



Supplement of

Secondary organic aerosol formation from α -pinene, alkanes, and oil-sands-related precursors in a new oxidation flow reactor

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S1. OFR comparisons

The design of the ECCC-OFR (Environment and Climate Change Canada oxidation flow reactor) was partially based upon recent OFRs designs (Lambe et al., 2011;Huang et al., 2017;Simonen et al., 2017) with several small specific differences. The specific differences and similarities between the various reported OFRs is described below.

Comparison with the PAM (Potential Aerosol Mass) reactor (Lambe et al., 2011):

The ECCC-OFR utilizes a conical diffusion inlet, while PAM employs a straight inlet. The straight inlet is likely to lead to some jetting and recirculation, while cone inlet should have improved fluid dynamics (Huang et al., 2017;Mitroo et al., 2018). The lamps of the ECCC-OFR are located on the outside of the reactor, while the lamps for the PAM are located inside the reactor which can increase surface-to-volume ratio and hence wall losses. Finally, both OFRs sample from the center with appropriate side flows as exhaust, however the ECCC-OFR uses a sampling tube which is 12.7 cm offset from the end of the OFR.

Comparison with CPOT (Caltech Photooxidation Flow Tube) (Huang et al., 2017):

Both of these OFRs use a conical diffusion inlet, with the lamps of both located out of the reactor. The full cone angle of CPOT (15°) is smaller than ECCC-OFR (30°) . The ECCC-OFR samples from a center port, while CPOT samples all gases at the exit cone. The CPOT has a larger surface-to-volume ratio and a longer residence time (~1500 s) compared to the ECCC-OFR, which may lead to larger wall losses of particles and organic vapors.

Comparison with TSAR (TUT Secondary Aerosol Reactor) (Simonen et al., 2017):

Both of these OFRs make use of a conical inlet, with lamps located on the outside of the reactor. These OFRs both sample from the OFR center-line, with sampling tubes offset from the end of the reactors. The TSAR is designed for rapidly changing sources, with a volume that is smaller (3.3 L) and a residence time which is shorter (37 s) (Simonen et al., 2017). As a result, the OH concentration within the TSAR will be higher at the same OH exposure. The LVOCs inside the reactor can be consumed by high concentration of OH, or exit the OFR because due to insufficient time to condense on aerosols (Simonen et al., 2017).

Comparison with PEAR (Photochemical Emission Aging Flow Tube Reactor) (Ihalainen et al., 2019):

Both of these OFRs use a conical inlet, and they both sample from the center-line with appropriate side flows as exhaust. The full cone angle of PEAR (28°) is similar to ECCC-OFR (30°). The lamps of the ECCC-OFR are located on the outside of the reactor, while the lamps for the PEAR are located inside the reactor. The volume of PEAR (139 L) is larger than ECCC-OFR, and the flow rate is also larger, resulting in similar residence time.

S2. CFD simulations and RTD measurements

To assess the near laminar flow of the ECCC-OFR, computational fluid dynamics (CFD) simulations were performed using ANSYS Fluent software (Version 2019 R2) in three dimensions to characterize the flow field inside the ECCC-OFR. Hybrid tetrahedral–hexahedral mesh consisting of 5.7×10^5 computation cells were used.

Turbulence was modeled using a realizable k-epsilon model. The simulation results are shown in Fig. S4. It is shown in Fig. S4a that the flow velocity distribution in the reactor is generally uniform. A high velocity is observed only near the inlet, but reduces to the average velocity in the conical diffuser. The velocity distribution here indicates that jetting is much weaker in ECCC-OFR compared to PAM (Mitroo et al., 2018). Fig. S4b indicates that the flow field is quite good in ECCC-OFR, with a small recirculation zone, similar to previous studies using a conical diffusion inlet (Huang et al., 2017;Ihalainen et al., 2019), but much better than PAM (Mitroo et al., 2018).

The residence time distribution (RTD) was measured for ECCC-OFR and compared to ideal laminar flow in Fig. S5. The RTD was characterized by injecting a constant flow rate of CO₂ (10 s) into the OFR. The CO₂ concentration was then monitored from the sampling outlet of OFR with a CO₂ analyzer (Li-Cor LI-840A). The RTD was calculated from the differential CO₂ as a function of time elapsed since the start of injection. Fig. S5 indicates that the residence time associated with the CO₂ maximum intensity for the measured RTD and the ideal laminar flow RTD are in good agreement, and improved over the PAM and TPOT (Lambe et al., 2011). The shape of measured RTD before ~100 s is similar to the ideal laminar flow RTD, but slightly wider, which is likely due to dispersion. After ~100 s, the measured decrease in CO₂ is slower than for ideal laminar flow, which is likely due to recirculation in the OFR (Fig. S4).

S3. OS sample details

OS ore samples were collected directly from the CNRL-Horizon OPP (Ore Preparation Plant) surge bin #1 on 24-Sep-2016, and stored in a freezer (-10 °C). Once the oil sands is hauled out of the mine, the ore is first processed in an Ore Preparation Plant (OPP), where clumps of oil sands are broken down (OPP-dry) and mixed with water to produce a pumpable slurry (OPP-wet). Mined oil sands can contain large chucks of bitumen, ice and fine solids. The crusher and sizer of the OPP-dry process breaks these clumps down into a loose, unconsolidated material (OSM, 2019). The OS ore sample here is from the surge bin after the OPP-dry process, which can be considered as "unprocessed" OS ore.

Bitumen was collected from the bitumen froth from the CNRL-Horizon plant on 25-Sep-2016, and stored in a freezer (-10 °C). Bitumen froth is extracted from the oil sands slurry (after OPP-wet) through a simple water-based gravity separation process, which occurs in a large cone-bottomed vessel. The bitumen attaches to free air bubbles and rises to the top of the vessel, forming an intermediate froth product (the bitumen samples here) (OSM, 2019). The heavy sand sinks to the bottom and is pumped out to the tailings plant (OSM, 2019). Bitumen froth contains about 50-60% bitumen, 30-40% water and 10-15% fine solids.

Naphtha solvent was collected from the CNRL-Horizon facility on 25-Sep-2016, and stored in a refrigerator (~4 °C). Naphtha, a type of solvent/diluent used in the extraction and dilution of bitumen, is a mixture of C_3 - C_{14} hydrocarbons with major fractions of *n*-alkanes (e.g., heptanes, octane, nonane) and aromatics (e.g., benzene, toluene, ethylbenzene, xylenes) (Siddique et al., 2006).

Dilbit is diluted bitumen, and was collected from the CNRL-Horizon facility on 25-Sep-2016, and stored in a refrigerator (~4 °C). Though bitumen froth contains only ~60% bitumen, it acts almost like a single-phase fluid. The water and bitumen are closely intermixed, with fine solids trapped within the viscous mixture (OSM, 2019). Therefore, bitumen froth is further processed by froth treatment, which is accomplished through the addition of a solvent or diluent (e.g., naphtha). This solvent/diluent dilutes the bitumen, producing a less viscous, lighter product, with a density lower than water. The viscosity of the diluted bitumen also drops significantly, which releases the trapped fines. The diluted bitumen floats to the top of the gravity separation vessel, leaving the fines to settle to the bottom of the water phase (OSM, 2019). Bitumen itself is extremely heavy and viscous, and cannot be transported in pipelines. The diluted bitumen also makes the transport in pipelines possible for bitumen products.

Tailings pond water was collected from Suncor pond 2/3 on 23-Aug-2017, and stored in a refrigerator (~4 °C). Tailings are a mixture of water, fine silts, residual bitumen, salts and soluble organic compounds. They also include solvents (e.g., naphtha) that are added to the oil sands during the separation process. Suncor tailings pond 2/3 is considered as one of the most active tailings ponds in the Alberta OS region (Small et al., 2015).

S4. Volatility distribution of OS-related precursors

The total ion chromatogram of the OS-related precursors as a function of retention time for GC-MS are shown in Fig. S3. The retention time for *n*-alkane standards are also shown. Here we assume that the unresolved complex mixture between C_n to C_{n+1} alkanes have lower volatility than C_n and higher volatility than C_{n+1} , and bin the data (after response factor correction for compounds with different volatilities) by carbon number in Fig. 1a. According to the relationship between carbon number and volatility (Donahue et al., 2006), the data are also binned by effective saturation concentration (C^*) in Fig. 1b. For OS ore, the volatility covers a wide range of carbon number from 6 to 15. Based on the mass fraction of each carbon number, we calculated the average carbon number of OS ore, which is ~10 (9.79). As a commonly used solvent in OS processing, the naphtha has a narrow distribution mainly at C_6 - C_8 , with a peak at C_7 . The VD of OS ore and bitumen are very similar at a retention time of > 4 min (Fig. S3), which corresponds to approximately C₈, and indicates that the processed bitumen is essentially oil sands mixed with some solvents during the processing. Bitumen, dilbit and tailings pond water contain varying amounts of the similar solvents, consistent with the dominant volatility at C_7 for these samples. Based on the VDs in Fig. 1, the fraction of solvent within the emissions follows the order of naphtha (100% solvent) > dilbit > tailings pond > bitumen > OS ore (0% solvent). This corresponds to a non-solvent fraction in the sample vapors in the order of OS ore > bitumen >tailings pond > dilbit > naphtha (0% heavy oil). From the VDs in Fig. 1, it is also found that the vapors from OS ore and bitumen contain ~15% and ~10% IVOCs (carbon number ≥ 12 , C* $\leq 10^6 \ \mu g \ m^{-3}$), respectively, while other precursors are almost 100% VOCs.

We note that VD of OS ore measured here is somewhat different from that previously measured (Liggio et al., 2016), which were mainly in the C_{12} to C_{16} range. However, the VD for OS ore in this study is mainly from C_8 to C_{13} with an average value of C_{10} , which are more volatile than the previous study. This difference is likely due to the nature

of the samples themselves, as the previous OS ore sample was collected off-site, in an area not associated with active mining, and exposed to the atmosphere for a long period. It is highly likely that the majority of the more volatile components had long since evaporated. Conversely, the current study utilized samples taken directly from the active mining operations (>50 m below ground) and immediately archived at -10 °C. In addition, the VD derived here are consistent with those from more recent aircraft measurements around active mining operations.

S5. Secondary particle yield calculations

The yield of sulfuric acid $(Y_{H_2SO_4})$ was calculated using the mass fraction of H₂SO₄ in particles $(x_{H_2SO_4})$, the SMPSmeasured particle volume $(V_{H_2SO_4:H_2O}, \text{ nm}^3 \text{ cm}^{-3})$, the density of the particles $(\rho_{H_2SO_4:H_2O}, \text{ g cm}^{-3})$, and the measurement of the reacted SO₂ (ΔSO_2 , ppb), using an approach which has been described in detail previously (Lambe et al., 2011):

$$Y_{H_2SO_4} = \frac{x_{H_2SO_4} \times V_{H_2SO_4 \cdot H_2O} \times \rho_{H_2SO_4 \cdot H_2O}}{3.95 \times \Delta SO_2}$$

The $x_{H_2SO_4}$ and $\rho_{H_2SO_4:H_2O}$ were estimated using the Extended Aerosol Inorganic Thermodynamic Model (E-AIM) I (http://www.aim.env.uea.ac.uk/aim/aim.php) (Carslaw et al., 1995). The dry yield of H₂SO₄ from OH oxidation of SO₂ is 3.95 µg m⁻³ per ppb SO₂ reacted (Lambe et al., 2011); hence $Y_{H_2SO_4}$ is expected to be 100% without wall losses.

The SOA yields (Y) are calculated using the mass concentration of organic aerosols (ΔM_0) and reacted parent hydrocarbons (ΔHC):

$$Y = \Delta M_0 / \Delta HC$$

Here, ΔM_0 is calculated by multiplying the integrated volume concentration by the effective particle density (see Methods). The calculation for ΔHC is based on the measured THC. From the measured total carbon (converted CO₂) mixing ratio, one can derive the carbon mass concentration [C]. For single precursors (which contains only C and H), the precursor mass concentration [HC] is calculated by:

$$[HC] = [C] \times (1 + \frac{H/C}{12})$$

The reacted hydrocarbon mass concentration is then calculated by:

$$\Delta HC = [HC] \times (1 - e^{(-k \cdot [OH] \cdot t)})$$

where k is the second-order rate constant of the precursor with OH; [OH]·t is the OH exposure, which is measured off-line via the decay of CO (see Methods).

For OS-related precursors, the molecular composition (and hence H/C and k) is unknown, hence there are several assumptions when applying the above steps. The H/C of the OS precursors (H/C_{OS}) is estimated via the intercept of the linear fit of the Van Krevelen diagram of OS SOA (Fig. S10, Table S1). The intercept represents the H/C ratio

when O/C ratio is zero, which is expected to be similar to the precursor H/C. However, based on the intercept and the precursor H/C of the alkanes (Table S1), we find that the intercept generally underestimates the precursor H/C, with a correlation of H/C = $0.606 + 0.768 \times$ Intercept (R² = 0.97). The H/C_{os} are then calculated based on this equation, and are shown in Table S1. The uncertainty of this approach is estimated to be within ±0.1, which corresponds to minor uncertainty (less than 1%) in the calculated mass of OS precursors.

Based on the VDs of OS precursors as a function of carbon number (Fig. 1a), the reacted OS precursor mass concentration is estimated by two approaches. First, using the rate constants of the *n*-alkanes ($C_6 - C_{15}$) to calculate the reacted precursor in each bin, and adding them together to obtain the total reacted mass concentration. Second, using the rate constant of the *n*-alkane similar to the average carbon number (e.g., C_{10} for OS ore, see Sect. S4) as the rate constant of the OS precursor and calculating the reacted mass concentration. Figure S7 shows the calculated yields using these two approaches for OS ore. These two approaches give very similar yields, with both of them higher than the SOA potential (assuming all precursor reacted) at photochemical age of < 3 days. After 3 days, all of the curves are identical since the precursors are 100% reacted. In addition, we also calculate the yields assuming the OS ore has a rate constant of cyclodecane, with calculated yields slightly lower than assuming it to be *n*-decane. For other OS-related precursors, the results are similar to OS ore. As these approaches give very similar results and the average carbon method is simpler to conduct, the SOA yields of OS precursors shown in this study are all calculated based on the average carbon method using the rate constant of corresponding *n*-alkanes.

The carbon and oxygen yields (Y_c and Y_o) are also shown in this study, which are calculated by the method described previously (Kroll et al., 2009;Lambe et al., 2012):

$$Y_C = Y \times \frac{12}{12 + 16 \times 0/C + H/C}$$
$$Y_O = \frac{Y_C \times 0/C \times 16}{12}$$

S6. LVOCs fate correction

The fate of the LVOCs in the OFR include condensation on the reactor wall, exiting the reactor, and reacting with OH to form higher volatility compounds that are not condensable. These three losses may influence the LVOC fate, which in the atmosphere is expected to be condensation to pre-existing aerosol. Here, the method developed by Palm et al. (2016) was used to correct for these losses. Briefly, the lifetimes of LVOCs associated with these processes were used to estimate the fractional loss of each process. These lifetimes are $\tau_{aerosol}$ (condense on aerosol), τ_{wall} (condense on reactor wall), and τ_{OH} (react with OH to form non-condensable compounds) and their parametrization is described below:

1. $\tau_{aerosol}$ is estimated by (Pirjola et al., 1999):

$$\tau_{aerosol} = 1/(4\pi \cdot CS \cdot D)$$

where D is the diffusion coefficient, which is estimated to be 7×10^{-6} m² s⁻¹ for oxidized organic vapor with a molecular weight of 200 g mol⁻¹ (Tang et al., 2015); CS is the "condensational sink", which represents the sink associated with aerosols and is related to particle size distribution:

$$CS = \sum_{i} r_i \,\beta_i \,N_i$$

where r_i and N_i are the particle radius and number concentration of each size bin of SMPS. The β term is the correction factor for gaseous diffusion to the particle surface:

$$\beta = \frac{Kn+1}{0.377Kn+1 + \frac{4}{3}\alpha^{-1}Kn^2 + \frac{4}{3}\alpha^{-1}Kn}$$

where α is the mass accommodation coefficient (also known as the sticking coefficient) of condensing vapor, which is assumed to be 1; *Kn* is the Knudsen number:

$$Kn = 3 \sqrt{\frac{\pi M}{8RT}} \frac{D}{r}$$

where *M* is the molecular weight of the condensing vapor, which is assumed to be 200 g mol⁻¹; *R* is the gas constant; *T* is the temperature.

2. τ_{wall} is estimated by (McMurry and Grosjean, 1985):

$$\tau_{wall} = \frac{\pi}{2 \; \frac{A}{V} \; \sqrt{k_e D}}$$

where A/V is the surface-area-to-volume ratio, which is 22.3 m⁻¹ for our OFR; k_e is the eddy diffusion coefficient (Krechmer et al., 2016):

$$k_{\rho} = 0.004 + 0.0056 \cdot V^{0.74}$$

3. It is assumed that after reacting five times with OH, the LVOCs are fragmented to small molecules that are too volatile to condense (Palm et al., 2016). Hence, τ_{OH} is estimated by:

$$\tau_{OH} = \frac{5}{k_{OH} \cdot [OH]}$$

where k_{OH} is the rate constant for reaction with OH, which is assumed to be 1×10^{-11} cm³ molec⁻¹ s⁻¹ (Ziemann and Atkinson, 2012).

Using the three lifetimes above, the total lifetime of these three pathways (τ_{total}) can be estimated by:

$$\frac{1}{\tau_{total}} = \frac{1}{\tau_{aerosol}} + \frac{1}{\tau_{wall}} + \frac{1}{\tau_{OH}}$$

Combined with the residence time (τ_{OFR}), the fraction that exit the OFR (F_{exit}), condense on aerosol ($F_{aerosol}$), condense on reactor wall (F_{wall}), and react with OH (F_{OH}) can be estimated using the following equations:

$$F_{exit} = e^{-\frac{\tau_{OFR}}{\tau_{total}}}$$

$$F_{aerosol} = (1 - F_{exit}) \cdot \frac{\tau_{total}}{\tau_{aerosol}}$$

$$F_{wall} = (1 - F_{exit}) \cdot \frac{\tau_{total}}{\tau_{wall}}$$

$$F_{OH} = (1 - F_{exit}) \cdot \frac{\tau_{total}}{\tau_{OH}}$$

According to the sensitivity analysis performed previously (Palm et al., 2016), variations in most of parameters above have little influence on the results with the exception of CS and α . The CS that used in this method is the average value at the beginning and the end of the reactor. Using a CS at the beginning of the reactor will largely enhance the correction factor. However, as organic aerosols are formed through the entire reactor, using the average CS is more reasonable. Varying α from 1 to 0.1 also largely enhances the correction factor. However, according to a recent study (Krechmer et al., 2017), the accommodation coefficient was quantified to be ~1. Hence the influence from α is likely to be small. Overall, we assume that the uncertainty associated with this correction approach is within ±30%.

When applying the above correction to SOA yields, one needs to know the LVOCs fraction in SOA. The LVOCs and ELVOCs (extremely low volatility VOCs) in the atmosphere are ~100% in the particle phase, while the S/IVOCs may be partially in the gas phase, depending on the organic aerosol concentration. As a result, a OFR fate correction for S/IVOCs is not feasible. Previous field measurements and laboratory studies demonstrated that SOA from various sources were mainly ~40%-80% L/ELVOCs and ~20%-60% S/IVOCs (Hong et al., 2017;Saha et al., 2017;D'Ambro et al., 2018;Sato et al., 2018;Saha et al., 2018). In our experiments, the low-NO_x yields are significantly higher than the high-NO_x yields (paper in preparation), indicating a lower volatility for SOA formed under low-NO_x conditions. Hence, the upper limit of the fraction (80%) is used for the LVOCs correction of low-NO_x yields. The 20% S/IVOCs in SOA remain unchanged. Previous studies have assumed SOA to be 100% LVOCs (Ortega et al., 2016;Palm et al., 2016), and the 80% LVOCs used here leads to slightly lower correction factors.

S7. Conversion efficiency determination of the THC system

The conversion efficiency of the catalyst in the THC system is determined by injecting a small constant volume of liquid hydrocarbon into a constant flow of zero air using a syringe pump (Harvard apparatus 11A Plus). In our measurement, the liquid flow rate was 0.005-0.1 μ L min⁻¹ depending on the volatility, and the flow rate of zero air was 5-10 L min⁻¹. The airflow was at room temperature for high volatility compounds (e.g., toluene), while it is maintained at 60-100 °C for low volatility compounds (e.g., octadecane) to ensure that the hydrocarbon was 100% evaporated. The mixing ratio of the hydrocarbon was then calculated by:

$$HC(ppb) = \frac{RT\rho F_l \times 10^9}{MPF_g}$$

where *R* is the gas constant; *T* is the temperature; ρ is the density of the liquid; F_i is the flow rate of the liquid; *M* is the molecular weight; *P* is the atmospheric pressure; F_g is the zero air flow rate.

Figures and Tables



Figure S1. Schematic diagram of the ECCC-OFR and associated experimental setup.



Figure S2. Catalyst efficiencies of the THC system for toluene and C₁₀, C₁₃, C₁₄, C₁₆, and C₁₈ *n*-alkanes.



Figure S3. GC-MS chromatogram of the OS-related precursors. Dashed lines indicate the retention time of the *n*-alkanes from C_7 to C_{13} .



Figure S4. CFD simulation results: (a) velocity distribution; (b) vectors showing flow field. The red lines in (b) indicate the areas with recirculation.



Figure S5. Residence time distribution (RTD) of CO₂ in ECCC-OFR compared to ideal laminar flow.



Figure S6. H_2SO_4 yields as a function of particle diameter in the ECCC-OFR and previous OFRs (Lambe et al., 2011).



Figure S7. OS ore SOA potential and yield based on the reaction rate constant from the VD, *n*-decane and cyclodecane. The inset shows the first three photochemical ages.



Figure S8. Comparison of LVOCs fate corrected SOA yields for α -pinene with previous smog chamber studies (Ng et al., 2007;Eddingsaas et al., 2012;Bruns et al., 2015;Han et al., 2016).



Figure S9. Representative AMS mass spectra for each OS-related oxidation experiment at OH exposure of ~ 1.5×10^{12} molec cm⁻³ s.



Figure S10. Linear regression of the H/C versus O/C ratios of SOA formed from OS-related precursors.

Precursor	Intercept	Slope	R ²	Precursor H/C
<i>n</i> -Heptane (C7)	2.17	-0.63	0.983	2.29
<i>n</i> -Decane (C10)	2.12	-0.72	0.996	2.2
<i>n</i> -Dodecane (C12)	2.04	-0.70	0.996	2.17
Cyclodecane	1.76	-0.43	0.995	2
Decalin	1.59	-0.32	0.999	1.8
Naphtha	2.10	-0.66	0.989	2.22 ^a
Dilbit	2.07	-0.65	0.998	2.2 ^a
Tailings pond	1.93	-0.47	0.999	2.09 ^a
Bitumen	1.80	-0.48	0.997	1.99 ^a
OS ore	1.77	-0.43	0.996	1.96 ^a

Table S1. Intercept and slope of the linear regression of the H/C versus O/C ratios in Fig. 7, and the H/C ratio of the precursors.

a. Calculated based on the linear fitting result of precursor H/C and intercept for alkanes (see Sect. S5).

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