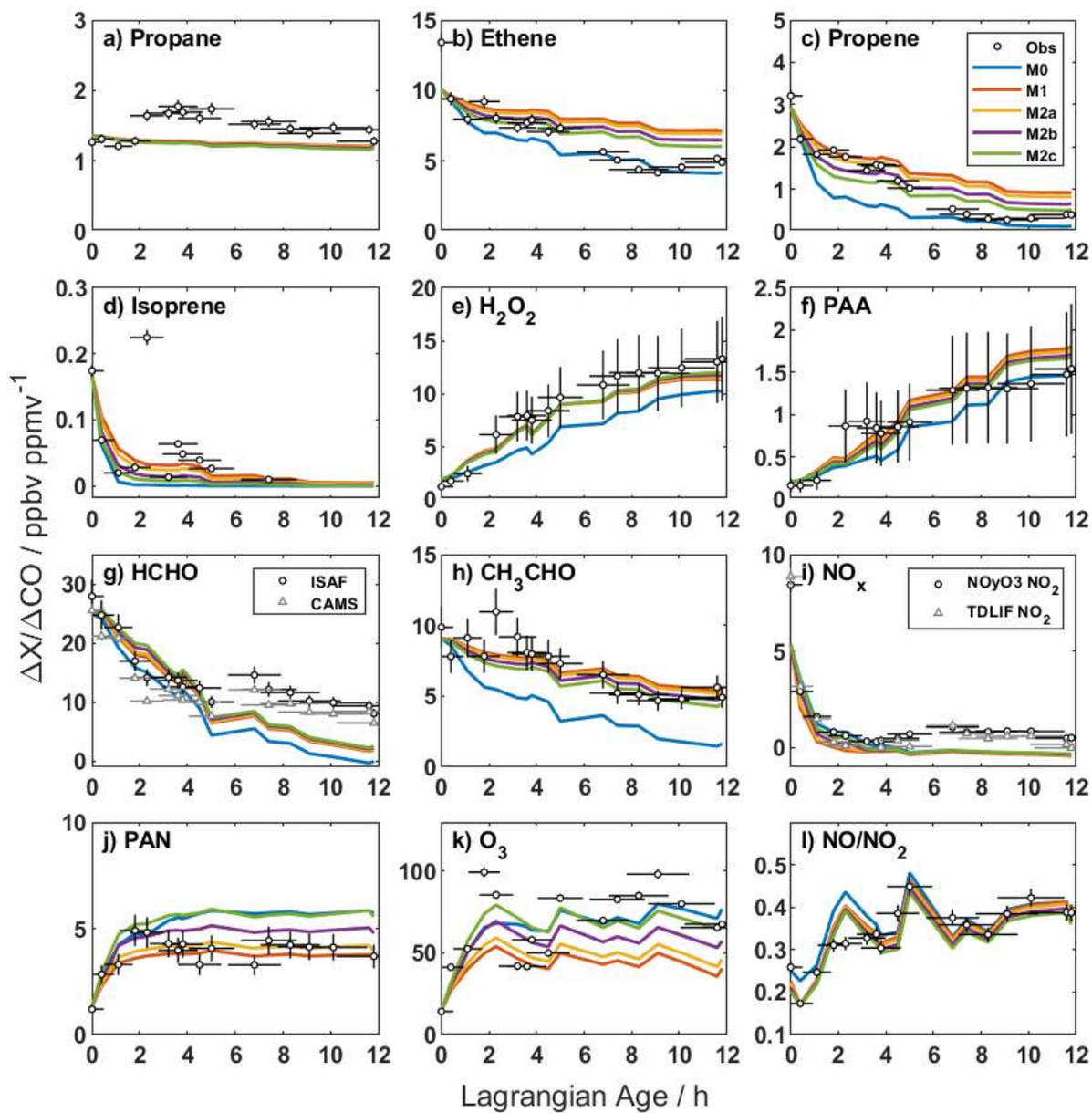


Figure S21. Age evolution of NEMRs for sensitivity simulations to assumed initial HONO concentration. Black circles are observations with their corresponding uncertainty due to measurement accuracy and age. Colored lines are model output from the base simulation (M0, blue), addition of unmeasured VOC (M1, red), and addition of unmeasured VOC plus primary HONO at mixing ratios of 5, 15, and 25 ppbv (yellow, purple, and green, respectively).

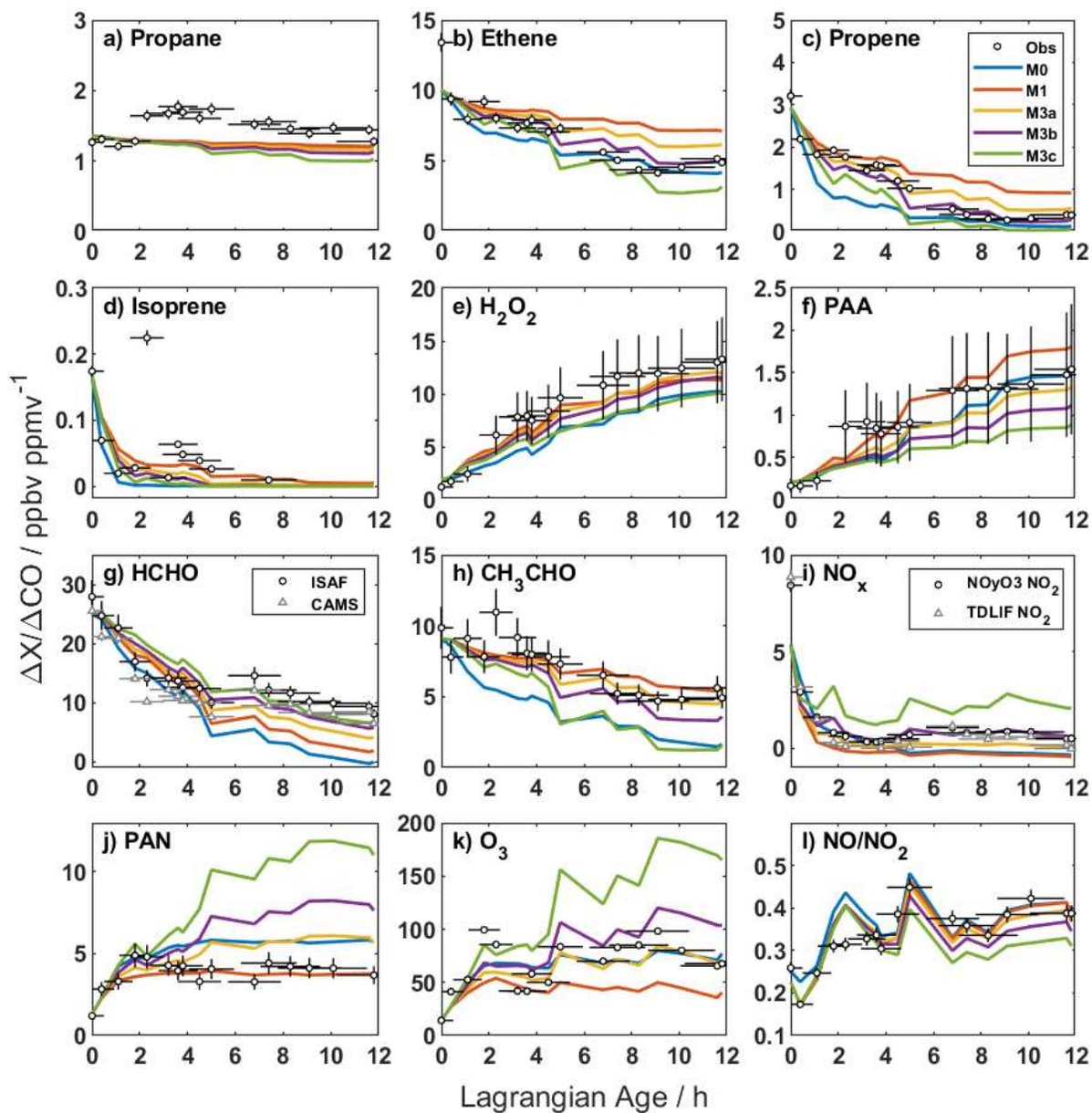


Figure S22. Age evolution of NEMRs for sensitivity simulations to particulate nitrate photolysis. Black circles are observations with their corresponding uncertainty due to measurement accuracy and age. Colored lines are model output from the base simulation (M0, blue), addition of unmeasured VOC (M1, red), and addition of unmeasured VOC plus  $\text{pNO}_3^-$  photolysis with rate multipliers of 0.5, 1, and 2 (yellow, purple, and green, respectively).

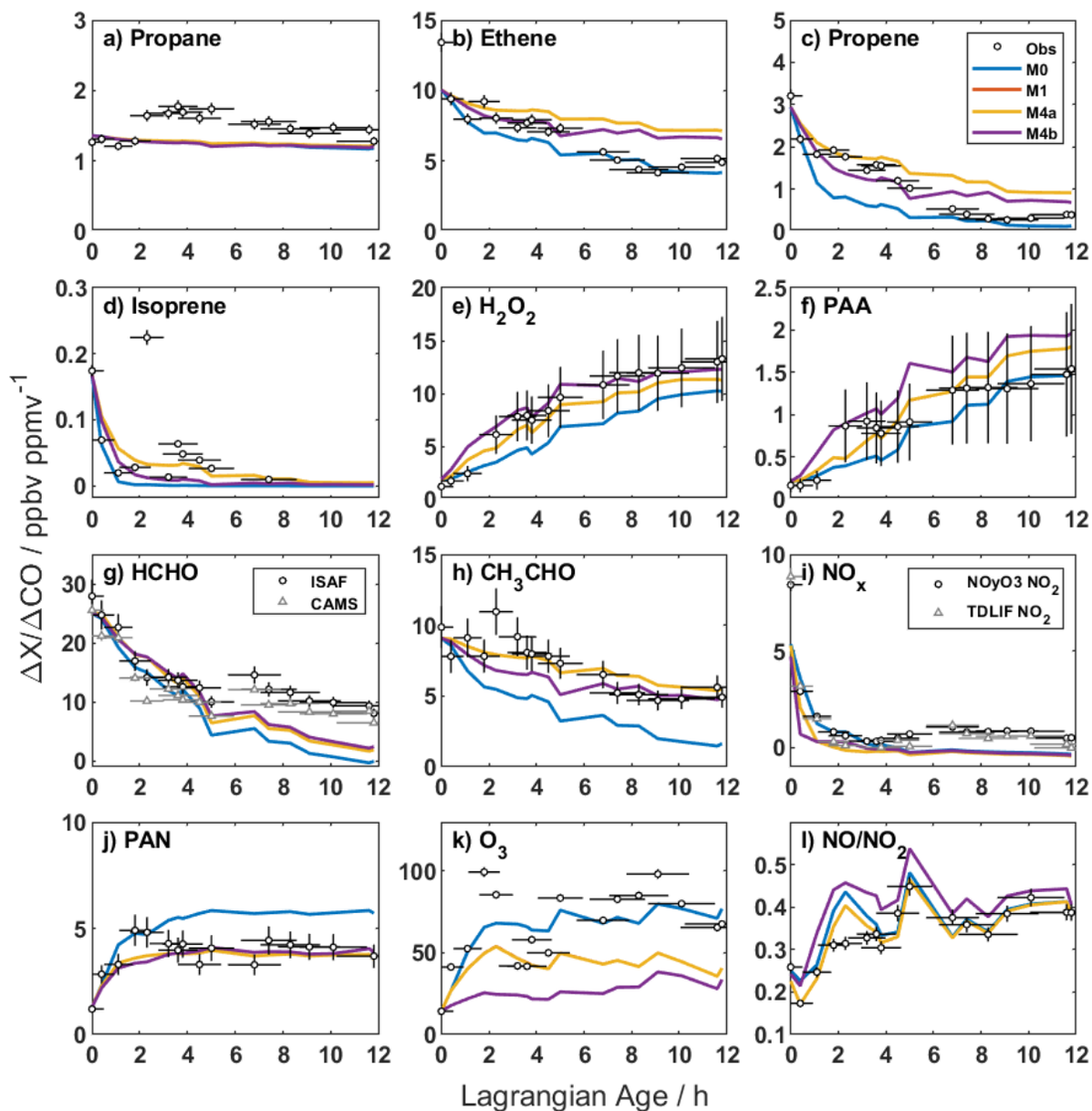


Figure S23. Age evolution of NEMRs for sensitivity simulations to heterogeneous reaction of  $\text{NO}_2$ . Black circles are observations with their corresponding uncertainty due to measurement accuracy and age. Colored lines are model output from the base simulation (M0, blue), addition of unmeasured VOC (M1, red), and addition of unmeasured VOC plus  $\text{NO}_2$  heterogeneous reaction with rate multipliers of 1 and 1000 (yellow and purple, respectively). Note that there is no visible difference in model output for simulations M1 and M4a.

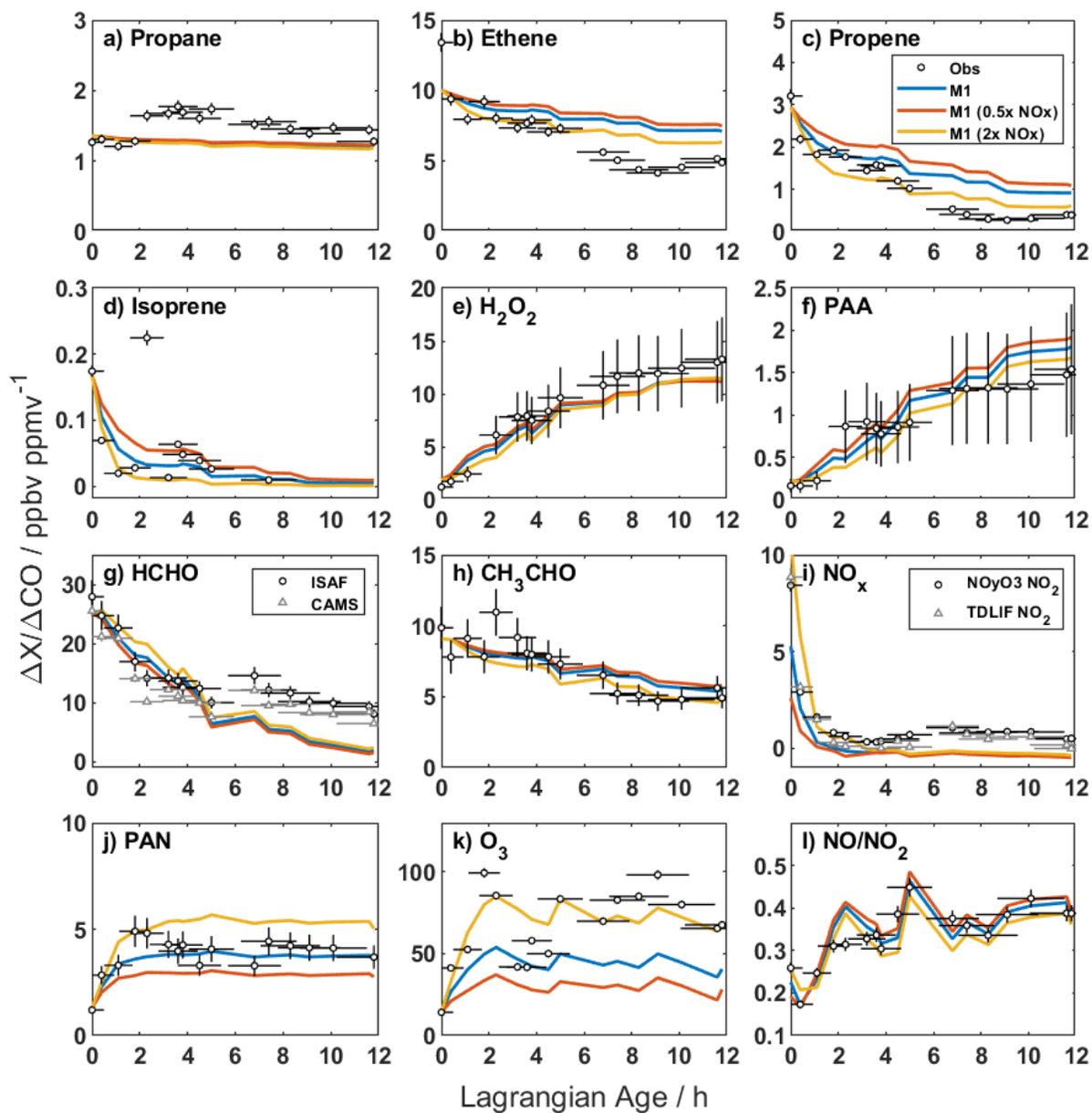


Figure S24. Age evolution of NEMRs for sensitivity simulations to initial  $\text{NO}_x$ . Simulation M1 (blue line) is modified by multiplying initial  $\text{NO}$  and  $\text{NO}_2$  mixing ratios by a factor of 0.5 (red) or 2 (yellow). Other details are as described in Fig. 2 of the main text.

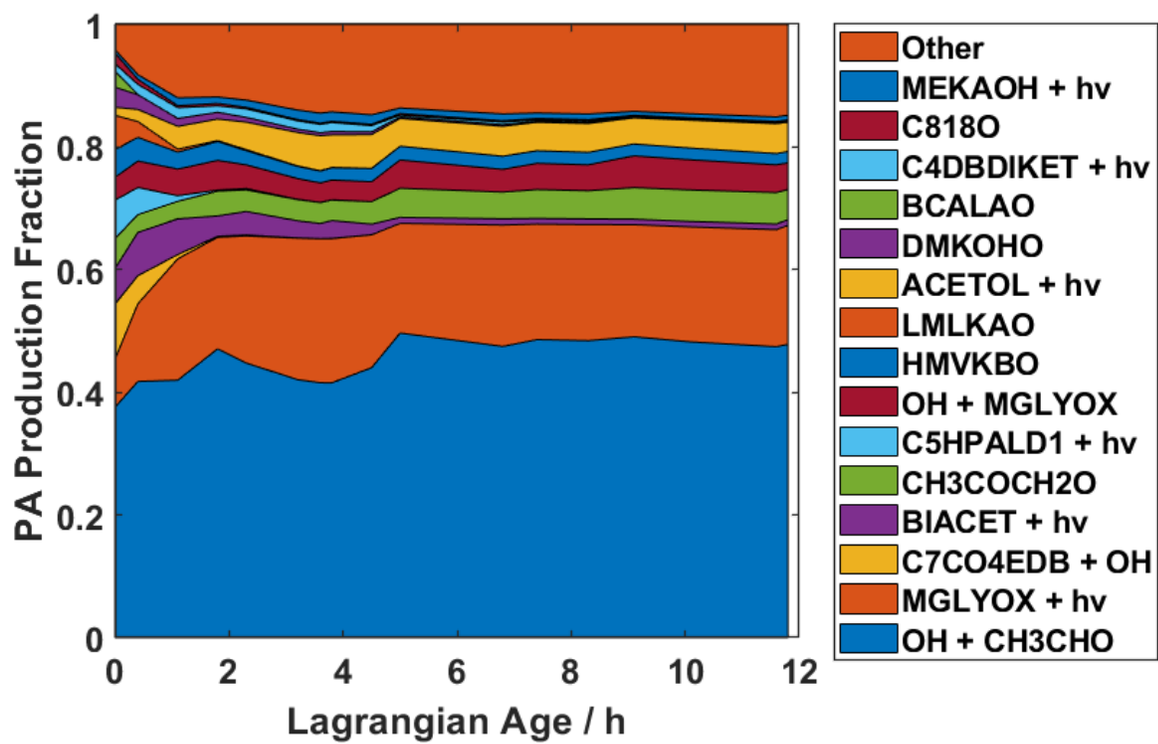


Figure S25. Fractional contributions to production of peroxyacetyl radical in simulation M3b.

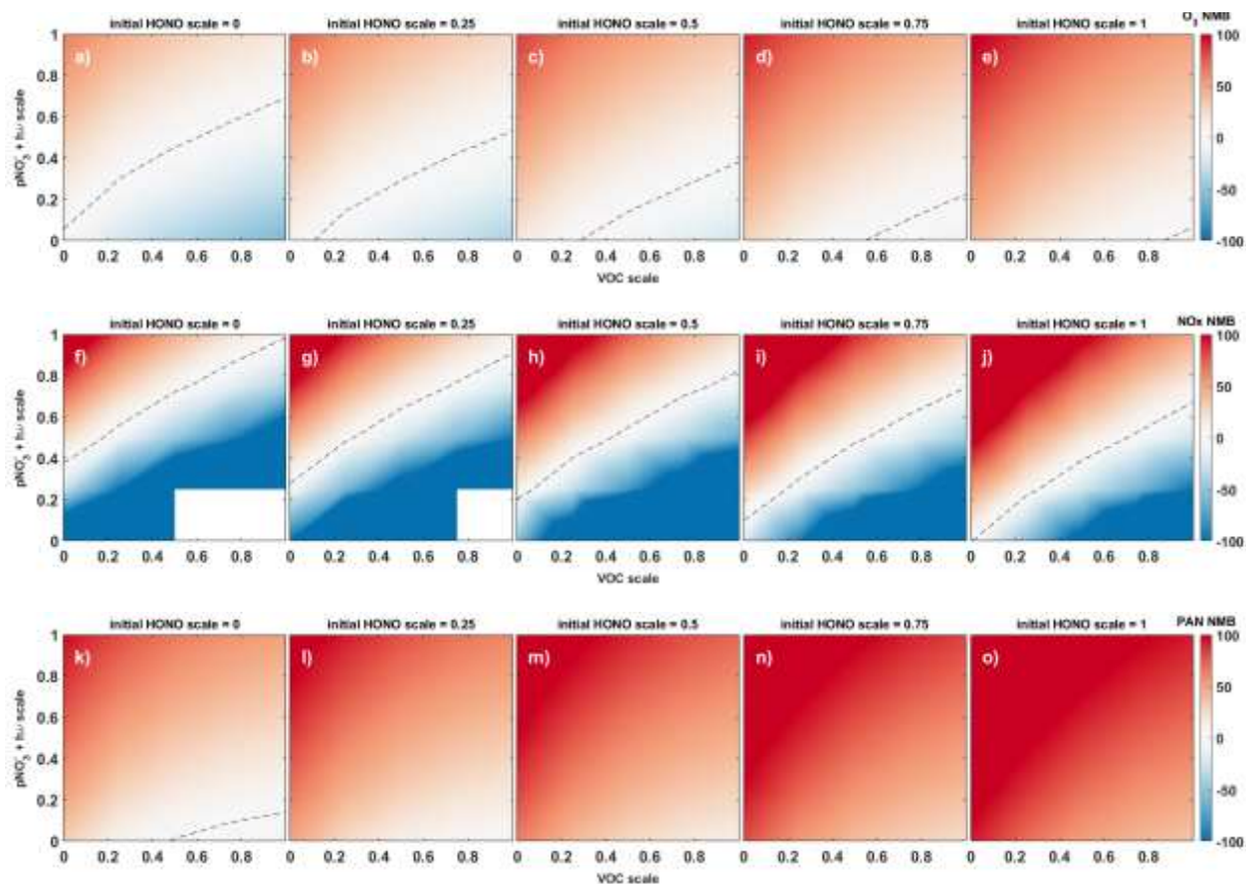


Figure S26. NMB of NEMRs for ozone (a-e), NO<sub>x</sub> (f-j), and PAN (k-o) for the sensitivity simulations described in SI Text S4. Simulations involve iteratively scaling unmeasured VOC (x-axis), pNO<sub>3</sub><sup>-</sup> photolysis (y-axis), and initial HONO (columns) by factors of 0, 0.25, 0.5, 0.75, and 1. Shading indicates NMB of simulation NEMRs against observations, ranging from negative (blue) to positive (red) values. Dashed lines indicate interpolated contours for NMB of zero, corresponding to values shown in Fig. S27.

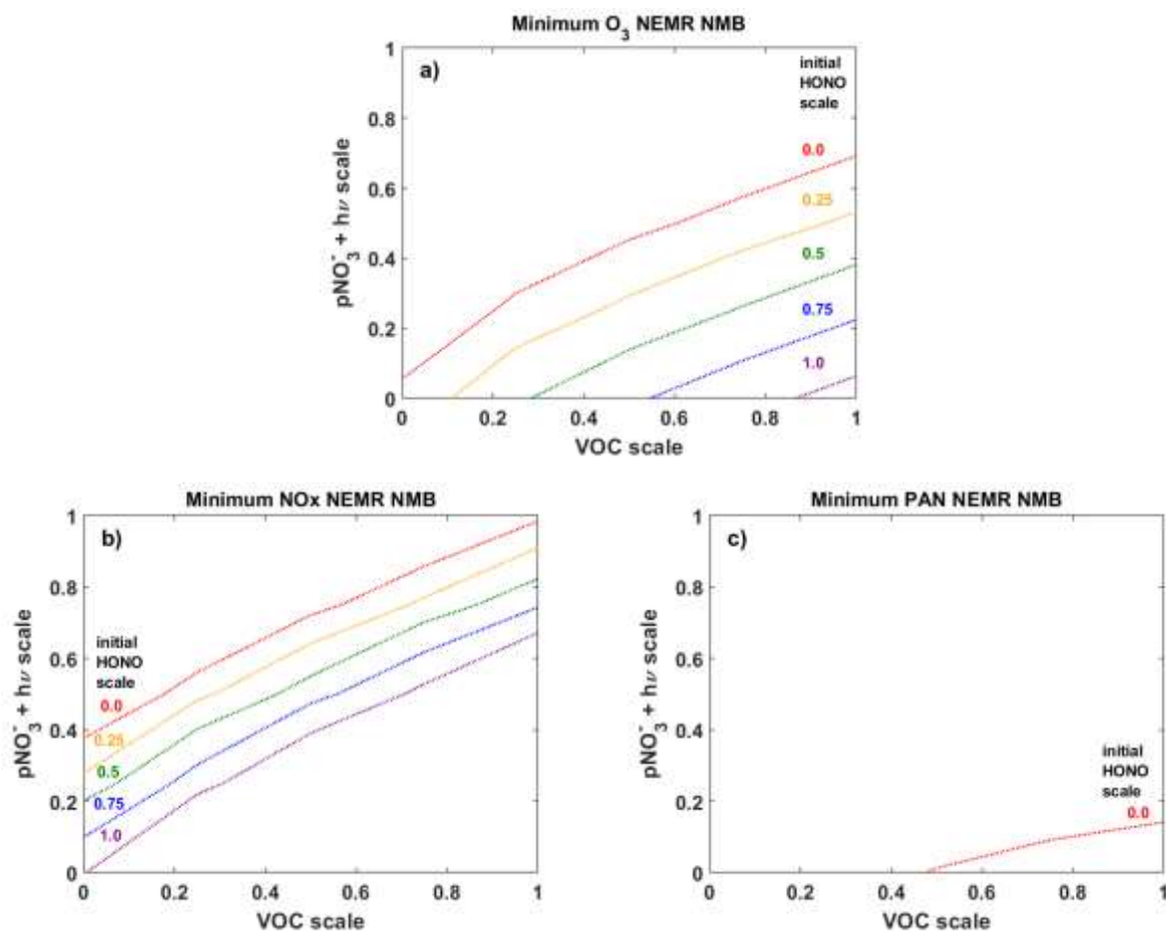


Figure S27. Isopleths for net-zero values of the normalized mean bias (NMB) for NEMRs of ozone (a), NO<sub>x</sub> (b), and PAN (c). Each colored dotted line represents a fixed scaling factor for initial HONO mixing ratios. The x-y coordinates for a point on a given line represent a combination of VOC and pNO<sub>3</sub><sup>-</sup> photolysis scaling factors that minimize the O<sub>3</sub> NEMR NMB. Isopleths are based on interpolation of results from the optimization simulations (Fig. S26).

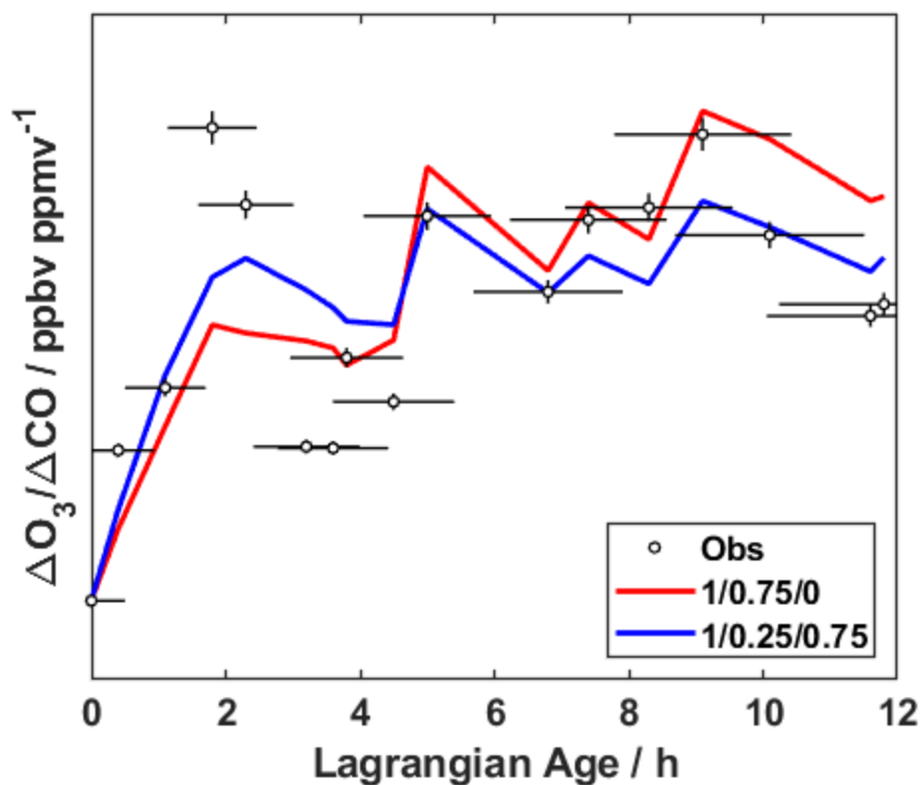


Figure S28. Comparison of observed O<sub>3</sub> NEMRs with those from two optimization simulations (SI Text S4) with near-zero ozone NMB (red: VOC scale = 1, pNO<sub>3</sub><sup>-</sup> + hv scale = 0.75, initial HONO scale = 0; purple: VOC scale = 1, pNO<sub>3</sub><sup>-</sup> + hv scale = 0.25, initial HONO scale = 0.75).