Volatile Organic Compounds and Ozone in Rocky Mountain National Park during FRAPPÉ

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Supplemental Information.

More detailed methods:

15 In situ GC

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The VOC analytical system was similar to those used in previous studies (Abeleira et al., 2017; Sive et al., 2005) and utilized a Qdrive 2s102K cryocooler (Chart, Inc., Troy, NY) that was capable of cooling a sample enrichment loop from 100 $^{\circ}$ C to -180 $^{\circ}$ C in 25 minutes for sample concentration. The loop was made from a 4 inch × 3/16 inch piece of stainless steel tubing and filled with 1 mm diameter deactivated glass beads (Ohio Valley, Marietta, OH) to provide an inert area for sample concentration during the trapping stage. The loop was heated to ~100 $^{\circ}$ C during the desorption, injection, and bake out stages

of the system.

The trapping procedure was initiated when the sample enrichment loop reached its initial set point temperature (-180 $^{\circ}$ C). A 1000 cm³ aliquot of ambient air was sampled through a heated stainless steel inlet line (70 $^{\circ}$ C) to remove O₃ prior to the sample passing through the enrichment loop at a rate of 200 cm³ min⁻¹. After the ambient air trapping procedure was

completed, 100 cm³ of ultra-high purity (UHP) helium was passed through the loop at a rate of 100 cm³ min⁻¹. The helium sweep was used to remove any residual O_3 , because condensed phase chemistry with O_3 may occur before or during desorption

(Koppmann et al., 1995). After the helium sweep was completed, the sample enrichment loop was isolated, and rapidly heated to 100 °C. Once the desorption temperature was reached, the sample was injected, and the GCs were triggered to start the temperature programs. UHP helium carrier gas flushed the contents of the sample enrichment loop through a 0.53 mm i.d. Silonite-coated transfer line to the splitter box. At the splitter box, the sample was quantitatively and reproducibly split into

5 four sample sub-streams. Each of the four sub-streams was transferred to a separate GC separation column/detector pair.

Two Shimadzu GC-17A gas chromatographs (Shimadzu Scientific, Columbia, MD) each housed two different separation columns. One 50 m × 0.53 mm i.d., 10 μ m film thickness CP-Al₂O₃/Na₂SO₄ PLOT column, one 60 m × 0.32 mm i.d., 1 μ m film thickness VF-1ms column, one 60 m × 0.25 mm i.d., 3 μ m film thickness CP-PoraBond Q column, and one 60 m × 0.25 mm i.d., 1.0 μ m film thickness VF-1701 column (Varian Inc.) were used for trace gas separation. The PLOT, VF-1ms, and PoraBond Q columns were each connected to a Flame Ionization Detector (FID) for analysis of C₂-C₇ and C₆-C₁₀ non-methane hydrocarbons (NMHC), and selected oxidized VOCs (OVOCs: methanol, ethanol, acetaldehyde, acetone and MEK) measurements, respectively. The VF-1701 column was plumbed into an Electron Capture Detector (ECD) and used for measuring C₁-C₂ halocarbons and C₁-C₅ alkyl nitrates.

Two different whole air standards were alternately analysed every 10 samples throughout the campaign. The measurement

15 precision, represented by the relative standard deviation (RSD) of the peak areas for each compound in the standards, was 1-8% for the NMHCs, 3-11% for halocarbons, and 3-5% for alkyl nitrates. A list of VOCs measured during this study using the *in situ* GC is shown in Table 1.

Canister GC System

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20 The analytical system and methodology used for this study are similar to those used in previous studies (Russo et al., 2010; Sive, 1998; Zhou et al., 2010). For each sample, a 1265 cm³ (STP) aliquot of air was trapped on a glass bead filled loop immersed in liquid nitrogen. After the sample was trapped, the loop was isolated, warmed to 80 °C and injected. The carrier gas (UHP helium) flushed the contents of the loop and the stream was split into five, with each sub-stream feeding a separate GC column/detector pair as follows: (1) a CP-Al₂O₃/Na₂SO₄ PLOT column (Varian-Chrompack; 50 m × 0.53 mm i.d., 10 μm film thickness) connected to an FID was used to measure C₂-C₇ NMHCs; (2) a VF-1ms column (Varian-Chrompack; 60m × 0.53 mm i.d., 10 μm

0.32 mm i.d., 1 μ m film thickness) connected to a FID measured C₄-C₁₀ NMHCs; (3) a CP-PoraBond Q column (Varian-Chrompack; 25 m × 0.25mm i.d., 3 μ m thickness) coupled to a Restek XTI-5 column (Restek; 30 m × 0.25 mm i.d., 0.25 μ m film thickness) connected to an FID was used to measure selected OVOCs; (4) an OV-1701 column (Ohio Valley Specialty Chemical; 60 m × 0.25 mm i.d. 1 μ m thickness) connected to an ECD was used to measure C₁-C₅ alkyl nitrates and C₁-C₂ halocarbons; (5) and an OV-624 column (Ohio Valley Specialty Chemical; 60 m × 0.25 mm i.d., 1.4 μ m thickness) connected

5 halocarbons; (5) and an OV-624 column (Ohio Valley Specialty Chemical; 60 m \times 0.25 mm i.d., 1.4 μ m thickness) connected to an MS measured C₆-C₁₀ NMHCs, C₁-C₂ halocarbons, and was used to measure select OVOCs and reduced sulphur compounds.

Trail Ridge Road Ozone Calibrations

10 Calibrations of the POMS unit were consistent with the US EPA guidance and were performed at Air Resource Specialists, Inc. (ARS) immediately prior to deployment and again after the campaign. The calibrations were performed by comparing the instrument output to an ultraviolet (UV) photometer transfer standard that is traceable to EPA. The instruments were challenged at multiple O₃ concentrations (including zeros) and evaluated by the acceptance criteria which includes the percent difference of the analyser's response during the multipoint challenges and the instrument's linearity. The ARS calibration 15 acceptance criteria for O₃ is ≤3% for the multipoint challenges and ≤1% for the linearity. Further details regarding the calibrations of the O₃ and meteorological measurements can be found in the Gaseous Pollutant Monitoring Program 2015 Quality Assurance Project Plan (https://ard-request.air-resource.com/Project/documents.aspx).

Details for calculating air mass photochemical age

The dominant source of alkyl nitrates is the photochemical production from hydrocarbons. Equations 1-6 show the chemical mechanism for alkyl nitrate production.

$$RH+OH \rightarrow R+H_2O (k_1, \alpha_1)$$
(1)

$$R+O_2 \rightarrow RO_2 (k_2)$$
(2)

$$RO_2+NO \rightarrow RO+NO_2 (k_3, 1-\alpha_4)$$
(3)

$$RO_2+NO \rightarrow RONO_2 (k_4, \alpha_4)$$
(4)

$$RONO_2 + hv \rightarrow RO+NO_2 (j_5)$$
(5)

$$RONO_2+OH \rightarrow \text{ products } (k_6)$$
(6)

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 k_1, k_2, k_3, k_4, k_6 are reaction rate constants, α_1 and α_4 are the reaction branching rates and j_5 is the photolysis rate constant.

$$\frac{(\text{RONO}_2)}{(\text{RH})} = \frac{\beta k_A}{(k_B - k_A)} (1 - e^{(k_A - k_B)t}) + \frac{(RONO_2)_0}{(RH)_0} e^{(k_A - k_B)t}$$
(8)

15 $k_A = k_1[OH], k_B = k_6[OH] + J_5, \beta = \alpha_1 \alpha_4, (RONO_2)_o/(RH)_o$ is the initial alkyl nitrate/parent hydrocarbon ratio. If (RONO_2)_o /(RH)_o =0, the solution to equation describes the time evolution of the alkyl nitrate/parent hydrocarbon ratio based solely on gas phase hydrocarbon chemistry (Bertman et al., 1995).



Figure 1S Time series of VOCs measured at a ROMO site during FRAPPÉ. a) ethane and propane, b) i-butane and n-butane, c) ipentane and n-pentane, d) alkyl nitrates, e) C₂Cl₄ and C₂HCl₃, f) ethyne, ethane, and propene, g) benzene and toluene



Figure 2S Time series of VOCs measured at a ROMO site during FRAPPE. a) isoprene and PTR-MS isoprene, b) α-pinene and βpinene, c) CHBr₃ and CH₂Br₂, d) CH₃I and CHBrCl₂, e) ethylbenzene, m+p-xylene, o-xylene, f) acetone, g) acetaldehylde



Figure S3a. VOC canister data plotted by longitude and date with 2-BuONO₂ mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3b. VOC canister data plotted by longitude and date with 2-PenONO₂ mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3c. VOC canister data plotted by longitude and date with α -pinene mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3d. VOC canister data plotted by longitude and date with benzene mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3e. VOC canister data plotted by longitude and date with C_2Cl_4 mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3f. VOC canister data plotted by longitude and date with C₂HCl₃ mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3g. VOC canister data plotted by longitude and date with DMS mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3h. VOC canister data plotted by longitude and date with ethylbenzene mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3i. VOC canister data plotted by longitude and date with ethyne mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3j. VOC canister data plotted by longitude and date with isoprene mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3k. VOC canister data plotted by longitude and date with propane mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.



Figure S3I. VOC canister data plotted by longitude and date with toluene mixing ratio represented by the size of each point and the i- to n-pentane represented by the colour. Locations correspond to transect samples shown in Figure 2.

	Alpine	Visitors	Forest (Canyon	Rainbov	w Q.Ve	Many Par	ks Curve	ГО.	Ş	Beaver N	/eadows	Eastes Pa	rk Resort	Eof	Take	BIG Tho	mpson	Camp sit	e CDPHE	City Park Depiter
	Average	Stdev	Averee	Striev	<u>Åversce</u>	C+C-V	<u>Averace</u>	Ctolev	Avera	Striev	Average	Ctules.	<u>Averece</u>	2 dev	AVE FOR	Strieu	Averes e	Std av	Averace	Stdev	
e nerte		100		464.6		404.0	1604.0	OVE O	1720 F	726.1	3077.0	1100 7		0150	0000	1000 E	1100.0	10010	7630 F	200 6	1704.0
	21221	2880	10011	570.2	10101		- CE0	0.000	1143.7	675.7	1394.8	1214.0	876 B	674.1		1607.7	ander o	2512.2	1100.7	167.4	
i-butane	60.1 1	36.5	78	49.7	80.2	6.66	253.6	149.0	126.6	105.5	220.6	232.2	158.0	115.0	355.0	388.1	519.6	428.8	216.8	65.4	225.4
n-butane	129.4	95.7	253.2	216.2	198.8	90.5	638.5	411.3	300.5	278.3	565.3	628.5	400.5	316.9	87 9.9	914.7	1390,9	1219.5	579.9	219.1	385.5
ethyne	100.5	28,9	107.2	29.5	110.3	43,0	459.8	604.9	110.8	30.2	145.5	75.3	189.3	150.4	178,4	78.0	292.8	229,4	9 88 0	135.1	303.0
í-pentane	43.5	22.7	260.7	395.8	71.1	26.0	629.8	589.1	97.8	72.0	239.2	2184	362.1	385.0	436.1	577.9	512.6	334,9	687.7	464.7	395.3
n-pentane	38.7	24.6	92.6	80.3	57.4	17.3	375.0	320.0	95.1	78.2	191.7	192.8	220.8	282.5	341.9	388,8	494.2	417.4	355.6	228.3	233.8
n-hexane	13.5	9.2	28.2	14,9	17.4	5.7	175.6	177.2	26.0	18.3	58.4	41.0	65.9	71.4	87.6	90,7	136,9	105.2	153.5	96.2	82.4
n-heptane	5.5	2.3	12.3	с,9 1	9.7		781	74,8	11.2	4.1	16.8	11.9	26.9	28.3	31.7	25.8	4 ∿	29.2	301.8	355.0	37.0
n-nonane	16.2	4.2	15.2	6.3	16.7	9.5	42.4	12.9	243	12.7	20.2	7.4	23.0	10.1	25.1	13.0	27.5	10.3	37.3	26.0	17.7
benzene	22.4	7.8	34.6	13.0	26.3	11.1	123.3	92.5	31.3	9.9	82.4	71.2	62.7	348 8	63.6	6 %	102.5	71.8	231.8	117.2	107.9
toluene	17.4	6.4	38.0	28.9	26.2	82	339.4	400.3	346 8	14,9	75.0	45.8	114.7	95.9	77.0	50.9	1984	382.9	1608.0	1758.6	189.9
e thy lb en zene	3.1	1.4	5.5	47	3.2	0.7	56.4	66.5	40	0.9	11.7	8 6	16.4	15.2	10.0	6.2	20.9	24.5	69.7	45.5	31.2
m+p-xylene	7.9	6'8	15.8	14,4	0 0 00	2.7	201.9	225.7	11.4	5.0	38.1	31.3	541	53.0	249	19.7	63.6	97.9	228.1	165.1	104.2
o-xylene	3.6	1;8 1	7.0	6.1	4	1.8	76.1	83	5.2	1.8	14.5	11.4	20.5	189	11.1	7.8	28.9	26.8	8 .9	61.9	37,4
styrene	2.8	1.3	8.2	2.1	1 1 1	0.7	8 9 10	3.6	8.2	0.6	4	4	3.7	2.1	4	8.5 2	6.9	6.8 9	12.5	7.9	5,9 2,9
Isoprene	287.6	66,6	272.7	174.3	316,3	200.6	191.2	172.9	332.1	295.1	506.9	486.2	153.8	103.6	117.8	102.3	599,9	748.1	37.9	4 0	159.2
a-pinene	49.9	31.0	45.8	7.0	81.7	27.6	69,4	4	₽ 9	17.6	80.5	8. 6	34.1	31.7	53.9	128,8	346 8	58.0	42	2,4	36,9
b-pinene	346 8	22.7	38.5	9,1	101.9	69.0	40.9	14,4	66.7	30.0	74.3	49.5	36.6	38.8	en K	131.3	33,4	74.0	1.2	80	51.1
DMS	0.7	0.2	1.2	0.2	0.6	0.3	0'0	0.2	1.0	0.4 4	1.6	0.2	1.7	1.3	1,8	1.3	0.4	3.2	1.0	0.1	3.2
CH ₂ Cl ₂	35.2	9'8	37.3	3,7	36,9	6.7	37.6	8,4	30.2	5.8 9	35.5	4.6	9°8	5.0	36.6	e,4	46.0	41.4	37.6	1.4	6 8 6
C ₂ HCl ₃	0.2	0'0	0.2	0.1	0.1	0.1	0.3	0.2	0.3	0.4	0.5	0.3	0.5	0.5	0,6	0,6	1.1	1.2	1.4	0.2	8 0 0
c₂d₄	2.1	0,4	2.1	0.5	2.6	0,6	2,6	0'9	2.2	0'9	2.7	1.1	3,0	1,4	45	3.1	5.6	2.7	10.3	6'0	46
MeONO ₂	4,6	4.0	5.0	0. 4	4 0	0.8	4 80	0.5	6.1	0.9	5.1	0.7	6.0	1.2	6,4	1.7	6.7	2.3	7.7	1.5	6.2
EtoNo ₂	2.7	80	8.0 8	1.0	3.1	80	80 ന്	1.5	4	1.0	ъ.	1.9	4 4	2.2	ы С	2.5	6.0	2.9	6.9 9	4	ຍ ຍໍຍ
I-Pr@NO2	7.2	3.1	ຕ ຜ	4 0	9.9	4	10.9	6.2	13,4	г. 4	13.4	8 21	12.2	9.1	17.7	9.5	21.8	13.0	20.0	7.0	10.4
n-PrONO ₂	1.6	0.7	1.8	80	1.8	0.6	2.2	1.0	3.0	4.1	2 .6	1.1	2.8 1.8	2.3	4	2,4	5.1	3.2	46	1.7	2.2
2-BUONO2	7.4	4 8	7.7	5.1	9.9	5.7	11.5	8.0	14.6	9.0	17.3	13.8	12.4	10.6	22.7	16.7	30.7	20.4	19.7	1.2	13.2
3-PenONO2	1.4	0.6	2.4	1.9	2.0	0.8	3.1	2.1	3.6	2.3	4 0	2.2	3.1	2.6	6'S	4,9	7.6	4.7	5.0	0.0	2.6
2-PenONO2	1.7	1.4	2.3	1.6	9. 1	1.1	3'0 '	2.0	9'B	8.0	5.5	3,0	3,4	3,1	7.8	5.7	10.5	6'9	5.8 8	0.1	6,0
acetone (ppbv)	9.7	9'8	13.2	6,6	10.6	5.4	22.0	11.4	16.2	5,6	14.3	40	15.4	7.7	17.1	7.0	15.1	7.4	16.0	9.1	6,9

Table S1: Statistics of VOCs measured in transect samples during FRAPPE by site.

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