Impact of HO₂/RO₂ ratio on highly oxygenated α-pinene photooxidation products and secondary organic aerosol formation potential

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Abstract. Highly oxygenated molecules (HOMs) from the atmospheric oxidation of biogenic volatile organic compounds are important contributors to secondary organic aerosol (SOA). Organic peroxy radicals (RO₂) and hydroperoxy radicals (HO₂) are key species influencing the HOM product distribution. In laboratory studies, experimental requirements often result in overemphasis on RO₂ cross-reactions compared to reactions of RO₂ with HO₂. We analyzed the photochemical formation of HOMs from α-pinene and their potential to contribute to SOA formation under high (≈ 1/1) and low (≈ 1/100) HO₂/RO₂ conditions. As HO₂/RO₂ > 1 is prevalent in the daytime atmosphere, sufficiently high HO₂/RO₂ is crucial to mimic atmospheric conditions and to prevent biases by low HO₂/RO₂ on the HOM product distribution and thus SOA yield. Experiments were performed under steady-state conditions in the new, continuously stirred tank reactor SAPHIR-STAR at Forschungszentrum Jülich. The HO₂/RO₂ ratio was increased by adding CO while keeping the OH concentration constant. We determined the HOM’s SOA formation potential, considering its fraction remaining in the gas phase after seeding with (NH₄)₂SO₄ aerosol. An increase in HO₂/RO₂ led to a reduction in SOA formation potential, with the main driver being a ∼ 60 % reduction in HOM-accretion products. We also observed a shift in HOM-monomer functionalization from carbonyl to hydroperoxide groups. We determined a reduction of the HOM’s SOA formation potential by ∼ 30 % at HO₂/RO₂ ≈ 1/1 compared to HO₂/RO₂ ≈ 1/100. Particle-phase observations measured a similar decrease in SOA mass and yield. Our study shows that too low HO₂/RO₂ ratios compared to the atmosphere can lead to an overestimation of SOA yields.
1 Introduction

In the atmosphere, highly oxidized products from the oxidation of biogenic or anthropogenic volatile organic compounds (VOCs) are an important source of secondary organic aerosol (SOA) (Roldin et al., 2019; Mohr et al., 2019). SOA is an important contributor to the overall ambient aerosol and of interest because of its impact on climate, visibility and human health (Hallquist et al., 2009).

Recently, many studies (Pullinen et al., 2020; Berndt et al., 2016; Bianchi et al., 2017) have focused on understanding the oxidation pathways of VOCs that yield highly oxygenated molecules (HOMs), as these are expected to be of low enough volatility to condense into the particle phase. One important tool for the investigation of VOC degradation and SOA formation is the utilization of experiments in atmospheric simulation chambers (Hidy, 2019). Such experiments have also helped to elucidate key processes in the HOM formation, i.e., the process of autoxidation.

After an initial oxidant attack and the formation of a peroxy radical (RO$_2$), autoxidation adds oxygen to the molecule via an internal H shift to the peroxy group, forming a hydroxy peroxy group and an alkyl radical, to which O$_2$ immediately adds, re-establishing the peroxy functionality. This process can be repeated multiple times, yielding almost instantaneously highly oxygenated peroxy radicals (HOM-RO$_2$) which are terminated to a series of HOM closed-shell products (Bianchi et al., 2019; Ehn et al., 2014; Crounse et al., 2013).

Chamber studies often work with a singular compound and operate at higher precursor concentrations than those observed in the atmosphere for experimental reasons. These experiments cannot represent the complex mixture of VOCs and oxidized VOCs present in the atmosphere (McFiggans et al., 2019). Higher precursor concentrations can lead, per se, to higher SOA yields than observed in the atmosphere (a well-characterized phenomenon; see Henry et al., 2012, and Shilling et al., 2009) and to a general preference for higher-order processes which may not be important in the atmosphere. One example is that chamber studies tend to overestimate the role of cross-reactions between organic peroxy radicals (RO$_2$), owing to high precursor concentrations of a single VOC. In chambers, reactions of HOM-RO$_2$ with other organic peroxy radicals terminate the autoxidation chain, leading typically to multifunctional carbonyl and alcohols compounds. In comparison, in the atmosphere termination by HO$_2$ is more likely, leading to multifunctional hydroperoxides. In the presence of sufficient NO, termination to multifunctional organic nitrates may be more important (Schervish and Donahue, 2021).

Another possible termination reaction of HOM-RO$_2$ with HOM-RO$_2$ and less oxidized RO$_2$ leads to the formation of accretion products, which are expected to be extremely low volatility organic compounds (ELVOCs) and are therefore expected to contribute to new particle formation and SOA formation (Ehn et al., 2014; Berndt et al., 2018). Schervish and Donahue (2021) raised awareness that chamber studies could overestimate the SOA formation potential from the oxidation of terpenes such as α-pinene compared to the atmosphere because of missing HO$_2$ and low-molecular-weight RO$_2$ (e.g., CH$_3$O$_2$), which favors accretion product formation. Previous studies of VOC ozonolysis with different OH scavengers by Docherty and Ziemann (2003) and Keywood et al. (2004) indicated a significant impact of the HO$_2$/RO$_2$ ratio on SOA yields.

In chamber studies, the use of higher VOC concentrations is often an unavoidable necessity either to match the sensitivity of the analytical instrumentation or to overcome chamber-related effects. The question remains, how can conditions dictated by the chamber be steered towards more realistic chemical pathways and higher atmospheric relevance?

In this study, we address this overestimated importance of peroxy radical cross-reactions. We studied the photooxidation of α-pinene in a series of steady-state experiments in the newly built, continuously stirred tank reactor SAPHIR-STAR (a modernized version of JPAC; see Mentel et al., 2009).

We compared two experimental conditions, a pure α-pinene photooxidation case, leading to low HO$_2$/RO$_2$ ratios and high importance of RO$_2$ cross-reactions, and a high HO$_2$/RO$_2$ case, representing more atmospherically relevant conditions with high importance of RO$_2$+HO$_2$ reactions. One important concept of the conducted experiments is the constant OH availability to α-pinene in order to prevent effects of different oxidant levels and allow for a direct comparison between the two chemical regimes. To this end, the OH concentration in the experiments was adjusted to keep the α-pinene OH turnover constant and to avoid changes due to oxidant scavenging.

Furthermore, the addition of seed particles ((NH$_4$)$_2$SO$_4$) allowed us to observe the condensation behavior of the HOM products and to compare our gas-phase observations directly with particulate-phase measurements of the condensed organic mass.

In this study, we will address two central questions. How does the shift in HO$_2$/RO$_2$ impact the oxidation mechanism of α-pinene, especially the HOM formation pathway? And what is the subsequent impact on the SOA formation potential of the α-pinene photooxidation system? As the central analysis tool, we will use high-resolution time-of-flight mass spectrometry with chemical ionization (HR-TOF-CIMS) with nitrate (NO$_3^-$) reagent ions as this ionization scheme is selective towards HOM compounds (Hyttinen et al., 2018).
2 Methods

2.1 Generic α-pinene HOM peroxy radical chemistry

The chemical mechanistic information for the basic oxidation scheme of α-pinene was taken from the Master Chemical Mechanism (MCM) v3.3.1 (Jenkin et al., 1997; Saunders et al., 2003) (http://mcm.york.ac.uk, last access: 20 October 2023). The main peroxy radicals expected from α-pinene photooxidation are C10H17O4 and C10H15O5. C10H17O4 is formed by the addition of OH to α-pinene, followed by O2 (starting RO2: C10H17O3) (MCM v3.3.1 (Jenkin et al., 1997; Saunders et al., 2003)). Studies showed that the autoxidation can start from C10H17O3 with the four-member ring in α-pinene opened (Berndt, 2021; Xu et al., 2019).

For C10H15O5, the autoxidation chain is assumed to start with C9H15O4, which can be formed directly from ozonolysis via the vinyl hydroperoxide path (Johnson and Marston, 2008; Iyer et al., 2021) or via H abstraction from first-generation oxidation products such as pinonaldehyde (C10H16O2) (MCM v3.3.1; Jenkin et al., 1997; Saunders et al., 2003; Fantechi et al., 2002). A recent study suggests direct H abstraction by OH from α-pinene (Shen et al., 2022) as a starting point for the autoxidation chain.

The autoxidation process is rapid with H-shift rates of about 0.01–0.1 s−1 and faster (Piletic and Kleindienst, 2022; Berndt, 2021; Xu et al., 2019; Vereecken et al., 2007). The autoxidation chain will run quickly, adding more oxygen to the molecule, until bimolecular termination reactions are able to compete with all available H-shift rates. The rate of an H shift is determined by the hydrogen’s position in relation to the peroxy radical and the functional groups near the hydrogen and peroxy radical (Otkjaer et al., 2018; Vereecken and Nozière, 2020). In the absence of NOx, the peroxy radicals have two major bimolecular termination channels: the reaction with another RO2 or with HO2. A third pathway is the intramolecular termination (Rissanan et al., 2014).

Based on the considerations above, we apply a simplified generic reaction scheme to analyze our observations. Figure 1 shows an overview of the reaction pathways for the main peroxy radical families in the α-pinene photooxidation and the resulting product groups and families. The compounds can be separated into four classes: peroxy radicals (HOM-RO2), monomers (HOM-Mon), accretion products (HOM-Acc) and fragments (HOM-Frag). The HOM-RO2 class consists of all detected HOM-RO2, with special focus on the analysis of the C10 HOM-RO2 families. The HOM-Mon class contains the closed-shell HOM-C10 products. The compounds in the fragment class contain less than 10 carbon atoms, while all HOM-Acc compounds contain more than 10 carbon atoms. The compound classes are further divided into groups and families. Here, the term group is used for compounds with the same carbon number, while a family contains all compounds with the same carbon and hydrogen number but a varying oxygen number.

The termination of RO2 with HO2 will lead to hydroperoxide formation:

\[ RO_2 + HO_2 \rightarrow ROOH + O_2. \] (R1)

In the case of C10H15O5, Reaction (R1) will lead to multifunctional C10H16O2 hydroperoxides (wherein the notation “hydroperoxides”, “carbonyls”, or “alcohols”, etc., here and in the following relates to the functionality of the group formed by the termination reaction). For C10H17O4, it will lead to the formation of C10H18O2 hydroperoxides. The termination via RO2+RO2 can either result in the formation of accretion products or in the formation of carbonyls and alcohols. For the accretion product formation, it is assumed that the two RO2 chemically bond, eliminating O2 from the molecule:

\[ RO_2 + R'O_2 \rightarrow R-O-O-R' + O_2. \] (R2)

Recombination reactions of the main peroxy radical families C10H15O4 and C10H17O5 lead to the product families C20H30O6 (combination of two C10H15O4), C20H32O7 (combination of C10H15O5 and C10H17O5), and C20H34O8 (combination of two C10H17O4).

However, due to reactions with smaller peroxy radicals, HOM-Acc families with smaller carbon and hydrogen numbers are also observed. Indeed, one reason why the RO2+RO2 termination is expected to affect the SOA formation potential is the formation of accretion products by scavenging of less oxidized and low-molecular-weight RO2 by HOM-RO2. Thus, the smaller RO2 will also contribute to the SOA mass which would otherwise not be the case. For the HOM-RO2 itself, it is expected that they contribute to SOA formation independently of the termination pathway, due to the low volatility of its expected termination products (Pullinen et al., 2020; McFiggans et al., 2019).

The second RO2+R’O2 termination pathway is the formation of a carbonyl and alcohol compound:

\[ RO_2 + R'O_2 \rightarrow R-OH + R'=O + O_2. \] (R3)

In this reaction, both radicals lose an oxygen atom, and a hydrogen atom is transferred to the RO2, forming the alcohol termination group. Preferences of RO2 to form an alcohol or carbonyl compound are possible for individual reactions, but statistically carbonyl and alcohols should be formed with the same fractions. Since mass spectrometry can only determine formula composition, we cannot distinguish alcohols and hydroperoxides, which arise from RO2 differing by one O atom. Therefore, details of balance of alcohol and carbonyl formation cannot be detected.

However, the formula composition can help to differentiate certain formation pathways. The C10H14O2 family contains only carbonyl formed from a C10H15O4 RO2, while the alcohol will be part of the C10H16O4 family. The C10H16O2 family also contains the carbonyl produced.
from the RO₂+R'O₂ monomer termination of C₁₀H₁₇O₃, while the alcohol from this RO₂ family will be found in the C₁₀H₁₅O₂ family. So, from a diagnostic point of view, C₁₀H₁₆O₂ as well as C₁₀H₁₈O₂ are uniquely related to a precursor radical family. The classification of the formation pathways of the monomers is helpful to analyze the effect of the HO₂/RO₂ ratio shift in the experiments. Considering the termination pathways, a decrease in the C₁₀H₁₄O₂ family and an increase in the C₁₀H₁₈O₂ family are expected with increasing HO₂/RO₂ because of increasing termination by HO₂ and decreasing termination by RO₂. In the case of C₁₀H₁₈O₂, the increase in hydroperoxides is partially compensated by a decrease in the alcohol channel. For C₁₀H₁₆O₂, the situation is more complicated as it contains contributions from all termination pathways.

Besides closed-shell products, HOM-RO₂ can also form alkoxy radicals (HOM-RO). In general, alkoxy radicals (RO) are important intermediates in the oxidation scheme of organics and are formed via (Reaction R4) and probably also via (Reaction R5) for specific RO₂ (Jenkin et al., 2019):

\[
\text{RO}_2 + \text{R'}O_2 \rightarrow \text{RO} + \text{R'O} + \text{O}_2, \quad \text{(R4)}
\]

\[
\text{RO}_2 + \text{HO}_2 \rightarrow \text{RO} + \text{OH} + \text{O}_2. \quad \text{(R5)}
\]

In Reaction (R5), OH will be formed. The importance of Reaction (R5) compared to Reaction (R1) is still unclear in the literature, but functionalization of the RO₂ close to the peroxy functionality possibly enables this reaction (Iyer et al., 2018; Eddingsaas et al., 2012; Hasson et al., 2005; Jenkin et al., 2019). If Reaction (R5) is of negligible importance, the reaction scheme will simplify, and the effect of increased HO₂/RO₂ is easier to diagnose.

We are interested in the importance of alkoxy radical formation as (HOM)-RO tends to fragment, leading to the formation of smaller products (Vereecken et al., 2007). In the context of SOA formation, these fragments are less likely to contribute to SOA mass because of their higher volatility. Since alkoxy radicals are too unstable to be detected directly, we use two diagnosis tools to judge the importance of HOM-RO. Firstly, HOM-RO fragmentation can lead to HOM-RO₂ with less than 10 carbon atoms, which may also continue the autoxidation chain. Therefore, the abundance of HOM with less than 10 carbon atoms (HOM-Frag) indicates the importance of alkoxy steps. Secondly, with increasing functionalization, H shifts retaining the carbon backbone become more likely (Vereecken et al., 2007), which will lead to the generation of new C₁₀-HOM-RO₂. Such alkoxy peroxy steps can continue the autoxidation chain (Mentel et al., 2015). Interestingly, by coupling of an alkoxy and a peroxy step, the parity of the number of oxygen atoms in the HOM-RO₂ changes, while in pure autoxidation steps the oxygen parity remains the same. Therefore, a parity change of the oxygen number can be used as an indication of alkoxy step abundance (Kang, 2021).

In summary, we will use the changes in contribution and relative signal of the different families and classes to judge the impact of shifting from low to high HO₂/RO₂ on the α-pinene photooxidation pathway.

2.2 Control of α-pinene OH turnover

After the initial α-pinene photooxidation phase as a reference, CO was added to the oxidation system. The idea is to represent small, oxidized VOCs in the atmosphere that can produce HO₂ by reaction with OH (compare Schervish and Donahue, 2021). The presence of CO shifts the HO₂/RO₂ ratio, increasing the importance of the termination of RO₂ by HO₂. McFiggans et al. (2019) showed that one limiting factor in mixture experiments is oxidant scavenging: the products and their yields in mixed systems change, because there is less OH available to the individual VOCs. Thus, after the CO addition the OH production in the chamber was increased to compensate for the OH consumed by CO. The OH levels in the system before and after the CO addition were approxi-
mately the same, keeping the \(\alpha\)-pinene OH turnover approximately constant. This OH adjustment ensures that the primary \(\alpha\)-pinene chemistry was kept the same, avoiding effects by different oxidant levels and enabling a direct comparison.

However, since experiments could only be performed at about the same OH levels, a normalization by the actual \(\alpha\)-pinene OH turnover is applied to the data. This compensates for the slight experimental imperfections and enables better comparison of experiment series with different boundary conditions. The turnover in steady state is given in Eq. (1). Here the subscript “SS” denotes steady-state condition for the concentrations of \(\alpha\)-pinene and OH, and \(k_{\text{OH}}\) is the \(\alpha\)-pinene OH reaction rate constant.

\[
\text{turnover}_{\alpha\text{-pinene}+\text{OH}} = k_{\text{OH}} \cdot [\alpha\text{-pinene}]_{\text{SS}} \cdot [\text{OH}]_{\text{SS}}
\] (1)

This normalization also directly shows the yield of certain oxidation products or product groups per \(\alpha\)-pinene consumed by OH.

### 2.3 Derivation of effect on condensable mass from gas-phase measurement

A simple proxy for the condensable mass from HOM products can be calculated from the steady-state HOM signals measured by the NO\(_3\)-CIMS, assuming condensation for all low-volatility HOM compounds and no back-evaporation into the gas phase. To only take low-volatility products into account, we used all detected formula compositions with \(M > 230\ \text{g mol}^{-1}\) and weighted them with their molar mass. The reasoning behind this threshold can be found in Sect. 4.4. All contributions were summed up and normalized with the \(\alpha\)-pinene OH turnover for the comparison between the low and high \(\text{HO}_2/\text{RO}_2\) cases (Eq. 2).

\[
\text{mass-weighted signal sum} = \frac{\sum_{i=0}^{i} S_i \cdot M_i}{\text{turnover}_{\alpha\text{-pinene}+\text{OH}}}
\] (2)

We also estimated the expected SOA mass formed using the calibration factor obtained for sulfuric acid for our NO\(_3\)-CIMS instrument in a calibration setup (see Supplement Sect. S1). From this, we calculated an upper boundary concentration of detected HOM compounds in the gas phase under the assumption that sulfuric acid clusters with nitrate at the collision limit, yielding maximum sensitivity (a common approach; see, for example, Ehn et al., 2014, and Pullinen et al., 2020).

The calculated gas-phase concentration was then used in the steady-state equation describing the relationship between gas- and particle-phase concentrations of a single compound \(i\) shown in Eq. (3).

\[
m_{\text{seed}}(p) = \frac{m_{\text{seed}}(g) \cdot k_{\text{cond},i}}{k_{\text{particleLoss}} + k_{\text{evap},i}}
\] (3)

Equation (3) shows that the steady-state particle-phase (mass) concentration \(m_{\text{seed}}(p)\) of compound \(i\) in the presence of seed particles in the chamber is only dependent on the steady-state gas-phase concentration \(m_{\text{seed}}(g)\), the condensation rate and evaporation rate constants \(k_{\text{cond},i}\) and \(k_{\text{evap},i}\) of \(i\) (to and from the particles) and the particle-loss rate constant \(k_{\text{particleLoss}}\) in the chamber. The condensation rate can be calculated (see Supplement Sect. S8), and the particle-loss rate constant was measured by observation of the particle loss in the chamber after ending the seed addition (details in the Supplement Sect. S2). The evaporation rate was assumed to be negligible for the investigated HOM compounds.

For the SOA yield calculation, we calculated a corrected organic mass \(m_{\text{SOA}}\) from the organic mass \(m_{\text{AMS}}\) measured by aerosol mass spectrometry (AMS) and the fraction expected to be lost on the seed particles compared to the overall loss on particles and chamber wall as shown in Eq. (4) (McFiggans et al., 2019).

\[
m_{\text{SOA}} = m_{\text{AMS}} \cdot \frac{k_{\text{cond}} + k_{\text{wall}}}{k_{\text{cond}}}
\] (4)

In Eq. (4), we use the condensation rate constant \(k_{\text{cond}}\) calculated for one major HOM product (\(\text{C}_{16}\text{H}_{26}\text{O}_{7}\)) and the average HOM-Mon wall loss rate \(k_{\text{wall}}\), which was determined by switching off the UV-C light and observing the decay of photooxidation products in the NO\(_3\)-CIMS. The wall loss determination, as well as SOA mass correction, was described before in Sarrafzadeh et al. (2016) and McFiggans et al. (2019).

### 3 Experimental methods

#### 3.1 Chamber setup

Experiments were conducted in the Jülich SAPHIR-STAR chamber, which is the modern successor of the JPAC setup (Mentel et al., 2009). The basic concepts are the same as in JPAC, but each parameter is set, controlled and monitored in a program. The chamber was operated as a continuously stirred tank reactor. It is a borosilicate glass cylinder (\(l = 2.5\ \text{m}, d = 1\ \text{m}\)) with a volume of close to 2000 L, and all equipment inside the chamber is either glass or glass-coated steel (SilcoTek GmbH).

With an inflow of 32 L min\(^{-1}\), the residence time in the chamber was approximately 61 min with a fan ensuring mixing within minutes. In contrast to the JPAC chamber, the stirring is conducted perpendicular to the cylinder axis, as opposed to coaxial. Chamber inflow is split into two humidified clean air flows (mixed from \(\text{N}_2\) and \(\text{O}_2\)) of about equal volume: one with added oxidant (here \(\text{O}_3\)) and the other with added VOC and other trace gases (here \(\alpha\)-pinene and CO). All experiments were performed at a relative humidity of 50% and temperature of 20 °C. Temperature stability is ensured by the climate-controlled surroundings of the chamber.

\(\alpha\)-Pinene (\(\geq 99\%\) purity, Sigma-Aldrich, Merck KGaA) was introduced via liquid injection with a syringe pump...
(Fusion 4000, CHEMYX Inc.) into a heated glass bulb and flushed by a 1 L min\(^{-1}\) \(\text{N}_2\) stream into the chamber. CO was added from a gas bottle (10 % CO in \(\text{N}_2\), Messer SE & Co. KGaA). Ozone was directly produced photolytically before injection with a self-built ozone generator.

OH is produced in the chamber by ozone photolysis using two UV-C lamps with a wavelength of 254 nm and subsequent reaction of \(\text{O}(^1\text{D})\) with water vapor. The lamps are mounted in closed quartz cylinders in the middle of the chamber, vertically to the cylinder axis, and light intensity can be varied with a movable shielding installed around the lamps. The shielding allows for an exact percentage of the lamp to be covered, thus controlling the amount of OH produced in the chamber.

The OH radical concentration after CO addition was adjusted by setting the shielding of the UV-C lamps and a slight adjustment of \(\text{O}_3\) inflow. The applied \(J(\text{O}(^1\text{D}))\) values in different phases were calculated to be in the range of \(8 \times 10^{-3}\) to \(2.4 \times 10^{-3}\) s\(^{-1}\).

In some of the experiments, ammonium sulfate (\(\geq 99\%\) purity, Merck KGaA) seed particles were added to the system to provide a surface for the condensation of organic material. The aerosol was produced with a modified TSI atomizer (model 3076, TSI GmbH) and dried to 50 % relative humidity.

VOC concentrations in the chamber were measured using proton-transfer-reaction mass spectrometry (PTR-TOF-MS; Ionicom GmbH). \(\text{CO}_2\), \(\text{CO}\) and \(\text{H}_2\text{O}\) (G2401 cavity ringdown spectrometer, Picarro Inc.); \(\text{NO}\) and \(\text{NO}_x\) (NCLD899, Eco Physics GmbH with a home-built photolytic converter); and \(\text{O}_3\) (O342e, Envea GmbH) were additionally monitored. Particle distribution and concentration were measured with a condensation particle counter (CPC, model 3788, TSI GmbH) and a scanning mobility particle sizer (SMPS; model 3080, TSI GmbH) with a CPC (model 3788, TSI GmbH). The aerosol composition was measured with a high-resolution aerosol mass spectrometer (HR-TOF AMS; Aerodyne Inc.).

In all experiments, VOC, \(\text{O}_3\) and SMPS + CPC sampling switched between inlet and outlet of the chamber to measure the input concentrations as well as the concentrations in the reactor. The flow control system of the chamber adapts to these switches so that the inflow into the chamber stays constant.

All results discussed here were observed under steady-state conditions when all parameters were constant. For each steady state, the OH concentration was calculated from the decay of \(\alpha\)-pinene as described by Kiendler-Scharr et al. (2009). Equation (5) is derived from the mass balance of \(\alpha\)-pinene at steady state. The steady-state OH concentration \([\text{OH}]_{\text{SS}}\) depends on the amount of \(\alpha\)-pinene consumed by reaction with OH and the reaction with \(\text{O}_3\), as well as the flush out.

\[
[\text{OH}]_{\text{SS}} = \frac{F \cdot \frac{[\text{VOC}]_{\text{in}} - [\text{VOC}]_{\text{SS}}}{[\text{VOC}]_{\text{SS}}} - k_{\text{O}_3} \cdot [\text{O}_3]_{\text{SS}}}{k_{\text{OH}}}
\]

Here, \(F\) is the total flow and \(V\) the volume of the chamber. The subscript “SS” indicates steady-state concentrations, while \([\text{VOC}]_{\text{in}}\) represents the \(\alpha\)-pinene concentration entering the chamber. \(k_{\text{O}_3}\) and \(k_{\text{OH}}\) represent the reaction rate constants of \(\alpha\)-pinene with the corresponding oxidant. We applied rate coefficients of \(k_{\text{OH}} = 5.36 \times 10^{-11}\) cm\(^3\) s\(^{-1}\) (Atkinson and Arey, 2003) and \(k_{\text{O}_3} = 9.25 \times 10^{-17}\) cm\(^3\) s\(^{-1}\) (Cox et al., 2020) at 20 °C. The uncertainty of the OH calculation was estimated as 20 % by Wildt et al. (2014).

### 3.2 Experiment conditions

An overview of the experiments and their boundary conditions can be found in Table 1. Four experiments were performed in total, leading to one repetition of each studied condition. In two of the experiments, ammonium sulfate seeds were added, leading to a total particle surface in the chamber on the order of \(8 \times 10^{-4}\) m\(^2\) m\(^{-3}\) and organic loadings of about 3 \(\mu\)g m\(^{-3}\) in the photooxidation stage. In the unseeded experiments, no significant nucleation was observed, leading to pure gas-phase conditions. The Exp2 experiment is a consecutive combination of a seeded followed by a non-seeded experiment to provide direct insight into the effect of the seed presence on the system.

As the OH radical is produced by photolysis of ozone and as \(\alpha\)-pinene reacts with ozone, it is important to know the relative contribution of the \(\alpha\)-pinene consumption by OH and by \(\text{O}_3\). This is achieved by comparing the turnover of \(\alpha\)-pinene with OH and \(\text{O}_3\), respectively. The results can be found in Table 1. The listed results are for the low \(\text{HO}_2/\text{RO}_2\) conditions, but nearly identical values were reached after the \(\text{HO}_2/\text{RO}_2\) shift.

### 3.3 Experimental procedure

All experiments started with \(\alpha\)-pinene ozonolysis followed by illumination with the UV-C lights to induce the reaction with OH. A general flow scheme of the experiment can be found in Fig. 2, together with one exemplary time series of the unseeded experiment (Exp1). After the photooxidation steady state, \(\text{CO}\) was added to the system. In the displayed Exp1, the OH level was adjusted in three steps to approach the same concentration as before the CO addition. First the UV-C light opening was adjusted and then \(\text{O}_3\) was added, and the UV-C light opening was adjusted again. In some experiments initially the effect of CO on the unchanged system was observed before the adjustment of OH. In other experiments (Exp2-2, Exp3), the adjustment of the \(\alpha\)-pinene OH turnover via ozone concentration and UV-C light opening were made simultaneously with the CO addition. Highlighted in green...
Table 1. Overview of experimental conditions.

<table>
<thead>
<tr>
<th>Name</th>
<th>Experiment description</th>
<th>[VOC]_{in}</th>
<th>[CO]_{in}</th>
<th>[OH]_{ss} at low HO\textsubscript{2}/RO\textsubscript{2}</th>
<th>Contribution of OH to turnover at low HO\textsubscript{2}/RO\textsubscript{2}</th>
<th>Particle surface at low HO\textsubscript{2}/RO\textsubscript{2}</th>
<th>Organic mass concentration at low HO\textsubscript{2}/RO\textsubscript{2}</th>
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</thead>
<tbody>
<tr>
<td>Exp1</td>
<td>unseeded (1)</td>
<td>10 ppbv</td>
<td>2.5 ppmv</td>
<td>4.8 × 10\textsuperscript{6} cm\textsuperscript{-3}</td>
<td>80 %</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Exp2.1</td>
<td>seeded (1)</td>
<td>10 ppbv</td>
<td>2.5 ppmv</td>
<td>1.1 × 10\textsuperscript{7} cm\textsuperscript{-3}</td>
<td>92 % 8.7 × 10\textsuperscript{-4} m\textsuperscript{2} m\textsuperscript{-3}</td>
<td>3.4 µg m\textsuperscript{-3}</td>
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</tr>
<tr>
<td>Exp2.2</td>
<td>unseeded (2)</td>
<td>10 ppbv</td>
<td>2.5 ppmv</td>
<td>1.5 × 10\textsuperscript{7} cm\textsuperscript{-3}</td>
<td>94 %</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Exp3</td>
<td>seeded (2)</td>
<td>10 ppbv</td>
<td>2.5 ppmv</td>
<td>1.1 × 10\textsuperscript{7} cm\textsuperscript{-3}</td>
<td>92 % 6.8 × 10\textsuperscript{-4} m\textsuperscript{2} m\textsuperscript{-3}</td>
<td>2.7 µg m\textsuperscript{-3}</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2. (a) Experiment flow scheme. (b) Exemplary time series of Exp1, showing HOM-Mon and HOM-Acc product sum (top panel); calculated OH concentration and BrHO\textsubscript{2} signal (middle panel), and ozone, α-pinene, and CO concentrations (bottom panel). Background color represents light intensity. Highlighted in green are the low HO\textsubscript{2}/RO\textsubscript{2} steady state and the steady state at high HO\textsubscript{2}/RO\textsubscript{2} (addition of CO and adjusted oxidant level).

are the steady states with the “same” OH concentration characterized by low and high HO\textsubscript{2}/RO\textsubscript{2}, which were used for analysis and interpretation.

3.4 Model calculation for HO\textsubscript{2}/RO\textsubscript{2} ratio estimation

Box-model calculations were performed by applying the MCM v3.3.1 chemistry (Jenkin et al., 1997; Saunders et al., 2003) under the boundary conditions of the SAPHIR-STAR chamber. All calculations were performed with the institute software package EASY, which uses FACSIMILE to solve the differential equations (EASY Version 5.69b). More details about the model parameters can be found in the Supplement Sect. S3. The model calculations reproduced the primary observables α-pinene, O\textsubscript{3}, CO and OH within the experimental uncertainties. The box-model results were used to characterize the HO\textsubscript{2}/RO\textsubscript{2} ratio of the chemical systems, as no direct measurement of these parameters was available. The observed cluster signal BrHO\textsubscript{2} follows the modeled HO\textsubscript{2} concentration (Fig. 3).

The model predicts a shift of the HO\textsubscript{2}/RO\textsubscript{2} ratio from about 0.01 to about 1 by CO addition and oxidant adjustment, an increase by 2 orders of magnitude. Owing to a lack of observations to verify model results, we will consider only the magnitude of HO\textsubscript{2}/RO\textsubscript{2} here. The model results show that indeed a major shift from RO\textsubscript{2} + RO\textsubscript{2} to RO\textsubscript{2} + HO\textsubscript{2} reactions can be expected.

We further used the modeled RO\textsubscript{2} and HO\textsubscript{2} concentrations to estimate the relative importance of pathways for individual (observed) HOM-RO\textsubscript{2}. For that, we applied two generic rate coefficients \( k_{RO2HO2} \) and \( k_{RO2RO2} \). As the rate coefficient for the RO\textsubscript{2} + HO\textsubscript{2} termination to a hydroperoxide (\( k_{RO2HO2} \)), we used the value specified in the MCM (2.46 × 10\textsuperscript{-11} cm\textsuperscript{3} s\textsuperscript{-1} at 20 °C (Jenkin et al., 1997; Saunders et al., 2003)). We chose a \( k_{RO2RO2} \) of 5 × 10\textsuperscript{-12} cm\textsuperscript{3} s\textsuperscript{-1} as the approximated reaction rate of the RO\textsubscript{2} + RO\textsubscript{2} reactions. This value applies to all possible reactions (accretion product, monomer, and alkoxy formation) and is in the range of \( k_{RO2RO2} \) utilized by Roldin et al. (2019) in the PRAM model.

3.5 Determination of oxidized VOCs, HOMs and HO\textsubscript{2}

Chemical ionization mass spectrometry (HR-TOF-CIMS) techniques were used to detect a range of gaseous compounds. For this, two atmospheric-pressure-interface time-of-flight mass spectrometers (API-TOF-MS; Tofwerk AG) with different inlet systems were used simultaneously. General information about the API-TOF-MS instrument can be found in Junninen et al. (2010).

A long TOF (LTOF, Tofwerk AG) (resolution of ~8500 for peaks at > m/z 200) was coupled with the multi-scheme...
ionization inlet (MION; Karsa Oy). The setup of the inlet is described in detail by Rissanen et al. (2019). The distinctive feature of the MION inlet is the switching between two reagent ions. Here, nitrate was used to detect closed-shell HOMs, as well as HOM-RO2. Bianchi et al. (2019) suggested to define HOMs as products stemming from autoxidation containing more than six oxygen atoms. In our overall analysis, we decided to also include fragments and monomers containing five or in a few cases four oxygens (see the peak list in Supplement Sect. S5) as we are interested to see if the importance of these less oxidized (but still with NO3-CIMS detectable) products increases at higher HO2/RO2. However, in all considerations regarding SOA formation, we furthermore set a molar weight threshold which automatically excluded any products with less than six oxygens.

As the second reagent ion, bromide was used to detect less oxidized products and the HO2 radical (Albrecht et al., 2019; Sanchez et al., 2016). The nitrate ion source had a reaction time of 600 ms, while the bromide ion source had a shorter reaction time of 60 ms. For all experiments, an inlet flow of 10 L min⁻¹ was used, and the ionization scheme was switched every 10 min.

In the data evaluation, the first step was the separation of the time series of the two reagent ions. The data were subsequently processed with Tofware (Version 3.2.3, Tofwerk AG), using the high-resolution time series workflow. No transmission correction was performed as previous measurements showed an approximately flat relative transmission curve in the mass region of interest. The ion signals were normalized with the reagent ion signal (NO3⁻ and HNO3NO3⁻ for nitrate and Br⁻ and BrH2O⁻ for bromide).

Since no direct HO2 calibration was available, the HO2 signal in the Br-MION-CIMS was used to compare the levels of HO2 relative to each other in the different phases of the experiment. The comparison of the measured HO2 signal to the modeled HO2 concentration shows a good linear relation between the model predictions and observations.

Figure 3 illustrates this for the example of the Exp2 experiment. A background signal of around ~1 x 10⁻⁵ is observed as soon as VOC and ozone are present in the reactor. The background HO2 signal was not observed when only O3 or only VOC were in the system. As shown by the MCM modeling results, HO2 production of this strength is not expected in the α-pinene ozonolysis phase, but this background phenomenon was observed before (Albrecht et al., 2019) and is not fully understood.

For the HOM molecules measured by the NO3-MION-CIMS, the relative changes between different experiment phases are compared. For all detected HOM products, the same detection sensitivity is assumed. Hyttinen et al. (2018) showed in quantum chemical calculations that HOMs containing six or more oxygen atoms have comparable sensitivity with the nitrate reagent ion. At this degree of oxidation, it can be expected that the HOMs already contain multiple hydroperoxyl and/or hydroxy functional groups (Bianchi et al., 2019) prior to the termination step, making it unlikely that the sensitivity is strongly influenced by the termination group. Thus, the signal strength reflects the correct ranking of the observations, and relative comparisons do not require calibration. Pullinen et al. (2020) studied the mass balance between condensable HOMs and formed particle mass and were able to find closure within a factor of 2.

A second CI-APi-TOF was used to measure less oxidized species. It was configured with a CI inlet based on the design of Eisele and Tanner (1993) coupled to a TOF-MS (HTOF Tofwerk AG, resolution ≈2700 for peaks at >m/z 200) and was operated in positive mode with propylamine (C3H7NH2, Sigma-Aldrich, purity ≥ 99%) to detect the early-generation RO2 and oxidation products (Berndt et al., 2018). The propylamine was purified and added as an amine–N2 mixture (flow: 0.12 mL min⁻¹) to the 30 L min⁻¹ sheath flow. Furthermore, the sheath flow air is humidified to optimize ionization. The instrument sampled 0.1 L min⁻¹ from the chamber, which was diluted with 9.9 L min⁻¹ for a sample flow of 10 L min⁻¹. The dilution was necessary to reduce depletion of the primary ion (Hantschke, 2022).

4 Results and discussion

In order to understand the effect of HO2/RO2 on the gas-phase product composition, we will present and compare two cases: the steady state without CO (low HO2/RO2) and the steady state with CO addition and OH adjustment by J(O1D) and O3 (high HO2/RO2). The modeling results predicted HO2/RO2 of about 1/100 and of about 1/1 for these two cases, respectively. The modeled concentrations can be found in Supplement Sect. S4. The modeling results show that the HO2/RO2 ratio changes by 2 orders of magnitude, because [RO2] was reduced by about a factor of 3, while [HO2] was increased by a factor of 30. Consequently, HO2 reactions were almost negligible at low HO2/RO2 while RO2 + RO2 reactions can still contribute at high HO2/RO2.

HO2/RO2 ratios of around 1 are highly relevant for atmospheric conditions with significant OH oxidation, though it should be kept in mind that in atmospheric conditions the methyl peroxy radical and other small RO2 contribute a significant portion to the total of peroxy radicals (Khan et al., 2015). Field studies reporting HO2 and RO2 measurements for different environments can be found in the Supplement Table S5. These exemplary studies show that HO2/RO2 ratios around 1 are relevant in remote to urban environments with different VOC sources and NOx levels.

Assuming correctly modeled [HO2] and [RO2], we calculated the competition between HO2 and RO2 reactions for each (observed) RO2 expressed in the form of pseudo-first-order rate coefficients in kRO2HO2 · [HO2] or kRO2RO2 · [RO2]. Herein [RO2] is the sum of all RO2 species as defined in the MCM v3.3.1. For all experiments, the results of
our calculations indicate that the sink for HOM-RO₂ is dominated by RO₂ + RO₂ reactions at low HO₂/RO₂ (∼ 97 % contribution), while at high HO₂/RO₂ RO₂ + HO₂ contributed ∼ 80 %. As the rate coefficients are not well known and as we cannot verify the modeling results for HO₂ and RO₂, our calculations serve solely as an indication of expected trends in the chemical system.

4.1 Impact on overall HOM formation

The top panel of Fig. 2 shows the time series of HOM-Mon and HOM-Acc products. The HOM-Mon signal recovers after the oxidant adjustment, while the HOM-Acc signal is significantly suppressed at high HO₂/RO₂. This indicates that the shift from low to high HO₂/RO₂ substantially impacts the termination reactions, shifting formation from the HOM-Acc product channel (RO₂ + RO₂) to the HOM-Mon channel.

An overview of the results for the product classes defined in the method section is shown in Fig. 4. Plotted are the average ratios of signal in the NO₃-CIMS in the high and low HO₂/RO₂ steady state compared to the low HO₂/RO₂ steady state. For better comparison, all experiment phases were normalized to the actual α-pinene OH turnover. The overall HOM signal was lower at high HO₂/RO₂, showing a reduction of about 20 %. Most distinctive, the HOM-Acc compounds were strongly reduced by about 60 %. A reduction of HOM-Acc by addition of CO was observed before by McFiggans et al. (2019); however, there the OH concentration was not kept constant. The HOM-Frag signal (5 ≤ C < 10) also shows a reduction of about 20 %. At high HO₂/RO₂, the C₁₀-HOM-RO₂ signal was also reduced significantly by about 40 %.

The HOM-Mon signal level remained about the same at low and high HO₂/RO₂. Without changes in the rates and contributions of the different termination reactions, the observed reduction in the HOM-RO₂ precursors should lead to nearly the same reduction in HOM-Mon. However, the decrease in accretion product formation and fragmentation should lead to an increase in HOM-Mon. The presence of HO₂ could reduce the alkoxy formation and thus fragmentation of HOM-RO₂. This missing sink could lead to an additional HOM-Mon source compared to the low HO₂/RO₂ case. However, the distribution of the product classes at low and high HO₂/RO₂ (Fig. 5) shows that contributions are shifted from HOM-Acc to HOM-Mon, while the contribution of HOM-Frag remains constant. Each HOM-Acc is formed from one HOM-RO₂ (HOM-RO₂ + RO₂) or potentially even two HOM-RO₂ compounds (HOM-RO₂ + HOM-RO₂) and therefore each HOM-Acc not formed will lead to at least one HOM-Mon.

Further changes in the product distribution become evident when considering the individual HOM-Mon families as shown in Fig. 6. The C₁₀H₁₅O₄ peroxo radical family and the related C₁₀H₁₄O₂ family (carbonyl compounds) show the strongest suppression with a decrease of about 40 % at high HO₂/RO₂. For the C₁₀H₁₇O₂ peroxo radical family,
4.2 Impact on HOM-RO₂

C₁₀-HOM-RO₂ compounds are key to understand the changes in the HOM product distribution. Therefore, we will first discuss the changes in the HOM-RO₂ products and then the changes in the closed-shell products.

The C₁₀ peroxide radical class consists of the C₁₀H₁₅O₄ and C₁₀H₁₇O₄ families, which were reduced to 54 % and 82 %, respectively, when comparing the high and low HO₂/RO₂ cases (Fig. 6, light blue and green bars). The observed reduction in C₁₀-HOM-RO₂ is significantly smaller than the overall RO₂ concentration reduction predicted by the MCM model results (reduction to ∼ 30 %). In the following paragraphs, we present a plausibility consideration to assess if these observed changes are consistent with our expectations from modeling results and reaction rates.

The change in the steady-state concentration of a compound is always defined by the changes in its sources and sinks. The source of a HOM-RO₂ is the intramolecular reaction of a precursor RO₂; thus, the HOM-RO₂’s source is reduced if the steady-state concentration of the precursor RO₂ is reduced. However, assuming the source term of the precursor RO₂ is the same in low and high HO₂/RO₂ (due to the constant α-pinene OH turnover) and the precursor RO₂’s sink term is dominated by the fast autoxidation in both cases, then the RO₂’s steady-state concentration would not be significantly changed. This consideration is only applicable for RO₂ where autoxidation dominates the sink term at low and high HO₂/RO₂. However, the unchanged oxidation level of the HOM-Mon indicates that once the autoxidation is initiated it out-competes the possible termination reactions.

In this case, the change in steady-state concentration of the HOM-RO₂ will be defined by the changes in the sink terms. Owing to the faster reaction of RO₂ + HO₂ compared to RO₂ + RO₂ the chemical sink for all RO₂ including HOM-RO₂ with slower autoxidation rates increased, which leads to a reduction in the steady-state concentration of RO₂ in general, despite holding the primary RO₂ source term constant.

For steady-state conditions, we can estimate the expected effect on the RO₂ ratio between high and low HO₂/RO₂ conditions for those HOM-RO₂ with production directly linked to the primary production (k[OH]·[HO₂]·[α-pinene]) with negligible further autoxidation. The necessary equations and assumptions can be found in Supplement Sect. S7. We assume the same primary production at low and high HO₂/RO₂ and that the reaction with HO₂, the reaction with RO₂ and the wall loss are the only significant loss pathways. At high HO₂/RO₂, a reduction to 80 % is expected if the chosen bulk rate constants are used (k[RO₂HO₂] = 2.46 × 10⁻¹¹ cm³ s⁻¹ at 20 °C (Jenkin et al., 1997; Saunders et al., 2003); k[RO₂RO₂] = 5.0 × 10⁻¹² cm³ s⁻¹). A reduction to 60 % is expected if k[RO₂HO₂] is around 7 times faster than k[RO₂RO₂] (k[RO₂RO₂] = 3.3 × 10⁻¹² cm³ s⁻¹). These reductions are in the range of what is observed for the C₁₀-HOM-RO₂. Of course, the approach of using generalized bulk rate constants is limited, but the resulting values for k[RO₂RO₂] were clearly within the range of rate coefficients expected for HOM-RO₂ + RO₂ reactions (Roldin et al., 2019), showing that the increased chemical sink is a plausible explanation for our observations.
Since C$_{10}$H$_{15}$O$_x$ family is on average reduced by around 30% more than the C$_{10}$H$_{17}$O$_x$ family (see Fig. 6). C$_{10}$H$_{15}$O$_x$ peroxy radicals are either formed by sequential oxidation of α-pinene, e.g., from oxidation products like pinonaldehyde, or directly from α-pinene via the H-abstraction pathway (Shen et al., 2022). Formation of pinonaldehyde and even more so HOM formation via the H-abstraction channel involve alkoxy steps. However, alkoxy radicals should be reduced at high HO$_2$/RO$_2$ because they are mainly formed by RO$_2$+RO$_2$ reactions in the absence of NO$_x$. Thus, missing source terms add to the increased chemical sink by HO$_2$ for C$_{10}$H$_{15}$O$_x$ peroxy radicals.

Amine CIMS measurements enabled detection of the formula composition C$_{10}$H$_{16}$O$_2$ (e.g., pinonaldehyde). C$_{10}$H$_{16}$O$_2$ was reduced on average to 70% ± 1% at high HO$_2$/RO$_2$ compared to low HO$_2$/RO$_2$. This supports that a fraction of the C$_{10}$H$_{15}$O$_x$ radical decrease at high HO$_2$/RO$_2$ arose from suppression of C$_{10}$H$_{16}$O$_2$ first generation products. In addition, a further suppression of HOM formation via the H-abstraction channel is likely. It should be noted that the reduction of C$_{10}$H$_{16}$O$_2$ is smaller than that expected by the MCM model results. Modeling results can be found in Supplement Sect. S4. This might indicate that HO$_2$ can also enable alkoxy radical steps to a certain degree as summarized by Jenkin et al. (2019) and postulated by e.g., Eddingsaas et al. (2012) as a source of pinonaldehyde in HO$_2$ dominated systems.

According to the model calculations, the pseudo-first-order rate coefficient $k_{RO2HO2}$·[HO$_2$] is expected to be about 0.04 s$^{-1}$ for the RO$_2$+HO$_2$ reaction at high HO$_2$/RO$_2$. Consequently, only such HOM-RO$_2$ with autoxidation rates of ≤ 0.04 s$^{-1}$ will be significantly lost by reaction with HO$_2$ at the higher HO$_2$ concentrations. However, typical isomerization rates of peroxy radicals in autoxidation are on the order of 0.1 s$^{-1}$, and many are faster (Piletic and Kleindienst, 2022; Berndt, 2021). Therefore, reduction in a HOM-RO$_2$ is only expected when the faster termination rate of $k_{RO2HO2}$·[HO$_2$] can compete with the autoxidation rate, i.e. when the autoxidation slows as the degree of oxidation increases on the specific HOM-RO$_2$. This consideration shows that the smaller reduction in HOM-RO$_2$ compared to the less oxidized RO$_2$ in the model is compatible with fast autoxidation reactions that are missing in the MCM.

The increase in chemical sink strength by going from RO$_2$ termination to HO$_2$ termination is the main expected reason for the decrease in C$_{10}$H$_{17}$O$_{32}$. As discussed, the C$_{10}$H$_{15}$O$_x$ family is subject to an additional decrease in the precursors due to the alkoxy steps necessary in the formation pathway. Since C$_{10}$H$_{15}$O$_x$ compounds were the main contributors to the C$_{10}$-HOM-RO$_2$ class, their stronger reduction is reflected in the overall reduction of C$_{10}$-HOM-RO$_2$.

### 4.2.1 Contribution of C$_{10}$H$_{15}$O$_x$ and C$_{10}$H$_{17}$O$_x$ families to HOM-RO$_2$

In the unseeded, pure gas-phase experiments, the contribution of the C$_{10}$H$_{17}$O$_x$ family to the C$_{10}$-HOM-RO$_2$ class is 23% ± 2% on average in the low HO$_2$/RO$_2$ case. In the high HO$_2$/RO$_2$ case, the contribution increases to 31% ± 4% on average. As discussed above, the suggested pathways to C$_{10}$H$_{15}$O$_x$ HOM-RO$_2$ may be additionally suppressed due to a decrease in alkoxy steps at high HO$_2$/RO$_2$, reducing the entry channel into C$_{10}$H$_{15}$O$_x$ HOM-RO$_2$.

Nevertheless, the contribution of C$_{10}$H$_{15}$O$_x$ is substantial in both experiment stages. Kang (2021) and Shen et al. (2022) reported that, in the photooxidation of α-pinene, the HOM-RO$_2$ compounds detected by NO$_3$-CIMS are dominated by the C$_{10}$H$_{15}$O$_x$ family, while C$_{10}$H$_{17}$O$_x$ formation is the main expected OH reaction pathway described in the literature (Berndt, 2021; Berndt et al., 2016; Xu et al., 2019).

This hints towards an effective pathway to HOM via C$_{10}$H$_{15}$O$_x$. A reason may be the fast opening of both carbon rings in the bicyclic α-pinene (Shen et al., 2022) or a four-ring opening in pinonaldehyde or similar compounds for easy autoxidation. From our observations, increasing the HO$_2$/RO$_2$ ratio does increase the relative importance of the C$_{10}$H$_{17}$O$_x$ family, but the change is less than 10% in contribution.

Contribution of the two peroxy radical families to the HOM formation is also reflected in the composition of C$_{20}$ HOM-Acc. Figure 7 shows the average contributions of the C$_{20}$H$_{10}$O$_2$, C$_{20}$H$_{32}$O$_x$ and C$_{20}$H$_{34}$O$_x$ families in the low and high HO$_2$/RO$_2$ cases. Although the absolute amount of HOM-Acc was suppressed by 60% the family distribution was similar, C$_{20}$H$_{32}$O$_x$ dominated, while C$_{20}$H$_{30}$O$_2$ was lowest. C$_{20}$H$_{30}$O$_2$ is formed from two members of the C$_{10}$H$_{15}$O$_x$ family, while C$_{20}$H$_{34}$O$_x$ is formed by two members of the C$_{10}$H$_{17}$O$_x$ family. C$_{20}$H$_{32}$O$_x$ is then a combination of a C$_{10}$H$_{15}$O$_x$-RO$_2$ and C$_{10}$H$_{17}$O$_x$-RO$_2$.

Families that require one or two C$_{10}$H$_{17}$O$_x$ peroxy radicals for their formation have a higher contribution than the C$_{10}$H$_{15}$O$_x$ family’s contribution to C$_{10}$-HOM-RO$_2$. Here, it is important to note that not only HOM-RO$_2$ can participate in HOM-Acc formation but also traditional, less oxidized RO$_2$ radicals (Berndt et al., 2018; Pullinen et al., 2020; McFiggans et al., 2019), which are not detectable by NO$_3$-CIMS. However, more oxidized peroxy radicals exhibit faster accretion rates (Berndt et al., 2018).

The large contributions of C$_{20}$H$_{32}$O$_2$ and C$_{20}$H$_{34}$O$_2$ thus clearly show the general importance of the C$_{10}$H$_{17}$O$_x$ peroxy radicals. The largest fraction, the C$_{20}$H$_{32}$O$_2$ family, indicates the importance of HOM-C$_{10}$H$_{33}$O$_x$, and a high abundance of less oxidized C$_{10}$H$_{17}$O$_x$ peroxy radicals. Less oxidized C$_{10}$H$_{17}$O$_x$-RO$_2$ compounds were recently measured by Berndt (2021). The fraction of C$_{20}$H$_{34}$O$_x$ is smaller because their formation requires HOM-C$_{10}$H$_{17}$O$_x$ radicals which are less abundant compared to HOM-C$_{10}$H$_{15}$O$_x$. 

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Figure 7. Average contribution of the $C_{20}H_{30}O_2$, $C_{20}H_{32}O_2$, and $C_{20}H_{34}O_2$ families to the $C_{20}$ HOM-Acc group signal in the low and high $HO_2/RO_2$ cases (unseeded experiments). Not pictured is $C_{20}H_{28}O_2$ due to its negligible signal (contribution $\sim 1\%$).

while the small fraction of $C_{20}H_{30}O_2$ indicates that, despite the importance of HOM-$C_{10}H_{15}O_x$, less oxidized $C_{10}H_{15}O_x$ are less important.

These results indicate the importance of mixed HOM-Acc formation by cross-reactions of HOM-RO and a less oxidized RO$_2$. The importance of mixed HOM-Acc is supported by the relatively small fractions of HOM-Acc products with very high oxygen numbers, which more likely stem from HOM-RO$_2$+HOM-RO$_2$. For example, $C_{20}$-HOM-Acc compounds with 12 or more oxygen atoms contribute only around 30% (low $HO_2/RO_2$: 26% $\pm$ 4%; high $HO_2/RO_2$: 31% $\pm$ 2%) of the signal in the product group.

Although the effect of the changed $HO_2/RO_2$ ratio is small, a tendency to higher $C_{20}H_{34}O_2$ contribution was observed. This is consistent with the observation of a slightly higher $C_{10}H_{17}O_3$ contribution to the $C_{10}$-HOM RO$_2$. The stronger suppression of the $C_{10}H_{15}O_x$ family at high $HO_2/RO_2$ is the first indication for, and can be explained by a reduction in the alkoxy radical formation.

4.2.2 Impact on HOM-alkoxy radical formation

Alkoxy radicals (RO) are the second important radical type in the oxidation chain of $\alpha$-pinene. RO compounds cannot be detected directly as they are highly unstable and thus have very low concentrations. However, as explained in Sect. 2.2 the parity change in the HOM-RO$_2$ families can be used as a diagnosis tool for the abundance of alkoxy steps (Kang, 2021). A second indicator for alkoxy steps is the abundance of HOM products with less than 10 carbon atoms.

Figure 8 shows the average contribution of $C_{10}H_{15}O_x$ and $C_{10}H_{17}O_x$ with an even and odd number of oxygens at low and high $HO_2/RO_2$. $C_{10}H_{15}O_x$ radicals with an even number of oxygens contribute on average 32% at low $HO_2/RO_2$. For $C_{10}H_{15}O_x$, the antioxidation chain is expected to start from an even number of oxygens either from $C_{10}H_{15}O_4$ (pinonaldehyde-like) (MCM v3.3.1 (Jenkin et al., 1997; Saunders et al., 2003) or from $C_{10}H_{15}O_2$ (C10H6 H abstraction) (Berndt, 2021; Shen et al., 2022). Therefore, without the involvement of an alkoxy step, the parity of the oxygen number in the observed $C_{10}H_{15}O_x$ HOM-RO$_2$ is expected to be even. Due to the average contribution of $C_{10}H_{15}O_{odd}$ of 69%, we conclude that at least one alkoxy step (or any odd number of alkoxy steps) must have taken place in most of the cases at low $HO_2/RO_2$.

At high $HO_2/RO_2$, $C_{10}H_{15}O_{even}$ contributed 63%, and the $C_{10}H_{15}O_{odd}$ contribution was reduced to 37%. This demonstrates a change in the number of alkoxy steps along the formation pathway of the observed HOM-RO$_2$ radicals. The increased contribution of $C_{10}H_{15}O_{even}$ at high $HO_2/RO_2$ lets us infer an even number of alkoxy steps as more common (0, 2, 4, ...). In the simplest case, one alkoxy step takes place at low $HO_2/RO_2$ due to HOM-RO formation from HOM-RO$_2$+RO$_2$ reactions, while no alkoxy steps take place at high $HO_2/RO_2$, because HOM-RO$_2$+HO$_2$ produces none or less HOM-RO than HOM-RO$_2$+RO$_2$.

For $C_{10}H_{17}O_3$, the entry channel into antioxidation is $C_{10}H_{17}O_3$ with an odd number of oxygen atoms. Therefore, in antioxidation without alkoxy steps the oxygen parity is expected to be odd. At low $HO_2/RO_2$ $C_{10}H_{17}O_{odd}$ species contribute 75% to the total $C_{10}H_{17}O_3$ signal indicating that either none or an even number (2, 4, ...) of alkoxy steps occurred. At high $HO_2/RO_2$ the odd contribution increases to 84% (see Fig. 8). This result could indicate a low occurrence of alkoxy steps even at low $HO_2/RO_2$, with a further decrease in alkoxy formation at high $HO_2/RO_2$. However, the observed shift is minor.

In any case, the different responses of the $C_{10}H_{15}O_x$ and $C_{10}H_{17}O_x$ families to the reduction of HOM-RO$_2$ formation from HOM-RO$_2$+RO$_2$ at high $HO_2/RO_2$ indicate that there could be fundamental differences in the antioxidation chains of $C_{10}H_{15}O_x$ and $C_{10}H_{17}O_x$ (or the limit of the parity analysis). The parity analysis indicates a decrease in alkoxy steps at high $HO_2/RO_2$, but it cannot be directly inferred with certainty. However, the decrease in alkoxy steps at high $HO_2/RO_2$ is supported by the observation of changes in HOM-Frag products.

On average the sum of all HOM-Frag products (detected compounds with $5 \leq C < 10$ by NO$_3$-CIMS) showed a reduction of around 20% (unseeded experiments; see Fig. 4). Further trends become recognizable when separating the species

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according to their carbon number. Figure 9 shows the C5, C7, C8 and C9 HOM-Frag compounds at high HO2/RO2 compared to the low HO2/RO2 case, normalized to the α-pinene OH turnover. The fragment group with C6 compounds is not included, as it contributed less than 5% of the fragment signal and contained few detected compounds.

Figure 9 shows a significant reduction in HOM-Frag with shorter carbon chain length: C5 HOM-Frag compounds are reduced by around 60% compared to the low HO2/RO2 case. When we assume that the fragmentation of C10 compounds happens in consecutive steps via scission of HOM-RO radicals (analogously to the MCM), this observation is in accordance with decreasing importance of alkoxy radical formation at high HO2/RO2.

Overall, all observations indicate strong involvement of RO in HOM formation as well as a reduced, but still significant, involvement of RO at high HO2/RO2, when HO2 chemistry dominates. This is supported by the change of the oxygen parity in C10-HOM-RO2 and the decrease of fragmentation products, especially with lower carbon number, as well as the only moderate reduction in the observed C10H16O2 product (pinonaldehyde) and the still substantial importance of the C10H15O4 HOM-RO2 family at high HO2/RO2.

4.3 Impact on carbonyl and hydroperoxide formation

Increased HO2/RO2 should shift the product distribution by reduction of alcohol and carbonyl compounds from the so-called molecular channel in the RO2+RO2 reaction (see Reaction R3) in favor of hydroperoxide formation from RO2+HO2 termination (Reaction R1). This effect can be best observed in the C10H18O2 family, which contains the hydroperoxide and alcohol termination products arising from C10H17O4, C10H18O2 significantly increased to, on average, 159% (see Fig. 6). This supports an increased hydroperoxide formation but with some uncertainty due to the alcohol termination products from C10H17O4 (by reaction with RO2). To elucidate this further, the contribution of individual species to the C10H18O2 family was examined.

Formation of an alcohol via the molecular path (Reaction R3) leads to the loss of one oxygen atom compared to the precursor C10H17O2 radical, while in the hydroperoxide formation (Reaction R1) the oxygen number remains the same. The most abundant member of the C10H17O3 family was C10H17O7 with a contribution of 72% ± 6% at low HO2/RO2 and a contribution of 82% ± 1% at high HO2/RO2. C10H17O7 terminates to C10H18O2 products either as an alcohol with sum formula C10H18O6 or as a hydroperoxide with sum formula C10H18O7. These products have additional sources from C10H17O6 and C10H17O8, but due to the dominant contribution of C10H17O7 to the C10H17O family we expect any other production channels to be of minor importance.

Figure 10 shows the HOM product distribution within the C10H18O2 family at low and high HO2/RO2. The sum of the O6 and O7 product did not change significantly in the two regimes (about 88%), showing that these are the major products and agreeing well with the observation of C10H17O7 being the major C10H17O family signal at low and high HO2/RO2. At low HO2/RO2, the O6 product has a larger contribution of 64% ± 8%, while at high HO2/RO2 ∼ 30% of signal is shifted to the O7 product. This shows that the increase in C10H18O2 is matched with an increase in hydroperoxide formation.

An indicator for carbonyl formation is the C10H14O2 family as it only contains the carbonyl products arising from C10H15O2. The C10H14O2 family was reduced on average to 61% at high HO2/RO2; however, this decrease matches the decrease in the C10H15O4 precursor family. If the reaction of a C10H15O4-HOM-RO2 with a second RO2 were the main formation pathway for C10H14O4, a stronger reduction would be expected as both precursor species were decreased significantly. Instead, it appears that C10H14O2 is mainly impacted by the decrease in C10H15O4 as their reductions are similar. A possible explanation could be that intramolecular termination is a major reaction pathway for C10H15O4-RO2, forming C10H14O4 carbonyls. Intramolecular termination of the autoxidation chain has been discussed in the literature for different VOCs (Shen et al., 2021; Guo et al., 2022). Rissanen et al. (2014) discussed the possible importance of the unimolecular termination via an H shift, followed by formation of a carbonyl functional group and
OH loss in the autoxidation chain of cyclohexene. Piletic and Kleindienst (2022) calculated fast reaction rate constants in the range of 1–30 s$^{-1}$ for such intramolecular termination reactions to carbonyls for some C$_{10}$H$_{17}$O$_x$ in the $\alpha$-pinene photooxidation, indicating that this pathway could also be significant for C$_{10}$H$_{15}$O$_x$. However, more investigation is necessary.

The overall contributions of the C$_{10}$H$_{16}$O$_2$, C$_{10}$H$_{16}$O$_2$, and C$_{10}$H$_{18}$O$_2$ families to the HOM-Mon class at high HO$_2$/RO$_2$ are shifted as shown in Fig. 11.

The contribution of C$_{10}$H$_{16}$O$_2$ is largest and remains similar in both cases, matching the already shown unchanged signal level in Fig. 6. This is the case because the C$_{10}$H$_{16}$O$_2$ family contains the alcohols from C$_{10}$H$_{15}$O$_x$ + RO$_2$, carbonyls from C$_{10}$H$_{17}$O$_x$ + RO$_2$ and hydroperoxides from C$_{10}$H$_{15}$O$_x$ + HO$_2$ (see Fig. 1). A separation of the effects of enhanced HO$_2$ on this monomer family is difficult, as for the case where RO$_2$ termination dominates versus the case where HO$_2$ termination dominates, the loss of carbonyls and alcohols is partially compensated by the gain of hydroperoxides. A strong gain in hydroperoxides is clearly reflected in the strong increase in C$_{10}$H$_{18}$O$_2$ at high HO$_2$/RO$_2$.

Inspection of the C$_{10}$H$_{14}$O$_2$ and C$_{10}$H$_{15}$O$_2$ families shows that $\sim$13% of the contribution by C$_{10}$H$_{16}$O$_2$ is lost (carbonyls, 33% at low HO$_2$/RO$_2$) and is present instead as C$_{10}$H$_{18}$O$_2$ (hydroperoxides), giving C$_{10}$H$_{18}$O$_2$ a contribution of 38% at high HO$_2$/RO$_2$.

## 4.4 Impact on condensable organic mass

In the previous sections, we demonstrated a shift of the product distribution by the shift from low to high HO$_2$/RO$_2$ conditions. We also showed that the changes could be rationalized by generic mechanistic considerations. We added (NH$_4$)$_2$SO$_4$ seed aerosol in two experiments to determine how the shift in the product distribution affects the condensable organic mass by determining the fraction which remained in the gas phase after seeding.

Figure 12 shows the fraction remaining for the sum of all products as well as for the individual product classes for the high and the low HO$_2$/RO$_2$ case. In both cases a significant reduction of products in the gas phase was observed with seed present. Overall, the sum of all products was reduced by around 60%, with a slightly higher reduction in the low HO$_2$/RO$_2$ case. This can be attributed to the larger importance of HOM-Acc in the low HO$_2$/RO$_2$ case, as well as to a 10% lower reduction of the HOM-Frag in the high HO$_2$/RO$_2$ case. In both cases a reduction of the HOM-RO$_2$ is observed, which indicates that the provided particle sink could have affected HOM formation chemistry but only moderately.

The total organic particulate mass was determined by AMS measurements and was 2.0 and 3.4 µg m$^{-3}$ at high and low HO$_2$/RO$_2$ in the experiment (Exp2) displayed in Fig. 12. A reduction of condensed organic mass to 73% ± 3% at high HO$_2$/RO$_2$ (orange bar in Fig. 14) was observed on average. Since non-seeded and seeded experiments were conducted at otherwise the same conditions and we did not observe significant new particle formation, the gas-phase compositions can be directly compared. Therefore, we conclude that the shift in the product distribution led to a reduction of condensable material at the same $\alpha$-pinene turnover with OH (and O$_3$).

We calculated the wall-loss-corrected SOA yields with the corrected SOA mass as shown in Eq. (4) and as described by Sarrafzadeh et al. (2016). To this end, we used C$_{10}$H$_{16}$O$_7$ as the lead HOM compound. In the two experiments with seed present (Exp2.1 and Exp3) we had SOA yields of 7.3% and 10.0% at high HO$_2$/RO$_2$ and 10.0% and 12.8% at low HO$_2$/RO$_2$. The difference in the SOA yields between experiments can be explained by the slightly different OH concentrations and subsequent difference in contribution by photooxidation (see Table 1). Overall, our yields are in the lower range in comparison with the SOA yields reported by McFiggans et al. (2019) for the $\alpha$-pinene photooxidation. However, our experiments were also performed at 5 °C higher temperature (20 °C) compared to 15 °C in McFiggans et al. (2019). The SOA yields show an absolute reduction of $\sim$3% at high HO$_2$/RO$_2$ compared to low HO$_2$/RO$_2$ (relative a reduction of about 30%). A reduction of the SOA yield of $\alpha$-pinene by addition of CO was described before by McFiggans et al. (2019); however, there the $\alpha$-pinene OH turnover was not held constant.
The change from the low to high HO$_2$/RO$_2$ regime favored termination reactions to protic termination groups, as we observed less carbonyl compounds and more hydroperoxides. This could, overall, shift the product distribution to products with lower vapor pressures and favor SOA formation, since protic groups can act as hydrogen bond donors as well as hydrogen bond acceptors (as exemplified by the comparison of ethanol (boiling point (b.p.) 78 °C) and ethane hydroperoxide (b.p. 93–97 °C) with acetaldehyde (b.p. 20 °C) (Richter et al., 1955)). However, the effect of the termination group should be small for HOMs as they likely contain multiple hydroperoxide groups (compare Pullinen et al., 2020). The reduction in HOM-Acc is expected to decrease the condensable mass, since the HOM-Acc scavenge non-HOM-RO$_2$ that would otherwise not partition into the particle phase.

Which of the measured compounds contribute significantly to the organic particle mass can be inferred by comparing their signal from the pure gas-phase, unseeded cases to their signal with seed in the system. Under the assumptions that for most HOM compounds re-evaporation to the gas phase is negligible and that the precursor chemistry is not substantially disturbed by seed addition, the fraction of signal remaining with seed in the system reflects to which degree the compound is condensing. Figure 13 shows the fraction remaining with seed in the system plotted against the molar mass of each individual compound. The plot includes all closed-shell products that were detected with relative standard deviation <30% in all four experiment phases. Error bars represent result of error propagation (see Supplement Sect. S9).

Overall, in both cases we observed the same trend. Lighter compounds are not affected by the presence of seed particles, but with increasing molar mass the fraction remaining in the gas phase is reduced. A difference between the low and high HO$_2$/RO$_2$ cases can be observed in the low molar mass range: in the high HO$_2$/RO$_2$ case many fragmentation products show a higher gas-phase fraction remaining up to 1. (In some cases, values larger than 1 were observed but within the error limits. For the error estimation, see Supplement Sect. S9.) Fractions remaining larger than 1 beyond error could be an indication that such products have a particle-phase production source. Figure 13 also shows a critical semi-volatile to low-volatility organic compound molar mass region for molar masses between 175 and 250 g mol$^{-1}$ where neither a fraction remaining of 1 nor complete condensation was observed. The position of this region on the molar mass scale depends on the provided organic mass concentration. The large variation of the fraction remaining in this small range of molar masses shows that the partitioning coefficients are dependent on the detailed structure of the compounds and not simply on their molar mass. The semi-volatile and low-volatility products represent mainly higher oxidized fragments and HOM-Mon with less than eight oxygens.

For compounds with a molar mass larger than 250 g mol$^{-1}$, a constant fraction remaining is reached in steady state, which is due to an ongoing production of the compounds. From the condensation behavior shown in Fig. 13, we conclude that the compounds heavier than 230 g mol$^{-1}$ are expected to be of sufficiently low volatility to be mainly found in the particle phase for the organic mass present in the system and therefore contribute significantly to the SOA mass formation. Our finding agrees with the threshold used for low-volatility HOM products in Pullinen et al. (2020).

Therefore, the signal of all compounds with a molar mass heavier than 230 g mol$^{-1}$ was weighted with their molar mass and summed (see Eq. 2). The ratio of this weighted signal sum at low and high HO$_2$/RO$_2$ is then a measure of expected SOA mass loss. The calculation leads to an expected reduction to 72% (blue bar, Fig. 14). This simplified approach leads to a good agreement with the AMS measure-
ments and can thus explain the reduced particulate organic mass within the errors.

To test for closure between HOM lost and particulate organic mass measured, we approximated the upper limit of HOM concentration in the condensed phase. For this calculation, we used the calibration factor determined for sulfuric acid for our NO$_3$-MION-CIMS (7.0 x $10^9$ molecules cm$^{-3}$ ncps$^{-1}$, where ncps represents normalized counts per second) and the relationship between gas and particulate concentration of a compound in the SAPHIR-STAR chamber described in Eq. (3). Again, we considered all compounds with $M > 230$ g mol$^{-1}$ in our calculation. The summed mass concentration lost from the gas phase was then compared to the SOA mass measured in the AMS. This comparison yields a good agreement within the uncertainties. The detailed calculation results can be found in the Supplement (Fig. S3). Overall, an agreement within 40% is achieved for all measurement stages.

The comparisons presented above show that we understand the processes governing the SOA formation in our chamber and that the NO$_3$-CIMS measurements are well suited to observe the critical changes to understand the reduction in condensable organic material when shifting from low to high HO$_2$/RO$_2$.

5 Conclusions

In the presented series of experiments, we achieved a shift from a RO$_2$+RO$_2$-dominated chemistry to a more atmospherically relevant HO$_2$/RO$_2$ ratio under constant $\alpha$-pinene OH turnover. It was shown that moving towards atmospheric HO$_2$/RO$_2$ ratio affected the SOA formation potential, with the observed organic mass being reduced at high HO$_2$/RO$_2$. This is in support of the potential bias towards high SOA yields in chamber studies at low HO$_2$/RO$_2$ as discussed by Schervish and Donahue (2021). Our results confirm that too low HO$_2$/RO$_2$ is one important parameter that can lead to an underestimated SOA yield in laboratory studies. In a broader picture, the results show how important it is to consider the different contributions to the HOM-RO$_2$ sink (e.g., HO$_2$, RO$_2$, NO) when designing experiments and transferring laboratory results to the real atmosphere.

The gas-phase observations showed that the SOA reduction at high HO$_2$/RO$_2$ was mainly due to a reduced HOM-Acc formation, which were formed by RO$_2$+RO$_2$ cross-reactions in the low HO$_2$/RO$_2$ case. This prevented contribution to SOA by less oxidized RO$_2$ compounds which were scavenged in the HOM-Acc at low HO$_2$/RO$_2$. Under atmospheric conditions, such cross-reactions are less important, and such (mixed) accretion products would contribute less to SOA.

The overall observed HOM products were reduced slightly, showing that under certain circumstances RO$_2$+HO$_2$ termination can impede the HOM formation, mainly by reducing the precursor RO$_2$ levels and less by impeding the autoxidation itself. The autoxidation chain (once initiated) runs to a similar oxidation level at both high and low HO$_2$/RO$_2$. The observed HOM-Mon products shift significantly between monomer families due to the different termination reactions. A decrease in carbonyl and alcohol formation from RO$_2$+RO$_2$ and an increase in hydroperoxide formation from RO$_2$+HO$_2$ were observed at high HO$_2$/RO$_2$.

Furthermore, a reduction in HOM-Frag products, especially with lower carbon numbers, and a reduction in the parity of the C$_{10}$H$_{15}$O$_x$ HOM-RO$_2$ show a reduction in alkoxy radical formation at high HO$_2$/RO$_2$. The moderate reduction in larger HOM-Frag products and pinonaldehyde, however, suggests that some alkoxy radical steps are still important. This raises the question of whether alkoxy radical formation can be facilitated by HO$_2$. In the atmosphere such effects are most often overcome whenever RO$_2$+NO is the major alkoxy radical source.

Overall, the observed changes in the gas phase could be well explained with the presented generic mechanistic understanding of HOM formation in the $\alpha$-pinene system. The addition of seed particles demonstrated that the shift towards high HO$_2$/RO$_2$ reduced the condensable organic mass, stressing the importance of controlling higher-order reactions of peroxy radicals which lead to overemphasis on HOM-Acc product formation at low HO$_2$/RO$_2$ ratios.

Furthermore, the seed addition allowed us to determine which products were contributing to the SOA formation and show that their volatility is a function of molar mass and detailed molecular structure. This revealed a critical molar mass region in which compounds have significant fractions in gas and particulate phases. Based on absorptive partitioning theory, the volatilities at which this critical region is found should depend on the organic mass present in the system.

Valuable insight about the condensed phase can be gained from HOM gas-phase measurements. We inferred conclusions about the particulate phase from the gas-phase measurements and compared them to the direct particle-phase observations, finding good agreements between our expectations and the measurements.

Data availability. Information about all steady-state conditions utilized and the data necessary to reproduce the figures in this study are available at https://doi.org/10.26165/JUELICH-DATA/R8ITFF (Baker et al., 2024).

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