Impact of NO\textsubscript{x} and OH on secondary organic aerosol (SOA) formation from β-pinene photooxidation

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Abstract

In this study, the NO$_x$ dependence of secondary organic aerosol (SOA) formation from β-pinene photooxidation was comprehensively investigated in the Jülich Plant Atmosphere Chamber. Consistent with the results of previous NO$_x$ studies we found increases of SOA yields at low NO$_x$ conditions ([NO$_x$]$_0$ < 30 ppb, [BVOC]$_0$/[NO$_x$]$_0$ > 10 ppbC ppb$^{-1}$). Furthermore, increasing [NO$_x$] at high NO$_x$ conditions ([NO$_x$]$_0$ > 30 ppb, [BVOC]$_0$/[NO$_x$]$_0$ ~10 to ~2.6 ppbC ppb$^{-1}$) suppressed the SOA yield. The increase of SOA yield at low NO$_x$ conditions was attributed to increase of OH concentration, most probably by OH recycling in NO + HO$_2$ → NO$_2$ + OH reaction. Separate measurements without NO$_x$ addition but with different OH primary production rates confirmed the OH dependence of SOA yields. After removing the effect of OH concentration on SOA mass growth by keeping the OH concentration constant, SOA yields only decreased with increasing [NO$_x$]. Measuring the NO$_x$ dependence of SOA yields at lower [NO]/[NO$_2$] ratio showed less pronounced increase in both; OH concentration and SOA yield. This result was consistent to our assumption of OH recycling by NO and to SOA yields being dependent on OH concentrations. It furthermore indicated that NO$_x$ dependencies vary for different NO$_x$ compositions. A substantial fraction of the NO$_x$-induced decrease of SOA yields at high NO$_x$ conditions was caused by NO$_x$-induced suppression of new particle formation (NPF). This was shown by probing the NO$_x$ dependence of SOA formation in the presence of seed particles. After eliminating the effect of NO$_x$-induced suppression of NPF and NO$_x$ induced changes of OH concentrations, the overall effect of NO$_x$ on the SOA yield from β-pinene photooxidation was moderate. Comparing with β-pinene experiments, the SOA formation from α-pinene photooxidation was only suppressed by increasing NO$_x$. However, basic mechanisms of the NO$_x$ impacts were the same as that of β-pinene.
1. Introduction

Biogenic volatile organic compounds (BVOC), such as monoterpenes (C$_{10}$H$_{16}$) are emitted in large quantities into the atmosphere (Guenther et al., 1995, 2012; Griffin et al., 1999a). These BVOCs are oxidized in the atmosphere by hydroxyl radicals (OH), ozone (O$_3$), or nitrate radicals (NO$_3$) resulting in the formation of secondary organic aerosol (SOA). SOA contributes to a substantial fraction of ambient organic aerosol and is known to adversely affect visibility, climate and human health (Hallquist et al., 2009).

SOA formation potentials of BVOC species are represented by SOA yields which are generally defined as the ratio of the SOA mass produced from the oxidation of the SOA precursor to the mass of the precursor consumed (Odum et al., 1996). Despite the fact that many studies have focused on the production of SOA from a number of monoterpenes, reported SOA yields have shown high variability for a given precursor (Pandis et al., 1991; Hoffmann et al., 1997; Griffin et al., 1999b; Larsen et al., 2001; Presto et al., 2005; Kroll et al., 2006; Ng et al., 2007; Mentel et al., 2009; Eddingsaas et al., 2012a). For instance, the reported SOA mass yield for α-pinene photooxidation ranges from 8 % to 37 % (Eddingsaas et al., 2012a). This variability is likely related to the numerous factors that influence the SOA yields, such as the inorganic and organic mass loading, particle acidity, NO$_x$ (NO$_x$ = NO + NO$_2$) level, humidity, and temperature. Therefore, ambient SOA yields cannot be represented by a unique value for a given monoterpene as the yields are heavily dependent on the conditions under which the SOA is formed.

One of the critical factors is the impact of NO$_x$ on SOA formation. Results of the majority of studies indicate that SOA yields are lower at high NO$_x$ levels (Hatakeyama et al., 1991; Pandis et al., 1991; Presto et al., 2005; Kroll et al., 2006; Ng et al., 2007). It is generally assumed that the impact of NO$_x$ results from altering the balance between competing peroxy-radical (RO$_2$)...
reactions and thus from the changes in the distribution of oxidation products. Reaction (R1) is the dominant pathway for RO\textsubscript{2} radicals under low-NO\textsubscript{x} conditions which leads to the formation of low-volatility hydroperoxides that can participate in new particle formation (NPF) and contribute to SOA mass (Johnson et al., 2005; Camredon et al., 2007).

\[
\text{RO}_2 + \text{HO}_2 \rightarrow \text{ROOH} + \text{O}_2 \quad \text{(R1)}
\]

Under high-NO\textsubscript{x} conditions, RO\textsubscript{2} radicals react with NO resulting in the formation of organic nitrates (R2a) which are suggested to be relatively volatile as well as alkoxy radicals (R2b) that either fragment, or react to form more volatile products. This understanding implies that higher NO\textsubscript{x} concentrations will suppress the formation of low volatility products, and thereby suppress NPF and SOA mass formation.

\[
\text{NO} + \text{RO}_2 \rightarrow \text{RONO}_2 \quad \text{(R2a)}
\]

\[
\text{NO} + \text{RO}_2 \rightarrow \text{RO} + \text{NO}_2 \quad \text{(R2b)}
\]

Despite numerous studies of SOA formation from terpene ozonolysis, the SOA formation from OH oxidation of β-pinene has been scarcely investigated. In the present study we investigated the SOA formation from β-pinene photooxidation under varied NO\textsubscript{x} levels in the Jülich Plant Atmosphere Chamber (JPAC) to gain more insight into the impact of NO\textsubscript{x} on SOA yield. Since the [BVOC]/[NO\textsubscript{x}] ratio varies significantly in urban atmospheres, the quantification of the effect of NO\textsubscript{x} on SOA yield of biogenic precursors is needed to better predict air pollution and to improve the accuracy of current ambient models.
2. Experimental

The experimental setup of JPAC is described in detail elsewhere (Mentel et al., 2009, 2015). The chamber is 1450 litre in volume, made of borosilicate glass and set up in a climate-controlled housing. Temperature and relative humidity inside the chamber were held constant at 16 ± 1°C and 63 ± 2%, respectively over the course of the experiments. The chamber was operated as a continuously stirred tank reactor with a residence time of approximately 46 min. The flow into the chamber consisted of two purified air streams. One stream was passed through an ozonator and was humidified with double distilled water. The other stream contained β-pinene emitted from a diffusion/permeation source held at 38°C. Where necessary seed particles could be generated externally and introduced using a third air stream.

The chamber was equipped with several lamps; 12 discharge lamps (HQI 400 W/D; Osram) to simulate the solar light spectrum in the chamber, 12 discharge lamps (Phillips, TL 60 W/10-R, 60W, \(\lambda_{\text{max}} =365\) nm, from here on termed UVA lamps) for NO\(_2\) photolysis, and one internal UVC lamp (Philips, TUV 40W, \(\lambda_{\text{max}} =254\) nm; here termed as TUV lamp) for ozone photolysis to produce OH radicals by reaction of water vapour with O(1D) atoms. The TUV lamp could be shielded by glass tubes to control the amount of UV radiation entering the chamber. Thus, by altering the gap between these glass tubes, the OH production rate could be adjusted by varying the photolysis rate J(O(1D)). It has to be noted that the short wavelength cut off of the glass is around 350 nm and thus no light with wavelength short enough to produce O(1D) is in the chamber when the TUV lamp is off. Furthermore, the absorption cross section of NO\(_2\) at the wavelength of the TUV lamp is more than an order of magnitude lower than at wavelengths around 365 nm (Davidson et al., 1988). Together with the quite low energy of the TUV lamp compared to the energy of the UVA lamps, this allowed varying J(O(1D)) and J(NO\(_2\)) independent of each other.
A suite of instruments were used to measure both the gas and particle phase products. Ozone concentration was determined by UV photometric devices (Thermo Environmental 49 and Ansyco, O₃ 42M ozone analyzers), NO was measured by chemiluminescence (Eco Physics, CLD 770 AL ppt), NO₂ by chemiluminescence after photolysis (Eco Physics, PLC 760) and relative humidity was measured by dew point mirror (TS-2, Walz). Furthermore, a condensation particle counter (CPC, TSI 3783) and a scanning mobility particle sizer (SMPS, combination of a TSI 3081 electrostatic classifier and a TSI 3025 CPC) were used to count the total particle number greater than 3 nm and to measure the particle size distribution between 13 and 740 nm respectively. The total particle mass concentration was estimated from the measured total aerosol volume assuming a SOA density of ~ 1.2 g cm⁻³ and spherical particles. β-pinene mixing ratio in the chamber was determined by gas chromatography–mass spectrometry (GC-MS, Agilent GC-MSD system with HP6890 GC and 5973 MSD) and a proton transfer reaction mass spectrometer (PTR-MS, Ionicon). The GC-MS and PTR-MS were switched periodically between the outlet and the inlet of the chamber to quantify concentrations of β-pinene entering and exiting the chamber. The OH concentration was estimated from the decay of β-pinene in the chamber (Eq. 2) (Kiendler-Scharr et al., 2009).

\[
\frac{d[\beta p]}{dt} = \frac{F}{V} \left( [\beta p]_{in} - [\beta p] \right) - \left( k^{OH} \cdot [OH] + k^{O3} \cdot [O_3] \right) \cdot [\beta p]
\]

(1)

\[
[OH] = \frac{F \cdot [\beta p]_{in} - [\beta p]}{[\beta p]} \cdot \frac{k^{O3} \cdot [O_3]}{k^{OH}}
\]

(2)

Equation (1) is the basic rate equation for a continuously stirred tank reactor resulting from mass balance and Eq. (2) results from Eq. (1) under steady state conditions when solving for [OH]. In Eqs. (1) and (2), V is the volume of the chamber and F is the total air flow through the chamber.
[βp] in and [βp] are the concentrations of β-pinene in the inlet air and in the chamber, respectively. For a well-mixed continuously stirred tank reactor, [βp] is the concentration of β-pinene measured in the outlet flow. k\textsuperscript{OH} and k\textsuperscript{O3} are the rate constants of reactions of β-pinene with OH and with O\textsubscript{3}. Since β-pinene has a quite low rate constant with O\textsubscript{3} (k\textsuperscript{O3} = 1.5 \times 10^{-17} \text{ cm}^3 \text{ s}^{-1} (Atkinson and Arey, 2003)), the reaction of β-pinene with O\textsubscript{3} could be neglected from Eq. (2) for our ozone- (50-100 ppb) and OH concentrations (9 \times 10^6 – 1.6 \times 10^8 \text{ cm}^3 \text{ s}^{-1}). The uncertainty in OH concentration was estimated to be approximately 20% (Wildt et al., 2014).

SOA yields were determined as described in Mentel et al. (2009), by calculating incremental yields from the formed particle mass as a function of consumed precursor mass. However, different from the procedure described in Mentel et al. (2009), here we use particle masses corrected for wall losses of extremely low volatile organic compounds (ELVOC) that are direct particle precursors.

Briefly, wall losses and losses on particles were determined for the ELVOCs. Using the respective loss rates, the fraction of ELVOCs contributing to particle mass formation (dependent on particle surface) and the fraction of ELVOCs lost on chamber walls were calculated. The ratio of total loss rate divided by loss on particles was taken as correction factor allowing the determination of the particle mass that would have been obtained if there were no wall losses. Experiments with α-pinene and β-pinene verified the correction procedure for a wide range of particle surfaces. The derivation of the correction method is described in detail in the supplement. Note that this procedure is only valid for the chamber used in these experiments.

Additionally, we used another approach to determine the mass yields at steady state conditions. Correcting measured particle masses by the above mentioned procedure revealed that after an
induction time of ~ 20 minutes, the wall loss corrected particle mass was constant as long as the OH production rate was constant. This allowed determining yields using steady state assumptions. Yields were determined by dividing the particle mass formed in the experiment by the oxidation rate of the BVOC. At our method of [OH] determination, using the BVOC consumption according to Eq. (2), and at the negligible inflow of particle mass into the chamber, this equals the simple mass balance assumption: produced particle mass divided by BVOC consumption (Eq. 3).

\[ \frac{\text{Particle mass}}{\text{BVOC consumption}} \]

As was expected from the consistency of wall loss corrected particle masses, both procedure of yield determinations led to identical results. However, the method using steady state conditions had advantages since adjustments of [OH] were required during many experiments. The justification to use both type of yield determinations is given in the supplement.

2.1. Experimental procedure

A series of β-pinene photooxidation experiments were performed in the JPAC to investigate the SOA formation under low-NO\(_x\) (here defined as: [NO\(_x\)]\(_0\) < 30 ppb, [BVOC]\(_0\)/[NO\(_x\)]\(_0\) > 10) and high-NO\(_x\) (here defined as: [NO\(_x\)]\(_0\) > 30 ppb, [BVOC]\(_0\)/[NO\(_x\)]\(_0\) < 10) conditions. In these experiments, the inflow of β-pinene (95 %, Aldrich) to the chamber was kept constant, leading to initial mixing ratios of 37 ± 0.6 ppb. Initial O\(_3\) concentration was 40 ± 5 ppb. NO\(_2\) (Linde, 104 ± 3 ppm NO\(_2\) in nitrogen) was introduced into the β-pinene air stream. Initial NO\(_x\) concentrations, [NO\(_x\)]\(_0\), in the chamber were varied between < 1 ppb and 146 ppb. The chamber was illuminated with one of the HQI lamps and all the UVA lamps, resulting in an NO\(_2\) photolysis frequency (J(NO\(_2\))) of 4.3×10\(^{-3}\) s\(^{-1}\). When VOC-, NO\(_x\)- and O\(_3\) concentrations in the chamber were near to
steady state, the experiment was initiated by turning on the TUV lamp resulting in OH radical production. During the described experiments, a constant J(O¹D) was maintained. Experiments without NOₓ addition were performed between NOₓ experiments. In these cases, residual NOₓ concentrations from chamber walls were below 1 ppb.

After initiating the OH production, β-pinene and NOₓ concentrations decreased due to their reactions with OH radicals. The majority of the results presented in this study are from steady state measurements, when all physical and chemical parameters were constant in the chamber. However, for the purpose of comparison with the literature data, the initial concentration of NOₓ, [NOₓ]₀, and β-pinene, [BVOC]₀, were also used here.

To investigate the role of NOₓ on SOA formation in the presence of inorganic aerosol, β-pinene photooxidation/NOₓ experiments were repeated in the presence of seed aerosol. Seed particles were generated from a 40 mg/L aqueous (NH₄)₂SO₄ solution, passed through a diffusion drier and then introduced into the chamber. For these experiments, the organic particle mass was determined by subtracting the initial seed aerosol mass from the total particle mass.

3. Results and discussion

3.1. Impact of NOₓ on SOA formation

To determine the influence of NOₓ on SOA formation, a series of NOₓ experiments were conducted in the JPAC in which β-pinene was oxidized in the absence of inorganic seed aerosol. A summary of experimental conditions and results for the β-pinene/NOₓ photooxidation experiments is given in Table 1. Figure 1 shows SOA yields, calculated from wall-loss corrected maximum particle mass concentration (PMₘₐₓ), as a function of [BVOC]₀/[NOₓ]₀ ratio and [NOₓ]₀. The strong dependence of SOA yield on BVOC/NOₓ levels is evident. At low NOₓ
conditions an increase in the initial NO\textsubscript{x} concentration increases the SOA yield, whereas at high NO\textsubscript{x} concentrations the opposite SOA yield dependence on NO\textsubscript{x} was observed. Similar NO\textsubscript{x} dependencies of SOA yields have been observed in previous studies (Pandis et al., 1991; Kroll et al., 2006; Zhang et al., 1992; Camredon et al., 2007).

At high NO\textsubscript{x} conditions strong depletion of SOA yield as well as NPF was observed with increasing NO\textsubscript{x} (Fig. 1). Wildt et al. (2014) made similar observations for NPF during the photooxidation of a BVOC mix emitted from Mediterranean plants. However, the NO\textsubscript{x} dependence of SOA yield was differed from that shown by Wildt et al. (2014). At low NO\textsubscript{x} concentrations we observed an increasing SOA yield with increasing NO\textsubscript{x} (Fig. 1). At high NO\textsubscript{x} levels, the yields decreased. Having different NO\textsubscript{x} dependencies at different NO\textsubscript{x} regimes suggests multiple factors at play.

Kroll et al. (2006) suggested that the increase in SOA yield with NO\textsubscript{x} could be due to changes in the [NO]/[HO\textsubscript{2}] ratio. As experiments in batch reactors proceed, NO\textsubscript{x} concentrations decrease due to their reactions with OH resulting in a switch from high NO\textsubscript{x} to low NO\textsubscript{x} conditions. The lowered NO concentrations cause increasing HO\textsubscript{2} concentrations due to reaction (R3):

\[
\text{NO} + \text{HO}_2 \rightarrow \text{NO}_2 + \text{OH} \quad \text{(R3)}
\]

In such experiments, peroxy radicals initially react mainly with NO, whereas peroxy radicals formed later from first generation products, primarily react with HO\textsubscript{2}. Although the reason for the observed increase of SOA yield with increasing NO\textsubscript{x} at low NO\textsubscript{x} levels was not fully explored, Camredon et al. (2007) noted that this could be due to the influence of OH levels. However, in the majority of studies investigating the impact of NO\textsubscript{x} on SOA formation, OH
concentration was either not measured or the potential influence of OH was not discussed (Eddingsaas et al., 2012a, 2012b).

As the NO\textsubscript{x} concentration is changed in this study, the concentration of OH was found to change markedly (Fig. 2). OH concentrations passed through a maximum (~ 3.8 \times 10^7 \text{molecules cm}^{-3}) at [NO\textsubscript{x}]~40 \text{ppb} ([NO\textsubscript{x}]~70 \text{ppb}) which represented a fourfold increase over that in the absence of NO\textsubscript{x}. NO\textsubscript{x} enhanced OH production in two ways: by increasing [O\textsubscript{3}] and thereby the photolytic OH source, and by recycling OH through reaction (R3). However, at very high NO\textsubscript{x} concentrations, NO\textsubscript{x} is acting as a sink for OH due to Reaction (R4):

\[
\text{NO}_2 + \text{OH} (+\text{M}) \rightarrow \text{HNO}_3 (+\text{M})
\] (R4)

Therefore, as illustrated in Fig. 2, OH concentration increased rapidly with increasing NO\textsubscript{x}, reached a maximum value and then decreased gradually. In general terms this is consistent with the nonlinear dependence of OH concentration on NO\textsubscript{x} level in the lower Troposphere (Ehhalt and Rohrer, 1995).

It appeared in Fig. 2 that the SOA yields were somehow related to [OH]. Thus, we performed some experiments to further explore the dependence of SOA formation on OH concentration.

3.1.1. [OH] dependence of SOA mass formation

To examine whether or not SOA yield depends on the OH concentration, additional experiments were performed at two different OH production rates and the \(\beta\)-pinene concentration varied to give a range of SOA mass. The detailed experimental conditions are summarized in Table 2. Two different J(O\textsuperscript{1}D) conditions \((1.9 \pm 0.2 \times 10^3 \text{ s}^{-1} \text{ and } 5.4 \pm 0.5 \times 10^3 \text{ s}^{-1})\) were used to give significantly different OH production rates at otherwise unchanged conditions. Figure 3 shows
particle mass as a function of consumed β-pinene ranging from 20 to 140 µg m$^{-3}$. Approximately 90-95% of the total β-pinene was consumed in these experiments.

The OH concentrations were $4 \times 10^7$ – $1 \times 10^8$ molecules cm$^{-3}$ and $1.1 – 1.6 \times 10^8$ molecules cm$^{-3}$ under low and high OH conditions, respectively. The SOA yields (incremental yields, see Mentel et al., (2009) and the supplement to this paper) were higher at higher OH levels (31 ± 3% and 20 ± 1% for high and low OH conditions, respectively).

In discussing possible reasons for the OH impact, it is crucial to consider secondary reactions. As an example, the rate constant for the reaction of β-pinene + OH is higher than that of the reaction of nopinone + OH. Nopinone is a major product of β-pinene oxidation (Atkinson and Arey, 2003). Thus in the stirred flow reactor, where the oxidation of β-pinene does not go to completion, there is an appreciable concentration of nopinone. Increasing the OH concentration will therefore result in more nopinone consumption and, if nopinone oxidation also forms SOA mass, this additional oxidation forms more SOA mass. As a result, SOA mass will be higher at higher OH concentrations and thus, SOA yield based on the consumption of β-pinene will be higher. Similarly, such sequential OH reactions can also form SOA mass in reactions with other β-pinene oxidation products. Another possibility might be the OH dependence of ELVOCs formation. Formation of such molecules might require more than one OH reaction.

From our data we cannot decide which process plays a major role for the OH dependence of yields. Nevertheless, the results of these experiments affirm the importance of actual OH concentrations in SOA mass formation. This complicates the assignment of the observed changes in SOA yield to the impact of NO$_x$ on peroxy radical chemistry, as SOA yield was also likely
varied with [OH]. Thus, we undertook a series of experiments to decouple the impacts of OH and NO\textsubscript{x} on SOA formation.

### 3.1.2. Isolate the effect of [OH] on SOA formation

To examine the impact of NO\textsubscript{x} on SOA production independent of [OH]-changes, a series of experiments were performed where the steady state OH concentration was held constant by tuning the value of \( J(\text{O}^1\text{D}) \). This required constant [OH] monitoring as the system approached steady state, i.e. monitoring the consumption of \( \beta \)-pinene, to ensure the [OH] was adjusted to the desired level. Although there was a significant variation in initial OH concentrations on adding NO\textsubscript{x} if \( J(\text{O}^1\text{D}) \) was unchanged, it was possible to maintain the OH concentrations to within 5\% across all NO\textsubscript{x} concentrations by adjusting \( J(\text{O}^1\text{D}) \) (Fig. 4). Figure 5 shows the SOA yield as a function of [NO\textsubscript{x}] during the steady state before and after [OH] adjustment. Before adjusting [OH] the yield profile was consistent with our previous results; SOA yield increased with increasing NO\textsubscript{x} at low NO\textsubscript{x} levels and then dropped at high NO\textsubscript{x} levels. After adjusting [OH], the yield revealed no increase at low NO\textsubscript{x} levels. It was only suppressed by increasing NO\textsubscript{x}. This indicated that the observed increase in SOA yield without adjusting [OH] is a result of NO\textsubscript{x} enhancing [OH] and not a direct impact of NO\textsubscript{x} on the SOA production. Thus isolating the effect of [OH] revealed that increasing [NO\textsubscript{x}] only suppressed particle mass formation, and therefore also suppressed SOA mass yield.

### 3.2. Role of NO/NO\textsubscript{2} ratio in SOA formation

We also investigated the effect of the [NO]/[NO\textsubscript{2}] ratio on SOA formation from \( \beta \)-pinene/NO\textsubscript{x} mixtures. To change this ratio we changed [O\textsubscript{3}]. Ozone, NO and NO\textsubscript{2} are interrelated as illustrated in Reactions (R5) and (R6):
Neglecting reactions of NO with peroxy radicals the \([\text{NO}]/[\text{NO}_2]\) ratio can then be described by the photostationary steady state (Leighton, 1961):

\[
\frac{[\text{NO}]}{[\text{NO}_2]} = \frac{J(\text{NO}_2)}{k^5[\text{O}_3]} \tag{4}
\]

where \(J(\text{NO}_2)\) represents the photolysis rate of Reaction (R6) and \(k^5\) is the rate coefficient of Reaction (R5). Hence, adjusting \([\text{O}_3]\) in the chamber allowed varying \([\text{NO}]/[\text{NO}_2]\) ratios. Here, to probe the dependency of SOA mass formation on relative NO and NO\(_2\) concentrations, NO\(_x\) experiments were performed with approximately 50% higher \(\text{O}_3\) concentration (74 ± 7 ppb) than that of previous NO\(_x\) experiments. The remaining conditions maintained the same.

Results obtained from these experiments are shown in Fig. 6. The behaviour observed in the high-\(\text{O}_3\) experiments (equivalent to lower NO/NO\(_2\)) was different from that in the experiments with the lower \([\text{O}_3]\). When not adjusted, \([\text{OH}]\) increased with increasing NO\(_x\) but the increase was less pronounced. With a lower \([\text{NO}]/[\text{NO}_2]\) ratio, the maximum OH concentration increase was approximately twofold relative to the respective NO\(_x\) free experiments (Fig. 6), compared to the fourfold increase with the higher \([\text{NO}]/[\text{NO}_2]\) (Fig. 4). This is consistent to the assumption that reaction (R3) recycles OH. There was also a slight increase in SOA yield when \([\text{NO}_3]\) increased up to ~15 ppb. In addition, the increase in SOA yield was less pronounced in the high-\(\text{O}_3\) experiments again pointing to the role of \([\text{OH}]\) in SOA formation.

The role of \([\text{OH}]\) was also confirmed by another observation. By comparing the data without NO\(_x\) addition, it can be seen that \([\text{OH}]\) as well as the yields are higher in the high-\(\text{O}_3\)
experiments. The higher [O₃] caused a higher OH production rate and at otherwise same conditions (J(O¹D), [H₂O], etc.), higher [OH] and thereby higher yields. In summary, the results of the high-O₃ experiments confirmed our interpretations of the results from the low-O₃ experiments and again supported the key role of actual OH concentrations in SOA mass formation.

After adjusting [OH] to the same level as in the NOₓ free experiments (Fig. 6), no increase in SOA yield was observed and increasing NOₓ only suppressed SOA formation (Fig. 7). These results were consistent with those found earlier in the low-O₃ experiments indicating that, after isolating the effect of [OH], SOA yield was only suppressed with increasing NOₓ also at higher [O₃]. Comparing the yield profiles obtained from the low-O₃ and the high-O₃ experiments respectively (blue circles in Fig. 5 and Fig. 7), it can be seen that the decrease in SOA yield was~35 % in the high-O₃ experiments while it was roughly 70 % in the low-O₃ experiments. This shows that NOₓ dependencies itself depend on the composition of NOₓ. As the suppression of yield was more pronounced in the low-O₃ experiments (= higher [NO]/[NO₂]) it seemed that NO is the molecule mainly responsible for the SOA yield diminishing effect of NOₓ.

3.3. Comparison of the impact of NOₓ on SOA yield from α-pinene and β-pinene photooxidation

A series of α-pinene/NOₓ experiments were performed in the same chamber to compare the NOₓ dependencies of SOA formation of α-pinene to that of β-pinene. These experiments were performed with 12 ± 1.2 ppb α-pinene, 78 ± 14 ppb O₃ and with [NOₓ]₀ up to 126 ppb.

When using α-pinene as the SOA precursor, no increase in aerosol mass formation was observed at low NOₓ, with only suppression of the particle mass formation and the SOA yield (Fig. 8).
Furthermore, presumably due to the lower α-pinene concentrations, no particle formation at all was observed when [NO$_x$]$_0$ was above 60 ppb.

At high NO$_x$ levels, the differences between the NO$_x$ dependencies of α-pinene (Fig. 8) and β-pinene (Fig. 1) photooxidation were not very strong, both showing a decrease with increasing NO$_x$. At low NO$_x$ levels there were substantial differences, in that β-pinene showed a distinctive increase and maximum of the yield, while the yield of α-pinene is almost unaffected and monotonically decreasing. These differences can be explained by the differences in [O$_3$] during α-pinene and β-pinene photooxidation. The lower [O$_3$] in the β-pinene experiments caused higher [NO]/[NO$_2$] ratios and thus more effective conversion of HO$_2$ to OH by NO (reaction R3).

This is supported by the experiments with β-pinene performed at higher [O$_3$] that caused less NO$_x$ induced increase of [OH] (Fig. 6, black squares), as well as less NO$_x$ induced increase of SOA yield (Fig. 7, black circles). Restricting focus to the same [BVOC]/[NO$_x$] where the substantial increase in β-pinene SOA yield was observed (> 100 to ~20 ppbC ppb$^{-1}$), such increases were neither observed for α-pinene nor for the monoterpene mix emitted from Mediterranean species (Wildt et al., 2014). The NO$_x$ dependence of SOA formation therefore is different in different chemical systems. As the SOA yield was dependent on the actual OH concentrations, differences in OH recycling may be involved here as well. However at high NO$_x$ conditions, the principle behaviour of all systems was identical; a general SOA yield suppression with increasing NO$_x$.

3.4. Impact of NO$_x$ on SOA formation in the presence of seed aerosol

From Fig. 2, it can be seen that the SOA yield measured at [NO$_x$]$_{ss}$ ~ 86 ppb ([NO$_x$]$_0$ ~ 146 ppb) was lower than that at [NO$_x$]$_0$ < 1 ppb while [OH] was higher. Hence, there must be another effect of NO$_x$ addition besides its impact on [OH]. As illustrated in Fig. 9, particle number
concentrations were suppressed significantly when adding NO\textsubscript{x}. The strong decrease of particle number concentration with increasing NO\textsubscript{x} indicates that NPF is suppressed by NO\textsubscript{x} (see also Wildt et al., 2014). Hence, there is a high chance that the SOA formation is hindered by low particle phase condensational sinks at high NO\textsubscript{x} levels and thus, the observed suppression of SOA yield may be due to the suppression of NPF. At high NO\textsubscript{x} conditions, particle numbers and surfaces were quite low. Therefore, high correction factors had to be used to correct the particle mass for wall losses of ELVOCs which includes high uncertainties (see supplement). These uncertainties can be diminished by using seed particles since they provide a surface onto which the low volatile organics may condense. In the presence of seed aerosol, the growth of particles would not be limited by the surface of particles and would be much less affected by losses of SOA-precursors on the chamber walls.

Therefore, experiments with variations of [NO\textsubscript{x}] were also conducted in the presence of ammonium sulfate ((NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4}) seed particles (average seed mass and surface were approximately 9 ± 1 µg m\textsuperscript{-3} and 1.3 × 10\textsuperscript{3} m\textsuperscript{2} m\textsuperscript{-3}, respectively). These experiments were carried out in the same manner as previous experiments; by adjusting [OH] during steady state to the same level as during the NO\textsubscript{x} free experiments. Figure 10 presents a comparison of SOA yield before and after [OH] adjustment for seeded NO\textsubscript{x} experiments (seed mass is subtracted from the total particle mass to determine the organic mass used in the yield calculation). The results from the seeded experiments showed the same general features as the unseeded experiments. Without adjustment of [OH], yields increased with increasing NO\textsubscript{x} at low levels and were slightly suppressed with further increasing NO\textsubscript{x}. The most evident difference here to the experiments without seed is the fairly high yield at high [NO\textsubscript{x}] (Fig. 10). Even after [OH] was adjusted, the decrease in SOA yield was not as significant as in the experiments without seed particles. This
suggests that in the absence of seed particles, the accumulation of mass was indeed limited by low particle surface caused by the \( \text{NO}_x \) induced suppression of NPF.

In the presence of seed particles and at constant \([\text{OH}]\), the decrease in yield was only moderate. This indicates that other \( \text{NO}_x \) impacts such as formation of organic nitrates were moderate as well. The difference between yields determined with and without addition of seed particles indicates that at very small particle surface, our correction procedure underestimates wall losses of precursors. This might be due to either possible differences in uptake of the ELVOC by particles (mainly organic particles versus ammonium sulfate particles), or the differences in the size of particles. However, the real reason for this underestimation is not known yet.

Our correction procedure may involve uncertainties and errors. Nevertheless, it had to be applied; otherwise the \( \text{NO}_x \) dependence would have been overestimated. \( \text{NO}_x \) suppresses NPF and thereby limits mass formation in the absence of seed particles. As both, the impacts of wall losses and impacts of suppressed NPF on SOA mass formation are certainly diminished in the presence of seed, we assume that the experiments with seed particles give the most reliable results on direct \( \text{NO}_x \) impacts on SOA mass formation from \( \beta \)-pinene.

4. Summary and Conclusions

We investigated the effect of \( \text{NO}_x \) on SOA formation from \( \beta \)-pinene photooxidation under low \( \text{NO}_x \) and high \( \text{NO}_x \) conditions and found a very similar behaviour as that observed in other studies (Pandis et al., 1991; Zhang et al., 1992; Presto et al., 2005; Kroll et al., 2006; Camredon et al., 2007; Pathak et al., 2007; Chan et al., 2010; Hoyle et al., 2011; Loza et al., 2014). At low \( \text{NO}_x \) levels SOA yields increased with increasing \( \text{NO}_x \) and then decreased at higher \( \text{NO}_x \) concentrations. The increase of yield at low \([\text{NO}_x]\) was caused by the \( \text{NO}_x \) induced increase of
[OH]. The decrease of yield at higher NO\textsubscript{x} levels was mainly a consequence of NPF suppression and thereby decreasing particle condensational sink with increasing NO\textsubscript{x}. Eliminating the impacts of NO\textsubscript{x} on NPF and on [OH], showed that the impacts of NO\textsubscript{x} on mass formation were only moderate. Even at the highest NO\textsubscript{x} level ([NO\textsubscript{x}]\textsubscript{ss} ~ 86 ppb, [BVOC]/[NO\textsubscript{x}]\textsubscript{ss} ~1.1) suppression of mass yield was only 20 – 30 %. VOC/NO\textsubscript{x} ratios in typical urban air are often much higher than the [BVOC]/[NO\textsubscript{x}] range scanned here (Cai et al., 2011; Pollack et al., 2013; Zou et al., 2015). Therefore dependent on the conditions, impacts of NO\textsubscript{x} on SOA formation in the real atmosphere may be far less than 20-30 %.

Our study also showed that SOA yield is dependent on OH concentration. Although the exact mechanism for this [OH] dependence is still unknown, our results show that besides yield dependencies on the amount of pre-existing matter and effects like partitioning there is also a dependence on reaction conditions, in particular on oxidant levels. Although SOA yields measured in laboratory chambers may not be indicative of the yields in the real atmosphere, their variations as a consequence of different conditions could provide a more comprehensive description of SOA in global and climate model.

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References


Ng, N. L., Chhabra, P. S., Chan, A. W. H., Surratt, J. D., Kroll, J. H., Kwan, A. J., McCabe, D. C., Wennberg, P. O., Sorooshian, A., Murphy, S. M., Dalleska, N. F., Flagan, R. C., and


Table 1. Experimental conditions and results for β-pinene/NO\textsubscript{x} photooxidation experiments.

<table>
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<th>β-pinene reacted (ppb)</th>
<th>[NO\textsubscript{x}]\textsubscript{0} (ppb)\textsuperscript{a}</th>
<th>[NO\textsubscript{x}]\textsubscript{ss} (ppb)</th>
<th>[OH]/10\textsuperscript{7} (cm\textsuperscript{-3})</th>
<th>PM\textsubscript{max} (µg m\textsuperscript{-3})\textsuperscript{c}</th>
<th>PM\textsubscript{ss} (µg m\textsuperscript{-3})\textsuperscript{d}</th>
<th>J\textsubscript{3} (cm\textsuperscript{-3}s\textsuperscript{-1})\textsuperscript{e}</th>
<th>[BVOC]\textsubscript{0}/[NO\textsubscript{x}]\textsubscript{0} (ppbC ppb\textsuperscript{-1})</th>
<th>SOA yield\textsuperscript{f} (%)</th>
<th>SOA yield\textsuperscript{g} (%)</th>
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\textsuperscript{a}Initial NO\textsubscript{x} concentration before OH production
\textsuperscript{b}NO\textsubscript{x} concentration during steady state, in cases without NO\textsubscript{x} addition we found increasing NO\textsubscript{x} which is assumed to be produced from residual HNO\textsubscript{3} that is photolyzed by the TUV lamp.
\textsuperscript{c}Maximum formed particle mass concentration, assuming an SOA density of 1.2 g cm\textsuperscript{-3}. These values have been adjusted for wall losses and losses on particles.
\textsuperscript{d}Particle mass concentration during steady state, assuming an SOA density of 1.2 g cm\textsuperscript{-3}. These values have been adjusted for wall losses and losses on particles.
\textsuperscript{e}Rates of new particle formation for particles greater than 3 nm
\textsuperscript{f}SOA yields determined from PM\textsubscript{max}
\textsuperscript{g}SOA yields determined from PM\textsubscript{ss}
Table 2. Experimental conditions and results for β-pinene photooxidation experiments at two OH levels.

<table>
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<th>J(O¹D)/10^3 (s⁻¹)ᵃ</th>
<th>Initial β-pinene (ppb)</th>
<th>β-pinene reacted (ppb)</th>
<th>[OH]/10^7 (cm⁻³)</th>
<th>PMₘₐₓ (µg m⁻³)ᵇ</th>
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ᵃ J(O¹D) was varied by altering the TUV gap
ᵇ Maximum formed particle mass concentration, assuming an SOA density of 1.2 g cm⁻³. These values have been adjusted for wall losses and losses on particles.
Figure 1. Measured SOA yield from $\text{PM}_{\text{max}}$ (black circles) and rates of new particle formation (blue squares) for the $\beta$-pinene photooxidation as a function of the ratio of the initial hydrocarbon to the initial NO$_x$ concentration and as a function of the initial NO$_x$ concentration. Each point corresponds to one experiment. The errors in nucleation rate and [NO$_x$] were estimated to be $\pm$ 10%. The error in SOA yield was estimated from error propagation using the sum of the systematic error, correction procedure error and error in BVOC data. Note that the horizontal error bars is associated with the BVOC/NO$_x$ axis.

Figure 2. Measured SOA yield from $\text{PM}_{\text{max}}$ and from steady state PM (black and blue circles respectively) and measured OH concentration (red squares) as a function of initial ([NO$_x$]$_0$) and steady state ([NO$_x$]$_{ss}$) NO$_x$ concentrations. The errors in [OH] and [NO$_x$] were estimated to be $\pm$ 20% and $\pm$ 10% respectively. The error in SOA yield was estimated from error propagation using the sum of the systematic error, correction procedure error and error in BVOC data.
Figure 3. Total aerosol mass concentration as a function of the amount of reacted β-pinene under low [OH] (blue open circles) and high [OH] (red open squares) conditions. The SOA yield was estimated from the aerosol mass linear regression slope as a function of consumed β-pinene which resulted in approximately 20 ± 1 % and 31 ± 3 % for low and high OH conditions respectively. The error in [consumed β-pinene] was estimated to be ± 10 % and the error in particle mass was estimated from the sum of the systematic error and the correction procedure error.

Figure 4. Comparison of [OH] before (black squares) and after (blue squares) adjusting OH concentration during steady state in NOx experiments. The errors in [OH] and [NOx] were estimated to be ± 20 % and ± 10 % respectively.
Figure 5. Comparison of SOA yield before (black circles) and after (blue circles) adjusting OH concentration during steady state in NO$_x$ experiments. The error in [NO$_x$] was estimated to be ± 10 % and the error in SOA yield was estimated from error propagation using the sum of the systematic error, correction procedure error and error in BVOC data.

Figure 6. Comparison of [OH] before (black squares) and after (blue squares) adjusting OH concentration during steady state in NO$_x$ experiments performed under lower [NO]/[NO$_2$] ratio. The errors in [OH] and [NO$_x$] were estimated to be ± 20 % and ± 10 % respectively.
Figure 7. Comparison of SOA yield before (black circles) and after (blue circles) adjusting OH concentration during steady state in NO\textsubscript{x} experiments performed under lower [NO]/[NO\textsubscript{2}] ratio. The error in [NO\textsubscript{x}] was estimated to be ± 10 % and the error in SOA yield was estimated from error propagation using the sum of the systematic error, correction procedure error and error in BVOC data.

Figure 8. Measured SOA yield (black diamonds) and measured OH concentration (blue squares) as a function of initial NO\textsubscript{x} concentration for α-pinene/NO\textsubscript{x} photooxidation experiments. Note that, due to the lower α-pinene concentrations, the x-axis is not directly comparable to the x-axes of Fig. 5 and Fig. 7. In sense of BVOC/NO\textsubscript{x} ratios, the NO\textsubscript{x} range scanned here is ~ 3 times higher. The errors in [OH] and [NO\textsubscript{x}] were estimated to be ± 15 % and ± 10 % respectively. The error in SOA yield was estimated from error propagation using the sum of the systematic error, correction procedure error and error in BVOC data.
Figure 9. Comparison of measured particle mass and particle number concentration after adjusting [OH] as a function of [NO\textsubscript{x}] during steady state in the absence of seed aerosol. The error in particle number concentration was estimated to be ±10%.

Figure 10. Comparison of SOA yield before (black circles) and after (blue circles) adjusting OH concentration during steady state in NO\textsubscript{x} experiments performed in the presence of seed aerosol. The error in [NO\textsubscript{x}] was estimated to be ±10% and the error in SOA yield was estimated from error propagation using the sum of the systematic error, correction procedure error and error in BVOC data.